

19th Edition

HARRISON'S™

PRINCIPLES OF
INTERNAL
MEDICINE

KASPER

FAUCI

HAUSER

LONGO

JAMESON

LOSCALZO

Mc
Graw
Hill
Education

PATIENT MANAGEMENT ALGORITHMS

AGING

- FIG. 11-16** Algorithm depicting assessment and management of delirium in hospitalized older patients 79
- FIG. 11-17** Algorithm depicting assessment and management of falls in older patients 80

ALLERGY, IMMUNOLOGY, RHEUMATOLOGY

- FIG. 376-4** The diagnosis and management of rhinitis 2123
- FIG. 378-2** The diagnosis and initial therapy of systemic lupus erythematosus 2128
- FIG. 383-1** Treatment for Sjögren's syndrome 2168
- FIG. 385-1** Approach to the patient with a suspected diagnosis of vasculitis 2181
- FIG. 390-8** Patient management for sarcoidosis 2210
- FIG. 390-9** Management of acute sarcoidosis 2211
- FIG. 390-10** Management of chronic sarcoidosis 2211
- FIG. 393-1** The diagnosis of musculoskeletal complaints 2217
- FIG. 393-2** The consideration of the most common musculoskeletal conditions 2218
- FIG. 393-6** Approach to synovial fluid aspiration and analysis 2224

ALTERATIONS IN CIRCULATORY AND RESPIRATORY FUNCTIONS

- FIG. 50-1** Clinical conditions in which a decrease in cardiac output and a systemic vascular resistance cause arterial underfilling. 251
- FIG. 51e-5** Differential diagnosis of a holosystolic murmur 51e-5

CARDIOLOGY

- FIG. 9-1** Cardiac risk assessment and stratification in patients undergoing noncardiac surgery 52
- FIG. 51e-9** Strategy for evaluating heart murmurs 51e-8
- FIG. 264-1** Approach to the evaluation of a heart murmur 1441
- FIG. 276-9** Treatment algorithm for patients presenting with hemodynamically stable paroxysmal supraventricular tachycardia 1483
- FIG. 283-2** Management strategy for patients with aortic stenosis 1533
- FIG. 283-6** Management of patients with aortic regurgitation 1537
- FIG. 284-1** Management of rheumatic mitral stenosis 1541
- FIG. 284-4** Management of mitral regurgitation 1545
- FIG. 285-1** Management of tricuspid regurgitation 1549
- FIG. 287-18** Treatment algorithm for hypertrophic cardiomyopathy 1570

- FIG. 293-2** Evaluation of the patient with known or suspected ischemic heart disease 1583
- FIG. 293-3** Management of the patient with ischemic heart disease 1591
- FIG. 294-3** Algorithm for evaluation and management of patients with suspected acute coronary syndrome 1595
- FIG. 295-5** The assessment of need for implantation of a cardioverter-defibrillator. 1609

CLINICAL GENETICS

- FIG. 84-2** Approach to genetic testing 448
- FIG. 85e-6** Clinical and laboratory investigation of a suspected mitochondrial DNA (mtDNA) disorder. 85e-7
- FIG. 101e-6** Genetic testing in a family with cancer predisposition 101e-8

CLINICAL NUTRITION

- FIG. 98e-1** Decision-making for the implementation of specialized nutritional support (SNS) 98e-2

CORONARY AND PERIPHERAL VASCULAR DISEASE

- FIG. 295-4** Reperfusion therapy for patients with T-segment elevation myocardial infarction (STEMI) 1604
- FIG. 296e-4** Algorithm for patients requiring revascularization: factors to consider when choosing between bare metal stents, drug-eluting stent, or CABG 296e-3

DISORDERS OF THE URINARY TRACT

- FIG. 337-2** Typical algorithm for early posttransplant care of a kidney recipient 1830

EMERGENCY AND CRITICAL CARE

- FIG. 263e-1** General guidelines for treatment of radiation casualties 263e-4
- FIG. 263e-2** Evacuation in a multicasualties radiologic event 263e-5
- FIG. 321-2** Approach to the patient in shock 1731
- FIG. 322-5** Algorithm for the initial management of acute respiratory distress syndrome 1739
- FIG. 326-2** Emergency management of patients with cardiogenic shock, acute pulmonary edema, or both 1761
- FIG. 327-3A** Cardiac arrest management: ventricular fibrillation 1769

FIG. 327-3B Cardiac arrest management: bradyarrhythmia/
asystole 1769

ENDOCRINOLOGY AND METABOLISM

FIG. 68-2 The evaluation and differential diagnosis
of hirsutism 334

FIG. 402-1 Management of adult growth hormone (GH)
deficiency 2260

FIG. 403-3 Management of prolactinoma 2268

FIG. 403-5 Management of acromegaly 2270

FIG. 403-6 Management of Cushing's syndrome 2273

FIG. 403-7 Management of a nonfunctioning
pituitary mass 2274

FIG. 405-13 Approach to the patient with a thyroid
nodule 2308

FIG. 406-10 Management of the patient with suspected
Cushing's syndrome 2316

FIG. 406-12 Diagnostic flowchart for evaluating patients
with possible mineralocorticoid excess 2320

FIG. 406-13 Management of the patient with an incidentally
discovered adrenal mass 2322

FIG. 406-16 Diagnostic flowchart for evaluating patients
with suspected adrenal insufficiency 2327

FIG. 411-5 Evaluation of gynecomastia 2367

FIG. 411-6 Evaluation of hypogonadism 2367

FIG. 416-1 Algorithm for the treatment of obesity 2395

FIG. 418-2 Essential elements in comprehensive
care of type 2 diabetes 2413

FIG. 418-3 Glycemic management of type 2 diabetes 2416

FIG. 419-4 Screening for microalbuminuria 2425

FIG. 420-2 Hypoglycemia-associated autonomic failure
in insulin-deficient diabetes 2432

FIG. 424-6 The evaluation of patients with hypercalcemia 2479

FIG. 428-3 Algorithm for screening for *HFE*-associated
hemochromatosis 2518

GASTROENTEROLOGY AND HEPATOLOGY

FIG. 53-2 Approach to the patient with dysphagia 257

FIG. 55-2 Management of acute diarrhea 267

FIG. 55-3 Management of chronic diarrhea 271

FIG. 55-4 Management of constipation 273

FIG. 57-1 Management of acute upper gastrointestinal
bleeding 278

FIG. 57-2 Algorithm for patients with acute lower
gastrointestinal bleeding 279

FIG. 58-1 Evaluation of the patient with jaundice 281

FIG. 348-12 Overview of new-onset dyspepsia 1924

FIG. 357-1 Algorithm for evaluation of abnormal
liver tests 1994

FIG. 358-1 Algorithm for the evaluation of chronically
abnormal liver tests 1996

FIG. 363-2 Treatment algorithm for alcoholic hepatitis 2054

FIG. 365-3 Management of recurrent variceal hemorrhage 2064

FIG. 365-5 Treatment of refractory ascites 2065

FIG. 370-1 A stepwise diagnostic approach to the patient
with suspected chronic pancreatitis 2088

HEMATOLOGY AND ONCOLOGY

FIG. 77-17 The physiologic classification of anemia 398

FIG. 77-18 An approach to diagnosing patients with
polycythemia 400

FIG. 104-2 Diagnosis and treatment of fever
and neutropenia 490

FIG. 106-1 Evaluation of a patient with cervical
adenopathy 503

FIG. 107-3 Algorithm for management of non-small-cell
lung cancer 512

FIG. 107-5 Algorithm for management of small-cell lung
cancer 515

FIG. 107-6 Approach to a solitary pulmonary nodule 517

FIG. 107-7 Management of recurrent small-cell lung
cancer 522

FIG. 107-8 Approach to first-line therapy in a patient
with stage IV non-small-cell lung cancer 523

FIG. 108-1 Approach to a palpable breast mass 525

FIG. 108-2 The "triple diagnosis" technique in suspected
breast cancer 525

FIG. 108-3 Management of a breast cyst 525

FIG. 108-4 Approaches to abnormalities detected by
mammogram 526

FIG. 111-2 Treatment approach to patients with
hepatocellular carcinoma 548

FIG. 120e-2 Treatment algorithm for adenocarcinoma and
poorly differentiated adenocarcinoma of
unknown primary 120e-3

FIG. 120e-3 Treatment algorithm for squamous cell
carcinoma of unknown primary 120e-4

FIG. 127-3 Pathophysiology of sickle cell crisis 634

FIG. 132-2 The therapy of newly diagnosed acute
myeloid leukemia 684

FIG. 137-1 Algorithm for the diagnosis of amyloidosis
and determination of type 720

FIG. 140-2 Evaluating the thrombocytopenic patient 726

FIG. 143-1 Classification of antithrombotic drugs 745

FIG. 331-2 Management of cancer patients with back pain 1791

FIG. 331-4 Management of patients at high risk for the
tumor lysis syndrome 1796

INFECTIOUS DISEASES

FIG. 44-2 Diagnosis and treatment of acute pharyngitis 232

FIG. 152e-1 Syndromic approach to differential diagnosis
of suspected infection in a veteran who has
returned from a foreign war 152e-7

FIG. 155-4 Diagnostic use of transesophageal and transtracheal
echocardiography in infective endocarditis 821

FIG. 159-3	Management of patients with intraabdominal abscesses using percutaneous drainage	850
FIG. 160-1	Approach to patients with community-acquired infectious diarrhea or bacterial food poisoning	854
FIG. 162-4	Diagnostic approach to urinary tract infection	865
FIG. 177-1	Clinical and pathologic progression of tetanus	985
FIG. 188-1	Schematic of the relationships between colonization with <i>Helicobacter pylori</i> and diseases of the upper gastrointestinal tract among persons in developed countries	1039
FIG. 188-2	Management of <i>Helicobacter pylori</i> infection	1041
FIG. 206-4	Interpretation of results from syphilis enzyme immunoassays used for screening	1137
FIG. 209-3	Treatment of relapsing fever	1148
FIG. 210-2	Treatment of the various early or late manifestations of Lyme borreliosis	1153
FIG. 226-32	Acute HIV infection	1249
FIG. 226-36	Evaluation of diarrhea in patients with HIV infection	1257
FIG. 232-5	Rabies postexposure prophylaxis	1303

NEPHROLOGY

FIG. 61-1	Approach to the patient with azotemia	290
FIG. 61-2	Approach to the patient with hematuria	292
FIG. 61-3	Approach to the patient with proteinuria	293
FIG. 61-4	Approach to the patient with polyuria	294
FIG. 63-5	The diagnostic approach to hyponatremia	299
FIG. 63-6	The diagnostic approach to hypernatremia	303
FIG. 63-7	The diagnostic approach to hypokalemia	308
FIG. 63-8	The diagnostic approach to hyperkalemia	311
FIG. 67-3	The evaluation and management of patients with erectile dysfunction	327
FIG. 337-2	A typical algorithm for early posttransplant care of the kidney recipient	1830
FIG. 340-1	Algorithm for the treatment of allergic and other immune-mediated acute interstitial nephritis	1857
FIG. 343-1	Diagnostic approach for urinary tract obstruction in unexplained renal failure	1873

NEUROLOGY AND PSYCHIATRY

FIG. 164-1	The management of patients with suspected central nervous system infection	884
-------------------	--	-----

FIG. 164-2	The pathophysiology of the neurologic complications of bacterial meningitis	886
FIG. 445-2	Evaluation of the adult patient with a seizure	2549
FIG. 445-3	Pharmacologic treatment of generalized tonic-clonic status epilepticus in adults	2558
FIG. 446-1	Medical management of stroke and transient ischemic attack	2560
FIG. 449-7	Treatment algorithm for the management of Parkinson's disease	2619
FIG. 458-4	Therapeutic decision-making for multiple sclerosis	2671
FIG. 459-1	Approach to the evaluation of peripheral neuropathies	2675
FIG. 461-2	The management of myasthenia gravis	2704
FIG. 462e-1	Diagnostic evaluation of intermittent weakness	462e-1
FIG. 462e-2	Diagnostic evaluation of persistent weakness	462e-2
FIG. 466-1	Medical management of major depressive disorder	2716

PULMONOLOGY

FIG. 47e-2	The evaluation of the patient with dyspnea	47e-3
FIG. 48-2	Algorithm for the evaluation of hemoptysis	246
FIG. 300-3	How to decide whether diagnostic imaging is needed in suspected pulmonary thromboembolism and deep venous thrombosis	1632
FIG. 300-6	Imaging tests to diagnose deep venous thrombosis and pulmonary embolism	1634
FIG. 300-7	Acute management of pulmonary thromboembolism	1634
FIG. 316-1	Approach to the diagnosis of pleural effusions	1717

SYSTEMIC CONDITIONS

FIG. 10-2	Common and uncommon clinical courses in the last days of terminally ill patients	68
FIG. 26-1	Structured approach to patients with fever of unknown origin	139
FIG. 30-3	The initial workup of a patient with weakness	156
FIG. 43-2	An algorithm for the approach to hearing loss	218
FIG. 69-2	Algorithm for evaluation of amenorrhea	337
FIG. 75-1	An algorithm for the diagnosis of a patient with photosensitivity	390

19th Edition



HARRISON'STM
P R I N C I P L E S O F
**INTERNAL
MEDICINE**

EDITORS OF PREVIOUS EDITIONS

T. R. Harrison

Editor-in-Chief, Editions 1, 2, 3, 4, 5

W. R. Resnick

Editor, Editions 1, 2, 3, 4, 5

M. M. Wintrobe

Editor, Editions 1, 2, 3, 4, 5
Editor-in-Chief, Editions 6, 7

G. W. Thorn

Editor, Editions 1, 2, 3, 4, 5, 6, 7
Editor-in-Chief, Edition 8

R. D. Adams

Editor, Editions 2, 3, 4, 5, 6, 7, 8, 9, 10

P. B. Beeson

Editor, Editions 1, 2

I. L. Bennett, Jr.

Editor, Editions 3, 4, 5, 6

E. Braunwald

Editor, Editions 6, 7, 8, 9, 10,
12, 13, 14, 16, 17
Editor-in-Chief, Editions 11, 15

K. J. Isselbacher

Editor, Editions 6, 7, 8, 10, 11, 12, 14
Editor-in-Chief, Editions 9, 13

R. G. Petersdorf

Editor, Editions 6, 7, 8, 9, 11, 12
Editor-in-Chief, Edition 10

J. D. Wilson

Editor, Editions 9, 10, 11, 13, 14
Editor-in-Chief, Edition 12

J. B. Martin

Editor, Editions 10, 11, 12, 13, 14

A. S. Fauci

Editor, Editions 11, 12, 13, 15, 16, 18
Editor-in-Chief, Editions 14, 17

R. Root

Editor, Edition 12

D. L. Kasper

Editor, Editions 13, 14, 15, 17, 18
Editor-in-Chief, Editions 16, 19

S. L. Hauser

Editor, Editions 14, 15, 16, 17, 18

D. L. Longo

Editor, Editions 14, 15, 16, 17
Editor-in-Chief, Edition 18

J. L. Jameson

Editor, Editions 15, 16, 17, 18

J. Loscalzo

Editor, Editions 17, 18

19th Edition



HARRISON'STM
P R I N C I P L E S O F
**INTERNAL
MEDICINE**

EDITORS

Dennis L. Kasper, MD

William Ellery Channing Professor of Medicine, Professor of Microbiology and Immunobiology, Department of Microbiology and Immunobiology, Harvard Medical School; Division of Infectious Diseases, Brigham and Women's Hospital
Boston, Massachusetts

Anthony S. Fauci, MD

Chief, Laboratory of Immunoregulation;
Director, National Institute of Allergy and Infectious Diseases,
National Institutes of Health
Bethesda, Maryland

Stephen L. Hauser, MD

Robert A. Fishman Distinguished Professor and Chairman,
Department of Neurology, University of California, San Francisco
San Francisco, California

Dan L. Longo, MD

Professor of Medicine, Harvard Medical School; Senior Physician,
Brigham and Women's Hospital; Deputy Editor, *New England
Journal of Medicine*, Boston, Massachusetts

J. Larry Jameson, MD, PhD

Robert G. Dunlop Professor of Medicine; Dean, Perelman School
of Medicine at the University of Pennsylvania; Executive Vice
President, University of Pennsylvania for the Health System
Philadelphia, Pennsylvania

Joseph Loscalzo, MD, PhD

Hersey Professor of the Theory and Practice of Medicine,
Harvard Medical School; Chairman, Department of Medicine, and
Physician-in-Chief, Brigham and Women's Hospital
Boston, Massachusetts



New York Chicago San Francisco Athens London Madrid Mexico City
Milan New Delhi Singapore Sydney Toronto

Copyright © 2015 by McGraw-Hill Education. Previous editions copyright © 2012, 2008, 2005, 2001, 1998, 1994, 1991, 1987, 1983, 1980, 1977, 1974, 1970, 1966, 1962, 1958 by The McGraw-Hill Companies, Inc. All rights reserved. Except as permitted under the United States Copyright Act of 1976, no part of this publication may be reproduced or distributed in any form or by any means, or stored in a database or retrieval system, without the prior written permission of the publisher, with the exception that the program listings may be entered, stored, and executed in a computer system, but they may not be reproduced for publication.

ISBN: 978-0-07-180216-1

MHID: 0-07-180216-9

The material in this eBook also appears in the print version of this title: ISBN: 978-0-07-180215-4,
MHID: 0-07-180215-0.

eBook conversion by codeMantra
Version 1.0

All trademarks are trademarks of their respective owners. Rather than put a trademark symbol after every occurrence of a trademarked name, we use names in an editorial fashion only, and to the benefit of the trademark owner, with no intention of infringement of the trademark. Where such designations appear in this book, they have been printed with initial caps.

McGraw-Hill Education eBooks are available at special quantity discounts to use as premiums and sales promotions or for use in corporate training programs. To contact a representative, please visit the Contact Us page at www.mhprofessional.com.

Note: Dr. Fauci's work as editor and author was performed outside the scope of his employment as a U.S. government employee. This work represents his personal and professional views and not necessarily those of the U.S. government.

TERMS OF USE

This is a copyrighted work and McGraw-Hill Education and its licensors reserve all rights in and to the work. Use of this work is subject to these terms. Except as permitted under the Copyright Act of 1976 and the right to store and retrieve one copy of the work, you may not decompile, disassemble, reverse engineer, reproduce, modify, create derivative works based upon, transmit, distribute, disseminate, sell, publish or sublicense the work or any part of it without McGraw-Hill Education's prior consent. You may use the work for your own noncommercial and personal use; any other use of the work is strictly prohibited. Your right to use the work may be terminated if you fail to comply with these terms.

THE WORK IS PROVIDED "AS IS." McGRAW-HILL EDUCATION AND ITS LICENSORS MAKE NO GUARANTEES OR WARRANTIES AS TO THE ACCURACY, ADEQUACY OR COMPLETENESS OF OR RESULTS TO BE OBTAINED FROM USING THE WORK, INCLUDING ANY INFORMATION THAT CAN BE ACCESSED THROUGH THE WORK VIA HYPERLINK OR OTHERWISE, AND EXPRESSLY DISCLAIM ANY WARRANTY, EXPRESS OR IMPLIED, INCLUDING BUT NOT LIMITED TO IMPLIED WARRANTIES OF MERCHANTABILITY OR FITNESS FOR A PARTICULAR PURPOSE. McGraw-Hill Education and its licensors do not warrant or guarantee that the functions contained in the work will meet your requirements or that its operation will be uninterrupted or error free. Neither McGraw-Hill Education nor its licensors shall be liable to you or anyone else for any inaccuracy, error or omission, regardless of cause, in the work or for any damages resulting therefrom. McGraw-Hill Education has no responsibility for the content of any information accessed through the work. Under no circumstances shall McGraw-Hill Education and/or its licensors be liable for any indirect, incidental, special, punitive, consequential or similar damages that result from the use of or inability to use the work, even if any of them has been advised of the possibility of such damages. This limitation of liability shall apply to any claim or cause whatsoever whether such claim or cause arises in contract, tort or otherwise.

Material from the disk that accompanies the printed version of this eBook may be obtained from McGraw-Hill Professional's MediaCenter at <http://www.mhprofessional.com/mediacenter/hpim19>

Some material may require a desktop or laptop computer for full access.

NOTICE

Medicine is an ever-changing science. As new research and clinical experience broaden our knowledge, changes in treatment and drug therapy are required. The authors and the publisher of this work have checked with sources believed to be reliable in their efforts to provide information that is complete and generally in accord with the standards accepted at the time of publication. However, in view of the possibility of human error or changes in medical sciences, neither the authors nor the publisher nor any other party who has been involved in the preparation or publication of this work warrants that the information contained herein is in every respect accurate or complete, and they disclaim all responsibility for any errors or omissions or for the results obtained from use of the information contained in this work. Readers are encouraged to confirm the information contained herein with other sources. For example and in particular, readers are advised to check the product information sheet included in the package of each drug they plan to administer to be certain that the information contained in this work is accurate and that changes have not been made in the recommended dose or in the contraindications for administration. This recommendation is of particular importance in connection with new or infrequently used drugs.

COVER ILLUSTRATION

Budding HIV particle, computer artwork (*Credit: Animate4.com Ltd. / Science Source*)

This page intentionally left blank

CONTRIBUTORS xix
PREFACE xxxvii

PART 1 General Considerations in Clinical Medicine

1 The Practice of Medicine 1
The Editors

2 Global Issues in Medicine 7
Paul Farmer, Joseph Rhatigan

3 Decision-Making in Clinical Medicine 18
Daniel B. Mark, John B. Wong

4 Screening and Prevention of Disease 26
Katrina Armstrong, Gary J. Martin

5 Principles of Clinical Pharmacology 31
Dan M. Roden

 **6e Women's Health** 45
Andrea Dunaif

 **7e Men's Health** 45
Shalender Bhasin, Shehzad Basaria

8 Medical Disorders During Pregnancy 45
Robert L. Barbieri, John T. Repke

9 Medical Evaluation of the Surgical Patient 51
Wei C. Lau, Kim A. Eagle

10 Palliative and End-of-Life Care 55
Ezekiel J. Emanuel

11 Clinical Problems of Aging 70
Luigi Ferrucci, Stephanie Studenski

 **12e The Safety and Quality of Health Care** 85
David W. Bates

 **13e Primary Care in Low- and Middle-Income Countries** 85
Tim Evans, Kumanan Rasanathan

 **14e Complementary, Alternative, and Integrative Health Practices** 86
Josephine P. Briggs

 **15e The Economics of Medical Care** 86
Joseph P. Newhouse

 **16e Racial and Ethnic Disparities in Health Care** 86
Joseph R. Betancourt, Alexander R. Green

 **17e Ethical Issues in Clinical Medicine** 86
Bernard Lo, Christine Grady

PART 2 Cardinal Manifestations and Presentation of Diseases

SECTION 1 PAIN

18 Pain: Pathophysiology and Management 87
James P. Rathmell, Howard L. Fields

19 Chest Discomfort 95
David A. Morrow

20 Abdominal Pain 103
Danny O. Jacobs, William Silen


21 Headache 107
Peter J. Goadsby, Neil H. Raskin

22 Back and Neck Pain 111
John W. Engstrom, Richard A. Deyo

SECTION 2 ALTERATIONS IN BODY TEMPERATURE

23 Fever 123
Charles A. Dinarello, Reuven Porat

24 Fever and Rash 127
Elaine T. Kaye, Kenneth M. Kaye

 **25e Atlas of Rashes Associated with Fever** 135
Kenneth M. Kaye, Elaine T. Kaye

26 Fever of Unknown Origin 135
Chantal P. Bleeker-Rovers, Jos W. M. van der Meer

SECTION 3 NERVOUS SYSTEM DYSFUNCTION

27 Syncope 142
Roy Freeman

28 Dizziness and Vertigo 148
Mark F. Walker, Robert B. Daroff

29 Fatigue 151
Jeffrey M. Gelfand, Vanja C. Douglas

30 Neurologic Causes of Weakness and Paralysis 154
Michael J. Aminoff

31 Numbness, Tingling, and Sensory Loss 158
Michael J. Aminoff

32 Gait and Balance Disorders 162
Lewis Sudarsky

 **33e Video Library of Gait Disorders** 165
Gail Kang, Nicholas B. Galifianakis, Michael D. Geschwind

34 Confusion and Delirium 166
S. Andrew Josephson, Bruce L. Miller

35 Dementia 170
William W. Seeley, Bruce L. Miller


36 Aphasia, Memory Loss, and Other Focal Cerebral Disorders 176
M.-Marsel Mesulam


 **37e Primary Progressive Aphasia, Memory Loss, and Other Focal Cerebral Disorders** 184
Maria Luisa Gorno-Tempini, Jennifer Ogar, Joel Kramer, Bruce L. Miller, Gil Rabinovici, Maria Carmela Tartaglia

38 Sleep Disorders 184
Charles A. Czeisler, Thomas E. Scammell, Clifford B. Saper

SECTION 4 DISORDERS OF EYES, EARS, NOSE, AND THROAT

39 Disorders of the Eye 195
Jonathan C. Horton

 **40e Use of the Hand-Held Ophthalmoscope** 211
Homayoun Tabandeh, Morton F. Goldberg

 **41e Video Library of Neuro-Ophthalmology** 211
Shirley H. Wray

42 Disorders of Smell and Taste 211
Richard L. Doty, Steven M. Bromley

43 Disorders of Hearing 217
Anil K. Lalwani

44 Sore Throat, Earache, and Upper Respiratory Symptoms 225
Michael A. Rubin, Larry C. Ford, Ralph Gonzales

- 45 Oral Manifestations of Disease** 235
Samuel C. Durso
- 46e Atlas of Oral Manifestations of Disease** 242
Samuel C. Durso, Janet A. Yellowitz

SECTION 5 ALTERATIONS IN CIRCULATORY AND RESPIRATORY FUNCTIONS

- 47e Dyspnea** 243
Richard M. Schwartzstein
- 48 Cough and Hemoptysis** 243
Patricia A. Kritek, Christopher H. Fanta
- 49 Hypoxia and Cyanosis** 247
Joseph Loscalzo
- 50 Edema** 250
Eugene Braunwald, Joseph Loscalzo
- 51e Approach to the Patient with a Heart Murmur** 253
Patrick T. O'Gara, Joseph Loscalzo
- 52 Palpitations** 254
Joseph Loscalzo

SECTION 6 ALTERATIONS IN GASTROINTESTINAL FUNCTION

- 53 Dysphagia** 254
Ikuo Hirano, Peter J. Kahrilas
- 54 Nausea, Vomiting, and Indigestion** 258
William L. Hasler
- 55 Diarrhea and Constipation** 264
Michael Camilleri, Joseph A. Murray
- 56 Involuntary Weight Loss** 274
Russell G. Robertson, J. Larry Jameson
- 57 Gastrointestinal Bleeding** 276
Loren Laine
- 58 Jaundice** 279
Savio John, Daniel S. Pratt
- 59 Abdominal Swelling and Ascites** 285
Kathleen E. Corey, Lawrence S. Friedman

SECTION 7 ALTERATIONS IN RENAL AND URINARY TRACT FUNCTION

- 60e Dysuria, Bladder Pain, and the Interstitial Cystitis/Bladder Pain Syndrome** 288
John W. Warren
- 61 Azotemia and Urinary Abnormalities** 289
Julie Lin, Bradley M. Denker
- 62e Atlas of Urinary Sediments and Renal Biopsies** 295
Agnes B. Fogo, Eric G. Neilson
- 63 Fluid and Electrolyte Disturbances** 295
David B. Mount
- 64e Fluid and Electrolyte Imbalances and Acid-Base Disturbances: Case Examples** 312
David B. Mount, Thomas D. DuBose, Jr.
- 65 Hypercalcemia and Hypocalcemia** 313
Sundeep Khosla
- 66 Acidosis and Alkalosis** 315
Thomas D. DuBose, Jr.

SECTION 8 ALTERATIONS IN SEXUAL FUNCTION AND REPRODUCTION

- 67 Sexual Dysfunction** 324
Kevin T. McVary

- 68 Hirsutism** 331
David A. Ehrmann
- 69 Menstrual Disorders and Pelvic Pain** 335
Janet E. Hall

SECTION 9 ALTERATIONS IN THE SKIN

- 70 Approach to the Patient with a Skin Disorder** 339
Thomas J. Lawley, Kim B. Yancey
- 71 Eczema, Psoriasis, Cutaneous Infections, Acne, and Other Common Skin Disorders** 344
Leslie P. Lawley, Calvin O. McCall, Thomas J. Lawley
- 72 Skin Manifestations of Internal Disease** 353
Jean L. Bologna, Irwin M. Braverman
- 73 Immunologically Mediated Skin Diseases** 370
Kim B. Yancey, Thomas J. Lawley
- 74 Cutaneous Drug Reactions** 377
Kanade Shinkai, Robert S. Stern, Bruce U. Wintroub
- 75 Photosensitivity and Other Reactions to Light** 385
Alexander G. Marneros, David R. Bickers
- 76e Atlas of Skin Manifestations of Internal Disease** 391
Thomas J. Lawley, Calvin O. McCall, Robert A. Swerlick

SECTION 10 HEMATOLOGIC ALTERATIONS

- 77 Anemia and Polycythemia** 392
John W. Adamson, Dan L. Longo
- 78 Bleeding and Thrombosis** 400
Barbara A. Konkle
- 79 Enlargement of Lymph Nodes and Spleen** 407
Patrick H. Henry, Dan L. Longo
- 80 Disorders of Granulocytes and Monocytes** 413
Steven M. Holland, John I. Gallin
- 81e Atlas of Hematology and Analysis of Peripheral Blood Smears** 424
Dan L. Longo


PART 3 Genes, the Environment, and Disease

- 82 Principles of Human Genetics** 425
J. Larry Jameson, Peter Kopp
- 83e Chromosome Disorders** 445
Nancy B. Spinner, Laura K. Conlin
- 84 The Practice of Genetics in Clinical Medicine** 446
Susan M. Domchek, J. Larry Jameson, Susan Miesfeldt
- 85e Mitochondrial DNA and Heritable Traits and Diseases** 451
Karl Skorecki, Doron Behar
- 86e The Human Microbiome** 451
Jeffrey I. Gordon, Rob Knight
- 87e Network Medicine: Systems Biology in Health and Disease** 451
Joseph Loscalzo




PART 4 Regenerative Medicine

- 88 Stem Cell Biology** 453
Minoru S. H. Ko
-  **89e Hematopoietic Stem Cells** 455
David T. Scadden, Dan L. Longo
-  **90e Applications of Stem Cell Biology in Clinical Medicine** 455
John A. Kessler
-  **91e Gene Therapy in Clinical Medicine** 456
Katherine A. High
-  **92e Tissue Engineering** 456
Anthony Atala

PART 5 Aging




-  **93e World Demography of Aging** 457
Richard M. Suzman, John G. Haaga
-  **94e The Biology of Aging** 457
Rafael de Cabo, David G. Le Couteur





PART 6 Nutrition and Weight Loss

-  **95e Nutrient Requirements and Dietary Assessment** 459
Johanna Dwyer
-  **96e Vitamin and Trace Mineral Deficiency and Excess** 459
Robert M. Russell, Paolo M. Suter
- 97 Malnutrition and Nutritional Assessment** 459
Douglas C. Heimburger
-  **98e Enteral and Parenteral Nutrition Therapy** 465
Bruce R. Bistrain, L. John Hoffer, David F. Driscoll

PART 7 Oncology and Hematology

SECTION 1 NEOPLASTIC DISORDERS

- 99 Approach to the Patient with Cancer** 467
Dan L. Longo
- 100 Prevention and Early Detection of Cancer** 475
Jennifer M. Croswell, Otis W. Brawley, Barnett S. Kramer
-  **101e Cancer Genetics** 483
Pat J. Morin, Jeffrey M. Trent, Francis S. Collins, Bert Vogelstein
-  **102e Cancer Cell Biology** 483
Jeffrey W. Clark, Dan L. Longo
-  **103e Principles of Cancer Treatment** 483
Edward A. Sausville, Dan L. Longo
- 104 Infections in Patients with Cancer** 484
Robert W. Finberg
- 105 Cancer of the Skin** 493
Walter J. Urba, Brendan D. Curti
- 106 Head and Neck Cancer** 502
Everett E. Vokes
- 107 Neoplasms of the Lung** 506
Leora Horn, Christine M. Lovly, David H. Johnson

- 108 Breast Cancer** 523
Marc E. Lippman
- 109 Upper Gastrointestinal Tract Cancers** 532
Robert J. Mayer
- 110 Lower Gastrointestinal Cancers** 537
Robert J. Mayer
- 111 Tumors of the Liver and Biliary Tree** 544
Brian I. Carr
- 112 Pancreatic Cancer** 554
Elizabeth Smyth, David Cunningham
- 113 Endocrine Tumors of the Gastrointestinal Tract and Pancreas** 557
Robert T. Jensen
- 114 Bladder and Renal Cell Carcinomas** 575
Howard I. Scher, Jonathan E. Rosenberg, Robert J. Motzer
- 115 Benign and Malignant Diseases of the Prostate** 579
Howard I. Scher, James A. Eastham
- 116 Testicular Cancer** 588
Robert J. Motzer, Darren R. Feldman, George J. Bosl
- 117 Gynecologic Malignancies** 592
Michael V. Seiden
- 118 Primary and Metastatic Tumors of the Nervous System** 598
Lisa M. DeAngelis, Patrick Y. Wen
-  **119e Soft Tissue and Bone Sarcomas and Bone Metastases** 608
Shreyaskumar R. Patel, Robert S. Benjamin
-  **120e Carcinoma of Unknown Primary** 608
Gauri R. Varadhachary, James L. Abbruzzese
- 121 Paraneoplastic Syndromes: Endocrinologic/Hematologic** 608
J. Larry Jameson, Dan L. Longo
- 122 Paraneoplastic Neurologic Syndromes and Autoimmune Encephalitis** 614
Josep Dalmau, Myrna R. Rosenfeld
-  **123e Thymoma** 619
Dan L. Longo
-  **124e Neoplasia During Pregnancy** 620
Michael F. Greene, Dan L. Longo
- 125 Late Consequences of Cancer and Its Treatment** 620
Carl E. Freter, Dan L. Longo

SECTION 2 HEMATOPOIETIC DISORDERS

- 126 Iron Deficiency and Other Hypoproliferative Anemias** 625
John W. Adamson
- 127 Disorders of Hemoglobin** 631
Edward J. Benz, Jr.
- 128 Megaloblastic Anemias** 640
A. Victor Hoffbrand
- 129 Hemolytic Anemias and Anemia Due to Acute Blood Loss** 649
Lucio Luzzatto
- 130 Bone Marrow Failure Syndromes Including Aplastic Anemia and Myelodysplasia** 662
Neal S. Young
- 131 Polycythemia Vera and Other Myeloproliferative Neoplasms** 672
Jerry L. Spivak
- 132 Acute Myeloid Leukemia** 678
Guido Marcucci, Clara D. Bloomfield

- 133 Chronic Myeloid Leukemia** 687
Hagop Kantarjian, Jorge Cortes
- 134 Malignancies of Lymphoid Cells** 695
Dan L. Longo
- 135e Less Common Hematologic Malignancies** 710
Ayalew Tefferi, Dan L. Longo
- 136 Plasma Cell Disorders** 710
Nikhil C. Munshi, Dan L. Longo, Kenneth C. Anderson
- 137 Amyloidosis** 719
David C. Seldin, John L. Berk
- 138e Transfusion Biology and Therapy** 724
Jeffery S. Dzieczkowski, Kenneth C. Anderson
- 139e Hematopoietic Cell Transplantation** 725
Frederick R. Appelbaum

SECTION 3 DISORDERS OF HEMOSTASIS

- 140 Disorders of Platelets and Vessel Wall** 725
Barbara A. Konkle
- 141 Coagulation Disorders** 732
Valder R. Arruda, Katherine A. High
- 142 Arterial and Venous Thrombosis** 740
Jane E. Freedman, Joseph Loscalzo
- 143 Antiplatelet, Anticoagulant, and Fibrinolytic Drugs** 745
Jeffrey I. Weitz
- 155 Infective Endocarditis** 816
Adolf W. Karchmer
- 156 Infections of the Skin, Muscles, and Soft Tissues** 827
Dennis L. Stevens
- 157 Infectious Arthritis** 833
Lawrence C. Madoff
- 158 Osteomyelitis** 838
Werner Zimmerli
- 159 Intraabdominal Infections and Abscesses** 846
Miriam Baron Barshak, Dennis L. Kasper
- 160 Acute Infectious Diarrheal Diseases and Bacterial Food Poisoning** 852
Regina C. LaRocque, Edward T. Ryan, Stephen B. Calderwood
- 161 Clostridium difficile Infection, Including Pseudomembranous Colitis** 857
Dale N. Gerding, Stuart Johnson
- 162 Urinary Tract Infections, Pyelonephritis, and Prostatitis** 861
Kalpna Gupta, Barbara W. Trautner
- 163 Sexually Transmitted Infections: Overview and Clinical Approach** 869
Jeanne M. Marrasso, King K. Holmes
- 164 Meningitis, Encephalitis, Brain Abscess, and Empyema** 883
Karen L. Roos, Kenneth L. Tyler
- 165 Chronic and Recurrent Meningitis** 906
Walter J. Koroshetz, Avindra Nath

- 166e Infectious Complications of Burns** 911
Lawrence C. Madoff, Florencia Pereyra
- 167e Infectious Complications of Bites** 911
Lawrence C. Madoff, Florencia Pereyra

PART 8 Infectious Diseases

SECTION 1 BASIC CONSIDERATIONS IN INFECTIOUS DISEASES

- 144 Approach to the Patient with an Infectious Disease** 761
Neeraj K. Surana, Dennis L. Kasper
- 145e Molecular Mechanisms of Microbial Pathogenesis** 768
Gerald B. Pier
- 146 Genomics and Infectious Disease** 768
Roby P. Bhattacharyya, Yonatan H. Grad, Deborah T. Hung
- 147 Approach to the Acutely Ill Infected Febrile Patient** 779
Tamar F. Barlam, Dennis L. Kasper
- 148 Immunization Principles and Vaccine Use** 785
Anne Schuchat, Lisa A. Jackson
- 149 Health Recommendations for International Travel** 793
Jay S. Keystone, Phyllis E. Kozarsky
- 150e Laboratory Diagnosis of Infectious Diseases** 802
Alexander J. McAdam, Andrew B. Onderdonk
- 151e Climate Change and Infectious Disease** 803
Aaron S. Bernstein
- 152e Infections in Veterans Returning from Foreign Wars** 803
Andrew W. Artenstein

SECTION 2 CLINICAL SYNDROMES: COMMUNITY-ACQUIRED INFECTIONS

- 153 Pneumonia** 803
Lionel A. Mandell, Richard G. Wunderink
- 154 Lung Abscess** 813
Rebecca M. Baron, Miriam Baron Barshak

SECTION 3 CLINICAL SYNDROMES: HEALTH CARE-ASSOCIATED INFECTIONS

- 168 Infections Acquired in Health Care Facilities** 911
Robert A. Weinstein
- 169 Infections in Transplant Recipients** 919
Robert W. Finberg, Joyce Fingerhuth

SECTION 4 APPROACH TO THERAPY FOR BACTERIAL DISEASES

- 170 Treatment and Prophylaxis of Bacterial Infections** 930
David C. Hooper, Erica S. Shenoy, Christy A. Varughese

SECTION 5 DISEASES CAUSED BY GRAM-POSITIVE BACTERIA

- 171 Pneumococcal Infections** 946
David Goldblatt, Katherine L. O'Brien
- 172 Staphylococcal Infections** 954
Franklin D. Lowy
- 173 Streptococcal Infections** 963
Michael R. Wessels
- 174 Enterococcal Infections** 971
Cesar A. Arias, Barbara E. Murray
- 175 Diphtheria and Other Corynebacterial Infections** 977
William R. Bishai, John R. Murphy
- 176 Listeria monocytogenes Infections** 982
Elizabeth L. Hohmann, Daniel A. Portnoy
- 177 Tetanus** 984
C. Louise Thwaites, Lam Minh Yen
- 178 Botulism** 987
Susan Maslanka, Agam K. Rao
- 179 Gas Gangrene and Other Clostridial Infections** 990
Amy E. Bryant, Dennis L. Stevens

SECTION 6 DISEASES CAUSED BY GRAM-NEGATIVE BACTERIA

- 180 Meningococcal Infections** 995
Andrew J. Pollard
- 181 Gonococcal Infections** 1003
Sanjay Ram, Peter A. Rice
- 182 Haemophilus and Moraxella Infections** 1010
Timothy F. Murphy
- 183e Infections Due to the HACEK Group and Miscellaneous Gram-Negative Bacteria** 1014
Tamar F. Barlam, Dennis L. Kasper
- 184 Legionella Infections** 1014
Victor L. Yu, M. Luisa Pedro-Botet, Yusen E. Lin
- 185 Pertussis and Other Bordetella Infections** 1021
Karina A. Top, Scott A. Halperin
- 186 Diseases Caused by Gram-Negative Enteric Bacilli** 1025
Thomas A. Russo, James R. Johnson
- 187 Acinetobacter Infections** 1036
David L. Paterson, Anton Y. Peleg
- 188 Helicobacter pylori Infections** 1038
John C. Atherton, Martin J. Blaser
- 189 Infections Due to Pseudomonas Species and Related Organisms** 1042
Reuben Ramphal
- 190 Salmonellosis** 1049
David A. Pegues, Samuel I. Miller
- 191 Shigellosis** 1055
Philippe J. Sansonetti, Jean Bergounioux
- 192 Infections Due to Campylobacter and Related Organisms** 1058
Martin J. Blaser
- 193 Cholera and Other Vibrios** 1061
Matthew K. Waldor, Edward T. Ryan
- 194e Brucellosis** 1066
Nicholas J. Beeching, Michael J. Corbel
- 195 Tularemia** 1066
Richard F. Jacobs, Gordon E. Schutze
- 196 Plague and Other Yersinia Infections** 1070
Michael B. Prentice
- 197 Bartonella Infections, Including Cat-Scratch Disease** 1078
Michael Giladi, Moshe Ephros
- 198e Donovanosis** 1083
Nigel O'Farrell

SECTION 7 MISCELLANEOUS BACTERIAL INFECTIONS

- 199 Nocardiosis** 1084
Gregory A. Filice
- 200 Actinomycosis and Whipple's Disease** 1088
Thomas A. Russo
- 201 Infections Due to Mixed Anaerobic Organisms** 1094
Ronit Cohen-Poradosu, Dennis L. Kasper

SECTION 8 MYCOBACTERIAL DISEASES

- 202 Tuberculosis** 1102
Mario C. Raviglione
- 203 Leprosy** 1122
Robert H. Gelber
- 204 Nontuberculous Mycobacterial Infections** 1128
Steven M. Holland
- 205e Antimycobacterial Agents** 1132
Max R. O'Donnell, Divya Reddy, Jussi J. Saukkonen

SECTION 9 SPIROCHETAL DISEASES

- 206 Syphilis** 1132
Sheila A. Lukehart
- 207e Endemic Treponematoses** 1140
Sheila A. Lukehart
- 208 Leptospirosis** 1140
Rudy A. Hartskeerl, Jiri F. P. Wagenaar
- 209 Relapsing Fever** 1146
Alan G. Barbour
- 210 Lyme Borreliosis** 1149
Allen C. Steere

SECTION 10 DISEASES CAUSED BY RICKETTSIAE, MYCOPLASMAS, AND CHLAMYDIAE

- 211 Rickettsial Diseases** 1154
David H. Walker, J. Stephen Dumler, Thomas Marrie
- 212 Infections Due to Mycoplasmas** 1163
R. Doug Hardy
- 213 Chlamydial Infections** 1165
Charlotte A. Gaydos, Thomas C. Quinn

SECTION 11 VIRAL DISEASES: GENERAL CONSIDERATIONS

- 214e Medical Virology** 1174
Fred Wang, Elliott Kieff
- 215e Antiviral Chemotherapy, Excluding Antiretroviral Drugs** 1174
Lindsey R. Baden, Raphael Dolin

SECTION 12 INFECTIONS DUE TO DNA VIRUSES

- 216 Herpes Simplex Virus Infections** 1175
Lawrence Corey
- 217 Varicella-Zoster Virus Infections** 1183
Richard J. Whitley
- 218 Epstein-Barr Virus Infections, Including Infectious Mononucleosis** 1186
Jeffrey I. Cohen
- 219 Cytomegalovirus and Human Herpesvirus Types 6, 7, and 8** 1190
Camille Nelson Kotton, Martin S. Hirsch
- 220e Molluscum Contagiosum, Monkeypox, and Other Poxvirus Infections** 1194
Fred Wang
- 221 Parvovirus Infections** 1195
Kevin E. Brown
- 222 Human Papillomavirus Infections** 1197
Aaron C. Ermel, Darron R. Brown

SECTION 13 INFECTIONS DUE TO DNA AND RNA RESPIRATORY VIRUSES

- 223 Common Viral Respiratory Infections** 1202
Raphael Dolin
- 224 Influenza** 1209
Yehuda Z. Cohen, Raphael Dolin

SECTION 14 INFECTIONS DUE TO HUMAN IMMUNODEFICIENCY VIRUS AND OTHER HUMAN RETROVIRUSES

- 225e The Human Retroviruses** 1215
Dan L. Longo, Anthony S. Fauci

- 226 Human Immunodeficiency Virus Disease: AIDS and Related Disorders** 1215
Anthony S. Fauci, H. Clifford Lane



SECTION 15 INFECTIONS DUE TO RNA VIRUSES

- 227 Viral Gastroenteritis** 1285
Umesh D. Parashar, Roger I. Glass
- 228 Enterovirus, Parechovirus, and Reovirus Infections** 1289
Jeffrey I. Cohen
- 229 Measles (Rubeola)** 1295
Kaitlin Rainwater-Lovett, William J. Moss
-  **230e Rubella (German Measles)** 1299
Laura A. Zimmerman, Susan E. Reef
-  **231e Mumps** 1299
Steven A. Rubin, Kathryn M. Carbone
- 232 Rabies and Other Rhabdovirus Infections** 1299
Alan C. Jackson
- 233 Arthropod-Borne and Rodent-Borne Virus Infections** 1304
Jens H. Kuhn, Clarence J. Peters
- 234 Ebolavirus and Marburgvirus Infections** 1323
Jens H. Kuhn

SECTION 16 FUNGAL INFECTIONS

- 235 Diagnosis and Treatment of Fungal Infections** 1329
John E. Edwards, Jr.
- 236 Histoplasmosis** 1332
Chadi A. Hage, L. Joseph Wheat
- 237 Coccidioidomycosis** 1334
Neil M. Ampel
- 238 Blastomycosis** 1337
Donna C. Sullivan, Rathel L. Nolan, III
- 239 Cryptococcosis** 1339
Arturo Casadevall
- 240 Candidiasis** 1342
John E. Edwards, Jr.
- 241 Aspergillosis** 1345
David W. Denning
- 242 Mucormycosis** 1350
Brad Spellberg, Ashraf S. Ibrahim
- 243 Superficial Mycoses and Less Common Systemic Mycoses** 1353
Carol A. Kauffman
- 244 Pneumocystis Infections** 1358
Henry Masur, Alison Morris

SECTION 17 PROTOZOAL AND HELMINTHIC INFECTIONS: GENERAL CONSIDERATIONS

-  **245e Laboratory Diagnosis of Parasitic Infections** 1362
Sharon L. Reed, Charles E. Davis
-  **246e Agents Used to Treat Parasitic Infections** 1363
Thomas A. Moore

SECTION 18 PROTOZOAL INFECTIONS

- 247 Amebiasis and Infection with Free-Living Amebas** 1363
Rosa M. Andrade, Sharon L. Reed
- 248 Malaria** 1368
Nicholas J. White, Joel G. Breman

- 249 Babesiosis** 1384
Edouard G. Vannier, Peter J. Krause

-  **250e Atlas of Blood Smears of Malaria and Babesiosis** 1387
Nicholas J. White, Joel G. Breman

- 251 Leishmaniasis** 1387
Shyam Sundar

- 252 Chagas Disease and African Trypanosomiasis** 1394
Louis V. Kirchhoff, Anis Rassi Jr.

- 253 Toxoplasma Infections** 1398
Kami Kim, Lloyd H. Kasper

- 254 Protozoal Intestinal Infections and Trichomoniasis** 1405
Peter F. Weller

SECTION 19 HELMINTHIC INFECTIONS

-  **255e Introduction to Helminthic Infections** 1409
Peter F. Weller

- 256 Trichinellosis and Other Tissue Nematode Infections** 1410
Peter F. Weller

- 257 Intestinal Nematode Infections** 1413
Peter F. Weller, Thomas B. Nutman

- 258 Filarial and Related Infections** 1417
Thomas B. Nutman, Peter F. Weller

- 259 Schistosomiasis and Other Trematode Infections** 1423
Charles H. King, Adel A. F. Mahmoud

- 260 Cestode Infections** 1430
A. Clinton White, Jr., Peter F. Weller

PART 9 Terrorism and Clinical Medicine

-  **261e Microbial Bioterrorism** 1437
H. Clifford Lane, Anthony S. Fauci

-  **262e Chemical Terrorism** 1437
Charles G. Hurst, Jonathan Newmark, James A. Romano, Jr.

-  **263e Radiation Terrorism** 1437
Christine E. Hill-Kayser, Eli Glatstein, Zelig A. Tochner

PART 10 Disorders of the Cardiovascular System

SECTION 1 INTRODUCTION TO CARDIOVASCULAR DISORDERS

- 264 Approach to the Patient with Possible Cardiovascular Disease** 1439
Joseph Loscalzo


-  **265e Basic Biology of the Cardiovascular System** 1441
Joseph Loscalzo, Peter Libby, Jonathan A. Epstein

-  **266e Epidemiology of Cardiovascular Disease** 1442
Thomas A. Gaziano, J. Michael Gaziano

SECTION 2 DIAGNOSIS OF CARDIOVASCULAR DISORDERS

- 267 Physical Examination of the Cardiovascular System** 1442
Patrick T. O'Gara, Joseph Loscalzo

- 268 Electrocardiography** 1450
Ary L. Goldberger
- 269e Atlas of Electrocardiography** 1459
Ary L. Goldberger
- 270e Noninvasive Cardiac Imaging: Echocardiography, Nuclear Cardiology, and Magnetic Resonance/Computed Tomography Imaging** 1459
Marcelo F. Di Carli, Raymond Y. Kwong, Scott D. Solomon
- 271e Atlas of Noninvasive Imaging** 1460
Marcelo F. Di Carli, Raymond Y. Kwong, Scott D. Solomon
- 272 Diagnostic Cardiac Catheterization and Coronary Angiography** 1460
Jane A. Leopold, David P. Faxon
- SECTION 3 DISORDERS OF RHYTHM**
- 273e Principles of Electrophysiology** 1466
David D. Spragg, Gordon F. Tomaselli
- 274 The Bradyarrhythmias: Disorders of the Sinoatrial Node** 1466
David D. Spragg, Gordon F. Tomaselli
- 275 The Bradyarrhythmias: Disorders of the Atrioventricular Node** 1470
David D. Spragg, Gordon F. Tomaselli
- 276 Supraventricular Tachyarrhythmias** 1476
Gregory F. Michaud, William G. Stevenson
- 277 Ventricular Arrhythmias** 1489
Roy M. John, William G. Stevenson
- 278e Atlas of Cardiac Arrhythmias** 1500
Ary L. Goldberger
- SECTION 4 DISORDERS OF THE HEART**
- 279 Heart Failure: Pathophysiology and Diagnosis** 1500
Douglas L. Mann, Murali Chakinala
- 280 Heart Failure: Management** 1507
Mandeep R. Mehra
- 281 Cardiac Transplantation and Prolonged Assisted Circulation** 1516
Sharon A. Hunt, Hari R. Mallidi
- 282 Congenital Heart Disease in the Adult** 1519
Jamil A. Aboulhosn, John S. Child
- 283 Aortic Valve Disease** 1528
Patrick T. O’Gara, Joseph Loscalzo
- 284 Mitral Valve Disease** 1539
Patrick T. O’Gara, Joseph Loscalzo
- 285 Tricuspid and Pulmonic Valve Disease** 1547
Patrick T. O’Gara, Joseph Loscalzo
- 286 Multiple and Mixed Valvular Heart Disease** 1550
Patrick T. O’Gara, Joseph Loscalzo
- 287 Cardiomyopathy and Myocarditis** 1553
Neal K. Lakdawala, Lynne Warner Stevenson, Joseph Loscalzo
- 288 Pericardial Disease** 1571
Eugene Braunwald
- 289e Tumors and Trauma of the Heart** 1577
Eric H. Awtry, Wilson S. Colucci
- 290e Cardiac Manifestations of Systemic Disease** 1577
Eric H. Awtry, Wilson S. Colucci
- SECTION 5 CORONARY AND PERIPHERAL VASCULAR DISEASE**
- 291e The Pathogenesis, Prevention, and Treatment of Atherosclerosis** 1578
Peter Libby
- 292e Atlas of Atherosclerosis** 1578
Peter Libby
- 293 Ischemic Heart Disease** 1578
Elliott M. Antman, Joseph Loscalzo
- 294 Non-ST-Segment Elevation Acute Coronary Syndrome (Non-ST-Segment Elevation Myocardial Infarction and Unstable Angina)** 1593
Christopher P. Cannon, Eugene Braunwald
- 295 ST-Segment Elevation Myocardial Infarction** 1599
Elliott M. Antman, Joseph Loscalzo
- 296e Percutaneous Coronary Interventions and Other Interventional Procedures** 1611
David P. Faxon, Deepak L. Bhatt
- 297e Atlas of Percutaneous Revascularization** 1611
Jane A. Leopold, Deepak L. Bhatt, David P. Faxon
- 298 Hypertensive Vascular Disease** 1611
Theodore A. Kotchen
- 299 Renovascular Disease** 1627
Stephen C. Textor
- 300 Deep Venous Thrombosis and Pulmonary Thromboembolism** 1631
Samuel Z. Goldhaber
- 301 Diseases of the Aorta** 1637
Mark A. Creager, Joseph Loscalzo
- 302 Arterial Diseases of the Extremities** 1643
Mark A. Creager, Joseph Loscalzo
- 303 Chronic Venous Disease and Lymphedema** 1650
Mark A. Creager, Joseph Loscalzo
- 304 Pulmonary Hypertension** 1655
Aaron B. Waxman, Joseph Loscalzo
- PART 11 Disorders of the Respiratory System**
- SECTION 1 DIAGNOSIS OF RESPIRATORY DISORDERS**
- 305 Approach to the Patient with Disease of the Respiratory System** 1661
Patricia A. Kritek, Augustine M. K. Choi
- 306e Disturbances of Respiratory Function** 1663
Edward T. Naureckas, Julian Solway
- 307 Diagnostic Procedures in Respiratory Disease** 1663
Anne L. Fuhlbrigge, Augustine M. K. Choi
- 308e Atlas of Chest Imaging** 1669
Patricia A. Kritek, John J. Reilly, Jr.
- SECTION 2 DISEASES OF THE RESPIRATORY SYSTEM**
- 309 Asthma** 1669
Peter J. Barnes
- 310 Hypersensitivity Pneumonitis and Pulmonary Infiltrates with Eosinophilia** 1681
Praveen Akuthota, Michael E. Wechsler
- 311 Occupational and Environmental Lung Disease** 1687
John R. Balmes, Frank E. Speizer
- 312 Bronchiectasis** 1694
Rebecca M. Baron, Miriam Baron Barshak
- 313 Cystic Fibrosis** 1697
Eric J. Sorscher
- 314 Chronic Obstructive Pulmonary Disease** 1700
John J. Reilly, Jr., Edwin K. Silverman, Steven D. Shapiro

315	Interstitial Lung Diseases	1708
	<i>Talmadge E. King, Jr.</i>	
316	Disorders of the Pleura	1716
	<i>Richard W. Light</i>	
317	Disorders of the Mediastinum	1719
	<i>Richard W. Light</i>	
318	Disorders of Ventilation	1720
	<i>John F. McConville, Babak Mokhlesi, Julian Solway</i>	
319	Sleep Apnea	1723
	<i>Andrew Wellman, Susan Redline</i>	
	320e Lung Transplantation	1727
	<i>Elbert P. Trulock</i>	

PART 12 Critical Care Medicine


SECTION 1 RESPIRATORY CRITICAL CARE

321	Approach to the Patient with Critical Illness	1729
	<i>John P. Kress, Jesse B. Hall</i>	
322	Acute Respiratory Distress Syndrome	1736
	<i>Bruce D. Levy, Augustine M. K. Choi</i>	
323	Mechanical Ventilatory Support	1740
	<i>Bartolome R. Celli</i>	

SECTION 2 SHOCK AND CARDIAC ARREST

324	Approach to the Patient with Shock	1744
	<i>Ronald V. Maier</i>	
325	Severe Sepsis and Septic Shock	1751
	<i>Robert S. Munford</i>	
326	Cardiogenic Shock and Pulmonary Edema	1759
	<i>Judith S. Hochman, David H. Ingbar</i>	
327	Cardiovascular Collapse, Cardiac Arrest, and Sudden Cardiac Death	1764
	<i>Robert J. Myerburg, Agustin Castellanos</i>	



SECTION 3 NEUROLOGIC CRITICAL CARE

328	Coma	1771
	<i>Allan H. Ropper</i>	
	329e Examination of the Comatose Patient	1777
	<i>S. Andrew Josephson</i>	
330	Neurologic Critical Care, Including Hypoxic-Ischemic Encephalopathy, and Subarachnoid Hemorrhage	1777
	<i>J. Claude Hemphill, III, Wade S. Smith, Daryl R. Gress</i>	

SECTION 4 ONCOLOGIC EMERGENCIES

331	Oncologic Emergencies	1787
	<i>Rasim Gucalp, Janice P. Dutcher</i>	

PART 13 Disorders of the Kidney and Urinary Tract

	332e Cellular and Molecular Biology of the Kidney	1799
	<i>Alfred L. George, Jr., Eric G. Neilson</i>	
	333e Adaptation of the Kidney to Injury	1799
	<i>Joseph V. Bonventre</i>	
334	Acute Kidney Injury	1799
	<i>Sushrut S. Waikar, Joseph V. Bonventre</i>	

335	Chronic Kidney Disease	1811
	<i>Joanne M. Bargman, Karl Skorecki</i>	
336	Dialysis in the Treatment of Renal Failure	1822
	<i>Kathleen D. Liu, Glenn M. Chertow</i>	
337	Transplantation in the Treatment of Renal Failure	1825
	<i>Jamil Azzi, Edgar L. Milford, Mohamed H. Sayegh, Anil Chandraker</i>	
338	Glomerular Diseases	1831
	<i>Julia B. Lewis, Eric G. Neilson</i>	
339	Polycystic Kidney Disease and Other Inherited Disorders of Tubule Growth and Development	1850
	<i>Jing Zhou, Martin R. Pollak</i>	
340	Tubulointerstitial Diseases of the Kidney	1856
	<i>Laurence H. Beck, David J. Salant</i>	
341	Vascular Injury to the Kidney	1863
	<i>Nelson Leung, Stephen C. Textor</i>	
342	Nephrolithiasis	1866
	<i>Gary C. Curhan</i>	
343	Urinary Tract Obstruction	1871
	<i>Julian L. Seifter</i>	

PART 14 Disorders of the Gastrointestinal System

SECTION 1 DISORDERS OF THE ALIMENTARY TRACT

344	Approach to the Patient with Gastrointestinal Disease	1875
	<i>William L. Hasler, Chung Owyang</i>	
345	Gastrointestinal Endoscopy	1880
	<i>Louis Michel Wong Kee Song, Mark Topazian</i>	
	346e Video Atlas of Gastrointestinal Endoscopy	1900
	<i>Louis Michel Wong Kee Song, Mark Topazian</i>	
347	Diseases of the Esophagus	1900
	<i>Peter J. Kahrilas, Ikuo Hirano</i>	
348	Peptic Ulcer Disease and Related Disorders	1911
	<i>John Del Valle</i>	
349	Disorders of Absorption	1932
	<i>Henry J. Binder</i>	
	350e The Schilling Test	1946
	<i>Henry J. Binder</i>	
351	Inflammatory Bowel Disease	1947
	<i>Sonia Friedman, Richard S. Blumberg</i>	
352	Irritable Bowel Syndrome	1965
	<i>Chung Owyang</i>	
353	Diverticular Disease and Common Anorectal Disorders	1971
	<i>Rizwan Ahmed, Susan L. Gearhart</i>	
354	Mesenteric Vascular Insufficiency	1978
	<i>Rizwan Ahmed, Mahmoud Malas</i>	
355	Acute Intestinal Obstruction	1981
	<i>Danny O. Jacobs</i>	
356	Acute Appendicitis and Peritonitis	1985
	<i>Danny O. Jacobs</i>	

SECTION 2 LIVER AND BILIARY TRACT DISEASE

357	Approach to the Patient with Liver Disease	1989
	<i>Marc G. Ghany, Jay H. Hoofnagle</i>	
358	Evaluation of Liver Function	1995
	<i>Daniel S. Pratt</i>	

359 The Hyperbilirubinemias 1999
Allan W. Wolkoff

360 Acute Viral Hepatitis 2004
Jules L. Dienstag

361 Toxic and Drug-Induced Hepatitis 2023
William M. Lee, Jules L. Dienstag

362 Chronic Hepatitis 2031
Jules L. Dienstag

363 Alcoholic Liver Disease 2052
Mark E. Mailliard, Michael F. Sorrell

364 Nonalcoholic Fatty Liver Diseases and Nonalcoholic Steatohepatitis 2054
Manal F. Abdelmalek, Anna Mae Diehl

365 Cirrhosis and Its Complications 2058
Bruce R. Bacon

366e Atlas of Liver Biopsies 2067
Jules L. Dienstag, Atul K. Bhan

367e Genetic, Metabolic, and Infiltrative Diseases Affecting the Liver 2067
Bruce R. Bacon

368 Liver Transplantation 2067
Raymond T. Chung, Jules L. Dienstag

369 Diseases of the Gallbladder and Bile Ducts 2075
Norton J. Greenberger, Gustav Paumgartner

SECTION 3 DISORDERS OF THE PANCREAS

370 Approach to the Patient with Pancreatic Disease 2086
Darwin L. Conwell, Norton J. Greenberger, Peter A. Banks

371 Acute and Chronic Pancreatitis 2090
Darwin L. Conwell, Peter A. Banks, Norton J. Greenberger

PART 15 Immune-Mediated, Inflammatory, and Rheumatologic Disorders

SECTION 1 THE IMMUNE SYSTEM IN HEALTH AND DISEASE

372e Introduction to the Immune System 2103
Barton F. Haynes, Kelly A. Soderberg, Anthony S. Fauci

373e The Major Histocompatibility Complex 2103
Gerald T. Nepom

374 Primary Immune Deficiency Diseases 2103
Alain Fischer

375e Primary Immunodeficiencies Associated with (or Secondary to) Other Diseases 2113
Alain Fischer

SECTION 2 DISORDERS OF IMMUNE-MEDIATED INJURY

376 Allergies, Anaphylaxis, and Systemic Mastocytosis 2113
Joshua A. Boyce, K. Frank Austen

377e Autoimmunity and Autoimmune Diseases 2124
Betty Diamond, Peter E. Lipsky

378 Systemic Lupus Erythematosus 2124
Bevra Hannahs Hahn

379 Antiphospholipid Syndrome 2134
Haralampos M. Moutsopoulos, Panayiotis G. Vlachoyiannopoulos

380 Rheumatoid Arthritis 2136
Ankooor Shah, E. William St. Clair

381 Acute Rheumatic Fever 2149
Jonathan R. Carapetis

382 Systemic Sclerosis (Scleroderma) and Related Disorders 2154
John Varga

383 Sjögren’s Syndrome 2166
Haralampos M. Moutsopoulos, Athanasios G. Tzioufas

384 The Spondyloarthritides 2169
Joel D. Taurog, John D. Carter

385 The Vasculitis Syndromes 2179
Carol A. Langford, Anthony S. Fauci

386e Atlas of the Vasculitic Syndromes 2193
Carol A. Langford, Anthony S. Fauci

387 Behçet’s Syndrome 2194
Haralampos M. Moutsopoulos

388 Polymyositis, Dermatomyositis, and Inclusion Body Myositis 2194
Marinos C. Dalakas

389 Relapsing Polychondritis 2202
Carol A. Langford

390 Sarcoidosis 2205
Robert P. Baughman, Elyse E. Lower

391e IgG4-Related Disease 2212
John H. Stone

392 Familial Mediterranean Fever and Other Hereditary Autoinflammatory Diseases 2212
Daniel L. Kastner

SECTION 3 DISORDERS OF THE JOINTS AND ADJACENT TISSUES

393 Approach to Articular and Musculoskeletal Disorders 2216
John J. Cush

394 Osteoarthritis 2226
David T. Felson

395 Gout and Other Crystal-Associated Arthropathies 2233
H. Ralph Schumacher, Lan X. Chen

396 Fibromyalgia 2238
Leslie J. Crofford

397 Arthritis Associated with Systemic Disease, and Other Arthritides 2240
Carol A. Langford, Brian F. Mandell

398 Periarticular Disorders of the Extremities 2247
Carol A. Langford

PART 16 Endocrinology and Metabolism

SECTION 1 ENDOCRINOLOGY

399 Approach to the Patient with Endocrine Disorders 2251
J. Larry Jameson

400e Mechanisms of Hormone Action 2255
J. Larry Jameson


401e Anterior Pituitary: Physiology of Pituitary Hormones 2255
Shlomo Melmed, J. Larry Jameson

402 Hypopituitarism	2255
<i>Shlomo Melmed, J. Larry Jameson</i>	
403 Anterior Pituitary Tumor Syndromes	2261
<i>Shlomo Melmed, J. Larry Jameson</i>	
404 Disorders of the Neurohypophysis	2274
<i>Gary L. Robertson</i>	
405 Disorders of the Thyroid Gland	2283
<i>J. Larry Jameson, Susan J. Mandel, Anthony P. Weetman</i>	
406 Disorders of the Adrenal Cortex	2309
<i>Wiebke Arlt</i>	
407 Pheochromocytoma	2329
<i>Hartmut P. H. Neumann</i>	
408 Multiple Endocrine Neoplasia	2335
<i>Rajesh V. Thakker</i>	
409 Autoimmune Polyendocrine Syndromes	2344
<i>Peter A. Gottlieb</i>	

SECTION 2 REPRODUCTIVE ENDOCRINOLOGY


410 Disorders of Sex Development	2349
<i>John C. Achermann, J. Larry Jameson</i>	
411 Disorders of the Testes and Male Reproductive System	2357
<i>Shalender Bhasin, J. Larry Jameson</i>	
412 Disorders of the Female Reproductive System	2375
<i>Janet E. Hall</i>	
413 Menopause and Postmenopausal Hormone Therapy	2381
<i>JoAnn E. Manson, Shari S. Bassuk</i>	
414 Infertility and Contraception	2387
<i>Janet E. Hall</i>	

SECTION 3 OBESITY, DIABETES MELLITUS, AND METABOLIC SYNDROME





 415e Biology of Obesity	2392
<i>Jeffrey S. Flier, Eleftheria Maratos-Flier</i>	
416 Evaluation and Management of Obesity	2392
<i>Robert F. Kushner</i>	
417 Diabetes Mellitus: Diagnosis, Classification, and Pathophysiology	2399
<i>Alvin C. Powers</i>	
418 Diabetes Mellitus: Management and Therapies	2407
<i>Alvin C. Powers</i>	
419 Diabetes Mellitus: Complications	2422
<i>Alvin C. Powers</i>	
420 Hypoglycemia	2430
<i>Philip E. Cryer, Stephen N. Davis</i>	
421 Disorders of Lipoprotein Metabolism	2435
<i>Daniel J. Rader, Helen H. Hobbs</i>	
422 The Metabolic Syndrome	2449
<i>Robert H. Eckel</i>	

SECTION 4 DISORDERS OF BONE AND MINERAL METABOLISM

423 Bone and Mineral Metabolism in Health and Disease	2454
<i>F. Richard Bringhurst, Marie B. Demay, Stephen M. Krane, Henry M. Kronenberg</i>	
424 Disorders of the Parathyroid Gland and Calcium Homeostasis	2466
<i>John T. Potts, Jr., Harald Jüppner</i>	








425 Osteoporosis	2488
<i>Robert Lindsay, Felicia Cosman</i>	
 426e Paget's Disease and Other Dysplasias of Bone	2504
<i>Murray J. Favus, Tamara J. Vokes</i>	

SECTION 5 DISORDERS OF INTERMEDIARY METABOLISM

427 Heritable Disorders of Connective Tissue	2504
<i>Darwin J. Prockop, John F. Bateman</i>	
428 Hemochromatosis	2514
<i>Lawrie W. Powell</i>	
429 Wilson's Disease	2519
<i>George J. Brewer</i>	
430 The Porphyrrias	2521
<i>Robert J. Desnick, Manisha Balwani</i>	
 431e Disorders of Purine and Pyrimidine Metabolism	2533
<i>Christopher M. Burns, Robert L. Wortmann</i>	
 432e Lysosomal Storage Diseases	2533
<i>Robert J. Hopkin, Gregory A. Grabowski</i>	
 433e Glycogen Storage Diseases and Other Inherited Disorders of Carbohydrate Metabolism	2534
<i>Priya S. Kishmani, Yuan-Tsong Chen</i>	
 434e Inherited Disorders of Amino Acid Metabolism in Adults	2534
<i>Nicola Longo</i>	
 435e Inherited Defects of Membrane Transport	2534
<i>Nicola Longo</i>	
 436e Atlas of Clinical Manifestations of Metabolic Diseases	2534
<i>J. Larry Jameson</i>	

PART 17 Neurologic Disorders

SECTION 1 DIAGNOSIS OF NEUROLOGIC DISORDERS

437 Approach to the Patient with Neurologic Disease	2535
<i>Daniel H. Lowenstein, Joseph B. Martin, Stephen L. Hauser</i>	
 438e The Neurologic Screening Exam	2540
<i>Daniel H. Lowenstein</i>	
 439e Video Atlas of the Detailed Neurologic Examination	2540
<i>Martin A. Samuels</i>	
 440e Neuroimaging in Neurologic Disorders	2540
<i>William P. Dillon</i>	
 441e Atlas of Neuroimaging	2541
<i>Andre D. Furtado, William P. Dillon</i>	
 442e Electrodiagnostic Studies of Nervous System Disorders: EEG, Evoked Potentials, and EMG	2541
<i>Michael J. Aminoff</i>	
 443e Technique of Lumbar Puncture	2541
<i>Elizabeth Robbins, Stephen L. Hauser</i>	
 444e Biology of Neurologic Diseases	2541
<i>Stephen L. Hauser, Stanley B. Prusiner, M. Flint Beal</i>	

SECTION 2 DISEASES OF THE CENTRAL NERVOUS SYSTEM

445 Seizures and Epilepsy	2542
<i>Daniel H. Lowenstein</i>	

446	Cerebrovascular Diseases	2559		469e Cocaine and Other Commonly Abused Drugs	2729
	<i>Wade S. Smith, S. Claiborne Johnston, J. Claude Hemphill, III</i>			<i>Nancy K. Mello, Jack H. Mendelson</i>	
447	Migraine and Other Primary Headache Disorders	2586	470	Nicotine Addiction	2729
	<i>Peter J. Goadsby, Neil H. Raskin</i>			<i>David M. Burns</i>	
448	Alzheimer's Disease and Other Dementias	2598		471e Neuropsychiatric Illnesses in War Veterans	2732
	<i>William W. Seeley, Bruce L. Miller</i>			<i>Charles W. Hoge</i>	
449	Parkinson's Disease and Other Movement Disorders	2609			
	<i>C. Warren Olanow, Anthony H.V. Schapira, Jose A. Obeso</i>				
450	Ataxic Disorders	2626			
	<i>Roger N. Rosenberg</i>				
	451e Classification of the Spinocerebellar Ataxias	2631			
	<i>Roger N. Rosenberg</i>				
452	Amyotrophic Lateral Sclerosis and Other Motor Neuron Diseases	2631		472e Heavy Metal Poisoning	2733
	<i>Robert H. Brown, Jr.</i>			<i>Howard Hu</i>	
	453e Prion Diseases	2637		473e Poisoning and Drug Overdose	2733
	<i>Stanley B. Prusiner, Bruce L. Miller</i>			<i>Mark B. Mycyk</i>	
454	Disorders of the Autonomic Nervous System	2637	474	Disorders Caused by Venomous Snakebites and Marine Animal Exposures	2733
	<i>Phillip A. Low, John W. Engstrom</i>			<i>Charles Lei, Natalie J. Badowski, Paul S. Auerbach, Robert L. Norris</i>	
455	Trigeminal Neuralgia, Bell's Palsy, and Other Cranial Nerve Disorders	2645	475	Ectoparasite Infestations and Arthropod Injuries	2744
	<i>M. Flint Beal, Stephen L. Hauser</i>			<i>Richard J. Pollack, Scott A. Norton</i>	
456	Diseases of the Spinal Cord	2650			
	<i>Stephen L. Hauser, Allan H. Ropper</i>				
	457e Concussion and Other Traumatic Brain Injuries	2660			
	<i>Allan H. Ropper</i>				
458	Multiple Sclerosis and Other Demyelinating Diseases	2661			
	<i>Stephen L. Hauser, Douglas S. Goodin</i>				
SECTION 3 NERVE AND MUSCLE DISORDERS					
459	Peripheral Neuropathy	2674			
	<i>Anthony A. Amato, Richard J. Barohn</i>				
460	Guillain-Barré Syndrome and Other Immune-Mediated Neuropathies	2694			
	<i>Stephen L. Hauser, Anthony A. Amato</i>				
461	Myasthenia Gravis and Other Diseases of the Neuromuscular Junction	2701			
	<i>Daniel B. Drachman, Anthony A. Amato</i>				
	462e Muscular Dystrophies and Other Muscle Diseases	2707			
	<i>Anthony A. Amato, Robert H. Brown, Jr.</i>				
	463e Special Issues in Inpatient Neurologic Consultation	2707			
	<i>S. Andrew Josephson, Martin A. Samuels</i>				
SECTION 4 CHRONIC FATIGUE SYNDROME					
	464e Chronic Fatigue Syndrome	2707			
	<i>Gijs Bleijenberg, Jos W. M. van der Meer</i>				
SECTION 5 PSYCHIATRIC AND ADDICTION DISORDERS					
	465e Biology of Psychiatric Disorders	2708			
	<i>Robert O. Messing, Eric J. Nestler</i>				
466	Mental Disorders	2708			
	<i>Victor I. Reus</i>				
467	Alcohol and Alcoholism	2723			
	<i>Marc A. Schuckit</i>				
	468e Opioid-Related Disorders	2728			
	<i>Thomas R. Kosten, Colin N. Haile</i>				
				PART 18 Poisoning, Drug Overdose, and Envenomation	
				472e Heavy Metal Poisoning	2733
				<i>Howard Hu</i>	
				473e Poisoning and Drug Overdose	2733
				<i>Mark B. Mycyk</i>	
			474	Disorders Caused by Venomous Snakebites and Marine Animal Exposures	2733
				<i>Charles Lei, Natalie J. Badowski, Paul S. Auerbach, Robert L. Norris</i>	
			475	Ectoparasite Infestations and Arthropod Injuries	2744
				<i>Richard J. Pollack, Scott A. Norton</i>	
				PART 19 Disorders Associated with Environmental Exposures	
				476e Altitude Illness	2753
				<i>Buddha Basnyat, Geoffrey Tabin</i>	
				477e Hyperbaric and Diving Medicine	2753
				<i>Michael H. Bennett, Simon J. Mitchell</i>	
				478e Hypothermia and Frostbite	2753
				<i>Daniel F. Danzl</i>	
				479e Heat-Related Illnesses	2753
				<i>Daniel F. Danzl</i>	
				APPENDIX: Laboratory Values of Clinical Importance	2754
				<i>Alexander Kratz, Michael A. Pesce, Robert C. Basner, Andrew J. Einstein</i>	
				480e The Clinical Laboratory in Modern Health Care	2770
				<i>Anthony A. Killeen</i>	
				481e Clinical Procedure Tutorial: Central Venous Catheter Placement	2770
				<i>Maria A. Yialamas, William E. Corcoran, Gyorgy Frenzl, Kurt Fink</i>	
				482e Clinical Procedure Tutorial: Thoracentesis	2770
				<i>Charles A. Morris, Andrea S. Wolf</i>	
				483e Clinical Procedure Tutorial: Abdominal Paracentesis	2770
				<i>Maria A. Yialamas, Anna E. Rutherford, Lindsay King</i>	
				484e Clinical Procedure Tutorial: Endotracheal Intubation	2770
				<i>Charles A. Morris, Emily Page Nelson</i>	
				485e Clinical Procedures Tutorial: Percutaneous Arterial Blood Gas Sampling	2770
				<i>Christian D. Becker</i>	
				<i>Medical Editors: Sean Sadikot, Jeremy Matloff</i>	
				486e Clinical Procedures Tutorial: Lumbar Puncture	2770
				<i>Beth Rapaport, Stephen Krieger, Corey McGraw</i>	
				<i>Medical Editors: Sean Sadikot, Jeremy Matloff</i>	
				Index	I-1

This page intentionally left blank

James L. Abbruzzese, MD

Chief, Division of Medical Oncology, Department of Medicine; Associate Director, Clinical Research, Duke Cancer Institute, Durham, North Carolina [120e]

Manal F. Abdelmalek, MD, MPH

Associate Professor of Medicine, Division of Gastroenterology and Hepatology, Duke University, Durham, North Carolina [364]

Jamil Aboulhosn, MD

Assistant Professor of Medicine; Director Ahmanson/UCLA Adult Congenital Heart Disease Center, David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, California [282]

John C. Achermann, MD, PhD, MB

Wellcome Trust Senior Research Fellow in Clinical Science, University College London; Professor of Paediatric Endocrinology, UCL Institute of Child Health, University College London, London, United Kingdom [410]

John W. Adamson, MD

Clinical Professor, Division of Hematology/Oncology, Department of Medicine, University of California at San Diego, San Diego, California [77, 126]

Rizwan Ahmed, MD

General Surgery Resident, Department of General Surgery, Johns Hopkins Hospital, Baltimore, Maryland [353, 354]

Praveen Akuthota, MD

Assistant Professor of Medicine, Harvard Medical School; Beth Israel Deaconess Medical Center, Boston, Massachusetts [310]

Anthony A. Amato, MD

Professor of Neurology, Harvard Medical School; Vice-Chairman, Department of Neurology, Brigham and Women's Hospital, Boston, Massachusetts [459-462e]

Michael J. Aminoff, MD, DSc, FRCP

Professor of Neurology, School of Medicine, University of California, San Francisco, San Francisco, California [30, 31, 442e]

Neil M. Ampel, MD

Professor of Medicine, University of Arizona; Staff Physician, Southern Arizona Veterans Affairs Health Care System, Tucson, Arizona [237]

Kenneth C. Anderson, MD

Kraft Family Professor of Medicine, Harvard Medical School; Chief, Jerome Lipper Multiple Myeloma Center, Dana-Farber Cancer Institute, Boston, Massachusetts [136, 138e]

Rosa M. Andrade, MD

Department of Medicine, Division of Infectious Diseases, University of California, San Diego, San Diego, California [247]

Elliott M. Antman, MD

Professor of Medicine, Cardiovascular Division, Department of Medicine, Brigham and Women's Hospital; Associate Dean for Clinical/Translational Research, Harvard Medical School, Boston, Massachusetts [293, 295]

Frederick R. Appelbaum, MD

Director, Division of Clinical Research, Fred Hutchinson Cancer Research Center, Seattle, Washington [139e]

Cesar A. Arias, MD, PhD, MSC

Associate Professor of Medicine, Microbiology and Molecular Genetics; Director, Laboratory for Antimicrobial Research, University of Texas Medical School at Houston, Houston, Texas; Director, Molecular Genetics and Antimicrobial Unit; Co-Director, International Center for Microbial Genomics, Universidad, El Bosque, Bogota, Colombia [174]

Wiebke Arlt, MD, DSc, FRCP, FMedSci

Professor of Medicine, Centre for Endocrinology, Diabetes and Metabolism, School of Clinical and Experimental Medicine, University of Birmingham; Consultant Endocrinologist, University Hospital Birmingham, Birmingham, United Kingdom [406]

Katrina Armstrong, MD, MS

Jackson Professor of Clinical Medicine, Physician in Chief, Massachusetts General Hospital, Boston Massachusetts [4]

Valder R. Arruda, MD, PhD

Associate Professor, Division of Hematology, Department of Pediatrics, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [141]

Andrew W. Artenstein, MD

Professor of Medicine, Tufts University School of Medicine, Boston, Massachusetts; Adjunct Professor of Medicine and Health Services, Policy and Practice, Alpert Medical School, Brown University, Providence, Rhode Island; Chair, Department of Medicine, Baystate Health, Springfield, Massachusetts [152e]

Anthony Atala, MD

Professor and Director, Wake Forest Institute for Regenerative Medicine, Wake Forest School of Medicine, Winston-Salem, North Carolina [92e]

John C. Atherton, MD, FRCP

Professor of Gastroenterology and Dean of the School of Medicine, University of Nottingham, Nottingham, United Kingdom [188]

Paul S. Auerbach, MD, MS, FACEP, FAWM

Redich Family Professor of Surgery, Division of Emergency Medicine, Stanford University School of Medicine, Stanford, California [474]

K. Frank Austen, MD

AstraZeneca Professor of Respiratory and Inflammatory Diseases, Director, Inflammation and Allergic Diseases Research Section, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [376]

Eric H. Awtry, MD

Cardiology Division, Boston Medical Center, Boston, Massachusetts [289e, 290e]

Jamil Azzi, MD

Instructor in Medicine, Harvard Medical School; Associate Physician, Brigham and Women's Hospital, Boston, Massachusetts [337]

Bruce R. Bacon, MD

James F. King, MD Endowed Chair in Gastroenterology; Professor of Internal Medicine, Saint Louis University Liver Center, Saint Louis University School of Medicine, St. Louis, Missouri [365, 367e]

Lindsey R. Baden, MD

Associate Professor of Medicine, Harvard Medical School; Dana-Farber Cancer Institute, Brigham and Women's Hospital, Boston, Massachusetts [215e]

Natalie J. Badowski, MD

Division of Emergency Medicine, Stanford University School of Medicine, Stanford, California [474]

John R. Balmes, MD

Professor of Medicine, University of California, San Francisco; Professor, School of Public Health, University of California, Berkeley; Chief, Division of Occupational and Environmental Medicine, San Francisco General Hospital, San Francisco, California [311]

Manisha Balwani, MD, MS

Assistant Professor, Department of Genetics and Genomic Sciences, Mount Sinai School of Medicine of New York University, New York, New York [430]

Peter A. Banks, MD

Professor of Medicine, Harvard Medical School; Senior Physician, Division of Gastroenterology, Brigham and Women's Hospital, Boston, Massachusetts [370, 371]

Robert L. Barbieri, MD

Kate Macy Ladd Professor of Obstetrics, Gynecology and Reproductive Biology, Harvard Medical School; Chair, Department of Obstetrics and Gynecology, Brigham and Women's Hospital, Boston, Massachusetts [8]

Alan G. Barbour, MD

Professor of Medicine and Microbiology and Molecular Genetics, University of California Irvine, Irvine, California [209]

Joanne M. Bargman, MD, FRCP

Professor of Medicine, University of Toronto; Staff Nephrologist, University Health Network, Toronto, Canada [335]

Tamar F. Barlam, MD, MSc

Associate Professor of Medicine, Infectious Disease Section, Boston University School of Medicine, Boston, Massachusetts [147, 183e]

Peter J. Barnes, DM, DSc, FMedSci, FRS

Head of Respiratory Medicine, Imperial College, London, United Kingdom [309]

Richard J. Barohn, MD

Chairman, Department of Neurology; Gertrude and Dewey Ziegler Professor of Neurology, University of Kansas Medical Center, Kansas City, Kansas [459]

Rebecca M. Baron, MD

Assistant Professor of Medicine, Harvard Medical School; Associate Physician, Brigham and Women's Hospital, Division of Pulmonary and Critical Care Medicine, Department of Medicine, Boston, Massachusetts [154, 312]

Miriam Baron Barshak, MD

Assistant Professor, Harvard Medical School; Associate Physician, Massachusetts General Hospital, Boston, Massachusetts [154, 159, 312]

Shehzad Basaria, MD

Men's Health: Aging and Metabolism, Brigham and Women's Hospital, Boston, Massachusetts [7e]

Robert C. Basner, MD

Professor of Clinical Medicine, Division of Pulmonary, Allergy, and Critical Care Medicine, Columbia University College of Physicians and Surgeons, New York, New York [Appendix]

Buddha Basnyat, MD, MSc, FACP, FRCP (Edinburgh)

Director, Oxford University Clinical Research Unit, Patan Academy of Health Sciences; Medical Director, Nepal International Clinic, Kathmandu, Nepal [476e]

Shari S. Bassuk, ScD

Epidemiologist, Division of Preventive Medicine, Brigham and Women's Hospital, Boston, Massachusetts [413]

John F. Bateman, PhD

Director, Cell Biology, Murdoch Children's Research Institute, Melbourne, Victoria, Australia; Murdoch Children's Research Institute, Parkville, Victoria, Australia [427]

David W. Bates, MD, MSc

Professor of Medicine, Harvard Medical School; Chief, General Internal Medicine and Primary Care Division, Brigham and Women's Hospital; Medical Director, Clinical and Quality Analysis, Partners HealthCare System, Inc., Boston, Massachusetts [12e]

Robert P. Baughman, MD

Department of Internal Medicine, University of Cincinnati Medical Center, Cincinnati, Ohio [390]

M. Flint Beal, MD

University Professor of Neurology and Neuroscience; Neurologist, New York Presbyterian Hospital; Weill Cornell Medical College, New York, New York [444e, 455]

Laurence H. Beck, MD, PhD

Assistant Professor of Medicine, Boston University School of Medicine, Boston, Massachusetts [340]

Christian D. Becker, MD, PhD, FCCP

Assistant Professor, Department of Internal Medicine, Division of Pulmonary, Critical Care and Sleep Medicine, Icahn School of Medicine at Mount Sinai New York, New York [485e]

Nicholas J. Beeching, MA, BM BCh, FRCP, FRACP, FFTM RCPS(Glasg), DCH, DTM&H

Senior Lecturer (Clinical) in Infectious Diseases, Liverpool School of Tropical Medicine; Clinical Director, Tropical and Infectious Disease Unit, Royal Liverpool University Hospital; NIHR Health Protection Research Unit in Emerging and Zoonotic Infections, Liverpool; Honorary Consultant, Public Health England and Honorary Civilian Consultant in Infectious Diseases, Army Medical Directorate, United Kingdom [194e]

Doron Behar, MD, PhD

Institute of Genetics, Rambam Health Care Campus, Haifa, Israel [85e]

Robert S. Benjamin, MD

P. H. and Faye E. Robinson Distinguished Professor of Medicine, Department of Sarcoma Medical Oncology, The University of Texas M.D. Anderson Cancer Center, Houston, Texas [119e]

Michael H. Bennett, MD, MBBS, MM (Clin Epi)

Conjoint Associate Professor in Anesthesia and Hyperbaric Medicine; Faculty of Medicine, University of New South Wales; Academic Head of Department, Wales Anaesthesia, Prince of Wales Hospital, Sydney, Australia [477e]

Edward J. Benz, Jr., MD

Richard and Susan Smith Professor of Medicine; Professor of Genetics, Harvard Medical School; President and CEO, Dana-Farber Cancer Institute; Director and Principal Investigator, Dana-Farber/Harvard Cancer Center; Boston, Massachusetts [127]

Jean Bergounioux, MD, PhD, PhC

Pediatric Intensive Care Unit, Hôpital Raymond-Poincaré, Université de Versailles-Saint Quentin, Garches, France [191]

John L. Berk, MD

Associate Professor of Medicine, Boston University School of Medicine; Clinical Director, Amyloidosis Center, Boston Medical Center, Boston, Massachusetts [137]

Aaron S. Bernstein, MD, MPH

Instructor, Harvard Medical School; Associate Director, Center for Health and the Global Environment, Harvard School of Public Health; Pediatric Hospitalist, Boston Children's Hospital, Boston, Massachusetts [151e]

Joseph R. Betancourt, MD, MPH

Associate Professor of Medicine, Harvard Medical School; Director, The Disparities Solutions Center, Massachusetts General Hospital, Boston, Massachusetts [16e]

Atul K. Bhan, MD, MBBS

Professor of Pathology, Harvard Medical School, Department of Pathology, Massachusetts General Hospital, Boston, Massachusetts [366e]

Shalender Bhasin, MBBS

Professor of Medicine, Harvard Medical School; Director, Research Program in Men's Health: Aging and Metabolism; Director, Boston Claude D. Pepper Older Americans Independence Center; Site Director, Harvard Catalyst Clinical Research Center at BWH, Brigham and Women's Hospital, Boston, Massachusetts [7e, 411]

Deepak L. Bhatt, MD, MPH

Professor of Medicine, Harvard Medical School; Chief of Cardiology, VA Boston Healthcare System; Senior Physician, Brigham and Women's Hospital; Senior Investigator, TIMI Study Group, Boston, Massachusetts [296e, 297e]

Roby P. Bhattacharyya, MD, PhD

Instructor in Medicine, Harvard Medical School; Assistant in Medicine, Division of Infectious Disease, Massachusetts General Hospital, Boston, Massachusetts [146]

David R. Bickers, MD

Carl Truman Nelson Professor and Chair, Department of Dermatology, Columbia University Medical Center, New York, New York [75]

Henry J. Binder, MD

Professor Emeritus of Medicine, Senior Research Scientist, Yale University, New Haven, Connecticut [349, 350e]

William R. Bishai, MD, PhD

Professor and Co-Director, Center for Tuberculosis Research, Department of Medicine, Division of Infectious Diseases, Johns Hopkins University School of Medicine, Baltimore, Maryland [175]

Bruce R. Bistrian, MD, PhD, MPH

Professor of Medicine, Harvard Medical School; Chief, Clinical Nutrition, Beth Israel Deaconess Medical Center, Boston, Massachusetts [98e]

Martin J. Blaser, MD

Muriel and George Singer Professor of Medicine; Professor of Microbiology; Director, Human Microbiome Program, New York University Langone Medical Center, New York, New York [188, 192]

Chantal P. Bleeker-Rovers, MD, PhD

Department of Internal Medicine, Radboud University Nijmegen Medical Center, Nijmegen, The Netherlands [26]

Gijs Bleijenberg, PhD

Professor Emeritus, Expert Centre for Chronic Fatigue, Radboud University Medical Centre, Nijmegen, The Netherlands [464e]

Clara D. Bloomfield, MD

Distinguished University Professor; William G. Pace, III Professor of Cancer Research; Cancer Scholar and Senior Advisor, The Ohio State University Comprehensive Cancer Center; Arthur G. James Cancer Hospital and Richard J. Solove Research Institute, Columbus, Ohio [132]

Richard S. Blumberg, MD

Chief, Division of Gastroenterology, Hepatology and Endoscopy, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts [351]

Jean L. Bolognia, MD

Professor, Department of Dermatology, Yale University School of Medicine, New Haven, Connecticut [72]

Joseph V. Bonventre, MD, PhD

Samuel A. Levine Professor of Medicine, Harvard Medical School; Chief, Renal Division; Chief, Division of Biomedical Engineering, Brigham and Women's Hospital, Boston, Massachusetts [333e, 334]

George J. Bosl, MD

Professor of Medicine, Weill Cornell Medical College; Chair, Department of Medicine; Patrick M. Byrne Chair in Clinical Oncology, Memorial Sloan-Kettering Cancer Center, New York, New York [116]

Joshua A. Boyce, MD

Professor of Medicine and Pediatrics; Albert L. Sheffer Professor of Medicine, Harvard Medical School; Director, Inflammation and Allergic Disease Research Section, Brigham and Women's Hospital, Boston, Massachusetts [376]

Eugene Braunwald, MD, MA (Hon), ScD (Hon), FRCP

Distinguished Hersey Professor of Medicine, Harvard Medical School; Founding Chairman, TIMI Study Group, Brigham and Women's Hospital, Boston, Massachusetts [50, 288, 294]

Irwin M. Braverman, MD

Professor Emeritus; Senior Research Scientist, Department of Dermatology, Yale University School of Medicine, New Haven, Connecticut [72]

Otis W. Brawley, MD, FACP

Professor of Hematology, Medical Oncology, Medicine and Epidemiology, Emory University; Chief Medical and Scientific Officer, American Cancer Society, Atlanta, Georgia [100]

Joel G. Breman, MD, DTPH

Senior Scientific Advisor, Fogarty International Center, National Institutes of Health, Bethesda, Maryland [248, 250e]

George J. Brewer, MD

Morton S. and Henrietta K. Sellner Emeritus, Professor of Human Genetics, Emeritus Professor of Internal Medicine, University of Michigan Medical School; Senior Vice President for Research and Development, Adeona Pharmaceuticals, Inc., Ann Arbor, Michigan [429]

Josephine P. Briggs, MD

Director, National Center for Complementary and Alternative Medicine (NCCAM) at the National Institutes of Health (NIH), Bethesda, Maryland [14e]

F. Richard Bringhurst, MD

Associate Professor of Medicine, Harvard Medical School; Physician, Massachusetts General Hospital, Boston, Massachusetts [423]

Steven M. Bromley, MD

Director, Outpatient Services, Virtua Neuroscience, Voorhees, New Jersey; Director, Bromley Neurology, PC, Audubon, New Jersey [42]

Darron R. Brown, MD

Professor of Medicine, Microbiology and Immunology, Division of Infectious Diseases, Simon Cancer Center, Indiana University School of Medicine, Indianapolis, Indiana [222]

Kevin E. Brown, MD, MRCp, FRCPath

Virus Reference Department, Public Health England, London, United Kingdom [221]

Robert H. Brown, Jr., MD, PhD

Chairman, Department of Neurology, University of Massachusetts Medical School, Worcester, Massachusetts [452, 462e]

Amy E. Bryant, PhD

Affiliate Assistant Professor, University of Washington School of Medicine, Seattle, Washington; Research Scientist, Veterans Affairs Medical Center, Boise, Idaho [179]

Christopher M. Burns, MD

Associate Professor, Department of Medicine, Section of Rheumatology, Geisel School of Medicine at Dartmouth, Dartmouth Hitchcock Medical Center, Lebanon, New Hampshire [431e]

David M. Burns, MD

Professor Emeritus, Department of Family and Preventive Medicine, University of California, San Diego School of Medicine, San Diego, California [470]

Stephen B. Calderwood, MD

Morton N. Swartz, MD Academy Professor of Medicine (Microbiology and Immunobiology), Harvard Medical School; Chief, Division of Infectious Diseases, Massachusetts General Hospital, Boston, Massachusetts [160]

Michael Camilleri, MD

Atherton and Winifred W. Bean Professor; Professor of Medicine, Pharmacology, and Physiology, Mayo Clinic College of Medicine, Rochester, Minnesota [55]

Christopher P. Cannon, MD

Professor of Medicine, Harvard Medical School; Senior Physician, Cardiovascular Division, Brigham and Women's Hospital, Boston, Massachusetts [294]

Jonathan R. Carapetis, MBBS, PhD, FRACP, FAFPHM

Director, Telethon Kids Institute, The University of Western Australia, Crawley, Western Australia [381]

Kathryn M. Carbone, MD

Deputy Scientific Director, Division of Intramural Research, National Institute of Dental and Craniofacial Research, Bethesda, Maryland [231e]

Brian I. Carr, MD, PhD, FRCP

IRCCS de Bellis National Center for GI Diseases, Castellana Grotte, BA, Italy [111]

John D. Carter, MD

Professor of Medicine, Division of Rheumatology; Director, University of South Florida Morsani College of Medicine, Tampa, Florida [384]

Arturo Casadevall, MD, PhD

Chair, Department of Microbiology and Immunology, Albert Einstein College of Medicine, Bronx, New York [239]

Agustin Castellanos, MD, FACC, FAHA

Professor of Medicine; Director, Clinical Electrophysiology, University of Miami Miller School of Medicine, Cardiovascular Division, Miami, Florida [327]

Bartolome R. Celli, MD

Professor of Medicine, Harvard Medical School; Staff Physician, Division of Pulmonary and Critical Care Medicine, Brigham and Women's Hospital, Boston, Massachusetts [323]

Murali Chakinala, MD

Associate Professor of Medicine, Division of Pulmonary and Critical Care Medicine, Washington University School of Medicine, St. Louis, Missouri [279]

Anil Chandraker, MBChB, FRCP

Associate Professor of Medicine, Harvard Medical School; Medical Director of Kidney and Pancreas Transplantation; Interim Director, Schuster Family Transplantation Research Center, Brigham and Women's Hospital, Boston, Massachusetts [337]

Lan X. Chen, MD, PhD

Penn Presbyterian Medical Center, Philadelphia, Pennsylvania [395]

Yuan-Tsong Chen, MD, PhD

Duke University Medical Center, Division of Medical Genetics, Department of Pediatrics, Durham, North Carolina [433e]

Glenn M. Chertow, MD, MPH

Norman S. Coplun/Satellite Healthcare Professor of Medicine; Chief, Division of Nephrology, Stanford University School of Medicine, Palo Alto, California [336]

John S. Child, MD, FACC, FAHA, FASE

Streisand Professor of Medicine and Cardiology; Director, Ahmanson-UCLA Adult Congenital Heart Disease Center; Director, UCLA Adult Noninvasive Cardiodiagnostics Laboratory Ronald Reagan-UCLA Medical Center, Geffen School of Medicine, University of California, Los Angeles (UCLA), Los Angeles, California [282]

Augustine M. K. Choi, MD

Chairman, Department of Medicine, Weill Cornell Medical College, Physician-in-Chief, New York-Presbyterian Hospital-Weill Cornell Medical Center, New York, New York [305, 307, 322]

Raymond T. Chung, MD

Associate Professor of Medicine, Harvard Medical School; Director of Hepatology and Liver Center; Vice Chief, Gastroenterology, Massachusetts General Hospital, Boston, Massachusetts [368]

Jeffrey W. Clark, MD

Associate Professor of Medicine, Harvard Medical School; Medical Director, Clinical Trials Core, Dana-Farber Harvard Cancer Center; Massachusetts General Hospital, Boston, Massachusetts [102e]

Jeffrey I. Cohen, MD

Chief, Laboratory of Clinical Infectious Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [218, 228]

Yehuda Z. Cohen, MD

Clinical Fellow, Department of Medicine, Division of Infectious Diseases and Center for Virology and Vaccine Research, Beth Israel Deaconess Medical Center; Harvard Medical School, Boston, Massachusetts [224]

Ronit Cohen-Poradosu, MD

Senior Physician, Infectious Diseases Unit, Tel Aviv Sourasky Medical Center, Tel Aviv, Israel [201]

Francis S. Collins, MD, PhD

Director, National Institutes of Health, Bethesda, Maryland [101e]

Wilson S. Colucci, MD, FAHA, FACC

Thomas J. Ryan Professor of Medicine, Boston University School of Medicine; Chief of Cardiovascular Medicine, Boston Medical Center, Boston, Massachusetts [289e, 290e]

Laura K. Conlin, PhD

Scientific Director, CytoGenomics Lab, The Children's Hospital of Philadelphia, Assistant Professor of Pathology and Laboratory Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [83e]

Darwin L. Conwell, MD, MS

Professor of Medicine, The Ohio State University College of Medicine; Director, Division of Gastroenterology, Hepatology and Nutrition; The Ohio State University Wexner Medical Center, Columbus, Ohio [370, 371]

Michael J. Corbel, PhD, DSc, FRCPATH

Retired (previously Head, Division of Bacteriology, National Institute for Biological Standards and Control, Hertfordshire, United Kingdom) [194e]

William E. Corcoran, V, MD

Staff Anesthesiologist, Allcare Clinical Associates; Medical Director of Analgesia and Sedation, Mission Health; Patient Safety Officer, Mission Health, Asheville, North Carolina [481e]

Kathleen E. Corey, MD, MPH

Clinical and Research Fellow, Harvard Medical School; Fellow, Gastrointestinal Unit, Massachusetts General Hospital, Boston, Massachusetts [59]

Lawrence Corey, MD

Professor, Medicine and Laboratory Medicine, University of Washington; President Emeritus, Fred Hutchinson Cancer Research Center; Member, Vaccine and Infectious Disease Division; Principal Investigator, HIV Vaccine Trials Network, Fred Hutchinson Cancer Research Center, Seattle, Washington [216]

Jorge Cortes, MD

D. B. Lane Cancer Research Distinguished Professor for Leukemia Research; Deputy Chairman; Section Chief of AML and CML, The University of Texas M.D. Anderson Cancer Center, Houston, Texas [133]

Felicia Cosman, MD

Professor of Medicine, Columbia University College of Physicians and Surgeons, New York, New York [425]

Mark A. Creager, MD

Professor of Medicine, Harvard Medical School; Simon C. Fireman Scholar in Cardiovascular Medicine; Director, Vascular Center, Brigham and Women's Hospital, Boston, Massachusetts [301-303]

Leslie J. Crofford, MD

Professor, Division of Rheumatology and Immunology, Vanderbilt University, Nashville, Tennessee; Chief, Division of Rheumatology, University of Kentucky, Lexington, Kentucky [396]

Jennifer M. Crowell, MD, MPH

Medical Officer, Center for Oncology Prevention Trials Research Group, Division of Cancer Prevention, National Cancer Institute, Bethesda, Maryland [100]

Philip E. Cryer, MD

Professor of Medicine Emeritus, Washington University in St. Louis; Physician, Barnes-Jewish Hospital, St. Louis, Missouri [420]

David Cunningham, MD, MB, ChB, FRCP

Professor, Head of Gastrointestinal/Lymphoma Unit; Director of Clinical Research, Royal Marsden NHS Trust, London, United Kingdom [112]

Gary C. Curhan, MD

Professor of Medicine, Harvard Medical School, Professor of Epidemiology, Harvard School of Public Health, Channing Division of Network Medicine/Renal Division, Brigham and Women's Hospital, Boston, Massachusetts [342]

Brendan D. Curti, MD

Director, Biotherapy Program, Robert W. Franz Cancer Research Center, Providence Portland Medical Center, Portland, Oregon [105]

John J. Cush, MD

Professor of Medicine and Rheumatology, Baylor University Medical Center; Director of Clinical Rheumatology, Baylor Research Institute, Dallas, Texas [393]

Charles A. Czeisler, MD, PhD

Frank Baldino, Jr., PhD Professor of Sleep Medicine, Professor of Medicine and Director, Division of Sleep Medicine, Harvard Medical School; Chief, Division of Sleep and Circadian Disorders, Departments of Medicine and Neurology, Brigham and Women's Hospital, Boston, Massachusetts [38]

Marinos C. Dalakas, MD

Professor of Neurology, University of Athens Medical School, Athens, Greece; Thomas Jefferson University, Philadelphia, Pennsylvania [388]

Josep Dalmau, MD, PhD

ICREA Professor, Institut d'Investigació Biomèdica August Pi i Sunyer, University of Barcelona, Barcelona, Spain; Adjunct Professor, University of Pennsylvania, Philadelphia, Pennsylvania [122]

Daniel F. Danzl, MD

University of Louisville, Department of Emergency Medicine, Louisville, Kentucky [478e, 479e]

Robert B. Daroff, MD

Professor and Chair Emeritus, Department of Neurology, Case Western Reserve University School of Medicine; University Hospitals-Case Medical Center, Cleveland, Ohio [28]

Charles E. Davis, MD

Professor of Pathology and Medicine, Emeritus, University of California, San Diego School of Medicine; Director Emeritus, Microbiology, University of California, San Diego Medical Center, San Diego, California [245e]

Stephen N. Davis, MBBS, FRCP

Theodore E. Woodward Professor and Chairman of the Department of Medicine, University of Maryland School of Medicine; Physician-in-Chief, University of Maryland Medical Center, Baltimore, Maryland [420]

Rafael de Cabo, PhD

Senior Investigator, Experimental Gerontology Section, TGB, National Institute on Aging, National Institutes of Health, Baltimore, Maryland [94e]

Lisa M. DeAngelis, MD

Professor of Neurology, Weill Cornell Medical College; Chair, Department of Neurology, Memorial Sloan Kettering Cancer Center, New York, New York [118]

John Del Valle, MD

Professor and Senior Associate Chair of Medicine, Department of Internal Medicine, University of Michigan School of Medicine, Ann Arbor, Michigan [348]

Marie B. Demay, MD

Professor of Medicine, Harvard Medical School; Physician, Massachusetts General Hospital, Boston, Massachusetts [423]

Bradley M. Denker, MD

Associate Professor of Medicine, Harvard Medical School; Renal Division, Beth Israel Deaconess Medical Center; Chief of Nephrology and Associate Chief of Medical Specialties, Harvard Vanguard Medical Associates, Boston, Massachusetts [61]

David W. Denning, MBBS, FRCP, FRCPath, FMedSci

Professor of Medicine and Medical Mycology; Director, National Aspergillosis Centre, The University of Manchester and Wythenshawe Hospital, Manchester, United Kingdom [241]

Robert J. Desnick, MD, PhD

Dean for Genetics and Genomics, Professor and Chairman Emeritus, Department of Human Genetics and Genomic Sciences, Icahn School of Medicine at Mount Sinai, New York, New York [430]

Richard A. Deyo, MD, MPH

Kaiser Permanente Professor of Evidence-Based Family Medicine, Department of Family Medicine, Department of Medicine, Department of Public Health and Preventive Medicine, Oregon Institute of Occupational Health Sciences; Oregon Health and Science University; Clinical Investigator, Kaiser Permanente Center for Health Research, Portland, Oregon [22]

Betty Diamond, MD

The Feinstein Institute for Medical Research, North Shore LIJ Health System; Center for Autoimmunity and Musculoskeletal Diseases, Manhasset, New York [377e]

Marcelo F. Di Carli, MD

Professor, Department of Radiology, Harvard Medical School; Chief, Division of Nuclear Medicine and Molecular Imaging; Executive Director, Noninvasive Cardiovascular Imaging Program, Brigham and Women's Hospital, Boston, Massachusetts [270e, 271e]

Anna Mae Diehl, MD

Florence McAlister Professor of Medicine; Chief, Division of Gastroenterology, Duke University, Durham, North Carolina [364]

Jules L. Dienstag, MD

Carl W. Walter Professor of Medicine and Dean for Medical Education, Harvard Medical School; Physician, Gastrointestinal Unit, Department of Medicine, Massachusetts General Hospital, Boston, Massachusetts [360-362, 366e, 368]

William P. Dillon, MD

Professor and Executive Vice-Chair; Chief, Section of Neuroradiology, Department of Radiology and Biomedical Imaging, University of California, San Francisco, San Francisco, California [440e, 441e]

Charles A. Dinarello, MD

Professor of Medicine and Immunology, University of Colorado Denver, Aurora, Colorado; Professor of Experimental Medicine, Radboud University Medical Center, Nijmegen, The Netherlands [23]

Raphael Dolin, MD

Maxwell Finland Professor of Medicine (Microbiology and Molecular Genetics), Harvard Medical School; Beth Israel Deaconess Medical Center; Brigham and Women's Hospital, Boston, Massachusetts [215e, 223, 224]

Susan M. Domchek, MD

Basser Professor of Oncology, Abramson Cancer Center, University of Pennsylvania, Philadelphia, Pennsylvania [84]

Richard L. Doty, PhD, MA

Director, Smell and Taste Center; Professor, Department of Otorhinolaryngology: Head and Neck Surgery, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [42]

Vanja C. Douglas, MD

Assistant Professor of Clinical Neurology and Sara and Evan Williams Foundation Endowed Neurohospitalist Chair, University of California, San Francisco, San Francisco, California [29]

Daniel B. Drachman, MD

Professor of Neurology and Neuroscience, W. W. Smith Charitable Trust Professor of Neuroimmunology, Department of Neurology, Johns Hopkins School of Medicine, Baltimore, Maryland [461]

David F. Driscoll, PhD

Associate Professor of Medicine, University of Massachusetts Medical School, Worcester, Massachusetts [98e]

Thomas D. DuBose, Jr., MD, MACP

Emeritus Professor of Internal Medicine and Nephrology, Wake Forest University School of Medicine, Winston-Salem, North Carolina [64e, 66]

J. Stephen Dumler, MD

Professor, Division of Medical Microbiology, Department of Pathology, Johns Hopkins University School of Medicine, Baltimore, Maryland [211]

Andrea Dunaif, MD

Charles F. Kettering Professor of Endocrinology and Metabolism and Vice-Chair for Research, Department of Medicine, Feinberg School of Medicine, Northwestern University, Chicago, Illinois [6e]

Samuel C. Durso, MD, MBA

Mason F. Lord Professor of Medicine; Director, Division of Geriatric Medicine and Gerontology, Johns Hopkins University School of Medicine, Baltimore, Maryland [45, 46e]

Janice P. Dutcher, MD

Associate Director, Cancer Research Foundation of New York, Chappaqua, New York; Former Professor, New York Medical College, Valhalla, New York [331]

Johanna Dwyer, DSc, RD

Jean Mayer USDA Human Nutrition Research Center on Aging; Professor, Tufts Medical Center and Director, Frances Stern Nutrition Center, Tufts Medical Center, Boston, Massachusetts [95e]

Jeffrey S. Dzieczkowski, MD

Physician, St. Alphonsus Regional Medical Center; Medical Director, Coagulation Clinic, Saint Alphonsus Medical Group, International Medicine and Travel Medicine, Boise, Idaho [138e]

Kim A. Eagle, MD

Albion Walter Hewlett Professor of Internal Medicine; Chief of Clinical Cardiology; Director, Frankel Cardiovascular Center, University of Michigan Health System, Ann Arbor, Michigan [9]

James A. Eastham, MD

Chief, Urology Service, Florence and Theodore Baumritter/Enid Ancell Chair of Urologic Oncology, Department of Surgery, Sidney Kimmel Center for Prostate and Urologic Cancers, Memorial Sloan Kettering Cancer Center, New York, New York [115]

Robert H. Eckel, MD

Professor of Medicine, Division of Endocrinology, Metabolism and Diabetes, Division of Cardiology; Professor of Physiology and Biophysics, Charles A. Boettcher, II Chair in Atherosclerosis, University of Colorado School of Medicine, Anschutz Medical Campus, Director Lipid Clinic, University of Colorado Hospital, Aurora, Colorado [422]

John E. Edwards, Jr., MD

Professor of Medicine, David Geffen School of Medicine, University of California, Los Angeles (UCLA), Los Angeles, California; Chief, Division of Infectious Diseases, Harbor/UCLA Medical Center, Torrance, California [235, 240]

David A. Ehrmann, MD

Professor, Department of Medicine, Section of Endocrinology, Diabetes, and Metabolism, The University of Chicago Pritzker School of Medicine, Chicago, Illinois [68]

Andrew J. Einstein, MD, PhD

Victoria and Esther Aboodi Assistant Professor of Medicine; Director, Cardiac CT Research; Co-Director, Cardiac CT and MRI, Department of Medicine, Cardiology Division, Department of Radiology, Columbia University College of Physicians and Surgeons, New York-Presbyterian Hospital, New York, New York [Appendix]

Ezekiel J. Emanuel, MD, PhD

Chair, Department of Medical Ethics and Health Policy, Levy University Professor, Perelman School of Medicine and Wharton School, University of Pennsylvania, Philadelphia, Pennsylvania [10]

John W. Engstrom, MD

Betty Anker Fife Distinguished Professor and Vice-Chairman; Neurology Residency Program Director, University of California, San Francisco, San Francisco, California [22, 454]

Moshe Ephros, MD

Clinical Associate Professor, Faculty of Medicine, Technion-Israel Institute of Technology; Pediatric Infectious Disease Unit, Carmel Medical Center; Haifa, Israel [197]

Jonathan A. Epstein, MD

William Wikoff Smith Professor; Chair, Department of Cell and Developmental Biology; Scientific Director, Penn Cardiovascular Institute, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [265e]

Aaron C. Ermel, MD

Assistant Research Professor; Assistant Professor of Clinical Medicine, Department of Internal Medicine, Division of Infectious Disease, Indiana University School of Medicine, Indianapolis, Indiana [222]

Tim Evans, MD, PhD

Senior Director, Health, Nutrition and Population, The World Bank Group, Washington, DC [13e]

Christopher H. Fanta, MD

Professor of Medicine, Harvard Medical School; Pulmonary and Critical Care Division, Brigham and Women's Hospital; Director, Partners Asthma Center, Boston, Massachusetts [48]

Paul Farmer, MD, PhD

Kolokotronis University Professor, Harvard University; Chair, Department of Global Health and Social Medicine, Harvard Medical School; Chief, Division of Global Health Equity, Brigham and Women's Hospital; Co-Founder, Partners In Health, Boston, Massachusetts [2]

Anthony S. Fauci, MD

Chief, Laboratory of Immunoregulation; Director, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [1, 225e, 226, 261e, 372e, 385, 386e]

Murray J. Favus, MD

Professor of Medicine, Department of Medicine, Section of Endocrinology, Diabetes and Metabolism, Director Bone Program, University of Chicago Pritzker School of Medicine, Chicago, Illinois [426e]

David P. Faxon, MD

Vice Chair of Medicine for Strategic Planning, Department of Medicine, Brigham and Women's Hospital; Senior Lecturer, Harvard Medical School, Boston, Massachusetts [272, 296e, 297]

Darren R. Feldman, MD

Associate Professor in Medicine, Weill Cornell Medical Center; Assistant Attending, Genitourinary Oncology Service, Memorial Sloan-Kettering Cancer Center, New York, New York [116]

David T. Felson, MD, MPH

Professor of Medicine and Epidemiology; Chair, Clinical Epidemiology Unit, Boston University School of Medicine, Boston, Massachusetts [394]

Luigi Ferrucci, MD, PhD

Scientific Director, National Institute of Aging, National Institutes of Health, Baltimore, Maryland [11]

Howard L. Fields, MD, PhD

Professor, Department of Neurology, University of California, San Francisco, San Francisco, California [18]

Gregory A. Filice, MD

Professor of Medicine, Medical School, and Adjunct Professor of Epidemiology and Community Health, School of Public Health, University of Minnesota; Chief, Infectious Disease Section, Veterans Affairs Healthcare System, Minneapolis, Minnesota [199]

Robert W. Finberg, MD

Chair, Department of Medicine, University of Massachusetts Medical School, Worcester, Massachusetts [104, 169]

Joyce Fingerroth, MD

Professor of Medicine and MAPS, Division of Infectious Disease, University of Massachusetts Medical School, Worcester, Massachusetts [169]

Kurt Fink, MD

Palo Alto Medical Foundation, Palo Alto, California [481e]

Alain Fischer, MD, PhD

Director of INSERM U768; Director of Imagine Institute; Professor of Immunology and Pediatric Hematology; Université Paris Descartes, Paris, France [374, 375e]

Jeffrey S. Flier, MD

Caroline Shields Walker Professor of Medicine and Dean, Harvard Medical School, Boston, Massachusetts [415e]

Agnes B. Fogo, MD

John L. Shapiro Professor of Pathology; Professor of Medicine and Pediatrics, Vanderbilt University Medical Center, Nashville, Tennessee [62e]

Larry C. Ford, MD

Clinical Infectious Diseases, Intermountain Healthcare, Provo, Utah [44]

Jane E. Freedman, MD

Professor of Medicine, University of Massachusetts Medical School, Worcester, Massachusetts [142]

Roy Freeman, MD

Professor of Neurology, Harvard Medical School; Director, Center for Autonomic and Peripheral Nerve Disorders, Beth Israel Deaconess Medical Center, Boston, Massachusetts [27]

Gyorgy Frendl, MD, PhD, FCCM

Assistant Professor; Director of Surgical Critical Care Research Center, Department of Anesthesiology, Perioperative Critical Care and Pain Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [481e]

Carl E. Freter, MD, PhD, FACP

Professor of Medicine; Director, Division of Hematology and Oncology; Associate Director, Cancer Center, Saint Louis University, St. Louis, Missouri [125]

Lawrence S. Friedman, MD

Professor of Medicine, Harvard Medical School; Professor of Medicine, Tufts University School of Medicine; Assistant Chief of Medicine, Massachusetts General Hospital, Boston, Massachusetts; Anton R. Fried, MD Chair, Department of Medicine, Newton-Wellesley Hospital, Newton, Massachusetts [59]

Sonia Friedman, MD

Associate Professor of Medicine, Harvard Medical School; Associate Physician, Brigham and Women's Hospital, Boston, Massachusetts [351]

Anne L. Fuhlbrigge, MD, MS

Assistant Professor of Medicine, Harvard Medical School, Pulmonary and Critical Care Division; Brigham and Women's Hospital, Boston, Massachusetts [307]

Andre D. Furtado, MD

Assistant Professor, Department of Radiology, School of Medicine, University of Pittsburgh, Pittsburgh, Pennsylvania [441e]

Nicholas B. Galifianakis, MD, MPH

Assistant Clinical Professor, Surgical Movement Disorders Center, Department of Neurology, University of California, San Francisco, San Francisco, California [33e]

John I. Gallin, MD

Director, Clinical Center, National Institutes of Health, Bethesda, Maryland [80]

Charlotte A. Gaydos, DrPh

Professor of Medicine, Johns Hopkins University, Division of Infectious Diseases, Baltimore, Maryland [213]

J. Michael Gaziano, MD, MPH

Professor of Medicine, Harvard Medical School; Chief, Division of Aging, Brigham and Women's Hospital; Director, Massachusetts Veterans Epidemiology Center, Boston VA Healthcare System, Boston, Massachusetts [266e]

Thomas A. Gaziano, MD, MSc

Assistant Professor of Medicine, Harvard Medical School; Assistant Professor, Health Policy and Management, Center for Health Decision Sciences, Harvard School of Public Health; Faculty Co-Leader, Chronic and Cardiovascular Diseases Working Group, Harvard Institute for Global Health, Harvard University; Associate Physician in Cardiovascular Medicine, Department of Cardiology, Brigham and Women's Hospital, Boston, Massachusetts [266e]

Susan L. Gearhart, MD

Associate Professor, Surgery, Johns Hopkins Medical Institutions, Baltimore, Maryland [353]

Robert H. Gelber, MD

Clinical Professor of Medicine and Dermatology, University of California, San Francisco, San Francisco, California [203]

Jeffrey M. Gelfand, MD, MAS

Assistant Professor of Clinical Neurology, Department of Neurology, University of California, San Francisco, San Francisco, California [29]

Alfred L. George, Jr., MD

Magerstadt Professor and Chair, Department of Pharmacology, Feinberg School of Medicine, Northwestern University, Chicago, Illinois [332e]

Dale N. Gerding, MD

Professor of Medicine, Department of Medicine, Loyola University Chicago Stritch School of Medicine, Maywood, Illinois; Research Physician, Edward Hines Jr. Veterans Affairs Hospital, Hines, Illinois [161]

Michael D. Geschwind, MD, PhD

Associate Professor of Neurology, Memory and Aging Center, University of California, San Francisco, San Francisco, California [33e]

Marc G. Ghany, MD, MHSC

Staff Physician, Liver Diseases Branch, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, Maryland [357]

Michael Giladi, MD, MSc

Associate Professor of Medicine, Sackler Faculty of Medicine, Tel Aviv University; The Infectious Disease Unit and the Bernard Pridan Laboratory for Molecular Biology of Infectious Diseases, Tel Aviv Medical Center, Tel Aviv, Israel [197]

Roger I. Glass, MD, PhD

Director, Fogarty International Center, Bethesda, Maryland [227]

Eli Glatstein, MD

Professor and Vice Chairman, Department of Radiation Oncology, Hospital of the University of Pennsylvania, Philadelphia, Pennsylvania [263e]

Peter J. Goadsby, MD, PhD, DSc, FRACp, FRCP

Professor, NIHR-Wellcome Trust Clinical Research Facility, King's College, London, United Kingdom; Professor, Department of Neurology, University of California, San Francisco, San Francisco, California [21, 447]

Morton F. Goldberg, MD, FACS, FAOS

Director Emeritus and Joseph Green Professor of Ophthalmology, Wilmer Eye Institute, Johns Hopkins University School of Medicine and Johns Hopkins Hospital, Baltimore, Maryland [40e]

Ary L. Goldberger, MD

Professor of Medicine, Harvard Medical School; Wyss Institute for Biologically Inspired Engineering, Harvard University; Beth Israel Deaconess Medical Center, Boston, Massachusetts [268, 269e, 278e]

David Goldblatt, MB, ChB, PhD

Professor of Vaccinology and Immunology; Consultant in Paediatric Immunology; Director of Clinical Research and Development; Director, NIHR Biomedical Research Centre, Institute of Child Health; University College London; Great Ormond Street Hospital for Children NHS Trust, London, United Kingdom [171]

Samuel Z. Goldhaber, MD

Professor of Medicine, Harvard Medical School; Director, Thrombosis Research Group, Brigham and Women's Hospital, Boston, Massachusetts [300]

Ralph Gonzales, MD, MSPH

Professor of Medicine, University of California, San Francisco, San Francisco, California [44]

Douglas S. Goodin, MD

Professor, Department of Neurology, School of Medicine, University of California, San Francisco, San Francisco, California [458]

Jeffrey I. Gordon, MD

Dr. Robert J. Glaser Distinguished University Professor and Director, Center for Genome Sciences and Systems Biology, Washington University School of Medicine, St. Louis, Missouri [86e]

Maria Luisa Gorno-Tempini, MD, PhD

Professor, Department of Neurology; Language Neurobiology Lab, Memory and Aging Center; Dyslexia Center, University of California, San Francisco, San Francisco, California [37e]

Peter A. Gottlieb, MD

Professor of Pediatrics and Medicine, Barbara Davis Center, University of Colorado School of Medicine, Aurora, Colorado [409]

Gregory A. Grabowski, MD

Adjunct Professor of Pediatrics and Molecular Genetics, Biochemistry, and Microbiology, University of Cincinnati College of Medicine; Division of Human Genetics Cincinnati Children's Hospital Medical Center, Cincinnati, Ohio; Chief Scientific Officer, Synageva BioPharma Corp., Lexington, Massachusetts [432e]

Yonatan H. Grad, MD, PhD

Assistant Professor of Immunology and Infectious Diseases, Harvard School of Public Health; Associate Physician, Division of Infectious Diseases, Brigham and Women's Hospital, Boston, Massachusetts [146]

Christine Grady, RN, PhD

Chief, Department of Bioethics, National Institutes of Health Clinical Center, Bethesda, Maryland [17e]

Alexander R. Green, MD, MPH

Assistant Professor of Medicine, Harvard Medical School; Associate Director, The Disparities Solutions Center, Massachusetts General Hospital, Boston, Massachusetts [16e]

Norton J. Greenberger, MD

Clinical Professor of Medicine, Harvard Medical School; Senior Physician, Division of Gastroenterology, Brigham and Women's Hospital, Boston, Massachusetts [369-371]

Michael F. Greene, MD

Professor of Obstetrics, Gynecology and Reproductive Biology, Harvard Medical School; Vincent Department of Obstetrics and Gynecology, Massachusetts General Hospital, Boston, Massachusetts [124e]

Daryl R. Gress, MD, FAAN, FCCM, FNCS

Associate Professor of Neurology, University of Virginia, Charlottesville, Virginia [330]

Rasim Gucalp, MD

Professor of Clinical Medicine, Albert Einstein College of Medicine; Associate Chairman for Educational Programs, Department of Oncology; Director, Hematology/Oncology Fellowship, Montefiore Medical Center, Bronx, New York [331]

Kalpana Gupta, MD, MPH

Associate Professor, Department of Medicine, Boston University School of Medicine; Chief, Section of Infectious Diseases, VA Boston Healthcare System, Boston, Massachusetts [162]

John G. Haaga, PhD

Deputy Associate Director, Behavioral and Social Research Program, National Institute on Aging, National Institutes of Health, Bethesda, Maryland [93e]

Chadi A. Hage, MD

Assistant Professor of Medicine, Thoracic Transplant Program, Indiana University Health, Indianapolis, Indiana [236]

Bevra Hannahs Hahn, MD

Professor Emerita Division of Rheumatology, University of California, Los Angeles, Los Angeles, California [378]

Colin N. Haile, MD, PhD

Assistant Professor, Menninger Department of Psychiatry and Behavioral Sciences, Baylor College of Medicine; Michael E. DeBakey VA Medical Center, Houston, Texas [468e]

Janet E. Hall, MD, MSc

Professor of Medicine, Harvard Medical School and Associate Chief, Reproductive Endocrine Unit, Massachusetts General Hospital, Boston, Massachusetts [69, 412, 414]

Jesse B. Hall, MD, FCCP

Professor of Medicine, Anesthesia and Critical Care; Chief, Section of Pulmonary and Critical Care Medicine, University of Chicago, Chicago, Illinois [321]

Scott A. Halperin, MD

Professor of Pediatrics and Microbiology and Immunology Head, Pediatric Infectious Diseases, Director, Canadian Center for Vaccinology, Dalhousie University, Halifax, Nova Scotia, Canada [185]

R. Doug Hardy, MD

Infectious Diseases Specialists, PA; Medical City Dallas Hospital and Medical City Children's Hospital, Dallas; Baylor Regional Medical Center, Plano, Texas [212]

Rudy A. Hartskeerl, PhD

Director WHO/FAO/OIE and National Leptospirosis Reference Centre, KIT Biomedical Research, KIT (Royal Tropical Institute), Amsterdam, The Netherlands [208]

William L. Hasler, MD

Professor, Division of Gastroenterology, University of Michigan Health System, Ann Arbor, Michigan [54, 344]

Stephen L. Hauser, MD

Robert A. Fishman Distinguished Professor and Chairman, Department of Neurology, University of California, San Francisco, San Francisco, California [1, 437, 443e, 444e, 455, 456, 458, 460]

Barton F. Haynes, MD

Frederic M. Hanes Professor of Medicine and Immunology, Departments of Medicine and Immunology; Director, Duke Human Vaccine Institute, Duke University School of Medicine, Durham, North Carolina [372e]

Douglas C. Heimbarger, MD, MS

Professor of Medicine, Associate Director for Education and Training, Vanderbilt Institute for Global Health, Vanderbilt University School of Medicine, Nashville, Tennessee [97]

J. Claude Hemphill, III, MD, MAS

Professor of Neurology and Neurological Surgery, University of California, San Francisco; Chief of Neurology, San Francisco General Hospital, San Francisco, California [330, 446]

Patrick H. Henry, MD

Clinical Adjunct Professor of Medicine, University of Iowa, Iowa City, Iowa [79]

Katherine A. High, MD

William H. Bennett Professor of Pediatrics, Perelman School of Medicine, University of Pennsylvania; Investigator, Howard Hughes Medical Institute, The Children's Hospital of Philadelphia, Philadelphia, Pennsylvania [91e, 141]

Christine E. Hill-Kayser, MD

Assistant Professor of Radiation Oncology, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [263e]

Ikuo Hirano, MD

Professor of Medicine, Division of Gastroenterology, Northwestern University Feinberg School of Medicine, Chicago, Illinois [53, 347]

Martin S. Hirsch, MD

Professor of Medicine, Harvard Medical School; Professor of Immunology and Infectious Diseases, Harvard School of Public Health; Physician, Massachusetts General Hospital, Boston, Massachusetts [219]

Helen H. Hobbs, MD

Professor, Internal Medicine and Molecular Genetics, University of Texas Southwestern Medical Center; Investigator, Howard Hughes Medical Institute, Dallas, Texas [421]

Judith S. Hochman, MD

Harold Snyder Family Professor of Cardiology, Clinical Chief, Leon Charney Division of Cardiology, Co-Director, NYU-HHC Clinical and Translational Science Institute; Director, Cardiovascular Clinical Research Center, New York University School of Medicine, New York, New York [326]

A. Victor Hoffbrand, DM

Emeritus Professor of Haematology, University College, London; Honorary Consultant Haematologist, Royal Free Hospital, London, United Kingdom [128]

L. John Hoffer, MD, PhD

Professor, Faculty of Medicine, McGill University; Senior Physician, Divisions of Internal Medicine and Endocrinology, Lady Davis Institute for Medical Research, Jewish General Hospital, Montreal, Quebec, Canada [98e]

Charles W. Hoge, MD

Senior Scientist, Center for Psychiatry and Neuroscience, Walter Reed Army Institute of Research, Silver Spring, Maryland [471e]

Elizabeth L. Hohmann, MD

Associate Professor of Medicine and Infectious Diseases, Harvard Medical School; Massachusetts General Hospital, Boston, Massachusetts [176]

Steven M. Holland, MD

Chief, Laboratory of Clinical Infectious Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [80, 204]

King K. Holmes, MD, PhD

Chair, Global Health; Professor of Medicine and Global Health; Adjunct Professor, Epidemiology; Director, Center for AIDS and STD; University of Washington School of Medicine; Head, Infectious Diseases Section, Harborview Medical Center, Seattle, Washington [163]

Jay H. Hoofnagle, MD

Director, Liver Diseases Research Branch, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Bethesda, Maryland [357]

David C. Hooper, MD

Professor, Harvard Medical School; Chief, Infection Control Unit; Associate Chief, Division of Infectious Diseases, Massachusetts General Hospital, Boston, Massachusetts [170]

Robert J. Hopkin, MD

Associate Professor, Cincinnati Children's Hospital Medical Center, Cincinnati, Ohio [432e]

Leora Horn, MD, MSc

Assistant Professor, Division of Hematology and Medical Oncology, Vanderbilt University School of Medicine, Nashville, Tennessee [107]

Jonathan C. Horton, MD, PhD

William F. Hoyt Professor of Neuro-ophthalmology, Professor of Ophthalmology, Neurology and Physiology, University of California, San Francisco School of Medicine, San Francisco, California [39]

Howard Hu, MD, MPH, ScD

Dean; Professor of Environmental Health, Epidemiology and Global Health, Dalla Lana School of Public Health; Professor of Medicine, University of Toronto, Toronto, Ontario, Canada [472e]

Deborah T. Hung, MD, PhD

Associate Professor of Microbiology and Molecular Genetics, Assistant Professor of Medicine, Harvard Medical School; Brigham and Women's Hospital; Massachusetts General Hospital, Boston, Massachusetts; Co-director, Infectious Disease Initiative, Broad Institute of Harvard University and Massachusetts Institute of Technology, Cambridge, Massachusetts [146]

Sharon A. Hunt, MD, FACC

Professor of Medicine, Cardiovascular Medicine, Stanford University, Palo Alto, California [281]

Charles G. Hurst, MD

Chief, Chemical Casualty Care Division, United States Medical Research Institute of Chemical Defense, APG-Edgewood Area, Maryland [262e]

Ashraf S. Ibrahim, PhD

Professor, Department of Medicine, Geffen School of Medicine, University of California, Los Angeles (UCLA); Division of Infectious Diseases, Los Angeles Biomedical Research Institute at Harbor-UCLA Medical Center, Torrance, California [242]

David H. Ingbar, MD

Professor of Medicine, Pediatrics, and Physiology; Director, Pulmonary Allergy, Critical Care and Sleep Division, University of Minnesota School of Medicine, Minneapolis, Minnesota [326]

Alan C. Jackson, MD, FRCPC

Professor of Medicine (Neurology) and of Medical Microbiology, University of Manitoba; Section Head of Neurology, Winnipeg Regional Health Authority, Winnipeg, Manitoba, Canada [232]

Lisa A. Jackson, MD, MPH

Senior Investigator, Group Health Research Institute, Seattle, Washington [148]

Danny O. Jacobs, MD, MPH, FACS

Executive Vice President, Provost, and Dean of the School of Medicine; Thomas N. and Gleaves T. James Distinguished Chair, The University of Texas Medical Branch at Galveston, Galveston, Texas [20, 355, 356]

Richard F. Jacobs, MD

Robert H. Fiser, Jr., MD Endowed Chair in Pediatrics; Professor and Chairman, Department of Pediatrics, University of Arkansas for Medical Sciences; President, Arkansas Children's Hospital Research Institute, Little Rock, Arkansas [195]

J. Larry Jameson, MD, PhD

Robert G. Dunlop Professor of Medicine; Dean, Perelman School of Medicine at the University of Pennsylvania; Executive Vice President, University of Pennsylvania for the Health System, Philadelphia, Pennsylvania [1, 56, 82, 84, 121, 399-403, 405, 410, 411, 436e]

Robert T. Jensen, MD

Chief, Cell Biology Section, National Institutes of Diabetes, Digestive and Kidney Diseases, National Institutes of Health, Bethesda, Maryland [113]

Roy M. John, MBBS, PhD, FRCP

Assistant Professor of Medicine, Harvard Medical School; Department of Medicine, Brigham and Women's Hospital, Boston, Massachusetts [277]

Savio John, MD

Assistant Professor of Medicine, Division of Gastroenterology and Hepatology, State University of New York Upstate Medical University, Syracuse, New York [58]

David H. Johnson, MD

Donald W. Seldin Distinguished Chair in Internal Medicine; Professor and Chairman, Department of Internal Medicine, University of Texas Southwestern School of Medicine, Dallas, Texas [107]

James R. Johnson, MD

Professor of Medicine, University of Minnesota, Minneapolis, Minnesota [186]

Stuart Johnson, MD

Associate Professor of Medicine, Loyola University Chicago Stritch School of Medicine; Staff Physician, Edward Hines Jr. VA Hospital, Hines, Illinois [161]

S. Clairborne Johnston, MD, PhD

Dean, Dell Medical School; Frank Denius Distinguished Dean's Chair in Medical Leadership; Vice President for Medical Affairs, University of Texas, Austin, Austin, Texas [446]

S. Andrew Josephson, MD

Associate Professor; Vice Chairman, Department of Neurology, University of California, San Francisco, San Francisco, California [34, 329e, 463e]

Harald Jüppner, MD

Professor of Pediatrics, Endocrine Unit and Pediatric Nephrology Unit, Massachusetts General Hospital, Boston, Massachusetts [424]

Peter J. Kahrilas, MD

Gilbert H. Marquardt Professor of Medicine, Feinberg School of Medicine, Northwestern University, Chicago, Illinois [53, 347]

Gail Kang, MD

San Francisco, California [33e]

Hagop Kantarjian, MD

Chairman, Leukemia Department; Professor of Leukemia, The University of Texas M.D. Anderson Cancer Center, Houston, Texas [133]

Adolf W. Karchmer, MD

Professor of Medicine, Harvard Medical School, Division of Infectious Diseases, Beth Israel Deaconess Medical Center, Boston, Massachusetts [155]

Dennis L. Kasper, MD, MA

William Ellery Channing Professor of Medicine, Professor of Microbiology and Immunobiology, Department of Microbiology and Immunobiology, Harvard Medical School; Division of Infectious Diseases, Brigham and Women's Hospital, Boston, Massachusetts [1, 144, 147, 159, 183e, 201]

Lloyd H. Kasper, MD

Professor of Microbiology/Immunology and Medicine, Geisel School of Medicine, Dartmouth College, Hanover, New Hampshire [253]

Daniel L. Kastner, MD, PhD

Scientific Director, National Human Genome Research Institute, National Institutes of Health, Bethesda, Maryland [392]

Carol A. Kauffman, MD

Professor of Internal Medicine, University of Michigan Medical School; Chief, Infectious Diseases Section, Veterans Affairs Ann Arbor Healthcare System, Ann Arbor, Michigan [243]

Elaine T. Kaye, MD

Assistant Clinical Professor of Dermatology, Harvard Medical School; Boston Children's Hospital, Boston, Massachusetts [24, 25e]

Kenneth M. Kaye, MD

Associate Professor of Medicine, Harvard Medical School; Division of Infectious Diseases, Brigham and Women's Hospital, Boston, Massachusetts [24, 25e]

John A. Kessler, MD

Davee Professor of Stem Cell Biology, Department of Neurology, Feinberg School of Medicine, Northwestern University, Chicago, Illinois [90e]

Jay S. Keystone, MD, FRCPC, MSc(CTM)

Professor of Medicine, University of Toronto, Toronto, Ontario, Canada [149]

Sundeep Khosla, MD

Professor of Medicine and Physiology, College of Medicine, Mayo Clinic, Rochester, Minnesota [65]

Elliott Kieff, MD, PhD

Harriet Ryan Albee Professor of Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [214e]

Anthony A. Killeen, MD, PhD

Professor, Department of Laboratory Medicine and Pathology, University of Minnesota, Minneapolis, Minnesota [480e]

Kami Kim, MD

Professor, Departments of Medicine, Pathology, and Microbiology and Immunology, Albert Einstein College of Medicine, Bronx, New York [253]

Charles H. King, MD, MS

Professor, Center for Global Health and Diseases, School of Medicine, Case Western Reserve University, Cleveland, Ohio [259]

Lindsay King, MD, MPH

Advanced Transplant/Hepatology Fellow, Department of Medicine, Gastrointestinal Unit, Massachusetts General Hospital, Boston, Massachusetts [483e]

Talmadge E. King, Jr., MD

Professor and Chair, Department of Medicine, University of California, San Francisco, San Francisco, California [315]

Louis V. Kirchhoff, MD, MPH

Professor, Departments of Internal Medicine (Infectious Diseases) and Epidemiology, University of Iowa; Staff Physician, Department of Veterans Affairs Medical Center, Iowa City, Iowa [252]

Priya S. Kishnani, MD

Professor of Pediatrics, Division Chief, Medical Genetics, Duke University Medical Center, Durham, North Carolina [433e]

Rob Knight, PhD

Professor, Howard Hughes Medical Institute; Departments of Chemistry and Biochemistry and Computer Science, Biofrontiers Institute, University of Colorado, Boulder, Colorado [86e]

Minoru S. H. Ko, MD, PhD

Mitsunada Sakaguchi Professor and Chair, Department of Systems Medicine, Keio University School of Medicine, Tokyo, Japan [88]

Barbara A. Konkle, MD

Professor of Medicine, Hematology, University of Washington; Director, Translational Research, Puget Sound Blood Center, Seattle, Washington [78, 140]

Peter Kopp, MD

Associate Professor, Division of Endocrinology, Metabolism and Molecular Science and Center for Genetic Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois [82]

Walter J. Koroshetz, MD

National Institute of Neurological Disorders and Stroke, National Institutes of Health, Bethesda, Maryland [165]

Thomas R. Kosten, MD

J. H. Waggoner Professor of Psychiatry, Pharmacology, Immunology, Neuroscience, Baylor College of Medicine, Houston, Texas [468e]

Theodore A. Kotchen, MD

Professor Emeritus, Department of Medicine; Associate Dean for Clinical Research, Medical College of Wisconsin, Milwaukee, Wisconsin [298]

Camille Nelson Kotton, MD, FIDSA

Clinical Director, Transplant and Immunocompromised Host Infectious Diseases, Infectious Diseases Division, Massachusetts General Hospital; Harvard Medical School, Boston, Massachusetts [219]

Phyllis E. Kozarsky, MD

Professor of Medicine and Infectious Diseases, Emory University School of Medicine, Atlanta, Georgia [149]

Barnett S. Kramer, MD, MPH, FACP

Director, Division of Cancer Prevention, National Cancer Institute, Bethesda, Maryland [100]

Joel Kramer, PsyD

Professor of Neuropsychology in Neurology; Director of Neuropsychology, Memory and Aging Center, University of California, San Francisco, San Francisco, California [37e]

Stephen M. Krane, MD

Persis, Cyrus and Marlow B. Harrison Distinguished Professor of Medicine, Harvard Medical School; Massachusetts General Hospital, Boston, Massachusetts [423]

Alexander Kratz, MD, MPH, PhD

Associate Professor of Clinical Pathology and Cell Biology, Columbia University College of Physicians and Surgeons; Director, Core Laboratory, Columbia University Medical Center and the New York Presbyterian Hospital; Director, the Allen Hospital Laboratory, New York, New York [Appendix]

Peter J. Krause, MD

Senior Research Scientist, Yale School of Public Health; Yale School of Medicine, New Haven, Connecticut [249]

John P. Kress, MD

Professor of Medicine, Director, Medical Intensive Care Unit, University of Chicago, Chicago, Illinois [321]

Stephen Krieger, MD

Assistant Professor, Department of Neurology; Director, Neurology Residency Program, Icahn School of Medicine at Mount Sinai; Attending Physician, The Corinne Goldsmith Dickinson Center for MS, New York, New York [486e]

Patricia A. Kritek, MD, EdM

Associate Professor, Division of Pulmonary and Critical Care Medicine, University of Washington, Seattle, Washington [48, 305, 308e]

Henry M. Kronenberg, MD

Professor of Medicine, Harvard Medical School; Chief, Endocrine Unit, Massachusetts General Hospital, Boston, Massachusetts [423]

Jens H. Kuhn, MD, PhD, MS

Principal, Tunnell Government Services (TGS), Inc.; Lead Virologist, Integrated Research Facility at Fort Detrick (IRF-Frederick); TGS IRF-Frederick Team Leader, NIH/NIAID/DCR, Fort Detrick, Frederick, Maryland [233, 234]

Robert F. Kushner, MD, MS

Professor of Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois [416]

Raymond Y. Kwong, MD, MPH

Associate Professor of Medicine, Harvard Medical School; Director of Cardiac Magnetic Resonance Imaging, Cardiovascular Division, Department of Medicine, Brigham and Women's Hospital, Boston, Massachusetts [270e, 271e]

Loren Laine, MD

Professor of Medicine, Yale University School of Medicine, New Haven, Connecticut; VA Connecticut Healthcare System, West Haven, Connecticut [57]

Neil K. Lakdawala, MD

Instructor in Medicine, Harvard Medical School; Associate Physician, Cardiovascular Medicine, Brigham and Women's Hospital; Boston VA Healthcare; Boston, Massachusetts [287]

Anil K. Lalwani, MD

Professor and Vice Chair for Research; Director, Division of Otolaryngology, Neurotology and Skull Base Surgery; Director, Columbia Cochlear Implant Center, Columbia University College of Physicians and Surgeons, New York, New York [43]

H. Clifford Lane, MD

Clinical Director, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [226, 261e]

Carol A. Langford, MD, MHS

Harold C. Schott Endowed Chair; Director, Center for Vasculitis Care and Research, Department of Rheumatic and Immunologic Diseases, Cleveland Clinic, Cleveland, Ohio [385, 386e, 389, 397, 398]

Regina C. LaRocque, MD, MPH

Assistant Professor of Medicine, Harvard Medical School; Assistant Physician, Massachusetts General Hospital, Boston, Massachusetts [160]

Wei C. Lau, MD, FAHA

Emeritus Associate Professor, Department of Anesthesiology, Section Cardiovascular Anesthesiology, University of Michigan Health System Cardiovascular Center, Ann Arbor, Michigan [9]

Leslie P. Lawley, MD

Assistant Professor, Department of Dermatology, School of Medicine, Emory University, Atlanta, Georgia [71]

Thomas J. Lawley, MD

William P. Timmie Professor of Dermatology, Dean, Emory University School of Medicine, Atlanta, Georgia [70, 71, 73, 76e]

David G. Le Couteur, MD, PhD, FRACP

Professor of Geriatric Medicine, Director of the Centre for Education and Research on Ageing, University of Sydney and Sydney Research, Sydney, Australia [94e]

William M. Lee, MD

Professor of Internal Medicine; Meredith Mosle Chair in Liver Diseases, University of Texas Southwestern Medical Center at Dallas, Dallas, Texas [361]

Charles Lei, MD

Assistant Professor, Department of Emergency Medicine, Vanderbilt University Medical Center, Nashville, Tennessee [474]

Jane A. Leopold, MD

Associate Professor of Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [272, 297e]

Nelson Leung, MD

Associate Professor of Medicine, Division of Nephrology and Hypertension, Division of Hematology, Mayo Clinic Rochester, Rochester, Minnesota [341]

Bruce D. Levy, MD

Associate Professor of Medicine, Harvard Medical School; Pulmonary and Critical Care Medicine, Brigham and Women's Hospital, Boston, Massachusetts [322]

Julia B. Lewis, MD

Professor, Department of Medicine, Division of Nephrology, Vanderbilt University Medical Center, Nashville, Tennessee [338]

Peter Libby, MD

Mallinckrodt Professor of Medicine, Harvard Medical School; Chief, Cardiovascular Medicine, Brigham and Women's Hospital, Boston, Massachusetts [265e, 291e, 292e]

Richard W. Light, MD

Professor of Medicine, Division of Allergy, Pulmonary, and Critical Care Medicine, Vanderbilt University, Nashville, Tennessee [316, 317]

Julie Lin, MD, MPH

Lecturer on Medicine, Harvard Medical School; Associate Physician, Renal Division, Brigham and Women's Hospital, Boston, Massachusetts, Boston, Massachusetts [61]

Yusen E. Lin, PhD, MBA

Professor and Director, Center for Environmental Laboratory Services; National Kaohsiung Normal University, Kaohsiung, Taiwan [184]

Robert Lindsay, MD, PhD

Chief, Internal Medicine; Professor of Clinical Medicine, Helen Hayes Hospital, West Haverstraw, New York [425]

Marc E. Lippman, MD, MACP, FRCP

Kathleen and Stanley Glaser Professor, Department of Medicine, Deputy Director, Sylvester Comprehensive Cancer Center, University of Miami Miller School of Medicine, Miami, Florida [108]

Peter E. Lipsky, MD

Charlottesville, Virginia [377e]

Kathleen D. Liu, MD, PhD, MAS

Associate Professor, Division of Nephrology, Department of Medicine, Division of Critical Care Medicine, Department of Anesthesiology, University of California, San Francisco, San Francisco, California [336]

Bernard Lo, MD

President, The Greenwall Foundation, New York; Professor of Medicine Emeritus and Director Emeritus of the Program in Medical Ethics, University of California, San Francisco, San Francisco, California [17e]

Dan L. Longo, MD

Professor of Medicine, Harvard Medical School; Senior Physician, Brigham and Women's Hospital; Deputy Editor, *New England Journal of Medicine*, Boston, Massachusetts [1, 77, 79, 81e, 89e, 99, 102e, 103e, 121, 123e, 124e, 125, 134, 135e, 136, 225e]

Nicola Longo, MD, PhD

Professor and Chief, Division of Medical Genetics, Departments of Pediatrics and Pathology; Medical Co-Director, Biochemical Genetics Laboratory, ARUP Laboratories, University of Utah, Salt Lake City, Utah [434e, 435e]

Joseph Loscalzo, MD, PhD

Hersey Professor of the Theory and Practice of Medicine, Harvard Medical School; Chairman, Department of Medicine; Physician-in-Chief, Brigham and Women's Hospital, Boston, Massachusetts [1, 49-52, 87e, 142, 264, 265e, 267, 283-287, 293, 295, 301-304]

Christine M. Lovly, MD, PhD

Academic, Vanderbilt Ingram Cancer Center, Vanderbilt University School of Medicine, Nashville, Tennessee [107]

Phillip A. Low, MD, FRACP, FRCP(Hon)

Robert D. and Patricia E. Kern Professor of Neurology, Mayo Clinic, College of Medicine, Rochester, Minnesota [454]

Daniel H. Lowenstein, MD

Dr. Robert B. and Mrs. Ellinor Aird Professor of Neurology; Director, Epilepsy Center, University of California, San Francisco, San Francisco, California [437, 438e, 445]

Elyse E. Lower, MD

Medical Oncology and Hematology, University of Cincinnati, Oncology Hematology Care, Inc., Cincinnati, Ohio [390]

Franklin D. Lowy, MD

Professor of Medicine and Pathology, Columbia University College of Physicians and Surgeons, New York, New York [172]

Sheila A. Lukehart, PhD

Professor, Departments of Medicine and Global Health, University of Washington, Seattle, Washington [206, 207e]

Lucio Luzzatto, MD, FRCP, FRCPATH

Professor of Hematology, University of Genova, Genova; Scientific Director, Istituto Toscano Tumori, Florence, Italy [129]

Lawrence C. Madoff, MD

Professor of Medicine, University of Massachusetts Medical School, Worcester, Massachusetts; Director, Division of Epidemiology and Immunization, Massachusetts Department of Public Health, Jamaica Plain, Massachusetts [157, 166e, 167e]

Adel A. F. Mahmoud, MD, PhD

Professor in Molecular Biology and Public Policy, Princeton University, Princeton, New Jersey [259]

Ronald V. Maier, MD

Jane and Donald D. Trunkey Professor and Vice-Chair, Surgery, University of Washington; Surgeon-in-Chief, Harborview Medical Center, Seattle, Washington [324]

Mark E. Mailliard, MD

Frederick F. Paustian Professor; Chief, Division of Gastroenterology and Hepatology, Department of Internal Medicine, University of Nebraska College of Medicine, Omaha, Nebraska [363]

Mahmoud Malas, MD, MHS, FACS

Associate Professor of Surgery, Johns Hopkins University; Director of Endovascular Surgery; Director of The Vascular and Endovascular Clinical Research Center, Johns Hopkins Bayview Medical Center, Baltimore, Maryland [354]

Hari R. Mallidi, MD

Associate Professor of Surgery and Chief, Division of Transplant and Assist Devices; Lester and Sue Smith Endowed Chair in Surgery, Baylor College of Medicine, Houston, Texas [281]

Susan J. Mandel, MD, MPH

Professor of Medicine; Associate Chief, Division of Endocrinology, Diabetes and Metabolism, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [405]

Brian F. Mandell, MD, PhD

Professor and Chairman of Medicine, Cleveland Clinic Lerner College of Medicine, Department of Rheumatic and Immunologic Disease, Cleveland Clinic, Cleveland, Ohio [397]

Lionel A. Mandell, MD, FRCPC

Professor of Medicine, McMaster University, Hamilton, Ontario, Canada [153]

Douglas L. Mann, MD

Lewin Chair and Chief, Cardiovascular Division; Professor of Medicine, Cell Biology and Physiology, Washington University School of Medicine, Cardiologist-in-Chief, Barnes Jewish Hospital, St. Louis, Missouri [279]

JoAnn E. Manson, MD, DrPH

Professor of Medicine and the Elizabeth Fay Brigham Professor of Women's Health, Harvard Medical School; Chief, Division of Preventive Medicine, Brigham and Women's Hospital, Boston, Massachusetts [413]

Eleftheria Maratos-Flier, MD

Professor of Medicine, Harvard Medical School; Division of Endocrinology, Beth Israel Deaconess Medical Center, Boston, Massachusetts [415e]

Guido Marcucci, MD

Professor of Medicine; John B. and Jane T. McCoy Chair in Cancer Research; Associate Director of Translational Research, Comprehensive Cancer Center, The Ohio State University College of Medicine, Columbus, Ohio [132]

Daniel B. Mark, MD, MPH

Professor of Medicine, Duke University Medical Center; Director, Outcomes Research, Duke Clinical Research Institute, Durham, North Carolina [3]

Alexander G. Marneros, MD, PhD

Assistant Professor, Department of Dermatology, Harvard Medical School; Cutaneous Biology Research Center, Massachusetts General Hospital, Boston, Massachusetts [75]

Jeanne M. Marrazzo, MD, MPH

Professor of Medicine, Division of Allergy and Infectious Diseases, University of Washington, Seattle, Washington [163]

Thomas Marrie, MD

Dean, Faculty of Medicine; Professor, Department of Medicine, Dalhousie University, Halifax, Nova Scotia, Canada [211]

Gary J. Martin, MD

Raymond J. Langenbach, MD Professor of Medicine; Vice Chairman for Faculty Affairs, Department of Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois [4]

Joseph B. Martin, MD, PhD

Edward R. and Anne G. Lefler Professor, Department of Neurobiology, Harvard Medical School, Boston, Massachusetts [437]

Susan Maslanka, PhD

Enteric Diseases Laboratory Branch, Centers for Disease Control and Prevention, Atlanta, Georgia [178]

Henry Masur, MD

Chief, Critical Care Medicine Department, Clinical Center, National Institutes of Health, Bethesda, Maryland [244]

Jeremy Matloff, MD

Fellow, Department of Gastroenterology, Icahn School of Medicine at Mount Sinai, New York, New York [485e, 486e]

Robert J. Mayer, MD

Faculty Vice President for Academic Affairs, Dana-Farber Cancer Institute; Stephen B. Kay Family Professor of Medicine, Harvard Medical School, Boston, Massachusetts [109, 110]

Alexander J. McAdam, MD, PhD

Associate Professor of Pathology, Harvard Medical School; Medical Director, Infectious Diseases Diagnostic Laboratory, Children's Hospital of Boston, Boston, Massachusetts [150e]

Calvin O. McCall, MD

Associate Professor, Department of Dermatology, Virginia Commonwealth University Medical Center, Richmond, Virginia; Chief, Dermatology Section, Hunter Holmes McGuire Veterans Affairs Medical Center, Richmond, Virginia [71, 76e]

John F. McConville, MD

Associate Professor of Medicine and Director, Internal Medicine Residency Program, University of Chicago, Chicago, Illinois [318]

Corey A. McGraw, MD

Assistant Professor, The Saul R. Korey Department of Neurology, Albert Einstein College of Medicine, Yeshiva University, New York, New York [486e]

Kevin T. McVary, MD, FACS

Professor and Chairman, Division of Urology, Southern Illinois University School of Medicine, Springfield, Illinois [67]

Mandeep R. Mehra, MD, FACC, FACP

Professor of Medicine, Harvard Medical School; Executive Director, Center for Advanced Heart Disease, Brigham and Women's Hospital; Co-Director, Brigham and Women's Hospital Heart and Vascular Center, Boston, Massachusetts [280]

Nancy K. Mello,[†] PhD

Professor of Psychology (Neuroscience), Harvard Medical School, Boston, Massachusetts; Director, Alcohol and Drug Abuse Research Center, McLean Hospital, Belmont, Massachusetts [469e]

Shlomo Melmed, MD

Senior Vice President and Dean of the Medical Faculty, Cedars-Sinai Medical Center, Los Angeles, California [401e-403]

Jack H. Mendelson,[†] MD

Professor of Psychiatry (Neuroscience), Harvard Medical School, Belmont, Massachusetts [469e]

Robert O. Messing, MD

Professor, Division of Pharmacology and Toxicology, College of Pharmacy; Associate Director, Waggoner Center for Alcohol and Addiction Research, University of Texas at Austin, Austin, Texas [465e]

M.-Marsel Mesulam, MD

Professor of Neurology, Psychiatry and Psychology, Cognitive Neurology and Alzheimer's Disease Center, Northwestern University Feinberg School of Medicine, Chicago, Illinois [36]

Gregory F. Michaud, MD

Assistant Professor of Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [276]

Susan Miesfeldt, MD

Medical Oncology, Medical Director, Cancer Risk and Prevention Clinic, Maine Medical Center, Scarborough, Maine [84]

Edgar L. Milford, MD

Associate Professor of Medicine, Harvard Medical School; Director, Tissue Typing Laboratory, Brigham and Women's Hospital, Boston, Massachusetts [337]

Bruce L. Miller, MD

A. W. and Mary Margaret Clausen Distinguished Professor of Neurology, University of California, San Francisco School of Medicine, San Francisco, California [34, 35, 37e, 448, 453e]

Samuel I. Miller, MD

Professor, Departments of Microbiology, Medicine and Genome Sciences, University of Washington, Seattle, Washington [190]

Simon J. Mitchell, MB ChB, PhD, FUHM, FANZCA

Associate Professor, Department of Anaesthesiology, University of Auckland and Auckland City Hospital, Auckland, New Zealand [477e]

Babak Mokhlesi, MD, MSc

Professor of Medicine, Department of Medicine, Section of Pulmonary and Critical Care; Director, Sleep Disorders Center and Sleep Fellowship Program, University of Chicago, Chicago, Illinois [318]

Thomas A. Moore, MD, FACP, FIDSA

Chairman, Department of Infectious Diseases, Ochsner Health System, New Orleans, Louisiana [246e]

Pat J. Morin, PhD

Senior Director, Scientific Review and Grants Administration, American Association for Cancer Research, Philadelphia, Pennsylvania [101e]

Alison Morris, MD, MS

Associate Professor, Departments of Medicine and Immunology; Director, University of Pittsburgh HIV Lung Research Center, Division of Pulmonary, Allergy, and Critical Care Medicine, University of Pittsburgh School of Medicine, Pittsburgh, Pennsylvania [244]

Charles A. Morris, MD, MPH

Instructor in Medicine, Harvard Medical School; Staff Physician, Brigham and Women's Hospital, Boston, Massachusetts [482e, 484e]

David A. Morrow, MD, MPH

Associate Professor of Medicine, Harvard Medical School; Director, Levine Cardiac Intensive Care Unit; Senior Investigator, TIMI Study Group, Brigham and Women's Hospital, Boston, Massachusetts [19]

William J. Moss, MD, MPH

Professor, Departments of Epidemiology, International Health, and Molecular Microbiology and Immunology, Johns Hopkins Bloomberg School of Public Health, Baltimore, Maryland [229]

Robert J. Motzer, MD

Professor of Medicine, Joan and Sanford Weill College of Medicine of Cornell University D. Attending Physician, Genitourinary Oncology Service, Memorial Sloan-Kettering Cancer Center, New York, New York [114, 116]

David B. Mount, MD

Assistant Professor of Medicine, Harvard Medical School; Renal Division, Brigham and Women's Hospital, Renal Division, Boston VA Healthcare System, Boston, Massachusetts [63, 64e]

Haralampos M. Moutsopoulos, MD, FACP, FRCP(hc), Master ACR

Professor and Director, Department of Pathophysiology, Medical School, National University of Athens, Athens, Greece [379, 383, 387]

Robert S. Munford, MD

Senior Clinician, Laboratory of Clinical Infectious Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [325]

Nikhil C. Munshi, MD

Professor of Medicine, Harvard Medical School; Boston VA Healthcare System; Director of Basic and Correlative Sciences; Associate Director, Jerome Lipper Myeloma Center, Dana-Farber Cancer Institute, Boston, Massachusetts [136]

John R. Murphy, PhD

Professor of Medicine and Microbiology; Director ad interim, National Emerging Infectious Diseases Laboratories Institute, Boston University School of Medicine, Boston, Massachusetts [175]

Timothy F. Murphy, MD

SUNY Distinguished Professor; Director, Clinical and Translational Research Center, University at Buffalo, the State University of New York, Buffalo, New York [182]

Barbara E. Murray, MD

J. Ralph Meadows Professor and Director, Division of Infectious Diseases, University of Texas Medical School, Houston, Texas [174]

Joseph A. Murray, MD

Professor of Medicine, Departments of Internal Medicine and Immunology, Mayo Clinic, Rochester, Minnesota [55]

Mark B. Mycyk, MD

Associate Professor, Department of Emergency Medicine, Northwestern University Feinberg School of Medicine; Associate Professor, Department of Emergency Medicine, Rush University School of Medicine; Research Director, Toxikon Consortium; Attending Physician, Department of Emergency Medicine, Cook County Hospital, Chicago, Illinois [473e]

Robert J. Myerburg, MD

Professor, Departments of Medicine and Physiology, Division of Cardiology; AHA Chair in Cardiovascular Research, University of Miami Miller School of Medicine, Miami, Florida [327]

Avindra Nath, MD

Chief, Section of Infections of the Nervous System; Clinical Director, National Institute of Neurological Disorders and Stroke (NINDS), National Institutes of Health, Bethesda, Maryland [165]

Edward T. Naureckas, MD

Professor of Medicine, and Director, Pulmonary Function Laboratory, and Adult Cystic Fibrosis Laboratory, Section of Pulmonary and Critical Care Medicine, University of Chicago, Chicago, Illinois [306e]

Eric G. Neilson, MD

Lewis Landsberg Dean, and Vice President, Medical Affairs, Feinberg School of Medicine, Northwestern University, Chicago, Illinois [62e, 332e, 338]

Emily Page Nelson, MD

Clinical Instructor, Department of Anesthesiology, Perioperative, and Pain Medicine, Brigham and Women's Hospital, Boston, Massachusetts [484e]

Gerald T. Nepom, MD, PhD

Professor (Affiliate), University of Washington School of Medicine; Director, Benaroya Research Institute at Virginia Mason; Director, Immune Tolerance Network, Seattle, Washington [373e]

[†]Deceased

Eric J. Nestler, MD, PhD

Nash Family Professor and Chair, Department of Neuroscience; Director, Friedman Brain Institute, Ichan School of Medicine at Mount Sinai, New York, New York [465e]

Hartmut P. H. Neumann, MD

Universitaet Freiburg, Medizinische Universitaetsklinik, Freiburg im Breisgau, Germany [407]

Joseph P. Newhouse, PhD

John D. MacArthur Professor of Health Policy and Management, Department of Health Care Policy, Harvard Medical School; Faculty, John F. Kennedy School of Government, Harvard School of Public Health, Faculty of Arts and Sciences, Harvard University, Boston, Massachusetts [15e]

Jonathan Newmark, MD

Colonel, Medical Corps, U.S. Army; Deputy Joint Program Executive Officer, Medical Systems, Joint Program Executive Office for Chemical/Biological Defense, U.S. Department of Defense, Falls Church, Virginia; Adjunct Professor of Neurology, F. Edward Hebert School of Medicine, Uniformed Services University of the Health Sciences, Bethesda, Maryland [262e]

Rathel L. Nolan, III, MD

Professor, Department of Medicine, Division of Infectious Diseases, University of Mississippi Medical Center, Jackson, Mississippi [238]

Robert L. Norris, MD

Professor, Department of Surgery; Chief, Division of Emergency Medicine, Stanford University School of Medicine, Stanford, California [474]

Scott A. Norton, MD, MPH, MSc

Chief of Dermatology, Children's National Health Systems, Washington, DC [475]

Thomas B. Nutman, MD

Head, Helminth Immunology Section, Head, Clinical Parasitology Unit, Laboratory of Parasitic Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [257, 258]

Jose A. Obeso, MD

Professor of Neurology and Director, CIINAC, Hospital de Madrid; Medical School, CEU-San Pablo, Madrid, Spain [449]

Katherine L. O'Brien, MD, MPH

Professor, Department of International Health, Bloomberg School of Public Health, Johns Hopkins University, Baltimore, Maryland [171]

Max R. O'Donnell, MD, MPH

Assistant Professor of Medicine and Epidemiology, Division of Pulmonary, Allergy, and Critical Care Medicine, Columbia University Medical Center, New York, New York [205e]

Nigel O'Farrell, MD, MSc, FRCP

Ealing Hospital, London, United Kingdom [198e]

Jennifer Ogar, MS CCC-SLP

Speech-Language Pathologist, Memory and Aging Center, University of California, San Francisco, San Francisco, California [37e]

Patrick T. O'Gara, MD

Professor of Medicine, Harvard Medical School; Director, Clinical Cardiology, Brigham and Women's Hospital, Boston, Massachusetts [51e, 267, 283-286]

C. Warren Olanow, MD, FRCP, FRCP(hon)

Henry P. and Georgette Goldschmidt Professor and Chairman Emeritus, Department of Neurology; Professor, Department of Neuroscience, Mount Sinai School of Medicine, New York, New York [449]

Andrew B. Onderdonk, PhD

Professor of Pathology, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [150e]

Chung Owyang, MD

H. Marvin Pollard Professor of Internal Medicine; Chief, Division of Gastroenterology, University of Michigan Health System, Ann Arbor, Michigan [344, 352]

Umesh D. Parashar, MBBS, MPH

Lead, Viral Gastroenteritis Epidemiology Team, Division of Viral Diseases, National Center for Immunization and Respiratory Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia [227]

Shreyaskumar R. Patel, MD

Robert R. Herring Distinguished Professor of Medicine; Center Medical Director, Sarcoma Center, The University of Texas M.D. Anderson Cancer Center, Houston, Texas [119e]

David L. Paterson, MD, PhD

Professor of Medicine, University of Queensland Centre for Clinical Research; Royal Brisbane and Women's Hospital, Brisbane, Australia [187]

Gustav Paumgartner, MD

Professor Emeritus of Medicine, University of Munich, Munich, Germany [369]

M. Luisa Pedro-Botet, MD, PhD

Professor of Medicine, Autonomous University of Barcelona; Infectious Diseases Section (Senior Consultant), Germans Trias i Pujol University Hospital, Badalona, Barcelona, Spain [184]

David A. Pegues, MD

Professor of Medicine, Division of Infectious Diseases, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania [190]

Anton Y. Peleg, MBBS, PhD, MPH, FRACP

Associate Professor, Department of Infectious Diseases and Microbiology, The Alfred Hospital and Monash University, Melbourne, Victoria, Australia [187]

Florencia Pereyra, MD

Instructor in Medicine, Harvard Medical School; Associate Physician, Infectious Disease Division, Brigham and Women's Hospital, Boston, Massachusetts [166e, 167e]

Michael A. Pesce, PhD

Professor Emeritus of Pathology and Cell Biology, Columbia University College of Physicians and Surgeons; Director, Biochemical Genetics Laboratory, Columbia University Medical Center, New York Presbyterian Hospital, New York, New York [Appendix]

Clarence J. Peters, MD

John Sealy Distinguished University Chair in Tropical and Emerging Virology; Professor, Department of Microbiology and Immunology; Department of Pathology; Director for Biodefense, Center for Biodefense and Emerging Infectious Diseases, University of Texas Medical Branch, Galveston, Texas [233]

Gerald B. Pier, PhD

Professor of Medicine (Microbiology and Immunobiology), Harvard Medical School; Brigham and Women's Hospital, Division of Infectious Diseases, Boston, Massachusetts [145e]

Richard J. Pollack, PhD

Instructor, Department of Immunology and Infectious Disease, Harvard School of Public Health, Boston, Massachusetts; Senior Environmental Public Health Officer, Department of Environmental Health and Safety, Harvard University, Cambridge, Massachusetts; President and Chief Scientific Officer, IdentifyUS LLC, Newton, Massachusetts [475]

Martin R. Pollak, MD

Professor of Medicine, Harvard Medical School; Beth Israel Deaconess Medical Center, Boston, Massachusetts [339]

Andrew J. Pollard, PhD, FRCPCH

Professor of Paediatric Infection and Immunity, Department of Paediatrics, University of Oxford, Oxford, United Kingdom [180]

Reuven Porat, MD

Professor of Medicine, Department of Internal Medicine, Tel Aviv Sourasky Medical Center; Sackler Faculty of Medicine, Tel Aviv University, Tel Aviv, Israel [23]

Daniel A. Portnoy, PhD

Professor, Department of Molecular and Cell Biology and the School of Public Health, University of California, Berkeley, Berkeley, California [176]

John T. Potts, Jr., MD

Jackson Distinguished Professor of Clinical Medicine, Harvard Medical School; Physician-in-Chief and Director of Research Emeritus, Massachusetts General Hospital, Boston, Massachusetts [424]

Lawrie W. Powell, MD, PhD

Professor of Medicine, The University of Queensland; Director, Centre for the Advancement of Clinical Research, Royal Brisbane and Women's Hospital, Brisbane, Australia [428]

Alvin C. Powers, MD

Joe C. Davis Chair in Biomedical Science; Professor of Medicine, Molecular Physiology and Biophysics; Director, Vanderbilt Diabetes Center; Chief, Division of Diabetes, Endocrinology, and Metabolism, Vanderbilt University School of Medicine, Nashville, Tennessee [417-419]

Daniel S. Pratt, MD

Assistant Professor of Medicine, Harvard Medical Center; Massachusetts General Hospital, Boston, Massachusetts [58, 358]

Michael B. Prentice, MB ChB, PhD, MRCP(UK), FRCPath, FFPRCP

Professor of Medical Microbiology, Departments of Microbiology and Pathology, University College Cork, Cork, Ireland [196]

Darwin J. Prockop, MD, PhD

Director and Professor, Institute for Regenerative Medicine, Texas A&M Health Science Center College of Medicine at Scott & White, Temple, Texas [427]

Stanley B. Prusiner, MD

Director, Institute for Neurodegenerative Diseases; Professor, Department of Neurology, University of California, San Francisco, San Francisco, California [444e, 453e]

Thomas C. Quinn, MD

Professor of Medicine, Johns Hopkins University, Baltimore, Maryland; Senior Investigator, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland [213]

Gil Rabinovici, MD

Associate Professor in Neurology, Memory and Aging Center, University of California, San Francisco, San Francisco, California [37e]

Daniel J. Rader, MD

Seymour Gray Professor of Molecular Medicine; Chair, Department of Genetics; Chief, Division of Translational Medicine and Human Genetics, Department of Medicine, Perelman School of Medicine at the University of Pennsylvania, Philadelphia, Pennsylvania [421]

Kaitlin Rainwater-Lovett, PhD, MPH

Research Fellow, Division of Infectious Diseases, Department of Pediatrics, Johns Hopkins University School of Medicine, Baltimore, Maryland [229]

Sanjay Ram, MBBS

Associate Professor of Medicine, Division of Infectious Diseases and Immunology, University of Massachusetts Medical School, Worcester, Massachusetts [181]

Reuben Ramphal, MD

Adjunct Professor of Medicine, Division of Infectious Diseases and Global Medicine, University of Florida College of Medicine, Gainesville, Florida [189]

Agam K. Rao, MD

Medical Officer, Division of Foodborne, Waterborne, and Environmental Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia [178]

Beth Rapaport, MD

Attending Physician, Elmhurst Hospital Center, Mount Sinai Medical Affiliate, Elmhurst, New York [486e]

Kumanan Rasanathan, MBChB, MPH, FAFPHM

Technical Officer, Department of Ethics, Equity, Trade, and Human Rights, World Health Organization, Geneva, Switzerland [13e]

Neil H. Raskin, MD

Department of Neurology, University of California, San Francisco, San Francisco, California [21, 447]

Anis Rassi, Jr., MD, PhD, FACC, FACP, FAHA

Scientific Director, Anis Rassi Hospital, Goiânia, Brazil [252]

James P. Rathmell, MD

Henry Knowles Beecher Professor of Anesthesiology, Harvard Medical School; Executive Vice Chair and Chief, Division of Pain Medicine, Department of Anesthesia, Critical Care and Pain Medicine, Massachusetts General Hospital, Boston, Massachusetts [18]

Mario C. Raviglione, MD

Director, Global TB Programme, World Health Organization, Geneva, Switzerland [202]

Divya Reddy, MBBS, MPH

Faculty, Department of Medicine, Pulmonary Division, Albert Einstein College of Medicine; Montefiore Medical Center, Bronx, New York [205e]

Susan Redline, MD, MPH

Peter C. Farrell Professor of Sleep Medicine, Harvard Medical School; Brigham and Women's Hospital; Beth Israel Deaconess Medical Center, Boston, Massachusetts [319]

Sharon L. Reed, MD, MSCTM, D(ABMM)

Professor of Pathology and Medicine; Director, Microbiology Laboratory, University of California, San Diego School of Medicine, La Jolla, California [245e, 247]

Susan E. Reef, MD

Medical Epidemiologist, Centers for Disease Control and Prevention, Atlanta, Georgia [230e]

John J. Reilly, Jr., MD

Jack D. Myers Professor of Medicine and Chair, Department of Medicine; Vice Chair for Clinical Affairs, University of Pittsburgh, Pittsburgh, Pennsylvania [308e, 314]

John T. Repke, MD, FACOG

University Professor and Chairman, Department of Obstetrics and Gynecology, Pennsylvania State University College of Medicine; Obstetrician-Gynecologist In-Chief, The Milton S. Hershey Medical Center, Hershey, Pennsylvania [8]

Victor I. Reus, MD

Department of Psychiatry, University of California, San Francisco School of Medicine; Langley Porter Neuropsychiatric Institute, San Francisco, California [466]

Joseph J. Rhatigan, MD

Assistant Professor, Harvard Medical School, Harvard School of Public Health; Associate Chief, Division of Global Health Equity, Brigham and Women's Hospital, Boston, Massachusetts [2]

Peter A. Rice, MD

Professor of Medicine, Division of Infectious Diseases and Immunology, University of Massachusetts Medical School, Worcester, Massachusetts [181]

Elizabeth Robbins, MD

Clinical Professor of Pediatrics, University of California, San Francisco, San Francisco, California [443e]

Gary L. Robertson, MD

Emeritus Professor of Medicine, Northwestern University School of Medicine, Chicago, Illinois [404]

Russell G. Robertson, MD

Professor of Family Medicine, Chicago Medical School, Rosalind Franklin University of Medicine and Science, Chicago, Illinois [56]

Dan M. Roden, MD

William Stokes Professor of Experimental Therapeutics; Professor of Medicine and Pharmacology, Assistant Vice-Chancellor for Personalized Medicine, Director, Oates Institute for Experimental Therapeutics, Vanderbilt University School of Medicine, Nashville, Tennessee [5]

James A. Romano, Jr., PhD, DABT

Principal Senior Life Scientist Advisor, Tunnell Government Services, Inc., Rockville, Maryland [262e]

Karen L. Roos, MD

John and Nancy Nelson Professor of Neurology; Professor of Neurological Surgery, Indiana University School of Medicine, Indianapolis, Indiana [164]

Allan H. Ropper, MD, FRCP, FACP

Professor of Neurology, Harvard Medical School; Raymond D. Adams Master Clinician; Executive Vice Chair, Department of Neurology, Brigham and Women's Hospital, Boston, Massachusetts [328, 456, 457e]

Jonathan E. Rosenberg, MD

Associate Attending; Section Chief, Non-Prostate Program, Division of Solid Tumor Oncology, Department of Medicine, Memorial Sloan-Kettering Cancer Center, New York, New York [114]

Roger N. Rosenberg, MD

Zale Distinguished Chair and Professor of Neurology, Department of Neurology, University of Texas Southwestern Medical Center, Dallas, Texas [450, 451e]

Myrna R. Rosenfeld, MD, PhD

Department of Neurology, Hospital Clínic/IDIBAPS, Barcelona, Spain [122]

Michael A. Rubin, MD, PhD

Assistant Professor of Medicine, University of Utah School of Medicine, Salt Lake City, Utah [44]

Steven A. Rubin, PhD

Acting Principal Investigator, Center for Biologics Evaluation and Research, Food and Drug Administration, Bethesda, Maryland [231e]

Robert M. Russell, MD

Professor Emeritus of Medicine and Nutrition, Tufts University, Boston, Massachusetts; Office of Dietary Supplements, National Institutes of Health, Bethesda, Maryland [96e]

Thomas A. Russo, MD, CM

Staff Physician, Western New York VA Healthcare System; Professor of Medicine and Microbiology and Immunology; Vice Chair of Medicine; Head, Division of Infectious Disease, University at Buffalo, State University of New York, Buffalo, New York [186, 200]

Anna E. Rutherford, MD, MPH

Assistant Professor, Harvard Medical School; Associate Physician, Brigham and Women's Hospital, Boston, Massachusetts [483e]

Edward T. Ryan, MD, FACP, FIDSA, FASTMH

Professor of Medicine, Harvard Medical School; Professor of Immunology and Infectious Diseases, Harvard School of Public Health; Director, Global Infectious Diseases, Division of Infectious Diseases, Massachusetts General Hospital, Boston, Massachusetts [160, 193]

Sean Sadikot, MD

Department of Internal Medicine, Critical Care and Pulmonary Disease, Hackensack University Medical Center, Hackensack, New Jersey [485e, 486e]

David J. Salant, MD

Professor of Medicine, Boston University School of Medicine; Chief, Section of Nephrology, Boston Medical Center, Boston, Massachusetts [340]

Martin A. Samuels, MD

Professor of Neurology, Harvard Medical School; Chair, Department of Neurology, Brigham and Women's Hospital, Boston, Massachusetts [439e, 463e]

Philippe J. Sansonetti, MD

Professor, Collège de France; Institut Pasteur, Paris, France [191]

Clifford B. Saper, MD, PhD

James Jackson Putnam Professor of Neurology and Neuroscience, Harvard Medical School; Chairman, Department of Neurology, Beth Israel Deaconess Medical Center, Boston, Massachusetts [38]

Jussi J. Saukkonen, MD

Associate Professor of Medicine, Section of Pulmonary, Allergy, and Critical Care Medicine, Boston University School of Medicine, Boston, Massachusetts [205e]

Edward A. Sausville, MD, PhD

Professor of Medicine, University of Maryland School of Medicine; Associate Director for Clinical Research, Marlene and Stewart Greenbaum Cancer Center, Baltimore, Maryland [103e]

Mohamed H. Sayegh, MD

Raja N. Khuri Dean, Faculty of Medicine; Professor of Medicine and Immunology; Vice President of Medical Affairs, American University of Beirut, Beirut, Lebanon; Senior Lecturer, Harvard Medical School; Schuster Family Transplantation Center, Brigham and Women's Hospital, Boston, Massachusetts [337]

David T. Scadden, MD

Gerald and Darlene Professor of Medicine; Co-Chair, Harvard Stem Cell Institute; Co-chair, Department of Stem Cell and Regenerative Biology, Harvard Medical School; Director, Center for Regenerative Medicine; Chief, Hematologic Malignancies, Cancer Center, Massachusetts General Hospital, Boston, Massachusetts [89e]

Thomas E. Scammell, MD

Professor, Harvard Medical School; Beth Israel Deaconess Medical Center; Boston Children's Hospital, Boston, Massachusetts [38]

Anthony H. V. Schapira, MD, DSc, FRCP, FMedSci

Chair and Professor of Clinical Neurosciences, UCL Institute of Neurology, London, United Kingdom [449]

Howard I. Scher, MD

Professor of Medicine, Joan and Sanford Weill College of Medicine of Cornell University; D. Wayne Calloway Chair in Urologic Oncology; Attending Physician and Chief, Genitourinary Oncology Service, Department of Medicine, Memorial Sloan-Kettering Cancer Center, New York, New York [114, 115]

Anne Schuchat, MD

RADM, U.S. Public Health Service; Assistant Surgeon General, National Center for Immunization and Respiratory Diseases, Centers for Disease Control and Prevention, Atlanta, Georgia [148]

Marc A. Schuckit, MD

Distinguished Professor of Psychiatry, University of California, San Diego School of Medicine, La Jolla, California [467]

H. Ralph Schumacher, Jr., MD

Professor of Medicine, Division of Rheumatology, University of Pennsylvania, School of Medicine, Philadelphia, Pennsylvania [395]

Gordon E. Schutz, MD, FAAP

Professor of Pediatrics; Vice-Chairman for Educational Affairs; Martin I. Lorin, MD Chair in Medical Education, Department of Pediatrics, Section of Retrovirology, Vice President, Baylor International Pediatric AIDS Initiative at Texas Children's Hospital, Baylor College of Medicine, Texas Children's Hospital, Houston, Texas [195]

Richard M. Schwartzstein, MD

Ellen and Melvin Gordon Professor of Medicine and Medical Education, Harvard Medical School; Associate Division Chief, Division of Pulmonary, Critical Care, and Sleep Medicine, Beth Israel Deaconess Medical Center, Boston, Massachusetts [47e]

William W. Seeley, MD

Associate Professor of Neurology and Pathology, Memory and Aging Center, University of California, San Francisco, San Francisco, California [35, 448]

Michael V. Seiden, MD, PhD

Chief Medical Officer, McKesson Specialty Health, The Woodlands, Texas [117]

Julian L. Seifter, MD

Associate Professor of Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [343]

David C. Seldin, MD, PhD

Professor, Departments of Medicine and Microbiology; Chief, Section of Hematology-Oncology; Director, Amyloidosis Center, Boston University School of Medicine; Boston Medical Center, Boston, Massachusetts [137]

Ankoor Shah, MD

Assistant Professor, Department of Medicine, Division of Rheumatology and Immunology, Duke University Medical Center, Durham, North Carolina [380]

Steven D. Shapiro, MD

Jack D. Myers Professor and Chair, Department of Medicine, University of Pittsburgh, Pittsburgh, Pennsylvania [314]

Erica S. Shenoy, MD, PhD

Instructor in Medicine, Harvard Medical School; Assistant Chief, Infection Control Unit, Massachusetts General Hospital, Boston, Massachusetts [170]

Kanade Shinkai, MD, PhD

Assistant Professor, Department of Dermatology, University of California, San Francisco, San Francisco, California [74]

William Silen, MD

Johnson and Johnson Professor Emeritus of Surgery, Harvard Medical School, Aurburndale, Massachusetts [20]

Edwin K. Silverman, MD, PhD

Professor of Medicine, Harvard Medical School; Chief, Channing Division of Network Medicine, Department of Medicine, Brigham and Women's Hospital, Boston, Massachusetts [314]

Karl Skorecki, MD, FRCP(C), FASN

Annie Chutick Professor in Medicine (Nephrology); Director, Rappaport Research Institute, Technion-Israel Institute of Technology; Director of Medical and Research Development, Rambam Health Care Campus, Haifa, Israel [85e, 335]

Wade S. Smith, MD, PhD

Professor of Neurology, Daryl R. Gress Endowed Chair of Neurocritical Care and Stroke; Director, University of California, San Francisco Neurovascular Service, San Francisco, California [330, 446]

Elizabeth Smyth, MB BAO, MSc

Department of Gastrointestinal Oncology, Royal Marsden NHS Foundation Trust, London and Sutton, United Kingdom [112]

Kelly A. Soderberg, PhD, MPH

Associate Director, Duke Center for HIV/AIDS Vaccine Immunology-Immunogen Discovery, Duke Human Vaccine Institute, Duke University, Durham, North Carolina [372e]

Scott D. Solomon, MD

Professor, Harvard Medical School; Director, Noninvasive Cardiology, Brigham and Women's Hospital, Boston, Massachusetts [270e, 271e]

Julian Solway, MD

Walter L. Palmer Distinguished Service Professor of Medicine and Pediatrics; Associate Dean for Translational Medicine, Biological Sciences Division; Vice Chair for Research, Department of Medicine; Chair, Committee on Molecular Medicine, University of Chicago, Chicago, Illinois [306e, 318]

Michael F. Sorrell, MD

Robert L. Grissom Professor of Medicine, Department of Internal Medicine, University of Nebraska Medical Center, Omaha, Nebraska [363]

Eric J. Sorscher, MD

Professor, Departments of Medicine and Genetics, Gwaltney Chair for Medical Research; Director, Gregory Fleming James Cystic Fibrosis Research Center, University of Alabama at Birmingham, Birmingham, Alabama [313]

Frank E. Speizer, MD

E. H. Kass Distinguished Professor of Medicine, Harvard Medical School; Channing Division of Network Medicine, Department of Medicine, Brigham and Women's Hospital; Professor of Environmental Science, Department of Environmental Health, Harvard School of Public Health, Boston, Massachusetts [311]

Brad Spellberg, MD

Professor of Medicine; Associate Medical Director for Inpatient Services, Harbor-UCLA Medical Center and Los Angeles Biomedical Research Institute, Torrance, California [242]

Nancy B. Spinner, PhD, FACMG

Professor of Pathology and Laboratory Medicine, Perelman School of Medicine, University of Pennsylvania; Chief, Division of Genomic Diagnostics, Children's Hospital of Philadelphia, Philadelphia, Pennsylvania [83e]

Jerry L. Spivak, MD

Professor of Medicine and Oncology, Hematology Division, Johns Hopkins University School of Medicine, Baltimore, Maryland [131]

David D. Spragg, MD

Associate Professor, Department of Medicine, Johns Hopkins University; Director, Electrophysiology Laboratory, Johns Hopkins Bayview Medical Center, Baltimore, Maryland [273e-275]

E. William St. Clair, MD

Professor of Medicine and Immunology, Department of Medicine, Duke University Medical Center, Durham, North Carolina [380]

Allen C. Steere, MD

Professor of Medicine, Harvard Medical School; Massachusetts General Hospital, Boston, Massachusetts [210]

Robert S. Stern, MD

Carl J. Herzog Professor of Dermatology, Harvard Medical School; Chair, Department of Dermatology, Beth Israel Deaconess Medical Center, Boston, Massachusetts [74]

Dennis L. Stevens, MD, PhD

Professor of Medicine, University of Washington School of Medicine, Seattle, Washington [156, 179]

Lynne Warner Stevenson, MD

Professor of Medicine, Harvard Medical School; Director, Heart Failure Program, Brigham and Women's Hospital, Boston, Massachusetts [287]

William G. Stevenson, MD

Brigham and Women's Hospital; Cardiovascular Division, Department of Medicine, Harvard Medical School, Boston, Massachusetts [276, 277]

John H. Stone, MD, MPH

Professor of Medicine, Harvard Medical School; Director, Clinical Rheumatology, Massachusetts General Hospital, Boston, Massachusetts [391e]

Stephanie Studenski, MD, MPH

Director, Longitudinal Studies Section, Intramural Research Program, National Institute on Aging, National Institutes of Health, Baltimore, Maryland [11]

Lewis Sudarsky, MD

Associate Professor of Neurology, Harvard Medical School; Director of Movement Disorders, Brigham and Women's Hospital, Boston, Massachusetts [32]

Donna C. Sullivan, PhD

Professor, Department of Medicine, Division of Infectious Diseases, University of Mississippi Medical School, Jackson, Mississippi [238]

Shyam Sundar, MD, FRCP, FNA

Professor of Medicine, Institute of Medical Sciences, Banaras Hindu University, Varanasi, India [251]

Neeraj K. Surana, MD, PhD

Instructor in Pediatrics, Harvard Medical School; Assistant in Medicine, Boston Children's Hospital, Boston, Massachusetts [144]

Paolo M. Suter, MD, MS

Professor, Clinic and Polyclinic of Internal Medicine, University Hospital, Zurich, Switzerland [96e]

Richard M. Suzman, PhD

Director, Behavioral and Social Research Program, National Institute on Aging, National Institutes of Health, Bethesda, Maryland [93e]

Robert A. Swerlick, MD

Alicia Leizman Stonecipher Professor and Chair of Dermatology, Emory University School of Medicine, Atlanta, Georgia [76e]

Homayoun Tabandeh, MD

Retina-Vitreous Associates Medical Group, Los Angeles, California [40e]

Geoffrey Tabin, MD

John and Marva Warnock Presidential Professor, University of Utah School of Medicine; Director, International Ophthalmology Division, John A. Moran Eye Center; Director, Himalayan Cataract Project, Salt Lake City, Utah [476e]

Maria Carmela Tartaglia, MD

Assistant Professor, Tanz Centre for Research in Neurodegenerative Diseases, University of Toronto, Toronto, Ontario, Canada [37e]

Joel D. Taurog, MD

Professor of Internal Medicine, Rheumatic Diseases Division, University of Texas Southwestern Medical Center at Dallas, Dallas, Texas [384]

Ayalew Tefferi, MD

Professor of Medicine and Hematology, Mayo Clinic, Rochester, Minnesota [135e]

Stephen C. Textor, MD

Professor of Medicine, Division of Nephrology and Hypertension, Mayo Clinic, Rochester, Minnesota [299, 341]

Rajesh V. Thakker, MD, FMedSci, FR

May Professor of Medicine, Academic Endocrine Unit, University of Oxford; O.C.D.E.M., Churchill Hospital, Headington, Oxford, United Kingdom [408]

C. Louise Thwaites, MD, MBBS

Oxford University Clinical Research Unit, Hospital for Tropical Diseases, Ho Chi Minh City, Vietnam [177]

Zelig A. Tochner, MD

Professor of Radiation Oncology, University of Pennsylvania School of Medicine; Medical Director, Proton Therapy Center, Philadelphia, Pennsylvania [263e]

Gordon F. Tomaselli, MD

Michel Mirowski, MD Professor of Cardiology; Professor of Medicine and Cellular and Molecular Medicine; Chief, Division of Cardiology, Johns Hopkins University, Baltimore, Maryland [273e-275]

Karina A. Top, MD, MS

Assistant Professor of Pediatrics, Dalhousie University, Halifax, Nova Scotia, Canada [185]

Mark Topazian, MD

Professor of Medicine, Mayo Clinic, Rochester, Minnesota [345, 346e]

Barbara W. Trautner, MD, PhD

Assistant Professor, Section of Infectious Diseases, Department of Medicine, Baylor College of Medicine; Houston VA Health Services Research and Development Center of Excellence, Houston, Texas [162]

Jeffrey M. Trent, PhD, FACMG

President and Research Director, Translational Genomics Research Institute, Phoenix, Arizona; Van Andel Research Institute, Grand Rapids, Michigan [101e]

Elbert P. Trulock, III, MD

Rosemary and I. Jerome Flance Professor in Pulmonary Medicine, Washington University School of Medicine, St. Louis, Missouri [320e]

Kenneth L. Tyler, MD

Reuler-Lewin Family Professor and Chair of Neurology; Professor of Medicine, Immunology, and Microbiology, University of Colorado School of Medicine, Aurora, Colorado; Neurologist, Denver Veterans Affairs Medical Center, Denver, Colorado [164]

Athanasios G. Tzioufas, MD

Professor, Department of Pathophysiology, School of Medicine, National University of Athens, Athens, Greece [383]

Walter J. Urba, MD, PhD

Director of Research, Earle A. Chiles Research Institute, Providence Cancer Center, Portland, Oregon [105]

Jos W. M. van der Meer, MD, PhD

Professor of Medicine; Head, Department of General Internal Medicine, Radboud University Nijmegen Medical Centre, Nijmegen, The Netherlands [26, 464e]

Edouard G. Vannier, PharmD, PhD

Assistant Professor, Division of Geographic Medicine and Infectious Diseases, Department of Medicine, Tufts Medical Center and Tufts University School of Medicine, Boston, Massachusetts [249]

Gauri R. Varadhachary, MD

Professor, Department of Gastrointestinal Medical Oncology, The University of Texas M.D. Anderson Cancer Center, Houston, Texas [120e]

John Varga, MD

John Hughes Professor of Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois [382]

Christy A. Varughese, PharmD

Infectious Disease Specialist, Department of Pharmacy, Massachusetts General Hospital, Boston, Massachusetts [170]

Panayiotis G. Vlachoyiannopoulos, MD

Professor of Medicine-Immunology, Department of Pathophysiology, Medical School, National University of Athens, Athens, Greece [379]

Bert Vogelstein, MD

Investigator, Howard Hughes Medical Institute; Director, Ludwig Center at the Sidney Kimmel Comprehensive Cancer Center; Clayton Professor of Oncology and Pathology; Johns Hopkins Medical Institutions, Baltimore, Maryland [101e]

Everett E. Vokes, MD

John E. Ulmann Professor; Chairman, Department of Medicine; Physician-in-Chief, University of Chicago Medical Center, Chicago, Illinois [106]

Tamara J. Vokes, MD

Professor, Department of Medicine, Section of Endocrinology, University of Chicago, Chicago, Illinois [426e]

Jiří F. P. Wagenaar, MD, PhD

Senior Scientist, WHO/FAO/OIE and National Leptospirosis Reference Centre, KIT Biomedical Research, KIT (Royal Tropical Institute), Amsterdam, The Netherlands [208]

Sushrut S. Waikar, MD, MPH

Associate Professor, Harvard Medical School; Director, Renal Ambulatory Services, Brigham and Women's Hospital, Boston, Massachusetts [334]

Matthew K. Waldor, MD, PhD

Edward H. Kass Professor of Medicine, Channing Laboratory, Brigham and Women's Hospital; Harvard Medical School and Howard Hughes Medical Institute, Boston, Massachusetts [193]

David H. Walker, MD

The Carmage and Martha Walls Distinguished University Chair in Tropical Diseases; Professor and Chairman, Department of Pathology; Executive Director, Center for Biodefense and Emerging Infectious Diseases, University of Texas Medical Branch, Galveston, Texas [211]

Mark F. Walker, MD

Associate Professor, Neurology, Case Western Reserve University; Cleveland VA Medical Center, Cleveland, Ohio [28]

Fred Wang, MD

Professor of Medicine, Harvard Medical School and Brigham and Women's Hospital, Boston, Massachusetts [214e, 220e]

John W. Warren, MD

Professor of Medicine, University of Maryland School of Medicine, Baltimore, Maryland [60e]

Aaron B. Waxman, MD, PhD, FACP, FCCP

Associate Professor of Medicine, Harvard Medical School; Executive Director, Center for Pulmonary-Heart Diseases, Brigham and Women's Hospital Heart and Vascular Center, Boston, Massachusetts [304]

Michael E. Wechsler, MD, MMSc

Professor of Medicine; Director, Asthma Program, National Jewish Health, Denver, Colorado [310]

Anthony P. Weetman, MD, DSc

University of Sheffield, School of Medicine Sheffield, United Kingdom [405]

Robert A. Weinstein, MD

The C. Anderson Hedberg, MD Professor of Internal Medicine, Rush Medical College; Chief Academic Officer, Cook County Health and Hospitals System, Chicago, Illinois [168]

Jeffrey I. Weitz, MD, FRCP(C), FACP

Professor of Medicine and Biochemistry, McMaster University; Executive Director, Thrombosis and Atherosclerosis Research Institute, Hamilton, Ontario, Canada [143]

Peter F. Weller, MD

Chief, Infectious Disease Division; Chief, Allergy and Inflammation Division; Beth Israel Deaconess Medical Center, Boston, Massachusetts [254-258, 260]

Andrew Wellman, MD, PhD

Assistant Professor of Medicine, Division of Sleep Medicine, Harvard Medical School; Brigham and Women's Hospital, Boston, Massachusetts [319]

Patrick Y. Wen, MD

Professor of Neurology, Harvard Medical School; Director, Center for Neuro-Oncology, Dana-Farber Cancer Institute; Director, Division of Neuro-Oncology, Department of Neurology, Brigham and Women's Hospital; Dana-Farber Cancer Institute, Boston, Massachusetts [118]

Michael R. Wessels, MD

John F. Enders Professor of Pediatrics; Professor of Medicine, Harvard Medical School; Chief, Division of Infectious Diseases, Boston Children's Hospital, Boston, Massachusetts [173]

L. Joseph Wheat, MD

President and Medical Director, MiraVista Diagnostics, LLC, Indianapolis, Indiana [236]

A. Clinton White, Jr., MD

Paul R. Stalnaker Distinguished Professor; Director, Infectious Disease Division, Department of Internal Medicine, University of Texas Medical Branch, Galveston, Texas [260]

Nicholas J. White, DSc, MD, FRCP, F Med Sci, FRS

Professor of Tropical Medicine, Faculty of Tropical Medicine, Mahidol University, Mahidol-Oxford Research Unit, Bangkok, Thailand [248, 250e]

Richard J. Whitley, MD

Distinguished Professor of Pediatrics; Loeb Eminent Scholar Chair in Pediatrics; Professor of Microbiology, Medicine and Neurosurgery, The University of Alabama at Birmingham, Birmingham, Alabama [217]

Bruce U. Wintroub, MD

Professor and Chair, Department of Dermatology, University of California, San Francisco, San Francisco, California [74]

Andrea S. Wolf, MD, MPH

Assistant Professor, and Director, Women's Lung Cancer Program, Mt. Sinai Medical Center, New York, New York [482e]

Allan W. Wolkoff, MD

The Herman Lopata Chair in Liver Disease Research; Professor of Medicine and Anatomy and Structural Biology; Associate Chair of Medicine for Research; Chief, Division of Gastroenterology and Liver Diseases; Director, Marion Bessin Liver Research Center, Albert Einstein College of Medicine and Montefiore Medical Center, Bronx, New York [359]

John B. Wong, MD

Professor of Medicine, Tufts University School of Medicine; Chief, Division of Clinical Decision Making, Department of Medicine, Tufts Medical Center, Boston, Massachusetts [3]

Louis Michel Wong Kee Song, MD

Associate Professor, Division of Gastroenterology and Hepatology, Mayo Clinic College of Medicine, Rochester, Minnesota [345, 346e]

Robert L. Wortmann, MD, FACP, MACR

Professor Emeritus, Department of Medicine, Geisel School of Medicine at Dartmouth, Lebanon, New Hampshire [431e]

Shirley H. Wray, MD, PhD, FRCP

Professor of Neurology, Harvard Medical School; Master Clinician, Department of Neurology, Massachusetts General Hospital, Boston, Massachusetts [41e]

Richard G. Wunderink, MD

Professor, Pulmonary and Critical Care, Northwestern University Feinberg School of Medicine, Chicago, Illinois [153]

Kim B. Yancey, MD

Professor and Chair, Department of Dermatology, University of Texas Southwestern Medical Center in Dallas, Dallas, Texas [70, 73]

Janet A. Yellowitz, DMD, MPH

Associate Professor; Director, Geriatric Dentistry, University of Maryland Dental School, Baltimore, Maryland [46e]

Lam Minh Yen, MD

Director, Tetanus Intensive Care Unit, Hospital for Tropical Diseases, Ho Chi Minh City, Vietnam [177]

Maria A. Yialamas, MD

Assistant Professor of Medicine, Harvard Medical School; Associate Program Director, Internal Medicine Residency, Brigham and Women's Hospital, Boston, Massachusetts [481e, 483e]

Neal S. Young, MD

Chief, Hematology Branch, National Heart, Lung and Blood Institute; Director, NIH Center for Human Immunology, Autoimmunity and Inflammation, National Institutes of Health, Bethesda, Maryland [130]

Victor L. Yu, MD

Professor of Medicine, Department of Medicine, University of Pittsburgh Medical Center, Pittsburgh, Pennsylvania [184]

Jing Zhou, MD, PhD, FASN

Director, Laboratory of Molecular Genetics and Developmental Biology of Disease, Renal Division; Director, Center for Polycystic Kidney Disease Research, Brigham and Women's Hospital; Harvard Medical School, Boston, Massachusetts [339]

Werner Zimmerli, MD

Professor of Medicine, Basel University; Interdisciplinary Unit of Orthopaedic Infection, Kantonspital Baselland, Liestal, Switzerland [158]

Laura A. Zimmerman, MPH

Epidemiologist, Centers for Disease Control and Prevention, Atlanta, Georgia [230e]

The Editors are pleased to present the 19th edition of *Harrison's Principles of Internal Medicine*. Since the first edition was published 65 years ago, virtually every area of medicine and medical education has evolved remarkably, and many new fields have emerged.

While retaining the founding goals of *Harrison's*, this edition has been modified extensively in light of the varied needs of the book's readers and the diverse methods and formats by which information is now acquired and applied. The focus of medical education worldwide is shifting from the classic structure/function/disease approach to an integrated, often case-based approach in which basic and population sciences are specifically linked to the practical diagnosis and management of disease. Many of the updates and changes described here have been undertaken with the modern educational and clinical environments in mind.

This new edition offers a thoroughly updated presentation of the classic pathophysiologic basis of clinical medicine and details the cutting-edge methods and tools that are now available for the assessment of symptoms and the effective management of diseases in the modern patient-care environment. The text is supplemented by germane new photographs, radiographs, illustrations, atlases, patient-care algorithms, tables, and practical demonstrative videos.

In the interest of producing the most practical format possible for the 19th edition, a new system of referencing has been used. Detailed bibliographic listings, with summaries of articles' relevance to practice, appear in the online edition, replacing the general and limited collection of Suggested Readings that appeared in the prior print editions.

The 19th edition of *Harrison's* is designed for accessibility and flexibility. The print textbook is available in two volumes. Volume 1 focuses on the foundations of medicine and the understanding and assessment of cardinal disease manifestations; Volume 2 focuses on specific diseases, by system. This functional division will be helpful to students who are mastering the basis of clinical medicine and to clinicians who are more focused on gaining an advanced understanding of mechanisms and patient care in specific diseases. In terms of digital media, *Harrison's* is now available as an eBook on multiple platforms, as an enhanced "app" developed especially for tablet and smartphone devices that offer high-definition resolution of multimedia content and interactive features, and as an updated online edition. All of these options provide access to our numerous e-chapters as well as to videos and atlases. Additional resources include the *Harrison's Self-Assessment and Board Review*, a useful study guide based on information in the 19th edition, and the *Harrison's Manual of Medicine*, a pocket version of *Harrison's Principles of Internal Medicine*. A new *Harrison's* collection of case vignettes emphasizing differential diagnostic considerations in the assessment of cardinal manifestations will also be available.

Advances in medical science have been spectacular since *Harrison's* was first published in 1949. At that time, peptic ulcer disease was thought to be caused by stress, nearly every tumor that was not resected resulted in death, rheumatic heart disease was widely prevalent, and hepatitis B and HIV infection were unknown. In the intervening years, both the infectious cause of and the cure for peptic ulcer disease were identified; advances in diagnosis and treatment made it possible to cure two-thirds of cancers; rheumatic heart disease virtually disappeared; atherosclerotic coronary artery disease waxed and then—at least in part through management of modifiable risk factors—began to wane; hepatitis B and its consequences, cirrhosis and hepatocellular carcinoma, became preventable by a vaccine; and HIV, first viewed as a uniformly fatal worldwide scourge, became a treatable chronic disease. Notably, emerging and reemerging diseases have presented significant challenges to medical research and practice, while a new understanding of systems-wide concepts such as the microbiome offers exciting new possibilities for understanding and managing health and disease in ways never before possible.

Of particular note in this 19th edition are critical updates in the classic chapter on HIV/AIDS, which offers both a clinically pragmatic

focus and a comprehensive and analytical approach to pathogenesis. The updates cover the latest treatment protocols and address the issue of combination prevention modalities, making the chapter the most up-to-date and comprehensive treatise on HIV disease available.

Several other chapters likewise reflect the rapid pace of advancement in the field of immune-related diseases and their treatment. In this regard, Chapter 372e, "Introduction to the Immune System," serves as a mini-textbook of immunology for use in immunology courses. In addition, a new chapter on IgG4-mediated disease summarizes an important and newly recognized constellation of entities.

Readers will find expanded coverage of the neurodegenerative diseases, highlighting advances in their classification and management and delineating the new understanding of mechanisms responsible for the deposition and spread of pathogenic protein aggregates in these disorders. The chapter on chronic hepatitis discusses in detail the dramatic new discoveries in the use of direct-acting antiviral agents for the treatment of hepatitis C virus disease; these agents are responsible for some of the most exciting therapeutic advances in medicine today.

The rapidly expanding application of genetic knowledge to medicine is covered in many chapters, including a new chapter on microbial genomics and infectious diseases and substantially updated chapters on the human microbiome and chromosomal genetic disorders. Other new chapters address timely topics such as the impact of climate change on disease, infections in U.S. veterans returning from foreign wars, and advances in contraception and the treatment of infertility. Another topic of increasing interest, the impact of aging on health and disease, is addressed by several chapters, including a newly authored chapter on the biology of aging. A new chapter on men's health complements the updated chapter on women's health. New chapters also address diverse topics including the emerging field of tissue engineering, the examination of the comatose patient, the management of heart failure, the major characteristics of helminths and helminthic infections, specific cardiac valvular disorders, venous and lymphatic diseases of the extremities, renovascular disease, late complications of diabetes, chronic myeloid leukemia, heat-related illnesses, fatigue, polyglandular failure syndromes, and nonalcoholic fatty liver disease and nonalcoholic steatohepatitis.

Supplementing new content and updates on diseases and patient care, new videos in this edition cover sleep disorders, tissue engineering, noninvasive cardiac imaging, examination of the comatose patient, and myasthenia gravis and other diseases of the neuromuscular junction. New atlases for this edition address noninvasive imaging, percutaneous revascularization, and gastrointestinal endoscopy.

We have many people to thank for their efforts in producing this book. First, the authors have done a superb job of producing authoritative chapters that synthesize vast amounts of scientific and clinical data to create state-of-the-art descriptions of medical disorders encompassed by internal medicine. In today's information-rich, rapidly evolving environment, they have ensured that this information is current. Helpful suggestions and critical input have been provided by a number of colleagues; particularly notable was the advice of Chung Owyang on the Gastroenterology Section. We are most grateful to our colleagues in each of our editorial offices who have kept track of the work in its various phases and facilitated communication with the authors, with the McGraw-Hill Education staff, and among the editors: Patricia Conrad, Patricia L. Duffey, Gregory K. Folkers, Julie B. McCoy, Elizabeth Robbins, Anita Rodriguez, and Stephanie Tribuna.

The staff at McGraw-Hill Education has been a constant source of support and expertise. James Shanahan, Associate Publisher for McGraw-Hill Education's Professional Publishing Division, has been a superb and insightful partner to the editors, guiding the development of the book and

xxxviii its related products in new formats. Kim Davis, as Associate Managing Editor, has adeptly ensured that the complex production of this multi-authored textbook proceeded smoothly and efficiently. Dominik Pucek oversaw the production of the new procedural videos and Priscilla Beer expertly oversaw the production of our extensive DVD content. Jeffrey Herzich ably served as production manager for this new edition.

We are privileged to have compiled this 19th edition and are enthusiastic about all that it offers our readers. We learned much in the process of editing *Harrison's* and hope that you will find this edition a uniquely valuable educational resource.

THE EDITORS

1 The Practice of Medicine

The Editors

THE PHYSICIAN IN THE TWENTY-FIRST CENTURY

No greater opportunity, responsibility, or obligation can fall to the lot of a human being than to become a physician. In the care of the suffering, [the physician] needs technical skill, scientific knowledge, and human understanding.... Tact, sympathy, and understanding are expected of the physician, for the patient is no mere collection of symptoms, signs, disordered functions, damaged organs, and disturbed emotions. [The patient] is human, fearful, and hopeful, seeking relief, help, and reassurance.

—Harrison's Principles of Internal Medicine, 1950

The practice of medicine has changed in significant ways since the first edition of this book appeared more than 60 years ago. The advent of molecular genetics, molecular and systems biology, and molecular pathophysiology; sophisticated new imaging techniques; and advances in bioinformatics and information technology have contributed to an explosion of scientific information that has fundamentally changed the way physicians define, diagnose, treat, and attempt to prevent disease. This growth of scientific knowledge is ongoing and accelerating.

The widespread use of electronic medical records and the Internet have altered the way doctors practice medicine and access and exchange information (Fig. 1-1). As today's physicians strive to integrate copious amounts of scientific knowledge into everyday practice, it is critically important that they remember two things: first, that the ultimate goal of medicine is to prevent disease and treat patients; and second, that despite more than 60 years of scientific advances since the first edition of this text, cultivation of the intimate relationship between physician and patient still lies at the heart of successful patient care.

THE SCIENCE AND ART OF MEDICINE

Deductive reasoning and applied technology form the foundation for the solution to many clinical problems. Spectacular advances in biochemistry, cell biology, and genomics, coupled with newly developed imaging techniques, allow access to the innermost parts of the cell and provide a window into the most remote recesses of the body. Revelations about the nature of genes and single cells have opened a portal for formulating a new molecular basis for the physiology of systems. Increasingly, physicians are learning how subtle changes in many different genes can affect the function of cells and organisms. Researchers are deciphering the complex mechanisms by which genes are regulated. Clinicians have developed a new appreciation of the role of stem cells in normal tissue function; in the development of cancer, degenerative diseases, and other disorders; and in the treatment of certain diseases. Entirely new areas of research, including studies of the human microbiome, have become important in understanding both health and disease. The knowledge gleaned from the *science of medicine* continues to enhance physicians' understanding of complex disease processes and provide new approaches to treatment and prevention. Yet skill in the most sophisticated applications of laboratory technology and in the use of the latest therapeutic modality alone does not make a good physician.

When a patient poses challenging clinical problems, an effective physician must be able to identify the crucial elements in a complex history and physical examination; order the appropriate laboratory, imaging, and diagnostic tests; and extract the key results from densely populated computer screens to determine whether to treat or to "watch." As the number of tests increases, so does the likelihood that some incidental finding, completely unrelated to the clinical problem at hand, will be uncovered. Deciding whether a clinical clue is worth pursuing or should be dismissed as a "red herring" and weighing

whether a proposed test, preventive measure, or treatment entails a greater risk than the disease itself are essential judgments that a skilled clinician must make many times each day. This combination of medical knowledge, intuition, experience, and judgment defines the *art of medicine*, which is as necessary to the practice of medicine as is a sound scientific base.

CLINICAL SKILLS

History-Taking The written history of an illness should include all the facts of medical significance in the life of the patient. Recent events should be given the most attention. Patients should, at some early point, have the opportunity to tell their own story of the illness without frequent interruption and, when appropriate, should receive expressions of interest, encouragement, and empathy from the physician. Any event related by a patient, however trivial or seemingly irrelevant, may provide the key to solving the medical problem. In general, only patients who feel comfortable with the physician will offer complete information; thus putting the patient at ease to the greatest extent possible contributes substantially to obtaining an adequate history.

An informative history is more than an orderly listing of symptoms. By listening to patients and noting the way in which they describe their symptoms, physicians can gain valuable insight. Inflections of voice, facial expression, gestures, and attitude (i.e., "body language") may offer important clues to patients' perception of their symptoms. Because patients vary in their medical sophistication and ability to recall facts, the reported medical history should be corroborated whenever possible. The social history also can provide important insights into the types of diseases that should be considered. The family history not only identifies rare Mendelian disorders within a family but often reveals risk factors for common disorders, such as coronary heart disease, hypertension, and asthma. A thorough family history may require input from multiple relatives to ensure completeness and accuracy; once recorded, it can be updated readily. The process of history-taking provides an opportunity to observe the patient's behavior and to watch for features to be pursued more thoroughly during the physical examination.

The very act of eliciting the history provides the physician with an opportunity to establish or enhance the unique bond that forms the basis for the ideal patient-physician relationship. This process helps the physician develop an appreciation of the patient's view of the illness, the patient's expectations of the physician and the health care system, and the financial and social implications of the illness for the patient. Although current health care settings may impose time constraints on patient visits, it is important not to rush the history-taking. A hurried approach may lead patients to believe that what they are relating is not of importance to the physician, and thus they may withhold relevant information. The confidentiality of the patient-physician relationship cannot be overemphasized.

Physical Examination The purpose of the physical examination is to identify physical signs of disease. The significance of these objective indications of disease is enhanced when they confirm a functional or structural change already suggested by the patient's history. At times, however, physical signs may be the only evidence of disease.

The physical examination should be methodical and thorough, with consideration given to the patient's comfort and modesty. Although attention is often directed by the history to the diseased organ or part of the body, the examination of a new patient must extend from head to toe in an objective search for abnormalities. Unless the physical examination is systematic and is performed consistently from patient to patient, important segments may be omitted inadvertently. The results of the examination, like the details of the history, should be recorded at the time they are elicited—not hours later, when they are subject to the distortions of memory. Skill in physical diagnosis is acquired with experience, but it is not merely technique that determines success in eliciting signs of disease. The detection of a few

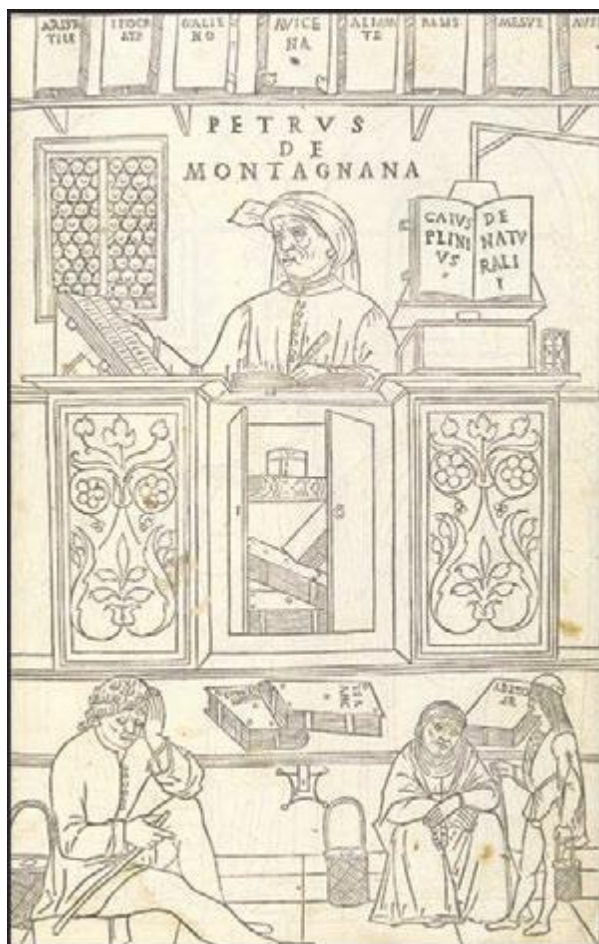


FIGURE 1-1 Woodcuts from Johannes de Ketham's *Fasciculus Medicinae*, the first illustrated medical text ever printed, show methods of information access and exchange in medical practice during the early Renaissance. Initially published in 1491 for use by medical students and practitioners, *Fasciculus Medicinae* appeared in six editions over the next 25 years. *Left:* Petrus de Montagnana, a well-known physician and teacher at the University of Padua and author of an anthology of instructive case studies, consults medical texts dating from antiquity up to the early Renaissance. *Right:* A patient with plague is attended by a physician and his attendants. (Courtesy, U.S. National Library of Medicine.)

scattered petechiae, a faint diastolic murmur, or a small mass in the abdomen is not a question of keener eyes and ears or more sensitive fingers but of a mind alert to those findings. Because physical findings can change with time, the physical examination should be repeated as frequently as the clinical situation warrants.

Given the many highly sensitive diagnostic tests now available (particularly imaging techniques), it may be tempting to place less emphasis on the physical examination. Indeed, many patients are seen by consultants after a series of diagnostic tests have been performed and the results are known. This fact should not deter the physician from performing a thorough physical examination since important clinical findings may have escaped detection by the barrage of prior diagnostic tests. The act of examining (touching) the patient also offers an opportunity for communication and may have reassuring effects that foster the patient-physician relationship.

Diagnostic Studies Physicians rely increasingly on a wide array of laboratory tests to solve clinical problems. However, accumulated laboratory data do not relieve the physician from the responsibility of carefully observing, examining, and studying the patient. It is also essential to appreciate the limitations of diagnostic tests. By virtue of their impersonal quality, complexity, and apparent precision, they often gain an aura of certainty regardless of the fallibility of the tests themselves, the instruments used in the tests, and the individuals performing or interpreting the tests. Physicians must weigh the expense involved in laboratory procedures against the value of the information these procedures are likely to provide.

Single laboratory tests are rarely ordered. Instead, physicians generally request “batteries” of multiple tests, which often prove useful.

For example, abnormalities of hepatic function may provide the clue to nonspecific symptoms such as generalized weakness and increased fatigability, suggesting a diagnosis of chronic liver disease. Sometimes a single abnormality, such as an elevated serum calcium level, points to a particular disease, such as hyperparathyroidism or an underlying malignancy.

The thoughtful use of screening tests (e.g., measurement of low-density lipoprotein cholesterol) may be of great value. A group of laboratory values can conveniently be obtained with a single specimen at relatively low cost. Screening tests are most informative when they are directed toward common diseases or disorders and when their results indicate whether other useful—but often costly—tests or interventions are needed. On the one hand, biochemical measurements, together with simple laboratory determinations such as blood count, urinalysis, and erythrocyte sedimentation rate, often provide a major clue to the presence of a pathologic process. On the other hand, the physician must learn to evaluate occasional screening-test abnormalities that do not necessarily connote significant disease. An in-depth workup after the report of an isolated laboratory abnormality in a person who is otherwise well is almost invariably wasteful and unproductive. Because so many tests are performed routinely for screening purposes, it is not unusual for one or two values to be slightly abnormal. Nevertheless, even if there is no reason to suspect an underlying illness, tests yielding abnormal results ordinarily are repeated to rule out laboratory error. If an abnormality is confirmed, it is important to consider its potential significance in the context of the patient's condition and other test results.

The development of technically improved imaging studies with greater sensitivity and specificity proceeds apace. These tests provide remarkably detailed anatomic information that can be a pivotal factor

in medical decision-making. Ultrasonography, a variety of isotopic scans, CT, MRI, and positron emission tomography have supplanted older, more invasive approaches and opened new diagnostic vistas. In light of their capabilities and the rapidity with which they can lead to a diagnosis, it is tempting to order a battery of imaging studies. All physicians have had experiences in which imaging studies revealed findings that led to an unexpected diagnosis. Nonetheless, patients must endure each of these tests, and the added cost of unnecessary testing is substantial. Furthermore, investigation of an unexpected abnormal finding may be associated with risk and/or expense and may lead to the diagnosis of an irrelevant or incidental problem. A skilled physician must learn to use these powerful diagnostic tools judiciously, always considering whether the results will alter management and benefit the patient.

PRINCIPLES OF PATIENT CARE

Evidence-Based Medicine *Evidence-based medicine* refers to the making of clinical decisions that are formally supported by data, preferably data derived from prospectively designed, randomized, controlled clinical trials. This approach is in sharp contrast to anecdotal experience, which is often biased. Unless they are attuned to the importance of using larger, more objective studies for making decisions, even the most experienced physicians can be influenced to an undue extent by recent encounters with selected patients. Evidence-based medicine has become an increasingly important part of routine medical practice and has led to the publication of many practice guidelines.

Practice Guidelines Many professional organizations and government agencies have developed formal clinical-practice guidelines to aid physicians and other caregivers in making diagnostic and therapeutic decisions that are evidence-based, cost-effective, and most appropriate to a particular patient and clinical situation. As the evidence base of medicine increases, guidelines can provide a useful framework for managing patients with particular diagnoses or symptoms. Clinical guidelines can protect patients—particularly those with inadequate health care benefits—from receiving substandard care. These guidelines also can protect conscientious caregivers from inappropriate charges of malpractice and society from the excessive costs associated with the overuse of medical resources. There are, however, caveats associated with clinical-practice guidelines since they tend to oversimplify the complexities of medicine. Furthermore, groups with different perspectives may develop divergent recommendations regarding issues as basic as the need for screening of women in their forties by mammography or of men over age 50 by serum prostate-specific antigen (PSA) assay. Finally, guidelines, as the term implies, do not—and cannot be expected to—account for the uniqueness of each individual and his or her illness. The physician's challenge is to integrate into clinical practice the useful recommendations offered by experts without accepting them blindly or being inappropriately constrained by them.

Medical Decision-Making Medical decision-making is an important responsibility of the physician and occurs at each stage of the diagnostic and therapeutic process. The decision-making process involves the ordering of additional tests, requests for consultations, and decisions about treatment and predictions concerning prognosis. This process requires an in-depth understanding of the pathophysiology and natural history of disease. As discussed above, medical decision-making should be evidence-based so that patients derive full benefit from the available scientific knowledge. Formulating a differential diagnosis requires not only a broad knowledge base but also the ability to assess the relative probabilities of various diseases. Application of the scientific method, including hypothesis formulation and data collection, is essential to the process of accepting or rejecting a particular diagnosis. Analysis of the differential diagnosis is an iterative process. As new information or test results are acquired, the group of disease processes being considered can be contracted or expanded appropriately.

Despite the importance of evidence-based medicine, much medical decision-making relies on good clinical judgment, an attribute that is difficult to quantify or even to assess qualitatively. Physicians must use

their knowledge and experience as a basis for weighing known factors, along with the inevitable uncertainties, and then making a sound judgment; this synthesis of information is particularly important when a relevant evidence base is not available. Several quantitative tools may be invaluable in synthesizing the available information, including diagnostic tests, Bayes' theorem, and multivariate statistical models. *Diagnostic tests* serve to reduce uncertainty about an individual's diagnosis or prognosis and help the physician decide how best to manage that individual's condition. The battery of diagnostic tests complements the history and the physical examination. The accuracy of a particular test is ascertained by determining its sensitivity (true-positive rate) and specificity (true-negative rate) as well as the predictive value of a positive and a negative result. *Bayes' theorem* uses information on a test's sensitivity and specificity, in conjunction with the pretest probability of a diagnosis, to determine mathematically the posttest probability of the diagnosis. More complex clinical problems can be approached with *multivariate statistical models*, which generate highly accurate information even when multiple factors are acting individually or together to affect disease risk, progression, or response to treatment. Studies comparing the performance of statistical models with that of expert clinicians have documented equivalent accuracy, although the models tend to be more consistent. Thus, multivariate statistical models may be particularly helpful to less experienced clinicians. [See Chap. 3 for a more thorough discussion of decision-making in clinical medicine.](#)

Electronic Medical Records Both the growing reliance on computers and the strength of information technology now play central roles in medicine. Laboratory data are accessed almost universally through computers. Many medical centers now have electronic medical records, computerized order entry, and bar-coded tracking of medications. Some of these systems are interactive, sending reminders or warning of potential medical errors.

Electronic medical records offer rapid access to information that is invaluable in enhancing health care quality and patient safety, including relevant data, historical and clinical information, imaging studies, laboratory results, and medication records. These data can be used to monitor and reduce unnecessary variations in care and to provide real-time information about processes of care and clinical outcomes. Ideally, patient records are easily transferred across the health care system. However, technologic limitations and concerns about privacy and cost continue to limit broad-based use of electronic health records in many clinical settings.

As valuable as it is, information technology is merely a tool and can never replace the clinical decisions that are best made by the physician. Clinical knowledge and an understanding of a patient's needs, supplemented by quantitative tools, still represent the best approach to decision-making in the practice of medicine.

Evaluation of Outcomes Clinicians generally use *objective* and readily measurable parameters to judge the outcome of a therapeutic intervention. These measures may oversimplify the complexity of a clinical condition as patients often present with a major clinical problem in the context of multiple complicating background illnesses. For example, a patient may present with chest pain and cardiac ischemia, but with a background of chronic obstructive pulmonary disease and renal insufficiency. For this reason, outcome measures such as mortality, length of hospital stay, or readmission rates are typically risk-adjusted. An important point is that patients usually seek medical attention for *subjective* reasons; they wish to obtain relief from pain, to preserve or regain function, and to enjoy life. The components of a patient's health status or quality of life can include bodily comfort, capacity for physical activity, personal and professional function, sexual function, cognitive function, and overall perception of health. Each of these important areas can be assessed through structured interviews or specially designed questionnaires. Such assessments provide useful parameters by which a physician can judge patients' subjective views of their disabilities and responses to treatment, particularly in chronic illness. The practice of medicine requires consideration and integration of both objective and subjective outcomes.

Women's Health and Disease Although past epidemiologic studies and clinical trials have often focused predominantly on men, more recent studies have included more women, and some, like the Women's Health Initiative, have exclusively addressed women's health issues. Significant sex-based differences exist in diseases that afflict both men and women. Much is still to be learned in this arena, and ongoing studies should enhance physicians' understanding of the mechanisms underlying these differences in the course and outcome of certain diseases. **For a more complete discussion of women's health, see Chap. 6e.**

Care of the Elderly The relative proportion of elderly individuals in the populations of developed nations has grown considerably over the past few decades and will continue to grow. The practice of medicine is greatly influenced by the health care needs of this growing demographic group. The physician must understand and appreciate the decline in physiologic reserve associated with aging; the differences in appropriate doses, clearance, and responses to medications; the diminished responses of the elderly to vaccinations such as those against influenza; the different manifestations of common diseases among the elderly; and the disorders that occur commonly with aging, such as depression, dementia, frailty, urinary incontinence, and fractures. **For a more complete discussion of medical care for the elderly, see Chap. 11 and Part 5, Chaps. 93e and 94e.**

Errors in the Delivery of Health Care A 1999 report from the Institute of Medicine called for an ambitious agenda to reduce medical error rates and improve patient safety by designing and implementing fundamental changes in health care systems. Adverse drug reactions occur in at least 5% of hospitalized patients, and the incidence increases with the use of a large number of drugs. Whatever the clinical situation, it is the physician's responsibility to use powerful therapeutic measures wisely, with due regard for their beneficial actions, potential dangers, and cost. It is the responsibility of hospitals and health care organizations to develop systems to reduce risk and ensure patient safety. Medication errors can be reduced through the use of ordering systems that rely on electronic processes or, when electronic options are not available, that eliminate misreading of handwriting. Implementation of infection control systems, enforcement of hand-washing protocols, and careful oversight of antibiotic use can minimize the complications of nosocomial infections. Central-line infection rates have been dramatically reduced at many centers by careful adherence of trained personnel to standardized protocols for introducing and maintaining central lines. Rates of surgical infection and wrong-site surgery can likewise be reduced by the use of standardized protocols and checklists. Falls by patients can be minimized by judicious use of sedatives and appropriate assistance with bed-to-chair and bed-to-bathroom transitions. Taken together, these and other measures are saving thousands of lives each year.

The Physician's Role in Informed Consent The fundamental principles of medical ethics require physicians to act in the patient's best interest and to respect the patient's autonomy. These requirements are particularly relevant to the issue of informed consent. Patients are required to sign a consent form for essentially any diagnostic or therapeutic procedure. Most patients possess only limited medical knowledge and must rely on their physicians for advice. Communicating in a clear and understandable manner, physicians must fully discuss the alternatives for care and explain the risks, benefits, and likely consequences of each alternative. In every case, the physician is responsible for ensuring that the patient thoroughly understands these risks and benefits; encouraging questions is an important part of this process. This is the very definition of *informed consent*. Full, clear explanation and discussion of the proposed procedures and treatment can greatly mitigate the fear of the unknown that commonly accompanies hospitalization. Excellent communication can also help alleviate misunderstandings in situations where complications of intervention occur. Often the patient's understanding is enhanced by repeatedly discussing the issues in an unthreatening and supportive way, answering new questions that occur to the patient as they arise.

Special care should be taken to ensure that a physician seeking a patient's informed consent has no real or apparent conflict of interest involving personal gain.

The Approach to Grave Prognoses and Death No circumstance is more distressing than the diagnosis of an incurable disease, particularly when premature death is inevitable. What should the patient and family be told? What measures should be taken to maintain life? What can be done to maintain the quality of life?

Honesty is absolutely essential in the face of a terminal illness. The patient must be given an opportunity to talk with the physician and ask questions. A wise and insightful physician uses such open communication as the basis for assessing what the patient wants to know and when he or she wants to know it. On the basis of the patient's responses, the physician can assess the right tempo for sharing information. Ultimately, the patient must understand the expected course of the disease so that appropriate plans and preparations can be made. The patient should participate in decision-making with an understanding of the goal of treatment (palliation) and its likely effects. The patient's religious beliefs must be taken into consideration. Some patients may find it easier to share their feelings about death with their physician, who is likely to be more objective and less emotional, than with family members.

The physician should provide or arrange for emotional, physical, and spiritual support and must be compassionate, unhurried, and open. In many instances, there is much to be gained by the laying on of hands. Pain should be controlled adequately, human dignity maintained, and isolation from family and close friends avoided. These aspects of care tend to be overlooked in hospitals, where the intrusion of life-sustaining equipment can detract from attention to the whole person and encourage concentration instead on the life-threatening disease, against which the battle ultimately will be lost in any case. In the face of terminal illness, the goal of medicine must shift from *cure* to *care* in the broadest sense of the term. *Primum succurrere*, first hasten to help, is a guiding principle. In offering care to a dying patient, a physician must be prepared to provide information to family members and deal with their grief and sometimes their feelings of guilt or even anger. It is important for the doctor to assure the family that everything reasonable has been done. A substantial problem in these discussions is that the physician often does not know how to gauge the prognosis. In addition, various members of the health care team may offer different opinions. Good communication among providers is essential so that consistent information is provided to patients. This is especially important when the best path forward is uncertain. Advice from experts in palliative and terminal care should be sought whenever necessary to ensure that clinicians are not providing patients with unrealistic expectations. **For a more complete discussion of end-of-life care, see Chap. 10.**

THE PATIENT-PHYSICIAN RELATIONSHIP

The significance of the intimate personal relationship between physician and patient cannot be too strongly emphasized, for in an extraordinarily large number of cases both the diagnosis and treatment are directly dependent on it. One of the essential qualities of the clinician is interest in humanity, for the secret of the care of the patient is in caring for the patient.

—Francis W. Peabody, October 21, 1925,
Lecture at Harvard Medical School

Physicians must never forget that patients are individual human beings with problems that all too often transcend their physical complaints. They are not "cases" or "admissions" or "diseases." Patients do not fail treatments; treatments fail to benefit patients. This point is particularly important in this era of high technology in clinical medicine. Most patients are anxious and fearful. Physicians should instill confidence and offer reassurance but must never come across as arrogant or patronizing. A professional attitude, coupled with warmth and openness, can do much to alleviate anxiety and to encourage patients to share all aspects of their medical history. Empathy and compassion are the essential features of a caring physician. The physician needs to consider the setting in which an illness occurs—in terms not only of patients themselves but also of their familial, social, and cultural backgrounds. The ideal patient-physician relationship is based on thorough knowledge of the patient, mutual trust, and the ability to communicate.

The Dichotomy of Inpatient and Outpatient Internal Medicine The hospital environment has changed dramatically over the last few decades. Emergency departments and critical care units have evolved to identify and manage critically ill patients, allowing them to survive formerly fatal diseases. At the same time, there is increasing pressure to reduce the length of stay in the hospital and to manage complex disorders in the outpatient setting. This transition has been driven not only by efforts to reduce costs but also by the availability of new outpatient technologies, such as imaging and percutaneous infusion catheters for long-term antibiotics or nutrition, minimally invasive surgical procedures, and evidence that outcomes often are improved by minimizing inpatient hospitalization.

In these circumstances, two important issues arise as physicians cope with the complexities of providing care for hospitalized patients. On the one hand, highly specialized health professionals are essential to the provision of optimal acute care in the hospital; on the other, these professionals—with their diverse training, skills, responsibilities, experiences, languages, and “cultures”—need to work as a team.

In addition to traditional medical beds, hospitals now encompass multiple distinct levels of care, such as the emergency department, procedure rooms, overnight observation units, critical care units, and palliative care units. A consequence of this differentiation has been the emergence of new trends, including specialties (e.g., emergency medicine and end-of-life care) and the provision of in-hospital care by hospitalists and intensivists. Most *hospitalists* are board-certified internists who bear primary responsibility for the care of hospitalized patients and whose work is limited entirely to the hospital setting. The shortened length of hospital stay that is now standard means that most patients receive only acute care while hospitalized; the increased complexities of inpatient medicine make the presence of a generalist with specific training, skills, and experience in the hospital environment extremely beneficial. *Intensivists* are board-certified physicians who are further certified in critical care medicine and who direct and provide care for very ill patients in critical care units. Clearly, then, an important challenge in internal medicine today is to ensure the continuity of communication and information flow between a patient’s primary care doctor and these physicians who are in charge of the patient’s hospital care. Maintaining these channels of communication is frequently complicated by patient “handoffs”—i.e., from the outpatient to the inpatient environment, from the critical care unit to a general medicine floor, and from the hospital to the outpatient environment. The involvement of many care providers in conjunction with these transitions can threaten the traditional one-to-one relationship between patient and primary care physician. Of course, patients can benefit greatly from effective collaboration among a number of health care professionals; however, *it is the duty of the patient’s principal or primary physician to provide cohesive guidance through an illness.* To meet this challenge, primary care physicians must be familiar with the techniques, skills, and objectives of specialist physicians and allied health professionals who care for their patients in the hospital. In addition, primary care doctors must ensure that their patients will benefit from scientific advances and from the expertise of specialists when they are needed both in and out of the hospital. Primary care physicians can also explain the role of these specialists to reassure patients that they are in the hands of the physicians best trained to manage an acute illness. However, the primary care physician should retain ultimate responsibility for making major decisions about diagnosis and treatment and should assure patients and their families that decisions are being made in consultation with these specialists by a physician who has an overall and complete perspective on the case.

A key factor in mitigating the problems associated with multiple care providers is a commitment to interprofessional teamwork. Despite the diversity in training, skills, and responsibilities among health care professionals, common values need to be reinforced if patient care is not to be adversely affected. This component of effective medical care is widely recognized, and several medical schools have integrated interprofessional teamwork into their curricula. The evolving concept of the “medical home” incorporates team-based primary care with linked subspecialty care in a cohesive environment that ensures smooth transitions of care cost-effectively.

Appreciation of the Patient’s Hospital Experience The hospital is an intimidating environment for most individuals. Hospitalized patients find themselves surrounded by air jets, buttons, and glaring lights; invaded by tubes and wires; and beset by the numerous members of the health care team—hospitalists, specialists, nurses, nurses’ aides, physicians’ assistants, social workers, technologists, physical therapists, medical students, house officers, attending and consulting physicians, and many others. They may be transported to special laboratories and imaging facilities replete with blinking lights, strange sounds, and unfamiliar personnel; they may be left unattended at times; and they may be obligated to share a room with other patients who have their own health problems. It is little wonder that a patient’s sense of reality may be compromised. Physicians who appreciate the hospital experience from the patient’s perspective and who make an effort to develop a strong relationship within which they can guide the patient through this experience may make a stressful situation more tolerable.

Trends in the Delivery of Health Care: A Challenge to the Humane Physician Many trends in the delivery of health care tend to make medical care impersonal. These trends, some of which have been mentioned already, include (1) vigorous efforts to reduce the escalating costs of health care; (2) the growing number of managed-care programs, which are intended to reduce costs but in which the patient may have little choice in selecting a physician or in seeing that physician consistently; (3) increasing reliance on technological advances and computerization for many aspects of diagnosis and treatment; and (4) the need for numerous physicians to be involved in the care of most patients who are seriously ill.

In light of these changes in the medical care system, it is a major challenge for physicians to maintain the *humane* aspects of medical care. The American Board of Internal Medicine, working together with the American College of Physicians–American Society of Internal Medicine and the European Federation of Internal Medicine, has published a *Charter on Medical Professionalism* that underscores three main principles in physicians’ contract with society: (1) the primacy of patient welfare, (2) patient autonomy, and (3) social justice. While medical schools appropriately place substantial emphasis on professionalism, a physician’s personal attributes, including integrity, respect, and compassion, also are extremely important. Availability to the patient, expression of sincere concern, willingness to take the time to explain all aspects of the illness, and a nonjudgmental attitude when dealing with patients whose cultures, lifestyles, attitudes, and values differ from those of the physician are just a few of the characteristics of a humane physician. Every physician will, at times, be challenged by patients who evoke strongly negative or positive emotional responses. Physicians should be alert to their own reactions to such patients and situations and should consciously monitor and control their behavior so that the patient’s best interest remains the principal motivation for their actions at all times.

An important aspect of patient care involves an appreciation of the patient’s “quality of life,” a subjective assessment of what each patient values most. This assessment requires detailed, sometimes intimate knowledge of the patient, which usually can be obtained only through deliberate, unhurried, and often repeated conversations. Time pressures will always threaten these interactions, but they should not diminish the importance of understanding and seeking to fulfill the priorities of the patient.

EXPANDING FRONTIERS IN MEDICAL PRACTICE

The Era of “Omics”: Genomics, Epigenomics, Proteomics, Microbiomics, Metagenomics, Metabolomics, Exposomics . . . In the spring of 2003, announcement of the complete sequencing of the human genome officially ushered in the genomic era. However, even before that landmark accomplishment, the practice of medicine had been evolving as a result of the insights into both the human genome and the genomes of a wide variety of microbes. The clinical implications of these insights are illustrated by the complete genome sequencing of H1N1 influenza virus in 2009 and the rapid identification of H1N1 influenza as a potentially fatal pandemic illness, with swift development and dissemination of an effective protective vaccine. Today, gene expression profiles are being

used to guide therapy and inform prognosis for a number of diseases, the use of genotyping is providing a new means to assess the risk of certain diseases as well as variations in response to a number of drugs, and physicians are better understanding the role of certain genes in the causality of common conditions such as obesity and allergies. Despite these advances, the use of complex genomics in the diagnosis, prevention, and treatment of disease is still in its early stages. The task of physicians is complicated by the fact that phenotypes generally are determined not by genes alone but by the interplay of genetic and environmental factors. Indeed, researchers have just begun to scratch the surface of the potential applications of genomics in the practice of medicine.

Rapid progress also is being made in other areas of molecular medicine. *Epigenomics* is the study of alterations in chromatin and histone proteins and methylation of DNA sequences that influence gene expression. Every cell of the body has identical DNA sequences; the diverse phenotypes a person's cells manifest are the result of epigenetic regulation of gene expression. Epigenetic alterations are associated with a number of cancers and other diseases. *Proteomics*, the study of the entire library of proteins made in a cell or organ and its complex relationship to disease, is enhancing the repertoire of the 23,000 genes in the human genome through alternate splicing, post-translational processing, and posttranslational modifications that often have unique functional consequences. The presence or absence of particular proteins in the circulation or in cells is being explored for diagnostic and disease-screening applications. *Microbiomics* is the study of the resident microbes in humans and other mammals. The human haploid genome has ~20,000 genes, while the microbes residing on and in the human body comprise over 3–4 million genes; the contributions of these resident microbes are likely to be of great significance with regard to health status. In fact, research is demonstrating that the microbes inhabiting human mucosal and skin surfaces play a critical role in maturation of the immune system, in metabolic balance, and in disease susceptibility. A variety of environmental factors, including the use and overuse of antibiotics, have been tied experimentally to substantial increases in disorders such as obesity, metabolic syndrome, atherosclerosis, and immune-mediated diseases in both adults and children. *Metagenomics*, of which microbiomics is a part, is the genomic study of environmental species that have the potential to influence human biology directly or indirectly. An example is the study of exposures to microorganisms in farm environments that may be responsible for the lower incidence of asthma among children raised on farms. *Metabolomics* is the study of the range of metabolites in cells or organs and the ways they are altered in disease states. The aging process itself may leave telltale metabolic footprints that allow the prediction (and possibly the prevention) of organ dysfunction and disease. It seems likely that disease-associated patterns will be sought in lipids, carbohydrates, membranes, mitochondria, and other vital components of cells and tissues. Finally, *exposomics* refers to efforts to catalogue and capture environmental exposures such as smoking, sunlight, diet, exercise, education, and violence, which together have an enormous impact on health. All of this new information represents a challenge to the traditional reductionist approach to medical thinking. The variability of results in different patients, together with the large number of variables that can be assessed, creates difficulties in identifying pre-clinical disease and defining disease states unequivocally. Accordingly, the tools of *systems biology* and *network medicine* are being applied to the enormous body of information now obtainable for every patient and may eventually provide new approaches to classifying disease. **For a more complete discussion of a complex systems approach to human disease, see Chap. 87e.**

The rapidity of these advances may seem overwhelming to practicing physicians. However, physicians have an important role to play in ensuring that these powerful technologies and sources of new information are applied with sensitivity and intelligence to the patient. Since “omics” are evolving so rapidly, physicians and other health care professionals must continue to educate themselves so that they can apply this new knowledge to the benefit of their patients' health and well-being. Genetic testing requires wise counsel based on an understanding of the value and limitations of the tests as well as the implications of

their results for specific individuals. **For a more complete discussion of genetic testing, see Chap. 84.**

The Globalization of Medicine Physicians should be cognizant of diseases and health care services beyond local boundaries. Global travel has implications for disease spread, and it is not uncommon for diseases endemic to certain regions to be seen in other regions after a patient has traveled to and returned from those regions. In addition, factors such as wars, the migration of refugees, and climate change are contributing to changing disease profiles worldwide. Patients have broader access to unique expertise or clinical trials at distant medical centers, and the cost of travel may be offset by the quality of care at those distant locations. As much as any other factor influencing global aspects of medicine, the Internet has transformed the transfer of medical information throughout the world. This change has been accompanied by the transfer of technological skills through telemedicine and international consultation—for example, regarding radiologic images and pathologic specimens. **For a complete discussion of global issues, see Chap. 2.**

Medicine on the Internet On the whole, the Internet has had a very positive effect on the practice of medicine; through personal computers, a wide range of information is available to physicians and patients almost instantaneously at any time and from anywhere in the world. This medium holds enormous potential for the delivery of current information, practice guidelines, state-of-the-art conferences, journal content, textbooks (including this text), and direct communications with other physicians and specialists, expanding the depth and breadth of information available to the physician regarding the diagnosis and care of patients. Medical journals are now accessible online, providing rapid sources of new information. By bringing them into direct and timely contact with the latest developments in medical care, this medium also serves to lessen the information gap that has hampered physicians and health care providers in remote areas.

Patients, too, are turning to the Internet in increasing numbers to acquire information about their illnesses and therapies and to join Internet-based support groups. Patients often arrive at a clinic visit with sophisticated information about their illnesses. In this regard, physicians are challenged in a positive way to keep abreast of the latest relevant information while serving as an “editor” as patients navigate this seemingly endless source of information, the accuracy and validity of which are not uniform.

A critically important caveat is that virtually anything can be published on the Internet, with easy circumvention of the peer-review process that is an essential feature of academic publications. Both physicians and patients who search the Internet for medical information must be aware of this danger. Notwithstanding this limitation, appropriate use of the Internet is revolutionizing information access for physicians and patients and in this regard represents a remarkable resource that was not available to practitioners a generation ago.

Public Expectations and Accountability The general public's level of knowledge and sophistication regarding health issues has grown rapidly over the last few decades. As a result, expectations of the health care system in general and of physicians in particular have risen. Physicians are expected to master rapidly advancing fields (the *science* of medicine) while considering their patients' unique needs (the *art* of medicine). Thus, physicians are held accountable not only for the technical aspects of the care they provide but also for their patients' satisfaction with the delivery and costs of care.

In many parts of the world, physicians increasingly are expected to account for the way in which they practice medicine by meeting certain standards prescribed by federal and local governments. The hospitalization of patients whose health care costs are reimbursed by the government and other third parties is subjected to utilization review. Thus, a physician must defend the cause for and duration of a patient's hospitalization if it falls outside certain “average” standards. Authorization for reimbursement increasingly is based on documentation of the nature and complexity of an illness, as reflected by recorded elements of the history and physical examination. A growing “pay-for-performance” movement seeks to link reimbursement to quality of

care. The goal of this movement is to improve standards of health care and contain spiraling health care costs. In many parts of the United States, managed (capitated) care contracts with insurers have replaced traditional fee-for-service care, placing the onus of managing the cost of all care directly on the providers and increasing the emphasis on preventive strategies. In addition, physicians are expected to give evidence of their current competence through mandatory continuing education, patient record audits, maintenance of certification, and relicensing.

Medical Ethics and New Technologies The rapid pace of technological advance has profound implications for medical applications that go far beyond the traditional goals of disease prevention, treatment, and cure. Cloning, genetic engineering, gene therapy, human-computer interfaces, nanotechnology, and use of designer drugs have the potential to modify inherited predispositions to disease, select desired characteristics in embryos, augment “normal” human performance, replace failing tissues, and substantially prolong life span. Given their unique training, physicians have a responsibility to help shape the debate on the appropriate uses of and limits placed on these new techniques and to consider carefully the ethical issues associated with the implementation of such interventions.

The Physician as Perpetual Student From the time doctors graduate from medical school, it becomes all too apparent that their lot is that of the “perpetual student” and that the mosaic of their knowledge and experiences is eternally unfinished. This realization is at the same time exhilarating and anxiety-provoking. It is exhilarating because doctors can apply constantly expanding knowledge to the treatment of their patients; it is anxiety-provoking because doctors realize that they will never know as much as they want or need to know. Ideally, doctors will translate the latter feeling into energy through which they can continue to improve themselves and reach their potential as physicians. It is the physician’s responsibility to pursue new knowledge continually by reading, attending conferences and courses, and consulting colleagues and the Internet. This is often a difficult task for a busy practitioner; however, a commitment to continued learning is an integral part of being a physician and must be given the highest priority.

The Physician as Citizen Being a physician is a privilege. The capacity to apply one’s skills for the benefit of one’s fellow human beings is a noble calling. The doctor-patient relationship is inherently unbalanced in the distribution of power. In light of their influence, physicians must always be aware of the potential impact of what they do and say and must always strive to strip away individual biases and preferences to find what is best for the patient. To the extent possible, physicians should also act within their communities to promote health and alleviate suffering. Meeting these goals begins by setting a healthy example and continues in taking action to deliver needed care even when personal financial compensation may not be available.

A goal for medicine and its practitioners is to strive to provide the means by which the poor can cease to be unwell.

Learning Medicine It has been a century since the publication of the Flexner Report, a seminal study that transformed medical education and emphasized the scientific foundations of medicine as well as the acquisition of clinical skills. In an era of burgeoning information and access to medical simulation and informatics, many schools are implementing new curricula that emphasize lifelong learning and the acquisition of competencies in teamwork, communication skills, system-based practice, and professionalism. These and other features of the medical school curriculum provide the foundation for many of the themes highlighted in this chapter and are expected to allow physicians to progress, with experience and learning over time, from competency to proficiency to mastery.

At a time when the amount of information that must be mastered to practice medicine continues to expand, increasing pressures both within and outside of medicine have led to the implementation of restrictions on the amount of time a physician-in-training can spend in the hospital. Because the benefits associated with continuity of medical care and observation of a patient’s progress over time were thought

to be outstripped by the stresses imposed on trainees by long hours and by the fatigue-related errors they made in caring for patients, strict limits were set on the number of patients that trainees could be responsible for at one time, the number of new patients they could evaluate in a day on call, and the number of hours they could spend in the hospital. In 1980, residents in medicine worked in the hospital more than 90 hours per week on average. In 1989, their hours were restricted to no more than 80 per week. Resident physicians’ hours further decreased by ~10% between 1996 and 2008, and in 2010 the Accreditation Council for Graduate Medical Education further restricted (i.e., to 16 hours per shift) consecutive in-hospital duty hours for first-year residents. The impact of these changes is still being assessed, but the evidence that medical errors have decreased as a consequence is sparse. An unavoidable by-product of fewer hours at work is an increase in the number of “handoffs” of patient responsibility from one physician to another. These transfers often involve a transition from a physician who knows the patient well, having evaluated that individual on admission, to a physician who knows the patient less well. It is imperative that these transitions of responsibility be handled with care and thoroughness, with all relevant information exchanged and acknowledged.

Research, Teaching, and the Practice of Medicine The word *doctor* is derived from the Latin *docere*, “to teach.” As teachers, physicians should share information and medical knowledge with colleagues, students of medicine and related professions, and their patients. The practice of medicine is dependent on the sum total of medical knowledge, which in turn is based on an unending chain of scientific discovery, clinical observation, analysis, and interpretation. Advances in medicine depend on the acquisition of new information through research, and improved medical care requires the transmission of that information. As part of their broader societal responsibilities, physicians should encourage patients to participate in ethical and properly approved clinical investigations if these studies do not impose undue hazard, discomfort, or inconvenience. However, physicians engaged in clinical research must be alert to potential conflicts of interest between their research goals and their obligations to individual patients. The best interests of the patient must always take priority.

To wrest from nature the secrets which have perplexed philosophers in all ages, to track to their sources the causes of disease, to correlate the vast stores of knowledge, that they may be quickly available for the prevention and cure of disease—these are our ambitions.

—William Osler, 1849–1919

2

Global Issues in Medicine

Paul Farmer, Joseph Rhatigan

WHY GLOBAL HEALTH?

Global health is not a discipline; it is, rather, a collection of problems. Some scholars have defined global health as the field of study and practice concerned with improving the health of all people and achieving health equity worldwide, with an emphasis on addressing transnational problems. No single review can do much more than identify the leading problems in applying evidence-based medicine in settings of great poverty or across national boundaries. However, this is a moment of opportunity: only recently, persistent epidemics, improved metrics, and growing interest have been matched by an unprecedented investment in addressing the health problems of poor people in the developing world. To ensure that this opportunity is not wasted, the facts need to be laid out for specialists and laypeople alike. This chapter introduces the major international bodies that address health problems; identifies the more significant barriers to improving the health of people who to date have not, by and large, had access

to modern medicine; and summarizes population-based data on the most common health problems faced by people living in poverty. Examining specific problems—notably HIV/AIDS (**Chap. 226**) but also tuberculosis (TB, **Chap. 202**), malaria (**Chap. 248**), and key “non-communicable” chronic diseases (NCDs)—helps sharpen the discussion of barriers to prevention, diagnosis, and care as well as the means of overcoming them. This chapter closes by discussing global health equity, drawing on notions of social justice that once were central to international public health but had fallen out of favor during the last decades of the twentieth century.

A BRIEF HISTORY OF GLOBAL HEALTH INSTITUTIONS

Concern about health across national boundaries dates back many centuries, predating the Black Plague and other pandemics. One of the first organizations founded explicitly to tackle cross-border health issues was the Pan American Sanitary Bureau, which was formed in 1902 by 11 countries in the Americas. The primary goal of what later became the Pan American Health Organization was the control of infectious diseases across the Americas. Of special concern was yellow fever, which had been running a deadly course through much of South and Central America and halted the construction of the Panama Canal. In 1948, the United Nations formed the first truly global health institution: the World Health Organization (WHO). In 1958, under the aegis of the WHO and in line with a long-standing focus on communicable diseases that cross borders, leaders in global health initiated the effort that led to what some see as the greatest success in international health: the eradication of smallpox. Naysayers were surprised when the smallpox eradication campaign, which engaged public health officials throughout the world, proved successful in 1979 despite the ongoing Cold War.

At the International Conference on Primary Health Care in Alma-Ata (in what is now Kazakhstan) in 1978, public health officials from around the world agreed on a commitment to “Health for All by the Year 2000,” a goal to be achieved by providing universal access to primary health care worldwide. Critics argued that the attainment of this goal by the proposed date was impossible. In the ensuing years, a strategy for the provision of selective primary health care emerged that included four inexpensive interventions collectively known as GOBI: growth monitoring, oral rehydration, breast-feeding, and immunizations for diphtheria, whooping cough, tetanus, polio, TB, and measles. GOBI later was expanded to GOBI-FFF, which also included female education, food, and family planning. Some public health figures saw GOBI-FFF as an interim strategy to achieve “health for all,” but others criticized it as a retreat from the bolder commitments of Alma-Ata.

The influence of the WHO waned during the 1980s. In the early 1990s, many observers argued that, with its vastly superior financial resources and its close—if unequal—relationships with the governments of poor countries, the World Bank had eclipsed the WHO as the most important multilateral institution working in the area of health. One of the stated goals of the World Bank was to help poor countries identify “cost-effective” interventions worthy of public funding and international support. At the same time, the World Bank encouraged many of those nations to reduce public expenditures in health and education in order to stimulate economic growth as part of (later discredited) structural adjustment programs whose restrictions were imposed as a condition for access to credit and assistance through international financial institutions such as the World Bank and the International Monetary Fund. There was a resurgence of many diseases, including malaria, trypanosomiasis, and schistosomiasis, in Africa. TB, an eminently curable disease, remained the world’s leading infectious killer of adults. Half a million women per year died in childbirth during the last decade of the twentieth century, and few of the world’s largest philanthropic or funding institutions focused on global health equity.

HIV/AIDS, first described in 1981, precipitated a change. In the United States, the advent of this newly described infectious killer marked the culmination of a series of events that discredited talk of “closing the book” on infectious diseases. In Africa, which would emerge as the global epicenter of the pandemic, HIV disease strained TB control programs, and malaria continued to take as many lives as

ever. At the dawn of the twenty-first century, these three diseases alone killed nearly 6 million people each year. New research, new policies, and new funding mechanisms were called for. The past decade has seen the rise of important multilateral global health financing institutions such as the Global Fund to Fight AIDS, Tuberculosis, and Malaria; bilateral efforts such as the U.S. President’s Emergency Plan for AIDS Relief (PEPFAR); and private philanthropic organizations such as the Bill & Melinda Gates Foundation. With its 193 member states and 147 country offices, the WHO remains important in matters relating to the cross-border spread of infectious diseases and other health threats. In the aftermath of the epidemic of severe acute respiratory syndrome in 2003, the WHO’s International Health Regulations—which provide a legal foundation for that organization’s direct investigation into a wide range of global health problems, including pandemic influenza, in any member state—were strengthened and brought into force in May 2007.

Even as attention to and resources for health problems in poor countries grow, the lack of coherence in and among global health institutions may undermine efforts to forge a more comprehensive and effective response. The WHO remains underfunded despite the ever-growing need to engage a wider and more complex range of health issues. In another instance of the paradoxical impact of success, the rapid growth of the Gates Foundation, which is one of the most important developments in the history of global health, has led some foundations to question the wisdom of continuing to invest their more modest resources in this field. This indeed may be what some have called “the golden age of global health,” but leaders of major organizations such as the WHO, the Global Fund, the United Nations Children’s Fund (UNICEF), the Joint United Nations Programme on HIV/AIDS (UNAIDS), PEPFAR, and the Gates Foundation must work together to design an effective architecture that will make the most of opportunities to link new resources for and commitments to global health equity with the emerging understanding of disease burden and unmet need. To this end, new and old players in global health must invest heavily in *discovery* (relevant basic science), *development* of new tools (preventive, diagnostic, and therapeutic), and modes of *delivery* that will ensure the equitable provision of health products and services to all who need them.

THE ECONOMICS OF GLOBAL HEALTH

Political and economic concerns have often guided global health interventions. As mentioned, early efforts to control yellow fever were tied to the completion of the Panama Canal. However, the precise nature of the link between economics and health remains a matter for debate. Some economists and demographers argue that improving the health status of populations must begin with economic development; others maintain that addressing ill health is the starting point for development in poor countries. In either case, investment in health care, especially the control of communicable diseases, should lead to increased productivity. The question is where to find the necessary resources to start the predicted “virtuous cycle.”

During the past two decades, spending on health in poor countries has increased dramatically. According to a study from the Institute for Health Metrics and Evaluation (IHME) at the University of Washington, total development assistance for health worldwide grew to \$28.2 billion in 2010—up from \$5.6 billion in 1990. In 2010, the leading contributors included U.S. bilateral agencies such as PEPFAR, the Global Fund, nongovernmental organizations (NGOs), the WHO, the World Bank, and the Gates Foundation. It appears, however, that total development assistance for health plateaued in 2010, and it is unclear whether growth will continue in the upcoming decade.

To reach the United Nations Millennium Development Goals, which include targets for poverty reduction, universal primary education, and gender equality, spending in the health sector must be increased above the 2010 levels. To determine by how much and for how long, it is imperative to improve the ability to assess the global burden of disease and to plan interventions that more precisely match need.

MORTALITY AND THE GLOBAL BURDEN OF DISEASE

Refining metrics is an important task for global health: only recently have there been solid assessments of the global burden of disease. The first study to look seriously at this issue, conducted in 1990, laid the foundation for the first report on *Disease Control Priorities in Developing Countries* and for the World Bank's 1993 World Development Report *Investing in Health*. Those efforts represented a major advance in the understanding of health status in developing countries. *Investing in Health* has been especially influential: it familiarized a broad audience with cost-effectiveness analysis for specific health interventions and with the notion of disability-adjusted life years (DALYs). The DALY, which has become a standard measure of the impact of a specific health condition on a population, combines absolute years of life lost and years lost due to disability for incident cases of a condition. (See Fig. 2-1 and Table 2-1 for an analysis of the global disease burden by DALYs.)

In 2012, the IHME and partner institutions began publishing results from the Global Burden of Diseases, Injuries, and Risk Factors Study 2010 (GBD 2010). GBD 2010 is the most comprehensive effort to date to produce longitudinal, globally complete, and comparable estimates of the burden of diseases, injuries, and risk factors. This report reflects the expansion of the available data on health in the poorest countries and of the capacity to quantify the impact of specific conditions on a population. It measures current levels and recent trends in all major diseases, injuries, and risk factors among 21 regions and for 20 age groups and both sexes. The GBD 2010 team revised and improved the health-state severity weight system, collated published data, and used household surveys to enhance the breadth and accuracy of disease burden data. As analytic methods and data quality improve, important trends can be identified in a comparison of global disease burden estimates from 1990 to 2010.

GLOBAL MORTALITY

Of the 52.8 million deaths worldwide in 2010, 24.6% (13 million) were due to communicable diseases, maternal and perinatal conditions, and nutritional deficiencies—a marked decrease compared with figures for 1990, when these conditions accounted for 34% of global mortality. Among the fraction of all deaths related to communicable diseases, maternal and perinatal conditions, and nutritional deficiencies, 76% occurred in sub-Saharan Africa and southern Asia. While the proportion of deaths due to these conditions has decreased significantly in the past decade, there has been a dramatic rise in the number of deaths from NCDs, which constituted the top five causes of death in 2010. The leading cause of death among adults in 2010 was ischemic heart disease, accounting for 7.3 million deaths (13.8% of total deaths) worldwide. In high-income countries ischemic heart disease accounted for 17.9% of total deaths, and in developing (low- and middle-income) countries it accounted for 10.1%. It is noteworthy that ischemic heart disease was responsible for just 2.6% of total deaths in sub-Saharan Africa (Table 2-2). In second place—causing 11.1% of global mortality—was cerebrovascular disease, which accounted for 9.9% of deaths in high-income countries, 10.5% in developing countries, and 4.0% in sub-Saharan Africa. Although the third leading cause of death in high-income countries was lung cancer (accounting for 5.6% of all deaths), this condition did not figure among the top 10 causes in low- and middle-income countries. Among the 10 leading causes of death in sub-Saharan Africa, 6 were infectious diseases, with malaria and HIV/AIDS ranking as the dominant contributors to disease burden. In high-income countries, however, only one infectious disease—lower respiratory infection—ranked among the top 10 causes of death.

The GBD 2010 found that the worldwide mortality figure among children <5 years of age dropped from 16.39 million in 1970 to 11.9 million in 1990 and to 6.8 million in 2010—a decrease that surpassed predictions. Of childhood deaths in 2010, 3.1 million (40%) occurred in the neonatal period. About one-third of deaths among children <5 years old occurred in southern Asia and almost one-half in sub-Saharan Africa; <1% occurred in high-income countries.

The global burden of death due to HIV/AIDS and malaria was on an upward slope until 2004; significant improvements have since been documented. Global deaths from HIV infection fell from 1.7 million

in 2006 to 1.5 million in 2010, while malaria deaths dropped from 1.2 million to 0.98 million over the same period. Despite these improvements, malaria and HIV/AIDS continue to be major burdens in particular regions, with global implications. Although it has a minor impact on mortality outside sub-Saharan Africa and Southeast Asia, malaria is the eleventh leading cause of death worldwide. HIV infection ranked thirty-third in global DALYs in 1990 but was the fifth leading cause of disease burden in 2010, with sub-Saharan Africa bearing the vast majority of this burden (Fig. 2-1).

The world's population is living longer: global life expectancy has increased significantly over the past 40 years from 58.8 years in 1970 to 70.4 years in 2010. This demographic change, accompanied by the fact that the prevalence of NCDs increases with age, is dramatically shifting the burden of disease toward NCDs, which have surpassed communicable, maternal, nutritional, and neonatal causes. By 2010, 65.5% of total deaths at all ages and 54% of all DALYs were due to NCDs. Increasingly, the global burden of disease comprises conditions and injuries that cause disability rather than death.

Worldwide, although both life expectancy and years of life lived in good health have risen, years of life lived with disability have also increased. Despite the higher prevalence of diseases common in older populations (e.g., dementia and musculoskeletal disease) in developed and high-income countries, best estimates from 2010 reveal that disability resulting from cardiovascular diseases, chronic respiratory diseases, and the long-term impact of communicable diseases was greater in low- and middle-income countries. In most developing countries, people lived shorter lives and experienced disability and poor health for a greater proportion of their lives. Indeed, 50% of the global burden of disease occurred in southern Asia and sub-Saharan Africa, which together account for only 35% of the world's population.

HEALTH AND WEALTH

Clear disparities in burden of disease (both communicable and noncommunicable) across country income levels are strong indicators that poverty and health are inherently linked. Poverty remains one of the most important root causes of poor health worldwide, and the global burden of poverty continues to be high. Among the 6.7 billion people alive in 2008, 19% (1.29 billion) lived on less than \$1.25 a day—one standard measurement of extreme poverty—and another 1.18 billion lived on \$1.25 to \$2 a day. Approximately 600 million children—more than 30% of those in low-income countries—lived in extreme poverty in 2005. Comparison of national health indicators with gross domestic product per capita among nations shows a clear relationship between higher gross domestic product and better health, with only a few outliers. Numerous studies have also documented the link between poverty and health within nations as well as across them.

RISK FACTORS FOR DISEASE BURDEN

The GBD 2010 study found that the three leading risk factors for global disease burden in 2010 were (in order of frequency) high blood pressure, tobacco smoking (including secondhand smoke), and alcohol use—a substantial change from 1990, when childhood undernutrition was ranked first. Though ranking eighth in 2010, childhood undernutrition remains the leading risk factor for death worldwide among children <5 years of age. In an era that has seen obesity become a major health concern in many developed countries—and the sixth leading risk factor worldwide—the persistence of undernutrition is surely cause for great consternation. Low body weight is still the dominant risk factor for disease burden in sub-Saharan Africa. Inability to feed the hungry reflects many years of failed development projects and must be addressed as a problem of the highest priority. Indeed, no health care initiative, however generously funded, will be effective without adequate nutrition.

In a 2006 publication that examined how specific diseases and injuries are affected by environmental risk, the WHO estimated that roughly one-quarter of the total global burden of disease, one-third of the global disease burden among children, and 23% of all deaths were due to modifiable environmental factors. Many of these factors lead to deaths from infectious diseases; others lead to deaths

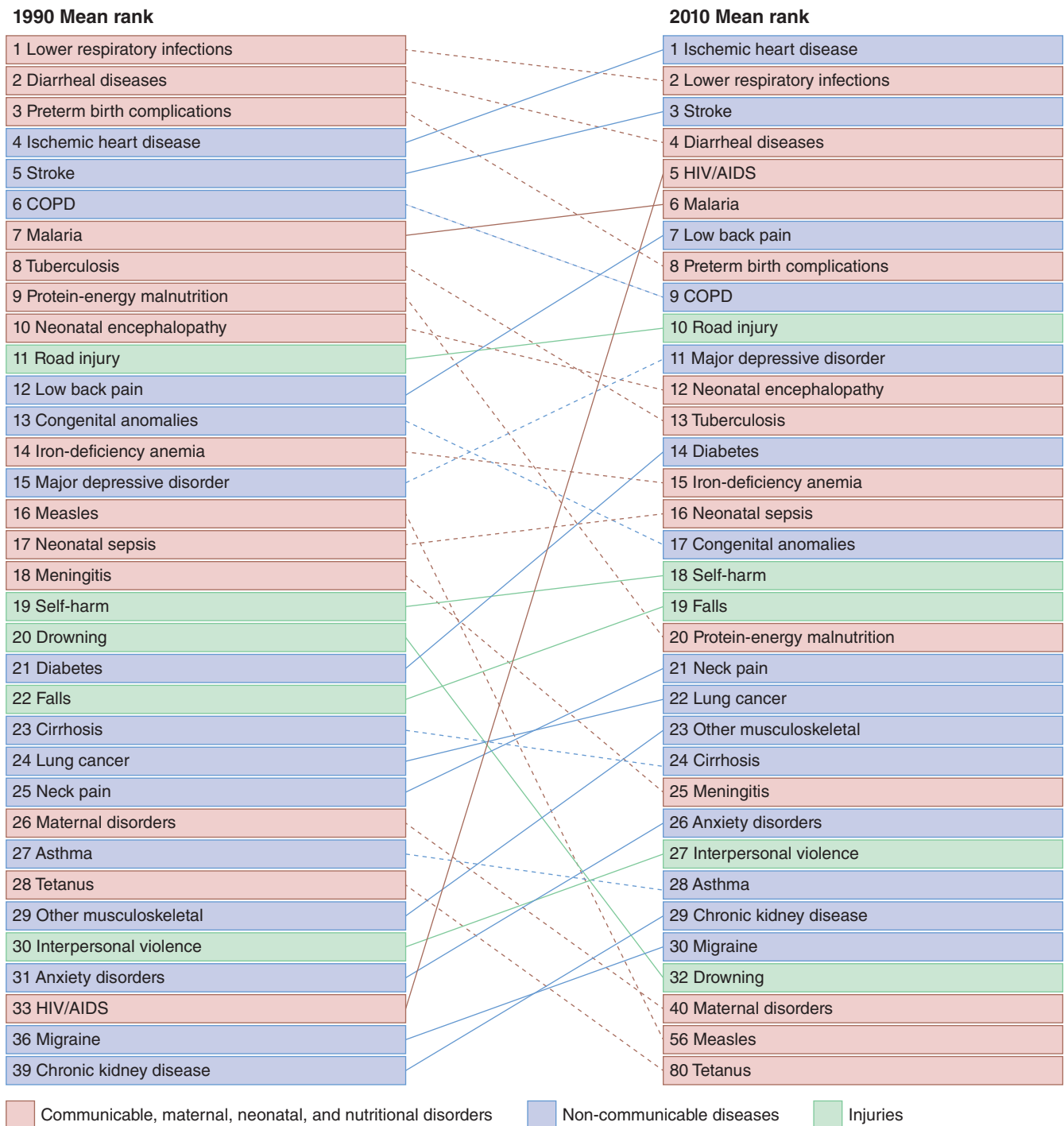


FIGURE 2-1 Global DALY (disability-adjusted life year) ranks for the top causes of disease burden in 1990 and 2010. COPD, chronic obstructive pulmonary disease. (Reproduced with permission from C Murray et al: *Disability-adjusted life years [DALYs] for 291 diseases and injuries in 21 regions, 1990–2010: A systematic analysis for the Global Burden of Disease Study 2010*. *Lancet* 380:2197–2223, 2012.)

from malignancies. Etiology and nosology are increasingly difficult to parse. As much as 94% of diarrheal disease, which is linked to unsafe drinking water and poor sanitation, can be attributed to environmental factors. Risk factors such as indoor air pollution due to use of solid fuels, exposure to secondhand tobacco smoke, and outdoor air pollution account for 20% of lower respiratory infections in developed countries and for as many as 42% of such infections in developing countries. Various forms of unintentional injury and malaria top the list of health problems to which environmental factors contribute. Some 4 million children die every year from causes related to unhealthy environments, and the number of infant deaths due to environmental factors in developing countries is 12 times that in developed countries.

The second edition of *Disease Control Priorities in Developing Countries*, published in 2006, is a document of great breadth and ambition, providing cost-effectiveness analyses for more than 100 interventions and including 21 chapters focused on strategies for strengthening health systems. Cost-effectiveness analyses that compare relatively equivalent interventions and facilitate the best choices under constraint are necessary; however, these analyses are often based on an incomplete knowledge of cost and evolving evidence of effectiveness. As both resources and objectives for global health grow, cost-effectiveness analyses (particularly those based on older evidence) must not hobble the increased worldwide commitment to providing resources and accessible health care services to all who need them. This is why we use the term *global health equity*. To illustrate these points, it

TABLE 2-1 LEADING CAUSES OF DISEASE BURDEN, 2010

Disease or Injury	DALYs (Millions)	Percent of Total DALYs	Disease or Injury	DALYs (Millions)	Percent of Total DALYs
World			High-income countries^b		
1 Ischemic heart disease	129.8	5.2	1 Ischemic heart disease	21.8	8.2
2 Lower respiratory infections	115.2	4.7	2 Low back pain	17.0	6.4
3 Cerebrovascular disease	102.2	4.1	3 Cerebrovascular disease	11.3	4.2
4 Diarrheal disease	89.5	3.6	4 Major depressive disorder	9.7	3.7
5 HIV/AIDS	81.5	3.3	5 Lung cancer	9.2	3.5
6 Malaria	82.7	3.3	6 COPD	8.6	3.2
7 Low back pain	80.7	3.2	7 Other musculoskeletal disorders	8.2	3.1
8 Preterm birth complications	77.0	3.1	8 Diabetes mellitus	7.3	2.8
9 COPD	76.8	3.1	9 Neck pain	7.2	2.7
10 Road injury	75.5	3.1	10 Falls	6.8	2.5
Developing countries^a			Sub-Saharan Africa		
1 Lower respiratory infections	109.0	5.2	1 Malaria	76.6	13.3
2 Diarrheal disease	88.0	4.2	2 HIV/AIDS	57.8	10.1
3 Ischemic heart disease	85.5	4.1	3 Lower respiratory infections	43.5	7.6
4 Malaria	82.7	3.9	4 Diarrheal diseases	39.2	6.8
5 Cerebrovascular disease	79.4	3.8	5 Protein-energy malnutrition	22.3	3.9
6 HIV/AIDS	77.0	3.7	6 Preterm birth complications	20.0	3.5
7 Preterm birth complications	74.4	3.5	7 Neonatal sepsis	18.9	3.3
8 Road injury	66.2	3.2	8 Meningitis	16.3	2.8
9 COPD	65.6	3.1	9 Neonatal encephalopathy	14.9	2.6
10 Low back pain	58.4	2.8	10 Road injury	13.9	2.5

^aThe term *developing countries* refers to low- and middle-income economies. See data.worldbank.org/about/country-classifications. ^bThe World Bank classifies high-income countries as those whose gross national income per capita is \$12,476 or more. See data.worldbank.org/about/country-classifications.

Abbreviations: COPD, chronic obstructive pulmonary disease; DALYs, disability-adjusted life years.

Source: Institute for Health Metrics and Evaluation, University of Washington (2013). Data are available through www.healthmetricsandevaluation.org/gbd/visualizations/country.

TABLE 2-2 LEADING CAUSES OF DEATH WORLDWIDE, 2010

Disease or Injury	Deaths (Millions)	Percent of Total Deaths	Disease or Injury	Deaths (Millions)	Percent of Total Deaths
World			High-income countries^b		
1 Ischemic heart disease	7.3	13.3	1 Ischemic heart disease	1.6	17.9
2 Cerebrovascular disease	5.9	11.1	2 Cerebrovascular disease	0.9	9.9
3 COPD	2.9	5.5	3 Lung cancer	0.5	5.6
4 Lower respiratory infections	2.8	5.3	4 Lower respiratory infections	0.4	4.7
5 Lung cancer	1.5	2.9	5 COPD	0.4	4.5
6 HIV/AIDS	1.5	2.8	6 Alzheimer's and other dementias	0.4	4.0
7 Diarrheal diseases	1.4	2.7	7 Colon and rectum cancers	0.3	3.3
8 Road injury	1.3	2.5	8 Diabetes	0.2	2.6
9 Diabetes	1.3	2.4	9 Other cardiovascular and circulatory diseases	0.2	2.5
10 Tuberculosis	1.2	2.3	10 Chronic kidney disease	0.2	2.0
Developing countries^a			Sub-Saharan Africa		
1 Cerebrovascular disease	4.2	10.5	1 Malaria	1.1	12.7
2 Ischemic heart disease	4.0	10.1	2 HIV/AIDS	1.0	12.0
3 COPD	2.4	6.1	3 Lower respiratory infections	0.8	9.3
4 Lower respiratory infections	2.3	5.9	4 Diarrheal diseases	0.5	6.6
5 Diarrheal diseases	1.4	3.6	5 Cerebrovascular disease	0.3	4.0
6 HIV/AIDS	1.4	3.4	6 Protein-energy malnutrition	0.3	4.0
7 Malaria	1.2	2.9	7 Tuberculosis	0.3	3.6
8 Road injury	1.2	2.9	8 Road injury	0.2	2.8
9 Tuberculosis	1.1	2.9	9 Preterm birth complications	0.2	2.8
10 Diabetes	1.0	2.6	10 Meningitis	0.2	2.6

^aThe term *developing countries* refers to low- and middle-income economies. See data.worldbank.org/about/country-classifications. ^bThe World Bank classifies high-income countries as those whose gross national income per capita is \$12,476 or more. See data.worldbank.org/about/country-classifications.

Abbreviation: COPD, chronic obstructive pulmonary disease.

Source: Institute for Health Metrics and Evaluation, University of Washington (2013). Data available through www.healthmetricsandevaluation.org/gbd/visualizations/country.

is instructive to look to HIV/AIDS, which in the course of the last three decades has become the world's leading infectious cause of adult death.

HIV INFECTION/AIDS

Chapter 226 provides an overview of the HIV epidemic in the world today. Here the discussion will be limited to HIV/AIDS in the developing world. Lessons learned from tackling HIV/AIDS in resource-constrained settings are highly relevant to discussions of other chronic diseases, including NCDs, for which effective therapies have been developed.

Approximately 34 million people in all countries worldwide were living with HIV infection in 2011; more than 8 million of those in low- and middle-income countries were receiving antiretroviral therapy (ART)—a number representing a 20-fold increase over the corresponding figure for 2003. By the end of 2011, 54% of people eligible for treatment were receiving ART. (It remains to be seen how many of these people are receiving ART regularly and with the requisite social support.)

In the United States, the availability of ART has transformed HIV/AIDS from an inescapably fatal destruction of cell-mediated immunity into a manageable chronic illness. In high-income countries, improved ART has prolonged life by an estimated average of 35 years per patient—up from 6.8 years in 1993 and 24 years in 2006. This success rate exceeds that obtained with almost any treatment for adulthood cancer or for complications of coronary artery disease. In developing countries, treatment has been offered broadly only since 2003, and only in 2009 did the number of patients receiving treatment exceed 40% of the number who needed it. Before 2003, many arguments were raised to justify not moving forward rapidly with ART programs for people living with HIV/AIDS in resource-limited settings. The standard litany included the price of therapy compared with the poverty of the patient, the complexity of the intervention, the lack of infrastructure for laboratory monitoring, and the lack of trained health care providers. Narrow cost-effectiveness arguments that created false dichotomies—prevention *or* treatment rather than both—too often went unchallenged. As a cumulative result of these delays in the face of health disparities, old and new, there were millions of premature deaths.

Disparities in access to HIV treatment gave rise to widespread moral indignation and a new type of health activism. In several middle-income countries, including Brazil, public programs have helped bridge the access gap. Other innovative projects pioneered by international NGOs in diverse settings such as Haiti and Rwanda have established that a simple approach to ART that is based on intensive community engagement and support can achieve remarkable results (**Fig. 2-2**).

During the past decade, the availability of ART has increased sharply in the low- and middle-income countries that have borne the greatest burden of the HIV/AIDS pandemic. In 2000, very few people living with HIV/AIDS in these nations had access to ART, whereas by 2011, as stated above, 8 million people, a majority of those deemed eligible, in these countries were receiving ART. This scale-up was made possible by a number of developments: a staggering drop in the cost of ART, the development of a standardized approach to treatment, substantial investments by funders, and the political commitment of governments to make ART available. Civil-society AIDS activists spurred many of these efforts.

Starting in the early 2000s, a combination of factors, including work by the Clinton Foundation HIV/AIDS Initiative and Médecins Sans Frontières, led to the availability of generic ART medications. While first-line ART cost more than \$10,000 per patient per year in 2000, first-line regimens in low- and middle-income countries are now available for less than \$100 per year. At the same time, fixed-dose combination drugs that are easier to administer have become more widely available.

Also around this time, the WHO began advocating a public health approach to the treatment of people with AIDS in resource-limited settings. This approach, derived from models of care pioneered by the NGO Partners In Health and other groups, proposed standard first-line treatment regimens based on a simple five-drug formulation, with a more complex (and more expensive) set of second-line options in reserve. Clinical protocols were standardized, and intensive training packages for health professionals and community health workers were developed and implemented in many countries. These efforts were supported by new funding from the World Bank, the Global Fund, and PEPFAR. In 2003, lack of access to ART was declared a global public health emergency by the WHO and UNAIDS, and those two agencies launched the “3 by 5 initiative,” setting an ambitious target: to have 3 million people in developing countries on treatment by the end of 2005. Worldwide funding for HIV/AIDS treatment increased dramatically during this period, rising from \$300 million in 1996 to over \$15 billion in 2010.

Many countries set corresponding national targets and have worked to integrate ART into their national AIDS programs and health systems and to harness the synergies between HIV/AIDS treatment and prevention activities. Further lessons with implications for policy and action have come from efforts now under way among lower-income countries. Rwanda provides an example: Over the past decade, mortality from HIV disease has fallen by >78% as the country—despite its relatively low gross national income (**Fig. 2-3**)—has provided almost universal access to ART. The reasons for this success include strong national leadership, evidence-based policy, cross-sector collaboration, community-based care, and a deliberate focus on a health system approach that embeds HIV/AIDS treatment and prevention in the primary health care service delivery platform. As we will discuss later in this chapter, these principles can be applied to other conditions, including NCDs.

TUBERCULOSIS

Chapter 202 provides a concise overview of the pathophysiology and treatment of TB. In 2011, an estimated 12 million people were living with active TB, and 1.4 million died from it. The disease is closely linked to HIV infection in much of the world: of the 8.7 million estimated new cases of TB in 2011, 1.2 million occurred among people living with HIV. Indeed, a substantial proportion of the TB resurgence



FIGURE 2-2 An HIV/TB-co-infected patient in Rwanda before (*left*) and after (*right*) 6 months of treatment.

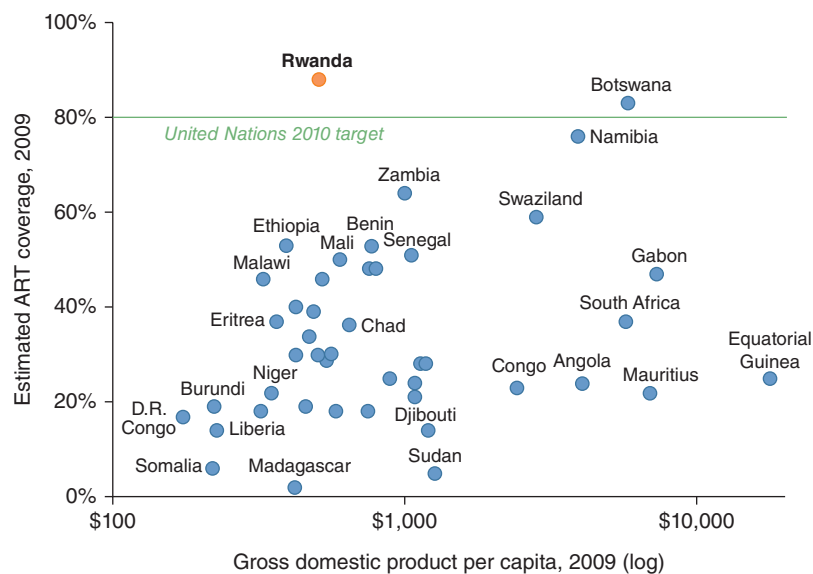


FIGURE 2-3 Antiretroviral therapy (ART) coverage in sub-Saharan Africa, 2009.

registered in southern Africa is attributed to HIV co-infection. Even before the advent of HIV, however, it was estimated that fewer than one-half of all cases of TB in developing countries were ever diagnosed, much less treated. Primarily because of the common failure to diagnose and treat TB, international authorities devised a single strategy to reduce the burden of disease. In the early 1990s, the World Bank, the WHO, and other international bodies promoted the DOTS strategy (directly observed therapy using short-course isoniazid- and rifampin-based regimens) as highly cost-effective. Passive case-finding of smear-positive patients was central to the strategy, and an uninterrupted drug supply was, of course, deemed necessary for cure.

DOTS was clearly effective for most uncomplicated cases of drug-susceptible TB, but a number of shortcomings were soon identified. First, the diagnosis of TB based solely on sputum smear microscopy—a method dating from the late nineteenth century—is not sensitive. Many cases of pulmonary TB and all cases of exclusively extrapulmonary TB are missed by smear microscopy, as are most cases of active disease in children. Second, passive case-finding relies on the availability of health care services, which is uneven in the settings where TB is most prevalent. Third, patients with multidrug-resistant TB (MDR-TB) are by definition infected with strains of *Mycobacterium tuberculosis* resistant to isoniazid and rifampin; thus exclusive reliance on these drugs is unwarranted in settings in which drug resistance is an established problem.

The crisis of antibiotic resistance registered in U.S. hospitals is not confined to the industrialized world or to common bacterial infections. The great majority of patients sick with and dying from TB are afflicted with strains susceptible to all first-line drugs. In some settings, however, a substantial minority of patients with TB are infected with *M. tuberculosis* strains resistant to at least one first-line anti-TB drug. A 2012 article in a leading journal reported that, in China, 10% of all patients with TB and 26% of all previously treated patients were sick with MDR strains of *M. tuberculosis*. Most of these cases were the result of primary transmission. To improve DOTS-based responses to MDR-TB, global health authorities adopted DOTS-Plus, which adds the diagnostics and drugs necessary to manage drug-resistant disease. Even as DOTS-Plus was being piloted in resource-constrained settings, however, new strains of extensively drug-resistant (XDR) *M. tuberculosis* (resistant to isoniazid and rifampin, any fluoroquinolone, and at least one injectable second-line drug) had already threatened the success of TB control programs in beleaguered South Africa, for example, where high rates of HIV infection have led to a doubling of TB incidence over the last decade. Despite the poor capacity for detection of MDR- and XDR-TB in most resource-limited settings, an estimated 630,000 cases of MDR-TB were thought to occur in 2011. Approximately 9% of these drug-resistant cases were caused by XDR strains. It is clear that poor

infection control in hospitals and clinics is associated with explosive and lethal epidemics due to these strains and that patients may be infected with multiple strains.

TUBERCULOSIS AND AIDS AS CHRONIC DISEASES: LESSONS LEARNED

Strategies effective against MDR-TB have implications for the management of drug-resistant HIV infection and even drug-resistant malaria, which, through repeated infections and a lack of effective therapy, has become a chronic disease in parts of Africa (see “Malaria,” below). As new therapies, whether for TB or for hepatitis C infection, become available, many of the problems encountered in the past will recur. Indeed, examining AIDS and TB as chronic diseases—instead of simply communicable diseases—makes it possible to draw a number of conclusions, many of them pertinent to global health in general.

First, the chronic infections discussed here are best treated with multidrug regimens to which the infecting strains are susceptible. This is true of chronic infections due to many bacteria, fungi, parasites, or viruses; even acute infections such as those caused by *Plasmodium* species are not reliably treated with a single drug.

Second, charging fees for AIDS prevention and care poses insurmountable problems for people living in poverty, many of whom are unable to pay even modest amounts for services or medications. Like efforts to battle airborne TB, such services might best be seen as a *public good* promoting public health. Initially, a subsidy approach will require sustained donor contributions, but many African countries have set targets for increased national investments in health—a pledge that could render ambitious programs sustainable in the long run, as the Rwanda experience suggests. Meanwhile, as local investments increase, the price of AIDS care is decreasing. The development of generic medications means that ART can now cost <\$0.25 per day; costs continue to decrease.

Third, the effective scale-up of pilot projects requires strengthening and sometimes rebuilding of health care systems, including those charged with delivering primary care. In the past, the lack of health care infrastructure has been cited as a barrier to providing ART in the world’s poorest regions; however, AIDS resources, which are at least considerable, may be marshaled to rebuild public health systems in sub-Saharan Africa and other HIV-burdened regions—precisely the settings in which TB is resurgent.

Fourth, the lack of trained health care personnel, most notably doctors and nurses, in resource-poor settings must be addressed. This personnel deficiency is invoked as a reason for the failure to treat AIDS in poor countries. In what is termed the *brain drain*, many physicians and nurses emigrate from their home countries to pursue opportunities abroad, leaving behind health systems that are understaffed and ill equipped to deal with the epidemic diseases that ravage local populations. The WHO recommends a minimum of 20 physicians and 100 nurses per 100,000 persons, but recent reports from that organization and others confirm that many countries, especially in sub-Saharan Africa, fall far short of those target numbers. Specifically, more than one-half of those countries register fewer than 10 physicians per 100,000 population. In contrast, the United States and Cuba register 279 and 596 doctors per 100,000 population, respectively. Similarly, the majority of sub-Saharan African countries do not have even half of the WHO-recommended minimum number of nurses. Further inequalities in health care staffing exist *within* countries. Rural–urban disparities in health care personnel mirror disparities of both wealth and health. For instance, nearly 90% of Malawi’s population lives in rural areas, but more than 95% of clinical officers work at urban facilities, and 47% of nurses work at tertiary care facilities. Even community health workers trained to provide first-line services to rural populations often transfer to urban districts.

One reason doctors and nurses leave sub-Saharan Africa and other resource-poor areas is that they lack the tools to practice their trade there. Funding for “vertical” (disease-specific) programs can be used

not only to strengthen health systems but to recruit and train physicians and nurses to underserved regions where they, in turn, can help to train and then work with community health workers in supervising care for patients with AIDS and many other diseases within their communities. Such training should be undertaken even where physicians are abundant, since close community-based supervision represents the highest standard of care for chronic disease, whether in developing or developed countries. The United States has much to learn from Rwanda.

Fifth, the barriers to adequate health care and patient adherence that are raised by extreme poverty can be removed only with the deployment of “wrap-around services”: food supplements for the hungry, help with transportation to clinics, child care, and housing. Extreme poverty makes it difficult for many patients to comply with therapy for chronic diseases, whether communicable or not. Indeed, poverty in its many dimensions is far and away the greatest barrier to the scale-up of treatment and prevention programs. In many rural regions of Africa, hunger is the major coexisting condition in patients with AIDS or TB, and those consumptive diseases cannot be treated effectively without adequate caloric intake.

Finally, there is a need for a renewed basic-science commitment to the discovery and development of vaccines; more reliable, less expensive diagnostic tools; and new classes of therapeutic agents. This need applies not only to the three leading infectious killers—against none of which is there an effective vaccine—but also to most other neglected diseases of poverty.

MALARIA

Chapter 248 reviews the etiology, pathogenesis, and clinical treatment of malaria, the world’s third-ranking infectious killer. Malaria’s human cost is enormous, with the highest toll among children—especially African children—living in poverty. In 2010, there were ~219 million cases of malaria, and the disease is thought to have killed 660,000 people; 86% of these deaths (~568,000) occurred among children <5 years old. The poor disproportionately experience the burden of malaria: more than 80% of estimated malaria deaths occur in just 14 countries, and mortality rates are highest in sub-Saharan Africa. The Democratic Republic of the Congo and Nigeria account for more than 40% of total estimated malaria deaths globally.

Microeconomic analyses focusing on direct and indirect costs estimate that malaria may consume >10% of a household’s annual income. A study in rural Kenya shows that mean direct-cost burdens vary between the wet and dry seasons (7.1% and 5.9% of total household expenditure, respectively) and that this proportion is >10% in the poorest households in both seasons. A Ghanaian study that categorized the population by income group highlighted the regressive nature of this cost: responding to malaria consumed only 1% of a wealthy family’s income but 34% of a poor household’s income.

Macroeconomic analyses estimate that malaria may reduce the per capita gross national product of a disease-endemic country by 50% relative to that of a non-malaria-endemic country. The causes of this drag include impaired cognitive development of children, decreased schooling, decreased savings, decreased foreign investment, and restriction of worker mobility. In light of this enormous cost, it is little wonder that an important review by the economists Sachs and Malaney concludes that “where malaria prospers most, human societies have prospered least.”

Rolling Back Malaria In part because of differences in vector distribution and climate, resource-rich countries offer few blueprints for malaria control and treatment that are applicable in tropical (and resource-poor) settings. In 2001, African heads of state endorsed the WHO Roll Back Malaria (RBM) campaign, which prescribes strategies appropriate for sub-Saharan African countries. In 2008, the RBM partnership launched the Global Malaria Action Plan (GMAP). This strategy *integrates prevention and care* and calls for an avoidance of single-dose regimens and an awareness of existing drug resistance. The GMAP recommends a number of key tools to reduce malaria-related morbidity and mortality rates: the use of insecticide-treated bed nets (ITNs), indoor residual spraying, and artemisinin-based combination

therapy (ACT) as well as intermittent preventive treatment during pregnancy, prompt diagnosis, and other vector control measures such as larviciding and environmental management.

INSECTICIDE-TREATED BED NETS ITNs are an efficacious and cost-effective public health intervention. A meta-analysis of controlled trials in seven sub-Saharan African countries indicates that parasitemia prevalence is reduced by 24% among children <5 years of age who sleep under ITNs compared with that among those who do not. Even untreated nets reduce malaria incidence by one-quarter. On an individual level, the utility of ITNs extends beyond protection from malaria. Several studies suggest that ITNs reduce all-cause mortality among children under age 5 to a greater degree than can be attributed to the reduction in malarial disease alone. Morbidity (specifically that due to anemia), which predisposes children to diarrheal and respiratory illnesses and pregnant women to the delivery of low-birth-weight infants, also is reduced in populations using ITNs. In some areas, ITNs offer a supplemental benefit by preventing transmission of lymphatic filariasis, cutaneous leishmaniasis, Chagas’ disease, and tick-borne relapsing fever. At the community level, investigators suggest that the use of an ITN in just one household may reduce the number of mosquito bites in households up to a hundred meters away by reducing mosquito density. The cost of ITNs per DALY saved—estimated at \$29—makes ITNs a good-value public health investment.

The WHO recommends that all individuals living in malaria-endemic areas sleep under protective ITNs. About 140 million long-lasting ITNs were distributed in high-burden African countries in 2006–2008, and rates of household ownership of ITNs in high-burden countries increased to 31%. Although the RBM partnership has seen modest success, the WHO’s 2009 World Malaria Report states that the percentage of children <5 years of age using an ITN (24%) remains well below the World Health Assembly’s target of 80%. Limited success in scaling up ITN coverage reflects the inadequately acknowledged economic barriers that prevent the destitute sick from gaining access to critical preventive technologies and the challenges faced in designing and implementing effective delivery platforms for these products. In other words, this is a *delivery* failure rather than a lack of knowledge of how best to reduce malaria deaths.

INDOOR RESIDUAL SPRAYING Indoor residual spraying is one of the most common interventions for preventing the transmission of malaria in endemic areas. Vector control using insecticides approved by the WHO, including DDT, can effectively reduce or even interrupt malaria transmission. However, studies have indicated that spraying is effective in controlling malaria transmission only if most (~80%) of the structures in the targeted community are treated. Moreover, since a successful program depends on well-trained spraying teams as well as on effective monitoring and planning, indoor residual spraying is difficult to employ and is often reliant on health systems with a strong infrastructure. Regardless of the limitations of indoor residual spraying, the WHO recommends its use in combination with ITNs. Neither intervention alone is sufficient to prevent transmission of malaria entirely.

ARTEMISININ-BASED COMBINATION THERAPY The emergence and spread of chloroquine resistance have increased the need for antimalarial combination therapy. To limit the spread of resistance, the WHO now recommends that only ACT (as opposed to artemisinin monotherapy) be used for uncomplicated falciparum malaria. Like that of other antimalarial interventions, the use of ACT has increased in the last few years, but coverage rates remain very low in several countries in sub-Saharan Africa. The RBM partnership has invested significantly in measures to enhance access to ACT by facilitating its delivery through the public health sector and developing innovative funding mechanisms (e.g., the Affordable Medicines Facility—malaria) that reduce its cost significantly so that ineffective monotherapies can be eliminated from the market.

In the last several years, resistance to antimalarial medicines and insecticides has become an even larger problem than in the past. In 2009, confirmation of artemisinin resistance was reported. Although the WHO has called for an end to the use of artemisinin monotherapy, the

marketing of such therapies continues in many countries. Ongoing use of artemisinin monotherapy increases the likelihood of drug resistance, a deadly prospect that will make malaria far more difficult to treat.

Between 2001 and 2011, global malaria deaths were reduced by an estimated 38%, with reductions of $\geq 50\%$ in 10 African countries as well as in most endemic countries in other regions. Again the experience in Rwanda is instructive: from 2005 to 2011, malaria deaths dropped by $>85\%$ for the same reasons mentioned earlier in recounting that nation's successes in battling HIV.

Meeting the challenge of malaria control will continue to require careful study of appropriate preventive and therapeutic strategies in the context of an increasingly sophisticated molecular understanding of pathogen, vector, and host. However, an appreciation of the economic and social devastation wrought by malaria—like that inflicted by diarrhea, AIDS, and TB—on the most vulnerable populations should heighten the level of commitment to critical analysis of ways to implement proven strategies for prevention and treatment.

Funding from the Global Fund, the Gates Foundation, the World Bank's International Development Association, and the U.S. President's Malaria Initiative, along with leadership from public health authorities, is critical to sustain the benefits of prevention and treatment. Building on the growing momentum of the last decade with adequate financial support, innovative strategies, and effective tools for prevention, diagnosis, and treatment, we may one day achieve the goal of a world free of malaria.

"NONCOMMUNICABLE" CHRONIC DISEASES

Although the burden of communicable diseases—especially HIV infection, TB, and malaria—still accounts for the majority of deaths in resource-poor regions such as sub-Saharan Africa, 63% of all deaths worldwide in 2008 were held to be due to NCDs. Although we will use this term to describe cardiovascular diseases, cancers, diabetes, and chronic lung diseases, this usage masks important distinctions. For instance, two significant NCDs in low-income countries, rheumatic heart disease (RHD) and cervical cancer, represent the chronic sequelae of infections with group A *Streptococcus* and human papillomavirus, respectively. It is in these countries that the burden of disease due to NCDs is rising most rapidly. Close to 80% of deaths attributable to NCDs occur in low- and middle-income countries, where 86% of the global population lives. The WHO reports that $\sim 25\%$ of global NCD-related deaths take place before the age of 60—a figure representing ~ 5.7 million people and exceeding the total number of deaths due to AIDS, TB, and malaria combined. In almost all high-income countries, the WHO reported that NCD deaths accounted for $\sim 70\%$ of total deaths in 2008. By 2020, NCDs will account for 80% of the global burden of disease and for 7 of every 10 deaths in developing countries. The recent increase in resources for and attention to communicable diseases is both welcome and long overdue, but developing countries are already carrying a “double burden” of communicable and noncommunicable diseases.

Diabetes, Cardiovascular Disease, and Cancer: A Global Perspective In contrast to TB, HIV infection, and malaria—diseases caused by single pathogens that damage multiple organs—cardiovascular diseases reflect injury to a single organ system downstream of a variety of insults, both infectious and noninfectious. Some of these insults result from rapid changes in diet and labor conditions.

Other insults are of a less recent vintage. The burden of cardiovascular disease in low-income countries represents one consequence of decades of neglect of health systems. Furthermore, cardiovascular research and investment have long focused on the ischemic conditions that are increasingly common in high- and middle-income countries. Meanwhile, despite awareness of its health impact in the early twentieth century, cardiovascular damage in response to infection and malnutrition has fallen out of view until recently.

The misperception of cardiovascular diseases as a problem primarily of elderly populations in middle- and high-income countries has contributed to the neglect of these diseases by global health institutions. Even in Eastern Europe and Central Asia, where the collapse of the Soviet Union was followed by a catastrophic surge in cardiovascular

disease deaths (mortality rates from ischemic heart disease nearly doubled between 1991 and 1994 in Russia, for example), the modest flow of overseas development assistance to the health sector focused on the communicable causes that accounted for <1 in 20 excess deaths during that period.

DIABETES The International Diabetes Federation reports that the number of diabetic patients in the world is expected to increase from 366 million in 2011 to 552 million by 2030. Already, a significant proportion of diabetic patients live in developing countries where, because those affected are far more frequently between ages 40 and 59, the complications of micro- and macrovascular disease take a far greater toll. Globally, these complications are a major cause of disability and reduced quality of life. A high fasting plasma glucose level alone ranks seventh among risks for disability and is the sixth leading risk factor for global mortality. The GBD 2010 estimates that diabetes accounted for 1.28 million deaths in 2010, with almost 80% of those deaths occurring in low- and middle-income countries.

Predictions of an imminent rise in the share of deaths and disabilities due to NCDs in developing countries have led to calls for preventive policies to improve diet, increase exercise, and restrict tobacco use, along with the prescription of multidrug regimens for persons at high-level vascular risk. Although this agenda could do much to prevent pandemic NCD, it will do little to help persons with established heart disease stemming from nonatherogenic pathologies.

CARDIOVASCULAR DISEASE Because systemic investigation of the causes of stroke and heart failure in sub-Saharan Africa has begun only recently, little is known about the impact of elevated blood pressure in this portion of the continent. Modestly elevated blood pressure in the absence of tobacco use in populations with low rates of obesity may confer little risk of adverse events in the short run. In contrast, persistently elevated blood pressure above 180/110 goes largely undetected, untreated, and uncontrolled in this part of the world. In the cohort of men assessed in the Framingham Heart Study, the prevalence of blood pressures above 210/120—severe hypertension—declined from 1.8% in the 1950s to 0.1% by the 1960s with the introduction of effective antihypertensive agents. Although debate continues about appropriate screening strategies and treatment thresholds, rural health centers staffed largely by nurses must quickly gain access to essential antihypertensive medications.

The epidemiology of heart failure reflects inequalities in risk factor prevalence and in treatment. The reported burden of this condition has remained unchanged since the 1950s, but the causes of heart failure and the age of the people affected vary across the globe. Heart failure as a consequence of pericardial, myocardial, endocardial, or valvular injury accounts for as many as 5% of all medical admissions to hospitals around the world. In high-income countries, coronary artery disease and hypertension among the elderly account for most cases of heart failure. For example, in the United States, coronary artery disease is present in 60% of patients with heart failure and hypertension in 70%. Among the world's poorest 1 billion people, however, heart failure reflects poverty-driven exposure of children and young adults to rheumatogenic strains of streptococci and cardiotropic microorganisms (e.g., HIV, *Trypanosoma cruzi*, enteroviruses, *M. tuberculosis*), untreated high blood pressure, and nutrient deficiencies. The mechanisms underlying other causes of heart failure common in these populations—such as idiopathic dilated cardiomyopathy, peripartum cardiomyopathy, and endomyocardial fibrosis—remain unclear.

In stark contrast to the extraordinary lengths to which clinicians in wealthy countries will go to treat ischemic cardiomyopathy, little attention has been paid to young patients with nonischemic cardiomyopathies in resource-poor settings. Nonischemic cardiomyopathies, such as those due to hypertension, RHD, and chronic lung disease, account for $>90\%$ of cases of cardiac failure in sub-Saharan Africa and include poorly understood entities such as peripartum cardiomyopathy (which has an incidence in rural Haiti of 1 per 300 live births) and HIV-associated cardiomyopathy. Multidrug regimens that include beta blockers, angiotensin-converting enzyme inhibitors, and other agents can dramatically reduce mortality risk and improve quality of life for these patients. Lessons learned in the scale-up of chronic care

for HIV infection and TB may be illustrative as progress is made in establishing the means to deliver heart-failure therapies.

Some of the lessons learned from the chronic infections discussed above are, of course, relevant to cardiovascular disease, especially those classified as NCDs but caused by infectious pathogens. Integration of prevention and care remains as important today as in 1960 when Paul Dudley White and his colleagues found little evidence of myocardial infarction in the region near the Albert Schweitzer Hospital in Lambaréné, Gabon, but reported that “the high prevalence of mitral stenosis is astonishing.... We believe strongly that it is a duty to help bring to these sufferers the benefits of better penicillin prophylaxis and of cardiac surgery when indicated. The same responsibility exists for those with correctable congenital cardiovascular defects.”

RHD affects more than 15 million people worldwide, with more than 470,000 new cases each year. Among the 2.4 million annual cases of pediatric RHD, an estimated 42% occur in sub-Saharan Africa. This disease, which may cause endocarditis or stroke, leads to more than 345,000 deaths per year—almost all occurring in developing countries. Researchers in Ethiopia have reported annual death rates as high as 12.5% in rural areas. In part because the prevention of RHD has not advanced since the disease’s disappearance in wealthy countries, no part of sub-Saharan Africa has eradicated RHD despite examples of success in Costa Rica, Cuba, and some Caribbean nations. A survey of acute heart failure among adults in sub-Saharan Africa showed that ~14.3% of these cases were due to RHD.

Strategies to eliminate rheumatic heart disease may depend on active case-finding, with confirmation by echocardiography, among high-risk groups as well as on efforts to expand access to surgical interventions among children with advanced valvular damage. Partnerships between established surgical programs and areas with limited or non-existent facilities may help expand the capacity to provide life-saving interventions to patients who otherwise would die early and painfully. A long-term goal is the establishment of regional centers of excellence equipped to provide consistent, accessible, high-quality services.

Clinicians from tertiary care centers in sub-Saharan Africa and elsewhere have continued to call for prevention and treatment of the cardiovascular conditions of the poor. The reconstruction of health services in response to pandemic infectious disease offers an opportunity to identify and treat patients with organ damage and to undertake the prevention of cardiovascular and other chronic conditions of poverty.

CANCER Cancers account for ~5% of the global burden of disease. Low- and middle-income countries accounted for more than two-thirds of

the 12.6 million cases and 7.6 million deaths due to cancer in 2008. By 2030, annual mortality from cancer will increase by 4 million—with developing countries experiencing a sharper increase than developed nations. “Western” lifestyle changes will be responsible for the increased incidence of cancers of the breast, colon, and prostate among populations in low- and middle-income countries, but historic realities, sociocultural and behavioral factors, genetics, and poverty itself also will have a profound impact on cancer-related mortality and morbidity rates. At least 2 million cancer cases per year—18% of the global cancer burden—are attributable to infectious causes, which are responsible for <10% of cancers in developed countries but account for up to 20% of all malignancies in low- and middle-income countries. Infectious causes of cancer such as human papillomavirus, hepatitis B virus, and *Helicobacter pylori* will continue to have a much larger impact in developing countries. Environmental and dietary factors, such as indoor air pollution and high-salt diets, also contribute to increased rates of certain cancers (e.g., lung and gastric cancers). Tobacco use (both smoking and chewing) is the most important source of increased mortality rates from lung and oral cancers. In contrast to decreasing tobacco use in many developed countries, the number of smokers is growing in developing countries, especially among women and young persons.

For many reasons, outcomes of malignancies are far worse in developing countries than in developed nations. As currently funded, overstretched health systems in poor countries are not capable of early detection; the majority of patients already have incurable malignancies at diagnosis. Treatment of cancers is available for only a very small number of mostly wealthy citizens in the majority of poor countries, and, even when treatment is available, the range and quality of services are often substandard. Yet this need not be the future. Only a decade ago, MDR-TB and HIV infection were considered untreatable in settings of great poverty. The feasibility of creating innovative programs that reduce technical and financial barriers to the provision of care for treatable malignancies among the world’s poorest populations is now clear (Fig. 2-4). Several middle-income countries, including Mexico, have expanded publicly funded cancer care to reach poorer populations. This commitment of resources has dramatically improved outcomes for cancers, from childhood leukemia to cervical cancer.

Prevention of Noncommunicable Diseases False debates, including those pitting prevention against care, continue in global health and reflect, in part, outmoded paradigms or a partial understanding of disease burden and etiology as well as the dramatic variations in risk within a single nation. Moreover, debates are sometimes politicized as a



FIGURE 2-4 An 11-year-old Rwandan patient with embryonal rhabdomyosarcoma before (left) and after (right) 48 weeks of chemotherapy plus surgery. Five years later, she is healthy with no evidence of disease.

result of vested interests. For example, in 2004, the WHO released its Global Strategy on Diet, Physical Activity, and Health, which focused on the population-wide promotion of healthy diet and regular physical activity in an effort to reduce the growing global problem of obesity. Passing this strategy at the World Health Assembly proved difficult because of strong opposition from the food industry and from a number of WHO member states, including the United States. Although globalization has had many positive effects, one negative effect has been the growth in both developed and developing countries of well-financed lobbies that have aggressively promoted unhealthy dietary changes and increased consumption of alcohol and tobacco. Foreign direct investment in tobacco, beverage, and food products in developing countries reached \$90.3 billion in 2010—a figure nearly 490 times greater than the \$185 million spent during that year to address NCDs by bilateral funding agencies, the WHO, the World Bank, and all other sources of development assistance for health combined. Investment in curbing NCDs remains disproportionately low despite the WHO's 2008–2013 Action Plan for the Global Strategy for the Prevention and Control of Noncommunicable Diseases.

The WHO estimates that 80% of all cases of cardiovascular disease and type 2 diabetes as well as 40% of all cancers can be prevented through healthier diets, increased physical activity, and avoidance of tobacco. These estimates mask large local variations. Although some evidence indicates that population-based measures can have some impact on these behaviors, it is sobering to note that increasing obesity levels have not been reversed in any population. Tobacco avoidance may be the most important and most difficult behavioral modification of all. In the twentieth century, 100 million people worldwide died of tobacco-related diseases; it is projected that more than 1 billion people will die of these diseases in the twenty-first century, with the vast majority of those deaths in developing countries. The WHO's 2003 Framework Convention on Tobacco Control represented a major advance, committing all of its signatories to a set of policy measures shown to reduce tobacco consumption. Today, ~80% of the world's 1 billion smokers live in low- and middle-income countries. If trends continue, tobacco-related deaths will increase to 8 million per year by 2030, with 80% of those deaths in low- and middle-income countries.

MENTAL HEALTH

The WHO reports that some 450 million people worldwide are affected by mental, neurologic, or behavioral problems at any given time and that ~877,000 people die by suicide every year. Major depression is the leading cause of years lost to disability in the world today. One in four patients visiting a health service has at least one mental, neurologic, or behavioral disorder, but most of these disorders are neither diagnosed nor treated. Most low- and middle-income countries devote <1% of their health expenditures to mental health.

Increasingly effective therapies exist for many of the major causes of mental disorders. Effective treatments for many neurologic diseases, including seizure disorders, have long been available. One of the greatest barriers to delivery of such therapies is the paucity of skilled personnel. Most sub-Saharan African countries have only a handful of psychiatrists, for example; most of them practice in cities and are unavailable within the public sector or to patients living in poverty.

Among the few patients who are fortunate enough to see a psychiatrist or neurologist, fewer still are able to adhere to treatment regimens: several surveys of already diagnosed patients ostensibly receiving daily therapy have revealed that, among the poor, multiple barriers prevent patients from taking their medications as prescribed. In one study from Kenya, no patients being seen in an epilepsy clinic had therapeutic blood levels of anti-seizure medications, even though all had had these drugs prescribed. Moreover, many patients had no detectable blood levels of these agents. The same barriers that prevent the poor from having reliable access to insulin or ART prevent them from benefiting from antidepressant, antipsychotic, and antiepileptic agents. To alleviate this problem, some authorities are proposing the training of health workers to provide community-based adherence support,

counseling services, and referrals for patients in need of mental health services. One such program instituted in Goa, India, used “lay” counselors and resulted in a significant reduction in symptoms of common mental disorders among the target population.

World Mental Health: Problems and Priorities in Low-Income Countries still offers a comprehensive analysis of the burden of mental, behavioral, and social problems in low-income countries and relates the mental health consequences of social forces such as violence, dislocation, poverty, and disenfranchisement of women to current economic, political, and environmental concerns. In the years since this report was published, however, a number of pilot projects designed to deliver community-based care to patients with chronic mental illness have been launched in settings as diverse as Goa, India; Banda Aceh, Indonesia; rural China; post-earthquake Haiti; and Fiji. Some of these programs have been school-based and have sought to link prevention to care.

CONCLUSION: TOWARD A SCIENCE OF IMPLEMENTATION

Public health strategies draw largely on quantitative methods—epidemiology, biostatistics, and economics. Clinical practice, including the practice of internal medicine, draws on a rapidly expanding knowledge base but remains focused on individual patient care; clinical interventions are rarely population-based. But global health equity depends on avoiding the false debates of the past: neither public health nor clinical approaches alone are adequate to address the problems of global health. There is a long way to go before evidence-based internal medicine is applied effectively among the world's poor. Complex infectious diseases such as HIV/AIDS and TB have proved difficult but not impossible to manage; drug resistance and lack of effective health systems have complicated such work. Beyond what is usually termed “communicable diseases”—i.e., in the arena of chronic diseases such as cardiovascular disease and mental illness—global health is a nascent endeavor. Efforts to address any one of these problems in settings of great scarcity need to be integrated into broader efforts to strengthen failing health systems and alleviate the growing personnel crisis within these systems.

Such efforts must include the building of “platforms” for care delivery that are robust enough to incorporate new preventive, diagnostic, and therapeutic technologies rapidly in response to changes both in the burden of disease and in the needs not met by dominant paradigms and systems of health delivery. Academic medical centers have tried to address this “know–do” gap as new technologies are introduced and assessed through clinical trials, but the reach of these institutions into settings of poverty is limited in rich and poor countries alike. When such centers link their capacities effectively to the public institutions charged with the delivery of health care to the poor, great progress can be made.

For these reasons, scholarly work and practice in the field once known as “international health” and now often designated “global health equity” are changing rapidly. That work is still informed by the tension between clinical practice and population-based interventions, between analysis and action, and between prevention and care. Once metrics are refined, how might they inform efforts to lessen premature morbidity and mortality rates among the world's poor? As in the nineteenth century, human rights perspectives have proved helpful in turning attention to the problems of the destitute sick; such perspectives may also inform strategies for delivering care equitably.

A number of university hospitals are developing training programs for physicians with an interest in global health. In medical schools across the United States and in other wealthy countries, interest in global health has exploded. One study has shown that more than 25% of medical students take part in at least one global health experience prior to graduation. Half a century or even a decade ago, such high levels of interest would have been unimaginable.

An estimated 12 million people die each year simply because they live in poverty. An absolute majority of these premature deaths occur in Africa, with the poorer regions of Asia not far behind. Most of these deaths occur because the world's poorest do not have access to the fruits of science. They include deaths from vaccine-preventable illness, deaths during childbirth, deaths from infectious diseases that might be

cured with access to antibiotics and other essential medicines, deaths from malaria that would have been prevented by bed nets and access to therapy, and deaths from waterborne illnesses. Other excess mortality is attributable to the inadequacy of efforts to develop new preventive, diagnostic, and therapeutic tools. Those funding the discovery and development of new tools typically neglect the concurrent need for strategies to make them available to the poor. Indeed, some would argue that the biggest challenge facing those who seek to address this outcome gap is the lack of practical means of distribution in the most heavily affected regions.

The development of tools must be followed quickly by their equitable distribution. When new preventive and therapeutic tools are developed without concurrent attention to delivery or implementation, one encounters what are sometimes termed *perverse effects*: even as new tools are developed, inequalities of outcome—lower morbidity and mortality rates among those who can afford access, with sustained high morbidity and mortality among those who cannot—will grow in the absence of an equity plan to deliver the tools to those most at risk. Preventing such a future is the most important goal of global health.

3

Decision-Making in Clinical Medicine

Daniel B. Mark, John B. Wong

INTRODUCTION

To a medical student who requires hours to collect a patient's history, perform a physical examination, and organize that information into a coherent presentation, an experienced clinician's ability to decide on a diagnosis and management plan in minutes may seem extraordinary. What separates the master clinician from the novice is an elusive quality called "expertise." The first part of this chapter provides an overview of our current understanding of expertise in clinical reasoning, what it is, and how it can be developed.

The proper use of diagnostic tests and the integration of the results into the patient's clinical assessment may also be equally bewildering to students. Hoping to hit the unknown diagnostic target, novice medical practitioners typically apply a "shotgun" approach to testing. The expert, in contrast, usually focuses her testing strategy to specific diagnostic hypotheses. The second part of the chapter reviews basic statistical concepts useful for interpreting diagnostic tests and quantitative tools useful for clinical decision-making.

Evidence-based medicine (EBM) constitutes the integration of the best available research evidence with clinical judgment as applied to the care of individual patients. The third part of the chapter provides an overview of the tools of EBM.

BRIEF INTRODUCTION TO CLINICAL REASONING

Clinical Expertise Defining "clinical expertise" remains surprisingly difficult. Chess has an objective ranking system based on skill and performance criteria. Athletics, similarly, have ranking systems to distinguish novices from Olympians. But in medicine, after physicians complete training and pass the boards, no further tests or benchmarks identify those who have attained the highest levels of clinical performance. Of course, physicians often consult a few "elite" clinicians for their "special problem-solving prowess" when particularly difficult or obscure cases have baffled everyone else. Yet despite their skill, even master clinicians typically cannot explain their exact processes and methods, thereby limiting the acquisition and dissemination of the expertise used to achieve their impressive results. Furthermore, clinical virtuosity appears not to be generalizable, e.g., an expert on hypertrophic cardiomyopathy may be no better (and possibly worse) than a first-year medical resident at diagnosing and managing a patient with neutropenia, fever, and hypotension.

Broadly construed, clinical expertise includes not only cognitive dimensions and the integration of verbal and visual cues or information but also complex fine-motor skills necessary for invasive and noninvasive procedures and tests. In addition, "the complete package" of expertise in medicine includes the ability to communicate effectively with patients and work well with members of the medical team. Research on medical expertise remains relatively sparse overall, with most of the work focused on diagnostic reasoning, and much less work focused on treatment decisions or the technical skills involved in the performance of procedures. Thus, in this chapter, we focus primarily on the cognitive elements of clinical reasoning.

Because clinical reasoning takes place in the heads of doctors, it is therefore not readily observable, making it obviously difficult to study. One method of research on reasoning asks doctors to "think out loud" as they receive increments of clinical information in a manner meant to simulate a clinical encounter. Another research approach has focused on how doctors *should* reason diagnostically rather than on how they actually *do* reason. Much of what is known about clinical reasoning comes from empirical studies of nonmedical problem-solving behavior. Because of the diverse perspectives contributing to this area, with important contributions from cognitive psychology, sociology, medical education, economics, informatics, and decision sciences, no single integrated model of clinical reasoning exists, and not infrequently, different terms and models describe similar phenomena.

Intuitive versus Analytic Reasoning A contemporary model of reasoning, *dual-process theory* distinguishes two general systems of cognitive processes. *Intuition* (System 1) provides rapid effortless judgments from memorized associations using pattern recognition and other simplifying "rules of thumb" (i.e., heuristics). For example, a very simple pattern that could be useful in certain situations is "African-American women plus hilar adenopathy equals sarcoid." Because no effort is involved in recalling the pattern, typically, the clinician is unable to say how those judgments were formulated. In contrast, *analysis* (System 2), the other form of reasoning in the dual-process model, is slow, methodical, deliberative, and effortful. These are, of course, idealized extremes of the cognitive continuum. How these systems interact in different decision problems, how experts use them differently from novices, and when their use can lead to errors in judgment remain the subject of considerable study and debate.

Pattern recognition is a complex cognitive process that appears largely effortless. One can recognize people's faces, the breed of a dog, or an automobile model without necessarily being able to say what specific features prompted the recognition. Analogously, experienced clinicians often recognize familiar diagnosis patterns quickly. In the absence of an extensive stored repertoire of diagnostic patterns, students (as well as more experienced clinicians operating outside their area of expertise) often use the more laborious System 2 analytic approach along with more intensive and comprehensive data collection to reach the diagnosis.

The following three brief scenarios of a patient with hemoptysis demonstrate three distinct patterns:

- A 46-year-old man presents to his internist with a chief complaint of hemoptysis. An otherwise healthy nonsmoker, he is recovering from an apparent viral bronchitis. This presentation pattern suggests that the small amount of blood-streaked sputum is due to acute bronchitis, so that a chest x-ray provides sufficient reassurance that a more serious disorder is absent.
- In the second scenario, a 46-year-old patient who has the same chief complaint but with a 100-pack-year smoking history, a productive morning cough, and episodes of blood-streaked sputum fits the pattern of carcinoma of the lung. Consequently, along with the chest x-ray, the physician obtains a sputum cytology examination and refers this patient for a chest computed tomography (CT) scan.
- In the third scenario, a 46-year-old patient with hemoptysis who immigrated from a developing country has an echocardiogram as well, because the physician hears a soft diastolic rumbling murmur at the apex on cardiac auscultation, suggesting rheumatic mitral stenosis and possibly pulmonary hypertension.

Although rapid, pattern recognition used without sufficient reflection can result in *premature closure*: mistakenly concluding that one already knows the correct diagnosis and therefore failing to complete the data collection that would demonstrate the lack of fit of the initial pattern selected. For example, a 45-year-old man presents with a 3-week history of a “flulike” upper respiratory infection (URI) including symptoms of dyspnea and a productive cough. On the basis of the presenting complaints, the clinician uses a “URI assessment form” to improve the quality and efficiency of care by standardizing the information gathered. After quickly acquiring the requisite structured examination components and noting in particular the absence of fever and a clear chest examination, the physician prescribes medication for acute bronchitis and sends the patient home with the reassurance that his illness was not serious. Following a sleepless night with significant dyspnea, the patient develops nausea and vomiting and collapses. He presents to the emergency department in cardiac arrest and is unable to be resuscitated. His autopsy shows a posterior wall myocardial infarction and a fresh thrombus in an atherosclerotic right coronary artery. What went wrong? The clinician had decided, based on the patient’s appearance, even before starting the history, that the patient’s complaints were not serious. Therefore, he felt confident that he could perform an abbreviated and focused examination by using the URI assessment protocol rather than considering the broader range of possibilities and performing appropriate tests to confirm or refute his initial hypotheses. In particular, by concentrating on the URI, the clinician failed to elicit the full dyspnea history, which would have suggested a far more serious disorder, and he neglected to search for other symptoms that could have directed him to the correct diagnosis.

Heuristics, also referred to as cognitive shortcuts or rules of thumb, are simplifying decision strategies that ignore part of the data available so as to provide an efficient path to the desired judgment. They are generally part of the intuitive system tools. Two major research programs have come to different conclusions about the value of heuristics in clinical judgment. The “heuristics and biases” program focused on understanding how heuristics in problem solving could be biased by testing the numerical intuition of psychology undergraduates against the rules of statistics. In contrast, the “fast and frugal heuristics” research program explored how and when decision makers’ reliance on simple heuristics can produce good decisions. Although many heuristics have relevance to clinical reasoning, only four will be mentioned here.

When assessing a particular patient, clinicians often weigh the similarity of that patient’s symptoms, signs, and risk factors against those of their mental representations of the diagnostic hypotheses being considered. In other words, among the diagnostic possibilities, clinicians identify the diagnosis for which the patient appears to be a representative example. Analogous to pattern recognition, this cognitive shortcut is called the *representativeness heuristic*. However, physicians using the representativeness heuristic can reach erroneous conclusions if they fail to consider the underlying prevalence (i.e., the prior, or pretest, probabilities) of the two competing diagnoses that could explain the patient’s symptoms. Consider a patient with hypertension and headache, palpitations, and diaphoresis. Inexperienced clinicians might judge pheochromocytoma to be quite likely based on the representativeness heuristic with this classic symptom triad suggesting pheochromocytoma. Doing so would be incorrect given that other causes of hypertension are much more common than pheochromocytoma, and this triad of symptoms can occur in patients who do not have pheochromocytoma. Less experience with a particular diagnosis and with the breadth of presentations (e.g., diseases that affect multiple organ systems such as sarcoid) may also lead to errors.

A second commonly used cognitive shortcut, the *availability heuristic*, involves judgments based on how easily prior similar cases or outcomes can be brought to mind. For example, an experienced clinician may recall 20 elderly patients seen over the last few years who presented with painless dyspnea of acute onset and were found to have acute myocardial infarction (MI). A novice clinician may spend valuable time seeking a pulmonary cause for the symptoms before

considering and then confirming the cardiac diagnosis. In this situation, the patient’s clinical pattern does not fit the most common pattern of acute MI, but experience with this atypical presentation, along with the ability to recall it, directs the physician to the diagnosis.

Errors with the availability heuristic arise from several sources of recall bias. Rare catastrophes are likely to be remembered with a clarity and force disproportionate to their likelihood for future diagnosis—for example, a patient with a sore throat eventually found to have leukemia or a young athlete with leg pain eventually found to have a sarcoma—and those publicized in the media or that are recent experiences are, of course, easier to recall and therefore more influential on clinical judgments.

The third commonly used cognitive shortcut, the *anchoring heuristic* (also called conservatism or stickiness), involves estimating a probability of disease (the anchor) and then insufficiently adjusting that probability up or down (compared with Bayes’ rule) when interpreting new data about the patient, i.e., sticking to their initial diagnosis. For example, a clinician may still judge the probability of coronary artery disease (CAD) to be high after a negative exercise thallium test and proceed to cardiac catheterization (see “Measures of Disease Probability and Bayes’ Rule,” below).

The fourth heuristic states that clinicians should use the simplest explanation possible that will account adequately for the patient’s symptoms or findings (Occam’s razor or, alternatively, the *simplicity heuristic*). Although this is an attractive and often used principle, it is important to remember that no biologic basis for it exists. Errors from the simplicity heuristic include premature closure leading to the neglect of unexplained significant symptoms or findings.

Even experienced physicians use analytic reasoning processes (System 2) when the problem they face is recognized to be complex or to involve important unfamiliar elements or features. In such situations, clinicians proceed much more methodically in what has been referred to as the hypothetico-deductive model of reasoning. From the outset, expert clinicians working analytically generate, refine, and discard diagnostic hypotheses. The hypotheses drive questions asked during history taking and may change based on the working hypotheses of the moment. Even the physical examination is focused by the working hypotheses. Is the spleen enlarged? How big is the liver? Is it tender? Are there any palpable masses or nodules? Each question must be answered (with the exclusion of all other inputs) before the examiner can move on to the next specific question. Each diagnostic hypothesis provides testable predictions and sets a context for the next question or step to follow. For example, if the enlarged and quite tender liver felt on physical examination is due to acute hepatitis (the hypothesis), certain specific liver function tests should be markedly elevated (the prediction). If the tests come back normal, the hypothesis may have to be discarded or substantially modified.

Negative findings often are neglected but are as important as positive ones because they often reduce the likelihood of the diagnostic hypotheses under consideration. Chest discomfort that is not provoked or worsened by exertion in an active patient reduces the likelihood that chronic ischemic heart disease is the underlying cause. The absence of a resting tachycardia and thyroid gland enlargement reduces the likelihood of hyperthyroidism in a patient with paroxysmal atrial fibrillation.

The acuity of a patient’s illness may override considerations of prevalence and the other issues described above. “Diagnostic imperatives” recognize the significance of relatively rare but potentially catastrophic diagnoses if undiagnosed and untreated. For example, clinicians are taught to consider aortic dissection routinely as a possible cause of acute severe chest discomfort. Even though the typical history of dissection differs from that of MI, dissection is far less prevalent, so diagnosing dissection remains challenging unless it is explicitly and routinely considered as a diagnostic imperative (Chap. 301). If the clinician fails to elicit any of the characteristic features of dissection by history and finds equivalent blood pressures in both arms and no pulse deficits, he may feel comfortable discarding the aortic dissection hypothesis. If, however, the chest x-ray shows a possible widened mediastinum, the hypothesis may be reinstated and an appropriate

imaging test ordered (e.g., thoracic CT scan, transesophageal echocardiogram) to evaluate more fully. In nonacute situations, the prevalence of potential alternative diagnoses should play a much more prominent role in diagnostic hypothesis generation.

Cognitive scientists studying the thought processes of expert clinicians have observed that clinicians group data into packets, or “chunks,” that are stored in short-term or “working memory” and manipulated to generate diagnostic hypotheses. Because short-term memory can typically retain only 5–9 items at a time, the number of packets that can be actively integrated into hypothesis-generating activities is similarly limited. For this reason, the cognitive shortcuts discussed above play a key role in the generation of diagnostic hypotheses, many of which are discarded as rapidly as they are formed (thereby demonstrating that the distinction between analytic and intuitive reasoning is an arbitrary and simplistic, but nonetheless useful, representation of cognition).

Research into the hypothetico-deductive model of reasoning has had surprising difficulty identifying the elements of the reasoning process that distinguish experts from novices. This has led to a shift from examining the problem-solving process of experts to analyzing the organization of their knowledge. For example, diagnosis may be based on the resemblance of a new case to prior individual instances (exemplars). Experts have a much larger store of memorized cases, for example, visual long-term memory in radiology. However, clinicians do not simply rely on literal recall of specific cases but have constructed elaborate conceptual networks of memorized information or models of disease to aid in arriving at their conclusions. That is, expertise involves an increased ability to connect symptoms, signs, and risk factors to one another in meaningful ways; relate those findings to possible diagnoses; and identify the additional information necessary to confirm the diagnosis.

No single theory accounts for all the key features of expertise in medical diagnosis. Experts have more knowledge about more things and a larger repertoire of cognitive tools to employ in problem solving than do novices. One definition of expertise highlights the ability to make powerful distinctions. In this sense, expertise involves a working knowledge of the diagnostic possibilities and what features distinguish one disease from another. Memorization alone is insufficient. Memorizing a medical textbook would not make one an expert. But having access to detailed and specific relevant information is critically important. Clinicians of the past primarily accessed their own remembered experience. Clinicians of the future will be able to access the experience of large numbers of clinicians using electronic tools, but, as with the memorized textbook, the data alone will not create an instant expert. The expert adds these data to an extensive internalized database of knowledge and experience not available to the novice (and nonexpert).

Despite all the work that has been done to understand expertise, in medicine and other disciplines, it remains uncertain whether there is any didactic program that can accelerate the progression from novice to expert or from experienced clinician to master clinician. Deliberate effortful practice (over an extended period of time, sometimes said to be 10 years or 10,000 practice hours) and personal coaching are two strategies that are often used outside medicine (e.g., music, athletics, chess) to promote expertise. Their use in developing medical expertise and maintaining or enhancing it has not yet been adequately explored.

DIAGNOSTIC VERSUS THERAPEUTIC DECISION MAKING

The modern ideal of medical therapeutic decision making is to “personalize” the recommendation. In the abstract, personalizing treatment involves combining the best available evidence about what works with an individual patient’s unique features (e.g., risk factors) and his or her preferences and health goals to craft an optimal treatment recommendation with the patient. Operationally, there are two different and complementary levels of personalization possible: individualizing the evidence for the specific patient based on relevant clinical and other characteristics, and personalizing the patient interaction by incorporating their values, often referred to as shared decision-making, which is critically important, but falls outside the scope of this chapter.

Individualizing the evidence about therapy **does not** mean relying on physician impressions of what works based on personal experience. Because of small sample sizes and rare events, the chance of drawing erroneous causal inferences from one’s own clinical experience is very high. For most chronic diseases, therapeutic effectiveness is only demonstrable statistically in patient populations. It would be incorrect to infer with any certainty, for example, that treating a hypertensive patient with angiotensin-converting enzyme (ACE) inhibitors necessarily prevented a stroke from occurring during treatment, or that an untreated patient would definitely have avoided a stroke had he or she been treated. For many chronic diseases, a majority of patients will remain event free regardless of treatment choices; some will have events regardless of which treatment is selected; and those who avoided having an event through treatment cannot be individually identified. Blood pressure lowering, a readily observable surrogate endpoint, does not have a tightly coupled relationship with strokes prevented. Consequently, demonstrating therapeutic effectiveness cannot rely simply on observing the outcome of an individual patient but should instead be based on large groups of patients carefully studied and properly analyzed.

Therapeutic decision making, therefore, should be based on the best available evidence from clinical trials and well-done outcome studies. Authoritative, well-done clinical practice guidelines that synthesize such evidence offer readily available, reliable, and trustworthy information relevant to many treatment decisions clinicians face. However, all guidelines recognize that their “one size fits all” recommendations may not apply to individual patients. Increased attention is now being paid to understand how best to adjust group-level clinical evidence of treatment harms and benefits to account for the absolute level of risks faced by subgroups and even individual patients, using, for example, validated clinical risk scores.

NONCLINICAL INFLUENCES ON CLINICAL DECISION-MAKING

More than a decade of research on variations in clinician practice patterns has shed much light on the forces that shape clinical decisions. These factors can be grouped conceptually into three overlapping categories: (1) factors related to physicians’ personal characteristics and practice style, (2) factors related to the practice setting, and (3) factors related to economic incentives.

Factors Related to Practice Style To ensure that necessary care is provided at a high level of quality, physicians fulfill a key role in medical care by serving as the patient’s agent. Factors that influence performance in this role include the physician’s knowledge, training, and experience. Clearly, physicians cannot practice EBM (described later in the chapter) if they are unfamiliar with the evidence. As would be expected, specialists generally know the evidence in their field better than do generalists. Beyond published evidence and practice guidelines, a major set of influences on physician practice can be subsumed under the general concept of “practice style.” The practice style serves to define norms of clinical behavior. Beliefs about effectiveness of different therapies and preferred patterns of diagnostic test use are examples of different facets of a practice style. The physician beliefs that drive these different practice styles may be based on personal experience, recollection, and interpretation of the available medical evidence. For example, heart failure specialists are much more likely than generalists to achieve target doses of ACE inhibitor therapy in their heart failure patients because they are more familiar with what the targets are (as defined by large clinical trials), have more familiarity with the specific drugs (including adverse effects), and are less likely to overreact to foreseeable problems in therapy such as a rise in creatinine levels or asymptomatic hypotension.

Beyond the patient’s welfare, physician perceptions about the risk of a malpractice suit resulting from either an erroneous decision or a bad outcome may drive clinical decisions and create a practice referred to as *defensive medicine*. This practice involves using tests and therapies with very small marginal benefit, ostensibly to preclude future criticism should an adverse outcome occur. Without any conscious awareness of a connection to the risk of litigation, however, over time such patterns of care may become accepted as part of the practice norm,

thereby perpetuating their overuse, e.g., annual cardiac exercise testing in asymptomatic patients.

Practice Setting Factors Factors in this category relate to the physical resources available to the physician's practice and the practice environment. *Physician-induced demand* is a term that refers to the repeated observation that once medical facilities and technologies are made available to physicians, they will use them. Other environmental factors that can influence decision-making include the local availability of specialists for consultations and procedures; "high-tech" advanced imaging or procedure facilities such as MRI machines and proton beam therapy centers; and fragmentation of care.

Economic Incentives Economic incentives are closely related to the other two categories of practice-modifying factors. Financial issues can exert both stimulatory and inhibitory influences on clinical practice. In general, physicians are paid on a fee-for-service, capitation, or salary basis. In fee-for-service, physicians who do more get paid more, thereby encouraging overuse, consciously or unconsciously. When fees are reduced (discounted reimbursement), doctors tend to increase the number of services provided to maintain revenue. Capitation, in contrast, provides a fixed payment per patient per year to encourage physicians to consider a global population budget in managing individual patients and ideally reducing the use of interventions with small marginal benefit. In contrast to inexpensive preventive services, however, this type of incentive is more likely to affect expensive interventions. To discourage volume-based excessive utilization, fixed salary compensation plans pay physicians the same regardless of the clinical effort expended, but may provide an incentive to see fewer patients.

INTERPRETATION OF DIAGNOSTIC TESTS IN THE CONTEXT OF DECISION-MAKING

Despite the great technological advances in medicine over the last century, uncertainty remains a key challenge in all aspects of medical decision-making. Compounding this challenge is the massive information overload that characterizes modern medicine. Today's clinician needs access to close to 2 million pieces of information to practice medicine. According to one estimate, doctors subscribe to an average of seven journals, representing over 2500 new articles each year. Of course, to be useful, this information must be sifted for applicability to and then integrated with patient-specific data. Although computers appear to offer an obvious solution both for information management and for quantification of medical care uncertainties, many practical problems must be solved before computerized decision support can be routinely incorporated into the clinical reasoning process in a way that demonstrably improves the quality of care. For the present, understanding the nature of diagnostic test information can help clinicians become more efficient users of such data. The next section reviews important concepts related to diagnostic testing.

DIAGNOSTIC TESTING: MEASURES OF TEST ACCURACY

The purpose of performing a test on a patient is to reduce uncertainty about the patient's diagnosis or prognosis in order to facilitate optimal management. Although diagnostic tests commonly are thought of as laboratory tests (e.g., blood count) or procedures (e.g., colonoscopy or bronchoscopy), any technology that changes a physician's understanding of the patient's problem qualifies as a diagnostic test. Thus, even the history and physical examination can be considered a form of diagnostic test. In clinical medicine, it is common to reduce the results of a test to a dichotomous outcome, such as positive or negative, normal or abnormal. Although this simplification ignores useful information (such as the degree of abnormality), such simplification does make it easier to demonstrate the fundamental principles of test interpretation discussed below.

The accuracy of diagnostic tests is defined in relation to an accepted "gold standard," which defines the presumably true state of the patient (Table 3-1). Characterizing the diagnostic performance of a new test requires identifying an appropriate population (ideally, patients in whom the new test would be used) and applying both the new and the gold standard tests to all subjects. Biased estimates of test performance

TABLE 3-1 MEASURES OF DIAGNOSTIC TEST ACCURACY

Test Result	Disease Status	
	Present	Absent
Positive	True positive (TP)	False positive (FP)
Negative	False negative (FN)	True negative (TN)
Test Characteristics in Patients with Disease		
True-positive rate (sensitivity) = $TP/(TP + FN)$		
False-negative rate = $FN/(TP + FN)$		
True-positive rate = $1 - \text{false-negative rate}$		
Test Characteristics in Patients without Disease		
True-negative rate (specificity) = $TN/(TN + FP)$		
False-positive rate = $FP/(TN + FP)$		
True-negative rate = $1 - \text{false-positive rate}$		

may occur from using an inappropriate population or from incompletely applying the gold standard test. By comparing the two tests, the characteristics of the new test are determined. The *sensitivity* or *true-positive rate* of the new test is the proportion of patients with disease (defined by the gold standard) who have a positive (new) test. This measure reflects how well the new test identifies patients with disease. The proportion of patients with disease who have a negative test is the *false-negative rate* and is calculated as $1 - \text{sensitivity}$. Among patients without disease, the proportion who have a negative test is the *specificity*, or *true-negative rate*. This measure reflects how well the new test correctly identifies patients without disease. Among patients without disease, the proportion who have a positive test is the *false-positive rate*, calculated as $1 - \text{specificity}$. A perfect test would have a sensitivity of 100% and a specificity of 100% and would completely distinguish patients with disease from those without it.

Calculating sensitivity and specificity requires selection of a threshold value or cut point above which the test is considered "positive." Making the cut point "stricter" (e.g., raising it) lowers sensitivity but improves specificity, whereas making it "laxer" (e.g., lowering it) raises sensitivity but lowers specificity. This dynamic trade-off between more accurate identification of subjects with disease versus those without disease is often displayed graphically as a receiver operating characteristic (ROC) curve (Fig. 3-1) by plotting sensitivity (y axis) versus $1 - \text{specificity}$ (x axis). Each point on the curve represents a potential cut point with an associated sensitivity and specificity value. The area under the ROC curve often is used as a quantitative measure of the information content of a test. Values range from 0.5 (no diagnostic information from testing at all; the test is equivalent to flipping a coin) to 1.0 (perfect test). The choice of cut point should depend on the relative harms and benefits of treatment for those without versus those with disease. For example, if treatment was safe with substantial benefit, then choosing a high-sensitivity cut point (upper right of the ROC curve) for a low-risk test may be appropriate (e.g., phenylketonuria in newborns), but if treatment had substantial risk for harm, then choosing a high-specificity cut point (lower left of the ROC curve) may be appropriate (e.g., amniocentesis that may lead to therapeutic abortion of a normal fetus). The choice of cut point may also depend on the likelihood of disease, with low likelihoods placing a greater emphasis on the harms of treating false-positive tests and higher likelihoods placing a greater emphasis on missed benefit by not treating false-negative tests.

MEASURES OF DISEASE PROBABILITY AND BAYES' RULE

Unfortunately, there are no perfect tests. After every test is completed, the true disease state of the patient remains uncertain. Quantifying this residual uncertainty can be done with Bayes' rule, which provides a simple way to calculate the likelihood of disease after a test result or posttest probability from three parameters: the pretest probability of disease, the test sensitivity, and the test specificity. The pretest probability is a quantitative estimate of the likelihood of the diagnosis before the test is performed and is usually the prevalence of the disease in the underlying population although occasionally it can be the disease

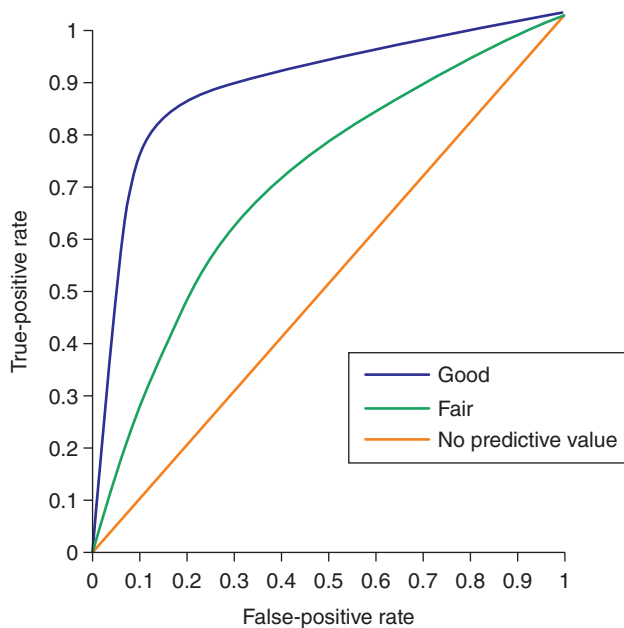


FIGURE 3-1 Each receiver operating characteristic (ROC) curve illustrates a trade-off that occurs between improved test sensitivity (accurate detection of patients with disease) and improved test specificity (accurate detection of patients without disease), because the test value defining when the test turns from “negative” to “positive” is varied. A 45° line would indicate a test with no predictive value (sensitivity = specificity at every test value). The area under each ROC curve is a measure of the information content of the test. Thus, a larger ROC area signifies increased diagnostic accuracy.

incidence. For some common conditions, such as CAD, nomograms and statistical models generate estimates of pretest probability that account for history, physical examination, and test findings. The posttest probability (also called the predictive value of the test) is a revised statement of the likelihood of the diagnosis, accounting for both pretest probability and test results. For the likelihood of disease following a positive test (i.e., positive predictive value), Bayes’ rule is calculated as:

$$\text{Posttest probability} = \frac{\text{Pretest probability} \times \text{Sensitivity}}{\text{Pretest probability} \times \text{Sensitivity} + (1 - \text{Pretest probability}) \times \text{False-positive Rate}}$$

For example, with a pretest probability of 0.50 and a “positive” diagnostic test result (test sensitivity = 0.90 and specificity = 0.90):

$$\text{Posttest probability} = \frac{0.50 \times 0.90}{0.50 \times 0.90 + (1 - 0.50) \times 0.10} = 0.90$$

The term *predictive value* often is used as a synonym for the posttest probability. Unfortunately, clinicians commonly misinterpret reported predictive values as intrinsic measures of test accuracy. Studies of diagnostic tests compound the confusion by calculating predictive values on the same sample used to measure sensitivity and specificity. Since all posttest probabilities are a function of the prevalence of disease in the tested population, such calculations may be misleading unless the test is applied subsequently to populations with the same disease prevalence. For these reasons, the term *predictive value* is best avoided in favor of the more informative *posttest probability* following a positive or a negative test result.

The nomogram version of Bayes’ rule (Fig. 3-2) helps us to conceptually understand how it estimates the posttest probability of disease. In this nomogram, the impact of the diagnostic test result is summarized by the *likelihood ratio*, which is defined as the ratio of

the probability of a given test result (e.g., “positive” or “negative”) in a patient with disease to the probability of that result in a patient without disease, thereby providing a measure of how well the test distinguishes those with from those without disease.

For a positive test, the likelihood ratio positive is calculated as the ratio of the true-positive rate to the false-positive rate (or sensitivity/[1 – specificity]). For example, a test with a sensitivity of 0.90 and a specificity of 0.90 has a likelihood ratio of 0.90/(1 – 0.90), or 9. Thus, for this hypothetical test, a “positive” result is nine times more likely in a patient with the disease than in a patient without it. Most tests in medicine have likelihood ratios for a positive result between 1.5 and 20. Higher values are associated with tests that more substantially increase the posttest likelihood of disease. A very high likelihood ratio positive (exceeding 10) usually implies high specificity, so a positive high-specificity test helps “rule in” disease. If sensitivity is excellent but specificity is less so, the likelihood ratio will be reduced substantially (e.g., with a 90% sensitivity but a 55% specificity, the likelihood ratio is 2.0).

For a negative test, the corresponding likelihood ratio negative is the ratio of the false-negative rate to the true-negative rate (or [1 – sensitivity]/specificity). Lower likelihood ratio values more substantially lower the posttest likelihood of disease. A very low likelihood ratio negative (falling below 0.10) usually implies high sensitivity, so a negative high-sensitivity test helps “rule out” disease. The hypothetical test considered above with a sensitivity of 0.9 and a specificity of 0.9 would have a likelihood ratio for a negative test result of (1 – 0.9)/0.9, or 0.11, meaning that a negative result is about one-tenth as likely in patients with disease than in those without disease (or 10 times more likely in those without disease than in those with disease).

APPLICATIONS TO DIAGNOSTIC TESTING IN CAD

Consider two tests commonly used in the diagnosis of CAD: an exercise treadmill test and an exercise single-photon emission CT (SPECT) myocardial perfusion imaging test (Chap. 270e). Meta-analysis has shown that a positive treadmill ST-segment response has an average sensitivity of 66% and an average specificity of 84%, yielding a likelihood ratio of 4.1 (0.66/[1 – 0.84]) (consistent with small discriminatory ability because it falls between 2 and 5). For a patient with a 10% pretest probability of CAD, the posttest probability of disease after a positive result rises to only about 30%. If a patient with a pretest probability of CAD of 80% has a positive test result, the posttest probability of disease is about 95%.

In contrast, exercise SPECT myocardial perfusion test is more accurate for CAD. For simplicity, assume that the finding of a reversible exercise-induced perfusion defect has both a sensitivity and a specificity of 90%, yielding a likelihood ratio for a positive test of 9.0 (0.90/[1 – 0.90]) (consistent with moderate discriminatory ability because it falls between 5 and 10). For the same 10% pretest probability patient, a positive test raises the probability of CAD to 50% (Fig. 3-2). However, despite the differences in posttest probabilities between these two tests (30% versus 50%), the more accurate test may not improve diagnostic likelihood enough to change patient management (e.g., decision to refer to cardiac catheterization) because the more accurate test has only moved the physician from being fairly certain that the patient did not have CAD to a 50:50 chance of disease. In a patient with a pretest probability of 80%, exercise SPECT test raises the posttest probability to 97% (compared with 95% for the exercise treadmill). Again, the more accurate test does not provide enough improvement in posttest confidence to alter management, and neither test has improved much on what was known from clinical data alone.

In general, positive results with an accurate test (e.g., likelihood ratio positive 10) when the pretest probability is low (e.g., 20%) do not move the posttest probability to a range high enough to rule in disease (e.g., 80%). In screening situations, pretest probabilities are often particularly low because patients are asymptomatic. In such cases, specificity becomes particularly important. For example, in screening first-time female blood donors without risk factors for HIV, a positive test raised the likelihood of HIV to only 67% despite a specificity of 99.995% because the prevalence was 0.01%. Conversely, with a high pretest

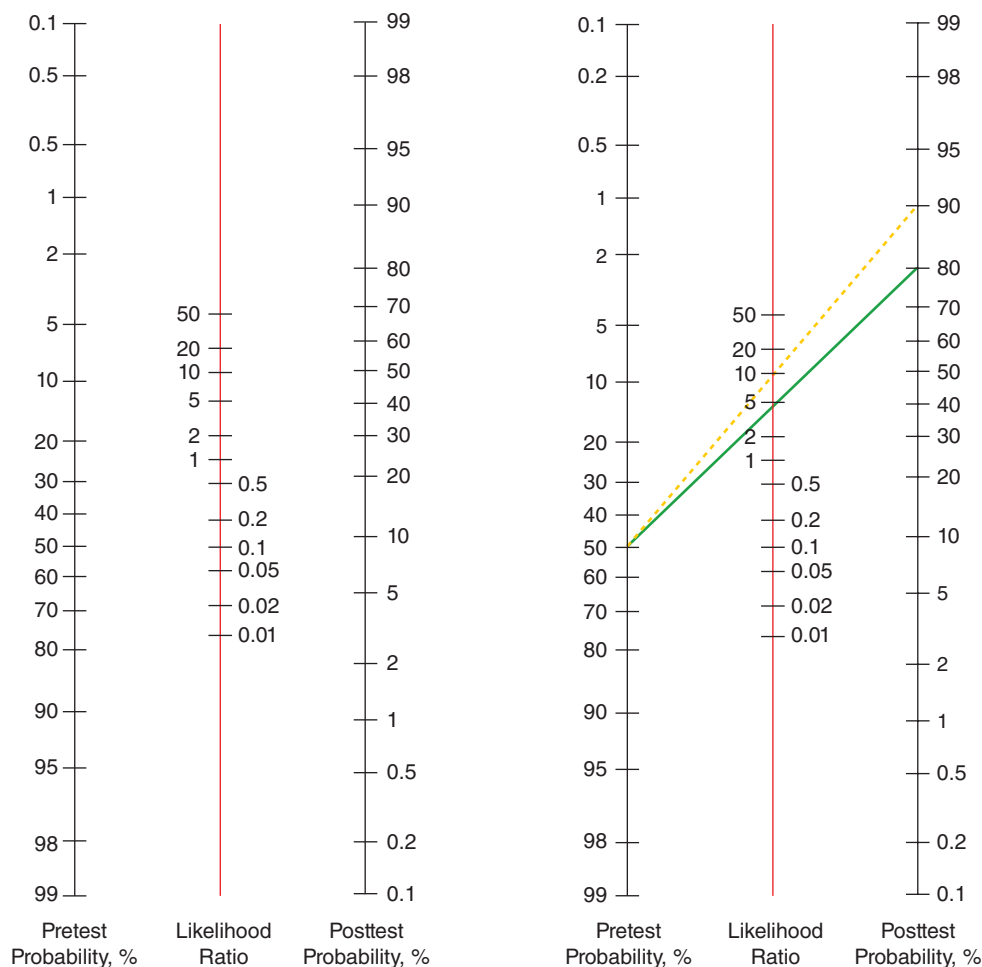


FIGURE 3-2 Nomogram version of Bayes' rule used to predict the posttest probability of disease (right-hand scale) using the pretest probability of disease (left-hand scale) and the likelihood ratio for a positive test (middle scale). See text for information on calculation of likelihood ratios. To use, place a straight edge connecting the pretest probability and the likelihood ratio and read off the posttest probability. The right-hand part of the figure illustrates the value of a positive exercise treadmill test (likelihood ratio 4, green line) and a positive exercise thallium single-photon emission computed tomography perfusion study (likelihood ratio 9, broken yellow line) in a patient with a pretest probability of coronary artery disease of 50%. (Adapted from *Centre for Evidence-Based Medicine: Likelihood ratios*. Available at <http://www.cebm.net/index.aspx?o=1043>.)

probability, a negative test may not rule out disease adequately if it is not sufficiently sensitive. Thus, the largest change in diagnostic likelihood following a test result occurs when the clinician is most uncertain (i.e., pretest probability between 30% and 70%). For example, if a patient has a pretest probability for CAD of 50%, a positive exercise treadmill test will move the posttest probability to 80% and a positive exercise SPECT perfusion test will move it to 90% (Fig. 3-2).

As presented above, Bayes' rule employs a number of important simplifications that should be considered. First, few tests have only positive or negative results, and many tests provide multiple outcomes (e.g., ST-segment depression and exercise duration with exercise testing). Although Bayes' rule can be adapted to this more detailed test result format, it is computationally more complex to do so. Similarly, when multiple tests are performed, the posttest probability may be used as the pretest probability to interpret the second test. However, this simplification assumes conditional independence—that is, that the results of the first test do not affect the likelihood of the second test result—and this is often not true.

Finally, it has long been asserted that sensitivity and specificity are prevalence-independent parameters of test accuracy, and many texts still make this statement. This statistically useful assumption, however, is clinically simplistic. A treadmill exercise test, for example, has a sensitivity in a population of patients with one-vessel CAD of around 30%, whereas its sensitivity in patients with severe three-vessel CAD approaches 80%. Thus, the best estimate of sensitivity to use in a particular decision may vary, depending on the severity of disease in the

local population. A hospitalized, symptomatic, or referral population typically has a higher prevalence of disease and, in particular, a higher prevalence of more advanced disease than does an outpatient population. Consequently, test sensitivity will likely be higher in hospitalized patients, and test specificity higher in outpatients.

STATISTICAL PREDICTION MODELS

Bayes' rule, while illustrative as presented above, provides an unrealistically simple solution to most problems a clinician faces. Predictions based on multivariable statistical models, however, can more accurately address these more complex problems by accounting for specific patient characteristics. In particular, these models explicitly account for multiple possibly overlapping pieces of patient-specific information and assign a relative weight to each on the basis of its unique contribution to the prediction in question. For example, a logistic regression model to predict the probability of CAD considers all the relevant independent factors from the clinical examination and diagnostic testing and their significance instead of the limited data that clinicians can manage in their heads or with Bayes' rule. However, despite this strength, prediction models are usually too complex computationally to use without a calculator or computer (although this limitation may be overcome once medicine is practiced from a fully computerized platform).

To date, only a handful of prediction models have been validated properly (for example, Wells criteria for pulmonary embolism) (Table 3-2). The importance of independent validation in a population

TABLE 3-2 WELLS CLINICAL PREDICTION RULE FOR PULMONARY EMBOLISM

Clinical Feature	Points
Clinical signs of deep vein thrombosis	3
Alternative diagnosis is less likely than pulmonary embolism	3
Heart rate >100 beats/min	1.5
Immobilization ≥ 3 days or surgery in previous 4 weeks	1.5
History of deep vein thrombosis or pulmonary embolism	1.5
Hemoptysis	1
Malignancy (with treatment within 6 months) or palliative	1
Interpretation	
Score >6.0	High
Score 2.0–6.0	Intermediate
Score <2.0	Low

separate from the one used to develop the model cannot be overstated. An unvalidated prediction model should be viewed with the skepticism appropriate for any new drug or medical device that has not had rigorous clinical trial testing.

When statistical models have been compared directly with expert clinicians, they have been found to be more consistent, as would be expected, but not significantly more accurate. Their biggest promise, then, may be in helping less-experienced clinicians identify critical discriminating patient characteristics and become more accurate in their predictions.

FORMAL DECISION SUPPORT TOOLS

DECISION SUPPORT SYSTEMS

Over the last 40 years, many attempts have been made to develop computer systems to aid clinical decision-making and patient management. Conceptually attractive because computers offer ready access to the vast information available to today's physicians, they may also support management decisions by making accurate predictions of outcome, simulating the whole decision process, or providing algorithmic guidance. Computer-based predictions using Bayesian or statistical regression models inform a clinical decision but do not actually reach a "conclusion" or "recommendation." Artificial intelligence systems attempt to simulate or replace human reasoning with a computer-based analogue. To date, such approaches have achieved only limited success. Reminder or protocol-directed systems do not make predictions but use existing algorithms, such as guidelines, to guide clinical practice. In general, however, decision support systems have had little impact on practice. Reminder systems, although not yet in widespread use, have shown the most promise, particularly in correcting drug dosing and promoting adherence to guidelines. Checklists, as used by pilots for example, have garnered recent support as an approach to avoid or reduce errors.

DECISION ANALYSIS

Compared with the decision support methods discussed above, decision analysis represents a prescriptive approach to decision making in the face of uncertainty. Its principal application is in complex decisions that involve substantial risk, abundant uncertainty, trade-offs in the outcomes emphasizing a role for preferences, or absence of evidence due to an idiosyncratic feature. For a public health example, [Fig. 3-3](#) displays a decision tree to evaluate strategies for screening for HIV infection. Infected individuals who are unaware of their illness cause up to 20,000 new cases of HIV infection annually in the United States, and about 40% of HIV-positive patients progress to AIDS within a year

of the initial diagnosis because of delayed diagnosis. Early identification offers the opportunity to prevent progression to AIDS through CD4 count and viral load monitoring and combination antiretroviral therapy and to reduce spread by reducing risky injection or sexual behaviors.

In 2003, the Centers for Disease Control and Prevention (CDC) proposed that routine universal HIV testing should be incorporated into standard adult medical care and, in part, cited a decision analysis model comparing HIV screening with usual care. Assuming a 1% prevalence of unidentified HIV infection in the population, routine screening of a cohort of 43-year-old men and women increased life expectancy by 5.5 days and lifetime costs by \$194 per person screened, yielding an incremental cost-effectiveness ratio for screening versus usual care of \$15,078 per quality-adjusted life-year (the additional cost to society to increase population health by 1 year of perfect health). Factors that influenced the results included assumptions about the effectiveness of behavior modification on subsequent sexual behavior, the benefits of early therapy for HIV infection, and the prevalence and incidence of HIV infection in the population targeted. This model, which required over 75 separate data points, provided novel insights into a public health problem in the absence of a randomized clinical trial and helped weigh the pros and cons of such a health policy recommendation. Although such models have been developed for selected clinical problems, their benefit and application to individual real-time clinical management have yet to be demonstrated.

DIAGNOSIS AS AN ELEMENT OF QUALITY OF CARE

High-quality medical care begins with accurate diagnosis. Recently, diagnostic errors have been re-envisioned: the old view was that they were caused by a lack of sufficient skill of an individual clinician; the new view is that they represent a quality of care patient-safety problem traceable to breakdowns in the health care system. Whether this conceptual shift will lead to new ways to improve diagnosis is uncertain. An annual rate of diagnostic errors of 10–15%, possibly leading to 40,000 deaths in the United States, is commonly cited, but these figures are imprecise.

Solutions to the "diagnostic errors as a system of care problem" have focused on system-level approaches, such as decision support and other tools integrated into electronic medical records. The use of checklists has been proposed as a means of reducing some of the cognitive errors discussed earlier in the chapter, such as premature closure. Although checklists have been shown to be useful in certain medical contexts, such as operating rooms and intensive care units, their value in preventing diagnostic errors that lead to patient adverse events remains to be shown.

EVIDENCE-BASED MEDICINE

Clinical medicine is defined traditionally as a practice combining medical knowledge (including scientific evidence), intuition, and judgment in the care of patients ([Chap. 1](#)). EBM updates this construct by placing much greater emphasis on the processes by which clinicians gain knowledge of the most up-to-date and relevant clinical research

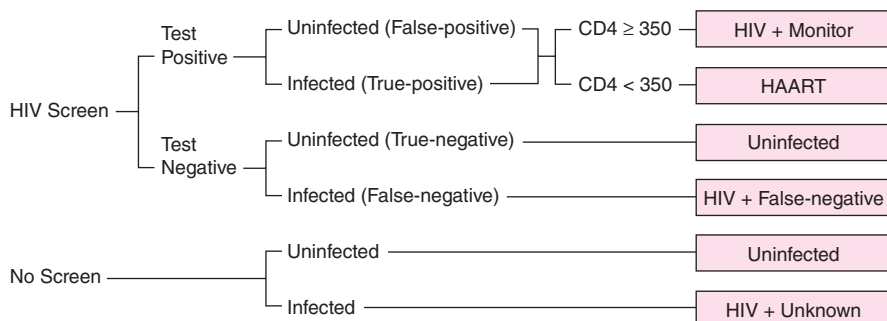


FIGURE 3-3 Basic structure of decision model used to evaluate strategies for screening for HIV in the general population. HAART, highly active antiretroviral therapy. (Provided courtesy of G. Sanders, with permission.)

to determine for themselves whether medical interventions alter the disease course and improve the length or quality of life. The meaning of practicing EBM becomes clearer through an examination of its four key steps:

1. Formulating the management question to be answered
2. Searching the literature and online databases for applicable research data
3. Appraising the evidence gathered with regard to its validity and relevance
4. Integrating this appraisal with knowledge about the unique aspects of the patient (including the patient's preferences about the possible outcomes)

The process of searching the world's research literature and appraising the quality and relevance of studies thus identified can be quite time-consuming and requires skills and training that most clinicians do not possess. Thus, identifying recent systematic overviews of the problem in question (Table 3-3) may offer the best starting point for most EBM searches.

Generally, the EBM tools listed in Table 3-3 provide access to research information in one of two forms. The first, primary research reports, is the original peer-reviewed research work that is published in medical journals and accessible through MEDLINE in abstract form. However, without training in using MEDLINE, quickly and efficiently locating reports that are on point in a huge sea of irrelevant or unhelpful citations may be difficult, and important studies could also be missed. The second form, systematic reviews, is the highest level of evidence in the hierarchy because it comprehensively summarizes the available evidence on a particular topic up to a certain date. To avoid the potential biases in review articles, predefined explicit search strategies and inclusion and exclusion criteria are used to find all of the relevant scientific research and grade its quality. The prototype for this kind of resource is the Cochrane Database of Systematic Reviews. When appropriate, a meta-analysis quantitatively summarizes the systematic review findings. The next two sections explicate the major types of clinical research reports available in the literature and the process of aggregating those data into meta-analyses.

SOURCES OF EVIDENCE: CLINICAL TRIALS AND REGISTRIES

The notion of learning from observation of patients is as old as medicine itself. Over the last 50 years, physicians' understanding of how best to turn raw observation into useful evidence has evolved considerably. Case reports, personal anecdotal experience, and small single-center case series are now recognized as having severe limitations in validity and generalizability, and although they may generate hypotheses or be the first reports of adverse events, they have no role

in formulating modern standards of practice. The major tools used to develop reliable evidence consist of the randomized clinical trial and the large observational registry. A registry or database typically is focused on a disease or syndrome (e.g., cancer, CAD, heart failure), a clinical procedure (e.g., bone marrow transplantation, coronary revascularization), or an administrative process (e.g., claims data used for billing and reimbursement).

By definition, in observational data, the investigator does not control patient care. Carefully collected *prospective* observational data, however, can achieve a level of evidence quality approaching that of major clinical trial data. At the other end of the spectrum, data collected retrospectively (e.g., chart review) are limited in form and content to what previous observers recorded, which may not include the specific research data being sought, e.g., claims data. Advantages of observational data include the inclusion of a broader population as encountered in practice than is typically represented in clinical trials because of their restrictive inclusion and exclusion criteria. In addition, observational data provide primary evidence for research questions when a randomized trial cannot be performed. For example, it would be difficult to randomize patients to test diagnostic or therapeutic strategies that are unproven but widely accepted in practice, and it would be unethical to randomize based on sex, racial/ethnic group, socioeconomic status, or country of residence or to randomize patients to a potentially harmful intervention, such as smoking or deliberately overeating to develop obesity.

A well-done prospective observational study of a particular management strategy differs from a well-done randomized clinical trial most importantly by its lack of protection from treatment selection bias. The use of observational data to compare diagnostic or therapeutic strategies assumes that sufficient uncertainty exists in clinical practice to ensure that similar patients will be managed differently by different physicians. In short, the analysis assumes that a sufficient element of randomness (in the sense of disorder rather than in the formal statistical sense) exists in clinical management. In such cases, statistical models attempt to adjust for important imbalances to "level the playing field" so that a fair comparison among treatment options can be made. When management is clearly not random (e.g., all eligible left main CAD patients are referred for coronary bypass surgery), the problem may be too confounded (biased) for statistical correction, and observational data may not provide reliable evidence.

In general, the use of concurrent controls is vastly preferable to that of historical controls. For example, comparison of current surgical management of left main CAD with left main CAD patients treated medically during the 1970s (the last time these patients were routinely treated with medicine alone) would be extremely misleading because "medical therapy" has substantially improved in the interim.

TABLE 3-3 SELECTED TOOLS FOR FINDING THE EVIDENCE IN EVIDENCE-BASED MEDICINE (EBM)

Name	Description	Web Address	Availability
Evidence-Based Medicine Reviews	Comprehensive electronic database that combines and integrates: 1. The Cochrane Database of Systematic Reviews 2. ACP Journal Club 3. The Database of Abstracts of Reviews of Effectiveness	www.ovid.com	Subscription required. Available through medical center libraries and other institutions.
Cochrane Library	Collection of EBM databases, including The Cochrane Database of Systematic Reviews—full text articles reviewing specific health care topics.	www.cochrane.org	Subscription required. Abstracts of systematic reviews available free online. Some countries have funding to provide free access to all residents.
ACP Journal Club	Collection of summaries of original studies and systematic reviews. Published bimonthly. All data since 1991 available on website, updated yearly.	www.acpjc.org	Subscription required.
Clinical Evidence	Monthly updated directory of concise overviews of common clinical interventions.	www.clinicalevidence.com	Subscription required. Free access for United Kingdom and developing countries.
MEDLINE	National Library of Medicine database with citations back to 1966.	www.nlm.nih.gov	Free via Internet.

Randomized controlled clinical trials include the careful prospective design features of the best observational data studies but also include the use of random allocation of treatment. This design provides the best protection against measured and unmeasured confounding due to treatment selection bias (a major aspect of internal validity). However, the randomized trial may not have good external validity (generalizability) if the process of recruitment into the trial resulted in the exclusion of many patients seen in clinical practice.

Consumers of medical evidence need to be aware that randomized trials vary widely in their quality and applicability to practice. The process of designing such a trial often involves many compromises. For example, trials designed to gain U.S. Food and Drug Administration (FDA) approval for an investigational drug or device must fulfill regulatory requirements that may result in a trial population and design that differs substantially from what practicing clinicians would find most useful.

META-ANALYSIS

The Greek prefix *meta* signifies something at a later or higher stage of development. Meta-analysis is research that combines and summarizes the available evidence quantitatively. Although occasionally used to examine nonrandomized studies, meta-analysis is used most typically to summarize all randomized trials examining a particular therapy. Ideally, unpublished trials should be identified and included to avoid publication bias (i.e., missing “negative” trials that may not be published). Furthermore, the best meta-analyses obtain and analyze individual patient-level data from all trials rather than working only the summary data in published reports of each trial. Nonetheless, not all published meta-analyses yield reliable evidence for a particular problem, so their methodology should be scrutinized carefully to ensure proper study design and analysis. The results of a well-done meta-analysis are likely to be most persuasive if they include at least several large-scale, properly performed randomized trials. Meta-analysis can especially help detect benefits when individual trials are inadequately powered (e.g., the benefits of streptokinase thrombolytic therapy in acute MI demonstrated by ISIS-2 in 1988 were evident by the early 1970s through meta-analysis). However, in cases in which the available trials are small or poorly done, meta-analysis should not be viewed as a remedy for the deficiency in primary trial data.

Meta-analyses typically focus on summary measures of relative treatment benefit, such as odds ratios or relative risks. Clinicians also should examine what absolute risk reduction (ARR) can be expected from the therapy. A useful summary metric of absolute treatment benefit is the number needed to treat (NNT) to prevent one adverse outcome event (e.g., death, stroke). NNT is simply $1/ARR$. For example, if a hypothetical therapy reduced mortality rates over a 5-year follow-up by 33% (the relative treatment benefit) from 12% (control arm) to 8% (treatment arm), the ARR would be $12\% - 8\% = 4\%$, and the NNT would be $1/0.04$, or 25. Thus, it would be necessary to treat 25 patients for 5 years to prevent 1 death. If the hypothetical treatment was applied to a lower-risk population, say, with a 6% 5-year mortality, the 33% relative treatment benefit would reduce absolute mortality by 2% (from 6% to 4%), and the NNT for the same therapy in this lower-risk group of patients would be 50. Although not always made explicit, comparisons of NNT estimates from different studies should account for the duration of follow-up used to create each estimate.

CLINICAL PRACTICE GUIDELINES

According to the 1990 Institute of Medicine definition, clinical practice guidelines are “systematically developed statements to assist practitioner and patient decisions about appropriate health care for specific clinical circumstances.” This definition emphasizes several crucial features of modern guideline development. First, guidelines are created by using the tools of EBM. In particular, the core of the development process is a systematic literature search followed by a review of the relevant peer-reviewed literature. Second, guidelines usually are focused on a clinical disorder (e.g., adult diabetes, stable angina pectoris) or a health care intervention (e.g., cancer screening). Third, the primary objective of guidelines is to improve the quality of medical care by

identifying care practices that should be routinely implemented, based on high-quality evidence and high benefit-to-harm ratios for the interventions. Guidelines are intended to “assist” decision-making, not to define explicitly what decisions should be made in a particular situation, in part because evidence alone is never sufficient for clinical decision-making (e.g., deciding whether to intubate and administer antibiotics for pneumonia in a terminally ill individual, in an individual with dementia, or in an otherwise healthy 30-year-old mother).

Guidelines are narrative documents constructed by expert panels whose composition often is determined by interested professional organizations. These panels vary in the degree to which they represent all relevant stakeholders. The guideline documents consist of a series of specific management recommendations, a summary indication of the quantity and quality of evidence supporting each recommendation, an assessment of the benefit-to-harm ratio for the recommendation, and a narrative discussion of the recommendations. Many recommendations simply reflect the expert consensus of the guideline panel because literature-based evidence is absent. The final step in guideline construction is peer review, followed by a final revision in response to the critiques provided. To improve the reliability and trustworthiness of guidelines, the Institute of Medicine has made methodologic recommendations for guideline development.

Guidelines are closely tied to the process of quality improvement in medicine through their identification of evidence-based best practices. Such practices can be used as quality indicators. Examples include the proportion of acute MI patients who receive aspirin upon admission to a hospital and the proportion of heart failure patients with a depressed ejection fraction treated with an ACE inhibitor.

CONCLUSIONS

In this era of EBM, it is tempting to think that all the difficult decisions practitioners face have been or soon will be solved and digested into practice guidelines and computerized reminders. However, EBM provides practitioners with an ideal rather than a finished set of tools with which to manage patients. Moreover, even with such evidence, it is always worth remembering that the response to therapy of the “average” patient represented by the summary clinical trial outcomes may not be what can be expected for the specific patient sitting in front of a physician in the clinic or hospital. In addition, meta-analyses cannot generate evidence when there are no adequate randomized trials, and most of what clinicians confront in practice will never be thoroughly tested in a randomized trial. For the foreseeable future, excellent clinical reasoning skills, experience supplemented by well-designed quantitative tools, and a keen appreciation for the role of individual patient preferences in their health care will continue to be of paramount importance in the practice of clinical medicine.


4

Screening and Prevention of Disease

Katrina Armstrong, Gary J. Martin

A primary goal of health care is to prevent disease or detect it early enough that intervention will be more effective. Tremendous progress has been made toward this goal over the last 50 years. Screening tests are available for many common diseases and encompass biochemical (e.g., cholesterol, glucose), physiologic (e.g., blood pressure, growth curves), radiologic (e.g., mammogram, bone densitometry), and cytologic (e.g., Pap smear) approaches. Effective preventive interventions have resulted in dramatic declines in mortality from many diseases, particularly infections. Preventive interventions include counseling about risk behaviors, vaccinations, medications, and, in some relatively uncommon settings, surgery. Preventive services (including screening tests, preventive interventions, and counseling) are different than

other medical interventions because they are proactively administered to healthy individuals instead of in response to a symptom, sign, or diagnosis. Thus, the decision to recommend a screening test or preventive intervention requires a particularly high bar of evidence that testing and intervention are both practical and effective.

 Because population-based screening and prevention strategies must be extremely low risk to have an acceptable benefit-to-harm ratio, the ability to target individuals who are more likely to develop disease could enable the application of a wider set of potential approaches and increase efficiency. Currently, there are many types of data that can predict disease incidence in an asymptomatic individual. Genomic data have received the most attention to date, at least in part because mutations in high-penetrance genes have clear implications for preventive care (Chap. 84). Women with mutations in either *BRCA1* or *BRCA2*, the two major breast cancer susceptibility genes identified to date, have a markedly increased risk (5- to 20-fold) of breast and ovarian cancer. Screening and prevention recommendations include prophylactic oophorectomy and breast magnetic resonance imaging (MRI), both of which are considered to incur too much harm for women at average cancer risk. Some women opt for prophylactic mastectomy to dramatically reduce their breast cancer risk. Although the proportion of common disease explained by high-penetrance genes appears to be relatively small (5–10% of most diseases), mutations in rare, moderate-penetrance genes, and variants in low-penetrance genes, also contribute to the prediction of disease risk. The advent of affordable whole exome/whole genome sequencing is likely to speed the dissemination of these tests into clinical practice and may transform the delivery of preventive care.

Other forms of “omic” data also have the potential to provide important predictive information, including proteomics and metabolomics. These fields are earlier in development and have yet to move into clinical practice. Imaging and other clinical data may also be integrated into a risk-stratified paradigm as evidence grows about the predictive ability of these data and the feasibility of their collection. Of course, all of these data may also be helpful in predicting the risk of harms from screening or prevention, such as the risk of a false-positive mammogram. To the degree that this information can be incorporated into personalized screening and prevention strategies, it could also improve delivery and efficiency.

In addition to advances in risk prediction, there are several other factors that are likely to promote the importance of screening and prevention in the near term. New imaging modalities are being developed that promise to detect changes at the cellular and subcellular levels, greatly increasing the probability that early detection improves outcomes. The rapidly growing understanding of the biologic pathways underlying initiation and progression of many common diseases has the potential to transform the development of preventive interventions, including chemoprevention. Furthermore, screening and prevention offer the promise of both improving health and sparing the costs of disease treatment, an issue that has gained national attention with the continued growth in health care costs.

This chapter will review the basic principles of screening and prevention in the primary care setting. Recommendations for specific disorders such as cardiovascular disease, diabetes, and cancer are provided in the chapters dedicated to those topics.

BASIC PRINCIPLES OF SCREENING

The basic principles of screening populations for disease were published by the World Health Organization in 1968 (Table 4-1).

In general, screening is most effective when applied to relatively common disorders that carry a large disease burden (Table 4-2). The five leading causes of mortality in the United States are heart diseases, malignant neoplasms, accidents, cerebrovascular diseases, and chronic obstructive pulmonary disease. Thus, many screening strategies are targeted at these conditions. From a global health perspective, these conditions are priorities, but malaria, malnutrition, AIDS, tuberculosis, and violence also carry a heavy disease burden (Chap. 2).


 Having an effective treatment for early disease has proven challenging for some common diseases. For example, although Alzheimer’s disease is the sixth leading cause of death in the

TABLE 4-1 PRINCIPLES OF SCREENING

The condition should be an important health problem.
There should be a treatment for the condition.
Facilities for diagnosis and treatment should be available.
There should be a latent stage of the disease.
There should be a test or examination for the condition.
The test should be acceptable to the population.
The natural history of the disease should be adequately understood.
There should be an agreed policy on whom to treat.
The cost of finding a case should be balanced in relation to overall medical expenditure.

United States, there are no curative treatments and no evidence that early treatment improves outcomes. Lack of facilities for diagnosis and treatment is a particular challenge for developing countries and may change screening strategies, including the development of “see and treat” approaches such as those currently used for cervical cancer screening in some countries. A long latent or preclinical phase where early treatment increases the chance of cure is a hallmark of many cancers; for example, polypectomy prevents progression to colon cancer. Similarly, early identification of hypertension or hyperlipidemia allows therapeutic interventions that reduce the long-term risk of cardiovascular or cerebrovascular events. In contrast, lung cancer screening has historically proven more challenging because most tumors are not curable by the time they can be detected on a chest x-ray. However, the length of the preclinical phase also depends on the level of resolution of the screening test, and this situation changed with the development of chest computed tomography (CT). Low-dose chest CT scanning can detect tumors earlier and was recently demonstrated to reduce lung cancer mortality by 20% in individuals who had at least a 30-pack-year history of smoking. The short interval between the ability to detect disease on a screening test and the development of incurable disease also contributes to the limited effectiveness of mammography screening in reducing breast cancer mortality among premenopausal women. Similarly, the early detection of prostate cancer may not lead to a difference in the mortality rate because the disease is often indolent and competing morbidities, such as coronary artery disease, may ultimately cause mortality (Chap. 100). This uncertainty about the natural history is also reflected in the controversy about treatment of prostate cancer, further contributing to the challenge of screening in this disease. Finally, screening programs can incur significant economic costs that must be considered in the context of the available resources and alternative strategies for improving health outcomes.

METHODS OF MEASURING HEALTH BENEFITS

Because screening and preventive interventions are recommended to asymptomatic individuals, they are held to a high standard for demonstrating a favorable risk-benefit ratio before implementation. In general, the principles of evidence-based medicine apply to demonstrating the efficacy of screening tests and preventive interventions, where randomized controlled trials (RCTs) with mortality outcomes are the gold standard. However, because RCTs are often not feasible, observational studies, such as case-control designs, have been used to assess the effectiveness of some interventions such as colorectal cancer screening. For some strategies, such as cervical cancer screening, the only data available are ecologic data demonstrating dramatic declines in mortality.

TABLE 4-2 LIFETIME CUMULATIVE RISK

Breast cancer for women	10%
Colon cancer	6%
Cancer of the cervix for women ^a	2%
Domestic violence for women	Up to 15%
Hip fracture for white women	16%

^aAssuming an unscreened population.

Irrespective of the study design used to assess the effectiveness of screening, it is critical that disease incidence or mortality is the primary endpoint rather than length of disease survival. This is important because lead time bias and length time bias can create the appearance of an improvement in disease survival from a screening test when there is no actual effect. Lead time bias occurs because screening identifies a case before it would have presented clinically, thereby creating the perception that a patient lived longer after diagnosis simply by moving the date of diagnosis earlier rather than the date of death later. Length time bias occurs because screening is more likely to identify slowly progressive disease than rapidly progressive disease. Thus, within a fixed period of time, a screened population will have a greater proportion of these slowly progressive cases and will appear to have better disease survival than an unscreened population.

A variety of endpoints are used to assess the potential gain from screening and preventive interventions.

1. *The absolute and relative impact of screening on disease incidence or mortality.* The absolute difference in disease incidence or mortality between a screened and nonscreened group allows the comparison of size of the benefit across preventive services. A meta-analysis of Swedish mammography trials (ages 40–70) found that ~1.2 fewer women per 1000 would die from breast cancer if they were screened over a 12-year period. By comparison, ~3 lives per 1000 would be saved from colon cancer in a population (ages 50–75) screened with annual fecal occult blood testing (FOBT) over a 13-year period. Based on this analysis, colon cancer screening may actually save more women's lives than does mammography. However, the relative impact of FOBT (30% reduction in colon cancer death) is similar to the relative impact of mammography (14–32% reduction in breast cancer death), emphasizing the importance of both relative and absolute comparisons.
2. *The number of subjects screened to prevent disease or death in one individual.* The inverse of the absolute difference in mortality is the number of subjects who would need to be screened or receive a preventive intervention to prevent one death. For example, 731 women ages 65–69 would need to be screened by dual-energy x-ray absorptiometry (DEXA) (and treated appropriately) to prevent one hip fracture from osteoporosis.
3. *Increase in average life expectancy for a population.* Predicted increases in life expectancy for various screening and preventive interventions are listed in [Table 4-3](#). It should be noted, however, that the increase in life expectancy is an average that applies to a population, not to an individual. In reality, the vast majority of the population does not derive any benefit from a screening test or preventive intervention. A small subset of patients, however, will benefit greatly. For example, Pap smears do not benefit the 98% of women who never develop cancer of the cervix. However, for the 2% who would have developed cervical cancer, Pap smears may add as much as 25 years to their lives. Some studies suggest that a 1-month gain of life expectancy is a reasonable goal for a population-based screening or prevention strategy.

ASSESSING THE HARMS OF SCREENING AND PREVENTION

Just as with most aspects of medical care, screening and preventive interventions also incur the possibility of adverse outcomes. These adverse outcomes include side effects from preventive medications and vaccinations, false-positive screening tests, overdiagnosis of disease from screening tests, anxiety, radiation exposure from some screening tests, and discomfort from some interventions and screening tests. The risk of side effects from preventive medications is analogous to the use of medications in therapeutic settings and is considered in the Food and Drug Administration (FDA) approval process. Side effects from currently recommended vaccinations are primarily limited to discomfort and minor immune reactions. However, the concern about associations between vaccinations and serious adverse outcomes continues to limit the acceptance of many vaccinations despite the lack of data supporting the causal nature of these associations.

The possibility of a false-positive test occurs with nearly all screening tests, although the definition of what constitutes a false-positive

TABLE 4-3 ESTIMATED AVERAGE INCREASE IN LIFE EXPECTANCY FOR A POPULATION

Screening or Preventive Intervention	Average Increase
Mammography:	
Women, 40–50 years	0–5 days
Women, 50–70 years	1 month
Pap smears, age 18–65	2–3 months
Getting a 35-year-old smoker to quit	3–5 years
Beginning regular exercise for a 40-year-old man (30 min, 3 times a week)	9 months–2 years

result often varies across settings. For some tests such as screening mammography and screening chest CT, a false-positive result occurs when an abnormality is identified that is not malignant, requiring either a biopsy diagnosis or short-term follow-up. For other tests such as Pap smears, a false-positive result occurs because the test identifies a wide range of potentially pre-malignant states, only a small percentage of which would ever progress to an invasive cancer. This risk is closely tied to the risk of overdiagnosis in which the screening test identifies disease that would not have presented clinically in the patient's lifetime. Assessing the degree of overdiagnosis from a screening test is very difficult given the need for long-term follow-up of an unscreened population to determine the true incidence of disease over time. Recent estimates suggest that as much as 15–25% of breast cancers identified by mammography screening and 15–37% of prostate cancers identified by prostate-specific antigen testing may never have presented clinically. Screening tests also have the potential to create unwarranted anxiety, particularly in conjunction with false-positive findings. Although multiple studies have documented increased anxiety through the screening process, there are few data suggesting this anxiety has long-term adverse consequences, including subsequent screening behavior. Screening tests that involve radiation (e.g., mammography, chest CT) add to the cumulative radiation exposure for the screened individual. The absolute amount of radiation is very small from any of these tests, but the overall impact of repeated exposure from multiple sources is still being determined. Some preventive interventions (e.g., vaccinations) and screening tests (e.g., mammography) may lead to discomfort at the time of administration, but again, there is little evidence of long-term adverse consequences.

WEIGHING THE BENEFITS AND HARMS

The decision to implement a population-based screening and prevention strategy requires weighing the benefits and harms, including the economic impact of the strategy. The costs include not only the expense of the intervention but also time away from work, downstream costs from false-positive results or adverse events, and other potential harms. Cost-effectiveness is typically assessed by calculating the cost per year of life saved, with adjustment for the quality of life impact of different interventions and disease states (i.e., quality-adjusted life-year). Typically, strategies that cost <\$50,000 to \$100,000 per quality-adjusted year of life saved are considered “cost-effective” ([Chap. 3](#)).

The U.S. Preventive Services Task Force (USPSTF) is an independent panel of experts in preventive care that provides evidence-based recommendations for screening and preventive strategies based on an assessment of the benefit-to-harm ratio ([Tables 4-4 and 4-5](#)). Because there are multiple advisory organizations providing recommendations for preventive services, the agreement among the organizations varies across the different services. For example, all advisory groups support screening for hyperlipidemia and colorectal cancer, whereas consensus is lower for breast cancer screening among women in their 40s and almost nonexistent for prostate cancer screening. Because the guidelines are only updated periodically, differences across advisory organizations may also reflect the data that were available when the guideline was issued. For example, multiple organizations have recently issued recommendations supporting lung cancer screening among heavy smokers based on the results of the National Lung Screening Trial (NLST) published in 2011, whereas the USPSTF did not review lung cancer screening until 2014.

TABLE 4-4 SCREENING TESTS RECOMMENDED BY THE U.S. PREVENTIVE SERVICES TASK FORCE FOR AVERAGE-RISK ADULTS

Disease	Test	Population	Frequency	Chapter
Abdominal aortic aneurysm	Ultrasound	Men 65–75 who have ever smoked	Once	
Alcohol misuse	Alcohol Use Disorders Identification Test	>18	Unknown	467
Breast cancer	Mammography with or without clinical breast examination	Women 50–75	Every 2 years	
Cervical cancer	Pap smear	Women 21–65	Every 3 years	100
	Pap smear and HPV testing	Women 30–65	Every 5 years if HPV negative	
Chlamydia/gonorrhea	Nucleic acid amplification test on urine or cervical swab	Sexually active women <25	Unknown	213
Colorectal cancer	Fecal occult blood testing	50–75	Every year	100, 110
	Sigmoidoscopy	50–75	Every 5 years	
	Colonoscopy	50–75	Every 10 years	
Depression	Screening questions	All adults	Periodically	
Diabetes	Fasting blood glucose	Adults with hypertension	Every 3 years	417
Hepatitis C	Anti-HCV antibody followed by confirmatory PCR	Adults born between 1945 and 1965	Once	
HIV	Reactive immunoassay or rapid HIV followed by confirmatory test	15–65	Once	
Hyperlipidemia	Cholesterol	Men >35	Every 5 years	291e
		Women >45	Every 5 years	
Hypertension	Blood pressure	All adults	Periodically	298
Intimate partner violence	Screening questions	Women of childbearing age	Unknown	
Obesity	Body mass index	All adults	Unknown	
Osteoporosis	DEXA	Women >65 or >60 with risk factors	Unknown	425

Abbreviations: DEXA, dual-energy x-ray absorptiometry; HCV, hepatitis C virus; HPV, human papillomavirus; PCR, polymerase chain reaction.

Source: Adapted from the U.S. Preventive Services Task Force 2013. <http://www.uspreventiveservicestaskforce.org/adultrec.htm>.

For many screening tests and preventive interventions, the balance of benefits and harms may be uncertain for the average-risk population but more favorable for individuals at higher risk for disease. Although age is the most commonly used risk factor for determining screening and prevention recommendations, the USPSTF also recommends some screening tests in populations with other risk factors for the disease (e.g., syphilis). In addition, being at increased risk for the disease often supports initiating screening at an earlier age than that recommended for the average-risk population. For example, when there is a significant family history of breast or colon cancer, it is prudent to initiate screening 10 years before the age at which the youngest family member was diagnosed with cancer.

Although informed consent is important for all aspects of medical care, shared decision-making may be a particularly important approach to decisions about preventive services when the benefit-to-harm ratio is uncertain for a specific population. For example, many expert groups, including the USPSTF, recommend an individualized discussion about prostate cancer screening, because the decision-making process is complex and relies heavily on personal issues. Some men may decline screening, whereas others may be more willing to accept the risks of an early detection strategy. Recent analysis suggests that many men may be better off not screening for prostate cancer because watchful waiting was the preferred strategy when quality-adjusted life-years were considered. Another example of

TABLE 4-5 PREVENTIVE INTERVENTIONS RECOMMENDED FOR AVERAGE-RISK ADULTS

Intervention	Disease	Population	Frequency	Chapter
Adult immunization				148, 149
Tetanus-diphtheria		>18	Every 10 years	
Varicella		Susceptibles only, >18	Two doses	
Measles-mumps-rubella		Women, childbearing age	One dose	
Pneumococcal		>65	One dose	
Influenza		>50	Yearly	
Human papillomavirus		Boys >21	If not done prior	
		Girls >26		
Zoster		>60	Once	
Chemoprevention				
Aspirin	Cardiovascular disease	Men 45–79		
		Women 55–79		
Folic acid	Neural tube defects in baby	Women planning or capable of pregnancy		
Tamoxifen/raloxifene	Breast cancer	Women at high risk for breast cancer		
Vitamin D	Fracture/falls	>65 at increased risk for falls		

shared decision-making involves the choice of techniques for colon cancer screening (Chap. 100). In controlled studies, the use of annual FOBT reduces colon cancer deaths by 15–30%. Flexible sigmoidoscopy reduces colon cancer deaths by ~60%. Colonoscopy offers the same benefit as or greater benefit than flexible sigmoidoscopy, but its use incurs additional costs and risks. These screening procedures have not been compared directly in the same population, but the estimated cost to society is similar: \$10,000–25,000 per year of life saved. Thus, although one patient may prefer the ease of preparation, less time disruption, and the lower risk of flexible sigmoidoscopy, others may prefer the sedation and thoroughness of colonoscopy.

COUNSELING ON HEALTHY BEHAVIORS

In considering the impact of preventive services, it is important to recognize that tobacco and alcohol use, diet, and exercise constitute the vast majority of factors that influence preventable deaths in developed countries. Perhaps the single greatest preventive health care measure is to help patients quit smoking (Chap. 470). However, efforts in these areas frequently involve behavior changes (e.g., weight loss, exercise, seat belts) or the management of addictive conditions (e.g., tobacco and alcohol use) that are often recalcitrant to intervention. Although these are challenging problems, evidence strongly supports the role of counseling by health care providers (Table 4-6) in effecting health behavior change. Educational campaigns, public policy changes, and community-based interventions have also proven to be important parts of a strategy for addressing these factors in some settings. Although the USPSTF found that the evidence was conclusive to recommend a relatively small set of counseling activities, counseling in areas such as physical activity and injury prevention (including seat belts and bicycle and motorcycle helmets) has become a routine part of primary care practice.

IMPLEMENTING DISEASE PREVENTION AND SCREENING

The implementation of disease prevention and screening strategies in practice is challenging. A number of techniques can assist physicians with the delivery of these services. An appropriately configured electronic health record can provide reminder systems that make it easier for physicians to track and meet guidelines. Some systems give patients secure access to their medical records, providing an additional means

TABLE 4-6 PREVENTIVE COUNSELING RECOMMENDED BY THE USPSTF

Topic	Chapter Reference
Alcohol and drug use	467, 468e
Genetic counseling for <i>BRCA1/2</i> testing among women at increased risk for deleterious mutations	
Nutrition and diet	
Sexually transmitted infections	163, 226
Sun exposure	75
Tobacco use	470

to enhance adherence to routine screening. Systems that provide nurses and other staff with standing orders are effective for smoking prevention and immunizations. The Agency for Healthcare Research and Quality and the Centers for Disease Control and Prevention have developed flow sheets and electronic tools as part of their “Put Prevention into Practice” program (<http://www.uspreventiveservicestaskforce.org/tools.htm>). Many of these tools use age categories to help guide implementation. Age-specific recommendations for screening and counseling are summarized in Table 4-7.

Many patients see a physician for ongoing care of chronic illnesses, and this visit provides an opportunity to include a “measure of prevention” for other health problems. For example, a patient seen for management of hypertension or diabetes can have breast cancer screening incorporated into one visit and a discussion about colon cancer screening at the next visit. Other patients may respond more favorably to a clearly defined visit that addresses all relevant screening and prevention interventions. Because of age or comorbidities, it may be appropriate with some patients to abandon certain screening and prevention activities, although there are fewer data about when to “sunset” these services. For many screening tests, the benefit of screening does not accrue until 5 to 10 years of follow-up, and there are generally few data to support continuing screening for most diseases past age 75. In addition, for patients with advanced diseases and limited life expectancy, there is considerable benefit from shifting the focus from screening procedures to the conditions and interventions more likely to affect quality and length of life.

TABLE 4-7 AGE-SPECIFIC CAUSES OF MORTALITY AND CORRESPONDING PREVENTIVE OPTIONS

Age Group	Leading Causes of Age-Specific Mortality	Screening Prevention Interventions to Consider for Each Specific Population
15–24	<ol style="list-style-type: none"> 1. Accident 2. Homicide 3. Suicide 4. Malignancy 5. Heart disease 	<ul style="list-style-type: none"> • Counseling on routine seat belt use, bicycle/motorcycle/ATV helmets (1) • Counseling on diet and exercise (5) • Discuss dangers of alcohol use while driving, swimming, boating (1) • Assess and update vaccination status (tetanus, diphtheria, hepatitis B, MMR, rubella, varicella, meningitis, HPV) • Ask about gun use and/or gun possession (2,3) • Assess for substance abuse history including alcohol (2,3) • Screen for domestic violence (2,3) • Screen for depression and/or suicidal/homicidal ideation (2,3) • Pap smear for cervical cancer screening after age 21 (4) • Discuss skin, breast awareness, and testicular self-exams (4) • Recommend UV light avoidance and regular sunscreen use (4) • Measurement of blood pressure, height, weight, and body mass index (5) • Discuss health risks of tobacco use, consider emphasis on cosmetic and economic issues to improve quit rates for younger smokers (4,5) • Chlamydia and gonorrhea screening and contraceptive counseling for sexually active females, discuss STD prevention • Hepatitis B, and syphilis testing if there is high-risk sexual behavior(s) or any prior history of sexually transmitted disease • HIV testing • Continue annual influenza vaccination

(Continued)

TABLE 4-7 AGE-SPECIFIC CAUSES OF MORTALITY AND CORRESPONDING PREVENTIVE OPTIONS (CONTINUED)

Age Group	Leading Causes of Age-Specific Mortality	Screening Prevention Interventions to Consider for Each Specific Population
25–44	<ol style="list-style-type: none"> 1. Accident 2. Malignancy 3. Heart disease 4. Suicide 5. Homicide 6. HIV 	<p><i>As above plus consider the following:</i></p> <ul style="list-style-type: none"> • Readdress smoking status, encourage cessation at every visit (2,3) • Obtain detailed family history of malignancies and begin early screening/prevention program if patient is at significant increased risk (2) • Assess all cardiac risk factors (including screening for diabetes and hyperlipidemia) and consider primary prevention with aspirin for patients at >3% 5-year risk of a vascular event (3) • Assess for chronic alcohol abuse, risk factors for viral hepatitis, or other risks for development of chronic liver disease • Consider individualized breast cancer screening with mammography at age 40 (2)
45–64	<ol style="list-style-type: none"> 1. Malignancy 2. Heart disease 3. Accident 4. Diabetes mellitus 5. Cerebrovascular disease 6. Chronic lower respiratory disease 7. Chronic liver disease and cirrhosis 8. Suicide 	<ul style="list-style-type: none"> • Consider prostate cancer screen with annual PSA and digital rectal exam at age 50 (or possibly earlier in African Americans or patients with family history) (1) • Begin colorectal cancer screening at age 50 with fecal occult blood testing, flexible sigmoidoscopy, or colonoscopy (1) • Reassess and update vaccination status at age 50 and vaccinate all smokers against <i>S. pneumoniae</i> at age 50 (6) • Consider screening for coronary disease in higher-risk patients (2,5) • Consider screening for hepatitis C in adults born between 1945 and 1965 (7) • Zoster vaccination at age 60 • Begin mammography screening by age 50
≥65	<ol style="list-style-type: none"> 1. Heart disease 2. Malignancy 3. Cerebrovascular disease 4. Chronic lower respiratory disease 5. Alzheimer's disease 6. Influenza and pneumonia 7. Diabetes mellitus 8. Kidney disease 9. Accidents 10. Septicemia 	<p><i>As above plus consider the following:</i></p> <ul style="list-style-type: none"> • Readdress smoking status, encourage cessation at every visit (1,2,3,4) • One-time ultrasound for AAA in men 65–75 who have ever smoked • Consider pulmonary function testing for all long-term smokers to assess for development of chronic obstructive pulmonary disease (4,6) • Screen all postmenopausal women (and all men with risk factors) for osteoporosis • Continue annual influenza vaccination and vaccinate against <i>S. pneumoniae</i> at age 65 (4, 6) • Screen for dementia and depression (5) • Screen for visual and hearing problems, home safety issues, and elder abuse (9)

Note: The numbers in parentheses refer to areas of risk in the mortality column affected by the specified intervention.

Abbreviations: AAA, abdominal aortic aneurysm; ATV, all-terrain vehicle; HPV, human papillomavirus; MMR, measles-mumps-rubella; PSA, prostate-specific antigen; STD, sexually transmitted disease; UV, ultraviolet.

5

Principles of Clinical Pharmacology

Dan M. Roden

Drugs are the cornerstone of modern therapeutics. Nevertheless, it is well recognized among physicians and in the lay community that the outcome of drug therapy varies widely among individuals. While this variability has been perceived as an unpredictable, and therefore inevitable, accompaniment of drug therapy, this is not the case. The goal of this chapter is to describe the principles of clinical pharmacology that can be used for the safe and optimal use of available and new drugs.

Drugs interact with specific target molecules to produce their beneficial and adverse effects. The chain of events between administration of a drug and production of these effects in the body can be divided into two components, both of which contribute to variability in drug actions. The first component comprises the processes that determine drug delivery to, and removal from, molecular targets. The resulting description of the relationship between drug concentration and time

is termed *pharmacokinetics*. The second component of variability in drug action comprises the processes that determine variability in drug actions despite equivalent drug delivery to effector drug sites. This description of the relationship between drug concentration and effect is termed *pharmacodynamics*. As discussed further below, pharmacodynamic variability can arise as a result of variability in function of the target molecule itself or of variability in the broad biologic context in which the drug-target interaction occurs to achieve drug effects.

Two important goals of the discipline of clinical pharmacology are (1) to provide a description of conditions under which drug actions vary among human subjects; and (2) to determine mechanisms underlying this variability, with the goal of improving therapy with available drugs as well as pointing to new drug mechanisms that may be effective in the treatment of human disease. The first steps in the discipline were empirical descriptions of the influence of disease on drug actions and of individuals or families with unusual sensitivities to adverse drug effects. These important descriptive findings are now being replaced by an understanding of the molecular mechanisms underlying variability in drug actions. Thus, the effects of disease, drug coadministration, or familial factors in modulating drug action can now be reinterpreted as variability in expression or function of specific genes whose products

determine pharmacokinetics and pharmacodynamics. Nevertheless, it is often the personal interaction of the patient with the physician or other health care provider that first identifies unusual variability in drug actions; maintained alertness to unusual drug responses continues to be a key component of improving drug safety.

Unusual drug responses, segregating in families, have been recognized for decades and initially defined the field of *pharmacogenetics*. Now, with an increasing appreciation of common and rare polymorphisms across the human genome, comes the opportunity to reinterpret descriptive mechanisms of variability in drug action as a consequence of specific DNA variants, or sets of variants, among individuals. This approach defines the field of *pharmacogenomics*, which may hold the opportunity of allowing practitioners to integrate a molecular understanding of the basis of disease with an individual's genomic makeup to prescribe personalized, highly effective, and safe therapies.

IDENTIFYING DRUG TARGETS

Drug therapy is an ancient feature of human culture. The first treatments were plant extracts discovered empirically to be effective for indications like fever, pain, or breathlessness. This symptom-based empiric approach to drug development was supplanted in the twentieth century by identification of compounds targeting more fundamental biologic processes such as bacterial growth or elevated blood pressure; the term “magic bullet,” coined by Paul Ehrlich to describe the search for effective compounds for syphilis, captures the essence of the hope that understanding basic biologic processes will lead to highly effective new therapies. An integral step in modern drug development follows identification of a chemical lead with biologic activity with increasingly sophisticated medicinal chemistry-based structural modifications to develop compounds with specificity for the chosen target, lack of “off-target” effects, and pharmacokinetic properties suitable for human use (e.g., consistent bioavailability, long elimination half-life, no high-risk pharmacokinetic features described further below).

A common starting point for contemporary drug development is basic biologic discovery that implicates potential target molecules: examples of such target molecules include HMG-CoA reductase or the *BRAF V600E* mutation in many malignant melanomas. The development of compounds targeting these molecules has not only revolutionized treatment for diseases such as hypercholesterolemia or malignant melanoma, but has also revealed new biologic features of disease. Thus, for example, initial spectacular successes with vemurafenib (which targets *BRAF V600E*) were followed by near-universal tumor relapse, strongly suggesting that inhibition of this pathway alone would be insufficient for tumor control. This reasoning, in turn, supports a view that many complex diseases will not lend themselves to cure by targeting a single magic bullet, but rather single drugs or combinations will need to attack multiple pathways whose perturbation results in disease. The use of combination therapy in settings such as hypertension, tuberculosis, HIV infection, and many cancers highlights potential for such a “systems biology” view of drug therapy.

GLOBAL CONSIDERATIONS



It is true across all cultures and diseases that factors such as compliance, genetic variants affecting pharmacokinetics or pharmacodynamics, and drug interactions contribute to drug responses. In addition, culture- or ancestry-specific factors play a role. For example, the frequency of specific genetic variants modulating drug responses often varies by ancestry, as discussed later. Cost issues or cultural factors may determine the likelihood that specific drugs, drug combinations, or over-the-counter (OTC) remedies are prescribed. The broad principles of clinical pharmacology enunciated here can be used to analyze the mechanisms underlying successful or unsuccessful therapy with any drug.

INDICATIONS FOR DRUG THERAPY: RISK VERSUS BENEFIT

It is self-evident that the benefits of drug therapy should outweigh the risks. Benefits fall into two broad categories: those designed to alleviate

a symptom and those designed to prolong useful life. An increasing emphasis on the principles of evidence-based medicine and techniques such as large clinical trials and meta-analyses have defined benefits of drug therapy in broad patient populations. Establishing the balance between risk and benefit is not always simple. An increasing body of evidence supports the idea, with which practitioners are very familiar, that individual patients may display responses that are not expected from large population studies and often have comorbidities that typically exclude them from large clinical trials. In addition, therapies that provide symptomatic benefits but shorten life may be entertained in patients with serious and highly symptomatic diseases such as heart failure or cancer. These considerations illustrate the continuing, highly personal nature of the relationship between the prescriber and the patient.

Adverse Effects Some adverse effects are so common and so readily associated with drug therapy that they are identified very early during clinical use of a drug. By contrast, serious adverse effects may be sufficiently uncommon that they escape detection for many years after a drug begins to be widely used. The issue of how to identify rare but serious adverse effects (that can profoundly affect the benefit-risk perception in an individual patient) has not been satisfactorily resolved. Potential approaches range from an increased understanding of the molecular and genetic basis of variability in drug actions to expanded postmarketing surveillance mechanisms. None of these have been completely effective, so practitioners must be continuously vigilant to the possibility that unusual symptoms may be related to specific drugs, or combinations of drugs, that their patients receive.

Therapeutic Index Beneficial and adverse reactions to drug therapy can be described by a series of dose-response relations (Fig. 5-1). Well-tolerated drugs demonstrate a wide margin, termed the *therapeutic ratio*, *therapeutic index*, or *therapeutic window*, between the doses required to produce a therapeutic effect and those producing toxicity. In cases where there is a similar relationship between plasma drug concentration and effects, monitoring plasma concentrations can be a highly effective aid in managing drug therapy by enabling concentrations to be maintained above the minimum required to produce an effect and below the concentration range likely to produce toxicity. Such monitoring has been widely used to guide therapy with specific agents, such as certain antiarrhythmics, anticonvulsants, and antibiotics. Many of the principles in clinical pharmacology and examples

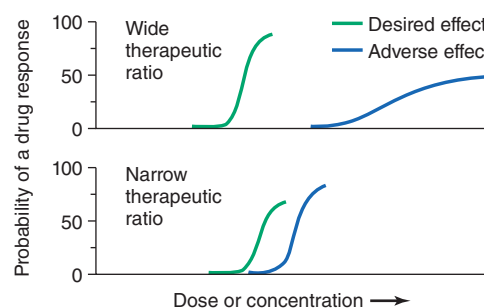


FIGURE 5-1 The concept of a therapeutic ratio. Each panel illustrates the relationship between increasing dose and cumulative probability of a desired or adverse drug effect. **Top.** A drug with a wide therapeutic ratio, i.e., a wide separation of the two curves. **Bottom.** A drug with a narrow therapeutic ratio; here, the likelihood of adverse effects at therapeutic doses is increased because the curves are not well separated. Further, a steep dose-response curve for adverse effects is especially undesirable, as it implies that even small dosage increments may sharply increase the likelihood of toxicity. When there is a definable relationship between drug concentration (usually measured in plasma) and desirable and adverse effect curves, concentration may be substituted on the abscissa. Note that not all patients necessarily demonstrate a therapeutic response (or adverse effect) at any dose, and that some effects (notably some adverse effects) may occur in a dose-independent fashion.

outlined below, which can be applied broadly to therapeutics, have been developed in these arenas.

PRINCIPLES OF PHARMACOKINETICS

The processes of absorption, distribution, metabolism, and excretion—collectively termed *drug disposition*—determine the concentration of drug delivered to target effector molecules.

ABSORPTION AND BIOAVAILABILITY

When a drug is administered orally, subcutaneously, intramuscularly, rectally, sublingually, or directly into desired sites of action, the amount of drug actually entering the systemic circulation may be less than with the intravenous route (Fig. 5-2A). The fraction of drug available to the systemic circulation by other routes is termed *bioavailability*. Bioavailability may be <100% for two main reasons: (1) absorption is reduced, or (2) the drug undergoes metabolism or elimination prior to entering the systemic circulation. Occasionally, the administered drug formulation is inconsistent or has degraded with time; for example, the anticoagulant dabigatran degrades rapidly (over weeks) once exposed to air, so the amount administered may be less than prescribed.

When a drug is administered by a nonintravenous route, the peak concentration occurs later and is lower than after the same dose given by rapid intravenous injection, reflecting absorption from the site of administration (Fig. 5-2). The extent of absorption may be reduced because a drug is incompletely released from its dosage form, undergoes destruction at its site of administration, or has physicochemical properties such as insolubility that prevent complete absorption from its site of administration. Slow absorption rates are deliberately designed into “slow-release” or “sustained-release” drug formulations in order to minimize variation in plasma concentrations during the interval between doses.

“First-Pass” Effect When a drug is administered orally, it must traverse the intestinal epithelium, the portal venous system, and the liver prior

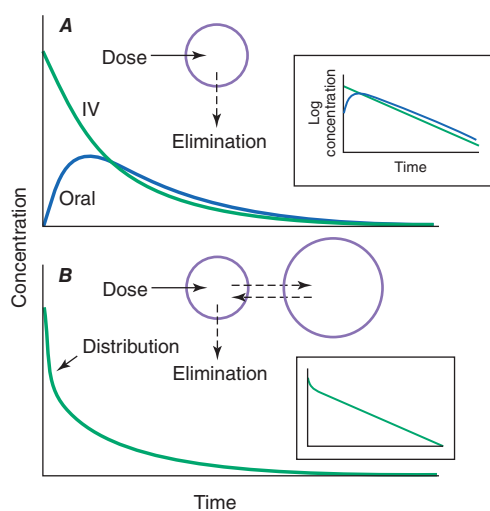


FIGURE 5-2 Idealized time-plasma concentration curves after a single dose of drug. **A.** The time course of drug concentration after an instantaneous IV bolus or an oral dose in the one-compartment model shown. The area under the time-concentration curve is clearly less with the oral drug than the IV, indicating incomplete bioavailability. Note that despite this incomplete bioavailability, concentration after the oral dose can be higher than after the IV dose at some time points. The inset shows that the decline of concentrations over time is linear on a log-linear plot, characteristic of first-order elimination, and that oral and IV drugs have the same elimination (parallel) time course. **B.** The decline of central compartment concentration when drug is distributed both to and from a peripheral compartment and eliminated from the central compartment. The rapid initial decline of concentration reflects not drug elimination but distribution.

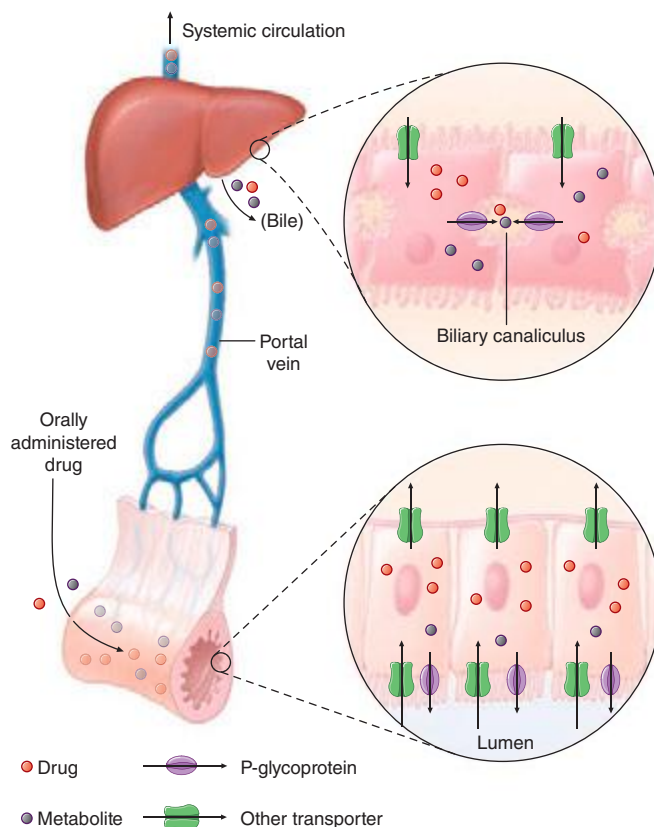


FIGURE 5-3 Mechanism of presystemic clearance. After drug enters the enterocyte, it can undergo metabolism, excretion into the intestinal lumen, or transport into the portal vein. Similarly, the hepatocyte may accomplish metabolism and biliary excretion prior to the entry of drug and metabolites to the systemic circulation. (Adapted by permission from DM Roden, in DP Zipes, J Jalife [eds]: *Cardiac Electrophysiology: From Cell to Bedside*, 4th ed. Philadelphia, Saunders, 2003. Copyright 2003 with permission from Elsevier.)

to entering the systemic circulation (Fig. 5-3). Once a drug enters the enterocyte, it may undergo metabolism, be transported into the portal vein, or be excreted back into the intestinal lumen. Both excretion into the intestinal lumen and metabolism decrease systemic bioavailability. Once a drug passes this enterocyte barrier, it may also be taken up into the hepatocyte, where bioavailability can be further limited by metabolism or excretion into the bile. This elimination in intestine and liver, which reduces the amount of drug delivered to the systemic circulation, is termed *presystemic elimination*, *presystemic extraction*, or *first-pass elimination*.

Drug movement across the membrane of any cell, including enterocytes and hepatocytes, is a combination of passive diffusion and active transport, mediated by specific drug uptake and efflux molecules. One widely studied drug transport molecule is P-glycoprotein, the product of the *MDR1* gene. P-glycoprotein is expressed on the apical aspect of the enterocyte and on the canalicular aspect of the hepatocyte (Fig. 5-3). In both locations, it serves as an efflux pump, limiting availability of drug to the systemic circulation. P-glycoprotein-mediated drug efflux from cerebral capillaries limits drug brain penetration and is an important component of the blood-brain barrier.

Drug metabolism generates compounds that are usually more polar and, hence, more readily excreted than parent drug. Metabolism takes place predominantly in the liver but can occur at other sites such as kidney, intestinal epithelium, lung, and plasma. “Phase I” metabolism involves chemical modification, most often oxidation accomplished by members of the cytochrome P450 (CYP) monooxygenase superfamily. CYPs that are especially important for drug metabolism are presented in Table 5-1, and each drug may be a substrate for one or more of these enzymes. “Phase II” metabolism involves conjugation of specific endogenous compounds to drugs or their metabolites. The enzymes

TABLE 5-1 MOLECULAR PATHWAYS MEDIATING DRUG DISPOSITION

Molecule	Substrates ^a	Inhibitors ^a
CYP3A	Calcium channel blockers Antiarrhythmics (lidocaine, quinidine, mexiletine) HMG-CoA reductase inhibitors ("statins"; see text) Cyclosporine, tacrolimus Indinavir, saquinavir, ritonavir	Amiodarone Ketoconazole, itraconazole Erythromycin, clarithromycin Ritonavir
CYP2D6 ^b	Timolol, metoprolol, carvedilol Phenformin Codeine Propafenone, flecainide Tricyclic antidepressants Fluoxetine, paroxetine	Quinidine (even at ultra-low doses) Tricyclic antidepressants Fluoxetine, paroxetine
CYP2C9 ^b	Warfarin Phenytoin Glipizide Losartan	Amiodarone Fluconazole Phenytoin
CYP2C19 ^b	Omeprazole Mephenytoin Clopidogrel	Omeprazole
CYP2B6 ^b	Efavirenz	
Thiopurine S-methyltransferase ^b	6-Mercaptopurine, azathioprine	
N-acetyltransferase ^b	Isoniazid Procainamide Hydralazine Some sulfonamides	
UGT1A1 ^b	Irinotecan	
Pseudocholinesterase ^b	Succinylcholine	
P-glycoprotein	Digoxin HIV protease inhibitors Many CYP3A substrates	Quinidine Amiodarone Verapamil Cyclosporine Itraconazole Erythromycin
SLCO1B1 ^b	Simvastatins and some other statins	

^aInhibitors affect the molecular pathway, and thus may affect substrate. ^bClinically important genetic variants described; see Table 5-2.

Note: A listing of CYP substrates, inhibitors, and inducers is maintained at <http://medicine.jupui.edu/flockhart/table.htm>.

that accomplish phase II reactions include glucuronyl-, acetyl-, sulfo-, and methyltransferases. Drug metabolites may exert important pharmacologic activity, as discussed further below.

Clinical Implications of Altered Bioavailability Some drugs undergo near-complete presystemic metabolism and, thus, cannot be administered orally. Nitroglycerin cannot be used orally because it is completely extracted prior to reaching the systemic circulation. The drug is, therefore, used by the sublingual or transdermal routes, which bypass presystemic metabolism.

Some drugs with very extensive presystemic metabolism can still be administered by the oral route, using much higher doses than those required intravenously. Thus, a typical intravenous dose of verapamil is 1–5 mg, compared to the usual single oral dose of 40–120 mg. Administration of low-dose aspirin can result in exposure of cyclooxygenase in platelets in the portal vein to the drug, but systemic sparing

because of first-pass aspirin deacylation in the liver. This is an example of presystemic metabolism being exploited to therapeutic advantage.

PHARMACOKINETIC CONCEPTS

Most pharmacokinetic processes, such as elimination, are first-order; that is, the rate of the process depends on the amount of drug present. Elimination can occasionally be zero-order (fixed amount eliminated per unit time), and this can be clinically important (see "Principles of Dose Selection"). In the simplest pharmacokinetic model (Fig. 5-2A), a drug bolus (D) is administered instantaneously to a central compartment, from which drug elimination occurs as a first-order process. Occasionally, central and other compartments correspond to physiologic spaces (e.g., plasma volume), whereas in others they are simply mathematical functions used to describe drug disposition. The first-order nature of drug elimination leads directly to the relationship describing drug concentration (C) at any time (t) following the bolus:

$$C = \frac{D}{V_c} \cdot e^{(-0.69t/t_{1/2})}$$

where V_c is the volume of the compartment into which drug is delivered and $t_{1/2}$ is elimination half-life. As a consequence of this relationship, a plot of the logarithm of concentration versus time is a straight line (Fig. 5-2A, inset). *Half-life* is the time required for 50% of a first-order process to be complete. Thus, 50% of drug elimination is achieved after one drug-elimination half-life, 75% after two, 87.5% after three, etc. In practice, first-order processes such as elimination are near-complete after four–five half-lives.

In some cases, drug is removed from the central compartment not only by elimination but also by distribution into peripheral compartments. In this case, the plot of plasma concentration versus time after a bolus may demonstrate two (or more) exponential components (Fig. 5-2B). In general, the initial rapid drop in drug concentration represents not elimination but drug distribution into and out of peripheral tissues (also first-order processes), while the slower component represents drug elimination; the initial precipitous decline is usually evident with administration by intravenous but not by other routes. Drug concentrations at peripheral sites are determined by a balance between drug distribution to and redistribution from those sites, as well as by elimination. Once distribution is near-complete (four–five distribution half-lives), plasma and tissue concentrations decline in parallel.

Clinical Implications of Half-Life Measurements The elimination half-life not only determines the time required for drug concentrations to fall to near-immeasurable levels after a single bolus, it is also the sole determinant of the time required for steady-state plasma concentrations to be achieved after any change in drug dosing (Fig. 5-4). This applies to the initiation of chronic drug therapy (whether by multiple oral doses or by continuous intravenous infusion), a change in chronic drug dose or dosing interval, or discontinuation of drug.

Steady state describes the situation during chronic drug administration when the amount of drug administered per unit time equals drug eliminated per unit time. With a continuous intravenous infusion, plasma concentrations at steady state are stable, while with chronic oral drug administration, plasma concentrations vary during the dosing interval but the time-concentration profile between dosing intervals is stable (Fig. 5-4).

DRUG DISTRIBUTION

In a typical 70-kg human, plasma volume is ~3 L, blood volume is ~5.5 L, and extracellular water outside the vasculature is ~20 L. The volume of distribution of drugs extensively bound to plasma proteins but not to tissue components approaches plasma volume; warfarin is one such example. By contrast, for drugs highly bound to tissues, the volume of distribution can be far greater than any physiologic space. For example, the volume of distribution of digoxin and tricyclic antidepressants is hundreds of liters, obviously exceeding total-body volume. Such drugs are not readily removed by dialysis, an important consideration in overdose.

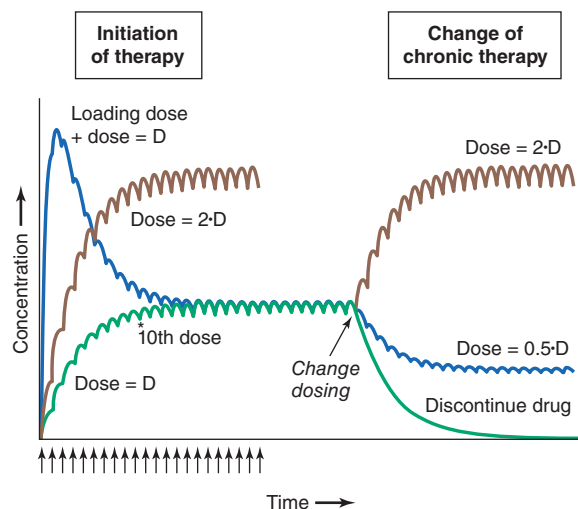


FIGURE 5-4 Drug accumulation to steady state. In this simulation, drug was administered (arrows) at intervals = 50% of the elimination half-life. Steady state is achieved during initiation of therapy after ~5 elimination half-lives, or 10 doses. A loading dose did not alter the eventual steady state achieved. A doubling of the dose resulted in a doubling of the steady state but the same time course of accumulation. Once steady state is achieved, a change in dose (increase, decrease, or drug discontinuation) results in a new steady state in ~5 elimination half-lives. (Adapted by permission from DM Roden, in DP Zipes, J Jalife [eds]: *Cardiac Electrophysiology: From Cell to Bedside*, 4th ed. Philadelphia, Saunders, 2003. Copyright 2003 with permission from Elsevier.)

Clinical Implications of Drug Distribution In some cases, pharmacologic effects require drug distribution to peripheral sites. In this instance, the time course of drug delivery to and removal from these sites determines the time course of drug effects; anesthetic uptake into the central nervous system (CNS) is an example.

LOADING DOSES For some drugs, the indication may be so urgent that administration of “loading” dosages is required to achieve rapid elevations of drug concentration and therapeutic effects earlier than with chronic maintenance therapy (Fig. 5-4). Nevertheless, the time required for true steady state to be achieved is still determined only by the elimination half-life.

RATE OF INTRAVENOUS ADMINISTRATION Although the simulations in Fig. 5-2 use a single intravenous bolus, this is usually inappropriate in practice because side effects related to transiently very high concentrations can result. Rather, drugs are more usually administered orally or as a slower intravenous infusion. Some drugs are so predictably lethal when infused too rapidly that special precautions should be taken to prevent accidental boluses. For example, solutions of potassium for intravenous administration >20 mEq/L should be avoided in all but the most exceptional and carefully monitored circumstances. This minimizes the possibility of cardiac arrest due to accidental increases in infusion rates of more concentrated solutions.

Transiently high drug concentrations after rapid intravenous administration can occasionally be used to advantage. The use of midazolam for intravenous sedation, for example, depends upon its rapid uptake by the brain during the distribution phase to produce sedation quickly, with subsequent egress from the brain during the redistribution of the drug as equilibrium is achieved.

Similarly, adenosine must be administered as a rapid bolus in the treatment of reentrant supraventricular tachycardias (Chap. 276) to prevent elimination by very rapid ($t_{1/2}$ of seconds) uptake into erythrocytes and endothelial cells before the drug can reach its clinical site of action, the atrioventricular node.

Clinical Implications of Altered Protein Binding Many drugs circulate in the plasma partly bound to plasma proteins. Since only unbound (free) drug can distribute to sites of pharmacologic action, drug

response is related to the free rather than the total circulating plasma drug concentration. In chronic kidney or liver disease, protein binding may be decreased and thus drug actions increased. In some situations (myocardial infarction, infection, surgery), acute phase reactants transiently increase drug binding and thus decrease efficacy. These changes assume the greatest clinical importance for drugs that are highly protein-bound since even a small change in protein binding can result in large changes in free drug; for example, a decrease in binding from 99% to 98% doubles the free drug concentration from 1% to 2%. For some drugs (e.g., phenytoin), monitoring free rather than total drug concentrations can be useful.

ELIMINATION

Drug elimination reduces the amount of drug in the body over time. An important approach to quantifying this reduction is to consider that drug concentrations at the beginning and end of a time period are unchanged and that a specific volume of the body has been “cleared” of the drug during that time period. This defines clearance as volume/time. Clearance includes both drug metabolism and excretion.

Clinical Implications of Altered Clearance While elimination half-life determines the time required to achieve steady-state plasma concentration (C_{ss}), the *magnitude* of that steady state is determined by clearance (Cl) and dose alone. For a drug administered as an intravenous infusion, this relationship is:

$$C_{ss} = \text{dosing rate}/Cl \quad \text{or} \quad \text{dosing rate} = Cl \cdot C_{ss}$$

When drug is administered orally, the average plasma concentration within a dosing interval ($C_{avg,ss}$) replaces C_{ss} , and the dosage (dose per unit time) must be increased if bioavailability (F) is less than 1:

$$\text{Dose/time} = Cl \cdot C_{avg,ss}/F$$

Genetic variants, drug interactions, or diseases that reduce the activity of drug-metabolizing enzymes or excretory mechanisms lead to decreased clearance and, hence, a requirement for downward dose adjustment to avoid toxicity. Conversely, some drug interactions and genetic variants increase the function of drug elimination pathways, and hence, increased drug dosage is necessary to maintain a therapeutic effect.

ACTIVE DRUG METABOLITES

Metabolites may produce effects similar to, overlapping with, or distinct from those of the parent drug. Accumulation of the major metabolite of procainamide, *N*-acetylprocainamide (NAPA), likely accounts for marked QT prolongation and torsades des pointes ventricular tachycardia (Chap. 276) during therapy with procainamide. Neurotoxicity during therapy with the opioid analgesic meperidine is likely due to accumulation of normeperidine, especially in renal disease.

Prodrugs are inactive compounds that require metabolism to generate active metabolites that mediate the drug effects. Examples include many angiotensin-converting enzyme (ACE) inhibitors, the angiotensin receptor blocker losartan, the antineoplastic irinotecan, the anti-estrogen tamoxifen, the analgesic codeine (whose active metabolite morphine probably underlies the opioid effect during codeine administration), and the antiplatelet drug clopidogrel. Drug metabolism has also been implicated in bioactivation of procarcinogens and in generation of reactive metabolites that mediate certain adverse drug effects (e.g., acetaminophen hepatotoxicity, discussed below).

THE CONCEPT OF HIGH-RISK PHARMACOKINETICS

When plasma concentrations of active drug depend exclusively on a single metabolic pathway, any condition that inhibits that pathway (be it disease-related, genetic, or due to a drug interaction) can lead to dramatic changes in drug concentrations and marked variability in drug action. This problem of high-risk pharmacokinetics is especially pronounced in two settings. *First*, variability in bioactivation of a prodrug can lead to striking variability in drug action; examples include decreased CYP2D6 activity, which prevents analgesia by codeine, and decreased CYP2C19 activity, which reduces the antiplatelet effects of clopidogrel. The *second* setting is drug elimination that relies on

a single pathway. In this case, inhibition of the elimination pathway by genetic variants or by administration of inhibiting drugs leads to marked elevation of drug concentration and, for drugs with a narrow therapeutic window, an increased likelihood of dose-related toxicity. Individuals with loss-of-function alleles in CYP2C9, responsible for metabolism of the active S-enantiomer of warfarin, appear to be at increased risk for bleeding. When drugs undergo elimination by multiple-drug metabolizing or excretory pathways, absence of one pathway (due to a genetic variant or drug interaction) is much less likely to have a large impact on drug concentrations or drug actions.

PRINCIPLES OF PHARMACODYNAMICS

The Onset of Drug Action For drugs used in the urgent treatment of acute symptoms, little or no delay is anticipated (or desired) between the drug-target interaction and the development of a clinical effect. Examples of such acute situations include vascular thrombosis, shock, or status epilepticus.

For many conditions, however, the indication for therapy is less urgent, and a delay between the interaction of a drug with its pharmacologic target(s) and a clinical effect is clinically acceptable. Common pharmacokinetic mechanisms that can contribute to such a delay include slow elimination (resulting in slow accumulation to steady state), uptake into peripheral compartments, or accumulation of active metabolites. Another common explanation for such a delay is that the clinical effect develops as a downstream consequence of the initial molecular effect the drug produces. Thus, administration of a proton pump inhibitor or an H₂-receptor blocker produces an immediate increase in gastric pH but ulcer healing that is delayed. Cancer chemotherapy similarly produces delayed therapeutic effects.

Drug Effects May Be Disease Specific A drug may produce no action or a different spectrum of actions in unaffected individuals compared to patients with underlying disease. Further, concomitant disease can complicate interpretation of response to drug therapy, especially adverse effects. For example, high doses of anticonvulsants such as phenytoin may cause neurologic symptoms, which may be confused with the underlying neurologic disease. Similarly, increasing dyspnea in a patient with chronic lung disease receiving amiodarone therapy could be due to drug, underlying disease, or an intercurrent cardiopulmonary problem. Thus, the presence of chronic lung disease may argue against the use of amiodarone.

While drugs interact with specific molecular receptors, drug effects may vary over time, even if stable drug and metabolite concentrations are maintained. The drug-receptor interaction occurs in a complex biologic milieu that can vary to modulate the drug effect. For example, ion channel blockade by drugs, an important anticonvulsant and antiarrhythmic effect, is often modulated by membrane potential, itself a function of factors such as extracellular potassium or local ischemia. Receptors may be up- or downregulated by disease or by the drug itself. For example, β -adrenergic blockers upregulate β -receptor density during chronic therapy. While this effect does not usually result in resistance to the therapeutic effect of the drugs, it may produce severe agonist-mediated effects (such as hypertension or tachycardia) if the blocking drug is abruptly withdrawn.

PRINCIPLES OF DOSE SELECTION

The desired goal of therapy with any drug is to maximize the likelihood of a beneficial effect while minimizing the risk of adverse effects. Previous experience with the drug, in controlled clinical trials or in postmarketing use, defines the relationships between dose or plasma concentration and these dual effects (Fig. 5-1) and has important implications for initiation of drug therapy:

1. *The target drug effect should be defined when drug treatment is started.* With some drugs, the desired effect may be difficult to measure objectively, or the onset of efficacy can be delayed for weeks or months; drugs used in the treatment of cancer and psychiatric disease are examples. Sometimes a drug is used to treat a symptom, such as pain or palpitations, and here it is the patient who will report whether the selected dose is effective. In yet other settings,

such as anticoagulation or hypertension, the desired response can be repeatedly and objectively assessed by simple clinical or laboratory tests.

2. *The nature of anticipated toxicity often dictates the starting dose.* If side effects are minor, it may be acceptable to start chronic therapy at a dose highly likely to achieve efficacy and down-titrate if side effects occur. However, this approach is rarely, if ever, justified if the anticipated toxicity is serious or life-threatening; in this circumstance, it is more appropriate to initiate therapy with the lowest dose that may produce a desired effect. In cancer chemotherapy, it is common practice to use maximum-tolerated doses.
3. *The above considerations do not apply if these relationships between dose and effects cannot be defined.* This is especially relevant to some adverse drug effects (discussed in further detail below) whose development are not readily related to drug dose.
4. *If a drug dose does not achieve its desired effect, a dosage increase is justified only if toxicity is absent and the likelihood of serious toxicity is small.*

Failure of Efficacy Assuming the diagnosis is correct and the correct drug is prescribed, explanations for failure of efficacy include drug interactions, noncompliance, or unexpectedly low drug dosage due to administration of expired or degraded drug. These are situations in which measurement of plasma drug concentrations, if available, can be especially useful. Noncompliance is an especially frequent problem in the long-term treatment of diseases such as hypertension and epilepsy, occurring in $\geq 25\%$ of patients in therapeutic environments in which no special effort is made to involve patients in the responsibility for their own health. Multidrug regimens with multiple doses per day are especially prone to noncompliance.

Monitoring response to therapy, by physiologic measures or by plasma concentration measurements, requires an understanding of the relationships between plasma concentration and anticipated effects. For example, measurement of QT interval is used during treatment with sotalol or dofetilide to avoid marked QT prolongation that can herald serious arrhythmias. In this setting, evaluating the electrocardiogram at the time of anticipated peak plasma concentration and effect (e.g., 1–2 h postdose at steady state) is most appropriate. Maintained high vancomycin levels carry a risk of nephrotoxicity, so dosages should be adjusted on the basis of plasma concentrations measured at trough (predose). Similarly, for dose adjustment of other drugs (e.g., anticonvulsants), concentration should be measured at its lowest during the dosing interval, just prior to a dose at steady state (Fig. 5-4), to ensure a maintained therapeutic effect.

Concentration of Drugs in Plasma as a Guide to Therapy Factors such as interactions with other drugs, disease-induced alterations in elimination and distribution, and genetic variation in drug disposition combine to yield a wide range of plasma levels in patients given the same dose. Hence, if a predictable relationship can be established between plasma drug concentration and beneficial or adverse drug effect, measurement of plasma levels can provide a valuable tool to guide selection of an optimal dose, especially when there is a narrow range between the plasma levels yielding therapeutic and adverse effects. Monitoring is commonly used with certain types of drugs including many anticonvulsants, antiemetic agents, antiarrhythmics, and antibiotics. By contrast, if no such relationship can be established (e.g., if drug access to important sites of action outside plasma is highly variable), monitoring plasma concentration may not provide an accurate guide to therapy (Fig. 5-5A).

The common situation of first-order elimination implies that average, maximum, and minimum steady-state concentrations are related linearly to the dosing rate. Accordingly, the maintenance dose may be adjusted on the basis of the ratio between the desired and measured concentrations *at steady state*; for example, if a doubling of the steady-state plasma concentration is desired, the dose should be doubled. This does not apply to drugs eliminated by zero-order kinetics (fixed amount per unit time), where small dosage increases will produce disproportionate increases in plasma concentration; examples include phenytoin and theophylline.

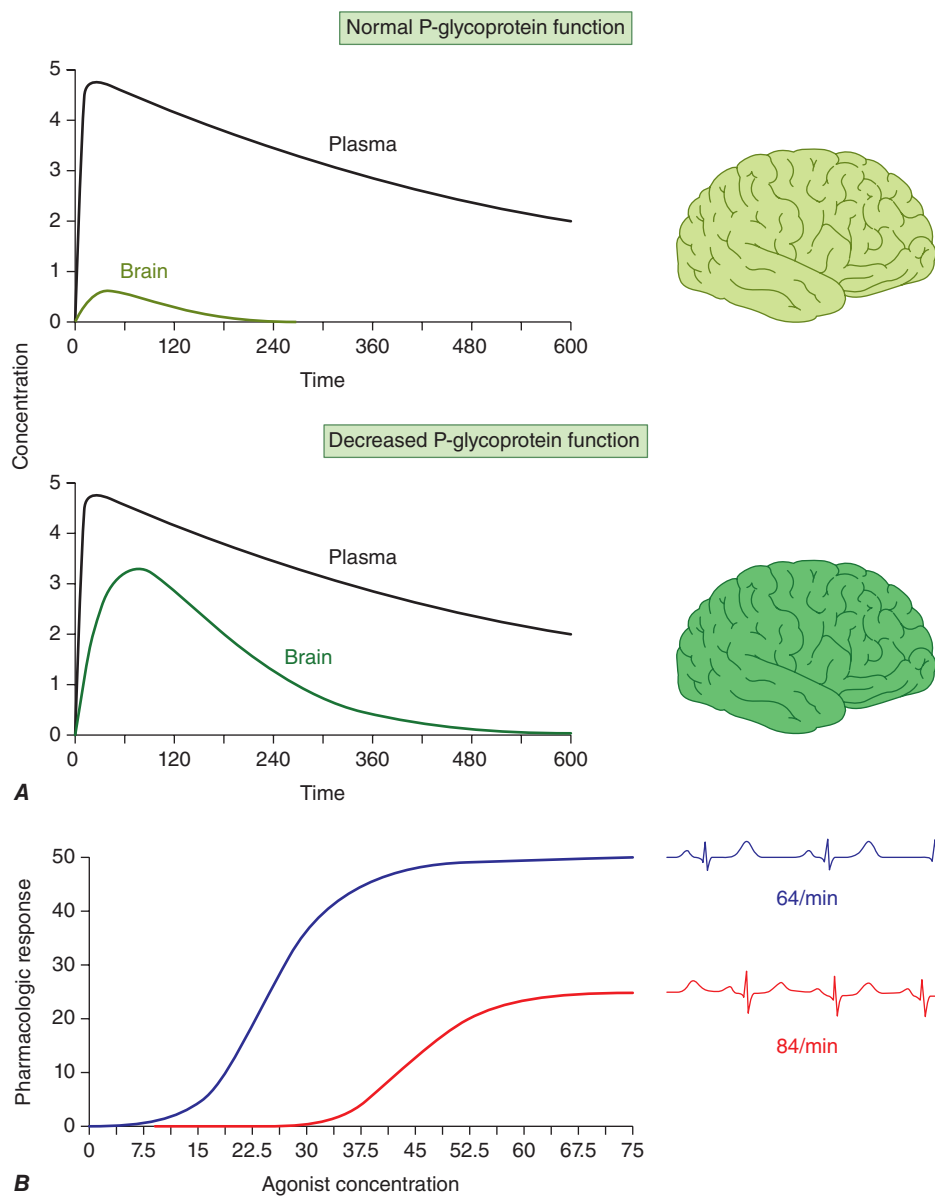


FIGURE 5-5 **A.** The efflux pump P-glycoprotein excludes drugs from the endothelium of capillaries in the brain and so constitutes a key element of the blood-brain barrier. Thus, reduced P-glycoprotein function (e.g., due to drug interactions or genetically determined variability in gene transcription) increases penetration of substrate drugs into the brain, even when plasma concentrations are unchanged. **B.** The graph shows an effect of a β_1 -receptor polymorphism on receptor function in vitro. Patients with the hypofunctional variant (red) may display lesser heart-rate slowing or blood pressure lowering on exposure to a receptor blocking agent.

An increase in dosage is usually best achieved by changing the drug dose but not the dosing interval (e.g., by giving 200 mg every 8 h instead of 100 mg every 8 h). However, this approach is acceptable only if the resulting maximum concentration is not toxic and the trough value does not fall below the minimum effective concentration for an undesirable period of time. Alternatively, the steady state may be changed by altering the frequency of intermittent dosing but not the size of each dose. In this case, the magnitude of the fluctuations around the average steady-state level will change—the shorter the dosing interval, the smaller the difference between peak and trough levels.

EFFECTS OF DISEASE ON DRUG CONCENTRATION AND RESPONSE

RENAL DISEASE

Renal excretion of parent drug and metabolites is generally accomplished by glomerular filtration and by specific drug transporters. If a drug or its metabolites are primarily excreted through the kidneys

and increased drug levels are associated with adverse effects, drug dosages must be reduced in patients with renal dysfunction to avoid toxicity. The antiarrhythmics dofetilide and sotalol undergo predominant renal excretion and carry a risk of QT prolongation and arrhythmias if doses are not reduced in renal disease. In end-stage renal disease, sotalol has been given as 40 mg after dialysis (every second day), compared to the usual daily dose, 80–120 mg every 12 h. The narcotic analgesic meperidine undergoes extensive hepatic metabolism, so that renal failure has little effect on its plasma concentration. However, its metabolite, normeperidine, does undergo renal excretion, accumulates in renal failure, and probably accounts for the signs of CNS excitation, such as irritability, twitching, and seizures, that appear when multiple doses of meperidine are administered to patients with renal disease. Protein binding of some drugs (e.g., phenytoin) may be altered in uremia, so measuring free drug concentration may be desirable.

In non-end-stage renal disease, changes in renal drug clearance are generally proportional to those in creatinine clearance, which may be measured directly or estimated from the serum creatinine (**Chap. 333e**). This estimate, coupled with the knowledge of how much drug is normally excreted renally versus nonrenally, allows an estimate of the dose adjustment required. In practice, most decisions involving dosing adjustment in patients with renal failure use published recommended adjustments in dosage or dosing interval based on the severity of renal dysfunction indicated by creatinine clearance. Any such modification of dose is a first approximation and should be followed by plasma concentration data (if available) and clinical observation to further optimize therapy for the individual patient.

LIVER DISEASE

Standard tests of liver function are not useful in adjusting doses in diseases like hepatitis or cirrhosis. First-pass metabolism may decrease, leading to increased oral bioavailability as a consequence of disrupted hepatocyte function, altered liver architecture, and portacaval shunts. The oral bioavailability for high first-pass drugs such as morphine, meperidine, midazolam, and nifedipine is almost doubled in patients with cirrhosis, compared to those with normal liver function. Therefore, the size of the oral dose of such drugs should be reduced in this setting.

HEART FAILURE AND SHOCK

Under conditions of decreased tissue perfusion, the cardiac output is redistributed to preserve blood flow to the heart and brain at the expense of other tissues (**Chap. 279**). As a result, drugs may be distributed into a smaller volume of distribution, higher drug concentrations will be present in the plasma, and the tissues that are best perfused (the brain and heart) will be exposed to these higher concentrations, resulting in increased CNS or cardiac effects. As well, decreased perfusion of the kidney and liver may impair drug clearance. Another consequence of severe heart failure is decreased gut perfusion, which may reduce drug absorption and, thus, lead to reduced or absent effects of orally administered therapies.

DRUG USE IN THE ELDERLY

In the elderly, multiple pathologies and medications used to treat them result in more drug interactions and adverse effects. Aging also results in changes in organ function, especially of the organs involved in drug disposition. Initial doses should be less than the usual adult dosage and should be increased slowly. The number of medications, and doses per day, should be kept as low as possible.

Even in the absence of kidney disease, renal clearance may be reduced by 35–50% in elderly patients. Dosages should be adjusted on the basis of creatinine clearance. Aging also results in a decrease in the size of, and blood flow to, the liver and possibly in the activity of hepatic drug-metabolizing enzymes; accordingly, the hepatic clearance of some drugs is impaired in the elderly. As with liver disease, these changes are not readily predicted.

Elderly patients may display altered drug sensitivity. Examples include increased analgesic effects of opioids, increased sedation from benzodiazepines and other CNS depressants, and increased risk of bleeding while receiving anticoagulant therapy, even when clotting parameters are well controlled. Exaggerated responses to cardiovascular drugs are also common because of the impaired responsiveness of normal homeostatic mechanisms. Conversely, the elderly display decreased sensitivity to β -adrenergic receptor blockers.

Adverse drug reactions are especially common in the elderly because of altered pharmacokinetics and pharmacodynamics, the frequent use of multidrug regimens, and concomitant disease. For example, use of long half-life benzodiazepines is linked to the occurrence of hip fractures in elderly patients, perhaps reflecting both a risk of falls from these drugs (due to increased sedation) and the increased incidence of osteoporosis in elderly patients. In population surveys of the noninstitutionalized elderly, as many as 10% had at least one adverse drug reaction in the previous year.

DRUG USE IN CHILDREN

While most drugs used to treat disease in children are the same as those in adults, there are few studies that provide solid data to guide dosing. Drug metabolism pathways mature at different rates after birth, and disease mechanisms may be different in children. In practice, doses are adjusted for size (weight or body surface area) as a first approximation unless age-specific data are available.

GENETIC DETERMINANTS OF THE RESPONSE TO DRUGS

PRINCIPLES OF GENETIC VARIATION AND HUMAN TRAITS

(SEE ALSO CHAPS. 82 AND 84)

The concept that genetically determined variations in drug metabolism might be associated with variable drug levels and hence, effect, was advanced at the end of the nineteenth century, and the examples of familial clustering of unusual drug responses were noted in the mid-twentieth century. A goal of traditional Mendelian genetics is to identify DNA variants associated with a distinct phenotype in multiple related family members (Chap. 84). However, it is unusual for a drug response phenotype to be accurately measured in more than one family member, let alone across a kindred. Thus, non-family-based approaches are generally used to

identify and validate DNA variants contributing to variable drug actions.

Candidate Gene Studies in Pharmacogenetics Most studies to date have used an understanding of the molecular mechanisms modulating drug action to identify candidate genes in which variants could explain variable drug responses. One very common scenario is that variable drug actions can be attributed to variability in plasma drug concentrations. When plasma drug concentrations vary widely (e.g., more than an order of magnitude), especially if their distribution is non-unimodal as in Fig. 5-6, variants in single genes controlling drug concentrations often contribute. In this case, the most obvious candidate genes are those responsible for drug metabolism and elimination. Other candidate genes are those encoding the target molecules with which drugs interact to produce their effects or molecules modulating that response, including those involved in disease pathogenesis.

Genome-Wide Association Studies in Pharmacogenomics The field has also had some success with “unbiased” approaches such as genome-wide association (GWA) (Chap. 82), particularly in identifying single variants associated with high risk for certain forms of drug toxicity (Table 5-2). GWA studies have identified variants in the HLA-B locus that are associated with high risk for severe skin rashes during treatment with the anticonvulsant carbamazepine and the antiretroviral abacavir. A GWA study of simvastatin-associated myopathy identified a single non-coding single nucleotide polymorphism (SNP) in *SLCO1B1*, encoding OATP1B1, a drug transporter known to modulate simvastatin

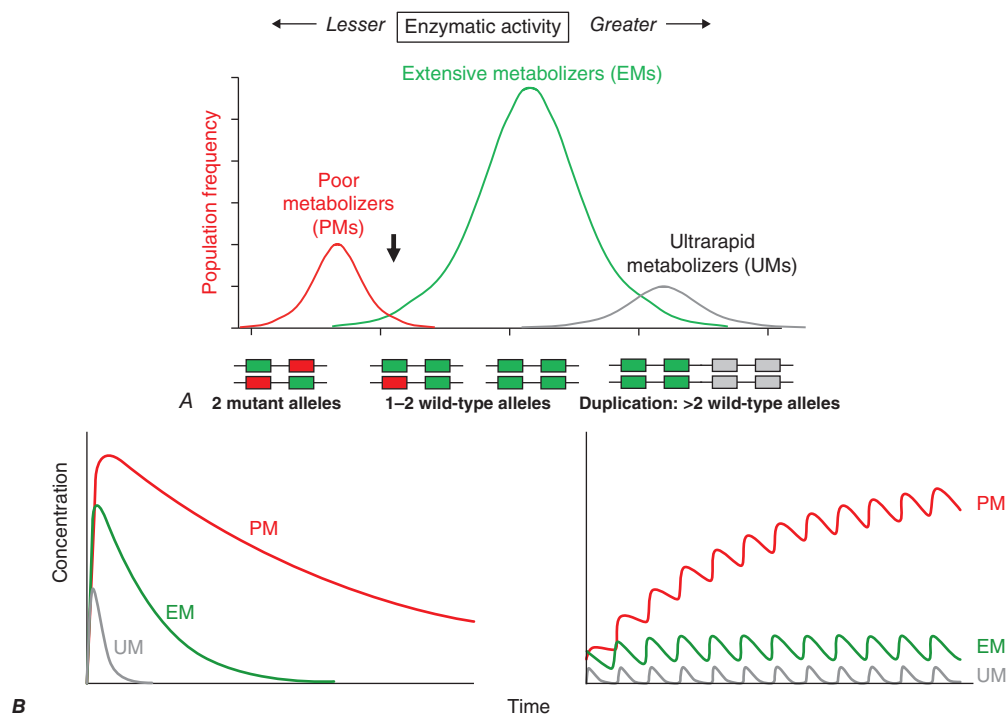


FIGURE 5-6 A. CYP2D6 metabolic activity was assessed in 290 subjects by administration of a test dose of a probe substrate and measurement of urinary formation of the CYP2D6-generated metabolite. The heavy arrow indicates a clear antimode, separating poor metabolizer subjects (PMs, red), with two loss-of-function CYP2D6 alleles, indicated by the intron-exon structures below the chart. Individuals with one or two functional alleles are grouped together as extensive metabolizers (EMs, green). Also shown are ultra-rapid metabolizers (UMs), with 2–12 functional copies of the gene (gray), displaying the greatest enzyme activity. (Adapted from M-L Dahl et al. *J Pharmacol Exp Ther* 274:516, 1995.) B. These simulations show the predicted effects of CYP2D6 genotype on disposition of a substrate drug. With a single dose (left), there is an inverse “gene-dose” relationship between the number of active alleles and the areas under the time-concentration curves (smallest in UM subjects; highest in PM subjects); this indicates that clearance is greatest in UM subjects. In addition, elimination half-life is longest in PM subjects. The right panel shows that these single dose differences are exaggerated during chronic therapy: steady-state concentration is much higher in PM subjects (decreased clearance), as is the time required to achieve steady state (longer elimination half-life).

TABLE 5-2 GENETIC VARIANTS AND DRUG RESPONSES

Gene	Drugs	Effect of Genetic Variants ^a
Variants in Drug Metabolism Pathways		
CYP2C9	Losartan	Decreased bioactivation and effects (PMs)
	Warfarin	Decreased dose requirements; possible increased bleeding risk (PMs)
CYP2C19	Omeprazole, voriconazole	Decreased effect in extensive metabolizers (EMs)
	Celecoxib	Exaggerated effect in PMs
CYP2D6	Clopidogrel	Decreased effect in PMs
	Codeine, tamoxifen	Decreased bioactivation and drug effects in PMs
	Codeine	Morphine-like adverse effects in UMs
	Tricyclic antidepressants	Increased adverse effects in PMs; decreased therapeutic effects in UMs
CYP3A5	Metoprolol, carvedilol, timolol, propafenone	Increased beta blockade in PMs
	Tacrolimus, vincristine	Decreased drug concentrations and effect
Dihydropyrimidine dehydrogenase	Capecitabine, fluorouracil	Possible severe toxicity (PMs)
NAT2	Rifampin, isoniazid, pyrazinamide, hydralazine, procainamide	Increased risk of toxicity in PMs
Thiopurine S-methyltransferase (<i>TPMT</i>)	Azathioprine, 6-mercaptopurine	*3A/*3A (PMs): increased risk of bone marrow aplasia; wild-type homozygote: possible decreased drug action at usual dosages
Uridine diphosphate glucuronosyltransferase (<i>UGT1A1</i>)	Irinotecan	*28/*28 PM homozygotes: increased risk of severe adverse effects (diarrhea, bone marrow aplasia)
Variants in Other Genes		
Glucose 6-phosphate dehydrogenase (G6PD)	Rasburicase, primaquine, chloroquine	Increased risk of hemolytic anemia in G6PD-deficient subjects
HLA-B*1501	Carbamazepine	Carriers (1 or 2 alleles) at increased risk of severe skin toxicity
HLA-B*5701	Abacavir	Carriers (1 or 2 alleles) at increased risk of severe skin toxicity
IL28B	Interferon	Variable response in hepatitis C therapy
IL15	Childhood leukemia therapy	Variability in response
SLCO1B1	Simvastatin	Encodes a drug uptake transporter; variant non-synonymous single nucleotide polymorphism increases myopathy risk
VKORC1	Warfarin	Decreased dose requirements with variant promoter haplotype
ITPA	Ribavirin	Variants modulate risk for hemolytic anemia
Variants in Other Genomes (Infectious Agents, Tumors)		
Chemokine C-C motif receptor (CCR5)	Maraviroc	Drug effective only in HIV strains with CCR5 detectible
C-KIT	Imatinib	In gastrointestinal stromal tumors, drug indicated only with c-kit–positive cases
Epidermal growth factor receptor (EGFR)	Cetuximab	Clinical trials conducted in patients with EGFR-positive tumors
Her2/neu overexpression	Trastuzumab, lapatinib	Drugs indicated only with tumor overexpression
K-ras mutation	Panitumumab, cetuximab	Lack of efficacy with <i>KRAS</i> mutation
Philadelphia chromosome	Busulfan, dasatinib, nilotinib, imatinib	Decreased efficacy in Philadelphia chromosome–negative chronic myelogenous leukemia

^aDrug effect in homozygotes unless otherwise specified.

Note: EM, extensive metabolizer (normal enzymatic activity); PM, poor metabolizer (homozygote for reduced or loss of function allele); UM, ultra-rapid metabolizer (enzymatic activity much greater than normal, e.g., with gene duplication, Fig. 5-6). Further data at U.S. Food and Drug Administration: <http://www.fda.gov/Drugs/ScienceResearch/ResearchAreas/Pharmacogenetics/ucm083378.htm>; or Pharmacogenetics Research Network/Knowledge Base: <http://www.pharmgkb.org>.

uptake into the liver, which accounts for 60% of myopathy risk. GWA approaches have also implicated interferon variants in antileukemic responses and in response to therapy in hepatitis C. Ribavirin, used as therapy in hepatitis C, causes hemolytic anemia, and this has been linked to variants in *ITPA*, encoding inosine triphosphatase.

GENETIC VARIANTS AFFECTING PHARMACOKINETICS

Clinically important genetic variants have been described in multiple molecular pathways of drug disposition (Table 5-2). A distinct multimodal distribution of drug disposition (as shown in Fig. 5-6) argues for a predominant effect of variants in a single gene in the metabolism of

that substrate. Individuals with two alleles (variants) encoding for non-functional protein make up one group, often termed *poor metabolizers* (PM phenotype); for some genes, many variants can produce such a loss of function, complicating the use of genotyping in clinical practice. Individuals with one functional allele make up a second (*intermediate metabolizers*) and may or may not be distinguishable from those with two functional alleles (*extensive metabolizers*, EMs). *Ultra-rapid metabolizers* with especially high enzymatic activity (occasionally due to gene duplication; Fig. 5-6) have also been described for some traits. Many drugs in widespread use can inhibit specific drug disposition pathways (Table 5-1), and so EM individuals receiving such inhibitors can respond like PM patients (*phenocopying*). Polymorphisms in genes encoding drug uptake or drug efflux transporters may be other contributors to variability in drug delivery to target sites and, hence, in drug effects.

CYP Variants Members of the CYP3A family (CYP3A4, 3A5) metabolize the greatest number of drugs in therapeutic use. CYP3A4 activity is highly variable (up to an order of magnitude) among individuals, but the underlying mechanisms are not well understood. In whites, but not African Americans, there is a common loss-of-function polymorphism in the closely related CYP3A5 gene. Decreased efficacy of the antirejection agent tacrolimus in African-American subjects has been attributed to more rapid elimination due to relatively greater CYP3A5 activity. A lower risk of vincristine-associated neuropathy has been reported in CYP3A5 “expressers.”

CYP2D6 is second to CYP3A4 in the number of commonly used drugs that it metabolizes. CYP2D6 activity is polymorphically distributed, with about 7% of European- and African-derived populations (but very few Asians) displaying the PM phenotype (Fig. 5-6). Dozens of loss-of-function variants in the CYP2D6 gene have been described; the PM phenotype arises in individuals with two such alleles. In addition, ultra-rapid metabolizers with multiple functional copies of the CYP2D6 gene have been identified.

Codeine is biotransformed by CYP2D6 to the potent active metabolite morphine, so its effects are blunted in PMs and exaggerated in ultra-rapid metabolizers. In the case of drugs with beta-blocking properties metabolized by CYP2D6, greater signs of beta blockade (e.g., bronchospasm, bradycardia) are seen in PM subjects than in EMs. This can be seen not only with orally administered beta blockers such as metoprolol and carvedilol, but also with ophthalmic timolol and with the sodium channel-blocking antiarrhythmic propafenone, a CYP2D6 substrate with beta-blocking properties. Ultra-rapid metabolizers may require very high dosages of tricyclic antidepressants to achieve a therapeutic effect and, with codeine, may display transient euphoria and nausea due to very rapid generation of morphine. Tamoxifen undergoes CYP2D6-mediated biotransformation to an active metabolite, so its efficacy may be in part related to this polymorphism. In addition, the widespread use of selective serotonin reuptake inhibitors (SSRIs) to treat tamoxifen-related hot flashes may also alter the drug's effects because many SSRIs, notably fluoxetine and paroxetine, are also CYP2D6 inhibitors.

The PM phenotype for CYP2C19 is common (20%) among Asians and rarer (2–3%) in European-derived populations. The impact of polymorphic CYP2C19-mediated metabolism has been demonstrated with the proton pump inhibitor omeprazole, where ulcer cure rates with “standard” dosages were much lower in EM patients (29%) than in PMs (100%). Thus, understanding the importance of this polymorphism would have been important in developing the drug, and knowing a patient's CYP2C19 genotype should improve therapy. CYP2C19 is responsible for bioactivation of the antiplatelet drug clopidogrel, and several large studies have documented decreased efficacy (e.g., increased myocardial infarction after placement of coronary stents) among Caucasian subjects with reduction of function alleles. In addition, some studies suggest that omeprazole and possibly other proton pump inhibitors phenocopy this effect.

There are common variants of CYP2C9 that encode proteins with loss of catalytic function. These variant alleles are associated with increased rates of neurologic complications with phenytoin, hypoglycemia with glipizide, and reduced warfarin dose required to maintain stable anticoagulation. The angiotensin-receptor blocker losartan is a prodrug that is bioactivated by CYP2C9; as a result, PMs and those receiving inhibitor drugs may display little response to therapy.

Transferase Variants One of the most extensively studied phase II polymorphisms is the PM trait for thiopurine S-methyltransferase (TPMT). TPMT bioinactivates the antileukemic drug 6-mercaptopurine. Further, 6-mercaptopurine is itself an active metabolite of the immunosuppressive azathioprine. Homozygotes for alleles encoding the inactive TPMT (1 in 300 individuals) predictably exhibit severe and potentially fatal pancytopenia on standard doses of azathioprine or 6-mercaptopurine. On the other hand, homozygotes for fully functional alleles may display less anti-inflammatory or antileukemic effect with the drugs.

N-acetylation is catalyzed by hepatic N-acetyl transferase (NAT), which represents the activity of two genes, NAT-1 and NAT-2. Both enzymes transfer an acetyl group from acetyl coenzyme A to the drug; polymorphisms in NAT-2 are thought to underlie individual differences in the rate at which drugs are acetylated and thus define “rapid acetylators” and “slow acetylators.” Slow acetylators make up ~50% of European- and African-derived populations but are less common among Asians. Slow acetylators have an increased incidence of the drug-induced lupus syndrome during procainamide and hydralazine therapy and of hepatitis with isoniazid. Induction of CYPs (e.g., by rifampin) also increases the risk of isoniazid-related hepatitis, likely reflecting generation of reactive metabolites of acetylhydrazine, itself an isoniazid metabolite.

Individuals homozygous for a common promoter polymorphism that reduces transcription of uridine diphosphate glucuronosyltransferase (*UGT1A1*) have benign hyperbilirubinemia (Gilbert's syndrome; Chap. 358). This variant has also been associated with diarrhea and increased bone marrow depression with the antineoplastic prodrug irinotecan, whose active metabolite is normally detoxified by UGT1A1-mediated glucuronidation. The antiretroviral atazanavir is a UGT1A1 inhibitor, and individuals with the Gilbert's variant develop higher bilirubin levels during treatment.

VARIABILITY IN THE MOLECULAR TARGETS WITH WHICH DRUGS INTERACT

Multiple polymorphisms identified in the β_2 -adrenergic receptor appear to be linked to specific phenotypes in asthma and congestive heart failure, diseases in which β_2 -receptor function might be expected to determine prognosis. Polymorphisms in the β_2 -receptor gene have also been associated with response to inhaled β_2 -receptor agonists, while those in the β_1 -adrenergic receptor gene have been associated with variability in heart rate slowing and blood pressure lowering (Fig. 5-5B). In addition, in heart failure, a common polymorphism in the β_1 -adrenergic receptor gene has been implicated in variable clinical outcome during therapy with the investigational beta blocker bucindolol. Response to the 5-lipoxygenase inhibitor zileuton in asthma has been linked to polymorphisms that determine the expression level of the 5-lipoxygenase gene.

Drugs may also interact with genetic pathways of disease to elicit or exacerbate symptoms of the underlying conditions. In the porphyrias, CYP inducers are thought to increase the activity of enzymes proximal to the deficient enzyme, exacerbating or triggering attacks (Chap. 430). Deficiency of glucose-6-phosphate dehydrogenase (G6PD), most often in individuals of African, Mediterranean, or South Asian descent, increases the risk of hemolytic anemia in response to the antimalarial primaquine (Chap. 129) and the uric acid-lowering agent rasburicase, which do not cause hemolysis in patients with normal amounts of the enzyme. Patients with mutations in the ryanodine receptor, which controls intracellular calcium in skeletal muscle and

other tissues, may be asymptomatic until exposed to certain general anesthetics, which trigger the rare syndrome of malignant hyperthermia. Certain antiarrhythmics and other drugs can produce marked QT prolongation and torsades des pointes (Chap. 276), and in some patients, this adverse effect represents unmasking of previously subclinical congenital long QT syndrome. Up to 50% of the variability in steady-state warfarin dose requirement is attributable to polymorphisms in the promoter of *VKORC1*, which encodes the warfarin target, and in the coding region of *CYP2C9*, which mediates its elimination.

Tumor and Infectious Agent Genomes The actions of drugs used to treat infectious or neoplastic disease may be modulated by variants in these nonhuman germline genomes. Genotyping tumors is a rapidly evolving approach to target therapies to underlying mechanisms and to avoid potentially toxic therapy in patients who would derive no benefit (Chap. 101e). Trastuzumab, which potentiates anthracycline-related cardiotoxicity, is ineffective in breast cancers that do not express the herceptin receptor. Imatinib targets a specific tyrosine kinase, BCR-Abl1, that is generated by the translocation that creates the Philadelphia chromosome typical of chronic myelogenous leukemia (CML). BCR-Abl1 is not only active but may be central to the pathogenesis of CML; its use in BCR-Abl1-positive tumors has resulted in remarkable antitumor efficacy. Similarly, the anti-epidermal growth factor receptor (EGFR) antibodies cetuximab and panitumumab appear especially effective in colon cancers in which K-ras, a G protein in the EGFR pathway, is not mutated. Vemurafenib does not inhibit wild-type *BRAF* but is active against the V600E mutant form of the kinase.

PROSPECTS FOR INCORPORATING PHARMACOGENETIC INFORMATION INTO CLINICAL PRACTICE

The description of genetic variants linked to variable drug responses naturally raises the question of if and how to use this information in practice. Indeed, the U.S. Food and Drug Administration (FDA) now incorporates pharmacogenetic data into information (“package inserts”) meant to guide prescribing. A decision to adopt pharmacogenetically guided dosing for a given drug depends on multiple factors. The most important are the magnitude and clinical importance of the genetic effect and the strength of evidence linking genetic variation to variable drug effects (e.g., anecdote versus post-hoc analysis of clinical trial data versus randomized prospective clinical trial). The evidence can be strengthened if statistical arguments from clinical trial data are complemented by an understanding of underlying physiologic mechanisms. Cost versus expected benefit may also be a factor.

When the evidence is compelling, alternate therapies are not available, and there are clear recommendations for dosage adjustment in subjects with variants, there is a strong argument for deploying genetic testing as a guide to prescribing. The association between HLA-B*5701 and severe skin toxicity with abacavir is an example. In other situations, the arguments are less compelling: the magnitude of the genetic effect may be smaller, the consequences may be less serious, alternate therapies may be available, or the drug effect may be amenable to monitoring by other approaches. Ongoing clinical trials are addressing the utility of preprescription genotyping in large populations exposed to drugs with known pharmacogenetic variants (e.g., warfarin). Importantly, technological advances are now raising the possibility of inexpensive whole genome sequencing. Incorporating a patient’s whole genome sequence into their electronic medical record would allow the information to be accessed as needed for many genetic and pharmacogenetic applications, and the argument has been put forward that this approach would lower logistic barriers to use of pharmacogenomic variant data in prescribing. There are multiple issues (e.g., economic, technological, and ethical) that need to be addressed if such a paradigm is to be adopted (Chap. 82). While barriers to bringing genomic and pharmacogenomic information to the bedside seem daunting, the field is very young and evolving rapidly. Indeed, one major result of understanding the role of genetics in drug action has

been improved screening of drugs during the development process to reduce the likelihood of highly variable metabolism or unanticipated toxicity.

INTERACTIONS BETWEEN DRUGS

Drug interactions can complicate therapy by increasing or decreasing the action of a drug; interactions may be based on changes in drug disposition or in drug response in the absence of changes in drug levels. *Interactions must be considered in the differential diagnosis of any unusual response occurring during drug therapy.* Prescribers should recognize that patients often come to them with a legacy of drugs acquired during previous medical experiences, often with multiple physicians who may not be aware of all the patient’s medications. A meticulous drug history should include examination of the patient’s medications and, if necessary, calls to the pharmacist to identify prescriptions. It should also address the use of agents not often volunteered during questioning, such as OTC drugs, health food supplements, and topical agents such as eye drops. Lists of interactions are available from a number of electronic sources. While it is unrealistic to expect the practicing physician to memorize these, certain drugs consistently run the risk of generating interactions, often by inhibiting or inducing specific drug elimination pathways. Examples are presented below and in Table 5-3. Accordingly, when these drugs are started or stopped, prescribers must be especially alert to the possibility of interactions.

PHARMACOKINETIC INTERACTIONS CAUSING DECREASED DRUG EFFECTS

Gastrointestinal absorption can be reduced if a drug interaction results in drug binding in the gut, as with aluminum-containing antacids, kaolin-pectin suspensions, or bile acid sequestrants. Drugs such as histamine H₂-receptor antagonists or proton pump inhibitors that alter gastric pH may decrease the solubility and hence absorption of weak bases such as ketoconazole.

Expression of some genes responsible for drug elimination, notably *CYP3A* and *MDR1*, can be markedly increased by inducing drugs, such as rifampin, carbamazepine, phenytoin, St. John’s wort, and glutethimide, and by smoking, exposure to chlorinated insecticides such as DDT (*CYP1A2*), and chronic alcohol ingestion. Administration of inducing agents lowers plasma levels, and thus effects, over 2–3 weeks as gene expression is increased. If a drug dose is stabilized in the presence of an inducer that is subsequently stopped, major toxicity can occur as clearance returns to preinduction levels and drug concentrations rise. Individuals vary in the extent to which drug metabolism can be induced, likely through genetic mechanisms.

Interactions that inhibit the bioactivation of prodrugs will decrease drug effects (Table 5-1).

Interactions that decrease drug delivery to intracellular sites of action can decrease drug effects: tricyclic antidepressants can blunt the antihypertensive effect of clonidine by decreasing its uptake into adrenergic neurons. Reduced CNS penetration of multiple HIV protease inhibitors (with the attendant risk of facilitating viral replication in a sanctuary site) appears attributable to P-glycoprotein-mediated exclusion of the drug from the CNS; indeed, inhibition of P-glycoprotein has been proposed as a therapeutic approach to enhance drug entry to the CNS (Fig. 5-5A).

PHARMACOKINETIC INTERACTIONS CAUSING INCREASED DRUG EFFECTS

The most common mechanism here is inhibition of drug elimination. In contrast to induction, new protein synthesis is not involved, and the effect develops as drug and any inhibitor metabolites accumulate (a function of their elimination half-lives). Since shared substrates of a single enzyme can compete for access to the active site of the protein, many CYP substrates can also be considered inhibitors. However, some drugs are especially potent as inhibitors (and occasionally may not even be substrates) of specific drug elimination pathways, and so it is in the use of these agents that clinicians must be most alert to the potential for interactions (Table 5-3). Commonly implicated interacting drugs of this type include amiodarone, cimetidine,

TABLE 5-3 DRUGS WITH A HIGH RISK OF GENERATING PHARMACOKINETIC INTERACTIONS

Drug	Mechanism	Examples
Antacids	Reduced absorption	Antacids/tetracyclines
Bile acid sequestrants		Cholestyramine/digoxin
Proton pump inhibitors	Altered gastric pH	Ketoconazole absorption decreased
H ₂ -receptor blockers		
Rifampin	Induction of CYPs and/or P-glycoprotein	Decreased concentration and effects of warfarin
Carbamazepine		quinidine
Barbiturates		cyclosporine
Phenytoin		losartan
St. John's wort		oral contraceptives
Glutethimide		methadone, dabigatran
Nevirapine (CYP3A; CYP2B6)		
Tricyclic antidepressants	Inhibitors of CYP2D6	Increased effect of many β blockers
Fluoxetine		Decreased codeine effect; possible decreased tamoxifen effect
Quinidine		
Cimetidine	Inhibitor of multiple CYPs	Increased concentration and effects of warfarin theophylline phenytoin
Ketoconazole, itraconazole	Inhibitor of CYP3A	Increased concentration and toxicity of some HMG-CoA reductase inhibitors
Erythromycin, clarithromycin		cyclosporine, cisapride, terfenadine (now withdrawn)
Calcium channel blockers		Increased concentration and effects of indinavir (with ritonavir)
Ritonavir		Decreased clearance and dose requirement for cyclosporine (with calcium channel blockers)
Allopurinol	Xanthine oxidase inhibitor	Azathioprine and 6-mercaptopurine toxicity
Amiodarone	Inhibitor of many CYPs and of P-glycoprotein	Decreased clearance (risk of toxicity) for warfarin digoxin quinidine
Gemfibrozil (and other fibrates)	CYP3A inhibition	Rhabdomyolysis when co-prescribed with some HMG-CoA reductase inhibitors
Quinidine	P-glycoprotein inhibition	Risk of toxicity with P-glycoprotein substrates (e.g., digoxin, dabigatran)
Amiodarone		
Verapamil		
Cyclosporine		
Itraconazole		
Erythromycin		
Phenylbutazone	Inhibition of renal tubular transport	Increased risk of methotrexate toxicity with salicylates
Probenecid		
Salicylates		

erythromycin and some other macrolide antibiotics (clarithromycin but not azithromycin), ketoconazole and otherazole antifungals, the antiretroviral agent ritonavir, and high concentrations of grapefruit juice (Table 5-3). The consequences of such interactions will depend on the drug whose elimination is being inhibited (see "The Concept of High-Risk Pharmacokinetics," above). Examples include CYP3A inhibitors increasing the risk of cyclosporine toxicity or of rhabdomyolysis with some HMG-CoA reductase inhibitors (lovastatin, simvastatin, atorvastatin, but not pravastatin), and P-glycoprotein inhibitors increasing the risk of toxicity with digoxin therapy or of bleeding with the thrombin inhibitor dabigatran.

These interactions can occasionally be exploited to therapeutic benefit. The antiviral ritonavir is a very potent CYP3A4 inhibitor that is sometimes added to anti-HIV regimens, not because of its antiviral effects but because it decreases clearance, and hence increases efficacy, of other anti-HIV agents. Similarly, calcium channel blockers have been deliberately coadministered with cyclosporine to reduce its clearance and thus its maintenance dosage and cost.

Phenytoin, an inducer of many systems, including CYP3A, inhibits CYP2C9. CYP2C9 metabolism of losartan to its active metabolite is inhibited by phenytoin, with potential loss of antihypertensive effect.

Grapefruit (but not orange) juice inhibits CYP3A, especially at high doses; patients receiving drugs where even modest CYP3A inhibition may increase the risk of adverse effects (e.g., cyclosporine, some HMG-CoA reductase inhibitors) should therefore avoid grapefruit juice.

CYP2D6 is markedly inhibited by quinidine, a number of neuroleptic drugs (chlorpromazine and haloperidol), and the SSRIs fluoxetine and paroxetine. The clinical consequences of fluoxetine's interaction with CYP2D6 substrates may not be apparent for weeks after the drug is started, because of its very long half-life and slow generation of a CYP2D6-inhibiting metabolite.

6-Mercaptopurine is metabolized not only by TPMT but also by xanthine oxidase. When allopurinol, an inhibitor of xanthine oxidase, is administered with standard doses of azathioprine or 6-mercaptopurine, life-threatening toxicity (bone marrow suppression) can result.

A number of drugs are secreted by the renal tubular transport systems for organic anions. Inhibition of these systems can cause excessive drug accumulation. Salicylate, for example, reduces the renal clearance of methotrexate, an interaction that may lead to methotrexate toxicity. Renal tubular secretion contributes substantially to the elimination of penicillin, which can be inhibited (to increase its therapeutic effect) by

probenecid. Similarly, inhibition of the tubular cation transport system by cimetidine decreases the renal clearance of dofetilide.

DRUG INTERACTIONS NOT MEDIATED BY CHANGES IN DRUG DISPOSITION

Drugs may act on separate components of a common process to generate effects greater than either has alone. Antithrombotic therapy with combinations of antiplatelet agents (glycoprotein IIb/IIIa inhibitors, aspirin, clopidogrel) and anticoagulants (warfarin, heparins) is often used in the treatment of vascular disease, although such combinations carry an increased risk of bleeding.

Nonsteroidal anti-inflammatory drugs (NSAIDs) cause gastric ulcers, and in patients treated with warfarin, the risk of upper gastrointestinal bleeding is increased almost threefold by concomitant use of an NSAID.

Indomethacin, piroxicam, and probably other NSAIDs antagonize the antihypertensive effects of β -adrenergic receptor blockers, diuretics, ACE inhibitors, and other drugs. The resulting elevation in blood pressure ranges from trivial to severe. This effect is not seen with aspirin and sulindac but has been found with the cyclooxygenase 2 (COX-2) inhibitor celecoxib.

Torsades des pointes ventricular tachycardia during administration of QT-prolonging antiarrhythmics (quinidine, sotalol, dofetilide) occurs much more frequently in patients receiving diuretics, probably reflecting hypokalemia. In vitro, hypokalemia not only prolongs the QT interval in the absence of drug but also potentiates drug block of ion channels that results in QT prolongation. Also, some diuretics have direct electrophysiologic actions that prolong QT.

The administration of supplemental potassium leads to more frequent and more severe hyperkalemia when potassium elimination is reduced by concurrent treatment with ACE inhibitors, spironolactone, amiloride, or triamterene.

The pharmacologic effects of sildenafil result from inhibition of the phosphodiesterase type 5 isoform that inactivates cyclic guanosine monophosphate (GMP) in the vasculature. Nitroglycerin and related nitrates used to treat angina produce vasodilation by elevating cyclic GMP. Thus, coadministration of these nitrates with sildenafil can cause profound hypotension, which can be catastrophic in patients with coronary disease.

Sometimes, combining drugs can increase overall efficacy and/or reduce drug-specific toxicity. Such therapeutically useful interactions are described in chapters dealing with specific disease entities.

ADVERSE REACTIONS TO DRUGS

The beneficial effects of drugs are coupled with the inescapable risk of untoward effects. The morbidity and mortality from these adverse effects often present diagnostic problems because they can involve every organ and system of the body and may be mistaken for signs of underlying disease. As well, some surveys have suggested that drug therapy for a range of chronic conditions such as psychiatric disease or hypertension does not achieve its desired goal in up to half of treated patients; thus, the most common “adverse” drug effect may be failure of efficacy.

Adverse reactions can be classified in two broad groups. One type results from exaggeration of an intended pharmacologic action of the drug, such as increased bleeding with anticoagulants or bone marrow suppression with antineoplastics. The second type of adverse reaction ensues from toxic effects unrelated to the intended pharmacologic actions. The latter effects are often unanticipated (especially with new drugs) and frequently severe and may result from recognized as well as previously undescribed mechanisms.

Drugs may increase the frequency of an event that is common in a general population, and this may be especially difficult to recognize; an excellent example is the increase in myocardial infarctions with the COX-2 inhibitor rofecoxib. Drugs can also cause rare and serious adverse effects, such as hematologic abnormalities, arrhythmias, severe skin reactions, or hepatic or renal dysfunction. Prior to regulatory approval and marketing, new drugs are tested in relatively few patients who tend to be less sick and to have fewer concomitant diseases than those patients who subsequently receive the drug therapeutically.

Because of the relatively small number of patients studied in clinical trials and the selected nature of these patients, rare adverse effects are generally not detected prior to a drug’s approval; indeed, if they are detected, the new drugs are generally not approved. Therefore, physicians need to be cautious in the prescription of new drugs and alert for the appearance of previously unrecognized adverse events.

Elucidating mechanisms underlying adverse drug effects can assist development of safer compounds or allow a patient subset at especially high risk to be excluded from drug exposure. National adverse reaction reporting systems, such as those operated by the FDA (suspected adverse reactions can be reported online at <http://www.fda.gov/safety/medwatch/default.htm>) and the Committee on Safety of Medicines in Great Britain, can prove useful. The publication or reporting of a newly recognized adverse reaction can in a short time stimulate many similar such reports of reactions that previously had gone unrecognized.

Occasionally, “adverse” effects may be exploited to develop an entirely new indication for a drug. Unwanted hair growth during minoxidil treatment of severely hypertensive patients led to development of the drug for hair growth. Sildenafil was initially developed as an antianginal, but its effects to alleviate erectile dysfunction not only led to a new drug indication but also to increased understanding of the role of type 5 phosphodiesterase in erectile tissue. These examples further reinforce the concept that prescribers must remain vigilant to the possibility that unusual symptoms may reflect unappreciated drug effects.

Some 25–50% of patients make errors in self-administration of prescribed medicines, and these errors can be responsible for adverse drug effects. Similarly, patients commit errors in taking OTC drugs by not reading or following the directions on the containers. Health care providers must recognize that providing directions with prescriptions does not always guarantee compliance.

In hospitals, drugs are administered in a controlled setting, and patient compliance is, in general, ensured. Errors may occur nevertheless—the wrong drug or dose may be given or the drug may be given to the wrong patient—and improved drug distribution and administration systems are addressing this problem.

SCOPE OF THE PROBLEM

Patients receive, on average, 10 different drugs during each hospitalization. The sicker the patient, the more drugs are given, and there is a corresponding increase in the likelihood of adverse drug reactions. When <6 different drugs are given to hospitalized patients, the probability of an adverse reaction is ~5%, but if >15 drugs are given, the probability is >40%. Retrospective analyses of ambulatory patients have revealed adverse drug effects in 20%. Serious adverse reactions are also well-recognized with “herbal” remedies and OTC compounds; examples include kava-associated hepatotoxicity, L-tryptophan-associated eosinophilia-myalgia, and phenylpropanolamine-associated stroke, each of which has caused fatalities.

A small group of widely used drugs accounts for a disproportionate number of reactions. Aspirin and other NSAIDs, analgesics, digoxin, anticoagulants, diuretics, antimicrobials, glucocorticoids, antineoplastics, and hypoglycemic agents account for 90% of reactions.

TOXICITY UNRELATED TO A DRUG’S PRIMARY PHARMACOLOGIC ACTIVITY

Drugs or more commonly reactive metabolites generated by CYPs can covalently bind to tissue macromolecules (such as proteins or DNA) to cause tissue toxicity. Because of the reactive nature of these metabolites, covalent binding often occurs close to the site of production, typically the liver.

The most common cause of drug-induced hepatotoxicity is acetaminophen overdose (Chap. 361). Normally, reactive metabolites are detoxified by combining with hepatic glutathione. When glutathione becomes depleted, the metabolites bind instead to hepatic protein, with resultant hepatocyte damage. The hepatic necrosis produced by the ingestion of acetaminophen can be prevented or attenuated by the administration of substances such as *N*-acetylcysteine that reduce the binding of electrophilic metabolites to hepatic proteins. The risk of acetaminophen-related hepatic necrosis is increased in patients receiving drugs such as phenobarbital or phenytoin, which increase

the rate of drug metabolism, or ethanol, which exhausts glutathione stores. Such toxicity has even occurred with therapeutic dosages, so patients at risk through these mechanisms should be warned.

Most pharmacologic agents are small molecules with low molecular weights (<2000) and thus are poor immunogens. Generation of an immune response to a drug therefore usually requires *in vivo* activation and covalent linkage to protein, carbohydrate, or nucleic acid.

Drug stimulation of antibody production may mediate tissue injury by several mechanisms. The antibody may attack the drug when the drug is covalently attached to a cell and thereby destroy the cell. This occurs in penicillin-induced hemolytic anemia. Antibody-drug-antigen complexes may be passively adsorbed by a bystander cell, which is then destroyed by activation of complement; this occurs in quinine- and quinidine-induced thrombocytopenia. Heparin-induced thrombocytopenia arises when antibodies against complexes of platelet factor 4 peptide and heparin generate immune complexes that activate platelets; thus, the thrombocytopenia is accompanied by “paradoxical” thrombosis and is treated with thrombin inhibitors. Drugs or their reactive metabolites may alter a host tissue, rendering it antigenic and eliciting autoantibodies. For example, hydralazine and procainamide (or their reactive metabolites) can chemically alter nuclear material, stimulating the formation of antinuclear antibodies and occasionally causing lupus erythematosus. Drug-induced pure red cell aplasia (**Chap. 130**) is due to an immune-based drug reaction.

Serum sickness (**Chap. 376**) results from the deposition of circulating drug-antibody complexes on endothelial surfaces. Complement activation occurs, chemotactic factors are generated locally, and an inflammatory response develops at the site of complex entrapment. Arthralgias, urticaria, lymphadenopathy, glomerulonephritis, or cerebritis may result. Foreign proteins (vaccines, streptokinase, therapeutic antibodies) and antibiotics are common causes. Many drugs, particularly antimicrobial agents, ACE inhibitors, and aspirin, can elicit anaphylaxis with production of IgE, which binds to mast cell membranes. Contact with a drug antigen initiates a series of biochemical events in the mast cell and results in the release of mediators that can produce the characteristic urticaria, wheezing, flushing, rhinorrhea, and (occasionally) hypotension.

Drugs may also elicit cell-mediated immune responses. Topically administered substances may interact with sulfhydryl or amino groups in the skin and react with sensitized lymphocytes to produce the rash characteristic of contact dermatitis. Other types of rashes may also result from the interaction of serum factors, drugs, and sensitized lymphocytes.

DIAGNOSIS AND TREATMENT OF ADVERSE DRUG REACTIONS

The manifestations of drug-induced diseases frequently resemble those of other diseases, and a given set of manifestations may be produced by different and dissimilar drugs. Recognition of the role of a drug or drugs in an illness depends on appreciation of the possible adverse reactions to drugs in any disease, on identification of the temporal relationship between drug administration and development of the illness, and on familiarity with the common manifestations of the drugs. A suspected adverse drug reaction developing after introduction of a new drug naturally implicates that drug; however, it is also important to remember that a drug interaction may be responsible. Thus, for example, a patient on a chronic stable warfarin dose may develop a bleeding complication after introduction of amiodarone; this does not reflect a direct reaction to amiodarone but rather its effect to inhibit warfarin metabolism. Many associations between particular drugs and specific reactions have been described, but there is always a “first time” for a novel association, and any drug should be suspected of causing an adverse effect if the clinical setting is appropriate.

Illness related to a drug's intended pharmacologic action is often more easily recognized than illness attributable to immune or other mechanisms. For example, side effects such as cardiac arrhythmias in patients receiving digitalis, hypoglycemia in patients given insulin, or bleeding in patients receiving anticoagulants are more readily related to a specific drug than are symptoms such as fever or rash, which may be caused by many drugs or by other factors.

Electronic listings of adverse drug reactions can be useful. However, exhaustive compilations often provide little sense of perspective in terms of frequency and seriousness, which can vary considerably among patients.

Eliciting a drug history from each patient is important for diagnosis. Attention must be directed to OTC drugs and herbal preparations as well as to prescription drugs. Each type can be responsible for adverse drug effects, and adverse interactions may occur between OTC drugs and prescribed drugs. Loss of efficacy of oral contraceptives or cyclosporine with concurrent use of St. John's wort (a P-glycoprotein inducer) is an example. In addition, it is common for patients to be cared for by several physicians, and duplicative, additive, antagonistic, or synergistic drug combinations may therefore be administered if the physicians are not aware of the patients' drug histories. Every physician should determine what drugs a patient has been taking, for the previous month or two ideally, before prescribing any medications. Medications stopped for inefficacy or adverse effects should be documented to avoid pointless and potentially dangerous reexposure. A frequently overlooked source of additional drug exposure is topical therapy; for example, a patient complaining of bronchospasm may not mention that an ophthalmic beta blocker is being used unless specifically asked. A history of previous adverse drug effects in patients is common. Since these patients have shown a predisposition to drug-induced illnesses, such a history should dictate added caution in prescribing new drugs.

Laboratory studies may include demonstration of serum antibody in some persons with drug allergies involving cellular blood elements, as in agranulocytosis, hemolytic anemia, and thrombocytopenia. For example, both quinine and quinidine can produce platelet agglutination *in vitro* in the presence of complement and the serum from a patient who has developed thrombocytopenia following use of this drug. Biochemical abnormalities such as G6PD deficiency, serum pseudocholinesterase level, or genotyping may also be useful in diagnosis, often after an adverse effect has occurred in the patient or a family member.

Once an adverse reaction is suspected, discontinuation of the suspected drug followed by disappearance of the reaction is presumptive evidence of a drug-induced illness. Confirming evidence may be sought by cautiously reintroducing the drug and seeing if the reaction reappears. However, that should be done only if confirmation would be useful in the future management of the patient and if the attempt would not entail undue risk. With concentration-dependent adverse reactions, lowering the dosage may cause the reaction to disappear, and raising it may cause the reaction to reappear. When the reaction is thought to be allergic, however, readministration of the drug may be hazardous, since anaphylaxis may develop.

If the patient is receiving many drugs when an adverse reaction is suspected, the drugs likeliest to be responsible can usually be identified; this should include both potential culprit agents as well as drugs that alter their elimination. All drugs may be discontinued at once or, if this is not practical, discontinued one at a time, starting with the ones most suspect, and the patient observed for signs of improvement. The time needed for a concentration-dependent adverse effect to disappear depends on the time required for the concentration to fall below the range associated with the adverse effect; that, in turn, depends on the initial blood level and on the rate of elimination or metabolism of the drug. Adverse effects of drugs with long half-lives or those not directly related to serum concentration may take a considerable time to disappear.

SUMMARY

Modern clinical pharmacology aims to replace empiricism in the use of drugs with therapy based on in-depth understanding of factors that determine an individual's response to drug treatment. Molecular pharmacology, pharmacokinetics, genetics, clinical trials, and the educated prescriber all contribute to this process. No drug response should ever be termed *idiosyncratic*; all responses have a mechanism whose understanding will help guide further therapy with that drug or successors. This rapidly expanding understanding of variability in drug

actions makes the process of prescribing drugs increasingly daunting for the practitioner. However, fundamental principles should guide this process:

- The benefits of drug therapy, however defined, should always outweigh the risk.
- The smallest dosage necessary to produce the desired effect should be used.
- The number of medications and doses per day should be minimized.
- Although the literature is rapidly expanding, accessing it is becoming easier; electronic tools to search databases of literature and unbiased opinion will become increasingly commonplace.
- Genetics play a role in determining variability in drug response and may become a part of clinical practice.
- Electronic medical record and pharmacy systems will increasingly incorporate prescribing advice, such as indicated medications not used; unindicated medications being prescribed; and potential dosing errors, drug interactions, or genetically determined drug responses.
- Prescribers should be particularly wary when adding or stopping specific drugs that are especially liable to provoke interactions and adverse reactions.
- Prescribers should use only a limited number of drugs, with which they are thoroughly familiar.

6e

Women's Health

Andrea Dunaif

The National Institutes of Health's Office of Research on Women's Health celebrated its twentieth anniversary in 2010 with a new strategic plan recognizing the study of the biologic basis of sex differences as a distinct scientific discipline. It has become clear that both sex chromosomes and sex hormones contribute to these differences. Indeed, it is recommended that the term *sex difference* be used for biologic processes that differ between males and females and the term *gender difference* be used for features related to social influences. The clinical discipline of women's health emphasizes greater attention to patient education and involvement in disease prevention and medical decision-making and has become a model for patient-centered health care.

DISEASE RISK: REALITY AND PERCEPTION

The leading causes of death are the same in women and men: (1) heart disease, and (2) cancer (Table 6e-1; Fig. 6e-1). The leading cause of cancer death, lung cancer, is the same in both sexes. Breast cancer is the second leading cause of cancer death in women, but it causes about 60% fewer deaths than does lung cancer. Men are substantially more likely to die from suicide and accidents than are women.

Women's risk for many diseases increases at menopause, which occurs at a median age of 51.4 years. In the industrialized world, women spend one-third of their lives in the postmenopausal period. Estrogen levels fall abruptly at menopause, inducing a variety of physiologic and metabolic responses. Rates of cardiovascular disease (CVD) increase and bone density begins to decrease rapidly after menopause. In the United States, women live on average about 5 years longer than men, with a life expectancy at birth in 2011 of 81.1 years compared with 76.3 years in men. Elderly women outnumber elderly men, so that age-related conditions such as hypertension have a female preponderance. However, the difference in life expectancy between men and women has decreased an average of 0.1 year every year since its peak of 7.8 years in 1979. If this convergence in mortality figures continues, it is projected that mortality rates will be similar by 2054.

Public awareness campaigns have resulted in a marked increase in the percentage of U.S. women knowing that CVD is the leading cause

of death in women. In 1997, the majority of U.S. women surveyed thought that cancer (35%) rather than heart disease (30%) was the leading cause of death in women (Fig. 6e-2). In 2012, these perceptions were reversed, with 56% of U.S. women surveyed recognizing that heart disease rather than cancer (24%) was the leading cause of death in women (Fig. 6e-2). Although awareness of heart disease has improved substantially among black and Hispanic women over this time period, these groups were 66% less likely than white women to recognize that heart disease is the leading cause of death in women.

Nevertheless, women younger than 65 years still consider breast cancer to be their leading health risk, despite the fact that death rates from breast cancer have been falling since the 1990s. In any specific decade of life, a woman's risk for breast cancer never exceeds 1 in 34. Although a woman's lifetime risk of developing breast cancer if she lives past 85 years is about 1 in 9, it is much more likely that she will die from CVD than from breast cancer. In other words, many elderly women have breast cancer but die from other causes. Similarly, a minority of women are aware that lung cancer is the leading cause of cancer death in women. Physicians are also less likely to recognize women's risk for CVD. Even in 2012, only 21% of U.S. women surveyed reported that their physicians had counseled them about their risk for heart disease. These misconceptions are unfortunate as they perpetuate inadequate attention to modifiable risk factors such as dyslipidemia, hypertension, and cigarette smoking.

SEX DIFFERENCES IN HEALTH AND DISEASE

ALZHEIMER'S DISEASE

(See also Chap. 448) Alzheimer's disease (AD) affects approximately twice as many women as men. Because the risk for AD increases with age, part of this sex difference is accounted for by the fact that women live longer than men. However, additional factors probably contribute to the increased risk for AD in women, including sex differences in brain size, structure, and functional organization. There is emerging evidence for sex-specific differences in gene expression, not only for genes on the X and Y chromosomes but also for some autosomal genes. Estrogens have pleiotropic genomic and nongenomic effects on the central nervous system, including neurotrophic actions in key areas involved in cognition and memory. Women with AD have lower endogenous estrogen levels than do women without AD. These observations have led to the hypothesis that estrogen is neuroprotective.

TABLE 6e-1 DEATHS AND PERCENTAGE OF TOTAL DEATHS FOR THE LEADING CAUSES OF DEATH BY SEX IN THE UNITED STATES IN 2010

Cause of Death	Female			Male		
	Rank	Deaths	Deaths, %	Rank	Deaths	Deaths, %
Diseases of heart	1	290,305	23.5	1	307,384	24.9
Malignant neoplasms	2	273,706	22.1	2	301,037	24.4
Cerebrovascular diseases	3	77,109	6.2	5	52,367	4.2
Chronic lower respiratory diseases	4	72,657	5.9	4	65,423	5.3
Alzheimer's disease	5	58,130	4.7	8	25,364	2.1
Accidents (unintentional injuries)	6	44,938	3.6	3	75,921	6.2
Diabetes mellitus	7	33,581	2.7	6	35,490	2.9
Influenza and pneumonia	8	26,482	2.1	10	23,615	1.9
Nephritis, nephrotic syndrome, and nephrosis	9	25,611	2.1	9	24,865	2.0
Septicemia	10	18,743	1.5	12	16,069	1.3
Essential hypertension and hypertensive renal disease	11	15,788	1.3	16	10,846	0.9
Chronic liver disease and cirrhosis	12	11,105	0.9	11	20,798	1.7
Parkinson's disease	13	9161	0.7	14	12,871	1.0
Intentional self-harm (suicide)	14	8087	0.7	7	30,277	2.5
Pneumonitis due to solids and liquids	15	7803	0.6	17	9208	0.7

Note: Category titles beginning with "other" or "all other" are not ranked when determining the leading causes of death.

Source: Data from Centers for Disease Control and Prevention: National Vital Statistics Reports, Vol. 61, No. 4, May 8, 2013, Table 12, http://www.cdc.gov/nchs/data/nvsr/nvsr61/nvsr61_04.pdf.

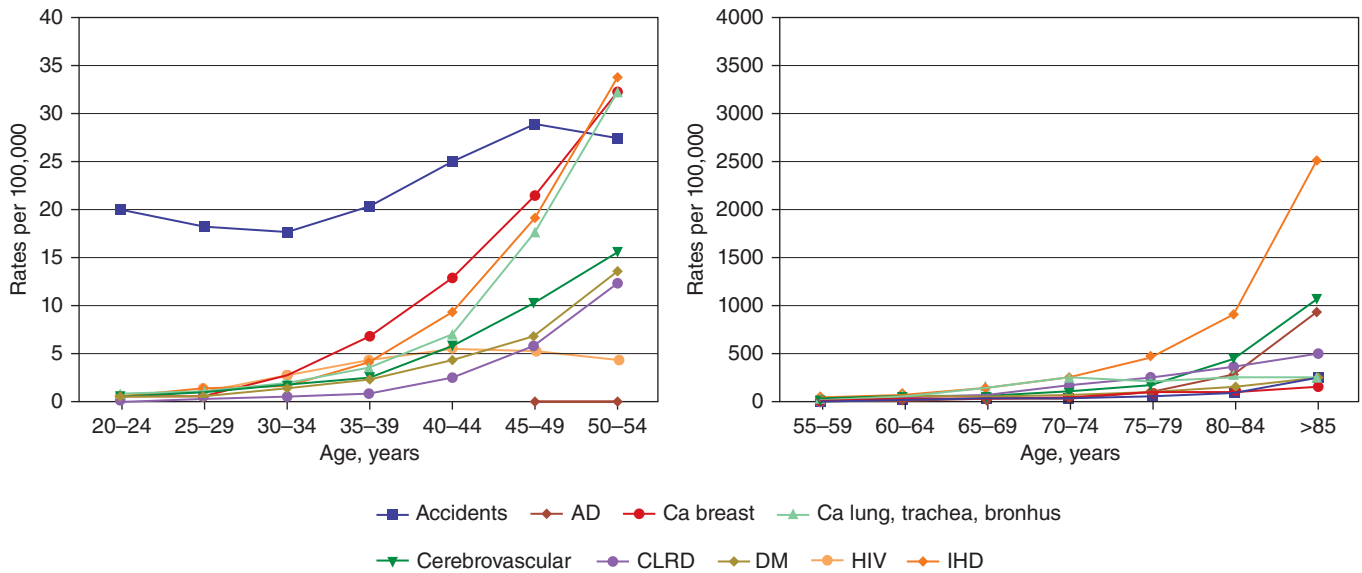


FIGURE 6e-1 Death rates per 100,000 population for 2007 by 5-year age groups in U.S. women. Note that the scale of the y axis is increased in the graph on the right compared with that on the left. Accidents and HIV/AIDS are the leading causes of death in young women 20–34 years of age. Accidents, breast cancer, and ischemic heart disease (IHD) are the leading causes of death in women 35–49 years of age. IHD becomes the leading cause of death in women beginning at age 50 years. In older women, IHD remains the leading cause of death, cerebrovascular disease becomes the second leading cause of death, and lung cancer is the leading cause of cancer-related deaths. At age 85 years and beyond, Alzheimer's disease (AD) becomes the third leading cause of death. Ca, cancer; CLRD, chronic lower respiratory disease; DM, diabetes mellitus. (Data adapted from Centers for Disease Control and Prevention, http://www.cdc.gov/nchs/data/dvs/MortFinal2007_WorkTable210R.pdf.)

Some studies have suggested that estrogen administration improves cognitive function in nondemented postmenopausal women as well as in women with AD, and several observational studies have suggested that postmenopausal hormone therapy (HT) may decrease the risk of AD. However, HT placebo-controlled trials have found no improvement in disease progression or cognitive function in women with AD. Further, the Women's Health Initiative Memory Study (WHIMS), an ancillary study in the Women's Health Initiative (WHI), found no benefit compared with placebo of estrogen alone [combined continuous equine estrogen (CEE), 0.625 mg daily] or estrogen with progestin [CEE, 0.625 mg daily, and medroxyprogesterone acetate (MPA), 2.5 mg daily] on cognitive function or the development of dementia in women ≥ 65 years. Indeed, there was a significantly increased risk for both dementia and mild cognitive impairment in women receiving HT. However, preliminary findings from the Kronos Early Estrogen Prevention Study (KEEPS), a randomized clinical trial of early initiation of HT after menopause that compared CEE 0.45 mg daily, 50 μ g of weekly transdermal estradiol (both estrogen arms included cyclic

oral micronized progesterone 200 mg daily for 12 days each month), or placebo, found no adverse effects on cognitive function.

CARDIOVASCULAR DISEASE AND STROKE

(See also Chap. 293) There are major sex differences in CVD, the leading cause of death in men and women in developed countries. A greater number of U.S. women than men die annually of CVD and stroke. Deaths from CVD have decreased markedly in men since 1980, whereas CVD deaths only began to decrease substantially in women beginning in 2000. However, in middle-aged women, the prevalence rates of both coronary heart disease (CHD) and stroke have increased in the 1999–2004 National Health and Nutrition Survey (NHANES) compared to the 1988–1994 NHANES, whereas prevalence rates have decreased or remained unchanged, respectively, in men. These increases were paralleled by an increasing prevalence of abdominal obesity and other components of metabolic syndrome in women.

Sex steroids have major effects on the cardiovascular system and lipid metabolism. Estrogen increases high-density lipoprotein (HDL)

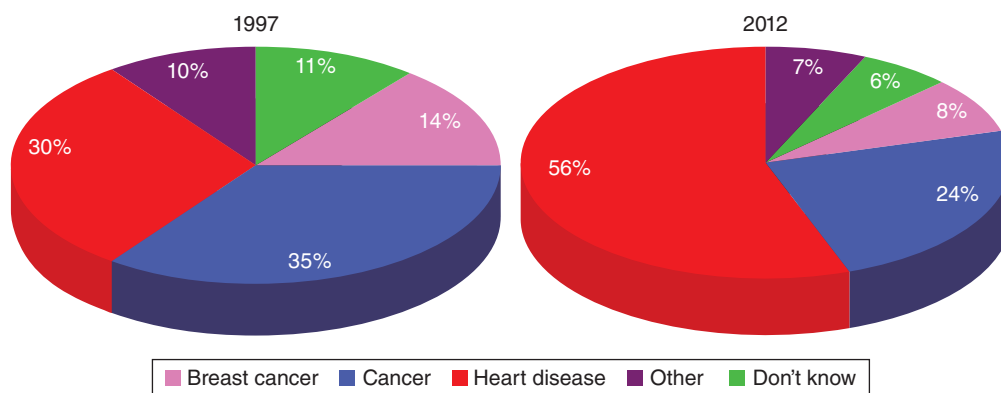


FIGURE 6e-2 Changes in perceived leading causes of death among women surveyed in 1997 compared with those surveyed in 2012. In 1997, cancer was cited as the leading cause of death in women, not heart disease. In 2012, this trend had reversed. The rate of awareness that heart disease is the leading cause of death in women was significantly higher in 2012 (56% vs 30%, $p < .001$) than in 1997. (Data adapted from L Mosca et al: *Circulation* 127:1254, 2013.)

and lowers low-density lipoprotein (LDL), whereas androgens have the opposite effect. Estrogen has direct vasodilatory effects on the vascular endothelium, enhances insulin sensitivity, and has antioxidant and anti-inflammatory properties. There is a striking increase in CHD after both natural and surgical menopause, suggesting that endogenous estrogens are cardioprotective. Women also have longer QT intervals on electrocardiograms, and this increases their susceptibility to certain arrhythmias.

CHD presents differently in women, who are usually 10–15 years older than their male counterparts and are more likely to have comorbidities such as hypertension, congestive heart failure, and diabetes mellitus (DM). In the Framingham study, angina was the most common initial symptom of CHD in women, whereas myocardial infarction (MI) was the most common initial presentation in men. Women more often have atypical symptoms such as nausea, vomiting, indigestion, and upper back pain. Although awareness that heart disease is the leading cause of death in women has nearly doubled over the last 15 years, women remain less aware that its symptoms are often atypical, and they are less likely to contact 9-1-1 when they experience such symptoms.

Women with MI are more likely to present with cardiac arrest or cardiogenic shock, whereas men are more likely to present with ventricular tachycardia. Further, younger women with MI are more likely to die than are men of similar age. However, this mortality gap has decreased substantially in recent years because younger women have experienced greater improvements in survival after MI than men (Fig. 6e-3). The improvement in survival is due largely to a reduction in comorbidities, suggesting a greater attention to modifiable risk factors in women.

Nevertheless, physicians are less likely to suspect heart disease in women with chest pain and less likely to perform diagnostic and therapeutic cardiac procedures in women. Women are less likely to receive therapies such as angioplasty, thrombolytic therapy, coronary artery bypass grafts (CABGs), beta blockers, and aspirin. There are also sex differences in outcomes when women with CHD do receive therapeutic interventions. Women undergoing CABG surgery have more advanced disease, a higher perioperative mortality rate, less relief of angina, and less graft patency; however, 5- and 10-year survival rates are similar. Women undergoing percutaneous transluminal coronary angioplasty have lower rates of initial angiographic and clinical success than men, but they also have a lower rate of restenosis and a better long-term outcome. Women may benefit less and have more frequent serious bleeding complications from thrombolytic therapy compared with men. Factors such as older age, more comorbid conditions,

smaller body size, and more severe CHD in women at the time of events or procedures account in part for the observed sex differences.

Elevated cholesterol levels, hypertension, smoking, obesity, low HDL cholesterol levels, DM, and lack of physical activity are important risk factors for CHD in both men and women. Total triglyceride levels are an independent risk factor for CHD in women but not in men. Low HDL cholesterol and DM are more important risk factors for CHD in women than in men. Smoking is an important risk factor for CHD in women—it accelerates atherosclerosis, exerts direct negative effects on cardiac function, and is associated with an earlier age of menopause. Cholesterol-lowering drugs are equally effective in men and women for primary and secondary prevention of CHD. However, because of perceptions that women are at lower risk for CHD, they receive fewer interventions for modifiable risk factors than do men. In contrast to men, randomized trials showed that aspirin was not effective in the primary prevention of CHD in women; it did significantly reduce the risk of ischemic stroke.

The sex differences in CHD prevalence, beneficial biologic effects of estrogen on the cardiovascular system, and reduced risk for CHD in observational studies led to the hypothesis that HT was cardioprotective. However, the WHI, which studied more than 16,000 women on CEE plus MPA or placebo and more than 10,000 women with hysterectomy on CEE alone or placebo, did not demonstrate a benefit of HT for the primary or secondary prevention of CHD. In addition, CEE plus MPA was associated with an increased risk for CHD, particularly in the first year of therapy, whereas CEE alone neither increased nor decreased CHD risk. Both CEE plus MPA and CEE alone were associated with an increased risk for ischemic stroke.

In the WHI, there was a suggestion of a reduction in CHD risk in women who initiated HT closer to menopause. This finding suggests that the time at which HT is initiated is critical for cardioprotection. According to this “timing” hypothesis, HT has differential effects, depending on the stage of atherosclerosis; adverse effects are seen with advanced, unstable lesions. A recent study using data from the Danish Osteoporosis Prevention Study (DOPS), an open-label randomized trial of triphasic oral estradiol compared with no treatment in recently menopausal or perimenopausal women (a cyclic oral synthetic progestin, norethisterone acetate, was added in women who had a uterus), found significantly reduced mortality and CVD after 10 years of HT. However, DOPS was designed to investigate HT for the primary prevention of osteoporotic bone fractures, and CVD outcomes were not prespecified endpoints. Further, there were relatively few CVD events in the study groups.

KEEPS was designed to directly test the “timing” hypothesis. Seven hundred twenty-seven recently menopausal women age 42–58 years (mean 52.7 years) were randomized to oral CEE (lower dose than WHI), transdermal estradiol, or placebo for 4 years; both estrogen arms included oral cyclical micronized progesterone (see above section on AD for dosing details). There were no significant beneficial or deleterious effects on the progression of atherosclerosis by computed tomography assessment of coronary artery calcification in either HT arm. Adverse events including stroke, MI, venous thromboembolism, and breast cancer were not increased in the HT arms compared with the placebo arm. There were improvements in hot flashes, night sweats, mood, sexual function, and bone density in the HT arms. This relatively small study does not suggest that early HT administration, transdermally or orally, reduces atherosclerosis. However, the study suggests that short-term HT may be safely administered for symptom relief in recently menopausal women. [HT is discussed further in Chap. 413.](#)

DIABETES MELLITUS

(See also [Chap. 417](#)) Women are more sensitive to insulin than men are. Despite this, the prevalence of type 2 DM is similar in men and women. There is a sex difference in the relationship between endogenous androgen levels and DM risk. Higher bioavailable testosterone levels are associated with increased risk in women, whereas lower bioavailable testosterone levels are associated with increased risk in men. Polycystic ovary syndrome and gestational DM—common

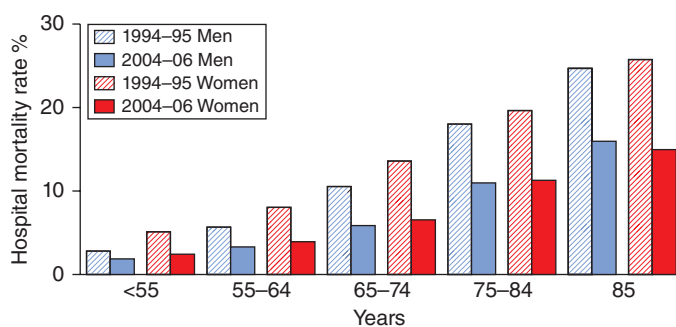


FIGURE 6e-3 Hospital mortality rates in men and women for acute myocardial infarction (MI) in 1994–1995 compared with 2004–2006. Women younger than age 65 years had substantially greater mortality than men of similar age in 1994–1995. Mortality rates declined markedly for both sexes across all age groups in 2004–2006 compared with 1994–1995. However, there was a more striking decrease in mortality in women younger than age 75 years compared with men of similar age. The mortality rate reduction was largest in women less than age 55 years (52.9%) and lowest in men of similar age (33.3%). (Data adapted from V Vaccarino et al: *Arch Intern Med* 169:1767, 2009.)

conditions in premenopausal women—are associated with a significantly increased risk for type 2 DM. Premenopausal women with DM lose the cardioprotective effect of female sex and have rates of CHD identical to those in males. These women have impaired endothelial function and reduced coronary vasodilatory responses, which may predispose to cardiovascular complications. Among individuals with DM, women have a greater risk for MI than do men. Women with DM are more likely to have left ventricular hypertrophy. Women with DM receive less aggressive treatment for modifiable CHD risk factors than men with DM. In the WHI, CEE plus MPA significantly reduced the incidence of DM, whereas with CEE alone, there was only a trend toward decreased DM incidence.

HYPERTENSION

(See also Chap. 298) After age 60, hypertension is more common in U.S. women than in men, largely because of the high prevalence of hypertension in older age groups and the longer survival of women. Isolated systolic hypertension is present in 30% of women >60 years old. Sex hormones affect blood pressure. Both normotensive and hypertensive women have higher blood pressure levels during the follicular phase than during the luteal phase. In the Nurses' Health Study, the relative risk of hypertension was 1.8 in current users of oral contraceptives, but this risk is lower with the newer low-dose contraceptive preparations. HT is not associated with hypertension. Among secondary causes of hypertension, there is a female preponderance of renal artery fibromuscular dysplasia.

The benefits of treatment for hypertension have been dramatic in both women and men. A meta-analysis of the effects of hypertension treatment, the Individual Data Analysis of Antihypertensive Intervention Trial, found a reduction of risk for stroke and for major cardiovascular events in women. The effectiveness of various antihypertensive drugs appears to be comparable in women and men; however, women may experience more side effects. For example, women are more likely to develop cough with angiotensin-converting enzyme inhibitors.

AUTOIMMUNE DISORDERS

(See also Chap. 377e) Most autoimmune disorders occur more commonly in women than in men; they include autoimmune thyroid and liver diseases, lupus, rheumatoid arthritis (RA), scleroderma, multiple sclerosis (MS), and idiopathic thrombocytopenic purpura. However, there is no sex difference in the incidence of type 1 DM, and ankylosing spondylitis occurs more commonly in men. Women may be more resistant to bacterial infections than men. Sex differences in both immune responses and adverse reactions to vaccines have been reported. For example, there is a female preponderance of postvaccination arthritis.

Adaptive immune responses are more robust in women than in men; this may be explained by the stimulatory actions of estrogens and the inhibitory actions of androgens on the cellular mediators of immunity. Consistent with an important role for sex hormones, there is variation in immune responses during the menstrual cycle, and the activity of certain autoimmune disorders is altered by castration or pregnancy (e.g., RA and MS may remit during pregnancy). Nevertheless, the majority of studies show that exogenous estrogens and progestins in the form of HT or oral contraceptives do not alter autoimmune disease incidence or activity. Exposure to fetal antigens, including circulating fetal cells that persist in certain tissues, has been speculated to increase the risk of autoimmune responses. There is clearly an important genetic component to autoimmunity, as indicated by the familial clustering and HLA association of many such disorders. X chromosome genes also contribute to sex differences in immunity. Indeed, nonrandom X chromosome inactivation may be a risk factor for autoimmune diseases.

HIV INFECTION

(See also Chap. 226) Women account for almost 50% of the 34 million persons infected with HIV-1 worldwide. AIDS is an important cause of death in younger women (Fig. 6e-1). Heterosexual contact

with an at-risk partner is the fastest-growing transmission category, and women are more susceptible to HIV infection than are men. This increased susceptibility is accounted for in part by an increased prevalence of sexually transmitted diseases in women. Some studies have suggested that hormonal contraceptives may increase the risk of HIV transmission. Progesterone has been shown to increase susceptibility to infection in nonhuman primate models of HIV. Women are also more likely to be infected by multiple variants of the virus than are men. Women with HIV have more rapid decreases in their CD4 cell counts than do men. Compared with men, HIV-infected women more frequently develop candidiasis, but Kaposi's sarcoma is less common than it is in men. Women have more adverse reactions, such as lipodystrophy, dyslipidemia, and rash, with antiretroviral therapy than do men. This observation is explained in part by sex differences in the pharmacokinetics of certain antiretroviral drugs, resulting in higher plasma concentrations in women.

OBESITY

(See also Chap. 416) The prevalence of both obesity (body mass index ≥ 30 kg/m²) and abdominal obesity (waist circumference ≥ 88 cm in women) is higher in U.S. women than in men. However, between 1999 and 2008, the prevalence of obesity increased significantly in men but not in women. The prevalence of abdominal obesity increased over this time period in both sexes. More than 80% of patients who undergo bariatric surgery are women. Pregnancy and menopause are risk factors for obesity.

There are major sex differences in body fat distribution. Women characteristically have gluteal and femoral or gynoid pattern of fat distribution, whereas men typically have a central or android pattern. Women have more subcutaneous fat than men. In women, endogenous androgen levels are positively associated with abdominal obesity, and androgen administration increases visceral fat. In contrast, there is an inverse relationship between endogenous androgen levels and abdominal obesity in men. Further, androgen administration decreases visceral fat in these obese men. The reasons for these sex differences in the relationship between visceral fat and androgens are unknown. Studies in humans also suggest that sex steroids play a role in modulating food intake and energy expenditure.

In men and women, abdominal obesity characterized by increased visceral fat is associated with an increased risk for CVD and DM. Obesity increases a woman's risk for certain cancers, in particular postmenopausal breast and endometrial cancer, in part because adipose tissue provides an extragonadal source of estrogen through aromatization of circulating adrenal and ovarian androgens, especially the conversion of androstenedione to estrone. Obesity increases the risk of infertility, miscarriage, and complications of pregnancy.

OSTEOPOROSIS

(See also Chap. 425) Osteoporosis is about five times more common in postmenopausal women than in age-matched men, and osteoporotic hip fractures are a major cause of morbidity in elderly women. Men accumulate more bone mass and lose bone more slowly than do women. Sex differences in bone mass are found as early as infancy. Calcium intake, vitamin D, and estrogen all play important roles in bone formation and bone loss. Particularly during adolescence, calcium intake is an important determinant of peak bone mass. Vitamin D deficiency is surprisingly common in elderly women, occurring in >40% of women living in northern latitudes. Receptors for estrogens and androgens have been identified in bone. Estrogen deficiency is associated with increased osteoclast activity and a decreased number of bone-forming units, leading to net bone loss. The aromatase enzyme, which converts androgens to estrogens, is also present in bone. Estrogen is an important determinant of bone mass in men (derived from the aromatization of androgens) as well as in women.

PHARMACOLOGY

On average, women have lower body weights, smaller organs, a higher percentage of body fat, and lower total-body water than men. There are also important sex differences in drug action and metabolism that are

not accounted for by these differences in body size and composition. Sex steroids alter the binding and metabolism of a number of drugs. Further, menstrual cycle phase and pregnancy can alter drug action. Two-thirds of cases of drug-induced torsades des pointes, a rare, life-threatening ventricular arrhythmia, occur in women because they have a longer, more vulnerable QT interval. These drugs, which include certain antihistamines, antibiotics, antiarrhythmics, and antipsychotics, can prolong cardiac repolarization by blocking cardiac voltage-gated potassium channels. Women require lower doses of neuroleptics to control schizophrenia. Women awaken from anesthesia faster than do men given the same doses of anesthetics. Women also take more medications than men, including over-the-counter formulations and supplements. The greater use of medications combined with these biologic differences may account for the reported higher frequency of adverse drug reactions in women than in men.

PSYCHOLOGICAL DISORDERS

(See also Chap. 466) Depression, anxiety, and affective and eating disorders (bulimia and anorexia nervosa) are more common in women than in men. Epidemiologic studies from both developed and developing nations consistently find major depression to be twice as common in women as in men, with the sex difference becoming evident in early adolescence. Depression occurs in 10% of women during pregnancy and in 10–15% of women during the postpartum period. There is a high likelihood of recurrence of postpartum depression with subsequent pregnancies. The incidence of major depression diminishes after age 45 years and does not increase with the onset of menopause. Depression in women appears to have a worse prognosis than does depression in men; episodes last longer, and there is a lower rate of spontaneous remission. Schizophrenia and bipolar disorders occur at equal rates in men and women, although there may be sex differences in symptoms.

Both biologic and social factors account for the greater prevalence of depressive disorders in women. Men have higher levels of the neurotransmitter serotonin. Sex steroids also affect mood, and fluctuations during the menstrual cycle have been linked to symptoms of premenstrual syndrome. Sex hormones differentially affect the hypothalamic-pituitary-adrenal responses to stress. Testosterone appears to blunt cortisol responses to corticotropin-releasing hormone. Both low and high levels of estrogen can activate the hypothalamic-pituitary-adrenal axis.

SLEEP DISORDERS

(See also Chap. 38) There are striking sex differences in sleep and its disorders. During sleep, women have an increased amount of slow-wave activity, differences in timing of delta activity, and an increase in the number of sleep spindles. Testosterone modulates neural control of breathing and upper airway mechanics. Men have a higher prevalence of sleep apnea. Testosterone administration to hypogonadal men as well as to women increases apneic episodes during sleep. Women with the hyperandrogenic disorder polycystic ovary syndrome have an increased prevalence of obstructive sleep apnea, and apneic episodes are positively correlated with their circulating testosterone levels. In contrast, progesterone accelerates breathing, and in the past, progestins were used for treatment of sleep apnea.

SUBSTANCE ABUSE AND TOBACCO

(See also Chaps. 467 and 470) Substance abuse is more common in men than in women. However, one-third of Americans who

suffer from alcoholism are women. Women alcoholics are less likely to be diagnosed than men. A greater proportion of men than women seek help for alcohol and drug abuse. Men are more likely to go to an alcohol or drug treatment facility, whereas women tend to approach a primary care physician or mental health professional for help under the guise of a psychosocial problem. Late-life alcoholism is more common in women than in men. On average, alcoholic women drink less than alcoholic men but exhibit the same degree of impairment. Blood alcohol levels are higher in women than in men after drinking equivalent amounts of alcohol, adjusted for body weight. This greater bioavailability of alcohol in women is due to both the smaller volume of distribution and the slower gastric metabolism of alcohol secondary to lower activity of gastric alcohol dehydrogenase than is the case in men. In addition, alcoholic women are more likely to abuse tranquilizers, sedatives, and amphetamines. Women alcoholics have a higher mortality rate than do nonalcoholic women and alcoholic men. Women also appear to develop alcoholic liver disease and other alcohol-related diseases with shorter drinking histories and lower levels of alcohol consumption. Alcohol abuse also poses special risks to a woman, adversely affecting fertility and the health of the baby (fetal alcohol syndrome). Even moderate alcohol use increases the risk of breast cancer, hypertension, and stroke in women.

More men than women smoke tobacco, but this sex difference continues to decrease. Women have a much larger burden of smoking-related disease. Smoking markedly increases the risk of CVD in premenopausal women and is also associated with a decrease in the age of menopause. Women who smoke are more likely to develop chronic obstructive pulmonary disease and lung cancer than men and at lower levels of tobacco exposure. Postmenopausal women who smoke have lower bone density than women who never smoked. Smoking during pregnancy increases the risk of preterm deliveries and low birth weight infants.

VIOLENCE AGAINST WOMEN

More than one in three women in the United States have experienced rape, physical violence, and/or stalking by an intimate partner. Adult women are much more likely to be raped by a spouse, ex-spouse, or acquaintance than by a stranger. Domestic or intimate partner violence is a leading cause of death among young women. Domestic violence may be an unrecognized feature of certain clinical presentations, such as chronic abdominal pain, headaches, and eating disorders, in addition to more obvious manifestations such as trauma. Intimate partner violence is an important risk factor for depression, substance abuse, and suicide in women. Screening instruments can accurately identify women experiencing intimate partner violence. Such screening by health care providers is acceptable to women in settings ensuring adequate privacy and safety.

SUMMARY

Women's health is now a mature discipline, and the importance of sex differences in biologic processes is well recognized. There has been a striking reduction in the excess mortality rate from MI in younger women. Nevertheless, ongoing misperceptions about disease risk, not only among women but also among their physicians, result in inadequate attention to modifiable risk factors. Research into the fundamental mechanisms of sex differences will provide important biologic insights. Further, those insights will have an impact on both women's and men's health.

7e

Men's Health

Shalender Bhasin, Shehzad Basaria

The emergence of men's health as a distinct discipline within internal medicine is founded on the evidence that men and women differ across their life span in their susceptibility to disease, in the clinical manifestations of the disease, and in their response to treatment. Furthermore, men and women weigh the health consequences of illness differently and have different motivations for seeking care. Men and women experience different types of disparities in access to health care services and in the manner in which health care is delivered to them because of a complex array of socioeconomic and cultural factors. Attitudinal and institutional barriers to accessing care, fear and embarrassment due to the perception by some that it is not manly to seek medical help, and reticence on the part of patients and physicians to discuss issues related to sexuality, drug use, and aging have heightened the need for programs tailored to address the specific health needs of men.

Sex differences in disease prevalence, susceptibility, and clinical manifestations of disease were discussed in [Chap. 6e](#) ("Women's Health"). It is notable that the two leading causes of death in both men and women—heart disease and cancer—are the same. However, men have a higher prevalence of neurodevelopmental and degenerative disorders; substance abuse disorders, including the use of performance-enhancing drugs and alcohol dependence; diabetes; and cardiovascular disease; and women have a higher prevalence of autoimmune disorders, depression, rheumatologic disorders, and osteoporosis. Men are substantially more likely to die from accidents, suicides, and homicides than women. Among men 15–34 years of age, unintentional injuries, homicides, and suicides account for over three-fourths of all deaths. Among men 35–64 years of age, heart disease, cancer, and unintentional injuries are the leading causes of death. Among men 65 years of age or older, heart disease, cancer, lower respiratory tract infections, and stroke are the major causes of death.

The biologic bases of sex differences in disease susceptibility, progression, and manifestation remain incompletely understood and are likely multifactorial. Undoubtedly, sex-specific differences in the genetic architecture and circulating sex hormones influence disease phenotype; additionally, epigenetic effects of sex hormones during fetal life, early childhood, and pubertal development may imprint sexual and nonsexual behaviors, body composition, and disease susceptibility. Reproductive load and physiologic changes during pregnancy, including profound hormonal and metabolic shifts and microchimerism (transfer of cells from the mother to the fetus and from the fetus to the mother), may affect disease susceptibility and disease severity in women. Sociocultural norms of child-rearing practices, societal expectations of gender roles, and the long-term economic impact of these practices and gender roles also may affect disease risk and its clinical manifestation. The trajectories of age-related changes in sex hormones during the reproductive and postreproductive years vary substantially between men and women and may influence the sex differences in the temporal evolution of age-related conditions such as osteoporosis, breast cancer, and autoimmune disease.

In a reflection of the growing attention to issues related to men's health, health clinics focused on the health problems of men are being established with increasing frequency. Although the major threats to men's health have not changed—heart disease, cancer, and unintentional injury continue to dominate the list of major medical causes of morbidity and mortality in men—the men who attend men's health clinics do so largely for sexual, reproductive, and urologic health concerns involving common conditions such as androgen deficiency syndromes, age-related decline in testosterone levels, sexual dysfunction, muscle dysmorphia and anabolic-androgenic steroid use, lower urinary tract symptoms, and medical complications of prostate cancer therapy, which are the focus of this chapter. Additionally, new categories of body image disorders have emerged in men that had not been recognized until the 1980s, such as body dysmorphia syndrome and

the use of performance-enhancing drugs to increase muscularity and lean appearance. Although menopause in women has been the subject of intense investigation for more than five decades, the issues that are specific to men's health are just beginning to gain the attention that they deserve because of their high prevalence and impact on overall health, well-being, and quality of life.

AGING-RELATED CHANGES IN MALE REPRODUCTIVE FUNCTION (SEE CHAP. 411)

A number of studies have established that testosterone concentrations decrease with advancing age. This age-related decline starts in the third decade of life and progresses thereafter ([Fig. 7e-1](#)). Low total and bioavailable testosterone concentrations are associated with decreased skeletal muscle mass and strength, higher visceral fat mass, insulin resistance, and increased risk of coronary artery disease and mortality ([Table 7e-1](#)). Most studies suggest that these symptoms and signs develop with total testosterone levels below 320 ng/dL and free testosterone levels below 64 pg/mL in older men. Testing for low testosterone in older men should be limited to those with symptoms or signs attributable to androgen deficiency.

Testosterone therapy of healthy older men with low testosterone increases lean body mass, grip strength, and self-reported physical function ([Fig. 7e-2](#)). Testosterone therapy also increases vertebral but not femoral bone mineral density. In men with sexual dysfunction and low testosterone levels, testosterone therapy improves libido, but effects on erectile function and response to selective phosphodiesterase inhibitors are variable ([Chap. 67](#)). As discussed in [Chap. 411](#), there is concern that testosterone therapy may stimulate the growth of prostate cancers.

Sexual Dysfunction (See [Chap. 67](#)) Various forms of sexual dysfunction are a major motivating factor for men seeking care at men's health clinics. The landmark descriptions of the human sexual response cycle by Masters and Johnson, demonstrating that men and women display predictable physiologic responses after sexual stimulation, provided the basis for rational classification of human sexual disorders. Accordingly, sexual disorders have been classified into four categories depending on phase of sexual response cycle in which the abnormality exists:

1. Hypoactive sexual desire disorder
2. Erectile dysfunction
3. Ejaculatory and orgasmic disorders
4. Disorders of pain

Classification of the patient's disorder into these categories is important because the etiologic factors, diagnostic tests, and therapeutic

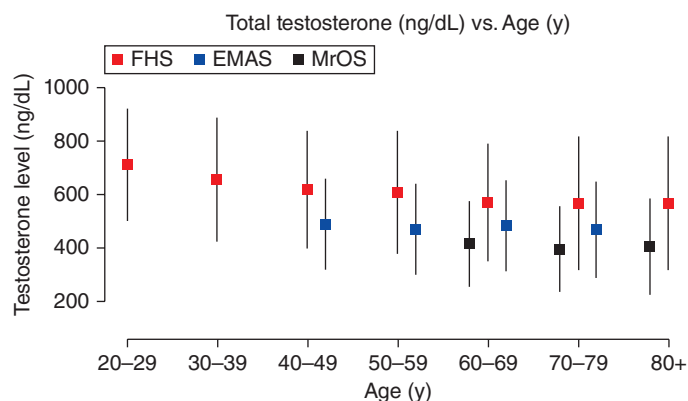


FIGURE 7e-1 Age-related decline in total testosterone levels. Total testosterone levels measured using liquid chromatography tandem mass spectrometry in men of the Framingham Heart Study (FHS), the European Male Aging Study (EMAS), and the Osteoporotic Fractures in Men Study (MrOS). (Reproduced with permission from S Bhasin et al: *J Clin Endocrinol Metab* 96:2430, 2011.)

TABLE 7e-1 ASSOCIATION OF TESTOSTERONE LEVELS WITH OUTCOMES IN OLDER MEN

1. Positively associated with:
• Muscle mass and muscle strength
• Physical function
• Sexual desire
• Bone mineral density, bone geometry, and volumetric bone mineral density
2. Negatively associated with:
• Coronary artery disease
• Visceral fat
• Diabetes mellitus
• Metabolic syndrome
• Mortality
• Falls and fracture risk
• Frailty
3. Not associated with:
• Lower urinary tract symptoms
• Erectile dysfunction
• Dementia
• Major depression

strategies vary for each class of sexual disorder. Historically, the classification and nomenclature for sexual disorders used criteria identified in the *Diagnostic and Statistical Manual of Mental Disorders* (DSM), based on the erroneous belief that sexual disorders in men are largely psychogenic in their origin. However, the recognition of erectile dysfunction as a manifestation of systemic disease and the availability of easy-to-use oral selective phosphodiesterase-5 inhibitors have placed sexual disorders in men within the purview of the primary care provider.

MUSCLE DYSMORPHIA SYNDROME IN MEN: A BODY IMAGE DISORDER

Muscle dysmorphia is a form of body image disorder characterized by a pathologic preoccupation with muscularity and leanness. The men with muscle dysmorphia express a strong desire to be more muscular and lean. These men describe shame and embarrassment about their body size and shape and often report adverse symptoms such as dissatisfaction with appearance, preoccupation with bodybuilding and muscularity, and functional impairment. Patients with muscle dysmorphia also report higher rates of mood and anxiety disorders, as well as obsessive and compulsive behaviors. These men often experience impairment of social and occupational functioning.

Patients with muscle dysmorphia syndrome—nearly all men—are almost always engaged in weightlifting and body building and are more likely to use performance-enhancing drugs, especially anabolic-androgenic steroids. Muscle dysmorphia disorder predisposes men to an increased risk of disease due to the combined interactive effects of the intensity of physical exercise, the use of performance-enhancing drugs, and other lifestyle factors associated with weightlifting and the use of performance-enhancing drugs. No randomized trials of any treatment modalities have been conducted; anecdotally, behavioral and cognitive therapies have been tried with varying degrees of success.

Anabolic-Androgenic Steroid Abuse by Athletes and Recreational Body-Builders

The illicit use of anabolic-androgenic steroids (AAS) to enhance athletic performance first surfaced in the 1950s among powerlifters and spread rapidly to other sports and to professional as well as high school athletes and recreational bodybuilders. In the early 1980s, the use of AAS spread beyond the athletic community into the general population. As many as 3 million Americans, most of them men, have likely used these compounds. Most AAS users are not athletes, but rather recreational weightlifters who use these drugs to look lean and more muscular.

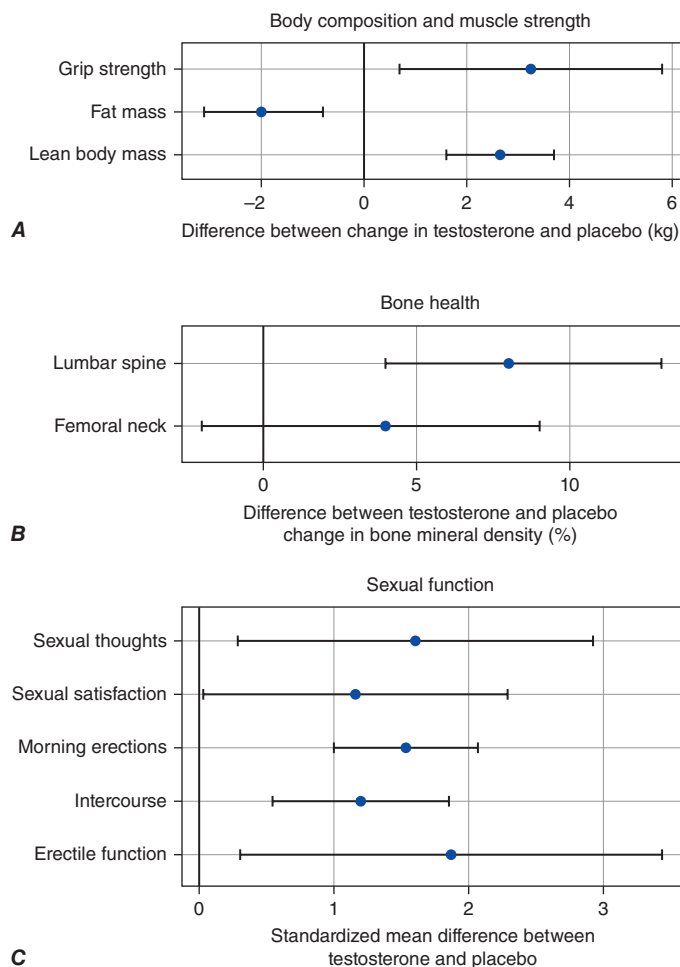


FIGURE 7e-2 The effects of testosterone therapy on body composition, muscle strength, bone mineral density, and sexual function in intervention trials. The point estimates and the associated 95% confidence intervals are shown. **A.** The effects of testosterone therapy on lean body mass, grip strength, and fat mass in a meta-analysis of randomized trials. (Data derived from S Bhasin et al: *Nat Clin Pract Endocrinol Metab* 2:146, 2006.) **B.** The effects of testosterone therapy on lumbar and femoral bone mineral density in a meta-analysis of randomized trials. (Data derived from a meta-analysis by MJ Tracz et al: *J Clin Endocrinol Metab* 91:2011, 2006.) **C.** The effects of testosterone therapy on measures of sexual function in men with baseline testosterone less than 10 nmol/L (290 ng/dL). (Data derived from a meta-analysis by AM Isidori et al: *Clin Endocrinol (Oxf)* 63:381, 2005.) (Reproduced with permission from M Spitzer et al: *Nat Rev Endocrinol* 9:414, 2013.)

The most commonly used AAS include testosterone esters, nandrolone, stanozolol, methandienone, and methenolone. AAS users generally use increasing doses of multiple steroids in a practice known as stacking.

The adverse effects of long-term AAS abuse remain poorly understood. Most of the information about the adverse effects of AAS has emerged from case reports, uncontrolled studies, or clinical trials that used replacement doses of testosterone (Table 7e-2). Of note, AAS users may administer 10–100 times the replacement doses of testosterone over many years, making it unjustifiable to extrapolate from trials using replacement doses. A substantial fraction of AAS users also use other drugs that are perceived to be muscle-building or performance-enhancing, such as growth hormone; erythropoiesis-stimulating agents; insulin; and stimulants such as amphetamine, clenbuterol, cocaine, ephedrine, and thyroxine; and drugs perceived to reduce adverse effects such as human chorionic gonadotropin, aromatase inhibitors, or estrogen antagonists. The men who abuse AAS are

TABLE 7e-2 POTENTIAL ADVERSE EFFECTS ASSOCIATED WITH THE USE OF ANABOLIC-ANDROGENIC STEROIDS (AAS)

Organ System Effect	
Cardiovascular	Dyslipidemia Atherosclerotic disease Sudden death Myocardial fibrosis, cardiomyopathy Cardiac conduction abnormalities Hypertension
Neuroendocrine	HPT axis suppression Hypogonadism after AAS withdrawal Gynecomastia
Females	Virilizing effects
Neuropsychiatric	Major mood disorders (mania, hypomania, depression) Aggression, violence AAS dependence Neuronal apoptosis Cognitive deficits
Hematologic	Polycythemia Hypercoagulability and thrombosis
Hepatic	Inflammatory and cholestatic effects Peliosis hepatitis (rare) Neoplasms (rare)
Musculoskeletal	Premature epiphyseal closure (in adolescents) Tendon rupture
Kidney	Renal failure secondary to rhabdomyolysis Focal segmental glomerulosclerosis
Dermatologic	Acne Striae

Abbreviation: HPT axis, hypothalamic-pituitary-testicular axis.

Source: Modified with permission from HG Pope Jr et al: Adverse health consequences of performance-enhancing drugs: an endocrine society scientific statement. *Endocr Rev* 35:341, 2014.

more likely to engage in other high-risk behaviors than nonusers. The adverse events associated with AAS use may be due to AAS themselves, concomitant use of other drugs, high-risk behaviors, and host characteristics that may render these individuals more susceptible to AAS use or to other high-risk behaviors.

The high rates of mortality and morbidities observed in AAS users are alarming. The risk of death among elite powerlifters has been reported to be fivefold greater than in age-matched men from the general population. The causes of death among powerlifters included suicides, myocardial infarction, hepatic coma, and non-Hodgkin's lymphoma.

Numerous reports of cardiac death among young AAS users raise concerns about the adverse cardiovascular effects of AAS. High doses of AAS may induce proatherogenic dyslipidemia, increase thrombosis risk via effects on clotting factors and platelets, induce vasospasm through their effects on vascular nitric oxide, and induce myocardial hypertrophy and fibrosis.

Replacement doses of testosterone, when administered parenterally, are associated with only a small decrease in high-density lipoprotein (HDL) cholesterol and little or no effect on total cholesterol, low-density lipoprotein (LDL) cholesterol, and triglyceride levels. In contrast, supraphysiologic doses of testosterone and orally administered, 17- α -alkylated, nonaromatizable AAS are associated with marked reductions in HDL cholesterol and increases in LDL cholesterol.

Long-term AAS use may be associated with myocardial hypertrophy and fibrosis as well as shortening of QT intervals. AAS use suppresses LH and FSH secretion and inhibits endogenous testosterone production and spermatogenesis. Consequently, stopping AAS may be associated with sexual dysfunction, fatigue, infertility, and depressive symptoms.

In some AAS users, hypothalamic-pituitary-testicular axis suppression may last more than a year, and in a few individuals, complete recovery may not occur. The symptoms of androgen deficiency during AAS withdrawal may cause some men to revert back to using AAS, leading to continued use and AAS dependence. As many as 30% of AAS users develop a syndrome of AAS dependence, characterized by long-term AAS use, despite adverse medical and psychiatric effects. Supraphysiologic doses of testosterone may also impair insulin sensitivity, predisposing to diabetes. Elevated liver enzymes, cholestatic jaundice, hepatic neoplasms, and peliosis hepatitis have been reported with oral 17- α -alkylated AAS. AAS use may cause muscle hypertrophy without compensatory adaptations in tendons, ligaments, and joints, thus increasing the risk of tendon and joint injuries. AAS use is associated with acne, baldness, and increased body hair.

Unsafe injection practices, high-risk behaviors, and increased rates of incarceration render AAS users at increased risk of HIV and hepatitis B and C. In one survey, nearly 1 in 10 gay men had injected AAS or other substances, and AAS users were more likely to report high-risk unprotected anal sex than other men.

Some AAS users develop hypomanic and manic symptoms during AAS exposure (irritability, aggressiveness, reckless behavior, and occasional psychotic symptoms, sometimes associated with violence) and major depression (sometimes associated with suicidality) during AAS withdrawal. Users may also develop other forms of illicit drug use, which may be potentiated or exacerbated by AAS.

APPROACH TO THE PATIENT: AAS Use

AAS users generally mistrust physicians and seek medical help infrequently; when they do seek medical help, it is often for the treatment of AAS withdrawal syndrome, infertility, gynecomastia, or other medical or psychiatric complications of AAS use. The suspicion of AAS use should be raised by increased hemoglobin and hematocrit levels; suppressed luteinizing hormone (LH), follicle-stimulating hormone (FSH), and testosterone levels; low HDL cholesterol; and low testicular volume and sperm density in a person who looks highly muscular (Table 7e-3). A combination of these findings and a self-report of AAS use by the patient, which usually can be elicited by a tactful interview, are often sufficient to establish a diagnosis in clinical practice.

Accredited laboratories use gas chromatography and mass spectrometry or liquid chromatography and mass spectrometry to detect AAS abuse. In recent years, the availability of high-resolution mass spectrometry and tandem mass spectrometry has further improved the sensitivity of detecting AAS abuse. Illicit testosterone use is most often detected by the urinary testosterone-to-epitestosterone ratio and further confirmed by the use of the ^{13}C : ^{12}C

TABLE 7e-3 DETECTION OF THE USE OF ANABOLIC-ANDROGENIC STEROIDS

Clinical indicators that should raise suspicion of anabolic-androgenic steroid use
Very muscular phenotype
Reduced testicular volume (<15 mL)
Laboratory indicators
Suppressed LH and FSH levels
Increased hematocrit
Detection of anabolic-androgenic steroids
LC-MS/MS analysis of urine
Detection of exogenous testosterone use
Urinary testosterone-to-epitestosterone ratio
Isotope ratio mass spectrometry analysis to detect differences in ^{13}C : ^{12}C ratio in exogenous and endogenous testosterone

Abbreviations: FSH, follicle-stimulating hormone; LC-MS/MS, liquid chromatography and tandem mass spectrometry; LH, luteinizing hormone.

ratio in testosterone by the use of isotope ratio combustion mass spectrometry. Exogenous testosterone administration increases urinary testosterone glucuronide excretion and, consequently, the testosterone-to-epitestosterone ratio. Ratios above 4 suggest exogenous testosterone use but can also reflect genetic variation. Genetic variations in the uridine diphosphoglucuronyl transferase 2B17 (*UGT2B17*), the major enzyme for testosterone glucuronidation, affect the testosterone-to-epitestosterone ratio. Synthetic testosterone has a lower $^{13}\text{C}:^{12}\text{C}$ ratio than endogenously produced testosterone, and these differences can be detected by isotope ratio combustion mass spectrometry.

TREATMENT COMPLICATIONS ASSOCIATED WITH AAS USE

The nonathlete weightlifters who abuse AAS rarely seek medical treatment and do not typically view these drugs and the associated lifestyle as deleterious to their health. In turn, many internists erroneously view AAS abuse as largely a problem of cheating in competitive sports, whereas, in fact, most AAS users are not athletes. Also, physicians often have a poor understanding of the factors motivating the use of these performance-enhancing drugs, the long-term health effects of AAS, and the associated psychopathologies that may affect treatment choices.

In addition to treating the underlying body dysmorphia disorder that motivates the use of these drugs, the treatment should be directed at the symptoms or the condition for which the patient seeks therapy, such as infertility, sexual dysfunction, gynecomastia, or depressive symptoms. Accordingly, therapy may include some combination of cognitive and behavioral therapy for the muscle dysmorphia syndrome, antidepressant therapy for depression, selective phosphodiesterase-5 inhibitors for erectile dysfunction, selective estrogen receptor modulators or aromatase inhibitors to reactivate the hypothalamic-pituitary-testicular axis, or hCG to restore testosterone levels. Clomiphene citrate, a partial estrogen receptor agonist, administered in a dose of 25–50 mg on alternate days, can increase LH and FSH levels and restore testosterone levels in a vast majority of men with AAS withdrawal syndrome. However, the recovery of sexual function during clomiphene administration is variable despite improvements in testosterone levels. Anecdotally, other aromatase inhibitors, such as anastrozole, have also been used. hCG, administered by intramuscular injections of 750–1500 IU three times each week, can raise testosterone levels into the normal range. Some patients may not respond to either clomiphene or hCG therapy, raising the possibility of irreversible long-term toxic effects of AAS on Leydig cell function.

MALE LOWER URINARY TRACT SYMPTOMS

Lower urinary tract symptoms (LUTS) in men include storage symptoms (urgency, daytime and nighttime frequency, and urgency incontinence), voiding disturbances (slow or intermittent stream, difficulty in initiating micturition, straining to void, pain or discomfort during the passage of urine, and terminal dribbling), or postmicturition symptoms (a sense of incomplete voiding after passing urine and postmicturition dribble). The overactive bladder syndrome refers to urgency with or without urgency incontinence, usually with urinary frequency and nocturia, and is often due to detrusor muscle overactivity. LUTS have historically been attributed to benign prostatic hyperplasia, although it has become apparent that the pathophysiologic mechanisms of LUTS are complex and multifactorial and may include structural or functional abnormalities of the bladder, bladder neck, prostate, distal sphincter mechanism, and urethra, as well as abnormalities in the neural control to the lower urinary tract. A presumptive diagnosis of benign prostatic hyperplasia should be made only in men with LUTS who have demonstrable evidence of prostate enlargement and obstruction based on the size of the prostate. Diuretics, antihistamines, antidepressants, and other medications that have anticholinergic properties

can cause or exacerbate LUTS in older men. The intensity of LUTS symptoms tends to fluctuate over time.

LUTS is highly prevalent in older men, affecting nearly 50% of men over the age of 65 and 70% of men over the age of 80. LUTS adversely affects quality of life because of its impact on sleep, ability to perform activities of daily living, and depressive symptoms. LUTS is often associated with erectile dysfunction.

APPROACH TO THE PATIENT: LUTS

Medical evaluation should include assessment of the symptom severity using the International Prostate Symptom Score and, in some patients, a frequency-volume chart. The impact of LUTS on sleep and activities of daily living and quality of life should be evaluated. Evaluation should also include verification of medications that may contribute to LUTS, digital prostate examination, neurologic examination focused on perineum and lower extremities, urinalysis, fasting blood glucose, electrolytes, creatinine, and prostate-specific antigen (PSA). Urodynamic studies are not required in most patients but are recommended when invasive surgical therapies are being considered.

TREATMENT LUTS

Men who have mild symptoms can be reassured and followed. Men with mild to moderate LUTS can be treated effectively using α -adrenergic antagonists, phosphodiesterase-5 (PDE5) inhibitors, steroid 5 α -reductase inhibitors, or anticholinergic agents alone or in combination. Selective α -adrenergic antagonists are typically the first line of therapy. In men with probable benign prostate obstruction with gland enlargement and LUTS, therapy using a steroid 5 α -reductase inhibitor, such as finasteride or dutasteride, for 1 or more years improves urinary symptoms and flow rate and reduces prostatic volume. Long-term treatment with 5 α -reductase inhibitors can reduce progression to acute urinary retention and need for prostate surgery. Combined administration of a steroid 5 α -reductase inhibitor and α_1 -adrenergic blocker can rapidly improve urinary symptoms and reduce the relative risk of acute urinary retention and surgery. PDE5 inhibitors, when administered chronically alone or in combination with α -adrenergic blockers, are effective in improving LUTS and erectile dysfunction through their effects on nitric oxide-cyclic guanosine monophosphate (cGMP) in the bladder, urethra, and prostate. PDE5 inhibitors do not improve urinary flow parameters. Anticholinergic drugs are used for the treatment of overactive bladder in men with prominent urgency symptoms and no evidence of elevated postvoid residual urine. Surgery is indicated when medical therapy fails or if symptoms progress despite medical therapy.

MEDICAL COMPLICATIONS OF PROSTATE CANCER THERAPY

Prostate cancer is the most common malignancy in American men, accounting for 29% of all diagnosed cancers and approximately 13% of all cancer deaths; its incidence is on the rise, partly due to increased screening with PSA. In 2013, approximately 233,000 new cases of prostate cancer were diagnosed in the United States and there were 29,480 deaths related to prostate cancer. The majority of these men have low-grade, organ-confined prostate cancer and excellent prospects of long-term survival. Substantial improvement in survival in men with prostate cancer has focused attention on the high prevalence of sexual dysfunction, physical dysfunction, and low vitality, which are important contributors to poor quality of life among patients treated for prostate cancer. The pathophysiology of these symptoms after radical prostatectomy is multifactorial, but denervation and androgen deficiency are important contributors to these symptoms.

Androgen deficiency is common in men with prostate cancer. Testosterone levels decline with age, and men with prostate cancer are at risk of having low testosterone levels simply by virtue of their age.

However, total and free testosterone levels are even lower in men with prostate cancer, who have undergone prostatectomy, when compared with age-matched controls without cancer. Androgen deficiency in men with prostate cancer is associated with distressing symptoms such as fatigue, sexual dysfunction, hot flashes, mobility limitation, and decreased physical function. Even with a bilateral nerve-sparing procedure, more than 50% of men develop sexual dysfunction after surgery. Although there is some recovery of sexual function with passage of time, 40–50% of men undergoing radical prostatectomy find their sexual performance to be problematic 18 months after surgery. Sexual performance problems are a source of psychosocial distress in men with localized prostate cancer. In addition to its causal contribution to distressing symptoms, androgen deficiency in men with prostate cancer increases the risk of bone fractures, diabetes, coronary heart disease, and frailty.

Testosterone Therapy in Men with History of Prostate Cancer A history of prostate cancer has historically been considered a contraindication for testosterone therapy. This guidance is based on observations that testosterone promotes the growth of metastatic prostate cancer. Metastatic prostate cancer generally regresses after orchidectomy and androgen deprivation therapy. Androgen receptor signaling plays a central role in maintaining growth of normal prostate and prostate cancer. PSA levels are lower in hypogonadal men and increase after testosterone therapy. Prostate volume is lower in hypogonadal men and increases after testosterone therapy to levels seen in age-matched controls.

However, the role of testosterone in prostate cancer is complex. Epidemiologic studies have not revealed a consistent relationship between serum testosterone and prostate cancer. In a landmark randomized trial, testosterone therapy of older men with low testosterone did not affect intraprostatic androgen levels or the expression of androgen-dependent prostatic genes. The suppression of circulating testosterone levels by a gonadotropin-releasing hormone (GnRH) antagonist also does not affect intraprostatic androgen concentrations. Open-label trials and retrospective analyses of testosterone therapy in

men with prostate cancer, who have undergone radical prostatectomy and have undetectable PSA levels after radical prostatectomy, have found very low rates of PSA recurrence. Even in men with high-grade prostatic intraepithelial neoplasia (HGPIN)—a group at high risk of developing prostate cancer—testosterone therapy for 1 year did not increase PSA or rates of prostate cancer.

After radical prostatectomy, in the absence of residual cancer, PSA becomes undetectable within a month. An undetectable PSA after radical prostatectomy is a good indicator of biochemical recurrence-free survival at 5 years. Therefore, men with organ-confined prostate cancer (pT2), Gleason score ≤ 6 , and a preoperative PSA of <10 ng/mL, who have had undetectable PSA levels (<0.1 ng/mL) for >2 years after radical prostatectomy, have very low risk of disease recurrence ($<0.5\%$ at 10 years) and may be considered for testosterone therapy on an individualized basis. If testosterone therapy is instituted, it should be associated with careful monitoring of PSA levels and done in consultation with a urologist.

MEDICAL COMPLICATIONS OF ANDROGEN DEPRIVATION THERAPY

In patients with prostate cancer and distant metastases, androgen deprivation therapy (ADT) improves survival. In patients with locally advanced disease, ADT in combination with external-beam radiation or as an adjuvant therapy (after prostatectomy and pelvic lymphadenectomy) also has been shown to improve survival. However, ADT is being increasingly used as primary therapy in men with localized disease and in men encountering biochemical recurrence without clear evidence of survival advantage. Because most men with prostate cancer die of conditions other than their primary malignancy, recognition and management of these adverse effects is paramount.

Profound hypogonadism resulting from ADT is associated with sexual dysfunction, vasomotor symptoms, gynecomastia, decreased muscle mass and strength, frailty, increased fat mass, anemia, fatigue, bone loss, loss of body hair, depressive symptoms, and reduced quality of life. Diabetes and cardiovascular disease have recently been added to the list of these complications (Fig. 7e-3). Treatment with GnRH

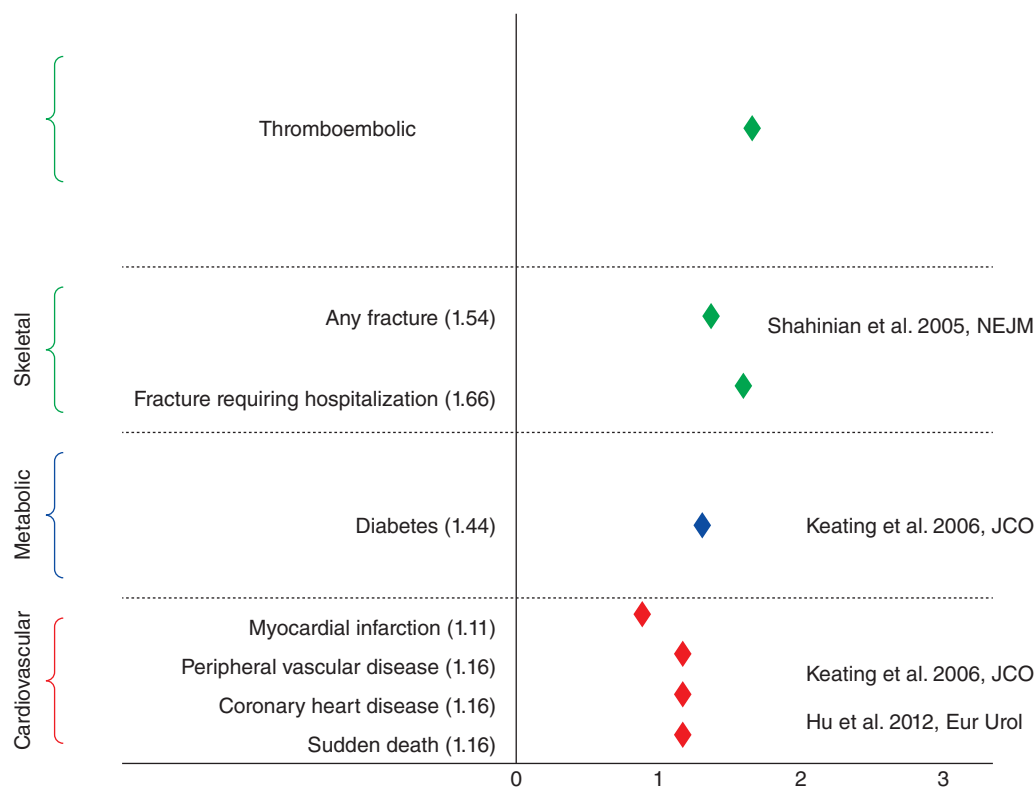


FIGURE 7e-3 Adverse cardiometabolic and skeletal effects of androgen deprivation therapy (ADT) in men receiving ADT for prostate cancer. Administration of ADT has been associated with increased risk of thromboembolic events, fractures, and diabetes. Some, but not all, studies have reported increased risk of cardiovascular events in men receiving ADT. (Data on relative risk were derived from VB Shahinian et al: *N Engl J Med* 352:154, 2005; NL Keating et al: *J Clin Oncol* 24:4448, 2006; and JC Hu et al: *Eur Urol* 61:1119, 2012.)

TABLE 7e-4 CHECKLIST FOR MEN UNDERGOING ANDROGEN DEPRIVATION THERAPY (ADT)

1. Weigh the risks and benefits of ADT and whether intermittent ADT is a feasible and safe option.
2. Perform a baseline assessment including fasting glucose, plasma lipids, blood pressure, bone mineral density, and FRAX[®] score.
3. Optimize calcium and vitamin D intake, encourage structured physical activity and exercise, and consider pharmacologic therapy in men with a previous minimal trauma fracture and those with a 10-year risk of a major osteoporotic fracture >20%, unless contraindicated.
4. Monitor body weight, fasting glucose, plasma lipids, blood pressure, and bone mineral density, and encourage smoking cessation and physical activity.
5. In men who are receiving ADT and who experience bothersome hot flashes, as indicated by sleep disturbance or interference with work or activities of daily living, consider initial therapy with venlafaxine. If in effective, add medroxyprogesterone acetate.
6. In men who experience painful breast enlargement, consider therapy with an estrogen receptor antagonist, such as tamoxifen.

agonists in men with prostate cancer is associated with rapid induction of insulin resistance, hyperinsulinemia, and a significant increase in the risk of incident diabetes. Metabolic syndrome is prevalent in over 50% of men undergoing long-term ADT. Some but not all studies have reported an increased risk of cardiovascular events, death due to cardiovascular events, and peripheral vascular disease in men undergoing ADT. Men receiving ADT are also at increased risk of thromboembolic events. The rates of acute kidney injury are higher in men currently receiving ADT than in men not receiving ADT; the increased risk appears to be particularly associated with the use of combined regimens of a GnRH agonist plus an antiandrogen. ADT also is associated with substantially increased risk of osteoporosis and bone fractures.

APPROACH TO THE PATIENT: Men Receiving ADT

The benefits of ADT in treating nonmetastatic prostate cancer should be carefully weighed against the risks of ADT-induced adverse events (Table 7e-4). If ADT is medically indicated, consider whether intermittent ADT is a feasible option. Men being considered for ADT should undergo assessment of cardiovascular, diabetes, and fracture risk; this assessment may include measurement of blood glucose, plasma lipids, and bone mineral density (BMD) by dual-energy x-ray absorptiometry. Institute measures to prevent bone loss, including physical activity, adequate calcium and vitamin D intake, and pharmacologic therapy in men with a previous minimal trauma fracture and those with a 10-year risk of a major osteoporotic fracture >20%, unless contraindicated. Men with prostate cancer who are receiving ADT should be monitored for weight gain and diabetes. Encourage lifestyle interventions, including physical activity and exercise, and attention to weight, blood pressure, lipid profile, blood glucose, and smoking cessation, to reduce the risk of cardiometabolic complications. In randomized trials, medroxyprogesterone, cyproterone acetate, and the selective serotonin reuptake inhibitor venlafaxine have been shown to be more efficacious than placebo in alleviating hot flashes. The side effects of these medications, including increased appetite and weight gain with medroxyprogesterone, gynecomastia with estrogenic compounds, and dry mouth with venlafaxine, should be weighed against their relative efficacy. Acupuncture, soy products, vitamin E, and herbal medicines have been used empirically for the treatment of vasomotor symptoms without clear evidence of efficacy. Gynecomastia can be prevented by local radiation therapy or the use of an antiestrogen or an aromatase inhibitor; these therapies are effective in alleviating pain and tenderness but are less effective in reducing established gynecomastia.

8 Medical Disorders During Pregnancy

Robert L. Barbieri, John T. Repke

Each year, approximately 4 million births occur in the United States, and more than 130 million births occur worldwide. A significant proportion of births are complicated by medical disorders. In the past, many medical disorders were contraindications to pregnancy. Advances in obstetrics, neonatology, obstetric anesthesiology, and medicine have increased the expectation that pregnancy will result in a positive outcome for both mother and fetus despite most of these conditions. A successful pregnancy requires important physiologic adaptations, such as a marked increase in cardiac output. Medical problems that interfere with the physiologic adaptations of pregnancy increase the risk for poor pregnancy outcome; conversely, in some instances, pregnancy may adversely impact an underlying medical disorder.

HYPERTENSION

(See also Chap. 298) In pregnancy, cardiac output increases by 40%, with most of the increase due to an increase in stroke volume. Heart rate increases by ~10 beats/min during the third trimester. In the second trimester, systemic vascular resistance decreases, and this decline is associated with a fall in blood pressure. During pregnancy, a blood pressure of 140/90 mmHg is considered to be abnormally elevated and is associated with an increase in perinatal morbidity and mortality. In all pregnant women, the measurement of blood pressure should be performed in the sitting position, because the lateral recumbent position may result in a blood pressure lower than that recorded in the sitting position. The diagnosis of hypertension requires the measurement of two elevated blood pressures at least 6 h apart. Hypertension during pregnancy is usually caused by preeclampsia, chronic hypertension, gestational hypertension, or renal disease.

PREECLAMPSIA

Approximately 5–7% of all pregnant women develop *preeclampsia*, the new onset of hypertension (blood pressure >140/90 mmHg) and proteinuria (either a 24 hour urinary protein >300 mg/24 h, or a protein-creatinine ratio ≥ 0.3) after 20 weeks of gestation. Although the precise pathophysiology of preeclampsia remains unknown, recent studies show excessive placental production of antagonists to both vascular epithelial growth factor (VEGF) and transforming growth factor β (TGF- β). These antagonists to VEGF and TGF- β disrupt endothelial and renal glomerular function resulting in edema, hypertension, and proteinuria. The renal histological feature of preeclampsia is glomerular endotheliosis. Glomerular endothelial cells are swollen and encroach on the vascular lumen. Preeclampsia is associated with abnormalities of cerebral circulatory autoregulation, which increase the risk of stroke at mildly and moderately elevated blood pressures. Risk factors for the development of preeclampsia include nulliparity, diabetes mellitus, a history of renal disease or chronic hypertension, a prior history of preeclampsia, extremes of maternal age (>35 years or <15 years), obesity, antiphospholipid antibody syndrome, and multiple gestation. Low-dose aspirin (81 mg daily, initiated at the end of the first trimester) may reduce the risk of preeclampsia in pregnant women at high risk of developing the disease.

In December, 2013 The American College of Obstetricians and Gynecologists issued a report summarizing the findings and recommendations of their Task Force on Hypertension in Pregnancy. With respect to preeclampsia several pertinent revisions to the diagnostic criteria were made including: proteinuria is no longer an absolute requirement for making the diagnosis; the terms mild and severe preeclampsia have been replaced, and the disease is now termed preeclampsia either with or without severe features; removal of fetal growth restriction as a defining criterion for severe preeclampsia.

Preeclampsia with severe features is the presence of new-onset hypertension and proteinuria accompanied by end-organ damage. Features may include severe elevation of blood pressure (>160/110 mmHg), evidence of central nervous system (CNS) dysfunction (headaches, blurred vision, seizures, coma), renal dysfunction (oliguria or creatinine >1.5 mg/dL), pulmonary edema, hepatocellular injury (serum alanine aminotransferase level more than twofold the upper limit of normal), hematologic dysfunction (platelet count <100,000/L or disseminated intravascular coagulation [DIC]). The *HELLP syndrome* (hemolysis, elevated liver enzymes, low platelets) is a special subtype of severe preeclampsia and is a major cause of morbidity and mortality in this disease. Platelet dysfunction and coagulation disorders further increase the risk of stroke.

TREATMENT PREECLAMPSIA

Preeclampsia resolves within a few weeks after delivery. For pregnant women with preeclampsia prior to 37 weeks of gestation, delivery reduces the mother's morbidity but exposes the fetus to the risk of premature birth. The management of preeclampsia is challenging because it requires the clinician to balance the health of the mother and fetus simultaneously. In general, prior to term, women with *mild* preeclampsia without severe features may be managed conservatively with limited physical activity, although bed rest is not recommended, close monitoring of blood pressure and renal function, and careful fetal surveillance. For women with preeclampsia with severe features, delivery is recommended unless the patient is eligible for expectant management in a tertiary hospital setting. Expectant management of preeclampsia with severe features remote from term affords some benefits for the fetus but significant risks for the mother.

The definitive treatment of preeclampsia is delivery of the fetus and placenta. For women with preeclampsia with severe features, aggressive management of blood pressures >160/110 mmHg reduces the risk of cerebrovascular accidents. IV labetalol or hydralazine is most commonly used to acutely manage severe hypertension in preeclampsia; labetalol is associated with fewer episodes of maternal hypotension. Oral nifedipine and labetalol are commonly used to manage hypertension in pregnancy. Elevated arterial pressure should be reduced slowly to avoid hypotension and a decrease in blood flow to the fetus. Angiotensin-converting enzyme (ACE) inhibitors as well as angiotensin-receptor blockers should be avoided in the second and third trimesters of pregnancy because of their adverse effects on fetal development.

Magnesium sulfate is the preferred agent for the prevention and treatment of eclamptic seizures. Large, randomized clinical trials have demonstrated the superiority of magnesium sulfate over phenytoin and diazepam in reducing the risk of seizure and, possibly, the risk of maternal death. Magnesium may prevent seizures by interacting with *N*-methyl-D-aspartate (NMDA) receptors in the CNS. Given the difficulty of predicting eclamptic seizures on the basis of disease severity, once the decision to proceed with delivery is made, most patients carrying a diagnosis of preeclampsia should be treated with magnesium sulfate. Women who have had preeclampsia appear to be at increased risk of cardiovascular and renal disease later in life.

CHRONIC ESSENTIAL HYPERTENSION

Pregnancy complicated by chronic essential hypertension is associated with intrauterine growth restriction and increased perinatal mortality.

Pregnant women with chronic hypertension are at increased risk for superimposed preeclampsia and abruptio placentae. Women with chronic hypertension should have a thorough prepregnancy evaluation, both to identify remediable causes of hypertension and to ensure that the prescribed antihypertensive agents (e.g., ACE inhibitors, angiotensin-receptor blockers) are not associated with an adverse outcome of pregnancy. α -Methyldopa, labetalol, and nifedipine are the most commonly used medications for the treatment of chronic hypertension in pregnancy. The target blood pressure is in the range of 130–150 mmHg systolic and 80–100 mmHg diastolic. Should hypertension worsen during pregnancy, baseline evaluation of renal function (see below) is necessary to help differentiate the effects of chronic hypertension from those of superimposed preeclampsia. There are no convincing data that the treatment of mild chronic hypertension improves perinatal outcome.

GESTATIONAL HYPERTENSION

The development of elevated blood pressure during pregnancy or in the first 24 h post-partum in the absence of preexisting chronic hypertension or proteinuria is referred to as *gestational hypertension*. Mild gestational hypertension that does not progress to preeclampsia has not been associated with adverse pregnancy outcome or adverse long-term prognosis.

RENAL DISEASE

(See also Chaps. 333 and 341) Normal pregnancy is characterized by an increase in glomerular filtration rate and creatinine clearance. This increase occurs secondary to a rise in renal plasma flow and increased glomerular filtration pressures. Patients with underlying renal disease and hypertension may expect a worsening of hypertension during pregnancy. If superimposed preeclampsia develops, the additional endothelial injury results in a capillary leak syndrome that may make management challenging. In general, patients with underlying renal disease and hypertension benefit from aggressive management of blood pressure. Preconception counseling is also essential for these patients so that accurate risk assessment and medication changes can occur prior to pregnancy. In general, a prepregnancy serum creatinine level <133 μ mol/L (<1.5 mg/dL) is associated with a favorable prognosis. When renal disease worsens during pregnancy, close collaboration between the internist and the maternal-fetal medicine specialist is essential so that decisions regarding delivery can be weighed to balance the sequelae of prematurity for the neonate versus long-term sequelae for the mother with respect to future renal function.

CARDIAC DISEASE

VALVULAR HEART DISEASE

(See also Chaps. 283–286) Valvular heart disease is the most common cardiac problem complicating pregnancy.

Mitral Stenosis This is the valvular disease most likely to cause death during pregnancy. The pregnancy-induced increase in blood volume, cardiac output, and tachycardia can increase the transmitral pressure gradient and cause pulmonary edema in women with mitral stenosis. Women with moderate to severe mitral stenosis who are planning pregnancy and have either symptomatic disease or pulmonary hypertension should undergo valvuloplasty prior to conception. Pregnancy associated with long-standing mitral stenosis may result in pulmonary hypertension. Sudden death has been reported when hypovolemia occurs. Careful control of heart rate, especially during labor and delivery, minimizes the impact of tachycardia and reduced ventricular filling times on cardiac function. Pregnant women with mitral stenosis are at increased risk for the development of atrial fibrillation and other tachyarrhythmias. Medical management of severe mitral stenosis and atrial fibrillation with digoxin and beta blockers is recommended. Balloon valvulotomy can be carried out during pregnancy. The immediate postpartum period is a time of particular concern secondary to rapid volume shifts. Careful monitoring of cardiac and fluid status should be observed.

Mitral Regurgitation and Aortic Regurgitation and Stenosis The pregnancy-induced decrease in systemic vascular resistance reduces

the risk of cardiac failure with these conditions. As a rule, mitral valve prolapse does not present problems for the pregnant patient, and aortic stenosis, unless very severe, is well tolerated. In the most severe cases of aortic stenosis, limitation of activity or balloon valvuloplasty may be indicated.

CONGENITAL HEART DISEASE

(See also Chap. 282) Reparative surgery has markedly increased the number of women with surgically repaired congenital heart disease. Maternal morbidity and mortality are greater among these women than among those without surgical repairs. When pregnant, these patients should be jointly managed by a cardiologist and an obstetrician familiar with these problems. The presence of a congenital cardiac lesion in the mother increases the risk of congenital cardiac disease in the newborn. Prenatal screening of the fetus for congenital cardiac disease with ultrasound is recommended. Atrial or ventricular septal defect is usually well tolerated during pregnancy in the absence of pulmonary hypertension, provided that the woman's prepregnancy cardiac status is favorable. Use of air filters on IV sets during labor and delivery in patients with intracardiac shunts is recommended.

OTHER CARDIAC DISORDERS

Supraventricular tachycardia (Chap. 276) is a common cardiac complication of pregnancy. Treatment is the same as in the nonpregnant patient, and fetal tolerance of medications such as adenosine and calcium channel blockers is acceptable. When necessary, pharmacologic or electric cardioversion may be performed to improve cardiac performance and reduce symptoms. This intervention is generally well tolerated by mother and fetus.

Peripartum cardiomyopathy (Chap. 287) is an uncommon disorder of pregnancy associated with myocarditis, and its etiology remains unknown. Treatment is directed toward symptomatic relief and improvement of cardiac function. Many patients recover completely; others are left with progressive dilated cardiomyopathy. Recurrence in a subsequent pregnancy has been reported, and women who do not have normal baseline left-ventricular function after an episode of peripartum cardiomyopathy should be counseled to avoid pregnancy.

SPECIFIC HIGH-RISK CARDIAC LESIONS

Marfan Syndrome (See also Chap. 427) This autosomal dominant disease is associated with a high risk of maternal morbidity. Approximately 15% of pregnant women with Marfan syndrome develop a major cardiovascular manifestation during pregnancy, with almost all women surviving. An aortic root diameter <40 mm is associated with a favorable outcome of pregnancy. Prophylactic therapy with beta blockers has been advocated, although large-scale clinical trials in pregnancy have not been performed. *Ehlers-Danlos syndrome* (EDS) may be associated with premature labor, and in type IV EDS there is increased risk of organ or vascular rupture that may cause death.

Pulmonary Hypertension (See also Chap. 304) Maternal mortality in the setting of severe pulmonary hypertension is high, and primary pulmonary hypertension is a contraindication to pregnancy. Termination of pregnancy may be advisable in these circumstances to preserve the life of the mother. In the *Eisenmenger syndrome*, i.e., the combination of pulmonary hypertension with right-to-left shunting due to congenital abnormalities (Chap. 282), maternal and fetal deaths occur frequently. Systemic hypotension may occur after blood loss, prolonged Valsalva maneuver, or regional anesthesia; sudden death secondary to hypotension is a dreaded complication. Management of these patients is challenging, and invasive hemodynamic monitoring during labor and delivery is recommended in severe cases.

In patients with pulmonary hypertension, vaginal delivery is less stressful hemodynamically than cesarean section, which should be reserved for accepted obstetric indications.

DEEP VENOUS THROMBOSIS AND PULMONARY EMBOLISM

(See also Chap. 300) A hypercoagulable state is characteristic of pregnancy, and deep venous thrombosis (DVT) occurs in about 1 in

500 pregnancies. In pregnant women, most unilateral DVTs occur in the left leg because the left iliac vein is compressed by the right iliac artery and the uterus compresses the inferior vena cava. Pregnancy is associated with an increase in procoagulants such as factors V and VII and a decrease in anticoagulant activity, including proteins C and S. Pulmonary embolism is one of the most common causes of maternal death in the United States. Activated protein C resistance caused by the factor V Leiden mutation increases the risk for DVT and pulmonary embolism during pregnancy. Approximately 25% of women with DVT during pregnancy carry the factor V Leiden allele. Additional genetic mutations associated with DVT during pregnancy include the prothrombin G20210A mutation (heterozygotes and homozygotes) and the methylenetetrahydrofolate reductase C677T mutation (homozygotes).

TREATMENT DEEP VENOUS THROMBOSIS

Aggressive diagnosis and management of DVT and suspected pulmonary embolism optimize the outcome for mother and fetus. In general, all diagnostic and therapeutic modalities afforded the nonpregnant patient should be utilized in pregnancy except for D-dimer measurement, in which values are elevated in normal pregnancy. Anticoagulant therapy with low-molecular-weight heparin (LMWH) or unfractionated heparin is indicated in pregnant women with DVT. LMWH may be associated with an increased risk of epidural hematoma in women receiving an epidural anesthetic in labor. Four weeks prior to anticipated delivery, LMWH should be switched to unfractionated heparin. Warfarin therapy is contraindicated in the first trimester due to its association with fetal chondrodysplasia punctata. In the second and third trimesters, warfarin may cause fetal optic atrophy and mental retardation. When DVT occurs in the postpartum period, LMWH therapy for 7–10 days may be followed by warfarin therapy for 3–6 months. Warfarin is not contraindicated in breast-feeding women. For women at moderate or high risk of DVT who have a cesarean delivery, mechanical and/or pharmacologic prophylaxis is warranted.

ENDOCRINE DISORDERS

DIABETES MELLITUS

(See also Chaps. 417–419) In pregnancy, the fetoplacental unit induces major metabolic changes, the purpose of which is to shunt glucose and amino acids to the fetus while the mother uses ketones and triglycerides to fuel her metabolic needs. These metabolic changes are accompanied by maternal insulin resistance caused in part by placental production of steroids, a growth hormone variant, and placental lactogen. Although pregnancy has been referred to as a state of “accelerated starvation,” it is better characterized as “accelerated ketosis.” In pregnancy, after an overnight fast, plasma glucose is lower by 0.8–1.1 mmol/L (15–20 mg/dL) than in the nonpregnant state. This difference is due to the use of glucose by the fetus. In early pregnancy, fasting may result in circulating glucose concentrations in the range of 2.2 mmol/L (40 mg/dL) and may be associated with symptoms of hypoglycemia. In contrast to the decrease in maternal glucose concentration, plasma hydroxybutyrate and acetoacetate levels rise to two to four times normal after a fast.

TREATMENT DIABETES MELLITUS IN PREGNANCY

Pregnancy complicated by diabetes mellitus is associated with higher maternal and perinatal morbidity and mortality rates. Preconception counseling and treatment are important for the diabetic patient contemplating pregnancy and can reduce the risk of congenital malformations and improve pregnancy outcome. Folate supplementation reduces the incidence of fetal neural tube defects, which occur with greater frequency in fetuses of diabetic mothers. In addition, optimizing glucose control during

key periods of organogenesis reduces other congenital anomalies, including sacral agenesis, caudal dysplasia, renal agenesis, and ventricular septal defect.

Once pregnancy is established, glucose control should be managed more aggressively than in the nonpregnant state. In addition to dietary changes, this enhanced management requires more frequent blood glucose monitoring and often involves additional injections of insulin or conversion to an insulin pump. Fasting blood glucose levels should be maintained at <5.8 mmol/L (<105 mg/dL), with avoidance of values >7.8 mmol/L (140 mg/dL). Commencing in the third trimester, regular surveillance of maternal glucose control as well as assessment of fetal growth (obstetric sonography) and fetoplacental oxygenation (fetal heart rate monitoring or biophysical profile) optimize pregnancy outcome. Pregnant diabetic patients without vascular disease are at greater risk for delivering a macrosomic fetus, and attention to fetal growth via clinical and ultrasound examination is important. Fetal macrosomia is associated with an increased risk of maternal and fetal birth trauma, including permanent newborn *Erb's palsy*. Pregnant women with diabetes have an increased risk of developing preeclampsia, and those with vascular disease are at greater risk for developing intrauterine growth restriction, which is associated with an increased risk of fetal and neonatal death. Excellent pregnancy outcomes in patients with diabetic nephropathy and proliferative retinopathy have been reported with aggressive glucose control and intensive maternal and fetal surveillance.

As pregnancy progresses, glycemic control may become more difficult to achieve due to an increase in insulin resistance. Because of delayed pulmonary maturation of the fetuses of diabetic mothers, early delivery should be avoided unless there is biochemical evidence of fetal lung maturity. In general, efforts to control glucose and avoid preterm delivery result in the best overall outcome for both mother and newborn. Preterm delivery is generally performed only for the usual obstetric indications (e.g., preeclampsia, fetal growth restriction, non-reassuring fetal testing) or for worsening maternal renal or active proliferative retinopathy.

GESTATIONAL DIABETES

Gestational diabetes occurs in approximately 4% of pregnancies. All pregnant women should be screened for gestational diabetes unless they are in a low-risk group. Women at low risk for gestational diabetes are those <25 years of age; those with a body mass index <25 kg/m², no maternal history of macrosomia or gestational diabetes, and no diabetes in a first-degree relative; and those who are not members of a high-risk ethnic group (African American, Hispanic, Native American). A typical two-step strategy for establishing the diagnosis of gestational diabetes involves administration of a 50-g oral glucose challenge with a single serum glucose measurement at 60 min. If the plasma glucose is <7.8 mmol/L (<130 mg/dL), the test is considered normal. Plasma glucose >7.8 mmol/L (>130 mg/dL) warrants administration of a 100-g oral glucose challenge with plasma glucose measurements obtained in the fasting state and at 1, 2, and 3 h. Normal plasma glucose concentrations at these time points are <5.8 mmol/L (<105 mg/dL), 10.5 mmol/L (190 mg/dL), 9.1 mmol/L (165 mg/dL), and 8.0 mmol/L (145 mg/dL), respectively. Some centers have adopted more sensitive criteria, using values of <5.3 mmol/L (<95 mg/dL), <10 mmol/L (<180 mg/dL), <8.6 mmol/L (<155 mg/dL), and <7.8 mmol/L (<140 mg/dL) as the upper norms for a 3-h glucose tolerance test. Two elevated glucose values indicate a positive test. Adverse pregnancy outcomes for mother and fetus appear to increase with glucose as a continuous variable; thus it is challenging to define the optimal threshold for establishing the diagnosis of gestational diabetes.

Pregnant women with gestational diabetes are at increased risk of stillbirth, preeclampsia, and delivery of infants who are large for their gestational age, with resulting birth lacerations, shoulder dystocia, and birth trauma including brachial plexus injury. These fetuses are at risk of hypoglycemia, hyperbilirubinemia, and polycythemia. Tight control of blood sugar during pregnancy and labor can reduce these risks.

TREATMENT GESTATIONAL DIABETES

Treatment of gestational diabetes with a two-step strategy—dietary intervention followed by insulin injections if diet alone does not adequately control blood sugar [fasting glucose <5.6 mmol/L (<100 mg/dL) and 2-h postprandial glucose <7.0 mmol/L (<126 mg/dL)]—is associated with a decreased risk of birth trauma for the fetus. Oral hypoglycemic agents such as glyburide and metformin have become more commonly utilized for managing gestational diabetes refractory to nutritional management, but many experts favor insulin therapy. For women with gestational diabetes, there is a 40% risk of being diagnosed with diabetes within the 10 years after the index pregnancy. In women with a history of gestational diabetes, exercise, weight loss, and treatment with metformin reduce the risk of developing diabetes. All women with a history of gestational diabetes should be counseled about prevention strategies and evaluated regularly for diabetes.

OBESITY

(See also Chap. 416) Pregnant women who are obese have an increased risk of stillbirth, congenital fetal malformations, gestational diabetes, preeclampsia, urinary tract infections, post-date delivery, and cesarean delivery. Women contemplating pregnancy should attempt to attain a healthy weight prior to conception. For morbidly obese women who have not been able to lose weight with lifestyle changes, bariatric surgery may result in weight loss and improve pregnancy outcomes. Following bariatric surgery, women should delay conception for 1 year to avoid pregnancy during an interval of rapid metabolic changes.

THYROID DISEASE

(See also Chap. 405) In pregnancy, the estrogen-induced increase in thyroxine-binding globulin increases circulating levels of total T₃ and total T₄. The normal range of circulating levels of free T₄, free T₃, and thyroid-stimulating hormone (TSH) remain unaltered by pregnancy.

The thyroid gland normally enlarges during pregnancy. Many physiologic adaptations to pregnancy may mimic subtle signs of *hyperthyroidism*. Maternal hyperthyroidism occurs at a rate of ~ 2 per 1000 pregnancies and is generally well tolerated by pregnant women. Clinical signs and symptoms should alert the physician to the occurrence of this condition. Hyperthyroidism in pregnancy is most commonly caused by Graves' disease, but autonomously functioning nodules and gestational trophoblastic disease should also be considered. Although pregnant women are able to tolerate mild hyperthyroidism without adverse sequelae, more severe hyperthyroidism can cause spontaneous abortion or premature labor, and thyroid storm is associated with a significant risk of maternal death.

Testing for *hypothyroidism* using TSH measurements before or early in pregnancy may be warranted in symptomatic women and in women with a personal or family history of thyroid disease. With use of this case-finding approach, about 30% of pregnant women with mild hypothyroidism remain undiagnosed, leading some to recommend universal screening. Children born to women with an elevated serum TSH (and a normal total thyroxine) during pregnancy may have impaired performance on neuropsychologic tests.

TREATMENT HYPERTHYROIDISM IN PREGNANCY

HYPERTHYROIDISM

Methimazole crosses the placenta to a greater degree than propylthiouracil and has been associated with fetal aplasia cutis. However, propylthiouracil can be associated with liver failure. Some experts recommend propylthiouracil in the first trimester and methimazole thereafter. Radioiodine should not be used during pregnancy, either for scanning or for treatment, because of effects on the fetal thyroid. In emergent circumstances, additional treatment with beta blockers may be necessary. Hyperthyroidism is most difficult to control in the first trimester of pregnancy and easiest to control in the third trimester.

HYPOTHYROIDISM

The goal of therapy for hypothyroidism is to maintain the serum TSH in the normal range, and thyroxine is the drug of choice. During pregnancy, the dose of thyroxine required to keep the TSH in the normal range rises. In one study, the mean replacement dose of thyroxine required to maintain the TSH in the normal range was 0.1 mg daily before pregnancy and increased to 0.15 mg daily during pregnancy. Since the increased thyroxine requirement occurs as early as the fifth week of pregnancy, one approach is to increase the thyroxine dose by 30% (two additional pills weekly) as soon as pregnancy is diagnosed and then adjust the dose by serial measurements of TSH.

HEMATOLOGIC DISORDERS

Pregnancy has been described as a state of physiologic anemia. Part of the reduction in hemoglobin concentration is dilutional, but iron and folate deficiencies are major causes of correctable anemia during pregnancy.

In populations at high risk for hemoglobinopathies (Chap. 127), hemoglobin electrophoresis should be performed as part of the prenatal screen. Hemoglobinopathies can be associated with increased maternal and fetal morbidity and mortality. Management is tailored to the specific hemoglobinopathy and is generally the same for both pregnant and nonpregnant women. Prenatal diagnosis of hemoglobinopathies in the fetus is readily available and should be discussed with prospective parents either prior to or early in pregnancy.

Thrombocytopenia occurs commonly during pregnancy. The majority of cases are benign gestational thrombocytopenias, but the differential diagnosis should include immune thrombocytopenia (Chap. 140), thrombotic thrombocytopenic purpura, and preeclampsia. Maternal thrombocytopenia may also be caused by DIC, which is a consumptive coagulopathy characterized by thrombocytopenia, prolonged prothrombin time (PT) and activated partial thromboplastin time (aPTT), elevated fibrin degradation products, and a low fibrinogen concentration. Several catastrophic obstetric events are associated with the development of DIC, including retention of a dead fetus, sepsis, abruptio placentae, and amniotic fluid embolism.

NEUROLOGIC DISORDERS

Headache appearing during pregnancy is usually due to migraine (Chap. 21), a condition that may worsen, improve, or be unaffected by pregnancy. A new or worsening headache, particularly if associated with visual blurring, may signal eclampsia (above) or pseudotumor cerebri (benign intracranial hypertension); diplopia due to a sixth-nerve palsy suggests pseudotumor cerebri (Chap. 39). The risk of seizures in patients with epilepsy increases in the postpartum period but not consistently during pregnancy; management is discussed in Chap. 445. The risk of stroke is generally thought to increase during pregnancy because of a hypercoagulable state; however, studies suggest that the period of risk occurs primarily in the postpartum period and that both ischemic and hemorrhagic strokes may occur at this time. Guidelines for use of heparin therapy are summarized above (see “Deep Venous Thrombosis and Pulmonary Embolism”); warfarin is teratogenic and should be avoided.

The onset of a new movement disorder during pregnancy suggests *chorea gravidarum*, a variant of Sydenham’s chorea associated with rheumatic fever and streptococcal infection (Chap. 381); the chorea may recur with subsequent pregnancies. Patients with preexisting *multiple sclerosis* (Chap. 458) experience a gradual decrease in the risk of relapses as pregnancy progresses and, conversely, an increase in attack risk during the postpartum period. Disease-modifying agents, including interferon β , should *not* be administered to pregnant multiple sclerosis patients, but moderate or severe relapses can be safely treated with pulse glucocorticoid therapy. Finally, certain tumors, particularly pituitary adenoma and meningioma (Chap. 403), may manifest during pregnancy because of accelerated growth, possibly driven by hormonal factors.

Peripheral nerve disorders associated with pregnancy include *Bell’s palsy* (idiopathic facial paralysis) (Chap. 459), which is approximately threefold more likely to occur during the third trimester and immediate postpartum period than in the general population. Therapy with glucocorticoids should follow the guidelines established for nonpregnant patients. Entrapment neuropathies are common in the later stages of pregnancy, presumably as a result of fluid retention. *Carpal tunnel syndrome* (median nerve) presents first as pain and paresthesia in the hand (often worse at night) and later with weakness in the thenar muscles. Treatment is generally conservative; wrist splints may be helpful, and glucocorticoid injections or surgical section of the carpal tunnel can usually be postponed. *Meralgia paresthetica* (lateral femoral cutaneous nerve entrapment) consists of pain and numbness in the lateral aspect of the thigh without weakness. Patients are usually reassured to learn that these symptoms are benign and can be expected to remit spontaneously after the pregnancy has been completed. *Restless leg syndrome* is the most common peripheral nerve and movement disorder in pregnancy. Disordered iron metabolism is the suspected etiology. Management is expectant in most cases.

GASTROINTESTINAL AND LIVER DISEASE

Up to 90% of pregnant women experience nausea and vomiting during the first trimester of pregnancy. *Hyperemesis gravidarum* is a severe form that prevents adequate fluid and nutritional intake and may require hospitalization to prevent dehydration and malnutrition.

Crohn’s disease may be associated with exacerbations in the second and third trimesters. Ulcerative colitis is associated with disease exacerbations in the first trimester and during the early postpartum period. Medical management of these diseases during pregnancy is similar to management in the nonpregnant state (Chap. 351).

Exacerbation of gallbladder disease is common during pregnancy. In part, this aggravation may be due to pregnancy-induced alteration in the metabolism of bile and fatty acids. Intrahepatic cholestasis of pregnancy is generally a third-trimester event. Profound pruritus may accompany this condition, and it may be associated with increased fetal mortality. Placental bile salt deposition may contribute to progressive uteroplacental insufficiency. Therefore, regular fetal surveillance should be undertaken once the diagnosis of intrahepatic cholestasis is made, and delivery should be planned once the fetus reaches about 37 weeks of gestation. Favorable results with ursodiol have been reported.

Acute fatty liver is a rare complication of pregnancy. Frequently confused with the HELLP syndrome (see “Preeclampsia” above) and severe preeclampsia, the diagnosis of acute fatty liver of pregnancy may be facilitated by imaging studies and laboratory evaluation. Acute fatty liver of pregnancy is generally characterized by markedly increased serum levels of bilirubin and ammonia and by hypoglycemia. Management of acute fatty liver of pregnancy is supportive; recurrence in subsequent pregnancies has been reported.

All pregnant women should be screened for hepatitis B. This information is important for pediatricians after delivery of the infant. All infants receive hepatitis B vaccine. Infants born to mothers who are carriers of hepatitis B surface antigen should also receive hepatitis B immune globulin as soon after birth as possible and preferably within the first 72 h. Screening for hepatitis C is recommended for individuals at high risk for exposure.

INFECTIONS

BACTERIAL INFECTIONS

Other than bacterial vaginosis, the most common bacterial infections during pregnancy involve the urinary tract (Chap. 162). Many pregnant women have asymptomatic bacteriuria, most likely due to stasis caused by progesterational effects on ureteral and bladder smooth muscle and later in pregnancy due to compression effects of the enlarging uterus. In itself, this condition is not associated with an adverse outcome of pregnancy. However, if asymptomatic bacteriuria is left untreated, symptomatic pyelonephritis may occur. Indeed, ~75% of pregnancy-associated pyelonephritis cases are the result of untreated asymptomatic bacteriuria. All pregnant women should be screened

with a urine culture for asymptomatic bacteriuria at the first prenatal visit. Subsequent screening with nitrite/leukocyte esterase strips is indicated for high-risk women, such as those with sickle cell trait or a history of urinary tract infections. All women with positive screens should be treated. Pregnant women who develop pyelonephritis need careful monitoring, including inpatient IV antibiotic administration due to the elevated risk of urosepsis and acute respiratory distress syndrome in pregnancy.

Abdominal pain and fever during pregnancy create a clinical dilemma. The diagnosis of greatest concern is intrauterine amniotic infection. While amniotic infection most commonly follows rupture of the membranes, this is not always the case. In general, antibiotic therapy is not recommended as a temporizing measure in these circumstances. If intrauterine infection is suspected, induced delivery with concomitant antibiotic therapy is generally indicated. Intrauterine amniotic infection is most often caused by pathogens such as *Escherichia coli* and group B *Streptococcus* (GBS). In high-risk patients at term or in preterm patients, routine intrapartum prophylaxis of GBS disease is recommended. Penicillin G and ampicillin are the drugs of choice. In penicillin-allergic patients with a low risk of anaphylaxis, cefazolin is recommended. If the patient is at high risk of anaphylaxis, vancomycin is recommended. If the organism is known to be sensitive to clindamycin, this antibiotic may be used. For the reduction of neonatal morbidity due to GBS, universal screening of pregnant women for GBS between 35 and 37 weeks of gestation, with intrapartum antibiotic treatment of infected women, is recommended.

Postpartum infection is a significant cause of maternal morbidity and mortality. Postpartum endomyometritis is more common after cesarean delivery than vaginal delivery and develops in 2% of women after elective repeat cesarean section and in up to 10% after emergency cesarean section following prolonged labor. To reduce the risk of endomyometritis, prophylactic antibiotics should be given to all patients undergoing cesarean section, and administration 30–60 min prior to skin incision is preferable to administration at the time of umbilical cord clamping. As most cases of postpartum endomyometritis are polymicrobial, broad-spectrum antibiotic coverage with a penicillin, an aminoglycoside, and metronidazole is recommended (Chap. 201). Most cases resolve within 72 h. Women who do not respond to antibiotic treatment for postpartum endomyometritis should be evaluated for septic pelvic thrombophlebitis. Imaging studies may be helpful in establishing the diagnosis, which is primarily a clinical diagnosis of exclusion. Patients with septic pelvic thrombophlebitis generally have tachycardia out of proportion to their fever and respond rapidly to IV administration of heparin.

All pregnant patients are screened prenatally for gonorrhea and chlamydial infections, and the detection of either should result in prompt treatment. Ceftriaxone and azithromycin are the agents of choice (Chaps. 181 and 213).

VIRAL INFECTIONS

Influenza (See also Chap. 224) Pregnant women with influenza are at increased risk of serious complications and death. All women who are pregnant or plan to become pregnant in the near future should receive inactivated influenza vaccine. The prompt initiation of antiviral treatment is recommended for pregnant women in whom influenza is suspected. Treatment can be reconsidered once the results of high-sensitivity tests are available. Prompt initiation of treatment lowers the risk of admission to an intensive care unit and death.

Cytomegalovirus Infection The most common cause of congenital viral infection in the United States is cytomegalovirus (CMV) (Chap. 219). As many as 50–90% of women of childbearing age have antibodies to CMV, but only rarely does CMV reactivation result in neonatal infection. More commonly, primary CMV infection during pregnancy creates a risk of congenital CMV. No currently accepted treatment of CMV infection during pregnancy has been demonstrated to protect the fetus effectively. Moreover, it is difficult to predict which fetus will sustain a life-threatening CMV infection. Severe CMV disease in the newborn is characterized most often by petechiae, hepatosplenomegaly, and jaundice. Chorioretinitis, microcephaly, intracranial calcifications,

hepatitis, hemolytic anemia, and purpura may also develop. CNS involvement, resulting in the development of psychomotor, ocular, auditory, and dental abnormalities over time, has been described.

Rubella (See also Chap. 230e) Rubella virus is a known teratogen; first-trimester rubella carries a high risk of fetal anomalies, though the risk significantly decreases later in pregnancy. Congenital rubella may be diagnosed by percutaneous umbilical-blood sampling with the detection of IgM antibodies in fetal blood. All pregnant women and all women of childbearing age should be tested for their immune status to rubella. All nonpregnant women who are not immune to rubella should be vaccinated. The incidence of congenital rubella in the United States is extremely low.

Herpesvirus Infection (See also Chap. 216) The acquisition of genital herpes during pregnancy is associated with spontaneous abortion, prematurity, and congenital and neonatal herpes. A cohort study of pregnant women without evidence of previous herpesvirus infection demonstrated that ~2% acquired a new herpesvirus infection during the pregnancy. Approximately 60% of the newly infected women had no clinical symptoms. Infection occurred with equal frequency in all three trimesters. If herpesvirus seroconversion occurred early in pregnancy, the risk of transmission to the newborn was very low. In women who acquired genital herpes shortly before delivery, the risk of transmission was high. The risk of active genital herpes lesions at term can be reduced by prescribing acyclovir for the last 4 weeks of pregnancy to women who have had their first episode of genital herpes during the pregnancy.

Herpesvirus infection in the newborn can be devastating. Disseminated neonatal herpes carries with it high mortality and morbidity rates from CNS involvement. It is recommended that pregnant women with active genital herpes lesions at the time of presentation in labor be delivered by cesarean section.

Parvovirus Infection (See also Chap. 221) Parvovirus infection (caused by human parvovirus B19) may occur during pregnancy. It rarely causes sequelae, but susceptible women infected during pregnancy may be at risk for fetal hydrops secondary to erythroid aplasia and profound anemia.

HIV Infection (See also Chap. 226) The predominant cause of HIV infection in children is transmission of the virus from mother to newborn during the perinatal period. All pregnant women should be screened for HIV infection. Factors that increase the risk of mother-to-newborn transmission include high maternal viral load, low maternal CD4+ T cell count, prolonged labor, prolonged duration of membrane rupture, and the presence of other genital tract infections, such as syphilis or herpes. Prior to the widespread use of antiretroviral treatment, the perinatal transmission rate was in the range of 20%. In women with a good response to antiretroviral treatment, the transmission rate is about 1%. Measurement of maternal plasma HIV RNA copy number guides the decision for vaginal versus cesarean delivery. For women with <1000 copies of plasma HIV RNA/ml who are receiving combination antiretroviral therapy, the risk of transmission to the newborn is approximately 1% regardless of mode of delivery or duration of membrane rupture. These women may elect to attempt a vaginal birth following the spontaneous onset of labor. For women with a viral load of ≥1000 copies/ml prior to 38 weeks of gestation, a scheduled prelabor cesarean at 38 weeks is recommended to reduce the risk of HIV transmission to the newborn. To reduce the risk of mother-to-newborn transmission, women with >400 copies of HIV RNA/ml should be treated during the intrapartum interval with zidovudine. All newborns of HIV-infected mothers should be treated with zidovudine for 6 months after birth. Women who are HIV-positive may transmit the virus through their breast milk. In developed countries, HIV-infected mothers are advised not to breast-feed.

VACCINATIONS

(See also Chap. 148) For rubella-nonimmune individuals contemplating pregnancy, measles-mumps-rubella vaccine should be administered, ideally at least 3 months prior to conception but otherwise in

the immediate postpartum period. In addition, pregnancy is not a contraindication for vaccination against influenza, tetanus, diphtheria, and pertussis (Tdap), and these vaccines are recommended for appropriate individuals.

MATERNAL MORTALITY

Maternal death is defined as death occurring during pregnancy or within 42 days of completion of pregnancy from a cause related to or aggravated by pregnancy, but not due to accident or incidental causes. From 1935 to 2007, the U.S. maternal death rate decreased from nearly 600/100,000 births to 12.7/100,000 births. There are significant health disparities in the maternal mortality rate, with the highest rates among non-Hispanic black women. In 2007, maternal mortality rates (per 100,000) by race were 10.5 among non-Hispanic white women, 8.9 among Hispanic women, and 28.4 among non-Hispanic black women. The most common causes of maternal death in the United States today are pulmonary embolism, obstetric hemorrhage, hypertension, sepsis, cardiovascular conditions (including peripartum cardiomyopathy), and ectopic pregnancy.



As stated above, the maternal mortality rate in the United States is about 12.7/100,000 births. In some countries in sub-Saharan Africa and southern Asia, the maternal mortality rate is about 500/100,000 live births. The most common cause of maternal death in these countries is maternal hemorrhage. The high maternal death rates are due in part to inadequate contraceptive and family-planning services, an insufficient number of skilled birth attendants, and difficulty in accessing birthing centers and emergency obstetrical care units. Maternal death is a global public-health tragedy that could be mitigated with the application of modest resources.

SUMMARY

With improved diagnostic and therapeutic modalities as well as advances in the treatment of infertility, more patients with medical complications will be seeking and will require complex obstetric care. Improved outcomes of pregnancy in these women will be best attained by a team of internists, maternal-fetal medicine (high-risk obstetrics) specialists, and anesthesiologists assembled to counsel these patients about the risks of pregnancy and to plan their treatment prior to conception. The importance of preconception counseling cannot be overstated. It is the responsibility of all physicians caring for women in the reproductive age group to assess their patients' reproductive plans as part of their overall health evaluation.

9

Medical Evaluation of the Surgical Patient

Wei C. Lau, Kim A. Eagle

Cardiovascular and pulmonary complications continue to account for major morbidity and mortality in patients undergoing noncardiac surgery. Emerging evidence-based practices dictate that the internist should perform an individualized evaluation of the surgical patient to provide an accurate preoperative risk assessment and stratification that will guide optimal perioperative risk-reduction strategies. This chapter reviews cardiovascular and pulmonary preoperative risk assessment, targeting intermediate- and high-risk patients with the goal of improving outcome. It also reviews perioperative management and prophylaxis of diabetes mellitus, endocarditis, and venous thromboembolism.

EVALUATION OF INTERMEDIATE- AND HIGH-RISK PATIENTS

Simple, standardized preoperative screening questionnaires, such as the one shown in [Table 9-1](#), have been developed for the purpose of identifying patients at intermediate or high risk who may benefit

TABLE 9-1 STANDARDIZED PREOPERATIVE QUESTIONNAIRE^a

1. Age, weight, height
2. Are you:
 - Female and 55 years of age or older or male and 45 years of age or older?
 - If yes, are you 70 years of age or older?
3. Do you take anticoagulant medications ("blood thinners")?
4. Do you have or have you had any of the following heart-related conditions?
 - Heart disease
 - Heart attack within the last 6 months
 - Angina (chest pain)
 - Irregular heartbeat
 - Heart failure
5. Do you have or have you ever had any of the following?
 - Rheumatoid arthritis
 - Kidney disease
 - Liver disease
 - Diabetes
6. Do you get short of breath when you lie flat?
7. Are you currently on oxygen treatment?
8. Do you have a chronic cough that produces any discharge or fluid?
9. Do you have lung problems or diseases?
10. Have you or any blood member of your family ever had a problem other than nausea with any anesthesia?
 - If yes, describe:
11. If female, is it possible that you are pregnant?
 - Pregnancy test:
 - Please list date of last menstrual period:

^aUniversity of Michigan Health System patient information report. Patients who answer yes to any of questions 2–9 should receive a more detailed clinical evaluation.

Source: Adapted from KK Tremper, P Benedict: *Anesthesiology* 92:1212, 2000; with permission.

from a more detailed clinical evaluation. Evaluation of such patients for surgery should always begin with a thorough history and physical examination and with a 12-lead resting electrocardiogram (ECG), in accordance with the American College of Cardiology/American Heart Association (ACC/AHA) guidelines. The history should focus on symptoms of occult cardiac or pulmonary disease. The urgency of the surgery should be determined, as true emergency procedures are associated with unavoidably higher morbidity and mortality risk. Preoperative laboratory testing should be carried out only for specific clinical conditions, as noted during clinical examination. Thus, healthy patients of any age who are undergoing elective surgical procedures without coexisting medical conditions should not require any testing unless the degree of surgical stress may result in unusual changes from the baseline state.

PREOPERATIVE CARDIAC RISK ASSESSMENT

A stepwise approach to cardiac risk assessment and stratification in patients undergoing noncardiac surgery is illustrated in [Fig. 9-1](#). Assessment of exercise tolerance in the prediction of in-hospital perioperative risk is most helpful in patients who self-report worsening exercise-induced cardiopulmonary symptoms; those who may benefit from noninvasive or invasive cardiac testing regardless of a scheduled surgical procedure; and those with known coronary artery disease (CAD) or with multiple risk factors who are able to exercise. For predicting perioperative events, poor exercise tolerance has been defined as the inability to walk four blocks or climb two flights of stairs at a normal pace or to meet a metabolic equivalent (MET) level of 4 (e.g., carrying objects of 15–20 lb or playing golf or doubles tennis) because of the development of dyspnea, angina, or excessive fatigue ([Table 9-2](#)).

Previous studies have compared several cardiac risk indices. The American College of Surgeons' National Surgical Quality Improvement

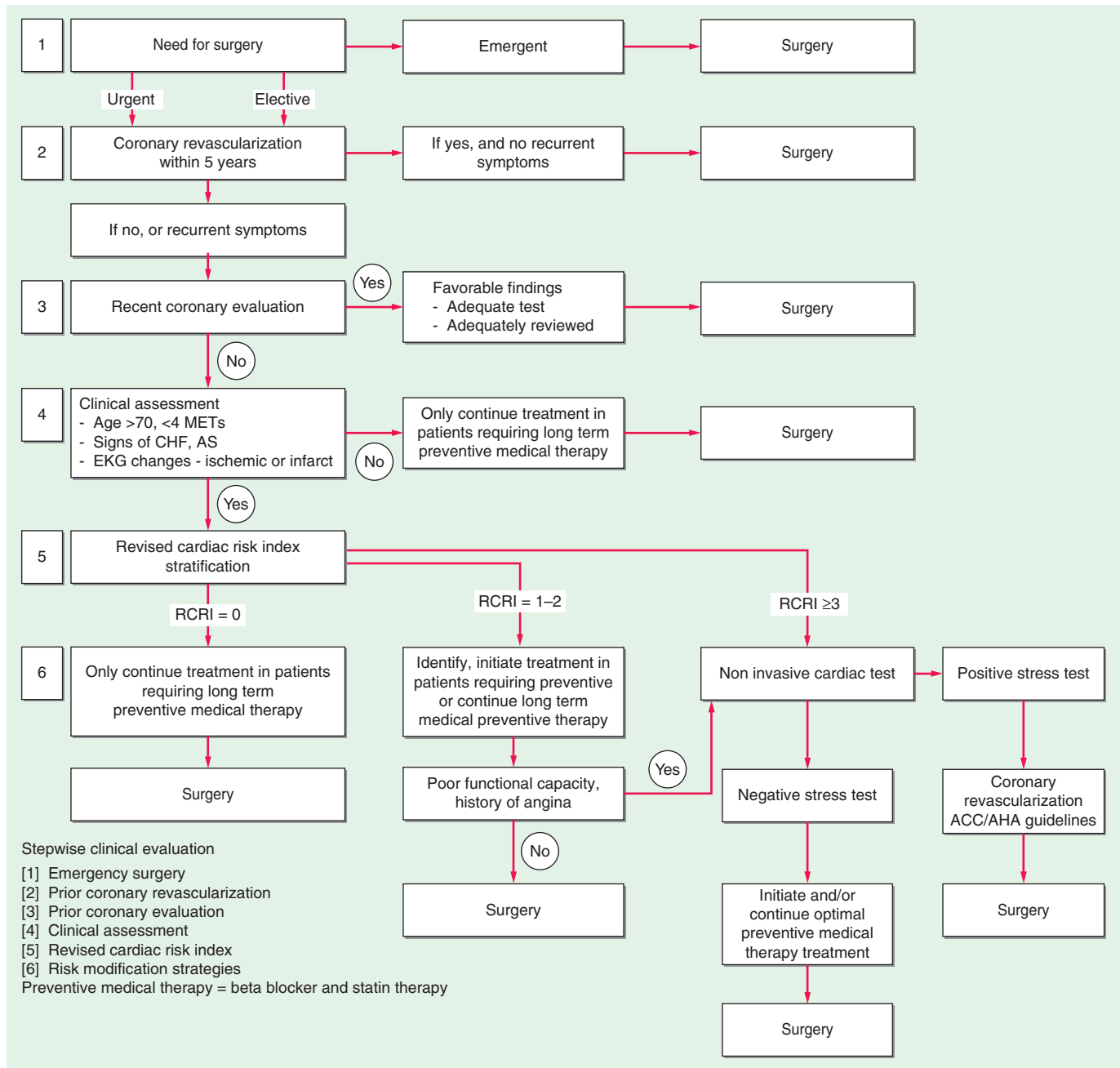


FIGURE 9-1 Composite algorithm for cardiac risk assessment and stratification in patients undergoing noncardiac surgery. Stepwise clinical evaluation: [1] emergency surgery; [2] prior coronary revascularization; [3] prior coronary evaluation; [4] clinical assessment; [5] RCRI; [6] risk modification strategies. Preventive medical therapy = beta blocker and statin therapy. RCRI, revised cardiac risk index. (Adapted from LA Fleisher et al: *Circulation* 116:1971, 2007, with permission.)

Program prospective database has identified five predictors of perioperative myocardial infarction (MI) and cardiac arrest based on increasing age, American Society of Anesthesiologists class, type of surgery, dependent functional status, and abnormal serum creatinine level. However, given its accuracy and simplicity, the revised cardiac risk

TABLE 9-2 ASSESSMENT OF CARDIAC RISK BY FUNCTIONAL STATUS

Risk	Higher	<ul style="list-style-type: none"> • Has difficulty with adult activities of daily living • Cannot walk four blocks or up two flights of stairs or does not meet a MET level of 4
	Lower	<ul style="list-style-type: none"> • Is inactive but has no limitations • Is active: easily does vigorous tasks • Performs regular vigorous exercises

Source: From LA Fleisher et al: *Circulation* 116:1971, 2007.

index (RCRI) (Table 9-3) is favored. The RCRI relies on the presence or absence of six identifiable predictive factors: high-risk surgery, ischemic heart disease, congestive heart failure, cerebrovascular disease, diabetes mellitus, and renal dysfunction. Each of these predictors is assigned one point. The risk of major cardiac events—defined as myocardial infarction, pulmonary edema, ventricular fibrillation or primary cardiac arrest, and complete heart block—can then be predicted. Based on the presence of none, one, two, three, or more of these clinical predictors, the rate of development of one of these four major cardiac events is estimated to be 0.4, 0.9, 7, and 11%, respectively (Fig. 9-2). An RCRI score of 0 signifies a 0.4–0.5% risk of cardiac events; RCRI 1, 0.9–1.3%; RCRI 2, 4–7%; and RCRI ≥ 3 , 9–11%. The clinical utility of the RCRI is to identify patients with three or more predictors who are at very high risk ($\geq 11\%$) for cardiac complications and who may benefit from further risk stratification with noninvasive cardiac testing or initiation of preoperative preventive medical management.

TABLE 9-3 CLINICAL MARKERS INCLUDED IN THE REVISED CARDIAC RISK INDEX

High-Risk Surgical Procedures
Vascular surgery
Major intraperitoneal or intrathoracic procedures
Ischemic Heart Disease
History of myocardial infarction
Current angina considered to be ischemic
Requirement for sublingual nitroglycerin
Positive exercise test
Pathological Q-waves on ECG
History of PCI and/or CABG with current angina considered to be ischemic
Congestive Heart Failure
Left ventricular failure by physical examination
History of paroxysmal nocturnal dyspnea
History of pulmonary edema
S ₃ gallop on cardiac auscultation
Bilateral rales on pulmonary auscultation
Pulmonary edema on chest x-ray
Cerebrovascular Disease
History of transient ischemic attack
History of cerebrovascular accident
Diabetes Mellitus
Treatment with insulin
Chronic Renal Insufficiency
Serum creatinine >2 mg/dL

Abbreviations: CABG, coronary artery bypass grafting; ECG, electrocardiogram; PCI, percutaneous coronary interventions.

Source: Adapted from TH Lee et al: *Circulation* 100:1043, 1999.

PREOPERATIVE NONINVASIVE CARDIAC TESTING FOR RISK STRATIFICATION

There is little evidence to support widespread application of preoperative noninvasive cardiac testing for all patients undergoing major surgery. Rather, a discriminative approach based on clinical risk categorization appears to be both clinically useful and cost-effective. There is potential benefit in identifying asymptomatic but high-risk patients, such as those with left main or left main–equivalent CAD or those with three-vessel CAD and poor left ventricular function, who may benefit from coronary revascularization (Chap. 293). However, evidence does not support aggressive attempts to identify patients at intermediate risk who have asymptomatic but advanced coronary artery disease, in whom coronary revascularization appears to offer little advantage over medical therapy.

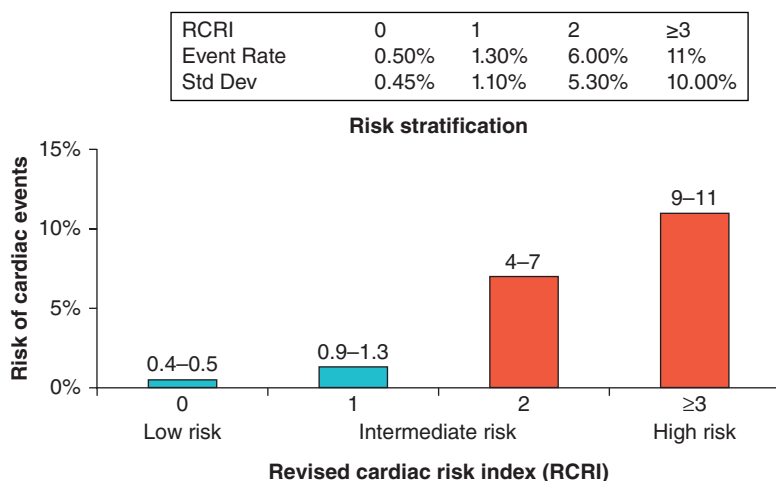


FIGURE 9-2 Risk stratification based on the RCRI: derivation and prospective validation of a simple index for prediction of cardiac risk in patients undergoing major noncardiac surgery. Cardiac events include myocardial infarction, pulmonary edema, ventricular fibrillation, cardiac asystole, and complete heart block. (Adapted from TH Lee et al: *Circulation* 100:1043, 1999.)

An RCRI score ≥ 3 in patients with severe myocardial ischemia on stress testing should lead to consideration of coronary revascularization prior to noncardiac surgery. Noninvasive cardiac testing is most appropriate if it is anticipated that, in the event of a strongly positive test, a patient will meet guidelines for coronary angiography and coronary revascularization. Pharmacologic stress tests are more useful than exercise testing in patients with functional limitations. Dobutamine echocardiography and persantine, adenosine, or dobutamine nuclear perfusion testing (Chap. 270e) have excellent negative predictive values (near 100%) but poor positive predictive values (<20%) in the identification of patients at risk for perioperative MI or death. Thus, a negative study is reassuring, but a positive study is a relatively weak predictor of a “hard” perioperative cardiac event.

RISK MODIFICATION: PREVENTIVE STRATEGIES TO REDUCE CARDIAC RISK

Perioperative Coronary Revascularization Currently, potential options for reducing perioperative cardiovascular risk include coronary artery revascularization and/or perioperative preventive medical therapies (Chap. 293). *Prophylactic* coronary revascularization with either coronary artery bypass grafting (CABG) or percutaneous coronary intervention (PCI) provides no short- or midterm survival benefit for patients *without* left main CAD or three-vessel CAD in the presence of poor left ventricular systolic function and is *not recommended* for patients with stable CAD before noncardiac surgery. Although PCI is associated with lower procedural risk than is CABG in the perioperative setting, the placement of a coronary artery stent soon before noncardiac surgery may increase the risk of bleeding during surgery if dual antiplatelet therapy (aspirin and thienopyridine) is administered; moreover, stent placement shortly before noncardiac surgery increases the perioperative risk of MI and cardiac death due to stent thrombosis if such therapy is withdrawn prematurely (Chap. 296e). It is recommended that, if possible, noncardiac surgery be delayed 30–45 days after placement of a bare metal coronary stent and for 365 days after a drug-eluting stent. For patients who *must* undergo noncardiac surgery early (>14 days) after PCI, balloon angioplasty without stent placement appears to be a reasonable alternative because dual antiplatelet therapy is not necessary in such patients. One recent clinical trial further suggests that after 6 months, bare metal and drug eluting stents may not pose a threat.

PERIOPERATIVE PREVENTIVE MEDICAL THERAPIES The goal of perioperative preventive medical therapies with β -adrenergic antagonists, HMG-CoA reductase inhibitors (statins), antiplatelet agents, and α_2 agonists is to reduce perioperative adrenergic stimulation, ischemia, and inflammation, which are triggered during the perioperative period.

β -ADRENERGIC ANTAGONISTS The use of perioperative beta blockade should be based on a thorough assessment of a patient’s perioperative clinical and surgery-specific cardiac risk (RCRI ≥ 2). For patients with or without mild to moderate reactive airway disease, the cardioselective beta blocker of choice should be used and titrated to maintain a target heart rate of 60–80 beats/min in the absence of hypotension in the operative and perioperative period. In RCRI ≥ 2 patients without a long-term indication for beta blockers, the medications can be administered IV as a preoperative medication on the day of surgery, with a targeted heart rate of 60–80 beats/min without hypotension, and continued for >7 days (preferably 30 days) postoperatively. Abrupt perioperative beta blocker withdrawal should be avoided unless necessary because of the associated increase in risk of MI and angina. IV preparations should be substituted for oral medications if patients are unable to swallow or absorb pills in the perioperative period. The results from the Perioperative Ischemic Evaluation (POISE) trial showed that, although cardiac death, nonfatal myocardial infarction, or cardiac arrest was reduced among patients who received metoprolol rather than placebo, there was an increased incidence of death and stroke among metoprolol recipients because of a high and rapidly loading dose of this drug.

The POISE trial highlights the importance of a clear risk-and-benefit assessment, with careful initiation and titration to therapeutic efficacy of preoperative beta blockers in patients undergoing noncardiac surgery. A recent meta-analysis which included the POISE study further supports that excessive beta blocker dosing is, in fact, harmful.

The ACC/AHA guidelines recommend the following: (1) Beta blockers *should be continued* in patients with active cardiac conditions who are undergoing surgery and are receiving beta blockers. (2) Beta blockers titrated to heart rate and blood pressure are *probably recommended* for patients undergoing vascular surgery who are at high cardiac risk defined by CAD or cardiac ischemia on preoperative testing. (3) Beta blockers are *reasonable* for high-risk patients (RCRI ≥ 2) who undergo vascular surgery. (4) Beta blockers are *reasonable* for patients with known CAD or high risk (RCRI ≥ 2) who undergo intermediate-risk surgery. (5) Nondiscriminant administration of high-dose beta blockers without dose titration to effectiveness is *contraindicated* for patients who have never been treated with a beta blocker.

HMG-CoA REDUCTASE INHIBITORS (STATINS) A number of prospective and retrospective studies support the perioperative prophylactic use of statins for reduction of cardiac complications in patients with established atherosclerosis. The ACC/AHA Guidelines support the protective efficacy of perioperative statins on cardiac complications in intermediate risk patients undergoing major noncardiac surgery. For patients undergoing noncardiac surgery and currently taking statins, statin therapy *should be continued* to reduce perioperative cardiac risk. Statins are *reasonable* for patients undergoing vascular surgery with or without clinical risk factors (RCRI ≥ 1).

ANGIOTENSIN-CONVERTING ENZYME (ACE) INHIBITORS Evidence supports the discontinuation of ACE inhibitors and angiotensin receptor blockers for 24 h prior to noncardiac surgery due to adverse circulatory effects after induction of anesthesia.

ORAL ANTIPLATELET AGENTS Evidence-based recommendations regarding perioperative use of aspirin and/or thienopyridine to reduce cardiac risk currently lack clarity. A substantial increase in perioperative bleeding and in the need for transfusion in patients receiving dual antiplatelet therapy has been observed. The discontinuation of thienopyridine and aspirin for 5–7 days prior to major surgery to minimize the risk of perioperative bleeding and transfusion must be balanced with the potential increased risk of an acute coronary syndrome and of subacute stent thrombosis in patients with recent coronary stent implantation. If clinicians elect to withhold antiplatelet agents prior to surgery, these agents should be restarted as soon as possible postoperatively.

α_2 AGONISTS Several prospective and retrospective meta-analyses of perioperative α_2 agonists (clonidine and mivazerol) demonstrated a reduction of cardiac death rates among patients with known coronary artery disease who underwent noncardiac surgery. α_2 agonists thus *may be considered* for perioperative control of hypertension in patients with known coronary artery disease or an RCRI score ≥ 2 .

CALCIUM CHANNEL BLOCKERS Evidence is lacking to support the use of calcium channel blockers as a prophylactic strategy to decrease perioperative risk in major noncardiac surgery.

ANESTHETICS Mortality risk is low with safe delivery of modern anesthesia, especially among low-risk patients undergoing low-risk surgery (Table 9-4). Inhaled anesthetics have predictable circulatory and respiratory effects: all decrease arterial pressure in a dose-dependent manner by reducing sympathetic tone and causing systemic vasodilation, myocardial depression, and decreased cardiac output. Inhaled anesthetics also cause respiratory depression, with diminished responses to both hypercapnia and hypoxemia, in a dose-dependent manner; in addition, these agents have a variable effect on heart rate. Prolonged residual neuromuscular blockade also increases the risk of postoperative pulmonary complications due to reduction in functional residual lung capacity, loss of diaphragmatic and intercostal muscle function, atelectasis, and arterial hypoxemia from ventilation-perfusion mismatch.

TABLE 9-4 GRADATION OF MORTALITY RISK OF COMMON NONCARDIAC SURGICAL PROCEDURES

Higher	<ul style="list-style-type: none"> Emergent major operations, especially in the elderly Aortic and other noncarotid major vascular surgery (endovascular and nonendovascular) Prolonged surgery associated with large fluid shift and/or blood loss
Intermediate	<ul style="list-style-type: none"> Major thoracic surgery Major abdominal surgery Carotid endarterectomy surgery Head/neck surgery Orthopedic surgery Prostate surgery
Lower	<ul style="list-style-type: none"> Eye, skin, and superficial surgery Endoscopic procedures

Source: From LA Fleisher et al: Circulation 116:1971, 2007, with permission.

Several meta-analyses have shown that rates of pneumonia and respiratory failure are lower among patients receiving neuroaxial anesthesia (epidural or spinal) rather than general anesthesia (inhaled). However, there were no significant differences in cardiac events between the two approaches. Evidence from a meta-analysis of randomized controlled trials supports postoperative epidural analgesia for >24 h for the purpose of pain relief. However, the risk of epidural hematoma in the setting of systemic anticoagulation for venous thromboembolism prophylaxis (see below) and postoperative epidural catheterization must be considered.

PREOPERATIVE PULMONARY RISK ASSESSMENT

Perioperative pulmonary complications occur frequently and lead to significant morbidity and mortality. The guidelines from the American College of Physicians recommend the following:

- All patients undergoing noncardiac surgery should be assessed for risk of pulmonary complications (Table 9-5).
- Patients undergoing emergency or prolonged (3- to 4-h) surgery; aortic aneurysm repair; vascular surgery; major abdominal, thoracic, neurologic, head, or neck surgery; and general anesthesia

TABLE 9-5 PREDISPOSING RISK FACTORS FOR PULMONARY COMPLICATIONS

- Upper respiratory tract infection: cough, dyspnea
- Age >60 years
- Chronic obstructive pulmonary disease
- Cigarette use
- American Society of Anesthesiologists Class ≥ 2
- Functional dependence
- Congestive heart failure
- Serum albumin <3.5 g/dL
- Obstructive sleep apnea
- Impaired sensorium (confusion, delirium, or mental status changes)
- Abnormal findings on chest examination
- Alcohol use
- Weight loss
- Spirometry threshold before lung resection
 - FEV₁ <2 L
 - MVV <50% of predicted
 - PEF <100 L or 50% predicted value
 - PCO₂ ≥ 45 mmHg
 - PO₂ ≤ 50 mmHg

Abbreviations: FEV₁, forced expiratory volume in 1 s; MVV, maximal voluntary ventilation; PEF, peak expiratory flow rate; PCO₂, partial pressure of carbon dioxide; PO₂, partial pressure of oxygen.

Source: A Qaseem et al: Ann Intern Med 144:575-80. Modified from GW Smetana et al: Ann Intern Med 144:581, 2006, and from DN Mohr et al: Postgrad Med 100:247, 1996.

TABLE 9-6 RISK MODIFICATION TO REDUCE PERIOPERATIVE PULMONARY COMPLICATIONS

Preoperatively
<ul style="list-style-type: none"> • Cessation of smoking for at least 8 weeks before and until at least 10 days after surgery • Training in proper lung expansion techniques • Inhalation bronchodilator and/or steroid therapy, when indicated • Control of infection and secretion, when indicated • Weight reduction, when appropriate
Intraoperatively
<ul style="list-style-type: none"> • Limited duration of anesthesia • Avoidance of long-acting neuromuscular blocking drugs, when indicated • Prevention of aspiration and maintenance of optimal bronchodilation
Postoperatively
<ul style="list-style-type: none"> • Optimization of inspiratory capacity maneuvers, with attention to: <ul style="list-style-type: none"> Mobilization of secretions Early ambulation Encouragement of coughing Selective use of a nasogastric tube • Adequate pain control without excessive narcotics

Source: From VA Lawrence et al: *Ann Intern Med* 144:596, 2006, and WF Dunn, PD Scanlon: *Mayo Clin Proc* 68:371, 1993.

should be considered to be at elevated risk for postoperative pulmonary complications.

3. Patients at higher risk of pulmonary complications should undergo incentive spirometry, deep-breathing exercises, cough encouragement, postural drainage, percussion and vibration, suctioning and ambulation, intermittent positive-pressure breathing, continuous positive airway pressure, and selective use of a nasogastric tube for postoperative nausea, vomiting, or symptomatic abdominal distention to reduce postoperative risk (Table 9-6).
4. Routine preoperative spirometry and chest radiography should not be used routinely for predicting risk of postoperative pulmonary complications but may be appropriate for patients with chronic obstructive pulmonary disease or asthma.
5. Spirometry is of value before lung resection in determining candidacy for coronary artery bypass; however, it does not provide a spirometric threshold for extrathoracic surgery below which the risks of surgery are unacceptable.
6. Pulmonary artery catheterization, administration of total parenteral nutrition (as opposed to no supplementation), or total enteral nutrition has no benefit in reducing postoperative pulmonary complications.

PERIOPERATIVE MANAGEMENT AND PROPHYLAXIS

DIABETES MELLITUS

(See also Chaps. 417–419) Many patients with diabetes mellitus have significant symptomatic or asymptomatic CAD and may have silent myocardial ischemia due to autonomic dysfunction. Evidence supports intensive perioperative glycemic control to achieve near-normal glucose levels (90–110 mg/dL) rather than moderate glycemic control (120–200 mg/dL), using insulin infusion. This practice must be balanced against the risk of hypoglycemic complications. Oral hypoglycemic agonists should not be given on the morning of surgery. Perioperative hyperglycemia should be treated with IV infusion of short-acting insulin or SC sliding-scale insulin. Patients whose diabetes is diet controlled may proceed to surgery with close postoperative monitoring.

INFECTIVE ENDOCARDITIS

(See also Chap. 155) Perioperative prophylactic antibiotics should be administered to patients with congenital or valvular heart disease, prosthetic valves, mitral valve prolapse, or other cardiac abnormalities, in accordance with ACC/AHA practice guidelines.

VENOUS THROMBOEMBOLISM

(See also Chap. 300) Perioperative prophylaxis of venous thromboembolism should follow established guidelines of the American College of Chest Physicians. Aspirin is not supported as a single agent for thromboprophylaxis. Low-dose unfractionated heparin (≤ 5000 units SC bid), low-molecular weight heparin (e.g., enoxaparin, 30 mg bid or 40 mg qd), or a pentasaccharide (fondaparinux, 2.5 mg qd) is appropriate for patients at moderate risk; unfractionated heparin (5000 units SC tid) is appropriate for patients at high risk. Graduated compression stockings and pneumatic compression devices are useful supplements to anticoagulant therapy.

10

Palliative and End-of-Life Care

Ezekiel J. Emanuel

EPIDEMIOLOGY

In 2010, according to the Centers for Disease Control and Prevention, 2,468,435 individuals died in the United States (Table 10-1). Approximately 73% of all deaths occur in those >65 years of age. The epidemiology of mortality is similar in most developed countries; cardiovascular diseases and cancer are the predominant causes of death, a marked change since 1900, when heart disease caused $\sim 8\%$ of all deaths and cancer accounted for $<4\%$ of all deaths. In 2010, the year with the most recent available data, AIDS did not rank among the top 15 causes of death, causing just 8369 deaths. Even among people age 35–44, heart disease, cancer, chronic liver disease, and accidents all cause more deaths than AIDS.

It is estimated that in developed countries $\sim 70\%$ of all deaths are preceded by a disease or condition, making it reasonable to plan for dying in the foreseeable future. Cancer has served as the paradigm for terminal care, but it is not the only type of illness with a recognizable and predictable terminal phase. Because heart failure, chronic obstructive pulmonary disease (COPD), chronic liver failure, dementia, and many other conditions have recognizable terminal phases, a systematic approach to end-of-life care should be part of all medical specialties. Many patients with illness-related suffering also can benefit from palliative care regardless of prognosis. Ideally, palliative care should be considered part of comprehensive care for all patients. Palliative care can be improved by coordination between caregivers, doctors, and patients for advance care planning, as well as dedicated teams of physicians, nurses, and other providers.

The rapid increases in life expectancy in developed countries over the last century have been accompanied by new difficulties facing individuals, families, and society as a whole in addressing the needs of an aging population. These challenges include both more complicated conditions and technologies to address them at the end of life. The development of technologies that can prolong life without restoring full health has led many Americans to seek out alternative end-of-life care settings and approaches that relieve suffering for those with terminal diseases. Over the last few decades in the United States, a significant change in the site of death has occurred that coincides with patient and family preferences. Nearly 60% of Americans died as inpatients in hospitals in 1980. By 2000, the trend was reversing, with $\sim 31\%$ of Americans dying as hospital inpatients (Fig. 10-1). This shift has been most dramatic for those dying from cancer and COPD and for younger and very old individuals. In the last decade, it has been associated with the increased use of hospice care; in 2008, approximately 39% of all decedents in the United States received such care. Cancer patients currently constitute $\sim 36.9\%$ of hospice users. About 79% of patients receiving hospice care die out of the hospital, and around 42% of those receiving hospice care die in a private residence. In addition, in 2008, for the first time, the American Board of Medical Specialties (ABMS) offered certification in hospice and palliative medicine. With shortening of hospital stays,

TABLE 10-1 TEN LEADING CAUSES OF DEATH IN THE UNITED STATES AND BRITAIN

Cause of Death	United States			Britain	
	Number of Deaths	Percentage of Total	Number of Deaths Among People ≥ 65 Years of Age	Number of Deaths	Percentage of Total
All deaths	2,468,435	100	1,798,276	499,331	100
Heart disease	597,689	24.2	477,338	141,362	28.3
Malignant neoplasms	574,743	23.3	396,670	142,107	28.5
Chronic lower respiratory diseases	138,080	5.6	118,031	27,132	5.4
Cerebrovascular diseases	129,476	5.2	109,990	35,846	7.2
Accidents	120,859	4.9	41,300	11,256	2.3
Alzheimer's disease	83,494	3.4	82,616	8859	1.8
Diabetes mellitus	69,071	2.8	49,191	4931	1.0
Nephritis, nephritic syndrome, nephrosis	50,476	2.0	41,994	4102	0.8
Influenza and pneumonia	50,097	2.0	42,846	26,138	5.2
Intentional self-harm	38,364	1.6	6008	3671	0.7

Source: National Center for Health Statistics (data for all age groups from 2010), <http://www.cdc.gov/nchs>; National Statistics (England and Wales, 2012), <http://www.statistics.gov.uk>.

many serious conditions are being treated at home or on an outpatient basis. Consequently, providing optimal palliative and end-of-life care requires ensuring that appropriate services are available in a variety of settings, including noninstitutional settings.

HOSPICE AND THE PALLIATIVE CARE FRAMEWORK

Central to this type of care is an interdisciplinary team approach that typically encompasses pain and symptom management, spiritual and psychological care for the patient, and support for family caregivers during the patient's illness and the bereavement period.

Terminally ill patients have a wide variety of advanced diseases, often with multiple symptoms that demand relief, and require non-invasive therapeutic regimens to be delivered in flexible care settings. Fundamental to ensuring quality palliative and end-of-life care is a focus on four broad domains: (1) physical symptoms; (2) psychological symptoms; (3) social needs that include interpersonal relationships, caregiving, and economic concerns; and (4) existential or spiritual needs.

A comprehensive assessment screens for and evaluates needs in each of these four domains. Goals for care are established in discussions with the patient and/or family, based on the assessment in each of the domains. Interventions then are aimed at improving or managing symptoms and needs. Although physicians are responsible

for certain interventions, especially technical ones, and for coordinating the interventions, they cannot be responsible for providing all of them. Because failing to address any one of the domains is likely to preclude a good death, a well-coordinated, effectively communicating interdisciplinary team takes on special importance in end-of-life care. Depending on the setting, critical members of the interdisciplinary team will include physicians, nurses, social workers, chaplains, nurse's aides, physical therapists, bereavement counselors, and volunteers.

ASSESSMENT AND CARE PLANNING

Comprehensive Assessment Standardized methods for conducting a comprehensive assessment focus on evaluating the patient's condition in all four domains affected by illness: physical, psychological, social, and spiritual. The assessment of physical and mental symptoms should follow a modified version of the traditional medical history and physical examination that emphasizes symptoms. Questions should aim at elucidating symptoms and discerning sources of suffering and gauging how much those symptoms interfere with the patient's quality of life. Standardized assessment is critical. Currently, there are 21 symptom assessment instruments for cancer alone. Further research on and validation of these assessment tools, especially taking into account patient perspectives, could improve their effectiveness. Instruments with good psychometric properties that assess a wide range of symptoms include the Memorial Symptom Assessment Scale (MSAS), the Rotterdam Symptom Checklist, the Worthing Chemotherapy Questionnaire, and the Computerized Symptom Assessment Instrument. These instruments are long and may be useful for initial clinical or for research assessments. Shorter instruments are useful for patients whose performance status does not permit comprehensive assessments. Suitable shorter instruments include the Condensed Memorial Symptom Assessment Scale, the Edmonton Symptom Assessment System, the M.D. Anderson Symptom Assessment Inventory, and the Symptom Distress Scale. Using such instruments ensures that the assessment is comprehensive and does not focus only on pain and a few other physical symptoms. Invasive tests are best avoided in end-of-life care, and even minimally invasive tests should be evaluated carefully for their benefit-to-burden ratio for the patient. Aspects of the physical examination that are uncomfortable and unlikely to yield useful information can be omitted.

Regarding social needs, health care providers should assess the status of important relationships, financial burdens, caregiving needs, and access to medical care. Relevant questions will include the following: *How often is there someone to feel close to? How has this illness been for your family? How has it affected your relationships? How much help do you need with things like getting meals and getting around? How much trouble do you have getting the medical care you need?* In the area of existential needs, providers should assess distress and the

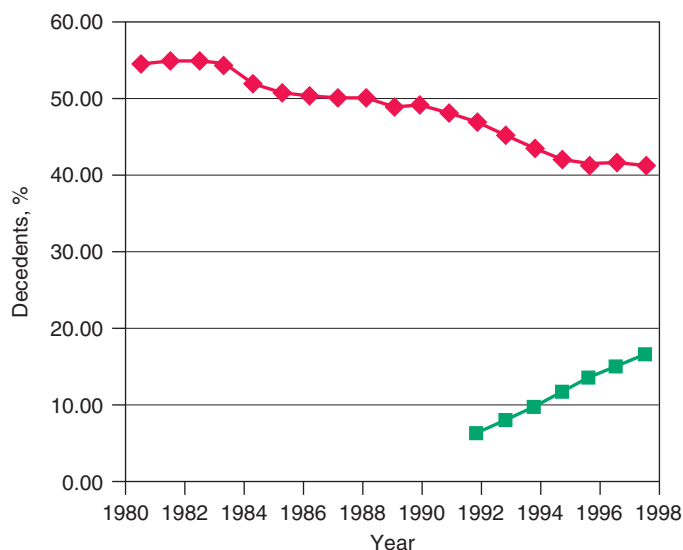


FIGURE 10-1 Graph showing trends in the site of death in the last two decades. \blacklozenge , percentage of hospital inpatient deaths; \blacksquare , percentage of decedents enrolled in a hospice.

patient's sense of being emotionally and existentially settled and of finding purpose or meaning. Helpful assessment questions can include the following: *How much are you able to find meaning since your illness began? What things are most important to you at this stage?* In addition, it can be helpful to ask how the patient perceives his or her care: *How much do you feel your doctors and nurses respect you? How clear is the information from us about what to expect regarding your illness? How much do you feel that the medical care you are getting fits with your goals?* If concern is detected in any of these areas, deeper evaluative questions are warranted.

Communication Especially when an illness is life-threatening, there are many emotionally charged and potentially conflict-creating moments, collectively called “bad news” situations, in which empathic and effective communication skills are essential. Those moments include communicating with the patient and/or family about a terminal diagnosis, the patient's prognosis, any treatment failures, deemphasizing efforts to cure and prolong life while focusing more on symptom management and palliation, advance care planning, and the patient's death. Although these conversations can be difficult and lead to tension, research indicates that end-of-life discussions can lead to earlier

hospice referrals rather than overly aggressive treatment, benefiting quality of life for patients and improving the bereavement process for families.

Just as surgeons plan and prepare for major operations and investigators rehearse a presentation of research results, physicians and health care providers caring for patients with significant or advanced illness can develop a practiced approach to sharing important information and planning interventions. In addition, families identify as important both how well the physician was prepared to deliver bad news and the setting in which it was delivered. For instance, 27% of families making critical decisions for patients in an intensive care unit (ICU) desired better and more private physical space to communicate with physicians, and 48% found having clergy present reassuring.

An organized and effective seven-step procedure for communicating bad news goes by the acronym P-SPIKES: (1) **p**repare for the discussion, (2) **s**et up a suitable environment, (3) **b**egin the discussion by finding out what the **p**atient and/or family understand, (4) **d**etermine how they will comprehend new **i**nformation best and how much they want to know, (5) **p**rovide needed new **k**nowledge accordingly, (6) **a**llow for **e**motional responses, and (7) **s**hare plans for the next steps in care. **Table 10-2** provides a summary of these steps along with

TABLE 10-2 ELEMENTS OF COMMUNICATING BAD NEWS—THE P-SPIKES APPROACH

Acronym	Steps	Aim of the Interaction	Preparations, Questions, or Phrases
P	Preparation	Mentally prepare for the interaction with the patient and/or family.	Review what information needs to be communicated. Plan how you will provide emotional support. Rehearse key steps and phrases in the interaction.
S	Setting of the interaction	Ensure the appropriate setting for a serious and potentially emotionally charged discussion.	Ensure that patient, family, and appropriate social supports are present. Devote sufficient time. Ensure privacy and prevent interruptions by people or beeper. Bring a box of tissues.
P	Patient's perception and preparation	Begin the discussion by establishing the baseline and whether the patient and family can grasp the information. Ease tension by having the patient and family contribute.	Start with open-ended questions to encourage participation. Possible phrases to use: <i>What do you understand about your illness?</i> <i>When you first had symptom X, what did you think it might be?</i> <i>What did Dr. X tell you when he or she sent you here?</i> <i>What do you think is going to happen?</i>
I	Invitation and information needs	Discover what information needs the patient and/or family have and what limits they want regarding the bad information.	Possible phrases to use: <i>If this condition turns out to be something serious, do you want to know?</i> <i>Would you like me to tell you all the details of your condition? If not, who would you like me to talk to?</i>
K	Knowledge of the condition	Provide the bad news or other information to the patient and/or family sensitively.	Do not just dump the information on the patient and family. Check for patient and family understanding. Possible phrases to use: <i>I feel badly to have to tell you this, but . . .</i> <i>Unfortunately, the tests showed . . .</i> <i>I'm afraid the news is not good . . .</i>
E	Empathy and exploration	Identify the cause of the emotions—e.g., poor prognosis. Empathize with the patient and/or family's feelings. Explore by asking open-ended questions.	Strong feelings in reaction to bad news are normal. Acknowledge what the patient and family are feeling. Remind them such feelings are normal, even if frightening. Give them time to respond. Remind patient and family you won't abandon them. Possible phrases to use: <i>I imagine this is very hard for you to hear.</i> <i>You look very upset. Tell me how you are feeling.</i> <i>I wish the news were different.</i> <i>We'll do whatever we can to help you.</i>
S	Summary and planning	Delineate for the patient and the family the next steps, including additional tests or interventions.	It is the unknown and uncertain that can increase anxiety. Recommend a schedule with goals and landmarks. Provide your rationale for the patient and/or family to accept (or reject). If the patient and/or family are not ready to discuss the next steps, schedule a follow-up visit.

Source: Adapted from R Buckman: *How to Break Bad News: A Guide for Health Care Professionals*. Baltimore, Johns Hopkins University Press, 1992.

suggested phrases and underlying rationales for each one. Additional research that further considers the response of patients to systematic methods of delivering bad news could build the evidence base for even more effective communication procedures.

Continuous Goal Assessment Major barriers to ensuring quality palliative and end-of-life care include difficulty providing an accurate prognosis and emotional resistance of patients and their families to accepting the implications of a poor prognosis. There are two practical solutions to these barriers. One is to integrate palliative care with curative care regardless of prognosis. With this approach, palliative care no longer conveys the message of failure, having no more treatments, or “giving up hope.” Fundamental to integrating palliative care with curative therapy is to include continuous goal assessment as part of the routine patient reassessment that occurs at most patient-physician encounters. Alternatively, some practices may find it useful to implement a standard point in the clinical course to address goals of care and advance care planning. For example, some oncology practices ask all patients whose Eastern Cooperative Oncology Group (ECOG) performance status is 3 or less—meaning they spend 50% or more of the day in bed—or those who develop metastatic disease about their goals of care and advance care preferences.

Goals for care are numerous, ranging from cure of a specific disease, to prolonging life, to relief of a symptom, to delaying the course of an incurable disease, to adapting to progressive disability without disrupting the family, to finding peace of mind or personal meaning, to dying in a manner that leaves loved ones with positive memories. Discernment of goals for care can be approached through a seven-step protocol: (1) ensure that medical and other information is as complete as reasonably possible and is understood by all relevant parties (see above); (2) explore what the patient and/or family are hoping for while identifying relevant and realistic goals; (3) share all the options with the patient and family; (4) respond with empathy as they adjust to changing expectations; (5) make a plan, emphasizing what can be done toward achieving the realistic goals; (6) follow through with the plan; and (7) review and revise the plan periodically, considering at every encounter whether the goals of care should be reviewed with the patient and/or family. Each of these steps need not be followed in rote order, but together they provide a helpful framework for interactions with patients and their families about goals for care. It can be especially challenging if a patient or family member has difficulty letting go of an unrealistic goal. One strategy is to help them refocus on more realistic goals and also suggest that while hoping for the best, it is still prudent to plan for other outcomes as well.

Advance Care Planning • PRACTICES Advance care planning is a process of planning for future medical care in case the patient becomes incapable of making medical decisions. A 2010 study of adults 60 or older who died between 2000 and 2006 found that 42% required decision making about treatment in the final days of life but 70% lacked decision-making capacity. Among those lacking decision-making capacity, around one-third did not have advance planning directives. Ideally, such planning would occur before a health care crisis or the terminal phase of an illness. Diverse barriers prevent this. Polls suggest 80% of Americans endorse advance care planning and completing living wills. However, data suggest between 33 and 42% have actually completed one. Other countries have even lower completion rates. Most patients expect physicians to initiate advance care planning and will wait for physicians to broach the subject. Patients also wish to discuss advance care planning with their families. Yet patients with unrealistic expectations are significantly more likely to prefer aggressive treatments. Fewer than one-third of health care providers have completed advance care planning for themselves. Hence, a good first step is for health care providers to complete their own advance care planning. This makes providers aware of the critical choices in the process and the issues that are especially charged and allows them to tell their patients truthfully that they personally have done advance planning. Lessons from behavioral economics suggest that setting this kind of social norming helps people view completing an advance directive as acceptable and even expected.

Steps in effective advance care planning center on (1) introducing the topic, (2) structuring a discussion, (3) reviewing plans that have been discussed by the patient and family, (4) documenting the plans, (5) updating them periodically, and (6) implementing the advance care directives (Table 10-3). Two of the main barriers to advance care planning are problems in raising the topic and difficulty in structuring a succinct discussion. Raising the topic can be done efficiently as a routine matter, noting that it is recommended for all patients, analogous to purchasing insurance or estate planning. Many of the most difficult cases have involved unexpected, acute episodes of brain damage in young individuals.

Structuring a focused discussion is a central communication skill. Identify the health care proxy and recommend his or her involvement in the process of advance care planning. Select a worksheet, preferably one that has been evaluated and demonstrated to produce reliable and valid expressions of patient preferences, and orient the patient and proxy to it. Such worksheets exist for both general and disease-specific situations. Discuss with the patient and proxy one scenario as an example to demonstrate how to think about the issues. It is often helpful to begin with a scenario in which the patient is likely to have settled preferences for care, such as being in a persistent vegetative state. Once the patient's preferences for interventions in this scenario are determined, suggest that the patient and proxy discuss and complete the worksheet for the others. If appropriate, suggest that they involve other family members in the discussion. On a return visit, go over the patient's preferences, checking and resolving any inconsistencies. After having the patient and proxy sign the document, place it in the medical chart and be sure that copies are provided to relevant family members and care sites. Because patients' preferences can change, these documents have to be reviewed periodically.

TYPES OF DOCUMENTS Advance care planning documents are of three broad types. The first includes living wills or instructional directives; these are advisory documents that describe the types of decisions that should direct care. Some are more specific, delineating different scenarios and interventions for the patient to choose from. Among these, some are for general use and others are designed for use by patients with a specific type of disease, such as cancer or HIV. A second type is a less specific directive that provides general statements of not wanting life-sustaining interventions or forms that describe the values that should guide specific discussions about terminal care. These can be problematic because, when critical decisions about specific treatments are needed, they require assessments by people other than the patient of whether a treatment fulfills a particular wish. The third type of advance directive allows the designation of a health care proxy (sometimes also referred to as a durable attorney for health care), who is an individual selected by the patient to make decisions. The choice is not either/or; a combined directive that includes a living will and designates a proxy is often used, and the directive should indicate clearly whether the specified patient preferences or the proxy's choice takes precedence if they conflict. The Five Wishes and the Medical Directive are such combined forms. Some states have begun to put into practice a “Physician Orders for Life-Sustaining Treatment (POLST)” paradigm, which builds on communication between providers and patients to include guidance for end-of-life care in a color-coordinated form that follows the patient across treatment settings. The procedures for completing advance care planning documents vary according to state law.

A potentially misleading distinction relates to statutory as opposed to advisory documents. Statutory documents are drafted to fulfill relevant state laws. Advisory documents are drafted to reflect the patient's wishes. Both are legal, the first under state law and the latter under common or constitutional law.

LEGAL ASPECTS The U.S. Supreme Court has ruled that patients have a constitutional right to decide about refusing and terminating medical interventions, including life-sustaining interventions, and that mentally incompetent patients can exercise this right by providing “clear and convincing evidence” of their preferences. Because advance care directives permit patients to provide such evidence, commentators agree that

TABLE 10-3 STEPS IN ADVANCE CARE PLANNING

Step	Goals to be Achieved and Measures to Cover	Useful Phrases or Points to Make
Introducing advance care planning	Ask the patient what he or she knows about advance care planning and if he or she has already completed an advance care directive.	<i>I'd like to talk with you about something I try to discuss with all my patients. It's called advance care planning. In fact, I feel that this is such an important topic that I have done this myself. Are you familiar with advance care planning or living wills?</i>
	Indicate that you as a physician have completed advance care planning.	<i>Have you thought about the type of care you would want if you ever became too sick to speak for yourself? That is the purpose of advance care planning.</i>
	Indicate that you try to perform advance care planning with all patients regardless of prognosis.	<i>There is no change in health that we have not discussed. I am bringing this up now because it is sensible for everyone, no matter how well or ill, old or young.</i>
	Explain the goals of the process as empowering the patient and ensuring that you and the proxy understand the patient's preferences.	Have many copies of advance care directives available, including in the waiting room, for patients and families.
	Provide the patient relevant literature, including the advance care directive that you prefer to use.	Know resources for state-specific forms (available at www.nhpco.org).
Structured discussion of scenarios and patient	Recommend the patient identify a proxy decision-maker who should attend the next meeting.	
	Affirm that the goal of the process is to follow the patient's wishes if the patient loses decision-making capacity.	Use a structured worksheet with typical scenarios.
	Elicit the patient's overall goals related to health care.	Begin the discussion with persistent vegetative state and consider other scenarios, such as recovery from an acute event with serious disability, asking the patient about his or her preferences regarding specific interventions, such as ventilators, artificial nutrition, and CPR, and then proceeding to less invasive interventions, such as blood transfusions and antibiotics.
	Elicit the patient's preferences for specific interventions in a few salient and common scenarios.	
Review the patient's preferences	Help the patient define the threshold for withdrawing and withholding interventions.	
	Define the patient's preference for the role of the proxy.	
Document the patient's preferences	After the patient has made choices of interventions, review them to ensure they are consistent and the proxy is aware of them.	
Update the directive	Formally complete the advance care directive and have a witness sign it.	
	Provide a copy for the patient and the proxy.	
Apply the directive	Insert a copy into the patient's medical record and summarize in a progress note.	
	Periodically, and with major changes in health status, review the directive with the patient and make any modifications.	
	The directive goes into effect only when the patient becomes unable to make medical decisions for himself or herself.	
Apply the directive	Reread the directive to be sure about its content.	
	Discuss your proposed actions based on the directive with the proxy.	

Abbreviation: CPR, cardiopulmonary resuscitation.

they are constitutionally protected. Most commentators believe that a state is required to honor any clear advance care directive whether or not it is written on an "official" form. Many states have enacted laws explicitly to honor out-of-state directives. If a patient is not using a statutory form, it may be advisable to attach a statutory form to the advance care directive being used. State-specific forms are readily available free of charge for health care providers and patients and families through the National Hospice and Palliative Care Organization's "Caring Connections" website (<http://www.caringinfo.org>).

In January 2014, Texas judge R. H. Wallace ruled that a brain dead woman who was 23 weeks pregnant should be removed from life support. This was after several months of disagreement between the woman's family and the hospital providing care. The hospital cited Texas law that states that life-sustaining treatment must be administered to a pregnant woman, but the judge sided with the woman's family saying that the law did not apply because the patient was legally dead.

As of 2013, advance directives are legal in all states and the District of Columbia either through state specific legislation, state judicial rulings, or United States Supreme Court rulings. Many states have their own statutory forms. Massachusetts and Michigan do not have living will laws, although both have health care proxy laws. In 27 states, the laws state that the living will is not valid if a woman is pregnant. However, like all other states except Alaska, these states have enacted durable power of attorney for health care laws that permit patients to designate a proxy decision-maker with authority to terminate life-sustaining treatments. Only in Alaska does the law prohibit proxies from terminating life-sustaining treatments. The health reform

legislation, the Affordable Care Act of 2010, raised substantial controversy when early versions of the law included Medicare reimbursement for advance care planning consultations. These provisions were withdrawn over accusations that they would lead to the rationing of care for the elderly.

INTERVENTIONS

PHYSICAL SYMPTOMS AND THEIR MANAGEMENT

Great emphasis has been placed on addressing dying patients' pain. Some institutions have made pain assessment a fifth vital sign to emphasize its importance. This also has been advocated by large health care systems such as the Veterans' Administration and accrediting bodies such as the Joint Commission. Although this embrace of pain as the fifth vital sign has been symbolically important, no data document that it has improved pain management practices. Although good palliative care requires good pain management, it also requires more. The frequency of symptoms varies by disease and other factors. The most common physical and psychological symptoms among all terminally ill patients include pain, fatigue, insomnia, anorexia, dyspnea, depression, anxiety, and nausea and vomiting. In the last days of life, terminal delirium is also common. Assessments of patients with advanced cancer have shown that patients experienced an average of 11.5 different physical and psychological symptoms (**Table 10-4**).

Evaluations to determine the etiology of these symptoms usually can be limited to the history and physical examination. In some cases, radiologic or other diagnostic examinations will provide sufficient

TABLE 10-4 COMMON PHYSICAL AND PSYCHOLOGICAL SYMPTOMS OF TERMINALLY ILL PATIENTS

Physical Symptoms	Psychological Symptoms
Pain	Anxiety
Fatigue and weakness	Depression
Dyspnea	Hopelessness
Insomnia	Meaninglessness
Dry mouth	Irritability
Anorexia	Impaired concentration
Nausea and vomiting	Confusion
Constipation	Delirium
Cough	Loss of libido
Swelling of arms or legs	
Itching	
Diarrhea	
Dysphagia	
Dizziness	
Fecal and urinary incontinence	
Numbness/tingling in hands/feet	

benefit in directing optimal palliative care to warrant the risks, potential discomfort, and inconvenience, especially to a seriously ill patient. Only a few of the common symptoms that present difficult management issues will be addressed in this chapter. **Additional information on the management of other symptoms, such as nausea and vomiting, insomnia, and diarrhea, can be found in Chaps. 54 and 99, Chap. 38, and Chap. 55, respectively.**

Pain • FREQUENCY The frequency of pain among terminally ill patients varies widely. Substantial pain occurs in 36–90% of patients with advanced cancer. In the SUPPORT study of hospitalized patients with diverse conditions and an estimated survival ≤ 6 months, 22% reported moderate to severe pain, and caregivers of those patients noted that 50% had similar levels of pain during the last few days of life. A meta-analysis found pain prevalence of 58–69% in studies that included patients characterized as having advanced, metastatic, or terminal cancer; 44–73% in studies that included patients characterized as undergoing cancer treatment; and 21–46% in studies that included posttreatment individuals.

ETIOLOGY *Nociceptive pain* is the result of direct mechanical or chemical stimulation of nociceptors and normal neural signaling to the brain. It tends to be localized, aching, throbbing, and cramping. The classic example is bone metastases. *Visceral pain* is caused by nociceptors in gastrointestinal, respiratory, and other organ systems. It is a deep or colicky type of pain classically associated with pancreatitis, myocardial infarction, or tumor invasion of viscera. *Neuropathic pain* arises from disordered nerve signals. It is described by patients as burning, electrical, or shocklike pain. Classic examples are poststroke pain, tumor invasion of the brachial plexus, and herpetic neuralgia.

ASSESSMENT Pain is a subjective experience. Depending on the patient's circumstances, perspective, and physiologic condition, the same physical lesion or disease state can produce different levels of reported pain and need for pain relief. Systematic assessment includes eliciting the following: (1) type: throbbing, cramping, burning, etc.; (2) periodicity: continuous, with or without exacerbations, or incident; (3) location; (4) intensity; (5) modifying factors; (6) effects of treatments; (7) functional impact; and (8) impact on patient. Several validated pain assessment measures may be used, such as the Visual Analogue Scale, the Brief Pain Inventory, and the pain component of one of the more comprehensive symptom assessment instruments. Frequent reassessments are essential to assess the effects of interventions.

INTERVENTIONS Interventions for pain must be tailored to each individual, with the goal of preempting chronic pain and relieving breakthrough pain. At the end of life, there is rarely reason to doubt a

patient's report of pain. Pain medications are the cornerstone of management. If they are failing and nonpharmacologic interventions—including radiotherapy and anesthetic or neurosurgical procedures such as peripheral nerve blocks or epidural medications—are required, a pain consultation is appropriate.

Pharmacologic interventions follow the World Health Organization three-step approach involving nonopioid analgesics, mild opioids, and strong opioids, with or without adjuvants (Chap. 18). Nonopioid analgesics, especially nonsteroidal anti-inflammatory drugs (NSAIDs), are the initial treatments for mild pain. They work primarily by inhibiting peripheral prostaglandins and reducing inflammation but also may have central nervous system (CNS) effects. They have a ceiling effect. Ibuprofen, up to a total dose of 1600 mg/d given in four doses of 400 mg each, has a minimal risk of causing bleeding and renal impairment and is a good initial choice. In patients with a history of severe gastrointestinal (GI) or other bleeding, it should be avoided. In patients with a history of mild gastritis or gastroesophageal reflux disease (GERD), acid-lowering therapy such as a proton pump inhibitor should be used. Acetaminophen is an alternative in patients with a history of GI bleeding and can be used safely at up to 4 g/d given in four doses of 1 g each. In patients with liver dysfunction due to metastases or other causes and in patients with heavy alcohol use, doses should be reduced.

If nonopioid analgesics are insufficient, opioids should be introduced. They work by interacting with μ opioid receptors in the CNS to activate pain-inhibitory neurons; most are receptor antagonists. The mixed agonist/antagonist opioids useful for postacute pain should not be used for the chronic pain in end-of-life care. Weak opioids such as codeine can be used initially. However, if they are escalated and fail to relieve pain, strong opioids such as morphine, 5–10 mg every 4 h, should be used. Nonopioid analgesics should be combined with opioids because they potentiate the effect of opioids.

For continuous pain, opioids should be administered on a regular, around-the-clock basis consistent with their duration of analgesia. They should not be provided only when the patient experiences pain; the goal is to prevent patients from experiencing pain. Patients also should be provided rescue medication, such as liquid morphine, for breakthrough pain, generally at 20% of the baseline dose. Patients should be informed that using the rescue medication does not obviate the need to take the next standard dose of pain medication. If after 24 h the patient's pain remains uncontrolled and recurs before the next dose, requiring the patient to use the rescue medication, the daily opioid dose can be increased by the total dose of rescue medications used by the patient, or by 50% for moderate pain and 100% for severe pain of the standing opioid daily dose.

It is inappropriate to start with extended-release preparations. Instead, an initial focus on using short-acting preparations to determine how much is required in the first 24–48 h will allow clinicians to determine opioid needs. Once pain relief is obtained with short-acting preparations, one should switch to extended-release preparations. Even with a stable extended-release preparation regimen, the patient may have incident pain, such as during movement or dressing changes. Short-acting preparations should be taken before such predictable episodes. Although less common, patients may have “end-of-dose failure” with long-acting opioids, meaning that they develop pain after 8 h in the case of an every-12-h medication. In these cases, a trial of giving an every-12-h medication every 8 h is appropriate.

Because of differences in opioid receptors, cross-tolerance among opioids is incomplete, and patients may experience different side effects with different opioids. Therefore, if a patient is not experiencing pain relief or is experiencing too many side effects, a change to another opioid preparation is appropriate. When switching, one should begin with 50–75% of the published equianalgesic dose of the new opioid.

Unlike NSAIDs, opioids have no ceiling effect; therefore, there is no maximum dose no matter how many milligrams the patient is receiving. The appropriate dose is the dose needed to achieve pain relief. This is an important point for clinicians to explain to patients and families. Addiction or excessive respiratory depression is extremely unlikely in the terminally ill; fear of these side effects should neither prevent escalating opioid medications when the patient is experiencing insufficient pain relief nor justify using opioid antagonists.

Opioid side effects should be anticipated and treated preemptively. Nearly all patients experience constipation that can be debilitating (see below). Failure to prevent constipation often results in noncompliance with opioid therapy. Methylnaltrexone is a drug that targets opioid-induced constipation by blocking peripheral opioid receptors but not central receptors for analgesia. In placebo-controlled trials, it has been shown to cause laxation within 24 h of administration. As with the use of opioids, about a third of patients using methylnaltrexone experience nausea and vomiting, but unlike constipation, tolerance develops, usually within a week. Therefore, when one is beginning opioids, an antiemetic such as metoclopramide or a serotonin antagonist often is prescribed prophylactically and stopped after 1 week. Olanzapine also has anti-nausea properties and can be effective in countering delirium or anxiety, with the advantage of some weight gain.

Drowsiness, a common side effect of opioids, also usually abates within a week. During this period, drowsiness can be treated with psychostimulants such as dextroamphetamine, methylphenidate, and modafinil. Modafinil has the advantage of everyday dosing. Pilot reports suggest that donepezil may also be helpful for opiate-induced drowsiness as well as relieving fatigue and anxiety. Metabolites of morphine and most opioids are cleared renally; doses may have to be adjusted for patients with renal failure.

Seriously ill patients who require chronic pain relief rarely if ever become addicted. Suspicion of addiction should not be a reason to withhold pain medications from terminally ill patients. Patients and families may withhold prescribed opioids for fear of addiction or dependence. Physicians and health care providers should reassure patients and families that the patient will not become addicted to opioids if they are used as prescribed for pain relief; this fear should not prevent the patient from taking the medications around the clock. However, diversion of drugs for use by other family members or illicit sale may occur. It may be necessary to advise the patient and caregiver about secure storage of opioids. Contract writing with the patient and family can help. If that fails, transfer to a safe facility may be necessary.

Tolerance is the need to increase medication dosage for the same pain relief without a change in disease. In the case of patients with advanced disease, the need for increasing opioid dosage for pain relief usually is caused by disease progression rather than tolerance. Physical dependence is indicated by symptoms from the abrupt withdrawal of opioids and should not be confused with addiction.

Adjuvant analgesic medications are nonopioids that potentiate the analgesic effects of opioids. They are especially important in the management of neuropathic pain. Gabapentin and pregabalin, calcium channel alpha 2-delta ligands, are now the first-line treatments for neuropathic pain from a variety of causes. Gabapentin is begun at 100–300 mg bid or tid, with 50–100% dose increments every 3 days. Usually 900–3600 mg/d in two or three doses is effective. The combination of gabapentin and nortriptyline may be more effective than gabapentin alone. One potential side effect of gabapentin to be aware of is confusion and drowsiness, especially in the elderly. Pregabalin has the same mechanism of action as gabapentin but is absorbed more efficiently from the GI tract. It is started at 75 mg bid and increased to 150 mg bid. The maximum dose is 225 mg bid. Carbamazepine, a first-generation agent, has been proved effective in randomized trials for neuropathic pain. Other potentially effective anticonvulsant adjuvants include topiramate (25–50 mg qd or bid, rising to 100–300 mg/d) and oxcarbazepine (75–300 mg bid, rising to 1200 mg bid). Glucocorticoids, preferably dexamethasone given once a day, can be useful in reducing inflammation that causes pain while elevating mood, energy, and appetite. Its main side effects include confusion, sleep difficulties, and fluid retention. Glucocorticoids are especially effective for bone pain and abdominal pain from distention of the GI tract or liver. Other drugs, including clonidine and baclofen, can be effective in pain relief. These drugs are adjuvants and generally should be used in conjunction with—not instead of—opioids. Methadone, carefully dosed because of its unpredictable half-life in many patients, has activity at the *N*-methyl-D-aspartate (NMDA) receptor and is useful for complex pain syndromes and neuropathic pain. It generally is reserved for cases in which first-line opioids (morphine, oxycodone, hydromorphone) are either ineffective or unavailable.

Radiation therapy can treat bone pain from single metastatic lesions. Bone pain from multiple metastases can be amenable to radiopharmaceuticals such as strontium-89 and samarium-153. Bisphosphonates (such as pamidronate [90 mg every 4 weeks]) and calcitonin (200 IU intranasally once or twice a day) also provide relief from bone pain but have an onset of action of days.

Constipation • FREQUENCY Constipation is reported in up to 87% of patients requiring palliative care.

ETIOLOGY Although hypercalcemia and other factors can cause constipation, it is most frequently a predictable consequence of the use of opioids for the relief of pain and dyspnea and of tricyclic antidepressants, from their anticholinergic effects, and of the inactivity and poor diet that are common among seriously ill patients. If untreated, constipation can cause substantial pain and vomiting and also is associated with confusion and delirium. Whenever opioids and other medications known to cause constipation are used, preemptive treatment for constipation should be instituted.

ASSESSMENT The physician should establish the patient's previous bowel habits, including the frequency, consistency, and volume. Abdominal and rectal examinations should be performed to exclude impaction or acute abdomen. A number of constipation assessment scales are available, although guidelines issued in the *Journal of Palliative Medicine* did not recommend them for routine practice. Radiographic assessments beyond a simple flat plate of the abdomen in cases in which obstruction is suspected are rarely necessary.

INTERVENTION Intervention to reestablish comfortable bowel habits and relieve pain and discomfort should be the goals of any measures to address constipation during end-of-life care. Although physical activity, adequate hydration, and dietary treatments with fiber can be helpful, each is limited in its effectiveness for most seriously ill patients, and fiber may exacerbate problems in the setting of dehydration and if impaired motility is the etiology. Fiber is contraindicated in the presence of opioid use. Stimulant and osmotic laxatives, stool softeners, fluids, and enemas are the mainstays of therapy (Table 10-5). In preventing constipation from opioids and other medications, a combination of a laxative and a stool softener (such as senna and docusate) should be used. If after several days of treatment, a bowel movement has not occurred, a rectal examination to remove impacted stool and place a suppository is necessary. For patients with impending bowel obstruction or gastric stasis, octreotide to reduce secretions can be helpful. For patients in whom the suspected mechanism is dysmotility, metoclopramide can be helpful.

Nausea • FREQUENCY Up to 70% of patients with advanced cancer have nausea, defined as the subjective sensation of wanting to vomit.

ETIOLOGY Nausea and vomiting are both caused by stimulation at one of four sites: the GI tract, the vestibular system, the chemoreceptor trigger zone (CTZ), and the cerebral cortex. Medical treatments for nausea are aimed at receptors at each of these sites: the GI tract contains mechanoreceptors, chemoreceptors, and 5-hydroxytryptamine type 3 (5-HT₃) receptors; the vestibular system probably contains histamine and acetylcholine receptors; and the CTZ contains chemoreceptors, dopamine type 2 receptors, and 5-HT₃ receptors. An example of nausea that most likely is mediated by the cortex is anticipatory nausea before a dose of chemotherapy or other noxious stimuli.

Specific causes of nausea include metabolic changes (liver failure, uremia from renal failure, hypercalcemia), bowel obstruction, constipation, infection, GERD, vestibular disease, brain metastases, medications (including antibiotics, NSAIDs, proton pump inhibitors, opioids, and chemotherapy), and radiation therapy. Anxiety can also contribute to nausea.

INTERVENTION Medical treatment of nausea is directed at the anatomic and receptor-mediated cause that a careful history and physical examination reveals. When a single specific cause is not found, many advocate beginning treatment with a dopamine antagonist such as haloperidol or prochlorperazine. Prochlorperazine is usually more

TABLE 10-5 MEDICATIONS FOR THE MANAGEMENT OF CONSTIPATION

Intervention	Dose	Comment
Stimulant laxatives		These agents directly stimulate peristalsis and may reduce colonic absorption of water.
Prune juice	120–240 mL/d	Work in 6–12 h.
Senna (Senokot)	2–8 tablets PO bid	
Bisacodyl	5–15 mg/d PO, PR	
Osmotic laxatives		These agents are not absorbed. They attract and retain water in the gastrointestinal tract.
Lactulose	15–30 mL PO q4–8h	Lactulose may cause flatulence and bloating.
Magnesium hydroxide (Milk of Magnesia)	15–30 mL/d PO	Lactulose works in 1 day, magnesium products in 6 h.
Magnesium citrate	125–250 mL/d PO	
Stool softeners		These agents work by increasing water secretion and as detergents, increasing water penetration into the stool.
Sodium docusate (Colace)	300–600 mg/d PO	Work in 1–3 days.
Calcium docusate	300–600 mg/d PO	
Suppositories and enemas		
Bisacodyl	10–15 PR qd	
Sodium phosphate enema	PR qd	Fixed dose, 4.5 oz, Fleet's.

sedating than haloperidol. When decreased motility is suspected, metoclopramide can be an effective treatment. When inflammation of the GI tract is suspected, glucocorticoids such as dexamethasone are an appropriate treatment. For nausea that follows chemotherapy and radiation therapy, one of the 5-HT₃ receptor antagonists (ondansetron, granisetron, dolasetron, palonosetron) is recommended. Studies suggest palonosetron has higher receptor binding affinity and clinical superiority to the other 5-HT₃ receptor antagonists. Clinicians should attempt prevention of postchemotherapy nausea rather than provide treatment after the fact. Current clinical guidelines recommend tailoring the strength of treatments to the specific emetic risk posed by a specific chemotherapy drug. When a vestibular cause (such as “motion sickness” or labyrinthitis) is suspected, antihistamines such as meclizine (whose primary side effect is drowsiness) or anticholinergics such as scopolamine can be effective. In anticipatory nausea, a benzodiazepine such as lorazepam is indicated. As with antihistamines, drowsiness and confusion are the main side effects.

Dyspnea • FREQUENCY Dyspnea is a subjective experience of being short of breath. Frequencies vary among causes of death, but it can affect 80–90% of dying patients with lung cancer, COPD, and heart disease. Dyspnea is among the most distressing physical symptoms and can be even more distressing than pain.

ASSESSMENT As with pain, dyspnea is a subjective experience that may not correlate with objective measures of PO₂, PCO₂, or respiratory rate. Consequently, measurements of oxygen saturation through pulse oximetry or blood gases are rarely helpful in guiding therapy. Despite the limitations of existing assessment methods, physicians should regularly assess and document patients' experience of dyspnea and its intensity. Guidelines recommend visual or analogue dyspnea scales to assess the severity of symptoms and the effects of treatment. Potentially reversible or treatable causes of dyspnea include infection, pleural effusions, pulmonary emboli, pulmonary edema, asthma, and tumor encroachment on the airway. However, the risk-versus-benefit ratio of the diagnostic and therapeutic interventions for patients with little time left to live must be considered carefully before one undertakes

TABLE 10-6 MEDICATIONS FOR THE MANAGEMENT OF DYSPNEA

Intervention	Dose	Comments
Weak opioids		For patients with mild dyspnea
Codeine (or codeine with 325 mg acetaminophen)	30 mg PO q4h	For opioid-naïve patients
Hydrocodone	5 mg PO q4h	
Strong opioids		For opioid-naïve patients with moderate to severe dyspnea
Morphine	5–10 mg PO q4h 30–50% of baseline opioid dose q4h	For patients already taking opioids for pain or other symptoms
Oxycodone	5–10 mg PO q4h	
Hydromorphone	1–2 mg PO q4h	
Anxiolytics		Give a dose every hour until the patient is relaxed, then provide a dose for maintenance
Lorazepam	0.5–2.0 mg PO/SL/IV qh then q4–6h	
Clonazepam	0.25–2.0 mg PO q12h	
Midazolam	0.5 mg IV q15min	

diagnostic steps. Frequently, the specific etiology cannot be identified, and dyspnea is the consequence of progression of the underlying disease that cannot be treated. The anxiety caused by dyspnea and the choking sensation can significantly exacerbate the underlying dyspnea in a negatively reinforcing cycle.

INTERVENTIONS When reversible or treatable etiologies are diagnosed, they should be treated as long as the side effects of treatment, such as repeated drainage of effusions or anticoagulants, are less burdensome than the dyspnea itself. More aggressive treatments such as stenting a bronchial lesion may be warranted if it is clear that the dyspnea is due to tumor invasion at that site and if the patient and family understand the risks of such a procedure. Usually, treatment will be symptomatic (Table 10-6). A dyspnea scale and careful monitoring should guide dose adjustment. Low-dose opioids reduce the sensitivity of the central respiratory center and the sensation of dyspnea. If patients are not receiving opioids, weak opioids can be initiated; if patients are already receiving opioids, morphine or other strong opioids should be used. Controlled trials do not support the use of nebulized opioids for dyspnea at the end of life. Phenothiazines and chlorpromazine may be helpful when combined with opioids. Benzodiazepines can be helpful if anxiety is present but should be neither used as first-line therapy nor used alone in the treatment of dyspnea. If the patient has a history of COPD or asthma, inhaled bronchodilators and glucocorticoids may be helpful. If the patient has pulmonary edema due to heart failure, diuresis with a medication such as furosemide is indicated. Excess secretions can be dried with scopolamine, transdermally or intravenously. Use of oxygen is controversial. There are conflicting data on its effectiveness for patients with proven hypoxemia. But there is no clear benefit of oxygen compared to room air for nonhypoxemic patients. Noninvasive positive-pressure ventilation using a facemask or nasal plugs may be used for some patients for symptom relief. For some families and patients, oxygen is distressing; for others, it is reassuring. More general interventions that medical staff can do include sitting the patient upright, removing smoke or other irritants such as perfume, ensuring a supply of fresh air with sufficient humidity, and minimizing other factors that can increase anxiety.

Fatigue • FREQUENCY More than 90% of terminally ill patients experience fatigue and/or weakness. Fatigue is one of the most commonly reported symptoms of cancer treatment as well as in the palliative care of multiple sclerosis, COPD, heart failure, and HIV. Fatigue frequently is cited as among the most distressing symptoms.

ETIOLOGY The multiple causes of fatigue in the terminally ill can be categorized as resulting from the underlying disease; from disease-induced factors such as tumor necrosis factor and other cytokines; and from secondary factors such as dehydration, anemia, infection, hypothyroidism, and drug side effects. Apart from low caloric intake, loss of muscle mass and changes in muscle enzymes may play an important role in fatigue of terminal illness. The importance of changes in the CNS, especially the reticular activating system, have been hypothesized based on reports of fatigue in patients receiving cranial radiation, experiencing depression, or having chronic pain in the absence of cachexia or other physiologic changes. Finally, depression and other causes of psychological distress can contribute to fatigue.

ASSESSMENT Like pain and dyspnea, fatigue is subjective. Objective changes, even in body mass, may be absent. Consequently, assessment must rely on patient self-reporting. Scales used to measure fatigue, such as the Edmonton Functional Assessment Tool, the Fatigue Self-Report Scales, and the Rhoten Fatigue Scale, are usually appropriate for research rather than clinical purposes. In clinical practice, a simple performance assessment such as the Karnofsky Performance Status or the ECOG's question "How much of the day does the patient spend in bed?" may be the best measure. In this 0–4 performance status assessment, 0 = normal activity; 1 = symptomatic without being bedridden; 2 = requiring some, but <50%, bed time; 3 = bedbound more than half the day; and 4 = bedbound all the time. Such a scale allows for assessment over time and correlates with overall disease severity and prognosis. A 2008 review by the European Association of Palliative Care also described several longer assessment tools with 9–20 items, including the Piper Fatigue Inventory, the Multidimensional Fatigue Inventory, and the Brief Fatigue Inventory (BFI).

INTERVENTIONS For some patients, there are reversible causes such as anemia, but for most patients at the end of life, fatigue will not be "cured." The goal is to ameliorate it and help patients and families adjust expectations. Behavioral interventions should be used to avoid blaming the patient for inactivity and to educate both the family and the patient that the underlying disease causes physiologic changes that produce low energy levels. Understanding that the problem is physiologic and not psychological can help alter expectations regarding the patient's level of physical activity. Practically, this may mean reducing routine activities such as housework and cooking or social events outside the house and making it acceptable to receive guests lying on a couch. At the same time, institution of exercise regimens and physical therapy can raise endorphins, reduce muscle wasting, and reduce the risk of depression. In addition, ensuring good hydration without worsening edema may help reduce fatigue. Discontinuing medications that worsen fatigue may help, including cardiac medications, benzodiazepines, certain antidepressants, or opioids if pain is well-controlled. As end-of-life care proceeds into its final stages, fatigue may protect patients from further suffering, and continued treatment could be detrimental.

There are woefully few pharmacologic interventions that target fatigue and weakness. Glucocorticoids can increase energy and enhance mood. Dexamethasone is preferred for its once-a-day dosing and minimal mineralocorticoid activity. Benefit, if any, usually is seen within the first month. Psychostimulants such as dextroamphetamine (5–10 mg PO) and methylphenidate (2.5–5 mg PO) may also enhance energy levels, although a randomized trial did not show methylphenidate beneficial compared with placebo in cancer fatigue. Doses should be given in the morning and at noon to minimize the risk of counterproductive insomnia. Modafinil, developed for narcolepsy, has shown some promise in the treatment of severe fatigue and has the advantage of once-daily dosing. Its precise role in fatigue at the end of life has not been determined. Anecdotal evidence suggests that L-carnitine may improve fatigue, depression, and sleep disruption. Similarly, some studies suggest ginseng can reduce fatigue.

PSYCHOLOGICAL SYMPTOMS AND THEIR MANAGEMENT

Depression • FREQUENCY Depression at the end of life presents an apparently paradoxical situation. Many people believe that depression

is normal among seriously ill patients because they are dying. People frequently say, "Wouldn't you be depressed?" However, depression is not a necessary part of terminal illness and can contribute to needless suffering. Although sadness, anxiety, anger, and irritability are normal responses to a serious condition, they are typically of modest intensity and transient. Persistent sadness and anxiety and the physically disabling symptoms that they can lead to are abnormal and suggestive of major depression. Although as many as 75% of terminally ill patients experience emotional distress and depressive symptoms, <30% of terminally ill patients have major depression. Depression is not limited to cancer patients but found in patients with end-stage renal disease, Parkinson's disease, multiple sclerosis, and other terminal conditions.

ETIOLOGY Previous history of depression, family history of depression or bipolar disorder, and prior suicide attempts are associated with increased risk for depression among terminally ill patients. Other symptoms, such as pain and fatigue, are associated with higher rates of depression; uncontrolled pain can exacerbate depression, and depression can cause patients to be more distressed by pain. Many medications used in the terminal stages, including glucocorticoids, and some anticancer agents, such as tamoxifen, interleukin 2, interferon α , and vincristine, also are associated with depression. Some terminal conditions, such as pancreatic cancer, certain strokes, and heart failure, have been reported to be associated with higher rates of depression, although this is controversial. Finally, depression may be attributable to grief over the loss of a role or function, social isolation, or loneliness.

ASSESSMENT Diagnosing depression among seriously ill patients is complicated because many of the vegetative symptoms in the DSM-V (*Diagnostic and Statistical Manual of Mental Disorders*) criteria for clinical depression—insomnia, anorexia and weight loss, fatigue, decreased libido, and difficulty concentrating—are associated with the dying process itself. The assessment of depression in seriously ill patients therefore should focus on the dysphoric mood, helplessness, hopelessness, and lack of interest and enjoyment and concentration in normal activities. The single questions "How often do you feel downhearted and blue?" (more than a good bit of the time or similar responses) and "Do you feel depressed most of the time?" are appropriate for screening. Visual Analog Scales can also be useful in screening.

INTERVENTIONS Physicians must treat any physical symptom, such as pain, that may be causing or exacerbating depression. Fostering adaptation to the many losses that the patient is experiencing can also be helpful. Nonpharmacologic interventions, including group or individual psychological counseling, and behavioral therapies such as relaxation and imagery can be helpful, especially in combination with drug therapy.

Pharmacologic interventions remain the core of therapy. The same medications are used to treat depression in terminally ill as in nonterminally ill patients. Psychostimulants may be preferred for patients with a poor prognosis or for those with fatigue or opioid-induced somnolence. Psychostimulants are comparatively fast acting, working within a few days instead of the weeks required for selective serotonin reuptake inhibitors (SSRIs). Dextroamphetamine or methylphenidate should be started at 2.5–5.0 mg in the morning and at noon, the same starting doses used for treating fatigue. The dose can be escalated up to 15 mg bid. Modafinil is started at 100 mg qd and can be increased to 200 mg if there is no effect at the lower dose. Pemoline is a nonamphetamine psychostimulant with minimal abuse potential. It is also effective as an antidepressant beginning at 18.75 mg in the morning and at noon. Because it can be absorbed through the buccal mucosa, it is preferred for patients with intestinal obstruction or dysphagia. If it is used for prolonged periods, liver function must be monitored. The psychostimulants can also be combined with more traditional antidepressants while waiting for the antidepressants to become effective and then tapered after a few weeks if necessary. Psychostimulants have side effects, particularly initial anxiety, insomnia, and rarely paranoia, which may necessitate lowering the dose or discontinuing treatment.

Mirtazapine, an antagonist at the postsynaptic serotonin receptors, is a promising psychostimulant. It should be started at 7.5 mg before

bed. It has sedating, antiemetic, and anxiolytic properties with few drug interactions. Its side effect of weight gain may be beneficial for seriously ill patients; it is available in orally disintegrating tablets.

For patients with a prognosis of several months or longer, SSRIs, including fluoxetine, sertraline, paroxetine, citalopram, escitalopram, and fluvoxamine, and serotonin-noradrenaline reuptake inhibitors such as venlafaxine, are the preferred treatment because of their efficacy and comparatively few side effects. Because low doses of these medications may be effective for seriously ill patients, one should use half the usual starting dose for healthy adults. The starting dose for fluoxetine is 10 mg once a day. In most cases, once-a-day dosing is possible. The choice of which SSRI to use should be driven by (1) the patient's past success or failure with the specific medication, (2) the most favorable side effect profile for that specific agent, and (3) the time it takes to reach steady-state drug levels. For instance, for a patient in whom fatigue is a major symptom, a more activating SSRI (fluoxetine) would be appropriate. For a patient in whom anxiety and sleeplessness are major symptoms, a more sedating SSRI (paroxetine) would be appropriate.

Atypical antidepressants are recommended only in selected circumstances, usually with the assistance of a specialty consultation. Trazodone can be an effective antidepressant but is sedating and can cause orthostatic hypotension and, rarely, priapism. Therefore, it should be used only when a sedating effect is desired and is often used for patients with insomnia, at a dose starting at 25 mg. In addition to its antidepressant effects, bupropion is energizing, making it useful for depressed patients who experience fatigue. However, it can cause seizures, preventing its use for patients with a risk of CNS neoplasms or terminal delirium. Finally, alprazolam, a benzodiazepine, starting at 0.25–1.0 mg tid, can be effective in seriously ill patients who have a combination of anxiety and depression. Although it is potent and works quickly, it has many drug interactions and may cause delirium, especially among very ill patients, because of its strong binding to the benzodiazepine- γ -aminobutyric acid (GABA) receptor complex.

Unless used as adjuvants for the treatment of pain, tricyclic antidepressants are not recommended. Similarly, monoamine oxidase (MAO) inhibitors are not recommended because of their side effects and dangerous drug interactions.

Delirium (See Chap. 34) • FREQUENCY In the weeks or months before death, delirium is uncommon, although it may be significantly underdiagnosed. However, delirium becomes relatively common in the hours and days immediately before death. Up to 85% of patients dying from cancer may experience terminal delirium.

ETIOLOGY Delirium is a global cerebral dysfunction characterized by alterations in cognition and consciousness. It frequently is preceded by anxiety, changes in sleep patterns (especially reversal of day and night), and decreased attention. In contrast to dementia, delirium has an acute onset, is characterized by fluctuating consciousness and inattention, and is reversible, although reversibility may be more theoretical than real for patients near death. Delirium may occur in a patient with dementia; indeed, patients with dementia are more vulnerable to delirium.

Causes of delirium include metabolic encephalopathy arising from liver or renal failure, hypoxemia, or infection; electrolyte imbalances such as hypercalcemia; paraneoplastic syndromes; dehydration; and primary brain tumors, brain metastases, or leptomeningeal spread of tumor. Commonly, among dying patients, delirium can be caused by side effects of treatments, including radiation for brain metastases, and medications, including opioids, glucocorticoids, anticholinergic drugs, antihistamines, antiemetics, benzodiazepines, and chemotherapeutic agents. The etiology may be multifactorial; e.g., dehydration may exacerbate opioid-induced delirium.

ASSESSMENT Delirium should be recognized in any terminally ill patient with new onset of disorientation, impaired cognition, somnolence, fluctuating levels of consciousness, or delusions with or without agitation. Delirium must be distinguished from acute anxiety and depression as well as dementia. The central distinguishing feature is

altered consciousness, which usually is not noted in anxiety, depression, and dementia. Although “hyperactive” delirium characterized by overt confusion and agitation is probably more common, patients also should be assessed for “hypoactive” delirium characterized by sleep-wake reversal and decreased alertness.

In some cases, use of formal assessment tools such as the Mini-Mental Status Examination (which does not distinguish delirium from dementia) and the Delirium Rating Scale (which does distinguish delirium from dementia) may be helpful in distinguishing delirium from other processes. The patient's list of medications must be evaluated carefully. Nonetheless, a reversible etiologic factor for delirium is found in fewer than half of terminally ill patients. Because most terminally ill patients experiencing delirium will be very close to death and may be at home, extensive diagnostic evaluations such as lumbar punctures and neuroradiologic examinations are usually inappropriate.

INTERVENTIONS One of the most important objectives of terminal care is to provide terminally ill patients the lucidity to say goodbye to the people they love. Delirium, especially with agitation during the final days, is distressing to family and caregivers. A strong determinant of bereavement difficulties is witnessing a difficult death. Thus, terminal delirium should be treated aggressively.

At the first sign of delirium, such as day-night reversal with slight changes in mentation, the physician should let the family members know that it is time to be sure that everything they want to say has been said. The family should be informed that delirium is common just before death.

If medications are suspected of being a cause of the delirium, unnecessary agents should be discontinued. Other potentially reversible causes, such as constipation, urinary retention, and metabolic abnormalities, should be treated. Supportive measures aimed at providing a familiar environment should be instituted, including restricting visits only to individuals with whom the patient is familiar and eliminating new experiences; orienting the patient, if possible, by providing a clock and calendar; and gently correcting the patient's hallucinations or cognitive mistakes.

Pharmacologic management focuses on the use of neuroleptics and, in the extreme, anesthetics (Table 10-7). Haloperidol remains first-line therapy. Usually, patients can be controlled with a low dose (1–3 mg/d), usually given every 6 h, although some may require as much as 20 mg/d. It can be administered PO, SC, or IV. IM injections should not be used, except when this is the only way to get a patient under control. Newer atypical neuroleptics, such as olanzapine, risperidone, and quetiapine, have shown significant effectiveness in completely resolving delirium in cancer patients. These drugs also have fewer side effects than haloperidol, along with other beneficial effects for terminally ill patients, including antiemesis, antianxiety, and weight gain. They are useful for patients with longer anticipated life expectancy because they are less likely to cause dysphoria and have a lower risk of dystonic reactions. Also, because they are metabolized through multiple pathways, they can be used in patients with hepatic and renal

TABLE 10-7 MEDICATIONS FOR THE MANAGEMENT OF DELIRIUM

Interventions	Dose
Neuroleptics	
Haloperidol	0.5–5 mg q2–12h, PO/IV/SC/IM
Thioridazine	10–75 mg q4–8h, PO
Chlorpromazine	12.5–50 mg q4–12h, PO/IV/IM
Atypical neuroleptics	
Olanzapine	2.5–5 mg qd or bid, PO
Risperidone	1–3 mg q12h, PO
Quetiapine	50 mg qd, PO
Anxiolytics	
Lorazepam	0.5–2 mg q1–4h, PO/IV/IM
Midazolam	1–5 mg/h continuous infusion, IV/SC
Anesthetics	
Propofol	0.3–2.0 mg/h continuous infusion, IV

dysfunction. Olanzapine has the disadvantage that it is available only orally and that it takes a week to reach steady state. The usual dose is 2.5–5 mg PO bid. Chlorpromazine (10–25 mg every 4–6 h) can be useful if sedation is desired and can be administered IV or PR in addition to PO. Dystonic reactions resulting from dopamine blockade are a side effect of neuroleptics, although they are reported to be rare when these drugs are used to treat terminal delirium. If patients develop dystonic reactions, benztropine should be administered. Neuroleptics may be combined with lorazepam to reduce agitation when the delirium is the result of alcohol or sedative withdrawal.

If no response to first-line therapy is seen, a specialty consultation should be obtained with a change to a different medication. If patients fail to improve after a second neuroleptic, sedation with an anesthetic such as propofol or continuous-infusion midazolam may be necessary. By some estimates, at the very end of life, as many as 25% of patients experiencing delirium, especially restless delirium with myoclonus or convulsions, may require sedation.

Physical restraints should be used with great reluctance and only when the patient's violence is threatening to self or others. If they are used, their appropriateness should be reevaluated frequently.

Insomnia • FREQUENCY Sleep disorders, defined as difficulty initiating sleep or maintaining sleep, sleep difficulty at least 3 nights a week, or sleep difficulty that causes impairment of daytime functioning, occur in 19–63% of patients with advanced cancer. Some 30–74% of patients with other end-stage conditions, including AIDS, heart disease, COPD, and renal disease, experience insomnia.

ETIOLOGY Patients with cancer may have changes in sleep efficiency such as an increase in stage I sleep. Other etiologies of insomnia are coexisting physical illness such as thyroid disease and coexisting psychological illnesses such as depression and anxiety. Medications, including antidepressants, psychostimulants, steroids, and β agonists, are significant contributors to sleep disorders, as are caffeine and alcohol. Multiple over-the-counter medications contain caffeine and antihistamines, which can contribute to sleep disorders.

ASSESSMENT Assessment should include specific questions concerning sleep onset, sleep maintenance, and early-morning waking as these will provide clues to the causative agents and to management. Patients should be asked about previous sleep problems, screened for depression and anxiety, and asked about symptoms of thyroid disease. Caffeine and alcohol are prominent causes of sleep problems, and a careful history of the use of these substances should be obtained. Both excessive use and withdrawal from alcohol can be causes of sleep problems.

INTERVENTIONS The mainstays of intervention include improvement of sleep hygiene (encouragement of regular time for sleep, decreased nighttime distractions, elimination of caffeine and other stimulants and alcohol), intervention to treat anxiety and depression, and treatment for the insomnia itself. For patients with depression who have insomnia and anxiety, a sedating antidepressant such as mirtazapine can be helpful. In the elderly, trazodone, beginning at 25 mg at nighttime, is an effective sleep aid at doses lower than those which cause its antidepressant effect. Zolpidem may have a decreased incidence of delirium in patients compared with traditional benzodiazepines, but this has not been clearly established. When benzodiazepines are prescribed, short-acting ones (such as lorazepam) are favored over longer-acting ones (such as diazepam). Patients who receive these medications should be observed for signs of increased confusion and delirium.

SOCIAL NEEDS AND THEIR MANAGEMENT

Financial Burdens • FREQUENCY Dying can impose substantial economic strains on patients and families, causing distress. In the United States, with one of the least comprehensive health insurance systems among the developed countries, ~20% of terminally ill patients and their families spend >10% of family income on health care costs over and above health insurance premiums. Between 10 and 30% of families

sell assets, use savings, or take out a mortgage to pay for the patient's health care costs. Nearly 40% of terminally ill patients in the United States report that the cost of their illness is a moderate or great economic hardship for their family.

The patient is likely to reduce and eventually stop working. In 20% of cases, a family member of the terminally ill patient also stops working to provide care. The major underlying causes of economic burden are related to poor physical functioning and care needs, such as the need for housekeeping, nursing, and personal care. More debilitated patients and poor patients experience greater economic burdens.

INTERVENTION This economic burden should not be ignored as a private matter. It has been associated with a number of adverse health outcomes, including preferring comfort care over life-prolonging care as well as consideration of euthanasia or physician-assisted suicide. Economic burdens increase the psychological distress of families and caregivers of terminally ill patients, and poverty is associated with many adverse health outcomes. Importantly, recent studies found that "patients with advanced cancer who reported having end-of-life conversations with physicians had significantly lower health care costs in their final week of life. Higher costs were associated with worse quality of death." Assistance from a social worker, early on if possible, to ensure access to all available benefits may be helpful. Many patients, families, and health care providers are unaware of options for long-term care insurance, respite care, the Family Medical Leave Act (FMLA), and other sources of assistance. Some of these options (such as respite care) may be part of a formal hospice program, but others (such as the FMLA) do not require enrollment in a hospice program.

Relationships • FREQUENCY Settling personal issues and closing the narrative of lived relationships are universal needs. When asked if sudden death or death after an illness is preferable, respondents often initially select the former but soon change to the latter as they reflect on the importance of saying goodbye. Bereaved family members who have not had the chance to say goodbye often have a more difficult grief process.

INTERVENTIONS Care of seriously ill patients requires efforts to facilitate the types of encounters and time spent with family and friends that are necessary to meet those needs. Family and close friends may need to be accommodated with unrestricted visiting hours, which may include sleeping near the patient even in otherwise regimented institutional settings. Physicians and other health care providers may be able to facilitate and resolve strained interactions between the patient and other family members. Assistance for patients and family members who are unsure about how to create or help preserve memories, whether by providing materials such as a scrapbook or memory box or by offering them suggestions and informational resources, can be deeply appreciated. Taking photographs and creating videos can be especially helpful to terminally ill patients who have younger children or grandchildren.

Family Caregivers • FREQUENCY Caring for seriously ill patients places a heavy burden on families. Families frequently are required to provide transportation and homemaking as well as other services. Typically, paid professionals such as home health nurses and hospice workers supplement family care; only about a quarter of all caregiving consists of exclusively paid professional assistance. The trend toward more out-of-hospital deaths will increase reliance on families for end-of-life care. Increasingly, family members are being called upon to provide physical care (such as moving and bathing patients) and medical care (such as assessing symptoms and giving medications) in addition to emotional care and support.

Three-quarters of family caregivers of terminally ill patients are women—wives, daughters, sisters, and even daughters-in-law. Because many are widowed, women tend to be able to rely less on family for caregiving assistance and may need more paid assistance. About 20% of terminally ill patients report substantial unmet needs for nursing and personal care. The impact of caregiving on family caregivers is substantial: both bereaved and current caregivers have a higher mortality rate than that of non-caregiving controls.

INTERVENTIONS It is imperative to inquire about unmet needs and to try to ensure that those needs are met either through the family or by paid professional services when possible. Community assistance through houses of worship or other community groups often can be mobilized by telephone calls from the medical team to someone the patient or family identifies. Sources of support specifically for family caregivers should be identified through local sources or nationally through groups such as the National Family Caregivers Association (www.nfcares.org), the American Cancer Society (www.cancer.org), and the Alzheimer's Association (www.alz.org).

EXISTENTIAL NEEDS AND THEIR MANAGEMENT

Frequency Religion and spirituality are often important to dying patients. Nearly 70% of patients report becoming more religious or spiritual when they became terminally ill, and many find comfort in religious or spiritual practices such as prayer. However, ~20% of terminally ill patients become less religious, frequently feeling cheated or betrayed by becoming terminally ill. For other patients, the need is for existential meaning and purpose that is distinct from and may even be antithetical to religion or spirituality. When asked, patients and family caregivers frequently report wanting their professional caregivers to be more attentive to religion and spirituality.

ASSESSMENT Health care providers are often hesitant about involving themselves in the religious, spiritual, and existential experiences of their patients because it may seem private or not relevant to the current illness. But physicians and other members of the care team should be able at least to detect spiritual and existential needs. Screening questions have been developed for a physician's spiritual history taking. Spiritual distress can amplify other types of suffering and even masquerade as intractable physical pain, anxiety, or depression. The screening questions in the comprehensive assessment are usually sufficient. Deeper evaluation and intervention are rarely appropriate for the physician unless no other member of a care team is available or suitable. Pastoral care providers may be helpful, whether from the medical institution or from the patient's own community.

INTERVENTIONS Precisely how religious practices, spirituality, and existential explorations can be facilitated and improve end-of-life care is not well established. What is clear is that for physicians, one main intervention is to inquire about the role and importance of spirituality and religion in a patient's life. This will help a patient feel heard and help physicians identify specific needs. In one study, only 36% of respondents indicated that a clergy member would be comforting. Nevertheless, the increase in religious and spiritual interest among a substantial fraction of dying patients suggests inquiring of individual patients how this need can be addressed. Some evidence supports specific methods of addressing existential needs in patients, ranging from establishing a supportive group environment for terminal patients to individual treatments emphasizing a patient's dignity and sources of meaning.

MANAGING THE LAST STAGES

WITHDRAWING AND WITHHOLDING LIFE-SUSTAINING TREATMENT

LEGAL ASPECTS For centuries, it has been deemed ethical to withhold or withdraw life-sustaining interventions. The current legal consensus in the United States and most developed countries is that patients have a moral as well as constitutional or common law right to refuse medical interventions. American courts also have held that incompetent patients have a right to refuse medical interventions. For patients who are incompetent and terminally ill and who have not completed an advance care directive, next of kin can exercise that right, although this may be restricted in some states, depending how clear and convincing the evidence is of the patient's preferences. Courts have limited families' ability to terminate life-sustaining treatments in patients who are conscious, incompetent, but not terminally ill. In theory, patients' right to refuse medical therapy can be limited by four countervailing interests: (1) preservation of life, (2) prevention of suicide, (3) protection of third parties such as children, and (4) preservation of the integrity of

the medical profession. In practice, these interests almost never override the right of competent patients and incompetent patients who have left explicit and advance care directives.

For incompetent patients who either appointed a proxy without specific indications of their wishes or never completed an advance care directive, three criteria have been suggested to guide the decision to terminate medical interventions. First, some commentators suggest that ordinary care should be administered but extraordinary care could be terminated. Because the ordinary/extraordinary distinction is too vague, courts and commentators widely agree that it should not be used to justify decisions about stopping treatment. Second, many courts have advocated the use of the substituted-judgment criterion, which holds that the proxy decision-makers should try to imagine what the incompetent patient would do if he or she were competent. However, multiple studies indicate that many proxies, even close family members, cannot accurately predict what the patient would have wanted. Therefore, substituted judgment becomes more of a guessing game than a way of fulfilling the patient's wishes. Finally, the best-interests criterion holds that proxies should evaluate treatments by balancing their benefits and risks and select those treatments in which the benefits maximally outweigh the burdens of treatment. Clinicians have a clear and crucial role in this by carefully and dispassionately explaining the known benefits and burdens of specific treatments. Yet even when that information is as clear as possible, different individuals can have very different views of what is in the patient's best interests, and families may have disagreements or even overt conflicts. This criterion has been criticized because there is no single way to determine the balance between benefits and burdens; it depends on a patient's personal values. For instance, for some people, being alive even if mentally incapacitated is a benefit, whereas for others, it may be the worst possible existence. As a matter of practice, physicians rely on family members to make decisions that they feel are best and object only if those decisions seem to demand treatments that the physicians consider not beneficial.

PRACTICES Withholding and withdrawing acutely life-sustaining medical interventions from terminally ill patients are now standard practice. More than 90% of American patients die without cardiopulmonary resuscitation (CPR), and just as many forgo other potentially life-sustaining interventions. For instance, in ICUs in the period 1987–1988, CPR was performed 49% of the time, but it was performed only 10% of the time in 1992–1993. On average, 3.8 interventions, such as vasopressors and transfusions, were stopped for each dying ICU patient. However, up to 19% of decedents in hospitals received interventions such as extubation, ventilation, and surgery in the 48 h preceding death. However, practices vary widely among hospitals and ICUs, suggesting an important element of physician preferences rather than objective data.

Mechanical ventilation may be the most challenging intervention to withdraw. The two approaches are *terminal extubation*, which is the removal of the endotracheal tube, and *terminal weaning*, which is the gradual reduction of the or ventilator rate. One-third of ICU physicians prefer to use the terminal weaning technique, and 13% extubate; the majority of physicians use both techniques. The American Thoracic Society's 2008 clinical policy guidelines note that there is no single correct process of ventilator withdrawal and that physicians use and should be proficient in both methods but that the chosen approach should carefully balance benefits and burdens as well as patient and caregiver preferences. Physicians' assessment of patients' likelihood of survival, their prediction of possible cognitive damage, and patients' preferences about the use of life support are primary factors in determining the likelihood of withdrawal of mechanical ventilation. Some recommend terminal weaning because patients do not develop upper airway obstruction and the distress caused by secretions or stridor; however, terminal weaning can prolong the dying process and not allow a patient's family to be with him or her unencumbered by an endotracheal tube. To ensure comfort for conscious or semiconscious patients before withdrawal of the ventilator, neuromuscular blocking agents should be terminated and sedatives

and analgesics administered. Removing the neuromuscular blocking agents permits patients to show discomfort, facilitating the titration of sedatives and analgesics; it also permits interactions between patients and their families. A common practice is to inject a bolus of midazolam (2–4 mg) or lorazepam (2–4 mg) before withdrawal, followed by 5–10 mg of morphine and continuous infusion of morphine (50% of the bolus dose per hour) during weaning. In patients who have significant upper airway secretions, IV scopolamine at a rate of 100 µg/h can be administered. Additional boluses of morphine or increases in the infusion rate should be administered for respiratory distress or signs of pain. Higher doses will be needed for patients already receiving sedatives and opioids. Families need to be reassured about treatments for common symptoms after withdrawal of ventilatory support, such as dyspnea and agitation, and warned about the uncertainty of length of survival after withdrawal of ventilatory support: up to 10% of patients unexpectedly survive for 1 day or more after mechanical ventilation is stopped.

FUTILE CARE

Beginning in the late 1980s, some commentators argued that physicians could terminate futile treatments demanded by the families of terminally ill patients. Although no objective definition or standard of futility exists, several categories have been proposed. Physiologic futility means that an intervention will have no physiologic effect. Some have defined qualitative futility as applying to procedures that “fail to end a patient’s total dependence on intensive medical care.” Quantitative futility occurs “when physicians conclude (through personal experience, experiences shared with colleagues, or consideration of reported empiric data) that in the last 100 cases, a medical treatment has been useless.” The term conceals subjective value judgments about when a treatment is “not beneficial.” Deciding whether a treatment that obtains an additional 6 weeks of life or a 1% survival advantage confers benefit depends on patients’ preferences and goals. Furthermore, physicians’ predictions of when treatments were futile deviated markedly from the quantitative definition. When residents thought CPR was quantitatively futile, more than one in five patients had a >10% chance of survival to hospital discharge. Most studies that purport to guide determinations of futility are based on insufficient data to provide statistical confidence for clinical decision making. Quantitative futility rarely applies in ICU settings. Many commentators reject using futility as a criterion for withdrawing care, preferring instead to consider futility situations as ones that represent conflict that calls for careful negotiation between families and health care providers.

In the wake of a lack of consensus over quantitative measures of futility, many hospitals adopted process-based approaches to resolve disputes over futility and enhance communication with patients and surrogates, including focusing on interests and alternatives rather than opposing positions and generating a wide range of options. Some hospitals have enacted “unilateral do not resuscitate (DNR)” policies to allow clinicians to provide a DNR order in cases in which consensus cannot be reached with families and medical opinion is that resuscitation would be futile if attempted. This type of a policy is not a replacement for careful and patient communication and negotiation but recognizes that agreement cannot always be reached. Over the last 15 years, many states, such as Texas, Virginia, Maryland, and California, have enacted so-called medical futility laws that provide physicians a “safe harbor” from liability if they refuse a patient or family’s request for life-sustaining interventions. For instance, in Texas when a disagreement about terminating interventions between the medical team and the family has not been resolved by an ethics consultation, the hospital is supposed to try to facilitate transfer of the patient to an institution willing to provide treatment. If this fails after 10 days, the hospital and physician may unilaterally withdraw treatments determined to be futile. The family may appeal to a state court. Early data suggest that the law increases futility consultations for the ethics committee and that although most families concur with withdrawal, about 10–15% of families refuse to withdraw treatment. Approximately 12 cases have gone to court in Texas in the 7 years since the adoption of the law. As of 2007, there had been 974 ethics committee consultations on medical

futility cases and 65 in which committees ruled against families and gave notice that treatment would be terminated. Treatment was withdrawn for 27 of those patients, and the remainder were transferred to other facilities or died while awaiting transfer.

EUTHANASIA AND PHYSICIAN-ASSISTED SUICIDE

Euthanasia and physician-assisted suicide are defined in [Table 10-8](#). Terminating life-sustaining care and providing opioid medications to manage symptoms have long been considered ethical by the medical profession and legal by courts and should not be confused with euthanasia or physician-assisted suicide.

LEGAL ASPECTS Euthanasia is legal in the Netherlands, Belgium, and Luxembourg. It was legalized in the Northern Territory of Australia in 1995, but that legislation was repealed in 1997. Euthanasia is not legal in any state in the United States. With certain conditions, in Switzerland, a layperson can legally assist suicide. In the United States, physician-assisted suicide is legal in four states: Oregon, Vermont, and Washington State by legislation and Montana by court ruling. In jurisdictions where physician-assisted suicide is legal, physicians wishing to prescribe the necessary medication must fulfill multiple criteria and complete processes that include a waiting period. In other countries and all other states in the United States, physician-assisted suicide and euthanasia are illegal explicitly or by common law.

PRACTICES Fewer than 10–20% of terminally ill patients actually consider euthanasia and/or physician-assisted suicide for themselves. In the Netherlands and Oregon, >70% of patients using these interventions are dying of cancer; in Oregon, in 2013, just 1.2% of physician-assisted suicide cases involved patients with HIV/AIDS and 7.2% involved patients with amyotrophic lateral sclerosis. In the Netherlands, the share of deaths attributable to euthanasia or physician-assisted suicide declined from around 2.8% of all deaths in 2001 to around 1.8% in 2005. In 2013, the last year with complete data, around 71 patients in Oregon (just 0.2% of all deaths) died by physician-assisted suicide, although this may be an underestimate. In Washington State, between March 2009 (when the law allowing physician-assisted suicide went into force) and December 2009, 36 individuals died from prescribed lethal doses.

Pain is not a primary motivator for patients’ requests for or interest in euthanasia and/or physician-assisted suicide. Fewer than 25% of all patients in Oregon cite inadequate pain control as the reason for desiring physician-assisted suicide. Depression, hopelessness, and, more profoundly, concerns about loss of dignity or autonomy or being a burden on family members appear to be primary factors motivating a desire for euthanasia or physician-assisted suicide. Over 75% cite loss of autonomy or dignity and inability to engage in enjoyable activities as the reason for wanting physician-assisted suicide. About 40% cite being a burden on family. A study from the Netherlands showed that depressed terminally ill cancer patients were four times more likely

TABLE 10-8 DEFINITIONS OF ASSISTED SUICIDE AND EUTHANASIA

Term	Definition	Legal Status
Voluntary active euthanasia	Intentionally administering medications or other interventions to cause the patient’s death with the patient’s informed consent	Netherlands, Belgium
Involuntary active euthanasia	Intentionally administering medications or other interventions to cause the patient’s death when the patient was competent to consent but did not—e.g., the patient may not have been asked	Nowhere
Passive euthanasia	Withholding or withdrawing life-sustaining medical treatments from a patient to let him or her die (terminating life-sustaining treatments)	Everywhere
Physician-assisted suicide	A physician provides medications or other interventions to a patient with the understanding that the patient can use them to commit suicide	Oregon, Netherlands, Belgium, Switzerland

to request euthanasia and confirmed that uncontrolled pain was not associated with greater interest in euthanasia. Interestingly, despite the importance of emotional distress in motivating requests for euthanasia and physician-assisted suicide, few patients receive psychiatric care. For instance, in Oregon, only 5.9% of patients have been referred for psychiatric evaluation.

Euthanasia and physician-assisted suicide are no guarantee of a painless, quick death. Data from the Netherlands indicate that in as many as 20% of cases technical and other problems arose, including patients waking from coma, not becoming comatose, regurgitating medications, and experiencing a prolonged time to death. Data from Oregon indicate that between 1997 and 2013, 22 patients (~5%) regurgitated after taking prescribed medication, 1 patient awakened, and none experienced seizures. Problems were significantly more common in physician-assisted suicide, sometimes requiring the physician to intervene and provide euthanasia.

Whether practicing in a setting where euthanasia is legal or not, over a career, 12–54% of physicians receive a request for euthanasia or physician-assisted suicide from a patient. Competency in dealing with such a request is crucial. Although challenging, the request can also provide a chance to address intense suffering. After receiving a request for euthanasia and/or physician-assisted suicide, health care providers should carefully clarify the request with empathic, open-ended questions to help elucidate the underlying cause for the request, such as “What makes you want to consider this option?” Endorsing either moral opposition or moral support for the act tends to be counterproductive, giving an impression of being judgmental or of endorsing the idea that the patient’s life is worthless. Health care providers must reassure the patient of continued care and commitment. The patient should be educated about alternative, less controversial options, such as symptom management and withdrawing any unwanted treatments and the reality of euthanasia and/or physician-assisted suicide, because the patient may have misconceptions about their effectiveness as well as the legal implications of the choice. Depression, hopelessness, and other symptoms of psychological distress as well as physical suffering and economic burdens are likely factors motivating the request, and such factors should be assessed and treated aggressively. After these interventions and clarification of options, most patients proceed with another approach, declining life-sustaining interventions, possibly including refusal of nutrition and hydration.

CARE DURING THE LAST HOURS

Most laypersons have limited experiences with the actual dying process and death. They frequently do not know what to expect of the final hours and afterward. The family and other caregivers must be prepared, especially if the plan is for the patient to die at home.

Patients in the last days of life typically experience extreme weakness and fatigue and become bedbound; this can lead to pressure sores. The issue of turning patients who are near the end of life, however, must be balanced against the potential discomfort that movement may cause. Patients stop eating and drinking with drying of mucosal membranes and dysphagia. Careful attention to oral swabbing, lubricants for lips, and use of artificial tears can provide a form of care to substitute for attempts at feeding the patient. With loss of the gag reflex and dysphagia, patients may also experience accumulation of oral secretions, producing noises during respiration sometimes called “the death rattle.” Scopolamine can reduce the secretions. Patients also experience changes in respiration with periods of apnea or Cheyne-Stokes breathing. Decreased intravascular volume and cardiac output cause tachycardia, hypotension, peripheral coolness, and livedo reticularis (skin mottling). Patients can have urinary and, less frequently, fecal incontinence. Changes in consciousness and neurologic function generally lead to two different paths to death (Fig. 10-2).

Each of these terminal changes can cause patients and families distress, requiring reassurance and targeted interventions (Table 10-9). Informing families that these changes might occur and providing them with an information sheet can help preempt problems and minimize distress. Understanding that patients stop eating because they are dying, not dying because they have stopped eating, can reduce family

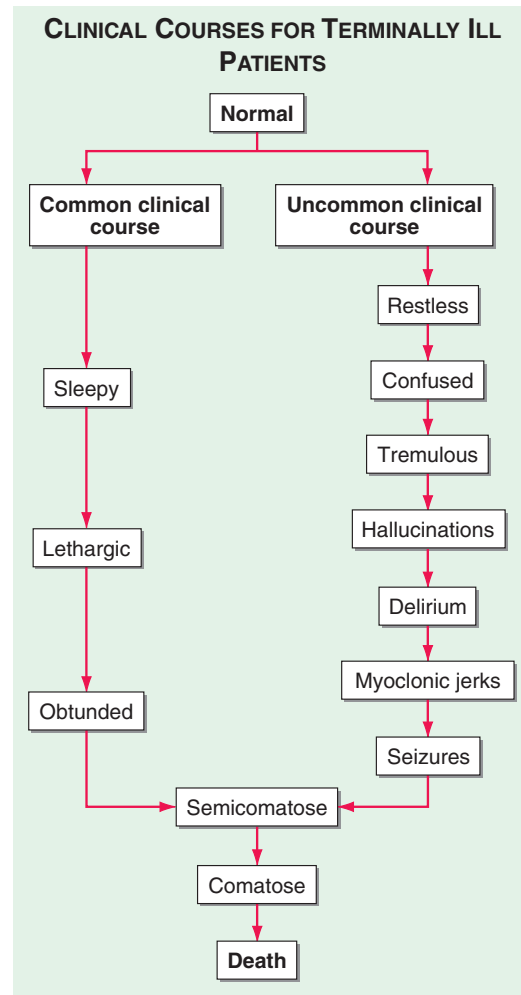


FIGURE 10-2 Common and uncommon clinical courses in the last days of terminally ill patients. (Adapted from FD Ferris et al: *Module 4: Palliative care, in Comprehensive Guide for the Care of Persons with HIV Disease*. Toronto: Mt. Sinai Hospital and Casey Hospice, 1995, <http://www.cpsonline.info/content/resources/hivmodule/module4complete.pdf>.)

and caregiver anxiety. Similarly, informing the family and caregivers that the “death rattle” may occur and that it is not indicative of suffocation, choking, or pain can reduce their worry from the breathing sounds.

Families and caregivers may also feel guilty about stopping treatments, fearing that they are “killing” the patient. This may lead to demands for interventions, such as feeding tubes, that may be ineffective. In such cases, the physician should remind the family and caregivers about the inevitability of events and the palliative goals. Interventions may prolong the dying process and cause discomfort. Physicians also should emphasize that withholding treatments is both legal and ethical and that the family members are not the cause of the patient’s death. This reassurance may have to be provided multiple times.

Hearing and touch are said to be the last senses to stop functioning. Whether this is the case or not, families and caregivers can be encouraged to communicate with the dying patient. Encouraging them to talk directly to the patient, even if he or she is unconscious, and hold the patient’s hand or demonstrate affection in other ways can be an effective way to channel their urge “to do something” for the patient.

When the plan is for the patient to die at home, the physician must inform the family and caregivers how to determine that the patient has died. The cardinal signs are cessation of cardiac function and respiration; the pupils become fixed; the body becomes cool; muscles relax; and incontinence may occur. Remind the family and caregivers that the eyes may remain open even after the patient has died because

TABLE 10-9 MANAGING CHANGES IN THE PATIENT'S CONDITION DURING THE FINAL DAYS AND HOURS

Changes in the Patient's Condition	Potential Complication	Family's Possible Reaction and Concern	Advice and Intervention
Profound fatigue	Bedbound with development of pressure ulcers that are prone to infection, malodor, and pain, and joint pain	Patient is lazy and giving up.	Reassure family and caregivers that terminal fatigue will not respond to interventions and should not be resisted. Use an air mattress if necessary.
Anorexia	None	Patient is giving up; patient will suffer from hunger and will starve to death.	Reassure family and caregivers that the patient is not eating because he or she is dying; not eating at the end of life does not cause suffering or death. Forced feeding, whether oral, parenteral, or enteral, does not reduce symptoms or prolong life.
Dehydration	Dry mucosal membranes (see below)	Patient will suffer from thirst and die of dehydration.	Reassure family and caregivers that dehydration at the end of life does not cause suffering because patients lose consciousness before any symptom distress. Intravenous hydration can worsen symptoms of dyspnea by pulmonary edema and peripheral edema as well as prolong dying process.
Dysphagia	Inability to swallow oral medications needed for palliative care		Do not force oral intake. Discontinue unnecessary medications that may have been continued, including antibiotics, diuretics, antidepressants, and laxatives. If swallowing pills is difficult, convert essential medications (analgesics, antiemetics, anxiolytics, and psychotropics) to oral solutions, buccal, sublingual, or rectal administration.
"Death rattle"—noisy breathing		Patient is choking and suffocating.	Reassure the family and caregivers that this is caused by secretions in the oropharynx and the patient is not choking. Reduce secretions with scopolamine (0.2–0.4 mg SC q4h or 1–3 patches q3d). Reposition patient to permit drainage of secretions. Do not suction. Suction can cause patient and family discomfort and is usually ineffective.
Apnea, Cheyne-Stokes respirations, dyspnea		Patient is suffocating.	Reassure family and caregivers that unconscious patients do not experience suffocation or air hunger. Apneic episodes are frequently a premonitory change. Opioids or anxiolytics may be used for dyspnea. Oxygen is unlikely to relieve dyspneic symptoms and may prolong the dying process.
Urinary or fecal incontinence	Skin breakdown if days until death Potential transmission of infectious agents to caregivers	Patient is dirty, malodorous, and physically repellent.	Remind family and caregivers to use universal precautions. Frequent changes of bedclothes and bedding. Use diapers, urinary catheter, or rectal tube if diarrhea or high urine output.
Agitation or delirium	Day/night reversal Hurt self or caregivers	Patient is in horrible pain and going to have a horrible death.	Reassure family and caregivers that agitation and delirium do not necessarily connote physical pain. Depending on the prognosis and goals of treatment, consider evaluating for causes of delirium and modify medications. Manage symptoms with haloperidol, chlorpromazine, diazepam, or midazolam.
Dry mucosal membranes	Cracked lips, mouth sores, and candidiasis can also cause pain. Odor	Patient may be malodorous or physically repellent.	Use baking soda mouthwash or saliva preparation q15–30min. Use topical nystatin for candidiasis. Coat lips and nasal mucosa with petroleum jelly q60–90min. Use ophthalmic lubricants q4h or artificial tears q30min.

the retroorbital fat pad may be depleted, permitting the orbit to fall posteriorly, which makes it difficult for the eyelids to cover the eyeball.

The physician should establish a plan for who the family or caregivers will contact when the patient is dying or has died. Without a plan, they may panic and call 911, unleashing a cascade of unwanted events, from arrival of emergency personnel and resuscitation to hospital admission. The family and caregivers should be instructed to contact the hospice (if one is involved), the covering physician, or the on-call member of the palliative care team. They should also be told that the medical examiner need not be called unless the state requires it for all deaths. Unless foul play is suspected, the health care team need not contact the medical examiner either.

Just after the patient dies, even the best-prepared family may experience shock and loss and be emotionally distraught. They need time to assimilate the event and be comforted. Health care providers are likely to find it meaningful to write a bereavement card or letter to the family. The purpose is to communicate about the patient, perhaps emphasizing the patient's virtues and the honor it was to care for the patient, and to express concern for the family's hardship. Some physicians attend the funerals of their patients. Although this is beyond any medical obligation, the presence of the physician can be a source of support to the grieving family and provides an opportunity for closure for the physician.

Death of a spouse is a strong predictor of poor health, and even mortality, for the surviving spouse. It may be important to alert the

spouse's physician about the death so that he or she is aware of symptoms that might require professional attention.

PALLIATIVE CARE SERVICES: HOW AND WHERE

Determining the best approach to providing palliative care to patients will depend on patient preferences, the availability of caregivers and specialized services in close proximity, institutional resources, and reimbursement. Hospice is a leading, but not the only, model of palliative care services. In the United States, a plurality—41.5%—of hospice care is provided in residential homes. In 2012, just over 17% of hospice care was provided in nursing homes. In the United States, Medicare pays for hospice services under Part A, the hospital insurance part of reimbursement. Two physicians must certify that the patient has a prognosis of ≤ 6 months if the disease runs its usual course. Prognoses are probabilistic by their nature; patients are not required to die within 6 months but rather to have a condition from which half the individuals with it would not be alive within 6 months. Patients sign a hospice enrollment form that states their intent to forgo curative services related to their terminal illness, but they can still receive medical services for other comorbid conditions. Patients also can withdraw enrollment and reenroll later; the hospice Medicare benefit can be revoked later to secure traditional Medicare benefits. Payments to the hospice are per diem (or capitated), not fee-for-service. Payments are intended to cover physician services for the medical direction of the care team; regular home care visits by registered nurses and licensed practical nurses; home health aid and homemaker services; chaplain services; social work services; bereavement counseling; and medical equipment, supplies, and medications. No specific therapy is excluded, and the goal is for each therapy to be considered for its symptomatic (as opposed to disease-modifying) effect. Additional clinical care, including services of the primary physician, is covered by Medicare Part B even while the hospice Medicare benefit is in place. The health reform legislation signed into law in March 2010—the Affordable Care Act—directs the Secretary of Health and Human Services to gather data on Medicare hospice reimbursement with the goal of reforming payment rates to account for resource use over an entire episode of care. The legislation also requires additional evaluations and reviews of eligibility for hospice care by hospice physicians or nurses. Finally, the legislation establishes a demonstration project for concurrent hospice care in Medicare, which would test and evaluate allowing patients to remain eligible for regular Medicare during hospice care.

By 2012, the mean length of enrollment in a hospice was around 71.8 days, with the median being 18.7 days. Such short stays create barriers to establishing high-quality palliative services in patients' homes and also place financial strains on hospice providers because the initial assessments are resource intensive. Physicians should initiate early referrals to the hospice to allow more time for patients to receive palliative care.

Hospice care has been the main method in the United States for securing palliative services for terminally ill patients. However, efforts are being made to ensure continuity of palliative care across settings and through time. Palliative care services are becoming available as consultative services and more rarely as palliative care units in hospitals, in day care and other outpatient settings, and in nursing homes. Palliative care consultations for nonhospice patients can be billed as for other consultations under Medicare Part B, the physician reimbursement part. Many believe palliative care should be offered to patients regardless of their prognosis. A patient, his or her family, and physicians should not have to make a "curative versus palliative care" decision because it is rarely possible to make such a decisive switch to embracing mortality.

FUTURE DIRECTIONS

OUTCOME MEASURES

Care near the end of life cannot be measured by most of the available validated outcome measures because palliative care does not consider death a bad outcome. Similarly, the family and patients receiving end-of-life care may not desire the elements elicited in current quality-of-life measurements. Symptom control, enhanced family relationships, and quality of bereavement are difficult to measure and are

rarely the primary focus of carefully developed or widely used outcome measures. Nevertheless, outcomes are as important in end-of-life care as in any other field of medical care. Specific end-of-life care instruments are being developed both for assessment, such as The Brief Hospice Inventory and NEST (*needs near the end of life screening tool*), and for outcome measures, such as the Palliative Care Outcomes Scale, as well as for prognosis, such as the Palliative Prognostic Index. The field of end-of-life care is entering an era of evidence-based practice and continuous improvement through clinical trials.

11 Clinical Problems of Aging

Luigi Ferrucci, Stephanie Studenski

While an in-depth understanding of internal medicine serves as a foundation, proper care of older adults should be complemented by insight into the multidimensional effects of aging on disease manifestations, consequences, and response to treatment. In younger adults, individual diseases tend to have a more distinct pathophysiology with well-defined risk factors; the same diseases in older persons may have a less distinct pathophysiology and are often the result of failed homeostatic mechanisms. Causes and clinical manifestations are less specific and can vary widely between individuals. Therefore, the care of older patients demands an understanding of the effects of aging on human physiology and a broader perspective that incorporates geriatric syndromes, disability, social contexts, and goals of care. For example, care planning for the older patient should account for the substantial portion of the wide variability in life expectancy across individuals of the same age that can be predicted by simple and inexpensive measures such as walking speed. Estimation of the expected remaining years of life can guide recommendations about appropriate preventive and other long-term interventions and can shape discussions about treatment alternatives.

DEMOGRAPHY

(See also Chap. 93e) Population aging emerged as a worldwide phenomenon for the first time in history within the past century. Since aging influences many facets of life, governments and societies—as well as families and communities—now face new social and economic challenges that affect health care. Fig. 11-1 highlights recent and predicted changes in U.S. population structure.

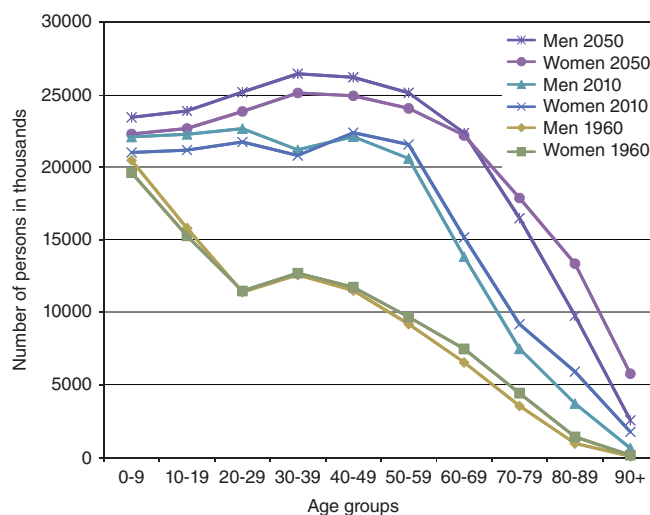


FIGURE 11-1 Change in the structure of the U.S. population between 1960 and 2050. (From United Nations World Population Prospects: The 2008 Revision, http://www.un.org/esa/population/publications/wpp2008/wpp2008_highlights.pdf.)

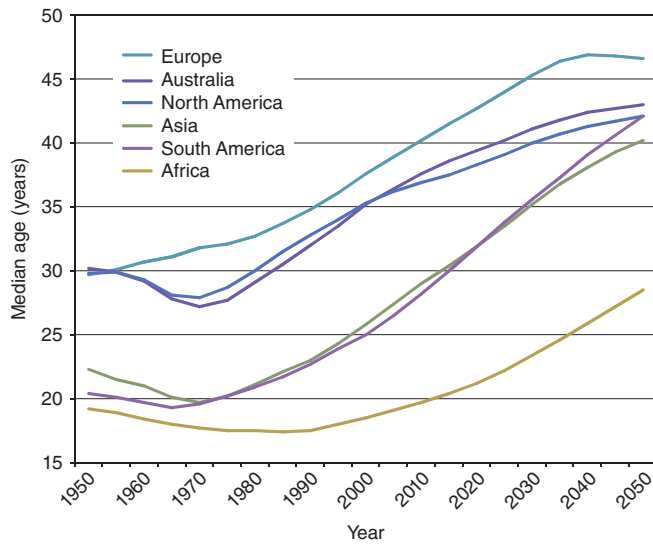


FIGURE 11-2 Population aging in different geographic regions. (From United Nations World Population Prospects: The 2008 Revision, http://www.un.org/esa/population/publications/wpp2008/wpp2008_highlights.pdf)

The overall number of children has remained relatively stable, but explosive growth has occurred among older populations. The percentage of growth is particularly dramatic among the oldest of the old. For example, the number of persons aged 80–89 years more than tripled between 1960 and 2010 and will increase over tenfold between 1960 and 2050. Women already outlive men by many years, and the sex discrepancy in longevity is projected to increase further in the future.

Population aging occurs at different rates in varying geographic regions of the world. Over the past century, Europe, Australia, and North America have had the populations with the greatest proportions of older persons, but the populations of Asia and South America are aging rapidly, and the population structure on these continents will resemble that of “older” countries by around 2050 (Fig. 11-2). Among older persons, the oldest old (those >80 years of age) are the fastest-growing segment of the population (Fig. 11-3), and the pace of population aging is projected to accelerate in most countries over the next 50 years. There is no evidence that the rate of population aging is decreasing.

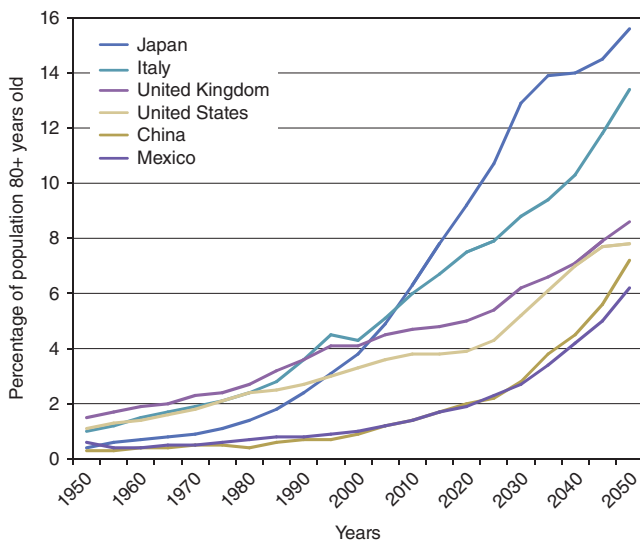


FIGURE 11-3 Percentage of the population age >80 years from 1950 to 2050 in representative nations. The pace of population aging will accelerate. (From United Nations World Population Prospects: The 2008 Revision, http://www.un.org/esa/population/publications/wpp2008/wpp2008_highlights.pdf)

POPULATION AGING AND HEALTH

Many chronic diseases increase in prevalence with age. It is not unusual for older persons to have multiple chronic diseases (Fig. 11-4), although some seem more susceptible than others to co-occurring problems. Functional problems that pose difficulties or require help in performing basic activities of daily living (ADLs) (Table 11-1) increase with age and are more common among women than among men. In recent decades, the age-specific prevalence of disability has declined, especially in the oldest old. Estimated rates are shown in Fig. 11-5 as the percentage of persons who reported severe difficulty or needed help in bathing, and data on other basic ADLs show similar trends. Although the age-specific prevalence of disability is decreasing, the magnitude of this decline is small compared to the overwhelming effect of population aging. Thus, the number of people with disability in the United States and other countries is rapidly expanding. Rates of cognitive impairments, such as memory problems, also increase with aging (Fig. 11-6). Chronic disease and disability lead to increased use of health care resources. Health care expenditures increase with age, increase more with disability, and are highest in the last year of life. However, new medical technologies and expensive medications are greater influences on health care costs than population aging itself. General practitioners and internists with little specific training in geriatric medicine provide the bulk of care for older persons.

SYSTEMIC EFFECTS OF AGING

Systemic consequences of aging are widespread but can be clustered into four main domains or processes (Fig. 11-7): (1) body composition; (2) balance between energy availability and energy demand; (3) signaling networks that maintain homeostasis; and (4) neurodegeneration. Each domain can be assessed with routine clinical tests, although more detailed research techniques are also available (Table 11-2).

Body Composition Profound changes in body composition may be the most evident and inescapable effect of aging (Fig. 11-8). Over the life span, body weight tends to increase through childhood, puberty, and adulthood until late middle age. Weight tends to decline in men between ages 65 and 70 years and in women somewhat later. Lean body mass, composed predominantly of muscle and visceral organs, decreases steadily after the third decade. In muscle, this atrophy is greater in fast-twitch than in slow-twitch fibers. The origin of this change is unknown, but several lines of evidence suggest that progressive loss of motor neurons probably plays an important role. Fat mass tends to increase in middle age and then declines in late life, reflecting the trajectory of weight change. Waist circumference continues to increase across the life span, a pattern suggesting that visceral fat, which is responsible for most of the pathologic consequences of obesity, continues to accumulate. In some individuals, fat also accumulates inside muscle, affecting muscle quality and function. With age, fibroconnective tissue tends to increase in many organ systems. In muscle, fibroconnective tissue buildup also affects muscle quality and function. In combination, the loss of muscle mass and quality result in reduced muscle strength, which ultimately affects functional capacity and mobility. Muscle strength declines with aging; this decrease not only affects functional status but also is a strong independent predictor of mortality (Fig. 11-9). Progressive demineralization and architectural modification occur in bone, resulting in a decline of bone strength. Loss of bone strength increases the risk of fracture. Sex differences in the effects of aging on bone mass are due to differences in peak bone mass and the effects of gonadal hormones on bone. Overall, compared with men, women tend to lose bone mass at a younger age and more quickly reach the threshold of low bone strength that increases fracture risk.

All of these changes in body composition can be attributed to disruptions in the links between synthesis, degradation, and repair that normally serve to remodel tissues. Such changes in body composition are influenced not only by aging and illness but also by lifestyle factors such as physical activity and diet. Body composition can be approximated in clinical practice on the basis of weight, height, body mass index (BMI; weight in kilograms divided by height in meters squared), and waist circumference or, more precisely, with dual-energy x-ray

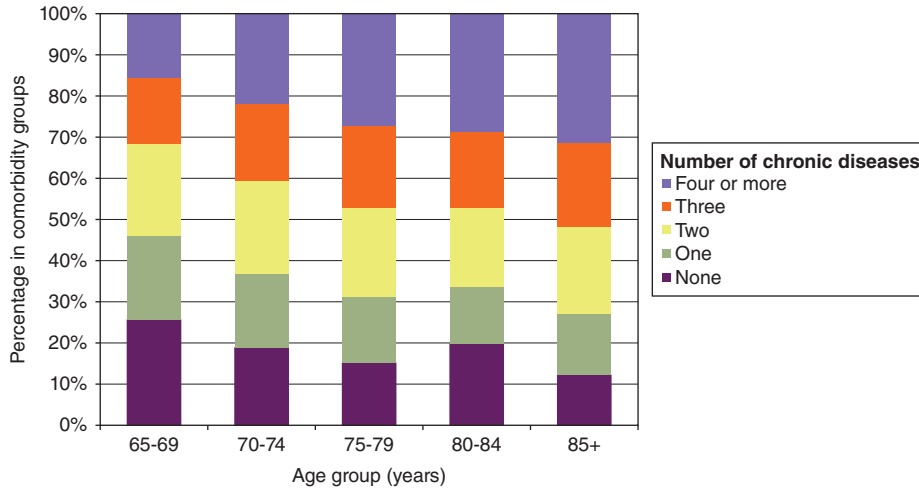


FIGURE 11-4 Prevalence of comorbidity by age group in persons ≥ 65 years old living in the United States and enrolled in Medicare parts A and B in 1999. (From JL Wolff et al: *Arch Intern Med* 162:2269, 2002.)

TABLE 11-1 BASIC AND INSTRUMENTAL ACTIVITIES OF DAILY LIVING

Basic Activities of Daily Living: Self-Care Tasks

- Personal hygiene
- Dressing and undressing
- Eating
- Transferring from bed to chair and back
- Voluntarily controlling urinary and fecal discharge
- Using the toilet
- Moving around (as opposed to being bedridden)

Instrumental Activities of Daily Living: Not Necessary for Fundamental Functioning, but Permit an Individual to Live Independently in a Community

- Doing light housework
- Preparing meals
- Taking medications
- Shopping for groceries or clothes
- Using the telephone
- Managing money
- Using technology^a

^aWith the recognition that older persons may not be that technologically savvy since they were not as extensively exposed to technology during their lifetime.)

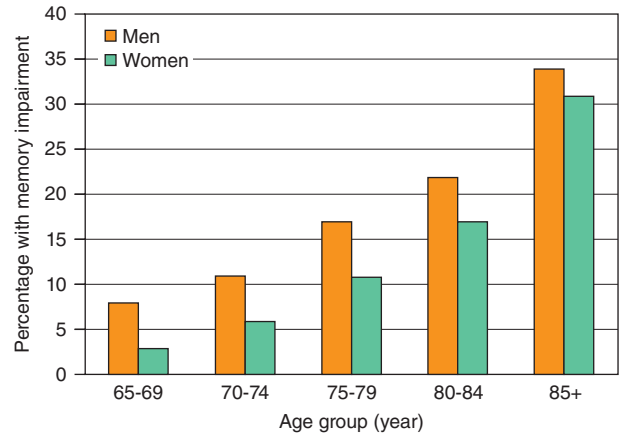


FIGURE 11-6 Rates of memory impairment in different age groups. The definition of “moderate or severe memory impairment” is 4 or fewer words recalled out of 20. (Source: *Health and Retirement Survey*. Accessed November 15, 2013, at aqa.gov/agingstatsdotnet/Main_Site/Data/2000_Documents/healthstatus.aspx.)

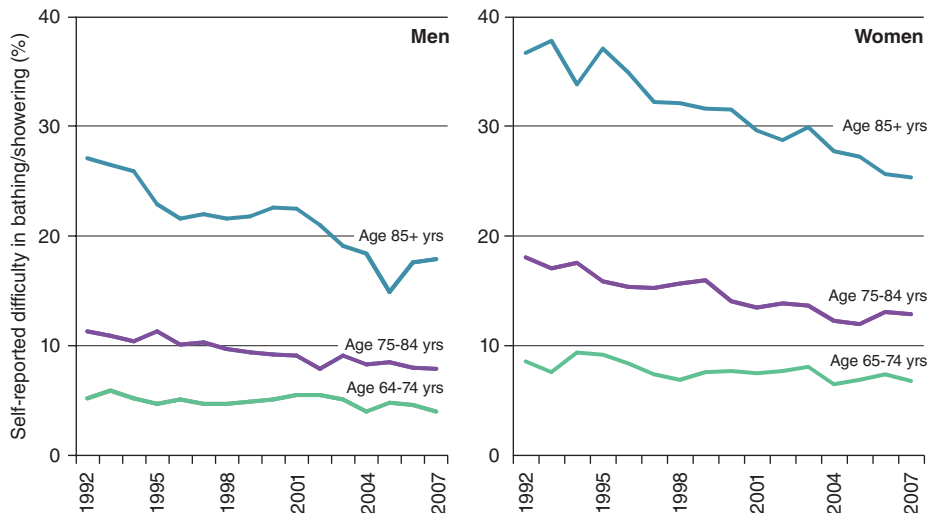


FIGURE 11-5 Self-reported prevalence of disability (severe difficulty) in bathing/showering between 1992 and 2007, according to age and sex. (From *Medicare Current Beneficiary Survey 1992–2007*.)

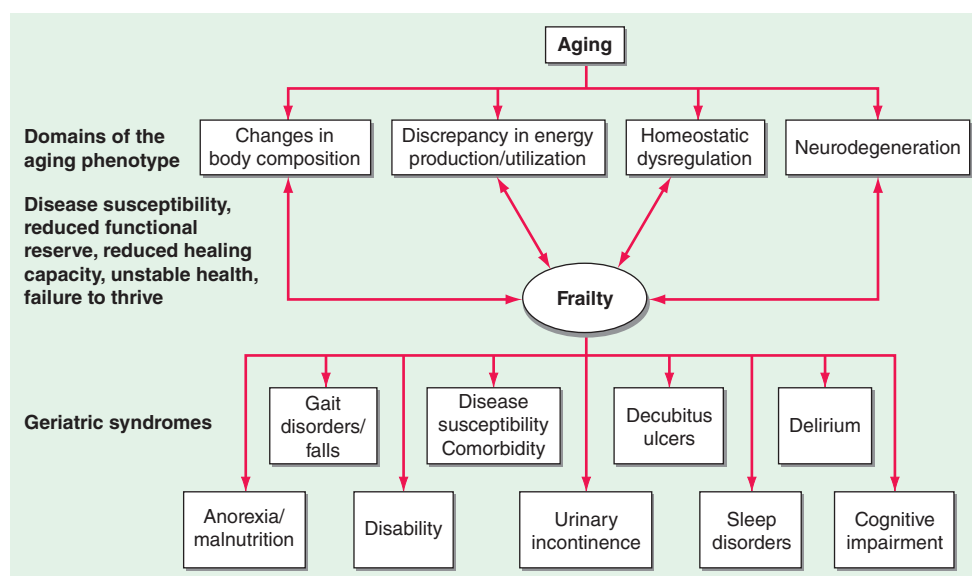


FIGURE 11-7 A unifying model of aging, frailty, and the geriatric syndromes.

absorptiometry, CT, or MRI. In healthy men and women in their twenties, lean body mass is, on average, 85% of body weight, with roughly 50% of lean mass represented by skeletal muscle. With aging, both the percentage of lean mass and the percentage represented by muscle decline rapidly, and these changes have important health and functional consequences.

Balance Between Energy Availability and Energy Demand Release of phosphate from ATP provides every living cell with the energy required for life. However, the storage of ATP is only enough for 6 sec; therefore, ATP is constantly resynthesized. Although ATP can be resynthesized by anaerobic glycolysis, most of the energy used in the body is generated through aerobic metabolism. Therefore, energy consumption is usually estimated indirectly by oxygen consumption (indirect

calorimetry). There is currently no method to measure true “fitness,” which is the maximal energy that can be produced by an organism over extended periods. Thus, fitness is estimated indirectly from peak oxygen consumption ($MVO_{2\text{peak}}$), often during a maximal treadmill test. Longitudinal studies have demonstrated that $MVO_{2\text{peak}}$ declines progressively with aging (Fig. 11-10), and the rate of decline is accelerated in persons who are sedentary and in those affected by chronic diseases.

A large portion of energy is consumed as the “resting metabolic rate” (RMR)—i.e., the amount of energy expended at rest in a neutral temperature environment and in a postabsorptive state. In healthy men and women, RMR declines with aging, and such decline is only partially explained by the parallel decline in the highly metabolically active tissues that make up lean body mass (Fig. 11-11). However, persons with unstable homeostasis due to illness require additional

TABLE 11-2 EXAMPLES OF ASSESSMENT OF THE FOUR DOMAINS OF THE AGING PHENOTYPE

Approach to Assessment	Body Composition	Energetics	Homeostatic Regulation	Neurodegeneration
Self-report		Self-reported questionnaires investigating physical activity, sense of fatigue/exhaustion, exercise tolerance		
Physical examination	Muscle strength testing (isometric and isokinetic) Anthropometrics (weight, height, BMI, waist circumference, arm and leg circumference, skin folds)	Performance-based tests of physical function		Objective assessment of gait, balance, reaction time, coordination Standard neurologic exam, including assessment of global cognition ^a
Laboratory values	Biomarkers (24-h creatinuria or 3-methyl-histidine)		Nutritional biomarkers (e.g., vitamins, antioxidants) Baseline levels of biomarkers and hormone levels Inflammatory markers (e.g., ESR, CRP, IL-6, TNF- α)	
Imaging	CT and MRI, DEXA	Magnetic resonance spectroscopy		MRI, fMRI, PET, and other dynamic imaging techniques
Other	Hydrostatic weighing	Resting metabolic rate Treadmill testing of oxygen consumption during walking Objective measures of physical activity (accelerometers, double-labeled water)	Stress response Response to provocative tests, such as oral glucose tolerance test, dexamethasone test, and others	Evoked potentials Electroneurography and electromyography

^aMini Mental State; Montreal Cognitive Assessment.

Abbreviations: BMI, body mass index; CRP, C-reactive protein; DEXA, dual-energy x-ray absorptiometry; ESR, erythrocyte sedimentation rate; fMRI, functional MRI; IL-6, interleukin 6; PET, positron emission tomography; TNF α , tumor necrosis factor α .

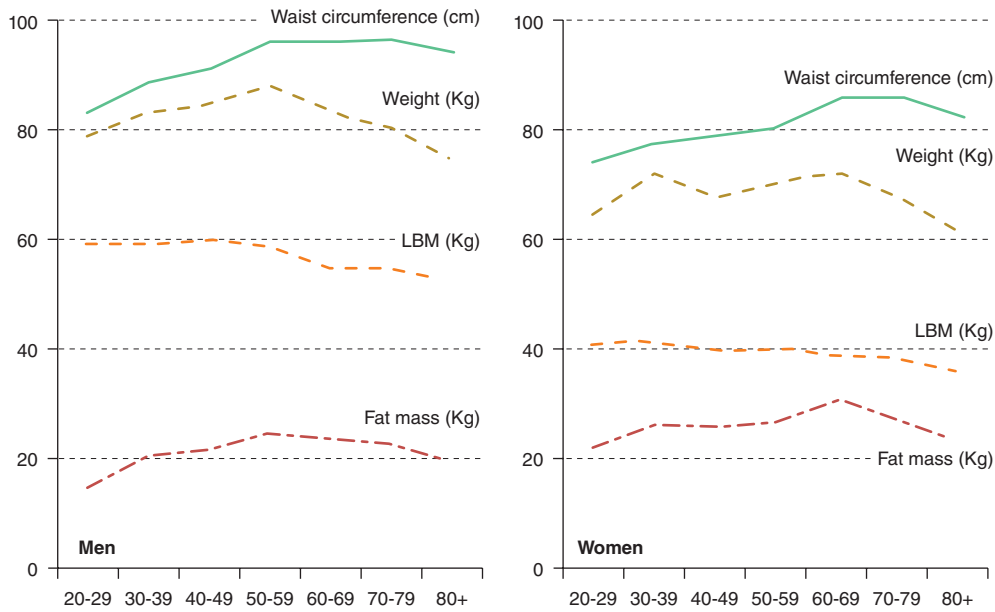


FIGURE 11-8 Longitudinal changes of weight, body composition, and waist circumference over the life span, estimated in 1167 participants in the Baltimore Longitudinal Study of Aging. Lean body mass (LBM) and fat mass were estimated with dual-energy x-ray absorptiometry. (Source: *The Baltimore Longitudinal study of Aging 2010*; unpublished data.)

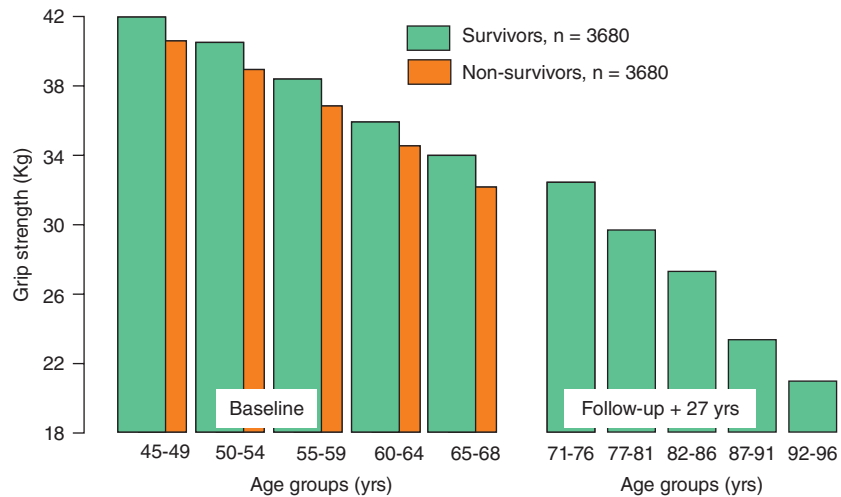


FIGURE 11-9 Cross-sectional differences and longitudinal changes in muscle strength over a 27-year follow-up. Note that persons who died during the follow-up had lower baseline muscle strength. (From *T Rantanen et al: J Appl Physiol 85:2047, 1998*.)

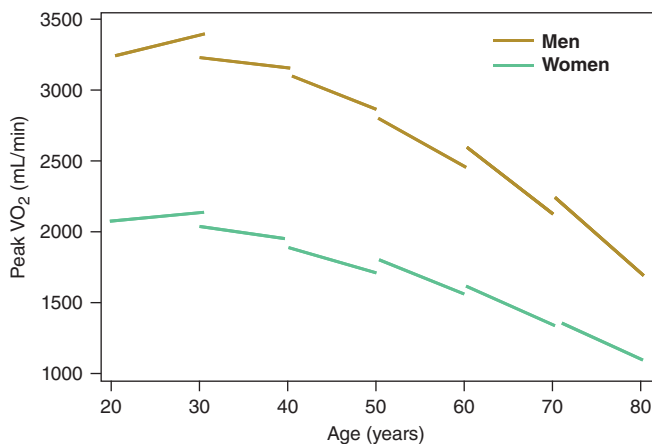


FIGURE 11-10 Longitudinal changes in aerobic capacity in participants in the Baltimore Longitudinal Study of Aging. (From *JL Fleg: Circulation 112:674, 2005*.)

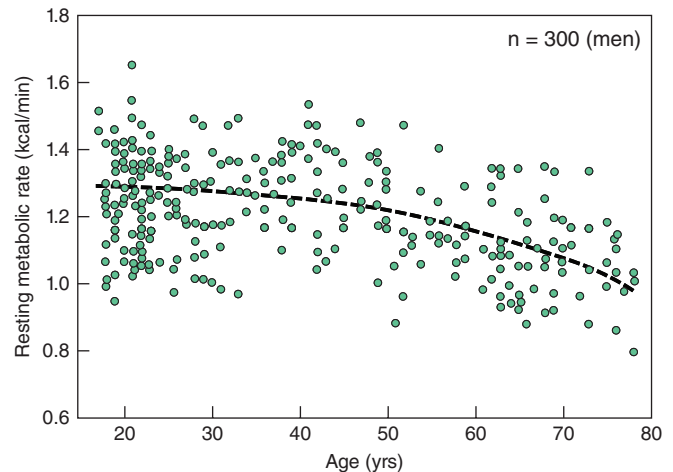


FIGURE 11-11 Changes in resting metabolic rate with aging. (Unpublished data from the Baltimore Longitudinal Study of Aging.)

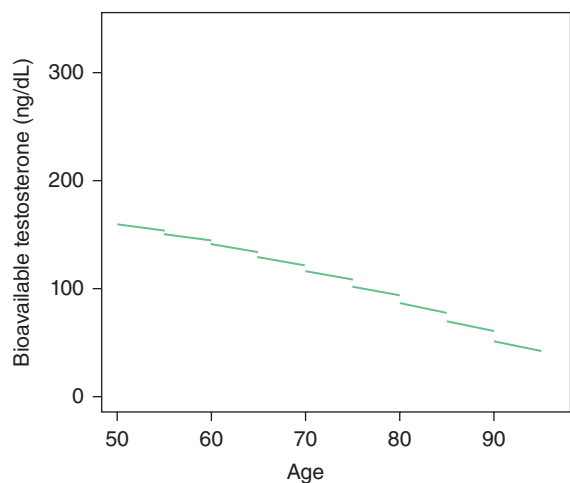


FIGURE 11-12 Longitudinal trajectory of bioavailable testosterone plasma concentration in the Baltimore Longitudinal Study of Aging (BLSA). The plot is based on 584 men who were 50 years or older with a total of 1455 data points. The average follow up for each subject was 3.2 years. (Figure created using unpublished data from the BLSA.)

energy for compensatory mechanisms. Indeed, observational studies have demonstrated (1) that older persons with poor health status and substantial morbidity have a higher RMR than healthier individuals of the same age and sex and (2) that a high RMR is an independent risk factor for mortality and may contribute to the weight loss that often accompanies severe illness. Finally, for reasons that are not yet completely clear but certainly involve changes in the biomechanical characteristics of movement, older age, pathology, and physical impairment increase the energy cost of motor activities such as walking. Overall, older individuals with multiple chronic conditions have low available energy levels and require more energy both at rest and during physical activity. Thus, sick older people may consume all their available energy performing the most basic ADLs, and consequent fatigue and restriction may lead to a sedentary existence. Energy status can be assessed clinically by simply asking patients about their perceived level of fatigue during daily activities such as walking or dressing. Energy capacity can be assessed more precisely by exercise tolerance during a walking test or a treadmill test coupled with spirometry.

The main signaling pathways that control homeostasis involve hormones, inflammatory mediators, and antioxidants; all are profoundly affected by aging. Sex hormone levels, such as testosterone in men (Fig. 11-12) and estrogen in women, decrease with age, while other hormone systems may change more subtly (Table 11-3). Most aging individuals, even those who remain healthy and fully functional, tend to develop a mild proinflammatory state characterized by high levels of proinflammatory markers, including interleukin 6 (IL-6) and C-reactive protein (CRP) (Fig. 11-13). Aging is also thought to be associated with increased oxidative stress damage, either because the production of reactive oxygen species increases or because antioxidant buffers are less effective. Since hormones, inflammatory markers, and antioxidants are integrated into complex signaling networks, levels of individual biomarkers may well reflect adaptation within homeostatic feedback loops rather than true causative factors. Thus, the therapeutic strategy of single-molecule replacement may be ineffective or even counterproductive. The presence of such signaling networks and feedback loops may help explain why single-hormone “replacement therapy” for problems of aging has demonstrated little benefit. The focus of research in this area is now on multiple-hormone

TABLE 11-3 HORMONES THAT DECREASE, REMAIN STABLE, AND INCREASE WITH AGING

Decrease	No Change	Increase
Growth hormone	Prolactin	Cholecystokinin
Luteinizing hormone (men)	Thyrotropin	Luteinizing hormone (women)
Insulin growth factor 1	Thyroid hormones	Follicle-stimulating hormone
Testosterone	Epinephrine	Cortisol
Estradiol	Glucagon-like peptide 1	Prolactin
Dehydroepiandrosterone	Gastric inhibitory polypeptide	Norepinephrine
Pregnenolone		Insulin
25-Hydroxyvitamin D		Parathormone
Aldosterone		
Vasoactive intestinal peptide		
Melatonin		

dysregulation. For example, taken one at a time, levels of testosterone, dehydroepiandrosterone (DHEA), and insulin-like growth factor 1 (IGF-1) do not predict mortality, but in combination they are highly predictive of longevity. This combination effect is especially strong in the setting of congestive heart failure. Similarly, several micronutrients, such as vitamins (especially vitamin D), minerals (selenium and magnesium), and antioxidants (vitamins D and E), also regulate aspects of metabolism. Low levels of these micronutrients have been associated with accelerated aging and a high risk of adverse outcomes. However, except for vitamin D, no clear evidence suggests that supplementation has positive effects on health. Unfortunately, no standard criteria exist that allow the detection and quantification of homeostatic dysregulation as a general phenomenon.

Neurodegeneration It was long generally believed that neurons stop reproducing shortly after birth and that their number declines throughout life. However, results from animal models and even some studies in humans suggest that neurogenesis in the hippocampus continues at low levels throughout life. Brain atrophy occurs with aging after the age of 60 years. Atrophy proceeds at varying rates in different parts of the brain (Fig. 11-14) and is often accompanied by an inflammatory response and microglial activation. Age-associated brain atrophy may contribute to age-related declines in cognitive and motor function. Atrophy may also be a factor in some brain diseases that can occur with aging, such as mild cognitive impairment, in which persons have mild but detectable impairments on tests of cognition but no severe disability in daily activities. In mild cognitive impairment, atrophy has been found mostly in the prefrontal cortex and hippocampus,

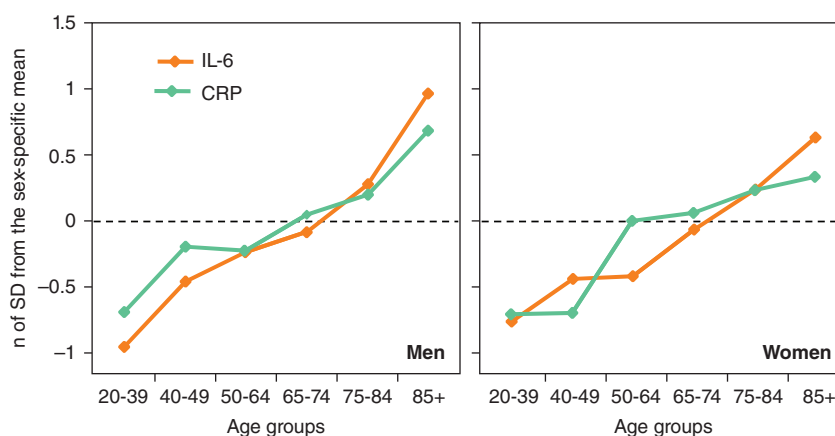


FIGURE 11-13 Change in interleukin 6 (IL-6) and C-reactive protein (CRP) with aging. Values are expressed as Z-scores to make them comparable. (From L Ferrucci et al: *Blood* 105:2294, 2005.)

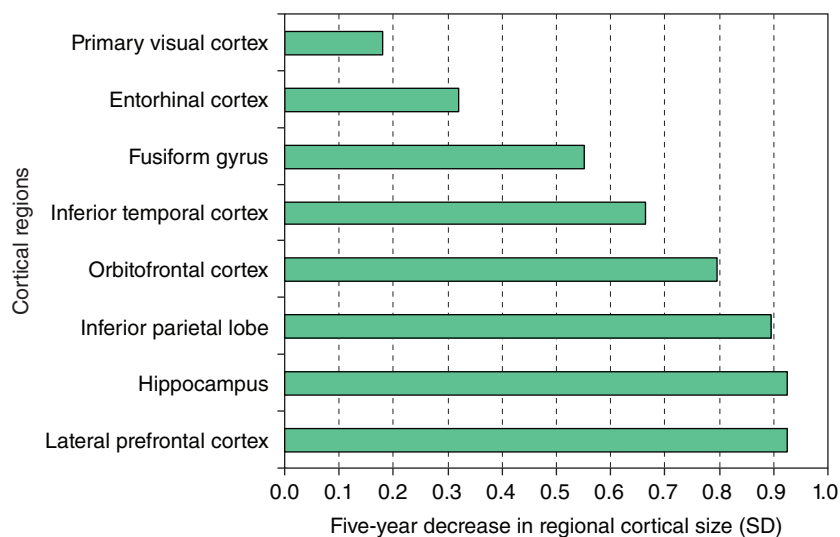


FIGURE 11-14 Five-year decline in mean volumes of different brain regions, measured in standard deviation (SD) units (Cohen's *d*). The primary visual cortex shows the least average shrinkage, and the prefrontal and inferior parietal cortex and hippocampus show the most average shrinkage. (From N Raz et al: *Ann N Y Acad Sci* 1097:84, 2007.)

but these findings are not specific and their diagnostic utility is unclear (Fig. 11-15). Other neurophysiologic changes in the brain frequently occur with aging and may contribute to cognitive decline. Functional imaging studies have shown that some older people have diminished coordination between the brain regions responsible for higher-order cognitive functions and that such diminished coordination is correlated with poor cognitive performance. In young healthy individuals, the brain activity associated with executive cognitive functions (e.g., problem-solving, decision-making) is very well localized; in contrast, in healthy older individuals, the pattern of cortical activation is more diffuse. Brain pathology has typically been associated with specific diseases; amyloid plaques and neurofibrillary tangles are considered the pathologic hallmarks of Alzheimer's disease. However, these pathologic markers have been found at autopsy in many older individuals who had normal cognition, as assessed by extensive testing in the year before death.

Taken together, trends in brain changes with aging suggest that some neurophysiologic manifestations are compensatory adaptations rather than primary contributors to age-related declines. Because the brain is capable of reorganization and compensation, extensive neurodegeneration may not be clinically evident. Therefore, early detection requires careful testing. Clinically, cortical and subcortical changes are reflected in the high prevalence of "soft," nonspecific neurologic signs, often reflected in slow and unstable gait, poor balance, and slow reaction times. These movement changes can be elicited more overtly with "dual tasks," in which a cognitive and a motor task are performed simultaneously. In a simple version of a dual task, when an older adult has to stop walking in order to talk, an increased risk of falls can be predicted. Poor dual-task performance has been interpreted as a marker of reduced overall capacity for central processing, so that simultaneous processing is more constrained. Beyond the brain, the spinal cord also experiences changes after the age of 60 years, including reduced numbers of motor neurons and damage to myelin. The motor neurons that survive compensate by increased branching complexity and by service to larger motor units. As motor units become larger, they decline

in number at a rate of ~1% per year, starting after the third decade. These larger motor units contribute to reductions in fine-motor control and manual dexterity. Age-related changes also occur in the autonomic nervous system, affecting cardiovascular and splanchnic function.

Systemic Changes Coexisting with and Affecting One Another • THE PHENOTYPE OF AGING: THE FINAL COMMON PATHWAY OF SYSTEMIC INTERACTION While age-related system changes have been described individually, in reality, these changes develop in parallel and affect one another through many feed-forward and feedback loops. Some systemic interactions are well understood, while others are under investigation. For example, body composition interacts with energy balance and signaling. Higher lean body mass increases energy consumption and improves insulin sensitivity and carbohydrate metabolism. Higher fat mass, especially visceral fat mass, is the culprit in the metabolic syndrome and is associated with low testosterone levels, high sex hormone-binding globulin levels, and increased levels of proinflammatory markers such as CRP and IL-6. Altered signaling can affect neurodegeneration; insulin resistance and adipokines such as leptin and adiponectin are associated with declines in

cognitive function. Combined with loss of motor neurons and dysfunction of the motor unit, a state of inflammation and reduced levels of testosterone and IGF-1 have been linked to accelerated decline of muscle mass and strength. Normal intersystem coordination is also affected by aging. The hypothalamus normally functions as a central regulator of metabolism and energy use and coordinates physiologic responses of the entire organism through hormonal signaling; aging-related changes in the hypothalamus alter this control. The central nervous system (CNS) also controls adaptive sympathetic/parasympathetic activity, so that age-related CNS degeneration may have implications for autonomic function.

The phenotype that results from the aging process is characterized by increased susceptibility to diseases, high risk of multiple coexisting diseases, impaired response to stress (including limited ability to heal or recover after an acute disease), emergence of "geriatric syndromes" (characterized by stereotyped clinical manifestations but multifactorial causes), altered response to treatment, high risk of disability, and loss of personal autonomy with all its psychological and social consequences. In addition, these key aging processes may interfere with the typical pathophysiology of specific diseases, thereby altering expected clinical manifestations and confounding diagnosis. Clinically, patients may present with obvious problems within only one of these domains, but, since systems interact, all four main domains should be evaluated

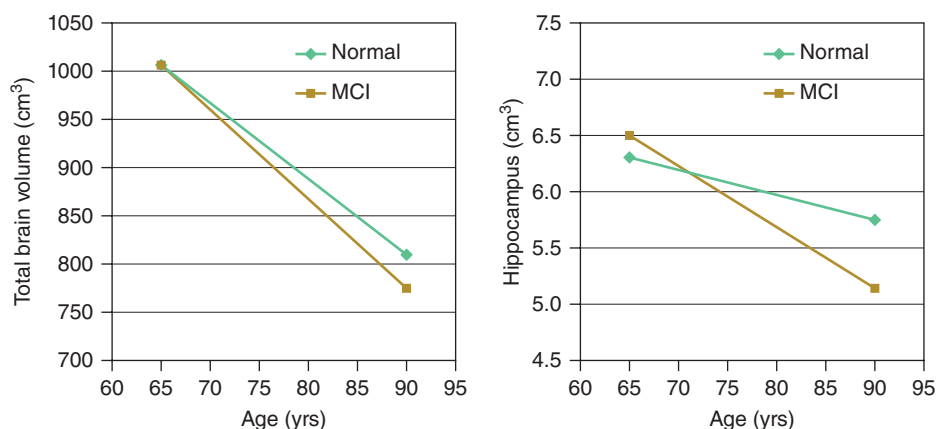


FIGURE 11-15 Longitudinal changes of regional brain volumes in normal aging and mild cognitive impairment (MCI). (From I Driscoll et al: *Neurology* 72:1906, 2009.)

and considered potential therapeutic targets. When patients present with obvious problems in multiple main systems affected by aging, they tend toward extreme degrees of susceptibility and loss of resilience, a condition that is globally referred to as *frailty*.

BIOLOGIC UNDERPINNINGS OF THE DOMAINS OF THE AGING PHENOTYPE The changes that occur with aging encompass multiple physiologic systems. Although they are often described in isolation, they are likely attributable to the progressive dysfunction of a unique mechanism that affects some fundamental housekeeping mechanism of cellular physiology. An important goal of future research is to connect the aging phenotype in humans to theories of aging that have largely been developed from studies in cell or animal models. If the main theories of aging could be operationalized into assessments that are feasible in humans, it would be possible to test the hypothesis that some of these processes are correlated with all the domains of the aging phenotype, above and beyond chronological age. Review of the biologic theories (hallmarks) of aging provides an excellent template for a working hypothesis that, at least theoretically, could be tested in longitudinal studies. Candidate mechanisms of mammalian aging include genomic instability, telomere attrition, epigenetic alterations, loss of proteostasis, deregulated nutrient sensing, mitochondrial dysfunction, cellular senescence, stem cell exhaustion, and altered intercellular communication.

Frailty Frailty has been described as a physiologic syndrome that is characterized by decreased reserve and diminished resistance to stressors, that results from cumulative decline across multiple physiologic systems, and that causes vulnerability to adverse outcomes and a high risk of death. A proposed “phenotype” definition characterized by weight loss, fatigue, impaired grip strength, diminished physical activity, and slow gait has shown good internal consistency and strong predictive validity and has been used in many clinical and epidemiologic studies. An alternative approach, the Frailty Index, assesses cumulative physiologic and functional burden. When combined with a structured clinical assessment (the Comprehensive Geriatric Assessment), the Frailty Index can be applied in clinical settings and has low rates of missing data; it predicts survival in community-dwelling older people as well as survival, length of stay and discharge location in acute-care settings. Regardless of the definition, an extensive body of literature shows that older persons who are considered frail by any definition have overt changes in the same four main processes: body composition, homeostatic dysregulation, energetic failure, and neurodegeneration—the characteristics of the aging phenotype. A classic clinical case would be an older woman with sarcopenic obesity characterized by increased body fat and decreased muscle (body composition changes); extremely low exercise tolerance and extreme fatigue (energetic failure); high insulin levels, low IGF-1 levels, inadequate intake of calories, and low levels of vitamins D and E and carotenoids (signal dysregulation); and memory problems, slow gait, and unstable balance (neurodegeneration). This woman is likely to exhibit all the manifestations of frailty, including a high risk of multiple diseases, disability, urinary incontinence, falls, delirium, depression, and other geriatric syndromes. It is expected that the biologic process underlying a particular “aging theory” would be more advanced in this woman than would be expected on the basis of chronological age.

A goal of future research in geriatric medicine that has strong potential for clinical translation is to demonstrate that the hypothetical patient described above is biologically older, according to some robust biomarkers of biologic aging, than would be estimated from chronological age alone. Conceptualizing frailty through the four main underlying processes is a step in this direction that stems from accumulated evidence and recognizes the heterogeneity and dynamic nature of the aging phenotype. Aging is universal but proceeds at highly variable rates, with wide heterogeneity in the emergence of the aging phenotype. Thus, the question is not whether an older patient is frail, but rather whether the severity of frailty is beyond the threshold of clinical and behavioral relevance. Understanding frailty through the lens of four interacting underlying processes also provides an interface with diseases that, like aging itself, affect the aging phenotype. For example, congestive heart failure is associated with low energy availability,

multiple hormonal derangements, and a proinflammatory state, thereby contributing to frailty severity. Parkinson’s disease provides an example of neurodegeneration that, in an advanced state, affects body composition, energy metabolism, and homeostatic signaling, resulting in a syndrome that closely resembles frailty. Diabetes is especially important to aging and frailty because it harms body composition, energy metabolism, homeostatic dysregulation, and neuronal integrity. Accordingly, a number of studies have found that type 2 diabetes is a strong risk factor for frailty and for many of its consequences. Since disease and aging interact, careful and appropriate treatment of disease is critical to prevent or reduce frailty.

CONSEQUENCES OF AGING PROCESSES, THE AGING PHENOTYPE, AND FRAILITY

While the pathophysiology of frailty is still being elucidated, its consequences have been well characterized in prospective studies. Four main consequences are important for clinical practice: (1) ineffective or incomplete homeostatic response to stress, (2) multiple coexisting diseases (multi- or comorbidity) and polypharmacy, (3) physical disability, and (4) the so-called geriatric syndromes. We will briefly address each one of these consequences.

Low Resistance to Stress Frailty can be considered a progressive loss of reserve in multiple physiologic functions. At an early stage and in the absence of stress, mildly frail older individuals may appear to be normal. However, they have reduced ability to cope with challenges, such as acute diseases, traumas, surgical procedures, or chemotherapy. Acute illness involving a hospital stay is associated with undernutrition and inactivity, which sometimes may be of such magnitude that the residual muscle mass fails to meet the minimal requirement for walking. Even when nutrition is reinstated, energy reserves may be insufficient to adequately rebuild muscle mass. Older persons have a reduced ability to tolerate infections, in part because they are less able than younger people to build a dynamic inflammatory response to vaccination or infectious exposure; thus, infections are more likely to become severe and systemic and to resolve more slowly. In the context of tolerance to stress, assessing aspects of frailty can help estimate the individual’s ability to withstand the rigors of aggressive treatments and to respond to interventions aimed at infection as well as the caregiver’s ability to anticipate and prevent complications of hospitalization and generally to estimate prognosis. Accordingly, treatment plans may be adjusted to improve tolerance and safety; bed rest and hospitalization should be used sparingly; and infections should be prevented, anticipated, and managed assertively.

Comorbidity and Polypharmacy Older age is associated with high rates of many chronic diseases (Fig. 11-4). Thus, not unexpectedly, the percentage of individuals affected by multiple medical conditions (co- or multimorbidity) also increases with age. In frail older individuals, comorbidity occurs at higher rates than would be expected from the combined probability of the component conditions. It is likely that frailty and comorbidity affect each other, so that multiple diseases contribute to frailty and frailty increases susceptibility to diseases.

Clinically, patients with multiple conditions present unique diagnostic and treatment challenges. Standard diagnostic criteria may not be informative because there are additional confusing signs and symptoms. A classic example is the coexistence of deficiencies in iron and vitamin B₁₂, creating an apparently normocytic anemia. The risk/benefit ratio for many medical and surgical treatment options may be reduced in the face of other diseases. Drug treatment planning is made more complex because comorbid diseases may affect the absorption, volume of distribution, protein binding, and, especially, elimination of many drugs, leading to fluctuation in therapeutic levels and increased risk of under- or overdosing. Drug excretion is affected by renal and hepatic changes with aging that may not be detectable with the usual clinical tests. Formulas for estimating glomerular filtration rate in older patients are available, whereas the estimation of changes in hepatic excretion remains a challenge.

Patients with many diseases are usually prescribed multiple drugs, especially when they are cared for by multiple specialists who do not communicate. The risk of adverse drug reactions, drug–drug interactions, and poor compliance increases geometrically with the number

of drugs prescribed and with the severity of frailty. Some general rules to minimize the chances of adverse drug events are as follows: (1) Always ask patients to bring in all medications, including prescription drugs, over-the-counter products, vitamin supplements, and herbal preparations (the “brown bag test”). (2) Screen for unnecessary drugs; those without a clear indication should be discontinued. (3) Simplify the regimen in terms of number of agents and schedules, try to avoid frequent changes, and use single-daily-dose regimens whenever possible. (4) Avoid drugs that are expensive or not covered by insurance whenever possible. (5) Minimize the number of drugs to those that are absolutely essential, and always check for possible interactions. (6) Make sure that the patient or an available caregiver understands the administered regimen, and provide legible written instructions. (7) Schedule periodic medication reviews.

Disability and Impaired Recovery from Acute-Onset Disability The prevalence of disability in self-care and home management increases steeply with aging and tends to be higher among women than among men (Fig. 11-5). Physical and cognitive function in older persons reflects overall health status and predicts health care utilization, institutionalization, and mortality more accurately than any other known biomedical measure. Thus, assessment of function and disability and prediction of the risk of disability are cornerstones of geriatric medicine. Frailty, regardless of the criteria used for its definition, is a robust and powerful risk factor for disability. Because of this strong relationship, measures of physical function and mobility have been proposed as standard criteria for frailty. However, disability occurs late in the frailty process, after reserve and compensation are exhausted. Early in the development of frailty, body composition changes, reductions in fitness, homeostatic deregulation, and neurodegeneration can begin without affecting daily function. As opposed to disability in younger persons, in which the rule is to look for a clear dominant cause, disability in frail older persons is almost always multifactorial. Multiple disrupted aging processes are usually involved, even when the precipitating cause seems unique. Excess fat mass, poor muscle strength, reduced lean body mass, poor fitness, reduced energy efficiency, poor nutritional intake, low circulating levels of antioxidant micronutrients, high levels of proinflammatory markers,

objective signs of neurologic dysfunction, and cognitive impairment all contribute to disability. The multifactorial nature of disability in frail older persons reduces the capacity for compensation and interferes with functional recovery. For example, a small lacunar stroke that causes problems with balance in a young hypertensive individual can be overcome by standing and walking with the feet further apart, a strategy that requires brain adaptation, strong muscles, and a high energy capacity. The same small lacunar stroke may cause catastrophic disability in an older person already affected by neurodegeneration and weakness, who is less able to compensate. As a consequence, interventions aimed at preventing and reducing disability in older persons should have a dual focus on both the precipitating cause and the systems needed for compensation. In the case of the lacunar stroke, interventions to promote mobility function might include stroke prevention, balance rehabilitation, and strength training.

As a rule of thumb, the assessment of contributing causes and the design of intervention strategies for disability in older persons should always consider the four main aging processes that contribute to frailty. One of the most popular approaches to disability measurement is a modification of the International Classification of Impairments, Disabilities and Handicaps (World Health Organization, 1980) proposed by the Institute of Medicine (1992). This classification infers a causal pathway in four steps: pathology (diseases), impairment (the physical manifestation of diseases), functional limitation (global functions such as walking, grasping, climbing stairs), and disability (ability to fulfill social roles in the environment). In practice, the assessment of functional limitation and disability is performed either by (1) self-reported questionnaire concerning the degree of ability to perform basic self-care or more complex ADLs or by (2) performance-based measures of physical function that assess specific domains, such as balance, gait, manual dexterity, coordination, flexibility, and endurance. A concise list of standard tools that can be used to assess physical function in older persons is provided in Table 11-4. In 2001, the WHO officially endorsed a new classification system, the International Classification of Functioning, Disability and Health, known more commonly as the ICF. In the ICF, health measures are classified from bodily, individual, and societal perspectives by means of two lists: a list of body functions and

TABLE 11-4 TOOLS FOR FUNCTIONAL ASSESSMENT IN OLDER PATIENTS

Measurement Instrument	Evaluation	Activities/Reference	Notes
Index of independence in ADLs	Self-reported	Difficulty/need for help in bathing, dressing, using toilet, transferring, continence, feeding (<i>S Katz et al: JAMA 185:914, 1963</i>)	Short and simple, but subjective
Instrumental ADLs	Self-reported	Difficulty using the telephone, using a car/public transportation, shopping, preparing meals, housework, managing medications, financial management (<i>MP Lawton et al: Psychopharmacol Bull 24:609, 1988</i>)	Short and simple; sex-biased and culturally biased items
Functional Independence Measure	Consensus multidisciplinary team	Motor (eating, grooming, bathing, dressing, toileting, managing bladder/bowels, transferring, walking, climbing stairs); cognitive (auditory comprehension, verbal expression, social interaction, problem solving, memory) (<i>RA Keith et al: Adv Clin Rehabil 1:6, 1987</i>)	Administered by trained health professionals
Barthel Index	Professionally evaluated	Independence and need for help in feeding, transferring from bed to chair and back, grooming, transferring to and from toilet, bathing, walking, climbing stairs, dressing, continence (<i>FI Mahoney et al: Md State Med J 14:61, 1965</i>)	Administered by trained health professionals
Mobility Questionnaire	Self-reported	Severe difficulty walking ¼ mile and/or climbing stairs	Short and simple
Short Physical Performance Battery	Objective performance based	Time required to walk 4 m, rise from a chair 5 times, maintain balance for 10 sec in the side-by-side, semi-tandem, and tandem positions (<i>JM Guralnik et al: J Gerontol 49:M85, 1994</i>)	Some training required
Berg Balance Scale	Objective and professionally evaluated	Performance in 14 tasks related to balance (<i>KO Berg et al: Arch Phys Med Rehabil 73:1073, 1992</i>)	Typically used by physical therapists
Walking speed	Objective performance	Measure walking speed over a 4 m course (<i>S Studenski: J Nutr Health Aging 13:878, 2009</i>)	Simple and powerful, but limited to patients who can walk
6-Minute walk	Objective performance based	Distance covered in 6 min (<i>GH Guyatt: Can Med Assoc J 132:919, 1985</i>)	Good measure of fitness, walking capacity/endurance
Long-Distance Corridor Walk (400 m)	Objective performance based	Time to fast-walk 400 m (<i>AB Newman et al: JAMA 295:2018, 2006</i>)	More challenging than the 6-min walk

Abbreviation: ADLs, activities of daily living.

structure and a list of domains of activity and participation. Since an individual's functioning and disability occur in a context, the ICF also includes a list of environmental factors. A detailed list of codes that allow the classification of body functions, activities, and participation is being developed. The ICF system is widely implemented in Europe and is gaining popularity in the United States. Whatever classification system is used, the health care provider should try to identify factors that can be modified to minimize disability. Many of these factors are discussed in this chapter. Important issues related to aging that are not addressed in this chapter but are covered elsewhere include dementia (Chap. 35) and other cognitive disorders including aphasia, memory loss, and other focal cerebral disorders (Chap. 36).

Geriatric Syndromes The term *geriatric syndrome* encompasses clinical conditions that are frequently encountered in older persons; have a deleterious effect on function and quality of life; have a multifactorial pathophysiology, often involving systems unrelated to the apparent chief symptom; and are manifested by stereotypical clinical presentations. The list of geriatric syndromes includes incontinence, delirium, falls, pressure ulcers, sleep disorders, problems with eating or feeding, pain, and depressed mood. In addition, dementia and physical disability are sometimes considered to be geriatric syndromes. The term *syndrome* is somewhat misleading in this context since it is most commonly used to describe a pattern of symptoms and signs that have a single underlying cause. The term *geriatric syndromes*, by contrast, refers to “multifactorial health conditions that occur when the accumulated effects of impairments in multiple systems render an older person vulnerable to situational challenges.” According to this definition, geriatric syndromes reflect the complex interactions between an individual's vulnerabilities and exposure to stressors or challenges. This definition aligns well with the concept that geriatric syndromes should be considered as phenotypic consequences of frailty and that a limited number of shared risk factors contribute to their etiology. Indeed, in various combinations and frequencies, virtually all geriatric syndromes are characterized by body composition changes, energy gaps, signaling disequilibria, and neurodegeneration. For example, detrusor (bladder) underactivity is a multifactorial geriatric condition that contributes to urinary retention in the frail elderly. It is characterized by detrusor muscle loss, fibrosis, and axonal degeneration. A proinflammatory state and a lack of estrogen signaling cause bladder muscle loss and detrusor underactivity, while a chronic urinary tract infection may cause detrusor hyperactivity; all of these factors may contribute to urinary incontinence.

Because of limited space, only delirium, falls, chronic pain, incontinence, and anorexia are addressed here. Interested readers are referred to textbooks on geriatric medicine for a discussion of other geriatric syndromes.

Delirium (See also Chap. 34) Delirium is an acute disorder of disturbed attention that fluctuates with time. It affects 15–55% of hospitalized older patients. Delirium has previously been considered to be transient and reversible and a normal consequence of surgery, chronic disease, or infections in older people. Delirium may be associated with a substantially increased risk for dementia and is an independent risk factor for morbidity, prolonged hospitalization, and death. These associations are particularly strong in the oldest old. Fig. 11-16 shows an algorithm for assessment and management of delirium in hospitalized older patients. The clinical presentation of delirium is heterogeneous, but frequent features are (1) a rapid decline in the level of consciousness, with difficulty focusing, shifting, or sustaining attention; (2) cognitive change (rumbling incoherent speech, memory gaps, disorientation, hallucinations) not explained by dementia; and (3) a medical history suggestive of preexisting cognitive impairment, frailty, and comorbidity. The strongest predisposing factors for delirium are dementia, any other condition associated with chronic or transient neurologic dysfunction (neurologic diseases, dehydration, alcohol consumption, psychoactive drugs), and sensory (visual and hearing) deprivation; these associations suggest that delirium is a condition of brain function susceptibility (neurodegeneration or transient neuronal impairment) that precludes the avoidance of decompensation in the face of a stressful event. Many stressful conditions have been implicated as precipitating factors, including surgery;

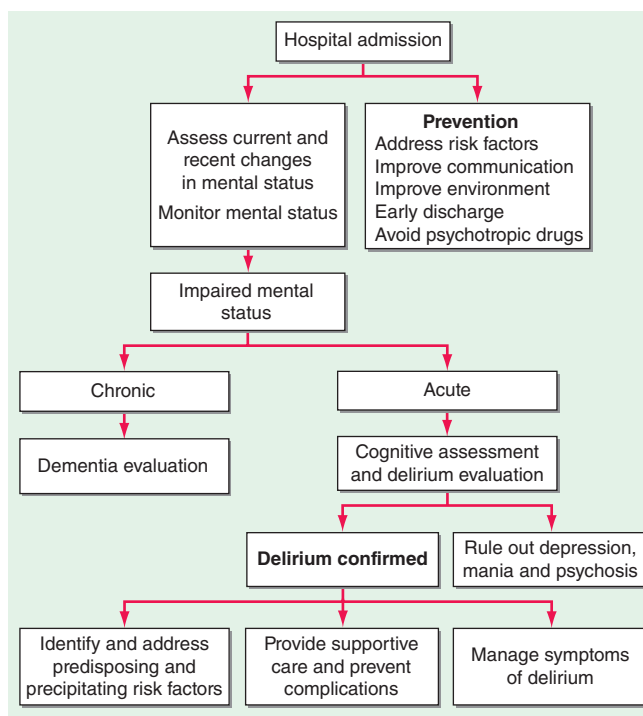


FIGURE 11-16 Algorithm depicting assessment and management of delirium in hospitalized older patients. (Modified from SK Inouye: *N Engl J Med* 354:1157, 2006.)

anesthesia; persistent pain; treatment with opiates, narcotics, or anticholinergics; sleep deprivation, immobilization; hypoxia; malnutrition; and metabolic and electrolyte derangements. Both the occurrence and the severity of delirium can be reduced by anticipatory screening and preventive strategies targeting the precipitating causes. The Confusion Assessment Method is a simple, validated tool for screening in the hospital setting. The three pillars of treatment are (1) immediate identification and treatment of precipitating factors, (2) withdrawal of drugs that may have promoted the onset of delirium, and (3) supportive care, including management of hypoxia, hydration and nutrition, mobilization, and environmental modifications. Whether patients who are cared for in special delirium units have better outcomes than those who are not is still in question. Physical restraints should be avoided because they tend to increase agitation and injury. Whenever possible, drug treatment should be avoided because it may prolong or aggravate delirium in some cases. The treatment of choice is low-dose haloperidol. It remains difficult to reduce delirium in patients with acute illness or other stressful conditions. Interventions based on dietary supplementation or careful use of pain medications and sedatives in pre- and postoperative older patients have been only partially successful.

Falls and Balance Disorders Unstable gait and falls are serious concerns in the older adult because they lead not only to injury but also to restricted activity, increased health care utilization, and even death. Like all geriatric syndromes, problems with balance and falls tend to be multifactorial and are strongly connected with the disrupted aging systems that contribute to frailty. Poor muscle strength, neural damage in the basal ganglia and cerebellum, diabetes, and peripheral neuropathy are all recognized risk factors for falls. Therefore, evaluation and management require a structured multisystem approach that spans the entire frailty spectrum and beyond. Accordingly, interventions to prevent or reduce instability and falls usually require a mix of medical, rehabilitative, and environmental modification approaches. Guidelines for the evaluation and management of falls, released by the American Geriatrics Society, recommend asking all older adults about falls and perceived gait instability (Fig. 11-17). Patients with a positive history of multiple falls as well as persons who have sustained one or more injurious falls should undergo an evaluation of gait and balance as well as a targeted history and physical examination to detect

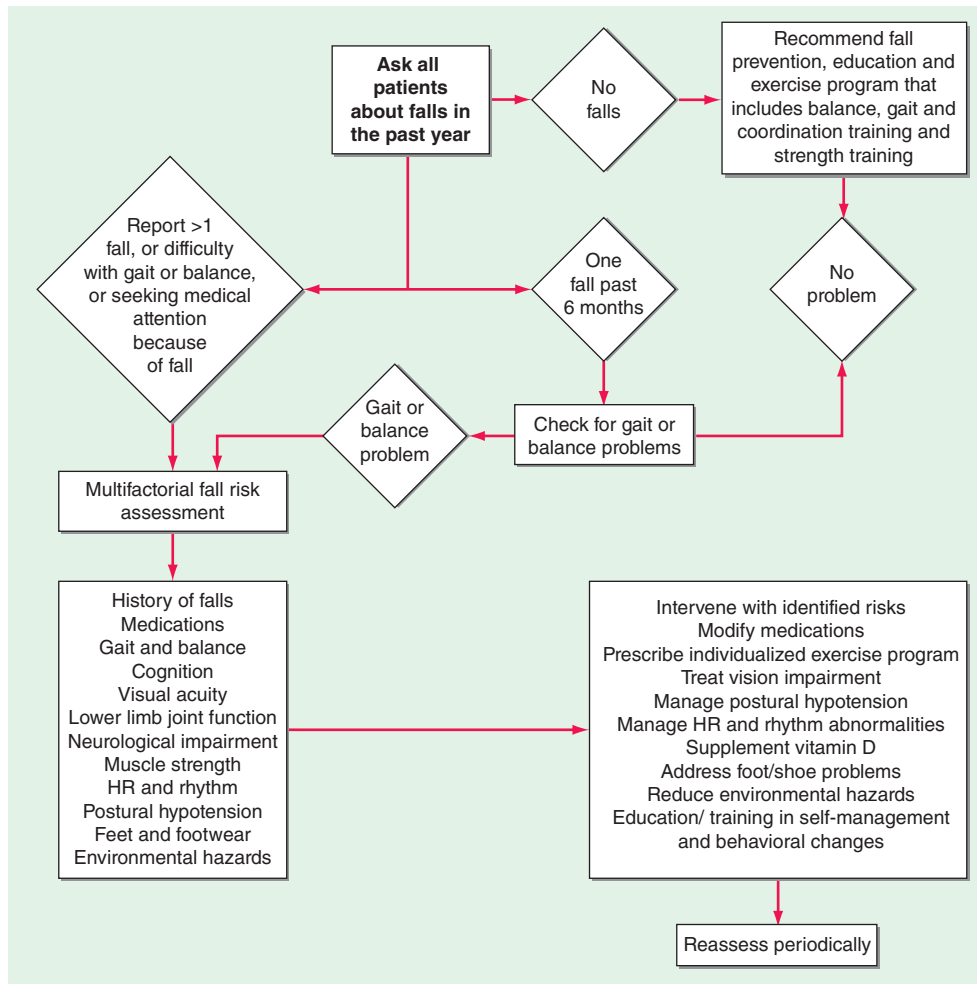


FIGURE 11-17 Algorithm depicting assessment and management of falls in older patients. HR, heart rate. (From American Geriatrics Society and British Geriatrics Society: *Clinical Practice Guideline for the Prevention of Falls in Older Persons*. New York, American Geriatric Society, 2010.)

sensory, nervous system, brain, cardiovascular, and musculoskeletal contributors. Interventions depend on the factors identified but often include medication adjustment, physical therapy, and home modifications. Meta-analyses of strategies to reduce the risk of falls have found that multifactorial risk assessment and management as well as individually targeted therapeutic exercise are effective. Supplementation with vitamin D at 800 IU daily may also help reduce falls, especially in older persons with low vitamin D levels.

Persistent Pain Pain from multiple sources is the most common symptom reported by older adults in primary care settings and is also common in acute-care, long-term-care, and palliative-care settings. Acute pain and cancer pain are beyond the scope of this chapter. Persistent pain results in restricted activity, depression, sleep disorders, and social isolation and increases the risk of adverse events due to medication. The most common causes of persistent pain are musculoskeletal problems, but neuropathic pain and ischemic pain occur frequently, and multiple concurrent causes are often found. Alterations in mechanical and structural elements of the skeleton commonly lead to secondary problems in other parts of the body, especially soft tissue or myofascial components. A structured history should elicit information about the quality, severity, and temporal patterns of pain. Physical examination should focus on the back and joints, on trigger points and periarticular areas, and on possible evidence of radicular neurologic patterns and peripheral vascular disease. Pharmacologic management should follow standard progressions, as recommended by the World Health Organization (Chap. 18), and adverse effects on the CNS, which are especially likely in this population, must be monitored. For persistent pain, regular analgesic schedules are appropriate and should

be combined with nonpharmacologic approaches such as splints, physical exercise, heat, and other modalities. A variety of adjuvant analgesics such as antidepressants and anticonvulsants may be used; again, however, effects on reaction time and alertness may be dose limiting, especially in older persons with cognitive impairment. Joint or soft tissue injections may be helpful. Education of the patient and mutually agreed-upon goal setting are important since pain usually is not fully eliminated but rather is controlled to a tolerable level that maximizes function while minimizing adverse effects.

Urinary Incontinence Urinary incontinence—the involuntary leakage of urine—is highly prevalent among older persons (especially women) and has a profound negative impact on quality of life. Approximately 50% of American women will experience some form of urinary incontinence over a lifetime. Increasing age, white race, childbirth, obesity, and medical comorbidity are all risk factors for urinary incontinence. The three main clinical forms of urinary incontinence are as follows: (1) *Stress* incontinence is the failure of the sphincteric mechanism to remain closed when there is a sudden increase in intraabdominal pressure, such as a cough or sneeze. In women this condition is due to insufficient strength of the pelvic floor muscles, while in men it is almost exclusively secondary to prostate surgery. (2) *Urge* incontinence is the loss of urine accompanied by a sudden sensation of need to urinate and inability to control it and is due to detrusor muscle overactivity (lack of inhibition) caused by loss of neurologic control or local irritation. (3) *Overflow* incontinence is characterized by urinary dribbling, either constantly or for some period after urination. This condition is due to impaired detrusor contractility (due usually to denervation, for example, in

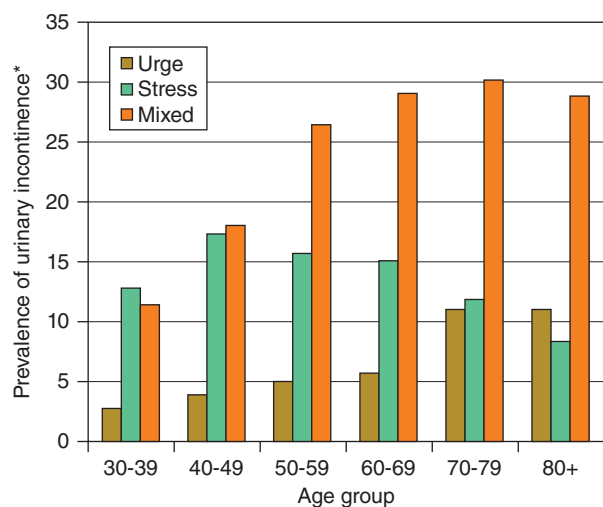


FIGURE 11-18 Rates of urge, stress, and mixed incontinence, by age group, in a sample of 3552 women. *Based on a sample of 3553 participants. (From JL Melville et al: *Arch Intern Med* 165:537, 2005.)

diabetes) or bladder outlet obstruction (prostate hypertrophy in men and cystocele in women). Thus, it is not surprising that the pathogenesis of urinary incontinence is connected to the disrupted aging systems that contribute to frailty, body composition changes (atrophy of the bladder and pelvic floor muscle), and neurodegeneration (both central and peripheral nervous systems). Frailty is a strong risk factor for urinary incontinence. Indeed, older women are more likely to have mixed (urge + stress) incontinence than any pure form (Fig. 11-18). In analogy with the other geriatric syndromes, urinary incontinence derives from a predisposing condition superimposed on a stressful precipitating factor. Accordingly, treatment of urinary incontinence should address both. The first line of treatment is bladder training associated with pelvic muscle exercise (Kegel exercises) that sometimes should be associated with electrical stimulation. Women with possible vaginal or uterine prolapse should be referred to a specialist. Urinary tract infections should be investigated and eventually treated. A long list of medications can precipitate urinary incontinence, including diuretics, antidepressants, sedative hypnotics, adrenergic agonists or blockers, anticholinergics, and calcium channel blockers. Whenever possible, these medications should be discontinued. Until recently, it was believed that oral or local estrogen treatment alleviated the symptoms of urinary incontinence in postmenopausal women, but this notion is now controversial. Antimuscarinic drugs such as tolterodine, darifenacin, and fesoterodine are modestly effective for mixed-etiology incontinence, but all of these drugs can affect cognition and so must be used with caution and with monitoring of cognitive status. In some cases, surgical treatment should be considered. Chronic catheterization has many adverse effects and should be limited to chronic urinary retention that cannot be managed in any other way. Bacteriuria always occurs and should be treated only if it is symptomatic. Bacterial communities isolated from the urine of women with urinary incontinence appear to differ with the type of incontinence; this observation suggests that the bladder microbiota may play a role in urinary incontinence. If so, this microbial population would be a potential target for treatment.

Undernutrition and Anorexia There is strong evidence that the healthy mammalian life span is greatly affected by changes in the activity of central nutrient-sensing mechanisms, especially those that involve the rapamycin (mTOR) network. Polymorphic variations in the gene that encodes mTOR in humans are associated with longevity; this association suggests that the role of nutrient signaling in healthy aging may be conserved in humans. Normal aging is associated with a decline in food intake that is more marked in men than in women. To some extent, food intake is reduced because energy demand declines as a

result of the combination of a lower level of physical activity, a decline in lean body mass, and slowed rates of protein turnover. Other contributors to decreased food intake include losses of taste sensation, reduced stomach compliance, higher circulating levels of cholecystokinin, and, in men, low testosterone levels associated with increased leptin. When food intake decreases to a level below the reduced energy demand, the result is energy malnutrition.

Malnutrition in older persons should be considered a geriatric syndrome because it is the result of intrinsic susceptibility due to aging, complicated by multiple superimposed precipitating causes. Many older individuals tend to consume a monotonous diet that lacks sufficient fresh food, fruits, and vegetables, so that intake of important micronutrients is inadequate. Undernutrition in older people is associated with multiple adverse health consequences, including impaired muscle function, decreased bone mass, immune dysfunction, anemia, reduced cognitive function, poor wound healing, delayed recovery from surgery, and increased risk of falls, disability, and death. Despite these serious potential consequences, undernutrition often remains unrecognized until it is well advanced because weight loss tends to be ignored by both patients and physicians. Muscle wasting is a frequent feature of weight loss and malnutrition that is often associated with loss of subcutaneous fat. The main causes of weight loss are anorexia, cachexia, sarcopenia, malabsorption, hypermetabolism, and dehydration, almost always in various combinations. Many of these causes can be detected and corrected. Cancer accounts for only 10–15% of cases of weight loss and anorexia in older people. Other important causes include a recent move to a long-term-care setting, acute illness (often with inflammation), hospitalization with bed rest for as little as 1–2 days, depression, drugs that cause anorexia and nausea (e.g., digoxin and antibiotics), swallowing problems, oral infections, dental problems, gastrointestinal pathology, thyroid and other hormonal problems, poverty, and isolation, with reduced access to food. Weight loss may also result from dehydration, possibly related to excess sweating, diarrhea, vomiting, or reduced fluid intake. Early identification is paramount and requires careful weight monitoring. Patients or caregivers should be taught to record weight regularly at home, the patient should be weighed at each clinical encounter, and a record of serial weights should be maintained in the medical record. If malnutrition is suspected, formal assessment should begin with a standardized screening instrument such as the Mini Nutritional Assessment, the Malnutrition Universal Screening Tool, or the Simplified Nutritional Appetite Questionnaire. The Mini Nutritional Assessment includes questions on appetite, timing of eating, frequency of meals, and taste. Its sensitivity and specificity are >75% for future weight loss of ≥5% of body weight in older people. Many nutritional supplements are available, and their use should be initiated early to prevent more severe weight loss and its consequences. When an older patient has malnutrition, the diet should be liberalized and dietary restrictions should be lifted as much as possible. Nutritional supplements should be given between meals to avoid interference with food intake at mealtime. Limited evidence supports the use of any pharmacologic intervention to treat weight loss. The two anti-anorexic drugs most often prescribed in older persons are megestrol and dronabinol. Both can increase weight; however, the gain is mostly fat, not muscle, and both drugs have serious side effects. Dronabinol is an excellent drug for use in the palliative-care setting. There is little evidence that intentional weight loss in overweight older people prolongs life. Weight loss after the age of 70 should probably be limited to persons with extreme obesity and should always be medically supervised.

HOW THE AGING PHENOTYPE AFFECTS DISEASE PRESENTATION

Common diseases in older adults may have unexpected and atypical clinical features. Most age-related changes in clinical presentation, evolution, and response to treatment are due to interaction of disease pathophysiology with age-related system dysregulation. Some diseases, such as Parkinson's disease (PD) and diabetes, directly affect aging systems and therefore have a devastating impact on frailty and its consequences.

Parkinson's Disease (See also Chap. 449) Most cases of PD begin after the age of 60 years, and the incidence increases up to the age of ~80 years. Brain aging and PD have long been thought to be related. The nigrostriatal system deteriorates with aging, and many older persons tend to develop a mild form of movement disorder characterized by bradykinesia and stooped posture that mimics mild PD. It is interesting that, in PD, older age at presentation is associated with a more severe and rapid decline in gait, balance, posture, and cognition. These age-related motor and cognitive manifestations of PD tend to be poorly responsive to levodopa or dopamine agonist treatments, especially in the oldest old. In contrast, age at presentation does not correlate with the severity and progression of other classic PD symptoms, such as tremor, rigidity, and bradykinesia, nor does it affect the response of these symptoms to levodopa. The pattern of PD features in older persons suggests that late-life PD may reflect a failure of the normal cellular compensatory mechanisms in vulnerable brain regions and that this vulnerability is increased by age-related neurodegeneration, making PD symptoms particularly resistant to levodopa treatment. In addition to motor symptoms, older PD patients tend to have reduced muscle mass (sarcopenia), eating disorders, and poor levels of fitness. Accordingly, PD is a powerful risk factor for frailty and its consequences, including disability, comorbidity, falls, incontinence, chronic pain, and delirium. Use of levodopa and dopaminergic agonists by older PD patients requires complex dosing schedules; therefore, slow-release preparations are preferred. Both dopaminergic and anticholinergic agents increase the risk of confusion and hallucinations. Use of anticholinergic agents should generally be avoided. For dopaminergic agents, cognitive side effects can be dose limiting.

Diabetes (See also Chaps. 417–419) Both the incidence and the prevalence of diabetes mellitus increase with aging. Among persons ≥ 65 years old, the prevalence is ~12% (with higher figures among African Americans and Hispanics), reflecting the effects of population aging and the obesity epidemic. Diabetes affects all four main aging systems that contribute to frailty. Obesity, especially visceral obesity, is a strong risk factor for insulin resistance, the metabolic syndrome, and diabetes. Diabetes is associated with both reduced muscle mass and accelerated rates of muscle wasting. Diabetic patients have an elevated RMR and a poor degree of fitness. Diabetes is associated with multiple hormone dysregulation, a proinflammatory state, and excess oxidative stress. Finally, diabetes-induced neurodegeneration involves both the central and peripheral nervous systems. Given these characteristics, it is not surprising that patients with diabetes mellitus are more likely to be frail and at high risk of developing physical disability, depression, delirium, cognitive impairment, urinary incontinence, injurious falls, and persistent pain. Thus, the assessment of older diabetic patients should always include screening and risk factor evaluation for these conditions.

In young and adult patients, the main treatment goal has been strict glycemic control aimed at bringing the hemoglobin A1c level to within normal values (i.e., $\leq 6\%$). However, the risk/benefit ratio is optimized by the use of less aggressive glycemic targets. In fact, in the context of a randomized clinical trial, strict glycemic control was associated with a higher mortality rate. Thus, a more reasonable goal for hemoglobin A1c is 7% or slightly below. Treatment goals are altered further in frail older adults who have a high risk of complications of hypoglycemia and a life expectancy of < 5 years. In these cases, an even less stringent target (e.g., 7–8%) should be considered, with A1c monitored every 6 or 12 months. Hypoglycemia is particularly difficult to identify in older diabetic patients because autonomic and nervous system symptoms occur at a lower blood sugar level than in younger diabetics, although the metabolic reactions and neurologic injury effects are similar in the two age groups. The autonomic symptoms of hypoglycemia are often masked by beta blockers. Frail older adults are at even higher risk for serious hypoglycemia than are healthier, higher-functioning older adults. In older patients with type 2 diabetes, a history of severe hypoglycemic episodes is associated with higher mortality risk, more severe microvascular complications, and greater risk of dementia. Thus, patients with suspected or documented episodes of hypoglycemia, especially those who are frail or disabled, need more liberal glucose-control goals, careful education about

hypoglycemia, and close follow-up by the health care provider, possibly in the presence of a caregiver. Chlorpropamide has a prolonged half-life, particularly in older adults, and should be avoided because it is associated with a high risk of hypoglycemia. Metformin should be used with caution and only in patients free of severe renal insufficiency. Renal insufficiency should be assessed by a calculated glomerular filtration rate or, in very old patients who have reduced muscle mass, by a direct measure of creatinine clearance from a 24-h urine collection. Lifestyle changes in diet and exercise and a little weight loss can prevent or delay diabetes in high-risk individuals and are substantially more effective than metformin treatment. The risk of type 2 diabetes decreased by 58% in a study of diet and exercise, and this effect was similar at all ages and in all ethnic groups. The risk reduction with standard care plus metformin was 31%.

APPROACH TO THE CARE OF OLDER PERSONS

Effects of Altered Pathophysiology and Multimorbidity on Clinical Decision-Making

The fact that older people are more likely to have atypical manifestations of disease and multiple coexisting conditions has serious consequences for the availability of high-quality evidence for medical practice and clinical decision-making. Randomized clinical trials—the basis for high-quality evidence—have tended to exclude older persons with atypical manifestations of disease, multimorbidity, or functional limitations. Across a wide range of conditions, the average age of a clinical trial participant is 20 years younger than the average age of the population with the condition. Clinical practice guidelines and care-quality metrics are focused on one condition at a time and tend not to consider the impact of comorbid conditions on the safety and feasibility of each set of recommendations. These disease-centric recommendations tend to result in fragmented care. Therefore, clinical decision-making with regard to an older person with multiple chronic conditions must be based on the weighing of several influential factors, including the patient's priorities and preferences, potential beneficial and harmful interactions among the several conditions and their treatments, life expectancy, and practical issues such as transportation, or ability to cooperate with the test or treatment.

Organization of Health Care for Older Adults The complex underlying physiology of aging leads to multiple coexisting medical problems and functional consequences that are often chronic, with recurrent exacerbations and remissions. Combined with the social consequences of aging (e.g., widowhood or lack of an available caregiver), these medical and functional factors mandate that older adults must sometimes use non-medical services to meet functional needs. The end result of these medical, functional, and social factors is that older adults use many health care and social support services in a variety of settings. Thus, it is incumbent on the internist, whether a generalist or specialist, to be familiar with the scope of settings and services that are used by their patients.

For many settings, Medicare reimbursement requires a medical order based on specific indications, so the hospitalist or referring physician must be familiar with eligibility requirements. Table 11-5 summarizes the types of services and payment sources for common settings of care. Older adults who have experienced new disability during a hospitalization are eligible for rehabilitation services. Inpatient rehabilitation requires at least 3 h per day of active rehabilitative activity and is limited to specific diagnoses. More and more rehabilitative services are provided in postacute settings, where the required intensity of service is less stringent. Postacute settings are also used for complex nursing services such as provision and supervision of long-term parenteral medication use or wound care. Under current policy, Medicare covers postacute care only if there is an eligible medical, nursing, or rehabilitation service. Otherwise, nursing home care is not covered by Medicare and must be paid for with personal assets until all resources have been consumed, at which time Medicaid coverage becomes available.

Medicaid is a state–federal partnership whose greatest single expenditure is nursing home care. Thus, the need for chronic daily assistance with personal care in a nursing home consumes a large part of most state Medicaid budgets as well as personal assets. Accordingly, alternatives to chronic nursing-home care are of great interest to states, patients, and families. Some states have developed Medicaid-funded day-care programs, sometimes based on the Program for All-Inclusive

TABLE 11-5 CARE SETTINGS FOR OLDER PATIENTS

Setting	Services	Payment Source
Hospital acute care	Medical, surgical, and psychiatric services that cannot be provided in less complex settings	Medicare, Medicaid, and private insurance
Emergency room	Resuscitation, stabilization, triage, disposition	Medicare, Medicaid, and private insurance
Inpatient rehabilitation	Hospital-based residential program providing team-based, physician-supervised, intensive therapeutic rehabilitation for specific diagnoses	Medicare, Medicaid, and private insurance
Outpatient clinic	Chronic, urgent, and preventive services	Medicare, Medicaid, and private insurance
Postacute care	Medical, nursing, and rehabilitative services after hospitalization, often based in hospitals or nursing homes	Medicare up to 100 days with eligibility requirements
Long-term care	Residential program with daily nursing and aide care for persons who are dependent in self-care	Medicaid, private payment, long-term care insurance
Assisted living	Residential program with daily aide care and housing for persons who are dependent in household management	Private payment
Home health care	Nursing and rehabilitative services for episodes of care provided to persons in the community	Medicare, Medicaid
Day programs	Supervised settings providing nursing and aide care for scheduled hours	Private payment, Medicaid

Care of the Elderly (PACE) model. In this situation, older adults who are eligible for both Medicare and Medicaid and who are otherwise eligible for chronic nursing-home care can receive coordinated medical and functional services in conjunction with a day-care program.

For most older adults, a caregiver must be available to provide assistance on weeknights and weekends. Under current policy, home health services do not provide chronic functional assistance in the home but rather are targeted at episodes of care supplied by medical or rehabilitative services for older adults who are considered home bound. Some community agencies, whether private or public, can provide homemaker and home aide services to assist the home-bound older adult with functional needs, but there may be income requirements or expensive private payment may be needed.

Within the past decade, there has been tremendous growth in a broad spectrum of assisted-living settings. Such settings do not offer the degree of 24-h nursing supervision or personal aide care that is provided in traditional nursing homes, although distinctions are becoming blurred. Most assisted-living settings provide meals, medication supervision, and homemaking services, but they often require that residents be capable of transporting themselves to a congregate meal site. Moreover, most of these settings accept only private payment from residents and their families and thus are hard to access for older adults with limited resources. Some states are exploring coverage for lower-cost residential-care services such as family care homes.

Models of Care Coordination The complexity and fragmentation of care for older adults results in both increased costs and increased risk of iatrogenic complications such as missed diagnoses, adverse medication events, further worsening of function, and even death. These serious consequences have led to a strong interest in care coordination through teams of providers, with the goals to reduce unnecessary costs and to prevent adverse events. [Table 11-6](#) lists examples of evidence-based models of care coordination that were recommended in a 2009

TABLE 11-6 EVIDENCE-BASED MODELS OF CARE COORDINATION FOR OLDER PATIENTS (INSTITUTE OF MEDICINE, 2009)

Model	Team Members	Services
Interdisciplinary primary care "medical home"	Primary care physician plus social worker, nurse, nurse practitioner, or other care coordinators	Coordination of medical and social needs across settings
Case management	Nurse or social worker	Education of and information dissemination to patients and families; in some cases, communication with providers and settings
Disease management	Nurse	Health education and follow-up support for specific chronic diseases
Preventive home visits	Physician, nurse, social worker, and others	Structured assessment of physical, mental, functional, and social status in the home setting, with recommendations for care and prevention
Comprehensive outpatient geriatric assessment and management	Physician, nurse, social worker, and sometimes others (e.g., pharmacist, rehabilitation therapist, psychologist)	Structured assessment of physical, mental, functional, and social status in the outpatient setting, with recommendations for care and prevention. Some programs also take responsibility for implementing the recommendations.
Pharmaceutical care management	Pharmacist	Review and recommendations, in any setting, regarding the total medication regimen
Chronic disease self-management	Nurse, health educator, or other health professional	Health education and coaching for specific chronic conditions
Prevention rehabilitation	Rehabilitation therapist	Anticipatory evaluation, therapeutic exercise, and assistive technology in the home or outpatient setting for older adults with physical disability
Caregiver services	Social worker, psychologist, or other health professional	Education, counseling, and resource referral for caregivers of older adults with chronic functional and mental health problems
Hospital discharge/transition coordination	Nurse, nurse practitioner	Care planning and education for patient and family before and after hospital discharge
Hospital at home	Physician, nurse, pharmacist	Diagnostic testing and medical treatments that can replace hospitalization or reduce length of hospital stay for target conditions
Nursing home care coordination	Nurse practitioner or physician assistant	Scheduled assessment and care planning as well as education for health workers in chronic-care settings
Hospital delirium comprehensive care	Physician, nurse	Prevention, screening, and management of delirium in the hospital setting
Comprehensive inpatient geriatric assessment and management	Physician, nurse, social worker, and sometimes others (e.g., pharmacist, rehabilitation therapist, psychologist)	Specialized inpatient settings such as acute care of the elderly (ACE) units or roving multidisciplinary teams that provide evaluation and recommendations for medical, mental health, functional, and social needs. ACE units and some teams take responsibility for implementation of recommendations.

Source: Reproduced with permission from C Boulton et al: *J Am Geriatr Soc* 57:2328, 2009.

Institute of Medicine report. While not mentioned as a specific type of team care, modern information technology offers substantial promise in providing consistent, readily available information across settings and providers. All such team programs are targeted at prevention and management of chronic and complex problems. Evidence from clinical trials or quasi-experimental studies supports the benefit of each model, and for some models data are sufficient to support meta-analyses. The evidence for benefit is not always consistent between studies or types of care but includes some support for improved quality of care, quality of life, function, survival, and health care costs and use. Some models of care are disease-specific and focus on common chronic conditions such as diabetes mellitus, congestive heart failure, chronic obstructive pulmonary disease, and stroke. One challenge in the use of these models is that a majority of older adults will have multiple simultaneous conditions and thus will need services from multiple programs that may not communicate among themselves.

Most models of care are difficult to implement in today's health care system because nonphysician services are not reimbursed, nor is physician effort that is not incorporated into "face-to-face" time. Thus, several models have been developed largely by the Department of Veterans Affairs Health Care System, Medicare Managed Care providers, and other sponsoring agencies. Medicare has developed a series of demonstration projects that can expand the evidence base and serve policy makers. More recently, there has been an effort to promote coordinated care through Accountable Care Organizations and patient-centered "medical homes." However, the processes and outcomes of such care must evolve from disease-specific indicators to more general markers, such as optimizing functional status, focusing on outcomes that are important to patients, and minimizing inappropriate care.

SCREENING AND PREVENTION IN OLDER PERSONS

In older adults, prevention tests and interventions are less consistently recommended for all asymptomatic patients. The guidelines fail to address the influence of health status and life expectancy on recommendations, although the benefits of prevention are clearly affected by life expectancy. For example, in most types of cancer, screening provides no benefit in patients with a life expectancy of ≤ 5 years. More research is needed to build an appropriate evidence base for age- and life expectancy-adapted preventive services. Health behavior modification, especially increasing physical activity and improving nutrition, probably has the greatest potential to promote healthy aging.

Screening Tests

- Osteoporosis: Bone mineral density (BMD) should be measured at least once after the age of 65 years. There is little evidence that regular monitoring of BMD improves the prediction of fractures. Because of limitations in the precision of dual-energy x-ray absorptiometry, the minimal interval between evaluations should be 2–3 years.
- Hypertension: Blood pressure should be determined at least once a year or more often in patients with hypertension.
- Diabetes: Serum glucose and hemoglobin A1c should be checked every 3 years or more often in patients who are obese or hypertensive.
- Lipid disorders: A lipid panel should be done every 5 years or more often in patients with diabetes or any cardiovascular disease.
- Colorectal cancer: A fecal occult blood test and a sigmoidoscopy or colonoscopy should be done on a regular schedule up to the age of 75 years. No consensus guidelines exist for these tests >75 years of age.
- Breast cancer: Mammography should be done every 2 years between the ages of 50 and 74 years. No consensus guidelines exist for mammography after the age of 75 years.
- Cervical cancer: A Pap smear should be done every 3 years up to the age of 65 years.

Preventive Interventions

- Influenza: Immunize annually.
- Shingles: Administer herpes zoster vaccine once after the age of 50 years.
- Pneumonia: Administer pneumococcal vaccine once at the age of 65 years.

- Myocardial infarction: Prescribe daily aspirin for patients with prevalent cardiovascular disease or with a poor cardiovascular risk profile.
- Osteoporosis: Prescribe calcium at 1200 mg daily and vitamin D at ≥ 800 IU daily.

Exercise Rates of regular physical activity decrease with age and are lowest in older persons. This situation is unfortunate because increased physical activity has clear benefits in older adults, improving physical function, muscle strength, mood, sleep, and metabolic risk profile. Some studies suggest that exercise can improve cognition and prevent dementia, but this association is still controversial. Exercise programs, both aerobic and strength training, are feasible and beneficial even in very old and frail individuals. Regular, moderate-intensity exercise can reduce the rate of age-associated decline in physical function. The Centers for Disease Control and Prevention recommends that older persons should spend at least 150 min per week in moderate-intensity aerobic activity (e.g., brisk walking) and should engage in muscle-strengthening activities that work all major muscle groups (legs, hips, back, abdomen, chest, shoulders, and arms) at least 2 days a week. In the absence of contraindications, more intense and prolonged physical activity provides greater benefits. Frail and sedentary persons may need supervision, at least at the start of the exercise program, to avoid falls and exercise-related injuries.

Nutrition Older persons are particularly vulnerable to malnutrition, and many problems that affect older patients can be addressed by dietary modification. As mentioned above, nutrient sensing is the major factor associated with differential longevity in several animal models, including mammals. Treatment with rapamycin, the only pharmacologic intervention that has been associated with longevity, affects nutrient sensing. Nevertheless, there are almost no evidence-based guidelines for individualizing dietary modifications based on differing health outcomes in the elderly. Even when guidelines exist, older people tend to be poorly compliant with dietary recommendations. Basic principles of a healthy diet that are also valid for older persons are as follows:

- Encourage the consumption of fruits and vegetables; they are rich in micronutrients, mineral, and fibers. Whole grains are also a good source of fiber. Keep in mind that some of these foods are costly and thus less accessible to low-income persons.
- Emphasize that good hydration is essential. Fluid intake should be at least 1000 mL daily.
- Encourage the use of fat-free and low-fat dairy products, legumes, poultry, and lean meats. Encourage consumption of fish at least once a week, since there is strong epidemiologic evidence that fish consumption is associated with a lowered risk of Alzheimer's disease.
- Match intake of energy (calories) to overall energy needs in order to maintain a healthy weight and BMI (20–27). Recommend moderate (5–10%) caloric restriction only when the BMI is >27 .
- Limit consumption of foods with high caloric density, high sugar content, and high salt content.
- Limit the intake of foods with a high content of saturated fatty acids and cholesterol.
- Limit alcohol consumption (one drink per day or less).
- Introduce vitamin D–fortified foods and/or vitamin D supplements into the diet. Older persons who have little exposure to UVB radiation are at risk of vitamin D insufficiency.
- Make sure that the diet includes adequate food-related intake of magnesium, vitamin A, and vitamin B₁₂.
- Monitor daily protein intake, which, in healthy older persons, should be in the range of 1.0–1.2 g/kg of body weight. Higher daily protein intake (i.e., ≥ 1.2 –1.5 g/kg) is advised for those who are exercising or are affected by chronic diseases, especially if these conditions are associated with chronic inflammation. Older people with severe kidney disease (i.e., an estimated glomerular filtration rate of <30 mL/min per 1.73 m²) who are not on dialysis should limit protein intake.

- For constipation, increase dietary fiber intake to 10–25 g/d and fluid intake to 1500 mL/d. A bulk laxative (methylcellulose or psyllium) can be added.

Novel Interventions to Modify Aging Processes Aging is a complex process with multiple manifestations at the molecular, cellular, organ, and whole-organism level. The nature of the aging process is still not fully understood, but aging and its effects may be modulated by appropriate interventions. Dietary and genetic alterations can increase healthy life span and prevent the development of dysregulated systems and the aging phenotype in laboratory model organisms. The mechanisms responsible for life span expansion are “food” sensors typically activated in situations of food shortage, such as IGF/insulin and the TOR (target of rapamycin) pathways. Accordingly, a reduction in food intake without malnutrition extends the life span by 10–50% in diverse organisms, from yeasts to rhesus monkeys. Mechanisms that mediate the effects of caloric restriction are under intensive study because they are potential targets for interventions aimed at counteracting the emergence of the aging phenotype and its deleterious effects in humans. For example, resveratrol, a natural compound found in grape skin that mimics some of the effects of dietary restriction, increases longevity and improves health in mice fed a high-fat diet but has little effect on mice fed a standard diet. Other compounds that potentially mimic caloric restriction are being developed and tested. A high prevalence of IGF-1 receptor gene mutation has been found in Ashkenazi Jewish centenarians and in other long-lived individuals, suggesting that the downregulation of IGF-1 signaling may promote human longevity. A 20-year period of 30% dietary restriction applied to adult rhesus monkeys was associated with reduced cardiovascular and cancer morbidity, reduced signs of aging, and greater longevity, although a second such study did not find increased longevity. In humans, dietary restriction is effective against obesity and reduces insulin resistance, inflammation, blood pressure, CRP level, and intima-media thickness of the carotid arteries. However, the beneficial effects of dietary restriction in humans are still controversial, and some potential negative effects have not been sufficiently studied. An interesting effect of caloric restriction in humans is mitochondrial biogenesis. Mitochondrial dysfunction has emerged as a potentially important underlying contributor to aging. Reduced expression of mitochondrial genes is a strongly conserved feature of aging across different species. Mitochondria are the machinery for chemical energy production, and brain and muscle are particularly susceptible to defective mitochondrial function. Thus, declining mitochondrial function may be a direct cause of at least three of the main dysregulated systems contributing to the phenotype of aging.

OTHER ASPECTS OF AGING

This chapter has touched on some of the fundamental aspects of human aging, focusing mostly on those that are relevant to the care of older patients. Many aspects of geriatric medicine have not been addressed because of space limitations. Valuable topics not considered include details of comprehensive geriatric assessment, depression and anxiety, hypertension, orthostatic hypotension, dementia, vision and hearing impairment, osteoporosis, palliative care, prostate disorders, foot problems, and women’s health. Some of these topics are treated extensively elsewhere in this text, sometimes with comments on age-specific issues.

CONCLUSIONS

The universal process of aging is becoming better understood. There appear to be shared underlying cellular and molecular processes that induce widespread dysregulation in key systems. This dysregulation contributes to clinical manifestations of a frailty phenotype and can be used to understand how to evaluate and manage the older patient.

ACKNOWLEDGMENT

We would like to thank our colleagues who provided criticisms and suggestions for improvement of this chapter. We are particularly indebted to Dr. John Morley for his valuable suggestions regarding the section on undernutrition and anorexia.

This page intentionally left blank

12e The Safety and Quality of Health Care

David W. Bates

Safety and quality are two of the central dimensions of health care. In recent years it has become easier to measure safety and quality, and it is increasingly clear that performance in both dimensions could be much better. The public is—with good justification—demanding measurement and accountability, and payment for services will increasingly be based on performance in these areas. Thus, physicians must learn about these two domains, how they can be improved, and the relative strengths and limitations of the current ability to measure them.

Safety and quality are closely related but do not completely overlap. The Institute of Medicine has suggested in a seminal series of reports that safety is the first part of quality and that the health care system must first and foremost guarantee that it will deliver safe care, although quality is also pivotal. In the end, it is likely that more net clinical benefit will be derived from improving quality than from improving safety, though both are important and safety is in many ways more tangible to the public. The first section of this chapter will address issues relating to the safety of care and the second will cover quality of care.

SAFETY IN HEALTH CARE

Safety Theory and Systems Theory *Safety theory* clearly points out that individuals make errors all the time. Think of driving home from the hospital: you intend to stop and pick up a quart of milk on the way home but find yourself entering your driveway without realizing how you got there. Everybody uses low-level, semiautomatic behavior for many activities in daily life; this kind of error is called a *slip*. Slips occur often during care delivery—e.g., when people intend to write an order but forget because they have to complete another action first. *Mistakes*, by contrast, are errors of a higher level; they occur in new or nonstereotypic situations in which conscious decisions are being made. An example would be dosing of a medication with which a physician is not familiar. The strategies used to prevent slips and mistakes are often different.

Systems theory suggests that most accidents occur as the result of a series of small failures that happen to line up in an individual instance so that an accident can occur (Fig. 12e-1). It also suggests that most individuals in an industry such as health care are trying to do the right thing (e.g., deliver safe care) and that most accidents thus can be seen as resulting from defects in systems. Systems should be designed both to make errors less likely and to identify those that do inevitably occur.

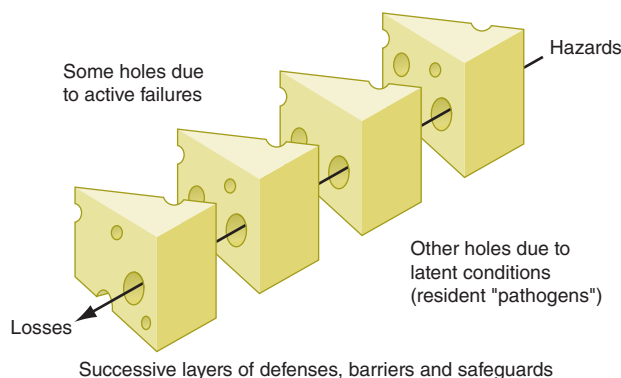


FIGURE 12e-1 “Swiss cheese” diagram. Reason argues that most accidents occur when a series of “latent failures” are present in a system and happen to line up in a given instance, resulting in an accident. Examples of latent failures in the case of a fall might be that the unit is unusually busy and the floor happens to be wet. (Adapted from *J Reason: BMJ* 320:768, 2000; with permission.)

Factors that Increase the Likelihood of Errors Many factors ubiquitous in health care systems can increase the likelihood of errors, including fatigue, stress, interruptions, complexity, and transitions. The effects of fatigue in other industries are clear, but its effects in health care have been more controversial until recently. For example, the accident rate among truck drivers increases dramatically if they work over a certain number of hours in a week, especially with prolonged shifts. A recent study of house officers in the intensive care unit demonstrated that they were about one-third more likely to make errors when they were on a 24-h shift than when they were on a schedule that allowed them to sleep 8 h the previous night. The American College of Graduate Medical Education has moved to address this issue by putting in place the 80-h workweek. Although this stipulation is a step forward, it does not address the most important cause of fatigue-related errors: extended-duty shifts. High levels of stress and heavy workloads also can increase error rates. Thus, in extremely high-pressure situations, such as cardiac arrests, errors are more likely to occur. Strategies such as using protocols in these settings can be helpful, as can simple recognition that the situation is stressful.

Interruptions also increase the likelihood of error and occur frequently in health care delivery. It is common to forget to complete an action when one is interrupted partway through it by a page, for example. Approaches that may be helpful in this area include minimizing interruptions and setting up tools that help define the urgency of an interruption.

Complexity represents a key issue that contributes to errors. Providers are confronted by streams of data (e.g., laboratory tests and vital signs), many of which provide little useful information but some of which are important and require action or suggest a specific diagnosis. Tools that emphasize specific abnormalities or combinations of abnormalities may be helpful in this area.

Transitions between providers and settings are also common in health care, especially with the advent of the 80-h workweek, and generally represent points of vulnerability. Tools that provide structure in exchanging information—for example, when transferring care between providers—may be helpful.

The Frequency of Adverse Events in Health Care Most large studies focusing on the frequency and consequences of adverse events have been performed in the inpatient setting; some data are available for nursing homes, but much less information is available about the outpatient setting. The Harvard Medical Practice Study, one of the largest studies to address this issue, was performed with hospitalized patients in New York. The primary outcome was the adverse event: an injury caused by medical management rather than by the patient’s underlying disease. In this study, an event either resulted in death or disability at discharge or prolonged the length of hospital stay by at least 2 days. Key findings were that the adverse event rate was 3.7% and that 58% of the adverse events were considered preventable. Although New York is not representative of the United States as a whole, the study was replicated later in Colorado and Utah, where the rates were essentially similar. Since then, other studies using analogous methodologies have been performed in various developed nations, and the rates of adverse events in these countries appear to be ~10%. Rates of safety issues appear to be even higher in developing and transitional countries; thus, this is clearly an issue of global proportions. The World Health Organization has focused on this area, forming the World Alliance for Patient Safety.

In the Harvard Medical Practice Study, adverse drug events (ADEs) were most common, accounting for 19% of all adverse events, and were followed in frequency by wound infections (14%) and technical complications (13%). Almost half of adverse events were associated with a surgical procedure. Among nonoperative events, 37% were ADEs, 15% were diagnostic mishaps, 14% were therapeutic mishaps, 13% were procedure-related mishaps, and 5% were falls.

ADEs have been studied more than any other error category. Studies focusing specifically on ADEs have found that they appear to be much more common than was suggested by the Harvard Medical Practice Study, although most other studies use more inclusive criteria. Detection approaches in the research setting include chart review and

the use of a computerized ADE monitor, a tool that explores the database and identifies signals that suggest an ADE may have occurred. Studies that use multiple approaches find more ADEs than does any individual approach, and this discrepancy suggests that the true underlying rate in the population is higher than would be identified by a single approach. About 6–10% of patients admitted to U.S. hospitals experience an ADE.

Injuries caused by drugs are also common in the outpatient setting. One study found a rate of 21 ADEs per every 100 patients per year when patients were called to assess whether they had had a problem with one of their medications. The severity level was lower than in the inpatient setting, but approximately one-third of these ADEs were preventable.

The period immediately after a patient is discharged from the hospital appears to be very risky. A recent study of patients hospitalized on a medical service found an adverse event rate of 19%; about one-third of those events were preventable, and another one-third were ameliorable (i.e., they could have been made less severe). ADEs were the single leading error category.

Prevention Strategies Most work on strategies to prevent adverse events has targeted specific types of events in the inpatient setting, with nosocomial infections and ADEs having received the most attention. Nosocomial infection rates have been reduced greatly in intensive care settings, especially through the use of checklists. For ADEs, several strategies have been found to reduce the medication error rate, although it has been harder to demonstrate that they reduce the ADE rate overall, and no studies with adequate power to show a clinically meaningful reduction have been published.

Implementation of checklists to ensure that specific actions are carried out has had a major impact on rates of catheter-associated bloodstream infection and ventilator-associated pneumonia, two of the most serious complications occurring in intensive care units. The checklist concept is based on the premise that several specific actions can reduce the frequency of these issues; when these actions are all taken for every patient, the result has been an extreme reduction in the frequency of the associated complication. These practices have been disseminated across wide areas, in particular in the state of Michigan.

Computerized physician order entry (CPOE) linked with clinical decision support reduces the rate of serious medication errors, defined as those that harm someone or have the potential to do so. In one study, CPOE, even with limited decision support, decreased the serious medication error rate by 55%. CPOE can prevent medication errors by suggesting a default dose, ensuring that all orders are complete (e.g., that they include dose, route, and frequency), and checking orders for allergies, drug–drug interactions, and drug–laboratory issues. In addition, clinical decision support can suggest the right dose for a patient, tailoring it to level of renal function and age. In one study, patients with renal insufficiency received the appropriate dose only one-third of the time without decision support, whereas that fraction increased to approximately two-thirds with decision support; moreover, with such support, patients with renal insufficiency were discharged from the hospital half a day earlier. As of 2009, only ~15% of U.S. hospitals had implemented CPOE, but many plan to do so and will receive major financial incentives for achieving this goal.

Another technology that can improve medication safety is bar coding linked with an electronic medication administration record. Bar coding can help ensure that the right patient gets the right medication at the right time. Electronic medication administration records can make it much easier to determine what medications a patient has received. Studies to assess the impact of bar coding on medication safety are under way, and the early results are promising. Another technology to improve medication safety is “smart pumps.” These pumps can be set according to which medication is being given and at what dose; the health care professional will receive a warning if too high a dose is about to be administered.

The National Safety Picture Several organizations, including the National Quality Forum and the Joint Commission, have made recommendations for improving safety. In particular, the National Quality

Forum has released recommendations to U.S. hospitals about what practices will most improve the safety of care, and all hospitals are expected to implement these recommendations. Many of these practices arise frequently in routine care. One example is “readback,” the practice of recording all verbal orders and immediately reading them back to the physician to verify the accuracy of what was heard. Another is the consistent use of standard abbreviations and dose designations; some abbreviations and dose designations are particularly prone to error (e.g., 7U may be read as 70).

Measurement of Safety Measuring the safety of care is difficult and expensive, since adverse events are, fortunately, rare. Most hospitals rely on spontaneous reporting to identify errors and adverse events, but the sensitivity of this approach is very low, with only ~1 in 20 ADEs reported. Promising research techniques involve searching the electronic record for signals suggesting that an adverse event has occurred. These methods are not yet in wide use but will probably be used routinely in the future. Claims data have been used to identify the frequency of adverse events; this approach works much better for surgical care than for medical care and requires additional validation. The net result is that, except for a few specific types of events (e.g., falls and nosocomial infections), hospitals have little idea about the true frequency of safety issues.

Nonetheless, all providers have the responsibility to report problems with safety as they are identified. All hospitals have spontaneous reporting systems, and, if providers report events as they occur, those events can serve as lessons for subsequent improvement.

Conclusions about Safety It is abundantly clear that the safety of health care can be improved substantially. As more areas are studied closely, more problems are identified. Much more is known about the epidemiology of safety in the inpatient setting than in outpatient settings. A number of effective strategies for improving inpatient safety have been identified and are increasingly being applied. Some effective strategies are also available for the outpatient setting. Transitions appear to be especially risky. The solutions to improving care often entails the consistent use of systematic techniques such as checklists and often involves leveraging of information technology. Nevertheless, solutions will also include many other domains, such as human factors techniques, team training, and a culture of safety.

QUALITY IN HEALTH CARE

Assessment of quality of care has remained somewhat elusive, although the tools for this purpose have increasingly improved. Selection of health care and measurement of its quality are components of a complex process.

Quality Theory Donabedian has suggested that quality of care can be categorized by type of measurement into structure, process, and outcome. *Structure* refers to whether a particular characteristic is applicable in a particular setting—e.g., whether a hospital has a catheterization laboratory or whether a clinic uses an electronic health record. *Process* refers to the way care is delivered; examples of process measures are whether a Pap smear was performed at the recommended interval or whether an aspirin was given to a patient with a suspected myocardial infarction. *Outcome* refers to what actually happens—e.g., the mortality rate in myocardial infarction. It is important to note that good structure and process do not always result in a good outcome. For instance, a patient may present with a suspected myocardial infarction to an institution with a catheterization laboratory and receive recommended care, including aspirin, but still die because of the infarction.

Quality theory also suggests that overall quality will be improved more in the aggregate if the performance level of all providers is raised rather than if a few poor performers are identified and punished. This view suggests that systems changes are especially likely to be helpful in improving quality, since large numbers of providers may be affected simultaneously.

The theory of *continuous quality improvement* suggests that organizations should be evaluating the care they deliver on an ongoing basis and continually making small changes to improve their individual processes. This approach can be very powerful if embraced over time.

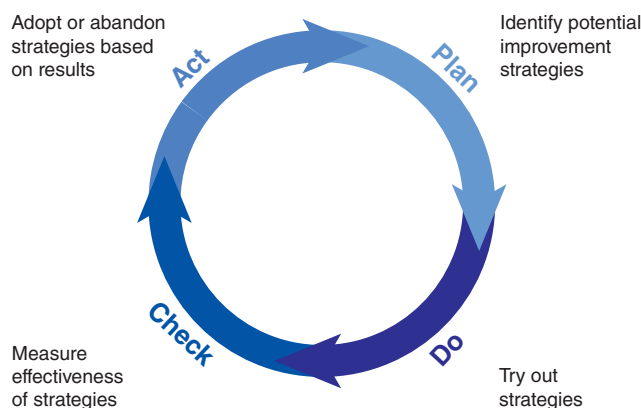


FIGURE 12e-2 Plan-Do-Check-Act cycle. This approach can be used to improve a specific process rapidly. First, planning is undertaken, and several potential improvement strategies are identified. Next, these strategies are evaluated in small “tests of change.” “Checking” entails measuring whether the strategies have appeared to make a difference, and “acting” refers to acting on the results.

A number of specific tools have been developed to help improve process performance. One of the most important is the Plan-Do-Check-Act cycle (Fig. 12e-2). This approach can be used for “rapid cycle” improvement of a process—e.g., the time that elapses between a diagnosis of pneumonia and administration of antibiotics to the patient. Specific statistical tools, such as control charts, are often used in conjunction to determine whether progress is being made. Because most medical care includes one or many processes, this tool is especially important for improvement.

Factors Relating to Quality Many factors can decrease the level of quality, including stress to providers, high or low levels of production pressure, and poor systems. Stress can have an adverse effect on quality because it can lead providers to omit important steps, as can a high level of production pressure. Low levels of production pressure sometimes can result in worse quality, as providers may be bored or have little experience with a specific problem. Poor systems can have a tremendous impact on quality, and even extremely dedicated providers typically cannot achieve high levels of performance if they are operating within a poor system.

Data about the Current State of Quality A study published by the RAND Corporation in 2006 provided the most complete picture of quality of care delivered in the United States to date. The results were sobering. The authors found that, across a wide range of quality parameters, patients in the United States received only 55% of recommended care overall; there was little variation by subtype, with scores of 54% for preventive care, 54% for acute care, and 56% for care of chronic conditions. The authors concluded that, in broad terms, the chances of getting high-quality care in the United States were little better than those of winning a coin flip.

Work from the Dartmouth Atlas of Health Care evaluating geographic variation in use and quality of care demonstrates that, despite large variations in utilization, there is no positive correlation between the two variables at the regional level. An array of data demonstrate, however, that providers with larger volumes for specific conditions, especially for surgical conditions, do have better outcomes.

Strategies for Improving Quality and Performance A number of specific strategies can be used to improve quality at the individual level, including rationing, education, feedback, incentives, and penalties. *Rationing* has been effective in some specific areas, such as persuading physicians to prescribe within a formulary, but it generally has been resisted. *Education* is effective in the short run and is necessary for changing opinions, but its effect decays fairly rapidly with time. *Feedback* on performance can be given at either the group or the individual level. Feedback is most effective if it is individualized and is given in close temporal proximity to the original events. *Incentives* can be effective,

and many believe that they will prove to be a key to improving quality, especially if pay-for-performance with sufficient incentives is broadly implemented (see below). *Penalties* produce provider resentment and are rarely used in health care.

Another set of strategies for improving quality involves changing the systems of care. An example would be introducing reminders about which specific actions needed to be taken at a visit for a specific patient—a strategy that has been demonstrated to improve performance in certain situations, such as the delivery of preventive services. Another approach that has been effective is the development of “bundles” or groups of quality measures that can be implemented together with a high degree of fidelity. A number of hospitals have implemented a bundle for ventilator-associated pneumonia in the intensive care unit that includes five measures (e.g., ensuring that the head of the bed is elevated). These hospitals have been able to improve performance substantially.

Perhaps the most pressing need is to improve the quality of care for chronic diseases. The Chronic Care Model has been developed by Wagner and colleagues (Fig. 12e-3); it suggests that a combination of strategies is necessary (including self-management support, changes in delivery system design, decision support, and information systems) and that these strategies must be delivered by a practice team composed of several providers, not just a physician.

Available evidence about the relative efficacy of strategies in reducing hemoglobin A1c (HbA1c) in outpatient diabetes care supports this general premise. It is especially notable that the outcome was the HbA1c level, as it has generally been much more difficult to improve outcome measures than process measures (such as whether HbA1c was measured). In this meta-analysis, a variety of strategies were effective, but the most effective ones were the use of team changes and the use of a case manager. When cost-effectiveness is considered in addition, it appears likely that an amalgam of strategies will be needed. However, the more expensive strategies, such as the use of case managers, probably will be implemented widely only if pay-for-performance takes hold.

National State of Quality Measurement In the inpatient setting, quality measurement is now being performed by a very large proportion of hospitals for several conditions, including myocardial infarction, congestive heart failure, pneumonia, and surgical infection prevention; 20 measures are included in all. This is the result of the Hospital Quality Initiative, which represents a collaboration among many entities,



FIGURE 12e-3 The Chronic Care Model, which focuses on improving care for chronic diseases, suggests that (1) delivery of high-quality care requires a range of strategies that must closely involve and engage the patient and (2) team care is essential. (From EH Wagner et al: *Eff Clin Pract* 1:2, 1998.)

including the Hospital Quality Alliance, the Joint Commission, the National Quality Forum, and the Agency for Healthcare Research and Quality. The data are housed at the Center for Medicare and Medicaid Services, which publicly releases performance data on the measures on a website called *Hospital Compare* (www.cms.gov/Medicare/Quality-Initiatives-Patient-Assessment-Instruments/HospitalQualityInits/HospitalCompare.html). These data are reported voluntarily and are available for a very high proportion of the nation's hospitals. Analyses demonstrate substantial regional variation in quality and important differences among hospitals. Analyses by the Joint Commission for similar indicators reveal that performance on measures by hospitals has improved over time and that, as might be hoped, lower performers have improved more than higher performers.

Public Reporting Overall, public reporting of quality data is becoming increasingly common. There are now commercial websites that have quality-related data for most regions of the United States, and these data can be accessed for a fee. Similarly, national data for hospitals are available. The evidence to date indicates that patients have not made much use of such data but that the data have had an important effect on provider and organization behavior. Instead, patients have relied on provider reputation to make choices, partly because little information was available until very recently and the information that was available was not necessarily presented in ways that were easy for patients to access. Many authorities think that, as more information about quality becomes available, it will become increasingly central to patients' choices about where to access care.

Pay-for-Performance Currently, providers in the United States get paid exactly the same amount for a specific service, regardless of the quality of care delivered. The pay-for-performance theory suggests that, if providers are paid more for higher-quality care, they will invest in strategies that enable them to deliver that care. The current key issues in the pay-for-performance debate relate to (1) how effective it is, (2) what levels of incentives are needed, and (3) what perverse consequences are produced. The evidence on effectiveness is fairly limited, although a number of studies are ongoing. With respect to incentive levels, most quality-based performance incentives have accounted for merely 1–2% of total payment in the United States to

date. In the United Kingdom, however, 40% of general practitioners' salaries have been placed at risk according to performance across a wide array of parameters; this approach has been associated with substantial improvements in reported quality performance, although it is still unclear to what extent this change represents better performance versus better reporting. The potential for perverse consequences exists with any incentive scheme. One problem is that, if incentives are tied to outcomes, there may be a tendency to transfer the sickest patients to other providers and systems. Another concern is that providers will pay too much attention to quality measures with incentives and ignore the rest of the quality picture. The validity of these concerns remains to be determined. Nonetheless, it appears likely that, under health care reform, the use of various pay-for-performance schemes is likely to increase.

CONCLUSIONS

The safety and quality of care in the United States could be improved substantially. A number of available interventions have been shown to improve the safety of care and should be used more widely; others are undergoing evaluation or soon will be. Quality also could be dramatically better, and the science of quality improvement continues to mature. Implementation of pay-for-performance should make it much easier for organizations to justify investments in improving safety and quality parameters, including health information technology. However, many improvements will also require changing the structure of care—e.g., moving to a more team-oriented approach and ensuring that patients are more involved in their own care. Health care reform is likely to result in increased use of pay-for-performance. Measures of safety are still relatively immature and could be made much more robust; it would be particularly useful if organizations had measures they could use in routine operations to assess safety at a reasonable cost. Although the quality measures available are more robust than those for safety, they still cover a relatively small proportion of the entire domain of quality, and more measures need to be developed. The public and payers are demanding better information about safety and quality as well as better performance in these areas. The clear implication is that these domains will have to be addressed directly by providers.

13e Primary Care in Low- and Middle-Income Countries

Tim Evans, Kumanan Rasanathan

The twentieth century witnessed the rise of an unprecedented global health divide. Industrialized or high-income countries experienced rapid improvement in standards of living, nutrition, health, and health care. Meanwhile, in low- and middle-income countries with much less favorable conditions, health and health care progressed much more slowly. The scale of this divide is reflected in the current extremes of life expectancy at birth, with Japan at the high end (83 years) and Sierra Leone at the low end (47 years). This nearly 40-year difference reflects the daunting range of health challenges faced by low- and middle-income countries. These nations must deal not only with a complex mixture of diseases (both infectious and chronic) and illness-promoting conditions but also, and more fundamentally, with the fragility of the foundations underlying good health (e.g., sufficient food, water, sanitation, and education) and of the systems necessary for universal access to good-quality health care. In the last decades of the twentieth century, the need to bridge this global health divide and establish health equity was increasingly recognized. The Declaration of Alma Ata in 1978 crystallized a vision of justice in health, regardless of income, gender, ethnicity, or education, and called for “health for all by the year 2000” through primary health care. While much progress has been made since the declaration, at the end of the first decade and a half of the twenty-first century, much remains to be done to achieve global health equity.

This chapter looks first at the nature of the health challenges in low- and middle-income countries that underlie the health divide. It then outlines the values and principles of a primary health care approach, with a focus on primary care services. Next, the chapter reviews the experience of low- and middle-income countries in addressing health challenges through primary care and a primary health care approach. Finally, the chapter identifies how current challenges and global context provide an agenda and opportunities for the renewal of primary health care and primary care.

PRIMARY CARE AND PRIMARY HEALTH CARE

The term *primary care* has been used in many different ways: to describe a level of care or the setting of the health system, a set of treatment and prevention activities carried out by specific personnel, a set of attributes for the way care is delivered, or an approach to organizing health systems that is synonymous with the term *primary health care*. In 1996, the U.S. Institute of Medicine encompassed many of these different usages, defining primary care as “the provision of integrated, accessible health care services by clinicians who are accountable for addressing a large majority of personal health care needs, developing a sustained partnership with patients, and practicing in the context of family and community.”¹ We use this definition of *primary care* in this chapter. Primary care performs an essential function for health systems, providing the first point of contact when people seek health care, dealing with most problems, and referring patients onward to other services when necessary. As is increasingly evident in countries of all income levels, without strong primary care, health systems cannot function properly or address the health challenges of the communities they serve.

Primary care is only one part of a primary health care approach. The Declaration of Alma Ata, drafted in 1978 at the International Conference on Primary Health Care in Alma Ata (now Almaty in Kazakhstan), identified many features of primary care as being essential to achieving the goal of “health for all by the year 2000.” However, it also identified the need to work across different sectors, address

the social and economic factors that determine health, mobilize the participation of communities in health systems, and ensure the use and development of technology that was appropriate in terms of setting and cost. The declaration drew from the experiences of low- and middle-income countries in trying to improve the health of their people following independence. Commonly, these countries had built hospital-based systems similar to those in high-income countries. This effort had resulted in the development of high-technology services in urban areas while leaving the bulk of the population without access to health care unless they traveled great distances to these urban facilities. Furthermore, much of the population lacked access to basic public health measures. Primary health care efforts aimed to move care closer to where people lived, to ensure their involvement in decisions about their own health care, and to address key aspects of the physical and social environment essential to health, such as water, sanitation, and education.

After the Declaration of Alma Ata, many countries implemented reforms of their health systems based on primary health care. Most progress involved strengthening of primary care services; unexpectedly, however, much of this progress was seen in high-income countries, most of which constructed systems that made primary care available at low or no cost to their entire populations and that delivered the bulk of services in primary care settings. This endeavor also saw the reinforcement of family medicine as a specialty to provide primary care services. Even in the United States (an obvious exception to this trend), it became clear that the populations of states with more primary care physicians and services were healthier than those with fewer such resources.

Progress was also made in many low- and middle-income countries. However, the target of “health for all by the year 2000” was missed by a large margin. The reasons were complex but partly entailed a general failure to implement all aspects of the primary health care approach, particularly work across sectors to address social and economic factors that affect health and provision of sufficient human and other resources to make possible the access to primary care attained in high-income countries. Furthermore, despite the consensus in Alma Ata in 1978, the global health community rapidly became fractured in its commitment to the far-reaching measures called for by the declaration. Economic recession tempered enthusiasm for primary health care, and momentum shifted to programs concentrating on a few priority measures such as immunization, oral rehydration, breastfeeding, and growth monitoring for child survival. Success with these initiatives supported the continued movement of health development efforts away from the comprehensive approach of primary health care and toward programs that targeted specific public health priorities. This approach was reinforced by the need to address the HIV/AIDS epidemic. By the 1990s, primary health care had fallen out of favor in many global-health policy circles, and low- and middle-income countries were being encouraged to reduce public sector spending on health and to focus on cost-effectiveness analysis to provide a package of health care measures thought to offer the greatest health benefits.

HEALTH CHALLENGES IN LOW- AND MIDDLE-INCOME COUNTRIES

Low- and middle-income countries, defined by a per capita gross national income of <\$12,476 (U.S.) per person per year, account for >80% of the world’s population. Average life expectancy in these countries lags far behind that in high-income countries: whereas the average life expectancy at birth in high-income countries is 74 years, it is only 68 years in middle-income countries and 58 years in low-income countries. This discrepancy has received growing attention over the past 40 years. Initially, the situation in poor countries was characterized primarily in terms of high fertility and high infant, child, and maternal mortality rates, with most deaths and illnesses attributable to infectious or tropical diseases among remote, largely rural populations. With growing adult (and especially elderly) populations and changing lifestyles linked to global forces of urbanization, a new set of health challenges characterized by chronic diseases, environmental overcrowding, and road traffic injuries has emerged rapidly

¹Institute of Medicine. Primary Care: America’s Health in a New Era (1996).

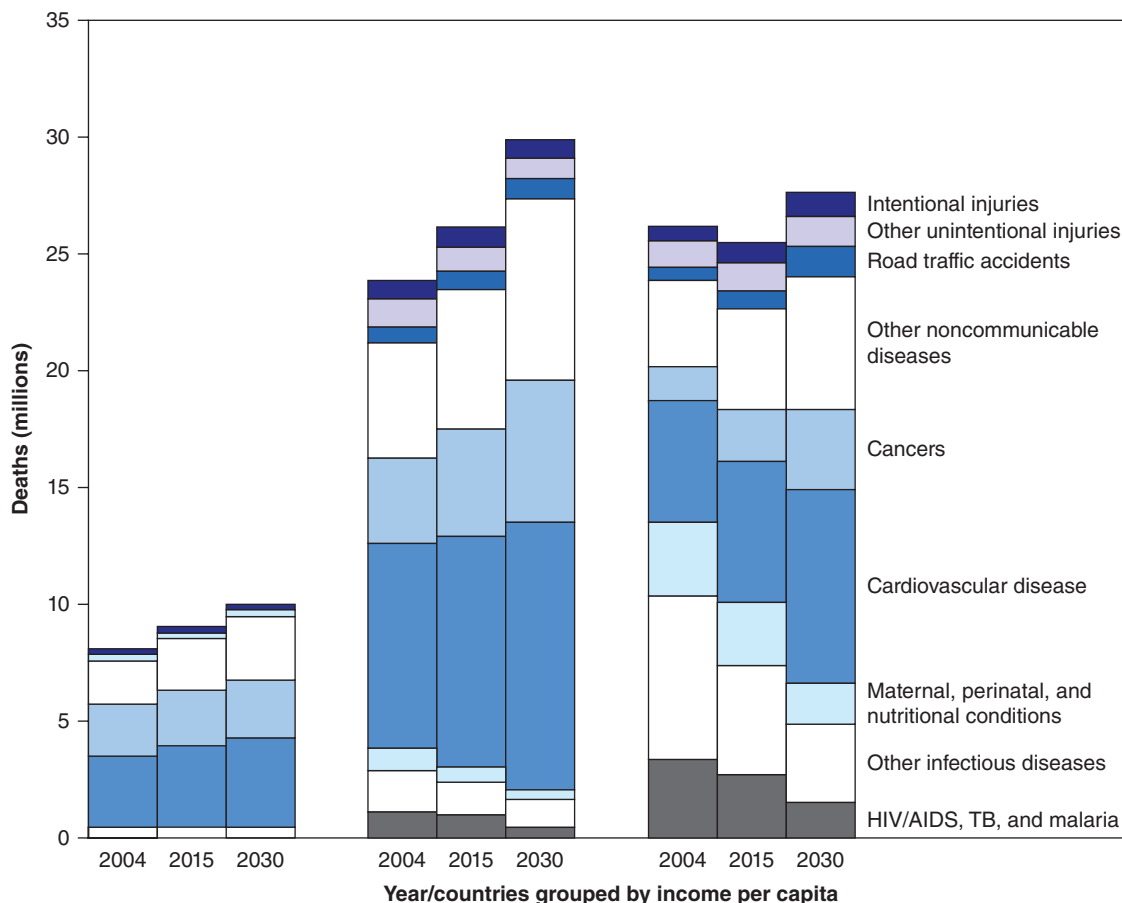


FIGURE 13e-1 Projections of disease burden to 2030 for high-, middle-, and low-income countries (left, center, and right, respectively). TB, tuberculosis. (Source: World Health Organization: *The Global Burden of Disease 2004 Update*, 2008.)

(Fig. 13e-1). The majority of tobacco-related deaths globally now occur in low- and middle-income countries, and the risk of a child’s dying from a road traffic injury in Africa is more than twice that in Europe. Hence, low- and middle-income countries in the twenty-first century face a full spectrum of health challenges—infectious, chronic, and injury-related—at much higher incidences and prevalences than are documented in high-income countries and with many fewer resources to address these challenges.

Addressing these challenges, however, does not mean simply waiting for economic growth. Analysis of the association between wealth and health across countries reveals that, for any given level of wealth, there is a substantial variation in life expectancy at birth that has persisted despite overall global progress in life expectancy during the past 30 years (Fig. 13e-2). Health status in low- and middle-income countries varies enormously. Nations such as Cuba and Costa Rica have life expectancies and childhood mortality rates similar to or even better than those in high-income countries; in contrast, countries in sub-Saharan Africa and the former Soviet bloc have experienced significant reverses in these health markers in the past 20 years.

As Angus Deaton stated in the World Institute for Development Economics Research annual lecture on September 29, 2006, “People in poor countries are sick not primarily because they are poor but because of other social organizational failures, including health delivery, which are not automatically ameliorated by higher income.” This analysis concurs with classic studies of the array of societal factors explaining good health in poor

settings such as Cuba and Kerala State in India. Analyses conducted over the past three decades indeed show that rapid health improvement is possible in very different contexts. That some countries continue to lag far behind can be understood through a comparison of regional differences in progress in terms of life expectancy over this period (Fig. 13e-3). While most regions have made impressive

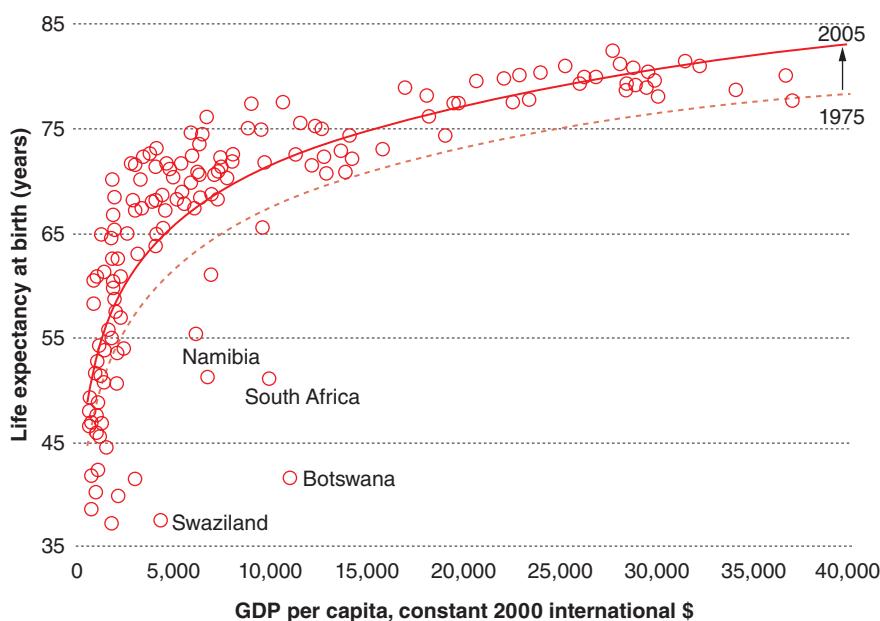


FIGURE 13e-2 Gross domestic product (GDP) per capita and life expectancy at birth in 169 countries, 1975 and 2005. Only outlying countries are named. (Source: World Health Organization: *Primary Health Care: Now More Than Ever*. World Health Report 2008.)

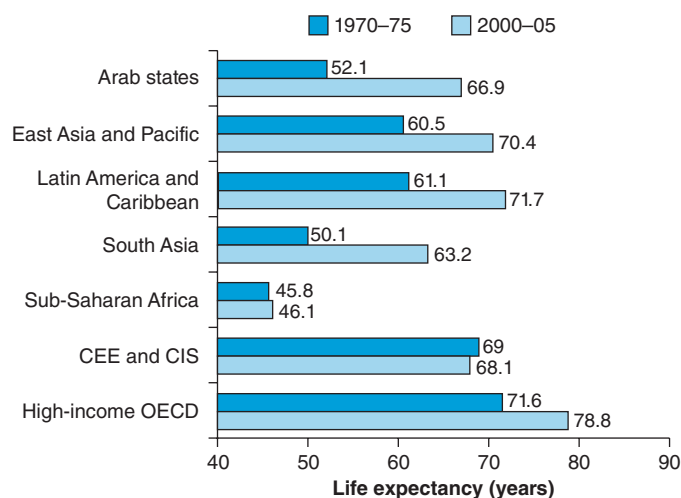


FIGURE 13e-3 Regional trends in life expectancy. CEE and CIS, Central and Eastern Europe and the Commonwealth of Independent States; OECD, Organization for Economic Co-operation and Development. (Source: World Health Organization: *Closing the Gap in a Generation: Health Equity Through Action on the Social Determinants of Health. Commission on Social Determinants of Health Final Report, 2008.*)

progress, sub-Saharan Africa and the former Soviet states have seen stagnation and even reversals.

As average levels of health vary across regions and countries, so too do they vary within countries (Fig. 13e-4). Indeed, disparities within countries are often greater than those between high-income and low-income countries. For example, if low- and middle-income countries could reduce their overall childhood mortality rate to that of the richest one-fifth of their populations, global childhood mortality could be decreased by 40%. Disparities in health are mostly a result of social and economic factors such as daily living conditions, access to resources, and ability to participate in life-affecting decisions. In most countries, the health care sector actually tends to exacerbate health inequalities (the “inverse-care law”); because of neglect and discrimination, poor and marginalized communities are much less likely to benefit from public health services than those that are better off. Reforming health systems toward people-centered primary care provides an opportunity to reverse these negative trends.

Health services have failed to make their contribution to reducing these pervasive social inequalities by ensuring universal access to existing, scientifically validated, low-cost interventions such as insecticide-treated bed nets for malaria, taxes on cigarettes, short-course

chemotherapy for tuberculosis, antibiotic treatment for pneumonia, dietary modification and secondary prevention measures for high blood pressure and high cholesterol levels, and water treatment and oral rehydration therapy for diarrhea. Despite decades of “essential packages” and “basic” health campaigns, the effective implementation of what is already known to work appears (deceptively) to be difficult.

Recent analyses have begun to focus on “the how” (as opposed to “the what”) of health care delivery, exploring why health progress is slow and sluggish despite the abundant availability of proven interventions for health conditions in low- and middle-income countries. Three general categories of reasons are being identified: (1) shortfalls in performance of health systems; (2) stratifying social conditions; and (3) skews in science.

SHORTFALLS IN PERFORMANCE OF HEALTH SYSTEMS

Specific health problems often require the development of specific health interventions (e.g., tuberculosis requires short-course chemotherapy). However, the delivery of different interventions is often facilitated by a common set of resources or functions: money or financing, trained health workers, and facilities with reliable supplies fit for multiple purposes. Unfortunately, health systems in most low- and middle-income countries are largely dysfunctional at present.

In the large majority of low- and middle-income countries, the level of public financing for health is woefully insufficient: whereas high-income countries spend, on average, 7% of the gross domestic product on health, middle-income countries spend <4% and low-income countries <3%. External financing for health through various donor channels has grown significantly over time. While these funds for health are significant (~\$20 billion [U.S.] in 2008 for low- and middle-income countries), they represent <2% of total health expenditures in low- and middle-income countries and hence are neither a sufficient nor a long-term solution to chronic underfinancing. In Africa, 70% of health expenditures come from domestic sources. The predominant form of health care financing—charging patients at the point of service—is the least efficient and the most inequitable, tipping millions of households into poverty annually.

Health workers, who represent another critical resource, are often inadequately trained and supported in their work. Recent estimates indicate a shortage of >4 million health workers, constituting a crisis that is greatly exacerbated by the migration of health workers from low- and middle-income countries to high-income countries. Sub-Saharan Africa carries 24% of the global disease burden but has only 3% of the health workforce (Fig. 13e-5). The International Organization for Migration estimated in 2006 that there were more Ethiopian physicians practicing in Chicago than in Ethiopia itself.

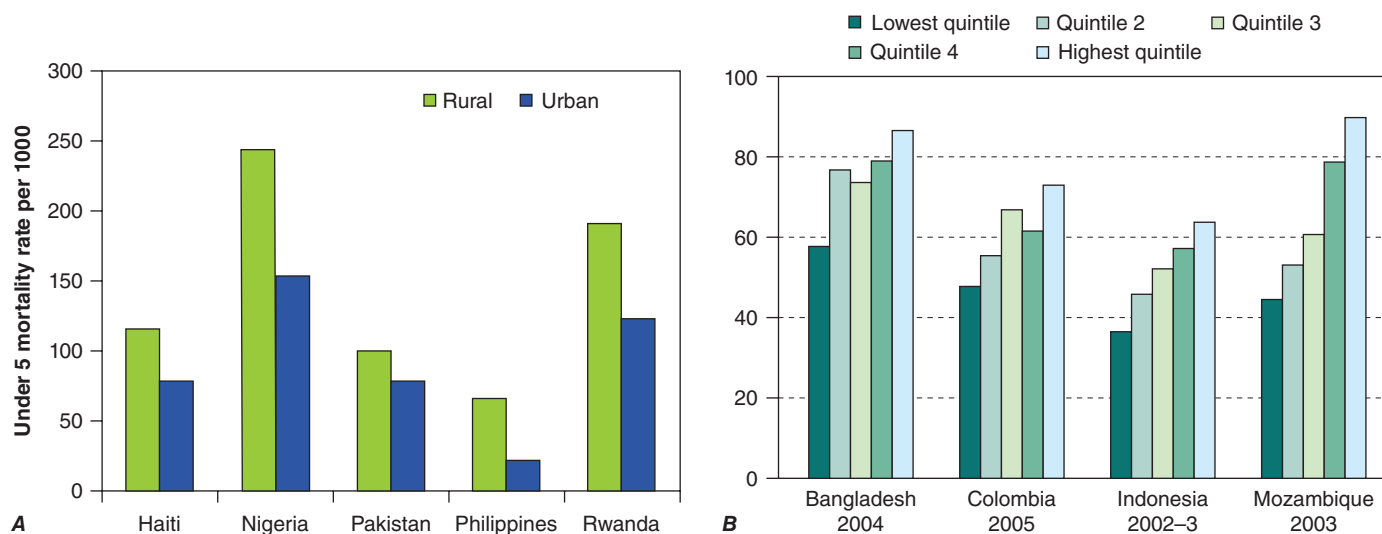


FIGURE 13e-4 A. Mortality of children under 5 years old, by place of residence, in five countries. (Source: Data from the World Health Organization.) B. Full basic immunization coverage (%), by income group. (Source: *Primary Health Care: Now More Than Ever. World Health Report 2008.*)

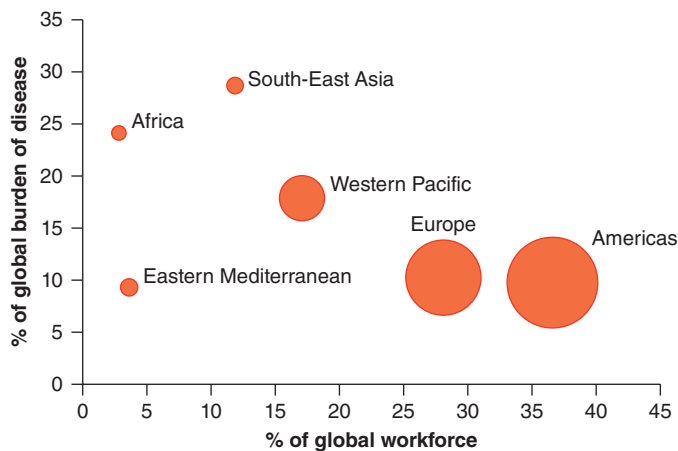


FIGURE 13e-5 Global burden of disease and health workforce.

(Source: World Health Organization: *Working Together for Health*, 2006.)

Critical diagnostics and drugs often do not reach patients in need because of supply-chain failures. Moreover, facilities fail to provide safe care: new evidence suggests much higher rates of adverse events among hospitalized patients in low- and middle-income countries than in high-income countries. Weak government planning, regulatory, monitoring, and evaluation capacities are associated with rampant, unregulated commercialization of health services and chaotic fragmentation of these services as donors “push” their respective priority programs. With such fragile foundations, it is not surprising that low-cost, affordable, validated interventions are not reaching those who need them.

STRATIFYING SOCIAL CONDITIONS

Health care delivery systems do not exist in a vacuum but rather are embedded in a complex of social and economic forces that often stratify opportunities for health unfairly. Most worrisome are the pervasive forces of social inequality that serve to marginalize populations with disproportionately large health needs (e.g., the urban poor; illiterate mothers). Why should a poor slum dweller with no income be expected to come up with the money for a bus fare needed to travel to a clinic to learn the results of a sputum test for tuberculosis? How can a mother living in a remote rural village and caring for an infant with febrile convulsions find the means to get her child to appropriate care? Shaky or nonexistent social security systems, dangerous work environments, isolated communities with little or no infrastructure, and systematic discrimination against minorities are among the myriad forces with which efforts for more equitable health care delivery must contend.

SKEWS IN SCIENCE

While science has yielded enormous breakthroughs in health in high-income countries, with some spillover to low- and middle-income countries, many important health problems continue to affect primarily low- and middle-income countries whose research and development investments are deplorably inadequate. The past decade has seen growing efforts to right this imbalance with research and development investment for new drugs, vaccines, and diagnostics that effectively cater to the specific health needs of populations in low- and middle-income countries. For example, the Medicines for Malaria Venture has revitalized a previously “dry” pipeline for new malaria drugs. This is but one of many such efforts, but much more needs to be done.

As discussed above, the primary constraint on better health in low- and middle-income countries is related less to the availability of health technologies and more to their effective delivery. Underlying these systems and social challenges to greater equity in health is a major bias regarding what constitutes legitimate “science” to improve health equity. The lion’s share of health research financing is channeled toward the development of new technologies—drugs, vaccines, and diagnostics; in contrast, virtually no resources are directed toward

research on how health care delivery systems can become more reliable and overcome adverse social conditions. The complexity of systems and social context is such that this issue of delivery requires an enormous investment in terms not only of money but also of scientific rigor, with the development of new research methods and measures and the attainment of greater legitimacy in the mainstream scientific establishment.

These common challenges to low- and middle-income countries partly explain the resurgence of interest in the primary health care approach. In some countries (mostly middle-income), significant progress has been made in expanding coverage by health systems based on primary care and even in improving indicators of population health. More countries are embarking on the creation of primary care services despite the challenges that exist, particularly in low-income countries. Even when these challenges are acknowledged, there are many reasons for optimism that low- and middle-income countries can accelerate progress in building primary care.

PRIMARY HEALTH CARE IN THE TWENTY-FIRST CENTURY

The new millennium has seen a resurgence of interest in primary health care as a means of addressing global health challenges. This interest has been driven by many of the same issues that led to the Declaration of Alma Ata: rapidly increasing disparities in health between and within countries, spiraling costs of health care at a time when many people lack quality care, dissatisfaction of communities with the care they are able to access, and failure to address changes in health threats, especially noncommunicable disease epidemics. These challenges require a comprehensive approach and strong health systems with effective primary care. Global health development agencies have recognized that sustaining gains in public health priorities such as HIV/AIDS requires not only robust health systems but also the tackling of social and economic factors related to disease incidence and progression. Weak health systems have proved a major obstacle to delivering new technologies, such as antiretroviral therapy, to all who need them. Changing disease patterns have led to a demand for health systems that can treat people as individuals whether or not they present to a health facility with the public health “priority” (e.g., HIV/AIDS or tuberculosis) to which that facility is targeted. We discuss experiences in low- and middle-income countries in relation to primary care in greater detail below. First, we consider the features of primary health care and primary care as currently understood.

REVITALIZATION OF PRIMARY HEALTH CARE

At the 2009 World Health Assembly (an annual meeting of all countries to discuss the work of the World Health Organization [WHO]), a resolution was passed reaffirming the principles of the Declaration of Alma Ata and the need for national health systems to be based on primary health care. This resolution did not suggest that nothing had changed in the intervening 30 years since the declaration, nor did it dispute that its prescription needed reframing in light of changing public health needs. The 2008 WHO World Health Report describes how a primary health care approach is necessary “now more than ever” to address global health priorities, especially in terms of disparities and new health challenges. As discussed below, this report highlights four broad areas in which reform is required (Fig. 13e-6). One of these areas—the need to organize health care so that it places the needs of people first—essentially relates to the necessity for strong primary care in health systems and what this requirement entails. The other three areas also relate to primary care. All four areas require action to move health systems in a direction that will reduce disparities and increase the satisfaction of those they serve. The World Health Report’s recommendations present a vision of primary health care that is based on the principles of Alma Ata but that differs from many attempts to implement primary health care in the 1970s and 1980s.

Universal Coverage Reforms to Improve Health Equity Despite progress in many countries, most people in the world can receive health care services only if they can pay at the point of service. Disparities in health are caused not only by a lack of access to necessary health



FIGURE 13e-6 The four reforms of primary health care renewal.

(Source: World Health Organization: *Primary Health Care: Now More Than Ever*. World Health Report 2008.)

services but also by the impact of expenditure on health. More than 100 million people are driven into poverty each year by health care costs, with countless others deterred from accessing services at all. Moving toward prepayment financing systems for universal coverage, which ensure access to a comprehensive package of services according to need without precipitating economic ruin, is therefore emerging as a major priority in low- and middle-income countries. Increasing coverage of health services can be considered in terms of three axes: the proportion of the population covered, the range of services underwritten, and the percentage of costs paid (Fig. 13e-7). Moving toward universal coverage requires ensuring the availability of health care services to all, eliminating barriers to access, and organizing pooled financing mechanisms, such as taxation or insurance, to remove user fees at the point of service. It also requires measures beyond financing, including expansion of health services in poorly served areas, improvement in the quality of services provided to marginalized communities, and increased coverage of other social services that significantly affect health (e.g., education).

Service Delivery Reforms to Make Health Systems People-Centered Health systems have often been organized around the needs of those who provide health care services, such as clinicians and policymakers. The result is a centralization of services or the provision of vertical programs that target single diseases. The principles of primary health

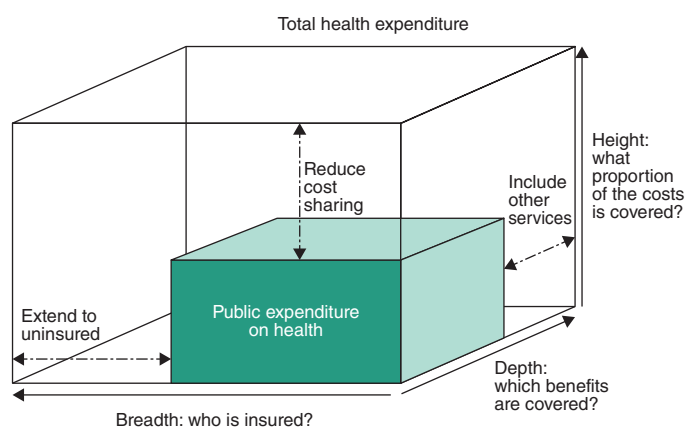


FIGURE 13e-7 Three ways of moving toward universal coverage.

care, including the development of primary care, reorient care around the needs of the people to whom services cater. This “people-centered” approach aims to provide health care that is both more effective and appropriate.

The increase in noncommunicable diseases in low- and middle-income countries offers a further stimulus for urgent reform of service delivery to improve chronic disease care. As discussed above, large numbers of people currently fail to receive relatively low-cost interventions that have reduced the incidence of these diseases in high-income countries. Delivery of these interventions requires health systems that can address multiple problems and manage people over a long period within their own communities, yet many low- and middle-income countries are only now starting to adapt and build primary care services that can address noncommunicable diseases and communicable diseases requiring chronic care. Even some countries (e.g., Iran) that have had significant success in reducing communicable diseases and improving child survival have been slow to adapt their health systems to rapidly accelerating noncommunicable disease epidemics.

People-centered care requires a safe, comprehensive, and integrated response to the needs of those presenting to health systems, with treatment at the first point of contact or referral to appropriate services. Because no discrete boundary separates people’s needs for health promotion, curative interventions, and rehabilitation services across different diseases, primary care services must address all presenting problems in a unified way. Meeting people’s needs also involves improved communication between patients and their clinicians, who must take the time to understand the impact of the patients’ social context on the problems they present with. This enhanced understanding is made possible by improvements in the continuity of care so that responsibility transcends the limited time people spend in health care facilities. Primary care plays a vital role in navigating people through the health system; when people are referred elsewhere for services, primary care providers must monitor the resulting consultations and perform follow-up. All too often, people do not receive the benefit of complex interventions undertaken in hospitals because they lose contact with the health care system once discharged. Comprehensiveness and continuity of care are best achieved by ensuring that people have an ongoing personal relationship with a care team.

Public Policy Reforms to Promote and Protect the Health of Communities

Public policies in sectors other than health care are essential to reduce disparities in health and to make progress toward global public health targets. The 2008 final report of the WHO Commission on Social Determinants of Health provides an exhaustive review of the intersectoral policies required to address health inequities at the local, national, and global levels. Advances against major challenges such as HIV/AIDS, tuberculosis, emerging pandemics, cardiovascular disease, cancers, and injuries require effective collaboration with sectors such as transport, housing, labor, agriculture, urban planning, trade, and energy. While tobacco control provides a striking example of what is possible if different sectors work together toward health goals, the lack of implementation of many evidence-based tobacco control measures in most countries just as clearly illustrates the difficulties encountered in such intersectoral work and the unrealized potential of public policies to improve health. At the local level, primary care services can help enact health-promoting public policies in other sectors.

Leadership Reforms to Make Health Authorities More Responsive

The Declaration of Alma Ata emphasized the importance of participation by people in their own health care. In fact, participation is important at all levels of decision-making. Contemporary health challenges require new models of leadership that acknowledge the role of government in reducing disparities in health but that also recognize the many types of organizations that provide health care services. Governments need to guide and negotiate among these different groups, including nongovernmental organizations (NGOs) and the private sector, and to provide strong regulation where necessary. This difficult task requires a massive reinvestment in leadership and governance capacity, especially if action by different sectors is to be effectively implemented. Moreover, disadvantaged groups and other actors are increasingly expecting that

their voices and health needs will be included in the decision-making process. The complex landscape for leadership at the national level is mirrored in many ways at the international or global level. The transnational character of health and the increasing interdependence of countries with respect to outbreak diseases, climate change, security, migration, and agriculture place a premium on more effective global health governance.

EXPERIENCES WITH PRIMARY CARE IN LOW- AND MIDDLE-INCOME COUNTRIES

Aspects of the primary health care approach described above, with an emphasis on primary care services, have been implemented to varying degrees in many low- and middle-income countries over the past half-century. As discussed above, some of these experiences inspired and informed the Declaration of Alma Ata, which itself led many more countries to attempt to implement primary health care. This section describes the experiences of a selection of low- and middle-income countries in improving primary care services that have enhanced the health of their populations.

Before Alma Ata, few countries had attempted to develop primary health care on a national level. Rather, most focused on expanding primary care services to specific communities (often rural villages), making use of community volunteers to compensate for the absence of facility-based care. In contrast, in the post–World War II period, China invested in primary care on a national scale, and life expectancy doubled within roughly 20 years. The Chinese expansion of primary care services included a massive investment in infrastructure for public health (e.g., water and sanitation systems) linked to innovative use of community health workers. These “barefoot doctors” lived in and expanded care to rural villages. They received a basic level of training that enabled them to provide immunizations, maternal care, and basic medical interventions, including the use of antibiotics. Through the work of the barefoot doctors, China brought low-cost universal basic health care coverage to its entire population, most of which had previously had no access to these services.

In 1982, the Rockefeller Foundation convened a conference to review the experiences of China along with those of Costa Rica, Sri Lanka, and the state of Kerala in India. In all of these locations, good health care at low cost appeared to have been achieved. Despite lower levels of economic development and health spending, all of these jurisdictions, along with Cuba, had health indicators approaching—or in some cases exceeding—those of developed countries. Analysis of these experiences revealed a common emphasis on primary care services, with expansion of care to the entire population free of charge or at low cost, combined with community participation in decision-making about health services and coordinated work in different sectors (especially education) toward health goals. During the three decades since the Rockefeller meeting, some of these countries have built on this progress, while others have experienced setbacks. Recent experiences in developing primary care services show that the same combination of features is necessary for success. For example, Brazil—a large country with a dispersed population—has made major strides in increasing the availability of health care in the past quarter century. In this millennium, the Brazilian Family Health Program has expanded progressively across the country, with almost all areas now covered. This program provides communities with free access to primary care teams made up of primary care physicians, community health workers, nurses, dentists, obstetricians, and pediatricians. These teams are responsible for the provision of primary care to all people in a specified geographic area—not only those who access health clinics. Moreover, individual community health workers are responsible for a named list of people within the area covered by the primary care team. Problems with access to health care persist in Brazil, especially in isolated areas and urban slums. However, solid evidence indicates that the Family

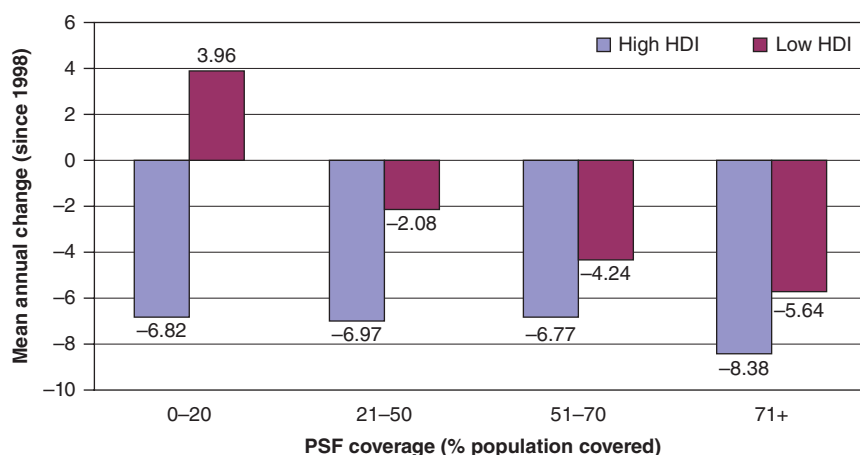


FIGURE 13e-8 Improvements in childhood mortality following the Family Health Program in Brazil. HDI, Human Development Index; PSF, Program Saúde da Família (Family Health Program). (Source: Ministry of Health, Brazil.)

Health Program has already contributed to impressive gains in population health, particularly in terms of childhood mortality and health inequities. In fact, this program has already had an especially marked impact on childhood mortality reduction in less developed areas (Fig. 13e-8).

Chile has also built on its existing primary care services in the past decade, aiming to improve the quality of care and the extent of coverage in remote areas, above all for disadvantaged populations. This effort has been made in concert with measures aimed at reducing social inequalities and fostering development, including social welfare benefits for families and disadvantaged groups and increased access to early-childhood educational facilities. As in Brazil, these steps have improved maternal and child health and have reduced health inequities. In addition to directly enhancing primary care services, Brazil and Chile have instituted measures to increase both the accountability of health providers and the participation of communities in decision-making. In Brazil, national and regional health assemblies with high levels of public participation are integral parts of the health policy-making process. Chile has instituted a patient’s charter that explicitly specifies the rights of patients in terms of the range of services to which they are entitled.

Other countries that have made recent progress with primary health care include Bangladesh, one of the poorest countries in the world. Since achieving its independence from Pakistan in 1971, Bangladesh has seen a dramatic increase in life expectancy, and childhood mortality rates are now lower than those in neighboring nations such as India and Pakistan. The expansion of access to primary health care services has played a major role in these achievements. This progress has been spearheaded by a vibrant NGO community that has focused its attention on improving the lives and livelihoods of poor women and their families through innovative and integrated microcredit, education, and primary care programs.

The above examples, along with others from the past 30 years in countries such as Thailand, Malaysia, Portugal, and Oman, illustrate how the implementation of a primary health care approach, with a greater emphasis on primary care, has led to better access to health care services—a trend that has not been seen in many other low- and middle-income countries. This trend, in turn, has contributed to improvements in population health and reductions in health inequities. However, as these nations have progressed, other countries have shown how previous gains in primary care can easily be eroded. In sub-Saharan Africa, undermining of primary care services has contributed to catastrophic reversals in health outcomes catalyzed by the HIV/AIDS epidemic. Countries such as Botswana and Zimbabwe implemented primary health care strategies in the 1980s, increasing access to care and making impressive gains in child health. Both countries have since been severely affected by HIV/AIDS, with pronounced

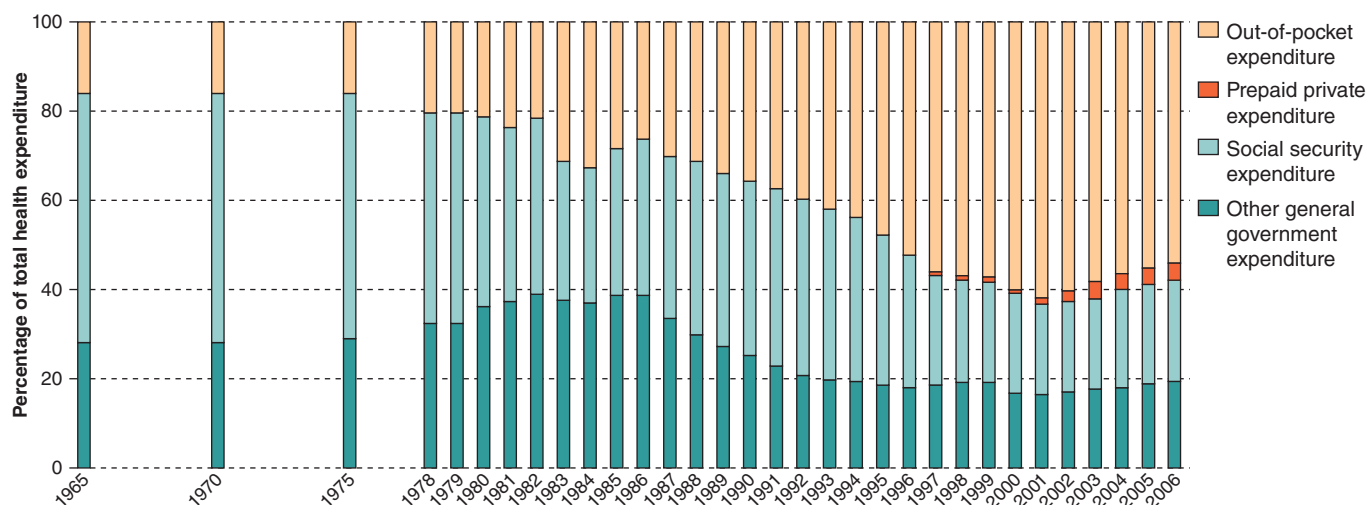


FIGURE 13e-9 Changes in source of health expenditure in China over the past 40 years. (Source: World Health Organization: *Primary Health Care: Now More Than Ever*. World Health Report 2008.)

decreases in life expectancy. However, Zimbabwe has also seen political turmoil, a decline of health and other social services, and the flight of health personnel, whereas Botswana has maintained primary care services to a greater extent and has managed to organize widespread access to antiretroviral therapy for people living with HIV/AIDS. Zimbabwe's health situation has therefore become more desperate than that in Botswana.

China provides a particularly striking example of how changes in health policy relevant to the organization of health systems (Fig. 13e-9) can have rapid, far-reaching consequences for population health. Even as the 1982 Rockefeller conference was celebrating China's achievements in primary care, its health system was unraveling. The decision to open up the economy in the early 1980s led to rapid privatization of the health sector and the breakdown of universal health coverage. As a result, by the end of the 1980s, most people, especially the poorer segments of the population, were paying directly out of pocket for health care, and almost no Chinese had insurance—a dramatic transformation. The “barefoot doctor” schemes collapsed, and the population either turned to care paid for at hospitals or simply became unable to access care. This undermining of access to primary care services in the Chinese system and the resulting increase in impoverishment due to illness contributed to the stagnation of progress in health in China at the same time that incomes in that country increased at an unprecedented rate. Reversals in primary care have meant that China now increasingly faces health care issues similar to those faced by India. In both countries, rapid economic growth has been linked to lifestyle changes and noncommunicable disease epidemics. The health care systems of the two nations share two negative features that are common when primary care is weak: a disproportionate focus on specialty services provided in hospitals and unregulated commercialization of health services. China and India have both seen expansion of private hospital services that cater to middle-class and urban populations who can afford care; at the same time, hundreds of millions of people in rural areas now struggle to access the most basic services. Even in the former groups, a lack of primary care services has been associated with late presentation with illness and with insufficient investment in primary prevention approaches. This neglect of prevention poses a risk of large-scale epidemics of cardiovascular disease, which could endanger continued economic growth. In addition, the health systems of both countries now depend for the majority of their funding on out-of-pocket payments by people when they use services. Thus substantial proportions of the population must sacrifice other essential goods as a result of health expenditure and may even be driven into poverty by this cost. The commercial nature of health services with inadequate or no regulation has also led to the proliferation of charlatan providers, inappropriate care, and pressure for people to pay for expensive and sometimes unnecessary care. Commercial providers have limited incentives

to use interventions (including public health measures) that cannot be charged for or that are what the person who is paying can afford.

Faced with these problems, China and India have implemented measures to strengthen primary health care. China has increased government funding of health care, has taken steps toward restoring health insurance, and has enacted a target of universal access to primary care services. India has similarly mobilized funding to greatly expand primary care services in rural areas and is now duplicating this process in urban settings. Both countries are increasingly using public resources from their growing economies to fund primary care services.

These encouraging trends are illustrative of new opportunities to implement a primary health care approach and strengthen primary care services in low- and middle-income countries. Brazil, India, China, and Chile are being joined by many other low- and middle-income countries, including Indonesia, Mexico, the Philippines, Turkey, Rwanda, Ethiopia, South Africa, and Ghana, in ambitious initiatives mobilizing new resources to move toward universal coverage of health services at affordable cost.

OPPORTUNITIES TO BUILD PRIMARY CARE IN LOW- AND MIDDLE-INCOME COUNTRIES

Global public health targets will not be met unless health systems are significantly strengthened. More money is currently being spent on health than ever before. In 2005, global health spending totaled \$5.1 trillion (U.S.)—double the amount spent a decade earlier. Although most expenditure occurs in high-income countries, spending in many emerging middle-income countries has rapidly accelerated, as has the allocation of monies for this purpose by both governments in and donors to low-income countries. These twin trends—greater emphasis on building health systems based on primary care and allotment of more money for health care—provide opportunities to address many of the challenges discussed above in low- and middle-income countries.

Accelerating progress requires a better understanding of how global health initiatives can more effectively facilitate the development of primary care in low-income countries. A review by the WHO Maximizing Positive Synergies Collaborative Group looked at programs funded by the Global Fund to Fight AIDS, Tuberculosis and Malaria; the Global Alliance for Vaccines and Immunisation (GAVI); the U.S. President's Emergency Plan for AIDS Relief (PEPFAR); and the World Bank (on HIV/AIDS). This group found that global health initiatives had improved access to and quality of the targeted health services and had led to better information systems and more adequate financing. The review also identified the need for better alignment of global health initiatives with other national health priorities and systematic exploitation of potential synergies. If global health initiatives implement programs that work in tandem with other components of national health systems without undermining staffing and procurement of supplies,

they have the potential to contribute substantially to the capacity of health systems to provide comprehensive primary care services.

Even in the aftermath of the global financial crisis, global health initiatives continue to draw significant funding. In 2009, for example, U.S. President Barack Obama announced increasing development assistance from the United States for global health, earmarking \$63 billion over the period 2009–2014 for a Global Health Initiative. New funding is also promised through a range of other initiatives focusing particularly on maternal and child health in low-income countries. The general trend is to coordinate this funding in order to reduce fragmentation of national health systems and to concentrate more on strengthening these systems. Comprehensive primary care in low-income countries must inevitably deal with the rapid emergence of chronic diseases and the growing prominence of injury-related health problems; thus, international health development assistance must become more responsive to these needs.

Beyond the new streams of funding for health services, other opportunities exist. Increased social participation in health systems can help build primary care services. In many countries, political pressure from community advocates for more holistic and accountable care as well as entrepreneurial initiatives to scale up community-based services through NGOs have accelerated progress in primary care without major increases in funding. Participation of the population in the provision of health care services and in relevant decision-making often drives services to cater to people's needs as a whole rather than to narrow public health priorities.

Participation and innovation can help address critical issues related to the health workforce in low- and middle-income countries by establishing effective people-centered primary care services. Many primary care services do not need to be delivered by a physician or a nurse. Multidisciplinary teams can include paid community workers who have access to a physician if necessary but who can provide a range of health services on their own. In Ethiopia, more than 30,000 community health workers have been trained and deployed to improve access to primary care services, and there is increasing evidence that this measure is contributing to better health outcomes. In India, more than 600,000 community health advocates have been recruited as part of expanded rural primary care services. In Niger, the deployment of community health workers to deliver essential child health interventions (as a component of integrated community case management) has had impressive results in reducing childhood mortality and decreasing disparities. After the Declaration of Alma Ata, experiences with community health workers were mixed, with particular problems about levels of training and lack of payment. Current endeavors are not immune from these concerns. However, with access to physician support and the deployment of teams, some of these concerns may be addressed. Growing evidence from many countries indicates that shifting appropriate tasks to primary care workers who have had shorter,

less expensive training than physicians will be essential to address the human resources crisis.

Finally, recent improvements in information and communication technologies, particularly mobile phone and Internet systems, have created the potential for systematic implementation of e-health, telemedicine, and improved health data initiatives in low- and middle-income countries. These developments raise the tantalizing possibility that health systems in these countries, which have long lagged behind those in high-income countries but are less encumbered by legacy systems that have proved hard to modernize in many settings, could leapfrog their wealthier counterparts in exploiting these technologies. Although the challenges posed by poor or absent infrastructure and investment in many low- and middle-income countries cannot be underestimated and will need to be addressed to make this possibility a reality, the rapid rollout of mobile networks and their use for health and other social services in many low-income countries where access to fixed telephone lines was previously very limited offer great promise in building primary care services in low- and middle-income countries.

CONCLUSION

As concern continues to mount about glaring inequities in global health, there is a growing commitment to redress these egregious shortfalls, as exemplified by global mobilization around the United Nations' Millennium Development Goals and the early discussions on what targets should build on these goals in the post-2015 era. This commitment begins first and foremost with a clear vision of the fundamental importance of health in all countries, regardless of income. The values of health and health equity are shared across all borders, and primary health care provides a framework for their effective translation across all contexts.

The translation of these fundamental values has its roots in four types of reforms that reflect the distinct but interlinked challenges of (re)orienting a society's resources on the basis of its citizens' health needs: (1) organizing health care services around the needs of people and communities; (2) harnessing services and sectors beyond health care to promote and protect health more effectively; (3) establishing sustainable and equitable financing mechanisms for universal coverage; and (4) investing in effective leadership of the whole of society. This common primary health care agenda highlights the striking similarity, despite enormous differences in context, in the nature and direction of the reforms that national health systems must undertake to promote greater equity in health. This shared agenda is complemented by the growing reality of global health interconnectedness due, for example, to shared microbial threats, bridging of ethnolinguistic diversity, flows in migrant health workers, and mobilization of global funds to support the neediest populations. Embracing solidarity in global health while strengthening health systems through a primary health care approach is fundamental to sustained progress in global health.

14e Complementary, Alternative, and Integrative Health Practices

Josephine P. Briggs

The search for health includes many beliefs and practices that are outside conventional medicine. Physicians are important sources for information and guidance about health matters, but our patients also rely on a wide range of other sources including family and friends, cultural traditions, alternative practitioners, and increasingly the Internet, popular media, and advertising. It is essential for physicians to understand what patients are doing to seek health, as this understanding is important to harness potential benefits and to help patients avoid harm.

DEFINITIONS AND SCOPE

The phrase *complementary and alternative medicine* is used to describe a group of diverse medical and health care systems, practices, and products that have historic origins outside mainstream medicine. Most of these practices are used together with conventional therapies and

therefore have been called *complementary* to distinguish them from *alternative* practices, those used as a substitute for standard care. Use of dietary supplements; mind-body practices such as acupuncture, massage, meditation, and hypnosis; and care from a traditional healer all fall under this umbrella. Brief definitions for some of the common complementary and alternative health practices are provided in **Table 14e-1**. Although some complementary health practices are implemented by a complementary health care provider such as a chiropractor, acupuncturist, or naturopathic practitioner, or by a physician, many of these practices are undertaken as “self-care.” Most are paid for out of pocket.

In the last decade or so, the terms *integrative care* and *integrative medicine* have entered the dialogue. A 2007 national survey conducted by the Centers for Disease Control and Prevention’s National Center for Health Statistics found that 42% of hospices had integrated complementary health practices into the care they provide. Integration of select complementary approaches is also common in Veterans Administration and Department of Defense facilities, particularly as part of management of pain and post-traumatic stress disorder.

The term *integrative medicine* is usually used to refer to a style of practice that places strong emphasis on a holistic approach to patient care while focusing on reduced use of technology. Physicians advocating

TABLE 14e-1 TERMINOLOGY OF COMPLEMENTARY AND ALTERNATIVE MEDICAL PRACTICES

Mind-Body Practices	
Acupuncture and acupressure	A family of procedures involving stimulation of defined anatomic points, a component of the major Asian medical traditions; most common application involves the insertion and manipulation of thin metallic needles
Alexander technique	A movement therapy that uses guidance and education to improve posture, movement, and efficient use of muscles for improvement of overall body functioning
Guided imagery	The use of relaxation techniques followed by the visualization of images, usually calm and peaceful in nature, to invoke specific images to alter neurologic function or physiologic states
Hypnosis	The induction of an altered state of consciousness characterized by increased responsiveness to suggestion
Massage	Manual therapies that manipulate muscle and connective tissues to promote muscle relaxation, healing, and sense of well-being
Meditation	A group of practices, largely based in Eastern spiritual traditions, intended to focus or control attention and obtain greater awareness of the present moment, or mindfulness
Reflexology	Manual stimulation of points on hands or feet that are believed to affect organ function
Rolfing/structural integration	A manual therapy that attempts to realign the body by deep tissue manipulation of fascia
Spinal manipulation	A range of manual techniques, employed by chiropractors and osteopaths, for adjustments of the spine to affect neuromuscular function and other health outcomes
Tai chi	A mind-body practice originating in China that involves slow, gentle movements and sometimes is described as “moving meditation”
Therapeutic touch	Secular version of the laying on of hands, described as “healing meditation”
Yoga	An exercise practice, originally East Indian, that combines breathing exercises, physical postures, and meditation
Traditional Medical Systems	
Ayurvedic medicine	The major East Indian traditional medicine system; treatment includes meditation, diet, exercise, herbs, and elimination regimens (using emetics and diarrheals)
Curanderismo	A spiritual healing tradition common in Latin American communities that uses ritual cleansing, herbs, and incantations
Native American medicine	Diverse traditional systems that incorporate chanting, shaman healing ceremonies, herbs, laying on of hands, and smudging (ritual cleansing with smoke from sacred plants)
Siddha medicine	An East Indian medical system (prevalent among Tamil-speaking people)
Tibetan medicine	A medical system that uses diagnosis by pulse and urine examination; therapies include herbs, diet, and massage
Traditional Chinese medicine	A medical system that uses acupuncture, herbal mixtures, massage, exercise, and diet
Unani medicine	An East Indian medical system, derived from Persian medicine, practiced primarily in the Muslim community; also called “hikmat”
“Modern” Medical Systems	
Anthroposophic medicine	A spiritually based system of medicine that incorporates herbs, homeopathy, diet, and a movement therapy called eurythmy
Chiropractic	Chiropractic care involves the adjustment of the spine and joints to alleviate pain and improve general health; primarily used to treat back problems, musculoskeletal complaints, and headaches
Homeopathy	A medical system with origins in Germany that is based on a core belief in the theory of “like cures like”—compounds that produce certain syndromes, if administered in very diluted solutions, will be curative
Naturopathy	A clinical discipline that emphasizes a holistic approach to the patient, herbal medications, diet, and exercise; practitioners have degrees as doctors of naturopathy
Osteopathy	A clinical discipline, now incorporated into mainstream medicine, that historically emphasized spinal manipulative techniques to relieve pain, restore function, and promote overall health

this approach generally include selected complementary health practices in the care they offer patients, and many have established practice settings that include complementary health practitioners. Although this approach appears to be attractive to many patients, the heavy use of dietary supplements and the weaknesses in the evidence base for a number of the interventions offered in integrative practices continue to attract substantial concern and controversy.

Until a decade ago or so, “complementary and alternative medicine” could be defined as practices that are neither taught in medical schools nor reimbursed, but this definition is no longer workable, since medical students increasingly seek and receive some instruction about complementary health practices, and some practices are reimbursed by third-party payers. Another definition, practices that lack an evidence base, is also not useful, since there is a growing body of research on some of these modalities, and some aspects of standard care do not have a strong evidence base.

By its nature, the demarcation between mainstream medicine and complementary health practices is porous, varying from culture to culture and over time. Traditional Chinese medicine and the Indian practice of Ayurvedic medicine were once the dominant health teachings in those cultures. Certain health practices that arose as challenges to the mainstream have been integrated gradually into conventional care. Examples include the teachings of Fernand Lamaze that led to the widespread use of relaxation techniques during childbirth, the promotion of lactation counseling by the La Leche League, and the teaching of Cicely Saunders and Elizabeth Kübler-Ross that established the hospice movement.

The late nineteenth century saw the development of a number of healing philosophies by care providers who were critical of the medicine of the time. Of these, naturopathy and homeopathy, which arose in Germany, and chiropractic and osteopathy, which developed in the United States, have continued to endure. Osteopathic medicine is currently thoroughly integrated into conventional medicine, although the American Medical Association (AMA) labeled it a cult as late as 1960. The other three traditions have remained resolutely separate from mainstream medicine, although chiropractic care is available in some conventional care settings.

PATTERNS OF USE

The first large survey of use of these practices was performed by David Eisenberg and associates in 1993. It surprised the medical community by showing that more than 30% of Americans use complementary or alternative health approaches. Many studies since that time have extended those conclusions. Subsequently, the National Health Interview Survey (NHIS), a large, national survey conducted by the National Center for Health Statistics, a component of the Centers for Disease Control and Prevention, has addressed the use of complementary health practices and largely confirmed those results. The NHIS is a household survey of many kinds of health practices in the civilian population; it uses methods that create a nationally representative sample and has a sample size large enough to permit valid estimates about some subgroups. In 2002, 2007, and 2012, the survey included a set of questions that addressed complementary and alternative health approaches. Information was obtained from 31,000 adults in 2002 and 23,300 adults and 9400 children in 2007. Only preliminary data are available from the 2012 survey. In all three surveys, approximately 40% of adults report using some form of complementary therapy or health practice. In the 2007 study, 38% of adults and 12% of children had used one or more modalities. These surveys yield the estimate that nonvitamin, nonmineral dietary supplements are used by approximately 18% of the population. The most prevalent mind-body practices are relaxation techniques and meditation, chiropractic, and therapeutic massage. Americans are willing to pay for these services; the estimated out-of-pocket expenditure for complementary health practices in 2007 was \$34 billion, representing 1.5% of total health expenditures and 11% of out-of-pocket costs.

The appeal of unproven complementary health approaches continues to perplex many physicians. Many factors contribute to these choices. Some patients seek out complementary health practitioners

because they offer optimism or greater personal attention. For others, alternative approaches reflect a “self-help” approach to health and wellness or satisfy a search for “natural” or less invasive alternatives, since dietary supplements and other natural products are believed, correctly or not, to be inherently healthier and safer than standard pharmaceuticals. In NHIS surveys, the most common health conditions cited by patients for use of complementary health practices involve management of symptoms often poorly controlled by conventional care, particularly back pain and other painful musculoskeletal complaints, anxiety, and insomnia.

PRACTITIONER-BASED DISCIPLINES

Licensure and Accreditation At present, six fields of complementary health practice—osteopathic manipulation, chiropractic, acupuncture and traditional Chinese medicine, therapeutic massage, naturopathy, and homeopathy—are subject to some form of educational accreditation and state licensure. Accreditation of educational programs is the responsibility of professional organizations or commissions under federal oversight by the Department of Education. Licensure, in contrast, is strictly a state matter, generally determined by state legislatures. Legal recognition establishes public access to therapies even when there is no scientific consensus about their clinical value.

Osteopathic Manipulative Therapy Founded in 1892 by the physician Andrew Taylor Still, osteopathic medicine was originally based on the belief that manipulation of soft tissue and bone can correct a wide range of diseases of the musculoskeletal and other organ systems. Over the ensuing century, the osteopathic profession has welcomed increasing integration with conventional medicine. Today, the postgraduate training, practice, credentialing, and licensure of osteopathic physicians are virtually indistinguishable from those of allopathic physicians. Osteopathic medical schools, however, include training in manual therapies, particularly spinal manipulation. Approximately 70% of family practice osteopathic physicians perform manipulative therapies on some of their patients.

Chiropractic The practice of chiropractic care, founded by David Palmer in 1895, is the most widespread practitioner-based complementary health practice in the United States. Chiropractic practice emphasizes manual therapies for treatment of musculoskeletal complaints, although the scope of practice varies widely, and in some rural areas, chiropractors may serve a primary care role, due in part to the lack of other providers. According to the NHIS, approximately 8% of Americans receive chiropractic manipulation in a given year.

Since the mid-1970s, chiropractors have been licensed in all 50 states and reimbursed by Medicare. Chiropractic educational standards mandate 2 years of undergraduate training, 4 years of training at an accredited school of chiropractic, and in most states, successful completion of a standardized board examination. Postgraduate training is not required. The U.S. Department of Labor estimates that there are 52,000 licensed chiropractors (2010 figure). There is substantial geographic variation, with greater numbers of practitioners and greater use in the midwest, particularly in rural areas, and lower use in the southeast.

Historically, the relationship between the medical and chiropractic professions has been strained. Extending through the 1970s, the AMA set forth standards prohibiting physicians consulting or entering into professional relationships with chiropractors, but in 1987, after a decade of complex litigation, the U.S. District Court found the AMA in violation of antitrust laws. An uneasy truce has followed, with continued physician skepticism, but also evidence for robust patient demand and satisfaction.

The role of both osteopathic and chiropractic spinal manipulative therapies (SMTs) in back pain management has been the subject of a number of carefully performed trials and many systematic reviews. Conclusions are not consistent, but the most recent guidelines from the American College of Physicians and the American Pain Society conclude that SMT is associated with small to moderate benefit for low-back pain of less than 4 weeks in duration (evidence level B/C) and moderate benefit (evidence level B) for subacute or chronic low-

back pain. The evidence of benefit for neck pain is not as extensive, and continued concern that cervical manipulation may occasionally precipitate vascular injury clouds a contentious debate.

Naturopathy Naturopathy is a discipline that emerged in central Europe in the nineteenth century as part of the Natural Cure movement and was introduced to the United States in the early twentieth century by Benjamin Lust. Fifteen states currently license naturopathic physicians, with considerable variation in the scope of practice. The naturopathic profession is actively seeking licensure in other states. There are estimated to be approximately 3000 licensed naturopathic physicians in the United States. There is also a robust naturopathy presence in Canada. Conventional and unconventional diagnostic tests and medications are prescribed, with an emphasis on relatively low doses of drugs, herbal medicines, healthy diet, and exercise. While there is some support for success of naturopathic practitioners in motivating healthy behaviors, concern exists about the heavy promotion of dietary supplements, most with little rigorous evidence.

Homeopathy Homeopathy was widespread in the United States in the late nineteenth and early twentieth centuries and continues to be a common alternative practice in many European countries, but estimates from the NHIS suggest that less than 1.5% of Americans visit a homeopathic practitioner in any given year. In the United States, licensure as a homeopathic physician is only possible in three states (Arizona, Connecticut, and Nevada) where it is restricted to licensed physicians. The number of practitioners is uncertain, however, because some states include homeopathy within the scope of practice of other fields, including chiropractic and naturopathy, and some practitioners may self-identify as homeopathic practitioners. As discussed below, the regulatory framework for homeopathic remedies differs from other dietary supplements. Homeopathic remedies are widely available and commonly recommended by naturopathic physicians, chiropractors, and other licensed and unlicensed practitioners.

Therapeutic Massage The field of therapeutic massage is growing rapidly, as use by the public is increasing. According to U.S. Department of Labor statistics, there are approximately 155,000 licensed massage therapists employed in the United States, and by 2020, this number is projected to grow by 20%. Forty-three states and the District of Columbia currently have laws regulating massage therapy; however, there is little consistency, and in some states, regulation is by town ordinance. States that do provide licensure for massage therapists typically require a minimum of 500 hours of training at an accredited institution, as well as meeting specific continuing education requirements and carrying malpractice insurance. Massage training programs generally are approved by a state board, but some may also be accredited by an independent agency, such as the Commission on Massage Therapy Accreditation (COMTA). The development of regulatory standards for therapeutic massage has not yet caught up with the evolution of the field or the high demand. Many techniques used are also employed by physical therapists.

Acupuncture and Traditional Chinese Medicine A venerable component of traditional Chinese medicine, with a history of use that extends at least 2000 years, acupuncture became better known in the United States in 1971, when *New York Times* reporter James Reston wrote about how doctors in China used needles to ease his pain after surgery. More than 3 million adults in the United States use acupuncture, according to NHIS data. In a number of European countries, acupuncture is performed primarily by physicians. In the United States, the training and licensure processes for physicians and nonphysicians differ. Currently, acupuncture is licensed in 42 states and the District of Columbia, with licensure standards varying within the scope of practice of each state. Licensure for nonphysicians generally requires 3 years of accredited training and the successful completion of a standardized examination. The main accrediting organization is the Accreditation Commission for Acupuncture and Oriental Medicine. Acupuncture is included in doctor of medicine (MD) and doctor of osteopathic medicine (DO) licensure in 31 states, with 11 states requiring additional training for physicians performing acupuncture.

MIND-BODY INTERVENTIONS

Mind-body practices are a large and diverse group of techniques that are administered or taught to others by a trained practitioner or teacher. Examples include acupuncture, massage therapy, meditation, relaxation techniques, spinal manipulation, and yoga. These approaches are being used more frequently in mainstream health care facilities for both patients and health care providers. Mind-body practices such as meditation and yoga are not licensed in any state, and training in those practices is not subject to national accreditation.

Americans often turn to complementary approaches for help in managing health conditions associated with physical and psychological pain—especially back pain, headache, musculoskeletal complaints, and functional pain syndromes. Chronic pain management is often refractory to conventional medical approaches, and standard pharmacologic approaches have substantial drawbacks. Health care guidelines of the American Pain Society and other professional organizations recognize the value of certain complementary approaches as adjuncts to pharmacologic management.

The evidence base for the effectiveness of these modalities is still relatively incomplete, but a few rigorous examples where there is promise of usefulness and safety include acupuncture for osteoarthritis pain; tai chi for fibromyalgia pain; and massage, yoga, and spinal manipulation for chronic back pain. In addition, new research is shedding light on the effects of meditation and acupuncture on central mechanisms of pain processing and perception and regulation of emotion and attention. Although many unanswered questions remain about these effects, findings are pointing to scientifically plausible mechanisms by which these modalities might yield benefit.

DIETARY SUPPLEMENTS

Regulation The Dietary Supplements Health and Education Act (DSHEA), passed in 1994, gives authority to the U.S. Food and Drug Administration (FDA) to regulate dietary supplements, but with expectations that differ in many respects from the regulation of drugs or food additives. Purveyors of dietary supplements cannot claim that they prevent or treat any disease. They can, however, claim that they maintain “normal structure and function” of body systems. For example, a product cannot claim to treat arthritis, but it can claim to maintain “normal joint health.” Homeopathic products predate FDA drug regulations and are sold with no requirement that they be proved effective. Although homeopathic products are widely believed to be safe because they are highly dilute, one product, a nasal spray called Zicam, was withdrawn from the market when it was found to produce anosmia, probably because of a significant zinc content. Homeopathic products, and indeed other complementary health products and practices, also convey the very significant risk that individuals will use them instead of effective conventional modalities.

Regulation of advertising and marketing claims is the purview of the Federal Trade Commission (FTC). The FTC does take legal action against promoters or websites that advertise or sell dietary supplements with false or deceptive statements.

Inherent Toxicity Although the public may believe that “natural” equates with “safe,” it is abundantly clear that natural products can be toxic. Misidentification of medicinal mushrooms has led to liver failure. Contamination of tryptophan supplements caused the eosinophilia-myalgia syndrome. Herbal products containing particular species of *Aristolochia* were associated with genitourinary malignancies and interstitial nephritis. In 2013, dietary supplements containing 1,3-dimethylamylamine (DMAA), often touted as a “natural” stimulant, led to cardiovascular problems, including heart attacks. Among the most controversial dietary supplements is *Ephedra sinica*, or ma huang, a product used in traditional Chinese medicine for short-term treatment of asthma and bronchial congestion. The scientific basis for these indications was revealed when ephedra was shown to contain the ephedrine alkaloids, especially ephedrine and pseudoephedrine. With the promulgation of the DSHEA regulations, supplements containing ephedra and herbs rich in caffeine sold widely in the U.S. marketplace because of their claims to promote weight loss and enhance athletic

performance. Reports of severe and fatal adverse events associated with use of ephedra-containing products led to an evidence-based review of the data surrounding them, and in 2004, the FDA banned their sale in the United States.

Another major current concern with dietary supplements is adulteration with pharmacologic active compounds. Multi-ingredient products marketed for weight loss, body building, “sexual health,” and athletic performance are of particular concern. Recent FDA recalls have involved contamination with steroids, diuretics, stimulants, and phosphodiesterase type 5 inhibitors.

Herb-Drug Interactions A number of herbal products have potential impact on the metabolism of drugs. This effect was illustrated most compellingly with the demonstration in 2000 that consumption of St. John’s wort interferes with the bioavailability of the HIV protease inhibitor indinavir. Later studies showed its similar interference with metabolism of topoisomerase inhibitors such as irinotecan, with cyclosporine, and with many other drugs. The breadth of interference stems from the ability of hyperforin in St. John’s wort to upregulate expression of the pregnane X receptor, a promiscuous nuclear regulatory factor that promotes the expression of many hepatic oxidative, conjugative, and efflux enzymes involved in drug and food metabolism.

Because of the large number of compounds that alter drug metabolism and the large number of agents some patients are taking, identification of all potential interactions can be a daunting task. Several useful Web resources are available as information sources (Table 14e-2). Clearly, attention to this problem is particularly important with drugs with a narrow therapeutic index, such as anticoagulants, antiseizure medications, antibiotics, immunosuppressants, and cancer chemotherapeutic agents.

PATIENT AND PROVIDER RESOURCES

Physicians regularly face difficult challenges in providing patients with advice and education about complementary practices. Of particular concern to all physicians are practices of uncertain safety and practices that raise inappropriate hopes. Cancer therapies, antiaging regimens, weight-loss programs, sexual function, and athletic performance are frequently targeted for excessive claims and irresponsible marketing. A number of Internet resources provide critical tools for patient education (Table 14e-3). Because many complementary health products and practices are used as self-care and because many patients research these approaches extensively on the Internet, directing patients to responsible websites can often be very helpful.

TABLE 14e-2 RESOURCES FOR DIETARY SUPPLEMENT–DRUG INTERACTIONS

Medscape

<http://www.medscape.com/druginfo/druginterchecker?cid=med>

This website is maintained by WebMD and includes a free drug interaction checker tool that provides information on interactions between two or more drugs, herbals, and/or dietary supplements.

Natural Medicines Comprehensive Database

<http://naturaldatabase.therapeuticresearch.com>

This website provides an interactive natural product–drug interaction checker tool that identifies interactions between drugs and natural products, including herbals and dietary supplements. This service is available by subscription. A PDA version is available.

Natural Standard

<http://www.naturalstandard.com/tools/>

This website provides an interactive tool for checking drug and herb/supplement interactions. This service is available by subscription. A PDA version is available.

Abbreviation: PDA, personal digital assistant.

TABLE 14e-3 INTERNET RESOURCES ON COMPLEMENTARY HEALTH APPROACHES

The Cochrane Collaboration Complementary Medicine Reviews

This website offers rigorous systematic reviews of mainstream and complementary health interventions using standardized methods. It includes more than 300 reviews of complementary health practices. Complete reviews require institutional or individual subscription, but summaries are available to the public.

<http://www.cochrane.org/cochrane-reviews>

MedlinePlus All Herbs and Supplements, A–Z List

MedlinePlus Complementary and Alternative Medicine

NLM FAQ: Dietary Supplements, Complementary or Alternative Medicines

These National Library of Medicine (NLM) Web pages provide an A–Z database of science-based information on herbal and dietary supplements; basic facts about complementary health practices; and federal government sources on information about using natural products, dietary supplements, medicinal plants, and other complementary health modalities.

http://www.nlm.nih.gov/medlineplus/druginfo/herb_All.html

<http://www.cochrane.org/cochrane-reviews>

<http://www.nlm.nih.gov/medlineplus/dietarysupplements.html>

National Institutes of Health National Center for Complementary and Alternative Medicine (NCCAM)

This National Institutes of Health NCCAM website contains information for consumers and health care providers on many aspects of complementary health products and practices. Downloadable information sheets include short summaries of complementary health approaches, uses and risks of herbal therapies, and advice on wise use of dietary supplements.

<http://www.nccam.nih.gov>

Resources for Health Care Providers: <http://www.nccam.nih.gov/health/providers>

NCCAM Clinical Digest e-Newsletter: <http://www.nccam.nih.gov/health/providers/digest>

Continuing medical education lectures: <http://www.nccam.nih.gov/training/videolectures>

The scientific evidence regarding complementary therapies is fragmentary and incomplete. Nonetheless, in some areas, particularly pain management, it is increasingly possible to perform the kind of rigorous systematic reviews of complementary health approaches that are the cornerstone of evidence-based medicine. A particularly valuable resource in this respect is the Cochrane Collaboration, which has performed more than 300 systematic reviews of complementary health practices. Practitioners will find this a valuable source to answer patient questions. Practice guidelines, particularly for pain management, are also available from several professional organizations. Links to these resources are provided in Table 14e-3.

SUMMARY

The use of complementary and alternative health practices reflects an active interest in improved health. An array of unproven modalities will always be used by our patients. While some of these choices need to be actively discouraged, many are in fact innocuous and can be accommodated. Some may be genuinely helpful, particularly in the management of troublesome symptoms. The dialogue with patients about complementary health practices is an opportunity to understand patients’ beliefs and expectations and use those insights to help guide health-seeking practices in a constructive way.

ACKNOWLEDGMENT

The late Dr. Stephen Straus contributed this chapter in prior editions, and some material from his chapter has been retained here.

15e The Economics of Medical Care

Joseph P. Newhouse

15e-1

CHAPTER 15e

The Economics of Medical Care

The purpose of this chapter is to explain to physicians how economists think about physicians' decision-making with regard to the treatment of patients. Economists' mode of thinking has shaped health care policy and institutions and thus the environment in which physicians practice, not only in the United States but in many other countries as well. It may prove useful for physicians to understand some aspects of the economists' way of thinking, even if it sometimes seems foreign or uncongenial.

Physicians see themselves as professionals and as healers, assisting their patients with their health care needs. When economists are patients, they probably see physicians the same way, but when they view doctors through the lens of economics as a discipline, they see physicians—and their patients as well—as economic agents. In other words, economists are interested in the degree to which physicians and patients respond to various incentives in deciding how to deploy the resources over which they exercise choice. Examples of issues that would concern an economist include how much time physicians devote to seeing a patient; which tests they order; what drugs, if any, they prescribe; whether they recommend a procedure; whether they refer a patient; and whether they admit a patient to the hospital. In addition, patients consider the cost when they make a decision about whether to seek care.

To say that economists view physicians and patients as economic agents is not to say that economists consider financial incentives the predominant factor in the decisions that either physicians or patients make about treatment; it is to say only that these incentives have some influence on these decisions. In fact, the role played by financial incentives in medical decision-making may often be dwarfed by the roles played by scientific knowledge, by professional norms and ethics, and by the influence of peers. However, economic policy greatly influences financial incentives, and economists tend to focus on this domain. Their interest stems from fundamental economic questions: What goods and services are produced and consumed? In particular, how much medical care is available, and how much of other goods and services? How is medical care produced? For example, what mix of specific services is used in treating a particular episode of illness? Who receives particular treatments?

Physicians in all societies live and function in economic markets, although those markets differ greatly both from the simple competitive markets depicted in introductory economics textbooks and from country to country, depending on national institutions. Many of the differences between actual medical markets and textbook competitive markets cause what economists term *market failure*, a condition in which some individuals can be made better off without making anyone else worse off.

This chapter explains two features of health care financing that cause market failure: selection and moral hazard. A common response to market failure in medical care is what economists refer to as *administered prices*, which this chapter also describes. Unfortunately, administered prices exact a cost, leading to what economists call *regulatory or government failure*. All societies seek a balance between market failure and regulatory failure, a topic addressed in this chapter's conclusion.

SELECTION

In the idealized competitive market found in economic textbooks, buyers and sellers have the same knowledge about the goods or services they are buying and selling. When one party knows more—or when goods of different quality are being sold at the same price, which is analytically similar—markets can break down in the following sense: There may be a price at which an equally well informed buyer and seller could make a transaction that would make them both better off, but that transaction does not occur because one party knows more than the other. Hence, both the potential buyer and the potential seller are worse off.

The used car market is a classic example of differential information. Owners of used cars (potential sellers) know more about the quality of their cars than do potential buyers. At any specific market price for a certain make and model of car, the only used cars offered will be those whose sellers value them at less than that price. Assuming that quality varies among used cars, those that are offered for sale will differentially be of low quality (“lemons”) relative to the given price. Owners of higher-quality cars may simply hold on to them. If, however, a potential buyer knew that the car was of higher quality, the buyer might be willing to pay enough so that the owner of the higher-quality car would sell. It is for this reason that sellers may offer warranties and guarantees, although these are uncommon (but not unknown) in medical care.

The same thing happens when goods of different quality are sold at the supermarket at the same price. Shoppers are happy to take any box of a particular brand of breakfast cereal or any bottle of a particular soft drink on the shelf because the quality of the contents of any box or bottle is the same; however, that is not the case in the produce section, where shoppers will inspect the fruit they pick up to ensure that the apple is not bruised or the banana overly ripe. At the end of the day, it is the bruised apples and overly ripe bananas that are left in the store. In effect, the seller has not used all the available information in pricing the produce, and buyers exploit that information differential.

Selection affects markets for individual—and, to some degree, small-group—health insurance in a fashion similar to that seen in the used car market and at the produce stand, but in this case it is the buyer of insurance who has more knowledge than the seller. Individuals who use above-average amounts of care—for example, those with a chronic disease or a strong proclivity to seek care for a symptom—will value health insurance more than will those who are healthy or who for various reasons shun medical care even if they are symptomatic. However, the insurer does not necessarily know the risk of those it insures, and so it gears insurance premiums to an average risk, which in some instances is conditional on certain observable characteristics, such as age. Just as shoppers do not want the bruised apples and used car buyers do not want lemons, many healthy people will not want to buy insurance voluntarily if its price mainly reflects its use by those who are sick. (Healthy but very risk-averse individuals still may be willing to pay premiums well above their expected use.) In an extreme case, healthy people drop out of the insurance pool, premiums rise because the average person left in the pool is sicker, that rise causes still more people to drop out of the pool, premiums rise further, and so forth, until few people remain to buy insurance.

For this reason, no developed country relies primarily on voluntary individual insurance to finance health care, although many countries use it in the supplemental insurance market, and selection is, in fact, often a feature of that market. Instead, governments and/or employers provide or heavily subsidize the purchase of either mandated or voluntary health insurance (e.g., in Canada or Germany, the Medicare and Medicaid programs in the United States and the purchase of insurance in exchanges by lower-income persons) or provide health services directly (e.g., the United Kingdom and the U.S. Veterans Health Administration). In addition, governments or third parties administering individual insurance markets with competing insurers may “risk-adjust” payments to insurers; that is, transfer monies from insurers who enroll better risks (as measured by observable features, such as diagnoses that are not used to rate premiums) to insurers who enroll worse risks. This feature is found in the American Medicare Advantage program and American insurance exchanges as well as in Germany and the Netherlands. The idea is to reduce insurers' incentives to structure their products in order to appeal to good risks, especially when insurers are making choices about networks and formularies.

Moreover, countries that rely on employment-based health insurance, such as the United States and Germany, either mandate taxes to finance that insurance or provide large tax subsidies for its purchase; otherwise, many healthy employees would prefer that the employer give them the money the employer uses to subsidize the insurance as cash wages. Because an employer that offers health insurance will pay lower cash wages than an otherwise equivalent employer that does

not, larger American employers that, before the Affordable Care Act was implemented in 2014, were not required to offer insurance may not, in fact, have offered it if they had many low-wage employees; the reason is that, if they had offered insurance, the cash wage they could afford to pay would have been below the minimum wage. (For the same reason, these employers typically do not offer a pension benefit.) Many low-wage employers, however, are small businesses that might not be viable if they had to subsidize health insurance. As a result, the Affordable Care Act exempted firms with fewer than 50 employees from any penalties if their employees received a public subsidy and purchased insurance in the exchange. Some self-employed individuals or those who work at small firms may belong to a trade association or a professional society through which they can purchase insurance, but because that purchase is voluntary, it is subject to selection.

How does this situation affect the practice of medicine? Prior to the Affordable Care Act, individual and small-group insurance policies typically had preexisting condition clauses to protect the insurer against selection—that is, to protect the insurer against a person's purchasing insurance (or more complete insurance) after that person had been diagnosed with a disease that is expensive to treat. Even though there is now a penalty for remaining uninsured, some individuals still choose to do so, and others purchase insurance with substantial amounts of cost sharing that they may not be able to pay if they become sick. Caring for such patients may give the physician a choice between making do with less than clinically optimal treatment and proceeding in a clinically optimal way but leaving the patient with a large bill and possible bankruptcy—and potentially leaving the physician with bill collection issues or unpaid bills.

Selection can arise in a different guise when physicians are reimbursed a fixed amount per patient (i.e., *capitation*) rather than receiving fee-for-service payments. Depending on the adequacy of any adjustments in the capitated amount for the resources that a specific patient will require ("risk adjustment"), physicians who receive a fixed amount have a financial incentive to avoid caring for sicker patients. Similarly, physicians who receive a capitated amount for their own services but are not financially responsible for hospital care or the services of other physicians may make an excessive number of referrals, just as physicians reimbursed in a fee-for-service arrangement may make too few.

MORAL HAZARD

The term *moral hazard* comes from the actuarial literature; it originally referred to the weaker incentives of an insured individual to prevent the loss against which he or she is insured. A classic example is failure of homeowners in areas prone to brush fires to cut the brush around their houses or possibly install fire-resistant shingles on their roofs because of their expectation that insurance will compensate them if their houses burn down. In some lines of insurance, however, moral hazard is not a substantive issue. Persons who buy life insurance on their own lives are not likely to commit suicide so that their heirs can receive the proceeds. Moreover, despite the brush fire example, homeowner's insurance probably has little moral hazard associated with it because individuals often cannot replace some valued personal items when a house burns down. In short, if moral hazard is negligible, insured persons take appropriate precautions against the potential loss.

In the context of health insurance, this classic form of moral hazard refers to potentially reduced incentives to prevent illness, but that is probably not a major issue. Sickness and disease generally imply some pain and suffering, not to mention possibly shortened life expectancy. Because there is no insurance for pain and suffering, individuals have strong incentives to try to remain healthy regardless of how much health insurance they have. Put another way, having better health insurance probably does not alter those incentives much.

Instead of weakened incentives to prevent illness, *moral hazard* in the health insurance context typically refers to the incentives for better-insured individuals to use more medical services. For instance, a patient with back pain or shoulder pain might seek an MRI if it costs him or her little or nothing, even if the physician feels the clinical value of the MRI is negligible. Conversely, the physician may be more

cautious in ordering a test that seems likely to produce little information if there are severe financial consequences for the patient.

Some of the strongest evidence on this point comes from the randomized RAND Health Insurance Experiment conducted in the late 1970s and early 1980s. Families whose members were under 65 years of age were randomized to insurance plans in which the amount they had to pay when using services ("cost sharing") varied from nothing (fully insured care) to a large deductible (catastrophic insurance). All the plans capped families' annual out-of-pocket payments, with a reduced cap for low-income families. Families with complete insurance used ~40% more services in a year than did families with catastrophic insurance, a figure that did not vary much across the six geographically dispersed sites in which the experiment was run. Although these data come from the era before managed care in the United States, subsequent observational studies in this country and elsewhere have largely confirmed the experiment's findings with respect to the relationship between variations in care use and variations in patient payment at the point of service. The difference among the plans was almost entirely related to the likelihood that a patient would seek care. Once care was sought, there appeared to be little difference in how physicians treated their patients in different plans.

One might assume that the additional care provided to fully insured patients would have resulted in improved outcomes, but by and large it did not. In fact, there was little or no difference in average health outcomes among the different health plans, with the important exception that hypertension, especially in patients with low incomes, was better controlled when care was free.

A possible explanation for the paucity of beneficial effects attributable to the additional medical services used by fully insured patients lies in the observations that (1) the additional care targeted both problems for which care can be efficacious and those for which it is not and (2) the population in the experiment, which consisted of non-elderly community-dwelling individuals, was mostly healthy. Perhaps the additional two visits and the greater number of hospitalizations when care was free were as likely to lead to poor outcomes as to good outcomes in that population. Certainly, the subsequent literature on quality of care and medical error rates has implied that a good deal of inappropriate care was—and is—provided to patients. For example, more than half of the antibiotics prescribed to the experiment's participants were prescribed for viral conditions. Moreover, about one-quarter of patients who were hospitalized (in all plans) were admitted for procedures that could have been performed equally well outside the hospital, in line with the substantial decrease in hospital use over the last three decades. In short, the additional inappropriate care provided when care was free was not necessarily innocuous; if a mainly healthy person saw a physician, he or she could have been made worse off. The literature on inappropriate care is mostly American in origin, but the finding probably holds elsewhere as well.

Finally, patients' health habits did not change in response to insurance status. This finding is consistent with the intuition that moral hazard does not much affect incentives to prevent illness.

Recently, another randomized experiment was conducted in Oregon among low-income, childless adults who were uninsured. Many people who gained insurance coverage in 2014 when the United States implemented the Affordable Care Act are from this group. Some of the uninsured childless adults won a lottery that made them eligible for Medicaid; those who did not win became the comparison group. After ~2 years, the results suggested that the use of services by persons on Medicaid had increased by ~25–35%. Medicaid served its purpose of providing protection against large medical bills; there was an 81% reduction in the proportion of families who spent >30% of their income on medical care, and there were large reductions in both medical debt and borrowing to pay for medical care. Turning to health outcomes, there was a 30% reduction in depression among the uninsured who received Medicaid relative to the comparison group as well as an increase in the numbers of diagnosed diabetics and of diabetics taking medication. Although there were no statistically significant changes in measures of blood pressure, lipids, or glycosylated

hemoglobin, confidence intervals were sufficiently large that clinically meaningful effects could not be ruled out.

In sum, insurance is certainly desirable to protect families against the financial risk of large medical expenses and in some instances to address underuse of valuable medical services (e.g., by a patient with cardiovascular disease who fails to take medications for financial reasons). Thus, the remedy for moral hazard is not to abolish insurance but instead to strike the right balance between financial protection and incentives to seek care. Moreover, it is probably useful to vary the amounts that patients pay out of pocket, depending on the specific service and the patient's clinical condition. Health outcomes after myocardial infarction, for example, were better among patients who were randomized to have no copayments for statins, beta blockers, angiotensin-converting enzyme inhibitors, and angiotensin receptor blockers than among those who had to pay for these drugs.

ADMINISTERED PRICES

Because insurers, whether public or private, cannot pay any price that a physician sets, prices in medical markets with widespread insurance are either set administratively or negotiated. In the simple textbook model of a competitive market, prices approximate the cost of production, but this is not necessarily the case when prices are administered. In the traditional American Medicare program, for example, the government sets a take-it-or-leave-it price. Because of the market share represented by the program, the great majority of physicians choose to take the government's price rather than leave the program. In some countries (e.g., Canada and Germany), medical societies negotiate fees for all physicians in the nation or in a subnational area. In the United States, commercial insurers negotiate fees with individual physicians or groups of physicians.

The principal problem with administered prices arises because someone must set them and that person has an imperfect knowledge of cost. If the price that is set departs markedly from incremental cost, distortions inevitably result. Because the price-setter typically has little information about incremental cost, the set price could be, and often is, far from the actual cost. If the regulator sets the price below cost, the service may not be available or, if it is, will have to be cross-subsidized from a profitable service. If the price is set above cost, there may well be excess entry and too many services being offered on too small a scale. A beneficent regulator in theory could approximate a competitive price by trial and error if technology did not change, but that condition clearly does not hold in medical care. Not only do new goods and services appear continually, but physicians often become more skilled at delivering a service that is already available or at developing new tools to deliver that service at a different (and frequently lower) cost. For example, cataract surgery, which took upward of 8 h when first introduced, can now be completed in <30 min.

The distortions between price and cost when prices are administered have consequences for the way medical care is produced. There may well be too much capacity in "profitable" areas of medicine, such as cardiac services and sports medicine, and too little in less profitable areas, such as primary care. A fee above incremental cost for a procedure encourages more procedures.

Conversely, payment methods that attempt to cover many services with one fixed payment, such as capitation and a per-admission payment, pay nothing for doing more and therefore may result in too few services and in choices by providers to reduce the number of unprofitable patients under their care, much as a hospital may shutter an emergency room if it becomes a magnet for the uninsured. These phenomena also reflect the asymmetry of information between patients and physicians and, in the case of fee-for-service payment, the incentive for insured patients to go along with a recommendation for

additional services ("I am pretty sure I know what the problem is, but let's just carry out this additional test to be sure").

There is good evidence that physicians respond to prices that are set. For example, if there is a general reduction in fees that, other things being equal, would lower practice income, some physicians order more services, whereas the opposite pertains if all fees increase. This behavior is sufficiently well established empirically that the U.S. Medicare program's actuaries account for it in their estimates of what various changes in fees will cost or save. The outcome is different if the fee for one procedure or service changes and that procedure accounts for a modest proportion of income. In that case, another service can be substituted. For example, if the fee for a mastectomy increases relative to that for breast-conserving surgery, there will be a higher proportion of mastectomies. A particularly striking example of this type of behavior occurred when Medicare sharply reduced the fees it paid oncologists for chemotherapy in 2005. The proportion of lung cancer patients who received chemotherapy rose by 10%. Margins for some chemotherapeutic agents, however, were cut more than those for other agents, and thereafter oncologists made less use of the agents whose margins had fallen more.

Negotiated prices may get closer to cost than administered prices that are set, but they are not a panacea. First, negotiated prices may well exceed cost when there is no effective competition among similar physicians in a particular market. Because medical care markets are typically local, there may only be one group or a few groups in any particular specialty in a smaller market, in which case physicians will have considerable market power to obtain more favorable reimbursement. Further increasing physicians' market power is the fact that many, and probably most, patients are reluctant to change physicians because their current physician knows their medical history, because they are uncertain whether a new physician would be an improvement, and because insurance may shield them from most of the cost differences among physicians. Finally, in the United States, commercial insurers often negotiate fees as a multiple of the Medicare fee schedule; thus, any distortion in administratively determined relative fees is carried over into negotiated fees. For example, in the Medicare fee schedule, procedures generally are more profitable than cognitive services known as "evaluation and management," and this difference probably plays a role in the numerical insufficiency of primary care physicians in the United States.

CONCLUSION

One branch of economics—*positive economics*—seeks to explain actual phenomena without making a judgment about the desirability of those phenomena. Another branch—*normative economics*—seeks to prescribe what should happen and, in particular, what public policy should be developed to ensure that it happens. The main result of the application of normative economics is that, under certain very special assumptions, competitive markets result in a system in which no one can be made better off without another person's being made worse off. These assumptions do not hold in medical care, in part because of selection and moral hazard; economists term the result a *market failure*. By contrast, as the discussion of administered prices in this chapter indicates, even a beneficent regulator will introduce distortions from lack of sufficient information; moreover, there is no guarantee that a regulator will be beneficent, as periodic corruption scandals show. Economists term this phenomenon *regulatory* or *government failure*. Economists see decisions about the proper form and amount of public intervention and regulation in medical care as a matter of finding the right balance between various types of market failures and various types of regulatory failures—a balance that different societies may choose to strike differently.

16e Racial and Ethnic Disparities in Health Care

Joseph R. Betancourt, Alexander R. Green

Over the course of its history, the United States has experienced dramatic improvements in overall health and life expectancy, largely as a result of initiatives in public health, health promotion, disease prevention, and chronic care management. Our ability to prevent, detect, and treat diseases in their early stages has allowed us to target and reduce rates of morbidity and mortality. Despite interventions that have improved the overall health of the majority of Americans, racial and ethnic minorities (blacks, Hispanics/Latinos, Native Americans/Alaskan Natives, Asian/Pacific Islanders) have benefited less from these advances than whites and have suffered poorer health outcomes from many major diseases, including cardiovascular disease, cancer, and diabetes. Research has revealed that minorities may receive less care and lower-quality care than whites, even when confounders such as stage of presentation, comorbidities, and health insurance are controlled. These differences in quality are called *racial and ethnic disparities in health care*. Addressing these disparities has taken on greater importance with the significant transformation of the U.S. health care system and the implementation of health care reform and value-based purchasing. The shift toward creating financial incentives and disincentives to achieve quality goals makes focusing on those who receive lower-quality care more important than ever before. This chapter will provide an overview of racial and ethnic disparities in health and health care, identify root causes, and provide key recommendations to address these disparities at both the clinical and health system levels.

NATURE AND EXTENT OF DISPARITIES

Minority Americans have poorer health outcomes than whites from preventable and treatable conditions such as cardiovascular disease, diabetes, asthma, cancer, and HIV/AIDS (Fig. 16e-1). Multiple factors contribute to these racial and ethnic disparities in health. First and foremost, social determinants—such as lower socioeconomic status (SES; e.g., lower income, less wealth, and lower educational attainment), inadequate and unsafe housing, and racism—are strongly linked to poor health outcomes. These factors disproportionately impact minority populations. In fact, SES has consistently been found to be one of the strongest predictors of health outcomes. While the mechanisms are complex (i.e., does poverty cause poor health, or does

poor health cause poverty?), it is clear that low-SES populations experience disparities in health and that low SES is a major factor in racial/ethnic disparities.

Racial/ethnic disparities are documented globally, although their assessment has centered more on the comparison of individuals by SES in other countries than in the United States. Similar to the U.S. pattern, low-SES residents of other nations tend to have poorer health outcomes. It is noteworthy that results are mixed when the health status of nations is compared by SES. High-SES nations such as the United States do not necessarily have health outcomes that correlate with their high expenditures for health care. For example, as of 2011, the United States ranks 34th in the world—just behind Cuba—on basic public health measures such as infant mortality. This ranking may be due in part to the correlation between wealth distribution and SES rather than just absolute SES. This area of active research is outside the scope of this chapter.

Racism has more recently been shown to predict poorer health outcomes. The physiologic impact of the stress imposed by racism (and poverty), including increased cortisol levels, can lead to chronic adverse effects on health. Lack of access to care also takes a significant toll. Uninsured individuals are less likely to have a regular source of care and are more likely to delay seeking care and to go without needed care; this limited access results in avoidable hospitalizations, emergency hospital care, and adverse health outcomes.

In addition to racial and ethnic disparities in *health*, there are racial and ethnic disparities in the *quality of care* for persons with access to the health care system. For instance, disparities have been found in the treatment of pneumonia (Fig. 16e-2) and congestive heart failure, with blacks receiving less optimal care than whites when hospitalized for these conditions. Moreover, blacks with end-stage renal disease are referred less often to the transplant list than are their white counterparts (Fig. 16e-3). Disparities have been found, for example, in the use of cardiac diagnostic and therapeutic procedures (with blacks being referred less often than whites for cardiac catheterization and bypass grafting), prescription of analgesia for pain control (with blacks and Latinos receiving less pain medication than whites for long-bone fractures and cancer), and surgical treatment of lung cancer (with blacks receiving less curative surgery than whites for non-small-cell lung cancer). Again, many of these disparities have occurred even when variations in factors such as insurance status, income, age, comorbid conditions, and symptom expression are taken into account. However, one additional factor—disparities in the quality of care provided at the sites where minorities tend to receive care—has been shown to be an important contributor to overall disparities.

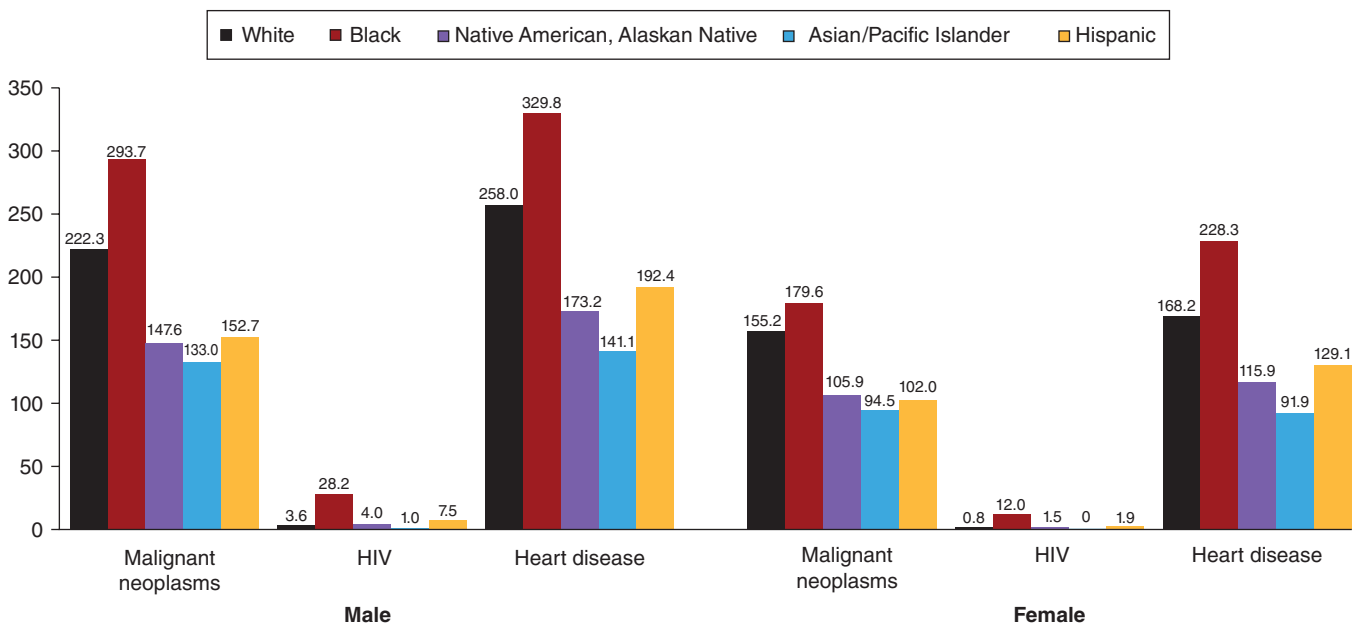


FIGURE 16e-1 Age-adjusted death rates for selected causes by race and Hispanic origin, 2005. (From the U.S. Census Bureau, 2009.)

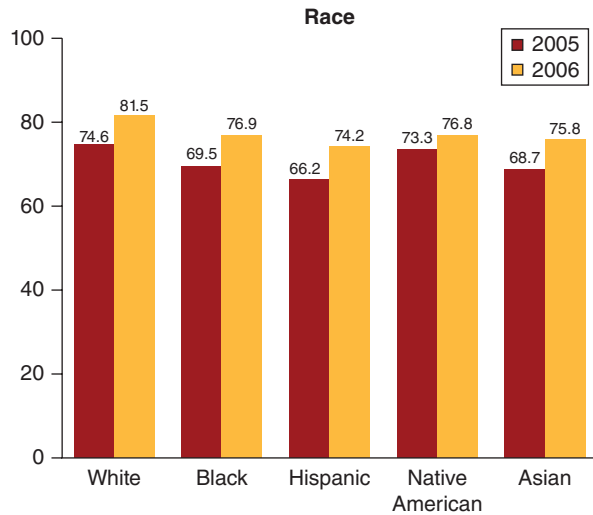


FIGURE 16e-2 Recommended hospital care received by Medicare patients with pneumonia, by race/ethnicity, 2006. The reference population consisted of Medicare beneficiaries with pneumonia who were hospitalized. The composite was calculated by averaging the percentage of the population that received each of the five incorporated components of care. (Adapted from Agency for Healthcare Research and Quality: *The 2008 National Health Care Disparities Report*.)

Little progress has been made in addressing racial/ethnic disparities in cardiovascular procedures and other advanced surgical procedures, whereas some progress has been made in eliminating disparities in primary-care process measures. Data from the National Registry of Myocardial Infarction found evidence of continued disparities in guideline-based admission, procedural, and discharge therapy use from 1994 to 2006. Black patients were less likely than white patients to receive percutaneous coronary intervention/coronary artery bypass grafting (PCI/CABG), a disparity that has improved little since 1994. Further, compared with whites, black patients were less likely to receive lipid-lowering medications at discharge, with a gap that has widened since 1998 (Fig. 16e-4). A 2009 study showed that blacks had worse post-myocardial infarction outcomes than whites but that the difference could be explained by site of care and patient factors (such as socioeconomic status and comorbid conditions).

The Centers for Disease Control and Prevention (CDC) analyzed national and state rates of total knee replacement (TKR) for Medicare enrollees for the period 2000–2006, with patients stratified by sex, age, and black or white race. TKR rates overall in the United States

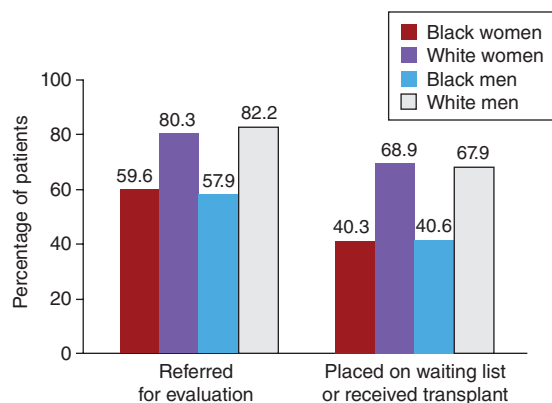


FIGURE 16e-3 Referral for evaluation at a transplantation center or placement on a waiting list/receipt of a renal transplant within 18 months after the start of dialysis among patients who wanted a transplant, according to race and sex. The reference population consisted of 239 black women, 280 white women, 271 black men, and 271 white men. Racial differences were statistically significant among both the women and the men ($p < .0001$ for each comparison). (From JZ Ayanian et al: *N Engl J Med* 341:1661, 1999.)

increased 58%, with similar increases among whites (61%) and blacks (56%). However, the TKR rate for blacks was 37% lower than the rate for whites in 2000 and 39% lower in 2006; i.e., the disparity not only did not improve but even worsened slightly (Fig. 16e-5). More recent data (up to 2010) show no apparent change in these figures. Data for enrollees in Medicare managed-care plans provides evidence for a narrowing in racial disparities between 1997 and 2003 in several “report card” preventive care measures, such as mammography and glucose and cholesterol testing. However, racial disparities in more complex measures, such as glucose control in diabetic patients and cholesterol levels in patients after a heart attack, actually worsened during that interval.

The 2012 National Healthcare Disparities Report, released by the Agency for Healthcare Research and Quality, found little improvement in disparities for core measures of quality over the previous decade. In fact, for blacks, Asians, Native Americans/Alaskan Natives, Hispanics, and poor people, the vast majority of core quality measures (87–92%) stayed the same, and a small proportion (2–8%) got worse, including measures of effectiveness, patient safety, and timeliness of care. While a small number of these measures improved, disparities were eliminated in no measured area. This annual report is particularly important, given that most studies of disparities have not been repeated with the same methodology used to document possible trends. Some studies have tracked disparities using specific disease and treatment registries. For example, by 2008, the use of acute and discharge medications for myocardial infarction had largely been equalized among racial and ethnic groups; however, African-American and Hispanic patients still experienced longer delays before reperfusion, with door-to-balloon times of <90 min for 83% of white patients as opposed to 75% and 76% of black and Hispanic patients, respectively.

ROOT CAUSES OF DISPARITIES

The Institute of Medicine (IOM) report *Unequal Treatment*, released in March 2002, remains the preeminent study of racial and ethnic disparities in health care in the United States. The IOM was charged with assessing the extent of racial/ethnic differences in health care that are not otherwise attributable to known factors, such as access to care. To provide recommendations regarding interventions aimed at eliminating health care disparities, the IOM studied health system, provider, and patient factors. The study found the following:

- Racial and ethnic disparities in health care exist and, because they are associated with worse health outcomes, are unacceptable.
- Racial and ethnic disparities in health care occur in the context of (1) broader historic and contemporary social and economic inequality and (2) evidence of persistent racial and ethnic discrimination in many sectors of American life.
- Many sources—including health systems, health care providers, patients, and utilization managers—may contribute to racial and ethnic disparities in health care.
- Bias, stereotyping, prejudice, and clinical uncertainty on the part of health care providers may contribute to racial and ethnic disparities in health care.
- A small number of studies suggest that minority patients may be more likely to refuse treatments, yet these refusal rates are generally small and do not fully explain health care disparities.

Unequal Treatment went on to identify a set of root causes that included the following:

- **Health system factors:** These include issues related to the complexity of the health care system, the difficulty that minority patients may have in navigating this complex system, and the lack of availability of interpreter services to assist patients with limited English proficiency. In addition, health care systems are generally ill prepared to identify and address disparities.
- **Provider-level factors:** These include issues related to the health care provider, including stereotyping, the impact of race/ethnicity on clinical decision-making, and clinical uncertainty due to poor communication.

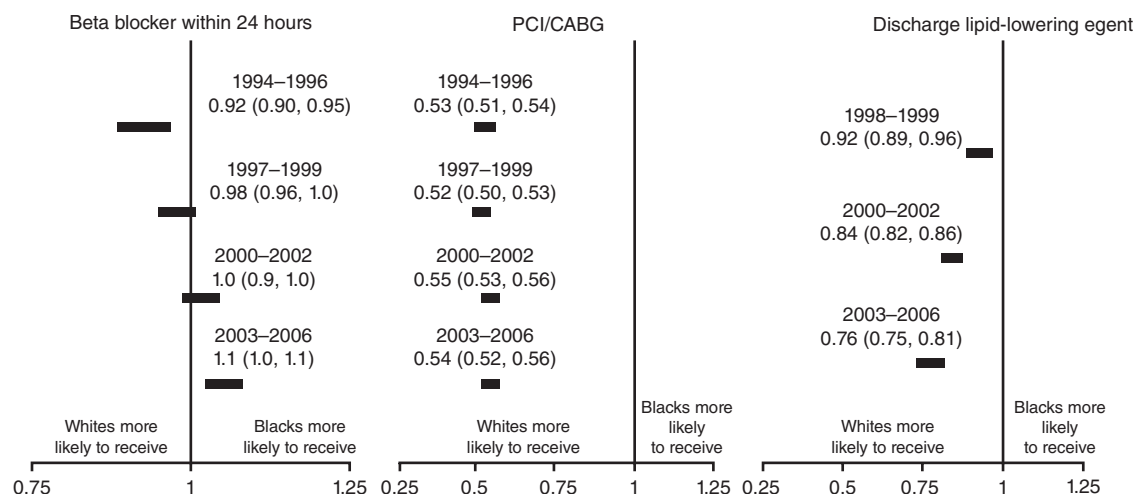


FIGURE 16e-4 Racial differences in guideline-based treatments for acute myocardial infarction (AMI). The reference population consisted of 2,515,106 patients with AMI admitted to U.S. hospitals between July 1990 and December 2006. CABG, coronary artery bypass grafting; PCI, percutaneous coronary intervention. (From ED Peterson et al. *Am Heart J* 156:1045, 2008.)

- **Patient-level factors:** These include patients' mistrust of the health care system leading to refusal of services, poor adherence to treatment, and delay in seeking care.

A more detailed analysis of these root causes is presented below.

Health System Factors • HEALTH SYSTEM COMPLEXITY Even among persons who are insured and educated and who have a high degree of health literacy, navigating the U.S. health care system can be complicated and confusing. Some individuals may be at higher risk for receiving substandard care because of their difficulty navigating the system's complexities. These individuals may include those from cultures unfamiliar with the Western model of health care delivery, those with limited English proficiency, those with low health literacy, and those who are mistrustful of the health care system. These individuals may have difficulty knowing how and where to go for a referral to a specialist; how to prepare for a procedure such as a colonoscopy; or how to follow up on an abnormal test result such as a mammogram. Since people of color in the United States tend to be overrepresented among the groups listed above, the inherent complexity of navigating the health care system has been seen as a root cause for racial/ethnic disparities in health care.

OTHER HEALTH SYSTEM FACTORS Racial/ethnic disparities are due not only to differences in care provided within hospitals but also to where and

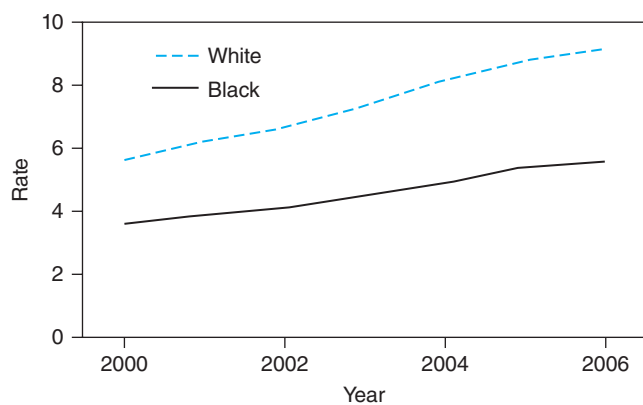


FIGURE 16e-5 Racial trends in age-adjusted total knee replacement in Medicaid enrollees from 2000 to 2006. The reference population consisted of Medicaid part A enrollees ≥ 65 years of age who were not members of a managed-care plan. (From the Centers for Disease Control and Prevention, 2009.)

from whom minorities receive their care; i.e., certain specific providers, geographic regions, or hospitals are lower-performing on certain aspects of quality. For example, one study showed that 25% of hospitals cared for 90% of black Medicare patients in the United States and that these hospitals tended to have lower performance scores on certain quality measures than other hospitals. That said, health systems generally are not well prepared to measure, report, and intervene to reduce disparities in care. Few hospitals or health plans stratify their quality data by race/ethnicity or language to measure disparities, and even fewer use data of this type to develop disparity-targeted interventions. Similarly, despite regulations concerning the need for professional interpreters, research demonstrates that many health care organizations and providers fail to routinely provide this service for patients with limited English proficiency. Despite the link between limited English proficiency and health-care quality and safety, few providers or institutions monitor performance for patients in these areas.

Provider-Level Factors • PROVIDER-PATIENT COMMUNICATION Significant evidence highlights the impact of sociocultural factors, race, ethnicity, and limited English proficiency on health and clinical care. Health care professionals frequently care for diverse populations with varied perspectives, values, beliefs, and behaviors regarding health and well-being. The differences include variations in the recognition of symptoms, thresholds for seeking care, comprehension of management strategies, expectations of care (including preferences for or against diagnostic and therapeutic procedures), and adherence to preventive measures and medications. In addition, sociocultural differences between patient and provider influence communication and clinical decision-making and are especially pertinent: evidence clearly links provider-patient communication to improved patient satisfaction, regimen adherence, and better health outcomes (Fig. 16e-6). Thus, when sociocultural differences between patient and provider are not appreciated, explored, understood, or communicated effectively during the medical encounter, patient dissatisfaction, poor adherence, poorer health outcomes, and racial/ethnic disparities in care may result.

A survey of 6722 Americans ≥ 18 years of age is particularly relevant to this important link between provider-patient communication and health outcomes. Whites, blacks, Hispanics/Latinos, and Asian Americans who had made a medical visit in the past 2 years were asked whether they had trouble understanding their doctors; whether they felt the doctors did not listen; and whether they had medical questions they were afraid to ask. The survey found that 19% of all patients experienced one or more of these problems, yet whites experienced them 16% of the time as opposed to 23% of the time for blacks, 33% for Hispanics/Latinos, and 27% for Asian Americans (Fig. 16e-7).

How do we link communication to outcomes?



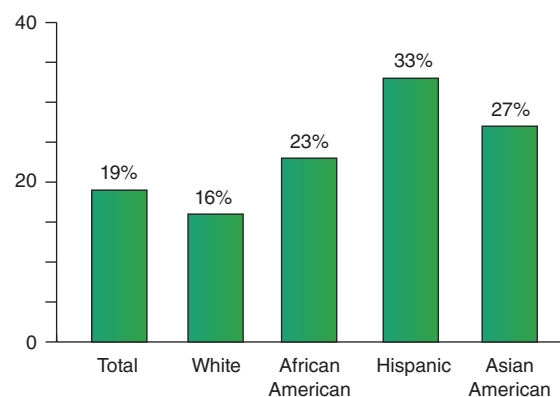
FIGURE 16e-6 The link between effective communication and patient satisfaction, adherence, and health outcomes. (From the Institute of Medicine: *Unequal Treatment: Confronting Racial and Ethnic Disparities in Health Care*. Washington, DC, National Academy Press, 2002.)

In addition, in the setting of even a minimal language barrier, provider–patient communication without an interpreter is recognized as a major challenge to effective health care delivery. These communication barriers for patients with limited English proficiency lead to frequent misunderstanding of diagnosis, treatment, and follow-up plans; inappropriate use of medications; lack of informed consent for surgical procedures; high rates of serious adverse events; and a lower-quality health care experience than is provided to patients who speak fluent English. Physicians who have access to trained interpreters report a significantly higher quality of patient–physician communication than physicians who use other methods. Communication issues related to discordant language disproportionately affect minorities and likely contribute to racial/ethnic disparities in health care.

CLINICAL DECISION-MAKING Theory and research suggest that variations in clinical decision-making may contribute to racial and ethnic disparities in health care. Two factors are central to this process: clinical uncertainty and stereotyping.

First, a doctor’s decision-making process is nested in *clinical uncertainty*. Doctors depend on inferences about severity based on what they understand about illness and the information obtained from the patient. A doctor caring for a patient whose symptoms he or she has difficulty understanding and whose “signals”—the set of clues and indications that physicians rely on to make clinical decisions—are hard to read may make a decision different from the one that would be made

Percent of adults with one or more communication problems*



Base: Adults with health care visit in past two years

*Problems include understanding doctor, feeling doctor listened, had questions but did not ask.

FIGURE 16e-7 Communication difficulties with physicians, by race/ethnicity. The reference population consisted of 6722 Americans ≥ 18 years of age who had made a medical visit in the previous 2 years and were asked whether they had had trouble understanding their doctors, whether they felt that the doctors had not listened, and whether they had had medical questions they were afraid to ask. (From the Commonwealth Fund Health Care Quality Survey, 2001.)

for another patient who presents with exactly the same clinical condition. Given that the expression of symptoms may differ among cultural and racial groups, doctors—the overwhelming majority of whom are white—may understand symptoms best when expressed by patients of their own racial/ethnic groups. The consequence is that white patients may be treated differently from minority patients. Differences in clinical decisions can arise from this mechanism even when the doctor has the same regard for each patient (i.e., is not prejudiced).

Second, the literature on social cognitive theory highlights how natural tendencies to stereotype may influence clinical decision-making. *Stereotyping* can be defined as the way in which people use social categories (e.g., race, gender, age) in acquiring, processing, and recalling information about others. Faced with enormous information loads and the need to make many decisions, people often subconsciously simplify the decision-making process and lessen cognitive effort by using “categories” or “stereotypes” that bundle information into groups or types that can be processed more quickly. Although functional, stereotyping can be systematically biased, as people are automatically classified into social categories based on dimensions such as *race*, *gender*, and *age*. Many people may not be aware of their attitudes, may not consciously endorse specific stereotypes, and paradoxically may consider themselves egalitarian and not prejudiced.

Stereotypes may be strongly influenced by the messages presented consciously and unconsciously in society. For instance, if the media and our social/professional contacts tend to present images of minorities as being less educated, more violent, and nonadherent to health care recommendations, these impressions may generate stereotypes that unnaturally and unjustly impact clinical decision-making. As signs of racism, classism, gender bias, and ageism are experienced (consciously or unconsciously) in our society, stereotypes may be created that impact the way doctors manage patients from these groups. On the basis of training or practice location, doctors may develop certain perceptions about race/ethnicity, culture, and class that may evolve into stereotypes. For example, many medical students and residents are often trained—and minorities cared for—in academic health centers or public hospitals located in socioeconomically disadvantaged areas. As a result, doctors may begin to equate certain races and ethnicities with specific health beliefs and behaviors (e.g., “these patients” engage in risky behaviors, “those patients” tend to be noncompliant) that are more associated with the social environment (e.g., poverty) than with a patient’s racial/ethnic background or cultural traditions. This “conditioning” phenomenon may also be operative if doctors are faced with certain racial/ethnic patient groups who frequently do not choose aggressive forms of diagnostic or therapeutic intervention. The result over time may be that doctors begin to believe that “these patients” do not like invasive procedures; thus they may not offer these procedures as options. A wide range of studies have documented the potential for provider biases to contribute to racial/ethnic disparities in health care. For example, one study measured physicians’ unconscious (or implicit) biases and showed that these were related to differences in decisions to provide thrombolysis for a hypothetical black or white patient with a myocardial infarction.

It is important to differentiate stereotyping from prejudice and discrimination. *Prejudice* is a conscious prejudgment of individuals that may lead to disparate treatment, and *discrimination* is conscious and intentional disparate treatment. All individuals *stereotype* subconsciously, yet, if left unquestioned, these subconscious assumptions may lead to lower-quality care for certain groups because of differences in clinical decision-making or differences in communication and patient-centeredness. For example, one study tested physicians’ unconscious racial/ethnic biases and showed that patients perceived more biased physicians as being less patient-centered in their communication. What is particularly salient is that stereotypes tend to be activated most in environments where the individual is stressed, multitasking, and under time pressure—the hallmarks of the clinical encounter.

Patient-Level Factors Lack of trust has become a major concern for many health care institutions today. For example, an IOM report, *To Err Is Human: Building a Safer Health System*, documented alarming

rates of medical errors that made patients feel vulnerable and less trustful of the U.S. health care system. The increased media and academic attention to problems related to quality of care (and of disparities themselves) has clearly diminished trust in doctors and nurses.

Trust is a crucial element in the therapeutic alliance between patient and health care provider. It facilitates open communication and is directly correlated with adherence to the physician's recommendations and the patient's satisfaction. In other words, patients who mistrust their health care providers are less satisfied with the care they receive, and mistrust of the health care system greatly affects patients' use of services. Mistrust can also result in inconsistent care, "doctor-shopping," self-medication, and an increased demand by patients for referrals and diagnostic tests.

On the basis of historic factors such as discrimination, segregation, and medical experimentation, blacks may be especially mistrustful of providers. The exploitation of blacks by the U.S. Public Health Service during the Tuskegee syphilis study from 1932 to 1972 left a legacy of mistrust that persists even today among this population. Other populations, including Native Americans/Alaskan Natives, Hispanics/Latinos, and Asian Americans, also harbor significant mistrust of the health care system. A national Kaiser Family Foundation survey of 3884 individuals found that 36% of Hispanics and 35% of blacks (compared with 15% of whites) felt they had been treated unfairly in the health care system in the past because of their race/ethnicity. Perhaps even more alarming, 65% of blacks and 58% of Hispanics (compared with 22% of whites) were afraid of being treated unfairly in the future on that basis (Fig. 16e-8).

This mistrust may contribute to wariness in accepting or following recommendations, undergoing invasive procedures, or participating in clinical research, and these choices, in turn, may lead to misunderstanding and the perpetuation of stereotypes among health professionals.

KEY RECOMMENDATIONS TO ADDRESS RACIAL/ETHNIC DISPARITIES IN HEALTH CARE

The publication *Unequal Treatment* provides a series of recommendations to address racial and ethnic disparities in health care, focusing on a broad set of stakeholders. These recommendations include *health system interventions*, *provider interventions*, *patient interventions*, and *general recommendations*, which are described in more detail below.

Health System Interventions • COLLECTION AND REPORTING OF DATA ON HEALTH CARE ACCESS AND USE, BY PATIENTS' RACE/ETHNICITY *Unequal Treatment* found that the appropriate systems to track and monitor racial and ethnic disparities in health care are lacking and that less is known about the disparities affecting minority groups other than African

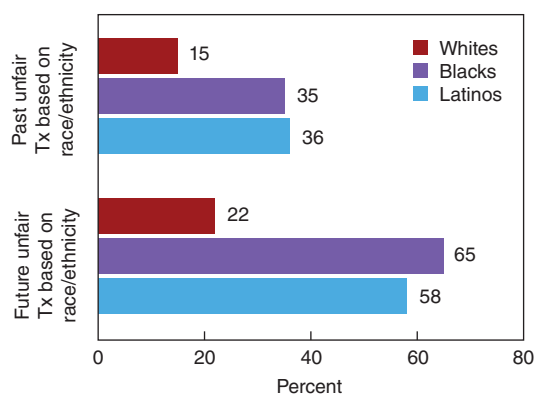


FIGURE 16e-8 Patient perspectives regarding unfair treatment (Tx) based on race/ethnicity. The reference population consisted of 3884 individuals surveyed about how fairly they had been treated in the health care system in the past and how fairly they felt they would be treated in the future on the basis of their race/ethnicity. (From *Race, Ethnicity & Medical Care: A Survey of Public Perceptions and Experiences*. Kaiser Family Foundation, 2005.)

Americans (Hispanics, Asian Americans, Pacific Islanders, Native Americans, and Alaskan Natives). For instance, only in the mid-1980s did the Medicare database begin to collect data on patient groups outside the standard categories of "white," "black," and "other." Federal, private, and state-supported data-collection efforts are scattered and unsystematic, and many health care systems and hospitals still do not collect data on the race, ethnicity, or primary language of enrollees or patients. A survey by Regenstein and Sickler in 2006 found that 78% of 501 U.S. hospitals collected information on race, 50% collected data on ethnicity, and 50% collected data on primary language. However, the information was not collected by standard categories or collection methods and thus was of questionable accuracy. Surveys by America's Health Insurance Plans in 2003 and 2006 showed that the proportion of enrollees in plans that collected race/ethnicity data of some type increased from 54% to 67%; however, the total percentage of plan enrollees whose race/ethnicity and language are recorded is still much lower than these figures.

ENCOURAGEMENT OF THE USE OF EVIDENCE-BASED GUIDELINES AND QUALITY IMPROVEMENT *Unequal Treatment* highlights the subjectivity of clinical decision-making as a potential cause of racial and ethnic disparities in health care by describing how clinicians—despite the existence of well-delineated practice guidelines—may offer (consciously or unconsciously) different diagnostic and therapeutic options to different patients on the basis of their race or ethnicity. Therefore, the widespread adoption and implementation of evidence-based guidelines is a key recommendation in eliminating disparities. For instance, evidence-based guidelines are now available for the management of diabetes, HIV/AIDS, cardiovascular diseases, cancer screening and management, and asthma—all areas where significant disparities exist. As part of ongoing quality-improvement efforts, particular attention should be paid to the implementation of evidence-based guidelines for all patients, regardless of their race and ethnicity.

SUPPORT FOR THE USE OF LANGUAGE INTERPRETATION SERVICES IN THE CLINICAL SETTING As described previously, a lack of efficient and effective interpreter services in a health care system can lead to patient dissatisfaction, to poor comprehension and adherence, and thus to ineffective/lower-quality care for patients with limited English proficiency. *Unequal Treatment's* recommendation to support the use of interpretation services has clear implications for delivery of quality health care by improving doctors' ability to communicate effectively with these patients.

INCREASES IN THE PROPORTION OF UNDERREPRESENTED MINORITIES IN THE HEALTH CARE WORKFORCE Data for 2004 from the Association of American Medical Colleges indicate that, of the 72.4% of U.S. physicians whose race and ethnicity are known, Hispanics make up 2.8%, blacks 3.3%, and Native American and Alaskan Natives 0.3%. Furthermore, U.S. national data show that minorities (excluding Asians) compose just 7.5% of medical school faculty. In addition, minority faculty in 2007 were more likely to be at or below the rank of assistant professor, while whites composed the highest proportion of full professors. Despite representing ~26% of the U.S. population (a number projected to almost double by 2050), minority students are still underrepresented in medical schools. In 2007, matriculants to U.S. medical schools were 7.2% Latino, 6.4% African American, 0.2% Native Hawaiian or Other Pacific Islander, and 0.3% Native American or Alaskan Native. These percentages have decreased or remained the same since 2007. It will be difficult to develop a diverse health-care workforce that can meet the needs of an increasingly diverse population without dramatic changes in the racial and ethnic composition of medical student bodies.

Provider Interventions • INTEGRATION OF CROSS-CULTURAL EDUCATION INTO THE TRAINING OF ALL HEALTH CARE PROFESSIONALS The goal of cross-cultural education is to improve providers' ability to understand, communicate with, and care for patients from diverse backgrounds. Such education focuses on enhancing awareness of sociocultural influences on health beliefs and behaviors and on building skills to facilitate understanding and management of these factors in the medical encounter. Cross-cultural education includes curricula on health care disparities,

use of interpreters, and effective communication and negotiation across cultures. These curricula can be incorporated into health-professions training in medical schools, residency programs, and nursing schools and can be offered as a component of continuing education. Despite the importance of this area of education and the attention it has attracted from medical education accreditation bodies, a national survey of senior resident physicians by Weissman and colleagues found that up to 28% felt unprepared to deal with cross-cultural issues, including caring for patients who have religious beliefs that may affect treatment, patients who use complementary medicine, patients who have health beliefs at odds with Western medicine, patients who mistrust the health care system, and new immigrants. In a study at one medical school, 70% of fourth-year students felt inadequately prepared to care for patients with limited English proficiency. Efforts to incorporate cross-cultural education into medical education will contribute to improving communication and to providing a better quality of care for all patients.

INCORPORATION OF TEACHING ON THE IMPACT OF RACE, ETHNICITY, AND CULTURE ON CLINICAL DECISION-MAKING *Unequal Treatment* and more recent studies found that stereotyping by health care providers can lead to disparate treatment based on a patient's race or ethnicity. The Liaison Committee on Medical Education, which accredits medical schools, issued a directive that medical education should include instruction on how a patient's race, ethnicity, and culture might unconsciously impact communication and clinical decision-making.

Patient Interventions Difficulty navigating the health care system and obtaining access to care can be a hindrance to all populations, particularly to minorities. Similarly, lack of empowerment or involvement in the medical encounter by minorities can be a barrier to care. Patients need to be educated on how to navigate the health care system and how best to access care. Interventions should be used to increase patients' participation in treatment decisions.

General Recommendations • INCREASE AWARENESS OF RACIAL/ETHNIC DISPARITIES IN HEALTH CARE Efforts to raise awareness of racial/ethnic health care disparities have done little for the general public but have been fairly successful among physicians, according to a Kaiser Family Foundation report. In 2006, nearly 6 in 10 people surveyed believed that blacks received the same quality of care as whites, and 5 in 10 believed that Latinos received the same quality of care as whites. These estimates are similar to findings in a 1999 survey. Despite this lack of awareness, most people believed that all Americans deserve quality care, regardless of their background. In contrast, the level of awareness among physicians has risen sharply. In 2002, the majority (69%) of physicians said that the health care system "rarely or never" treated people unfairly on the basis of their racial/ethnic background. In 2005, less than one-quarter (24%) of physicians disagreed with the statement that "minority patients generally receive lower-quality care than white patients." Increasing awareness of racial and ethnic health disparities among health care professionals and the public is an important first step in addressing these disparities. The ultimate goals are to generate discourse and to mobilize action to address disparities at multiple levels, including health policy makers, health systems, and the community.

CONDUCT FURTHER RESEARCH TO IDENTIFY SOURCES OF DISPARITIES AND PROMISING INTERVENTIONS While the literature that formed the basis for the findings reported and recommendations made in *Unequal Treatment* provided significant evidence for racial and ethnic disparities, additional research is needed in several areas. First, most of the literature on disparities focuses on black-versus-white differences; much less is known about the experiences of other minority groups. Improving the ability to collect racial and ethnic patient data should facilitate this process. However, in instances where the necessary systems are not yet in place, racial and ethnic patient data may be collected prospectively in the setting of clinical or health services research to more fully elucidate disparities for other populations. Second, much of the literature on disparities to date has focused on defining areas in which these disparities exist, but less has been done to identify the multiple factors that

contribute to the disparities or to test interventions to address these factors. There is clearly a need for research that identifies promising practices and solutions to disparities.

IMPLICATIONS FOR CLINICAL PRACTICE

Individual health care providers can do several things in the clinical encounter to address racial and ethnic disparities in health care.

Be Aware that Disparities Exist Increasing awareness of racial and ethnic disparities among health care professionals is an important first step in addressing disparities in health care. Only with greater awareness can care providers be attuned to their behavior in clinical practice and thus monitor that behavior and ensure that all patients receive the highest quality of care, regardless of race, ethnicity, or culture.

Practice Culturally Competent Care Previous efforts have been made to teach clinicians about the attitudes, values, beliefs, and behaviors of certain cultural groups—the key practice “dos and don’ts” in caring for “the Hispanic patient” or the “Asian patient,” for example. In certain situations, learning about a particular local community or cultural group, with a goal of following the principles of community-oriented primary care, can be helpful; when broadly and uncritically applied, however, this approach can actually lead to stereotyping and oversimplification of culture, without respect for its complexity.

Cultural competence has thus evolved from merely learning information and making assumptions about patients on the basis of their backgrounds to focusing on the development of skills that follow the principles of patient-centered care. *Patient-centeredness* encompasses the qualities of compassion, empathy, and responsiveness to the needs, values, and expressed preferences of the individual patient. *Cultural competence* aims to take things a step further by expanding the repertoire of knowledge and skills classically defined as “patient-centered” to include those that are especially useful in cross-cultural interactions (and that, in fact, are vital in all clinical encounters). This repertoire includes effectively using interpreter services, eliciting the patient's understanding of his or her condition, assessing decision-making preferences and the role of family, determining the patient's views about biomedicine versus complementary and alternative medicine, recognizing sexual and gender issues, and building trust. For example, while it is important to understand all patients' beliefs about health, it may be particularly crucial to understand the health beliefs of patients who come from a different culture or have a different health care experience. With the individual patient as teacher, the physician can adjust his or her practice style to meet the patient's specific needs.

Avoid Stereotyping Several strategies can allow health care providers to counteract, both systemically and individually, the normal tendency to stereotype. For example, when racially/ethnically/culturally/socially diverse teams in which each member is given equal power are assembled and are tasked to achieve a common goal, a sense of camaraderie develops and prevents the development of stereotypes based on race/ethnicity, gender, culture, or class. Thus, health care providers should aim to gain experiences working with and learning from a diverse set of colleagues. In addition, simply being aware of the operation of social cognitive factors allows providers to actively check up on or monitor their behavior. Physicians can constantly reevaluate to ensure that they are offering the same things, in the same ways, to all patients. Understanding one's own susceptibility to stereotyping—and how disparities may result—is essential in providing equitable, high-quality care to all patients.

Work to Build Trust Patients' mistrust of the health care system and of health care providers impacts multiple facets of the medical encounter, with effects ranging from decreased patient satisfaction to delayed care. Although the historic legacy of discrimination can never be erased, several steps can be taken to build trust with patients and to address disparities. First, providers must be aware that mistrust exists and is more prevalent among minority populations, given the history of discrimination in the United States and other countries. Second, providers must reassure patients that they come first, that everything possible

will be done to ensure that they always get the best care available, and that their caregivers will serve as their advocates. Third, interpersonal skills and communication techniques that demonstrate honesty, openness, compassion, and respect on the part of the health care provider are essential tools in dismantling mistrust. Finally, patients indicate that trust is built when there is shared, participatory decision-making and the provider makes a concerted effort to understand the patient's background. When the doctor-patient relationship is reframed as one of solidarity, the patient's sense of vulnerability can be transformed into one of trust. The successful elimination of disparities requires trust-building interventions and strengthening of this relationship.

CONCLUSION

The issue of racial and ethnic disparities in health care has gained national prominence, both with the release of the IOM report *Unequal Treatment* and with more recent articles that have confirmed their

persistence and explored their root causes. Furthermore, another influential IOM report, *Crossing the Quality Chasm*, has highlighted the importance of equity—i.e., no variations in quality of care due to personal characteristics, including race and ethnicity—as a central principle of quality. Current efforts in health care reform and transformation, including a greater focus on value (high-quality care and cost-control), will sharpen the nation's focus on the care of populations who experience low-quality, costly care. Addressing disparities will become a major focus, and there will be many obvious opportunities for interventions to eliminate them. Greater attention to addressing the root causes of disparities will improve the care provided to all patients, not just those who belong to racial and ethnic minorities.

ACKNOWLEDGMENT

The authors thank Marina Cervantes for her contributions to this chapter.

17e Ethical Issues in Clinical Medicine

Bernard Lo, Christine Grady

Twenty-first-century physicians face novel ethical dilemmas that can be perplexing and emotionally draining. For example, electronic medical records, handheld personal devices, and provision of care by interdisciplinary teams all hold the promise of more coordinated and comprehensive care but also raise new concerns about confidentiality, appropriate boundaries of the doctor–patient relationship, and responsibility. **Chapter 1** puts the practice of medicine into a professional and historical context. The current chapter presents approaches and principles that physicians can use to address the ethical issues they encounter in their work. Physicians make ethical judgments about clinical situations every day. Traditional professional codes and ethical principles provide instructive guidance for physicians but need to be interpreted and applied to each situation. Physicians need to be prepared for lifelong learning about ethical issues and dilemmas as well as about new scientific and clinical developments. When struggling with difficult ethical issues, physicians may need to reevaluate their basic convictions, tolerate uncertainty, and maintain their integrity while respecting the opinions of others. Discussing perplexing ethical issues with other members of the health care team, ethics consultation services, or the hospital ethics committee can clarify issues and reveal strategies for resolution, including improving communication and dealing with strong or conflicting emotions.

APPROACHES TO ETHICAL PROBLEMS

Several approaches may be useful for resolving ethical issues. Among these approaches are those based on ethical principles, virtue ethics, professional oaths, and personal values. These various sources of guidance encompass precepts that may conflict in a particular case, leaving the physician in a quandary. In a diverse society, different individuals may turn to different sources of moral guidance. In addition, general moral precepts often need to be interpreted and applied in the context of a particular clinical situation. When facing an ethical challenge, physicians should articulate their concerns and reasoning, discuss and listen to the views of others involved in the case, and call on available resources as needed. Through these efforts, physicians can gain deeper insight into the ethical issues they face and often can reach mutually acceptable resolutions to complex problems.

ETHICAL PRINCIPLES

Ethical principles can serve as general guidelines to help physicians determine the right thing to do.

Respecting Patients Physicians should always treat patients with respect, which entails understanding patients' goals, communicating effectively, obtaining informed and voluntary consent for interventions, respecting informed refusals, and protecting confidentiality. Different clinical goals and approaches are often feasible, and interventions can cause both benefit and harm. Individuals place different values on health and medical care and weigh the benefits and risks of medical interventions differently. Generally, the values and informed choices of patients should be respected.

OBTAINING INFORMED CONSENT To help patients make informed decisions, physicians should discuss with them the nature of the proposed care; the alternatives; and the risks, benefits, and likely consequences of each option. Informed consent involves more than obtaining signatures on consent forms. Physicians should promote shared decision-making by educating patients, answering their questions, making recommendations, and helping them deliberate. Patients can be overwhelmed by medical jargon, needlessly complicated explanations, or the provision of too much information at once. Patients can make informed decisions only if they receive honest and understandable information. Competent, informed patients may refuse recommended interventions and choose among reasonable alternatives. If patients

cannot give consent in an emergency and if delay of treatment while surrogates are contacted will place their lives or health in peril, treatment can be given without informed consent. People are presumed to want such emergency care unless they have previously indicated otherwise.

Respect for patients does not entitle the patients to insist on any care they want. Physicians are not obligated to provide interventions that have no physiologic rationale, that have already failed, or that are contrary to evidence-based practice recommendations, good clinical judgment, or public policies. National policies and laws also dictate certain decisions—e.g., allocating cadaveric organs for transplantation and, in most states, prohibiting physician-assisted suicide.

Physicians should disclose and discuss relevant and accurate information about diagnosis, prognosis, and treatment options. To help patients cope with bad news, doctors can adjust the pace of disclosure, offer empathy and hope, provide emotional support, and call on other resources such as spiritual care or social work. Physicians may be tempted to withhold a serious diagnosis, misrepresent it by using ambiguous terms, or limit discussions of prognosis or risks for fear that certain information will make patients anxious or depressed. Providing honest information about clinical situations preserves patients' autonomy and trust and promotes sound communication with patients and colleagues. However, patients may choose not to receive such information, asking surrogates to make decisions on their behalf, as is common with serious diagnoses in some traditional cultures.

AVOIDING DECEPTION Health care providers sometimes consider using lies or deception to obtain benefits for patients. *Lying* refers to statements known to be false and intended to mislead the listener. *Deception* includes statements and actions intended to mislead the listener, whether or not they are literally true. For example, a physician might sign a disability form for a patient who does not meet disability criteria. Although motivated by a desire to help the patient, such deception is ethically problematic because it undermines physicians' credibility and trustworthiness.

MAINTAINING CONFIDENTIALITY Maintaining confidentiality is essential in respecting patients' autonomy and privacy, encourages them to seek treatment and to discuss problems candidly, and prevents discrimination. However, confidentiality may be overridden to prevent serious harm to third parties or to the patient. Exceptions to confidentiality are justified if the risk is serious and probable, if there are no less restrictive measures by which to avert risk, if the adverse effects of overriding confidentiality are minimized, and if these adverse effects are deemed acceptable by society. For example, the law requires physicians to report cases of tuberculosis, sexually transmitted infection, elder or child abuse, and domestic violence.

CARING FOR PATIENTS WHO LACK DECISION-MAKING CAPACITY Some patients are not able to make informed decisions because of unconsciousness, dementia, delirium, or other medical conditions. Although only the courts have the legal authority to determine that a patient is incompetent for making medical decisions, in practice, physicians determine when patients lack the capacity to make health care decisions and arrange for surrogates to make decisions for them, without involving the courts. Patients with decision-making capacity can express a choice and appreciate the medical situation; the nature of the proposed care; the alternatives; and the risks, benefits, and consequences of each alternative. Their choices should be consistent with their values and not the result of delusions or hallucinations. Psychiatrists may help assess decision-making capacity in difficult cases. When impairments are fluctuating or reversible, decisions should be postponed if possible until the patient recovers decision-making capacity.

If a patient lacks decision-making capacity, physicians should ask: Who is the appropriate surrogate, and what would the patient want done? Patients may designate someone to serve as their health care proxy or to assume durable power of attorney for health care; such choices should be respected. (See **Chap. 10** for further details about advance care planning.) Unless a patient without decision-making capacity has previously designated a health care proxy, physicians

usually ask family members to serve as surrogates. Many patients want family members as surrogates, and family members generally have the patient's best interests at heart. Statutes in most U.S. states delineate a prioritized list of relatives who may serve as surrogates if the patient has not designated a proxy. Surrogates' decisions should be guided by the patient's values, goals, and previously expressed preferences. However, it may be appropriate to override previous preferences in favor of the patient's current best interests if an intervention is highly likely to provide a significant benefit, if previous statements do not fit the situation well, or if the patient expressed a desire for the surrogate to have leeway in making decisions.

ACTING IN THE BEST INTERESTS OF PATIENTS Respect for patients is broader than respecting their autonomy to make informed choices about their medical care and promoting shared decision-making. Physicians should also be compassionate and dedicated and should act in the best interests of their patients.

The principle of *beneficence* requires physicians to act for the patient's benefit. Patients typically lack medical expertise and may be vulnerable because of their illness. They rely on physicians to provide sound recommendations and to promote their well-being. Physicians encourage such trust and have a fiduciary duty to act in the best interests of the patient, which should prevail over the physicians' own self-interest or the interests of third parties, such as hospitals or insurers. Physicians' fiduciary obligations contrast sharply with business relationships, which are characterized by "let the buyer beware," not by reliance and trust. A related principle, "first do no harm," forbids physicians to provide ineffective interventions or to act without due care. Although often cited, this precept alone provides only limited guidance because many beneficial interventions pose serious risks. Physicians should prevent unnecessary harm by recommending interventions that maximize benefit and minimize harm.

MANAGING CONFLICTS BETWEEN RESPECTING PATIENTS AND ACTING IN THEIR BEST INTERESTS Conflicts can arise when patients' refusal of interventions thwarts their own goals for care or causes them serious harm. For example, if a young woman with asthma refuses mechanical ventilation for reversible respiratory failure, simple acceptance of this decision by the physician, in the name of respecting autonomy, is morally constricted. Physicians should elicit patients' expectations and concerns, correct their misunderstandings, and try to persuade them to accept beneficial therapies. If disagreements persist after such efforts, patients' informed choices and views of their own best interests should prevail. While refusing recommended care does not render a patient incompetent, it may lead the physician to probe further to ensure that the patient has the capacity to make informed decisions.

Acting Justly The principle of *justice* provides guidance to physicians about how to ethically treat patients and to make decisions about allocating important resources, including their own time. *Justice* in a general sense means fairness: people should receive what they deserve. In addition, it is important to act consistently in cases that are similar in ethically relevant ways. Otherwise, decisions may be arbitrary, biased, and unfair. *Justice* forbids discrimination in health care based on race, religion, gender, sexual orientation, or other personal characteristics (Chap. 16e).

Justice also requires that limited health care resources be allocated fairly. Universal access to medically needed health care remains an unrealized moral aspiration in the United States and much of the rest of the world. Patients without health insurance often cannot afford health care and lack access to safety-net services. Even among insured patients, insurers may deny coverage for interventions recommended by the physician. In this situation, physicians should advocate for patients and try to help them obtain needed care. Doctors might consider—or patients might request—the use of deception to obtain such benefits. However, avoiding deception is a basic ethical guideline that sets limits on advocating for patients.

Allocation of health care resources is unavoidable because these resources are limited. Ideally, decisions about allocation are made at the level of public policy, with physician input. For example, the United Network for Organ Sharing (www.unos.org) provides criteria

for allocating scarce organs. Ad hoc resource allocation at the bedside is problematic because it may be inconsistent, unfair, and ineffective. Physicians have an important role, however, in avoiding unnecessary interventions. Evidence-based lists of tests and procedures that physicians and patients should question and discuss were developed through the recent initiative *Choosing Wisely* (<http://www.choosingwisely.org/>). At the bedside, physicians should act as patient advocates within constraints set by society, reasonable insurance coverage, and evidence-based practice. For example, if a patient's insurer has a higher copayment for nonformulary drugs, it still may be reasonable for physicians to advocate for nonformulary products for good reasons (e.g., when the formulary drugs are ineffective or not tolerated).

VIRTUE ETHICS

Virtue ethics focuses on physicians' character and qualities, with the expectation that doctors will cultivate such virtues as compassion, dedication, altruism, humility, and integrity. Proponents argue that, if such characteristics become ingrained, they help guide physicians in novel situations. Moreover, merely following ethical precepts or principles without these virtues leads to uncaring doctor-patient relationships.

PROFESSIONAL OATHS AND CODES

Professional oaths and codes are useful guides for physicians. Most physicians take oaths at student white-coat ceremonies and at medical school graduation, and many are members of professional societies that have professional codes. Members of the profession pledge to the public and to their patients that they will be guided by the principles and values in these oaths or codes. Oaths and codes focus physicians on ethical ideals rather than on daily pragmatic concerns. However, professional oaths and codes—even the Hippocratic tradition—have been criticized for lack of patient or public input and the limited role given to patients in making decisions.

PERSONAL VALUES

Personal values, cultural traditions, and religious beliefs are important sources of personal morality that help physicians address ethical issues and cope with the moral distress they may experience in practice. While essential, personal morality is a limited ethical guide in clinical practice. Physicians have role-specific ethical obligations that go beyond their obligations as good people, including the duties to obtain informed consent and maintain confidentiality discussed earlier in this chapter. Furthermore, in a culturally and religiously diverse world, patients and colleagues have personal moral beliefs that commonly differ from their physicians'.

Claims of Conscience Some physicians have conscientious objections to providing or referring patients for certain treatments, such as contraception. While physicians should not be asked to violate deeply held moral beliefs or religious convictions, patients need to receive medically appropriate, timely care. Institutions such as clinics and hospitals have a collective duty to provide care that patients need while making reasonable attempts to accommodate health care workers' conscientious objections—for example, by arranging for another professional to provide the service in question. Patients seeking a relationship with a doctor or health care institution should be notified in advance of any conscientious objections to the provision of specific interventions. Since patients commonly must select providers for insurance purposes, switching providers when a specific service is needed would be burdensome. There are important limits on claims of conscience. Health care workers may not insist that patients receive unwanted medical interventions and may not refuse to treat patients because of their race, ethnicity, national origin, gender, or religion. Such discrimination is illegal and violates the physician's duty to respect patients.

Moral Distress Physicians and other health care providers may experience moral distress when they feel they know the ethically correct action to take in a particular situation but are constrained by institutional policies, limited resources, or a position subordinate to the ultimate decision-maker. Moral distress can lead to anger, anxiety,

frustration, fatigue, and work dissatisfaction. Discussing complex clinical situations with colleagues and seeking assistance with difficult decisions helps to alleviate moral distress, as does a healthy work environment characterized by open communication and mutual respect.

These various sources of guidance contain precepts that may conflict in a particular case, leaving the physician in a quandary. In a diverse society, different individuals may turn to different sources of moral guidance. In addition, general moral precepts often need to be interpreted and applied in the context of a particular clinical situation. When facing an ethical challenge, physicians should articulate their concerns and reasoning, discuss and listen to the views of others involved in the case, and call on available resources as needed. Through these efforts, physicians can gain deeper insight into the ethical issues they face and often reach mutually acceptable resolutions to complex problems.

CHANGES IN ORGANIZATION AND DELIVERY OF CARE

Recent changes in the organization and delivery of health care have led to new ethical challenges for physicians.

WORK HOURS

The Accreditation Council for Graduate Medical Education requires medical students and residents to observe work-hour limitations, which are intended to help prevent physician burnout, reduce mistakes, and create a better balance between work and private life. In addition to continuing controversy over their effectiveness, some ethical concerns are raised by work-hour limitations. One concern is that physicians may develop a shift-worker mentality that undermines their dedication to the well-being of patients. Forced handoffs to colleagues may actually increase the risk of errors, and inflexibility can be detrimental. In some cases, trainees could provide an irreplaceable benefit to a patient or family by going beyond work-hour limits, especially if there is rapport with the patient or family that is not easily transferred to another provider. For example, a resident may want to discuss decisions about life-sustaining interventions or to comfort a family member over a patient's death (Chap. 10). Thus strict adherence to work-hour limits is not always consistent with the ideal of acting for the good of the patient and with compassion. Exceptions to work-hour limits, however, should remain exceptions and should not be allowed to undercut work-hour policies.

TEAM-BASED CARE

Physicians' roles are changing as care is increasingly provided by multidisciplinary teams. The traditional hierarchy in which the physician is the "captain of the ship" may be inappropriate, particularly in areas such as prevention, disease management and its coordination, and patient education. Physicians should respect team members and acknowledge the expertise of those from other disciplinary backgrounds. Team-based care promises to provide more comprehensive and higher-quality care. However, regular communication and planning are critical to avoid diffusion of responsibility and to ensure that someone is accountable for the completion of patient-care tasks.

CLINICAL PRACTICE GUIDELINES

The increasing use of evidence-based practice guidelines and benchmarking of performance raises the overall quality of care. However, practice guideline recommendations may be inappropriate for an individual patient, while another option may provide substantially greater benefits. In such situations, physicians' duty to act in the patient's best interests should take priority over benefits to society as a whole. Physicians need to understand practice guidelines, to recognize situations in which exceptions might be reasonable, and to be prepared to justify an exception.

GLOBAL HEALTH FIELD EXPERIENCES



With the growing importance of and interest in global health, many physicians and trainees work overseas for various periods. Typically, physicians gain valuable experience while providing service to patients in need. Such arrangements, however, can

raise ethical challenges—for example, because of differences in beliefs about health and illness, expectations regarding health care and the physician's role, standards of clinical practice, and norms for disclosure of serious diagnoses. Additional dilemmas arise if visiting physicians take on responsibilities beyond their level of training or if donated drugs and equipment are not appropriate to local needs. Visiting physicians and trainees should exercise due diligence in obtaining needed information about the cultural and clinical practices in the host community, should work closely with local professionals and team members, and should be explicit about their skills, knowledge, and limits. In addition, these arrangements can pose risks. The visiting physician may face personal risk from infectious disease or motor vehicle accident. The host institution incurs administrative and supervisory costs. Advance preparation for these possibilities minimizes harm, distress, and misunderstanding.

PHYSICIANS' USE OF SOCIAL MEDIA

Increasingly, physicians use social and electronic media to share information with patients and other providers. Social networking may be especially useful in reaching young or otherwise hard-to-access patients. However, the use of social media, including blogs, social networks, and websites, raises ethical challenges and can have harmful consequences if not approached prudently. Injudicious use of social media can pose risks to patient confidentiality, expose patients to intimate details of physicians' personal lives, cross professional boundaries, and jeopardize therapeutic relationships. Posts may be considered unprofessional and lead to adverse consequences for a provider's reputation, safety, or even employment, especially if they express frustration or anger over work incidents, disparage patients or colleagues, use offensive or discriminatory language, reveal highly personal information, or picture a physician intoxicated, using illegal drugs, or in sexually suggestive poses. Physicians should remember that, in the absence of highly restrictive privacy settings, postings on the Internet in general and on social networking sites in particular are usually permanent and may be accessible to the public, their employers, and their patients. Physicians should separate professional from personal websites, social networking accounts, and blogs and should follow guidelines developed by institutions and professional societies on using social media to communicate with patients.

CONFLICTS OF INTEREST

Acting in patients' best interests may conflict with the physician's self-interest or the interests of third parties such as insurers or hospitals. From an ethical viewpoint, patients' interests should be paramount. Even the appearance of a conflict of interest may undermine trust in the profession.

FINANCIAL INCENTIVES

Health care providers may be offered financial incentives to improve the quality or efficiency of care. Such pay-for-performance incentives, however, could lead physicians to avoid sicker patients with more complicated cases or to focus on benchmarked outcomes even when such a focus is not in the best interests of an individual patient. In contrast, fee-for-service payments offer physicians incentives to order more interventions than may be necessary or to refer patients to laboratory or imaging facilities in which they have a financial stake. Regardless of financial incentives, physicians should recommend available care that is in the patient's best interests—no more and no less.

RELATIONSHIPS WITH PHARMACEUTICAL COMPANIES

Financial relationships between physicians and industry are increasingly scrutinized. Gifts from drug and device companies may create an inappropriate risk of undue influence, induce subconscious feelings of reciprocity, impair public trust, and increase the cost of health care. Many academic medical centers have banned drug-company gifts of pens, notepads, and meals to physicians. Under the new Physician Payment Sunshine Act, companies must disclose publicly the names of physicians to whom they have made payments or transferred material goods and the amount of those payments or transfers. The challenge will be to distinguish payments for scientific consulting and research

contracts—which are consistent with professional and academic missions and should be encouraged—from those for promotional speaking and consulting whose goal is to increase sales of company products.

OCCUPATIONAL RISKS

Some health care workers, fearing fatal occupational infections, have refused to care for certain patients, such as those with HIV infection or severe acute respiratory syndrome (SARS). Such fears about personal safety need to be acknowledged. Health care institutions should reduce occupational risk by providing proper training, protective equipment, and supervision. To fulfill their mission of helping patients, physicians should provide appropriate care within their clinical expertise, despite some personal risk.

RESPONSE TO MEDICAL ERRORS

Errors are inevitable in clinical medicine, and some errors cause serious adverse events that harm patients. Most errors are caused by lapses of attention or flaws in the system of delivering health care; only a few result from blameworthy individual behavior (**Chaps. 3 and 12e**). Physicians and students may fear that disclosing errors will damage their careers. However, patients appreciate being told when an error occurs, receiving an apology, and being informed about efforts to prevent similar errors in the future. Physicians and health care institutions show respect for patients by disclosing errors, offering appropriate compensation for harm done, and using errors as opportunities to improve the quality of care. Overall, patient safety is more likely to be improved through a quality improvement approach to errors rather than a punitive one except in cases of gross incompetence, physician impairment, boundary violations, or repeated violations of standard procedures.

LEARNING CLINICAL SKILLS

Physicians' interest in learning, which fosters the long-term goal of benefiting future patients, may conflict with the short-term goal of providing optimal care to current patients. When trainees learn to carry out procedures on patients, they lack the proficiency of experienced physicians, and patients may experience inconvenience,

discomfort, longer procedures, or even increased risk. Although patients' consent for trainee participation in their care is always important, it is particularly important for intimate examinations, such as pelvic, rectal, breast, and testicular examinations, and for invasive procedures. To ensure patients' cooperation, some care providers introduce students as physicians or do not tell patients that trainees will be performing procedures. Such misrepresentation undermines trust, may lead to more elaborate deception, and makes it difficult for patients to make informed choices about their care. Patients should be told who is providing care and how trainees are supervised. Most patients, when informed, allow trainees to play an active role in their care.

PHYSICIAN IMPAIRMENT

Physicians may hesitate to intervene when colleagues impaired by alcohol abuse, drug abuse, or psychiatric or medical illness place patients at risk. However, society relies on physicians to regulate themselves. If colleagues of an impaired physician do not take steps to protect patients, no one else may be in a position to do so.

ETHICAL ISSUES IN CLINICAL RESEARCH

Clinical research is essential to translate scientific discoveries into beneficial tests and therapies for patients. However, clinical research raises ethical concerns because participants face inconvenience and risks in research that is designed not specifically to benefit them but rather to advance scientific knowledge. Ethical guidelines for researchers require them to obtain informed and voluntary consent from participants and approval from an institutional review board, which determines that risks to participants are acceptable and have been minimized and recommends appropriate additional protections when research includes vulnerable participants. Physicians may be involved as clinical research investigators or may be in a position to refer or recommend clinical trial participation to their patients. Physicians should be critical consumers of clinical research results and keep up with advances that change standards of practice. Courses and guidance on the ethics of clinical research are widely available.

SECTION 1 PAIN

18 Pain: Pathophysiology and Management

James P. Rathmell, Howard L. Fields

The province of medicine is to preserve and restore health and to relieve suffering. Understanding pain is essential to both of these goals. Because pain is universally understood as a signal of disease, it is the most common symptom that brings a patient to a physician's attention. The function of the pain sensory system is to protect the body and maintain homeostasis. It does this by detecting, localizing, and identifying potential or actual tissue-damaging processes. Because different diseases produce characteristic patterns of tissue damage, the quality, time course, and location of a patient's pain lend important diagnostic clues. It is the physician's responsibility to provide rapid and effective pain relief.

THE PAIN SENSORY SYSTEM

Pain is an unpleasant sensation localized to a part of the body. It is often described in terms of a penetrating or tissue-destructive process (e.g., stabbing, burning, twisting, tearing, squeezing) and/or of a bodily or emotional reaction (e.g., terrifying, nauseating, sickening). Furthermore, any pain of moderate or higher intensity is accompanied by anxiety and the urge to escape or terminate the feeling. These properties illustrate the duality of pain: it is both sensation and emotion. When it is acute, pain is characteristically associated with behavioral arousal and a stress response consisting of increased blood pressure, heart rate, pupil diameter, and plasma cortisol levels. In addition, local muscle contraction (e.g., limb flexion, abdominal wall rigidity) is often present.

PERIPHERAL MECHANISMS

The Primary Afferent Nociceptor A peripheral nerve consists of the axons of three different types of neurons: primary sensory afferents, motor neurons, and sympathetic postganglionic neurons (Fig. 18-1). The cell bodies of primary sensory afferents are located in the dorsal root ganglia within the vertebral foramina. The primary afferent axon has two branches: one projects centrally into the spinal cord and the other projects peripherally to innervate tissues. Primary afferents are classified by their diameter, degree of myelination, and conduction velocity. The largest diameter afferent fibers, A-beta ($A\beta$), respond maximally to light touch and/or moving stimuli; they are present primarily in nerves that innervate the skin. In normal individuals, the activity of these fibers does not produce pain. There are two other classes of primary afferent nerve fibers: the small diameter myelinated A-delta ($A\delta$) and the unmyelinated (C) axons (Fig. 18-1). These fibers are present in nerves to the skin and to deep somatic and visceral

structures. Some tissues, such as the cornea, are innervated only by $A\delta$ and C fiber afferents. Most $A\delta$ and C fiber afferents respond maximally only to intense (painful) stimuli and produce the subjective experience of pain when they are electrically stimulated; this defines them as *primary afferent nociceptors (pain receptors)*. The ability to detect painful stimuli is completely abolished when conduction in $A\delta$ and C fiber axons is blocked.

Individual primary afferent nociceptors can respond to several different types of noxious stimuli. For example, most nociceptors respond to heat; intense cold; intense mechanical distortion, such as a pinch; changes in pH, particularly an acidic environment; and application of chemical irritants including adenosine triphosphate (ATP), serotonin, bradykinin, and histamine.

Sensitization When intense, repeated, or prolonged stimuli are applied to damaged or inflamed tissues, the threshold for activating primary afferent nociceptors is lowered, and the frequency of firing is higher for all stimulus intensities. Inflammatory mediators such as bradykinin, nerve-growth factor, some prostaglandins, and leukotrienes contribute to this process, which is called *sensitization*. Sensitization occurs at the level of the peripheral nerve terminal (*peripheral sensitization*) as well as at the level of the dorsal horn of the spinal cord (*central sensitization*). Peripheral sensitization occurs in damaged or inflamed tissues, when inflammatory mediators activate intracellular signal transduction in nociceptors, prompting an increase in the production, transport, and membrane insertion of chemically gated and voltage-gated ion channels. These changes increase the excitability of nociceptor terminals and lower their threshold for activation by mechanical, thermal, and chemical stimuli. Central sensitization occurs when activity, generated by nociceptors during inflammation, enhances the excitability of nerve cells in the dorsal horn of the spinal cord. Following injury and resultant sensitization, normally innocuous stimuli can produce pain (termed *allodynia*). Sensitization is a clinically important process that contributes to tenderness, soreness, and *hyperalgesia* (increased pain intensity in response to the same noxious stimulus; e.g., moderate pressure causes severe pain). A striking example of sensitization is

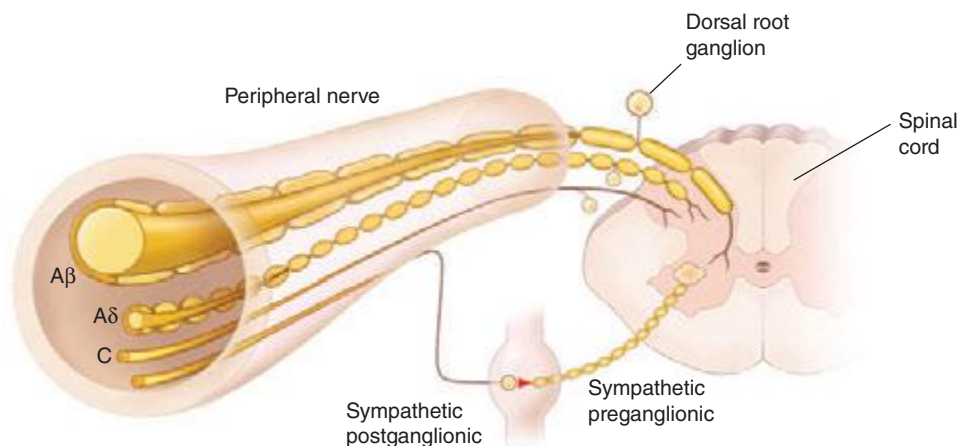


FIGURE 18-1 Components of a typical cutaneous nerve. There are two distinct functional categories of axons: primary afferents with cell bodies in the dorsal root ganglion, and sympathetic postganglionic fibers with cell bodies in the sympathetic ganglion. Primary afferents include those with large-diameter myelinated ($A\beta$), small-diameter myelinated ($A\delta$), and unmyelinated (C) axons. All sympathetic postganglionic fibers are unmyelinated.

sunburned skin, in which severe pain can be produced by a gentle slap on the back or a warm shower.

Sensitization is of particular importance for pain and tenderness in deep tissues. Viscera are normally relatively insensitive to noxious mechanical and thermal stimuli, although hollow viscera do generate significant discomfort when distended. In contrast, when affected by a disease process with an inflammatory component, deep structures such as joints or hollow viscera characteristically become exquisitely sensitive to mechanical stimulation.

A large proportion of A δ and C fiber afferents innervating viscera are completely insensitive in normal noninjured, noninflamed tissue. That is, they cannot be activated by known mechanical or thermal stimuli and are not spontaneously active. However, in the presence of inflammatory mediators, these afferents become sensitive to mechanical stimuli. Such afferents have been termed *silent nociceptors*, and their characteristic properties may explain how, under pathologic conditions, the relatively insensitive deep structures can become the source of severe and debilitating pain and tenderness. Low pH, prostaglandins, leukotrienes, and other inflammatory mediators such as bradykinin play a significant role in sensitization.

Nociceptor-Induced Inflammation Primary afferent nociceptors also have a neuroeffector function. Most nociceptors contain polypeptide mediators that are released from their peripheral terminals when they are activated (Fig. 18-2). An example is substance P, an 11-amino-acid peptide. Substance P is released from primary afferent nociceptors and has multiple biologic activities. It is a potent vasodilator, degranulates mast cells, is a chemoattractant for leukocytes, and increases the production and release of inflammatory mediators. Interestingly, depletion of substance P from joints reduces the severity of experimental arthritis. Primary afferent nociceptors are not simply passive messengers of threats to tissue injury but also play an active role in tissue protection through these neuroeffector functions.

CENTRAL MECHANISMS

The Spinal Cord and Referred Pain The axons of primary afferent nociceptors enter the spinal cord via the dorsal root. They terminate in the dorsal horn of the spinal gray matter (Fig. 18-3). The terminals of primary afferent axons contact spinal neurons that transmit the pain signal to brain sites involved in pain perception. When primary afferents are activated by noxious stimuli, they release neurotransmitters from their terminals that excite the spinal cord neurons. The major neurotransmitter released is glutamate, which rapidly excites dorsal horn neurons. Primary afferent nociceptor terminals also release peptides, including substance P and calcitonin gene-related peptide, which produce a slower and longer-lasting excitation of the dorsal horn neurons. The axon of each primary afferent contacts many spinal neurons, and each spinal neuron receives convergent inputs from many primary afferents.

The convergence of sensory inputs to a single spinal pain-transmission neuron is of great importance because it underlies the phenomenon of referred pain. All spinal neurons that receive input from the viscera and deep musculoskeletal structures also receive input from the skin. The convergence patterns are determined by the spinal segment of the dorsal root ganglion that supplies the afferent innervation of a structure. For example, the afferents that supply the central diaphragm are derived from the third and fourth cervical dorsal root ganglia. Primary afferents with cell bodies in these same ganglia supply the skin of the shoulder and lower neck. Thus, sensory inputs from both the shoulder skin and the central diaphragm converge on pain-transmission neurons in the third and fourth cervical spinal segments. *Because of this convergence and the fact that the spinal neurons are most often activated by inputs from the skin, activity evoked in spinal neurons by input from deep structures is mislocalized by the patient to a place that roughly corresponds with the region of skin innervated by the same spinal segment.* Thus, inflammation near the central diaphragm is often reported as shoulder discomfort. This spatial displacement of pain sensation from the site of the injury that produces it is known as *referred pain*.

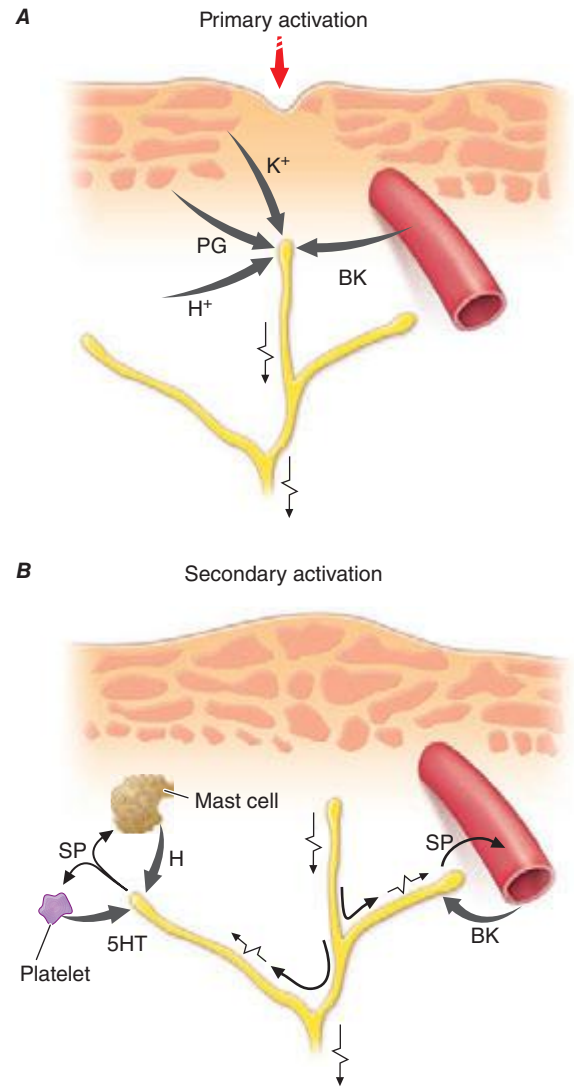


FIGURE 18-2 Events leading to activation, sensitization, and spread of sensitization of primary afferent nociceptor terminals. **A.** Direct activation by intense pressure and consequent cell damage. Cell damage induces lower pH (H^+) and leads to release of potassium (K^+) and to synthesis of prostaglandins (PG) and bradykinin (BK). Prostaglandins increase the sensitivity of the terminal to bradykinin and other pain-producing substances. **B.** Secondary activation. Impulses generated in the stimulated terminal propagate not only to the spinal cord but also into other terminal branches where they induce the release of peptides, including substance P (SP). Substance P causes vasodilation and neurogenic edema with further accumulation of bradykinin (BK). Substance P also causes the release of histamine (H) from mast cells and serotonin (5HT) from platelets.

Ascending Pathways for Pain A majority of spinal neurons contacted by primary afferent nociceptors send their axons to the contralateral thalamus. These axons form the contralateral spinothalamic tract, which lies in the anterolateral white matter of the spinal cord, the lateral edge of the medulla, and the lateral pons and midbrain. The spinothalamic pathway is crucial for pain sensation in humans. Interruption of this pathway produces permanent deficits in pain and temperature discrimination.

Spinothalamic tract axons ascend to several regions of the thalamus. There is tremendous divergence of the pain signal from these thalamic sites to several distinct areas of the cerebral cortex that subservise different aspects of the pain experience (Fig. 18-4). One of the thalamic projections is to the somatosensory cortex. This projection mediates the purely sensory aspects of pain, i.e., its location, intensity, and quality. Other thalamic neurons project to cortical regions that are linked

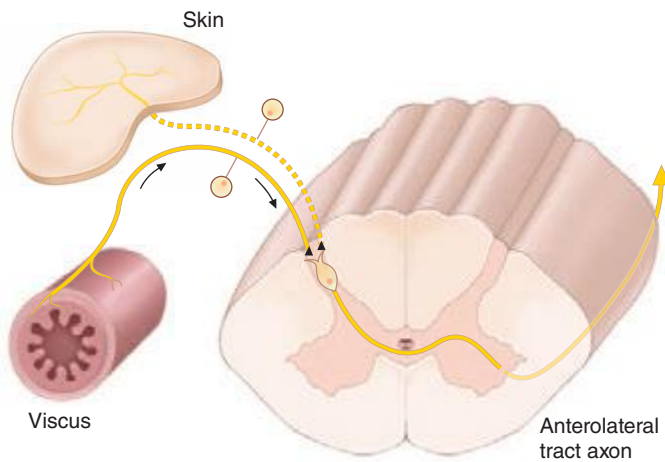


FIGURE 18-3 The convergence-projection hypothesis of referred pain. According to this hypothesis, visceral afferent nociceptors converge on the same pain-projection neurons as the afferents from the somatic structures in which the pain is perceived. The brain has no way of knowing the actual source of input and mistakenly “projects” the sensation to the somatic structure.

to emotional responses, such as the cingulate gyrus and other areas of the frontal lobes, including the insular cortex. These pathways to the frontal cortex subserve the affective or unpleasant emotional dimension of pain. This affective dimension of pain produces suffering and exerts potent control of behavior. Because of this dimension, fear is

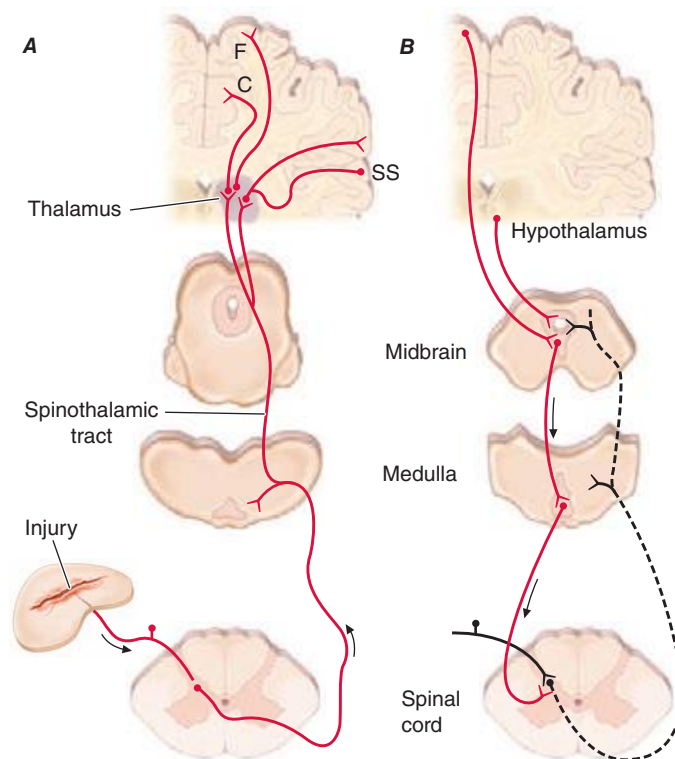


FIGURE 18-4 Pain transmission and modulatory pathways.

A. Transmission system for nociceptive messages. Noxious stimuli activate the sensitive peripheral ending of the primary afferent nociceptor by the process of transduction. The message is then transmitted over the peripheral nerve to the spinal cord, where it synapses with cells of origin of the major ascending pain pathway, the spinothalamic tract. The message is relayed in the thalamus to the anterior cingulate (C), frontal insular (F), and somatosensory cortex (SS). **B.** Pain-modulation network. Inputs from frontal cortex and hypothalamus activate cells in the midbrain that control spinal pain-transmission cells via cells in the medulla.

a constant companion of pain. As a consequence, injury or surgical lesions to areas of the frontal cortex activated by painful stimuli can diminish the emotional impact of pain while largely preserving the individual’s ability to recognize noxious stimuli as painful.

PAIN MODULATION

The pain produced by injuries of similar magnitude is remarkably variable in different situations and in different individuals. For example, athletes have been known to sustain serious fractures with only minor pain, and Beecher’s classic World War II survey revealed that many soldiers in battle were unbothered by injuries that would have produced agonizing pain in civilian patients. Furthermore, even the suggestion that a treatment will relieve pain can have a significant analgesic effect (the *placebo effect*). On the other hand, many patients find even minor injuries (such as venipuncture) frightening and unbearable, and the expectation of pain can induce pain even without a noxious stimulus. The suggestion that pain will worsen following administration of an inert substance can increase its perceived intensity (the *nocebo effect*).

The powerful effect of expectation and other psychological variables on the perceived intensity of pain is explained by brain circuits that modulate the activity of the pain-transmission pathways. One of these circuits has links to the hypothalamus, midbrain, and medulla, and it selectively controls spinal pain-transmission neurons through a descending pathway (Fig. 18-4).

Human brain-imaging studies have implicated this pain-modulating circuit in the pain-relieving effect of attention, suggestion, and opioid analgesic medications (Fig. 18-5). Furthermore, each of the component structures of the pathway contains opioid receptors and is sensitive to the direct application of opioid drugs. In animals, lesions of this descending modulatory system reduce the analgesic effect of systemically administered opioids such as morphine. Along with the opioid receptor, the component nuclei of this pain-modulating circuit contain endogenous opioid peptides such as the enkephalins and β -endorphin.

The most reliable way to activate this endogenous opioid-mediated modulating system is by suggestion of pain relief or by intense emotion directed away from the pain-causing injury (e.g., during severe threat or an athletic competition). In fact, pain-relieving endogenous opioids are released following surgical procedures and in patients given a placebo for pain relief.

Pain-modulating circuits can enhance as well as suppress pain. Both pain-inhibiting and pain-facilitating neurons in the medulla project to and control spinal pain-transmission neurons. Because pain-transmission neurons can be activated by modulatory neurons, it is theoretically possible to generate a pain signal with no peripheral noxious stimulus. In fact, human functional imaging studies have demonstrated increased activity in this circuit during migraine headaches. A central circuit that facilitates pain could account for the finding that pain can be induced by suggestion or enhanced by expectation and provides a framework for understanding how psychological factors can contribute to chronic pain.

NEUROPATHIC PAIN

Lesions of the peripheral or central nociceptive pathways typically result in a loss or impairment of pain sensation. Paradoxically, damage to or dysfunction of these pathways can also produce pain. For example, damage to peripheral nerves, as occurs in diabetic neuropathy, or to primary afferents, as in herpes zoster infection, can result in pain that is referred to the body region innervated by the damaged nerves. Pain may also be produced by damage to the central nervous system (CNS), for example, in some patients following trauma or vascular injury to the spinal cord, brainstem, or thalamic areas that contain central nociceptive pathways. Such neuropathic pains are often severe and are typically resistant to standard treatments for pain.

Neuropathic pain typically has an unusual burning, tingling, or electric shock-like quality and may be triggered by very light touch. These features are rare in other types of pain. On examination, a sensory deficit is characteristically present in the area of the patient’s pain. *Hyperpathia*, a greatly exaggerated pain sensation to innocuous

Pattern of Brain Activity During Placebo Analgesia

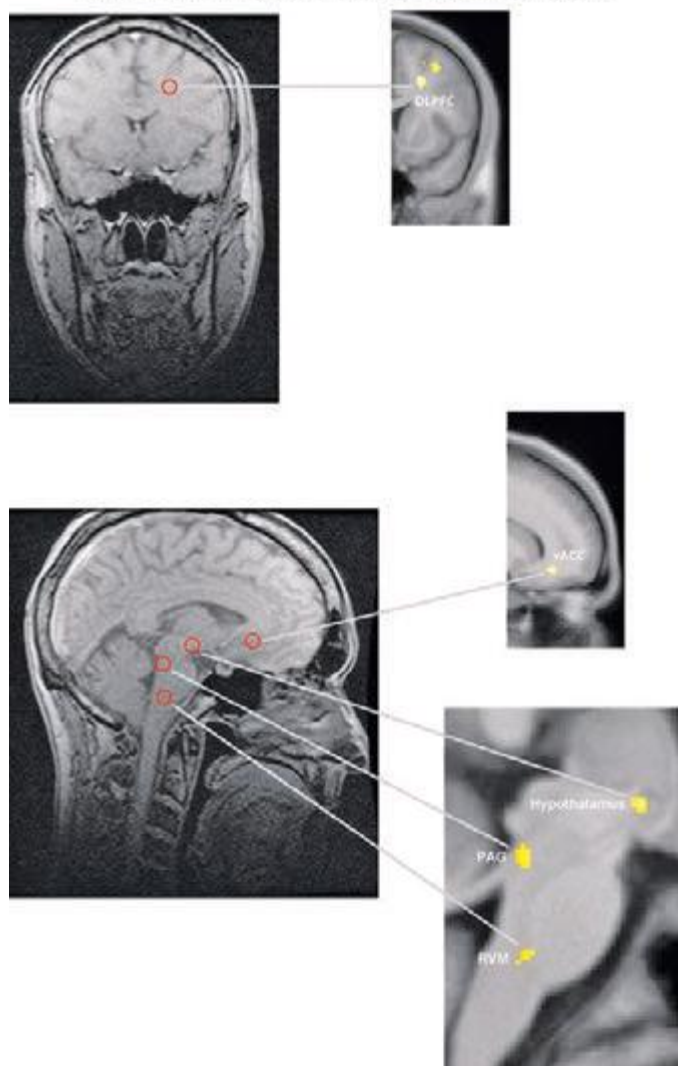


FIGURE 18-5 Functional magnetic resonance imaging (fMRI) demonstrates placebo-enhanced brain activity in anatomic regions correlating with the opioidergic descending pain control system. *Top panel:* Frontal fMRI image shows placebo-enhanced brain activity in the dorsal lateral prefrontal cortex (DLPFC). *Bottom panel:* Sagittal fMRI images show placebo-enhanced responses in the rostral anterior cingulate cortex (rACC), the rostral ventral medullae (RVM), the periaqueductal gray (PAG) area, and the hypothalamus. The placebo-enhanced activity in all areas was reduced by naloxone, demonstrating the link between the descending opioidergic system and the placebo analgesic response. (Adapted with permission from F Eippert et al: *Neuron* 63:533, 2009.)

or mild nociceptive stimuli, is also characteristic of neuropathic pain; patients often complain that the very lightest moving stimulus evokes exquisite pain (allodynia). In this regard, it is of clinical interest that a topical preparation of 5% lidocaine in patch form is effective for patients with postherpetic neuralgia who have prominent allodynia.

A variety of mechanisms contribute to neuropathic pain. As with sensitized primary afferent nociceptors, damaged primary afferents, including nociceptors, become highly sensitive to mechanical stimulation and may generate impulses in the absence of stimulation. Increased sensitivity and spontaneous activity are due, in part, to an increased concentration of sodium channels in the damaged nerve fiber. Damaged primary afferents may also develop sensitivity to norepinephrine. Interestingly, spinal cord pain-transmission neurons cut off from their normal input may also become spontaneously active.

Thus, both CNS and peripheral nervous system hyperactivity contribute to neuropathic pain.

Sympathetically Maintained Pain Patients with peripheral nerve injury occasionally develop spontaneous pain in the region innervated by the nerve. This pain is often described as having a burning quality. The pain typically begins after a delay of hours to days or even weeks and is accompanied by swelling of the extremity, periarticular bone loss, and arthritic changes in the distal joints. The pain may be relieved by a local anesthetic block of the sympathetic innervation to the affected extremity. Damaged primary afferent nociceptors acquire adrenergic sensitivity and can be activated by stimulation of the sympathetic outflow. This constellation of spontaneous pain and signs of sympathetic dysfunction following injury has been termed *complex regional pain syndrome* (CRPS). When this occurs after an identifiable nerve injury, it is termed CRPS type II (also known as posttraumatic neuralgia or, if severe, *causalgia*). When a similar clinical picture appears without obvious nerve injury, it is termed CRPS type I (also known as *reflex sympathetic dystrophy*). CRPS can be produced by a variety of injuries, including fractures of bone, soft tissue trauma, myocardial infarction, and stroke (**Chap. 446**). CRPS type I typically resolves with symptomatic treatment; however, when it persists, detailed examination often reveals evidence of peripheral nerve injury. Although the pathophysiology of CRPS is poorly understood, the pain and the signs of inflammation, when acute, can be rapidly relieved by blocking the sympathetic nervous system. This implies that sympathetic activity can activate undamaged nociceptors when inflammation is present. Signs of sympathetic hyperactivity should be sought in patients with post-traumatic pain and inflammation and no other obvious explanation.

TREATMENT ACUTE PAIN

The ideal treatment for any pain is to remove the cause; thus, while treatment can be initiated immediately, efforts to establish the underlying etiology should always proceed as treatment begins. Sometimes, treating the underlying condition does not immediately relieve pain. Furthermore, some conditions are so painful that rapid and effective analgesia is essential (e.g., the postoperative state, burns, trauma, cancer, or sickle cell crisis). Analgesic medications are a first line of treatment in these cases, and all practitioners should be familiar with their use.

ASPIRIN, ACETAMINOPHEN, AND NONSTEROIDAL ANTI-INFLAMMATORY AGENTS (NSAIDs)

These drugs are considered together because they are used for similar problems and may have a similar mechanism of action (**Table 18-1**). All these compounds inhibit cyclooxygenase (COX), and, except for acetaminophen, all have anti-inflammatory actions, especially at higher dosages. They are particularly effective for mild to moderate headache and for pain of musculoskeletal origin.

Because they are effective for these common types of pain and are available without prescription, COX inhibitors are by far the most commonly used analgesics. They are absorbed well from the gastrointestinal tract and, with occasional use, have only minimal side effects. With chronic use, gastric irritation is a common side effect of aspirin and NSAIDs and is the problem that most frequently limits the dose that can be given. Gastric irritation is most severe with aspirin, which may cause erosion and ulceration of the gastric mucosa leading to bleeding or perforation. Because aspirin irreversibly acetylates platelet cyclooxygenase and thereby interferes with coagulation of the blood, gastrointestinal bleeding is a particular risk. Older age and history of gastrointestinal disease increase the risks of aspirin and NSAIDs. In addition to the well-known gastrointestinal toxicity of NSAIDs, nephrotoxicity is a significant problem for patients using these drugs on a chronic basis. Patients at risk for renal insufficiency, particularly those with significant contraction of their intravascular volume as occurs with chronic diuretic use or

TABLE 18-1 DRUGS FOR RELIEF OF PAIN

Generic Name	Dose, mg	Interval	Comments					
Nonnarcotic analgesics: usual doses and intervals								
Acetylsalicylic acid	650 PO	q4h	Enteric-coated preparations available					
Acetaminophen	650 PO	q4h	Side effects uncommon					
Ibuprofen	400 PO	q4–6h	Available without prescription					
Naproxen	250–500 PO	q12h	Delayed effects may be due to long half-life					
Fenoprofen	200 PO	q4–6h	Contraindicated in renal disease					
Indomethacin	25–50 PO	q8h	Gastrointestinal side effects common					
Ketorolac	15–60 IM/IV	q4–6h	Available for parenteral use					
Celecoxib	100–200 PO	q12–24h	Useful for arthritis					
Valdecoxib	10–20 PO	q12–24h	Removed from U.S. market in 2005					
Generic Name	Parenteral Dose, mg	PO Dose, mg	Comments					
Narcotic analgesics: usual doses and intervals								
Codeine	30–60 q4h	30–60 q4h	Nausea common					
Oxycodone	—	5–10 q4–6h	Usually available with acetaminophen or aspirin					
Morphine	5 q4h	30 q4h						
Morphine sustained release	—	15–60 bid to tid	Oral slow-release preparation					
Hydromorphone	1–2 q4h	2–4 q4h	Shorter acting than morphine sulfate					
Levorphanol	2 q6–8h	4 q6–8h	Longer acting than morphine sulfate; absorbed well PO					
Methadone	5–10 q6–8h	5–20 q6–8h	Delayed sedation due to long half-life; therapy should not be initiated with >40 mg/d, and dose escalation should be made no more frequently than every 3 days					
Meperidine	50–100 q3–4h	300 q4h	Poorly absorbed PO; normeperidine is a toxic metabolite; routine use of this agent is not recommended					
Butorphanol	—	1–2 q4h	Intranasal spray					
Fentanyl	25–100 µg/h	—	72-h transdermal patch					
Buprenorphine	5–20 µg/h	—	7-day transdermal patch					
Buprenorphine	0.3 q6–8h	—	Parenteral administration					
Tramadol	—	50–100 q4–6h	Mixed opioid/adrenergic action					
Generic Name	Uptake Blockade		Sedative Potency	Anticholinergic Potency	Orthostatic Hypotension	Cardiac Arrhythmia	Ave. Dose, mg/d	Range, mg/d
	5-HT	NE						
Antidepressants^a								
Doxepin	++	+	High	Moderate	Moderate	Less	200	75–400
Amitriptyline	++++	++	High	Highest	Moderate	Yes	150	25–300
Imipramine	++++	++	Moderate	Moderate	High	Yes	200	75–400
Nortriptyline	+++	++	Moderate	Moderate	Low	Yes	100	40–150
Desipramine	+++	++++	Low	Low	Low	Yes	150	50–300
Venlafaxine	+++	++	Low	None	None	No	150	75–400
Duloxetine	+++	+++	Low	None	None	No	40	30–60
Generic Name	PO Dose, mg	Interval	Generic Name	PO Dose, mg	Interval			
Anticonvulsants and antiarrhythmics^a								
Phenytoin	300	daily/qhs	Clonazepam	1	q6h			
Carbamazepine	200–300	q6h	Gabapentin ^b	600–1200	q8h			
Oxcarbazepine	300	bid	Pregabalin	150–600	bid			

^aAntidepressants, anticonvulsants, and antiarrhythmics have not been approved by the U.S. Food and Drug Administration (FDA) for the treatment of pain. ^bGabapentin in doses up to 1800 mg/d is FDA approved for postherpetic neuralgia.

Abbreviations: 5-HT, serotonin; NE, norepinephrine.

acute hypovolemia, should be monitored closely. NSAIDs can also increase blood pressure in some individuals. Long-term treatment with NSAIDs requires regular blood pressure monitoring and treatment if necessary. Although toxic to the liver when taken in high doses, acetaminophen rarely produces gastric irritation and does not interfere with platelet function.

The introduction of parenteral forms of NSAIDs, ketorolac and diclofenac, extends the usefulness of this class of compounds in the management of acute severe pain. Both agents are sufficiently potent and rapid in onset to supplant opioids for many patients with acute severe headache and musculoskeletal pain.

There are two major classes of COX: COX-1 is constitutively expressed, and COX-2 is induced in the inflammatory state. COX-2-selective drugs have similar analgesic potency and produce less gastric irritation than the nonselective COX inhibitors. The use of COX-2-selective drugs does not appear to lower the risk of nephrotoxicity compared to nonselective NSAIDs. On the other hand, COX-2-selective drugs offer a significant benefit in the management of acute postoperative pain because they do not affect blood coagulation. Nonselective COX inhibitors are usually contraindicated postoperatively because they impair platelet-mediated blood clotting and are thus associated with increased bleeding at the

operative site. COX-2 inhibitors, including celecoxib (Celebrex), are associated with increased cardiovascular risk. It appears that this is a class effect of NSAIDs, excluding aspirin. These drugs are contraindicated in patients in the immediate period after coronary artery bypass surgery and should be used with caution in elderly patients and those with a history of or significant risk factors for cardiovascular disease.

OPIOID ANALGESICS

Opioids are the most potent pain-relieving drugs currently available. Of all analgesics, they have the broadest range of efficacy and provide the most reliable and effective method for rapid pain relief. Although side effects are common, most are reversible: nausea, vomiting, pruritus, and constipation are the most frequent and bothersome side effects. Respiratory depression is uncommon at standard analgesic doses, but can be life-threatening. Opioid-related side effects can be reversed rapidly with the narcotic antagonist naloxone. Many physicians, nurses, and patients have a certain trepidation about using opioids that is based on an exaggerated fear of addiction. In fact, there is a vanishingly small chance of patients becoming addicted to narcotics as a result of their appropriate medical use. The physician should not hesitate to use opioid analgesics in patients with acute severe pain. Table 18-1 lists the most commonly used opioid analgesics.

Opioids produce analgesia by actions in the CNS. They activate pain-inhibitory neurons and directly inhibit pain-transmission neurons. Most of the commercially available opioid analgesics act at the same opioid receptor (μ -receptor), differing mainly in potency, speed of onset, duration of action, and optimal route of administration. Some side effects are due to accumulation of nonopioid metabolites that are unique to individual drugs. One striking example of this is normeperidine, a metabolite of meperidine. At higher doses of meperidine, typically greater than 1 g/d, accumulation of normeperidine can produce hyperexcitability and seizures that are not reversible with naloxone. Normeperidine accumulation is increased in patients with renal failure.

The most rapid pain relief is obtained by intravenous administration of opioids; relief with oral administration is significantly slower. Because of the potential for respiratory depression, patients with any form of respiratory compromise must be kept under close observation following opioid administration; an oxygen-saturation monitor may be useful, but only in a setting where the monitor is under constant surveillance. Opioid-induced respiratory depression is typically accompanied by sedation and a reduction in respiratory rate. A fall in oxygen saturation represents a critical level of respiratory depression and the need for immediate intervention to prevent life-threatening hypoxemia. Ventilatory assistance should be maintained until the opioid-induced respiratory depression has resolved. The opioid antagonist naloxone should be readily available whenever opioids are used at high doses or in patients with compromised pulmonary function. Opioid effects are dose-related, and there is great variability among patients in the doses that relieve pain and produce side effects. Synergistic respiratory depression is common when opioids are administered with other CNS depressants, most commonly the benzodiazepines. Because of this, initiation of therapy requires titration to optimal dose and interval. The most important principle is to provide adequate pain relief. This requires determining whether the drug has adequately relieved the pain and frequent reassessment to determine the optimal interval for dosing. *The most common error made by physicians in managing severe pain with opioids is to prescribe an inadequate dose. Because many patients are reluctant to complain, this practice leads to needless suffering.* In the absence of sedation at the expected time of peak effect, a physician should not hesitate to repeat the initial dose to achieve satisfactory pain relief.

An innovative approach to the problem of achieving adequate pain relief is the use of patient-controlled analgesia (PCA). PCA uses a microprocessor-controlled infusion device that can deliver a baseline continuous dose of an opioid drug as well as preprogrammed

additional doses whenever the patient pushes a button. The patient can then titrate the dose to the optimal level. This approach is used most extensively for the management of postoperative pain, but there is no reason why it should not be used for any hospitalized patient with persistent severe pain. PCA is also used for short-term home care of patients with intractable pain, such as that caused by metastatic cancer.

It is important to understand that the PCA device delivers small, repeated doses to maintain pain relief; in patients with severe pain, the pain must first be brought under control with a loading dose before transitioning to the PCA device. The bolus dose of the drug (typically 1 mg of morphine, 0.2 mg of hydromorphone, or 10 μ g of fentanyl) can then be delivered repeatedly as needed. To prevent overdosing, PCA devices are programmed with a lockout period after each demand dose is delivered (5–10 min) and a limit on the total dose delivered per hour. Although some have advocated the use of a simultaneous continuous or basal infusion of the PCA drug, this increases the risk of respiratory depression and has not been shown to increase the overall efficacy of the technique.

The availability of new routes of administration has extended the usefulness of opioid analgesics. Most important is the availability of spinal administration. Opioids can be infused through a spinal catheter placed either intrathecally or epidurally. By applying opioids directly to the spinal or epidural space adjacent to the spinal cord, regional analgesia can be obtained using relatively low total doses. Indeed, the dose required to produce effective localized analgesia when using morphine intrathecally (0.1–0.3 mg) is a fraction of that required to produce similar analgesia when administered intravenously (5–10 mg). In this way, side effects such as sedation, nausea, and respiratory depression can be minimized. This approach has been used extensively during labor and delivery and for postoperative pain relief following surgical procedures. Continuous intrathecal delivery via implanted spinal drug-delivery systems is now commonly used, particularly for the treatment of cancer-related pain that would require sedating doses for adequate pain control if given systemically. Opioids can also be given intranasally (butorphanol), rectally, and transdermally (fentanyl and buprenorphine), or through the oral mucosa (fentanyl), thus avoiding the discomfort of frequent injections in patients who cannot be given oral medication. The fentanyl and buprenorphine transdermal patches have the advantage of providing fairly steady plasma levels, which maximizes patient comfort.

Recent additions to the armamentarium for treating opioid-induced side effects are the peripherally acting opioid antagonists alvimopan (Entereg) and methylnaltrexone (Relistor). Alvimopan is available as an orally administered agent that is restricted to the intestinal lumen by limited absorption; methylnaltrexone is available in a subcutaneously administered form that has virtually no penetration into the CNS. Both agents act by binding to peripheral μ -receptors, thereby inhibiting or reversing the effects of opioids at these peripheral sites. The action of both agents is restricted to receptor sites outside of the CNS; thus, these drugs can reverse the adverse effects of opioid analgesics that are mediated through their peripheral receptors without reversing their analgesic effects. Alvimopan has proven effective in lowering the duration of persistent ileus following abdominal surgery in patients receiving opioid analgesics for postoperative pain control. Methylnaltrexone has proven effective for relief of opioid-induced constipation in patients taking opioid analgesics on a chronic basis.

Opioid and COX Inhibitor Combinations When used in combination, opioids and COX inhibitors have additive effects. Because a lower dose of each can be used to achieve the same degree of pain relief and their side effects are nonadditive, such combinations are used to lower the severity of dose-related side effects. However, fixed-ratio combinations of an opioid with acetaminophen carry an important risk. Dose escalation as a result of increased severity of pain or decreased opioid effect as a result of tolerance may lead to ingestion of levels of acetaminophen that are toxic to the liver. Although

acetaminophen-related hepatotoxicity is uncommon, it remains a significant cause for liver failure. Thus, many practitioners have moved away from the use of opioid-acetaminophen combination analgesics to avoid the risk of excessive acetaminophen exposure as the dose of the analgesic is escalated.

CHRONIC PAIN

Managing patients with chronic pain is intellectually and emotionally challenging. The patient's problem is often difficult or impossible to diagnose with certainty; such patients are demanding of the physician's time and often appear emotionally distraught. The traditional medical approach of seeking an obscure organic pathology is usually unhelpful. On the other hand, psychological evaluation and behaviorally based treatment paradigms are frequently helpful, particularly in the setting of a multidisciplinary pain-management center. Unfortunately, this approach, while effective, remains largely underused in current medical practice.

There are several factors that can cause, perpetuate, or exacerbate chronic pain. First, of course, the patient may simply have a disease that is characteristically painful for which there is presently no cure. Arthritis, cancer, chronic daily headaches, fibromyalgia, and diabetic neuropathy are examples of this. Second, there may be secondary perpetuating factors that are initiated by disease and persist after that disease has resolved. Examples include damaged sensory nerves, sympathetic efferent activity, and painful reflex muscle contraction (spasm). Finally, a variety of psychological conditions can exacerbate or even cause pain.

There are certain areas to which special attention should be paid in a patient's medical history. Because depression is the most common emotional disturbance in patients with chronic pain, patients should be questioned about their mood, appetite, sleep patterns, and daily activity. A simple standardized questionnaire, such as the Beck Depression Inventory, can be a useful screening device. It is important to remember that major depression is a common, treatable, and potentially fatal illness.

Other clues that a significant emotional disturbance is contributing to a patient's chronic pain complaint include pain that occurs in multiple, unrelated sites; a pattern of recurrent, but separate, pain problems beginning in childhood or adolescence; pain beginning at a time of emotional trauma, such as the loss of a parent or spouse; a history of physical or sexual abuse; and past or present substance abuse.

On examination, special attention should be paid to whether the patient guards the painful area and whether certain movements or postures are avoided because of pain. Discovering a mechanical component to the pain can be useful both diagnostically and therapeutically. Painful areas should be examined for deep tenderness, noting whether this is localized to muscle, ligamentous structures, or joints. Chronic myofascial pain is very common, and, in these patients, deep palpation may reveal highly localized trigger points that are firm bands or knots in muscle. Relief of the pain following injection of local anesthetic into these trigger points supports the diagnosis. A neuropathic component to the pain is indicated by evidence of nerve damage, such as sensory impairment, exquisitely sensitive skin (allodynia), weakness, and muscle atrophy, or loss of deep tendon reflexes. Evidence suggesting sympathetic nervous system involvement includes the presence of diffuse swelling, changes in skin color and temperature, and hypersensitive skin and joint tenderness compared with the normal side. Relief of the pain with a sympathetic block supports the diagnosis, but once the condition becomes chronic, the response to sympathetic blockade is of variable magnitude and duration; the role for repeated sympathetic blocks in the overall management of CRPS is not established.

A guiding principle in evaluating patients with chronic pain is to assess both emotional and organic factors before initiating therapy. Addressing these issues together, rather than waiting to address emotional issues after organic causes of pain have been ruled out, improves compliance in part because it assures patients that a psychological evaluation does not mean that the physician is questioning the validity

of their complaint. Even when an organic cause for a patient's pain can be found, it is still wise to look for other factors. For example, a cancer patient with painful bony metastases may have additional pain due to nerve damage and may also be depressed. Optimal therapy requires that each of these factors be looked for and treated.

TREATMENT CHRONIC PAIN

Once the evaluation process has been completed and the likely causative and exacerbating factors identified, an explicit treatment plan should be developed. An important part of this process is to identify specific and realistic functional goals for therapy, such as getting a good night's sleep, being able to go shopping, or returning to work. A multidisciplinary approach that uses medications, counseling, physical therapy, nerve blocks, and even surgery may be required to improve the patient's quality of life. There are also some newer, relatively invasive procedures that can be helpful for some patients with intractable pain. These include image-guided interventions such as epidural injection of glucocorticoids for acute radicular pain and radiofrequency treatment of the facet joints for chronic facet-related back and neck pain. For patients with severe and persistent pain that is unresponsive to more conservative treatment, placement of electrodes within the spinal canal overlying the dorsal columns of the spinal cord (spinal cord stimulation) or implantation of intrathecal drug-delivery systems has shown significant benefit. The criteria for predicting which patients will respond to these procedures continue to evolve. They are generally reserved for patients who have not responded to conventional pharmacologic approaches. Referral to a multidisciplinary pain clinic for a full evaluation should precede any invasive procedure. Such referrals are clearly not necessary for all chronic pain patients. For some, pharmacologic management alone can provide adequate relief.

ANTIDEPRESSANT MEDICATIONS

The tricyclic antidepressants (TCAs), particularly nortriptyline and desipramine (Table 18-1), are useful for the management of chronic pain. Although developed for the treatment of depression, the TCAs have a spectrum of dose-related biologic activities that include analgesia in a variety of chronic clinical conditions. Although the mechanism is unknown, the analgesic effect of TCAs has a more rapid onset and occurs at a lower dose than is typically required for the treatment of depression. Furthermore, patients with chronic pain who are not depressed obtain pain relief with antidepressants. There is evidence that TCAs potentiate opioid analgesia, so they may be useful adjuncts for the treatment of severe persistent pain such as occurs with malignant tumors. **Table 18-2** lists some of the painful conditions that respond to TCAs. TCAs are of particular value in the management of neuropathic pain such as occurs in diabetic neuropathy and postherpetic neuralgia, for which there are few other therapeutic options.

The TCAs that have been shown to relieve pain have significant side effects (Table 18-1; **Chap. 466**). Some of these side effects, such as orthostatic hypotension, drowsiness, cardiac conduction delay, memory impairment, constipation, and urinary retention,

TABLE 18-2 PAINFUL CONDITIONS THAT RESPOND TO TRICYCLIC ANTIDEPRESSANTS

Postherpetic neuralgia ^a
Diabetic neuropathy ^a
Tension headache ^a
Migraine headache ^a
Rheumatoid arthritis ^{a,b}
Chronic low back pain ^b
Cancer
Central poststroke pain

^aControlled trials demonstrate analgesia. ^bControlled studies indicate benefit but not analgesia.

are particularly problematic in elderly patients, and several are additive to the side effects of opioid analgesics. The selective serotonin reuptake inhibitors such as fluoxetine (Prozac) have fewer and less serious side effects than TCAs, but they are much less effective for relieving pain. It is of interest that venlafaxine (Effexor) and duloxetine (Cymbalta), which are nontricyclic antidepressants that block both serotonin and norepinephrine reuptake, appear to retain most of the pain-relieving effect of TCAs with a side effect profile more like that of the selective serotonin reuptake inhibitors. These drugs may be particularly useful in patients who cannot tolerate the side effects of TCAs.

ANTICONVULSANTS AND ANTIARRHYTHMICS

These drugs are useful primarily for patients with neuropathic pain. Phenytoin (Dilantin) and carbamazepine (Tegretol) were first shown to relieve the pain of trigeminal neuralgia. This pain has a characteristic brief, shooting, electric shock–like quality. In fact, anticonvulsants seem to be particularly helpful for pains that have such a lancinating quality. Newer anticonvulsants, gabapentin (Neurontin) and pregabalin (Lyrica), are effective for a broad range of neuropathic pains. Furthermore, because of their favorable side effect profile, these newer anticonvulsants are often used as first-line agents.

CHRONIC OPIOID MEDICATION

The long-term use of opioids is accepted for patients with pain due to malignant disease. Although opioid use for chronic pain of nonmalignant origin is controversial, it is clear that, for many patients, opioids are the only option that produces meaningful pain relief. This is understandable because opioids are the most potent and have the broadest range of efficacy of any analgesic medications. Although addiction is rare in patients who first use opioids for pain relief, some degree of tolerance and physical dependence is likely with long-term use. Furthermore, animal studies suggest that long-term opioid therapy may worsen pain in some individuals. Therefore, before embarking on opioid therapy, other options should be explored, and the limitations and risks of opioids should be explained to the patient. It is also important to point out that some opioid analgesic medications have mixed agonist-antagonist properties (e.g., butorphanol and buprenorphine). From a practical standpoint, this means that they may worsen pain by inducing an abstinence syndrome in patients who are physically dependent on other opioid analgesics.

With long-term outpatient use of orally administered opioids, it is desirable to use long-acting compounds such as levorphanol, methadone, sustained-release morphine, or transdermal fentanyl (Table 18-1). The pharmacokinetic profiles of these drug preparations enable the maintenance of sustained analgesic blood levels, potentially minimizing side effects such as sedation that are associated with high peak plasma levels, and reducing the likelihood of rebound pain associated with a rapid fall in plasma opioid concentration. Although long-acting opioid preparations may provide superior pain relief in patients with a continuous pattern of ongoing pain, others suffer from intermittent severe episodic pain and experience superior pain control and fewer side effects with the periodic use of short-acting opioid analgesics. Constipation is a virtually universal side effect of opioid use and should be treated expectantly. As noted above in the discussion of acute pain treatment, a recent advance for patients is the development of peripherally acting opioid antagonists that can reverse the constipation associated with opioid use without interfering with analgesia.

Soon after the introduction of a controlled-release oxycodone formulation (OxyContin) in the late 1990s, a dramatic rise in emergency department visits and deaths associated with oxycodone ingestion appeared, focusing public attention on misuse of prescription pain medications. The magnitude of prescription opioid abuse has grown over the last decade, leading the Centers for Disease Control and Prevention to classify prescription opioid analgesic abuse as an epidemic. This appears to be due in large part to individuals using a prescription drug nonmedically, most often an opioid analgesic.

TABLE 18-3 GUIDELINES FOR SELECTING AND MONITORING PATIENTS RECEIVING CHRONIC OPIOID THERAPY (COT) FOR THE TREATMENT OF CHRONIC, NONCANCER PAIN

Patient Selection

- Conduct a history, physical examination, and appropriate testing, including an assessment of risk of substance abuse, misuse, or addiction.
- Consider a trial of COT if pain is moderate or severe, pain is having an adverse impact on function or quality of life, and potential therapeutic benefits outweigh potential harms.
- A benefit-to-harm evaluation, including a history, physical examination, and appropriate diagnostic testing, should be performed and documented before and on an ongoing basis during COT.

Informed Consent and Use of Management Plans

- Informed consent should be obtained. A continuing discussion with the patient regarding COT should include goals, expectations, potential risks, and alternatives to COT.
- Consider using a written COT management plan to document patient and clinician responsibilities and expectations and assist in patient education.

Initiation and Titration

- Initial treatment with opioids should be considered as a therapeutic trial to determine whether COT is appropriate.
- Opioid selection, initial dosing, and titration should be individualized according to the patient's health status, previous exposure to opioids, attainment of therapeutic goals, and predicted or observed harms.

Monitoring

- Reassess patients on COT periodically and as warranted by changing circumstances. Monitoring should include documentation of pain intensity and level of functioning, assessments of progress toward achieving therapeutic goals, presence of adverse events, and adherence to prescribed therapies.
- In patients on COT who are at high risk or who have engaged in aberrant drug-related behaviors, clinicians should periodically obtain urine drug screens or other information to confirm adherence to the COT plan of care.
- In patients on COT not at high risk and not known to have engaged in aberrant drug-related behaviors, clinicians should consider periodically obtaining urine drug screens or other information to confirm adherence to the COT plan of care.

Source: Adapted with permission from R Chou et al: *J Pain* 10:113, 2009.

Drug-induced deaths have rapidly risen and are now the second leading cause of death in Americans, just behind motor vehicle fatalities. In 2011, the Office of National Drug Control Policy established a multifaceted approach to address prescription drug abuse, including Prescription Drug Monitoring Programs that allow practitioners to determine if patients are receiving prescriptions from multiple providers and use of law enforcement to eliminate improper prescribing practices. This increased scrutiny leaves many practitioners hesitant to prescribe opioid analgesics, other than for brief periods to control pain associated with illness or injury. For now, the choice to begin chronic opioid therapy for a given patient is left to the individual practitioner. Pragmatic guidelines for properly selecting and monitoring patients receiving chronic opioid therapy are shown in **Table 18-3**.

TREATMENT OF NEUROPATHIC PAIN

It is important to individualize treatment for patients with neuropathic pain. Several general principles should guide therapy: the first is to move quickly to provide relief, and the second is to minimize drug side effects. For example, in patients with postherpetic neuralgia and significant cutaneous hypersensitivity, topical lidocaine (Lidoderm patches) can provide immediate relief without side effects. Anticonvulsants (gabapentin or pregabalin; see above) or antidepressants (nortriptyline, desipramine, duloxetine, or venlafaxine) can be used as first-line drugs for patients with neuropathic pain. Systemically administered antiarrhythmic drugs such as lidocaine and mexiletine are less likely to be effective; although intravenous infusion of lidocaine can provide analgesia for patients with different types of neuropathic pain, the relief is usually transient, typically lasting just hours after the cessation of the infusion.

The oral lidocaine congener mexiletine is poorly tolerated, producing frequent gastrointestinal adverse effects. There is no consensus on which class of drug should be used as a first-line treatment for any chronically painful condition. However, because relatively high doses of anticonvulsants are required for pain relief, sedation is very common. Sedation is also a problem with TCAs but is much less of a problem with serotonin/norepinephrine reuptake inhibitors (SNRIs; e.g., venlafaxine and duloxetine). Thus, in the elderly or in patients whose daily activities require high-level mental activity, these drugs should be considered the first line. In contrast, opioid medications should be used as a second- or third-line drug class. Although highly effective for many painful conditions, opioids are sedating, and their effect tends to lessen over time, leading to dose escalation and, occasionally, a worsening of pain due to physical dependence. Drugs of different classes can be used in combination to optimize pain control.

It is worth emphasizing that many patients, especially those with chronic pain, seek medical attention primarily because they are suffering and because only physicians can provide the medications required for pain relief. A primary responsibility of all physicians is to minimize the physical and emotional discomfort of their patients. Familiarity with pain mechanisms and analgesic medications is an important step toward accomplishing this aim.

19 Chest Discomfort

David A. Morrow

Chest discomfort is among the most common reasons for which patients present for medical attention at either an emergency department (ED) or an outpatient clinic. The evaluation of nontraumatic chest discomfort is inherently challenging owing to the broad variety of possible causes, a minority of which are life-threatening conditions that should not be missed. It is helpful to frame the initial diagnostic assessment and triage of patients with acute chest discomfort around three categories: (1) myocardial ischemia; (2) other cardiopulmonary causes (pericardial disease, aortic emergencies, and pulmonary conditions); and (3) non-cardiopulmonary causes. Although rapid identification of high-risk conditions is a priority of the initial assessment, strategies that incorporate routine liberal use of testing carry the potential for adverse effects of unnecessary investigations.

EPIDEMIOLOGY AND NATURAL HISTORY

Chest discomfort is the third most common reason for visits to the ED in the United States, resulting in 6 to 7 million emergency visits each year. More than 60% of patients with this presentation are hospitalized for further testing, and the rest undergo additional investigation in the ED. Fewer than 25% of evaluated patients are eventually diagnosed with acute coronary syndrome (ACS), with rates of 5–15% in most series of unselected populations. In the remainder, the most common diagnoses are gastrointestinal causes (Fig. 19-1), and fewer than 10% are other life-threatening cardiopulmonary conditions. In a large proportion of patients with transient acute chest discomfort, ACS or another acute cardiopulmonary cause is excluded but the cause is not determined. Therefore, the resources and time devoted to the evaluation of chest discomfort *in the absence of a severe cause* are substantial. Nevertheless, a disconcerting 2–6% of patients with chest discomfort of presumed non-ischemic etiology who are discharged from the ED are later deemed to have had a missed myocardial infarction (MI). Patients with a missed diagnosis of MI have a 30-day risk of death that is double that of their counterparts who are hospitalized.

The natural histories of ACS, acute pericardial diseases, pulmonary embolism, and aortic emergencies are discussed in Chaps. 288, 294 and 295, 300, and 301, respectively. In a study of more than 350,000 patients with unspecified presumed non-cardiopulmonary chest discomfort, the mortality rate 1 year after discharge was <2% and did not differ significantly from age-adjusted mortality in the general population. The estimated rate of major cardiovascular events through 30 days in patients with acute chest pain who had been stratified as low risk was 2.5% in a large population-based study that excluded patients with ST-segment elevation or definite noncardiac chest pain.

CAUSES OF CHEST DISCOMFORT

The major etiologies of chest discomfort are discussed in this section and summarized in Table 19-1. Additional elements of the history, physical examination, and diagnostic testing that aid in distinguishing these causes are discussed in a later section (see “Approach to the Patient”).

MYOCARDIAL ISCHEMIA/INJURY

Myocardial ischemia causing chest discomfort, termed *angina pectoris*, is a primary clinical concern in patients presenting with chest symptoms. Myocardial ischemia is precipitated by an imbalance between myocardial oxygen requirements and myocardial oxygen supply, resulting in insufficient delivery of oxygen to meet the heart’s metabolic demands. Myocardial oxygen consumption may be elevated by increases in heart rate, ventricular wall stress, and myocardial contractility, whereas myocardial oxygen supply is determined by coronary

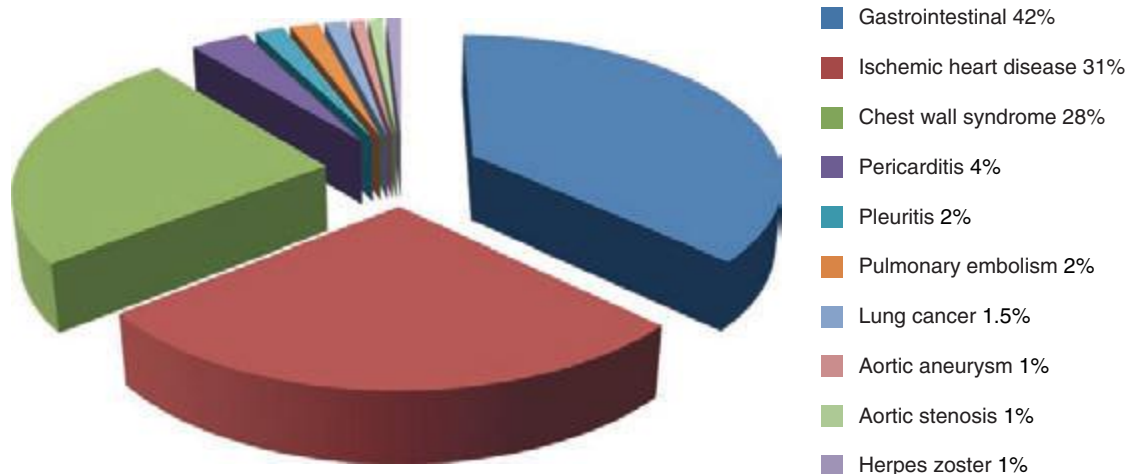


FIGURE 19-1 Distribution of final discharge diagnoses in patients with nontraumatic acute chest pain. (Figure prepared from data in P Fruergaard et al: *Eur Heart J* 17:1028, 1996.)

TABLE 19-1 TYPICAL CLINICAL FEATURES OF MAJOR CAUSES OF ACUTE CHEST DISCOMFORT

System	Condition	Onset/Duration	Quality	Location	Associated Features
Cardiopulmonary					
Cardiac	Myocardial ischemia	<i>Stable angina:</i> Precipitated by exertion, cold, or stress; 2–10 min	Pressure, tightness, squeezing, heaviness, burning	Retrosternal; often radiation to neck, jaw, shoulders, or arms; sometimes epigastric	S ₄ gallop or mitral regurgitation murmur (rare) during pain; S ₃ or rales if severe ischemia or complication of myocardial infarction
		<i>Unstable angina:</i> Increasing pattern or at rest <i>Myocardial infarction:</i> Usually >30 min			
	Pericarditis	Variable; hours to days; may be episodic	Pleuritic, sharp	Retrosternal or toward cardiac apex; may radiate to left shoulder	May be relieved by sitting up and leaning forward; pericardial friction rub
Vascular	Acute aortic syndrome	Sudden onset of unremitting pain	Tearing or ripping; knifelike	Anterior chest, often radiating to back, between shoulder blades	Associated with hypertension and/or underlying connective tissue disorder; murmur of aortic insufficiency; loss of peripheral pulses
	Pulmonary embolism	Sudden onset	Pleuritic; may manifest as heaviness with massive pulmonary embolism	Often lateral, on the side of the embolism	Dyspnea, tachypnea, tachycardia, and hypotension
	Pulmonary hypertension	Variable; often exertional	Pressure	Substernal	Dyspnea, signs of increased venous pressure
Pulmonary	Pneumonia or pleuritis	Variable	Pleuritic	Unilateral, often localized	Dyspnea, cough, fever, rales, occasional rub
	Spontaneous pneumothorax	Sudden onset	Pleuritic	Lateral to side of pneumothorax	Dyspnea, decreased breath sounds on side of pneumothorax
Non-cardiopulmonary					
Gastrointestinal	Esophageal reflux	10–60 min	Burning	Substernal, epigastric	Worsened by postprandial recumbency; relieved by antacids
	Esophageal spasm	2–30 min	Pressure, tightness, burning	Retrosternal	Can closely mimic angina
	Peptic ulcer	Prolonged; 60–90 min after meals	Burning	Epigastric, substernal	Relieved with food or antacids
	Gallbladder disease	Prolonged	Aching or colicky	Epigastric, right upper quadrant; sometimes to the back	May follow meal
Neuromuscular	Costochondritis	Variable	Aching	Sternal	Sometimes swollen, tender, warm over joint; may be reproduced by localized pressure on examination
	Cervical disk disease	Variable; may be sudden	Aching; may include numbness	Arms and shoulders	May be exacerbated by movement of neck
	Trauma or strain	Usually constant	Aching	Localized to area of strain	Reproduced by movement or palpation
	Herpes zoster	Usually prolonged	Sharp or burning	Dermatomal distribution	Vesicular rash in area of discomfort
Psychological	Emotional and psychiatric conditions	Variable; may be fleeting or prolonged	Variable; often manifests as tightness and dyspnea with feeling of panic or doom	Variable; may be retrosternal	Situational factors may precipitate symptoms; history of panic attacks, depression

blood flow and coronary arterial oxygen content. When myocardial ischemia is sufficiently severe and prolonged in duration (as little as 20 min), irreversible cellular injury occurs, resulting in MI.

Ischemic heart disease is most commonly caused by atheromatous plaque that obstructs one or more of the epicardial coronary arteries. Stable ischemic heart disease (Chap. 293) usually results from the gradual atherosclerotic narrowing of the coronary arteries. *Stable angina* is characterized by ischemic episodes that are typically precipitated by a superimposed increase in oxygen demand during physical exertion and relieved upon resting. Ischemic heart disease becomes unstable most commonly when rupture or erosion of one or more atherosclerotic lesions triggers coronary thrombosis (Chap. 291e). Unstable ischemic heart disease is classified clinically by the presence or absence of detectable myocardial injury and the presence or absence of ST-segment elevation on the patient's electrocardiogram (ECG). When acute coronary atherothrombosis occurs, the intracoronary thrombus may be partially obstructive, generally leading to myocardial ischemia in the absence of ST-segment elevation. Marked by ischemic

symptoms at rest, with minimal activity, or in an accelerating pattern, unstable ischemic heart disease is classified as *unstable angina* when there is no detectable myocardial injury and as *non-ST elevation MI* (NSTEMI) when there is evidence of myocardial necrosis (Chap. 294). When the coronary thrombus is acutely and completely occlusive, transmural myocardial ischemia usually ensues, with ST-segment elevation on the ECG and myocardial necrosis leading to a diagnosis of *ST elevation MI* (STEMI, see Chap. 295).

Clinicians should be aware that unstable ischemic symptoms may also occur predominantly because of increased myocardial oxygen demand (e.g., during intense psychological stress or fever) or because of decreased oxygen delivery due to anemia, hypoxia, or hypotension. However, the term *acute coronary syndrome*, which encompasses unstable angina, NSTEMI, and STEMI, is in general reserved for ischemia precipitated by acute coronary atherothrombosis. In order to guide therapeutic strategies, a standardized system for classification of MI has been expanded to discriminate MI resulting from acute coronary thrombosis (type 1) from MI occurring secondary to other

imbalances of myocardial oxygen supply and demand (type 2; see [Chap. 294](#)).

Other contributors to stable and unstable ischemic heart disease, such as endothelial dysfunction, microvascular disease, and vasospasm, may exist alone or in combination with coronary atherosclerosis and may be the dominant cause of myocardial ischemia in some patients. Moreover, non-atherosclerotic processes, including congenital abnormalities of the coronary vessels, myocardial bridging, coronary arteritis, and radiation-induced coronary disease, can lead to coronary obstruction. In addition, conditions associated with extreme myocardial oxygen demand and impaired endocardial blood flow, such as aortic valve disease ([Chap. 301](#)), hypertrophic cardiomyopathy, or idiopathic dilated cardiomyopathy ([Chap. 287](#)), can precipitate myocardial ischemia in patients with or without underlying obstructive atherosclerosis.

Characteristics of Ischemic Chest Discomfort The clinical characteristics of angina pectoris, often referred to simply as “angina,” are highly similar whether the ischemic discomfort is a manifestation of stable ischemic heart disease, unstable angina, or MI; the exceptions are differences in the pattern and duration of symptoms associated with these syndromes (Table 19-1). Heberden initially described angina as a sense of “strangling and anxiety.” Chest discomfort characteristic of myocardial ischemia is typically described as aching, heavy, squeezing, crushing, or constricting. However, in a substantial minority of patients, the quality of discomfort is extremely vague and may be described as a mild tightness, or merely an uncomfortable feeling, that sometimes is experienced as numbness or a burning sensation. The site of the discomfort is usually retrosternal, but radiation is common and generally occurs down the ulnar surface of the left arm; the right arm, both arms, neck, jaw, or shoulders may also be involved. These and other characteristics of ischemic chest discomfort pertinent to discrimination from other causes of chest pain are discussed later in this chapter (see “Approach to the Patient”).

Stable angina usually begins gradually and reaches its maximal intensity over a period of minutes before dissipating within several minutes with rest or with nitroglycerin. The discomfort typically occurs predictably at a characteristic level of exertion or psychological stress. By definition, unstable angina is manifest by self-limited anginal chest discomfort that is exertional but occurs at increased frequency with progressively lower intensity of physical activity or even at rest. Chest discomfort associated with MI is typically more severe, is prolonged (usually lasting ≥ 30 min), and is not relieved by rest.

Mechanisms of Cardiac Pain The neural pathways involved in ischemic cardiac pain are poorly understood. Ischemic episodes are thought to excite local chemosensitive and mechanoreceptive receptors that, in turn, stimulate release of adenosine, bradykinin, and other substances that activate the sensory ends of sympathetic and vagal afferent fibers. The afferent fibers traverse the nerves that connect to the upper five thoracic sympathetic ganglia and upper five distal thoracic roots of the spinal cord. From there, impulses are transmitted to the thalamus. Within the spinal cord, cardiac sympathetic afferent impulses may converge with impulses from somatic thoracic structures, and this convergence may be the basis for referred cardiac pain. In addition, cardiac vagal afferent fibers synapse in the nucleus tractus solitarius of the medulla and then descend to the upper cervical spinothalamic tract, and this route may contribute to anginal pain experienced in the neck and jaw.

OTHER CARDIOPULMONARY CAUSES

Pericardial and Other Myocardial Diseases (See also [Chap. 288](#)) Inflammation of the pericardium due to infectious or noninfectious causes can be responsible for acute or chronic chest discomfort. The visceral surface and most of the parietal surface of the pericardium are insensitive to pain. Therefore, the pain of pericarditis is thought to arise principally from associated pleural inflammation and is more common with infectious causes of pericarditis, which typically involve the pleura. Because of this pleural association, the discomfort of pericarditis is usually pleuritic pain that is exacerbated by breathing,

coughing, or changes in position. Moreover, owing to the overlapping sensory supply of the central diaphragm via the phrenic nerve with somatic sensory fibers originating in the third to fifth cervical segments, the pain of pleural pericarditis is often referred to the shoulder and neck. Involvement of the pleural surface of the lateral diaphragm can lead to pain in the upper abdomen.

Acute inflammatory and other non-ischemic myocardial diseases can also produce chest discomfort. The symptoms of *Takotsubo* (stress-related) *cardiomyopathy* often start abruptly with chest pain and shortness of breath. This form of cardiomyopathy, in its most recognizable form, is triggered by an emotionally or physically stressful event and may mimic acute MI because of its commonly associated ECG abnormalities, including ST-segment elevation, and elevated biomarkers of myocardial injury. Observational studies support a predilection for women >50 years of age. The symptoms of acute myocarditis are highly varied. Chest discomfort may either originate with inflammatory injury of the myocardium or be due to severe increases in wall stress related to poor ventricular performance.

Diseases of the Aorta (See also [Chap. 301](#)) Acute aortic dissection (Fig. 19-1) is a less common cause of chest discomfort but is important because of the catastrophic natural history of certain subsets of cases when recognized late or left untreated. Acute aortic syndromes encompass a spectrum of acute aortic diseases related to disruption of the media of the aortic wall. *Aortic dissection* involves a tear in the aortic intima, resulting in separation of the media and creation of a separate “false” lumen. A *penetrating ulcer* has been described as ulceration of an aortic atheromatous plaque that extends through the intima and into the aortic media, with the potential to initiate an intramural dissection or rupture into the adventitia. *Intramural hematoma* is an aortic wall hematoma with no demonstrable intimal flap, no radiologically apparent intimal tear, and no false lumen. Intramural hematoma can occur due to either rupture of the vasa vasorum or, less commonly, a penetrating ulcer.

Each of these subtypes of acute aortic syndrome typically presents with chest discomfort that is often severe, sudden in onset, and sometimes described as “tearing” in quality. Acute aortic syndromes involving the *ascending* aorta tend to cause pain in the midline of the anterior chest, whereas *descending* aortic syndromes most often present with pain in the back. Therefore, dissections that begin in the ascending aorta and extend to the descending aorta tend to cause pain in the front of the chest that extends toward the back, between the shoulder blades. Proximal aortic dissections that involve the ascending aorta (type A in the Stanford nomenclature) are at high risk for major complications that may influence the clinical presentation, including (1) compromise of the aortic ostia of the coronary arteries, resulting in MI; (2) disruption of the aortic valve, causing acute aortic insufficiency; and (3) rupture of the hematoma into the pericardial space, leading to pericardial tamponade.

Knowledge of the epidemiology of acute aortic syndromes can be helpful in maintaining awareness of this relatively uncommon group of disorders (estimated annual incidence, 3 cases per 100,000 population). Nontraumatic aortic dissections are very rare in the absence of hypertension or conditions associated with deterioration of the elastic or muscular components of the aortic media, including pregnancy, bicuspid aortic disease, or inherited connective tissue diseases, such as Marfan and Ehlers-Danlos syndromes.

Although aortic aneurysms are most often asymptomatic, thoracic aortic aneurysms can cause chest pain and other symptoms by compressing adjacent structures. This pain tends to be steady, deep, and occasionally severe. Aortitis, whether of noninfectious or infectious etiology, in the absence of aortic dissection is a rare cause of chest or back discomfort.

Pulmonary Conditions Pulmonary and pulmonary-vascular conditions that cause chest discomfort usually do so in conjunction with dyspnea and often produce symptoms that have a pleuritic nature.

PULMONARY EMBOLISM (See also [Chap. 300](#)) Pulmonary emboli (annual incidence, ~ 1 per 1000) can produce dyspnea and chest discomfort

that is sudden in onset. Typically pleuritic in pattern, the chest discomfort associated with pulmonary embolism may result from (1) involvement of the pleural surface of the lung adjacent to a resultant pulmonary infarction; (2) distention of the pulmonary artery; or (3) possibly, right ventricular wall stress and/or subendocardial ischemia related to acute pulmonary hypertension. The pain associated with small pulmonary emboli is often lateral and pleuritic and is believed to be related to the first of these three possible mechanisms. In contrast, massive pulmonary emboli may cause severe substernal pain that may mimic an MI and that is plausibly attributed to the second and third of these potential mechanisms. Massive or submassive pulmonary embolism may also be associated with syncope, hypotension, and signs of right heart failure. Other typical characteristics that aid in the recognition of pulmonary embolism are discussed later in this chapter (see “Approach to the Patient”).

PNEUMOTHORAX (See also Chap. 317) Primary spontaneous pneumothorax is a rare cause of chest discomfort, with an estimated annual incidence in the United States of 7 per 100,000 among men and <2 per 100,000 among women. Risk factors include male sex, smoking, family history, and Marfan syndrome. The symptoms are usually sudden in onset, and dyspnea may be mild; thus, presentation to medical attention is sometimes delayed. Secondary spontaneous pneumothorax may occur in patients with underlying lung disorders, such as chronic obstructive pulmonary disease, asthma, or cystic fibrosis, and usually produces symptoms that are more severe. Tension pneumothorax is a medical emergency caused by trapped intrathoracic air that precipitates hemodynamic collapse.

Other Pulmonary Parenchymal, Pleural, or Vascular Disease (See also Chaps. 304, 305, and 316) Most pulmonary diseases that produce chest pain, including pneumonia and malignancy, do so because of involvement of the pleura or surrounding structures. Pleurisy is typically described as a knifelike pain that is worsened by inspiration or coughing. In contrast, chronic pulmonary hypertension can manifest as chest pain that may be very similar to angina in its characteristics, suggesting right ventricular myocardial ischemia in some cases. Reactive airways diseases similarly can cause chest tightness associated with breathlessness rather than pleurisy.

NON-CARDIOPULMONARY CAUSES

Gastrointestinal Conditions (See also Chap. 344) Gastrointestinal disorders are the most common cause of nontraumatic chest discomfort and often produce symptoms that are difficult to discern from more serious causes of chest pain, including myocardial ischemia. Esophageal disorders, in particular, may simulate angina in the character and location of the pain. Gastroesophageal reflux and disorders of esophageal motility are common and should be considered in the differential diagnosis of chest pain (Fig. 19-1 and Table 19-1). Acid reflux often causes a burning discomfort. The pain of esophageal spasm, in contrast, is commonly an intense, squeezing discomfort that is retrosternal in location and, like angina, may be relieved by nitroglycerin or dihydropyridine calcium channel antagonists. Chest pain can also result from injury to the esophagus, such as a Mallory-Weiss tear or even an esophageal rupture (Boerhaave syndrome) caused by severe vomiting. Peptic ulcer disease is most commonly epigastric in location but can radiate into the chest (Table 19-1).

Hepatobiliary disorders, including cholecystitis and biliary colic, may mimic acute cardiopulmonary diseases. Although the pain arising from these disorders usually localizes to the right upper quadrant of the abdomen, it is variable and may be felt in the epigastrium and radiate to the back and lower chest. This discomfort is sometimes referred to the scapula or may in rare cases be felt in the shoulder, suggesting diaphragmatic irritation. The pain is steady, usually lasts several hours, and subsides spontaneously, without symptoms between attacks. Pain resulting from pancreatitis is typically aching epigastric pain that radiates to the back.

Musculoskeletal and Other Causes (See also Chap. 393) Chest discomfort can be produced by any musculoskeletal disorder involving the chest wall or the nerves of the chest wall, neck, or upper limbs.

Costochondritis causing tenderness of the costochondral junctions (*Tietze's syndrome*) is relatively common. Cervical radiculitis may manifest as a prolonged or constant aching discomfort in the upper chest and limbs. The pain may be exacerbated by motion of the neck. Occasionally, chest pain can be caused by compression of the brachial plexus by the cervical ribs, and tendinitis or bursitis involving the left shoulder may mimic the radiation of angina. Pain in a dermatomal distribution can also be caused by cramping of intercostal muscles or by herpes zoster (Chap. 217).

Emotional and Psychiatric Conditions As many as 10% of patients who present to emergency departments with acute chest discomfort have a panic disorder or related condition (Table 19-1). The symptoms may include chest tightness or aching that is associated with a sense of anxiety and difficulty breathing. The symptoms may be prolonged or fleeting.

APPROACH TO THE PATIENT: Chest Discomfort

Given the broad set of potential causes and the heterogeneous risk of serious complications in patients who present with acute nontraumatic chest discomfort, the priorities of the initial clinical encounter include assessment of (1) the patient's clinical stability and (2) the probability that the patient has an underlying cause of the discomfort that may be life-threatening. The high-risk conditions of principal concern are acute cardiopulmonary processes, including ACS, acute aortic syndrome, pulmonary embolism, tension pneumothorax, and pericarditis with tamponade. Among non-cardiopulmonary causes of chest pain, esophageal rupture likely holds the greatest urgency for diagnosis. Patients with these conditions may deteriorate rapidly despite initially appearing well. The remaining population with non-cardiopulmonary conditions has a more favorable prognosis during completion of the diagnostic work-up. A rapid targeted assessment for a serious cardiopulmonary cause is of particular relevance for patients with acute ongoing pain who have presented for emergency evaluation. Among patients presenting in the outpatient setting with chronic pain or pain that has resolved, a general diagnostic assessment is reasonably undertaken (see “Outpatient Evaluation of Chest Discomfort,” below). A series of questions that can be used to structure the clinical evaluation of patients with chest discomfort is shown in Table 19-2.

TABLE 19-2 CONSIDERATIONS IN THE ASSESSMENT OF THE PATIENT WITH CHEST DISCOMFORT

1. Could the chest discomfort be due to an acute, potentially life-threatening condition that warrants urgent evaluation and management?			
Unstable ischemic heart disease	Aortic dissection	Pneumothorax	Pulmonary embolism
2. If not, could the discomfort be due to a chronic condition likely to lead to serious complications?			
Stable angina	Aortic stenosis	Pulmonary hypertension	
3. If not, could the discomfort be due to an acute condition that warrants specific treatment?			
Pericarditis	Pneumonia/pleuritis	Herpes zoster	
4. If not, could the discomfort be due to another treatable chronic condition?			
Esophageal reflux	Cervical disk disease		
Esophageal spasm	Arthritis of the shoulder or spine		
Peptic ulcer disease	Costochondritis		
Gallbladder disease	Other musculoskeletal disorders		
Other gastrointestinal conditions	Anxiety state		

Source: Developed by Dr. Thomas H. Lee for the 18th edition of *Harrison's Principles of Internal Medicine*.

HISTORY

The evaluation of nontraumatic chest discomfort relies heavily on the clinical history and physical examination to direct subsequent diagnostic testing. The evaluating clinician should assess the quality, location (including radiation), and pattern (including onset and duration) of the pain as well as any provoking or alleviating factors. The presence of associated symptoms may also be useful in establishing a diagnosis.

Quality of Pain The quality of chest discomfort alone is never sufficient to establish a diagnosis. However, the characteristics of the pain are pivotal in formulating an initial clinical impression and assessing the likelihood of a serious cardiopulmonary process (Table 19-1), including ACS in particular (Fig. 19-2). Pressure or tightness is consistent with a typical presentation of myocardial ischemic pain. Nevertheless, the clinician must remember that some patients with ischemic chest symptoms deny any “pain” but rather complain of dyspnea or a vague sense of anxiety. The severity of the discomfort has poor diagnostic accuracy. It is often helpful to ask about the similarity of the discomfort to previous definite ischemic symptoms. It is unusual for angina to be sharp, as in knifelike, stabbing, or pleuritic; however, patients sometimes use the word “sharp” to convey the intensity of discomfort rather than the quality. Pleuritic discomfort is suggestive of a process involving the pleura, including pericarditis, pulmonary embolism, or pulmonary parenchymal processes. Less frequently, the pain of pericarditis or massive pulmonary embolism is a steady severe pressure or aching that can be difficult to discriminate from myocardial ischemia. “Tearing” or “ripping” pain is often described by patients with acute aortic dissection. However, acute aortic emergencies also present commonly with severe, knifelike pain. A burning quality can suggest acid reflux or peptic ulcer disease but may also occur with myocardial ischemia. Esophageal pain, particularly with spasm, can be a severe squeezing discomfort identical to angina.

Location of Discomfort A substernal location with radiation to the neck, jaw, shoulder, or arms is typical of myocardial ischemic discomfort. Some patients present with aching in sites of radiated pain

as their only symptoms of ischemia. However, pain that is highly localized—e.g., that which can be demarcated by the tip of one finger—is highly unusual for angina. A retrosternal location should prompt consideration of esophageal pain; however, other gastrointestinal conditions usually present with pain that is most intense in the abdomen or epigastrium, with possible radiation into the chest. Angina may also occur in an epigastric location. However, pain that occurs solely above the mandible or below the epigastrium is rarely angina. Severe pain radiating to the back, particularly between the shoulder blades, should prompt consideration of an acute aortic syndrome. Radiation to the trapezius ridge is characteristic of pericardial pain and does not usually occur with angina.

Pattern Myocardial ischemic discomfort usually builds over minutes and is exacerbated by activity and mitigated by rest. In contrast, pain that reaches its peak intensity immediately is more suggestive of aortic dissection, pulmonary embolism, or spontaneous pneumothorax. Pain that is fleeting (lasting only a few seconds) is rarely ischemic in origin. Similarly, pain that is constant in intensity for a prolonged period (many hours to days) is unlikely to represent myocardial ischemia if it occurs in the absence of other clinical consequences, such as abnormalities of the ECG, elevation of cardiac biomarkers, or clinical sequelae (e.g., heart failure or hypotension). Both myocardial ischemia and acid reflux may have their onset in the morning, the latter because of the absence of food to absorb gastric acid.

Provoking and Alleviating Factors Patients with myocardial ischemic pain usually prefer to rest, sit, or stop walking. However, clinicians should be aware of the phenomenon of “warm-up angina” in which some patients experience relief of angina as they continue at the same or even a greater level of exertion without symptoms (Chap. 293). Alterations in the intensity of pain with changes in position or movement of the upper extremities and neck are less likely with myocardial ischemia and suggest a musculoskeletal etiology. The pain of pericarditis, however, often is worse in the supine position and relieved by sitting upright and leaning forward. Gastroesophageal reflux may be exacerbated by alcohol, some foods, or by a reclined position. Relief can occur with sitting.

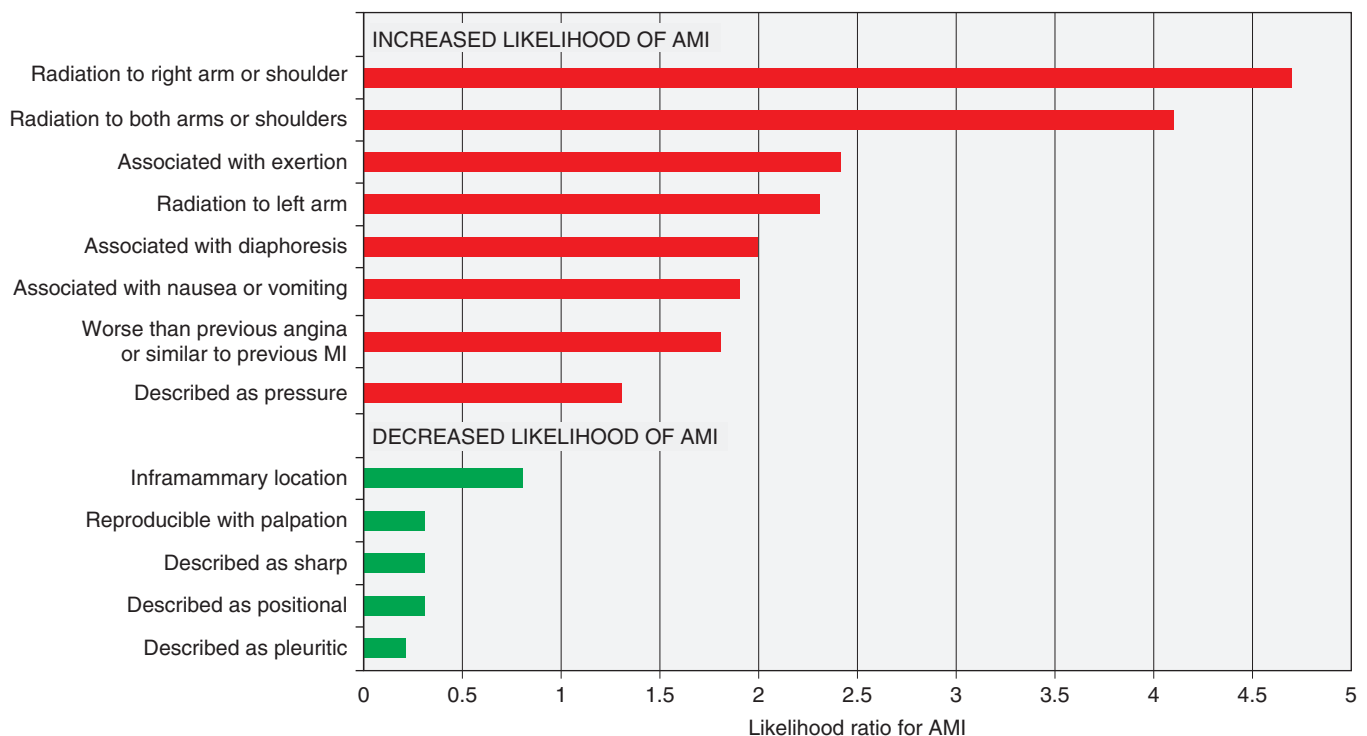


FIGURE 19-2 Association of chest pain characteristics with the probability of acute myocardial infarction (AMI). (Figure prepared from data in CJ Swap, JT Nagurney: JAMA 294:2623, 2005.)

Exacerbation by eating suggests a gastrointestinal etiology such as peptic ulcer disease, cholecystitis, or pancreatitis. Peptic ulcer disease tends to become symptomatic 60–90 min after meals. However, in the setting of severe coronary atherosclerosis, redistribution of blood flow to the splanchnic vasculature after eating can trigger postprandial angina. The discomfort of acid reflux and peptic ulcer disease is usually diminished promptly by acid-reducing therapies. In contrast with its impact in some patients with angina, physical exertion is very unlikely to alter symptoms from gastrointestinal causes of chest pain. Relief of chest discomfort within minutes after administration of nitroglycerin is suggestive of but not sufficiently sensitive or specific for a definitive diagnosis of myocardial ischemia. Esophageal spasm may also be relieved promptly with nitroglycerin. A delay of >10 min before relief is obtained after nitroglycerin suggests that the symptoms either are not caused by ischemia or are caused by severe ischemia, such as during acute MI.

Associated Symptoms Symptoms that accompany myocardial ischemia may include diaphoresis, dyspnea, nausea, fatigue, faintness, and eructations. In addition, these symptoms may exist in isolation as anginal equivalents (i.e., symptoms of myocardial ischemia other than typical angina), particularly in women and the elderly. Dyspnea may occur with multiple conditions considered in the differential diagnosis of chest pain and thus is not discriminative, but the presence of dyspnea is important because it suggests a cardiopulmonary etiology. Sudden onset of significant respiratory distress should lead to consideration of pulmonary embolism and spontaneous pneumothorax. Hemoptysis may occur with pulmonary embolism, or as blood-tinged frothy sputum in severe heart failure but usually points toward a pulmonary parenchymal etiology of chest symptoms. Presentation with syncope or pre-syncope should prompt consideration of hemodynamically significant pulmonary embolism or aortic dissection as well as ischemic arrhythmias. Although nausea and vomiting suggest a gastrointestinal disorder, these symptoms may occur in the setting of MI (more commonly inferior MI), presumably because of activation of the vagal reflex or stimulation of left ventricular receptors as part of the Bezold-Jarisch reflex.

Past Medical History The past medical history is useful in assessing the patient for risk factors for coronary atherosclerosis (Chap. 291e) and venous thromboembolism (Chap. 300) as well as for conditions that may predispose the patient to specific disorders. For example, a history of connective tissue diseases such as marfan syndrome should heighten the clinician's suspicion of an acute aortic syndrome or spontaneous pneumothorax. A careful history may elicit clues about depression or prior panic attacks.

PHYSICAL EXAMINATION

In addition to providing an initial assessment of the patient's clinical stability, the physical examination of patients with chest discomfort can provide direct evidence of specific etiologies of chest pain (e.g., unilateral absence of lung sounds) and can identify potential precipitants of acute cardiopulmonary causes of chest pain (e.g., uncontrolled hypertension), relevant comorbid conditions (e.g., obstructive pulmonary disease), and complications of the presenting syndrome (e.g., heart failure). However, because the findings on physical examination may be normal in patients with unstable ischemic heart disease, an unremarkable physical exam is not definitively reassuring.

General The patient's general appearance is helpful in establishing an initial impression of the severity of illness. Patients with acute MI or other acute cardiopulmonary disorders often appear anxious, uncomfortable, pale, cyanotic, or diaphoretic. Patients who are massaging or clutching their chests may describe their pain with a clenched fist held against the sternum (*Levine's sign*). Occasionally, body habitus is helpful—e.g., in patients with Marfan syndrome or the prototypical young, tall, thin man with spontaneous pneumothorax.

Vital Signs Significant tachycardia and hypotension are indicative of important hemodynamic consequences of the underlying cause of chest discomfort and should prompt a rapid survey for the most severe conditions, such as acute MI with cardiogenic shock, massive pulmonary embolism, pericarditis with tamponade, or tension pneumothorax. Acute aortic emergencies usually present with severe hypertension but may be associated with profound hypotension when there is coronary arterial compromise or dissection into the pericardium. Sinus tachycardia is an important manifestation of submassive pulmonary embolism. Tachypnea and hypoxemia point toward a pulmonary cause. The presence of low-grade fever is non-specific because it may occur with MI and with thromboembolism in addition to infection.

Pulmonary Examination of the lungs may localize a primary pulmonary cause of chest discomfort, as in cases of pneumonia, asthma, or pneumothorax. Left ventricular dysfunction from severe ischemia/infarction as well as acute valvular complications of MI or aortic dissection can lead to pulmonary edema, which is an indicator of high risk.

Cardiac The jugular venous pulse is often normal in patients with acute myocardial ischemia but may reveal characteristic patterns with pericardial tamponade or acute right ventricular dysfunction (Chaps. 267 and 288). Cardiac auscultation may reveal a third or, more commonly, a fourth heart sound, reflecting myocardial systolic or diastolic dysfunction. Murmurs of mitral regurgitation or a harsh murmur of a ventricular-septal defect may indicate mechanical complications of STEMI. A murmur of aortic insufficiency may be a complication of proximal aortic dissection. Other murmurs may reveal underlying cardiac disorders contributory to ischemia (e.g., aortic stenosis or hypertrophic cardiomyopathy). Pericardial friction rubs reflect pericardial inflammation.

Abdominal Localizing tenderness on the abdominal exam is useful in identifying a gastrointestinal cause of the presenting syndrome. Abdominal findings are infrequent with purely acute cardiopulmonary problems, except in the case of underlying chronic cardiopulmonary disease or severe right ventricular dysfunction leading to hepatic congestion.

Vascular Pulse deficits may reflect underlying chronic atherosclerosis, which increases the likelihood of coronary artery disease. However, evidence of acute limb ischemia with loss of the pulse and pallor, particularly in the upper extremities, can indicate catastrophic consequences of aortic dissection. Unilateral lower-extremity swelling should raise suspicion about venous thromboembolism.

Musculoskeletal Pain arising from the costochondral and chondrosternal articulations may be associated with localized swelling, redness, or marked localized tenderness. Pain on palpation of these joints is usually well localized and is a useful clinical sign, though deep palpation may elicit pain in the absence of costochondritis. Although palpation of the chest wall often elicits pain in patients with various musculoskeletal conditions, it should be appreciated that chest wall tenderness does not exclude myocardial ischemia. Sensory deficits in the upper extremities may be indicative of cervical disk disease.

ELECTROCARDIOGRAPHY

Electrocardiography is crucial in the evaluation of nontraumatic chest discomfort. The ECG is pivotal for identifying patients with ongoing ischemia as the principal reason for their presentation as well as secondary cardiac complications of other disorders. Professional society guidelines recommend that an ECG be obtained within 10 min of presentation, with the primary goal of identifying patients with ST-segment elevation diagnostic of MI who are candidates for immediate interventions to restore flow in the occluded coronary artery. ST-segment depression and symmetric T-wave inversions at least 0.2 mV in depth are useful for

detecting myocardial ischemia in the absence of STEMI and are also indicative of higher risk of death or recurrent ischemia. Serial performance of ECGs (every 30–60 min) is recommended in the ED evaluation of suspected ACS. In addition, an ECG with right-sided lead placement should be considered in patients with clinically suspected ischemia and a nondiagnostic standard 12-lead ECG. Despite the value of the resting ECG, its sensitivity for ischemia is poor—as low as 20% in some studies.

Abnormalities of the ST segment and T wave may occur in a variety of conditions, including pulmonary embolism, ventricular hypertrophy, acute and chronic pericarditis, myocarditis, electrolyte imbalance, and metabolic disorders. Notably, hyperventilation associated with panic disorder can also lead to nonspecific ST and T-wave abnormalities. Pulmonary embolism is most often associated with sinus tachycardia but can also lead to rightward shift of the ECG axis, manifesting as an S-wave in lead I, with a Q-wave and T-wave in lead III (Chaps. 268 and 300). In patients with ST-segment elevation, the presence of diffuse lead involvement not corresponding to a specific coronary anatomic distribution and PR-segment depression can aid in distinguishing pericarditis from acute MI.

CHEST RADIOGRAPHY

(See Chap. 308e) Plain radiography of the chest is performed routinely when patients present with acute chest discomfort and selectively when individuals who are being evaluated as outpatients have subacute or chronic pain. The chest radiograph is most useful for identifying pulmonary processes, such as pneumonia or pneumothorax. Findings are often unremarkable in patients with ACS, but pulmonary edema may be evident. Other specific findings include widening of the mediastinum in some patients with aortic dissection, Hampton's hump or Westermarck's sign in patients with pulmonary embolism (Chaps. 300 and 308e), or pericardial calcification in chronic pericarditis.

CARDIAC BIOMARKERS

Laboratory testing in patients with acute chest pain is focused on the detection of myocardial injury. Such injury can be detected by the presence of circulating proteins released from damaged myocardial cells. Owing to the time necessary for this release, initial biomarkers of injury may be in the normal range, even in patients with STEMI. Because of superior cardiac tissue-specificity compared with creatine kinase MB, cardiac troponin is the preferred biomarker for the diagnosis of MI and should be measured in all patients with suspected ACS at presentation and repeated in 3–6 h. Testing after 6 h is required only when there is uncertainty regarding the onset of pain or when stuttering symptoms have occurred. It is not necessary or advisable to measure troponin in patients without suspicion of ACS unless this test is being used specifically for risk stratification (e.g., in pulmonary embolism or heart failure).

The development of cardiac troponin assays with progressively greater analytical sensitivity has facilitated detection of substantially lower blood concentrations of troponin than was previously possible. This evolution permits earlier detection of myocardial injury, enhances the overall accuracy of a diagnosis of MI, and improves risk stratification in suspected ACS. The greater negative predictive value of a negative troponin result with current-generation assays is an advantage in the evaluation of chest pain in the ED. Rapid rule-out protocols that use serial testing and changes in troponin concentration over as short a period as 1–2 h appear promising and remain under investigation. However, with these advantages has come a trade-off: myocardial injury is detected in a larger proportion of patients who have non-ACS cardiopulmonary conditions than with previous, less sensitive assays. This evolution in testing for myocardial necrosis has rendered other aspects of the clinical evaluation critical to the practitioner's determination of the probability that the symptoms represent ACS. In addition, observation of a change in cardiac troponin concentration between serial samples is useful in discriminating acute causes of myocardial injury from chronic elevation due to underlying structural heart disease,

end-stage renal disease, or interfering antibodies. The diagnosis of MI is reserved for acute myocardial injury that is marked by a rising and/or falling pattern—with at least one value exceeding the 99th percentile reference limit—and that is caused by ischemia. Other non-ischemic insults, such as myocarditis, may result in myocardial injury but should not be labeled MI.

Other laboratory assessments may include the D-dimer test to aid in exclusion of pulmonary embolism (Chap. 300). Measurement of a B-type natriuretic peptide is useful when considered in conjunction with the clinical history and exam for the diagnosis of heart failure. B-type natriuretic peptides also provide prognostic information regarding patients with ACS and those with pulmonary embolism. Other putative biomarkers of acute myocardial ischemia or ACS, such as myeloperoxidase, have not been adopted in routine use.

INTEGRATIVE DECISION-AIDS

Multiple clinical algorithms have been developed to aid in decision-making during the evaluation and disposition of patients with acute nontraumatic chest pain. Such decision-aids have been derived on the basis of their capacity to estimate either of two closely related but not identical probabilities: (1) the probability of a final diagnosis of ACS and (2) the probability of major cardiac events during short-term follow-up. Such decision-aids are used most commonly to identify patients with a low clinical probability of ACS who are candidates either for early provocative testing for ischemia or for discharge from the ED. Goldman and Lee developed one of the first such decision-aids, using only the ECG and risk indicators—hypotension, pulmonary rales, and known ischemic heart disease—to categorize patients into four risk categories ranging from a <1% to a >16% probability of a major cardiovascular complication. The Acute Cardiac Ischemia Time-Insensitive Predictive Instrument (ACI-TIPI) combines age, sex, chest pain presence, and ST-segment abnormalities to define a probability of ACS. More recently developed decision-aids are shown in Fig. 19-3. Elements common to each of these tools are (1) symptoms typical for ACS; (2) older age; (3) risk factors for or known atherosclerosis; (4) ischemic ECG abnormalities; and (5) elevated cardiac troponin levels. Although, because of very low specificity, the overall diagnostic performance of such decision-aids is poor (area under the receiver operating curve, 0.55–0.65), they can help identify patients with a very low probability of ACS (e.g., <1%). Nevertheless, no such decision-aid (or single clinical factor) is sufficiently sensitive and well validated to use as a sole tool for clinical decision-making.

Clinicians should differentiate between the algorithms discussed above and risk scores derived for stratification of prognosis (e.g., the TIMI and GRACE risk scores, Chap. 295) in patients *who already have an established diagnosis of ACS*. The latter risk scores were not designed to be used for *diagnostic* assessment.

PROVOCATIVE TESTING FOR ISCHEMIA

Exercise electrocardiography (“stress testing”) is commonly employed for completion of risk stratification of patients who have undergone an initial evaluation that has not revealed a specific cause of chest discomfort and has identified them as being at low or selectively intermediate risk of ACS. Early exercise testing is safe in patients without high-risk findings after 8–12 h of observation and can assist in refining their prognostic assessment. For example, of low-risk patients who underwent exercise testing in the first 48 h after presentation, those without evidence of ischemia had a 2% rate of cardiac events through 6 months, whereas the rate was 15% among patients with either clear evidence of ischemia or an equivocal result. Patients who are unable to exercise may undergo pharmacological stress testing with either nuclear perfusion imaging or echocardiography. Notably, some experts have deemed the routine use of stress testing for low-risk patients unsupported by direct clinical evidence and a potentially unnecessary source of cost.

Professional society guidelines identify ongoing chest pain as a contraindication to stress testing. In selected patients with persistent pain and nondiagnostic ECG and biomarker data, resting

HEART Score		
History	Highly suspicious	2
	Moderately suspicious	1
	Slightly suspicious	0
ECG	Significant ST-depression	2
	Non-specific abnormality	1
	Normal	0
Age	≥65 y	2
	45–<65 y	1
	<45 y	0
Risk factors	≥3 risk factors	2
	1–2 risk factors	1
	None	0
Troponin (serial)	≥3 × 99th percentile	2
	1–<3 × 99th percentile	1
	≤99th percentile	0
TOTAL		
Low-risk: 0–3		
Not low-risk: ≥4		

North American Chest Pain Rule	
High Risk Criteria	Y/N
Typical symptoms for ischemia	
ECG: acute ischemic changes	
Age ≥50 y	
Known coronary artery disease	
Troponin (serial) >99th percentile	
Low-risk: All No	
Not Low-risk: Any Yes	

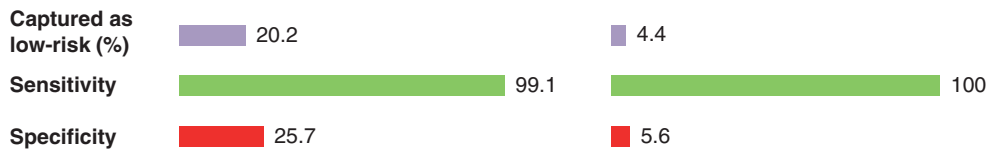


FIGURE 19-3 Examples of decision-aids used in conjunction with serial measurement of cardiac troponin for evaluation of acute chest pain. (Figure prepared from data in SA Mahler et al: *Int J Cardiol* 168:795, 2013.)

myocardial perfusion images can be obtained; the absence of any perfusion abnormality substantially reduces the likelihood of coronary artery disease. In some centers, early myocardial perfusion imaging is performed as part of a routine strategy for evaluating patients at low or intermediate risk of ACS in parallel with other testing. Management of patients with normal perfusion images can be expedited with earlier discharge and outpatient stress testing, if indicated. Those with abnormal rest perfusion imaging, which cannot discriminate between old or new myocardial defects, must undergo additional in-hospital evaluation.

OTHER NONINVASIVE STUDIES

Other noninvasive imaging studies of the chest can be used selectively to provide additional diagnostic and prognostic information on patients with chest discomfort.

Echocardiography Echocardiography is not necessarily routine in patients with chest discomfort. However, in patients with an uncertain diagnosis, particularly those with nondiagnostic ST elevation, ongoing symptoms, or hemodynamic instability, detection of abnormal regional wall motion provides evidence of possible ischemic dysfunction. Echocardiography is diagnostic in patients with mechanical complications of MI or in patients with pericardial tamponade. Transthoracic echocardiography is poorly sensitive for aortic dissection, although an intimal flap may sometimes be detected in the ascending aorta.

CT Angiography (See Chap. 270e) CT angiography is emerging as a modality for the evaluation of patients with acute chest discomfort. Coronary CT angiography is a sensitive technique for detection of obstructive coronary disease, particularly in the proximal third of the major epicardial coronary arteries. CT appears to enhance the speed to disposition of patients with a low-intermediate probability for ACS; its major strength being the negative predictive value of a finding of

no significant disease. In addition, contrast-enhanced CT can detect focal areas of myocardial injury in the acute setting as decreased areas of enhancement. At the same time, CT angiography can exclude aortic dissection, pericardial effusion, and pulmonary embolism. Balancing factors in the consideration of the emerging role of coronary CT angiography in low-risk patients are radiation exposure and additional testing prompted by nondiagnostic abnormal results.

MRI (See Chap. 270e) Cardiac magnetic resonance (CMR) imaging is an evolving, versatile technique for structural and functional evaluation of the heart and the vasculature of the chest. CMR accurately measures ventricular dimensions and function and can be performed as a modality for pharmacologic stress perfusion imaging. Gadolinium-enhanced CMR can provide early detection of MI, defining areas of myocardial necrosis accurately, and can delineate patterns of myocardial disease that are often useful in discriminating ischemic from non-ischemic myocardial injury. Although usually not practical for the urgent evaluation of acute chest discomfort, CMR can be a useful modality for cardiac structural evaluation of patients with elevated cardiac troponin levels in the absence of definite coronary artery disease. CMR coronary angiography is in its early stages. MRI also permits highly accurate assessment for aortic dissection but is infrequently used as the first test because CT and transesophageal echocardiography are usually more practical.

CRITICAL PATHWAYS FOR ACUTE CHEST DISCOMFORT

Because of the challenges inherent in reliably identifying the small proportion of patients with serious causes of acute chest discomfort while not exposing the larger number of low-risk patients to unnecessary testing and extended ED or hospital evaluations, many medical centers have adopted critical pathways to expedite the assessment and management of patients with nontraumatic chest pain, often in dedicated chest pain units. Such pathways are generally aimed at

(1) rapid identification, triage, and treatment of high-risk cardiopulmonary conditions (e.g., STEMI); (2) accurate identification of low-risk patients who can be safely observed in units with less intensive monitoring, undergo early exercise testing, or be discharged home; and (3) through more efficient and systematic accelerated diagnostic protocols, safe reduction in costs associated with overuse of testing and unnecessary hospitalizations. In some studies, provision of protocol-driven care in chest pain units has decreased costs and overall duration of hospital evaluation with no detectable excess of adverse clinical outcomes.

OUTPATIENT EVALUATION OF CHEST DISCOMFORT

Chest pain is common in outpatient practice, with a lifetime prevalence of 20–40% in the general population. More than 25% of patients with MI have had a related visit with a primary care physician in the previous month. The diagnostic principles are the same as in the ED. However, the pretest probability of an acute cardiopulmonary cause is significantly lower. Therefore, testing paradigms are less intense, with an emphasis on the history, physical examination, and ECG. Moreover, decision-aids developed for settings with a high prevalence of significant cardiopulmonary disease have lower positive predictive value when applied in the practitioner's office. However, in general, if the level of clinical suspicion of ACS is sufficiently high to consider troponin testing, the patient should be referred to the ED for evaluation.

20

Abdominal Pain

Danny O. Jacobs, William Silen

Correctly interpreting acute abdominal pain can be quite challenging. Few clinical situations require greater judgment, because the most catastrophic of events may be forecast by the subtlest of symptoms and signs. In every instance, the clinician must distinguish those conditions that require urgent intervention from those that do not and can best be managed nonoperatively. A meticulously executed, detailed history and physical examination are critically important for focusing the differential diagnosis, where necessary, and allowing the diagnostic evaluation to proceed expeditiously (Table 20-1).

The etiologic classification in Table 20-2, although not complete, provides a useful framework for evaluating patients with abdominal pain.

The most common causes of abdominal pain on admission are acute appendicitis, nonspecific abdominal pain, pain of urologic origin, and intestinal obstruction. A diagnosis of “acute or surgical abdomen” is not acceptable because of its often misleading and erroneous connotations. Most patients who present with acute abdominal pain will have self-limited disease processes. However, it is important to remember that pain severity does not necessarily correlate with the severity of the underlying condition. The most obvious of “acute abdomens” may

not require operative intervention, and the mildest of abdominal pains may herald an urgently correctable lesion. Any patient with abdominal pain of recent onset requires early and thorough evaluation and accurate diagnosis.

SOME MECHANISMS OF PAIN ORIGINATING IN THE ABDOMEN

Inflammation of the Parietal Peritoneum The pain of parietal peritoneal inflammation is steady and aching in character and is located directly over the inflamed area, its exact reference being possible because it is transmitted by somatic nerves supplying the parietal peritoneum. The intensity of the pain is dependent on the type and amount of material to which the peritoneal surfaces are exposed in a given time period. For example, the sudden release into the peritoneal cavity of a small quantity of sterile acid gastric juice causes much more pain than the same amount of grossly contaminated neutral feces. Enzymatically active pancreatic juice incites more pain and inflammation than does the same amount of sterile bile containing no potent enzymes. Blood is normally only a mild irritant and the response to urine can be bland, so exposure of blood and urine to the peritoneal cavity may go unnoticed unless it is sudden and massive. Bacterial contamination, such as may occur with pelvic inflammatory disease or perforated distal intestine, causes low-intensity pain until multiplication causes a significant amount of inflammatory mediators to be released. Patients with perforated upper gastrointestinal ulcers may present entirely differently depending on how quickly gastric juices enter the peritoneal cavity. Thus, the rate at which any inflammatory material irritates the peritoneum is important.

The pain of peritoneal inflammation is invariably accentuated by pressure or changes in tension of the peritoneum, whether produced by palpation or by movement such as with coughing or sneezing. The patient with peritonitis characteristically lies quietly in bed, preferring to avoid motion, in contrast to the patient with colic, who may be thrashing in discomfort.

Another characteristic feature of peritoneal irritation is tonic reflex spasm of the abdominal musculature, localized to the involved body segment. Its intensity depends on the integrity of the nervous system, the location of the inflammatory process, and the rate at which it develops. Spasm over a perforated retrocecal appendix or perforation into the lesser peritoneal sac may be minimal or absent because of the protective effect of overlying viscera. Catastrophic abdominal emergencies may be associated with minimal or no detectable pain or muscle spasm in obtunded, seriously ill, debilitated, immunosuppressed, or psychotic patients. A slowly developing process also often greatly attenuates the degree of muscle spasm.

Obstruction of Hollow Viscera Intraluminal obstruction classically elicits intermittent or colicky abdominal pain that is not as well localized as the pain of parietal peritoneal irritation. However, the absence of cramping discomfort should not be misleading because distention of a hollow viscus may also produce steady pain with only rare paroxysms.

Small-bowel obstruction often presents as poorly localized, intermittent periumbilical or supraumbilical pain. As the intestine progressively dilates and loses muscular tone, the colicky nature of the pain may diminish. With superimposed strangulating obstruction, pain may spread to the lower lumbar region if there is traction on the root of the mesentery. The colicky pain of colonic obstruction is of lesser intensity, is commonly located in the infraumbilical area, and may often radiate to the lumbar region.

Sudden distention of the biliary tree produces a steady rather than colicky type of pain; hence, the term *biliary colic* is misleading. Acute distention of the gallbladder usually causes pain in the right upper quadrant with radiation to the right posterior region of the thorax or to the tip of the right scapula, but it is also not uncommonly found near the midline. Distention of the common bile duct often causes epigastric pain that may radiate to the upper lumbar region. Considerable variation is common, however, so that differentiation between these may be impossible. The typical subscapular pain or lumbar radiation is frequently absent. Gradual dilatation of the biliary tree, as can occur with carcinoma of the head of the pancreas, may cause no pain

TABLE 20-1 SOME KEY COMPONENTS OF THE PATIENT'S HISTORY

Age
Time and mode of onset of the pain
Pain characteristics
Duration of symptoms
Location of pain and sites of radiation
Associated symptoms and their relationship to the pain
Nausea, emesis, and anorexia
Diarrhea, constipation, or other changes in bowel habits
Menstrual history

TABLE 20-2 SOME IMPORTANT CAUSES OF ABDOMINAL PAIN**Pain Originating in the Abdomen**

Parietal peritoneal inflammation	Vascular disturbances
Bacterial contamination	Embolism or thrombosis
Perforated appendix or other perforated viscus	Vascular rupture
Pelvic inflammatory disease	Pressure or torsional occlusion
Chemical irritation	Sickle cell anemia
Perforated ulcer	Abdominal wall
Pancreatitis	Distortion or traction of mesentery
Mittelschmerz	Trauma or infection of muscles
Mechanical obstruction of hollow viscera	Distension of visceral surfaces, e.g., by hemorrhage
Obstruction of the small or large intestine	Hepatic or renal capsules
Obstruction of the biliary tree	Inflammation
Obstruction of the ureter	Appendicitis
	Typhoid fever
	Neutropenic enterocolitis or “typhlitis”

Pain Referred from Extraabdominal Source

Cardiothoracic	Pleurodynia
Acute myocardial infarction	Pneumothorax
Myocarditis, endocarditis, pericarditis	Empyema
Congestive heart failure	Esophageal disease, including spasm, rupture, or inflammation
Pneumonia (especially lower lobes)	Genitalia
Pulmonary embolus	Torsion of the testis

Metabolic Causes

Diabetes	Acute adrenal insufficiency
Uremia	Familial Mediterranean fever
Hyperlipidemia	Porphyria
Hyperparathyroidism	C1 esterase inhibitor deficiency (angioneurotic edema)

Neurologic/Psychiatric Causes

Herpes zoster	Spinal cord or nerve root compression
Tabes dorsalis	Functional disorders
Causalgia	Psychiatric disorders
Radiculitis from infection or arthritis	

Toxic Causes

Lead poisoning
Insect or animal envenomation
Black widow spider bites
Snake bites

Uncertain Mechanisms

Narcotic withdrawal
Heat stroke

or only a mild aching sensation in the epigastrium or right upper quadrant. The pain of distention of the pancreatic ducts is similar to that described for distention of the common bile duct but, in addition, is very frequently accentuated by recumbency and relieved by the upright position.

Obstruction of the urinary bladder usually causes dull, low-intensity pain in the suprapubic region. Restlessness without specific complaint of pain may be the only sign of a distended bladder in an obtunded patient. In contrast, acute obstruction of the intravesicular portion of the ureter is characterized by severe suprapubic and flank pain that radiates to the penis, scrotum, or inner aspect of the upper thigh. Obstruction of the ureteropelvic junction manifests as pain near the costovertebral angle, whereas obstruction of the remainder of the ureter is associated with flank pain that often extends into the same side of the abdomen.

Vascular Disturbances A frequent misconception is that pain due to intraabdominal vascular disturbances is sudden and catastrophic in nature. Certain disease processes, such as embolism or thrombosis of the superior mesenteric artery or impending rupture of an abdominal

aortic aneurysm, can certainly be associated with diffuse, severe pain. Yet, just as frequently, the patient with occlusion of the superior mesenteric artery only has mild continuous or cramping diffuse pain for 2 or 3 days before vascular collapse or findings of peritoneal inflammation appear. The early, seemingly insignificant discomfort is caused by hyperperistalsis rather than peritoneal inflammation. Indeed, absence of tenderness and rigidity in the presence of continuous, diffuse pain (e.g., “pain out of proportion to physical findings”) in a patient likely to have vascular disease is quite characteristic of occlusion of the superior mesenteric artery. Abdominal pain with radiation to the sacral region, flank, or genitalia should always signal the possible presence of a rupturing abdominal aortic aneurysm. This pain may persist over a period of several days before rupture and collapse occur.

Abdominal Wall Pain arising from the abdominal wall is usually constant and aching. Movement, prolonged standing, and pressure accentuate the discomfort and associated muscle spasm. In the case of hematoma of the rectus sheath, now most frequently encountered in association with anticoagulant therapy, a mass may be present in the lower quadrants of the abdomen. Simultaneous involvement of

muscles in other parts of the body usually serves to differentiate myositis of the abdominal wall from other processes that might cause pain in the same region.

REFERRED PAIN IN ABDOMINAL DISEASE

Pain referred to the abdomen from the thorax, spine, or genitalia may present a vexing diagnostic challenge because diseases of the upper part of the abdominal cavity such as acute cholecystitis or perforated ulcer may be associated with intrathoracic complications. A most important, yet often forgotten, dictum is that the possibility of intrathoracic disease must be considered in every patient with abdominal pain, especially if the pain is in the upper abdomen.

Systematic questioning and examination directed toward detecting myocardial or pulmonary infarction, pneumonia, pericarditis, or esophageal disease (the intrathoracic diseases that most often masquerade as abdominal emergencies) will often provide sufficient clues to establish the proper diagnosis. Diaphragmatic pleuritis resulting from pneumonia or pulmonary infarction may cause pain in the right upper quadrant and pain in the supraclavicular area, the latter radiation to be distinguished from the referred subscapular pain caused by acute distention of the extrahepatic biliary tree. The ultimate decision as to the origin of abdominal pain may require deliberate and planned observation over a period of several hours, during which repeated questioning and examination will provide the diagnosis or suggest the appropriate studies.

Referred pain of thoracic origin is often accompanied by splinting of the involved hemithorax with respiratory lag and decrease in excursion more marked than that seen in the presence of intraabdominal disease. In addition, apparent abdominal muscle spasm caused by referred pain will diminish during the inspiratory phase of respiration, whereas it persists throughout both respiratory phases if it is of abdominal origin. Palpation over the area of referred pain in the abdomen also does not usually accentuate the pain and, in many instances, actually seems to relieve it.

Thoracic disease and abdominal disease frequently coexist and may be difficult or impossible to differentiate. For example, the patient with known biliary tract disease often has epigastric pain during myocardial infarction, or biliary colic may be referred to the precordium or left shoulder in a patient who has suffered previously from angina pectoris.

For an explanation of the radiation of pain to a previously diseased area, see Chap. 18.

Referred pain from the spine, which usually involves compression or irritation of nerve roots, is characteristically intensified by certain motions such as cough, sneeze, or strain and is associated with hyperesthesia over the involved dermatomes. Pain referred to the abdomen from the testes or seminal vesicles is generally accentuated by the slightest pressure on either of these organs. The abdominal discomfort experienced is of dull, aching character and is poorly localized.

METABOLIC ABDOMINAL CRISES

Pain of metabolic origin may simulate almost any other type of intraabdominal disease. Several mechanisms may be at work. In certain instances, such as hyperlipidemia, the metabolic disease itself may be accompanied by an intraabdominal process such as pancreatitis, which can lead to unnecessary laparotomy unless recognized. C1 esterase deficiency associated with angioneurotic edema is often associated with episodes of severe abdominal pain. Whenever the cause of abdominal pain is obscure, a metabolic origin always must be considered. Abdominal pain is also the hallmark of familial Mediterranean fever (**Chap. 392**).

The problem of differential diagnosis is often not readily resolved. The pain of porphyria and of lead colic is usually difficult to distinguish from that of intestinal obstruction, because severe hyperperistalsis is a prominent feature of both. The pain of uremia or diabetes is nonspecific, and the pain and tenderness frequently shift in location and intensity. Diabetic acidosis may be precipitated by acute appendicitis or intestinal obstruction, so if prompt resolution of the abdominal pain does not result from correction of the metabolic abnormalities, an underlying organic problem should be suspected. Black widow spider

bites produce intense pain and rigidity of the abdominal muscles and back, an area infrequently involved in intraabdominal disease.

IMMUNOCOMPROMISE

Evaluating and diagnosing causes of abdominal pain in immunosuppressed or otherwise immunocompromised patients is very difficult. This includes those who have undergone organ transplantation; who are receiving immunosuppressive treatments for autoimmune diseases, chemotherapy, or glucocorticoids; who have AIDS; and who are very old. In these circumstances, normal physiologic responses may be absent or masked. In addition, unusual infections may cause abdominal pain where the etiologic agents include cytomegalovirus, mycobacteria, protozoa, and fungi. These pathogens may affect all gastrointestinal organs, including the gallbladder, liver, and pancreas, as well as the gastrointestinal tract, causing occult or overtly symptomatic perforations of the latter. Splenic abscesses due to *Candida* or *Salmonella* infection should also be considered, especially when evaluating patients with left upper quadrant or left flank pain. Acalculous cholecystitis is a relative common complication in patients with AIDS, where it is often associated with cryptosporidiosis or cytomegalovirus infection.

Neutropenic enterocolitis is often identified as a cause of abdominal pain and fever in some patients with bone marrow suppression due to chemotherapy. Acute graft-versus-host disease should be considered. Optimal management of these patients may require meticulous follow-up including serial examinations to be certain that surgical intervention is not required to treat an underlying disease process.

NEUROGENIC CAUSES

Diseases that injure sensory nerves may cause causalgic pain. It has a burning character and is usually limited to the distribution of a given peripheral nerve. Normal nonpainful stimuli such as touch or a change in temperature may be causalgic and may frequently be present even at rest. The demonstration of irregularly spaced cutaneous pain spots may be the only indication that an old nerve injury exists. Even though the pain may be precipitated by gentle palpation, rigidity of the abdominal muscles is absent, and the respirations are not disturbed. Distention of the abdomen is uncommon, and the pain has no relationship to the intake of food.

Pain arising from spinal nerves or roots comes and goes suddenly and is of a lancinating type (**Chap. 22**). It may be caused by herpes zoster, impingement by arthritis, tumors, a herniated nucleus pulposus, diabetes, or syphilis. It is not associated with food intake, abdominal distention, or changes in respiration. Severe muscle spasm, as in the gastric crises of tabes dorsalis, is common but is either relieved or not accentuated by abdominal palpation. The pain is made worse by movement of the spine and is usually confined to a few dermatomes. Hyperesthesia is very common.

Pain due to functional causes conforms to none of the aforementioned patterns. Mechanisms of disease are not clearly established. Irritable bowel syndrome (IBS) is a functional gastrointestinal disorder characterized by abdominal pain and altered bowel habits. The diagnosis is made on the basis of clinical criteria (**Chap. 352**) and after exclusion of demonstrable structural abnormalities. The episodes of abdominal pain are often brought on by stress, and the pain varies considerably in type and location. Nausea and vomiting are rare. Localized tenderness and muscle spasm are inconsistent or absent. The causes of IBS or related functional disorders are not known.

APPROACH TO THE PATIENT: Abdominal Pain

Few abdominal conditions require such urgent operative intervention that an orderly approach need be abandoned, no matter how ill the patient. Only patients with exsanguinating intraabdominal hemorrhage (e.g., ruptured aneurysm) must be rushed to the operating room immediately, but in such instances, only a few minutes are required to assess the critical nature of the problem. Under these

circumstances, all obstacles must be swept aside, adequate venous access for fluid replacement obtained, and the operation begun. Many of these patients have died in the radiology department or the emergency room while awaiting unnecessary examinations such as electrocardiograms or computed tomography (CT) scans. *There are no contraindications to operation when massive intraabdominal hemorrhage is present.* Fortunately, this situation is relatively rare. This statement does not necessarily apply to patients with intraluminal gastrointestinal hemorrhage, who can often be managed by other means (Chap. 57). Nothing will supplant an orderly, painstakingly detailed history, which is far more valuable than any laboratory or radiographic examination. This kind of history is laborious and time-consuming, making it not especially popular, even though a reasonably accurate diagnosis can be made on the basis of the history alone in the majority of cases.

In cases of acute abdominal pain, a diagnosis is readily established in most instances, whereas success is not so frequent in patients with chronic pain. IBS is one of the most common causes of abdominal pain and must always be kept in mind (Chap. 352). The location of the pain can assist in narrowing the differential diagnosis (Table 20-3); however, the chronological sequence of events in the patient's history is often more important than the pain's location. If the examiner is sufficiently open-minded and unhurried, asks the proper questions, and listens, the patient will usually provide the diagnosis. Careful attention should be paid to the extraabdominal regions. Narcotics or analgesics should not be withheld until a definitive diagnosis or a definitive plan has been formulated; obfuscation of the diagnosis by adequate analgesia is unlikely.

An accurate menstrual history in a female patient is essential. It is important to remember that normal anatomic relationships can be significantly altered by the gravid uterus. Abdominal and pelvic pain may occur during pregnancy due to conditions that do

not require surgery. Lastly, some otherwise noteworthy laboratory values (e.g., leukocytosis) may represent the normal physiologic changes of pregnancy.

In the examination, simple critical inspection of the patient, e.g., of facies, position in bed, and respiratory activity, provides valuable clues. The amount of information to be gleaned is directly proportional to the gentleness and thoroughness of the examiner. Once a patient with peritoneal inflammation has been examined brusquely, accurate assessment by the next examiner becomes almost impossible. Eliciting rebound tenderness by sudden release of a deeply palpating hand in a patient with suspected peritonitis is cruel and unnecessary. The same information can be obtained by gentle percussion of the abdomen (rebound tenderness on a miniature scale), a maneuver that can be far more precise and localizing. Asking the patient to cough will elicit true rebound tenderness without the need for placing a hand on the abdomen. Furthermore, the forceful demonstration of rebound tenderness will startle and induce protective spasm in a nervous or worried patient in whom true rebound tenderness is not present. A palpable gallbladder will be missed if palpation is so aggressive that voluntary muscle spasm becomes superimposed on involuntary muscular rigidity. As with history taking, sufficient time should be spent in the examination. Abdominal signs may be minimal but nevertheless, if accompanied by consistent symptoms, may be exceptionally meaningful. Abdominal signs may be virtually or totally absent in cases of pelvic peritonitis, so careful pelvic and rectal examinations are mandatory in every patient with abdominal pain. Tenderness on pelvic or rectal examination in the absence of other abdominal signs can be caused by operative indications such as perforated appendicitis, diverticulitis, twisted ovarian cyst, and many others. Much attention has been paid to the presence or absence of peristaltic sounds, their quality, and their frequency. Auscultation of the abdomen is one of the least revealing aspects of the physical examination of a patient with abdominal pain. Catastrophes such as a strangulating small intestinal obstruction or perforated appendicitis may occur in the presence of normal peristaltic sounds. Conversely, when the proximal part of the intestine above obstruction becomes markedly distended and edematous, peristaltic sounds may lose the characteristics of borborygmi and become weak or absent, even when peritonitis is not present. It is usually the severe chemical peritonitis of sudden onset that is associated with the truly silent abdomen.

Laboratory examinations may be valuable in assessing the patient with abdominal pain, yet, with few exceptions, they rarely establish a diagnosis. Leukocytosis should never be the single deciding factor as to whether or not operation is indicated. A white blood cell count >20,000/ μ L may be observed with perforation of a viscus, but pancreatitis, acute cholecystitis, pelvic inflammatory disease, and intestinal infarction may also be associated with marked leukocytosis. A normal white blood cell count is not rare in cases of perforation of abdominal viscera. The diagnosis of anemia may be more helpful than the white blood cell count, especially when combined with the history.

The urinalysis may reveal the state of hydration or rule out severe renal disease, diabetes, or urinary infection. Blood urea nitrogen, glucose, and serum bilirubin levels may be helpful. Serum amylase levels may be increased by many diseases other than pancreatitis, e.g., perforated ulcer, strangulating intestinal obstruction, and acute cholecystitis; thus, elevations of serum amylase do not rule out the need for an operation.

Plain and upright or lateral decubitus radiographs of the abdomen may be of value in cases of intestinal obstruction, perforated ulcer, and a variety of other conditions. They are usually unnecessary in patients with acute appendicitis or strangulated external hernias. In rare instances, barium or water-soluble contrast study of the upper part of the gastrointestinal tract may demonstrate partial intestinal obstruction that may elude diagnosis by other means. If there is any question of obstruction of the colon, oral administration of barium sulfate should be avoided. On the other hand, in

TABLE 20-3 DIFFERENTIAL DIAGNOSES OF ABDOMINAL PAIN BY LOCATION

Right Upper Quadrant	Epigastric	Left Upper Quadrant
Cholecystitis	Peptic ulcer disease	Splenic infarct
Cholangitis	Gastritis	Splenic rupture
Pancreatitis	GERD	Splenic abscess
Pneumonia/empyema	Pancreatitis	Gastritis
Pleurisy/pleurodynia	Myocardial infarction	Gastric ulcer
Subdiaphragmatic abscess	Pericarditis	Pancreatitis
Hepatitis	Ruptured aortic aneurysm	Subdiaphragmatic abscess
Budd-Chiari syndrome	Esophagitis	
Right Lower Quadrant	Periumbilical	Left Lower Quadrant
Appendicitis	Early appendicitis	Diverticulitis
Salpingitis	Gastroenteritis	Salpingitis
Inguinal hernia	Bowel obstruction	Inguinal hernia
Ectopic pregnancy	Ruptured aortic aneurysm	Ectopic pregnancy
Nephrolithiasis		Nephrolithiasis
Inflammatory bowel disease		Irritable bowel syndrome
Mesenteric lymphadenitis		Inflammatory bowel disease
Typhlitis		
Diffuse Nonlocalized Pain		
Gastroenteritis	Malaria	
Mesenteric ischemia	Familial Mediterranean fever	
Bowel obstruction		
Irritable bowel syndrome	Metabolic diseases	
Peritonitis	Psychiatric disease	
Diabetes		

Abbreviation: GERD, gastroesophageal reflux disease.

cases of suspected colonic obstruction (without perforation), a contrast enema may be diagnostic.

In the absence of trauma, peritoneal lavage has been replaced as a diagnostic tool by CT scanning and laparoscopy. Ultrasonography has proved to be useful in detecting an enlarged gallbladder or pancreas, the presence of gallstones, an enlarged ovary, or a tubal pregnancy. Laparoscopy is especially helpful in diagnosing pelvic conditions, such as ovarian cysts, tubal pregnancies, salpingitis, and acute appendicitis.

Radioisotopic hepatobiliary iminodiacetic acid scans (HIDAs) may help differentiate acute cholecystitis or biliary colic from acute pancreatitis. A CT scan may demonstrate an enlarged pancreas, ruptured spleen, or thickened colonic or appendiceal wall and streaking of the mesocolon or mesoappendix characteristic of diverticulitis or appendicitis.

Sometimes, even under the best circumstances with all available aids and with the greatest of clinical skill, a definitive diagnosis cannot be established at the time of the initial examination. Nevertheless, even in the absence of a clear anatomic diagnosis, it may be abundantly clear to an experienced and thoughtful physician and surgeon that operation is indicated on clinical grounds alone. Should that decision be questionable, watchful waiting with repeated questioning and examination will often elucidate the true nature of the illness and indicate the proper course of action.

21 Headache

Peter J. Goadsby, Neil H. Raskin

Headache is among the most common reasons patients seek medical attention, on a global basis being responsible for more disability than any other neurologic problem. Diagnosis and management are based on a careful clinical approach augmented by an understanding of the anatomy, physiology, and pharmacology of the nervous system pathways mediating the various headache syndromes. This chapter will focus on the general approach to a patient with headache; migraine and other primary headache disorders are discussed in [Chap. 447](#).

GENERAL PRINCIPLES

A classification system developed by the International Headache Society (www.ihs-headache.org/) characterizes headache as primary or secondary ([Table 21-1](#)). *Primary headaches* are those in which headache and its associated features are the disorder in itself, whereas *secondary headaches* are those caused by exogenous disorders (Headache Classification Committee of the International Headache Society, 2013). Primary headache often results in considerable disability and a decrease in the patient's quality of life. Mild secondary headache, such as that seen in association with upper respiratory tract infections, is

TABLE 21-1 COMMON CAUSES OF HEADACHE

Primary Headache		Secondary Headache	
Type	%	Type	%
Tension-type	69	Systemic infection	63
Migraine	16	Head injury	4
Idiopathic stabbing	2	Vascular disorders	1
Exertional	1	Subarachnoid hemorrhage	<1
Cluster	0.1	Brain tumor	0.1

Source: After J Olesen et al: *The Headaches*. Philadelphia, Lippincott Williams & Wilkins, 2005.

common but rarely worrisome. Life-threatening headache is relatively uncommon, but vigilance is required in order to recognize and appropriately treat such patients.

ANATOMY AND PHYSIOLOGY OF HEADACHE

Pain usually occurs when peripheral nociceptors are stimulated in response to tissue injury, visceral distension, or other factors ([Chap. 18](#)). In such situations, pain perception is a normal physiologic response mediated by a healthy nervous system. Pain can also result when pain-producing pathways of the peripheral or central nervous system (CNS) are damaged or activated inappropriately. Headache may originate from either or both mechanisms. Relatively few cranial structures are pain-producing; these include the scalp, middle meningeal artery, dural sinuses, falx cerebri, and proximal segments of the large pial arteries. The ventricular ependyma, choroid plexus, pial veins, and much of the brain parenchyma are not pain-producing.

The key structures involved in primary headache appear to be the following:

- The large intracranial vessels and dura mater and the peripheral terminals of the trigeminal nerve that innervate these structures
- The caudal portion of the trigeminal nucleus, which extends into the dorsal horns of the upper cervical spinal cord and receives input from the first and second cervical nerve roots (the trigeminocervical complex)
- Rostral pain-processing regions, such as the ventroposteromedial thalamus and the cortex
- The pain-modulatory systems in the brain that modulate input from trigeminal nociceptors at all levels of the pain-processing pathways and influence vegetative functions, such as hypothalamus and brainstem structures

The innervation of the large intracranial vessels and dura mater by the trigeminal nerve is known as the *trigeminovascular system*. Cranial autonomic symptoms, such as *lacrimation*, *conjunctival injection*, *nasal congestion*, *rhinorrhea*, *periorbital swelling*, *aural fullness*, and *ptosis*, are prominent in the trigeminal autonomic cephalalgias, including cluster headache and paroxysmal hemicrania, and may also be seen in migraine, even in children. These autonomic symptoms reflect activation of cranial parasympathetic pathways, and functional imaging studies indicate that vascular changes in migraine and cluster headache, when present, are similarly driven by these cranial autonomic systems. Moreover, they can often be mistaken for symptoms or signs of cranial sinus inflammation, which is thus overdiagnosed and inappropriately managed. Migraine and other primary headache types are not “vascular headaches”; these disorders do not reliably manifest vascular changes, and treatment outcomes cannot be predicted by vascular effects. Migraine is a brain disorder and is best understood and managed as such.

CLINICAL EVALUATION OF ACUTE, NEW-ONSET HEADACHE

The patient who presents with a new, severe headache has a differential diagnosis that is quite different from the patient with recurrent headaches over many years. In new-onset and severe headache, the probability of finding a potentially serious cause is considerably greater than in recurrent headache. Patients with recent onset of pain require prompt evaluation and appropriate treatment. Serious causes to be considered include meningitis, subarachnoid hemorrhage, epidural or subdural hematoma, glaucoma, tumor, and purulent sinusitis. When worrisome symptoms and signs are present ([Table 21-2](#)), rapid diagnosis and management are critical.

A careful neurologic examination is an essential first step in the evaluation. In most cases, patients with an abnormal examination or a history of recent-onset headache should be evaluated by a computed tomography (CT) or magnetic resonance imaging (MRI) study. As an initial screening procedure for intracranial pathology in this setting, CT and MRI methods appear to be equally sensitive. In some circumstances, a lumbar puncture (LP) is also required, unless a benign etiology can be otherwise established. A general evaluation of acute headache might include cranial arteries by palpation; cervical spine by

TABLE 21-2 HEADACHE SYMPTOMS THAT SUGGEST A SERIOUS UNDERLYING DISORDER

Sudden-onset headache
First severe headache
“Worst” headache ever
Vomiting that precedes headache
Subacute worsening over days or weeks
Pain induced by bending, lifting, cough
Pain that disturbs sleep or presents immediately upon awakening
Known systemic illness
Onset after age 55
Fever or unexplained systemic signs
Abnormal neurologic examination
Pain associated with local tenderness, e.g., region of temporal artery

the effect of passive movement of the head and by imaging; the investigation of cardiovascular and renal status by blood pressure monitoring and urine examination; and eyes by funduscopy, intraocular pressure measurement, and refraction.

The psychological state of the patient should also be evaluated because a relationship exists between head pain and depression. This is intended to identify comorbidity rather than provide an explanation for the headache, because troublesome headache is seldom simply caused by mood change. Although it is notable that medicines with antidepressant actions are also effective in the prophylactic treatment of both tension-type headache and migraine, each symptom must be treated optimally.

Underlying recurrent headache disorders may be activated by pain that follows otologic or endodontic surgical procedures. Thus, pain about the head as the result of diseased tissue or trauma may reawaken an otherwise quiescent migraine syndrome. Treatment of the headache is largely ineffective until the cause of the primary problem is addressed.

Serious underlying conditions that are associated with headache are described below. Brain tumor is a rare cause of headache and even less commonly a cause of severe pain. The vast majority of patients presenting with severe headache have a benign cause.

SECONDARY HEADACHE

The management of secondary headache focuses on diagnosis and treatment of the underlying condition.

MENINGITIS

Acute, severe headache with stiff neck and fever suggests meningitis. LP is mandatory. Often there is striking accentuation of pain with eye movement. Meningitis can be easily mistaken for migraine in that the cardinal symptoms of pounding headache, photophobia, nausea, and vomiting are frequently present, perhaps reflecting the underlying biology of some of the patients.

Meningitis is discussed in Chaps. 164 and 165.

INTRACRANIAL HEMORRHAGE

Acute, severe headache with stiff neck but without fever suggests subarachnoid hemorrhage. A ruptured aneurysm, arteriovenous malformation, or intraparenchymal hemorrhage may also present with headache alone. Rarely, if the hemorrhage is small or below the foramen magnum, the head CT scan can be normal. Therefore, LP may be required to definitively diagnose subarachnoid hemorrhage.

Intracranial hemorrhage is discussed in Chap. 330.

BRAIN TUMOR

Approximately 30% of patients with brain tumors consider headache to be their chief complaint. The head pain is usually nondescript—an intermittent deep, dull aching of moderate intensity, which may worsen with exertion or change in position and may be associated with nausea and vomiting. This pattern of symptoms results from

migraine far more often than from brain tumor. The headache of brain tumor disturbs sleep in about 10% of patients. Vomiting that precedes the appearance of headache by weeks is highly characteristic of posterior fossa brain tumors. A history of amenorrhea or galactorrhea should lead one to question whether a prolactin-secreting pituitary adenoma (or the polycystic ovary syndrome) is the source of headache. Headache arising de novo in a patient with known malignancy suggests either cerebral metastases or carcinomatous meningitis, or both. Head pain appearing abruptly after bending, lifting, or coughing can be due to a posterior fossa mass, a Chiari malformation, or low cerebrospinal fluid (CSF) volume.

Brain tumors are discussed in Chap. 118.

TEMPORAL ARTERITIS

(See also Chaps. 39 and 385) Temporal (giant cell) arteritis is an inflammatory disorder of arteries that frequently involves the extracranial carotid circulation. It is a common disorder of the elderly; its annual incidence is 77 per 100,000 individuals age 50 and older. The average age of onset is 70 years, and women account for 65% of cases. About half of patients with untreated temporal arteritis develop blindness due to involvement of the ophthalmic artery and its branches; indeed, the ischemic optic neuropathy induced by giant cell arteritis is the major cause of rapidly developing bilateral blindness in patients >60 years. Because treatment with glucocorticoids is effective in preventing this complication, prompt recognition of the disorder is important.

Typical presenting symptoms include headache, polymyalgia rheumatica (Chap. 385), jaw claudication, fever, and weight loss. Headache is the dominant symptom and often appears in association with malaise and muscle aches. Head pain may be unilateral or bilateral and is located temporally in 50% of patients but may involve any and all aspects of the cranium. Pain usually appears gradually over a few hours before peak intensity is reached; occasionally, it is explosive in onset. The quality of pain is only seldom throbbing; it is almost invariably described as dull and boring, with superimposed episodic stabbing pains similar to the sharp pains that appear in migraine. Most patients can recognize that the origin of their head pain is superficial, external to the skull, rather than originating deep within the cranium (the pain site for migraineurs). Scalp tenderness is present, often to a marked degree; brushing the hair or resting the head on a pillow may be impossible because of pain. Headache is usually worse at night and often aggravated by exposure to cold. Additional findings may include reddened, tender nodules or red streaking of the skin overlying the temporal arteries, and tenderness of the temporal or, less commonly, the occipital arteries.

The erythrocyte sedimentation rate (ESR) is often, although not always, elevated; a normal ESR does not exclude giant cell arteritis. A temporal artery biopsy followed by immediate treatment with prednisone 80 mg daily for the first 4–6 weeks should be initiated when clinical suspicion is high. The prevalence of migraine among the elderly is substantial, considerably higher than that of giant cell arteritis. Migraineurs often report amelioration of their headaches with prednisone; thus, caution must be used when interpreting the therapeutic response.

GLAUCOMA

Glaucoma may present with a prostrating headache associated with nausea and vomiting. The headache often starts with severe eye pain. On physical examination, the eye is often red with a fixed, moderately dilated pupil.

Glaucoma is discussed in Chap. 39.

PRIMARY HEADACHE DISORDERS

Primary headaches are disorders in which headache and associated features occur in the absence of any exogenous cause. The most common are migraine, tension-type headache, and the trigeminal autonomic cephalalgias, notably cluster headache. These entities are discussed in detail in Chap. 447.

TABLE 21-3 CLASSIFICATION OF CHRONIC DAILY HEADACHE

Primary		
>4 h Daily	<4 h Daily	Secondary
Chronic migraine ^a	Chronic cluster headache ^b	Posttraumatic Head injury Iatrogenic Postinfectious
Chronic tension-type headache ^a	Chronic paroxysmal hemicrania	Inflammatory, such as Giant cell arteritis Sarcoidosis Behçet's syndrome
Hemicrania continua ^a	SUNCT/SUNA	Chronic CNS infection
New daily persistent headache ^a	Hypnic headache	Medication-overuse headache ^a

^aMay be complicated by medication overuse. ^bSome patients may have headache >4 h/d.

Abbreviations: CNS, central nervous system; SUNA, short-lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms; SUNCT, short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing.

CHRONIC DAILY HEADACHE

The broad diagnosis of chronic daily headache (CDH) can be applied when a patient experiences headache on 15 days or more per month. CDH is not a single entity; it encompasses a number of different headache syndromes, both primary and secondary (Table 21-3). In aggregate, this group presents considerable disability and is thus specially dealt with here. Population-based estimates suggest that about 4% of adults have daily or near-daily headache.

APPROACH TO THE PATIENT: Chronic Daily Headache

The first step in the management of patients with CDH is to diagnose any secondary headache and treat that problem (Table 21-3). This can sometimes be a challenge where the underlying cause triggers a worsening of a primary headache. For patients with primary headaches, diagnosis of the headache type will guide therapy. Preventive treatments such as tricyclics, either amitriptyline or nortriptyline at doses up to 1 mg/kg, are very useful in patients with CDH arising from migraine or tension-type headache or where the secondary cause has activated the underlying primary headache. Tricyclics are started in low doses (10–25 mg) daily and may be given 12 h before the expected time of awakening in order to avoid excess morning sleepiness. Anticonvulsants, such as topiramate, valproate, flunarizine (not available in the United States), and candesartan are also useful in migraine.

MANAGEMENT OF MEDICALLY INTRACTABLE DISABLING PRIMARY CHRONIC DAILY HEADACHE

The management of medically intractable headache is difficult. Currently there are a number of promising neuromodulatory approaches, such as occipital nerve stimulation, which appears to modulate thalamic processing in migraine, and has also shown promise in chronic cluster headache, short-lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms (SUNA), short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing (SUNCT), and hemicrania continua (Chap. 447). Single-pulse transcranial magnetic stimulation is in use in Europe and is approved for migraine with aura in the United States. Other modalities are discussed in Chap. 447.

MEDICATION-OVERUSE HEADACHE

Overuse of analgesic medication for headache can aggravate headache frequency, markedly impair the effect of preventive medicines, and induce a state of refractory daily or near-daily headache called *medication-overuse headache*. A proportion of patients who stop

taking analgesics will experience substantial improvement in the severity and frequency of their headache. However, even after cessation of analgesic use, many patients continue to have headache, although they may feel clinically improved in some way, especially if they have been using opioids or barbiturates regularly. The residual symptoms probably represent the underlying primary headache disorder, and most commonly, this issue occurs in patients prone to migraine.

Management of Medication Overuse: Outpatients For patients who overuse medications, it is essential that analgesic use be reduced and eliminated. One approach is to reduce the medication dose by 10% every 1–2 weeks. Immediate cessation of analgesic use is possible for some patients, provided there is no contraindication. Both approaches are facilitated by the use of a medication diary maintained during the month or two before cessation; this helps to identify the scope of the problem. A small dose of a nonsteroidal anti-inflammatory drug (NSAID) such as naproxen, 500 mg bid, if tolerated, will help relieve residual pain as analgesic use is reduced. NSAID overuse is not usually a problem for patients with daily headache when a NSAID with a longer half-life is taken once or twice daily; however, overuse problems may develop with more frequent dosing schedules or shorter acting NSAIDs. Once the patient has substantially reduced analgesic use, a preventive medication should be introduced. It must be emphasized that *preventives generally do not work in the presence of analgesic overuse*. The most common cause of unresponsiveness to treatment is the use of a preventive when analgesics continue to be used regularly. For some patients, discontinuing analgesics is very difficult; often the best approach is to directly inform the patient that some degree of pain is inevitable during this initial period.

Management of Medication Overuse: Inpatients Some patients will require hospitalization for detoxification. Such patients have typically failed efforts at outpatient withdrawal or have a significant medical condition, such as diabetes mellitus, which would complicate withdrawal as an outpatient. Following admission to the hospital, acute medications are withdrawn completely on the first day, in the absence of a contraindication. Antiemetics and fluids are administered as required; clonidine is used for opioid withdrawal symptoms. For acute intolerable pain during the waking hours, aspirin, 1 g IV (not approved in United States), is useful. IM chlorpromazine can be helpful at night; patients must be adequately hydrated. Three to 5 days into the admission, as the effect of the withdrawn substance wears off, a course of IV dihydroergotamine (DHE) can be used. DHE, administered every 8 h for 5 consecutive days, can induce a significant remission that allows a preventive treatment to be established. 5-HT₃ antagonists, such as ondansetron or granisetron, or the neurokinin receptor antagonist, aprepitant, may be required with DHE to prevent significant nausea, and domperidone (not approved in the United States) orally or by suppository can be very helpful. Avoiding sedating or otherwise side effect prone antiemetics is helpful.

NEW DAILY PERSISTENT HEADACHE

New daily persistent headache (NDPH) is a clinically distinct syndrome; its causes are listed in Table 21-4.

TABLE 21-4 DIFFERENTIAL DIAGNOSIS OF NEW DAILY PERSISTENT HEADACHE

Primary	Secondary
Migrainous-type	Subarachnoid hemorrhage
Featureless (tension-type)	Low cerebrospinal fluid (CSF) volume headache
	Raised CSF pressure headache
	Posttraumatic headache ^a
	Chronic meningitis

^aIncludes postinfectious forms.

Clinical Presentation The patient with NDPH presents with headache on most if not all days, and the patient can clearly, and often vividly, recall the moment of onset. The headache usually begins abruptly, but onset may be more gradual; evolution over 3 days has been proposed as the upper limit for this syndrome. Patients typically recall the exact day and circumstances of the onset of headache; the new, persistent head pain does not remit. The first priority is to distinguish between a primary and a secondary cause of this syndrome. Subarachnoid hemorrhage is the most serious of the secondary causes and must be excluded either by history or appropriate investigation (**Chap. 330**).

Secondary NDPH • LOW CSF VOLUME HEADACHE In these syndromes, head pain is positional: it begins when the patient sits or stands upright and resolves upon reclining. The pain, which is occipito-frontal, is usually a dull ache but may be throbbing. Patients with chronic low CSF volume headache typically present with a history of headache from one day to the next that is generally not present on waking but worsens during the day. Recumbency usually improves the headache within minutes, and it can take only minutes to an hour for the pain to return when the patient resumes an upright position.

The most common cause of headache due to persistent low CSF volume is CSF leak following LP. Post-LP headache usually begins within 48 h but may be delayed for up to 12 days. Its incidence is between 10 and 30%. Beverages with caffeine may provide temporary relief. Besides LP, index events may include epidural injection or a vigorous Valsalva maneuver, such as from lifting, straining, coughing, clearing the eustachian tubes in an airplane, or multiple orgasms. Spontaneous CSF leaks are well recognized, and the diagnosis should be considered whenever the headache history is typical, even when there is no obvious index event. As time passes from the index event, the postural nature may become less apparent; cases in which the index event occurred several years before the eventual diagnosis have been recognized. Symptoms appear to result from low volume rather than low pressure: although low CSF pressures, typically 0–50 mmH₂O, are usually identified, a pressure as high as 140 mmH₂O has been noted with a documented leak.

Postural orthostatic tachycardia syndrome (POTS; **Chap. 454**) can present with orthostatic headache similar to low CSF volume headache and is a diagnosis that needs consideration in this setting.

When imaging is indicated to identify the source of a presumed leak, an MRI with gadolinium is the initial study of choice (**Fig. 21-1**). A striking pattern of diffuse meningeal enhancement is so typical that in the appropriate clinical context the diagnosis is established. Chiari malformations may sometimes be noted on MRI; in such cases, surgery to decompress the posterior fossa usually worsens the headache. Spinal MRI with T2 weighting may reveal a leak, and spinal MRI may demonstrate spinal meningeal cysts whose role in these syndromes is yet to be elucidated. The source of CSF leakage may be identified by spinal MRI with appropriate sequences, by CT, or increasingly by MR myelography. Less used now, ¹¹¹In-DTPA CSF studies in the absence of a directly identified site of leakage, may demonstrate early emptying of ¹¹¹In-DTPA tracer into the bladder or slow progress of tracer across the brain suggesting a CSF leak.

Initial treatment for low CSF volume headache is bed rest. For patients with persistent pain, IV caffeine (500 mg in 500 mL of saline administered over 2 h) can be very effective. An electrocardiogram (ECG) to screen for arrhythmia should be performed before administration. It is reasonable to administer at least two infusions of caffeine before embarking on additional tests to identify the source of the CSF leak. Because IV caffeine is safe and can be curative, it spares many patients the need for further investigations. If unsuccessful, an abdominal binder may be helpful. If a leak can be identified, an autologous blood patch is usually curative. A blood patch is also effective for post-LP headache; in this setting, the location is empirically determined to be the site of the LP. In patients with intractable pain, oral theophylline is a useful alternative; however, its effect is less rapid than caffeine.

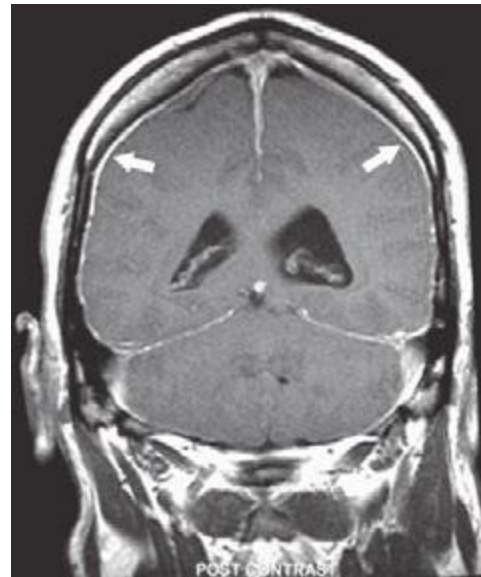


FIGURE 21-1 Magnetic resonance image showing diffuse meningeal enhancement after gadolinium administration in a patient with low cerebrospinal fluid (CSF) volume headache.

RAISED CSF PRESSURE HEADACHE Raised CSF pressure is well recognized as a cause of headache. Brain imaging can often reveal the cause, such as a space-occupying lesion. NDPH due to raised CSF pressure can be the presenting symptom for patients with idiopathic intracranial hypertension (pseudotumor cerebri) without visual problems, particularly when the fundi are normal. Persistently raised intracranial pressure can trigger chronic migraine. These patients typically present with a history of generalized headache that is present on waking and improves as the day goes on. It is generally worse with recumbency. Visual obscurations are frequent. The diagnosis is relatively straightforward when papilledema is present, but the possibility must be considered even in patients without fundoscopic changes. Formal visual field testing should be performed even in the absence of overt ophthalmic involvement. Headache on rising in the morning or nocturnal headache is also characteristic of obstructive sleep apnea or poorly controlled hypertension.

Evaluation of patients suspected to have raised CSF pressure requires brain imaging. It is most efficient to obtain an MRI, including an MR venogram, as the initial study. If there are no contraindications, the CSF pressure should be measured by LP; this should be done when the patient is symptomatic so that both the pressure and the response to removal of 20–30 mL of CSF can be determined. An elevated opening pressure and improvement in headache following removal of CSF are diagnostic.

Initial treatment is with acetazolamide (250–500 mg bid); the headache may improve within weeks. If ineffective, topiramate is the next treatment of choice; it has many actions that may be useful in this setting, including carbonic anhydrase inhibition, weight loss, and neuronal membrane stabilization, likely mediated via effects on phosphorylation pathways. Severely disabled patients who do not respond to medical treatment require intracranial pressure monitoring and may require shunting.

POSTTRAUMATIC HEADACHE A traumatic event can trigger a headache process that lasts for many months or years after the event. The term *trauma* is used in a very broad sense: headache can develop following an injury to the head, but it can also develop after an infectious episode, typically viral meningitis, a fluke illness, or a parasitic infection. Complaints of dizziness, vertigo, and impaired memory can accompany the headache. Symptoms may remit after several weeks or persist for months and even years after the injury. Typically the neurologic examination is normal and CT or MRI studies are unrevealing. Chronic subdural hematoma may on occasion mimic this disorder. Posttraumatic headache may also be seen

after carotid dissection and subarachnoid hemorrhage and after intracranial surgery. The underlying theme appears to be that a traumatic event involving the pain-producing meninges can trigger a headache process that lasts for many years.

OTHER CAUSES In one series, one-third of patients with NDPH reported headache beginning after a transient flulike illness characterized by fever, neck stiffness, photophobia, and marked malaise. Evaluation typically reveals no apparent cause for the headache. There is no convincing evidence that persistent Epstein-Barr virus infection plays a role in NDPH. A complicating factor is that many patients undergo LP during the acute illness; iatrogenic low CSF volume headache must be considered in these cases.

TREATMENT Treatment is largely empirical. Tricyclic antidepressants, notably amitriptyline, and anticonvulsants, such as topiramate, valproate, and gabapentin, have been used with reported benefit. The monoamine oxidase inhibitor phenelzine may also be useful in carefully selected patients. The headache usually resolves within 3–5 years, but it can be quite disabling.

PRIMARY CARE AND HEADACHE MANAGEMENT

Most patients with headache will be seen first in a primary care setting. The task of the primary care physician is to identify the very few worrisome secondary headaches from the very great majority of primary and less troublesome secondary headaches (Table 21-2).

Absent any warning signs, a reasonable approach is to treat when a diagnosis is established. As a general rule, the investigation should focus on identifying worrisome causes of headache or on gaining confidence if no primary headache diagnosis can be made.

After treatment has been initiated, follow-up care is essential to identify whether progress has been made against the headache complaint. Not all headaches will respond to treatment, but, in general, worrisome headaches will progress and will be easier to identify.

When a primary care physician feels the diagnosis is a primary headache disorder, it is worth noting that more than 90% of patients who present to primary care with a complaint of headache will have migraine (Chap. 447).

In general, patients who do not have a clear diagnosis, have a primary headache disorder other than migraine or tension-type headache, or are unresponsive to two or more standard therapies for the considered headache type should be considered for referral to a specialist. In a practical sense, the threshold for referral is also determined by the experience of the primary care physician in headache medicine and the availability of secondary care options.

central gelatinous nucleus pulposus surrounded by a tough cartilaginous ring, the annulus fibrosus. Disks are responsible for 25% of spinal column length and allow the bony vertebrae to move easily upon each other (Figs. 22-1 and 22-2). Desiccation of the nucleus pulposus and degeneration of the annulus fibrosus increase with age and result in loss of disk height. The disks are largest in the cervical and lumbar regions where movements of the spine are greatest. The anterior spine absorbs the shock of bodily movements such as walking and running and, with the posterior spine, protects the spinal cord and nerve roots in the spinal canal.

The posterior spine consists of the vertebral arches and processes. Each arch consists of paired cylindrical pedicles anteriorly and paired lamina posteriorly. The vertebral arch also gives rise to two transverse processes laterally, one spinous process posteriorly, plus two superior and two inferior articular facets. The apposition of a superior and inferior facet constitutes a *facet joint*. The posterior spine provides an anchor for the attachment of muscles and ligaments. The contraction of muscles attached to the spinous and transverse processes and lamina works like a system of pulleys and levers that results in flexion, extension, and lateral bending movements of the spine.

Nerve root injury (*radiculopathy*) is a common cause of neck, arm, low back, buttock, and leg pain (see Figs. 31-2 and 31-3). The nerve roots exit at a level above their respective vertebral bodies in the cervical region (e.g., the C7 nerve root exits at the C6-C7 level) and below their respective vertebral bodies in the thoracic and lumbar regions (e.g., the T1 nerve root exits at the T1-T2 level). The cervical nerve roots follow a short intraspinal course before exiting. By contrast, because the spinal cord ends at the vertebral L1 or L2 level, the lumbar nerve roots follow a long intraspinal course and can be injured anywhere from the upper lumbar spine to their exit at the intervertebral foramen. For example, disk herniation at the L4-L5 level can produce not only L5 root compression, but also compression of the traversing S1 nerve root (Fig. 22-3). The lumbar nerve roots are mobile in the spinal canal, but eventually pass through the narrow *lateral recess* of the spinal canal and *intervertebral foramen* (Figs. 22-2 and 22-3). Neuroimaging of the spine must include both sagittal and axial views to assess possible compression in either the lateral recess or intervertebral foramen.

Pain-sensitive structures of the spine include the periosteum of the vertebrae, dura, facet joints, annulus fibrosus of the intervertebral disk, epidural veins and arteries, and the longitudinal ligaments. Disease of these diverse structures may explain many cases of back pain without nerve root compression. Under normal circumstances, the nucleus pulposus of the intervertebral disk is not pain sensitive.

APPROACH TO THE PATIENT:

Back Pain

TYPES OF BACK PAIN

Delineating the type of pain reported by the patient is the essential first step. Attention is also focused on identification of risk factors for a serious underlying etiology. The most frequent causes of back pain are radiculopathy, fracture, tumor, infection, or referred pain from visceral structures (Table 22-1).

Local pain is caused by injury to pain-sensitive structures that compress or irritate sensory nerve endings. The site of the pain is near the affected part of the back.

Pain referred to the back may arise from abdominal or pelvic viscera. The pain is usually described as primarily abdominal or pelvic, accompanied by back pain and usually unaffected by posture. The patient may occasionally complain of back pain only.

Pain of spine origin may be located in the back or referred to the buttocks or legs. Diseases affecting the upper lumbar spine tend to refer pain to the lumbar region, groin, or anterior thighs. Diseases affecting the lower lumbar spine tend to produce pain referred to the buttocks, posterior thighs, calves, or feet. Referred pain can explain pain syndromes that cross multiple dermatomes without evidence of nerve root compression.

22

Back and Neck Pain

John W. Engstrom, Richard A. Deyo

The importance of back and neck pain in our society is underscored by the following: (1) the cost of back pain in the United States exceeds \$100 billion annually; approximately one-third of these costs are direct health care expenses, and two-thirds are indirect costs resulting from loss of wages and productivity; (2) back symptoms are the most common cause of disability in those <45 years; (3) low back pain is the second most common reason for visiting a physician in the United States; and (4) 70% of persons will have back pain at some point in their lives.

ANATOMY OF THE SPINE

The anterior spine consists of cylindrical vertebral bodies separated by intervertebral disks and held together by the anterior and posterior longitudinal ligaments. The intervertebral disks are composed of a

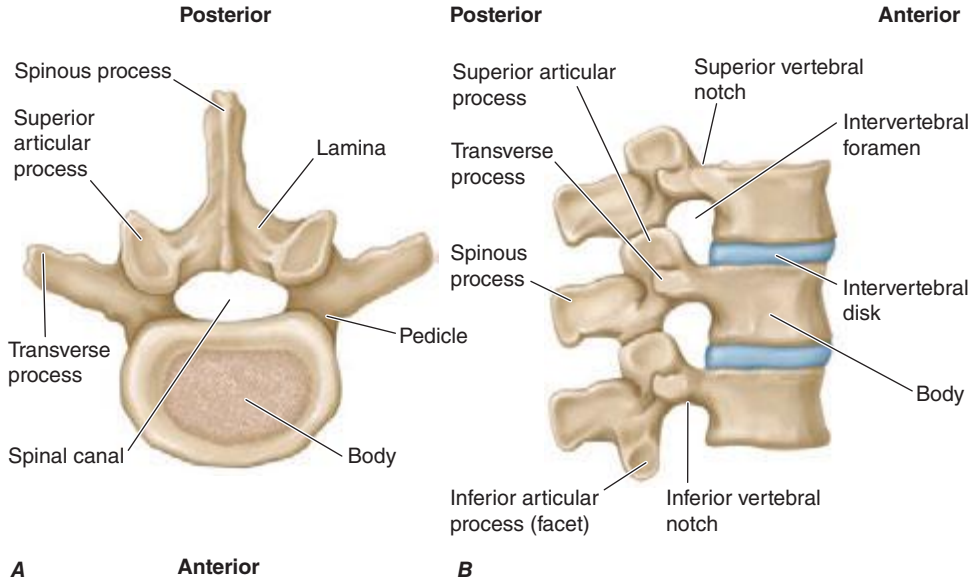


FIGURE 22-1 Vertebral anatomy. (From A Gauthier Cornuelle, DH Gronefeld: *Radiographic Anatomy Positioning*. New York, McGraw-Hill, 1998; with permission.)

Radicular pain is typically sharp and radiates from the low back to a leg within the territory of a nerve root (see “Lumbar Disk Disease,” below). Coughing, sneezing, or voluntary contraction of abdominal muscles (lifting heavy objects or straining at stool) may elicit the radiating pain. The pain may increase in postures that stretch the nerves and nerve roots. Sitting with the leg outstretched places traction on the sciatic nerve and L5 and S1 roots because the nerve passes posterior to the hip. The femoral nerve (L2, L3, and L4 roots) passes anterior to the hip and is not stretched by sitting.

The description of the pain alone often fails to distinguish between referred pain and radiculopathy, although a burning or electric quality favors radiculopathy.

Pain associated with muscle spasm, although of obscure origin, is commonly associated with many spine disorders. The spasms are accompanied by abnormal posture, tense paraspinal muscles, and dull or achy pain in the paraspinal region.

Knowledge of the circumstances associated with the onset of back pain is important when weighing possible serious underlying causes for the pain. Some patients involved in accidents or work-related injuries may exaggerate their pain for the purpose of compensation or for psychological reasons.

EXAMINATION OF THE BACK

A physical examination that includes the abdomen and rectum is advisable. Back pain referred from visceral organs may be reproduced during palpation of the abdomen (pancreatitis, abdominal aortic aneurysm [AAA]) or percussion over the costovertebral angles (pyelonephritis).

The normal spine has a cervical and lumbar lordosis and a thoracic kyphosis. Exaggeration of these normal alignments may result in hyperkyphosis of the thoracic spine or hyperlordosis of the lumbar spine. Inspection may reveal a lateral curvature of the spine (scoliosis). An asymmetry in the prominence of the paraspinal muscles suggests muscle spasm. Spine pain reproduced by palpation over the spinous process reflects injury of the affected vertebrae or adjacent pain-sensitive structures.

Forward bending is often limited by paraspinal muscle spasm; the latter may flatten the usual lumbar lordosis. Flexion at the hips is normal in patients with lumbar spine disease, but flexion of the lumbar spine is limited and sometimes painful. Lateral bending to the side opposite the injured spinal element may stretch the damaged tissues, worsen pain, and limit motion. Hyperextension of the spine (with the patient prone or standing) is limited when nerve root compression, facet joint pathology, or other bony spine disease is present.

Pain from hip disease may mimic the pain of lumbar spine disease. Hip pain can be reproduced by internal and external rotation at the hip with the knee and hip in flexion or by compressing the heel with the examiner’s palm while the leg is extended (heel percussion sign).

The *straight leg-raising (SLR)* maneuver is a simple bedside test for nerve root disease. With the patient supine, passive flexion of the extended leg at the hip stretches the L5 and S1 nerve roots and

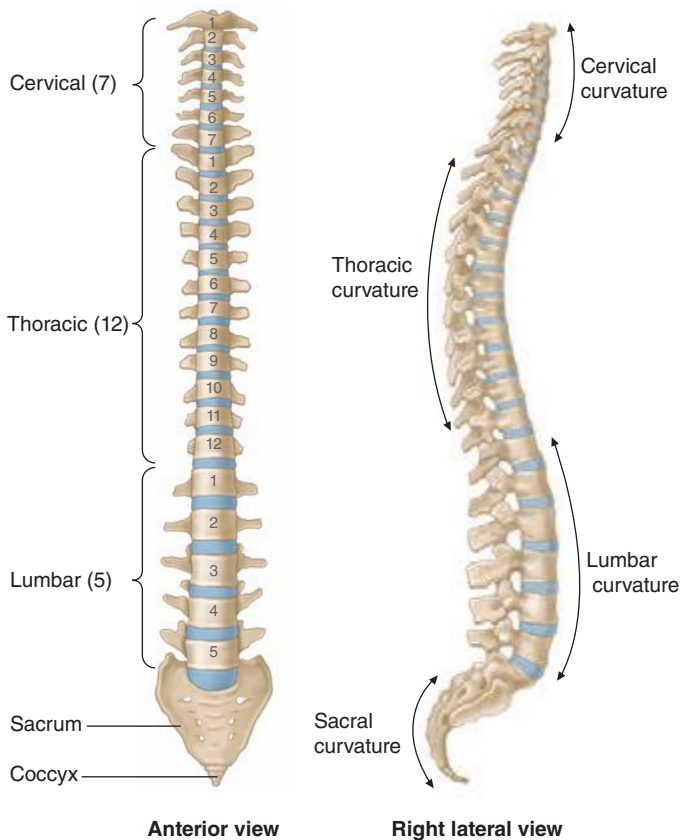


FIGURE 22-2 Spinal column. (From A Gauthier Cornuelle, DH Gronefeld: *Radiographic Anatomy Positioning*. New York, McGraw-Hill, 1998; with permission.)

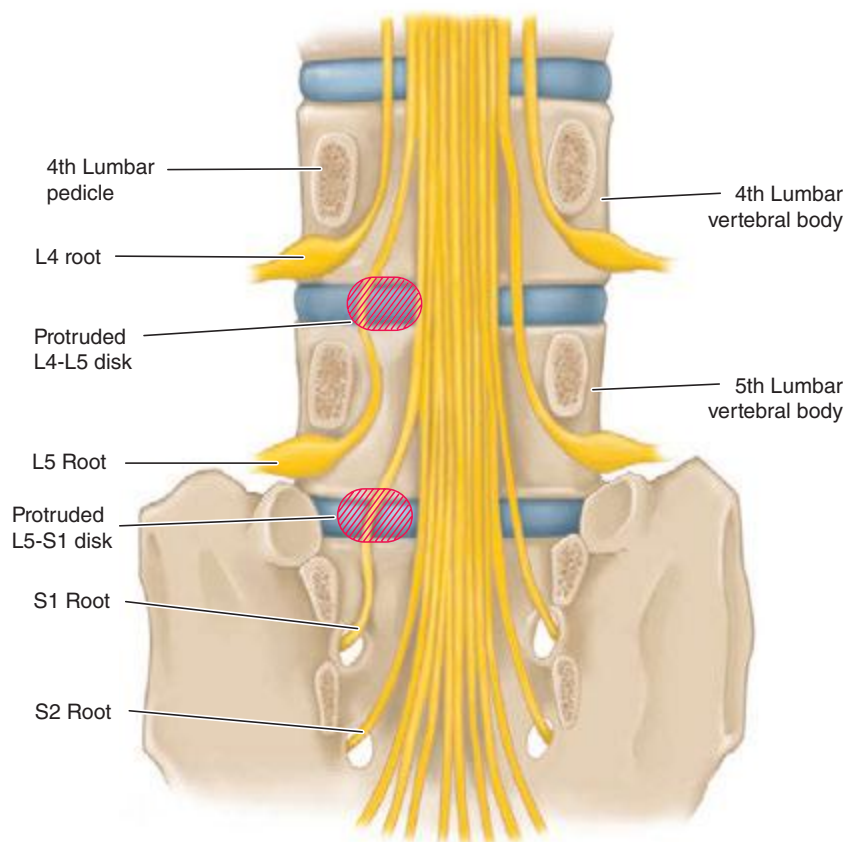


FIGURE 22-3 Compression of L5 and S1 roots by herniated disks. (From AH Ropper, MA Samuels: *Adams and Victor's Principles of Neurology*, 9th ed. New York, McGraw-Hill, 2009; with permission.)

the sciatic nerve. Passive dorsiflexion of the foot during the maneuver adds to the stretch. In healthy individuals, flexion to at least 80° is normally possible without causing pain, although a tight, stretching sensation in the hamstring muscles is common. The SLR test is positive if the maneuver reproduces the patient's usual back or limb pain. Eliciting the SLR sign in both the supine and sitting positions can help determine if the finding is reproducible. The patient may describe pain in the low back, buttocks, posterior thigh, or lower leg, but the *key feature is reproduction of the patient's usual pain*.

TABLE 22-1 ACUTE LOW BACK PAIN: RISK FACTORS FOR AN IMPORTANT STRUCTURAL CAUSE

History

Pain worse at rest or at night
 Prior history of cancer
 History of chronic infection (especially lung, urinary tract, skin)
 History of trauma
 Incontinence
 Age >70 years
 Intravenous drug use
 Glucocorticoid use
 History of a rapidly progressive neurologic deficit

Examination

Unexplained fever
 Unexplained weight loss
 Percussion tenderness over the spine
 Abdominal, rectal, or pelvic mass
 Internal/external rotation of the leg at the hip; heel percussion sign
 Straight leg– or reverse straight leg–raising signs
 Progressive focal neurologic deficit

The *crossed SLR sign* is present when flexion of one leg reproduces the usual pain in the opposite leg or buttocks. In disk herniation, the crossed SLR sign is less sensitive but more specific than the SLR sign. The *reverse SLR sign* is elicited by standing the patient next to the examination table and passively extending each leg with the knee fully extended. This maneuver, which stretches the L2-L4 nerve roots, lumbosacral plexus, and femoral nerve, is considered positive if the patient's usual back or limb pain is reproduced. For all of these tests, the nerve or nerve root lesion is always on the side of the pain.

The neurologic examination includes a search for focal weakness or muscle atrophy, focal reflex changes, diminished sensation in the legs, or signs of spinal cord injury. The examiner should be alert to the possibility of breakaway weakness, defined as fluctuations in the maximum power generated during muscle testing. Breakaway weakness may be due to pain or a combination of pain and an underlying true weakness. Breakaway weakness without pain is almost always due to a lack of effort. In uncertain cases, electromyography (EMG) can determine if true weakness due to nerve tissue injury is present. Findings with specific lumbosacral nerve root lesions are shown in [Table 22-2](#) and are discussed below.

LABORATORY, IMAGING, AND EMG STUDIES

Laboratory studies are rarely needed for the initial evaluation of nonspecific acute (<3 months in duration) low back pain (ALBP). Risk factors for a serious underlying cause and for infection, tumor, or fracture, in particular, should be sought by history and exam. If risk factors are present ([Table 22-1](#)), then laboratory studies (complete blood count [CBC], erythrocyte sedimentation rate [ESR], urinalysis) are indicated. If risk factors are absent, then management is conservative (see “Treatment,” below)

Computed tomography (CT) scanning is superior to routine x-rays for the detection of fractures involving posterior spine structures, craniocervical and cervicothoracic junctions, C1 and C2

TABLE 22-2 LUMBOSACRAL RADICULOPATHY: NEUROLOGIC FEATURES

Lumbosacral Nerve Roots	Examination Findings			
	Reflex	Sensory	Motor	Pain Distribution
L2 ^a	—	Upper anterior thigh	Psoas (hip flexors)	Anterior thigh
L3 ^a	—	Lower anterior thigh Anterior knee	Psoas (hip flexors) Quadriceps (knee extensors) Thigh adductors	Anterior thigh, knee
L4 ^a	Quadriceps (knee)	Medial calf	Quadriceps (knee extensors) ^b Thigh adductors	Knee, medial calf Anterolateral thigh
L5 ^c	—	Dorsal surface—foot Lateral calf	Peronei (foot evertors) ^b Tibialis anterior (foot dorsiflexors) Gluteus medius (hip abductors) Toe dorsiflexors	Lateral calf, dorsal foot, posterolateral thigh, buttocks
S1 ^c	Gastrocnemius/soleus (ankle)	Plantar surface—foot Lateral aspect—foot	Gastrocnemius/soleus (foot plantar flexors) ^b Abductor hallucis (toe flexors) ^b Gluteus maximus (hip extensors)	Bottom foot, posterior calf, posterior thigh, buttocks

^aReverse straight leg-raising sign present—see “Examination of the Back.” ^bThese muscles receive the majority of innervation from this root. ^cStraight leg-raising sign present—see “Examination of the Back.”

vertebrae, bone fragments within the spinal canal, or misalignment. CT scans are increasingly used as a primary screening modality for moderate to severe acute trauma. Magnetic resonance imaging (MRI) or CT myelography is the radiologic test of choice for evaluation of most serious diseases involving the spine. MRI is superior for the definition of soft tissue structures, whereas CT myelography provides optimal imaging of the lateral recess of the spinal canal and is better tolerated by claustrophobic patients.

Annual population surveys in the United States suggest that patients with back pain have reported progressively worse functional limitations in recent years, rather than progressive improvements, despite rapid increases in spine imaging, opioid prescribing, injections, and spine surgery. This suggests that more selective use of diagnostic and treatment modalities may be appropriate.

Spine imaging often reveals abnormalities of dubious clinical relevance that may alarm clinicians and patients alike and prompt further testing and unnecessary therapy. Both randomized trials and observational studies have suggested such a “cascade effect” of imaging may create a gateway to other unnecessary care. Based in part on such evidence, the American College of Physicians has made parsimonious spine imaging a high priority in its “Choosing Wisely” campaign, aimed at reducing unnecessary care. Successful efforts to reduce unnecessary imaging have typically been multifaceted. Some include physician education by clinical leaders and computerized decision support, to identify any recent relevant imaging tests and require approved indications for ordering an imaging test. Other strategies have included audit and feedback regarding individual rates of ordering and indications, and more rapid access to physical therapy or consultation for patients without imaging indications.

When imaging tests are reported, it may be useful to indicate that certain degenerative findings are common in normal, pain-free individuals. In an observational study, this strategy was associated with lower rates of repeat imaging, opioid therapy, and physical therapy referral.

Electrodiagnostic studies can be used to assess the functional integrity of the peripheral nervous system (**Chap. 442e**). Sensory nerve conduction studies are normal when focal sensory loss confirmed by examination is due to nerve root damage because the nerve roots are proximal to the nerve cell bodies in the dorsal root ganglia. Injury to nerve tissue distal to the dorsal root ganglion (e.g., plexus or peripheral nerve) results in reduced sensory nerve signals. Needle EMG complements nerve conduction studies by detecting denervation or reinnervation changes in a myotomal (segmental)

distribution. Multiple muscles supplied by different nerve roots and nerves are sampled; the pattern of muscle involvement indicates the nerve root(s) responsible for the injury. Needle EMG provides objective information about motor nerve fiber injury when clinical evaluation of weakness is limited by pain or poor effort. EMG and nerve conduction studies will be normal when sensory nerve root injury or irritation is the pain source.

CAUSES OF BACK PAIN

(Table 22-3)

LUMBAR DISK DISEASE

This is a common cause of acute, chronic, or recurrent low back and leg pain (**Figs. 22-3 and 22-4**). Disk disease is most likely to occur at the L4-L5 or L5-S1 levels, but upper lumbar levels are involved occasionally. The cause is often unknown, but the risk is increased in overweight individuals. Disk herniation is unusual prior to age 20 years and is rare in the fibrotic disks of the elderly. Complex genetic factors may play a role in predisposing some patients to disk disease. The pain may be located in the low back only or referred to a leg, buttock, or hip. A sneeze, cough, or trivial movement may cause the nucleus pulposus to prolapse, pushing the frayed and weakened annulus posteriorly. With severe disk disease, the nucleus may protrude through the annulus (herniation) or become extruded to lie as a free fragment in the spinal canal.

The mechanism by which intervertebral disk injury causes back pain is controversial. The inner annulus fibrosus and nucleus pulposus are normally devoid of innervation. Inflammation and production of proinflammatory cytokines within a ruptured nucleus pulposus may trigger or perpetuate back pain. Ingrowth of nociceptive (pain) nerve fibers into inner portions of a diseased disk may be responsible for some chronic “diskogenic” pain. Nerve root injury (radiculopathy) from disk herniation is usually due to inflammation, but lateral herniation may produce compression in the lateral recess or at the intervertebral foramen.

A ruptured disk may be asymptomatic or cause back pain, abnormal posture, limitation of spine motion (particularly flexion), a focal neurologic deficit, or radicular pain. A dermatomal pattern of sensory loss or a reduced or absent deep tendon reflex is more suggestive of a specific root lesion than is the pattern of pain. Motor findings (focal weakness, muscle atrophy, or fasciculations) occur less frequently than focal sensory or reflex changes. Symptoms and signs are usually unilateral, but bilateral involvement does occur with large central disk herniations that compress multiple roots or cause inflammation of

TABLE 22-3 CAUSES OF BACK OR NECK PAIN

Lumbar Disk Disease
Degenerative Spine Disease
Lumbar spinal stenosis without or with neurogenic claudication
Intervertebral foraminal or lateral recess narrowing
Disk-osteophyte complex
Facet or uncovertebral joint hypertrophy
Lateral disk protrusion
Spondylosis (osteoarthritis) and spondylolisthesis
Spine Infection
Vertebral osteomyelitis
Spinal epidural abscess
Septic disk (diskitis)
Meningitis
Lumbar arachnoiditis
Neoplasms—Metastatic, Hematologic, Primary Bone Tumors
Fractures
Trauma/falls, motor vehicle accidents
Atraumatic fractures: osteoporosis, neoplastic infiltration, osteomyelitis
Minor Trauma
Strain or sprain
Whiplash injury
Metabolic Spine Disease
Osteoporosis—hyperparathyroidism, immobility
Osteosclerosis (e.g., Paget's disease)
Congenital/Developmental
Spondylolysis
Kyphoscoliosis
Spina bifida occulta
Tethered spinal cord
Autoimmune Inflammatory Arthritis
Other Causes of Back Pain
Referred pain from visceral disease (e.g., abdominal aortic aneurysm)
Postural
Psychiatric, malingering, chronic pain syndromes

nerve roots within the spinal canal. Clinical manifestations of specific nerve root lesions are summarized in Table 22-2.

The differential diagnosis covers a variety of serious and treatable conditions, including epidural abscess, hematoma, fracture, or tumor.

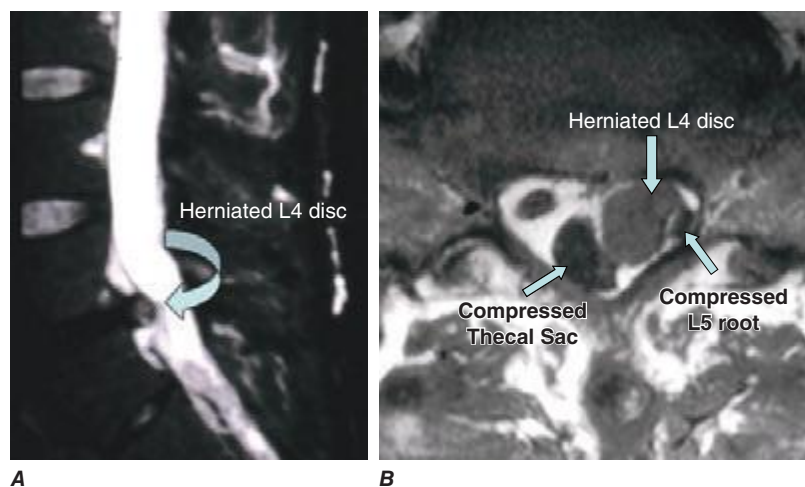


FIGURE 22-4 Left L5 radiculopathy. **A.** Sagittal T2-weighted image on the left reveals disk herniation at the L4-L5 level. **B.** Axial T1-weighted image shows paracentral disk herniation with displacement of the thecal sac medially and the left L5 nerve root posteriorly in the left lateral recess.

Fever, constant pain uninfluenced by position, sphincter abnormalities, or signs of spinal cord disease suggest an etiology other than lumbar disk disease. Absence of ankle reflexes can be a normal finding in persons older than age 60 years or a sign of bilateral S1 radiculopathy. An absent deep tendon reflex or focal sensory loss may indicate injury to a nerve root, but other sites of injury along the nerve must also be considered. For example, an absent knee reflex may be due to a femoral neuropathy or an L4 nerve root injury. A loss of sensation over the foot and lateral lower calf may result from a peroneal or lateral sciatic neuropathy or an L5 nerve root injury. Focal muscle atrophy may reflect injury to the anterior horn cells of the spinal cord, a nerve root, peripheral nerve, or disuse.

A lumbar spine MRI scan or CT myelogram is necessary to establish the location and type of pathology. Spine MRIs yield exquisite views of intraspinal and adjacent soft tissue anatomy. Bony lesions of the lateral recess or intervertebral foramen are optimally visualized by CT myelography. The correlation of neuroradiologic findings to symptoms, particularly pain, is not simple. Contrast-enhancing tears in the annulus fibrosus or disk protrusions are widely accepted as common sources of back pain; however, studies have found that many asymptomatic adults have similar findings. Asymptomatic disk protrusions are also common and may enhance with contrast. Furthermore, in patients with known disk herniation treated either medically or surgically, persistence of the herniation 10 years later had no relationship to the clinical outcome. In summary, MRI findings of disk protrusion, tears in the annulus fibrosus, or hypertrophic facet joints are common incidental findings that, by themselves, should not dictate management decisions for patients with back pain.

The diagnosis of nerve root injury is most secure when the history, examination, results of imaging studies, and the EMG are concordant. The correlation between CT and EMG for localization of nerve root injury is between 65 and 73%. Up to one-third of asymptomatic adults have a lumbar disk protrusion detected by CT or MRI scans.

Management of lumbar disk disease is discussed below.

Cauda equina syndrome (CES) signifies an injury of multiple lumbosacral nerve roots within the spinal canal distal to the termination of the spinal cord at L1-L2. Low back pain, weakness and areflexia in the legs, saddle anesthesia, or loss of bladder function may occur. The problem must be distinguished from disorders of the lower spinal cord (conus medullaris syndrome), acute transverse myelitis (Chap. 456), and Guillain-Barré syndrome (Chap. 460). Combined involvement of the conus medullaris and cauda equina can occur. CES is commonly due to a ruptured lumbosacral intervertebral disk, lumbosacral spine fracture, hematoma within the spinal canal (e.g., following lumbar puncture in patients with coagulopathy), compressive tumor, or other mass lesion. Treatment options include surgical decompression, sometimes urgently in an attempt to restore or preserve motor or sphincter function, or radiotherapy for metastatic tumors (Chap. 118).

DEGENERATIVE CONDITIONS

Lumbar spinal stenosis (LSS) describes a narrowed lumbar spinal canal and is frequently asymptomatic. Typical is *neurogenic claudication*, consisting of back and buttock or leg pain induced by walking or standing and relieved by sitting. Symptoms in the legs are usually bilateral. Unlike vascular claudication, symptoms are often provoked by standing without walking. Unlike lumbar disk disease, symptoms are usually relieved by sitting. Patients with neurogenic claudication can often walk much farther when leaning over a shopping cart and can pedal a stationary bike with ease while sitting. These flexed positions increase the anteroposterior spinal canal diameter and reduce intraspinal venous hypertension, resulting in pain relief. Focal weakness, sensory loss, or reflex changes may occur when spinal stenosis is associated with neural foraminal narrowing and radiculopathy. Severe neurologic deficits, including paralysis and urinary incontinence, occur only rarely.

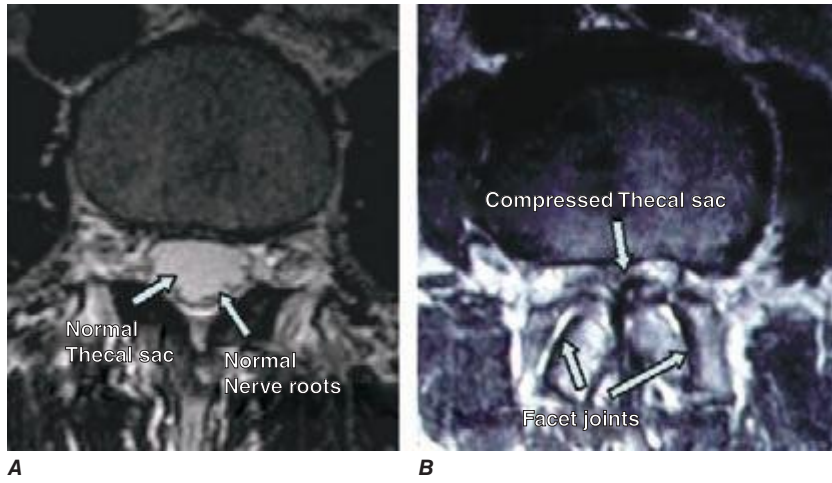


FIGURE 22-5 Axial T2-weighted images of the lumbar spine. **A.** The image shows a normal thecal sac within the lumbar spinal canal. The thecal sac is bright. The lumbar roots are dark punctuate dots in the posterior thecal sac with the patient supine. **B.** The thecal sac is not well visualized due to severe lumbar spinal canal stenosis, partially the result of hypertrophic facet joints.

LSS by itself is frequently asymptomatic, and the correlation between the severity of symptoms and degree of stenosis of the spinal canal is variable. LSS can be acquired (75%), congenital, or both. Congenital forms (achondroplasia, idiopathic) are characterized by short, thick pedicles that produce both spinal canal and lateral recess stenosis. Acquired factors that contribute to spinal stenosis include degenerative diseases (spondylosis, spondylolisthesis, scoliosis), trauma, spine surgery, metabolic or endocrine disorders (epidural lipomatosis, osteoporosis, acromegaly, renal osteodystrophy, hypoparathyroidism), and Paget's disease. MRI provides the best definition of the abnormal anatomy (Fig. 22-5).

Conservative treatment of symptomatic LSS includes nonsteroidal anti-inflammatory drugs (NSAIDs), acetaminophen, exercise programs, and symptomatic treatment of acute pain episodes. There is

insufficient evidence to support the routine use of epidural glucocorticoid injections. Surgical therapy is considered when medical therapy does not relieve symptoms sufficiently to allow for resumption of activities of daily living or when focal neurologic signs are present. Most patients with neurogenic claudication who are treated medically do not improve over time. Surgical management can produce significant relief of back and leg pain within 6 weeks, and pain relief persists for at least 2 years. However, up to one-quarter develop recurrent stenosis at the same spinal level or an adjacent level 7–10 years after the initial surgery; recurrent symptoms usually respond to a second surgical decompression.

Neural foraminal narrowing with radiculopathy is a common consequence of osteoarthritic processes that cause lumbar spinal stenosis (Figs. 22-1 and 22-6), including osteophytes, lateral disk protrusion, calcified disk-osteophytes, facet joint hypertrophy, uncovertebral joint hypertrophy (cervical spine), congenitally shortened pedicles, or, frequently, a combination of these processes. Neoplasms (primary or metastatic), fractures, infections (epidural abscess), or hematomas are other considerations. These conditions can produce unilateral nerve root symptoms or signs due to compression at the intervertebral foramen or in the lateral recess; symptoms are indistinguishable from disk-related radiculopathy, but treatment may differ depending on the specific etiology. The history and neurologic examination alone cannot distinguish between these possibilities. A spine neuroimaging (CT or MRI) procedure is required to identify the anatomic cause. Neurologic findings from the examination and EMG can help direct the attention of the radiologist to specific nerve roots, especially on axial images. For *facet joint hypertrophy*, surgical foraminotomy produces long-term relief of leg and back pain in 80–90% of patients. The usefulness of therapeutic facet joint blocks for pain is controversial. Medical causes of lumbar or cervical radiculopathy unrelated to anatomic spine disease include infections

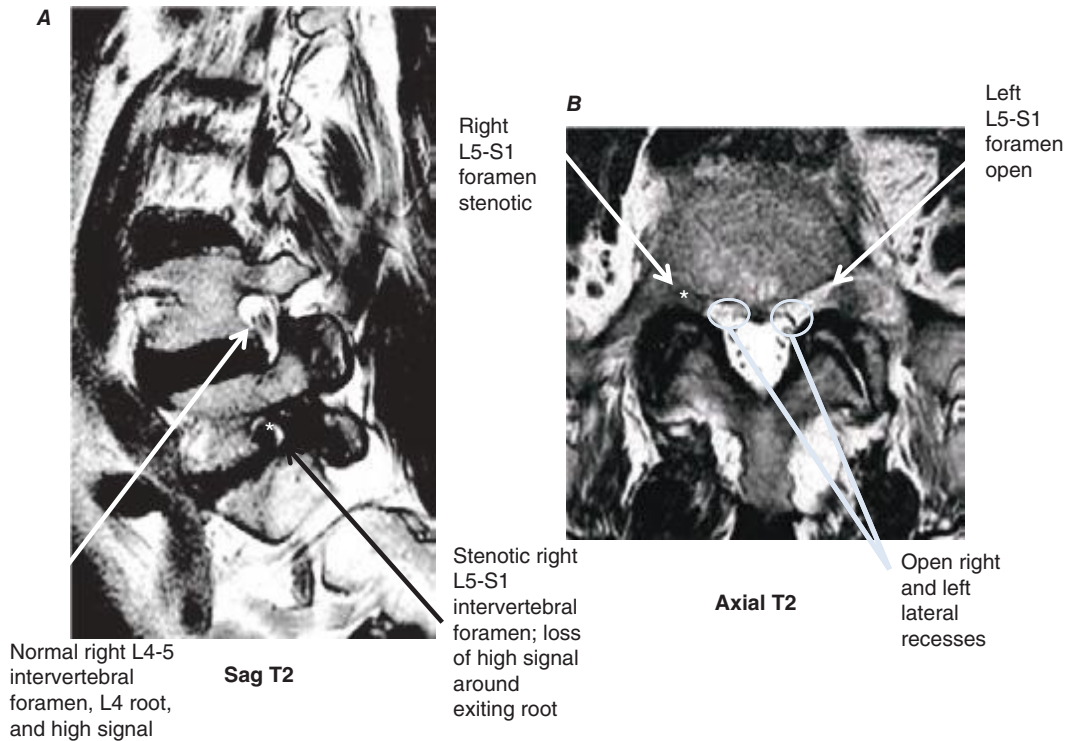


FIGURE 22-6 Right L5 radiculopathy. **A.** Sagittal T2-weighted image. There is normal high signal around the exiting right L4 nerve root in the right neural foramen at L4-L5; effacement of the high signal in the right L5-S1 foramen is present one level caudal on the right at L5-S1. **B.** Axial T2-weighted image. The lateral recesses are normal bilaterally; the intervertebral foramen is normal on the left, but severely stenotic on the right. *Severe right L5-S1 foraminal stenosis.

(e.g., herpes zoster, Lyme disease), carcinomatous meningitis, and root avulsion or traction (severe trauma).

SPONDYLOSIS AND SPONDYLOLISTHESIS

Spondylosis, or osteoarthritic spine disease, typically occurs in later life and primarily involves the cervical and lumbosacral spine. Patients often complain of back pain that increases with movement, is associated with stiffness, and is better when inactive. The relationship between clinical symptoms and radiologic findings is usually not straightforward. Pain may be prominent when x-ray, CT, or MRI findings are minimal, and prominent degenerative spine disease can be seen in asymptomatic patients. Osteophytes or combined disk-osteophytes may cause or contribute to central spinal canal stenosis, lateral recess stenosis, or neural foraminal narrowing.

Spondylolisthesis is the anterior slippage of the vertebral body, pedicles, and superior articular facets, leaving the posterior elements behind. Spondylolisthesis can be associated with spondylolysis, congenital anomalies, degenerative spine disease, or other causes of mechanical weakness of the pars (e.g., infection, osteoporosis, tumor, trauma, prior surgery). The slippage may be asymptomatic or may cause low back pain and hamstring tightness, nerve root injury (the L5 root most frequently), symptomatic spinal stenosis, or CES in severe cases. Tenderness may be elicited near the segment that has “slipped” forward (most often L4 on L5 or occasionally L5 on S1). Focal anterolisthesis or retrolisthesis can occur at any cervical or lumbar level and be the source of neck or low back pain. Plain x-rays with the neck or low back in flexion and extension will reveal the movement at the abnormal spinal segment. Surgery is considered for pain symptoms that do not respond to conservative measures (e.g., rest, physical therapy) and in cases with progressive neurologic deficit, postural deformity, slippage >50%, or scoliosis.

NEOPLASMS

Back pain is the most common neurologic symptom in patients with systemic cancer and is the presenting symptom in 20%. The cause is usually vertebral body metastasis but can also result from spread of cancer through the intervertebral foramen (especially with lymphoma), from carcinomatous meningitis, or from metastasis to the spinal cord. Cancer-related back pain tends to be constant, dull, unrelieved by rest, and worse at night. By contrast, mechanical low back pain usually improves with rest. MRI, CT, and CT myelography are the studies of choice when spinal metastasis is suspected. Once a metastasis is found, imaging of the entire spine reveals additional tumor deposits in one-third of patients. MRI is preferred for soft tissue definition, but the most rapidly available imaging modality is best because the patient's condition may worsen quickly without intervention. Fewer than 5% of patients who are nonambulatory at the time of diagnosis ever regain the ability to walk; thus, early diagnosis is crucial. [The management of spinal metastasis is discussed in detail in Chap. 118.](#)

INFECTIONS/INFLAMMATION

Vertebral osteomyelitis is often caused by staphylococci, but other bacteria or tuberculosis (Pott's disease) may be responsible. The primary source of infection is usually the urinary tract, skin, or lungs. Intravenous drug use is a well-recognized risk factor. Whenever pyogenic osteomyelitis is found, the possibility of bacterial endocarditis should be considered. Back pain unrelieved by rest, spine tenderness over the involved spine segment, and an elevated ESR are the most common findings in vertebral osteomyelitis. Fever or an elevated white blood cell count is found in a minority of patients. MRI and CT are sensitive and specific for early detection of osteomyelitis; CT may be more readily available in emergency settings and better tolerated by some patients with severe back pain. The intervertebral disk can also be affected by infection (diskitis) and, very rarely, by tumor.

Spinal epidural abscess ([Chap. 456](#)) presents with back pain (aggravated by movement or palpation), fever, radiculopathy, or signs of spinal cord compression. The subacute development of two or more of these findings should increase the index of suspicion for spinal epidural abscess. The abscess may track over multiple spinal levels and is best delineated by spine MRI.

Lumbar adhesive arachnoiditis with radiculopathy is due to fibrosis following inflammation within the subarachnoid space. The fibrosis results in nerve root adhesions and presents as back and leg pain associated with focal motor, sensory, or reflex changes. Causes of arachnoiditis include multiple lumbar operations, chronic spinal infections (especially tuberculosis in the developing world), spinal cord injury, intrathecal hemorrhage, myelography (rare), intrathecal injections (glucocorticoids, anesthetics, or other agents), and foreign bodies. The MRI shows clumped nerve roots or loculations of cerebrospinal fluid within the thecal sac. Clumped nerve roots may also occur with demyelinating polyneuropathy or neoplastic infiltration. Treatment is usually unsatisfactory. Microsurgical lysis of adhesions, dorsal rhizotomy, dorsal root ganglionectomy, and epidural glucocorticoids have been tried, but outcomes have been poor. Dorsal column stimulation for pain relief has produced varying results.

TRAUMA

A patient complaining of back pain and an inability to move the legs may have a spine fracture or dislocation; with fractures above L1 the spinal cord is at risk for compression. Care must be taken to avoid further damage to the spinal cord or nerve roots by immobilizing the back or neck pending the results of radiologic studies. Vertebral fractures frequently occur in the absence of trauma in association with osteoporosis, glucocorticoid use, osteomyelitis, or neoplastic infiltration.

Sprains and Strains The terms *low back sprain*, *strain*, and *mechanically induced muscle spasm* refer to minor, self-limited injuries associated with lifting a heavy object, a fall, or a sudden deceleration such as in an automobile accident. These terms are used loosely and do not clearly describe a specific anatomic lesion. The pain is usually confined to the lower back, and there is no radiation to the buttocks or legs. Patients with paraspinal muscle spasm often assume unusual postures.

Traumatic Vertebral Fractures Most traumatic fractures of the lumbar vertebral bodies result from injuries producing anterior wedging or compression. With severe trauma, the patient may sustain a fracture-dislocation or a “burst” fracture involving the vertebral body and posterior elements. Traumatic vertebral fractures are caused by falls from a height, sudden deceleration in an automobile accident, or direct injury. Neurologic impairment is common, and early surgical treatment is indicated. In victims of blunt trauma, CT scans of the chest, abdomen, or pelvis can be reformatted to detect associated vertebral fractures.

METABOLIC CAUSES

Osteoporosis and Osteosclerosis Immobilization, osteomalacia, the postmenopausal state, renal disease, multiple myeloma, hyperparathyroidism, hyperthyroidism, metastatic carcinoma, or glucocorticoid use may accelerate osteoporosis and weaken the vertebral body, leading to compression fractures and pain. Up to two-thirds of compression fractures seen on radiologic imaging are asymptomatic. The most common nontraumatic vertebral body fractures are due to postmenopausal or senile osteoporosis ([Chap. 425](#)). The risk of an additional vertebral fracture at 1 year following a first vertebral fracture is 20%. The presence of fever, weight loss, fracture at a level above T4, or the conditions described above should increase suspicion for a cause other than senile osteoporosis. The sole manifestation of a compression fracture may be localized back or radicular pain exacerbated by movement and often reproduced by palpation over the spinous process of the affected vertebra.

Relief of acute pain can often be achieved with acetaminophen or a combination of opioids and acetaminophen. The role of NSAIDs is controversial. Both pain and disability are improved with bracing. Antiresorptive drugs, especially bisphosphonates (e.g., alendronate), have been shown to reduce the risk of osteoporotic fractures and are the preferred treatment to prevent additional fractures. Less than one-third of patients with prior compression fractures are adequately treated for osteoporosis despite the increased risk for future fractures; even fewer at-risk patients without a history of fracture are adequately treated. Given the negative results of sham-controlled studies of percutaneous vertebroplasty (PVP) and of kyphoplasty for osteoporotic

compression fractures associated with debilitating pain, these procedures are not routinely recommended.

Osteosclerosis, an abnormally increased bone density often due to Paget's disease, is readily identifiable on routine x-ray studies and can sometimes be a source of back pain. It may be associated with an isolated increase in alkaline phosphatase in an otherwise healthy older person. Spinal cord or nerve root compression can result from bony encroachment. The diagnosis of Paget's disease as the cause of a patient's back pain is a diagnosis of exclusion.

For further discussion of these bone disorders, see Chaps. 424, 425, and 426e.

AUTOIMMUNE INFLAMMATORY ARTHRITIS

Autoimmune inflammatory disease of the spine can present with the insidious onset of low back, buttock, or neck pain. Examples include rheumatoid arthritis (**Chap 380**), ankylosing spondylitis, reactive arthritis, psoriatic arthritis, or inflammatory bowel disease (**Chap. 384**).

CONGENITAL ANOMALIES OF THE LUMBAR SPINE

Spondylolysis is a bony defect in the vertebral pars interarticularis (a segment near the junction of the pedicle with the lamina); the cause is usually a stress microfracture in a congenitally abnormal segment. It occurs in up to 6% of adolescents. The defect (usually bilateral) is best visualized on plain x-rays, CT scan, or bone scan and is frequently asymptomatic. Symptoms may occur in the setting of a single injury, repeated minor injuries, or during a growth spurt. Spondylolysis is the most common cause of persistent low back pain in adolescents and is often associated with sports-related activities.

Scoliosis refers to an abnormal curvature in the coronal (lateral) plane of the spine. With *kyphoscoliosis*, there is, in addition, a forward curvature of the spine. The abnormal curvature may be congenital due to abnormal spine development, acquired in adulthood due to degenerative spine disease, or occasionally progressive due to neuromuscular disease. The deformity can progress until ambulation or pulmonary function is compromised.

Spina bifida occulta is a failure of closure of one or several vertebral arches posteriorly; the meninges and spinal cord are normal. A dimple or small lipoma may overlie the defect. Most cases are asymptomatic and discovered incidentally during an evaluation for back pain.

Tethered cord syndrome usually presents as a progressive cauda equina disorder (see below), although myelopathy may also be the initial manifestation. The patient is often a young adult who complains of perineal or perianal pain, sometimes following minor trauma. MRI studies reveal a low-lying conus (below L1 and L2) and a short and thickened filum terminale.

REFERRED PAIN FROM VISCERAL DISEASE

Diseases of the thorax, abdomen, or pelvis may refer pain to the posterior portion of the spinal segment that innervates the diseased organ. Occasionally, back pain may be the first and only manifestation. Upper abdominal diseases generally refer pain to the lower thoracic or upper lumbar region (eighth thoracic to the first and second lumbar vertebrae), lower abdominal diseases to the midlumbar region (second to fourth lumbar vertebrae), and pelvic diseases to the sacral region. Local signs (pain with spine palpation, paraspinal muscle spasm) are absent, and little or no pain accompanies routine movements of the spine.

Low Thoracic or Lumbar Pain with Abdominal Disease Tumors of the posterior wall of the stomach or duodenum typically produce epigastric pain (**Chaps. 109 and 348**), but midline back or paraspinal pain may occur if retroperitoneal extension is present. Fatty foods occasionally induce back pain associated with biliary disease. Diseases of the pancreas can produce right or left paraspinal back pain. Pathology in retroperitoneal structures (hemorrhage, tumors, pyelonephritis) can produce paraspinal pain that radiates to the lower abdomen, groin, or anterior thighs. A mass in the iliopsoas region can produce unilateral lumbar pain with radiation toward the groin, labia, or testicle. The sudden appearance of lumbar pain in a patient receiving anticoagulants suggests retroperitoneal hemorrhage.

Isolated low back pain occurs in some patients with a contained rupture of an abdominal aortic aneurysm (AAA). The classic clinical triad of abdominal pain, shock, and back pain occurs in <20% of patients. The typical patient at risk is an elderly male smoker with back pain. The diagnosis may be missed because the symptoms and signs can be nonspecific. Misdiagnoses include nonspecific back pain, diverticulitis, renal colic, sepsis, and myocardial infarction. A careful abdominal examination revealing a pulsatile mass (present in 50–75% of patients) is an important physical finding. Patients with suspected AAA should be evaluated with abdominal ultrasound, CT, or MRI (**Chap. 301**).

Sacral Pain with Gynecologic and Urologic Disease Pelvic organs rarely cause low back pain, except for gynecologic disorders involving the uterosacral ligaments. The pain is referred to the sacral region. Endometriosis or uterine cancers may invade the uterosacral ligaments. Pain associated with endometriosis is typically premenstrual and often continues until it merges with menstrual pain. Uterine malposition may cause uterosacral ligament traction (retroversion, descensus, and prolapse) or produce sacral pain after prolonged standing.

Menstrual pain may be felt in the sacral region sometimes with poorly localized, cramping pain radiating down the legs. Pain due to neoplastic infiltration of nerves is typically continuous, progressive in severity, and unrelieved by rest at night. Less commonly, radiation therapy of pelvic tumors may produce sacral pain from late radiation necrosis of tissue. Low back pain that radiates into one or both thighs is common in the last weeks of pregnancy.

Urologic sources of lumbosacral back pain include chronic prostatitis, prostate cancer with spinal metastasis (**Chap. 115**), and diseases of the kidney or ureter. Lesions of the bladder and testes do not often produce back pain. Infectious, inflammatory, or neoplastic renal diseases may produce ipsilateral lumbosacral pain, as can renal artery or vein thrombosis. Paraspinal lumbar pain may be a symptom of ureteral obstruction due to nephrolithiasis.

OTHER CAUSES OF BACK PAIN

Postural Back Pain There is a group of patients with nonspecific chronic low back pain (CLBP) in whom no specific anatomic lesion can be found despite exhaustive investigation. These individuals complain of vague, diffuse back pain with prolonged sitting or standing that is relieved by rest. Exercises to strengthen the paraspinal and abdominal muscles are sometimes helpful.

Psychiatric Disease CLBP may be encountered in patients who seek financial compensation; in malingerers; or in those with concurrent substance abuse. Many patients with CLBP have a history of psychiatric illness (depression, anxiety states) or childhood trauma (physical or sexual abuse) that antedates the onset of back pain. Preoperative psychological assessment has been used to exclude patients with marked psychological impairments that predict a poor surgical outcome from spine surgery.

IDIOPATHIC

The cause of low back pain occasionally remains unclear. Some patients have had multiple operations for disk disease but have persistent pain and disability. The original indications for surgery may have been questionable, with back pain only, no definite neurologic signs, or a minor disk bulge noted on CT or MRI. Scoring systems based on neurologic signs, psychological factors, physiologic studies, and imaging studies have been devised to minimize the likelihood of unsuccessful surgery.

TREATMENT BACK PAIN

HEALTH CARE FOR POPULATIONS OF BACK PAIN PATIENTS: A CLINICAL CARE SYSTEMS VIEW

There are increasing pressures to contain health care costs, especially when expensive care is not based on sound evidence. Physicians, patients, the insurance industry, and government providers of

health care will need to work together to ensure cost-effective care for patients with back pain.

Surveys in the United States indicate that patients with back pain have reported progressively worse functional limitations in recent years, despite rapid increases in spine imaging, opioid prescribing, injections, and spine surgery. This suggests that more selective use of diagnostic and treatment modalities may be appropriate.

Spine imaging often reveals abnormalities of dubious clinical relevance that may alarm clinicians and patients and prompt further testing and unnecessary therapy. Both randomized trials and observational studies have suggested a “cascade effect” of imaging, which may create a gateway to other unnecessary care. Based in part on such evidence, the American College of Physicians has made parsimonious spine imaging a high priority in its “Choosing Wisely” campaign, aimed at reducing unnecessary care. Successful efforts to reduce unnecessary imaging have included physician education by clinical leaders, computerized decision support to identify recent imaging tests and eliminate duplication, and requiring an approved indication to order an imaging test. Other strategies have included audit and feedback regarding individual practitioners’ rates of ordering and indications and facilitating rapid access to physical therapy for patients who do not need imaging. When imaging tests are reported, it may also be useful to routinely note that some degenerative findings are common in normal, pain-free individuals. In an observational study, this strategy was associated with lower rates of repeat imaging, opioid therapy, and referral for physical therapy.

Mounting evidence of morbidities from long-term opioid therapy (including overdose, dependency, addiction, falls, fractures, accident risk, and sexual dysfunction) has prompted efforts to reduce use for chronic pain, including back pain (Chap. 18). Safety may be improved with automated reminders for high doses, early refills, or overlapping opioid and benzodiazepine prescriptions. Greater access to alternative treatments for chronic pain, such as tailored exercise programs and cognitive-behavioral therapy, may also reduce opioid prescribing.

The high cost, wide geographic variations, and rapidly increasing rates of spinal fusion surgery have prompted scrutiny over appropriate indications. Some insurance carriers have begun to limit coverage for the most controversial indications, such as low back pain without radiculopathy. Finally, educating patients and the public about the risks of imaging and excessive therapy may be necessary. A successful media campaign in Australia provides a successful model for this approach.

ALBP WITHOUT RADICULOPATHY

ALBP is defined as pain of <3 months in duration. Full recovery can be expected in more than 85% of adults with ALBP without leg pain. Most have purely “mechanical” symptoms (i.e., pain that is aggravated by motion and relieved by rest).

The initial assessment excludes serious causes of spine pathology that require urgent intervention including infection, cancer, or trauma. Risk factors for a serious cause of ALBP are shown in Table 22-1. Laboratory and imaging studies are unnecessary if risk factors are absent. CT, MRI, or plain spine films are rarely indicated in the first month of symptoms unless a spine fracture, tumor, or infection is suspected.

The prognosis is generally excellent. Many patients do not seek medical care and improve on their own. Even among those seen in primary care, two-thirds report being substantially improved after 7 weeks. This spontaneous improvement can mislead clinicians and researchers about the efficacy of treatment interventions unless subjected to rigorous prospective trials. Many treatments commonly used in the past but now known to be ineffective, including bed rest, lumbar traction, and coccygectomy, have been largely abandoned.

Clinicians should reassure patients that improvement is very likely and instruct them in self-care. Education is an important part of treatment. Satisfaction and the likelihood of follow-up increase

when patients are educated about prognosis, treatment methods, activity modifications, and strategies to prevent future exacerbations. Patients who report that they did not receive an adequate explanation for their symptoms are likely to request further diagnostic tests. In general, bed rest should be avoided for relief of severe symptoms or kept to a day or two at most. Several randomized trials suggest that bed rest does not hasten the pace of recovery. In general, the best activity recommendation is for early resumption of normal physical activity, avoiding only strenuous manual labor. Possible advantages of early ambulation for ALBP include maintenance of cardiovascular conditioning, improved disk and cartilage nutrition, improved bone and muscle strength, and increased endorphin levels. Specific back exercises or early vigorous exercise have not shown benefits for acute back pain, but may be useful for chronic pain. Use of heating pads or blankets is sometimes helpful.

Evidence-based guidelines recommend over-the-counter medicines such as acetaminophen and NSAIDs as first-line options for treatment of ALBP. In otherwise healthy patients, a trial of acetaminophen can be followed by NSAIDs for time-limited periods. In theory, the anti-inflammatory effects of NSAIDs might provide an advantage over acetaminophen to suppress inflammatory changes that accompany many causes of ALBP, but in practice, there is no clinical evidence to support the superiority of NSAIDs. The risk of renal and gastrointestinal toxicity with NSAIDs is increased in patients with preexisting medical comorbidities (e.g., renal insufficiency, cirrhosis, prior gastrointestinal hemorrhage, use of anticoagulants or steroids, heart failure). Skeletal muscle relaxants, such as cyclobenzaprine or methocarbamol, may be useful, but sedation is a common side effect. Limiting the use of muscle relaxants to nighttime only may be an option for patients with back pain that interferes with sleep.

There is no good evidence to support the use of opioid analgesics or tramadol as first-line therapy for ALBP. Their use is best reserved for patients who cannot tolerate acetaminophen or NSAIDs or for those with severe refractory pain. As with muscle relaxants, these drugs are often sedating, so it may be useful to prescribe them at nighttime only. Side effects of short-term opioid use include nausea, constipation, and pruritus; risks of long-term opioid use include hypersensitivity to pain, hypogonadism, and dependency. Falls, fractures, driving accidents, and fecal impaction are other risks. Clinical efficacy of opioids beyond 16 weeks of use is unproven.

There is no evidence to support use of oral or injected glucocorticoids for ALBP without radiculopathy. Similarly, therapies for neuropathic pain, such as gabapentin or tricyclic antidepressants, are not indicated for ALBP.

Nonpharmacologic treatments for ALBP include spinal manipulation, exercise, physical therapy, massage, acupuncture, transcutaneous electrical nerve stimulation, and ultrasound. Spinal manipulation appears to be roughly equivalent to conventional medical treatments and may be a useful alternative for patients who wish to avoid or who cannot tolerate drug therapy. There is little evidence to support the use of physical therapy, massage, acupuncture, laser therapy, therapeutic ultrasound, corsets, or lumbar traction. Although important for chronic pain, back exercises for ALBP are generally not supported by clinical evidence. There is no convincing evidence regarding the value of ice or heat applications for ALBP; however, many patients report temporary symptomatic relief from ice or frozen gel packs, and heat may produce a short-term reduction in pain after the first week. Patients often report improved satisfaction with the care that they receive when they actively participate in the selection of symptomatic approaches that are tried.

CLBP WITHOUT RADICULOPATHY

CLBP is defined as pain lasting >12 weeks; it accounts for 50% of total back pain costs. Risk factors include obesity, female gender, older age, prior history of back pain, restricted spinal mobility, pain radiating into a leg, high levels of psychological distress, poor self-rated health, minimal physical activity, smoking, job dissatisfaction, and widespread pain. In general, the same treatments that are

recommended for ALBP can be useful for patients with CLBP. In this setting, however, the long-term benefit of opioid therapy or muscle relaxants is less clear.

Evidence supports the use of exercise therapy, and this can be one of the mainstays of treatment for CLBP. Effective regimens have generally included a combination of gradually increasing aerobic exercise, strengthening exercises, and stretching exercises. Motivating patients is sometimes challenging, and in this setting, a program of supervised exercise can improve compliance. In general, activity tolerance is the primary goal, while pain relief is secondary. Supervised intensive physical exercise or “work hardening” regimens have been effective in returning some patients to work, improving walking distance, and reducing pain. In addition, some forms of yoga have been evaluated in randomized trials and may be helpful for patients who are interested. A long-term benefit of spinal manipulation or massage for CLBP is unproven.

Medications for CLBP may include acetaminophen, NSAIDs, and tricyclic antidepressants. Trials of tricyclics suggest benefit even for patients without evidence of depression. Trials do not support the efficacy of selective serotonin reuptake inhibitors (SSRIs) for CLBP. However, depression is common among patients with chronic pain and should be appropriately treated.

Cognitive-behavioral therapy is based on evidence that psychological and social factors, as well as somatic pathology, are important in the genesis of chronic pain and disability. Cognitive-behavioral therapy includes efforts to identify and modify patients’ thinking about their pain and disability. A systematic review concluded that such treatments are more effective than a waiting list control group for short-term pain relief; however, long-term results remain unclear. Behavioral treatments may have effects similar in magnitude to exercise therapy.

Back pain is the most frequent reason for seeking complementary and alternative treatments. The most common of these for back pain are spinal manipulation, acupuncture, and massage. The role of most complementary and alternative medicine approaches remains unclear. Biofeedback has not been studied rigorously. There is no convincing evidence that either spinal manipulation or transcutaneous electrical nerve stimulation (TENS) is effective in treating CLBP. Rigorous recent trials of acupuncture suggest that true acupuncture is not superior to sham acupuncture, but that both may offer an advantage over routine care. Whether this is due entirely to placebo effects provided even by sham acupuncture is uncertain. Some trials of massage therapy have been encouraging, but this has been less well studied than spinal manipulation or acupuncture.

Various injections, including epidural glucocorticoid injections, facet joint injections, and trigger point injections, have been used for treating CLBP. However, in the absence of radiculopathy, there is no evidence that these approaches are effective.

Injection studies are sometimes used diagnostically to help determine the anatomic source of back pain. The use of discography to provide evidence that a specific disk is the pain generator is not recommended. Pain relief following a glucocorticoid injection into a facet is commonly used as evidence that the facet joint is the pain source; however, the possibility that the response was a placebo effect or due to systemic absorption of the glucocorticoids is difficult to exclude.

Another category of intervention for chronic back pain is electrothermal and radiofrequency therapy. Intradiskal therapy has been proposed using both types of energy to thermocoagulate and destroy nerves in the intervertebral disk, using specially designed catheters or electrodes. Current evidence does not support the use of these intradiskal therapies.

Radiofrequency denervation is sometimes used to destroy nerves that are thought to mediate pain, and this technique has been used for facet joint pain (with the target nerve being the medial branch of the primary dorsal ramus), for back pain thought to arise from the intervertebral disk (ramus communicans), and radicular back pain (dorsal root ganglia). A few small trials have produced conflicting results for facet joint and diskogenic pain. A trial in patients with chronic radicular pain found no difference between radiofrequency

denervation of the dorsal root ganglia and sham treatment. These interventional therapies have not been studied in sufficient detail to draw conclusions of their value for CLBP.

Surgical intervention for CLBP without radiculopathy has been evaluated in a small number of randomized trials, all conducted in Europe. Each of these studies included patients with back pain and a degenerative disk, but no sciatica. Three of the four trials concluded that lumbar fusion surgery was no more effective than highly structured, rigorous rehabilitation combined with cognitive-behavioral therapy. The fourth trial found an advantage of fusion surgery over haphazard “usual care,” which appeared to be less effective than the structured rehabilitation in other trials. Given conflicting evidence, indications for surgery for CLBP without radiculopathy have remained controversial. Both U.S. and British guidelines suggest considering referral for an opinion on spinal fusion for people who have completed an optimal nonsurgical treatment program (including combined physical and psychological treatment) and who have persistent severe back pain for which they would consider surgery.

Lumbar disk replacement with prosthetic disks is U.S. Food and Drug Administration approved for uncomplicated patients needing single-level surgery at the L3-S1 levels. The disks are generally designed as metal plates with a polyethylene cushion sandwiched in between. The trials that led to approval of these devices compared them to spine fusion and concluded that the artificial disks were “not inferior.” Serious complications are somewhat more likely with the artificial disk. This treatment remains controversial for CLBP.

Intensive multidisciplinary rehabilitation programs may involve daily or frequent physical therapy, exercise, cognitive-behavioral therapy, a workplace evaluation, and other interventions. For patients who have not responded to other approaches, such programs appear to offer some benefit. Systematic reviews suggest that the evidence is limited and benefits are incremental.

Some observers have raised concern that CLBP may often be overtreated. For CLBP without radiculopathy, new British guidelines explicitly recommend against use of SSRIs, any type of injection, TENS, lumbar supports, traction, radiofrequency facet joint denervation, intradiskal electrothermal therapy, or intradiskal radiofrequency thermocoagulation. These treatments are also not recommended in guidelines from the American College of Physicians and the American Pain Society. On the other hand, exercise therapy and treatment of depression appear to be useful and underused.

LOW BACK PAIN WITH RADICULOPATHY

A common cause of back pain with radiculopathy is a herniated disk with nerve root impingement, resulting in back pain with radiation down the leg. The term sciatica is used when the leg pain radiates posteriorly in a sciatic or L5/S1 distribution. The prognosis for acute low back and leg pain with radiculopathy due to disk herniation is generally favorable, with most patients showing substantial improvement over months. Serial imaging studies suggest spontaneous regression of the herniated portion of the disk in two-thirds of patients over 6 months. Nonetheless, there are several important treatment options to provide symptomatic relief while this natural healing process unfolds.

Resumption of normal activity is recommended. Randomized trial evidence suggests that bed rest is ineffective for treating sciatica as well as back pain alone. Acetaminophen and NSAIDs are useful for pain relief, although severe pain may require short courses of opioid analgesics.

Epidural glucocorticoid injections have a role in providing temporary symptom relief for sciatica due to a herniated disk. However, there does not appear to be a benefit in terms of reducing subsequent surgical interventions. Diagnostic nerve root blocks have been advocated to determine if pain originates from a specific nerve root. However, improvement may result even when the nerve root is not responsible for the pain; this may occur as a placebo effect, from a pain-generating lesion located distally along the peripheral nerve, or from effects of systemic absorption. The utility of diagnostic nerve root blocks remains a subject of debate.

Surgical intervention is indicated for patients who have progressive motor weakness due to nerve root injury demonstrated on clinical examination or EMG. Urgent surgery is recommended for patients who have evidence of CES or spinal cord compression, generally suggested by bowel or bladder dysfunction, diminished sensation in a saddle distribution, a sensory level on the trunk, and bilateral leg weakness or spasticity.

Surgery is also an important option for patients who have disabling radicular pain despite optimal conservative treatment. Sciatica is perhaps the most common reason for recommending spine surgery. Because patients with a herniated disk and sciatica generally experience rapid improvement over a matter of weeks, most experts do not recommend considering surgery unless the patient has failed to respond to 6–8 weeks of maximum nonsurgical management. For patients who have not improved, randomized trials indicate that, compared to nonsurgical treatment, surgery results in more rapid pain relief. However, after the first year or two of follow-up, patients with sciatica appear to have much the same level of pain relief and functional improvement with or without surgery. Thus, both treatment approaches are reasonable, and patient preferences and needs (e.g., rapid return to employment) strongly influence decision making. Some patients will want the fastest possible relief and find surgical risks acceptable. Others will be more risk-averse and more tolerant of symptoms and will choose watchful waiting if they understand that improvement is likely in the end.

The usual surgical procedure is a partial hemilaminectomy with excision of the prolapsed disk (discectomy). Fusion of the involved lumbar segments should be considered only if significant spinal instability is present (i.e., degenerative spondylolisthesis). The costs associated with lumbar interbody fusion have increased dramatically in recent years. There are no large prospective, randomized trials comparing fusion to other types of surgical intervention. In one study, patients with persistent low back pain despite an initial discectomy fared no better with spine fusion than with a conservative regimen of cognitive intervention and exercise. Artificial disks have been in use in Europe for the past decade; their utility remains controversial in the United States.

PAIN IN THE NECK AND SHOULDER

Neck pain, which usually arises from diseases of the cervical spine and soft tissues of the neck, is common. Neck pain arising from the cervical spine is typically precipitated by movement and may be accompanied

by focal tenderness and limitation of motion. Many of the prior comments made regarding causes of low back pain also apply to disorders of the cervical spine. The text below will emphasize differences. Pain arising from the brachial plexus, shoulder, or peripheral nerves can be confused with cervical spine disease (Table 22-4), but the history and examination usually identify a more distal origin for the pain. Cervical spine trauma, disk disease, or spondylosis with intervertebral foraminal narrowing may be asymptomatic or painful and can produce a myelopathy, radiculopathy, or both. The same risk factors for serious causes of low back pain also apply to neck pain with the additional feature that neurologic signs of myelopathy (incontinence, sensory level, spastic legs) may also occur. Lhermitte's sign, an electrical shock down the spine with neck flexion, suggests involvement of the cervical spinal cord.

TRAUMA TO THE CERVICAL SPINE

Trauma to the cervical spine (fractures, subluxation) places the spinal cord at risk for compression. Motor vehicle accidents, violent crimes, or falls account for 87% of cervical spinal cord injuries (Chap. 456). Immediate immobilization of the neck is essential to minimize further spinal cord injury from movement of unstable cervical spine segments. The decision to obtain imaging should be based on the nature of the injury. The NEXUS low-risk criteria established that normally alert patients without palpation tenderness in the midline; intoxication; neurologic deficits; or painful distracting injuries were very unlikely to have sustained a clinically significant traumatic injury to the cervical spine. The Canadian C-spine rule recommends that imaging should be obtained following neck region trauma if the patient is >65 years old or has limb paresthesias or if there was a dangerous mechanism for the injury (e.g., bicycle collision with tree or parked car, fall from height >3 feet or five stairs, diving accident). These guidelines are helpful but must be tailored to individual circumstances; for example, patients with advanced osteoporosis, glucocorticoid use, or cancer may warrant imaging after even mild trauma. A CT scan is the diagnostic procedure of choice for detection of acute fractures following severe trauma; plain x-rays can be used for lesser degrees of trauma. When traumatic injury to the vertebral arteries or cervical spinal cord is suspected, visualization by MRI with magnetic resonance angiography is preferred.

Whiplash injury is due to rapid flexion and extension of the neck, usually in automobile accidents. The exact mechanism of the injury is unclear. This diagnosis should not be applied to patients with fractures, disk herniation, head injury, focal neurologic findings, or altered consciousness. Up to 50% of persons reporting whiplash injury acutely have persistent neck pain 1 year later. Once personal

TABLE 22-4 CERVICAL RADICULOPATHY: NEUROLOGIC FEATURES

Cervical Nerve Roots	Examination Findings			
	Reflex	Sensory	Motor	Pain Distribution
C5	Biceps	Lateral deltoid	Rhomboids ^a (elbow extends backward with hand on hip) Infraspinatus ^a (arm rotates externally with elbow flexed at the side) Deltoid ^a (arm raised laterally 30–45° from the side)	Lateral arm, medial scapula
C6	Biceps	Thumb/index finger; Dorsal hand/lateral forearm	Biceps ^a (arm flexed at the elbow in supination) Pronator teres (forearm pronated)	Lateral forearm, thumb/index fingers
C7	Triceps	Middle fingers Dorsal forearm	Triceps ^a (forearm extension, flexed at elbow) Wrist/finger extensors ^a	Posterior arm, dorsal forearm, dorsal hand
C8	Finger flexors	Palmar surface of little finger Medial hand and forearm	Abductor pollicis brevis (abduction of thumb) First dorsal interosseous (abduction of index finger) Abductor digiti minimi (abduction of little finger)	Fourth and fifth fingers, medial hand and forearm
T1	Finger flexors	Axilla and medial arm	Abductor pollicis brevis (abduction of thumb) First dorsal interosseous (abduction of index finger) Abductor digiti minimi (abduction of little finger)	Medial arm, axilla

^aThese muscles receive the majority of innervation from this root.

compensation for pain and suffering was removed from the Australian health care system, the prognosis for recovery at 1 year from whiplash injury improved also. Imaging of the cervical spine is not cost-effective acutely but is useful to detect disk herniations when symptoms persist for >6 weeks following the injury. Severe initial symptoms have been associated with a poor long-term outcome.

CERVICAL DISK DISEASE

Herniation of a lower cervical disk is a common cause of pain or tingling in the neck, shoulder, arm, or hand. Neck pain, stiffness, and a range of motion limited by pain are the usual manifestations. Herniated cervical disks are responsible for ~25% of cervical radiculopathies. Extension and lateral rotation of the neck narrow the ipsilateral intervertebral foramen and may reproduce radicular symptoms (Spurling's sign). In young adults, acute nerve root compression from a ruptured cervical disk is often due to trauma. Cervical disk herniations are usually posterolateral near the lateral recess. Typical patterns of reflex, sensory, and motor changes that accompany cervical nerve root lesions are summarized in Table 22-4. Although the classic patterns are clinically helpful, there are numerous exceptions because (1) there is overlap in sensory function between adjacent nerve roots, (2) symptoms and signs may be evident in only part of the injured nerve root territory, and (3) the location of pain is the most variable of the clinical features.

CERVICAL SPONDYLOSIS

Osteoarthritis of the cervical spine may produce neck pain that radiates into the back of the head, shoulders, or arms, or may be the source of headaches in the posterior occipital region (supplied by the C2-C4 nerve roots). Osteophytes, disk protrusions, or hypertrophic facet or uncovertebral joints may alone or in combination compress one or several nerve roots at the intervertebral foramina; these causes together account for 75% of cervical radiculopathies. The roots most commonly affected are C7 and C6. Narrowing of the spinal canal by osteophytes, ossification of the posterior longitudinal ligament (OPLL), or a large central disk may compress the cervical spinal cord and produce signs of radiculopathy and myelopathy in combination (myeloradiculopathy). When little or no neck pain accompanies cervical cord involvement, other diagnoses to be considered include amyotrophic lateral sclerosis (Chap. 452), multiple sclerosis (Chap. 458), spinal cord tumors, or syringomyelia (Chap. 456). The possibility of cervical spondylosis should be considered even when the patient presents with symptoms or signs in the legs only. MRI is the study of choice to define anatomic abnormalities of soft tissues in the cervical region including the spinal cord, but plain CT is adequate to assess bony spurs, foraminal narrowing, lateral recess stenosis, or OPLL. EMG and nerve conduction studies can localize and assess the severity of nerve root injury.

OTHER CAUSES OF NECK PAIN

Rheumatoid arthritis (RA) (Chap. 380) of the cervical facet joints produces neck pain, stiffness, and limitation of motion. Synovitis of the atlantoaxial joint (C1-C2; Fig. 22-2) may damage the transverse ligament of the atlas, producing forward displacement of the atlas on the axis (atlantoaxial subluxation). Radiologic evidence of atlantoaxial subluxation occurs in up to 30% of patients with RA. The degree of subluxation correlates with the severity of erosive disease. When subluxation is present, careful assessment is important to identify early signs of myelopathy. Occasional patients develop high spinal cord compression leading to quadriplegia, respiratory insufficiency, and death. Surgery should be considered when myelopathy or spinal instability is present. MRI is the imaging modality of choice. *Ankylosing spondylitis* can cause neck pain and less commonly atlantoaxial subluxation; surgery may be required to prevent spinal cord compression.

Acute *herpes zoster* can present as acute posterior occipital or neck pain prior to the outbreak of vesicles. *Neoplasms* metastatic to the cervical spine, *infections* (osteomyelitis and epidural abscess), and *metabolic bone diseases* may be the cause of neck pain, as discussed above

among causes of low back pain. Neck pain may also be referred from the heart with coronary artery ischemia (cervical angina syndrome).

THORACIC OUTLET SYNDROMES

The thoracic outlet contains the first rib, the subclavian artery and vein, the brachial plexus, the clavicle, and the lung apex. Injury to these structures may result in postural or movement-induced pain around the shoulder and supraclavicular region, classified as follows.

True neurogenic thoracic outlet syndrome (TOS) is an uncommon disorder resulting from compression of the lower trunk of the brachial plexus or ventral rami of the C8 or T1 nerve roots, caused most often by an anomalous band of tissue connecting an elongate transverse process at C7 with the first rib. Pain is mild or may be absent. Signs include weakness and wasting of intrinsic muscles of the hand and diminished sensation on the palmar aspect of the fifth digit. An anteroposterior cervical spine x-ray will show an elongate C7 transverse process (an anatomic marker for the anomalous cartilaginous band), and EMG and nerve conduction studies confirm the diagnosis. Treatment consists of surgical resection of the anomalous band. The weakness and wasting of intrinsic hand muscles typically does not improve, but surgery halts the insidious progression of weakness.

Arterial TOS results from compression of the subclavian artery by a cervical rib, resulting in poststenotic dilatation of the artery and in some cases secondary thrombus formation. Blood pressure is reduced in the affected limb, and signs of emboli may be present in the hand. Neurologic signs are absent. Ultrasound can confirm the diagnosis noninvasively. Treatment is with thrombolysis or anticoagulation (with or without embolectomy) and surgical excision of the cervical rib compressing the subclavian artery.

Venous TOS is due to subclavian vein thrombosis resulting in swelling of the arm and pain. The vein may be compressed by a cervical rib or anomalous scalene muscle. Venography is the diagnostic test of choice.

Disputed TOS accounts for 95% of patients diagnosed with TOS; chronic arm and shoulder pain are prominent and of unclear cause. The lack of sensitive and specific findings on physical examination or specific markers for this condition results in diagnostic uncertainty. The role of surgery in disputed TOS is controversial. Multidisciplinary pain management is a conservative approach, although treatment is often unsuccessful.

BRACHIAL PLEXUS AND NERVES

Pain from injury to the brachial plexus or peripheral nerves of the arm can occasionally mimic referred pain of cervical spine origin including cervical radiculopathy. Neoplastic infiltration of the lower trunk of the brachial plexus may produce shoulder or supraclavicular pain radiating down the arm, numbness of the fourth and fifth fingers or medial forearm, and weakness of intrinsic hand muscles innervated by the ulnar and median nerves. Delayed radiation injury may produce similar findings, although pain is less often present and almost always less severe. A Pancoast tumor of the lung (Chap. 107) is another cause and should be considered, especially when a concurrent Horner's syndrome is present. *Suprascapular neuropathy* may produce severe shoulder pain, weakness, and wasting of the supraspinatus and infraspinatus muscles. *Acute brachial neuritis* is often confused with radiculopathy; the acute onset of severe shoulder or scapular pain is followed typically over days by weakness of the proximal arm and shoulder girdle muscles innervated by the upper brachial plexus. The onset may be preceded by an infection, vaccination, or minor surgical procedure. The long thoracic nerve may be affected resulting in a winged scapula. Brachial neuritis may also present as an isolated paralysis of the diaphragm with or without involvement of other nerves of the upper limb. Recovery may take up to 3 years.

Occasional cases of carpal tunnel syndrome produce pain and paresthesias extending into the forearm, arm, and shoulder resembling a C5 or C6 root lesion. Lesions of the radial or ulnar nerve can mimic a radiculopathy at C7 or C8, respectively. EMG and nerve conduction

studies can accurately localize lesions to the nerve roots, brachial plexus, or peripheral nerves.

For further discussion of peripheral nerve disorders, see Chap. 459.

SHOULDER

Pain arising from the shoulder can on occasion mimic pain from the spine. If symptoms and signs of radiculopathy are absent, then the differential diagnosis includes mechanical shoulder pain (tendonitis, bursitis, rotator cuff tear, dislocation, adhesive capsulitis, or rotator cuff impingement under the acromion) and referred pain (subdiaphragmatic irritation, angina, Pancoast tumor). Mechanical pain is often worse at night, associated with local shoulder tenderness and aggravated by passive abduction, internal rotation, or extension of the arm. Pain from shoulder disease may radiate into the arm or hand, but focal neurologic signs (sensory, motor, or reflex changes) are absent.

TREATMENT NECK PAIN WITHOUT RADICULOPATHY

The evidence regarding treatment for neck pain is less complete than that for low back pain, but the approach is remarkably similar in many respects. As with low back pain, spontaneous improvement is the norm for acute neck pain. The usual goals of therapy are to promote a rapid return to normal function and provide symptom relief while healing proceeds.

The evidence in support of nonsurgical treatments for whiplash-associated disorders is generally of limited quality and neither supports nor refutes the common treatments used for symptom relief. Gentle mobilization of the cervical spine combined with exercise programs may be beneficial. Evidence is insufficient to recommend for or against the routine use of acupuncture, cervical traction, TENS, ultrasound, diathermy, or massage. Some patients obtain modest relief using a soft neck collar; there is little risk or cost.

For patients with neck pain unassociated with trauma, supervised exercise with or without mobilization appears to be effective. Exercises often include shoulder rolls and neck stretches. The evidence for the use of muscle relaxants, analgesics, and NSAIDs in acute and chronic neck pain is of lower quality and less consistent than for low back pain.

Low-level laser therapy directed at areas of tenderness, local acupuncture points, or a grid of predetermined points is a controversial approach to the treatment of neck pain. A 2009 meta-analysis suggested that this treatment may provide greater pain relief than sham

therapy for both acute and chronic neck pain, but comparison to other conservative and less expensive treatment measures is needed.

Although some surgical studies have proposed a role for anterior discectomy and fusion in patients with neck pain, these studies generally have not been rigorously conducted. A systematic review suggested that there was no valid clinical evidence to support either cervical fusion or cervical disk arthroplasty in patients with neck pain without radiculopathy. Similarly, there is no evidence to support radiofrequency neurotomy or cervical facet injections for neck pain without radiculopathy.

TREATMENT NECK PAIN WITH RADICULOPATHY

The natural history of neck pain with acute radiculopathy due to disk disease is favorable, and many patients will improve without specific therapy. Although there are no randomized trials of NSAIDs for neck pain, a course of NSAIDs, acetaminophen, or both, with or without muscle relaxants, is reasonable as initial therapy. Other nonsurgical treatments are commonly used, including opioid analgesics, oral glucocorticoids, cervical traction, and immobilization with a hard or soft cervical collar. However, there are no randomized trials that establish the effectiveness of these treatments. Soft cervical collars can be modestly helpful by limiting spontaneous and reflex neck movements that exacerbate pain.

As for lumbar radiculopathy, epidural glucocorticoids appear to provide short-term symptom relief in cervical radiculopathy, but rigorous studies addressing this question have not been conducted. If cervical radiculopathy is due to bony compression from cervical spondylosis with foraminal narrowing, periodic follow-up to assess for progression is indicated and consideration of surgical decompression is reasonable.

Surgical treatment can produce rapid pain relief, although it is unclear whether long-term outcomes are improved over nonsurgical therapy. Indications for cervical disk surgery include a progressive radicular motor deficit, functionally limiting pain that fails to respond to conservative management, or spinal cord compression.

Surgical treatments include anterior cervical discectomy alone, laminectomy with discectomy, or discectomy with fusion. The risk of subsequent radiculopathy or myelopathy at cervical segments adjacent to a fusion is ~3% per year and 26% per decade. Although this risk is sometimes portrayed as a late complication of surgery, it may also reflect the natural history of degenerative cervical disk disease.

SECTION 2 ALTERATIONS IN BODY TEMPERATURE

23

Fever

Charles A. Dinarello, Reuven Porat

Body temperature is controlled by the hypothalamus. Neurons in both the preoptic anterior hypothalamus and the posterior hypothalamus receive two kinds of signals: one from peripheral nerves that transmit information from warmth/cold receptors in the skin and the other from the temperature of the blood bathing the region. These two types of signals are integrated by the thermoregulatory center of the hypothalamus to maintain normal temperature. In a neutral temperature environment, the human metabolic rate produces more heat than is necessary to maintain the core body temperature in the range of 36.5–37.5°C (97.7–99.5°F).

A normal body temperature is ordinarily maintained despite environmental variations because the hypothalamic thermoregulatory center balances the excess heat production derived from metabolic activity in muscle and the liver with heat dissipation from the skin and lungs. According to studies of healthy individuals 18–40 years of age, the mean oral temperature is 36.8° ± 0.4°C (98.2° ± 0.7°F), with low levels at 6 A.M. and higher levels at 4–6 P.M. The maximal normal oral temperature is 37.2°C (98.9°F) at 6 A.M. and 37.7°C (99.9°F) at 4 P.M.; these values define the 99th percentile for healthy individuals. In light of these studies, *an A.M. temperature of >37.2°C (>98.9°F) or a P.M. temperature of >37.7°C (>99.9°F) would define a fever.* The normal daily temperature variation is typically 0.5°C (0.9°F). However, in some individuals recovering from a febrile illness, this daily variation can be as great as 1.0°C. During a febrile illness, the diurnal variation is usually maintained, but at higher, febrile levels. The daily temperature

variation appears to be fixed in early childhood; in contrast, elderly individuals can exhibit a reduced ability to develop fever, with only a modest fever even in severe infections.

Rectal temperatures are generally 0.4°C (0.7°F) higher than oral readings. The lower oral readings are probably attributable to mouth breathing, which is a factor in patients with respiratory infections and rapid breathing. Lower-esophageal temperatures closely reflect core temperature. Tympanic membrane thermometers measure radiant heat from the tympanic membrane and nearby ear canal and display that absolute value (*unadjusted mode*) or a value automatically calculated from the absolute reading on the basis of nomograms relating the radiant temperature measured to actual core temperatures obtained in clinical studies (*adjusted mode*). These measurements, although convenient, may be more variable than directly determined oral or rectal values. Studies in adults show that readings are lower with unadjusted-mode than with adjusted-mode tympanic membrane thermometers and that unadjusted-mode tympanic membrane values are 0.8°C (1.6°F) lower than rectal temperatures.

In women who menstruate, the A.M. temperature is generally lower in the 2 weeks before ovulation; it then rises by ~0.6°C (1°F) with ovulation and remains at that level until menses occur. Body temperature can be elevated in the postprandial state. Pregnancy and endocrinologic dysfunction also affect body temperature.

FEVER VERSUS HYPERTHERMIA

Fever is an elevation of body temperature that exceeds the normal daily variation and occurs *in conjunction with an increase in the hypothalamic set point* (e.g., from 37°C to 39°C). This shift of the set point from “normothermic” to febrile levels very much resembles the resetting of the home thermostat to a higher level in order to raise the ambient temperature in a room. Once the hypothalamic set point is raised, neurons in the vasomotor center are activated and vasoconstriction commences. The individual first notices vasoconstriction in the hands and feet. Shunting of blood away from the periphery to the internal organs essentially decreases heat loss from the skin, and the person feels cold. For most fevers, body temperature increases by 1–2°C. Shivering, which increases heat production from the muscles, may begin at this time; however, shivering is not required if heat conservation mechanisms raise blood temperature sufficiently. Nonshivering heat production from the liver also contributes to increasing core temperature. Behavioral adjustments (e.g., putting on more clothing or bedding) help raise body temperature by decreasing heat loss.

The processes of heat conservation (vasoconstriction) and heat production (shivering and increased nonshivering thermogenesis) continue until the temperature of the blood bathing the hypothalamic neurons matches the new thermostat setting. Once that point is reached, the hypothalamus maintains the temperature at the febrile level by the same mechanisms of heat balance that function in the afebrile state. When the hypothalamic set point is again reset downward (in response to either a reduction in the concentration of pyrogens or the use of antipyretics), the processes of heat loss through vasodilation and sweating are initiated. Loss of heat by sweating and vasodilation continues until the blood temperature at the hypothalamic level matches the lower setting. Behavioral changes (e.g., removal of clothing) facilitate heat loss.

A fever of >41.5°C (>106.7°F) is called *hyperpyrexia*. This extraordinarily high fever can develop in patients with severe infections but most commonly occurs in patients with central nervous system (CNS) hemorrhages. In the preantibiotic era, fever due to a variety of infectious diseases rarely exceeded 106°F, and there has been speculation that this natural “thermal ceiling” is mediated by neuropeptides functioning as central antipyretics.

In rare cases, the hypothalamic set point is elevated as a result of local trauma, hemorrhage, tumor, or intrinsic hypothalamic malfunction. The term *hypothalamic fever* is sometimes used to describe elevated temperature caused by abnormal hypothalamic function. However, most patients with hypothalamic damage have *subnormal*, not *supra-normal*, body temperatures.

Although most patients with elevated body temperature have fever, there are circumstances in which elevated temperature represents not fever but *hyperthermia* (*heat stroke*). Hyperthermia is characterized by an uncontrolled increase in body temperature that exceeds the body’s ability to lose heat. The setting of the hypothalamic thermoregulatory center is unchanged. In contrast to fever in infections, hyperthermia does not involve pyrogenic molecules. Exogenous heat exposure and endogenous heat production are two mechanisms by which hyperthermia can result in dangerously high internal temperatures. Excessive heat production can easily cause hyperthermia despite physiologic and behavioral control of body temperature. For example, work or exercise in hot environments can produce heat faster than peripheral mechanisms can lose it. [For a detailed discussion of hyperthermia, see Chap. 479e.](#)

It is important to distinguish between fever and hyperthermia since hyperthermia can be rapidly fatal and characteristically does not respond to antipyretics. In an emergency situation, however, making this distinction can be difficult. For example, in systemic sepsis, fever (hyperpyrexia) can be rapid in onset, and temperatures can exceed 40.5°C (104.9°F). Hyperthermia is often diagnosed on the basis of the events immediately preceding the elevation of core temperature—e.g., heat exposure or treatment with drugs that interfere with thermoregulation. In patients with heat stroke syndromes and in those taking drugs that block sweating, the skin is hot but dry, whereas in fever the skin can be cold as a consequence of vasoconstriction. Antipyretics do not reduce the elevated temperature in hyperthermia, whereas in fever—and even in hyperpyrexia—adequate doses of either aspirin or acetaminophen usually result in some decrease in body temperature.

PATHOGENESIS OF FEVER

PYROGENS

The term *pyrogen* (Greek *pyro*, “fire”) is used to describe any substance that causes fever. *Exogenous* pyrogens are derived from outside the patient; most are microbial products, microbial toxins, or whole microorganisms (including viruses). The classic example of an exogenous pyrogen is the lipopolysaccharide (endotoxin) produced by all gram-negative bacteria. Pyrogenic products of gram-positive organisms include the enterotoxins of *Staphylococcus aureus* and the groups A and B streptococcal toxins, also called *superantigens*. One staphylococcal toxin of clinical importance is that associated with isolates of *S. aureus* from patients with toxic shock syndrome. These products of staphylococci and streptococci cause fever in experimental animals when injected intravenously at concentrations of 1–10 µg/kg. Endotoxin is a highly pyrogenic molecule in humans: when injected intravenously into volunteers, a dose of 2–3 ng/kg produces fever, leukocytosis, acute-phase proteins, and generalized symptoms of malaise.

PYROGENIC CYTOKINES

Cytokines are small proteins (molecular mass, 10,000–20,000 Da) that regulate immune, inflammatory, and hematopoietic processes. For example, the elevated leukocytosis seen in several infections with an absolute neutrophilia is attributable to the cytokines interleukin (IL) 1 and IL-6. Some cytokines also cause fever; formerly referred to as *endogenous pyrogens*, they are now called *pyrogenic cytokines*. The pyrogenic cytokines include IL-1, IL-6, tumor necrosis factor (TNF), and ciliary neurotropic factor, a member of the IL-6 family. Interferons (IFNs), particularly IFN-α, also are pyrogenic cytokines; fever is a prominent side effect of IFN-α used in the treatment of hepatitis. Each pyrogenic cytokine is encoded by a separate gene, and each has been shown to cause fever in laboratory animals and in humans. When injected into humans at low doses (10–100 ng/kg), IL-1 and TNF produce fever; in contrast, for IL-6, a dose of 1–10 µg/kg is required for fever production.

A wide spectrum of bacterial and fungal products induce the synthesis and release of pyrogenic cytokines. However, fever can be a manifestation of disease in the absence of microbial infection. For example, inflammatory processes, trauma, tissue necrosis, and antigen-antibody complexes induce the production of IL-1, TNF,

and/or IL-6; individually or in combination, these cytokines trigger the hypothalamus to raise the set point to febrile levels.

ELEVATION OF THE HYPOTHALAMIC SET POINT BY CYTOKINES

During fever, levels of prostaglandin E₂ (PGE₂) are elevated in hypothalamic tissue and the third cerebral ventricle. The concentrations of PGE₂ are highest near the circumventricular vascular organs (organum vasculosum of lamina terminalis)—networks of enlarged capillaries surrounding the hypothalamic regulatory centers. Destruction of these organs reduces the ability of pyrogens to produce fever. Most studies in animals have failed to show, however, that pyrogenic cytokines pass from the circulation into the brain itself. Thus, it appears that both exogenous pyrogens and pyrogenic cytokines interact with the endothelium of these capillaries and that this interaction is the first step in initiating fever—i.e., in raising the set point to febrile levels.

The key events in the production of fever are illustrated in Fig. 23-1. Myeloid and endothelial cells are the primary cell types that produce pyrogenic cytokines. Pyrogenic cytokines such as IL-1, IL-6, and TNF are released from these cells and enter the systemic circulation. Although these circulating cytokines lead to fever by inducing the synthesis of PGE₂, they also induce PGE₂ in peripheral tissues. The increase in PGE₂ in the periphery accounts for the nonspecific myalgias and arthralgias that often accompany fever. It is thought that some systemic PGE₂ escapes destruction by the lung and gains access to the hypothalamus via the internal carotid. However, it is the elevation of PGE₂ in the brain that starts the process of raising the hypothalamic set point for core temperature.

There are four receptors for PGE₂, and each signals the cell in different ways. Of the four receptors, the third (EP-3) is essential for fever: when the gene for this receptor is deleted in mice, no fever follows the injection of IL-1 or endotoxin. Deletion of the other PGE₂ receptor genes leaves the fever mechanism intact. Although PGE₂ is essential for fever, it is not a neurotransmitter. Rather, the release of PGE₂ from the brain side of the hypothalamic endothelium triggers the PGE₂ receptor on glial cells, and this stimulation results in the rapid release of cyclic adenosine 5'-monophosphate (cAMP), which is a neurotransmitter. As shown in Fig. 23-1, the release of cAMP from glial cells activates neuronal endings from the thermoregulatory center that extend into the area. The elevation of cAMP is thought to account for changes in the hypothalamic set point either directly or indirectly (by inducing the release of neurotransmitters). Distinct receptors for microbial products are located on the hypothalamic endothelium. These receptors are called *Toll-like receptors* and are similar in many ways to IL-1 receptors. IL-1 receptors and Toll-like receptors share the same signal-transducing mechanism. Thus, the direct activation of Toll-like receptors or IL-1 receptors results in PGE₂ production and fever.

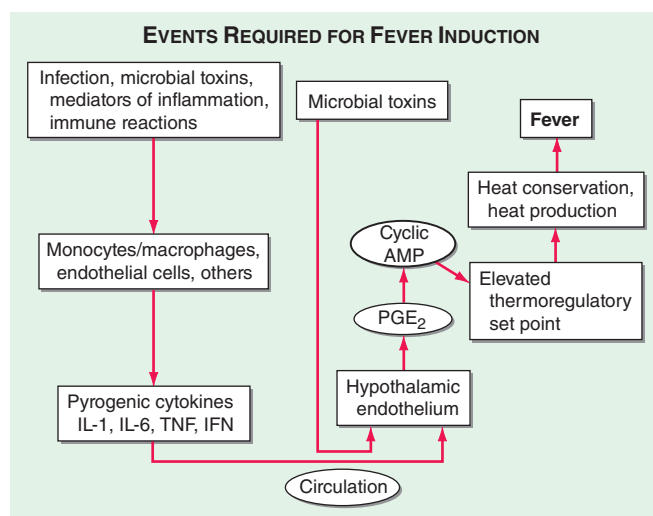


FIGURE 23-1 Chronology of events required for the induction of fever. AMP, adenosine 5'-monophosphate; IFN, interferon; IL, interleukin; PGE₂, prostaglandin E₂; TNF, tumor necrosis factor.

PRODUCTION OF CYTOKINES IN THE CNS

Cytokines produced in the brain may account for the hyperpyrexia of CNS hemorrhage, trauma, or infection. Viral infections of the CNS induce microglial and possibly neuronal production of IL-1, TNF, and IL-6. In experimental animals, the concentration of a cytokine required to cause fever is several orders of magnitude lower with direct injection into the brain substance or brain ventricles than with systemic injection. Therefore, cytokines produced in the CNS can raise the hypothalamic set point, bypassing the circumventricular organs. CNS cytokines likely account for the hyperpyrexia of CNS hemorrhage, trauma, or infection.

APPROACH TO THE PATIENT: Fever

PHYSICAL EXAMINATION

The chronology of events preceding fever, including exposure to other infected individuals or to vectors of disease, should be ascertained. Electronic devices for measuring oral, tympanic membrane, or rectal temperatures are reliable, but the same site should be used consistently to monitor a febrile disease. Moreover, physicians should be aware that newborns, elderly patients, patients with chronic liver or renal failure, and patients taking glucocorticoids or being treated with an anticytokine may have active infection in the absence of fever due to a blunted febrile response.

LABORATORY TESTS

The workup should include a complete blood count; a differential count should be performed manually or with an instrument sensitive to the identification of juvenile or band forms, toxic granulations, and Döhle bodies, which are suggestive of bacterial infection. Neutropenia may be present with some viral infections.

Measurement of circulating cytokines in patients with fever is not helpful since levels of cytokines such as IL-1 and TNF in the circulation often are below the detection limit of the assay or do not coincide with fever. However, in patients with low-grade fevers or possible disease, the most valuable measurements are the C-reactive protein level and the erythrocyte sedimentation rate. These markers of inflammatory processes are particularly helpful in detecting occult disease. Measurement of circulating IL-6 is useful because IL-6 induces C-reactive protein. **Acute-phase reactants are discussed in Chap. 325.**

FEVER IN PATIENTS RECEIVING ANTICYTOKINE THERAPY

Patients receiving long-term treatment with anticytokine-based regimens are at a disadvantage because of lowered host defense against infection. Even when the results of tests for latent *Mycobacterium tuberculosis* infection are negative, active tuberculosis can develop in patients receiving anti-TNF therapy. With the increasing use of anticytokines to reduce the activity of IL-1, IL-6, IL-12, or TNF in patients with Crohn's disease, rheumatoid arthritis, or psoriasis, the possibility that these therapies blunt the febrile response must be kept in mind.

The blocking of cytokine activity has the distinct clinical drawback of lowering the level of host defenses against both routine bacterial and opportunistic infections. The opportunistic infections reported in patients treated with agents that neutralize TNF- α are similar to those reported in the HIV-1-infected population (e.g., a new infection with or reactivation of *Mycobacterium tuberculosis*, with dissemination).

In nearly all reported cases of infection associated with anticytokine therapy, fever is among the presenting signs. However, the extent to which the febrile response is blunted in these patients remains unknown. A similar situation is seen in patients receiving high-dose glucocorticoid therapy or anti-inflammatory agents such as ibuprofen. Therefore, low-grade fever is of considerable concern in patients receiving anticytokine therapies. The physician should conduct an early and rigorous diagnostic evaluation in these patients.

THE DECISION TO TREAT FEVER

Most fevers are associated with self-limited infections, such as common viral diseases. The use of antipyretics is not contraindicated in these infections: no significant clinical evidence indicates either that antipyretics delay the resolution of viral or bacterial infections or that fever facilitates recovery from infection or acts as an adjuvant to the immune system. In short, treatment of fever and its symptoms with routine antipyretics does no harm and does not slow the resolution of common viral and bacterial infections.

However, in bacterial infections, the withholding of antipyretic therapy can be helpful in evaluating the effectiveness of a particular antibiotic, especially in the absence of positive cultures of the infecting organism, and the routine use of antipyretics can mask an inadequately treated bacterial infection. Withholding antipyretics in some cases may facilitate the diagnosis of an unusual febrile disease. Temperature-pulse dissociation (*relative bradycardia*) occurs in typhoid fever, brucellosis, leptospirosis, some drug-induced fevers, and factitious fever. As stated earlier, in newborns, elderly patients, patients with chronic liver or kidney failure, and patients taking glucocorticoids, fever may not be present despite infection. Hypothermia can develop in patients with septic shock.

Some infections have characteristic patterns in which febrile episodes are separated by intervals of normal temperature. For example, *Plasmodium vivax* causes fever every third day, whereas fever occurs every fourth day with *P. malariae*. Another relapsing fever is related to *Borrelia* infection, with days of fever followed by a several-day afebrile period and then a relapse into additional days of fever. In the Pel-Ebstein pattern, fever lasting 3–10 days is followed by afebrile periods of 3–10 days; this pattern can be classic for Hodgkin's disease and other lymphomas. In cyclic neutropenia, fevers occur every 21 days and accompany the neutropenia. There is no periodicity of fever in patients with familial Mediterranean fever. However, these patterns have limited or no diagnostic value compared with specific and rapid laboratory tests.

ANTICYTOKINE THERAPY TO REDUCE FEVER IN AUTOIMMUNE AND AUTOINFLAMMATORY DISEASES

Recurrent fever is documented at some point in most autoimmune diseases and nearly all autoinflammatory diseases. Although fever can be a manifestation of autoimmune diseases, recurrent fevers are characteristic of autoinflammatory diseases (Table 23-1), including adult and juvenile Still's disease, familial Mediterranean fever, and hyper-IgD syndrome. In addition to recurrent fevers, neutrophilia and serosal inflammation characterize autoinflammatory diseases. The fevers associated with these illnesses are dramatically reduced

TABLE 23-1 AUTOINFLAMMATORY DISEASES IN WHICH FEVER IS CHARACTERISTIC

Adult and juvenile Still's disease
Cryopyrin-associated periodic syndromes (CAPS)
Familial Mediterranean fever
Hyper-IgD syndrome
Behçet's syndrome
Macrophage activation syndrome
Normocomplementemic urticarial vasculitis
Antisynthetase myositis
PAPA ^a syndrome
Blau syndrome
Gouty arthritis

^aPyogenic arthritis, pyoderma gangrenosum, and acne.

by blocking of IL-1 β activity. Anticytokines therefore reduce fever in autoimmune and autoinflammatory diseases. Although fevers in autoinflammatory diseases are mediated by IL-1 β , patients also respond to antipyretics.

MECHANISMS OF ANTIPYRETIC AGENTS

The reduction of fever by lowering of the elevated hypothalamic set point is a direct function of reduction of the PGE₂ level in the thermoregulatory center. The synthesis of PGE₂ depends on the constitutively expressed enzyme cyclooxygenase. The substrate for cyclooxygenase is arachidonic acid released from the cell membrane, and this release is the rate-limiting step in the synthesis of PGE₂. Therefore, inhibitors of cyclooxygenase are potent antipyretics. The antipyretic potency of various drugs is directly correlated with the inhibition of brain cyclooxygenase. Acetaminophen is a poor cyclooxygenase inhibitor in peripheral tissue and lacks noteworthy anti-inflammatory activity; in the brain, however, acetaminophen is oxidized by the p450 cytochrome system, and the oxidized form inhibits cyclooxygenase activity. Moreover, in the brain, the inhibition of another enzyme, COX-3, by acetaminophen may account for the antipyretic effect of this agent. However, COX-3 is not found outside the CNS.

Oral aspirin and acetaminophen are equally effective in reducing fever in humans. Nonsteroidal anti-inflammatory drugs (NSAIDs) such as ibuprofen and specific inhibitors of COX-2 also are excellent antipyretics. Chronic, high-dose therapy with antipyretics such as aspirin or any NSAID does not reduce normal core body temperature. Thus, PGE₂ appears to play no role in normal thermoregulation.

As effective antipyretics, glucocorticoids act at two levels. First, similar to the cyclooxygenase inhibitors, glucocorticoids reduce PGE₂ synthesis by inhibiting the activity of phospholipase A₂, which is needed to release arachidonic acid from the cell membrane. Second, glucocorticoids block the transcription of the mRNA for the pyrogenic cytokines. Limited experimental evidence indicates that ibuprofen and COX-2 inhibitors reduce IL-1-induced IL-6 production and may contribute to the antipyretic activity of NSAIDs.

REGIMENS FOR THE TREATMENT OF FEVER

The objectives in treating fever are first to reduce the elevated hypothalamic set point and second to facilitate heat loss. Reducing fever with antipyretics also reduces systemic symptoms of headache, myalgias, and arthralgias.

Oral aspirin and NSAIDs effectively reduce fever but can adversely affect platelets and the gastrointestinal tract. Therefore, acetaminophen is preferred as an antipyretic. In children, acetaminophen or oral ibuprofen must be used because aspirin increases the risk of Reye's syndrome. If the patient cannot take oral antipyretics, parenteral preparations of NSAIDs and rectal suppositories of various antipyretics can be used.

Treatment of fever in some patients is highly recommended. Fever increases the demand for oxygen (i.e., for every increase of 1°C over 37°C, there is a 13% increase in oxygen consumption) and can aggravate the condition of patients with preexisting impairment of cardiac, pulmonary, or CNS function. Children with a history of febrile or nonfebrile seizure should be aggressively treated to reduce fever. However, it is unclear what triggers the febrile seizure, and there is no correlation between absolute temperature elevation and onset of a febrile seizure in susceptible children.

In hyperpyrexia, the use of cooling blankets facilitates the reduction of temperature; however, cooling blankets should not be used without oral antipyretics. In hyperpyretic patients with CNS disease or trauma (CNS bleeding), reducing core temperature mitigates the detrimental effects of high temperature on the brain.

For a discussion of treatment for hyperthermia, see Chap. 479e.

24

Fever and Rash

Elaine T. Kaye, Kenneth M. Kaye

The acutely ill patient with fever and rash may present a diagnostic challenge for physicians. However, the distinctive appearance of an eruption in concert with a clinical syndrome can facilitate a prompt diagnosis and the institution of life-saving therapy or critical infection-control interventions. **Representative images of many of the rashes discussed in this chapter are included in Chap. 25e.**

APPROACH TO THE PATIENT: Fever and Rash

A thorough history of patients with fever and rash includes the following relevant information: immune status, medications taken within the previous month, specific travel history, immunization status, exposure to domestic pets and other animals, history of animal (including arthropod) bites, recent dietary exposures, existence of cardiac abnormalities, presence of prosthetic material, recent exposure to ill individuals, and exposure to sexually transmitted diseases. The history should also include the site of onset of the rash and its direction and rate of spread.

A thorough physical examination entails close attention to the rash, with an assessment and precise definition of its salient features. First, it is critical to determine what *type* of lesions make up the eruption. *Macules* are flat lesions defined by an area of changed color (i.e., a blanchable erythema). *Papules* are raised, solid lesions <5 mm in diameter; *plaques* are lesions >5 mm in diameter with a flat, plateau-like surface; and *nodules* are lesions >5 mm in diameter with a more rounded configuration. *Wheals* (urticaria, hives) are papules or plaques that are pale pink and may appear annular (ringlike) as they enlarge; classic (nonvasculitic) wheals are transient, lasting only 24 h in any defined area. *Vesicles* (<5 mm) and *bullae* (>5 mm) are circumscribed, elevated lesions containing fluid. *Pustules* are raised lesions containing purulent exudate; vesicular processes such as varicella or herpes simplex may evolve to pustules. *Nonpalpable purpura* is a flat lesion that is due to bleeding into the skin. If <3 mm in diameter, the purpuric lesions are termed *petechiae*; if >3 mm, they are termed *ecchymoses*. *Palpable purpura* is a raised lesion that is due to inflammation of the vessel wall (vasculitis) with subsequent hemorrhage. An *ulcer* is a defect in the skin extending at least into the upper layer of the dermis, and an *eschar* (tâche noire) is a necrotic lesion covered with a black crust.

Other pertinent features of rashes include their *configuration* (i.e., annular or target), the *arrangement* of their lesions, and their *distribution* (i.e., central or peripheral).

For further discussion, see Chaps. 70, 72, and 147.

CLASSIFICATION OF RASH

This chapter reviews rashes that reflect systemic disease, but it does not include localized skin eruptions (i.e., cellulitis, impetigo) that may also be associated with fever (Chap. 156). The chapter is not intended to be all-inclusive, but it covers the most important and most common diseases associated with fever and rash. Rashes are classified herein on the basis of lesion morphology and distribution. For practical purposes, this classification system is based on the most typical disease presentations. However, morphology may vary as rashes evolve, and the presentation of diseases with rashes is subject to many variations (Chap. 72). For instance, the classic petechial rash of Rocky Mountain spotted fever (Chap. 211) may initially consist of blanchable erythematous macules distributed peripherally; at times, however, the rash associated with this disease may not be predominantly acral, or no rash may develop at all.

Diseases with fever and rash may be classified by type of eruption: centrally distributed maculopapular, peripheral, confluent desquamative erythematous, vesiculobullous, urticaria-like, nodular, purpuric, ulcerated, or with eschars. Diseases are listed by these categories in Table 24-1, and many are highlighted in the text. However, for a more detailed discussion of each disease associated with a rash, the reader is referred to the chapter dealing with that specific disease. (Reference chapters are cited in the text and listed in Table 24-1.)

CENTRALLY DISTRIBUTED MACULOPAPULAR ERUPTIONS

Centrally distributed rashes, in which lesions are primarily truncal, are the most common type of eruption. The rash of *rubeola* (measles) starts at the hairline 2–3 days into the illness and moves down the body, typically sparing the palms and soles (Chap. 229). It begins as discrete erythematous lesions, which become confluent as the rash spreads. Koplik's spots (1- to 2-mm white or bluish lesions with an erythematous halo on the buccal mucosa) are pathognomonic for measles and are generally seen during the first 2 days of symptoms. They should not be confused with Fordyce's spots (ectopic sebaceous glands), which have no erythematous halos and are found in the mouth of healthy individuals. Koplik's spots may briefly overlap with the measles exanthem.

Rubella (German measles) also spreads from the hairline downward; unlike that of measles, however, the rash of rubella tends to clear from originally affected areas as it migrates, and it may be pruritic (Chap. 230e). Forchheimer spots (palatal petechiae) may develop but are nonspecific because they also develop in *infectious mononucleosis* (Chap. 218) and *scarlet fever* (Chap. 173). Postauricular and suboccipital adenopathy and arthritis are common among adults with rubella. Exposure of pregnant women to ill individuals should be avoided, as rubella causes severe congenital abnormalities. Numerous strains of *enteroviruses* (Chap. 228), primarily echoviruses and coxsackieviruses, cause nonspecific syndromes of fever and eruptions that may mimic rubella or measles. Patients with *infectious mononucleosis* caused by Epstein-Barr virus (Chap. 218) or with *primary HIV infection* (Chap. 226) may exhibit pharyngitis, lymphadenopathy, and a nonspecific maculopapular exanthem.

The rash of *erythema infectiosum* (fifth disease), which is caused by human parvovirus B19, primarily affects children 3–12 years old; it develops after fever has resolved as a bright blanchable erythema on the cheeks ("slapped cheeks") with perioral pallor (Chap. 221). A more diffuse rash (often pruritic) appears the next day on the trunk and extremities and then rapidly develops into a lacy reticular eruption that may wax and wane (especially with temperature change) over 3 weeks. Adults with fifth disease often have arthritis, and fetal hydrops can develop in association with this condition in pregnant women.

Exanthem subitum (roseola) is caused by human herpesvirus 6 and is most common among children <3 years of age (Chap. 219). As in *erythema infectiosum*, the rash usually appears after fever has subsided. It consists of 2- to 3-mm rose-pink macules and papules that coalesce only rarely, occur initially on the trunk and sometimes on the extremities (sparing the face), and fade within 2 days.

Although drug reactions have many manifestations, including urticaria, exanthematous *drug-induced eruptions* (Chap. 74) are most common and are often difficult to distinguish from viral exanthems. Eruptions elicited by drugs are usually more intensely erythematous and pruritic than viral exanthems, but this distinction is not reliable. A history of new medications and an absence of prostration may help to distinguish a drug-related rash from an eruption of another etiology. Rashes may persist for up to two weeks after administration of the offending agent is discontinued. Certain populations are more prone than others to drug rashes. Of HIV-infected patients, 50–60% develop a rash in response to sulfa drugs; 90% of patients with mononucleosis due to Epstein-Barr virus develop a rash when given ampicillin.

Rickettsial illnesses (Chap. 211) should be considered in the evaluation of individuals with centrally distributed maculopapular eruptions. The usual setting for *epidemic typhus* is a site of war or natural disaster in which people are exposed to body lice. *Endemic typhus* or *leptospirosis* (the latter caused by a spirochete) (Chap. 208) may be

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Centrally Distributed Maculopapular Eruptions					
Acute meningococemia ^a	—	—	—	—	180
Drug reaction with eosinophilia and systemic symptoms (DRESS) (also termed drug-induced hypersensitivity syndrome; DIHS) ^b	—	—	—	—	74
Rubeola (measles, first disease)	Paramyxovirus	Discrete lesions that become confluent as rash spreads from hairline downward, usually sparing palms and soles; lasts ≥3 days; Koplik's spots	Nonimmune individuals	Cough, conjunctivitis, coryza, severe prostration	229
Rubella (German measles, third disease)	Togavirus	Spreads from hairline downward, clearing as it spreads; Forschheimer spots	Nonimmune individuals	Adenopathy, arthritis	230e
Erythema infectiosum (fifth disease)	Human parvovirus B19	Bright-red "slapped-cheeks" appearance followed by lacy reticular rash that waxes and wanes over 3 weeks; rarely, papular-purpuric "gloves-and-socks" syndrome on hands and feet	Most common among children 3–12 years old; occurs in winter and spring	Mild fever; arthritis in adults; rash following resolution of fever	221
Exanthem subitum (roseola, sixth disease)	Human herpesvirus 6	Diffuse maculopapular eruption over trunk and neck; resolves within 2 days	Usually affects children <3 years old	Rash following resolution of fever; similar to Boston exanthem (echovirus 16); febrile seizures may occur	219
Primary HIV infection	HIV	Nonspecific diffuse macules and papules; less commonly, urticarial or vesicular oral or genital ulcers	Individuals recently infected with HIV	Pharyngitis, adenopathy, arthralgias	226
Infectious mononucleosis	Epstein-Barr virus	Diffuse maculopapular eruption (5% of cases; 90% if ampicillin is given); urticaria, petechiae in some cases; periorbital edema (50%); palatal petechiae (25%)	Adolescents, young adults	Hepatosplenomegaly, pharyngitis, cervical lymphadenopathy, atypical lymphocytosis, heterophile antibody	218
Other viral exanthems	Echoviruses 2, 4, 9, 11, 16, 19, 25; coxsackieviruses A9, B1, B5; etc.	Wide range of skin findings that may mimic rubella or measles	Affect children more commonly than adults	Nonspecific viral syndromes	228
Exanthematous drug-induced eruption	Drugs (antibiotics, anticonvulsants, diuretics, etc.)	Intensely pruritic, bright-red macules and papules, symmetric on trunk and extremities; may become confluent	Occurs 2–3 days after exposure in previously sensitized individuals; otherwise, after 2–3 weeks (but can occur anytime, even shortly after drug is discontinued)	Variable findings: fever and eosinophilia	74
Epidemic typhus	<i>Rickettsia prowazekii</i>	Maculopapular eruption appearing in axillae, spreading to trunk and later to extremities; usually spares face, palms, soles; evolves from blanchable macules to confluent eruption with petechiae; rash evanescent in recrudescent typhus (Brill-Zinsser disease)	Exposure to body lice; occurrence of recrudescent typhus as relapse after 30–50 years	Headache, myalgias; mortality rates 10–40% if untreated; milder clinical presentation in recrudescent form	211
Endemic (murine) typhus	<i>Rickettsia typhi</i>	Maculopapular eruption, usually sparing palms, soles	Exposure to rat or cat fleas	Headache, myalgias	211
Scrub typhus	<i>Orientia tsutsugamushi</i>	Diffuse macular rash starting on trunk; eschar at site of mite bite	Endemic in South Pacific, Australia, Asia; transmitted by mites	Headache, myalgias; regional adenopathy; mortality rates up to 30% if untreated	211
Rickettsial spotted fevers	<i>Rickettsia conorii</i> (boutonneuse fever), <i>Rickettsia australis</i> (North Queensland tick typhus), <i>Rickettsia sibirica</i> (Siberian tick typhus), and others	Eschar common at bite site; maculopapular (rarely, vesicular and petechial) eruption on proximal extremities, spreading to trunk and face	Exposure to ticks; <i>R. conorii</i> in Mediterranean region, India, Africa; <i>R. australis</i> in Australia; <i>R. sibirica</i> in Siberia, Mongolia	Headache, myalgias, regional adenopathy	211

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Centrally Distributed Maculopapular Eruptions (Continued)					
Human monocytotropic ehrlichiosis ^c	<i>Ehrlichia chaffeensis</i>	Maculopapular eruption (40% of cases), involves trunk and extremities; may be petechial	Tick-borne; most common in U.S. Southeast, southern Midwest, and mid-Atlantic regions	Headache, myalgias, leukopenia	211
Leptospirosis	<i>Leptospira interrogans</i>	Maculopapular eruption; conjunctivitis; scleral hemorrhage in some cases	Exposure to water contaminated with animal urine	Myalgias; aseptic meningitis; <i>fulminant form</i> : icterohemorrhagic fever (Weil's disease)	208
Lyme disease	<i>Borrelia burgdorferi</i> (sole cause in U.S.), <i>Borrelia afzelii</i> , <i>Borrelia garinii</i>	Papule expanding to erythematous annular lesion with central clearing (erythema migrans; average diameter, 15 cm), sometimes with concentric rings, sometimes with indurated or vesicular center; multiple secondary erythema migrans lesions in some cases	Bite of <i>Ixodes</i> tick vector	Headache, myalgias, chills, photophobia occurring acutely; CNS disease, myocardial disease, arthritis weeks to months later in some cases	210
Southern tick-associated rash illness (STAR), Master's disease)	Unknown (possibly <i>Borrelia lonestari</i> or other <i>Borrelia</i> spirochetes)	Similar to erythema migrans of Lyme disease with several differences, including: multiple secondary lesions less likely; lesions tending to be smaller (average diameter, ~8 cm); central clearing more likely	Bite of tick vector <i>Amblyomma americanum</i> (Lone Star tick); often found in regions where Lyme disease is uncommon, including southern United States	Compared with Lyme disease: fewer constitutional symptoms, tick bite more likely to be recalled; other Lyme disease sequelae lacking	210
Typhoid fever	<i>Salmonella typhi</i>	Transient, blanchable erythematous macules and papules, 2–4 mm, usually on trunk (rose spots)	Ingestion of contaminated food or water (rare in U.S.)	Variable abdominal pain and diarrhea; headache, myalgias, hepatosplenomegaly	190
Dengue fever ^d	Dengue virus (4 serotypes; flaviviruses)	Rash in 50% of cases; initially diffuse flushing; midway through illness, onset of maculopapular rash, which begins on trunk and spreads centrifugally to extremities and face; pruritus, hyperesthesia in some cases; after defervescence, petechiae on extremities in some cases	Occurs in tropics and subtropics; transmitted by mosquito	Headache, musculoskeletal pain ("breakbone fever"); leukopenia; occasionally biphasic ("saddle-back") fever	233
Rat-bite fever (sodoku)	<i>Spirillum minus</i>	Eschar at bite site; then blotchy violaceous or red-brown rash involving trunk and extremities	Rat bite; primarily found in Asia; rare in U.S.	Regional adenopathy; recurrent fevers if untreated	167e
Relapsing fever	<i>Borrelia</i> species	Central rash at end of febrile episode; petechiae in some cases	Exposure to ticks or body lice	Recurrent fever, headache, myalgias, hepatosplenomegaly	209
Erythema marginatum (rheumatic fever)	Group A <i>Streptococcus</i>	Erythematous annular papules and plaques occurring as polycyclic lesions in waves over trunk, proximal extremities; evolving and resolving within hours	Patients with rheumatic fever	Pharyngitis preceding polyarthritis, carditis, subcutaneous nodules, chorea	381
Systemic lupus erythematosus (SLE)	Autoimmune disease	Macular and papular erythema, often in sun-exposed areas; discoid lupus lesions (local atrophy, scale, pigmentary changes); periungual telangiectasis; malar rash; vasculitis sometimes causing urticaria, palpable purpura; oral erosions in some cases	Most common in young to middle-aged women; flares precipitated by sun exposure	Arthritis; cardiac, pulmonary, renal, hematologic, and vasculitic disease	378
Still's disease	Autoimmune disease	Transient 2- to 5-mm erythematous papules appearing at height of fever on trunk, proximal extremities; lesions evanescent	Children and young adults	High spiking fever, polyarthritis, splenomegaly; erythrocyte sedimentation rate, >100 mm/h	398
African trypanosomiasis	<i>Trypanosoma brucei rhodesiense</i> / <i>gambiense</i>	Blotchy or annular erythematous macular and papular rash (trypanid), primarily on trunk; pruritus; chancre at site of tsetse fly bite may precede rash by several weeks	Tsetse fly bite in eastern (<i>T. brucei rhodesiense</i>) or western (<i>T. brucei gambiense</i>) Africa	Hemolymphatic disease followed by meningoencephalitis; Winterbottom's sign (posterior cervical lymphadenopathy) (<i>T. brucei gambiense</i>)	252
Arcanobacterial pharyngitis	<i>Arcanobacterium</i> (<i>Corynebacterium</i>) <i>haemolyticum</i>	Diffuse, erythematous, maculopapular eruption involving trunk and proximal extremities; may desquamate	Children and young adults	Exudative pharyngitis, lymphadenopathy	175
West Nile fever	West Nile virus	Maculopapular eruption involving the trunk, extremities, and head or neck; rash in 20–50% of cases	Mosquito bite; rarely, blood transfusion or transplanted organ	Headache, weakness, malaise, myalgias, neuroinvasive disease (encephalitis, meningitis, flaccid paralysis)	233

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Peripheral Eruptions					
Chronic meningococemia, disseminated gonococcal infection, ^a human parvovirus B19 infection ^e	—	—	—	—	180, 181, 221
Rocky Mountain spotted fever	<i>Rickettsia rickettsii</i>	Rash beginning on wrists and ankles and spreading centripetally; appears on palms and soles later in disease; lesion evolution from blanchable macules to petechiae	Tick vector; widespread but more common in southeastern and south-west-central U.S.	Headache, myalgias, abdominal pain; mortality rates up to 40% if untreated	211
Secondary syphilis	<i>Treponema pallidum</i>	Coincident primary chancre in 10% of cases; copper-colored, scaly papular eruption, diffuse but prominent on palms and soles; rash never vesicular in adults; condyloma latum, mucous patches, and alopecia in some cases	Sexually transmitted	Fever, constitutional symptoms	206
Chikungunya fever	Chikungunya virus	Maculopapular eruption; prominent on upper extremities and face, but can also occur on trunk and lower extremities	<i>Aedes aegypti</i> and <i>A. albopictus</i> mosquito bites; primarily in Africa and Indian Ocean region	Severe polyarticular, migratory arthralgias, especially involving small joints (e.g., hands, wrists, ankles)	233
Hand-foot-and-mouth disease	Coxsackievirus A16 most common cause	Tender vesicles, erosions in mouth; 0.25-cm papules on hands and feet with rim of erythema evolving into tender vesicles	Summer and fall; primarily children <10 years old; multiple family members	Transient fever	228
Erythema multiforme (EM)	Infection, drugs, idiopathic causes	Target lesions (central erythema surrounded by area of clearing and another rim of erythema) up to 2 cm; symmetric on knees, elbows, palms, soles; spreads centripetally; papular, sometimes vesicular; when extensive and involving mucous membranes, termed <i>EM major</i>	Herpes simplex virus or <i>Mycoplasma pneumoniae</i> infection; drug intake (i.e., sulfa, phenytoin, penicillin)	50% of patients <20 years old; fever more common in most severe form, EM major, which can be confused with Stevens-Johnson syndrome (but EM major lacks prominent skin sloughing)	— ^f
Rat-bite fever (Haverhill fever)	<i>Streptobacillus moniliformis</i>	Maculopapular eruption over palms, soles, and extremities; tends to be more severe at joints; eruption sometimes becoming generalized; may be purpuric; may desquamate	Rat bite, ingestion of contaminated food	Myalgias; arthritis (50%); fever recurrence in some cases	167e
Bacterial endocarditis	<i>Streptococcus</i> , <i>Staphylococcus</i> , etc.	<i>Subacute course</i> : Osler's nodes (tender pink nodules on finger or toe pads); petechiae on skin and mucosa; splinter hemorrhages. <i>Acute course</i> (e.g., <i>Staphylococcus aureus</i>): Janeway lesions (painless erythematous or hemorrhagic macules, usually on palms and soles)	Abnormal heart valve (e.g., viridans group streptococci), intravenous drug use	New or changing heart murmur	155
Confluent Desquamative Erythemas					
Scarlet fever (second disease)	Group A <i>Streptococcus</i> (pyrogenic exotoxins A, B, C)	Diffuse blanchable erythema beginning on face and spreading to trunk and extremities; circumoral pallor; "sandpaper" texture to skin; accentuation of linear erythema in skin folds (Pastia's lines); enanthem of white evolving into red "strawberry" tongue; desquamation in second week	Most common among children 2–10 years old; usually follows group A streptococcal pharyngitis	Fever, pharyngitis, headache	173
Kawasaki disease	Idiopathic causes	Rash similar to scarlet fever (scarlatiniform) or EM; fissuring of lips, strawberry tongue; conjunctivitis; edema of hands, feet; desquamation later in disease	Children <8 years old	Cervical adenopathy, pharyngitis, coronary artery vasculitis	72, 385
Streptococcal toxic shock syndrome	Group A <i>Streptococcus</i> (associated with pyrogenic exotoxin A and/or B or certain M types)	When present, rash often scarlatiniform	May occur in setting of severe group A streptococcal infections (e.g., necrotizing fasciitis, bacteremia, pneumonia)	Multorgan failure, hypotension; mortality rate 30%	173
Staphylococcal toxic shock syndrome	<i>S. aureus</i> (toxic shock syndrome toxin 1, enterotoxins B and others)	Diffuse erythema involving palms; pronounced erythema of mucosal surfaces; conjunctivitis; desquamation 7–10 days into illness	Colonization with toxin-producing <i>S. aureus</i>	Fever >39°C (>102°F), hypotension, multiorgan dysfunction	172

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Confluent Desquamative Erythemas (Continued)					
Staphylococcal scalded-skin syndrome	<i>S. aureus</i> , phage group II	Diffuse tender erythema, often with bullae and desquamation; Nikolsky's sign	Colonization with toxin-producing <i>S. aureus</i> ; occurs in children <10 years old (termed <i>Ritter's disease</i> in neonates) or adults with renal dysfunction	Irritability; nasal or conjunctival secretions	172
Exfoliative erythroderma syndrome	Underlying psoriasis, eczema, drug eruption, mycosis fungoides	Diffuse erythema (often scaling) interspersed with lesions of underlying condition	Usually occurs in adults over age 50; more common among men	Fever, chills (i.e., difficulty with thermoregulation); lymphadenopathy	72, 74
DRESS (DIHS)	Aromatic anticonvulsants; other drugs, including sulfonamides, minocycline	Maculopapular eruption (mimicking exanthematous drug rash), sometimes progressing to exfoliative erythroderma; profound edema, especially facial; pustules may occur	Individuals genetically unable to detoxify arene oxides (anticonvulsants), patients with slow <i>N</i> -acetylating capacity (sulfonamides)	Lymphadenopathy, multiorgan failure (especially hepatic), eosinophilia, atypical lymphocytes; mimics sepsis	74
Stevens-Johnson syndrome (SJS), toxic epidermal necrolysis (TEN)	Drugs (80% of cases; often allopurinol, anticonvulsants, antibiotics), infection, idiopathic	Erythematous and purpuric macules, sometimes targetoid, or diffuse erythema progressing to bullae, with sloughing and necrosis of entire epidermis; Nikolsky's sign; involves mucosal surfaces; TEN (>30% epidermal necrosis) is maximal form; SJS involves <10% of epidermis; SJS/TEN overlap involves 10–30% of epidermis	Uncommon among children; more common among patients with HIV infection, SLE, certain HLA types, or slow acetylators	Dehydration, sepsis sometimes resulting from lack of normal skin integrity; mortality rates up to 30%	74
Vesiculobullous or Pustular Eruptions					
Hand-foot-and-mouth syndrome ^a ; staphylococcal scalded-skin syndrome; TEN ^b ; DRESS ^b	—	—	—	—	— ^f
Varicella (chickenpox)	VZV	Macules (2–3 mm) evolving into papules, then vesicles (sometimes umbilicated), on an erythematous base ("dewdrops on a rose petal"); pustules then forming and crusting; lesions appearing in crops; may involve scalp, mouth; intensely pruritic	Usually affects children; 10% of adults susceptible; most common in late winter and spring; incidence down by 90% in U.S. as a result of varicella vaccination	Malaise; generally mild disease in healthy children; more severe disease with complications in adults and immunocompromised children	217
<i>Pseudomonas</i> "hot-tub" folliculitis	<i>Pseudomonas aeruginosa</i>	Pruritic erythematous follicular, papular, vesicular, or pustular lesions that may involve axillae, buttocks, abdomen, and especially areas occluded by bathing suits; can manifest as tender isolated nodules on palmar or plantar surfaces (the latter designated " <i>Pseudomonas</i> hot-foot syndrome")	Bathers in hot tubs or swimming pools; occurs in outbreaks	Earache, sore eyes and/or throat; fever may be absent; generally self-limited	189
Variola (smallpox)	Variola major virus	Red macules on tongue and palate evolving to papules and vesicles; skin macules evolving to papules, then vesicles, then pustules over 1 week, with subsequent lesion crusting; lesions initially appearing on face and spreading centrifugally from trunk to extremities; differs from varicella in that (1) skin lesions in any given area are at same stage of development and (2) there is a prominent distribution of lesions on face and extremities (including palms, soles)	Nonimmune individuals exposed to smallpox	Prodrome of fever, headache, backache, myalgias; vomiting in 50% of cases	261e
Primary herpes simplex virus (HSV) infection	HSV	Erythema rapidly followed by hallmark painful <i>grouped vesicles</i> that may evolve into pustules that ulcerate, especially on mucosal surfaces; lesions at site of inoculation: commonly gingivostomatitis for HSV-1 and genital lesions for HSV-2; recurrent disease milder (e.g., herpes labialis does not involve oral mucosa)	Primary infection most common among children and young adults for HSV-1 and among sexually active young adults for HSV-2; no fever in recurrent infection	Regional lymphadenopathy	216

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Vesiculobullous or Pustular Eruptions (Continued)					
Disseminated herpesvirus infection	Varicella-zoster virus (VZV) or HSV	Generalized vesicles that can evolve to pustules and ulcerations; individual lesions similar for VZV and HSV. <i>Zoster cutaneous dissemination</i> : >25 lesions extending outside involved dermatome. <i>HSV</i> : extensive, progressive mucocutaneous lesions that may occur in absence of dissemination, sometimes disseminate in eczematous skin (eczema herpeticum); HSV visceral dissemination may occur with only localized mucocutaneous disease; in disseminated neonatal disease, skin lesions diagnostically helpful when present, but rash absent in a substantial minority of cases	Patients with immunosuppression, eczema; neonates	Visceral organ involvement (e.g., liver, lungs) in some cases; neonatal disease particularly severe	164, 216, 217
Rickettsialpox	<i>Rickettsia akari</i>	Eschar found at site of mite bite; generalized rash involving face, trunk, extremities; may involve palms and soles; <100 papules and plaques (2–10 mm); tops of lesions developing vesicles that may evolve into pustules	Seen in urban settings; transmitted by mouse mites	Headache, myalgias, regional adenopathy; mild disease	211
Acute generalized eruptive pustulosis	Drugs (mostly anti-convulsants or antimicrobials); also viral	Tiny sterile nonfollicular pustules on erythematous, edematous skin; begins on face and in body folds, then becomes generalized	Appears 2–21 days after start of drug therapy, depending on whether patient has been sensitized	Acute fever, pruritus, leukocytosis	74
Disseminated <i>Vibrio vulnificus</i> infection	<i>V. vulnificus</i>	Erythematous lesions evolving into hemorrhagic bullae and then into necrotic ulcers	Patients with cirrhosis, diabetes, renal failure; exposure by ingestion of contaminated saltwater, seafood	Hypotension; mortality rate 50%	193
Ecthyma gangrenosum	<i>P. aeruginosa</i> , other gram-negative rods, fungi	Indurated plaque evolving into hemorrhagic bulla or pustule that sloughs, resulting in eschar formation; erythematous halo; most common in axillary, groin, perianal regions	Usually affects neutropenic patients; occurs in up to 28% of individuals with <i>Pseudomonas</i> bacteremia	Clinical signs of sepsis	189
Urticaria-Like Eruptions					
Urticarial vasculitis	Serum sickness, often due to infection (including hepatitis B viral, enteroviral, parasitic), drugs; connective tissue disease	Erythematous, edematous “urticaria-like” plaques, pruritic or burning; unlike urticaria: typical lesion duration >24 h (up to 5 days) and lack of complete lesion blanching with compression due to hemorrhage	Patients with serum sickness (including hepatitis B), connective tissue disease	Fever variable; arthralgias/ arthritis	385 ^f
Nodular Eruptions					
Disseminated infection	Fungal infections (e.g., candidiasis, histoplasmosis, cryptococcosis, sporotrichosis, coccidioidomycosis); mycobacteria	Subcutaneous nodules (up to 3 cm); fluctuance, draining common with mycobacteria; necrotic nodules (extremities, periorbital or nasal regions) common with <i>Aspergillus</i> , <i>Mucor</i>	Immunocompromised hosts (i.e., bone marrow transplant recipients, patients undergoing chemotherapy, HIV-infected patients, alcoholics)	Features vary with organism	— ^f
Erythema nodosum (septal panniculitis)	Infections (e.g., streptococcal, fungal, mycobacterial, yersinia); drugs (e.g., sulfas, penicillins, oral contraceptives); sarcoidosis; idiopathic causes	Large, violaceous, nonulcerative, subcutaneous nodules; exquisitely tender; usually on lower legs but also on upper extremities	More common among girls and women 15–30 years old	Arthralgias (50%); features vary with associated condition	— ^f
Sweet syndrome (acute febrile neutrophilic dermatosis)	Yersinia; upper respiratory infection; inflammatory bowel disease; pregnancy; malignancy (usually hematologic); drugs (G-CSF)	Tender red or blue edematous nodules giving impression of vesiculation; usually on face, neck, upper extremities; when on lower extremities, may mimic erythema nodosum	More common among women and among persons 30–60 years old; 20% of cases associated with malignancy (men and women equally affected in this group)	Headache, arthralgias, leukocytosis	72

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Nodular Eruption (Continued)					
Bacillary angiomatosis	<i>Bartonella henselae</i> , <i>B. quintana</i>	Many forms, including erythematous, smooth vascular nodules; friable, exophytic lesions; erythematous plaques (may be dry, scaly); subcutaneous nodules (may be erythematous)	Immunosuppressed individuals, especially those with advanced HIV infection	Peliosis of liver and spleen in some cases; lesions sometimes involving multiple organs; bacteremia	197
Purpuric Eruptions					
Rocky Mountain spotted fever, rat-bite fever, endocarditis ^a ; epidemic typhus ^a ; dengue fever ^{d,e} ; human parvovirus B19 infection ^e	—	—	—	—	— ^f
Acute meningococemia	<i>Neisseria meningitidis</i>	Initially pink maculopapular lesions evolving into petechiae; petechiae rapidly becoming numerous, sometimes enlarging and becoming vesicular; trunk, extremities most commonly involved; may appear on face, hands, feet; may include purpura fulminans (see below) reflecting DIC	Most common among children, individuals with asplenia or terminal complement deficiency (C5–C8)	Hypotension, meningitis (sometimes preceded by upper respiratory infection)	180
Purpura fulminans	Severe DIC	Large ecchymoses with sharply irregular shapes evolving into hemorrhagic bullae and then into black necrotic lesions	Individuals with sepsis (e.g., involving <i>N. meningitidis</i>), malignancy, or massive trauma; asplenic patients at high risk for sepsis	Hypotension	180, 325
Chronic meningococemia	<i>N. meningitidis</i>	Variety of recurrent eruptions, including pink maculopapular; nodular (usually on lower extremities); petechial (sometimes developing vesicular centers); purpuric areas with pale blue-gray centers	Individuals with complement deficiencies	Fevers, sometimes intermittent; arthritis, myalgias, headache	180
Disseminated gonococcal infection	<i>Neisseria gonorrhoeae</i>	Papules (1–5 mm) evolving over 1–2 days into hemorrhagic pustules with gray necrotic centers; hemorrhagic bullae occurring rarely; lesions (usually <40) distributed peripherally near joints (more commonly on upper extremities)	Sexually active individuals (more often females), some with complement deficiency	Low-grade fever, tenosynovitis, arthritis	181
Enteroviral petechial rash	Usually echovirus 9 or coxsackievirus A9	Disseminated petechial lesions (may also be maculopapular, vesicular, or urticarial)	Often occurs in outbreaks	Pharyngitis, headache; aseptic meningitis with echovirus 9	228
Viral hemorrhagic fever	Arboviruses (including dengue) and arenaviruses	Petechial rash	Residence in or travel to endemic areas, other virus exposure	Triad of fever, shock, hemorrhage from mucosa or gastrointestinal tract	233, 234
Thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome	Idiopathic, bloody diarrhea caused by Shiga toxin–generating bacteria (e.g., <i>Escherichia coli</i> O157:H7), deficiency in ADAMTS13 (cleaves von Willebrand factor), drugs (e.g., quinine, chemotherapy, immunosuppression)	Petechiae	Individuals with <i>E. coli</i> O157:H7 gastroenteritis (especially children), cancer chemotherapy, HIV infection, autoimmune diseases, pregnant/postpartum women	Fever (not always present), microangiopathic hemolytic anemia, thrombocytopenia, renal dysfunction, neurologic dysfunction; coagulation studies normal	72, 129, 140, 186, 191
Cutaneous small-vessel vasculitis (leukocytoclastic vasculitis)	Infections (including that caused by group A <i>Streptococcus</i> , viral hepatitis), drugs, idiopathic causes	Palpable purpuric lesions appearing in crops on legs or other dependent areas; may become vesicular or ulcerative	Occurs in a wide spectrum of diseases, including connective tissue disease, cryoglobulinemia, malignancy, Henoch-Schönlein purpura (HSP); more common among children	Fever (not always present), malaise, arthralgias, myalgias; systemic vasculitis in some cases; renal, joint, and gastrointestinal involvement common in HSP	72
Eruptions with Ulcers and/or Eschars					
Scrub typhus, rickettsial spotted fevers, rat-bite fever ^a ; rickettsialpox, ecthyma gangrenosum ^b	—	—	—	—	— ^f

(Continued)

TABLE 24-1 DISEASES ASSOCIATED WITH FEVER AND RASH (CONTINUED)

Disease	Etiology	Description	Group Affected/ Epidemiologic Factors	Clinical Syndrome	Chapter
Eruptions with Ulcers and/or Eschars (Continued)					
Tularemia	<i>Francisella tularensis</i>	Ulceroglandular form: erythematous, tender papule evolves into necrotic, tender ulcer with raised borders; in 35% of cases, eruptions (maculopapular, vesiculopapular, acneiform, or urticarial; erythema nodosum; or EM) may occur	Exposure to ticks, biting flies, infected animals	Fever, headache, lymphadenopathy	195
Anthrax	<i>Bacillus anthracis</i>	Pruritic papule enlarging and evolving into a 1- by 3-cm painless ulcer surrounded by vesicles and then developing a central eschar with edema; residual scar	Exposure to infected animals or animal products, other exposure to anthrax spores	Lymphadenopathy, headache	261e

*See "Purpuric Eruptions." †See "Confluent Desquamative Erythemas." ‡In human granulocytotropic ehrlichiosis or anaplasmosis (caused by *Anaplasma phagocytophila*; most common in the upper midwestern and northeastern United States), rash is rare. §See "Viral hemorrhagic fever" under "Purpuric Eruptions" for dengue hemorrhagic fever/dengue shock syndrome. ¶See "Centrally Distributed Maculopapular Eruptions." ††See etiology-specific chapters. †††See "Peripheral Eruptions." ††††See "Vesiculobullous or Pustular Eruptions."

Abbreviations: CNS, central nervous system; DIC, disseminated intravascular coagulation; G-CSF, granulocyte colony-stimulating factor; HLA, human leukocyte antigen.

seen in urban environments where rodents proliferate. Outside the United States, other rickettsial diseases cause a spotted-fever syndrome and should be considered in residents of or travelers to endemic areas. Similarly, *typhoid fever*, a nonrickettsial disease caused by *Salmonella typhi* (Chap. 190), is usually acquired during travel outside the United States. *Dengue fever*, caused by a mosquito-transmitted flavivirus, occurs in tropical and subtropical regions of the world (Chap. 233).

Some centrally distributed maculopapular eruptions have distinctive features. Erythema migrans, the rash of *Lyme disease* (Chap. 210), typically manifests as single or multiple annular plaques. Untreated erythema migrans lesions usually fade within a month but may persist for more than a year. *Southern tick-associated rash illness* (STARI) (Chap. 210) has an erythema migrans–like rash but is less severe than *Lyme disease* and often occurs in regions where *Lyme* is not endemic. *Erythema marginatum*, the rash of *acute rheumatic fever* (Chap. 381), has a distinctive pattern of enlarging and shifting transient annular lesions.

Collagen vascular diseases may cause fever and rash. Patients with *systemic lupus erythematosus* (Chap. 378) typically develop a sharply defined, erythematous eruption in a butterfly distribution on the cheeks (malar rash) as well as many other skin manifestations. *Still's disease* (Chap. 398) presents as an evanescent, salmon-colored rash on the trunk and proximal extremities that coincides with fever spikes.

PERIPHERAL ERUPTIONS

These rashes are alike in that they are most prominent peripherally or begin in peripheral (acral) areas before spreading centripetally. Early diagnosis and therapy are critical in *Rocky Mountain spotted fever* (Chap. 211) because of its grave prognosis if untreated. Lesions evolve from macular to petechial, start on the wrists and ankles, spread centripetally, and appear on the palms and soles only later in the disease. The rash of *secondary syphilis* (Chap. 206), which may be generalized but is prominent on the palms and soles, should be considered in the differential diagnosis of pityriasis rosea, especially in sexually active patients. *Chikungunya fever* (Chap. 233), which is transmitted by mosquito bite in Africa and the Indian Ocean region, is associated with a maculopapular eruption and severe polyarticular small-joint arthralgias. *Hand-foot-and-mouth disease* (Chap. 228), most commonly caused by coxsackievirus A16, is distinguished by tender vesicles distributed peripherally and in the mouth; outbreaks commonly occur within families. The classic target lesions of *erythema multiforme* appear symmetrically on the elbows, knees, palms, soles, and face. In severe cases, these lesions spread diffusely and involve mucosal surfaces. Lesions may develop on the hands and feet in *endocarditis* (Chap. 155).

CONFLUENT DESQUAMATIVE ERYTHEMAS

These eruptions consist of diffuse erythema frequently followed by desquamation. The eruptions caused by group A *Streptococcus* or

Staphylococcus aureus are toxin-mediated. *Scarlet fever* (Chap. 173) usually follows pharyngitis; patients have a facial flush, a "strawberry" tongue, and accentuated petechiae in body folds (Pastia's lines). *Kawasaki disease* (Chaps. 72 and 385) presents in the pediatric population as fissuring of the lips, a strawberry tongue, conjunctivitis, adenopathy, and sometimes cardiac abnormalities. *Streptococcal toxic shock syndrome* (Chap. 173) manifests with hypotension, multiorgan failure, and, often, a severe group A streptococcal infection (e.g., necrotizing fasciitis). *Staphylococcal toxic shock syndrome* (Chap. 172) also presents with hypotension and multiorgan failure, but usually only *S. aureus* colonization—not a severe *S. aureus* infection—is documented. *Staphylococcal scalded-skin syndrome* (Chap. 172) is seen primarily in children and in immunocompromised adults. Generalized erythema is often evident during the prodrome of fever and malaise; profound tenderness of the skin is distinctive. In the exfoliative stage, the skin can be induced to form bullae with light lateral pressure (Nikolsky's sign). In a mild form, a scarlatiniform eruption mimics scarlet fever, but the patient does not exhibit a strawberry tongue or circumoral pallor. In contrast to the staphylococcal scalded-skin syndrome, in which the cleavage plane is superficial in the epidermis, *toxic epidermal necrolysis* (Chap. 74), a maximal variant of *Stevens-Johnson syndrome*, involves sloughing of the entire epidermis, resulting in severe disease. *Exfoliative erythroderma syndrome* (Chaps. 72 and 74) is a serious reaction associated with systemic toxicity that is often due to eczema, psoriasis, a drug reaction, or mycosis fungoides. *Drug rash with eosinophilia and systemic symptoms* (DRESS), often due to antiepileptic and antibiotic agents (Chap. 74), initially appears similar to an exanthematous drug reaction but may progress to exfoliative erythroderma; it is accompanied by multi-organ failure and has an associated mortality rate of ~10%.

VESICULOBULLOUS OR PUSTULAR ERUPTIONS

Varicella (Chap. 217) is highly contagious, often occurring in winter or spring. At any point in time, within a given region of the body, varicella lesions are in different stages of development. In immunocompromised hosts, varicella vesicles may lack the characteristic erythematous base or may appear hemorrhagic. Lesions of *Pseudomonas "hot-tub" folliculitis* (Chap. 189) are also pruritic and may appear similar to those of varicella. However, hot-tub folliculitis generally occurs in outbreaks after bathing in hot tubs or swimming pools, and lesions occur in regions occluded by bathing suits. Lesions of *variola (smallpox)* (Chap. 261e) also appear similar to those of varicella but are all at the same stage of development in a given region of the body. Variola lesions are most prominent on the face and extremities, while varicella lesions are most prominent on the trunk. *Herpes simplex virus infection* (Chap. 216) is characterized by hallmark grouped vesicles on an erythematous base. Primary herpes infection is accompanied by fever and toxicity, while recurrent disease is milder. *Rickettsialpox* (Chap. 211) is often documented in urban settings and is characterized

by vesicles followed by pustules. It can be distinguished from varicella by an eschar at the site of the mouse-mite bite and the papule/plaque base of each vesicle. *Acute generalized eruptive pustulosis* should be considered in individuals who are acutely febrile and are taking new medications, especially anticonvulsant or antimicrobial agents (Chap. 74). Disseminated *Vibrio vulnificus* infection (Chap. 193) or *ecthyma gangrenosum* due to *Pseudomonas aeruginosa* (Chap. 189) should be considered in immunosuppressed individuals with sepsis and hemorrhagic bullae.

URTICARIA-LIKE ERUPTIONS

Individuals with classic urticaria (“hives”) usually have a hypersensitivity reaction without associated fever. In the presence of fever, urticaria-like eruptions are most often due to *urticarial vasculitis* (Chap. 385). Unlike individual lesions of classic urticaria, which last up to 24 h, these lesions may last 3–5 days. Etiologies include serum sickness (often induced by drugs such as penicillins, sulfas, salicylates, or barbiturates), connective-tissue disease (e.g., systemic lupus erythematosus or Sjögren’s syndrome), and infection (e.g., with hepatitis B virus, enteroviruses, or parasites). Malignancy, especially lymphoma, may be associated with fever and chronic urticaria (Chap. 72).

NODULAR ERUPTIONS

In immunocompromised hosts, nodular lesions often represent disseminated infection. Patients with disseminated *candidiasis* (often due to *Candida tropicalis*) may have a triad of fever, myalgias, and eruptive nodules (Chap. 240). Disseminated *cryptococcosis* lesions (Chap. 239) may resemble molluscum contagiosum (Chap. 220e). Necrosis of nodules should raise the suspicion of *aspergillosis* (Chap. 241) or *mucormycosis* (Chap. 242). *Erythema nodosum* presents with exquisitely tender nodules on the lower extremities. *Sweet syndrome* (Chap. 72) should be considered in individuals with multiple nodules and plaques, often so edematous that they give the appearance of vesicles or bullae. Sweet syndrome may occur in individuals with infection, inflammatory bowel disease, or malignancy and can also be induced by drugs.

PURPURIC ERUPTIONS

Acute meningococemia (Chap. 180) classically presents in children as a petechial eruption, but initial lesions may appear as blanchable macules or urticaria. Rocky Mountain spotted fever should be considered in the differential diagnosis of acute meningococemia. *Echovirus 9 infection* (Chap. 228) may mimic acute meningococemia; patients should be treated as if they have bacterial sepsis because prompt differentiation of these conditions may be impossible. Large ecchymotic areas of *purpura fulminans* (Chaps. 180 and 325) reflect severe underlying disseminated intravascular coagulation, which may be due to infectious or noninfectious causes. The lesions of *chronic meningococemia* (Chap. 180) may have a variety of morphologies, including petechial. Purpuric nodules may develop on the legs and resemble erythema nodosum but lack its exquisite tenderness. Lesions of *disseminated gonococemia* (Chap. 181) are distinctive, sparse, countable hemorrhagic pustules, usually located near joints. The lesions of chronic meningococemia and those of gonococemia may be indistinguishable in terms of appearance and distribution. *Viral hemorrhagic fever* (Chaps. 233 and 234) should be considered in patients with an appropriate travel history and a petechial rash. *Thrombotic thrombocytopenic purpura* (Chaps. 72, 129, and 140) and *hemolytic-uremic syndrome* (Chaps. 140, 186, and 191) are closely related and are noninfectious causes of fever and petechiae. *Cutaneous small-vessel vasculitis* (*leukocytoclastic vasculitis*) typically manifests as palpable purpura and has a wide variety of causes (Chap. 72).

ERUPTIONS WITH ULCERS OR ESCHARS

The presence of an ulcer or eschar in the setting of a more widespread eruption can provide an important diagnostic clue. For example, the presence of an eschar may suggest the diagnosis of *scrub typhus* or *rickettsialpox* (Chap. 211) in the appropriate setting. In other illnesses (e.g., anthrax) (Chap. 261e), an ulcer or eschar may be the only skin manifestation.

Given the extremely broad differential diagnosis, the presentation of a patient with fever and rash often poses a thorny diagnostic challenge for even the most astute and experienced clinician. Rapid narrowing of the differential by prompt recognition of a rash's key features can result in appropriate and sometimes life-saving therapy. This atlas presents high-quality images of a variety of rashes that have an infectious etiology and are commonly associated with fever.



A



B

FIGURE 25e-1 **A.** Erythema leading to “slapped cheeks” appearance in erythema infectiosum (fifth disease) caused by parvovirus B19. **B.** Lacy reticular rash of erythema infectiosum. (Panel A reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)

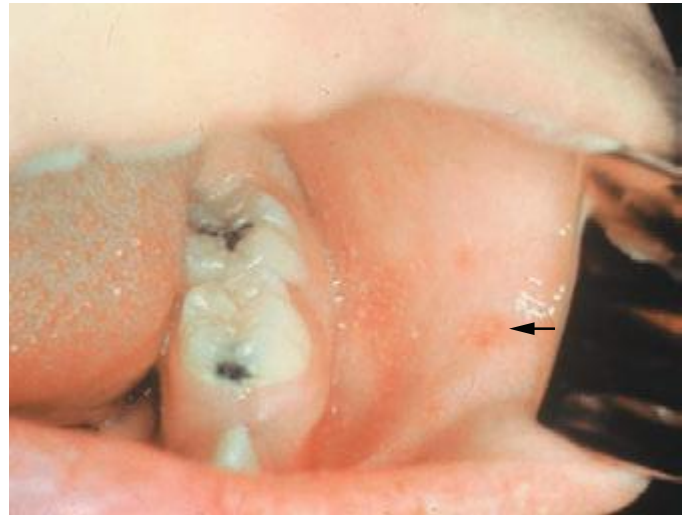


FIGURE 25e-2 Koplik's spots, which manifest as white or bluish lesions with an erythematous halo on the buccal mucosa, usually occur in the first 2 days of measles symptoms and may briefly overlap the measles exanthem. The presence of the erythematous halo (*arrow indicates one example*) differentiates Koplik's spots from Fordyce's spots (ectopic sebaceous glands), which occur in the mouths of healthy individuals. (Courtesy of the Centers for Disease Control and Prevention.)



FIGURE 25e-3 In measles, discrete erythematous lesions become confluent on the face and neck over 2–3 days as the rash spreads downward to the trunk and arms, where lesions remain discrete. (Reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.)



FIGURE 25e-4 In rubella, an erythematous exanthem spreads from the hairline downward and clears as it spreads. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-7 This exanthematous, drug-induced eruption consists of brightly erythematous macules and papules, some of which are confluent, distributed symmetrically on the trunk and extremities. Ampicillin caused this rash. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.)



FIGURE 25e-5 Exanthem subitum (roseola) occurs most commonly in young children. A diffuse maculopapular exanthem follows resolution of fever. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-6 Erythematous macules and papules are apparent on the trunk and arm of this patient with primary HIV infection. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.)



FIGURE 25e-8 Erythema migrans is the early cutaneous manifestation of Lyme disease and is characterized by erythematous annular patches, often with a central erythematous focus at the tick-bite site. (Reprinted from RP Usatine et al: *Color Atlas of Family Medicine*, 2nd ed. New York, McGraw-Hill, 2013. Courtesy of Thomas Corson, MD.)



FIGURE 25e-9 Rose spots are evident as erythematous macules on the trunk of this patient with typhoid fever. (Courtesy of the Centers for Disease Control and Prevention.)



FIGURE 25e-12 Chronic discoid lupus erythematosus. Violaceous, hyperpigmented, atrophic plaques, often with evidence of follicular plugging (which may result in scarring), are characteristic of this cutaneous form of lupus. (Reprinted from K Wolff, RA Johnson, AP Saavedra: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 7th ed. New York, McGraw-Hill, 2013.)



FIGURE 25e-10 Systemic lupus erythematosus showing prominent malar erythema and minimal scaling. Involvement of other sun-exposed sites is also common. (Reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-11 Subacute lupus erythematosus on the upper chest, with brightly erythematous and slightly edematous coalescent papules and plaques. (Reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-13 The rash of Still's disease typically exhibits evanescent, erythematous papules that appear at the height of fever on the trunk and proximal extremities. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-14 Impetigo is a superficial group A streptococcal or *Staphylococcus aureus* infection consisting of honey-colored crusts and erythematous weeping erosions. (Reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-15 Erysipelas is a group A streptococcal infection of the superficial dermis and consists of well-demarcated, erythematous, edematous, warm plaques. (Reprinted from K Wolff, RA Johnson, AP Saavedra: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 7th ed. New York, McGraw-Hill, 2013.)

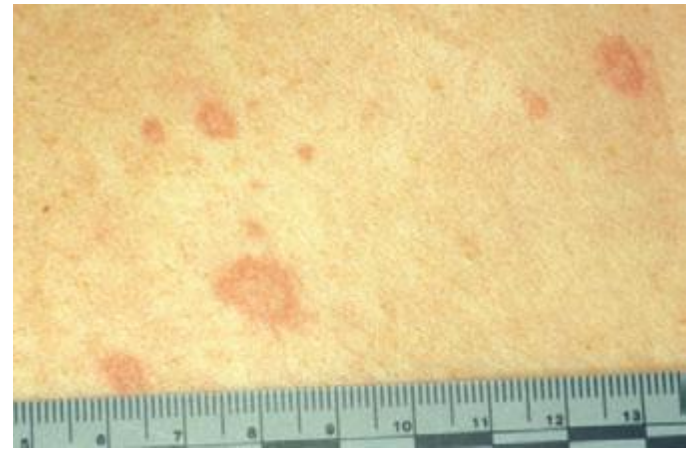


FIGURE 25e-16 **Top:** Petechial lesions of Rocky Mountain spotted fever on the lower legs and soles of a young, otherwise healthy patient. **Bottom:** Close-up of lesions from the same patient. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-17 Primary syphilis with firm, nontender chancres. (Courtesy of M. Rein and the Centers for Disease Control and Prevention.)



FIGURE 25e-18 Secondary syphilis, demonstrating the papulosquamous truncal eruption.



FIGURE 25e-21 Mucous patches on the tongue of a patient with secondary syphilis. (Courtesy of Ron Roddy; with permission.)



FIGURE 25e-19 Secondary syphilis commonly affects the palms and soles with scaling, firm, red-brown papules.



FIGURE 25e-22 Petechial lesions in a patient with atypical measles. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-20 Condylomata lata are moist, somewhat verrucous intertriginous plaques seen in secondary syphilis.

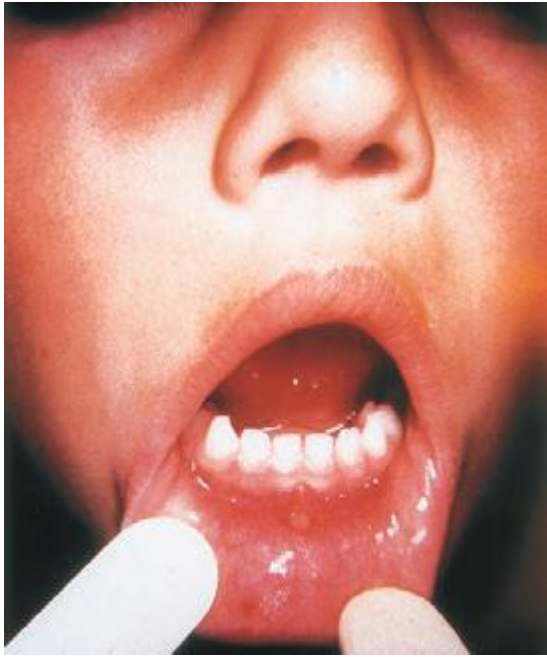


FIGURE 25e-23 Tender vesicles and erosions in the mouth of a patient with hand-foot-and-mouth disease. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-24 Septic emboli with hemorrhage and infarction due to acute *Staphylococcus aureus* endocarditis. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-25 Erythema multiforme is characterized by erythematous plaques with a target or iris morphology, sometimes with a vesicle in the center. It usually represents a hypersensitivity reaction to infections (especially herpes simplex virus or *Mycoplasma pneumoniae*) or drugs. (Reprinted from K Wolff, RA Johnson: *Fitzpatrick's Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-26 Scarlet fever exanthem. Finely punctuated erythema has become confluent (scarlatiniform); accentuation of linear erythema in body folds (Pastia's lines) is seen here. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-27 Erythema progressing to bullae with resulting sloughing of the entire thickness of the epidermis occurs in toxic epidermal necrolysis. This reaction was due to a sulfonamide. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.)



FIGURE 25e-29 This infant with staphylococcal scalded skin syndrome demonstrates generalized desquamation. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)

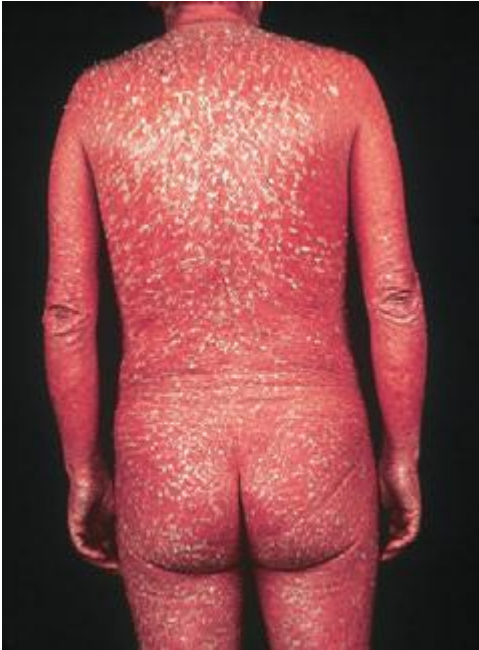


FIGURE 25e-28 Diffuse erythema and scaling are present in this patient with psoriasis and the exfoliative erythroderma syndrome. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-30 Fissuring of the lips and an erythematous exanthem are evident in this patient with Kawasaki disease. (Courtesy of Stephen E. Gellis, MD; with permission.)



FIGURE 25e-31 Numerous varicella lesions at various stages of evolution: vesicles on an erythematous base and umbilicated vesicles, which then develop into crusting lesions. (Courtesy of the Centers for Disease Control and Prevention.)



FIGURE 25e-32 Lesions of disseminated zoster at different stages of evolution, including pustules and crusting, similar to varicella. Note nongrouping of lesions, in contrast to herpes simplex or zoster. (Reprinted from K Wolff, RA Johnson, AP Saavedra: *Color Atlas and Synopsis of Clinical Dermatology*, 7th ed. New York, McGraw-Hill, 2013.)



A



B

FIGURE 25e-33 Herpes zoster is seen in this patient taking prednisone. Grouped vesicles and crusted lesions are seen in the T2 dermatome on the back and arm (**A**) and on the right side of the chest (**B**). (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-34 *Top:* Eschar at the site of the mite bite in a patient with rickettsialpox. *Middle:* Papulovesicular lesions on the trunk of the same patient. *Bottom:* Close-up of lesions from the same patient. (Reprinted from A Krusell et al: *Emerg Infect Dis* 8:727, 2002.)



FIGURE 25e-35 Ecthyma gangrenosum in a neutropenic patient with *Pseudomonas aeruginosa* bacteremia.



FIGURE 25e-36 Urticaria showing characteristic discrete and confluent, edematous, erythematous papules and plaques. (Reprinted from K Wolff, RA Johnson, AP Saavedra: *Color Atlas and Synopsis of Clinical Dermatology*, 7th ed. New York, McGraw-Hill, 2013.)



FIGURE 25e-37 Disseminated cryptococcal infection. A liver transplant recipient developed six cutaneous lesions similar to the one shown. Biopsy and serum antigen testing demonstrated *Cryptococcus*. Important features of the lesion include a benign-appearing fleshy papule with central umbilication resembling molluscum contagiosum. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-38 Disseminated candidiasis. Tender, erythematous, nodular lesions developed in a neutropenic patient with leukemia who was undergoing induction chemotherapy. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-39 Disseminated *Aspergillus* infection. Multiple necrotic lesions developed in this neutropenic patient undergoing hematopoietic stem cell transplantation. The lesion in the photograph is on the inner thigh and is several centimeters in diameter. Biopsy demonstrated infarction caused by *Aspergillus fumigatus*. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-40 Erythema nodosum is a panniculitis characterized by tender, deep-seated nodules and plaques usually located on the lower extremities. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 25e-41 Sweet syndrome is an erythematous indurated plaque with a pseudovesicular border. (Courtesy of Robert Swerlick, MD; with permission.)

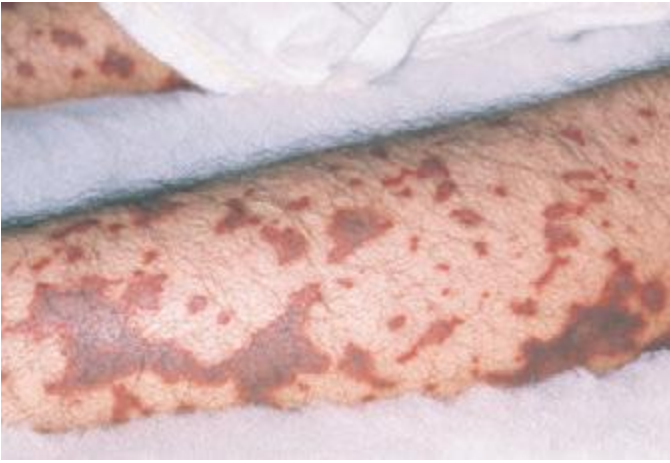


FIGURE 25e-42 Fulminant meningococemia with extensive angular purpuric patches. (Courtesy of Stephen E. Gellis, MD; with permission.)

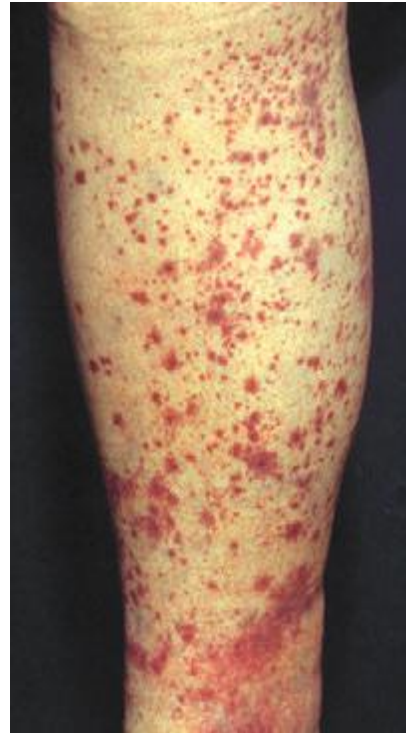


FIGURE 25e-45 Palpable purpuric papules on the lower leg are seen in this patient with cutaneous small-vessel hypersensitivity vasculitis. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)



FIGURE 25e-43 Erythematous papular lesions are seen on the leg of this patient with chronic meningococemia (arrow indicates a lesion).



FIGURE 25e-46 The thumb of a patient with a necrotic ulcer of tularemia. (Courtesy of the Centers for Disease Control and Prevention.)

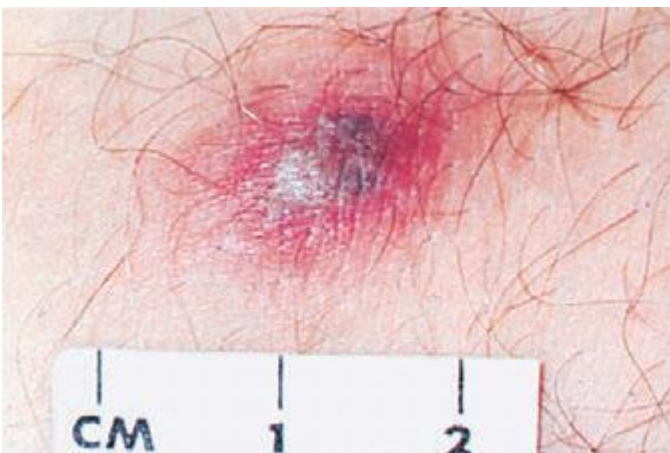


FIGURE 25e-44 Disseminated gonococemia in the skin is seen as hemorrhagic papules and pustules with purpuric centers in a centrifugal distribution. (Courtesy of Daniel M. Musher, MD; with permission.)



FIGURE 25e-47 This 50-year-old man developed high fever and massive inguinal lymphadenopathy after a small ulcer healed on his foot. Tularemia was diagnosed. (Courtesy of Lindsey Baden, MD; with permission.)



FIGURE 25e-48 This painful trypanosomal chancre developed at the site of a tsetse fly bite on the dorsum of the foot. *Trypanosoma brucei* was diagnosed from an aspirate of the ulcer. (Courtesy of Edward T. Ryan, MD. *N Engl J Med* 346:2069, 2002; with permission.)



FIGURE 25e-49 Drug reaction with eosinophilia and systemic symptoms/drug-induced hypersensitivity syndrome (DRESS/DIHS). This patient developed a progressive eruption exhibiting early desquamation after taking phenobarbital. There was also associated lymphadenopathy and hepatomegaly. (Courtesy of Peter Lio, MD; with permission.)



FIGURE 25e-50 Many small, nonfollicular pustules are seen against a background of erythema in this patient with acute generalized eruptive pustulosis (AGEP). The rash began in body folds and progressed to cover the trunk and face. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)

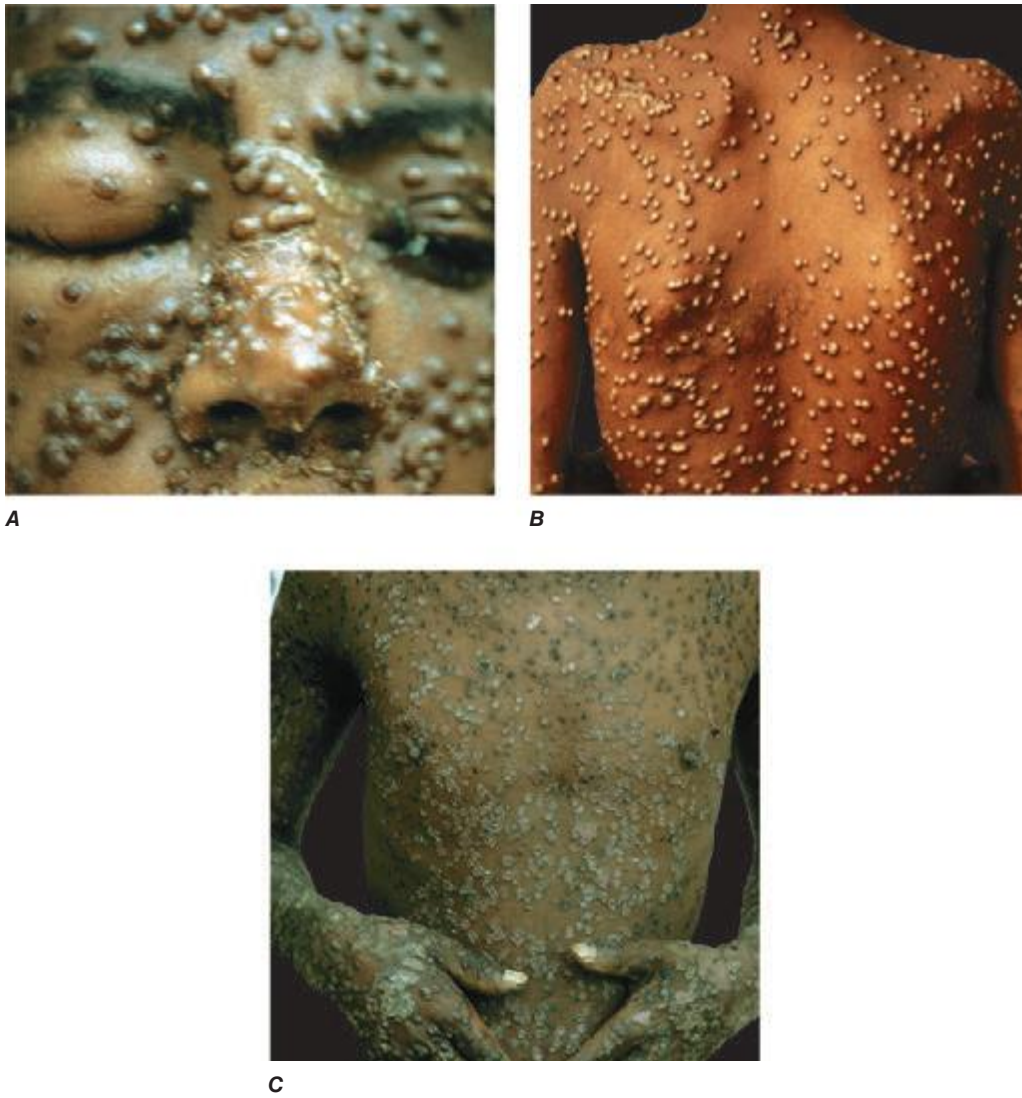


FIGURE 25e-51 Smallpox is shown with many pustules on the face, becoming confluent (**A**), and on the trunk (**B**). Pustules are all in the same stage of development. **C**. Crusting, healing lesions are noted on the trunk, arms, and hands. (Reprinted from K Wolff, RA Johnson: *Color Atlas and Synopsis of Clinical Dermatology*, 6th ed. New York, McGraw-Hill, 2009.)

This page intentionally left blank

26

Fever of Unknown Origin

Chantal P. Bleeker-Rovers, Jos W. M. van der Meer

DEFINITION

Clinicians commonly refer to any febrile illness without an initially obvious etiology as *fever of unknown origin* (FUO). Most febrile illnesses either resolve before a diagnosis can be made or develop distinguishing characteristics that lead to a diagnosis. The term *FUO* should be reserved for prolonged febrile illnesses without an established etiology despite intensive evaluation and diagnostic testing. This chapter focuses on classic FUO in the adult patient.

FUO was originally defined by Petersdorf and Beeson in 1961 as an illness of >3 weeks' duration with fever of $\geq 38.3^{\circ}\text{C}$ (101°F) on two occasions and an uncertain diagnosis despite 1 week of inpatient evaluation. Nowadays, most patients with FUO are hospitalized if their clinical condition requires it, but not for diagnostic purposes only; thus the in-hospital evaluation requirement has been eliminated from the definition. The definition of FUO has been further modified by the exclusion of immunocompromised patients, whose workup requires an entirely different diagnostic and therapeutic approach. For the optimal comparison of patients with FUO in different geographic areas, it has been proposed that the quantitative criterion (diagnosis uncertain after 1 week of evaluation) be changed to a qualitative criterion that requires the performance of a specific list of investigations. Accordingly, FUO is now defined as:

1. Fever $>38.3^{\circ}\text{C}$ (101°F) on at least two occasions
2. Illness duration of ≥ 3 weeks
3. No known immunocompromised state
4. Diagnosis that remains uncertain after a thorough history-taking, physical examination, and the following obligatory investigations: determination of erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP) level; platelet count; leukocyte count and differential; measurement of levels of hemoglobin, electrolytes, creatinine, total protein, alkaline phosphatase, alanine aminotransferase, aspartate aminotransferase, lactate dehydrogenase, creatine kinase, ferritin, antinuclear antibodies, and rheumatoid factor; protein electrophoresis; urinalysis; blood cultures ($n = 3$); urine culture; chest x-ray; abdominal ultrasonography; and tuberculin skin test (TST).


ETIOLOGY AND EPIDEMIOLOGY

The range of FUO etiologies has evolved over time as a result of changes in the spectrum of diseases causing FUO, the widespread

TABLE 26-1 ETIOLOGY OF FEVER OF UNKNOWN ORIGIN (FUO) OVER THE PAST 20 YEARS: FINDINGS FROM LARGE FUO STUDIES

First Author (Country, Year of Publication)	No. of Patients (Recruitment Period)	Percentage of Cases Due to Indicated Cause				
		Infections	Noninfectious Inflammatory Diseases	Neoplasms	Miscellaneous	Unknown
Western Countries						
De Kleijn et al. (Netherlands, 1997)	167 (1992–1994)	26	24	13	8	30
Vanderschueren et al. (Belgium, 2003)	185 (1990–1999)	11	18	10	8	53
Zenone et al. (France, 2006)	144 (1999–2005)	23	26	10	15	26
Bleeker-Rovers (Netherlands, 2007)	73 (2003–2005)	16	22	7	4	51
Mansueto et al. (Italy, 2008)	91 (1991–2002)	32	12	14	10	32
Efstathiou et al. (Greece, 2010)	112 (2001–2007)	30	33	11	5	21
Total	772	22	23	11	9	36
Other Geographic Locations						
Tabak et al. (Turkey, 2003)	117 (1984–2001)	34	29	19	4	14
Saltoglu et al. (Turkey, 2004)	87 (1994–2002)	59	18	14	2	7
Ergonul et al. (Turkey, 2005)	80 (1993–1999)	52	16	18	3	11
Chin et al. (Taiwan, 2006)	94 (2001–2002)	57	7	9	9	18
Colpan et al. (Turkey, 2007)	71 (2001–2004)	45	27	14	6	9
Hu et al. (China, 2008)	142 (2002–2003)	36	32	13	5	14
Kucukardali et al. (Turkey, 2008)	154 (2003–2004)	34	31	14	5	16
Ali-Eldin et al. (Egypt, 2011)	93 (2009–2010)	42	15	30	0	12
Total	838	43	23	16	4	13

use of antibiotics, and the availability of new diagnostic techniques. The proportion of cases caused by intraabdominal abscesses and tumors, for example, has decreased because of earlier detection by CT and ultrasound. In addition, infective endocarditis is a less frequent cause because blood culture and echocardiographic techniques have improved. Conversely, some diagnoses, such as acute HIV infection, were unknown four decades ago.

 **Table 26-1** summarizes the findings of several large studies on FUO conducted over the past 20 years. In general, infection accounts for about 20–25% of cases of FUO in Western countries; next in frequency are neoplasms and noninfectious inflammatory diseases (NIIDs), the latter including “collagen or rheumatic diseases,” vasculitis syndromes, and granulomatous disorders. In geographic areas outside the West, infections are a much more common cause of FUO (43% vs 22%), while the proportions of cases due to NIIDs and neoplasms are similar. Up to 50% of cases caused by infections in patients with FUO outside Western nations are due to tuberculosis, which is a less common cause in the United States and Western Europe. The number of FUO patients diagnosed with NIIDs probably will not decrease in the near future, as fever may precede more typical manifestations or serologic evidence by months in these diseases. Moreover, many NIIDs can be diagnosed only after prolonged observation and exclusion of other diseases.

In the West, the percentage of undiagnosed cases of FUO has increased in more recent studies. An important factor contributing to

the seemingly high diagnostic failure rate is that a diagnosis is more often being established before 3 weeks have elapsed, given that patients with fever tend to seek medical advice earlier and better diagnostic techniques, such as CT and MRI, are widely available; therefore, only the cases that are more difficult to diagnose continue to meet the criteria for FUO. Furthermore, most patients who have FUO without a diagnosis currently do well, and thus a less aggressive diagnostic approach may be used in clinically stable patients once diseases with immediate therapeutic or prognostic consequences have been ruled out to a reasonable extent. This factor may be especially relevant to patients with recurrent fever who are asymptomatic in between febrile episodes. In patients with recurrent fever (defined as repeated episodes of fever interspersed with fever-free intervals of at least 2 weeks and apparent remission of the underlying disease), the chance of attaining an etiologic diagnosis is <50%.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis for FUO is extensive, but it is important to remember that FUO is far more often caused by an atypical presentation of a rather common disease than by a very rare disease. **Table 26-2** presents an overview of possible causes of FUO. An atypical presentation of endocarditis, diverticulitis, vertebral osteomyelitis, and extrapulmonary tuberculosis are the more common infectious disease diagnoses. Q fever and Whipple’s disease are quite rare but should always be kept in mind as a cause of FUO since the presenting symptoms

TABLE 26-2 ALL REPORTED CAUSES OF FUO^a

Infections	
Bacterial, nonspecific	Abdominal abscess, adnexitis, apical granuloma, appendicitis, cholangitis, cholecystitis, diverticulitis, endocarditis, endometritis, epidural abscess, infected vascular catheter, infected joint prosthesis, infected vascular prosthesis, infectious arthritis, infective myonecrosis, intracranial abscess, liver abscess, lung abscess, malakoplakia, mastoiditis, mediastinitis, mycotic aneurysm, osteomyelitis, pelvic inflammatory disease, prostatitis, pyelonephritis, pylephlebitis, renal abscess, septic phlebitis, sinusitis, spondylodiscitis, xanthogranulomatous urinary tract infection
Bacterial, specific	Actinomycosis, atypical mycobacterial infection, bartonellosis, brucellosis, <i>Campylobacter</i> infection, <i>Chlamydia pneumoniae</i> infection, chronic meningococcemia, ehrlichiosis, gonococcemia, legionellosis, leptospirosis, listeriosis, louse-borne relapsing fever (<i>Borrelia recurrentis</i>), Lyme disease, melioidosis (<i>Pseudomonas pseudomallei</i>), <i>Mycoplasma</i> infection, nocardiosis, psittacosis, Q fever (<i>Coxiella burnetii</i>), rickettsiosis, <i>Spirillum minor</i> infection, <i>Streptobacillus moniliformis</i> infection, syphilis, tick-borne relapsing fever (<i>Borrelia duttonii</i>), tuberculosis, tularemia, typhoid fever and other salmonellosis, Whipple's disease (<i>Tropheryma whipplei</i>), yersiniosis
Fungal	Aspergillosis, blastomycosis, candidiasis, coccidioidomycosis, cryptococcosis, histoplasmosis, <i>Malassezia furfur</i> infection, paracoccidioidomycosis, <i>Pneumocystis jirovecii</i> pneumonia, sporotrichosis, zygomycosis
Parasitic	Amebiasis, babesiosis, echinococcosis, fascioliasis, malaria, schistosomiasis, strongyloidiasis, toxocarosis, toxoplasmosis, trichinellosis, trypanosomiasis, visceral leishmaniasis
Viral	Colorado tick fever, coxsackievirus infection, cytomegalovirus infection, dengue, Epstein-Barr virus infection, hantavirus infection, hepatitis (A, B, C, D, E), herpes simplex, HIV infection, human herpesvirus 6 infection, parvovirus infection, West Nile virus infection
Noninfectious Inflammatory Diseases	
Systemic rheumatic and autoimmune diseases	Ankylosing spondylitis, antiphospholipid syndrome, autoimmune hemolytic anemia, autoimmune hepatitis, Behçet's disease, cryoglobulinemia, dermatomyositis, Felty syndrome, gout, mixed connective-tissue disease, polymyositis, pseudogout, reactive arthritis, relapsing polychondritis, rheumatic fever, rheumatoid arthritis, Sjögren's syndrome, systemic lupus erythematosus, Vogt-Koyanagi-Harada syndrome
Vasculitis	Allergic vasculitis, Churg-Strauss syndrome, giant cell vasculitis/polymyalgia rheumatica, granulomatosis with polyangiitis, hypersensitivity vasculitis, Kawasaki's disease, polyarteritis nodosa, Takayasu arteritis, urticarial vasculitis
Granulomatous diseases	Idiopathic granulomatous hepatitis, sarcoidosis
Autoinflammatory syndromes	Adult-onset Still's disease, Blau syndrome, CAPS ^b (cryopyrin-associated periodic syndromes), Crohn's disease, DIRA (deficiency of the interleukin 1 receptor antagonist), familial Mediterranean fever, hemophagocytic syndrome, hyper-IgD syndrome (HIDS, also known as mevalonate kinase deficiency), juvenile idiopathic arthritis, PAPA syndrome (pyogenic sterile arthritis, pyoderma gangrenosum, and acne), PFAPA syndrome (periodic fever, aphthous stomatitis, pharyngitis, adenitis), recurrent idiopathic pericarditis, SAPHO (synovitis, acne, pustulosis, hyperostosis, osteomyelitis), Schnitzler's syndrome, TRAPS (tumor necrosis factor receptor-associated periodic syndrome)
Neoplasms	
Hematologic malignancies	Amyloidosis, angioimmunoblastic lymphoma, Castleman's disease, Hodgkin's disease, hypereosinophilic syndrome, leukemia, lymphomatoid granulomatosis, malignant histiocytosis, multiple myeloma, myelodysplastic syndrome, myelofibrosis, non-Hodgkin's lymphoma, plasmacytoma, systemic mastocytosis, vaso-occlusive crisis in sickle cell disease
Solid tumors	Most solid tumors and metastases can cause fever. Those most commonly causing FUO are breast, colon, hepatocellular, lung, pancreatic, and renal cell carcinomas.
Benign tumors	Angiomyolipoma, cavernous hemangioma of the liver, craniopharyngioma, necrosis of dermoid tumor in Gardner's syndrome
Miscellaneous	
	ADEM (acute disseminated encephalomyelitis), adrenal insufficiency, aneurysms, anomalous thoracic duct, aortic dissection, aortic-enteral fistula, aseptic meningitis (Mollaret's syndrome), atrial myxoma, brewer's yeast ingestion, Caroli disease, cholesterol emboli, cirrhosis, complex partial status epilepticus, cyclic neutropenia, drug fever, Erdheim-Chester disease, extrinsic allergic alveolitis, Fabry's disease, factitious disease, fire-eater's lung, fraudulent fever, Gaucher's disease, Hamman-Rich syndrome (acute interstitial pneumonia), Hashimoto's encephalopathy, hematoma, hypersensitivity pneumonitis, hypertriglyceridemia, hypothalamic hypopituitarism, idiopathic normal-pressure hydrocephalus, inflammatory pseudotumor, Kikuchi's disease, linear IgA dermatosis, mesenteric fibromatosis, metal fume fever, milk protein allergy, myotonic dystrophy, nonbacterial osteitis, organic dust toxic syndrome, panniculitis, POEMS (polyneuropathy, organomegaly, endocrinopathy, monoclonal protein, skin changes), polymer fume fever, post-cardiac injury syndrome, primary biliary cirrhosis, primary hyperparathyroidism, pulmonary embolism, pyoderma gangrenosum, retroperitoneal fibrosis, Rosai-Dorfman disease, sclerosing mesenteritis, silicone embolization, subacute thyroiditis (de Quervain's), Sweet syndrome (acute febrile neutrophilic dermatosis), thrombosis, tubulointerstitial nephritis and uveitis syndrome (TINU), ulcerative colitis
Thermoregulatory Disorders	
Central	Brain tumor, cerebrovascular accident, encephalitis, hypothalamic dysfunction
Peripheral	Anhidrotic ectodermal dysplasia, exercise-induced hyperthermia, hyperthyroidism, pheochromocytoma

^aThis table includes all causes of FUO that have been described in the literature. ^bCAPS includes chronic infantile neurologic cutaneous and articular syndrome (CINCA, also known as neonatal-onset multisystem inflammatory disease, or NOMID), familial cold autoinflammatory syndrome (FCAS), and Muckle-Wells syndrome.

can be nonspecific. Serologic testing for Q fever, which results from exposure to animals or animal products, should be performed when the patient lives in a rural area or has a history of heart valve disease, an aortic aneurysm, or a vascular prosthesis. In patients with unexplained symptoms localized to the central nervous system (CNS), gastrointestinal tract, or joints, polymerase chain reaction (PCR) testing for *Tropheryma whipplei* should be performed. Travel to or (former) residence in tropical countries or the American Southwest should lead

to consideration of infectious diseases such as malaria, leishmaniasis, histoplasmosis, or coccidioidomycosis. Fever with signs of endocarditis and negative blood culture results poses a special problem. Culture-negative endocarditis may be due to difficult-to-culture bacteria such as nutritionally variant bacteria, HACEK organisms (*Haemophilus parainfluenzae*, *H. paraphrophilus*, *Aggregatibacter* species [*actinomycetemcomitans*, *aphrophilus*], *Cardiobacterium* species [*hominis*, *valvarum*], *Eikenella corrodens*, and *Kingella kingae*; discussed below),

Coxiella burnetii (as indicated above), *T. whipplei*, and *Bartonella* species. Marantic endocarditis is a sterile thrombotic disease that occurs as a paraneoplastic phenomenon, especially with adenocarcinomas. Sterile endocarditis is also seen in the context of systemic lupus erythematosus and antiphospholipid syndrome.

Of the NIIDs, large-vessel vasculitis, polymyalgia rheumatica, sarcoidosis, familial Mediterranean fever, and adult-onset Still's disease are rather common diagnoses in patients with FUO. The hereditary autoinflammatory syndromes are very rare and usually present in young patients. Schnitzler's syndrome, which can present at any age, is uncommon but can often be diagnosed easily in a patient with FUO who presents with urticaria, bone pain, and monoclonal gammopathy.

Although most tumors can present with fever, malignant lymphoma is by far the most common diagnosis of FUO among the neoplasms. Sometimes the fever even precedes lymphadenopathy detectable by physical examination.

Apart from drug-induced fever and exercise-induced hyperthermia, none of the miscellaneous causes of fever is found very frequently in patients with FUO. Virtually all drugs can cause fever, even that commencing after long-term use. *Drug-induced fever*, including DRESS (drug reaction with eosinophilia and systemic symptoms; Fig. 25e-49), is often accompanied by eosinophilia and also by lymphadenopathy, which can be extensive. More common causes of drug-induced fever are allopurinol, carbamazepine, lamotrigine, phenytoin, sulfasalazine, furosemide, antimicrobial drugs (especially sulfonamides, minocycline, vancomycin, β -lactam antibiotics, and isoniazid), some cardiovascular drugs (e.g., quinidine), and some antiretroviral drugs (e.g., nevirapine). *Exercise-induced hyperthermia* (Chap. 479e) is characterized by an elevated body temperature that is associated with moderate to strenuous exercise lasting from half an hour up to several hours without an increase in CRP level or ESR; typically these patients sweat during the temperature elevation. *Factitious fever* (fever artificially induced by the patient—for example, by IV injection of contaminated water) should be considered in all patients but is more common among young women in health care professions. In *fraudulent fever*, the patient is normothermic but manipulates the thermometer. Simultaneous measurements at different body sites (rectum, ear, mouth) should rapidly identify this diagnosis. Another clue to fraudulent fever is a dissociation between pulse rate and temperature.

Previous studies of FUO have shown that a diagnosis is more likely in elderly patients than in younger age groups. In many cases, FUO in the elderly results from an atypical manifestation of a common disease, among which giant cell arteritis and polymyalgia rheumatica are most frequently involved. Tuberculosis is the most common infectious disease associated with FUO in elderly patients, occurring much more often than in younger patients. As many of these diseases are treatable, it is well worth pursuing the cause of fever in elderly patients.

APPROACH TO THE PATIENT: Fever of Unknown Origin

FIRST-STAGE DIAGNOSTIC TESTS

Figure 26-1 shows a structured approach to patients presenting with FUO. The most important step in the diagnostic workup is the search for potentially diagnostic clues (PDCs) through complete and repeated history-taking and physical examination and the obligatory investigations listed above. PDCs are defined as all localizing signs, symptoms, and abnormalities potentially pointing toward a diagnosis. Although PDCs are often misleading, only with their help can a concise list of probable diagnoses be made. The history should include information about the fever pattern (continuous or recurrent) and duration, previous medical history, present and recent drug use, family history, sexual history, country of origin, recent and remote travel, unusual environmental exposures associated with travel or hobbies, and animal contacts. A complete physical examination should be performed, with special attention to the eyes, lymph nodes, temporal arteries, liver, spleen,

sites of previous surgery, entire skin surface, and mucous membranes. Before further diagnostic tests are initiated, antibiotic and glucocorticoid treatment, which can mask many diseases, should be stopped. For example, blood and other cultures are not reliable when samples are obtained during antibiotic treatment, and the size of enlarged lymph nodes usually decreases during glucocorticoid treatment, regardless of the cause of the lymphadenopathy. Despite the high number of false-positive ultrasounds and the relatively low sensitivity of chest x-rays, the performance of these simple, low-cost diagnostic tests remains obligatory in all patients with FUO in order to separate cases that are caused by easily diagnosed diseases from those that are not. Abdominal ultrasound is preferred to abdominal CT as an obligatory test because of relatively low cost, lack of radiation burden, and absence of side effects.

Only rarely do biochemical tests (beyond the obligatory tests needed to classify a patient's fever as FUO) lead directly to a definitive diagnosis in the absence of PDCs. The diagnostic yield of immunologic serology other than that included in the obligatory tests is relatively low. These tests more often yield false-positive rather than true-positive results and are of little use without PDCs pointing to specific immunologic disorders. Given the absence of specific symptoms in many patients and the relatively low cost of the test, investigation of cryoglobulins appears to be a valuable screening test in patients with FUO.

Multiple blood samples should be cultured in the laboratory long enough to ensure ample growth time for any fastidious organisms, such as HACEK organisms. It is critical to inform the laboratory of the intent to test for unusual organisms. Specialized media should be used when the history suggests uncommon microorganisms, such as *Histoplasma* or *Legionella*. Performing more than three blood cultures or more than one urine culture is useless in patients with FUO in the absence of PDCs (e.g., a high level of clinical suspicion of endocarditis). Repeating blood or urine cultures is useful only when previously cultured samples were collected during antibiotic treatment or within 1 week after its discontinuation. FUO with headache should prompt microbiologic examination of cerebrospinal fluid (CSF) for organisms including herpes simplex virus (HSV; especially HSV-2), *Cryptococcus neoformans*, and *Mycobacterium tuberculosis*. In CNS tuberculosis, the CSF typically has elevated protein and lowered glucose concentrations, with a mononuclear pleocytosis. CSF protein levels range from 100 to 500 mg/dL in most patients, the CSF glucose concentration is <45 mg/dL in 80% of cases, and the usual CSF cell count is between 100 and 500 cells/ μ L.

Microbiologic serology should not be included in the diagnostic workup in patients without PDCs for specific infections. A TST is included in the obligatory investigations, but it may yield false-negative results in patients with miliary tuberculosis, malnutrition, or immunosuppression. Although the interferon γ release assay is less influenced by prior vaccination with bacille Calmette-Guérin or by infection with nontuberculous mycobacteria, its sensitivity is similar to that of the TST; a negative TST or interferon γ release assay therefore does not exclude a diagnosis of tuberculosis. Miliary tuberculosis is especially difficult to diagnose. Granulomatous disease in liver or bone marrow biopsy samples, for example, should always lead to a (re)consideration of this diagnosis. If miliary tuberculosis is suspected, liver biopsy for acid-fast smear, culture, and PCR probably still has the highest diagnostic yield; however, biopsies of bone marrow, lymph nodes, or other involved organs also can be considered.

The diagnostic yield of echocardiography, sinus radiography, radiologic or endoscopic evaluation of the gastrointestinal tract, and bronchoscopy is very low in the absence of PDCs. Therefore, these tests should not be used as screening procedures.

After identification of all PDCs retrieved from the history, physical examination, and obligatory tests, a limited list of the most probable diagnoses should be made. Since most investigations are helpful only for patients who have PDCs for the diagnoses sought, further diagnostic procedures should be limited to specific investigations aimed at confirming or excluding diseases on this list. In FUO, the

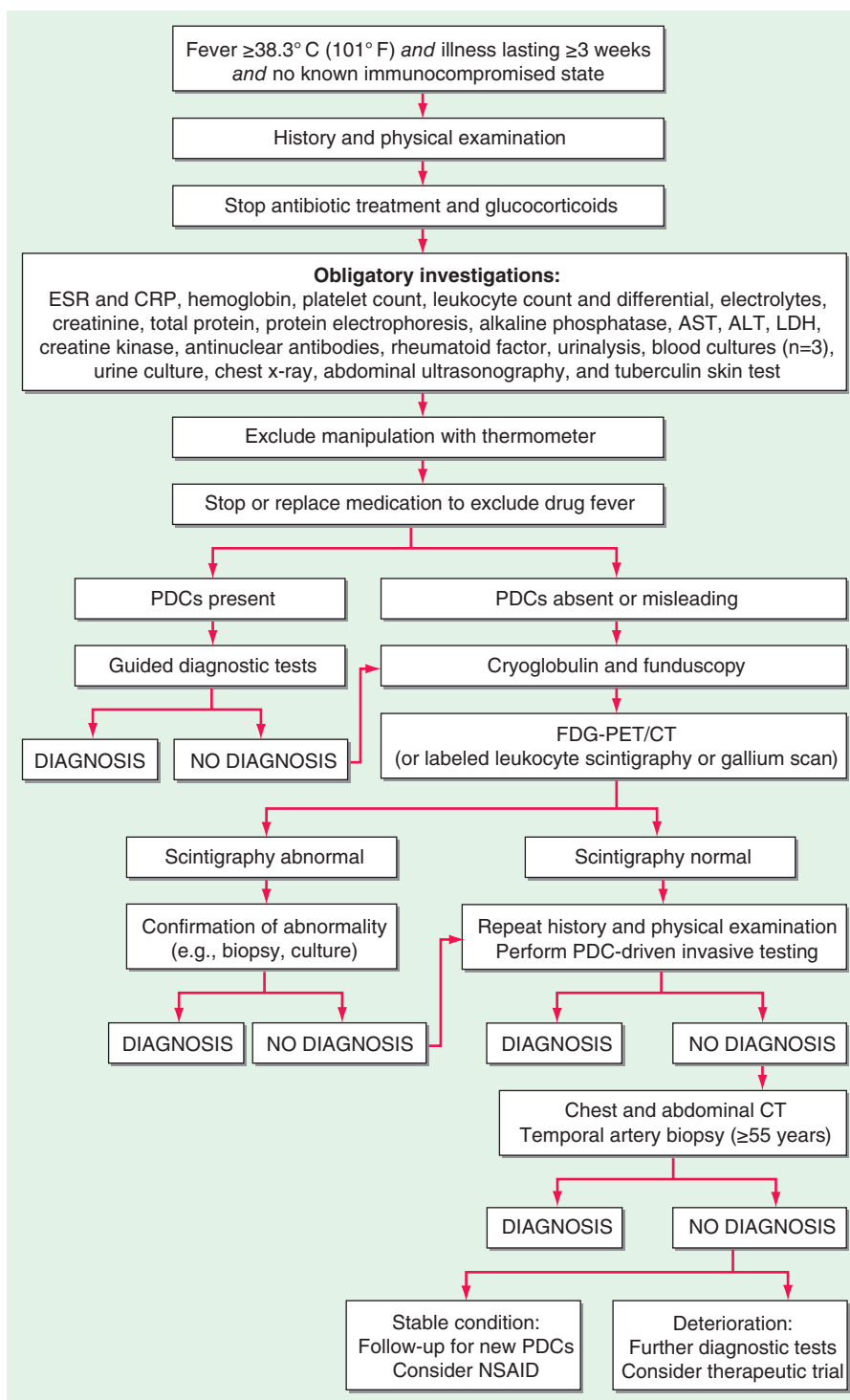


FIGURE 26-1 Structured approach to patients with FUO. ALT, alanine aminotransferase; AST, aspartate aminotransferase; CRP, C-reactive protein; ESR, erythrocyte sedimentation rate; FDG-PET/CT, ^{18}F -fluorodeoxyglucose positron emission tomography combined with low-dose computed tomography; LDH, lactate dehydrogenase; PDCs, potentially diagnostic clues (all localizing signs, symptoms, and abnormalities potentially pointing toward a diagnosis); NSAID, nonsteroidal anti-inflammatory drug.

diagnostic pointers are numerous and diverse but may be missed on initial examination, often being detected only by a very careful examination performed subsequently. In the absence of PDCs, the history and physical examination should therefore be repeated regularly. One of the first steps should be to rule out factitious or fraudulent fever, particularly in patients without signs of inflammation in laboratory tests. All medications, including nonprescription drugs and nutritional supplements, should be discontinued early in the evaluation to exclude drug fever. If fever persists beyond 72 h after discontinuation of the suspected drug, it is unlikely that this

drug is the cause. In patients without PDCs or with only misleading PDCs, funduscopy by an ophthalmologist may be useful in the early stage of the diagnostic workup. When the first-stage diagnostic tests do not lead to a diagnosis, scintigraphy should be performed, especially when the ESR or CRP level is elevated.

Recurrent Fever In patients with recurrent fever, the diagnostic workup should consist of thorough history-taking, physical examination, and obligatory tests. The search for PDCs should be directed to clues matching known recurrent syndromes (Table 26-3). Patients

TABLE 26-3 ALL REPORTED CAUSES OF RECURRENT FEVER^a

Infections	
Bacterial, nonspecific	Apical granuloma, diverticulitis, prostatitis, recurrent bacteremia caused by colonic neoplasia or persistent focal infection, recurrent cellulitis, recurrent cholangitis or cholecystitis, recurrent pneumonia, recurrent sinusitis, recurrent urinary tract infection
Bacterial, specific	Bartonellosis, brucellosis, chronic gonococcemia, chronic meningococcemia, louse-borne relapsing fever (<i>Borrelia recurrentis</i>), melioidosis (<i>Pseudomonas pseudomallei</i>), Q fever (<i>Coxiella burnetii</i>), salmonellosis, <i>Spirillum minor</i> infection, <i>Streptobacillus moniliformis</i> infection, syphilis, tick-borne relapsing fever (<i>Borrelia duttonii</i>), tularemia, Whipple's disease (<i>Tropheryma whipplei</i>), yersiniosis
Fungal	Coccidioidomycosis, paracoccidioidomycosis
Parasitic	Babesiosis, malaria, toxoplasmosis, trypanosomiasis, visceral leishmaniasis
Viral	Cytomegalovirus infection, Epstein-Barr virus infection, herpes simplex
Noninfectious Inflammatory Diseases	
Systemic rheumatic and autoimmune diseases	Ankylosing spondylitis, antiphospholipid syndrome, autoimmune hemolytic anemia, autoimmune hepatitis, Behçet's disease, cryoglobulinemia, gout, polymyositis, pseudogout, reactive arthritis, relapsing polychondritis, systemic lupus erythematosus
Vasculitis	Churg-Strauss syndrome, giant cell vasculitis/polymyalgia rheumatica, hypersensitivity vasculitis, polyarteritis nodosa, urticarial vasculitis
Granulomatous diseases	Idiopathic granulomatous hepatitis, sarcoidosis
Autoinflammatory syndromes	Adult-onset Still's disease, Blau syndrome, CAPS ^b (cryopyrin-associated periodic syndrome), Crohn's disease, DIRA (deficiency of the IL-1 receptor antagonist), familial Mediterranean fever, hemophagocytic syndrome, hyper-IgD syndrome (HIDS, also known as mevalonate kinase deficiency), juvenile idiopathic arthritis, PAPA syndrome (pyogenic sterile arthritis, pyoderma gangrenosum, and acne), PFAPA syndrome (periodic fever, aphthous stomatitis, pharyngitis, adenitis), recurrent idiopathic pericarditis, SAPHO (synovitis, acne, pustulosis, hyperostosis, osteomyelitis), Schnitzler's syndrome, TRAPS (tumor necrosis factor receptor-associated periodic syndrome)
Neoplasms	
	Angioimmunoblastic lymphoma, Castleman's disease, colon carcinoma, craniopharyngioma, Hodgkin's disease, non-Hodgkin lymphoma, malignant histiocytosis, mesothelioma
Miscellaneous	
	Adrenal insufficiency, aortic-enteral fistula, aseptic meningitis (Mollaret's syndrome), atrial myxoma, brewer's yeast ingestion, cholesterol emboli, cyclic neutropenia, drug fever, extrinsic allergic alveolitis, Fabry's disease, factitious disease, fraudulent fever, Gaucher's disease, hypersensitivity pneumonitis, hypertriglyceridemia, hypothalamic hypopituitarism, inflammatory pseudotumor, metal fume fever, milk protein allergy, polymer fume fever, pulmonary embolism, sclerosing mesenteritis
Thermoregulatory Disorders	
Central	Hypothalamic dysfunction
Peripheral	Anhidrotic ectodermal dysplasia, exercise-induced hyperthermia, pheochromocytoma

^aThis table includes all causes of recurrent fever that have been described in the literature. ^bCAPS includes chronic infantile neurologic cutaneous and articular syndrome (CINCA, also known as neonatal-onset multisystem inflammatory disease, or NOMID), familial cold autoinflammatory syndrome (FCAS), and Muckle-Wells syndrome.

should be asked to return during a febrile episode so that the history, physical examination, and laboratory tests can be repeated during a symptomatic phase. Further diagnostic tests, such as scintigraphic imaging (see below), should be performed only during a febrile episode because abnormalities may be absent between episodes. In patients with recurrent fever lasting >2 years, it is very unlikely that the fever is caused by infection or malignancy. Further diagnostic tests in that direction should be considered only when PDCs for infections, vasculitis syndromes, or malignancy are present or when the patient's clinical condition is deteriorating.

Scintigraphy Scintigraphic imaging is a noninvasive method allowing delineation of foci in all parts of the body on the basis of functional changes in tissues. This procedure plays an important role in the diagnosis of patients with FUO in clinical practice. Conventional scintigraphic methods used in clinical practice are ⁶⁷Ga-citrate scintigraphy and ¹¹¹In- or ^{99m}Tc-labeled leukocyte scintigraphy. Focal infectious and inflammatory processes can also be detected by several radiologic techniques, such as CT, MRI, and ultrasound. However, because of the lack of substantial pathologic changes in the early phase, infectious and inflammatory foci cannot be detected at this time. Furthermore, distinguishing active infectious or inflammatory lesions from residual changes due to cured processes or surgery remains critical. Finally, CT and MRI routinely provide information only on part of the body, while scintigraphy readily allows whole-body imaging.

Fluorodeoxyglucose Positron Emission Tomography ¹⁸F-fluorodeoxyglucose (FDG) positron emission tomography (PET) has become an

established imaging procedure in FUO. FDG accumulates in tissues with a high rate of glycolysis, which occurs not only in malignant cells but also in activated leukocytes, and thus permits the imaging of acute and chronic inflammatory processes. Normal uptake may obscure pathologic foci in the brain, heart, bowel, kidneys, and bladder. In patients with fever, bone marrow uptake is frequently increased in a nonspecific way due to cytokine activation, which upregulates glucose transporters in bone marrow cells. Compared with conventional scintigraphy, FDG-PET offers the advantages of higher resolution, greater sensitivity in chronic low-grade infections, and a high degree of accuracy in the central skeleton. Furthermore, vascular uptake of FDG is increased in patients with vasculitis. The mechanisms responsible for FDG uptake do not allow differentiation among infection, sterile inflammation, and malignancy. However, in patients with FUO, since all of these disorders are causes of FUO, FDG-PET can be used to guide additional diagnostic tests (e.g., targeted biopsies) that may yield the final diagnosis. Improved anatomic resolution by direct integration with CT (FDG-PET/CT) has further improved the accuracy of this modality.

Overall rates of helpfulness in final diagnosis of FUO are 40% for FDG-PET and 54% for FDG-PET/CT. In one study, FDG-PET was never helpful in diagnosing FUO in patients with a normal CRP level and a normal ESR. In two prospective studies in patients with FUO, FDG-PET was superior to ⁶⁷Ga-citrate scintigraphy, with a similar or better diagnostic yield and results that were available within hours instead of days. In one study, the sensitivity of FDG-PET was greater than that of ¹¹¹In-granulocyte scintigraphy (86% vs 20%) in patients with FUO.

Although scintigraphic techniques do not directly provide a definitive diagnosis, they often identify the anatomic location of a particular ongoing metabolic process and, with the help of other techniques such as biopsy and culture, facilitate timely diagnosis and treatment. Pathologic FDG uptake is quickly eradicated by treatment with glucocorticoids in many diseases, including vasculitis and lymphoma; therefore, glucocorticoid use should be stopped or postponed until after FDG-PET is performed. Results reported in the literature and the advantages offered by FDG-PET indicate that conventional scintigraphic techniques should be replaced by FDG-PET/CT in the investigation of patients with FUO at institutions where this technique is available. FDG-PET/CT is a relatively expensive procedure whose availability is still limited compared with that of CT and conventional scintigraphy. Nevertheless, FDG-PET/CT can be cost-effective in the FUO diagnostic workup if used at an early stage, helping to establish an early diagnosis, reducing days of hospitalization for diagnostic purposes, and obviating unnecessary and unhelpful tests.

LATER-STAGE DIAGNOSTIC TESTS

In some cases, more invasive tests are appropriate. Abnormalities found with scintigraphic techniques often need to be confirmed by pathology and/or culture of biopsy specimens. If lymphadenopathy is found, lymph node biopsy is necessary, even when the affected lymph nodes are hard to reach. In the case of skin lesions, skin biopsy should be undertaken. In one study, pulmonary wedge excision, histologic examination of an excised tonsil, and biopsy of the peritoneum were performed in light of PDCs or abnormal FDG-PET results and yielded a diagnosis.

If no diagnosis is reached despite scintigraphic and PDC-driven histologic investigations or culture, second-stage screening diagnostic tests should be considered (Fig. 26-1). In three studies, the diagnostic yield of screening chest and abdominal CT in patients with FUO was ~20%. The specificity of chest CT was ~80%, but that of abdominal CT varied between 63% and 80%. Despite the relatively limited specificity of abdominal CT and the probably limited additional value of chest CT after normal FDG-PET, chest and abdominal CT may be used as screening procedures at a later stage of the diagnostic protocol because of their noninvasive nature and high sensitivity. Bone marrow aspiration is seldom useful in the absence of PDCs for bone marrow disorders. With addition of FDG-PET, which is very sensitive in detecting lymphoma, carcinoma, and osteomyelitis, the value of bone marrow biopsy as a screening procedure is probably further reduced. Several studies have shown a high prevalence of giant cell arteritis among patients with FUO, with rates up to 17% among elderly patients. Giant cell arteritis often involves large arteries and in most cases can be diagnosed by FDG-PET. However, temporal artery biopsy is still recommended for patients ≥ 55 years of age in a later stage of the diagnostic protocol: FDG-PET will not be useful in vasculitis limited to the temporal arteries because of the small diameter of these vessels and the high levels of FDG uptake in the brain that overlies them. In the past, liver biopsies have often been performed as a screening procedure in patients with FUO. In each of two recent studies, liver biopsy as part of the later stage of a screening diagnostic protocol was helpful in only one patient. Moreover, abnormal liver tests are not predictive of a diagnostic liver biopsy in FUO. Liver biopsy is an invasive procedure that carries the possibility of complications and even death. Therefore, it should not be used for screening purposes in patients with FUO except in those with PDCs for liver disease.

In patients with unexplained fever after all of the above procedures, the last step in the diagnostic workup—with only a marginal diagnostic yield—comes at an extraordinarily high cost in terms of both expense and discomfort for the patient. Repetition of a thorough history-taking and physical examination and review of laboratory results and imaging studies (including those from other hospitals) are recommended. Diagnostic delay often results from a failure to recognize PDCs in the available information. In these

patients with persisting FUO, waiting for new PDCs to appear probably is better than ordering more screening investigations. Only when a patient's condition deteriorates without providing new PDCs should a further diagnostic workup be performed.

TREATMENT FEVER OF UNKNOWN ORIGIN

Empirical therapeutic trials with antibiotics, glucocorticoids, or antituberculous agents should be avoided in FUO except when a patient's condition is rapidly deteriorating after the aforementioned diagnostic tests have failed to provide a definite diagnosis.

ANTIBIOTICS AND ANTITUBERCULOUS THERAPY

Antibiotic or antituberculous therapy may irrevocably diminish the ability to culture fastidious bacteria or mycobacteria. However, hemodynamic instability or neutropenia is a good indication for empirical antibiotic therapy. If the TST is positive or if granulomatous disease is present with anergy and sarcoidosis seems unlikely, a therapeutic trial for tuberculosis should be started. Especially in military tuberculosis, it may be very difficult to obtain a rapid diagnosis. If the fever does not respond after 6 weeks of empirical antituberculous treatment, another diagnosis should be considered.

COLCHICINE, NONSTEROIDAL ANTI-INFLAMMATORY DRUGS, AND GLUCOCORTICOIDS

Colchicine is highly effective in preventing attacks of familial Mediterranean fever but is not always effective once an attack is well under way. When familial Mediterranean fever is suspected, the response to colchicine is not a completely reliable diagnostic tool in the acute phase, but with colchicine treatment most patients show remarkable improvements in the frequency and severity of subsequent febrile episodes within weeks to months. If the fever persists and the source remains elusive after completion of the later-stage investigations, supportive treatment with nonsteroidal anti-inflammatory drugs (NSAIDs) can be helpful. The response of adult-onset Still's disease to NSAIDs is dramatic in some cases. The effects of glucocorticoids on giant cell arteritis and polymyalgia rheumatica are equally impressive. Early empirical trials with glucocorticoids, however, decrease the chances of reaching a diagnosis for which more specific and sometimes life-saving treatment might be more appropriate, such as malignant lymphoma. The ability of NSAIDs and glucocorticoids to mask fever while permitting the spread of infection or lymphoma dictates that their use should be avoided unless infectious diseases and malignant lymphoma have been largely ruled out and inflammatory disease is probable and is likely to be debilitating or threatening.

ANAKINRA

Interleukin (IL) 1 is a key cytokine in local and systemic inflammation and the febrile response. The availability of specific IL-1-targeting agents has revealed a pathologic role of IL-1-mediated inflammation in a growing list of diseases. Anakinra, a recombinant form of the naturally occurring IL-1 receptor antagonist (IL-1Ra), blocks the activity of both IL-1 α and IL-1 β . Anakinra is extremely effective in the treatment of many autoinflammatory syndromes, such as familial Mediterranean fever, cryopyrin-associated periodic syndrome, tumor necrosis factor receptor-associated periodic syndrome, hyper-IgD syndrome, and Schnitzler's syndrome. There is a growing list of other chronic inflammatory disorders in which the reduction of IL-1 activity can be highly effective. A therapeutic trial with anakinra can be considered in patients whose FUO has not been diagnosed after later-stage diagnostic tests. Although most chronic inflammatory conditions without a known basis can be controlled with glucocorticoids, monotherapy with IL-1 blockade can provide improved control without the metabolic, immunologic, and gastrointestinal side effects of glucocorticoid administration.

FUO-related mortality rates have continuously declined over recent decades. The majority of fevers are caused by treatable diseases, and the risk of death related to FUO is, of course, dependent on the underlying disease. In a study by our group (Table 26-1), none of 37 FUO patients without a diagnosis died during a follow-up period of at least 6 months; 4 of 36 patients with a diagnosis died during follow-up due to infection ($n = 1$) or malignancy ($n = 3$). Other studies have also

shown that malignancy accounts for most FUO-related deaths. Non-Hodgkin's lymphoma carries a disproportionately high death toll. In nonmalignant FUO, fatality rates are very low. The good outcome in patients without a diagnosis confirms that potentially lethal occult diseases are very unusual and that empirical therapy with antibiotics, antituberculous agents, or glucocorticoids is rarely required in stable patients. In less affluent regions, infectious diseases are still a major cause of FUO, and outcomes may be different.

SECTION 3 NERVOUS SYSTEM DYSFUNCTION

27 Syncope

Roy Freeman

Syncope is a transient, self-limited loss of consciousness due to acute global impairment of cerebral blood flow. The onset is rapid, duration brief, and recovery spontaneous and complete. Other causes of transient loss of consciousness need to be distinguished from syncope; these include seizures, vertebrobasilar ischemia, hypoxemia, and hypoglycemia. A syncopal prodrome (*presyncope*) is common, although loss of consciousness may occur without any warning symptoms. Typical presyncopal symptoms include dizziness, lightheadedness or faintness, weakness, fatigue, and visual and auditory disturbances. The causes of syncope can be divided into three general categories: (1) neurally mediated syncope (also called *reflex or vasovagal syncope*), (2) orthostatic hypotension, and (3) cardiac syncope.

Neurally mediated syncope comprises a heterogeneous group of functional disorders that are characterized by a transient change in the reflexes responsible for maintaining cardiovascular homeostasis. Episodic vasodilation (or loss of vasoconstrictor tone) and bradycardia occur in varying combinations, resulting in temporary failure of blood pressure control. In contrast, in patients with orthostatic hypotension due to autonomic failure, these cardiovascular homeostatic reflexes are chronically impaired. Cardiac syncope may be due to arrhythmias or structural cardiac diseases that cause a decrease in cardiac output. The clinical features, underlying pathophysiologic mechanisms, therapeutic interventions, and prognoses differ markedly among these three causes.

EPIDEMIOLOGY AND NATURAL HISTORY

Syncope is a common presenting problem, accounting for approximately 3% of all emergency room visits and 1% of all hospital admissions. The annual cost for syncope-related hospitalization in the United States is ~\$2.4 billion. Syncope has a lifetime cumulative incidence of up to 35% in the general population. The peak incidence in the young occurs between ages 10 and 30 years, with a median peak around 15 years. Neurally mediated syncope is the etiology in the vast majority of these cases. In elderly adults, there is a sharp rise in the incidence of syncope after 70 years.

In population-based studies, neurally mediated syncope is the most common cause of syncope. The incidence is slightly higher in females than males. In young subjects, there is often a family history in first-degree relatives. Cardiovascular disease due to structural disease or arrhythmias is the next most common cause in most series, particularly in emergency room settings and in older patients. Orthostatic hypotension also increases in prevalence with age because of the reduced baroreflex responsiveness, decreased cardiac compliance, and attenuation of the vestibulosympathetic reflex associated with aging. In the elderly, orthostatic hypotension is substantially more common in institutionalized (54–68%) than community-dwelling (6%) individuals, an observation most likely explained by the greater prevalence of

TABLE 27-1 HIGH-RISK FEATURES INDICATING HOSPITALIZATION OR INTENSIVE EVALUATION OF SYNCOPES

Chest pain suggesting coronary ischemia
Features of congestive heart failure
Moderate or severe valvular disease
Moderate or severe structural cardiac disease
Electrocardiographic features of ischemia
History of ventricular arrhythmias
Prolonged QT interval (>500 ms)
Repetitive sinoatrial block or sinus pauses
Persistent sinus bradycardia
Bi- or trifascicular block or intraventricular conduction delay with QRS duration ≥ 120 ms
Atrial fibrillation
Nonsustained ventricular tachycardia
Family history of sudden death
Preexcitation syndromes
Brugada pattern on ECG
Palpitations at time of syncope
Syncope at rest or during exercise

predisposing neurologic disorders, physiologic impairment, and vasoactive medication use among institutionalized patients.

The prognosis after a single syncopal event for all age groups is generally benign. In particular, syncope of noncardiac and unexplained origin in younger individuals has an excellent prognosis; life expectancy is unaffected. By contrast, syncope due to a cardiac cause, either structural heart disease or primary arrhythmic disease, is associated with an increased risk of sudden cardiac death and mortality from other causes. Similarly, mortality rate is increased in individuals with syncope due to orthostatic hypotension related to age and the associated comorbid conditions (Table 27-1).

PATHOPHYSIOLOGY

The upright posture imposes a unique physiologic stress upon humans; most, although not all, syncopal episodes occur from a standing position. Standing results in pooling of 500–1000 mL of blood in the lower extremities and splanchnic circulation. There is a decrease in venous return to the heart and reduced ventricular filling that result in diminished cardiac output and blood pressure. These hemodynamic changes provoke a compensatory reflex response, initiated by the baroreceptors in the carotid sinus and aortic arch, resulting in increased sympathetic outflow and decreased vagal nerve activity (Fig. 27-1). The reflex increases peripheral resistance, venous return to the heart, and cardiac output and thus limits the fall in blood pressure. If this response fails, as is the case chronically in orthostatic hypotension and transiently in neurally mediated syncope, cerebral hypoperfusion occurs.

Syncope is a consequence of global cerebral hypoperfusion and thus represents a failure of cerebral blood flow autoregulatory mechanisms.

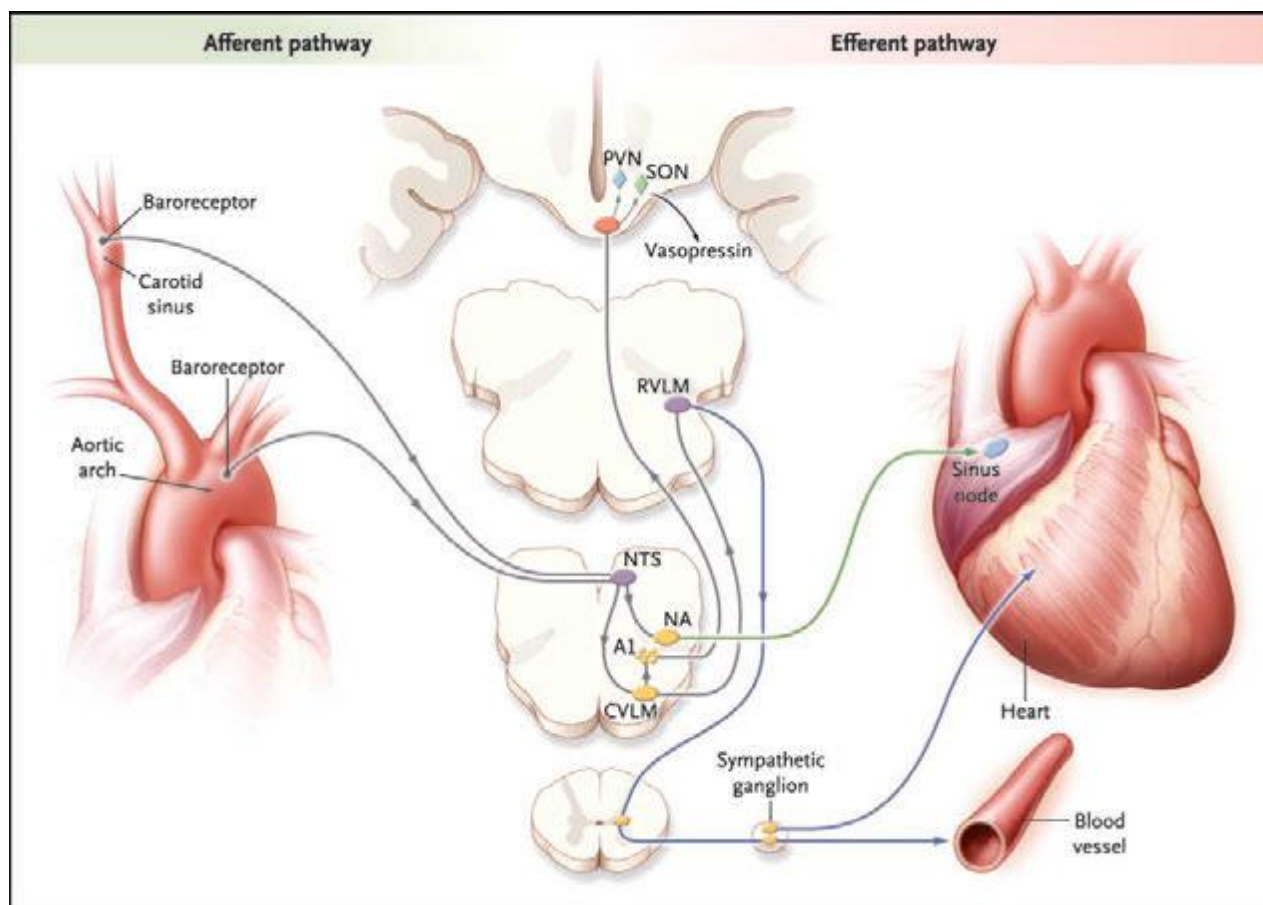


FIGURE 27-1 The baroreflex. A decrease in arterial pressure unloads the baroreceptors—the terminals of afferent fibers of the glossopharyngeal and vagus nerves—that are situated in the carotid sinus and aortic arch. This leads to a reduction in the afferent impulses that are relayed from these mechanoreceptors through the glossopharyngeal and vagus nerves to the nucleus of the tractus solitarius (NTS) in the dorsomedial medulla. The reduced baroreceptor afferent activity produces a decrease in vagal nerve input to the sinus node that is mediated via connections of the NTS to the nucleus ambiguus (NA). There is an increase in sympathetic efferent activity that is mediated by the NTS projections to the caudal ventrolateral medulla (CVLM) (an excitatory pathway) and from there to the rostral ventrolateral medulla (RVLN) (an inhibitory pathway). The activation of RVLN presympathetic neurons in response to hypotension is thus predominantly due to disinhibition. In response to a sustained fall in blood pressure, vasopressin release is mediated by projections from the A1 noradrenergic cell group in the ventrolateral medulla. This projection activates vasopressin-synthesizing neurons in the magnocellular portion of the paraventricular nucleus (PVN) and the supraoptic nucleus (SON) of the hypothalamus. *Blue* denotes sympathetic neurons, and *green* denotes parasympathetic neurons. (From R Freeman: *N Engl J Med* 358:615, 2008.)

Myogenic factors, local metabolites, and to a lesser extent autonomic neurovascular control are responsible for the autoregulation of cerebral blood flow (Chap. 330). The latency of the autoregulatory response is 5–10 s. Typically cerebral blood flow ranges from 50 to 60 mL/min per 100 g brain tissue and remains relatively constant over perfusion pressures ranging from 50 to 150 mmHg. Cessation of blood flow for 6–8 s will result in loss of consciousness, while impairment of consciousness ensues when blood flow decreases to 25 mL/min per 100 g brain tissue.

From the clinical standpoint, a fall in systemic systolic blood pressure to ~50 mmHg or lower will result in syncope. A decrease in cardiac output and/or systemic vascular resistance—the determinants of blood pressure—thus underlies the pathophysiology of syncope. Common causes of impaired cardiac output include decreased effective circulating blood volume; increased thoracic pressure; massive pulmonary embolus; cardiac brady- and tachyarrhythmias; valvular heart disease; and myocardial dysfunction. Systemic vascular resistance may be decreased by central and peripheral autonomic nervous system diseases, sympatholytic medications, and transiently during neurally mediated syncope. Increased cerebral vascular resistance, most frequently due to hypocarbia induced by hyperventilation, may also contribute to the pathophysiology of syncope.

Two patterns of electroencephalographic (EEG) changes occur in syncopal subjects. The first is a “slow-flat-slow” pattern (Fig. 27-2)

in which normal background activity is replaced with high-amplitude slow delta waves. This is followed by sudden flattening of the EEG—a cessation or attenuation of cortical activity—followed by the return of slow waves, and then normal activity. A second pattern, the “slow pattern,” is characterized by increasing and decreasing slow wave activity only. The EEG flattening that occurs in the slow-flat-slow pattern is a marker of more severe cerebral hypoperfusion. Despite the presence of myoclonic movements and other motor activity during some syncopal events, EEG seizure discharges are not detected.

CLASSIFICATION

NEURALLY MEDIATED SYNCOPÉ

Neurally mediated (reflex; vasovagal) syncope is the final pathway of a complex central and peripheral nervous system reflex arc. There is a sudden, transient change in autonomic efferent activity with increased parasympathetic outflow, plus sympathoinhibition (the vasodepressor response), resulting in bradycardia, vasodilation, and/or reduced vasoconstrictor tone. The resulting fall in systemic blood pressure can then reduce cerebral blood flow to below the compensatory limits of autoregulation (Fig. 27-3). In order to elicit neurally mediated syncope, a functioning autonomic nervous system is necessary, in contrast to syncope resulting from autonomic failure (discussed below).

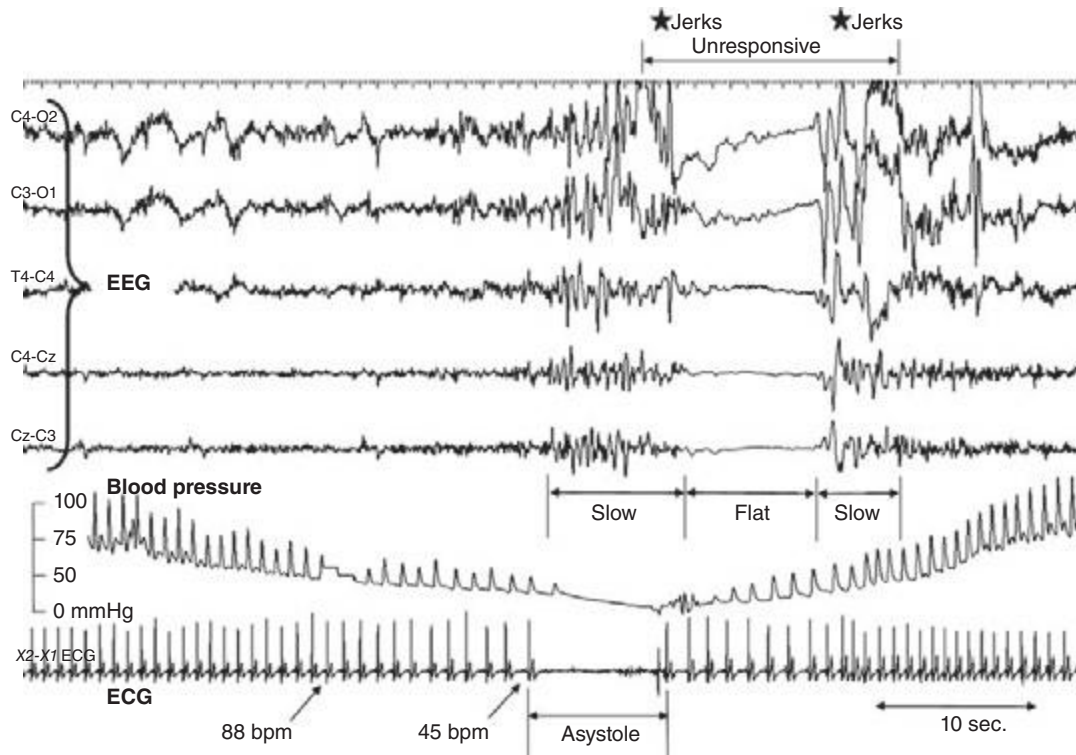


FIGURE 27-2 The electroencephalogram (EEG) in vasovagal syncope. A 1-min segment of a tilt-table test with typical vasovagal syncope demonstrating the “slow-flat-slow” EEG pattern. Finger beat-to-beat blood pressure, electrocardiogram (ECG), and selected EEG channels are shown. EEG slowing starts when systolic blood pressure drops to ~50 mmHg; heart rate is then approximately 45 beats/min (bpm). Asystole occurred, lasting about 8 s. The EEG flattens for a similar period, but with a delay. A transient loss of consciousness, lasting 14 s, was observed. There were muscle jerks just before and just after the flat period of the EEG. (Figure reproduced with permission from W Wieling et al: *Brain* 132:2630, 2009.)

Multiple triggers of the afferent limb of the reflex arc can result in neurally mediated syncope. In some situations, these can be clearly defined, e.g., the carotid sinus, the gastrointestinal tract, or the bladder. Often, however, the trigger is less easily recognized and the cause is multifactorial. Under these circumstances, it is likely that different afferent pathways converge on the central autonomic

network within the medulla that integrates the neural impulses and mediates the vasodepressor-bradycardic response.

Classification of Neurally Mediated Syncope Neurally mediated syncope may be subdivided based on the afferent pathway and provocative trigger. Vasovagal syncope (the common faint) is provoked by intense emotion, pain, and/or orthostatic stress, whereas the situational reflex syncopes have specific localized stimuli that provoke the reflex vasodilation and bradycardia that leads to syncope. The underlying mechanisms have been identified and pathophysiology delineated for most of these situational reflex syncopes. The afferent trigger may originate in the pulmonary system, gastrointestinal system, urogenital system, heart, and carotid artery (Table 27-2). Hyperventilation leading to hypocarbia and cerebral vasoconstriction, and raised intrathoracic pressure that impairs venous return to the heart, play a central role in many of the situational reflex syncopes. The afferent pathway of the reflex arc differs among these disorders, but the efferent response via the vagus and sympathetic pathways is similar.

Alternately, neurally mediated syncope may be subdivided based on the predominant efferent pathway. Vasodepressor syncope describes syncope predominantly due to efferent, sympathetic, vasoconstrictor failure; cardioinhibitory syncope describes syncope predominantly associated with bradycardia or asystole due

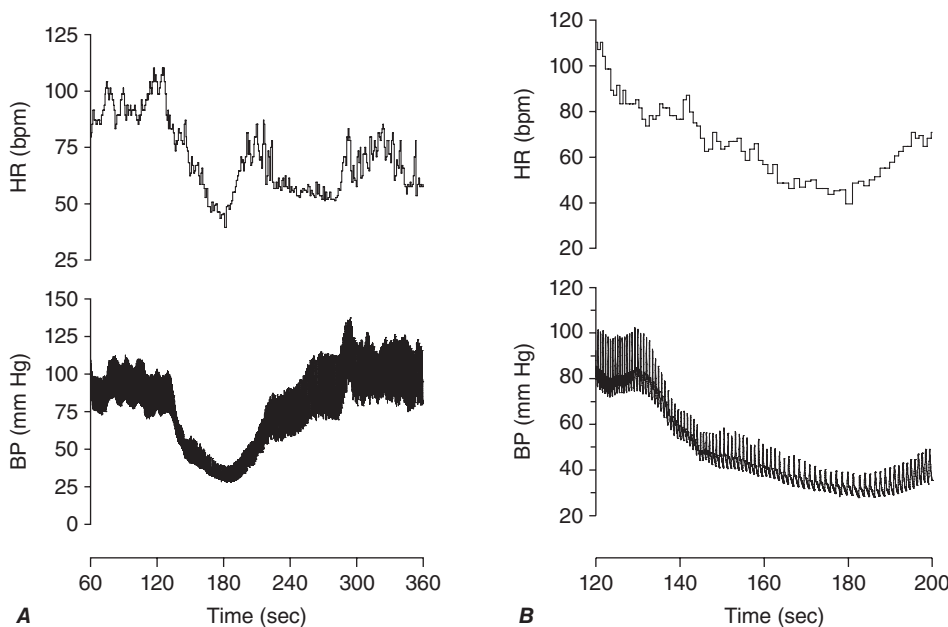


FIGURE 27-3 **A.** The paroxysmal hypotensive-bradycardic response that is characteristic of neurally mediated syncope. Noninvasive beat-to-beat blood pressure and heart rate are shown over 5 min (from 60 to 360 s) of an upright tilt on a tilt table. **B.** The same tracing expanded to show 80 s of the episode (from 80 to 200 s). BP, blood pressure; bpm, beats per minute; HR, heart rate.

TABLE 27-2 CAUSES OF SYNCOPE**A. Neurally Mediated Syncope**

- Vasovagal syncope
 - Provoked fear, pain, anxiety, intense emotion, sight of blood, unpleasant sights and odors, orthostatic stress
- Situational reflex syncope
 - Pulmonary
 - Cough syncope, wind instrument player's syncope, weightlifter's syncope, "mess trick"^a and "fainting lark,"^b sneeze syncope, airway instrumentation
 - Urogenital
 - Postmicturition syncope, urogenital tract instrumentation, prostatic massage
 - Gastrointestinal
 - Swallow syncope, glossopharyngeal neuralgia, esophageal stimulation, gastrointestinal tract instrumentation, rectal examination, defecation syncope
 - Cardiac
 - Bezold-Jarisch reflex, cardiac outflow obstruction
 - Carotid sinus
 - Carotid sinus sensitivity, carotid sinus massage
 - Ocular
 - Ocular pressure, ocular examination, ocular surgery

B. Orthostatic Hypotension

- Primary autonomic failure due to idiopathic central and peripheral neurodegenerative diseases—the "synucleinopathies"
 - Lewy body diseases
 - Parkinson's disease
 - Lewy body dementia
 - Pure autonomic failure
 - Multiple system atrophy (the Shy-Drager syndrome)
- Secondary autonomic failure due to autonomic peripheral neuropathies
 - Diabetes
 - Hereditary amyloidosis (familial amyloid polyneuropathy)
 - Primary amyloidosis (AL amyloidosis; immunoglobulin light chain associated)
 - Hereditary sensory and autonomic neuropathies (HSAN) (especially type III—familial dysautonomia)
 - Idiopathic immune-mediated autonomic neuropathy
 - Autoimmune autonomic ganglionopathy
 - Sjögren's syndrome
 - Paraneoplastic autonomic neuropathy
 - HIV neuropathy
- Postprandial hypotension
- Iatrogenic (drug-induced)
- Volume depletion

C. Cardiac Syncope

- Arrhythmias
 - Sinus node dysfunction
 - Atrioventricular dysfunction
 - Supraventricular tachycardias
 - Ventricular tachycardias
 - Inherited channelopathies
- Cardiac structural disease
 - Valvular disease
 - Myocardial ischemia
 - Obstructive and other cardiomyopathies
 - Atrial myxoma
 - Pericardial effusions and tamponade

^aHyperventilation for ~1 minute, followed by sudden chest compression. ^bHyperventilation (~20 breaths) in a squatting position, rapid rise to standing, then Valsalva.

to increased vagal outflow; and mixed syncope describes syncope in which there are both vagal and sympathetic reflex changes.

Features of Neurally Mediated Syncope In addition to symptoms of orthostatic intolerance such as dizziness, lightheadedness, and fatigue, premonitory features of autonomic activation may be present in patients with neurally mediated syncope. These include diaphoresis, pallor, palpitations, nausea, hyperventilation, and yawning. During the syncopal event, proximal and distal myoclonus (typically arrhythmic and multifocal) may occur, raising the possibility of epilepsy. The eyes typically remain open and usually deviate upward. Pupils are usually dilated. Roving eye movements may occur. Grunting, moaning, snorting, and stertorous breathing may be present. Urinary incontinence may occur. Fecal incontinence is very rare. Postictal confusion is also rare, although visual and auditory hallucinations and near death and out-of-body experiences are sometimes reported.

Although some predisposing factors and provocative stimuli are well established (for example, motionless upright posture, warm ambient temperature, intravascular volume depletion, alcohol ingestion, hypoxemia, anemia, pain, the sight of blood, venipuncture, and intense emotion), the underlying basis for the widely different thresholds for syncope among individuals exposed to the same provocative stimulus is not known. A genetic basis for neurally mediated syncope may exist; several studies have reported an increased incidence of syncope in first-degree relatives of fainters, but no gene or genetic marker has been identified, and environmental, social, and cultural factors have not been excluded by these studies.

TREATMENT NEURALLY MEDIATED SYNCOPE

Reassurance, avoidance of provocative stimuli, and plasma volume expansion with fluid and salt are the cornerstones of the management of neurally mediated syncope. Isometric counterpressure maneuvers of the limbs (leg crossing or handgrip and arm tensing) may raise blood pressure by increasing central blood volume and cardiac output. By maintaining pressure in the autoregulatory zone, these maneuvers avoid or delay the onset of syncope. Randomized controlled trials support this intervention.

Fludrocortisone, vasoconstricting agents, and beta-adrenoreceptor antagonists are widely used by experts to treat refractory patients, although there is no consistent evidence from randomized controlled trials for any pharmacotherapy to treat neurally mediated syncope. Because vasodilation is the dominant pathophysiologic syncopal mechanism in most patients, use of a cardiac pacemaker is rarely beneficial. Possible exceptions are older patients (>40 years) in whom syncope is associated with asystole or severe bradycardia and patients with prominent cardioinhibition due to carotid sinus syndrome. In these patients, dual-chamber pacing may be helpful.

ORTHOSTATIC HYPOTENSION

Orthostatic hypotension, defined as a reduction in systolic blood pressure of at least 20 mmHg or diastolic blood pressure of at least 10 mmHg within 3 min of standing or head-up tilt on a tilt table, is a manifestation of sympathetic vasoconstrictor (autonomic) failure (Fig. 27-4). In many (but not all) cases, there is no compensatory increase in heart rate despite hypotension; with partial autonomic failure, heart rate may increase to some degree but is insufficient to maintain cardiac output. A variant of orthostatic hypotension is "delayed" orthostatic hypotension, which occurs beyond 3 min of standing; this may reflect a mild or early form of sympathetic adrenergic dysfunction. In some cases, orthostatic hypotension occurs within 15 s of standing (so-called "initial" orthostatic hypotension), a finding that may reflect a transient mismatch between cardiac output and peripheral vascular resistance and does not represent autonomic failure.

Characteristic symptoms of orthostatic hypotension include lightheadedness, dizziness, and presyncope (near-faintness) occurring in response to sudden postural change. However, symptoms may be

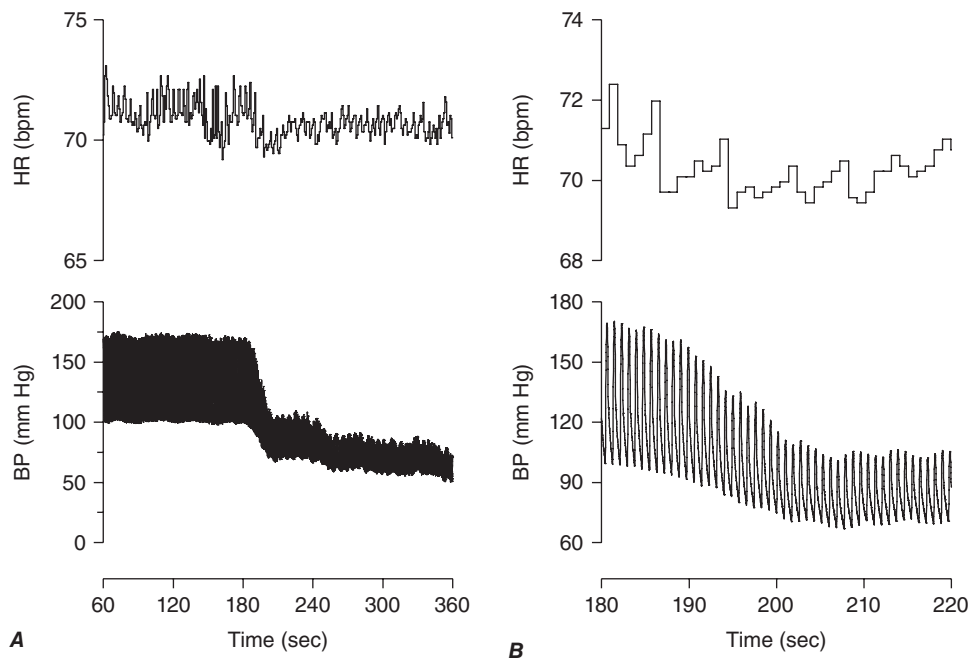


FIGURE 27-4 **A.** The gradual fall in blood pressure without a compensatory heart rate increase that is characteristic of orthostatic hypotension due to autonomic failure. Blood pressure and heart rate are shown over 5 min (from 60 to 360 s) of an upright tilt on a tilt table. **B.** The same tracing expanded to show 40 s of the episode (from 180 to 220 s). BP, blood pressure; bpm, beats per minute; HR, heart rate.

absent or nonspecific, such as generalized weakness, fatigue, cognitive slowing, leg buckling, or headache. Visual blurring may occur, likely due to retinal or occipital lobe ischemia. Neck pain, typically in the suboccipital, posterior cervical, and shoulder region (the “coat-hanger headache”), most likely due to neck muscle ischemia, may be the only symptom. Patients may report orthostatic dyspnea (thought to reflect ventilation-perfusion mismatch due to inadequate perfusion of ventilated lung apices) or angina (attributed to impaired myocardial perfusion even with normal coronary arteries). Symptoms may be exacerbated by exertion, prolonged standing, increased ambient temperature, or meals. Syncope is usually preceded by warning symptoms, but may occur suddenly, suggesting the possibility of a seizure or cardiac cause.

Supine hypertension is common in patients with orthostatic hypotension due to autonomic failure, affecting over 50% of patients in some series. Orthostatic hypotension may present after initiation of therapy for hypertension, and supine hypertension may follow treatment of orthostatic hypotension. However, in other cases, the association of the two conditions is unrelated to therapy; it may in part be explained by baroreflex dysfunction in the presence of residual sympathetic outflow, particularly in patients with central autonomic degeneration.

Causes of Neurogenic Orthostatic Hypotension Causes of neurogenic orthostatic hypotension include central and peripheral autonomic nervous system dysfunction (**Chap. 454**). Autonomic dysfunction of other organ systems (including the bladder, bowels, sexual organs, and sudomotor system) of varying severity frequently accompanies orthostatic hypotension in these disorders (**Table 27-2**).

The primary autonomic degenerative disorders are multiple system atrophy (the Shy-Drager syndrome; **Chap. 454**), Parkinson’s disease (**Chap. 449**), dementia with Lewy bodies (**Chap. 448**), and pure autonomic failure (**Chap. 454**). These are often grouped together as “synucleinopathies” due to the presence of alpha-synuclein, a small protein that precipitates predominantly in the cytoplasm of neurons in the Lewy body disorders (Parkinson’s disease, dementia with Lewy bodies, and pure autonomic failure) and in the glia in multiple system atrophy.

Peripheral autonomic dysfunction may also accompany small-fiber peripheral neuropathies such as those seen in diabetes, amyloid, immune-mediated neuropathies, hereditary sensory and autonomic neuropathies (HSAN; particularly HSAN type III, familial dysautonomia),

and inflammatory neuropathies (**Chaps. 459 and 460**). Less frequently, orthostatic hypotension is associated with the peripheral neuropathies that accompany vitamin B₁₂ deficiency, neurotoxic exposure, HIV and other infections, and porphyria.

Patients with autonomic failure and the elderly are susceptible to falls in blood pressure associated with meals. The magnitude of the blood pressure fall is exacerbated by large meals, meals high in carbohydrate, and alcohol intake. The mechanism of postprandial syncope is not fully elucidated.

Orthostatic hypotension is often iatrogenic. Drugs from several classes may lower peripheral resistance (e.g., alpha-adrenoreceptor antagonists used to treat hypertension and prostatic hypertrophy; antihypertensive agents of several classes; nitrates and other vasodilators; tricyclic agents and phenothiazines). Iatrogenic volume depletion due to diuresis and volume depletion due to medical causes (hemorrhage, vomiting, diarrhea, or decreased fluid intake) may also result in decreased effective circulatory volume, orthostatic hypotension, and syncope.

TREATMENT ORTHOSTATIC HYPOTENSION

The first step is to remove reversible causes—usually vasoactive medications (**Table 454-6**). Next, nonpharmacologic interventions should be introduced. These interventions include patient education regarding staged moves from supine to upright; warnings about the hypotensive effects of large meals; instructions about the isometric counterpressure maneuvers that increase intravascular pressure (see above); and raising the head of the bed to reduce supine hypertension. Intravascular volume should be expanded by increasing dietary fluid and salt. If these nonpharmacologic measures fail, pharmacologic intervention with fludrocortisone acetate and vasoconstricting agents such as midodrine, L-dihydroxyphenylserine, and pseudoephedrine should be introduced. Some patients with intractable symptoms require additional therapy with supplementary agents that include pyridostigmine, yohimbine, desmopressin acetate (DDAVP), and erythropoietin (**Chap. 454**).

CARDIAC SYNCOPE

Cardiac (or cardiovascular) syncope is caused by arrhythmias and structural heart disease. These may occur in combination because structural disease renders the heart more vulnerable to abnormal electrical activity.

Arrhythmias Bradyarrhythmias that cause syncope include those due to severe sinus node dysfunction (e.g., sinus arrest or sinoatrial block) and atrioventricular (AV) block (e.g., Mobitz type II, high-grade, and complete AV block). The bradyarrhythmias due to sinus node dysfunction are often associated with an atrial tachyarrhythmia, a disorder known as the tachycardia-bradycardia syndrome. A prolonged pause following the termination of a tachycardic episode is a frequent cause of syncope in patients with the tachycardia-bradycardia syndrome. Medications of several classes may also cause bradyarrhythmias of sufficient severity to cause syncope. Syncope due to bradycardia or asystole is referred to as a Stokes-Adams attack.

Ventricular tachyarrhythmias frequently cause syncope. The likelihood of syncope with ventricular tachycardia is in part dependent on

the ventricular rate; rates below 200 beats/min are less likely to cause syncope. The compromised hemodynamic function during ventricular tachycardia is caused by ineffective ventricular contraction, reduced diastolic filling due to abbreviated filling periods, loss of AV synchrony, and concurrent myocardial ischemia.

Several disorders associated with cardiac electrophysiologic instability and arrhythmogenesis are due to mutations in ion channel subunit genes. These include the long QT syndrome, Brugada syndrome, and catecholaminergic polymorphic ventricular tachycardia. The long QT syndrome is a genetically heterogeneous disorder associated with prolonged cardiac repolarization and a predisposition to ventricular arrhythmias. Syncope and sudden death in patients with long QT syndrome result from a unique polymorphic ventricular tachycardia called *torsades des pointes* that degenerates into ventricular fibrillation. The long QT syndrome has been linked to genes encoding K⁺ channel α -subunits, K⁺ channel β -subunits, voltage-gated Na⁺ channel, and a scaffolding protein, ankyrin B (ANK2). Brugada syndrome is characterized by idiopathic ventricular fibrillation in association with right ventricular electrocardiogram (ECG) abnormalities without structural heart disease. This disorder is also genetically heterogeneous, although it is most frequently linked to mutations in the Na⁺ channel α -subunit, SCN5A. Catecholaminergic polymorphic tachycardia is an inherited, genetically heterogeneous disorder associated with exercise- or stress-induced ventricular arrhythmias, syncope, or sudden death. Acquired QT interval prolongation, most commonly due to drugs, may also result in ventricular arrhythmias and syncope. **These disorders are discussed in detail in Chap. 277.**

Structural Disease Structural heart disease (e.g., valvular disease, myocardial ischemia, hypertrophic and other cardiomyopathies, cardiac masses such as atrial myxoma, and pericardial effusions) may lead to syncope by compromising cardiac output. Structural disease may also contribute to other pathophysiologic mechanisms of syncope. For example, cardiac structural disease may predispose to arrhythmogenesis; aggressive treatment of cardiac failure with diuretics and/or vasodilators may lead to orthostatic hypotension; and inappropriate reflex vasodilation may occur with structural disorders such as aortic stenosis and hypertrophic cardiomyopathy, possibly provoked by increased ventricular contractility.

TREATMENT CARDIAC SYNCOPES

Treatment of cardiac disease depends on the underlying disorder. Therapies for arrhythmias include cardiac pacing for sinus node disease and AV block, and ablation, antiarrhythmic drugs, and cardioverter-defibrillators for atrial and ventricular tachyarrhythmias. These disorders are best managed by physicians with specialized skills in this area.

APPROACH TO THE PATIENT:

Syncope

DIFFERENTIAL DIAGNOSIS

Syncope is easily diagnosed when the characteristic features are present; however, several disorders with transient real or apparent loss of consciousness may create diagnostic confusion.

Generalized and partial seizures may be confused with syncope; however, there are a number of differentiating features. Whereas tonic-clonic movements are the hallmark of a generalized seizure, myoclonic and other movements also may occur in up to 90% of syncopal episodes. Myoclonic jerks associated with syncope may be multifocal or generalized. They are typically arrhythmic and of short duration (<30 s). Mild flexor and extensor posturing also may occur. Partial or partial-complex seizures with secondary generalization are usually preceded by an aura, commonly an unpleasant smell; fear; anxiety; abdominal discomfort; or other visceral sensations. These phenomena should be differentiated from the premonitory features of syncope.

Autonomic manifestations of seizures (autonomic epilepsy) may provide a more difficult diagnostic challenge. Autonomic seizures have cardiovascular, gastrointestinal, pulmonary, urogenital, pupillary, and cutaneous manifestations that are similar to the premonitory features of syncope. Furthermore, the cardiovascular manifestations of autonomic epilepsy include clinically significant tachycardias and bradycardias that may be of sufficient magnitude to cause loss of consciousness. The presence of accompanying non-autonomic auras may help differentiate these episodes from syncope.

Loss of consciousness associated with a seizure usually lasts longer than 5 min and is associated with prolonged postictal drowsiness and disorientation, whereas reorientation occurs almost immediately after a syncopal event. Muscle aches may occur after both syncope and seizures, although they tend to last longer and be more severe following a seizure. Seizures, unlike syncope, are rarely provoked by emotions or pain. Incontinence of urine may occur with both seizures and syncope; however, fecal incontinence occurs very rarely with syncope.

Hypoglycemia may cause transient loss of consciousness, typically in individuals with type 1 or type 2 diabetes treated with insulin. The clinical features associated with impending or actual hypoglycemia include tremor, palpitations, anxiety, diaphoresis, hunger, and paresthesias. These symptoms are due to autonomic activation to counter the falling blood glucose. Hunger, in particular, is not a typical premonitory feature of syncope. Hypoglycemia also impairs neuronal function, leading to fatigue, weakness, dizziness, and cognitive and behavioral symptoms. Diagnostic difficulties may occur in individuals in strict glycemic control; repeated hypoglycemia impairs the counterregulatory response and leads to a loss of the characteristic warning symptoms that are the hallmark of hypoglycemia.

Patients with cataplexy experience an abrupt partial or complete loss of muscular tone triggered by strong emotions, typically anger or laughter. Unlike syncope, consciousness is maintained throughout the attacks, which typically last between 30 s and 2 min. There are no premonitory symptoms. Cataplexy occurs in 60–75% of patients with narcolepsy.

The clinical interview and interrogation of eyewitnesses usually allow differentiation of syncope from falls due to vestibular dysfunction, cerebellar disease, extrapyramidal system dysfunction, and other gait disorders. If the fall is accompanied by head trauma, a postconcussive syndrome, amnesia for the precipitating events, and/or the presence of loss of consciousness may contribute to diagnostic difficulty.

Apparent loss of consciousness can be a manifestation of psychiatric disorders such as generalized anxiety, panic disorders, major depression, and somatization disorder. These possibilities should be considered in individuals who faint frequently without prodromal symptoms. Such patients are rarely injured despite numerous falls. There are no clinically significant hemodynamic changes concurrent with these episodes. In contrast, transient loss of consciousness due to vasovagal syncope precipitated by fear, stress, anxiety, and emotional distress is accompanied by hypotension, bradycardia, or both.

INITIAL EVALUATION

The goals of the initial evaluation are to determine whether the transient loss of consciousness was due to syncope; to identify the cause; and to assess risk for future episodes and serious harm (Table 27-1). The initial evaluation should include a detailed history, thorough questioning of eyewitnesses, and a complete physical and neurologic examination. Blood pressure and heart rate should be measured in the supine position and after 3 min of standing to determine whether orthostatic hypotension is present. An ECG should be performed if there is suspicion of syncope due to an arrhythmia or underlying cardiac disease. Relevant electrocardiographic abnormalities include bradyarrhythmias or tachyarrhythmias, AV block, ischemia, old myocardial infarction, long QT syndrome, and bundle branch block. This initial assessment will lead to the identification of a cause of syncope in approximately

50% of patients and also allows stratification of patients at risk for cardiac mortality.

Laboratory Tests Baseline laboratory blood tests are rarely helpful in identifying the cause of syncope. Blood tests should be performed when specific disorders, e.g., myocardial infarction, anemia, and secondary autonomic failure, are suspected (Table 27-2).

Autonomic Nervous System Testing (Chap. 454) Autonomic testing, including tilt-table testing, can be performed in specialized centers. Autonomic testing is helpful to uncover objective evidence of autonomic failure and also to demonstrate a predisposition to neurally mediated syncope. Autonomic testing includes assessments of parasympathetic autonomic nervous system function (e.g., heart rate variability to deep respiration and a Valsalva maneuver), sympathetic cholinergic function (e.g., thermoregulatory sweat response and quantitative sudomotor axon reflex test), and sympathetic adrenergic function (e.g., blood pressure response to a Valsalva maneuver and a tilt-table test with beat-to-beat blood pressure measurement). The hemodynamic abnormalities demonstrated on tilt-table test (Figs. 27-3 and 27-4) may be useful in distinguishing orthostatic hypotension due to autonomic failure from the hypotensive bradycardic response of neurally mediated syncope. Similarly, the tilt-table test may help identify patients with syncope due to immediate or delayed orthostatic hypotension.

Carotid sinus massage should be considered in patients with symptoms suggestive of carotid sinus syncope and in patients over age 50 years with recurrent syncope of unknown etiology. This test should only be carried out under continuous ECG and blood pressure monitoring and should be avoided in patients with carotid bruits, plaques, or stenosis.

Cardiac Evaluation ECG monitoring is indicated for patients with a high pretest probability of arrhythmia causing syncope. Patients should be monitored in hospital if the likelihood of a life-threatening arrhythmia is high, e.g., patients with severe structural or coronary artery disease, nonsustained ventricular tachycardia, trifascicular heart block, prolonged QT interval, Brugada syndrome ECG pattern, or family history of sudden cardiac death (Table 27-1). Outpatient Holter monitoring is recommended for patients who experience frequent syncopal episodes (one or more per week), whereas loop recorders, which continually record and erase cardiac rhythm, are indicated for patients with suspected arrhythmias with low risk of sudden cardiac death. Loop recorders may be external (recommended for evaluation of episodes that occur at a frequency of greater than one per month) or implantable (if syncope occurs less frequently).

Echocardiography should be performed in patients with a history of cardiac disease or if abnormalities are found on physical examination or the ECG. Echocardiographic diagnoses that may be responsible for syncope include aortic stenosis, hypertrophic cardiomyopathy, cardiac tumors, aortic dissection, and pericardial tamponade. Echocardiography also has a role in risk stratification based on the left ventricular ejection fraction.

Treadmill exercise testing with ECG and blood pressure monitoring should be performed in patients who have experienced syncope during or shortly after exercise. Treadmill testing may help identify exercise-induced arrhythmias (e.g., tachycardia-related AV block) and exercise-induced exaggerated vasodilation.

Electrophysiologic studies are indicated in patients with structural heart disease and ECG abnormalities in whom noninvasive investigations have failed to yield a diagnosis. Electrophysiologic studies have low sensitivity and specificity and should only be performed when a high pretest probability exists. Currently, this test is rarely performed to evaluate patients with syncope.

Psychiatric Evaluation Screening for psychiatric disorders may be appropriate in patients with recurrent unexplained syncope episodes. Tilt-table testing, with demonstration of symptoms in the absence of hemodynamic change, may be useful in reproducing syncope in patients with suspected psychogenic syncope.

28

Dizziness and Vertigo

Mark F. Walker, Robert B. Daroff

Dizziness is an imprecise symptom used to describe a variety of sensations that include vertigo, light-headedness, faintness, and imbalance. When used to describe a sense of spinning or other motion, dizziness is designated as *vertigo*. Vertigo may be physiologic, occurring during or after a sustained head rotation, or it may be pathologic, due to vestibular dysfunction. The term *light-headedness* is commonly applied to presyncopal sensations due to brain hypoperfusion but also may refer to disequilibrium and imbalance. A challenge to diagnosis is that patients often have difficulty distinguishing among these various symptoms, and the words they choose do not reliably indicate the underlying etiology.

There are a number of potential causes of dizziness. Vascular disorders cause presyncopal dizziness as a result of cardiac dysrhythmia, orthostatic hypotension, medication effects, or other causes. Such presyncopal sensations vary in duration; they may increase in severity until loss of consciousness occurs, or they may resolve before loss of consciousness if the cerebral ischemia is corrected. Faintness and syncope, which are discussed in detail in [Chap. 27](#), should always be considered when one is evaluating patients with brief episodes of dizziness or dizziness that occurs with upright posture.

Vestibular causes of dizziness (vertigo or imbalance) may be due to peripheral lesions that affect the labyrinths or vestibular nerves or to involvement of the central vestibular pathways. They may be paroxysmal or due to a fixed unilateral or bilateral vestibular deficit. Acute unilateral lesions cause vertigo due to a sudden imbalance in vestibular inputs from the two labyrinths. Bilateral lesions cause imbalance and instability of vision (*oscillopsia*) when the head moves. Other causes of dizziness include nonvestibular imbalance and gait disorders (e.g., loss of proprioception from sensory neuropathy, parkinsonism) and anxiety.

When evaluating patients with dizziness, questions to consider include the following: (1) Is it dangerous (e.g., arrhythmia, transient ischemic attack/stroke)? (2) Is it vestibular? (3) If vestibular, is it peripheral or central? A careful history and examination often provide sufficient information to answer these questions and determine whether additional studies or referral to a specialist is necessary.

APPROACH TO THE PATIENT:

Dizziness

HISTORY

When a patient presents with dizziness, the first step is to delineate more precisely the nature of the symptom. In the case of vestibular disorders, the physical symptoms depend on whether the lesion is unilateral or bilateral, and whether it is acute or chronic and progressive. Vertigo, an illusion of self or environmental motion, implies asymmetry of vestibular inputs from the two labyrinths or in their central pathways that is usually acute. Symmetric bilateral vestibular hypofunction causes imbalance but no vertigo. Because of the ambiguity in patients' descriptions of their symptoms, diagnosis based simply on symptom characteristics is typically unreliable. The history should focus closely on other features, including whether this is the first attack, the duration of this and any prior episodes, provoking factors, and accompanying symptoms.

Dizziness can be divided into episodes that last for seconds, minutes, hours, or days. Common causes of brief dizziness (seconds) include benign paroxysmal positional vertigo (BPPV) and orthostatic hypotension, both of which typically are provoked by changes in head and body position. Attacks of vestibular migraine and Ménière's disease often last hours. When episodes are of intermediate duration (minutes), transient ischemic attacks of the posterior circulation should be considered, although migraine and a number of other causes are also possible.

Symptoms that accompany vertigo may be helpful in distinguishing peripheral vestibular lesions from central causes. Unilateral hearing loss and other aural symptoms (ear pain, pressure, fullness) typically point to a peripheral cause. Because the auditory pathways quickly become bilateral upon entering the brainstem, central lesions are unlikely to cause unilateral hearing loss, unless the lesion lies near the root entry zone of the auditory nerve. Symptoms such as double vision, numbness, and limb ataxia suggest a brainstem or cerebellar lesion.

EXAMINATION

Because dizziness and imbalance can be a manifestation of a variety of neurologic disorders, the neurologic examination is important in the evaluation of these patients. Particular focus should be given to assessment of eye movements, vestibular function, and hearing. The range of eye movements and whether they are equal in each eye should be observed. Peripheral eye movement disorders (e.g., cranial neuropathies, eye muscle weakness) are usually disconjugate (different in the two eyes). One should check pursuit (the ability to follow a smoothly moving target) and saccades (the ability to look back and forth accurately between two targets). Poor pursuit or inaccurate (dysmetric) saccades usually indicates central pathology, often involving the cerebellum. Finally, one should look for spontaneous nystagmus, an involuntary back-and-forth movement of the eyes. Nystagmus is most often of the jerk type, in which a slow drift (slow phase) in one direction alternates with a rapid saccadic movement (quick phase or fast phase) in the opposite direction that resets the position of the eyes in the orbits. Except in the case of acute vestibulopathy (e.g., vestibular neuritis), if primary position nystagmus is easily seen in the light, it is probably due to a central cause. Two forms of nystagmus that are characteristic of lesions of the cerebellar pathways are vertical nystagmus with downward fast phases (downbeat nystagmus) and horizontal nystagmus that changes direction with gaze (gaze-evoked nystagmus). By contrast, peripheral lesions typically cause unidirectional horizontal nystagmus. Use of Frenzel eyeglasses (self-illuminated goggles with convex lenses that blur the patient's vision but allow the examiner to see the eyes greatly magnified) can aid in the detection of peripheral vestibular nystagmus, because they reduce the patient's ability to use visual fixation to suppress nystagmus. **Table 28-1** outlines key findings that help distinguish peripheral from central causes of vertigo.

The most useful bedside test of peripheral vestibular function is the head impulse test, in which the vestibuloocular reflex (VOR) is assessed with small-amplitude (~20 degrees) rapid head rotations. While the patient fixates on a target, the head is rotated to the left or right. If the VOR is deficient, the rotation is followed by a catch-up saccade in the opposite direction (e.g., a leftward saccade after a rightward rotation). The head impulse test can identify both unilateral (catch-up saccades after rotations toward the weak side) and bilateral vestibular hypofunction (catch-up saccades after rotations in both directions).

All patients with episodic dizziness, especially if provoked by positional change, should be tested with the Dix-Hallpike maneuver. The patient begins in a sitting position with the head turned

45 degrees; holding the back of the head, the examiner then lowers the patient into a supine position with the head extended backward by about 20 degrees while watching the eyes. Posterior canal BPPV can be diagnosed confidently if transient upbeating-torsional nystagmus is seen. If no nystagmus is observed after 15–20 s, the patient is raised to the sitting position, and the procedure is repeated with the head turned to the other side. Again, Frenzel goggles may improve the sensitivity of the test.

Dynamic visual acuity is a functional test that can be useful in assessing vestibular function. Visual acuity is measured with the head still and when the head is rotated back and forth by the examiner (about 1–2 Hz). A drop in visual acuity during head motion of more than one line on a near card or Snellen chart is abnormal and indicates vestibular dysfunction.

ANCILLARY TESTING

The choice of ancillary tests should be guided by the history and examination findings. Audiometry should be performed whenever a vestibular disorder is suspected. Unilateral sensorineural hearing loss supports a peripheral disorder (e.g., vestibular schwannoma). Predominantly low-frequency hearing loss is characteristic of Ménière's disease. Electronystagmography or videonystagmography includes recordings of spontaneous nystagmus (if present) and measurement of positional nystagmus. Caloric testing assesses the responses of the two horizontal semicircular canals. The test battery often includes recording of saccades and pursuit to assess central ocular motor function. Neuroimaging is important if a central vestibular disorder is suspected. In addition, patients with unexplained unilateral hearing loss or vestibular hypofunction should undergo magnetic resonance imaging (MRI) of the internal auditory canals, including administration of gadolinium, to rule out a schwannoma.

DIFFERENTIAL DIAGNOSIS AND TREATMENT

Treatment of vestibular symptoms should be driven by the underlying diagnosis. Simply treating dizziness with vestibular suppressant medications is often not helpful and may make the symptoms worse and prolong recovery. The diagnostic and specific treatment approaches for the most commonly encountered vestibular disorders are discussed below.

ACUTE PROLONGED VERTIGO (VESTIBULAR NEURITIS)

An acute unilateral vestibular lesion causes constant vertigo, nausea, vomiting, oscillopsia (motion of the visual scene), and imbalance. These symptoms are due to a sudden asymmetry of inputs from the two labyrinths or in their central connections, simulating a continuous rotation of the head. Unlike BPPV, continuous vertigo persists even when the head remains still.

When a patient presents with an acute vestibular syndrome, the most important question is whether the lesion is central (e.g., a cerebellar or brainstem infarct or hemorrhage), which may be life-threatening, or peripheral, affecting the vestibular nerve or labyrinth (vestibular neuritis). Attention should be given to any symptoms or signs that point to central dysfunction (diplopia, weakness or numbness, dysarthria). The pattern of spontaneous nystagmus, if present, may be helpful (Table 28-1). If the head impulse test is normal, an acute peripheral vestibular lesion is unlikely. A central lesion cannot always be excluded with certainty based on symptoms and examination alone; thus, older patients with vascular risk factors who present with an acute vestibular syndrome should be evaluated for the possibility of stroke even when there are no specific findings that indicate a central lesion.

Most patients with vestibular neuritis recover spontaneously, but glucocorticoids can improve outcome if administered within 3 days of symptom onset. Antiviral medications are of no proven benefit and are not typically given unless there is evidence to suggest herpes zoster oticus (Ramsay Hunt syndrome). Vestibular suppressant medications may reduce acute symptoms but should be avoided after the first several days because they may impede central compensation and recovery. Patients should be encouraged to resume a normal level of activity as soon as possible, and directed vestibular rehabilitation therapy may accelerate improvement.

TABLE 28-1 FEATURES OF PERIPHERAL AND CENTRAL VERTIGO

- Nystagmus from an acute peripheral lesion is unidirectional, with fast phases beating away from the ear with the lesion. Nystagmus that changes direction with gaze is due to a central lesion.
- Transient mixed vertical-torsional nystagmus occurs in BPPV, but pure vertical or pure torsional nystagmus is a central sign.
- Nystagmus from a peripheral lesion may be inhibited by visual fixation, whereas central nystagmus is not suppressed.
- Absence of a head impulse sign in a patient with acute prolonged vertigo should suggest a central cause.
- Unilateral hearing loss suggests peripheral vertigo. Findings such as diplopia, dysarthria, and limb ataxia suggest a central disorder.

BENIGN PAROXYSMAL POSITIONAL VERTIGO

BPPV is a common cause of recurrent vertigo. Episodes are brief (<1 min and typically 15–20 s) and are always provoked by changes in head position relative to gravity, such as lying down, rolling over in bed, rising from a supine position, and extending the head to look upward. The attacks are caused by free-floating otoconia (calcium carbonate crystals) that have been dislodged from the utricular macula and have moved into one of the semicircular canals, usually the posterior canal. When head position changes, gravity causes the otoconia to move within the canal, producing vertigo and nystagmus. With posterior canal BPPV, the nystagmus beats upward and torsionally (the upper poles of the eyes beat toward the affected lower ear). Less commonly, the otoconia enter the horizontal canal, resulting in a horizontal nystagmus when the patient is lying with either ear down. Superior (also called anterior) canal involvement is rare. BPPV is treated with repositioning maneuvers that use gravity to remove the otoconia from the semicircular canal. For posterior canal BPPV, the Epley maneuver (Fig. 28-1) is the most commonly used procedure. For more refractory cases of BPPV, patients can be taught a variant of this maneuver that they can perform alone at home. A demonstration of the Epley maneuver is available online (<http://www.dizziness-and-balance.com/disorders/bppv/bppv.html>).

VESTIBULAR MIGRAINE

Vestibular symptoms occur frequently in migraineurs, sometimes as a headache aura but usually independent of headache. The duration of vertigo may be from minutes to hours, and some patients also experience more prolonged periods of disequilibrium (lasting days to weeks). Motion sensitivity and sensitivity to visual motion (e.g., movies) are common in patients with vestibular migraine. Although data from controlled studies are generally lacking, vestibular migraine typically is treated with medications that are used for prophylaxis of migraine headaches. Antiemetics may be helpful to relieve symptoms at the time of an attack.

MÉNIÈRE'S DISEASE

Attacks of Ménière's disease consist of vertigo and hearing loss, as well as pain, pressure, and/or fullness in the affected ear. The low-frequency

hearing loss and aural symptoms are key features that distinguish Ménière's disease from other peripheral vestibulopathies and from vestibular migraine. Audiometry at the time of an attack shows a characteristic asymmetric low-frequency hearing loss; hearing commonly improves between attacks, although permanent hearing loss may eventually occur. Ménière's disease is thought to be due to excess fluid (endolymph) in the inner ear; hence the term *endolymphatic hydrops*. Patients suspected of having Ménière's disease should be referred to an otolaryngologist for further evaluation. Diuretics and sodium restriction are the initial treatments. If attacks persist, injections of gentamicin into the middle ear are typically the next line of therapy. Full ablative procedures (vestibular nerve section, labyrinthectomy) are seldom required.

VESTIBULAR SCHWANNOMA

Vestibular schwannomas (sometimes termed *acoustic neuromas*) and other tumors at the cerebellopontine angle cause slowly progressive unilateral sensorineural hearing loss and vestibular hypofunction. These patients typically do not have vertigo, because the gradual vestibular deficit is compensated centrally as it develops. The diagnosis often is not made until there is sufficient hearing loss to be noticed. The examination will show a deficient response to the head impulse test when the head is rotated toward the affected side. As noted above, patients with unexplained unilateral sensorineural hearing loss or vestibular hypofunction require MRI of the internal auditory canals to look for a schwannoma.

BILATERAL VESTIBULAR HYPOFUNCTION

Patients with bilateral loss of vestibular function also typically do not have vertigo, because vestibular function is lost on both sides simultaneously, and there is no asymmetry of vestibular input. Symptoms include loss of balance, particularly in the dark, where vestibular input is most critical, and oscillopsia during head movement, such as while walking or riding in a car. Bilateral vestibular hypofunction may be (1) idiopathic and progressive, (2) part of a neurodegenerative disorder, or (3) iatrogenic, due to medication ototoxicity (most commonly

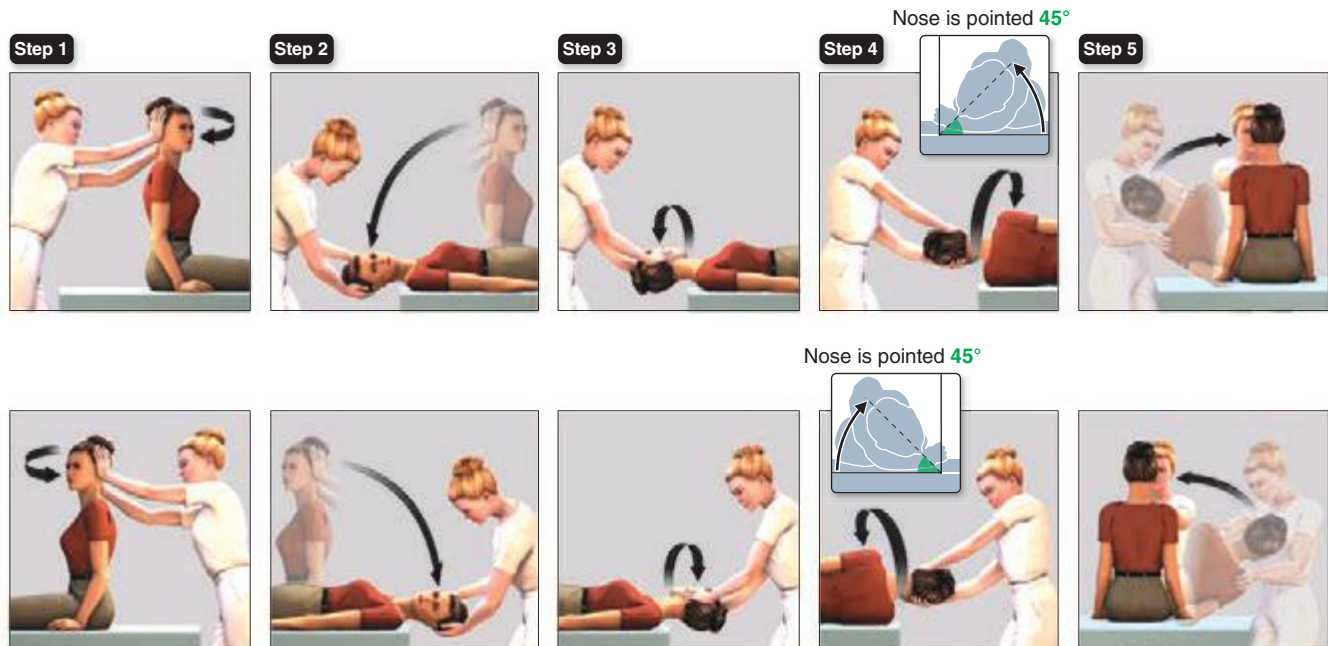


FIGURE 28-1 Modified Epley maneuver for treatment of benign paroxysmal positional vertigo of the right (*top panels*) and left (*bottom panels*) posterior semicircular canals. **Step 1.** With the patient seated, turn the head 45 degrees toward the affected ear. **Step 2.** Keeping the head turned, lower the patient to the head-hanging position and hold for at least 30 s and until nystagmus disappears. **Step 3.** Without lifting the head, turn it 90 degrees toward the other side. Hold for another 30 s. **Step 4.** Rotate the patient onto her side while turning the head another 90 degrees, so that the nose is pointed down 45 degrees. Hold again for 30 s. **Step 5.** Have the patient sit up on the side of the table. After a brief rest, the maneuver should be repeated to confirm successful treatment. (Figure adapted from <http://www.dizziness-and-balance.com/disorders/bppv/movies/Epley-480x640.avi>.)

gentamicin or other aminoglycoside antibiotics). Other causes include bilateral vestibular schwannomas (neurofibromatosis type 2), autoimmune disease, superficial siderosis, and meningeal-based infection or tumor. It also may occur in patients with peripheral polyneuropathy; in these patients, both vestibular loss and impaired proprioception may contribute to poor balance. Finally, unilateral processes such as vestibular neuritis and Ménière's disease may involve both ears sequentially, resulting in bilateral vestibulopathy.

Examination findings include diminished *dynamic visual acuity* (see above) due to loss of stable vision when the head is moving, abnormal head impulse responses in both directions, and a Romberg sign. Responses to caloric testing are reduced. Patients with bilateral vestibular hypofunction should be referred for vestibular rehabilitation therapy. Vestibular suppressant medications should not be used, as they will increase the imbalance. Evaluation by a neurologist is important not only to confirm the diagnosis but also to consider any other associated neurologic abnormalities that may clarify the etiology.

CENTRAL VESTIBULAR DISORDERS

Central lesions causing vertigo typically involve vestibular pathways in the brainstem and/or cerebellum. They may be due to discrete lesions, such as from ischemic or hemorrhagic stroke (Chap. 446), demyelination (Chap. 458), or tumors (Chap. 118), or they may be due to neurodegenerative conditions that include the vestibulocerebellum (Chap. 448). Subacute cerebellar degeneration may be due to immune, including paraneoplastic, processes (Chaps. 122 and 450). Table 28-1 outlines important features of the history and examination that help to identify central vestibular disorders. Acute central vertigo is a medical emergency, due to the possibility of life-threatening stroke or hemorrhage. All patients with suspected central vestibular disorders should undergo brain MRI, and the patient should be referred for full neurologic evaluation.

PSYCHOSOMATIC DIZZINESS/VERTIGO

Psychological factors play an important role in chronic dizziness. First, dizziness may be a somatic manifestation of a psychiatric condition such as major depression, anxiety, or panic disorder (Chap. 465e). Second, patients may develop anxiety and autonomic symptoms as a consequence or comorbidity of an independent vestibular disorder. One particular form of this has been termed variously *phobic postural vertigo*, *psychophysiologic vertigo*, or *chronic subjective dizziness*. These patients have a chronic feeling (months or longer) of dizziness and disequilibrium, an increased sensitivity to self-motion and visual motion (e.g., movies), and a particular intensification of symptoms when moving through complex visual environments such as supermarkets (visual vertigo). Although there may be a past history of an acute vestibular disorder (e.g., vestibular neuritis), the neurootologic examination and vestibular testing are normal or indicative of a compensated vestibular deficit, indicating that the ongoing subjective dizziness cannot be explained by a primary vestibular disorder. Anxiety disorders are particularly common in patients with chronic dizziness and contribute substantially to the morbidity. Thus, treatment with anti-anxiety medications (selective serotonin reuptake inhibitors [SSRIs]) and cognitive-behavioral therapy may be helpful. Vestibular rehabilitation therapy is also sometimes beneficial. Vestibular suppressant medications generally should be avoided. This condition should be suspected when the patient states, "My dizziness is so bad, I'm afraid to leave my house" (agoraphobia).

TREATMENT VERTIGO

Table 28-2 provides a list of commonly used medications for suppression of vertigo. As noted, these medications should be reserved for short-term control of active vertigo, such as during the first few days of acute vestibular neuritis, or for acute attacks of Ménière's disease. They are less helpful for chronic dizziness and, as previously stated, may hinder central compensation. An exception is that benzodiazepines may attenuate psychosomatic dizziness and the associated anxiety, although SSRIs are generally preferable in such patients.

TABLE 28-2 TREATMENT OF VERTIGO

Agent ^a	Dose ^b
Antihistamines	
Meclizine	25–50 mg 3 times daily
Dimenhydrinate	50 mg 1–2 times daily
Promethazine	25 mg 2–3 times daily (also can be given rectally and IM)
Benzodiazepines	
Diazepam	2.5 mg 1–3 times daily
Clonazepam	0.25 mg 1–3 times daily
Anticholinergic	
Scopolamine transdermal ^f	Patch
Physical therapy	
Repositioning maneuvers ^d	
Vestibular rehabilitation	
Other	
Diuretics and/or low-sodium (1000 mg/d) diet ^e	
Antimigrainous drugs ^f	
Methylprednisolone ^g	100 mg daily days 1–3; 80 mg daily days 4–6; 60 mg daily days 7–9; 40 mg daily days 10–12; 20 mg daily days 13–15; 10 mg daily days 16–18, 20, 22
Selective serotonin reuptake inhibitors ^h	

^aAll listed drugs are approved by the U.S. Food and Drug Administration, but most are not approved for the treatment of vertigo. ^bUsual oral (unless otherwise stated) starting dose in adults; a higher maintenance dose can be reached by a gradual increase. ^cFor motion sickness only. ^dFor benign paroxysmal positional vertigo. ^eFor Ménière's disease. ^fFor vestibular migraine. ^gFor acute vestibular neuritis (started within 3 days of onset). ^hFor psychosomatic vertigo.

Vestibular rehabilitation therapy promotes central adaptation processes that compensate for vestibular loss and also may help habituate motion sensitivity and other symptoms of psychosomatic dizziness. The general approach is to use a graded series of exercises that progressively challenge gaze stabilization and balance.

29 Fatigue

Jeffrey M. Gelfand, Vanja C. Douglas

Fatigue is one of the most common symptoms in clinical medicine. It is a prominent manifestation of a number of systemic, neurologic, and psychiatric syndromes, although a precise cause will not be identified in a substantial minority of patients. Fatigue refers to an inherently subjective human experience of physical and mental weariness, sluggishness, and exhaustion. In the context of clinical medicine, fatigue is most typically and practically defined as difficulty initiating or maintaining voluntary mental or physical activity. Nearly everyone who has ever been ill with a self-limited infection has experienced this near-universal symptomatology, and fatigue is usually brought to medical attention only when it is either of unclear cause or the severity is out of proportion with what would be expected for the associated trigger. *Fatigue* should be distinguished from *muscle weakness*, a reduction of neuromuscular power (Chap. 30); most patients complaining of fatigue are not truly weak when direct muscle power is tested. By definition, fatigue is also distinct from *somnolence* and *dyspnea on exertion*, although patients may use the word fatigue to describe those two symptoms. The task facing clinicians when a patient presents with fatigue is to identify an underlying cause if one exists and to develop

a therapeutic alliance, the goal of which is to spare patients expensive and fruitless diagnostic workups and steer them toward effective therapy.

EPIDEMIOLOGY AND GLOBAL CONSIDERATIONS



Variability in the definitions of fatigue and the survey instruments used in different studies makes it difficult to arrive at precise figures about the global burden of fatigue. The point prevalence of fatigue was 6.7% and the lifetime prevalence was 25% in a large National Institute of Mental Health survey of the U.S. general population. In primary care clinics in Europe and the United States, between 10 and 25% of patients surveyed endorsed symptoms of prolonged (present for >1 month) or chronic (present for >6 months) fatigue, but fatigue was the primary reason for seeking medical attention in only a minority of patients. In a community survey of women in India, 12% reported chronic fatigue. By contrast, the prevalence of chronic fatigue syndrome, as defined by the U.S. Centers for Disease Control and Prevention, is low (**Chap. 464e**).

DIFFERENTIAL DIAGNOSIS

Psychiatric Disease Fatigue is a common somatic manifestation of many major psychiatric syndromes, including depression, anxiety, and somatoform disorders. Psychiatric symptoms are reported in more than three-quarters of patients with unexplained chronic fatigue. Even in patients with systemic or neurologic syndromes in which fatigue is independently recognized as a manifestation of disease, comorbid psychiatric symptoms or disease may still be an important source of interaction.

Neurologic Disease Patients complaining of fatigue often say they feel weak, but upon careful examination, objective muscle weakness is rarely discernible. If found, muscle weakness must then be localized to the central nervous system, peripheral nervous system, neuromuscular junction, or muscle and the appropriate follow-up studies obtained (**Chap. 30**). *Fatigability* of muscle power is a cardinal manifestation of some neuromuscular disorders such as myasthenia gravis and can be distinguished from *fatigue* by finding clinically apparent diminution of the amount of force that a muscle generates upon repeated contraction (**Chap. 461**). Fatigue is one of the most common and bothersome symptoms reported in multiple sclerosis (MS) (**Chap. 458**), affecting nearly 90% of patients; fatigue in MS can persist between MS attacks and does not necessarily correlate with magnetic resonance imaging (MRI) disease activity. Fatigue is also increasingly identified as a troublesome feature of many other neurodegenerative diseases, including Parkinson's disease, central dysautonomias, and amyotrophic lateral sclerosis. Poststroke fatigue is a well-described but poorly understood entity with a widely varying prevalence. Episodic fatigue can be a premonitory symptom of migraine. Fatigue is also a frequent result of traumatic brain injury, often occurring in association with depression and sleep disorders.

Sleep Disorders Obstructive sleep apnea is an important cause of excessive daytime sleepiness in association with fatigue and should be investigated using overnight polysomnography, particularly in those with prominent snoring, obesity, or other predictors of obstructive sleep apnea (**Chap. 319**). Whether the cumulative sleep deprivation that is common in modern society contributes to clinically apparent fatigue is not known (**Chap. 38**).

Endocrine Disorders Fatigue, sometimes in association with true muscle weakness, can be a heralding symptom of hypothyroidism, particularly in the context of hair loss, dry skin, cold intolerance, constipation, and weight gain. Fatigue in association with heat intolerance, sweating, and palpitations is typical of hyperthyroidism. Adrenal insufficiency can also manifest with unexplained fatigue as a primary or prominent symptom, often in association with anorexia, weight loss, nausea, myalgias, and arthralgias; hyponatremia and hyperkalemia may be present at time of diagnosis. Mild hypercalcemia can cause fatigue, which may be relatively vague, whereas severe hypercalcemia can lead

to lethargy, stupor, and coma. Both hypoglycemia and hyperglycemia can cause lethargy, often in association with confusion; chronic diabetes, particularly type 1 diabetes, is also associated with fatigue independent of glucose levels. Fatigue may also accompany Cushing's disease, hypoadosteronism, and hypogonadism.

Liver and Kidney Disease Both chronic liver failure and chronic kidney disease can cause fatigue. Over 80% of hemodialysis patients complain of fatigue, which makes this one of the most common patient-reported symptoms in chronic kidney disease.

Obesity Obesity is associated with fatigue and sleepiness independent of the presence of obstructive sleep apnea. Obese patients undergoing bariatric surgery experience improvement in daytime sleepiness sooner than would be expected if the improvement were solely the result of weight loss and resolution of sleep apnea. A number of other factors common in obese patients are likely contributors as well, including depression, physical inactivity, and diabetes.

Malnutrition Although fatigue can be a presenting feature of malnutrition, nutritional status may also be an important comorbidity and contributor to fatigue in other chronic illnesses, including cancer-associated fatigue.

Infection Both acute and chronic infections commonly lead to fatigue as part of the broader infectious syndrome. Evaluation for undiagnosed infection as the cause of unexplained fatigue, and particularly prolonged or chronic fatigue, should be guided by the history, physical examination, and infectious risk factors, with particular attention to risk for tuberculosis, HIV, chronic hepatitis B and C, and endocarditis. Infectious mononucleosis may cause prolonged fatigue that persists for weeks to months following the acute illness, but infection with the Epstein-Barr virus is only very rarely the cause of unexplained chronic fatigue.

Drugs Many medications, drug use, drug withdrawal, and chronic alcohol use can all lead to fatigue. Medications that are more likely to be causative in this context include antidepressants, antipsychotics, anxiolytics, opiates, antispasticity agents, antiseizure agents, and beta blockers.

Cardiovascular and Pulmonary Fatigue is one of the most taxing patient-reported symptoms of congestive heart failure and chronic obstructive pulmonary disease and negatively affects quality of life.

Malignancy Fatigue, particularly in association with unexplained unintended weight loss, can be a sign of occult malignancy, but this is only rarely identified as causative in patients with unexplained chronic fatigue in the absence of other telltale signs or symptoms. Cancer-related fatigue is experienced by 40% of patients at time of diagnosis and greater than 80% of patients later in the disease course.

Hematologic Chronic or progressive anemia may present with fatigue, sometimes in association with exertional tachycardia and breathlessness. Anemia may also contribute to fatigue in chronic illness. Low serum ferritin in the absence of anemia may also cause fatigue that is reversible with iron replacement.

Systemic Inflammatory/Rheumatologic Disorders Fatigue is a prominent complaint in many chronic inflammatory disorders, including systemic lupus erythematosus, polymyalgia rheumatica, rheumatoid arthritis, inflammatory bowel disease, antineutrophil cytoplasmic antibody (ANCA)-associated vasculitis, sarcoidosis, and Sjögren's syndrome, but is not usually an isolated symptom.

Pregnancy Fatigue is very commonly reported by women during all stages of pregnancy and postpartum.

Disorders of Unclear Cause Chronic fatigue syndrome (**Chap. 464e**) and fibromyalgia (**Chap. 396**) incorporate chronic fatigue as part of the syndromic definition when present in association with a number of other inclusion and exclusion criteria, as discussed in detail in

their respective chapters. The pathophysiology of each is unknown. Idiopathic chronic fatigue is used to describe the syndrome of unexplained chronic fatigue in the absence of enough additional clinical features to meet the diagnostic criteria for chronic fatigue syndrome.

APPROACH TO THE PATIENT: Fatigue

A detailed history focusing on the quality, pattern, time-course, associated symptoms, and alleviating factors of fatigue is critical in defining the syndrome, determining whether fatigue is the appropriate designation, determining whether the symptoms are acute or chronic, and determining whether fatigue is primarily mental, physical, or both in order to direct further evaluation and treatment. The review of systems should attempt to distinguish fatigue from excessive daytime sleepiness, dyspnea on exertion, exercise intolerance, and muscle weakness. The presence of fever, chills, night sweats, or weight loss should raise suspicion for an occult infection or malignancy. A careful review of prescription, over-the-counter, herbal, and recreational drug and alcohol use is mandatory. Circumstances surrounding the onset of symptoms and potential triggers should be investigated. The social history is important, with attention paid to job stress and work hours, the social support network, and domestic affairs including a screen for intimate partner violence. Sleep habits and sleep hygiene should be questioned. The impact of fatigue on daily functioning is important to understand the patient's experience and gauge recovery and the success of treatment.

The physical examination of patients with fatigue is guided by the history and differential diagnosis. A detailed mental status examination should be performed with particular attention to symptoms of depression and anxiety. A formal neurologic examination is required to determine whether objective muscle weakness is present. This is usually a straightforward exercise, although occasionally patients with fatigue have difficulty sustaining effort against resistance and sometimes report that generating full power requires substantial mental effort. On confrontational testing, they are able to generate full power for only a brief period before suddenly giving way to the examiner. This type of weakness is often referred to as *breakaway weakness* and may or may not be associated with pain. This is contrasted with weakness due to lesions in the motor tracts or lower motor unit, in which the patient's resistance can be overcome in a smooth and steady fashion and full power can never be generated. Occasionally, a patient may demonstrate fatigable weakness, in which power is full when first tested but becomes weak upon repeat evaluation without interval rest. Fatigable weakness, which usually indicates a problem of neuromuscular transmission, never has the sudden breakaway quality that one occasionally observes in patients with fatigue. If the presence or absence of muscle weakness cannot be determined with the physical examination, electromyography with nerve conduction studies can be a helpful ancillary test.

The general physical examination should screen for signs of cardiopulmonary disease, malignancy, lymphadenopathy, organomegaly, infection, liver failure, kidney disease, malnutrition, endocrine abnormalities, and connective tissue disease. Although the diagnostic yield of the general physical examination may be relatively low in the context of evaluation of unexplained chronic fatigue, elucidating the cause of 2% of cases in one prospective analysis, the yield of a detailed neuropsychiatric and mental status evaluation is likely to be much higher, revealing a potential explanation for fatigue in up to 75–80% of patients in some series. Furthermore, the rite of physical examination demonstrates a thorough and systematic approach to the patient's complaint and helps build trust and a therapeutic alliance.

Laboratory testing is likely to identify the cause of chronic fatigue in only about 5% of cases. Beyond a few standard screening tests, laboratory evaluation should be guided by the history and physical examination; extensive testing is more likely to lead to false-positive

results that require explanation and unnecessary investigation and should be avoided in lieu of frequent clinical follow-up. A reasonable approach to screening includes a complete blood count with differential (to screen for anemia, infection, and malignancy), electrolytes (including sodium, potassium, and calcium), glucose, renal function, liver function, and thyroid function. Testing for HIV and adrenal function can also be considered. Published guidelines defining chronic fatigue syndrome also recommend an erythrocyte sedimentation rate (ESR) as part of the evaluation for mimics, but unless the value is very high, such nonspecific testing in the absence of other features is unlikely to clarify the situation. Routine screening with an antinuclear antibody (ANA) test is also unlikely to be informative in isolation and is frequently positive at low titers in otherwise healthy adults. Additional unfocused studies, such as whole-body imaging scans, are usually not indicated; in addition to their inconvenience, potential risk, and cost, they often reveal unrelated incidental findings that can prolong the workup unnecessarily.

TREATMENT FATIGUE

The first priority of treatment is to address the underlying disorder or disorders that account for fatigue, because this can be curative in select contexts and palliative in others. Unfortunately, in many chronic illnesses, fatigue may be refractory to traditional disease-modifying therapies, and it is important in such cases to evaluate for other potential contributors, because the cause may be multifactorial. Antidepressant treatment ([Chap. 466](#)) may be helpful for treatment of chronic fatigue when symptoms of depression are present and may be most effective in the context of a multimodal approach. However, antidepressants can also cause fatigue and should be discontinued if they are not clearly effective. Cognitive-behavioral therapy has also been demonstrated to be helpful in the context of chronic fatigue syndrome as well as cancer-associated fatigue. Graded exercise therapy in which physical exercise, most typically walking, is gradually increased with attention to target heart rates to avoid overexertion, was shown to modestly improve walking times and self-reported fatigue measures in patients in the United Kingdom with chronic fatigue syndrome in the large 2011 randomized controlled PACE trial. Psychostimulants such as amphetamines, modafinil, and armodafinil can help increase alertness and concentration and reduce excessive daytime sleepiness in certain clinical contexts, which may in turn help with symptoms of fatigue in a minority of patients, but they have generally proven to be unhelpful in randomized trials for treating fatigue in posttraumatic brain injury, Parkinson's disease, and MS.

Development of more effective therapy for fatigue is hampered by limited knowledge of the biologic basis of this symptom. Tentative data suggests that proinflammatory cytokines, such as interleukin 1 β and tumor necrosis factor α , might mediate fatigue in some patients; thus, cytokine antagonists represent one possible future approach.

PROGNOSIS

Acute fatigue significant enough to require medical evaluation is more likely to lead to an identifiable medical, neurologic, or psychiatric cause than unexplained chronic fatigue. Evaluation of unexplained chronic fatigue most commonly leads to diagnosis of a psychiatric condition or remains unexplained. Identification of a previously undiagnosed serious or life-threatening culprit etiology is rare on longitudinal follow-up in patients with unexplained chronic fatigue. Complete resolution of unexplained chronic fatigue is uncommon, at least over the short term, but multidisciplinary treatment approaches can lead to symptomatic improvements that can substantially improve quality of life.

Michael J. Aminoff

Normal motor function involves integrated muscle activity that is modulated by the activity of the cerebral cortex, basal ganglia, cerebellum, red nucleus, brainstem reticular formation, lateral vestibular nucleus, and spinal cord. Motor system dysfunction leads to weakness or paralysis, discussed in this chapter, or to ataxia (Chap. 450) or abnormal movements (Chap. 449). *Weakness* is a reduction in the power that can be exerted by one or more muscles. It must be distinguished from increased *fatigability* (i.e., the inability to sustain the performance of an activity that should be normal for a person of the same age, sex, and size), limitation in function due to pain or articular stiffness, or impaired motor activity because severe *proprioceptive sensory loss* prevents adequate feedback information about the direction and power of movements. It is also distinct from *bradykinesia* (in which increased time is required for full power to be exerted) and *apraxia*, a disorder of planning and initiating a skilled or learned movement unrelated to a significant motor or sensory deficit (Chap. 36).

Paralysis or the suffix “-plegia” indicates weakness so severe that a muscle cannot be contracted at all, whereas *paresis* refers to less severe weakness. The prefix “hemi-” refers to one-half of the body, “para-” to both legs, and “quadri-” to all four limbs.

The *distribution* of weakness helps to localize the underlying lesion. Weakness from involvement of upper motor neurons occurs particularly in the extensors and abductors of the upper limb and the flexors of the lower limb. Lower motor neuron weakness depends on whether involvement is at the level of the anterior horn cells, nerve root, limb plexus, or peripheral nerve—only muscles supplied by the affected structure are weak. Myopathic weakness is generally most marked in proximal muscles. Weakness from impaired neuromuscular transmission has no specific pattern of involvement.

Weakness often is accompanied by other neurologic abnormalities that help indicate the site of the responsible lesion (Table 30-1).

Tone is the resistance of a muscle to passive stretch. Increased tone may be of several types. *Spasticity* is the increase in tone associated with disease of upper motor neurons. It is velocity-dependent, has a sudden release after reaching a maximum (the “clasp-knife” phenomenon), and predominantly affects the antigravity muscles (i.e., upper-limb flexors and lower-limb extensors). *Rigidity* is hypertonia that is present throughout the range of motion (a “lead pipe” or “plastic” stiffness) and affects flexors and extensors equally; it sometimes has a cogwheel quality that is enhanced by voluntary movement of the contralateral limb (reinforcement). Rigidity occurs with certain extrapyramidal disorders, such as Parkinson’s disease. *Paratonia* (or *gegenhalten*) is increased tone that varies irregularly in a manner seemingly related to the degree of relaxation, is present throughout the range of motion, and affects flexors and extensors equally; it usually results from disease of the frontal lobes. Weakness with *decreased tone* (*flaccidity*) or normal tone occurs with disorders of *motor units*. A motor unit consists of a single lower motor neuron and all the muscle fibers that it innervates.

Muscle bulk generally is not affected by upper motor neuron lesions, although mild disuse atrophy eventually may occur. By contrast, atrophy is often conspicuous when a lower motor neuron lesion is responsible for weakness and also may occur with advanced muscle disease.

Muscle stretch (tendon) reflexes are usually increased with upper motor neuron lesions, but may be decreased or absent for a variable period immediately after onset of an acute lesion. Hyperreflexia is usually—but not invariably—accompanied by loss of *cutaneous reflexes* (such as superficial abdominals; Chap. 437) and, in particular, by an extensor plantar (Babinski) response. The muscle stretch reflexes are depressed with lower motor neuron lesions directly involving specific reflex arcs. They generally are preserved in patients with myopathic weakness except in advanced stages, when they sometimes are attenuated. In disorders of the neuromuscular junction, reflex responses may be affected by preceding voluntary activity of affected muscles; such activity may lead to enhancement of initially depressed reflexes in Lambert-Eaton myasthenic syndrome and, conversely, to depression of initially normal reflexes in myasthenia gravis (Chap. 461).

The distinction of *neuropathic* (lower motor neuron) from *myopathic* weakness is sometimes difficult clinically, although distal weakness is likely to be neuropathic, and symmetric proximal weakness myopathic. *Fasciculations* (visible or palpable twitch within a muscle due to the spontaneous discharge of a motor unit) and early atrophy indicate that weakness is neuropathic.

PATHOGENESIS

Upper Motor Neuron Weakness Lesions of the upper motor neurons or their descending axons to the spinal cord (Fig. 30-1) produce weakness through decreased activation of lower motor neurons. In general, distal muscle groups are affected more severely than proximal ones, and axial movements are spared unless the lesion is severe and bilateral. Spasticity is typical but may not be present acutely. Rapid repetitive movements are slowed and coarse, but normal rhythmicity is maintained. With corticobulbar involvement, weakness occurs in the lower face and tongue; extraocular, upper facial, pharyngeal, and jaw muscles are typically spared. Bilateral corticobulbar lesions produce a *pseudobulbar palsy*: dysarthria, dysphagia, dysphonia, and emotional lability accompany bilateral facial weakness and a brisk jaw jerk. Electromyogram (EMG) (Chap. 442e) shows that with weakness of the upper motor neuron type, motor units have a diminished maximal discharge frequency.

Lower Motor Neuron Weakness This pattern results from disorders of lower motor neurons in the brainstem motor nuclei and the anterior horn of the spinal cord or from dysfunction of the axons of these neurons as they pass to skeletal muscle (Fig. 30-2). Weakness is due to a decrease in the number of muscle fibers that can be activated through a loss of a motor neurons or disruption of their connections to muscle. Loss of γ motor neurons does not cause weakness but decreases tension on the muscle spindles, which decreases muscle tone and attenuates the stretch reflexes. An absent stretch reflex suggests involvement of spindle afferent fibers.

When a motor unit becomes diseased, especially in anterior horn cell diseases, it may discharge spontaneously, producing *fasciculations*. When a motor neurons or their axons degenerate, the denervated muscle fibers also may discharge spontaneously. These single muscle

TABLE 30-1 SIGNS THAT DISTINGUISH THE ORIGIN OF WEAKNESS

Sign	Upper Motor Neuron	Lower Motor Neuron	Myopathic	Psychogenic
Atrophy	None	Severe	Mild	None
Fasciculations	None	Common	None	None
Tone	Spastic	Decreased	Normal/decreased	Variable/paratonia
Distribution of weakness	Pyramidal/regional	Distal/segmental	Proximal	Variable/inconsistent with daily activities
Muscle stretch reflexes	Hyperactive	Hypoactive/absent	Normal/hypoactive	Normal
Babinski sign	Present	Absent	Absent	Absent

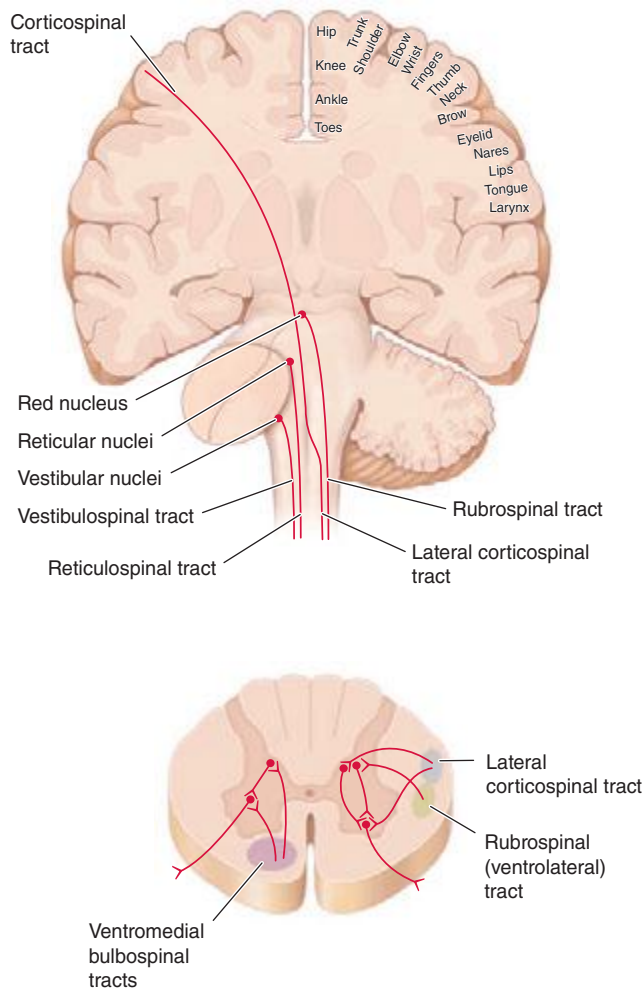


FIGURE 30-1 The corticospinal and bulbospinal upper motor neuron pathways. Upper motor neurons have their cell bodies in layer V of the primary motor cortex (the precentral gyrus, or Brodmann's area 4) and in the premotor and supplemental motor cortex (area 6). The upper motor neurons in the primary motor cortex are somatotopically organized (right side of figure).

Axons of the upper motor neurons descend through the subcortical white matter and the posterior limb of the internal capsule. Axons of the *pyramidal* or *corticospinal* system descend through the brainstem in the cerebral peduncle of the midbrain, the basis pontis, and the medullary pyramids. At the cervicomedullary junction, most corticospinal axons decussate into the contralateral corticospinal tract of the lateral spinal cord, but 10–30% remain ipsilateral in the anterior spinal cord. Corticospinal neurons synapse on premotor interneurons, but some—especially in the cervical enlargement and those connecting with motor neurons to distal limb muscles—make direct monosynaptic connections with lower motor neurons. They innervate most densely the lower motor neurons of hand muscles and are involved in the execution of learned, fine movements. Corticobulbar neurons are similar to corticospinal neurons but innervate brainstem motor nuclei.

Bulbospinal upper motor neurons influence strength and tone but are not part of the pyramidal system. The descending *ventromedial bulbospinal pathways* originate in the tectum of the midbrain (tectospinal pathway), the vestibular nuclei (vestibulospinal pathway), and the reticular formation (reticulospinal pathway). These pathways influence axial and proximal muscles and are involved in the maintenance of posture and integrated movements of the limbs and trunk. The descending *ventrolateral bulbospinal pathways*, which originate predominantly in the red nucleus (rubrospinal pathway), facilitate distal limb muscles. The bulbospinal system sometimes is referred to as the *extrapyramidal upper motor neuron system*. In all figures, nerve cell bodies and axon terminals are shown, respectively, as closed circles and forks.

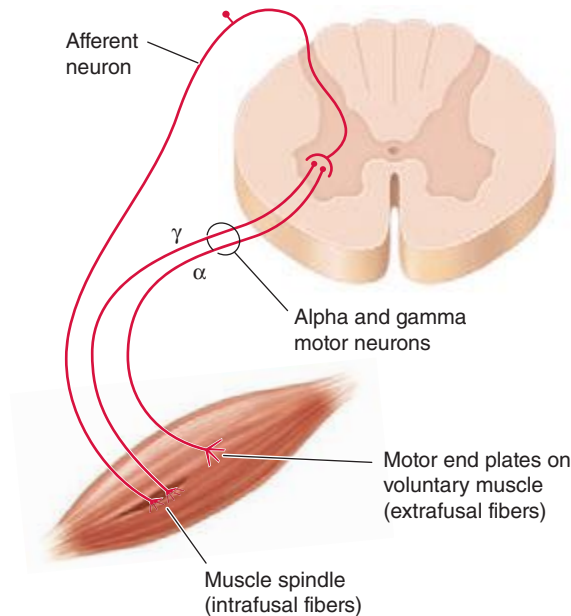


FIGURE 30-2 Lower motor neurons are divided into α and γ types.

The larger α motor neurons are more numerous and innervate the extrafusal muscle fibers of the motor unit. Loss of a motor neurons or disruption of their axons produces lower motor neuron weakness. The smaller, less numerous γ motor neurons innervate the intrafusal muscle fibers of the muscle spindle and contribute to normal tone and stretch reflexes. The α motor neuron receives direct excitatory input from corticomotoneurons and primary muscle spindle afferents. The α and γ motor neurons also receive excitatory input from other descending upper motor neuron pathways, segmental sensory inputs, and interneurons. The α motor neurons receive direct inhibition from Renshaw cell interneurons, and other interneurons indirectly inhibit the α and γ motor neurons.

A muscle stretch (tendon) reflex requires the function of all the illustrated structures. A tap on a tendon stretches muscle spindles (which are tonically activated by γ motor neurons) and activates the primary spindle afferent neurons. These neurons stimulate the α motor neurons in the spinal cord, producing a brief muscle contraction, which is the familiar tendon reflex.

fiber discharges, or *fibrillation potentials*, cannot be seen but can be recorded with EMG. Weakness leads to delayed or reduced recruitment of motor units, with fewer than normal activated at a particular discharge frequency.

Neuromuscular Junction Weakness Disorders of the neuromuscular junctions produce weakness of variable degree and distribution. The number of muscle fibers that are activated varies over time, depending on the state of rest of the neuromuscular junctions. Strength is influenced by preceding activity of the affected muscle. In myasthenia gravis, for example, sustained or repeated contractions of affected muscle decline in strength despite continuing effort (**Chap. 461**). Thus, fatigable weakness is suggestive of disorders of the neuromuscular junction, which cause functional loss of muscle fibers due to failure of their activation.

Myopathic Weakness Myopathic weakness is produced by a decrease in the number or contractile force of muscle fibers activated within motor units. With muscular dystrophies, inflammatory myopathies, or myopathies with muscle fiber necrosis, the number of muscle fibers is reduced within many motor units. On EMG, the size of each motor unit action potential is decreased, and motor units must be recruited more rapidly than normal to produce the desired power. Some myopathies produce weakness through loss of contractile force of muscle fibers or through relatively selective involvement of type II (fast)

fibers. These myopathies may not affect the size of individual motor unit action potentials and are detected by a discrepancy between the electrical activity and force of a muscle.

Psychogenic Weakness Weakness may occur without a recognizable organic basis. It tends to be variable, inconsistent, and with a pattern of distribution that cannot be explained on a neuroanatomic basis. On formal testing, antagonists may contract when the patient is supposedly activating the agonist muscle. The severity of weakness is out of keeping with the patient's daily activities.

Hemiparesis Hemiparesis results from an upper motor neuron lesion above the midcervical spinal cord; most such lesions are above the foramen magnum. The presence of other neurologic deficits helps localize the lesion. Thus, language disorders, for example, point to a cortical lesion. Homonymous visual field defects reflect either a cortical or a subcortical hemispheric lesion. A "pure motor" hemiparesis of the face, arm, and leg often is due to a small, discrete lesion in the posterior limb of the internal capsule, cerebral peduncle, or upper pons. Some brainstem lesions produce "crossed paralyzes," consisting of ipsilateral cranial nerve signs and contralateral hemiparesis (Chap. 446). The absence of cranial nerve signs or facial weakness suggests that a hemiparesis is due to a lesion in the high cervical spinal cord, especially if associated with the Brown-Séquard syndrome (Chap. 456).

Acute or episodic hemiparesis usually results from focal structural lesions, particularly rapidly expanding lesions, or an inflammatory process. *Subacute hemiparesis* that evolves over days or weeks may relate to subdural hematoma, infectious or inflammatory disorders (e.g., cerebral abscess, fungal granuloma or meningitis, parasitic infection, multiple sclerosis, sarcoidosis), or primary and metastatic neoplasms. AIDS may present with subacute hemiparesis due to toxoplasmosis or primary central nervous system (CNS) lymphoma. *Chronic hemiparesis* that evolves over months usually is due to a neoplasm or vascular malformation, a chronic subdural hematoma, or a degenerative disease.

Investigation of hemiparesis (Fig. 30-3) of acute origin starts with a computed tomography (CT) scan of the brain and laboratory studies. If the CT is normal, or in subacute or chronic cases of hemiparesis, magnetic resonance imaging (MRI) of the brain and/or cervical spine (including the foramen magnum) is performed, depending on the clinical accompaniments.

Paraparesis *Acute paraparesis* is caused most commonly by an intraspinal lesion, but its spinal origin may not be recognized initially if the legs are flaccid and areflexic. Usually, however, there is sensory loss in the legs with an upper level on the trunk, a dissociated sensory loss suggestive of a central cord syndrome (Chap. 456), or hyperreflexia in the legs with normal reflexes in the arms. Imaging the spinal cord (Fig. 30-3) may reveal compressive lesions, infarction (proprioception usually is spared), arteriovenous fistulas or other vascular anomalies, or transverse myelitis (Chap. 456).

Diseases of the cerebral hemispheres that produce acute paraparesis include anterior cerebral artery ischemia (shoulder shrug also is affected), superior sagittal sinus or cortical venous thrombosis, and acute hydrocephalus.

Paraparesis may result from a cauda equina syndrome, for example, after trauma to the low back, a midline disk herniation, or an intraspinal tumor; although the sphincters are often affected, hip flexion often is spared, as is sensation over the anterolateral thighs. Rarely, paraparesis is caused by a rapidly evolving anterior horn cell disease (such as poliovirus or West Nile virus infection), peripheral neuropathy (such as Guillain-Barré syndrome; Chap. 460), or myopathy (Chap. 462e).

Subacute or chronic spastic paraparesis is caused by upper motor neuron disease. When associated with lower-limb sensory loss and sphincter involvement, a chronic spinal cord disorder should be considered (Chap. 456). If hemispheric signs are present, a parasagittal meningioma or chronic hydrocephalus is likely. The absence of spasticity in a long-standing paraparesis suggests a lower motor neuron or myopathic etiology.

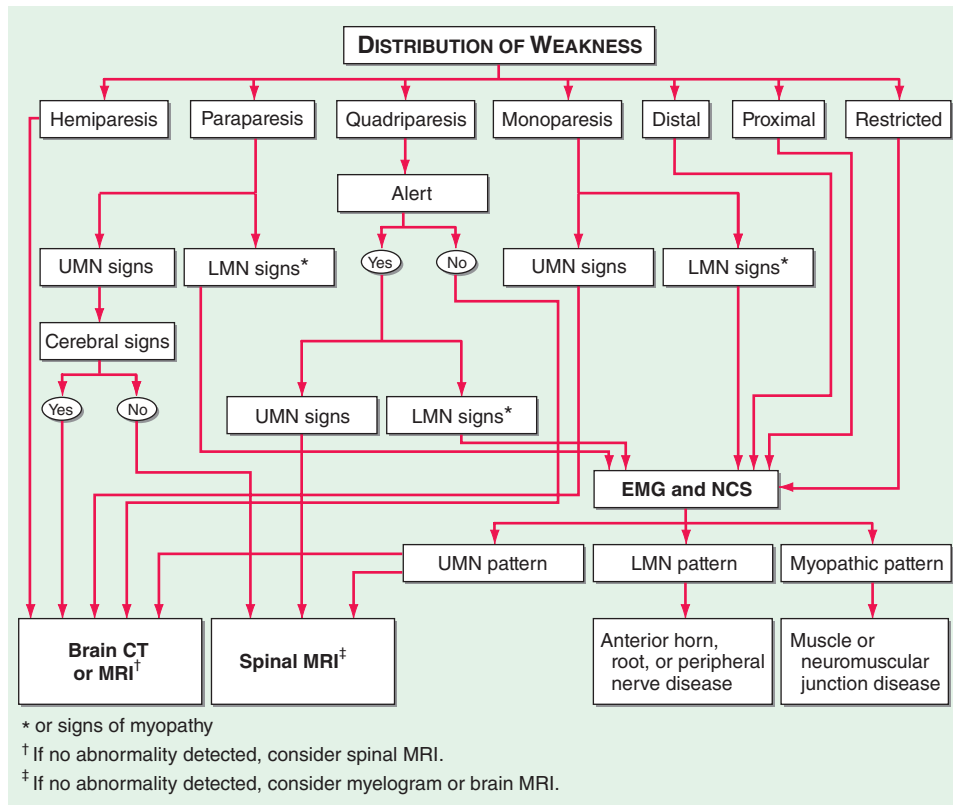


FIGURE 30-3 An algorithm for the initial workup of a patient with weakness. CT, computed tomography; EMG, electromyography; LMN, lower motor neuron; MRI, magnetic resonance imaging; NCS, nerve conduction studies; UMN, upper motor neuron.

Investigations typically begin with spinal MRI, but when upper motor neuron signs are associated with drowsiness, confusion, seizures, or other hemispheric signs, brain MRI should also be performed, sometimes as the initial investigation. Electrophysiologic studies are diagnostically helpful when clinical findings suggest an underlying neuromuscular disorder.

Quadripareisis or Generalized Weakness Generalized weakness may be due to disorders of the CNS or the motor unit. Although the terms often are used interchangeably, *quadripareisis* is commonly used when an upper motor neuron cause is suspected, and *generalized weakness* is used when a disease of the motor units is likely. Weakness from CNS disorders usually is associated with changes in consciousness or cognition and accompanied by spasticity, hyperreflexia, and sensory disturbances. Most neuromuscular causes of generalized weakness are associated with normal mental function, hypotonia, and hypoactive muscle stretch reflexes. The major causes of intermittent weakness are listed in **Table 30-2**. A patient with generalized fatigability without objective weakness may have the chronic fatigue syndrome (**Chap. 464e**).

ACUTE QUADRIPAREISIS Quadripareisis with onset over minutes may result from disorders of upper motor neurons (such as from anoxia, hypotension, brainstem or cervical cord ischemia, trauma, and systemic metabolic abnormalities) or muscle (electrolyte disturbances, certain inborn errors of muscle energy metabolism, toxins, and periodic paralyses). Onset over hours to weeks may, in addition to these disorders, be due to lower motor neuron disorders such as Guillain-Barré syndrome (**Chap. 460**).

In obtunded patients, evaluation begins with a CT scan of the brain. If upper motor neuron signs are present but the patient is alert, the initial test is usually an MRI of the cervical cord. If weakness is lower motor neuron, myopathic, or uncertain in origin, the clinical approach begins with blood studies to determine the level of muscle enzymes and electrolytes and with EMG and nerve conduction studies.

SUBACUTE OR CHRONIC QUADRIPAREISIS Quadripareisis due to upper motor neuron disease may develop over weeks to years from chronic myelopathies, multiple sclerosis, brain or spinal tumors, chronic subdural hematomas, and various metabolic, toxic, and infectious disorders. It may also result from lower motor neuron disease, a chronic neuropathy (in which weakness is often most profound distally), or myopathic weakness (typically proximal).

When *quadripareisis* develops acutely in obtunded patients, evaluation begins with a CT scan of the brain. If upper motor neuron signs have developed acutely but the patient is alert, the initial test is usually

an MRI of the cervical cord. When onset has been gradual, disorders of the cerebral hemispheres, brainstem, and cervical spinal cord can usually be distinguished clinically, and imaging is directed first at the clinically suspected site of pathology. If weakness is lower motor neuron, myopathic, or uncertain in origin, laboratory studies to determine the levels of muscle enzymes and electrolytes, and EMG and nerve conduction studies help to localize the pathologic process.

Monoparesis Monoparesis usually is due to lower motor neuron disease, with or without associated sensory involvement. Upper motor neuron weakness occasionally presents as a monoparesis of distal and nonantigravity muscles. Myopathic weakness rarely is limited to one limb.

ACUTE MONOPARESIS If weakness is predominantly distal and of upper motor neuron type and is not associated with sensory impairment or pain, focal cortical ischemia is likely (**Chap. 446**); diagnostic possibilities are similar to those for acute hemiparesis. Sensory loss and pain usually accompany acute lower motor neuron weakness; the weakness commonly localizes to a single nerve root or peripheral nerve, but occasionally reflects plexus involvement. If lower motor neuron weakness is likely, evaluation begins with EMG and nerve conduction studies.

SUBACUTE OR CHRONIC MONOPARESIS Weakness and atrophy that develop over weeks or months are usually of lower motor neuron origin. When associated with sensory symptoms, a peripheral cause (nerve, root, or plexus) is likely; otherwise, anterior horn cell disease should be considered. In either case, an electrodiagnostic study is indicated. If weakness is of the upper motor neuron type, a discrete cortical (precentral gyrus) or cord lesion may be responsible, and appropriate imaging is performed.

Distal Weakness Involvement of two or more limbs distally suggests lower motor neuron or peripheral nerve disease. Acute distal lower-limb weakness results occasionally from an acute toxic polyneuropathy or cauda equina syndrome. Distal symmetric weakness usually develops over weeks, months, or years and, when associated with numbness, is due to peripheral neuropathy (**Chap. 459**). Anterior horn cell disease may begin distally but is typically asymmetric and without accompanying numbness (**Chap. 452**). Rarely, myopathies present with distal weakness (**Chap. 462e**). Electrodiagnostic studies help localize the disorder (**Fig. 30-3**).

Proximal Weakness Myopathy often produces symmetric weakness of the pelvic or shoulder girdle muscles (**Chap. 462e**). Diseases of the neuromuscular junction, such as myasthenia gravis (**Chap. 461**), may present with symmetric proximal weakness often associated with ptosis, diplopia, or bulbar weakness and fluctuating in severity during the day. In anterior horn cell disease, proximal weakness is usually asymmetric, but it may be symmetric if familial. Numbness does not occur with any of these diseases. The evaluation usually begins with determination of the serum creatine kinase level and electrophysiologic studies.

Weakness in a Restricted Distribution Weakness may not fit any of these patterns, being limited, for example, to the extraocular, hemifacial, bulbar, or respiratory muscles. If it is unilateral, restricted weakness usually is due to lower motor neuron or peripheral nerve disease, such as in a facial palsy. Weakness of part of a limb is commonly due to a peripheral nerve lesion such as an entrapment neuropathy. Relatively symmetric weakness of extraocular or bulbar muscles frequently is due to a myopathy (**Chap. 462e**) or neuromuscular junction disorder (**Chap. 461**). Bilateral facial palsy with areflexia suggests Guillain-Barré syndrome (**Chap. 460**). Worsening of relatively symmetric weakness with fatigue is characteristic of neuromuscular junction disorders. Asymmetric bulbar weakness usually is due to motor neuron disease. Weakness limited to respiratory muscles is uncommon and usually is due to motor neuron disease, myasthenia gravis, or polymyositis/dermatomyositis (**Chap. 388**).

TABLE 30-2 CAUSES OF EPISODIC GENERALIZED WEAKNESS

1. Electrolyte disturbances, e.g., hypokalemia, hyperkalemia, hypercalcemia, hyponatremia, hyponatremia, hypophosphatemia, hypermagnesemia
2. Muscle disorders
 - a. Channelopathies (periodic paralyses)
 - b. Metabolic defects of muscle (impaired carbohydrate or fatty acid utilization; abnormal mitochondrial function)
3. Neuromuscular junction disorders
 - a. Myasthenia gravis
 - b. Lambert-Eaton myasthenic syndrome
4. Central nervous system disorders
 - a. Transient ischemic attacks of the brainstem
 - b. Transient global cerebral ischemia
 - c. Multiple sclerosis
5. Lack of voluntary effort
 - a. Anxiety
 - b. Pain or discomfort
 - c. Somatization disorder

31 Numbness, Tingling, and Sensory Loss

Michael J. Aminoff

Normal somatic sensation reflects a continuous monitoring process, little of which reaches consciousness under ordinary conditions. By contrast, disordered sensation, particularly when experienced as painful, is alarming and dominates the patient's attention. Physicians should be able to recognize abnormal sensations by how they are described, know their type and likely site of origin, and understand their implications. **Pain is considered separately in Chap. 18.**

POSITIVE AND NEGATIVE SYMPTOMS

Abnormal sensory symptoms can be divided into two categories: positive and negative. The prototypical positive symptom is tingling (pins and needles); other positive sensory phenomena include itch and altered sensations that are described as pricking, bandlike, lightning-like shooting feelings (lancinations), aching, knifelike, twisting, drawing, pulling, tightening, burning, searing, electrical, or raw feelings. Such symptoms are often painful.

Positive phenomena usually result from trains of impulses generated at sites of lowered threshold or heightened excitability along a peripheral or central sensory pathway. The nature and severity of the abnormal sensation depend on the number, rate, timing, and distribution of ectopic impulses and the type and function of nervous tissue in which they arise. Because positive phenomena represent excessive activity in sensory pathways, they are not necessarily associated with a sensory deficit (loss) on examination.

Negative phenomena represent loss of sensory function and are characterized by diminished or absent feeling that often is experienced as numbness and by abnormal findings on sensory examination. In disorders affecting peripheral sensation, at least one-half the afferent axons innervating a particular site are probably lost or functionless before a sensory deficit can be demonstrated by clinical examination. If the rate of loss is slow, however, lack of cutaneous feeling may be unnoticed by the patient and difficult to demonstrate on examination, even though few sensory fibers are functioning; if it is rapid, both positive and negative phenomena are usually conspicuous. Subclinical degrees of sensory dysfunction may be revealed by sensory nerve conduction studies or somatosensory evoked potentials (**Chap. 442e**).

Whereas sensory symptoms may be either positive or negative, sensory signs on examination are always a measure of negative phenomena.

TERMINOLOGY

Paresthesias and dysesthesias are general terms used to denote positive sensory symptoms. The term *paresthesias* typically refers to tingling or pins-and-needles sensations but may include a wide variety of other abnormal sensations, except pain; it sometimes implies that the abnormal sensations are perceived spontaneously. The more general term *dysesthesias* denotes all types of abnormal sensations, including painful ones, regardless of whether a stimulus is evident.

Another set of terms refers to sensory abnormalities found on examination. *Hypesthesia* or *hypoesthesia* refers to a reduction of cutaneous sensation to a specific type of testing such as pressure, light touch, and warm or cold stimuli; *anesthesia*, to a complete absence of skin sensation to the same stimuli plus pinprick; and *hypalgesia* or *analgesia*, to reduced or absent pain perception (nociception). *Hyperesthesia* means pain or increased sensitivity in response to touch. Similarly, *allodynia* describes the situation in which a nonpainful stimulus, once perceived, is experienced as painful, even excruciating. An example is elicitation of a painful sensation by application of a vibrating tuning fork. *Hyperalgesia* denotes severe pain in response to a mildly noxious stimulus, and *hyperpathia*, a broad term, encompasses all the phenomena described by hyperesthesia, allodynia, and hyperalgesia. With hyperpathia, the threshold for a sensory stimulus is increased and perception is delayed, but once felt, it is unduly painful.

Disorders of deep sensation arising from muscle spindles, tendons, and joints affect proprioception (position sense). Manifestations include imbalance (particularly with eyes closed or in the dark), clumsiness of precision movements, and unsteadiness of gait, which are referred to collectively as *sensory ataxia*. Other findings on examination usually, but not invariably, include reduced or absent joint position and vibratory sensibility and absent deep tendon reflexes in the affected limbs. The Romberg sign is positive, which means that the patient sways markedly or topples when asked to stand with feet close together and eyes closed. In severe states of deafferentation involving deep sensation, the patient cannot walk or stand unaided or even sit unsupported. Continuous involuntary movements (*pseudoathetosis*) of the outstretched hands and fingers occur, particularly with eyes closed.

ANATOMY OF SENSATION

Cutaneous receptors are classified by the type of stimulus that optimally excites them. They consist of naked nerve endings (nociceptors, which respond to tissue-damaging stimuli, and thermoreceptors, which respond to noninjurious thermal stimuli) and encapsulated terminals (several types of mechanoreceptor, activated by physical deformation of the skin). Each type of receptor has its own set of sensitivities to specific stimuli, size and distinctness of receptive fields, and adaptational qualities.

Afferent fibers in peripheral nerve trunks traverse the dorsal roots and enter the dorsal horn of the spinal cord (**Fig. 31-1**). From there, the polysynaptic projections of the smaller fibers (unmyelinated and small myelinated), which subserve mainly nociception, itch, temperature sensibility, and touch, cross and ascend in the opposite anterior and lateral columns of the spinal cord, through the brainstem, to the ventral posterolateral (VPL) nucleus of the thalamus and ultimately project to the postcentral gyrus of the parietal cortex (**Chap. 18**). This is the *spinothalamic pathway* or *anterolateral system*. The larger fibers, which subserve tactile and position sense and kinesthesia, project rostrally in the posterior and posterolateral columns on the same side of the spinal cord and make their first synapse in the gracile or cuneate nucleus of the lower medulla. Axons of second-order neurons decussate and ascend in the medial lemniscus located medially in the medulla and in the tegmentum of the pons and midbrain and synapse in the VPL nucleus; third-order neurons project to parietal cortex as well as to other cortical areas. This large-fiber system is referred to as the *posterior column–medial lemniscal pathway* (lemniscal, for short). Although the fiber types and functions that make up the spinothalamic and lemniscal systems are relatively well known, many other fibers, particularly those associated with touch, pressure, and position sense, ascend in a diffusely distributed pattern both ipsilaterally and contralaterally in the anterolateral quadrants of the spinal cord. This explains why a complete lesion of the posterior columns of the spinal cord may be associated with little sensory deficit on examination.

Nerve conduction studies and nerve biopsy are important means of investigating the peripheral nervous system, but they do not evaluate the function or structure of cutaneous receptors and free nerve endings or of unmyelinated or thinly myelinated nerve fibers in the nerve trunks. Skin biopsy can be used to evaluate these structures in the dermis and epidermis.

CLINICAL EXAMINATION OF SENSATION

The main components of the sensory examination are tests of primary sensation (pain, touch, vibration, joint position, and thermal sensation) (**Table 31-1**). The examiner must depend on patient responses, and this complicates interpretation. Further, examination may be limited in some patients. In a stuporous patient, for example, sensory examination is reduced to observing the briskness of withdrawal in response to a pinch or another noxious stimulus. Comparison of responses on the two sides of the body is essential. In an alert but uncooperative patient, it may not be possible to examine cutaneous sensation, but some idea of proprioceptive function may be gained by noting the patient's best performance of movements requiring balance and precision.

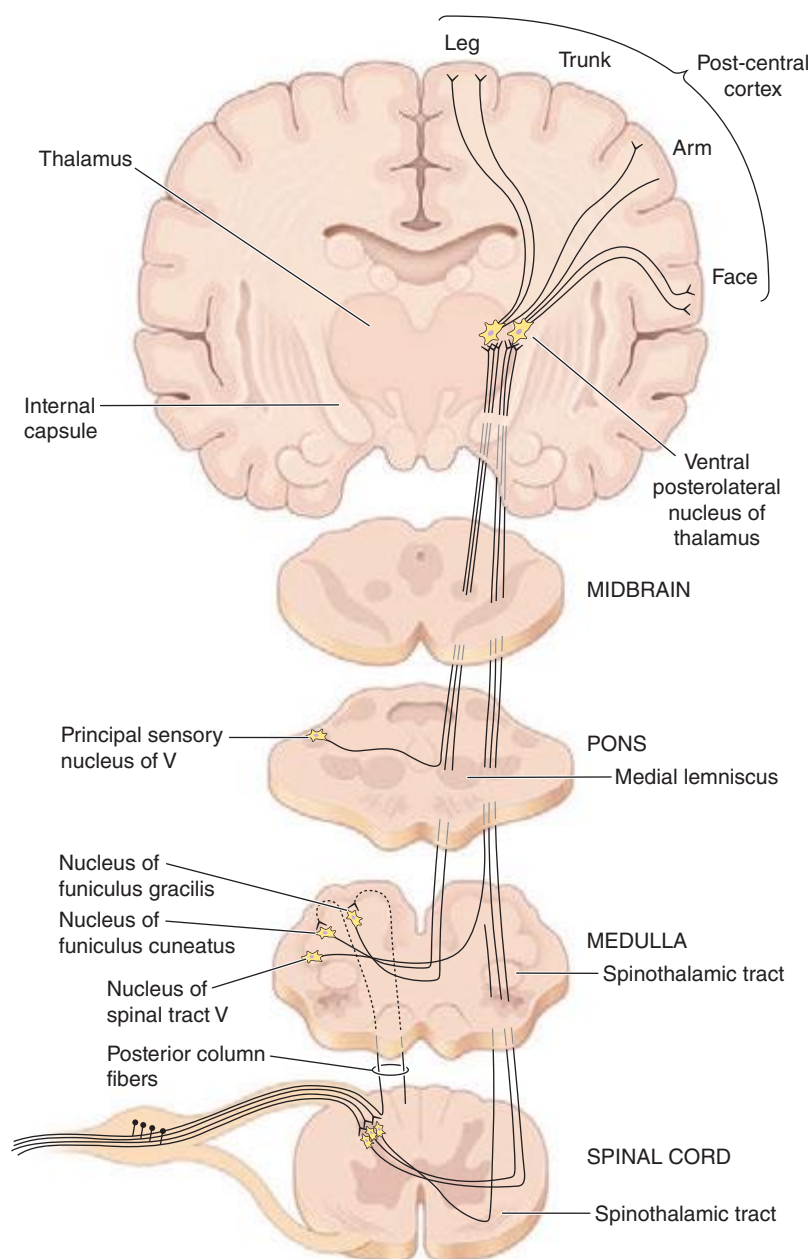


FIGURE 31-1 The main somatosensory pathways. The spinothalamic tract (pain, thermal sense) and the posterior column–lemniscal system (touch, pressure, joint position) are shown. Offshoots from the ascending anterolateral fasciculus (spinothalamic tract) to nuclei in the medulla, pons, and mesencephalon and nuclear terminations of the tract are indicated. (From AH Ropper, MA Samuels: *Adams and Victor's Principles of Neurology*, 9th ed. New York, McGraw-Hill, 2009.)

In patients with sensory complaints, testing should begin in the center of the affected region and proceed radially until sensation is perceived as normal. The distribution of any abnormality is defined and compared to root and peripheral nerve territories (Figs. 31-2 and 31-3). Some patients present with sensory symptoms that do not fit an anatomic localization and are accompanied by either no abnormalities or gross inconsistencies on examination. The examiner should consider whether the sensory symptoms are a disguised request for help with psychologic or situational problems. Sensory examination of a patient who has no neurologic complaints can be brief and consist of pinprick, touch, and vibration testing in the hands and feet plus evaluation of stance and gait, including the Romberg maneuver (Chap. 438). Evaluation of stance and gait also tests the integrity of motor and cerebellar systems.

Primary Sensation The sense of pain usually is tested with a clean pin, which is then discarded. The patient is asked to close the eyes and focus on the pricking or unpleasant quality of the stimulus, not just the pressure or touch sensation elicited. Areas of hypalgesia should be mapped by proceeding radially from the most hypalgesic site. Temperature sensation to both hot and cold is best tested with small containers filled with water of the desired temperature. An alternative way to test cold sensation is to touch a metal object, such as a tuning fork at room temperature, to the skin. For testing warm temperatures, the tuning fork or another metal object may be held under warm water of the desired temperature and then used. The appreciation of both cold and warmth should be tested because different receptors respond to each. Touch usually is tested with a wisp of cotton or a fine camel hair brush, minimizing pressure on the skin. In general, it is better to avoid testing touch on hairy skin because of the profusion of the sensory endings that surround each hair follicle. The patient is tested with the eyes closed and should indicate as soon as the stimulus is perceived, indicating its location.

Joint position testing is a measure of proprioception. With the patient's eyes closed, joint position is tested in the distal interphalangeal joint of the great toe and fingers. The digit is held by its sides, distal to the joint being tested, and moved passively while more proximal joints are stabilized—the patient indicates the change in position or direction of movement. If errors are made, more proximal joints are tested. A test of proximal joint position sense, primarily at the shoulder, is performed by asking the patient to bring the two index fingers together with arms extended and eyes closed. Normal individuals can do this accurately, with errors of 1 cm or less.

The sense of vibration is tested with an oscillating tuning fork that vibrates at 128 Hz. Vibration is tested over bony points, beginning distally; in the feet, it is tested over the dorsal surface of the distal phalanx of the big toes and at the malleoli of the ankles, and in the hands, it is tested dorsally at the distal phalanx of the fingers. If abnormalities are found, more proximal sites should be examined. Vibratory thresholds at the same site in the patient and the examiner may be compared for control purposes.

Quantitative Sensory Testing Effective sensory testing devices are commercially available. Quantitative sensory testing is particularly useful for serial evaluation of cutaneous sensation in clinical trials. Threshold testing for touch and vibratory and thermal sensation is the most widely used application.

Cortical Sensation The most commonly used tests of cortical function are two-point discrimination, touch localization, and bilateral simultaneous stimulation and tests for graphesthesia and stereognosis. Abnormalities of these sensory tests, in the presence of normal primary sensation in an alert cooperative patient, signify a lesion of the parietal cortex or thalamocortical projections. If primary sensation is altered, these cortical discriminative functions usually will be abnormal also. Comparisons should always be made between analogous sites on the two sides of the body because the deficit with a specific parietal lesion is likely to be unilateral.

Two-point discrimination is tested with special calipers, the points of which may be set from 2 mm to several centimeters apart and then applied simultaneously to the test site. On the fingertips, a normal individual can distinguish about a 3-mm separation of points.

Touch localization is performed by light pressure for an instant with the examiner's fingertip or a wisp of cotton wool; the patient,

TABLE 31-1 TESTING PRIMARY SENSATION

Sense	Test Device	Endings Activated	Fiber Size Mediating	Central Pathway
Pain	Pinprick	Cutaneous nociceptors	Small	SpTh, also D
Temperature, heat	Warm metal object	Cutaneous thermoreceptors for hot	Small	SpTh
Temperature, cold	Cold metal object	Cutaneous thermoreceptors for cold	Small	SpTh
Touch	Cotton wisp, fine brush	Cutaneous mechanoreceptors, also naked endings	Large and small	Lem, also D and SpTh
Vibration	Tuning fork, 128 Hz	Mechanoreceptors, especially pacinian corpuscles	Large	Lem, also D
Joint position	Passive movement of specific joints	Joint capsule and tendon endings, muscle spindles	Large	Lem, also D

Abbreviations: D, diffuse ascending projections in ipsilateral and contralateral anterolateral columns; Lem, posterior column and lemniscal projection, ipsilateral; SpTh, spinothalamic projection, contralateral.

whose eyes are closed, is required to identify the site of touch. *Bilateral simultaneous stimulation* at analogous sites (e.g., the dorsum of both hands) can be carried out to determine whether the perception of touch is extinguished consistently on one side (*extinction* or *neglect*). *Graphesthesia* refers to the capacity to recognize, with eyes closed, letters or numbers drawn by the examiner's fingertip on the palm of the hand. Once again, interside comparison is of prime importance. Inability to recognize numbers or letters is termed *agraphesthesia*.

Stereognosis refers to the ability to identify common objects by palpation, recognizing their shape, texture, and size. Common standard objects such as keys, paper clips, and coins are best used. Patients with normal stereognosis should be able to distinguish a dime from a penny and a nickel from a quarter without looking. Patients should feel the object with only one hand at a time. If they are unable to identify it in one hand, it

should be placed in the other for comparison. Individuals who are unable to identify common objects and coins in one hand but can do so in the other are said to have *astereognosis* of the abnormal hand.

LOCALIZATION OF SENSORY ABNORMALITIES

Sensory symptoms and signs can result from lesions at many different levels of the nervous system from the parietal cortex to the peripheral sensory receptor. Noting their distribution and nature is the most important way to localize their source. Their extent, configuration, symmetry, quality, and severity are the key observations.

Dysesthesias without sensory findings by examination may be difficult to interpret. To illustrate, tingling dysesthesias in an acral distribution (hands and feet) can be systemic in origin, e.g., secondary to hyperventilation, or induced by a medication such as acetazolamide.

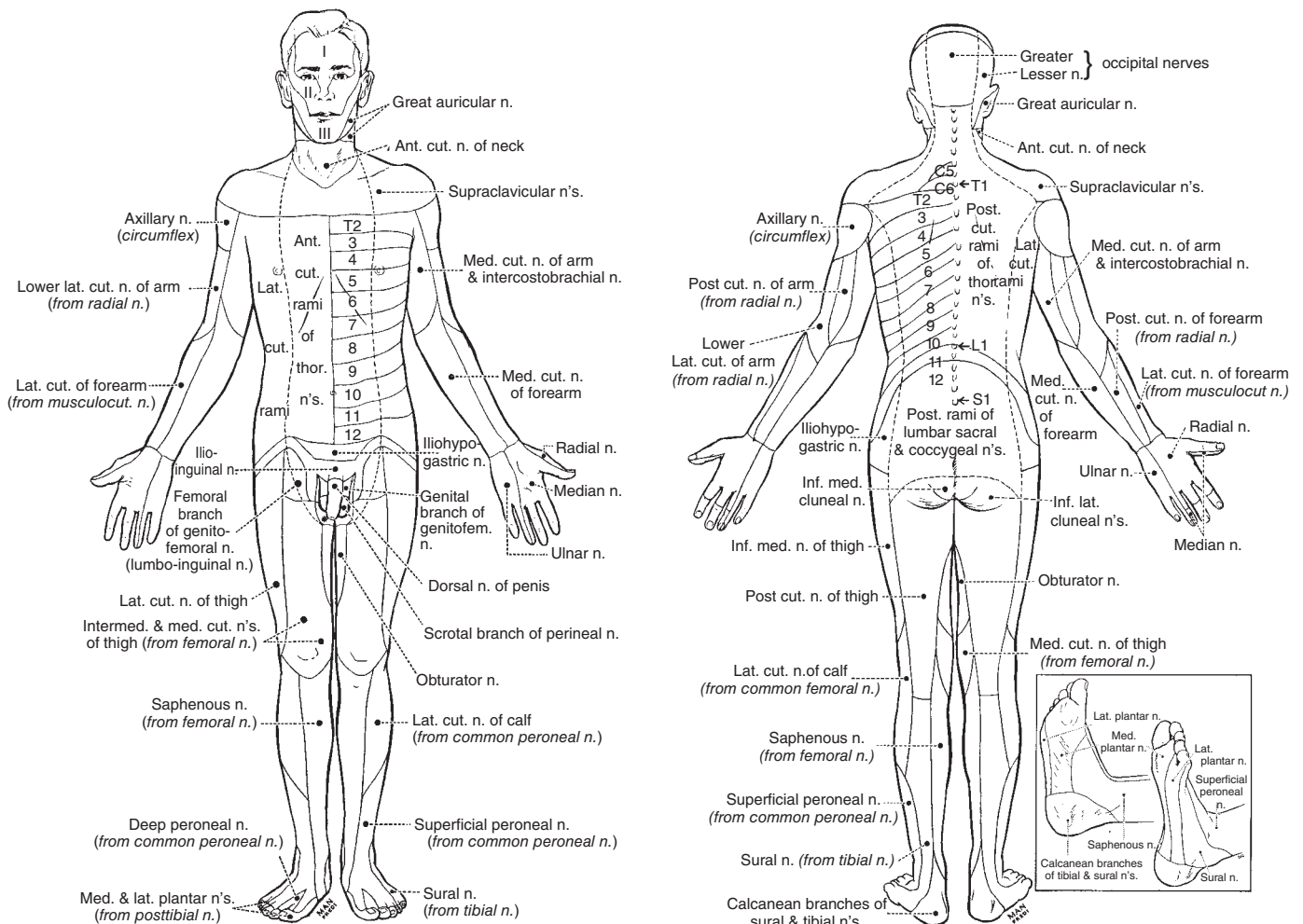


FIGURE 31-2 The cutaneous fields of peripheral nerves. (Reproduced by permission from W Haymaker, B Woodhall: *Peripheral Nerve Injuries*, 2nd ed. Philadelphia, Saunders, 1953.)

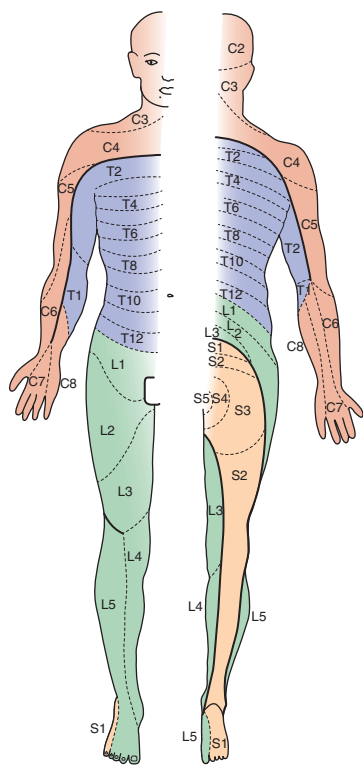


FIGURE 31-3 Distribution of the sensory spinal roots on the surface of the body (dermatomes). (From D Sinclair: *Mechanisms of Cutaneous Sensation*. Oxford, UK, Oxford University Press, 1981; with permission from Dr. David Sinclair.)

Distal dysesthesias can also be an early event in an evolving polyneuropathy or may herald a myelopathy, such as from vitamin B₁₂ deficiency. Sometimes distal dysesthesias have no definable basis. In contrast, dysesthesias that correspond in distribution to that of a particular peripheral nerve structure denote a lesion at that site. For instance, dysesthesias restricted to the fifth digit and the adjacent one-half of the fourth finger on one hand reliably point to disorder of the ulnar nerve, most commonly at the elbow.

Nerve and Root In focal nerve trunk lesions, sensory abnormalities are readily mapped and generally have discrete boundaries (Figs. 31-2 and 31-3). Root (“radicular”) lesions frequently are accompanied by deep, aching pain along the course of the related nerve trunk. With compression of a fifth lumbar (L5) or first sacral (S1) root, as from a ruptured intervertebral disk, sciatica (radicular pain relating to the sciatic nerve trunk) is a common manifestation (Chap. 22). With a lesion affecting a single root, sensory deficits may be minimal or absent because adjacent root territories overlap extensively.

Isolated mononeuropathies may cause symptoms beyond the territory supplied by the affected nerve, but abnormalities on examination typically are confined to appropriate anatomic boundaries. In multiple mononeuropathies, symptoms and signs occur in discrete territories supplied by different individual nerves and—as more nerves are affected—may simulate a polyneuropathy if deficits become confluent. With polyneuropathies, sensory deficits are generally graded, distal, and symmetric in distribution (Chap. 459). Dysesthesias, followed by numbness, begin in the toes and ascend symmetrically. When dysesthesias reach the knees, they usually also have appeared in the fingertips. The process is nerve length-dependent, and the deficit is often described as “stocking-glove” in type. Involvement of both hands and feet also occurs with lesions of the upper cervical cord or the brainstem, but an upper level of the sensory disturbance may then be found on the trunk and other evidence of a central lesion may be present, such as sphincter involvement or signs of an upper motor neuron lesion (Chap. 30). Although most polyneuropathies are pansensory and affect all modalities of sensation, selective sensory dysfunction

according to nerve fiber size may occur. Small-fiber polyneuropathies are characterized by burning, painful dysesthesias with reduced pinprick and thermal sensation but with sparing of proprioception, motor function, and deep tendon reflexes. Touch is involved variably; when it is spared, the sensory pattern is referred to as exhibiting *sensory dissociation*. Sensory dissociation may occur also with spinal cord lesions as well as small-fiber neuropathies. Large-fiber polyneuropathies are characterized by vibration and position sense deficits, imbalance, absent tendon reflexes, and variable motor dysfunction but preservation of most cutaneous sensation. Dysesthesias, if present at all, tend to be tingling or bandlike in quality.

Sensory neuronopathy (or ganglionopathy) is characterized by widespread but asymmetric sensory loss occurring in a non-length-dependent manner so that it may occur proximally or distally and in the arms, legs, or both. Pain and numbness progress to sensory ataxia and impairment of all sensory modalities with time. This condition is usually paraneoplastic or idiopathic in origin (Chaps. 122 and 459) or related to an autoimmune disease, particularly Sjögren’s syndrome.

Spinal Cord (See also Chap. 456) If the spinal cord is transected, all sensation is lost below the level of transection. Bladder and bowel function also are lost, as is motor function. Lateral hemisection of the spinal cord produces the Brown-Séquard syndrome, with absent pain and temperature sensation contralaterally and loss of proprioceptive sensation and power ipsilaterally below the lesion (see Figs. 31-1 and 456-1).

Numbness or paresthesias in both feet may arise from a spinal cord lesion; this is especially likely when the upper level of the sensory loss extends to the trunk. When all extremities are affected, the lesion is probably in the cervical region or brainstem unless a peripheral neuropathy is responsible. The presence of upper motor neuron signs (Chap. 30) supports a central lesion; a hyperesthetic band on the trunk may suggest the level of involvement.

A dissociated sensory loss can reflect spinothalamic tract involvement in the spinal cord, especially if the deficit is unilateral and has an upper level on the torso. Bilateral spinothalamic tract involvement occurs with lesions affecting the center of the spinal cord, such as in syringomyelia. There is a dissociated sensory loss with impairment of pinprick and temperature appreciation but relative preservation of light touch, position sense, and vibration appreciation.

Dysfunction of the posterior columns in the spinal cord or of the posterior root entry zone may lead to a bandlike sensation around the trunk or a feeling of tight pressure in one or more limbs. Flexion of the neck sometimes leads to an electric shock–like sensation that radiates down the back and into the legs (Lhermitte’s sign) in patients with a cervical lesion affecting the posterior columns, such as from multiple sclerosis, cervical spondylosis, or recent irradiation to the cervical region.

Brainstem Crossed patterns of sensory disturbance, in which one side of the face and the opposite side of the body are affected, localize to the lateral medulla. Here a small lesion may damage both the ipsilateral descending trigeminal tract and the ascending spinothalamic fibers subserving the opposite arm, leg, and hemitorso (see “Lateral medullary syndrome” in Fig. 446-10). A lesion in the tegmentum of the pons and midbrain, where the lemniscal and spinothalamic tracts merge, causes pansensory loss contralaterally.

Thalamus Hemisensory disturbance with tingling numbness from head to foot is often thalamic in origin but also can arise from the anterior parietal region. If abrupt in onset, the lesion is likely to be due to a small stroke (lacunar infarction), particularly if localized to the thalamus. Occasionally, with lesions affecting the VPL nucleus or adjacent white matter, a syndrome of thalamic pain, also called *Déjerine-Roussy syndrome*, may ensue. The persistent, unremitting unilateral pain often is described in dramatic terms.

Cortex With lesions of the parietal lobe involving either the cortex or the subjacent white matter, the most prominent symptoms are contralateral hemineglect, hemi-inattention, and a tendency not to use

the affected hand and arm. On cortical sensory testing (e.g., two-point discrimination, graphesthesia), abnormalities are often found but primary sensation is usually intact. Anterior parietal infarction may present as a pseudothalamic syndrome with contralateral loss of primary sensation from head to toe. Dysesthesias or a sense of numbness and, rarely, a painful state may also occur.

Focal Sensory Seizures These seizures generally are due to lesions in the area of the postcentral or precentral gyrus. The principal symptom of focal sensory seizures is tingling, but additional, more complex sensations may occur, such as a rushing feeling, a sense of warmth, or a sense of movement without detectable motion. Symptoms typically are unilateral; commonly begin in the arm or hand, face, or foot; and often spread in a manner that reflects the cortical representation of different bodily parts, as in a Jacksonian march. Their duration is variable; seizures may be transient, lasting only for seconds, or persist for an hour or more. Focal motor features may supervene, often becoming generalized with loss of consciousness and tonic-clonic jerking.

ACKNOWLEDGMENT

Arthur Asbury authored or co-authored this chapter in earlier editions of this book.

32

Gait and Balance Disorders

Lewis Sudarsky

PREVALENCE, MORBIDITY, AND MORTALITY

Gait and balance problems are common in the elderly and contribute to the risk of falls and injury. Gait disorders have been described in 15% of individuals older than 65. By age 80 one person in four will use a mechanical aid to assist with ambulation. Among those 85 and older, the prevalence of gait abnormality approaches 40%. In epidemiologic studies, gait disorders are consistently identified as a major risk factor for falls and injury.

A substantial number of older persons report insecure balance and experience falls and fear of falling. Prospective studies indicate that 30% of those older than 65 fall each year. The proportion is even higher in frail elderly and nursing home patients. Each year, 8% of individuals older than 75 suffer a serious fall-related injury. Hip fractures result in hospitalization, can lead to nursing home admission, and are associated with an increased mortality risk in the subsequent year. For each person who is physically disabled, there are others whose functional independence is limited by anxiety and fear of falling. Nearly one in five elderly individuals voluntarily restricts his or her activity because of fear of falling. With loss of ambulation, the quality of life diminishes, and rates of morbidity and mortality increase.

ANATOMY AND PHYSIOLOGY

An upright bipedal gait depends on the successful integration of postural control and locomotion. These functions are widely distributed in the central nervous system. The biomechanics of bipedal walking are complex, and the performance is easily compromised by a neurologic deficit at any level. Command and control centers in the brainstem, cerebellum, and forebrain modify the action of spinal pattern generators to promote stepping. While a form of “fictive locomotion” can be elicited from quadrupedal animals after spinal transection, this capacity is limited in primates. Step generation in primates is dependent on locomotor centers in the pontine tegmentum, midbrain, and subthalamic region. Locomotor synergies are executed through the reticular formation and descending pathways in the ventromedial spinal cord. Cerebral control provides a goal and purpose for walking

and is involved in avoidance of obstacles and adaptation of locomotor programs to context and terrain.

Postural control requires the maintenance of the center of mass over the base of support through the gait cycle. Unconscious postural adjustments maintain standing balance: long latency responses are measurable in the leg muscles, beginning 110 milliseconds after a perturbation. Forward motion of the center of mass provides propulsive force for stepping, but failure to maintain the center of mass within stability limits results in falls. The anatomic substrate for dynamic balance has not been well defined, but the vestibular nucleus and midline cerebellum contribute to balance control in animals. Patients with damage to these structures have impaired balance while standing and walking.

Standing balance depends on good-quality sensory information about the position of the body center with respect to the environment, support surface, and gravitational forces. Sensory information for postural control is primarily generated by the visual system, the vestibular system, and proprioceptive receptors in the muscle spindles and joints. A healthy redundancy of sensory afferent information is generally available, but loss of two of the three pathways is sufficient to compromise standing balance. Balance disorders in older individuals sometimes result from multiple insults in the peripheral sensory systems (e.g., visual loss, vestibular deficit, peripheral neuropathy) that critically degrade the quality of afferent information needed for balance stability.

Older patients with cognitive impairment from neurodegenerative diseases appear to be particularly prone to falls and injury. There is a growing body of literature on the use of attentional resources to manage gait and balance. Walking is generally considered to be unconscious and automatic, but the ability to walk while attending to a cognitive task (*dual-task walking*) may be compromised in frail elderly individuals with a history of falls. Older patients with deficits in executive function may have particular difficulty in managing the attentional resources needed for dynamic balance when distracted.

DISORDERS OF GAIT

Disorders of gait may be attributed to frailty, fatigue, arthritis, and orthopedic deformity, but neurologic causes are disabling and important to address. The heterogeneity of gait disorders observed in clinical practice reflects the large network of neural systems involved in the task. Walking is vulnerable to neurologic disease at every level. Gait disorders have been classified descriptively on the basis of abnormal physiology and biomechanics. One problem with this approach is that many failing gaits look fundamentally similar. This overlap reflects common patterns of adaptation to threatened balance stability and declining performance. *The gait disorder observed clinically must be viewed as the product of a neurologic deficit and a functional adaptation.* Unique features of the failing gait are often overwhelmed by the adaptive response. Some common patterns of abnormal gait are summarized next. Gait disorders can also be classified by etiology (Table 32-1).

TABLE 32-1 ETIOLOGY OF GAIT DISORDERS

Etiology	No. of Cases	Percent
Sensory deficits	22	18.3
Myelopathy	20	16.7
Multiple infarcts	18	15.0
Parkinsonism	14	11.7
Cerebellar degeneration	8	6.7
Hydrocephalus	8	6.7
Toxic/metabolic causes	3	2.5
Psychogenic causes	4	3.3
Other	6	5.0
Unknown causes	17	14.2
Total	120	100

Source: Reproduced with permission from J Masdeu, L Sudarsky, L Wolfson: *Gait Disorders of Aging*. Lippincott Raven, 1997.

CAUTIOUS GAIT

The term *cautious gait* is used to describe the patient who walks with an abbreviated stride and lowered center of mass, as if walking on a slippery surface. This disorder is both common and nonspecific. It is, in essence, an adaptation to a perceived postural threat. There may be an associated fear of falling. This disorder can be observed in more than one-third of older patients with gait impairment. Physical therapy often improves walking to the degree that follow-up observation may reveal a more specific underlying disorder.

STIFF-LEGGED GAIT

Spastic gait is characterized by stiffness in the legs, an imbalance of muscle tone, and a tendency to circumduct and scuff the feet. The disorder reflects compromise of corticospinal command and overactivity of spinal reflexes. The patient may walk on the toes. In extreme instances, the legs cross due to increased tone in the adductors. Upper motor neuron signs are present on physical examination. Shoes often reflect an uneven pattern of wear across the outside. The disorder may be cerebral or spinal in origin.

Myelopathy from cervical spondylosis is a common cause of spastic or spastic-ataxic gait in the elderly. Demyelinating disease and trauma are the leading causes of myelopathy in younger patients. In chronic progressive myelopathy of unknown cause, a workup with laboratory and imaging tests may establish a diagnosis. A family history should suggest hereditary spastic paraplegia (**Chap. 452**); genetic testing is now available for some of the common mutations responsible for this disorder. Tropical spastic paraparesis related to the retrovirus human T-cell lymphotropic virus 1 (HTLV-1) is endemic in parts of the Caribbean and South America. A structural lesion, such as a tumor or a spinal vascular malformation, should be excluded with appropriate testing. **Spinal cord disorders are discussed in detail in Chap. 456.**

With cerebral spasticity, asymmetry is common, the upper extremities are usually involved, and dysarthria is often an associated feature. Common causes include vascular disease (stroke), multiple sclerosis, and perinatal injury to the nervous system (cerebral palsy).

Other stiff-legged gaits include dystonia (**Chap. 449**) and stiff-person syndrome (**Chap. 122**). Dystonia is a disorder characterized by sustained muscle contractions resulting in repetitive twisting movements and abnormal posture. It often has a genetic basis. Dystonic spasms can produce plantar flexion and inversion of the feet, sometimes with torsion of the trunk. In autoimmune stiff-person syndrome, exaggerated lordosis of the lumbar spine and overactivation of antagonist muscles restrict trunk and lower-limb movement and result in a wooden or fixed posture.

PARKINSONISM AND FREEZING GAIT

Parkinson's disease (**Chap. 449**) is common, affecting 1% of the population >55 years of age. The stooped posture and shuffling gait are characteristic and distinctive features. Patients sometimes accelerate (festinate) with walking, display retropulsion, or exhibit a tendency to turn en bloc. A National Institutes of Health workshop defined freezing of gait as "brief, episodic absence of forward progression of the feet, despite the intention to walk." Gait freezing occurs in 26% of Parkinson's patients by the end of 5 years and develops in most such patients eventually. Postural instability and falling occur as the disease progresses; some falls are precipitated by freezing of gait.

Freezing of gait is even more common in some Parkinson's-related neurodegenerative disorders, such as progressive supranuclear palsy, multiple-system atrophy, and corticobasal degeneration. Patients with these disorders frequently present with axial stiffness, postural instability, and a shuffling, freezing gait while lacking the characteristic pill-rolling tremor of Parkinson's disease. Falls within the first year suggest the possibility of progressive supranuclear palsy.

Hyperkinetic movement disorders also produce characteristic and recognizable disturbances in gait. In Huntington's disease (**Chap. 449**), the unpredictable occurrence of choreic movements gives the gait a dancing quality. Tardive dyskinesia is the cause of many odd, stereotypic gait disorders seen in patients chronically exposed to antipsychotics and other drugs that block the D₂ dopamine receptor.

FRONTAL GAIT DISORDER

Frontal gait disorder, sometimes known as *gait apraxia*, is common in the elderly and has a variety of causes. The term is used to describe a shuffling, freezing gait with imbalance and other signs of higher cerebral dysfunction. Typical features include a wide base of support, a short stride, shuffling along the floor, and difficulty with starts and turns. Many patients exhibit a difficulty with gait initiation that is descriptively characterized as the "slipping clutch" syndrome or gait ignition failure. The term *lower-body parkinsonism* is also used to describe such patients. Strength is generally preserved, and patients are able to make stepping movements when not standing and maintaining their balance at the same time. This disorder is best considered a higher-level motor control disorder, as opposed to an apraxia (**Chap. 36**).

The most common cause of frontal gait disorder is vascular disease, particularly subcortical small-vessel disease. Lesions are frequently found in the deep frontal white matter and centrum ovale. Gait disorder may be the salient feature in hypertensive patients with ischemic lesions of the deep-hemisphere white matter (*Binswanger's disease*). The clinical syndrome includes mental changes (variable in degree), dysarthria, pseudobulbar affect (emotional disinhibition), increased tone, and hyperreflexia in the lower limbs.

Communicating hydrocephalus in adults also presents with a gait disorder of this type. Other features of the diagnostic triad (mental changes, incontinence) may be absent in the initial stages. MRI demonstrates ventricular enlargement, an enlarged flow void about the aqueduct, and a variable degree of periventricular white-matter change. A lumbar puncture or dynamic test is necessary to confirm hydrocephalus.

CEREBELLAR GAIT ATAXIA

Disorders of the cerebellum have a dramatic impact on gait and balance. Cerebellar gait ataxia is characterized by a wide base of support, lateral instability of the trunk, erratic foot placement, and decompensation of balance when attempting to walk on a narrow base. Difficulty maintaining balance when turning is often an early feature. Patients are unable to walk tandem heel to toe and display truncal sway in narrow-based or tandem stance. They show considerable variation in their tendency to fall in daily life.

Causes of cerebellar ataxia in older patients include stroke, trauma, tumor, and neurodegenerative disease such as multiple-system atrophy (**Chaps. 449 and 454**) and various forms of hereditary cerebellar degeneration (**Chap. 450**). A short expansion at the site of the fragile X mutation (*fragile X pre-mutation*) has been associated with gait ataxia in older men. Alcoholic cerebellar degeneration can be screened by history and often confirmed by MRI. In patients with ataxia, MRI demonstrates the extent and topography of cerebellar atrophy.

SENSORY ATAXIA

As reviewed earlier in this chapter, balance depends on high-quality afferent information from the visual and the vestibular systems and proprioception. When this information is lost or degraded, balance during locomotion is impaired and instability results. The sensory ataxia of tabetic neurosyphilis is a classic example. The contemporary equivalent is the patient with neuropathy affecting large fibers. Vitamin B₁₂ deficiency is a treatable cause of large-fiber sensory loss in the spinal cord and peripheral nervous system. Joint position and vibration sense are diminished in the lower limbs. The stance in such patients is destabilized by eye closure; they often look down at their feet when walking and do poorly in the dark. **Table 32-2** compares sensory ataxia with cerebellar ataxia and frontal gait disorder. Some frail older patients exhibit a syndrome of imbalance from the combined effect of multiple sensory deficits. Such patients have disturbances in proprioception, vision, and vestibular sense that impair postural support.

NEUROMUSCULAR DISEASE

Patients with neuromuscular disease often have an abnormal gait, occasionally as a presenting feature. With distal weakness (peripheral neuropathy), the step height is increased to compensate for footdrop, and the sole of the foot may slap on the floor during weight acceptance. Neuropathy may be associated with a degree of sensory imbalance, as described earlier. Patients with myopathy or muscular dystrophy more

TABLE 32-2 FEATURES OF CEREBELLAR ATAXIA, SENSORY ATAXIA, AND FRONTAL GAIT DISORDERS

Feature	Cerebellar Ataxia	Sensory Ataxia	Frontal Gait
Base of support	Wide-based	Narrow base, looks down	Wide-based
Velocity	Variable	Slow	Very slow
Stride	Irregular, lurching	Regular with path deviation	Short, shuffling
Romberg test	+/-	Unsteady, falls	+/-
Heel → shin	Abnormal	+/-	Normal
Initiation	Normal	Normal	Hesitant
Turns	Unsteady	+/-	Hesitant, multistep
Postural instability	+	+++	++++ Poor postural synergies rising from a chair
Falls	Late event	Frequent	Frequent

typically exhibit proximal weakness. Weakness of the hip girdle may result in some degree of excess pelvic sway during locomotion.

TOXIC AND METABOLIC DISORDERS

Alcohol intoxication is the most common cause of acute walking difficulty. Chronic toxicity from medications and metabolic disturbances can impair motor function and gait. Mental status changes may be found, and examination may reveal asterixis or myoclonus. Static equilibrium is disturbed, and such patients are easily thrown off balance. Disequilibrium is particularly evident in patients with chronic renal disease and those with hepatic failure, in whom asterixis may impair postural support. Sedative drugs, especially neuroleptics and long-acting benzodiazepines, affect postural control and increase the risk for falls. These disorders are especially important to recognize because they are often treatable.

PSYCHOGENIC GAIT DISORDER

Psychogenic disorders are common in neurologic practice, and the presentation often involves gait. Some patients with extreme anxiety or phobia walk with exaggerated caution with abduction of the arms, as if walking on ice. This inappropriately overcautious gait differs in degree from the gait of the patient who is insecure and making adjustments for imbalance. Depressed patients exhibit primarily slowness, a manifestation of psychomotor retardation, and lack of purpose in their stride. Hysterical gait disorders are among the most spectacular encountered. Odd gyrations of posture with wastage of muscular energy (astasia-abasia), extreme slow motion, and dramatic fluctuations over time may be observed in patients with somatoform disorders and conversion reactions.

APPROACH TO THE PATIENT:

Slowly Progressive Disorder of Gait

When reviewing the history, it is helpful to inquire about the onset and progression of disability. Initial awareness of an unsteady gait often follows a fall. Stepwise evolution or sudden progression suggests vascular disease. Gait disorder may be associated with urinary urgency and incontinence, particularly in patients with cervical spine disease or hydrocephalus. It is always important to review the use of alcohol and medications that affect gait and balance. Information on localization derived from the neurologic examination can be helpful in narrowing the list of possible diagnoses.

Gait observation provides an immediate sense of the patient's degree of disability. Arthritic and antalgic gaits are recognized by observation, though neurologic and orthopedic problems may coexist. Characteristic patterns of abnormality are sometimes seen, though, as stated previously, failing gaits often look fundamentally similar. Cadence (steps per minute), velocity, and stride length can

be recorded by timing a patient over a fixed distance. Watching the patient rise from a chair provides a good functional assessment of balance.

Brain imaging studies may be informative in patients with an undiagnosed disorder of gait. MRI is sensitive for cerebral lesions of vascular or demyelinating disease and is a good screening test for occult hydrocephalus. Patients with recurrent falls are at risk for subdural hematoma. As mentioned earlier, many elderly patients with gait and balance difficulty have white matter abnormalities in the periventricular region and centrum semiovale. While these lesions may be an incidental finding, a substantial burden of white matter disease will ultimately impact cerebral control of locomotion.

DISORDERS OF BALANCE

DEFINITION, ETIOLOGY, AND MANIFESTATIONS

Balance is the ability to maintain equilibrium—a state in which opposing physical forces cancel one another out. In physiology, this term is taken to mean the ability to control the center of mass with respect to gravity and the support surface. In reality, people are not consciously aware of their center of mass, but everyone (particularly gymnasts, figure skaters, and platform divers, for example) move so as to manage it. Disorders of balance present as difficulty maintaining posture while standing and walking and as a subjective sense of disequilibrium, which is a form of dizziness.

The cerebellum and vestibular system organize antigravity responses needed to maintain an upright posture. These responses are physiologically complex, and the anatomic representation they entail is not well understood. Failure, resulting in disequilibrium, can occur at several levels: cerebellar, vestibular, somatosensory, and higher-level disequilibrium.

Patients with *cerebellar ataxia* do not generally complain of dizziness, though balance is visibly impaired. Neurologic examination reveals a variety of cerebellar signs. Postural compensation may prevent falls early on, but falls are inevitable with disease progression. The progression of neurodegenerative ataxia is often measured by the number of years to loss of stable ambulation.

Vestibular disorders (Chap. 28) have symptoms and signs that fall into three categories: (1) vertigo (the subjective inappropriate perception or illusion of movement); (2) nystagmus (involuntary eye movements); and (3) impaired standing balance. Not every patient has all manifestations. Patients with vestibular deficits related to ototoxic drugs may lack vertigo or obvious nystagmus, but their balance is impaired on standing and walking, and they cannot navigate in the dark. Laboratory testing is available to investigate vestibular deficits.

Somatosensory deficits also produce imbalance and falls. There is often a subjective sense of insecure balance and fear of falling. Postural control is compromised by eye closure (*Romberg's sign*); these patients also have difficulty navigating in the dark. A dramatic example is provided by the patient with autoimmune subacute sensory neuropathy, which is sometimes a paraneoplastic disorder (Chap. 122). Compensatory strategies enable such patients to walk in the virtual absence of proprioception, but the task requires active visual monitoring.

Patients with *higher-level disorders of equilibrium* have difficulty maintaining balance in daily life and may present with falls. Their awareness of balance impairment may be reduced. Patients taking sedating medications are in this category. In prospective studies, dementia and sedating medications substantially increase the risk for falls.

FALLS

Falls are common in the elderly; 30% of people older than 65 who are living in the community fall each year. Modest changes in balance function have been described in fit older individuals as a result of normal aging. Subtle deficits in sensory systems, attention, and motor reaction time contribute to the risk, and environmental hazards abound. Many falls by older adults are episodes of tripping or slipping, often designated *mechanical falls*. A fall is not a neurologic problem per se, but there are events for which neurologic evaluation is appropriate.

It is important to distinguish falls associated with loss of consciousness (syncope, seizure), which require appropriate evaluation and intervention (Chaps. 27 and 445). In most prospective studies, a small subset of individuals experience a large number of fall events. These individuals with recurrent falls often have gait and balance issues that need to be addressed.

Fall Patterns: The Event Description The history of a fall is often problematic or incomplete, and the underlying mechanism or cause may be difficult to establish in retrospect. The patient and family may have limited information about what triggered the fall. Injuries can complicate the physical examination. While there is no standard nosology of falls, some common clinical patterns may emerge and provide a clue.

DROP ATTACKS AND COLLAPSING FALLS Drop attacks are sudden collapsing falls without loss of consciousness. Patients who collapse from lack of postural tone present a diagnostic challenge. Patients may report that their legs just “gave out” underneath them; their families may describe these patients as “collapsing in a heap.” Orthostatic hypotension may be a factor in some such falls, and this possibility should be thoroughly evaluated. Rarely, a colloid cyst of the third ventricle can present with intermittent obstruction of the foramen of Monro, with a consequent drop attack. While collapsing falls are more common among older patients with vascular risk factors, they should not be confused with vertebrobasilar ischemic attacks.

TOPPLING FALLS Some patients maintain tone in antigravity muscles but fall over like a tree trunk, as if postural defenses had disengaged. There may be a consistent direction to such falls. The patient with cerebellar pathology may lean and topple over toward the side of the lesion. Patients with lesions of the vestibular system or its central pathways may experience lateral pulsion and toppling falls. Patients with progressive supranuclear palsy often fall over backward. Falls of this nature occur in patients with advanced Parkinson’s disease once postural instability has developed.

FALLS DUE TO GAIT FREEZING Another fall pattern in Parkinson’s disease and related disorders is the fall due to freezing of gait. The feet stick to the floor and the center of mass keeps moving, resulting in a disequilibrium from which the patient has difficulty recovering. This sequence of events can result in a forward fall. Gait freezing can also occur as the patient attempts to turn and change direction. Similarly, patients with Parkinson’s disease and festinating gait may find their feet unable to keep up and may thus fall forward.

FALLS RELATED TO SENSORY LOSS Patients with somatosensory, visual, or vestibular deficits are prone to falls. These patients have particular difficulty dealing with poor illumination or walking on uneven ground. They often report subjective imbalance, apprehension, and fear of falling. Deficits in joint position and vibration sense are apparent on physical examination. These patients may be especially responsive to a rehabilitation-based intervention.

WEAKNESS AND FRAILTY Patients who lack strength in antigravity muscles have difficulty rising from a chair, tire easily when walking, and have difficulty maintaining their balance after a perturbation. These patients are often unable to get up after a fall and may have to remain on the floor for a prolonged period until help arrives. Deconditioning of this sort is often treatable. Resistance strength training can increase muscle mass and leg strength, even for people in their eighties and nineties.

RISK FACTORS FOR FALLS

The most productive approach is to identify the high-risk patient prospectively, before there is a serious injury. Patients at particular risk include hospitalized patients with mental status changes, nursing home residents, patients with dementia, and those taking medications that compromise attention and alertness. Patients with Parkinson’s disease and other gait disorders are also at increased risk. (Table 32-3) summarizes a meta-analysis of prospective studies establishing the principal risk factors for falls. It is often possible to address and mitigate some of the major risk factors. Medication overuse may be the most important remediable risk factor for falls.

TABLE 32-3 META-ANALYSIS OF RISK FACTORS FOR FALLS: SUMMARY OF 16 CONTROLLED STUDIES

Risk Factor	Mean RR (OR)	Range
Weakness	4.9	1.9–10.3
Balance deficit	3.2	1.6–5.4
Gait disorder	3.0	1.7–4.8
Visual deficit	2.8	1.1–7.4
Mobility limitation	2.5	1.0–5.3
Cognitive impairment	2.4	2.0–4.7
Impaired functional status	2.0	1.0–3.1
Postural hypotension	1.9	1.0–3.4

Abbreviations: OR, odds ratio from retrospective studies; RR, relative risk from prospective studies.

Source: Reproduced with permission from J Masdeu, L Sudarsky, L Wolfson: *Gait Disorders of Aging*. Lippincott Raven, 1997.

TREATMENT INTERVENTIONS TO REDUCE THE RISK OF FALLS AND INJURY

Efforts should be made to define the etiology of the gait disorder and the mechanism underlying the falls by a given patient. Orthostatic changes in blood pressure and pulse should be recorded. Rising from a chair and walking should be evaluated for safety. Specific treatment may be possible once a diagnosis is established. Therapeutic intervention is often recommended for older patients at substantial risk for falls, even if no neurologic disease is identified. A home visit to look for environmental hazards can be helpful. A variety of modifications may be recommended to improve safety, including improved lighting and the installation of grab bars and nonslip surfaces.

Rehabilitative interventions aim to improve muscle strength and balance stability and to make the patient more resistant to injury. High-intensity resistance strength training with weights and machines is useful to improve muscle mass, even in frail older patients. Improvements realized in posture and gait should translate to reduced risk of falls and injury. Sensory balance training is another approach to improving balance stability. Measurable gains can be made in a few weeks of training, and benefits can be maintained over 6 months by a 10- to 20-min home exercise program. This strategy is particularly successful in patients with vestibular and somatosensory balance disorders. A Tai Chi exercise program has been demonstrated to reduce the risk of falls and injury in patients with Parkinson’s disease.



33e

Video Library of Gait Disorders

Gail Kang, Nicholas B. Galifianakis, Michael D. Geschwind

Problems with gait and balance are major causes of falls, accidents, and resulting disability, especially in later life, and are often harbingers of neurologic disease. Early diagnosis is essential, especially for treatable conditions, because it may permit the institution of prophylactic measures to prevent dangerous falls and also to reverse or ameliorate the underlying cause. In this video, examples of gait disorders due to Parkinson's disease, other extrapyramidal disorders, and ataxias, as well as other common gait disorders, are presented.

Confusion, a mental and behavioral state of reduced comprehension, coherence, and capacity to reason, is one of the most common problems encountered in medicine, accounting for a large number of emergency department visits, hospital admissions, and inpatient consultations. *Delirium*, a term used to describe an acute confusional state, remains a major cause of morbidity and mortality, costing over \$150 billion dollars yearly in health care costs in the United States alone. Despite increased efforts targeting awareness of this condition, delirium often goes unrecognized in the face of evidence that it is usually the cognitive manifestation of serious underlying medical or neurologic illness.

CLINICAL FEATURES OF DELIRIUM

A multitude of terms are used to describe patients with delirium, including *encephalopathy*, *acute brain failure*, *acute confusional state*, and *postoperative* or *intensive care unit (ICU) psychosis*. Delirium has many clinical manifestations, but is defined as a relatively acute decline in cognition that fluctuates over hours or days. The hallmark of delirium is a deficit of attention, although all cognitive domains—including memory, executive function, visuospatial tasks, and language—are variably involved. Associated symptoms that may be present in some cases include altered sleep-wake cycles, perceptual disturbances such as hallucinations or delusions, affect changes, and autonomic findings that include heart rate and blood pressure instability.

Delirium is a clinical diagnosis that is made only at the bedside. Two subtypes have been described—hyperactive and hypoactive—based on differential psychomotor features. The cognitive syndrome associated with severe alcohol withdrawal (i.e., “delirium tremens”) remains the classic example of the hyperactive subtype, featuring prominent hallucinations, agitation, and hyperarousal, often accompanied by life-threatening autonomic instability. In striking contrast is the hypoactive subtype, exemplified by benzodiazepine intoxication, in which patients are withdrawn and quiet, with prominent apathy and psychomotor slowing.

This dichotomy between subtypes of delirium is a useful construct, but patients often fall somewhere along a spectrum between the hyperactive and hypoactive extremes, sometimes fluctuating from one to the other. Therefore, clinicians must recognize this broad range of presentations of delirium to identify all patients with this potentially reversible cognitive disturbance. Hyperactive patients are often easily recognized by their characteristic severe agitation, tremor, hallucinations, and autonomic instability. Patients who are quietly hypoactive are more often overlooked on the medical wards and in the ICU.

The reversibility of delirium is emphasized because many etiologies, such as systemic infection and medication effects, can be treated easily. The long-term cognitive effects of delirium remain largely unknown. Some episodes of delirium continue for weeks, months, or even years. The persistence of delirium in some patients and its high recurrence rate may be due to inadequate initial treatment of the underlying etiology. In other instances, delirium appears to cause permanent neuronal damage and cognitive decline. Even if an episode of delirium completely resolves, there may be lingering effects of the disorder; a patient’s recall of events after delirium varies widely, ranging from complete amnesia to repeated re-experiencing of the frightening period of confusion, similar to what is seen in patients with posttraumatic stress disorder.

RISK FACTORS

An effective primary prevention strategy for delirium begins with identification of patients at high risk for this disorder, including those preparing for elective surgery or being admitted to the hospital. Although no single validated scoring system has been widely accepted as a screen for asymptomatic patients, there are multiple well-established risk factors for delirium.

The two most consistently identified risks are older age and baseline cognitive dysfunction. Individuals who are over age 65 or exhibit low scores on standardized tests of cognition develop delirium upon hospitalization at a rate approaching 50%. Whether age and baseline cognitive dysfunction are truly independent risk factors is uncertain. Other predisposing factors include sensory deprivation, such as preexisting hearing and visual impairment, as well as indices for poor overall health, including baseline immobility, malnutrition, and underlying medical or neurologic illness.

In-hospital risks for delirium include the use of bladder catheterization, physical restraints, sleep and sensory deprivation, and the addition of three or more new medications. Avoiding such risks remains a key component of delirium prevention as well as treatment. Surgical and anesthetic risk factors for the development of postoperative delirium include specific procedures such as those involving cardiopulmonary bypass, inadequate or excessive treatment of pain in the immediate postoperative period, and perhaps specific agents such as inhalational anesthetics.

The relationship between delirium and dementia ([Chap. 448](#)) is complicated by significant overlap between the two conditions, and it is not always simple to distinguish between them. Dementia and preexisting cognitive dysfunction serve as major risk factors for delirium, and at least two-thirds of cases of delirium occur in patients with coexisting underlying dementia. A form of dementia with parkinsonism, termed *dementia with Lewy bodies*, is characterized by a fluctuating course, prominent visual hallucinations, parkinsonism, and an attentional deficit that clinically resembles hyperactive delirium; patients with this condition are particularly vulnerable to delirium. Delirium in the elderly often reflects an insult to the brain that is vulnerable due to an underlying neurodegenerative condition. Therefore, the development of delirium sometimes heralds the onset of a previously unrecognized brain disorder.

EPIDEMIOLOGY

Delirium is common, but its reported incidence has varied widely with the criteria used to define this disorder. Estimates of delirium in hospitalized patients range from 18 to 64%, with higher rates reported for elderly patients and patients undergoing hip surgery. Older patients in the ICU have especially high rates of delirium that approach 75%. The condition is not recognized in up to one-third of delirious inpatients, and the diagnosis is especially problematic in the ICU environment, where cognitive dysfunction is often difficult to appreciate in the setting of serious systemic illness and sedation. Delirium in the ICU should be viewed as an important manifestation of organ dysfunction not unlike liver, kidney, or heart failure. Outside the acute hospital setting, delirium occurs in nearly one-quarter of patients in nursing homes and in 50 to 80% of those at the end of life. These estimates emphasize the remarkably high frequency of this cognitive syndrome in older patients, a population expected to grow in the upcoming decades.

Until recently, an episode of delirium was viewed as a transient condition that carried a benign prognosis. It is now recognized as a disorder with a substantial morbidity rate and increased mortality rate and often represents the first manifestation of a serious underlying illness. Recent estimates of in-hospital mortality rates among delirious patients have ranged from 25 to 33%, a rate similar to that of patients with sepsis. Patients with an in-hospital episode of delirium have a fivefold higher mortality rate in the months after their illness compared with age-matched nondelirious hospitalized patients. Delirious hospitalized patients have a longer length of stay, are more likely to be discharged to a nursing home, and are more likely to experience subsequent episodes of delirium and cognitive decline; as a result, this condition has enormous economic implications.

PATHOGENESIS

The pathogenesis and anatomy of delirium are incompletely understood. The attentional deficit that serves as the neuropsychological hallmark of delirium has a diffuse localization within the brainstem, thalamus, prefrontal cortex, and parietal lobes. Rarely, focal lesions

such as ischemic strokes have led to delirium in otherwise healthy persons; right parietal and medial dorsal thalamic lesions have been reported most commonly, pointing to the importance of these areas to delirium pathogenesis. In most cases, delirium results from widespread disturbances in cortical and subcortical regions rather than a focal neuroanatomic cause. Electroencephalogram (EEG) data in persons with delirium usually show symmetric slowing, a nonspecific finding that supports diffuse cerebral dysfunction.

Multiple neurotransmitter abnormalities, proinflammatory factors, and specific genes likely play a role in the pathogenesis of delirium. Deficiency of acetylcholine may play a key role, and medications with anticholinergic properties also can precipitate delirium. Dementia patients are susceptible to episodes of delirium, and those with Alzheimer's pathology and dementia with Lewy bodies or Parkinson's disease dementia are known to have a chronic cholinergic deficiency state due to degeneration of acetylcholine-producing neurons in the basal forebrain. Additionally, other neurotransmitters are also likely to be involved in this diffuse cerebral disorder. For example, increases in dopamine can also lead to delirium. Patients with Parkinson's disease treated with dopaminergic medications can develop a delirium-like state that features visual hallucinations, fluctuations, and confusion.

Not all individuals exposed to the same insult will develop signs of delirium. A low dose of an anticholinergic medication may have no cognitive effects on a healthy young adult but produce a florid delirium in an elderly person with known underlying dementia, although even healthy young persons develop delirium with very high doses of anticholinergic medications. This concept of delirium developing as the result of an insult in predisposed individuals is currently the most widely accepted pathogenic construct. Therefore, if a previously healthy individual with no known history of cognitive illness develops delirium in the setting of a relatively minor insult such as elective surgery or hospitalization, an unrecognized underlying neurologic illness such as a neurodegenerative disease, multiple previous strokes, or another diffuse cerebral cause should be considered. In this context, delirium can be viewed as a "stress test for the brain" whereby exposure to known inciting factors such as systemic infection and offending drugs can unmask a decreased cerebral reserve and herald a serious underlying and potentially treatable illness.

APPROACH TO THE PATIENT: Delirium

Because the diagnosis of delirium is clinical and is made at the bedside, a careful history and physical examination are necessary in evaluating patients with possible confusional states. Screening tools can aid physicians and nurses in identifying patients with delirium, including the Confusion Assessment Method (CAM) (Table 34-1); the Organic Brain Syndrome Scale; the Delirium Rating Scale; and, in the ICU, the ICU version of the CAM and the Delirium Detection Score. Using the well-validated CAM, a diagnosis of delirium is made if there is (1) an acute onset and fluctuating course and (2) inattention accompanied by either (3) disorganized thinking or (4) an altered level of consciousness. These scales may not identify the full spectrum of patients with delirium, and all patients who are acutely confused should be presumed delirious regardless of their presentation due to the wide variety of possible clinical features. A course that fluctuates over hours or days and may worsen at night (termed *sundowning*) is typical but not essential for the diagnosis. Observation of the patient usually will reveal an altered level of consciousness or a deficit of attention. Other features that are sometimes present include alteration of sleep-wake cycles, thought disturbances such as hallucinations or delusions, autonomic instability, and changes in affect.

HISTORY

It may be difficult to elicit an accurate history in delirious patients who have altered levels of consciousness or impaired attention. Information from a collateral source such as a spouse or another

TABLE 34-1 THE CONFUSION ASSESSMENT METHOD (CAM) DIAGNOSTIC ALGORITHM^a

The diagnosis of delirium requires the presence of features 1 and 2 and of either feature 3 or 4.

Feature 1. Acute onset and fluctuating course

This feature is satisfied by positive responses to the following questions: Is there evidence of an acute change in mental status from the patient's baseline? Did the (abnormal) behavior fluctuate during the day, that is, tend to come and go, or did it increase and decrease in severity?

Feature 2. Inattention

This feature is satisfied by a positive response to the following question: Did the patient have difficulty focusing attention, for example, being easily distractible, or have difficulty keeping track of what was being said?

Feature 3. Disorganized thinking

This feature is satisfied by a positive response to the following question: Was the patient's thinking disorganized or incoherent, such as rambling or irrelevant conversation, unclear or illogical flow of ideas, or unpredictable switching from subject to subject?

Feature 4. Altered level of consciousness

This feature is satisfied by any answer other than "alert" to the following question: Overall, how would you rate the patient's level of consciousness: alert (normal), vigilant (hyperalert), lethargic (drowsy, easily aroused), stupor (difficult to arouse), or coma (unarousable)?

^aInformation is usually obtained from a reliable reporter, such as a family member, caregiver, or nurse.

Source: Modified from SK Inouye et al: Clarifying confusion: The Confusion Assessment Method. A new method for detection of delirium. *Ann Intern Med* 113:941, 1990.

family member is therefore invaluable. The three most important pieces of history are the patient's baseline cognitive function, the time course of the present illness, and current medications.

Premorbid cognitive function can be assessed through the collateral source or, if needed, via a review of outpatient records. Delirium by definition represents a change that is relatively acute, usually over hours to days, from a cognitive baseline. As a result, an acute confusional state is nearly impossible to diagnose without some knowledge of baseline cognitive function. Without this information, many patients with dementia or depression may be mistaken as delirious during a single initial evaluation. Patients with a more hypoactive, apathetic presentation with psychomotor slowing may be identified as being different from baseline only through conversations with family members. A number of validated instruments have been shown to diagnose cognitive dysfunction accurately using a collateral source, including the modified Blessed Dementia Rating Scale and the Clinical Dementia Rating (CDR). Baseline cognitive impairment is common in patients with delirium. Even when no such history of cognitive impairment is elicited, there should still be a high suspicion for a previously unrecognized underlying neurologic disorder.

Establishing the time course of cognitive change is important not only to make a diagnosis of delirium but also to correlate the onset of the illness with potentially treatable etiologies such as recent medication changes or symptoms of systemic infection.

Medications remain a common cause of delirium, especially compounds with anticholinergic or sedative properties. It is estimated that nearly one-third of all cases of delirium are secondary to medications, especially in the elderly. Medication histories should include all prescription as well as over-the-counter and herbal substances taken by the patient and any recent changes in dosing or formulation, including substitution of generics for brand-name medications.

Other important elements of the history include screening for symptoms of organ failure or systemic infection, which often contributes to delirium in the elderly. A history of illicit drug use, alcoholism, or toxin exposure is common in younger delirious patients. Finally, asking the patient and collateral source about other symptoms that may accompany delirium, such as depression, may help identify potential therapeutic targets.

PHYSICAL EXAMINATION

The general physical examination in a delirious patient should include careful screening for signs of infection such as fever, tachypnea, pulmonary consolidation, heart murmur, and stiff neck. The patient's fluid status should be assessed; both dehydration and fluid overload with resultant hypoxemia have been associated with delirium, and each is usually easily rectified. The appearance of the skin can be helpful, showing jaundice in hepatic encephalopathy, cyanosis in hypoxemia, or needle tracks in patients using intravenous drugs.

The neurologic examination requires a careful assessment of mental status. Patients with delirium often present with a fluctuating course; therefore, the diagnosis can be missed when one relies on a single time point of evaluation. Some but not all patients exhibit the characteristic pattern of sundowning, a worsening of their condition in the evening. In these cases, assessment only during morning rounds may be falsely reassuring.

An altered level of consciousness ranging from hyperarousal to lethargy to coma is present in most patients with delirium and can be assessed easily at the bedside. In a patient with a relatively normal level of consciousness, a screen for an attentional deficit is in order, because this deficit is the classic neuropsychological hallmark of delirium. Attention can be assessed while taking a history from the patient. Tangential speech, a fragmentary flow of ideas, or inability to follow complex commands often signifies an attentional problem. There are formal neuropsychological tests to assess attention, but a simple bedside test of digit span forward is quick and fairly sensitive. In this task, patients are asked to repeat successively longer random strings of digits beginning with two digits in a row, said to the patient at 1-second intervals. Healthy adults can repeat a string of five to seven digits before faltering; a digit span of four or less usually indicates an attentional deficit unless hearing or language barriers are present, and many patients with delirium have digit spans of three or fewer digits.

More formal neuropsychological testing can be helpful in assessing a delirious patient, but it is usually too cumbersome and time-consuming in the inpatient setting. A Mini-Mental State Examination (MMSE) provides information regarding orientation, language, and visuospatial skills; however, performance of many tasks on the MMSE, including the spelling of "world" backward and serial subtraction of digits, will be impaired by delirious patients' attentional deficits, rendering the test unreliable.

The remainder of the screening neurologic examination should focus on identifying new focal neurologic deficits. Focal strokes or mass lesions in isolation are rarely the cause of delirium, but patients with underlying extensive cerebrovascular disease or neurodegenerative conditions may not be able to cognitively tolerate even relatively small new insults. Patients should be screened for other signs of neurodegenerative conditions such as parkinsonism, which is seen not only in idiopathic Parkinson's disease but also in other dementing conditions such as Alzheimer's disease, dementia with Lewy bodies, and progressive supranuclear palsy. The presence of multifocal myoclonus or asterixis on the motor examination is nonspecific but usually indicates a metabolic or toxic etiology of the delirium.

ETIOLOGY

Some etiologies can be easily discerned through a careful history and physical examination, whereas others require confirmation with laboratory studies, imaging, or other ancillary tests. A large, diverse group of insults can lead to delirium, and the cause in many patients is often multifactorial. Common etiologies are listed in [Table 34-2](#).

Prescribed, over-the-counter, and herbal medications all can precipitate delirium. Drugs with anticholinergic properties, narcotics, and benzodiazepines are particularly common offenders, but nearly any compound can lead to cognitive dysfunction in a predisposed patient. Whereas an elderly patient with baseline dementia

TABLE 34-2 COMMON ETIOLOGIES OF DELIRIUM

Toxins
Prescription medications: especially those with anticholinergic properties, narcotics, and benzodiazepines
Drugs of abuse: alcohol intoxication and alcohol withdrawal, opiates, ecstasy, LSD, GHB, PCP, ketamine, cocaine, "bath salts," marijuana and its synthetic forms
Poisons: inhalants, carbon monoxide, ethylene glycol, pesticides
Metabolic conditions
Electrolyte disturbances: hypoglycemia, hyperglycemia, hyponatremia, hypernatremia, hypercalcemia, hypocalcemia, hypomagnesemia
Hypothermia and hyperthermia
Pulmonary failure: hypoxemia and hypercarbia
Liver failure/hepatic encephalopathy
Renal failure/uremia
Cardiac failure
Vitamin deficiencies: B ₁₂ , thiamine, folate, niacin
Dehydration and malnutrition
Anemia
Infections
Systemic infections: urinary tract infections, pneumonia, skin and soft tissue infections, sepsis
CNS infections: meningitis, encephalitis, brain abscess
Endocrine conditions
Hyperthyroidism, hypothyroidism
Hyperparathyroidism
Adrenal insufficiency
Cerebrovascular disorders
Global hypoperfusion states
Hypertensive encephalopathy
Focal ischemic strokes and hemorrhages (rare): especially nondominant parietal and thalamic lesions
Autoimmune disorders
CNS vasculitis
Cerebral lupus
Neurologic paraneoplastic syndromes
Seizure-related disorders
Nonconvulsive status epilepticus
Intermittent seizures with prolonged postictal states
Neoplastic disorders
Diffuse metastases to the brain
Gliomatosis cerebri
Carcinomatous meningitis
CNS lymphoma
Hospitalization
Terminal end-of-life delirium

Abbreviations: CNS, central nervous system; GHB, γ -hydroxybutyrate; LSD, lysergic acid diethylamide; PCP, phencyclidine.

may become delirious upon exposure to a relatively low dose of a medication, less susceptible individuals may become delirious only with very high doses of the same medication. This observation emphasizes the importance of correlating the timing of recent medication changes, including dose and formulation, with the onset of cognitive dysfunction.

In younger patients, illicit drugs and toxins are common causes of delirium. In addition to more classic drugs of abuse, the recent rise in availability of methylenedioxymethamphetamine (MDMA, ecstasy), γ -hydroxybutyrate (GHB), "bath salts," synthetic cannabis, and the phencyclidine (PCP)-like agent ketamine, has led to an increase in delirious young persons presenting to acute care settings ([Chap. 469e](#)). Many common prescription drugs such as oral

narcotics and benzodiazepines are often abused and readily available on the street. Alcohol abuse leading to high serum levels causes confusion, but more commonly, it is withdrawal from alcohol that leads to a hyperactive delirium. Alcohol and benzodiazepine withdrawal should be considered in all cases of delirium because even patients who drink only a few servings of alcohol every day can experience relatively severe withdrawal symptoms upon hospitalization.

Metabolic abnormalities such as electrolyte disturbances of sodium, calcium, magnesium, or glucose can cause delirium, and mild derangements can lead to substantial cognitive disturbances in susceptible individuals. Other common metabolic etiologies include liver and renal failure, hypercarbia and hypoxemia, vitamin deficiencies of thiamine and B₁₂, autoimmune disorders including central nervous system (CNS) vasculitis, and endocrinopathies such as thyroid and adrenal disorders.

Systemic infections often cause delirium, especially in the elderly. A common scenario involves the development of an acute cognitive decline in the setting of a urinary tract infection in a patient with baseline dementia. Pneumonia, skin infections such as cellulitis, and frank sepsis also lead to delirium. This so-called septic encephalopathy, often seen in the ICU, is probably due to the release of proinflammatory cytokines and their diffuse cerebral effects. CNS infections such as meningitis, encephalitis, and abscess are less common etiologies of delirium; however, in light of the high mortality rates associated with these conditions when they are not treated quickly, clinicians must always maintain a high index of suspicion.

In some susceptible individuals, exposure to the unfamiliar environment of a hospital itself can lead to delirium. This etiology usually occurs as part of a multifactorial delirium and should be considered a diagnosis of exclusion after all other causes have been thoroughly investigated. Many primary prevention and treatment strategies for delirium involve relatively simple methods to address the aspects of the inpatient setting that are most confusing.

Cerebrovascular etiologies of delirium are usually due to global hypoperfusion in the setting of systemic hypotension from heart failure, septic shock, dehydration, or anemia. Focal strokes in the right parietal lobe and medial dorsal thalamus rarely can lead to a delirious state. A more common scenario involves a new focal stroke or hemorrhage causing confusion in a patient who has decreased cerebral reserve. In these individuals, it is sometimes difficult to distinguish between cognitive dysfunction resulting from the new neurovascular insult itself and delirium due to the infectious, metabolic, and pharmacologic complications that can accompany hospitalization after stroke.

Because a fluctuating course often is seen in delirium, intermittent seizures may be overlooked when one is considering potential etiologies. Both nonconvulsive status epilepticus and recurrent focal or generalized seizures followed by postictal confusion can cause delirium; EEG remains essential for this diagnosis. Seizure activity spreading from an electrical focus in a mass or infarct can explain global cognitive dysfunction caused by relatively small lesions.

It is very common for patients to experience delirium at the end of life in palliative care settings. This condition, sometimes described as *terminal restlessness*, must be identified and treated aggressively because it is an important cause of patient and family discomfort at the end of life. It should be remembered that these patients also may be suffering from more common etiologies of delirium such as systemic infection.

LABORATORY AND DIAGNOSTIC EVALUATION

A cost-effective approach to the diagnostic evaluation of delirium allows the history and physical examination to guide further tests. No established algorithm for workup will fit all delirious patients due to the staggering number of potential etiologies, but one stepwise approach is detailed in [Table 34-3](#). If a clear precipitant is

TABLE 34-3 STEPWISE EVALUATION OF A PATIENT WITH DELIRIUM

Initial evaluation
History with special attention to medications (including over-the-counter and herbals)
General physical examination and neurologic examination
Complete blood count
Electrolyte panel including calcium, magnesium, phosphorus
Liver function tests, including albumin
Renal function tests
First-tier further evaluation guided by initial evaluation
Systemic infection screen
Urinalysis and culture
Chest radiograph
Blood cultures
Electrocardiogram
Arterial blood gas
Serum and/or urine toxicology screen (perform earlier in young persons)
Brain imaging with MRI with diffusion and gadolinium (preferred) or CT
Suspected CNS infection: lumbar puncture after brain imaging
Suspected seizure-related etiology: electroencephalogram (EEG) (if high suspicion, should be performed immediately)
Second-tier further evaluation
Vitamin levels: B ₁₂ , folate, thiamine
Endocrinologic laboratories: thyroid-stimulating hormone (TSH) and free T ₄ ; cortisol
Serum ammonia
Sedimentation rate
Autoimmune serologies: antinuclear antibodies (ANA), complement levels; p-ANCA, c-ANCA. consider paraneoplastic serologies
Infectious serologies: rapid plasmin reagin (RPR); fungal and viral serologies if high suspicion; HIV antibody
Lumbar puncture (if not already performed)
Brain MRI with and without gadolinium (if not already performed)

Abbreviations: c-ANCA, cytoplasmic antineutrophil cytoplasmic antibody; CNS, central nervous system; CT, computed tomography; MRI, magnetic resonance imaging; p-ANCA, perinuclear antineutrophil cytoplasmic antibody.

identified, such as an offending medication, further testing may not be required. If, however, no likely etiology is uncovered with initial evaluation, an aggressive search for an underlying cause should be initiated.

Basic screening labs, including a complete blood count, electrolyte panel, and tests of liver and renal function, should be obtained in all patients with delirium. In elderly patients, screening for systemic infection, including chest radiography, urinalysis and culture, and possibly blood cultures, is important. In younger individuals, serum and urine drug and toxicology screening may be appropriate early in the workup. Additional laboratory tests addressing other autoimmune, endocrinologic, metabolic, and infectious etiologies should be reserved for patients in whom the diagnosis remains unclear after initial testing.

Multiple studies have demonstrated that brain imaging in patients with delirium is often unhelpful. If, however, the initial workup is unrevealing, most clinicians quickly move toward imaging of the brain to exclude structural causes. A noncontrast computed tomography (CT) scan can identify large masses and hemorrhages but is otherwise unlikely to help determine an etiology of delirium. The ability of magnetic resonance imaging (MRI) to identify most acute ischemic strokes as well as to provide neuroanatomic detail that gives clues to possible infectious, inflammatory, neurodegenerative, and neoplastic conditions makes it the test of choice. Because MRI techniques are limited by availability, speed of imaging, patient cooperation, and contraindications, many clinicians begin with CT scanning and proceed to MRI if the etiology of delirium remains elusive.

Lumbar puncture (LP) must be obtained immediately after appropriate neuroimaging in all patients in whom CNS infection is suspected. Spinal fluid examination can also be useful in identifying inflammatory and neoplastic conditions. As a result, LP should be considered in any delirious patient with a negative workup. EEG does not have a routine role in the workup of delirium, but it remains invaluable if seizure-related etiologies are considered.

TREATMENT DELIRIUM

Management of delirium begins with treatment of the underlying inciting factor (e.g., patients with systemic infections should be given appropriate antibiotics, and underlying electrolyte disturbances judiciously corrected). These treatments often lead to prompt resolution of delirium. Blindly targeting the symptoms of delirium pharmacologically only serves to prolong the time patients remain in the confused state and may mask important diagnostic information.

Relatively simple methods of supportive care can be highly effective in treating patients with delirium. Reorientation by the nursing staff and family combined with visible clocks, calendars, and outside-facing windows can reduce confusion. Sensory isolation should be prevented by providing glasses and hearing aids to patients who need them. Sundowning can be addressed to a large extent through vigilance to appropriate sleep-wake cycles. During the day, a well-lit room should be accompanied by activities or exercises to prevent napping. At night, a quiet, dark environment with limited interruptions by staff can assure proper rest. These sleep-wake cycle interventions are especially important in the ICU setting as the usual constant 24-h activity commonly provokes delirium. Attempting to mimic the home environment as much as possible also has been shown to help treat and even prevent delirium. Visits from friends and family throughout the day minimize the anxiety associated with the constant flow of new faces of staff and physicians. Allowing hospitalized patients to have access to home bedding, clothing, and nightstand objects makes the hospital environment less foreign and therefore less confusing. Simple standard nursing practices such as maintaining proper nutrition and volume status as well as managing incontinence and skin breakdown also help alleviate discomfort and resulting confusion.

In some instances, patients pose a threat to their own safety or to the safety of staff members, and acute management is required. Bed alarms and personal sitters are more effective and much less disorienting than physical restraints. Chemical restraints should be avoided, but only when necessary, very-low-dose typical or atypical antipsychotic medications administered on an as-needed basis are effective. The recent association of antipsychotic use in the elderly with increased mortality rates underscores the importance of using these medications judiciously and only as a last resort. Benzodiazepines often worsen confusion through their sedative properties. Although many clinicians still use benzodiazepines to treat acute confusion, their use should be limited to cases in which delirium is caused by alcohol or benzodiazepine withdrawal.

PREVENTION

In light of the high morbidity associated with delirium and the tremendously increased health care costs that accompany it, development of an effective strategy to prevent delirium in hospitalized patients is extremely important. Successful identification of high-risk patients is the first step, followed by initiation of appropriate interventions. Simple standardized protocols used to manage risk factors for delirium, including sleep-wake cycle reversal, immobility, visual impairment, hearing impairment, sleep deprivation, and dehydration, have been shown to be effective. Recent trials in the ICU have focused both on identifying sedatives, such as dexmedetomidine, that are less likely to lead to delirium in critically ill patients and on developing protocols for daily awakenings in which infusions of sedative medications are interrupted and the patient is reoriented by the staff. All hospitals and health care systems should work toward decreasing the incidence of delirium.

35

Dementia

William W. Seeley, Bruce L. Miller

Dementia, a syndrome with many causes, affects >5 million people in the United States and results in a total annual health care cost between \$157 and \$215 billion. Dementia is defined as an acquired deterioration in cognitive abilities that impairs the successful performance of activities of daily living. Episodic memory, the ability to recall events specific in time and place, is the cognitive function most commonly lost; 10% of persons age >70 years and 20–40% of individuals age >85 years have clinically identifiable memory loss. In addition to memory, dementia may erode other mental faculties, including language, visuospatial, praxis, calculation, judgment, and problem-solving abilities. Neuropsychiatric and social deficits also arise in many dementia syndromes, manifesting as depression, apathy, anxiety, hallucinations, delusions, agitation, insomnia, sleep disturbances, compulsions, or disinhibition. The clinical course may be slowly progressive, as in Alzheimer's disease (AD); static, as in anoxic encephalopathy; or may fluctuate from day to day or minute to minute, as in dementia with Lewy bodies. Most patients with AD, the most prevalent form of dementia, begin with episodic memory impairment, although in other dementias, such as frontotemporal dementia, memory loss is not typically a presenting feature. **Focal cerebral disorders are discussed in Chap. 36 and illustrated in a video library in Chap. 37e; the pathogenesis of AD and related disorders is discussed in Chap. 448.**

FUNCTIONAL ANATOMY OF THE DEMENTIAS

Dementia syndromes result from the disruption of specific large-scale neuronal networks; the location and severity of synaptic and neuronal loss combine to produce the clinical features (**Chap. 36**). Behavior, mood, and attention are modulated by ascending noradrenergic, serotonergic, and dopaminergic pathways, whereas cholinergic signaling is critical for attention and memory functions. The dementias differ in the relative neurotransmitter deficit profiles; accordingly, accurate diagnosis guides effective pharmacologic therapy.

AD begins in the entorhinal region of the medial temporal lobe, spreads to the hippocampus, and then moves to lateral and posterior temporal and parietal neocortex, eventually causing a more widespread degeneration. *Vascular dementia* is associated with focal damage in a variable patchwork of cortical and subcortical regions or white matter tracts that disconnect nodes within distributed networks. In keeping with its anatomy, AD typically presents with episodic memory loss accompanied later by aphasia or navigational problems. In contrast, dementias that begin in frontal or subcortical regions, such as *frontotemporal dementia* (FTD) or *Huntington's disease* (HD), are less likely to begin with memory problems and more likely to present with difficulties with judgment, mood, executive control, movement, and behavior.

Lesions of frontal-striatal¹ pathways produce specific and predictable effects on behavior. The dorsolateral prefrontal cortex has connections with a central band of the caudate nucleus. Lesions of either the caudate or dorsolateral prefrontal cortex, or their connecting white matter pathways, may result in executive dysfunction, manifesting as poor organization and planning, decreased cognitive flexibility, and impaired working memory. The lateral orbital frontal cortex connects with the ventromedial caudate, and lesions of this system cause impulsiveness, distractibility, and disinhibition. The anterior cingulate cortex and adjacent medial prefrontal cortex project to the nucleus accumbens, and interruption of this system produces apathy, poverty of speech, emotional blunting, or even akinetic mutism. All cortico-striatal systems also include topographically organized projections through the globus pallidus and thalamus, and damage to these nodes

¹The striatum comprises the caudate/putamen.

can likewise reproduce the clinical syndrome of cortical or striatal injury.

THE CAUSES OF DEMENTIA

The single strongest risk factor for dementia is increasing age. The prevalence of disabling memory loss increases with each decade over age 50 and is usually associated with the microscopic changes of AD at autopsy. Yet some centenarians have intact memory function and no evidence of clinically significant dementia. Whether dementia is an inevitable consequence of normal human aging remains controversial.

The many causes of dementia are listed in [Table 35-1](#). The frequency of each condition depends on the age group under study, access of the group to medical care, country of origin, and perhaps racial or ethnic background. AD is the most common cause of dementia in Western

countries, accounting for more than half of all patients. Vascular disease is considered the second most frequent cause for dementia and is particularly common in elderly patients or populations with limited access to medical care, where vascular risk factors are undertreated. Often, vascular brain injury is mixed with neurodegenerative disorders, making it difficult, even for the neuropathologist, to estimate the contribution of cerebrovascular disease to the cognitive disorder in an individual patient. Dementias associated with Parkinson's disease (PD) ([Chap. 449](#)) are common and may develop years after onset of a parkinsonian disorder, as seen with PD-related dementia (PDD), or can occur concurrently with or preceding the motor syndrome, as in dementia with Lewy bodies (DLB). In patients under the age of 65, FTD rivals AD as the most common cause of dementia. Chronic intoxications, including those resulting from alcohol and prescription drugs, are an important and often treatable cause of dementia. Other disorders listed in [Table 35-1](#) are uncommon but important because many are reversible. The classification of dementing illnesses into reversible and irreversible disorders is a useful approach to differential diagnosis. When effective treatments for the neurodegenerative conditions emerge, this dichotomy will become obsolete.

In a study of 1000 persons attending a memory disorders clinic, 19% had a potentially reversible cause of the cognitive impairment and 23% had a potentially reversible concomitant condition that may have contributed to the patient's impairment. The three most common potentially reversible diagnoses were depression, normal pressure hydrocephalus (NPH), and alcohol dependence; medication side effects are also common and should be considered in every patient ([Table 35-1](#)).

Subtle cumulative decline in episodic memory is a common part of aging. This frustrating experience, often the source of jokes and humor, is referred to as *benign forgetfulness of the elderly*. *Benign* means that it is not so progressive or serious that it impairs reasonably successful and productive daily functioning, although the distinction between benign and more significant memory loss can be difficult to make. At age 85, the average person is able to learn and recall approximately one-half of the items (e.g., words on a list) that he or she could at age 18. A measurable cognitive problem that does not seriously disrupt daily activities is often referred to as *mild cognitive impairment* (MCI). Factors that predict progression from MCI to an AD dementia include a prominent memory deficit, family history of dementia, presence of an apolipoprotein $\epsilon 4$ (Apo $\epsilon 4$) allele, small hippocampal volumes, an AD-like signature of cortical atrophy, low cerebrospinal fluid A β , and elevated tau or evidence of brain amyloid deposition on positron emission tomography (PET) imaging.

The major degenerative dementias include AD, DLB, FTD and related disorders, HD, and prion diseases, including Creutzfeldt-Jakob disease (CJD). These disorders are all associated with the abnormal aggregation of a specific protein: A β_{42} and tau in AD; α -synuclein in DLB; tau, TAR DNA-binding protein of 43 kDa (TDP-43), or fused in sarcoma (FUS) in FTD; huntingtin in HD; and misfolded prion protein (PrP^{sc}) in CJD ([Table 35-2](#)).

TABLE 35-1 DIFFERENTIAL DIAGNOSIS OF DEMENTIA

Most Common Causes of Dementia

Alzheimer's disease	Alcoholism ^a
Vascular dementia	PDD/LBD spectrum
Multi-infarct	Drug/medication intoxication ^a
Diffuse white matter disease (Binswanger's)	

Less Common Causes of Dementia

Vitamin deficiencies	Toxic disorders
Thiamine (B ₁): Wernicke's encephalopathy ^a	Drug, medication, and narcotic poisoning ^a
B ₁₂ (subacute combined degeneration) ^a	Heavy metal intoxication ^a
Nicotinic acid (pellagra) ^a	Organic toxins
Endocrine and other organ failure	Psychiatric
Hypothyroidism ^a	Depression (pseudodementia) ^a
Adrenal insufficiency and Cushing's syndrome ^a	Schizophrenia ^a
Hypo- and hyperparathyroidism ^a	Conversion disorder ^a
Renal failure ^a	Degenerative disorders
Liver failure ^a	Huntington's disease
Pulmonary failure ^a	Multisystem atrophy
Chronic infections	Hereditary ataxias (some forms)
HIV	Frontotemporal lobar degeneration spectrum
Neurosyphilis ^a	Multiple sclerosis
Papovavirus (JC virus) (progressive multifocal leukoencephalopathy)	Adult Down's syndrome with Alzheimer's disease
Tuberculosis, fungal, and protozoal ^a	ALS-parkinsonism-dementia complex of Guam
Whipple's disease ^a	Prion (Creutzfeldt-Jakob and Gerstmann-Sträussler-Scheinker diseases)
Head trauma and diffuse brain damage	Miscellaneous
Chronic traumatic encephalopathy	Sarcoidosis ^a
Chronic subdural hematoma ^a	Vasculitis ^a
Postanoxia	CADASIL, etc.
Postencephalitis	Acute intermittent porphyria ^a
Normal-pressure hydrocephalus ^a	Recurrent nonconvulsive seizures ^a
Intracranial hypotension	Additional conditions in children or adolescents
Neoplastic	Pantothenate kinase-associated neurodegeneration
Primary brain tumor ^a	Subacute sclerosing panencephalitis
Metastatic brain tumor ^a	Metabolic disorders (e.g., Wilson's and Leigh's diseases, leukodystrophies, lipid storage diseases, mitochondrial mutations)
Paraneoplastic/autoimmune limbic encephalitis ^a	

^aPotentially reversible dementia.

Abbreviations: ALS, amyotrophic lateral sclerosis; CADASIL, cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; LBD, Lewy body disease; PDD, Parkinson's disease dementia.

APPROACH TO THE PATIENT: Dementias

Three major issues should be kept at the forefront: (1) What is the best fit for a clinical diagnosis? (2) What component of the dementia syndrome is treatable or reversible? (3) Can the physician help to alleviate the burden on caregivers? A broad overview of the approach to dementia is shown in [Table 35-3](#). The major degenerative dementias can usually be distinguished by the initial symptoms; neuropsychological, neuropsychiatric, and neurologic findings; and neuroimaging features ([Table 35-4](#)).

HISTORY

The history should concentrate on the onset, duration, and tempo of progression. An acute or subacute onset of confusion may be due to delirium ([Chap. 34](#)) and should trigger the search for intoxication, infection, or metabolic derangement. An elderly person

TABLE 35-2 THE MOLECULAR BASIS FOR DEGENERATIVE DEMENTIA

Dementia	Molecular Basis	Causal Genes (Chromosome)	Susceptibility Genes	Pathologic Findings
AD	A β /tau	<i>APP</i> (21), <i>PS-1</i> (14), <i>PS-2</i> (1) (<2% carry these mutations, most often in <i>PS-1</i>)	<i>Apo E4</i> (19)	Amyloid plaques, neurofibrillary tangles, and neuropil threads
FTD	Tau	<i>MAPT</i> exon and intron mutations (17) (about 10% of familial cases)	H1 <i>MAPT</i> haplotype	Tau neuronal and glial inclusions varying in morphology and distribution
	TDP-43	<i>GRN</i> (10% of familial cases), <i>C9ORF72</i> (20–30% of familial cases), rare <i>VCP</i> , very rare <i>TARDBP</i>		TDP-43 neuronal and glial inclusions varying in morphology and distribution
	FUS	Very rare <i>FUS</i>		FUS neuronal and glial inclusions varying in morphology and distribution
DLB	α -Synuclein	Very rare <i>SNCA</i> (4)	Unknown	α -Synuclein neuronal inclusions (Lewy bodies)
CJD	PrP ^{Sc}	<i>PRNP</i> (20) (up to 15% of patients carry these dominant mutations)	Codon 129 homozygosity for methionine or valine	PrP ^{Sc} deposition, panlaminar spongiosis

Abbreviations: AD, Alzheimer's disease; CJD, Creutzfeldt-Jakob disease; DLB, dementia with Lewy bodies; FTD, frontotemporal dementia.

with slowly progressive memory loss over several years is likely to suffer from AD. Nearly 75% of patients with AD begin with memory symptoms, but other early symptoms include difficulty with managing money, driving, shopping, following instructions, finding words, or navigating. Personality change, disinhibition, and weight gain or compulsive eating suggest FTD, not AD. FTD is also suggested by prominent apathy, compulsivity, loss of empathy for others, or progressive loss of speech fluency or single-word comprehension and by a relative sparing of memory and visuo-spatial abilities. The diagnosis of DLB is suggested by early visual hallucinations; parkinsonism; proneness to delirium or sensitivity to psychoactive medications; rapid eye movement (REM) behavior disorder (RBD; the loss of skeletal muscle paralysis during dreaming); or Capgras syndrome, the delusion that a familiar person has been replaced by an impostor.

A history of stroke with irregular stepwise progression suggests vascular dementia. Vascular dementia is also commonly seen in the setting of hypertension, atrial fibrillation, peripheral vascular disease, and diabetes. In patients suffering from cerebrovascular disease, it can be difficult to determine whether the dementia is due to AD, vascular disease, or a mixture of the two because many of the risk factors for vascular dementia, including diabetes, high cholesterol, elevated homocysteine, and low exercise, are also putative risk factors for AD. Moreover, many patients with a major vascular contribution to their dementia lack a history of stepwise decline. Rapid progression with motor rigidity and myoclonus suggests CJD (**Chap. 453e**). Seizures may indicate strokes or neoplasm but also occur in AD, particularly early-age-of-onset AD. Gait disturbance is common in vascular dementia, PD/DLB, or NPH. A history of high-risk sexual behaviors or intravenous drug use

TABLE 35-3 EVALUATION OF THE PATIENT WITH DEMENTIA

Routine Evaluation	Optional Focused Tests	Occasionally Helpful Tests
History	Psychometric testing	EEG
Physical examination	Chest x-ray	Parathyroid function
Laboratory tests	Lumbar puncture	Adrenal function
Thyroid function (TSH)	Liver function	Urine heavy metals
Vitamin B ₁₂	Renal function	RBC sedimentation rate
Complete blood count	Urine toxin screen	Angiogram
Electrolytes	HIV	Brain biopsy
CT/MRI	Apolipoprotein E	SPECT
	RPR or VDRL	PET
		Lab screen for autoantibodies
Diagnostic Categories		
Reversible Causes	Irreversible/Degenerative Dementias	Psychiatric Disorders
Examples	Examples	Depression
Hypothyroidism	Alzheimer's	Schizophrenia
Thiamine deficiency	Frontotemporal dementia	Conversion reaction
Vitamin B ₁₂ deficiency	Huntington's	
Normal-pressure hydrocephalus	Dementia with Lewy bodies	
Subdural hematoma	Vascular	
Chronic infection	Leukoencephalopathies	
Brain tumor	Parkinson's	
Drug intoxication		
Autoimmune encephalopathy		
Associated Treatable Conditions		
	Depression	Agitation
	Seizures	Caregiver "burnout"
	Insomnia	Drug side effects

Abbreviations: CT, computed tomography; EEG, electroencephalogram; MRI, magnetic resonance imaging; PET, positron emission tomography; RBC, red blood cell; RPR, rapid plasma reagin (test); SPECT, single-photon emission computed tomography; TSH, thyroid-stimulating hormone; VDRL, Venereal Disease Research Laboratory (test for syphilis).

TABLE 35-4 CLINICAL DIFFERENTIATION OF THE MAJOR DEMENTIAS

Disease	First Symptom	Mental Status	Neuropsychiatry	Neurology	Imaging
AD	Memory loss	Episodic memory loss	Irritability, anxiety, depression	Initially normal	Entorhinal cortex and hippocampal atrophy
FTD	Apathy; poor judgment/insight, speech/language; hyperorality	Frontal/executive and/or language; spares drawing	Apathy, disinhibition, overeating, compulsivity	May have vertical gaze palsy, axial rigidity, dystonia, alien hand, or MND	Frontal, insular, and/or temporal atrophy; usually spares posterior parietal lobe
DLB	Visual hallucinations, REM sleep behavior disorder, delirium, Capgras syndrome, parkinsonism	Drawing and frontal/executive; spares memory; delirium-prone	Visual hallucinations, depression, sleep disorder, delusions	Parkinsonism	Posterior parietal atrophy; hippocampi larger than in AD
CJD	Dementia, mood, anxiety, movement disorders	Variable, frontal/executive, focal cortical, memory	Depression, anxiety, psychosis in some	Myoclonus, rigidity, parkinsonism	Cortical ribboning and basal ganglia or thalamus hyperintensity on diffusion/FLAIR MRI
Vascular	Often but not always sudden; variable; apathy, falls, focal weakness	Frontal/executive, cognitive slowing; can spare memory	Apathy, delusions, anxiety	Usually motor slowing, spasticity; can be normal	Cortical and/or subcortical infarctions, confluent white matter disease

Abbreviations: AD, Alzheimer's disease; CBD, cortical basal degeneration; CJD, Creutzfeldt-Jakob disease; DLB, dementia with Lewy bodies; FLAIR, fluid-attenuated inversion recovery; FTD, frontotemporal dementia; MND, motor neuron disease; MRI, magnetic resonance imaging; PSP, progressive supranuclear palsy; REM, rapid eye movement.

should trigger a search for central nervous system (CNS) infection, especially HIV or syphilis. A history of recurrent head trauma could indicate chronic subdural hematoma, chronic traumatic encephalopathy (a progressive dementia best characterized in contact sport athletes such as boxers and American football players), intracranial hypotension, or NPH. Subacute onset of severe amnesia and psychosis with mesial temporal T2/fluid-attenuated inversion recovery (FLAIR) hyperintensities on magnetic resonance imaging (MRI) should raise concern for paraneoplastic limbic encephalitis, especially in a long-term smoker or other patients at risk for cancer. Related autoimmune conditions, such as voltage-gated potassium channel (VGKC)- or N-methyl-D-aspartate (NMDA)-receptor antibody-mediated encephalopathy, can present with a similar tempo and imaging signature with or without characteristic motor manifestations such as myokymia (anti-VGKC) and faciobrachial dystonic seizures (anti-NMDA). Alcohol abuse creates risk for malnutrition and thiamine deficiency. Veganism, bowel irradiation, an autoimmune diathesis, a remote history of gastric surgery, and chronic antihistamine therapy for dyspepsia or gastroesophageal reflux predispose to B₁₂ deficiency. Certain occupations, such as working in a battery or chemical factory, might indicate heavy metal intoxication. Careful review of medication intake, especially for sedatives and analgesics, may raise the issue of chronic drug intoxication. An autosomal dominant family history is found in HD and in familial forms of AD, FTD, DLB, or prion disorders. A history of mood disorders, the recent death of a loved one, or depressive signs, such as insomnia or weight loss, raise the possibility of depression-related cognitive impairments.

PHYSICAL AND NEUROLOGIC EXAMINATION

A thorough general and neurologic examination is essential to document dementia, to look for other signs of nervous system involvement, and to search for clues suggesting a systemic disease that might be responsible for the cognitive disorder. Typical AD spares motor systems until later in the course. In contrast, FTD patients often develop axial rigidity, supranuclear gaze palsy, or a motor neuron disease reminiscent of amyotrophic lateral sclerosis (ALS). In DLB, the initial symptoms may include the new onset of a parkinsonian syndrome (resting tremor, cogwheel rigidity, bradykinesia, festinating gait), but DLB often starts with visual hallucinations or dementia. Symptoms referable to the lower brainstem (RBD, gastrointestinal or autonomic problems) may arise years or even decades before parkinsonism or dementia. Corticobasal syndrome (CBS) features asymmetric akinesia and rigidity, dystonia, myoclonus, alien limb phenomena, pyramidal signs, and prefrontal deficits such as nonfluent aphasia with or without motor speech

impairment, executive dysfunction, apraxia, or a behavioral disorder. Progressive supranuclear palsy (PSP) is associated with unexplained falls, axial rigidity, dysphagia, and vertical gaze deficits. CJD is suggested by the presence of diffuse rigidity, an akinetic-mute state, and prominent, often startle-sensitive myoclonus.

Hemiparesis or other focal neurologic deficits suggest vascular dementia or brain tumor. Dementia with a myelopathy and peripheral neuropathy suggests vitamin B₁₂ deficiency. Peripheral neuropathy could also indicate another vitamin deficiency, heavy metal intoxication, thyroid dysfunction, Lyme disease, or vasculitis. Dry, cool skin, hair loss, and bradycardia suggest hypothyroidism. Fluctuating confusion associated with repetitive stereotyped movements may indicate ongoing limbic, temporal, or frontal seizures. In the elderly, hearing impairment or visual loss may produce confusion and disorientation misinterpreted as dementia. Profound bilateral sensorineural hearing loss in a younger patient with short stature or myopathy, however, should raise concern for a mitochondrial disorder.

COGNITIVE AND NEUROPSYCHIATRIC EXAMINATION

Brief screening tools such as the Mini-Mental State Examination (MMSE), the Montreal Cognitive Assessment (MOCA), and Cognistat can be used to capture dementia and follow progression. None of these tests is highly sensitive to early-stage dementia or discriminates between dementia syndromes. The MMSE is a 30 point test of cognitive function, with each correct answer being scored as 1 point. It includes tests in the areas of: orientation (e.g., identify season/date/month/year/floor/hospital/town/state/country); registration (e.g., name and restate 3 objects); recall (e.g., remember the same three objects 5 minutes later); and language (e.g., name pencil and watch; repeat "no if's and's or but's"; follow a 3-step command; obey a written command; and write a sentence and copy a design). In most patients with MCI and some with clinically apparent AD, bedside screening tests may be normal, and a more challenging and comprehensive set of neuropsychological tests will be required. When the etiology for the dementia syndrome remains in doubt, a specially tailored evaluation should be performed that includes tasks of working and episodic memory, executive function, language, and visuospatial and perceptual abilities. In AD, the early deficits involve episodic memory, category generation ("name as many animals as you can in 1 minute"), and visuoconstructive ability. Usually deficits in verbal or visual episodic memory are the first neuropsychological abnormalities detected, and tasks that require the patient to recall a long list of words or a series of pictures after a predetermined delay will demonstrate deficits in most patients. In FTD, the earliest

deficits on cognitive testing involve executive control or language (speech or naming) function, but some patients lack either finding despite profound social-emotional deficits. PDD or DLB patients have more severe deficits in visuospatial function but do better on episodic memory tasks than patients with AD. Patients with vascular dementia often demonstrate a mixture of executive control and visuospatial deficits, with prominent psychomotor slowing. In delirium, the most prominent deficits involve attention, working memory, and executive function, making the assessment of other cognitive domains challenging and often uninformative.

A functional assessment should also be performed to help the physician determine the day-to-day impact of the disorder on the patient's memory, community affairs, hobbies, judgment, dressing, and eating. Knowledge of the patient's functional abilities will help the clinician and the family to organize a therapeutic approach.

Neuropsychiatric assessment is important for diagnosis, prognosis, and treatment. In the early stages of AD, mild depressive features, social withdrawal, and irritability or anxiety are the most prominent psychiatric changes, but patients often maintain core social graces into the middle or late stages, when delusions, agitation, and sleep disturbance may emerge. In FTD, dramatic personality change with apathy, overeating, compulsions, disinhibition, euphoria, and loss of empathy are early and common. DLB is associated with visual hallucinations, delusions related to person or place identity, RBD, and excessive daytime sleepiness. Dramatic fluctuations occur not only in cognition but also in arousal. Vascular dementia can present with psychiatric symptoms such as depression, anxiety, delusions, disinhibition, or apathy.

LABORATORY TESTS

The choice of laboratory tests in the evaluation of dementia is complex and should be tailored to the individual patient. The physician must take measures to avoid missing a reversible or treatable cause, yet no single treatable etiology is common; thus, a screen must use multiple tests, each of which has a low yield. Cost/benefit ratios are difficult to assess, and many laboratory screening algorithms for dementia discourage multiple tests. Nevertheless, even a test with only a 1–2% positive rate is worth undertaking if the alternative is missing a treatable cause of dementia. Table 35-3 lists most screening tests for dementia. The American Academy of Neurology recommends the routine measurement of a complete blood count, electrolytes, renal and thyroid function, a vitamin B₁₂ level, and a neuroimaging study (computed tomography [CT] or MRI).

Neuroimaging studies, especially MRI, help to rule out primary and metastatic neoplasms, locate areas of infarction or inflammation, detect subdural hematomas, and suggest NPH or diffuse white matter disease. They also help to establish a regional pattern of atrophy. Support for the diagnosis of AD includes hippocampal atrophy in addition to posterior-predominant cortical atrophy (Fig. 35-1). Focal frontal, insular, and/or anterior temporal atrophy suggests FTD (Chap. 448). DLB often features less prominent atrophy, with greater involvement of amygdala than hippocampus. In CJD, magnetic resonance (MR) diffusion-weighted imaging reveals restricted diffusion within the cortical ribbon and basal ganglia in most patients. Extensive white matter abnormalities correlate with a vascular etiology (Fig. 35-2). Communicating hydrocephalus with vertex effacement (crowding of dorsal convexity gyri/sulci), gapping Sylvian fissures despite minimal cortical atrophy, and additional features shown in Fig. 35-3 suggest NPH. Single-photon emission computed tomography (SPECT) and PET scanning show temporal-parietal hypoperfusion or hypometabolism in AD and frontotemporal deficits in FTD, but these changes often reflect atrophy and can therefore be detected with MRI alone in many patients. Recently, amyloid imaging has shown promise for the diagnosis of AD, and Pittsburgh Compound-B (PiB) (not available outside of research settings) and ¹⁸F-AV-45 (florbetapir; approved by the U.S. Food and Drug Administration in 2013) are reliable radioligands for detecting brain amyloid associated with amyloid

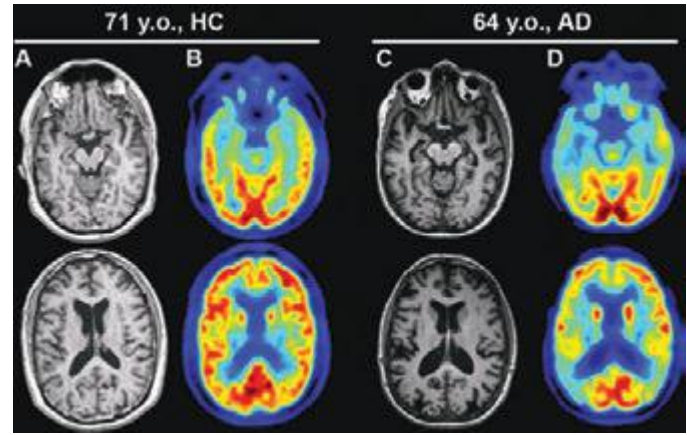


FIGURE 35-1 Alzheimer's disease (AD). Axial T1-weighted magnetic resonance images of a healthy 71-year-old (A) and a 64-year-old with AD (C). Note the reduction in medial temporal lobe volume in the patient with AD. Fluorodeoxyglucose positron emission tomography scans of the same individuals (B and D) demonstrate reduced glucose metabolism in the posterior temporoparietal regions bilaterally in AD, a typical finding in this condition. HC, healthy control. (Images courtesy of Gil Rabinovici, University of California, San Francisco and William Jagust, University of California, Berkeley.)

angiopathy or neuritic plaques of AD (Fig. 35-4). Because these abnormalities can be seen in cognitively normal older persons, however (~25% of individuals at age 65), amyloid imaging may also detect preclinical or incidental AD in patients lacking an AD-like dementia syndrome. Currently, the main clinical value of amyloid imaging is to exclude AD as the likely cause of dementia in patients who have negative scans. Once disease-modifying therapies become available, use of these biomarkers may help to identify treatment



FIGURE 35-2 Diffuse white matter disease. Axial fluid-attenuated inversion recovery (FLAIR) magnetic resonance image through the lateral ventricles reveals multiple areas of hyperintensity (arrows) involving the periventricular white matter as well as the corona radiata and striatum. Although seen in some individuals with normal cognition, this appearance is more pronounced in patients with dementia of a vascular etiology.

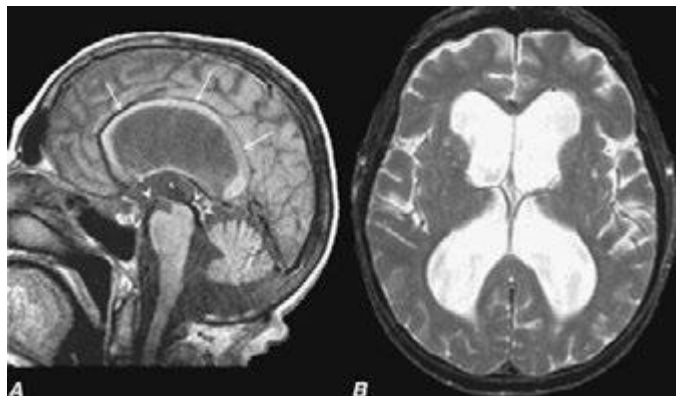


FIGURE 35-3 Normal-pressure hydrocephalus. **A.** Sagittal T1-weighted magnetic resonance image (MRI) demonstrates dilation of the lateral ventricle and stretching of the corpus callosum (arrows), depression of the floor of the third ventricle (single arrowhead), and enlargement of the aqueduct (double arrowheads). Note the diffuse dilation of the lateral, third, and fourth ventricles with a patent aqueduct, typical of communicating hydrocephalus. **B.** Axial T2-weighted MRIs demonstrate dilation of the lateral ventricles. This patient underwent successful ventriculoperitoneal shunting.

candidates before irreversible brain injury has occurred. In the meantime, the significance of detecting brain amyloid in an asymptomatic elder remains a topic of vigorous investigation. Similarly, MRI perfusion and structural/functional connectivity methods are being explored as potential treatment-monitoring strategies.

Lumbar puncture need not be done routinely in the evaluation of dementia, but it is indicated when CNS infection or inflammation are credible diagnostic possibilities. Cerebrospinal fluid (CSF) levels of $A\beta_{42}$ and tau proteins show differing patterns with the various dementias, and the presence of low $A\beta_{42}$ and mildly elevated CSF tau is highly suggestive of AD. The routine use of lumbar puncture in the diagnosis of dementia is debated, but the sensitivity and specificity of AD diagnostic measures are not yet high enough to warrant routine use. Formal psychometric testing helps to document the severity of cognitive disturbance, suggest psychogenic causes, and provide a more formal method for following the disease course. Electroencephalogram (EEG) is not routinely used but can help to suggest CJD (repetitive bursts of diffuse high-amplitude sharp waves, or “periodic complexes”) or an underlying nonconvulsive seizure disorder (epileptiform discharges). Brain biopsy (including meninges) is not advised except

to diagnose vasculitis, potentially treatable neoplasms, or unusual infections when the diagnosis is uncertain. Systemic disorders with CNS manifestations, such as sarcoidosis, can usually be confirmed through biopsy of lymph node or solid organ rather than brain. MR angiography should be considered when cerebral vasculitis or cerebral venous thrombosis is a possible cause of the dementia.

TREATMENT DEMENTIA

The major goals of dementia management are to treat reversible causes and to provide comfort and support to the patient and caregivers. Treatment of underlying causes includes thyroid replacement for hypothyroidism; vitamin therapy for thiamine or B_{12} deficiency or for elevated serum homocysteine; antimicrobials for opportunistic infections or antiretrovirals for HIV; ventricular shunting for NPH; or appropriate surgical, radiation, and/or chemotherapeutic treatment for CNS neoplasms. Removal of cognition-impairing drugs or medications is frequently useful. If the patient’s cognitive complaints stem from a psychiatric disorder, vigorous treatment of this condition should seek to eliminate the cognitive complaint or confirm that it persists despite adequate resolution of the mood or anxiety symptoms. Patients with degenerative diseases may also be depressed or anxious, and those aspects of their condition often respond to therapy. Antidepressants, such as selective serotonin reuptake inhibitors (SSRIs) or serotonin-norepinephrine reuptake inhibitors (SNRIs) (Chap. 465e), which feature anxiolytic properties but few cognitive side effects, provide the mainstay of treatment when necessary. Anticonvulsants are used to control seizures. Levetiracetam may be particularly useful, but there have as yet been no randomized trials for treatment of AD-associated seizures.

Agitation, hallucinations, delusions, and confusion are difficult to treat. These behavioral problems represent major causes for nursing home placement and institutionalization. Before treating these behaviors with medications, the clinician should aggressively seek out modifiable environmental or metabolic factors. Hunger, lack of exercise, toothache, constipation, urinary tract or respiratory infection, electrolyte imbalance, and drug toxicity all represent easily correctable causes that can be remedied without psychoactive drugs. Drugs such as phenothiazines and benzodiazepines may ameliorate the behavior problems but have untoward side effects such as sedation, rigidity, dyskinesia, and occasionally paradoxical disinhibition (benzodiazepines). Despite their unfavorable side effect profile, second-generation antipsychotics such as quetiapine (starting dose, 12.5–25 mg daily) can be used for patients with agitation, aggression, and

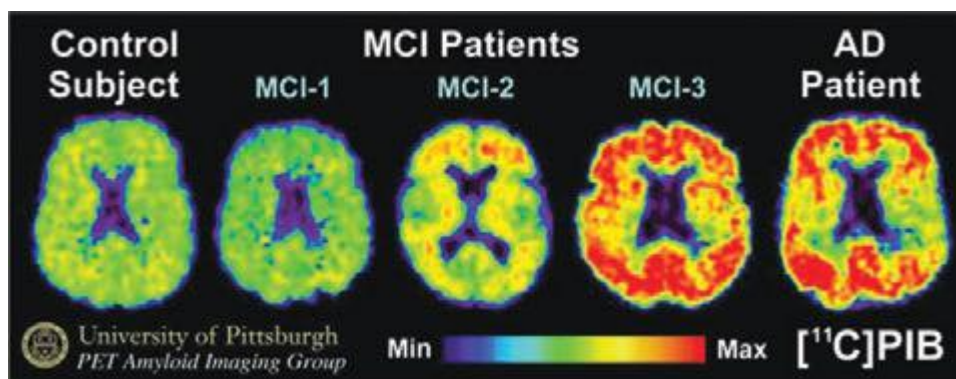


FIGURE 35-4 Positron emission tomography (PET) images obtained with the amyloid-imaging agent Pittsburgh Compound-B ($[^{11}C]PIB$) in a normal control (left); three different patients with mild cognitive impairment (MCI; center); and a patient with mild Alzheimer’s disease (AD; right). Some MCI patients have control-like levels of amyloid, some have AD-like levels of amyloid, and some have intermediate levels. (Images courtesy of William Klunk and Chester Mathis, University of Pittsburgh.)

psychosis, although the risk profile for these compounds is significant. When patients do not respond to treatment, it is usually a mistake to advance to higher doses or to use anticholinergic drugs or sedatives (such as barbiturates or benzodiazepines). It is important to recognize and treat depression; treatment can begin with a low dose of an SSRI (e.g., escitalopram, starting dose 5 mg daily, target dose 5–10 mg daily) while monitoring for efficacy and toxicity. Sometimes apathy, visual hallucinations, depression, and other psychiatric symptoms respond to the cholinesterase inhibitors, especially in DLB, obviating the need for other more toxic therapies.

Cholinesterase inhibitors are being used to treat AD (donepezil, rivastigmine, galantamine) and PDD (rivastigmine). Recent work has focused on developing antibodies against $A\beta_{42}$ as a treatment for AD. Although the initial randomized controlled trials failed, there was some evidence for efficacy in the mildest patient groups. Therefore, researchers have begun to focus on patients with very mild disease and asymptomatic individuals at risk for AD, such as those who carry autosomal dominantly inherited genetic mutations or healthy elders with CSF or amyloid imaging biomarker evidence supporting presymptomatic AD. Memantine proves useful when treating some patients with moderate to severe AD; its major benefit relates to decreasing caregiver burden, most likely by decreasing resistance to dressing and grooming support. In moderate to severe AD, the combination of memantine and a cholinesterase inhibitor delayed nursing home placement in several studies, although other studies have not supported the efficacy of adding memantine to the regimen.

A proactive strategy has been shown to reduce the occurrence of delirium in hospitalized patients. This strategy includes frequent orientation, cognitive activities, sleep-enhancement measures, vision and hearing aids, and correction of dehydration.

Nondrug behavior therapy has an important place in dementia management. The primary goals are to make the patient's life comfortable, uncomplicated, and safe. Preparing lists, schedules, calendars, and labels can be helpful in the early stages. It is also useful to stress familiar routines, walks, and simple physical exercises. For many demented patients, memory for events is worse than their ability to carry out routine activities, and they may still be able to take part in activities such as walking, bowling, dancing, singing, bingo, and golf. Demented patients often object to losing control over familiar tasks such as driving, cooking, and handling finances. Attempts to help or take over may be greeted with complaints, depression, or anger. Hostile responses on the part of the caregiver are counterproductive and sometimes even harmful. Reassurance, distraction, and calm positive statements are more productive in this setting. Eventually, tasks such as finances and driving must be assumed by others, and the patient will conform and adjust. Safety is an important issue that includes not only driving but controlling the kitchen, bathroom, and sleeping area environments, as well as stairways. These areas need to be monitored, supervised, and made as safe as possible. A move to a retirement complex, assisted-living center, or nursing home can initially increase confusion and agitation. Repeated reassurance, reorientation, and careful introduction to the new personnel will help to smooth the process. Providing activities that are known to be enjoyable to the patient can be of considerable benefit.

The clinician must pay special attention to frustration and depression among family members and caregivers. Caregiver guilt and burnout are common. Family members often feel overwhelmed and helpless and may vent their frustrations on the patient, each other, and health care providers. Caregivers should be encouraged to take advantage of day-care facilities and respite services. Education and counseling about dementia are important. Local and national support groups, such as the Alzheimer's Association (www.alz.org), can provide considerable help.

36

Aphasia, Memory Loss, and Other Focal Cerebral Disorders

M.-Marsel Mesulam

The cerebral cortex of the human brain contains approximately 20 billion neurons spread over an area of 2.5 m². The primary sensory and motor areas constitute 10% of the cerebral cortex. The rest is subsumed by modality-selective, heteromodal, paralimbic, and limbic areas collectively known as the *association cortex* (Fig. 36-1). The association cortex mediates the integrative processes that subserve cognition, emotion, and comportment. A systematic testing of these mental functions is necessary for the effective clinical assessment of the association cortex and its diseases. According to current thinking, there are no centers for “hearing words,” “perceiving space,” or “storing memories.”

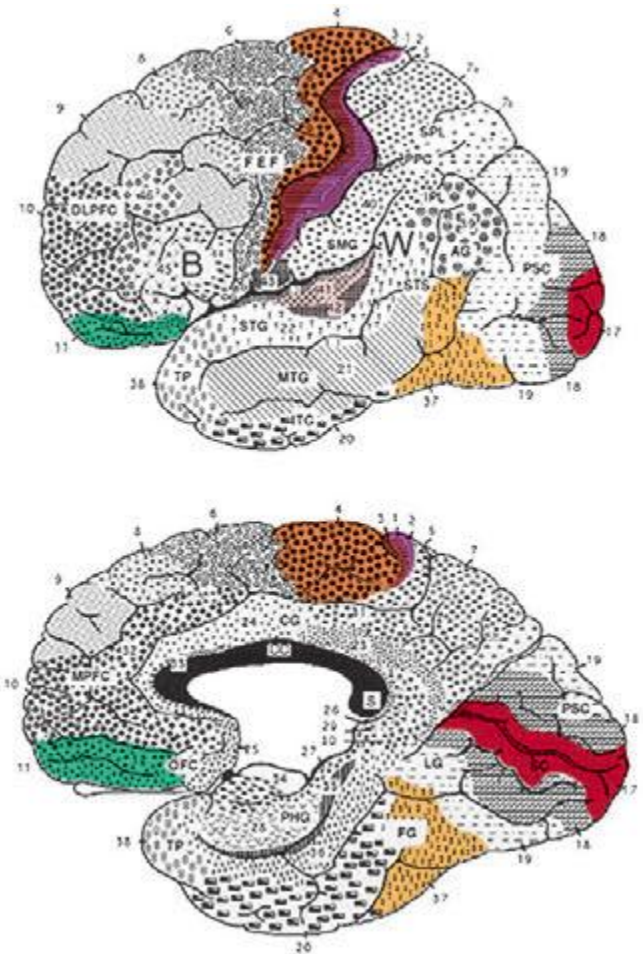


FIGURE 36-1 Lateral (*top*) and medial (*bottom*) views of the cerebral hemispheres. The numbers refer to the Brodmann cytoarchitectonic designations. Area 17 corresponds to the primary visual cortex, 41–42 to the primary auditory cortex, 1–3 to the primary somatosensory cortex, and 4 to the primary motor cortex. The rest of the cerebral cortex contains association areas. AG, angular gyrus; B, Broca's area; CC, corpus callosum; CG, cingulate gyrus; DLPFC, dorsolateral prefrontal cortex; FEF, frontal eye fields (premotor cortex); FG, fusiform gyrus; IPL, inferior parietal lobule; ITG, inferior temporal gyrus; LG, lingual gyrus; MPFC, medial prefrontal cortex; MTG, middle temporal gyrus; OFC, orbitofrontal cortex; PHG, parahippocampal gyrus; PPC, posterior parietal cortex; PSC, peristriate cortex; SC, striate cortex; SMG, supramarginal gyrus; SPL, superior parietal lobule; STG, superior temporal gyrus; STS, superior temporal sulcus; TP, temporopolar cortex; W, Wernicke's area.

Cognitive and behavioral functions (domains) are coordinated by intersecting *large-scale neural networks* that contain interconnected cortical and subcortical components. Five anatomically defined large-scale networks are most relevant to clinical practice: (1) a perisylvian network for language, (2) a parietofrontal network for spatial orientation, (3) an occipitotemporal network for face and object recognition, (4) a limbic network for retentive memory, and (5) a prefrontal network for the executive control of cognition and comportment.

THE LEFT PERISYLVIAN NETWORK FOR APHASIAS

The areas that are critical for language make up a distributed network located along the perisylvian region of the *left* hemisphere. One hub, located in the inferior frontal gyrus, is known as Broca's area. Damage to this region impairs phonology, fluency, and the grammatical structure of sentences. The location of a second hub, known as *Wernicke's area*, is less clearly settled but is traditionally thought to include the posterior parts of the temporal lobe. Cerebrovascular accidents that damage this area interfere with the ability to understand spoken or written sentences as well as the ability to express thoughts through meaningful words and statements. These two hubs are interconnected with each other and with surrounding parts of the frontal, parietal, and temporal lobes. Damage to this network gives rise to language impairments known as aphasia. Aphasia should be diagnosed only when there are deficits in the formal aspects of language, such as word finding, word choice, comprehension, spelling, or grammar. Dysarthria and mutism do not by themselves lead to a diagnosis of aphasia. In approximately 90% of right-handers and 60% of left-handers, aphasia occurs only after lesions of the left hemisphere.

CLINICAL EXAMINATION

The clinical examination of language should include the assessment of naming, spontaneous speech, comprehension, repetition, reading, and writing. A deficit of naming (*anomia*) is the single most common finding in aphasic patients. When asked to name a common object, the patient may fail to come up with the appropriate word, may provide a circumlocutious description of the object ("the thing for writing"), or may come up with the wrong word (*paraphasia*). If the patient offers an incorrect but related word ("pen" for "pencil"), the naming error is known as a *semantic paraphasia*; if the word approximates the correct answer but is phonetically inaccurate ("plentil" for "pencil"), it is known as a *phonemic paraphasia*. In most anomias, the patient cannot retrieve the appropriate name when shown an object but can point to the appropriate object when the name is provided by the examiner. This is known as a one-way (or retrieval-based) naming deficit. A two-way (comprehension-based) naming deficit exists if the patient can neither provide nor recognize the correct name. *Spontaneous speech* is described as "fluent" if it maintains appropriate output volume, phrase length, and melody or as "nonfluent" if it is sparse and halting and average utterance length is below four words. The

examiner also should note the integrity of *grammar* as manifested by word order (syntax), tenses, suffixes, prefixes, plurals, and possessives. *Comprehension* can be tested by assessing the patient's ability to follow conversation, asking yes-no questions ("Can a dog fly?" "Does it snow in summer?"), asking the patient to point to appropriate objects ("Where is the source of illumination in this room?"), or asking for verbal definitions of single words. *Repetition* is assessed by asking the patient to repeat single words, short sentences, or strings of words such as "No ifs, ands, or buts." The testing of repetition with tongue twisters such as "hippopotamus" and "Irish constabulary" provides a better assessment of dysarthria and palilalia than of aphasia. It is important to make sure that the number of words does not exceed the patient's attention span. Otherwise, the failure of repetition becomes a reflection of the narrowed attention span (working memory) rather than an indication of an aphasic deficit. *Reading* should be assessed for deficits in reading aloud as well as comprehension. *Alexia* describes an inability to either read aloud or comprehend single words and simple sentences; *agraphia* (or dysgraphia) is used to describe an acquired deficit in spelling.

Aphasias can arise acutely in cerebrovascular accidents (CVAs) or gradually in neurodegenerative diseases. The syndromes listed in [Table 36-1](#) are most applicable to the former group, where gray matter and white matter at the lesion site are abruptly and jointly destroyed. Progressive neurodegenerative diseases can have cellular, laminar, and regional specificity, giving rise to a different set of aphasias that will be described separately. The syndromes outlined below are idealizations and rarely occur in pure form.

Wernicke's Aphasia Comprehension is impaired for spoken and written words and sentences. Language output is fluent but is highly paraphasic and circumlocutious. Paraphasic errors may lead to strings of neologisms, which lead to "jargon aphasia." Speech contains few substantive nouns. The output is therefore voluminous but uninformative. For example, a patient attempts to describe how his wife accidentally threw away something important, perhaps his dentures: "We don't need it anymore, she says. And with it when that was downstairs was my teeth-tick ... a ... den ... dentith ... my dentist. And they happened to be in that bag ... see? ... Where my two ... two little pieces of dentist that I use ... that I ... all gone. If she throws the whole thing away ... visit some friends of hers and she can't throw them away."

Gestures and pantomime do not improve communication. The patient may not realize that his or her language is incomprehensible and may appear angry and impatient when the examiner fails to decipher the meaning of a severely paraphasic statement. In some patients this type of aphasia can be associated with severe agitation and paranoia. The ability to follow commands aimed at axial musculature may be preserved. The dissociation between the failure to understand simple questions ("What is your name?") in a patient who rapidly closes his or her eyes, sits up, or rolls over when asked to do so is characteristic of Wernicke's aphasia and helps differentiate it from

TABLE 36-1 CLINICAL FEATURES OF APHASIAS AND RELATED CONDITIONS COMMONLY SEEN IN CEREBROVASCULAR ACCIDENTS

	Comprehension	Repetition of Spoken Language	Naming	Fluency
Wernicke's	Impaired	Impaired	Impaired	Preserved or increased
Broca's	Preserved (except grammar)	Impaired	Impaired	Decreased
Global	Impaired	Impaired	Impaired	Decreased
Conduction	Preserved	Impaired	Impaired	Preserved
Nonfluent (anterior) transcortical	Preserved	Preserved	Impaired	Impaired
Fluent (posterior) transcortical	Impaired	Preserved	Impaired	Preserved
Isolation	Impaired	Echolalia	Impaired	No purposeful speech
Anomic	Preserved	Preserved	Impaired	Preserved except for word-finding pauses
Pure word deafness	Impaired only for spoken language	Impaired	Preserved	Preserved
Pure alexia	Impaired only for reading	Preserved	Preserved	Preserved

deafness, psychiatric disease, or malingering. Patients with Wernicke's aphasia cannot express their thoughts in meaning-appropriate words and cannot decode the meaning of words in any modality of input. This aphasia therefore has expressive as well as receptive components. Repetition, naming, reading, and writing also are impaired.

The lesion site most commonly associated with Wernicke's aphasia is the posterior portion of the language network. An embolus to the inferior division of the middle cerebral artery, to the posterior temporal or angular branches in particular, is the most common etiology (Chap. 446). Intracerebral hemorrhage, head trauma, and neoplasm are other causes of Wernicke's aphasia. A coexisting right hemianopia or superior quadrantanopia is common, and mild right nasolabial flattening may be found, but otherwise, the examination is often unrevealing. The paraphasic, neologistic speech in an agitated patient with an otherwise unremarkable neurologic examination may lead to the suspicion of a primary psychiatric disorder such as schizophrenia or mania, but the other components characteristic of acquired aphasia and the absence of prior psychiatric disease usually settle the issue. Prognosis for recovery of language function is guarded.

Broca's Aphasia Speech is nonfluent, labored, interrupted by many word-finding pauses, and usually dysarthric. It is impoverished in function words but enriched in meaning-appropriate nouns. Abnormal word order and the inappropriate deployment of *bound morphemes* (word endings used to denote tenses, possessives, or plurals) lead to a characteristic agrammatism. Speech is telegraphic and pithy but quite informative. In the following passage, a patient with Broca's aphasia describes his medical history: "I see ... the doctor, doctor sent me ... Bosson. Go to hospital. Doctor ... kept me beside. Two, tee days, doctor send me home."

Output may be reduced to a grunt or single word ("yes" or "no"), which is emitted with different intonations in an attempt to express approval or disapproval. In addition to fluency, naming and repetition are impaired. Comprehension of spoken language is intact except for syntactically difficult sentences with a passive voice structure or embedded clauses, indicating that Broca's aphasia is not just an "expressive" or "motor" disorder and that it also may involve a comprehension deficit in decoding syntax. Patients with Broca's aphasia can be tearful, easily frustrated, and profoundly depressed. Insight into their condition is preserved, in contrast to Wernicke's aphasia. Even when spontaneous speech is severely dysarthric, the patient may be able to display a relatively normal articulation of words when singing. This dissociation has been used to develop specific therapeutic approaches (melodic intonation therapy) for Broca's aphasia. Additional neurologic deficits include right facial weakness, hemiparesis or hemiplegia, and a buccofacial apraxia characterized by an inability to carry out motor commands involving oropharyngeal and facial musculature (e.g., patients are unable to demonstrate how to blow out a match or suck through a straw). The cause is most often infarction of Broca's area (the inferior frontal convolution; "B" in Fig. 36-1) and surrounding anterior perisylvian and insular cortex due to occlusion of the superior division of the middle cerebral artery (Chap. 446). Mass lesions, including tumor, intracerebral hemorrhage, and abscess, also may be responsible. When the cause of Broca's aphasia is stroke, recovery of language function generally peaks within 2 to 6 months, after which time further progress is limited. Speech therapy is more successful than in Wernicke's aphasia.

Conduction Aphasia Speech output is fluent but contains many phonemic paraphasias, comprehension of spoken language is intact, and repetition is severely impaired. Naming elicits phonemic paraphasias, and spelling is impaired. Reading aloud is impaired, but reading comprehension is preserved. The lesion sites spare the functionality of Broca's and Wernicke's areas but may induce a disconnection between the two. Occasionally, a transient Wernicke's aphasia may rapidly resolve into a conduction aphasia. The paraphasic output in conduction aphasia interferes with the ability to express meaning, but this deficit is not nearly as severe as the one displayed by patients with Wernicke's aphasia. Associated neurologic signs in conduction aphasia vary according to the primary lesion site.

Transcortical Aphasias: Fluent and Nonfluent Clinical features of *fluent (posterior) transcortical aphasia* are similar to those of Wernicke's aphasia, but repetition is intact. The lesion site disconnects the intact core of the language network from other temporoparietal association areas. Associated neurologic findings may include hemianopia. Cerebrovascular lesions (e.g., infarctions in the posterior watershed zone) and neoplasms that involve the temporoparietal cortex posterior to Wernicke's area are common causes. The features of *nonfluent (anterior) transcortical aphasia* are similar to those of Broca's aphasia, but repetition is intact and agrammatism is less pronounced. The neurologic examination may be otherwise intact, but a right hemiparesis also can exist. The lesion site disconnects the intact language network from prefrontal areas of the brain and usually involves the anterior watershed zone between anterior and middle cerebral artery territories or the supplementary motor cortex in the territory of the anterior cerebral artery.

Global and Isolation Aphasias *Global aphasia* represents the combined dysfunction of Broca's and Wernicke's areas and usually results from strokes that involve the entire middle cerebral artery distribution in the left hemisphere. Speech output is nonfluent, and comprehension of language is severely impaired. Related signs include right hemiplegia, hemisensory loss, and homonymous hemianopia. *Isolation aphasia* represents a combination of the two transcortical aphasias. Comprehension is severely impaired, and there is no purposeful speech output. The patient may parrot fragments of heard conversations (*echolalia*), indicating that the neural mechanisms for repetition are at least partially intact. This condition represents the pathologic function of the language network when it is isolated from other regions of the brain. Broca's and Wernicke's areas tend to be spared, but there is damage to the surrounding frontal, parietal, and temporal cortex. Lesions are patchy and can be associated with anoxia, carbon monoxide poisoning, or complete watershed zone infarctions.

Anomic Aphasia This form of aphasia may be considered the "minimal dysfunction" syndrome of the language network. Articulation, comprehension, and repetition are intact, but confrontation naming, word finding, and spelling are impaired. Word-finding pauses are uncommon, so language output is fluent but paraphasic, circumlocutious, and uninformative. The lesion sites can be anywhere within the left hemisphere language network, including the middle and inferior temporal gyri. *Anomic aphasia is the single most common language disturbance seen in head trauma, metabolic encephalopathy, and Alzheimer's disease.*

Pure Word Deafness The most common causes are either bilateral or left-sided middle cerebral artery (MCA) strokes affecting the superior temporal gyrus. The net effect of the underlying lesion is to interrupt the flow of information from the auditory association cortex to the language network. Patients have no difficulty understanding written language and can express themselves well in spoken or written language. They have no difficulty interpreting and reacting to environmental sounds since primary auditory cortex and auditory association areas of the right hemisphere are spared. Because auditory information cannot be conveyed to the language network, however, it cannot be decoded into neural word representations, and the patient reacts to speech as if it were in an alien tongue that cannot be deciphered. Patients cannot repeat spoken language but have no difficulty naming objects. In time, patients with pure word deafness teach themselves lipreading and may appear to have improved. There may be no additional neurologic findings, but agitated paranoid reactions are common in the acute stages. Cerebrovascular lesions are the most common cause.

Pure Alexia Without Agraphia This is the visual equivalent of pure word deafness. The lesions (usually a combination of damage to the left occipital cortex and to a posterior sector of the corpus callosum—the splenium) interrupt the flow of visual input into the language network. There is usually a right hemianopia, but the core language network remains unaffected. The patient can understand and produce spoken language, name objects in the left visual hemifield, repeat, and write. However, the patient acts as if illiterate when asked to read even the

simplest sentence because the visual information from the written words (presented to the intact left visual hemifield) cannot reach the language network. Objects in the left hemifield may be named accurately because they activate nonvisual associations in the right hemisphere, which in turn can access the language network through transcallosal pathways anterior to the splenium. Patients with this syndrome also may lose the ability to name colors, although they can match colors. This is known as a *color anomia*. The most common etiology of pure alexia is a vascular lesion in the territory of the posterior cerebral artery or an infiltrating neoplasm in the left occipital cortex that involves the optic radiations as well as the crossing fibers of the splenium. Because the posterior cerebral artery also supplies medial temporal components of the limbic system, a patient with pure alexia also may experience an amnesia, but this is usually transient because the limbic lesion is unilateral.

Apraxia and Aphemia *Apraxia* designates a complex motor deficit that cannot be attributed to pyramidal, extrapyramidal, cerebellar, or sensory dysfunction and that does not arise from the patient's failure to understand the nature of the task. *Apraxia of speech* is used to designate articulatory abnormalities in the duration, fluidity, and stress of syllables that make up words. Intoning the words may improve articulation. It can arise with CVAs in the posterior part of Broca's area or in the course of frontotemporal lobar degeneration (FTLD) with tauopathy. *Aphemia* is a severe form of acute speech apraxia that presents with severely impaired fluency (often mutism). Recovery is the rule and involves an intermediate stage of hoarse whispering. Writing, reading, and comprehension are intact, and so this is not a true aphasic syndrome. CVAs in parts of Broca's area or subcortical lesions that undercut its connections with other parts of the brain may be present. Occasionally, the lesion site is on the medial aspects of the frontal lobes and may involve the supplementary motor cortex of the left hemisphere. *Ideomotor apraxia* is diagnosed when commands to perform a specific motor act ("cough," "blow out a match") or pantomime the use of a common tool (a comb, hammer, straw, or toothbrush) in the absence of the real object cannot be followed. The patient's ability to comprehend the command is ascertained by demonstrating multiple movements and establishing that the correct one can be recognized. Some patients with this type of apraxia can imitate the appropriate movement (when it is demonstrated by the examiner) and show no impairment when handed the real object, indicating that the sensorimotor mechanisms necessary for the movement are intact. Some forms of ideomotor apraxia represent a disconnection of the language network from pyramidal motor systems so that commands to execute complex movements are understood but cannot be conveyed to the appropriate motor areas. *Buccofacial apraxia* involves apraxic deficits in movements of the face and mouth. *Limb apraxia* encompasses apraxic deficits in movements of the arms and legs. Ideomotor apraxia almost always is caused by lesions in the left hemisphere and is commonly associated with aphasic syndromes, especially Broca's aphasia and conduction aphasia. Because the handling of real objects is not impaired, ideomotor apraxia by itself causes no major limitation of daily living activities. Patients with lesions of the anterior corpus callosum can display ideomotor apraxia confined to the left side of the body, a sign known as *sympathetic dyspraxia*. A severe form of sympathetic dyspraxia, known as the *alien hand* syndrome, is characterized by additional features of motor disinhibition on the left hand. *Ideational apraxia* refers to a deficit in the sequencing of goal-directed movements in patients who have no difficulty executing the individual components of the sequence. For example, when the patient is asked to pick up a pen and write, the sequence of uncapping the pen, placing the cap at the opposite end, turning the point toward the writing surface, and writing may be disrupted, and the patient may be seen trying to write with the wrong end of the pen or even with the removed cap. These motor sequencing problems usually are seen in the context of confusional states and dementias rather than focal lesions associated with aphasic conditions. *Limb-kinetic apraxia* involves clumsiness in the use of tools or objects that cannot be attributed to sensory, pyramidal, extrapyramidal, or cerebellar dysfunction. This condition can

emerge in the context of focal premotor cortex lesions or *corticobasal degeneration*.

Gerstmann's Syndrome The combination of *acalculia* (impairment of simple arithmetic), *dysgraphia* (impaired writing), *finger anomia* (an inability to name individual fingers such as the index and thumb), and *right-left confusion* (an inability to tell whether a hand, foot, or arm of the patient or examiner is on the right or left side of the body) is known as Gerstmann's syndrome. In making this diagnosis, it is important to establish that the finger and left-right naming deficits are not part of a more generalized anomia and that the patient is not otherwise aphasic. When Gerstmann's syndrome arises acutely and in isolation, it is commonly associated with damage to the inferior parietal lobule (especially the angular gyrus) in the left hemisphere.

Pragmatics and Prosody *Pragmatics* refers to aspects of language that communicate attitude, affect, and the figurative rather than literal aspects of a message (e.g., "green thumb" does not refer to the actual color of the finger). One component of pragmatics, *prosody*, refers to variations of melodic stress and intonation that influence attitude and the inferential aspect of verbal messages. For example, the two statements "He is clever." and "He is *clever*?" contain an identical word choice and syntax but convey vastly different messages because of differences in the intonation with which the statements are uttered. Damage to right hemisphere regions corresponding to Broca's area impairs the ability to introduce meaning-appropriate prosody into spoken language. The patient produces grammatically correct language with accurate word choice, but the statements are uttered in a monotone that interferes with the ability to convey the intended stress and affect. Patients with this type of *aprosodia* give the mistaken impression of being depressed or indifferent. Other aspects of pragmatics, especially the ability to infer the figurative aspect of a message, become impaired by damage to the right hemisphere or frontal lobes.

Subcortical Aphasia Damage to subcortical components of the language network (e.g., the striatum and thalamus of the left hemisphere) also can lead to aphasia. The resulting syndromes contain combinations of deficits in the various aspects of language but rarely fit the specific patterns described in Table 36-1. In a patient with a CVA, an anomic aphasia accompanied by dysarthria or a fluent aphasia with hemiparesis should raise the suspicion of a subcortical lesion site.

Progressive Aphasias Aphasias caused by major cerebrovascular accidents start suddenly and display maximal deficits at the onset. These are the "classic" aphasias described above. Aphasias caused by neurodegenerative diseases have an insidious onset and relentless progression. The neuropathology can be selective not only for gray matter but also for specific layers and cell types. The clinico-anatomic patterns are therefore different from those described in Table 36-1.

CLINICAL PRESENTATION AND DIAGNOSIS OF PRIMARY PROGRESSIVE APHASIA (PPA) Several neurodegenerative syndromes, such as typical Alzheimer-type (amnestic) and frontal-type (behavioral) dementias, can also undermine language as the disease progresses. In these cases, the aphasia is an ancillary component of the overall syndrome. When a neurodegenerative language disorder arises in relative isolation and becomes the primary concern that brings the patient to medical attention, a diagnosis of PPA is made.

LANGUAGE IN PPA The impairments of language in PPA have slightly different patterns from those seen in CVA-caused aphasias. Three major subtypes of PPA can be recognized. The *agrammatic variant* is characterized by consistently low fluency and impaired grammar but intact word comprehension. It most closely resembles Broca's aphasia or anterior transcortical aphasia but usually lacks the right hemiparesis or dysarthria and has more profound impairments of grammar. Peak sites of neuronal loss (gray matter atrophy) include the left inferior frontal gyrus where Broca's area is located. The neuropathology is usually an FTLD with tauopathy but can also be an atypical form of Alzheimer's disease (AD) pathology. The *semantic variant* is characterized by preserved fluency and syntax but poor single-word comprehension and profound two-way naming impairments. This

kind of aphasia is not seen with CVAs. It differs from Wernicke's aphasia or posterior transcortical aphasia because speech is usually informative, repetition is intact, and comprehension of conversation is relatively preserved, as long as the meaning is not too dependent on words that the patient fails to understand. Peak atrophy sites are located in the left anterior temporal lobe, indicating that this part of the brain plays a critical role in the comprehension of words, especially words that denote concrete objects. The neuropathology is frequently an FTLN with abnormal precipitates of the 43-kDa transactive response DNA-binding protein TDP-43. The *logopenic variant* is characterized by preserved syntax and comprehension but frequent and severe word-finding pauses, anomia, circumlocutions, and simplifications during spontaneous speech. Peak atrophy sites are located in the temporoparietal junction and posterior temporal lobe, partially overlapping with traditional location of Wernicke's area. However, the comprehension impairment of *Wernicke's aphasia* is absent, perhaps because the underlying white matter, frequently damaged by cerebrovascular accidents, remains relatively intact in PPA. In contrast to Broca's aphasia or agrammatic PPA, the interruption of fluency is variable so that speech may appear entirely normal if the patient is allowed to engage in small talk. Logopenic PPA resembles the anomic aphasia of Table 36-1 but usually has longer and more frequent word-finding pauses. Patients may also have poor phrase and word repetition, in which case the aphasia resembles the *conduction aphasia* in Table 36-1. Of all PPA subtypes, this is the one most commonly associated with the pathology of AD, but FTLN can also be the cause. In addition to these three major subtypes, PPA can also present in the form of *pure word deafness* or *Gerstmann's syndrome*.

THE PARIETOFRONTAL NETWORK FOR NEGLECT AND RELATED CONDITIONS

Adaptive spatial orientation is subserved by a large-scale network containing three major cortical components. The *cingulate cortex* provides access to a motivational mapping of the extrapersonal space, the *posterior parietal cortex* to a sensorimotor representation of salient extrapersonal events, and the *frontal eye fields* to motor strategies for attentional behaviors (Fig. 36-2). Subcortical components of this network include the striatum and the thalamus. Damage to this network can undermine the distribution of attention within the extrapersonal space, giving rise to hemispatial neglect, simultanagnosia and object finding failures. The integration of egocentric (self-centered) with allocentric (object-centered) coordinates can also be disrupted, giving rise to impairments in route finding, the ability to avoid obstacles, and the ability to dress.

HEMISPATIAL NEGLECT

Contralesional hemispatial neglect represents one outcome of damage to the cortical or subcortical components of this network. *The traditional view that hemispatial neglect always denotes a parietal lobe lesion is inaccurate.* According to one model of spatial cognition, the right hemisphere directs attention within the *entire* extrapersonal space, whereas the left hemisphere directs attention mostly within the contralateral right hemispace. Consequently, left hemisphere lesions do not give rise to much contralesional neglect because the global attentional mechanisms of the right hemisphere can compensate for the loss of the *contralaterally* directed attentional functions of the left hemisphere. Right hemisphere lesions, however, give rise to severe contralesional left hemispatial neglect because the unaffected left hemisphere does not contain ipsilateral attentional mechanisms. This model is consistent with clinical experience, which shows that contralesional neglect is more common, more severe, and longer lasting after damage to the right hemisphere than after damage to the left hemisphere. Severe neglect for the right hemispace is rare, even in left-handers with left hemisphere lesions.

Clinical Examination Patients with severe neglect may fail to dress, shave, or groom the left side of the body; fail to eat food placed on the

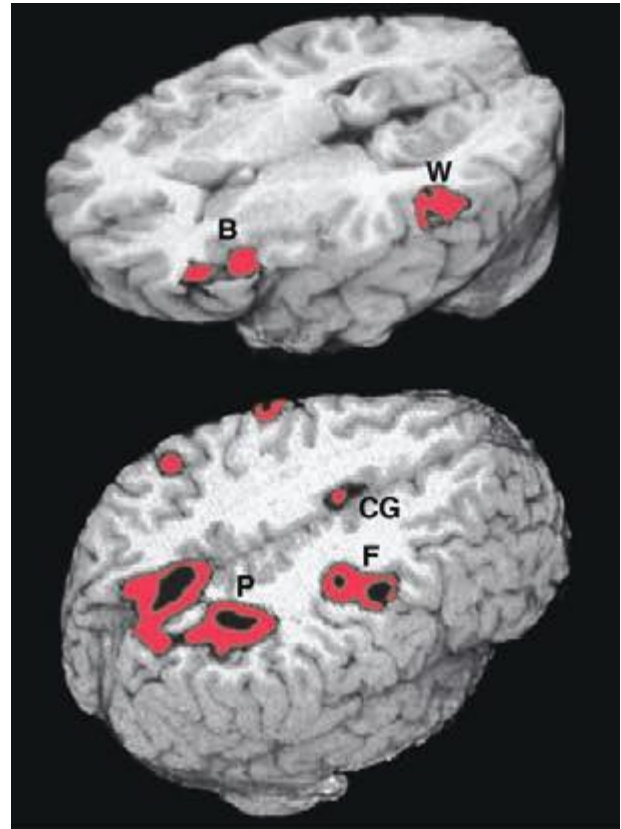
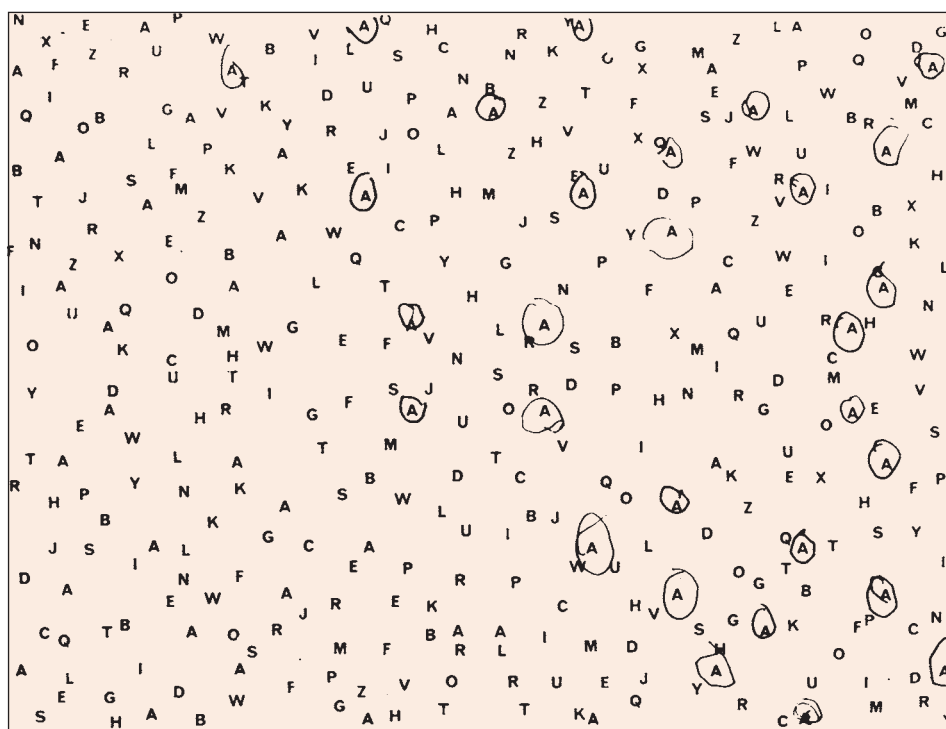


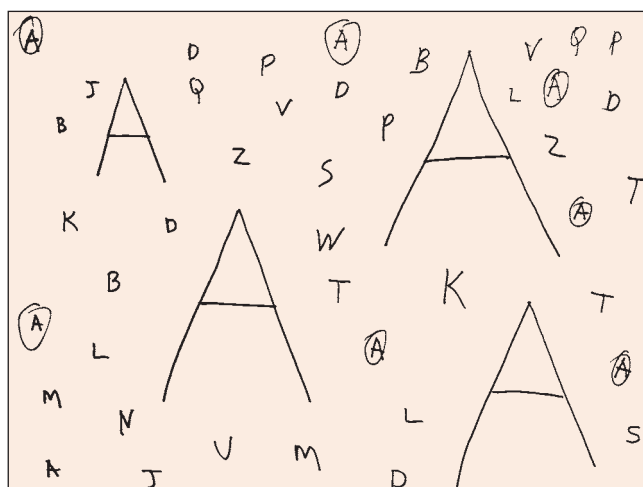
FIGURE 36-2 Functional magnetic resonance imaging of language and spatial attention in neurologically intact subjects.

The red and black areas show regions of task-related significant activation. (Top) The subjects were asked to determine if two words were synonymous. This language task led to the simultaneous activation of the two epicenters of the language network, Broca's area (B) and Wernicke's area (W). The activations are exclusively in the left hemisphere. (Bottom) The subjects were asked to shift spatial attention to a peripheral target. This task led to the simultaneous activation of the three epicenters of the attentional network: the posterior parietal cortex (P), the frontal eye fields (F), and the cingulate gyrus (CG). The activations are predominantly in the right hemisphere. (Courtesy of Darren Gitelman, MD; with permission.)

left side of the tray; and fail to read the left half of sentences. When asked to copy a simple line drawing, the patient fails to copy detail on the left, and when the patient is asked to write, there is a tendency to leave an unusually wide margin on the left. Two bedside tests that are useful in assessing neglect are *simultaneous bilateral stimulation* and *visual target cancellation*. In the former, the examiner provides either unilateral or simultaneous bilateral stimulation in the visual, auditory, and tactile modalities. After right hemisphere injury, patients who have no difficulty detecting unilateral stimuli on either side experience the bilaterally presented stimulus as coming only from the right. This phenomenon is known as *extinction* and is a manifestation of the sensory-representational aspect of hemispatial neglect. In the target detection task, targets (e.g., A's) are interspersed with foils (e.g., other letters of the alphabet) on a 21.5- to 28.0-cm (8.5 to 11 in.) sheet of paper, and the patient is asked to circle all the targets. A failure to detect targets on the left is a manifestation of the exploratory (motor) deficit in hemispatial neglect (Fig. 36-3A). Hemianopia is not by itself sufficient to cause the target detection failure because the patient is free to turn the head and eyes to the left. Target detection failures therefore reflect a distortion of spatial attention, not just of sensory input. Some patients with neglect also may deny the existence of hemiparesis and may even deny ownership of the paralyzed limb, a condition known as *anosognosia*.



A



B

FIGURE 36-3 **A.** A 47-year-old man with a large frontoparietal lesion in the right hemisphere was asked to circle all the A's. Only targets on the right are circled. This is a manifestation of left hemispatial neglect. **B.** A 70-year-old woman with a 2-year history of degenerative dementia was able to circle most of the small targets but ignored the larger ones. This is a manifestation of simultanagnosia.

BÁLINT'S SYNDROME, SIMULTANAGNOSIA, DRESSING APRAXIA, CONSTRUCTION APRAXIA, AND ROUTE FINDING

Bilateral involvement of the network for spatial attention, especially its parietal components, leads to a state of severe spatial disorientation known as *Bálint's syndrome*. Bálint's syndrome involves deficits in the orderly visuomotor scanning of the environment (*oculomotor apraxia*), accurate manual reaching toward visual targets (*optic ataxia*), and the ability to integrate visual information in the center of gaze with more peripheral information (*simultanagnosia*). A patient with simultanagnosia “misses the forest for the trees.” For example, a patient who is shown a table lamp and asked to name the object may look at its circular base and call it an ashtray. Some patients with simultanagnosia report that objects they look at may vanish suddenly, probably indicating an inability to look back at the original point of gaze after brief saccadic displacements. Movement and distracting stimuli greatly exacerbate the difficulties of visual perception. Simultanagnosia can occur without the other two components of Bálint's syndrome.

A modification of the letter cancellation task described above can be used for the bedside diagnosis of simultanagnosia. In this modification, some of the targets (e.g., A's) are made to be much larger than the others (7.5 to 10 cm vs 2.5 cm [3 to 4 in. vs 1 in.] in height), and all targets are embedded among foils. Patients with simultanagnosia display a counterintuitive but characteristic tendency to miss the larger targets (Fig. 36-3B). This occurs because the information needed for the identification of the larger targets cannot be confined to the immediate line of gaze and requires the integration of visual information across multiple fixation points. The greater difficulty in the detection of the larger targets also indicates that poor acuity is not responsible for the impairment of visual function and that the problem is central rather than peripheral. The test shown in Fig. 36-3B is not by itself sufficient to diagnose simultanagnosia because some patients with a frontal network syndrome may omit the large letters, perhaps because they lack the mental flexibility needed to realize that the two types of targets are symbolically identical despite being superficially different.

Bilateral parietal lesions can impair the integration of egocentric with allocentric spatial coordinates. One manifestation is *Dressing apraxia*. A patient with this condition is unable to align the body axis with the axis of the garment and can be seen struggling as he or she holds a coat from its bottom or extends his or her arm into a fold of the garment rather than into its sleeve. Lesions that involve the posterior parietal cortex also lead to severe difficulties in copying simple line drawings. This is known as a *construction apraxia* and is much more severe if the lesion is in the right hemisphere. In some patients with right hemisphere lesions, the drawing difficulties are confined to the left side of the figure and represent a manifestation of hemispatial neglect; in others, there is a more universal deficit in reproducing contours and three-dimensional perspective. Impairments of route finding can be included in this group of disorders, which reflect an inability to orient the self with respect to external objects and landmarks.

Causes of Spatial Disorientation Cerebrovascular lesions and neoplasms in the right hemisphere are common causes of hemispatial neglect. Depending on the site of the lesion, a patient with neglect also may have hemiparesis, hemihypesthesia, and hemianopia on the left, but these are not invariant findings. The majority of these patients display considerable improvement of hemispatial neglect, usually within the first several weeks. Bálint's syndrome, dressing apraxia, and route finding impairments are more likely to result from bilateral dorsal parietal lesions; common settings for acute onset include watershed infarction between the middle and posterior cerebral artery territories, hypoglycemia, and sagittal sinus thrombosis.

A progressive form of spatial disorientation, known as the *posterior cortical atrophy* syndrome, most commonly represents a variant of AD with unusual concentrations of neurofibrillary degeneration in the parieto-occipital cortex and the superior colliculus. The patient displays a progressive hemispatial neglect or Bálint's syndrome, usually accompanied by dressing and construction apraxia. The corticobasal syndrome, which can be caused by AD or FTLN pathology, can also lead to a progressive left hemineglect syndrome. Both syndromes can impair route finding.

THE OCCIPITOTEMPORAL NETWORK FOR FACE AND OBJECT RECOGNITION

A patient with *prosopagnosia* cannot recognize familiar faces, including, sometimes, the reflection of his or her own face in the mirror. This is not a perceptual deficit because prosopagnosic patients easily can tell whether two faces are identical. Furthermore, a prosopagnosic patient who cannot recognize a familiar face by visual inspection alone can use auditory cues to reach appropriate recognition if allowed to listen to the person's voice. The deficit in prosopagnosia is therefore modality-specific and reflects the existence of a lesion that prevents the activation of otherwise intact multimodal templates by relevant visual input. Prosopagnosic patients characteristically have no difficulty with the generic identification of a face as a face or a car as a car, but may not recognize the identity of an individual face or the make of an individual car. This reflects a visual recognition deficit for proprietary features that characterize individual members of an object class. When recognition problems become more generalized and extend to the generic identification of common objects, the condition is known as *visual object agnosia*. A patient with anomia cannot name the object but can describe its use. In contrast, a patient with visual agnosia is unable either to name a visually presented object or to describe its use. Face and object recognition disorders also can result from the simultanagnosia of Bálint's syndrome, in which case they are known as *apperceptive* agnosias as opposed to the *associative* agnosias that result from inferior temporal lobe lesions.

CAUSES

The characteristic lesions in prosopagnosia and visual object agnosia of acute onset consist of bilateral infarctions in the territory of the posterior cerebral arteries. Associated deficits can include visual field defects (especially superior quadrantanopias) and a centrally based

color blindness known as achromatopsia. Rarely, the responsible lesion is unilateral. In such cases, prosopagnosia is associated with lesions in the right hemisphere, and object agnosia with lesions in the left. Degenerative diseases of anterior and inferior temporal cortex can cause progressive associative prosopagnosia and object agnosia. The combination of progressive associative agnosia and a fluent aphasia is known as *semantic dementia*. Patients with semantic dementia fail to recognize faces and objects and cannot understand the meaning of words denoting objects. This needs to be differentiated from the semantic type of PPA where there is severe impairment in understanding words that denote objects and in naming faces and objects but a relative preservation of face and object recognition.

LIMBIC NETWORK FOR MEMORY AND AMNESIA

Limbic and paralimbic areas (such as the hippocampus, amygdala, and entorhinal cortex), the anterior and medial nuclei of the thalamus, the medial and basal parts of the striatum, and the hypothalamus collectively constitute a distributed network known as the *limbic system*. The behavioral affiliations of this network include the coordination of emotion, motivation, autonomic tone, and endocrine function. An additional area of specialization for the limbic network and the one that is of most relevance to clinical practice is that of declarative (explicit) memory for recent episodes and experiences. A disturbance in this function is known as an *amnesic state*. In the absence of deficits in motivation, attention, language, or visuospatial function, the clinical diagnosis of a persistent global amnesic state is always associated with bilateral damage to the limbic network, usually within the hippocampo-entorhinal complex or the thalamus. Damage to the limbic network does not necessarily destroy memories but interferes with their conscious recall in coherent form. The individual fragments of information remain preserved despite the limbic lesions and can sustain what is known as *implicit memory*. For example, patients with amnesic states can acquire new motor or perceptual skills even though they may have no conscious knowledge of the experiences that led to the acquisition of these skills.

The memory disturbance in the amnesic state is multimodal and includes retrograde and anterograde components. The *retrograde amnesia* involves an inability to recall experiences that occurred before the onset of the amnesic state. Relatively recent events are more vulnerable to retrograde amnesia than are more remote and more extensively consolidated events. A patient who comes to the emergency room complaining that he cannot remember his or her identity but can remember the events of the previous day almost certainly does not have a neurologic cause of memory disturbance. The second and most important component of the amnesic state is the *anterograde amnesia*, which indicates an inability to store, retain, and recall new knowledge. Patients with amnesic states cannot remember what they ate a few hours ago or the details of an important event they may have experienced in the recent past. In the acute stages, there also may be a tendency to fill in memory gaps with inaccurate, fabricated, and often implausible information. This is known as *confabulation*. Patients with the amnesic syndrome forget that they forget and tend to deny the existence of a memory problem when questioned. Confabulation is more common in cases where the underlying lesion also interferes with parts of the frontal network, as in the case of the Wernicke-Korsakoff syndrome or traumatic head injury.

CLINICAL EXAMINATION

A patient with an amnesic state is almost always disoriented, especially to time, and has little knowledge of current news. The anterograde component of an amnesic state can be tested with a list of four to five words read aloud by the examiner up to five times or until the patient can immediately repeat the entire list without an intervening delay. The next phase of the recall occurs after a period of 5 to 10 min during which the patient is engaged in other tasks. Amnesic patients fail this phase of the task and may even forget that they were given a list of words to remember. Accurate recognition of the words by multiple choice in a patient who cannot recall them indicates a less severe memory disturbance that affects mostly the retrieval stage of memory.

The retrograde component of an amnesia can be assessed with questions related to autobiographical or historic events. The anterograde component of amnesic states is usually much more prominent than the retrograde component. In rare instances, occasionally associated with temporal lobe epilepsy or herpes simplex encephalitis, the retrograde component may dominate. Confusional states caused by toxic-metabolic encephalopathies and some types of frontal lobe damage lead to secondary memory impairments, especially at the stages of encoding and retrieval, even in the absence of limbic lesions. This sort of memory impairment can be differentiated from the amnesic state by the presence of additional impairments in the attention-related tasks described below in the section on the frontal lobes.

CAUSES, INCLUDING ALZHEIMER'S DISEASE

Neurologic diseases that give rise to an amnesic state include tumors (of the sphenoid wing, posterior corpus callosum, thalamus, or medial temporal lobe), infarctions (in the territories of the anterior or posterior cerebral arteries), head trauma, herpes simplex encephalitis, Wernicke-Korsakoff encephalopathy, paraneoplastic limbic encephalitis, and degenerative dementias such as AD and Pick's disease. The one common denominator of all these diseases is the presence of bilateral lesions within one or more components in the limbic network. Occasionally, unilateral left-sided hippocampal lesions can give rise to an amnesic state, but the memory disorder tends to be transient. Depending on the nature and distribution of the underlying neurologic disease, the patient also may have visual field deficits, eye movement limitations, or cerebellar findings. AD and its prodromal state of mild cognitive impairment (MCI) are the most common causes of progressive memory impairments. The predilection of the entorhinal cortex and hippocampus for early neurofibrillary degeneration by typical AD pathology is responsible for the initially selective impairment of episodic memory. In time, additional impairments in language, attention, and visuospatial skills emerge as the neurofibrillary degeneration spreads to additional neocortical areas.

Transient global amnesia is a distinctive syndrome usually seen in late middle age. Patients become acutely disoriented and repeatedly ask who they are, where they are, and what they are doing. The spell is characterized by anterograde amnesia (inability to retain new information) and a retrograde amnesia for relatively recent events that occurred before the onset. The syndrome usually resolves within 24 to 48 h and is followed by the filling in of the period affected by the retrograde amnesia, although there is persistent loss of memory for the events that occurred during the ictus. Recurrences are noted in approximately 20% of patients. Migraine, temporal lobe seizures, and perfusion abnormalities in the posterior cerebral territory have been postulated as causes of transient global amnesia. The absence of associated neurologic findings occasionally may lead to the incorrect diagnosis of a psychiatric disorder.

THE PREFRONTAL NETWORK FOR EXECUTIVE FUNCTION AND BEHAVIOR

The frontal lobes can be subdivided into motor-premotor, dorsolateral prefrontal, medial prefrontal, and orbitofrontal components. The terms *frontal lobe syndrome* and *prefrontal cortex* refer only to the last three of these four components. These are the parts of the cerebral cortex that show the greatest phylogenetic expansion in primates, especially in humans. The dorsolateral prefrontal, medial prefrontal, and orbitofrontal areas, along with the subcortical structures with which they are interconnected (i.e., the head of the caudate and the dorsomedial nucleus of the thalamus), collectively make up a large-scale network that coordinates exceedingly complex aspects of human cognition and behavior.

The prefrontal network plays an important role in behaviors that require multitasking and the integration of thought with emotion. Cognitive operations impaired by prefrontal cortex lesions often are referred to as "executive functions." The most common clinical manifestations of damage to the prefrontal network take the form of two relatively distinct syndromes. In the *frontal abulic syndrome*, the patient shows a loss of initiative, creativity, and curiosity and displays

a pervasive emotional blandness, apathy, and lack of empathy. In the *frontal disinhibition syndrome*, the patient becomes socially disinhibited and shows severe impairments of judgment, insight, foresight, and the ability to mind rules of conduct. The dissociation between intact intellectual function and a total lack of even rudimentary common sense is striking. Despite the preservation of all essential memory functions, the patient cannot learn from experience and continues to display inappropriate behaviors without appearing to feel emotional pain, guilt, or regret when those behaviors repeatedly lead to disastrous consequences. The impairments may emerge only in real-life situations when behavior is under minimal external control and may not be apparent within the structured environment of the medical office. Testing judgment by asking patients what they would do if they detected a fire in a theater or found a stamped and addressed envelope on the road is not very informative because patients who answer these questions wisely in the office may still act very foolishly in real-life settings. The physician must therefore be prepared to make a diagnosis of frontal lobe disease based on historic information alone even when the mental state is quite intact in the office examination.

CLINICAL EXAMINATION

The emergence of developmentally primitive reflexes, also known as frontal release signs, such as grasping (elicited by stroking the palm) and sucking (elicited by stroking the lips) are seen primarily in patients with large structural lesions that extend into the premotor components of the frontal lobes or in the context of metabolic encephalopathies. The vast majority of patients with prefrontal lesions and frontal lobe behavioral syndromes do not display these reflexes. Damage to the frontal lobe disrupts a variety of attention-related functions, including working memory (the transient online holding and manipulation of information), concentration span, the scanning and retrieval of stored information, the inhibition of immediate but inappropriate responses, and mental flexibility. Digit span (which should be seven forward and five reverse) is decreased, reflecting poor working memory; the recitation of the months of the year in reverse order (which should take less than 15 s) is slowed as another indication of poor working memory; and the fluency in producing words starting with the letter a, f, or s that can be generated in 1 min (normally ≥ 12 per letter) is diminished even in nonaphasic patients, indicating an impairment in the ability to search and retrieve information from long-term stores. In "go-no go" tasks (where the instruction is to raise the finger upon hearing one tap but keep it still upon hearing two taps), the patient shows a characteristic inability to inhibit the response to the "no go" stimulus. Mental flexibility (tested by the ability to shift from one criterion to another in sorting or matching tasks) is impoverished; distractibility by irrelevant stimuli is increased; and there is a pronounced tendency for impersistence and perseveration. The ability for abstracting similarities and interpreting proverbs is also undermined.

The attentional deficits disrupt the orderly registration and retrieval of new information and lead to *secondary* memory deficits. The distinction of the underlying neural mechanisms is illustrated by the observation that severely amnesic patients who cannot remember events that occurred a few minutes ago may have intact if not superior working memory capacity as shown in tests of digit span.

CAUSES: TRAUMA, NEOPLASM, AND FRONTOTEMPORAL DEMENTIA

The abulic syndrome tends to be associated with damage in dorsolateral or dorsomedial prefrontal cortex, and the disinhibition syndrome with damage in orbitofrontal or ventromedial cortex. These syndromes tend to arise almost exclusively after bilateral lesions. Unilateral lesions confined to the prefrontal cortex may remain silent until the pathology spreads to the other side; this explains why thromboembolic CVA is an unusual cause of the frontal lobe syndrome. Common settings for frontal lobe syndromes include head trauma, ruptured aneurysms, hydrocephalus, tumors (including metastases, glioblastoma, and falx or olfactory groove meningiomas), and focal degenerative diseases. A major clinical form of FTLD known as the behavioral variant of frontotemporal dementia (bvFTD) causes a progressive frontal lobe syndrome. The behavioral changes can range

from apathy to shoplifting, compulsive gambling, sexual indiscretions, remarkable lack of common sense, new ritualistic behaviors, and alterations in dietary preferences, usually leading to increased taste for sweets or rigid attachment to specific food items. In many patients with AD, neurofibrillary degeneration eventually spreads to prefrontal cortex and gives rise to components of the frontal lobe syndrome, but almost always on a background of severe memory impairment. Rarely, the bvFTD syndrome can arise in isolation in the context of an atypical form of AD pathology.

Lesions in the caudate nucleus or in the dorsomedial nucleus of the thalamus (subcortical components of the prefrontal network) also can produce a frontal lobe syndrome. This is one reason why the changes in mental state associated with degenerative basal ganglia diseases such as Parkinson's disease and Huntington's disease display components of the frontal lobe syndrome. Bilateral multifocal lesions of the cerebral hemispheres, none of which are individually large enough to cause specific cognitive deficits such as aphasia and neglect, can collectively interfere with the connectivity and therefore integrating (executive) function of the prefrontal cortex. A frontal lobe syndrome is therefore the single most common behavioral profile associated with a variety of bilateral multifocal brain diseases, including metabolic encephalopathy, multiple sclerosis, and vitamin B₁₂ deficiency, among others. Many patients with the clinical diagnosis of a frontal lobe syndrome tend to have lesions that do not involve prefrontal cortex but involve either the subcortical components of the prefrontal network or its connections with other parts of the brain. To avoid making a diagnosis of "frontal lobe syndrome" in a patient with no evidence of frontal cortex disease, it is advisable to use the diagnostic term *frontal network syndrome*, with the understanding that the responsible lesions can lie anywhere within this distributed network. A patient with frontal lobe disease raises potential dilemmas in differential diagnosis: the abulia and blandness may be misinterpreted as depression, and the disinhibition as idiopathic mania or acting out. Appropriate intervention may be delayed while a treatable tumor keeps expanding.

CARING FOR PATIENTS WITH DEFICITS OF HIGHER CEREBRAL FUNCTION

Brain damage may cause a dissociation between feeling states and their expression so that a patient who may superficially appear jocular could still be suffering from an underlying depression that needs to be treated. If neuroleptics become absolutely necessary for the control of agitation, atypical neuroleptics are preferable because of their lower extrapyramidal side effects. Treatment with neuroleptics in elderly patients with dementia requires weighing the potential benefits against the potentially serious side effects.

Spontaneous improvement of cognitive deficits due to acute neurologic lesions is common. It is most rapid in the first few weeks but may continue for up to 2 years, especially in young individuals with single brain lesions. Some of the initial deficits appear to arise from remote dysfunction (diaschisis) in parts of the brain that are interconnected with the site of initial injury. Improvement in these patients may reflect, at least in part, a normalization of the remote dysfunction. Other mechanisms may involve functional reorganization in surviving neurons adjacent to the injury or the compensatory use of homologous structures, e.g., the right superior temporal gyrus with recovery from Wernicke's aphasia. Cognitive rehabilitation procedures have been used in the treatment of higher cortical deficits. There are few controlled studies, but some show a benefit of rehabilitation in the recovery from hemispatial neglect and aphasia. Determining driving competence is challenging, especially in the early stages of dementing diseases. The diagnosis of a neurodegenerative disease is not by itself sufficient for asking the patient to stop driving. An on-the-road driving test and reports from family members may help time decisions related to this very important activity.

Some of the deficits described in this chapter are so complex that they may bewilder not only the patient and family but also the physician. It is imperative to carry out a systematic clinical evaluation to characterize the nature of the deficits and explain them in lay terms to

the patient and family. An enlightened approach to patients with damage to the cerebral cortex requires an understanding of the principles that link neural networks to higher cerebral functions in health and disease.

37e

Primary Progressive Aphasia, Memory Loss, and Other Focal Cerebral Disorders

Maria Luisa Gorno-Tempini, Jennifer Ogar, Joel Kramer, Bruce L. Miller, Gil Rabinovici, Maria Carmela Tartaglia

Language and memory are essential human functions. For the experienced clinician, the recognition of different types of language and memory disturbances often provides essential clues to the anatomic localization and diagnosis of neurologic disorders. This video illustrates classic disorders of language and speech (including the aphasias), memory (the amnesias), and other disorders of cognition that are commonly encountered in clinical practice.



38 Sleep Disorders

Charles A. Czeisler, Thomas E. Scammell,
Clifford B. Saper

Disturbed sleep is among the most frequent health complaints that physicians encounter. More than one-half of adults in the United States experience at least intermittent sleep disturbance, and only 30% of adult Americans report consistently obtaining a sufficient amount of sleep. The Institute of Medicine has estimated that 50–70 million Americans suffer from a chronic disorder of sleep and wakefulness, which can adversely affect daytime functioning as well as physical and mental health. Over the last 20 years, the field of sleep medicine has emerged as a distinct specialty in response to the impact of sleep disorders and sleep deficiency on overall health.

PHYSIOLOGY OF SLEEP AND WAKEFULNESS

Given the opportunity, most healthy young adults will sleep 7–8 h per night, although the timing, duration, and internal structure of sleep vary among individuals. In the United States, adults tend to have one consolidated sleep episode each night, although in some cultures sleep may be divided into a mid-afternoon nap and a shortened night sleep. This pattern changes considerably over the life span, as infants and young children sleep considerably more than older people.

The stages of human sleep are defined on the basis of characteristic patterns in the electroencephalogram (EEG), the electrooculogram (EOG—a measure of eye-movement activity), and the surface electromyogram (EMG) measured on the chin, neck, and legs. The continuous recording of these electrophysiologic parameters to define sleep and wakefulness is termed *polysomnography*.

Polysomnographic profiles define two basic states of sleep: (1) rapid eye movement (REM) sleep and (2) non-rapid eye movement (NREM) sleep. NREM sleep is further subdivided into three stages: N1,

N2, and N3, characterized by increasing arousal threshold and slowing of the cortical EEG. REM sleep is characterized by a low-amplitude, mixed-frequency EEG similar to that of NREM stage N1 sleep. The EOG shows bursts of rapid eye movements similar to those seen during eyes-open wakefulness. EMG activity is absent in nearly all skeletal muscles, reflecting the brainstem-mediated muscle atonia that is characteristic of REM sleep.

ORGANIZATION OF HUMAN SLEEP

Normal nocturnal sleep in adults displays a consistent organization from night to night (Fig. 38-1). After sleep onset, sleep usually progresses through NREM stages N1–N3 sleep within 45–60 min. NREM stage N3 sleep (also known as slow-wave sleep) predominates in the first third of the night and comprises 15–25% of total nocturnal sleep time in young adults. Sleep deprivation increases the rapidity of sleep onset and both the intensity and amount of slow-wave sleep.

The first REM sleep episode usually occurs in the second hour of sleep. NREM and REM sleep alternate through the night with an average period of 90–110 min (the “ultradian” sleep cycle). Overall, in a healthy young adult, REM sleep constitutes 20–25% of total sleep, and NREM stages N1 and N2 constitute 50–60%.

Age has a profound impact on sleep state organization (Fig. 38-1). N3 sleep is most intense and prominent during childhood, decreasing with puberty and across the second and third decades of life. N3 sleep declines during adulthood to the point where it may be completely absent in older adults. The remaining NREM sleep becomes more fragmented, with many more frequent awakenings from NREM sleep. It is the increased frequency of awakenings, rather than a decreased ability to fall back asleep, that accounts for the increased wakefulness during the sleep episode in older people. While REM sleep may account for 50% of total sleep time in infancy, the percentage falls off sharply over the first postnatal year as a mature REM-NREM cycle develops; thereafter, REM sleep occupies about 25% of total sleep time.

Sleep deprivation degrades cognitive performance, particularly on tests that require continual vigilance. Paradoxically, older people are less vulnerable to the neurobehavioral performance impairment induced by acute sleep deprivation than young adults, maintaining their reaction time and sustaining vigilance with fewer lapses of attention. However, it is more difficult for older adults to obtain recovery sleep after staying awake all night, as the ability to sleep during the daytime declines with age.

After sleep deprivation, NREM sleep is generally recovered first, followed by REM sleep. However, because REM sleep tends to be most prominent in the second half of the night, sleep truncation (e.g., by an alarm clock) results in selective REM sleep deprivation. This may

increase REM sleep pressure to the point where the first REM sleep may occur much earlier in the nightly sleep episode. Because several disorders (see below) also cause sleep fragmentation, it is important that the patient have sufficient sleep opportunity (at least 8 h per night) for several nights prior to a diagnostic polysomnogram.

There is growing evidence that sleep deficiency in humans may cause glucose intolerance and contribute to the development of diabetes, obesity, and the metabolic syndrome, as well as impaired immune responses, accelerated atherosclerosis, and increased risk of cardiac disease and stroke. For these reasons, the Institute of Medicine declared sleep deficiency and sleep disorders “an unmet public health problem.”

WAKE AND SLEEP ARE REGULATED BY BRAIN CIRCUITS

Two principal neural systems govern the expression of the sleep and wakefulness. The ascending arousal system, illustrated in green in Fig. 38-2, consists of clusters of nerve cells extending from the upper pons to the hypothalamus and basal forebrain that activate the cerebral cortex, thalamus (which is necessary to relay sensory information to the cortex), and other forebrain regions. The ascending arousal neurons use monoamines (norepinephrine, dopamine, serotonin, and histamine), glutamate, or acetylcholine as neurotransmitters to activate their target neurons. Additional arousal-promoting neurons in the hypothalamus use the peptide neurotransmitter orexin (also known as hypocretin, shown in blue) to reinforce activity in the other arousal cell groups.

Damage to the arousal system at the level of the rostral pons and lower midbrain causes coma, indicating that the ascending arousal influence from this level is critical in maintaining wakefulness. Damage to the hypothalamic branch of the arousal system causes profound sleepiness, but usually not coma. Specific loss of the orexin neurons produces the sleep disorder narcolepsy (see below).

The arousal system is turned off during sleep by inhibitory inputs from cell groups in the sleep-promoting system, shown in Fig. 38-2 in red. These neurons in the preoptic area, lateral hypothalamus, and pons use γ -aminobutyric acid (GABA) to inhibit the arousal system. Many sleep-promoting neurons are themselves inhibited by inputs from the arousal system. This mutual inhibition between the arousal and sleep-promoting systems forms a neural circuit akin to what electrical engineers call a “flip-flop switch.” A switch of this type tends to promote rapid transitions between the on (wake) and off (sleep) states, while avoiding intermediate states. The relatively rapid transitions between waking and sleeping states, as seen in the EEG of humans and animals, is consistent with this model.

Neurons in the ventrolateral preoptic nucleus, one of the key sleep-promoting sites, are lost during normal human aging, correlating with reduced ability to maintain sleep (sleep fragmentation). The ventrolateral preoptic neurons are also injured in Alzheimer’s disease, which may in part account for the poor sleep quality in those patients.

Transitions between NREM and REM sleep appear to be governed by a similar switch in the brainstem. GABAergic REM-Off neurons have been identified in the lower midbrain that inhibit REM-On neurons in the upper pons. The REM-On group contains both GABAergic neurons that inhibit the REM-Off group (thus satisfying the conditions for a REM flip-flop switch) as well as glutamatergic neurons that project widely in the central nervous system (CNS) to cause the key phenomena associated with REM sleep. REM-On neurons that project to the medulla and spinal cord activate inhibitory (GABA and glycine-containing) interneurons, which in turn hyperpolarize the motor neurons, producing the atonia of REM sleep. REM-On neurons that project to the forebrain may be important in producing dreams.

The REM sleep switch receives cholinergic input, which favors transitions to REM sleep, and monoaminergic (norepinephrine and serotonin) input that prevents REM sleep. As a result, drugs that

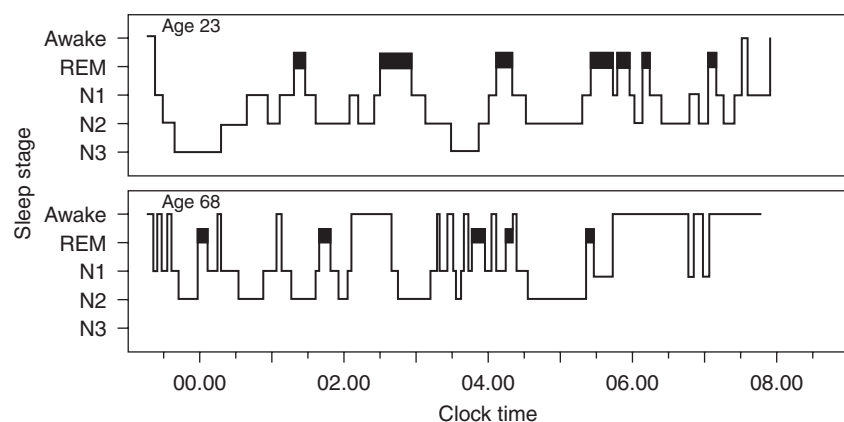


FIGURE 38-1 Wake-sleep architecture. Alternating stages of wakefulness, the three stages of NREM sleep (N1–N3), and REM sleep (solid bars) occur over the course of the night for representative young and older adult men. Characteristic features of sleep in older people include reduction of N3 slow-wave sleep, frequent spontaneous awakenings, early sleep onset, and early morning awakening. NREM, non-rapid eye movement; REM, rapid eye movement. (From the *Division of Sleep and Circadian Disorders, Brigham and Women’s Hospital*.)

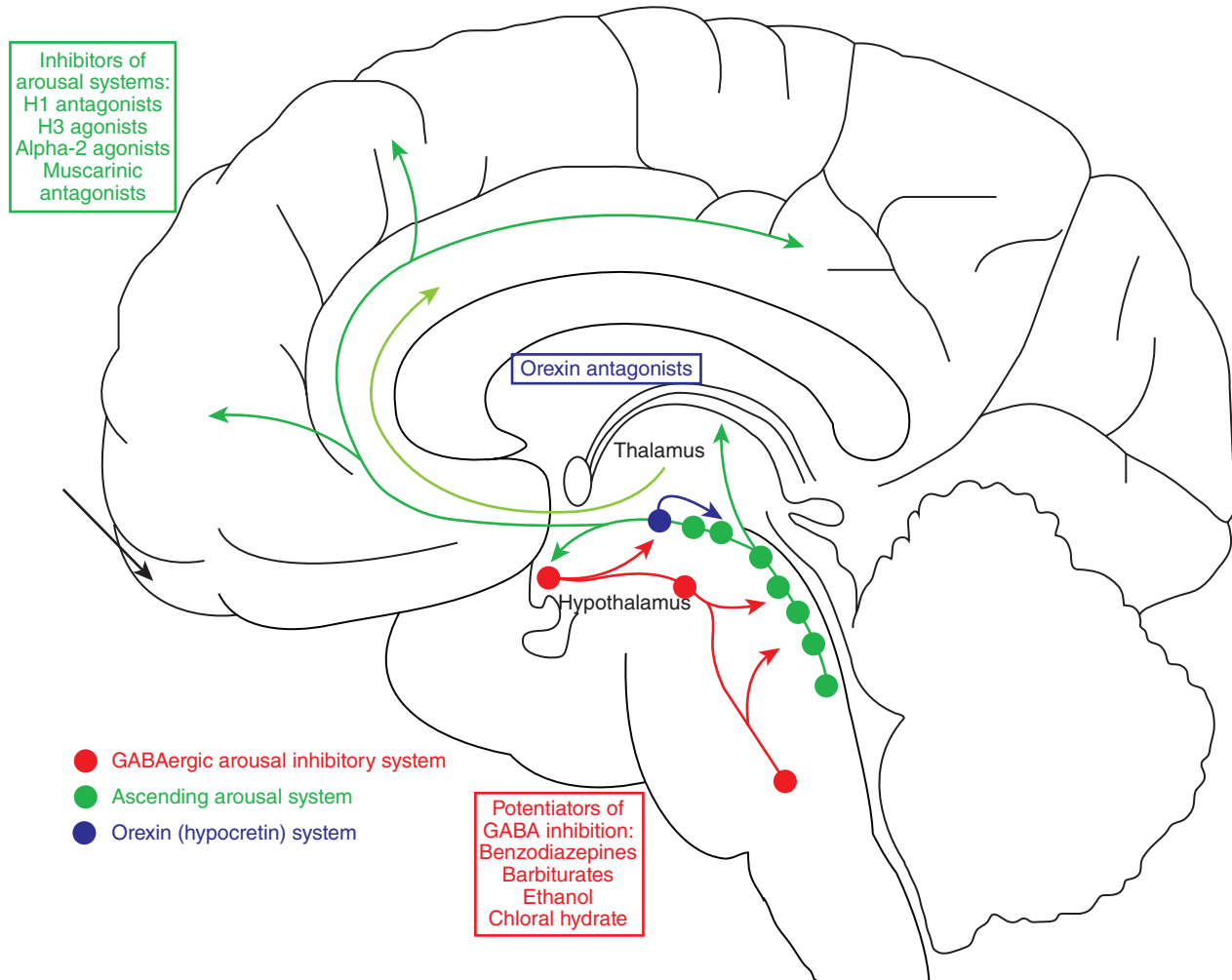


FIGURE 38-2 Relationship of drugs for insomnia with wake-sleep systems. The arousal system in the brain (*green*) includes monoaminergic, glutamatergic, and cholinergic neurons in the brainstem that activate neurons in the hypothalamus, thalamus, basal forebrain, and cerebral cortex. Orexin neurons (*blue*) in the hypothalamus, which are lost in narcolepsy, reinforce and stabilize arousal by activating other components of the arousal system. The sleep-promoting system (*red*) consists of GABAergic neurons in the preoptic area, lateral hypothalamus, and brainstem that inhibit the components of the arousal system, thus allowing sleep to occur. Drugs used to treat insomnia include those that block the effects of arousal system neurotransmitters (*green and blue*) and those that enhance the effects of γ -aminobutyric acid (GABA) produced by the sleep system (*red*).

increase monoamine tone (e.g., serotonin or norepinephrine reuptake inhibitors) tend to reduce the amount of REM sleep. Damage to the neurons that promote REM sleep atonia can produce REM sleep behavior disorder, a condition in which patients act out their dreams (see below).

SLEEP-WAKE CYCLES ARE DRIVEN BY HOMEOSTATIC, ALLOSTATIC, AND CIRCADIAN INPUTS

The gradual increase in sleep drive with prolonged wakefulness, followed by deeper slow-wave sleep and prolonged sleep episodes, demonstrates that there is a *homeostatic* mechanism that regulates sleep. The neurochemistry of sleep homeostasis is only partially understood, but with prolonged wakefulness, adenosine levels rise in parts of the brain. Adenosine may act through A₁ receptors to directly inhibit many arousal-promoting brain regions. In addition, adenosine promotes sleep through A_{2a} receptors; inhibition of these receptors by caffeine is one of the chief ways in which people fight sleepiness. Other humoral factors, such as prostaglandin D₂, have also been implicated in this process. Both adenosine and prostaglandin D₂ activate the sleep-promoting neurons in the ventrolateral preoptic nucleus.

Allostasis is the physiological response to a threat that cannot be managed by homeostatic mechanisms (e.g., the presence of physical danger or psychological threat). These stress responses can severely impact the need for and ability to sleep. For example, insomnia is very common

in patients with anxiety and other psychiatric disorders. Stress-induced insomnia is even more common, affecting most people at some time in their lives. Positron emission tomography (PET) studies in patients with chronic insomnia show hyperactivation of the components of the ascending arousal system, as well as their targets in the limbic system in the forebrain (e.g., cingulate cortex and amygdala). The limbic areas are not only targets for the arousal system, but they also send excitatory outputs back to the arousal system, which contributes to a vicious cycle of anxiety about wakefulness that makes it more difficult to sleep. Approaches to treating insomnia rely on drugs that either inhibit the output of the ascending arousal system (*green and blue* in Fig. 38-2) or potentiate the output of the sleep-promoting system (*red* in Fig. 38-2). However, behavioral approaches (cognitive behavioral therapy and sleep hygiene) that may reduce forebrain limbic activity at bedtime are often equally or more successful.

Sleep is also regulated by a strong *circadian* timing signal, driven by the suprachiasmatic nuclei (SCN) of the hypothalamus, as described below. The SCN sends outputs to key sites in the hypothalamus, which impose 24-h rhythms on a wide range of behaviors and body systems, including the wake-sleep cycle.

PHYSIOLOGY OF CIRCADIAN RHYTHMICITY

The wake-sleep cycle is the most evident of many 24-h rhythms in humans. Prominent daily variations also occur in endocrine,

thermoregulatory, cardiac, pulmonary, renal, immune, gastrointestinal, and neurobehavioral functions. At the molecular level, endogenous circadian rhythmicity is driven by self-sustaining transcriptional/translational feedback loops. In evaluating daily rhythms in humans, it is important to distinguish between diurnal components passively evoked by periodic environmental or behavioral changes (e.g., the increase in blood pressure and heart rate that occurs upon assumption of the upright posture) and circadian rhythms actively driven by an endogenous oscillatory process (e.g., the circadian variations in adrenal cortisol and pineal melatonin secretion that persist across a variety of environmental and behavioral conditions).

While it is now recognized that most cells in the body have circadian clocks that regulate diverse physiologic processes, most of these disparate clocks are unable to maintain the synchronization with each other that is required to produce useful 24-h rhythms aligned with the external light-dark cycle. The neurons in the SCN are interconnected with one another in such a way as to produce a near-24-h synchronous rhythm of neural activity that is then transmitted to the rest of the body. Bilateral destruction of the SCN results in a loss of most endogenous circadian rhythms including wake-sleep behavior and rhythms in endocrine and metabolic systems. The genetically determined period of this endogenous neural oscillator, which averages ~24.15 h in humans, is normally synchronized to the 24-h period of the environmental light-dark cycle through direct input from intrinsically photosensitive ganglion cells in the retina to the SCN. Humans are exquisitely sensitive to the resetting effects of light, particularly the shorter wavelengths (~460–500 nm) of the visible spectrum. Small differences in circadian period contribute to variations in diurnal preference in young adults (with the circadian period shorter in those who typically go to bed and rise earlier compared to those who typically go to bed and wake up later), whereas changes in homeostatic sleep regulation may underlie the age-related tendency toward earlier sleep-wake timing.

The timing and internal architecture of sleep are directly coupled to the output of the endogenous circadian pacemaker. Paradoxically, the endogenous circadian rhythm for wake propensity peaks just before the habitual bedtime, whereas that of sleep propensity peaks near the habitual wake time. These rhythms are thus timed to oppose the rise of sleep tendency throughout the usual waking day and the decline of sleep propensity during the habitual sleep episode, respectively. Misalignment of the endogenous circadian pacemaker with the desired wake-sleep cycle can, therefore, induce insomnia, decreased alertness, and impaired performance evident in night-shift workers and airline travelers.

BEHAVIORAL AND PHYSIOLOGIC CORRELATES OF SLEEP STATES AND STAGES

Polysomnographic staging of sleep correlates with behavioral changes during specific states and stages. During the transitional state (stage N1) between wakefulness and deeper sleep, individuals may respond to faint auditory or visual signals. Formation of short-term memories is inhibited at the onset of NREM stage N1 sleep, which may explain why individuals aroused from that transitional sleep stage frequently lack situational awareness. After sleep deprivation, such transitions may intrude upon behavioral wakefulness notwithstanding attempts to remain continuously awake (see “Shift-Work Disorder,” below).

Awakenings from REM sleep are associated with recall of vivid dream imagery over 80% of the time, especially later in the night. Imagery may also be reported after NREM sleep interruptions. Certain disorders may occur during specific sleep stages and are described below under “Parasomnias.” These include sleepwalking, night terrors, and enuresis (bed wetting), which occur most commonly in children during deep (N3) NREM sleep, and REM sleep behavior disorder, which occurs mainly among older men who fail to maintain full atonia during REM sleep, and often call out, thrash around, or even act out entire dreams.

All major physiologic systems are influenced by sleep. Blood pressure and heart rate decrease during NREM sleep, particularly during N3 sleep. During REM sleep, bursts of eye movements are associated with large variations in both blood pressure and heart rate mediated by the autonomic nervous system. Cardiac dysrhythmias may occur

selectively during REM sleep. Respiratory function also changes. In comparison to relaxed wakefulness, respiratory rate becomes slower but more regular during NREM sleep (especially N3 sleep) and becomes irregular during bursts of eye movements in REM sleep. Decreases in minute ventilation during NREM sleep are out of proportion to the decrease in metabolic rate, resulting in a slightly higher P_{CO_2} .

Endocrine function also varies with sleep. N3 sleep is associated with secretion of growth hormone in men, while sleep in general is associated with augmented secretion of prolactin in both men and women. Sleep has a complex effect on the secretion of luteinizing hormone (LH): during puberty, sleep is associated with increased LH secretion, whereas sleep in the postpubertal female inhibits LH secretion in the early follicular phase of the menstrual cycle. Sleep onset (and probably N3 sleep) is associated with inhibition of thyroid-stimulating hormone and of the adrenocorticotrophic hormone–cortisol axis, an effect that is superimposed on the prominent circadian rhythms in the two systems.

The pineal hormone melatonin is secreted predominantly at night in both day- and night-active species, reflecting the direct modulation of pineal activity by a circuitous neural pathway that links the SCN to the sympathetic nervous system, which innervates the pineal gland. Melatonin secretion does not require sleep, but melatonin secretion is inhibited by ambient light, an effect mediated by the neural connection from the retina to the pineal gland via the SCN. Sleep efficiency is highest when the sleep episode coincides with endogenous melatonin secretion. Administration of exogenous melatonin can hasten sleep onset and increase sleep efficiency when administered at a time when endogenous melatonin levels are low, such as in the afternoon or evening or at the desired bedtime in patients with delayed sleep-wake phase disorder, but it does not increase sleep efficiency if administered when endogenous melatonin levels are elevated. This may explain why melatonin is often ineffective in the treatment of patients with primary insomnia.

Sleep is accompanied by alterations of thermoregulatory function. NREM sleep is associated with an increase in the firing of warm-responsive neurons in the preoptic area and a fall in body temperature; conversely, skin warming without increasing core body temperature has been found to increase NREM sleep. REM sleep is associated with reduced thermoregulatory responsiveness.

DISORDERS OF SLEEP AND WAKEFULNESS

APPROACH TO THE PATIENT: Sleep Disorders

Patients may seek help from a physician because of: (1) sleepiness or tiredness during the day; (2) difficulty initiating or maintaining sleep at night (insomnia); or (3) unusual behaviors during sleep itself (parasomnias).

Obtaining a careful history is essential. In particular, the duration, severity, and consistency of the symptoms are important, along with the patient's estimate of the consequences of the sleep disorder on waking function. Information from a bed partner or family member is often helpful because some patients may be unaware of symptoms such as heavy snoring or may underreport symptoms such as falling asleep at work or while driving. Physicians should inquire about when the patient typically goes to bed, when they fall asleep and wake up, whether they awaken during sleep, whether they feel rested in the morning, and whether they nap during the day. Depending on the primary complaint, it may be useful to ask about snoring, witnessed apneas, restless sensations in the legs, movements during sleep, depression, anxiety, and behaviors around the sleep episode. The physical exam may provide evidence of a small airway, large tonsils, or a neurologic or medical disorder that contributes to the main complaint.

It is important to remember that, rarely, seizures may occur exclusively during sleep, mimicking a primary sleep disorder; such sleep-related seizures typically occur during episodes of NREM sleep and may take the form of generalized tonic-clonic movements

(sometimes with urinary incontinence or tongue biting) or stereotyped movements in partial complex epilepsy (**Chap. 445**).

It is often helpful for the patient to complete a daily sleep log for 1–2 weeks to define the timing and amounts of sleep. When relevant, the log can also include information on levels of alertness, work times, and drug and alcohol use, including caffeine and hypnotics.

Polysomnography is necessary for the diagnosis of several disorders such as sleep apnea, narcolepsy, and periodic limb movement disorder. A conventional polysomnogram performed in a clinical sleep laboratory allows measurement of sleep stages, respiratory effort and airflow, oxygen saturation, limb movements, heart rhythm, and additional parameters. A home sleep test usually focuses on just respiratory measures and is helpful in patients with a moderate to high likelihood of having obstructive sleep apnea. The multiple sleep latency test (MSLT) is used to measure a patient's propensity to sleep during the day and can provide crucial evidence for diagnosing narcolepsy and some other causes of sleepiness. The maintenance of wakefulness test is used to measure a patient's ability to sustain wakefulness during the daytime and can provide important evidence for evaluating the efficacy of therapies for improving sleepiness in conditions such as narcolepsy and obstructive sleep apnea.

EVALUATION OF DAYTIME SLEEPINESS

Up to 25% of the adult population has persistent daytime sleepiness that impairs an individual's ability to perform optimally in school, at work, while driving, and in other conditions that require alertness. Sleepy students often have trouble staying alert and performing well in school, and sleepy adults struggle to stay awake and focused on their work. More than half of Americans have fallen asleep while driving. An estimated 1.2 million motor vehicle crashes per year are due to drowsy drivers, causing about 20% of all serious crash injuries and deaths. One needn't fall asleep to have an accident, as the inattention and slowed responses of drowsy drivers are a major contributor. Reaction time is equally impaired by 24 h of sleep loss as by a blood alcohol concentration of 0.10 g/dL.

Identifying and quantifying sleepiness can be challenging. First, patients may describe themselves as “sleepy,” “fatigued,” or “tired,” and the meanings of these words may differ between patients. For clinical purposes, it is best to use the term “sleepiness” to describe a propensity to fall asleep; whereas “fatigue” is best used to describe a feeling of low physical or mental energy but without a tendency to actually sleep. Sleepiness is usually most evident when the patient is sedentary, whereas fatigue may interfere with more active pursuits. Sleepiness generally occurs with disorders that reduce the quality or quantity of sleep or that interfere with the neural mechanisms of arousal, whereas fatigue

is more common in inflammatory disorders such as cancer, multiple sclerosis (**Chap. 458**), fibromyalgia (**Chap. 396**), chronic fatigue syndrome (**Chap. 464e**), or endocrine deficiencies such as hypothyroidism (**Chap. 405**) or Addison's disease (**Chap. 406**). Second, sleepiness can affect judgment in a manner analogous to ethanol, such that patients may have limited insight into the condition and the extent of their functional impairment. Finally, patients may be reluctant to admit that sleepiness is a problem because they may have become unfamiliar with feeling fully alert and because sleepiness is sometimes viewed pejoratively as reflecting poor motivation or bad sleep habits.

Table 38-1 outlines the diagnostic and therapeutic approach to the patient with a complaint of excessive daytime sleepiness.

To determine the extent and impact of sleepiness on daytime function, it is helpful to ask patients about the occurrence of sleep episodes during normal waking hours, both intentional and unintentional. Specific areas to be addressed include the occurrence of inadvertent sleep episodes while driving or in other safety-related settings, sleepiness while at work or school (and the relationship of sleepiness to work and school performance), and the effect of sleepiness on social and family life. Standardized questionnaires such as the Epworth Sleepiness Scale are often used clinically to measure sleepiness.

Eliciting a history of daytime sleepiness is usually adequate, but objective quantification is sometimes necessary. The MSLT measures a patient's propensity to sleep under quiet conditions. The test is performed after an overnight polysomnogram to establish that the patient has had an adequate amount of good-quality nighttime sleep. The MSLT consists of five 20-min nap opportunities every 2 h across the day. The patient is instructed to try to fall asleep, and the major endpoints are the average latency to sleep and the occurrence of REM sleep during the naps. An average sleep latency across the naps of less than 8 min is considered objective evidence of excessive daytime sleepiness. REM sleep normally occurs only during the nighttime sleep episode, and the occurrence of REM sleep in two or more of the MSLT naps provides support for the diagnosis of narcolepsy.

For the safety of the individual and the general public, physicians have a responsibility to help manage issues around driving in patients with sleepiness. Legal reporting requirements vary from state to state, but at a minimum, physicians should inform sleepy patients about their increased risk of having an accident and advise such patients not to drive a motor vehicle until the sleepiness has been treated effectively. This discussion is especially important for professional drivers, and it should be documented in the patient's medical record.

INSUFFICIENT SLEEP

Insufficient sleep is probably the most common cause of excessive daytime sleepiness. The average adult needs 7.5–8 h of sleep, but on

TABLE 38-1 EVALUATION OF THE PATIENT WITH EXCESSIVE DAYTIME SLEEPINESS

Findings on History and Physical Examination	Diagnostic Evaluation	Diagnosis	Therapy
Difficulty waking in the morning, rebound sleep on weekends and vacations with improvement in sleepiness	Sleep log	Insufficient sleep	Sleep education and behavioral modification to increase amount of sleep
Obesity, snoring, hypertension	Polysomnogram	Obstructive sleep apnea (Chap. 319)	Continuous positive airway pressure; upper airway surgery (e.g., uvulopalatopharyngoplasty); dental appliance; weight loss
Cataplexy, hypnagogic hallucinations, sleep paralysis	Polysomnogram and multiple sleep latency test	Narcolepsy	Stimulants (e.g., modafinil, methylphenidate); REM sleep-suppressing antidepressants (e.g., venlafaxine); sodium oxybate
Restless legs, kicking movements during sleep	Assessment for predisposing medical conditions (e.g., iron deficiency or renal failure)	Restless legs syndrome with or without periodic limb movements	Treatment of predisposing condition if possible; dopamine agonists (e.g., pramipexole, ropinirole)
Sedating medications, stimulant withdrawal, head trauma, systemic inflammation, Parkinson's disease and other neurodegenerative disorders, hypothyroidism, encephalopathy	Thorough medical history and exam including detailed neurologic exam	Sleepiness due to a drug or medical condition	Change medications, treat underlying condition, consider stimulants

weeknights, the average U.S. adult gets only 6.75 h of sleep. Only 30% of the U.S. adult population reports consistently obtaining sufficient sleep. Insufficient sleep is especially common among shift workers, individuals working multiple jobs, and people in lower socioeconomic groups. Most teenagers need ≥ 9 h of sleep, but many fail to get enough sleep because of circadian phase delay, or social pressures to stay up late coupled with early school start times. Late evening light exposure, television viewing, video-gaming, social media, texting, and smartphone use often delay bedtimes despite the fixed, early wake times required for work or school. As is typical with any disorder that causes sleepiness, individuals with chronically insufficient sleep may feel inattentive, irritable, unmotivated, and depressed, and have difficulty with school, work, and driving. Individuals differ in their optimal amount of sleep, and it can be helpful to ask how much sleep the patient obtains on a quiet vacation when he or she can sleep without restrictions. Some patients may think that a short amount of sleep is normal or advantageous, and they may not appreciate their biological need for more sleep, especially if coffee and other stimulants mask the sleepiness. A 2-week sleep log documenting the timing of sleep and daily level of alertness is diagnostically useful and provides helpful feedback for the patient. Extending sleep to the optimal amount on a regular basis can resolve the sleepiness and other symptoms. As with any lifestyle change, extending sleep requires commitment and adjustments, but the improvements in daytime alertness make this change worthwhile.

SLEEP APNEA SYNDROMES

Respiratory dysfunction during sleep is a common, serious cause of excessive daytime sleepiness as well as of disturbed nocturnal sleep. At least 24% of middle-aged men and 9% of middle-aged women in the United States have a reduction or cessation of breathing dozens or more times each night during sleep, with 9% of men and 4% of women doing so more than a hundred times per night. These episodes may be due to an occlusion of the airway (*obstructive sleep apnea*), absence of respiratory effort (*central sleep apnea*), or a combination of these factors (*mixed sleep apnea*). Failure to recognize and treat these conditions appropriately may lead to impairment of daytime alertness, increased risk of sleep-related motor vehicle crashes, depression, hypertension, myocardial infarction, diabetes, stroke, and increased mortality. Sleep apnea is particularly prevalent in overweight men and in the elderly, yet it is estimated to go undiagnosed in most affected individuals. This is unfortunate because several effective treatments are available. **Readers are referred to Chap. 319 for a comprehensive review of the diagnosis and treatment of patients with sleep apnea.**

NARCOLEPSY

Narcolepsy is characterized by difficulty sustaining wakefulness, poor regulation of REM sleep, and disturbed nocturnal sleep. All patients with narcolepsy have excessive daytime sleepiness. This sleepiness is often severe, but in some, it is mild. In contrast to patients with disrupted sleep (e.g., sleep apnea), people with narcolepsy usually feel well rested upon awakening and then feel tired throughout much of the day. In addition, they often experience symptoms related to an intrusion of REM sleep characteristics. REM sleep is characterized by dreaming and muscle paralysis, and people with narcolepsy can have: (1) sudden muscle weakness without a loss of consciousness, which is usually triggered by strong emotions (cataplexy; **Video 38-1**); (2) dream-like hallucinations at sleep onset (hypnagogic hallucinations) or upon awakening (hypnopompic hallucinations); and (3) muscle paralysis upon awakening (sleep paralysis). With severe cataplexy, an individual may be laughing at a joke and then suddenly collapse to the ground, immobile but awake for 1–2 min. With milder episodes, patients may have mild weakness

of the face or neck. Narcolepsy is one of the more common causes of chronic sleepiness and affects about 1 in 2000 people in the United States. Narcolepsy typically begins between age 10 and 20; once established, the disease persists for life.

Narcolepsy is caused by loss of the hypothalamic neurons that produce the orexin neuropeptides (also known as hypocretins). Research in mice and dogs first demonstrated that a loss of orexin signaling due to null mutations of either the orexin neuropeptides or one of the orexin receptors causes sleepiness and cataplexy nearly identical to that seen in people with narcolepsy. Although genetic mutations rarely cause human narcolepsy, researchers soon discovered that patients with narcolepsy had very low or undetectable levels of orexins in their cerebrospinal fluid, and autopsy studies showed a nearly complete loss of the orexin-producing neurons in the hypothalamus. The orexins normally promote long episodes of wakefulness and suppress REM sleep, and thus, loss of orexin signaling results in frequent intrusions of sleep during the usual waking episode, with REM sleep and fragments of REM sleep at any time of day (**Fig. 38-3**).

Extensive evidence suggests that an autoimmune process likely causes this selective loss of the orexin-producing neurons. Certain human leukocyte antigens (HLAs) can increase the risk of autoimmune disorders (**Chap. 373e**), and narcolepsy has the strongest known HLA association. HLA DQB1*06:02 is found in about 90% of people with narcolepsy, whereas it occurs in only 12–25% of the general population. Researchers now hypothesize that in people with DQB1*06:02, an immune response against influenza, *Streptococcus*, or other infections may also damage the orexin-producing neurons through a process of molecular mimicry. This mechanism may account for the 8- to 12-fold increase in new cases of narcolepsy among children in Europe who received a particular brand of H1N1 influenza A vaccine (Pandemrix).

On rare occasions, narcolepsy can occur with neurologic disorders such as tumors or strokes that directly damage the orexin-producing neurons in the hypothalamus or their projections.

Diagnosis Narcolepsy is most commonly diagnosed by the history of chronic sleepiness plus cataplexy or other symptoms. Many disorders can cause feelings of weakness, but with true cataplexy, patients will describe definite functional weakness (e.g., slurred speech, dropping a cup, slumping into a chair) that has consistent emotional triggers such as heartfelt mirth when laughing at a great joke, happy surprise at unexpectedly seeing a friend, or intense anger. Cataplexy occurs in about half of all narcolepsy patients and is diagnostically very helpful because it occurs in almost no other disorder. In contrast, occasional hypnagogic hallucinations and sleep paralysis occur in about 20% of the general population, and these symptoms are not as diagnostically specific.

When narcolepsy is suspected, the diagnosis should be firmly established with a polysomnogram followed by an MSLT. The

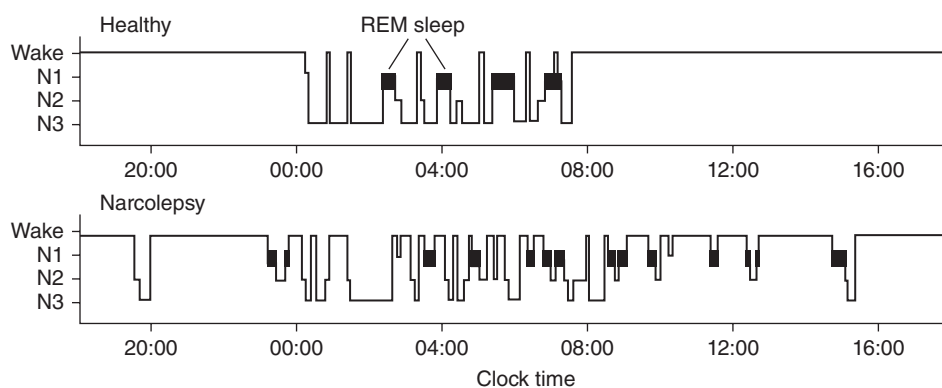


FIGURE 38-3 Polysomnographic recordings of a healthy individual and a patient with narcolepsy. The individual with narcolepsy enters rapid eye movement (REM) sleep quickly at night and has moderately fragmented sleep. During the day, the healthy subject stays awake from 8:00 AM until midnight, but the patient with narcolepsy dozes off frequently, with many daytime naps that include REM sleep.

polysomnogram helps rule out other possible causes of sleepiness such as sleep apnea, and the MSLT provides essential, objective evidence of sleepiness plus REM sleep dysregulation. Across the five naps of the MSLT, most patients with narcolepsy will fall asleep in less than 8 min on average, and they will have episodes of REM sleep in at least two of the naps. Abnormal regulation of REM sleep is also manifested by the appearance of REM sleep within 15 min of sleep onset at night, which is rare in healthy individuals sleeping at their habitual bedtime. Stimulants should be stopped 1 week before the MSLT and antidepressants should be stopped 3 weeks prior, because these medications can affect the MSLT. In addition, patients should be encouraged to obtain a fully adequate amount of sleep each night for the week prior to the test to eliminate any effects of insufficient sleep.

TREATMENT NARCOLEPSY

The treatment of narcolepsy is symptomatic. Most patients with narcolepsy feel more alert after sleep, and they should be encouraged to get adequate sleep each night and to take a 15- to 20-min nap in the afternoon. This nap may be sufficient for some patients with mild narcolepsy, but most also require treatment with wake-promoting medications. Modafinil is used quite often because it has fewer side effects than amphetamines and a relatively long half-life; for most patients, 200–400 mg each morning is very effective. Methylphenidate (10–20 mg bid) or dextroamphetamine (10 mg bid) are often effective, but sympathomimetic side effects, anxiety, and the potential for abuse can be concerns. These medications are available in slow-release formulations, extending their duration of action and allowing easier dosing. Sodium oxybate (gamma hydroxybutyrate) is given twice each night and is often very valuable in improving alertness, but it can produce excessive sedation, nausea, and confusion.

Cataplexy is usually much improved with antidepressants that increase noradrenergic or serotonergic tone because these medications strongly suppress REM sleep and cataplexy. Venlafaxine (37.5–150 mg each morning) and fluoxetine (10–40 mg each morning) are often quite effective. The tricyclic antidepressants, such as protriptyline (10–40 mg/d) or clomipramine (25–50 mg/d) are potent suppressors of cataplexy, but their anticholinergic effects, including sedation and dry mouth, make them less attractive.¹ Sodium oxybate, given at bedtime and 3–4 h later, is also very helpful in reducing cataplexy.

EVALUATION OF INSOMNIA

Insomnia is the complaint of poor sleep and usually presents as difficulty initiating or maintaining sleep. People with insomnia are dissatisfied with their sleep and feel that it impairs their ability to function well in work, school, and social situations. Affected individuals often experience fatigue, decreased mood, irritability, malaise, and cognitive impairment.

Chronic insomnia, lasting more than 3 months, occurs in about 10% of adults and is more common in women, older adults, people of lower socioeconomic status, and individuals with medical, psychiatric, and substance abuse disorders. Acute or short-term insomnia affects over 30% of adults and is often precipitated by stressful life events such as a major illness or loss, change of occupation, medications, and substance abuse. If the acute insomnia triggers maladaptive behaviors such as increased nocturnal light exposure, frequently checking the clock, or attempting to sleep more by napping, it can lead to chronic insomnia.

Most insomnia begins in adulthood, but many patients may be predisposed and report easily disturbed sleep predating the insomnia, suggesting that their sleep is lighter than usual. Clinical studies and animal models indicate that insomnia is associated with activation

during sleep of brain areas normally active only during wakefulness. The polysomnogram is rarely used in the evaluation of insomnia, as it typically confirms the patient's subjective report of long latency to sleep and numerous awakenings but usually adds little new information. Many patients with insomnia have increased fast (beta) activity in the EEG during sleep; this fast activity is normally present only during wakefulness, which may explain why some patients report feeling awake for much of the night. The MSLT is rarely used in the evaluation of insomnia because, despite their feelings of low energy, most people with insomnia do not easily fall asleep during the day, and on the MSLT, their average sleep latencies are usually longer than normal.

Many factors can contribute to insomnia, and obtaining a careful history is essential so one can select therapies targeting the underlying factors. The assessment should focus on identifying predisposing, precipitating, and perpetuating factors.

Psychophysiologic Factors Many patients with insomnia have negative expectations and conditioned arousal that interfere with sleep. These individuals may worry about their insomnia during the day and have increasing anxiety as bedtime approaches if they anticipate a poor night of sleep. While attempting to sleep, they may frequently check the clock, which only heightens anxiety and frustration. They may find it easier to sleep in a new environment rather than their bedroom, as it lacks the negative associations.

Inadequate Sleep Hygiene Patients with insomnia sometimes develop counterproductive behaviors that contribute to their insomnia. These can include daytime napping that reduces sleep drive at night; an irregular sleep-wake schedule that disrupts their circadian rhythms; use of wake-promoting substances (e.g., caffeine, tobacco) too close to bedtime; engaging in alerting or stressful activities close to bedtime (e.g., arguing with a partner, work-related emailing and texting while in bed, sleeping with a smartphone or tablet at the bedside); and routinely using the bedroom for activities other than sleep or sex (e.g., TV, work), so the bedroom becomes associated with arousing or stressful feelings.

Psychiatric Conditions About 80% of patients with psychiatric disorders have sleep complaints, and about half of all chronic insomnia occurs in association with a psychiatric disorder. Depression is classically associated with early morning awakening, but it can also interfere with the onset and maintenance of sleep. Mania and hypomania can disrupt sleep and often are associated with substantial reductions in the total amount of sleep. Anxiety disorders can lead to racing thoughts and rumination that interfere with sleep and can be very problematic if the patient's mind becomes active midway through the night. Panic attacks can occur during sleep and need to be distinguished from other parasomnias. Insomnia is common in schizophrenia and other psychoses, often resulting in fragmented sleep, less deep NREM sleep, and sometimes reversal of the day-night sleep pattern.

Medications and Drugs of Abuse A wide variety of psychoactive drugs can interfere with sleep. Caffeine, which has a half-life of 6–9 h, can disrupt sleep for up to 8–14 h, depending on the dose, variations in metabolism, and an individual's caffeine sensitivity. Insomnia can also result from use of prescription medications too close to bedtime (e.g., theophylline, stimulants, antidepressants, glucocorticoids). Conversely, withdrawal of sedating medications such as alcohol, narcotics, or benzodiazepines can cause insomnia. Alcohol taken just before bed can shorten sleep latency, but it often produces rebound insomnia 2–3 h later as it wears off. This same problem with sleep maintenance can occur with short-acting benzodiazepines such as alprazolam.

Medical Conditions A large number of medical conditions disrupt sleep. Pain from rheumatologic disorders or a painful neuropathy commonly disrupts sleep. Some patients may sleep poorly because of respiratory conditions such as asthma, chronic obstructive pulmonary disease, cystic fibrosis, congestive heart failure, or restrictive lung disease, and some of these disorders are worse at night in bed due to circadian variations in airway resistance and postural changes that can result in

¹No antidepressant has been approved by the U.S. Food and Drug Administration (FDA) for treating narcolepsy.

paroxysmal nocturnal dyspnea. Many women experience poor sleep with the hormonal changes of menopause. Gastroesophageal reflux is also a common cause of difficulty sleeping.

Neurologic Disorders Dementia (Chap. 35) is often associated with poor sleep, probably due to a variety of factors, including napping during the day, altered circadian rhythms, and perhaps a weakened output of the brain's sleep-promoting mechanisms. In fact, insomnia and nighttime wandering are some of the most common causes for institutionalization of patients with dementia, because they place a larger burden on caregivers. Conversely, in cognitively intact elderly men, fragmented sleep and poor sleep quality are associated with subsequent cognitive decline. Patients with Parkinson's disease may sleep poorly due to rigidity, dementia, and other factors. Fatal familial insomnia is a very rare neurodegenerative condition caused by mutations in the prion protein gene, and although insomnia is a common early symptom, most patients present with other obvious neurologic signs such as dementia, myoclonus, dysarthria, or autonomic dysfunction.

TREATMENT INSOMNIA

Treatment of insomnia improves quality of life and can promote long-term health. With improved sleep, patients often report less daytime fatigue, improved cognition, and more energy. Treating the insomnia can also improve the comorbid disease. For example, management of insomnia at the time of diagnosis of major depression often improves the response to antidepressants and reduces the risk of relapse. Sleep loss can heighten the perception of pain, so a similar approach is warranted in acute and chronic pain management.

The treatment plan should target all putative contributing factors: establish good sleep hygiene, treat medical disorders, use behavioral therapies for anxiety and negative conditioning, and use pharmacotherapy and/or psychotherapy for psychiatric disorders. Behavioral therapies should be the first-line treatment, followed by judicious use of sleep-promoting medications if needed.

TREATMENT OF MEDICAL AND PSYCHIATRIC DISEASE

If the history suggests that a medical or psychiatric disease contributes to the insomnia, then it should be addressed by, for example, treating the pain, improving breathing, and switching or adjusting the timing of medications.

IMPROVE SLEEP HYGIENE

Attention should be paid to improving sleep hygiene and avoiding counterproductive, arousing behaviors before bedtime. Patients should establish a regular bedtime and wake time, even on weekends, to help synchronize their circadian rhythms and sleep patterns. The amount of time allocated for sleep should not be more than their actual total amount of sleep. In the 30 min before bedtime, patients should establish a relaxing "wind-down" routine that can include a warm bath, listening to music, meditation, or other relaxation techniques. The bedroom should be off-limits to computers, televisions, radios, smartphones, videogames, and tablets. Once in bed, patients should try to avoid thinking about anything stressful or arousing such as problems with relationships or work. If they cannot fall asleep within 20 min, it often helps to get out of bed and read or listen to relaxing music in dim light as a form of distraction from any anxiety, but artificial light, including light from a television, cell phone, or computer, should be avoided, because light itself suppresses melatonin secretion and is arousing.

Table 38-2 outlines some of the key aspects of good sleep hygiene to improve insomnia.

COGNITIVE BEHAVIORAL THERAPY (CBT)

CBT uses a combination of the techniques above plus additional methods to improve insomnia. A trained therapist may use cognitive psychology techniques to reduce excessive worrying about sleep and to reframe faulty beliefs about the insomnia and its daytime consequences. The therapist may also teach the patient relaxation

TABLE 38-2 METHODS TO IMPROVE SLEEP HYGIENE IN INSOMNIA PATIENTS

Helpful Behaviors	Behaviors to Avoid
Use the bed only for sleep and sex	Avoid behaviors that interfere with sleep physiology, including:
<ul style="list-style-type: none"> If you cannot sleep within 20 min, get out of bed and read or do other relaxing activities in dim light before returning to bed 	<ul style="list-style-type: none"> Napping, especially after 3:00 PM Attempting to sleep too early Caffeine after lunchtime
Make quality sleep a priority	In the 2–3 h before bedtime, avoid:
<ul style="list-style-type: none"> Go to bed and get up at the same time each day Ensure a restful environment (comfortable bed, bedroom quiet and dark) 	<ul style="list-style-type: none"> Heavy eating Smoking or alcohol Vigorous exercise
Develop a consistent bedtime routine. For example:	When trying to fall asleep, avoid:
<ul style="list-style-type: none"> Prepare for sleep with 20–30 min of relaxation (e.g., soft music, meditation, yoga, pleasant reading) Take a warm bath 	<ul style="list-style-type: none"> Solving problems Thinking about life issues Reviewing events of the day

techniques, such as progressive muscle relaxation or meditation, to reduce autonomic arousal, intrusive thoughts, and anxiety.

MEDICATIONS FOR INSOMNIA

If insomnia persists after treatment of these contributing factors, pharmacotherapy is often used on a nightly or intermittent basis. A variety of sedatives can improve sleep.

Antihistamines, such as diphenhydramine, are the primary active ingredient in most over-the-counter sleep aids. These may be of benefit when used intermittently, but often produce rapid tolerance and can produce anticholinergic side effects such as dry mouth and constipation, which limit their use, particularly in the elderly.

Benzodiazepine receptor agonists (BzRAs) are an effective and well-tolerated class of medications for insomnia. BzRAs bind to the GABA_A receptor and potentiate the postsynaptic response to GABA. GABA_A receptors are found throughout the brain, and BzRAs may globally reduce neural activity and may enhance the activity of specific sleep-promoting GABAergic pathways. Classic BzRAs include lorazepam, triazolam, and clonazepam, whereas newer agents such as zolpidem and zaleplon have more selective affinity for the α_1 subunit of the GABA_A receptor.

Specific BzRAs are often chosen based on the desired duration of action. The most commonly prescribed agents in this family are zaleplon (5–20 mg), with a half-life of 1–2 h; zolpidem (5–10 mg) and triazolam (0.125–0.25 mg), with half-lives of 2–4 h; eszopiclone (1–3 mg), with a half-life of 5–8 h; and temazepam (15–30 mg), with a half-life of 8–20 h. Generally, side effects are minimal when the dose is kept low and the serum concentration is minimized during the waking hours (by using the shortest-acting effective agent). For chronic insomnia, intermittent use is recommended, unless the consequences of untreated insomnia outweigh concerns regarding chronic use.

The heterocyclic *antidepressants* (trazodone, amitriptyline,² and doxepin) are the most commonly prescribed alternatives to BzRAs due to their lack of abuse potential and lower cost. Trazodone (25–100 mg) is used more commonly than the tricyclic antidepressants, because it has a much shorter half-life (5–9 h) and less anticholinergic activity.

Medications for insomnia are now among the most commonly prescribed medications, but they should be used cautiously. All sedatives increase the risk of injurious falls and confusion in the elderly, and therefore if needed, these medications should be used at the lowest effective dose. Morning sedation can interfere with driving and judgment, and when selecting a medication, one should

²Trazodone and amitriptyline have not been approved by the FDA for treating insomnia.

consider the duration of action. Benzodiazepines carry a risk of addiction and abuse, especially in patients with a history of alcohol or sedative abuse. Like alcohol, some sleep-promoting medications can worsen sleep apnea. Sedatives can also produce complex behaviors during sleep, such as sleep walking and sleep eating, although this seems more likely at higher doses.

RESTLESS LEGS SYNDROME

Patients with restless legs syndrome (RLS) report an irresistible urge to move the legs. Many patients report a creepy-crawly or unpleasant deep ache within the thighs or calves, and those with more severe RLS may have discomfort in the arms as well. For most patients with RLS, these dysesthesias and restlessness are much worse in the evening and first half of the night. The symptoms appear with inactivity and can make sitting still in an airplane or when watching a movie a miserable experience. The sensations are temporarily relieved by movement, stretching, or massage. This nocturnal discomfort usually interferes with sleep, and patients may report daytime sleepiness as a consequence. RLS is very common, affecting 5–10% of adults and is more common in women and older adults.

A variety of factors can cause RLS. Iron deficiency is the most common treatable cause, and iron replacement should be considered if the ferritin level is less than 50 ng/mL. RLS can also occur with peripheral neuropathies and uremia and can be worsened by pregnancy, caffeine, alcohol, antidepressants, lithium, neuroleptics, and antihistamines. Genetic factors contribute to RLS, and polymorphisms in a variety of genes (*BTBD9*, *MEIS1*, *MAP2K5/LBXCOR*, and *PTPRD*) have been linked to RLS, although as yet, the mechanism through which they cause RLS remains unknown. Roughly one-third of patients (particularly those with an early age of onset) have multiple affected family members.

RLS is treated by addressing the underlying cause such as iron deficiency if present. Otherwise, treatment is symptomatic, and dopamine agonists are used most frequently. Agonists of dopamine $D_{2/3}$ receptors such as pramipexole (0.25–0.5 mg q7PM) or ropinirole (0.5–4 mg q7PM) are considered first-line agents. Augmentation is a worsening of RLS such that symptoms begin earlier in the day and can spread to other body regions, and it can occur in about 25% of patients taking dopamine agonists. Other possible side effects of dopamine agonists include nausea, morning sedation, and increases in rewarding behavior such as gambling and sex. Opioids, benzodiazepines, pregabalin, and gabapentin may also be of therapeutic value. Most patients with restless legs also experience periodic limb movement disorder, although the reverse is not the case.

PERIODIC LIMB MOVEMENT DISORDER

Periodic limb movement disorder (PLMD) involves rhythmic twitches of the legs that disrupt sleep. The movements resemble a triple flexion reflex with extensions of the great toe and dorsiflexion of the foot for 0.5 to 5.0 s, which recur every 20–40 s during NREM sleep, in episodes lasting from minutes to hours. PLMD is diagnosed by a polysomnogram that includes recordings of the anterior tibialis and sometimes other muscles. The EEG shows that the movements of PLMD frequently cause brief arousals that disrupt sleep and can cause insomnia and daytime sleepiness. PLMD can be caused by the same factors that cause RLS (see above), and the frequency of leg movements improves with the same medications as used for RLS, including dopamine agonists. Recent genetic studies identified polymorphisms associated with RLS/PLMD, suggesting that they may have a common pathophysiology.

PARASOMNIAS

Parasomnias are abnormal behaviors or experiences that arise from or occur during sleep. A variety of parasomnias can occur during NREM sleep, from brief confusional arousals to sleepwalking and night terrors. The presenting complaint is usually related to the behavior itself, but the parasomnias can disturb sleep continuity or lead to mild impairments in daytime alertness. Two main parasomnias occur in REM sleep: REM sleep behavior disorder (RBD) and nightmares.

Sleepwalking (Somnambulism) Patients affected by this disorder carry out automatic motor activities that range from simple to complex. Individuals may walk, urinate inappropriately, eat, exit the house, or drive a car with minimal awareness. Full arousal may be difficult, and occasional individuals may respond to attempted awakening with agitation or violence. Sleepwalking arises from NREM stage N3 sleep, usually in the first few hours of the night, and the EEG usually shows the slow cortical activity of deep NREM sleep even when the patient is moving about. Sleepwalking is most common in children and adolescents, when these sleep stages are most robust. About 15% of children have occasional sleepwalking, and it persists in about 1% of adults. Episodes are usually isolated but may be recurrent in 1–6% of patients. The cause is unknown, although it has a familial basis in roughly one-third of cases. Sleepwalking can be worsened by insufficient sleep, which subsequently causes an increase in deep NREM sleep; alcohol; and stress. These should be addressed if present. Small studies have shown some efficacy of antidepressants and benzodiazepines; relaxation techniques and hypnosis can also be helpful. Patients and their families should improve home safety (e.g., replace glass doors, remove low tables to avoid tripping) to minimize the chance of injury if sleepwalking occurs.

Sleep Terrors This disorder occurs primarily in young children during the first few hours of sleep during NREM stage N3 sleep. The child often sits up during sleep and screams, exhibiting autonomic arousal with sweating, tachycardia, large pupils, and hyperventilation. The individual may be difficult to arouse and rarely recalls the episode on awakening in the morning. Treatment usually consists of reassuring the parents that the condition is self-limited and benign, and like sleepwalking, it may improve by avoiding insufficient sleep.

Sleep Bruxism Bruxism is an involuntary, forceful grinding of teeth during sleep that affects 10–20% of the population. The patient is usually unaware of the problem. The typical age of onset is 17–20 years, and spontaneous remission usually occurs by age 40. Sex distribution appears to be equal. In many cases, the diagnosis is made during dental examination, damage is minor, and no treatment is indicated. In more severe cases, treatment with a tooth guard is necessary to prevent tooth injury. Stress management or, in some cases, biofeedback can be useful when bruxism is a manifestation of psychological stress. There are anecdotal reports of benefit with benzodiazepines.

Sleep Enuresis Bedwetting, like sleepwalking and night terrors, is another parasomnia that occurs during sleep in the young. Before age 5 or 6 years, nocturnal enuresis should be considered a normal feature of development. The condition usually improves spontaneously by puberty, has a prevalence in late adolescence of 1–3%, and is rare in adulthood. Treatment consists of bladder training exercises and behavioral therapy. Symptomatic pharmacotherapy is usually accomplished in adults with desmopressin (0.2 mg qhs), oxybutynin chloride (5 mg qhs), or imipramine (10–25 mg qhs). Important causes of nocturnal enuresis in patients who were previously continent for 6–12 months include urinary tract infections or malformations, cauda equina lesions, emotional disturbances, epilepsy, sleep apnea, and certain medications.

REM Sleep Behavior Disorder (RBD) RBD ([Video 38-2](#)) is distinct from other parasomnias in that it occurs during REM sleep. The patient or the bed partner usually reports agitated or violent behavior during sleep, and upon awakening, the patient can often report a dream that accompanied the movements. During normal REM sleep, nearly all skeletal muscles are paralyzed, but in patients with RBD, the polysomnogram often shows limb movements during REM sleep, lasting for seconds to minutes. The movements can be dramatic, and it is not uncommon for the patient or the bed partner to be injured.

RBD primarily afflicts older men, and most either have or will develop a neurodegenerative disorder. In longitudinal studies of RBD, half of the patients developed a synucleinopathy such as Parkinson's disease ([Chap. 449](#)) or dementia with Lewy bodies ([Chap. 448](#)), or occasionally multiple system atrophy ([Chap. 454](#)), within 12 years, and over 80% developed a synucleinopathy by 20 years. RBD can occur

in patients taking antidepressants, and in some, these medications may unmask this early indicator of neurodegeneration. Synucleinopathies probably cause neuronal loss in brainstem regions that regulate muscle atonia during REM sleep, and loss of these neurons permits movements to break through during REM sleep. RBD also occurs in about 30% of patients with narcolepsy, but the underlying cause is probably different, as they seem to be at no increased risk of a neurodegenerative disorder.

Many patients with RBD have sustained improvement with clonazepam (0.5–2.0 mg qhs).³ Melatonin at doses up to 9 mg nightly may also prevent attacks.

CIRCADIAN RHYTHM SLEEP DISORDERS

A subset of patients presenting with either insomnia or hypersomnia may have a disorder of sleep *timing* rather than sleep *generation*. Disorders of sleep timing can be either organic (i.e., due to an abnormality of circadian pacemaker[s]) or environmental/behavioral (i.e., due to a disruption of environmental synchronizers). Effective therapies aim to entrain the circadian rhythm of sleep propensity to an appropriate phase.

Delayed Sleep-Wake Phase Disorder Delayed sleep-wake phase disorder (DSWPD) is characterized by: (1) reported sleep onset and wake times intractably later than desired; (2) actual sleep times at nearly the same clock hours daily; and (3) if conducted at the habitual delayed sleep time, essentially normal sleep on polysomnography (except for delayed sleep onset). Patients with DSWPD exhibit an abnormally delayed endogenous circadian phase, which can be assessed by measuring, in a dimly lit environment, the onset of secretion of the endogenous circadian rhythm of pineal melatonin in either the blood or saliva, as light suppresses melatonin secretion. Dim-light melatonin onset (DLMO) in DSWPD patients typically occurs later in the evening than normal, which is about 8:00–9:00 PM (i.e., about 1–2 h before habitual bedtime). Patients tend to be young adults. The delayed circadian phase could be due to: (1) an abnormally long, genetically determined intrinsic period of the endogenous circadian pacemaker; (2) reduced phase-advancing capacity of the pacemaker; (3) slower rate of buildup of homeostatic sleep drive during wakefulness; or (4) an irregular prior sleep-wake schedule, characterized by frequent nights when the patient chooses to remain awake while exposed to artificial light well past midnight (for personal, social, school, or work reasons). In most cases, it is difficult to distinguish among these factors, as patients with either a behaviorally induced or biologically driven circadian phase delay may both exhibit a similar circadian phase delay in DLMO, making it difficult for both to fall asleep at the desired hour. DSWPD is a self-perpetuating condition that can persist for years and may not respond to attempts to reestablish normal bedtime hours. Treatment methods involving phototherapy with blue-enriched light during the morning hours and/or melatonin administration in the evening hours show promise in these patients, although the relapse rate is high. Patients with this circadian rhythm sleep disorder can be distinguished from those who have sleep-onset insomnia because DSWPD patients show late onset of dim-light melatonin secretion.

Advanced Sleep-Wake Phase Disorder Advanced sleep-wake phase disorder (ASWPD) is the converse of DSWPD. Most commonly, this syndrome occurs in older people, 15% of whom report that they cannot sleep past 5:00 AM, with twice that number complaining that they wake up too early at least several times per week. Patients with ASWPD are sleepy during the evening hours, even in social settings. Sleep-wake timing in ASWPD patients can interfere with a normal social life. Patients with this circadian rhythm sleep disorder can be distinguished from those who have early wakening due to insomnia because ASWPD patients show early onset of dim-light melatonin secretion.

In addition to age-related ASWPD, an early-onset familial variant of this condition has also been reported. In two families in which ASWPD was inherited in an autosomal dominant pattern, the syndrome was

due to missense mutations in a circadian clock component (in the casein kinase binding domain of *PER2* in one family, and in casein kinase I delta in the other) that altered the circadian period. Patients with ASWPD may benefit from bright-light and/or blue enriched phototherapy during the evening hours to reset the circadian pacemaker to a later hour.

Non-24-h Sleep-Wake Rhythm Disorder Non-24-h sleep-wake rhythm disorder (N24SWRD) can occur when the primary synchronizing input (i.e., the light-dark cycle) from the environment to the circadian pacemaker is compromised (as occurs in many blind people with no light perception) or when the maximal phase-advancing capacity of the circadian pacemaker cannot accommodate the difference between the 24-h geophysical day and the intrinsic period of the patient's circadian pacemaker, resulting in loss of entrainment to the 24-h day. Rarely, self-selected exposure to artificial light may, in some sighted patients, inadvertently entrain the circadian pacemaker to a >24-h schedule. Affected patients with N24SWRD have difficulty maintaining a stable phase relationship between the output of the pacemaker and the 24-h day. Such patients typically present with an incremental pattern of successive delays in sleep propensity, progressing in and out of phase with local time. When the N24SWRD patient's endogenous circadian rhythms are out of phase with the local environment, nighttime insomnia coexists with excessive daytime sleepiness. Conversely, when the endogenous circadian rhythms are in phase with the local environment, symptoms remit. The interval between symptomatic phases may last several weeks to several months in N24SWRD, depending on the period of the underlying nonentrained rhythm and the 24-h day. Nightly low-dose (0.5 mg) melatonin administration may improve sleep and, in some cases, induce synchronization of the circadian pacemaker.

Shift-Work Disorder More than 7 million workers in the United States regularly work at night, either on a permanent or rotating schedule. Many more begin the commute to work or school between 4:00 AM and 7:00 AM, requiring them to commute and then work during the time of day that they would otherwise be asleep. In addition, each week, millions of “day” workers and students elect to remain awake at night or awaken very early in the morning to work or study to meet work or school deadlines, drive long distances, compete in sporting events, or participate in recreational activities. Such schedules can result in both sleep loss and misalignment of circadian rhythms with respect to the sleep-wake cycle.

The circadian timing system usually fails to adapt successfully to the inverted schedules required by overnight work or the phase advance required by early morning (4:00 AM to 7:00 AM) start times. This leads to a misalignment between the desired work-rest schedule and the output of the pacemaker and to disturbed daytime sleep in most individuals. Excessive work hours (per day or per week), insufficient time off between consecutive days of work or school, and transmeridian travel may be contributing factors. Sleep deficiency, increased length of time awake prior to work, and misalignment of circadian phase produce decreased alertness and performance, increased reaction time, and increased risk of performance lapses, thereby resulting in greater safety hazards among night workers and other sleep-deprived individuals. Sleep disturbance nearly doubles the risk of a fatal work accident. Long-term night shift workers have higher rates of breast, colorectal, and prostate cancer and of cardiac, gastrointestinal, and reproductive disorders. The World Health Organization has added night-shift work to its list of probable carcinogens.

Sleep onset begins in local brain regions before gradually sweeping over the entire brain as sensory thresholds rise and consciousness is lost. A sleepy individual struggling to remain awake may attempt to continue performing routine and familiar motor tasks during the transition state between wakefulness and stage N1 sleep, while unable to adequately process sensory input from the environment. Motor vehicle operators who fail to heed the warning signs of sleepiness are especially vulnerable to sleep-related accidents, as sleep processes can intrude involuntarily upon the waking brain, causing catastrophic consequences. Such sleep-related attentional failures typically last only seconds but are known on occasion to persist for longer durations.

³No medications have been approved by the FDA for the treatment of RBD.

There is a significant increase in the risk of sleep-related, fatal-to-the-driver highway crashes in the early morning and late afternoon hours, coincident with bimodal peaks in the daily rhythm of sleep tendency.

Resident physicians constitute another group of workers at greater risk for accidents and other adverse consequences of lack of sleep and misalignment of the circadian rhythm. Recurrent scheduling of resident physicians to work shifts of ≥ 24 consecutive hours impairs psychomotor performance to a degree that is comparable to alcohol intoxication, doubles the risk of attentional failures among intensive care unit resident physicians working at night, and significantly increases the risk of serious medical errors in intensive care units, including a fivefold increase in the risk of serious diagnostic mistakes. Some 20% of hospital resident physicians report making a fatigue-related mistake that injured a patient, and 5% admit making a fatigue-related mistake that resulted in the death of a patient. Moreover, working for >24 consecutive hours increases the risk of percutaneous injuries and more than doubles the risk of motor vehicle crashes on the commute home. For these reasons, in 2008, the Institute of Medicine concluded that the practice of scheduling resident physicians to work for more than 16 consecutive hours without sleep is hazardous for both resident physicians and their patients.

From 5 to 15% of individuals scheduled to work at night or in the early morning hours have much greater-than-average difficulties remaining awake during night work and sleeping during the day; these individuals are diagnosed with chronic and severe shift-work disorder (SWD). Patients with this disorder have a level of excessive sleepiness during work at night or in the early morning and insomnia during day sleep that the physician judges to be clinically significant; the condition is associated with an increased risk of sleep-related accidents and with some of the illnesses associated with night-shift work. Patients with chronic and severe SWD are profoundly sleepy at work. In fact, their sleep latencies during night work average just 2 min, comparable to mean daytime sleep latency durations of patients with narcolepsy or severe sleep apnea.

TREATMENT SHIFT-WORK DISORDER

Caffeine is frequently used by night workers to promote wakefulness. However, it cannot forestall sleep indefinitely, and it does not shield users from sleep-related performance lapses. Postural changes, exercise, and strategic placement of nap opportunities can sometimes temporarily reduce the risk of fatigue-related performance lapses. Properly timed exposure to blue-enriched light or bright white light can directly enhance alertness and facilitate more rapid adaptation to night-shift work.

Modafinil (200 mg) or armodafinil (150 mg) 30–60 min before the start of each night shift is an effective treatment for the excessive sleepiness during night work in patients with SWD. Although treatment with modafinil or armodafinil significantly improves performance and reduces sleep propensity and the risk of lapses of attention during night work, affected patients remain excessively sleepy.

Fatigue risk management programs for night shift workers should promote education about sleep, increase awareness of the hazards associated with sleep deficiency and night work, and screen for common sleep disorders. Work schedules should be designed to minimize: (1) exposure to night work; (2) the frequency of shift rotations; (3) the number of consecutive night shifts; and (4) the duration of night shifts.

Jet Lag Disorder Each year, more than 60 million people fly from one time zone to another, often resulting in excessive daytime sleepiness, sleep-onset insomnia, and frequent arousals from sleep, particularly in the latter half of the night. The syndrome is transient, typically lasting 2–14 d depending on the number of time zones crossed, the direction of travel, and the traveler's age and phase-shifting capacity. Travelers

who spend more time outdoors at their destination reportedly adapt more quickly than those who remain in hotel rooms, presumably due to brighter (outdoor) light exposure. Avoidance of antecedent sleep loss and obtaining naps on the afternoon prior to overnight travel can reduce the difficulties associated with extended wakefulness. Laboratory studies suggest that low doses of melatonin can enhance sleep efficiency, but only if taken when endogenous melatonin concentrations are low (i.e., during the biologic daytime).

In addition to jet lag associated with travel across time zones, many patients report a behavioral pattern that has been termed *social jet lag*, in which bedtimes and wake times on weekends or days off occur 4–8 h later than during the week. Such recurrent displacement of the timing of the sleep-wake cycle is common in adolescents and young adults and is associated with sleep-onset insomnia, poorer academic performance, increased risk of depressive symptoms, and excessive daytime sleepiness.

MEDICAL IMPLICATIONS OF CIRCADIAN RHYTHMICITY

Prominent circadian variations have been reported in the incidence of acute myocardial infarction, sudden cardiac death, and stroke, the leading causes of death in the United States. Platelet aggregability is increased in the early morning hours, coincident with the peak incidence of these cardiovascular events. Recurrent circadian disruption combined with chronic sleep deficiency, such as occurs during night-shift work, is associated with increased plasma glucose concentrations after a meal due to inadequate pancreatic insulin secretion. Night shift workers with elevated fasting glucose have an increased risk of progressing to diabetes. Blood pressure of night workers with sleep apnea is higher than that of day workers. A better understanding of the possible role of circadian rhythmicity in the acute destabilization of a chronic condition such as atherosclerotic disease could improve the understanding of its pathophysiology.

Diagnostic and therapeutic procedures may also be affected by the time of day at which data are collected. Examples include blood pressure, body temperature, the dexamethasone suppression test, and plasma cortisol levels. The timing of chemotherapy administration has been reported to have an effect on the outcome of treatment. In addition, both the toxicity and effectiveness of drugs can vary with time of day. For example, more than a fivefold difference has been observed in mortality rates following administration of toxic agents to experimental animals at different times of day. Anesthetic agents are particularly sensitive to time-of-day effects. Finally, the physician must be aware of the public health risks associated with the ever-increasing demands made by the 24/7 schedules in our round-the-clock society.

ACKNOWLEDGMENT

John W. Winkelman, MD, PhD and Gary S. Richardson, MD contributed to this chapter in the prior edition and some material from that chapter has been retained here.

VIDEO 38-1 A typical episode of severe cataplexy. The patient is joking and then falls to the ground with an abrupt loss of muscle tone. The electromyogram recordings (four lower traces on the right) show reductions in muscle activity during the period of paralysis. The electroencephalogram (top two traces) shows wakefulness throughout the episode. (Video courtesy of Giuseppe Plazzi, University of Bologna.)

VIDEO 38-2 Typical aggressive movements in rapid eye movement (REM) sleep behavior disorder. (Video courtesy of Dr. Carlos Schenck, University of Minnesota Medical School.)

SECTION 4 DISORDERS OF EYES, EARS, NOSE, AND THROAT

39 Disorders of the Eye

Jonathan C. Horton

THE HUMAN VISUAL SYSTEM

The visual system provides a supremely efficient means for the rapid assimilation of information from the environment to aid in the guidance of behavior. The act of seeing begins with the capture of images focused by the cornea and lens on a light-sensitive membrane in the back of the eye called the *retina*. The retina is actually part of the brain, banished to the periphery to serve as a transducer for the conversion of patterns of light energy into neuronal signals. Light is absorbed by pigment in two types of photoreceptors: rods and cones. In the human retina there are 100 million rods and 5 million cones. The rods operate in dim (scotopic) illumination. The cones function under daylight (photopic) conditions. The cone system is specialized for color perception and high spatial resolution. The majority of cones are within the macula, the portion of the retina that serves the central 10° of vision. In the middle of the macula a small pit termed the *fovea*, packed exclusively with cones, provides the best visual acuity.

Photoreceptors hyperpolarize in response to light, activating bipolar, amacrine, and horizontal cells in the inner nuclear layer. After processing of photoreceptor responses by this complex retinal circuit, the flow of sensory information ultimately converges on a final common pathway: the ganglion cells. These cells translate the visual image impinging on the retina into a continuously varying barrage of action potentials that propagates along the primary optic pathway to visual centers within the brain. There are a million ganglion cells in each retina and hence a million fibers in each optic nerve.

Ganglion cell axons sweep along the inner surface of the retina in the nerve fiber layer, exit the eye at the optic disc, and travel through the optic nerve, optic chiasm, and optic tract to reach targets in the brain. The majority of fibers synapse on cells in the lateral geniculate body, a thalamic relay station. Cells in the lateral geniculate body project in turn to the primary visual cortex. This afferent retinogeniculocortical sensory pathway provides the neural substrate for visual perception. Although the lateral geniculate body is the main target of the retina, separate classes of ganglion cells project to other subcortical visual nuclei involved in different functions. Ganglion cells that mediate pupillary constriction and circadian rhythms are light sensitive owing to a novel visual pigment, melanopsin. Pupil responses are mediated by input to the pretectal olivary nuclei in the midbrain. The pretectal nuclei send their output to the Edinger-Westphal nuclei, which in turn provide parasympathetic innervation to the iris sphincter via an interneuron in the ciliary ganglion. Circadian rhythms are timed by a retinal projection to the suprachiasmatic nucleus. Visual orientation and eye movements are served by retinal input to the superior colliculus. Gaze stabilization and optokinetic reflexes are governed by a group of small retinal targets known collectively as the *brainstem accessory optic system*.

The eyes must be rotated constantly within their orbits to place and maintain targets of visual interest on the fovea. This activity, called *foveation*, or looking, is governed by an elaborate efferent motor system. Each eye is moved by six extraocular muscles that are supplied by cranial nerves from the oculomotor (III), trochlear (IV), and abducens (VI) nuclei. Activity in these ocular motor nuclei is coordinated by pontine and midbrain mechanisms for smooth pursuit, saccades, and gaze stabilization during head and body movements. Large regions of the frontal and parietooccipital cortex control these brainstem eye movement centers by providing descending supranuclear input.

CLINICAL ASSESSMENT OF VISUAL FUNCTION

REFRACTIVE STATE

In approaching a patient with reduced vision, the first step is to decide whether refractive error is responsible. In *emmetropia*, parallel rays from infinity are focused perfectly on the retina. Sadly, this condition is enjoyed by only a minority of the population. In *myopia*, the globe is too long, and light rays come to a focal point in front of the retina. Near objects can be seen clearly, but distant objects require a diverging lens in front of the eye. In *hyperopia*, the globe is too short, and hence a converging lens is used to supplement the refractive power of the eye. In *astigmatism*, the corneal surface is not perfectly spherical, necessitating a cylindrical corrective lens. As an alternative to eyeglasses or contact lenses, refractive error can be corrected by performing laser in situ keratomileusis (LASIK) or photorefractive keratectomy (PRK) to alter the curvature of the cornea.

With the onset of middle age, *presbyopia* develops as the lens within the eye becomes unable to increase its refractive power to accommodate on near objects. To compensate for presbyopia an emmetropic patient must use reading glasses. A patient already wearing glasses for distance correction usually switches to bifocals. The only exception is a myopic patient, who may achieve clear vision at near simply by removing glasses containing the distance prescription.

Refractive errors usually develop slowly and remain stable after adolescence, except in unusual circumstances. For example, the acute onset of diabetes mellitus can produce sudden myopia because of lens edema induced by hyperglycemia. Testing vision through a pinhole aperture is a useful way to screen quickly for refractive error. If visual acuity is better through a pinhole than it is with the unaided eye, the patient needs refraction to obtain best corrected visual acuity.

VISUAL ACUITY

The Snellen chart is used to test acuity at a distance of 6 m (20 ft). For convenience, a scale version of the Snellen chart called the Rosenbaum card is held at 36 cm (14 in.) from the patient (Fig. 39-1). All subjects should be able to read the 6/6 m (20/20 ft) line with each eye using their refractive correction, if any. Patients who need reading glasses because of presbyopia must wear them for accurate testing with the Rosenbaum card. If 6/6 (20/20) acuity is not present in each eye, the deficiency in vision must be explained. If it is worse than 6/240 (20/800), acuity should be recorded in terms of counting fingers, hand motions, light perception, or no light perception. Legal blindness is defined by the Internal Revenue Service as a best corrected acuity of 6/60 (20/200) or less in the better eye or a binocular visual field subtending 20° or less. For driving the laws vary by state, but most states require a corrected acuity of 6/12 (20/40) in at least one eye for unrestricted privileges. Patients with a homonymous hemianopia should not drive.

PUPILS

The pupils should be tested individually in dim light with the patient fixating on a distant target. There is no need to check the near response if the pupils respond briskly to light, because isolated loss of constriction (miosis) to accommodation does not occur. For this reason, the ubiquitous abbreviation PERRLA (pupils equal, round, and reactive to light and accommodation) implies a wasted effort with the last step. However, it is important to test the near response if the light response is poor or absent. Light-near dissociation occurs with neurosyphilis (Argyll Robertson pupil), with lesions of the dorsal midbrain (*Parinaud's syndrome*), and after aberrant regeneration (oculomotor nerve palsy, Adie's tonic pupil).

ROSENBAUM POCKET VISION SCREENER

95	distance equivalent	
	$\frac{20}{800}$	
874	Point Jaeger	$\frac{20}{400}$
2843	26 16	$\frac{20}{200}$
638 E W M X O O	14 10	$\frac{20}{100}$
8 7 4 5 E M W O X O	10 7	$\frac{20}{70}$
6 3 9 2 5 M E E X O X	8 5	$\frac{20}{60}$
4 2 8 3 6 5 W E M O X O	6 3	$\frac{20}{40}$
3 7 4 2 5 8 E W M X X O	5 2	$\frac{20}{30}$
9 3 7 8 2 6 W M E X O O	4 1	$\frac{20}{25}$
4 2 8 7 3 5 E W M O O X	3 1+	$\frac{20}{20}$

14" TESTING DISTANCE

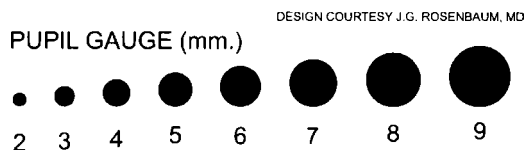


FIGURE 39-1 The Rosenbaum card is a miniature, scale version of the Snellen chart for testing visual acuity at near. When the visual acuity is recorded, the Snellen distance equivalent should bear a notation indicating that vision was tested at near, not at 6 m (20 ft), or else the Jaeger number system should be used to report the acuity.

An eye with no light perception has no pupillary response to direct light stimulation. If the retina or optic nerve is only partially injured, the direct pupillary response will be weaker than the consensual pupillary response evoked by shining a light into the healthy fellow eye. A *relative afferent pupillary defect* (Marcus Gunn pupil) can be elicited with the swinging flashlight test (Fig. 39-2). It is an extremely useful sign in retrobulbar optic neuritis and other optic nerve diseases, in which it may be the sole objective evidence for disease. In bilateral optic neuropathy, no afferent pupil defect is present if the optic nerves are affected equally.

Subtle inequality in pupil size, up to 0.5 mm, is a fairly common finding in normal persons. The diagnosis of essential or physiologic anisocoria is secure as long as the relative pupil asymmetry remains constant as ambient lighting varies. Anisocoria that increases in dim light indicates a sympathetic paresis of the iris dilator muscle. The triad of miosis with ipsilateral ptosis and anhidrosis constitutes *Horner's syndrome*, although anhidrosis is an inconstant feature. Brainstem stroke, carotid dissection, and neoplasm impinging on the sympathetic chain occasionally are identified as the cause of Horner's syndrome, but most cases are idiopathic.

Anisocoria that increases in bright light suggests a parasympathetic palsy. The first concern is an oculomotor nerve paresis. This possibility is excluded if the eye movements are full and the patient has no ptosis or diplopia. Acute pupillary dilation (mydriasis) can result from

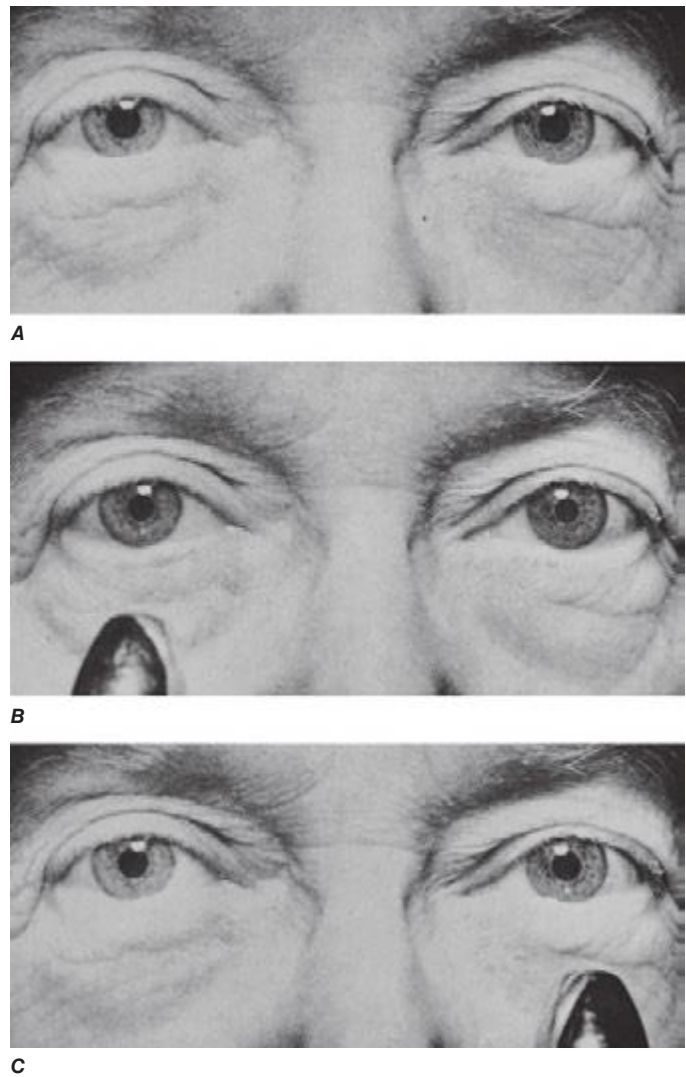


FIGURE 39-2 Demonstration of a relative afferent pupil defect (Marcus Gunn pupil) in the left eye, done with the patient fixating on a distant target. **A.** With dim background lighting, the pupils are equal and relatively large. **B.** Shining a flashlight into the right eye evokes equal, strong constriction of both pupils. **C.** Swinging the flashlight over to the damaged left eye causes dilation of both pupils, although they remain smaller than in **A.** Swinging the flashlight back over to the healthy right eye would result in symmetric constriction back to the appearance shown in **B.** Note that the pupils always remain equal; the damage to the left retina/optic nerve is revealed by weaker bilateral pupil constriction to a flashlight in the left eye compared with the right eye. (From P Levatin: *Arch Ophthalmol* 62:768, 1959. Copyright © 1959 American Medical Association. All rights reserved.)

damage to the ciliary ganglion in the orbit. Common mechanisms are infection (herpes zoster, influenza), trauma (blunt, penetrating, surgical), and ischemia (diabetes, temporal arteritis). After denervation of the iris sphincter the pupil does not respond well to light, but the response to near is often relatively intact. When the near stimulus is removed, the pupil redilates very slowly compared with the normal pupil, hence the term *tonic pupil*. In *Adie's syndrome* a tonic pupil is present, sometimes in conjunction with weak or absent tendon reflexes in the lower extremities. This benign disorder, which occurs predominantly in healthy young women, is assumed to represent a mild dysautonomia. Tonic pupils are also associated with Shy-Drager syndrome, segmental hypohidrosis, diabetes, and amyloidosis. Occasionally, a tonic pupil is discovered incidentally in an otherwise completely

normal, asymptomatic individual. The diagnosis is confirmed by placing a drop of dilute (0.125%) pilocarpine into each eye. Denervation hypersensitivity produces pupillary constriction in a tonic pupil, whereas the normal pupil shows no response. Pharmacologic dilatation from accidental or deliberate instillation of anticholinergic agents (atropine, scopolamine drops) into the eye also can produce pupillary mydriasis. In this situation, normal strength (1%) pilocarpine causes no constriction.

Both pupils are affected equally by systemic medications. They are small with narcotic use (morphine, heroin) and large with anticholinergics (scopolamine). Parasympathetic agents (pilocarpine, demecarium bromide) used to treat glaucoma produce miosis. In any patient with an unexplained pupillary abnormality, a slit-lamp examination is helpful to exclude surgical trauma to the iris, an occult foreign body, perforating injury, intraocular inflammation, adhesions (synechia), angle-closure glaucoma, and iris sphincter rupture from blunt trauma.

EYE MOVEMENTS AND ALIGNMENT

Eye movements are tested by asking the patient, with both eyes open, to pursue a small target such as a penlight into the cardinal fields of gaze. Normal ocular versions are smooth, symmetric, full, and maintained in all directions without nystagmus. Saccades, or quick refixation eye movements, are assessed by having the patient look back and forth between two stationary targets. The eyes should move rapidly and accurately in a single jump to their target. Ocular alignment can be judged by holding a penlight directly in front of the patient at about 1 m. If the eyes are straight, the corneal light reflex will be centered in the middle of each pupil. To test eye alignment more precisely, the cover test is useful. The patient is instructed to look at a small fixation target in the distance. One eye is covered suddenly while the second eye is observed. If the second eye shifts to fixate on the target, it was misaligned. If it does not move, the first eye is uncovered and the test is repeated on the second eye. If neither eye moves the eyes are aligned orthotropically. If the eyes are orthotropic in primary gaze but the patient complains of diplopia, the cover test should be performed with the head tilted or turned in whatever direction elicits diplopia. With practice, the examiner can detect an ocular deviation (heterotropia) as small as 1–2° with the cover test. In a patient with vertical diplopia, a small deviation can be difficult to detect and easy to dismiss. The magnitude of the deviation can be measured by placing a prism in front of the misaligned eye to determine the power required to neutralize the fixation shift evoked by covering the other eye. Temporary press-on plastic Fresnel prisms, prism eyeglasses, or eye muscle surgery can be used to restore binocular alignment.

STEREOPSIS

Stereoacuity is determined by presenting targets with retinal disparity separately to each eye by using polarized images. The most popular office tests measure a range of thresholds from 800–40 seconds of arc. Normal stereoacuity is 40 seconds of arc. If a patient achieves this level of stereoacuity, one is assured that the eyes are aligned orthotropically and that vision is intact in each eye. Random dot stereograms have no monocular depth cues and provide an excellent screening test for strabismus and amblyopia in children.

COLOR VISION

The retina contains three classes of cones, with visual pigments of differing peak spectral sensitivity: red (560 nm), green (530 nm), and blue (430 nm). The red and green cone pigments are encoded on the X chromosome, and the blue cone pigment on chromosome 7. Mutations of the blue cone pigment are exceedingly rare. Mutations of the red and green pigments cause congenital X-linked color blindness in 8% of males. Affected individuals are not truly color blind; rather, they differ from normal subjects in the way they perceive color and how they combine primary monochromatic lights to match a particular color. Anomalous trichromats have three cone types, but a mutation in one cone pigment (usually red or green) causes a shift in peak spectral sensitivity, altering the proportion of primary colors required to achieve a color match. Dichromats have only two cone types and

therefore will accept a color match based on only two primary colors. Anomalous trichromats and dichromats have 6/6 (20/20) visual acuity, but their hue discrimination is impaired. Ishihara color plates can be used to detect red-green color blindness. The test plates contain a hidden number that is visible only to subjects with color confusion from red-green color blindness. Because color blindness is almost exclusively X-linked, it is worth screening only male children.

The Ishihara plates often are used to detect acquired defects in color vision, although they are intended as a screening test for congenital color blindness. Acquired defects in color vision frequently result from disease of the macula or optic nerve. For example, patients with a history of optic neuritis often complain of color desaturation long after their visual acuity has returned to normal. Color blindness also can result from bilateral strokes involving the ventral portion of the occipital lobe (cerebral achromatopsia). Such patients can perceive only shades of gray and also may have difficulty recognizing faces (prosopagnosia). Infarcts of the dominant occipital lobe sometimes give rise to color anomia. Affected patients can discriminate colors but cannot name them.

VISUAL FIELDS

Vision can be impaired by damage to the visual system anywhere from the eyes to the occipital lobes. One can localize the site of the lesion with considerable accuracy by mapping the visual field deficit by finger confrontation and then correlating it with the topographic anatomy of the visual pathway (Fig. 39-3). Quantitative visual field mapping is performed by computer-driven perimeters that present a target of variable intensity at fixed positions in the visual field (Fig. 39-3A). By generating an automated printout of light thresholds, these static perimeters provide a sensitive means of detecting scotomas in the visual field. They are exceedingly useful for serial assessment of visual function in chronic diseases such as glaucoma and pseudotumor cerebri.

The crux of visual field analysis is to decide whether a lesion is before, at, or behind the optic chiasm. If a scotoma is confined to one eye, it must be due to a lesion anterior to the chiasm, involving either the optic nerve or the retina. Retinal lesions produce scotomas that correspond optically to their location in the fundus. For example, a superior-nasal retinal detachment results in an inferior-temporal field cut. Damage to the macula causes a central scotoma (Fig. 39-3B).

Optic nerve disease produces characteristic patterns of visual field loss. Glaucoma selectively destroys axons that enter the superotemporal or inferotemporal poles of the optic disc, resulting in arcuate scotomas shaped like a Turkish scimitar, which emanate from the blind spot and curve around fixation to end flat against the horizontal meridian (Fig. 39-3C). This type of field defect mirrors the arrangement of the nerve fiber layer in the temporal retina. Arcuate or nerve fiber layer scotomas also result from optic neuritis, ischemic optic neuropathy, optic disc drusen, and branch retinal artery or vein occlusion.

Damage to the entire upper or lower pole of the optic disc causes an altitudinal field cut that follows the horizontal meridian (Fig. 39-3D). This pattern of visual field loss is typical of ischemic optic neuropathy but also results from retinal vascular occlusion, advanced glaucoma, and optic neuritis.

About half the fibers in the optic nerve originate from ganglion cells serving the macula. Damage to papillomacular fibers causes a cecentral scotoma that encompasses the blind spot and macula (Fig. 39-3E). If the damage is irreversible, pallor eventually appears in the temporal portion of the optic disc. Temporal pallor from a cecentral scotoma may develop in optic neuritis, nutritional optic neuropathy, toxic optic neuropathy, Leber's hereditary optic neuropathy, Kjer's dominant optic atrophy, and compressive optic neuropathy. It is worth mentioning that the temporal side of the optic disc is slightly paler than the nasal side in most normal individuals. Therefore, it sometimes can be difficult to decide whether the temporal pallor visible on fundus examination represents a pathologic change. Pallor of the nasal rim of the optic disc is a less equivocal sign of optic atrophy.

At the optic chiasm, fibers from nasal ganglion cells decussate into the contralateral optic tract. Crossed fibers are damaged more by

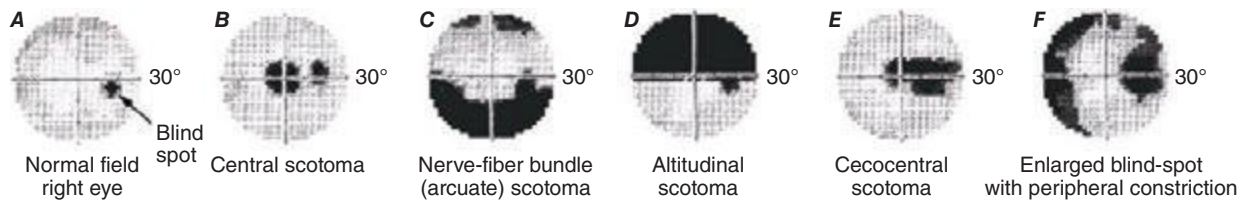
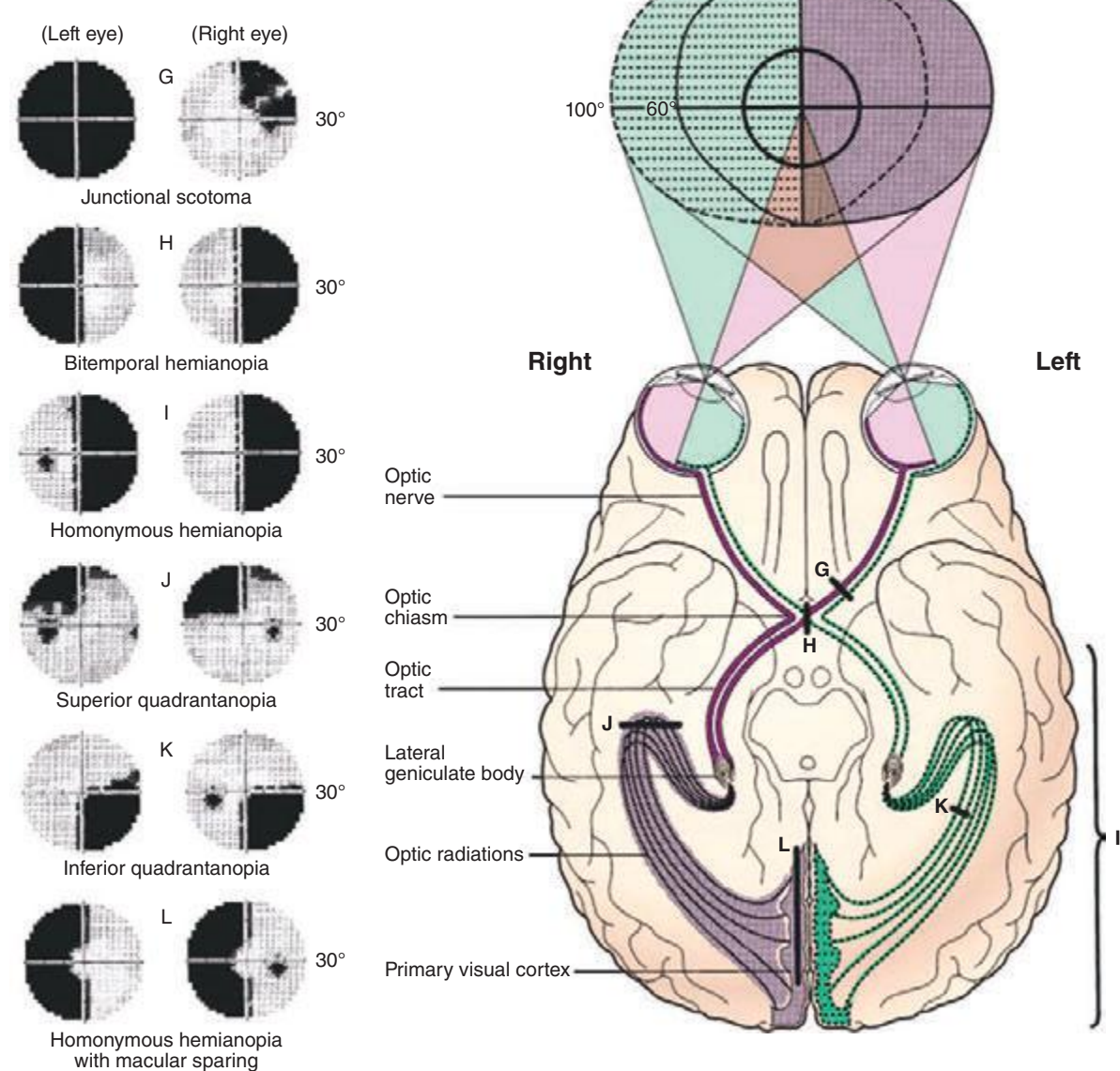
Monocular prechiasmal field defects:**Binocular chiasmal or postchiasmal field defects:**

FIGURE 39-3 Ventral view of the brain, correlating patterns of visual field loss with the sites of lesions in the visual pathway. The visual fields overlap partially, creating 120° of central binocular field flanked by a 40° monocular crescent on either side. The visual field maps in this figure were done with a computer-driven perimeter (Humphrey Instruments, Carl Zeiss, Inc.). It plots the retinal sensitivity to light in the central 30° by using a gray scale format. Areas of visual field loss are shown in black. The examples of common monocular, prechiasmal field defects are all shown for the right eye. By convention, the visual fields are always recorded with the left eye's field on the left and the right eye's field on the right, just as the patient sees the world.

compression than are uncrossed fibers. As a result, mass lesions of the sellar region cause a temporal hemianopia in each eye. Tumors anterior to the optic chiasm, such as meningiomas of the tuberculum sellae, produce a junctional scotoma characterized by an optic neuropathy in one eye and a superior-temporal field cut in the other eye (Fig. 39-3G). More symmetric compression of the optic chiasm by a pituitary adenoma (see Fig. 403-1), meningioma, craniopharyngioma, glioma, or aneurysm results in a bitemporal hemianopia (Fig. 39-3H).

The insidious development of a bitemporal hemianopia often goes unnoticed by the patient and will escape detection by the physician unless each eye is tested separately.

It is difficult to localize a postchiasmal lesion accurately, because injury anywhere in the optic tract, lateral geniculate body, optic radiations, or visual cortex can produce a homonymous hemianopia (i.e., a temporal hemifield defect in the contralateral eye and a matching nasal hemifield defect in the ipsilateral eye) (Fig. 39-3I). A unilateral

postchiasmal lesion leaves the visual acuity in each eye unaffected, although the patient may read the letters on only the left or right half of the eye chart. Lesions of the optic radiations tend to cause poorly matched or incongruous field defects in each eye. Damage to the optic radiations in the temporal lobe (Meyer's loop) produces a superior quadrantic homonymous hemianopia (Fig. 39-3J), whereas injury to the optic radiations in the parietal lobe results in an inferior quadrantic homonymous hemianopia (Fig. 39-3K). Lesions of the primary visual cortex give rise to dense, congruous hemianopic field defects. Occlusion of the posterior cerebral artery supplying the occipital lobe is a common cause of total homonymous hemianopia. Some patients with hemianopia after occipital stroke have macular sparing, because the macular representation at the tip of the occipital lobe is supplied by collaterals from the middle cerebral artery (Fig. 39-3L). Destruction of both occipital lobes produces cortical blindness. This condition can be distinguished from bilateral prechiasmal visual loss by noting that the pupil responses and optic fundi remain normal.

DISORDERS

RED OR PAINFUL EYE

Corneal Abrasions Corneal abrasions are seen best by placing a drop of fluorescein in the eye and looking with the slit lamp, using a cobalt-blue light. A penlight with a blue filter will suffice if a slit lamp is not available. Damage to the corneal epithelium is revealed by yellow fluorescence of the exposed basement membrane underlying the epithelium. It is important to check for foreign bodies. To search the conjunctival fornices, the lower lid should be pulled down and the upper lid everted. A foreign body can be removed with a moistened cotton-tipped applicator after a drop of a topical anesthetic such as proparacaine has been placed in the eye. Alternatively, it may be possible to flush the foreign body from the eye by irrigating copiously with saline or artificial tears. If the corneal epithelium has been abraded, antibiotic ointment and a patch should be applied to the eye. A drop of an intermediate-acting cycloplegic such as cyclopentolate hydrochloride 1% helps reduce pain by relaxing the ciliary body. The eye should be reexamined the next day. Minor abrasions may not require patching, antibiotics, or cycloplegia.

Subconjunctival Hemorrhage This results from rupture of small vessels bridging the potential space between the episclera and the conjunctiva. Blood dissecting into this space can produce a spectacular red eye, but vision is not affected and the hemorrhage resolves without treatment. Subconjunctival hemorrhage is usually spontaneous but can result from blunt trauma, eye rubbing, or vigorous coughing. Occasionally it is a clue to an underlying bleeding disorder.

Pinguecula Pinguecula is a small, raised conjunctival nodule at the temporal or nasal limbus. In adults such lesions are extremely common and have little significance unless they become inflamed (pingueculitis). They are more apt to occur in workers with frequent outdoor exposure. A *pterygium* resembles a pinguecula but has crossed the limbus to encroach on the corneal surface. Removal is justified when symptoms of irritation or blurring develop, but recurrence is a common problem.

Blepharitis This refers to inflammation of the eyelids. The most common form occurs in association with acne rosacea or seborrheic dermatitis. The eyelid margins usually are colonized heavily by staphylococci. Upon close inspection, they appear greasy, ulcerated, and crusted with scaling debris that clings to the lashes. Treatment consists of strict eyelid hygiene, using warm compresses and eyelash scrubs with baby shampoo. An external *hordeolum* (sty) is caused by staphylococcal infection of the superficial accessory glands of Zeis or Moll located in the eyelid margins. An internal hordeolum occurs after suppurative infection of the oil-secreting meibomian glands within the tarsal plate of the eyelid. Topical antibiotics such as bacitracin/polymyxin B ophthalmic ointment can be applied. Systemic antibiotics, usually tetracyclines or azithromycin, sometimes are necessary for treatment of meibomian gland inflammation (meibomitis) or chronic, severe blepharitis. A *chalazion* is a painless, chronic granulomatous

inflammation of a meibomian gland that produces a pealike nodule within the eyelid. It can be incised and drained or injected with glucocorticoids. Basal cell, squamous cell, or meibomian gland carcinoma should be suspected with any nonhealing ulcerative lesion of the eyelids.

Dacryocystitis An inflammation of the lacrimal drainage system, dacryocystitis can produce epiphora (tearing) and ocular injection. Gentle pressure over the lacrimal sac evokes pain and reflux of mucus or pus from the tear puncta. Dacryocystitis usually occurs after obstruction of the lacrimal system. It is treated with topical and systemic antibiotics, followed by probing, silicone stent intubation, or surgery to reestablish patency. *Entropion* (inversion of the eyelid) or *ectropion* (sagging or eversion of the eyelid) can also lead to epiphora and ocular irritation.

Conjunctivitis Conjunctivitis is the most common cause of a red, irritated eye. Pain is minimal, and visual acuity is reduced only slightly. The most common viral etiology is adenovirus infection. It causes a watery discharge, a mild foreign-body sensation, and photophobia. Bacterial infection tends to produce a more mucopurulent exudate. Mild cases of infectious conjunctivitis usually are treated empirically with broad-spectrum topical ocular antibiotics such as sulfacetamide 10%, polymyxin-bacitracin, or a trimethoprim-polymyxin combination. Smears and cultures usually are reserved for severe, resistant, or recurrent cases of conjunctivitis. To prevent contagion, patients should be admonished to wash their hands frequently, not to touch their eyes, and to avoid direct contact with others.

Allergic Conjunctivitis This condition is extremely common and often is mistaken for infectious conjunctivitis. Itching, redness, and epiphora are typical. The palpebral conjunctiva may become hypertrophic with giant excrescences called cobblestone papillae. Irritation from contact lenses or any chronic foreign body also can induce formation of cobblestone papillae. *Atopic conjunctivitis* occurs in subjects with atopic dermatitis or asthma. Symptoms caused by allergic conjunctivitis can be alleviated with cold compresses, topical vasoconstrictors, antihistamines, and mast cell stabilizers such as cromolyn sodium. Topical glucocorticoid solutions provide dramatic relief of immune-mediated forms of conjunctivitis, but their long-term use is ill advised because of the complications of glaucoma, cataract, and secondary infection. Topical nonsteroidal anti-inflammatory drugs (NSAIDs) (e.g., ketorolac tromethamine) are better alternatives.

Keratoconjunctivitis Sicca Also known as dry eye, this produces a burning foreign-body sensation, injection, and photophobia. In mild cases the eye appears surprisingly normal, but tear production measured by wetting of a filter paper (Schirmer strip) is deficient. A variety of systemic drugs, including antihistaminic, anticholinergic, and psychotropic medications, result in dry eye by reducing lacrimal secretion. Disorders that involve the lacrimal gland directly, such as sarcoidosis and Sjögren's syndrome, also cause dry eye. Patients may develop dry eye after radiation therapy if the treatment field includes the orbits. Problems with ocular drying are also common after lesions affecting cranial nerve V or VII. Corneal anesthesia is particularly dangerous, because the absence of a normal blink reflex exposes the cornea to injury without pain to warn the patient. Dry eye is managed by frequent and liberal application of artificial tears and ocular lubricants. In severe cases the tear puncta can be plugged or cauterized to reduce lacrimal outflow.

Keratitis Keratitis is a threat to vision because of the risk of corneal clouding, scarring, and perforation. Worldwide, the two leading causes of blindness from keratitis are trachoma from chlamydial infection and vitamin A deficiency related to malnutrition. In the United States, contact lenses play a major role in corneal infection and ulceration. They should not be worn by anyone with an active eye infection. In evaluating the cornea, it is important to differentiate between a superficial infection (*keratoconjunctivitis*) and a deeper, more serious ulcerative process. The latter is accompanied by greater visual loss, pain, photophobia, redness, and discharge. Slit-lamp examination shows disruption of the corneal epithelium, a cloudy infiltrate or abscess in the

stroma, and an inflammatory cellular reaction in the anterior chamber. In severe cases, pus settles at the bottom of the anterior chamber, giving rise to a hypopyon. Immediate empirical antibiotic therapy should be initiated after corneal scrapings are obtained for Gram's stain, Giemsa stain, and cultures. Fortified topical antibiotics are most effective, supplemented with subconjunctival antibiotics as required. A fungal etiology should always be considered in a patient with keratitis. Fungal infection is common in warm humid climates, especially after penetration of the cornea by plant or vegetable material.

Herpes Simplex The *herpesviruses* are a major cause of blindness from keratitis. Most adults in the United States have serum antibodies to herpes simplex, indicating prior viral infection (Chap. 216). Primary ocular infection generally is caused by herpes simplex type 1 rather than type 2. It manifests as a unilateral follicular blepharoconjunctivitis that is easily confused with adenoviral conjunctivitis unless telltale vesicles appear on the periocular skin or conjunctiva. A dendritic pattern of corneal epithelial ulceration revealed by fluorescein staining is pathognomonic for herpes infection but is seen in only a minority of primary infections. Recurrent ocular infection arises from reactivation of the latent herpesvirus. Viral eruption in the corneal epithelium may result in the characteristic herpes dendrite. Involvement of the corneal stroma produces edema, vascularization, and iridocyclitis. Herpes keratitis is treated with topical antiviral agents, cycloplegics, and oral acyclovir. Topical glucocorticoids are effective in mitigating corneal scarring but must be used with extreme caution because of the danger of corneal melting and perforation. Topical glucocorticoids also carry the risk of prolonging infection and inducing glaucoma.

Herpes Zoster Herpes zoster from reactivation of latent varicella (chickenpox) virus causes a dermatomal pattern of painful vesicular dermatitis. Ocular symptoms can occur after zoster eruption in any branch of the trigeminal nerve but are particularly common when vesicles form on the nose, reflecting nasociliary (V1) nerve involvement (Hutchinson's sign). Herpes zoster ophthalmicus produces corneal dendrites, which can be difficult to distinguish from those seen in herpes simplex. Stromal keratitis, anterior uveitis, raised intraocular pressure, ocular motor nerve palsies, acute retinal necrosis, and postherpetic scarring and neuralgia are other common sequelae. Herpes zoster ophthalmicus is treated with antiviral agents and cycloplegics. In severe cases, glucocorticoids may be added to prevent permanent visual loss from corneal scarring.

Episcleritis This is an inflammation of the episclera, a thin layer of connective tissue between the conjunctiva and the sclera. Episcleritis resembles conjunctivitis, but it is a more localized process and discharge is absent. Most cases of episcleritis are idiopathic, but some occur in the setting of an autoimmune disease. *Scleritis* refers to a deeper, more severe inflammatory process that frequently is associated with a connective tissue disease such as rheumatoid arthritis, lupus erythematosus, polyarteritis nodosa, granulomatosis with polyangiitis (Wegener's), or relapsing polychondritis. The inflammation and thickening of the sclera can be diffuse or nodular. In anterior forms of scleritis, the globe assumes a violet hue and the patient complains of severe ocular tenderness and pain. With posterior scleritis, the pain and redness may be less marked, but there is often proptosis, choroidal effusion, reduced motility, and visual loss. Episcleritis and scleritis should be treated with NSAIDs. If these agents fail, topical or even systemic glucocorticoid therapy may be necessary, especially if an underlying autoimmune process is active.

Uveitis Involving the anterior structures of the eye, uveitis also is called *iritis* or *iridocyclitis*. The diagnosis requires slit-lamp examination to identify inflammatory cells floating in the aqueous humor or deposited on the corneal endothelium (keratic precipitates). Anterior uveitis develops in sarcoidosis, ankylosing spondylitis, juvenile rheumatoid arthritis, inflammatory bowel disease, psoriasis, reactive arthritis, and Behçet's disease. It also is associated with herpes infections, syphilis, Lyme disease, onchocerciasis, tuberculosis, and leprosy. Although anterior uveitis can occur in conjunction with many diseases, no cause is found to explain the majority of cases. For this reason,

laboratory evaluation usually is reserved for patients with recurrent or severe anterior uveitis. Treatment is aimed at reducing inflammation and scarring by judicious use of topical glucocorticoids. Dilatation of the pupil reduces pain and prevents the formation of synechiae.

Posterior Uveitis This is diagnosed by observing inflammation of the vitreous, retina, or choroid on fundus examination. It is more likely than anterior uveitis to be associated with an identifiable systemic disease. Some patients have panuveitis, or inflammation of both the anterior and posterior segments of the eye. Posterior uveitis is a manifestation of autoimmune diseases such as sarcoidosis, Behçet's disease, Vogt-Koyanagi-Harada syndrome, and inflammatory bowel disease. It also accompanies diseases such as toxoplasmosis, onchocerciasis, cysticercosis, coccidioidomycosis, toxocariasis, and histoplasmosis; infections caused by organisms such as *Candida*, *Pneumocystis carinii*, *Cryptococcus*, *Aspergillus*, herpes, and cytomegalovirus (see Fig. 219-1); and other diseases, such as syphilis, Lyme disease, tuberculosis, cat-scratch disease, Whipple's disease, and brucellosis. In multiple sclerosis, chronic inflammatory changes can develop in the extreme periphery of the retina (pars planitis or intermediate uveitis).

Acute Angle-Closure Glaucoma This is an unusual but frequently misdiagnosed cause of a red, painful eye. Asian populations have a particularly high risk of angle-closure glaucoma. Susceptible eyes have a shallow anterior chamber because the eye has either a short axial length (hyperopia) or a lens enlarged by the gradual development of cataract. When the pupil becomes mid-dilated, the peripheral iris blocks aqueous outflow via the anterior chamber angle and the intraocular pressure rises abruptly, producing pain, injection, corneal edema, obscurations, and blurred vision. In some patients, ocular symptoms are overshadowed by nausea, vomiting, or headache, prompting a fruitless workup for abdominal or neurologic disease. The diagnosis is made by measuring the intraocular pressure during an acute attack or by performing gonioscopy, a procedure that allows one to observe a narrow chamber angle with a mirrored contact lens. Acute angle closure is treated with acetazolamide (PO or IV), topical beta blockers, prostaglandin analogues, α_2 -adrenergic agonists, and pilocarpine to induce miosis. If these measures fail, a laser can be used to create a hole in the peripheral iris to relieve pupillary block. Many physicians are reluctant to dilate patients routinely for fundus examination because they fear precipitating an angle-closure glaucoma. The risk is actually remote and more than outweighed by the potential benefit to patients of discovering a hidden fundus lesion visible only through a fully dilated pupil. Moreover, a single attack of angle closure after pharmacologic dilatation rarely causes any permanent damage to the eye and serves as an inadvertent provocative test to identify patients with narrow angles who would benefit from prophylactic laser iridectomy.

Endophthalmitis This results from bacterial, viral, fungal, or parasitic infection of the internal structures of the eye. It usually is acquired by hematogenous seeding from a remote site. Chronically ill, diabetic, or immunosuppressed patients, especially those with a history of indwelling IV catheters or positive blood cultures, are at greatest risk for endogenous endophthalmitis. Although most patients have ocular pain and injection, visual loss is sometimes the only symptom. Septic emboli from a diseased heart valve or a dental abscess that lodge in the retinal circulation can give rise to endophthalmitis. White-centered retinal hemorrhages known as Roth's spots (Fig. 39-4) are considered pathognomonic for subacute bacterial endocarditis, but they also appear in leukemia, diabetes, and many other conditions. Endophthalmitis also occurs as a complication of ocular surgery, especially glaucoma filtering, occasionally months or even years after the operation. An occult penetrating foreign body or unrecognized trauma to the globe should be considered in any patient with unexplained intraocular infection or inflammation.

TRANSIENT OR SUDDEN VISUAL LOSS

Amaurosis Fugax This term refers to a transient ischemic attack of the retina (Chap. 446). Because neural tissue has a high rate of metabolism, interruption of blood flow to the retina for more than a few seconds



FIGURE 39-4 Roth's spot, cotton-wool spot, and retinal hemorrhages in a 48-year-old liver transplant patient with candidemia from immunosuppression.

results in *transient monocular blindness*, a term used interchangeably with amaurosis fugax. Patients describe a rapid fading of vision like a curtain descending, sometimes affecting only a portion of the visual field. Amaurosis fugax usually results from an embolus that becomes stuck within a retinal arteriole (Fig. 39-5). If the embolus breaks up or passes, flow is restored and vision returns quickly to normal without permanent damage. With prolonged interruption of blood flow, the inner retina suffers infarction. Ophthalmoscopy reveals zones of whitened, edematous retina following the distribution of branch retinal arterioles. Complete occlusion of the central retinal artery produces arrest of blood flow and a milky retina with a cherry-red fovea (Fig. 39-6). Emboli are composed of cholesterol (Hollenhorst plaque), calcium, or platelet-fibrin debris. The most common source is an atherosclerotic plaque in the carotid artery or aorta, although emboli also can arise from the heart, especially in patients with diseased valves, atrial fibrillation, or wall motion abnormalities.

In rare instances, amaurosis fugax results from low central retinal artery perfusion pressure in a patient with a critical stenosis of the ipsilateral carotid artery and poor collateral flow via the circle of Willis. In this situation, amaurosis fugax develops when there is a dip in systemic blood pressure or a slight worsening of the carotid stenosis. Sometimes there is contralateral motor or sensory loss, indicating concomitant hemispheric cerebral ischemia.



FIGURE 39-5 Hollenhorst plaque lodged at the bifurcation of a retinal arteriole proves that a patient is shedding emboli from the carotid artery, great vessels, or heart.



FIGURE 39-6 Central retinal artery occlusion in a 78-year-old man reducing acuity to counting fingers in the right eye. Note the splinter hemorrhage on the optic disc and the slightly milky appearance to the macula with a cherry-red fovea.

Retinal arterial occlusion also occurs rarely in association with retinal migraine, lupus erythematosus, anticardiolipin antibodies, anticoagulant deficiency states (protein S, protein C, and antithrombin deficiency), pregnancy, IV drug abuse, blood dyscrasias, dysproteinemias, and temporal arteritis.

Marked *systemic hypertension* causes sclerosis of retinal arterioles, splinter hemorrhages, focal infarcts of the nerve fiber layer (cotton-wool spots), and leakage of lipid and fluid (hard exudate) into the macula (Fig. 39-7). In hypertensive crisis, sudden visual loss can result from vasospasm of retinal arterioles and retinal ischemia. In addition, acute hypertension may produce visual loss from ischemic swelling of the optic disc. Patients with acute hypertensive retinopathy should be treated by lowering the blood pressure. However, the blood pressure should not be reduced precipitously, because there is a danger of optic disc infarction from sudden hypoperfusion.

Impending *branch or central retinal vein occlusion* can produce prolonged visual obscurations that resemble those described by patients with amaurosis fugax. The veins appear engorged and phlebitic, with numerous retinal hemorrhages (Fig. 39-8). In some patients, venous blood flow recovers spontaneously, whereas others evolve a frank obstruction with extensive retinal bleeding ("blood and thunder" appearance), infarction, and visual loss. Venous occlusion of the retina is often idiopathic, but hypertension, diabetes, and glaucoma

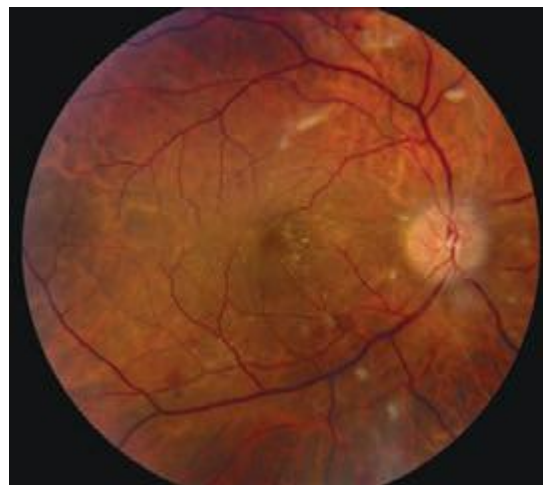


FIGURE 39-7 Hypertensive retinopathy with blurred optic disc, scattered hemorrhages, cotton-wool spots (nerve fiber layer infarcts), and foveal exudate in a 62-year-old man with chronic renal failure and a systolic blood pressure of 220.

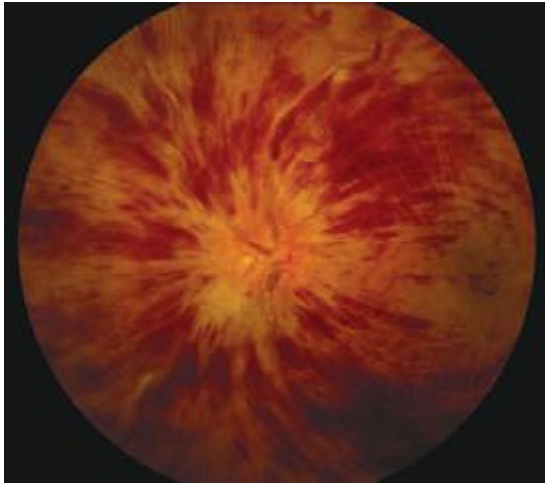


FIGURE 39-8 Central retinal vein occlusion can produce massive retinal hemorrhage (“blood and thunder”), ischemia, and vision loss.

are prominent risk factors. Polycythemia, thrombocythemia, or other factors leading to an underlying hypercoagulable state should be corrected; aspirin treatment may be beneficial.

Anterior Ischemic Optic Neuropathy (AION) This is caused by insufficient blood flow through the posterior ciliary arteries that supply the optic disc. It produces painless monocular visual loss that is sudden in onset, followed sometimes by stuttering progression. The optic disc appears swollen and surrounded by nerve fiber layer splinter hemorrhages (Fig. 39-9). AION is divided into two forms: arteritic and nonarteritic. The nonarteritic form is most common. No specific cause can be identified, although diabetes and hypertension are common risk factors. A crowded disc architecture and small optic cup predispose to the development of nonarteritic AION. No treatment is available. About 5% of patients, especially those age >60, develop the arteritic form of AION in conjunction with giant-cell (temporal) arteritis (Chap. 385). It is urgent to recognize arteritic AION so that high doses of glucocorticoids can be instituted immediately to prevent blindness in the second eye. Symptoms of polymyalgia rheumatica may be present; the sedimentation rate and C-reactive protein level are usually elevated. In a patient with visual loss from suspected arteritic AION, temporal artery biopsy is mandatory to confirm the diagnosis. Glucocorticoids should be started immediately, without waiting for the biopsy to be completed. The diagnosis of arteritic AION is difficult to sustain in the face of a negative temporal artery biopsy, but such cases do occur



FIGURE 39-9 Anterior ischemic optic neuropathy from temporal arteritis in a 67-year-old woman with acute disc swelling, splinter hemorrhages, visual loss, and an erythrocyte sedimentation rate of 70 mm/h.

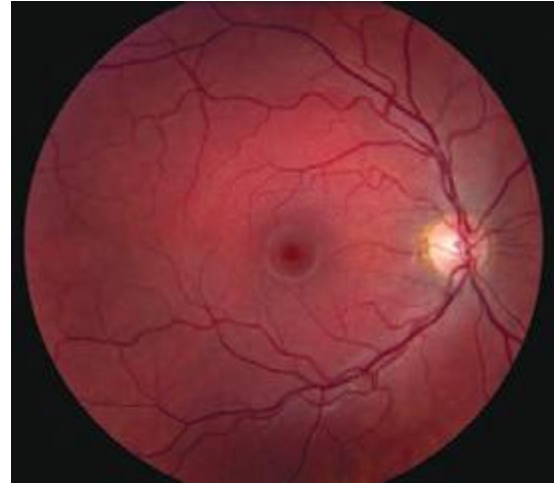


FIGURE 39-10 Retrobulbar optic neuritis is characterized by a normal fundus examination initially, hence the rubric “the doctor sees nothing, and the patient sees nothing.” Optic atrophy develops after severe or repeated attacks.

rarely. It is important to biopsy an arterial segment of at least 3 cm and to examine a sufficient number of tissue sections prepared from the specimen.

Posterior Ischemic Optic Neuropathy This is an uncommon cause of acute visual loss, induced by the combination of severe anemia and hypotension. Cases have been reported after major blood loss during surgery (especially in patients undergoing cardiac or lumbar spine operations), exsanguinating trauma, gastrointestinal bleeding, and renal dialysis. The fundus usually appears normal, although optic disc swelling develops if the process extends anteriorly far enough to reach the globe. Vision can be salvaged in some patients by prompt blood transfusion and reversal of hypotension.

Optic Neuritis This is a common inflammatory disease of the optic nerve. In the Optic Neuritis Treatment Trial (ONTT), the mean age of patients was 32 years, 77% were female, 92% had ocular pain (especially with eye movements), and 35% had optic disc swelling. In most patients, the demyelinating event was retrobulbar and the ocular fundus appeared normal on initial examination (Fig. 39-10), although optic disc pallor slowly developed over subsequent months.

Virtually all patients experience a gradual recovery of vision after a single episode of optic neuritis, even without treatment. This rule is so reliable that failure of vision to improve after a first attack of optic neuritis casts doubt on the original diagnosis. Treatment with high-dose IV methylprednisolone (250 mg every 6 h for 3 days) followed by oral prednisone (1 mg/kg per day for 11 days) makes no difference in ultimate acuity 6 months after the attack, but the recovery of visual function occurs more rapidly. Therefore, when visual loss is severe (worse than 20/100), IV followed by PO glucocorticoids are often recommended.

For some patients, optic neuritis remains an isolated event. However, the ONTT showed that the 15-year cumulative probability of developing clinically definite multiple sclerosis after optic neuritis is 50%. A brain magnetic resonance (MR) scan is advisable in every patient with a first attack of optic neuritis. If two or more plaques are present on initial imaging, treatment should be considered to prevent the development of additional demyelinating lesions (Chap. 458).

LEBER'S HEREDITARY OPTIC NEUROPATHY

This disease usually affects young men, causing gradual, painless, severe central visual loss in one eye, followed weeks to years later by the same process in the other eye. Acutely, the optic disc appears mildly plethoric with surface capillary telangiectasias but no vascular leakage on fluorescein angiography. Eventually optic atrophy ensues. Leber's optic neuropathy is caused by a point mutation at codon 11778 in the mitochondrial gene encoding nicotinamide adenine dinucleotide

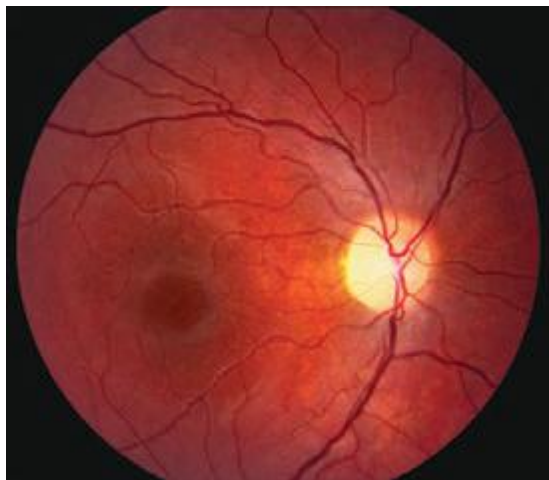


FIGURE 39-11 Optic atrophy is not a specific diagnosis but refers to the combination of optic disc pallor, arteriolar narrowing, and nerve fiber layer destruction produced by a host of eye diseases, especially optic neuropathies.

dehydrogenase (NADH) subunit 4. Additional mutations responsible for the disease have been identified, most in mitochondrial genes that encode proteins involved in electron transport. Mitochondrial mutations that cause Leber's neuropathy are inherited from the mother by all her children, but usually only sons develop symptoms.

Toxic Optic Neuropathy This can result in acute visual loss with bilateral optic disc swelling and central or cecentral scotomas. Such cases have been reported to result from exposure to ethambutol, methyl alcohol (moonshine), ethylene glycol (antifreeze), or carbon monoxide. In toxic optic neuropathy, visual loss also can develop gradually and produce optic atrophy (Fig. 39-11) without a phase of acute optic disc edema. Many agents have been implicated as a cause of toxic optic neuropathy, but the evidence supporting the association for many is weak. The following is a partial list of potential offending drugs or toxins: disulfiram, ethchlorvynol, chloramphenicol, amiodarone, monoclonal anti-CD3 antibody, ciprofloxacin, digitalis, streptomycin, lead, arsenic, thallium, D-penicillamine, isoniazid, emetine, sildenafil, tadalafil, vardenafil, and sulfonamides. Deficiency states induced by starvation, malabsorption, or alcoholism can lead to insidious visual loss. Thiamine, vitamin B₁₂, and folate levels should be checked in any patient with unexplained bilateral central scotomas and optic pallor.

Papilledema This connotes bilateral optic disc swelling from raised intracranial pressure (Fig. 39-12). Headache is a common but not



FIGURE 39-12 Papilledema means optic disc edema from raised intracranial pressure. This young woman developed acute papilledema, with hemorrhages and cotton-wool spots, as a rare side effect of treatment with tetracycline for acne.

invariable accompaniment. All other forms of optic disc swelling (e.g., from optic neuritis or ischemic optic neuropathy) should be called "optic disc edema". This convention is arbitrary but serves to avoid confusion. Often it is difficult to differentiate papilledema from other forms of optic disc edema by fundus examination alone. Transient visual obscurations are a classic symptom of papilledema. They can occur in only one eye or simultaneously in both eyes. They usually last seconds but can persist longer. Obscurations follow abrupt shifts in posture or happen spontaneously. When obscurations are prolonged or spontaneous, the papilledema is more threatening. Visual acuity is not affected by papilledema unless the papilledema is severe, long-standing, or accompanied by macular edema and hemorrhage. Visual field testing shows enlarged blind spots and peripheral constriction (Fig. 39-3F). With unremitting papilledema, peripheral visual field loss progresses in an insidious fashion while the optic nerve develops atrophy. In this setting, reduction of optic disc swelling is an ominous sign of a dying nerve rather than an encouraging indication of resolving papilledema.

Evaluation of papilledema requires neuroimaging to exclude an intracranial lesion. MR angiography is appropriate in selected cases to search for a dural venous sinus occlusion or an arteriovenous shunt. If neuroradiologic studies are negative, the subarachnoid opening pressure should be measured by lumbar puncture. An elevated pressure, with normal cerebrospinal fluid, points by exclusion to the diagnosis of *pseudotumor cerebri* (idiopathic intracranial hypertension). The majority of patients are young, female, and obese. Treatment with a carbonic anhydrase inhibitor such as acetazolamide lowers intracranial pressure by reducing the production of cerebrospinal fluid. Weight reduction is vital: bariatric surgery should be considered in patients who cannot lose weight by diet control. If vision loss is severe or progressive, a shunt should be performed without delay to prevent blindness. Occasionally, emergency surgery is required for sudden blindness caused by fulminant papilledema.

Optic Disc Drusen These are refractile deposits within the substance of the optic nerve head (Fig. 39-13). They are unrelated to drusen of the retina, which occur in age-related macular degeneration. Optic disc drusen are most common in people of northern European descent. Their diagnosis is obvious when they are visible as glittering particles on the surface of the optic disc. However, in many patients they are hidden beneath the surface, producing pseudopapilledema. It is important to recognize optic disc drusen to avoid an unnecessary evaluation for papilledema. Ultrasound or computed tomography (CT) scanning is sensitive for detection of buried optic disc drusen because they contain calcium. In most patients, optic disc drusen are an incidental, innocuous finding, but they can produce visual obscurations. On perimetry they give rise to enlarged blind spots and arcuate

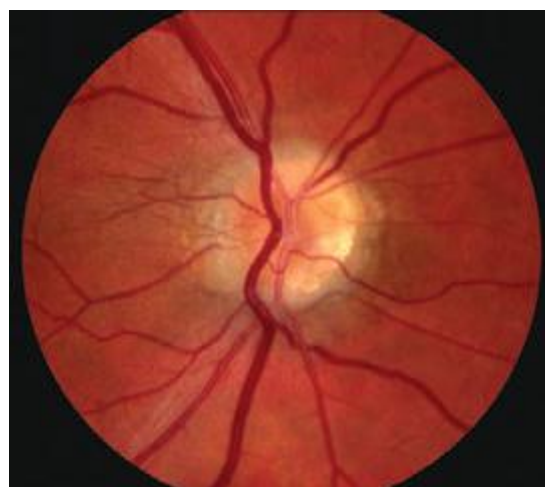


FIGURE 39-13 Optic disc drusen are calcified, mulberry-like deposits of unknown etiology within the optic disc, giving rise to "pseudopapilledema."

scotomas from damage to the optic disc. With increasing age, drusen tend to become more exposed on the disc surface as optic atrophy develops. Hemorrhage, choroidal neovascular membrane, and AION are more likely to occur in patients with optic disc drusen. No treatment is available.

Vitreous Degeneration This occurs in all individuals with advancing age, leading to visual symptoms. Opacities develop in the vitreous, casting annoying shadows on the retina. As the eye moves, these distracting “floaters” move synchronously, with a slight lag caused by inertia of the vitreous gel. Vitreous traction on the retina causes mechanical stimulation, resulting in perception of flashing lights. This photopsia is brief and is confined to one eye, in contrast to the bilateral, prolonged scintillations of cortical migraine. Contraction of the vitreous can result in sudden separation from the retina, heralded by an alarming shower of floaters and photopsia. This process, known as *vitreous detachment*, is a common involuntional event in the elderly. It is not harmful unless it damages the retina. A careful examination of the dilated fundus is important in any patient complaining of floaters or photopsia to search for peripheral tears or holes. If such a lesion is found, laser application can forestall a retinal detachment. Occasionally a tear ruptures a retinal blood vessel, causing vitreous hemorrhage and sudden loss of vision. On attempted ophthalmoscopy the fundus is hidden by a dark haze of blood. Ultrasound is required to examine the interior of the eye for a retinal tear or detachment. If the hemorrhage does not resolve spontaneously, the vitreous can be removed surgically. Vitreous hemorrhage also results from the fragile neovascular vessels that proliferate on the surface of the retina in diabetes, sickle cell anemia, and other ischemic ocular diseases.

Retinal Detachment This produces symptoms of floaters, flashing lights, and a scotoma in the peripheral visual field corresponding to the detachment (Fig. 39-14). If the detachment includes the fovea, there is an afferent pupil defect and the visual acuity is reduced. In most eyes, retinal detachment starts with a hole, flap, or tear in the peripheral retina (rhegmatogenous retinal detachment). Patients with peripheral retinal thinning (lattice degeneration) are particularly vulnerable to this process. Once a break has developed in the retina, liquefied vitreous is free to enter the subretinal space, separating the retina from the pigment epithelium. The combination of vitreous traction on the retinal surface and passage of fluid behind the retina leads inexorably to detachment. Patients with a history of myopia, trauma, or prior cataract extraction are at greatest risk for retinal detachment. The diagnosis is confirmed by ophthalmoscopic examination of the dilated eye.

Classic Migraine (See also Chap. 447) This usually occurs with a visual aura lasting about 20 min. In a typical attack, a small central disturbance in the field of vision marches toward the periphery, leaving a transient

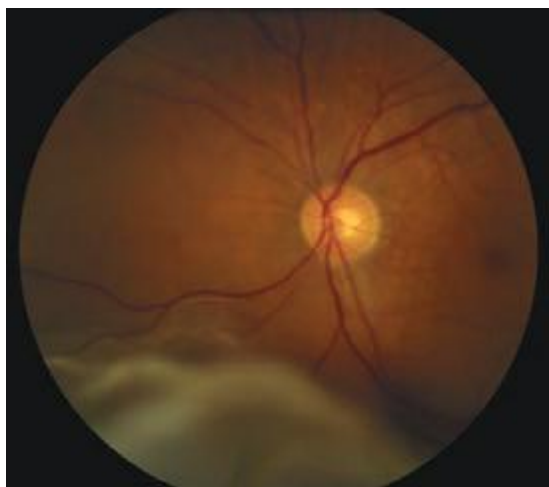


FIGURE 39-14 Retinal detachment appears as an elevated sheet of retinal tissue with folds. In this patient, the fovea was spared, so acuity was normal, but an inferior detachment produced a superior scotoma.

scotoma in its wake. The expanding border of migraine scotoma has a scintillating, dancing, or zigzag edge, resembling the bastions of a fortified city, hence the term *fortification spectra*. Patients' descriptions of fortification spectra vary widely and can be confused with amaurosis fugax. Migraine patterns usually last longer and are perceived in both eyes, whereas amaurosis fugax is briefer and occurs in only one eye. Migraine phenomena also remain visible in the dark or with the eyes closed. Generally they are confined to either the right or the left visual hemifield, but sometimes both fields are involved simultaneously. Patients often have a long history of stereotypic attacks. After the visual symptoms recede, headache develops in most patients.

Transient Ischemic Attacks Vertebrobasilar insufficiency may result in acute homonymous visual symptoms. Many patients mistakenly describe symptoms in the left or right eye when in fact the symptoms are occurring in the left or right hemifield of both eyes. Interruption of blood supply to the visual cortex causes a sudden fogging or graying of vision, occasionally with flashing lights or other positive phenomena that mimic migraine. Cortical ischemic attacks are briefer in duration than migraine, occur in older patients, and are not followed by headache. There may be associated signs of brainstem ischemia, such as diplopia, vertigo, numbness, weakness, and dysarthria.

Stroke Stroke occurs when interruption of blood supply from the posterior cerebral artery to the visual cortex is prolonged. The only finding on examination is a homonymous visual field defect that stops abruptly at the vertical meridian. Occipital lobe stroke usually is due to thrombotic occlusion of the vertebrobasilar system, embolus, or dissection. Lobar hemorrhage, tumor, abscess, and arteriovenous malformation are other common causes of hemianopic cortical visual loss.

Factitious (Functional, Nonorganic) Visual Loss This is claimed by hysterics or malingerers. The latter account for the vast majority, seeking sympathy, special treatment, or financial gain by feigning loss of sight. The diagnosis is suspected when the history is atypical, physical findings are lacking or contradictory, inconsistencies emerge on testing, and a secondary motive can be identified. In our litigious society, the fraudulent pursuit of recompense has spawned an epidemic of factitious visual loss.

CHRONIC VISUAL LOSS

Cataract Cataract is a clouding of the lens sufficient to reduce vision. Most cataracts develop slowly as a result of aging, leading to gradual impairment of vision. The formation of cataract occurs more rapidly in patients with a history of ocular trauma, uveitis, or diabetes mellitus. Cataracts are acquired in a variety of genetic diseases, such as myotonic dystrophy, neurofibromatosis type 2, and galactosemia. Radiation therapy and glucocorticoid treatment can induce cataract as a side effect. The cataracts associated with radiation or glucocorticoids have a typical posterior subcapsular location. Cataract can be detected by noting an impaired red reflex when viewing light reflected from the fundus with an ophthalmoscope or by examining the dilated eye with the slit lamp.

The only treatment for cataract is surgical extraction of the opacified lens. Millions of cataract operations are performed each year around the globe. The operation generally is done under local anesthesia on an outpatient basis. A plastic or silicone intraocular lens is placed within the empty lens capsule in the posterior chamber, substituting for the natural lens and leading to rapid recovery of sight. More than 95% of patients who undergo cataract extraction can expect an improvement in vision. In some patients, the lens capsule remaining in the eye after cataract extraction eventually turns cloudy, causing secondary loss of vision. A small opening, called a posterior capsulotomy, is made in the lens capsule with a laser to restore clarity.

Glaucoma Glaucoma is a slowly progressive, insidious optic neuropathy that usually is associated with chronic elevation of intraocular pressure. After cataract, it is the most common cause of blindness in the world. It is especially prevalent in people of African descent. The mechanism by which raised intraocular pressure injures the optic



FIGURE 39-15 Glaucoma results in “cupping” as the neural rim is destroyed and the central cup becomes enlarged and excavated. The cup-to-disc ratio is about 0.8 in this patient.

nerve is not understood. Axons entering the inferotemporal and superotemporal aspects of the optic disc are damaged first, producing typical nerve fiber bundle or arcuate scotomas on perimetric testing. As fibers are destroyed, the neural rim of the optic disc shrinks and the physiologic cup within the optic disc enlarges (Fig. 39-15). This process is referred to as pathologic “cupping.” The cup-to-disc diameter is expressed as a fraction (e.g., 0.2). The cup-to-disc ratio ranges widely in normal individuals, making it difficult to diagnose glaucoma reliably simply by observing an unusually large or deep optic cup. Careful documentation of serial examinations is helpful. In a patient with physiologic cupping the large cup remains stable, whereas in a patient with glaucoma it expands relentlessly over the years. Observation of progressive cupping and detection of an arcuate scotoma or a nasal step on computerized visual field testing is sufficient to establish the diagnosis of glaucoma. Optical coherence tomography reveals corresponding loss of fibers along the arcuate pathways in the nerve fiber layer.

About 95% of patients with glaucoma have open anterior chamber angles. In most affected individuals the intraocular pressure is elevated. The cause of elevated intraocular pressure is unknown, but it is associated with gene mutations in the heritable forms. Surprisingly, a third of patients with open-angle glaucoma have an intraocular pressure within the normal range of 10–20 mmHg. For this so-called normal or low-tension form of glaucoma, high myopia is a risk factor.

Chronic angle-closure glaucoma and chronic open-angle glaucoma are usually asymptomatic. Only acute angle-closure glaucoma causes a red or painful eye, from abrupt elevation of intraocular pressure. In all forms of glaucoma, foveal acuity is spared until end-stage disease is reached. For these reasons, severe and irreversible damage can occur before either the patient or the physician recognizes the diagnosis. Screening of patients for glaucoma by noting the cup-to-disc ratio on ophthalmoscopy and by measuring intraocular pressure is vital. Glaucoma is treated with topical adrenergic agonists, cholinergic agonists, beta blockers, and prostaglandin analogues. Occasionally, systemic absorption of beta blocker from eyedrops can be sufficient to cause side effects of bradycardia, hypotension, heart block, bronchospasm, or depression. Topical or oral carbonic anhydrase inhibitors are used to lower intraocular pressure by reducing aqueous production. Laser treatment of the trabecular meshwork in the anterior chamber angle improves aqueous outflow from the eye. If medical or laser treatments fail to halt optic nerve damage from glaucoma, a filter must be constructed surgically (trabeculectomy) or a drainage device placed to release aqueous from the eye in a controlled fashion.

Macular Degeneration This is a major cause of gradual, painless, bilateral central visual loss in the elderly. It occurs in a nonexudative (dry) form and an exudative (wet) form. Inflammation may be important in both forms of macular degeneration; susceptibility is associated

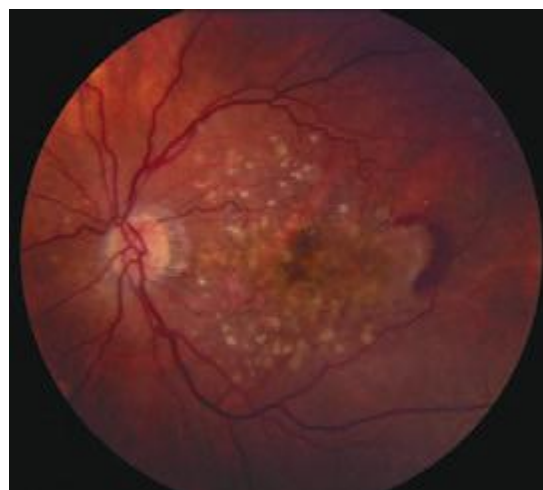


FIGURE 39-16 Age-related macular degeneration consisting of scattered yellow drusen in the macula (dry form) and a crescent of fresh hemorrhage temporal to the fovea from a subretinal neovascular membrane (wet form).

with variants in the gene for complement factor H, an inhibitor of the alternative complement pathway. The nonexudative process begins with the accumulation of extracellular deposits called drusen underneath the retinal pigment epithelium. On ophthalmoscopy, they are pleomorphic but generally appear as small discrete yellow lesions clustered in the macula (Fig. 39-16). With time they become larger, more numerous, and confluent. The retinal pigment epithelium becomes focally detached and atrophic, causing visual loss by interfering with photoreceptor function. Treatment with vitamins C and E, beta-carotene, and zinc may retard dry macular degeneration.

Exudative macular degeneration, which develops in only a minority of patients, occurs when neovascular vessels from the choroid grow through defects in Bruch’s membrane and proliferate underneath the retinal pigment epithelium or the retina. Leakage from these vessels produces elevation of the retina, with distortion (metamorphopsia) and blurring of vision. Although the onset of these symptoms is usually gradual, bleeding from a subretinal choroidal neovascular membrane sometimes causes acute visual loss. Neovascular membranes can be difficult to see on fundus examination because they are located beneath the retina. Fluorescein angiography and optical coherence tomography, a technique for acquiring images of the retina in cross-section, are extremely useful for their detection. Major or repeated hemorrhage under the retina from neovascular membranes results in fibrosis, development of a round (disciform) macular scar, and permanent loss of central vision.

A major therapeutic advance has occurred with the discovery that exudative macular degeneration can be treated with intraocular injection of antagonists to vascular endothelial growth factor. Bevacizumab, ranibizumab, or aflibercept is administered by direct injection into the vitreous cavity, beginning on a monthly basis. These antibodies cause the regression of neovascular membranes by blocking the action of vascular endothelial growth factor, thereby improving visual acuity.

Central Serous Chorioretinopathy This primarily affects males between the ages of 20 and 50 years. Leakage of serous fluid from the choroid causes small, localized detachment of the retinal pigment epithelium and the neurosensory retina. These detachments produce acute or chronic symptoms of metamorphopsia and blurred vision when the macula is involved. They are difficult to visualize with a direct ophthalmoscope because the detached retina is transparent and only slightly elevated. Optical coherence tomography shows fluid beneath the retina, and fluorescein angiography shows dye streaming into the subretinal space. The cause of central serous chorioretinopathy is unknown. Symptoms may resolve spontaneously if the retina reattaches, but recurrent detachment is common. Laser photocoagulation has benefited some patients with this condition.

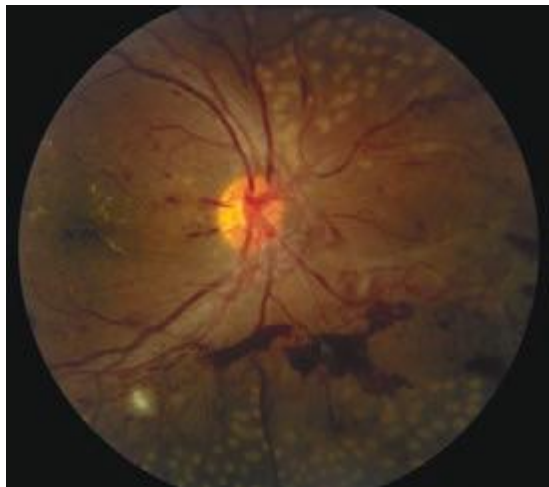


FIGURE 39-17 Proliferative diabetic retinopathy in a 25-year-old man with an 18-year history of diabetes, showing neovascular vessels emanating from the optic disc, retinal and vitreous hemorrhage, cotton-wool spots, and macular exudate. Round spots in the periphery represent recently applied panretinal photocoagulation.

Diabetic Retinopathy A rare disease until 1921, when the discovery of insulin resulted in a dramatic improvement in life expectancy for patients with diabetes mellitus, diabetic retinopathy is now a leading cause of blindness in the United States. The retinopathy takes years to develop but eventually appears in nearly all cases. Regular surveillance of the dilated fundus is crucial for any patient with diabetes. In advanced diabetic retinopathy, the proliferation of neovascular vessels leads to blindness from vitreous hemorrhage, retinal detachment, and glaucoma (Fig. 39-17). These complications can be avoided in most patients by administration of panretinal laser photocoagulation at the appropriate point in the evolution of the disease. **For further discussion of the manifestations and management of diabetic retinopathy, see Chaps. 417–419.**

Retinitis Pigmentosa This is a general term for a disparate group of rod-cone dystrophies characterized by progressive night blindness, visual field constriction with a ring scotoma, loss of acuity, and an abnormal electroretinogram (ERG). It occurs sporadically or in an autosomal recessive, dominant, or X-linked pattern. Irregular black deposits of clumped pigment in the peripheral retina, called *bone spicules* because of their vague resemblance to the spicules of cancellous bone, give the disease its name (Fig. 39-18). The name is actually a



FIGURE 39-18 Retinitis pigmentosa with black clumps of pigment known as “bone spicules.” The patient had peripheral visual field loss with sparing of central (macular) vision.

misnomer because retinitis pigmentosa is not an inflammatory process. Most cases are due to a mutation in the gene for rhodopsin, the rod photopigment, or in the gene for peripherin, a glycoprotein located in photoreceptor outer segments. Vitamin A (15,000 IU/d) slightly retards the deterioration of the ERG in patients with retinitis pigmentosa but has no beneficial effect on visual acuity or fields.

Leber’s congenital amaurosis, a rare cone dystrophy, has been treated by replacement of the missing RPE65 protein through gene therapy, resulting in modest improvement in visual function. Some forms of retinitis pigmentosa occur in association with rare, hereditary systemic diseases (olivopontocerebellar degeneration, Bassen-Kornzweig disease, Kearns-Sayre syndrome, Refsum’s disease). Chronic treatment with chloroquine, hydroxychloroquine, and phenothiazines (especially thioridazine) can produce visual loss from a toxic retinopathy that resembles retinitis pigmentosa.

Epiretinal Membrane This is a fibrocellular tissue that grows across the inner surface of the retina, causing metamorphopsia and reduced visual acuity from distortion of the macula. A crinkled, cellophane-like membrane is visible on the retinal examination. Epiretinal membrane is most common in patients over 50 years of age and is usually unilateral. Most cases are idiopathic, but some occur as a result of hypertensive retinopathy, diabetes, retinal detachment, or trauma. When visual acuity is reduced to the level of about 6/24 (20/80), vitrectomy and surgical peeling of the membrane to relieve macular puckering are recommended. Contraction of an epiretinal membrane sometimes gives rise to a *macular hole*. Most macular holes, however, are caused by local vitreous traction within the fovea. Vitrectomy can improve acuity in selected cases.

Melanoma and Other Tumors Melanoma is the most common primary tumor of the eye (Fig. 39-19). It causes photopsia, an enlarging scotoma, and loss of vision. A small melanoma is often difficult to differentiate from a benign choroidal nevus. Serial examinations are required to document a malignant pattern of growth. Treatment of melanoma is controversial. Options include enucleation, local resection, and irradiation. *Metastatic tumors* to the eye outnumber primary tumors. Breast and lung carcinomas have a special propensity to spread to the choroid or iris. Leukemia and lymphoma also commonly invade ocular tissues. Sometimes their only sign on eye examination is cellular debris in the vitreous, which can masquerade as a chronic posterior uveitis. *Retrolbulbar tumor* of the optic nerve (meningioma, glioma) or *chiasmal tumor* (pituitary adenoma, meningioma) produces



FIGURE 39-19 Melanoma of the choroid, appearing as an elevated dark mass in the inferior fundus, with overlying hemorrhage. The black line denotes the plane of the optical coherence tomography scan (*below*) showing the subretinal tumor.

gradual visual loss with few objective findings except for optic disc pallor. Rarely, sudden expansion of a pituitary adenoma from infarction and bleeding (*pituitary apoplexy*) causes acute retrobulbar visual loss, with headache, nausea, and ocular motor nerve palsies. In any patient with visual field loss or optic atrophy, CT or MR scanning should be considered if the cause remains unknown after careful review of the history and thorough examination of the eye.

PROPTOSIS

When the globes appear asymmetric, the clinician must first decide which eye is abnormal. Is one eye recessed within the orbit (*enophthalmos*), or is the other eye protuberant (*exophthalmos*, or *proptosis*)? A small globe or a Horner's syndrome can give the appearance of enophthalmos. True enophthalmos occurs commonly after trauma, from atrophy of retrobulbar fat, or from fracture of the orbital floor. The position of the eyes within the orbits is measured by using a Hertel exophthalmometer, a handheld instrument that records the position of the anterior corneal surface relative to the lateral orbital rim. If this instrument is not available, relative eye position can be judged by bending the patient's head forward and looking down upon the orbits. A proptosis of only 2 mm in one eye is detectable from this perspective. The development of proptosis implies a space-occupying lesion in the orbit and usually warrants CT or MR imaging.

Graves' Ophthalmopathy This is the leading cause of proptosis in adults (Chap. 405). The proptosis is often asymmetric and can even appear to be unilateral. Orbital inflammation and engorgement of the extraocular muscles, particularly the medial rectus and the inferior rectus, account for the protrusion of the globe. Corneal exposure, lid retraction, conjunctival injection, restriction of gaze, diplopia, and visual loss from optic nerve compression are cardinal symptoms. Graves' eye disease is a clinical diagnosis, but laboratory testing can be useful. The serum level of thyroid-stimulating immunoglobulins is often elevated. Orbital imaging usually reveals enlarged extraocular eye muscles, but not always. Graves' ophthalmopathy can be treated with oral prednisone (60 mg/d) for 1 month, followed by a taper over several months. Worsening of symptoms upon glucocorticoid withdrawal is common. Topical lubricants, taping the eyelids closed at night, moisture chambers, and eyelid surgery are helpful to limit exposure of ocular tissues. Radiation therapy is not effective. Orbital decompression should be performed for severe, symptomatic exophthalmos or if visual function is reduced by optic nerve compression. In patients with diplopia, prisms or eye muscle surgery can be used to restore ocular alignment in primary gaze.

Orbital Pseudotumor This is an idiopathic, inflammatory orbital syndrome that is distinguished from Graves' ophthalmopathy by the prominent complaint of pain. Other symptoms include diplopia, ptosis, proptosis, and orbital congestion. Evaluation for sarcoidosis, granulomatosis with polyangiitis, and other types of orbital vasculitis or collagen-vascular disease is negative. Imaging often shows swollen eye muscles (orbital myositis) with enlarged tendons. By contrast, in Graves' ophthalmopathy, the tendons of the eye muscles usually are spared. The Tolosa-Hunt syndrome (Chap. 455) may be regarded as an extension of orbital pseudotumor through the superior orbital fissure into the cavernous sinus. The diagnosis of orbital pseudotumor is difficult. Biopsy of the orbit frequently yields nonspecific evidence of fat infiltration by lymphocytes, plasma cells, and eosinophils. A dramatic response to a therapeutic trial of systemic glucocorticoids indirectly provides the best confirmation of the diagnosis.

Orbital Cellulitis This causes pain, lid erythema, proptosis, conjunctival chemosis, restricted motility, decreased acuity, afferent pupillary defect, fever, and leukocytosis. It often arises from the paranasal sinuses, especially by contiguous spread of infection from the ethmoid sinus through the lamina papyracea of the medial orbit. A history of recent upper respiratory tract infection, chronic sinusitis, thick mucus secretions, or dental disease is significant in any patient with suspected orbital cellulitis. Blood cultures should be obtained, but they are usually negative. Most patients respond to empirical therapy with broad-spectrum

IV antibiotics. Occasionally, orbital cellulitis follows an overwhelming course, with massive proptosis, blindness, septic cavernous sinus thrombosis, and meningitis. To avert this disaster, orbital cellulitis should be managed aggressively in the early stages, with immediate imaging of the orbits and antibiotic therapy that includes coverage of methicillin-resistant *Staphylococcus aureus* (MRSA). Prompt surgical drainage of an orbital abscess or paranasal sinusitis is indicated if optic nerve function deteriorates despite antibiotics.

Tumors Tumors of the orbit cause painless, progressive proptosis. The most common primary tumors are cavernous hemangioma, lymphangioma, neurofibroma, schwannoma, dermoid cyst, adenoid cystic carcinoma, optic nerve glioma, optic nerve meningioma, and benign mixed tumor of the lacrimal gland. Metastatic tumor to the orbit occurs frequently in breast carcinoma, lung carcinoma, and lymphoma. Diagnosis by fine-needle aspiration followed by urgent radiation therapy sometimes can preserve vision.

Carotid Cavernous Fistulas With anterior drainage through the orbit, these fistulas produce proptosis, diplopia, glaucoma, and corkscrew, arterialized conjunctival vessels. Direct fistulas usually result from trauma. They are easily diagnosed because of the prominent signs produced by high-flow, high-pressure shunting. Indirect fistulas, or dural arteriovenous malformations, are more likely to occur spontaneously, especially in older women. The signs are more subtle, and the diagnosis frequently is missed. The combination of slight proptosis, diplopia, enlarged muscles, and an injected eye often is mistaken for thyroid ophthalmopathy. A bruit heard upon auscultation of the head or reported by the patient is a valuable diagnostic clue. Imaging shows an enlarged superior ophthalmic vein in the orbits. Carotid cavernous shunts can be eliminated by intravascular embolization.

PTOSIS

Blepharoptosis This is an abnormal drooping of the eyelid. Unilateral or bilateral ptosis can be congenital, from dysgenesis of the levator palpebrae superioris, or from abnormal insertion of its aponeurosis into the eyelid. Acquired ptosis can develop so gradually that the patient is unaware of the problem. Inspection of old photographs is helpful in dating the onset. A history of prior trauma, eye surgery, contact lens use, diplopia, systemic symptoms (e.g., dysphagia or peripheral muscle weakness), or a family history of ptosis should be sought. Fluctuating ptosis that worsens late in the day is typical of myasthenia gravis. Examination should focus on evidence for proptosis, eyelid masses or deformities, inflammation, pupil inequality, or limitation of motility. The width of the palpebral fissures is measured in primary gaze to determine the degree of ptosis. The ptosis will be underestimated if the patient compensates by lifting the brow with the frontalis muscle.

Mechanical Ptosis This occurs in many elderly patients from stretching and redundancy of eyelid skin and subcutaneous fat (dermatochalasis). The extra weight of these sagging tissues causes the lid to droop. Enlargement or deformation of the eyelid from infection, tumor, trauma, or inflammation also results in ptosis on a purely mechanical basis.

Aponeurotic Ptosis This is an acquired dehiscence or stretching of the aponeurotic tendon, which connects the levator muscle to the tarsal plate of the eyelid. It occurs commonly in older patients, presumably from loss of connective tissue elasticity. Aponeurotic ptosis is also a common sequela of eyelid swelling from infection or blunt trauma to the orbit, cataract surgery, or contact lens use.

Myogenic Ptosis The causes of *myogenic ptosis* include myasthenia gravis (Chap. 461) and a number of rare myopathies that manifest with ptosis. The term *chronic progressive external ophthalmoplegia* refers to a spectrum of systemic diseases caused by mutations of mitochondrial DNA. As the name implies, the most prominent findings are symmetric, slowly progressive ptosis and limitation of eye movements. In general, diplopia is a late symptom because all eye movements are reduced equally. In the *Kearns-Sayre* variant, retinal pigmentary changes and abnormalities of cardiac conduction develop.

Peripheral muscle biopsy shows characteristic “ragged-red fibers.” *Oculopharyngeal dystrophy* is a distinct autosomal dominant disease with onset in middle age, characterized by ptosis, limited eye movements, and trouble swallowing. *Myotonic dystrophy*, another autosomal dominant disorder, causes ptosis, ophthalmoparesis, cataract, and pigmentary retinopathy. Patients have muscle wasting, myotonia, frontal balding, and cardiac abnormalities.

Neurogenic Ptosis This results from a lesion affecting the innervation to either of the two muscles that open the eyelid: Müller’s muscle or the levator palpebrae superioris. Examination of the pupil helps distinguish between these two possibilities. In Horner’s syndrome, the eye with ptosis has a smaller pupil and the eye movements are full. In an oculomotor nerve palsy, the eye with the ptosis has a larger or a normal pupil. If the pupil is normal but there is limitation of adduction, elevation, and depression, a pupil-sparing oculomotor nerve palsy is likely (see next section). Rarely, a lesion affecting the small, central subnucleus of the oculomotor complex will cause bilateral ptosis with normal eye movements and pupils.

DOUBLE VISION (DIPLOPIA)

The first point to clarify is whether diplopia persists in either eye after the opposite eye is covered. If it does, the diagnosis is monocular diplopia. The cause is usually intrinsic to the eye and therefore has no dire implications for the patient. Corneal aberrations (e.g., keratoconus, pterygium), uncorrected refractive error, cataract, or foveal traction may give rise to monocular diplopia. Occasionally it is a symptom of malingering or psychiatric disease. Diplopia alleviated by covering one eye is binocular diplopia and is caused by disruption of ocular alignment. Inquiry should be made into the nature of the double vision (purely side-by-side versus partial vertical displacement of images), mode of onset, duration, intermittency, diurnal variation, and associated neurologic or systemic symptoms. If the patient has diplopia while being examined, motility testing should reveal a deficiency corresponding to the patient’s symptoms. However, subtle limitation of ocular excursions is often difficult to detect. For example, a patient with a slight left abducens nerve paresis may appear to have full eye movements despite a complaint of horizontal diplopia upon looking to the left. In this situation, the cover test provides a more sensitive method for demonstrating the ocular misalignment. It should be conducted in primary gaze and then with the head turned and tilted in each direction. In the above example, a cover test with the head turned to the right will maximize the fixation shift evoked by the cover test.

Occasionally, a cover test performed in an asymptomatic patient during a routine examination will reveal an ocular deviation. If the eye movements are full and the ocular misalignment is equal in all directions of gaze (concomitant deviation), the diagnosis is strabismus. In this condition, which affects about 1% of the population, fusion is disrupted in infancy or early childhood. To avoid diplopia, vision is suppressed from the nonfixating eye. In some children, this leads to impaired vision (amblyopia, or “lazy” eye) in the deviated eye.

Binocular diplopia results from a wide range of processes: infectious, neoplastic, metabolic, degenerative, inflammatory, and vascular. One must decide whether the diplopia is neurogenic in origin or is due to restriction of globe rotation by local disease in the orbit. Orbital pseudotumor, myositis, infection, tumor, thyroid disease, and muscle entrapment (e.g., from a blowout fracture) cause restrictive diplopia. The diagnosis of restriction is usually made by recognizing other associated signs and symptoms of local orbital disease. Omission of high-resolution orbital imaging is a common mistake in the evaluation of diplopia.

Myasthenia Gravis (See also Chap. 461) This is a major cause of diplopia. The diplopia is often intermittent, variable, and not confined to any single ocular motor nerve distribution. The pupils are always normal. Fluctuating ptosis may be present. Many patients have a purely ocular form of the disease, with no evidence of systemic muscular weakness. The diagnosis can be confirmed by an IV edrophonium injection, which produces a transient reversal of eyelid or eye muscle

weakness. Blood tests for antibodies against the acetylcholine receptor or the MuSK protein can establish the diagnosis but are frequently negative in the purely ocular form of myasthenia gravis. *Botulism* from food or wound poisoning can mimic ocular myasthenia.

After restrictive orbital disease and myasthenia gravis are excluded, a lesion of a cranial nerve supplying innervation to the extraocular muscles is the most likely cause of binocular diplopia.

Oculomotor Nerve The third cranial nerve innervates the medial, inferior, and superior recti; inferior oblique; levator palpebrae superioris; and the iris sphincter. Total palsy of the oculomotor nerve causes ptosis, a dilated pupil, and leaves the eye “down and out” because of the unopposed action of the lateral rectus and superior oblique. This combination of findings is obvious. More challenging is the diagnosis of early or partial oculomotor nerve palsy. In this setting any combination of ptosis, pupil dilation, and weakness of the eye muscles supplied by the oculomotor nerve may be encountered. Frequent serial examinations during the evolving phase of the palsy help ensure that the diagnosis is not missed. The advent of an oculomotor nerve palsy with a pupil involvement, especially when accompanied by pain, suggests a compressive lesion, such as a tumor or circle of Willis aneurysm. Neuroimaging should be obtained, along with a CT or MR angiogram. Occasionally, a catheter arteriogram must be done to exclude an aneurysm.

A lesion of the oculomotor nucleus in the rostral midbrain produces signs that differ from those caused by a lesion of the nerve itself. There is bilateral ptosis because the levator muscle is innervated by a single central subnucleus. There is also weakness of the contralateral superior rectus, because it is supplied by the oculomotor nucleus on the other side. Occasionally both superior recti are weak. Isolated nuclear oculomotor palsy is rare. Usually neurologic examination reveals additional signs that suggest brainstem damage from infarction, hemorrhage, tumor, or infection.

Injury to structures surrounding fascicles of the oculomotor nerve descending through the midbrain has given rise to a number of classic eponymic designations. In *Nothnagel’s syndrome*, injury to the superior cerebellar peduncle causes ipsilateral oculomotor palsy and contralateral cerebellar ataxia. In *Benedikt’s syndrome*, injury to the red nucleus results in ipsilateral oculomotor palsy and contralateral tremor, chorea, and athetosis. *Claude’s syndrome* incorporates features of both of these syndromes, by injury to both the red nucleus and the superior cerebellar peduncle. Finally, in *Weber’s syndrome*, injury to the cerebral peduncle causes ipsilateral oculomotor palsy with contralateral hemiparesis.

In the subarachnoid space the oculomotor nerve is vulnerable to aneurysm, meningitis, tumor, infarction, and compression. In cerebral herniation, the nerve becomes trapped between the edge of the tentorium and the uncus of the temporal lobe. Oculomotor palsy also can result from midbrain torsion and hemorrhages during herniation. In the cavernous sinus, oculomotor palsy arises from carotid aneurysm, carotid cavernous fistula, cavernous sinus thrombosis, tumor (pituitary adenoma, meningioma, metastasis), herpes zoster infection, and the Tolosa-Hunt syndrome.

The etiology of an isolated, pupil-sparing oculomotor palsy often remains an enigma even after neuroimaging and extensive laboratory testing. Most cases are thought to result from microvascular infarction of the nerve somewhere along its course from the brainstem to the orbit. Usually the patient complains of pain. Diabetes, hypertension, and vascular disease are major risk factors. Spontaneous recovery over a period of months is the rule. If this fails to occur or if new findings develop, the diagnosis of microvascular oculomotor nerve palsy should be reconsidered. Aberrant regeneration is common when the oculomotor nerve is injured by trauma or compression (tumor, aneurysm). Miswiring of sprouting fibers to the levator muscle and the rectus muscles results in elevation of the eyelid upon downgaze or adduction. The pupil also constricts upon attempted adduction, elevation, or depression of the globe. Aberrant regeneration is not seen after oculomotor palsy from microvascular infarct and hence vitiates that diagnosis.

Trochlear Nerve The fourth cranial nerve originates in the midbrain, just caudal to the oculomotor nerve complex. Fibers exit the brainstem dorsally and cross to innervate the contralateral superior oblique. The principal actions of this muscle are to depress and intort the globe. A palsy therefore results in hypertropia and excyclotorsion. The cyclotorsion seldom is noticed by patients. Instead, they complain of vertical diplopia, especially upon reading or looking down. The vertical diplopia also is exacerbated by tilting the head toward the side with the muscle palsy and alleviated by tilting it away. This “head tilt test” is a cardinal diagnostic feature.

Isolated trochlear nerve palsy results from all the causes listed above for the oculomotor nerve except aneurysm. The trochlear nerve is particularly apt to suffer injury after closed head trauma. The free edge of the tentorium is thought to impinge on the nerve during a concussive blow. Most isolated trochlear nerve palsies are idiopathic and hence are diagnosed by exclusion as “microvascular.” Spontaneous improvement occurs over a period of months in most patients. A base-down prism (conveniently applied to the patient’s glasses as a stick-on Fresnel lens) may serve as a temporary measure to alleviate diplopia. If the palsy does not resolve, the eyes can be realigned by weakening the inferior oblique muscle.

Abducens Nerve The sixth cranial nerve innervates the lateral rectus muscle. A palsy produces horizontal diplopia, worse on gaze to the side of the lesion. A nuclear lesion has different consequences, because the abducens nucleus contains interneurons that project via the medial longitudinal fasciculus to the medial rectus subnucleus of the contralateral oculomotor complex. Therefore, an abducens nuclear lesion produces a complete lateral gaze palsy from weakness of both the ipsilateral lateral rectus and the contralateral medial rectus. *Foville’s syndrome* after dorsal pontine injury includes lateral gaze palsy, ipsilateral facial palsy, and contralateral hemiparesis incurred by damage to descending corticospinal fibers. *Millard-Gubler syndrome* from ventral pontine injury is similar except for the eye findings. There is lateral rectus weakness only, instead of gaze palsy, because the abducens fascicle is injured rather than the nucleus. Infarct, tumor, hemorrhage, vascular malformation, and multiple sclerosis are the most common etiologies of brainstem abducens palsy.

After leaving the ventral pons, the abducens nerve runs forward along the clivus to pierce the dura at the petrous apex, where it enters the cavernous sinus. Along its subarachnoid course it is susceptible to meningitis, tumor (meningioma, chordoma, carcinomatous meningitis), subarachnoid hemorrhage, trauma, and compression by aneurysm or dolichoectatic vessels. At the petrous apex, mastoiditis can produce deafness, pain, and ipsilateral abducens palsy (*Gradenigo’s syndrome*). In the cavernous sinus, the nerve can be affected by carotid aneurysm, carotid cavernous fistula, tumor (pituitary adenoma, meningioma, nasopharyngeal carcinoma), herpes infection, and Tolosa-Hunt syndrome.

Unilateral or bilateral abducens palsy is a classic sign of raised intracranial pressure. The diagnosis can be confirmed if papilledema is observed on fundus examination. The mechanism is still debated but probably is related to rostral-caudal displacement of the brainstem. The same phenomenon accounts for abducens palsy from Chiari malformation or low intracranial pressure (e.g., after lumbar puncture, spinal anesthesia, or spontaneous dural cerebrospinal fluid leak).

Treatment of abducens palsy is aimed at prompt correction of the underlying cause. However, the cause remains obscure in many instances despite diligent evaluation. As was mentioned above for isolated trochlear or oculomotor palsy, most cases are assumed to represent microvascular infarcts because they often occur in the setting of diabetes or other vascular risk factors. Some cases may develop as a postinfectious mononeuritis (e.g., after a viral flu). Patching one eye, occluding one eyeglass lens with tape, or applying a temporary prism will provide relief of diplopia until the palsy resolves. If recovery is incomplete, eye muscle surgery nearly always can realign the eyes, at least in primary position. A patient with an abducens palsy that fails to improve should be reevaluated for an occult etiology (e.g., chordoma,

carcinomatous meningitis, carotid cavernous fistula, myasthenia gravis). Skull base tumors are easily missed even on contrast-enhanced neuroimaging studies.

Multiple Ocular Motor Nerve Palsies These should not be attributed to spontaneous microvascular events affecting more than one cranial nerve at a time. This remarkable coincidence does occur, especially in diabetic patients, but the diagnosis is made only in retrospect after all other diagnostic alternatives have been exhausted. Neuroimaging should focus on the cavernous sinus, superior orbital fissure, and orbital apex, where all three ocular motor nerves are in close proximity. In a diabetic or immunocompromised host, fungal infection (*Aspergillus*, *Mucorales*, *Cryptococcus*) is a common cause of multiple nerve palsies. In a patient with systemic malignancy, carcinomatous meningitis is a likely diagnosis. Cytologic examination may be negative despite repeated sampling of the cerebrospinal fluid. The cancer-associated Lambert-Eaton myasthenic syndrome also can produce ophthalmoplegia. Giant cell (temporal) arteritis occasionally manifests as diplopia from ischemic palsies of extraocular muscles. Fisher’s syndrome, an ocular variant of Guillain-Barré, produces ophthalmoplegia with areflexia and ataxia. Often the ataxia is mild, and the reflexes are normal. Antiganglioside antibodies (GQ1b) can be detected in about 50% of cases.

Supranuclear Disorders of Gaze These are often mistaken for multiple ocular motor nerve palsies. For example, Wernicke’s encephalopathy can produce nystagmus and a partial deficit of horizontal and vertical gaze that mimics a combined abducens and oculomotor nerve palsy. The disorder occurs in malnourished or alcoholic patients and can be reversed by thiamine. Infarct, hemorrhage, tumor, multiple sclerosis, encephalitis, vasculitis, and Whipple’s disease are other important causes of supranuclear gaze palsy. Disorders of vertical gaze, especially downward saccades, are an early feature of progressive supranuclear palsy. Smooth pursuit is affected later in the course of the disease. Parkinson’s disease, Huntington’s disease, and olivopontocerebellar degeneration also can affect vertical gaze.

The *frontal eye field* of the cerebral cortex is involved in generation of saccades to the contralateral side. After hemispheric stroke, the eyes usually deviate toward the lesioned side because of the unopposed action of the frontal eye field in the normal hemisphere. With time, this deficit resolves. Seizures generally have the opposite effect: the eyes deviate conjugately away from the irritative focus. *Parietal lesions* disrupt smooth pursuit of targets moving toward the side of the lesion. Bilateral parietal lesions produce *Bálint’s syndrome*, which is characterized by impaired eye-hand coordination (optic ataxia), difficulty initiating voluntary eye movements (ocular apraxia), and visuospatial disorientation (simultanagnosia).

Horizontal Gaze Descending cortical inputs mediating horizontal gaze ultimately converge at the level of the pons. Neurons in the paramedian pontine reticular formation are responsible for controlling conjugate gaze toward the same side. They project directly to the ipsilateral abducens nucleus. A lesion of either the paramedian pontine reticular formation or the abducens nucleus causes an ipsilateral conjugate gaze palsy. Lesions at either locus produce nearly identical clinical syndromes, with the following exception: vestibular stimulation (oculocephalic maneuver or caloric irrigation) will succeed in driving the eyes conjugately to the side in a patient with a lesion of the paramedian pontine reticular formation but not in a patient with a lesion of the abducens nucleus.

INTERNUCLEAR OPHTHALMOPLEGIA This results from damage to the medial longitudinal fasciculus ascending from the abducens nucleus in the pons to the oculomotor nucleus in the midbrain (hence, “internuclear”). Damage to fibers carrying the conjugate signal from abducens interneurons to the contralateral medial rectus motoneurons results in a failure of adduction on attempted lateral gaze. For example, a patient with a left internuclear ophthalmoplegia (INO) will have slowed or absent adducting movements of the left eye (Fig. 39-20). A patient



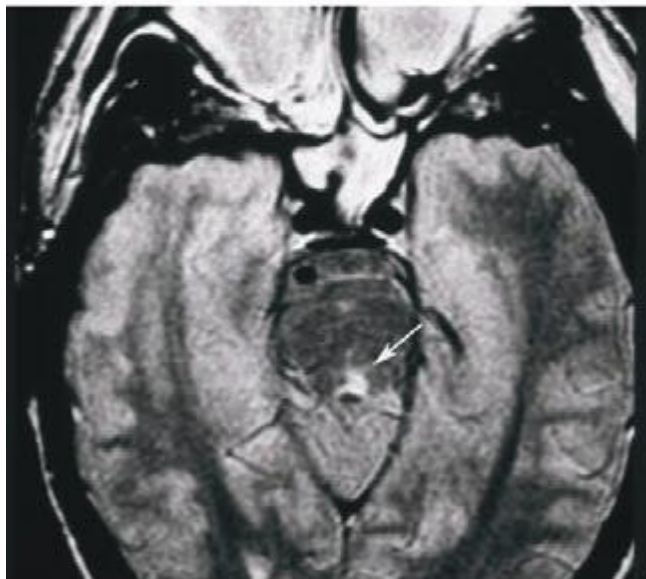
A



B



C



D

FIGURE 39-20 Left internuclear ophthalmoplegia (INO). **A.** In primary position of gaze, the eyes appear normal. **B.** Horizontal gaze to the left is intact. **C.** On attempted horizontal gaze to the right, the left eye fails to adduct. In mildly affected patients, the eye may adduct partially or more slowly than normal. Nystagmus is usually present in the abducted eye. **D.** T2-weighted axial magnetic resonance image through the pons showing a demyelinating plaque in the left medial longitudinal fasciculus (arrow).

with bilateral injury to the medial longitudinal fasciculus will have bilateral INO. Multiple sclerosis is the most common cause, although tumor, stroke, trauma, or any brainstem process may be responsible. *One-and-a-half syndrome* is due to a combined lesion of the medial longitudinal fasciculus and the abducens nucleus on the same side. The patient's only horizontal eye movement is abduction of the eye on the other side.

Vertical Gaze This is controlled at the level of the midbrain. The neuronal circuits affected in disorders of vertical gaze are not fully elucidated, but lesions of the rostral interstitial nucleus of the medial longitudinal fasciculus and the interstitial nucleus of Cajal cause supranuclear paresis of upgaze, downgaze, or all vertical eye movements. Distal basilar artery ischemia is the most common etiology. *Skew deviation* refers to a vertical misalignment of the eyes, usually constant in all positions of gaze. The finding has poor localizing value because skew deviation has been reported after lesions in widespread regions of the brainstem and cerebellum.

PARINAUD'S SYNDROME Also known as dorsal midbrain syndrome, this is a distinct supranuclear vertical gaze disorder caused by damage to the posterior commissure. It is a classic sign of hydrocephalus from aqueductal stenosis. Pineal region or midbrain tumors, cysticercosis, and stroke also cause Parinaud's syndrome. Features include loss of upgaze (and sometimes downgaze), convergence-retraction nystagmus on attempted upgaze, downward ocular deviation ("setting sun" sign), lid retraction (Collier's sign), skew deviation, pseudoabducens palsy, and light-near dissociation of the pupils.

Nystagmus This is a rhythmic oscillation of the eyes, occurring physiologically from vestibular and optokinetic stimulation or pathologically in a wide variety of diseases (Chap. 28). Abnormalities of the eyes or optic nerves, present at birth or acquired in childhood, can produce a complex, searching nystagmus with irregular pendular (sinusoidal) and jerk features. Examples are albinism, Leber's congenital amaurosis, and bilateral cataract. This nystagmus is commonly referred to as *congenital sensory nystagmus*. This is a poor term because even in children with congenital lesions, the nystagmus does not appear until weeks after birth. *Congenital motor nystagmus*, which looks similar to congenital sensory nystagmus, develops in the absence of any abnormality of the sensory visual system. Visual acuity also is reduced in congenital motor nystagmus, probably by the nystagmus itself, but seldom below a level of 20/200.

JERK NYSTAGMUS This is characterized by a slow drift off the target, followed by a fast corrective saccade. By convention, the nystagmus is named after the quick phase. Jerk nystagmus can be downbeat, upbeat, horizontal (left or right), and torsional. The pattern of nystagmus may vary with gaze position. Some patients will be oblivious to their nystagmus. Others will complain of blurred vision or a subjective to-and-fro movement of the environment (oscillopsia) corresponding to the nystagmus. Fine nystagmus may be difficult to see on gross examination of the eyes. Observation of nystagmoid movements of the optic disc on ophthalmoscopy is a sensitive way to detect subtle nystagmus.

GAZE-EVOKED NYSTAGMUS This is the most common form of jerk nystagmus. When the eyes are held eccentrically in the orbits, they have a natural tendency to drift back to primary position. The subject compensates by making a corrective saccade to maintain the deviated eye position. Many normal patients have mild gaze-evoked nystagmus. Exaggerated gaze-evoked nystagmus can be induced by drugs (sedatives, anticonvulsants, alcohol); muscle paresis; myasthenia gravis; demyelinating disease; and cerebellopontine angle, brainstem, and cerebellar lesions.

VESTIBULAR NYSTAGMUS *Vestibular nystagmus* results from dysfunction of the labyrinth (Ménière's disease), vestibular nerve, or vestibular nucleus in the brainstem. Peripheral vestibular nystagmus often occurs in discrete attacks, with symptoms of nausea and vertigo. There may be associated tinnitus and hearing loss. Sudden shifts in head position may provoke or exacerbate symptoms.

DOWNBEAT NYSTAGMUS *Downbeat nystagmus* results from lesions near the craniocervical junction (Chiari malformation, basilar invagination). It also has been reported in brainstem or cerebellar stroke, lithium or anticonvulsant intoxication, alcoholism, and multiple sclerosis. *Upbeat nystagmus* is associated with damage to the pontine tegmentum from stroke, demyelination, or tumor.

Opsoclonus This rare, dramatic disorder of eye movements consists of bursts of consecutive saccades (saccadomania). When the saccades are confined to the horizontal plane, the term *ocular flutter* is preferred. It can result from viral encephalitis, trauma, or a paraneoplastic effect of neuroblastoma, breast carcinoma, and other malignancies. It has also been reported as a benign, transient phenomenon in otherwise healthy patients.

Examination of the living human retina provides a unique opportunity for the direct study of nervous, vascular, and connective tissues. Many systemic disorders have retinal manifestations that are valuable for screening, diagnosis, and management of these conditions. Furthermore, retinal involvement in systemic disorders, such as diabetes mellitus, is a major cause of morbidity. Early recognition by ophthalmoscopic screening is a key factor in effective treatment. Ophthalmoscopy has the potential to be one of the most “high-yield” elements of the physical examination. Effective ophthalmoscopy requires a basic understanding of ocular structures and ophthalmoscopic techniques and recognition of abnormal findings.

OVERVIEW OF OCULAR STRUCTURES

The eye consists of a shell (cornea and sclera), lens, iris diaphragm, ciliary body, choroid, and retina. The anterior chamber is the space between the cornea and the lens, and it is filled with aqueous humor. The space between the posterior aspect of the lens and the retina is filled by vitreous gel. The choroid and the retina cover the posterior two-thirds of the sclera internally. The cornea and the lens form the focusing system of the eye, while the retina functions as the photoreceptor system, translating light to neuronal signals that are in turn transmitted to the brain via the optic nerve and visual pathways. The choroid is a layer of highly vascularized tissue that nourishes the retina and is located between the sclera and the retina. The retinal pigment epithelium (RPE) layer is a monolayer of pigmented cells that are adherent to the overlying retinal photoreceptor cells. RPE plays a major role in retinal photoreceptor metabolism.

NORMAL FUNDUS

The important areas that are visible by ophthalmoscopy include the macula, optic disc, retinal blood vessels, and retinal periphery (Fig. 40e-1).

THE MACULA

The macula is the central part of the retina and is responsible for detailed vision (acuity) and perception of color. The macula is defined clinically as the area of the retina centered on the posterior pole of the fundus, measuring about 5 disc diameters (DD) (7–8 mm) and bordered by the optic disc nasally and the temporal vascular arcades superiorly and inferiorly. Temporally, the macula extends for about 2.5 DD from its center. The fovea, in the central part of the macula, corresponds to the site of sharpest visual acuity. It is approximately 1 DD in size and appears darker in color than the surrounding area. The center of the fovea, the foveola, has a depressed pit-like configuration measuring about 350 μm .

THE OPTIC DISC

The optic disc measures about 1.5 mm and is located about 4 mm (2.5 DD) nasal to the fovea. It contains the central retinal artery and vein as they branch, a central excavation (cup), and a peripheral neural rim. Normally, the cup-to-disc ratio is less than 0.6. The cup is located temporal to the entry of the disc vessels. The normal optic disc is yellow/pink in color. It has clear and well-defined margins and is in the same plane as the retina (Fig. 40e-2). Pathologic findings include pallor (atrophy), swelling, and enlarged cupping.

THE EQUATOR AND PERIPHERAL RETINA

The equator of the fundus is clinically defined as the area that includes the internal opening of the vortex veins. The peripheral retina extends from the equator anteriorly to the ora serrata.

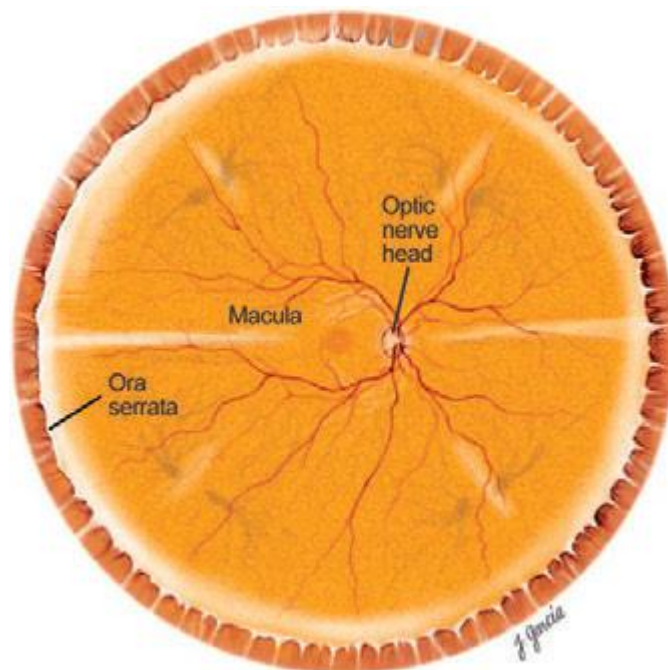


FIGURE 40e-1 Diagram showing the landmarks of the normal fundus. The macula is bounded by the superior and inferior vascular arcades and extends for 5 disc diameters (DD) temporal to the optic disc (optic nerve head). The central part of the macula (fovea) is located 2.5 DD temporal to the optic disc. The peripheral fundus is arbitrarily defined as the area extending anteriorly from the opening of the vortex veins to the ora serrata (the juncture between the retina and ciliary body). (Drawing courtesy of Juan R. Garcia. Used with permission from Johns Hopkins University.)



FIGURE 40e-2 Photograph of a normal left optic disc illustrating branching of the central retinal vein and artery, a physiologic cup, surface capillaries, and distinct margin. The cup is located temporal to the entry of the disc vessels. (From H Tabandeh, MF Goldberg: *Retina in Systemic Disease: A Color Manual of Ophthalmoscopy*. New York, Thieme, 2009.)

There are a number of ways to visualize the retina, including direct ophthalmoscopy, binocular indirect ophthalmoscopy, and slit-lamp biomicroscopy. Most nonophthalmologists prefer direct ophthalmoscopy, performed with a hand-held ophthalmoscope, because the technique is simple to master and the device is very portable. Ophthalmologists often use slit-lamp biomicroscopy and indirect ophthalmoscopy to obtain a more extensive view of the fundus.

DIRECT OPHTHALMOSCOPE

Direct ophthalmoscopes are simple hand-held devices that include a small light source for illumination, a viewing aperture through which the examiner looks at the retina, and a lens dial used for correction of the examiner's and the patient's refractive errors. A more recent design, the PanOptic ophthalmoscope, provides a wider field of view.

How to Use a Direct Ophthalmoscope Good alignment is the key. The goal is to align the examiner's eye with the viewing aperture of the ophthalmoscope, the patient's pupil, and the area of interest on the retina. Both the patient and the examiner should be in a comfortable position (sitting or lying for the patient, sitting or standing for the examiner). Dilating the pupil and dimming the room lights make the examination easier. Steps for performing direct ophthalmoscopy are summarized in [Table 40e-1](#).

PANOPTIC OPHTHALMOSCOPE

The PanOptic ophthalmoscope is a type of direct ophthalmoscope that is designed to provide a wider view of the fundus and has slightly more magnification than the standard direct ophthalmoscope. Steps for using the PanOptic Ophthalmoscope are summarized in [Table 40e-2](#).

TABLE 40e-1 GUIDELINES FOR PERFORMING DIRECT OPHTHALMOSCOPY

- Instruct the patient to remove glasses, keep the head straight, and to look steadily at a distant target straight in front. You may keep or remove your own glasses. Position your head at the same level as the patient's head.
- Use your right eye and right hand to examine the patient's right eye, and use your left eye and left hand to examine the patient's left eye.
- Using the ophthalmoscope light as a pen light, briefly examine the external features of the eye, including lashes, lid margins, conjunctiva, sclera, iris, and pupil shape, size, and reactivity.
- Shine the ophthalmoscope light into the patient's pupil at arm's length and observe the red reflex. Note abnormalities of the red reflex such as an opacity of the media.
- Dialing up a +10 D lens in the lens wheel, while examining the eye from 10 cm, allows magnified viewing of the anterior segment of the eye.
- Reduce the power of the lens in the wheel to zero, and move closer to the patient. Identify the optic disc by pointing the ophthalmoscope about 15° nasally or by following a blood vessel toward the apex of any branching. If the retina is out of focus, turn the lens dial either way, without moving your head. If the disc becomes clearer, keep turning until best focus is achieved; if it becomes more blurred, turn the dial the other way.
- Once you visualize the optic nerve, note its shape, size, color, margins, and the cup. Also note the presence of any venous pulsation or surrounding pigment, such as a choroidal or scleral crescent.
- Next, examine the macula. The macula is the area between the superior and inferior temporal vascular arcades, and its center is the fovea. You can examine the macula by pointing your ophthalmoscope about 15° temporal to the optic disc. Alternatively, ask the patient to look into the center of the light. Note the foveal reflex and the presence of any hemorrhage, exudate, abnormal blood vessels, scars, deposits, or other abnormalities.
- Examine the retinal blood vessels by re-identifying the optic disc and following each of the four main branches away from the disc. The veins are dark red and relatively large. The arteries are narrower and bright red.
- Ask the patient to look in the eight cardinal directions to allow you to view the peripheral fundus. In a patient with a well-dilated pupil, it is possible to visualize as far as the equator.

TABLE 40e-2 HOW TO USE A PANOPTIC OPHTHALMOSCOPE

- Focus the ophthalmoscope: Look through the scope at an object that is at least 10 to 15 feet away. Sharpen the image of the object by using the focusing wheel. Set the aperture dial to "small" or home position.
- Turn the scope on, and adjust the light intensity to "Maximum."
- Instruct the patient to look straight ahead. Move the ophthalmoscope close to the patient until the eyecup touches the patient's brow. The eyecup should be compressed about half its length to optimize the view.
- Visualize the optic disc.
- Examine the fundus as described in [Table 40e-1](#).

RETINAL SIGNS ASSOCIATED WITH SYSTEMIC DISEASES

AGE-RELATED CHANGES

Common age-related changes include diminished foveal light reflex, drusen (small yellow subretinal deposits), mild RPE atrophy, and pigment clumping.

RETINAL HEMORRHAGES

Retinal hemorrhages may take various shapes and sizes depending on their location within the retina ([Figs. 40e-3 and 40e-4](#)). Flame-shaped hemorrhages are located at the level of the superficial nerve fiber layer and represent bleeding from the inner capillary network of the retina. A white-centered hemorrhage is a superficial flame-shaped hemorrhage with an area of central whitening, often representing edema, focal necrosis, or cellular infiltration. Causes of white-centered hemorrhage include bacterial endocarditis and septicemia (Roth spots), lymphoproliferative disorders, diabetes mellitus, hypertension, anemia, and collagen vascular disorders. Dot hemorrhages are small, round, superficial hemorrhages that also originate from the superficial capillary network of the retina. They resemble microaneurysms. Blot hemorrhages are slightly larger in size, dark, and intraretinal. They represent bleeding from the deep capillary network of the retina. Subhyaloid hemorrhages are variable in shape and size and tend to be larger than other types of hemorrhages. They often have a fluid level ("boat-shaped" hemorrhage) and are located within the space between the vitreous and the retina. Subretinal hemorrhages are located deep (external) to the retina. The retinal vessels can be seen crossing over (internal to) such hemorrhages. Subretinal hemorrhages are variable in size and most commonly are caused by choroidal neovascularization (e.g., wet macular degeneration).



FIGURE 40e-3 Superficial flame-shaped hemorrhages, dot hemorrhages, and microaneurysms in a patient with nonproliferative diabetic retinopathy.

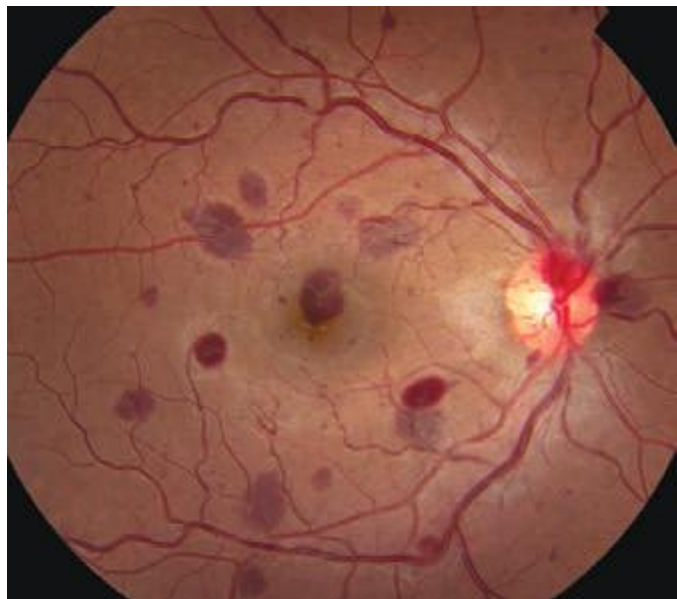


FIGURE 40e-4 Deep and superficial retinal hemorrhages in a patient with chronic leukemia.

Conditions associated with retinal hemorrhages include diseases causing retinal microvasculopathy (Table 40e-3), retinitis, retinal macroaneurysm, papilledema, subarachnoid hemorrhage (Terson's syndrome), Valsalva retinopathy, trauma (ocular injury, head injury, compression injuries of chest and abdomen, shaken baby syndrome, strangulation), macular degeneration, and posterior vitreous detachment. Hyperviscosity states may produce dot and blot hemorrhages, dilated veins ("string of sausages" appearance), optic disc edema, and exudates; similar changes can occur with adaptation to high altitude in mountain climbers.

MICROANEURYSMS

Microaneurysms are outpouchings of the retinal capillaries, appearing as red dots (similar to dot hemorrhages) and measuring 15–50 μm . Microaneurysms have increased permeability and may bleed or leak, resulting in localized retinal hemorrhage or edema. A microaneurysm ultimately thromboses and disappears within 3–6 months. Microaneurysms may occur in any condition that causes retinal microvasculopathy (Table 40e-3).

TABLE 40e-3 DISEASES ASSOCIATED WITH RETINAL MICROVASCULOPATHY

- Diabetes mellitus
- Systemic hypertension
- Retinal vein occlusion
- Retinal artery occlusion
- Multiple microemboli, e.g., talc retinopathy secondary to intravenous drug abuse, septicemia, endocarditis, Purtscher's retinopathy
- Carotid artery disease, carotid-cavernous fistula, aortic arch syndrome
- Sickle cell retinopathy
- Radiation retinopathy, head/neck irradiation
- HIV retinopathy
- Retinal vasculitis
- Anemia
- Thrombocytopenia
- Lymphoproliferative disorders
- Coagulopathy
- Hyperviscosity syndromes
- Retinopathy of prematurity

HARD EXUDATES

Hard exudates are well-circumscribed, shiny, yellow deposits located within the retina. They arise at the margins of areas of retinal edema and indicate increased capillary permeability. Hard exudates contain lipoproteins and lipid-laden macrophages. They may clear spontaneously or following laser photocoagulation, often within 6 months. Hard exudates may occur in isolation or may be scattered throughout the fundus. They may occur in a circular (circinate) pattern centered around an area of leaking microaneurysms. A macular star consists of a radiating, star-shaped pattern of hard exudates that is characteristically seen in severe systemic hypertension and in neuroretinitis associated with cat-scratch disease. Conditions associated with hard exudates include those causing retinal microvasculopathy (Table 40e-3), papilledema, neuroretinitis such as cat-scratch disease and Lyme disease, retinal vascular lesions (macroaneurysm, retinal capillary hemangioma, Coats' disease), intraocular tumors, and wet age-related macular degeneration. Drusen may be mistaken for hard exudates on ophthalmoscopy. Unlike hard exudates, drusen are nonrefractile subretinal deposits with blurred margins. They are usually seen in association with age-related macular degeneration.

COTTON-WOOL SPOTS

Cotton-wool spots are yellow/white superficial retinal lesions with indistinct feathery borders measuring 0.25–1 DD in size (Fig. 40e-5). They represent areas of edema within the retinal nerve fiber layer due to focal ischemia. Cotton-wool spots usually resolve spontaneously within 3 months. If the underlying ischemic condition persists, new lesions can develop in different locations. Cotton-wool spots often occur in conjunction with retinal hemorrhages and microaneurysms and represent retinal microvasculopathy caused by a number of systemic conditions (Table 40e-3). They may occur in isolation in HIV retinopathy, systemic lupus erythematosus, anemia, bodily trauma, other systemic conditions (Purtscher's/Purtscher's-like retinopathy), and interferon therapy.

RETINAL NEOVASCULARIZATION

Retinal neovascular complexes are irregular meshworks of fine blood vessels that grow in response to severe retinal ischemia or chronic inflammation (Fig. 40e-6). They may occur on or adjacent to the optic disc or elsewhere in the retina. Neovascular complexes are very

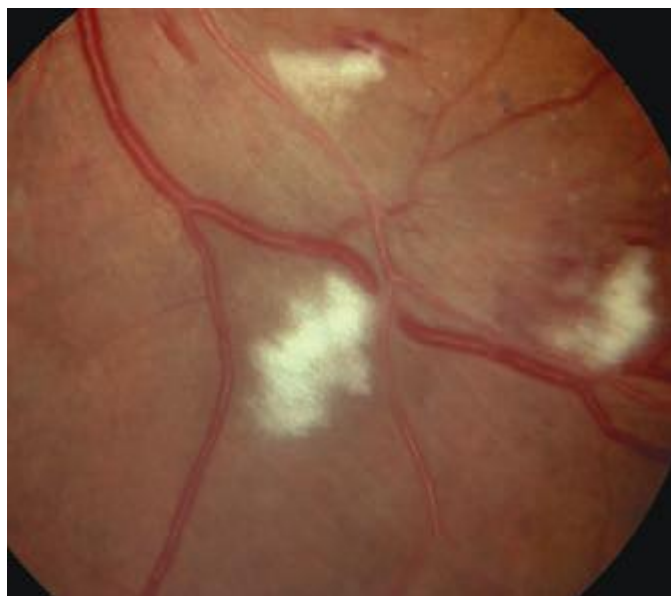


FIGURE 40e-5 Cotton-wool spots, yellow-white superficial lesions with characteristic feathery borders, in a patient with hypertensive retinopathy. (From H Tabandeh, MF Goldberg: *Retina in Systemic Disease: A Color Manual of Ophthalmoscopy*. New York, Thieme, 2009.)

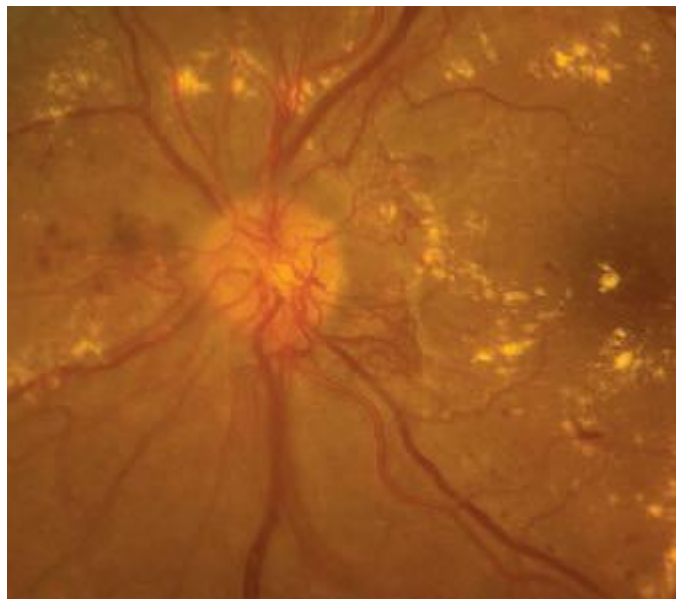


FIGURE 40e-6 Optic disc neovascularization in a patient with severe proliferative diabetic retinopathy. Multiple hard exudates are also present.

fragile and have a high risk for hemorrhaging, often causing visual loss. Diseases associated with retinal neovascularization include conditions that cause severe retinal microvasculopathy, especially diabetic and sickle cell retinopathies (Table 40e-3), intraocular tumors, intraocular inflammation (sarcoidosis, chronic uveitis), and chronic retinal detachment.

RETINAL EMBOLI

Common sources of retinal emboli include carotid artery atheromatous plaque, cardiac valve and septal abnormalities, cardiac arrhythmias, atrial myxoma, bacterial endocarditis, septicemia, fungemia, and intravenous drug abuse.

Platelet emboli are yellowish in appearance and conform to the shape of the blood vessel. They usually originate from an atheromatous plaque within the carotid artery and can cause transient loss of vision (amaurosis fugax). Cholesterol emboli, otherwise termed Hollenhorst plaques, are yellow crystalline deposits that are commonly found at the bifurcations of the retinal arteries and may be associated with amaurosis fugax. Calcific emboli have a pearly white appearance, are larger than the platelet and cholesterol emboli, and tend to lodge in the larger retinal arteries in or around the optic disc. Calcific emboli often result in retinal arteriolar occlusion. Septic emboli can cause white-centered retinal hemorrhages (Roth spots), retinal microabscesses, and endogenous endophthalmitis. Fat embolism and amniotic fluid embolism are characterized by multiple small vessel occlusions, typically causing cotton-wool spots and few hemorrhages (Purtscher's-like retinopathy). Talc embolism occurs with intravenous drug abuse and is characterized by multiple refractile deposits within the small retinal vessels. Any severe form of retinal artery embolism may result in retinal ischemia and its sequelae, including retinal neovascularization.

CHERRY RED SPOT AT THE MACULA

Cherry red spot at the macula is the term used to describe the dark red appearance of the central foveal area in comparison to the surrounding macular region (Fig. 40e-7). This appearance is most commonly due to a relative loss of transparency of the parafoveal retina resulting from ischemic cloudy swelling or storage of macromolecules within the ganglion cell layer. Diseases associated with a cherry red spot at the macula include central retinal artery occlusion, sphingolipidoses, and mucopolipidoses.

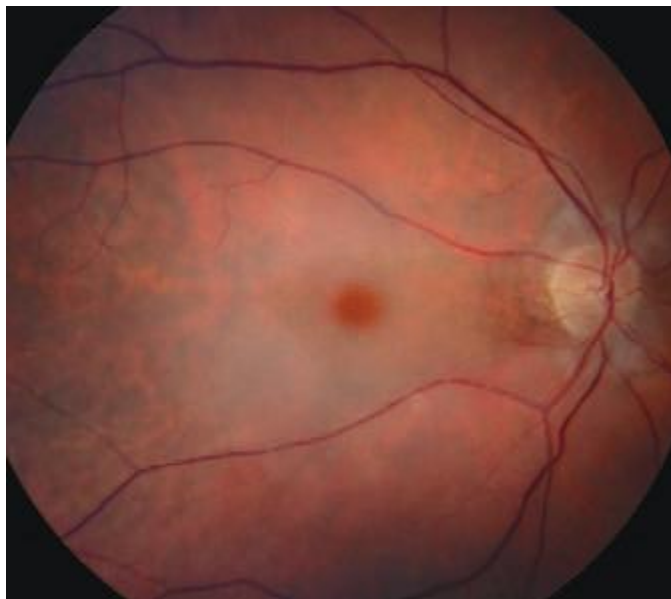


FIGURE 40e-7 Cherry red spot at the macula and cloudy swelling of the macula in a patient with central retinal artery occlusion due to embolus originating from a carotid artery atheromatous plaque.

RETINAL CRYSTAL DEPOSITION

Retinal crystals appear as fine, refractile, yellow-white deposits. Associated conditions include infantile cystinosis, primary hyperoxaluria, secondary oxalosis, Sjögren-Larson syndrome, intravenous drug abuse (talc retinopathy), and drugs such as tamoxifen, canthaxanthin, nitrofurantoin, methoxyflurane, and ethylene glycol. Crystals may also be seen in primary retinal diseases such as juxtafoveal telangiectasia, gyrate atrophy, and Bietti's crystalline degeneration. Old microemboli may mimic retinal crystals.

RETINAL VASCULAR SHEATHING

Vascular sheathing appears as a yellow-white cuff surrounding a retinal artery or vein (Fig. 40e-8). Diseases associated with retinal vascular sheathing include sarcoidosis, tuberculosis, toxoplasmosis, syphilis, HIV, retinitis (cytomegalovirus, herpes zoster, and herpes simplex),



FIGURE 40e-8 Vascular sheathing over the optic disc in a patient with neurosarcoidosis.

Lyme disease, cat-scratch disease, multiple sclerosis, chronic leukemia, amyloidosis, Behçet's disease, retinal vasculitis, retinal vascular occlusion, and chronic uveitis.

RETINAL DETACHMENT

Retinal detachment is the separation of the retina from the underlying RPE. There are three main types: (1) serous/exudative, (2) tractional, and (3) rhegmatogenous retinal detachment.

In serous retinal detachment, the location of the subretinal fluid is position-dependent, characteristically gravitating to the lowermost part of the fundus (shifting fluid sign), and retinal breaks are absent. Diseases associated with serous/exudative retinal detachment include severe systemic hypertension, dural arteriovenous shunt, retinal vascular anomalies, hyperviscosity syndromes, papilledema, posterior uveitis, scleritis, orbital inflammation, and intraocular neoplasms such as choroidal melanoma, choroidal metastasis, lymphoma, and multiple myeloma.

Tractional retinal detachment is caused by internal traction on the retina in the absence of a retinal break. The retina in the area of detachment is immobile and concaved internally. Fibrovascular proliferation is a frequent associated finding. Conditions associated with tractional retinal detachment include vascular proliferative retinopathies such as severe proliferative diabetic retinopathy, branch retinal vein occlusion, sickle cell retinopathy, and retinopathy of prematurity. Ocular trauma, proliferative vitreoretinopathy, and intraocular inflammation are other causes of a tractional retinal detachment.

Rhegmatogenous retinal detachment is caused by the presence of a retinal break, allowing fluid from the vitreous cavity to gain access to the subretinal space. The surface of the retina is usually convex forward. Rhegmatogenous retinal detachment has a corrugated appearance, and undulates with eye movement. Causes of retinal breaks include posterior vitreous detachment, severe vitreoretinal traction, trauma, intraocular surgery, retinitis, and atrophic holes.

OPTIC DISC SWELLING

Optic disc swelling is abnormal elevation of the optic disc with blurring of its margins (Fig. 40e-9). The term “papilledema” is used to describe swelling of the optic disc secondary to elevation of intracranial pressure. In papilledema, the normal venous pulsation at the disc is characteristically absent. The differential diagnosis of optic disc swelling includes papilledema, anterior optic neuritis (papillitis), central retinal vein occlusion, anterior ischemic optic neuropathy,

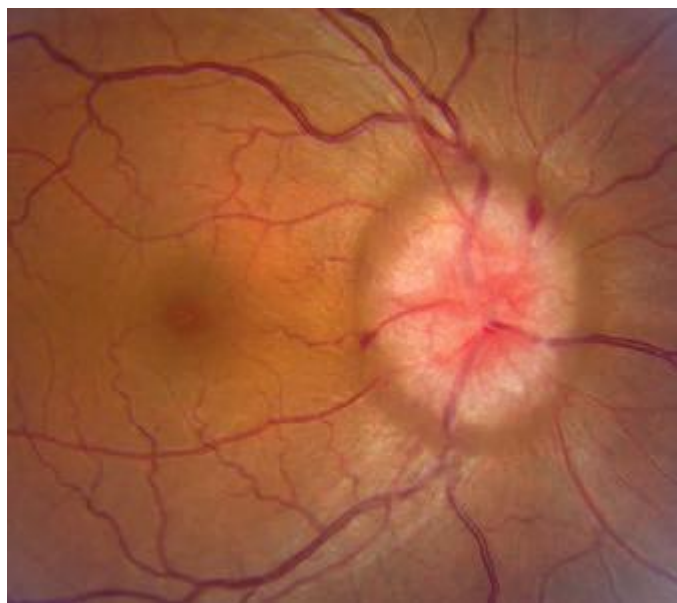


FIGURE 40e-9 Optic disc swelling in a patient with papilledema due to idiopathic intracranial hypertension. The optic disc is hyperemic, with indistinct margins. Superficial hemorrhages are present.



FIGURE 40e-10 Optic disc edema and retinal hemorrhages in a patient with malignant hypertension.

toxic optic neuropathy, hereditary optic neuropathy, neuroretinitis, diabetic papillopathy, hypertension (Fig. 40e-10), respiratory failure, carotid-cavernous fistula, optic disc nerve infiltration (glioma, lymphoma, leukemia, sarcoidosis, and granulomatous infections), ocular hypotony, chronic intraocular inflammation, optic disc drusen (pseudopapilledema), and high hypermetropia (pseudopapilledema).

CHORIORETINAL MASS LESIONS

Choroidal mass lesions appear thickened and may or may not be associated with increased pigmentation. Pigmented mass lesions include choroidal nevus (usually flat), choroidal malignant melanoma (Fig. 40e-11), and melanocytoma. Nonpigmented lesions include amelanotic choroidal melanoma, choroidal metastasis, retinoblastoma, capillary hemangioma, granuloma (e.g., *Toxocara canis*), choroidal detachment, choroidal hemorrhage, and wet age-related macular degeneration. Other rare tumors that may be visible on ophthalmoscopy include

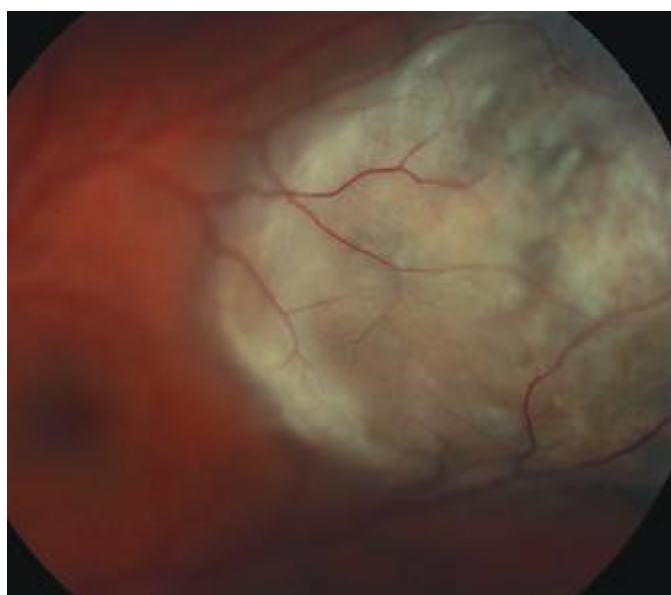


FIGURE 40e-11 Choroidal malignant melanoma. The lesion is highly elevated and pigmented, and has subretinal orange pigment deposits characteristic for malignant melanoma.

40e-6 osteoma, astrocytoma (e.g., tuberous sclerosis), neurilemmoma, and leiomyoma.

PIGMENTED LESIONS

The differential diagnosis of flat pigmented lesions of the fundus is summarized in [Table 40e-4](#). The appearance of chorioretinal scarring from old *Toxoplasma* chorioretinitis is shown in [Fig. 40e-12](#).

TABLE 40e-4 DIFFERENTIAL DIAGNOSIS OF FLAT PIGMENTED LESIONS OF THE FUNDUS

Bone spicule pigmentation

- Retinitis pigmentosa and its variants
- Pigmentary retinopathy in systemic diseases: Usher's syndrome, abetalipoproteinemia, Refsum's disease, Kearns-Sayre syndrome, Alström's syndrome, Cockayne's syndrome, Friedreich's ataxia, mucopolysaccharidoses, paraneoplastic syndrome
- Infections: congenital rubella (salt and pepper retinopathy), congenital syphilis
- Resolved choroidal/retinal detachment
- Age-related reticular pigmentary degeneration

Patchy pigmented lesions

- Chorioretinal scars
 - Infections: *Toxoplasma gondii*, *Toxocara canis*, syphilis, cytomegalovirus, herpes zoster and herpes simplex viruses, west Nile virus, histoplasmosis, parasitic infection
 - Choroiditis: sarcoidosis, sympathetic ophthalmia, Vogt-Koyanagi-Harada syndrome
 - Choroidal infarct: severe hypertension, sickle cell hemoglobinopathies
 - Trauma, cryotherapy, laser photocoagulation scars
 - Age-related macular degeneration
- Drugs: chloroquine/hydroxychloroquine, thioridazine, chlorpromazine, desferrioxamine
- Choroidal nevus
- Congenital hypertrophy of the retinal pigment epithelium

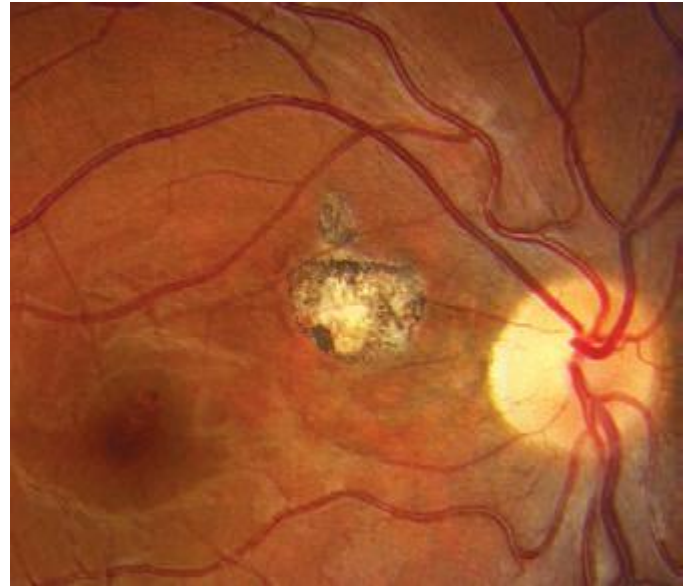


FIGURE 40e-12 Chorioretinal scarring due to old *Toxoplasma* chorioretinitis. The lesion is flat and pigmented. Areas of hypopigmentation are also present.

41e Video Library of Neuro-Ophthalmology

Shirley H. Wray

The proper control of eye movements requires the coordinated activity of many different anatomic structures in the peripheral and central nervous system, and in turn, manifestations of a diverse array of neurologic and medical disorders are revealed as disorders of eye movement. In this remarkable video collection, an introduction to distinctive eye movement disorders encountered in the context of neuromuscular, paraneoplastic, demyelinating, neurovascular, and neurodegenerative disorders is presented.

Cases with Multiple Sclerosis

- VIDEO 41e-1** Fisher's One-and-a-Half Syndrome (ID164-2)
VIDEO 41e-2 A Case of Ocular Flutter (ID166-2)
VIDEO 41e-3 Downbeat Nystagmus and Periodic Alternating Nystagmus (ID168-6)
VIDEO 41e-4 Bilateral Internuclear Ophthalmoplegia (ID933-1)

Cases with Myasthenia Gravis or Mitochondrial Myopathy

- VIDEO 41e-5** Unilateral Ptosis: Myasthenia Gravis (Thymic Tumor) (ID163-1)
VIDEO 41e-6 Progressive External Ophthalmoplegia (Progressive External Ophthalmoplegia: Mitochondrial Cytopathy) (ID906-2)

Cases with Paraneoplastic Disease

- VIDEO 41e-7** Paraneoplastic Upbeat Nystagmus, Cancer of the Pancreas, Positive Anti-Hu Antibody (ID212-3)
VIDEO 41e-8 Paraneoplastic Ocular Flutter, Small-Cell Adenocarcinoma of the Lung, Negative Marker (ID936-7)
VIDEO 41e-9 Opsoclonus/Flutter, Bilateral Sixth Nerve Palsy, Adenocarcinoma of the Breast, Negative Marker (ID939-8)

Cases with Fisher's Syndrome

- VIDEO 41e-10** Bilateral Ptosis: Facial Diplegia, Total External Ophthalmoplegia, Positive Anti-GQ_{1b} Antibody (ID944-1)

Cases with Vascular Disease

- VIDEO 41e-11** Retinal Emboli (Film or Fundus) (ID16-1)
VIDEO 41e-12 Third Nerve Palsy (Microinfarct) (ID939-2)

Case with Neurodegenerative Disease

- VIDEO 41e-13** Apraxia of Eyelid Opening (Progressive Supranuclear Palsy) (ID932-3)

Case of Thyroid-Associated Ophthalmopathy

- VIDEO 41e-14** Restrictive Orbitopathy of Graves' Disease, Bilateral Exophthalmos (ID925-4)

Case with Wernicke's Encephalopathy

- VIDEO 41e-15** Bilateral Sixth Nerve Palsies (ID 163-3)

Case with the Locked-in-Syndrome

- VIDEO 41e-16** Ocular Dipping (ID 4-1)

Case in a Comatose Patient

- VIDEO 41e-17** Downbeat Nystagmus (ID 166-11)

Case Two Years After a Pontine Hemorrhage

- VIDEO 41e-18** Palatal Tremor (ID 936-4)

The Video Library of Neuro-Ophthalmology shows a number of cases with eye movement disorders. All the clips are taken from Dr. Shirley Wray's collection on the NOVEL website.

To access go to:

<http://NOVEL.utah.edu/Wray>

<http://Respiratory.Countway.Harvard.edu/Wray>

and/or to her book

Eye Movement Disorders in Clinical Practice

Shirley H. Wray, MD, PhD, Oxford University Press, 2014.

42 Disorders of Smell and Taste

Richard L. Doty, Steven M. Bromley

All environmental chemicals necessary for life enter the body by the nose and mouth. The senses of smell (olfaction) and taste (gustation) monitor such chemicals, determine the flavor and palatability of foods and beverages, and warn of dangerous environmental conditions, including fire, air pollution, leaking natural gas, and bacteria-laden foodstuffs. These senses contribute significantly to quality of life and, when dysfunctional, can have untoward physical and psychological consequences. A basic understanding of these senses in health and disease is critical for the physician, because thousands of patients present to doctors' offices each year with complaints of chemosensory dysfunction. Among the more important recent developments in neurology is the discovery that decreased smell function is among the first signs, if not the first sign, of such neurodegenerative diseases as Parkinson's disease (PD) and Alzheimer's disease (AD), signifying their "presymptomatic" phase.

ANATOMY AND PHYSIOLOGY

Olfactory System Odorous chemicals enter the front of nose during inhalation and active sniffing, as well as the back of the nose (nasopharynx) during deglutition. After reaching the highest recesses of the nasal cavity, they dissolve in the olfactory mucus and diffuse or are actively transported by specialized proteins to receptors located on the cilia of olfactory receptor cells. The cilia, dendrites, cell bodies, and proximal axonal segments of these bipolar cells are located within a unique neuroepithelium covering the cribriform plate, the superior nasal septum, superior turbinate, and sectors of the middle turbinate (Fig. 42-1). Each of the ~6 million bipolar receptor cells expresses only one of ~450 receptor protein types, most of which respond to more than a single chemical. When damaged, the receptor cells can be replaced by stem cells near the basement membrane. Unfortunately, such replacement is often incomplete.

After coalescing into bundles surrounded by glia-like ensheathing cells (termed fila), the receptor cell axons pass through the cribriform plate to the olfactory bulbs, where they synapse with dendrites of other cell types within the glomeruli (Fig. 42-2). These spherical structures, which make up a distinct layer of the olfactory bulb, are a site of convergence of information, because many more fibers enter than leave them. Receptor cells that express the same type of receptor project to the same glomeruli, effectively making each glomerulus a functional unit. The major projection neurons of the olfactory system—the mitral and tufted cells—send primary dendrites into the glomeruli, connecting not only with the incoming receptor cell axons, but with dendrites of periglomerular cells. The activity of the mitral/tufted cells is modulated by the periglomerular cells, secondary dendrites from other mitral/tufted cells, and granule cells, the most numerous cells of the bulb. The latter cells, which are largely GABAergic, receive inputs from central brain structures and modulate the output of the mitral/tufted cells. Interestingly, like the olfactory receptor cells, some cells within the bulb undergo replacement. Thus, neuroblasts formed within the anterior subventricular zone of the brain migrate along the rostral migratory stream, ultimately becoming granule and periglomerular cells.

The axons of the mitral and tufted cells synapse within the primary olfactory cortex (POC) (Fig. 42-3). The POC is defined as those cortical structures that receive direct projections from the olfactory bulb, most notably the piriform and entorhinal cortices. Although olfaction is unique in that its initial afferent projections bypass the thalamus, persons with damage to the thalamus can exhibit olfactory deficits, particularly ones of odor identification. Such deficits likely reflect the involvement of thalamic connections between the primary olfactory cortex and the orbitofrontal cortex (OFC), where odor identification occurs. The close anatomic ties between the olfactory system and the

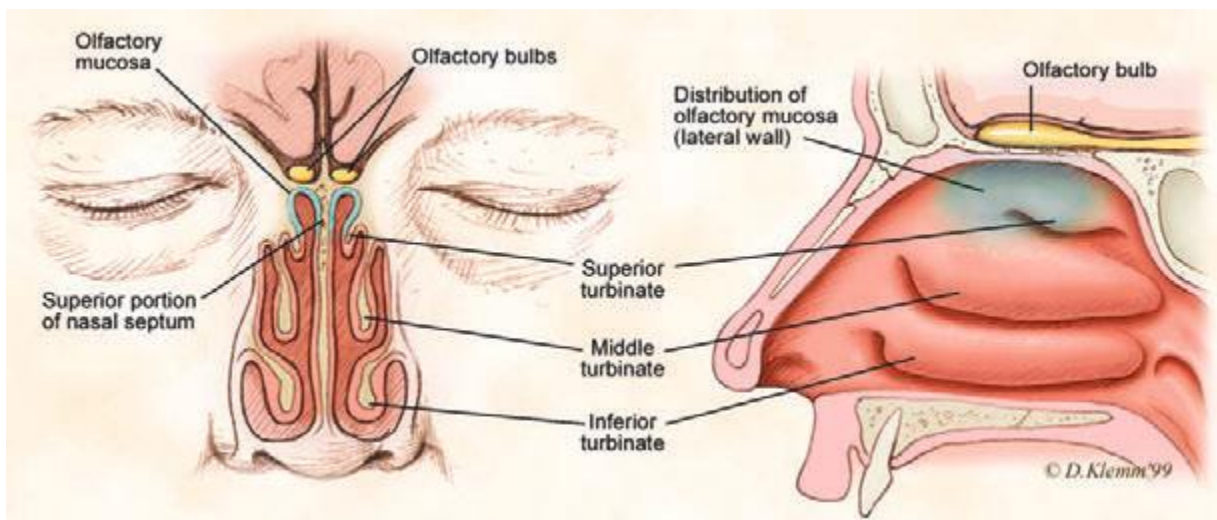


FIGURE 42-1 Anatomy of the olfactory neural pathways, showing the distribution of olfactory receptors in the roof of the nasal cavity. (Copyright David Klemm, Faculty and Curriculum Support [FACS], Georgetown University Medical Center; used with permission.)

amygdala, hippocampus, and hypothalamus help to explain the intimate associations between odor perception and cognitive functions such as memory, motivation, arousal, autonomic activity, digestion, and sex.

Taste System Tastants are sensed by specialized receptor cells present within taste buds—small grapefruit-like segmented structures located on the lateral margins and dorsum of the tongue, roof of the mouth, pharynx, larynx, and superior esophagus (Fig. 42-4). Lingual taste buds are imbedded in well-defined protuberances, termed fungiform, foliate, and circumvallate papillae. After dissolving in a liquid, tastants enter the opening of the taste bud—the taste pore—and bind to receptors on microvilli, small extensions of receptor cells within each taste bud. Such binding changes the electrical potential across the taste cell, resulting in neurotransmitter release onto the first-order taste neurons. Although humans have ~7500 taste buds, not all harbor

taste-sensitive cells; some contain only one class of receptor (e.g., cells responsive only to sugars), whereas others contain cells sensitive to more than one class. The number of taste receptor cells per taste bud ranges from zero to well over 100. A small family of three G-protein-coupled receptors (GPCRs), namely T1R1, T1R2, and T1R3, mediate sweet and umami taste sensations. Bitter sensations, on the other hand, depend on T2R receptors, a family of ~30 GPCRs expressed on cells different from those that express the sweet and umami receptors. T2Rs sense a wide range of bitter substances but do not distinguish among them. Sour tastants are sensed by the PKD2L1 receptor, a member of the transient receptor potential protein (TRP) family. Perception of salty sensations, such as induced by sodium chloride, arises from the entry of Na^+ ions into the cells via specialized membrane channels, such as the amiloride-sensitive Na^+ channel.

Recent studies have found that both bitter and sweet taste-related receptors are also present elsewhere in the body, most notably in the

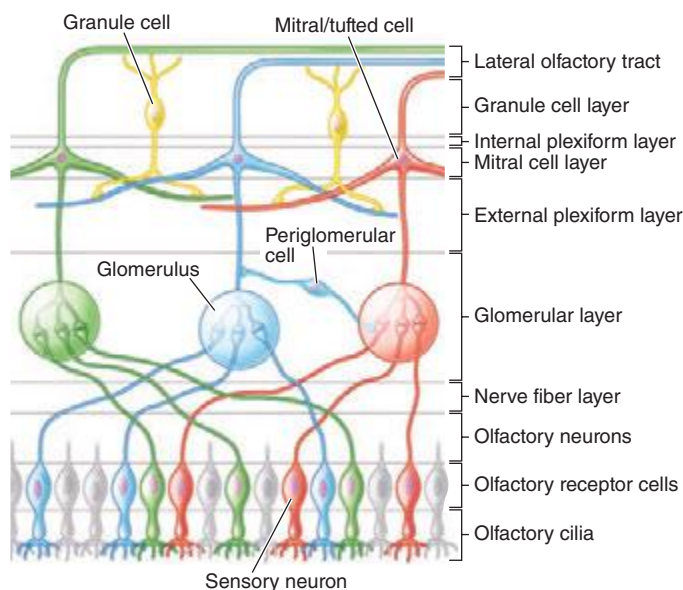


FIGURE 42-2 Schematic of the layers and wiring of the olfactory bulb. Each receptor type (red, green, blue) projects to a common glomerulus. The neural activity within each glomerulus is modulated by periglomerular cells. The activity of the primary projection cells, the mitral and tufted cells, is modulated by granule cells, periglomerular cells, and secondary dendrites from adjacent mitral and tufted cells. (From www.med.yale.edu/neurosurg/treloar/index.html.)

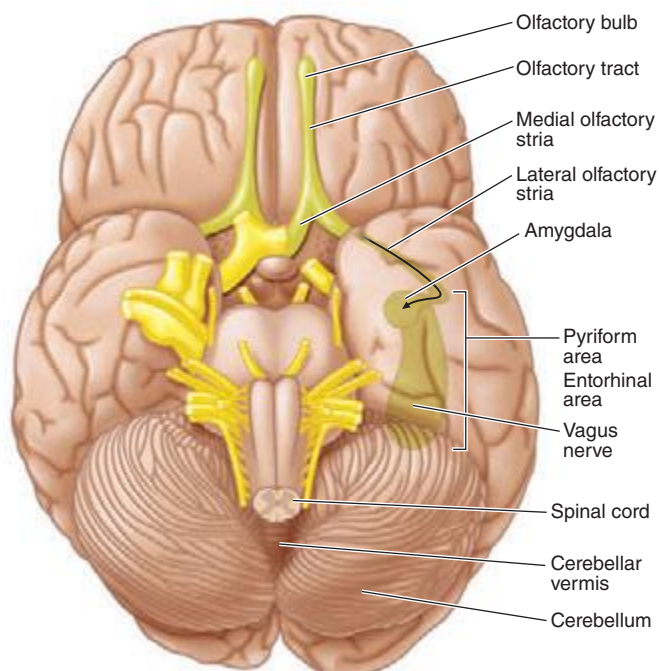


FIGURE 42-3 Anatomy of the base of the brain showing the primary olfactory cortex.

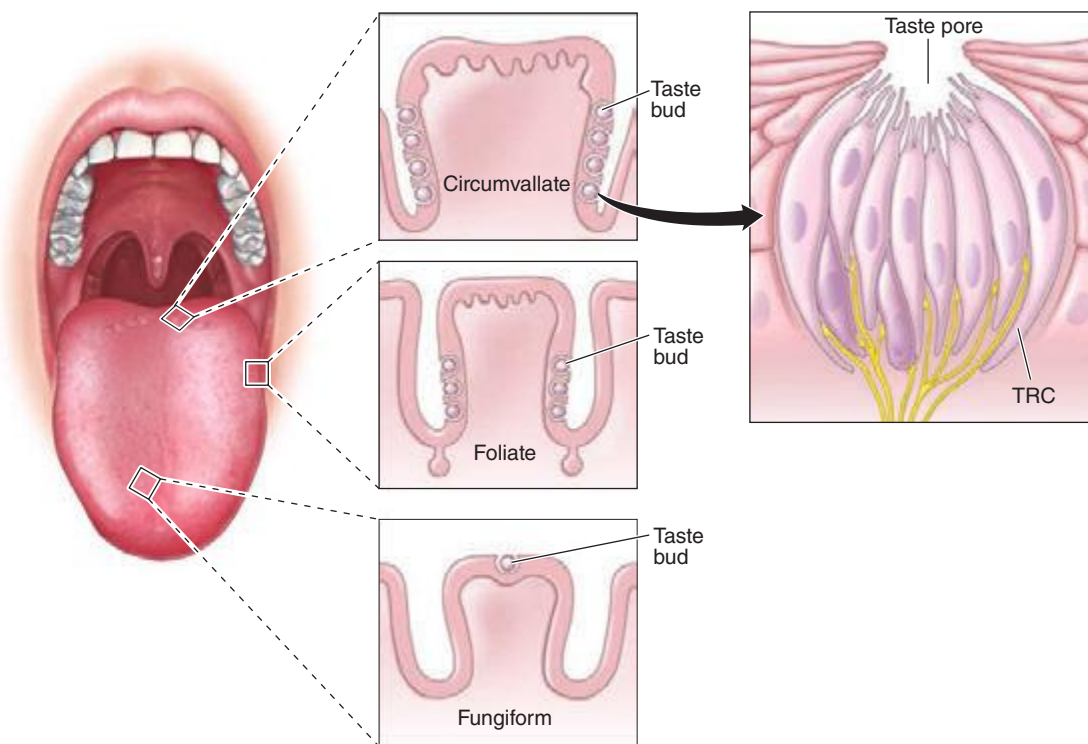


FIGURE 42-4 Schematic of the taste bud and its opening (pore), as well as the location of buds on the three major types of papillae: fungiform (anterior), foliate (lateral), and circumvallate (posterior).

alimentary and respiratory tracts. This important discovery generalizes the concept of taste-related chemoreception to areas of the body beyond the mouth and throat, with α -gustducin, the taste-specific G-protein α -subunit, expressed in so-called brush cells found specifically within the human trachea, lung, pancreas, and gallbladder. These brush cells are rich in nitric oxide (NO) synthase, known to defend against xenobiotic organisms, protect the mucosa from acid-induced lesions, and, in the case of the gastrointestinal tract, stimulate vagal and splanchnic afferent neurons. NO further acts on nearby cells, including enteroendocrine cells, absorptive or secretory epithelial cells, mucosal blood vessels, and cells of the immune system. Members of the T2R family of bitter receptors and the sweet receptors of the T1R family have been identified within the gastrointestinal tract and in enteroendocrine cell lines. In some cases, these receptors are important for metabolism, with the T1R3 receptors and gustducin playing decisive roles in the sensing and transport of dietary sugars from the intestinal lumen into absorptive enterocytes via a sodium-dependent glucose transporter and in regulation of hormone release from gut enteroendocrine cells. In other cases, these receptors may be important for airway protection, with a number of T2R bitter receptors in the motile cilia of the human airway that responded to bitter compounds by increasing their beat frequency. One specific T2R38 taste receptor is expressed in human upper respiratory epithelia and responds to acyl-monoserine lactone quorum-sensing molecules secreted by *Pseudomonas aeruginosa* and other gram-negative bacteria. Differences in T2R38 functionality, as related to TAS2R38 genotype, correlate with susceptibility to upper respiratory infections in humans.

Taste information is sent to the brain via three cranial nerves (CNs): CN VII (the *facial nerve*, which involves the intermediate nerve with its branches, the greater petrosal and chorda tympani nerves), CN IX (the *glossopharyngeal nerve*), and CN X (the *vagus nerve*) (Fig. 42-5). CN VII innervates the anterior tongue and all of the soft palate, CN IX innervates the posterior tongue, and CN X innervates the laryngeal surface of the epiglottis, larynx, and proximal portion of the esophagus. The mandibular branch of CN V (V_3) conveys somatosensory information (e.g., touch, burning, cooling, irritation) to the brain. Although not technically a gustatory nerve, CN V shares primary nerve routes with many of the gustatory nerve fibers and adds temperature, texture,

pungency, and spiciness to the taste experience. The chorda tympani nerve is famous for taking a recurrent course through the facial canal in the petrosal portion of the temporal bone, passing through the middle ear, and then exiting the skull via the petrotympanic fissure, where it joins the lingual nerve (a division of CN V) near the tongue. This nerve also carries parasympathetic fibers to the submandibular and sublingual glands, whereas the greater petrosal nerve supplies the palatine glands, thereby influencing saliva production.

The axons of the projection cells, which synapse with taste buds, enter the rostral portion of the nucleus of the solitary tract (NTS)

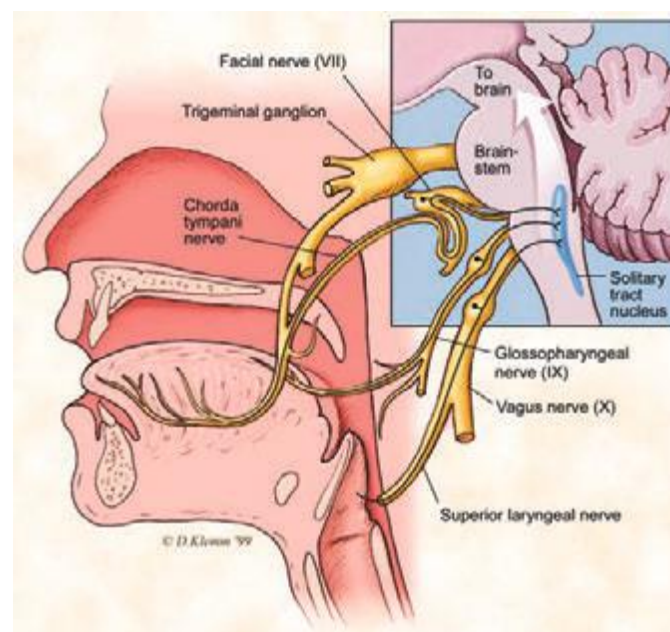


FIGURE 42-5 Schematic of the cranial nerves (CNs) that mediate taste function, including the chorda tympani nerve (CN VII), the glossopharyngeal nerve (CN IX), and the vagus nerve (CN X).

within the medulla of the brainstem (Fig. 42-5). From the NTS, neurons then project to a division of the ventroposteromedial thalamic nucleus (VPM) via the medial lemniscus. From here, projections are made to the rostral part of the frontal operculum and adjoining insula, a brain region considered the *primary taste cortex* (PTC). Projections from the PTC then go to the *secondary taste cortex*, namely the caudolateral OFC. This brain region is involved in the conscious recognition of taste qualities. Moreover, because it contains cells that are activated by several sensory modalities, it is likely a center for establishing “flavor.”

DISORDERS OF OLFACTION

The ability to smell is influenced, in everyday life, by such factors as age, gender, general health, nutrition, smoking, and reproductive state. Women typically outperform men on tests of olfactory function and retain normal smell function to a later age than do men. Significant decrements in the ability to smell are present in over 50% of the population between 65 and 80 years of age and in 75% of those 80 years of age and older (Fig. 42-6). Such presbyosmia helps to explain why many elderly report that food has little flavor, a problem that can result in nutritional disturbances. This also helps to explain why a disproportionate number of elderly die in accidental gas poisonings. A relatively complete listing of conditions and disorders that have been associated with olfactory dysfunction is presented in Table 42-1.

Aside from aging, the three most common identifiable causes of long-lasting or permanent smell loss seen in the clinic are, in order of frequency, severe upper respiratory infections, head trauma, and chronic rhinosinusitis. The physiologic basis for most head trauma-related losses is the shearing and subsequent scarring of the olfactory fila as they pass from the nasal cavity into the brain cavity. The cribriform plate does not have to be fractured or show pathology for smell loss to be present. Severity of trauma, as indexed by a poor Glasgow Coma Scale score on presentation and the length of posttraumatic amnesia, is associated with higher risk of olfactory impairment. Less than 10% of posttraumatic anosmic patients will recover age-related normal function over time. This increases to nearly 25% of those with less-than-total loss. Upper respiratory infections, such as those associated with the common cold, influenza, pneumonia, or HIV, can directly and permanently harm the olfactory epithelium by decreasing receptor cell number, damaging cilia on remaining receptor cells, and inducing the replacement of sensory epithelium with respiratory epithelium. The smell loss associated with chronic rhinosinusitis is

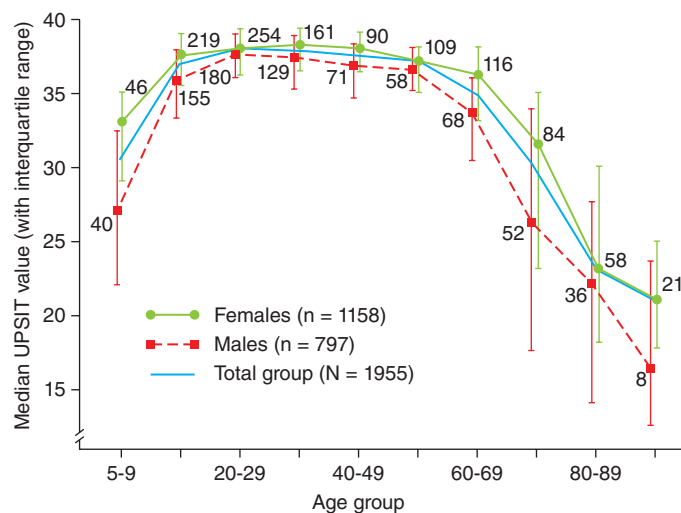


FIGURE 42-6 Scores on the University of Pennsylvania Smell Identification Test (UPSIT) as a function of subject age and sex. Numbers by each data point indicate sample sizes. Note that women identify odorants better than men at all ages. (From RL Doty et al: *Science* 226:1421, 1984. Copyright © 1984 American Association for the Advancement of Science.)

TABLE 42-1 DISORDERS AND CONDITIONS ASSOCIATED WITH COMPROMISED OLFACTION FUNCTION, AS MEASURED BY OLFACTION TESTING

22q11 deletion syndrome	Liver disease
AIDS/HIV infection	Lubag disease
Adenoid hypertrophy	Medications
Adrenal cortical insufficiency	Migraine
Age	Multiple sclerosis
Alcoholism	Multi-infarct dementia
Allergies	Myasthenia gravis
Alzheimer's disease	Narcolepsy with cataplexy
Amyotrophic lateral sclerosis (ALS)	Neoplasms, cranial/nasal
Anorexia nervosa	Nutritional deficiencies
Asperger's syndrome	Obstructive pulmonary disease
Ataxias	Obesity
Attention deficit/hyperactivity disorder	Obsessive compulsive disorder
Bardet-Biedl syndrome	Orthostatic tremor
Chagas' disease	Panic disorder
Chemical exposure	Parkinson's disease (PD)
Chronic obstructive pulmonary disease	Pick's disease
Congenital	Posttraumatic stress disorder
Cushing's syndrome	Pregnancy
Cystic fibrosis	Pseudohypoparathyroidism
Degenerative ataxias	Psychopathy
Diabetes	Radiation (therapeutic, cranial)
Down's syndrome	REM behavior disorder
Epilepsy	Refsum's disease
Facial paralysis	Renal failure/end-stage kidney disease
Frontotemporal lobe degeneration	Restless leg syndrome
Gonadal dysgenesis (Turner's syndrome)	Rhinosinusitis/polyposis
Guamanian ALS/PD/dementia syndrome	Schizophrenia
Head trauma	Seasonal affective disorder
Herpes simplex encephalitis	Sjögren's syndrome
Hypothyroidism	Stroke
Huntington's disease	Tobacco smoking
Itrogenesis	Toxic chemical exposure
Kallmann's syndrome	Upper respiratory infections
Korsakoff's psychosis	Usher syndrome
Leprosy	Vitamin B ₁₂ deficiency

related to disease severity, with most loss occurring in cases where rhinosinusitis and polyposis are both present. Although systemic glucocorticoid therapy can usually induce short-term functional improvement, it does not, on average, return smell test scores to normal, implying that chronic permanent neural loss is present and/or that short-term administration of systemic glucocorticoids does not completely mitigate the inflammation. It is well established that microinflammation in an otherwise seemingly normal epithelium can influence smell function.

A number of neurodegenerative diseases are accompanied by olfactory impairment, including PD, AD, Huntington's disease, Down's syndrome, parkinsonism-dementia complex of Guam, dementia with Lewy bodies (DLB), multiple system atrophy, corticobasal degeneration, and frontotemporal dementia; smell loss can also occur in multiple sclerosis (MS) and idiopathic rapid eye movement (REM) behavioral sleep disorder (iRBD). Olfactory impairment in PD often predates the clinical diagnosis by at least 4 years. In staged cases, studies of the sequence of formation of abnormal α -synuclein aggregates and Lewy bodies suggest that the olfactory bulbs may be, along with the dorsomotor nucleus of the vagus, the first site of neural damage

in PD. In postmortem studies of patients with very mild “presymptomatic” signs of AD, poorer smell function has been associated with higher levels of AD-related pathology. Smell loss is more marked in patients with early clinical manifestations of DLB than in those with mild AD. Interestingly, smell loss is minimal or nonexistent in progressive supranuclear palsy and 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)-induced parkinsonism. In MS, the olfactory disturbance varies as a function of the plaque activity within the frontal and temporal lobes.

The smell loss seen in iRBD is of the same magnitude as that found in PD. This is of particular interest because patients with iRBD frequently develop PD and hyposmia. There is some evidence that iRBD may actually represent an early associated condition of PD. REM behavior disorder is not only seen in its idiopathic form, but can also be associated with narcolepsy. This led to a recent study of narcoleptic patients with and without REM behavior disorder, which demonstrated that narcolepsy, independent of REM behavior disorder, was associated with impairments in olfactory function. Orexin A, also known as hypocretin-1, is dramatically diminished or undetectable in the cerebrospinal fluid of patients with narcolepsy and cataplexy (Chap. 38). The orexin-containing neurons in the hypothalamus project throughout the entire olfactory system (from the olfactory epithelium to the olfactory cortex), and damage to these orexin-containing projections may be one underlying mechanism for impaired olfactory performance in narcoleptic patients. The administration of intranasal orexin A (hypocretin-1) appears to result in improved olfactory function, supporting the notion that mild olfactory impairment is not only a primary feature of narcolepsy with cataplexy, but that central nervous system orexin deficiency may be a fundamental part of the mechanism for this loss.

DISORDERS OF TASTE

The majority of patients who present with taste dysfunction exhibit olfactory, not taste, loss. This is because most flavors attributed to taste actually depend on retronasal stimulation of the olfactory receptors during deglutition. As noted earlier, taste buds only mediate basic tastes such as sweet, sour, bitter, salty, and umami. Significant impairment of whole-mouth gustatory function is rare outside of generalized metabolic disturbances or systemic use of some medications, because taste bud regeneration occurs and peripheral damage alone would require the involvement of multiple cranial nerve pathways. Nonetheless, taste can be influenced by (1) the release of foul-tasting materials from the oral cavity from oral medical conditions or appliances (e.g., gingivitis, purulent sialadenitis), (2) transport problems of tastants to the taste buds (e.g., drying of the orolingual mucosa, infections, inflammatory conditions), (3) damage to the taste buds themselves (e.g., local trauma, invasive carcinomas), (4) damage to the neural pathways innervating the taste buds (e.g., middle ear infections), (5) damage to central structures (e.g., multiple sclerosis, tumor, epilepsy, stroke), and (6) systemic disturbances of metabolism (e.g., diabetes, thyroid disease, medications). Unlike CN VII, CN IX is relatively protected along its path, although iatrogenic interventions such as tonsillectomy, bronchoscopy, laryngoscopy, endotracheal intubation, and radiation therapy can result in selective injury. CN VII damage commonly results from mastoidectomy, tympanoplasty, and stapedectomy, in some cases inducing persistent metallic sensations. Bell’s palsy (Chap. 455) is one of the most common causes of CN VII injury that results in taste disturbance. On rare occasions, migraines (Chap. 447) are associated with a gustatory prodrome or aura, and in some cases, tastants can trigger a migraine attack. Interestingly, dysgeusia occurs in some cases of burning mouth syndrome (BMS; also termed *glossodynia* or *glossalgia*), as do dry mouth and thirst. BMS is likely associated with dysfunction of the trigeminal nerve (CN V). Some of the etiologies suggested for this poorly understood syndrome are amenable to treatment, including (1) nutritional deficiencies (e.g., iron, folic acid, B vitamins, zinc), (2) diabetes mellitus (possibly predisposing to oral candidiasis), (3) denture allergy, (4) mechanical irritation from dentures or oral devices, (5) repetitive movements of the mouth (e.g., tongue thrusting, teeth grinding, jaw clenching),

(6) tongue ischemia as a result of temporal arteritis, (7) periodontal disease, (8) reflux esophagitis, and (9) geographic tongue.

Although both taste and smell can be adversely influenced by pharmacologic agents, drug-related taste alterations are more common. Indeed, over 250 medications have been reported to alter the ability to taste. Major offenders include antineoplastic agents, antirheumatic drugs, antibiotics, and blood pressure medications. Terbinafine, a commonly used antifungal, has been linked to taste disturbance lasting up to 3 years. In a recent controlled trial, nearly two-thirds of individuals taking eszopiclone (Lunesta) experienced a bitter dysgeusia that was stronger in women, systematically related to the time since drug administration, and positively correlated with both blood and saliva levels of the drug. Intranasal use of nasal gels and sprays containing zinc, which are common over-the-counter prophylactics for upper respiratory viral infections, has been implicated in loss of smell function. Whether their efficacy in preventing such infections, which are the most common cause of anosmia and hyposmia, outweighs their potential detriment to smell function requires study. Dysgeusia occurs commonly in the context of drugs used to treat or minimize symptoms of cancer, with a weighted prevalence from 56–76% depending on the type of cancer treatment. Attempts to prevent taste problems from such drugs using prophylactic zinc sulfate or amifostine have proven to be minimally beneficial. Although antiepileptic medications are occasionally used to treat smell or taste disturbances, the use of topiramate has been reported to result in a reversible loss of an ability to detect and recognize tastes and odors during treatment.

As with olfaction, a number of systemic disorders can affect taste. These include chronic renal failure, end-stage liver disease, vitamin and mineral deficiencies, diabetes mellitus, and hypothyroidism (to name a few). In diabetes, there appears to be a progressive loss of taste beginning with glucose and then extending to other sweeteners, salty stimuli, and then all stimuli. Psychiatric conditions can be associated with chemosensory alterations (e.g., depression, schizophrenia, bulimia). A recent review of tactile, gustatory, and olfactory hallucinations demonstrated that no one type of hallucinatory experience is pathognomonic to any given diagnosis.

Pregnancy proves to be a unique condition with regard to taste function. There appears to be an increase in dislike and intensity of bitter tastes during the first trimester that may help to ensure that pregnant women avoid poisons during a critical phase of fetal development. Similarly, a relative increase in the preference for salt and bitter in the second and third trimesters may support the ingestion of much needed electrolytes to expand fluid volume and support a varied diet.

CLINICAL EVALUATION

In most cases, a careful clinical history will establish the probable etiology of a chemosensory problem, including questions about its nature, onset, duration, and pattern of fluctuations. *Sudden loss* suggests the possibility of head trauma, ischemia, infection, or a psychiatric condition. *Gradual loss* can reflect the development of a progressive obstructive lesion. *Intermittent loss* suggests the likelihood of an inflammatory process. The patient should be asked about potential precipitating events, such as cold or flu infections prior to symptom onset, because these often go underappreciated. Information regarding head trauma, smoking habits, drug and alcohol abuse (e.g., intranasal cocaine, chronic alcoholism in the context of Wernicke’s and Korsakoff’s syndromes), exposures to pesticides and other toxic agents, and medical interventions is also informative. A determination of all the medications that the patient was taking before and at the time of symptom onset is important, because many can cause chemosensory disturbances. Comorbid medical conditions associated with smell impairment, such as renal failure, liver disease, hypothyroidism, diabetes, or dementia, should be assessed. Delayed puberty in association with anosmia (with or without midline craniofacial abnormalities, deafness, and renal anomalies) suggests the possibility of Kallmann’s syndrome. Recollection of epistaxis, discharge (clear, purulent, or bloody), nasal obstruction, allergies, and somatic symptoms, including headache or irritation, may have localizing value. Questions related to memory, parkinsonian signs, and seizure activity (e.g., automatisms, blackouts,

auras, déjà vu) should be posed. Pending litigation and the possibility of malingering should be considered. Modern forced-choice olfactory tests can detect malingering from improbable responses.

Neurologic and otorhinolaryngologic (ORL) examinations, along with appropriate brain and nasosinus imaging, aid in the evaluation of patients with olfactory or gustatory complaints. The neural evaluation should focus on cranial nerve function, with particular attention to possible skull base and intracranial lesions. Visual acuity, field, and optic disc examinations aid in the detection of intracranial mass lesions that induce intracranial pressure (papilledema) and optic atrophy, especially when considering Foster Kennedy syndrome. The ORL examination should thoroughly assess the intranasal architecture and mucosal surfaces. Polyps, masses, and adhesions of the turbinates to the septum may compromise the flow of air to the olfactory receptors, because less than a fifth of the inspired air traverses the olfactory cleft in the unobstructed state. Blood tests may be helpful to identify such conditions as diabetes, infection, heavy metal exposure, nutritional deficiency (e.g., vitamin B₆ or B₁₂), allergy, and thyroid, liver, and kidney disease.

As with other sensory disorders, quantitative sensory testing is advised. Self-reports of patients can be misleading, and a number of patients who complain of chemosensory dysfunction have normal function for their age and gender. Quantitative smell and taste testing provides valid information for worker's compensation and other legal claims, as well as a way to accurately assess treatment interventions. A number of standardized olfactory and taste tests are commercially available. Most evaluate the ability of patients to detect and identify odors or tastes. For example, the most widely used of these tests, the 40-item University of Pennsylvania Smell Identification Test (UPSIT), uses norms based on nearly 4000 normal subjects. A determination is made of both absolute dysfunction (i.e., mild loss, moderate loss, severe loss, total loss, probable malingering) and relative dysfunction (percentile rank for age and gender). Although electrophysiologic testing is available at some smell and taste centers (e.g., odor event-related potentials), they require complex stimulus presentation and recording equipment and rarely provide additional diagnostic information. With the exception of electrogustometers, commercially available taste tests have only recently become available. Most use filter paper strips impregnated with tastants, so no stimulus preparation is required.

TREATMENT AND MANAGEMENT

Given the various mechanisms by which olfactory and gustatory disturbance can occur, management of patients tends to be condition specific. For example, patients with hypothyroidism, diabetes, or infections often benefit from specific treatments to correct the underlying disease process that is adversely influencing chemoreception. For most patients who present primarily with obstructive/transport loss affecting the nasal and paranasal regions (e.g., allergic rhinitis, polypoid, intranasal neoplasms, nasal deviations), medical and/or surgical intervention is often beneficial. Antifungal and antibiotic treatments may reverse taste problems secondary to candidiasis or other oral infections. Chlorhexidine mouthwash mitigates some salty or bitter dysgeusias, conceivably as a result of its strong positive charge. Excessive dryness of the oral mucosa is a problem with many medications and conditions, and artificial saliva (e.g., Xerolube) or oral pilocarpine treatments may prove beneficial. Other methods to improve salivary flow include the use of mints, lozenges, or sugarless gum. Flavor enhancers may make food more palatable (e.g., monosodium glutamate), but caution is advised to avoid overusing ingredients containing sodium or sugar, particularly in circumstances when a patient also has underlying hypertension or diabetes. Medications that induce distortions of taste can often be discontinued and replaced with other types of medications or modes of therapy. As mentioned earlier, pharmacologic agents result in taste disturbances much more frequently than smell disturbances, and over 250 medications have been reported to alter the sense of taste. It is important to note, however, that many drug-related effects are long lasting and not reversed by short-term drug discontinuance.

A recent study of endoscopic sinus surgery in patients with chronic rhinosinusitis and hyposmia revealed that patients with severe olfactory dysfunction prior to the surgery had a more dramatic and sustained improvement over time compared to patients with more mild olfactory dysfunction prior to intervention. In the case of intranasal and sinus-related inflammatory conditions, such as seen with allergy, viruses, and traumas, the use of intranasal or systemic glucocorticoids may also be helpful. One common approach is to use a tapering course of oral prednisone. The utility of restoring olfaction with either topical or systemic glucocorticoids has been studied. Topical intranasal administration was found to be less effective in general than systemic administration; however, the effects of different nasal administration techniques were not analyzed; for example, intranasal glucocorticoids are more effective if administered in the Moffett's position (head in the inverted position such as over the edge of the bed with the bridge of the nose perpendicular to the floor). After head trauma, an initial trial of glucocorticoids may help to reduce local edema and the potential deleterious deposition of scar tissue around olfactory fila at the level of the cribriform plate.

Treatments are limited for patients with chemosensory loss or primary injury to neural pathways. Nonetheless, spontaneous recovery can occur. In a follow-up study of 542 patients presenting to our center with smell loss from a variety of causes, modest improvement occurred over an average time period of 4 years in about half of the participants. However, only 11% of the anosmic and 23% of the hyposmic patients regained normal age-related function. Interestingly, the amount of dysfunction present at the time of presentation, not etiology, was the best predictor of prognosis. Other predictors were age and the duration of dysfunction prior to initial testing.

A nonblinded study has reported that patients with hyposmia may benefit from smelling strong odors (e.g., eucalyptol, citronella, eugenol, and phenyl ethyl alcohol) before going to bed and immediately upon awakening each day over the course of several months. The rationale for such an approach comes from animal studies demonstrating that prolonged exposure to odorants can induce increased neural activity within the olfactory bulb. In an uncontrolled study, α -lipoic acid (400 mg/d), an essential cofactor for many enzyme complexes with possible antioxidant effects, was reported to be beneficial in mitigating smell loss following viral infection of the upper respiratory tract; controlled studies are needed to confirm this observation. This agent has also been suggested to be useful in some cases of hypogeusia and BMS.

The use of zinc and vitamin A in treating olfactory disturbances is controversial, and there does not appear to be much benefit beyond replenishing established deficiencies. However, zinc has been shown to improve taste function secondary to hepatic deficiencies, and retinoids (bioactive vitamin A derivatives) are known to play an essential role in the survival of olfactory neurons. One protocol in which zinc was infused with chemotherapy treatments suggested a possible protective effect against developing taste impairment. Diseases of the alimentary tract can not only influence chemoreceptive function, but also occasionally influence vitamin B₁₂ absorption. This can result in a relative deficiency of vitamin B₁₂, theoretically contributing to olfactory nerve disturbance. Vitamin B₂ (riboflavin) and magnesium supplements are reported in the alternative literature to aid in the management of migraine that, in turn, may be associated with smell dysfunction. Because vitamin D deficiency is a cofactor of chemotherapy-induced mucocutaneous toxicity and dysgeusia, adding vitamin D₃, 1000–2000 units per day, may benefit some patients with smell and taste complaints during or following chemotherapy.

A number of medications have reportedly been used with success in ameliorating olfactory symptoms, although strong scientific evidence for efficacy is generally lacking. A report that theophylline improved smell function was uncontrolled and failed to account for the fact that some meaningful improvement occurs without treatment; indeed, the percentage of responders was about the same (~50%) as that noted by others to show spontaneous improvement over a similar time period. Antiepileptics and some antidepressants (e.g., amitriptyline) have been used to treat dysosmias and smell distortions, particularly following head trauma. Ironically, amitriptyline is also frequently on the list of medications that can ultimately distort smell and taste function,

possibly from its anticholinergic effects. A recent study suggests that the use of the centrally acting acetylcholinesterase inhibitor donepezil in AD resulted in improvements on smell identification measures that correlated with overall clinician-based impressions of change in dementia severity scores.

Alternative therapies, such as acupuncture, meditation, cognitive-behavioral therapy, and yoga, can help patients manage uncomfortable experiences associated with chemosensory disturbance and oral pain syndromes and to cope with the psychosocial stressors surrounding the impairment. Additionally, modification of diet and eating habits is also important. By accentuating the other sensory experiences of a meal, such as food texture, aroma, temperature, and color, one can optimize the overall eating experience for a patient. In some cases, a flavor enhancer like monosodium glutamate (MSG) can be added to foods to increase palatability and encourage intake.

Proper oral and nasal hygiene and routine dental care are extremely important ways for patients to protect themselves from disorders of the mouth and nose that can ultimately result in chemosensory disturbance. Patients should be warned not to overcompensate for their taste loss by adding excessive amounts of sugar or salt. Smoking cessation and the discontinuance of oral tobacco use are essential in the management of any patient with smell and/or taste disturbance and should be repeatedly emphasized.

A major and often overlooked element of therapy comes from chemosensory testing itself. Confirmation or lack of confirmation of loss is beneficial to patients who come to believe, in light of unsupportive family members and medical providers, that they may be “crazy.” In cases where the loss is minor, patients can be informed of the likelihood of a more positive prognosis. Importantly, quantitative testing places the patient’s problem into overall perspective. Thus, it is often therapeutic for an older person to know that, while his or her smell function is not what it used to be, it still falls above the average of his or her peer group. Without testing, many such patients are simply told they are getting old and nothing can be done for them, leading in some cases to depression and decreased self-esteem.

low-frequency sounds, the point of maximal displacement is toward the apex of the cochlea.

The inner and outer hair cells of the organ of Corti have different innervation patterns, but both are mechanoreceptors. The afferent innervation relates principally to the inner hair cells, and the efferent innervation relates principally to outer hair cells. The motility of the outer hair cells alters the micromechanics of the inner hair cells, creating a cochlear amplifier, which explains the exquisite sensitivity and frequency selectivity of the cochlea.

Beginning in the cochlea, the frequency specificity is maintained at each point of the central auditory pathway: dorsal and ventral cochlear nuclei, trapezoid body, superior olivary complex, lateral lemniscus, inferior colliculus, medial geniculate body, and auditory cortex. At low frequencies, individual auditory nerve fibers can respond more or less synchronously with the stimulating tone. At higher frequencies, phase-locking occurs so that neurons alternate in response to particular phases of the cycle of the sound wave. Intensity is encoded by the amount of neural activity in individual neurons, the number of neurons that are active, and the specific neurons that are activated.

There is evidence that the right and left ears as well as the central nervous system may process speech asymmetrically. Generally, a sound is processed symmetrically from the peripheral to the central auditory system. However, a “right ear advantage” exists for dichotic listening tasks, in which subjects are asked to report on competing sounds presented to each ear. In most individuals, a perceptual right ear advantage for consonant-vowel syllables, stop consonants, and words also exists. Similarly, whereas central auditory processing for sounds is symmetric with minimal lateral specialization for the most part, speech processing is lateralized. There is specialization of the left auditory cortex for speech recognition and production, and of the right hemisphere for emotional and tonal aspects of speech. Left hemisphere dominance for speech is found in 95–98% of right-handed persons and 70–80% of left-handed persons.

DISORDERS OF THE SENSE OF HEARING

Hearing loss can result from disorders of the auricle, external auditory canal, middle ear, inner ear, or central auditory pathways (Fig. 43-2). *In general, lesions in the auricle, external auditory canal, or middle ear that impede the transmission of sound from the external environment to the inner ear cause conductive hearing loss, whereas lesions that impair mechanotransduction in the inner ear or transmission of the electrical signal along the eighth nerve to the brain cause sensorineural hearing loss.*

Conductive Hearing Loss The external ear, the external auditory canal, and the middle ear apparatus is designed to collect and amplify sound and efficiently transfer the mechanical energy of the sound wave to the fluid-filled cochlea. Factors that obstruct the transmission of sound or serve to dampen the acoustical energy result in conductive hearing loss. Conductive hearing loss can occur from obstruction of the external auditory canal by cerumen, debris, and foreign bodies; swelling of the lining of the canal; atresia or neoplasms of the canal; perforations of the tympanic membrane; disruption of the ossicular chain, as occurs with necrosis of the long process of the incus in trauma or infection; otosclerosis; or fluid, scarring, or neoplasms in the middle ear. Rarely, inner ear malformations or pathologies, such as superior semicircular canal dehiscence, lateral semicircular canal dysplasia, incomplete partition of the inner ear, and large vestibular aqueduct, may also be associated with conductive hearing loss.

Eustachian tube dysfunction is extremely common in adults and may predispose to acute otitis media (AOM) or serous otitis media (SOM). Trauma, AOM, and chronic otitis media are the usual factors responsible for tympanic membrane perforation. While small perforations often heal spontaneously, larger defects usually require surgical intervention. Tympanoplasty is highly effective (>90%) in the repair of tympanic membrane perforations. Otoscopy is usually sufficient to diagnose AOM, SOM, chronic otitis media, cerumen impaction, tympanic membrane perforation, and eustachian tube dysfunction; tympanometry can be useful to confirm the clinical suspicion of these conditions.

43

Disorders of Hearing

Anil K. Lalwani

Hearing loss is one of the most common sensory disorders in humans and can present at any age. Nearly 10% of the adult population has some hearing loss, and one-third of individuals age >65 years have a hearing loss of sufficient magnitude to require a hearing aid.

PHYSIOLOGY OF HEARING

The function of the external and middle ear is to amplify sound to facilitate conversion of the mechanical energy of the sound wave into an electrical signal by the inner ear hair cells, a process called mechanotransduction (Fig. 43-1). Sound waves enter the external auditory canal and set the tympanic membrane (eardrum) in motion, which in turn moves the malleus, incus, and stapes of the middle ear. Movement of the footplate of the stapes causes pressure changes in the fluid-filled inner ear, eliciting a traveling wave in the basilar membrane of the cochlea. The tympanic membrane and the ossicular chain in the middle ear serve as an impedance-matching mechanism, improving the efficiency of energy transfer from air to the fluid-filled inner ear.

Stereocilia of the hair cells of the organ of Corti, which rests on the basilar membrane, are in contact with the tectorial membrane and are deformed by the traveling wave. A point of maximal displacement of the basilar membrane is determined by the frequency of the stimulating tone. High-frequency tones cause maximal displacement of the basilar membrane near the base of the cochlea, whereas for

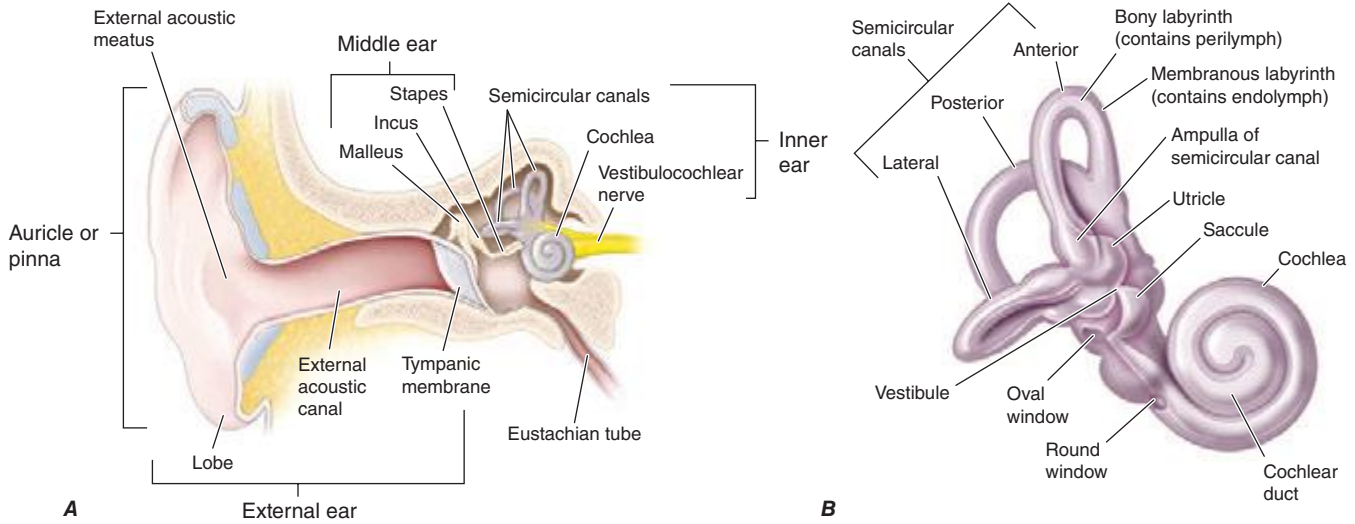


FIGURE 43-1 Ear anatomy. **A.** Drawing of modified coronal section through external ear and temporal bone, with structures of the middle and inner ear demonstrated. **B.** High-resolution view of inner ear.

Cholesteatoma, a benign tumor composed of stratified squamous epithelium in the middle ear or mastoid, occurs frequently in adults. This is a slowly growing lesion that destroys bone and normal ear tissue. Theories of pathogenesis include traumatic immigration and invasion of squamous epithelium through a retraction pocket, implantation of squamous epithelia in the middle ear through a perforation or surgery, and metaplasia following chronic infection and irritation.

On examination, there is often a perforation of the tympanic membrane filled with cheesy white squamous debris. The presence of an aural polyp obscuring the tympanic membrane is highly suggestive of an underlying cholesteatoma. A chronically draining ear that fails to respond to appropriate antibiotic therapy should raise suspicion of a cholesteatoma. Conductive hearing loss secondary to ossicular erosion is common. Surgery is required to remove this destructive process.

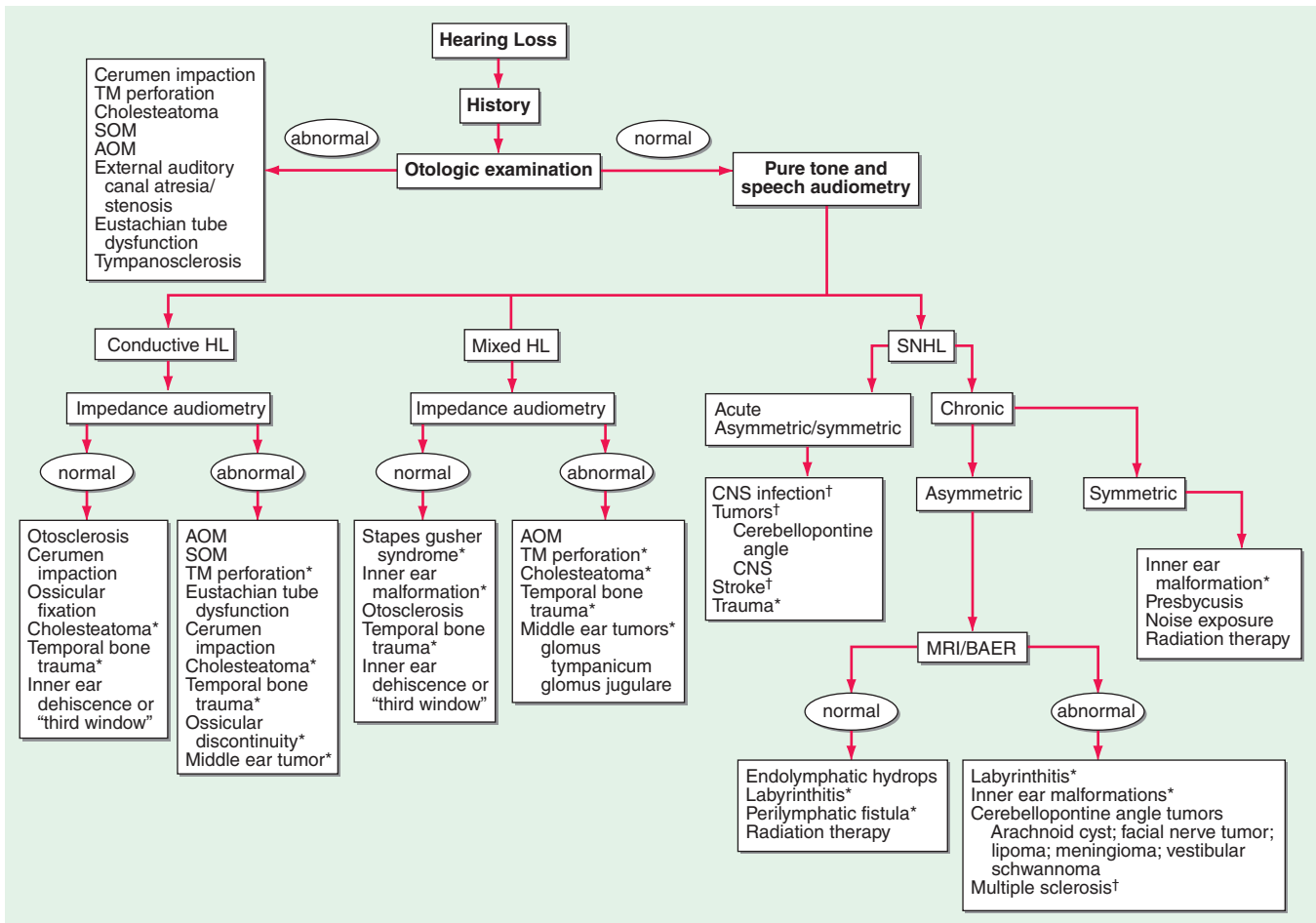


FIGURE 43-2 An algorithm for the approach to hearing loss. AOM, acute otitis media; BAER, brainstem auditory evoked response; CNS, central nervous system; HL, hearing loss; SNHL, sensorineural hearing loss; SOM, serous otitis media; TM, tympanic membrane. *Computed tomography scan of temporal bone. †Magnetic resonance imaging (MRI) scan.

Conductive hearing loss with a normal ear canal and intact tympanic membrane suggests either ossicular pathology or the presence of “third window” in the inner ear (see below). Fixation of the stapes from *otosclerosis* is a common cause of low-frequency conductive hearing loss. It occurs equally in men and women and is inherited as an autosomal dominant trait with incomplete penetrance; in some cases, it may be a manifestation of *osteogenesis imperfecta*. Hearing impairment usually presents between the late teens and the forties. In women, the otosclerotic process is accelerated during pregnancy, and the hearing loss is often first noticeable at this time. A hearing aid or a simple outpatient surgical procedure (stapedectomy) can provide adequate auditory rehabilitation. Extension of otosclerosis beyond the stapes footplate to involve the cochlea (cochlear otosclerosis) can lead to mixed or sensorineural hearing loss. Fluoride therapy to prevent hearing loss from cochlear otosclerosis is of uncertain value.

Disorders that lead to the formation of a pathologic “third window” in the inner ear can be associated with conductive hearing loss. There are normally two major openings, or windows, that connect the inner ear with the middle ear and serve as conduits for transmission of sound; these are, respectively, the oval and round windows. A third window is formed where the normally hard otic bone surrounding the inner ear is eroded; dissipation of the acoustic energy at the third window is responsible for the “inner ear conductive hearing loss.” The superior semicircular canal dehiscence syndrome resulting from erosion of the otic bone over the superior circular canal can present with conductive hearing loss that mimics otosclerosis. A common symptom is vertigo evoked by loud sounds (Tullio phenomenon), by Valsalva maneuvers that change middle ear pressure, or by applying positive pressure on the tragus (the cartilage anterior to the external opening of the ear canal). Patients with this syndrome also complain of being able to hear the movement of their eyes and neck. A large jugular bulb or jugular bulb diverticulum can create a “third window” by eroding into the vestibular aqueduct or posterior semicircular canal; the symptoms are similar to those of the superior semicircular canal dehiscence syndrome.

Sensorineural Hearing Loss Sensorineural hearing loss results from either damage to the mechanotransduction apparatus of the cochlea or disruption of the electrical conduction pathway from the inner ear to the brain. Thus, injury to hair cells, supporting cells, auditory neurons, or the central auditory pathway can cause sensorineural hearing loss. Damage to the hair cells of the organ of Corti may be caused by intense noise, viral infections, ototoxic drugs (e.g., salicylates, quinine and its synthetic analogues, aminoglycoside antibiotics, loop diuretics such as furosemide and ethacrynic acid, and cancer chemotherapeutic agents such as cisplatin), fractures of the temporal bone, meningitis, cochlear otosclerosis (see above), Ménière’s disease, and aging. Congenital malformations of the inner ear may be the cause of hearing loss in some adults. Genetic predisposition alone or in concert with environmental exposures may also be responsible (see below).

Presbycusis (age-associated hearing loss) is the most common cause of sensorineural hearing loss in adults. In the early stages, it is characterized by symmetric, gentle to sharply sloping high-frequency hearing loss (Fig. 43-3). With progression, the hearing loss involves all frequencies. More importantly, the hearing impairment is associated with significant loss in clarity. There is a loss of discrimination for phonemes, recruitment (abnormal growth of loudness), and particular difficulty in understanding speech in noisy environments such as at restaurants and social events. Hearing aids are helpful in enhancing the signal-to-noise ratio by amplifying sounds that are close to the listener. *Although hearing aids are able to amplify sounds, they cannot restore the clarity of hearing.* Thus, amplification with hearing aids may provide only limited rehabilitation once the word recognition score deteriorates below 50%. Cochlear implants are the treatment of choice when hearing aids prove inadequate, even when hearing loss is incomplete (see below).

Ménière’s disease is characterized by episodic vertigo, fluctuating sensorineural hearing loss, tinnitus, and aural fullness. Tinnitus and/or deafness may be absent during the initial attacks of vertigo, but it

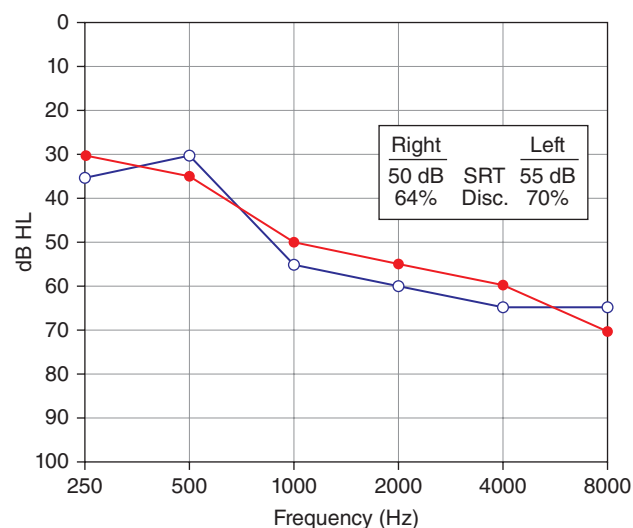


FIGURE 43-3 Presbycusis or age-related hearing loss. The audiogram shows a moderate to severe downsloping sensorineural hearing loss typical of presbycusis. The loss of high-frequency hearing is associated with a decreased speech discrimination score; consequently, patients complain of lack of clarity of hearing, especially in a noisy background. HL, hearing threshold level; SRT, speech reception threshold.

invariably appears as the disease progresses and increases in severity during acute attacks. The annual incidence of Ménière’s disease is 0.5–7.5 per 1000; onset is most frequently in the fifth decade of life but may also occur in young adults or the elderly. Histologically, there is distention of the endolymphatic system (endolymphatic hydrops) leading to degeneration of vestibular and cochlear hair cells. This may result from endolymphatic sac dysfunction secondary to infection, trauma, autoimmune disease, inflammatory causes, or tumor; an idiopathic etiology constitutes the largest category and is most accurately referred to as Ménière’s disease. Although any pattern of hearing loss can be observed, typically, low-frequency, unilateral sensorineural hearing impairment is present. Magnetic resonance imaging (MRI) should be obtained to exclude retrocochlear pathology such as a cerebellopontine angle tumor or demyelinating disorder. Therapy is directed toward the control of vertigo. A 2-g/d low-salt diet is the mainstay of treatment for control of rotatory vertigo. Diuretics, a short course of glucocorticoids, and intratympanic gentamicin may also be useful adjuncts in recalcitrant cases. Surgical therapy of vertigo is reserved for unresponsive cases and includes endolymphatic sac decompression, labyrinthectomy, and vestibular nerve section. Both labyrinthectomy and vestibular nerve section abolish rotatory vertigo in >90% of cases. Unfortunately, there is no effective therapy for hearing loss, tinnitus, or aural fullness from Ménière’s disease.

Sensorineural hearing loss may also result from any neoplastic, vascular, demyelinating, infectious, or degenerative disease or trauma affecting the central auditory pathways. HIV leads to both peripheral and central auditory system pathology and is associated with sensorineural hearing impairment.

Primary diseases of the central nervous system can also present with hearing impairment. Characteristically, a reduction in clarity of hearing and speech comprehension is much greater than the loss of the ability to hear pure tone. Auditory testing is consistent with an auditory neuropathy; normal otoacoustic emissions (OAE) and an abnormal auditory brainstem response (ABR) is typical (see below). Hearing loss can accompany hereditary sensorimotor neuropathies and inherited disorders of myelin. Tumors of the cerebellopontine angle such as vestibular schwannoma and meningioma usually present with asymmetric sensorineural hearing loss with greater deterioration of speech understanding than pure tone hearing. Multiple sclerosis may present with acute unilateral or bilateral hearing loss; typically, pure tone testing remains relatively stable while speech understanding

fluctuates. Isolated labyrinthine infarction can present with acute hearing loss and vertigo due to a cerebrovascular accident involving the posterior circulation, usually the anterior inferior cerebellar artery; it may also be the heralding sign of impending catastrophic basilar artery infarction (**Chap. 446**).

A finding of conductive and sensory hearing loss in combination is termed *mixed hearing loss*. Mixed hearing losses are due to pathology of both the middle and inner ear, as can occur in otosclerosis involving the ossicles and the cochlea, head trauma, chronic otitis media, cholesteatoma, middle ear tumors, and some inner ear malformations.

Trauma resulting in temporal bone fractures may be associated with conductive, sensorineural, or mixed hearing loss. If the fracture spares the inner ear, there may simply be conductive hearing loss due to rupture of the tympanic membrane or disruption of the ossicular chain. These abnormalities can be surgically corrected. Profound hearing loss and severe vertigo are associated with temporal bone fractures involving the inner ear. A perilymphatic fistula associated with leakage of inner ear fluid into the middle ear can occur and may require surgical repair. An associated facial nerve injury is not uncommon. Computed tomography (CT) is best suited to assess fracture of the traumatized temporal bone, evaluate the ear canal, and determine the integrity of the ossicular chain and the involvement of the inner ear. Cerebrospinal fluid leaks that accompany temporal bone fractures are usually self-limited; the value of prophylactic antibiotics is uncertain.

Tinnitus is defined as the perception of a sound when there is no sound in the environment. It may have a buzzing, roaring, or ringing quality and may be pulsatile (synchronous with the heartbeat). Tinnitus is often associated with either a conductive or sensorineural hearing loss. The pathophysiology of tinnitus is not well understood. The cause of the tinnitus can usually be determined by finding the cause of the associated hearing loss. Tinnitus may be the first symptom of a serious condition such as a vestibular schwannoma. Pulsatile tinnitus requires evaluation of the vascular system of the head to exclude vascular tumors such as glomus jugulare tumors, aneurysms, dural arteriovenous fistulas, and stenotic arterial lesions; it may also occur with SOM. It is most commonly associated with some abnormality of the jugular bulb such as a large jugular bulb or jugular bulb diverticulum.

GENETIC CAUSES OF HEARING LOSS



More than half of childhood hearing impairment is thought to be hereditary; hereditary hearing impairment (HHI) can also manifest later in life. HHI may be classified as either nonsyndromic, when hearing loss is the only clinical abnormality, or syndromic, when hearing loss is associated with anomalies in other organ systems. Nearly two-thirds of HHIs are nonsyndromic, and the remaining one-third are syndromic. Between 70 and 80% of nonsyndromic HHI is inherited in an autosomal recessive manner and designated DFNB; another 15–20% is autosomal dominant (DFNA). Less than 5% is X-linked (DFNX) or maternally inherited via the mitochondria.

More than 150 loci harboring genes for nonsyndromic HHI have been mapped, with recessive loci outnumbering dominant; numerous genes have now been identified (**Table 43-1**). The hearing genes fall into the categories of structural proteins (*MYH9*, *MYO7A*, *MYO15*, *TECTA*, *DIAPH1*), transcription factors (*POU3F4*, *POU4F3*), ion channels (*KCNQ4*, *SLC26A4*), and gap junction proteins (*GJB2*, *GJB3*, *GJB6*). Several of these genes, including *GJB2*, *TECTA*, and *TMCI*, cause both autosomal dominant and recessive forms of nonsyndromic HHI. In general, the hearing loss associated with dominant genes has its onset in adolescence or adulthood, varies in severity, and progresses with age, whereas the hearing loss associated with recessive inheritance is congenital and profound. Connexin 26, product of the *GJB2* gene, is particularly important because it is responsible for nearly 20% of all cases of childhood deafness; half of genetic deafness in children is *GJB2*-related. Two frameshift mutations, 35delG and 167delT, account for >50% of the cases; however, screening for these two mutations alone is insufficient, and sequencing of the entire gene is required to diagnose *GJB2*-related recessive deafness. The 167delT mutation is highly prevalent in Ashkenazi Jews; ~1 in 1765 individuals in this

population are homozygous and affected. The hearing loss can also vary among the members of the same family, suggesting that other genes or factors influence the auditory phenotype.

In addition to *GJB2*, several other nonsyndromic genes are associated with hearing loss that progresses with age. The contribution of genetics to presbycusis is also becoming better understood. Sensitivity to aminoglycoside ototoxicity can be maternally transmitted through a mitochondrial mutation. Susceptibility to noise-induced hearing loss may also be genetically determined.

There are >400 syndromic forms of hearing loss. These include Usher's syndrome (retinitis pigmentosa and hearing loss), Waardenburg's syndrome (pigmentary abnormality and hearing loss), Pendred's syndrome (thyroid organification defect and hearing loss), Alport's syndrome (renal disease and hearing loss), Jervell and Lange-Nielsen syndrome (prolonged QT interval and hearing loss), neurofibromatosis type 2 (bilateral acoustic schwannoma), and mitochondrial disorders (mitochondrial encephalopathy, lactic acidosis, and stroke-like episodes [MELAS]; myoclonic epilepsy and ragged red fibers [MERRF]; progressive external ophthalmoplegia [PEO]) (**Table 43-2**).

APPROACH TO THE PATIENT: Disorders of the Sense of Hearing

The goal in the evaluation of a patient with auditory complaints is to determine (1) the nature of the hearing impairment (conductive vs sensorineural vs mixed), (2) the severity of the impairment (mild, moderate, severe, or profound), (3) the anatomy of the impairment (external ear, middle ear, inner ear, or central auditory pathway), and (4) the etiology. The history should elicit characteristics of the hearing loss, including the duration of deafness, unilateral versus bilateral involvement, nature of onset (sudden vs insidious), and rate of progression (rapid vs slow). Symptoms of tinnitus, vertigo, imbalance, aural fullness, otorrhea, headache, facial nerve dysfunction, and head and neck paresthesias should be noted. Information regarding head trauma, exposure to ototoxins, occupational or recreational noise exposure, and family history of hearing impairment may also be important. A sudden onset of unilateral hearing loss, with or without tinnitus, may represent a viral infection of the inner ear, vestibular schwannoma, or a stroke. Patients with unilateral hearing loss (sensory or conductive) usually complain of reduced hearing, poor sound localization, and difficulty hearing clearly with background noise. Gradual progression of a hearing deficit is common with otosclerosis, noise-induced hearing loss, vestibular schwannoma, or Ménière's disease. Small vestibular schwannomas typically present with asymmetric hearing impairment, tinnitus, and imbalance (rarely vertigo); cranial neuropathy, in particular of the trigeminal or facial nerve, may accompany larger tumors. In addition to hearing loss, Ménière's disease may be associated with episodic vertigo, tinnitus, and aural fullness. Hearing loss with otorrhea is most likely due to chronic otitis media or cholesteatoma.

Examination should include the auricle, external ear canal, and tympanic membrane. The external ear canal of the elderly is often dry and fragile; it is preferable to clean cerumen with wall-mounted suction or cerumen loops and to avoid irrigation. In examining the eardrum, the topography of the tympanic membrane is more important than the presence or absence of the light reflex. In addition to the pars tensa (the lower two-thirds of the tympanic membrane), the pars flaccida (upper one-third of the tympanic membrane) above the short process of the malleus should also be examined for retraction pockets that may be evidence of chronic eustachian tube dysfunction or cholesteatoma. Insufflation of the ear canal is necessary to assess tympanic membrane mobility and compliance. Careful inspection of the nose, nasopharynx, and upper respiratory tract is indicated. Unilateral serous effusion should prompt a fiberoptic examination of the nasopharynx to exclude neoplasms. Cranial nerves should be evaluated with special attention to facial and trigeminal nerves, which are commonly affected with tumors involving the cerebellopontine angle.

TABLE 43-1 HEREDITARY HEARING IMPAIRMENT GENES

Designation	Gene	Function	Designation	Gene	Function
Autosomal Dominant			DFNB12	<i>CDH23</i>	Intercellular adherence protein
DFNA1	<i>CRYM</i>	Thyroid hormone-binding protein	DFNB15/72/95	<i>GIPC3</i>	PDZ domain containing protein
DFNA2A	<i>DIAPH1</i>	Cytoskeletal protein	DFNB16	<i>STRC</i>	Stereocilia protein
DFNA2B	<i>KCNQ4</i>	Potassium channel	DFNB18	<i>USH1C</i>	Unknown
DFNA3A	<i>GJB3 (Cx31)</i>	Gap junction	DFNB21	<i>TECTA</i>	Tectorial membrane protein
DFNA3B	<i>GJB2 (Cx26)</i>	Gap junction	DFNB22	<i>OTOA</i>	Gel attachment to nonsensory cell
DFNA4	<i>GJB6 (Cx30)</i>	Gap junction	DFNB23	<i>PCDH15</i>	Morphogenesis and cohesion
	<i>MYH14</i>	Class II nonmuscle myosin	DFNB24	<i>RDX</i>	Cytoskeletal protein
	<i>CEACAM16</i>	Cell adhesion molecule	DFNB25	<i>GRXCR1</i>	Reversible S-glutathionylation of proteins
DFNA5	<i>DFNA5</i>	Unknown	DFNB28	<i>TRIOBP</i>	Cytoskeletal-organizing protein
DFNA6/14/38	<i>WFS1</i>	Transmembrane protein	DFNB29	<i>CLDN14</i>	Tight junctions
DFNA8/12	<i>TECTA</i>	Tectorial membrane protein	DFNB30	<i>MYO3A</i>	Hybrid motor-signaling myosin
DFNA9	<i>COCH</i>	Unknown	DFNB31	<i>WHRN</i>	PDZ domain-containing protein
DFNA10	<i>EYA4</i>	Developmental gene	DFNB35	<i>ESRRB</i>	Estrogen-related receptor beta protein
DFNA11	<i>MYO7A</i>	Cytoskeletal protein	DFNB36	<i>ESPN</i>	Ca-insensitive actin-bundling protein
DFNA13	<i>COL11A2</i>	Cytoskeletal protein	DFNB37	<i>MYO6</i>	Unconventional myosin
DFNA15	<i>POU4F3</i>	Transcription factor	DFNB39	<i>HFG</i>	Hepatocyte growth factor
DFNA17	<i>MYH9</i>	Cytoskeletal protein	DFNB42	<i>ILDR1</i>	Ig-like domain-containing receptor
DFNA20/26	<i>ACTG1</i>	Cytoskeletal protein	DFNB48	<i>CIB2</i>	Calcium and integrin binding protein
DFNA22	<i>MYO6</i>	Unconventional myosin	DFNB49	<i>MARVELD2</i>	Tight junction protein
DFNA23	<i>SIX1</i>	Developmental gene	DFNB53	<i>COL11A2</i>	Collagen protein
DFNA25	<i>SLC17A8</i>	Vesicular glutamate transporter	DFNB59	<i>PJVK</i>	Zn-binding protein
DFNA28	<i>TFCP2L3</i>	Transcription factor	DFNB61	<i>SLC26A5</i>	Motor protein
DFNA36	<i>TMC1</i>	Transmembrane protein	DFNB63	<i>LRTOMT/COMT2</i>	Putative methyltransferase
DFNA41	<i>P2RX2</i>	Purinergic receptor	DFNB66/67	<i>LHFPL5</i>	Tetraspan protein
DFNA44	<i>CCDC50</i>	Effector of epidermal growth factor-mediated signaling	DFNB70	<i>PNPT1</i>	Mitochondrial-RNA-import protein
DFNA48	<i>MYO1A</i>	Unconventional myosin	DFNB74	<i>MSRB3</i>	Methionine sulfoxide reductase
DFNA50	<i>MIRN96</i>	MicroRNA	DFNB77	<i>LOXHD1</i>	Stereociliary protein
DFNA51	<i>TJP2</i>	Tight junction protein	DFNB79	<i>TPRN</i>	Unknown
DFNA56	<i>TNC</i>	Extracellular matrix protein	DFNB82	<i>GPSM2</i>	G protein signaling modulator
DFNA64	<i>SMAC/DIABLO</i>	Mitochondrial proapoptotic protein	DFNB84	<i>PTPRQ</i>	Type III receptor-like protein-tyrosine phosphatase family
Autosomal Recessive			DFNB86	<i>TBC1D24</i>	GTPase-activating protein
DFNB1A	<i>GJB2 (CX26)</i>	Gap junction	DFNB88	<i>ELMOD3</i>	GTPase-activating protein
DFNB1B	<i>GJB6 (CX30)</i>	Gap junction	DFNB89	<i>KARS</i>	Lysyl-tRNA synthetase
DFNB2	<i>MYO7A</i>	Cytoskeletal protein	DFNB91	<i>GJB3</i>	Gap junction
DFNB3	<i>MYO15</i>	Cytoskeletal protein	DFNB93	<i>CABP2</i>	Calcium binding protein
DFNB4	<i>PDS (SLC26A4)</i>	Chloride/iodide transporter	DFNB98	<i>TSPEAR</i>	Epilepsy-associated repeats containing protein
DFNB6	<i>TMIE</i>	Transmembrane protein		<i>SERPIN6</i>	Protease inhibitor
DFNB7/B11	<i>TMC1</i>	Transmembrane protein			
DFNB9	<i>OTOF</i>	Trafficking of membrane vesicles			
DFNB8/10	<i>TMPRSS3</i>	Transmembrane serine protease			

The Rinne and Weber tuning fork tests, with a 512-Hz tuning fork, are used to screen for hearing loss, differentiate conductive from sensorineural hearing losses, and confirm the findings of audiologic evaluation. The Rinne test compares the ability to hear by air conduction with the ability to hear by bone conduction. The tines of a vibrating tuning fork are held near the opening of the external auditory canal, and then the stem is placed on the mastoid process; for direct contact, it may be placed on teeth or dentures. The patient is asked to indicate whether the tone is louder by air conduction or bone conduction. Normally, and in the presence of sensorineural hearing loss, a tone is heard louder by air conduction than by bone conduction; however, with conductive hearing loss of ≥ 30 dB (see "Audiologic Assessment," below), the bone-conduction stimulus is perceived as louder than the air-conduction stimulus. For the Weber test, the stem of a vibrating tuning fork is placed on

the head in the midline and the patient is asked whether the tone is heard in both ears or better in one ear than in the other. With a unilateral conductive hearing loss, the tone is perceived in the affected ear. With a unilateral sensorineural hearing loss, the tone is perceived in the unaffected ear. A 5-dB difference in hearing between the two ears is required for lateralization.

LABORATORY ASSESSMENT OF HEARING

Audiologic Assessment The minimum audiologic assessment for hearing loss should include the measurement of pure tone air-conduction and bone-conduction thresholds, speech reception threshold, word recognition score, tympanometry, acoustic reflexes, and acoustic-reflex decay. This test battery provides a screening evaluation of the entire auditory system and allows one to determine whether further

TABLE 43-2 SYNDROMIC HEREDITARY HEARING IMPAIRMENT GENES

Syndrome	Gene	Function
Alport's syndrome	COL4A3-5	Cytoskeletal protein
BOR syndrome	EYA1	Developmental gene
	SIX5	Developmental gene
	SIX1	Developmental gene
Jervell and Lange-Nielsen syndrome	KCNQ1	Delayed rectifier K ⁺ channel
	KCNE1	Delayed rectifier K ⁺ channel
Norrie's disease	NDP	Cell-cell interactions
Pendred's syndrome	SLC26A4	Chloride/iodide transporter
	FOXI1	Transcriptional activator of SLC26A4
	KCNJ10	Inwardly rectifying K ⁺ channel
Treacher Collins syndrome	TCOF1	Nucleolar-cytoplasmic transport
	POLR1D	Subunit of RNA polymerases I and III
	POLR1C	Subunit of RNA polymerases I and III
Usher's syndrome	MYO7A	Cytoskeletal protein
	USH1C	Unknown
	CDH23	Intercellular adherence protein
	PCDH15	Cell adhesion molecule
	SANS	Harmonin-associated protein
	CIB2	Calcium and integrin binding protein
	USH2A	Cell adhesion molecule
	VLGR1	G protein-coupled receptor
	WHRN	PDZ domain-containing protein
	CLRN1	Cellular synapse protein?
PDZD7	PDZ domain containing protein	
WS type I, III	PAX3	Transcription factor
WS type II	MITF	Transcription factor
	SNAI2	Transcription factor
WS type IV	EDNRB	Endothelin B receptor
	EDN3	Endothelin B receptor ligand
	SOX10	Transcription factor

Abbreviations: BOR, branchio-oto-renal syndrome; WS, Waardenburg's syndrome.

differentiation of a sensory (cochlear) from a neural (retrocochlear) hearing loss is indicated.

Pure tone audiometry assesses hearing acuity for pure tones. The test is administered by an audiologist and is performed in a sound-attenuated chamber. The pure tone stimulus is delivered with an audiometer, an electronic device that allows the presentation of specific frequencies (generally between 250 and 8000 Hz) at specific intensities. Air- and bone-conduction thresholds are established for each ear. Air-conduction thresholds are determined by presenting the stimulus in air with the use of headphones. Bone-conduction thresholds are determined by placing the stem of a vibrating tuning fork or an oscillator of an audiometer in contact with the head. In the presence of a hearing loss, broad-spectrum noise is presented to the nontest ear for *masking* purposes so that responses are based on perception from the ear under test.

The responses are measured in decibels. An *audiogram* is a plot of intensity in decibels of hearing threshold versus frequency. A decibel (dB) is equal to 20 times the logarithm of the ratio of the sound pressure required to achieve threshold in the patient to the sound pressure required to achieve threshold in a normal-hearing person. Therefore, a change of 6 dB represents doubling of sound pressure, and a change of 20 dB represents a tenfold change in sound pressure. Loudness, which depends on the frequency, intensity, and duration of a sound, doubles with approximately each 10-dB increase in sound pressure level. Pitch,

on the other hand, does not directly correlate with frequency. The perception of pitch changes slowly in the low and high frequencies. In the middle tones, which are important for human speech, pitch varies more rapidly with changes in frequency.

Pure tone audiometry establishes the presence and severity of hearing impairment, unilateral versus bilateral involvement, and the type of hearing loss. Conductive hearing losses with a large mass component, as is often seen in middle ear effusions, produce elevation of thresholds that predominate in the higher frequencies. Conductive hearing losses with a large stiffness component, as in fixation of the footplate of the stapes in early otosclerosis, produce threshold elevations in the lower frequencies. Often, the conductive hearing loss involves all frequencies, suggesting involvement of both stiffness and mass. In general, sensorineural hearing losses such as presbycusis affect higher frequencies more than lower frequencies (Fig. 43-3). An exception is Ménière's disease, which is characteristically associated with low-frequency sensorineural hearing loss. Noise-induced hearing loss has an unusual pattern of hearing impairment in which the loss at 4000 Hz is greater than at higher frequencies. Vestibular schwannomas characteristically affect the higher frequencies, but any pattern of hearing loss can be observed.

Speech recognition requires greater synchronous neural firing than is necessary for appreciation of pure tones. *Speech audiometry* tests the clarity with which one hears. The *speech reception threshold* (SRT) is defined as the intensity at which speech is recognized as a meaningful symbol and is obtained by presenting two-syllable words with an equal accent on each syllable. The intensity at which the patient can repeat 50% of the words correctly is the SRT. Once the SRT is determined, discrimination or word recognition ability is tested by presenting one-syllable words at 25–40 dB above the SRT. The words are phonetically balanced in that the phonemes (speech sounds) occur in the list of words at the same frequency that they occur in ordinary conversational English. An individual with normal hearing or conductive hearing loss can repeat 88–100% of the phonetically balanced words correctly. Patients with a sensorineural hearing loss have variable loss of discrimination. As a general rule, neural lesions produce greater deficits in discrimination than do cochlear lesions. For example, in a patient with mild asymmetric sensorineural hearing loss, a clue to the diagnosis of vestibular schwannoma is the presence of greater than expected deterioration in discrimination ability. Deterioration in discrimination ability at higher intensities above the SRT also suggests a lesion in the eighth nerve or central auditory pathways.

Tympanometry measures the impedance of the middle ear to sound and is useful in diagnosis of middle ear effusions. A *tympanogram* is the graphic representation of change in impedance or compliance as the pressure in the ear canal is changed. Normally, the middle ear is most compliant at atmospheric pressure, and the compliance decreases as the pressure is increased or decreased (type A); this pattern is seen with normal hearing or in the presence of sensorineural hearing loss. Compliance that does not change with change in pressure suggests middle ear effusion (type B). With a negative pressure in the middle ear, as with eustachian tube obstruction, the point of maximal compliance occurs with negative pressure in the ear canal (type C). A tympanogram in which no point of maximal compliance can be obtained is most commonly seen with discontinuity of the ossicular chain (type A_d). A reduction in the maximal compliance peak can be seen in otosclerosis (type A_s).

During tympanometry, an intense tone elicits contraction of the stapedius muscle. The change in compliance of the middle ear with contraction of the stapedius muscle can be detected. The presence or absence of this *acoustic reflex* is important in determining the etiology of hearing loss as well as in the anatomic localization of facial nerve paralysis. The acoustic reflex can help differentiate between conductive hearing loss due to otosclerosis and that caused by an inner ear "third window"; it is absent in otosclerosis and present in inner ear conductive hearing loss. Normal or elevated acoustic reflex thresholds in an individual with sensorineural hearing impairment suggest a cochlear hearing loss. An absent acoustic reflex in the setting of sensorineural

hearing loss is not helpful in localizing the site of lesion. Assessment of *acoustic reflex decay* helps differentiate sensory from neural hearing losses. In neural hearing loss, such as with vestibular schwannoma, the reflex adapts or decays with time.

OAEs generated by outer hair cells only can be measured with microphones inserted into the external auditory canal. The emissions may be spontaneous or evoked with sound stimulation. The presence of OAEs indicates that the outer hair cells of the organ of Corti are intact and can be used to assess auditory thresholds and to distinguish sensory from neural hearing losses.

Evoked Responses *Electrocochleography* measures the earliest evoked potentials generated in the cochlea and the auditory nerve. Receptor potentials recorded include the cochlear microphonic, generated by the outer hair cells of the organ of Corti, and the summing potential, generated by the inner hair cells in response to sound. The whole nerve action potential representing the composite firing of the first-order neurons can also be recorded during electrocochleography. Clinically, the test is useful in the diagnosis of Ménière's disease, where an elevation of the ratio of summing potential to action potential is seen.

Brainstem auditory evoked responses (BAERs), also known as *auditory brainstem responses* (ABRs), are useful in differentiating the site of sensorineural hearing loss. In response to sound, five distinct electrical potentials arising from different stations along the peripheral and central auditory pathway can be identified using computer averaging from scalp surface electrodes. BAERs are valuable in situations in which patients cannot or will not give reliable voluntary thresholds. They are also used to assess the integrity of the auditory nerve and brainstem in various clinical situations, including intraoperative monitoring, and in determination of brain death.

The *vestibular-evoked myogenic potential (VEMP) test* elicits a vestibulocolic reflex whose afferent limb arises from acoustically sensitive cells in the saccule, with signals conducted via the inferior vestibular nerve. VEMP is a biphasic, short-latency response recorded from the tonically contracted sternocleidomastoid muscle in response to loud auditory clicks or tones. VEMPs may be diminished or absent in patients with early and late Ménière's disease, vestibular neuritis, benign paroxysmal positional vertigo, and vestibular schwannoma. On the other hand, the threshold for VEMPs may be lower in cases of superior canal dehiscence, other inner ear dehiscence, and perilymphatic fistula.

Imaging Studies The choice of radiologic tests is largely determined by whether the goal is to evaluate the bony anatomy of the external, middle, and inner ear or to image the auditory nerve and brain. Axial and coronal CT of the temporal bone with fine 0.3- to 0.6-mm cuts is ideal for determining the caliber of the external auditory canal, integrity of the ossicular chain, and presence of middle ear or mastoid disease; it can also detect inner ear malformations. CT is also ideal for the detection of bone erosion with chronic otitis media and cholesteatoma. Pöschl reformatting in the plane of the superior semicircular canal is required for the identification of dehiscence or absence of bone over the superior semicircular canal. MRI is superior to CT for imaging of retrocochlear pathology such as vestibular schwannoma, meningioma, other lesions of the cerebellopontine angle, demyelinating lesions of the brainstem, and brain tumors. Both CT and MRI are equally capable of identifying inner ear malformations and assessing cochlear patency for preoperative evaluation of patients for cochlear implantation.

TREATMENT DISORDERS OF THE SENSE OF HEARING

In general, conductive hearing losses are amenable to surgical correction, whereas sensorineural hearing losses are usually managed medically. Atresia of the ear canal can be surgically repaired, often with significant improvement in hearing. Tympanic membrane perforations due to chronic otitis media or trauma can be repaired with an outpatient tympanoplasty. Likewise, conductive hearing loss associated with otosclerosis can be treated by stapedectomy, which is successful in >95% of cases. Tympanostomy tubes allow

the prompt return of normal hearing in individuals with middle ear effusions. Hearing aids are effective and well tolerated in patients with conductive hearing losses.

Patients with mild, moderate, and severe sensorineural hearing losses are regularly rehabilitated with hearing aids of varying configuration and strength. Hearing aids have been improved to provide greater fidelity and have been miniaturized. The current generation of hearing aids can be placed entirely within the ear canal, thus reducing any stigma associated with their use. In general, the more severe the hearing impairment, the larger the hearing aid required for auditory rehabilitation. Digital hearing aids lend themselves to individual programming, and multiple and directional microphones at the ear level may be helpful in noisy surroundings. Because all hearing aids amplify noise as well as speech, the only absolute solution to the problem of noise is to place the microphone closer to the speaker than the noise source. This arrangement is not possible with a self-contained, cosmetically acceptable device. A significant limitation of rehabilitation with a hearing aid is that although it is able to enhance detection of sound with amplification, it cannot restore clarity of hearing that is lost with presbycusis.

Patients with unilateral deafness have difficulty with sound localization and reduced clarity of hearing in background noise. They may benefit from a CROS (contralateral routing of signal) hearing aid in which a microphone is placed on the hearing-impaired side and the sound is transmitted to the receiver placed on the contralateral ear. The same result may be obtained with a bone-anchored hearing aid (BAHA), in which a hearing aid clamps to a screw integrated into the skull on the hearing-impaired side. Like the CROS hearing aid, the BAHA transfers the acoustic signal to the contralateral hearing ear, but it does so by vibrating the skull. Patients with profound deafness on one side and some hearing loss in the better ear are candidates for a BICROS hearing aid; it differs from the CROS hearing aid in that the patient wears a hearing aid, and not simply a receiver, in the better ear. Unfortunately, while CROS and BAHA devices provide benefit, they do not restore hearing in the deaf ear. Only cochlear implants can restore hearing (see below). Increasingly, cochlear implants are being investigated for the treatment of patients with single-sided deafness; early reports show great promise in not only restoring hearing but also improving sound localization and performance in background noise.

In many situations, including lectures and the theater, hearing-impaired persons benefit from assistive devices that are based on the principle of having the speaker closer to the microphone than any source of noise. Assistive devices include infrared and frequency-modulated (FM) transmission as well as an electromagnetic loop around the room for transmission to the individual's hearing aid. Hearing aids with telecoils can also be used with properly equipped telephones in the same way.

In the event that the hearing aid provides inadequate rehabilitation, cochlear implants may be appropriate (Fig. 43-4). Criteria for implantation include severe to profound hearing loss with open-set sentence cognition of $\leq 40\%$ under best aided conditions. Worldwide, more than 300,000 hearing-impaired individuals have received cochlear implants. Cochlear implants are neural prostheses that convert sound energy to electrical energy and can be used to stimulate the auditory division of the eighth nerve directly. In most cases of profound hearing impairment, the auditory hair cells are lost but the ganglionic cells of the auditory division of the eighth nerve are preserved. Cochlear implants consist of electrodes that are inserted into the cochlea through the round window, speech processors that extract acoustical elements of speech for conversion to electrical currents, and a means of transmitting the electrical energy through the skin. Patients with implants experience sound that helps with speech reading, allows open-set word recognition, and helps in modulating the person's own voice. Usually, within the first 3–6 months after implantation, adult patients can understand speech without visual cues. With the current generation of multichannel cochlear implants, nearly 75% of patients are able to converse on the telephone.

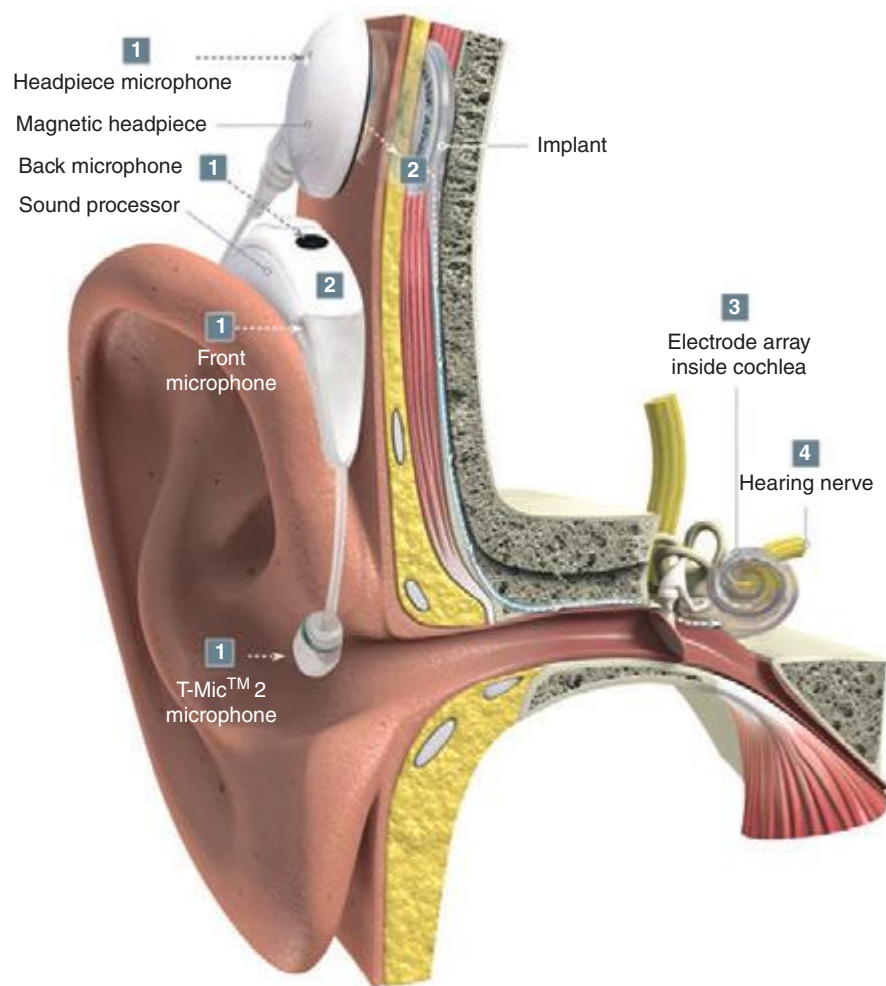


FIGURE 43-4 A cochlear implant is composed of an external microphone and speech processor worn on the ear and a receiver implanted underneath the temporalis muscle. The internal receiver is attached to an electrode that is placed surgically in the cochlea.

The U.S. Food and Drug Administration recently approved the first hybrid cochlear implant for the treatment of high-frequency hearing loss. Patients with presbycusis typically have normal low-frequency hearing, while suffering from high-frequency hearing loss associated with loss of clarity that cannot always be adequately rehabilitated with a hearing aid. However, these patients are not candidates for conventional cochlear implants because they have too much residual hearing. The hybrid implant has been specifically designed for this patient population; it has a shorter electrode than a conventional cochlear implant and can be introduced into the cochlea atraumatically, thus preserving low-frequency hearing. Individuals with a hybrid implant use their own natural low-frequency “acoustic” hearing and rely on the implant for providing “electrical” high-frequency hearing. Patients who have received the hybrid implant perform better on speech testing in both quiet and noisy backgrounds.

For individuals who have had both eighth nerves destroyed by trauma or bilateral vestibular schwannomas (e.g., neurofibromatosis type 2), brainstem auditory implants placed near the cochlear nucleus may provide auditory rehabilitation.

Tinnitus often accompanies hearing loss. As for background noise, tinnitus can degrade speech comprehension in individuals with hearing impairment. Therapy for tinnitus is usually directed toward minimizing the appreciation of tinnitus. Relief of the tinnitus may be obtained by masking it with background music. Hearing aids are also helpful in tinnitus suppression, as are tinnitus maskers, devices that present a sound to the affected ear that is more pleasant to listen to than the tinnitus. The use of a tinnitus

masker is often followed by several hours of inhibition of the tinnitus. Antidepressants have been shown to be beneficial in helping patients cope with tinnitus.

Hard-of-hearing individuals often benefit from a reduction in unnecessary noise in the environment (e.g., radio or television) to enhance the signal-to-noise ratio. Speech comprehension is aided by lip reading; therefore, the impaired listener should be seated so that the face of the speaker is well illuminated and easily seen. Although speech should be in a loud, clear voice, one should be aware that in sensorineural hearing losses in general and in hard-of-hearing elderly in particular, recruitment (abnormal perception of loud sounds) may be troublesome. Above all, optimal communication cannot take place without both parties giving it their full and undivided attention.

PREVENTION

Conductive hearing losses may be prevented by prompt antibiotic therapy of adequate duration for AOM and by ventilation of the middle ear with tympanostomy tubes in middle ear effusions lasting ≥ 12 weeks. Loss of vestibular function and deafness due to aminoglycoside antibiotics can largely be prevented by careful monitoring of serum peak and trough levels.

Some 10 million Americans have noise-induced hearing loss, and 20 million are exposed to hazardous noise in their employment. Noise-induced hearing loss can be prevented by avoidance of exposure to loud noise or by regular use of ear plugs or fluid-filled ear muffs to attenuate intense sound. [Table 43-3](#) lists loudness levels for a variety of

TABLE 43-3 DECIBEL (LOUDNESS) LEVEL OF COMMON ENVIRONMENTAL NOISE

Source	Decibel (dB)
Weakest sound heard	0
Whisper	30
Normal conversation	55–65
City traffic inside car	85
OSHA monitoring requirement begins	90
Jackhammer	95
Subway train at 200 ft	95
Power mower	107
Power saw	110
Painful sound	125
Jet engine at 100 feet	140
12-gauge shotgun blast	165
Loudest sound that can occur	194

Abbreviation: OSHA, Occupational Safety and Health Administration.

environmental sounds. High-risk activities for noise-induced hearing loss include use of electrical equipment for wood and metal working and target practice or hunting with small firearms. All internal-combustion and electric engines, including snow and leaf blowers, snowmobiles, outboard motors, and chainsaws, require protection of the user with hearing protectors. Virtually all noise-induced hearing loss is preventable through education, which should begin before the teenage years. Programs for conservation of hearing in the workplace are required by the Occupational Safety and Health Administration (OSHA) whenever the exposure over an 8-h period averages 85 dB. OSHA mandates that workers in such noisy environments have hearing monitoring and protection programs that include a preemployment screen, an annual audiologic assessment, and the mandatory use of hearing protectors. Exposure to loud sounds above 85 dB in the work environment is restricted by OSHA, with halving of allowed exposure time for each increment of 5 dB above this threshold; for example, exposure to 90 dB is permitted for 8 h; 95 dB for 4 h, and 100 dB for 2 h (Table 43-4).

TABLE 43-4 OSHA DAILY PERMISSIBLE NOISE LEVEL EXPOSURE

Sound Level (dB)	Duration per Day (h)
90	8
92	6
95	4
97	3
100	2
102	1.5
105	1
110	0.5
115	≤0.25

Note: Exposure to impulsive or impact noise should not exceed 140-dB peak sound pressure level.

Source: From https://www.osha.gov/pls/oshaweb/owadisp.show_document?p_table=standards&p_id=9735.

44 Sore Throat, Earache, and Upper Respiratory Symptoms

Michael A. Rubin, Larry C. Ford, Ralph Gonzales

Infections of the upper respiratory tract (URIs) have a tremendous impact on public health. They are among the most common reasons for visits to primary care providers, and although the illnesses are typically mild, their high incidence and transmission rates place them among the leading causes of time lost from work or school. Even though a minority (~25%) of cases are caused by bacteria, URIs are the leading diagnoses for which antibiotics are prescribed on an outpatient basis in the United States. The enormous consumption of antibiotics for these illnesses has contributed to the rise in antibiotic resistance among common community-acquired pathogens such as *Streptococcus pneumoniae*—a trend that in itself has an enormous influence on public health.

Although most URIs are caused by viruses, distinguishing patients with primary viral infection from those with primary bacterial infection is difficult. Signs and symptoms of bacterial and viral URIs are typically indistinguishable. Until consistent, inexpensive, and rapid testing becomes available and is used widely, acute infections will be diagnosed largely on clinical grounds. The judicious use and potential for misuse of antibiotics in this setting pose definite challenges.

NONSPECIFIC INFECTIONS OF THE UPPER RESPIRATORY TRACT

Nonspecific URIs are a broadly defined group of disorders that collectively constitute the leading cause of ambulatory care visits in the United States. By definition, nonspecific URIs have no prominent localizing features. They are identified by a variety of descriptive names, including *acute infective rhinitis*, *acute rhinopharyngitis/nasopharyngitis*, *acute coryza*, and *acute nasal catarrh*, as well as by the inclusive label *common cold*.

ETIOLOGY

The large assortment of URI classifications reflects the wide variety of causative infectious agents and the varied manifestations of common pathogens. Nearly all nonspecific URIs are caused by viruses spanning multiple virus families and many antigenic types. For instance, there are at least 100 immunotypes of rhinovirus (Chap. 223), the most common cause of URI (~30–40% of cases); other causes include influenza virus (three immunotypes; Chap. 224) as well as parainfluenza virus (four immunotypes), coronavirus (at least three immunotypes), and adenovirus (47 immunotypes) (Chap. 223). Respiratory syncytial virus (RSV), a well-established pathogen in pediatric populations, is also a recognized cause of significant disease in elderly and immunocompromised individuals. A host of additional viruses, including some viruses not typically associated with URIs (e.g., enteroviruses, rubella virus, and varicella-zoster virus), account for a small percentage of cases in adults each year. Although new diagnostic modalities (e.g., nasopharyngeal swab for polymerase chain reaction [PCR]) can assign a viral etiology, there are few specific treatment options, and no pathogen is identified in a substantial proportion of cases. A specific diagnostic workup beyond a clinical diagnosis is generally unnecessary in an otherwise healthy adult.

CLINICAL MANIFESTATIONS

The signs and symptoms of nonspecific URI are similar to those of other URIs but lack a pronounced localization to one particular anatomic location, such as the sinuses, pharynx, or lower airway. Nonspecific URI commonly presents as an acute, mild, and self-limited catarrhal syndrome with a median duration of ~1 week (range, 2–10 days). Signs and symptoms are diverse and frequently variable across patients, even when caused by the same virus. The principal

signs and symptoms of nonspecific URI include rhinorrhea (with or without purulence), nasal congestion, cough, and sore throat. Other manifestations, such as fever, malaise, sneezing, lymphadenopathy, and hoarseness, are more variable, with fever more common among infants and young children. This varying presentation may reflect differences in host response as well as in infecting organisms; myalgias and fatigue, for example, sometimes are seen with influenza and parainfluenza infections, whereas conjunctivitis may suggest infection with adenovirus or enterovirus. Findings on physical examination are frequently nonspecific and unimpressive. Between 0.5% and 2% of colds are complicated by secondary bacterial infections (e.g., rhinosinusitis, otitis media, and pneumonia), particularly in higher-risk populations such as infants, elderly persons, and chronically ill or immunosuppressed individuals. Secondary bacterial infections usually are associated with a prolonged course of illness, increased severity of illness, and localization of signs and symptoms, often as a rebound after initial clinical improvement (the “double-dip” sign). Purulent secretions from the nares or throat often are misinterpreted as an indication of bacterial sinusitis or pharyngitis. These secretions, however, can be seen in nonspecific URI and, in the absence of other clinical features, are poor predictors of bacterial infection.

TREATMENT NONSPECIFIC UPPER RESPIRATORY INFECTIONS

Antibiotics have no role in the treatment of uncomplicated nonspecific URI, and their misuse facilitates the emergence of antimicrobial resistance; in healthy volunteers, a single course of a commonly prescribed antibiotic like azithromycin can result in macrolide resistance in oral streptococci many months later. In the absence of clinical evidence of bacterial infection, treatment remains entirely symptom based, with use of decongestants and nonsteroidal anti-inflammatory drugs. Clinical trials of zinc, vitamin C, echinacea, and other alternative remedies have revealed no consistent benefit in the treatment of nonspecific URI.

INFECTIONS OF THE SINUS

Rhinosinusitis refers to an inflammatory condition involving the nasal sinuses. Although most cases of sinusitis involve more than one sinus, the maxillary sinus is most commonly involved; next, in order of frequency, are the ethmoid, frontal, and sphenoid sinuses. Each sinus is lined with a respiratory epithelium that produces mucus, which is transported out by ciliary action through the sinus ostium and into the nasal cavity. Normally, mucus does not accumulate in the sinuses, which remain mostly sterile despite their adjacency to the bacterium-filled nasal passages. When the sinus ostia are obstructed or when ciliary clearance is impaired or absent, the secretions can be retained, producing the typical signs and symptoms of sinusitis. As these secretions accumulate with obstruction, they become more susceptible to infection with a variety of pathogens, including viruses, bacteria, and fungi. Sinusitis affects a tremendous proportion of the population, accounts for millions of visits to primary care physicians each year, and is the fifth leading diagnosis for which antibiotics are prescribed. It typically is classified by duration of illness (acute vs. chronic); by etiology (infectious vs. noninfectious); and, when infectious, by the offending pathogen type (viral, bacterial, or fungal).

ACUTE RHINOSINUSITIS

Acute rhinosinusitis—defined as sinusitis of <4 weeks’ duration—constitutes the vast majority of sinusitis cases. Most cases are diagnosed in the ambulatory care setting and occur primarily as a consequence of a preceding viral URI. Differentiating acute bacterial from viral sinusitis on clinical grounds is difficult. Therefore, it is perhaps not surprising that antibiotics are prescribed frequently (in 85–98% of all cases) for this condition.

Etiology The ostial obstruction in rhinosinusitis can arise from both infectious and noninfectious causes. Noninfectious etiologies include

allergic rhinitis (with either mucosal edema or polyp obstruction), barotrauma (e.g., from deep-sea diving or air travel), and exposure to chemical irritants. Obstruction can also occur with nasal and sinus tumors (e.g., squamous cell carcinoma) or granulomatous diseases (e.g., granulomatosis with polyangiitis, rhinoscleroma), and conditions leading to altered mucus content (e.g., cystic fibrosis) can cause sinusitis through impaired mucus clearance. In ICUs, nasotracheal intubation and nasogastric tubes are major risk factors for nosocomial sinusitis.

Viral rhinosinusitis is far more common than bacterial sinusitis, although relatively few studies have sampled sinus aspirates for the presence of different viruses. In the studies that have done so, the viruses most commonly isolated—both alone and with bacteria—have been rhinovirus, parainfluenza virus, and influenza virus. Bacterial causes of sinusitis have been better described. Among community-acquired cases, *S. pneumoniae* and nontypable *Haemophilus influenzae* are the most common pathogens, accounting for 50–60% of cases. *Moraxella catarrhalis* causes disease in a significant percentage (20%) of children but a lesser percentage of adults. Other streptococcal species and *Staphylococcus aureus* cause only a small percentage of cases, although there is increasing concern about methicillin-resistant *S. aureus* (MRSA) as an emerging cause. It is difficult to assess whether a cultured bacterium represents a true infecting organism, an insufficiently deep sample (which would not be expected to be sterile), or—especially in the case of previous sinus surgeries—a colonizing organism. Anaerobes occasionally are found in association with infections of the roots of premolar teeth that spread to the adjacent maxillary sinuses. The role of atypical organisms like *Chlamydia pneumoniae* and *Mycoplasma pneumoniae* in the pathogenesis of acute sinusitis is unclear. Nosocomial cases commonly are associated with bacteria prevalent in the hospital environment, including *S. aureus*, *Pseudomonas aeruginosa*, *Serratia marcescens*, *Klebsiella pneumoniae*, and *Enterobacter* species. Often, these infections are polymicrobial and can involve organisms that are highly resistant to numerous antibiotics. Fungi also are established causes of sinusitis, although most acute cases are in immunocompromised patients and represent invasive, life-threatening infections. The best-known example is rhinocerebral mucormycosis caused by fungi of the order Mucorales, which includes *Rhizopus*, *Rhizomucor*, *Mucor*, *Lichtheimia* (formerly *Mycocladius*, formerly *Absidia*), and *Cunninghamella* (Chap. 242). These infections classically occur in diabetic patients with ketoacidosis but can also develop in transplant recipients, patients with hematologic malignancies, and patients receiving chronic glucocorticoid or deferoxamine therapy. Other hyaline molds, such as *Aspergillus* and *Fusarium* species, also are occasional causes of this disease.

Clinical Manifestations Most cases of acute sinusitis present after or in conjunction with a viral URI, and it can be difficult to discriminate the clinical features of one from the other, with timing becoming important in diagnosis (see below). A large proportion of patients with colds have sinus inflammation, although, as previously stated, true bacterial sinusitis complicates only 0.2–2% of these viral infections. Common presenting symptoms of sinusitis include nasal drainage and congestion, facial pain or pressure, and headache. Thick, purulent or discolored nasal discharge is often thought to indicate bacterial sinusitis but also occurs early in viral infections such as the common cold and is not specific to bacterial infection. Other nonspecific manifestations include cough, sneezing, and fever. Tooth pain, most often involving the upper molars, as well as halitosis are occasionally associated with bacterial sinusitis.

In acute sinusitis, sinus pain or pressure often localizes to the involved sinus (particularly the maxillary sinus) and can be worse when the patient bends over or is supine. Although rare, manifestations of advanced sphenoid or ethmoid sinus infection can be profound, including severe frontal or retroorbital pain radiating to the occiput, thrombosis of the cavernous sinus, and signs of orbital cellulitis. Acute focal sinusitis is uncommon but should be considered with severe symptoms involving the maxillary sinus and fever, regardless of illness duration. Similarly, patients with advanced frontal sinusitis can

present with a condition known as *Pott's puffy tumor*, with soft tissue swelling and pitting edema over the frontal bone from a communicating subperiosteal abscess. Life-threatening complications of sinusitis include meningitis, epidural abscess, and cerebral abscess.

Patients with acute fungal rhinosinusitis (such as mucormycosis; [Chap. 242](#)) often present with symptoms related to pressure effects, particularly when the infection has spread to the orbits and cavernous sinus. Signs such as orbital swelling and cellulitis, proptosis, ptosis, and decreased extraocular movement are common, as is retro- or periorbital pain. Nasopharyngeal ulcerations, epistaxis, and headaches are also common, and involvement of cranial nerves V and VII has been described in more advanced cases. Bony erosion may be evident on examination or endoscopy. Often the patient does not appear seriously ill despite the rapidly progressive nature of these infections.

Patients with acute nosocomial sinusitis are often critically ill and thus do not manifest the typical clinical features of sinus disease. This diagnosis should be suspected, however, when hospitalized patients with appropriate risk factors (e.g., nasotracheal intubation) develop fever without another apparent cause.

Diagnosis Distinguishing viral from bacterial rhinosinusitis in the ambulatory setting is usually difficult because of the relatively low sensitivity and specificity of the common clinical features. One clinical feature that has been used to help guide diagnostic and therapeutic decision-making is illness duration. Because acute bacterial sinusitis is uncommon in patients whose symptoms have lasted <10 days, expert panels now recommend reserving this diagnosis for patients with “persistent” symptoms (i.e., symptoms lasting >10 days in adults or >10–14 days in children) accompanied by the three cardinal signs of purulent nasal discharge, nasal obstruction, and facial pain ([Table 44-1](#)). Even among patients who meet these criteria, only 40–50% have true bacterial sinusitis. The use of CT or sinus radiography is not recommended for acute disease, particularly early in the course of illness (i.e., at <10 days) in light of the high prevalence of similar findings among patients with acute viral rhinosinusitis. In the evaluation of persistent, recurrent, or chronic sinusitis, CT of the sinuses becomes the radiographic study of choice.

The clinical history and/or setting often can identify cases of acute anaerobic bacterial sinusitis, acute fungal sinusitis, or sinusitis from noninfectious causes (e.g., allergic rhinosinusitis). In the case of an immunocompromised patient with acute fungal sinus infection,

immediate examination by an otolaryngologist is required. Biopsy specimens from involved areas should be examined by a pathologist for evidence of fungal hyphal elements and tissue invasion. Cases of suspected acute nosocomial sinusitis should be confirmed by sinus CT. Because therapy should target the offending organism, a sinus aspirate for culture and susceptibility testing should be obtained, whenever possible, before the initiation of antimicrobial therapy.

TREATMENT ACUTE RHINOSINUSITIS

Most patients with a clinical diagnosis of acute rhinosinusitis improve without antibiotic therapy. The preferred initial approach in patients with mild to moderate symptoms of short duration is therapy aimed at symptom relief and facilitation of sinus drainage, such as with oral and topical decongestants, nasal saline lavage, and—at least in patients with a history of chronic sinusitis or allergies—nasal glucocorticoids. Newer studies have cast doubt on the role of antibiotics and nasal glucocorticoids in acute rhinosinusitis. In one notable double-blind, randomized, placebo-controlled trial, neither antibiotics nor topical glucocorticoids had a significant impact on cure in the study population of patients, the majority of whom had had symptoms for <7 days. Similarly, another high-profile randomized trial comparing antibiotics to placebo in patients with acute rhinosinusitis demonstrated no significant improvement in symptoms by the third day of therapy. Still, antibiotic therapy can be considered for adult patients whose condition does not improve after 10 days, and patients with more severe symptoms (regardless of duration) should be treated with antibiotics ([Table 44-1](#)). However, watchful waiting remains a viable option in many cases.

Empirical antibiotic therapy for adults with community-acquired sinusitis should consist of the narrowest-spectrum agent active against the most common bacterial pathogens, including *S. pneumoniae* and *H. influenzae*—e.g., amoxicillin or amoxicillin/clavulanate (with the decision guided by local rates of β -lactamase-producing *H. influenzae*). No clinical trials support the use of broader-spectrum agents for routine cases of bacterial sinusitis, even in the current era of drug-resistant *S. pneumoniae*. For those patients who do not respond to initial antimicrobial therapy, sinus aspiration and/or lavage by an otolaryngologist should be considered. Antibiotic prophylaxis to prevent episodes of recurrent acute bacterial sinusitis is not recommended.

Surgical intervention and IV antibiotic administration usually are reserved for patients with severe disease or those with intracranial complications such as abscess and orbital involvement. Immunocompromised patients with acute invasive fungal sinusitis usually require extensive surgical debridement and treatment with IV antifungal agents active against fungal hyphal forms, such as amphotericin B. Specific therapy should be individualized according to the fungal species and its susceptibilities as well as the individual patient's characteristics.

Treatment of nosocomial sinusitis should begin with broad-spectrum antibiotics to cover common and often resistant pathogens such as *S. aureus* and gram-negative bacilli. Therapy then should be tailored to the results of culture and susceptibility testing of sinus aspirates.

CHRONIC SINUSITIS

Chronic sinusitis is characterized by symptoms of sinus inflammation lasting >12 weeks. This illness is most commonly associated with either bacteria or fungi, and clinical cure in most cases is very difficult. Many patients have undergone treatment with repeated courses of antibacterial agents and multiple sinus surgeries, increasing their risk of colonization with antibiotic-resistant pathogens and of surgical complications. These patients often have high rates of morbidity, sometimes over many years.

In *chronic bacterial sinusitis*, infection is thought to be due to the impairment of mucociliary clearance from repeated infections rather than to persistent bacterial infection. The pathogenesis of this

TABLE 44-1 GUIDELINES FOR THE DIAGNOSIS AND TREATMENT OF ACUTE SINUSITIS IN ADULTS

Diagnostic Criteria	Treatment Recommendations ^a
Moderate symptoms (e.g., nasal purulence/congestion or cough) for >10 d or	<i>Initial therapy:</i> Amoxicillin, 500 mg PO tid; or Amoxicillin/clavulanate, 500/125 mg PO tid or 875/125 mg PO bid ^b
Severe symptoms of any duration, including unilateral/focal facial swelling or tooth pain	<i>Penicillin allergy:</i> Doxycycline, 100 mg PO bid; or Clindamycin, 300 mg PO tid <i>Exposure to antibiotics within 30 d or >30% prevalence of penicillin-resistant Streptococcus pneumoniae:</i> Amoxicillin/clavulanate (extended release), 2000/125 mg PO bid; or An antipneumococcal fluoroquinolone (e.g., moxifloxacin, 400 mg PO daily) <i>Recent treatment failure:</i> Amoxicillin/clavulanate (extended release), 2000 mg PO bid; or An antipneumococcal fluoroquinolone (e.g., moxifloxacin, 400 mg PO daily)

^aThe duration of therapy is generally 7–10 days (with consideration of a 5-day course), with appropriate follow-up. Severe disease may warrant IV antibiotics and consideration of hospital admission. ^bAlthough the evidence is not as strong, amoxicillin/clavulanate may be considered for initial use, particularly if local rates of penicillin resistance or β -lactamase production are high.

condition, however, is poorly understood. Although certain conditions (e.g., cystic fibrosis) can predispose patients to chronic bacterial sinusitis, most patients with chronic rhinosinusitis do not have obvious underlying conditions that result in the obstruction of sinus drainage, the impairment of ciliary action, or immune dysfunction. Patients experience constant nasal congestion and sinus pressure, with intermittent periods of greater severity, which may persist for years. CT can be helpful in determining the extent of disease, detecting an underlying anatomic defect or obstructing process (e.g., a polyp), and assessing the response to therapy. Management should involve an otolaryngologist to conduct endoscopic examinations and obtain tissue samples for histologic examination and culture. An endoscopy-derived culture not only has a higher yield but also allows direct visualization for abnormal anatomy.

Chronic fungal sinusitis is a disease of immunocompetent hosts and is usually noninvasive, although slowly progressive invasive disease is sometimes seen. Noninvasive disease, which typically is associated with hyaline molds such as *Aspergillus* species and dematiaceous molds such as *Curvularia* or *Bipolaris* species, can present as a number of different scenarios. In mild, indolent disease, which usually occurs in the setting of repeated failures of antibacterial therapy, only nonspecific mucosal changes may be seen on sinus CT. Although there is some controversy on this point, endoscopic surgery is usually curative in these cases, with no need for antifungal therapy. Another form of disease presents as long-standing, often unilateral symptoms and opacification of a single sinus on imaging studies as a result of a mycetoma (fungus ball) within the sinus. Treatment for this condition also is surgical, although systemic antifungal therapy may be warranted in the rare case in which bony erosion occurs. A third form of disease, known as *allergic fungal sinusitis*, is seen in patients with a history of nasal polyposis and asthma, who often have had multiple sinus surgeries. Patients with this condition produce a thick, eosinophil-laden mucus with the consistency of peanut butter that contains sparse fungal hyphae on histologic examination. These patients often present with pansinusitis.

TREATMENT CHRONIC SINUSITIS

Treatment of chronic bacterial sinusitis can be challenging and consists primarily of repeated culture-guided courses of antibiotics, sometimes for 3–4 weeks or longer at a time; administration of intranasal glucocorticoids; and mechanical irrigation of the sinus with sterile saline solution. When this management approach fails, sinus surgery may be indicated and sometimes provides significant, albeit short-term, alleviation. Treatment of chronic fungal sinusitis consists of surgical removal of impacted mucus. Recurrence, unfortunately, is common.

INFECTIONS OF THE EAR AND MASTOID

Infections of the ear and associated structures can involve both the middle and the external ear, including the skin, cartilage, periosteum, ear canal, and tympanic and mastoid cavities. Both viruses and bacteria are known causes of these infections, some of which result in significant morbidity if not treated appropriately.

INFECTIONS OF EXTERNAL EAR STRUCTURES

Infections involving the structures of the external ear are often difficult to differentiate from noninfectious inflammatory conditions with similar clinical manifestations. Clinicians should consider inflammatory disorders as possible causes of external ear irritation, particularly in the absence of local or regional adenopathy. Aside from the more salient causes of inflammation, such as trauma, insect bite, and overexposure to sunlight or extreme cold, the differential diagnosis should include less common conditions such as autoimmune disorders (e.g., lupus or relapsing polychondritis) and vasculitides (e.g., granulomatosis with polyangiitis).

Auricular Cellulitis Auricular cellulitis is an infection of the skin overlying the external ear and typically follows minor local trauma. It

presents as the typical signs and symptoms of cellulitis, with tenderness, erythema, swelling, and warmth of the external ear (particularly the lobule) but without apparent involvement of the ear canal or inner structures. Treatment consists of warm compresses and oral antibiotics such as cephalexin or dicloxacillin that are active against typical skin and soft tissue pathogens (specifically, *S. aureus* and streptococci). IV antibiotics such as a first-generation cephalosporin (e.g., cefazolin) or a penicillinase-resistant penicillin (e.g., nafcillin) occasionally are needed for more severe cases, with consideration of MRSA if either risk factors or failure of therapy point to this organism.

Perichondritis Perichondritis, an infection of the perichondrium of the auricular cartilage, typically follows local trauma (e.g., piercings, burns, or lacerations). Occasionally, when the infection spreads down to the cartilage of the pinna itself, patients may develop chondritis. The infection may closely resemble auricular cellulitis, with erythema, swelling, and extreme tenderness of the pinna, although the lobule is less often involved in perichondritis. The most common pathogens are *P. aeruginosa* and *S. aureus*, although other gram-negative and gram-positive organisms occasionally are involved. Treatment consists of systemic antibiotics active against both *P. aeruginosa* and *S. aureus*. An antipseudomonal penicillin (e.g., piperacillin) or a combination of a penicillinase-resistant penicillin and an antipseudomonal quinolone (e.g., nafcillin plus ciprofloxacin) is typically used. Incision and drainage may be helpful for culture and for resolution of infection, which often takes weeks. When perichondritis fails to respond to adequate antimicrobial therapy, clinicians should consider a noninfectious inflammatory etiology such as relapsing polychondritis.

Otitis Externa The term *otitis externa* refers to a collection of diseases involving primarily the auditory meatus. Otitis externa usually results from a combination of heat and retained moisture, with desquamation and maceration of the epithelium of the outer ear canal. The disease exists in several forms: localized, diffuse, chronic, and invasive. All forms are predominantly bacterial in origin, with *P. aeruginosa* and *S. aureus* the most common pathogens.

Acute localized otitis externa (furunculosis) can develop in the outer third of the ear canal, where skin overlies cartilage and hair follicles are numerous. As in furunculosis elsewhere on the body, *S. aureus* is the usual pathogen, and treatment typically consists of an oral antistaphylococcal penicillin (e.g., dicloxacillin or cephalexin), with incision and drainage in cases of abscess formation.

Acute diffuse otitis externa is also known as *swimmer's ear*, although it can develop in patients who have not recently been swimming. Heat, humidity, and the loss of protective cerumen lead to excessive moisture and elevation of the pH in the ear canal, which in turn lead to skin maceration and irritation. Infection may then follow; the predominant pathogen is *P. aeruginosa*, although other gram-negative and gram-positive organisms—and rarely yeasts—have been recovered from patients with this condition. The illness often starts with itching and progresses to severe pain, which is usually elicited by manipulation of the pinna or tragus. The onset of pain is generally accompanied by the development of an erythematous, swollen ear canal, often with scant white, clumpy discharge. Treatment consists of cleansing the canal to remove debris and enhance the activity of topical therapeutic agents—usually hypertonic saline or mixtures of alcohol and acetic acid. Inflammation can also be decreased by adding glucocorticoids to the treatment regimen or by using Burow's solution (aluminum acetate in water). Antibiotics are most effective when given topically. Otic mixtures provide adequate pathogen coverage; these preparations usually combine neomycin with polymyxin, with or without glucocorticoids. Systemic antimicrobial agents typically are reserved for severe disease or infections in immunocompromised hosts.

Chronic otitis externa is caused primarily by repeated local irritation, most commonly arising from persistent drainage from a chronic middle-ear infection. Other causes of repeated irritation, such as insertion of cotton swabs or other foreign objects into the ear canal, can lead to this condition, as can rare chronic infections such as syphilis, tuberculosis, and leprosy. Chronic otitis externa typically presents as erythematous, scaling dermatitis in which the predominant symptom

is pruritus rather than pain; this condition must be differentiated from several others that produce a similar clinical picture, such as atopic dermatitis, seborrheic dermatitis, psoriasis, and dermatomycosis. Therapy consists of identifying and treating or removing the offending process, although successful resolution is frequently difficult.

Invasive otitis externa, also known as *malignant* or *necrotizing* otitis externa, is an aggressive and potentially life-threatening disease that occurs predominantly in elderly diabetic patients and other immunocompromised persons. The disease begins in the external canal as a soft tissue infection that progresses slowly over weeks to months and often is difficult to distinguish from a severe case of chronic otitis externa because of the presence of purulent otorrhea and an erythematous swollen ear and external canal. Severe, deep-seated otalgia, frequently out of proportion to findings on examination, is often noted and can help differentiate invasive from chronic otitis externa. The characteristic finding on examination is granulation tissue in the posteroinferior wall of the external canal, near the junction of bone and cartilage. If left unchecked, the infection can migrate to the base of the skull (resulting in skull-base osteomyelitis) and onward to the meninges and brain, with a high mortality rate. Cranial nerve involvement is seen occasionally, with the facial nerve usually affected first and most often. Thrombosis of the sigmoid sinus can occur if the infection extends to the area. CT, which can reveal osseous erosion of the temporal bone and skull base, can be used to help determine the extent of disease, as can gallium and technetium-99 scintigraphy studies. *P. aeruginosa* is by far the most common offender, although *S. aureus*, *Staphylococcus epidermidis*, *Aspergillus*, *Actinomyces*, and some gram-negative bacteria also have been associated with this disease. In all cases, the external ear canal should be cleaned and a biopsy specimen of the granulation tissue within the canal (or of deeper tissues) obtained for culture of the offending organism. IV antibiotic therapy should be given for a prolonged course (6–8 weeks) and directed specifically toward the recovered pathogen. For *P. aeruginosa*, the regimen typically includes an antipseudomonal penicillin or cephalosporin (e.g., piperacillin or cefepime), often with an aminoglycoside or a fluoroquinolone, the latter of which can even be administered orally given its excellent

bioavailability. In addition, antibiotic drops containing an agent active against *Pseudomonas* (e.g., ciprofloxacin) are usually prescribed and are combined with glucocorticoids to reduce inflammation. Cases of invasive *Pseudomonas* otitis externa recognized in the early stages can sometimes be treated with oral and otic fluoroquinolones alone, albeit with close follow-up. Extensive surgical debridement, once an important component of the treatment approach, is now rarely indicated.

In *necrotizing otitis externa*, recurrence is documented up to 20% of the time. Aggressive glycemic control in diabetics is important not only for effective treatment but also for prevention of recurrence. The role of hyperbaric oxygen has not been clearly established.

INFECTIONS OF MIDDLE-EAR STRUCTURES

Otitis media is an inflammatory condition of the middle ear that results from dysfunction of the eustachian tube in association with a number of illnesses, including URIs and chronic rhinosinusitis. The inflammatory response in these conditions leads to the development of a sterile transudate within the middle ear and mastoid cavities. Infection may occur if bacteria or viruses from the nasopharynx contaminate this fluid, producing an acute (or sometimes chronic) illness.

Acute Otitis Media Acute otitis media results when pathogens from the nasopharynx are introduced into the inflammatory fluid collected in the middle ear (e.g., by nose blowing during a URI). Pathogenic proliferation in this space leads to the development of the typical signs and symptoms of acute middle-ear infection. The diagnosis of acute otitis media requires the demonstration of fluid in the middle ear (with tympanic membrane [TM] immobility) and the accompanying signs or symptoms of local or systemic illness (Table 44-2).

ETIOLOGY Acute otitis media typically follows a viral URI. The causative viruses (most commonly RSV, influenza virus, rhinovirus, and enterovirus) can themselves cause subsequent acute otitis media; more often, they predispose the patient to bacterial otitis media. Studies using tympanocentesis have consistently found *S. pneumoniae* to be the most important bacterial cause, isolated in up to 35% of cases. *H. influenzae* (nontypable strains) and *M. catarrhalis* also are

TABLE 44-2 GUIDELINES FOR THE DIAGNOSIS AND TREATMENT OF ACUTE OTITIS MEDIA

Illness Severity	Diagnostic Criteria	Treatment Recommendations
Mild to moderate	>2 yrs or 6 mo to 2 yrs without middle-ear effusion	<i>Observation alone</i> (deferring antibiotic therapy for 48–72 h and limiting management to symptom relief)
	<6 mo; or 6 mo to 2 yrs with middle-ear effusion (fluid in the middle ear, evidenced by decreased TM mobility, air/fluid level behind TM, bulging TM, purulent otorrhea) and acute onset of signs and symptoms of middle-ear inflammation, including fever, otalgia, decreased hearing, tinnitus, vertigo, erythematous TM; or	<i>Initial therapy</i> ^a Amoxicillin, 80–90 mg/kg qd (up to 2 g) PO in divided doses (bid or tid); or Cefdinir, 14 mg/kg qd PO in 1 dose or divided doses (bid); or Cefuroxime, 30 mg/kg qd PO in divided doses (bid); or Azithromycin, 10 mg/kg qd PO on day 1 followed by 5 mg/kg qd PO for 4 d
	>2 yrs with bilateral disease, TM perforation, high fever, immunocompromise, emesis	<i>Exposure to antibiotics within 30 d or recent treatment failure</i> ^{a,b} : Amoxicillin, 90 mg/kg qd (up to 2 g) PO in divided doses (bid), plus clavulanate, 6.4 mg/kg qd PO in divided doses (bid); or Ceftriaxone, 50 mg/kg IV/IM qd for 3 d; or Clindamycin, 30–40 mg/kg qd PO in divided doses (tid)
Severe	As above, with temperature $\geq 39.0^{\circ}\text{C}$ (102°F); or Moderate to severe otalgia	<i>Initial therapy</i> ^a Amoxicillin, 90 mg/kg qd (up to 2 g) PO in divided doses (bid), plus clavulanate, 6.4 mg/kg qd PO in divided doses (bid); or Ceftriaxone, 50 mg/kg IV/IM qd for 3 d <i>Exposure to antibiotics within 30 d or recent treatment failure</i> ^{a,b} Ceftriaxone, 50 mg/kg IV/IM qd for 3 d; or Clindamycin, 30–40 mg/kg qd PO in divided doses (tid); or Consider tympanocentesis with culture

^aDuration (unless otherwise specified): 10 days for patients <6 years old and patients with severe disease; 5–7 days (with consideration of observation only in previously healthy individuals with mild disease) for patients ≥ 6 years old. ^bFailure to improve and/or clinical worsening after 48–72 h of observation or treatment.

Abbreviation: TM, tympanic membrane.

Source: American Academy of Pediatrics Subcommittee on Management of Acute Otitis Media, 2004.

common bacterial causes of acute otitis media, and concern is increasing with MRSA as an emerging etiologic agent. Viruses, such as those mentioned above, have been recovered either alone or with bacteria in 17–40% of cases.

CLINICAL MANIFESTATIONS Fluid in the middle ear is typically demonstrated or confirmed with pneumatic otoscopy. In the absence of fluid, the TM moves visibly with the application of positive and negative pressure, but this movement is dampened when fluid is present. With bacterial infection, the TM can also be erythematous, bulging, or retracted and occasionally can perforate spontaneously. The signs and symptoms accompanying infection can be local or systemic, including otalgia, otorrhea, diminished hearing, and fever. Erythema of the TM is often evident but is nonspecific as it frequently is seen in association with inflammation of the upper respiratory mucosa. Other signs and symptoms occasionally reported include vertigo, nystagmus, and tinnitus.

TREATMENT ACUTE OTITIS MEDIA

There has been considerable debate on the usefulness of antibiotics for the treatment of acute otitis media. A higher proportion of treated than untreated patients are free of illness 3–5 days after diagnosis. The difficulty of predicting which patients will benefit from antibiotic therapy has led to different approaches. In the Netherlands, for instance, physicians typically manage acute otitis media with initial observation, administering anti-inflammatory agents for aggressive pain management and reserving antibiotics for high-risk patients, patients with complicated disease, or patients whose condition does not improve after 48–72 h. In contrast, many experts in the United States continue to recommend antibiotic therapy for children <6 months old in light of the higher frequency of secondary complications in this young and functionally immunocompromised population. However, observation without antimicrobial therapy is now the recommended option in the United States for acute otitis media in children >2 years of age and for mild to moderate disease without middle-ear effusion in children 6 months to 2 years of age. Treatment is typically indicated for patients <6 months old; for children 6 months to 2 years old who have middle-ear effusion and signs/symptoms of middle-ear inflammation; for all patients >2 years old who have bilateral disease, TM perforation, immunocompromise, or emesis; and for any patient who has severe symptoms, including a fever $\geq 39^{\circ}\text{C}$ or moderate to severe otalgia (Table 44-2).

Because most studies of the etiologic agents of acute otitis media consistently document similar pathogen profiles, therapy is generally empirical except in those few cases in which tympanocentesis is warranted—e.g., cases refractory to therapy and cases in patients who are severely ill or immunodeficient. Despite resistance to penicillin and amoxicillin in roughly one-quarter of *S. pneumoniae* isolates, one-third of *H. influenzae* isolates, and nearly all *M. catarrhalis* isolates, outcome studies continue to find that amoxicillin is as successful as any other agent, and it remains the drug of first choice in recommendations from multiple sources (Table 44-2). Therapy for uncomplicated acute otitis media typically is administered for 5–7 days to patients ≥ 6 years old; longer courses (e.g., 10 days) should be reserved for patients with severe disease, in whom short-course therapy may be inadequate.

A switch in regimen is recommended if there is no clinical improvement by the third day of therapy, given the possibility of infection with a β -lactamase-producing strain of *H. influenzae* or *M. catarrhalis* or with a strain of penicillin-resistant *S. pneumoniae*. Decongestants and antihistamines are frequently used as adjunctive agents to reduce congestion and relieve obstruction of the eustachian tube, but clinical trials have yielded no significant evidence of benefit with either class of agents.


Recurrent Acute Otitis Media Recurrent acute otitis media (more than three episodes within 6 months or four episodes within 12 months)

generally is due to relapse or reinfection, although data indicate that the majority of early recurrences are new infections. In general, the same pathogens responsible for acute otitis media cause recurrent disease; even so, the recommended treatment consists of antibiotics active against β -lactamase-producing organisms. Antibiotic prophylaxis (e.g., with trimethoprim-sulfamethoxazole [TMP-SMX] or amoxicillin) can reduce recurrences in patients with recurrent acute otitis media by an average of one episode per year, but this benefit is small compared with the high likelihood of colonization with antibiotic-resistant pathogens. Other approaches, including placement of tympanostomy tubes, adenoidectomy, and tonsillectomy plus adenoidectomy, are of questionable overall value in light of the relatively small benefit compared with the potential for complications.

Serous Otitis Media In serous otitis media (otitis media with effusion), fluid is present in the middle ear for an extended period in the absence of signs and symptoms of infection. In general, acute effusions are self-limited; most resolve in 2–4 weeks. In some cases, however (in particular after an episode of acute otitis media), effusions can persist for months. These chronic effusions are often associated with significant hearing loss in the affected ear. The great majority of cases of otitis media with effusion resolve spontaneously within 3 months without antibiotic therapy. Antibiotic therapy or myringotomy with insertion of tympanostomy tubes typically is reserved for patients in whom bilateral effusion (1) has persisted for at least 3 months and (2) is associated with significant bilateral hearing loss. With this conservative approach and the application of strict diagnostic criteria for acute otitis media and otitis media with effusion, it is estimated that 6–8 million courses of antibiotics could be avoided each year in the United States.

Chronic Otitis Media Chronic suppurative otitis media is characterized by persistent or recurrent purulent otorrhea in the setting of TM perforation. Usually, there is also some degree of conductive hearing loss. This condition can be categorized as active or inactive. Inactive disease is characterized by a central perforation of the TM, which allows drainage of purulent fluid from the middle ear. When the perforation is more peripheral, squamous epithelium from the auditory canal may invade the middle ear through the perforation, forming a mass of keratinaceous debris (*cholesteatoma*) at the site of invasion. This mass can enlarge and has the potential to erode bone and promote further infection, which can lead to meningitis, brain abscess, or paralysis of cranial nerve VII. Treatment of chronic active otitis media is surgical; mastoidectomy, myringoplasty, and tympanoplasty can be performed as outpatient surgical procedures, with an overall success rate of ~80%. Chronic inactive otitis media is more difficult to cure, usually requiring repeated courses of topical antibiotic drops during periods of drainage. Systemic antibiotics may offer better cure rates, but their role in the treatment of this condition remains unclear.

Mastoiditis Acute mastoiditis was relatively common among children before the introduction of antibiotics. Because the mastoid air cells connect with the middle ear, the process of fluid collection and infection is usually the same in the mastoid as in the middle ear. Early and frequent treatment of acute otitis media is most likely the reason that the incidence of acute mastoiditis has declined to only 1.2–2.0 cases per 100,000 person-years in countries with high prescribing rates for acute otitis media.

 In countries such as the Netherlands, where antibiotics are used sparingly for acute otitis media, the incidence rate of acute mastoiditis is roughly twice that in countries like the United States. However, neighboring Denmark has a rate of acute mastoiditis similar to that in the Netherlands but an antibiotic-prescribing rate for acute otitis media more similar to that in the United States.

In typical acute mastoiditis, purulent exudate collects in the mastoid air cells (Fig. 44-1), producing pressure that may result in erosion of the surrounding bone and formation of abscess-like cavities that are



FIGURE 44-1 Acute mastoiditis. Axial CT image shows an acute fluid collection within the mastoid air cells on the left.

usually evident on CT. Patients typically present with pain, erythema, and swelling of the mastoid process along with displacement of the pinna, usually in conjunction with the typical signs and symptoms of acute middle-ear infection. Rarely, patients can develop severe complications if the infection tracks under the periosteum of the temporal bone to cause a subperiosteal abscess, erodes through the mastoid tip to cause a deep neck abscess, or extends posteriorly to cause septic thrombosis of the lateral sinus.

Purulent fluid should be cultured whenever possible to help guide antimicrobial therapy. Initial empirical therapy usually is directed against the typical organisms associated with acute otitis media, such as *S. pneumoniae*, *H. influenzae*, and *M. catarrhalis*. Patients with more severe or prolonged courses of illness should be treated for infection with *S. aureus* and gram-negative bacilli (including *Pseudomonas*). Broad empirical therapy should be narrowed once culture results become available. Most patients can be treated conservatively with IV antibiotics; surgery (cortical mastoidectomy) is reserved for complicated cases and those in which conservative treatment has failed.

INFECTIONS OF THE PHARYNX AND ORAL CAVITY

Oropharyngeal infections range from mild, self-limited viral illnesses to serious, life-threatening bacterial infections. The most common presenting symptom is sore throat—one of the most common reasons for ambulatory care visits by both adults and children. Although sore throat is a symptom in many noninfectious illnesses as well, the overwhelming majority of patients with a new sore throat have acute pharyngitis of viral or bacterial etiology.

ACUTE PHARYNGITIS

Millions of visits to primary care providers each year are for sore throat; the majority of cases of acute pharyngitis are caused by typical respiratory viruses. The most important source of concern is infection with group A β -hemolytic *Streptococcus* (*S. pyogenes*) that is associated with acute glomerulonephritis and acute rheumatic fever. The risk of rheumatic fever can be reduced by timely penicillin therapy.

Etiology A wide variety of organisms cause acute pharyngitis. The relative importance of the different pathogens can only be estimated, since a significant proportion of cases (~30%) have no identified cause.

Together, respiratory viruses are the most common identifiable cause of acute pharyngitis, with rhinoviruses and coronaviruses accounting for large proportions of cases (~20% and at least 5%, respectively). Influenza virus, parainfluenza virus, and adenovirus also account for a measurable share of cases, with the former two more seasonal and the latter as part of the more clinically severe syndrome of pharyngoconjunctival fever. Other important but less common viral causes include herpes simplex virus (HSV) types 1 and 2, coxsackievirus A, cytomegalovirus (CMV), and Epstein-Barr virus (EBV). Acute HIV infection can present as acute pharyngitis and should be considered in at-risk populations.

Acute bacterial pharyngitis is typically caused by *S. pyogenes*, which accounts for ~5–15% of all cases of acute pharyngitis in adults; rates vary with the season and with utilization of the health care system. Group A streptococcal pharyngitis is primarily a disease of children 5–15 years of age; it is uncommon among children <3 years old, as is rheumatic fever. Streptococci of groups C and G account for a minority of cases, although these serogroups are nonrheumatogenic. *Fusobacterium necrophorum* has been increasingly recognized as a cause of pharyngitis in adolescents and young adults and is isolated nearly as often as group A streptococci. This organism is important because of the rare but life-threatening *Lemierre's disease*, which is generally associated with *F. necrophorum* and is usually preceded by pharyngitis (see “Oral Infections,” below). The remaining bacterial causes of acute pharyngitis are seen infrequently (<1% of cases each) but should be considered in appropriate exposure groups because of the severity of illness if left untreated; these etiologic agents include *Neisseria gonorrhoeae*, *Corynebacterium diphtheriae*, *Corynebacterium ulcerans*, *Yersinia enterocolitica*, and *Treponema pallidum* (in secondary syphilis). Anaerobic bacteria also can cause acute pharyngitis (*Vincent's angina*) and can contribute to more serious polymicrobial infections, such as peritonsillar or retropharyngeal abscesses (see below). Atypical organisms such as *M. pneumoniae* and *C. pneumoniae* have been recovered from patients with acute pharyngitis; whether these agents are commensals or causes of acute infection is debatable.

Clinical Manifestations Although the signs and symptoms accompanying acute pharyngitis are not reliable predictors of the etiologic agent, the clinical presentation occasionally suggests one etiology over another. Acute pharyngitis due to respiratory viruses such as rhinovirus or coronavirus usually is not severe and typically is associated with a constellation of coryzal symptoms better characterized as nonspecific URI. Findings on physical examination are uncommon; fever is rare, and tender cervical adenopathy and pharyngeal exudates are not seen. In contrast, acute pharyngitis from influenza virus can be severe and is much more likely to be associated with fever as well as with myalgias, headache, and cough. The presentation of pharyngoconjunctival fever due to adenovirus infection is similar. Since pharyngeal exudate may be present on examination, this condition can be difficult to differentiate from streptococcal pharyngitis. However, adenoviral pharyngitis is distinguished by the presence of conjunctivitis in one-third to one-half of patients. Acute pharyngitis from primary HSV infection can also mimic streptococcal pharyngitis in some cases, with pharyngeal inflammation and exudate, but the presence of vesicles and shallow ulcers on the palate can help differentiate the two diseases. This HSV syndrome is distinct from pharyngitis caused by coxsackievirus (*herpangina*), which is associated with small vesicles that develop on the soft palate and uvula and then rupture to form shallow white ulcers. Acute exudative pharyngitis coupled with fever, fatigue, generalized lymphadenopathy, and (on occasion) splenomegaly is characteristic of infectious mononucleosis due to EBV or CMV. Acute primary infection with HIV is frequently associated with fever and acute pharyngitis as well as with myalgias, arthralgias, malaise, and occasionally a nonpruritic maculopapular rash, which may be followed by lymphadenopathy and mucosal ulcerations without exudate.

The clinical features of acute pharyngitis caused by streptococci of groups A, C, and G are similar, ranging from a relatively mild illness without many accompanying symptoms to clinically severe cases with profound pharyngeal pain, fever, chills, and abdominal pain.

A hyperemic pharyngeal membrane with tonsillar hypertrophy and exudate is usually seen, along with tender anterior cervical adenopathy. Coryzal manifestations, including cough, are typically absent; when present, they suggest a viral etiology. Strains of *S. pyogenes* that generate erythrogenic toxin can also produce scarlet fever characterized by an erythematous rash and strawberry tongue. The other types of acute bacterial pharyngitis (e.g., gonococcal, diphtherial, and yersinial) often present as exudative pharyngitis with or without other clinical features. Their etiologies are often suggested only by the clinical history.

Diagnosis The primary goal of diagnostic testing is to separate acute streptococcal pharyngitis from pharyngitis of other etiologies

(particularly viral) so that antibiotics can be prescribed more efficiently for patients in whom they may be beneficial. The most appropriate standard for the diagnosis of streptococcal pharyngitis, however, has not been established definitively. Throat swab culture is generally regarded as the most appropriate but cannot distinguish between infection and colonization and requires 24–48 h to yield results that vary with technique and culture conditions. Rapid antigen-detection tests offer good specificity (>90%) but lower sensitivity when implemented in routine practice. The sensitivity has also been shown to vary across the clinical spectrum of disease (65–90%). Several clinical prediction systems (Fig. 44-2) can increase the sensitivity of rapid antigen-detection tests to >90% in controlled settings. Since the

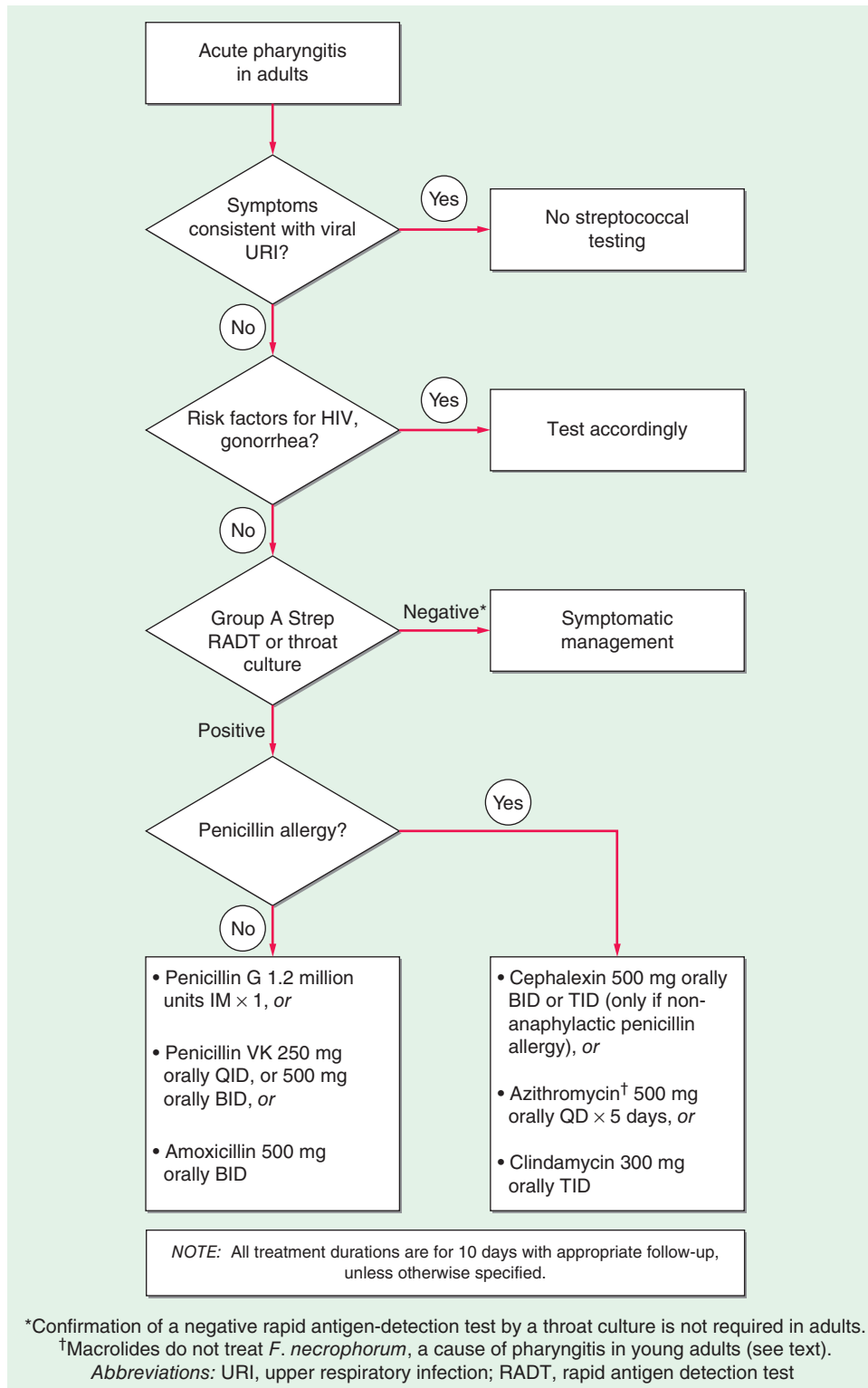


FIGURE 44-2 Algorithm for the diagnosis and treatment of acute pharyngitis.

sensitivities achieved in routine clinical practice are often lower, several medical and professional societies continue to recommend that all negative rapid antigen-detection tests in children be confirmed by a throat culture to limit transmission and complications of illness caused by group A streptococci. The Centers for Disease Control and Prevention, the Infectious Diseases Society of America, and the American Academy of Family Physicians do not recommend backup culture when adults have negative results from a highly sensitive rapid antigen-detection test, however, because of the lower prevalence and smaller benefit in this age group.

Cultures and rapid diagnostic tests for other causes of acute pharyngitis, such as influenza virus, adenovirus, HSV, EBV, CMV, and *M. pneumoniae*, are available in many locations and can be used when suspected. The diagnosis of acute EBV infection depends primarily on the detection of antibodies to the virus with a heterophile agglutination assay (monospot slide test) or enzyme-linked immunosorbent assay. Testing for HIV RNA or antigen (p24) should be performed when acute primary HIV infection is suspected. If other bacterial causes are suspected (particularly *N. gonorrhoeae*, *C. diphtheriae*, or *Y. enterocolitica*), specific cultures should be requested since these organisms may be missed on routine throat swab culture.

TREATMENT PHARYNGITIS

Antibiotic treatment of pharyngitis due to *S. pyogenes* confers numerous benefits, including a decrease in the risk of rheumatic fever, the primary focus of treatment. The magnitude of this benefit is fairly small, since rheumatic fever is now a rare disease, even among untreated patients. Nevertheless, when therapy is started within 48 h of illness onset, symptom duration is decreased modestly. An additional benefit of therapy is the potential to reduce the transmission of streptococcal pharyngitis, particularly in areas of overcrowding or close contact. Antibiotic therapy for acute pharyngitis is therefore recommended in cases in which *S. pyogenes* is confirmed as the etiologic agent by rapid antigen-detection test or throat swab culture. Otherwise, antibiotics should be given in routine cases only when another bacterial cause has been identified. Effective therapy for streptococcal pharyngitis consists of either a single dose of IM benzathine penicillin or a full 10-day course of oral penicillin (Fig. 44-2).



Azithromycin can be used in place of penicillin, although resistance to azithromycin among *S. pyogenes* strains in some parts of the world (particularly Europe) can prohibit the use of this drug. Newer (and more expensive) antibiotics also are active against streptococci but offer no greater efficacy than the agents mentioned above. Testing for cure is unnecessary and may reveal only chronic colonization. There is no evidence to support antibiotic treatment of group C or G streptococcal pharyngitis or pharyngitis in which mycoplasmas or chlamydiae have been recovered. Cultures can be of benefit because *F. necrophorum*, an increasingly common cause of bacterial pharyngitis in young adults, is not covered by macrolide therapy. Long-term penicillin prophylaxis (benzathine penicillin G, 1.2 million units IM every 3–4 weeks; or penicillin VK, 250 mg PO bid) is indicated for patients at risk of recurrent rheumatic fever.

Treatment of viral pharyngitis is entirely symptom based except in infection with influenza virus or HSV. For influenza, the armamentarium includes the adamantanes amantadine and rimantadine and the neuraminidase inhibitors oseltamivir and zanamivir. Administration of all these agents needs to be started within 48 h of symptom onset to reduce illness duration meaningfully. Among these agents, only oseltamivir and zanamivir are active against both influenza A and influenza B and therefore can be used when local patterns of infection and antiviral resistance are unknown. Oropharyngeal HSV infection sometimes responds to treatment with antiviral agents such as acyclovir, although these drugs are often reserved for immunosuppressed patients.

Complications Although rheumatic fever is the best-known complication of acute streptococcal pharyngitis, the risk of its following acute infection remains quite low. Other complications include acute glomerulonephritis and numerous suppurative conditions, such as peritonsillar abscess (*quinsy*), otitis media, mastoiditis, sinusitis, bacteremia, and pneumonia—all of which occur at low rates. Although antibiotic treatment of acute streptococcal pharyngitis can prevent the development of rheumatic fever, there is no evidence that it can prevent acute glomerulonephritis. Some evidence supports antibiotic use to prevent the suppurative complications of streptococcal pharyngitis, particularly peritonsillar abscess, which can also involve oral anaerobes such as *Fusobacterium*. Abscesses usually are accompanied by severe pharyngeal pain, dysphagia, fever, and dehydration; in addition, medial displacement of the tonsil and lateral displacement of the uvula are often evident on examination. Although early use of IV antibiotics (e.g., clindamycin, penicillin G with metronidazole) may obviate the need for surgical drainage in some cases, treatment typically involves needle aspiration or incision and drainage.

ORAL INFECTIONS

Aside from periodontal disease such as gingivitis, infections of the oral cavity most commonly involve HSV or *Candida* species. In addition to causing painful cold sores on the lips, HSV can infect the tongue and buccal mucosa, causing the formation of irritating vesicles. Although topical antiviral agents (e.g., acyclovir and penciclovir) can be used externally for cold sores, oral or IV acyclovir is often needed for primary infections, extensive oral infections, and infections in immunocompromised patients. Oropharyngeal candidiasis (*thrush*) is caused by a variety of *Candida* species, most often *C. albicans*. Thrush occurs predominantly in neonates, immunocompromised patients (especially those with AIDS), and recipients of prolonged antibiotic or glucocorticoid therapy. In addition to sore throat, patients often report a burning tongue, and physical examination reveals friable white or gray plaques on the gingiva, tongue, and oral mucosa. Treatment, which usually consists of an oral antifungal suspension (nystatin or clotrimazole) or oral fluconazole, is typically successful. In the uncommon cases of fluconazole-refractory thrush that are seen in some patients with HIV/AIDS, other therapeutic options include oral formulations of itraconazole or voriconazole as well as an IV echinocandin (caspofungin, micafungin, or anidulafungin) or amphotericin B deoxycholate, if needed. In these cases, therapy based on culture and susceptibility test results is ideal.

Vincent's angina, also known as *acute necrotizing ulcerative gingivitis* or *trench mouth*, is a unique and dramatic form of gingivitis characterized by painful, inflamed gingiva with ulcerations of the interdental papillae that bleed easily. Since oral anaerobes are the cause, patients typically have halitosis and frequently present with fever, malaise, and lymphadenopathy. Treatment consists of debridement and oral administration of penicillin plus metronidazole, with clindamycin or doxycycline alone as an alternative.

Ludwig's angina is a rapidly progressive, potentially fulminant form of cellulitis that involves the bilateral sublingual and submandibular spaces and that typically originates from an infected or recently extracted tooth, most commonly the lower second and third molars. Improved dental care has reduced the incidence of this disorder substantially. Infection in these areas leads to dysphagia, odynophagia, and “woody” edema in the sublingual region, forcing the tongue up and back with the potential for airway obstruction. Fever, dysarthria, and drooling also may be noted, and patients may speak in a “hot potato” voice. Intubation or tracheostomy may be necessary to secure the airway, as asphyxiation is the most common cause of death. Patients should be monitored closely and treated promptly with IV antibiotics directed against streptococci and oral anaerobes. Recommended agents include ampicillin/sulbactam, clindamycin, or high-dose penicillin plus metronidazole.

Postanginal septicemia (*Lemierre's disease*) is a rare anaerobic oropharyngeal infection caused predominantly by *F. necrophorum*. The illness typically starts as a sore throat (most commonly in adolescents and young adults), which may present as exudative tonsillitis or peritonsillar abscess. Infection of the deep pharyngeal tissue allows


organisms to drain into the lateral pharyngeal space, which contains the carotid artery and internal jugular vein. Septic thrombophlebitis of the internal jugular vein can result, with associated pain, dysphagia, and neck swelling and stiffness. Sepsis usually occurs 3–10 days after the onset of sore throat and is often coupled with metastatic infection to the lung and other distant sites. Occasionally, the infection can extend along the carotid sheath and into the posterior mediastinum, resulting in mediastinitis, or it can erode into the carotid artery, with the early sign of repeated small bleeds into the mouth. The mortality rate from these invasive infections can be as high as 50%. Treatment consists of IV antibiotics (clindamycin or ampicillin/sulbactam) and surgical drainage of any purulent collections. The concomitant use of anticoagulants to prevent embolization remains controversial and is sometimes advised, with careful consideration of both the risks and the benefits.

INFECTIONS OF THE LARYNX AND EPIGLOTTITIS

LARYNGITIS

Laryngitis is defined as any inflammatory process involving the larynx and can be caused by a variety of infectious and noninfectious processes. The vast majority of laryngitis cases seen in clinical practice in developed countries are acute. Acute laryngitis is a common syndrome caused predominantly by the same viruses responsible for many other URIs. In fact, most cases of acute laryngitis occur in the setting of a viral URI.

Etiology Nearly all major respiratory viruses have been implicated in acute viral laryngitis, including rhinovirus, influenza virus, parainfluenza virus, adenovirus, coxsackievirus, coronavirus, and RSV. Acute laryngitis can also be associated with acute bacterial respiratory infections such as those caused by group A *Streptococcus* or *C. diphtheriae* (although diphtheria has been virtually eliminated in the United States). Another bacterial pathogen thought to play a role (albeit unclear) in the pathogenesis of acute laryngitis is *M. catarrhalis*, which has been recovered from nasopharyngeal culture in a significant percentage of cases.

 Chronic laryngitis of infectious etiology is much less common in developed than in developing countries. Laryngitis due to *Mycobacterium tuberculosis* is often difficult to distinguish from laryngeal cancer, in part because of the frequent absence of signs, symptoms, and radiographic findings typical of pulmonary disease. *Histoplasma* and *Blastomyces* may cause laryngitis, often as a complication of systemic infection. *Candida* species can cause laryngitis as well, often in association with thrush or esophagitis and particularly in immunosuppressed patients. Rare cases of chronic laryngitis are due to *Coccidioides* and *Cryptococcus*.

Clinical Manifestations Laryngitis is characterized by hoarseness and also can be associated with reduced vocal pitch or aphonia. As acute laryngitis is caused predominantly by respiratory viruses, these symptoms usually occur in association with other symptoms and signs of URI, including rhinorrhea, nasal congestion, cough, and sore throat. Direct laryngoscopy often reveals diffuse laryngeal erythema and edema, along with vascular engorgement of the vocal folds. In addition, chronic disease (e.g., tuberculous laryngitis) often includes mucosal nodules and ulcerations visible on laryngoscopy; these lesions are sometimes mistaken for laryngeal cancer.

TREATMENT LARYNGITIS

Acute laryngitis is usually treated with humidification and voice rest alone. Antibiotics are not recommended except when group A *Streptococcus* is cultured, in which case penicillin is the drug of choice. The choice of therapy for chronic laryngitis depends on the pathogen, whose identification usually requires biopsy with culture. Patients with laryngeal tuberculosis are highly contagious because of the large number of organisms that are easily aerosolized. These patients should be managed in the same way as patients with active pulmonary disease.

CROUP

The term *croup* actually denotes a group of diseases collectively referred to as “croup syndrome,” all of which are acute and predominantly viral respiratory illnesses characterized by marked swelling of the subglottic region of the larynx. Croup primarily affects children <6 years old. For a detailed discussion of this entity, the reader should consult a textbook of pediatric medicine.

EPIGLOTTITIS

Acute epiglottitis (supraglottitis) is an acute, rapidly progressive form of cellulitis of the epiglottis and adjacent structures that can result in complete—and potentially fatal—airway obstruction in both children and adults. Before the widespread use of *H. influenzae* type b (Hib) vaccine, this entity was much more common among children, with a peak incidence at ~3.5 years of age. In some countries, mass vaccination against Hib has reduced the annual incidence of acute epiglottitis in children by >90%; in contrast, the annual incidence in adults has changed little since the introduction of Hib vaccine. Because of the danger of airway obstruction, acute epiglottitis constitutes a medical emergency, particularly in children, and prompt diagnosis and airway protection are of the utmost importance.

Etiology After the introduction of the Hib vaccine in the mid-1980s, disease incidence among children in the United States declined dramatically. Nevertheless, lack of vaccination or vaccine failure has meant that many pediatric cases seen today are still due to Hib. In adults and (more recently) in children, a variety of other bacterial pathogens have been associated with epiglottitis, the most common being group A *Streptococcus*. Other pathogens—seen less frequently—include *S. pneumoniae*, *Haemophilus parainfluenzae*, and *S. aureus* (including MRSA). Viruses have not been established as causes of acute epiglottitis.

Clinical Manifestations and Diagnosis Epiglottitis typically presents more acutely in young children than in adolescents or adults. On presentation, most children have had symptoms for <24 h, including high fever, severe sore throat, tachycardia, systemic toxicity, and (in many cases) drooling while sitting forward. Symptoms and signs of respiratory obstruction also may be present and may progress rapidly. The somewhat milder illness in adolescents and adults often follows 1–2 days of severe sore throat and is commonly accompanied by dyspnea, drooling, and stridor. Physical examination of patients with acute epiglottitis may reveal moderate or severe respiratory distress, with inspiratory stridor and retractions of the chest wall. These findings *diminish* as the disease progresses and the patient tires. Conversely, oropharyngeal examination reveals infection that is much less severe than would be predicted from the symptoms—a finding that should alert the clinician to a cause of symptoms and obstruction that lies beyond the tonsils. The diagnosis often is made on clinical grounds, although direct fiberoptic laryngoscopy is frequently performed in a controlled environment (e.g., an operating room) to visualize and culture the typical edematous “cherry-red” epiglottis and facilitate placement of an endotracheal tube. Direct visualization in an examination room (i.e., with a tongue blade and indirect laryngoscopy) is not recommended because of the risk of immediate laryngospasm and complete airway obstruction. Lateral neck radiographs and laboratory tests can assist in the diagnosis but may delay the critical securing of the airway and cause the patient to be moved or repositioned more than is necessary, thereby increasing the risk of further airway compromise. Neck radiographs typically reveal an enlarged edematous epiglottis (the “thumbprint sign,” Fig. 44-3), usually with a dilated hypopharynx and normal subglottic structures. Laboratory tests characteristically document mild to moderate leukocytosis with a predominance of neutrophils. Blood cultures are positive in a significant proportion of cases.

TREATMENT EPIGLOTTITIS

Security of the airway is always of primary concern in acute epiglottitis, even if the diagnosis is only suspected. Mere observation for signs of impending airway obstruction is not routinely recommended,



FIGURE 44-3 Acute epiglottitis. In this lateral soft tissue radiograph of the neck, the arrow indicates the enlarged edematous epiglottis (the “thumbprint sign”).

particularly in children. Many adults have been managed with observation only since the illness is perceived to be milder in this age group, but some data suggest that this approach may be risky and probably should be reserved only for adult patients who have yet to develop dyspnea or stridor. Once the airway has been secured and specimens of blood and epiglottis tissue have been obtained for culture, treatment with IV antibiotics should be given to cover the most likely organisms, particularly *H. influenzae*. Because rates of ampicillin resistance in this organism have risen significantly in recent years, therapy with a β -lactam/ β -lactamase inhibitor combination or a second- or third-generation cephalosporin is recommended. Typically, ampicillin/sulbactam, cefuroxime, cefotaxime, or ceftriaxone is given, with clindamycin and TMP-SMX reserved for patients allergic to β -lactams. Antibiotic therapy should be continued for 7–10 days and should be tailored to the organism recovered in culture. If the household contacts of a patient with *H. influenzae* epiglottitis include an unvaccinated child under age 4, all members of the household (including the patient) should receive prophylactic rifampin for 4 days to eradicate carriage of *H. influenzae*.

INFECTIONS OF DEEP NECK STRUCTURES

Deep neck infections are usually extensions of infection from other primary sites, most often within the pharynx or oral cavity. Many of these infections are life threatening but are difficult to detect at early stages, when they may be more easily managed. Three of the most clinically relevant spaces in the neck are the submandibular (and sublingual) space, the lateral pharyngeal (or parapharyngeal) space, and the retropharyngeal space. These spaces communicate with one another and with other important structures in the head, neck, and thorax, providing pathogens with easy access to areas that include the mediastinum, carotid sheath, skull base, and meninges. Once infection reaches these sensitive areas, mortality rates can be as high as 20–50%.

Infection of the submandibular and/or sublingual space typically originates from an infected or recently extracted lower tooth. The result is the severe, life-threatening infection referred to as Ludwig’s angina (see “Oral Infections,” above). Infection of the lateral pharyngeal (or parapharyngeal) space is most often a complication of common infections of the oral cavity and upper respiratory tract, including tonsillitis, peritonsillar abscess, pharyngitis, mastoiditis, and periodontal infection. This space, situated deep in the lateral wall of

the pharynx, contains a number of sensitive structures, including the carotid artery, internal jugular vein, cervical sympathetic chain, and portions of cranial nerves IX through XII; at its distal end, it opens into the posterior mediastinum. Involvement of this space with infection can therefore be rapidly fatal. Examination may reveal some tonsillar displacement, trismus, and neck rigidity, but swelling of the lateral pharyngeal wall can easily be missed. The diagnosis can be confirmed by CT. Treatment consists of airway management, operative drainage of fluid collections, and at least 10 days of IV therapy with an antibiotic active against streptococci and oral anaerobes (e.g., ampicillin/sulbactam). A particularly severe form of this infection involving the components of the carotid sheath (postanginal septicemia, Lemierre’s disease) is described above (see “Oral Infections”). Infection of the retropharyngeal space also can be extremely dangerous, as this space runs posterior to the pharynx from the skull base to the superior mediastinum. Infections in this space are more common among children <5 years old because of the presence of several small retropharyngeal lymph nodes that typically atrophy by age 4 years. Infection is usually a consequence of extension from another site of infection—most commonly, acute pharyngitis. Other sources include otitis media, tonsillitis, dental infections, Ludwig’s angina, and anterior extension of vertebral osteomyelitis. Retropharyngeal space infection also can follow penetrating trauma to the posterior pharynx (e.g., from an endoscopic procedure). Infections are commonly polymicrobial, involving a mixture of aerobes and anaerobes; group A β -hemolytic streptococci and *S. aureus* are the most common pathogens. *M. tuberculosis* was a common cause in the past but now is rarely involved in the United States.

Patients with retropharyngeal abscess typically present with sore throat, fever, dysphagia, and neck pain and are often drooling because of difficulty and pain with swallowing. Examination may reveal tender cervical adenopathy, neck swelling, and diffuse erythema and edema of the posterior pharynx as well as a bulge in the posterior pharyngeal wall that may not be obvious on routine inspection. A soft tissue mass is usually demonstrable by lateral neck radiography or CT. Because of the risk of airway obstruction, treatment begins with securing of the airway, followed by a combination of surgical drainage and IV antibiotic administration. Initial empirical therapy should cover streptococci, oral anaerobes, and *S. aureus*; ampicillin/sulbactam, clindamycin plus ceftriaxone, or meropenem is usually effective. Complications result primarily from extension to other areas (e.g., rupture into the posterior pharynx may lead to aspiration pneumonia and empyema). Extension may also occur to the lateral pharyngeal space and mediastinum, resulting in mediastinitis and pericarditis, or into nearby major blood vessels. All these events are associated with a high mortality rate.

45

Oral Manifestations of Disease

Samuel C. Durso

As primary care physicians and consultants, internists are often asked to evaluate patients with disease of the oral soft tissues, teeth, and pharynx. Knowledge of the oral milieu and its unique structures is necessary to guide preventive services and recognize oral manifestations of local or systemic disease (**Chap. 46e**). Furthermore, internists frequently collaborate with dentists in the care of patients who have a variety of medical conditions that affect oral health or who undergo dental procedures that increase their risk of medical complications.

DISEASES OF THE TEETH AND PERIODONTAL STRUCTURES

Tooth formation begins during the sixth week of embryonic life and continues through 17 years of age. Teeth start to develop in utero and continue to develop until after the tooth erupts. Normally, all 20 deciduous teeth have erupted by age 3 and have been shed by age 13. Permanent teeth, eventually totaling 32, begin to erupt by age 6 and

have completely erupted by age 14, though third molars (“wisdom teeth”) may erupt later.

The erupted tooth consists of the visible *crown* covered with enamel and the root submerged below the gum line and covered with bonelike *cementum*. *Dentin*, a material that is denser than bone and exquisitely sensitive to pain, forms the majority of the tooth substance, surrounding a core of myxomatous *pulp* containing the vascular and nerve supply. The tooth is held firmly in the alveolar socket by the *periodontium*, supporting structures that consist of the gingivae, alveolar bone, cementum, and periodontal ligament. The periodontal ligament tenaciously binds the tooth’s cementum to the alveolar bone. Above this ligament is a collar of attached gingiva just below the crown. A few millimeters of unattached or free gingiva (1–3 mm) overlap the base of the crown, forming a shallow sulcus along the gum-tooth margin.

Dental Caries, Pulpal and Periapical Disease, and Complications Dental caries usually begin asymptotically as a destructive infectious process of the enamel. Bacteria—principally *Streptococcus mutans*—colonize the organic buffering biofilm (*plaque*) on the tooth surface. If not removed by brushing or by the natural cleansing and antibacterial action of saliva, bacterial acids can demineralize the enamel. Fissures and pits on the occlusal surfaces are the most frequent sites of early decay. Surfaces between the teeth, adjacent to tooth restorations and exposed roots, are also vulnerable, particularly as individuals age. Over time, dental caries extend to the underlying dentin, leading to cavitation of the enamel. Without management, the caries will penetrate to the tooth pulp, producing *acute pulpitis*. At this stage, when the pulp infection is limited, the tooth may become sensitive to percussion and to hot or cold, and pain resolves immediately when the irritating stimulus is removed. Should the infection spread throughout the pulp, *irreversible pulpitis* occurs, leading to *pulp necrosis*. At this later stage, pain can be severe and has a sharp or throbbing visceral quality that may be worse when the patient lies down. Once pulp necrosis is complete, pain may be constant or intermittent, but cold sensitivity is lost.

Treatment of caries involves removal of the softened and infected hard tissue and restoration of the tooth structure with silver amalgam, glass ionomer, composite resin, or gold. Once irreversible pulpitis occurs, root canal therapy becomes necessary; removal of the contents of the pulp chamber and root canals is followed by thorough cleaning and filling with an inert material. Alternatively, the tooth may be extracted.

Pulpal infection leads to *periapical abscess* formation, which can produce pain on chewing. If the infection is mild and chronic, a *periapical granuloma* or eventually a *periapical cyst* forms, either of which produces radiolucency at the root apex. When unchecked, a periapical abscess can erode into the alveolar bone, producing osteomyelitis; penetrate and drain through the gingivae, producing a parulis (gumboil); or track along deep fascial planes, producing virulent cellulitis (Ludwig’s angina) involving the submandibular space and floor of the mouth (**Chap. 201**). Elderly patients, patients with diabetes mellitus, and patients taking glucocorticoids may experience little or no pain or fever as these complications develop.

Periodontal Disease Periodontal disease and dental caries are the primary causes of tooth loss. Like dental caries, chronic infection of the gingiva and anchoring structures of the tooth begins with formation of bacterial plaque. The process begins at the gum line. Plaque and *calculus* (calcified plaque) are preventable by appropriate daily oral hygiene, including periodic professional cleaning. Left undisturbed, chronic inflammation can ensue and produce hyperemia of the free and attached gingivae (*gingivitis*), which then typically bleed with brushing. If this issue is ignored, severe *periodontitis* can develop, leading to deepening of the physiologic sulcus and destruction of the periodontal ligament. Gingival pockets develop around the teeth. As the periodontium (including the supporting bone) is destroyed, the teeth loosen. A role for chronic inflammation due to chronic periodontal disease in promoting coronary heart disease and stroke has been proposed. Epidemiologic studies have demonstrated a moderate but significant association between chronic periodontal inflammation and atherosclerosis, though a causal role remains unproven.

Acute and aggressive forms of periodontal disease are less common than the chronic forms described above. However, if the host is stressed or exposed to a new pathogen, rapidly progressive and destructive disease of the periodontal tissue can occur. A virulent example is *acute necrotizing ulcerative gingivitis*. Stress and poor oral hygiene are risk factors. The presentation includes sudden gingival inflammation, ulceration, bleeding, interdental gingival necrosis, and fetid halitosis. *Localized juvenile periodontitis*, which is seen in adolescents, is particularly destructive and appears to be associated with impaired neutrophil chemotaxis. *AIDS-related periodontitis* resembles acute necrotizing ulcerative gingivitis in some patients and a more destructive form of adult chronic periodontitis in others. It may also produce a gangrene-like destructive process of the oral soft tissues and bone that resembles *noma*, an infectious condition seen in severely malnourished children in developing nations.

Prevention of Tooth Decay and Periodontal Infection Despite the reduced prevalences of dental caries and periodontal disease in the United States (due in large part to water fluoridation and improved dental care, respectively), both diseases constitute a major public health problem worldwide, particularly in certain groups. The internist should promote preventive dental care and hygiene as part of health maintenance. Populations at high risk for dental caries and periodontal disease include those with hyposalivation and/or xerostomia, diabetics, alcoholics, tobacco users, persons with Down syndrome, and those with gingival hyperplasia. Furthermore, patients lacking access to dental care (e.g., as a result of low socioeconomic status) and patients with a reduced ability to provide self-care (e.g., individuals with disabilities, nursing home residents, and persons with dementia or upper-extremity disability) suffer at a disproportionate rate. It is important to provide counseling regarding regular dental hygiene and professional cleaning, use of fluoride-containing toothpaste, professional fluoride treatments, and (for patients with limited dexterity) use of electric toothbrushes and also to instruct persons caring for those who are not capable of self-care. Cost, fear of dental care, and differences in language and culture create barriers that prevent some people from seeking preventive dental services.

Developmental and Systemic Disease Affecting the Teeth and Periodontium

In addition to posing cosmetic issues, *malocclusion*, the most common developmental oral problem, can interfere with mastication unless corrected through orthodontic and surgical techniques. Impacted third molars are common and can become infected or erupt into an insufficient space. Acquired prognathism due to *acromegaly* may also lead to malocclusion, as may deformity of the maxilla and mandible due to *Paget’s disease* of the bone. Delayed tooth eruption, a receding chin, and a protruding tongue are occasional features of *cretinism* and *hypopituitarism*. Congenital syphilis produces tapering, notched (*Hutchinson’s*) incisors and finely nodular (*mulberry*) molar crowns. *Enamel hypoplasia* results in crown defects ranging from pits to deep fissures of primary or permanent teeth. Intrauterine infection (syphilis, rubella), vitamin deficiency (A, C, or D), disorders of calcium metabolism (malabsorption, vitamin D-resistant rickets, hypoparathyroidism), prematurity, high fever, and rare inherited defects (*amelogenesis imperfecta*) are all causes. Tetracycline, given in sufficiently high doses during the first 8 years of life, may produce enamel hypoplasia and discoloration. Exposure to endogenous pigments can discolor developing teeth; etiologies include *erythroblastosis fetalis* (green or bluish-black), congenital liver disease (green or yellow-brown), and porphyria (red or brown that fluoresces with ultraviolet light). *Mottled enamel* occurs if excessive fluoride is ingested during development. Worn enamel is seen with age, bruxism, or excessive acid exposure (e.g., chronic gastric reflux or bulimia). Celiac disease is associated with nonspecific enamel defects in children but not in adults.

Total or partial tooth loss resulting from periodontitis is seen with cyclic neutropenia, Papillon-Lefèvre syndrome, Chédiak-Higashi syndrome, and leukemia. Rapid focal tooth loosening is most often due to infection, but rarer causes include Langerhans cell histiocytosis, Ewing’s sarcoma, osteosarcoma, and Burkitt’s lymphoma. Early loss of

primary teeth is a feature of *hypophosphatasia*, a rare congenital error of metabolism.

Pregnancy may produce gingivitis and localized *pyogenic granulomas*. Severe periodontal disease occurs in uncontrolled diabetes mellitus. *Gingival hyperplasia* may be caused by phenytoin, calcium channel blockers (e.g., nifedipine), and cyclosporine, though excellent daily oral care can prevent or reduce its occurrence. *Idiopathic familial gingival fibromatosis* and several syndrome-related disorders cause similar conditions. Discontinuation of the medication may reverse the drug-induced form, though surgery may be needed to control both of the latter entities. *Linear gingival erythema* is variably seen in patients with advanced HIV infection and probably represents immune deficiency and decreased neutrophil activity. Diffuse or focal gingival swelling may be a feature of early or late acute myelomonocytic leukemia as well as of other lymphoproliferative disorders. A rare but pathognomonic sign of granulomatosis with polyangiitis is a red-purple, granular gingivitis (*strawberry gums*).

DISEASES OF THE ORAL MUCOSA

Infections Most oral mucosal diseases involve microorganisms (Table 45-1).

Pigmented Lesions See Table 45-2.

Dermatologic Diseases See Tables 45-1, 45-2, and 45-3 and Chaps. 70-74.

Diseases of the Tongue See Table 45-4.

HIV Disease and AIDS See Tables 45-1, 45-2, 45-3, and 45-5; Chap. 226; and Fig. 218-3.

Ulcers Ulceration is the most common oral mucosal lesion. Although there are many causes, the host and the pattern of lesions, including the presence of organ system features, narrow the differential diagnosis (Table 45-1). Most acute ulcers are painful and self-limited. Recurrent aphthous ulcers and herpes simplex account for the majority. Persistent and deep aphthous ulcers can be idiopathic or can accompany HIV/AIDS. Aphthous lesions are often the presenting symptom in *Behçet's syndrome* (Chap. 387). Similar-appearing, though less painful, lesions may occur in reactive arthritis, and aphthous ulcers are occasionally present during phases of *discoid* or *systemic lupus erythematosus* (Chap. 382). Aphthous-like ulcers are seen in *Crohn's disease* (Chap. 351), but, unlike the common aphthous variety, they may exhibit granulomatous inflammation on histologic examination. Recurrent aphthae are more prevalent in patients with *celiac disease* and have been reported to remit with elimination of gluten.

Of major concern are chronic, relatively painless ulcers and mixed red/white patches (erythroplakia and leukoplakia) of >2 weeks' duration. Squamous cell carcinoma and premalignant dysplasia should be considered early and a diagnostic biopsy performed. This awareness and this procedure are critically important because early-stage malignancy is vastly more treatable than late-stage disease. High-risk sites include the lower lip, floor of the mouth, ventral and lateral tongue, and soft palate-tonsillar pillar complex. Significant risk factors for oral cancer in Western countries include sun exposure (lower lip), tobacco and alcohol use, and human papillomavirus infection. In India and some other Asian countries, smokeless tobacco mixed with betel nut, slaked lime, and spices is a common cause of oral cancer. Rarer causes of chronic oral ulcer, such as tuberculosis, fungal infection, granulomatosis with polyangiitis, and midline granuloma may look identical to carcinoma. Making the correct diagnosis depends on recognizing other clinical features and performing a biopsy of the lesion. The syphilitic chancre is typically painless and therefore easily missed. Regional lymphadenopathy is invariably present. The syphilitic etiology is confirmed with appropriate bacterial and serologic tests.

Disorders of mucosal fragility often produce painful oral ulcers that fail to heal within 2 weeks. *Mucous membrane pemphigoid* and *pemphigus vulgaris* are the major acquired disorders. While their clinical features are often distinctive, a biopsy or immunohistochemical

examination should be performed to diagnose these entities and to distinguish them from *lichen planus* and drug reactions.

Hematologic and Nutritional Disease Internists are more likely to encounter patients with acquired, rather than congenital, bleeding disorders. Bleeding should stop 15 min after minor trauma and within an hour after tooth extraction if local pressure is applied. More prolonged bleeding, if not due to continued injury or rupture of a large vessel, should lead to investigation for a clotting abnormality. In addition to bleeding, petechiae and ecchymoses are prone to occur at the vibrating line between the soft and hard palates in patients with platelet dysfunction or thrombocytopenia.

All forms of leukemia, but particularly *acute myelomonocytic leukemia*, can produce gingival bleeding, ulcers, and gingival enlargement. Oral ulcers are a feature of agranulocytosis, and ulcers and mucositis are often severe complications of chemotherapy and radiation therapy for hematologic and other malignancies. *Plummer-Vinson syndrome* (iron deficiency, angular stomatitis, glossitis, and dysphagia) raises the risk of oral squamous cell cancer and esophageal cancer at the postcricoid tissue web. Atrophic papillae and a red, burning tongue may occur with pernicious anemia. Deficiencies in B-group vitamins produce many of these same symptoms as well as oral ulceration and cheilosis. Consequences of *scurvy* include swollen, bleeding gums; ulcers; and loosening of the teeth.

NONDENTAL CAUSES OF ORAL PAIN

Most, but not all, oral pain emanates from inflamed or injured tooth pulp or periodontal tissues. Nonodontogenic causes are often overlooked. In most instances, toothache is predictable and proportional to the stimulus applied, and an identifiable condition (e.g., caries, abscess) is found. Local anesthesia eliminates pain originating from dental or periodontal structures, but not referred pains. The most common nondental source of pain is myofascial pain referred from muscles of mastication, which become tender and ache with increased use. Many sufferers exhibit *bruxism* (grinding of the teeth) secondary to stress and anxiety. *Temporomandibular joint disorder* is closely related. It affects both sexes, with a higher prevalence among women. Features include pain, limited mandibular movement, and temporomandibular joint sounds. The etiologies are complex; malocclusion does not play the primary role once attributed to it. *Osteoarthritis* is a common cause of masticatory pain. Anti-inflammatory medication, jaw rest, soft foods, and heat provide relief. The temporomandibular joint is involved in 50% of patients with *rheumatoid arthritis*, and its involvement is usually a late feature of severe disease. Bilateral preauricular pain, particularly in the morning, limits range of motion.

Migrainous neuralgia may be localized to the mouth. Episodes of pain and remission without an identifiable cause and a lack of relief with local anesthesia are important clues. *Trigeminal neuralgia* (*tic douloureux*) can involve the entire branch or part of the mandibular or maxillary branch of the fifth cranial nerve and can produce pain in one or a few teeth. Pain may occur spontaneously or may be triggered by touching the lip or gingiva, brushing the teeth, or chewing. *Glossopharyngeal neuralgia* produces similar acute neuropathic symptoms in the distribution of the ninth cranial nerve. Swallowing, sneezing, coughing, or pressure on the tragus of the ear triggers pain that is felt in the base of the tongue, pharynx, and soft palate and may be referred to the temporomandibular joint. *Neuritis* involving the maxillary and mandibular divisions of the trigeminal nerve (e.g., maxillary sinusitis, neuroma, and leukemic infiltrate) is distinguished from ordinary toothache by the neuropathic quality of the pain. Occasionally, *phantom pain* follows tooth extraction. Pain and hyperalgesia behind the ear and on the side of the face in the day or so before facial weakness develops often constitute the earliest symptom of *Bell's palsy*. Likewise, similar symptoms may precede visible lesions of herpes zoster infecting the seventh nerve (*Ramsey-Hunt syndrome*) or trigeminal nerve. *Postherpetic neuralgia* may follow either condition. *Coronary ischemia* may produce pain exclusively in the face and jaw; as in typical angina pectoris, this pain is usually reproducible with increased myocardial demand. Aching in several upper molar or

TABLE 45-1 VESICULAR, BULLOUS, OR ULCERATIVE LESIONS OF THE ORAL MUCOSA

Condition	Usual Location	Clinical Features	Course
Viral Diseases			
Primary acute herpetic gingivostomatitis (HSV type 1; rarely type 2)	Lip and oral mucosa (buccal, gingival, lingual mucosa)	Labial vesicles that rupture and crust, and intraoral vesicles that quickly ulcerate; extremely painful; acute gingivitis, fever, malaise, foul odor, and cervical lymphadenopathy; occurs primarily in infants, children, and young adults	Heals spontaneously in 10–14 days; unless secondarily infected, lesions lasting >3 weeks are not due to primary HSV infection
Recurrent herpes labialis	Mucocutaneous junction of lip, perioral skin	Eruption of groups of vesicles that may coalesce, then rupture and crust; painful to pressure or spicy foods	Lasts ~1 week, but condition may be prolonged if secondarily infected; if severe, topical or oral antiviral treatment may reduce healing time
Recurrent intraoral herpes simplex	Palate and gingiva	Small vesicles on keratinized epithelium that rupture and coalesce; painful	Heals spontaneously in ~1 week; if severe, topical or oral antiviral treatment may reduce healing time
Chickenpox (VZV)	Gingiva and oral mucosa	Skin lesions may be accompanied by small vesicles on oral mucosa that rupture to form shallow ulcers; may coalesce to form large bullous lesions that ulcerate; mucosa may have generalized erythema	Lesions heal spontaneously within 2 weeks
Herpes zoster (VZV reactivation)	Cheek, tongue, gingiva, or palate	Unilateral vesicular eruptions and ulceration in linear pattern following sensory distribution of trigeminal nerve or one of its branches	Gradual healing without scarring unless secondarily infected; postherpetic neuralgia is common; oral acyclovir, famciclovir, or valacyclovir reduces healing time and postherpetic neuralgia
Infectious mononucleosis (Epstein-Barr virus)	Oral mucosa	Fatigue, sore throat, malaise, fever, and cervical lymphadenopathy; numerous small ulcers usually appear several days before lymphadenopathy; gingival bleeding and multiple petechiae at junction of hard and soft palates	Oral lesions disappear during convalescence; no treatment is given, though glucocorticoids are indicated if tonsillar swelling compromises the airway
Herpangina (coxsackievirus A; also possibly coxsackievirus B and echovirus)	Oral mucosa, pharynx, tongue	Sudden onset of fever, sore throat, and oropharyngeal vesicles, usually in children <4 years old, during summer months; diffuse pharyngeal congestion and vesicles (1–2 mm), grayish-white surrounded by red areola; vesicles enlarge and ulcerate	Incubation period of 2–9 days; fever for 1–4 days; recovery uneventful
Hand-foot-and-mouth disease (most commonly coxsackievirus A16)	Oral mucosa, pharynx, palms, and soles	Fever, malaise, headache with oropharyngeal vesicles that become painful, shallow ulcers; highly infectious; usually affects children under age 10	Incubation period 2–18 days; lesions heal spontaneously in 2–4 weeks
Primary HIV infection	Gingiva, palate, and pharynx	Acute gingivitis and oropharyngeal ulceration, associated with febrile illness resembling mononucleosis and including lymphadenopathy	Followed by HIV seroconversion, asymptomatic HIV infection, and usually ultimately by HIV disease
Bacterial or Fungal Diseases			
Acute necrotizing ulcerative gingivitis (“trench mouth”)	Gingiva	Painful, bleeding gingiva characterized by necrosis and ulceration of gingival papillae and margins plus lymphadenopathy and foul breath	Debridement and diluted (1:3) peroxide lavage provide relief within 24 h; antibiotics in acutely ill patients; relapse may occur
Prenatal (congenital) syphilis	Palate, jaws, tongue, and teeth	Gummatous involvement of palate, jaws, and facial bones; Hutchinson’s incisors, mulberry molars, glossitis, mucous patches, and fissures at corner of mouth	Tooth deformities in permanent dentition irreversible
Primary syphilis (chancere)	Lesion appearing where organism enters body; may occur on lips, tongue, or tonsillar area	Small papule developing rapidly into a large, painless ulcer with indurated border; unilateral lymphadenopathy; chancre and lymph nodes containing spirochetes; serologic tests positive by third to fourth weeks	Healing of chancre in 1–2 months, followed by secondary syphilis in 6–8 weeks
Secondary syphilis	Oral mucosa frequently involved with mucous patches, which occur primarily on palate and also at commissures of mouth	Maculopapular lesions of oral mucosa, 5–10 mm in diameter with central ulceration covered by grayish membrane; eruptions occurring on various mucosal surfaces and skin, accompanied by fever, malaise, and sore throat	Lesions may persist from several weeks to a year
Tertiary syphilis	Palate and tongue	Gummatous infiltration of palate or tongue followed by ulceration and fibrosis; atrophy of tongue papillae produces characteristic bald tongue and glossitis	Gumma may destroy palate, causing complete perforation
Gonorrhea	Lesions may occur in mouth at site of inoculation or secondarily by hematogenous spread from a primary focus	Most pharyngeal infection is asymptomatic; may produce burning or itching sensation; oropharynx and tonsils may be ulcerated and erythematous; saliva viscous and fetid	More difficult to eradicate than urogenital infection, though pharyngitis usually resolves with appropriate antimicrobial treatment
Tuberculosis	Tongue, tonsillar area, soft palate	Painless, solitary, 1- to 5-cm, irregular ulcer covered with persistent exudate; ulcer has firm undermined border	Autoinoculation from pulmonary infection is usual; lesions resolve with appropriate antimicrobial therapy

(Continued)

TABLE 45-1 VESICULAR, BULLOUS, OR ULCERATIVE LESIONS OF THE ORAL MUCOSA (CONTINUED)

Condition	Usual Location	Clinical Features	Course
Cervicofacial actinomycosis	Swellings in region of face, neck, and floor of mouth	Infection may be associated with extraction, jaw fracture, or eruption of molar tooth; in acute form, resembles acute pyogenic abscess, but contains yellow "sulfur granules" (gram-positive mycelia and their hyphae)	Typically, swelling is hard and grows painlessly; multiple abscesses with draining tracts develop; penicillin first choice; surgery usually necessary
Histoplasmosis	Any area of the mouth, particularly tongue, gingiva, or palate	Nodular, verrucous, or granulomatous lesions; ulcers are indurated and painful; usual source hematogenous or pulmonary, but may be primary	Systemic antifungal therapy necessary
Candidiasis ^a			
Dermatologic Diseases			
Mucous membrane pemphigoid	Typically produces marked gingival erythema and ulceration; other areas of oral cavity, esophagus, and vagina may be affected	Painful, grayish-white collapsed vesicles or bullae of full-thickness epithelium with peripheral erythematous zone; gingival lesions desquamate, leaving ulcerated area	Protracted course with remissions and exacerbations; involvement of different sites develops slowly; glucocorticoids may temporarily reduce symptoms but do not control disease
EM minor and EM major (Stevens-Johnson syndrome)	Primarily oral mucosa and skin of hands and feet	Intraoral ruptured bullae surrounded by inflammatory area; lips may show hemorrhagic crusts; "iris" or "target" lesion on skin is pathognomonic; patient may have severe signs of toxicity	Onset very rapid; usually idiopathic, but may be associated with trigger such as drug reaction; condition may last 3–6 weeks; mortality rate for untreated EM major is 5–15%
Pemphigus vulgaris	Oral mucosa and skin; sites of mechanical trauma (soft/hard palate, frenulum, lips, buccal mucosa)	Usually (>70%) presents with oral lesions; fragile, ruptured bullae and ulcerated oral areas; mostly in older adults	With repeated occurrence of bullae, toxicity may lead to cachexia, infection, and death within 2 years; often controllable with oral glucocorticoids
Lichen planus	Oral mucosa and skin	White striae in mouth; purplish nodules on skin at sites of friction; occasionally causes oral mucosal ulcers and erosive gingivitis	White striae alone usually asymptomatic; erosive lesions often difficult to treat, but may respond to glucocorticoids
Other Conditions			
Recurrent aphthous ulcers	Usually on nonkeratinized oral mucosa (buccal and labial mucosa, floor of mouth, soft palate, lateral and ventral tongue)	Single or clustered painful ulcers with surrounding erythematous border; lesions may be 1–2 mm in diameter in crops (herpetiform), 1–5 mm (minor), or 5–15 mm (major)	Lesions heal in 1–2 weeks but may recur monthly or several times a year; protective barrier with benzocaine and topical glucocorticoids relieve symptoms; systemic glucocorticoids may be needed in severe cases
Behçet's syndrome	Oral mucosa, eyes, genitalia, gut, and CNS	Multiple aphthous ulcers in mouth; inflammatory ocular changes, ulcerative lesions on genitalia; inflammatory bowel disease and CNS disease	Oral lesions often first manifestation; persist several weeks and heal without scarring
Traumatic ulcers	Anywhere on oral mucosa; dentures frequently responsible for ulcers in vestibule	Localized, discrete ulcerated lesions with red border; produced by accidental biting of mucosa, penetration by foreign object, or chronic irritation by dentures	Lesions usually heal in 7–10 days when irritant is removed, unless secondarily infected
Squamous cell carcinoma	Any area of mouth, most commonly on lower lip, lateral borders of tongue, and floor of mouth	Red, white, or red and white ulcer with elevated or indurated border; failure to heal; pain not prominent in early lesions	Invades and destroys underlying tissues; frequently metastasizes to regional lymph nodes
Acute myeloid leukemia (usually monocytic)	Gingiva	Gingival swelling and superficial ulceration followed by hyperplasia of gingiva with extensive necrosis and hemorrhage; deep ulcers may occur elsewhere on mucosa, complicated by secondary infection	Usually responds to systemic treatment of leukemia; occasionally requires local irradiation
Lymphoma	Gingiva, tongue, palate, and tonsillar area	Elevated, ulcerated area that may proliferate rapidly, giving appearance of traumatic inflammation	Fatal if untreated; may indicate underlying HIV infection
Chemical or thermal burns	Any area in mouth	White slough due to contact with corrosive agents (e.g., aspirin, hot cheese) applied locally; removal of slough leaves raw, painful surface	Lesion heals in several weeks if not secondarily infected

^aSee Table 45-3.

Abbreviations: CNS, central nervous system; EM, erythema multiforme; HSV, herpes simplex virus; VZV, varicella-zoster virus.

premolar teeth that is unrelieved by anesthetizing the teeth may point to *maxillary sinusitis*.

Giant cell arteritis is notorious for producing headache, but it may also produce facial pain or sore throat without headache. Jaw and tongue claudication with chewing or talking is relatively common. Tongue infarction is rare. Patients with subacute thyroiditis often experience pain referred to the face or jaw before the tenderness of the thyroid gland and transient hyperthyroidism are appreciated.

"Burning mouth syndrome" (*glossodynia*) occurs in the absence of an identifiable cause (e.g., vitamin B₁₂ deficiency, iron deficiency, diabetes mellitus, low-grade *Candida* infection, food sensitivity, or subtle xerostomia) and predominantly affects postmenopausal women. The

etiology may be neuropathic. Clonazepam, α -lipoic acid, and cognitive behavioral therapy have benefited some patients. Some cases associated with an angiotensin-converting enzyme inhibitor have remitted when treatment with the drug was discontinued.

DISEASES OF THE SALIVARY GLANDS

Saliva is essential to oral health. Its absence leads to dental caries, periodontal disease, and difficulties in wearing dental prostheses, masticating, and speaking. Its major components, water and mucin, serve as a cleansing solvent and lubricating fluid. In addition, saliva contains antimicrobial factors (e.g., lysozyme, lactoperoxidase, secretory IgA), epidermal growth factor, minerals, and buffering systems. The

TABLE 45-2 PIGMENTED LESIONS OF THE ORAL MUCOSA

Condition	Usual Location	Clinical Features	Course
Oral melanotic macule	Any area of mouth	Discrete or diffuse, localized, brown to black macule	Remains indefinitely; no growth
Diffuse melanin pigmentation	Any area of mouth	Diffuse pale to dark-brown pigmentation; may be physiologic ("racial") or due to smoking	Remains indefinitely
Nevi	Any area of mouth	Discrete, localized, brown to black pigmentation	Remains indefinitely
Malignant melanoma	Any area of mouth	Can be flat and diffuse, painless, brown to black; or can be raised and nodular	Expands and invades early; metastasis leads to death
Addison's disease	Any area of mouth, but mostly buccal mucosa	Blotches or spots of bluish-black to dark-brown pigmentation occurring early in disease, accompanied by diffuse pigmentation of skin; other symptoms of adrenal insufficiency	Condition controlled by adrenal steroid replacement
Peutz-Jeghers syndrome	Any area of mouth	Dark-brown spots on lips, buccal mucosa, with characteristic distribution of pigment around lips, nose, and eyes and on hands; concomitant intestinal polyposis	Oral pigmented lesions remain indefinitely; gastrointestinal polyps may become malignant
Drug ingestion (neuroleptics, oral contraceptives, minocycline, zidovudine, quinine derivatives)	Any area of mouth	Brown, black, or gray areas of pigmentation	Gradually disappears following cessation of drug intake
Amalgam tattoo	Gingiva and alveolar mucosa	Small blue-black pigmented areas associated with embedded amalgam particles in soft tissues; may show up on radiographs as radiopaque particles in some cases	Remains indefinitely
Heavy metal pigmentation (bismuth, mercury, lead)	Gingival margin	Thin blue-black pigmented line along gingival margin; rarely seen except in children exposed to lead-based paint	Indicative of systemic absorption; no significance for oral health
Black hairy tongue	Dorsum of tongue	Elongation of filiform papillae of tongue, which become stained by coffee, tea, tobacco, or pigmented bacteria	Improves within 1–2 weeks with gentle brushing of tongue or (if due to bacterial overgrowth) discontinuation of antibiotic
Fordyce spots	Buccal and labial mucosa	Numerous small yellowish spots just beneath mucosal surface; no symptoms; due to hyperplasia of sebaceous glands	Benign; remains without apparent change
Kaposi's sarcoma	Palate most common, but may occur at any other site	Red or blue plaques of variable size and shape; often enlarge, become nodular, and may ulcerate	Usually indicative of HIV infection or non-Hodgkin's lymphoma; rarely fatal, but may require treatment for comfort or cosmesis
Mucous retention cysts	Buccal and labial mucosa	Bluish, clear fluid-filled cyst due to extravasated mucus from injured minor salivary gland	Benign; painless unless traumatized; may be removed surgically

TABLE 45-3 WHITE LESIONS OF ORAL MUCOSA

Condition	Usual Location	Clinical Features	Course
Lichen planus	Buccal mucosa, tongue, gingiva, and lips; skin	Striae, white plaques, red areas, ulcers in mouth; purplish papules on skin; may be asymptomatic, sore, or painful; lichenoid drug reactions may look similar	Protracted; responds to topical glucocorticoids
White sponge nevus	Oral mucosa, vagina, anal mucosa	Painless white thickening of epithelium; adolescence/early adulthood onset; familial	Benign and permanent
Smoker's leukoplakia and smokeless tobacco lesions	Any area of oral mucosa, sometimes related to location of habit	White patch that may become firm, rough, or red-fissured and ulcerated; may become sore and painful but is usually painless	May or may not resolve with cessation of habit; 2% of patients develop squamous cell carcinoma; early biopsy essential
Erythroplakia with or without white patches	Floor of mouth commonly affected in men; tongue and buccal mucosa in women	Velvety, reddish plaque; occasionally mixed with white patches or smooth red areas	High risk of squamous cell cancer; early biopsy essential
Candidiasis	Any area in mouth	<i>Pseudomembranous type</i> ("thrush"): creamy white curdlike patches that reveal a raw, bleeding surface when scraped; found in sick infants, debilitated elderly patients receiving high-dose glucocorticoids or broad-spectrum antibiotics, and patients with AIDS <i>Erythematous type</i> : flat, red, sometimes sore areas in same groups of patients <i>Candidal leukoplakia</i> : nonremovable white thickening of epithelium due to <i>Candida</i> <i>Angular cheilitis</i> : sore fissures at corner of mouth	Responds favorably to antifungal therapy and correction of predisposing causes where possible Course same as for pseudomembranous type Responds to prolonged antifungal therapy Responds to topical antifungal therapy
Hairy leukoplakia	Usually on lateral tongue, rarely elsewhere on oral mucosa	White areas ranging from small and flat to extensive accentuation of vertical folds; found in HIV carriers (all risk groups for AIDS)	Due to Epstein-Barr virus; responds to high-dose acyclovir but recurs; rarely causes discomfort unless secondarily infected with <i>Candida</i>
Warts (human papillomavirus)	Anywhere on skin and oral mucosa	Single or multiple papillary lesions with thick, white, keratinized surfaces containing many pointed projections; cauliflower lesions covered with normal-colored mucosa or multiple pink or pale bumps (focal epithelial hyperplasia)	Lesions grow rapidly and spread; squamous cell carcinoma must be ruled out with biopsy; excision or laser therapy; may regress in HIV-infected patients receiving antiretroviral therapy

TABLE 45-4 ALTERATIONS OF THE TONGUE

Type of Change	Clinical Features
Size or Morphology	
Macroglossia	Enlarged tongue that may be part of a syndrome found in developmental conditions such as Down syndrome, Simpson-Golabi-Behmel syndrome, or Beckwith-Wiedemann syndrome; may be due to tumor (hemangioma or lymphangioma), metabolic disease (e.g., primary amyloidosis), or endocrine disturbance (e.g., acromegaly or cretinism); may occur when all teeth are removed
Fissured ("scrotal") tongue	Dorsal surface and sides of tongue covered by painless shallow or deep fissures that may collect debris and become irritated
Median rhomboid glossitis	Congenital abnormality with ovoid, denuded area in median posterior portion of tongue; may be associated with candidiasis and may respond to antifungal treatment
Color	
"Geographic" tongue (benign migratory glossitis)	Asymptomatic inflammatory condition of tongue, with rapid loss and regrowth of filiform papillae leading to appearance of denuded red patches "wandering" across surface
Hairy tongue	Elongation of filiform papillae of medial dorsal surface area due to failure of keratin layer of papillae to desquamate normally; brownish-black coloration may be due to staining by tobacco, food, or chromogenic organisms
"Strawberry" and "raspberry" tongue	Appearance of tongue during scarlet fever due to hypertrophy of fungiform papillae as well as changes in filiform papillae
"Bald" tongue	Atrophy may be associated with xerostomia, pernicious anemia, iron-deficiency anemia, pellagra, or syphilis; may be accompanied by painful burning sensation; may be an expression of erythematous candidiasis and respond to antifungal treatment

TABLE 45-5 ORAL LESIONS ASSOCIATED WITH HIV INFECTION

Lesion Morphology	Etiologies
Papules, nodules, plaques	Candidiasis (hyperplastic and pseudomembranous) ^a Condyloma acuminatum (human papillomavirus infection) Squamous cell carcinoma (preinvasive and invasive) Non-Hodgkin's lymphoma ^a Hairy leukoplakia ^a
Ulcers	Recurrent aphthous ulcers ^a Angular cheilitis Squamous cell carcinoma Acute necrotizing ulcerative gingivitis ^a Necrotizing ulcerative periodontitis ^a Necrotizing ulcerative stomatitis Non-Hodgkin's lymphoma ^a Viral infection (herpes simplex, herpes zoster, cytomegalovirus infection) Infection caused by <i>Mycobacterium tuberculosis</i> or <i>Mycobacterium avium-intracellulare</i> Fungal infection (histoplasmosis, cryptococcosis, candidiasis, geotrichosis, aspergillosis) Bacterial infection (<i>Escherichia coli</i> , <i>Enterobacter cloacae</i> , <i>Klebsiella pneumoniae</i> , <i>Pseudomonas aeruginosa</i>) Drug reactions (single or multiple ulcers)
Pigmented lesions	Kaposi's sarcoma ^a Bacillary angiomatosis (skin and visceral lesions more common than oral) Zidovudine pigmentation (skin, nails, and occasionally oral mucosa) Addison's disease
Miscellaneous	Linear gingival erythema ^a

^aStrongly associated with HIV infection.

major salivary glands secrete intermittently in response to autonomic stimulation, which is high during a meal but low otherwise. Hundreds of minor glands in the lips and cheeks secrete mucus continuously throughout the day and night. Consequently, oral function becomes impaired when salivary function is reduced. The sensation of a dry mouth (*xerostomia*) is perceived when salivary flow is reduced by 50%. The most common etiology is medication, especially drugs with anticholinergic properties but also alpha and beta blockers, calcium channel blockers, and diuretics. Other causes include Sjögren's syndrome, chronic parotitis, salivary duct obstruction, diabetes mellitus, HIV/AIDS, and radiation therapy that includes the salivary glands in the field (e.g., for Hodgkin's disease and for head and neck cancer). Management involves the elimination or limitation of drying medications, preventive dental care, and supplementation with oral liquid or salivary substitutes. Sugarless mints or chewing gum may stimulate salivary secretion if dysfunction is mild. When sufficient exocrine tissue remains, pilocarpine or cevimeline has been shown to increase secretions. Commercial saliva substitutes or gels relieve dryness. Fluoride supplementation is critical to prevent caries.

Sialolithiasis presents most often as painful swelling but in some instances as only swelling or only pain. Conservative therapy consists of local heat, massage, and hydration. Promotion of salivary secretion with mints or lemon drops may flush out small stones. Antibiotic treatment is necessary when bacterial infection is suspected. In adults, *acute bacterial parotitis* is typically unilateral and most commonly affects postoperative, dehydrated, and debilitated patients. *Staphylococcus aureus* (including methicillin-resistant strains) and anaerobic bacteria are the most common pathogens. Chronic bacterial *sialadenitis* results from lowered salivary secretion and recurrent bacterial infection. When suspected bacterial infection is not responsive to therapy, the differential diagnosis should be expanded to include benign and malignant neoplasms, lymphoproliferative disorders, Sjögren's syndrome, sarcoidosis, tuberculosis, lymphadenitis, actinomycosis, and granulomatosis with polyangiitis. Bilateral nontender parotid enlargement occurs with diabetes mellitus, cirrhosis, bulimia, HIV/AIDS, and drugs (e.g., iodide, propylthiouracil).

Pleomorphic adenoma comprises two-thirds of all salivary neoplasms. The parotid is the principal salivary gland affected, and the tumor presents as a firm, slow-growing mass. Although this tumor is benign, its recurrence is common if resection is incomplete. Malignant tumors such as mucoepidermoid carcinoma, adenoid cystic carcinoma, and adenocarcinoma tend to grow relatively fast, depending upon grade. They may ulcerate and invade nerves, producing numbness and facial paralysis. Surgical resection is the primary treatment. Radiation therapy (particularly neutron-beam therapy) is used when surgery is not feasible and as post-resection for certain histologic types with a high risk of recurrence. Malignant salivary gland tumors have a 5-year survival rate of ~68%.

DENTAL CARE FOR MEDICALLY COMPLEX PATIENTS

Routine dental care (e.g., uncomplicated extraction, scaling and cleaning, tooth restoration, and root canal) is remarkably safe. The most common concerns regarding care of dental patients with medical disease are excessive bleeding for patients taking anticoagulants, infection of the heart valves and prosthetic devices from hematogenous seeding by the oral flora, and cardiovascular complications resulting from vasopressors used with local anesthetics during dental treatment. Experience confirms that the risk of any of these complications is very low.

Patients undergoing tooth extraction or alveolar and gingival surgery rarely experience uncontrolled bleeding when warfarin anticoagulation is maintained within the therapeutic range currently recommended for prevention of venous thrombosis, atrial fibrillation, or mechanical heart valve. Embolic complications and death, however, have been reported during subtherapeutic anticoagulation. Therapeutic anticoagulation should be confirmed before and continued through the procedure. Likewise, low-dose aspirin (e.g., 81–325 mg) can safely be continued. For patients taking aspirin and another antiplatelet medication (e.g., clopidogrel), the decision to continue the second antiplatelet

medication should be based on individual consideration of the risks of thrombosis and bleeding.

Patients at risk for bacterial endocarditis (**Chap. 155**) should maintain optimal oral hygiene, including flossing, and have regular professional cleanings. Currently, guidelines recommend that prophylactic antibiotics be restricted to those patients at high risk for bacterial endocarditis who undergo dental and oral procedures involving significant manipulation of gingival or periapical tissue or penetration of the oral mucosa. If unexpected bleeding occurs, antibiotics given within 2 h after the procedure provide effective prophylaxis.

Hematogenous bacterial seeding from oral infection can undoubtedly produce late prosthetic-joint infection and therefore requires removal of the infected tissue (e.g., drainage, extraction, root canal) and appropriate antibiotic therapy. However, evidence that late prosthetic-joint infection follows routine dental procedures is lacking. For this reason, antibiotic prophylaxis is not recommended before dental surgery for patients with orthopedic pins, screws, and plates. Antibiotic prophylaxis is recommended for patients within the first 2 years after joint replacement who have inflammatory arthropathies, immunosuppression, type 1 diabetes mellitus, previous prosthetic-joint infection, hemophilia, or malnourishment.

Concern often arises regarding the use of vasoconstrictors to treat patients with hypertension and heart disease. Vasoconstrictors enhance the depth and duration of local anesthesia, thus reducing the anesthetic dose and potential toxicity. If intravascular injection is avoided, 2% lidocaine with 1:100,000 epinephrine (limited to a total of 0.036 mg of epinephrine) can be used safely in patients with controlled hypertension and stable coronary heart disease, arrhythmia, or congestive heart failure. Precautions should be taken with patients taking tricyclic antidepressants and nonselective beta blockers because these drugs may potentiate the effect of epinephrine.

Elective dental treatments should be postponed for at least 1 month and preferably for 6 months after myocardial infarction, after which the risk of reinfarction is low provided the patient is medically stable (e.g., stable rhythm, stable angina, and no heart failure). Patients who have suffered a stroke should have elective dental care deferred for 6 months. In both situations, effective stress reduction requires good pain control, including the use of the minimal amount of vasoconstrictor necessary to provide good hemostasis and local anesthesia.

Bisphosphonate therapy is associated with *osteonecrosis* of the jaw. However, the risk with oral bisphosphonate therapy is very low. Most patients affected have received high-dose aminobisphosphonate therapy for multiple myeloma or metastatic breast cancer and have undergone tooth extraction or dental surgery. Intraoral lesions, of which two-thirds are painful, appear as exposed yellow-white hard bone involving the mandible or maxilla. Screening tests for determining risk of osteonecrosis are unreliable. Patients slated for aminobisphosphonate therapy should receive preventive dental care that reduces the risk of infection and the need for future dentoalveolar surgery.

HALITOSIS

Halitosis typically emanates from the oral cavity or nasal passages. Volatile sulfur compounds resulting from bacterial decay of food and cellular debris account for the malodor. Periodontal disease, caries, acute forms of gingivitis, poorly fitting dentures, oral abscess, and tongue coating are common causes. Treatment includes correcting poor hygiene, treating infection, and tongue brushing. Hyposalivation can produce and exacerbate halitosis. Pockets of decay in the tonsillar crypts, esophageal diverticulum, esophageal stasis (e.g., achalasia, stricture), sinusitis, and lung abscess account for some instances. A few systemic diseases produce distinctive odors: renal failure (ammoniacal), hepatic (fishy), and ketoacidosis (fruity). *Helicobacter pylori* gastritis can also produce ammoniacal breath. If a patient presents because of concern about halitosis but no odor is detectable, then pseudohalitosis or halitophobia must be considered.

AGING AND ORAL HEALTH

While tooth loss and dental disease are not normal consequences of aging, a complex array of structural and functional changes that occur with age can affect oral health. Subtle changes in tooth structure (e.g., diminished pulp space and volume, sclerosis of dentinal tubules, and altered proportions of nerve and vascular pulp content) result in the elimination or diminution of pain sensitivity and a reduction in the reparative capacity of the teeth. In addition, age-associated fatty replacement of salivary acini may reduce physiologic reserve, thus increasing the risk of hyposalivation. In healthy older adults, there is minimal, if any, reduction in salivary flow.

Poor oral hygiene often results when general health fails or when patients lose manual dexterity and upper-extremity flexibility. This situation is particularly common among frail older adults and nursing home residents and must be emphasized because regular oral cleaning and dental care reduce the incidence of pneumonia and oral disease as well as the mortality risk in this population. Other risks for dental decay include limited lifetime fluoride exposure. Without assiduous care, decay can become quite advanced yet remain asymptomatic. Consequently, much of a tooth—or the entire tooth—can be destroyed before the patient is aware of the process.

Periodontal disease, a leading cause of tooth loss, is indicated by loss of alveolar bone height. More than 90% of the U.S. population has some degree of periodontal disease by age 50. Healthy adults who have not had significant alveolar bone loss by the sixth decade of life do not typically experience significant worsening with advancing age.

Complete edentulousness with advanced age, though less common than in previous decades, still affects <50% of the U.S. population 85 years of age. Speech, mastication, and facial contours are dramatically affected. Edentulousness may also exacerbate obstructive sleep apnea, particularly in asymptomatic individuals who wear dentures. Dentures can improve verbal articulation and restore diminished facial contours. Mastication can also be restored; however, patients expecting dentures to facilitate oral intake are often disappointed. Accommodation to dentures requires a period of adjustment. Pain can result from friction or traumatic lesions produced by loose dentures. Poor fit and poor oral hygiene may permit the development of candidiasis. This fungal infection may be either asymptomatic or painful and is suggested by erythematous smooth or granular tissue conforming to an area covered by the appliance. Individuals with dentures and no natural teeth need regular (annual) professional oral examinations.

The health status of the oral cavity is linked to cardiovascular disease, diabetes, and other systemic illnesses. Thus, examining the oral cavity for signs of disease is a key part of the physical exam. This chapter presents numerous outstanding clinical photographs illustrating many of the conditions discussed in [Chap. 45](#), Oral Manifestations of Disease. Conditions affecting the teeth, periodontal tissues, and oral mucosa are all represented.



FIGURE 46e-1 Gingival overgrowth secondary to calcium channel blocker use.



FIGURE 46e-2 Oral lichen planus.



FIGURE 46e-3 Erosive lichen planus.



FIGURE 46e-4 Stevens-Johnson syndrome—reaction to nevirapine.

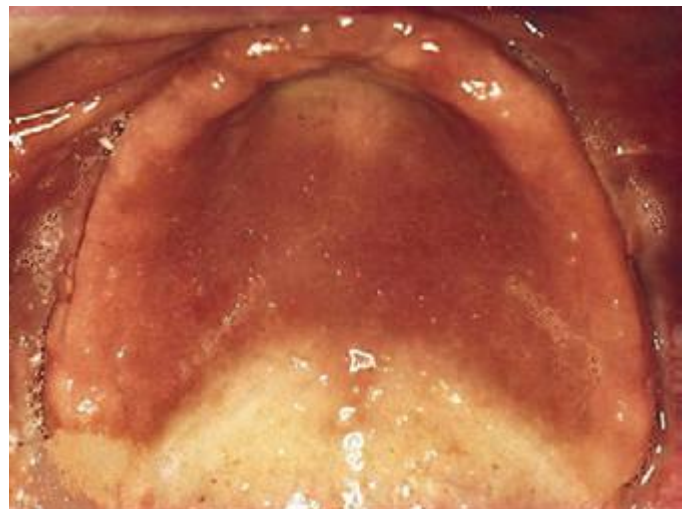


FIGURE 46e-5 Erythematous candidiasis under a denture (i.e., the patient should be treated for this fungal infection).



FIGURE 46e-6 Severe periodontitis.



FIGURE 46e-8 Sublingual leukoplakia.



FIGURE 46e-7 Angular cheilitis.



A



B

FIGURE 46e-9 A. Epulis (gingival hypertrophy) under denture. B. Epulis fissuratum.

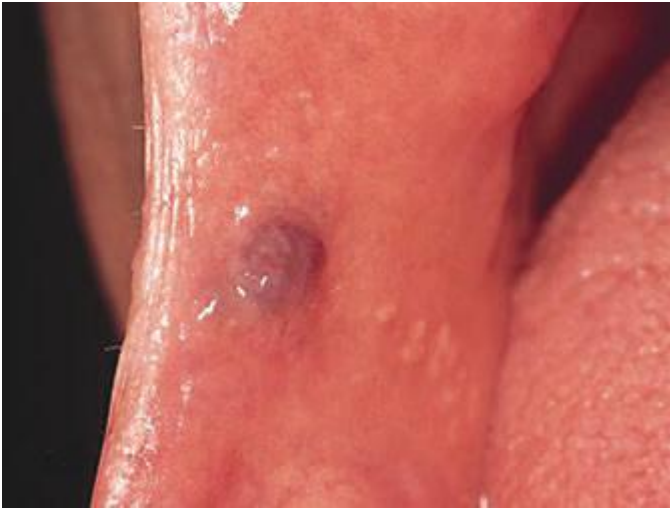


FIGURE 46e-10 Traumatic lesion inside of cheek.



FIGURE 46e-13 Healthy mouth.

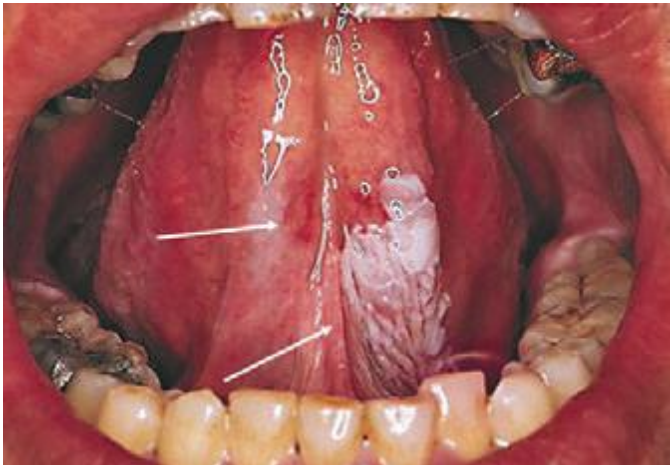


FIGURE 46e-11 Oral leukoplakia, subtype homogenous leukoplakia.



FIGURE 46e-14 Geographic tongue.



FIGURE 46e-12 Oral carcinoma.

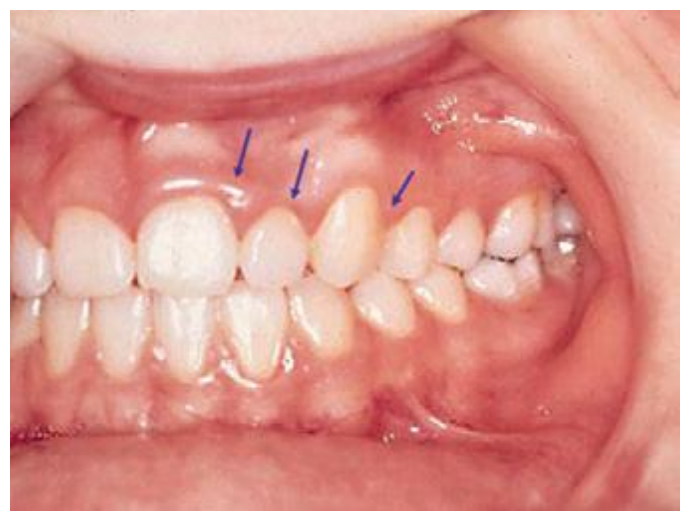


FIGURE 46e-15 Moderate gingivitis.

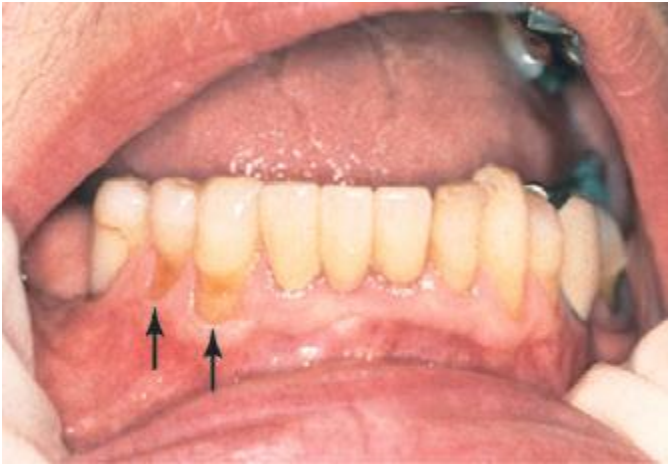


FIGURE 46e-16 Gingival recession.



FIGURE 46e-19 Root cavity in presence of severe periodontal disease.



FIGURE 46e-17 Heavy calculus and gingival inflammation.



FIGURE 46e-20 Ulcer on lateral border of tongue —potential carcinoma.



FIGURE 46e-18 Severe gingival inflammation and heavy calculus.



FIGURE 46e-21 Osteonecrosis.



FIGURE 46e-22 Severe periodontal disease, missing tooth, very mobile teeth.



FIGURE 46e-25 Traumatic ulcer.



FIGURE 46e-23 Salivary stone.



FIGURE 46e-26 Fissured tongue.



FIGURE 46e-27 White coated tongue—likely candidiasis.



FIGURE 46e-24 A. Calculus. B. Teeth cleaned.

ACKNOWLEDGMENT

Dr. Jane Atkinson was a co-author of this chapter in the 17th edition. Some of the materials have been carried over into this edition.

SECTION 5 ALTERATIONS IN CIRCULATORY AND RESPIRATORY FUNCTIONS

47e Dyspnea

Richard M. Schwartzstein

DYSPNEA

The American Thoracic Society defines *dyspnea* as a “subjective experience of breathing discomfort that consists of qualitatively distinct sensations that vary in intensity. The experience derives from interactions among multiple physiological, psychological, social, and environmental factors and may induce secondary physiological and behavioral responses.” Dyspnea, a symptom, can be perceived only by the person experiencing it and must be distinguished from the signs of increased work of breathing.

MECHANISMS OF DYSPNEA

Respiratory sensations are the consequence of interactions between the *efferent*, or outgoing, motor output from the brain to the ventilatory muscles (feed-forward) and the *afferent*, or incoming, sensory input from receptors throughout the body (feedback) as well as the integrative processing of this information that we infer must be occurring in the brain (Fig. 47e-1). In contrast to painful sensations, which can often be attributed to the stimulation of a single nerve ending, dyspnea sensations are more commonly viewed as holistic, more akin to hunger or thirst. A given disease state may lead to dyspnea by one or more mechanisms, some of which may be operative under some circumstances (e.g., exercise) but not others (e.g., a change in position).

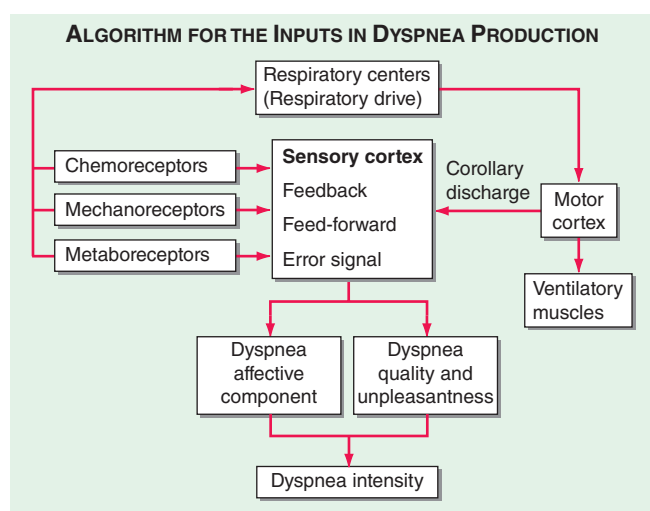


FIGURE 47e-1 Hypothetical model for integration of sensory inputs in the production of dyspnea. Afferent information from the receptors throughout the respiratory system projects directly to the sensory cortex to contribute to primary qualitative sensory experiences and to provide feedback on the action of the ventilatory pump. Afferents also project to the areas of the brain responsible for control of ventilation. The motor cortex, responding to input from the control centers, sends neural messages to the ventilatory muscles and a corollary discharge to the sensory cortex (feed-forward with respect to the instructions sent to the muscles). If the feed-forward and feedback messages do not match, an error signal is generated and the intensity of dyspnea increases. An increasing body of data supports the contribution of affective inputs to the ultimate perception of unpleasant respiratory sensations. (Adapted from MA Gillette, RM Schwartzstein, in SH Ahmedzai, MF Muer [eds]. *Supportive Care in Respiratory Disease*. Oxford, UK, Oxford University Press, 2005.)

Motor Efferents Disorders of the ventilatory pump—most commonly, increased airway resistance or stiffness (decreased compliance) of the respiratory system—are associated with increased work of breathing or the sense of an increased effort to breathe. When the muscles are weak or fatigued, greater effort is required, even though the mechanics of the system are normal. The increased neural output from the motor cortex is sensed via a *corollary discharge*, a neural signal that is sent to the sensory cortex at the same time that motor output is directed to the ventilatory muscles.

Sensory Afferents *Chemoreceptors* in the carotid bodies and medulla are activated by hypoxemia, acute hypercapnia, and acidemia. Stimulation of these receptors and of others that lead to an increase in ventilation produce a sensation of “air hunger.” *Mechanoreceptors* in the lungs, when stimulated by bronchospasm, lead to a sensation of chest tightness. J-receptors, which are sensitive to interstitial edema, and pulmonary vascular receptors, which are activated by acute changes in pulmonary artery pressure, appear to contribute to air hunger. Hyperinflation is associated with the sensation of increased work of breathing, an inability to get a deep breath, or an unsatisfying breath. *Metaboreceptors*, which are located in skeletal muscle, are believed to be activated by changes in the local biochemical milieu of the tissue active during exercise and, when stimulated, contribute to breathing discomfort.

Integration: Efferent-Reafferent Mismatch A discrepancy or mismatch between the feed-forward message to the ventilatory muscles and the feedback from receptors that monitor the response of the ventilatory pump increases the intensity of dyspnea. This mismatch is particularly important when there is a mechanical derangement of the ventilatory pump, as in asthma or chronic obstructive pulmonary disease (COPD).

Contribution of Emotional or Affective Factors to Dyspnea Acute anxiety or fear may increase the severity of dyspnea either by altering the interpretation of sensory data or by leading to patterns of breathing that heighten physiologic abnormalities in the respiratory system. In patients with expiratory flow limitation, for example, the increased respiratory rate that accompanies acute anxiety leads to hyperinflation, increased work and effort of breathing, and the sense of an unsatisfying breath.

ASSESSING DYSPNEA

Quality of Sensation Like pain assessment, dyspnea assessment begins with a determination of the quality of the patient’s discomfort (Table 47e-1). Dyspnea questionnaires or lists of phrases commonly used

TABLE 47e-1 ASSOCIATION OF QUALITATIVE DESCRIPTORS, CLINICAL CHARACTERISTICS, AND PATHOPHYSIOLOGIC MECHANISMS OF SHORTNESS OF BREATH

Descriptor	Clinical Examples	Pathophysiology
Chest tightness or constriction	Asthma, CHF	Bronchoconstriction, interstitial edema
Increased work or effort of breathing	COPD, asthma, neuromuscular disease, chest wall restriction	Airway obstruction, neuromuscular disease
“Air hunger,” need to breathe, urge to breathe	CHF, PE, COPD, asthma, pulmonary fibrosis	Increased drive to breathe
Inability to get a deep breath, unsatisfying breath	Moderate to severe asthma and COPD, pulmonary fibrosis, chest wall disease	Hyperinflation and restricted tidal volume
Heavy breathing, rapid breathing, breathing more	Sedentary status in healthy individual or patient with cardiopulmonary disease	Deconditioning

Abbreviations: CHF, congestive heart failure; COPD, chronic obstructive pulmonary disease; PE, pulmonary embolism.

by patients assist those who have difficulty describing their breathing sensations.

Sensory Intensity A modified Borg scale or visual analogue scale can be utilized to measure dyspnea at rest, immediately following exercise, or on recall of a reproducible physical task, such as climbing the stairs at home. An alternative approach is to gain a sense of the patient's disability by inquiring about what activities are possible. These methods indirectly assess dyspnea and may be affected by nonrespiratory factors, such as leg arthritis or weakness. The Baseline Dyspnea Index and the Chronic Respiratory Disease Questionnaire are commonly used tools for this purpose.

Affective Dimension For a sensation to be reported as a symptom, it must be perceived as unpleasant and interpreted as abnormal. Laboratory studies have demonstrated that air hunger evokes a stronger affective response than does increased effort or work of breathing. Some therapies for dyspnea, such as pulmonary rehabilitation, may reduce breathing discomfort, in part, by altering this dimension.

DIFFERENTIAL DIAGNOSIS

Dyspnea most often results from deviations from normal function in the cardiovascular and respiratory systems. These deviations produce breathlessness as a consequence of increased drive to breathe; increased effort or work of breathing; and/or stimulation of receptors in the heart, lungs, or vascular system. Most diseases of the respiratory system are associated with alterations in the mechanical properties of the lungs and/or chest wall, and some stimulate pulmonary receptors. In contrast, disorders of the cardiovascular system more commonly lead to dyspnea by causing gas-exchange abnormalities or stimulating pulmonary and/or vascular receptors (Table 47e-2).

Respiratory System Dyspnea • DISEASES OF THE AIRWAYS Asthma and COPD, the most common obstructive lung diseases, are characterized by expiratory airflow obstruction, which typically leads to dynamic hyperinflation of the lungs and chest wall. Patients with moderate to severe disease have both increased resistive and elastic loads (a term that relates to the stiffness of the system) on the ventilatory muscles and experience increased work of breathing. Patients with acute bronchoconstriction also report a sense of tightness, which can exist even when lung function is still within the normal range. These patients are commonly tachypneic; this condition leads to hyperinflation and reduced respiratory system compliance and also limits tidal volume. Both the chest tightness and the tachypnea are probably due to stimulation of pulmonary receptors. Both asthma and COPD may lead to hypoxemia and hypercapnia from ventilation-perfusion (\dot{V}/Q) mismatch (and diffusion limitation during exercise with emphysema); hypoxemia is much more common than hypercapnia as a consequence of the different ways in which oxygen and carbon dioxide bind to hemoglobin.

DISEASES OF THE CHEST WALL Conditions that stiffen the chest wall, such as kyphoscoliosis, or that weaken ventilatory muscles, such as myasthenia gravis or the Guillain-Barré syndrome, are also associated with

an increased effort to breathe. Large pleural effusions may contribute to dyspnea, both by increasing the work of breathing and by stimulating pulmonary receptors if there is associated atelectasis.

DISEASES OF THE LUNG PARENCHYMA Interstitial lung diseases, which may arise from infections, occupational exposures, or autoimmune disorders, are associated with increased stiffness (decreased compliance) of the lungs and increased work of breathing. In addition, \dot{V}/Q mismatch and the destruction and/or thickening of the alveolar-capillary interface may lead to hypoxemia and an increased drive to breathe. Stimulation of pulmonary receptors may further enhance the hyper-ventilation characteristic of mild to moderate interstitial disease.

Cardiovascular System Dyspnea • DISEASES OF THE LEFT HEART Diseases of the myocardium resulting from coronary artery disease and nonischemic cardiomyopathies cause a greater left-ventricular end-diastolic volume and an elevation of the left-ventricular end-diastolic as well as pulmonary capillary pressures. These elevated pressures lead to interstitial edema and stimulation of pulmonary receptors, thereby causing dyspnea; hypoxemia due to \dot{V}/Q mismatch may also contribute to breathlessness. Diastolic dysfunction, characterized by a very stiff left ventricle, may lead to severe dyspnea with relatively mild degrees of physical activity, particularly if it is associated with mitral regurgitation.

DISEASES OF THE PULMONARY VASCULATURE Pulmonary thromboembolic disease and primary diseases of the pulmonary circulation (primary pulmonary hypertension, pulmonary vasculitis) cause dyspnea via increased pulmonary-artery pressure and stimulation of pulmonary receptors. Hyperventilation is common, and hypoxemia may be present. However, in most cases, use of supplemental oxygen has only a minimal impact on the severity of dyspnea and hyperventilation.

DISEASES OF THE PERICARDIUM Constrictive pericarditis and cardiac tamponade are both associated with increased intracardiac and pulmonary vascular pressures, which are the likely cause of dyspnea in these conditions. To the extent that cardiac output is limited (at rest or with exercise) metaboreceptors may be stimulated if cardiac output is compromised to the degree that lactic acidosis develops; chemoreceptors will also be activated.

Dyspnea with Normal Respiratory and Cardiovascular Systems Mild to moderate anemia is associated with breathing discomfort during exercise. This symptom is thought to be related to stimulation of metaboreceptors; oxygen saturation is normal in patients with anemia. The breathlessness associated with obesity is probably due to multiple mechanisms, including high cardiac output and impaired ventilatory pump function (decreased compliance of the chest wall). Cardiovascular deconditioning (poor fitness) is characterized by the early development of anaerobic metabolism and the stimulation of chemoreceptors and metaboreceptors. Dyspnea that is medically unexplained has been associated with increased sensitivity to the unpleasantness of acute hypercapnia.

TABLE 47e-2 MECHANISMS OF DYSPNEA IN COMMON DISEASES

Disease	↑ Work of Breathing	↑ Drive to Breathe	Hypoxemia ^a	Acute Hypercapnia ^a	Stimulation of Pulmonary Receptors	Stimulation of Vascular Receptors	Metaboreceptors
COPD	•		•	•			
Asthma	•	•	•	•	•		
ILD	•	•	•	•	•		
PVD		•	•			•	
CPE	•	•	•		•	•	•
NCPE	•	•	•		•		
Anemia							•
Deconditioning							•

^aHypoxemia and hypercapnia are not always present in these conditions. When hypoxemia is present, dyspnea usually persists, albeit at a reduced intensity, with correction of hypoxemia by the administration of supplemental oxygen.

Abbreviations: COPD, chronic obstructive pulmonary disease; CPE, cardiogenic pulmonary edema; ILD, interstitial lung disease; NCPE, noncardiogenic pulmonary edema; PVD, pulmonary vascular disease.

APPROACH TO THE PATIENT:

Dyspnea (see Fig. 47e-2)

HISTORY

The patient should be asked to describe in his/her own words what the discomfort feels like as well as the effect of position, infections, and environmental stimuli on the dyspnea. Orthopnea is a common indicator of congestive heart failure (CHF), mechanical impairment of the diaphragm associated with obesity, or asthma triggered by esophageal reflux. Nocturnal dyspnea suggests CHF or asthma. Acute, intermittent episodes of dyspnea are more likely to reflect episodes of myocardial ischemia, bronchospasm, or pulmonary embolism, while chronic persistent dyspnea is typical of COPD, interstitial lung disease, and chronic thromboembolic disease. Information on risk factors for occupational lung disease and for coronary artery disease should be elicited. Left atrial myxoma or hepatopulmonary syndrome should be considered when the patient complains of *platypnea*—i.e., dyspnea in the upright position with relief in the supine position.

PHYSICAL EXAMINATION

The physical examination should begin during the interview of the patient. Inability of the patient to speak in full sentences before stopping to get a deep breath suggests a condition that leads to stimulation of the controller or impairment of the ventilatory pump with reduced vital capacity. Evidence of increased work of breathing (supraclavicular retractions; use of accessory muscles of ventilation;

and the tripod position, characterized by sitting with the hands braced on the knees) is indicative of increased airway resistance or stiffness of the lungs and the chest wall. When measuring the vital signs, the physician should accurately assess the respiratory rate and measure the pulsus paradoxus (Chap. 288); if the systolic pressure decreases by >10 mmHg, the presence of COPD, acute asthma, or pericardial disease should be considered. During the general examination, signs of anemia (pale conjunctivae), cyanosis, and cirrhosis (spider angiomas, gynecomastia) should be sought. Examination of the chest should focus on symmetry of movement; percussion (dullness is indicative of pleural effusion; hyperresonance is a sign of emphysema); and auscultation (wheezes, rhonchi, prolonged expiratory phase, and diminished breath sounds are clues to disorders of the airways; rales suggest interstitial edema or fibrosis). The cardiac examination should focus on signs of elevated right heart pressures (jugular venous distention, edema, accentuated pulmonic component to the second heart sound); left ventricular dysfunction (S3 and S4 gallops); and valvular disease (murmurs). When examining the abdomen with the patient in the supine position, the physician should note whether there is paradoxical movement of the abdomen: inward motion during inspiration is a sign of diaphragmatic weakness, and rounding of the abdomen during exhalation is suggestive of pulmonary edema. Clubbing of the digits may be an indication of interstitial pulmonary fibrosis, and joint swelling or deformation as well as changes consistent with Raynaud's disease may be indicative of a collagen-vascular process that can be associated with pulmonary disease.

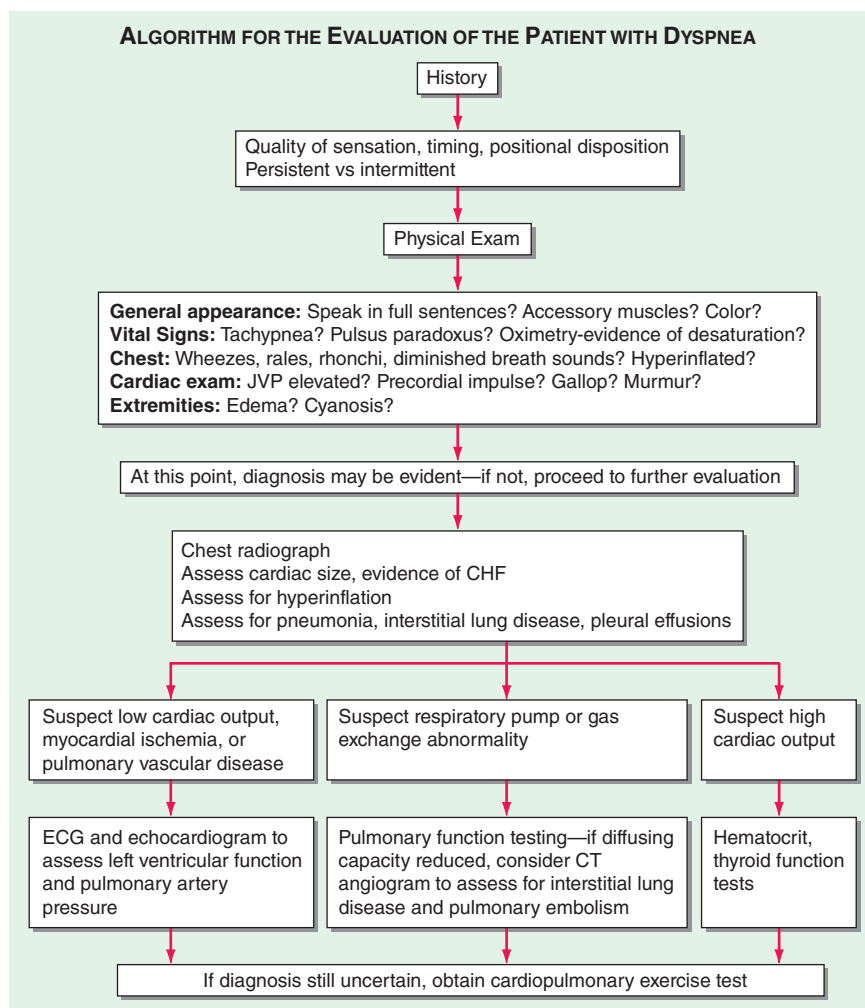


FIGURE 47e-2 Algorithm for the evaluation of the patient with dyspnea. JVP, jugular venous pulse; CHF, congestive heart failure; ECG, electrocardiogram; CT, computed tomography. (Adapted from RM Schwartzstein, D Feller-Kopman, in E Braunwald, L Goldman [eds]. *Primary Cardiology*, 2nd ed. Philadelphia, WB Saunders, 2003.)

Patients with exertional dyspnea should be asked to walk under observation in order to reproduce the symptoms. The patient should be examined during and at the end of exercise for new findings that were not present at rest and for changes in oxygen saturation.

CHEST IMAGING

After the history elicitation and the physical examination, a chest radiograph should be obtained. The lung volumes should be assessed: hyperinflation indicates obstructive lung disease, whereas low lung volumes suggest interstitial edema or fibrosis, diaphragmatic dysfunction, or impaired chest wall motion. The pulmonary parenchyma should be examined for evidence of interstitial disease and emphysema. Prominent pulmonary vasculature in the upper zones indicates pulmonary venous hypertension, while enlarged central pulmonary arteries suggest pulmonary arterial hypertension. An enlarged cardiac silhouette suggests dilated cardiomyopathy or valvular disease. Bilateral pleural effusions are typical of CHF and some forms of collagen-vascular disease. Unilateral effusions raise the specter of carcinoma and pulmonary embolism but may also occur in heart failure. CT of the chest is generally reserved for further evaluation of the lung parenchyma (interstitial lung disease) and possible pulmonary embolism.

LABORATORY STUDIES

Laboratory studies should include electrocardiography to seek evidence of ventricular hypertrophy and prior myocardial infarction. Echocardiography is indicated when systolic dysfunction, pulmonary hypertension, or valvular heart disease is suspected. Bronchoprovocation testing is useful in patients with intermittent symptoms suggestive of asthma but normal physical examination and lung function; up to one-third of patients with the clinical diagnosis of asthma do not have reactive airways disease when formally tested. Measurement of brain natriuretic peptide levels in serum is increasingly used to assess for CHF in patients presenting with acute dyspnea but may be elevated in the presence of right ventricular strain as well.

DISTINGUISHING CARDIOVASCULAR FROM RESPIRATORY SYSTEM DYSPNEA

If a patient has evidence of both pulmonary and cardiac disease, a cardiopulmonary exercise test should be carried out to determine which system is responsible for the exercise limitation. If, at peak exercise, the patient achieves predicted maximal ventilation, demonstrates an increase in dead space or hypoxemia, or develops bronchospasm, the respiratory system is probably the cause of the problem. Alternatively, if the heart rate is >85% of the predicted maximum, if the anaerobic threshold occurs early, if the blood pressure becomes excessively high or decreases during exercise, if the O_2 pulse (O_2 consumption/heart rate, an indicator of stroke volume) falls, or if there are ischemic changes on the electrocardiogram, an abnormality of the cardiovascular system is likely the explanation for the breathing discomfort.

TREATMENT DYSPNEA

The first goal is to correct the underlying problem responsible for the symptom. If this is not possible, an effort is made to lessen the intensity of the symptom and its effect on the patient's quality of life. Supplemental O_2 should be administered if the resting O_2 saturation is $\leq 89\%$ or if the patient's saturation drops to these levels with activity. For patients with COPD, pulmonary rehabilitation programs have demonstrated positive effects on dyspnea, exercise capacity, and rates of hospitalization. Studies of anxiolytics and antidepressants have not documented consistent benefit. Experimental interventions—e.g., cold air on the face, chest wall vibration, and inhaled furosemide—aimed at modulating the afferent information from receptors throughout the respiratory system are being studied. Morphine has been shown to reduce dyspnea out of proportion to the change in ventilation in laboratory models.

PULMONARY EDEMA

MECHANISMS OF FLUID ACCUMULATION

The extent to which fluid accumulates in the interstitium of the lung depends on the balance of hydrostatic and oncotic forces within the pulmonary capillaries and in the surrounding tissue. Hydrostatic pressure favors movement of fluid from the capillary into the interstitium. The oncotic pressure, which is determined by the protein concentration in the blood, favors movement of fluid into the vessel. Levels of albumin, the primary protein in the plasma, may be low in conditions such as cirrhosis and nephrotic syndrome. While hypoalbuminemia favors movement of fluid into the tissue for any given hydrostatic pressure in the capillary, it is usually not sufficient by itself to cause interstitial edema. In a healthy individual, the tight junctions of the capillary endothelium are impermeable to proteins, and the lymphatics in the tissue carry away the small amounts of protein that may leak out; together, these factors result in an oncotic force that maintains fluid in the capillary. Disruption of the endothelial barrier, however, allows protein to escape the capillary bed and enhances the movement of fluid into the tissue of the lung.

CARDIOGENIC PULMONARY EDEMA

(See also Chap. 326) Cardiac abnormalities that lead to an increase in pulmonary venous pressure shift the balance of forces between the capillary and the interstitium. Hydrostatic pressure is increased and fluid exits the capillary at an increased rate, resulting in interstitial and, in more severe cases, alveolar edema. The development of pleural effusions may further compromise respiratory system function and contribute to breathing discomfort.

Early signs of pulmonary edema include exertional dyspnea and orthopnea. Chest radiographs show peribronchial thickening, prominent vascular markings in the upper lung zones, and Kerley B lines. As the pulmonary edema worsens, alveoli fill with fluid; the chest radiograph shows patchy alveolar filling, typically in a perihilar distribution, which then progresses to diffuse alveolar infiltrates. Increasing airway edema is associated with rhonchi and wheezes.

NONCARDIOGENIC PULMONARY EDEMA

In noncardiogenic pulmonary edema, lung water increases due to damage of the pulmonary capillary lining with consequent leakage of proteins and other macromolecules into the tissue; fluid follows the protein as oncotic forces are shifted from the vessel to the surrounding lung tissue. This process is associated with dysfunction of the surfactant lining the alveoli, increased surface forces, and a propensity for the alveoli to collapse at low lung volumes. Physiologically, noncardiogenic pulmonary edema is characterized by intrapulmonary shunt with hypoxemia and decreased pulmonary compliance leading to lower functional residual capacity. On pathologic examination, hyaline membranes are evident in the alveoli, and inflammation leading to pulmonary fibrosis may be seen. Clinically, the picture ranges from mild dyspnea to respiratory failure. Auscultation of the lungs may be relatively normal despite chest radiographs that show diffuse alveolar infiltrates. CT scans demonstrate that the distribution of alveolar edema is more heterogeneous than was once thought. Although normal intracardiac pressures are considered by many to be part of the definition of noncardiogenic pulmonary edema, the pathology of the process, as described above, is distinctly different, and a combination of cardiogenic and noncardiogenic pulmonary edema is observed in some patients.

It is useful to categorize the causes of noncardiogenic pulmonary edema in terms of whether the injury to the lung is likely to result from direct, indirect, or pulmonary vascular causes (Table 47e-3). Direct injuries are mediated via the airways (e.g., aspiration) or as the consequence of blunt chest trauma. Indirect injury is the consequence of mediators that reach the lung via the bloodstream. The third category includes conditions that may result from acute changes in pulmonary vascular pressures, possibly due to sudden autonomic discharge (in the case of neurogenic and high-altitude pulmonary edema) or sudden swings of pleural pressure as well as transient damage to the pulmonary capillaries (in the case of reexpansion pulmonary edema).

TABLE 47e-3 COMMON CAUSES OF NONCARDIOGENIC PULMONARY EDEMA**Direct Injury to Lung**

Chest trauma, pulmonary contusion
 Aspiration
 Smoke inhalation
 Pneumonia
 Oxygen toxicity
 Pulmonary embolism, reperfusion

Hematogenous Injury to Lung

Sepsis
 Pancreatitis
 Nonthoracic trauma
 Leukoagglutination reactions
 Multiple transfusions
 Intravenous drug use (e.g., heroin)
 Cardiopulmonary bypass

Possible Lung Injury Plus Elevated Hydrostatic Pressures

High-altitude pulmonary edema
 Neurogenic pulmonary edema
 Reexpansion pulmonary edema

DISTINGUISHING CARDIOGENIC FROM NONCARDIOGENIC PULMONARY EDEMA

The *history* is essential for assessing the likelihood of underlying cardiac disease as well as for identification of one of the conditions associated with noncardiogenic pulmonary edema. The *physical examination* in cardiogenic pulmonary edema is notable for evidence of increased intracardiac pressures (S3 gallop, elevated jugular venous pulse, peripheral edema) and rales and/or wheezes on auscultation of the chest. In contrast, the physical examination in noncardiogenic pulmonary edema is dominated by the findings of the precipitating condition; pulmonary findings may be relatively normal in the early stages. The *chest radiograph* in cardiogenic pulmonary edema typically shows an enlarged cardiac silhouette, vascular redistribution, interstitial thickening, and perihilar alveolar infiltrates; pleural effusions are common. In noncardiogenic pulmonary edema, heart size is normal, alveolar infiltrates are distributed more uniformly throughout the lungs, and pleural effusions are uncommon. Finally, the *hypoxemia* of cardiogenic pulmonary edema is due largely to \dot{V}/Q mismatch and responds to the administration of supplemental oxygen. In contrast, hypoxemia in noncardiogenic pulmonary edema is due primarily to intrapulmonary shunting and typically persists despite high concentrations of inhaled oxygen.

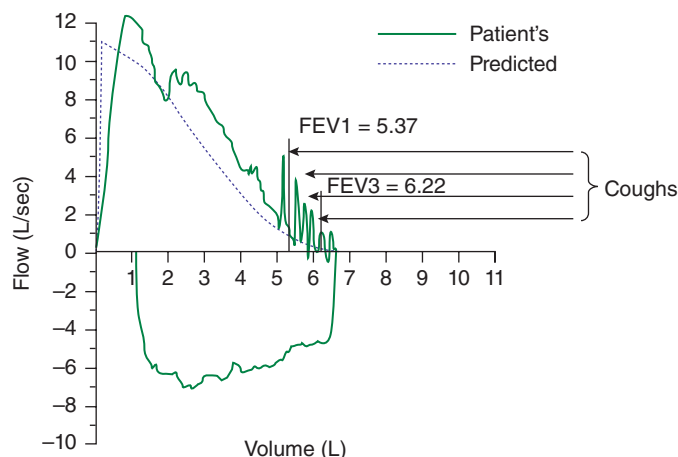


FIGURE 48-1 Flow-volume curve shows spikes of high expiratory flow achieved with cough. FEV₁, forced expiratory volume in 1 s.

exceeding the normal “envelope” of maximal expiratory flow seen on the flow-volume curve (Fig. 48-1). Bronchial smooth-muscle contraction together with dynamic compression of airways narrows airway lumens and maximizes the velocity of exhalation. The kinetic energy available to dislodge mucus from the inside of airway walls is directly proportional to the square of the velocity of expiratory airflow. A deep breath preceding a cough optimizes the function of the expiratory muscles; a series of repetitive coughs at successively lower lung volumes sweeps the point of maximal expiratory velocity progressively further into the lung periphery.

IMPAIRED COUGH

Weak or ineffective cough compromises the ability to clear lower respiratory tract infections, predisposing to more serious infections and their sequelae. Weakness, paralysis, or pain of the expiratory (abdominal and intercostal) muscles is foremost on the list of causes of impaired cough (Table 48-1). Cough strength is generally assessed qualitatively; peak expiratory flow or maximal expiratory pressure at the mouth can be used as a surrogate marker for cough strength. A variety of assistive devices and techniques have been developed to improve cough strength, running the gamut from simple (splinting of the abdominal muscles with a tightly-held pillow to reduce postoperative pain while coughing) to complex (a mechanical cough-assist device supplied via face mask or tracheal tube that applies a cycle of positive pressure followed rapidly by negative pressure). Cough may fail to clear secretions despite a preserved ability to generate normal expiratory velocities; such failure may be due to either abnormal airway secretions (e.g., bronchiectasis due to cystic fibrosis) or structural abnormalities of the airways (e.g., tracheomalacia with expiratory collapse during cough).

SYMPTOMATIC COUGH

The cough of chronic bronchitis in long-term cigarette smokers rarely leads the patient to seek medical advice. It lasts for only seconds to a

48 Cough and Hemoptysis

Patricia A. Kritek, Christopher H. Fanta

COUGH

Cough performs an essential protective function for human airways and lungs. Without an effective cough reflex, we are at risk for retained airway secretions and aspirated material predisposing to infection, atelectasis, and respiratory compromise. At the other extreme, excessive coughing can be exhausting; can be complicated by emesis, syncope, muscular pain, or rib fractures; and can aggravate abdominal or inguinal hernias and urinary incontinence. Cough is often a clue to the presence of respiratory disease. In many instances, cough is an expected and accepted manifestation of disease, as in acute respiratory tract infection. However, persistent cough in the absence of other respiratory symptoms commonly causes patients to seek medical attention.

COUGH MECHANISM

Spontaneous cough is triggered by stimulation of sensory nerve endings that are thought to be primarily rapidly adapting receptors and C fibers. Both chemical (e.g., capsaicin) and mechanical (e.g., particulates in air pollution) stimuli may initiate the cough reflex. A cationic ion channel—the type 1 vanilloid receptor—found on rapidly adapting receptors and C fibers is the receptor for capsaicin, and its expression is increased in patients with chronic cough. Afferent nerve endings richly innervate the pharynx, larynx, and airways to the level of the terminal bronchioles and extend into the lung parenchyma. They may also be located in the external auditory meatus (the auricular branch of the vagus nerve, or the Arnold nerve) and in the esophagus. Sensory signals travel via the vagus and superior laryngeal nerves to a region of the brainstem in the nucleus tractus solitarius vaguely identified as the “cough center.” The cough reflex involves a highly orchestrated series of involuntary muscular actions, with the potential for input from cortical pathways as well. The vocal cords adduct, leading to transient upper-airway occlusion. Expiratory muscles contract, generating positive intrathoracic pressures as high as 300 mmHg. With sudden release of the laryngeal contraction, rapid expiratory flows are generated,

TABLE 48-1 CAUSES OF IMPAIRED COUGH

Decreased expiratory-muscle strength
Decreased inspiratory-muscle strength
Chest wall deformity
Impaired glottic closure or tracheostomy
Tracheomalacia
Abnormal airway secretions
Central respiratory depression (e.g., anesthesia, sedation, or coma)

few minutes, is productive of benign-appearing mucoid sputum, and generally does not cause discomfort. Cough may occur in the context of other respiratory symptoms that together point to a diagnosis; for example, cough accompanied by wheezing, shortness of breath, and chest tightness after exposure to a cat or other sources of allergens suggests asthma. At times, however, cough is the dominant or sole symptom of disease, and it may be of sufficient duration and severity that relief is sought. The duration of cough is a clue to its etiology. Acute cough (<3 weeks) is most commonly due to a respiratory tract infection, aspiration, or inhalation of noxious chemicals or smoke. Subacute cough (3–8 weeks in duration) is a common residuum of tracheobronchitis, as in pertussis or “postviral tussive syndrome.” Chronic cough (>8 weeks) may be caused by a wide variety of cardiopulmonary diseases, including those of inflammatory, infectious, neoplastic, and cardiovascular etiologies. When initial assessment with chest examination and radiography is normal, cough-variant asthma, gastroesophageal reflux, nasopharyngeal drainage, and medications (angiotensin-converting enzyme [ACE] inhibitors) are the most common causes of chronic cough.

ASSESSMENT OF CHRONIC COUGH

Details as to the sound, the time of occurrence during the day, and the pattern of coughing infrequently provide useful etiologic clues. Regardless of cause, cough often worsens upon first lying down at night, with talking, or with the hyperpnea of exercise; it frequently improves with sleep. An exception may involve the cough that occurs only with certain allergic exposures or exercise in cold air, as in asthma. Useful historical questions include what circumstances surround the onset of cough, what makes the cough better or worse, and whether or not the cough produces sputum.

The physical examination seeks clues suggesting the presence of cardiopulmonary disease, including findings such as wheezing or crackles on chest examination. Examination of the auditory canals and tympanic membranes (for irritation of the latter resulting in stimulation of Arnold’s nerve), the nasal passageways (for rhinitis or polyps), and the nails (for clubbing) may also provide etiologic clues. Because cough can be a manifestation of a systemic disease such as sarcoidosis or vasculitis, a thorough general examination is equally important.

In virtually all instances, evaluation of chronic cough merits a chest radiograph. The list of diseases that can cause persistent cough without other symptoms and without detectable abnormalities on physical examination is long. It includes serious illnesses such as sarcoidosis or Hodgkin’s disease in young adults, lung cancer in older patients, and (worldwide) pulmonary tuberculosis. An abnormal chest film prompts an evaluation aimed at explaining the cough. In a patient with chronic productive cough, examination of expectorated sputum is warranted. Purulent-appearing sputum should be sent for routine bacterial culture and, in certain circumstances, mycobacterial culture as well. Cytologic examination of mucoid sputum may be useful to assess for malignancy and to distinguish neutrophilic from eosinophilic bronchitis. Expectoration of blood—whether streaks of blood, blood mixed with airway secretions, or pure blood—deserves a special approach to assessment and management (see “Hemoptysis,” below).

CHRONIC COUGH WITH A NORMAL CHEST RADIOGRAPH

It is commonly held that (alone or in combination) the use of an ACE inhibitor; postnasal drainage; gastroesophageal reflux; and asthma account for more than 90% of cases of chronic cough with a normal or noncontributory chest radiograph. However, clinical experience does not support this contention, and strict adherence to this concept discourages the search for alternative explanations by both clinicians and researchers. ACE inhibitor–induced cough occurs in 5–30% of patients taking these agents and is not dose dependent. ACE metabolizes bradykinin and other tachykinins, such as substance P. The mechanism of ACE inhibitor–associated cough may involve sensitization of sensory nerve endings due to accumulation of bradykinin. In support of this hypothesis, polymorphisms in the neurokinin-2 receptor gene are associated with ACE inhibitor–induced cough. Any patient with

chronic unexplained cough who is taking an ACE inhibitor should have a trial period off the medication, regardless of the timing of the onset of cough relative to the initiation of ACE inhibitor therapy. In most instances, a safe alternative is available; angiotensin-receptor blockers do not cause cough. Failure to observe a decrease in cough after 1 month off medication argues strongly against this etiology. Postnasal drainage of any etiology can cause cough as a response to stimulation of sensory receptors of the cough-reflex pathway in the hypopharynx or aspiration of draining secretions into the trachea. Clues suggesting this etiology include postnasal drip, frequent throat clearing, and sneezing and rhinorrhea. On speculum examination of the nose, excess mucoid or purulent secretions, inflamed and edematous nasal mucosa, and/or polyps may be seen; in addition, secretions or a cobblestoned appearance of the mucosa along the posterior pharyngeal wall may be noted. Unfortunately, there is no means by which to quantitate postnasal drainage. In many instances, this diagnosis must rely on subjective information provided by the patient. This assessment must also be counterbalanced by the fact that many people who have chronic postnasal drainage do not experience cough.

Linking gastroesophageal reflux to chronic cough poses similar challenges. It is thought that reflux of gastric contents into the lower esophagus may trigger cough via reflex pathways initiated in the esophageal mucosa. Reflux to the level of the pharynx (laryngopharyngeal reflux), with consequent aspiration of gastric contents, causes a chemical bronchitis and possibly pneumonitis that can elicit cough for days afterward. Retrosternal burning after meals or on recumbency, frequent eructation, hoarseness, and throat pain may be indicative of gastroesophageal reflux. Nevertheless, reflux may also elicit minimal or no symptoms. Glottic inflammation detected on laryngoscopy may be a manifestation of recurrent reflux to the level of the throat, but it is a nonspecific finding. Quantification of the frequency and level of reflux requires a somewhat invasive procedure to measure esophageal pH directly (either nasopharyngeal placement of a catheter with a pH probe into the esophagus for 24 h or endoscopic placement of a radio-transmitter capsule into the esophagus). The precise interpretation of test results that permits an etiologic linking of reflux events and cough remains debated. Again, assigning the cause of cough to gastroesophageal reflux must be weighed against the observation that many people with symptomatic reflux do not experience chronic cough.

Cough alone as a manifestation of asthma is common among children but not among adults. Cough due to asthma in the absence of wheezing, shortness of breath, and chest tightness is referred to as “cough-variant asthma.” A history suggestive of cough-variant asthma ties the onset of cough to exposure to typical triggers for asthma and the resolution of cough to discontinuation of exposure. Objective testing can establish the diagnosis of asthma (airflow obstruction on spirometry that varies over time or reverses in response to a bronchodilator) or exclude it with certainty (a negative response to a bronchoprovocation challenge—e.g., with methacholine). In a patient capable of taking reliable measurements, home expiratory peak flow monitoring can be a cost-effective method to support or discount a diagnosis of asthma.

Chronic eosinophilic bronchitis causes chronic cough with a normal chest radiograph. This condition is characterized by sputum eosinophilia in excess of 3% without airflow obstruction or bronchial hyperresponsiveness and is successfully treated with inhaled glucocorticoids.

Treatment of chronic cough in a patient with a normal chest radiograph is often empirical and is targeted at the most likely cause(s) of cough as determined by history, physical examination, and possibly pulmonary-function testing. Therapy for postnasal drainage depends on the presumed etiology (infection, allergy, or vasomotor rhinitis) and may include systemic antihistamines; antibiotics; nasal saline irrigation; and nasal pump sprays with glucocorticoids, antihistamines, or anticholinergics. Antacids, histamine type 2 (H₂) receptor antagonists, and proton-pump inhibitors are used to neutralize or decrease the production of gastric acid in gastroesophageal reflux disease; dietary changes, elevation of the head and torso during sleep, and medications to improve gastric emptying are additional therapeutic measures.

Cough-variant asthma typically responds well to inhaled glucocorticoids and intermittent use of inhaled β -agonist bronchodilators.

Patients who fail to respond to treatment targeting the common causes of chronic cough or who have had these causes excluded by appropriate diagnostic testing should undergo chest CT. Diseases causing cough that may be missed on chest x-ray include tumors, early interstitial lung disease, bronchiectasis, and atypical mycobacterial pulmonary infection. On the other hand, patients with chronic cough who have normal findings on chest examination, lung function testing, oxygenation assessment, and chest CT can be reassured as to the absence of serious pulmonary pathology.

SYMPTOM-BASED TREATMENT OF COUGH

Chronic idiopathic cough, also called *cough hypersensitivity syndrome*, is distressingly common. It is often experienced as a tickle or sensitivity in the throat, occurs more often in women, and is typically “dry” or at most productive of scant amounts of mucoid sputum. It can be exhausting, interfere with work, and cause social embarrassment. Once serious underlying cardiopulmonary pathology has been excluded, an attempt at cough suppression is appropriate. Most effective are narcotic cough suppressants, such as codeine or hydrocodone, which are thought to act in the “cough center” in the brainstem. The tendency of narcotic cough suppressants to cause drowsiness and constipation and their potential for addictive dependence limit their appeal for long-term use. *Dextromethorphan* is an over-the-counter, centrally acting cough suppressant with fewer side effects and less efficacy than the narcotic cough suppressants. *Dextromethorphan* is thought to have a different site of action than narcotic cough suppressants and can be used in combination with them if necessary. *Benzonate* is thought to inhibit neural activity of sensory nerves in the cough-reflex pathway. It is generally free of side effects; however, its effectiveness in suppressing cough is variable and unpredictable. Case series have reported benefit from off-label use of *gabapentin* or *amitriptyline* for chronic idiopathic cough. Novel cough suppressants without the limitations of currently available agents are greatly needed. Approaches that are being explored include the development of neurokinin receptor antagonists, type 1 vanilloid receptor antagonists, and novel opioid and opioid-like receptor agonists.

HEMOPTYSIS

Hemoptysis, the expectoration of blood from the respiratory tract, can arise at any location from the alveoli to the glottis. It is important to distinguish hemoptysis from *epistaxis* (bleeding from the nasopharynx) and *hematemesis* (bleeding from the upper gastrointestinal tract). Hemoptysis can range from the expectoration of blood-tinged sputum to that of life-threatening large volumes of bright red blood. For most patients, any degree of hemoptysis can cause anxiety and often prompts medical evaluation.

While precise epidemiologic data are lacking, the most common etiology of hemoptysis is infection of the medium-sized airways. In the United States, the cause is usually viral or bacterial bronchitis. Hemoptysis can arise in the setting of acute bronchitis or during an exacerbation of chronic bronchitis. Worldwide, the most common cause of hemoptysis is infection with *Mycobacterium tuberculosis*, presumably because of the high prevalence of tuberculosis and its predilection for cavity formation. While these are the most common causes, the differential diagnosis for hemoptysis is extensive, and a step-wise approach to evaluation is appropriate.

ETIOLOGY

One way to approach the source of hemoptysis is to search systematically for potential sites of bleeding from the alveolus to the mouth. Diffuse bleeding in the alveolar space, often referred to as *diffuse alveolar hemorrhage* (DAH), may present as hemoptysis. Causes of DAH can be inflammatory or noninflammatory. Inflammatory DAH is due to small-vessel vasculitis/capillaritis from a variety of diseases, including granulomatosis with polyangiitis and microscopic polyangiitis. Similarly,

systemic autoimmune diseases such as systemic lupus erythematosus can manifest as pulmonary capillaritis. Antibodies to the alveolar basement membrane, as are seen in Goodpasture’s disease, can also result in alveolar hemorrhage. In the early period after bone marrow transplantation, patients can develop a form of inflammatory DAH that can be catastrophic and life-threatening. The exact pathophysiology of this process is not well understood, but DAH should be suspected in patients with sudden-onset dyspnea and hypoxemia in the first 100 days after bone marrow transplantation.

Alveoli can also bleed due to direct inhalational injury, including thermal injury from fires, inhalation of illicit substances (e.g., cocaine), and inhalation of toxic chemicals. If alveoli are irritated from any process, patients with thrombocytopenia, coagulopathy, or antiplatelet or anticoagulant use will be at increased risk of hemoptysis.

Bleeding in hemoptysis most commonly arises from the small- to medium-sized airways. Irritation and injury of the bronchial mucosa can lead to small-volume bleeding. More significant hemoptysis can result from the proximity of the bronchial artery and vein to the airway, with these vessels and the bronchus running together in what is often referred to as the *bronchovascular bundle*. In the smaller airways, these blood vessels are close to the airspace, and lesser degrees of inflammation or injury can therefore result in their rupture into the airways. While alveolar hemorrhage arises from capillaries that are part of the low-pressure pulmonary circulation, bronchial bleeding generally originates from bronchial arteries, which are under systemic pressure and thus are predisposed to larger-volume bleeding.

Any infection of the airways can result in hemoptysis, although acute bronchitis is most commonly caused by viral infection. In patients with a history of chronic bronchitis, bacterial superinfection with organisms such as *Streptococcus pneumoniae*, *Haemophilus influenzae*, or *Moraxella catarrhalis* can also result in hemoptysis. Patients with bronchiectasis (a permanent dilation of the airways with loss of mucosal integrity) are particularly prone to hemoptysis due to chronic inflammation and anatomic abnormalities that bring the bronchial arteries closer to the mucosal surface. One common presentation of patients with advanced cystic fibrosis—the prototypical bronchiectatic lung disease—is hemoptysis, which can be life-threatening.

Pneumonias of any sort can cause hemoptysis. Tuberculous infection, which can lead to bronchiectasis or cavitary pneumonia, is a very common cause of hemoptysis worldwide. Patients may present with a chronic cough productive of blood-streaked sputum or with larger-volume bleeding. *Rasmussen’s aneurysm* (the dilation of a pulmonary artery in a cavity formed by previous tuberculous infection) remains a source of massive, life-threatening hemoptysis in the developing world. Community-acquired pneumonia and lung abscess can also result in bleeding. Once again, if the infection results in cavitation, there is a greater likelihood of bleeding due to erosion into blood vessels. Infections with *Staphylococcus aureus* and gram-negative rods (e.g., *Klebsiella pneumoniae*) are especially likely to cause necrotizing lung infections and thus to be associated with hemoptysis.



While not common in North America, pulmonary paragonimiasis (i.e., infection with the lung fluke *Paragonimus westermani*) often presents as fever, cough, and hemoptysis. This infection is a public health issue in Southeast Asia and China and is frequently confused with active tuberculosis, in which the clinical picture can be similar. Paragonimiasis should be considered in recent immigrants from endemic areas who have new or recurrent hemoptysis. In addition, pulmonary paragonimiasis has been reported secondary to ingestion of crayfish or small crabs in the United States.

Other causes of airway irritation resulting in hemoptysis include inhalation of toxic chemicals, thermal injury, and direct trauma from suctioning of the airways (particularly in intubated patients). All of these etiologies should be considered in light of the individual patient’s history and exposures.

Perhaps the most feared cause of hemoptysis is bronchogenic lung cancer, although hemoptysis is a presenting symptom in only ~10% of patients. Cancers arising in the proximal airways are much more likely to cause hemoptysis, but any malignancy in the chest can do so. Because both squamous cell carcinomas and small-cell carcinomas are

more commonly in or adjacent to the proximal airways, and large at presentation, they are more often a cause of hemoptysis. These cancers can present with large-volume and life-threatening hemoptysis because of erosion into the hilar vessels. Carcinoid tumors, which are found almost exclusively as endobronchial lesions with friable mucosa, can also present with hemoptysis.

In addition to cancers arising in the lung, metastatic disease in the pulmonary parenchyma can bleed. Malignancies that commonly metastasize to the lungs include renal cell, breast, colon, testicular, and thyroid cancers as well as melanoma. While hemoptysis is not a common manifestation of pulmonary metastases, the combination of multiple pulmonary nodules and hemoptysis should raise suspicion of this etiology. Finally, disease of the pulmonary vasculature can cause hemoptysis. Perhaps most frequently, congestive heart failure with transmission of elevated left atrial pressures can lead to rupture of small alveolar capillaries. These patients rarely present with bright red blood but more commonly have pink, frothy sputum or blood-tinged secretions. Patients with a focal jet of mitral regurgitation can present with an upper-lobe opacity on chest radiography together with hemoptysis. This finding is thought to be due to focal increases in pulmonary capillary pressure due to the regurgitant jet. Pulmonary arteriovenous malformations are prone to bleeding. Pulmonary embolism can also lead to the development of hemoptysis, which is generally associated with pulmonary infarction. Pulmonary arterial hypertension from other causes rarely results in hemoptysis.

EVALUATION

As with most signs of possible illness, the initial step in the evaluation of hemoptysis is a thorough history and physical examination (Fig. 48-2). As already mentioned, initial questioning should focus on ascertaining whether the bleeding is truly from the respiratory tract and not the nasopharynx or gastrointestinal tract; bleeding from the latter sources requires different approaches to evaluation and treatment.

History and Physical Examination The specific characteristics of hemoptysis may be helpful in determining an etiology, such as whether the expectorated material consists of blood-tinged, purulent secretions; pink, frothy sputum; or pure blood. Information on specific triggers

of the bleeding (e.g., recent inhalation exposures) as well as any previous episodes of hemoptysis should be elicited during history-taking. Monthly hemoptysis in a woman suggests catamenial hemoptysis from pulmonary endometriosis. Moreover, the volume of blood expectorated is important not only in determining the cause but also in gauging the urgency for further diagnostic and therapeutic maneuvers. Patients rarely exsanguinate from hemoptysis but can effectively “drown” in aspirated blood. Large-volume hemoptysis, referred to as *massive hemoptysis*, is variably defined as hemoptysis of >200–600 mL in 24 h. Massive hemoptysis should be considered a medical emergency. All patients should be asked about current or former cigarette smoking; this behavior predisposes to chronic bronchitis and increases the likelihood of bronchogenic cancer. Practitioners should inquire about symptoms and signs suggestive of respiratory tract infection (including fever, chills, and dyspnea), recent inhalation exposures, recent use of illicit substances, and risk factors for venous thromboembolism.

A medical history of malignancy or treatment thereof, rheumatologic disease, vascular disease, or underlying lung disease (e.g., bronchiectasis) may be relevant to the cause of hemoptysis. Because many causes of DAH can be part of a pulmonary-renal syndrome, specific inquiry into a history of renal insufficiency is important.

The physical examination begins with an assessment of vital signs and oxygen saturation to gauge whether there is evidence of life-threatening bleeding. Tachycardia, hypotension, and decreased oxygen saturation mandate a more expedited evaluation of hemoptysis. A specific focus on respiratory and cardiac examinations is important; these examinations should include inspection of the nares, auscultation of the lungs and heart, assessment of the lower extremities for symmetric or asymmetric edema, and evaluation for jugular venous distention. Clubbing of the digits may suggest underlying lung diseases such as bronchogenic carcinoma or bronchiectasis, which predispose to hemoptysis. Similarly, mucocutaneous telangiectasias should raise the specter of pulmonary arterial-venous malformations.

Diagnostic Evaluation For most patients, the next step in evaluation of hemoptysis should be a standard chest radiograph. If a source of bleeding is not identified on plain film, CT of the chest should be performed. CT allows better delineation of bronchiectasis, alveolar filling, cavitory

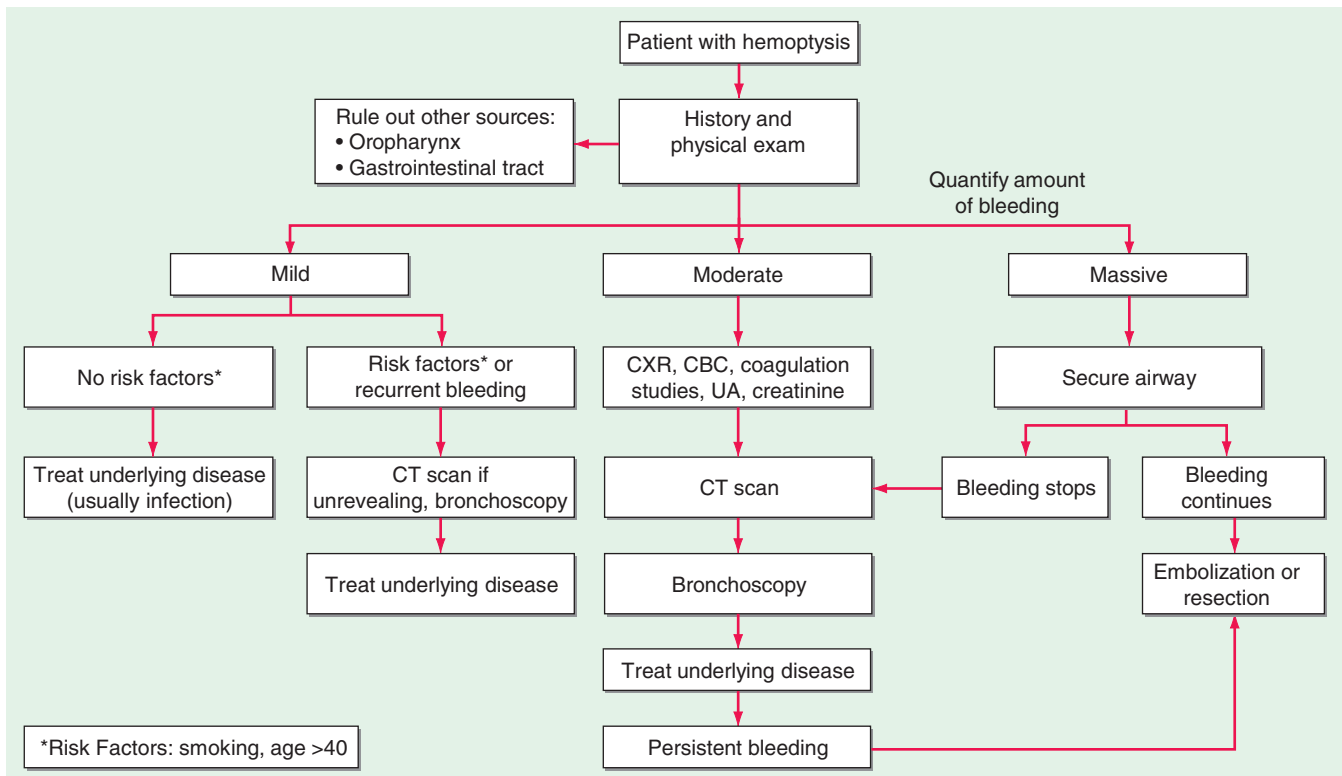


FIGURE 48-2 Decision tree for evaluation of hemoptysis. CBC, complete blood count; CT, computed tomography; CXR, chest x-ray; UA, urinalysis.

infiltrates, and masses than does chest radiograph. The practitioner should consider a CT protocol to assess for pulmonary embolism if the history or examination suggests venous thromboembolism as a cause of bleeding.

Laboratory studies should include a complete blood count to assess both the hematocrit and the platelet count as well as coagulation studies. Renal function should be evaluated and urinalysis conducted because of the possibility of pulmonary-renal syndromes presenting with hemoptysis. The documentation of acute renal insufficiency or the detection of red blood cells or their casts on urinalysis should elevate suspicion of small-vessel vasculitis, and studies such as anti-neutrophil cytoplasmic antibody, antiglomerular basement membrane antibody, and antinuclear antibody should be considered. If a patient is producing sputum, Gram's and acid-fast staining as well as culture should be undertaken.

If all of these studies are unrevealing, bronchoscopy should be considered. In any patient with a history of cigarette smoking, airway inspection should be part of the evaluation of new-onset hemoptysis as endobronchial lesions are not reliably visualized on CT.

TREATMENT HEMOPTYSIS

For the most part, the treatment of hemoptysis varies with its etiology. However, large-volume, life-threatening hemoptysis generally requires immediate intervention regardless of the cause. The first step is to establish a patent airway, usually by endotracheal intubation and subsequent mechanical ventilation. As large-volume hemoptysis usually arises from an airway lesion, it is ideal to identify the site of bleeding by either chest imaging or bronchoscopy (more commonly rigid rather than flexible). The goals are then to isolate the bleeding to one lung and not to allow the preserved airspaces in the other lung to be filled with blood so that gas exchange is further impaired. Patients should be placed with the bleeding lung in a dependent position (i.e., bleeding-side down), and, if possible, dual-lumen endotracheal tubes or an airway blocker should be placed in the proximal airway of the bleeding lung. These interventions generally require the assistance of anesthesiologists, interventional pulmonologists, or thoracic surgeons.

If the bleeding does not stop with treatment of the underlying cause and the passage of time, severe hemoptysis from bronchial arteries can be treated with angiographic embolization of the responsible bronchial artery. This intervention should be entertained only in the most severe and life-threatening cases of hemoptysis because of the risk of unintentional spinal-artery embolization and consequent paraplegia. Endobronchial lesions can be treated with a variety of bronchoscopically directed interventions, including cauterization and laser therapy. In extreme circumstances, surgical resection of the affected region of the lung is considered. Most cases of hemoptysis resolve with treatment of the infection or inflammatory process or with removal of the offending stimulus.

RESPONSES TO HYPOXIA

Decreased O_2 availability to cells results in an inhibition of oxidative phosphorylation and increased anaerobic glycolysis. This switch from aerobic to anaerobic metabolism, the Pasteur effect, maintains some, albeit reduced, adenosine 5'-triphosphate (ATP) production. In severe hypoxia, when ATP production is inadequate to meet the energy requirements of ionic and osmotic equilibrium, cell membrane depolarization leads to uncontrolled Ca^{2+} influx and activation of Ca^{2+} -dependent phospholipases and proteases. These events, in turn, cause cell swelling, activation of apoptotic pathways, and, ultimately, cell death.

The adaptations to hypoxia are mediated, in part, by the upregulation of genes encoding a variety of proteins, including glycolytic enzymes, such as phosphoglycerate kinase and phosphofructokinase, as well as the glucose transporters Glut-1 and Glut-2; and by growth factors, such as vascular endothelial growth factor (VEGF) and erythropoietin, which enhance erythrocyte production. The hypoxia-induced increase in expression of these key proteins is governed by the hypoxia-sensitive transcription factor, hypoxia-inducible factor-1 (HIF-1).

During hypoxia, systemic arterioles dilate, at least in part, by opening of K_{ATP} channels in vascular smooth-muscle cells due to the hypoxia-induced reduction in ATP concentration. By contrast, in pulmonary vascular smooth-muscle cells, inhibition of K^+ channels causes depolarization which, in turn, activates voltage-gated Ca^{2+} channels raising the cytosolic $[Ca^{2+}]$ and causing smooth-muscle cell contraction. Hypoxia-induced pulmonary arterial constriction shunts blood away from poorly ventilated portions toward better ventilated portions of the lung; however, it also increases pulmonary vascular resistance and right ventricular afterload.

Effects on the Central Nervous System Changes in the central nervous system (CNS), particularly the higher centers, are especially important consequences of hypoxia. Acute hypoxia causes impaired judgment, motor incoordination, and a clinical picture resembling acute alcohol intoxication. High-altitude illness is characterized by headache secondary to cerebral vasodilation, gastrointestinal symptoms, dizziness, insomnia, fatigue, or somnolence. Pulmonary arterial and sometimes venous constriction causes capillary leakage and high-altitude pulmonary edema (HAPE) (Chap. 47e), which intensifies hypoxia, further promoting vasoconstriction. Rarely, high-altitude cerebral edema (HACE) develops, which is manifest by severe headache and papilledema and can cause coma. As hypoxia becomes more severe, the regulatory centers of the brainstem are affected, and death usually results from respiratory failure.

Effects on the Cardiovascular System Acute hypoxia stimulates the chemoreceptor reflex arc to induce venoconstriction and systemic arterial vasodilation. These acute changes are accompanied by transiently increased myocardial contractility, which is followed by depressed myocardial contractility with prolonged hypoxia.

CAUSES OF HYPOXIA

Respiratory Hypoxia When hypoxia occurs from respiratory failure, P_{aO_2} declines, and when respiratory failure is persistent, the hemoglobin-oxygen ($Hb-O_2$) dissociation curve (see Fig. 127-2) is displaced to the right, with greater quantities of O_2 released at any level of tissue P_{O_2} . Arterial hypoxemia, i.e., a reduction of O_2 saturation of arterial blood (S_{aO_2}), and consequent cyanosis are likely to be more marked when such depression of P_{aO_2} results from pulmonary disease than when the depression occurs as the result of a decline in the fraction of oxygen in inspired air (F_{iO_2}). In this latter situation, P_{aCO_2} falls secondary to anoxia-induced hyperventilation and the $Hb-O_2$ dissociation curve is displaced to the left, limiting the decline in S_{aO_2} at any level of P_{aO_2} .

The most common cause of respiratory hypoxia is *ventilation-perfusion mismatch* resulting from perfusion of poorly ventilated alveoli. Respiratory hypoxemia may also be caused by *hypoventilation*, in which case it is associated with an elevation of P_{aCO_2} (Chap. 306e). These two forms of respiratory hypoxia are usually correctable by

49 Hypoxia and Cyanosis

Joseph Loscalzo

HYPOXIA

The fundamental purpose of the cardiorespiratory system is to deliver O_2 and nutrients to cells and to remove CO_2 and other metabolic products from them. Proper maintenance of this function depends not only on intact cardiovascular and respiratory systems, but also on an adequate number of red blood cells and hemoglobin and a supply of inspired gas containing adequate O_2 .

inspiring 100% O₂ for several minutes. A third cause of respiratory hypoxia is shunting of blood across the lung from the pulmonary arterial to the venous bed (*intrapulmonary right-to-left shunting*) by perfusion of nonventilated portions of the lung, as in pulmonary atelectasis or through pulmonary arteriovenous connections. The low PaO₂ in this situation is only partially corrected by an F_{IO}₂ of 100%.

Hypoxia Secondary to High Altitude As one ascends rapidly to 3000 m (~10,000 ft), the reduction of the O₂ content of inspired air (F_{IO}₂) leads to a decrease in alveolar PO₂ to approximately 60 mmHg, and a condition termed *high-altitude illness* develops (see above). At higher altitudes, arterial saturation declines rapidly and symptoms become more serious; and at 5000 m, unacclimated individuals usually cease to be able to function normally owing to the changes in CNS function described above.

Hypoxia Secondary to Right-to-Left Extrapulmonary Shunting From a physiologic viewpoint, this cause of hypoxia resembles intrapulmonary right-to-left shunting but is caused by congenital cardiac malformations, such as tetralogy of Fallot, transposition of the great arteries, and Eisenmenger's syndrome (Chap. 282). As in pulmonary right-to-left shunting, the PaO₂ cannot be restored to normal with inspiration of 100% O₂.

Anemic Hypoxia A reduction in hemoglobin concentration of the blood is accompanied by a corresponding decline in the O₂-carrying capacity of the blood. Although the PaO₂ is normal in anemic hypoxia, the absolute quantity of O₂ transported per unit volume of blood is diminished. As the anemic blood passes through the capillaries and the usual quantity of O₂ is removed from it, the PO₂ and saturation in the venous blood decline to a greater extent than normal.

Carbon Monoxide (CO) Intoxication (See also Chap. 472e) Hemoglobin that binds with CO (carboxy-hemoglobin, COHb) is unavailable for O₂ transport. In addition, the presence of COHb shifts the Hb-O₂ dissociation curve to the left (see Fig. 127-2) so that O₂ is unloaded only at lower tensions, further contributing to tissue hypoxia.

Circulatory Hypoxia As in anemic hypoxia, the PaO₂ is usually normal, but venous and tissue PO₂ values are reduced as a consequence of reduced tissue perfusion and greater tissue O₂ extraction. This pathophysiology leads to an increased arterial-mixed venous O₂ difference (a-v-O₂ difference), or gradient. Generalized circulatory hypoxia occurs in heart failure (Chap. 279) and in most forms of shock (Chap. 324).

Specific Organ Hypoxia Localized circulatory hypoxia may occur as a result of decreased perfusion secondary to arterial obstruction, as in localized atherosclerosis in any vascular bed, or as a consequence of vasoconstriction, as observed in Raynaud's phenomenon (Chap. 302). Localized hypoxia may also result from venous obstruction and the resultant expansion of interstitial fluid causing arteriolar compression and, thereby, reduction of arterial inflow. Edema, which increases the distance through which O₂ must diffuse before it reaches cells, can also cause localized hypoxia. In an attempt to maintain adequate perfusion to more vital organs in patients with reduced cardiac output secondary to heart failure or hypovolemic shock, vasoconstriction may reduce perfusion in the limbs and skin, causing hypoxia of these regions.

Increased O₂ Requirements If the O₂ consumption of tissues is elevated without a corresponding increase in perfusion, tissue hypoxia ensues and the PO₂ in venous blood declines. Ordinarily, the clinical picture of patients with hypoxia due to an elevated metabolic rate, as in fever or thyrotoxicosis, is quite different from that in other types of hypoxia: the skin is warm and flushed owing to increased cutaneous blood flow that dissipates the excessive heat produced, and cyanosis is usually absent.

Exercise is a classic example of increased tissue O₂ requirements. These increased demands are normally met by several mechanisms operating simultaneously: (1) increase in the cardiac output and ventilation and, thus, O₂ delivery to the tissues; (2) a preferential shift in blood flow to the exercising muscles by changing vascular resistances

in the circulatory beds of exercising tissues, directly and/or reflexly; (3) an increase in O₂ extraction from the delivered blood and a widening of the arteriovenous O₂ difference; and (4) a reduction in the pH of the tissues and capillary blood, shifting the Hb-O₂ curve to the right (see Fig. 127-2), and unloading more O₂ from hemoglobin. If the capacity of these mechanisms is exceeded, then hypoxia, especially of the exercising muscles, will result.

Improper Oxygen Utilization Cyanide (Chap. 473e) and several other similarly acting poisons cause cellular hypoxia. The tissues are unable to use O₂, and, as a consequence, the venous blood tends to have a high O₂ tension. This condition has been termed *histotoxic hypoxia*.

ADAPTATION TO HYPOXIA

An important component of the respiratory response to hypoxia originates in special chemosensitive cells in the carotid and aortic bodies and in the respiratory center in the brainstem. The stimulation of these cells by hypoxia increases ventilation, with a loss of CO₂, and can lead to respiratory alkalosis. When combined with the metabolic acidosis resulting from the production of lactic acid, the serum bicarbonate level declines (Chap. 66).

With the reduction of PaO₂, cerebrovascular resistance decreases and cerebral blood flow increases in an attempt to maintain O₂ delivery to the brain. However, when the reduction of PaO₂ is accompanied by hyperventilation and a reduction of PaCO₂, cerebrovascular resistance rises, cerebral blood flow falls, and tissue hypoxia intensifies.

The diffuse, systemic vasodilation that occurs in generalized hypoxia increases the cardiac output. In patients with underlying heart disease, the requirements of peripheral tissues for an increase of cardiac output with hypoxia may precipitate congestive heart failure. In patients with ischemic heart disease, a reduced PaO₂ may intensify myocardial ischemia and further impair left ventricular function.

One of the important compensatory mechanisms for chronic hypoxia is an increase in the hemoglobin concentration and in the number of red blood cells in the circulating blood, i.e., the development of polycythemia secondary to erythropoietin production (Chap. 131). In persons with chronic hypoxemia secondary to prolonged residence at a high altitude (>13,000 ft, 4200 m), a condition termed *chronic mountain sickness* develops. This disorder is characterized by a blunted respiratory drive, reduced ventilation, erythrocytosis, cyanosis, weakness, right ventricular enlargement secondary to pulmonary hypertension, and even stupor.

CYANOSIS

Cyanosis refers to a bluish color of the skin and mucous membranes resulting from an increased quantity of reduced hemoglobin (i.e., deoxygenated hemoglobin) or of hemoglobin derivatives (e.g., methemoglobin or sulfhemoglobin) in the small blood vessels of those tissues. It is usually most marked in the lips, nail beds, ears, and malar eminences. Cyanosis, especially if developed recently, is more commonly detected by a family member than the patient. The florid skin characteristic of polycythemia vera (Chap. 131) must be distinguished from the true cyanosis discussed here. A cherry-colored flush, rather than cyanosis, is caused by COHb (Chap. 473e).

The degree of cyanosis is modified by the color of the cutaneous pigment and the thickness of the skin, as well as by the state of the cutaneous capillaries. The accurate clinical detection of the presence and degree of cyanosis is difficult, as proved by oximetric studies. In some instances, central cyanosis can be detected reliably when the Sao₂ has fallen to 85%; in others, particularly in dark-skinned persons, it may not be detected until it has declined to 75%. In the latter case, examination of the mucous membranes in the oral cavity and the conjunctivae rather than examination of the skin is more helpful in the detection of cyanosis.

The increase in the quantity of reduced hemoglobin in the mucocutaneous vessels that produces cyanosis may be brought about either by an increase in the quantity of venous blood as a result of dilation of the venules (including precapillary venules) or by a reduction in the Sao₂ in the capillary blood. In general, cyanosis becomes apparent when

the concentration of reduced hemoglobin in capillary blood exceeds 40 g/L (4 g/dL).

It is the *absolute*, rather than the *relative*, quantity of reduced hemoglobin that is important in producing cyanosis. Thus, in a patient with severe anemia, the *relative* quantity of reduced hemoglobin in the venous blood may be very large when considered in relation to the total quantity of hemoglobin in the blood. However, since the concentration of the latter is markedly reduced, the *absolute* quantity of reduced hemoglobin may still be low, and, therefore, patients with severe anemia and even *marked* arterial desaturation may not display cyanosis. Conversely, the higher the total hemoglobin content, the greater the tendency toward cyanosis; thus, patients with marked polycythemia tend to be cyanotic at higher levels of Sao_2 than patients with normal hematocrit values. Likewise, local passive congestion, which causes an increase in the total quantity of reduced hemoglobin in the vessels in a given area, may cause cyanosis. Cyanosis is also observed when nonfunctional hemoglobin, such as methemoglobin (consequential or acquired) or sulfhemoglobin (Chap. 127), is present in blood.

Cyanosis may be subdivided into central and peripheral types. In *central* cyanosis, the Sao_2 is reduced or an abnormal hemoglobin derivative is present, and the mucous membranes and skin are both affected. *Peripheral* cyanosis is due to a slowing of blood flow and abnormally great extraction of O_2 from normally saturated arterial blood; it results from vasoconstriction and diminished peripheral blood flow, such as occurs in cold exposure, shock, congestive failure, and peripheral vascular disease. Often in these conditions, the mucous membranes of the oral cavity or those beneath the tongue may be spared. Clinical differentiation between central and peripheral cyanosis may not always be simple, and in conditions such as cardiogenic shock with pulmonary edema, there may be a mixture of both types.

DIFFERENTIAL DIAGNOSIS

Central Cyanosis (Table 49-1) Decreased Sao_2 results from a marked reduction in the Pao_2 . This reduction may be brought about by a decline in the Fio_2 without sufficient compensatory alveolar hyperventilation to maintain alveolar Po_2 . Cyanosis usually becomes manifest in an ascent to an altitude of 4000 m (13,000 ft).

Seriously *impaired pulmonary function*, through perfusion of unventilated or poorly ventilated areas of the lung or alveolar hypoventilation, is a common cause of central cyanosis (Chap. 306e).

TABLE 49-1 CAUSES OF CYANOSIS

Central Cyanosis

Decreased arterial oxygen saturation
Decreased atmospheric pressure—high altitude
Impaired pulmonary function
Alveolar hypoventilation
Inhomogeneity in pulmonary ventilation and perfusion (perfusion of hypoventilated alveoli)
Impaired oxygen diffusion
Anatomic shunts
Certain types of congenital heart disease
Pulmonary arteriovenous fistulas
Multiple small intrapulmonary shunts
Hemoglobin with low affinity for oxygen
Hemoglobin abnormalities
Methemoglobinemia—hereditary, acquired
Sulfhemoglobinemia—acquired
Carboxyhemoglobinemia (not true cyanosis)

Peripheral Cyanosis

Reduced cardiac output
Cold exposure
Redistribution of blood flow from extremities
Arterial obstruction
Venous obstruction

This condition may occur acutely, as in extensive pneumonia or pulmonary edema, or chronically, with chronic pulmonary diseases (e.g., emphysema). In the latter situation, secondary polycythemia is generally present and clubbing of the fingers (see below) may occur. Another cause of reduced Sao_2 is *shunting of systemic venous blood into the arterial circuit*. Certain forms of congenital heart disease are associated with cyanosis on this basis (see above and Chap. 282).

Pulmonary arteriovenous fistulae may be congenital or acquired, solitary or multiple, microscopic or massive. The severity of cyanosis produced by these fistulae depends on their size and number. They occur with some frequency in hereditary hemorrhagic telangiectasia. Sao_2 reduction and cyanosis may also occur in some patients with cirrhosis, presumably as a consequence of pulmonary arteriovenous fistulae or portal vein–pulmonary vein anastomoses.

In patients with cardiac or pulmonary right-to-left shunts, the presence and severity of cyanosis depend on the size of the shunt relative to the systemic flow as well as on the Hb- O_2 saturation of the venous blood. With increased extraction of O_2 from the blood by the exercising muscles, the venous blood returning to the right side of the heart is more unsaturated than at rest, and shunting of this blood intensifies the cyanosis. Secondary polycythemia occurs frequently in patients in this setting and contributes to the cyanosis.

Cyanosis can be caused by small quantities of circulating methemoglobin (Hb Fe^{3+}) and by even smaller quantities of sulfhemoglobin (Chap. 127); both of these hemoglobin derivatives impair oxygen delivery to the tissues. Although they are uncommon causes of cyanosis, these abnormal hemoglobin species should be sought by spectroscopy when cyanosis is not readily explained by malfunction of the circulatory or respiratory systems. Generally, digital clubbing does not occur with them.

Peripheral Cyanosis Probably the most common cause of peripheral cyanosis is the normal vasoconstriction resulting from exposure to cold air or water. When cardiac output is reduced, cutaneous vasoconstriction occurs as a compensatory mechanism so that blood is diverted from the skin to more vital areas such as the CNS and heart, and cyanosis of the extremities may result even though the arterial blood is normally saturated.

Arterial obstruction to an extremity, as with an embolus, or arteriolar constriction, as in cold-induced vasospasm (Raynaud's phenomenon) (Chap. 302), generally results in pallor and coldness, and there may be associated cyanosis. Venous obstruction, as in thrombophlebitis or deep venous thrombosis, dilates the subpapillary venous plexuses and thereby intensifies cyanosis.

APPROACH TO THE PATIENT:

Cyanosis

Certain features are important in arriving at the cause of cyanosis:

1. It is important to ascertain the time of onset of cyanosis. Cyanosis present since birth or infancy is usually due to congenital heart disease.
2. Central and peripheral cyanosis must be differentiated. Evidence of disorders of the respiratory or cardiovascular systems is helpful. Massage or gentle warming of a cyanotic extremity will increase peripheral blood flow and abolish peripheral, but not central, cyanosis.
3. The presence or absence of clubbing of the digits (see below) should be ascertained. The combination of cyanosis and clubbing is frequent in patients with congenital heart disease and right-to-left shunting and is seen occasionally in patients with pulmonary disease, such as lung abscess or pulmonary arteriovenous fistulae. In contrast, peripheral cyanosis or acutely developing central cyanosis is *not* associated with clubbed digits.
4. Pao_2 and Sao_2 should be determined, and, in patients with cyanosis in whom the mechanism is obscure, spectroscopic examination of the blood should be performed to look for abnormal types of hemoglobin (critical in the differential diagnosis of cyanosis).

The selective bulbous enlargement of the distal segments of the fingers and toes due to proliferation of connective tissue, particularly on the dorsal surface, is termed *clubbing*; there is also increased sponginess of the soft tissue at the base of the clubbed nail. Clubbing may be hereditary, idiopathic, or acquired and associated with a variety of disorders, including cyanotic congenital heart disease (see above), infective endocarditis, and a variety of pulmonary conditions (among them primary and metastatic lung cancer, bronchiectasis, asbestosis, sarcoidosis, lung abscess, cystic fibrosis, tuberculosis, and mesothelioma), as well as with some gastrointestinal diseases (including inflammatory bowel disease and hepatic cirrhosis). In some instances, it is occupational, e.g., in jackhammer operators.

Clubbing in patients with primary and metastatic lung cancer, mesothelioma, bronchiectasis, or hepatic cirrhosis may be associated with *hypertrophic osteoarthropathy*. In this condition, the subperiosteal formation of new bone in the distal diaphyses of the long bones of the extremities causes pain and symmetric arthritis-like changes in the shoulders, knees, ankles, wrists, and elbows. The diagnosis of hypertrophic osteoarthropathy may be confirmed by bone radiograph or magnetic resonance imaging (MRI). Although the mechanism of clubbing is unclear, it appears to be secondary to humoral substances that cause dilation of the vessels of the distal digits as well as growth factors released from platelet precursors in the digital circulation. In certain circumstances, clubbing is reversible, such as following lung transplantation for cystic fibrosis.

50

Edema

Eugene Braunwald, Joseph Loscalzo

STARLING FORCES AND FLUID EXCHANGE

About one-third of total-body water is confined to the extracellular space. Approximately 75% of the latter is interstitial fluid, and the remainder is the plasma. The forces that regulate the disposition of fluid between these two components of the extracellular compartment frequently are referred to as the *Starling forces*. The hydrostatic pressure within the capillaries and the colloid oncotic pressure in the interstitial fluid tend to promote movement of fluid from the vascular to the extravascular space. By contrast, the colloid oncotic pressure contributed by plasma proteins and the hydrostatic pressure within the interstitial fluid promote the movement of fluid into the vascular compartment. As a consequence, there is movement of water and diffusible solutes from the vascular space at the arteriolar end of the capillaries. Fluid is returned from the interstitial space into the vascular system at the venous end of the capillaries and by way of the lymphatics. These movements are usually balanced so that there is a steady state in the sizes of the intravascular and interstitial compartments, yet a large exchange between them occurs. However, if either the capillary hydrostatic pressure is increased and/or the oncotic pressure is reduced, a further net movement of fluid from intravascular to the interstitial spaces will take place.

Edema is defined as a clinically apparent increase in the interstitial fluid volume, which develops when Starling forces are altered so that there is increased flow of fluid from the vascular system into the interstitium. Edema due to an increase in capillary pressure may result from an elevation of venous pressure caused by obstruction to venous and/or lymphatic drainage. An increase in capillary pressure may be generalized, as occurs in heart failure, or it may be localized to one extremity when venous pressure is elevated due to unilateral thrombophlebitis (see below). The Starling forces also may be imbalanced when the colloid oncotic pressure of the plasma is reduced owing to any factor

that may induce hypoalbuminemia, as when large quantities of protein are lost in the urine such as in the nephrotic syndrome (see below), or when synthesis is reduced in a severe catabolic state.

CAPILLARY DAMAGE

Edema may also result from damage to the capillary endothelium, which increases its permeability and permits the transfer of proteins into the interstitial compartment. Injury to the capillary wall can result from drugs (see below), viral or bacterial agents, and thermal or mechanical trauma. Increased capillary permeability also may be a consequence of a hypersensitivity reaction and of immune injury. Damage to the capillary endothelium is presumably responsible for inflammatory edema, which is usually nonpitting, localized, and accompanied by other signs of inflammation—i.e., erythema, heat, and tenderness.

REDUCTION OF EFFECTIVE ARTERIAL VOLUME

In many forms of edema, despite the increase in extracellular fluid volume, the effective arterial blood volume, a parameter that represents the filling of the arterial tree and that effectively perfuses the tissues, is reduced. Underfilling of the arterial tree may be caused by a reduction of cardiac output and/or systemic vascular resistance, by pooling of blood in the splanchnic veins (as in cirrhosis), and by hypoalbuminemia (Fig. 50-1A). As a consequence of underfilling, a series of physiologic responses designed to restore the effective arterial volume to normal are set into motion. A key element of these responses is the renal retention of sodium and, therefore, water, thereby restoring effective arterial volume, but sometimes also leading to or intensifying edema.

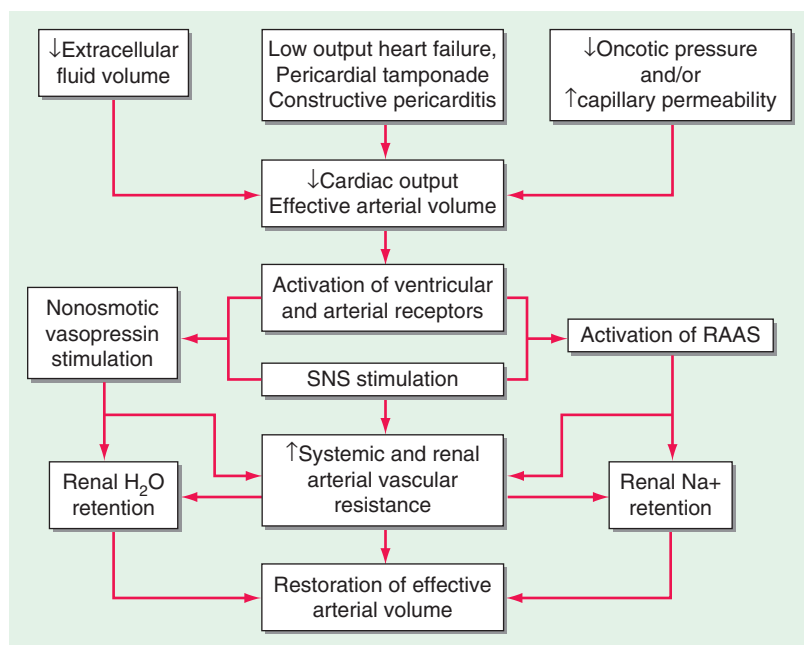
RENAL FACTORS AND THE RENIN-ANGIOTENSIN-ALDOSTERONE SYSTEM

The diminished renal blood flow characteristic of states in which the effective arterial blood volume is reduced is translated by the renal juxtaglomerular cells (specialized myoepithelial cells surrounding the afferent arteriole) into a signal for increased renin release. Renin is an enzyme with a molecular mass of about 40,000 Da that acts on its substrate, angiotensinogen, an α_2 -globulin synthesized by the liver, to release angiotensin I, a decapeptide, which in turn is converted to angiotensin II (AII), an octapeptide. AII has generalized vasoconstrictor properties, particularly on the renal efferent arterioles. This action reduces the hydrostatic pressure in the peritubular capillaries, whereas the increased filtration fraction raises the colloid osmotic pressure in these vessels, thereby enhancing salt and water reabsorption in the proximal tubule as well as in the ascending limb of the loop of Henle.

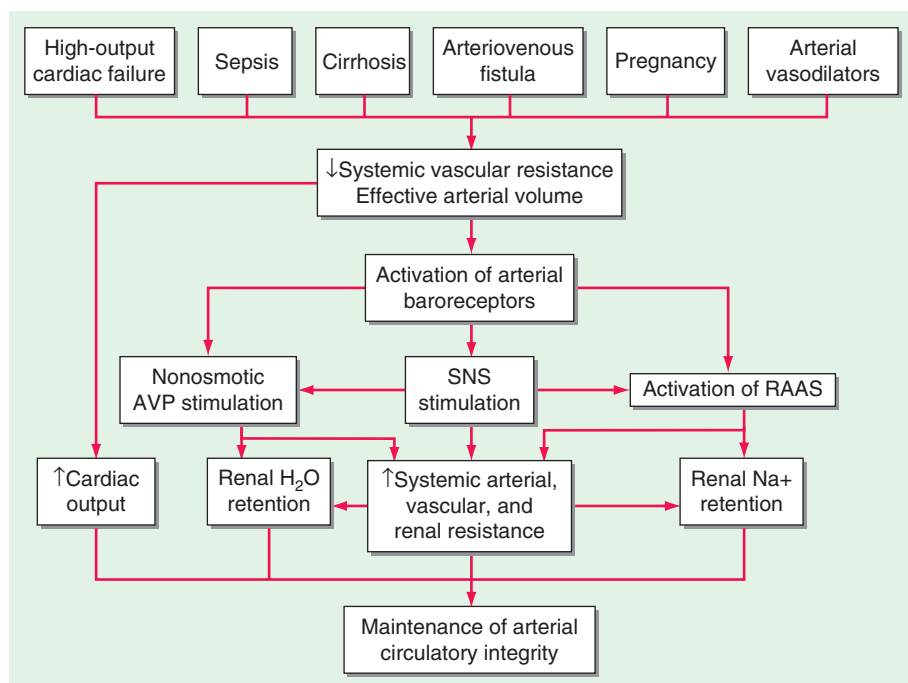
The renin-angiotensin-aldosterone system (RAAS) operates as both a hormonal and paracrine system. Its activation causes sodium and water retention and thereby contributes to edema formation. Blockade of the conversion of angiotensin I to AII and blockade of the AII receptor enhance sodium and water excretion and reduce many forms of edema. AII that enters the systemic circulation stimulates the production of aldosterone by the zona glomerulosa of the adrenal cortex. Aldosterone in turn enhances sodium reabsorption (and potassium excretion) by the collecting tubule, further favoring edema formation. In patients with heart failure, not only is aldosterone secretion elevated but the biologic half-life of the hormone is prolonged secondary to the depression of hepatic blood flow, which reduces its hepatic catabolism and increases further the plasma level of the hormone. Blockade of the action of aldosterone by spironolactone or eplerenone (aldosterone antagonists) or by amiloride (a blocker of epithelial sodium channels) often induces a moderate diuresis in edematous states.

ARGININE VASOPRESSIN

(See also Chap. 404) The secretion of arginine vasopressin (AVP) occurs in response to increased intracellular osmolar concentration, and, by stimulating V_2 receptors, AVP increases the reabsorption of free water in the distal tubules and collecting ducts of the kidneys, thereby increasing total-body water. Circulating AVP is elevated in many patients with heart failure secondary to a nonosmotic stimulus associated with decreased effective arterial volume and reduced compliance of the left atrium. Such patients fail to show the normal reduction of AVP with a reduction of osmolality, contributing to edema formation and hyponatremia.



A



B

FIGURE 50-1 Clinical conditions in which a decrease in cardiac output (A) and systemic vascular resistance (B) cause arterial underfilling with resulting neurohumoral activation and renal sodium and water retention. In addition to activating the neurohumoral axis, adrenergic stimulation causes renal vasoconstriction and enhances sodium and fluid transport by the proximal tubule epithelium. RAAS, renin-angiotensin aldosterone system; SNS, sympathetic nervous system. (Modified from RW Schrier: *Ann Intern Med* 113:155, 1990.)

ENDOTHELIN-1

This potent peptide vasoconstrictor is released by endothelial cells. Its concentration in the plasma is elevated in patients with severe heart failure and contributes to renal vasoconstriction, sodium retention, and edema.

NATRIURETIC PEPTIDES

Atrial distention causes release into the circulation of atrial natriuretic peptide (ANP), a polypeptide; a high-molecular-weight precursor of ANP is stored in secretory granules within atrial myocytes. The closely

related brain natriuretic peptide (pre-prohormone BNP) is stored primarily in ventricular myocytes and is released when ventricular diastolic pressure rises. Released ANP and BNP (which is derived from its precursor) bind to the natriuretic receptor-A, which causes: (1) excretion of sodium and water by augmenting glomerular filtration rate, inhibiting sodium reabsorption in the proximal tubule, and inhibiting release of renin and aldosterone; and (2) dilation of arterioles and venules by antagonizing the vasoconstrictor actions of AII, AVP, and sympathetic stimulation. Thus, elevated levels of natriuretic peptides have the capacity to oppose sodium retention in hypervolemic and edematous states.

Although circulating levels of ANP and BNP are elevated in heart failure and in cirrhosis with ascites, the natriuretic peptides are not sufficiently potent to prevent edema formation. Indeed, in edematous states, resistance to the actions of natriuretic peptides may be increased, further reducing their effectiveness.

Further discussion of the control of sodium and water balance is found in [Chap. 64e](#).

CLINICAL CAUSES OF EDEMA

A weight gain of several kilograms usually precedes overt manifestations of generalized edema, and a similar weight loss from diuresis can be induced in a slightly edematous patient before “dry weight” is achieved. *Anasarca* refers to gross, generalized edema. *Ascites* ([Chap. 59](#)) and *hydrothorax* refer to accumulation of excess fluid in the peritoneal and pleural cavities, respectively, and are considered special forms of edema.

Edema is recognized by the persistence of an indentation of the skin after pressure; this is known as “pitting” edema. In its more subtle form, edema may be detected by noting that after the stethoscope is removed from the chest wall, the rim of the bell leaves an indentation on the skin of the chest for a few minutes. Edema may be present when the ring on a finger fits more snugly than in the past or when a patient complains of difficulty putting on shoes, particularly in the evening. Edema may also be recognized by puffiness of the face, which is most readily apparent in the periorbital areas.

GENERALIZED EDEMA

The differences among the major causes of generalized edema are shown in [Table 50-1](#). Cardiac, renal, hepatic, or nutritional disorders are responsible for a majority of patients with generalized edema. Consequently, the

differential diagnosis of generalized edema should be directed toward identifying or excluding these several conditions.

Heart Failure (See also [Chap. 279](#)) In heart failure, the impaired systolic emptying of the ventricle(s) and/or the impairment of ventricular relaxation promotes an accumulation of blood in the venous circulation at the expense of the effective arterial volume. In addition, the heightened tone of the sympathetic nervous system causes renal vasoconstriction and reduction of glomerular filtration. In mild heart failure, a small increment of total blood volume may repair the deficit of

TABLE 50-1 PRINCIPAL CAUSES OF GENERALIZED EDEMA: HISTORY, PHYSICAL EXAMINATION, AND LABORATORY FINDINGS

Organ System	History	Physical Examination	Laboratory Findings
Cardiac	Dyspnea with exertion prominent—often associated with orthopnea—or paroxysmal nocturnal dyspnea	Elevated jugular venous pressure, ventricular (S ₃) gallop; occasionally with displaced or dyskinetic apical pulse; peripheral cyanosis, cool extremities, small pulse pressure when severe	Elevated urea nitrogen-to-creatinine ratio common; serum sodium often diminished; elevated natriuretic peptides
Hepatic	Dyspnea uncommon, except if associated with significant degree of ascites; most often a history of ethanol abuse	Frequently associated with ascites; jugular venous pressure normal or low; blood pressure lower than in renal or cardiac disease; one or more additional signs of chronic liver disease (jaundice, palmar erythema, Dupuytren's contracture, spider angiomas, male gynecomastia; asterix and other signs of encephalopathy) may be present	If severe, reductions in serum albumin, cholesterol, other hepatic proteins (transferrin, fibrinogen); liver enzymes elevated, depending on the cause and acuity of liver injury; tendency toward hypokalemia, respiratory alkalosis; macrocytosis from folate deficiency
Renal (CRF)	Usually chronic; may be associated with uremic signs and symptoms, including decreased appetite, altered (metallic or fishy) taste, altered sleep pattern, difficulty concentrating, restless legs, or myoclonus; dyspnea can be present, but generally less prominent than in heart failure	Elevated blood pressure; hypertensive retinopathy; nitrogenous fetor; pericardial friction rub in advanced cases with uremia	Elevation of serum creatinine and cystatin C; albuminuria; hyperkalemia, metabolic acidosis, hyperphosphatemia, hypocalcemia, anemia (usually normocytic)
Renal (NS)	Childhood diabetes mellitus; plasma cell dyscrasias	Periorbital edema; hypertension	Proteinuria (≥ 3.5 g/d); hypoalbuminemia; hypercholesterolemia; microscopic hematuria

Abbreviations: CRF, chronic renal failure; NS, nephrotic syndrome.

Source: Modified from GM Chertow: Approach to the patient with edema, in *Primary Cardiology*, 2nd ed, E Braunwald, L Goldman (eds). Philadelphia, Saunders, 2003, pp 117–128.

effective arterial volume through the operation of Starling's law of the heart, in which an increase in ventricular diastolic volume promotes a more forceful contraction and may thereby maintain the cardiac output. However, if the cardiac disorder is more severe, sodium and water retention continue, and the increment in blood volume accumulates in the venous circulation, raising venous pressure and causing edema (Fig. 50-1).

The presence of heart disease, as manifested by cardiac enlargement and/or ventricular hypertrophy, together with evidence of cardiac failure, such as dyspnea, basilar rales, venous distention, and hepatomegaly, usually indicates that edema results from heart failure. Noninvasive tests such as echocardiography may be helpful in establishing the diagnosis of heart disease. The edema of heart failure typically occurs in the dependent portions of the body.

Edema of Renal Disease (See also Chap. 338) The edema that occurs during the acute phase of glomerulonephritis is characteristically associated with hematuria, proteinuria, and hypertension. Although some evidence supports the view that the fluid retention is due to increased capillary permeability, in most instances, the edema results from primary retention of sodium and water by the kidneys owing to renal insufficiency. This state differs from most forms of heart failure in that it is characterized by a normal (or sometimes even increased) cardiac output. Patients with edema due to acute renal failure commonly have arterial hypertension as well as pulmonary congestion on chest roentgenogram, often without considerable cardiac enlargement, but they may not develop orthopnea. Patients with *chronic* renal failure may also develop edema due to primary renal retention of sodium and water.

Nephrotic Syndrome and other Hypoalbuminemic States The primary alteration in the nephrotic syndrome is a diminished colloid oncotic pressure due to losses of large quantities (≥ 3.5 g/d) of protein into the urine. With severe hypoalbuminemia (< 35 g/L) and the consequent reduced colloid osmotic pressure, the sodium and water that are retained cannot be restrained within the vascular compartment, and total and effective arterial blood volumes decline. This process initiates the edema-forming sequence of events described above, including activation of the RAAS. The nephrotic syndrome may occur during the course of a variety of kidney diseases, which include glomerulonephritis, diabetic glomerulosclerosis, and hypersensitivity reactions. The edema is diffuse, symmetric, and most prominent in the dependent

areas; as a consequence, periorbital edema is most prominent in the morning.

Hepatic Cirrhosis (See also Chap. 365) This condition is characterized in part by hepatic venous outflow blockade, which in turn expands the splanchnic blood volume and increases hepatic lymph formation. Intrahepatic hypertension acts as a stimulus for renal sodium retention and causes a reduction of effective arterial blood volume. These alterations are frequently complicated by hypoalbuminemia secondary to reduced hepatic synthesis of albumin, as well as peripheral arterial vasodilation. These effects reduce the effective arterial blood volume further, leading to activation of the RAAS and renal sympathetic nerves and to release of AVP, endothelin, and other sodium- and water-retaining mechanisms (Fig. 50-1B). The concentration of circulating aldosterone often is elevated by the failure of the liver to metabolize this hormone. Initially, the excess interstitial fluid is localized preferentially proximal (upstream) to the congested portal venous system and obstructed hepatic lymphatics, i.e., in the peritoneal cavity (causing ascites, Chap. 59). In later stages, particularly when there is severe hypoalbuminemia, peripheral edema may develop. A sizable accumulation of ascitic fluid may increase intraabdominal pressure and impede venous return from the lower extremities and contribute to the accumulation of edema of the lower extremities.

The excess production of prostaglandins (PGE₂ and PGI₂) in cirrhosis attenuates renal sodium retention. When the synthesis of these substances is inhibited by nonsteroidal anti-inflammatory drugs (NSAIDs), renal function may deteriorate, and this may increase sodium retention further.

Drug-Induced Edema A large number of widely used drugs can cause edema (Table 50-2). Mechanisms include renal vasoconstriction (NSAIDs and cyclosporine), arteriolar dilation (vasodilators), augmented renal sodium reabsorption (steroid hormones), and capillary damage.

Edema of Nutritional Origin A diet grossly deficient in protein over a prolonged period may produce hypoproteinemia and edema. The latter may be intensified by the development of beriberi heart disease, which also is of nutritional origin, in which multiple peripheral arteriovenous fistulae result in reduced effective systemic perfusion and effective arterial blood volume, thereby enhancing edema formation (Chap. 96e) (Fig. 50-1B). Edema may actually become intensified

TABLE 50-2 DRUGS ASSOCIATED WITH EDEMA FORMATION

Nonsteroidal anti-inflammatory drugs
Antihypertensive agents
Direct arterial/arteriolar vasodilators
Hydralazine
Clonidine
Methyldopa
Guanethidine
Minoxidil
Calcium channel antagonists
α-Adrenergic antagonists
Thiazolidinediones
Steroid hormones
Glucocorticoids
Anabolic steroids
Estrogens
Progestins
Cyclosporine
Growth hormone
Immunotherapies
Interleukin 2
OKT3 monoclonal antibody

Source: Modified from Chertow GM: Approach to the patient with edema, in *Primary Cardiology*, 2nd ed, E Braunwald, L Goldman (eds). Philadelphia, Saunders, 2003, pp 117–128.

when famished subjects are first provided with an adequate diet. The ingestion of more food may increase the quantity of sodium ingested, which is then retained along with water. So-called refeeding edema also may be linked to increased release of insulin, which directly increases tubular sodium reabsorption. In addition to hypoalbuminemia, hypokalemia and caloric deficits may be involved in the edema of starvation.

LOCALIZED EDEMA

In this condition, the hydrostatic pressure in the capillary bed upstream (proximal) of the obstruction increases so that an abnormal quantity of fluid is transferred from the vascular to the interstitial space. Since the alternative route (i.e., the lymphatic channels) also may be obstructed or maximally filled, an increased volume of interstitial fluid in the limb develops (i.e., there is trapping of fluid in the interstitium of the extremity). The displacement of large quantities of fluid into a limb may occur at the expense of the blood volume in the remainder of the body, thereby reducing effective arterial blood volume and leading to the retention of NaCl and H₂O until the deficit in plasma volume has been corrected.

Localized edema due to venous or lymphatic obstruction may be caused by thrombophlebitis, chronic lymphangitis, resection of regional lymph nodes, and filariasis, among other causes. Lymphedema is particularly intractable because restriction of lymphatic flow results in increased protein concentration in the interstitial fluid, a circumstance that aggravates fluid retention.

Other Causes of Edema These causes include hypothyroidism (myxedema) and hyperthyroidism (pretibial myxedema secondary to Graves' disease), the edema in which is typically nonpitting and due to deposition of hyaluronic acid and, in Graves' disease, lymphocytic infiltration and inflammation; exogenous hyperadrenocorticism; pregnancy; and administration of estrogens and vasodilators, particularly dihydropyridines such as nifedipine.

DISTRIBUTION OF EDEMA

The distribution of edema is an important guide to its cause. Edema associated with heart failure tends to be more extensive in the legs and to be accentuated in the evening, a feature also determined largely by

posture. When patients with heart failure are confined to bed, edema may be most prominent in the presacral region. Severe heart failure may cause ascites that may be distinguished from the ascites caused by hepatic cirrhosis by the jugular venous pressure, which is usually elevated in heart failure and normal in cirrhosis.

Edema resulting from hypoproteinemia, as occurs in the nephrotic syndrome, characteristically is generalized, but it is especially evident in the very soft tissues of the eyelids and face and tends to be most pronounced in the morning owing to the recumbent posture assumed during the night. Less common causes of facial edema include trichinosis, allergic reactions, and myxedema. Edema limited to one leg or to one or both arms is usually the result of venous and/or lymphatic obstruction. Unilateral paralysis reduces lymphatic and venous drainage on the affected side and may also be responsible for unilateral edema. In patients with obstruction of the superior vena cava, edema is confined to the face, neck, and upper extremities in which the venous pressure is elevated compared with that in the lower extremities.

APPROACH TO THE PATIENT:

Edema

An important first question is whether the edema is localized or generalized. If it is localized, the local phenomena that may be responsible should be considered. If the edema is generalized, one should first determine if there is serious hypoalbuminemia, e.g., serum albumin <25 g/L. If so, the history, physical examination, urinalysis, and other laboratory data will help evaluate the question of cirrhosis, severe malnutrition, or the nephrotic syndrome as the underlying disorder. If hypoalbuminemia is not present, it should be determined if there is evidence of heart failure severe enough to promote generalized edema. Finally, it should be ascertained as to whether or not the patient has an adequate urine output or if there is significant oliguria or anuria. **These abnormalities are discussed in Chaps. 61, 334, and 335.**

51e

Approach to the Patient with a Heart Murmur

Patrick T. O'Gara, Joseph Loscalzo

This is a digital-only chapter. It is available on the DVD that accompanies this book, as well as on Access Medicine/Harrison's Online, and the eBook and "app" editions of HPIIM 19e.

The differential diagnosis of a heart murmur begins with a careful assessment of its major attributes and response to bedside maneuvers. The history, clinical context, and associated physical examination findings provide additional clues by which the significance of a heart murmur can be established. Accurate bedside identification of a heart murmur can inform decisions regarding the indications for noninvasive testing and the need for referral to a cardiovascular specialist. Preliminary discussions can be held with the patient regarding antibiotic or rheumatic fever prophylaxis, the need to restrict various forms of physical activity, and the potential role for family screening.

The differential diagnosis of a heart murmur begins with a careful assessment of its major attributes and response to bedside maneuvers. The history, clinical context, and associated physical examination findings provide additional clues by which the significance of a heart murmur can be established. Accurate bedside identification of a heart murmur can inform decisions regarding the indications for noninvasive testing and the need for referral to a cardiovascular specialist. Preliminary discussions can be held with the patient regarding antibiotic or rheumatic fever prophylaxis, the need to restrict various forms of physical activity, and the potential role for family screening.

Heart murmurs are caused by audible vibrations that are due to increased turbulence from accelerated blood flow through normal or abnormal orifices, flow through a narrowed or irregular orifice into a dilated vessel or chamber, or backward flow through an incompetent valve, ventricular septal defect, or patent ductus arteriosus. They traditionally are defined in terms of their timing within the cardiac cycle (Fig. 51e-1). *Systolic murmurs* begin with or after the first heart sound (S_1) and terminate at or before the component (A_2 or P_2) of the second heart sound (S_2) that corresponds to their site of origin (left or right, respectively). *Diastolic murmurs* begin with or after the associated component of S_2 and end at or before the subsequent S_1 . *Continuous murmurs* are not confined to either phase of the cardiac cycle but instead begin in early systole and proceed through S_2 into all or part of diastole. The accurate timing of heart murmurs is the first step in their identification. The distinction between S_1 and S_2 and, therefore, systole and diastole is usually a straightforward process but can be difficult in the setting of a tachyarrhythmia, in which case the heart sounds can be distinguished by simultaneous palpation of the carotid upstroke, which should closely follow S_1 .

Duration and Character The duration of a heart murmur depends on the length of time over which a pressure difference exists between two cardiac chambers, the left ventricle and the aorta, the right ventricle and the pulmonary artery, or the great vessels. The magnitude and variability of this pressure difference, coupled with the geometry and compliance of the involved chambers or vessels, dictate the velocity of flow; the degree of turbulence; and the resulting frequency, configuration, and intensity of the murmur. The diastolic murmur of chronic aortic regurgitation (AR) is a blowing, high-frequency event, whereas the murmur of mitral stenosis (MS), indicative of the left atrial–left ventricular diastolic pressure gradient, is a low-frequency event, heard as a rumbling sound with the bell of the stethoscope. The frequency components of a heart murmur may vary at different sites of auscultation. The coarse systolic murmur of aortic stenosis (AS) may sound higher pitched and more acoustically pure at the apex, a phenomenon eponymously referred to as the *Gallavardin effect*. Some murmurs may have a distinct or unusual quality, such as the “honking” sound appreciated in some patients with mitral regurgitation (MR) due to mitral valve prolapse (MVP).

The configuration of a heart murmur may be described as crescendo, decrescendo, crescendo-decrescendo, or plateau. The decrescendo configuration of the murmur of chronic AR (Fig. 51e-1E) can be understood in terms of the progressive decline in the diastolic pressure gradient between the aorta and the left ventricle. The crescendo-decrescendo configuration of the murmur of AS reflects the changes in the systolic pressure gradient between the left ventricle and the aorta as ejection occurs, whereas the plateau configuration of the murmur of chronic MR (Fig. 51e-1B) is consistent with the large and nearly constant pressure difference between the left ventricle and the left atrium.

Intensity The intensity of a heart murmur is graded on a scale of 1–6 (or I–VI). A grade 1 murmur is very soft and is heard only with great

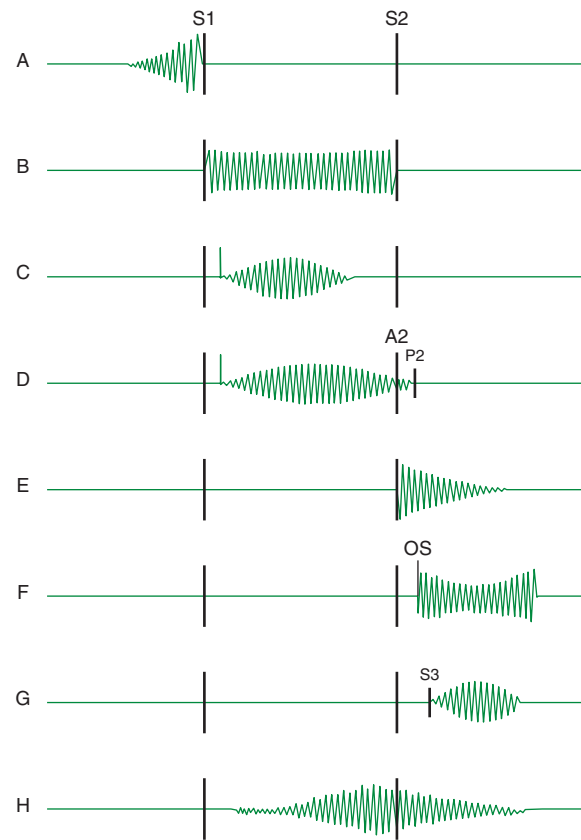


FIGURE 51e-1 Diagram depicting principal heart murmurs. **A.** Presystolic murmur of mitral or tricuspid stenosis. **B.** Holosystolic (pansystolic) murmur of mitral or tricuspid regurgitation or of ventricular septal defect. **C.** Aortic ejection murmur beginning with an ejection click and fading before the second heart sound. **D.** Systolic murmur in pulmonic stenosis spilling through the aortic second sound, pulmonic valve closure being delayed. **E.** Aortic or pulmonary diastolic murmur. **F.** Long diastolic murmur of mitral stenosis after the opening snap (OS). **G.** Short mid-diastolic inflow murmur after a third heart sound. **H.** Continuous murmur of patent ductus arteriosus. (Adapted from P Wood: *Diseases of the Heart and Circulation*, London, Eyre & Spottiswood, 1968. Permission granted courtesy of Antony and Julie Wood.)

effort. A grade 2 murmur is easily heard but not particularly loud. A grade 3 murmur is loud but is not accompanied by a palpable thrill over the site of maximal intensity. A grade 4 murmur is very loud and is accompanied by a thrill. A grade 5 murmur is loud enough to be heard with only the edge of the stethoscope touching the chest, whereas a grade 6 murmur is loud enough to be heard with the stethoscope slightly off the chest. Murmurs of grade 3 or greater intensity usually signify important structural heart disease and indicate high blood flow velocity at the site of murmur production. Small ventricular septal defects (VSDs), for example, are accompanied by loud, usually grade 4 or greater, systolic murmurs as blood is ejected at high velocity from the left ventricle to the right ventricle. Low-velocity events, such as left-to-right shunting across an atrial septal defect (ASD), are usually silent. The intensity of a heart murmur may be diminished by any process that increases the distance between the intracardiac source and the stethoscope on the chest wall, such as obesity, obstructive lung disease, and a large pericardial effusion. The intensity of a murmur also may be misleadingly soft when cardiac output is reduced significantly or when the pressure gradient between the involved cardiac structures is low.

Location and Radiation Recognition of the location and radiation of the murmur helps facilitate its accurate identification (Fig. 51e-2). Adventitious sounds, such as a systolic click or diastolic snap, or abnormalities of S_1 or S_2 may provide additional clues. Careful attention to the characteristics of the murmur and other heart sounds

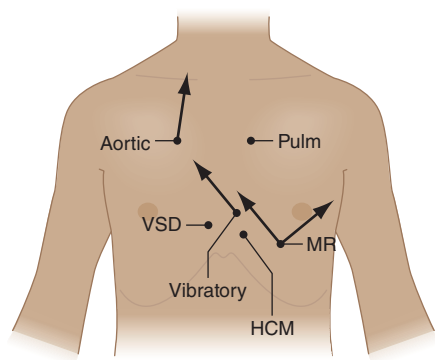


FIGURE 51e-2 Maximal intensity and radiation of six isolated systolic murmurs. HOCM, hypertrophic obstructive cardiomyopathy; MR, mitral regurgitation; Pulm, pulmonary stenosis; Aortic, aortic stenosis; VSD, ventricular septal defect. (From JB Barlow: *Perspectives on the Mitral Valve*. Philadelphia, FA Davis, 1987, p 140.)

during the respiratory cycle and the performance of simple bedside maneuvers complete the auscultatory examination. These features, along with recommendations for further testing, are discussed below in the context of specific systolic, diastolic, and continuous heart murmurs (Table 51e-1).

SYSTOLIC HEART MURMURS

Early Systolic Murmurs Early systolic murmurs begin with S_1 and extend for a variable period, ending well before S_2 . Their causes are relatively few in number. *Acute, severe MR* into a normal-sized, relatively noncompliant left atrium results in an early, decrescendo systolic murmur best heard at or just medial to the apical impulse. These characteristics reflect the progressive attenuation of the pressure gradient between the left ventricle and the left atrium during systole owing to the rapid rise in left atrial pressure caused by the sudden volume load into an unprepared, noncompliant chamber and contrast sharply with the auscultatory features of chronic MR. Clinical settings in which acute, severe MR occur include (1) papillary muscle rupture complicating acute myocardial infarction (MI) (Chap. 295), (2) rupture of chordae tendineae in the setting of myxomatous mitral valve disease (MVP, Chap. 283), (3) infective endocarditis (Chap. 155), and (4) blunt chest wall trauma.

Acute, severe MR from papillary muscle rupture usually accompanies an inferior, posterior, or lateral MI and occurs 2–7 days after presentation. It often is signaled by chest pain, hypotension, and pulmonary edema, but a murmur may be absent in up to 50% of cases. The posteromedial papillary muscle is involved 6 to 10 times more frequently than the anterolateral papillary muscle. The murmur is to be distinguished from that associated with post-MI ventricular septal rupture, which is accompanied by a systolic thrill at the left sternal border in nearly all patients and is holosystolic in duration. A new heart murmur after an MI is an indication for transthoracic echocardiography (TTE) (Chap. 270e), which allows bedside delineation of its etiology and pathophysiologic significance. The distinction between acute MR and ventricular septal rupture also can be achieved with right heart catheterization, sequential determination of oxygen saturations, and analysis of the pressure waveforms (tall v wave in the pulmonary artery wedge pressure in MR). Post-MI mechanical complications of this nature mandate aggressive medical stabilization and prompt referral for surgical repair.

Spontaneous chordal rupture can complicate the course of myxomatous mitral valve disease (MVP) and result in new-onset or “acute on chronic” severe MR. MVP may occur as an isolated phenomenon, or the lesion may be part of a more generalized connective tissue disorder as seen, for example, in patients with Marfan syndrome. Acute, severe MR as a consequence of infective endocarditis results from destruction of leaflet tissue, chordal rupture, or both. Blunt chest wall trauma is usually self-evident but may be disarmingly trivial; it

TABLE 51e-1 PRINCIPAL CAUSES OF HEART MURMURS

Systolic Murmurs	
Early systolic	
Mitral	Acute MR
VSD	
Muscular	Nonrestrictive with pulmonary hypertension
Tricuspid	TR with normal pulmonary artery pressure
Mid-systolic	
Aortic	
Obstructive	Supravalvular–supravalvular aortic stenosis, coarctation of the aorta
	Valvular–AS and aortic sclerosis
	Subvalvular–discrete, tunnel or HOCM
	Increased flow, hyperkinetic states, AR, complete heart block
	Dilation of ascending aorta, atheroma, aortitis
Pulmonary	
Obstructive	Supravalvular–pulmonary artery stenosis
	Valvular–pulmonic valve stenosis
	Subvalvular–infundibular stenosis (dynamic)
	Increased flow, hyperkinetic states, left-to-right shunt (e.g., ASD)
	Dilation of pulmonary artery
Late systolic	
Mitral	
	MVP, acute myocardial ischemia
Tricuspid	
	TVP
Holosystolic	
	Atrioventricular valve regurgitation (MR, TR)
	Left-to-right shunt at ventricular level (VSD)
Early Diastolic Murmurs	
Aortic regurgitation	
	Valvular: congenital (bicuspid valve), rheumatic deformity, endocarditis, prolapse, trauma, post-valvulotomy
	Dilation of valve ring: aorta dissection, annuloaortic ectasia, cystic medial degeneration, hypertension, ankylosing spondylitis
	Widening of commissures: syphilis
Pulmonic regurgitation	
	Valvular: post-valvulotomy, endocarditis, rheumatic fever, carcinoid
	Dilation of valve ring: pulmonary hypertension; Marfan syndrome
	Congenital: isolated or associated with tetralogy of Fallot, VSD, pulmonic stenosis
Mid-Diastolic Murmurs	
Mitral	
	Mitral stenosis
	Carey-Coombs murmur (mid-diastolic apical murmur in acute rheumatic fever)
	Increased flow across nonstenotic mitral valve (e.g., MR, VSD, PDA, high-output states, and complete heart block)
Tricuspid	
	Tricuspid stenosis
	Increased flow across nonstenotic tricuspid valve (e.g., TR, ASD, and anomalous pulmonary venous return)
	Left and right atrial tumors (myxoma)
	Severe AR (Austin Flint murmur)
Continuous Murmurs	
Patent ductus arteriosus	Proximal coronary artery stenosis
Coronary AV fistula	Mammary souffle of pregnancy
Ruptured sinus of Valsalva aneurysm	Pulmonary artery branch stenosis
Aortic septal defect	Bronchial collateral circulation
Cervical venous hum	Small (restrictive) ASD with MS
Anomalous left coronary artery	Intercostal AV fistula

Abbreviations: AR, aortic regurgitation; AS, aortic stenosis; ASD, atrial septal defect; AV, arteriovenous; HOCM, hypertrophic obstructive cardiomyopathy; MR, mitral regurgitation; MS, mitral stenosis; MVP, mitral valve prolapse; PDA, patent ductus arteriosus; TR, tricuspid regurgitation; TVP, tricuspid valve prolapse; VSD, ventricular septal defect.

Source: E Braunwald, JK Perloff, in D Zipes et al (eds): *Braunwald's Heart Disease*, 7th ed. Philadelphia, Elsevier, 2005; PJ Norton, RA O'Rourke, in E Braunwald, L Goldman (eds): *Primary Cardiology*, 2nd ed. Philadelphia, Elsevier, 2003.

can result in papillary muscle contusion and rupture, chordal detachment, or leaflet avulsion. TTE is indicated in all cases of suspected acute, severe MR to define its mechanism and severity, delineate left ventricular size and systolic function, and provide an assessment of suitability for primary valve repair.

A congenital, small muscular VSD (**Chap. 282**) may be associated with an early systolic murmur. The defect closes progressively during septal contraction, and thus, the murmur is confined to early systole. It is localized to the left sternal border (Fig. 51e-2) and is usually of grade 4 or 5 intensity. Signs of pulmonary hypertension or left ventricular volume overload are absent. Anatomically large and uncorrected VSDs, which usually involve the membranous portion of the septum, may lead to pulmonary hypertension. The murmur associated with the left-to-right shunt, which earlier may have been holosystolic, becomes limited to the first portion of systole as the elevated pulmonary vascular resistance leads to an abrupt rise in right ventricular pressure and an attenuation of the interventricular pressure gradient during the remainder of the cardiac cycle. In such instances, signs of pulmonary hypertension (right ventricular lift, loud and single or closely split S_2) may predominate. The murmur is best heard along the left sternal border but is softer. Suspicion of a VSD is an indication for TTE.

Tricuspid regurgitation (TR) with normal pulmonary artery pressures, as may occur with infective endocarditis, may produce an early systolic murmur. The murmur is soft (grade 1 or 2), is best heard at the lower left sternal border, and may increase in intensity with inspiration (Carvallo's sign). Regurgitant "c-v" waves may be visible in the jugular venous pulse. TR in this setting is not associated with signs of right heart failure.

Mid-Systolic Murmurs Mid-systolic murmurs begin at a short interval after S_1 , end before S_2 (Fig. 51e-1C), and are usually crescendo-decrescendo in configuration. Aortic stenosis is the most common cause of a mid-systolic murmur in an adult. The murmur of AS is usually loudest to the right of the sternum in the second intercostal space (aortic area, Fig. 51e-2) and radiates into the carotids. Transmission of the mid-systolic murmur to the apex, where it becomes higher-pitched, is common (Gallavardin effect; see above).

Differentiation of this apical systolic murmur from MR can be difficult. The murmur of AS will increase in intensity, or become louder, in the beat after a premature beat, whereas the murmur of MR will have constant intensity from beat to beat. The intensity of the AS murmur also varies directly with the cardiac output. With a normal cardiac output, a systolic thrill and a grade 4 or higher murmur suggest severe AS. The murmur is softer in the setting of heart failure and low cardiac output. Other auscultatory findings of severe AS include a soft or absent A_2 , paradoxical splitting of S_2 , an apical S_4 , and a late-peaking systolic murmur. In children, adolescents, and young adults with congenital valvular AS, an early ejection sound (click) is usually audible, more often along the left sternal border than at the base. Its presence signifies a flexible, noncalcified bicuspid valve (or one of its variants) and localizes the left ventricular outflow obstruction to the valvular (rather than sub- or supra-valvular) level.

Assessment of the volume and rate of rise of the carotid pulse can provide additional information. A small and delayed upstroke (*parvus et tardus*) is consistent with severe AS. The carotid pulse examination is less discriminatory, however, in older patients with stiffened arteries. The electrocardiogram (ECG) shows signs of left ventricular hypertrophy (LVH) as the severity of the stenosis increases. TTE is indicated to assess the anatomic features of the aortic valve, the severity of the stenosis, left ventricular size, wall thickness and function, and the size and contour of the aortic root and proximal ascending aorta.

The obstructive form of hypertrophic cardiomyopathy (HOCM) is associated with a mid-systolic murmur that is usually loudest along the left sternal border or between the left lower sternal border and the apex (**Chap. 287**, Fig. 51e-2). The murmur is produced by both dynamic left ventricular outflow tract obstruction and MR, and thus, its configuration is a hybrid between ejection and regurgitant phenomena. The intensity of the murmur may vary from beat to beat and after provocative maneuvers but usually does not exceed grade 3. The

murmur classically will increase in intensity with maneuvers that result in increasing degrees of outflow tract obstruction, such as a reduction in preload or afterload (Valsalva, standing, vasodilators), or with an augmentation of contractility (inotropic stimulation). Maneuvers that increase preload (squatting, passive leg raising, volume administration) or afterload (squatting, vasopressors) or that reduce contractility (β -adrenoreceptor blockers) decrease the intensity of the murmur. In rare patients, there may be reversed splitting of S_2 . A sustained left ventricular apical impulse and an S_4 may be appreciated. In contrast to AS, the carotid upstroke is rapid and of normal volume. Rarely, it is bisferiens or bifid in contour (see Fig. 267-2D) due to mid-systolic closure of the aortic valve. LVH is present on the ECG, and the diagnosis is confirmed by TTE. Although the systolic murmur associated with MVP behaves similarly to that due to HOCM in response to the Valsalva maneuver and to standing/squatting (Fig. 51e-3), these two lesions can be distinguished on the basis of their associated findings, such as the presence of LVH in HOCM or a nonejection click in MVP.

The mid-systolic, crescendo-decrescendo murmur of congenital pulmonic stenosis (PS, **Chap. 282**) is best appreciated in the second and third left intercostal spaces (pulmonic area) (Figs. 51e-2 and 51e-4). The duration of the murmur lengthens and the intensity of P_2 diminishes with increasing degrees of valvular stenosis (Fig. 51e-1D). An early ejection sound, the intensity of which decreases with inspiration, is heard in younger patients. A parasternal lift and ECG evidence of right ventricular hypertrophy indicate severe pressure overload. If obtained, the chest x-ray may show poststenotic dilation of the main pulmonary artery. TTE is recommended for complete characterization.

Significant left-to-right intracardiac shunting due to an ASD (**Chap. 282**) leads to an increase in pulmonary blood flow and a grade 2–3 mid-systolic murmur at the middle to upper left sternal border

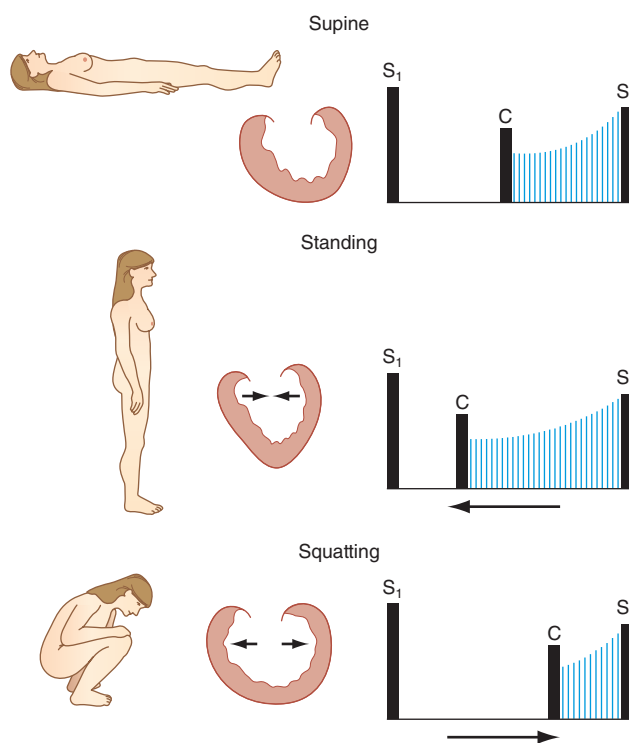


FIGURE 51e-3 A mid-systolic nonejection sound (C) occurs in mitral valve prolapse and is followed by a late systolic murmur that crescendos to the second heart sound (S_2). Standing decreases venous return; the heart becomes smaller; C moves closer to the first heart sound (S_1), and the mitral regurgitant murmur has an earlier onset. With prompt squatting, venous return and afterload increase; the heart becomes larger; C moves toward S_2 ; and the duration of the murmur shortens. (From JA Shaver, JJ Leonard, DF Leon: *Examination of the Heart, Part IV, Auscultation of the Heart*. Dallas, American Heart Association, 1990, p 13. Copyright, American Heart Association.)

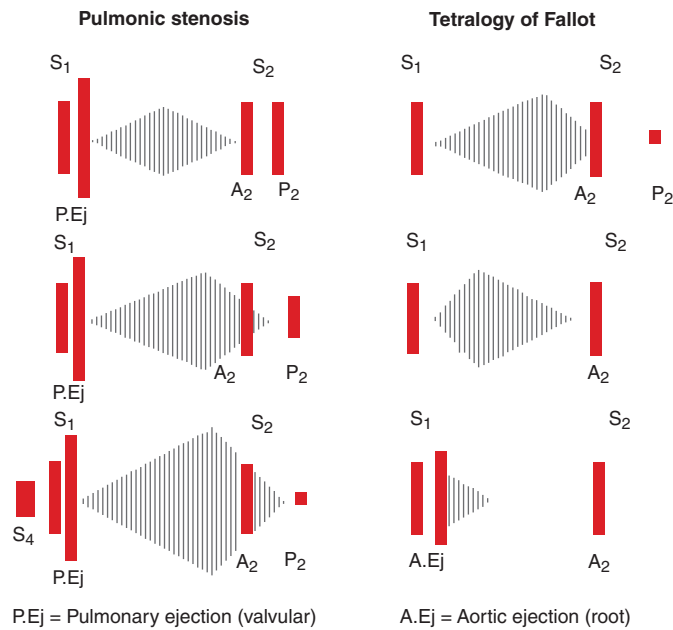


FIGURE 51e-4 *Left.* In valvular pulmonic stenosis with intact ventricular septum, right ventricular systolic ejection becomes progressively longer, with increasing obstruction to flow. As a result, the murmur becomes longer and louder, enveloping the aortic component of the second heart sound (A_2). The pulmonic component (P_2) occurs later, and splitting becomes wider but more difficult to hear because A_2 is lost in the murmur and P_2 becomes progressively fainter and lower pitched. As the pulmonic gradient increases, the isometric contraction phase shortens until the pulmonic valve ejection sound fuses with the first heart sound (S_1). In severe pulmonic stenosis with concentric hypertrophy and decreasing right ventricular compliance, a fourth heart sound appears. *Right.* In tetralogy of Fallot with increasing obstruction at the pulmonic infundibular area, an increasing amount of right ventricular blood is shunted across the silent ventricular septal defect and flow across the obstructed outflow tract decreases. Therefore, with increasing obstruction the murmur becomes shorter, earlier, and fainter. P_2 is absent in severe tetralogy of Fallot. A large aortic root receives almost all cardiac output from both ventricular chambers, and the aorta dilates and is accompanied by a root ejection sound that does not vary with respiration. (From JA Shaver, JJ Leonard, DF Leon: *Examination of the Heart, Part IV, Auscultation of the Heart*. Dallas, American Heart Association, 1990, p 45. Copyright, American Heart Association.)

attributed to increased flow rates across the pulmonic valve with fixed splitting of S_2 . Ostium secundum ASDs are the most common cause of these shunts in adults. Features suggestive of a primum ASD include the coexistence of MR due to a cleft anterior mitral valve leaflet and left axis deviation of the QRS complex on the ECG. With sinus venosus ASDs, the left-to-right shunt is usually not large enough to result in a systolic murmur, although the ECG may show abnormalities of sinus node function. A grade 2 or 3 mid-systolic murmur may also be heard best at the upper left sternal border in patients with idiopathic dilation of the pulmonary artery; a pulmonary ejection sound is also present in these patients. TTE is indicated to evaluate a grade 2 or 3 mid-systolic murmur when there are other signs of cardiac disease.

An isolated grade 1 or 2 mid-systolic murmur, heard in the absence of symptoms or signs of heart disease, is most often a benign finding for which no further evaluation, including TTE, is necessary. The most common example of a murmur of this type in an older adult patient is the crescendo-decrescendo murmur of aortic valve sclerosis, heard at the second right interspace (Fig. 51e-2). Aortic sclerosis is defined as focal thickening and calcification of the aortic valve to a degree that does not interfere with leaflet opening. The carotid upstrokes are normal, and electrocardiographic LVH is not present. A grade 1 or 2 mid-systolic murmur often can be heard at the left sternal border

with pregnancy, hyperthyroidism, or anemia, physiologic states that are associated with accelerated blood flow. *Still's murmur* refers to a benign grade 2, vibratory or musical mid-systolic murmur at the mid or lower left sternal border in normal children and adolescents, best heard in the supine position (Fig. 51e-2).

Late Systolic Murmurs A late systolic murmur that is best heard at the left ventricular apex is usually due to MVP (Chap. 283). Often, this murmur is introduced by one or more nonejection clicks. The radiation of the murmur can help identify the specific mitral leaflet involved in the process of prolapse or flail. The term *flail* refers to the movement made by an unsupported portion of the leaflet after loss of its chordal attachment(s). With posterior leaflet prolapse or flail, the resultant jet of MR is directed anteriorly and medially, as a result of which the murmur radiates to the base of the heart and masquerades as AS. Anterior leaflet prolapse or flail results in a posteriorly directed MR jet that radiates to the axilla or left infrascapular region. Leaflet flail is associated with a murmur of grade 3 or 4 intensity that can be heard throughout the precordium in thin-chested patients. The presence of an S_3 or a short, rumbling mid-diastolic murmur due to enhanced flow signifies severe MR.

Bedside maneuvers that decrease left ventricular preload, such as standing, will cause the click and murmur of MVP to move closer to the first heart sound, as leaflet prolapse occurs earlier in systole. Standing also causes the murmur to become louder and longer. With squatting, left ventricular preload and afterload are increased abruptly, leading to an increase in left ventricular volume, and the click and murmur move away from the first heart sound as leaflet prolapse is delayed; the murmur becomes softer and shorter in duration (Fig. 51e-3). As noted above, these responses to standing and squatting are directionally similar to those observed in patients with HOCM.

A late, apical systolic murmur indicative of MR may be heard transiently in the setting of acute myocardial ischemia; it is due to apical tethering and malcoaptation of the leaflets in response to structural and functional changes of the ventricle and mitral annulus. The intensity of the murmur varies as a function of left ventricular afterload and will increase in the setting of hypertension. TTE is recommended for assessment of late systolic murmurs.

Holosystolic Murmurs (Figs. 51e-1B and 51e-5) Holosystolic murmurs begin with S_1 and continue through systole to S_2 . They are usually indicative of chronic mitral or tricuspid valve regurgitation or a VSD and warrant TTE for further characterization. The holosystolic murmur of chronic MR is best heard at the left ventricular apex and radiates to the axilla (Fig. 51e-2); it is usually high-pitched and plateau in configuration because of the wide difference between left ventricular and left atrial pressure throughout systole. In contrast to acute MR, left atrial compliance is normal or even increased in chronic MR. As a result, there is only a small increase in left atrial pressure for any increase in regurgitant volume.

Several conditions are associated with chronic MR and an apical holosystolic murmur, including rheumatic scarring of the leaflets, mitral annular calcification, postinfarction left ventricular remodeling, and severe left ventricular chamber enlargement. The circumference of the mitral annulus increases as the left ventricle enlarges and leads to failure of leaflet coaptation with central MR in patients with dilated cardiomyopathy (Chap. 287). The severity of the MR is worsened by any contribution from apical displacement of the papillary muscles and leaflet tethering (remodeling). Because the mitral annulus is contiguous with the left atrial endocardium, gradual enlargement of the left atrium from chronic MR will result in further stretching of the annulus and more MR; thus, "MR begets MR." Chronic severe MR results in enlargement and leftward displacement of the left ventricular apex beat and, in some patients, a diastolic filling complex, as described previously.

The holosystolic murmur of chronic TR is generally softer than that of MR, is loudest at the left lower sternal border, and usually increases in intensity with inspiration (Carvallo's sign). Associated signs include *c-v* waves in the jugular venous pulse, an enlarged and pulsatile liver, ascites, and peripheral edema. The abnormal jugular venous waveforms are the predominant finding and are seen very often in the

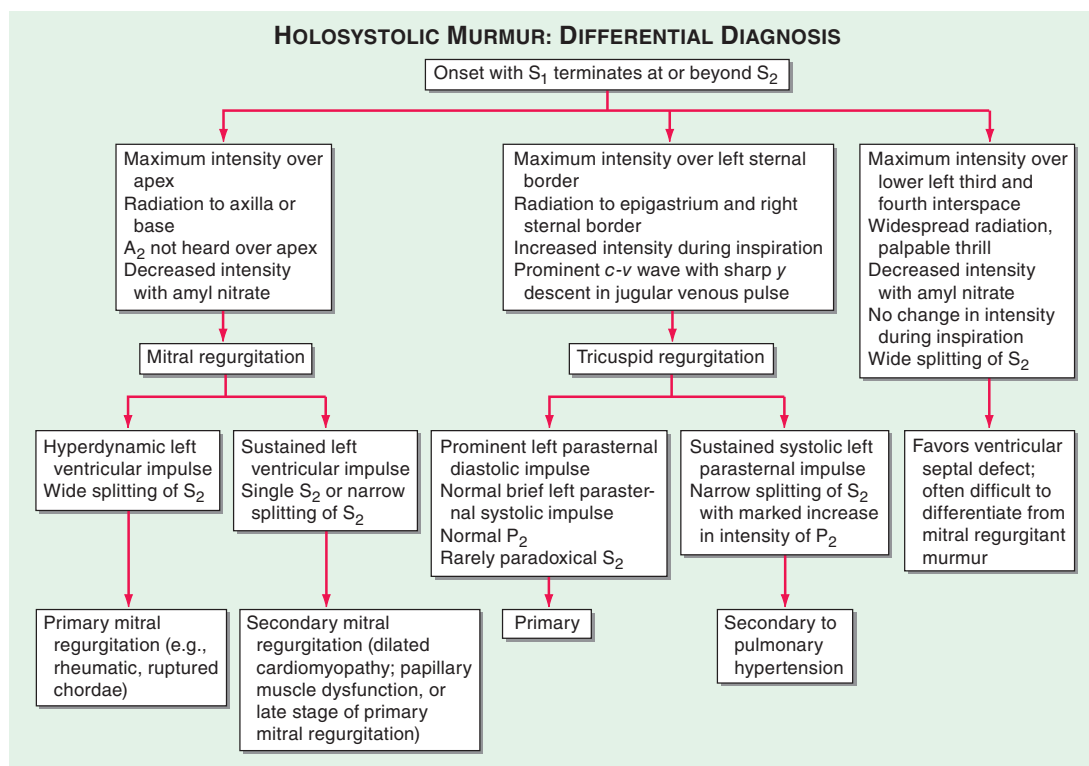


FIGURE 51e-5 Differential diagnosis of a holosystolic murmur.

absence of an audible murmur despite Doppler echocardiographic verification of TR. Causes of primary TR include myxomatous disease (prolapse), endocarditis, rheumatic disease, radiation, carcinoid, Ebstein's anomaly, and chordal detachment as a complication of right ventricular endomyocardial biopsy. TR is more commonly a passive process that results secondarily from annular enlargement due to right ventricular dilatation in the face of volume or pressure overload.

The holosystolic murmur of a VSD is loudest at the mid- to lower left sternal border (Fig. 51e-2) and radiates widely. A thrill is present at the site of maximal intensity in the majority of patients. There is no change in the intensity of the murmur with inspiration. The intensity of the murmur varies as a function of the anatomic size of the defect. Small, restrictive VSDs, as exemplified by the *maladie de Roger*, create a very loud murmur due to the significant and sustained systolic pressure gradient between the left and right ventricles. With large defects, the ventricular pressures tend to equalize, shunt flow is balanced, and a murmur is not appreciated. The distinction between post-MI ventricular septal rupture and MR has been reviewed previously.

DIASTOLIC HEART MURMURS

Early Diastolic Murmurs (Fig. 51e-1E) Chronic AR results in a high-pitched, blowing, decrescendo, early to mid-diastolic murmur that begins after the aortic component of S_2 (A_2) and is best heard at the second right interspace (Fig. 51e-6). The murmur may be soft and difficult to hear unless auscultation is performed with the patient leaning forward at end expiration. This maneuver brings the aortic root closer to the anterior chest wall. Radiation of the murmur may provide a clue to the cause of the AR. With primary valve disease, such as that due to congenital bicuspid disease, prolapse, or endocarditis, the diastolic murmur tends to radiate along the left sternal border, where it is often louder than appreciated in the second right interspace. When AR is caused by aortic root disease, the diastolic murmur may radiate along the right sternal border. Diseases of the aortic root cause dilation or distortion of the aortic annulus and failure of leaflet coaptation. Causes include Marfan syndrome with aneurysm formation, annuloaortic ectasia, ankylosing spondylitis, and aortic dissection.

Chronic, severe AR also may produce a lower-pitched mid to late, grade 1 or 2 diastolic murmur at the apex (Austin Flint murmur),

which is thought to reflect turbulence at the mitral inflow area from the admixture of regurgitant (aortic) and forward (mitral) blood flow (Fig. 51e-1G). This lower-pitched, apical diastolic murmur can be distinguished from that due to MS by the absence of an opening snap and the response of the murmur to a vasodilator challenge. Lowering afterload with an agent such as amyl nitrite will decrease the duration and magnitude of the aortic-left ventricular diastolic pressure gradient, and thus, the Austin Flint murmur of severe AR will become shorter and softer. The intensity of the diastolic murmur of mitral stenosis (Fig. 51e-6) may either remain constant or increase with afterload reduction because of the reflex increase in cardiac output and mitral valve flow.

Although AS and AR may coexist, a grade 2 or 3 crescendo-decrescendo mid-systolic murmur frequently is heard at the base of the heart in patients with isolated, severe AR and is due to an increased volume and rate of systolic flow. Accurate bedside identification of coexistent AS can be difficult unless the carotid pulse examination is abnormal or the mid-systolic murmur is of grade 4 or greater intensity. In the absence of heart failure, chronic severe AR is accompanied by several peripheral signs of significant diastolic run-off, including a wide pulse pressure, a "water-hammer" carotid upstroke (Corrigan's pulse), and Quincke's pulsations of the nail beds. The diastolic murmur of *acute, severe AR* is notably shorter in duration and lower pitched than the murmur of chronic AR. It can be very difficult to appreciate in the presence of a rapid heart rate. These attributes reflect the abrupt rate of rise of diastolic pressure within the unprepared and noncompliant left ventricle and the correspondingly rapid decline in the aortic-left ventricular diastolic pressure gradient. Left ventricular diastolic pressure may increase sufficiently to result in premature closure of the mitral valve and a soft first heart sound. Peripheral signs of significant diastolic run-off are not present.

Pulmonic regurgitation (PR) results in a decrescendo, early to mid-diastolic murmur (*Graham Steell murmur*) that begins after the pulmonic component of S_2 (P_2), is best heard at the second left interspace, and radiates along the left sternal border. The intensity of the murmur may increase with inspiration. PR is most commonly due to dilation of the valve annulus from chronic elevation of the pulmonary artery pressure. Signs of pulmonary hypertension, including a right ventricular

Diastolic Filling Murmur (Rumble) Mitral Stenosis

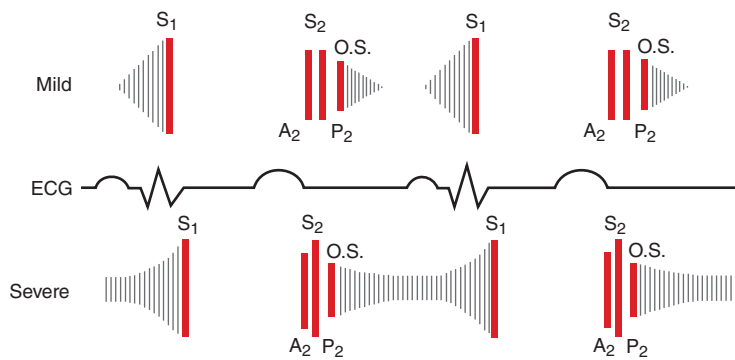


FIGURE 51e-6 Diastolic filling murmur (rumble) in mitral stenosis.

In mild mitral stenosis, the diastolic gradient across the valve is limited to the phases of rapid ventricular filling in early diastole and presystole. The rumble may occur during either or both periods. As the stenotic process becomes severe, a large pressure gradient exists across the valve during the entire diastolic filling period, and the rumble persists throughout diastole. As the left atrial pressure becomes greater, the interval between A_2 (or P_2) and the opening snap (O.S.) shortens. In severe mitral stenosis, secondary pulmonary hypertension develops and results in a loud P_2 and the splitting interval usually narrows. ECG, electrocardiogram. (From JA Shaver, JJ Leonard, DF Leon: *Examination of the Heart, Part IV, Auscultation of the Heart*. Dallas, American Heart Association, 1990, p 55. Copyright, American Heart Association.)

lift and a loud, single or narrowly split S_2 , are present. These features also help distinguish PR from AR as the cause of a decrescendo diastolic murmur heard along the left sternal border. PR in the absence of pulmonary hypertension can occur with endocarditis or a congenitally deformed valve. It is usually present after repair of tetralogy of Fallot in childhood. When pulmonary hypertension is not present, the diastolic murmur is softer and lower pitched than the classic Graham Steell murmur, and the severity of the PR can be difficult to appreciate.

TTE is indicated for the further evaluation of a patient with an early to mid-diastolic murmur. Longitudinal assessment of the severity of the valve lesion and ventricular size and systolic function help guide a potential decision for surgical management. TTE also can provide anatomic information regarding the root and proximal ascending aorta, although computed tomographic or magnetic resonance angiography may be indicated for more precise characterization (Chap. 270e).

Mid-Diastolic Murmurs (Figs. 51e-1G and 51e-1H) Mid-diastolic murmurs result from obstruction and/or augmented flow at the level of the mitral or tricuspid valve. Rheumatic fever is the most common cause of MS (Fig. 51e-6). In younger patients with pliable valves, S_1 is loud and the murmur begins after an opening snap, which is a high-pitched sound that occurs shortly after S_2 . The interval between the pulmonic component of the second heart sound (P_2) and the opening snap is inversely related to the magnitude of the left atrial–left ventricular pressure gradient. The murmur of MS is low-pitched and thus is best heard with the bell of the stethoscope. It is loudest at the left ventricular apex and often is appreciated only when the patient is turned in the left lateral decubitus position. It is usually of grade 1 or 2 intensity but may be absent when the cardiac output is severely reduced despite significant obstruction. The intensity of the murmur increases during maneuvers that increase cardiac output and mitral valve flow, such as exercise. The duration of the murmur reflects the length of time over which left atrial pressure exceeds left ventricular diastolic pressure. An increase in the intensity of the murmur just before S_1 , a phenomenon known as *presystolic accentuation* (Figs. 51e-1A and 51e-6), occurs in patients in sinus rhythm and is due to a late increase in transmitral flow with atrial contraction. Presystolic accentuation does not occur in patients with atrial fibrillation.

The mid-diastolic murmur associated with tricuspid stenosis is best heard at the lower left sternal border and increases in intensity with inspiration. A prolonged y descent may be visible in the jugular venous

waveform. This murmur is very difficult to hear and often is obscured by left-sided acoustical events.

There are several other causes of mid-diastolic murmurs. Large left atrial myxomas may prolapse across the mitral valve and cause variable degrees of obstruction to left ventricular inflow (Chap. 289e). The murmur associated with an atrial myxoma may change in duration and intensity with changes in body position. An opening snap is not present, and there is no presystolic accentuation. Augmented mitral diastolic flow can occur with isolated severe MR or with a large left-to-right shunt at the ventricular or great vessel level and produce a soft, rapid filling sound (S_3) followed by a short, low-pitched mid-diastolic apical murmur. The Austin Flint murmur of severe, chronic AR has already been described.

A short, mid-diastolic murmur is rarely heard during an episode of acute rheumatic fever (Carey-Coombs murmur) and probably is due to flow through an edematous mitral valve. An opening snap is not present in the acute phase, and the murmur dissipates with resolution of the acute attack. Complete heart block with dyssynchronous atrial and ventricular activation may be associated with intermittent mid- to late diastolic murmurs if atrial contraction occurs when the mitral valve is partially closed. Mid-diastolic murmurs indicative of increased tricuspid valve flow can occur with severe, isolated TR and with large ASDs and significant left-to-right shunting. Other signs of an ASD are present (Chap. 282), including fixed splitting of S_2 and a mid-systolic murmur at the mid- to upper left sternal border. TTE is indicated for evaluation of a patient with a mid- to late diastolic murmur. Findings specific to the diseases discussed above will help guide management.

CONTINUOUS MURMURS

(Figs. 51e-1H and 51e-7) Continuous murmurs begin in systole, peak near the second heart sound, and continue into all or part of diastole. Their presence throughout the cardiac cycle implies a pressure gradient between two chambers or vessels during both systole and diastole. The continuous murmur associated with a patent ductus arteriosus is best heard at the upper left sternal border. Large, uncorrected shunts may lead to pulmonary hypertension, attenuation or obliteration of the diastolic component of the murmur, reversal of shunt flow, and differential cyanosis of the lower extremities. A ruptured sinus of Valsalva aneurysm creates a continuous murmur of abrupt onset at the upper right sternal border. Rupture typically occurs into a right heart chamber, and the murmur is indicative of a continuous pressure difference between the aorta and either the right ventricle or the right atrium. A continuous murmur also may be audible along the left sternal border with a coronary arteriovenous fistula and at the site of an arteriovenous fistula used for hemodialysis access. Enhanced flow through enlarged intercostal collateral arteries in patients with aortic coarctation may produce a continuous murmur along the course of one or more ribs. A cervical bruit with both systolic and diastolic components (a to-fro murmur, Fig. 51e-7) usually indicates a high-grade carotid artery stenosis.

Not all continuous murmurs are pathologic. A continuous venous hum can be heard in healthy children and young adults, especially during pregnancy; it is best appreciated in the right supraclavicular fossa and can be obliterated by pressure over the right internal jugular vein or by having the patient turn his or her head toward the examiner. The continuous mammary souffle of pregnancy is created by enhanced arterial flow through engorged breasts and usually appears during the late third trimester or early puerperium. The murmur is louder in systole. Firm pressure with the diaphragm of the stethoscope can eliminate the diastolic portion of the murmur.

DYNAMIC AUSCULTATION

(Table 51e-2; see Table 267-1) Careful attention to the behavior of heart murmurs during simple maneuvers that alter cardiac hemodynamics can provide important clues to their cause and significance.

Respiration Auscultation should be performed during quiet respiration or with a modest increase in inspiratory effort, as more forceful

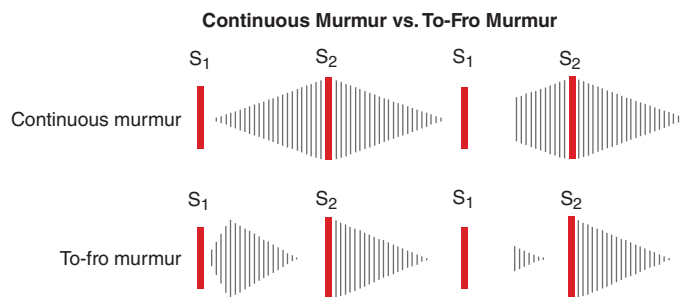


FIGURE 51e-7 Comparison of the continuous murmur and the to-fro murmur. During abnormal communication between high-pressure and low-pressure systems, a large pressure gradient exists throughout the cardiac cycle, producing a continuous murmur. A classic example is patent ductus arteriosus. At times, this type of murmur can be confused with a to-fro murmur, which is a combination of systolic ejection murmur and a murmur of semilunar valve incompetence. A classic example of a to-fro murmur is aortic stenosis and regurgitation. A continuous murmur crescendos to near the second heart sound (S_2), whereas a to-fro murmur has two components. The mid-systolic ejection component decrescendos and disappears as it approaches S_2 . (From JA Shaver, JJ Leonard, DF Leon: *Examination of the Heart, Part IV, Auscultation of the Heart*. Dallas, American Heart Association, 1990, p 55. Copyright, American Heart Association.)

movement of the chest tends to obscure the heart sounds. Left-sided murmurs may be best heard at end expiration, when lung volumes are minimized and the heart and great vessels are brought closer to the chest wall. This phenomenon is characteristic of the murmur of AR. Murmurs of right-sided origin, such as tricuspid or pulmonic regurgitation, increase in intensity during inspiration. The intensity of left-sided murmurs either remains constant or decreases with inspiration.

Beside assessment also should evaluate the behavior of S_2 with respiration and the dynamic relationship between the aortic and pulmonic components (Fig. 51e-8). Reversed splitting can be a feature of severe AS, HOCM, left bundle branch block, right ventricular pacing, or acute myocardial ischemia. Fixed splitting of S_2 in the presence of a grade 2 or 3 mid-systolic murmur at the mid- or upper left sternal border indicates an ASD. Physiologic but wide splitting during the respiratory cycle implies either premature aortic valve closure, as can occur with severe MR, or delayed pulmonic valve closure due to PS or right bundle branch block.

Alterations of Systemic Vascular Resistance Murmurs can change characteristics after maneuvers that alter systemic vascular resistance and left ventricular afterload. The systolic murmurs of MR and VSD become louder during sustained handgrip, simultaneous inflation of blood pressure cuffs on both upper extremities to pressures 20–40 mmHg above systolic pressure for 20 s, or infusion of a vasopressor agent. The murmurs associated with AS or HOCM will become softer or remain unchanged with these maneuvers. The diastolic murmur of AR becomes louder in response to interventions that raise systemic vascular resistance.

Opposite changes in systolic and diastolic murmurs may occur with the use of pharmacologic agents that lower systemic vascular resistance.

TABLE 51e-2 DYNAMIC AUSCULTATION: BEDSIDE MANEUVERS THAT CAN BE USED TO CHANGE THE INTENSITY OF CARDIAC MURMURS (SEE TEXT)

1. Respiration
2. Isometric exercise (handgrip)
3. Transient arterial occlusion
4. Pharmacologic manipulation of preload and/or afterload
5. Valsalva maneuver
6. Rapid standing/squatting
7. Passive leg raising
8. Post-premature beat

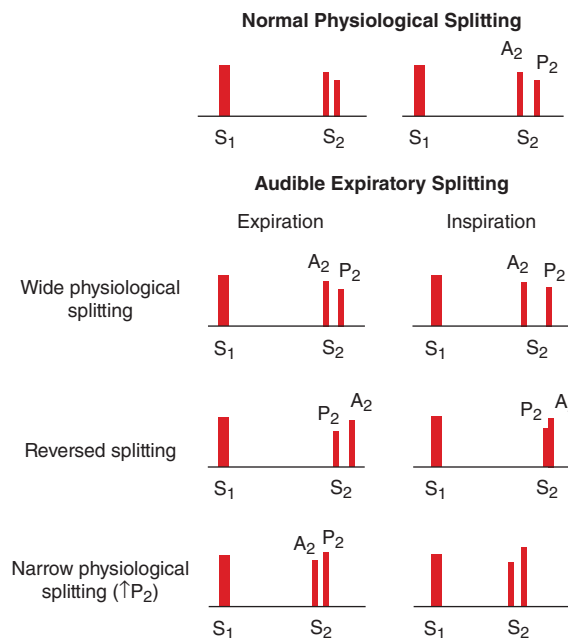


FIGURE 51e-8 Top. Normal physiologic splitting. During expiration, the aortic (A_2) and pulmonic (P_2) components of the second heart sound are separated by <30 ms and are appreciated as a single sound. During inspiration, the splitting interval widens, and A_2 and P_2 are clearly separated into two distinct sounds. Bottom. Audible expiratory splitting. Wide physiologic splitting is caused by a delay in P_2 . Reversed splitting is caused by a delay in A_2 , resulting in paradoxical movement; i.e., with inspiration P_2 moves toward A_2 , and the splitting interval narrows. Narrow physiologic splitting occurs in pulmonary hypertension, and both A_2 and P_2 are heard during expiration at a narrow splitting interval because of the increased intensity and high-frequency composition of P_2 . (From JA Shaver, JJ Leonard, DF Leon: *Examination of the Heart, Part IV, Auscultation of the Heart*. Dallas, American Heart Association, 1990, p 17. Copyright, American Heart Association.)

Inhaled amyl nitrite is now rarely used for this purpose but can help distinguish the murmur of AS or HOCM from that of either MR or VSD, if necessary. The former two murmurs increase in intensity, whereas the latter two become softer after exposure to amyl nitrite. As noted previously, the Austin Flint murmur of severe AR becomes softer, but the mid-diastolic rumble of MS becomes louder, in response to the abrupt lowering of systemic vascular resistance with amyl nitrite.

Changes in Venous Return The Valsalva maneuver results in an increase in intrathoracic pressure, followed by a decrease in venous return, ventricular filling, and cardiac output. The majority of murmurs decrease in intensity during the strain phase of the maneuver. Two notable exceptions are the murmurs associated with MVP and obstructive HOCM, both of which become louder during the Valsalva maneuver. The murmur of MVP may also become longer as leaflet prolapse occurs earlier in systole at smaller ventricular volumes. These murmurs behave in a similar and parallel fashion with standing. Both the click and the murmur of MVP move closer in timing to S_1 on rapid standing from a squatting position (Fig. 51e-3). The increase in the intensity of the murmur of HOCM is predicated on the augmentation of the dynamic left ventricular outflow tract gradient that occurs with reduced ventricular filling. Squatting results in abrupt increases in both venous return (preload) and left ventricular afterload that increase ventricular volume, changes that predictably cause a decrease in the intensity and duration of the murmurs associated with MVP and HOCM; the click and murmur of MVP move away from S_1 with squatting. Passive leg raising can be used to increase venous return in patients who are unable to squat and stand. This maneuver may lead to a decrease in the intensity of the murmur associated with HOCM but has less effect in patients with MVP.

Post-Premature Ventricular Contraction A change in the intensity of a systolic murmur in the first beat after a premature beat, or in the beat after a long cycle length in patients with atrial fibrillation, can help distinguish AS from MR, particularly in an older patient in whom the murmur of AS is well transmitted to the apex. Systolic murmurs due to left ventricular outflow obstruction, including that due to AS, increase in intensity in the beat after a premature beat because of the combined effects of enhanced left ventricular filling and post-extrasystolic potentiation of contractile function. Forward flow accelerates, causing an increase in the gradient and a louder murmur. The intensity of the murmur of MR does not change in the post-premature beat as there is relatively little further increase in mitral valve flow or change in the left ventricular–left atrial gradient.

THE CLINICAL CONTEXT

Additional clues to the etiology and importance of a heart murmur can be gleaned from the history and other physical examination findings. Symptoms suggestive of cardiovascular, neurologic, or pulmonary disease help focus the differential diagnosis, as do findings relevant to the jugular venous pressure and waveforms, the arterial pulses, other heart sounds, the lungs, the abdomen, the skin, and the extremities. In many instances, laboratory studies, an ECG, and/or a chest x-ray may have been obtained earlier and may contain valuable information. A patient with suspected infective endocarditis, for example, may have a murmur in the setting of fever, chills, anorexia, fatigue, dyspnea, splenomegaly, petechiae, and positive blood cultures. A new systolic murmur in a patient with a marked fall in blood pressure after a recent MI suggests myocardial rupture. By contrast, an isolated grade 1 or 2 mid-systolic murmur at the left sternal border in a healthy, active, and asymptomatic young adult is most likely a benign finding for which no further evaluation is indicated. The context in which the murmur is appreciated often dictates the need for further testing.

ECHOCARDIOGRAPHY

(Fig. 51e-9; Chaps. 267 and 270e) Echocardiography with color flow and spectral Doppler is a valuable tool for the assessment of cardiac murmurs. Information regarding valve structure and function, chamber size, wall thickness, ventricular function, estimated pulmonary artery pressures, intracardiac shunt flow, pulmonary and hepatic vein flow, and aortic flow can be ascertained readily. It is important to note that Doppler signals of trace or mild valvular regurgitation of no clinical consequence can be detected with structurally normal tricuspid,

pulmonic, and mitral valves. Such signals are not likely to generate enough turbulence to create an audible murmur.

Echocardiography is indicated for the evaluation of patients with early, late, or holosystolic murmurs and patients with grade 3 or louder mid-systolic murmurs. Patients with grade 1 or 2 mid-systolic murmurs but other symptoms or signs of cardiovascular disease, including those from ECG or chest x-ray, should also undergo echocardiography. Echocardiography is also indicated for the evaluation of any patient with a diastolic murmur and for patients with continuous murmurs not due to a venous hum or mammary souffle. Echocardiography also should be considered when there is a clinical need to verify normal cardiac structure and function in a patient whose symptoms and signs are probably noncardiac in origin. The performance of serial echocardiography to follow the course of asymptomatic individuals with valvular heart disease is a central feature of their longitudinal assessment and provides valuable information that may have an impact on decisions regarding the timing of surgery. Routine echocardiography is *not* recommended for asymptomatic patients with a grade 1 or 2 mid-systolic murmur without other signs of heart disease. For this category of patients, referral to a cardiovascular specialist should be considered if there is doubt about the significance of the murmur after the initial examination.

The selective use of echocardiography outlined above has not been subjected to rigorous analysis of its cost-effectiveness. For some clinicians, handheld or miniaturized cardiac ultrasound devices have replaced the stethoscope. Although several reports attest to the improved sensitivity of such devices for the detection of valvular heart disease, accuracy is highly operator-dependent, and incremental cost considerations and outcomes have not been addressed adequately. The use of electronic or digital stethoscopes with spectral display capabilities has also been proposed as a method to improve the characterization of heart murmurs and the mentored teaching of cardiac auscultation.

OTHER CARDIAC TESTING

(Chap. 270e, Fig. 51e-9) In relatively few patients, clinical assessment and TTE do not adequately characterize the origin and significance of a heart murmur. Transesophageal echocardiography (TEE) can be considered for further evaluation, especially when the TTE windows are limited by body size, chest configuration, or intrathoracic pathology. TEE offers enhanced sensitivity for the detection of a wide range of structural cardiac disorders. Electrocardiographically gated cardiac magnetic resonance (CMR) imaging, although limited in its ability to display valvular morphology, can provide quantitative information regarding valvular function, stenosis severity, regurgitant fraction, regurgitant volume, shunt flow, chamber and great vessel size, ventricular function, and myocardial perfusion. CMR has largely supplanted the need for cardiac catheterization and invasive hemodynamic assessment when there is a discrepancy between the clinical and echocardiographic findings. Invasive coronary angiography is performed routinely in most adult patients before valve surgery, especially when there is a suspicion of coronary artery disease predicated on symptoms, risk factors, and/or age. The use of computed tomography coronary angiography (CCTA) to exclude coronary artery disease in patients with a low pretest probability of disease before valve surgery is gaining wider acceptance.

INTEGRATED APPROACH

The accurate identification of a heart murmur begins with a systematic approach to cardiac auscultation. Characterization of its major attributes, as reviewed above, allows the examiner to construct a preliminary differential diagnosis, which is then refined by integration of information available from the history, associated cardiac findings, the

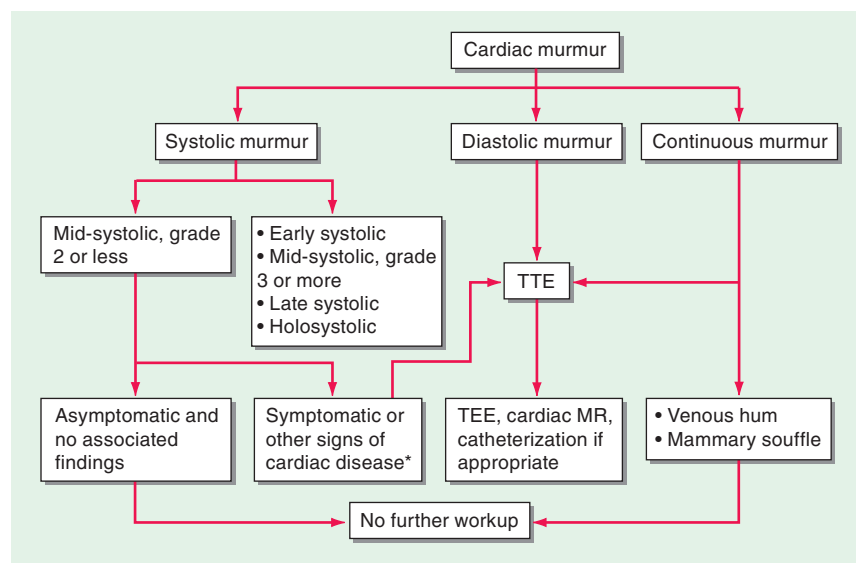


FIGURE 51e-9 Strategy for evaluating heart murmurs. *If an electrocardiogram or chest x-ray has been obtained and is abnormal, echocardiography is indicated. TTE, transthoracic echocardiography; TEE, transesophageal echocardiography; MR, magnetic resonance. (Adapted from RO Bonow et al: *J Am Coll Cardiol* 32:1486, 1998.)

general physical examination, and the clinical context. The need for and urgency of further testing follow sequentially. Correlation of the findings on auscultation with the noninvasive data provides an educational feedback loop and an opportunity for improving physical examination skills. Cost constraints mandate that noninvasive imaging

be justified on the basis of its incremental contribution to diagnosis, treatment, and outcome. Additional study is required to assess the cost-effective application of newer imaging technology in patients with heart murmurs.

Palpitations are extremely common among patients who present to their internists and can best be defined as an intermittent “thumping,” “pounding,” or “fluttering” sensation in the chest. This sensation can be either intermittent or sustained and either regular or irregular. Most patients interpret palpitations as an unusual awareness of the heartbeat and become especially concerned when they sense that they have had “skipped” or “missing” heartbeats. Palpitations are often noted when the patient is quietly resting, during which time other stimuli are minimal. Palpitations that are positional generally reflect a structural process within (e.g., atrial myxoma) or adjacent to (e.g., mediastinal mass) the heart.

Palpitations are brought about by cardiac (43%), psychiatric (31%), miscellaneous (10%), and unknown (16%) causes, according to one large series. Among the cardiovascular causes are premature atrial and ventricular contractions, supraventricular and ventricular arrhythmias, mitral valve prolapse (with or without associated arrhythmias), aortic insufficiency, atrial myxoma, and pulmonary embolism. Intermittent palpitations are commonly caused by premature atrial or ventricular contractions: the post-extrasystolic beat is sensed by the patient owing to the increase in ventricular end-diastolic dimension following the pause in the cardiac cycle and the increased strength of contraction (post-extrasystolic potentiation) of that beat. Regular, sustained palpitations can be caused by regular supraventricular and ventricular tachycardias. Irregular, sustained palpitations can be caused by atrial fibrillation. It is important to note that most arrhythmias are not associated with palpitations. In those that are, it is often useful either to ask the patient to “tap out” the rhythm of the palpitations or to take his/her pulse during palpitations. In general, hyperdynamic cardiovascular states caused by catecholaminergic stimulation from exercise, stress, or pheochromocytoma can lead to palpitations. Palpitations are common among athletes, especially older endurance athletes. In addition, the enlarged ventricle of aortic regurgitation and accompanying hyperdynamic precordium frequently lead to the sensation of palpitations. Other factors that enhance the strength of myocardial contraction, including tobacco, caffeine, aminophylline, atropine, thyroxine, cocaine, and amphetamines, can cause palpitations.

Psychiatric causes of palpitations include panic attacks or disorders, anxiety states, and somatization, alone or in combination. Patients with psychiatric causes for palpitations more commonly report a longer duration of the sensation (>15 min) and other accompanying symptoms than do patients with other causes. Among the miscellaneous causes of palpitations are thyrotoxicosis, drugs (see above) and ethanol, spontaneous skeletal muscle contractions of the chest wall, pheochromocytoma, and systemic mastocytosis.

APPROACH TO THE PATIENT:

Palpitations

The principal goal in assessing patients with palpitations is to determine whether the symptom is caused by a life-threatening arrhythmia. Patients with preexisting coronary artery disease (CAD) or risk factors for CAD are at greatest risk for ventricular arrhythmias (Chap. 276) as a cause for palpitations. In addition, the association of palpitations with other symptoms suggesting hemodynamic compromise, including syncope or lightheadedness, supports this diagnosis. Palpitations caused by sustained tachyarrhythmias in patients with CAD can be accompanied by angina pectoris or dyspnea, and, in patients with ventricular dysfunction (systolic or diastolic), aortic stenosis, hypertrophic cardiomyopathy, or mitral stenosis (with or without CAD), can be accompanied by dyspnea from increased left atrial and pulmonary venous pressure.

Key features of the physical examination that will help confirm or refute the presence of an arrhythmia as a cause for palpitations (as well as its adverse hemodynamic consequences) include measurement of the vital signs, assessment of the jugular venous pressure and pulse, and auscultation of the chest and precordium. A resting electrocardiogram can be used to document the arrhythmia. If exertion is known to induce the arrhythmia and accompanying palpitations, exercise electrocardiography can be used to make the diagnosis. If the arrhythmia is sufficiently infrequent, other methods must be used, including continuous electrocardiographic (Holter) monitoring; telephonic monitoring, through which the patient can transmit an electrocardiographic tracing during a sensed episode; loop recordings (external or implantable), which can capture the electrocardiographic event for later review; and mobile cardiac outpatient telemetry. Data suggest that Holter monitoring is of limited clinical utility, while the implantable loop recorder and mobile cardiac outpatient telemetry are safe and possibly more cost-effective in the assessment of patients with (infrequent) recurrent, unexplained palpitations.

Most patients with palpitations do not have serious arrhythmias or underlying structural heart disease. If sufficiently troubling to the patient, occasional benign atrial or ventricular premature contractions can often be managed with beta-blocker therapy. Palpitations incited by alcohol, tobacco, or illicit drugs need to be managed by abstinence, while those caused by pharmacologic agents should be addressed by considering alternative therapies when appropriate or possible. Psychiatric causes of palpitations may benefit from cognitive therapy or pharmacotherapy. The physician should note that palpitations are at the very least bothersome and, on occasion, frightening to the patient. Once serious causes for the symptom have been excluded, the patient should be reassured that the palpitations will not adversely affect prognosis.

SECTION 6

ALTERATIONS IN GASTROINTESTINAL FUNCTION

Dysphagia—difficulty with swallowing—refers to problems with the transit of food or liquid from the mouth to the hypopharynx or through the esophagus. Severe dysphagia can compromise nutrition, cause aspiration, and reduce quality of life. Additional terminology pertaining to swallowing dysfunction is as follows. *Aphagia* (inability to swallow) typically denotes complete esophageal obstruction, most

commonly encountered in the acute setting of a food bolus or foreign body impaction. *Odynophagia* refers to painful swallowing, typically resulting from mucosal ulceration within the oropharynx or esophagus. It commonly is accompanied by dysphagia, but the converse is not true. *Globus pharyngeus* is a foreign body sensation localized in the neck that does not interfere with swallowing and sometimes is relieved by swallowing. *Transfer dysphagia* frequently results in nasal regurgitation and pulmonary aspiration during swallowing and is characteristic of oropharyngeal dysphagia. *Phagophobia* (fear of swallowing) and *refusal to swallow* may be psychogenic or related to anticipatory anxiety about food bolus obstruction, odynophagia, or aspiration.

PHYSIOLOGY OF SWALLOWING

Swallowing begins with a voluntary (oral) phase that includes preparation during which food is masticated and mixed with saliva. This is followed by a transfer phase during which the bolus is pushed into the pharynx by the tongue. Bolus entry into the hypopharynx initiates the pharyngeal swallow response, which is centrally mediated and involves a complex series of actions, the net result of which is to propel food through the pharynx into the esophagus while preventing its entry into the airway. To accomplish this, the larynx is elevated and pulled forward, actions that also facilitate upper esophageal sphincter (UES) opening. Tongue pulsion then propels the bolus through the UES, followed by a peristaltic contraction that clears residue from the pharynx and through the esophagus. The lower esophageal sphincter (LES) relaxes as the food enters the esophagus and remains relaxed until the peristaltic contraction has delivered the bolus into the stomach. Peristaltic contractions elicited in response to a swallow are called *primary peristalsis* and involve sequenced inhibition followed by contraction of the musculature along the entire length of the esophagus. The inhibition that precedes the peristaltic contraction is called *deglutitive inhibition*. Local distention of the esophagus anywhere along its length, as may occur with gastroesophageal reflux, activates *secondary peristalsis* that begins at the point of distention and proceeds distally. Tertiary esophageal contractions are nonperistaltic, disordered esophageal contractions that may be observed to occur spontaneously during fluoroscopic observation.

The musculature of the oral cavity, pharynx, UES, and cervical esophagus is striated and directly innervated by lower motor neurons carried in cranial nerves (Fig. 53-1). Oral cavity muscles are innervated by the fifth (trigeminal) and seventh (facial) cranial nerves; the tongue, by the twelfth (hypoglossal) cranial nerve. Pharyngeal muscles are innervated by the ninth (glossopharyngeal) and tenth (vagus) cranial nerves.

Physiologically, the UES consists of the cricopharyngeus muscle, the adjacent inferior pharyngeal constrictor, and the proximal portion of the cervical esophagus. UES innervation is derived from the vagus nerve, whereas the innervation to the musculature acting on the UES to facilitate its opening during swallowing comes from the fifth, seventh, and twelfth cranial nerves. The UES remains closed at rest owing to both its inherent elastic properties and neurogenically mediated contraction of the cricopharyngeus muscle. UES opening during

swallowing involves both cessation of vagal excitation to the cricopharyngeus and simultaneous contraction of the suprahyoid and geniohyoid muscles that pull open the UES in conjunction with the upward and forward displacement of the larynx.

The neuromuscular apparatus for peristalsis is distinct in proximal and distal parts of the esophagus. The cervical esophagus, like the pharyngeal musculature, consists of striated muscle and is directly innervated by lower motor neurons of the vagus nerve. Peristalsis in the proximal esophagus is governed by the sequential activation of the vagal motor neurons in the nucleus ambiguus. In contrast, the distal esophagus and LES are composed of smooth muscle and are controlled by excitatory and inhibitory neurons within the esophageal myenteric plexus. Medullary preganglionic neurons from the dorsal motor nucleus of the vagus trigger peristalsis via these ganglionic neurons during primary peristalsis. Neurotransmitters of the excitatory ganglionic neurons are acetylcholine and substance P; those of the inhibitory neurons are vasoactive intestinal peptide and nitric oxide. Peristalsis results from the patterned activation of inhibitory followed by excitatory ganglionic neurons, with progressive dominance of the inhibitory neurons distally. Similarly, LES relaxation occurs with the onset of deglutitive inhibition and persists until the peristaltic sequence is complete. At rest, the LES is contracted because of excitatory ganglionic stimulation and its intrinsic myogenic tone, a property that distinguishes it from the adjacent esophagus. The function of the LES is supplemented by the surrounding muscle of the right diaphragmatic crus, which acts as an external sphincter during inspiration, cough, or abdominal straining.

PATHOPHYSIOLOGY OF DYSPHAGIA

Dysphagia can be subclassified both by location and by the circumstances in which it occurs. With respect to location, distinct considerations apply to oral, pharyngeal, or esophageal dysphagia. Normal transport of an ingested bolus depends on the consistency and size of the bolus, the caliber of the lumen, the integrity of peristaltic contraction, and deglutitive inhibition of both the UES and the LES. Dysphagia caused by an oversized bolus or a narrow lumen is called *structural dysphagia*, whereas dysphagia due to abnormalities of peristalsis or impaired sphincter relaxation after swallowing is called *propulsive* or

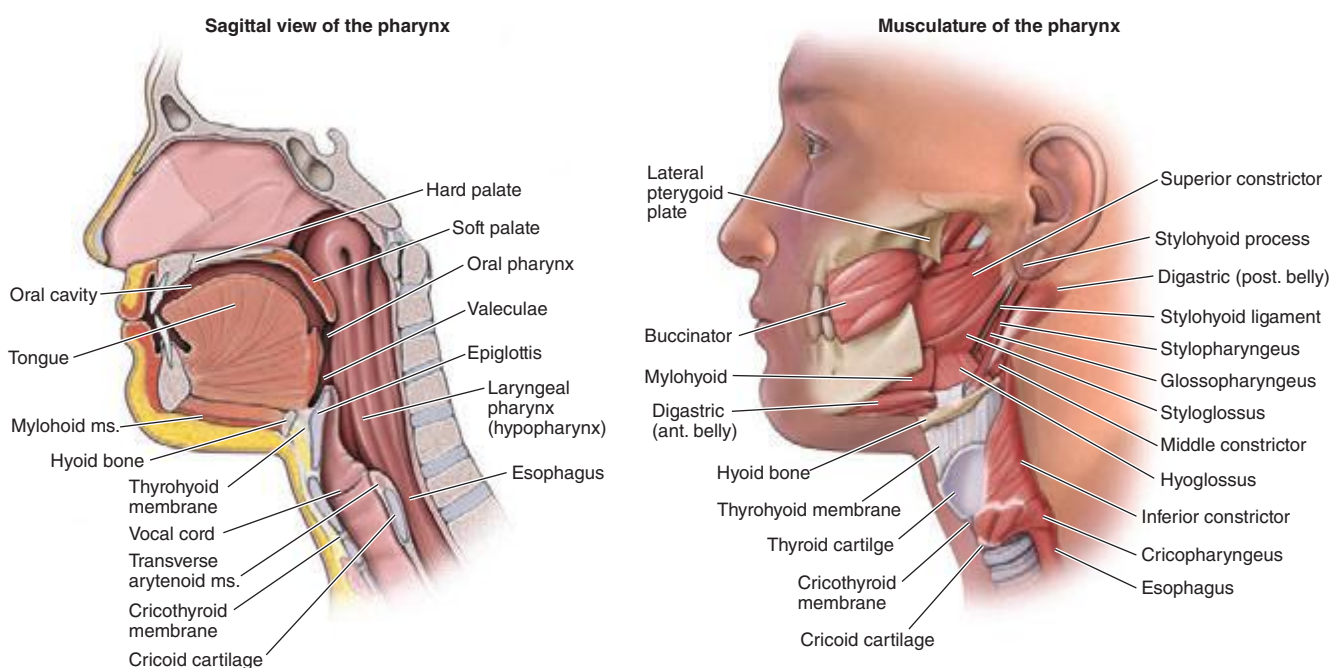


FIGURE 53-1 Sagittal and diagrammatic views of the musculature involved in enacting oropharyngeal swallowing. Note the dominance of the tongue in the sagittal view and the intimate relationship between the entrance to the larynx (airway) and the esophagus. In the resting configuration illustrated, the esophageal inlet is closed. This is transiently reconfigured such that the esophageal inlet is open and the laryngeal inlet closed during swallowing. (Adapted from PJ Kahrilas, in DW Gelfand and JE Richter [eds]: *Dysphagia: Diagnosis and Treatment*. New York: Igaku-Shoin Medical Publishers, 1989, pp. 11–28.)

motor dysphagia. More than one mechanism may be operative in a patient with dysphagia. Scleroderma commonly presents with absent peristalsis as well as a weakened LES that predisposes patients to peptic stricture formation. Likewise, radiation therapy for head and neck cancer may compound the functional deficits in the oropharyngeal swallow attributable to the tumor and cause cervical esophageal stenosis.

Oral and Pharyngeal (Oropharyngeal) Dysphagia Oral-phase dysphagia is associated with poor bolus formation and control so that food has prolonged retention within the oral cavity and may seep out of the mouth. Drooling and difficulty in initiating swallowing are other characteristic signs. Poor bolus control also may lead to premature spillage of food into the hypopharynx with resultant aspiration into the trachea or regurgitation into the nasal cavity. Pharyngeal-phase dysphagia is associated with retention of food in the pharynx due to poor tongue or pharyngeal propulsion or obstruction at the UES. Signs and symptoms of concomitant hoarseness or cranial nerve dysfunction may be associated with oropharyngeal dysphagia.

Oropharyngeal dysphagia may be due to neurologic, muscular, structural, iatrogenic, infectious, and metabolic causes. Iatrogenic, neurologic, and structural pathologies are most common. Iatrogenic causes include surgery and radiation, often in the setting of head and neck cancer. Neurogenic dysphagia resulting from cerebrovascular accidents, Parkinson's disease, and amyotrophic lateral sclerosis is a major source of morbidity related to aspiration and malnutrition. Medullary nuclei directly innervate the oropharynx. Lateralization of pharyngeal dysphagia implies either a structural pharyngeal lesion or a neurologic process that selectively targeted the ipsilateral brainstem nuclei or cranial nerve. Advances in functional brain imaging have elucidated an important role of the cerebral cortex in swallow function and dysphagia. Asymmetry in the cortical representation of the pharynx provides an explanation for the dysphagia that occurs as a consequence of unilateral cortical cerebrovascular accidents.

Oropharyngeal structural lesions causing dysphagia include Zenker's diverticulum, cricopharyngeal bar, and neoplasia. Zenker's diverticulum typically is encountered in elderly patients, with an estimated prevalence between 1:1000 and 1:10,000. In addition to dysphagia, patients may present with regurgitation of particulate food debris, aspiration, and halitosis. The pathogenesis is related to stenosis of the cricopharyngeus that causes diminished opening of the UES and results in increased hypopharyngeal pressure during swallowing with development of a pulsion diverticulum immediately above the cricopharyngeus in a region of potential weakness known as Killian's dehiscence. A cricopharyngeal bar, appearing as a prominent indentation behind the lower third of the cricoid cartilage, is related to Zenker's diverticulum in that it involves limited distensibility of the cricopharyngeus and can lead to the formation of a Zenker's diverticulum. However, a cricopharyngeal bar is a common radiographic finding, and most patients with transient cricopharyngeal bars are asymptomatic, making it important to rule out alternative etiologies of dysphagia before treatment. Furthermore, cricopharyngeal bars may be secondary to other neuromuscular disorders.

Since the pharyngeal phase of swallowing occurs in less than a second, rapid-sequence fluoroscopy is necessary to evaluate for functional abnormalities. Adequate fluoroscopic examination requires that the patient be conscious and cooperative. The study incorporates recordings of swallow sequences during ingestion of food and liquids of varying consistencies. The pharynx is examined to detect bolus retention, regurgitation into the nose, or aspiration into the trachea. Timing and integrity of pharyngeal contraction and opening of the UES with a swallow are analyzed to assess both aspiration risk and the potential for swallow therapy. Structural abnormalities of the oropharynx, especially those which may require biopsies, also should be assessed by direct laryngoscopic examination.

Esophageal Dysphagia The adult esophagus measures 18–26 cm in length and is anatomically divided into the cervical esophagus, extending from the pharyngoesophageal junction to the suprasternal notch, and the thoracic esophagus, which continues to the diaphragmatic hiatus. When distended, the esophageal lumen has internal dimensions of

about 2 cm in the anteroposterior plane and 3 cm in the lateral plane. Solid food dysphagia becomes common when the lumen is narrowed to <13 mm but also can occur with larger diameters in the setting of poorly masticated food or motor dysfunction. Circumferential lesions are more likely to cause dysphagia than are lesions that involve only a partial circumference of the esophageal wall. The most common structural causes of dysphagia are Schatzki's rings, eosinophilic esophagitis, and peptic strictures. Dysphagia also occurs in the setting of gastroesophageal reflux disease without a stricture, perhaps on the basis of altered esophageal sensation, distensibility, or motor function.

Propulsive disorders leading to esophageal dysphagia result from abnormalities of peristalsis and/or deglutitive inhibition, potentially affecting the cervical or thoracic esophagus. Since striated muscle pathology usually involves both the oropharynx and the cervical esophagus, the clinical manifestations usually are dominated by oropharyngeal dysphagia. Diseases affecting smooth muscle involve both the thoracic esophagus and the LES. A dominant manifestation of this, absent peristalsis, refers to either the complete absence of swallow-induced contraction or the presence of nonperistaltic, disordered contractions. Absent peristalsis and failure of deglutitive LES relaxation are the defining features of achalasia. In diffuse esophageal spasm (DES), LES function is normal, with the disordered motility restricted to the esophageal body. Absent peristalsis combined with severe weakness of the LES is a nonspecific pattern commonly found in patients with scleroderma.

APPROACH TO THE PATIENT: Dysphagia

Figure 53-2 shows an algorithm for the approach to a patient with dysphagia.

HISTORY

The patient history is extremely valuable in making a presumptive diagnosis or at least substantially restricting the differential diagnoses in most patients. Key elements of the history are the localization of dysphagia, the circumstances in which dysphagia is experienced, other symptoms associated with dysphagia, and progression. Dysphagia that localizes to the suprasternal notch may indicate either an oropharyngeal or an esophageal etiology as distal dysphagia is referred proximally about 30% of the time. Dysphagia that localizes to the chest is esophageal in origin. Nasal regurgitation and tracheobronchial aspiration manifest by coughing with swallowing are hallmarks of oropharyngeal dysphagia. Severe cough with swallowing may also be a sign of a tracheoesophageal fistula. The presence of hoarseness may be another important diagnostic clue. When hoarseness precedes dysphagia, the primary lesion is usually laryngeal; hoarseness that occurs after the development of dysphagia may result from compromise of the recurrent laryngeal nerve by a malignancy. The type of food causing dysphagia is a crucial detail. Intermittent dysphagia that occurs only with solid food implies structural dysphagia, whereas constant dysphagia with both liquids and solids strongly suggests a motor abnormality. Two caveats to this pattern are that despite having a motor abnormality, patients with scleroderma generally develop mild dysphagia for solids only and, somewhat paradoxically, that patients with oropharyngeal dysphagia often have greater difficulty managing liquids than solids. Dysphagia that is progressive over the course of weeks to months raises concern for neoplasia. Episodic dysphagia to solids that is unchanged over years indicates a benign disease process such as a Schatzki's ring or eosinophilic esophagitis. Food impaction with a prolonged inability to pass an ingested bolus even with ingestion of liquid is typical of a structural dysphagia. Chest pain frequently accompanies dysphagia whether it is related to motor disorders, structural disorders, or reflux disease. A prolonged history of heartburn preceding the onset of dysphagia is suggestive of peptic stricture and, infrequently, esophageal adenocarcinoma. A history of prolonged nasogastric intubation, esophageal or head and neck surgery, ingestion of caustic

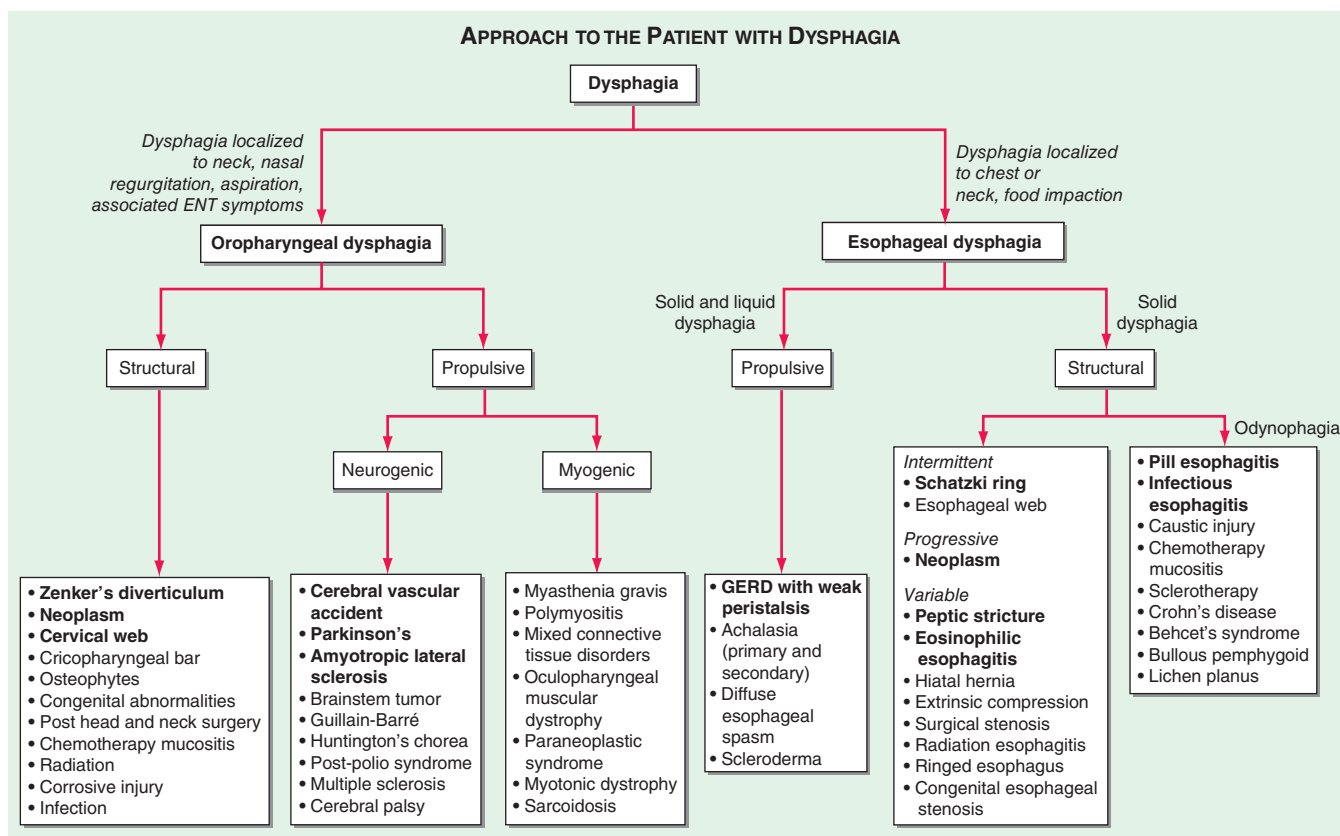


FIGURE 53-2 Approach to the patient with dysphagia. Etiologies in bold print are the most common. ENT, ear, nose, and throat; GERD, gastroesophageal reflux disease.

agents or pills, previous radiation or chemotherapy, or associated mucocutaneous diseases may help isolate the cause of dysphagia. With accompanying odynophagia, which usually is indicative of ulceration, infectious or pill-induced esophagitis should be suspected. In patients with AIDS or other immunocompromised states, esophagitis due to opportunistic infections such as *Candida*, herpes simplex virus, or cytomegalovirus and to tumors such as Kaposi's sarcoma and lymphoma should be considered. A strong history of atopy increases concerns for eosinophilic esophagitis.

PHYSICAL EXAMINATION

Physical examination is important in the evaluation of oral and pharyngeal dysphagia because dysphagia is usually only one of many manifestations of a more global disease process. Signs of bulbar or pseudobulbar palsy, including dysarthria, dysphonia, ptosis, tongue atrophy, and hyperactive jaw jerk, in addition to evidence of generalized neuromuscular disease, should be elicited. The neck should be examined for thyromegaly. A careful inspection of the mouth and pharynx should disclose lesions that may interfere with passage of food. Missing dentition can interfere with mastication and exacerbate an existing cause of dysphagia. Physical examination is less helpful in the evaluation of esophageal dysphagia as most relevant pathology is restricted to the esophagus. The notable exception is skin disease. Changes in the skin may suggest a diagnosis of scleroderma or mucocutaneous diseases such as pemphigoid, lichen planus and epidermolysis bullosa, all of which can involve the esophagus.

DIAGNOSTIC PROCEDURES

Although most instances of dysphagia are attributable to benign disease processes, dysphagia is also a cardinal symptom of several malignancies, making it an important symptom to evaluate. Cancer may result in dysphagia due to intraluminal obstruction (esophageal or proximal gastric cancer, metastatic deposits),

extrinsic compression (lymphoma, lung cancer), or paraneoplastic syndromes. Even when not attributable to malignancy, dysphagia is usually a manifestation of an identifiable and treatable disease entity, making its evaluation beneficial to the patient and gratifying to the practitioner. The specific diagnostic algorithm to pursue is guided by the details of the history (Fig. 53-2). If oral or pharyngeal dysphagia is suspected, a fluoroscopic swallow study, usually done by a swallow therapist, is the procedure of choice. Otolaryngoscopic and neurologic evaluation also can be important, depending on the circumstances. For suspected esophageal dysphagia, upper endoscopy is the single most useful test. Endoscopy allows better visualization of mucosal lesions than does barium radiography and also allows one to obtain mucosal biopsies. Endoscopic or histologic abnormalities are evident in the leading causes of esophageal dysphagia: Schatzki ring, gastroesophageal reflux disease and eosinophilic esophagitis. Furthermore, therapeutic intervention with esophageal dilation can be done as part of the procedure if it is deemed necessary. The emergence of eosinophilic esophagitis as a leading cause of dysphagia in both children and adults has led to the recommendation that esophageal mucosal biopsies be obtained routinely in the evaluation of unexplained dysphagia even if endoscopically identified esophageal mucosal lesions are absent. For cases of suspected esophageal motility disorders, endoscopy is still the appropriate initial evaluation as neoplastic and inflammatory conditions can secondarily produce patterns of either achalasia or esophageal spasm. Esophageal manometry is done if dysphagia is not adequately explained by endoscopy or to confirm the diagnosis of a suspected esophageal motor disorder. Barium radiography can provide useful adjunctive information in cases of subtle or complex esophageal strictures, prior esophageal surgery, esophageal diverticula, or paraesophageal herniation. In specific cases, computed tomography (CT) examination and endoscopic ultrasonography may be useful.

TREATMENT

Treatment of dysphagia depends on both the locus and the specific etiology. Oropharyngeal dysphagia most commonly results from functional deficits caused by neurologic disorders. In such circumstances, the treatment focuses on utilizing postures or maneuvers devised to reduce pharyngeal residue and enhance airway protection learned under the direction of a trained swallow therapist. Aspiration risk may be reduced by altering the consistency of ingested food and liquid. Dysphagia resulting from a cerebrovascular accident usually, but not always, spontaneously improves within the first few weeks after the event. More severe and persistent cases may require gastrostomy and enteral feeding. Patients with myasthenia gravis (**Chap. 461**) and polymyositis (**Chap. 388**) may respond to medical treatment of the primary neuromuscular disease. Surgical intervention with cricopharyngeal myotomy is usually not helpful, with the exception of specific disorders such as the idiopathic cricopharyngeal bar, Zenker's diverticulum, and oculopharyngeal muscular dystrophy. Chronic neurologic disorders such as Parkinson's disease and amyotrophic lateral sclerosis may manifest with severe oropharyngeal dysphagia. Feeding by a nasogastric tube or an endoscopically placed gastrostomy tube may be considered for nutritional support; however, these maneuvers do not provide protection against aspiration of salivary secretions or refluxed gastric contents.

Treatment of esophageal dysphagia is covered in detail in **Chap. 347**. The majority of causes of esophageal dysphagia are effectively managed by means of esophageal dilatation using bougie or balloon dilators. Cancer and achalasia are often managed surgically, although endoscopic techniques are available for both palliation and primary therapy, respectively. Infectious etiologies respond to antimicrobial medications or treatment of the underlying immunosuppressive state. Finally, eosinophilic esophagitis has emerged as an important cause of dysphagia that is amenable to treatment by elimination of dietary allergens or administration of swallowed, topically acting glucocorticoids.

54**Nausea, Vomiting, and Indigestion**

William L. Hasler

Nausea is the subjective feeling of a need to vomit. *Vomiting* (emesis) is the oral expulsion of gastrointestinal contents due to contractions of gut and thoracoabdominal wall musculature. Vomiting is contrasted with *regurgitation*, the effortless passage of gastric contents into the mouth. *Rumination* is the repeated regurgitation of food residue, which may be rechewed and reswallowed. In contrast to emesis, these phenomena may exhibit volitional control. *Indigestion* is a term encompassing a range of complaints including nausea, vomiting, heartburn, regurgitation, and dyspepsia (the presence of symptoms thought to originate in the gastroduodenal region). Some individuals with dyspepsia report predominantly epigastric burning, gnawing, or pain. Others experience postprandial fullness, early satiety (an inability to complete a meal due to premature fullness), bloating, eructation (belching), and anorexia.

NAUSEA AND VOMITING**MECHANISMS**

Vomiting is coordinated by the brainstem and is effected by responses in the gut, pharynx, and somatic musculature. Mechanisms underlying nausea are poorly understood but likely involve the cerebral cortex, as nausea requires conscious perception. This is supported by functional

brain imaging studies showing activation of a range of cerebral cortical regions during nausea.

Coordination of Emesis Brainstem nuclei—including the nucleus tractus solitarius; dorsal vagal and phrenic nuclei; medullary nuclei regulating respiration; and nuclei that control pharyngeal, facial, and tongue movements—coordinate initiation of emesis. Neurokinin NK₁, serotonin 5-HT₃, and vasopressin pathways participate in this coordination.

Somatic and visceral muscles respond stereotypically during emesis. Inspiratory thoracic and abdominal wall muscles contract, producing high intrathoracic and intraabdominal pressures that evacuate the stomach. The gastric cardia herniates above the diaphragm, and the larynx moves upward to propel the vomitus. Distally migrating gut contractions are normally regulated by an electrical phenomenon, the slow wave, which cycles at 3 cycles/min in the stomach and 11 cycles/min in the duodenum. During emesis, the slow wave is abolished and is replaced by orally propagating spikes that evoke retrograde contractions that assist in expulsion of gut contents.

Activators of Emesis Emetic stimuli act at several sites. Emesis evoked by unpleasant thoughts or smells originates in the brain, whereas cranial nerves mediate vomiting after gag reflex activation. Motion sickness and inner ear disorders act on the labyrinthine system. Gastric irritants and cytotoxic agents like cisplatin stimulate gastroduodenal vagal afferent nerves. Nongastric afferents are activated by intestinal and colonic obstruction and mesenteric ischemia. The area postrema, in the medulla, responds to bloodborne stimuli (emetogenic drugs, bacterial toxins, uremia, hypoxia, ketoacidosis) and is termed the *chemoreceptor trigger zone*.

Neurotransmitters mediating vomiting are selective for different sites. Labyrinthine disorders stimulate vestibular muscarinic M₁ and histaminergic H₁ receptors. Vagal afferent stimuli activate serotonin 5-HT₃ receptors. The area postrema is served by nerves acting on 5-HT₃, M₁, H₁, and dopamine D₂ subtypes. Cannabinoid CB₁ pathways may participate in the cerebral cortex. Optimal pharmacologic therapy of vomiting requires understanding of these pathways.

DIFFERENTIAL DIAGNOSIS

Nausea and vomiting are caused by conditions within and outside the gut as well as by drugs and circulating toxins (**Table 54-1**).

Intraperitoneal Disorders Visceral obstruction and inflammation of hollow and solid viscera may elicit vomiting. Gastric obstruction results from ulcers and malignancy, whereas small-bowel and colon blockage occur because of adhesions, benign or malignant tumors, volvulus, intussusception, or inflammatory diseases like Crohn's disease. The superior mesenteric artery syndrome, occurring after weight loss or prolonged bed rest, results when the duodenum is compressed by the overlying superior mesenteric artery. Abdominal irradiation impairs intestinal motor function and induces strictures. Biliary colic causes nausea by acting on local afferent nerves. Vomiting with pancreatitis, cholecystitis, and appendicitis is due to visceral irritation and induction of ileus. Enteric infections with viruses like norovirus or rotavirus or bacteria such as *Staphylococcus aureus* and *Bacillus cereus* often cause vomiting, especially in children. Opportunistic infections like cytomegalovirus or herpes simplex virus induce emesis in immunocompromised individuals.

Gut sensorimotor dysfunction often causes nausea and vomiting. *Gastroparesis* presents with symptoms of gastric retention with evidence of delayed gastric emptying and occurs after vagotomy or with pancreatic carcinoma, mesenteric vascular insufficiency, or organic diseases like diabetes, scleroderma, and amyloidosis. Idiopathic gastroparesis is the most common etiology. It occurs in the absence of systemic illness and may follow a viral illness, suggesting an infectious trigger. *Intestinal pseudoobstruction* is characterized by disrupted intestinal and colonic motor activity with retention of food residue and secretions; bacterial overgrowth; nutrient malabsorption; and symptoms of nausea, vomiting, bloating, pain, and altered defecation. Intestinal pseudoobstruction may be idiopathic, inherited as a familial visceral myopathy or neuropathy, result from systemic disease, or

TABLE 54-1 CAUSES OF NAUSEA AND VOMITING

Intraoperative	Extraperitoneal	Medications/ Metabolic Disorders
Obstructing disorders	Cardiopulmonary disease	Drugs
Pyloric obstruction	Cardiomyopathy	Cancer chemotherapy
Small-bowel obstruction	Myocardial infarction	Antibiotics
Colonic obstruction	Labyrinthine disease	Cardiac antiarrhythmics
Superior mesenteric artery syndrome	Motion sickness	Digoxin
Enteric infections	Labyrinthitis	Oral hypoglycemics
Viral	Malignancy	Oral contraceptives
Bacterial	Intracerebral disorders	Endocrine/metabolic disease
Inflammatory diseases	Malignancy	Pregnancy
Cholecystitis	Hemorrhage	Uremia
Pancreatitis	Abscess	Ketoacidosis
Appendicitis	Hydrocephalus	Thyroid and parathyroid disease
Hepatitis	Psychiatric illness	Adrenal insufficiency
Altered sensorimotor function	Anorexia and bulimia nervosa	Toxins
Gastroparesis	Depression	Liver failure
Intestinal pseudoobstruction	Postoperative vomiting	Ethanol
Gastroesophageal reflux		
Chronic idiopathic nausea		
Functional vomiting		
Cyclic vomiting syndrome		
Cannabinoid hyperemesis syndrome		
Rumination syndrome		
Biliary colic		
Abdominal irradiation		

occur as a paraneoplastic consequence of malignancy (e.g., small-cell lung carcinoma). Patients with gastroesophageal reflux may report nausea and vomiting, as do some with irritable bowel syndrome (IBS) or chronic constipation.

Other functional gastroduodenal disorders without organic abnormalities have been characterized in adults. *Chronic idiopathic nausea* is defined as nausea without vomiting occurring several times a week. *Functional vomiting* is defined as one or more vomiting episodes weekly in the absence of an eating disorder or psychiatric disease. *Cyclic vomiting syndrome* presents with periodic discrete episodes of relentless nausea and vomiting in children and adults and shows an association with migraine headaches, suggesting that some cases may be migraine variants. Some adult cases have been described in association with rapid gastric emptying. A related condition, *cannabinoid hyperemesis syndrome*, presents with cyclical vomiting with intervening well periods in individuals (mostly men) who use large quantities of cannabis over many years and resolves with its discontinuation. Pathologic behaviors such as taking prolonged hot baths or showers are associated with the syndrome. *Rumination syndrome*, characterized by repetitive regurgitation of recently ingested food, is often misdiagnosed as refractory vomiting.

Extraperitoneal Disorders Myocardial infarction and congestive heart failure may cause nausea and vomiting. Postoperative emesis occurs after 25% of surgeries, most commonly laparotomy and orthopedic surgery. Increased intracranial pressure from tumors, bleeding, abscess, or blockage of cerebrospinal fluid outflow produces vomiting with or without nausea. Patients with psychiatric illnesses including anorexia nervosa, bulimia nervosa, anxiety, and depression often report significant nausea that may be associated with delayed gastric emptying.

Medications and Metabolic Disorders Drugs evoke vomiting by action on the stomach (analgesics, erythromycin) or area postrema (opiates, anti-parkinsonian drugs). Other emetogenic agents include antibiotics,

cardiac antiarrhythmics, antihypertensives, oral hypoglycemics, antidepressants (selective serotonin and serotonin norepinephrine reuptake inhibitors), smoking cessation drugs (varenicline, nicotine), and contraceptives. Cancer chemotherapy causes vomiting that is acute (within hours of administration), delayed (after 1 or more days), or anticipatory. Acute emesis from highly emetogenic agents (e.g., cisplatin) is mediated by 5-HT₃ pathways, whereas delayed emesis is less dependent on 5-HT₃ mechanisms. Anticipatory nausea may respond to anxiolytic therapy rather than antiemetics.

Metabolic disorders elicit nausea and vomiting. Pregnancy is the most prevalent endocrinologic cause, and nausea affects 70% of women in the first trimester. Hyperemesis gravidarum is a severe form of nausea of pregnancy that produces significant fluid loss and electrolyte disturbances. Uremia, ketoacidosis, adrenal insufficiency, and parathyroid and thyroid disease are other metabolic etiologies.

Circulating toxins evoke emesis via effects on the area postrema. Endogenous toxins are generated in fulminant liver failure, whereas exogenous enterotoxins may be produced by enteric bacterial infection. Ethanol intoxication is a common toxic etiology of nausea and vomiting.

APPROACH TO THE PATIENT: Nausea and Vomiting

HISTORY AND PHYSICAL EXAMINATION

The history helps define the etiology of nausea and vomiting. Drugs, toxins, and infections often cause acute symptoms, whereas established illnesses evoke chronic complaints. Gastroparesis and pyloric obstruction elicit vomiting within an hour of eating. Emesis from intestinal blockage occurs later. Vomiting occurring within minutes of meal consumption prompts consideration of rumination syndrome. With severe gastric emptying delays, the vomitus may contain food residue ingested hours or days before. Hematemesis raises suspicion of an ulcer, malignancy, or Mallory-Weiss tear. Feculent emesis is noted with distal intestinal or colonic obstruction. Bileous vomiting excludes gastric obstruction, whereas emesis of undigested food is consistent with a Zenker's diverticulum or achalasia. Vomiting can relieve abdominal pain from a bowel obstruction, but has no effect in pancreatitis or cholecystitis. Profound weight loss raises concern about malignancy or obstruction. Fevers suggest inflammation. An intracranial source is considered if there are headaches or visual field changes. Vertigo or tinnitus indicates labyrinthine disease.

The physical examination complements the history. Orthostatic hypotension and reduced skin turgor indicate intravascular fluid loss. Pulmonary abnormalities raise concern for aspiration of vomitus. Abdominal auscultation may reveal absent bowel sounds with ileus. High-pitched rushes suggest bowel obstruction, whereas a succussion splash upon abrupt lateral movement of the patient is found with gastroparesis or pyloric obstruction. Tenderness or involuntary guarding raises suspicion of inflammation, whereas fecal blood suggests mucosal injury from ulcer, ischemia, or tumor. Neurologic disease presents with papilledema, visual field loss, or focal neural abnormalities. Neoplasm is suggested by palpation of masses or adenopathy.

DIAGNOSTIC TESTING

For intractable symptoms or an elusive diagnosis, selected screening tests can direct clinical care. Electrolyte replacement is indicated for hypokalemia or metabolic alkalosis. Iron-deficiency anemia mandates a search for mucosal injury. Pancreaticobiliary disease is indicated by abnormal pancreatic or liver biochemistries, whereas endocrinologic, rheumatologic, or paraneoplastic etiologies are suggested by hormone or serologic abnormalities. If bowel obstruction is suspected, supine and upright abdominal radiographs may show intestinal air-fluid levels with reduced colonic air. Ileus is characterized by diffusely dilated air-filled bowel loops.

Anatomic studies may be indicated if initial testing is nondiagnostic. Upper endoscopy detects ulcers, malignancy, and retained gastric food residue in gastroparesis. Small-bowel barium radiography or computed tomography (CT) diagnoses partial bowel obstruction. Colonoscopy or contrast enema radiography detects colonic obstruction. Ultrasound or CT defines intraperitoneal inflammation; CT and magnetic resonance imaging (MRI) enterography provide superior definition of inflammation in Crohn's disease. CT or MRI of the head can delineate intracranial disease. Mesenteric angiography, CT, or MRI is useful for suspected ischemia.

Gastrointestinal motility testing may detect an underlying motor disorder when anatomic abnormalities are absent. Gastroparesis commonly is diagnosed by gastric scintigraphy, by which emptying of a radiolabeled meal is measured. Isotopic breath tests and wireless motility capsule methods are alternative tests to define gastroparesis in different regions of the world. Intestinal pseudoobstruction often is suggested by abnormal barium transit and luminal dilation on small-bowel contrast radiography. Delayed small-bowel transit also may be detected by wireless capsule techniques. Small-intestinal manometry can confirm the diagnosis and further characterize the motor abnormality as neuropathic or myopathic based on contractile patterns. Such investigation can obviate the need for surgical intestinal biopsy to evaluate for smooth muscle or neuronal degeneration. Combined ambulatory esophageal pH/impedance testing and high-resolution manometry can facilitate diagnosis of rumination syndrome.

TREATMENT NAUSEA AND VOMITING

GENERAL PRINCIPLES

Therapy of vomiting is tailored to correcting remediable abnormalities if possible. Hospitalization is considered for severe dehydration, especially if oral fluid replenishment cannot be sustained. Once oral intake is tolerated, nutrients are restarted with low-fat liquids, because lipids delay gastric emptying. Foods high in indigestible

residue are avoided because these prolong gastric retention. Controlling blood glucose in poorly controlled diabetics can reduce hospitalizations in gastroparesis.

ANTIEMETIC MEDICATIONS

The most commonly used antiemetic agents act on central nervous system sites (Table 54-2). Antihistamines like dimenhydrinate and meclizine and anticholinergics like scopolamine act on labyrinthine pathways to treat motion sickness and inner ear disorders. Dopamine D₂ antagonists treat emesis evoked by area postrema stimuli and are used for medication, toxic, and metabolic etiologies. Dopamine antagonists cross the blood-brain barrier and cause anxiety, movement disorders, and hyperprolactinemic effects (galactorrhea, sexual dysfunction).

Other classes exhibit antiemetic properties. 5-HT₃ antagonists such as ondansetron and granisetron can prevent postoperative vomiting, radiation therapy–induced symptoms, and cancer chemotherapy–induced emesis, but also are used for other causes of emesis with limited evidence for efficacy. Tricyclic antidepressant agents provide symptomatic benefit in patients with chronic idiopathic nausea and functional vomiting as well as in long-standing diabetic patients with nausea and vomiting. Other antidepressants such as mirtazapine and olanzapine also may exhibit antiemetic effects.

GASTROINTESTINAL MOTOR STIMULANTS

Drugs that stimulate gastric emptying are used for gastroparesis (Table 54-2). Metoclopramide, a combined 5-HT₄ agonist and D₂ antagonist, is effective in gastroparesis, but antidopaminergic side effects, such as dystonias and mood and sleep disturbances, limit use in ~25% of cases. Erythromycin increases gastroduodenal motility by action on receptors for motilin, an endogenous stimulant of fasting motor activity. Intravenous erythromycin is useful for inpatients with refractory gastroparesis, but oral forms have some utility. Domperidone, a D₂ antagonist not available in the United States, exhibits prokinetic and antiemetic effects but does not cross into most brain regions; thus, anxiety and dystonic reactions are rare. The main side effects of domperidone relate to induction of

TABLE 54-2 TREATMENT OF NAUSEA AND VOMITING

Treatment	Mechanism	Examples	Clinical Indications
Antiemetic agents	Antihistaminergic	Dimenhydrinate, meclizine	Motion sickness, inner ear disease
	Anticholinergic	Scopolamine	Motion sickness, inner ear disease
	Antidopaminergic	Prochlorperazine, thiethylperazine	Medication-, toxin-, or metabolic-induced emesis
	5-HT ₃ antagonist	Ondansetron, granisetron	Chemotherapy- and radiation-induced emesis, postoperative emesis
	NK ₁ antagonist	Aprepitant	Chemotherapy-induced nausea and vomiting
	Tricyclic antidepressant	Amitriptyline, nortriptyline	Functional vomiting, chronic idiopathic nausea, cyclic vomiting syndrome, ?gastroparesis
Prokinetic agents	Other antidepressant	Mirtazapine, olanzapine	?Functional vomiting, ?chronic idiopathic nausea, ?gastroparesis
	5-HT ₄ agonist and antidopaminergic	Metoclopramide	Gastroparesis
	Motilin agonist	Erythromycin	Gastroparesis, ?intestinal pseudoobstruction
	Peripheral antidopaminergic	Domperidone	Gastroparesis
Special settings	Somatostatin analogue	Octreotide	Intestinal pseudoobstruction
	Acetylcholinesterase inhibitor	Pyridostigmine	?Small-intestinal dysmotility/pseudoobstruction
	Benzodiazepines	Lorazepam	Anticipatory nausea and vomiting with chemotherapy
	Glucocorticoids	Methylprednisolone, dexamethasone	Chemotherapy-induced emesis
	Cannabinoids	Tetrahydrocannabinol	Chemotherapy-induced emesis

Note: ?, indication is uncertain.

hyperprolactinemia via effects on pituitary regions served by a porous blood-brain barrier.

Refractory motility disorders pose significant challenges. Intestinal pseudoobstruction may respond to the somatostatin analogue octreotide, which induces propagative small-intestinal motor complexes. Acetylcholinesterase inhibitors such as pyridostigmine are also observed to benefit some patients with small-bowel dysmotility. Pyloric injections of botulinum toxin are reported in uncontrolled studies to reduce gastroparesis symptoms, but small controlled trials observe benefits no greater than sham treatments. Surgical pyloroplasty has improved symptoms in case series. Placing a feeding jejunostomy reduces hospitalizations and improves overall health in some patients with drug-refractory gastroparesis. Postvagotomy gastroparesis may improve with near-total gastric resection; similar operations are now being tried for other gastroparesis etiologies. Implanted gastric electrical stimulators may reduce symptoms, enhance nutrition, improve quality of life, and decrease health care expenditures in medication-refractory gastroparesis, but small controlled trials do not report convincing benefits.

SAFETY CONSIDERATIONS

Safety concerns about selected antiemetics have been emphasized. Centrally acting antidopaminergics, especially metoclopramide, can cause irreversible movement disorders such as tardive dyskinesia, particularly in older patients. This complication should be carefully explained and documented in the medical record. Some agents with antiemetic properties including domperidone, erythromycin, tricyclics, and 5-HT₃ antagonists can induce dangerous cardiac rhythm disturbances, especially in those with QTc interval prolongation on electrocardiography (ECG). Surveillance ECG testing has been advocated for some of these agents.

SELECTED CLINICAL SETTINGS

Some cancer chemotherapies are intensely emetogenic (Chap. 103e). Combining a 5-HT₃ antagonist, an NK₁ antagonist, and a glucocorticoid provides significant control of both acute and delayed vomiting after highly emetogenic chemotherapy. Unlike other drugs in the same class, the 5-HT₃ antagonist palonosetron exhibits efficacy at preventing delayed chemotherapy-induced vomiting. Benzodiazepines such as lorazepam can reduce anticipatory nausea and vomiting. Miscellaneous therapies with benefit in chemotherapy-induced emesis include cannabinoids, olanzapine, and alternative therapies like ginger. Most antiemetic regimens produce greater reductions in vomiting than in nausea.

Clinicians should exercise caution in managing pregnant patients with nausea. Studies of the teratogenic effects of antiemetic agents provide conflicting results. Few controlled trials have been performed in nausea of pregnancy. Antihistamines such as meclizine and doxylamine, antidopaminergics such as prochlorperazine, and antiserotonergics such as ondansetron demonstrate limited efficacy. Some obstetricians offer alternative therapies such as pyridoxine, acupressure, or ginger.

Managing cyclic vomiting syndrome is a challenge. Prophylaxis with tricyclic antidepressants, cyproheptadine, or β -adrenoceptor antagonists can reduce the severity and frequency of attacks. Intravenous 5-HT₃ antagonists combined with the sedating effects of a benzodiazepine like lorazepam are a mainstay of treatment of acute flares. Small studies report benefits with antimigraine agents, including the 5-HT₁ agonist sumatriptan, as well as selected anticonvulsants such as zonisamide and levetiracetam.

INDIGESTION

MECHANISMS

The most common causes of indigestion are gastroesophageal reflux and functional dyspepsia. Other cases are a consequence of organic illness.

Gastroesophageal Reflux Gastroesophageal reflux results from many physiologic defects. Reduced lower esophageal sphincter (LES) tone contributes to reflux in scleroderma and pregnancy and may be a factor in some patients without systemic illness. Others exhibit frequent transient LES relaxations (TLESRs) that cause bathing of the esophagus by acid or nonacidic fluid. Overeating and aerophagia override the barrier function of the LES, whereas reductions in esophageal body motility or salivary secretion prolong fluid exposure. Increased intragastric pressure promotes gastroesophageal reflux in obesity. The role of hiatal hernias is controversial—most reflux patients have hiatal hernias, but most with hiatal hernias do not have excess heartburn.

Gastric Motor Dysfunction Disturbed gastric motility may contribute to gastroesophageal reflux in up to one-third of cases. Delayed gastric emptying is also found in ~30% of functional dyspeptics. Conversely, some dyspeptics exhibit rapid gastric emptying. The relation of these defects to symptom induction is uncertain; studies show poor correlation between symptom severity and degrees of motor dysfunction. Impaired gastric fundus relaxation after eating (i.e., accommodation) may underlie selected dyspeptic symptoms like bloating, nausea, and early satiety in ~40% of patients.

Visceral Afferent Hypersensitivity Disturbed gastric sensation is another pathogenic factor in functional dyspepsia. Visceral hypersensitivity was first reported in IBS with demonstration of heightened perception of rectal balloon inflation without changes in compliance. Similarly, ~35% of dyspeptic patients note discomfort with fundic distention to lower pressures than healthy controls. Others with dyspepsia exhibit hypersensitivity to chemical stimulation with capsaicin or with acid or lipid exposure in the duodenum. Some individuals with functional heartburn without increased acid or nonacid reflux are believed to have heightened perception of normal esophageal pH and volume.

Other Factors *Helicobacter pylori* has a clear etiologic role in peptic ulcer disease, but ulcers cause a minority of dyspepsia cases. *H. pylori* is a minor factor in the genesis of functional dyspepsia. Functional dyspepsia is associated with chronic fatigue, produces reduced physical and mental well-being, and is exacerbated by stress. Anxiety, depression, and somatization may have contributing roles in some cases. Functional MRI studies show increased activation of several brain regions, emphasizing contributions from central nervous system factors. Analgesics cause dyspepsia, whereas nitrates, calcium channel blockers, theophylline, and progesterone promote gastroesophageal reflux. Other stimuli that induce reflux include ethanol, tobacco, and caffeine via LES relaxation. Genetic factors may promote development of reflux and dyspepsia.

DIFFERENTIAL DIAGNOSIS

Gastroesophageal Reflux Disease Gastroesophageal reflux disease (GERD) is prevalent. Heartburn is reported once monthly by 40% of Americans and daily by 7–10%. Most cases of heartburn occur because of excess acid reflux, but reflux of nonacidic fluid produces similar symptoms. Alkaline reflux esophagitis produces GERD-like symptoms most often in patients who have had surgery for peptic ulcer disease. Ten percent of patients with heartburn exhibit normal esophageal acid exposure and no increase in nonacidic reflux (functional heartburn).

Functional Dyspepsia Nearly 25% of the populace has dyspepsia at least six times yearly, but only 10–20% present to clinicians. Functional dyspepsia, the cause of symptoms in >60% of dyspeptic patients, is defined as ≥ 3 months of bothersome postprandial fullness, early satiety, or epigastric pain or burning with symptom onset at least 6 months before diagnosis in the absence of organic cause. Functional dyspepsia is subdivided into postprandial distress syndrome, characterized by meal-induced fullness, early satiety, and discomfort, and epigastric pain syndrome, which presents with epigastric burning pain unrelated to meals. Most cases follow a benign course, but some with *H. pylori* infection or on nonsteroidal anti-inflammatory drugs

(NSAIDs) develop ulcers. As with idiopathic gastroparesis, some cases of functional dyspepsia result from prior infection.

Ulcer Disease In most GERD patients, there is no destruction of the esophagus. However, 5% develop esophageal ulcers, and some form strictures. Symptoms cannot distinguish nonerosive from erosive or ulcerative esophagitis. A minority of cases of dyspepsia stem from gastric or duodenal ulcers. The most common causes of ulcer disease are *H. pylori* infection and use of NSAIDs. Other rare causes of gastroduodenal ulcers include Crohn's disease (Chap. 351) and Zollinger-Ellison syndrome (Chap. 348), resulting from gastrin overproduction by an endocrine tumor.

Malignancy Dyspeptic patients often seek care because of fear of cancer, but few cases result from malignancy. Esophageal squamous cell carcinoma occurs most often with long-standing tobacco or ethanol intake. Other risks include prior caustic ingestion, achalasia, and the hereditary disorder tylosis. Esophageal adenocarcinoma usually complicates prolonged acid reflux. Eight to 20% of GERD patients exhibit esophageal intestinal metaplasia, termed *Barrett's metaplasia*, a condition that predisposes to esophageal adenocarcinoma (Chap. 109). Gastric malignancies include adenocarcinoma, which is prevalent in certain Asian societies, and lymphoma.

Other Causes Opportunistic fungal or viral esophageal infections may produce heartburn but more often cause odynophagia. Other causes of esophageal inflammation include eosinophilic esophagitis and pill esophagitis. Biliary colic is in the differential diagnosis of unexplained upper abdominal pain, but most patients with biliary colic report discrete acute episodes of right upper quadrant or epigastric pain rather than the chronic burning, discomfort, and fullness of dyspepsia. Twenty percent of patients with gastroparesis report a predominance of pain or discomfort rather than nausea and vomiting. Intestinal lactase deficiency as a cause of gas, bloating, and discomfort occurs in 15–25% of whites of northern European descent but is more common in blacks and Asians. Intolerance of other carbohydrates (e.g., fructose, sorbitol) produces similar symptoms. Small-intestinal bacterial overgrowth may cause dyspepsia, often associated with bowel dysfunction, distention, and malabsorption. Eosinophilic infiltration of the duodenal mucosa is described in some dyspeptics, particularly with postprandial distress syndrome. Celiac disease, pancreatic disease (chronic pancreatitis, malignancy), hepatocellular carcinoma, Ménétrier's disease, infiltrative diseases (sarcoidosis, eosinophilic gastroenteritis), mesenteric ischemia, thyroid and parathyroid disease, and abdominal wall strain cause dyspepsia. Gluten sensitivity in the absence of celiac disease is reported to evoke unexplained upper abdominal symptoms. Extraperitoneal etiologies of indigestion include congestive heart failure and tuberculosis.

APPROACH TO THE PATIENT: Indigestion

HISTORY AND PHYSICAL EXAMINATION

Care of the indigestion patient requires a thorough interview. GERD classically produces heartburn, a substernal warmth that moves toward the neck. Heartburn often is exacerbated by meals and may awaken the patient. Associated symptoms include regurgitation of acid or nonacidic fluid and water brash, the reflex release of salty salivary secretions into the mouth. Atypical symptoms include pharyngitis, asthma, cough, bronchitis, hoarseness, and chest pain that mimics angina. Some patients with acid reflux on esophageal pH testing do not report heartburn, but note abdominal pain or other symptoms.

Dyspeptic patients typically report symptoms referable to the upper abdomen that may be meal-related, as with postprandial distress syndrome, or independent of food ingestion, as in epigastric pain syndrome. Functional dyspepsia overlaps with other disorders including GERD, IBS, and idiopathic gastroparesis.

TABLE 54-3 ALARM SYMPTOMS IN GASTROESOPHAGEAL REFLUX DISEASE

Odynophagia
Unexplained weight loss
Recurrent vomiting
Occult or gross gastrointestinal bleeding
Jaundice
Palpable mass or adenopathy
Family history of gastrointestinal malignancy

The physical exam with GERD and functional dyspepsia usually is normal. In atypical GERD, pharyngeal erythema and wheezing may be noted. Recurrent acid regurgitation may cause poor dentition. Dyspeptics may exhibit epigastric tenderness or distention.

Discriminating functional and organic causes of indigestion mandates excluding certain historic and exam features. Odynophagia suggests esophageal infection. Dysphagia is concerning for a benign or malignant esophageal blockage. Other alarm features include unexplained weight loss, recurrent vomiting, occult or gross bleeding, jaundice, palpable mass or adenopathy, and a family history of gastrointestinal neoplasm.

DIAGNOSTIC TESTING

Because indigestion is prevalent and most cases result from GERD or functional dyspepsia, a general principle is to perform only limited and directed diagnostic testing of selected individuals.

Once alarm factors are excluded (Table 54-3), patients with typical GERD do not need further evaluation and are treated empirically. Upper endoscopy is indicated to exclude mucosal injury in cases with atypical symptoms, symptoms unresponsive to acid suppression, or alarm factors. For heartburn >5 years in duration, especially in patients >50 years old, endoscopy is advocated to screen for Barrett's metaplasia. The benefits and cost-effectiveness of this approach have not been validated in controlled studies. Ambulatory esophageal pH testing using a catheter method or a wireless capsule endoscopically attached to the esophageal wall is considered for drug-refractory symptoms and atypical symptoms like unexplained chest pain. High-resolution esophageal manometry is ordered when surgical treatment of GERD is considered. A low LES pressure predicts failure of drug therapy and provides a rationale to proceed to surgery. Poor esophageal body peristalsis raises concern about postoperative dysphagia and directs the choice of surgical technique. Nonacidic reflux may be detected by combined esophageal impedance-pH testing in medication-unresponsive patients.

Upper endoscopy is recommended as the initial test in patients with unexplained dyspepsia who are >55 years old or who have alarm factors because of the purported elevated risks of malignancy and ulcer in these groups. However, endoscopic findings in unexplained dyspepsia include erosive esophagitis in 13%, peptic ulcer in 8%, and gastric or esophageal malignancy in only 0.3%. Management of patients <55 years old without alarm factors depends on the local prevalence of *H. pylori* infection. In regions with low *H. pylori* prevalence (<10%), a 4-week trial of an acid-suppressing medication such as a proton pump inhibitor (PPI) is recommended. If this fails, a "test and treat" approach is most commonly applied. *H. pylori* status is determined with urea breath testing, stool antigen measurement, or blood serology testing. Those who are *H. pylori* positive are given therapy to eradicate the infection. If symptoms resolve on either regimen, no further intervention is required. For patients in areas with high *H. pylori* prevalence (>10%), an initial test and treat approach is advocated, with a subsequent trial of an acid-suppressing regimen offered for those in whom *H. pylori* treatment fails or for those who are negative for the infection. In each of these patient subsets, upper endoscopy is reserved for those whose symptoms fail to respond to therapy.

Further testing is indicated in some settings. If bleeding is noted, a blood count can exclude anemia. Thyroid chemistries or calcium levels screen for metabolic disease, whereas specific serologies may

suggest celiac disease. Pancreatic and liver chemistries are obtained for possible pancreaticobiliary causes. Ultrasound, CT, or MRI is performed if abnormalities are found. Gastric emptying testing is considered to exclude gastroparesis for dyspeptic symptoms that resemble postprandial distress when drug therapy fails and in some GERD patients, especially if surgical intervention is an option. Breath testing after carbohydrate ingestion detects lactase deficiency, intolerance to other carbohydrates, or small-intestinal bacterial overgrowth.

TREATMENT INDIGESTION

GENERAL PRINCIPLES

For mild indigestion, reassurance that a careful evaluation revealed no serious organic disease may be the only intervention needed. Drugs that cause gastroesophageal reflux or dyspepsia should be stopped, if possible. Patients with GERD should limit ethanol, caffeine, chocolate, and tobacco use due to their effects on the LES. Other measures in GERD include ingesting a low-fat diet, avoiding snacks before bedtime, and elevating the head of the bed. Patients with functional dyspepsia also may be advised to reduce intake of fat, spicy foods, caffeine, and alcohol.

Specific therapies for organic disease should be offered when possible. Surgery is appropriate for biliary colic, whereas diet changes are indicated for lactase deficiency or celiac disease. Peptic ulcers may be cured by specific medical regimens. However, because most indigestion is caused by GERD or functional dyspepsia, medications that reduce gastric acid, modulate motility, or blunt gastric sensitivity are used.

ACID-SUPPRESSING OR -NEUTRALIZING MEDICATIONS

Drugs that reduce or neutralize gastric acid are often prescribed for GERD. Histamine H_2 antagonists like cimetidine, ranitidine, famotidine, and nizatidine are useful in mild to moderate GERD. For severe symptoms or for many cases of erosive or ulcerative esophagitis, PPIs such as omeprazole, lansoprazole, rabeprazole, pantoprazole, esomeprazole, or dexlansoprazole are needed. These drugs inhibit gastric H^+ , K^+ -ATPase and are more potent than H_2 antagonists. Up to one-third of GERD patients do not respond to standard PPI doses; one-third of these patients have nonacidic reflux, whereas 10% have persistent acid-related disease. Furthermore, heartburn typically responds better to PPI therapy than regurgitation or atypical GERD symptoms. Some individuals respond to doubling of the PPI dose or adding an H_2 antagonist at bedtime. Infrequent complications of long-term PPI therapy include infection, diarrhea (from *Clostridium difficile* infection or microscopic colitis), small-intestinal bacterial overgrowth, nutrient deficiency (vitamin B_{12} , iron, calcium), hypomagnesemia, bone demineralization, interstitial nephritis, and impaired medication absorption (e.g., clopidogrel). Many patients started on a PPI can be stepped down to an H_2 antagonist or be switched to an on-demand schedule.

Acid-suppressing drugs are also effective in selected patients with functional dyspepsia. A meta-analysis of eight controlled trials calculated a risk ratio of 0.86, with a 95% confidence interval of 0.78–0.95, favoring PPI therapy over placebo. H_2 antagonists also reportedly improve symptoms in functional dyspepsia; however, findings of trials of this drug class likely are influenced by inclusion of large numbers of GERD patients.

Antacids are useful for short-term control of mild GERD but have less benefit in severe cases unless given at high doses that cause side effects (diarrhea and constipation with magnesium- and aluminum-containing agents, respectively). Alginic acid combined with antacids forms a floating barrier to reflux in patients with upright symptoms. Sucralfate, a salt of aluminum hydroxide and sucrose octasulfate that buffers acid and binds pepsin and bile salts, shows efficacy in GERD similar to H_2 antagonists.

HELICOBACTER PYLORI ERADICATION

H. pylori eradication is definitively indicated only for peptic ulcer and mucosa-associated lymphoid tissue gastric lymphoma. The utility of eradication therapy in functional dyspepsia is limited, although some cases (particularly with the epigastric pain syndrome subtype) relate to this infection. A meta-analysis of 18 controlled trials calculated a relative risk reduction of 10%, with a 95% confidence interval of 6–14%, favoring *H. pylori* eradication over placebo. Most drug combinations (Chaps. 188 and 348) include 10–14 days of a PPI or bismuth subsalicylate in concert with two antibiotics. *H. pylori* infection is associated with reduced prevalence of GERD, especially in the elderly. However, eradication of the infection does not worsen GERD symptoms. No consensus recommendations regarding *H. pylori* eradication in GERD patients have been offered.

AGENTS THAT MODIFY GASTROINTESTINAL MOTOR ACTIVITY

Prokinetics like metoclopramide, erythromycin, and domperidone have limited utility in GERD. The γ -aminobutyric acid B (GABA-B) agonist baclofen reduces esophageal exposure to acid and non-acidic fluids by reducing TLESRs by 40%; this drug is proposed for refractory acid and nonacid reflux. Several studies have promoted the efficacy of motor-stimulating drugs in functional dyspepsia, but publication bias and small sample sizes raise questions about reported benefits of these agents. Some clinicians suggest that patients with the postprandial distress subtype may respond preferentially to prokinetic drugs. The 5-HT₁ agonist buspirone may improve some functional dyspepsia symptoms by enhancing meal-induced gastric accommodation. Acotiamide promotes gastric emptying and augments accommodation by enhancing gastric acetylcholine release via muscarinic receptor antagonism and acetylcholinesterase inhibition. This agent is approved for functional dyspepsia in Japan and is in testing elsewhere.

OTHER OPTIONS

Antireflux surgery (fundoplication) to increase LES pressure may be offered to GERD patients who are young and require lifelong therapy, have typical heartburn and regurgitation, are responsive to PPIs, and show evidence of acid reflux on pH monitoring. Surgery also is effective for some cases of nonacidic reflux. Individuals who respond less well to fundoplication include those with atypical symptoms or who have esophageal body motor disturbances. Dysphagia, gas-bloat syndrome, and gastroparesis are long-term complications of these procedures; ~60% develop recurrent GERD symptoms over time. The utility and safety of endoscopic therapies (radiofrequency ablation, transoral incisionless fundoplication) to enhance gastroesophageal barrier function have unproved durable benefits for refractory GERD.

Some patients with functional heartburn and functional dyspepsia refractory to standard therapies may respond to antidepressants in tricyclic and selective serotonin reuptake inhibitor classes, although studies are limited. Their mechanism of action may involve blunting of visceral pain processing in the brain. Gas and bloating are among the most troubling symptoms in some patients with indigestion and can be difficult to treat. Dietary exclusion of gas-producing foods such as legumes and use of simethicone or activated charcoal provide benefits in some cases. Low FODMAP (fermentable oligosaccharide, disaccharide, monosaccharide, and polyol) diets and therapies to modify gut flora (nonabsorbable antibiotics, probiotics) reduce gaseous symptoms in some IBS patients. The utility of low-FODMAP diets, antibiotics, and probiotics in functional dyspepsia is unproven. Herbal remedies such as STW 5 (Iberogast, a mixture of nine herbal agents) are useful in some dyspeptic patients. Psychological treatments (e.g., behavioral therapy, psychotherapy, hypnotherapy) may be offered for refractory functional dyspepsia, but no convincing data confirm their efficacy.

Diarrhea and constipation are exceedingly common and, together, exact an enormous toll in terms of mortality, morbidity, social inconvenience, loss of work productivity, and consumption of medical resources. Worldwide, >1 billion individuals suffer one or more episodes of acute diarrhea each year. Among the 100 million persons affected annually by acute diarrhea in the United States, nearly half must restrict activities, 10% consult physicians, ~250,000 require hospitalization, and ~5000 die (primarily the elderly). The annual economic burden to society may exceed \$20 billion. Acute infectious diarrhea remains one of the most common causes of mortality in developing countries, particularly among impoverished infants, accounting for 1.8 million deaths per year. Recurrent, acute diarrhea in children in tropical countries results in environmental enteropathy with long-term impacts on physical and intellectual development.

Constipation, by contrast, is rarely associated with mortality and is exceedingly common in developed countries, leading to frequent self-medication and, in a third of those, to medical consultation. Population statistics on chronic diarrhea and constipation are more uncertain, perhaps due to variable definitions and reporting, but the frequency of these conditions is also high. United States population surveys put prevalence rates for chronic diarrhea at 2–7% and for chronic constipation at 12–19%, with women being affected twice as often as men. Diarrhea and constipation are among the most common patient complaints presenting to internists and primary care physicians, and they account for nearly 50% of referrals to gastroenterologists.

Although diarrhea and constipation may present as mere nuisance symptoms at one extreme, they can be severe or life-threatening at the other. Even mild symptoms may signal a serious underlying gastrointestinal lesion, such as colorectal cancer, or systemic disorder, such as thyroid disease. Given the heterogeneous causes and potential severity of these common complaints, it is imperative for clinicians to appreciate the pathophysiology, etiologic classification, diagnostic strategies, and principles of management of diarrhea and constipation, so that rational and cost-effective care can be delivered.

NORMAL PHYSIOLOGY

While the primary function of the small intestine is the digestion and assimilation of nutrients from food, the small intestine and colon together perform important functions that regulate the secretion and absorption of water and electrolytes, the storage and subsequent transport of intraluminal contents aborally, and the salvage of some nutrients that are not absorbed in the small intestine after bacterial metabolism of carbohydrate allows salvage of short-chain fatty acids. The main motor functions are summarized in [Table 55-1](#). Alterations in fluid and electrolyte handling contribute significantly to diarrhea. Alterations in motor and sensory functions of the colon result in

TABLE 55-1 NORMAL GASTROINTESTINAL MOTILITY: FUNCTIONS AT DIFFERENT ANATOMIC LEVELS

Stomach and Small Bowel

- Synchronized MMC in fasting
- Accommodation, trituration, mixing, transit
 - Stomach ~3 h
 - Small bowel ~3 h
- Ileal reservoir empties boluses

Colon: Irregular Mixing, Fermentation, Absorption, Transit

- Ascending, transverse: reservoirs
- Descending: conduit
- Sigmoid/rectum: volitional reservoir

Abbreviation: MMC, migrating motor complex.

highly prevalent syndromes such as irritable bowel syndrome (IBS), chronic diarrhea, and chronic constipation.

NEURAL CONTROL

The small intestine and colon have intrinsic and extrinsic innervation. The *intrinsic innervation*, also called the enteric nervous system, comprises myenteric, submucosal, and mucosal neuronal layers. The function of these layers is modulated by interneurons through the actions of neurotransmitter amines or peptides, including acetylcholine, vasoactive intestinal peptide (VIP), opioids, norepinephrine, serotonin, adenosine triphosphate (ATP), and nitric oxide (NO). The myenteric plexus regulates smooth-muscle function through intermediary pacemaker-like cells called the interstitial cells of Cajal, and the submucosal plexus affects secretion, absorption, and mucosal blood flow. The enteric nervous system receives input from the extrinsic nerves, but it is capable of independent control of these functions.

The *extrinsic innervations* of the small intestine and colon are part of the autonomic nervous system and also modulate motor and secretory functions. The parasympathetic nerves convey visceral sensory pathways from and excitatory pathways to the small intestine and colon. Parasympathetic fibers via the vagus nerve reach the small intestine and proximal colon along the branches of the superior mesenteric artery. The distal colon is supplied by sacral parasympathetic nerves (S₂₋₄) via the pelvic plexus; these fibers course through the wall of the colon as ascending intracolonic fibers as far as, and in some instances including, the proximal colon. The chief excitatory neurotransmitters controlling motor function are acetylcholine and the tachykinins, such as substance P. The sympathetic nerve supply modulates motor functions and reaches the small intestine and colon alongside their arterial vessels. Sympathetic input to the gut is generally excitatory to sphincters and inhibitory to non-sphincteric muscle. Visceral afferents convey sensation from the gut to the central nervous system (CNS); initially, they course along sympathetic fibers, but as they approach the spinal cord they separate, have cell bodies in the dorsal root ganglion, and enter the dorsal horn of the spinal cord. Afferent signals are conveyed to the brain along the lateral spinothalamic tract and the nociceptive dorsal column pathway and are then projected beyond the thalamus and brainstem to the insula and cerebral cortex to be perceived. Other afferent fibers synapse in the prevertebral ganglia and reflexly modulate intestinal motility, blood flow, and secretion.

INTESTINAL FLUID ABSORPTION AND SECRETION

On an average day, 9 L of fluid enter the gastrointestinal (GI) tract, ~1 L of residual fluid reaches the colon, and the stool excretion of fluid constitutes about 0.2 L/d. The colon has a large capacitance and functional reserve and may recover up to four times its usual volume of 0.8 L/d, provided the rate of flow permits reabsorption to occur. Thus, the colon can partially compensate for excess fluid delivery to the colon that may result from intestinal absorptive or secretory disorders.

In the small intestine and colon, sodium absorption is predominantly electrogenic (i.e., it can be measured as an ionic current across the membrane because there is not an equivalent loss of a cation from the cell), and uptake takes place at the apical membrane; it is compensated for by the export functions of the basolateral sodium pump. There are several active transport proteins at the apical membrane, especially in the small intestine, whereby sodium ion entry is coupled to monosaccharides (e.g., glucose through the transporter SGLT1, or fructose through GLUT-5). Glucose then exits the basal membrane through a specific transport protein, GLUT-5, creating a glucose concentration gradient between the lumen and the intercellular space, drawing water and electrolytes passively from the lumen. A variety of neural and nonneural mediators regulate colonic fluid and electrolyte balance, including cholinergic, adrenergic, and serotonergic mediators. Angiotensin and aldosterone also influence colonic absorption, reflecting the common embryologic development of the distal colonic epithelium and the renal tubules.

SMALL-INTESTINAL MOTILITY

During the fasting period, the motility of the small intestine is characterized by a cyclical event called the migrating motor complex (MMC),

which serves to clear nondigestible residue from the small intestine (the intestinal “housekeeper”). This organized, propagated series of contractions lasts, on average, 4 min, occurs every 60–90 min, and usually involves the entire small intestine. After food ingestion, the small intestine produces irregular, mixing contractions of relatively low amplitude, except in the distal ileum where more powerful contractions occur intermittently and empty the ileum by bolus transfers.

ILEOCOLONIC STORAGE AND SALVAGE

The distal ileum acts as a reservoir, emptying intermittently by bolus movements. This action allows time for salvage of fluids, electrolytes, and nutrients. Segmentation by haustra compartmentalizes the colon and facilitates mixing, retention of residue, and formation of solid stools. There is increased appreciation of the intimate interaction between the colonic function and the luminal ecology. The resident microorganisms, predominantly anaerobic bacteria, in the colon are necessary for the digestion of unabsorbed carbohydrates that reach the colon even in health, thereby providing a vital source of nutrients to the mucosa. Normal colonic flora also keeps pathogens at bay by a variety of mechanisms. In health, the ascending and transverse regions of colon function as reservoirs (average transit time, 15 h), and the descending colon acts as a conduit (average transit time, 3 h). The colon is efficient at conserving sodium and water, a function that is particularly important in sodium-depleted patients in whom the small intestine alone is unable to maintain sodium balance. Diarrhea or constipation may result from alteration in the reservoir function of the proximal colon or the propulsive function of the left colon. Constipation may also result from disturbances of the rectal or sigmoid reservoir, typically as a result of dysfunction of the pelvic floor, the anal sphincters, the coordination of defecation, or dehydration.

COLONIC MOTILITY AND TONE

The small intestinal MMC only rarely continues into the colon. However, short duration or phasic contractions mix colonic contents, and high-amplitude (>75 mmHg) propagated contractions (HAPCs) are sometimes associated with mass movements through the colon and normally occur approximately five times per day, usually on awakening in the morning and postprandially. Increased frequency of HAPCs may result in diarrhea or urgency. The predominant phasic contractions in the colon are irregular and non-propagated and serve a “mixing” function.

Colonic tone refers to the background contractility upon which phasic contractile activity (typically contractions lasting <15 s) is superimposed. It is an important cofactor in the colon’s capacitance (volume accommodation) and sensation.

COLONIC MOTILITY AFTER MEAL INGESTION

After meal ingestion, colonic phasic and tonic contractility increase for a period of ~2 h. The initial phase (~10 min) is mediated by the vagus nerve in response to mechanical distention of the stomach. The subsequent response of the colon requires caloric stimulation (e.g., intake of at least 500 kcal) and is mediated, at least in part, by hormones (e.g., gastrin and serotonin).

DEFECATION

Tonic contraction of the puborectalis muscle, which forms a sling around the rectoanal junction, is important to maintain continence; during defecation, sacral parasympathetic nerves relax this muscle, facilitating the straightening of the rectoanal angle (Fig. 55-1). Distention of the rectum results in transient relaxation of the internal anal sphincter via intrinsic and

reflex sympathetic innervation. As sigmoid and rectal contractions, as well as straining (Valsalva maneuver), which increases intraabdominal pressure, increase the pressure within the rectum, the rectosigmoid angle opens by >15°. Voluntary relaxation of the external anal sphincter (striated muscle innervated by the pudendal nerve) in response to the sensation produced by distention permits the evacuation of feces. Defecation can also be delayed voluntarily by contraction of the external anal sphincter.

DIARRHEA

DEFINITION

Diarrhea is loosely defined as passage of abnormally liquid or unformed stools at an increased frequency. For adults on a typical Western diet, stool weight >200 g/d can generally be considered diarrheal. Diarrhea may be further defined as *acute* if <2 weeks, *persistent* if 2–4 weeks, and *chronic* if >4 weeks in duration.

Two common conditions, usually associated with the passage of stool totaling <200 g/d, must be distinguished from diarrhea, because diagnostic and therapeutic algorithms differ. *Pseudodiarrhea*, or the frequent passage of small volumes of stool, is often associated with rectal urgency, tenesmus, or a feeling of incomplete evacuation, and accompanies IBS or proctitis. *Fecal incontinence* is the involuntary discharge of rectal contents and is most often caused by neuromuscular disorders or structural anorectal problems. Diarrhea and urgency, especially if severe, may aggravate or cause incontinence. Pseudodiarrhea and fecal incontinence occur at prevalence rates comparable to or higher than that of chronic diarrhea and should always be considered in patients complaining of “diarrhea.” Overflow diarrhea may occur in nursing home patients due to fecal impaction that is readily detectable by rectal examination. A careful history and physical examination generally allow these conditions to be discriminated from true diarrhea.

ACUTE DIARRHEA

More than 90% of cases of acute diarrhea are caused by infectious agents; these cases are often accompanied by vomiting, fever, and abdominal pain. The remaining 10% or so are caused by medications, toxic ingestions, ischemia, food indiscretions, and other conditions.

Infectious Agents Most infectious diarrheas are acquired by fecal-oral transmission or, more commonly, via ingestion of food or water contaminated with pathogens from human or animal feces. In the immunocompetent person, the resident fecal microflora, containing >500 taxonomically distinct species, are rarely the source of diarrhea

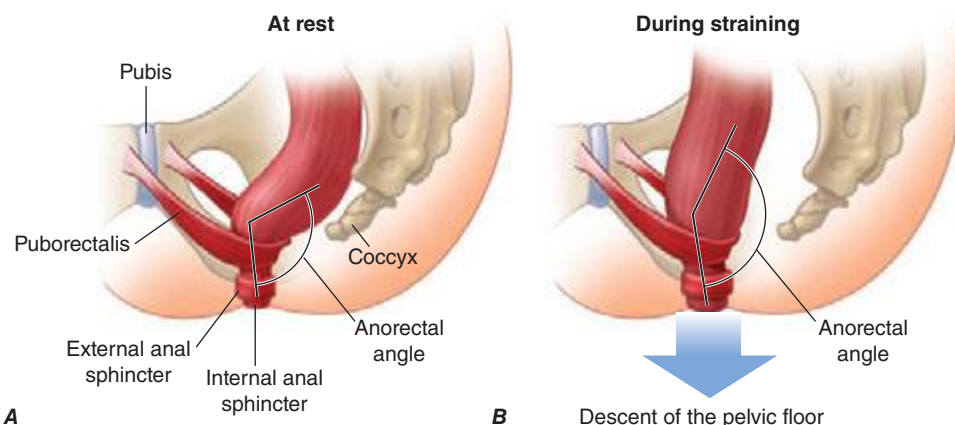


FIGURE 55-1 Sagittal view of the anorectum (A) at rest and (B) during straining to defecate.

Continence is maintained by normal rectal sensation and tonic contraction of the internal anal sphincter and the puborectalis muscle, which wraps around the anorectum, maintaining an anorectal angle between 80° and 110°. During defecation, the pelvic floor muscles (including the puborectalis) relax, allowing the anorectal angle to straighten by at least 15°, and the perineum descends by 1–3.5 cm. The external anal sphincter also relaxes and reduces pressure on the anal canal. (Reproduced with permission from A Lembo, M Camilleri: *N Engl J Med* 349:1360, 2003.)

and may actually play a role in suppressing the growth of ingested pathogens. Disturbances of flora by antibiotics can lead to diarrhea by reducing the digestive function or by allowing the overgrowth of pathogens, such as *Clostridium difficile* (Chap. 161). Acute infection or injury occurs when the ingested agent overwhelms or bypasses the host's mucosal immune and nonimmune (gastric acid, digestive enzymes, mucus secretion, peristalsis, and suppressive resident flora) defenses. Established clinical associations with specific enteropathogens may offer diagnostic clues.

In the United States, five high-risk groups are recognized:

1. **Travelers.** Nearly 40% of tourists to endemic regions of Latin America, Africa, and Asia develop so-called traveler's diarrhea, most commonly due to enterotoxigenic or enteroaggregative *Escherichia coli* as well as to *Campylobacter*, *Shigella*, *Aeromonas*, norovirus, *Coronavirus*, and *Salmonella*. Visitors to Russia (especially St. Petersburg) may have increased risk of *Giardia*-associated diarrhea; visitors to Nepal may acquire *Cyclospora*. Campers, backpackers, and swimmers in wilderness areas may become infected with *Giardia*. Cruise ships may be affected by outbreaks of gastroenteritis caused by agents such as norovirus.
2. **Consumers of certain foods.** Diarrhea closely following food consumption at a picnic, banquet, or restaurant may suggest infection with *Salmonella*, *Campylobacter*, or *Shigella* from chicken; enterohemorrhagic *E. coli* (O157:H7) from undercooked hamburger; *Bacillus cereus* from fried rice or other reheated food; *Staphylococcus aureus* or *Salmonella* from mayonnaise or creams; *Salmonella* from eggs; *Listeria* from uncooked foods or soft cheeses; and *Vibrio* species, *Salmonella*, or acute hepatitis A from seafood, especially if raw. State departments of public health issue communications regarding food-related illnesses, which may have originated domestically or been imported, but ultimately cause epidemics in the United States (e.g., the *Cyclospora* epidemic of 2013 in midwestern states that resulted from bagged salads).
3. **Immunodeficient persons.** Individuals at risk for diarrhea include those with either primary immunodeficiency (e.g., IgA deficiency, common variable hypogammaglobulinemia, chronic granulomatous disease) or the much more common secondary immunodeficiency states

(e.g., AIDS, senescence, pharmacologic suppression). Common enteric pathogens often cause a more severe and protracted diarrheal illness, and, particularly in persons with AIDS, opportunistic infections, such as by *Mycobacterium* species, certain viruses (cytomegalovirus, adenovirus, and herpes simplex), and protozoa (*Cryptosporidium*, *Isospora belli*, Microsporida, and *Blastocystis hominis*) may also play a role (Chap. 226). In patients with AIDS, agents transmitted venereally per rectum (e.g., *Neisseria gonorrhoeae*, *Treponema pallidum*, *Chlamydia*) may contribute to proctocolitis. Persons with hemochromatosis are especially prone to invasive, even fatal, enteric infections with *Vibrio* species and *Yersinia* infections and should avoid raw fish.

4. **Daycare attendees and their family members.** Infections with *Shigella*, *Giardia*, *Cryptosporidium*, rotavirus, and other agents are very common and should be considered.
5. **Institutionalized persons.** Infectious diarrhea is one of the most frequent categories of nosocomial infections in many hospitals and long-term care facilities; the causes are a variety of microorganisms but most commonly *C. difficile*. *C. difficile* can affect those with no history of antibiotic use and may be acquired in the community.

The pathophysiology underlying acute diarrhea by infectious agents produces specific clinical features that may also be helpful in diagnosis (Table 55-2). Profuse, watery diarrhea secondary to small-bowel hypersecretion occurs with ingestion of preformed bacterial toxins, enterotoxin-producing bacteria, and enteroadherent pathogens. Diarrhea associated with marked vomiting and minimal or no fever may occur abruptly within a few hours after ingestion of the former two types; vomiting is usually less, abdominal cramping or bloating is greater, and fever is higher with the latter. Cytotoxin-producing and invasive microorganisms all cause high fever and abdominal pain. Invasive bacteria and *Entamoeba histolytica* often cause bloody diarrhea (referred to as *dysentery*). *Yersinia* invades the terminal ileal and proximal colon mucosa and may cause especially severe abdominal pain with tenderness mimicking acute appendicitis.

Finally, infectious diarrhea may be associated with systemic manifestations. Reactive arthritis (formerly known as Reiter's syndrome), arthritis, urethritis, and conjunctivitis may accompany or follow

TABLE 55-2 ASSOCIATION BETWEEN PATHOBIOLOGY OF CAUSATIVE AGENTS AND CLINICAL FEATURES IN ACUTE INFECTIOUS DIARRHEA

Pathobiology/Agents	Incubation Period	Vomiting	Abdominal Pain	Fever	Diarrhea
Toxin producers					
Preformed toxin					
<i>Bacillus cereus</i> , <i>Staphylococcus aureus</i> , <i>Clostridium perfringens</i>	1–8 h 8–24 h	3–4+	1–2+	0–1+	3–4+, watery
Enterotoxin					
<i>Vibrio cholerae</i> , enterotoxigenic <i>Escherichia coli</i> , <i>Klebsiella pneumoniae</i> , <i>Aeromonas</i> species	8–72 h	2–4+	1–2+	0–1+	3–4+, watery
Enteroadherent					
Enteropathogenic and enteroadherent <i>E. coli</i> , <i>Giardia</i> organisms, cryptosporidiosis, helminths	1–8 d	0–1+	1–3+	0–2+	1–2+, watery, mushy
Cytotoxin producers					
<i>C. difficile</i>	1–3 d	0–1+	3–4+	1–2+	1–3+, usually watery, occasionally bloody
Hemorrhagic <i>E. coli</i>	12–72 h	0–1+	3–4+	1–2+	1–3+, initially watery, quickly bloody
Invasive organisms					
Minimal inflammation					
Rotavirus and norovirus	1–3 d	1–3+	2–3+	3–4+	1–3+, watery
Variable inflammation					
<i>Salmonella</i> , <i>Campylobacter</i> , and <i>Aeromonas</i> species, <i>Vibrio parahaemolyticus</i> , <i>Yersinia</i>	12 h–11 d	0–3+	2–4+	3–4+	1–4+, watery or bloody
Severe inflammation					
<i>Shigella</i> species, enteroinvasive <i>E. coli</i> , <i>Entamoeba histolytica</i>	12 h–8 d	0–1+	3–4+	3–4+	1–2+, bloody

Source: Adapted from DW Powell, in T Yamada (ed): *Textbook of Gastroenterology and Hepatology*, 4th ed. Philadelphia, Lippincott Williams & Wilkins, 2003.

infections by *Salmonella*, *Campylobacter*, *Shigella*, and *Yersinia*. Yersiniosis may also lead to an autoimmune-type thyroiditis, pericarditis, and glomerulonephritis. Both enterohemorrhagic *E. coli* (O157:H7) and *Shigella* can lead to the *hemolytic-uremic syndrome* with an attendant high mortality rate. The syndrome of postinfectious IBS has now been recognized as a complication of infectious diarrhea. Similarly, acute gastroenteritis may precede the diagnosis of celiac disease or Crohn's disease. Acute diarrhea can also be a major symptom of several systemic infections including *viral hepatitis*, *listeriosis*, *legionellosis*, and *toxic shock syndrome*.

Other Causes Side effects from medications are probably the most common noninfectious causes of acute diarrhea, and etiology may be suggested by a temporal association between use and symptom onset. Although innumerable medications may produce diarrhea, some of the more frequently incriminated include antibiotics, cardiac antidysrhythmics, antihypertensives, nonsteroidal anti-inflammatory drugs (NSAIDs), certain antidepressants, chemotherapeutic agents, bronchodilators, antacids, and laxatives. Occlusive or nonocclusive *ischemic colitis* typically occurs in persons >50 years; often presents as acute lower abdominal pain preceding watery, then bloody diarrhea; and generally results in acute inflammatory changes in the sigmoid or left colon while sparing the rectum. Acute diarrhea may accompany colonic *diverticulitis* and *graft-versus-host disease*. Acute diarrhea, often associated with systemic compromise, can follow ingestion of toxins including organophosphate insecticides; amanita and other mushrooms; arsenic; and preformed environmental toxins in seafood, such as ciguatera and scombroid. Acute anaphylaxis to food ingestion can have a similar presentation. Conditions causing chronic diarrhea can also be confused with acute diarrhea early in their course. This confusion may occur with inflammatory bowel disease (IBD) and some of the other inflammatory chronic diarrheas that may have an abrupt rather than insidious onset and exhibit features that mimic infection.

APPROACH TO THE PATIENT: Acute Diarrhea

The decision to evaluate acute diarrhea depends on its severity and duration and on various host factors (Fig. 55-2). Most episodes of acute diarrhea are mild and self-limited and do not justify the cost and potential morbidity rate of diagnostic or pharmacologic interventions. Indications for evaluation include profuse diarrhea with dehydration, grossly bloody stools, fever $\geq 38.5^{\circ}\text{C}$ ($\geq 101^{\circ}\text{F}$), duration >48 h without improvement, recent antibiotic use, new community outbreaks, associated severe abdominal pain in patients >50 years, and elderly (≥ 70 years) or immunocompromised patients. In some cases of moderately severe febrile diarrhea associated with fecal leukocytes (or increased fecal levels of the leukocyte proteins, such as calprotectin) or with gross blood, a diagnostic evaluation might be avoided in favor of an empirical antibiotic trial (see below).

The cornerstone of diagnosis in those suspected of severe acute infectious diarrhea is microbiologic analysis of the stool. Workup includes cultures for bacterial and viral pathogens, direct inspection for ova and parasites, and immunoassays for certain bacterial toxins (*C. difficile*), viral antigens (rotavirus), and protozoal antigens (*Giardia*, *E. histolytica*). The aforementioned clinical and epidemiologic associations may assist in focusing the evaluation. If a particular pathogen or set of possible pathogens is so implicated, then either the whole panel of routine studies may not be necessary or, in some instances, special cultures may be appropriate as for enterohemorrhagic and other types of *E. coli*, *Vibrio* species, and *Yersinia*. Molecular diagnosis of pathogens in stool can be made by identification of unique DNA sequences; and evolving microarray technologies have led to more rapid, sensitive, specific, and cost-effective diagnosis.

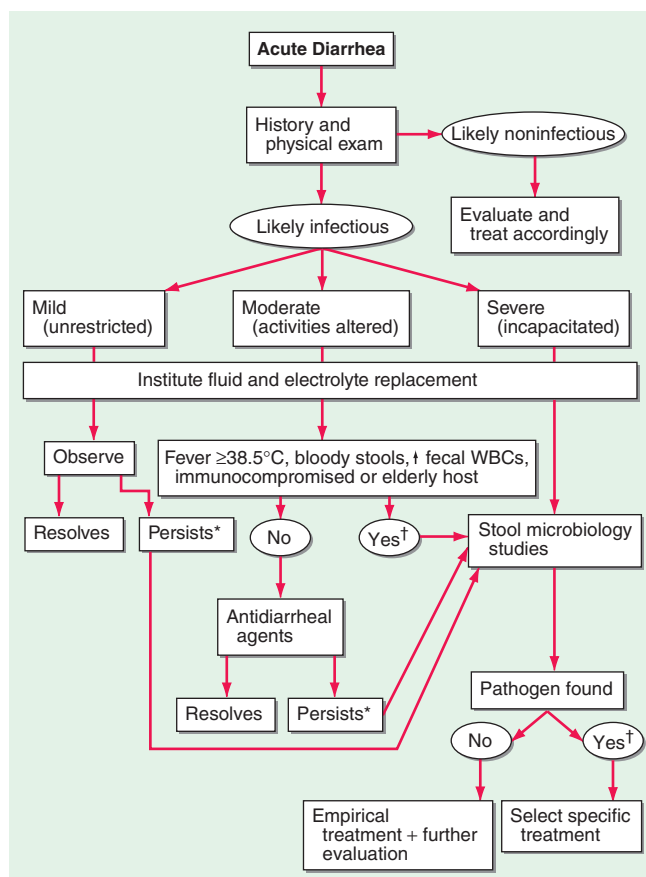


FIGURE 55-2 Algorithm for the management of acute diarrhea. Consider empirical treatment before evaluation with (*) metronidazole and with (†) quinolone. WBCs, white blood cells.

Persistent diarrhea is commonly due to *Giardia* (Chap. 247), but additional causative organisms that should be considered include *C. difficile* (especially if antibiotics had been administered), *E. histolytica*, *Cryptosporidium*, *Campylobacter*, and others. If stool studies are unrevealing, flexible sigmoidoscopy with biopsies and upper endoscopy with duodenal aspirates and biopsies may be indicated. Brainerd diarrhea is an increasingly recognized entity characterized by an abrupt-onset diarrhea that persists for at least 4 weeks, but may last 1–3 years, and is thought to be of infectious origin. It may be associated with subtle inflammation of the distal small intestine or proximal colon.

Structural examination by sigmoidoscopy, colonoscopy, or abdominal computed tomography (CT) scanning (or other imaging approaches) may be appropriate in patients with uncharacterized persistent diarrhea to exclude IBD or as an initial approach in patients with suspected noninfectious acute diarrhea such as might be caused by ischemic colitis, diverticulitis, or partial bowel obstruction.

TREATMENT ACUTE DIARRHEA

Fluid and electrolyte replacement are of central importance to all forms of acute diarrhea. Fluid replacement alone may suffice for mild cases. Oral sugar-electrolyte solutions (iso-osmolar sport drinks or designed formulations) should be instituted promptly with severe diarrhea to limit dehydration, which is the major cause of death. Profoundly dehydrated patients, especially infants and the elderly, require IV rehydration.

In moderately severe nonfebrile and nonbloody diarrhea, antimotility and antisecretory agents such as loperamide can be useful adjuncts to control symptoms. Such agents should be avoided with

febrile dysentery, which may be exacerbated or prolonged by them. Bismuth subsalicylate may reduce symptoms of vomiting and diarrhea but should not be used to treat immunocompromised patients or those with renal impairment because of the risk of bismuth encephalopathy.

Judicious use of antibiotics is appropriate in selected instances of acute diarrhea and may reduce its severity and duration (Fig. 55-2). Many physicians treat moderately to severely ill patients with febrile dysentery empirically without diagnostic evaluation using a quinolone, such as ciprofloxacin (500 mg bid for 3–5 d). Empirical treatment can also be considered for suspected giardiasis with metronidazole (250 mg qid for 7 d). Selection of antibiotics and dosage regimens are otherwise dictated by specific pathogens, geographic patterns of resistance, and conditions found (Chaps. 160, 186, and 190–196). Antibiotic coverage is indicated, whether or not a causative organism is discovered, in patients who are immunocompromised, have mechanical heart valves or recent vascular grafts, or are elderly. Bismuth subsalicylate may reduce the frequency of traveler's diarrhea. Antibiotic prophylaxis is only indicated for certain patients traveling to high-risk countries in whom the likelihood or seriousness of acquired diarrhea would be especially high, including those with immunocompromise, IBD, hemochromatosis, or gastric achlorhydria. Use of ciprofloxacin, azithromycin, or rifaximin may reduce bacterial diarrhea in such travelers by 90%, though rifaximin is not suitable for invasive disease, but rather as treatment for uncomplicated traveler's diarrhea. Finally, physicians should be vigilant to identify if an outbreak of diarrheal illness is occurring and to alert the public health authorities promptly. This may reduce the ultimate size of the affected population.

CHRONIC DIARRHEA

Diarrhea lasting >4 weeks warrants evaluation to exclude serious underlying pathology. In contrast to acute diarrhea, most of the causes of chronic diarrhea are noninfectious. The classification of chronic diarrhea by pathophysiologic mechanism facilitates a rational approach to management, although many diseases cause diarrhea by more than one mechanism (Table 55-3).

Secretory Causes Secretory diarrheas are due to derangements in fluid and electrolyte transport across the enterocolonic mucosa. They are characterized clinically by watery, large-volume fecal outputs that are typically painless and persist with fasting. Because there is no malabsorbed solute, stool osmolality is accounted for by normal endogenous electrolytes with no fecal osmotic gap.

MEDICATIONS Side effects from regular ingestion of drugs and toxins are the most common secretory causes of chronic diarrhea. Hundreds of prescription and over-the-counter medications (see earlier section, “Acute Diarrhea, Other Causes”) may produce diarrhea. Surreptitious or habitual use of stimulant laxatives (e.g., senna, cascara, bisacodyl, ricinoleic acid [castor oil]) must also be considered. Chronic ethanol consumption may cause a secretory-type diarrhea due to enterocyte injury with impaired sodium and water absorption as well as rapid transit and other alterations. Inadvertent ingestion of certain environmental toxins (e.g., arsenic) may lead to chronic rather than acute forms of diarrhea. Certain bacterial infections may occasionally persist and be associated with a secretory-type diarrhea.

BOWEL RESECTION, MUCOSAL DISEASE, OR ENTEROCOLIC FISTULA These conditions may result in a secretory-type diarrhea because of inadequate surface for reabsorption of secreted fluids and electrolytes. Unlike other secretory diarrheas, this subset of conditions tends to worsen with eating. With disease (e.g., Crohn's ileitis) or resection of <100 cm of terminal ileum, dihydroxy bile acids may escape absorption and stimulate colonic secretion (choleraic diarrhea). This mechanism may contribute to so-called *idiopathic secretory diarrhea or bile acid diarrhea (BAD)*, in which bile acids are functionally malabsorbed from a normal-appearing terminal ileum. This *idiopathic bile acid malabsorption (BAM)* may account for an average of 40% of unexplained chronic diarrhea. Reduced negative feedback regulation of bile acid

TABLE 55-3 MAJOR CAUSES OF CHRONIC DIARRHEA ACCORDING TO PREDOMINANT PATHOPHYSIOLOGIC MECHANISM

Secretory Causes
Exogenous stimulant laxatives
Chronic ethanol ingestion
Other drugs and toxins
Endogenous laxatives (dihydroxy bile acids)
Idiopathic secretory diarrhea or bile acid diarrhea
Certain bacterial infections
Bowel resection, disease, or fistula (↓ absorption)
Partial bowel obstruction or fecal impaction
Hormone-producing tumors (carcinoid, VIPoma, medullary cancer of thyroid, mastocytosis, gastrinoma, colorectal villous adenoma)
Addison's disease
Congenital electrolyte absorption defects
Osmotic Causes
Osmotic laxatives (Mg ²⁺ , PO ₄ ⁻³ , SO ₄ ⁻²)
Lactase and other disaccharide deficiencies
Nonabsorbable carbohydrates (sorbitol, lactulose, polyethylene glycol)
Gluten and FODMAP intolerance
Steatorrheal Causes
Intraluminal maldigestion (pancreatic exocrine insufficiency, bacterial overgrowth, bariatric surgery, liver disease)
Mucosal malabsorption (celiac sprue, Whipple's disease, infections, abetalipoproteinemia, ischemia, drug-induced enteropathy)
Postmucosal obstruction (1° or 2° lymphatic obstruction)
Inflammatory Causes
Idiopathic inflammatory bowel disease (Crohn's, chronic ulcerative colitis)
Lymphocytic and collagenous colitis
Immune-related mucosal disease (1° or 2° immunodeficiencies, food allergy, eosinophilic gastroenteritis, graft-versus-host disease)
Infections (invasive bacteria, viruses, and parasites, Brainerd diarrhea)
Radiation injury
Gastrointestinal malignancies
Dysmotile Causes
Irritable bowel syndrome (including postinfectious IBS)
Visceral neuromyopathies
Hyperthyroidism
Drugs (prokinetic agents)
Postvagotomy
Factitial Causes
Munchausen
Eating disorders
Iatrogenic Causes
Cholecystectomy
Ileal resection
Bariatric surgery
Vagotomy, fundoplication

Abbreviation: FODMAP, fermentable oligosaccharides, disaccharides, monosaccharides, and polyols.

synthesis in hepatocytes by fibroblast growth factor 19 (FGF-19) produced by ileal enterocytes results in a degree of bile-acid synthesis that exceeds the normal capacity for ileal reabsorption, producing BAD. An alternative cause of BAD is a genetic variation in the receptor proteins (β-klotho and fibroblast growth factor 4) on the hepatocyte that normally mediate the effect of FGF-19. Dysfunction of these proteins prevents FGF-19 inhibition of hepatocyte bile acid synthesis.

Partial bowel obstruction, ostomy stricture, or fecal impaction may paradoxically lead to increased fecal output due to fluid hypersecretion.

HORMONES Although uncommon, the classic examples of secretory diarrhea are those mediated by hormones. *Metastatic gastrointestinal*

carcinoid tumors or, rarely, *primary bronchial carcinoids* may produce watery diarrhea alone or as part of the carcinoid syndrome that comprises episodic flushing, wheezing, dyspnea, and right-sided valvular heart disease. Diarrhea is due to the release into the circulation of potent intestinal secretagogues including serotonin, histamine, prostaglandins, and various kinins. Pellagra-like skin lesions may rarely occur as the result of serotonin overproduction with niacin depletion. *Gastrinoma*, one of the most common neuroendocrine tumors, most typically presents with refractory peptic ulcers, but diarrhea occurs in up to one-third of cases and may be the only clinical manifestation in 10%. While other secretagogues released with gastrin may play a role, the diarrhea most often results from fat maldigestion owing to pancreatic enzyme inactivation by low intraduodenal pH. The watery diarrhea hypokalemia achlorhydria syndrome, also called *pancreatic cholera*, is due to a non- β cell pancreatic adenoma, referred to as a *VIPoma*, that secretes VIP and a host of other peptide hormones including pancreatic polypeptide, secretin, gastrin, gastrin-inhibitory polypeptide (also called glucose-dependent insulinotropic peptide), neurotensin, calcitonin, and prostaglandins. The secretory diarrhea is often massive with stool volumes >3 L/d; daily volumes as high as 20 L have been reported. Life-threatening dehydration; neuromuscular dysfunction from associated hypokalemia, hypomagnesemia, or hypercalcemia; flushing; and hyperglycemia may accompany a *VIPoma*. *Medullary carcinoma of the thyroid* may present with watery diarrhea caused by calcitonin, other secretory peptides, or prostaglandins. Prominent diarrhea is often associated with metastatic disease and poor prognosis. *Systemic mastocytosis*, which may be associated with the skin lesion urticaria pigmentosa, may cause diarrhea that is either secretory and mediated by histamine or inflammatory due to intestinal infiltration by mast cells. Large *colorectal villous adenomas* may rarely be associated with a secretory diarrhea that may cause hypokalemia, can be inhibited by NSAIDs, and are apparently mediated by prostaglandins.

CONGENITAL DEFECTS IN ION ABSORPTION Rarely, defects in specific carriers associated with ion absorption cause watery diarrhea from birth. These disorders include defective $\text{Cl}^-/\text{HCO}_3^-$ exchange (*congenital chloridorrhea*) with alkalosis (which results from a mutated *DRA* [down-regulated in adenoma] gene) and defective Na^+/H^+ exchange (*congenital sodium diarrhea*), which results from a mutation in the *NHE3* (sodium-hydrogen exchanger) gene and results in acidosis.

Some hormone deficiencies may be associated with watery diarrhea, such as occurs with adrenocortical insufficiency (Addison's disease) that may be accompanied by skin hyperpigmentation.

Osmotic Causes Osmotic diarrhea occurs when ingested, poorly absorbable, osmotically active solutes draw enough fluid into the lumen to exceed the reabsorptive capacity of the colon. Fecal water output increases in proportion to such a solute load. Osmotic diarrhea characteristically ceases with fasting or with discontinuation of the causative agent.

OSMOTIC LAXATIVES Ingestion of magnesium-containing antacids, health supplements, or laxatives may induce osmotic diarrhea typified by a stool osmotic gap (>50 mosmol/L): serum osmolarity (typically 290 mosmol/kg) $- (2 \times [\text{fecal sodium} + \text{potassium concentration}])$. Measurement of fecal osmolarity is no longer recommended because, even when measured immediately after evacuation, it may be erroneous because carbohydrates are metabolized by colonic bacteria, causing an increase in osmolarity.

CARBOHYDRATE MALABSORPTION Carbohydrate malabsorption due to acquired or congenital defects in brush-border disaccharidases and other enzymes leads to osmotic diarrhea with a low pH. One of the most common causes of chronic diarrhea in adults is *lactase deficiency*, which affects three-fourths of nonwhites worldwide and 5–30% of persons in the United States; the total lactose load at any one time influences the symptoms experienced. Most patients learn to avoid milk products without requiring treatment with enzyme supplements. Some sugars, such as sorbitol, lactulose, or fructose, are frequently malabsorbed, and diarrhea ensues with ingestion of medications, gum, or candies sweetened with these poorly or incompletely absorbed sugars.

WHEAT AND FODMAP INTOLERANCE Chronic diarrhea, bloating, and abdominal pain are recognized as symptoms of nonceliac gluten intolerance (which is associated with impaired intestinal or colonic barrier function) and intolerance of fermentable oligosaccharides, disaccharides, monosaccharides, and polyols (FODMAPs). The latter's effects represent the interaction between the GI microbiome and the nutrients.

Steatorrheal Causes Fat malabsorption may lead to greasy, foul-smelling, difficult-to-flush diarrhea often associated with weight loss and nutritional deficiencies due to concomitant malabsorption of amino acids and vitamins. Increased fecal output is caused by the osmotic effects of fatty acids, especially after bacterial hydroxylation, and, to a lesser extent, by the neutral fat. Quantitatively, steatorrhea is defined as stool fat exceeding the normal 7 g/d; rapid-transit diarrhea may result in fecal fat up to 14 g/d; daily fecal fat averages 15–25 g with small-intestinal diseases and is often >32 g with pancreatic exocrine insufficiency. Intraluminal maldigestion, mucosal malabsorption, or lymphatic obstruction may produce steatorrhea.

INTRALUMINAL MALDIGESTION This condition most commonly results from pancreatic exocrine insufficiency, which occurs when $>90\%$ of pancreatic secretory function is lost. *Chronic pancreatitis*, usually a sequel of ethanol abuse, most frequently causes pancreatic insufficiency. Other causes include *cystic fibrosis*; *pancreatic duct obstruction*; and, rarely, *somatostatinoma*. Bacterial overgrowth in the small intestine may deconjugate bile acids and alter micelle formation, impairing fat digestion; it occurs with stasis from a blind-loop, small-bowel diverticulum or dysmotility and is especially likely in the elderly. Finally, cirrhosis or biliary obstruction may lead to mild steatorrhea due to deficient intraluminal bile acid concentration.

MUCOSAL MALABSORPTION Mucosal malabsorption occurs from a variety of enteropathies, but it most commonly occurs from *celiac disease*. This gluten-sensitive enteropathy affects all ages and is characterized by villous atrophy and crypt hyperplasia in the proximal small bowel and can present with fatty diarrhea associated with multiple nutritional deficiencies of varying severity. Celiac disease is much more frequent than previously thought; it affects $\sim 1\%$ of the population, frequently presents without steatorrhea, can mimic IBS, and has many other GI and extraintestinal manifestations. *Tropical sprue* may produce a similar histologic and clinical syndrome but occurs in residents of or travelers to tropical climates; abrupt onset and response to antibiotics suggest an infectious etiology. *Whipple's disease*, due to the bacillus *Tropheryma whipplei* and histiocytic infiltration of the small-bowel mucosa, is a less common cause of steatorrhea that most typically occurs in young or middle-aged men; it is frequently associated with arthralgias, fever, lymphadenopathy, and extreme fatigue, and it may affect the CNS and endocardium. A similar clinical and histologic picture results from *Mycobacterium avium-intracellulare* infection in patients with AIDS. *Abetalipoproteinemia* is a rare defect of chylomicron formation and fat malabsorption in children, associated with acanthocytic erythrocytes, ataxia, and retinitis pigmentosa. Several other conditions may cause mucosal malabsorption including infections, especially with protozoa such as *Giardia*; numerous medications (e.g., olmesartan, mycophenolate mofetil, colchicine, cholestyramine, neomycin); amyloidosis; and chronic ischemia.

POSTMUCOSAL LYMPHATIC OBSTRUCTION The pathophysiology of this condition, which is due to the rare *congenital intestinal lymphangiectasia* or to *acquired lymphatic obstruction* secondary to trauma, tumor, cardiac disease or infection, leads to the unique constellation of fat malabsorption with enteric losses of protein (often causing edema) and lymphocytopenia. Carbohydrate and amino acid absorption are preserved.

Inflammatory Causes Inflammatory diarrheas are generally accompanied by pain, fever, bleeding, or other manifestations of inflammation. The mechanism of diarrhea may not only be exudation but, depending on lesion site, may include fat malabsorption, disrupted fluid/electrolyte absorption, and hypersecretion or hypermotility from

release of cytokines and other inflammatory mediators. The unifying feature on stool analysis is the presence of leukocytes or leukocyte-derived proteins such as calprotectin. With severe inflammation, exudative protein loss can lead to anasarca (generalized edema). Any middle-aged or older person with chronic inflammatory-type diarrhea, especially with blood, should be carefully evaluated to exclude a colorectal tumor.

IDIOPATHIC INFLAMMATORY BOWEL DISEASE The illnesses in this category, which include *Crohn's disease* and *chronic ulcerative colitis*, are among the most common organic causes of chronic diarrhea in adults and range in severity from mild to fulminant and life-threatening. They may be associated with uveitis, polyarthralgias, cholestatic liver disease (primary sclerosing cholangitis), and skin lesions (erythema nodosum, pyoderma gangrenosum). *Microscopic colitis*, including both lymphocytic and *collagenous colitis*, is an increasingly recognized cause of chronic watery diarrhea, especially in middle-aged women and those on NSAIDs, statins, proton pump inhibitors (PPIs), and selective serotonin reuptake inhibitors (SSRIs); biopsy of a normal-appearing colon is required for histologic diagnosis. It may coexist with symptoms suggesting IBS or with celiac sprue or drug-induced enteropathy. It typically responds well to anti-inflammatory drugs (e.g., bismuth), to the opioid agonist loperamide, or to budesonide.

PRIMARY OR SECONDARY FORMS OF IMMUNODEFICIENCY Immunodeficiency may lead to prolonged infectious diarrhea. With selective IgA deficiency or common variable *hypogammaglobulinemia*, diarrhea is particularly prevalent and often the result of giardiasis, bacterial overgrowth, or sprue.

EOSINOPHILIC GASTROENTERITIS Eosinophil infiltration of the mucosa, muscularis, or serosa at any level of the GI tract may cause diarrhea, pain, vomiting, or ascites. Affected patients often have an atopic history, Charcot-Leyden crystals due to extruded eosinophil contents may be seen on microscopic inspection of stool, and peripheral eosinophilia is present in 50–75% of patients. While hypersensitivity to certain foods occurs in adults, true food allergy causing chronic diarrhea is rare.

OTHER CAUSES Chronic inflammatory diarrhea may be caused by *radiation enterocolitis*, *chronic graft-versus-host disease*, *Behçet's syndrome*, and *Cronkhite-Canada syndrome*, among others.

Dysmotility Causes Rapid transit may accompany many diarrheas as a secondary or contributing phenomenon, but primary dysmotility is an unusual etiology of true diarrhea. Stool features often suggest a secretory diarrhea, but mild steatorrhea of up to 14 g of fat per day can be produced by maldigestion from rapid transit alone. *Hyperthyroidism*, *carcinoid syndrome*, and certain drugs (e.g., prostaglandins, prokinetic agents) may produce hypermotility with resultant diarrhea. Primary visceral neuromyopathies or idiopathic acquired intestinal pseudoobstruction may lead to stasis with secondary bacterial overgrowth causing diarrhea. *Diabetic diarrhea*, often accompanied by peripheral and generalized autonomic neuropathies, may occur in part because of intestinal dysmotility.

The exceedingly common IBS (10% point prevalence, 1–2% per year incidence) is characterized by disturbed intestinal and colonic motor and sensory responses to various stimuli. Symptoms of stool frequency typically cease at night, alternate with periods of constipation, are accompanied by abdominal pain relieved with defecation, and rarely result in weight loss.

Factitial Causes Factitial diarrhea accounts for up to 15% of unexplained diarrheas referred to tertiary care centers. Either as a form of *Munchausen syndrome* (deception or self-injury for secondary gain) or *eating disorders*, some patients covertly self-administer laxatives alone or in combination with other medications (e.g., diuretics) or surreptitiously add water or urine to stool sent for analysis. Such patients are typically women, often with histories of psychiatric illness, and disproportionately from careers in health care. Hypotension and hypokalemia are common co-presenting features. The evaluation of such patients may be difficult: contamination of the stool with water or urine is suggested by very low or high stool osmolarity, respectively.

Such patients often deny this possibility when confronted, but they do benefit from psychiatric counseling when they acknowledge their behavior.

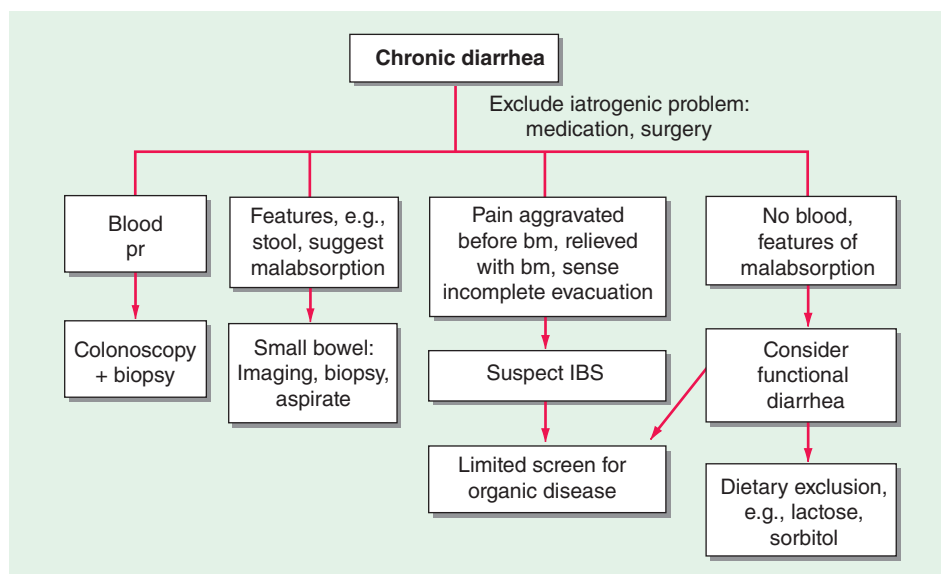
APPROACH TO THE PATIENT: Chronic Diarrhea

The laboratory tools available to evaluate the very common problem of chronic diarrhea are extensive, and many are costly and invasive. As such, the diagnostic evaluation must be rationally directed by a careful history, including medications, and physical examination (Fig. 55-3A). When this strategy is unrevealing, simple triage tests are often warranted to direct the choice of more complex investigations (Fig. 55-3B). The history, physical examination (Table 55-4), and routine blood studies should attempt to characterize the mechanism of diarrhea, identify diagnostically helpful associations, and assess the patient's fluid/electrolyte and nutritional status. Patients should be questioned about the onset, duration, pattern, aggravating (especially diet) and relieving factors, and stool characteristics of their diarrhea. The presence or absence of fecal incontinence, fever, weight loss, pain, certain exposures (travel, medications, contacts with diarrhea), and common extraintestinal manifestations (skin changes, arthralgias, oral aphthous ulcers) should be noted. A family history of IBD or sprue may indicate those possibilities. Physical findings may offer clues such as a thyroid mass, wheezing, heart murmurs, edema, hepatomegaly, abdominal masses, lymphadenopathy, mucocutaneous abnormalities, perianal fistulas, or anal sphincter laxity. Peripheral blood leukocytosis, elevated sedimentation rate, or C-reactive protein suggests inflammation; anemia reflects blood loss or nutritional deficiencies; or eosinophilia may occur with parasitoses, neoplasia, collagen-vascular disease, allergy, or eosinophilic gastroenteritis. Blood chemistries may demonstrate electrolyte, hepatic, or other metabolic disturbances. Measuring IgA tissue transglutaminase antibodies may help detect celiac disease. Bile acid diarrhea is confirmed by a scintigraphic radiolabeled bile acid retention test; however, this is not available in many countries. Alternative approaches are a screening blood test (serum C4 or FGF-19), measurement of fecal bile acids, or a therapeutic trial with a bile acid sequestrant (e.g., cholestyramine or colesevelam).

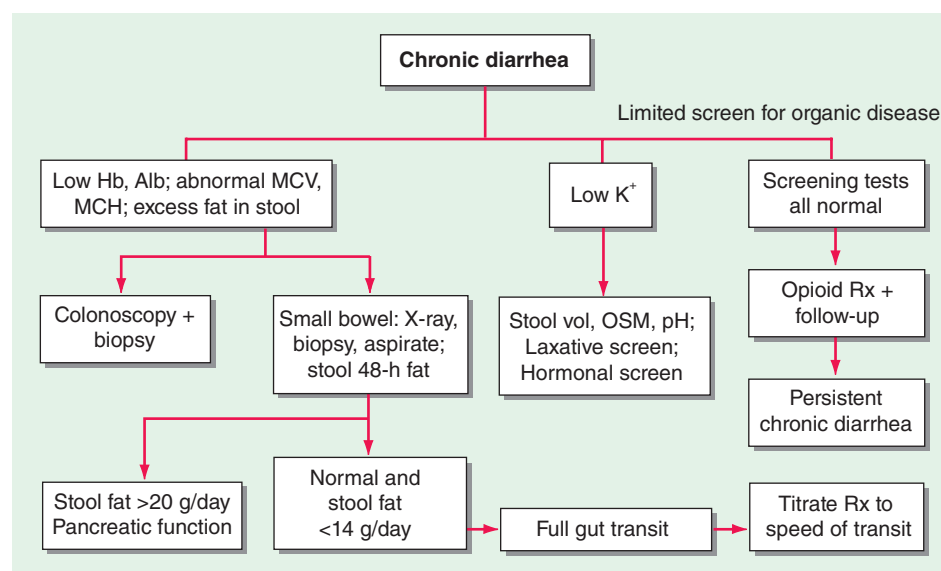
A therapeutic trial is often appropriate, definitive, and highly cost-effective when a specific diagnosis is suggested on the initial physician encounter. For example, chronic watery diarrhea, which ceases with fasting in an otherwise healthy young adult, may justify a trial of a lactose-restricted diet; bloating and diarrhea persisting since a mountain backpacking trip may warrant a trial of metronidazole for likely giardiasis; and postprandial diarrhea persisting following resection of terminal ileum might be due to bile acid malabsorption and be treated with cholestyramine or colesevelam before further evaluation. Persistent symptoms require additional investigation.

Certain diagnoses may be suggested on the initial encounter (e.g., idiopathic IBD); however, additional focused evaluations may be necessary to confirm the diagnosis and characterize the severity or extent of disease so that treatment can be best guided. Patients suspected of having IBS should be initially evaluated with flexible sigmoidoscopy with colorectal biopsies to exclude IBD, or particularly microscopic colitis, which is clinically indistinguishable from IBS with diarrhea; those with normal findings might be reassured and, as indicated, treated empirically with antispasmodics, antidiarrheals, or antidepressants (e.g., tricyclic agents). Any patient who presents with chronic diarrhea and hematochezia should be evaluated with stool microbiologic studies and colonoscopy.

In an estimated two-thirds of cases, the cause for chronic diarrhea remains unclear after the initial encounter, and further testing is required. Quantitative stool collection and analyses can yield



A



B

FIGURE 55-3 Chronic diarrhea. A. Initial management based on accompanying symptoms or features. **B.** Evaluation based on findings from a limited age-appropriate screen for organic disease. Alb, albumin; bm, bowel movement; Hb, hemoglobin; IBS, irritable bowel syndrome; MCH, mean corpuscular hemoglobin; MCV, mean corpuscular volume; OSM, osmolality; pr, per rectum. (Reprinted from M Camilleri: *Clin Gastroenterol Hepatol.* 2:198, 2004.)

important objective data that may establish a diagnosis or characterize the type of diarrhea as a triage for focused additional studies (Fig. 55-3B). If stool weight is >200 g/d, additional stool analyses should be performed that might include electrolyte concentration,

TABLE 55-4 PHYSICAL EXAMINATION IN PATIENTS WITH CHRONIC DIARRHEA

1. Are there general features to suggest malabsorption or inflammatory bowel disease (IBD) such as anemia, dermatitis herpetiformis, edema, or clubbing?
2. Are there features to suggest underlying autonomic neuropathy or collagen-vascular disease in the pupils, orthostasis, skin, hands, or joints?
3. Is there an abdominal mass or tenderness?
4. Are there any abnormalities of rectal mucosa, rectal defects, or altered anal sphincter functions?
5. Are there any mucocutaneous manifestations of systemic disease such as dermatitis herpetiformis (celiac disease), erythema nodosum (ulcerative colitis), flushing (carcinoid), or oral ulcers for IBD or celiac disease?

pH, occult blood testing, leukocyte inspection (or leukocyte protein assay), fat quantitation, and laxative screens.

For secretory diarrheas (watery, normal osmotic gap), possible medication-related side effects or surreptitious laxative use should be reconsidered. Microbiologic studies should be done including fecal bacterial cultures (including media for *Aeromonas* and *Plesiomonas*), inspection for ova and parasites, and *Giardia* antigen assay (the most sensitive test for giardiasis). Small-bowel bacterial overgrowth can be excluded by intestinal aspirates with quantitative cultures or with glucose or lactulose breath tests involving measurement of breath hydrogen, methane, or other metabolite. However, interpretation of these breath tests may be confounded by disturbances of intestinal transit. Upper endoscopy and colonoscopy with biopsies and small-bowel x-rays (formerly barium, but increasingly CT with enterography or magnetic resonance with enteroclysis) are helpful to rule out structural or occult inflammatory disease. When suggested by history or other findings, screens for peptide hormones should

be pursued (e.g., serum gastrin, VIP, calcitonin, and thyroid hormone/thyroid-stimulating hormone, urinary 5-hydroxyindol-acetic acid, histamine).

Further evaluation of osmotic diarrhea should include tests for lactose intolerance and magnesium ingestion, the two most common causes. Low fecal pH suggests carbohydrate malabsorption; lactose malabsorption can be confirmed by lactose breath testing or by a therapeutic trial with lactose exclusion and observation of the effect of lactose challenge (e.g., a liter of milk). Lactase determination on small-bowel biopsy is not generally available. If fecal magnesium or laxative levels are elevated, inadvertent or surreptitious ingestion should be considered and psychiatric help should be sought.

For those with proven fatty diarrhea, endoscopy with small-bowel biopsy (including aspiration for *Giardia* and quantitative cultures) should be performed; if this procedure is unrevealing, a small-bowel radiograph is often an appropriate next step. If small-bowel studies are negative or if pancreatic disease is suspected, pancreatic exocrine insufficiency should be excluded with direct tests, such as the secretin-cholecystokinin stimulation test or a variation that could be performed endoscopically. In general, indirect tests such as assay of fecal elastase or chymotrypsin activity or a bentiromide test have fallen out of favor because of low sensitivity and specificity.

Chronic inflammatory-type diarrheas should be suspected by the presence of blood or leukocytes in the stool. Such findings warrant stool cultures; inspection for ova and parasites; *C. difficile* toxin assay; colonoscopy with biopsies; and, if indicated, small-bowel contrast studies.

TREATMENT CHRONIC DIARRHEA

Treatment of chronic diarrhea depends on the specific etiology and may be curative, suppressive, or empirical. If the cause can be eradicated, treatment is curative as with resection of a colorectal cancer, antibiotic administration for Whipple's disease or tropical sprue, or discontinuation of a drug. For many chronic conditions, diarrhea can be controlled by suppression of the underlying mechanism. Examples include elimination of dietary lactose for lactase deficiency or gluten for celiac sprue, use of glucocorticoids or other anti-inflammatory agents for idiopathic IBDs, bile acid sequestrants for bile acid malabsorption, PPIs for the gastric hypersecretion of gastrinomas, somatostatin analogues such as octreotide for malignant carcinoid syndrome, prostaglandin inhibitors such as indomethacin for medullary carcinoma of the thyroid, and pancreatic enzyme replacement for pancreatic insufficiency. When the specific cause or mechanism of chronic diarrhea evades diagnosis, empirical therapy may be beneficial. Mild opiates, such as diphenoxylate or loperamide, are often helpful in mild or moderate watery diarrhea. For those with more severe diarrhea, codeine or tincture of opium may be beneficial. Such antimotility agents should be avoided with severe IBD, because toxic megacolon may be precipitated. Clonidine, an α_2 -adrenergic agonist, may allow control of diabetic diarrhea, although the medication may be poorly tolerated because it causes postural hypotension. The 5-HT₃ receptor antagonists (e.g., alosetron) may relieve diarrhea and urgency in patients with IBS diarrhea. For all patients with chronic diarrhea, fluid and electrolyte repletion is an important component of management (see "Acute Diarrhea," earlier). Replacement of fat-soluble vitamins may also be necessary in patients with chronic steatorrhea.

CONSTIPATION

DEFINITION

Constipation is a common complaint in clinical practice and usually refers to persistent, difficult, infrequent, or seemingly incomplete

defecation. Because of the wide range of normal bowel habits, constipation is difficult to define precisely. Most persons have at least three bowel movements per week; however, low stool frequency alone is not the sole criterion for the diagnosis of constipation. Many constipated patients have a normal frequency of defecation but complain of excessive straining, hard stools, lower abdominal fullness, or a sense of incomplete evacuation. The individual patient's symptoms must be analyzed in detail to ascertain what is meant by "constipation" or "difficulty" with defecation.

Stool form and consistency are well correlated with the time elapsed from the preceding defecation. Hard, pellety stools occur with slow transit, whereas loose, watery stools are associated with rapid transit. Both small pellety or very large stools are more difficult to expel than normal stools.

The perception of hard stools or excessive straining is more difficult to assess objectively, and the need for enemas or digital disimpaction is a clinically useful way to corroborate the patient's perceptions of difficult defecation.

Psychosocial or cultural factors may also be important. A person whose parents attached great importance to daily defecation will become greatly concerned when he or she misses a daily bowel movement; some children withhold stool to gain attention or because of fear of pain from anal irritation; and some adults habitually ignore or delay the call to have a bowel movement.

CAUSES

Pathophysiologically, chronic constipation generally results from inadequate fiber or fluid intake or from disordered colonic transit or anorectal function. These result from neurogastroenterologic disturbance, certain drugs, advancing age, or in association with a large number of systemic diseases that affect the GI tract (Table 55-5). Constipation of recent onset may be a symptom of significant organic disease such as tumor or stricture. In *idiopathic constipation*, a subset of patients exhibit delayed emptying of the ascending and transverse colon with prolongation of transit (often in the proximal colon) and a reduced frequency of propulsive HAPCs. *Outlet obstruction to defecation* (also called *evacuation disorders*) accounts for about a quarter of cases presenting with constipation in tertiary care and may cause delayed colonic transit, which is usually corrected by biofeedback retraining of the disordered defecation. Constipation of any cause may be exacerbated by hospitalization or chronic illnesses that lead to physical or mental impairment and result in inactivity or physical immobility.

TABLE 55-5 CAUSES OF CONSTIPATION IN ADULTS

Types of Constipation and Causes	Examples
Recent Onset	
Colonic obstruction	Neoplasm; stricture: ischemic, diverticular, inflammatory
Anal sphincter spasm	Anal fissure, painful hemorrhoids
Medications	
Chronic	
Irritable bowel syndrome	Constipation-predominant, alternating
Medications	Ca ²⁺ blockers, antidepressants
Colonic pseudoobstruction	Slow-transit constipation, megacolon (rare Hirschsprung's, Chagas' diseases)
Disorders of rectal evacuation	Pelvic floor dysfunction; anismus; descending perineum syndrome; rectal mucosal prolapse; rectocele
Endocrinopathies	Hypothyroidism, hypercalcemia, pregnancy
Psychiatric disorders	Depression, eating disorders, drugs
Neurologic disease	Parkinsonism, multiple sclerosis, spinal cord injury
Generalized muscle disease	Progressive systemic sclerosis

APPROACH TO THE PATIENT: Constipation

A careful history should explore the patient's symptoms and confirm whether he or she is indeed constipated based on frequency (e.g., fewer than three bowel movements per week), consistency (lumpy/hard), excessive straining, prolonged defecation time, or need to support the perineum or digitate the anorectum to facilitate stool evacuation. In the vast majority of cases (probably >90%), there is no underlying cause (e.g., cancer, depression, or hypothyroidism), and constipation responds to ample hydration, exercise, and supplementation of dietary fiber (15–25 g/d). A good diet and medication history and attention to psychosocial issues are key. Physical examination and, particularly, a rectal examination should exclude fecal impaction and most of the important diseases that present with constipation and possibly indicate features suggesting an evacuation disorder (e.g., high anal sphincter tone, failure of perineal descent, or paradoxical puborectalis contraction during straining to simulate stool evacuation).

The presence of weight loss, rectal bleeding, or anemia with constipation mandates either flexible sigmoidoscopy plus barium enema or colonoscopy alone, particularly in patients >40 years, to exclude structural diseases such as cancer or strictures. Colonoscopy alone is most cost-effective in this setting because it provides an opportunity to biopsy mucosal lesions, perform polypectomy, or dilate strictures. Barium enema has advantages over colonoscopy in the patient with isolated constipation because it is less costly and identifies colonic dilation and all significant mucosal lesions or strictures that are likely to present with constipation. Melanosis coli, or pigmentation of the colon mucosa, indicates the use of anthraquinone laxatives such as cascara or senna; however, this is usually apparent from a careful history. An unexpected disorder such as megacolon or cathartic colon may also be detected by colonic radiographs. Measurement of serum calcium, potassium, and thyroid-stimulating hormone levels will identify rare patients with metabolic disorders.

Patients with more troublesome constipation may not respond to fiber alone and may be helped by a bowel-training regimen, which involves taking an osmotic laxative (e.g., magnesium salts, lactulose, sorbitol, polyethylene glycol) and evacuating with enema or suppository (e.g., glycerine or bisacodyl) as needed. After breakfast, a distraction-free 15–20 min on the toilet without straining is encouraged. Excessive straining may lead to development of hemorrhoids, and, if there is weakness of the pelvic floor or injury to the pudendal nerve, may result in obstructed defecation from descending perineum syndrome several years later. Those few who do not benefit from the simple measures delineated above or require long-term treatment or fail to respond to potent laxatives should undergo further investigation (Fig. 55-4). Novel agents that induce secretion (e.g., lubiprostone, a chloride channel activator, or linaclotide, a guanylate cyclase C agonist that activates chloride secretion) are also available.

INVESTIGATION OF SEVERE CONSTIPATION

A small minority (probably <5%) of patients have severe or “intractable” constipation; about 25% have evacuation disorders. These are the patients most likely to require evaluation by gastroenterologists or in referral centers. Further observation of the patient may occasionally reveal a previously unrecognized cause, such as an evacuation disorder, laxative abuse, malingering, or psychological disorder. In these patients, evaluations of the physiologic function of the colon and pelvic floor and of psychological status aid in the rational choice of treatment. Even among these highly selected patients with severe constipation, a cause can be identified in only about one-third of tertiary referral patients, with the others being diagnosed with normal transit constipation.

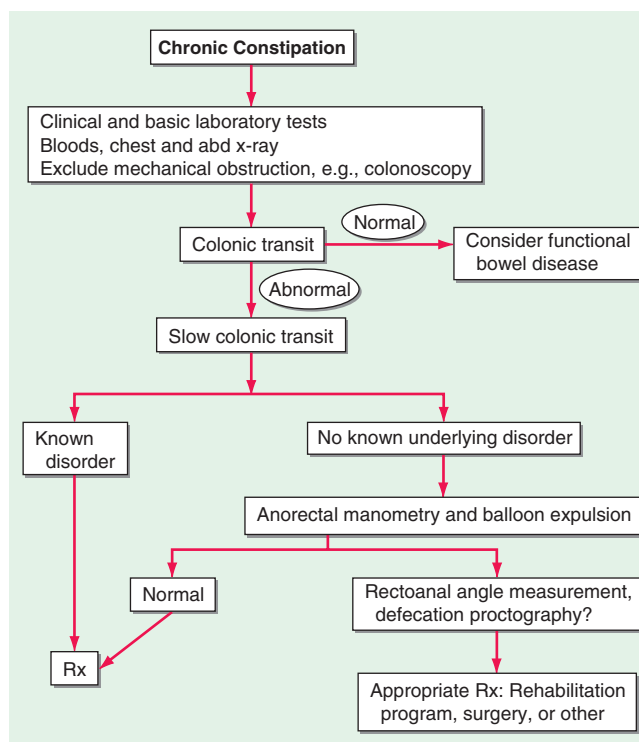


FIGURE 55-4 Algorithm for the management of constipation. abd, abdominal.

Measurement of Colonic Transit Radiopaque marker transit tests are easy, repeatable, generally safe, inexpensive, reliable, and highly applicable in evaluating constipated patients in clinical practice. Several validated methods are very simple. For example, radiopaque markers are ingested; an abdominal flat film taken 5 days later should indicate passage of 80% of the markers out of the colon without the use of laxatives or enemas. This test does not provide useful information about the transit profile of the stomach and small bowel.

Radioscintigraphy with a delayed-release capsule containing radio-labeled particles has been used to noninvasively characterize normal, accelerated, or delayed colonic function over 24–48 h with low radiation exposure. This approach simultaneously assesses gastric, small bowel (which may be important in ~20% of patients with delayed colonic transit because they reflect a more generalized GI motility disorder), and colonic transit. The disadvantages are the greater cost and the need for specific materials prepared in a nuclear medicine laboratory.

Anorectal and Pelvic Floor Tests Pelvic floor dysfunction is suggested by the inability to evacuate the rectum, a feeling of persistent rectal fullness, rectal pain, the need to extract stool from the rectum digitally, application of pressure on the posterior wall of the vagina, support of the perineum during straining, and excessive straining. These significant symptoms should be contrasted with the simple sense of incomplete rectal evacuation, which is common in IBS.

Formal psychological evaluation may identify eating disorders, “control issues,” depression, or post-traumatic stress disorders that may respond to cognitive or other intervention and may be important in restoring quality of life to patients who might present with chronic constipation.

A simple clinical test in the office to document a non-relaxing puborectalis muscle is to have the patient strain to expel the index finger during a digital rectal examination. Motion of the puborectalis posteriorly during straining indicates proper coordination of the pelvic floor muscles. Motion anteriorly with paradoxical contraction during simulated evacuation indicates pelvic floor dysfunction.

Measurement of perineal descent is relatively easy to gauge clinically by placing the patient in the left decubitus position and watching

the perineum to detect inadequate descent (<1.5 cm, a sign of pelvic floor dysfunction) or perineal ballooning during straining relative to bony landmarks (>4 cm, suggesting excessive perineal descent).

A useful overall test of evacuation is the balloon expulsion test. A balloon-tipped urinary catheter is placed and inflated with 50 mL of water. Normally, a patient can expel it while seated on a toilet or in the left lateral decubitus position. In the lateral position, the weight needed to facilitate expulsion of the balloon is determined; normally, expulsion occurs with <200 g added or unaided within 2 min.

Anorectal manometry, when used in the evaluation of patients with severe constipation, may find an excessively high resting (>80 mmHg) or squeeze anal sphincter tone, suggesting anismus (anal sphincter spasm). This test also identifies rare syndromes, such as adult Hirschsprung's disease, by the absence of the rectoanal inhibitory reflex.

Defecography (a dynamic barium enema including lateral views obtained during barium expulsion or a magnetic resonance defecogram) reveals "soft abnormalities" in many patients; the most relevant findings are the measured changes in rectoanal angle, anatomic defects of the rectum such as internal mucosal prolapse, and enteroceles or rectoceles. Surgically remediable conditions are identified in only a few patients. These include severe, whole-thickness intussusception with complete outlet obstruction due to funnel-shaped plugging at the anal canal or an extremely large rectocele that fills preferentially during attempts at defecation instead of expulsion of the barium through the anus. In summary, defecography requires an interested and experienced radiologist, and abnormalities are not pathognomonic for pelvic floor dysfunction. The most common cause of outlet obstruction is failure of the puborectalis muscle to relax; this is not identified by barium defecography, but can be demonstrated by magnetic resonance defecography, which provides more information about the structure and function of the pelvic floor, distal colorectum, and anal sphincters.

Neurologic testing (electromyography) is more helpful in the evaluation of patients with incontinence than of those with symptoms suggesting obstructed defecation. The absence of neurologic signs in the lower extremities suggests that any documented denervation of the puborectalis results from pelvic (e.g., obstetric) injury or from stretching of the pudendal nerve by chronic, long-standing straining. Constipation is common among patients with spinal cord injuries, neurologic diseases such as Parkinson's disease, multiple sclerosis, and diabetic neuropathy.

Spinal-evoked responses during electrical rectal stimulation or stimulation of external anal sphincter contraction by applying magnetic stimulation over the lumbosacral cord identify patients with limited sacral neuropathies with sufficient residual nerve conduction to attempt biofeedback training.

In summary, a balloon expulsion test is an important screening test for anorectal dysfunction. Rarely, an anatomic evaluation of the rectum or anal sphincters and an assessment of pelvic floor relaxation are the tools for evaluating patients in whom obstructed defecation is suspected and is associated with symptoms of rectal mucosal prolapse, pressure of the posterior wall of the vagina to facilitate defecation (suggestive of anterior rectocele), or prior pelvic surgery that may be complicated by enterocele.

TREATMENT CONSTIPATION

After the cause of constipation is characterized, a treatment decision can be made. Slow-transit constipation requires aggressive medical or surgical treatment; anismus or pelvic floor dysfunction usually responds to biofeedback management (Fig. 40-4). The remaining ~60% of patients with constipation has normal colonic transit and can be treated symptomatically. Patients with spinal cord injuries or other neurologic disorders require a dedicated bowel regimen that often includes rectal stimulation, enema therapy, and carefully timed laxative therapy.

Patients with constipation are treated with bulk, osmotic, prokinetic, secretory, and stimulant laxatives including fiber, psyllium, milk of magnesia, lactulose, polyethylene glycol (colonic lavage

solution), lubiprostone, linaclotide, and bisacodyl, or, in some countries, prucalopride, a 5-HT₄ agonist. If a 3-to 6-month trial of medical therapy fails, unassociated with obstructed defecation, the patients should be considered for laparoscopic colectomy with ileorectostomy; however, this should not be undertaken if there is continued evidence of an evacuation disorder or a generalized GI dysmotility. Referral to a specialized center for further tests of colonic motor function is warranted. The decision to resort to surgery is facilitated in the presence of megacolon and megarectum. The complications after surgery include small-bowel obstruction (11%) and fecal soiling, particularly at night during the first postoperative year. Frequency of defecation is 3–8 per day during the first year, dropping to 1–3 per day from the second year after surgery.

Patients who have a combined (evacuation and transit/motility) disorder should pursue pelvic floor retraining (biofeedback and muscle relaxation), psychological counseling, and dietetic advice first. If symptoms are intractable despite biofeedback and optimized medical therapy, colectomy and ileorectostomy could be considered as long as the evacuation disorder is resolved and optimized medical therapy is unsuccessful. In patients with pelvic floor dysfunction alone, biofeedback training has a 70–80% success rate, measured by the acquisition of comfortable stool habits. Attempts to manage pelvic floor dysfunction with operations (internal anal sphincter or puborectalis muscle division) or injections with botulinum toxin have achieved only mediocre success and have been largely abandoned.

56

Involuntary Weight Loss

Russell G. Robertson, J. Larry Jameson

Involuntary weight loss (IWL) is frequently insidious and can have important implications, often serving as a harbinger of serious underlying disease. Clinically important weight loss is defined as the loss of 10 pounds (4.5 kg) or >5% of one's body weight over a period of 6–12 months. IWL is encountered in up to 8% of all adult outpatients and 27% of frail persons age 65 years and older. There is no identifiable cause in up to one-quarter of patients despite extensive investigation. Conversely, up to half of people who claim to have lost weight have no documented evidence of weight loss. People with no known cause of weight loss generally have a better prognosis than do those with known causes, particularly when the source is neoplastic. Weight loss in older persons is associated with a variety of deleterious effects, including hip fracture, pressure ulcers, impaired immune function, and decreased functional status. Not surprisingly, significant weight loss is associated with increased mortality, which can range from 9% to as high as 38% within 1 to 2.5 years in the absence of clinical awareness and attention.

PHYSIOLOGY OF WEIGHT REGULATION WITH AGING

(See also Chaps. 94e and 415e) Among healthy aging people, total body weight peaks in the sixth decade of life and generally remains stable until the ninth decade, after which it gradually falls. In contrast, lean body mass (fat-free mass) begins to decline at a rate of 0.3 kg per year in the third decade, and the rate of decline increases further beginning at age 60 in men and age 65 in women. These changes in lean body mass largely reflect the age-dependent decline in growth hormone secretion and, consequently, circulating levels of insulin-like growth factor type I (IGF-I) that occur with normal aging. Loss of sex steroids, at menopause in women and more gradually with aging in men, also contributes to these changes in body composition. In the healthy elderly, an increase in fat tissue balances the loss in lean body mass until very old age, when loss of both fat and skeletal muscle occurs. Age-dependent changes also occur at the cellular level. Telomeres shorten, and body cell mass—the fat-free portion of cells—declines steadily with aging.

Between ages 20 and 80, mean energy intake is reduced by up to 1200 kcal/d in men and 800 kcal/d in women. Decreased hunger is a reflection of reduced physical activity and loss of lean body mass, producing lower demand for calories and food intake. Several important age-associated physiologic changes also predispose elderly persons to weight loss, such as declining chemosensory function (smell and taste), reduced efficiency of chewing, slowed gastric emptying, and alterations in the neuroendocrine axis, including changes in levels of leptin, cholecystokinin, neuropeptide Y, and other hormones and peptides. These changes are associated with early satiety and a decline in both appetite and the hedonistic appreciation of food. Collectively, they contribute to the “anorexia of aging.”

CAUSES OF INVOLUNTARY WEIGHT LOSS

Most causes of IWL belong to one of four categories: (1) malignant neoplasms, (2) chronic inflammatory or infectious diseases, (3) metabolic disorders (e.g., hyperthyroidism and diabetes), or (4) psychiatric disorders (Table 56-1). Not infrequently, more than one of these causes can be responsible for IWL. In most series, IWL is caused by malignant disease in a quarter of patients and by organic disease in one-third, with the remainder due to psychiatric disease, medications, or uncertain causes.

The most common malignant causes of IWL are gastrointestinal, hepatobiliary, hematologic, lung, breast, genitourinary, ovarian, and prostate. Half of all patients with cancer lose some body weight;

one-third lose more than 5% of their original body weight, and up to 20% of all cancer deaths are caused directly by cachexia (through immobility and/or cardiac/respiratory failure). The greatest incidence of weight loss is seen among patients with solid tumors. Malignancy that reveals itself through significant weight loss usually has a very poor prognosis.

In addition to malignancies, gastrointestinal causes are among the most prominent causes of IWL. Peptic ulcer disease, inflammatory bowel disease, dysmotility syndromes, chronic pancreatitis, celiac disease, constipation, and atrophic gastritis are some of the more common entities. Oral and dental problems are easily overlooked and may manifest with halitosis, poor oral hygiene, xerostomia, inability to chew, reduced masticatory force, nonocclusion, temporomandibular joint syndrome, edentulousness, and pain due to caries or abscesses.

Tuberculosis, fungal diseases, parasites, subacute bacterial endocarditis, and HIV are well-documented causes of IWL. Cardiovascular and pulmonary diseases cause unintentional weight loss through increased metabolic demand and decreased appetite and caloric intake. Uremia produces nausea, anorexia, and vomiting. Connective tissue diseases may increase metabolic demand and disrupt nutritional balance. As the incidence of diabetes mellitus increases with aging, the associated glucosuria can contribute to weight loss. Hyperthyroidism in the elderly may have less prominent sympathetic features and may present as “apathetic hyperthyroidism” or T₃ toxicosis (Chap. 405).

Neurologic injuries such as stroke, quadriplegia, and multiple sclerosis may lead to visceral and autonomic dysfunction that can impair caloric intake. Dysphagia from these neurologic insults is a common mechanism. Functional disability that compromises activities of daily living (ADLs) is a common cause of undernutrition in the elderly. Visual impairment from ophthalmic or central nervous system disorders such as a tremor can limit the ability of people to prepare and eat meals. IWL may be one of the earliest manifestations of Alzheimer’s dementia.

Isolation and depression are significant causes of IWL that may manifest as an inability to care for oneself, including nutritional needs. A cytokine-mediated inflammatory metabolic cascade can be both a cause of and a manifestation of depression. Bereavement can be a cause of IWL and, when present, is more pronounced in men. More intense forms of mental illness such as paranoid disorders may lead to delusions about food and cause weight loss. Alcoholism can be a significant source of weight loss and malnutrition.

Elderly persons living in poverty may have to choose whether to purchase food or use the money for other expenses, including medications. Institutionalization is an independent risk factor, as up to 30–50% of nursing home patients have inadequate food intake.

Medications can cause anorexia, nausea, vomiting, gastrointestinal distress, diarrhea, dry mouth, and changes in taste. This is particularly an issue in the elderly, many of whom take five or more medications.

ASSESSMENT

The four major manifestations of IWL are (1) anorexia (loss of appetite), (2) sarcopenia (loss of muscle mass), (3) cachexia (a syndrome that combines weight loss, loss of muscle and adipose tissue, anorexia, and weakness), and (4) dehydration. The current obesity epidemic adds complexity, as excess adipose tissue can mask the development of sarcopenia and delay awareness of the development of cachexia. If it is not possible to measure weight directly, a change in clothing size, corroboration of weight loss by a relative or friend, and a numeric estimate of weight loss provided by the patient are suggestive of true weight loss.

Initial assessment includes a comprehensive history and physical, a complete blood count, tests of liver enzyme levels, C-reactive protein, erythrocyte sedimentation rate, renal function studies, thyroid function tests, chest radiography, and an abdominal ultrasound (Table 56-2). Age, sex, and risk factor-specific cancer screening tests, such as mammography and colonoscopy, should be performed (Chap. 100). Patients at risk should have HIV testing. All elderly patients with weight loss should undergo screening for dementia and depression by using instruments such as the

TABLE 56-1 CAUSES OF INVOLUNTARY WEIGHT LOSS

Cancer	Medications
Colon	Sedatives
Hepatobiliary	Antibiotics
Hematologic	Nonsteroidal anti-inflammatory drugs
Lung	Serotonin reuptake inhibitors
Breast	Metformin
Genitourinary	Levodopa
Ovarian	Angiotensin-converting enzyme inhibitors
Prostate	Other drugs
Gastrointestinal disorders	Disorders of the mouth and teeth
Malabsorption	Caries
Peptic ulcer	Dysgeusia
Inflammatory bowel disease	
Pancreatitis	Age-related factors
Obstruction/constipation	Physiologic changes
Pernicious anemia	Visual impairment
Endocrine and metabolic	Decreased taste and smell
Hyperthyroidism	Functional disabilities
Diabetes mellitus	Neurologic
Pheochromocytoma	Stroke
Adrenal insufficiency	Parkinson’s disease
Cardiac disorders	Neuromuscular disorders
Chronic ischemia	Dementia
Chronic congestive heart failure	Social
Respiratory disorders	Isolation
Emphysema	Economic hardship
Chronic obstructive pulmonary disease	Psychiatric and behavioral
Renal insufficiency	Depression
Rheumatologic disease	Anxiety
Infections	Paranoia
HIV	Bereavement
Tuberculosis	Alcoholism
Parasitic infection	Eating disorders
Subacute bacterial endocarditis	Increased activity or exercise
	Idiopathic

TABLE 56-2 ASSESSMENT AND TESTING FOR INVOLUNTARY WEIGHT LOSS

Indications	Laboratory
5% weight loss in 30 d	Complete blood count
10% weight loss in 180 d	Comprehensive electrolyte and metabolic panel, including liver and renal function tests
Body mass index <21	Thyroid function tests
25% of food left uneaten after 7 d	Erythrocyte sedimentation rate
Change in fit of clothing	C-reactive protein
Change in appetite, smell, or taste	Ferritin
Abdominal pain, nausea, vomiting, diarrhea, constipation, dysphagia	HIV testing, if indicated
Assessment	Radiology
Complete physical exam, including dental evaluation	Chest x-ray
Medication review	Abdominal ultrasound
Recommended cancer screening	
Mini-Mental State Examination ^a	
Mini-Nutritional Assessment ^a	
Nutrition Screening Initiative ^a	
Simplified Nutritional Assessment Questionnaire ^a	
Observation of eating ^a	
Activities of daily living ^a	
Instrumental activities of daily living ^a	

^aMay be more specific to assess weight loss in the elderly.

Mini-Mental State Examination and the Geriatric Depression Scale, respectively (Chap. 11). The Mini Nutritional Assessment (www.mna-elderly.com) and the Nutrition Screening Initiative (<http://www.ncbi.nlm.nih.gov/pmc/articles/PMC1694757/>) are also available for the nutritional assessment of elderly patients. Almost all patients with a malignancy and >90% of those with other organic diseases have at least one laboratory abnormality. In patients presenting with substantial IWL, major organic and malignant diseases are unlikely when a baseline evaluation is completely normal. Careful follow-up rather than undirected testing is advised since the prognosis of weight loss of undetermined cause is generally favorable.

TREATMENT UNINTENTIONAL WEIGHT LOSS

The first priority in managing weight loss is to identify and treat the underlying causes systematically. Treatment of underlying metabolic, psychiatric, infectious, or other systemic disorders may be sufficient to restore weight and functional status gradually. Medications that cause nausea or anorexia should be withdrawn or changed, if possible. For those with unexplained IWL, oral nutritional supplements such as high-energy drinks sometimes reverse weight loss. Advising patients to consume supplements between meals rather than with a meal may help minimize appetite suppression and facilitate increased overall intake. Orexigenic, anabolic, and anticytokine agents are under investigation. In selected patients, the antidepressant mirtazapine results in a significant increase in body weight, body fat mass, and leptin concentration. Patients with wasting conditions who can comply with an appropriate exercise program gain muscle protein mass, strength, and endurance and may be more capable of performing ADLs.

57

Gastrointestinal Bleeding

Loren Laine

Gastrointestinal bleeding (GIB) accounts for ~150 hospitalizations per 100,000 population annually in the United States, with upper GIB (UGIB) ~1.5–2 times more common than lower GIB (LGIB). The incidence of GIB has decreased in recent decades, primarily due to a reduction in UGIB, and the mortality has also decreased to <5%. Patients today rarely die from exsanguination, but rather die due to decompensation of other underlying illnesses.

GIB presents as either overt or occult bleeding. *Overt GIB* is manifested by *hematemesis*, vomitus of red blood or “coffee-ground” material; *melenas*, black, tarry, foul-smelling stool; and/or *hematochezia*, passage of bright red or maroon blood from the rectum. *Occult GIB* may be identified in the absence of overt bleeding when patients present with *symptoms of blood loss or anemia* such as lightheadedness, syncope, angina, or dyspnea; or when routine diagnostic evaluation reveals iron deficiency anemia or a positive fecal occult blood test. GIB is also categorized by the site of bleeding as UGIB, LGIB, or obscure GIB if the source is unclear.

SOURCES OF GASTROINTESTINAL BLEEDING

Upper Gastrointestinal Sources of Bleeding (Table 57-1) Peptic ulcers are the most common cause of UGIB, accounting for ~50% of cases. Mallory-Weiss tears account for ~5–10% of cases. The proportion of patients bleeding from varices varies widely from ~5–40%, depending on the population. Hemorrhagic or erosive gastropathy (e.g., due to nonsteroidal anti-inflammatory drugs [NSAIDs] or alcohol) and erosive esophagitis often cause mild UGIB, but major bleeding is rare.

PEPTIC ULCERS Characteristics of an ulcer at endoscopy provide important prognostic information. One-third of patients with active bleeding or a nonbleeding visible vessel have further bleeding that requires urgent surgery if they are treated conservatively. These patients benefit from endoscopic therapy with bipolar electrocoagulation, heater probe, injection therapy (e.g., absolute alcohol, 1:10,000 epinephrine), and/or clips with reductions in bleeding, hospital stay, mortality, and costs. In contrast, patients with clean-based ulcers have rates of recurrent bleeding approaching zero. If stable with no other reason for hospitalization, such patients may be discharged home after endoscopy. Patients without clean-based ulcers usually remain in the hospital for 3 days because most episodes of recurrent bleeding occur within 3 days.

Randomized controlled trials document that high-dose, constant-infusion IV proton pump inhibitor (PPI) (80-mg bolus and 8-mg/h infusion), designed to sustain intragastric pH >6 and enhance clot stability, decreases further bleeding and mortality in patients with high-risk ulcers (active bleeding, nonbleeding visible vessel, adherent clot) when given after endoscopic therapy. Patients with lower-risk findings (flat pigmented spot or clean base) do not require endoscopic

TABLE 57-1 SOURCES OF BLEEDING IN PATIENTS HOSPITALIZED FOR UPPER GASTROINTESTINAL BLEEDING

Sources of Bleeding	Proportion of Patients, %
Ulcers	31–67
Varices	6–39
Mallory-Weiss tears	2–8
Gastroduodenal erosions	2–18
Erosive esophagitis	1–13
Neoplasm	2–8
Vascular ectasias	0–6
No source identified	5–14

Source: Data on hospitalizations from year 2000 onward from Am J Gastroenterol 98:1494, 2003; Gastrointest Endosc 57:AB147, 2003; 60:875, 2004; Eur J Gastroenterol Hepatol 16:177, 2004; 17:641, 2005; J Clin Gastroenterol 42:128, 2008; World J Gastroenterol 14:5046, 2008; Dig Dis Sci 54:333, 2009; Gut 60:1327, 2011; Endoscopy 44:998, 2012; J Clin Gastroenterol 48:113, 2014.

therapy and receive standard doses of oral PPI. Approximately one-third of patients with bleeding ulcers will rebleed within the next 1–2 years if no preventive strategies are employed. Prevention of recurrent bleeding focuses on the three main factors in ulcer pathogenesis, *Helicobacter pylori*, NSAIDs, and acid. Eradication of *H. pylori* in patients with bleeding ulcers decreases rates of rebleeding to <5%. If a bleeding ulcer develops in a patient taking NSAIDs, the NSAIDs should be discontinued. If NSAIDs must be given, a cyclooxygenase 2 (COX-2) selective inhibitor (coxib) plus a PPI should be used. PPI co-therapy alone or a coxib alone is associated with an annual rebleeding rate of ~10% in patients with a recent bleeding ulcer, whereas the combination of a coxib and PPI provides a further significant decrease in recurrent ulcer bleeding. Patients with established cardiovascular disease who develop bleeding ulcers while taking low-dose aspirin should restart aspirin as soon as possible after their bleeding episode (1–7 days). A randomized trial showed that failure to restart aspirin was associated with no significant difference in rebleeding (5% vs. 10% at 30 days) but a significant increase in mortality at 30 days (9% vs. 1%) and 8 weeks (13% vs. 1%) compared with immediate reinstitution of aspirin. Patients with bleeding ulcers unrelated to *H. pylori* or NSAIDs should remain on PPI therapy indefinitely. **Peptic ulcers are discussed in Chap. 348.**

MALLORY-WEISS TEARS The classic history is vomiting, retching, or coughing preceding hematemesis, especially in an alcoholic patient. Bleeding from these tears, which are usually on the gastric side of the gastroesophageal junction, stops spontaneously in 80–90% of patients and recurs in only 0–10%. Endoscopic therapy is indicated for actively bleeding Mallory-Weiss tears. Angiographic therapy with embolization and operative therapy with oversewing of the tear are rarely required. **Mallory-Weiss tears are discussed in Chap. 347.**

ESOPHAGEAL VARICES Patients with variceal hemorrhage have poorer outcomes than patients with other sources of UGIB. Urgent endoscopy within 12 h is recommended in cirrhotics with UGIB, and if esophageal varices are present, endoscopic ligation is performed and an IV vasoactive medication (e.g., octreotide 50 µg bolus and 50 µg/h infusion) is given for 2–5 days. Combination endoscopic and medical therapy appears to be superior to either therapy alone in decreasing rebleeding. In patients with advanced liver disease (e.g., Child-Pugh class C with score 10–13), a transjugular intrahepatic portosystemic shunt (TIPS) should be strongly considered within the first 1–2 days of hospitalization because randomized trials show significant decreases in rebleeding and mortality compared with standard endoscopic and medical therapy. Over the long term, treatment with nonselective beta blockers plus endoscopic ligation is recommended because the combination of endoscopic and medical therapy is more effective than either alone in reduction of recurrent esophageal variceal bleeding.

In patients who have persistent or recurrent bleeding despite endoscopic and medical therapy, TIPS is recommended. Decompressive surgery (e.g., distal splenorenal shunt) may be considered instead of TIPS in patients with well-compensated cirrhosis.

Portal hypertension is also responsible for bleeding from gastric varices, varices in the small and large intestine, and portal hypertensive gastropathy and enterocolopathy. Bleeding gastric varices due to cirrhosis are treated with endoscopic injection of tissue adhesive (e.g., *n*-butyl cyanoacrylate), if available; if not, TIPS is performed.

HEMORRHAGIC AND EROSIVE GASTROPATHY (“GASTRITIS”) Hemorrhagic and erosive gastropathy, often labeled gastritis, refers to endoscopically visualized subepithelial hemorrhages and erosions. These are mucosal lesions and do not cause major bleeding due to the absence of arteries and veins in the mucosa. Erosions develop in various clinical settings, the most important of which are NSAID use, alcohol intake, and stress. Half of patients who chronically ingest NSAIDs have erosions, whereas up to 20% of actively drinking alcoholic patients with symptoms of UGIB have evidence of subepithelial hemorrhages or erosions.

Stress-related gastric mucosal injury occurs only in extremely sick patients, such as those who have experienced serious trauma, major

surgery, burns covering more than one-third of the body surface area, major intracranial disease, or severe medical illness (i.e., ventilator dependence, coagulopathy). Severe bleeding should not develop unless ulceration occurs. The mortality rate in these patients is quite high because of their serious underlying illnesses.

The incidence of bleeding from stress-related gastric mucosal injury has decreased dramatically in recent years, most likely due to better care of critically ill patients. Pharmacologic prophylaxis for bleeding may be considered in the high-risk patients mentioned above. Meta-analyses of randomized trials indicate that PPIs are more effective than H₂ receptor antagonists in reduction of overt and clinically important UGIB without differences in mortality or nosocomial pneumonia.

OTHER CAUSES Other less frequent causes of UGIB include erosive duodenitis, neoplasms, aortoenteric fistulas, vascular lesions (including hereditary hemorrhagic telangiectasias [Osler-Weber-Rendu] and gastric antral vascular ectasia [“watermelon stomach”]), Dieulafoy’s lesion (in which an aberrant vessel in the mucosa bleeds from a pinpoint mucosal defect), prolapse gastropathy (prolapse of proximal stomach into esophagus with retching, especially in alcoholics), and hemobilia or hemosuccus pancreaticus (bleeding from the bile duct or pancreatic duct).

Small-Intestinal Sources of Bleeding Small-intestinal sources of bleeding (bleeding from sites beyond the reach of the standard upper endoscope) are often difficult to diagnose and are responsible for the majority of cases of obscure GIB. Fortunately, small-intestinal bleeding is uncommon. The most common causes in adults are vascular ectasias, tumors (e.g., GI stromal tumor, carcinoid, adenocarcinoma, lymphoma, metastases), and NSAID-induced erosions and ulcers. Other less common causes in adults include Crohn’s disease, infection, ischemia, vasculitis, small-bowel varices, diverticula, Meckel’s diverticulum, duplication cysts, and intussusception.

Meckel’s diverticulum is the most common cause of significant LGIB in children, decreasing in frequency as a cause of bleeding with age. In adults <40–50 years, small-bowel tumors often account for obscure GIB; in patients >50–60 years, vascular ectasias and NSAID-induced lesions are more commonly responsible.

Vascular ectasias should be treated with endoscopic therapy if possible. Although estrogen/progesterone compounds have been used for vascular ectasias, a large double-blind trial found no benefit in prevention of recurrent bleeding. Octreotide is also used, based on case series but no randomized trials. A randomized trial reported significant benefit of thalidomide and awaits further confirmation. Other isolated lesions, such as tumors, are generally treated with surgical resection.

Colonic Sources of Bleeding Hemorrhoids are probably the most common cause of LGIB; anal fissures also cause minor bleeding and pain. If these local anal processes, which rarely require hospitalization, are excluded, the most common causes of LGIB in adults are diverticula, vascular ectasias (especially in the proximal colon of patients >70 years), neoplasms (primarily adenocarcinoma), colitis (ischemic, infectious, idiopathic inflammatory bowel disease), and postpolypectomy bleeding. Less common causes include NSAID-induced ulcers or colitis, radiation proctopathy, solitary rectal ulcer syndrome, trauma, varices (most commonly rectal), lymphoid nodular hyperplasia, vasculitis, and aortocolic fistulas. In children and adolescents, the most common colonic causes of significant GIB are inflammatory bowel disease and juvenile polyps.

Diverticular bleeding is abrupt in onset, usually painless, sometimes massive, and often from the right colon; chronic or occult bleeding is not characteristic. Clinical reports suggest that bleeding colonic diverticula stop bleeding spontaneously in ~80% of patients and, on long-term follow-up, rebleed in ~15–25% of patients. Case series suggest endoscopic therapy may decrease recurrent bleeding in the uncommon case when colonoscopy identifies the specific bleeding diverticulum. When diverticular bleeding is found at angiography, transcatheter arterial embolization by superselective technique stops bleeding in a majority of patients. If bleeding persists or recurs, segmental surgical resection is indicated.

Bleeding from right colonic vascular ectasias in the elderly may be overt or occult; it tends to be chronic and only occasionally is hemodynamically significant. Endoscopic hemostatic therapy may be useful in the treatment of vascular ectasias, as well as discrete bleeding ulcers and postpolypectomy bleeding. Surgical therapy is generally required for major, persistent, or recurrent bleeding from the wide variety of colonic sources of GIB that cannot be treated medically, angiographically, or endoscopically.

APPROACH TO THE PATIENT: Gastrointestinal Bleeding

INITIAL ASSESSMENT

Measurement of the heart rate and blood pressure is the best way to initially assess a patient with GIB. Clinically significant bleeding leads to postural changes in heart rate or blood pressure, tachycardia, and, finally, recumbent hypotension. In contrast, the hemoglobin does not fall immediately with acute GIB, due to proportionate reductions in plasma and red cell volumes (i.e., “people bleed whole blood”). Thus, hemoglobin may be normal or only minimally decreased at the initial presentation of a severe bleeding episode. As extravascular fluid enters the vascular space to restore volume, the hemoglobin falls, but this process may take up to 72 h. Transfusion is recommended when the hemoglobin drops below 7 g/dL, based on a large randomized trial showing this restrictive transfusion strategy decreases rebleeding and death in acute UGIB compared with a transfusion threshold of 9 g/dL. Patients with slow, chronic GIB may have very low hemoglobin values despite normal blood pressure and heart rate. With the development of iron-deficiency anemia, the mean corpuscular volume will be low and red blood cell distribution width will increase.

DIFFERENTIATION OF UGIB FROM LGIB

Hematemesis indicates an upper GI source of bleeding (above the ligament of Treitz). Melena indicates blood has been present in the GI tract for at least 14 h, and as long as 3–5 days. The more proximal the bleeding site, the more likely melena will occur. Hematochezia usually represents a lower GI source of bleeding, although an upper GI lesion may bleed so briskly that blood transits the bowel before melena develops. When hematochezia is the presenting symptom of UGIB, it is associated with hemodynamic instability and dropping hemoglobin. Bleeding lesions of the small bowel may present as melena or hematochezia. Other clues to UGIB include hyperactive bowel sounds and an elevated blood urea nitrogen (due to volume depletion and blood proteins absorbed in the small intestine).

A nonbloody nasogastric aspirate may be seen in up to ~18% of patients with UGIB, usually from a duodenal source. Even a bile-stained appearance does not exclude a bleeding postpyloric lesion because reports of bile in the aspirate are incorrect in ~50% of cases. Testing of aspirates that are not grossly bloody for occult blood is not useful.

EVALUATION AND MANAGEMENT OF UGIB (FIG. 57-1)

At presentation, patients are generally stratified as higher or lower risk for further bleeding and death. Baseline characteristics predictive of rebleeding and death include hemodynamic compromise (tachycardia or hypotension), increasing age, and comorbidities. PPI infusion may be considered at presentation: it decreases high-risk ulcer stigmata (e.g., active bleeding) and need for endoscopic therapy but does not improve clinical outcomes such as further bleeding, surgery, or death. Treatment to improve endoscopic visualization with the promotility agent erythromycin, 250 mg intravenously ~30 min before endoscopy, also may be considered: it provides a small but significant increase in diagnostic yield and decrease in second endoscopies but is not documented to decrease further bleeding or death. Cirrhotic patients presenting with UGIB should be placed on antibiotics (e.g., quinolone, ceftriaxone) and started on a vasoactive medication (octreotide, terlipressin, somatostatin, vapreotide) upon presentation, even before endoscopy. Antibiotics decrease bacterial infections, rebleeding, and mortality in this population, and vasoactive medications appear to improve control of bleeding in the first 12 h after presentation.

Upper endoscopy should be performed within 24 h in most patients with UGIB. Patients at higher risk (e.g., hemodynamic instability, cirrhosis) may benefit from more urgent endoscopy within 12 h. Early endoscopy is also beneficial in low-risk patients for management decisions. Patients with major bleeding and high-risk endoscopic findings (e.g., varices, ulcers with active bleeding or a visible vessel) benefit from endoscopic hemostatic therapy, whereas patients with low-risk lesions (e.g., clean-based ulcers, nonbleeding Mallory-Weiss tears, erosive or hemorrhagic gastropathy) who have stable vital signs and hemoglobin and no other medical problems can be discharged home.

EVALUATION AND MANAGEMENT OF LGIB (FIG. 57-2)

Patients with hematochezia and hemodynamic instability should have upper endoscopy to rule out an upper GI source before evaluation of the lower GI tract.

Colonoscopy after an oral lavage solution is the procedure of choice in most patients admitted with LGIB unless bleeding is too massive,

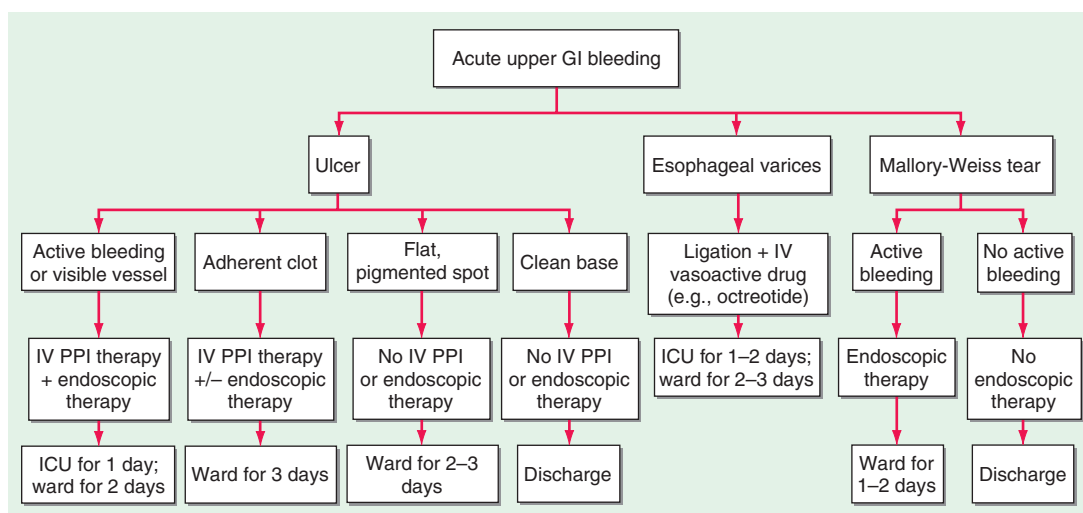


FIGURE 57-1 Suggested algorithm for patients with acute upper gastrointestinal (GI) bleeding. Recommendations on level of care and time of discharge assume patient is stabilized without further bleeding or other concomitant medical problems. ICU, intensive care unit; PPI, proton pump inhibitor.

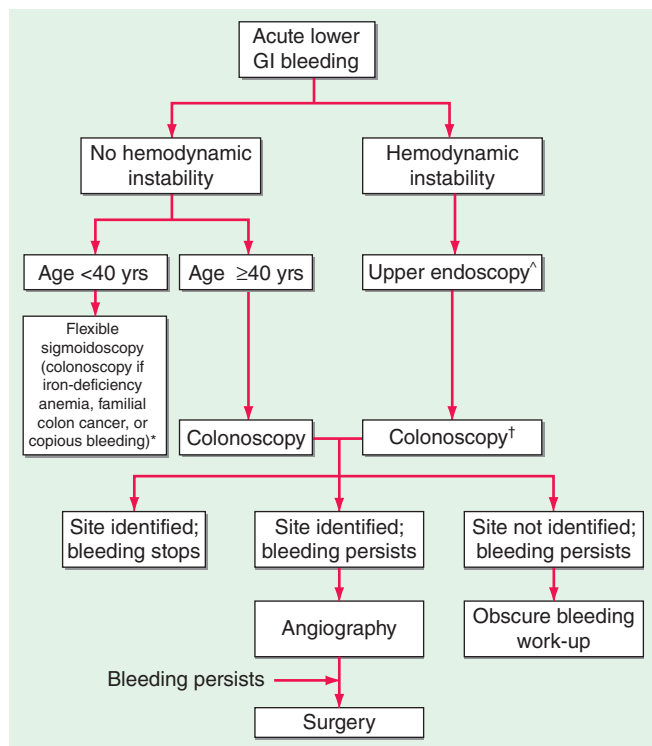


FIGURE 57-2 Suggested algorithm for patients with acute lower gastrointestinal (GI) bleeding. *Some suggest colonoscopy for any degree of rectal bleeding in patients <40 years as well. †If upper GI endoscopy reveals definite source, no further evaluation is needed. ‡If massive bleeding does not allow time for colonic lavage, proceed to angiography.

in which case angiography is recommended. Sigmoidoscopy is used primarily in patients <40 years old with minor bleeding. In patients with no source identified on colonoscopy, imaging studies may be employed. ^{99m}Tc -labeled red cell scan allows repeated imaging for up to 24 h and may identify the general location of bleeding. However, radionuclide scans should be interpreted with caution because results, especially from later images, are highly variable. Multidetector computed tomography (CT) “angiography” is an increasingly used technique that is likely superior to nuclear scintigraphy. In active LGIB, angiography can detect the site of bleeding (extravasation of contrast into the gut) and permits treatment with embolization. Even after bleeding has stopped, angiography may identify lesions with abnormal vasculature, such as vascular ectasias or tumors.

EVALUATION AND MANAGEMENT OF OBSCURE GIB

Obscure GIB is defined as persistent or recurrent bleeding for which no source has been identified by routine endoscopic and contrast x-ray studies; it may be overt (melena, hematochezia) or occult (iron-deficiency anemia). Current guidelines suggest angiography as the initial test for massive obscure bleeding, and video capsule endoscopy, which allows examination of the entire small intestine, for all others. Push enteroscopy, usually performed with a pediatric colonoscope, to inspect the entire duodenum and proximal jejunum also may be considered as an initial evaluation. A systematic review of 14 trials comparing push enteroscopy to capsule revealed “clinically significant findings” in 26% and 56% of patients, respectively. However, in contrast to enteroscopy, lack of control of the capsule prevents its manipulation and full visualization of the intestine; in addition, tissue cannot be sampled and therapy cannot be applied.

If capsule endoscopy is positive, management is dictated by the finding. If capsule endoscopy is negative, current recommendations

suggest patients may either be observed or, if their clinical course mandates (e.g., recurrent bleeding, need for transfusions or hospitalization), undergo further testing. “Deep” enteroscopy (e.g., double-balloon, single-balloon, and spiral enteroscopy) is commonly the next test undertaken in patients with clinically important obscure GIB because it allows the endoscopist to examine, obtain specimens from, and provide therapy to much or all of the small intestine. CT and magnetic resonance enterography also are used to examine the small intestine. Other imaging techniques sometimes used in evaluation of obscure GIB include ^{99m}Tc -labeled red blood cell scintigraphy, multidetector CT “angiography,” angiography, and ^{99m}Tc -pertechnetate scintigraphy for Meckel’s diverticulum (especially in young patients). If all tests are unrevealing, intraoperative endoscopy is indicated in patients with severe recurrent or persistent bleeding requiring repeated transfusions.

POSITIVE FECAL OCCULT BLOOD TEST

Fecal occult blood testing is recommended only for colorectal cancer screening and may be used beginning at age 50 in average-risk adults and beginning at age 40 in adults with a first-degree relative with colorectal neoplasm at ≥ 60 years or two second-degree relatives with colorectal cancer. A positive test necessitates colonoscopy. If evaluation of the colon is negative, further workup is not recommended unless iron-deficiency anemia or GI symptoms are present.

58

Jaundice

Savio John, Daniel S. Pratt

Jaundice, or icterus, is a yellowish discoloration of tissue resulting from the deposition of bilirubin. Tissue deposition of bilirubin occurs only in the presence of serum hyperbilirubinemia and is a sign of either liver disease or, less often, a hemolytic disorder. The degree of serum bilirubin elevation can be estimated by physical examination. Slight increases in serum bilirubin level are best detected by examining the sclerae, which have a particular affinity for bilirubin due to their high elastin content. The presence of scleral icterus indicates a serum bilirubin level of at least $51 \mu\text{mol/L}$ (3 mg/dL). The ability to detect scleral icterus is made more difficult if the examining room has fluorescent lighting. If the examiner suspects scleral icterus, a second site to examine is underneath the tongue. As serum bilirubin levels rise, the skin will eventually become yellow in light-skinned patients and even green if the process is long-standing; the green color is produced by oxidation of bilirubin to biliverdin.

The differential diagnosis for yellowing of the skin is limited. In addition to jaundice, it includes carotenoderma, the use of the drug quinacrine, and excessive exposure to phenols. Carotenoderma is the yellow color imparted to the skin of healthy individuals who ingest excessive amounts of vegetables and fruits that contain carotene, such as carrots, leafy vegetables, squash, peaches, and oranges. In jaundice the yellow coloration of the skin is uniformly distributed over the body, whereas in carotenoderma the pigment is concentrated on the palms, soles, forehead, and nasolabial folds. Carotenoderma can be distinguished from jaundice by the sparing of the sclerae. Quinacrine causes a yellow discoloration of the skin in 4–37% of patients treated with it.

Another sensitive indicator of increased serum bilirubin is darkening of the urine, which is due to the renal excretion of conjugated bilirubin. Patients often describe their urine as tea- or cola-colored. Bilirubinuria indicates an elevation of the direct serum bilirubin fraction and, therefore, the presence of liver disease.

Serum bilirubin levels increase when an imbalance exists between bilirubin production and clearance. A logical evaluation of the patient who is jaundiced requires an understanding of bilirubin production and metabolism.

PRODUCTION AND METABOLISM OF BILIRUBIN

(See also Chap. 359) Bilirubin, a tetrapyrrole pigment, is a breakdown product of heme (ferroprotoporphyrin IX). About 70–80% of the 250–300 mg of bilirubin produced each day is derived from the breakdown of hemoglobin in senescent red blood cells. The remainder comes from prematurely destroyed erythroid cells in bone marrow and from the turnover of hemoproteins such as myoglobin and cytochromes found in tissues throughout the body.

The formation of bilirubin occurs in reticuloendothelial cells, primarily in the spleen and liver. The first reaction, catalyzed by the microsomal enzyme heme oxygenase, oxidatively cleaves the α bridge of the porphyrin group and opens the heme ring. The end products of this reaction are biliverdin, carbon monoxide, and iron. The second reaction, catalyzed by the cytosolic enzyme biliverdin reductase, reduces the central methylene bridge of biliverdin and converts it to bilirubin. Bilirubin formed in the reticuloendothelial cells is virtually insoluble in water due to tight internal hydrogen bonding between the water-soluble moieties of bilirubin—i.e., the bonding of the propionic acid carboxyl groups of one dipyrrolic half of the molecule with the imino and lactam groups of the opposite half. This configuration blocks solvent access to the polar residues of bilirubin and places the hydrophobic residues on the outside. To be transported in blood, bilirubin must be solubilized. Solubilization is accomplished by the reversible, noncovalent binding of bilirubin to albumin. Unconjugated bilirubin bound to albumin is transported to the liver. There, the bilirubin—but not the albumin—is taken up by hepatocytes via a process that at least partly involves carrier-mediated membrane transport. No specific bilirubin transporter has yet been identified (Chap. 359, Fig. 359-1).

After entering the hepatocyte, unconjugated bilirubin is bound in the cytosol to a number of proteins including proteins in the glutathione-S-transferase superfamily. These proteins serve both to reduce efflux of bilirubin back into the serum and to present the bilirubin for conjugation. In the endoplasmic reticulum, bilirubin is solubilized by conjugation to glucuronic acid, a process that disrupts the internal hydrogen bonds and yields bilirubin monoglucuronide and diglucuronide. The conjugation of glucuronic acid to bilirubin is catalyzed by bilirubin uridine diphosphate-glucuronosyl transferase (UDPGT). The now-hydrophilic bilirubin conjugates diffuse from the endoplasmic reticulum to the canalicular membrane, where bilirubin monoglucuronide and diglucuronide are actively transported into canalicular bile by an energy-dependent mechanism involving the multidrug resistance-associated protein 2 (MRP2).

The conjugated bilirubin excreted into bile drains into the duodenum and passes unchanged through the proximal small bowel. Conjugated bilirubin is not taken up by the intestinal mucosa. When the conjugated bilirubin reaches the distal ileum and colon, it is hydrolyzed to unconjugated bilirubin by bacterial β -glucuronidases. The unconjugated bilirubin is reduced by normal gut bacteria to form a group of colorless tetrapyrroles called *urobilinogens*. About 80–90% of these products are excreted in feces, either unchanged or oxidized to orange derivatives called *urobilins*. The remaining 10–20% of the urobilinogens are passively absorbed, enter the portal venous blood, and are re-excreted by the liver. A small fraction (usually <3 mg/dL) escapes hepatic uptake, filters across the renal glomerulus, and is excreted in urine.

MEASUREMENT OF SERUM BILIRUBIN

The terms *direct* and *indirect* bilirubin—i.e., conjugated and unconjugated bilirubin, respectively—are based on the original van den Bergh reaction. This assay, or a variation of it, is still used in most clinical chemistry laboratories to determine the serum bilirubin level. In this assay, bilirubin is exposed to diazotized sulfanilic acid and splits into two relatively stable dipyrromethene azopigments that absorb

maximally at 540 nm, allowing photometric analysis. The direct fraction is that which reacts with diazotized sulfanilic acid in the absence of an accelerator substance such as alcohol. The direct fraction provides an approximation of the conjugated bilirubin level in serum. The *total* serum bilirubin is the amount that reacts after the addition of alcohol. The indirect fraction is the difference between the total and the direct bilirubin levels and provides an estimate of the unconjugated bilirubin in serum.

With the van den Bergh method, the normal serum bilirubin concentration usually is 17 $\mu\text{mol/L}$ (<1 mg/dL). Up to 30%, or 5.1 $\mu\text{mol/L}$ (0.3 mg/dL), of the total may be direct-reacting (conjugated) bilirubin. Total serum bilirubin concentrations are between 3.4 and 15.4 $\mu\text{mol/L}$ (0.2 and 0.9 mg/dL) in 95% of a normal population.

Several new techniques, although less convenient to perform, have added considerably to our understanding of bilirubin metabolism. First, studies using these methods demonstrate that, in normal persons or those with Gilbert's syndrome, almost 100% of the serum bilirubin is unconjugated; <3% is monoconjugated bilirubin. Second, in jaundiced patients with hepatobiliary disease, the total serum bilirubin concentration measured by these new, more accurate methods is lower than the values found with diazo methods. This finding suggests that there are diazo-positive compounds distinct from bilirubin in the serum of patients with hepatobiliary disease. Third, these studies indicate that, in jaundiced patients with hepatobiliary disease, monoglucuronides of bilirubin predominate over diglucuronides. Fourth, part of the direct-reacting bilirubin fraction includes conjugated bilirubin that is covalently linked to albumin. This albumin-linked bilirubin fraction (*delta fraction*, or *biliprotein*) represents an important fraction of total serum bilirubin in patients with cholestasis and hepatobiliary disorders. The delta fraction (delta bilirubin) is formed in serum when hepatic excretion of bilirubin glucuronides is impaired and the glucuronides accumulate in serum. By virtue of its tight binding to albumin, the clearance rate of delta bilirubin from serum approximates the half-life of albumin (12–14 days) rather than the short half-life of bilirubin (about 4 h).

The prolonged half-life of albumin-bound conjugated bilirubin accounts for two previously unexplained enigmas in jaundiced patients with liver disease: (1) that some patients with conjugated hyperbilirubinemia do not exhibit bilirubinuria during the recovery phase of their disease because the bilirubin is covalently bound to albumin and therefore not filtered by the renal glomeruli, and (2) that the elevated serum bilirubin level declines more slowly than expected in some patients who otherwise appear to be recovering satisfactorily. Late in the recovery phase of hepatobiliary disorders, all the conjugated bilirubin may be in the albumin-linked form.

MEASUREMENT OF URINE BILIRUBIN

Unconjugated bilirubin is always bound to albumin in the serum, is not filtered by the kidney, and is not found in the urine. Conjugated bilirubin is filtered at the glomerulus, and the majority is reabsorbed by the proximal tubules; a small fraction is excreted in the urine. Any bilirubin found in the urine is conjugated bilirubin. The presence of bilirubinuria implies the presence of liver disease. A urine dipstick test (Ictotest) gives the same information as fractionation of the serum bilirubin and is very accurate. A false-negative result is possible in patients with prolonged cholestasis due to the predominance of delta bilirubin, which is covalently bound to albumin and therefore not filtered by the renal glomeruli.

APPROACH TO THE PATIENT:

Jaundice

The goal of this chapter is not to provide an encyclopedic review of all of the conditions that can cause jaundice. Rather, the chapter is intended to offer a framework that helps a physician to evaluate the patient with jaundice in a logical way (Fig. 58-1).

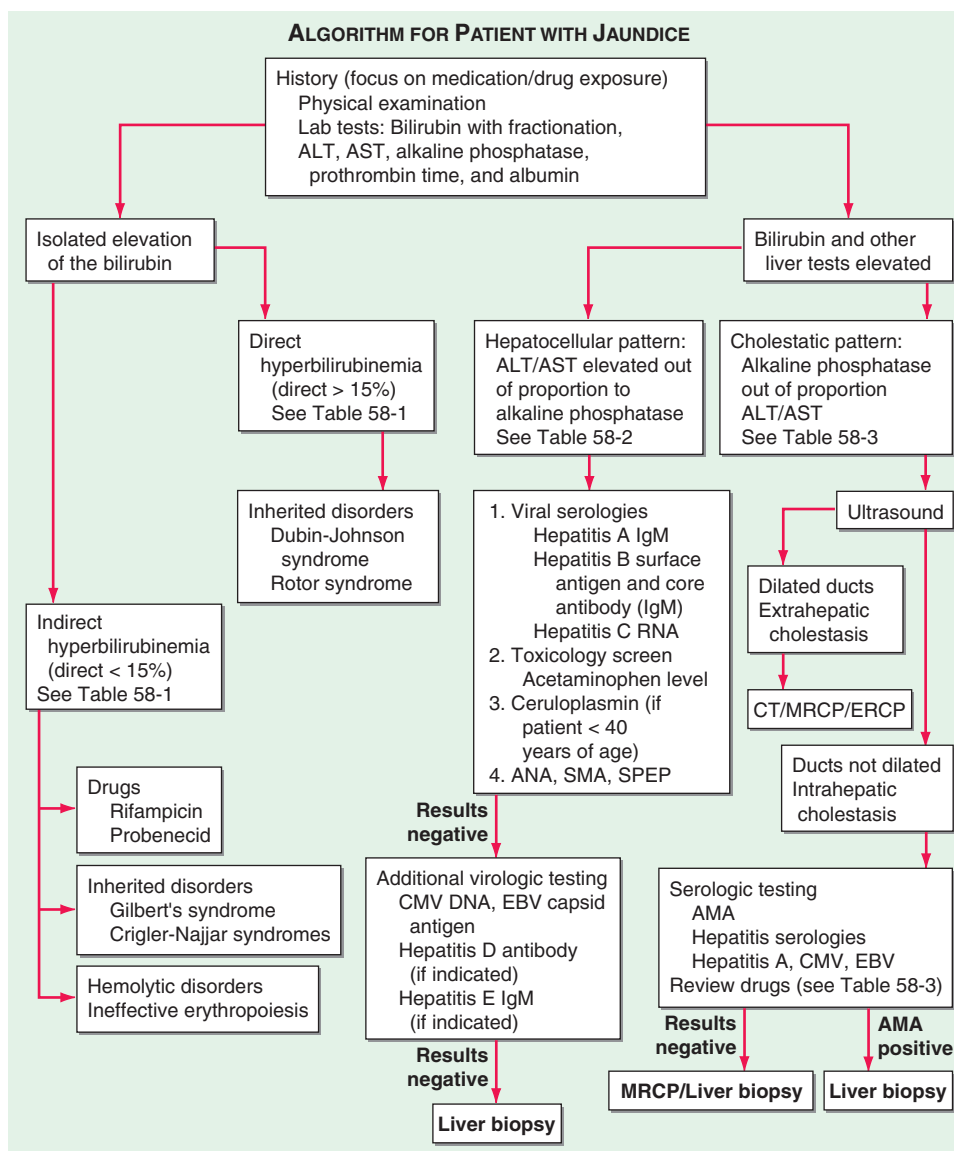


FIGURE 58-1 Evaluation of the patient with jaundice. ALT, alanine aminotransferase; AMA, antimitochondrial antibody; ANA, antinuclear antibody; AST, aspartate aminotransferase; CMV, cytomegalovirus; EBV, Epstein-Barr virus; LKM, liver-kidney microsomal antibody; MRCP, magnetic resonance cholangiopancreatography; SMA, smooth-muscle antibody; SPEP, serum protein electrophoresis.

Simply stated, the initial step is to perform appropriate blood tests in order to determine whether the patient has an isolated elevation of serum bilirubin. If so, is the bilirubin elevation due to an increased unconjugated or conjugated fraction? If the hyperbilirubinemia is accompanied by other liver test abnormalities, is the disorder hepatocellular or cholestatic? If cholestatic, is it intra- or extrahepatic? All of these questions can be answered with a thoughtful history, physical examination, and interpretation of laboratory and radiologic tests and procedures.

The bilirubin present in serum represents a balance between input from the production of bilirubin and hepatic/biliary removal of the pigment. Hyperbilirubinemia may result from (1) overproduction of bilirubin; (2) impaired uptake, conjugation, or excretion of bilirubin; or (3) regurgitation of unconjugated or conjugated bilirubin from damaged hepatocytes or bile ducts. An increase in unconjugated bilirubin in serum results from overproduction, impaired uptake, or conjugation of bilirubin. An increase in conjugated bilirubin is due to decreased excretion into the bile ductules or backward leakage of the pigment. The initial steps in evaluating the patient with jaundice are to determine (1) whether the hyperbilirubinemia is predominantly conjugated or unconjugated in nature and (2) whether other biochemical liver tests are abnormal.

The thoughtful interpretation of limited data permits a rational evaluation of the patient (Fig. 58-1). The following discussion will focus solely on the evaluation of the adult patient with jaundice.

ISOLATED ELEVATION OF SERUM BILIRUBIN

Unconjugated Hyperbilirubinemia The differential diagnosis of isolated unconjugated hyperbilirubinemia is limited (Table 58-1). The critical determination is whether the patient is suffering from a hemolytic process resulting in an overproduction of bilirubin (hemolytic disorders and ineffective erythropoiesis) or from impaired hepatic uptake/conjugation of bilirubin (drug effect or genetic disorders).

Hemolytic disorders that cause excessive heme production may be either inherited or acquired. Inherited disorders include spherocytosis, sickle cell anemia, thalassemia, and deficiency of red cell enzymes such as pyruvate kinase and glucose-6-phosphate dehydrogenase. In these conditions, the serum bilirubin level rarely exceeds $86 \mu\text{mol/L}$ (5 mg/dL). Higher levels may occur when there is coexistent renal or hepatocellular dysfunction or in acute hemolysis, such as a sickle cell crisis. In evaluating jaundice in patients with chronic hemolysis, it is important to remember the high incidence of pigmented (calcium bilirubinate) gallstones found in these

TABLE 58-1 CAUSES OF ISOLATED HYPERBILIRUBINEMIA

- I. Indirect hyperbilirubinemia
 - A. Hemolytic disorders
 1. Inherited
 - a. Spherocytosis, elliptocytosis, glucose-6-phosphate dehydrogenase and pyruvate kinase deficiencies
 - b. Sickle cell anemia
 2. Acquired
 - a. Microangiopathic hemolytic anemias
 - b. Paroxysmal nocturnal hemoglobinuria
 - c. Spur cell anemia
 - d. Immune hemolysis
 - e. Parasitic infections
 - (1) Malaria
 - (2) Babesiosis
 - B. Ineffective erythropoiesis
 1. Cobalamin, folate, and severe iron deficiencies
 2. Thalassemia
 - C. Increased bilirubin production
 1. Massive blood transfusion
 2. Resorption of hematoma
 - D. Drugs
 1. Rifampin
 2. Probenecid
 3. Ribavirin
 - E. Inherited conditions
 1. Crigler-Najjar types I and II
 2. Gilbert's syndrome
- II. Direct hyperbilirubinemia (inherited conditions)
 - A. Dubin-Johnson syndrome
 - B. Rotor syndrome

patients, which increases the likelihood of choledocholithiasis as an alternative explanation for hyperbilirubinemia.

Acquired hemolytic disorders include microangiopathic hemolytic anemia (e.g., hemolytic-uremic syndrome), paroxysmal nocturnal hemoglobinuria, spur cell anemia, immune hemolysis, and parasitic infections (e.g., malaria and babesiosis). Ineffective erythropoiesis occurs in cobalamin, folate, and iron deficiencies. Resorption of hematomas and massive blood transfusions both can result in increased hemoglobin release and overproduction of bilirubin.

In the absence of hemolysis, the physician should consider a problem with the hepatic uptake or conjugation of bilirubin. Certain drugs, including rifampin and probenecid, may cause unconjugated hyperbilirubinemia by diminishing hepatic uptake of bilirubin. Impaired bilirubin conjugation occurs in three genetic conditions: Crigler-Najjar syndrome types I and II and Gilbert's syndrome. *Crigler-Najjar type I* is an exceptionally rare condition found in neonates and characterized by severe jaundice (bilirubin $>342 \mu\text{mol/L}$ [$>20 \text{ mg/dL}$]) and neurologic impairment due to kernicterus, frequently leading to death in infancy or childhood. These patients have a complete absence of bilirubin UDPGT activity, usually due to mutations in the critical 3' domain of the *UDPGT* gene; are totally unable to conjugate bilirubin; and hence cannot excrete it.

Crigler-Najjar type II is somewhat more common than type I. Patients live into adulthood with serum bilirubin levels of $103\text{--}428 \mu\text{mol/L}$ ($6\text{--}25 \text{ mg/dL}$). In these patients, mutations in the bilirubin *UDPGT* gene cause the reduction—but not the complete eradication—of the enzyme's activity. Bilirubin *UDPGT* activity can be induced by the administration of phenobarbital, which can reduce serum bilirubin levels in these patients. Despite marked jaundice,

these patients usually survive into adulthood, although they may be susceptible to kernicterus under the stress of intercurrent illness or surgery.

Gilbert's syndrome is also marked by the impaired conjugation of bilirubin (to approximately one-third of normal) due to reduced bilirubin *UDPGT* activity. Patients with Gilbert's syndrome have mild unconjugated hyperbilirubinemia, with serum levels almost always $<103 \mu\text{mol/L}$ (6 mg/dL). The serum levels may fluctuate, and jaundice is often identified only during periods of fasting. The molecular defect in Gilbert's syndrome is linked to a reduction in transcription of the bilirubin *UDPGT* gene due to mutations in the promoter and, rarely, in the coding region. Unlike both Crigler-Najjar syndromes, Gilbert's syndrome is very common. The reported incidence is 3–7% of the population, with males predominating over females by a ratio of 2–7:1.

Conjugated Hyperbilirubinemia Elevated conjugated hyperbilirubinemia is found in two rare inherited conditions: *Dubin-Johnson syndrome* and *Rotor syndrome* (Table 58-1). Patients with either condition present with asymptomatic jaundice. The defect in Dubin-Johnson syndrome is the presence of mutations in the gene for MRP2. These patients have altered excretion of bilirubin into the bile ducts. Rotor syndrome may represent a deficiency of the major hepatic drug uptake transporters OATP1B1 and OATP1B3. Differentiating between these syndromes is possible but is clinically unnecessary due to their benign nature.

ELEVATION OF SERUM BILIRUBIN WITH OTHER LIVER TEST ABNORMALITIES

The remainder of this chapter will focus on the evaluation of patients with conjugated hyperbilirubinemia in the setting of other liver test abnormalities. This group of patients can be divided into those with a primary hepatocellular process and those with intra- or extrahepatic cholestasis. This distinction, which is based on the history and physical examination as well as the pattern of liver test abnormalities, guides the clinician's evaluation (Fig. 58-1).

History A complete medical history is perhaps the single most important part of the evaluation of the patient with unexplained jaundice. Important considerations include the use of or exposure to any chemical or medication, whether physician-prescribed, over-the-counter, complementary, or alternative medicines (e.g., herbal and vitamin preparations) or other drugs such as anabolic steroids. The patient should be carefully questioned about possible parenteral exposures, including transfusions, intravenous and intranasal drug use, tattooing, and sexual activity. Other important points include recent travel history; exposure to people with jaundice; exposure to possibly contaminated foods; occupational exposure to hepatotoxins; alcohol consumption; the duration of jaundice; and the presence of any accompanying signs and symptoms, such as arthralgias, myalgias, rash, anorexia, weight loss, abdominal pain, fever, pruritus, and changes in the urine and stool. While none of the latter manifestations is specific for any one condition, any of them can suggest a particular diagnosis. A history of arthralgias and myalgias predating jaundice suggests hepatitis, either viral or drug-related. Jaundice associated with the sudden onset of severe right-upper-quadrant pain and shaking chills suggests choledocholithiasis and ascending cholangitis.

Physical Examination The general assessment should include evaluation of the patient's nutritional status. Temporal and proximal muscle wasting suggests long-standing disease such as pancreatic cancer or cirrhosis. Stigmata of chronic liver disease, including spider nevi, palmar erythema, gynecomastia, caput medusae, Dupuytren's contractures, parotid gland enlargement, and testicular atrophy, are commonly seen in advanced alcoholic (Laennec's) cirrhosis and occasionally in other types of cirrhosis. An enlarged left supraclavicular node (Virchow's node) or a periumbilical nodule (Sister Mary Joseph's nodule) suggests an abdominal malignancy.

Jugular venous distention, a sign of right-sided heart failure, suggests hepatic congestion. Right pleural effusion in the absence of clinically apparent ascites may be seen in advanced cirrhosis.

The abdominal examination should focus on the size and consistency of the liver, on whether the spleen is palpable and hence enlarged, and on whether ascites is present. Patients with cirrhosis may have an enlarged left lobe of the liver, which is felt below the xiphoid, and an enlarged spleen. A grossly enlarged nodular liver or an obvious abdominal mass suggests malignancy. An enlarged tender liver could signify viral or alcoholic hepatitis; an infiltrative process such as amyloidosis; or, less often, an acutely congested liver secondary to right-sided heart failure. Severe right-upper-quadrant tenderness with respiratory arrest on inspiration (Murphy's sign) suggests cholecystitis. Ascites in the presence of jaundice suggests either cirrhosis or malignancy with peritoneal spread.

Laboratory Tests A battery of tests are helpful in the initial evaluation of a patient with unexplained jaundice. These include total and direct serum bilirubin measurement with fractionation; determination of serum aminotransferase, alkaline phosphatase, and albumin concentrations; and prothrombin time tests. Enzyme tests (alanine aminotransferase [ALT], aspartate aminotransferase [AST], and alkaline phosphatase [ALP]) are helpful in differentiating between a hepatocellular process and a cholestatic process (Table 58-1; Fig. 58-1)—a critical step in determining what additional workup is indicated. Patients with a hepatocellular process generally have a rise in the aminotransferases that is disproportionate to that in ALP, whereas patients with a cholestatic process have a rise in ALP that is disproportionate to that of the aminotransferases. The serum bilirubin can be prominently elevated in both hepatocellular and cholestatic conditions and therefore is not necessarily helpful in differentiating between the two.

In addition to enzyme tests, all jaundiced patients should have additional blood tests—specifically, an albumin level and a prothrombin time—to assess liver function. A low albumin level suggests a chronic process such as cirrhosis or cancer. A normal albumin level is suggestive of a more acute process such as viral hepatitis or choledocholithiasis. An elevated prothrombin time indicates either vitamin K deficiency due to prolonged jaundice and malabsorption of vitamin K or significant hepatocellular dysfunction. The failure of the prothrombin time to correct with parenteral administration of vitamin K indicates severe hepatocellular injury.

The results of the bilirubin, enzyme, albumin, and prothrombin time tests will usually indicate whether a jaundiced patient has a hepatocellular or a cholestatic disease and offer some indication of the duration and severity of the disease. The causes and evaluations of hepatocellular and cholestatic diseases are quite different.

Hepatocellular Conditions Hepatocellular diseases that can cause jaundice include viral hepatitis, drug or environmental toxicity, alcohol, and end-stage cirrhosis from any cause (Table 58-2). Wilson's disease occurs primarily in young adults. Autoimmune hepatitis is typically seen in young to middle-aged women but may affect men and women of any age. Alcoholic hepatitis can be differentiated from viral and toxin-related hepatitis by the pattern of the aminotransferases: patients with alcoholic hepatitis typically have an AST-to-ALT ratio of at least 2:1, and the AST level rarely exceeds 300 U/L. Patients with acute viral hepatitis and toxin-related injury severe enough to produce jaundice typically have aminotransferase levels >500 U/L, with the ALT greater than or equal to the AST. While ALT and AST values <8 times normal may be seen in either hepatocellular or cholestatic liver disease, values 25 times normal or higher are seen primarily in acute hepatocellular diseases. Patients with jaundice from cirrhosis can have normal or only slightly elevated aminotransferase levels.

When the clinician determines that a patient has a hepatocellular disease, appropriate testing for acute viral hepatitis includes

TABLE 58-2 HEPATOCELLULAR CONDITIONS THAT MAY PRODUCE JAUNDICE

Viral hepatitis
Hepatitis A, B, C, D, and E
Epstein-Barr virus
Cytomegalovirus
Herpes simplex virus
Alcoholic hepatitis
Drug toxicity
Predictable, dose-dependent (e.g., acetaminophen)
Unpredictable, idiosyncratic (e.g., isoniazid)
Environmental toxins
Vinyl chloride
Jamaica bush tea—pyrrolizidine alkaloids
Kava Kava
Wild mushrooms— <i>Amanita phalloides</i> , <i>A. verna</i>
Wilson's disease
Autoimmune hepatitis

a hepatitis A IgM antibody assay, a hepatitis B surface antigen and core IgM antibody assay, a hepatitis C viral RNA test, and, depending on the circumstances, a hepatitis E IgM antibody assay. Because it can take many weeks for hepatitis C antibody to become detectable, its assay is an unreliable test if acute hepatitis C is suspected. Studies for hepatitis D and E viruses, Epstein-Barr virus (EBV), and cytomegalovirus (CMV) may also be indicated. Ceruloplasmin is the initial screening test for Wilson's disease. Testing for autoimmune hepatitis usually includes an antinuclear antibody assay and measurement of specific immunoglobulins.

Drug-induced hepatocellular injury can be classified as either predictable or unpredictable. Predictable drug reactions are dose-dependent and affect all patients who ingest a toxic dose of the drug in question. The classic example is acetaminophen hepatotoxicity. Unpredictable or idiosyncratic drug reactions are not dose-dependent and occur in a minority of patients. A great number of drugs can cause idiosyncratic hepatic injury. Environmental toxins are also an important cause of hepatocellular injury. Examples include industrial chemicals such as vinyl chloride, herbal preparations containing pyrrolizidine alkaloids (Jamaica bush tea) or Kava Kava, and the mushrooms *Amanita phalloides* and *A. verna*, which contain highly hepatotoxic amatoxins.

Cholestatic Conditions When the pattern of the liver tests suggests a cholestatic disorder, the next step is to determine whether it is intra- or extrahepatic cholestasis (Fig. 58-1). Distinguishing intrahepatic from extrahepatic cholestasis may be difficult. History, physical examination, and laboratory tests often are not helpful. The next appropriate test is an ultrasound. The ultrasound is inexpensive, does not expose the patient to ionizing radiation, and can detect dilation of the intra- and extrahepatic biliary tree with a high degree of sensitivity and specificity. The absence of biliary dilation suggests intrahepatic cholestasis, while its presence indicates extrahepatic cholestasis. False-negative results occur in patients with partial obstruction of the common bile duct or in patients with cirrhosis or primary sclerosing cholangitis (PSC), in which scarring prevents the intrahepatic ducts from dilating.

Although ultrasonography may indicate extrahepatic cholestasis, it rarely identifies the site or cause of obstruction. The distal common bile duct is a particularly difficult area to visualize by ultrasound because of overlying bowel gas. Appropriate next tests include CT, magnetic resonance cholangiopancreatography (MRCP), endoscopic retrograde cholangiopancreatography (ERCP), and endoscopic ultrasound (EUS). CT scanning and MRCP are better than ultrasonography for assessing the head of the pancreas and for identifying choledocholithiasis in the distal

common bile duct, particularly when the ducts are not dilated. ERCP is the “gold standard” for identifying choledocholithiasis. Beyond its diagnostic capabilities, ERCP allows therapeutic interventions, including the removal of common bile duct stones and the placement of stents. MRCP has replaced ERCP as the initial diagnostic test in cases where the need for intervention is thought to be small. EUS displays sensitivity and specificity comparable to that of MRCP in the detection of bile duct obstruction. EUS also allows biopsy of suspected malignant lesions, but is invasive and requires sedation.

In patients with apparent *intrahepatic cholestasis*, the diagnosis is often made by serologic testing in combination with percutaneous liver biopsy. The list of possible causes of intrahepatic cholestasis is long and varied (Table 58-3). A number of conditions that typically cause a hepatocellular pattern of injury can also present as a cholestatic variant. Both hepatitis B and C viruses can cause cholestatic hepatitis (fibrosing cholestatic hepatitis). This disease variant has been reported in patients who have undergone solid organ transplantation. Hepatitis A and E, alcoholic hepatitis, and EBV or CMV infections may also present as cholestatic liver disease.

Drugs may cause intrahepatic cholestasis that is usually reversible after discontinuation of the offending agent, although it may take many months for cholestasis to resolve. Drugs most commonly associated with cholestasis are the anabolic and contraceptive steroids. Cholestatic hepatitis has been reported with chlorpromazine, imipramine, tolbutamide, sulindac, cimetidine, and erythromycin estolate. It also occurs in patients taking trimethoprim; sulfamethoxazole; and penicillin-based antibiotics such as ampicillin, dicloxacillin, and clavulanic acid. Rarely, cholestasis may be chronic and associated with progressive fibrosis despite early discontinuation of the offending drug. Chronic cholestasis has been associated with chlorpromazine and prochlorperazine.

Primary biliary cirrhosis is an autoimmune disease predominantly affecting middle-aged women and characterized by progressive destruction of interlobular bile ducts. The diagnosis is made by the detection of antimitochondrial antibody, which is found in 95% of patients. *Primary sclerosing cholangitis* is characterized by the destruction and fibrosis of larger bile ducts. The diagnosis of PSC is made with cholangiography (either MRCP or ERCP), which demonstrates the pathognomonic segmental strictures. Approximately 75% of patients with PSC have inflammatory bowel disease.

The *vanishing bile duct syndrome* and *adult bile ductopenia* are rare conditions in which a decreased number of bile ducts are seen in liver biopsy specimens. The histologic picture is similar to that in primary biliary cirrhosis. This picture is seen in patients who develop chronic rejection after liver transplantation and in those who develop graft-versus-host disease after bone marrow transplantation. Vanishing bile duct syndrome also occurs in rare cases of sarcoidosis, in patients taking certain drugs (including chlorpromazine), and idiopathically.

There are also familial forms of intrahepatic cholestasis. The familial intrahepatic cholestatic syndromes include *progressive familial intrahepatic cholestasis* (PFIC) types 1–3 and *benign recurrent cholestasis* (BRC). PFIC1 and BRC are autosomal recessive diseases that result from mutations in the *ATP8B1* gene that encodes a protein belonging to the subfamily of P-type ATPases; the exact function of this protein remains poorly defined. While PFIC1 is a progressive condition that manifests in childhood, BRC presents later and is marked by recurrent episodes of jaundice and pruritus; the episodes are self-limited but can be debilitating. PFIC2 is caused by mutations in the *ABCB11* gene, which encodes the bile salt export pump, and PFIC3 is caused by mutations in the multidrug-resistant P-glycoprotein 3. *Cholestasis of pregnancy* occurs in the second and third trimesters and resolves after delivery. Its cause is unknown, but the condition is probably inherited, and cholestasis can be triggered by estrogen administration.

TABLE 58-3 CHOLESTATIC CONDITIONS THAT MAY PRODUCE JAUNDICE

- I. Intrahepatic
 - A. Viral hepatitis
 1. Fibrosing cholestatic hepatitis—hepatitis B and C
 2. Hepatitis A, Epstein-Barr virus infection, cytomegalovirus infection
 - B. Alcoholic hepatitis
 - C. Drug toxicity
 1. Pure cholestasis—anabolic and contraceptive steroids
 2. Cholestatic hepatitis—chlorpromazine, erythromycin estolate
 3. Chronic cholestasis—chlorpromazine and prochlorperazine
 - D. Primary biliary cirrhosis
 - E. Primary sclerosing cholangitis
 - F. Vanishing bile duct syndrome
 1. Chronic rejection of liver transplants
 2. Sarcoidosis
 3. Drugs
 - G. Congestive hepatopathy and ischemic hepatitis
 - H. Inherited conditions
 1. Progressive familial intrahepatic cholestasis
 2. Benign recurrent cholestasis
 - I. Cholestasis of pregnancy
 - J. Total parenteral nutrition
 - K. Nonhepatobiliary sepsis
 - L. Benign postoperative cholestasis
 - M. Paraneoplastic syndrome
 - N. Venocclusive disease
 - O. Graft-versus-host disease
 - P. Infiltrative disease
 1. Tuberculosis
 2. Lymphoma
 3. Amyloidosis
 - Q. Infections
 1. Malaria
 2. Leptospirosis
- II. Extrahepatic
 - A. Malignant
 1. Cholangiocarcinoma
 2. Pancreatic cancer
 3. Gallbladder cancer
 4. Ampullary cancer
 5. Malignant involvement of the porta hepatis lymph nodes
 - B. Benign
 1. Choledocholithiasis
 2. Postoperative biliary strictures
 3. Primary sclerosing cholangitis
 4. Chronic pancreatitis
 5. AIDS cholangiopathy
 6. Mirizzi's syndrome
 7. Parasitic disease (ascariasis)

Other causes of intrahepatic cholestasis include total parenteral nutrition (TPN); nonhepatobiliary sepsis; benign postoperative cholestasis; and a paraneoplastic syndrome associated with a number of different malignancies, including Hodgkin's disease, medullary thyroid cancer, renal cell cancer, renal sarcoma, T cell lymphoma, prostate cancer, and several gastrointestinal malignancies. The term *Stauffer's syndrome* has been used for intrahepatic cholestasis specifically associated with renal cell cancer. In patients developing cholestasis in the intensive care unit, the major considerations

59 Abdominal Swelling and Ascites

Kathleen E. Corey, Lawrence S. Friedman

should be sepsis, ischemic hepatitis (“shock liver”), and TPN jaundice. Jaundice occurring after bone marrow transplantation is most likely due to veno-occlusive disease or graft-versus-host disease. In addition to hemolysis, sickle cell disease may cause intrahepatic and extrahepatic cholestasis. Jaundice is a late finding in heart failure caused by hepatic congestion and hepatocellular hypoxia. Ischemic hepatitis is a distinct entity of acute hypoperfusion characterized by an acute and dramatic elevation in the serum aminotransferases followed by a gradual peak in serum bilirubin.

Jaundice with associated liver dysfunction can be seen in severe cases of *Plasmodium falciparum* malaria. The jaundice in these cases is due to a combination of indirect hyperbilirubinemia from hemolysis and both cholestatic and hepatocellular jaundice. Weil’s disease, a severe presentation of leptospirosis, is marked by jaundice with renal failure, fever, headache, and muscle pain.

Causes of *extrahepatic cholestasis* can be split into malignant and benign (Table 58-3). Malignant causes include pancreatic, gallbladder, and ampullary cancers as well as cholangiocarcinoma. This last malignancy is most commonly associated with PSC and is exceptionally difficult to diagnose because its appearance is often identical to that of PSC. Pancreatic and gallbladder tumors as well as cholangiocarcinoma are rarely resectable and have poor prognoses. Ampullary carcinoma has the highest surgical cure rate of all the tumors that present as painless jaundice. Hilar lymphadenopathy due to metastases from other cancers may cause obstruction of the extrahepatic biliary tree.

Cholelithiasis is the most common cause of extrahepatic cholestasis. The clinical presentation can range from mild right-upper-quadrant discomfort with only minimal elevations of enzyme test values to ascending cholangitis with jaundice, sepsis, and circulatory collapse. PSC may occur with clinically important strictures limited to the extrahepatic biliary tree. IgG4-associated cholangitis is marked by stricturing of the biliary tree. It is critical that the clinician differentiate this condition from PSC as it is responsive to glucocorticoid therapy. In rare instances, chronic pancreatitis causes strictures of the distal common bile duct, where it passes through the head of the pancreas. AIDS cholangiopathy is a condition that is usually due to infection of the bile duct epithelium with CMV or cryptosporidia and has a cholangiographic appearance similar to that of PSC. The affected patients usually present with greatly elevated serum alkaline phosphatase levels (mean, 800 IU/L), but the bilirubin level is often near normal. These patients do not typically present with jaundice.

GLOBAL CONSIDERATIONS



While extrahepatic biliary obstruction and drugs are common causes of new-onset jaundice in developed countries, infections remain the leading cause in developing countries. Liver involvement and jaundice are observed with numerous infections, particularly malaria, babesiosis, severe leptospirosis, infections due to *Mycobacterium tuberculosis* and the *Mycobacterium avium* complex, typhoid fever, viral hepatitis secondary to infection with hepatitis viruses A–E, EBV and CMV infections, late phases of yellow fever, dengue hemorrhagic fever, schistosomiasis, fascioliasis, clonorchiasis, opisthorchiasis, ascariasis, echinococcosis, hepatosplenic candidiasis, disseminated histoplasmosis, cryptococcosis, coccidioidomycosis, ehrlichiosis, chronic Q fever, yersiniosis, brucellosis, syphilis, and leprosy. Bacterial infections that do not necessarily involve the liver and bile ducts may also lead to jaundice, as in cholestasis of sepsis.

ACKNOWLEDGMENT

This chapter is a revised version of chapters that have appeared in prior editions of *Harrison’s* in which Marshall M. Kaplan was a co-author together with Daniel Pratt.

ABDOMINAL SWELLING

Abdominal swelling is a manifestation of numerous diseases. Patients may complain of bloating or abdominal fullness and may note increasing abdominal girth on the basis of increased clothing or belt size. Abdominal discomfort is often reported, but pain is less frequent. When abdominal pain does accompany swelling, it is frequently the result of an intraabdominal infection, peritonitis, or pancreatitis. Patients with abdominal distention from *ascites* (fluid in the abdomen) may report the new onset of an inguinal or umbilical hernia. Dyspnea may result from pressure against the diaphragm and the inability to expand the lungs fully.

CAUSES

The causes of abdominal swelling can be remembered conveniently as the *six Fs*: flatus, fat, fluid, fetus, feces, or a “fatal growth” (often a neoplasm).

Flatus Abdominal swelling may be the result of increased intestinal gas. The normal small intestine contains approximately 200 mL of gas made up of nitrogen, oxygen, carbon dioxide, hydrogen, and methane. Nitrogen and oxygen are consumed (swallowed), whereas carbon dioxide, hydrogen, and methane are produced intraluminally by bacterial fermentation. Increased intestinal gas can occur in a number of conditions. *Aerophagia*, the swallowing of air, can result in increased amounts of oxygen and nitrogen in the small intestine and lead to abdominal swelling. *Aerophagia* typically results from gulping food; chewing gum; smoking; or as a response to anxiety, which can lead to repetitive belching. In some cases, increased intestinal gas is the consequence of bacterial metabolism of excess fermentable substances such as lactose and other oligosaccharides, which can lead to production of hydrogen, carbon dioxide, or methane. In many cases, the precise cause of abdominal distention cannot be determined. In some persons, particularly those with irritable bowel syndrome and bloating, the subjective sense of abdominal pressure is attributable to impaired intestinal transit of gas rather than increased gas volume. Abdominal distention—an objective increase in girth—is the result of a lack of coordination between diaphragmatic contraction and anterior abdominal wall relaxation, a response in some cases to an increase in intraabdominal volume loads. Occasionally, increased lumbar lordosis accounts for apparent abdominal distention.

Fat Weight gain with an increase in abdominal fat can result in an increase in abdominal girth and can be perceived as abdominal swelling. Abdominal fat may be caused by an imbalance between caloric intake and energy expenditure associated with a poor diet and sedentary lifestyle; it also can be a manifestation of certain diseases, such as Cushing’s syndrome. Excess abdominal fat has been associated with an increased risk of insulin resistance and cardiovascular disease.

Fluid The accumulation of fluid within the abdominal cavity (*ascites*) often results in abdominal distention and is discussed in detail below.

Fetus Pregnancy results in increased abdominal girth. Typically, an increase in abdominal size is first noted at 12–14 weeks of gestation, when the uterus moves from the pelvis into the abdomen. Abdominal distention may be seen before this point as a result of fluid retention and relaxation of the abdominal muscles.

Feces In the setting of severe constipation or intestinal obstruction, increased stool in the colon leads to increased abdominal girth. These conditions are often accompanied by abdominal discomfort or pain, nausea, and vomiting and can be diagnosed by imaging studies.

Fatal Growth An abdominal mass can result in abdominal swelling. Enlargement of the intraabdominal organs, specifically the liver (hepatomegaly) or spleen (splenomegaly), or an abdominal aortic aneurysm

can result in abdominal distention. Bladder distention also may result in abdominal swelling. In addition, malignancies, abscesses, or cysts can grow to sizes that lead to increased abdominal girth.

APPROACH TO THE PATIENT: Abdominal Swelling

HISTORY

Determining the etiology of abdominal swelling begins with history-taking and a physical examination. Patients should be questioned regarding symptoms suggestive of malignancy, including weight loss, night sweats, and anorexia. Inability to pass stool or flatus together with nausea or vomiting suggests bowel obstruction, severe constipation, or an ileus (lack of peristalsis). Increased eructation and flatus may point toward aerophagia or increased intestinal production of gas. Patients should be questioned about risk factors for or symptoms of chronic liver disease, including excessive alcohol use and jaundice, which suggest ascites. Patients should also be asked about other symptoms of medical conditions, including heart failure and tuberculosis, which may cause ascites.

PHYSICAL EXAMINATION

Physical examination should include an assessment for signs of systemic disease. The presence of lymphadenopathy, especially supraclavicular lymphadenopathy (*Virchow's node*), suggests metastatic abdominal malignancy. Care should be taken during the cardiac examination to evaluate for elevation of jugular venous pressure (JVP); *Kussmaul's sign* (elevation of the JVP during inspiration); a pericardial knock, which may be seen in heart failure or constrictive pericarditis; or a murmur of tricuspid regurgitation. Spider angiomata, palmar erythema, dilated superficial veins around the umbilicus (*caput medusae*), and gynecomastia suggest chronic liver disease.

The abdominal examination should begin with inspection for the presence of uneven distention or an obvious mass. Auscultation should follow. The absence of bowel sounds or the presence of high-pitched localized bowel sounds points toward an ileus or intestinal obstruction. An umbilical venous hum may suggest the presence of portal hypertension, and a harsh bruit over the liver is heard rarely in patients with hepatocellular carcinoma or alcoholic hepatitis. Abdominal swelling caused by intestinal gas can be differentiated from swelling caused by fluid or a solid mass by percussion; an abdomen filled with gas is tympanic, whereas an abdomen containing a mass or fluid is dull to percussion. The absence of abdominal dullness, however, does not exclude ascites, because a minimum of 1500 mL of ascitic fluid is required for detection on physical examination. Finally, the abdomen should be palpated to assess for tenderness, a mass, enlargement of the spleen or liver, or presence of a nodular liver suggesting cirrhosis or tumor. Light palpation of the liver may detect pulsations suggesting retrograde vascular flow from the heart in patients with right-sided heart failure, particularly tricuspid regurgitation.

IMAGING AND LABORATORY EVALUATION

Abdominal x-rays can be used to detect dilated loops of bowel suggesting intestinal obstruction or ileus. Abdominal ultrasonography can detect as little as 100 mL of ascitic fluid, hepatosplenomegaly, a nodular liver, or a mass. Ultrasonography is often inadequate to detect retroperitoneal lymphadenopathy or a pancreatic lesion because of overlying bowel gas. If malignancy or pancreatic disease is suspected, CT can be performed. CT may also detect changes associated with advanced cirrhosis and portal hypertension (Fig. 59-1).

Laboratory evaluation should include liver biochemical testing, serum albumin level measurement, and prothrombin time determination (international normalized ratio) to assess hepatic function as well as a complete blood count to evaluate for the presence of cytopenias that may result from portal hypertension or of leukocytosis, anemia, and thrombocytosis that may result from systemic infection. Serum amylase and lipase levels should be checked to evaluate the patient



FIGURE 59-1 CT of a patient with a cirrhotic, nodular liver (white arrow), splenomegaly (yellow arrow), and ascites (arrowheads).

for acute pancreatitis. Urinary protein quantitation is indicated when nephrotic syndrome, which may cause ascites, is suspected.

In selected cases, the hepatic venous pressure gradient (pressure across the liver between the portal and hepatic veins) can be measured via cannulation of the hepatic vein to confirm that ascites is caused by cirrhosis (Chap. 365). In some cases, a liver biopsy may be necessary to confirm cirrhosis.

ASCITES

PATHOGENESIS IN THE PRESENCE OF CIRRHOSIS

Ascites in patients with cirrhosis is the result of portal hypertension and renal salt and water retention. Similar mechanisms contribute to ascites formation in heart failure. Portal hypertension signifies elevation of the pressure within the portal vein. According to Ohm's law, pressure is the product of resistance and flow. Increased hepatic resistance occurs by several mechanisms. First, the development of hepatic fibrosis, which defines cirrhosis, disrupts the normal architecture of the hepatic sinusoids and impedes normal blood flow through the liver. Second, activation of hepatic stellate cells, which mediate fibrogenesis, leads to smooth-muscle contraction and fibrosis. Finally, cirrhosis is associated with a decrease in endothelial nitric oxide synthetase (eNOS) production, which results in decreased nitric oxide production and increased intrahepatic vasoconstriction.

The development of cirrhosis is also associated with increased systemic circulating levels of nitric oxide (contrary to the decrease seen intrahepatically) as well as increased levels of vascular endothelial growth factor and tumor necrosis factor that result in splanchnic arterial vasodilation. Vasodilation of the splanchnic circulation results in pooling of blood and a decrease in the effective circulating volume, which is perceived by the kidneys as hypovolemia. Compensatory vasoconstriction via release of antidiuretic hormone ensues; the consequences are free water retention and activation of the sympathetic nervous system and the renin angiotensin aldosterone system, which lead in turn to renal sodium and water retention.

PATHOGENESIS IN THE ABSENCE OF CIRRHOSIS

Ascites in the absence of cirrhosis generally results from peritoneal carcinomatosis, peritoneal infection, or pancreatic disease. Peritoneal carcinomatosis can result from primary peritoneal malignancies such as mesothelioma or sarcoma, abdominal malignancies such as gastric or colonic adenocarcinoma, or metastatic disease from breast or lung

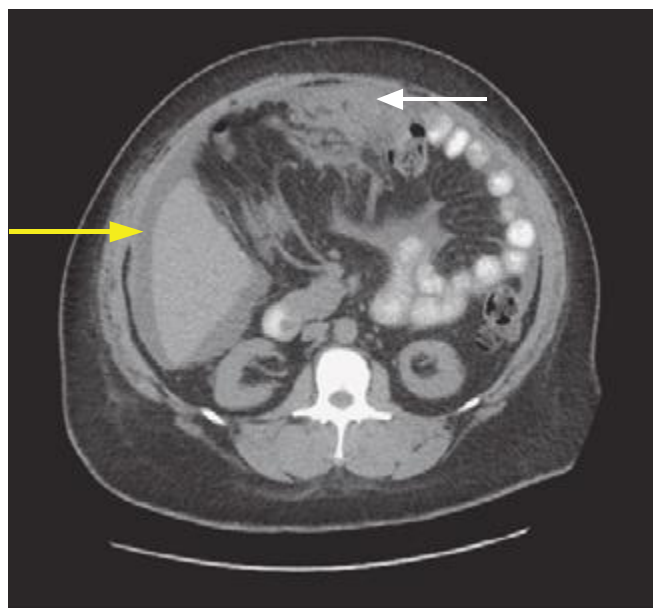


FIGURE 59-2 CT of a patient with peritoneal carcinomatosis (white arrow) and ascites (yellow arrow).

carcinoma or melanoma (Fig. 59-2). The tumor cells lining the peritoneum produce a protein-rich fluid that contributes to the development of ascites. Fluid from the extracellular space is drawn into the peritoneum, further contributing to the development of ascites. Tuberculous peritonitis causes ascites via a similar mechanism; tubercles deposited on the peritoneum exude a proteinaceous fluid. Pancreatic ascites results from leakage of pancreatic enzymes into the peritoneum.

CAUSES

Cirrhosis accounts for 84% of cases of ascites. Cardiac ascites, peritoneal carcinomatosis, and “mixed” ascites resulting from cirrhosis and a second disease account for 10–15% of cases. Less common causes of ascites include massive hepatic metastasis, infection (tuberculosis, *Chlamydia* infection), pancreatitis, and renal disease (nephrotic syndrome). Rare causes of ascites include hypothyroidism and familial Mediterranean fever.

EVALUATION

Once the presence of ascites has been confirmed, the etiology of the ascites is best determined by *paracentesis*, a bedside procedure in which a needle or small catheter is passed transcutaneously to extract ascitic fluid from the peritoneum. The lower quadrants are the most frequent sites for paracentesis. The left lower quadrant is preferred because of the greater depth of ascites and the thinner abdominal wall.

Paracentesis is a safe procedure even in patients with coagulopathy; complications, including abdominal wall hematomas, hypotension, hepatorenal syndrome, and infection, are infrequent.

Once ascitic fluid has been extracted, its gross appearance should be examined. Turbid fluid can result from the presence of infection or tumor cells. White, milky fluid indicates a triglyceride level >200 mg/dL (and often >1000 mg/dL), which is the hallmark of *chylous ascites*. Chylous ascites results from lymphatic disruption that may occur with trauma, cirrhosis, tumor, tuberculosis, or certain congenital abnormalities. Dark brown fluid can reflect a high bilirubin concentration and indicates biliary tract perforation. Black fluid may indicate the presence of pancreatic necrosis or metastatic melanoma.

The ascitic fluid should be sent for measurement of albumin and total protein levels, cell and differential counts, and, if infection is suspected, Gram’s stain and culture, with inoculation into blood culture bottles at the patient’s bedside to maximize the yield. A serum albumin level should be measured simultaneously to permit calculation of the *serum-ascites albumin gradient* (SAAG).

The SAAG is useful for distinguishing ascites caused by portal hypertension from nonportal hypertensive ascites (Fig. 59-3). The SAAG reflects the pressure within the hepatic sinusoids and correlates with the hepatic venous pressure gradient. The SAAG is calculated by subtracting the ascitic albumin concentration from the serum albumin level and does not change with diuresis. A SAAG ≥ 1.1 g/dL reflects the presence of portal hypertension and indicates that the ascites is due to increased pressure in the hepatic sinusoids. According to Starling’s law, a high SAAG reflects the oncotic pressure that counterbalances the portal pressure. Possible causes include cirrhosis, cardiac ascites, hepatic vein thrombosis (Budd-Chiari syndrome), sinusoidal obstruction syndrome (veno-occlusive disease), or massive liver metastases. A SAAG <1.1 g/dL indicates that the ascites is not related to portal hypertension, as in tuberculous peritonitis, peritoneal carcinomatosis, or pancreatic ascites.

For high-SAAG (≥ 1.1) ascites, the ascitic protein level can provide further clues to the etiology (Fig. 59-3). An ascitic protein level of ≥ 2.5 g/dL indicates that the hepatic sinusoids are normal and are allowing passage of protein into the ascites, as occurs in cardiac ascites, early Budd-Chiari syndrome, or sinusoidal obstruction syndrome. An ascitic protein level <2.5 g/dL indicates that the hepatic sinusoids have been damaged and scarred and no longer allow passage of protein, as occurs with cirrhosis, late Budd-Chiari syndrome, or massive liver metastases. Pro-brain-type natriuretic peptide (BNP) is a natriuretic hormone released by the heart as a result of increased volume and ventricular wall stretch. High levels of BNP in serum occur in heart failure and may be useful in identifying heart failure as the cause of high-SAAG ascites.

Further tests are indicated only in specific clinical circumstances. When secondary peritonitis resulting from a perforated hollow viscus is suspected, ascitic glucose and lactate dehydrogenase (LDH) levels can

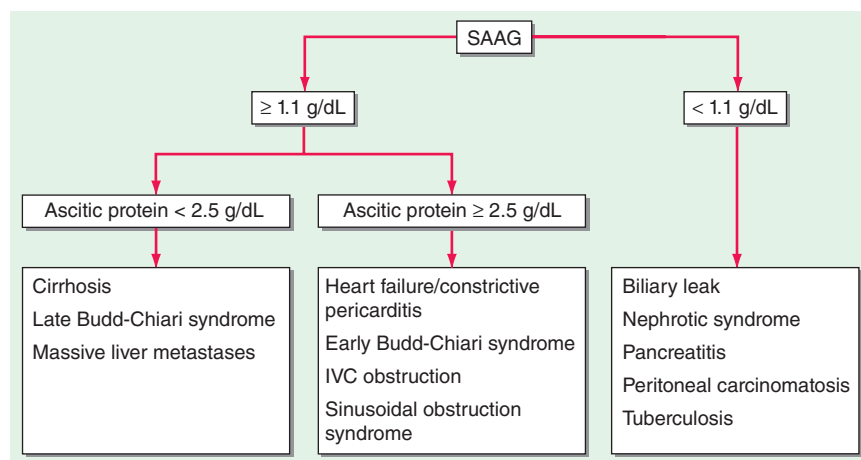


FIGURE 59-3 Algorithm for the diagnosis of ascites according to the serum-ascites albumin gradient (SAAG). IVC, inferior vena cava.

be measured. In contrast to “spontaneous” bacterial peritonitis, which may complicate cirrhotic ascites (see “Complications,” below), secondary peritonitis is suggested by an ascitic glucose level <50 mg/dL, an ascitic LDH level higher than the serum LDH level, and the detection of multiple pathogens on ascitic fluid culture. When pancreatic ascites is suspected, the ascitic amylase level should be measured and is typically >1000 mg/dL. Cytology can be useful in the diagnosis of peritoneal carcinomatosis. At least 50 mL of fluid should be obtained and sent for immediate processing. Tuberculous peritonitis is typically associated with ascitic fluid lymphocytosis but can be difficult to diagnose by paracentesis. A smear for acid-fast bacilli has a diagnostic sensitivity of only 0 to 3%; a culture increases the sensitivity to 35–50%. In patients without cirrhosis, an elevated ascitic adenosine deaminase level has a sensitivity of >90% when a cut-off value of 30–45 U/L is used. When the cause of ascites remains uncertain, laparotomy or laparoscopy with peritoneal biopsies for histology and culture remains the gold standard.

TREATMENT ASCITES

The initial treatment for cirrhotic ascites is restriction of sodium intake to 2 g/d. When sodium restriction alone is inadequate to control ascites, oral diuretics—typically the combination of spironolactone and furosemide—are used. Spironolactone is an aldosterone antagonist that inhibits sodium resorption in the distal convoluted tubule of the kidney. Use of spironolactone may be limited by hyponatremia, hyperkalemia, and painful gynecomastia. If the gynecomastia is distressing, amiloride (5–40 mg/d) may be substituted for spironolactone. Furosemide is a loop diuretic that is generally combined with spironolactone in a ratio of 40:100; maximal daily doses of spironolactone and furosemide are 400 mg and 160 mg, respectively.

Refractory cirrhotic ascites is defined by the persistence of ascites despite sodium restriction and maximal (or maximally tolerated) diuretic use. Pharmacologic therapy for refractory ascites includes the addition of midodrine, an α_1 -adrenergic antagonist, or clonidine, an α_2 -adrenergic antagonist, to diuretic therapy. These agents act as vasoconstrictors, counteracting splanchnic vasodilation. Midodrine alone or in combination with clonidine improves systemic hemodynamics and control of ascites over that obtained with diuretics alone. Although β -adrenergic blocking agents (beta blockers) are often prescribed to prevent variceal hemorrhage in patients with cirrhosis, the use of beta blockers in patients with refractory ascites is associated with decreased survival rates.

When medical therapy alone is insufficient, refractory ascites can be managed by repeated large-volume paracentesis (LVP) or a transjugular intrahepatic peritoneal shunt (TIPS)—a radiologically placed portosystemic shunt that decompresses the hepatic sinusoids. Intravenous infusion of albumin accompanying LVP decreases the risk of “post-paracentesis circulatory dysfunction” and death. Patients undergoing LVP should receive IV albumin infusions of

6–8 g/L of ascitic fluid removed. TIPS placement is superior to LVP in reducing the reaccumulation of ascites but is associated with an increased frequency of hepatic encephalopathy, with no difference in mortality rates.

Malignant ascites does not respond to sodium restriction or diuretics. Patients must undergo serial LVPs, transcutaneous drainage catheter placement, or, rarely, creation of a peritoneovenous shunt (a shunt from the abdominal cavity to the vena cava).

Ascites caused by tuberculous peritonitis is treated with standard antituberculosis therapy. Noncirrhotic ascites of other causes is treated by correction of the precipitating condition.

COMPLICATIONS

Spontaneous bacterial peritonitis (SBP; [Chap. 159](#)) is a common and potentially lethal complication of cirrhotic ascites. Occasionally, SBP also complicates ascites caused by nephrotic syndrome, heart failure, acute hepatitis, and acute liver failure but is rare in malignant ascites. Patients with SBP generally note an increase in abdominal girth; however, abdominal tenderness is found in only 40% of patients, and rebound tenderness is uncommon. Patients may present with fever, nausea, vomiting, or the new onset of or exacerbation of preexisting hepatic encephalopathy.

SBP is defined by a polymorphonuclear neutrophil (PMN) count of ≥ 250 /L in the ascitic fluid. Cultures of ascitic fluid typically reveal one bacterial pathogen. The presence of multiple pathogens in the setting of an elevated ascitic PMN count suggests *secondary peritonitis* from a ruptured viscus or abscess ([Chap. 159](#)). The presence of multiple pathogens without an elevated PMN count suggests bowel perforation from the paracentesis needle. SBP is generally the result of enteric bacteria that have translocated across an edematous bowel wall. The most common pathogens are gram-negative rods, including *Escherichia coli* and *Klebsiella*, as well as streptococci and enterococci.

Treatment of SBP with an antibiotic such as IV cefotaxime is effective against gram-negative and gram-positive aerobes. A 5-day course of treatment is sufficient if the patient improves clinically. Nosocomial or health care–acquired SBP is frequently caused by multidrug-resistant bacteria, and initial antibiotic therapy should be guided by the local bacterial epidemiology.

Cirrhotic patients with a history of SBP, an ascitic fluid total protein concentration <1 g/dL, or active gastrointestinal bleeding should receive prophylactic antibiotics to prevent SBP; oral daily norfloxacin is commonly used. Diuresis increases the activity of ascitic fluid protein opsonins and may decrease the risk of SBP.

Hepatic hydrothorax occurs when ascites, often caused by cirrhosis, migrates via fenestrae in the diaphragm into the pleural space. This condition can result in shortness of breath, hypoxia, and infection. Treatment is similar to that for cirrhotic ascites and includes sodium restriction, diuretics, and, if needed, thoracentesis or TIPS placement. Chest tube placement should be avoided.

SECTION 7 ALTERATIONS IN RENAL AND URINARY TRACT FUNCTION

60e Dysuria, Bladder Pain, and the Interstitial Cystitis/Bladder Pain Syndrome

John W. Warren

Dysuria and bladder pain are two symptoms that commonly call attention to the lower urinary tract.

DYSURIA

Dysuria, or pain that occurs during urination, is commonly perceived as burning or stinging in the urethra and is a symptom of several syndromes. The presence or absence of *other* symptoms is often helpful in distinguishing among these conditions. Some of these syndromes differ in men and women.

WOMEN

Approximately 50% of women experience dysuria at some time in their lives; ~20% report having had dysuria within the past year. Most dysuria syndromes in women can be categorized into two broad groups: bacterial cystitis and lower genital tract infections.

Bacterial cystitis is usually caused by *Escherichia coli*; a few other gram-negative rods and *Staphylococcus saprophyticus* can also be responsible. Bacterial cystitis is acute in onset and manifests not only as dysuria but also as urinary frequency, urinary urgency, suprapubic pain, and/or hematuria.

The lower genital tract infections include vaginitis, urethritis, and ulcerative lesions; many of these infections are caused by sexually transmitted organisms and should be considered particularly in young women who have new or multiple sexual partners or whose partner(s) do not use condoms. The onset of dysuria associated with these syndromes is more gradual than in bacterial cystitis and is thought (but not proven) to result from the flow of urine over damaged epithelium. Frequency, urgency, suprapubic pain, and hematuria are reported less frequently than in bacterial cystitis. Vaginitis, caused by *Candida albicans* or *Trichomonas vaginalis*, presents as vaginal discharge or irritation. Urethritis is a consequence of infection by *Chlamydia trachomatis* or *Neisseria gonorrhoeae*. Ulcerative genital lesions may be caused by herpes simplex virus and several other specific organisms.

Among women presenting with dysuria, the probability of bacterial cystitis is ~50%. This figure rises to >90% if four criteria are fulfilled: dysuria and frequency without vaginal discharge or irritation. Present standards suggest that women meeting these four criteria, if they are otherwise healthy, are not pregnant, and have an apparently normal urinary tract, can be diagnosed with uncomplicated bacterial cystitis and treated empirically with appropriate antibiotics. Other women with dysuria should be further evaluated by urine dipstick, urine culture, and a pelvic examination.

MEN

Dysuria is less common among men. The syndromes presenting as dysuria are similar to those in women but with some important distinctions.

In the majority of men with dysuria, frequency, urgency, and/or suprapubic, penile, and/or perineal pain, the prostate is involved, either as the source of infection or as an obstruction to urine flow. Bacterial prostatitis is usually caused by *E. coli* or another gram-negative rod, with one of two presentations. *Acute bacterial prostatitis* presents with fever and chills; prostate examination should be gentle or not performed at all, as massage may result in a wave of bacteremia. *Chronic bacterial prostatitis* presents as recurrent episodes of bacterial cystitis;

prostate examination with massage demonstrates prostatic bacteria and leukocytes. *Benign prostatic hyperplasia* (BPH) can obstruct urine flow, with consequent symptoms of weak stream, hesitancy, and dribbling. If a bacterial infection develops behind the obstructing prostate, dysuria and other symptoms of cystitis will occur. Men whose symptoms are consistent with bacterial cystitis should be evaluated with urinalysis and urine culture.

Several sexually transmitted infections can manifest as dysuria. Urethritis (usually without urinary frequency) presents as a urethral discharge and can be caused by *C. trachomatis*, *N. gonorrhoeae*, *Mycoplasma genitalium*, *Ureaplasma urealyticum*, or *T. vaginalis*. Herpes simplex, chancroid, and other ulcerous lesions may present as dysuria, again without urinary frequency.

For further discussion, see Chaps. 162 and 163.

EITHER WOMEN OR MEN

Other causes of dysuria may be found in patients of either sex. Some cases are acute and include lower urinary tract stones, trauma, and urethral exposure to topical chemicals. Others may be relatively chronic and attributable to lower urinary tract cancers, certain medications, Behçet's syndrome, reactive arthritis, a poorly understood entity known as *chronic urethral syndrome*, and interstitial cystitis/bladder pain syndrome (see below).

BLADDER PAIN

Studies indicate that patients perceive pain as coming from the urinary bladder if it is suprapubic in location, alters with bladder filling or emptying, and/or is associated with urinary symptoms such as urgency and frequency. Bladder pain occurring acutely (i.e., over hours or a day or two) is helpful in distinguishing bacterial cystitis from urethritis, vaginitis, and other genital infections. Chronic or recurrent bladder pain may accompany lower urinary tract stones; bladder, uterine, cervical, vaginal, urethral, or prostate cancer; urethral diverticulum; cystitis induced by radiation or certain medications; tuberculous cystitis; bladder neck obstruction; neurogenic bladder; urogenital prolapse; or BPH. In the absence of these conditions, the diagnosis of interstitial cystitis/bladder pain syndrome (IC/BPS) should be considered.

INTERSTITIAL CYSTITIS/BLADDER PAIN SYNDROME

Most clinicians with outpatient practices see undiagnosed cases of IC/BPS. This chronic condition is characterized by pain perceived to be from the urinary bladder, urinary urgency and frequency, and nocturia. The majority of cases are diagnosed in women. Symptoms wax and wane for months or years or possibly even for the rest of the patient's life. The spectrum of symptom intensity is broad. The pain can be excruciating, urgency can be distressing, frequency can be up to 60 times per 24 h, and nocturia can cause sleep deprivation. These symptoms can disrupt daily activities, work schedules, and personal relationships; patients with IC/BPS report less life satisfaction than do those with end-stage renal disease.

IC/BPS is not a new disease, having first been described in the late nineteenth century in a patient with the symptoms mentioned above and a single ulcer visible on cystoscopy (now called a *Hunner's lesion* after the urologist who first reported it). Over the ensuing decades, it became clear that many patients with similar symptoms had no ulcer. It is now appreciated that only up to 10% of patients with IC/BPS have a Hunner's lesion. The definition of IC/BPS, its diagnostic features, and even its name continue to evolve. The American Urological Association has defined IC/BPS as "an unpleasant sensation (pain, pressure, discomfort) perceived to be related to the urinary bladder, associated with lower urinary tract symptoms of more than six weeks' duration, in the absence of infection or other identifiable causes."

Many patients with IC/BPS also have other syndromes, such as fibromyalgia, chronic fatigue syndrome, irritable bowel syndrome, and migraine. These syndromes collectively are known as *functional somatic syndromes* (FSSs): chronic conditions in which pain and fatigue are prominent features but laboratory tests and histologic findings are normal. Like IC/BPS, the FSSs often are associated with depression and anxiety. The majority of FSSs affect more women than men, and more than one FSS can affect a single patient. Because of its similar features and comorbidity, IC/BPS sometimes is considered an FSS.

EPIDEMIOLOGY

Contemporary population studies of IC/BPS in the United States indicate a prevalence of 3–6% among women and 2–4% among men. For decades, it was thought that IC/BPS occurred mostly in women. These prevalence findings, however, have generated research aimed at determining the proportion of men who have symptoms usually diagnosed as chronic prostatitis (now known as chronic prostatitis/chronic pelvic pain syndrome) but who actually have IC/BPS.

Among women, the average age at onset of IC/BPS symptoms is the early forties, but the range is from childhood through the early sixties. Risk factors (antecedent features that distinguish cases from controls) primarily have been FSSs. Indeed, the odds of IC/BPS increase with the number of such syndromes present. Surgery was long thought to be a risk factor for IC/BPS, but analyses adjusting for FSSs refuted that association. About one-third of patients appear to have bacterial cystitis at the onset of IC/BPS.

The natural history of IC/BPS is not known. Although studies from urology and urogynecology practices have been interpreted as showing that IC/BPS lasts for the lifetime of the patient, population studies suggest that some individuals with IC/BPS do not consult specialists and may not seek medical care at all, and most prevalence studies do not show an upward trend with age—a pattern that would be expected with incident cases throughout adulthood followed by lifetime persistence of a nonfatal disease. It may be reasonable to conclude that patients in a urology practice represent those with the most severe and recalcitrant IC/BPS.

PATHOLOGY

For the $\leq 10\%$ of IC/BPS patients who have a Hunner's lesion, the term *interstitial cystitis* may indeed describe the histopathologic picture. Most of these patients have substantive inflammation, mast cells, and granulation tissue. However, in the 90% of patients without such lesions, the bladder mucosa and interstitium are relatively normal, with scant inflammation.

ETIOLOGY

Numerous hypotheses about the pathogenesis of IC/BPS have been put forward. It is not surprising that most early theories focused on the bladder. For instance, IC/BPS has been investigated as a chronic bladder infection. Sophisticated technologies have not identified a causative organism in urine or in bladder tissue; however, the patients studied by these methods had IC/BPS of long duration, and the results do not preclude the possibility that infection may trigger the syndrome or may be a feature of early IC/BPS. Other inflammatory factors, including a role for mast cells, have been postulated, but (as noted above) the 90% of patients without a Hunner's ulcer have little bladder inflammation and do not have a prominence of mast cells in bladder tissue. Autoimmunity has been considered, but autoantibodies are low in titer, nonspecific, and thought to be a result rather than a cause of IC/BPS. Increased permeability of the bladder mucosa due to defective epithelium or glycosaminoglycan (the bladder's mucous coating) has been studied frequently, but the findings have been inconclusive.

Investigations of causes outside the bladder have been prompted by the presence of comorbid FSSs. Many patients with FSSs have abnormal pain sensitivity as evidenced by (1) low pain thresholds in body areas unrelated to the diagnosed syndrome, (2) dysfunctional descending neurologic control of tactile signals, and (3) enhanced brain responses to touch in functional neuroimaging studies. Moreover, in patients with IC/BPS, body surfaces remote from the bladder are more sensitive to pain than is the case in individuals without IC/BPS. All

these findings are consistent with upregulation of sensory processing in the brain. Indeed, a prevailing theory is that these concomitantly occurring syndromes have in common an abnormality of brain processing of sensory input. However, antecedence is a critical criterion for causality, and no study has demonstrated that abnormal pain sensitivity precedes either IC/BPS or the FSSs.

CLINICAL PRESENTATION

In some patients, IC/BPS has a gradual onset, and/or the cardinal symptoms of pain, urgency, frequency, and nocturia appear sequentially in no consistent order. Other patients can identify the exact date of onset of IC/BPS symptoms. More than half of the latter patients describe dysuria beginning on that date. As stated, only a minority of IC/BPS patients who obtain medical care soon after symptom onset have uropathogenic bacteria or leukocytes in the urine. These patients—and many others with new-onset IC/BPS—are treated with antibiotics for presumptive bacterial cystitis or, if male, chronic bacterial prostatitis. Persistent or recurring symptoms without bacteriuria eventually prompt a differential diagnosis, and IC/BPS is considered. Traditionally, the diagnosis of IC/BPS has been delayed for years, but recent interest in the disease has shortened this interval.

The pain of IC/BPS includes suprapubic prominence and changes with the voiding cycle. Two-thirds of women with IC/BPS report two or more sites of pain. The most common site (involved in 80% of women) and generally the one with the most severe pain is the suprapubic area. About 35% of female patients have pain in the urethra, 25% in other parts of the vulva, and 30% in nonurogenital areas, mostly the low back and also the anterior or posterior thighs or the buttocks. The pain of IC/BPS is most commonly described as aching, pressing, throbbing, tender, and/or piercing. What may distinguish IC/BPS from other pelvic pain is that, in 95% of patients, bladder filling exacerbates the pain and/or bladder emptying relieves it. Almost as many patients report a puzzling pattern in which certain dietary substances worsen the pain of IC/BPS. Smaller proportions—but still the majority—of patients report that their IC/BPS pain is worsened by menstruation, stress, tight clothing, exercise, and riding in a car as well as during or after vaginal intercourse.

The urethral and vulvar pains of IC/BPS merit special mention. In addition to the descriptive adjectives for IC/BPS mentioned above, these pains commonly are described as burning, stinging, and sharp and as being worsened by touch, tampons, and vaginal intercourse. Patients report that urethral pain increases during urination and generally lessens afterward. These characteristics have commonly resulted in diagnosis of the urethral pain of IC/BPS as chronic urethral syndrome and the vulvar pain as vulvodynia.

In many patients with IC/BPS, there is a link between pain and urinary urgency; that is, two-thirds of patients describe the urge to urinate as a desire to relieve their bladder pain. Only 20% report that the urge stems from a desire to prevent incontinence; indeed, very few patients with IC/BPS are incontinent. As mentioned above, urinary frequency can be severe, with $\sim 85\%$ of patients voiding more than 10 times per 24 h and some as often as 60 times. Voiding continues through the night, and nocturia is common, frequent, and often associated with sleep deprivation.

Beyond these common symptoms of IC/BPS, additional urinary and other symptoms may be present. Among the urinary symptoms are difficulty in starting urine flow, perceptions of difficulty in emptying the bladder, and bladder spasms. Other symptoms include the manifestations of comorbid FSSs as well as symptoms that do not constitute recognized syndromes, such as numbness, muscle spasms, dizziness, ringing in the ears, and blurred vision.

The pain, urgency, and frequency of IC/BPS can be debilitating. Proximity to a bathroom is a continual focus, and patients report difficulties in the workplace, leisure activities, travel, and simply leaving home. Familial and sexual relationships can be strained.

DIAGNOSIS

Traditionally, IC/BPS has been considered a rare condition that is diagnosed by urologists at cystoscopy. However, this disorder is much

more common than once was thought; it is now being considered earlier in its course and is being diagnosed and managed more often by primary care clinicians. Results of physical examination, urinalysis, and urologic procedures are insensitive and/or nonspecific. Thus, diagnosis is based on the presence of appropriate symptoms and the exclusion of diseases with a similar presentation.

Three categories of disorders can be considered in the differential diagnosis of IC/BPS. The first comprises diseases that manifest as bladder pain (see above) or urinary symptoms. Among the latter diseases is *overactive bladder*, a chronic condition of women and men that presents as urgency and frequency and that can be distinguished from IC/BPS by the patient's history: pain is not a feature of overactive bladder, and its urgency arises from the need to avoid incontinence. Endometriosis is a special case: it can be asymptomatic or can cause pelvic pain, dysmenorrhea, and dyspareunia—i.e., types of pain that mimic IC/BPS. Endometrial implants on the bladder (although uncommon) can cause urinary symptoms, and the resulting syndrome can mimic IC/BPS. Even if endometriosis is identified, it is difficult in the absence of bladder implants to determine whether it is causative or incidental to the symptoms of IC/BPS in a specific woman.

The second category of disorders encompasses the FSSs that can accompany IC/BPS. IC/BPS can be misdiagnosed as gynecologic chronic pelvic pain, irritable bowel syndrome, or fibromyalgia. The correct diagnosis may be entertained only when either changes of pain with altered bladder volume or urinary symptoms become more prominent.

The third category involves syndromes that IC/BPS mimics by way of its referred pain, such as vulvodynia and chronic urethral syndrome. Therefore, IC/BPS should be considered in the differential diagnosis of persistent or recurrent "urinary tract infection" (UTI) with sterile urine cultures; overactive bladder with pain; chronic pelvic pain, endometriosis, vulvodynia, or FSSs with urinary symptoms; and "chronic prostatitis." As mentioned above, important clues to the diagnosis of IC/BPS are pain that changes with bladder volume or with certain foods or drinks. Common among these are chilies, chocolate, citrus fruits, tomatoes, alcohol, caffeinated drinks, and carbonated beverages; full lists of common trigger foods are available at the websites cited in the treatment section below.

Cystoscopy under anesthesia formerly was thought to be necessary for the diagnosis of IC/BPS because of its capacity to reveal a Hunner's lesion or—in the 90% of patients without an ulcer—petechial hemorrhages after bladder distention. However, because Hunner's lesions are uncommon in IC/BPS and petechiae are nonspecific, cystoscopy is no longer necessary for diagnosis. Accordingly, the indications for urologic referral have evolved toward the need to rule out other diseases or to administer more advanced treatment.

A typical patient presents to the primary clinician after days, weeks, or months of pain, urgency, frequency, and/or nocturia. The presence of urinary nitrites, leukocytes, or uropathogenic bacteria should prompt treatment for UTI in women and chronic bacterial prostatitis in men. Persistence or recurrence of symptoms in the absence of bacteriuria should prompt a pelvic examination for women, an assay for serum prostate-specific antigen for men, and urine cytology and inclusion of IC/BPS in the differential diagnosis for both sexes.

In the diagnosis of IC/BPS, inquiries about pain, pressure, and discomfort are useful; IC/BPS should be considered if any of these sensations are noted in one or more anterior or posterior sites between the umbilicus and the upper thighs. Nondirective questions about the effect of bladder volume changes include "As your next urination approaches, does this pain get better, get worse, or stay the same?" and "After you urinate, does this pain get better, get worse, or stay the same?" Establishing that the pain is exacerbated by the consumption of certain foods and drinks not only supports the diagnosis of IC/BPS but also serves as the basis for one of the first steps in managing this syndrome. A nondirective way to ask about urgency is to describe it to the patient as a compelling urge to urinate that is difficult to postpone; follow-up questions can determine whether this urge is intended to relieve pain or prevent incontinence. To assess severity and provide quantitative baseline measures, pain and urgency should be estimated

by the patient on a scale of 0–10, with 0 being none and 10 the worst imaginable. Frequency per 24-h period should be determined and nocturia assessed as the number of times per night the patient is awakened by the need to urinate.

About half of patients with IC/BPS have intermittent or persistent microscopic hematuria; this manifestation and the need to exclude bladder stones or cancer require urologic or urogynecologic referral. Initiation of therapy for IC/BPS does not hamper subsequent urologic evaluation.

TREATMENT INTERSTITIAL CYSTITIS/BLADDER PAIN SYNDROME

The goal of therapy is to relieve the symptoms of IC/BPS; the challenge lies in the fact that no treatment is uniformly successful. However, most patients eventually obtain relief, generally with a multifaceted approach. The American Urological Association's guidelines for management of IC/BPS are an excellent resource. The correct strategy is to begin with conservative therapies and proceed to riskier measures only if necessary and under the supervision of a urologist or urogynecologist. Conservative tactics include education, stress reduction, dietary changes, medications, pelvic-floor physical therapy, and treatment of associated FSSs.

Months or even years may have passed since the onset of symptoms, and the patient's life may have been disrupted continually, with repeated medical visits provoking frustration and dismay in both patient and physician. In this circumstance, simply giving a name to the syndrome is beneficial. The physician should discuss the disease, the diagnostic and therapeutic strategies, and the prognosis with the patient and with the spouse and/or other pertinent family members, who may need to be made aware that although IC/BPS has no visible manifestations, the patient is undergoing substantial pain and suffering. This information is particularly important for sexual partners, as exacerbation of pain during and after intercourse is a common feature of IC/BPS. Because stress can worsen IC/BPS symptoms, stress reduction and active measures such as yoga or meditation exercises may be suggested. The Interstitial Cystitis Association (<http://www.ichelp.com>) and the Interstitial Cystitis Network (<http://www.ic-network.com>) can be useful in this educational process.

In constructing a benign diet, some of the many patients who identify particular foods and drinks that exacerbate their symptoms find it useful to exclude all possible offenders and add items back into the diet one at a time to confirm which ones worsen their symptoms. Patients also should experiment with fluid volumes; some find relief with less fluid, others with more.

The pelvic floor is often tender in IC/BPS patients. Two randomized controlled trials showed that weekly physical therapy directed at relaxation of the pelvic muscles yielded significantly more relief than a similar schedule of general body massage. This intervention can be initiated under the direction of a knowledgeable physical therapist who recognizes that the objective is to relax the pelvic floor, not to strengthen it.

Among oral medications, nonsteroidal anti-inflammatory drugs are commonly used but are controversial and often unsuccessful. Two randomized controlled trials showed that amitriptyline can diminish IC/BPS symptoms if an adequate dose (≥ 50 mg per night) can be given. This drug is used not for its antidepressant activity but because of its proven effects on neuropathic pain; however, it is not approved by the U.S. Food and Drug Administration for treatment of IC/BPS. An initial dose of 10 mg at bedtime is increased weekly up to 75 mg (or less if a lower dose adequately relieves symptoms). Side effects can be expected and include dry mouth, weight gain, sedation, and constipation. If this regimen does not control symptoms adequately, pentosan polysulfate, a semisynthetic polysaccharide, can be added at a dose of 100 mg three times a day. Its theoretical effect is to replenish a possibly defective glycosaminoglycan layer over the bladder mucosa; randomized controlled trials suggest only

a modest benefit over placebo. Adverse reactions are uncommon and include gastrointestinal symptoms, headache, and alopecia. Pentosan polysulfate has weak anticoagulant effects and perhaps should be avoided by patients with coagulation abnormalities.

Anecdotal reports suggest that successful therapy for one FSS is accompanied by diminished symptoms of other FSSs. As has been noted here, IC/BPS often is associated with one or several FSSs. Thus, it seems reasonable to hope that, to the extent that accompanying FSSs are treated successfully, the symptoms of IC/BPS will be relieved as well.

If several months of these therapies in combination do not relieve symptoms adequately, the patient should be referred to a urologist or urogynecologist who has access to additional modalities. Cystoscopy under anesthesia allows distention of the bladder with

water, a procedure that provides ~40% of patients with several months of relief and can be repeated. For those few patients with a Hunner's lesion, fulguration may offer relief. Bladder instillation of solutions containing lidocaine or dimethyl sulfoxide can be administered. Physicians experienced in the care of IC/BPS patients have used anticonvulsants, narcotics, and cyclosporine as components of therapy. Pain specialists can be of assistance. Sacral neuromodulation with a temporary percutaneous electrode can be tested and, if effective, can then be performed with an implanted device. In a very small number of patients with recalcitrant symptoms, surgeries, including cystoplasty, partial or total cystectomy, and urinary diversion, may provide relief.

61

Azotemia and Urinary Abnormalities

Julie Lin, Bradley M. Denker

Normal kidney functions occur through numerous cellular processes to maintain body homeostasis. Disturbances in any of these functions can lead to abnormalities that may be detrimental to survival. Clinical manifestations of these disorders depend on the pathophysiology of renal injury and often are identified as a complex of symptoms, abnormal physical findings, and laboratory changes that constitute specific syndromes. These renal syndromes (Table 61-1) may arise from systemic illness or as primary renal disease. Nephrologic syndromes usually consist of several elements that reflect the underlying pathologic processes, typically including

one or more of the following: (1) reduction in glomerular filtration rate (GFR) (azotemia), (2) abnormalities of urine sediment (red blood cells [RBCs], white blood cells [WBCs], casts, and crystals), (3) abnormal excretion of serum proteins (proteinuria), (4) disturbances in urine volume (oliguria, anuria, polyuria), (5) presence of hypertension and/or expanded total body fluid volume (edema), (6) electrolyte abnormalities, and (7) in some syndromes, fever/pain. The specific combination of these findings should permit identification of one of the major nephrologic syndromes (Table 61-1) and allow differential diagnoses to be narrowed so that the appropriate diagnostic and therapeutic course can be determined. All these syndromes and their associated diseases are discussed in more detail in subsequent chapters. This chapter focuses on several aspects of renal abnormalities that are critically important for distinguishing among those processes: (1) reduction in GFR leading to azotemia, (2) alterations of the urinary sediment and/or protein excretion, and (3) abnormalities of urinary volume.

TABLE 61-1 INITIAL CLINICAL AND LABORATORY DATA BASE FOR DEFINING MAJOR SYNDROMES IN NEPHROLOGY

Syndrome	Important Clues to Diagnosis	Common Findings	Chap(s). Discussing Disease-Causing Syndrome
Acute or rapidly progressive renal failure	Anuria Oliguria Documented recent decline in GFR	Hypertension, hematuria Proteinuria, pyuria Casts, edema	334, 338, 340, 343
Acute nephritis	Hematuria, RBC casts Azotemia, oliguria Edema, hypertension	Proteinuria Pyuria Circulatory congestion	338
Chronic renal failure	Azotemia for >3 months Prolonged symptoms or signs of uremia Symptoms or signs of renal osteodystrophy Kidneys reduced in size bilaterally Broad casts in urinary sediment	Proteinuria Casts Polyuria, nocturia Edema, hypertension Electrolyte disorders	333e, 335
Nephrotic syndrome	Proteinuria, with >3.5 g/24 h per 1.73 m ² Hypoalbuminemia Edema Hyperlipidemia	Casts Lipiduria Hypercoagulable state	338
Asymptomatic urinary abnormalities	Hematuria Proteinuria (below nephrotic range) Sterile pyuria, casts		338
Urinary tract infection/pyelonephritis	Bacteriuria, with >10 ⁵ cfu/mL Other infectious agent documented in urine Pyuria, leukocyte casts Frequency, urgency Bladder tenderness, flank tenderness	Hematuria Mild azotemia Mild proteinuria Fever	162
Renal tubule defects	Electrolyte disorders Polyuria, nocturia Renal calcification Large kidneys Renal transport defects	Hematuria "Tubular" proteinuria (<1 g/24 h) Enuresis	339, 340
Hypertension	Systolic/diastolic hypertension	Proteinuria Casts Azotemia	298, 341
Nephrolithiasis	Previous history of stone passage or removal Previous history of stone seen by x-ray Renal colic	Hematuria Pyuria Frequency, urgency	342
Urinary tract obstruction	Azotemia, oliguria, anuria Polyuria, nocturia, urinary retention Slowing of urinary stream Large prostate, large kidneys Flank tenderness, full bladder after voiding	Hematuria Pyuria Enuresis, dysuria	343

Abbreviations: cfu, colony-forming units; GFR, glomerular filtration rate; RBC, red blood cell.

ASSESSMENT OF GFR

Monitoring the GFR is important in both hospital and outpatient settings, and several different methodologies are available. GFR is the primary metric for kidney “function,” and its direct measurement involves administration of a radioactive isotope (such as inulin or iothalamate) that is filtered at the glomerulus into the urinary space but is neither reabsorbed nor secreted throughout the tubule. GFR—i.e., the clearance of inulin or iothalamate in milliliters per minute—is calculated from the rate of appearance of the isotope in the urine over several hours. In most clinical circumstances, direct GFR measurement is not feasible, and the plasma creatinine level is used as a surrogate to estimate GFR. Plasma creatinine (P_{Cr}) is the most widely used marker for GFR, which is related directly to urine creatinine (U_{Cr}) excretion and inversely to P_{Cr} . On the basis of this relationship (with some important caveats, as discussed below), GFR will fall in roughly inverse proportion to the rise in P_{Cr} . Failure to account for GFR reductions in drug dosing can lead to significant morbidity and death from drug toxicities (e.g., digoxin, aminoglycosides). In the outpatient setting, P_{Cr} serves as an estimate for GFR (although much less accurate; see below). In patients with chronic progressive renal disease, there is an approximately linear relationship between $1/P_{Cr}$ (y axis) and time (x axis). The slope of that line will remain constant for an individual;

when values deviate, an investigation for a superimposed acute process (e.g., volume depletion, drug reaction) should be initiated. Signs and symptoms of uremia develop at significantly different levels of P_{Cr} , depending on the patient (size, age, and sex), underlying renal disease, existence of concurrent diseases, and true GFR. Generally, patients do not develop symptomatic uremia until renal insufficiency is severe (GFR <15 mL/min).

A significantly reduced GFR (either acute or chronic) is usually reflected in a rise in P_{Cr} , leading to retention of nitrogenous waste products (defined as azotemia) such as urea. Azotemia may result from reduced renal perfusion, intrinsic renal disease, or postrenal processes (ureteral obstruction; see below and Fig. 61-1). Precise determination of GFR is problematic, as both commonly measured indices (urea and creatinine) have characteristics that affect their accuracy as markers of clearance. Urea clearance may underestimate GFR significantly because of urea reabsorption by the tubule. In contrast, creatinine is derived from muscle metabolism of creatine, and its generation varies little from day to day.

Creatinine clearance (CrCl), an approximation of GFR, is measured from plasma and urinary creatinine excretion rates for a defined period (usually 24 h) and is expressed in milliliters per minute: $CrCl = (U_{vol} \times U_{Cr}) / (P_{Cr} \times T_{min})$. Creatinine is useful for estimating GFR because it is a small, freely filtered solute that is not reabsorbed by the tubules. P_{Cr} levels can increase acutely from

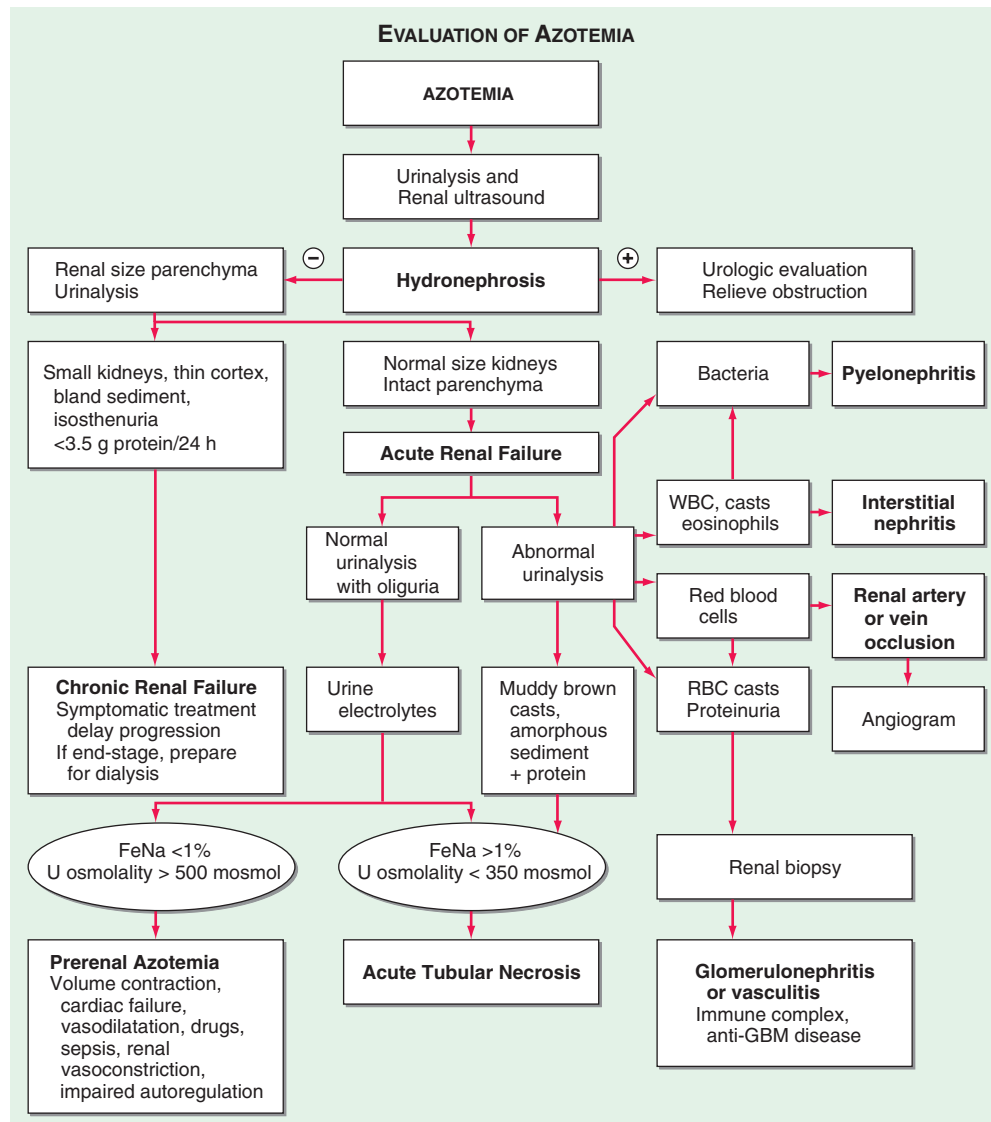


FIGURE 61-1 Approach to the patient with azotemia. FeNa, fractional excretion of sodium; GBM, glomerular basement membrane; RBC, red blood cell; WBC, white blood cell.

dietary ingestion of cooked meat, however, and creatinine can be secreted into the proximal tubule through an organic cation pathway (especially in advanced progressive chronic kidney disease), leading to overestimation of GFR. When a timed collection for CrCl is not available, decisions about drug dosing must be based on P_{Cr} alone. Two formulas are used widely to estimate kidney function from P_{Cr} : (1) Cockcroft-Gault and (2) four-variable MDRD (Modification of Diet in Renal Disease).

$$\text{Cockcroft-Gault: CrCl (mL/min)} = (140 - \text{age (years)} \times \text{weight (kg)} \\ \times [0.85 \text{ if female}]) / (72 \times P_{Cr} \text{ (mg/dL)})$$

$$\text{MDRD: eGFR (mL/min per 1.73 m}^2\text{)} = 186.3 \times P_{Cr} \text{ (e}^{-1.154}\text{)} \times \text{age (e}^{-0.203}\text{)} \\ \times (0.742 \text{ if female}) \times (1.21 \text{ if black}).$$

Numerous websites are available to assist with these calculations (www.kidney.org/professionals/kdoqi/gfr_calculator.cfm). A newer CKD-EPI eGFR, which was developed by pooling several cohorts with and without kidney disease who had data on directly measured GFR, appears to be more accurate:

$$\text{CKD-EPI: eGFR} = 141 \times \min(P_{Cr}/k, 1)^a \times \max(P_{Cr}/k, 1)^{-1.209} \\ \times 0.993^{Age} \times 1.018 \text{ [if female]} \times 1.159 \text{ [if black]},$$

where P_{Cr} is plasma creatinine, k is 0.7 for females and 0.9 for males, a is -0.329 for females and -0.411 for males, \min indicates the minimum of P_{Cr}/k or 1, and \max indicates the maximum of P_{Cr}/k or 1 (<http://www.qxmd.com/renal/Calculate-CKD-EPI-GFR.php>).

There are limitations to all creatinine-based estimates of GFR. Each equation, along with 24-h urine collection for measurement of creatinine clearance, is based on the assumption that the patient is in *steady state*, without daily increases or decreases in P_{Cr} as a result of rapidly changing GFR. The MDRD equation is better correlated with true GFR when the GFR is <60 mL/min per 1.73 m². The gradual loss of muscle from chronic illness, chronic use of glucocorticoids, or malnutrition can mask significant changes in GFR with small or imperceptible changes in P_{Cr} . Cystatin C, a member of the cystatin superfamily of cysteine protease inhibitors, is produced at a relatively constant rate from all nucleated cells. Serum cystatin C has been proposed to be a more sensitive marker of early GFR decline than is P_{Cr} ; however, like serum creatinine, cystatin C is influenced by the patient's age, race, and sex and also is associated with diabetes, smoking, and markers of inflammation.

APPROACH TO THE PATIENT: Azotemia

Once GFR reduction has been established, the physician must decide if it represents acute or chronic renal injury. The clinical situation, history, and laboratory data often make this an easy distinction. However, the laboratory abnormalities characteristic of chronic renal failure, including anemia, hypocalcemia, and hyperphosphatemia, often are present as well in patients presenting with acute renal failure. Radiographic evidence of renal osteodystrophy ([Chap. 335](#)) can be seen only in chronic renal failure but is a very late finding, and these patients are usually undergoing dialysis. The urinalysis and renal ultrasound can facilitate distinguishing acute from chronic renal failure. An approach to the evaluation of azotemic patients is shown in Fig. 61-1. Patients with advanced chronic renal insufficiency often have some proteinuria, nonconcentrated urine (isostenuria; isosmotic with plasma), and small kidneys on ultrasound, characterized by increased echogenicity and cortical thinning. Treatment should be directed toward slowing the progression of renal disease and providing symptomatic relief for edema, acidosis, anemia, and hyperphosphatemia, as discussed in [Chap. 335](#). Acute renal failure ([Chap. 334](#)) can result from processes that affect renal blood flow (prerenal azotemia), intrinsic renal diseases (affecting small vessels, glomeruli, or tubules), or postrenal processes (obstruction of urine flow in ureters, bladder, or urethra) ([Chap. 343](#)).

PRERENAL FAILURE

Decreased renal perfusion accounts for 40–80% of cases of acute renal failure and, if appropriately treated, is readily reversible. The etiologies of prerenal azotemia include any cause of decreased circulating blood volume (gastrointestinal hemorrhage, burns, diarrhea, diuretics), volume sequestration (pancreatitis, peritonitis, rhabdomyolysis), or decreased effective arterial volume (cardiogenic shock, sepsis). Renal perfusion also can be affected by reductions in cardiac output from peripheral vasodilation (sepsis, drugs) or profound renal vasoconstriction (severe heart failure, hepatorenal syndrome, agents such as nonsteroidal anti-inflammatory drugs [NSAIDs]). True or “effective” arterial hypovolemia leads to a fall in mean arterial pressure, which in turn triggers a series of neural and humoral responses, including activation of the sympathetic nervous and renin-angiotensin-aldosterone systems and antidiuretic hormone (ADH) release. GFR is maintained by prostaglandin-mediated relaxation of afferent arterioles and angiotensin II-mediated constriction of efferent arterioles. Once the mean arterial pressure falls below 80 mmHg, GFR declines steeply.

Blockade of prostaglandin production by NSAIDs can result in severe vasoconstriction and acute renal failure. Blocking angiotensin action with angiotensin-converting enzyme (ACE) inhibitors or angiotensin receptor blockers (ARBs) decreases efferent arteriolar tone and in turn decreases glomerular capillary perfusion pressure. Patients taking NSAIDs and/or ACE inhibitors/ARBs are most susceptible to hemodynamically mediated acute renal failure when blood volume is reduced for any reason. Patients with bilateral renal artery stenosis (or stenosis in a solitary kidney) are dependent on efferent arteriolar vasoconstriction for maintenance of glomerular filtration pressure and are particularly susceptible to a precipitous decline in GFR when given ACE inhibitors or ARBs.

Prolonged renal hypoperfusion may lead to acute tubular necrosis (ATN), an intrinsic renal disease that is discussed below. The urinalysis and urinary electrolyte measurements can be useful in distinguishing prerenal azotemia from ATN ([Table 61-2](#)). The urine Na and osmolality of patients with prerenal azotemia can be predicted from the stimulatory actions of norepinephrine, angiotensin II, ADH, and low tubule fluid flow rate. In prerenal conditions, the tubules are intact, leading to a concentrated urine (>500 mosmol), avid Na retention (urine Na concentration, <20 mmol/L; fractional excretion of Na, $<1\%$), and $U_{Cr}/P_{Cr} >40$ ([Table 61-2](#)). The prerenal urine sediment is usually normal or has hyaline and granular casts, whereas the sediment of ATN usually is filled with cellular debris, tubular epithelial casts, and dark (muddy brown) granular casts.

POSTRENAL AZOTEMIA

Urinary tract obstruction accounts for $<5\%$ of cases of acute renal failure but is usually reversible and must be ruled out early in the evaluation ([Fig. 61-1](#)). Since a single kidney is capable of adequate

TABLE 61-2 LABORATORY FINDINGS IN ACUTE RENAL FAILURE

Index	Prerenal Azotemia	Oliguric Acute Renal Failure
BUN/ P_{Cr} ratio	$>20:1$	10–15:1
Urine sodium U_{Na} , meq/L	<20	>40
Urine osmolality, mosmol/L H ₂ O	>500	<350
Fractional excretion of sodium ^a	$<1\%$	$>2\%$
Urine/plasma creatinine U_{Cr}/P_{Cr}	>40	<20
Urinalysis (casts)	None or hyaline/ granular	Muddy brown granular

$$^a \text{FE}_{Na} = \frac{U_{Na} \times P_{Cr} \times 100}{P_{Na} \times U_{Cr}}$$

Abbreviations: BUN, blood urea nitrogen; P_{Cr} , plasma creatinine concentration; P_{Na} , plasma sodium concentration; U_{Cr} , urine creatinine concentration; U_{Na} , urine sodium concentration.

clearance, obstructive acute renal failure requires obstruction at the urethra or bladder outlet, bilateral ureteral obstruction, or unilateral obstruction in a patient with a single functioning kidney. Obstruction is usually diagnosed by the presence of ureteral and renal pelvic dilation on renal ultrasound. However, early in the course of obstruction or if the ureters are unable to dilate (e.g., encasement by pelvic or periureteral tumors), the ultrasound examination may be negative. **The specific urologic conditions that cause obstruction are discussed in Chap. 343.**

INTRINSIC RENAL DISEASE

When prerenal and postrenal azotemia have been excluded as etiologies of renal failure, an intrinsic parenchymal renal disease is present. Intrinsic renal disease can arise from processes involving large renal vessels, intrarenal microvasculature and glomeruli, or the tubulointerstitium. Ischemic and toxic ATN account for ~90% of cases of acute intrinsic renal failure. As outlined in Fig. 61-1, the clinical setting and urinalysis are helpful in separating the possible etiologies. Prerenal azotemia and ATN are part of a spectrum of renal hypoperfusion; evidence of structural tubule injury is present in ATN, whereas prompt reversibility occurs with prerenal azotemia upon restoration of adequate renal perfusion. Thus, ATN often can be distinguished from prerenal azotemia by urinalysis and urine electrolyte composition (Table 61-2 and Fig. 61-1). Ischemic ATN is observed most frequently in patients who have undergone major surgery, trauma, severe hypovolemia, overwhelming sepsis, or extensive burns. Nephrotoxic ATN complicates the administration of many common medications, usually by inducing a combination of intrarenal vasoconstriction, direct tubule toxicity, and/or tubule obstruction. The kidney is vulnerable to toxic injury by virtue of its rich blood supply (25% of cardiac output) and its ability to concentrate and metabolize toxins. A diligent search for hypotension and nephrotoxins usually uncovers the specific etiology of ATN. Discontinuation of nephrotoxins and stabilization of blood pressure often suffice without the need for dialysis while the tubules recover. **An extensive list of potential drugs and toxins implicated in ATN is found in Chap. 334.**

Processes involving the tubules and interstitium can lead to acute kidney injury (AKI), a subtype of acute renal failure. These processes include drug-induced interstitial nephritis (especially by antibiotics, NSAIDs, and diuretics), severe infections (both bacterial and viral), systemic diseases (e.g., systemic lupus erythematosus), and infiltrative disorders (e.g., sarcoidosis, lymphoma, or leukemia). A list of drugs associated with allergic interstitial nephritis is found in **Chap. 340**. Urinalysis usually shows mild to moderate proteinuria, hematuria, and pyuria (~75% of cases) and occasionally WBC casts. The finding of RBC casts in interstitial nephritis has been reported but should prompt a search for glomerular diseases (Fig. 61-1). Occasionally, renal biopsy will be needed to distinguish among these possibilities. The finding of eosinophils in the urine is suggestive of allergic interstitial nephritis or atheroembolic renal disease and is optimally observed with Hansel staining. The absence of eosinophiluria, however, does not exclude these etiologies.

Occlusion of large renal vessels, including arteries and veins, is an uncommon cause of acute renal failure. A significant reduction in GFR by this mechanism suggests bilateral processes or, in a patient with a single functioning kidney, a unilateral process. Renal arteries can be occluded with atheroemboli, thromboemboli, in situ thrombosis, aortic dissection, or vasculitis. Atheroembolic renal failure can occur spontaneously but most often is associated with recent aortic instrumentation. The emboli are cholesterol-rich and lodge in medium and small renal arteries, with a consequent eosinophil-rich inflammatory reaction. Patients with atheroembolic acute renal failure often have a normal urinalysis, but the urine may contain eosinophils and casts. The diagnosis can be confirmed by renal biopsy, but this procedure is often unnecessary when other stigmata of atheroemboli are present (livedo reticularis, distal peripheral infarcts, eosinophilia). Renal artery thrombosis may lead

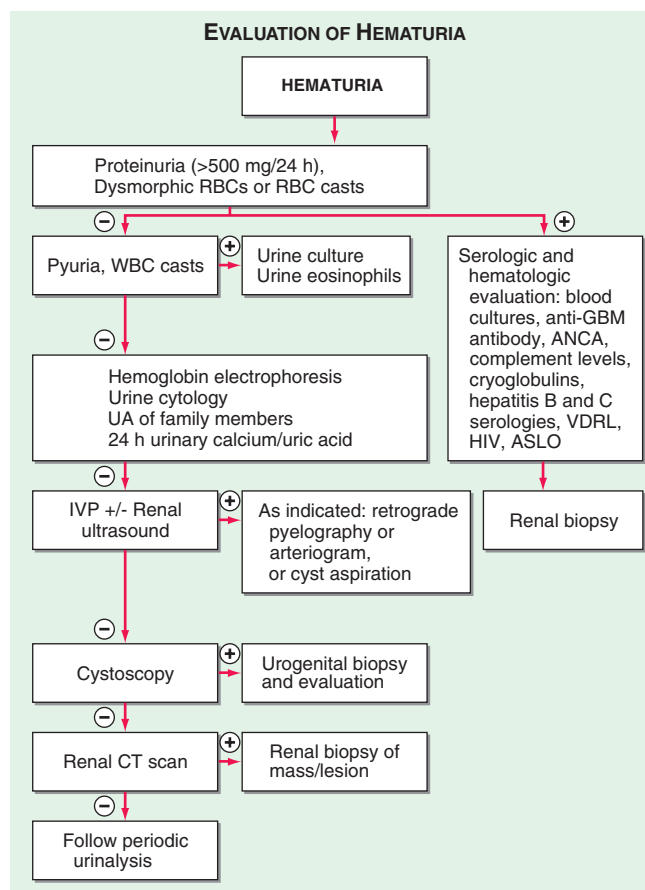


FIGURE 61-2 Approach to the patient with hematuria. ANCA, antineutrophil cytoplasmic antibody; ASLO, antistreptolysin O; CT, computed tomography; GBM, glomerular basement membrane; IVP, intravenous pyelography; RBC, red blood cell; UA, urinalysis; VDRL, Venereal Disease Research Laboratory; WBC, white blood cell.

to mild proteinuria and hematuria, whereas renal vein thrombosis typically induces heavy proteinuria and hematuria. **These vascular complications often require angiography for confirmation and are discussed in Chap. 341.**

Diseases of the glomeruli (glomerulonephritis and vasculitis) and the renal microvasculature (hemolytic-uremic syndromes, thrombotic thrombocytopenic purpura, and malignant hypertension) usually present with various combinations of glomerular injury: proteinuria, hematuria, reduced GFR, and alterations of sodium excretion that lead to hypertension, edema, and circulatory congestion (acute nephritic syndrome). These findings may occur as primary renal diseases or as renal manifestations of systemic diseases. The clinical setting and other laboratory data help distinguish primary renal diseases from systemic diseases. The finding of RBC casts in the urine is an indication for early renal biopsy (Fig. 61-1), as the pathologic pattern has important implications for diagnosis, prognosis, and treatment. Hematuria without RBC casts can also be an indication of glomerular disease; this evaluation is summarized in **Fig. 61-2**. **A detailed discussion of glomerulonephritis and diseases of the microvasculature is found in Chap. 340.**

OLIGURIA AND ANURIA

Oliguria refers to a 24-h urine output <400 mL, and *anuria* is the complete absence of urine formation (<100 mL). Anuria can be caused by total urinary tract obstruction, total renal artery or vein occlusion, and shock (manifested by severe hypotension and intense renal vasoconstriction). Cortical necrosis, ATN, and rapidly progressive glomerulonephritis occasionally cause anuria. Oliguria can accompany acute renal failure of any etiology and carries a

more serious prognosis for renal recovery in all conditions except prerenal azotemia. *Nonoliguria* refers to urine output >400 mL/d in patients with acute or chronic azotemia. With nonoliguric ATN, disturbances of potassium and hydrogen balance are less severe than in oliguric patients, and recovery to normal renal function is usually more rapid.

ABNORMALITIES OF THE URINE

PROTEINURIA

The evaluation of proteinuria is shown schematically in Fig. 61-3 and typically is initiated after detection of proteinuria by dipstick examination. The dipstick measurement detects only albumin and gives false-positive results at pH >7.0 or when the urine is very concentrated or contaminated with blood. Because the dipstick relies on urinary albumin concentration, a very dilute urine may obscure significant proteinuria on dipstick examination. Quantification of urinary albumin on a spot urine sample (ideally from a first morning void) by measurement of an albumin-to-creatinine ratio (ACR) is helpful in approximating a 24-h albumin excretion rate (AER), where ACR (mg/g) \approx AER (mg/24 h). Furthermore, proteinuria that is not predominantly due to albumin will be missed by dipstick screening. This information is particularly important for the detection of Bence-Jones proteins in the urine of patients with multiple myeloma. Tests to measure total urine protein concentration accurately rely on precipitation with sulfosalicylic or trichloroacetic acid (Fig. 61-3).

The magnitude of proteinuria and its composition in the urine depend on the mechanism of renal injury that leads to protein losses. Both charge and size selectivity normally prevent virtually all plasma albumin, globulins, and other high-molecular-weight proteins from crossing the glomerular wall; however, if this barrier is disrupted, plasma proteins may leak into the urine (glomerular proteinuria; Fig. 61-3). Smaller proteins (<20 kDa) are freely filtered but are readily reabsorbed by the proximal tubule. Traditionally, healthy individuals excrete <150 mg/d of total protein and <30 mg/d of albumin. However, even at albuminuria levels <30 mg/d, risk for progression to overt nephropathy or subsequent cardiovascular disease

is increased. The remainder of the protein in the urine is secreted by the tubules (Tamm-Horsfall, IgA, and urokinase) or represents small amounts of filtered β_2 -microglobulin, apoproteins, enzymes, and peptide hormones. Another mechanism of proteinuria entails excessive production of an abnormal protein that exceeds the capacity of the tubule for reabsorption. This situation most commonly occurs with plasma cell dyscrasias, such as multiple myeloma, amyloidosis, and lymphomas, that are associated with monoclonal production of immunoglobulin light chains.

The normal glomerular endothelial cell forms a barrier composed of pores of \sim 100 nm that retain blood cells but offer little impediment to passage of most proteins. The glomerular basement membrane traps most large proteins (>100 kDa), and the foot processes of epithelial cells (podocytes) cover the urinary side of the glomerular basement membrane and produce a series of narrow channels (slit diaphragms) to allow molecular passage of small solutes and water but not proteins. Some glomerular diseases, such as minimal change disease, cause fusion of glomerular epithelial cell foot processes, resulting in predominantly “selective” (Fig. 61-3) loss of albumin. Other glomerular diseases can present with disruption of the basement membrane and slit diaphragms (e.g., by immune complex deposition), resulting in losses of albumin and other plasma proteins. The fusion of foot processes causes increased pressure across the capillary basement membrane, resulting in areas with larger pore sizes (and more severe “nonselective” proteinuria (Fig. 61-3).

When the total daily urinary excretion of protein is >3.5 g, hypoalbuminemia, hyperlipidemia, and edema (nephrotic syndrome; Fig. 61-3) are often present as well. However, total daily urinary protein excretion >3.5 g can occur without the other features of the nephrotic syndrome in a variety of other renal diseases, including diabetes (Fig. 61-3). Plasma cell dyscrasias (multiple myeloma) can be associated with large amounts of excreted light chains in the urine, which may not be detected by dipstick. The light chains are filtered by the glomerulus and overwhelm the reabsorptive capacity of the proximal tubule. Renal failure from these disorders occurs through a variety of mechanisms, including proximal tubule injury, tubule obstruction (cast nephropathy), and light chain deposition (Chap. 340). However, not all excreted light chains are nephrotoxic.

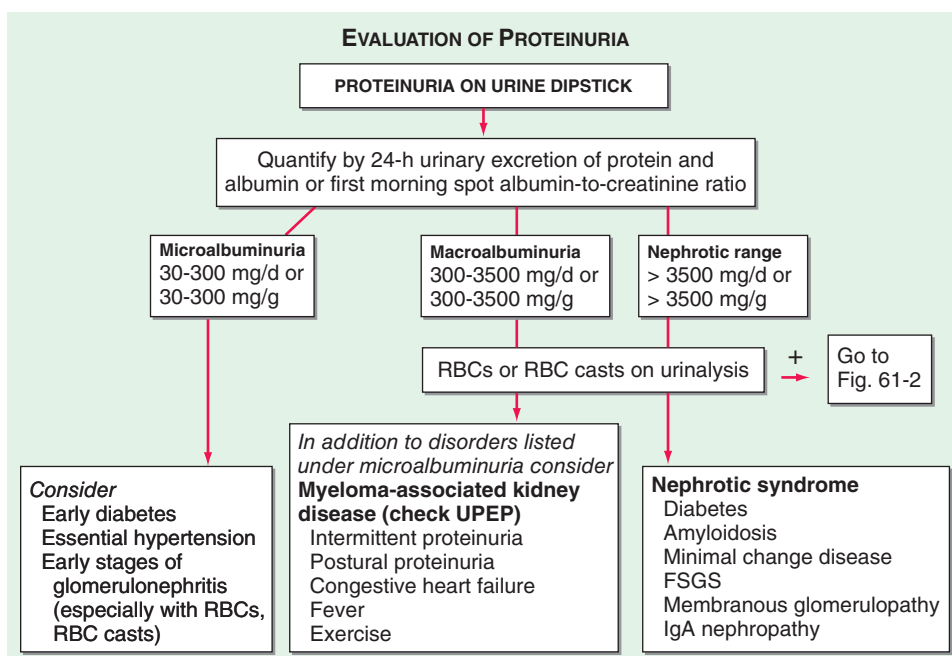


FIGURE 61-3 Approach to the patient with proteinuria. Investigation of proteinuria is often initiated by a positive dipstick on routine urinalysis. Conventional dipsticks detect predominantly albumin and provide a semiquantitative assessment (trace, 1+, 2+, or 3+), which is influenced by urinary concentration as reflected by urine specific gravity (minimum, <1.005; maximum, 1.030). However, more exact determination of proteinuria should employ a spot morning protein/creatinine ratio (mg/g) or a 24-h urine collection (mg/24 h). FSGS, focal segmental glomerulosclerosis; RBC, red blood cell; UPEP, urine protein electrophoresis.

Hypoalbuminemia in nephrotic syndrome occurs through excessive urinary losses and increased proximal tubule catabolism of filtered albumin. Edema forms from renal sodium retention and reduced plasma oncotic pressure, which favors fluid movement from capillaries to interstitium. To compensate for the perceived decrease in effective intravascular volume, activation of the renin-angiotensin system, stimulation of ADH, and activation of the sympathetic nervous system take place, promoting continued renal salt and water reabsorption and progressive edema. Despite these changes, hypertension is uncommon in primary kidney diseases resulting in the nephrotic syndrome (Fig. 61-3 and [Chap. 338](#)). The urinary loss of regulatory proteins and changes in hepatic synthesis contribute to the other manifestations of the nephrotic syndrome. A hypercoagulable state may arise from urinary losses of antithrombin III, reduced serum levels of proteins S and C, hyperfibrinogenemia, and enhanced platelet aggregation. Hypercholesterolemia may be severe and results from increased hepatic lipoprotein synthesis. Loss of immunoglobulins contributes to an increased risk of infection. Many diseases (some listed in Fig. 61-3) and drugs can cause the nephrotic syndrome; a complete list is found in [Chap. 338](#).

HEMATURIA, PYURIA, AND CASTS

Isolated hematuria without proteinuria, other cells, or casts is often indicative of bleeding from the urinary tract. Hematuria is defined as two to five RBCs per high-power field (HPF) and can be detected by dipstick. A false-positive dipstick for hematuria (where no RBCs are seen on urine microscopy) may occur when myoglobinuria is present, often in the setting of rhabdomyolysis. Common causes of isolated hematuria include stones, neoplasms, tuberculosis, trauma, and prostatitis. Gross hematuria with blood clots usually is not an intrinsic renal process; rather, it suggests a postrenal source in the urinary collecting system. Evaluation of patients presenting with microscopic hematuria is outlined in Fig. 61-2. A single urinalysis with hematuria is common and can result from menstruation, viral illness, allergy, exercise, or mild trauma. Persistent or significant hematuria (>3 RBCs/HPF on three urinalyses, a single urinalysis with >100 RBCs, or gross hematuria) is associated with significant renal or urologic lesions in 9.1% of cases. The level of suspicion for urogenital neoplasms in patients with isolated painless hematuria and nondysmorphic RBCs increases with age. Neoplasms are rare in the pediatric population, and isolated hematuria is more likely to be “idiopathic” or associated with a congenital anomaly. Hematuria with pyuria and bacteriuria is typical of infection and should be treated with antibiotics after appropriate cultures. Acute cystitis or urethritis in women can cause gross hematuria. Hypercalciuria and hyperuricosuria are also risk factors for unexplained isolated hematuria in both children and adults. In some of these patients (50–60%), reducing calcium and uric acid excretion through dietary interventions can eliminate the microscopic hematuria.

Isolated microscopic hematuria can be a manifestation of glomerular diseases. The RBCs of glomerular origin are often dysmorphic when examined by phase-contrast microscopy. Irregular shapes of RBCs may also result from pH and osmolarity changes produced along the distal nephron. Observer variability in detecting dysmorphic RBCs is common. The most common etiologies of isolated glomerular hematuria are IgA nephropathy, hereditary nephritis, and thin basement membrane disease. IgA nephropathy and hereditary nephritis can lead to episodic gross hematuria. A family history of renal failure is often present in hereditary nephritis, and patients with thin basement membrane disease often have family members with microscopic hematuria. A renal biopsy is needed for the definitive diagnosis of these disorders, which are discussed in more detail in [Chap. 338](#). Hematuria with dysmorphic RBCs, RBC casts, and protein excretion >500 mg/d is virtually diagnostic of glomerulonephritis. RBC casts form as RBCs that enter the tubule fluid and become trapped in a cylindrical mold of gelled Tamm-Horsfall protein. Even in the absence of azotemia, these patients should undergo serologic evaluation and renal biopsy as outlined in Fig. 61-2.

Isolated pyuria is unusual since inflammatory reactions in the kidney or collecting system also are associated with hematuria. The presence of bacteria suggests infection, and WBC casts with bacteria are indicative of pyelonephritis. WBCs and/or WBC casts also may be seen in acute glomerulonephritis as well as in tubulointerstitial processes such as interstitial nephritis and transplant rejection.

Casts can be seen in chronic renal diseases. Degenerated cellular casts called *waxy casts* or *broad casts* (arising in the dilated tubules that have undergone compensatory hypertrophy in response to reduced renal mass) may be seen in the urine.

ABNORMALITIES OF URINE VOLUME

POLYURIA

By history, it is often difficult for patients to distinguish urinary frequency (often of small volumes) from true polyuria (>3 L/d), and a quantification of volume by 24-h urine collection may be needed ([Fig. 61-4](#)). Polyuria results from two potential mechanisms: (1) excretion of nonabsorbable solutes (such as glucose) or (2) excretion of water (usually from a defect in ADH production or renal responsiveness). To distinguish a solute diuresis from a water diuresis and to determine whether the diuresis is appropriate for the clinical circumstances, urine osmolality is measured. The average person excretes between 600 and 800 mosmol of solutes per day, primarily as urea and electrolytes. If the urine output is >3 L/d and the urine is dilute (<250 mosmol/L), total mosmol excretion is normal and a water diuresis is present. This circumstance could arise from polydipsia, inadequate secretion of vasopressin (*central diabetes insipidus*), or failure of renal tubules to respond to vasopressin (*nephrogenic diabetes insipidus*). If the urine volume is >3 L/d and urine

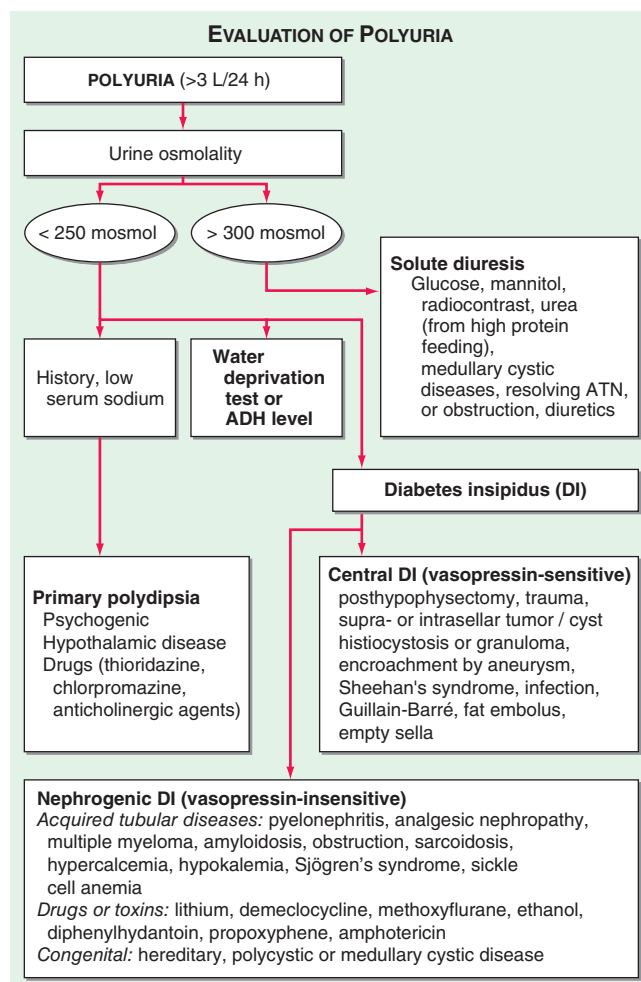


FIGURE 61-4 Approach to the patient with polyuria. ADH, antidiuretic hormone; ATN, acute tubular necrosis.

osmolality is >300 mosmol/L, a solute diuresis is clearly present and a search for the responsible solute(s) is mandatory.

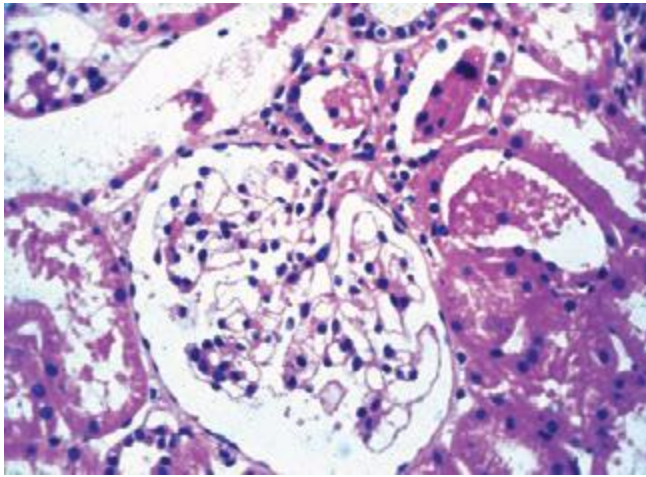
Excessive filtration of a poorly reabsorbed solute such as glucose or mannitol can depress reabsorption of NaCl and water in the proximal tubule and lead to enhanced excretion in the urine. Poorly controlled diabetes mellitus with glucosuria is the most common cause of a solute diuresis, leading to volume depletion and serum hypertonicity. Since the urine sodium concentration is less than that of blood, more water than sodium is lost, causing hypernatremia and hypertonicity. Common iatrogenic solute diuresis occurs in association with mannitol administration, radiocontrast media, and high-protein feedings (enteral or parenteral), leading to increased urea production and excretion. Less commonly, excessive sodium loss may result from cystic renal diseases or Bartter's syndrome or may develop during a tubulointerstitial process (such as resolving ATN). In these so-called salt-wasting disorders, the tubule damage results in direct impairment of sodium reabsorption and indirectly reduces the responsiveness of the tubule to aldosterone. Usually, the sodium losses are mild, and the obligatory urine output is <2 L/d; resolving ATN and postobstructive diuresis are exceptions and may be associated with significant natriuresis and polyuria.

Formation of large volumes of dilute urine is usually due to polydipsic states or diabetes insipidus. Primary polydipsia can result from habit, psychiatric disorders, neurologic lesions, or medications. During deliberate polydipsia, extracellular fluid volume is normal or expanded and plasma vasopressin levels are reduced because serum osmolality tends to be near the lower limits of normal. Urine osmolality is also maximally dilute at 50 mosmol/L.

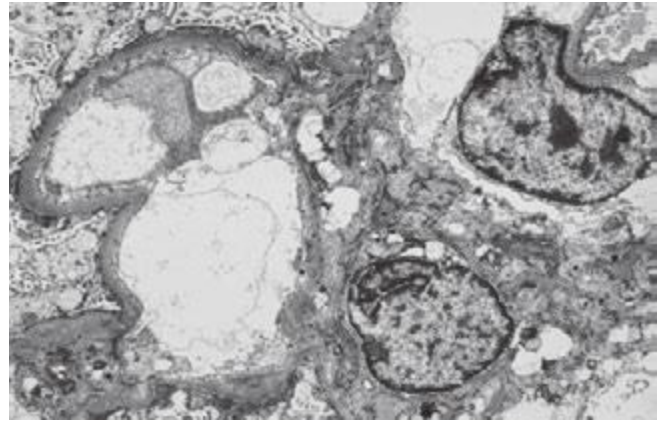
Central diabetes insipidus may be idiopathic in origin or secondary to a variety of conditions, including hypophysectomy, trauma, neoplastic, inflammatory, vascular, or infectious hypothalamic diseases. Idiopathic central diabetes insipidus is associated with selective destruction of the vasopressin-secreting neurons in the supraoptic and paraventricular nuclei and can either be inherited as an autosomal dominant trait or occur spontaneously. Nephrogenic diabetes insipidus can occur in a variety of clinical situations, as summarized in Fig. 61-4.

A plasma vasopressin level is recommended as the best method for distinguishing between central and nephrogenic diabetes insipidus. Alternatively, a water deprivation test plus exogenous vasopressin may distinguish primary polydipsia from central and nephrogenic diabetes insipidus. [For a detailed discussion, see Chap. 404.](#)

Key diagnostic features of selected diseases in renal biopsy are illustrated, with light, immunofluorescence, and electron microscopic images. Common urinalysis findings are also documented.



A



B

FIGURE 62e-1 Minimal-change disease. In minimal-change disease, light microscopy is unremarkable (A), whereas electron microscopy (B) reveals podocyte injury evidenced by complete foot process effacement. (ABF/Vanderbilt Collection.)

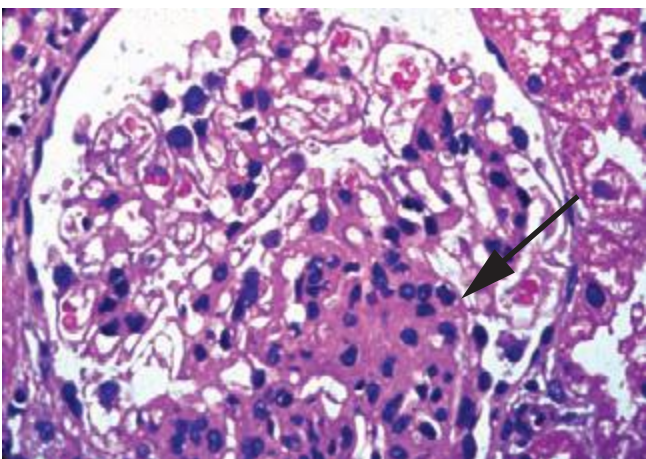


FIGURE 62e-2 Focal segmental glomerulosclerosis (FSGS). There is a well-defined segmental increase in matrix and obliteration of capillary loops (arrow), the sine qua non of segmental sclerosis not otherwise specified (NOS) type. (EGN/UPenn Collection.)

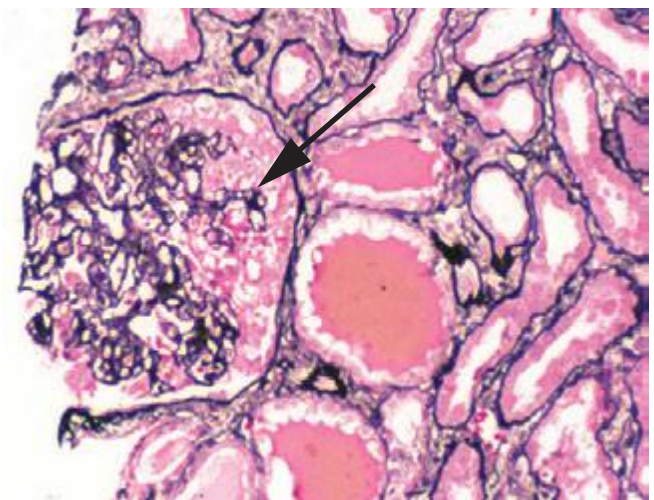


FIGURE 62e-3 Collapsing glomerulopathy. There is segmental collapse (arrow) of the glomerular capillary loops and overlying podocyte hyperplasia. This lesion may be idiopathic or associated with HIV infection and has a particularly poor prognosis. (ABF/Vanderbilt Collection.)

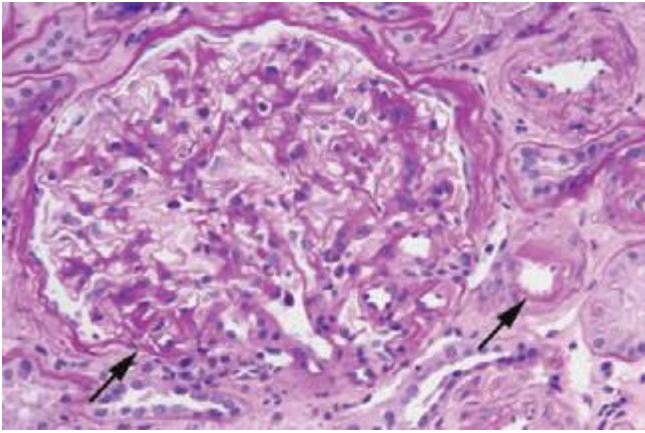


FIGURE 62e-4 Hilar variant of FSGS. There is segmental sclerosis of the glomerular tuft at the vascular pole with associated hyalinosis, also present in the afferent arteriole (*arrows*). This lesion often occurs as a secondary response when nephron mass is lost due to, e.g., scarring from other conditions. Patients usually have less proteinuria and less steroid response than FSGS, NOS type. (ABF/Vanderbilt Collection.)

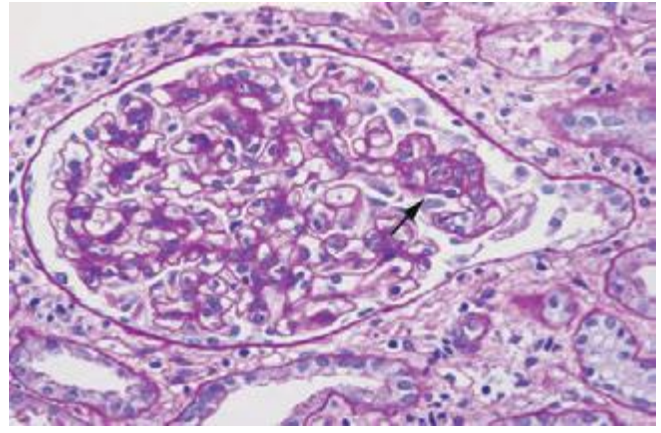
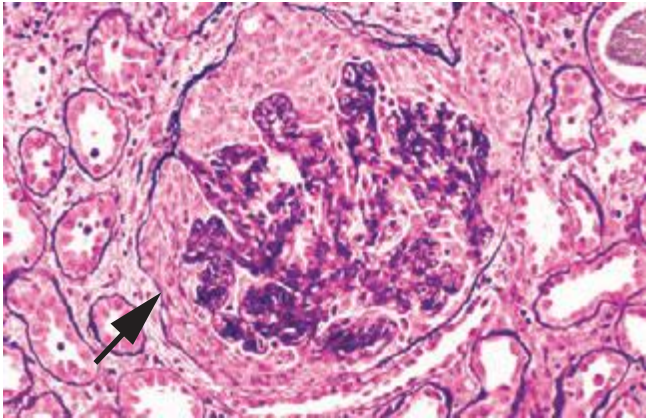
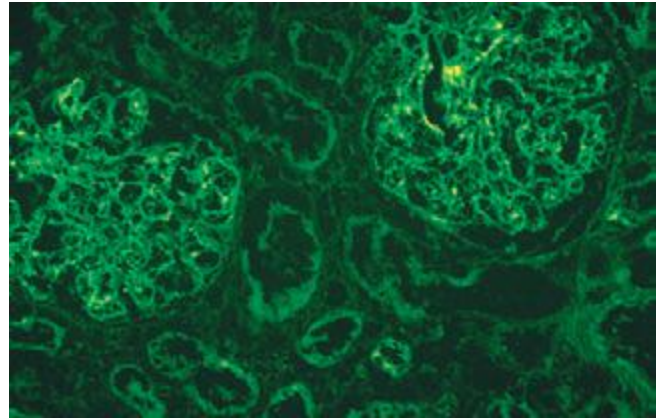


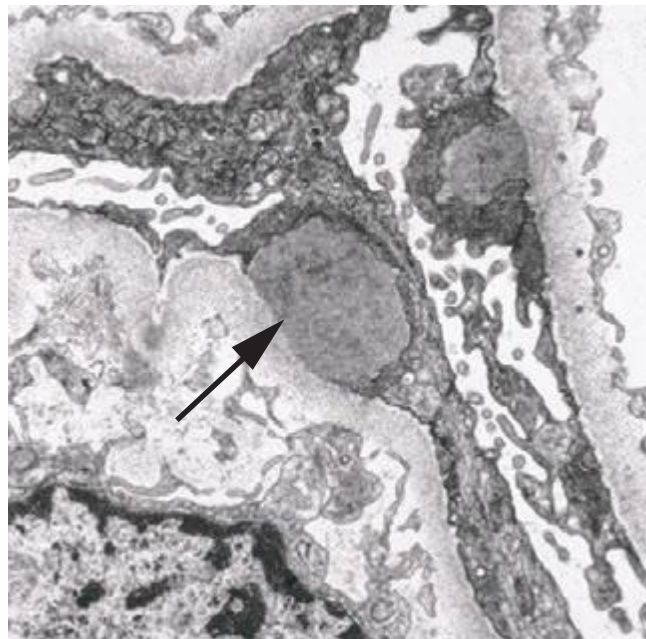
FIGURE 62e-5 Tip lesion variant of FSGS. There is segmental sclerosis of the glomerular capillary loops at the proximal tubular outlet (*arrow*). This lesion has a better prognosis than other types of FSGS. (ABF/Vanderbilt Collection.)



A

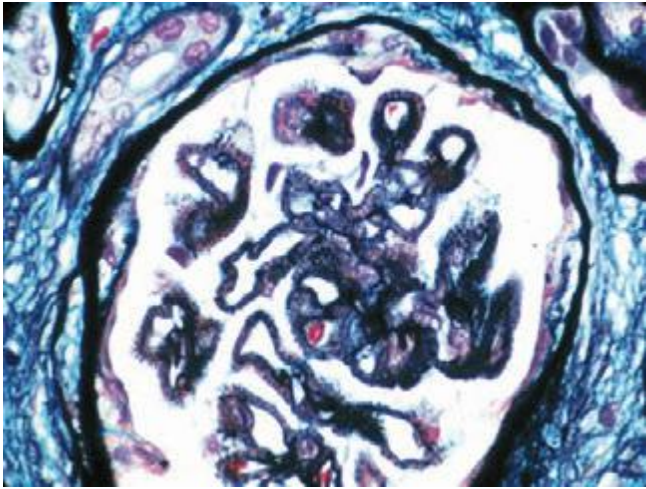


B

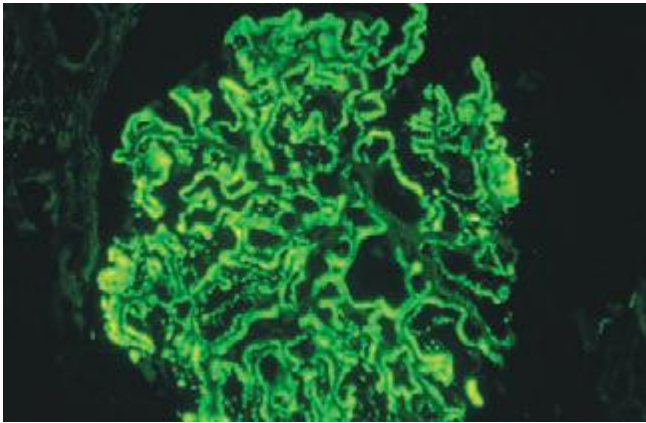


C

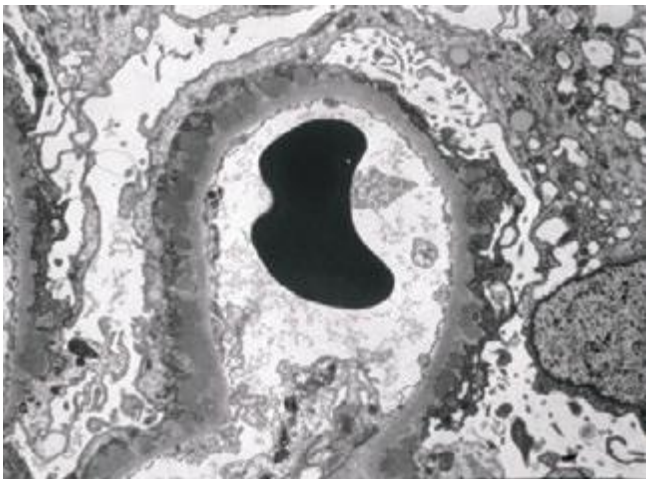
FIGURE 62e-6 Postinfectious (poststreptococcal) glomerulonephritis. The glomerular tuft shows proliferative changes with numerous polymorphonuclear leukocytes (PMNs), with a crescentic reaction (*arrow*) in severe cases (**A**). These deposits localize in the mesangium and along the capillary wall in a subepithelial pattern and stain dominantly for C3 and to a lesser extent for IgG (**B**). Subepithelial hump-shaped deposits are seen by electron microscopy (*arrow*) (**C**). (ABF/Vanderbilt Collection.)



A

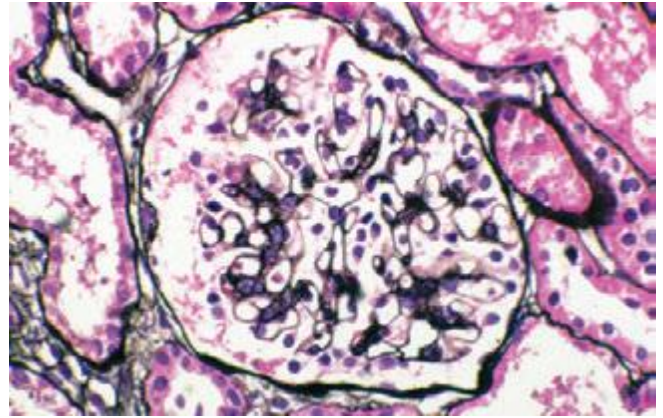


B

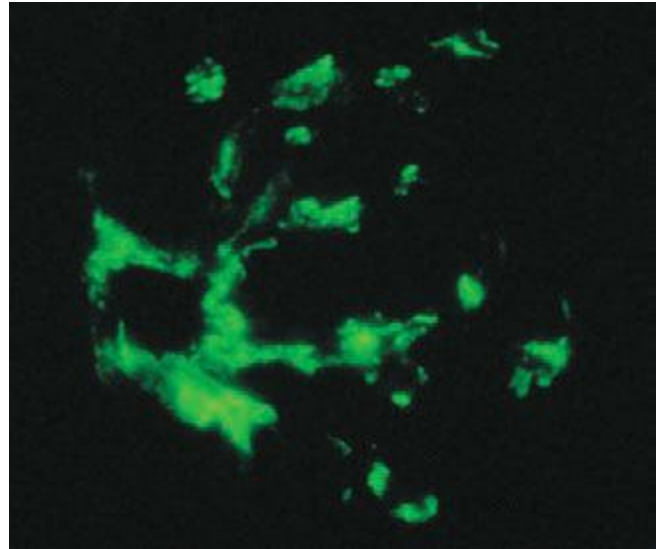


C

FIGURE 62e-7 Membranous glomerulopathy. Membranous glomerulopathy is due to subepithelial deposits, with resulting basement membrane reaction, resulting in the appearance of spike-like projections on silver stain (**A**). The deposits are directly visualized by fluorescent anti-IgG, revealing diffuse granular capillary loop staining (**B**). By electron microscopy, the subepithelial location of the deposits and early surrounding basement membrane reaction is evident, with overlying foot process effacement (**C**). (ABF/Vanderbilt Collection.)



A



B

FIGURE 62e-8 IgA nephropathy. There is variable mesangial expansion due to mesangial deposits, with some cases also showing endocapillary proliferation or segmental sclerosis (**A**). By immunofluorescence, mesangial IgA deposits are evident (**B**). (ABF/Vanderbilt Collection.)

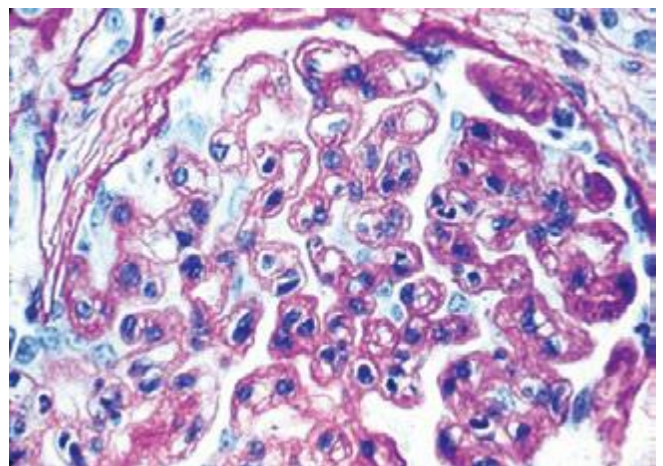


FIGURE 62e-9 Membranoproliferative glomerulonephritis. There is mesangial expansion and endocapillary proliferation with cellular interposition in response to subendothelial deposits, resulting in the "tram-track" of duplication of glomerular basement membrane. (EGN/UPenn Collection.)

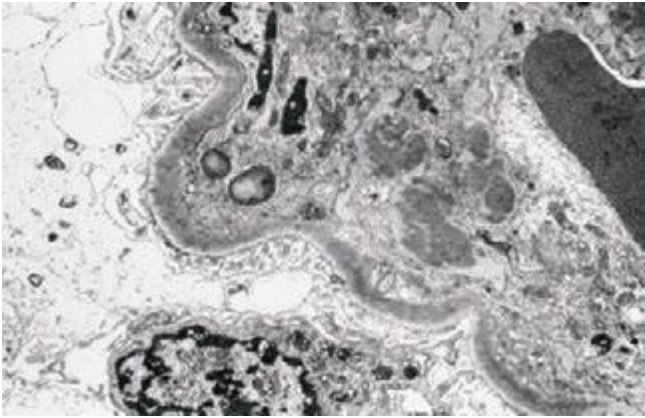


FIGURE 62e-10 Dense deposit disease (membranoproliferative glomerulonephritis type II). By light microscopy, there is a membranoproliferative pattern. By electron microscopy, there is a dense transformation of the glomerular basement membrane with round, globular deposits within the mesangium. By immunofluorescence, only C3 staining is usually present. Dense deposit disease is part of the group of renal diseases called C3 glomerulopathy, related to underlying complement dysregulation. (ABF/Vanderbilt Collection.)

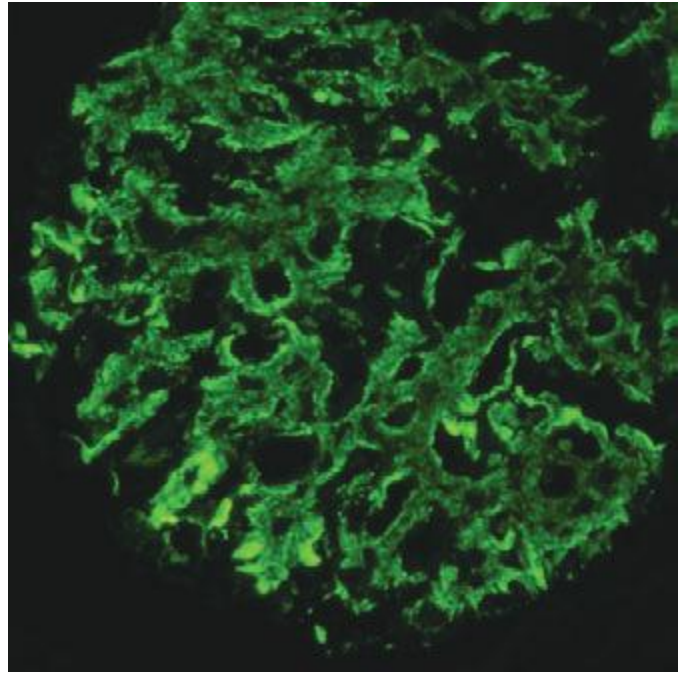


FIGURE 62e-12 C3 glomerulonephritis. By immunofluorescence, only C3 staining is usually present, with occasional minimal immunoglobulin, in an irregular capillary wall and mesangial distribution. (ABF/Vanderbilt Collection.)

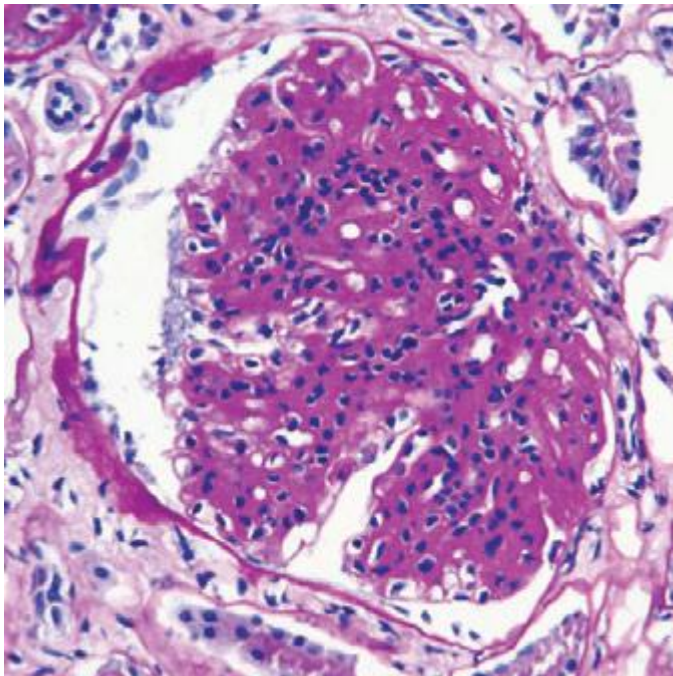


FIGURE 62e-11 C3 glomerulonephritis. By light microscopy, there is a membranoproliferative pattern. C3 glomerulonephritis is part of the group of renal diseases called C3 glomerulopathy, related to underlying complement dysregulation. (ABF/Vanderbilt Collection.)

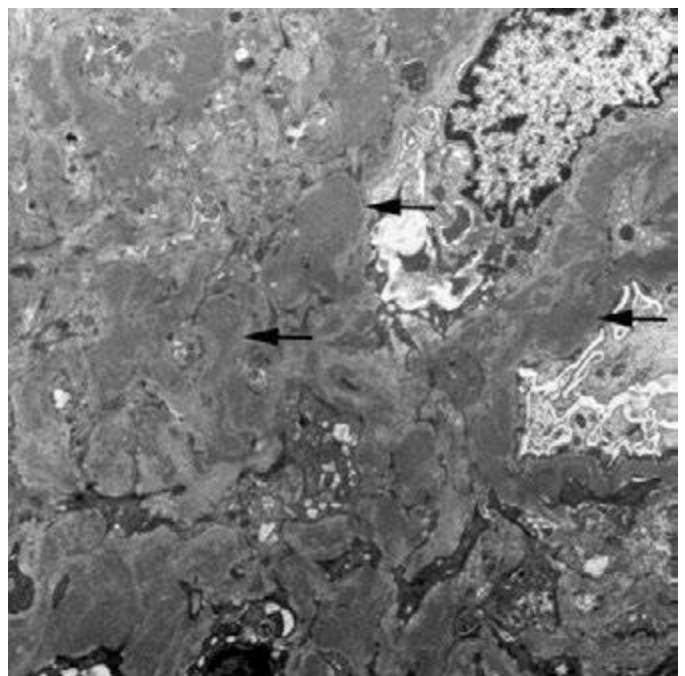


FIGURE 62e-13 C3 glomerulonephritis. By electron microscopy, usual density deposits are present (*arrows*), including mesangial, subendothelial, and occasional hump-type subepithelial deposits. (ABF/Vanderbilt Collection.)

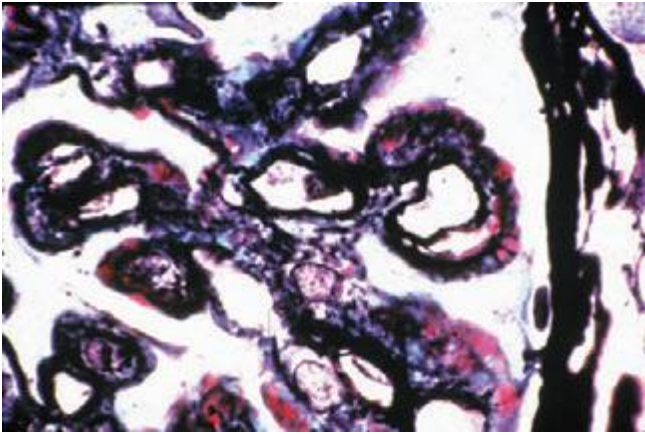


FIGURE 62e-14 Mixed proliferative and membranous glomerulonephritis. This specimen shows pink subepithelial deposits with spike reaction, and the "tram-track" sign of reduplication of glomerular basement membrane, resulting from subendothelial deposits, as may be seen in mixed membranous and proliferative lupus nephritis (International Society of Nephrology [ISN]/Renal Pathology Society [RPS] class V and IV). (EGN/UPenn Collection.)

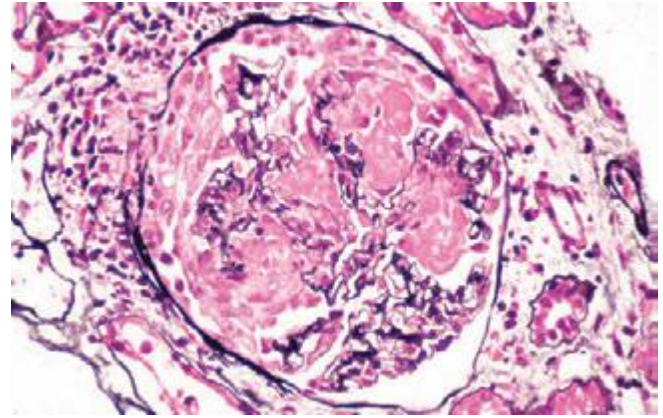
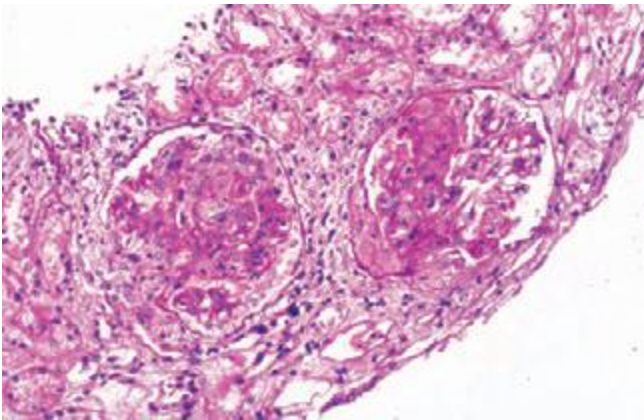
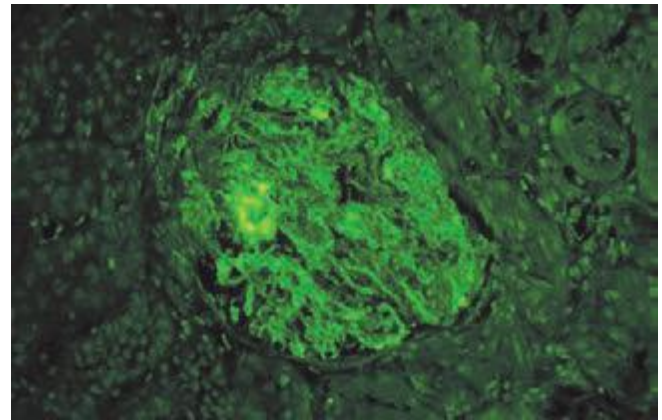


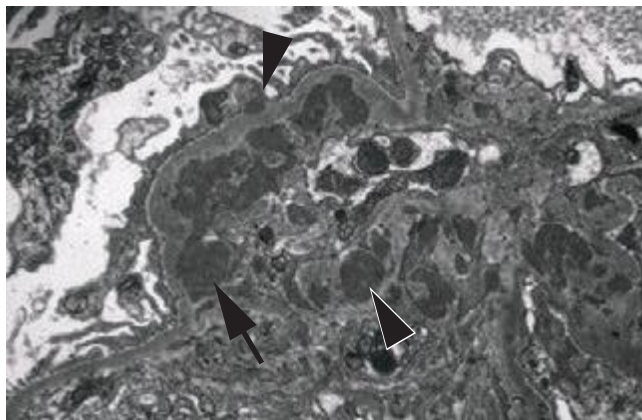
FIGURE 62e-16 Granulomatosis with polyangiitis (Wegener's). This pauci-immune necrotizing crescentic glomerulonephritis shows numerous breaks in the glomerular basement membrane with associated segmental fibrinoid necrosis and a crescent formed by proliferation of the parietal epithelium. Note that the uninvolved segment of the glomerulus (at ~5 o'clock) shows no evidence of proliferation or immune complexes. (ABF/Vanderbilt Collection.)



A

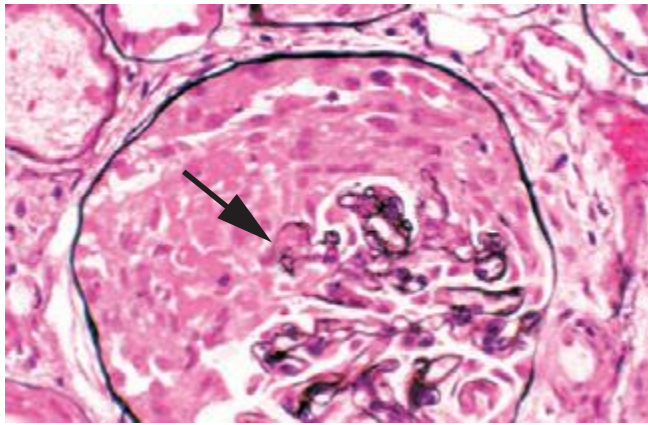


B

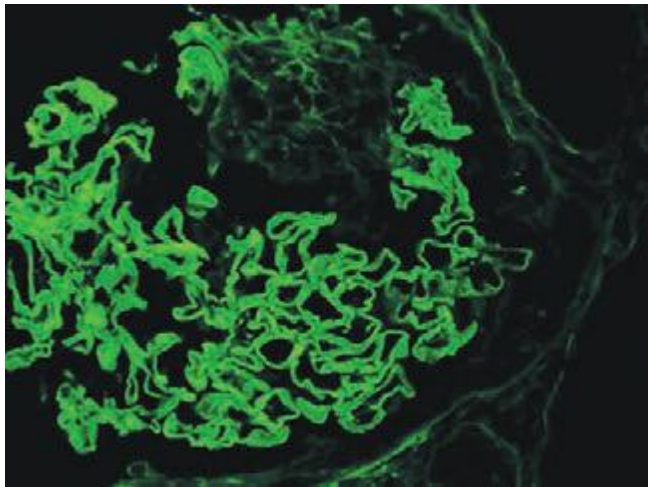


C

FIGURE 62e-15 Lupus nephritis. Proliferative lupus nephritis, ISN/RPS class III (focal) or IV (diffuse), manifests as endocapillary proliferation, which may result in segmental necrosis due to deposits, particularly in the subendothelial area (A). By immunofluorescence, chunky irregular mesangial and capillary loop deposits are evident, with some of the peripheral loop deposits having a smooth, molded outer contour due to their subendothelial location. These deposits typically stain for all three immunoglobulins, IgG, IgA, IgM, and both C3 and C1q (B). By electron microscopy, subendothelial (arrow), mesangial (white rim arrowhead), and rare subepithelial (black arrowhead) dense immune complex deposits are evident, along with extensive foot process effacement (C). (ABF/Vanderbilt Collection.)

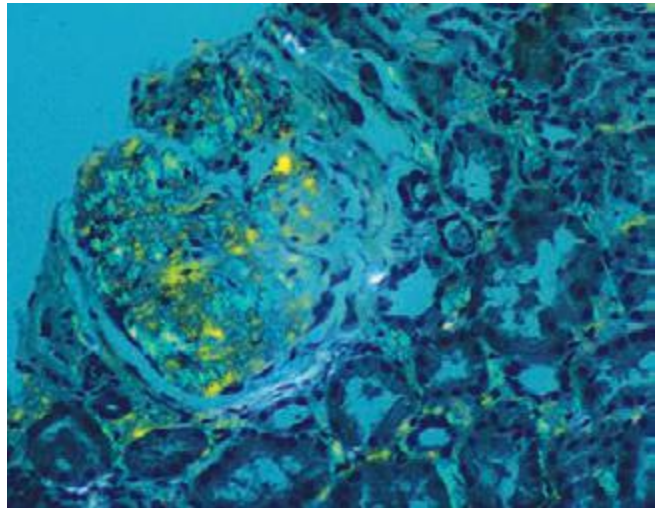


A

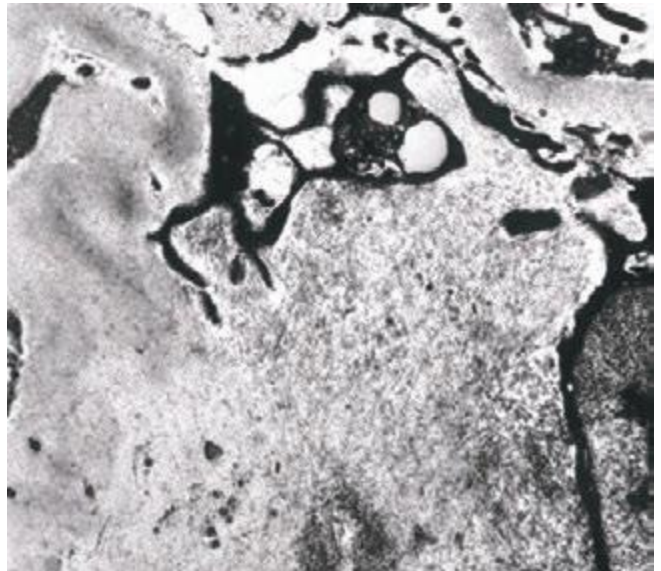


B

FIGURE 62e-17 Anti-glomerular basement membrane antibody-mediated glomerulonephritis. There is segmental necrosis with a break of the glomerular basement membrane (*arrow*) and a cellular crescent (**A**), and immunofluorescence for IgG shows linear staining of the glomerular basement membrane with a small crescent at ~1 o'clock (**B**). (ABF/Vanderbilt Collection.)

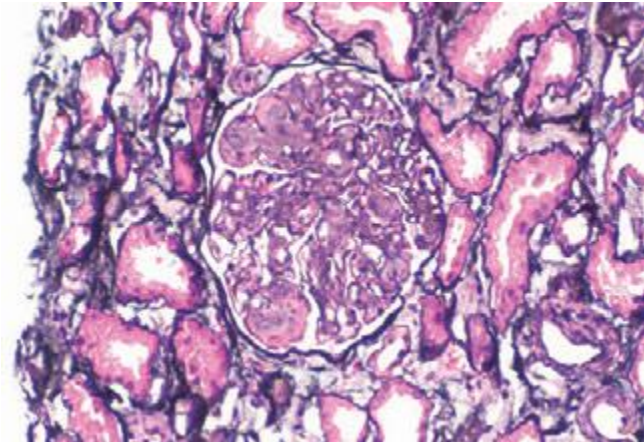


A

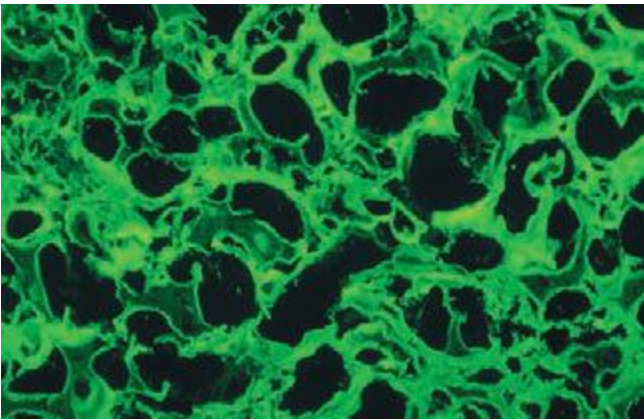


B

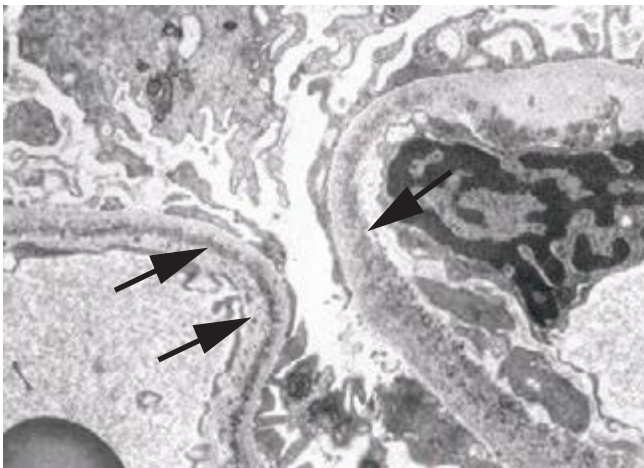
FIGURE 62e-18 Amyloidosis. Amyloidosis shows amorphous, acellular expansion of the mesangium, with material often also infiltrating glomerular basement membranes, vessels, and the interstitium, with apple-green birefringence by polarized Congo red stain (**A**). The deposits are composed of randomly organized 9- to 11-nm fibrils by electron microscopy (**B**). (ABF/Vanderbilt Collection.)



A



B



C

FIGURE 62e-19 Light chain deposition disease. There is mesangial expansion, often nodular by light microscopy (**A**), with immunofluorescence showing monoclonal staining, more commonly with kappa than lambda light chain, of tubules (**B**) and glomerular tufts. By electron microscopy (**C**), the deposits show an amorphous granular appearance and line the inside of the glomerular basement membrane (*arrows*) and are also found along the tubular basement membranes. (ABF/Vanderbilt Collection.)

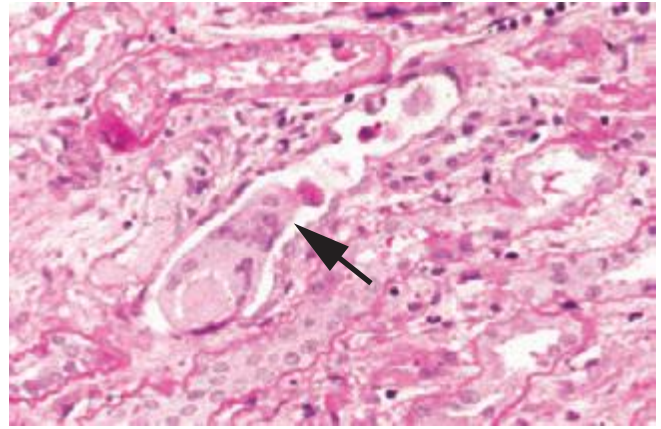
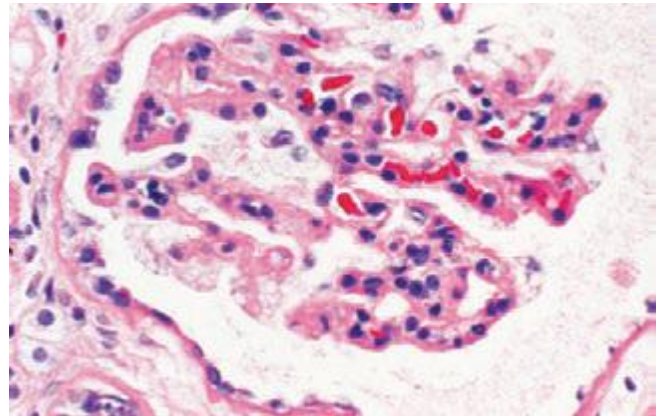
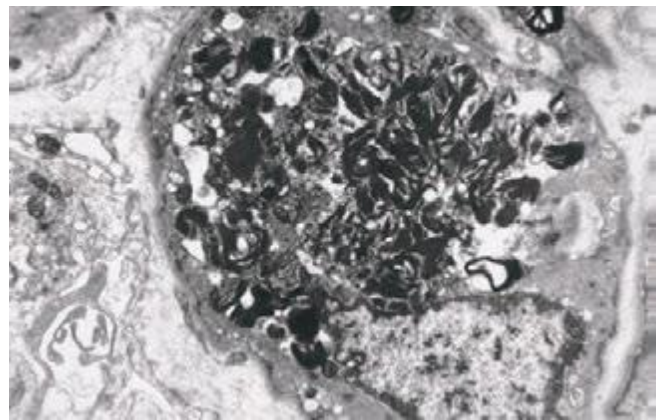


FIGURE 62e-20 Light chain cast nephropathy (myeloma kidney). Monoclonal light chains precipitate in tubules and result in a syncytial giant cell reaction (*arrow*) surrounding the casts and a surrounding chronic interstitial nephritis with tubulointerstitial fibrosis. (ABF/Vanderbilt Collection.)

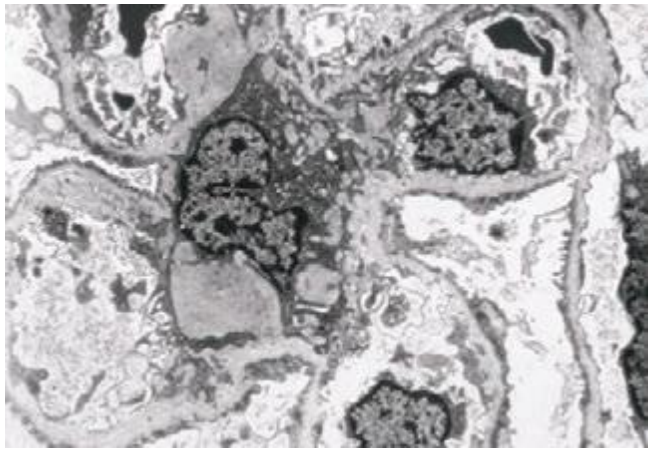


A

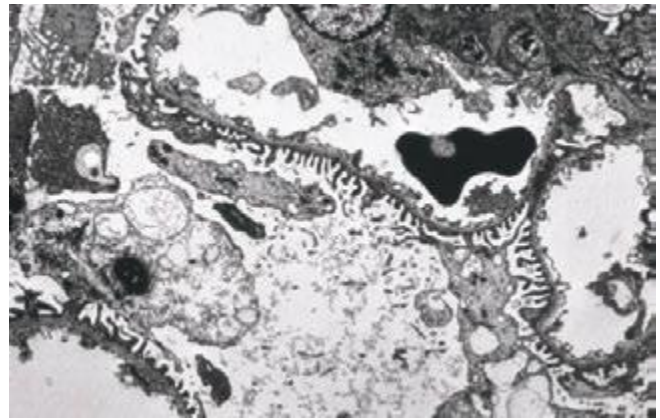


B

FIGURE 62e-21 Fabry's disease. Due to deficiency of α -galactosidase, there is abnormal accumulation of glycolipids, resulting in foamy podocytes by light microscopy (**A**). These deposits can be directly visualized by electron microscopy (**B**), where the glycosphingolipid appears as whorled so-called myeloid bodies, particularly in the podocytes. (ABF/Vanderbilt Collection.)

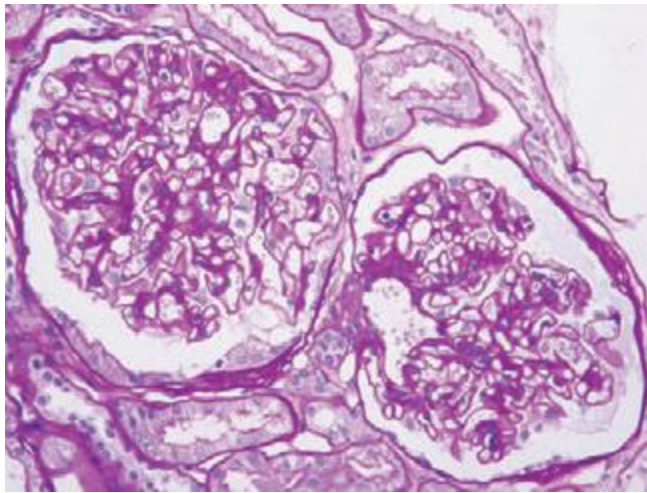


A

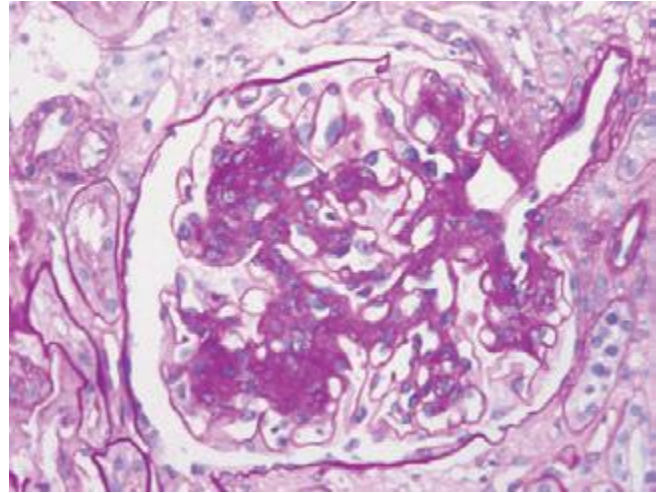


B

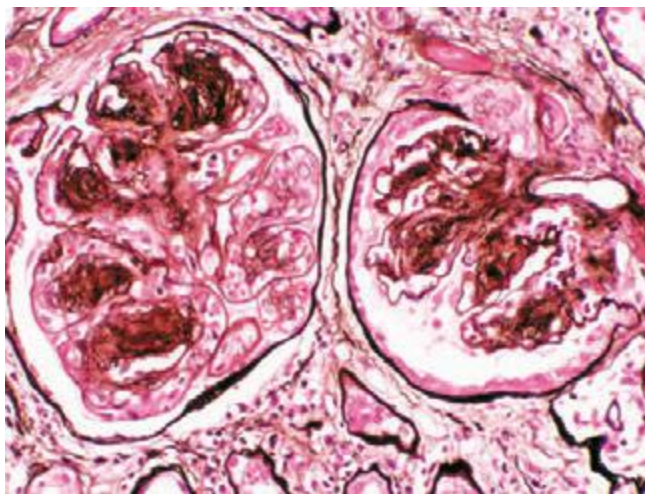
FIGURE 62e-22 Alport's syndrome and thin glomerular basement membrane lesion. In Alport's syndrome, there is irregular thinning alternating with thickened so-called basket-weaving abnormal organization of the glomerular basement membrane (**A**). In benign familial hematuria, or in early cases of Alport's syndrome or female carriers, only extensive thinning of the glomerular basement membrane is seen by electron microscopy (**B**). (ABF/Vanderbilt Collection.)



A

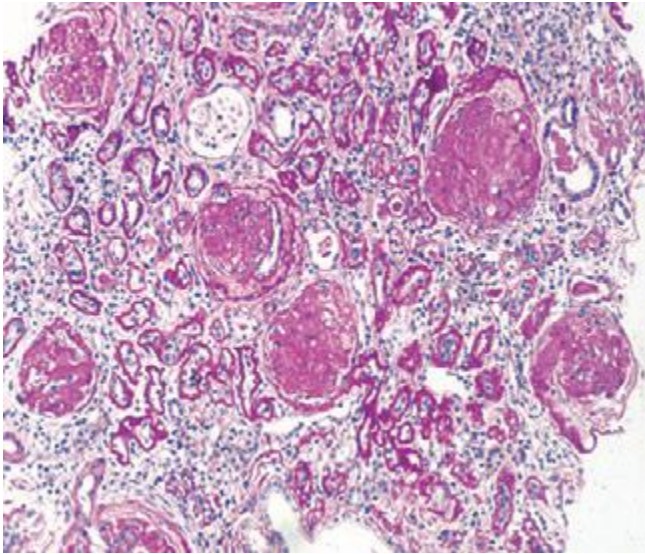


B

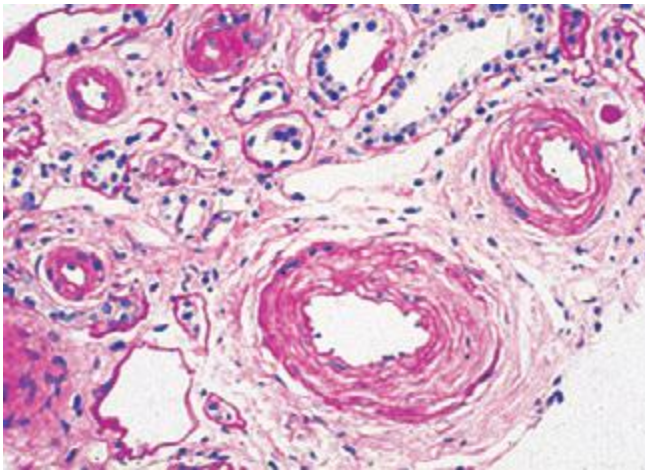


C

FIGURE 62e-23 Diabetic nephropathy. In the earliest stage of diabetic nephropathy, only mild mesangial increase and prominent glomerular basement membranes (confirmed to be thickened by electron microscopy) are present (**A**). In slightly more advanced stages, more marked mesangial expansion with early nodule formation develops, with evident arteriolar hyaline (**B**). In established diabetic nephropathy, there is nodular mesangial expansion, so-called Kimmelstiel-Wilson nodules, with increased mesangial matrix and cellularity, microaneurysm formation in the glomerulus on the left, and prominent glomerular basement membranes without evidence of immune deposits and arteriolar hyalinosis of both afferent and efferent arterioles (**C**). (ABF/Vanderbilt Collection.)



A



B

FIGURE 62e-24 Arterionephrosclerosis. Hypertension-associated injury often manifests extensive global sclerosis of glomeruli, with accompanying and proportional tubulointerstitial fibrosis and pericapsular fibrosis, and there may be segmental sclerosis (**A**). The vessels show disproportionately severe changes of intimal fibrosis, medial hypertrophy, and arteriolar hyaline deposits (**B**). (ABF/Vanderbilt Collection.)

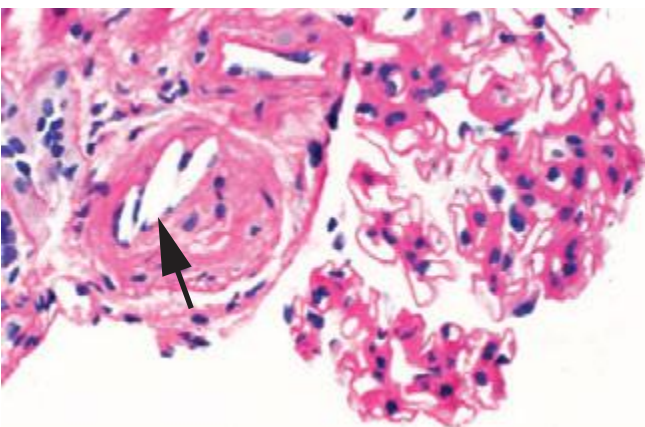


FIGURE 62e-25 Cholesterol emboli. Cholesterol emboli cause cleft-like spaces (*arrow*) where the lipid has been extracted during processing, with smooth outer contours and surrounding fibrotic and mononuclear cell reaction in these arterioles. (ABF/Vanderbilt Collection.)

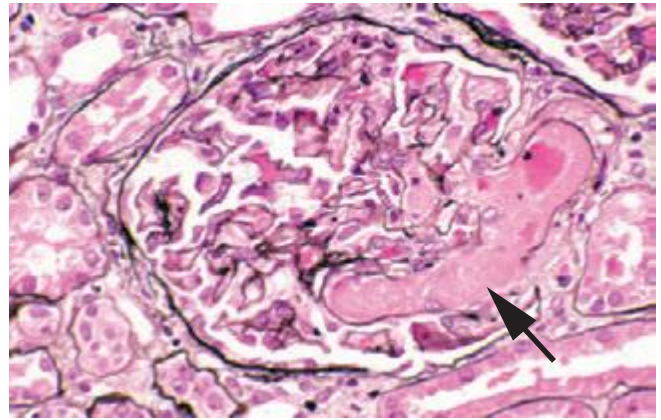
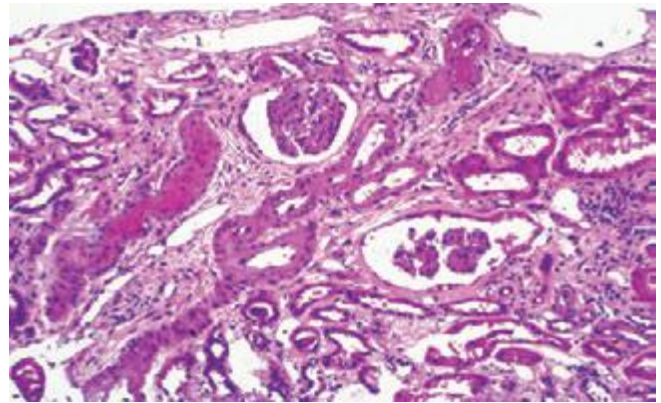
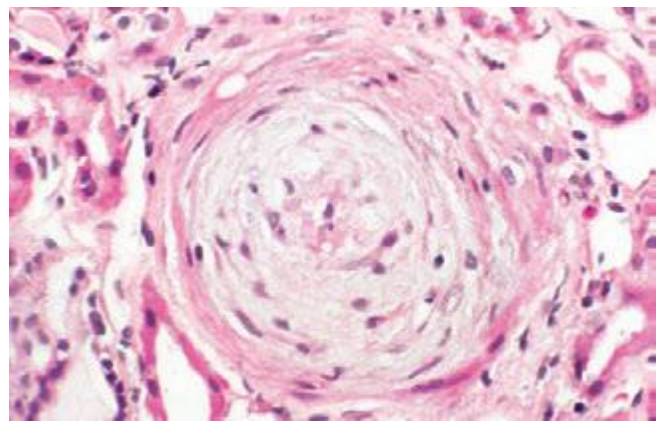


FIGURE 62e-26 Hemolytic-uremic syndrome. There are characteristic intraglomerular fibrin thrombi, with a chunky pink appearance (thrombotic microangiopathy) (*arrow*). The remaining portion of the capillary tuft shows corrugation of the glomerular basement membrane due to ischemia. (ABF/Vanderbilt Collection.)



A



B

FIGURE 62e-27 Progressive systemic sclerosis. Acutely, there is fibrinoid necrosis of interlobular and larger vessels, with intervening normal vessels and ischemic change in the glomeruli (**A**). Chronically, this injury leads to intimal proliferation, the so-called onion-skinning appearance (**B**). (ABF/Vanderbilt Collection.)

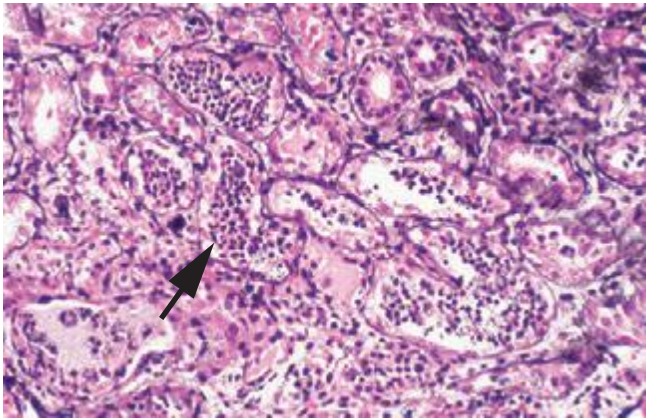


FIGURE 62e-28 Acute pyelonephritis. There are characteristic intra-tubular plugs and casts of PMNs (*arrow*) with inflammation extending into the surrounding interstitium and accompanying tubular injury. (ABF/Vanderbilt Collection.)

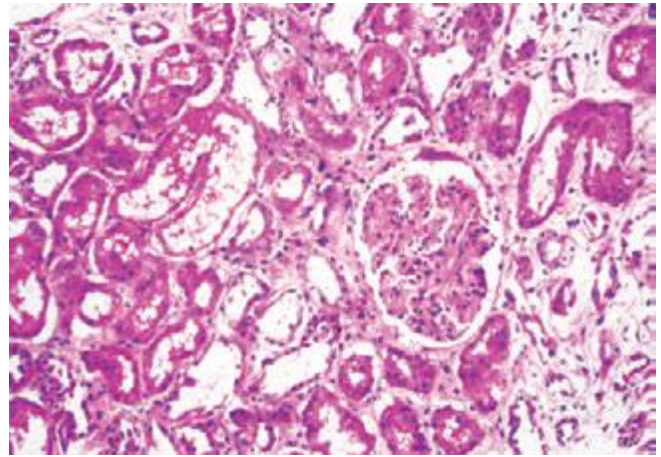
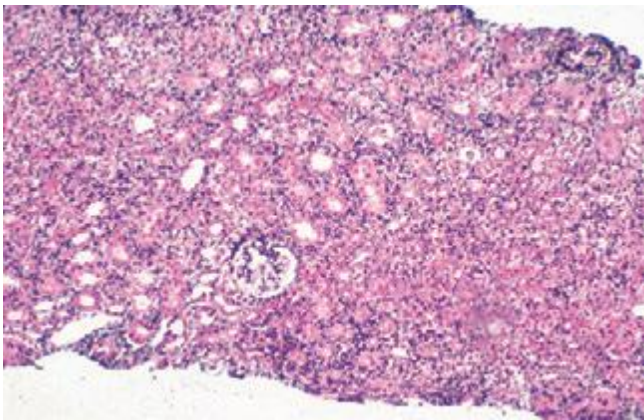
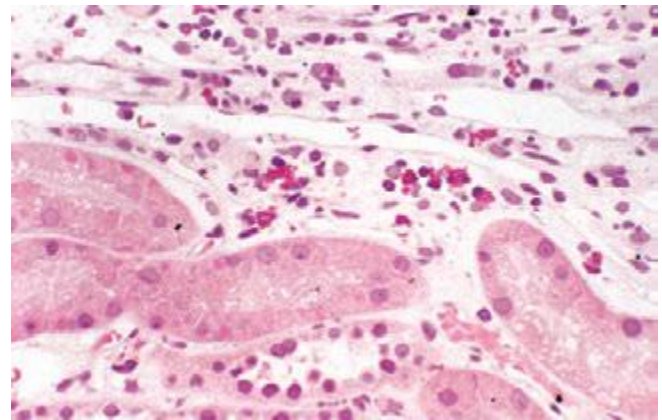


FIGURE 62e-29 Acute tubular injury. There is extensive flattening of the tubular epithelium and loss of the brush border, with mild interstitial edema, characteristic of acute tubular injury due to ischemia. (ABF/Vanderbilt Collection.)

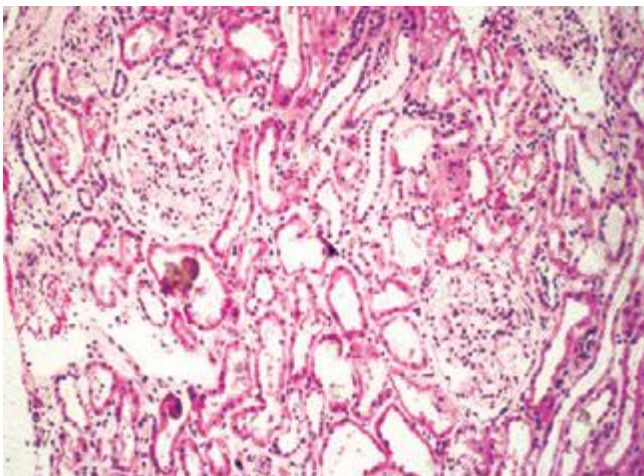


A

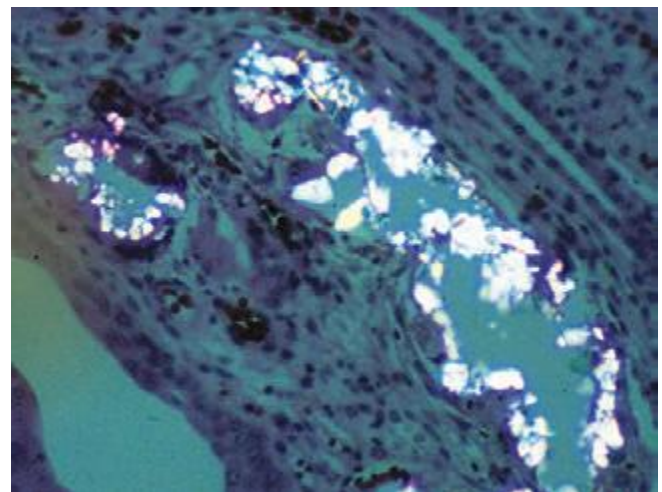


B

FIGURE 62e-30 Acute interstitial nephritis. There is extensive interstitial lymphoplasmocytic infiltrate with mild edema and associated tubular injury (**A**), which is frequently associated with interstitial eosinophils (**B**) when caused by a drug hypersensitivity reaction. (ABF/Vanderbilt Collection.)



A



B

FIGURE 62e-31 Oxalosis. Calcium oxalate crystals have caused extensive tubular injury, with flattening and regeneration of tubular epithelium (**A**). Crystals are well visualized as sheaves when viewed under polarized light (**B**). (ABF/Vanderbilt Collection.)

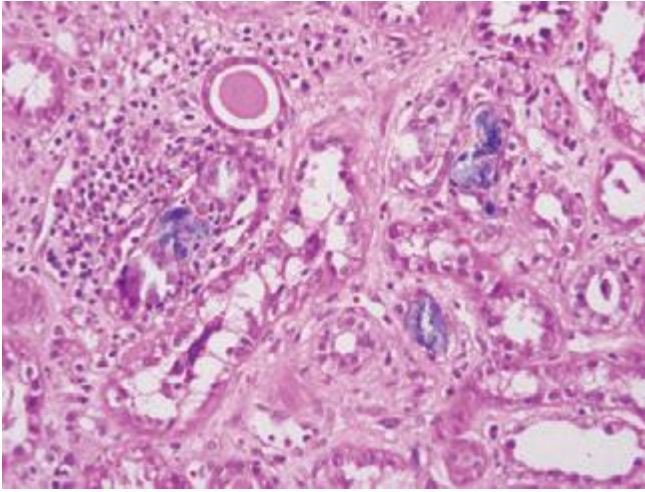


FIGURE 62e-32 Acute phosphate nephropathy. There is extensive acute tubular injury with intratubular nonpolarizable calcium phosphate crystals. (ABF/Vanderbilt Collection.)



FIGURE 62e-35 Coarse granular cast. (ABF/Vanderbilt Collection.)

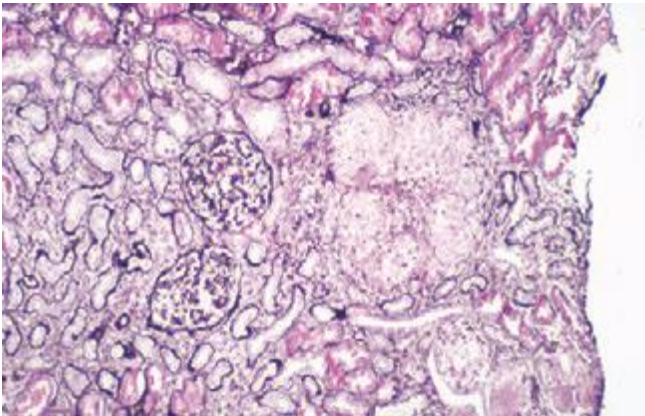


FIGURE 62e-33 Sarcoidosis. There is chronic interstitial nephritis with numerous, confluent, nonnecrotizing granulomas. The glomeruli are unremarkable, but there is moderate tubular atrophy and interstitial fibrosis. (ABF/Vanderbilt Collection.)

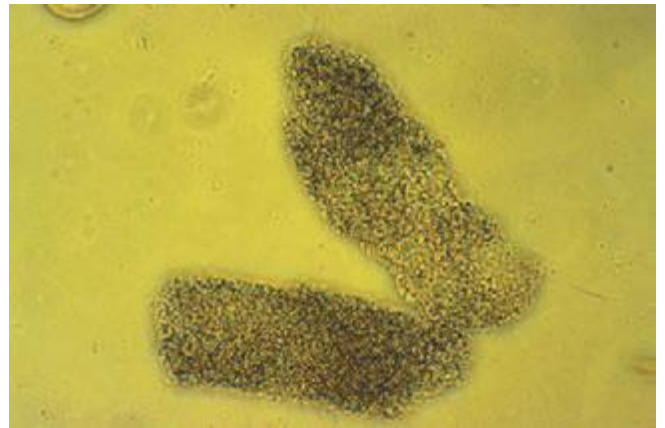


FIGURE 62e-36 Fine granular casts. (ABF/Vanderbilt Collection.)



FIGURE 62e-34 Hyaline cast. (ABF/Vanderbilt Collection.)



FIGURE 62e-37 Red blood cell cast. (ABF/Vanderbilt Collection.)



FIGURE 62e-38 White blood cell cast. (ABF/Vanderbilt Collection.)

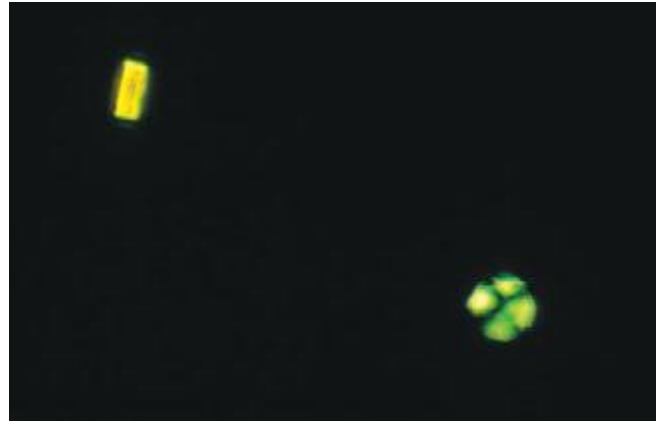


FIGURE 62e-40 "Maltese cross" formation in an oval fat body. (ABF/Vanderbilt Collection.)

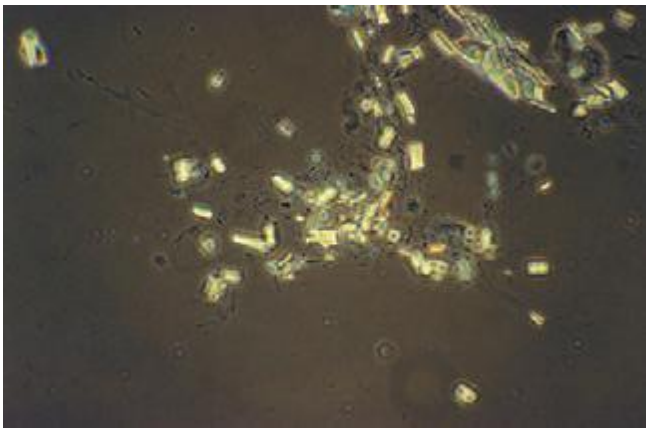


FIGURE 62e-39 Triple phosphate crystals. (ABF/Vanderbilt Collection.)

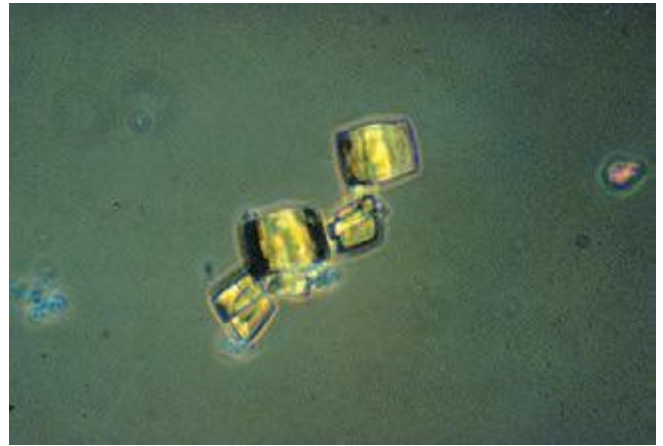


FIGURE 62e-41 Uric acid crystals. (ABF/Vanderbilt Collection.)

SODIUM AND WATER**COMPOSITION OF BODY FLUIDS**

Water is the most abundant constituent in the body, comprising approximately 50% of body weight in women and 60% in men. Total-body water is distributed in two major compartments: 55–75% is intracellular (intracellular fluid [ICF]), and 25–45% is extracellular (extracellular fluid [ECF]). The ECF is further subdivided into intravascular (plasma water) and extravascular (interstitial) spaces in a ratio of 1:3. Fluid movement between the intravascular and interstitial spaces occurs across the capillary wall and is determined by Starling forces, i.e., capillary hydraulic pressure and colloid osmotic pressure. The transcapillary hydraulic pressure gradient exceeds the corresponding oncotic pressure gradient, thereby favoring the movement of plasma ultrafiltrate into the extravascular space. The return of fluid into the intravascular compartment occurs via lymphatic flow.

The solute or particle concentration of a fluid is known as its osmolality, expressed as milliosmoles per kilogram of water (mOsm/kg). Water easily diffuses across most cell membranes to achieve osmotic equilibrium (ECF osmolality = ICF osmolality). Notably, the extracellular and intracellular solute compositions differ considerably owing to the activity of various transporters, channels, and ATP-driven membrane pumps. The major ECF particles are Na^+ and its accompanying anions Cl^- and HCO_3^- , whereas K^+ and organic phosphate esters (ATP, creatine phosphate, and phospholipids) are the predominant ICF osmoles. Solutes that are restricted to the ECF or the ICF determine the “tonicity” or effective osmolality of that compartment. Certain solutes, particularly urea, do not contribute to water shifts across most membranes and are thus known as *ineffective osmoles*.

Water Balance Vasopressin secretion, water ingestion, and renal water transport collaborate to maintain human body fluid osmolality between 280 and 295 mOsm/kg. Vasopressin (AVP) is synthesized in magnocellular neurons within the hypothalamus; the distal axons of these neurons project to the posterior pituitary or neurohypophysis, from which AVP is released into the circulation. A network of central “osmoreceptor” neurons, which includes the AVP-expressing magnocellular neurons themselves, sense circulating osmolality via nonselective, stretch-activated cation channels. These osmoreceptor neurons are activated or inhibited by modest increases and decreases in circulating osmolality, respectively; activation leads to AVP release and thirst.

AVP secretion is stimulated as systemic osmolality increases above a threshold level of ~285 mOsm/kg, above which there is a linear relationship between osmolality and circulating AVP (Fig. 63-1). Thirst and thus water ingestion are also activated at ~285 mOsm/kg, beyond which there is an equivalent linear increase in the perceived intensity of thirst as a function of circulating osmolality. Changes in blood volume and blood pressure are also direct stimuli for AVP release and thirst, albeit with a less sensitive response profile. Of perhaps greater clinical relevance to the pathophysiology of water homeostasis, ECF volume strongly modulates the relationship between circulating osmolality and AVP release, such that hypovolemia reduces the osmotic threshold and increases the slope of the response curve to osmolality; *hypervolemia* has an opposite effect, increasing the osmotic threshold and reducing the slope of the response curve (Fig. 63-1). Notably, AVP has a half-life in the circulation of only 10–20 minutes; thus, changes in ECF volume and/or circulating osmolality can rapidly affect water homeostasis. In addition to volume status, a number of other “nonosmotic” stimuli have potent activating effects on osmosensitive neurons and AVP release, including nausea, intracerebral angiotensin II, serotonin, and multiple drugs.

The excretion or retention of electrolyte-free water by the kidney is modulated by circulating AVP. AVP acts on renal, V_2 -type receptors in

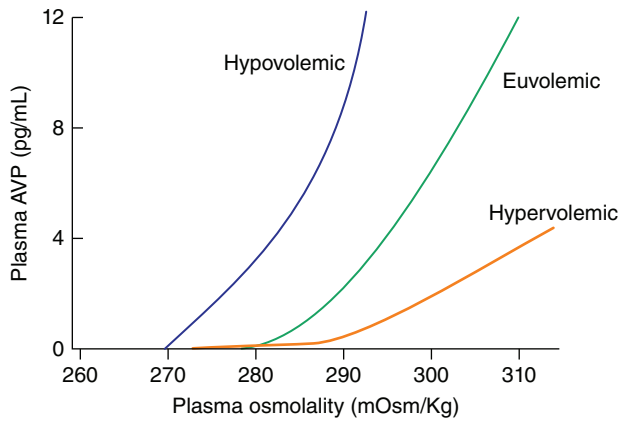


FIGURE 63-1 Circulating levels of vasopressin (AVP) in response to changes in osmolality. Plasma AVP becomes detectable in euvolemic, healthy individuals at a threshold of ~ 285 mOsm/kg, above which there is a linear relationship between osmolality and circulating AVP. The vasopressin response to osmolality is modulated strongly by volume status. The osmotic threshold is thus slightly lower in hypovolemia, with a steeper response curve; hypervolemia reduces the sensitivity of circulating AVP levels to osmolality.

the thick ascending limb of Henle and principal cells of the collecting duct (CD), increasing intracellular levels of cyclic AMP and activating protein kinase A (PKA)-dependent phosphorylation of multiple transport proteins. The AVP- and PKA-dependent activation of Na^+ - Cl^- and K^+ transport by the thick ascending limb of the loop of Henle (TALH) is a key participant in the countercurrent mechanism (Fig. 63-2). The countercurrent mechanism ultimately increases the interstitial osmolality in the inner medulla of the kidney, driving water absorption across the renal CD. However, water, salt, and solute transport by both proximal and distal nephron segments participates in the renal concentrating mechanism (Fig. 63-2). Water transport across apical and basolateral aquaporin-1 water channels in the descending thin limb of the loop of Henle is thus involved, as is passive absorption of Na^+ - Cl^-

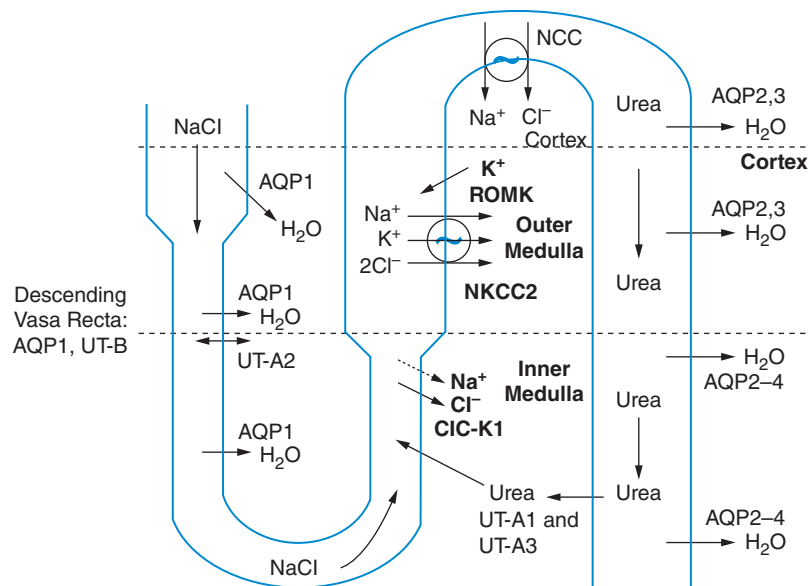


FIGURE 63-2 The renal concentrating mechanism. Water, salt, and solute transport by both proximal and distal nephron segments participates in the renal concentrating mechanism (see text for details). Diagram showing the location of the major transport proteins involved; a loop of Henle is depicted on the left, collecting duct on the right. AQP, aquaporin; CLC-K1, chloride channel; NKCC2, Na-K-2Cl cotransporter; ROMK, renal outer medullary K^+ channel; UT, urea transporter. (Used with permission from JM Sands: *Molecular approaches to urea transporters*. *J Am Soc Nephrol* 13:2795, 2002.)

by the thin ascending limb, via apical and basolateral CLC-K1 chloride channels and paracellular Na^+ transport. Renal urea transport in turn plays important roles in the generation of the medullary osmotic gradient and the ability to excrete solute-free water under conditions of both high and low protein intake (Fig. 63-2).

AVP-induced, PKA-dependent phosphorylation of the aquaporin-2 water channel in principal cells stimulates the insertion of active water channels into the lumen of the CD, resulting in transepithelial water absorption down the medullary osmotic gradient (Fig. 63-3). Under “antidiuretic” conditions, with increased circulating AVP, the kidney reabsorbs water filtered by the glomerulus, equilibrating the osmolality across the CD epithelium to excrete a hypertonic, “concentrated” urine (osmolality of up to 1200 mOsm/kg). In the absence of circulating AVP, insertion of aquaporin-2 channels and water absorption across the CD is essentially abolished, resulting in secretion of a hypotonic, dilute urine (osmolality as low as 30–50 mOsm/kg). Abnormalities in this “final common pathway” are involved in most disorders of water homeostasis, e.g., a reduced or absent insertion of active aquaporin-2 water channels into the membrane of principal cells in diabetes insipidus.

Maintenance of Arterial Circulatory Integrity Sodium is actively pumped out of cells by the Na^+ / K^+ -ATPase membrane pump. In consequence, 85–90% of body Na^+ is extracellular, and the ECF volume (ECFV) is a function of total-body Na^+ content. Arterial perfusion and circulatory integrity are, in turn, determined by renal Na^+ retention or excretion, in addition to the modulation of systemic arterial resistance. Within the kidney, Na^+ is filtered by the glomeruli and then sequentially reabsorbed by the renal tubules. The Na^+ cation is typically reabsorbed with the chloride anion (Cl^-), and, thus, chloride homeostasis also affects the ECFV. On a quantitative level, at a glomerular filtration rate (GFR) of 180 L/d and serum Na^+ of ~ 140 mM, the kidney filters some 25,200 mmol/d of Na^+ . This is equivalent to ~ 1.5 kg of salt, which would occupy roughly 10 times the extracellular space; 99.6% of filtered Na^+ - Cl^- must be reabsorbed to excrete 100 mM per day. Minute changes in renal Na^+ - Cl^- excretion will thus have significant effects on the ECFV, leading to edema syndromes or hypovolemia.

Approximately two-thirds of filtered Na^+ - Cl^- is reabsorbed by the renal proximal tubule, via both paracellular and transcellular mechanisms. The TALH subsequently reabsorbs another 25–30% of filtered Na^+ - Cl^- via the apical, furosemide-sensitive Na^+ - K^+ - 2Cl^- cotransporter. The adjacent aldosterone-sensitive distal nephron, comprising the distal convoluted tubule (DCT), connecting tubule (CNT), and CD, accomplishes the “fine-tuning” of renal Na^+ - Cl^- excretion. The thiazide-sensitive apical Na^+ - Cl^- cotransporter (NCC) reabsorbs 5–10% of filtered Na^+ - Cl^- in the DCT. Principal cells in the CNT and CD reabsorb Na^+ via electrogenic, amiloride-sensitive epithelial Na^+ channels (ENaC); Cl^- ions are primarily reabsorbed by adjacent intercalated cells, via apical Cl^- exchange (Cl^- - OH^- and Cl^- - HCO_3^- exchange, mediated by the SLC26A4 anion exchanger) (Fig. 63-4).

Renal tubular reabsorption of filtered Na^+ - Cl^- is regulated by multiple circulating and paracrine hormones, in addition to the activity of renal nerves. Angiotensin II activates proximal Na^+ - Cl^- reabsorption, as do adrenergic receptors under the influence of renal sympathetic innervation; locally generated dopamine, in contrast, has a *natriuretic* effect. Aldosterone primarily activates Na^+ - Cl^- reabsorption within the aldosterone-sensitive distal nephron. In particular, aldosterone activates the ENaC channel in principal cells, inducing Na^+ absorption and promoting K^+ excretion (Fig. 63-4).

Circulatory integrity is critical for the perfusion and function of vital organs. “Underfilling” of the arterial circulation is sensed by ventricular and vascular pressure receptors, resulting in a neurohumoral activation (increased sympathetic tone, activation of the

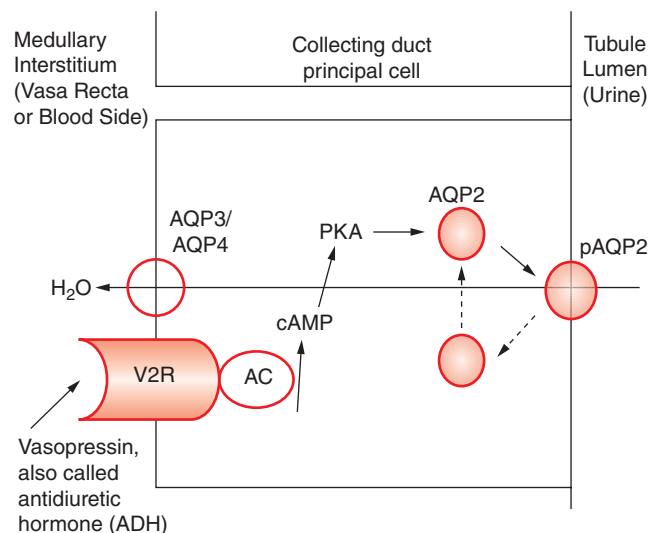


FIGURE 63-3 Vasopressin and the regulation of water permeability in the renal collecting duct. Vasopressin binds to the type 2 vasopressin receptor (V2R) on the basolateral membrane of principal cells, activates adenylyl cyclase (AC), increases intracellular cyclic adenosine monophosphate (cAMP), and stimulates protein kinase A (PKA) activity. Cytoplasmic vesicles carrying aquaporin-2 (AQP) water channel proteins are inserted into the luminal membrane in response to vasopressin, thereby increasing the water permeability of this membrane. When vasopressin stimulation ends, water channels are retrieved by an endocytic process and water permeability returns to its low basal rate. The AQP3 and AQP4 water channels are expressed on the basolateral membrane and complete the transcellular pathway for water reabsorption. pAQP2, phosphorylated aquaporin-2. (From JM Sands, DG Bichet: *Nephrogenic diabetes insipidus*. *Ann Intern Med* 144:186, 2006, with permission.)

renin-angiotensin-aldosterone axis, and increased circulating AVP) that synergistically increases renal Na^+ - Cl^- reabsorption, vascular resistance, and renal water reabsorption. This occurs in the context of decreased cardiac output, as occurs in hypovolemic states, low-output cardiac failure, decreased oncotic pressure, and/or increased capillary permeability. Alternatively, excessive arterial vasodilation results in relative arterial underfilling, leading to neurohumoral activation in the defense of tissue perfusion. These physiologic responses play important roles in many of the disorders discussed in this chapter. In particular, it is important to appreciate that AVP functions in the defense of circulatory integrity, inducing vasoconstriction, increasing sympathetic nervous system tone, increasing renal retention of both water and Na^+ - Cl^- , and modulating the arterial baroreceptor reflex. Most of these responses involve activation of systemic V_{1A} AVP receptors, but concomitant activation of V_2 receptors in the kidney can result in renal water retention and hyponatremia.

HYPOVOLEMIA

Etiology True volume depletion, or hypovolemia, generally refers to a state of combined salt and water loss, leading to contraction of the ECFV. The loss of salt and water may be renal or nonrenal in origin.

RENAL CAUSES Excessive urinary Na^+ - Cl^- and water loss is a feature of several conditions. A high filtered load of endogenous solutes, such as glucose and urea, can impair tubular reabsorption of Na^+ - Cl^- and water, leading to an osmotic diuresis. Exogenous mannitol, often used to decrease intracerebral pressure, is filtered by glomeruli but not reabsorbed by the proximal tubule, thus causing an osmotic diuresis. Pharmacologic diuretics selectively impair Na^+ - Cl^- reabsorption at specific sites along the nephron, leading to increased urinary Na^+ - Cl^- excretion. Other drugs can induce natriuresis as a side effect. For example, acetazolamide can inhibit proximal tubular Na^+ - Cl^- absorption via its inhibition of carbonic anhydrase; other drugs, such as the

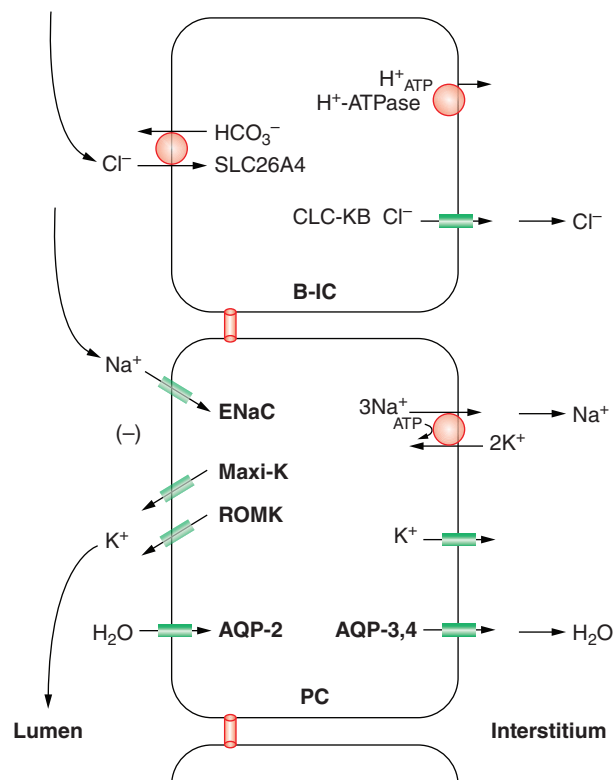


FIGURE 63-4 Sodium, water, and potassium transport in principal cells (PC) and adjacent β -intercalated cells (B-IC). The absorption of Na^+ via the amiloride-sensitive epithelial sodium channel (ENaC) generates a lumen-negative potential difference, which drives K^+ excretion through the apical secretory K^+ channel ROMK (renal outer medullary K^+ channel) and/or the flow-dependent BK channel. Transepithelial Cl^- transport occurs in adjacent β -intercalated cells, via apical Cl^- - HCO_3^- and Cl^- - OH^- exchange (SLC26A4 anion exchanger, also known as pendrin) basolateral CLC chloride channels. Water is absorbed down the osmotic gradient by principal cells, through the apical aquaporin-2 (AQP-2) and basolateral aquaporin-3 and aquaporin-4 (Fig. 63-3).

antibiotics trimethoprim and pentamidine, inhibit distal tubular Na^+ reabsorption through the amiloride-sensitive ENaC channel, leading to urinary Na^+ - Cl^- loss. Hereditary defects in renal transport proteins are also associated with reduced reabsorption of filtered Na^+ - Cl^- and/or water. Alternatively, mineralocorticoid deficiency, mineralocorticoid resistance, or inhibition of the mineralocorticoid receptor (MLR) can reduce Na^+ - Cl^- reabsorption by the aldosterone-sensitive distal nephron. Finally, tubulointerstitial injury, as occurs in interstitial nephritis, acute tubular injury, or obstructive uropathy, can reduce distal tubular Na^+ - Cl^- and/or water absorption.

Excessive excretion of free water, i.e., water without electrolytes, can also lead to hypovolemia. However, the effect on ECFV is usually less marked, given that two-thirds of the water volume is lost from the ICF. Excessive renal water excretion occurs in the setting of decreased circulating AVP or renal resistance to AVP (central and nephrogenic diabetes insipidus, respectively).

EXTRARENAL CAUSES Nonrenal causes of hypovolemia include fluid loss from the gastrointestinal tract, skin, and respiratory system. Accumulations of fluid within specific tissue compartments, typically the interstitium, peritoneum, or gastrointestinal tract, can also cause hypovolemia.

Approximately 9 L of fluid enter the gastrointestinal tract daily, 2 L by ingestion and 7 L by secretion; almost 98% of this volume is absorbed, such that daily fecal fluid loss is only 100–200 mL. Impaired gastrointestinal reabsorption or enhanced secretion of fluid can cause hypovolemia. Because gastric secretions have a low pH (high H^+

concentration), whereas biliary, pancreatic, and intestinal secretions are alkaline (high HCO_3^- concentration), vomiting and diarrhea are often accompanied by metabolic alkalosis and acidosis, respectively.

Evaporation of water from the skin and respiratory tract (so-called “insensible losses”) constitutes the major route for loss of solute-free water, which is typically 500–650 mL/d in healthy adults. This evaporative loss can increase during febrile illness or prolonged heat exposure. Hyperventilation can also increase insensible losses via the respiratory tract, particularly in ventilated patients; the humidity of inspired air is another determining factor. In addition, increased exertion and/or ambient temperature will increase insensible losses via sweat, which is hypotonic to plasma. Profuse sweating without adequate repletion of water and $\text{Na}^+\text{-Cl}^-$ can thus lead to both hypovolemia and hypertonicity. Alternatively, replacement of these insensible losses with a surfeit of free water, without adequate replacement of electrolytes, may lead to hypovolemic hyponatremia.

Excessive fluid accumulation in interstitial and/or peritoneal spaces can also cause intravascular hypovolemia. Increases in vascular permeability and/or a reduction in oncotic pressure (hypoalbuminemia) alter Starling forces, resulting in excessive “third spacing” of the ECFV. This occurs in sepsis syndrome, burns, pancreatitis, nutritional hypoalbuminemia, and peritonitis. Alternatively, distributive hypovolemia can occur due to accumulation of fluid within specific compartments, for example within the bowel lumen in gastrointestinal obstruction or ileus. Hypovolemia can also occur after extracorporeal hemorrhage or after significant hemorrhage into an expandable space, for example, the retroperitoneum.

Diagnostic Evaluation A careful history will usually determine the etiologic cause of hypovolemia. Symptoms of hypovolemia are non-specific and include fatigue, weakness, thirst, and postural dizziness; more severe symptoms and signs include oliguria, cyanosis, abdominal and chest pain, and confusion or obtundation. Associated electrolyte disorders may cause additional symptoms, for example, muscle weakness in patients with hypokalemia. On examination, diminished skin turgor and dry oral mucous membranes are less than ideal markers of a decreased ECFV in adult patients; more reliable signs of hypovolemia include a decreased jugular venous pressure (JVP), orthostatic tachycardia (an increase of >15–20 beats/min upon standing), and orthostatic hypotension (a >10–20 mmHg drop in blood pressure on standing). More severe fluid loss leads to hypovolemic shock, with hypotension, tachycardia, peripheral vasoconstriction, and peripheral hypoperfusion; these patients may exhibit peripheral cyanosis, cold extremities, oliguria, and altered mental status.

Routine chemistries may reveal an increase in blood urea nitrogen (BUN) and creatinine, reflective of a decrease in GFR. Creatinine is the more dependable measure of GFR, because BUN levels may be influenced by an increase in tubular reabsorption (“prerenal azotemia”), an increase in urea generation in catabolic states, hyperalimentation, or gastrointestinal bleeding, and/or a decreased urea generation in decreased protein intake. In hypovolemic shock, liver function tests and cardiac biomarkers may show evidence of hepatic and cardiac ischemia, respectively. Routine chemistries and/or blood gases may reveal evidence of acid-base disorders. For example, bicarbonate loss due to diarrheal illness is a very common cause of metabolic acidosis; alternatively, patients with severe hypovolemic shock may develop lactic acidosis with an elevated anion gap.

The neurohumoral response to hypovolemia stimulates an increase in renal tubular Na^+ and water reabsorption. Therefore, the urine Na^+ concentration is typically <20 mM in nonrenal causes of hypovolemia, with a urine osmolality of >450 mOsm/kg. The reduction in both GFR and distal tubular Na^+ delivery may cause a defect in renal potassium excretion, with an increase in plasma K^+ concentration. Of note, patients with hypovolemia and a hypochloremic alkalosis due to vomiting, diarrhea, or diuretics will typically have a urine Na^+ concentration >20 mM and urine pH of >7.0, due to the increase in filtered HCO_3^- ; the urine Cl^- concentration in this setting is a more accurate indicator of volume status, with a level <25 mM suggestive of hypovolemia. The urine Na^+ concentration is often >20 mM in patients with renal causes

of hypovolemia, such as acute tubular necrosis; similarly, patients with diabetes insipidus will have an inappropriately dilute urine.

TREATMENT HYPOVOLEMIA

The therapeutic goals in hypovolemia are to restore normovolemia and replace ongoing fluid losses. Mild hypovolemia can usually be treated with oral hydration and resumption of a normal maintenance diet. More severe hypovolemia requires intravenous hydration, tailoring the choice of solution to the underlying pathophysiology. Isotonic, “normal” saline (0.9% NaCl, 154 mM Na^+) is the most appropriate resuscitation fluid for normonatremic or hyponatremic patients with severe hypovolemia; colloid solutions such as intravenous albumin are not demonstrably superior for this purpose. Hypertatremic patients should receive a hypotonic solution, 5% dextrose if there has only been water loss (as in diabetes insipidus), or hypotonic saline (1/2 or 1/4 normal saline) if there has been water and $\text{Na}^+\text{-Cl}^-$ loss. Patients with bicarbonate loss and metabolic acidosis, as occur frequently in diarrhea, should receive intravenous bicarbonate, either an isotonic solution (150 meq of $\text{Na}^+\text{-HCO}_3^-$ in 5% dextrose) or a more hypotonic bicarbonate solution in dextrose or dilute saline. Patients with severe hemorrhage or anemia should receive red cell transfusions, without increasing the hematocrit beyond 35%.

SODIUM DISORDERS

Disorders of serum Na^+ concentration are caused by abnormalities in water homeostasis, leading to changes in the relative ratio of Na^+ to body water. Water intake and circulating AVP constitute the two key effectors in the defense of serum osmolality; defects in one or both of these two defense mechanisms cause most cases of hyponatremia and hypernatremia. In contrast, abnormalities in sodium homeostasis per se lead to a deficit or surplus of whole-body $\text{Na}^+\text{-Cl}^-$ content, a key determinant of the ECFV and circulatory integrity. Notably, volume status also modulates the release of AVP by the posterior pituitary, such that hypovolemia is associated with higher circulating levels of the hormone at each level of serum osmolality. Similarly, in “hypervolemic” causes of arterial underfilling, e.g., heart failure and cirrhosis, the associated neurohumoral activation is associated with an increase in circulating AVP, leading to water retention and hyponatremia. Therefore, a key concept in sodium disorders is that the absolute plasma Na^+ concentration tells one nothing about the volume status of a given patient, which furthermore must be taken into account in the diagnostic and therapeutic approach.

HYPONATREMIA

Hyponatremia, which is defined as a plasma Na^+ concentration <135 mM, is a very common disorder, occurring in up to 22% of hospitalized patients. This disorder is almost always the result of an increase in circulating AVP and/or increased renal sensitivity to AVP, combined with an intake of free water; a notable exception is hyponatremia due to low solute intake (see below). The underlying pathophysiology for the exaggerated or “inappropriate” AVP response differs in patients with hyponatremia as a function of their ECFV. Hyponatremia is thus subdivided diagnostically into three groups, depending on clinical history and volume status, i.e., “hypovolemic,” “euvoletic,” and “hypervolemic” (Fig. 63-5).

Hypovolemic Hyponatremia Hypovolemia causes a marked neurohumoral activation, increasing circulating levels of AVP. The increase in circulating AVP helps preserve blood pressure via vascular and baroreceptor V_{1A} receptors and increases water reabsorption via renal V_2 receptors; activation of V_2 receptors can lead to hyponatremia in the setting of increased free water intake. Nonrenal causes of hypovolemic hyponatremia include GI loss (e.g., vomiting, diarrhea, tube drainage) and insensible loss (sweating, burns) of $\text{Na}^+\text{-Cl}^-$ and water, in the absence of adequate oral replacement; urine Na^+ concentration is typically <20 mM. Notably, these patients may be clinically classified as euvoletic, with only the reduced urinary Na^+ concentration to

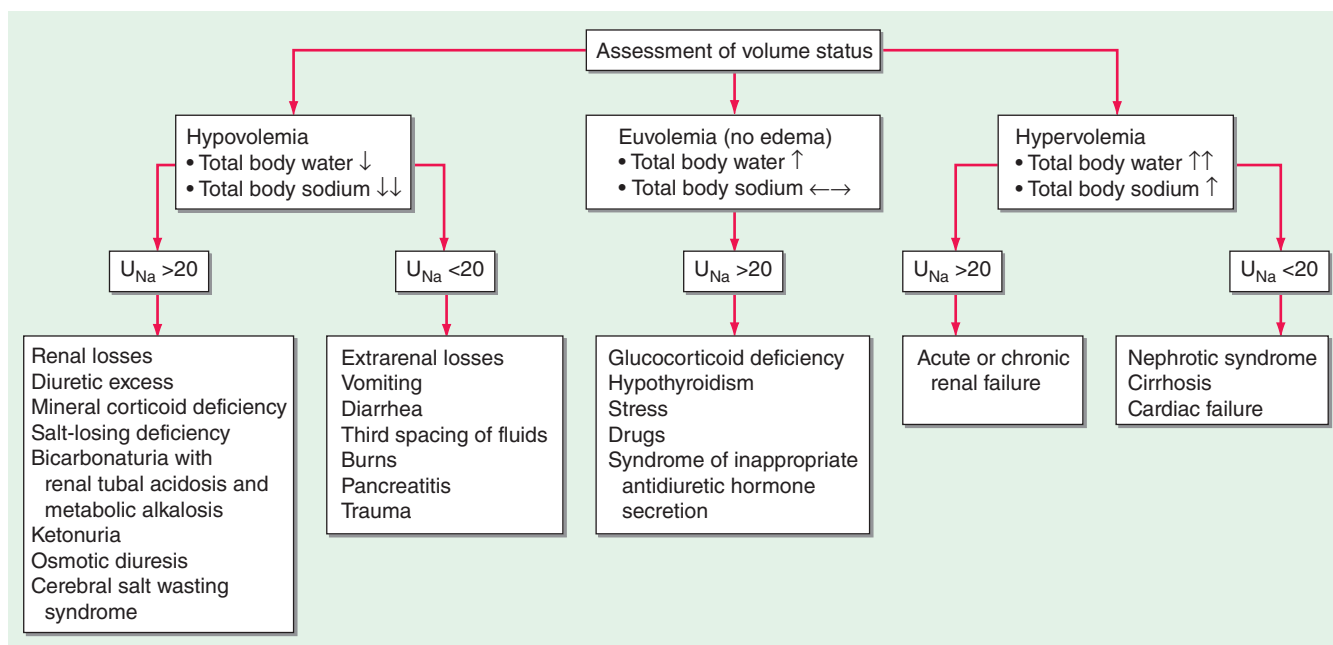


FIGURE 63-5 The diagnostic approach to hyponatremia. (From S Kumar, T Berl: *Diseases of water metabolism*, in *Atlas of Diseases of the Kidney*, RW Schrier [ed]. Philadelphia, Current Medicine, Inc, 1999; with permission.)

indicate the cause of their hyponatremia. Indeed, a urine Na^+ concentration <20 mM, in the absence of a cause of hypervolemic hyponatremia, predicts a rapid increase in plasma Na^+ concentration in response to intravenous normal saline; saline therapy thus induces a water diuresis in this setting, as circulating AVP levels plummet.

The renal causes of hypovolemic hyponatremia share an inappropriate loss of $\text{Na}^+\text{-Cl}^-$ in the urine, leading to volume depletion and an increase in circulating AVP; urine Na^+ concentration is typically >20 mM (Fig. 63-5). A deficiency in circulating aldosterone and/or its renal effects can lead to hyponatremia in primary adrenal insufficiency and other causes of hypoaldosteronism; hyperkalemia and hyponatremia in a hypotensive and/or hypovolemic patient with high urine Na^+ concentration (much greater than 20 mM) should strongly suggest this diagnosis. Salt-losing nephropathies may lead to hyponatremia when sodium intake is reduced, due to impaired renal tubular function; typical causes include reflux nephropathy, interstitial nephropathies, postobstructive uropathy, medullary cystic disease, and the recovery phase of acute tubular necrosis. Thiazide diuretics cause hyponatremia via a number of mechanisms, including polydipsia and diuretic-induced volume depletion. Notably, thiazides do not inhibit the renal concentrating mechanism, such that circulating AVP retains a full effect on renal water retention. In contrast, loop diuretics, which are less frequently associated with hyponatremia, inhibit $\text{Na}^+\text{-Cl}^-$ and K^+ absorption by the TALH, blunting the countercurrent mechanism and reducing the ability to concentrate the urine. Increased excretion of an osmotically active nonreabsorbable or poorly reabsorbable solute can also lead to volume depletion and hyponatremia; important causes include glycosuria, ketonuria (e.g., in starvation or in diabetic or alcoholic ketoacidosis), and bicarbonaturia (e.g., in renal tubular acidosis or metabolic alkalosis, where the associated bicarbonaturia leads to loss of Na^+).

Finally, the syndrome of “cerebral salt wasting” is a rare cause of hypovolemic hyponatremia, encompassing hyponatremia with clinical hypovolemia and inappropriate natriuresis in association with intracranial disease; associated disorders include subarachnoid hemorrhage, traumatic brain injury, craniotomy, encephalitis, and meningitis. Distinction from the more common syndrome of inappropriate antidiuresis is critical because cerebral salt wasting will typically respond to aggressive $\text{Na}^+\text{-Cl}^-$ repletion.

Hypervolemic Hyponatremia Patients with hypervolemic hyponatremia develop an increase in total-body $\text{Na}^+\text{-Cl}^-$ that is accompanied

by a proportionately *greater* increase in total-body water, leading to a reduced plasma Na^+ concentration. As in hypovolemic hyponatremia, the causative disorders can be separated by the effect on urine Na^+ concentration, with acute or chronic renal failure uniquely associated with an increase in urine Na^+ concentration (Fig. 63-5). The pathophysiology of hyponatremia in the sodium-avid edematous disorders (congestive heart failure [CHF], cirrhosis, and nephrotic syndrome) is similar to that in hypovolemic hyponatremia, except that arterial filling and circulatory integrity is decreased due to the specific etiologic factors (e.g., cardiac dysfunction in CHF, peripheral vasodilation in cirrhosis). Urine Na^+ concentration is typically very low, i.e., <10 mM, even after hydration with normal saline; this Na^+ -avid state may be obscured by diuretic therapy. The degree of hyponatremia provides an indirect index of the associated neurohumoral activation and is an important prognostic indicator in hypervolemic hyponatremia.

Euolemic Hyponatremia Euolemic hyponatremia can occur in moderate to severe hypothyroidism, with correction after achieving a euthyroid state. Severe hyponatremia can also be a consequence of secondary adrenal insufficiency due to pituitary disease; whereas the deficit in circulating aldosterone in primary adrenal insufficiency causes *hypovolemic* hyponatremia, the predominant glucocorticoid deficiency in secondary adrenal failure is associated with *euolemic* hyponatremia. Glucocorticoids exert a negative feedback on AVP release by the posterior pituitary such that hydrocortisone replacement in these patients will rapidly normalize the AVP response to osmolality, reducing circulating AVP.

The syndrome of inappropriate antidiuresis (SIAD) is the most frequent cause of euolemic hyponatremia (Table 63-1). The generation of hyponatremia in SIAD requires an intake of free water, with persistent intake at serum osmolalities that are lower than the usual threshold for thirst; as one would expect, the osmotic threshold and osmotic response curves for the sensation of thirst are shifted downward in patients with SIAD. Four distinct patterns of AVP secretion have been recognized in patients with SIAD, independent for the most part of the underlying cause. Unregulated, erratic AVP secretion is seen in about a third of patients, with no obvious correlation between serum osmolality and circulating AVP levels. Other patients fail to suppress AVP secretion at lower serum osmolalities, with a normal response curve to hyperosmolar conditions; others have a “reset osmostat,” with a lower threshold osmolality and a left-shifted osmotic response curve. Finally, the fourth subset of patients have essentially no detectable

TABLE 63-1 CAUSES OF THE SYNDROME OF INAPPROPRIATE ANTIDIURESIS (SIAD)

Malignant Diseases	Pulmonary Disorders	Disorders of the Central Nervous System	Drugs	Other Causes
Carcinoma	Infections	Infection	Drugs that stimulate release of AVP or enhance its action	Hereditary (gain-of-function mutations in the vasopressin V ₂ receptor)
Lung	Bacterial pneumonia	Encephalitis	Chlorpropamide	Idiopathic
Small cell	Viral pneumonia	Meningitis	SSRIs	Transient
Mesothelioma	Pulmonary abscess	Brain abscess	Tricyclic antidepressants	Endurance exercise
Oropharynx	Tuberculosis	Rocky Mountain spotted fever	Clofibrate	General anesthesia
Gastrointestinal tract	Aspergillosis	AIDS	Carbamazepine	Nausea
Stomach	Asthma	Bleeding and masses	Vincristine	Pain
Duodenum	Cystic fibrosis	Subdural hematoma	Nicotine	Stress
Pancreas	Respiratory failure associated with positive-pressure breathing	Subarachnoid hemorrhage	Narcotics	
Genitourinary tract		Cerebrovascular accident	Antipsychotic drugs	
Ureter		Brain tumors	Ifosfamide	
Bladder		Head trauma	Cyclophosphamide	
Prostate		Hydrocephalus	Nonsteroidal anti-inflammatory drugs	
Endometrium		Cavernous sinus thrombosis	MDMA (“ecstasy”)	
Endocrine thymoma		Other	AVP analogues	
Lymphomas		Multiple sclerosis	Desmopressin	
Sarcomas		Guillain-Barré syndrome	Oxytocin	
Ewing’s sarcoma		Shy-Drager syndrome	Vasopressin	
		Delirium tremens		
		Acute intermittent porphyria		

Abbreviations: AVP, vasopressin; MDMA; 3,4-methylenedioxymethamphetamine; SSRI, selective serotonin reuptake inhibitor.

Source: From DH Ellison, T Berl: Syndrome of inappropriate antidiuresis. *N Engl J Med* 356:2064, 2007.

circulating AVP, suggesting either a gain in function in renal water reabsorption or a circulating antidiuretic substance that is distinct from AVP. Gain-in-function mutations of a single specific residue in the V₂ AVP receptor have been described in some of these patients, leading to constitutive activation of the receptor in the absence of AVP and “nephrogenic” SIAD.

Strictly speaking, patients with SIAD are not euvolemic but are subclinically volume-expanded, due to AVP-induced water and Na⁺-Cl⁻ retention; “AVP escape” mechanisms invoked by sustained increases in AVP serve to limit distal renal tubular transport, preserving a modestly hypervolemic steady state. Serum uric acid is often low (<4 mg/dL) in patients with SIAD, consistent with suppressed proximal tubular transport in the setting of increased distal tubular Na⁺-Cl⁻ and water transport; in contrast, patients with hypovolemic hyponatremia will often be hyperuricemic, due to a shared activation of proximal tubular Na⁺-Cl⁻ and urate transport.

Common causes of SIAD include pulmonary disease (e.g., pneumonia, tuberculosis, pleural effusion) and central nervous system (CNS) diseases (e.g., tumor, subarachnoid hemorrhage, meningitis). SIAD also occurs with malignancies, most commonly with small-cell lung carcinoma (75% of malignancy-associated SIAD); ~10% of patients with this tumor will have a plasma Na⁺ concentration of <130 mM at presentation. SIAD is also a frequent complication of certain drugs, most commonly the selective serotonin reuptake inhibitors (SSRIs). Other drugs can potentiate the renal effect of AVP, without exerting direct effects on circulating AVP levels (Table 63-1).

Low Solute Intake and Hyponatremia Hyponatremia can occasionally occur in patients with a very low intake of dietary solutes. Classically, this occurs in alcoholics whose sole nutrient is beer, hence the diagnostic label of *beer potomania*; beer is very low in protein and salt content, containing only 1–2 mM of Na⁺. The syndrome has also been described in nonalcoholic patients with highly restricted solute intake due to nutrient-restricted diets, e.g., extreme vegetarian diets. Patients with hyponatremia due to low solute intake typically present with a very low urine osmolality (<100–200 mOsm/kg) with a urine Na⁺ concentration that is <10–20 mM. The fundamental abnormality

is the inadequate dietary intake of solutes; the reduced urinary solute excretion limits water excretion such that hyponatremia ensues after relatively modest polydipsia. AVP levels have not been reported in patients with beer potomania but are expected to be suppressed or rapidly suppressible with saline hydration; this fits with the overly rapid correction in plasma Na⁺ concentration that can be seen with saline hydration. Resumption of a normal diet and/or saline hydration will also correct the causative deficit in urinary solute excretion, such that patients with beer potomania typically correct their plasma Na⁺ concentration promptly after admission to the hospital.

Clinical Features of Hyponatremia Hyponatremia induces generalized cellular swelling, a consequence of water movement down the osmotic gradient from the hypotonic ECF to the ICF. The symptoms of hyponatremia are primarily neurologic, reflecting the development of cerebral edema within a rigid skull. The initial CNS response to acute hyponatremia is an increase in interstitial pressure, leading to shunting of ECF and solutes from the interstitial space into the cerebrospinal fluid and then on into the systemic circulation. This is accompanied by an efflux of the major intracellular ions, Na⁺, K⁺, and Cl⁻, from brain cells. Acute hyponatremic encephalopathy ensues when these volume regulatory mechanisms are overwhelmed by a rapid decrease in tonicity, resulting in acute cerebral edema. Early symptoms can include nausea, headache, and vomiting. However, severe complications can rapidly evolve, including seizure activity, brainstem herniation, coma, and death. A key complication of acute hyponatremia is normocapnic or hypercapnic respiratory failure; the associated hypoxia may amplify the neurologic injury. Normocapnic respiratory failure in this setting is typically due to noncardiogenic, “neurogenic” pulmonary edema, with a normal pulmonary capillary wedge pressure.

Acute symptomatic hyponatremia is a medical emergency, occurring in a number of specific settings (Table 63-2). Women, particularly before menopause, are much more likely than men to develop encephalopathy and severe neurologic sequelae. Acute hyponatremia often has an iatrogenic component, e.g., when hypotonic intravenous fluids are given to postoperative patients with an increase in circulating AVP. Exercise-associated hyponatremia, an important clinical issue at

TABLE 63-2 CAUSES OF ACUTE HYPONATREMIA

iatrogenic
Postoperative: premenopausal women
Hypotonic fluids with cause of ↑ vasopressin
Glycine irrigation: TURP, uterine surgery
Colonoscopy preparation
Recent institution of thiazides
Polydipsia
MDMA (“ecstasy,” “Molly”) ingestion
Exercise induced
Multifactorial, e.g., thiazide and polydipsia

Abbreviations: MDMA, 3,4-methylenedioxymethamphetamine; TURP, transurethral resection of the prostate.

marathons and other endurance events, has similarly been linked to both a “nonosmotic” increase in circulating AVP and excessive free water intake. The recreational drugs Molly and ecstasy, which share an active ingredient (MDMA, 3,4-methylenedioxymethamphetamine), cause a rapid and potent induction of both thirst and AVP, leading to severe acute hyponatremia.

Persistent, chronic hyponatremia results in an efflux of organic osmolytes (creatine, betaine, glutamate, myoinositol, and taurine) from brain cells; this response reduces intracellular osmolality and the osmotic gradient favoring water entry. This reduction in intracellular osmolytes is largely complete within 48 h, the time period that clinically defines chronic hyponatremia; this temporal definition has considerable relevance for the treatment of hyponatremia (see below). The cellular response to chronic hyponatremia does not fully protect patients from symptoms, which can include vomiting, nausea, confusion, and seizures, usually at plasma Na^+ concentration $<125 \text{ mM}$. Even patients who are judged “asymptomatic” can manifest subtle gait and cognitive defects that reverse with correction of hyponatremia; notably, chronic “asymptomatic” hyponatremia increases the risk of falls. Chronic hyponatremia also increases the risk of bony fractures owing to the associated neurologic dysfunction and to a hyponatremia-associated reduction in bone density. Therefore, every attempt should be made to correct safely the plasma Na^+ concentration in patients with chronic hyponatremia, even in the absence of overt symptoms (see the section on treatment of hyponatremia below).

The management of chronic hyponatremia is complicated significantly by the asymmetry of the cellular response to correction of plasma Na^+ concentration. Specifically, the *reaccumulation* of organic osmolytes by brain cells is attenuated and delayed as osmolality increases after correction of hyponatremia, sometimes resulting in degenerative loss of oligodendrocytes and an osmotic demyelination syndrome (ODS). Overly rapid correction of hyponatremia ($>8\text{--}10 \text{ mM}$ in 24 h or 18 mM in 48 h) is also associated with a disruption in integrity of the blood-brain barrier, allowing the entry of immune mediators that may contribute to demyelination. The lesions of ODS classically affect the pons, a structure wherein the delay in the reaccumulation of osmotic osmolytes is particularly pronounced; clinically, patients with central pontine myelinolysis can present 1 or more days after overcorrection of hyponatremia with paraparesis or quadriparesis, dysphagia, dysarthria, diplopia, a “locked-in syndrome,” and/or loss of consciousness. Other regions of the brain can also be involved in ODS, most commonly in association with lesions of the pons but occasionally in isolation; in order of frequency, the lesions of extrapontine myelinolysis can occur in the cerebellum, lateral geniculate body, thalamus, putamen, and cerebral cortex or subcortex. Clinical presentation of ODS can, therefore, vary as a function of the extent and localization of extrapontine myelinolysis, with the reported development of ataxia, mutism, parkinsonism, dystonia, and catatonia. Relowering of plasma Na^+ concentration after overly rapid correction can prevent or attenuate ODS (see the section on treatment of hyponatremia below). However, even appropriately slow correction can be associated with ODS, particularly in patients with additional risk

factors; these include alcoholism, malnutrition, hypokalemia, and liver transplantation.

Diagnostic Evaluation of Hyponatremia Clinical assessment of hyponatremic patients should focus on the underlying cause; a detailed drug history is particularly crucial (Table 63-1). A careful clinical assessment of volume status is obligatory for the classical diagnostic approach to hyponatremia (Fig. 63-5). Hyponatremia is frequently multifactorial, particularly when severe; clinical evaluation should consider *all* the possible causes for excessive circulating AVP, including volume status, drugs, and the presence of nausea and/or pain. Radiologic imaging may also be appropriate to assess whether patients have a pulmonary or CNS cause for hyponatremia. A screening chest x-ray may fail to detect a small-cell carcinoma of the lung; computed tomography (CT) scanning of the thorax should be considered in patients at high risk for this tumor (e.g., patients with a smoking history).

Laboratory investigation should include a measurement of serum osmolality to exclude pseudohyponatremia, which is defined as the coexistence of hyponatremia with a normal or increased plasma tonicity. Most clinical laboratories measure plasma Na^+ concentration by testing diluted samples with automated ion-sensitive electrodes, correcting for this dilution by assuming that plasma is 93% water. This correction factor can be inaccurate in patients with pseudohyponatremia due to extreme hyperlipidemia and/or hyperproteinemia, in whom serum lipid or protein makes up a greater percentage of plasma volume. The measured osmolality should also be converted to the effective osmolality (tonicity) by subtracting the measured concentration of urea (divided by 2.8, if in mg/dL); patients with hyponatremia have an effective osmolality of $<275 \text{ mOsm/kg}$.

Elevated BUN and creatinine in routine chemistries can also indicate renal dysfunction as a potential cause of hyponatremia, whereas hyperkalemia may suggest adrenal insufficiency or hypoaldosteronism. Serum glucose should also be measured; plasma Na^+ concentration falls by $\sim 1.6\text{--}2.4 \text{ mM}$ for every 100-mg/dL increase in glucose, due to glucose-induced water efflux from cells; this “true” hyponatremia resolves after correction of hyperglycemia. Measurement of serum uric acid should also be performed; whereas patients with SIAD-type physiology will typically be hypouricemic (serum uric acid $<4 \text{ mg/dL}$), volume-depleted patients will often be hyperuricemic. In the appropriate clinical setting, thyroid, adrenal, and pituitary function should also be tested; hypothyroidism and secondary adrenal failure due to pituitary insufficiency are important causes of euvolemic hyponatremia, whereas primary adrenal failure causes hypovolemic hyponatremia. A cosyntropin stimulation test is necessary to assess for primary adrenal insufficiency.

Urine electrolytes and osmolality are crucial tests in the initial evaluation of hyponatremia. A urine Na^+ concentration $<20\text{--}30 \text{ mM}$ is consistent with hypovolemic hyponatremia, in the clinical absence of a hypervolemic, Na^+ -avid syndrome such as CHF (Fig. 63-5). In contrast, patients with SIAD will typically excrete urine with an Na^+ concentration that is $>30 \text{ mM}$. However, there can be substantial overlap in urine Na^+ concentration values in patients with SIAD and hypovolemic hyponatremia, particularly in the elderly; the ultimate “gold standard” for the diagnosis of hypovolemic hyponatremia is the demonstration that plasma Na^+ concentration corrects after hydration with normal saline. Patients with thiazide-associated hyponatremia may also present with higher than expected urine Na^+ concentration and other findings suggestive of SIAD; one should defer making a diagnosis of SIAD in these patients until 1–2 weeks after discontinuing the thiazide. A urine osmolality $<100 \text{ mOsm/kg}$ is suggestive of polydipsia; urine osmolality $>400 \text{ mOsm/kg}$ indicates that AVP excess is playing a more dominant role, whereas intermediate values are more consistent with multifactorial pathophysiology (e.g., AVP excess with a significant component of polydipsia). Patients with hyponatremia due to decreased solute intake (beer potomania) typically have urine Na^+ concentration $<20 \text{ mM}$ and urine osmolality in the range of <100 to the low 200s. Finally, the measurement of urine K^+ concentration is required to calculate the urine-to-plasma electrolyte ratio, which is useful to predict the response to fluid restriction (see the section on treatment of hyponatremia below).

Three major considerations guide the therapy of hyponatremia. First, the presence and/or severity of symptoms determine the urgency and goals of therapy. Patients with acute hyponatremia (Table 63-2) present with symptoms that can range from headache, nausea, and/or vomiting, to seizures, obtundation, and central herniation; patients with chronic hyponatremia, present for >48 h, are less likely to have severe symptoms. Second, patients with chronic hyponatremia are at risk for ODS if plasma Na⁺ concentration is corrected by >8–10 mM within the first 24 h and/or by >18 mM within the first 48 h. Third, the response to interventions such as hypertonic saline, isotonic saline, or AVP antagonists can be highly unpredictable, such that frequent monitoring of plasma Na⁺ concentration during corrective therapy is imperative.

Once the urgency in correcting the plasma Na⁺ concentration has been established and appropriate therapy instituted, the focus should be on treatment or withdrawal of the underlying cause. Patients with euvolemic hyponatremia due to SIAD, hypothyroidism, or secondary adrenal failure will respond to successful treatment of the underlying cause, with an increase in plasma Na⁺ concentration. However, not all causes of SIAD are immediately reversible, necessitating pharmacologic therapy to increase the plasma Na⁺ concentration (see below). Hypovolemic hyponatremia will respond to intravenous hydration with isotonic normal saline, with a rapid reduction in circulating AVP and a brisk water diuresis; it may be necessary to reduce the rate of correction if the history suggests that hyponatremia has been chronic, i.e., present for more than 48 h (see below). Hypervolemic hyponatremia due to CHF will often respond to improved therapy of the underlying cardiomyopathy, e.g., following the institution or intensification of angiotensin-converting enzyme (ACE) inhibition. Finally, patients with hyponatremia due to beer potomania and low solute intake will respond very rapidly to intravenous saline and the resumption of a normal diet. Notably, patients with beer potomania have a very high risk of developing ODS, due to the associated hypokalemia, alcoholism, malnutrition, and high risk of overcorrecting the plasma Na⁺ concentration.

Water deprivation has long been a cornerstone of the therapy of chronic hyponatremia. However, patients who are excreting minimal electrolyte-free water will require aggressive fluid restriction; this can be very difficult for patients with SIAD to tolerate, given that their thirst is also inappropriately stimulated. The urine-to-plasma electrolyte ratio (urinary [Na⁺] + [K⁺]/plasma [Na⁺]) can be exploited as a quick indicator of electrolyte-free water excretion (Table 63-3); patients with a ratio of >1 should be more aggressively restricted

(<500 mL/d), those with a ratio of ~1 should be restricted to 500–700 mL/d, and those with a ratio <1 should be restricted to <1 L/d. In hypokalemic patients, potassium replacement will serve to increase plasma Na⁺ concentration, given that the plasma Na⁺ concentration is a functional of both exchangeable Na⁺ and exchangeable K⁺ divided by total-body water; a corollary is that aggressive repletion of K⁺ has the potential to overcorrect the plasma Na⁺ concentration even in the absence of hypertonic saline. Plasma Na⁺ concentration will also tend to respond to an increase in dietary solute intake, which increases the ability to excrete free water; however, the use of oral urea and/or salt tablets for this purpose is generally not practical or well tolerated.

Patients in whom therapy with fluid restriction, potassium replacement, and/or increased solute intake fails may merit pharmacologic therapy to increase their plasma Na⁺ concentration. Many patients with SIAD respond to combined therapy with oral furosemide, 20 mg twice a day (higher doses may be necessary in renal insufficiency), and oral salt tablets; furosemide serves to inhibit the renal countercurrent mechanism and blunt urinary concentrating ability, whereas the salt tablets counteract diuretic-associated natriuresis. Demeclocycline is a potent inhibitor of principal cells and can be used in patients whose Na levels do not increase in response to furosemide and salt tablets. However, this agent can be associated with a reduction in GFR, due to excessive natriuresis and/or direct renal toxicity; it should be avoided in cirrhotic patients in particular, who are at higher risk of nephrotoxicity due to drug accumulation.

AVP antagonists (vaptans) are highly effective in SIAD and in hypervolemic hyponatremia due to heart failure or cirrhosis, reliably increasing plasma Na⁺ concentration due to their “aquaretic” effects (augmentation of free water clearance). Most of these agents specifically antagonize the V₂ AVP receptor; tolvaptan is currently the only oral V₂ antagonist to be approved by the U.S. Food and Drug Administration. Conivaptan, the only available intravenous vaptan, is a mixed V_{1A}/V₂ antagonist, with a modest risk of hypotension due to V_{1A} receptor inhibition. Therapy with vaptans must be initiated in a hospital setting, with a liberalization of fluid restriction (>2 L/d) and close monitoring of plasma Na⁺ concentration. Although approved for the management of all but hypovolemic hyponatremia and acute hyponatremia, the clinical indications for these agents are not completely clear. Oral tolvaptan is perhaps most appropriate for the management of significant and persistent SIAD (e.g., in small-cell lung carcinoma) that has not responded to water restriction and/or oral furosemide and salt tablets. Abnormalities in liver function tests have been reported with chronic tolvaptan therapy; hence, the use of this agent should be restricted to <1–2 months.

Treatment of acute symptomatic hyponatremia should include hypertonic 3% saline (513 mM) to acutely increase plasma Na⁺ concentration by 1–2 mM/h to a total of 4–6 mM; this modest increase is typically sufficient to alleviate severe acute symptoms, after which corrective guidelines for chronic hyponatremia are appropriate (see below). A number of equations have been developed to estimate the required rate of hypertonic saline, which has an Na⁺-Cl⁻ concentration of 513 mM. The traditional approach is to calculate an Na⁺ deficit, where the Na⁺ deficit = 0.6 × body weight × (target plasma Na⁺ concentration – starting plasma Na⁺ concentration), followed by a calculation of the required rate. Regardless of the method used to determine the rate of administration, the increase in plasma Na⁺ concentration can be highly unpredictable during treatment with hypertonic saline, due to rapid changes in the underlying physiology; plasma Na⁺ concentration should be monitored every 2–4 h during treatment, with appropriate changes in therapy based on the observed rate of change. The administration of supplemental oxygen and ventilatory support is also critical in acute hyponatremia, in the event that patients develop acute pulmonary edema or hypercapnic respiratory failure. Intravenous loop diuretics will help treat acute pulmonary edema and will also increase free water excretion, by interfering with the renal countercurrent multiplication system. AVP antagonists do not have an approved role in the management of acute hyponatremia.

TABLE 63-3 MANAGEMENT OF HYPERNATREMIA

Water Deficit

1. Estimate total-body water (TBW): 50% of body weight in women and 60% in men
2. Calculate free-water deficit: $[(\text{Na}^+ - 140)/140] \times \text{TBW}$
3. Administer deficit over 48–72 h, without decrease in plasma Na⁺ concentration by >10 mM/24 h

Ongoing Water Losses

4. Calculate free-water clearance, C_eH₂O:

$$C_{eH_2O} = V \times \left(1 - \frac{U_{Na} + U_K}{P_{Na}} \right)$$

where V is urinary volume, U_{Na} is urinary [Na⁺], U_K is urinary [K⁺], and P_{Na} is plasma [Na⁺]

Insensible Losses

5. ~10 mL/kg per day: less if ventilated, more if febrile

Total

6. Add components to determine water deficit and ongoing water loss; correct the water deficit over 48–72 h and replace daily water loss. Avoid correction of plasma [Na⁺] by >10 mM/d.

The rate of correction should be comparatively slow in *chronic* hyponatremia (<8–10 mM in the first 24 h and <18 mM in the first 48 h), so as to avoid ODS; lower target rates are appropriate in patients at particular risk for ODS, such as alcoholics or hypokalemic patients. Overcorrection of the plasma Na⁺ concentration can occur when AVP levels rapidly normalize, for example following the treatment of patients with chronic hypovolemic hyponatremia with intravenous saline or following glucocorticoid replacement of patients with hypopituitarism and secondary adrenal failure. Approximately 10% of patients treated with vaptans will overcorrect; the risk is increased if water intake is not liberalized. In the event that the plasma Na⁺ concentration overcorrects following therapy, be it with hypertonic saline, isotonic saline, or a vaptan, hyponatremia can be safely reinduced or stabilized by the administration of the AVP *agonist* desmopressin acetate (DDAVP) and/or the administration of free water, typically intravenous D₅W; the goal is to prevent or reverse the development of ODS. Alternatively, the treatment of patients with marked hyponatremia can be initiated with the twice-daily administration of DDAVP to maintain constant AVP bioactivity, combined with the administration of hypertonic saline to slowly correct the serum sodium in a more controlled fashion, thus reducing upfront the risk of overcorrection.

HYPERNATREMIA

Etiology Hyponatremia is defined as an increase in the plasma Na⁺ concentration to >145 mM. Considerably less common than hyponatremia, hypernatremia is nonetheless associated with mortality rates of as high as 40–60%, mostly due to the severity of the associated underlying disease processes. Hypernatremia is usually the result of a combined water and electrolyte deficit, with losses of H₂O in excess of Na⁺. Less frequently, the ingestion or iatrogenic administration of excess Na⁺ can be causative, for example after IV administration of excessive hypertonic Na⁺-Cl⁻ or Na⁺-HCO₃⁻ (Fig. 63-6).

Elderly individuals with reduced thirst and/or diminished access to fluids are at the highest risk of developing hypernatremia. Patients with hypernatremia may rarely have a central defect in hypothalamic osmoreceptor function, with a mixture of both decreased thirst and reduced AVP secretion. Causes of this adipsic diabetes insipidus include primary or metastatic tumor, occlusion or ligation of the anterior communicating artery, trauma, hydrocephalus, and inflammation.

Hypernatremia can develop following the loss of water via both renal and nonrenal routes. Insensible losses of water may increase in the setting of fever, exercise, heat exposure, severe burns, or mechanical ventilation. Diarrhea is, in turn, the most common gastrointestinal cause of hypernatremia. Notably, osmotic diarrhea and viral gastroenteritis typically generate stools with Na⁺ and K⁺ <100 mM, thus leading to water loss and hypernatremia; in contrast, secretory diarrhea typically results in isotonic stool and thus hypovolemia with or without hypovolemic hyponatremia.

Common causes of renal water loss include osmotic diuresis secondary to hyperglycemia, excess urea, postobstructive diuresis, or mannitol; these disorders share an increase in urinary solute excretion and urinary osmolality (see “Diagnostic Approach,” below). Hypernatremia due to a water diuresis occurs in central or nephrogenic diabetes insipidus (DI).

Nephrogenic DI (NDI) is characterized by renal resistance to AVP, which can be partial or complete (see “Diagnostic Approach,” below). Genetic causes include loss-of-function mutations in the X-linked V₂ receptor; mutations in the AVP-responsive aquaporin-2 water channel can cause autosomal recessive and autosomal dominant NDI, whereas recessive deficiency of the aquaporin-1 water channel causes a more modest concentrating defect (Fig. 63-2). Hypercalcemia can also cause polyuria and NDI; calcium signals directly through the calcium-sensing receptor to downregulate Na⁺, K⁺, and Cl⁻ transport by the TALH and water transport in principal cells, thus reducing renal concentrating ability in hypercalcemia. Another common acquired cause of NDI is hypokalemia, which inhibits the renal response to AVP and downregulates aquaporin-2 expression. Several drugs can cause acquired NDI, in particular lithium, ifosfamide, and several anti-tumor agents. Lithium causes NDI by multiple mechanisms, including direct inhibition of renal glycogen synthase kinase-3 (GSK3), a kinase thought to be the pharmacologic target of lithium in bipolar disease; GSK3 is required for the response of principal cells to AVP. The entry of lithium through the amiloride-sensitive Na⁺ channel ENaC (Fig. 63-4) is required for the effect of the drug on principal cells, such that combined therapy within lithium and amiloride can mitigate lithium-associated NDI. However, lithium causes chronic tubulointerstitial scarring and chronic kidney disease after prolonged therapy, such that patients may have a persistent NDI long after stopping the drug, with a reduced therapeutic benefit from amiloride.

Finally, gestational DI is a rare complication of late-term pregnancy wherein increased activity of a circulating placental protease with “vasopressinase” activity leads to reduced circulating AVP and polyuria, often accompanied by hypernatremia. DDAVP is an effective therapy for this syndrome, given its resistance to the vasopressinase enzyme.

Clinical Features Hypernatremia increases osmolality of the ECF, generating an osmotic gradient between the ECF and ICF, an efflux of intracellular water, and cellular shrinkage. As in hyponatremia, the symptoms of hypernatremia are predominantly neurologic. Altered mental status is the most frequent manifestation, ranging from mild confusion and lethargy to deep coma. The sudden shrinkage of brain cells in acute hypernatremia may lead to parenchymal or subarachnoid hemorrhages and/or subdural hematomas; however, these vascular complications are primarily encountered in pediatric and neonatal patients. Osmotic damage to muscle membranes can also lead to hypernatremic rhabdomyolysis. Brain cells accommodate to a chronic increase in ECF osmolality (>48 h) by activating membrane transporters that mediate influx and intracellular accumulation of organic osmolytes (creatine, betaine, glutamate, myoinositol, and taurine); this results in an increase in ICF water and normalization of brain parenchymal volume. In consequence, patients with *chronic* hypernatremia are less likely to develop severe neurologic compromise. However, the cellular response to chronic hypernatremia predisposes these patients to the development of cerebral edema and seizures during overly rapid hydration (overcorrection of plasma Na⁺ concentration by >10 mM/d).

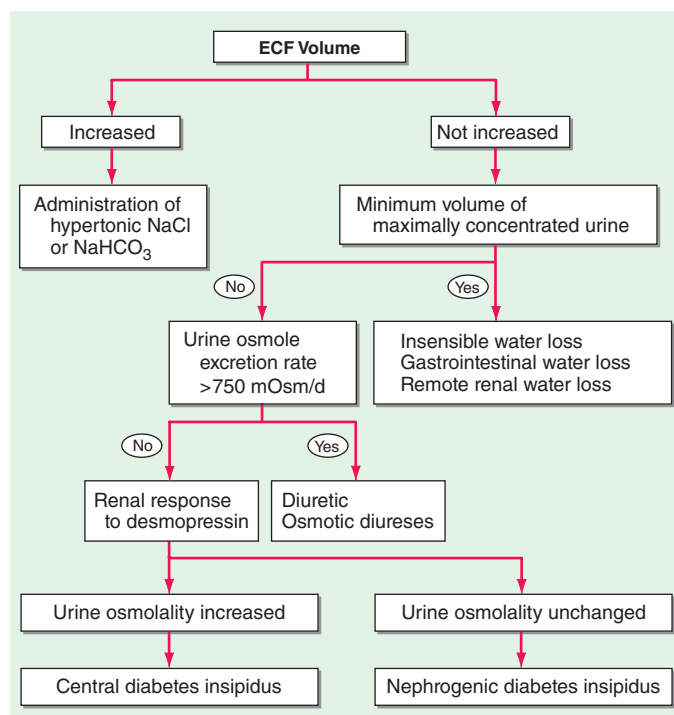


FIGURE 63-6 The diagnostic approach to hypernatremia. ECF, extracellular fluid.

Diagnostic Approach The history should focus on the presence or absence of thirst, polyuria, and/or an extrarenal source for water loss,

such as diarrhea. The physical examination should include a detailed neurologic exam and an assessment of the ECFV; patients with a particularly large water deficit and/or a combined deficit in electrolytes and water may be hypovolemic, with reduced JVP and orthostasis. Accurate documentation of daily fluid intake and daily urine output is also critical for the diagnosis and management of hypernatremia.

Laboratory investigation should include a measurement of serum and urine osmolality, in addition to urine electrolytes. The appropriate response to hypernatremia and a serum osmolality >295 mOsm/kg is an increase in circulating AVP and the excretion of low volumes (<500 mL/d) of maximally concentrated urine, i.e., urine with osmolality >800 mOsm/kg; should this be the case, then an extrarenal source of water loss is primarily responsible for the generation of hypernatremia. Many patients with hypernatremia are polyuric; should an osmotic diuresis be responsible, with excessive excretion of $\text{Na}^+\text{-Cl}^-$, glucose, and/or urea, then daily solute excretion will be $>750\text{--}1000$ mOsm/d (>15 mOsm/kg body water per day) (Fig. 63-6). More commonly, patients with hypernatremia and polyuria will have a predominant water diuresis, with excessive excretion of hypotonic, dilute urine.

Adequate differentiation between nephrogenic and central causes of DI requires the measurement of the response in urinary osmolality to DDAVP, combined with measurement of circulating AVP in the setting of hypertonicity. By definition, patients with baseline hypernatremia are hypertonic, with an adequate stimulus for AVP by the posterior pituitary. Therefore, in contrast to polyuric patients with a normal or reduced baseline plasma Na^+ concentration and osmolality, a water deprivation test (Chap. 61) is unnecessary in hypernatremia; indeed, water deprivation is absolutely contraindicated in this setting, given the risk for worsening the hypernatremia. Patients with NDI will fail to respond to DDAVP, with a urine osmolality that increases by $<50\%$ or <150 mOsm/kg from baseline, in combination with a normal or high circulating AVP level; patients with central DI will respond to DDAVP, with a reduced circulating AVP. Patients may exhibit a partial response to DDAVP, with a $>50\%$ rise in urine osmolality that nonetheless fails to reach 800 mOsm/kg; the level of circulating AVP will help differentiate the underlying cause, i.e., NDI versus central DI. In pregnant patients, AVP assays should be drawn in tubes containing the protease inhibitor 1,10-phenanthroline, to prevent in vitro degradation of AVP by placental vasopressinase.

For patients with hypernatremia due to renal loss of water, it is critical to quantify ongoing daily losses using the calculated electrolyte-free water clearance, in addition to calculation of the baseline water deficit (the relevant formulas are discussed in Table 63-3). This requires daily measurement of urine electrolytes, combined with accurate measurement of daily urine volume.

TREATMENT HYPERNATREMIA

The underlying cause of hypernatremia should be withdrawn or corrected, be it drugs, hyperglycemia, hypercalcemia, hypokalemia, or diarrhea. The approach to the correction of hypernatremia is outlined in Table 63-3. It is imperative to correct hypernatremia slowly to avoid cerebral edema, typically replacing the calculated free water deficit over 48 h. Notably, the plasma Na^+ concentration should be corrected by no more than 10 mM/d, which may take longer than 48 h in patients with severe hypernatremia (>160 mM). A rare exception is patients with acute hypernatremia (<48 h) due to sodium loading, who can safely be corrected rapidly at a rate of 1 mM/h.

Water should ideally be administered by mouth or by nasogastric tube, as the most direct way to provide free water, i.e., water without electrolytes. Alternatively, patients can receive free water in dextrose-containing IV solutions, such as 5% dextrose (D_5W); blood glucose should be monitored in case hyperglycemia occurs. Depending on the history, blood pressure, or clinical volume status, it may be appropriate to initially treat with hypotonic saline solutions (1/4 or 1/2 normal saline); normal saline is usually inappropriate in the absence of very severe hypernatremia, where normal saline is proportionally more hypotonic relative to plasma, or frank

hypotension. Calculation of urinary electrolyte-free water clearance (Table 63-3) is required to estimate daily, ongoing loss of free water in patients with NDI or central DI, which should be replenished daily.

Additional therapy may be feasible in specific cases. Patients with central DI should respond to the administration of intravenous, intranasal, or oral DDAVP. Patients with NDI due to lithium may reduce their polyuria with amiloride (2.5–10 mg/d), which decreases entry of lithium into principal cells by inhibiting ENaC (see above); in practice, however, most patients with lithium-associated DI are able to compensate for their polyuria by simply increasing their daily water intake. Thiazides may reduce polyuria due to NDI, ostensibly by inducing hypovolemia and increasing proximal tubular water reabsorption. Occasionally, nonsteroidal anti-inflammatory drugs (NSAIDs) have been used to treat polyuria associated with NDI, reducing the negative effect of intrarenal prostaglandins on urinary concentrating mechanisms; however, this assumes the risks of NSAID-associated gastric and/or renal toxicity. Furthermore, it must be emphasized that thiazides, amiloride, and NSAIDs are only appropriate for chronic management of polyuria from NDI and have no role in the acute management of associated hypernatremia, where the focus is on replacing free water deficits and ongoing free water loss.

POTASSIUM DISORDERS

Homeostatic mechanisms maintain plasma K^+ concentration between 3.5 and 5.0 mM, despite marked variation in dietary K^+ intake. In a healthy individual at steady state, the entire daily intake of potassium is excreted, approximately 90% in the urine and 10% in the stool; thus, the kidney plays a dominant role in potassium homeostasis. However, more than 98% of total-body potassium is intracellular, chiefly in muscle; buffering of extracellular K^+ by this large intracellular pool plays a crucial role in the regulation of plasma K^+ concentration. Changes in the exchange and distribution of intra- and extracellular K^+ can thus lead to marked hypo- or hyperkalemia. A corollary is that massive necrosis and the attendant release of tissue K^+ can cause severe hyperkalemia, particularly in the setting of acute kidney injury and reduced excretion of K^+ .

Changes in whole-body K^+ content are primarily mediated by the kidney, which reabsorbs filtered K^+ in hypokalemic, K^+ -deficient states and secretes K^+ in hyperkalemic, K^+ -replete states. Although K^+ is transported along the entire nephron, it is the principal cells of the connecting segment (CNT) and cortical CD that play a dominant role in renal K^+ secretion, whereas alpha-intercalated cells of the outer medullary CD function in renal tubular reabsorption of filtered K^+ in K^+ -deficient states. In principal cells, apical Na^+ entry via the amiloride-sensitive ENaC generates a lumen-negative potential difference, which drives passive K^+ exit through apical K^+ channels (Fig. 63-4). Two major K^+ channels mediate distal tubular K^+ secretion: the secretory K^+ channel ROMK (renal outer medullary K^+ channel; also known as Kir1.1 or Kcnj11) and the flow-sensitive big potassium (BK) or maxi- K^+ channel. ROMK is thought to mediate the bulk of constitutive K^+ secretion, whereas increases in distal flow rate and/or genetic absence of ROMK activate K^+ secretion via the BK channel.

An appreciation of the relationship between ENaC-dependent Na^+ entry and distal K^+ secretion (Fig. 63-4) is required for the bedside interpretation of potassium disorders. For example, decreased distal delivery of Na^+ , as occurs in hypovolemic, prerenal states, tends to blunt the ability to excrete K^+ , leading to hyperkalemia; on the other hand, an increase in distal delivery of Na^+ and distal flow rate, as occurs after treatment with thiazide and loop diuretics, can enhance K^+ secretion and lead to hypokalemia. Hyperkalemia is also a predictable consequence of drugs that directly inhibit ENaC, due to the role of this Na^+ channel in generating a lumen-negative potential difference. Aldosterone in turn has a major influence on potassium excretion, increasing the activity of ENaC channels and thus amplifying the driving force for K^+ secretion across the luminal membrane of principal cells. Abnormalities in the renin-angiotensin-aldosterone system can thus cause both hypokalemia and hyperkalemia. Notably,

however, potassium excess and potassium restriction have opposing, aldosterone-independent effects on the density and activity of apical K^+ channels in the distal nephron, i.e., factors other than aldosterone modulate the renal capacity to secrete K^+ . In addition, potassium restriction and hypokalemia activates aldosterone-independent distal *reabsorption* of filtered K^+ , activating apical H^+/K^+ -ATPase activity in intercalated cells within the outer medullary CD. Reflective perhaps of this physiology, changes in plasma K^+ concentration are not universal in disorders associated with changes in aldosterone activity.

HYPOKALEMIA

Hypokalemia, defined as a plasma K^+ concentration of <3.5 mM, occurs in up to 20% of hospitalized patients. Hypokalemia is associated with a 10-fold increase in in-hospital mortality, due to adverse effects on cardiac rhythm, blood pressure, and cardiovascular morbidity. Mechanistically, hypokalemia can be caused by redistribution of K^+ between tissues and the ECF or by renal and nonrenal loss of K^+ (Table 63-4). Systemic hypomagnesemia can also cause treatment-resistant hypokalemia, due to a combination of reduced cellular uptake of K^+ and exaggerated renal secretion. Spurious hypokalemia or “pseudohypokalemia” can occasionally result from *in vitro* cellular uptake of K^+ after venipuncture, for example, due to profound leukocytosis in acute leukemia.

Redistribution and Hypokalemia Insulin, β_2 -adrenergic activity, thyroid hormone, and alkalosis promote Na^+/K^+ -ATPase-mediated cellular uptake of K^+ , leading to hypokalemia. Inhibition of the passive *efflux* of K^+ can also cause hypokalemia, albeit rarely; this typically occurs in the setting of systemic inhibition of K^+ channels by toxic barium ions. Exogenous insulin can cause iatrogenic hypokalemia, particularly during the management of K^+ -deficient states such as diabetic ketoacidosis. Alternatively, the stimulation of *endogenous* insulin can provoke hypokalemia, hypomagnesemia, and/or hypophosphatemia in malnourished patients given a carbohydrate load. Alterations in the activity of the endogenous sympathetic nervous system can cause hypokalemia in several settings, including alcohol withdrawal, hyperthyroidism, acute myocardial infarction, and severe head injury. β_2 agonists, including both bronchodilators and tocolytics (ritodrine), are powerful activators of cellular K^+ uptake; “hidden” sympathomimetics, such as pseudoephedrine and ephedrine in cough syrup or dieting agents, may also cause unexpected hypokalemia. Finally, xanthine-dependent activation of cAMP-dependent signaling, downstream of the β_2 receptor, can lead to hypokalemia, usually in the setting of overdose (theophylline) or marked overingestion (dietary caffeine).

Redistributive hypokalemia can also occur in the setting of hyperthyroidism, with periodic attacks of hypokalemic paralysis (thyrotoxic periodic paralysis [TPP]). Similar episodes of hypokalemic weakness in the absence of thyroid abnormalities occur in *familial* hypokalemic periodic paralysis, usually caused by missense mutations of voltage sensor domains within the α_1 subunit of L-type calcium channels or the skeletal Na^+ channel; these mutations generate an abnormal gating pore current activated by hyperpolarization. TPP develops more frequently in patients of Asian or Hispanic origin; this shared predisposition has been linked to genetic variation in Kir2.6, a muscle-specific, thyroid hormone-responsive K^+ channel. Patients with TPP typically present with weakness of the extremities and limb girdles, with paralytic episodes that occur most frequently between 1 and 6 AM. Signs and symptoms of hyperthyroidism are not invariably present. Hypokalemia is usually profound and almost invariably accompanied by hypophosphatemia and hypomagnesemia. The hypokalemia in TPP is attributed to both direct and indirect activation of the Na^+/K^+ -ATPase, resulting in increased uptake of K^+ by muscle and other tissues. Increases in β -adrenergic activity play an important role in that high-dose propranolol (3 mg/kg) rapidly reverses the associated hypokalemia, hypophosphatemia, and paralysis.

Nonrenal Loss of Potassium The loss of K^+ in sweat is typically low, except under extremes of physical exertion. Direct gastric losses of K^+ due to vomiting or nasogastric suctioning are also minimal; however, the ensuing hypochloremic alkalosis results in persistent kaliuresis due to secondary hyperaldosteronism and bicarbonaturia, i.e., a *renal* loss

TABLE 63-4 CAUSES OF HYPOKALEMIA

- I. Decreased intake
 - A. Starvation
 - B. Clay ingestion
- II. Redistribution into cells
 - A. Acid-base
 1. Metabolic alkalosis
 - B. Hormonal
 1. Insulin
 2. Increased β_2 -adrenergic sympathetic activity: post-myocardial infarction, head injury
 3. β_2 -Adrenergic agonists – bronchodilators, tocolytics
 4. α -Adrenergic antagonists
 5. Thyrotoxic periodic paralysis
 6. Downstream stimulation of Na^+/K^+ -ATPase: theophylline, caffeine
 - C. Anabolic state
 1. Vitamin B_{12} or folic acid administration (red blood cell production)
 2. Granulocyte-macrophage colony-stimulating factor (white blood cell production)
 3. Total parenteral nutrition
 - D. Other
 1. Pseudohypokalemia
 2. Hypothermia
 3. Familial hypokalemic periodic paralysis
 4. Barium toxicity: systemic inhibition of “leak” K^+ channels
- III. Increased loss
 - A. Nonrenal
 1. Gastrointestinal loss (diarrhea)
 2. Integumentary loss (sweat)
 - B. Renal
 1. Increased distal flow and distal Na^+ delivery: diuretics, osmotic diuresis, salt-wasting nephropathies
 2. Increased secretion of potassium
 - a. Mineralocorticoid excess: primary hyperaldosteronism (aldosterone-producing adenomas, primary or unilateral adrenal hyperplasia, idiopathic hyperaldosteronism due to bilateral adrenal hyperplasia, and adrenal carcinoma), genetic hyperaldosteronism (familial hyperaldosteronism types I/II/III, congenital adrenal hyperplasias), secondary hyperaldosteronism (malignant hypertension, renin-secreting tumors, renal artery stenosis, hypovolemia), Cushing’s syndrome, Bartter’s syndrome, Gitelman’s syndrome
 - b. Apparent mineralocorticoid excess: genetic deficiency of 11 β -dehydrogenase-2 (syndrome of apparent mineralocorticoid excess), inhibition of 11 β -dehydrogenase-2 (glycyrrheticin/glycyrrhizinic acid and/or carbenoxolone; licorice, food products, drugs), Liddle’s syndrome (genetic activation of epithelial Na^+ channels)
 - c. Distal delivery of nonreabsorbed anions: vomiting, nasogastric suction, proximal renal tubular acidosis, diabetic ketoacidosis, glue-sniffing (toluene abuse), penicillin derivatives (penicillin, nafcillin, dicloxacillin, ticarcillin, oxacillin, and carbenicillin)
 3. Magnesium deficiency

of K^+ . Diarrhea is a globally important cause of hypokalemia, given the worldwide prevalence of infectious diarrheal disease. Noninfectious gastrointestinal processes such as celiac disease, ileostomy, villous adenomas, inflammatory bowel disease, colonic pseudo-obstruction (Ogilvie’s syndrome), VIPomas, and chronic laxative abuse can also cause significant hypokalemia; an exaggerated intestinal secretion of potassium by upregulated colonic BK channels has been directly implicated in the pathogenesis of hypokalemia in many of these disorders.

Renal Loss of Potassium Drugs can increase renal K^+ excretion by a variety of different mechanisms. Diuretics are a particularly common cause, due to associated increases in distal tubular Na^+ delivery and

distal tubular flow rate, in addition to secondary hyperaldosteronism. Thiazides have a greater effect on plasma K^+ concentration than loop diuretics, despite their lesser natriuretic effect. The diuretic effect of thiazides is largely due to inhibition of the Na^+-Cl^- cotransporter NCC in DCT cells. This leads to a direct increase in the delivery of luminal Na^+ to the principal cells immediately downstream in the CNT and cortical CD, which augments Na^+ entry via ENaC, increases the lumen-negative potential difference, and amplifies K^+ secretion. The higher propensity of thiazides to cause hypokalemia may also be secondary to thiazide-associated hypocalciuria, versus the *hypercalciuria* seen with loop diuretics; the increases in downstream luminal calcium in response to loop diuretics inhibit ENaC in principal cells, thus reducing the lumen-negative potential difference and attenuating distal K^+ excretion. High doses of penicillin-related antibiotics (nafcillin, dicloxacillin, ticarcillin, oxacillin, and carbenicillin) can increase obligatory K^+ excretion by acting as nonreabsorbable anions in the distal nephron. Finally, several renal tubular toxins cause renal K^+ and magnesium wasting, leading to hypokalemia and hypomagnesemia; these drugs include aminoglycosides, amphotericin, foscarnet, cisplatin, and ifosfamide (see also “Magnesium Deficiency and Hypokalemia,” below).

Aldosterone activates the ENaC channel in principal cells via multiple synergistic mechanisms, thus increasing the driving force for K^+ excretion. In consequence, increases in aldosterone bioactivity and/or gains in function of aldosterone-dependent signaling pathways are associated with hypokalemia. Increases in circulating aldosterone (hyperaldosteronism) may be primary or secondary. Increased levels of circulating renin in secondary forms of hyperaldosteronism lead to increased angiotensin II and thus aldosterone; renal artery stenosis is perhaps the most frequent cause (Table 63-4). Primary hyperaldosteronism may be genetic or acquired. Hypertension and hypokalemia, due to increases in circulating 11-deoxycorticosterone, occur in patients with congenital adrenal hyperplasia caused by defects in either steroid 11 β -hydroxylase or steroid 17 α -hydroxylase; deficient 11 β -hydroxylase results in associated virilization and other signs of androgen excess, whereas reduced sex steroids in 17 α -hydroxylase deficiency lead to hypogonadism.

The major forms of *isolated* primary genetic hyperaldosteronism are familial hyperaldosteronism type I (FH-I, also known as glucocorticoid-remediable hyperaldosteronism [GRA]) and familial hyperaldosteronism types II and III (FH-II and FH-III), in which aldosterone production is not repressible by exogenous glucocorticoids. FH-I is caused by a chimeric gene duplication between the homologous 11 β -hydroxylase (*CYP11B1*) and aldosterone synthase (*CYP11B2*) genes, fusing the adrenocorticotropic hormone (ACTH)-responsive 11 β -hydroxylase promoter to the coding region of aldosterone synthase; this chimeric gene is under the control of ACTH and thus repressible by glucocorticoids. FH-III is caused by mutations in the *KCNJ5* gene, which encodes the G-protein-activated inward rectifier K^+ channel 4 (GIRK4); these mutations lead to the acquisition of sodium permeability in the mutant GIRK4 channels, causing an exaggerated membrane depolarization in adrenal glomerulosa cells and the activation of voltage-gated calcium channels. The resulting calcium influx is sufficient to produce aldosterone secretion and cell proliferation, leading to adrenal adenomas and hyperaldosteronism.

Acquired causes of primary hyperaldosteronism include aldosterone-producing adenomas (APAs), primary or unilateral adrenal hyperplasia (PAH), idiopathic hyperaldosteronism (IHA) due to bilateral adrenal hyperplasia, and adrenal carcinoma; APA and IHA account for close to 60% and 40%, respectively, of diagnosed hyperaldosteronism. Acquired somatic mutations in *KCNJ5* or less frequently in the *ATP1A1* (an Na^+/K^+ ATPase α subunit) and *ATP2B3* (a Ca^{2+} ATPase) genes can be detected in APAs; as in FH-III (see above), the exaggerated depolarization of adrenal glomerulosa cells caused by these mutations is implicated in the excessive adrenal proliferation and the exaggerated release of aldosterone.

Random testing of plasma renin activity (PRA) and aldosterone is a helpful screening tool in hypokalemic and/or hypertensive patients, with an aldosterone:PRA ratio of >50 suggestive of primary

hyperaldosteronism. Hypokalemia and multiple antihypertensive drugs may alter the aldosterone:PRA ratio by suppressing aldosterone or increasing PRA, leading to a ratio of <50 in patients who do in fact have primary hyperaldosteronism; therefore, the clinical context should always be considered when interpreting these results.

The glucocorticoid cortisol has equal affinity for the mineralocorticoid receptor (MLR) to that of aldosterone, with resultant “mineralocorticoid-like” activity. However, cells in the aldosterone-sensitive distal nephron are protected from this “illicit” activation by the enzyme 11 β -hydroxysteroid dehydrogenase-2 (11 β HSD-2), which converts cortisol to cortisone; cortisone has minimal affinity for the MLR. Recessive loss-of-function mutations in the *11 β HSD-2* gene are thus associated with cortisol-dependent activation of the MLR and the syndrome of apparent mineralocorticoid excess (SAME), encompassing hypertension, hypokalemia, hypercalciuria, and metabolic alkalosis, with suppressed PRA and suppressed aldosterone. A similar syndrome is caused by biochemical inhibition of 11 β HSD-2 by glycyrrhetic/glycyrrhizic acid and/or carbenoxolone. Glycyrrhizic acid is a natural sweetener found in licorice root, typically encountered in licorice and its many guises or as a flavoring agent in tobacco and food products.

Finally, hypokalemia may also occur with systemic increases in glucocorticoids. In Cushing’s syndrome caused by increases in pituitary ACTH (Chap. 406), the incidence of hypokalemia is only 10%, whereas it is 60–100% in patients with ectopic secretion of ACTH, despite a similar incidence of hypertension. Indirect evidence suggests that the activity of renal 11 β HSD-2 is reduced in patients with ectopic ACTH compared with Cushing’s syndrome, resulting in SAME.

Finally, defects in multiple renal tubular transport pathways are associated with hypokalemia. For example, loss-of-function mutations in subunits of the acidifying H^+ -ATPase in alpha-intercalated cells cause hypokalemic distal renal tubular acidosis, as do many acquired disorders of the distal nephron. Liddle’s syndrome is caused by autosomal dominant gain-in-function mutations of ENaC subunits. Disease-associated mutations either activate the channel directly or abrogate aldosterone-inhibited retrieval of ENaC subunits from the plasma membrane; the end result is increased expression of activated ENaC channels at the plasma membrane of principal cells. Patients with Liddle’s syndrome classically manifest severe hypertension with hypokalemia, unresponsive to spironolactone yet sensitive to amiloride. Hypertension and hypokalemia are, however, variable aspects of the Liddle’s phenotype; more consistent features include a blunted aldosterone response to ACTH and reduced urinary aldosterone excretion.

Loss of the transport functions of the TALH and DCT nephron segments causes hereditary hypokalemic alkalosis, Bartter’s syndrome (BS) and Gitelman’s syndrome (GS), respectively. Patients with classic BS typically suffer from polyuria and polydipsia, due to the reduction in renal concentrating ability. They may have an increase in urinary calcium excretion, and 20% are hypomagnesemic. Other features include marked activation of the renin-angiotensin-aldosterone axis. Patients with antenatal BS suffer from a severe systemic disorder characterized by marked electrolyte wasting, polyhydramnios, and hypercalciuria with nephrocalcinosis; renal prostaglandin synthesis and excretion are significantly increased, accounting for much of the systemic symptoms. There are five disease genes for BS, all of them functioning in some aspect of regulated Na^+ , K^+ , and Cl^- transport by the TALH. In contrast, GS is genetically homogeneous, caused almost exclusively by loss-of-function mutations in the thiazide-sensitive Na^+-Cl^- cotransporter of the DCT. Patients with GS are uniformly hypomagnesemic and exhibit marked hypocalciuria, rather than the hypercalciuria typically seen in BS; urinary calcium excretion is thus a critical diagnostic test in GS. GS is a milder phenotype than BS; however, patients with GS may suffer from chondrocalcinosis, an abnormal deposition of calcium pyrophosphate dihydrate (CPPD) in joint cartilage (Chap. 339).

Magnesium Deficiency and Hypokalemia Magnesium depletion has inhibitory effects on muscle Na^+/K^+ -ATPase activity, reducing influx into muscle cells and causing a secondary kaliuresis. In addition,

magnesium depletion causes exaggerated K^+ secretion by the distal nephron; this effect is attributed to a reduction in the magnesium-dependent, intracellular block of K^+ efflux through the secretory K^+ channel of principal cells (ROMK; Fig. 63-4). In consequence, hypomagnesemic patients are clinically refractory to K^+ replacement in the absence of Mg^{2+} repletion. Notably, magnesium deficiency is also a common concomitant of hypokalemia because many disorders of the distal nephron may cause both potassium and magnesium wasting (Chap. 339).

Clinical Features Hypokalemia has prominent effects on cardiac, skeletal, and intestinal muscle cells. In particular, hypokalemia is a major risk factor for both ventricular and atrial arrhythmias. Hypokalemia predisposes to digoxin toxicity by a number of mechanisms, including reduced competition between K^+ and digoxin for shared binding sites on cardiac Na^+/K^+ -ATPase subunits. Electrocardiographic changes in hypokalemia include broad flat T waves, ST depression, and QT prolongation; these are most marked when serum K^+ is <2.7 mmol/L. Hypokalemia can thus be an important precipitant of arrhythmia in patients with additional genetic or acquired causes of QT prolongation. Hypokalemia also results in hyperpolarization of skeletal muscle, thus impairing the capacity to depolarize and contract; weakness and even paralysis may ensue. It also causes a skeletal myopathy and predisposes to rhabdomyolysis. Finally, the paralytic effects of hypokalemia on intestinal smooth muscle may cause intestinal ileus.

The functional effects of hypokalemia on the kidney can include Na^+ - Cl^- and HCO_3^- retention, polyuria, phosphaturia, hypocitraturia, and an activation of renal ammoniogenesis. Bicarbonate retention and other acid-base effects of hypokalemia can contribute to the generation of metabolic alkalosis. Hypokalemic polyuria is due to a combination of central polydipsia and an AVP-resistant renal concentrating defect. Structural changes in the kidney due to hypokalemia include a relatively specific vacuolizing injury to proximal tubular cells, interstitial nephritis, and renal cysts. Hypokalemia also predisposes to acute kidney injury and can lead to end-stage renal disease in patients with longstanding hypokalemia due to eating disorders and/or laxative abuse.

Hypokalemia and/or reduced dietary K^+ are implicated in the pathophysiology and progression of hypertension, heart failure, and stroke. For example, short-term K^+ restriction in healthy humans and patients with essential hypertension induces Na^+ - Cl^- retention and hypertension. Correction of hypokalemia is particularly important in hypertensive patients treated with diuretics, in whom blood pressure improves with the establishment of normokalemia.

Diagnostic Approach The cause of hypokalemia is usually evident from history, physical examination, and/or basic laboratory tests. The history should focus on medications (e.g., laxatives, diuretics, antibiotics), diet and dietary habits (e.g., licorice), and/or symptoms that suggest a particular cause (e.g., periodic weakness, diarrhea). The physical examination should pay particular attention to blood pressure, volume status, and signs suggestive of specific hypokalemic disorders, e.g., hyperthyroidism and Cushing's syndrome. Initial laboratory evaluation should include electrolytes, BUN, creatinine, serum osmolality, Mg^{2+} , Ca^{2+} , a complete blood count, and urinary pH, osmolality, creatinine, and electrolytes (Fig. 63-7). The presence of a non-anion gap acidosis suggests a distal, hypokalemic renal tubular acidosis or diarrhea; calculation of the urinary anion gap can help differentiate these two diagnoses. Renal K^+ excretion can be assessed with a 24-h urine collection; a 24-h K^+ excretion of <15 mmol is indicative of an extrarenal cause of hypokalemia (Fig. 63-7). If only a random, spot urine sample is available, serum and urine osmolality can be used to calculate the transtubular K^+ gradient (TTKG), which should be <3 in the presence of hypokalemia (see also "Hyperkalemia"). Alternatively, a urinary K^+ -to-creatinine ratio of >13 mmol/g creatinine (>1.5 mmol/mmol creatinine) is compatible with excessive renal K^+ excretion. Urine Cl^- is usually decreased in patients with hypokalemia from a nonreabsorbable anion, such as antibiotics or HCO_3^- . The most common causes of chronic hypokalemic alkalosis are surreptitious vomiting, diuretic abuse, and GS; these can be distinguished by the pattern of urinary electrolytes. Hypokalemic patients with vomiting due to

bulimia will thus have a urinary $Cl^- <10$ mmol/L; urine Na^+ , K^+ , and Cl^- are persistently elevated in GS, due to loss of function in the thiazide-sensitive Na^+ - Cl^- cotransporter, but less elevated in diuretic abuse and with greater variability. Urine diuretic screens for loop diuretics and thiazides may be necessary to further exclude diuretic abuse.

Other tests, such as urinary Ca^{2+} , thyroid function tests, and/or PRA and aldosterone levels, may also be appropriate in specific cases. A plasma aldosterone:PRA ratio of >50 , due to suppression of circulating renin and an elevation of circulating aldosterone, is suggestive of hyperaldosteronism. Patients with hyperaldosteronism or apparent mineralocorticoid excess may require further testing, for example adrenal vein sampling (Chap. 406) or the clinically available testing for specific genetic causes (e.g., FH-I, SAME, Liddle's syndrome). Patients with primary aldosteronism should thus be tested for the chimeric FH-I/GRA gene (see above) if they are younger than 20 years of age or have a family history of primary aldosteronism or stroke at a young age (<40 years). Preliminary differentiation of Liddle's syndrome due to mutant ENaC channels from SAME due to mutant 11β HSD-2 (see above), both of which cause hypokalemia and hypertension with aldosterone suppression, can be made on a clinical basis and then confirmed by genetic analysis; patients with Liddle's syndrome should respond to amiloride (ENaC inhibition) but not spironolactone, whereas patients with SAME will respond to spironolactone.

TREATMENT HYPOKALEMIA

The goals of therapy in hypokalemia are to prevent life-threatening and/or serious chronic consequences, to replace the associated K^+ deficit, and to correct the underlying cause and/or mitigate future hypokalemia. The urgency of therapy depends on the severity of hypokalemia, associated clinical factors (e.g., cardiac disease, digoxin therapy), and the rate of decline in serum K^+ . Patients with a prolonged QT interval and/or other risk factors for arrhythmia should be monitored by continuous cardiac telemetry during repletion. Urgent but cautious K^+ replacement should be considered in patients with severe redistributive hypokalemia (plasma K^+ concentration <2.5 mM) and/or when serious complications ensue; however, this approach has a risk of rebound hyperkalemia following acute resolution of the underlying cause. When excessive activity of the sympathetic nervous system is thought to play a dominant role in redistributive hypokalemia, as in TPP, theophylline overdose, and acute head injury, high-dose propranolol (3 mg/kg) should be considered; this nonspecific β -adrenergic blocker will correct hypokalemia without the risk of rebound hyperkalemia.

Oral replacement with K^+ - Cl^- is the mainstay of therapy in hypokalemia. Potassium phosphate, oral or IV, may be appropriate in patients with combined hypokalemia and hypophosphatemia. Potassium bicarbonate or potassium citrate should be considered in patients with concomitant metabolic acidosis. Notably, hypomagnesemic patients are refractory to K^+ replacement alone, such that concomitant Mg^{2+} deficiency should *always* be corrected with oral or intravenous repletion. The deficit of K^+ and the rate of correction should be estimated as accurately as possible; renal function, medications, and comorbid conditions such as diabetes should also be considered, so as to gauge the risk of overcorrection. In the absence of abnormal K^+ redistribution, the total deficit correlates with serum K^+ , such that serum K^+ drops by approximately 0.27 mM for every 100-mmol reduction in total-body stores; loss of 400–800 mmol of total-body K^+ results in a reduction in serum K^+ by approximately 2.0 mM. Notably, given the delay in redistributing potassium into intracellular compartments, this deficit must be replaced gradually over 24–48 h, with frequent monitoring of plasma K^+ concentration to avoid transient overrepletion and transient hyperkalemia.

The use of intravenous administration should be limited to patients unable to use the enteral route or in the setting of severe complications (e.g., paralysis, arrhythmia). Intravenous K^+ - Cl^- should always be administered in saline solutions, rather than dextrose, because the dextrose-induced increase in insulin can acutely

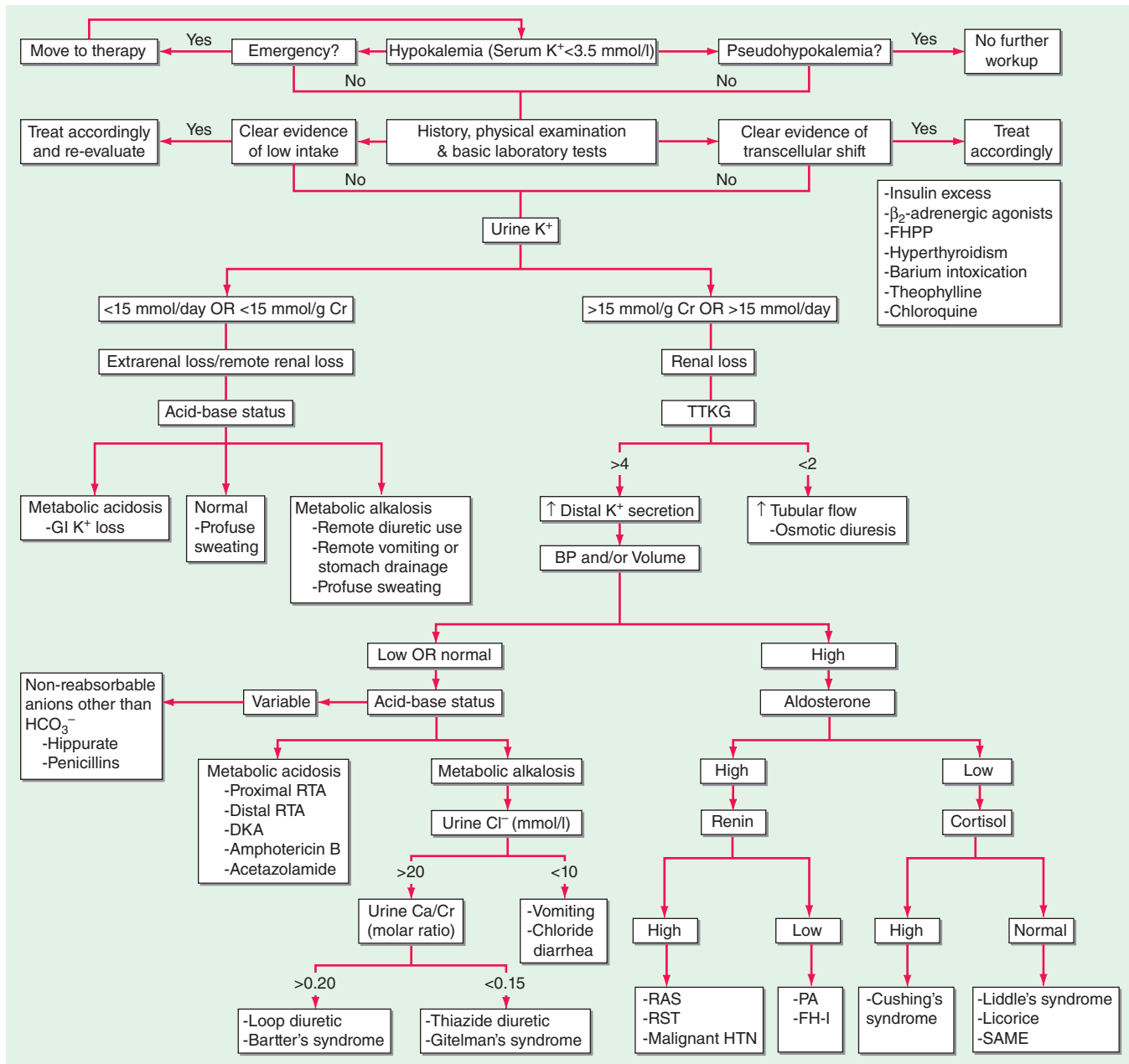


FIGURE 63-7 The diagnostic approach to hypokalemia. See text for details. AME, apparent mineralocorticoid excess; BP, blood pressure; CCD, cortical collecting duct; DKA, diabetic ketoacidosis; FH-I, familial hyperaldosteronism type I; FHPP, familial hypokalemic periodic paralysis; GI, gastrointestinal; GRA, glucocorticoid remediable aldosteronism; HTN, hypertension; PA, primary aldosteronism; RAS, renal artery stenosis; RST, renin-secreting tumor; RTA, renal tubular acidosis; SAME, syndrome of apparent mineralocorticoid excess; TTKG, transtubular potassium gradient. (Used with permission from DB Mount, K Zandi-Nejad K: Disorders of potassium balance, in Brenner and Rector's *The Kidney*, 8th ed, BM Brenner [ed]. Philadelphia, W.B. Saunders & Company, 2008, pp 547-587.)

exacerbate hypokalemia. The peripheral intravenous dose is usually 20–40 mmol of K^+Cl^- per liter; higher concentrations can cause localized pain from chemical phlebitis, irritation, and sclerosis. If hypokalemia is severe (< 2.5 mmol/L) and/or critically symptomatic, intravenous K^+Cl^- can be administered through a central vein with cardiac monitoring in an intensive care setting, at rates of 10–20 mmol/h; higher rates should be reserved for acutely life-threatening complications. The absolute amount of administered K^+ should be restricted (e.g., 20 mmol in 100 mL of saline solution) to prevent inadvertent infusion of a large dose. Femoral veins are preferable, because infusion through internal jugular or subclavian central lines can acutely increase the local concentration of K^+ and affect cardiac conduction.

Strategies to minimize K^+ losses should also be considered. These measures may include minimizing the dose of non- K^+ -sparing diuretics, restricting Na^+ intake, and using clinically appropriate

combinations of non- K^+ -sparing and K^+ -sparing medications (e.g., loop diuretics with angiotensin-converting enzyme inhibitors).

HYPERKALEMIA

Hyperkalemia is defined as a plasma potassium level of 5.5 mM, occurring in up to 10% of hospitalized patients; severe hyperkalemia (> 6.0 mM) occurs in approximately 1%, with a significantly increased risk of mortality. Although redistribution and reduced tissue uptake can acutely cause hyperkalemia, a decrease in renal K^+ excretion is the most frequent underlying cause (Table 63-5). Excessive intake of K^+ is a rare cause, given the adaptive capacity to increase renal secretion; however, dietary intake can have a major effect in susceptible patients, e.g., diabetics with hyporeninemic hypoaldosteronism and chronic kidney disease. Drugs that impact on the renin-angiotensin-aldosterone axis are also a major cause of hyperkalemia.

TABLE 63-5 CAUSES OF HYPERKALEMIA

- I. Pseudohyperkalemia
 - A. Cellular efflux; thrombocytosis, erythrocytosis, leukocytosis, in vitro hemolysis
 - B. Hereditary defects in red cell membrane transport
- II. Intra- to extracellular shift
 - A. Acidosis
 - B. Hyperosmolality; radiocontrast, hypertonic dextrose, mannitol
 - C. β_2 -Adrenergic antagonists (noncardioselective agents)
 - D. Digoxin and related glycosides (yellow oleander, foxglove, bufadienolide)
 - E. Hyperkalemic periodic paralysis
 - F. Lysine, arginine, and ϵ -aminocaproic acid (structurally similar, positively charged)
 - G. Succinylcholine; thermal trauma, neuromuscular injury, disuse atrophy, mucositis, or prolonged immobilization
 - H. Rapid tumor lysis
- III. Inadequate excretion
 - A. Inhibition of the renin-angiotensin-aldosterone axis; \uparrow risk of hyperkalemia when used in combination
 1. Angiotensin-converting enzyme (ACE) inhibitors
 2. Renin inhibitors; aliskiren (in combination with ACE inhibitors or angiotensin receptor blockers [ARBs])
 3. Angiotensin receptor blockers (ARBs)
 4. Blockade of the mineralocorticoid receptor: spironolactone, eplerenone, drospirenone
 5. Blockade of the epithelial sodium channel (ENaC): amiloride, triamterene, trimethoprim, pentamidine, nafamostat
 - B. Decreased distal delivery
 1. Congestive heart failure
 2. Volume depletion
 - C. Hyporeninemic hypoaldosteronism
 1. Tubulointerstitial diseases: systemic lupus erythematosus (SLE), sickle cell anemia, obstructive uropathy
 2. Diabetes, diabetic nephropathy
 3. Drugs: nonsteroidal anti-inflammatory drugs (NSAIDs), cyclooxygenase 2 (COX2) inhibitors, β -blockers, cyclosporine, tacrolimus
 4. Chronic kidney disease, advanced age
 5. Pseudohypoaldosteronism type II: defects in WNK1 or WNK4 kinases, Kelch-like 3 (KLHL3), or Cullin 3 (CUL3)
 - D. Renal resistance to mineralocorticoid
 1. Tubulointerstitial diseases: SLE, amyloidosis, sickle cell anemia, obstructive uropathy, post-acute tubular necrosis
 2. Hereditary: pseudohypoaldosteronism type I; defects in the mineralocorticoid receptor or the epithelial sodium channel (ENaC)
 - E. Advanced renal insufficiency
 1. Chronic kidney disease
 2. End-stage renal disease
 3. Acute oliguric kidney injury
 - F. Primary adrenal insufficiency
 1. Autoimmune: Addison's disease, polyglandular endocrinopathy
 2. Infectious: HIV, cytomegalovirus, tuberculosis, disseminated fungal infection
 3. Infiltrative: amyloidosis, malignancy, metastatic cancer
 4. Drug-associated: heparin, low-molecular-weight heparin
 5. Hereditary: adrenal hypoplasia congenita, congenital lipoid adrenal hyperplasia, aldosterone synthase deficiency
 6. Adrenal hemorrhage or infarction, including in antiphospholipid syndrome

Pseudohyperkalemia Hyperkalemia should be distinguished from factitious hyperkalemia or “pseudohyperkalemia,” an artifactual increase in serum K^+ due to the release of K^+ during or after venipuncture. Pseudohyperkalemia can occur in the setting of excessive muscle activity during venipuncture (e.g., fist clenching), a marked increase in cellular elements (thrombocytosis, leukocytosis, and/or erythrocytosis) with in vitro efflux of K^+ , and acute anxiety during venipuncture with respiratory alkalosis and redistributive hyperkalemia. Cooling of blood following venipuncture is another cause, due to reduced cellular uptake; the converse is the increased uptake of K^+ by cells at high ambient temperatures, leading to normal values for hyperkalemic patients and/or to spurious hypokalemia in normokalemic patients. Finally, there are multiple genetic subtypes of hereditary pseudohyperkalemia, caused by increases in the passive K^+ permeability of erythrocytes. For example, causative mutations have been described in the red cell anion exchanger (AE1, encoded by the *SLC4A1* gene), leading to reduced red cell anion transport, hemolytic anemia, the acquisition of a novel AE1-mediated K^+ leak, and pseudohyperkalemia.

Redistribution and Hyperkalemia Several different mechanisms can induce an efflux of intracellular K^+ and hyperkalemia. Acidemia is associated with cellular uptake of H^+ and an associated efflux of K^+ ; it is thought that this effective K^+ - H^+ exchange serves to help maintain extracellular pH. Notably, this effect of acidosis is limited to non-anion gap causes of metabolic acidosis and, to a lesser extent, respiratory causes of acidosis; hyperkalemia due to an acidosis-induced shift of potassium from the cells into the ECF does *not* occur in the anion gap acidoses lactic acidosis and ketoacidosis. Hyperkalemia due to hypertonic mannitol, hypertonic saline, and intravenous immune globulin is generally attributed to a “solvent drag” effect, as water moves out of cells along the osmotic gradient. Diabetics are also prone to osmotic hyperkalemia in response to intravenous hypertonic glucose, when given without adequate insulin. Cationic amino acids, specifically lysine, arginine, and the structurally related drug epsilon-aminocaproic acid, cause efflux of K^+ and hyperkalemia, through an effective cation- K^+ exchange of unknown identity and mechanism. Digoxin inhibits Na^+/K^+ -ATPase and impairs the uptake of K^+ by

skeletal muscle, such that digoxin overdose predictably results in hyperkalemia. Structurally related glycosides are found in specific plants (e.g., yellow oleander, foxglove) and in the cane toad, *Bufo marinus* (bufadienolide); ingestion of these substances and extracts thereof can also cause hyperkalemia. Finally, fluoride ions also inhibit Na^+/K^+ -ATPase, such that fluoride poisoning is typically associated with hyperkalemia.

Succinylcholine depolarizes muscle cells, causing an efflux of K^+ through acetylcholine receptors (AChRs). The use of this agent is contraindicated in patients who have sustained thermal trauma, neuromuscular injury, disuse atrophy, mucositis, or prolonged immobilization. These disorders share a marked increase and redistribution of AChRs at the plasma membrane of muscle cells; depolarization of these upregulated AChRs by succinylcholine leads to an exaggerated efflux of K^+ through the receptor-associated cation channels, resulting in acute hyperkalemia.

Hyperkalemia Caused by Excess Intake or Tissue Necrosis Increased intake of even small amounts of K^+ may provoke severe hyperkalemia in patients with predisposing factors; hence, an assessment of dietary intake is crucial. Foods rich in potassium include tomatoes, bananas, and citrus fruits; occult sources of K^+ , particularly K^+ -containing salt substitutes, may also contribute significantly. Iatrogenic causes include simple overreplacement with K^+-Cl^- or the administration of a potassium-containing medication (e.g., K^+ -penicillin) to a susceptible patient. Red cell transfusion is a well-described cause of hyperkalemia, typically in the setting of massive transfusions. Finally, severe tissue necrosis, as in acute tumor lysis syndrome and rhabdomyolysis, will predictably cause hyperkalemia from the release of intracellular K^+ .

Hypoaldosteronism and Hyperkalemia Aldosterone release from the adrenal gland may be reduced by hyporeninemic hypoaldosteronism, medications, primary hypoaldosteronism, or isolated deficiency of ACTH (secondary hypoaldosteronism). Primary hypoaldosteronism may be genetic or acquired (Chap. 406) but is commonly caused by autoimmunity, either in Addison's disease or in the context of a polyglandular endocrinopathy. HIV has surpassed tuberculosis as the most important infectious cause of adrenal insufficiency. The adrenal involvement in HIV disease is usually subclinical; however, adrenal insufficiency may be precipitated by stress, drugs such as ketoconazole that inhibit steroidogenesis, or the acute withdrawal of steroid agents such as megestrol.

Hyporeninemic hypoaldosteronism is a very common predisposing factor in several overlapping subsets of hyperkalemic patients: diabetics, the elderly, and patients with renal insufficiency. Classically, patients should have suppressed PRA and aldosterone; approximately 50% have an associated acidosis, with a reduced renal excretion of NH_4^+ , a positive urinary anion gap, and urine pH <5.5. Most patients are volume expanded, with secondary increases in circulating atrial natriuretic peptide (ANP) that inhibit both renal renin release and adrenal aldosterone release.

Renal Disease and Hyperkalemia Chronic kidney disease and end-stage kidney disease are very common causes of hyperkalemia, due to the associated deficit or absence of functioning nephrons. Hyperkalemia is more common in oliguric acute kidney injury; distal tubular flow rate and Na^+ delivery are less limiting factors in nonoliguric patients. Hyperkalemia out of proportion to GFR can also be seen in the context of tubulointerstitial disease that affects the distal nephron, such as amyloidosis, sickle cell anemia, interstitial nephritis, and obstructive uropathy.

Hereditary renal causes of hyperkalemia have overlapping clinical features with hypoaldosteronism, hence the diagnostic label *pseudohypoaldosteronism* (PHA). PHA type I (PHA-I) has both an autosomal recessive and an autosomal dominant form. The autosomal dominant form is due to loss-of-function mutations in the MLR; the recessive form is caused by various combinations of mutations in the three subunits of ENaC, resulting in impaired Na^+ channel activity in principal cells and other tissues. Patients with recessive PHA-I suffer from lifelong salt wasting, hypotension, and hyperkalemia, whereas the phenotype of autosomal dominant PHA-I due to MLR dysfunction improves in adulthood. PHA type II (PHA-II; also known as *hereditary hypertension with hyperkalemia*) is in every respect the mirror image of

GS caused by loss of function in NCC, the thiazide-sensitive Na^+-Cl^- cotransporter (see above); the clinical phenotype includes hypertension, hyperkalemia, hyperchloremic metabolic acidosis, suppressed PRA and aldosterone, hypercalciuria, and reduced bone density. PHA-II thus behaves like a gain of function in NCC, and treatment with thiazides results in resolution of the entire clinical phenotype. However, the NCC gene is not directly involved in PHA-II, which is caused by mutations in the WNK1 and WNK4 serine-threonine kinases or the upstream Kelch-like 3 (KLHL3) and Cullin 3 (CUL3), two components of an E3 ubiquitin ligase complex that regulates these kinases; these proteins collectively regulate NCC activity, with PHA-II-associated activation of the transporter.

Medication-Associated Hyperkalemia Most medications associated with hyperkalemia cause inhibition of some component of the renin-angiotensin-aldosterone axis. ACE inhibitors, angiotensin receptor blockers, renin inhibitors, and MLRs are predictable and common causes of hyperkalemia, particularly when prescribed in combination. The oral contraceptive agent Yasmin-28 contains the progestin drospirenone, which inhibits the MLR and can cause hyperkalemia in susceptible patients. Cyclosporine, tacrolimus, NSAIDs, and cyclooxygenase 2 (COX2) inhibitors cause hyperkalemia by multiple mechanisms, but share the ability to cause hyporeninemic hypoaldosteronism. Notably, most drugs that affect the renin-angiotensin-aldosterone axis also block the local adrenal response to hyperkalemia, thus attenuating the *direct* stimulation of aldosterone release by increased plasma K^+ concentration.

Inhibition of apical ENaC activity in the distal nephron by amiloride and other K^+ -sparing diuretics results in hyperkalemia, often with a voltage-dependent hyperchloremic acidosis and/or hypovolemic hyponatremia. Amiloride is structurally similar to the antibiotics trimethoprim (TMP) and pentamidine, which also block ENaC; risk factors for TMP-associated hyperkalemia include the administered dose, renal insufficiency, and hyporeninemic hypoaldosteronism. Indirect inhibition of ENaC at the plasma membrane is also a cause of drug-associated hyperkalemia; nafamostat, a protease inhibitor used in some countries for the management of pancreatitis, inhibits aldosterone-induced renal proteases that activate ENaC by proteolytic cleavage.

Clinical Features Hyperkalemia is a medical emergency due to its effects on the heart. Cardiac arrhythmias associated with hyperkalemia include sinus bradycardia, sinus arrest, slow idioventricular rhythms, ventricular tachycardia, ventricular fibrillation, and asystole. Mild increases in extracellular K^+ affect the repolarization phase of the cardiac action potential, resulting in changes in T-wave morphology; further increase in plasma K^+ concentration depresses intracardiac conduction, with progressive prolongation of the PR and QRS intervals. Severe hyperkalemia results in loss of the P wave and a progressive widening of the QRS complex; development of a sine-wave sinoventricular rhythm suggests impending ventricular fibrillation or asystole. Hyperkalemia can also cause a type I Brugada pattern in the electrocardiogram (ECG), with a pseudo-right bundle branch block and persistent coved ST segment elevation in at least two precordial leads. This hyperkalemic Brugada's sign occurs in critically ill patients with severe hyperkalemia and can be differentiated from genetic Brugada's syndrome by an absence of P waves, marked QRS widening, and an abnormal QRS axis. Classically, the electrocardiographic manifestations in hyperkalemia progress from tall peaked T waves (5.5–6.5 mM), to a loss of P waves (6.5–7.5 mM) to a widened QRS complex (7.0–8.0 mM), and, ultimately, to a sine wave pattern (>8.0 mM). However, these changes are notoriously insensitive, particularly in patients with chronic kidney disease or end-stage renal disease.

Hyperkalemia from a variety of causes can also present with ascending paralysis, denoted *secondary hyperkalemic paralysis* to differentiate it from familial hyperkalemic periodic paralysis (HYPP). The presentation may include diaphragmatic paralysis and respiratory failure. Patients with familial HYPP develop myopathic weakness during hyperkalemia induced by increased K^+ intake or rest after heavy exercise. Depolarization of skeletal muscle by hyperkalemia unmasks an inactivation defect in skeletal Na^+ channel; autosomal dominant mutations in the *SCN4A* gene encoding this channel are the predominant cause.

Within the kidney, hyperkalemia has negative effects on the ability to excrete an acid load, such that hyperkalemia per se can contribute to metabolic acidosis. This defect appears to be due in part to competition between K^+ and NH_4^+ for reabsorption by the TALH and subsequent countercurrent multiplication, ultimately reducing the medullary gradient for NH_3/NH_4 excretion by the distal nephron. Regardless of the underlying mechanism, restoration of normokalemia can, in many instances, correct hyperkalemic metabolic acidosis.

Diagnostic Approach The first priority in the management of hyperkalemia is to assess the need for emergency treatment, followed by a comprehensive workup to determine the cause (Fig. 63-8). History and physical examination should focus on medications, diet and dietary supplements, risk factors for kidney failure, reduction in

urine output, blood pressure, and volume status. Initial laboratory tests should include electrolytes, BUN, creatinine, serum osmolality, Mg^{2+} and Ca^{2+} , a complete blood count, and urinary pH, osmolality, creatinine, and electrolytes. A urine Na^+ concentration of <20 mM indicates that distal Na^+ delivery is a limiting factor in K^+ excretion; volume repletion with 0.9% saline or treatment with furosemide may be effective in reducing plasma K^+ concentration. Serum and urine osmolality are required for calculation of the transtubular K^+ gradient (TTKG) (Fig. 63-8). The expected values of the TTKG are largely based on historical data, and are <3 in the presence of hypokalemia and $>7-8$ in the presence of hyperkalemia.

$$TTKG = \frac{[K^+]_{urine} \times Osm_{serum}}{[K^+]_{serum} \times Osm_{urine}}$$

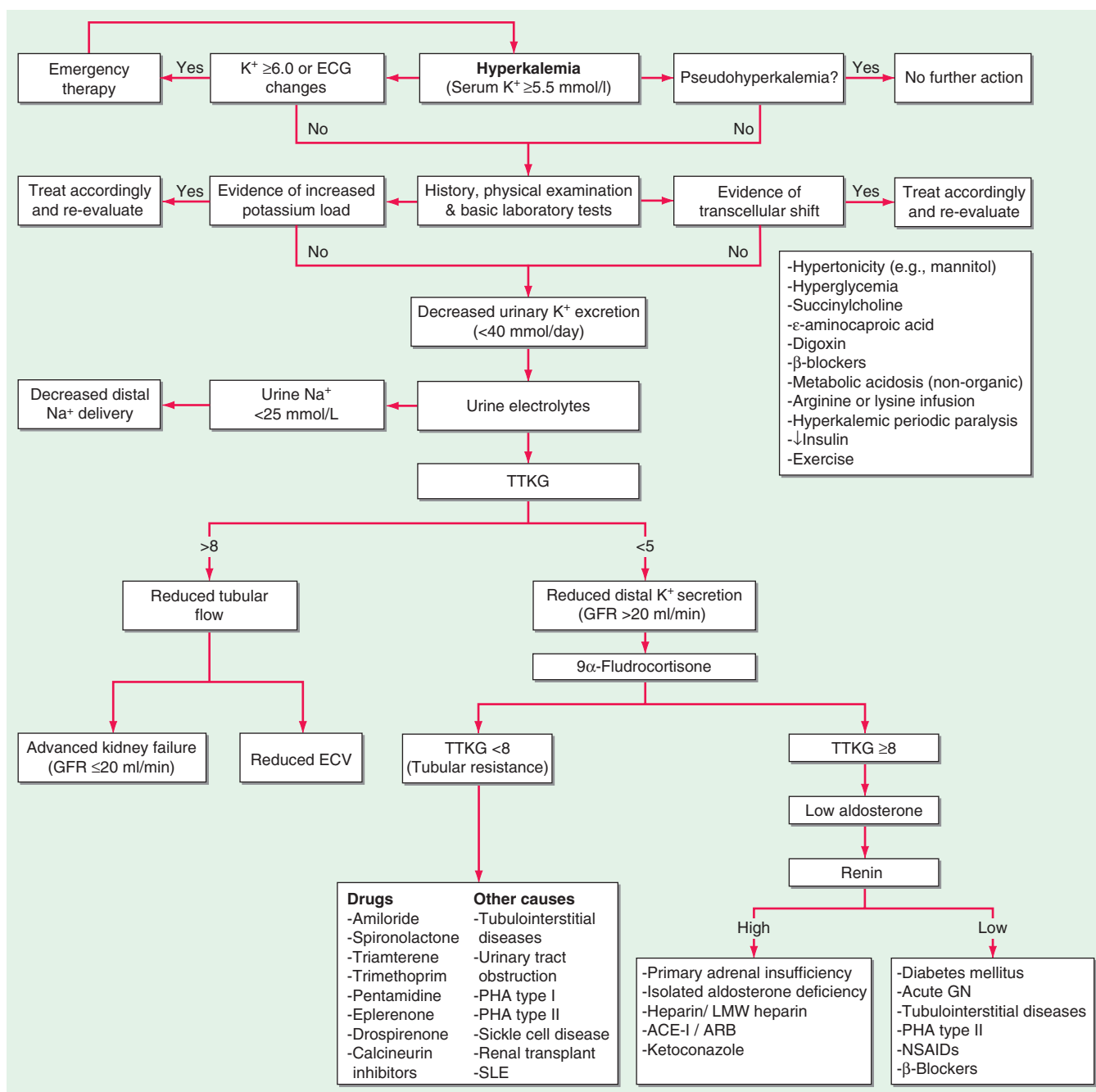


FIGURE 63-8 The diagnostic approach to hyperkalemia. See text for details. ACE-I, angiotensin-converting enzyme inhibitor; ARB, angiotensin II receptor blocker; CCD, cortical collecting duct; ECG, electrocardiogram; ECV, effective circulatory volume; GFR, glomerular filtration rate; GN, glomerulonephritis; HIV, human immunodeficiency virus; LMW heparin, low-molecular-weight heparin; NSAIDs, nonsteroidal anti-inflammatory drugs; PHA, pseudohypoaldosteronism; SLE, systemic lupus erythematosus; TTKG, transtubular potassium gradient. (Used with permission from DB Mount, K Zandi-Nejad K: Disorders of potassium balance, in Brenner and Rector's *The Kidney*, 8th ed, BM Brenner [ed]. Philadelphia, W.B. Saunders & Company, 2008, pp 547-587.)

Electrocardiographic manifestations of hyperkalemia should be considered a medical emergency and treated urgently. However, patients with significant hyperkalemia (plasma K^+ concentration ≥ 6.5 mM) in the absence of ECG changes should also be aggressively managed, given the limitations of ECG changes as a predictor of cardiac toxicity. Urgent management of hyperkalemia includes admission to the hospital, continuous cardiac monitoring, and immediate treatment. The treatment of hyperkalemia is divided into three stages:

1. *Immediate antagonism of the cardiac effects of hyperkalemia.* Intravenous calcium serves to protect the heart, whereas other measures are taken to correct hyperkalemia. Calcium raises the action potential threshold and reduces excitability, without changing the resting membrane potential. By restoring the difference between resting and threshold potentials, calcium reverses the depolarization blockade due to hyperkalemia. The recommended dose is 10 mL of 10% calcium gluconate (3–4 mL of calcium chloride), infused intravenously over 2–3 min with cardiac monitoring. The effect of the infusion starts in 1–3 min and lasts 30–60 min; the dose should be repeated if there is no change in ECG findings or if they recur after initial improvement. Hypercalcemia potentiates the cardiac toxicity of digoxin; hence, intravenous calcium should be used with extreme caution in patients taking this medication; if judged necessary, 10 mL of 10% calcium gluconate can be added to 100 mL of 5% dextrose in water and infused over 20–30 min to avoid acute hypercalcemia.
2. *Rapid reduction in plasma K^+ concentration by redistribution into cells.* Insulin lowers plasma K^+ concentration by shifting K^+ into cells. The recommended dose is 10 units of intravenous regular insulin followed immediately by 50 mL of 50% dextrose ($D_{50}W$, 25 g of glucose total); the effect begins in 10–20 min, peaks at 30–60 min, and lasts for 4–6 h. Bolus $D_{50}W$ without insulin is *never* appropriate, given the risk of acutely worsening hyperkalemia due to the osmotic effect of hypertonic glucose. Hypoglycemia is common with insulin plus glucose; hence, this should be followed by an infusion of 10% dextrose at 50–75 mL/h, with close monitoring of plasma glucose concentration. In hyperkalemic patients with glucose concentrations of ≥ 200 –250 mg/dL, insulin should be administered *without* glucose, again with close monitoring of glucose concentrations.

β_2 -agonists, most commonly albuterol, are effective but underused agents for the acute management of hyperkalemia. Albuterol and insulin with glucose have an additive effect on plasma K^+ concentration; however, ~20% of patients with end-stage renal disease (ESRD) are resistant to the effect of β_2 -agonists; hence, these drugs should not be used without insulin. The recommended dose for inhaled albuterol is 10–20 mg of nebulized albuterol in 4 mL of normal saline, inhaled over 10 min; the effect starts at about 30 min, reaches its peak at about 90 min, and lasts for 2–6 h. Hyperglycemia is a side effect, along with tachycardia. β_2 -Agonists should be used with caution in hyperkalemic patients with known cardiac disease.

Intravenous bicarbonate has no role in the acute treatment of hyperkalemia, but may slowly attenuate hyperkalemia with sustained administration over several hours. It should not be given repeatedly as a hypertonic intravenous bolus of undiluted ampules, given the risk of associated hypernatremia, but should instead be infused in an isotonic or hypotonic fluid (e.g., 150 mEq in 1 L of D_5W). In patients with metabolic acidosis, a delayed drop in plasma K^+ concentration can be seen after 4–6 h of isotonic bicarbonate infusion.

3. *Removal of potassium.* This is typically accomplished using cation exchange resins, diuretics, and/or dialysis. The cation exchange resin sodium polystyrene sulfonate (SPS) exchanges Na^+ for K^+ in the gastrointestinal tract and increases the fecal excretion of K^+ ; alternative calcium-based resins, when available, may

be more appropriate in patients with an increased ECFV. The recommended dose of SPS is 15–30 g of powder, almost always given in a premade suspension with 33% sorbitol. The effect of SPS on plasma K^+ concentration is slow; the full effect may take up to 24 h and usually requires repeated doses every 4–6 h. Intestinal necrosis, typically of the colon or ileum, is a rare but usually fatal complication of SPS. Intestinal necrosis is more common in patients administered SPS via enema and/or in patients with reduced intestinal motility (e.g., in the postoperative state or after treatment with opioids). The coadministration of SPS with sorbitol appears to increase the risk of intestinal necrosis; however, this complication can also occur with SPS alone. If SPS without sorbitol is not available, clinicians must consider whether treatment with SPS in sorbitol is absolutely necessary. The low but real risk of intestinal necrosis with SPS, which can sometimes be the only available or appropriate therapy for the removal of potassium, must be weighed against the delayed onset of efficacy. Whenever possible, alternative therapies for the acute management of hyperkalemia (i.e., aggressive redistributive therapy, isotonic bicarbonate infusion, diuretics, and/or hemodialysis) should be used instead of SPS.

Therapy with intravenous saline may be beneficial in hypovolemic patients with oliguria and decreased distal delivery of Na^+ , with the associated reductions in renal K^+ excretion. Loop and thiazide diuretics can be used to reduce plasma K^+ concentration in volume-replete or hypervolemic patients with sufficient renal function for a diuretic response; this may need to be combined with intravenous saline or isotonic bicarbonate to achieve or maintain euolemia.

Hemodialysis is the most effective and reliable method to reduce plasma K^+ concentration; peritoneal dialysis is considerably less effective. Patients with acute kidney injury require temporary, urgent venous access for hemodialysis, with the attendant risks; in contrast, patients with ESRD or advanced chronic kidney disease may have a preexisting venous access. The amount of K^+ removed during hemodialysis depends on the relative distribution of K^+ between ICF and ECF (potentially affected by prior therapy for hyperkalemia), the type and surface area of the dialyzer used, dialysate and blood flow rates, dialysate flow rate, dialysis duration, and the plasma-to-dialysate K^+ gradient.

David B. Mount, Thomas D. DuBose, Jr.

CASE 1

A 23-year-old woman was admitted with a 3-day history of fever, cough productive of blood-tinged sputum, confusion, and orthostasis. Past medical history included type 1 diabetes mellitus. A physical examination in the emergency department indicated postural hypotension, tachycardia, and Kussmaul respiration. The breath was noted to smell of “acetone.” Examination of the thorax suggested consolidation in the right lower lobe.

Laboratory Data	Value	Units
Sodium	130	meq/L
Potassium	5.0	meq/L
Chloride	96	meq/L
CO ₂	14	meq/L
Blood urea nitrogen (BUN)	20	mg/dL
Creatinine	1.3	mg/dL
Glucose	450	mg/dL
Arterial Blood Gases	On Room Air	
pH	7.39	
Pco ₂	24	mmHg
Pao ₂	89	mmHg
[HCO ₃ ⁻]	14	meq/L
Anion gap	20	meq/L
Urinalysis		
Urine ketones	Positive 4+	
Glucose	Positive 4+	
Serum Ketones	Strongly positive 1:8	
Chest X-Ray		
	Pneumonic infiltrate, right lower lobe	

APPROACH TO DIAGNOSIS

The diagnosis of the acid-base disorder should proceed in a stepwise fashion:

1. The normal anion gap (AG) is 8–10 meq/L, but in this case, the AG is elevated (20 meq/L). Therefore, the change in AG (Δ AG) = ~10 meq/L.
2. Compare the Δ AG and the Δ [HCO₃⁻]. In this case, the Δ AG, as noted above, is 10, and the Δ [HCO₃⁻] (25 – 14) is 11. Therefore, the *increment in the AG* is approximately equal to the *decrement in bicarbonate*.
3. Estimate the respiratory compensatory response. In this case, the predicted Paco₂ for an [HCO₃⁻] of 14 should be approximately 29 mmHg. This value is obtained by adding 15 to the measured [HCO₃⁻] (15 + 14 = 29) or by calculating the predicted Paco₂ from the Winter equation: $1.5 \times [\text{HCO}_3^-] + 8$. In either case, the predicted value for Paco₂ of 29 is significantly higher than the measured value of 24. Therefore, the prevailing Paco₂ exceeds the range for compensation alone and is too low, indicating a superimposed respiratory alkalosis.
4. Therefore, this patient has a *mixed acid-base disturbance* with two components: (a) high AG acidosis secondary to ketoacidosis and (b) respiratory alkalosis (which was secondary to community-acquired

pneumonia in this case). The latter resulted in an additional component of hyperventilation that exceeded the compensatory response driven by metabolic acidosis, explaining the normal pH. The finding of respiratory alkalosis in the setting of a high AG acidosis suggests another cause of the respiratory component. Respiratory alkalosis frequently accompanies community-acquired pneumonia.

The clinical features in this case include hyperglycemia, hypovolemia, ketoacidosis, central nervous system (CNS) signs of confusion, and superimposed pneumonia. This clinical scenario is consistent with diabetic ketoacidosis (DKA) developing in a patient with known type 1 diabetes mellitus. Infections in DKA are common and may be a precipitating feature in the development of ketoacidosis.

The diagnosis of DKA is usually not challenging but should be considered in all patients with an elevated AG and metabolic acidosis. Hyperglycemia and ketonemia (positive acetoacetate at a dilution of 1:8 or greater) are sufficient criteria for diagnosis in patients with type 1 diabetes mellitus. The Δ [HCO₃⁻] should approximate the increase in the plasma AG (Δ AG), but this equality can be modified by several factors. For example, the Δ AG will often decrease with IV hydration, as glomerular filtration increases and ketones are excreted into the urine. The decrement in plasma sodium is the result of hyperglycemia, which induces the movement of water into the extracellular compartment from the intracellular compartment of cells that require insulin for the transport of glucose. Additionally, a natriuresis occurs in response to an osmotic diuresis associated with hyperglycemia. Moreover, in patients with DKA, thirst is very common and water ingestion often continues. The plasma potassium concentration is usually mildly elevated, but in the face of acidosis, and as a result of the ongoing osmotic diuresis, a significant total-body deficit of potassium is almost always present. Recognition of the total-body deficit of potassium is critically important. The inclusion of potassium replacement in the therapeutic regimen at the appropriate time and with the appropriate indications (see below) is essential. Volume depletion is a very common finding in DKA and is a pivotal component in the pathogenesis of the disorder.

APPROACH TO MANAGEMENT

Patients with DKA often have a sustained and significant deficit of sodium, potassium, water, bicarbonate, and phosphate. The general approach to treatment requires attention to all of these abnormalities. Successful treatment of DKA involves a stepwise approach, as follows:

1. *Replace extracellular fluid (ECF) volume deficits.* Because most patients present with actual or relative hypotension and, at times, impending shock, the initial fluid administered should be 0.9% NaCl infused rapidly until the systolic blood pressure is >100 mmHg or until 2–3 L cumulatively have been administered. During the initial 2–3 h of infusion of saline, the decline in blood glucose can be accounted for by dilution and increased renal excretion. Glucose should be added to the infusion as D₅ normal saline (NS) or D₅ 0.45% NS once the plasma glucose declines to 230 mg/dL or below.
2. *Abate the production of ketoacids.* Regular insulin is required during DKA as an initial bolus of 0.1 U/kg body weight (BW) IV, followed immediately by a continuous infusion of 0.1 U/kg BW per hour in NS. The effectiveness of IV insulin (not subcutaneous) can be tracked by observing the decline in plasma ketones. Because the increment in the AG above the normal value of 10 meq/L represents accumulated ketoacids in DKA, the disappearance of ketoacid anions is reflected by the narrowing and eventual correction of the AG. Typically, the plasma AG returns to normal within 8–12 h.
3. *Replace potassium deficits.* Although patients with DKA often have hyperkalemia due to insulin deficiency, they are usually severely K⁺ depleted. KCl (20 meq/L) should be added to each liter of IV fluids when urine output is established and insulin has been administered.
4. *Correct the metabolic acidosis.* The plasma bicarbonate concentration will usually not increase for several hours because of dilution from administered IV NaCl. The plasma [HCO₃⁻] approaches 18 meq/L once ketoacidosis disappears. Sodium bicarbonate therapy is often not recommended or necessary and is contraindicated

- for children. Bicarbonate is administered to adults with DKA for extreme acidemia (pH <7.1); for elderly patients (>70 years old), a threshold pH of 7.20 is recommended. Sodium bicarbonate, if administered, should only be given in small amounts. Because ketoacids are metabolized in response to insulin therapy, bicarbonate will be added to the ECF as ketoacids are converted. Overshoot alkalosis may occur from the combination of exogenously administered sodium bicarbonate plus metabolic production of bicarbonate.
- Phosphate.** In the first 6–8 h of therapy, it may be necessary to infuse potassium with phosphate because of the unmasking of phosphate depletion during combined insulin and glucose therapy. The latter drives phosphate into the cell. Therefore, in patients with DKA, the plasma phosphate level should be followed closely, but phosphate should never be replaced empirically. Phosphate should be administered to patients with a declining plasma phosphate once the phosphate level declines into the low-normal level. Therapy is advisable in the form of potassium phosphate at a rate of 6 mmol/h.
 - Always seek underlying factors,** such as infection, myocardial infarction, pancreatitis, cessation of insulin therapy, or other events, responsible for initiating DKA. The case presented here is illustrative of this common scenario.
 - Volume overexpansion with IV fluid administration** is not uncommon and contributes to the development of hyperchloremic acidosis during the later stages of treatment of DKA. *Volume overexpansion should be avoided.*

CASE 2

A 25-year-old man with a 6-year history of HIV/AIDS complicated recently by *Pneumocystis jiroveci* pneumonia (PCP) was treated with intravenous trimethoprim-sulfamethoxazole (20 mg trimethoprim/kg per day). On day 4 of treatment, the following laboratory data were obtained:

Laboratory Data	Units	Plasma	Urine
Na ⁺	meq/L	135	60
K ⁺	meq/L	6.5	15
Cl ⁻	meq/L	110	43
HCO ₃ ⁻	meq/L	15	0
pH		7.30	5.5
BUN	mg/dL	14	—
Creatinine	mg/dL	0.9	—
Osmolality	mOsm/kg H ₂ O	268	270

APPROACH TO DIAGNOSIS

What caused the hyperkalemia and metabolic acidosis in this patient? What other medications may be associated with a similar presentation? How does one use the urine electrolyte data to determine if the hyperkalemia is of renal origin or due to a shift from the cell to the extracellular compartment?

Hyperkalemia occurs in 15–20% of hospitalized patients with HIV/AIDS. The usual causes are either adrenal insufficiency, the syndrome of hyporeninemic hypoaldosteronism, or one of several drugs, including trimethoprim, pentamidine, nonsteroidal anti-inflammatory drugs, angiotensin-converting enzyme (ACE) inhibitors, angiotensin II receptor blockers, spironolactone, and eplerenone. Trimethoprim is usually given in combination with sulfamethoxazole or dapsone for PCP and, on average, increases the plasma K⁺ concentration by about 1 meq/L; however, the hyperkalemia may be severe. Trimethoprim is structurally and chemically related to amiloride and triamterene and, in this way, may function as a potassium-sparing diuretic. This effect results in inhibition of the epithelial sodium channel (ENaC) in the principal cell of the collecting duct. By blocking the Na⁺ channel, K⁺ secretion is also inhibited; K⁺ secretion is dependent on the lumen-negative potential difference generated by Na⁺ entry through the ENaC (Fig. 64e-1).

Trimethoprim is associated with a non-AG acidosis that parallels development of hyperkalemia such that the co-occurrence of

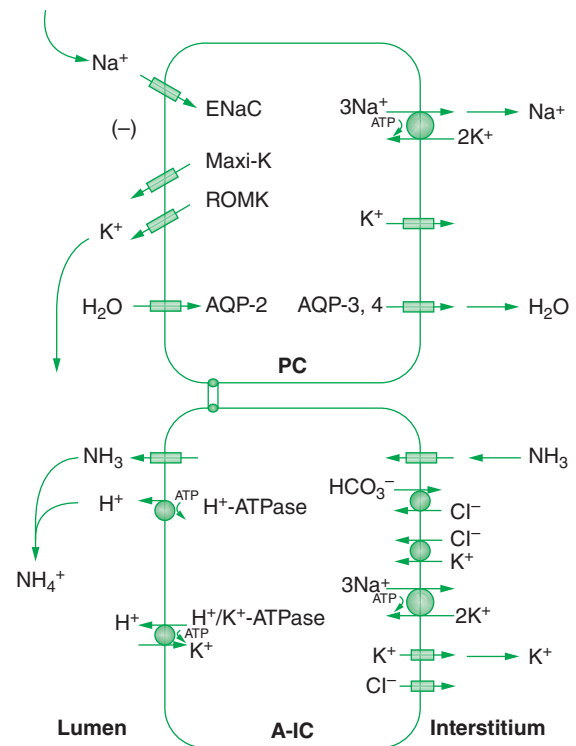


FIGURE 64e-1 Water, sodium, potassium, ammonia, and proton transport in principal cells (PC) and adjacent type A intercalated cells (A-IC). Water is absorbed down the osmotic gradient by principal cells, through the apical aquaporin-2 (AQP-2) and basolateral aquaporin-3 (AQP-3) and aquaporin-4 (AQP-4) channels. The absorption of Na⁺ via the amiloride-sensitive epithelial sodium channel (ENaC) generates a lumen-negative potential difference, which drives K⁺ excretion through the apical secretory K⁺ channel, ROMK (renal outer medullary K⁺ channel), and/or the flow-dependent maxi-K channel. Transepithelial ammonia (NH₃) transport and proton transport occur in adjacent type A intercalated cells, via apical and basolateral ammonia channels and apical H⁺-ATPase pumps, respectively; NH₄⁺ is ultimately excreted in the urine, in the defense of systemic pH. Electrogenic proton secretion by type A intercalated cells is also affected by the lumen-negative potential difference generated by the adjacent principal cells, such that reduction of this lumen-negative electrical gradient can reduce H⁺ excretion. Type A intercalated cells also reabsorb filtered K⁺ in potassium-deficient states, via apical H⁺/K⁺-ATPase.

hyperkalemia and metabolic acidosis is not uncommon in this setting. H⁺ secretion via apical H⁺-ATPase pumps in adjacent type A intercalated cells (Fig. 64e-1) is also electrogenic, such that the reduction in the lumen-negative potential difference due to trimethoprim inhibits distal H⁺ secretion; this is often referred to as a “voltage defect” form of dRTA. Systemic hyperkalemia also suppresses renal ammoniogenesis, ammonium excretion, and, thus, acid excretion; i.e., hyperkalemia per se has multiple effects on urinary acidification.

The inhibitory effect of trimethoprim on K⁺ and H⁺ secretion in the cortical collecting tubule follows a dose-response relationship, and therefore, the higher doses of this agent used in HIV/AIDS patients with PCP or in deep tissue infections with methicillin-resistant *Staphylococcus aureus* (MRSA) result in a higher prevalence of hyperkalemia and acidosis. Conventional doses of trimethoprim can also induce hyperkalemia and/or acidosis in predisposed patients, in particular the elderly, patients with renal insufficiency, and/or those with baseline hyporeninemic hypoaldosteronism.

One means by which to assess the role of the kidney in the development of hyperkalemia is to calculate, from a spot urine and coincident plasma sample, the transtubular potassium gradient (TTKG). The TTKG is calculated as $(P_{\text{osmol}} \times U_{\text{Potassium}}) / (P_{\text{Potassium}} \times U_{\text{osmol}})$. The expected values of the TTKG are <3 in the presence of hypokalemia (see also Case 7 and Case 8) and >7–8 in the presence of hyperkalemia.

In this case, the value for the TTKG of approximately 2 indicates that renal excretion of potassium is abnormally low for the prevailing hyperkalemia. Therefore, the inappropriately low TTKG indicates that the hyperkalemia is of renal tubular origin.

APPROACH TO MANAGEMENT

Knowledge of the factors controlling potassium secretion by the cortical collecting tubule principal cell can be helpful in understanding the basis for treatment of the hyperkalemia, especially if discontinuing the offending agent is not a reasonable clinical option. Potassium secretion is encouraged by a higher urine flow rate, increased distal delivery of sodium, distal delivery of a poorly reabsorbed anion (such as bicarbonate), and/or administration of a loop diuretic. Therefore, the approach to treatment in this patient should include intravenous 0.9% NaCl to expand the ECF and deliver more Na⁺ and Cl⁻ to the cortical collecting tubule. In addition, because the trimethoprim molecule must be protonated to inhibit ENaC, alkalinization of the renal tubule fluid enhances distal tubular K⁺ secretion. As an alternative to inducing bicarbonaturia to assist in potassium secretion, a carbonic anhydrase inhibitor may be administered to induce a kaliuresis. However, in the case presented here, for acetazolamide to be effective, the non-AG metabolic acidosis in this patient would first need to be corrected; Acetazolamide would, thus, require the coadministration of intravenous sodium bicarbonate for maximal benefit. Finally, systemic hyperkalemia directly suppresses renal ammoniogenesis, ammonium excretion, and, thus, acid excretion. Correcting the hyperkalemia with a potassium-binding resin (Kayexalate) is sometimes appropriate in these patients; the subsequent decline in the plasma K⁺ concentration will also increase urinary ammonium excretion, helping correct the acidosis.

CASE 3

A 63-year-old man was admitted to the intensive care unit (ICU) with a severe aspiration pneumonia. Past medical history included schizophrenia, for which he required institutional care; treatment had included neuroleptics and intermittent lithium, the latter restarted 6 months before admission. The patient was treated with antibiotics and intubated for several days, with the development of polyuria (3–5 L/d), hypernatremia, and acute renal insufficiency; the peak plasma Na⁺ concentration was 156 meq/L, and peak creatinine was 2.6 mg/dL. Urine osmolality was measured once and reported as 157 mOsm/kg, with a coincident plasma osmolality of 318 mOsm/kg. Lithium was stopped on admission to the ICU.

On physical examination, the patient was alert, extubated, and thirsty. Weight was 97.5 kg. Urine output for the previous 24 h had been 3.4 L, with an IV intake of 2 L/d of D₅W.

Laboratory Data					
Na 150	K 3.9	Cl 114	HCO ₃ ⁻ 26	BUN 8	Creat 1.7
Glu 95	Alb 3.1	Ca 8.1	Phos 2.6	Mg 2.0	Plasma Osm 315
Urine:	Na 34	K 5.2	Osm 137		

After 3 days of intravenous hydration, a water deprivation test was performed. A single dose of 2 µg IV desmopressin (DDAVP) was given at 9 h (+9):

Laboratory Data					
Time (h)	0	+6	+8	+12	+18
Na ⁺	145	148	150	152	149
K ⁺	5.4	5.3	3.9	3.9	3.9
Cl ⁻	111	110	118	120	114
HCO ₃ ⁻	24	27	25	242	25
Creat	1.3	1.3	1.4	1.3	1.3
S _{osmol}	300	311	315		
U _{osmol}	132	140	201	237	257
AVP		8.4	6.3		

APPROACH TO DIAGNOSIS

Why did the patient develop hypernatremia, polyuria, and acute renal insufficiency? What does the water deprivation test demonstrate? What is the underlying pathophysiology of this patient's hypernatremic syndrome?

This patient became polyuric after admission to the ICU with severe pneumonia, developing significant hypernatremia and acute renal insufficiency. Polyuria can result from either an osmotic diuresis or a water diuresis. An osmotic diuresis can be caused by excessive excretion of Na⁺-Cl⁻, mannitol, glucose, and/or urea, with a daily solute excretion of >750–1000 mOsm/d (>15 mOsm/kg body water per day). In this case, however, the patient was excreting large volumes of very hypotonic urine, with a urine osmolality that was substantially lower than that of plasma; this, by definition, was a water diuresis, resulting in inappropriate excretion of free water and hypernatremia. The appropriate response to hypernatremia and a plasma osmolality >295 mOsm/kg is an increase in circulating vasopressin (AVP) and the excretion of low volumes (<500 mL/d) of maximally concentrated urine, i.e., urine with osmolality >800 mOsm/kg. This patient's response to hypernatremia was clearly inappropriate, due to either a loss of circulating AVP (central diabetes insipidus [CDI]) or renal resistance to AVP (nephrogenic diabetes insipidus [NDI]). Ongoing loss of free water was sufficiently severe in this patient that absolute hypovolemia ensued, despite the fact that approximately two-thirds of the excreted water was derived from the intracellular fluid compartment rather than the ECF compartment. Hypovolemia led to an acute decrease in the glomerular filtration rate (GFR), i.e., acute renal insufficiency, with gradual improvement following hydration (see below).

Following the correction of hypernatremia and acute renal insufficiency with appropriate hydration (see below), the patient was subjected to a water deprivation test followed by administration of DDAVP. This test helps determine whether an inappropriate water diuresis is caused by CDI or NDI. The patient was water restricted beginning in the early morning, with careful monitoring of vital signs and urine output; overnight water deprivation of patients with diabetes insipidus is unsafe and clinically inappropriate, given the potential for severe hypernatremia. The plasma Na⁺ concentration, which is more accurate and more immediately available than plasma osmolality, was monitored hourly during water deprivation. A baseline AVP sample was drawn at the beginning of the test, with a second sample drawn once the plasma Na⁺ reached 148–150 meq/L. At this point, a single 2-µg dose of the V₂ AVP receptor agonist DDAVP was administered. An alternative approach would have been to measure AVP and administer DDAVP when the patient was initially hypernatremic; however, it would have been less safe to administer DDAVP in the setting of renal impairment because clearance of DDAVP is renal dependent.

The patient's water deprivation test was consistent with NDI, with an AVP level within the normal range in the setting of hypernatremia (i.e., no evidence of CDI) and an inappropriately low urine osmolality that failed to increase by >50% or >150 mOsm/kg after both water deprivation and the administration of DDAVP. This defect would be considered compatible with complete NDI; patients with partial NDI can achieve urine osmolalities of 500–600 mOsm/kg after DDAVP treatment but will not maximally concentrate their urine to 800 mOsm/kg or higher.

NDI has a number of genetic and acquired causes, which all share interference with some aspect of the renal concentrating mechanism. For example, loss-of-function mutations in the V₂ AVP receptor cause X-linked NDI. This patient suffered from NDI due to lithium therapy, perhaps the most common cause of NDI in adult medicine. Lithium causes NDI via direct inhibition of renal glycogen synthase kinase-3 (GSK3), a kinase thought to be the pharmacologic target of lithium in psychiatric disease; renal GSK3 is required for the response of principal cells to AVP. Lithium also induces the expression of cyclooxygenase-2 (COX2) in the renal medulla; COX2-derived prostaglandins inhibit AVP-stimulated salt transport by the thick ascending limb and AVP-stimulated water transport by the collecting duct, thereby

exacerbating lithium-associated polyuria. The entry of lithium through the amiloride-sensitive Na⁺ channel ENaC (Fig. 64e-1) is required for the effect of the drug on principal cells, such that combined therapy with lithium and amiloride can mitigate lithium-associated NDI. However, lithium causes chronic tubulointerstitial scarring and chronic kidney disease after prolonged therapy, such that patients may have a persistent NDI long after stopping the drug, with a reduced therapeutic benefit from amiloride. Notably, this particular patient had been treated intermittently for several years with lithium, with the development of chronic kidney disease (baseline creatinine of 1.3–1.4) and NDI that persisted after stopping the drug.

APPROACH TO MANAGEMENT

How should this patient be treated? What are the major pitfalls of therapy?

This patient developed severe hypernatremia due to a water diuresis from lithium-associated NDI. Treatment of hypernatremia must include both replacement of the existing free water deficit and daily replacement of ongoing free water loss. The first step is to estimate total-body water (TBW), typically estimated as 50% of the body weight in women and 60% in men. The free water deficit is then calculated as $([Na^+ - 140]/140) \times TBW$. In this patient, the free water deficit was 4.2 L at a weight of 97.5 kg and plasma Na⁺ concentration of 150 meq/L. This free water deficit should be replaced slowly over 48–72 h to avoid increasing the plasma Na⁺ concentration by >10 meq/L per 24 h. A common mistake is to replace this deficit while neglecting to replace ongoing losses of free water, such that plasma Na⁺ concentration either fails to correct or, in fact, increases.

Ongoing losses of free water can be estimated using the equation for electrolyte-free water clearance:

$$C_{eH_2O} = V (1 - [U_{Na} + U_K]/P_{Na})$$

where V is urinary volume, U_{Na} is urinary [Na⁺], U_K is urinary [K⁺], and P_{Na} is plasma [Na⁺].

For this patient, the C_{eH₂O} was 2.5 L/d when initially evaluated, i.e., with urine Na⁺ and K⁺ concentrations of 34 and 5.2 meq/L, plasma Na⁺ concentration of 150 meq/L, and a urinary volume of 3.4 L. Therefore, the patient was given 2.5 L of D₅W over the first 24 h to replace ongoing free water losses, along with 2.1 L of D₅W to replace half his free water deficit. Daily random urine electrolytes and urinary volume measurement can be used to monitor C_{eH₂O} and adjust daily fluid administration in this manner, while following plasma Na⁺ concentration. Physicians often calculate the free water deficit to guide therapy of hypernatremia, providing half the deficit in the first 24 h. This approach can be adequate in patients who do not have significant ongoing losses of free water, e.g., with hypernatremia due to decreased free water intake. This case illustrates how free water requirements can be grossly underestimated in hypernatremic patients if ongoing, daily free water losses are not taken into account.

CASE 4

A 78-year-old man was admitted with pneumonia and hyponatremia. Plasma Na⁺ concentration was initially 129 meq/L, decreasing within 3 days to 118–120 meq/L despite fluid restriction to 1 L/d. A chest computed tomography (CT) revealed a right 2.8 × 1.6 cm infrahilar mass and postobstructive pneumonia. The patient was an active smoker. Past medical history was notable for laryngeal carcinoma treated 15 years prior with radiation therapy, renal cell carcinoma, peripheral vascular disease, and hypothyroidism. On review of systems, he denied headache, nausea, and vomiting. He had chronic hip pain, managed with acetaminophen with codeine. Other medications included cilostazol, amoxicillin/clavulanate, digoxin, diltiazem, and thyroxine. He was euvolemic on examination, with no lymphadenopathy and a normal chest examination.

Laboratory Data

Na ⁺ 120	K ⁺ 4.3	Cl ⁻ 89	HCO ₃ ⁻ 23	BUN 8	Creat 1.0	Glu 93
Alb 3.1	Ca 8.9	Phos 2.8	Mg 2.0	Plasma osm 248	mOsm/kg	
Cortisol 25 μg/dL	TSH 2.6	Uric acid 2.7	mg/dL			
Urine:	Na ⁺ 97	K ⁺ 22	Cl ⁻ 86	Osm 597		

The patient was treated with furosemide, 20 mg PO bid, and salt tablets. The plasma Na⁺ concentration increased to 129 meq/L with this therapy; however, the patient developed orthostatic hypotension and dizziness. He was started on demeclocycline, 600 mg PO in the morning and 300 mg in the evening, just before discharge from hospital. Plasma Na⁺ concentration increased to 140 meq/L with a BUN of 23 and creatinine of 1.4, at which point demeclocycline was reduced to 300 mg PO bid. Bronchoscopic biopsy eventually showed small-cell lung cancer; the patient declined chemotherapy and was admitted to hospice.

APPROACH TO DIAGNOSIS AND MANAGEMENT

What factors contributed to this patient's hyponatremia? What are the therapeutic options?

This patient developed hyponatremia in the context of a central lung mass and postobstructive pneumonia. He was clinically euvolemic, with a generous urine Na⁺ concentration and low plasma uric acid concentration. He was euthyroid, with no evidence of pituitary dysfunction or secondary adrenal insufficiency. The clinical presentation is consistent with the syndrome of inappropriate antidiuresis (SIAD). Although pneumonia was a potential contributor to the SIAD, it was notable that the plasma Na⁺ concentration decreased despite a clinical response to antibiotics. It was suspected that this patient had SIAD due to small-cell lung cancer, with a central lung mass on chest CT and a significant smoking history. There was a history of laryngeal cancer and renal cancer but with no evidence of recurrent disease; these malignancies were not considered contributory to his SIAD. Biopsy of the lung mass ultimately confirmed the diagnosis of small-cell lung cancer, which is responsible for ~75% of malignancy-associated SIAD; ~10% of patients with this neuroendocrine tumor will have a plasma Na⁺ concentration of <130 meq/L at presentation. The patient had no other “nonosmotic” stimuli for an increase in AVP, with no medications associated with SIAD and minimal pain or nausea.

The patient had no symptoms attributable to hyponatremia but was judged at risk for worsening hyponatremia from severe SIAD. Persistent, chronic hyponatremia (duration >48 h) results in an efflux of organic osmolytes (creatine, betaine, glutamate, myoinositol, and taurine) from brain cells; this response reduces intracellular osmolality and the osmotic gradient favoring water entry. This cellular response does not fully protect patients from symptoms, which can include vomiting, nausea, confusion, and seizures, usually at plasma Na⁺ concentration <125 meq/L. Even patients who are judged “asymptomatic” can manifest subtle gait and cognitive defects that reverse with correction of hyponatremia. Chronic hyponatremia also increases the risk of bony fractures due to an increased risk of falls and to a hyponatremia-associated reduction in bone density. Therefore, every attempt should be made to correct plasma Na⁺ concentration safely in patients with chronic hyponatremia. This is particularly true in malignancy-associated SIAD, where it can take weeks to months for a tissue diagnosis and the subsequent reduction in AVP following initiation of chemotherapy, radiotherapy, and/or surgery.

What are the therapeutic options in SIAD? Water deprivation, a cornerstone of therapy for SIAD, had little effect on the plasma Na⁺ concentration in this patient. The urine:plasma electrolyte ratio (urinary [Na⁺] + [K⁺]/plasma [Na⁺]) can be used to estimate electrolyte-free water excretion and the required degree of water restriction; patients with a ratio of >1 should be more aggressively restricted (<500 mL/d), those with a ratio of ~1 should be restricted to 500–700 mL/d, and those with a ratio <1 should be restricted to <1 L/d. This patient had a urine:plasma electrolyte ratio of 1 and predictably did not respond to a moderate water restriction of ~1 L/d. A more

aggressive water restriction would have theoretically been successful; however, this can be very difficult for patients with SIAD to tolerate, given that their thirst is also inappropriately stimulated.

Combined therapy with furosemide and salt tablets can often increase the plasma Na⁺ concentration in SIAD; furosemide reduces maximal urinary concentrating ability by inhibiting the countercurrent mechanism, whereas the salt tablets mitigate diuretic-associated NaCl loss and amplify the ability to excrete free water by increasing urinary solute excretion. This regimen is not always successful and requires careful titration of salt tablets to avoid volume depletion; indeed, in this particular patient, the plasma Na⁺ concentration remained <130 meq/L and the patient became orthostatic. The principal cell toxin, demeclocycline, is an alternative oral agent in SIAD. Treatment with demeclocycline was very successful in this patient, with an increase in plasma Na⁺ concentration to 140 meq/L. However, demeclocycline can be natriuretic, leading to a prerenal decrease in GFR. Demeclocycline has also been implicated in nephrotoxic injury, particularly in patients with cirrhosis and chronic liver disease, in whom the drug accumulates. Notably, this particular patient developed a significant but stable decrease in GFR while on demeclocycline, necessitating a reduction in the administered dose.

A major advance in the management of hyponatremia was the clinical development of AVP antagonists (vaptans). These agents inhibit the effect of AVP on renal V₂ receptors, resulting in the excretion of electrolyte-free water and correction of hyponatremia. The specific indications for these agents are not as yet clear, despite U.S. Food and Drug Administration (FDA) approval for the management of both euvolemic and hypervolemic hyponatremia. It is, however, anticipated that the vaptans will have an increasing role in the management of SIAD and other causes of hyponatremia. Indeed, if this particular patient had continued with active therapy for his cancer, substitution of demeclocycline with oral tolvaptan (a V₂-specific oral vaptan) would have been the next appropriate step, given the development of renal insufficiency with demeclocycline. As with other measures to correct hyponatremia (e.g., hypertonic saline, demeclocycline), the vaptans have the potential to “overcorrect” plasma Na⁺ concentration (a rise of >8–10 meq/L per 24 h or 18 meq/L per 18 h), thus increasing the risk for osmotic demyelination (see Case 5). Therefore, the plasma Na⁺ concentration should be monitored closely during the initiation of therapy with these agents. In addition, long-term use of tolvaptan has been associated with abnormalities in liver function tests; hence, use of this agent should be restricted to only 1–2 months.

CASE 5

A 76-year-old woman presented with a several-month history of diarrhea, with marked worsening over the 2–3 weeks before admission (up to 12 stools a day). Review of systems was negative for fever, orthostatic dizziness, nausea and vomiting, or headache. Past medical history included hypertension, kidney stones, and hypercholesterolemia; medications included atenolol, spironolactone, and lovastatin. She also reliably consumed >2 L of liquid per day in management of the nephrolithiasis.

The patient received 1 L of saline over the first 5 h of her hospital admission. On examination at hour 6, the heart rate was 72 sitting and 90 standing, and blood pressure was 105/50 mmHg lying and standing. Her jugular venous pressure (JVP) was indistinct with no peripheral edema. On abdominal examination, the patient had a slight increase in bowel sounds but a nontender abdomen and no organomegaly.

The plasma Na⁺ concentration on admission was 113 meq/L, with a creatinine of 2.35 (Table 64e-1). At hospital hour 7, the plasma

Na⁺ concentration was 120 meq/L, potassium 5.4 meq/L, chloride 90 meq/L, bicarbonate 22 meq/L, BUN 32 mg/dL, creatinine 2.02 mg/dL, glucose 89 mg/dL, total protein 5.0, and albumin 1.9. The hematocrit was 33.9, white count 7.6, and platelets 405. A morning cortisol was 19.5, with thyroid-stimulating hormone (TSH) of 1.7. The patient was treated with 1 μg of intravenous DDAVP, along with 75 mL/h of intravenous half-normal saline. After the plasma Na⁺ concentration dropped to 116 meq/L, intravenous fluid was switched to normal saline at the same infusion rate. The subsequent results are shown in Table 64e-1.

APPROACH TO DIAGNOSIS

This patient presented with hypovolemic hyponatremia and a “prerenal” reduction in GFR, with an increase in serum creatinine. She had experienced diarrhea for some time and manifested an orthostatic tachycardia after a liter of normal saline. As expected for hypovolemic hyponatremia, the urine Na⁺ concentration was <20 meq/L in the absence of congestive heart failure or other causes of hypervolemic hyponatremia, and she responded to saline hydration with an increase in plasma Na⁺ concentration and a decrease in creatinine.

The initial hypovolemia increased the sensitivity of this patient’s AVP response to osmolality, both decreasing the osmotic threshold for AVP release and increasing the slope of the osmolality response curve. AVP has a half-life of only 10–20 min; therefore, the acute increase in intravascular volume after a liter of intravenous saline led to a rapid reduction in circulating AVP. The ensuing water diuresis is the primary explanation for the rapid increase in plasma Na⁺ concentration in the first 7 h of her hospitalization.

APPROACH TO MANAGEMENT

The key concern in this case was the evident chronicity of the patient’s hyponatremia, with several weeks of diarrhea followed by 2–3 days of acute exacerbation. This patient was judged to have chronic hyponatremia, i.e., with a suspected duration of >48 h; as such, she would be predisposed to osmotic demyelination were she to undergo too rapid a correction in her plasma Na⁺ concentration, i.e., by >8–10 meq/L in 24 h or 18 meq/L in 48 h. At presentation, she had no symptoms that one would typically attribute to acute hyponatremia, and the plasma Na⁺ concentration had already increased by a sufficient amount to protect from cerebral edema; however, she had corrected by 1 meq/L per hour within the first 7 h of admission, consistent with impending overcorrection. To reduce or halt the increase in plasma Na⁺ concentration, the patient received 1 μg of intravenous DDAVP along with intravenous free water. Given the hypovolemia and resolving acute renal insufficiency, a decision was made to administer half-normal saline as a source of free water, rather than D₅W; this was switched to normal saline when plasma Na⁺ concentration acutely dropped to 117 meq/L (Table 64e-1).

Overcorrection of chronic hyponatremia is a major risk factor for the development of osmotic demyelination syndrome (ODS). Animal studies show a neurologic and survival benefit in ODS of “re-lowering” plasma Na⁺ concentration with DDAVP and free water administration; this approach is demonstrably safe in patients with hyponatremia, with no evident risk of seizure or other sequelae. This combination can be used to prevent an overcorrection or to re-lower plasma Na⁺ concentration in patients who have already overcorrected. DDAVP is required because in most of these patients endogenous AVP levels have plummeted, resulting in a free water diuresis; the administration of free water alone has minimal effect in this setting, given the relative absence of circulating AVP. An alternative approach in patients who present with severe hyponatremia is to treat them

prospectively with twice-daily DDAVP to prevent changes in AVP bioactivity, coadministering hypertonic saline to increase slowly the plasma Na⁺ concentration in a more controlled fashion.

This patient’s plasma Na⁺ concentration remained depressed for several days after DDAVP administration. It is

TABLE 64e-1 SERIAL LABORATORY DATA FOR CASE 5

Hospital Hour	Baseline	0	3	7	11	14	24	48	72
Plasma Na ⁺ (meq/L)	137	113	115	120	117	116	117	124	130
Creatinine (mg/dL)	1.2	2.35	2.10	2.02	1.97	1.79	1.53	1.20	1.13
Urine osmolality (mOsm/kg)				319		415	397		
Urine Na ⁺ (meq/L)				17		23	47		

conceivable that residual hypovolemic hyponatremia attenuated the recovery of the plasma Na^+ concentration. Alternatively, attenuated recovery was due to persistent effects of the single dose of DDAVP. Of note, although the plasma half-life of DDAVP is only 1–2 h, pharmacodynamic studies indicate a much more prolonged effect on urine output and/or urine osmolality. One final consideration is the effect of the patient's initial renal dysfunction on the pharmacokinetics and pharmacodynamics of the administered DDAVP, which is renally excreted; DDAVP should be administered with caution for the reinduction of hyponatremia in patients with chronic kidney disease or acute renal dysfunction.

CASE 6

A 44-year-old woman was referred from a local hospital after presenting with flaccid paralysis. Severe hypokalemia was documented (2.0 meq/L), and an infusion containing KCl was initiated.

Laboratory Data	Value	Units
Sodium	140	meq/L
Potassium	2.6	meq/L
Chloride	115	meq/L
Bicarbonate	15	meq/L
Anion gap	10	meq/L
BUN	22	mg/dL
Creatinine	1.4	mg/dL
Arterial Blood Gases		
pH	7.32	U
Paco_2	30	mmHg
HCO_3^-	15	meq/L
Additional Laboratory Data		
Rheumatoid factor positive, anti-Ro/SS-A positive, and anti-La/SS-B positive		
Urinalysis		
pH = 6.0, normal sediment without white or red blood cell casts and no bacteria. The urine protein-to-creatinine ratio was 0.150 g/g. Urinary electrolyte values were: Na^+ 35, K^+ 40, Cl^- 18 meq/L. Therefore, the urine anion gap was positive, indicating low urine NH_4^+ excretion.		

APPROACH TO DIAGNOSIS

The diagnosis in this case is classic hypokalemic dRTA from Sjögren's syndrome. This patient presented with a non-AG metabolic acidosis. The urine AG was positive, indicating an abnormally low excretion of ammonium in the face of systemic acidosis. The urine pH was inappropriately alkaline, yet there was no evidence of hypercalciuria, nephrocalcinosis, or bone disease. The patient was subsequently shown to exhibit hyperglobulinemia. These findings, taken together, indicate that the cause of this patient's hypokalemia and non-AG metabolic acidosis was a renal tubular abnormality. The hypokalemia and abnormally low excretion of ammonium, as estimated by the urine AG, in the absence of glycosuria, phosphaturia, or aminoaciduria (Fanconi's syndrome), defines the entity of classic distal renal tubular acidosis (dRTA), also known as type 1 RTA. Because of the hyperglobulinemia, additional serology was obtained, providing evidence for the diagnosis of primary Sjögren's syndrome. Furthermore, additional history indicated a 5-year history of xerostomia and keratoconjunctivitis sicca but without synovitis, arthritis, or rash.

Classic dRTA occurs frequently in patients with Sjögren's syndrome and is a result of an immunologic attack on the collecting tubule, causing failure of the H^+ -ATPase to be inserted into the apical membrane of type A intercalated cells. Sjögren's syndrome is one of the best-documented acquired causes of classic dRTA. The loss of H^+ -ATPase function also occurs with certain inherited forms of classic dRTA. There was no family history in the present case, and other family members were not affected. A number of autoantibodies have been associated with Sjögren's syndrome; it is likely that these autoantibodies prevent trafficking or function of the H^+ -ATPase in the type A intercalated cell of the collecting tubule. Although proximal RTA

has also been reported in patients with Sjögren's syndrome, it is much less frequent, and there were no features of proximal tubule dysfunction (Fanconi's syndrome) in this patient. The hypokalemia is due to secondary hyperaldosteronism from volume depletion.

APPROACH TO MANAGEMENT

The long-term renal prognosis for patients with classic dRTA due to Sjögren's syndrome has not been established. Nevertheless, the metabolic acidosis and the hypokalemia respond to alkali replacement with either sodium citrate solution (Shohl's solution) or sodium bicarbonate tablets. Obviously, potassium deficits must be replaced initially, but potassium replacement is usually not required in dRTA patients long term because sodium bicarbonate (or citrate) therapy expands volume and corrects the secondary hyperaldosteronism. A consequence of the interstitial infiltrate seen in patients with Sjögren's syndrome and classic dRTA is progression of chronic kidney disease. Cytotoxic therapy plus glucocorticoids has been the mainstay of therapy in Sjögren's syndrome for many years, although B lymphocyte infiltration in salivary gland tissue subsides and urinary acidification improves after treatment with rituximab.

CASE 7

A 32-year-old man was admitted to the hospital with weakness and hypokalemia. The patient had been very healthy until 2 months previously when he developed intermittent leg weakness. His review of systems was otherwise negative. He denied drug or laxative abuse and was on no medications. Past medical history was unremarkable, with no history of neuromuscular disease. Family history was notable for a sister with thyroid disease. Physical examination was notable only for reduced deep tendon reflexes.

Laboratory Data	Admission	Baseline	Units
Sodium	139	143	meq/L
Potassium	2.0	3.8	meq/L
Chloride	105	107	meq/L
Bicarbonate	26	29	meq/L
BUN	11	16	mg/dL
Creatinine	0.6	1.0	mg/dL
Calcium	8.8	8.8	mg/dL
Phosphate	1.2		mg/dL
Albumin	3.8		mg/dL
Plasma osmolality	290		mOsm/kg
Urine osmolality	590		mOsm/kg
Urine potassium	10		meq/L
TSH 0.08 $\mu\text{IU/L}$ (normal 0.2–5.39)			
Free T_4 41 pmol/L (normal 10–27)			

APPROACH TO DIAGNOSIS

This patient developed hypokalemia due to a redistribution of potassium between the intracellular and extracellular compartments; this pathophysiology was readily apparent following calculation of the TTKG. The TTKG is calculated as $(\text{P}_{\text{osmol}} \times \text{U}_{\text{Potassium}}) / (\text{P}_{\text{Potassium}} \times \text{U}_{\text{osmol}})$. The expected values for the TTKG are <3 in the presence of hypokalemia and >7 – 8 in the presence of hyperkalemia (see also Case 2 and Case 8). Alternatively, a urinary K^+ -to-creatinine ratio of >13 mmol/g creatinine (>1.5 mmol/mmol creatinine) is compatible with excessive renal K^+ excretion. In this case, the calculated TTKG was 2.5, consistent with appropriate renal conservation of K^+ and a nonrenal cause for hypokalemia. In the absence of significant gastrointestinal loss of K^+ , the patient was diagnosed with a "redistributive" subtype of hypokalemia.

More than 98% of total-body potassium is intracellular; regulated buffering of extracellular K^+ by this large intracellular pool plays a crucial role in the maintenance of a stable plasma K^+ concentration. Clinically, changes in the exchange and distribution of intra- and

extracellular K^+ can cause significant hypo- or hyperkalemia. Insulin, β_2 -adrenergic activity, thyroid hormone, and alkalosis promote cellular uptake of K^+ by multiple interrelated mechanisms, leading to hypokalemia. In particular, alterations in the activity of the endogenous sympathetic nervous system can cause hypokalemia in several settings, including alcohol withdrawal, hyperthyroidism, acute myocardial infarction, and severe head injury.

Weakness is common in severe hypokalemia; hypokalemia causes hyperpolarization of muscle, thereby impairing the capacity to depolarize and contract. In this particular patient, Graves' disease caused hyperthyroidism and hypokalemic paralysis (thyrotoxic periodic paralysis [TPP]). TPP develops more frequently in patients of Asian or Hispanic origin. This predisposition has been linked to genetic variation in Kir2.6, a muscle-specific, thyroid hormone-induced K^+ channel; however, the pathophysiologic mechanisms that link dysfunction of this ion channel to TPP have yet to be elucidated. The hypokalemia in TPP is attributed to both direct and indirect activation of the Na^+/K^+ -ATPase by thyroid hormone, resulting in increased uptake of K^+ by muscle and other tissues. Thyroid hormone induces expression of multiple subunits of the Na^+/K^+ -ATPase in skeletal muscle, increasing the capacity for uptake of K^+ ; hyperthyroid increases in β -adrenergic activity are also thought to play an important role in TPP.

Clinically, patients with TPP present with weakness of the extremities and limb girdle, with paralytic episodes that occur most frequently between 1 and 6 AM. Precipitants of weakness include high carbohydrate loads and strenuous exercise. Signs and symptoms of hyperthyroidism are not always present, often leading to delays in diagnosis. Hypokalemia is often profound and usually accompanied by redistributive hypophosphatemia, as in this case. A TTKG of $<2-3$ separates patients with TPP from those with hypokalemia due to renal potassium wasting, who will have TTKG values that are >4 . This distinction is of considerable importance for therapy; patients with large potassium deficits require aggressive repletion with K^+-Cl^- , which has a significant risk of rebound hyperkalemia in TPP and related disorders.

APPROACH TO MANAGEMENT

Ultimately, definitive therapy for TPP requires treatment of the associated hyperthyroidism. In the short term, however, potassium replacement is necessary to hasten muscle recovery and prevent cardiac arrhythmias. The average recovery time of an acute attack is reduced by $\sim 50\%$ in patients treated with intravenous K^+-Cl^- at a rate of 10 meq/h; however, this incurs a significant risk of rebound hyperkalemia, with up to 70% developing a plasma K^+ concentration of >5.0 meq/L. This potential for rebound hyperkalemia is a general problem in the management of all causes of redistributive hypokalemia, resulting in the need to distinguish these patients accurately and rapidly from those with a large K^+ deficit due to renal or extrarenal loss of K^+ . An attractive alternative to K^+-Cl^- replacement in TPP is treatment with high-dose propranolol (3 mg/kg), which rapidly reverses the associated hypokalemia, hypophosphatemia, and paralysis. Notably, rebound hyperkalemia is not associated with this treatment.

CASE 8

A 66-year-old man was admitted to hospital with a plasma K^+ concentration of 1.7 meq/L and profound weakness. The patient had noted progressive weakness over several days, to the point that he was unable to rise from bed. Past medical history was notable for small-cell lung cancer with metastases to brain, liver, and adrenals. The patient had been treated with one cycle of cisplatin/etoposide 1 year before this admission, which was complicated by acute kidney injury (peak creatinine of 5, with residual chronic kidney disease), and three subsequent cycles of cyclophosphamide/doxorubicin/vincristine, in addition to 15 treatments with whole-brain radiation.

On physical examination, the patient was jaundiced. Blood pressure was 130/70 mmHg, increasing to 160/98 mmHg after 1 L of saline, with a JVP at 8 cm. There was generalized muscle weakness.

Laboratory Data	2 Months			Units
	PTA	Admission	HD2	
Sodium	143	149	144	meq/L
Potassium	3.7	1.7	3.5	meq/L
Chloride	103	84	96	meq/L
Bicarbonate	26	44	34	meq/L
Venous pH		7.47		pH
Venous P_{CO_2}		62		mmHg
BUN	21	41	40	mg/dL
Creatinine	2.8	2.9	2.3	mg/dL
Magnesium	1.3	1.6	2.4	mg/dL
CPK		183		U/L
ALT	8	75		U/L
Albumin	3.4	2.8	2.3	
Adjusted anion gap	15	24	18	
Total bilirubin	0.65	5.19		mg/dL
Alkaline phosphatase	93	217		U/L
Urine sodium		35	28	meq/L
Urine potassium		25	49	meq/L
Urine chloride		48	51	meq/L
Urine osmolality		391		mOsm/kg
Plasma osmolality		312		mOsm/kg
Urine pH		5.5		
Plasma ACTH		185		pg/mL (7–50 pg/mL)
Plasma cortisol		94		pg/mL (3–16 pg/mL)
24-h urine cortisol		1044		μ g/24 h (4–50 μ g/24 h)

Abbreviations: ACTH, adrenocorticotropic hormone; HD2, hospital day 2; PTA, prior to admission.

The patient's hospital course was complicated by acute respiratory failure attributed to pulmonary embolism; he died 2 weeks after admission.

APPROACH TO DIAGNOSIS

Why was this patient hypokalemic? Why was he weak? Why did he have an alkalosis?

This patient suffered from metastatic small-cell lung cancer, which was persistent despite several rounds of chemotherapy and radiotherapy. He presented with profound hypokalemia, alkalosis, hypertension, severe weakness, jaundice, and worsening liver function tests.

With respect to the hypokalemia, there was no evident cause of nonrenal potassium loss, e.g., diarrhea. The urinary TTKG was 11.7, at a plasma K^+ concentration of 1.7 meq/L; this TTKG value is consistent with inappropriate renal K^+ secretion, despite severe hypokalemia. The TTKG is calculated as $(P_{osmol} \times U_{Potassium}) / (P_{Potassium} \times U_{osmol})$. The expected values for the TTKG are <3 in the presence of hypokalemia and $>7-8$ in the presence of hyperkalemia (see also Case 2 and Case 6).

The patient had several explanations for excessive renal loss of potassium. First, he had had a history of cisplatin-associated acute kidney injury, with residual chronic kidney disease. Cisplatin can cause persistent renal tubular defects, with prominent hypokalemia and hypomagnesemia; however, this patient had not previously required potassium or magnesium repletion, suggesting that cisplatin-associated renal tubular defects did not play a major role in this presentation with severe hypokalemia. Second, he was hypomagnesemic on presentation, suggesting total-body magnesium depletion. Magnesium depletion has inhibitory effects on muscle Na^+/K^+ -ATPase activity, reducing influx into muscle cells and causing a secondary increase in K^+ excretion. Magnesium depletion also increases K^+ secretion by the distal nephron; this is attributed to a reduction in the magnesium-dependent, intracellular block of K^+ efflux through the secretory K^+ channel of principal cells (ROMK, Fig. 64e-1). Clinically, hypomagnesemic patients are

refractory to K^+ replacement in the absence of Mg^{2+} repletion. Again, however, this patient had not previously developed significant hypokalemia, despite periodic hypomagnesemia, such that other factors must have caused the severe hypokalemia.

The associated hypertension in this case suggested an increase in mineralocorticoid activity, causing increased activity of ENaC channels in principal cells, $NaCl$ retention, hypertension, and hypokalemia. The increase in ENaC-mediated Na^+ transport in principal cells would have led to an increase in the lumen-negative potential difference in the connecting tubule and cortical collecting duct, driving an increase in K^+ secretion through apical K^+ channels (Fig. 64e-1). This explanation is compatible with the very high TTKG, i.e., an increase in K^+ excretion that is inappropriate for the plasma K^+ concentration.

What caused an increase in mineralocorticoid activity in this patient? The patient had bilateral adrenal metastases, indicating that primary hyperaldosteronism was unlikely. The clinical presentation (hypokalemia, hypertension, and alkalosis) and the history of small-cell lung cancer suggested Cushing's syndrome, with a massive increase in circulating glucocorticoids, in response to ectopic adrenocorticotropic hormone (ACTH) secretion by his small-cell lung cancer tumor. Confirmation of this diagnosis was provided by a very high plasma cortisol level, high ACTH level, and increased urinary cortisol (see the laboratory data above).

Why would an increase in circulating cortisol cause an apparent increase in mineralocorticoid activity? Cortisol and aldosterone have equal affinity for the mineralocorticoid receptor (MLR); thus, cortisol has mineralocorticoid-like activity; however, cells in the aldosterone-sensitive distal nephron (the distal convoluted tubule [DCT]), connecting tubule (CNT), and collecting duct are protected from circulating cortisol by the enzyme 11β -hydroxysteroid dehydrogenase-2 (11β HSD-2), which converts cortisol to cortisone (Fig. 64e-2); cortisone has minimal affinity for the MLR. Activation of the MLR causes activation of the basolateral Na^+/K^+ -ATPase, activation of the thiazide-sensitive Na^+ - Cl^- cotransporter in the DCT, and activation of apical ENaC

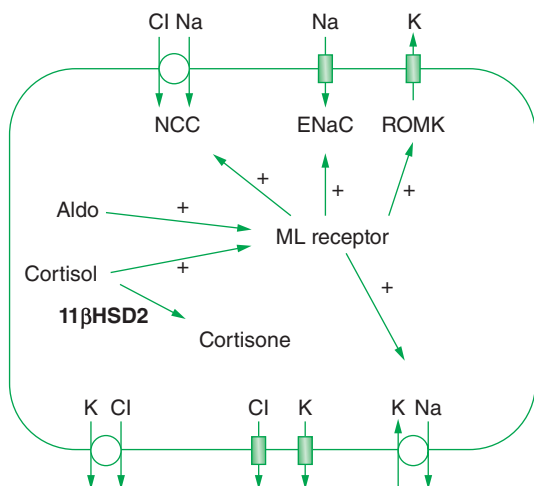


FIGURE 64e-2 11β -Hydroxysteroid dehydrogenase-2 (11β HSD-2) and syndromes of apparent mineralocorticoid excess. The enzyme 11β HSD-2 protects cells in the aldosterone-sensitive distal nephron (the distal convoluted tubule [DCT], connecting tubule [CNT], and collecting duct) from the illicit activation of mineralocorticoid receptors (MLR) by cortisol. Binding of aldosterone to the MLR leads to activation of the thiazide-sensitive Na^+ - Cl^- cotransporter in DCT cells and the amiloride-sensitive epithelial sodium channel (ENaC) in principal cells (CNT and collecting duct). Aldosterone also activates basolateral Na^+/K^+ -ATPase and, to a lesser extent, the apical secretory K^+ channel ROMK (renal outer medullary K^+ channel). Cortisol has equivalent affinity for the MLR to that of aldosterone; metabolism of cortisol to cortisone, which has no affinity for the MLR, prevents these cells from activation by circulating cortisol. Genetic deficiency of 11β HSD-2 or inhibition of its activity causes the syndromes of apparent mineralocorticoid excess (see Case 8).

channels in principal cells of the CNT and collecting duct (Fig. 64e-2). Recessive loss-of-function mutations in the 11β HSD-2 gene lead to cortisol-dependent activation of the MLR and the syndrome of apparent mineralocorticoid excess (SAME), comprising hypertension, hypokalemia, hypercalcemia, and metabolic alkalosis, with suppressed plasma renin activity (PRA) and suppressed aldosterone. A similar syndrome is caused by biochemical inhibition of 11β HSD-2 by glycyrrheticin/glycyrrhizinic acid (found in licorice, for example) and/or carbenoxolone.

In Cushing's syndrome caused by increases in pituitary ACTH, the incidence of hypokalemia is only 10%, whereas it is ~70% in patients with ectopic secretion of ACTH, despite a similar incidence of hypertension. The activity of renal 11β HSD-2 is reduced in patients with ectopic ACTH compared with Cushing's syndrome, resulting in SAME; the prevailing theory is that the much greater cortisol production in ectopic ACTH syndromes overwhelms the renal 11β HSD-2 enzyme, resulting in activation of renal MLRs by unmetabolized cortisol (Fig. 64e-2).

Why was the patient so weak? The patient was profoundly weak due to the combined effect of hypokalemia and increased cortisol. Hypokalemia causes hyperpolarization of muscle, thereby impairing the capacity to depolarize and contract. Weakness and even ascending paralysis can frequently complicate severe hypokalemia. Hypokalemia also causes a myopathy and predisposes to rhabdomyolysis; notably, however, the patient had a normal creatine phosphokinase (CPK) level. Cushing's syndrome is often accompanied by a proximal myopathy, due to the protein-wasting effects of cortisol excess.

The patient presented with a mixed acid-base disorder, with a significant metabolic alkalosis and a bicarbonate concentration of 44 meq/L. A venous blood gas was drawn soon after his presentation; venous and arterial blood gases demonstrate a high level of agreement in hemodynamically stable patients, allowing for the interpretation of acid-base disorders with venous blood gas results. In response to his metabolic alkalosis, the P_{CO_2} should have increased by 0.75 mmHg for each 1-meq/L increase in bicarbonate; the expected P_{CO_2} should have been ~55 mmHg. Given the P_{CO_2} of 62 mmHg, he had an additional respiratory acidosis, likely caused by respiratory muscle weakness from his acute hypokalemia and subacute hypercortisolism.

The patient's albumin-adjusted AG was $21 + ((4 - 2.8) \times 2.5) = 24$; this suggests a third acid-base disorder, AG acidosis. Notably, the measured AG can increase in alkalosis, due to both increases in plasma protein concentrations (in hypovolemic alkalosis) and to the alkalemia-associated increase in net negative charge of plasma proteins, both causing an increase in unmeasured anions; however, this patient was neither volume-depleted nor particularly alkalemic, suggesting that these effects played a minimal role in his increased AG. Alkalosis also stimulates an increase in lactic acid production, due to activation of phosphofructokinase and accelerated glycolysis; unfortunately, however, a lactic acid level was not measured in this patient. It should be noted in this regard that alkalosis typically increases lactic acid levels by a mere 1.5–3 meq/L and that the patient was not significantly alkalemic. Regardless of the underlying pathophysiology, the increased AG was likely related to the metabolic alkalosis, given that the AG had decreased to 18 by hospital day 2, coincident with a reduction in plasma bicarbonate.

Why did the patient have a metabolic alkalosis? The activation of MLRs in the distal nephron increases distal nephron acidification and net acid secretion. In consequence, mineralocorticoid excess causes a saline-resistant metabolic alkalosis, which is exacerbated significantly by the development of hypokalemia. Hypokalemia plays a key role in the generation of most forms of metabolic alkalosis, stimulating proximal tubular ammonium production, proximal tubular bicarbonate reabsorption, and distal tubular H^+/K^+ -ATPase activity.

APPROACH TO MANAGEMENT

The first priority in the management of this patient was to increase his plasma K^+ and magnesium concentrations rapidly; hypomagnesemic patients are refractory to K^+ replacement alone, resulting in the need to correct hypomagnesemia immediately. This was accomplished via the administration of both oral and intravenous K^+ - Cl^- , giving a total of 240 meq over the first 18 h; 5 g of intravenous magnesium sulfate was also administered. Multiple 100-mL "minibags" of saline containing

20 meq each were infused, with cardiac monitoring and frequent measurement of plasma electrolytes. Of note, intravenous K^+-Cl^- should always be given in saline solutions because dextrose-containing solutions can increase insulin levels and exacerbate hypokalemia.

This case illustrates the difficulty in predicting the whole-body deficit of K^+ in hypokalemic patients. In the absence of abnormal K^+ redistribution, the total deficit correlates with plasma K^+ concentration, which drops by approximately 0.27 mM for every 100-mmol reduction in total-body stores; this would suggest a deficit of ~650 meq of K^+ in this patient, at the admission plasma K^+ concentration of 1.7 meq/L. Notably, however, alkalemia induces a modest intracellular shift of circulating K^+ such that this patient's initial plasma K^+ concentration was not an ideal indicator of the total potassium deficit. Regardless of the underlying pathophysiology in this case, close monitoring of plasma K^+ concentration is always essential during the correction of severe hypokalemia in order to gauge the adequacy of repletion and to avoid overcorrection.

Subsequent management of this patient's Cushing's syndrome and ectopic ACTH secretion was complicated by the respiratory issues. The prognosis in patients with ectopic ACTH secretion depends on the tumor histology and the presence or absence of distant metastases. This patient had an exceptionally poor prognosis, with widely metastatic small-cell lung cancer that had failed treatment; other patients with ectopic ACTH secretion caused by more benign, isolated tumors, most commonly bronchial carcinoid tumors, have a much better prognosis. In the absence of successful surgical resection of the causative tumor, management of this syndrome can include surgical adrenalectomy or medical therapy to block adrenal steroid production.

CASE 9

A stuporous 22-year-old man was admitted with a history of behaving strangely. His friends indicated he experienced recent emotional problems stemming from a failed relationship and had threatened suicide. There was a history of alcohol abuse, but his friends were unaware of recent alcohol consumption. The patient was obtunded on admission, with no evident focal neurologic deficits. The remainder of the physical examination was unremarkable.

Laboratory Data	Value	Units
Na^+	140	meq/L
K^+	5	meq/L
Cl^-	95	meq/L
HCO_3^-	10	meq/L
Glucose	125	mg/dL
BUN	15	mg/dL
Creatinine	0.9	mg/dL
Ionized calcium	4.0	mg/dL
Plasma osmolality	325	mOsm/kg/ H_2O

Urinalysis revealed crystalluria, with a mixture of envelope-shaped and needle-shaped crystals.

APPROACH TO DIAGNOSIS

This patient presented with CNS manifestations and a history of suspicious behavior, suggesting ingestion of a toxin. The AG was strikingly elevated at 35 meq/L. The Δ AG of 25 significantly exceeded the ΔHCO_3^- of 15. The fact that the Δ values were significantly disparate indicates that the most likely acid-base diagnosis in this patient is a mixed high-AG metabolic acidosis and a metabolic alkalosis. The metabolic alkalosis in this case may have been the result of vomiting. Nevertheless, the most useful finding is that the osmolar gap is elevated. The osmolar gap of 33 (difference in measured and calculated osmolality or 325 – 292) in the face of a high-AG metabolic acidosis is diagnostic of an osmotically active metabolite in plasma; a difference of >10 mOsm/kg indicates a significant concentration of an unmeasured osmolyte. Examples of toxic osmolytes include ethylene glycol, diethylene glycol, methanol, and propylene glycol.

Several caveats apply to the interpretation of the osmolar gap and AG in the differential diagnosis of toxic alcohol ingestions. First, unmeasured, neutral osmolytes can also accumulate in lactic acidosis and alcoholic ketoacidosis; i.e., an elevated osmolar gap is not specific to AG acidoses associated with toxic alcohol ingestions. Second, patients can present having extensively metabolized the ingested toxin, with an insignificant osmolar gap but a large AG; i.e., the absence of an elevated osmolar gap does not rule out toxic alcohol ingestion. Third, the converse can also be seen in patients who present earlier after ingestion of the toxin, i.e., a large osmolar gap with minimal elevation of the AG. Finally, clinicians should be aware of the effect of co-ingested ethanol, which can itself elevate the osmolar gap and can reduce metabolism of the toxic alcohols via competitive inhibition of alcohol dehydrogenase (see below), thus attenuating the expected increase in the AG.

Ethylene glycol is commonly available as antifreeze or solvents and may be ingested accidentally or as a suicide attempt. The metabolism of ethylene glycol by alcohol dehydrogenase generates acids such as glycoaldehyde, glycolic acid, and oxalic acid. The initial effects of intoxication are on the CNS and, in the earliest stages, mimic inebriation, but may quickly progress to full-blown coma. Delay in treatment is one of the most common causes of mortality with toxic alcohol poisoning. The kidney shows evidence of acute tubular injury with widespread deposition of calcium oxalate crystals within tubular epithelial cells. Cerebral edema is common, as is crystal deposition in the brain; the latter is irreversible.

The co-occurrent crystalluria is typical of ethylene glycol intoxication; both needle-shaped monohydrate and envelope-shaped dihydrate calcium oxalate crystals can be seen in the urine as the process evolves. Circulating oxalate can also complex with plasma calcium, reducing the ionized calcium as in this case.

Although ethylene glycol intoxication should be verified by measuring ethylene glycol levels, therapy must be initiated immediately in this life-threatening situation. Although therapy can be initiated with confidence in cases with known or witnessed ingestions, such histories are rarely available. Therapy should thus be initiated in patients with severe metabolic acidosis and elevated anion and osmolar gaps. Other diagnostic features, such as hypocalcemia or acute renal failure with crystalluria, can provide important confirmation for urgent, empiric therapy.

APPROACH TO MANAGEMENT

Because all four osmotically active toxic alcohols—ethylene glycol, diethylene glycol, methanol, and propylene glycol—are metabolized by alcohol dehydrogenase to generate toxic products, competitive inhibition of this key enzyme is common to the treatment of all four intoxications. The most potent inhibitor of alcohol dehydrogenase, and the drug of choice in this circumstance, is fomepizole (4-methyl pyrazole). Fomepizole should be administered intravenously as a loading dose (15 mg/kg) followed by doses of 10 mg/kg every 12 h for four doses, and then 15 mg/kg every 12 h thereafter until ethylene glycol levels have been reduced to <20 mg/dL and the patient is asymptomatic with a normal pH. Additional important components of the treatment of toxic alcohol ingestion include fluid resuscitation, thiamine, pyridoxine, folate, sodium bicarbonate, and hemodialysis. Hemodialysis is used to remove both the parent compound and toxic metabolites, but it also removes administered fomepizole, necessitating adjustment of dosage frequency. Gastric aspiration, induced emesis, or the use of activated charcoal is only effective if initiated within 30–60 min after ingestion of the toxin. When fomepizole is not available, ethanol, which has more than 10-fold affinity for alcohol dehydrogenase compared to other alcohols, may be substituted and is quite effective. Ethanol must be administered IV to achieve a blood level of 22 meq/L (100 mg/dL). A disadvantage of ethanol is the obtundation that follows its administration, which is additive to the CNS effects of ethylene glycol. Furthermore, if hemodialysis is used, the infusion rate of ethanol must be increased because it is rapidly dialyzed. In general, hemodialysis is indicated for all patients with ethylene glycol intoxication when the arterial pH is <7.3 or the osmolar gap exceeds 20 mOsm/kg H_2O .

The calcium ion plays a critical role in normal cellular function and signaling, regulating diverse physiologic processes such as neuromuscular signaling, cardiac contractility, hormone secretion, and blood coagulation. Thus, extracellular calcium concentrations are maintained within an exquisitely narrow range through a series of feedback mechanisms that involve parathyroid hormone (PTH) and the active vitamin D metabolite 1,25-dihydroxyvitamin D [$1,25(\text{OH})_2\text{D}$]. These feedback mechanisms are orchestrated by integrating signals between the parathyroid glands, kidney, intestine, and bone (Fig. 65-1; Chap. 423). Disorders of serum calcium concentration are relatively common and often serve as a harbinger of underlying disease. This chapter provides a brief summary of the approach to patients with altered serum calcium levels. See Chap. 424 for a detailed discussion of this topic.

HYPERCALCEMIA

ETIOLOGY

The causes of hypercalcemia can be understood and classified based on derangements in the normal feedback mechanisms that regulate serum calcium (Table 65-1). Excess PTH production, which is not appropriately suppressed by increased serum calcium concentrations, occurs in primary neoplastic disorders of the parathyroid glands (parathyroid adenomas; hyperplasia; or, rarely, carcinoma) that are associated with

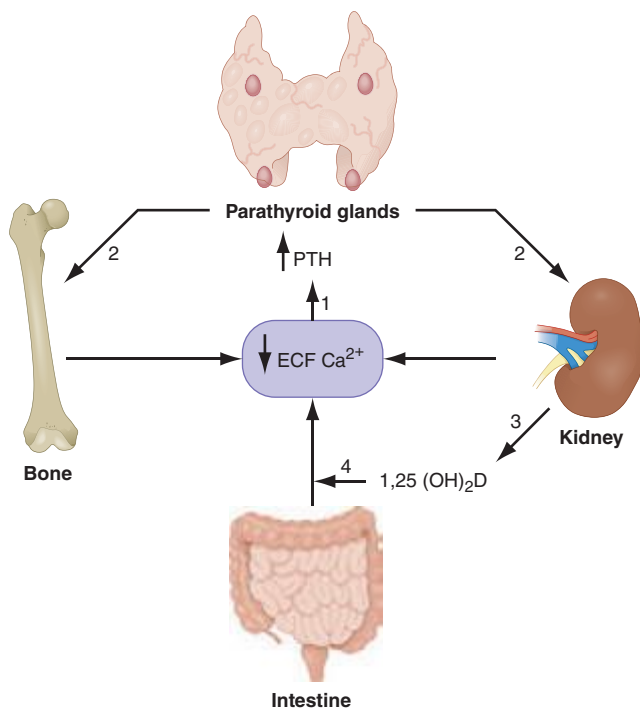


FIGURE 65-1 Feedback mechanisms maintaining extracellular calcium concentrations within a narrow, physiologic range (8.9–10.1 mg/dL [2.2–2.5 mM]). A decrease in extracellular (ECF) calcium (Ca^{2+}) triggers an increase in parathyroid hormone (PTH) secretion (1) via the calcium sensor receptor on parathyroid cells. PTH, in turn, results in increased tubular reabsorption of calcium by the kidney (2) and resorption of calcium from bone (2) and also stimulates renal $1,25(\text{OH})_2\text{D}$ production (3). $1,25(\text{OH})_2\text{D}$, in turn, acts principally on the intestine to increase calcium absorption (4). Collectively, these homeostatic mechanisms serve to restore serum calcium levels to normal.

TABLE 65-1 CAUSES OF HYPERCALCEMIA

Excessive PTH production
Primary hyperparathyroidism (adenoma, hyperplasia, rarely carcinoma)
Tertiary hyperparathyroidism (long-term stimulation of PTH secretion in renal insufficiency)
Ectopic PTH secretion (very rare)
Inactivating mutations in the CaSR or in G proteins (FHH)
Alterations in CaSR function (lithium therapy)
Hypercalcemia of malignancy
Overproduction of PTHrP (many solid tumors)
Lytic skeletal metastases (breast, myeloma)
Excessive $1,25(\text{OH})_2\text{D}$ production
Granulomatous diseases (sarcoidosis, tuberculosis, silicosis)
Lymphomas
Vitamin D intoxication
Primary increase in bone resorption
Hyperthyroidism
Immobilization
Excessive calcium intake
Milk-alkali syndrome
Total parenteral nutrition
Other causes
Endocrine disorders (adrenal insufficiency, pheochromocytoma, VIPoma)
Medications (thiazides, vitamin A, antiestrogens)

Abbreviations: CaSR, calcium sensor receptor; FHH, familial hypocalciuric hypercalcemia; PTH, parathyroid hormone; PTHrP, PTH-related peptide.

increased parathyroid cell mass and impaired feedback inhibition by calcium. Inappropriate PTH secretion for the ambient level of serum calcium also occurs with heterozygous inactivating calcium sensor receptor (CaSR) or G protein mutations, which impair extracellular calcium sensing by the parathyroid glands and the kidneys, resulting in familial hypocalciuric hypercalcemia (FHH). Although PTH secretion by tumors is extremely rare, many solid tumors produce PTH-related peptide (PTHrP), which shares homology with PTH in the first 13 amino acids and binds the PTH receptor, thus mimicking effects of PTH on bone and the kidney. In PTHrP-mediated hypercalcemia of malignancy, PTH levels are suppressed by the high serum calcium levels. Hypercalcemia associated with granulomatous disease (e.g., sarcoidosis) or lymphomas is caused by enhanced conversion of $25(\text{OH})\text{D}$ to the potent $1,25(\text{OH})_2\text{D}$. In these disorders, $1,25(\text{OH})_2\text{D}$ enhances intestinal calcium absorption, resulting in hypercalcemia and suppressed PTH. Disorders that directly increase calcium mobilization from bone, such as hyperthyroidism or osteolytic metastases, also lead to hypercalcemia with suppressed PTH secretion as does exogenous calcium overload, as in milk-alkali syndrome, or total parenteral nutrition with excessive calcium supplementation.

CLINICAL MANIFESTATIONS

Mild hypercalcemia (up to 11–11.5 mg/dL) is usually asymptomatic and recognized only on routine calcium measurements. Some patients may complain of vague neuropsychiatric symptoms, including trouble concentrating, personality changes, or depression. Other presenting symptoms may include peptic ulcer disease or nephrolithiasis, and fracture risk may be increased. More severe hypercalcemia (>12–13 mg/dL), particularly if it develops acutely, may result in lethargy, stupor, or coma, as well as gastrointestinal symptoms (nausea, anorexia, constipation, or pancreatitis). Hypercalcemia decreases renal concentrating ability, which may cause polyuria and polydipsia. With longstanding hyperparathyroidism, patients may present with bone pain or pathologic fractures. Finally, hypercalcemia can result in significant electrocardiographic changes, including bradycardia, AV block, and short QT interval; changes in serum calcium can be monitored by following the QT interval.

The first step in the diagnostic evaluation of hyper- or hypocalcemia is to ensure that the alteration in serum calcium levels is not due to abnormal albumin concentrations. About 50% of total calcium is ionized, and the rest is bound principally to albumin. Although direct measurements of ionized calcium are possible, they are easily influenced by collection methods and other artifacts; thus, it is generally preferable to measure total calcium and albumin to “correct” the serum calcium. When serum albumin concentrations are reduced, a corrected calcium concentration is calculated by adding 0.2 mM (0.8 mg/dL) to the total calcium level for every decrement in serum albumin of 1.0 g/dL below the reference value of 4.1 g/dL for albumin, and, conversely, for elevations in serum albumin.

A detailed history may provide important clues regarding the etiology of the hypercalcemia (Table 65-1). Chronic hypercalcemia is most commonly caused by primary hyperparathyroidism, as opposed to the second most common etiology of hypercalcemia, an underlying malignancy. The history should include medication use, previous neck surgery, and systemic symptoms suggestive of sarcoidosis or lymphoma.

Once true hypercalcemia is established, the second most important laboratory test in the diagnostic evaluation is a PTH level using a two-site assay for the intact hormone. Increases in PTH are often accompanied by hypophosphatemia. In addition, serum creatinine should be measured to assess renal function; hypercalcemia may impair renal function, and renal clearance of PTH may be altered depending on the fragments detected by the assay. If the PTH level is increased (or “inappropriately normal”) in the setting of elevated calcium and low phosphorus, the diagnosis is almost always primary hyperparathyroidism. Because individuals with FHH may also present with mildly elevated PTH levels and hypercalcemia, this diagnosis should be considered and excluded because parathyroid surgery is ineffective in this condition. A calcium/creatinine clearance ratio (calculated as urine calcium/serum calcium divided by urine creatinine/serum creatinine) of <0.01 is suggestive of FHH, particularly when there is a family history of mild, asymptomatic hypercalcemia. In addition, sequence analysis of the CaSR gene is now commonly performed for the definitive diagnosis of FHH, although in some families, FHH may be caused by mutations in G proteins that mediate signaling by the CaSR. Ectopic PTH secretion is extremely rare.

A suppressed PTH level in the face of hypercalcemia is consistent with non-parathyroid-mediated hypercalcemia, most often due to underlying malignancy. Although a tumor that causes hypercalcemia is generally overt, a PTHrP level may be needed to establish the diagnosis of hypercalcemia of malignancy. Serum 1,25(OH)₂D levels are increased in granulomatous disorders, and clinical evaluation in combination with laboratory testing will generally provide a diagnosis for the various disorders listed in Table 65-1.

TREATMENT HYPERCALCEMIA

Mild, asymptomatic hypercalcemia does not require immediate therapy, and management should be dictated by the underlying diagnosis. By contrast, significant, symptomatic hypercalcemia usually requires therapeutic intervention independent of the etiology of hypercalcemia. Initial therapy of significant hypercalcemia begins with volume expansion because hypercalcemia invariably leads to dehydration; 4–6 L of intravenous saline may be required over the first 24 h, keeping in mind that underlying comorbidities (e.g., congestive heart failure) may require the use of loop diuretics to enhance sodium and calcium excretion. However, loop diuretics should not be initiated until the volume status has been restored to normal. If there is increased calcium mobilization from bone (as in malignancy or severe hyperparathyroidism), drugs that inhibit bone resorption should be considered. Zoledronic acid (e.g., 4 mg intravenously over ~30 min), pamidronate (e.g., 60–90 mg intravenously over 2–4 h), and ibandronate (2 mg intravenously over 2 h) are bisphosphonates that are commonly used for the treatment of hypercalcemia of malignancy in adults. Onset of action is within

1–3 days, with normalization of serum calcium levels occurring in 60–90% of patients. Bisphosphonate infusions may need to be repeated if hypercalcemia relapses. An alternative to the bisphosphonates is gallium nitrate (200 mg/m² intravenously daily for 5 days), which is also effective, but has potential nephrotoxicity. In rare instances, dialysis may be necessary. Finally, although intravenous phosphate chelates calcium and decreases serum calcium levels, this therapy can be toxic because calcium-phosphate complexes may deposit in tissues and cause extensive organ damage.

In patients with 1,25(OH)₂D-mediated hypercalcemia, glucocorticoids are the preferred therapy, as they decrease 1,25(OH)₂D production. Intravenous hydrocortisone (100–300 mg daily) or oral prednisone (40–60 mg daily) for 3–7 days is used most often. Other drugs, such as ketoconazole, chloroquine, and hydroxychloroquine, may also decrease 1,25(OH)₂D production and are used occasionally.

HYPOCALCEMIA

ETIOLOGY

The causes of hypocalcemia can be differentiated according to whether serum PTH levels are low (hypoparathyroidism) or high (secondary hyperparathyroidism). Although there are many potential causes of hypocalcemia, impaired PTH production and impaired vitamin D production are the most common etiologies (Table 65-2) (Chap. 424). Because PTH is the main defense against hypocalcemia, disorders associated with deficient PTH production or secretion may be associated with profound, life-threatening hypocalcemia. In adults, hypoparathyroidism most commonly results from inadvertent damage to all four glands during thyroid or parathyroid gland surgery. Hypoparathyroidism is a cardinal feature of autoimmune endocrinopathies (Chap. 408); rarely, it

TABLE 65-2 CAUSES OF HYPOCALCEMIA

Low Parathyroid Hormone Levels (Hypoparathyroidism)

Parathyroid agenesis
Isolated
DiGeorge's syndrome
Parathyroid destruction
Surgical
Radiation
Infiltration by metastases or systemic diseases
Autoimmune
Reduced parathyroid function
Hypomagnesemia
Activating CaSR or G protein mutations

High Parathyroid Hormone Levels (Secondary Hyperparathyroidism)

Vitamin D deficiency or impaired 1,25(OH) ₂ D production/action
Nutritional vitamin D deficiency (poor intake or absorption)
Renal insufficiency with impaired 1,25(OH) ₂ D production
Vitamin D resistance, including receptor defects
Parathyroid hormone resistance syndromes
PTH receptor mutations
Pseudohypoparathyroidism (G protein mutations)
Drugs
Calcium chelators
Inhibitors of bone resorption (bisphosphonates, plicamycin)
Altered vitamin D metabolism (phenytoin, ketoconazole)
Miscellaneous causes
Acute pancreatitis
Acute rhabdomyolysis
Hungry bone syndrome after parathyroidectomy
Osteoblastic metastases with marked stimulation of bone formation (prostate cancer)

Abbreviations: CaSR, calcium sensor receptor; PTH, parathyroid hormone.

may be associated with infiltrative diseases such as sarcoidosis. Impaired PTH secretion may be secondary to magnesium deficiency or to activating mutations in the CaSR or in the G proteins that mediate CaSR signaling, which suppress PTH, leading to effects that are opposite to those that occur in FHH.

Vitamin D deficiency, impaired 1,25(OH)₂D production (primarily secondary to renal insufficiency), or vitamin D resistance also cause hypocalcemia. However, the degree of hypocalcemia in these disorders is generally not as severe as that seen with hypoparathyroidism because the parathyroids are capable of mounting a compensatory increase in PTH secretion. Hypocalcemia may also occur in conditions associated with severe tissue injury such as burns, rhabdomyolysis, tumor lysis, or pancreatitis. The cause of hypocalcemia in these settings may include a combination of low albumin, hyperphosphatemia, tissue deposition of calcium, and impaired PTH secretion.

CLINICAL MANIFESTATIONS

Patients with hypocalcemia may be asymptomatic if the decreases in serum calcium are relatively mild and chronic, or they may present with life-threatening complications. Moderate to severe hypocalcemia is associated with paresthesias, usually of the fingers, toes, and circumoral regions, and is caused by increased neuromuscular irritability. On physical examination, a Chvostek's sign (twitching of the circumoral muscles in response to gentle tapping of the facial nerve just anterior to the ear) may be elicited, although it is also present in ~10% of normal individuals. Carpal spasm may be induced by inflation of a blood pressure cuff to 20 mmHg above the patient's systolic blood pressure for 3 min (Trousseau's sign). Severe hypocalcemia can induce seizures, carpopedal spasm, bronchospasm, laryngospasm, and prolongation of the QT interval.

DIAGNOSTIC APPROACH

In addition to measuring serum calcium, it is useful to determine albumin, phosphorus, and magnesium levels. As for the evaluation of hypercalcemia, determining the PTH level is central to the evaluation of hypocalcemia. A suppressed (or "inappropriately low") PTH level in the setting of hypocalcemia establishes absent or reduced PTH secretion (hypoparathyroidism) as the cause of the hypocalcemia. Further history will often elicit the underlying cause (i.e., parathyroid agenesis vs. destruction). By contrast, an elevated PTH level (secondary hyperparathyroidism) should direct attention to the vitamin D axis as the cause of the hypocalcemia. Nutritional vitamin D deficiency is best assessed by obtaining serum 25-hydroxyvitamin D levels, which reflect vitamin D stores. In the setting of renal insufficiency or suspected vitamin D resistance, serum 1,25(OH)₂D levels are informative.

TREATMENT HYPOCALCEMIA

The approach to treatment depends on the severity of the hypocalcemia, the rapidity with which it develops, and the accompanying complications (e.g., seizures, laryngospasm). Acute, symptomatic hypocalcemia is initially managed with calcium gluconate, 10 mL 10% wt/vol (90 mg or 2.2 mmol) intravenously, diluted in 50 mL of 5% dextrose or 0.9% sodium chloride, given intravenously over 5 min. Continuing hypocalcemia often requires a constant intravenous infusion (typically 10 ampules of calcium gluconate or 900 mg of calcium in 1 L of 5% dextrose or 0.9% sodium chloride administered over 24 h). Accompanying hypomagnesemia, if present, should be treated with appropriate magnesium supplementation.

Chronic hypocalcemia due to hypoparathyroidism is treated with calcium supplements (1000–1500 mg/d elemental calcium in divided doses) and either vitamin D₂ or D₃ (25,000–100,000 U daily) or calcitriol [1,25(OH)₂D, 0.25–2 µg/d]. Other vitamin D metabolites (dihydroxycholesterol, alfalcidol) are now used less frequently. Vitamin D deficiency, however, is best treated using vitamin D supplementation, with the dose depending on the severity of the deficit and the underlying cause. Thus, nutritional vitamin D deficiency generally responds to relatively low doses of vitamin D (50,000 U, 2–3 times per week for several months), whereas vitamin

D deficiency due to malabsorption may require much higher doses (100,000 U/d or more). The treatment goal is to bring serum calcium into the low normal range and to avoid hypercalciuria, which may lead to nephrolithiasis.

GLOBAL CONSIDERATIONS



In countries with more limited access to health care or screening laboratory testing of serum calcium levels, primary hyperparathyroidism often presents in its severe form with skeletal complications (osteitis fibrosa cystica) in contrast to the asymptomatic form that is common in developed countries. In addition, vitamin D deficiency is paradoxically common in some countries despite extensive sunlight (e.g., India) due to avoidance of sun exposure and poor dietary vitamin D intake.

66 Acidosis and Alkalosis

Thomas D. DuBose, Jr.

NORMAL ACID-BASE HOMEOSTASIS

Systemic arterial pH is maintained between 7.35 and 7.45 by extracellular and intracellular chemical buffering together with respiratory and renal regulatory mechanisms. The control of arterial CO₂ tension (Paco₂) by the central nervous system (CNS) and respiratory system and the control of plasma bicarbonate by the kidneys stabilize the arterial pH by excretion or retention of acid or alkali. The metabolic and respiratory components that regulate systemic pH are described by the Henderson-Hasselbalch equation:

$$\text{pH} = 6.1 + \log \frac{[\text{HCO}_3^-]}{\text{Pa}_{\text{CO}_2} \times 0.0301}$$

Under most circumstances, CO₂ production and excretion are matched, and the usual steady-state Paco₂ is maintained at 40 mmHg. Underexcretion of CO₂ produces hypercapnia, and overexcretion causes hypocapnia. Nevertheless, production and excretion are again matched at a new steady-state Paco₂. Therefore, the Paco₂ is regulated primarily by neural respiratory factors and is not subject to regulation by the rate of CO₂ production. Hypercapnia is usually the result of hypoventilation rather than of increased CO₂ production. Increases or decreases in Paco₂ represent derangements of neural respiratory control or are due to compensatory changes in response to a primary alteration in the plasma [HCO₃⁻].

DIAGNOSIS OF GENERAL TYPES OF DISTURBANCES

The most common clinical disturbances are simple acid-base disorders; i.e., metabolic acidosis or alkalosis or respiratory acidosis or alkalosis.

SIMPLE ACID-BASE DISORDERS

Primary respiratory disturbances (primary changes in Paco₂) invoke compensatory metabolic responses (secondary changes in [HCO₃⁻]), and primary metabolic disturbances elicit predictable compensatory respiratory responses (secondary changes in Paco₂). Physiologic compensation can be predicted from the relationships displayed in **Table 66-1**. In general, with one exception, compensatory responses return the pH toward, but not to, the normal value. Chronic respiratory alkalosis when prolonged is an exception to this rule and often returns the pH to a normal value. Metabolic acidosis due to an increase in endogenous acids (e.g., ketoacidosis) lowers extracellular fluid [HCO₃⁻] and decreases extracellular pH. This stimulates the medullary chemoreceptors to increase ventilation and to return

TABLE 66-1 PREDICTION OF COMPENSATORY RESPONSES ON SIMPLE ACID-BASE DISTURBANCES AND PATTERN OF CHANGES

Disorder	Prediction of Compensation	Range of Values		
		pH	HCO ₃ ⁻	Paco ₂
Metabolic acidosis	Paco ₂ = (1.5 × HCO ₃ ⁻) + 8 ± 2 or Paco ₂ will ↓ 1.25 mmHg per mmol/L ↓ in [HCO ₃ ⁻] or Paco ₂ = [HCO ₃ ⁻] + 15	Low	Low	Low
Metabolic alkalosis	Paco ₂ will ↑ 0.75 mmHg per mmol/L ↑ in [HCO ₃ ⁻] or Paco ₂ will ↑ 6 mmHg per 10 mmol/L ↑ in [HCO ₃ ⁻] or Paco ₂ = [HCO ₃ ⁻] + 15	High	High	High
Respiratory alkalosis		High	Low	Low
Acute	[HCO ₃ ⁻] will ↓ 0.2 mmol/L per mmHg ↓ in Paco ₂			
Chronic	[HCO ₃ ⁻] will ↓ 0.4 mmol/L per mmHg ↓ in Paco ₂			
Respiratory acidosis		Low	High	High
Acute	[HCO ₃ ⁻] will ↑ 0.1 mmol/L per mmHg ↑ in Paco ₂			
Chronic	[HCO ₃ ⁻] will ↑ 0.4 mmol/L per mmHg ↑ in Paco ₂			

the ratio of [HCO₃⁻] to Paco₂, and thus pH, toward, but not to, normal. The degree of respiratory compensation expected in a simple form of metabolic acidosis can be predicted from the relationship: Paco₂ = (1.5 × [HCO₃⁻]) + 8 ± 2. Thus, a patient with metabolic acidosis and [HCO₃⁻] of 12 mmol/L would be expected to have a Paco₂ between 24 and 28 mmHg. Values for Paco₂ <24 or >28 mmHg define a mixed disturbance (metabolic acidosis and respiratory alkalosis or metabolic alkalosis and respiratory acidosis, respectively). Compensatory responses for primary metabolic disorders move the Paco₂ in the same direction as the change in [HCO₃⁻], whereas, conversely, compensation for primary respiratory disorders moves the [HCO₃⁻] in the same direction as the primary change in Paco₂ (Table 66-1). Therefore, changes in Paco₂ and [HCO₃⁻] in **opposite directions** (i.e., Paco₂ or [HCO₃⁻] is increased, whereas the other value is decreased) indicate a **mixed disturbance**. Another way to judge the appropriateness of the response in [HCO₃⁻] or Paco₂ is to use an acid-base nomogram (Fig. 66-1). While the shaded areas of the nomogram show the 95% confidence limits for normal compensation in simple disturbances, finding acid-base values within the shaded area does not necessarily rule out a mixed disturbance. Imposition of one disorder over another may result in values lying within the area of a third. Thus, the nomogram, while convenient, is not a substitute for the equations in Table 66-1.

MIXED ACID-BASE DISORDERS

Mixed acid-base disorders—defined as independently coexisting disorders, not merely compensatory responses—are often seen in patients in critical care units and can lead to dangerous extremes of pH (Table 66-2). A patient with diabetic ketoacidosis (metabolic acidosis) may develop an independent respiratory problem (e.g.,

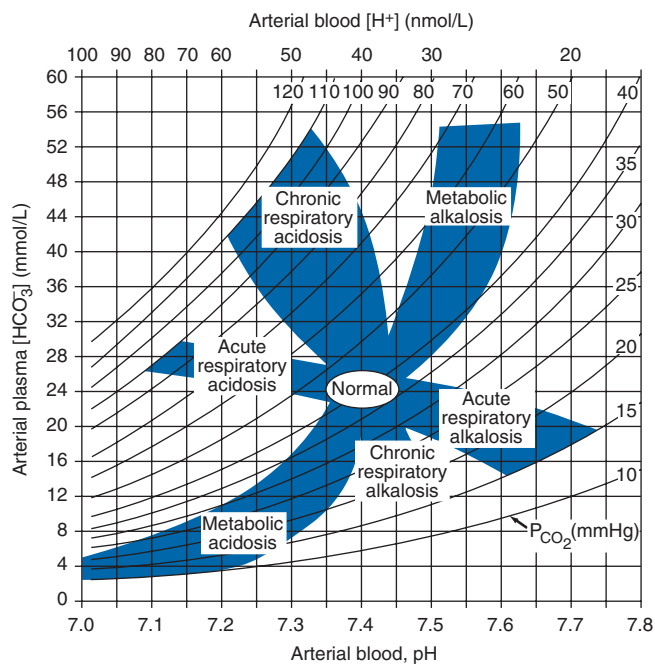


FIGURE 66-1 Acid-base nomogram. Shown are the 90% confidence limits (range of values) of the normal respiratory and metabolic compensations for primary acid-base disturbances. (From TD DuBose Jr: *Acid-base disorders*, in Brenner and Rector's *The Kidney*, 8th ed, BM Brenner [ed]. Philadelphia, Saunders, 2008, pp 505–546, with permission.)

TABLE 66-2 EXAMPLES OF MIXED ACID-BASE DISORDERS

Mixed Metabolic and Respiratory

Metabolic acidosis—respiratory alkalosis

Key: High- or normal-AG metabolic acidosis; prevailing Paco₂ below predicted value (Table 66-1)

Example: Na⁺, 140; K⁺, 4.0; Cl⁻, 106; HCO₃⁻, 14; AG, 20; Paco₂, 24; pH, 7.39 (lactic acidosis, sepsis in ICU)

Metabolic acidosis—respiratory acidosis

Key: High- or normal-AG metabolic acidosis; prevailing Paco₂ above predicted value (Table 66-1)

Example: Na⁺, 140; K⁺, 4.0; Cl⁻, 102; HCO₃⁻, 18; AG, 20; Paco₂, 38; pH, 7.30 (severe pneumonia, pulmonary edema)

Metabolic alkalosis—respiratory alkalosis

Key: Paco₂ does not increase as predicted; pH higher than expected

Example: Na⁺, 140; K⁺, 4.0; Cl⁻, 91; HCO₃⁻, 33; AG, 16; Paco₂, 38; pH, 7.55 (liver disease and diuretics)

Metabolic alkalosis—respiratory acidosis

Key: Paco₂ higher than predicted; pH normal

Example: Na⁺, 140; K⁺, 3.5; Cl⁻, 88; HCO₃⁻, 42; AG, 10; Paco₂, 67; pH, 7.42 (COPD on diuretics)

Mixed Metabolic Disorders

Metabolic acidosis—metabolic alkalosis

Key: Only detectable with high-AG acidosis; ΔAG >> ΔHCO₃⁻

Example: Na⁺, 140; K⁺, 3.0; Cl⁻, 95; HCO₃⁻, 25; AG, 20; Paco₂, 40; pH, 7.42 (uremia with vomiting)

Metabolic acidosis—metabolic acidosis

Key: Mixed high-AG—normal-AG acidosis; ΔHCO₃⁻ accounted for by combined change in ΔAG and ΔCl⁻

Example: Na⁺, 135; K⁺, 3.0; Cl⁻, 110; HCO₃⁻, 10; AG, 15; Paco₂, 25; pH, 7.20 (diarrhea and lactic acidosis, toluene toxicity, treatment of diabetic ketoacidosis)

Abbreviations: AG, anion gap; COPD, chronic obstructive pulmonary disease; ICU, intensive care unit.

pneumonia) leading to respiratory acidosis or alkalosis. Patients with underlying pulmonary disease (e.g., chronic obstructive pulmonary disease) may not respond to metabolic acidosis with an appropriate ventilatory response because of insufficient respiratory reserve. Such imposition of respiratory acidosis on metabolic acidosis can lead to severe acidemia. When metabolic acidosis and metabolic alkalosis coexist in the same patient, the pH may be normal or near normal. When the pH is normal, an elevated anion gap (AG; see below) reliably denotes the presence of an AG metabolic acidosis at a normal serum albumin of 4.5 g/dL. Assuming a normal AG of 10 mmol/L, a discrepancy in the Δ AG (prevailing minus normal AG) and the Δ HCO₃⁻ (normal value of 25 mmol/L minus abnormal HCO₃⁻ in the patient) indicates the presence of a mixed high-gap acidosis—metabolic alkalosis (see example below). A diabetic patient with ketoacidosis may have renal dysfunction resulting in simultaneous metabolic acidosis. Patients who have ingested an overdose of drug combinations such as sedatives and salicylates may have mixed disturbances as a result of the acid-base response to the individual drugs (metabolic acidosis mixed with respiratory acidosis or respiratory alkalosis, respectively). Triple acid-base disturbances are more complex. For example, patients with metabolic acidosis due to alcoholic ketoacidosis may develop metabolic alkalosis due to vomiting and superimposed respiratory alkalosis due to the hyperventilation of hepatic dysfunction or alcohol withdrawal.

APPROACH TO THE PATIENT: Acid-Base Disorders

A stepwise approach to the diagnosis of acid-base disorders follows (Table 66-3). Care should be taken when measuring blood gases to obtain the arterial blood sample without using excessive heparin. Blood for electrolytes and arterial blood gases should be drawn simultaneously prior to therapy, because an increase in [HCO₃⁻] occurs with metabolic alkalosis and respiratory acidosis. Conversely, a decrease in [HCO₃⁻] occurs in metabolic acidosis and respiratory alkalosis. In the determination of arterial blood gases by the clinical laboratory, both pH and PaCO₂ are measured, and the [HCO₃⁻] is calculated from the Henderson-Hasselbalch equation. This calculated value should be compared with the measured [HCO₃⁻] (total CO₂) on the electrolyte panel. These two values should agree within 2 mmol/L. If they do not, the values may not have been drawn simultaneously, a laboratory error may be present, or an error could have been made in calculating the [HCO₃⁻]. After verifying the blood acid-base values, the precise acid-base disorder can then be identified.

CALCULATE THE ANION GAP

All evaluations of acid-base disorders should include a simple calculation of the AG; it represents those unmeasured anions in plasma (normally 8–10 mmol/L) and is calculated as follows: $AG = Na^+ - (Cl^- + HCO_3^-)$. The unmeasured anions include anionic proteins (e.g., albumin), phosphate, sulfate, and organic anions. When acid anions, such as acetoacetate and lactate, accumulate in extracellular fluid, the AG increases, causing a high-AG

acidosis. An increase in the AG is most often due to an increase in unmeasured anions and, less commonly, is due to a decrease in unmeasured cations (calcium, magnesium, potassium). In addition, the AG may increase with an increase in anionic albumin, because of either increased albumin concentration or alkalosis, which alters albumin charge. A decrease in the AG can be due to (1) an increase in unmeasured cations; (2) the addition to the blood of abnormal cations, such as lithium (lithium intoxication) or cationic immunoglobulins (plasma cell dyscrasias); (3) a reduction in the major plasma anion albumin concentration (nephrotic syndrome); (4) a decrease in the effective anionic charge on albumin by acidosis; or (5) hyperviscosity and severe hyperlipidemia, which can lead to an underestimation of sodium and chloride concentrations. A fall in serum albumin by 1 g/dL from the normal value (4.5 g/dL) decreases the AG by 2.5 meq/L. Know the common causes of a high-AG acidosis (Table 66-3).

In the face of a normal serum albumin, a high AG is usually due to non-chloride-containing acids that contain inorganic (phosphate, sulfate), organic (ketoacids, lactate, uremic organic anions), exogenous (salicylate or ingested toxins with organic acid production), or unidentified anions. The high AG is significant even if an additional acid-base disorder is superimposed to modify the [HCO₃⁻] independently. Simultaneous metabolic acidosis of the high-AG variety plus either chronic respiratory acidosis or metabolic alkalosis represents such a situation in which [HCO₃⁻] may be normal or even high (Table 66-3). Compare the change in [HCO₃⁻] (Δ HCO₃⁻) and the change in the AG (Δ AG).

Similarly, normal values for [HCO₃⁻], PaCO₂, and pH do not ensure the absence of an acid-base disturbance. For instance, an alcoholic who has been vomiting may develop a metabolic alkalosis with a pH of 7.55, PaCO₂ of 47 mmHg, [HCO₃⁻] of 40 mmol/L, [Na⁺] of 135, [Cl⁻] of 80, and [K⁺] of 2.8. If such a patient were then to develop a superimposed alcoholic ketoacidosis with a β -hydroxybutyrate concentration of 15 mM, arterial pH would fall to 7.40, [HCO₃⁻] to 25 mmol/L, and the PaCO₂ to 40 mmHg. Although these blood gases are normal, the AG is elevated at 30 mmol/L, indicating a mixed metabolic alkalosis and metabolic acidosis. A mixture of high-gap acidosis and metabolic alkalosis is recognized easily by comparing the differences (Δ values) in the normal to prevailing patient values. In this example, the Δ HCO₃⁻ is 0 (25 – 25 mmol/L), but the Δ AG is 20 (30 – 10 mmol/L). Therefore, 20 mmol/L is unaccounted for in the Δ/Δ value (Δ AG to Δ HCO₃⁻).

METABOLIC ACIDOSIS

Metabolic acidosis can occur because of an increase in endogenous acid production (such as lactate and ketoacids), loss of bicarbonate (as in diarrhea), or accumulation of endogenous acids (as in renal failure). Metabolic acidosis has profound effects on the respiratory, cardiac, and nervous systems. The fall in blood pH is accompanied by a characteristic increase in ventilation, especially the tidal volume (Kussmaul respiration). Intrinsic cardiac contractility may be depressed, but inotropic function can be normal because of catecholamine release. Both peripheral arterial vasodilation and central venoconstriction can be present; the decrease in central and pulmonary vascular compliance predisposes to pulmonary edema with even minimal volume overload. CNS function is depressed, with headache, lethargy, stupor, and, in some cases, even coma. Glucose intolerance may also occur.

There are two major categories of clinical metabolic acidosis: high-AG and non-AG, or hyperchloremic, acidosis (Table 66-3 and Table 66-4).

TREATMENT METABOLIC ACIDOSIS

Treatment of metabolic acidosis with alkali should be reserved for severe acidemia except when the patient has no “potential HCO₃⁻” in plasma. Potential [HCO₃⁻] can be estimated from the increment (Δ) in the AG (Δ AG = patient’s AG – 10). It must be determined if

TABLE 66-3 STEPS IN ACID-BASE DIAGNOSIS

1. Obtain arterial blood gas (ABG) and electrolytes simultaneously.
2. Compare [HCO₃⁻] on ABG and electrolytes to verify accuracy.
3. Calculate anion gap (AG).
4. Know four causes of high-AG acidosis (ketoacidosis, lactic acid acidosis, renal failure, and toxins).
5. Know two causes of hyperchloremic or nongap acidosis (bicarbonate loss from gastrointestinal tract, renal tubular acidosis).
6. Estimate compensatory response (Table 66-1).
7. Compare Δ AG and Δ HCO₃⁻.
8. Compare change in [Cl⁻] with change in [Na⁺].

TABLE 66-4 CAUSES OF HIGH-ANION GAP METABOLIC ACIDOSIS

Lactic acidosis	Toxins
Ketoacidosis	Ethylene glycol
Diabetic	Methanol
Alcoholic	Salicylates
Starvation	Propylene glycol
	Pyroglutamic acid (5-oxoproline)
	Renal failure (acute and chronic)

the acid anion in plasma is metabolizable (i.e., β -hydroxybutyrate, acetoacetate, and lactate) or nonmetabolizable (anions that accumulate in chronic renal failure and after toxin ingestion). The latter requires return of renal function to replenish the $[\text{HCO}_3^-]$ deficit, a slow and often unpredictable process. Consequently, patients with a normal AG acidosis (hyperchloremic acidosis), a slightly elevated AG (mixed hyperchloremic and AG acidosis), or an AG attributable to a nonmetabolizable anion in the face of renal failure should receive alkali therapy, either PO (NaHCO_3 or Shohl's solution) or IV (NaHCO_3), in an amount necessary to slowly increase the plasma $[\text{HCO}_3^-]$ into the 20–22 mmol/L range. Overcorrection must be avoided.

Controversy exists, however, in regard to the use of alkali in patients with a pure AG acidosis owing to accumulation of a metabolizable organic acid anion (ketoacidosis or lactic acidosis). In general, severe acidosis ($\text{pH} < 7.10$) warrants the IV administration of 50–100 meq of NaHCO_3 over 30–45 min, during the initial 1–2 h of therapy. Provision of such modest quantities of alkali in this situation seems to provide an added measure of safety, but it is essential to monitor plasma electrolytes during the course of therapy, because the $[\text{K}^+]$ may decline as pH rises. The goal is to increase the $[\text{HCO}_3^-]$ to 10 meq/L and the pH to approximately 7.20, not to increase these values to normal.

HIGH-ANION GAP ACIDOSES

APPROACH TO THE PATIENT: High-Anion Gap Acidoses

There are four principal causes of a high-AG acidosis: (1) lactic acidosis, (2) ketoacidosis, (3) ingested toxins, and (4) acute and chronic renal failure (Table 66-4). Initial screening to differentiate the high-AG acidoses should include (1) a probe of the history for evidence of drug and toxin ingestion and measurement of arterial blood gas to detect coexistent respiratory alkalosis (salicylates); (2) determination of whether diabetes mellitus is present (diabetic ketoacidosis); (3) a search for evidence of alcoholism or increased levels of β -hydroxybutyrate (alcoholic ketoacidosis); (4) observation for clinical signs of uremia and determination of the blood urea nitrogen (BUN) and creatinine (uremic acidosis); (5) inspection of the urine for oxalate crystals (ethylene glycol); and (6) recognition of the numerous clinical settings in which lactate levels may be increased (hypotension, shock, cardiac failure, leukemia, cancer, and drug or toxin ingestion).

Lactic Acidosis An increase in plasma l-lactate may be secondary to poor tissue perfusion (type A)—circulatory insufficiency (shock, cardiac failure), severe anemia, mitochondrial enzyme defects, and inhibitors (carbon monoxide, cyanide)—or to aerobic disorders (type B)—malignancies, nucleoside analogue reverse transcriptase inhibitors in HIV, diabetes mellitus, renal or hepatic failure, thiamine deficiency, severe infections (cholera, malaria), seizures, or drugs/toxins (biguanides, ethanol, methanol, propylene glycol, isoniazid, and fructose). Unrecognized bowel ischemia or infarction in a patient with severe atherosclerosis or cardiac decompensation receiving vasopressors is a common cause of lactic acidosis. Pyroglutamic acidemia has been reported in critically ill patients receiving acetaminophen, which is associated with depletion of glutathione. D-Lactic acid acidosis, which

may be associated with jejunioleal bypass, short bowel syndrome, or intestinal obstruction, is due to formation of D-lactate by gut bacteria.

APPROACH TO THE PATIENT: L-Lactic Acid Acidosis

The underlying condition that disrupts lactate metabolism must first be corrected; tissue perfusion must be restored when inadequate. Vasoconstrictors should be avoided, if possible, because they may worsen tissue perfusion. Alkali therapy is generally advocated for acute, severe acidemia ($\text{pH} < 7.15$) to improve cardiac function and lactate use. However, NaHCO_3 therapy may paradoxically depress cardiac performance and exacerbate acidosis by enhancing lactate production (HCO_3^- stimulates phosphofructokinase). While the use of alkali in moderate lactic acidosis is controversial, it is generally agreed that attempts to return the pH or $[\text{HCO}_3^-]$ to normal by administration of exogenous NaHCO_3 are deleterious. A reasonable approach is to infuse sufficient NaHCO_3 to raise the arterial pH to no more than 7.2 over 30–40 min.

NaHCO_3 therapy can cause fluid overload and hypertension because the amount required can be massive when accumulation of lactic acid is relentless. Fluid administration is poorly tolerated because of central venoconstriction, especially in the oliguric patient. When the underlying cause of the lactic acidosis can be remedied, blood lactate will be converted to HCO_3^- and may result in an overshoot alkalosis.

Ketoacidosis • DIABETIC KETOACIDOSIS (DKA) This condition is caused by increased fatty acid metabolism and the accumulation of ketoacids (acetoacetate and β -hydroxybutyrate). DKA usually occurs in insulin-dependent diabetes mellitus in association with cessation of insulin or an intercurrent illness such as an infection, gastroenteritis, pancreatitis, or myocardial infarction, which increases insulin requirements temporarily and acutely. The accumulation of ketoacids accounts for the increment in the AG and is accompanied most often by hyperglycemia (glucose > 17 mmol/L [300 mg/dL]). The relationship between the ΔAG and ΔHCO_3^- is typically ~1:1 in DKA. It should be noted that, because insulin prevents production of ketones, bicarbonate therapy is rarely needed except with extreme acidemia ($\text{pH} < 7.1$), and then in only limited amounts. Patients with DKA are typically volume depleted and require fluid resuscitation with isotonic saline. Volume overexpansion with IV fluid administration is not uncommon, however, and contributes to the development of a hyperchloremic acidosis during treatment of DKA. The mainstay for treatment of this condition is IV regular insulin and is described in [Chap. 417](#) in more detail.

ALCOHOLIC KETOACIDOSIS (AKA) Chronic alcoholics can develop ketoacidosis when alcohol consumption is abruptly curtailed and nutrition is poor. AKA is usually associated with binge drinking, vomiting, abdominal pain, starvation, and volume depletion. The glucose concentration is variable, and acidosis may be severe because of elevated ketones, predominantly β -hydroxybutyrate. Hypoperfusion may enhance lactic acid production, chronic respiratory alkalosis may accompany liver disease, and metabolic alkalosis can result from vomiting (refer to the relationship between ΔAG and ΔHCO_3^-). Thus, mixed acid-base disorders are common in AKA. As the circulation is restored by administration of isotonic saline, the preferential accumulation of β -hydroxybutyrate is then shifted to acetoacetate. This explains the common clinical observation of an increasingly positive nitroprusside reaction as the patient improves. The nitroprusside ketone reaction (Acetest) can detect acetoacetic acid but not β -hydroxybutyrate, so that the degree of ketosis and ketonuria can not only change with therapy, but can be underestimated initially. Patients with AKA usually present with relatively normal renal function, as opposed to DKA, where renal function is often compromised because of volume depletion (osmotic diuresis) or diabetic nephropathy. The AKA patient with normal renal function may excrete relatively large quantities of ketoacids in the urine and, therefore, may have a relatively normal AG and a discrepancy in the $\Delta\text{AG}/\Delta\text{HCO}_3^-$ relationship.

TREATMENT ALCOHOLIC KETOACIDOSIS

Extracellular fluid deficits almost always accompany AKA and should be replenished by IV administration of saline and glucose (5% dextrose in 0.9% NaCl). Hypophosphatemia, hypokalemia, and hypomagnesemia may coexist and should be corrected. Hypophosphatemia usually emerges 12–24 h after admission, may be exacerbated by glucose infusion, and, if severe, may induce rhabdomyolysis or even respiratory arrest. Upper gastrointestinal hemorrhage, pancreatitis, and pneumonia may accompany this disorder.

Drug- and Toxin-Induced Acidosis • SALICYLATES (See also Chap. 472e)

Salicylate intoxication in adults usually causes respiratory alkalosis or a mixture of high-AG metabolic acidosis and respiratory alkalosis. Only a portion of the AG is due to salicylates. Lactic acid production is also often increased.

TREATMENT SALICYLATE-INDUCED ACIDOSIS

Vigorous gastric lavage with isotonic saline (not NaHCO_3) should be initiated immediately, followed by administration of activated charcoal per nasogastric tube. In the acidotic patient, to facilitate removal of salicylate, IV NaHCO_3 is administered in amounts adequate to alkalinize the urine and to maintain urine output (urine pH >7.5). While this form of therapy is straightforward in acidotic patients, a coexisting respiratory alkalosis may make this approach hazardous. Alkalemic patients should not receive NaHCO_3 . Acetazolamide may be administered in the face of alkalemia, when an alkaline diuresis cannot be achieved, or to ameliorate volume overload associated with NaHCO_3 administration, but this drug can cause systemic metabolic acidosis if HCO_3^- is not replaced. Hypokalemia should be anticipated with an alkaline diuresis and should be treated promptly and aggressively. Glucose-containing fluids should be administered because of the danger of hypoglycemia. Excessive insensible fluid losses may cause severe volume depletion and hypernatremia. If renal failure prevents rapid clearance of salicylate, hemodialysis can be performed against a bicarbonate dialysate.

ALCOHOLS Under most physiologic conditions, sodium, urea, and glucose generate the osmotic pressure of blood. Plasma osmolality is calculated according to the following expression: $P_{\text{osm}} = 2\text{Na}^+ + \text{Glu} + \text{BUN}$ (all in mmol/L), or, using conventional laboratory values in which glucose and BUN are expressed in milligrams per deciliter: $P_{\text{osm}} = 2\text{Na}^+ + \text{Glu}/18 + \text{BUN}/2.8$. The calculated and determined osmolality should agree within 10–15 mmol/kg H_2O . When the measured osmolality exceeds the calculated osmolality by >10–15 mmol/kg H_2O , one of two circumstances prevails. Either the serum sodium is spuriously low, as with hyperlipidemia or hyperproteinemia (pseudohyponatremia), or osmolytes other than sodium salts, glucose, or urea have accumulated in plasma. Examples of such osmolytes include mannitol, radiocontrast media, ethanol, isopropyl alcohol, ethylene glycol, propylene glycol, methanol, and acetone. In this situation, the difference between the calculated osmolality and the measured osmolality (*osmolar gap*) is proportional to the concentration of the unmeasured solute. With an appropriate clinical history and index of suspicion, identification of an osmolar gap is helpful in identifying the presence of poison-associated AG acidosis. Three alcohols may cause fatal intoxications: ethylene glycol, methanol, and isopropyl alcohol. All cause an elevated osmolar gap, but only the first two cause a high-AG acidosis.

ETHYLENE GLYCOL (See also Chap. 472e) Ingestion of ethylene glycol (commonly used in antifreeze) leads to a metabolic acidosis and severe damage to the CNS, heart, lungs, and kidneys. The increased AG and osmolar gap are attributable to ethylene glycol and its metabolites, oxalic acid, glycolic acid, and other organic acids. Lactic acid production increases secondary to inhibition of the tricarboxylic acid cycle and altered intracellular redox state. Diagnosis is facilitated by recognizing oxalate crystals in the urine, the presence of an osmolar gap

in serum, and a high-AG acidosis. Although use of a Wood's lamp to visualize the fluorescent additive to commercial antifreeze in the urine of patients with ethylene glycol ingestion, this is rarely reproducible. The combination of a high AG and high osmolar gap in a patient suspected of ethylene glycol ingestion should be taken as evidence of ethylene glycol toxicity. Treatment should not be delayed while awaiting measurement of ethylene glycol levels in this setting.

TREATMENT ETHYLENE GLYCOL-INDUCED ACIDOSIS

This includes the prompt institution of a saline or osmotic diuresis, thiamine and pyridoxine supplements, fomepizole, and usually, hemodialysis. The IV administration of the alcohol dehydrogenase inhibitor fomepizole (4-methylpyrazole; 15 mg/kg as a loading dose) is the agent of choice and offers the advantages of a predictable decline in ethylene glycol levels without excessive obtundation as seen during ethyl alcohol infusion. If used, ethanol IV should be infused to achieve a blood level of 22 mmol/L (100 mg/dL). Both fomepizole and ethanol reduce toxicity because they compete with ethylene glycol for metabolism by alcohol dehydrogenase. Hemodialysis is indicated when the arterial pH is <7.3 or the osmolar gap exceeds 20 mOsm/kg.

METHANOL (See also Chap. 472e) The ingestion of methanol (wood alcohol) causes metabolic acidosis, and its metabolites formaldehyde and formic acid cause severe optic nerve and CNS damage. Lactic acid, ketoacids, and other unidentified organic acids may contribute to the acidosis. Due to its low molecular mass (32 Da), an osmolar gap is usually present.

TREATMENT METHANOL-INDUCED ACIDOSIS

This is similar to that for ethylene glycol intoxication, including general supportive measures, fomepizole, and hemodialysis (as above).

PROPYLENE GLYCOL Propylene glycol is the vehicle used in IV administration of diazepam, lorazepam, phenobarbital, nitroglycerine, etomidate, enoximone, and phenytoin. Propylene glycol is generally safe for limited use in these IV preparations, but toxicity has been reported, most often in the setting of the intensive care unit in patients receiving frequent or continuous therapy. This form of high-gap acidosis should be considered in patients with unexplained high-gap acidosis, hyperosmolality, and clinical deterioration. Propylene glycol, like ethylene glycol and methanol, is metabolized by alcohol dehydrogenase. With intoxication by propylene glycol, the first response is to stop the offending infusion. Additionally, fomepizole should also be administered in acidotic patients.

ISOPROPYL ALCOHOL Ingested isopropanol is absorbed rapidly and may be fatal when as little as 150 mL of rubbing alcohol, solvent, or deicer is consumed. A plasma level >400 mg/dL is life-threatening. Isopropyl alcohol is metabolized by alcohol dehydrogenase to acetone. The characteristic features differ from ethylene glycol and methanol in that the parent compound, not the metabolites, causes toxicity, and an AG acidosis is *not* present because acetone is rapidly excreted. Both isopropyl alcohol and acetone increase the osmolar gap, and hypoglycemia is common. Alternative diagnoses should be considered if the patient does not improve significantly within a few hours. Patients with hemodynamic instability with plasma levels above 400 mg/dL should be considered for hemodialysis.

TREATMENT ISOPROPYL ALCOHOL TOXICITY

Isopropanol alcohol toxicity is treated by watchful waiting and supportive therapy, IV fluids, pressors, ventilatory support if needed, and occasionally hemodialysis for prolonged coma, hemodynamic instability, or levels >400 mg/dL.

PYROGLUTAMIC ACID Acetaminophen-induced high-AG metabolic acidosis is uncommon but is being recognized more often in either patients with acetaminophen overdose or malnourished or critically ill patients receiving acetaminophen in typical dosage. 5-Oxoproline accumulation after acetaminophen should be suspected in the setting of an unexplained high-AG acidosis without elevation of the osmolar gap in patients receiving acetaminophen. The first step in treatment is to immediately discontinue the drug. Additionally, sodium bicarbonate IV should be given. Although *N*-acetylcysteine has been suggested, it is not known if it hastens the metabolism of 5-oxoproline by increasing intracellular glutathione concentrations in this setting.

Renal Failure (See also Chap. 335) The hyperchloremic acidosis of moderate renal insufficiency is eventually converted to the high-AG acidosis of advanced renal failure. Poor filtration and reabsorption of organic anions contribute to the pathogenesis. As renal disease progresses, the number of functioning nephrons eventually becomes insufficient to keep pace with net acid production. Uremic acidosis is characterized, therefore, by a reduced rate of NH_4^+ production and excretion. The acid retained in chronic renal disease is buffered by alkaline salts from bone. Despite significant retention of acid (up to 20 mmol/d), the serum $[\text{HCO}_3^-]$ does not decrease further, indicating participation of buffers outside the extracellular compartment. Chronic metabolic acidosis results in significant loss of bone mass due to reduction in bone calcium carbonate. Chronic acidosis also increases urinary calcium excretion, proportional to cumulative acid retention.

TREATMENT RENAL FAILURE

Because of the association of renal failure acidosis with muscle catabolism and bone disease, both uremic acidosis and the hyperchloremic acidosis of renal failure require oral alkali replacement to maintain the $[\text{HCO}_3^-] > 22$ mmol/L. This can be accomplished with relatively modest amounts of alkali (1.0–1.5 mmol/kg body weight per day). Sodium citrate (Shohl's solution) or NaHCO_3 tablets (650-mg tablets contain 7.8 meq) are equally effective alkalinizing salts. Citrate enhances the absorption of aluminum from the gastrointestinal tract and should never be given together with aluminum-containing antacids because of the risk of aluminum intoxication.

NON-ANION GAP METABOLIC ACIDOSES

Alkali can be lost from the gastrointestinal tract from diarrhea or from the kidneys (renal tubular acidosis, RTA). In these disorders (Table 66-5), reciprocal changes in $[\text{Cl}^-]$ and $[\text{HCO}_3^-]$ result in a normal AG. In pure non-AG acidosis, therefore, the increase in $[\text{Cl}^-]$ above the normal value approximates the decrease in $[\text{HCO}_3^-]$. The absence of such a relationship suggests a mixed disturbance.

TREATMENT NON-ANION GAP METABOLIC ACIDOSES

In diarrhea, stools contain a higher $[\text{HCO}_3^-]$ and decomposed HCO_3^- than plasma so that metabolic acidosis develops along with volume depletion. Instead of an acid urine pH (as anticipated with systemic acidosis), urine pH is usually > 6 because metabolic acidosis and hypokalemia increase renal synthesis and excretion of NH_4^+ , thus providing a urinary buffer that increases urine pH. Metabolic acidosis due to gastrointestinal losses with a high urine pH can be differentiated from RTA because urinary NH_4^+ excretion is typically low in RTA and high with diarrhea. Urinary NH_4^+ levels can be estimated by calculating the urine anion gap (UAG): $\text{UAG} = [\text{Na}^+ + \text{K}^+]_u - [\text{Cl}^-]_u$. When $[\text{Cl}^-]_u > [\text{Na}^+ + \text{K}^+]_u$, the UAG is negative by definition. This indicates that the urine ammonium level is appropriately increased, suggesting an extrarenal cause of the acidosis. Conversely, when the UAG is positive, the urine ammonium level is low, suggesting a renal cause of the acidosis.

TABLE 66-5 CAUSES OF NON-ANION GAP ACIDOSIS

- I. Gastrointestinal bicarbonate loss
 - A. Diarrhea
 - B. External pancreatic or small-bowel drainage
 - C. Ureterosigmoidostomy, jejunal loop, ileal loop
 - D. Drugs
 1. Calcium chloride (acidifying agent)
 2. Magnesium sulfate (diarrhea)
 3. Cholestyramine (bile acid diarrhea)
- II. Renal acidosis
 - A. Hypokalemia
 1. Proximal RTA (type 2)
 - Drug-induced: acetazolamide, topiramate
 2. Distal (classic) RTA (type 1)
 - Drug induced: amphotericin B, ifosfamide
 - B. Hyperkalemia
 1. Generalized distal nephron dysfunction (type 4 RTA)
 - a. Mineralocorticoid deficiency
 - b. Mineralocorticoid resistance (PHA I, autosomal dominant)
 - c. Voltage defect (PHA I, autosomal recessive, and PHA II)
 - d. Tubulointerstitial disease
 - C. Normokalemia
 1. Chronic progressive kidney disease
- III. Drug-induced hyperkalemia (with renal insufficiency)
 - A. Potassium-sparing diuretics (amiloride, triamterene, spironolactone, eplerenone)
 - B. Trimethoprim
 - C. Pentamidine
 - D. ACE-Is and ARBs
 - E. Nonsteroidal anti-inflammatory drugs
 - F. Calcineurin inhibitors
- IV. Other
 - A. Acid loads (ammonium chloride, hyperalimentation)
 - B. Loss of potential bicarbonate: ketosis with ketone excretion
 - C. Expansion acidosis (rapid saline administration)
 - D. Hippurate
 - E. Cation exchange resins

Abbreviations: ACE-I, angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker; PHA, pseudohypoaldosteronism; RTA, renal tubular acidosis.

Proximal RTA (type 2 RTA) (Chap. 339) is most often due to generalized proximal tubular dysfunction manifested by glycosuria, generalized aminoaciduria, and phosphaturia (Fanconi syndrome). With a low plasma $[\text{HCO}_3^-]$, the urine pH is acid (pH < 5.5). The fractional excretion of $[\text{HCO}_3^-]$ may exceed 10–15% when the serum $\text{HCO}_3^- > 20$ mmol/L. Because HCO_3^- is not reabsorbed normally in the proximal tubule, therapy with NaHCO_3 will enhance renal potassium wasting and hypokalemia.

The typical findings in acquired or inherited forms of **classic distal RTA (type 1 RTA)** include hypokalemia, non-AG metabolic acidosis, low urinary NH_4^+ excretion (positive UAG, low urine $[\text{NH}_4^+]$), and inappropriately high urine pH (pH > 5.5). Most patients have hypocitraturia and hypercalciuria, so nephrolithiasis, nephrocalcinosis, and bone disease are common. In **generalized distal RTA (type 4 RTA)**, hyperkalemia is disproportionate to the reduction in glomerular filtration rate (GFR) because of coexisting dysfunction of potassium and acid secretion. Urinary ammonium excretion is invariably depressed, and renal function may be compromised, for example, due to diabetic nephropathy, obstructive uropathy, or chronic tubulointerstitial disease.

Hyporeninemic hypoaldosteronism typically causes non-AG metabolic acidosis, most commonly in older adults with diabetes mellitus or tubulointerstitial disease and renal insufficiency. Patients usually

have mild to moderate CKD (GFR, 20–50 mL/min) and acidosis, with elevation in serum $[K^+]$ (5.2–6.0 mmol/L), concurrent hypertension, and congestive heart failure. Both the metabolic acidosis and the hyperkalemia are out of proportion to impairment in GFR. Nonsteroidal anti-inflammatory drugs, trimethoprim, pentamidine, and angiotensin-converting enzyme (ACE) inhibitors can also cause non-AG metabolic acidosis in patients with renal insufficiency (Table 66-5).

METABOLIC ALKALOSIS

Metabolic alkalosis is manifested by an elevated arterial pH, an increase in the serum $[HCO_3^-]$, and an increase in $Paco_2$ as a result of compensatory alveolar hypoventilation (Table 66-1). It is often accompanied by hypochloremia and hypokalemia. The arterial pH establishes the diagnosis, because it is increased in metabolic alkalosis and decreased or normal in respiratory acidosis. Metabolic alkalosis frequently occurs in association with other disorders such as respiratory acidosis or alkalosis or metabolic acidosis.

PATHOGENESIS

Metabolic alkalosis occurs as a result of net gain of $[HCO_3^-]$ or loss of nonvolatile acid (usually HCl by vomiting) from the extracellular fluid. For HCO_3^- to be added to the extracellular fluid, it must be administered exogenously or synthesized endogenously, in part or entirely by the kidneys. Because it is unusual for alkali to be added to the body, the disorder involves a generative stage, in which the loss of acid usually causes alkalosis, and a maintenance stage, in which the kidneys fail to compensate by excreting HCO_3^- .

Maintenance of metabolic alkalosis represents a failure of the kidneys to eliminate HCO_3^- in the usual manner. The kidneys will retain, rather than excrete, the excess alkali and maintain the alkalosis if (1) volume deficiency, chloride deficiency, and K^+ deficiency exist in combination with a reduced GFR; or (2) hypokalemia exists because of autonomous hyperaldosteronism. In the first example, alkalosis is corrected by administration of NaCl and KCl, whereas, in the latter, it may be necessary to repair the alkalosis by pharmacologic or surgical intervention, not with saline administration.

DIFFERENTIAL DIAGNOSIS

To establish the cause of metabolic alkalosis (Table 66-6), it is necessary to assess the status of the extracellular fluid volume (ECFV), the recumbent and upright blood pressure, the serum $[K^+]$, and the renin-aldosterone system. For example, the presence of chronic hypertension and chronic hypokalemia in an alkalotic patient suggests either mineralocorticoid excess or that the hypertensive patient is receiving diuretics. Low plasma renin activity and normal urine $[Na^+]$ and $[Cl^-]$ in a patient who is not taking diuretics indicate a primary mineralocorticoid excess syndrome. The combination of hypokalemia and alkalosis in a normotensive, nonedematous patient can be due to Bartter's or Gitelman's syndrome, magnesium deficiency, vomiting, exogenous alkali, or diuretic ingestion. Determination of urine electrolytes (especially the urine $[Cl^-]$) and screening of the urine for diuretics may be helpful. If the urine is alkaline, with an elevated $[Na^+]$ and $[K^+]$ but low $[Cl^-]$, the diagnosis is usually either vomiting (overt or surreptitious) or alkali ingestion. If the urine is relatively acid and has low concentrations of Na^+ , K^+ , and Cl^- , the most likely possibilities are prior vomiting, the posthypercapnic state, or prior diuretic ingestion. If, on the other hand, neither the urine sodium, potassium, nor chloride concentrations are depressed, magnesium deficiency, Bartter's or Gitelman's syndrome, or current diuretic ingestion should be considered. Bartter's syndrome is distinguished from Gitelman's syndrome because of hypocalciuria and hypomagnesemia in the latter disorder.

Alkali Administration Chronic administration of alkali to individuals with normal renal function rarely causes alkalosis. However, in patients with coexistent hemodynamic disturbances, alkalosis can develop because the normal capacity to excrete HCO_3^- may be exceeded or there may be enhanced reabsorption of HCO_3^- . Such patients include those who receive HCO_3^- (PO or IV), acetate loads

TABLE 66-6 CAUSES OF METABOLIC ALKALOSIS

- I. Exogenous HCO_3^- loads
 - A. Acute alkali administration
 - B. Milk-alkali syndrome
- II. Effective ECFV contraction, normotension, K^+ deficiency, and secondary hyperreninemic hyperaldosteronism
 - A. Gastrointestinal origin
 1. Vomiting
 2. Gastric aspiration
 3. Congenital chloridorrhea
 4. Villous adenoma
 - B. Renal origin
 1. Diuretics
 2. Posthypercapnic state
 3. Hypercalcemia/hypoparathyroidism
 4. Recovery from lactic acidosis or ketoacidosis
 5. Nonreabsorbable anions including penicillin, carbenicillin
 6. Mg^{2+} deficiency
 7. K^+ depletion
 8. Bartter's syndrome (loss of function mutations of transporters and ion channels in TALH)
 9. Gitelman's syndrome (loss of function mutation in Na^+-Cl^- cotransporter in DCT)
- III. ECFV expansion, hypertension, K^+ deficiency, and mineralocorticoid excess
 - A. High renin
 1. Renal artery stenosis
 2. Accelerated hypertension
 3. Renin-secreting tumor
 4. Estrogen therapy
 - B. Low renin
 1. Primary aldosteronism
 - a. Adenoma
 - b. Hyperplasia
 - c. Carcinoma
 2. Adrenal enzyme defects
 - a. 11β -Hydroxylase deficiency
 - b. 17α -Hydroxylase deficiency
 3. Cushing's syndrome or disease
 4. Other
 - a. Licorice
 - b. Carbenoxolone
 - c. Chewer's tobacco
- IV. Gain-of-function mutation of renal sodium channel with ECFV expansion, hypertension, K^+ deficiency, and hyporeninemic-hypoaldosteronism
 - A. Liddle's syndrome

Abbreviations: DCT, distal convoluted tubule; ECFV, extracellular fluid volume; TALH, thick ascending limb of Henle's loop.

(parenteral hyperalimentation solutions), citrate loads (transfusions), or antacids plus cation-exchange resins (aluminum hydroxide and sodium polystyrene sulfonate). Nursing home patients receiving tube feedings have a higher incidence of metabolic alkalosis than nursing home patients receiving oral feedings.

METABOLIC ALKALOSIS ASSOCIATED WITH ECFV CONTRACTION, K^+ DEPLETION, AND SECONDARY HYPERRENINEMIC HYPERALDOSTERONISM

Gastrointestinal Origin Gastrointestinal loss of H^+ from vomiting or gastric aspiration results in retention of HCO_3^- . During active vomiting, the filtered load of bicarbonate is acutely increased to exceed the reabsorptive capacity of the proximal tubule for HCO_3^- so that the urine becomes alkaline and high in potassium. When vomiting ceases, the persistence of volume, potassium, and chloride depletion causes

maintenance of the alkalosis because of an enhanced capacity of the nephron to reabsorb HCO_3^- . Correction of the contracted ECFV with NaCl and repair of K^+ deficits corrects the acid-base disorder by restoring the ability of the kidney to excrete the excess bicarbonate.

Renal Origin • DIURETICS (See also Chap. 279) Drugs that induce diuresis, such as thiazides and loop diuretics (furosemide, bumetanide, torsemide, and ethacrynic acid), acutely diminish the ECFV without altering the total body bicarbonate content. The serum $[\text{HCO}_3^-]$ increases because the reduced ECFV “contracts” the $[\text{HCO}_3^-]$ in the plasma (contraction alkalosis). The chronic administration of diuretics tends to generate an alkalosis by increasing distal salt delivery, so that K^+ and H^+ secretion are stimulated. The alkalosis is maintained by persistence of the contraction of the ECFV, secondary hyperaldosteronism, K^+ deficiency, and the direct effect of the diuretic (as long as diuretic administration continues). Repair of the alkalosis is achieved by providing isotonic saline to correct the ECFV deficit.

SOLUTE LOSING DISORDERS: BARTTER'S SYNDROME AND GITELMAN'S SYNDROME See Chap. 339.

NONREABSORBABLE ANIONS AND MAGNESIUM DEFICIENCY Administration of large quantities of nonreabsorbable anions, such as penicillin or carbenicillin, can enhance distal acidification and K^+ secretion by increasing the transepithelial potential difference. Mg^{2+} deficiency results in hypokalemic alkalosis by enhancing distal acidification through stimulation of renin and hence aldosterone secretion.

POTASSIUM DEPLETION Chronic K^+ depletion may cause metabolic alkalosis by increasing urinary acid excretion. Both NH_4^+ production and absorption are enhanced and HCO_3^- reabsorption is stimulated. Chronic K^+ deficiency upregulates the renal H^+ , K^+ -ATPase to increase K^+ absorption at the expense of enhanced H^+ secretion. Alkalosis associated with severe K^+ depletion is resistant to salt administration, but repair of the K^+ deficiency corrects the alkalosis.

AFTER TREATMENT OF LACTIC ACIDOSIS OR KETOACIDOSIS When an underlying stimulus for the generation of lactic acid or ketoacid is removed rapidly, as with repair of circulatory insufficiency or with insulin therapy, the lactate or ketones are metabolized to yield an equivalent amount of HCO_3^- . Other sources of new HCO_3^- are additive with the original amount generated by organic anion metabolism to create a surfeit of HCO_3^- . Such sources include (1) new HCO_3^- added to the blood by the kidneys as a result of enhanced acid excretion during the preexisting period of acidosis, and (2) alkali therapy during the treatment phase of the acidosis. Acidosis-induced contraction of the ECFV and K^+ deficiency act to sustain the alkalosis.

POSTHYPERCAPNIA Prolonged CO_2 retention with chronic respiratory acidosis enhances renal HCO_3^- absorption and the generation of new HCO_3^- (increased net acid excretion). Metabolic alkalosis results from the effect of the persistently elevated $[\text{HCO}_3^-]$ when the elevated Paco_2 is abruptly returned toward normal.

METABOLIC ALKALOSIS ASSOCIATED WITH ECFV EXPANSION, HYPERTENSION, AND HYPERALDOSTERONISM

Increased aldosterone levels may be the result of autonomous primary adrenal overproduction or of secondary aldosterone release due to renal overproduction of renin. Mineralocorticoid excess increases net acid excretion and may result in metabolic alkalosis, which may be worsened by associated K^+ deficiency. ECFV expansion from salt retention causes hypertension. The kaliuresis persists because of mineralocorticoid excess and distal Na^+ absorption causing enhanced K^+ excretion, continued K^+ depletion with polydipsia, inability to concentrate the urine, and polyuria.

Liddle's syndrome (Chap. 339) results from increased activity of the collecting duct Na^+ channel (ENaC) and is a rare monogenic form of hypertension due to volume expansion manifested as hypokalemic alkalosis and normal aldosterone levels.

Symptoms With metabolic alkalosis, changes in CNS and peripheral nervous system function are similar to those of hypocalcemia (Chap. 423); symptoms include mental confusion; obtundation; and

a predisposition to seizures, paresthesia, muscular cramping, tetany, aggravation of arrhythmias, and hypoxemia in chronic obstructive pulmonary disease. Related electrolyte abnormalities include hypokalemia and hypophosphatemia.

TREATMENT METABOLIC ALKALOSIS

This is primarily directed at correcting the underlying stimulus for HCO_3^- generation. If primary aldosteronism, renal artery stenosis, or Cushing's syndrome is present, correction of the underlying cause will reverse the alkalosis. $[\text{H}^+]$ loss by the stomach or kidneys can be mitigated by the use of proton pump inhibitors or the discontinuation of diuretics. The second aspect of treatment is to remove the factors that sustain the inappropriate increase in HCO_3^- reabsorption, such as ECFV contraction or K^+ deficiency. K^+ deficits should always be repaired. Isotonic saline is usually sufficient to reverse the alkalosis if ECFV contraction is present.

If associated conditions preclude infusion of saline, renal HCO_3^- loss can be accelerated by administration of acetazolamide, a carbonic anhydrase inhibitor, which is usually effective in patients with adequate renal function but can worsen K^+ losses. Dilute hydrochloric acid (0.1 N HCl) is also effective but can cause hemolysis, and must be delivered slowly in a central vein.

RESPIRATORY ACIDOSIS

Respiratory acidosis can be due to severe pulmonary disease, respiratory muscle fatigue, or abnormalities in ventilatory control and is recognized by an increase in Paco_2 and decrease in pH (Table 66-7). In acute respiratory acidosis, there is an immediate compensatory elevation (due to cellular buffering mechanisms) in $[\text{HCO}_3^-]$, which increases 1 mmol/L for every 10-mmHg increase in Paco_2 . In chronic respiratory acidosis (>24 h), renal adaptation increases the $[\text{HCO}_3^-]$ by 4 mmol/L for every 10-mmHg increase in Paco_2 . The serum HCO_3^- usually does not increase above 38 mmol/L.

The clinical features vary according to the severity and duration of the respiratory acidosis, the underlying disease, and whether there is accompanying hypoxemia. A rapid increase in Paco_2 may cause anxiety, dyspnea, confusion, psychosis, and hallucinations and may progress to coma. Lesser degrees of dysfunction in chronic hypercapnia include sleep disturbances; loss of memory; daytime somnolence; personality changes; impairment of coordination; and motor disturbances such as tremor, myoclonic jerks, and asterixis. Headaches and other signs that mimic raised intracranial pressure, such as papilledema, abnormal reflexes, and focal muscle weakness, are due to vasoconstriction secondary to loss of the vasodilator effects of CO_2 .

Depression of the respiratory center by a variety of drugs, injury, or disease can produce respiratory acidosis. This may occur acutely with general anesthetics, sedatives, and head trauma or chronically with sedatives, alcohol, intracranial tumors, and the syndromes of sleep-disordered breathing including the primary alveolar and obesity-hypoventilation syndromes (Chaps. 318 and 319). Abnormalities or disease in the motor neurons, neuromuscular junction, and skeletal muscle can cause hypoventilation via respiratory muscle fatigue. Mechanical ventilation, when not properly adjusted and supervised, may result in respiratory acidosis, particularly if CO_2 production suddenly rises (because of fever, agitation, sepsis, or overfeeding) or alveolar ventilation falls because of worsening pulmonary function. High levels of positive end-expiratory pressure in the presence of reduced cardiac output may cause hypercapnia as a result of large increases in alveolar dead space (Chap. 306e). Permissive hypercapnia is being used with increasing frequency because of studies suggesting lower mortality rates than with conventional mechanical ventilation, especially with severe CNS or heart disease. The respiratory acidosis associated with permissive hypercapnia may require administration of NaHCO_3 to increase the arterial pH to 7.25, but overcorrection of the acidemia may be deleterious.

Acute hypercapnia follows sudden occlusion of the upper airway or generalized bronchospasm as in severe asthma, anaphylaxis,

TABLE 66-7 RESPIRATORY ACID-BASE DISORDERS

- I. Alkalosis
 - A. Central nervous system stimulation
 - 1. Pain
 - 2. Anxiety, psychosis
 - 3. Fever
 - 4. Cerebrovascular accident
 - 5. Meningitis, encephalitis
 - 6. Tumor
 - 7. Trauma
 - B. Hypoxemia or tissue hypoxia
 - 1. High altitude
 - 2. Pneumonia, pulmonary edema
 - 3. Aspiration
 - 4. Severe anemia
 - C. Drugs or hormones
 - 1. Pregnancy, progesterone
 - 2. Salicylates
 - 3. Cardiac failure
 - D. Stimulation of chest receptors
 - 1. Hemothorax
 - 2. Flail chest
 - 3. Cardiac failure
 - 4. Pulmonary embolism
 - E. Miscellaneous
 - 1. Septicemia
 - 2. Hepatic failure
 - 3. Mechanical hyperventilation
 - 4. Heat exposure
 - 5. Recovery from metabolic acidosis
- II. Acidosis
 - A. Central
 - 1. Drugs (anesthetics, morphine, sedatives)
 - 2. Stroke
 - 3. Infection
 - B. Airway
 - 1. Obstruction
 - 2. Asthma
 - C. Parenchyma
 - 1. Emphysema
 - 2. Pneumoconiosis
 - 3. Bronchitis
 - 4. Adult respiratory distress syndrome
 - 5. Barotrauma
 - D. Neuromuscular
 - 1. Poliomyelitis
 - 2. Kyphoscoliosis
 - 3. Myasthenia
 - 4. Muscular dystrophies
 - E. Miscellaneous
 - 1. Obesity
 - 2. Hypoventilation
 - 3. Permissive hypercapnia

inhalational burn, or toxin injury. Chronic hypercapnia and respiratory acidosis occur in end-stage obstructive lung disease. Restrictive disorders involving both the chest wall and the lungs can cause respiratory acidosis because the high metabolic cost of respiration causes ventilatory muscle fatigue. Advanced stages of intrapulmonary and extrapulmonary restrictive defects present as chronic respiratory acidosis.

The diagnosis of respiratory acidosis requires the measurement of P_{aCO_2} and arterial pH. A detailed history and physical examination often indicate the cause. Pulmonary function studies (Chap. 306e), including spirometry, diffusion capacity for carbon monoxide, lung volumes, and arterial P_{aCO_2} and O_2 saturation, usually make it possible to determine if respiratory acidosis is secondary to lung disease. The workup for nonpulmonary causes should include a detailed drug history, measurement of hematocrit, and assessment of upper airway, chest wall, pleura, and neuromuscular function.

TREATMENT RESPIRATORY ACIDOSIS

The management of respiratory acidosis depends on its severity and rate of onset. Acute respiratory acidosis can be life-threatening, and measures to reverse the underlying cause should be undertaken simultaneously with restoration of adequate alveolar ventilation. This may necessitate tracheal intubation and assisted mechanical ventilation. Oxygen administration should be titrated carefully in patients with severe obstructive pulmonary disease and chronic CO_2 retention who are breathing spontaneously (Chap. 314). When oxygen is used injudiciously, these patients may experience progression of the respiratory acidosis. Aggressive and rapid correction of hypercapnia should be avoided, because the falling P_{aCO_2} may provoke the same complications noted with acute respiratory alkalosis (i.e., cardiac arrhythmias, reduced cerebral perfusion, and seizures). The P_{aCO_2} should be lowered gradually in chronic respiratory acidosis, aiming to restore the P_{aCO_2} to baseline levels and to provide sufficient Cl^- and K^+ to enhance the renal excretion of HCO_3^- .

Chronic respiratory acidosis is frequently difficult to correct, but measures aimed at improving lung function (Chap. 314) can help some patients and forestall further deterioration in most.

RESPIRATORY ALKALOSIS

Alveolar hyperventilation decreases P_{aCO_2} and increases the HCO_3^-/P_{aCO_2} ratio, thus increasing pH (Table 66-7). Nonbicarbonate cellular buffers respond by consuming HCO_3^- . Hypocapnia develops when a sufficiently strong ventilatory stimulus causes CO_2 output in the lungs to exceed its metabolic production by tissues. Plasma pH and $[HCO_3^-]$ appear to vary proportionately with P_{aCO_2} over a range from 40–15 mmHg. The relationship between arterial $[H^+]$ concentration and P_{aCO_2} is ~ 0.7 mmol/L per mmHg (or 0.01 pH unit/mmHg), and that for plasma $[HCO_3^-]$ is 0.2 mmol/L per mmHg. Hypocapnia sustained for >2 –6 h is further compensated by a decrease in renal ammonium and titratable acid excretion and a reduction in filtered HCO_3^- reabsorption. Full renal adaptation to respiratory alkalosis may take several days and requires normal volume status and renal function. The kidneys appear to respond directly to the lowered P_{aCO_2} rather than to alkalosis per se. In chronic respiratory alkalosis a 1-mmHg decrease in P_{aCO_2} causes a 0.4- to 0.5-mmol/L drop in $[HCO_3^-]$ and a 0.3-mmol/L decrease (or 0.003 increase in pH) in $[H^+]$.

The effects of respiratory alkalosis vary according to duration and severity but are primarily those of the underlying disease. Reduced cerebral blood flow as a consequence of a rapid decline in P_{aCO_2} may cause dizziness, mental confusion, and seizures, even in the absence of hypoxemia. The cardiovascular effects of acute hypocapnia in the conscious human are generally minimal, but in the anesthetized or mechanically ventilated patient, cardiac output and blood pressure may fall because of the depressant effects of anesthesia and positive-pressure ventilation on heart rate, systemic resistance, and venous return. Cardiac arrhythmias may occur in patients with heart disease as a result of changes in oxygen unloading by blood from a left shift in the hemoglobin-oxygen dissociation curve (Bohr effect). Acute respiratory alkalosis causes intracellular shifts of Na^+ , K^+ , and PO_4^{2-} and reduces free $[Ca^{2+}]$ by increasing the protein-bound fraction. Hypocapnia-induced hypokalemia is usually minor.

Chronic respiratory alkalosis is the most common acid-base disturbance in critically ill patients and, when severe, portends a poor prognosis. Many cardiopulmonary disorders manifest respiratory alkalosis

in their early to intermediate stages, and the finding of normocapnia and hypoxemia in a patient with hyperventilation may herald the onset of rapid respiratory failure and should prompt an assessment to determine if the patient is becoming fatigued. Respiratory alkalosis is common during mechanical ventilation.

The hyperventilation syndrome may be disabling. Paresthesia; circumoral numbness; chest wall tightness or pain; dizziness; inability to take an adequate breath; and, rarely, tetany may be sufficiently stressful to perpetuate the disorder. Arterial blood-gas analysis demonstrates an acute or chronic respiratory alkalosis, often with hypocapnia in the range of 15–30 mmHg and no hypoxemia. CNS diseases or injury can produce several patterns of hyperventilation and sustained P_{aCO_2} levels of 20–30 mmHg. Hyperthyroidism, high caloric loads, and exercise raise the basal metabolic rate, but ventilation usually rises in proportion so that arterial blood gases are unchanged and respiratory alkalosis does not develop. Salicylates are the most common cause of drug-induced respiratory alkalosis as a result of direct stimulation of the medullary chemoreceptor (Chap. 472e). The methylxanthines, theophylline, and aminophylline stimulate ventilation and increase the ventilatory response to CO_2 . Progesterone increases ventilation and lowers arterial P_{aCO_2} by as much as 5–10 mmHg. Therefore, chronic respiratory alkalosis is a common feature of pregnancy. Respiratory alkalosis is also prominent in liver failure, and the severity correlates with the degree of hepatic insufficiency. Respiratory alkalosis is often an early finding in gram-negative septicemia, before fever, hypoxemia, or hypotension develops.

The diagnosis of respiratory alkalosis depends on measurement of arterial pH and P_{aCO_2} . The plasma $[K^+]$ is often reduced and the $[Cl^-]$ increased. In the acute phase, respiratory alkalosis is not associated with increased renal HCO_3^- excretion, but within hours net acid excretion is reduced. In general, the HCO_3^- concentration falls by 2.0 mmol/L for each 10-mmHg decrease in P_{aCO_2} . Chronic hypocapnia reduces the serum $[HCO_3^-]$ by 4.0 mmol/L for each 10-mmHg decrease in P_{aCO_2} . It is unusual to observe a plasma $HCO_3^- < 12$ mmol/L as a result of a pure respiratory alkalosis.

When a diagnosis of respiratory alkalosis is made, its cause should be investigated. The diagnosis of hyperventilation syndrome is made by exclusion. In difficult cases, it may be important to rule out other conditions such as pulmonary embolism, coronary artery disease, and hyperthyroidism.

TREATMENT RESPIRATORY ALKALOSIS

The management of respiratory alkalosis is directed toward alleviation of the underlying disorder. If respiratory alkalosis complicates ventilator management, changes in dead space, tidal volume, and frequency can minimize the hypocapnia. Patients with the hyperventilation syndrome may benefit from reassurance, rebreathing from a paper bag during symptomatic attacks, and attention to underlying psychological stress. Antidepressants and sedatives are not recommended. β -Adrenergic blockers may ameliorate peripheral manifestations of the hyperadrenergic state.

SECTION 8 ALTERATIONS IN SEXUAL FUNCTION AND REPRODUCTION

67 Sexual Dysfunction

Kevin T. McVary

Male sexual dysfunction affects 10–25% of middle-aged and elderly men, and female sexual dysfunction occurs with similar frequency. Demographic changes, the popularity of newer treatments, and greater awareness of sexual dysfunction by patients and society have led to increased diagnosis and associated health care expenditures for the management of this common disorder. Because many patients are reluctant to initiate discussion of their sex lives, physicians should address this topic directly to elicit a history of sexual dysfunction.

MALE SEXUAL DYSFUNCTION

PHYSIOLOGY OF MALE SEXUAL RESPONSE

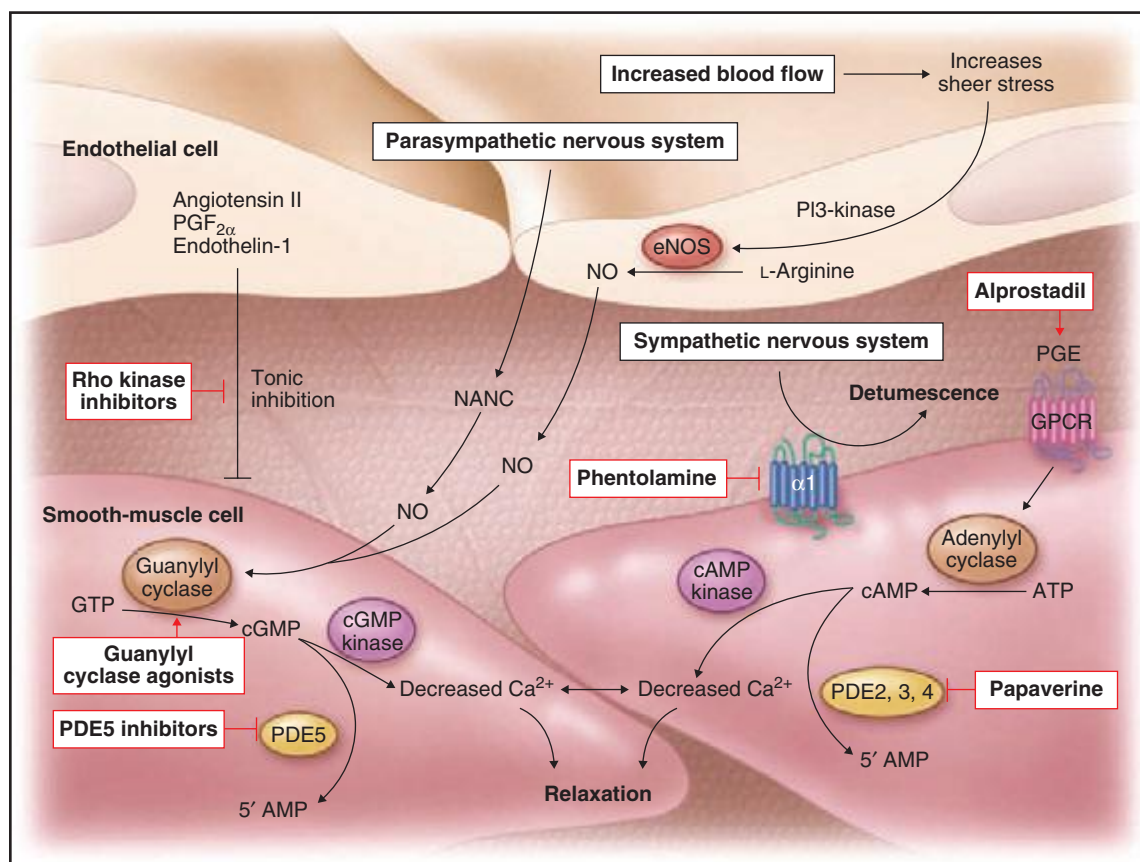
Normal male sexual function requires (1) an intact libido, (2) the ability to achieve and maintain penile erection, (3) ejaculation, and (4) detumescence. *Libido* refers to sexual desire and is influenced by a variety of visual, olfactory, tactile, auditory, imaginative, and hormonal stimuli. Sex steroids, particularly testosterone, act to increase libido. Libido can be diminished by hormonal or psychiatric disorders and by medications.

Penile tumescence leading to erection depends on an increased flow of blood into the lacunar network accompanied by complete relaxation of the arteries and corporal smooth muscle. The microarchitecture of the corpora is composed of a mass of smooth muscle (trabecula) that contains a network of endothelial-lined vessels (lacunar spaces). Subsequent compression of the trabecular smooth muscle against the fibroelastic tunica albuginea causes a passive closure of the emissary veins and accumulation of blood in the corpora. In the presence of a

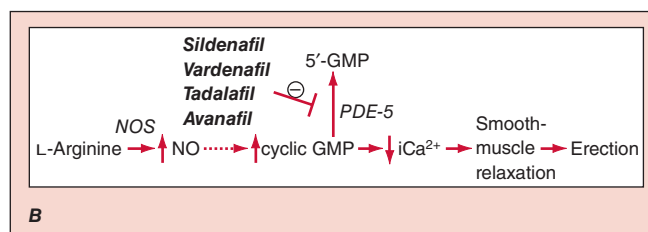
full erection and a competent valve mechanism, the corpora become noncompressible cylinders from which blood does not escape.

The central nervous system (CNS) exerts an important influence by either stimulating or antagonizing spinal pathways that mediate erectile function and ejaculation. The erectile response is mediated by a combination of central (psychogenic) innervation and peripheral (reflexogenic) innervation. Sensory nerves that originate from receptors in the penile skin and glans converge to form the dorsal nerve of the penis, which travels to the S2–S4 dorsal root ganglia via the pudendal nerve. Parasympathetic nerve fibers to the penis arise from neurons in the intermediolateral columns of the S2–S4 sacral spinal segments. Sympathetic innervation originates from the T11 to the L2 spinal segments and descends through the hypogastric plexus.

Neural input to smooth-muscle tone is crucial to the initiation and maintenance of an erection. There is also an intricate interaction between the corporal smooth-muscle cell and its overlying endothelial cell lining (Fig. 67-1). Nitric oxide, which induces vascular relaxation, promotes erection and is opposed by endothelin 1 (ET-1) and Rho kinase, which mediate vascular contraction. Nitric oxide is synthesized from l-arginine by nitric oxide synthase and is released from the nonadrenergic, noncholinergic (NANC) autonomic nerve supply to act postjunctionally on smooth-muscle cells. Nitric oxide increases the production of cyclic 3',5'-guanosine monophosphate (cyclic GMP), which induces relaxation of smooth muscle (Fig. 67-2). Cyclic GMP is gradually broken down by phosphodiesterase type 5 (PDE-5). Inhibitors of PDE-5, such as the oral medications sildenafil, vardenafil, and tadalafil, maintain erections by reducing the breakdown of cyclic GMP. However, if nitric oxide is not produced at some level, PDE-5 inhibitors are ineffective, as these drugs facilitate, but do not initiate, the initial enzyme cascade. In addition to nitric oxide, vasoactive prostaglandins (PGE_1 , $PGF_{2\alpha}$) are synthesized within the cavernosal tissue and increase cyclic AMP levels, also leading to relaxation of cavernosal smooth-muscle cells.



A



B

FIGURE 67-1 Pathways that regulate penile smooth-muscle relaxation and erection. **A.** Outflow from the parasympathetic nervous system leads to relaxation of the cavernous sinusoids in two ways, both of which increase the concentration of nitric oxide (NO) in smooth-muscle cells. First, NO is the neurotransmitter in nonadrenergic, noncholinergic (NANC) fibers; second, stimulation of endothelial nitric oxide synthase (eNOS) through cholinergic output causes increased production of NO. The NO produced in the endothelium then diffuses into the smooth-muscle cells and decreases its intracellular calcium concentration through a pathway mediated by cyclic guanosine monophosphate (cGMP), leading to relaxation. A separate mechanism that decreases the intracellular calcium level is mediated by cyclic adenosine monophosphate (cAMP). With increased cavernosal blood flow, as well as increased levels of vascular endothelial growth factor (VEGF), the endothelial release of NO is further sustained through the phosphatidylinositol 3 (PI3) kinase pathway. Active treatments (red boxes) include drugs that affect the cGMP pathway (phosphodiesterase [PDE] type 5 inhibitors and guanylyl cyclase agonists), the cAMP pathway (alprostadil), or both pathways (papaverine), along with neural-tone mediators (phentolamine and Rho kinase inhibitors). Agents that are being developed include guanylyl cyclase agonists (to bypass the need for endogenous NO) and Rho kinase inhibitors (to inhibit tonic contraction of smooth-muscle cells mediated through endothelin). α_1 , α -adrenergic receptor; GPCR, G-protein-coupled receptor, GTP, guanosine triphosphate; PGE, prostaglandin E; PGF, prostaglandin F. **B.** Biochemical pathways of NO synthesis and action. Sildenafil, vardenafil, and tadalafil enhance erectile function by inhibiting phosphodiesterase type 5 (PDE-5), thereby maintaining high levels of cyclic 3',5'-guanosine monophosphate (cyclic GMP). iCa^{2+} , intracellular calcium; NOS, nitric oxide synthase. (Part A from K McVary: *N Engl J Med* 357:2472, 2007; with permission.)

Ejaculation is stimulated by the sympathetic nervous system; this results in contraction of the epididymis, vas deferens, seminal vesicles, and prostate, causing seminal fluid to enter the urethra. Seminal fluid emission is followed by rhythmic contractions of the bulbocavernosus and ischiocavernosus muscles, leading to ejaculation. *Premature ejaculation* usually is related to anxiety or a learned behavior and is amenable to behavioral therapy or treatment with medications such as selective serotonin reuptake inhibitors (SSRIs). *Retrograde ejaculation* results when the internal urethral sphincter does not close; it may occur in men with diabetes or after surgery involving the bladder neck.

Detumescence is mediated by norepinephrine from the sympathetic nerves, endothelin from the vascular surface, and smooth-muscle contraction induced by postsynaptic α -adrenergic receptors and activation of Rho kinase. These events increase venous outflow and restore the flaccid state. Venous leak can cause premature detumescence and is caused by insufficient relaxation of the corporal smooth muscle rather than a specific anatomic defect. *Priapism* refers to a persistent and painful erection and may be associated with sickle cell anemia, hypercoagulable states, spinal cord injury, or injection of vasodilator agents into the penis.

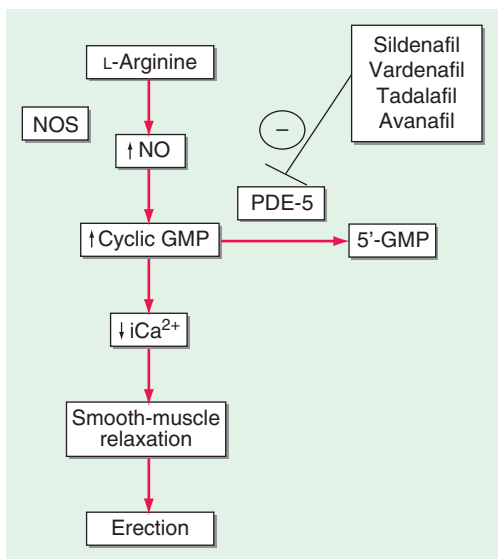


FIGURE 67-2 Biochemical pathways modified by phosphodiesterase type 5 (PDE-5) inhibitors. Sildenafil, vardenafil, tadalafil and avanafil enhance erectile function by inhibiting PDE-5, thereby maintaining high levels of cyclic 3',5'-guanosine monophosphate (cyclic GMP). iCa^{2+} , intracellular calcium; NO, nitric oxide; NOS, nitric oxide synthase.

ERECTILE DYSFUNCTION

Epidemiology Erectile dysfunction (ED) is not considered a normal part of the aging process. Nonetheless, it is associated with certain physiologic and psychological changes related to age. In the Massachusetts Male Aging Study (MMAS), a community-based survey of men age 40–70, 52% of responders reported some degree of ED. Complete ED occurred in 10% of respondents, moderate ED in 25%, and minimal ED in 17%. The incidence of moderate or severe ED more than doubled between the ages of 40 and 70. In the National Health and Social Life Survey (NHSLs), which included a sample of men and women age 18–59, 10% of men reported being unable to maintain an erection (corresponding to the proportion of men in the MMAS reporting severe ED). Incidence was highest among men in the age group 50–59 (21%) and men who were poor (14%), divorced (14%), and less educated (13%).

The incidence of ED is also higher among men with certain medical disorders, such as diabetes mellitus, obesity, lower urinary tract symptoms secondary to benign prostatic hyperplasia (BPH), heart disease, hypertension, decreased high-density lipoprotein (HDL) levels, and diseases associated with general systemic inflammation (e.g., rheumatoid arthritis). Cardiovascular disease and ED share etiologies as well as pathophysiology (e.g., endothelial dysfunction), and the degree of ED appears to correlate with the severity of cardiovascular disease. Consequently, ED represents a “sentinel symptom” in patients with occult cardiovascular and peripheral vascular disease.

Smoking is also a significant risk factor in the development of ED. Medications used in treating diabetes or cardiovascular disease are additional risk factors (see below). There is a higher incidence of ED among men who have undergone radiation or surgery for prostate cancer and in those with a lower spinal cord injury. Psychological causes of ED include depression, anger, stress from unemployment, and other stress-related causes.

Pathophysiology ED may result from three basic mechanisms: (1) failure to initiate (psychogenic, endocrinologic, or neurogenic), (2) failure to fill (arteriogenic), and (3) failure to store adequate blood volume within the lacunar network (venoocclusive dysfunction). These categories are not mutually exclusive, and multiple factors contribute to ED in many patients. For example, diminished filling pressure can lead secondarily to venous leak. Psychogenic factors frequently coexist with other etiologic factors and should be considered in all cases.

Diabetic, atherosclerotic, and drug-related causes account for >80% of cases of ED in older men.

VASCULOGENIC The most common organic cause of ED is a disturbance of blood flow to and from the penis. Atherosclerotic or traumatic arterial disease can decrease flow to the lacunar spaces, resulting in decreased rigidity and an increased time to full erection. Excessive outflow through the veins despite adequate inflow also may contribute to ED. Structural alterations to the fibroelastic components of the corpora may cause a loss of compliance and inability to compress the tunical veins. This condition may result from aging, increased cross-linking of collagen fibers induced by nonenzymatic glycosylation, hypoxemia, or altered synthesis of collagen associated with hypercholesterolemia.

NEUROGENIC Disorders that affect the sacral spinal cord or the autonomic fibers to the penis preclude nervous system relaxation of penile smooth muscle, thus leading to ED. In patients with spinal cord injury, the degree of ED depends on the completeness and level of the lesion. Patients with incomplete lesions or injuries to the upper part of the spinal cord are more likely to retain erectile capabilities than are those with complete lesions or injuries to the lower part. Although 75% of patients with spinal cord injuries have some erectile capability, only 25% have erections sufficient for penetration. Other neurologic disorders commonly associated with ED include multiple sclerosis and peripheral neuropathy. The latter is often due to either diabetes or alcoholism. Pelvic surgery may cause ED through disruption of the autonomic nerve supply.

ENDOCRINOLOGIC Androgens increase libido, but their exact role in erectile function is unclear. Individuals with castrate levels of testosterone can achieve erections from visual or sexual stimuli. Nonetheless, normal levels of testosterone appear to be important for erectile function, particularly in older males. Androgen replacement therapy can improve depressed erectile function when it is secondary to hypogonadism; however, it is not useful for ED when endogenous testosterone levels are normal. Increased prolactin may decrease libido by suppressing gonadotropin-releasing hormone (GnRH), and it also leads to decreased testosterone levels. Treatment of hyperprolactinemia with dopamine agonists can restore libido and testosterone.

DIABETIC ED occurs in 35–75% of men with diabetes mellitus. Pathologic mechanisms are related primarily to diabetes-associated vascular and neurologic complications. Diabetic macrovascular complications are related mainly to age, whereas microvascular complications correlate with the duration of diabetes and the degree of glycemic control (Chap. 417). Individuals with diabetes also have reduced amounts of nitric oxide synthase in both endothelial and neural tissues.

PSYCHOGENIC Two mechanisms contribute to the inhibition of erections in psychogenic ED. First, psychogenic stimuli to the sacral cord may inhibit reflexogenic responses, thereby blocking activation of vasodilator outflow to the penis. Second, excess sympathetic stimulation in an anxious man may increase penile smooth-muscle tone. The most common causes of psychogenic ED are performance anxiety, depression, relationship conflict, loss of attraction, sexual inhibition, conflicts over sexual preference, sexual abuse in childhood, and fear of pregnancy or sexually transmitted disease. Almost all patients with ED, even when it has a clear-cut organic basis, develop a psychogenic component as a reaction to ED.

MEDICATION-RELATED Medication-induced ED (Table 67-1) is estimated to occur in 25% of men seen in general medical outpatient clinics. The adverse effects related to drug therapy are additive, especially in older men. In addition to the drug itself, the disease being treated is likely to contribute to sexual dysfunction. Among the antihypertensive agents, the thiazide diuretics and beta blockers have been implicated most frequently. Calcium channel blockers and angiotensin converting-enzyme inhibitors are cited less frequently. These drugs may act directly at the corporal level (e.g., calcium channel blockers) or indirectly by reducing pelvic blood pressure, which is important in the development of penile rigidity. α -Adrenergic blockers are less likely to

TABLE 67-1 DRUGS ASSOCIATED WITH ERECTILE DYSFUNCTION

Classification	Drugs
Diuretics	Thiazides
	Spironolactone
Antihypertensives	Calcium channel blockers
	Methyldopa
	Clonidine
	Reserpine
	Beta blockers
	Guanethidine
Cardiac/antihyperlipidemics	Digoxin
	Gemfibrozil
	Clofibrate
Antidepressants	Selective serotonin reuptake inhibitors
	Tricyclic antidepressants
	Lithium
	Monoamine oxidase inhibitors
Tranquilizers	Butyrophenones
	Phenothiazines
H ₂ antagonists	Ranitidine
	Cimetidine
Hormones	Progesterone
	Estrogens
	Corticosteroids
	GnRH agonists
	5 α -Reductase inhibitors
	Cyproterone acetate
Cytotoxic agents	Cyclophosphamide
	Methotrexate
	Roferon-A
Anticholinergics	Disopyramide
	Anticonvulsants
Recreational	Ethanol
	Cocaine
	Marijuana

Abbreviation: GnRH, gonadotropin-releasing hormone.

cause ED. Estrogens, GnRH agonists, H₂ antagonists, and spironolactone cause ED by suppressing gonadotropin production or by blocking androgen action. Antidepressant and antipsychotic agents—particularly neuroleptics, tricyclics, and SSRIs—are associated with erectile, ejaculatory, orgasmic, and sexual desire difficulties.

If there is a strong association between the institution of a drug and the onset of ED, alternative medications should be considered. Otherwise, it is often practical to treat the ED without attempting multiple changes in medications, as it may be difficult to establish a causal role for a drug.

APPROACH TO THE PATIENT: Erectile Dysfunction

A good physician-patient relationship helps unravel the possible causes of ED, many of which require discussion of personal and sometimes embarrassing topics. For this reason, a primary care provider is often ideally suited to initiate the evaluation. However, a significant percentage of men experience ED and remain undiagnosed unless specifically questioned about this issue. By far the two most common reasons for underreporting of ED are patient embarrassment and perceptions of physicians' inattention to the disease. Once the topic is initiated by the physician, patients are more willing to discuss their potency issues. A complete medical and sexual

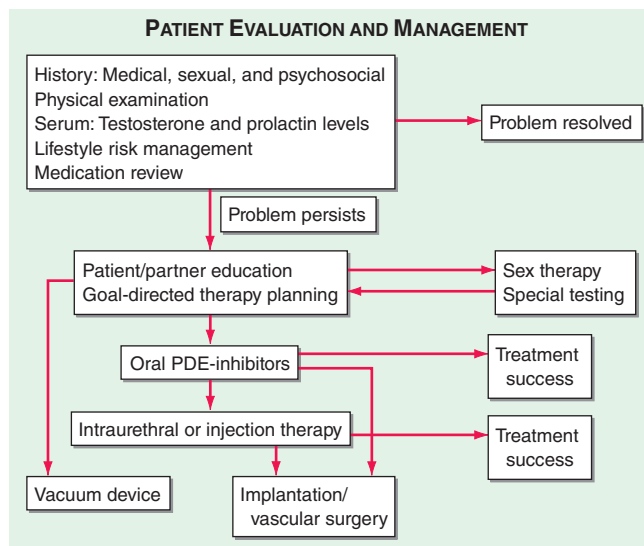


FIGURE 67-3 Algorithm for the evaluation and management of patients with erectile dysfunction. PDE, phosphodiesterase.

history should be taken in an effort to assess whether the cause of ED is organic, psychogenic, or multifactorial (Fig. 67-3).

Both the patient and his sexual partner should be interviewed regarding sexual history. ED should be distinguished from other sexual problems, such as premature ejaculation. Lifestyle factors such as sexual orientation, the patient's distress from ED, performance anxiety, and details of sexual techniques should be addressed. Standardized questionnaires are available to assess ED, including the International Index of Erectile Function (IIEF) and the more easily administered Sexual Health Inventory for Men (SHIM), a validated abridged version of the IIEF.

The initial evaluation of ED begins with a review of the patient's medical, surgical, sexual, and psychosocial histories. The history should note whether the patient has experienced pelvic trauma, surgery, or radiation. In light of the increasing recognition of the relationship between lower urinary tract symptoms and ED, it is advisable to evaluate for the presence of symptoms of bladder outlet obstruction. Questions should focus on the onset of symptoms, the presence and duration of partial erections, and the progression of ED. A history of nocturnal or early morning erections is useful for distinguishing physiologic ED from psychogenic ED. Nocturnal erections occur during rapid eye movement (REM) sleep and require intact neurologic and circulatory systems. Organic causes of ED generally are characterized by a gradual and persistent change in rigidity or the inability to sustain nocturnal, coital, or self-stimulated erections. The patient should be questioned about the presence of penile curvature or pain with coitus. It is also important to address libido, as decreased sexual drive and ED are sometimes the earliest signs of endocrine abnormalities (e.g., increased prolactin, decreased testosterone levels). It is useful to ask whether the problem is confined to coitus with one partner or also involves other partners; ED not uncommonly arises in association with new or extramarital sexual relationships. Situational ED, as opposed to consistent ED, suggests psychogenic causes. Ejaculation is much less commonly affected than erection, but questions should be asked about whether ejaculation is normal, premature, delayed, or absent. Relevant risk factors should be identified, such as diabetes mellitus, coronary artery disease (CAD), and neurologic disorders. The patient's surgical history should be explored with an emphasis on bowel, bladder, prostate, and vascular procedures. A complete drug history is also important. Social changes that may precipitate ED are also crucial to the evaluation, including health worries, spousal death, divorce, relationship difficulties, and financial concerns.

Because ED commonly involves a host of endothelial cell risk factors, men with ED report higher rates of overt and silent myocardial infarction. Therefore, ED in an otherwise asymptomatic male warrants consideration of other vascular disorders, including CAD.

The physical examination is an essential element in the assessment of ED. Signs of hypertension as well as evidence of thyroid, hepatic, hematologic, cardiovascular, or renal diseases should be sought. An assessment should be made of the endocrine and vascular systems, the external genitalia, and the prostate gland. The penis should be palpated carefully along the corpora to detect fibrotic plaques. Reduced testicular size and loss of secondary sexual characteristics are suggestive of hypogonadism. Neurologic examination should include assessment of anal sphincter tone, investigation of the bulbocavernosus reflex, and testing for peripheral neuropathy.

Although hyperprolactinemia is uncommon, a serum prolactin level should be measured, as decreased libido and/or ED may be the presenting symptoms of a prolactinoma or another mass lesion of the sella (**Chap. 403**). The serum testosterone level should be measured, and if it is low, gonadotropins should be measured to determine whether hypogonadism is primary (testicular) or secondary (hypothalamic-pituitary) in origin (**Chap. 411**). If not performed recently, serum chemistries, complete blood count (CBC), and lipid profiles may be of value, as they can yield evidence of anemia, diabetes, hyperlipidemia, or other systemic diseases associated with ED. Determination of serum prostate-specific antigen (PSA) should be conducted according to recommended clinical guidelines (**Chap. 115**).

Additional diagnostic testing is rarely necessary in the evaluation of ED. However, in selected patients, specialized testing may provide insight into pathologic mechanisms of ED and aid in the selection of treatment options. Optional specialized testing includes (1) studies of nocturnal penile tumescence and rigidity, (2) vascular testing (in-office injection of vasoactive substances, penile Doppler ultrasound, penile angiography, dynamic infusion cavernosography/cavernosometry), (3) neurologic testing (biothesiometry-graded vibratory perception, somatosensory evoked potentials), and (4) psychological diagnostic tests. The information potentially gained from these procedures must be balanced against their invasiveness and cost.

TREATMENT MALE SEXUAL DYSFUNCTION

PATIENT EDUCATION

Patient and partner education is essential in the treatment of ED. In goal-directed therapy, education facilitates understanding of the disease, the results of the tests, and the selection of treatment. Discussion of treatment options helps clarify how treatment is best offered and stratify first- and second-line therapies. Patients with high-risk lifestyle issues such as obesity, smoking, alcohol abuse, and recreational drug use should be counseled on the role those factors play in the development of ED.

Therapies currently employed for the treatment of ED include oral PDE-5 inhibitor therapy (most commonly used), injection therapies, testosterone therapy, penile devices, and psychological therapy. In addition, limited data suggest that treatments for underlying risk factors and comorbidities—for example, weight loss, exercise, stress reduction, and smoking cessation—may improve erectile function. Decisions regarding therapy should take into account the preferences and expectations of patients and their partners.

ORAL AGENTS

Sildenafil, tadalafil, vardenafil, and avanafil are the only approved and effective oral agents for the treatment of ED. These four medications have markedly improved the management of ED because they are effective for the treatment of a broad range of causes, including psychogenic, diabetic, vasculogenic, post-radical prostatectomy (nerve-sparing procedures), and spinal cord injury. They belong to a class of medications that are selective and potent inhibitors of PDE-5, the predominant phosphodiesterase

isoform found in the penis. They are administered in graduated doses and enhance erections after sexual stimulation. The onset of action is approximately 30–120 min, depending on the medication used and other factors, such as recent food intake. Reduced initial doses should be considered for patients who are elderly, are taking concomitant alpha blockers, have renal insufficiency, or are taking medications that inhibit the CYP3A4 metabolic pathway in the liver (e.g., erythromycin, cimetidine, ketoconazole, and possibly itraconazole and mibefradil), as they may increase the serum concentration of the PDE-5 inhibitors (PDE-5i) or promote hypotension.

Initially, there were concerns about the cardiovascular safety of PDE-5i drugs. These agents can act as a mild vasodilator, and warnings exist about orthostatic hypotension with concomitant use of alpha blockers. The use of PDE-5i is not contraindicated in men who are also receiving alpha blockers, but they must be stabilized on this blood pressure medication prior to initiating therapy. Concerns also existed that use of PDE-5i would increase cardiovascular events. However, the safety of these drugs has been confirmed in several controlled trials with no increase in myocardial ischemic events or overall mortality compared to the general population.

Several randomized trials have demonstrated the efficacy of this class of medications. There are no compelling data to support the superiority of one PDE-5i over another. Subtle differences between agents have variable clinical relevance (**Table 67-2**).

Patients may fail to respond to a PDE-5i for several reasons (**Table 67-3**). Some patients may not tolerate PDE-5i secondary to adverse events from vasodilation in nonpenile tissues expressing PDE-5 or from the inhibition of homologous nonpenile isozymes (i.e., PDE-6 found in the retina). Abnormal vision attributed to the effects of PDE-5i on retinal PDE-6 is of short duration, reported only with sildenafil and not thought to be clinically significant. A more serious concern is the possibility that PDE-5i may cause nonarteritic anterior ischemic optic neuropathy; although data to support that association are limited, it is prudent to avoid the use of these agents in men with a prior history of nonarteritic anterior ischemic optic neuropathy.

Testosterone supplementation combined with a PDE-5i may be beneficial in improving erectile function in hypogonadal men with ED who are unresponsive to PDE-5i alone. These drugs do not affect ejaculation, orgasm, or sexual drive. Side effects associated with PDE-5i include headaches (19%), facial flushing (9%), dyspepsia (6%), and nasal congestion (4%). Approximately 7% of men using sildenafil may experience transient altered color vision (blue halo effect), and 6% of men taking tadalafil may experience loin pain. PDE-5i is contraindicated in men receiving nitrate therapy for cardiovascular disease, including agents delivered by the oral, sublingual, transnasal, and topical routes. These agents can potentiate its hypotensive effect and may result in profound shock. Likewise, amyl/butyl nitrate “poppers” may have a fatal synergistic effect on blood pressure. PDE-5i also should be avoided in patients with congestive heart failure and cardiomyopathy because of the risk of vascular collapse. Because sexual activity leads to an increase in physiologic expenditure (5–6 metabolic equivalents [METs]), physicians have been advised to exercise caution in prescribing any drug for sexual activity to those with active coronary disease, heart failure, borderline hypotension, or hypovolemia and to those on complex antihypertensive regimens.

Although the various forms of PDE-5i have a common mechanism of action, there are a few differences among the four agents (**Table 67-2**). Tadalafil is unique in its longer half-life, whereas avanafil appears to have the most rapid onset of action. All four drugs are effective for patients with ED of all ages, severities, and etiologies. Although there are pharmacokinetic and pharmacodynamic differences among these agents, clinically relevant differences are not clear.

ANDROGEN THERAPY

Testosterone replacement is used to treat both primary and secondary causes of hypogonadism (**Chap. 411**). Androgen supplementation

TABLE 67-2 CHARACTERISTICS OF PDE-5i MEDICATIONS

Drug	Onset of Action	Half-Life	Dose	Adverse Effects	Contraindications
Sildenafil	T _{max} 30-120 min Duration, 4 h High-fat meal decreases absorption ETOH may affect efficacy	2-5 h	25-100 mg Starting dose, 50 mg	Headache, flushing, dyspepsia, nasal congestion, altered vision	Nitrates Hypotension Cardiovascular risk factors Retinitis pigmentosa Change dose with some antiretrovirals Should be on stable dose of alpha blockers
Vardenafil	T _{max} 30-120 min Duration, 4-5 h High-fat meal decreases absorption ETOH may affect efficacy	4.5 h	5-10 mg	Headache, flushing, rhinitis, dyspepsia	Same as sildenafil May have minor prolongation of QT interval Concomitant use of Class I antiarrhythmic
Tadalafil	T _{max} 30-60 min Duration, 12-36 h Plasma concentration Not affected by food or ETOH	17.5 h	10 mg, 20 mg; 2.5 or 5 mg for daily dose	Headache, dyspepsia, back pain, nasal congestion, myalgia	Same as sildenafil
Avanafil	T _{max} 30 min Duration, 2 h Plasma concentration not affected by food	3-5 h	50, 100, and 200 mg	Headache, flushing, nasal congestion, nasopharyngitis, back pain	Same as sildenafil

Abbreviations: ETOH, alcohol; T_{max}, time to maximum plasma concentration.

in the setting of normal testosterone is rarely efficacious in the treatment of ED and is discouraged. Methods of androgen replacement include transdermal patches and gels, parenteral administration of long-acting testosterone esters (enanthate and cypionate), and oral preparations (17 α -alkylated derivatives) (Chap. 411). Oral androgen preparations have the potential for hepatotoxicity and should be avoided.

Men who receive testosterone should be reevaluated after 1-3 months and at least annually thereafter for testosterone levels, erectile function, and adverse effects, which may include gynecomastia, sleep apnea, development or exacerbation of lower urinary tract symptoms or BPH, prostate cancer, lowering of HDL, erythrocytosis, elevations of liver function tests, and reduced fertility. Periodic reevaluation should include measurement of CBC and PSA and digital rectal exam. Therapy should be discontinued in patients who do not respond within 3 months.

VACUUM CONSTRICTION DEVICES

Vacuum constriction devices (VCDs) are a well-established noninvasive therapy. They are a reasonable treatment alternative for select patients who cannot take sildenafil or do not desire other interventions. VCDs draw venous blood into the penis and use a constriction ring to restrict venous return and maintain tumescence. Adverse events with VCD include pain, numbness, bruising, and altered ejaculation. Additionally, many patients complain that the devices are cumbersome and that the induced erections have a nonphysiologic appearance and feel.

INTRAURETHRAL ALPROSTADIL

If a patient fails to respond to oral agents, a reasonable next choice is intraurethral or self-injection of vasoactive substances.

TABLE 67-3 ISSUES TO CONSIDER IF PATIENTS REPORT FAILURE OF PDE-5i TO IMPROVE ERECTILE DYSFUNCTION

- A trial of medication on at least 6 different days at the maximal dose should be made before declaring patient nonresponsive to PDE-5i use
- Confirm that patient did not take medication after a high-fat meal
- Failure to include physical and psychic stimulation at the time of foreplay to induce endogenous NO
- Unrecognized hypogonadism

Abbreviations: NO, nitric oxide; PDE-5i, phosphodiesterase type 5 inhibitor.

Intraurethral prostaglandin E₁ (alprostadil), in the form of a semi-solid pellet (doses of 125-1000 μ g), is delivered with an applicator. Approximately 65% of men receiving intraurethral alprostadil respond with an erection when tested in the office, but only 50% achieve successful coitus at home. Intraurethral insertion is associated with a markedly reduced incidence of priapism in comparison to intracavernosal injection.

INTRACAVERNOSAL SELF-INJECTION

Injection of synthetic formulations of alprostadil is effective in 70-80% of patients with ED, but discontinuation rates are high because of the invasive nature of administration. Doses range between 1 and 40 μ g. Injection therapy is contraindicated in men with a history of hypersensitivity to the drug and men at risk for priapism (hypercoagulable states, sickle cell disease). Side effects include local adverse events, prolonged erections, pain, and fibrosis with chronic use. Various combinations of alprostadil, phentolamine, and/or papaverine sometimes are used.

SURGERY

A less frequently used form of therapy for ED involves the surgical implantation of a semirigid or inflatable penile prosthesis. The choice of prosthesis is dependent on patient preference and should take into account body habitus and manual dexterity, which may affect the ability of the patient to manipulate the device. Because of the permanence of prosthetic devices, patients should be advised to first consider less invasive options for treatment. These surgical treatments are invasive, are associated with potential complications, and generally are reserved for treatment of refractory ED. Despite their high cost and invasiveness, penile prostheses are associated with high rates of patient and partner satisfaction.

SEX THERAPY

A course of sex therapy may be useful for addressing specific interpersonal factors that may affect sexual functioning. Sex therapy generally consists of in-session discussion and at-home exercises specific to the person and the relationship. Psychosexual therapy involves techniques such as sensate focus (nongenital massage), sensory awareness exercises, correction of misconceptions about sexuality, and interpersonal difficulties therapy (e.g., open communication about sexual issues, physical intimacy scheduling, and

behavioral interventions). These approaches may be useful in patients who have psychogenic or social components to their ED, although data from randomized trials are scanty and inconsistent. It is preferable if therapy includes both partners if the patient is involved in an ongoing relationship.

FEMALE SEXUAL DYSFUNCTION

Female sexual dysfunction (FSD) has traditionally included disorders of desire, arousal, pain, and muted orgasm. The associated risk factors for FSD are similar to those in males: cardiovascular disease, endocrine disorders, hypertension, neurologic disorders, and smoking (Table 67-4).

EPIDEMIOLOGY

Epidemiologic data are limited, but the available estimates suggest that as many as 43% of women complain of at least one sexual problem. Despite the recent interest in organic causes of FSD, desire and arousal phase disorders (including lubrication complaints) remain the most common presenting problems when surveyed in a community-based population.

PHYSIOLOGY OF THE FEMALE SEXUAL RESPONSE

The female sexual response requires the presence of estrogens. A role for androgens is also likely but less well established. In the CNS, estrogens and androgens work synergistically to enhance sexual arousal and response. A number of studies report enhanced libido in women during preovulatory phases of the menstrual cycle, suggesting that hormones involved in the ovulatory surge (e.g., estrogens) increase desire.

Sexual motivation is heavily influenced by context, including the environment and partner factors. Once sufficient sexual desire is reached, sexual arousal is mediated by the central and autonomic nervous systems. Cerebral sympathetic outflow is thought to increase desire, and peripheral parasympathetic activity results in clitoral vasocongestion and vaginal secretion (lubrication).

The neurotransmitters for clitoral corporal engorgement are similar to those in the male, with a prominent role for neural, smooth-muscle, and endothelial released nitric oxide (NO). A fine network of vaginal nerves and arterioles promotes a vaginal transudate. The major transmitters of this complex vaginal response are not certain, but roles for NO and vasointestinal polypeptide (VIP) are suspected. Investigators studying the normal female sexual response have challenged the long-held construct of a linear and unmitigated relationship between initial desire, arousal, vasocongestion, lubrication, and eventual orgasm. Caregivers

should consider a paradigm of a positive emotional and physical outcome with one, many, or no orgasmic peak and release.

Although there are anatomic differences as well as variation in the density of vascular and neural beds in males and females, the primary effectors of sexual response are strikingly similar. Intact sensation is important for arousal. Thus, reduced levels of sexual functioning are more common in women with peripheral neuropathies (e.g., diabetes). Vaginal lubrication is a transudate of serum that results from the increased pelvic blood flow associated with arousal. Vascular insufficiency from a variety of causes may compromise adequate lubrication and result in dyspareunia. Cavernal and arteriole smooth-muscle relaxation occurs via increased nitric oxide synthase (NOS) activity and produces engorgement in the clitoris and the surrounding vestibule. Orgasm requires an intact sympathetic outflow tract; hence, orgasmic disorders are common in female patients with spinal cord injuries.

APPROACH TO THE PATIENT: Female Sexual Dysfunction

Many women do not volunteer information about their sexual response. Open-ended questions in a supportive atmosphere are helpful in initiating a discussion of sexual fitness in women who are reluctant to discuss such issues. Once a complaint has been voiced, a comprehensive evaluation should be performed, including a medical history, a psychosocial history, a physical examination, and limited laboratory testing.

The history should include the usual medical, surgical, obstetric, psychological, gynecologic, sexual, and social information. Past experiences, intimacy, knowledge, and partner availability should also be ascertained. Medical disorders that may affect sexual health should be delineated. They include diabetes, cardiovascular disease, gynecologic conditions, obstetric history, depression, anxiety disorders, and neurologic disease. Medications should be reviewed as they may affect arousal, libido, and orgasm. The need for counseling and recognizing life stresses should be identified. The physical examination should assess the genitalia, including the clitoris. Pelvic floor examination may identify prolapse or other disorders. Laboratory studies are needed, especially if menopausal status is uncertain. Estradiol, follicle-stimulating hormone (FSH), and luteinizing hormone (LH) are usually obtained, and dehydroepiandrosterone (DHEA) should be considered as it reflects adrenal androgen secretion. A CBC, liver function assessment, and lipid studies may be useful, if not otherwise obtained. Complicated diagnostic evaluations such as clitoral Doppler ultrasonography and biothesiometry require expensive equipment and are of uncertain utility. It is important for the patient to identify which symptoms are most distressing.

The evaluation of FSD previously occurred mainly in a psychosocial context. However, inconsistencies between diagnostic categories based only on psychosocial considerations and the emerging recognition of organic etiologies have led to a new classification of FSD. This diagnostic scheme is based on four components that are not mutually exclusive: (1) *hypoactive sexual desire*—the persistent or recurrent lack of sexual thoughts and/or receptivity to sexual activity, which causes personal distress; hypoactive sexual desire may result from endocrine failure or may be associated with psychological or emotional disorders; (2) *sexual arousal disorder*—the persistent or recurrent inability to attain or maintain sexual excitement, which causes personal distress; (3) *orgasmic disorder*—the persistent or recurrent loss of orgasmic potential after sufficient sexual stimulation and arousal, which causes personal distress; and (4) *sexual pain disorder*—persistent or recurrent genital pain associated with noncoital sexual stimulation, which causes personal distress. This newer classification emphasizes “personal distress” as a requirement for dysfunction and provides clinicians with an organized framework for evaluation before or in conjunction with more traditional counseling methods.

TABLE 67-4 RISK FACTORS FOR FEMALE SEXUAL DYSFUNCTION

Neurologic disease: stroke, spinal cord injury, parkinsonism
Trauma, genital surgery, radiation
Endocrinopathies: diabetes, hyperprolactinemia
Liver and/or renal failure
Cardiovascular disease
Psychological factors and interpersonal relationship disorders: sexual abuse, life stressors
Medications
Antiandrogens: cimetidine, spironolactone
Antidepressants, alcohol, hypnotics, sedatives
Antiandrogens or GnRH antagonists
Antihistamines, sympathomimetic amines
Antihypertensives: diuretics, calcium channel blockers
Alkylating agents
Anticholinergics

Abbreviation: GnRH, gonadotropin-releasing hormone.

TREATMENT FEMALE SEXUAL DYSFUNCTION

GENERAL

An open discussion with the patient is important as couples may need to be educated about normal anatomy and physiologic responses, including the role of orgasm, in sexual encounters. Physiologic changes associated with aging and/or disease should be explained. Couples may need to be reminded that clitoral stimulation rather than coital intromission may be more beneficial.

Behavioral modification and nonpharmacologic therapies should be a first step. Patient and partner counseling may improve communication and relationship strains. Lifestyle changes involving known risk factors can be an important part of the treatment process. Emphasis on maximizing physical health and avoiding lifestyles (e.g., smoking, alcohol abuse) and medications likely to produce FSD is important (Table 67-4). The use of topical lubricants may address complaints of dyspareunia and dryness. Contributing medications such as antidepressants may need to be altered, including the use of medications with less impact on sexual function, dose reduction, medication switching, or drug holidays.

HORMONAL THERAPY

In postmenopausal women, estrogen replacement therapy may be helpful in treating vaginal atrophy, decreasing coital pain, and improving clitoral sensitivity (Chap. 413). Estrogen replacement in the form of local cream is the preferred method, as it avoids systemic side effects. Androgen levels in women decline substantially before menopause. However, low levels of testosterone or DHEA are not effective predictors of a positive therapeutic outcome with androgen therapy. The widespread use of exogenous androgens is not supported by the literature except in select circumstances (premature ovarian failure or menopausal states) and in secondary arousal disorders.

ORAL AGENTS

The efficacy of PDE-5i in FSD has been a marked disappointment in light of the proposed role of nitric oxide-dependent physiology in the normal female sexual response. The use of PDE-5i for FSD should be discouraged pending proof that it is effective.

CLITORAL VACUUM DEVICE

In patients with arousal and orgasmic difficulties, the option of using a clitoral vacuum device may be explored. This handheld battery-operated device has a small soft plastic cup that applies a vacuum over the stimulated clitoris. This causes increased cavernosal blood flow, engorgement, and vaginal lubrication.

TABLE 68-1 CAUSES OF HIRSUTISM

Gonadal hyperandrogenism
Ovarian hyperandrogenism
Polycystic ovary syndrome/functional ovarian hyperandrogenism
Ovarian steroidogenic blocks
Syndromes of extreme insulin resistance (e.g., lipodystrophy)
Ovarian neoplasms
Adrenal hyperandrogenism
Premature adrenarche
Functional adrenal hyperandrogenism
Congenital adrenal hyperplasia (nonclassic and classic)
Abnormal cortisol action/metabolism
Adrenal neoplasms
Other endocrine disorders
Cushing's syndrome
Hyperprolactinemia
Acromegaly
Peripheral androgen overproduction
Obesity
Idiopathic
Pregnancy-related hyperandrogenism
Hyperreactio luteinalis
Thecoma of pregnancy
Drugs
Androgens
Oral contraceptives containing androgenic progestins
Minoxidil
Phenytoin
Diazoxide
Cyclosporine
True hermaphroditism

increased muscle bulk, clitoromegaly, and increased libido; virilization is an ominous sign that suggests the possibility of an ovarian or adrenal neoplasm.

HAIR FOLLICLE GROWTH AND DIFFERENTIATION

Hair can be categorized as either *vellus* (fine, soft, and not pigmented) or *terminal* (long, coarse, and pigmented). The number of hair follicles does not change over an individual's lifetime, but the follicle size and type of hair can change in response to numerous factors, particularly androgens. Androgens are necessary for terminal hair and sebaceous gland development and mediate differentiation of pilosebaceous units (PSUs) into either a terminal hair follicle or a sebaceous gland. In the former case, androgens transform the vellus hair into a terminal hair; in the latter case, the sebaceous component proliferates and the hair remains vellus.

There are three phases in the cycle of hair growth: (1) *anagen* (growth phase), (2) *catagen* (involution phase), and (3) *telogen* (rest phase). Depending on the body site, hormonal regulation may play an important role in the hair growth cycle. For example, the eyebrows, eyelashes, and vellus hairs are androgen-insensitive, whereas the axillary and pubic areas are sensitive to low levels of androgens. Hair growth on the face, chest, upper abdomen, and back requires higher levels of androgens and is therefore more characteristic of the pattern typically seen in men. Androgen excess in women leads to increased hair growth in most androgen-sensitive sites except in the scalp region, where hair loss occurs because androgens cause scalp hairs to spend less time in the anagen phase.

Although androgen excess underlies most cases of hirsutism, there is only a modest correlation between androgen levels and the quantity of hair growth. This is due to the fact that hair growth from the follicle also

68

Hirsutism

David A. Ehrmann

Hirsutism, which is defined as androgen-dependent excessive male-pattern hair growth, affects approximately 10% of women. Hirsutism is most often idiopathic or the consequence of androgen excess associated with the polycystic ovarian syndrome (PCOS). Less frequently, it may result from adrenal androgen overproduction as occurs in non-classic congenital adrenal hyperplasia (CAH) (Table 68-1). Rarely, it is a sign of a serious underlying condition. Cutaneous manifestations commonly associated with hirsutism include acne and male-pattern balding (androgenic alopecia). *Virilization* refers to a condition in which androgen levels are sufficiently high to cause additional signs and symptoms, such as deepening of the voice, breast atrophy,

depends on local growth factors, and there is variability in end organ (PSU) sensitivity. Genetic factors and ethnic background also influence hair growth. In general, dark-haired individuals tend to be more hirsute than blond or fair individuals. Asians and Native Americans have relatively sparse hair in regions sensitive to high androgen levels, whereas people of Mediterranean descent are more hirsute.

CLINICAL ASSESSMENT

Historic elements relevant to the assessment of hirsutism include the age at onset and rate of progression of hair growth and associated symptoms or signs (e.g., acne). Depending on the cause, excess hair growth typically is first noted during the second and third decades of life. The growth is usually slow but progressive. Sudden development and rapid progression of hirsutism suggest the possibility of an androgen-secreting neoplasm, in which case virilization also may be present.

The age at onset of menstrual cycles (menarche) and the pattern of the menstrual cycle should be ascertained; irregular cycles from the time of menarche onward are more likely to result from ovarian rather than adrenal androgen excess. Associated symptoms such as galactorrhea should prompt evaluation for hyperprolactinemia (**Chap. 403**) and possibly hypothyroidism (**Chap. 405**). Hypertension, striae, easy bruising, centripetal weight gain, and weakness suggest hypercortisolism (Cushing's syndrome; **Chap. 406**). Rarely, patients with growth hormone excess (i.e., acromegaly) present with hirsutism. Use of medications such as phenytoin, minoxidil, and cyclosporine may be associated with androgen-independent excess hair growth (i.e., hypertrichosis). A family history of infertility and/or hirsutism may indicate disorders such as nonclassic CAH (**Chap. 406**). Lipodystrophy is often associated with increased ovarian androgen production that occurs as a consequence of insulin resistance. Patients with lipodystrophy have a preponderance of central fat distribution together with scant subcutaneous adipose tissue in the upper and lower extremities.

Physical examination should include measurement of height and weight and calculation of body mass index (BMI). A BMI >25 kg/m² is indicative of excess weight for height, and values >30 kg/m² are often seen in association with hirsutism, probably the result of increased conversion of androgen precursors to testosterone. Notation should be made of blood pressure, as adrenal causes may be associated with hypertension. Cutaneous signs sometimes associated with androgen excess and insulin resistance include acanthosis nigricans and skin tags. Body fat distribution should also be noted.

An objective clinical assessment of hair distribution and quantity is central to the evaluation in any woman presenting with hirsutism. This assessment permits the distinction between hirsutism and hypertrichosis and provides a baseline reference point to gauge the response to treatment. A simple and commonly used method to grade hair growth is the modified scale of Ferriman and Gallwey (**Fig. 68-1**), in which each of nine androgen-sensitive sites is graded from 0 to 4. Approximately 95% of white women have a score below 8 on this scale; thus, it is normal for most women to have some hair growth in androgen-sensitive sites. Scores above 8 suggest excess androgen-mediated hair growth, a finding that should be assessed further by means of hormonal evaluation (see below). In racial/ethnic groups that are less likely to manifest hirsutism (e.g., Asian women), additional cutaneous evidence of androgen excess should be sought, including pustular acne and thinning scalp hair.

HORMONAL EVALUATION

Androgens are secreted by the ovaries and adrenal glands in response to their respective tropic hormones: luteinizing hormone (LH) and adrenocorticotropic hormone (ACTH). The principal circulating steroids involved in the etiology of hirsutism are testosterone, androstenedione, and dehydroepiandrosterone (DHEA) and its sulfated form (DHEAS). The ovaries and adrenal glands normally contribute about equally to testosterone production. Approximately half of the total testosterone originates from direct glandular secretion, and the remainder is derived from the peripheral conversion of androstenedione and DHEA (**Chap. 411**).

Although it is the most important circulating androgen, testosterone is in effect the penultimate androgen in mediating hirsutism; it is converted to the more potent dihydrotestosterone (DHT) by the enzyme 5 α -reductase, which is located in the PSU. DHT has a higher affinity for, and slower dissociation from, the androgen receptor. The local production of DHT allows it to serve as the primary mediator of androgen action at the level of the pilosebaceous unit. There are two isoenzymes of 5 α -reductase: Type 2 is found in the prostate gland and in hair follicles, and type 1 is found primarily in sebaceous glands.

One approach to the evaluation of hirsutism is depicted in **Fig. 68-2**. In addition to measuring blood levels of testosterone and DHEAS, it is important to measure the level of free (or unbound) testosterone. The fraction of testosterone that is not bound to its carrier protein, sex hormone-binding globulin (SHBG), is biologically available for conversion to DHT and binding to androgen receptors. Hyperinsulinemia and/or androgen excess decrease hepatic production of SHBG, resulting in levels of total testosterone within the high-normal range, whereas the unbound hormone is elevated more substantially. Although there is a decline in ovarian testosterone production after menopause, ovarian estrogen production decreases to an even greater extent, and the concentration of SHBG is reduced. Consequently, there is an increase in the relative proportion of unbound testosterone, and it may exacerbate hirsutism after menopause.

A baseline plasma total testosterone level >12 nmol/L (>3.5 ng/mL) usually indicates a virilizing tumor, whereas a level >7 nmol/L (>2 ng/mL) is suggestive. A basal DHEAS level >18.5 μ mol/L (>7000 μ g/L) suggests an adrenal tumor. Although DHEAS has been proposed as a "marker" of predominant adrenal androgen excess, it is not unusual to find modest elevations in DHEAS among women with PCOS. Computed tomography (CT) or magnetic resonance imaging (MRI) should be used to localize an adrenal mass, and transvaginal ultrasound usually suffices to identify an ovarian mass if clinical evaluation and hormonal levels suggest these possibilities.

PCOS is the most common cause of ovarian androgen excess (**Chap. 412**). An increased ratio of LH to follicle-stimulating hormone (FSH) is characteristic in carefully studied patients with PCOS. However, because of the pulsatile nature of gonadotropin secretion, this finding may be absent in up to half of women with PCOS. Therefore, measurement of plasma LH and FSH is not needed to make a diagnosis of PCOS. Transvaginal ultrasound classically shows enlarged ovaries and increased stroma in women with PCOS. However, cystic ovaries also may be found in women without clinical or laboratory features of PCOS.

It has been suggested that the measurement of circulating levels of antimüllerian hormone (AMH) may help in making the diagnosis of PCOS; however, this remains controversial. AMH levels reflect ovarian reserve and correlate with follicular number. Measurement of AMH can be useful when considering premature ovarian insufficiency in a patient who presents with oligomenorrhea, in which case a subnormal level of AMH will be present.

Because adrenal androgens are readily suppressed by low doses of glucocorticoids, the dexamethasone androgen-suppression test may broadly distinguish ovarian from adrenal androgen overproduction. A blood sample is obtained before and after the administration of dexamethasone (0.5 mg orally every 6 h for 4 days). An adrenal source is suggested by suppression of unbound testosterone into the normal range; incomplete suppression suggests ovarian androgen excess. An overnight 1-mg dexamethasone suppression test, with measurement of 8:00 a.m. serum cortisol, is useful when there is clinical suspicion of Cushing's syndrome (**Chap. 406**).

Nonclassic CAH is most commonly due to 21-hydroxylase deficiency but also can be caused by autosomal recessive defects in other steroidogenic enzymes necessary for adrenal corticosteroid synthesis (**Chap. 406**). Because of the enzyme defect, the adrenal gland cannot secrete glucocorticoids (especially cortisol) efficiently. This results in diminished negative feedback inhibition of ACTH, leading to compensatory adrenal hyperplasia and the accumulation of steroid precursors that subsequently are converted to androgen. Deficiency

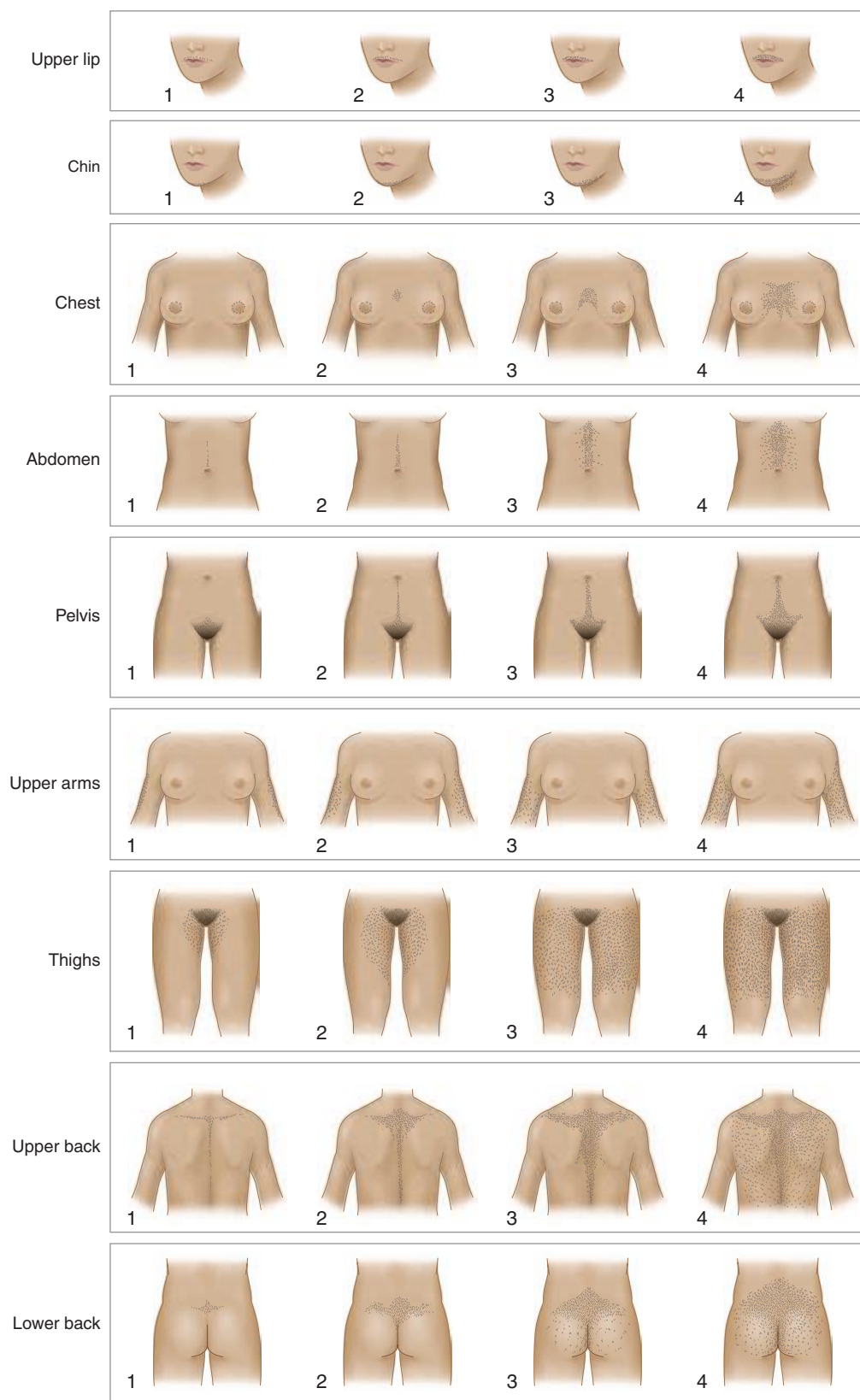


FIGURE 68-1 Hirsutism scoring scale of Ferriman and Gallwey. The nine body areas that have androgen-sensitive areas are graded from 0 (no terminal hair) to 4 (frankly virile) to obtain a total score. A normal hirsutism score is <8 . (Modified from DA Ehrmann et al: *Hyperandrogenism, hirsutism, and polycystic ovary syndrome*, in LJ DeGroot and JL Jameson [eds], *Endocrinology*, 5th ed. Philadelphia, Saunders, 2006; with permission.)

of 21-hydroxylase can be reliably excluded by determining a morning 17-hydroxyprogesterone level <6 nmol/L (<2 $\mu\text{g/L}$) (drawn in the follicular phase). Alternatively, 21-hydroxylase deficiency can be diagnosed by measurement of 17-hydroxyprogesterone 1 h after the administration of 250 μg of synthetic ACTH (cosyntropin) intravenously.

TREATMENT HIRSUTISM

Treatment of hirsutism may be accomplished pharmacologically or by mechanical means of hair removal. Nonpharmacologic treatments should be considered in all patients either as the only treatment or as an adjunct to drug therapy.

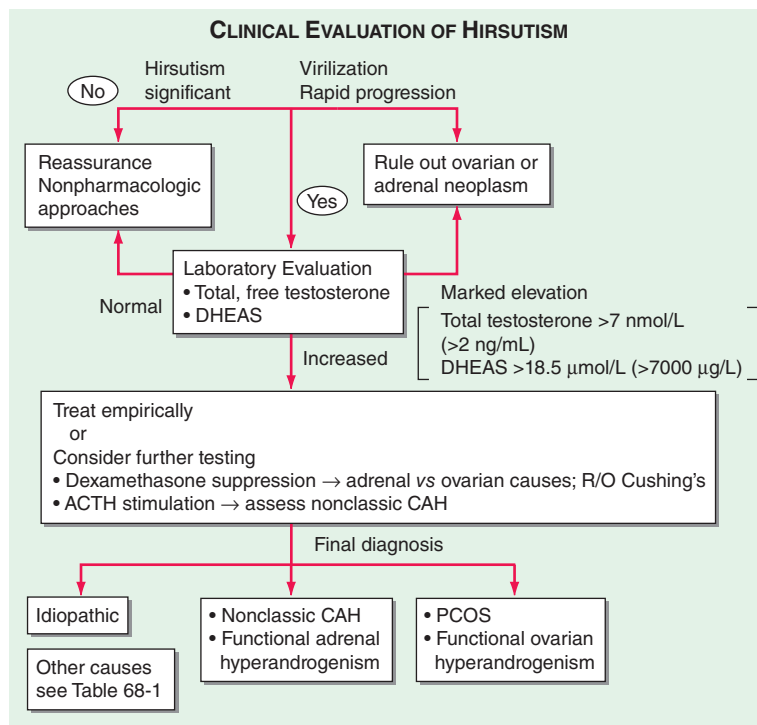


FIGURE 68-2 Algorithm for the evaluation and differential diagnosis of hirsutism. ACTH, adrenocorticotropic hormone; CAH, congenital adrenal hyperplasia; DHEAS, sulfated form of dehydroepiandrosterone; PCOS, polycystic ovarian syndrome.

Nonpharmacologic treatments include (1) bleaching; (2) depilatory (removal from the skin surface), such as shaving and chemical treatments; and (3) epilatory (removal of the hair including the root), such as plucking, waxing, electrolysis, and laser therapy. Despite perceptions to the contrary, shaving does not increase the rate or density of hair growth. Chemical depilatory treatments may be useful for mild hirsutism that affects only limited skin areas, though they can cause skin irritation. Wax treatment removes hair temporarily but is uncomfortable. Electrolysis is effective for more permanent hair removal, particularly in the hands of a skilled electrologist. Laser phototherapy appears to be efficacious for hair removal. It delays hair regrowth and causes permanent hair removal in most patients. The long-term effects and complications associated with laser treatment are being evaluated.

Pharmacologic therapy is directed at interrupting one or more of the steps in the pathway of androgen synthesis and action: (1) suppression of adrenal and/or ovarian androgen production; (2) enhancement of androgen-binding to plasma-binding proteins, particularly SHBG; (3) impairment of the peripheral conversion of androgen precursors to active androgen; and (4) inhibition of androgen action at the target tissue level. Attenuation of hair growth is typically not evident until 4–6 months after initiation of medical treatment and in most cases leads to only a modest reduction in hair growth.

Combination estrogen-progestin therapy in the form of an oral contraceptive is usually the first-line endocrine treatment for hirsutism and acne, after cosmetic and dermatologic management. The estrogenic component of most oral contraceptives currently in use is either ethinyl estradiol or mestranol. The suppression of LH leads to reduced production of ovarian androgens. The reduced androgen levels also result in a dose-related increase in SHBG, thus lowering the fraction of unbound plasma testosterone. Combination therapy also has been demonstrated to decrease DHEAS, perhaps by reducing ACTH levels. Estrogens also have a direct, dose-dependent suppressive effect on sebaceous cell function.

The choice of a specific oral contraceptive should be predicated on the progestational component, as progestins vary in their suppressive effect on SHBG levels and in their androgenic

potential. Ethynodiol diacetate has relatively low androgenic potential, whereas progestins such as norgestrel and levonorgestrel are particularly androgenic, as judged from their attenuation of the estrogen-induced increase in SHBG. Norgestimate exemplifies the newer generation of progestins that are virtually nonandrogenic. Drospirenone, an analogue of spironolactone that has both antiminerocorticoid and antiandrogenic activities, has been approved for use as a progestational agent in combination with ethinyl estradiol.

Oral contraceptives are contraindicated in women with a history of thromboembolic disease and women with increased risk of breast or other estrogen-dependent cancers (Chap. 413). There is a relative contraindication to the use of oral contraceptives in smokers and those with hypertension or a history of migraine headaches. In most trials, estrogen-progestin therapy alone improves the extent of acne by a maximum of 50–70%. The effect on hair growth may not be evident for 6 months, and the maximum effect may require 9–12 months owing to the length of the hair growth cycle. Improvements in hirsutism are typically in the range of 20%, but there may be an arrest of further progression of hair growth.

Adrenal androgens are more sensitive than cortisol to the suppressive effects of glucocorticoids. Therefore, glucocorticoids are the mainstay of treatment in patients with CAH. Although glucocorticoids have been reported to restore ovulatory function in some women with PCOS, this effect is highly variable. Because of side effects from excessive glucocorticoids, low doses should be used. Dexamethasone (0.2–0.5 mg) or prednisone (5–10 mg) should be taken at bedtime to achieve maximal suppression by inhibiting the nocturnal surge of ACTH.

Cyproterone acetate is the prototypic antiandrogen. It acts mainly by competitive inhibition of the binding of testosterone and DHT to the androgen receptor. In addition, it may enhance the metabolic clearance of testosterone by inducing hepatic enzymes. Although not available for use in the United States, cyproterone acetate is widely used in Canada, Mexico, and Europe. Cyproterone (50–100 mg) is given on days 1–15 and ethinyl estradiol (50 μg) is given on days 5–26 of the menstrual cycle. Side effects include irregular uterine bleeding, nausea, headache, fatigue, weight gain, and decreased libido.

Spironolactone, which usually is used as a mineralocorticoid antagonist, is also a weak antiandrogen. It is almost as effective as cyproterone acetate when used at high enough doses (100–200 mg daily). Patients should be monitored intermittently for hyperkalemia or hypotension, although these side effects are uncommon. Pregnancy should be avoided because of the risk of feminization of a male fetus. Spironolactone can also cause menstrual irregularity. It often is used in combination with an oral contraceptive, which suppresses ovarian androgen production and helps prevent pregnancy.

Flutamide is a potent nonsteroidal antiandrogen that is effective in treating hirsutism, but concerns about the induction of hepatocellular dysfunction have limited its use. Finasteride is a competitive inhibitor of 5 α -reductase type 2. Beneficial effects on hirsutism have been reported, but the predominance of 5 α -reductase type 1 in the PSU appears to account for its limited efficacy. Finasteride would also be expected to impair sexual differentiation in a male fetus, and it should not be used in women who may become pregnant.

Eflornithine cream (Vaniqa) has been approved as a novel treatment for unwanted facial hair in women, but long-term efficacy remains to be established. It can cause skin irritation under exaggerated conditions of use. Ultimately, the choice of any specific agent(s) must be tailored to the unique needs of the patient being treated. As noted previously, pharmacologic treatments for hirsutism should be used in conjunction with nonpharmacologic approaches. It is also helpful to review the pattern of female hair distribution in the normal population to dispel unrealistic expectations.

69 Menstrual Disorders and Pelvic Pain

Janet E. Hall

Menstrual dysfunction can signal an underlying abnormality that may have long-term health consequences. Although frequent or prolonged bleeding usually prompts a woman to seek medical attention, infrequent or absent bleeding may seem less troubling and the patient may not bring it to the attention of the physician. Thus, a focused menstrual history is a critical part of every encounter with a female patient. Pelvic pain is a common complaint that may relate to an abnormality of the reproductive organs but also may be of gastrointestinal, urinary tract, or musculoskeletal origin. Depending on its cause, pelvic pain may require urgent surgical attention.

MENSTRUAL DISORDERS

DEFINITION AND PREVALENCE

Amenorrhea refers to the absence of menstrual periods. Amenorrhea is classified as *primary* if menstrual bleeding has never occurred in the absence of hormonal treatment or *secondary* if menstrual periods cease for 3–6 months. Primary amenorrhea is a rare disorder that occurs in <1% of the female population. However, between 3 and 5% of women experience at least 3 months of secondary amenorrhea in any specific year. There is no evidence that race or ethnicity influences the prevalence of amenorrhea. However, because of the importance of adequate nutrition for normal reproductive function, both the age at menarche and the prevalence of secondary amenorrhea vary significantly in different parts of the world.

Oligomenorrhea is defined as a cycle length >35 days or <10 menses per year. Both the frequency and the amount of vaginal bleeding are irregular in oligomenorrhea, and minimal symptoms (premenstrual breast tenderness, food cravings, mood lability), suggestive of ovulation,

are variably present. Anovulation can also present with intermenstrual intervals <24 days or vaginal bleeding for >7 days. Frequent or heavy irregular bleeding is termed *dysfunctional uterine bleeding* if anatomic uterine and outflow tract lesions or a bleeding diathesis has been excluded.

Primary Amenorrhea The absence of menses by age 16 has been used traditionally to define primary amenorrhea. However, other factors, such as growth, secondary sexual characteristics, the presence of cyclic pelvic pain, and the secular trend toward an earlier age of menarche, particularly in African-American girls, also influence the age at which primary amenorrhea should be investigated. Thus, an evaluation for amenorrhea should be initiated by age 15 or 16 in the presence of normal growth and secondary sexual characteristics; age 13 in the absence of secondary sexual characteristics or if height is less than the third percentile; age 12 or 13 in the presence of breast development and cyclic pelvic pain; or within 2 years of breast development if menarche, defined by the first menstrual period, has not occurred.

Secondary Amenorrhea or Oligomenorrhea Anovulation and irregular cycles are relatively common for up to 2 years after menarche and for 1–2 years before the final menstrual period. In the intervening years, menstrual cycle length is ~28 days, with an intermenstrual interval normally ranging between 25 and 35 days. Cycle-to-cycle variability in an individual woman who is ovulating consistently is generally \pm 2 days. Pregnancy is the most common cause of amenorrhea and should be excluded early in any evaluation of menstrual irregularity. However, many women occasionally miss a single period. Three or more months of secondary amenorrhea should prompt an evaluation, as should a history of intermenstrual intervals >35 or <21 days or bleeding that persists for >7 days.

DIAGNOSIS

Evaluation of menstrual dysfunction depends on understanding the interrelationships between the four critical components of the reproductive tract: (1) the hypothalamus, (2) the pituitary, (3) the ovaries, and (4) the uterus and outflow tract (Fig. 69-1; Chap. 412). This system is maintained by complex negative and positive feedback loops involving the ovarian steroids (estradiol and progesterone) and peptides (inhibin B and inhibin A) and the hypothalamic (gonadotropin-releasing hormone [GnRH]) and pituitary (follicle-stimulating hormone [FSH] and luteinizing hormone [LH]) components of this system (Fig. 69-1).

Disorders of menstrual function can be thought of in two main categories: disorders of the uterus and outflow tract and disorders of ovulation. Many of the conditions that cause primary amenorrhea are congenital but go unrecognized until the time of normal puberty (e.g., genetic, chromosomal, and anatomic abnormalities). All causes of secondary amenorrhea also can cause primary amenorrhea.

Disorders of the Uterus or Outflow Tract Abnormalities of the uterus and outflow tract typically present as primary amenorrhea. In patients with normal pubertal development and a blind vagina, the differential diagnosis includes *obstruction* by a transverse vaginal septum or imperforate hymen; *müllerian agenesis* (Mayer-Rokitansky-Kuster-Hauser syndrome), which has been associated with mutations in the *WNT4* gene; and *androgen insensitivity syndrome* (AIS), which is an X-linked recessive disorder that accounts for ~10% of all cases of primary amenorrhea (Chap. 411). Patients with AIS have a 46,XY karyotype, but because of the lack of androgen receptor responsiveness, those with complete AIS have severe underandrogenization and female external genitalia. The absence of pubic and axillary hair distinguishes them clinically from patients with müllerian agenesis, as does an elevated testosterone level. *Asherman's syndrome* presents as secondary amenorrhea or hypomenorrhea and results from partial or complete obliteration of the uterine cavity by adhesions that prevent normal growth and shedding of the endometrium. Curettage performed for pregnancy complications accounts for >90% of cases; genital tuberculosis is an important cause in regions where it is endemic.

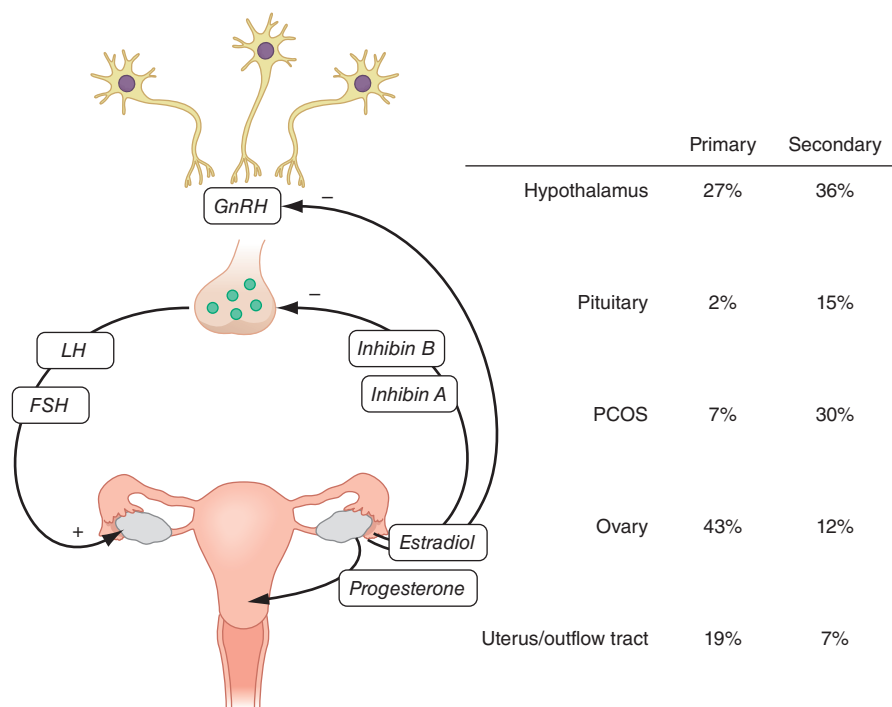


FIGURE 69-1 Role of the hypothalamic-pituitary-gonadal axis in the etiology of amenorrhea. Gonadotropin-releasing hormone (GnRH) secretion from the hypothalamus stimulates follicle-stimulating hormone (FSH) and luteinizing hormone (LH) secretion from the pituitary to induce ovarian folliculogenesis and steroidogenesis. Ovarian secretion of estradiol and progesterone controls the shedding of the endometrium, resulting in menses, and, in combination with the inhibins, provides feedback regulation of the hypothalamus and pituitary to control secretion of FSH and LH. The prevalence of amenorrhea resulting from abnormalities at each level of the reproductive system (hypothalamus, pituitary, ovary, uterus, and outflow tract) varies depending on whether amenorrhea is primary or secondary. PCOS, polycystic ovarian syndrome.

TREATMENT

DISORDERS OF THE UTERUS OR OUTFLOW TRACT

Obstruction of the outflow tract requires surgical correction. The risk of endometriosis is increased with this condition, perhaps because of retrograde menstrual flow. *Müllerian agenesis* also may require surgical intervention to allow sexual intercourse, although vaginal dilatation is adequate in some patients. Because ovarian function is normal, assisted reproductive techniques can be used with a surrogate carrier. *Androgen resistance syndrome* requires gonadectomy because there is risk of gonadoblastoma in the dysgenetic gonads. Whether this should be performed in early childhood or after completion of breast development is controversial. Estrogen replacement is indicated after gonadectomy, and vaginal dilatation may be required to allow sexual intercourse.

Disorders of Ovulation Once uterus and outflow tract abnormalities have been excluded, other causes of amenorrhea involve disorders of ovulation. The differential diagnosis is based on the results of initial tests, including a pregnancy test, an FSH level (to determine whether the cause is likely to be ovarian or central), and assessment of hyperandrogenism (Fig. 69-2).

HYPOGONADOTROPIC HYPOGONADISM Low estrogen levels in combination with normal or low levels of LH and FSH are seen with anatomic, genetic, or functional abnormalities that interfere with hypothalamic GnRH secretion or normal pituitary responsiveness to GnRH. Although relatively uncommon, tumors and infiltrative diseases should be considered in the differential diagnosis of hypogonadotropic hypogonadism (Chap. 403). These disorders may present with primary

or secondary amenorrhea. They may occur in association with other features suggestive of hypothalamic or pituitary dysfunction, such as short stature, diabetes insipidus, galactorrhea, and headache. Hypogonadotropic hypogonadism also may be seen after cranial irradiation. In the postpartum period, it may be caused by pituitary necrosis (Sheehan's syndrome) or lymphocytic hypophysitis. Because reproductive dysfunction is commonly associated with hyperprolactinemia from neuroanatomic lesions or medications, prolactin should be measured in all patients with hypogonadotropic hypogonadism (Chap. 403).

Isolated hypogonadotropic hypogonadism (IHH) occurs in women, although it is three times more common in men. IHH generally presents with primary amenorrhea, although 50% have some degree of breast development, and one to two menses have been described in ~10%. IHH is associated with anosmia in about 50% of women (termed Kallmann's syndrome). Genetic causes of IHH have been identified in ~60% of patients (Chaps. 411 and 412).

Functional hypothalamic amenorrhea (HA) is caused by a mismatch between energy expenditure and energy intake. Recent studies suggest that variants in genes associated with IHH may increase susceptibility to these environmental inputs, accounting in part for the clinical variability in this disorder. Leptin secretion may play a key role in transducing the signals from the periphery to the hypothalamus in HA. The hypothalamic-pituitary-adrenal axis also may play a role. The diagnosis of HA generally can be made on the basis of a careful history, a physical examination, and the demonstration of low levels of gonadotropins and normal prolactin levels.

Eating disorders and chronic disease must be specifically excluded. An atypical history, headache, signs of other hypothalamic dysfunction, or hyperprolactinemia, even if mild, necessitates cranial imaging with computed tomography (CT) or magnetic resonance imaging (MRI) to exclude a neuroanatomic cause.

HYPERGONADOTROPIC HYPOGONADISM Ovarian failure is considered premature when it occurs in women <40 years old and accounts for ~10% of secondary amenorrhea. *Primary ovarian insufficiency* (POI) has generally replaced the terms *premature menopause* and *premature ovarian failure* in recognition that this disorder represents a continuum of impaired ovarian function. Ovarian insufficiency is associated with the loss of negative-feedback restraint on the hypothalamus and pituitary, resulting in increased FSH and LH levels. FSH is a better marker of ovarian failure as its levels are less variable than those of LH. Antimüllerian hormone (AMH) levels will also be low in patients with POI, but are more frequently used in management of infertility. As with natural menopause, POI may wax and wane, and serial measurements may be necessary to establish the diagnosis.

Once the diagnosis of POI has been established, further evaluation is indicated because of other health problems that may be associated with POI. For example, POI occurs in association with a variety of chromosomal abnormalities, including Turner's syndrome, autoimmune polyglandular failure syndromes, radio- and chemotherapy, and galactosemia. The recognition that early ovarian failure occurs in premutation carriers of the fragile X syndrome is important because of the increased risk of severe mental retardation in male children with *FMRI* mutations. In the majority of cases, a cause for POI is not determined. Although there are increasing reports of genetic mutations in individuals and families with POI, testing for other than chromosomal abnormalities and *FMRI* mutations is not recommended.

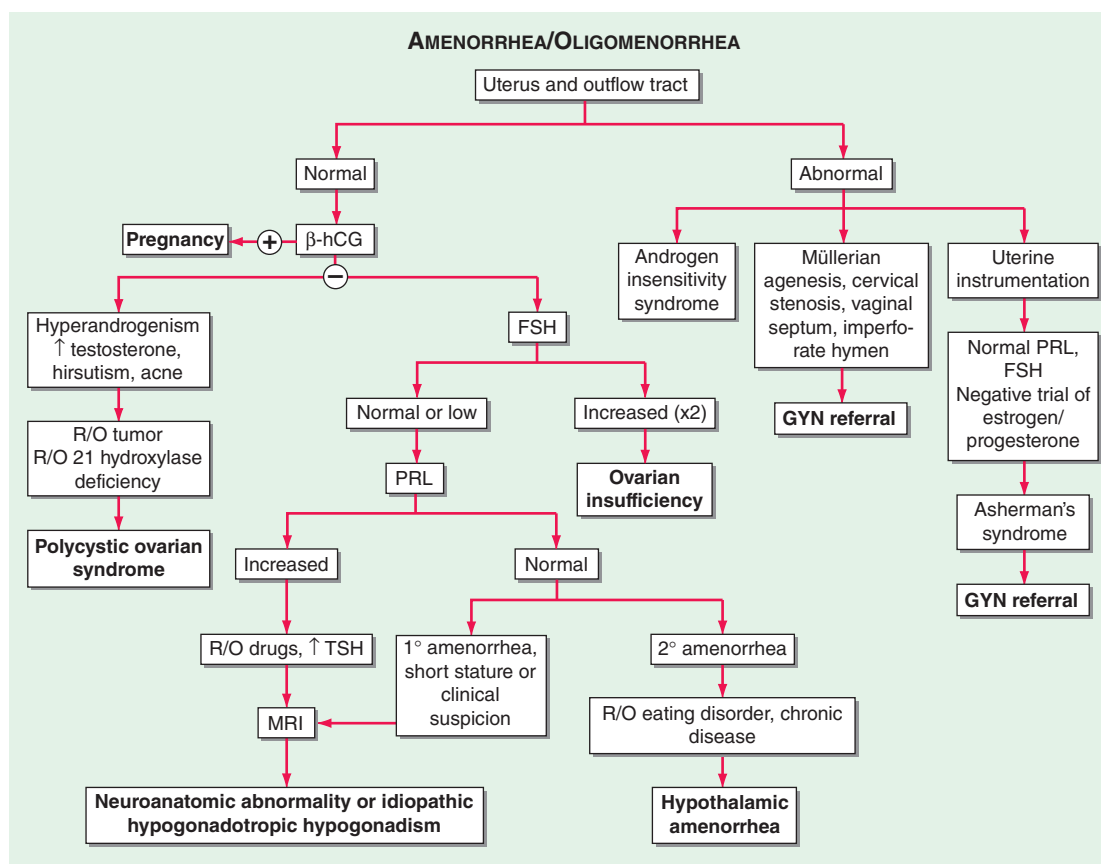


FIGURE 69-2 Algorithm for evaluation of amenorrhea. β -hCG, human chorionic gonadotropin; FSH, follicle-stimulating hormone; GYN, gynecologist; MRI, magnetic resonance imaging; PRL, prolactin; R/O, rule out; TSH, thyroid-stimulating hormone.

Hypergonadotropic hypogonadism occurs rarely in other disorders, such as mutations in the FSH or LH receptors. Aromatase deficiency and 17 α -hydroxylase deficiency are associated with decreased estrogen and elevated gonadotropins and with hyperandrogenism and hypertension, respectively. Gonadotropin-secreting tumors in women of reproductive age generally present with high, rather than low, estrogen levels and cause ovarian hyperstimulation or dysfunctional bleeding.

TREATMENT HYPO- AND HYPERGONADOTROPIC CAUSES OF AMENORRHEA

Amenorrhea almost always is associated with chronically low levels of estrogen, whether it is caused by hypogonadotropic hypogonadism or ovarian insufficiency. Development of secondary sexual characteristics requires gradual titration of estradiol replacement with eventual addition of progestin. Hormone replacement with either low-dose estrogen/progesterone regimens or oral contraceptive pills is recommended until the usual age of menopause for bone and cardiovascular protection. Patients with hypogonadotropic hypogonadism who are interested in fertility require treatment with exogenous FSH combined with LH or pulsatile GnRH. Patients with ovarian failure can consider oocyte donation, which has a high rate of success in this population, although its use in women with Turner's syndrome is limited by significant maternal cardiovascular risk.

POLYCYSTIC OVARIAN SYNDROME (PCOS) PCOS is diagnosed based on a combination of clinical or biochemical evidence of hyperandrogenism, amenorrhea or oligomenorrhea, and the ultrasound appearance of polycystic ovaries. Approximately half of patients with PCOS are obese, and abnormalities in insulin dynamics are common, as is metabolic syndrome. Symptoms generally begin shortly after menarche and are slowly progressive. Lean oligo-ovulatory patients with PCOS

generally have high LH levels in the presence of normal to low levels of FSH and estradiol. The LH/FSH ratio is less pronounced in obese patients in whom insulin resistance is a more prominent feature.

TREATMENT POLYCYSTIC OVARIAN SYNDROME

A major abnormality in patients with PCOS is the failure of regular, predictable ovulation. Thus, these patients are at risk for the development of dysfunctional bleeding and endometrial hyperplasia associated with unopposed estrogen exposure. Endometrial protection can be achieved with the use of oral contraceptives or progestins (medroxyprogesterone acetate, 5–10 mg, or promethrium, 200 mg daily for 10–14 days of each month). Oral contraceptives are also useful for management of hyperandrogenic symptoms, as are spironolactone and cyproterone acetate (not available in the United States), which function as weak androgen receptor blockers. Management of the associated metabolic syndrome may be appropriate for some patients (Chap. 422). For patients interested in fertility, weight control is a critical first step. Clomiphene citrate is highly effective as a first-line treatment, and there is increasing evidence that the aromatase inhibitor letrozole may also be effective. Exogenous gonadotropins can be used by experienced practitioners; a diagnosis of polycystic ovaries in the presence or absence of cycle abnormalities increases the risk of hyperstimulation.

PELVIC PAIN

The mechanisms that cause pelvic pain are similar to those that cause abdominal pain (Chap. 20) and include inflammation of the parietal peritoneum, obstruction of hollow viscera, vascular disturbances, and pain originating in the abdominal wall. Pelvic pain may reflect pelvic disease per se but also may reflect extrapelvic disorders that refer pain to the pelvis. In up to 60% of cases, pelvic pain can be attributed to

gastrointestinal problems, including appendicitis, cholecystitis, infections, intestinal obstruction, diverticulitis, and inflammatory bowel disease. Urinary tract and musculoskeletal disorders are also common causes of pelvic pain.

APPROACH TO THE PATIENT: Pelvic Pain

As with all types of abdominal pain, the first priority is to identify life-threatening conditions (shock, peritoneal signs) that may require emergent surgical management. The possibility of pregnancy should be identified as soon as possible by menstrual history and/or testing. A thorough history that includes the type, location, radiation, and status with respect to increasing or decreasing severity can help identify the cause of acute pelvic pain. Specific associations with vaginal bleeding, sexual activity, defecation, urination, movement, or eating should be specifically sought. Determination of whether the pain is acute versus chronic and cyclic versus noncyclic will direct further investigation (Table 69-1). However, disorders that cause cyclic pain occasionally may cause noncyclic pain, and the converse is also true.

ACUTE PELVIC PAIN

Pelvic inflammatory disease most commonly presents with bilateral lower abdominal pain. It is generally of recent onset and is exacerbated by intercourse or jarring movements. Fever is present in about half of these patients; abnormal uterine bleeding occurs in about one-third. New vaginal discharge, urethritis, and chills may be present but are less specific signs. *Adnexal pathology* can present acutely and may be due to rupture, bleeding or torsion of cysts, or, much less commonly, neoplasms of the ovary, fallopian tubes, or paraovarian areas. Fever may be present with ovarian torsion. *Ectopic pregnancy* is associated with right- or left-sided lower abdominal pain, with clinical signs generally appearing 6–8 weeks after the last normal menstrual period. Amenorrhea is present in ~75% of cases and vaginal bleeding in ~50% of cases. Orthostatic signs and fever may be present. Risk factors include the presence of known tubal disease, previous ectopic pregnancies, a history of infertility, diethylstilbestrol (DES) exposure of the mother in utero, or a history of pelvic infections. *Threatened abortion* may also present with amenorrhea, abdominal pain, and vaginal bleeding. Although more common than ectopic pregnancy, it is rarely associated with systemic signs. *Uterine pathology* includes endometritis and, less frequently, degenerating leiomyomas (fibroids). Endometritis often is associated with vaginal bleeding and systemic signs of infection. It occurs in the setting of sexually transmitted infections, uterine instrumentation, or postpartum infection.

TABLE 69-1 CAUSES OF PELVIC PAIN

	Acute	Chronic
Cyclic pelvic pain		Premenstrual symptoms Mittelschmerz Dysmenorrhea Endometriosis
Noncyclic pelvic pain	Pelvic inflammatory disease Ruptured or hemorrhagic ovarian cyst, endometrioma, or ovarian torsion Ectopic pregnancy Endometritis Acute growth or degeneration of uterine myoma Threatened abortion	Pelvic congestion syndrome Adhesions and retroversion of the uterus Pelvic malignancy Vulvodynia Chronic pelvic inflammatory disease Tuberculous salpingitis History of sexual abuse

A sensitive pregnancy test, complete blood count with differential, urinalysis, tests for chlamydial and gonococcal infections, and abdominal ultrasound aid in making the diagnosis and directing further management.

TREATMENT ACUTE PELVIC PAIN

Treatment of acute pelvic pain depends on the suspected etiology but may require surgical or gynecologic intervention. Conservative management is an important consideration for ovarian cysts, if torsion is not suspected, to avoid unnecessary pelvic surgery and the subsequent risk of infertility due to adhesions. Surgical treatment may be required for ectopic pregnancies; however, approximately 35% of ectopic pregnancies are unruptured and may be appropriate for treatment with methotrexate, which is effective in ~90% of cases.

CHRONIC PELVIC PAIN

Some women experience discomfort at the time of ovulation (*mittelschmerz*). The pain can be quite intense but is generally of short duration. The mechanism is thought to involve rapid expansion of the dominant follicle, although it also may be caused by peritoneal irritation by follicular fluid released at the time of ovulation. Many women experience premenstrual symptoms such as breast discomfort, food cravings, and abdominal bloating or discomfort. These moliminal symptoms are a good marker of prior ovulation, although their absence is less helpful.

Dysmenorrhea *Dysmenorrhea* refers to the crampy lower abdominal midline discomfort that begins with the onset of menstrual bleeding and gradually decreases over the next 12–72 h. It may be associated with nausea, diarrhea, fatigue, and headache and occurs in 60–93% of adolescents, beginning with the establishment of regular ovulatory cycles. Its prevalence decreases after pregnancy and with the use of oral contraceptives.

Primary dysmenorrhea results from increased stores of prostaglandin precursors, which are generated by sequential stimulation of the uterus by estrogen and progesterone. During menstruation, these precursors are converted to prostaglandins, which cause intense uterine contractions, decreased blood flow, and increased peripheral nerve hypersensitivity, resulting in pain.

Secondary dysmenorrhea is caused by underlying pelvic pathology. *Endometriosis* results from the presence of endometrial glands and stroma outside the uterus. These deposits of ectopic endometrium respond to hormonal stimulation and cause dysmenorrhea, which begins several days before menses. Endometriosis also may be associated with painful intercourse, painful bowel movements, and tender nodules in the uterosacral ligament. Fibrosis and adhesions can produce lateral displacement of the cervix. Transvaginal pelvic ultrasound is part of the initial workup and may detect an endometrioma within the ovary, rectovaginal or bladder nodules, or ureteral involvement. The CA125 level may be increased, but it has low negative predictive value. Definitive diagnosis requires laparoscopy. Symptomatology does not always predict the extent of endometriosis. The prevalence is lower in black and Hispanic women than in Caucasians and Asians. *Other secondary causes* of dysmenorrhea include adenomyosis, a condition caused by the presence of ectopic endometrial glands and stroma within the myometrium. Cervical stenosis may result from trauma, infection, or surgery.

TREATMENT DYSMENORRHEA

Local application of heat; dietary dairy intake; use of vitamins B₁, B₆, and E and fish oil; acupuncture; yoga; and exercise are of some benefit for the treatment of dysmenorrhea. Studies of vitamin D₃ are not yet adequate to provide a recommendation. However, nonsteroidal anti-inflammatory drugs (NSAIDs) are the most effective treatment and provide >80% sustained response rates. Ibuprofen, naproxen, ketoprofen, mefenamic acid, and nimesulide are all superior to

placebo. Treatment should be started a day before expected menses and generally is continued for 2–3 days. Oral contraceptives also reduce symptoms of dysmenorrhea. The use of tocolytics, antiphosphodiesterase inhibitors, and magnesium has been suggested, but

there are insufficient data to recommend them. Failure of response to NSAIDs and/or oral contraceptives is suggestive of a pelvic disorder such as endometriosis, and diagnostic laparoscopy should be considered to guide further treatment.

SECTION 9 ALTERATIONS IN THE SKIN

70 Approach to the Patient with a Skin Disorder

Thomas J. Lawley, Kim B. Yancey

The challenge of examining the skin lies in distinguishing normal from abnormal findings, distinguishing significant findings from trivial ones, and integrating pertinent signs and symptoms into an appropriate differential diagnosis. The fact that the largest organ in the body is visible is both an advantage and a disadvantage to those who examine it. It is advantageous because no special instrumentation is necessary and because the skin can be biopsied with little morbidity. However, the casual observer can be misled by a variety of stimuli and overlook important, subtle signs of skin or systemic disease. For instance, the sometimes minor differences in color and shape that distinguish a melanoma (Fig. 70-1) from a benign nevocmelanocytic nevus (Fig. 70-2) can be difficult to recognize. A variety of descriptive terms have been developed that characterize cutaneous lesions (Tables 70-1, 70-2, and Tables 70-3; Fig. 70-3), thereby aiding in their interpretation and in the formulation of a differential diagnosis (Table 70-4). For example, the finding of scaling papules, which are present in psoriasis or atopic dermatitis, places the patient in a different diagnostic category than would hemorrhagic papules, which may indicate vasculitis or sepsis (Figs. 70-4 and 70-5, respectively). It is also important to differentiate primary from secondary skin lesions. If the examiner focuses on linear erosions overlying an area of erythema and scaling, he or she may incorrectly assume that the erosion is the primary lesion and that the redness and scale are secondary, whereas the correct interpretation would be that the patient has a pruritic eczematous dermatitis with erosions caused by scratching.



FIGURE 70-1 Superficial spreading melanoma. This is the most common type of melanoma. Such lesions usually demonstrate asymmetry, border irregularity, color variegation (black, blue, brown, pink, and white), a diameter >6 mm, and a history of change (e.g., an increase in size or development of associated symptoms such as pruritus or pain).



FIGURE 70-2 Nevocmelanocytic nevus. Nevi are benign proliferations of nevocmelanocytes characterized by regularly shaped hyperpigmented macules or papules of a uniform color.

TABLE 70-1 DESCRIPTION OF PRIMARY SKIN LESIONS

Macule: A flat, colored lesion, <2 cm in diameter, not raised above the surface of the surrounding skin. A “freckle,” or ephelid, is a prototypical pigmented macule.
Patch: A large (>2-cm) flat lesion with a color different from the surrounding skin. This differs from a macule only in size.
Papule: A small, solid lesion, <0.5 cm in diameter, raised above the surface of the surrounding skin and thus palpable (e.g., a closed comedone, or white-head, in acne).
Nodule: A larger (0.5- to 5.0-cm), firm lesion raised above the surface of the surrounding skin. This differs from a papule only in size (e.g., a large dermal nevocmelanocytic nevus).
Tumor: A solid, raised growth >5 cm in diameter.
Plaque: A large (>1-cm), flat-topped, raised lesion; edges may either be distinct (e.g., in psoriasis) or gradually blend with surrounding skin (e.g., in eczematous dermatitis).
Vesicle: A small, fluid-filled lesion, <0.5 cm in diameter, raised above the plane of surrounding skin. Fluid is often visible, and the lesions are translucent (e.g., vesicles in allergic contact dermatitis caused by <i>Toxicodendron</i> [poison ivy]).
Pustule: A vesicle filled with leukocytes. Note: The presence of pustules does not necessarily signify the existence of an infection.
Bulla: A fluid-filled, raised, often translucent lesion >0.5 cm in diameter.
Wheal: A raised, erythematous, edematous papule or plaque, usually representing short-lived vasodilation and vasopermeability.
Telangiectasia: A dilated, superficial blood vessel.

TABLE 70-2 DESCRIPTION OF SECONDARY SKIN LESIONS

Lichenification: A distinctive thickening of the skin that is characterized by accentuated skin-fold markings.
Scale: Excessive accumulation of stratum corneum.
Crust: Dried exudate of body fluids that may be either yellow (i.e., serous crust) or red (i.e., hemorrhagic crust).
Erosion: Loss of epidermis without an associated loss of dermis.
Ulcer: Loss of epidermis and at least a portion of the underlying dermis.
Excoriation: Linear, angular erosions that may be covered by crust and are caused by scratching.
Atrophy: An acquired loss of substance. In the skin, this may appear as a depression with intact epidermis (i.e., loss of dermal or subcutaneous tissue) or as sites of shiny, delicate, wrinkled lesions (i.e., epidermal atrophy).
Scar: A change in the skin secondary to trauma or inflammation. Sites may be erythematous, hypopigmented, or hyperpigmented depending on their age or character. Sites on hair-bearing areas may be characterized by destruction of hair follicles.

TABLE 70-3 COMMON DERMATOLOGIC TERMS

Alopecia: Hair loss, partial or complete.
Annular: Ring-shaped.
Cyst: A soft, raised, encapsulated lesion filled with semisolid or liquid contents.
Herpetiform: In a grouped configuration.
Lichenoid eruption: Violaceous to purple, polygonal lesions that resemble those seen in lichen planus.
Milia: Small, firm, white papules filled with keratin.
Morbilliform rash: Generalized, small erythematous macules and/or papules that resemble lesions seen in measles.
Nummular: Coin-shaped.
Poikiloderma: Skin that displays variegated pigmentation, atrophy, and telangiectases.
Polycyclic lesions: A configuration of skin lesions formed from coalescing rings or incomplete rings.
Pruritus: A sensation that elicits the desire to scratch. Pruritus is often the predominant symptom of inflammatory skin diseases (e.g., atopic dermatitis, allergic contact dermatitis); it is also commonly associated with xerosis and aged skin. Systemic conditions that can be associated with pruritus include chronic renal disease, cholestasis, pregnancy, malignancy, thyroid disease, polycythemia vera, and delusions of parasitosis.

APPROACH TO THE PATIENT: Skin Disorder

In examining the skin it is usually advisable to assess the patient before taking an extensive history. This approach ensures that the entire cutaneous surface will be evaluated, and objective findings can be integrated with relevant historical data. Four basic features of a skin lesion must be noted and considered during a physical examination: the *distribution* of the eruption, the *types* of primary and secondary lesions, the *shape* of individual lesions, and the *arrangement* of the lesions. An ideal skin examination includes evaluation of the skin, hair, and nails as well as the mucous membranes of the mouth, eyes, nose, nasopharynx, and anogenital region. In the initial examination, it is important that the patient be disrobed as completely as possible to minimize chances of missing important individual skin lesions and permit accurate assessment of the distribution of the eruption. The patient should first be viewed from a distance of about 1.5–2 m (4–6 ft) so that the general character of the skin and the distribution of lesions can be evaluated. Indeed, the distribution of lesions often correlates highly with diagnosis (Fig. 70-6). For example, a hospitalized patient with a generalized erythematous exanthem is more likely to have a drug eruption than is a patient with a similar rash limited to the sun-exposed portions of the face. Once the distribution of the lesions has been established, the nature of the primary lesion

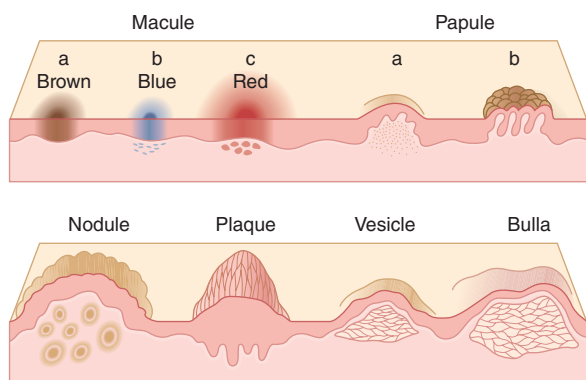


FIGURE 70-3 A schematic representation of several common primary skin lesions (see Table 70-1).

must be determined. Thus, when lesions are distributed on elbows, knees, and scalp, the most likely possibility based solely on distribution is psoriasis or dermatitis herpetiformis (Figs. 70-7 and 70-8, respectively). The primary lesion in psoriasis is a scaly papule that soon forms erythematous plaques covered with a white scale, whereas that of dermatitis herpetiformis is an urticarial papule that quickly becomes a small vesicle. In this manner, identification of the primary lesion directs the examiner toward the proper diagnosis. Secondary changes in skin can also be quite helpful. For example, scale represents excessive epidermis, while crust is the result of a discontinuous epithelial cell layer. Palpation of skin lesions can yield insight into the character of an eruption. Thus, red papules on the lower extremities that blanch with pressure can be a manifestation of many different diseases, but hemorrhagic red papules that do not blanch with pressure indicate palpable purpura characteristic of necrotizing vasculitis (Fig. 70-4).

The shape of lesions is also an important feature. Flat, round, erythematous papules and plaques are common in many cutaneous diseases. However, target-shaped lesions that consist in part of erythematous plaques are specific for erythema multiforme (Fig. 70-9). Likewise, the arrangement of individual lesions is important. Erythematous papules and vesicles can occur in many conditions, but their arrangement in a specific linear array suggests an external etiology such as allergic contact dermatitis (Fig. 70-10) or primary irritant dermatitis. In contrast, lesions with a generalized arrangement are common and suggest a systemic etiology.

As in other branches of medicine, a complete history should be obtained to emphasize the following features:

1. Evolution of lesions
 - a. Site of onset
 - b. Manner in which the eruption progressed or spread
 - c. Duration
 - d. Periods of resolution or improvement in chronic eruptions
2. Symptoms associated with the eruption
 - a. Itching, burning, pain, numbness
 - b. What, if anything, has relieved symptoms
 - c. Time of day when symptoms are most severe
3. Current or recent medications (prescribed as well as over-the-counter)
4. Associated systemic symptoms (e.g., malaise, fever, arthralgias)
5. Ongoing or previous illnesses
6. History of allergies
7. Presence of photosensitivity
8. Review of systems
9. Family history (particularly relevant for patients with melanoma, atopy, psoriasis, or acne)
10. Social, sexual, or travel history

DIAGNOSTIC TECHNIQUES

Many skin diseases can be diagnosed on the basis of gross clinical appearance, but sometimes relatively simple diagnostic procedures can yield valuable information. In most instances, they can be performed at the bedside with a minimum of equipment.

Skin Biopsy A skin biopsy is a straightforward minor surgical procedure; however, it is important to biopsy a lesion that is most likely to yield diagnostic findings. This decision may require expertise in skin diseases and knowledge of superficial anatomic structures in selected areas of the body. In this procedure, a small area of skin is anesthetized with 1% lidocaine with or without epinephrine. The skin lesion in question can be excised or saucerized with a scalpel or removed by punch biopsy. In the latter technique, a punch is pressed against the surface of the skin and rotated with downward pressure until it penetrates to the subcutaneous tissue. The circular biopsy is then lifted with forceps, and the bottom is cut with iris scissors. Biopsy sites may or may not need suture closure, depending on size and location.

TABLE 70-4 SELECTED COMMON DERMATOLOGIC CONDITIONS

Diagnosis	Common Distribution	Usual Morphology	Diagnosis	Common Distribution	Usual Morphology
Acne vulgaris	Face, upper back, chest	Open and closed comedones, erythematous papules, pustules, cysts	Seborrheic keratosis	Trunk, face	Brown plaques with adherent, greasy scale; “stuck on” appearance
Rosacea	Blush area of cheeks, nose, forehead, chin	Erythema, telangiectases, papules, pustules	Folliculitis Impetigo	Any hair-bearing area Anywhere	Follicular pustules Papules, vesicles, pustules, often with honey-colored crusts
Seborrheic dermatitis	Scalp, eyebrows, perinasal areas	Erythema with greasy yellow-brown scale	Herpes simplex	Lips, genitalia	Grouped vesicles progressing to crusted erosions
Atopic dermatitis	Antecubital and popliteal fossae; may be widespread	Patches and plaques of erythema, scaling, and lichenification; pruritus	Herpes zoster	Dermatomal, usually trunk but may be anywhere	Vesicles limited to a dermatome (often painful)
Stasis dermatitis	Ankles, lower legs over medial malleoli	Patches of erythema and scaling on background of hyperpigmentation associated with signs of venous insufficiency	Varicella	Face, trunk, relative sparing of extremities	Lesions arise in crops and quickly progress from erythematous macules, to papules, to vesicles, to pustules, to crusted sites.
Dyshidrotic eczema	Palms, soles, sides of fingers and toes	Deep vesicles	Pityriasis rosea	Trunk (Christmas tree pattern); herald patch followed by multiple smaller lesions	Symmetric erythematous patches with a collarette of scale
Allergic contact dermatitis	Anywhere	Localized erythema, vesicles, scale, and pruritus (e.g., fingers, earlobes—nickel; dorsal aspect of foot—shoe; exposed surfaces—poison ivy)	Tinea versicolor	Chest, back, abdomen, proximal extremities	Scaly hyper- or hypopigmented macules
Psoriasis	Elbows, knees, scalp, lower back, fingernails (may be generalized)	Papules and plaques covered with silvery scale; nails have pits	Candidiasis	Groin, beneath breasts, vagina, oral cavity	Erythematous macerated areas with satellite pustules; white, friable patches on mucous membranes
Lichen planus	Wrists, ankles, mouth (may be widespread)	Violaceous flat-topped papules and plaques	Dermatophytosis	Feet, groin, beard, or scalp	Varies with site, (e.g., tinea corporis—scaly annular plaque)
Keratosis pilaris	Extensor surfaces of arms and thighs, buttocks	Keratotic follicular papules with surrounding erythema	Scabies	Groin, axillae, between fingers and toes, beneath breasts	Excoriated papules, burrows, pruritus
Melasma	Forehead, cheeks, temples, upper lip	Tan to brown patches	Insect bites	Anywhere	Erythematous papules with central puncta
Vitiligo	Periorificial, trunk, extensor surfaces of extremities, flexor wrists, axillae	Chalk-white macules	Cherry angioma Keloid Dermatofibroma	Trunk Anywhere (site of previous injury) Anywhere	Red, blood-filled papules Firm tumor, pink, purple, or brown Firm red to brown nodule that shows dimpling of overlying skin with lateral compression
Actinic keratosis	Sun-exposed areas	Skin-colored or red-brown macule or papule with dry, rough, adherent scale	Acrochordons (skin tags)	Groin, axilla, neck	Fleshy papules
Basal cell carcinoma	Face	Papule with pearly, telangiectatic border on sun-damaged skin	Urticaria	Anywhere	Wheals, sometimes with surrounding flare; pruritus
Squamous cell carcinoma	Face, especially lower lip, ears	Indurated and possibly hyperkeratotic lesions often showing ulceration and/or crusting	Transient acantholytic dermatosis Xerosis	Trunk, especially anterior chest Extensor extremities, especially legs	Erythematous papules Dry, erythematous, scaling patches; pruritus

KOH Preparation A potassium hydroxide (KOH) preparation is performed on scaling skin lesions where a fungal infection is suspected. The edge of such a lesion is scraped gently with a no. 15 scalpel blade. The removed scale is collected on a glass microscope slide and then treated with 1 or 2 drops of a solution of 10–20% KOH. KOH dissolves keratin and allows easier visualization of fungal elements. Brief heating of the slide accelerates dissolution of keratin. When the preparation is viewed under the microscope, the refractile hyphae are seen more easily when the light intensity is reduced and the condenser is lowered. This technique can be used to identify

hyphae in dermatophyte infections, pseudohyphae and budding yeasts in *Candida* infections, and “spaghetti and meatballs” yeast forms in tinea versicolor. The same sampling technique can be used to obtain scale for culture of selected pathogenic organisms.

Tzanck Smear A Tzanck smear is a cytologic technique most often used in the diagnosis of herpesvirus infections (herpes simplex virus [HSV] or varicella zoster virus [VZV]) (see Figs. 217-1 and 217-3). An early vesicle, not a pustule or crusted lesion, is unroofed, and the base of the lesion is scraped gently with a scalpel



FIGURE 70-4 Necrotizing vasculitis. Palpable purpuric papules on the lower legs are seen in this patient with cutaneous small-vessel vasculitis. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 70-5 Meningococemia. An example of fulminant meningococemia with extensive angular purpuric patches. (Courtesy of Stephen E. Gellis, MD; with permission.)

blade. The material is placed on a glass slide, air-dried, and stained with Giemsa or Wright's stain. Multinucleated epithelial giant cells suggest the presence of HSV or VZV; culture, immunofluorescence microscopy, or genetic testing must be performed to identify the specific virus.

Diascopy Diascopy is designed to assess whether a skin lesion will blanch with pressure as, for example, in determining whether a red lesion is hemorrhagic or simply blood-filled. Urticaria (Fig. 70-11) will blanch with pressure, whereas a purpuric lesion caused by necrotizing vasculitis (Fig. 70-4) will not. Diascopy is performed by pressing a microscope slide or magnifying lens against a lesion and noting the amount of blanching that occurs. Granulomas often have an opaque to transparent, brown-pink "apple jelly" appearance on diascopy.

Wood's Light A Wood's lamp generates 360-nm ultraviolet ("black") light that can be used to aid the evaluation of certain skin disorders.

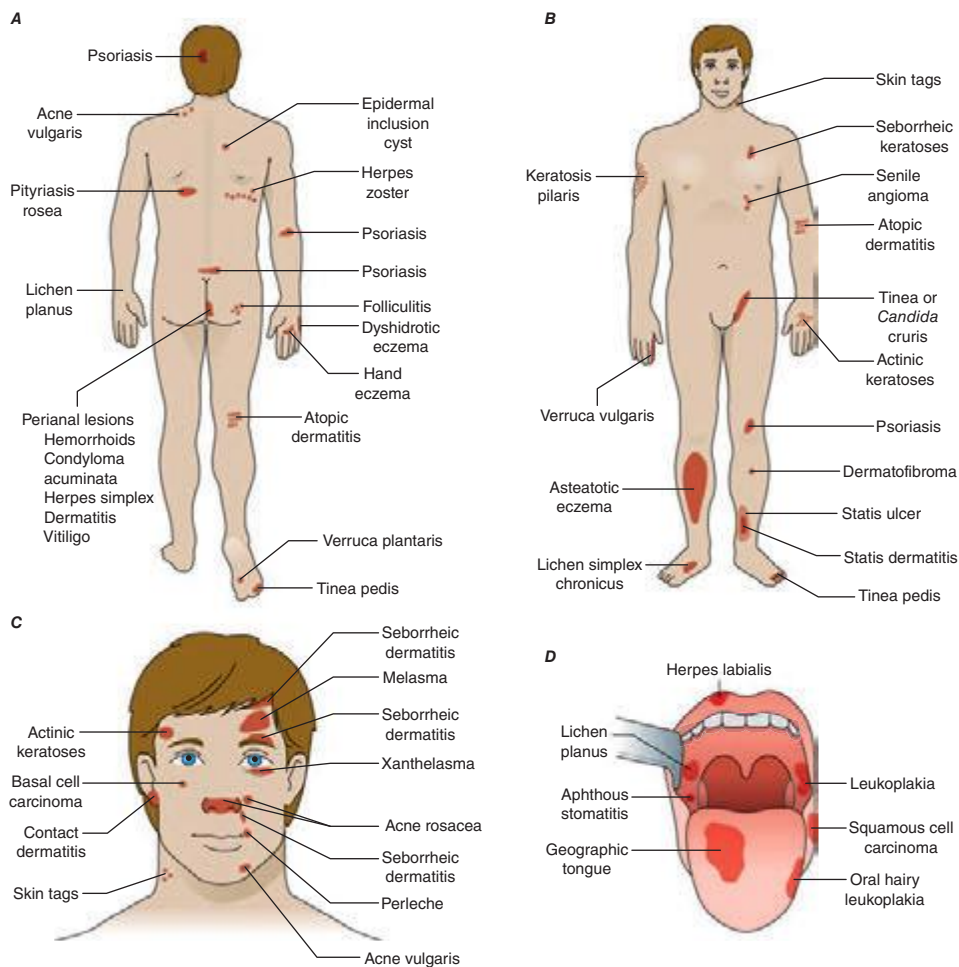


FIGURE 70-6 Distribution of some common dermatologic diseases and lesions.



FIGURE 70-7 Psoriasis. This papulosquamous skin disease is characterized by small and large erythematous papules and plaques with overlying adherent silvery scale.



FIGURE 70-8 Dermatitis herpetiformis. This disorder typically displays pruritic, grouped papulovesicles on elbows, knees, buttocks, and posterior scalp. Vesicles are often excoriated due to associated pruritus.



FIGURE 70-9 Erythema multiforme. This eruption is characterized by multiple erythematous plaques with a target or iris morphology. It usually represents a hypersensitivity reaction to drugs (e.g., sulfonamides) or infections (e.g., HSV). (Courtesy of the Yale Resident's Slide Collection; with permission.)



FIGURE 70-10 Allergic contact dermatitis (ACD). **A.** An example of ACD in its acute phase, with sharply demarcated, weeping, eczematous plaques in a perioral distribution. **B.** ACD in its chronic phase, with an erythematous, lichenified, weeping plaque on skin chronically exposed to nickel in a metal snap. (B, Courtesy of Robert Swerlick, MD; with permission.)

For example, a Wood's lamp will cause erythrasma (a superficial, intertriginous infection caused by *Corynebacterium minutissimum*) to show a characteristic coral pink color, and wounds colonized by *Pseudomonas* will appear pale blue. Tinea capitis caused by certain dermatophytes (e.g., *Microsporum canis* or *M. audouinii*) exhibits a yellow fluorescence. Pigmented lesions of the epidermis such as freckles are accentuated, while dermal pigment such as postinflammatory hyperpigmentation fades under a Wood's light. Vitiligo (Fig. 70-12) appears



FIGURE 70-11 Urticaria. Discrete and confluent, edematous, erythematous papules and plaques are characteristic of this whealing eruption.



FIGURE 70-12 Vitiligo. Characteristic lesions display an acral distribution and striking depigmentation as a result of loss of melanocytes.

totally white under a Wood's lamp, and previously unsuspected areas of involvement often become apparent. A Wood's lamp may also aid in the demonstration of tinea versicolor and in recognition of ash leaf spots in patients with tuberous sclerosis.

Patch Tests Patch testing is designed to document sensitivity to a specific antigen. In this procedure, a battery of suspected allergens is applied to the patient's back under occlusive dressings and allowed to remain in contact with the skin for 48 h. The dressings are removed, and the area is examined for evidence of delayed hypersensitivity reactions (e.g., erythema, edema, or papulovesicles). This test is best performed by physicians with special expertise in patch testing and is often helpful in the evaluation of patients with chronic dermatitis.

71 Eczema, Psoriasis, Cutaneous Infections, Acne, and Other Common Skin Disorders

Leslie P. Lawley, Calvin O. McCall, Thomas J. Lawley

ECZEMA AND DERMATITIS

Eczema is a type of dermatitis, and these terms are often used synonymously (e.g., atopic eczema or atopic dermatitis [AD]). Eczema is a reaction pattern that presents with variable clinical findings and the common histologic finding of *spongiosis* (intercellular edema of the epidermis). Eczema is the final common expression for a number of disorders, including those discussed in the following sections. Primary lesions may include erythematous macules, papules, and vesicles, which can coalesce to form patches and plaques. In severe eczema, secondary lesions from infection or excoriation, marked by weeping and crusting, may predominate. In chronic eczematous conditions, *lichenification* (cutaneous hypertrophy and accentuation of normal skin markings) may alter the characteristic appearance of eczema.

ATOPIC DERMATITIS

AD is the cutaneous expression of the atopic state, characterized by a family history of asthma, allergic rhinitis, or eczema. The prevalence of AD is increasing worldwide. Some of its features are shown in [Table 71-1](#).


 The etiology of AD is only partially defined, but there is a clear genetic predisposition. When both parents are affected by AD, >80% of their children manifest the disease. When only one parent is affected, the prevalence drops to slightly over 50%. A characteristic defect in AD that contributes to the pathophysiology is an

TABLE 71-1 CLINICAL FEATURES OF ATOPIC DERMATITIS

1. Pruritus and scratching
2. Course marked by exacerbations and remissions
3. Lesions typical of eczematous dermatitis
4. Personal or family history of atopy (asthma, allergic rhinitis, food allergies, or eczema)
5. Clinical course lasting >6 weeks
6. Lichenification of skin

impaired epidermal barrier. In many patients, a mutation in the gene encoding filaggrin, a structural protein in the stratum corneum, is responsible. Patients with AD may display a variety of immunoregulatory abnormalities, including increased IgE synthesis; increased serum IgE levels; and impaired, delayed-type hypersensitivity reactions.

The clinical presentation often varies with age. Half of patients with AD present within the first year of life, and 80% present by 5 years of age. About 80% ultimately coexpress allergic rhinitis or asthma. The infantile pattern is characterized by weeping inflammatory patches and crusted plaques on the face, neck, and extensor surfaces. The childhood and adolescent pattern is typified by dermatitis of flexural skin, particularly in the antecubital and popliteal fossae ([Fig. 71-1](#)). AD may resolve spontaneously, but approximately 40% of all individuals affected as children will have dermatitis in adult life. The distribution of lesions in adults may be similar to those seen in childhood; however, adults frequently have localized disease manifesting as lichen simplex chronicus or hand eczema (see below). In patients with localized disease, AD may be suspected because of a typical personal or family history or the presence of cutaneous stigmata of AD such as perioral pallor, an extra fold of skin beneath the lower eyelid (Dennie-Morgan folds), increased palmar skin markings, and an increased incidence of cutaneous infections, particularly with *Staphylococcus aureus*. Regardless of other manifestations, pruritus is a prominent characteristic of AD in all age groups and is exacerbated by dry skin. Many of the cutaneous findings in affected patients, such as lichenification, are secondary to rubbing and scratching.

TREATMENT ATOPIC DERMATITIS

Therapy for AD should include avoidance of cutaneous irritants, adequate moisturizing through the application of emollients, judicious use of topical anti-inflammatory agents, and prompt treatment of secondary infection. Patients should be instructed to bathe no more often than daily, using warm or cool water, and to use only mild bath soap. Immediately after bathing, while the skin is still moist, a topical anti-inflammatory agent in a cream or ointment base should be applied to areas of dermatitis, and all other skin



FIGURE 71-1 Atopic dermatitis. Hyperpigmentation, lichenification, and scaling in the antecubital fossae are seen in this patient with atopic dermatitis. (Courtesy of Robert Swerlick, MD; with permission.)

areas should be lubricated with a moisturizer. Approximately 30 g of a topical agent is required to cover the entire body surface of an average adult.

Low- to mid-potency topical glucocorticoids are employed in most treatment regimens for AD. Skin atrophy and the potential for systemic absorption are constant concerns, especially with more potent agents. Low-potency topical glucocorticoids or nonglucocorticoid anti-inflammatory agents should be selected for use on the face and in intertriginous areas to minimize the risk of skin atrophy. Two nonglucocorticoid anti-inflammatory agents are available: tacrolimus ointment and pimecrolimus cream. These agents are macrolide immunosuppressants that are approved by the U.S. Food and Drug Administration (FDA) for topical use in AD. Reports of broader effectiveness appear in the literature. These agents do not cause skin atrophy, nor do they suppress the hypothalamic-pituitary-adrenal axis. However, concerns have emerged regarding the potential for lymphomas in patients treated with these agents. Thus, caution should be exercised when these agents are considered. Currently, they are also more costly than topical glucocorticoids. Barrier-repair products that attempt to restore the impaired epidermal barrier are also nonglucocorticoid agents and are gaining popularity in the treatment of AD.

Secondary infection of eczematous skin may lead to exacerbation of AD. Crusted and weeping skin lesions may be infected with *S. aureus*. When secondary infection is suspected, eczematous lesions should be cultured and patients treated with systemic antibiotics active against *S. aureus*. The initial use of penicillinase-resistant penicillins or cephalosporins is preferable. Dicloxacillin or cephalexin (250 mg qid for 7–10 days) is generally adequate for adults; however, antibiotic selection must be directed by culture results and clinical response. More than 50% of *S. aureus* isolates are now methicillin resistant in some communities. Current recommendations for the treatment of infection with these community-acquired methicillin-resistant *S. aureus* (CA-MRSA) strains in adults include trimethoprim-sulfamethoxazole (1 double-strength tablet bid), minocycline (100 mg bid), doxycycline (100 mg bid), or clindamycin (300–450 mg qid). Duration of therapy should be 7–10 days. Inducible resistance may limit clindamycin's usefulness. Such resistance can be detected by the double-disk diffusion test, which should be ordered if the isolate is erythromycin resistant and clindamycin sensitive. As an adjunct, antibacterial washes or dilute sodium hypochlorite baths (0.005% bleach) and intermittent nasal mupirocin may be useful.

Control of pruritus is essential for treatment, because AD often represents "an itch that rashes." Antihistamines are most often used to control pruritus. Diphenhydramine (25 mg every 4–6 h), hydroxyzine (10–25 mg every 6 h), or doxepin (10–25 mg at bedtime) are useful primarily due to their sedating action. Higher doses of these agents may be required, but sedation can become bothersome. Patients need to be counseled about driving or operating heavy equipment after taking these medications. When used at bedtime, sedating antihistamines may improve the patient's sleep. Although they are effective in urticaria, non-sedating antihistamines and selective H₂ blockers are of little use in controlling the pruritus of AD.

Treatment with systemic glucocorticoids should be limited to severe exacerbations unresponsive to topical therapy. In the patient with chronic AD, therapy with systemic glucocorticoids will generally clear the skin only briefly, and cessation of the systemic therapy will invariably be accompanied by a return, if not a worsening, of the dermatitis. Patients who do not respond to conventional therapies should be considered for patch testing to rule out allergic contact dermatitis (ACD). The role of dietary allergens in AD is controversial, and there is little evidence that they play any role outside of infancy, during which a small percentage of patients with AD may be affected by food allergens.

LICHEN SIMPLEX CHRONICUS

Lichen simplex chronicus may represent the end stage of a variety of pruritic and eczematous disorders, including AD. It consists of a

circumscribed plaque or plaques of lichenified skin due to chronic scratching or rubbing. Common areas involved include the posterior nuchal region, dorsum of the feet, and ankles. Treatment of lichen simplex chronicus centers on breaking the cycle of chronic itching and scratching. High-potency topical glucocorticoids are helpful in most cases, but, in recalcitrant cases, application of topical glucocorticoids under occlusion, or intralesional injection of glucocorticoids may be required.

CONTACT DERMATITIS

Contact dermatitis is an inflammatory skin process caused by an exogenous agent or agents that directly or indirectly injure the skin. In *irritant* contact dermatitis (ICD), this injury is caused by an inherent characteristic of a compound—for example, a concentrated acid or base. Agents that cause ACD induce an antigen-specific immune response (e.g., poison ivy dermatitis). The clinical lesions of contact dermatitis may be acute (wet and edematous) or chronic (dry, thickened, and scaly), depending on the persistence of the insult (see Fig. 70-10).

Irritant Contact Dermatitis ICD is generally well demarcated and often localized to areas of thin skin (eyelids, intertriginous areas) or to areas where the irritant was occluded. Lesions may range from minimal skin erythema to areas of marked edema, vesicles, and ulcers. Prior exposure to the offending agent is not necessary, and the reaction develops in minutes to a few hours. Chronic low-grade irritant dermatitis is the most common type of ICD, and the most common area of involvement is the hands (see below). The most common irritants encountered are chronic wet work, soaps, and detergents. Treatment should be directed toward the avoidance of irritants and the use of protective gloves or clothing.

Allergic Contact Dermatitis ACD is a manifestation of delayed-type hypersensitivity mediated by memory T lymphocytes in the skin. Prior exposure to the offending agent is necessary to develop the hypersensitivity reaction, which may take as little as 12 h or as much as 72 h to develop. The most common cause of ACD is exposure to plants, especially to members of the family Anacardiaceae, including the genus *Toxicodendron*. Poison ivy, poison oak, and poison sumac are members of this genus and cause an allergic reaction marked by erythema, vesiculation, and severe pruritus. The eruption is often linear or angular, corresponding to areas where plants have touched the skin. The sensitizing antigen common to these plants is urushiol, an oleoresin containing the active ingredient pentadecylcatechol. The oleoresin may adhere to skin, clothing, tools, and pets, and contaminated articles may cause dermatitis even after prolonged storage. Blister fluid does not contain urushiol and is not capable of inducing skin eruption in exposed subjects.

TREATMENT CONTACT DERMATITIS

If contact dermatitis is suspected and an offending agent is identified and removed, the eruption will resolve. Usually, treatment with high-potency topical glucocorticoids is enough to relieve symptoms while the dermatitis runs its course. For those patients who require systemic therapy, daily oral prednisone—beginning at 1 mg/kg, but usually ≤60 mg/d—is sufficient. The dose should be tapered over 2–3 weeks, and each daily dose should be taken in the morning with food.

Identification of a contact allergen can be a difficult and time-consuming task. Allergic contact dermatitis should be suspected in patients with dermatitis unresponsive to conventional therapy or with an unusual and patterned distribution. Patients should be questioned carefully regarding occupational exposures and topical preparations, nickel sulfate, potassium dichromate, thimerosal, neomycin sulfate, fragrances, formaldehyde, and rubber-curing agents. Patch testing is helpful in identifying these agents but should not be attempted when patients have widespread active dermatitis or are taking systemic glucocorticoids.



FIGURE 71-2 Dyshidrotic eczema. This example is characterized by deep-seated vesicles and scaling on palms and lateral fingers, and the disease is often associated with an atopic diathesis.

HAND ECZEMA

Hand eczema is a very common, chronic skin disorder in which both exogenous and endogenous factors play important roles. It may be associated with other cutaneous disorders such as AD, and contact with various agents may be involved. Hand eczema represents a large proportion of cases of occupation-associated skin disease. Chronic, excessive exposure to water and detergents, harsh chemicals, or allergens may initiate or aggravate this disorder. It may present with dryness and cracking of the skin of the hands as well as with variable amounts of erythema and edema. Often, the dermatitis will begin under rings, where water and irritants are trapped. *Dyshidrotic eczema*, a variant of hand eczema, presents with multiple, intensely pruritic, small papules and vesicles on the thenar and hypothenar eminences and the sides of the fingers (Fig. 71-2). Lesions tend to occur in crops that slowly form crusts and then heal.

The evaluation of a patient with hand eczema should include an assessment of potential occupation-associated exposures. The history should be directed to identifying possible irritant or allergen exposures.

TREATMENT HAND ECZEMA

Therapy for hand eczema is directed toward avoidance of irritants, identification of possible contact allergens, treatment of coexistent infection, and application of topical glucocorticoids. Whenever possible, the hands should be protected by gloves, preferably vinyl. The use of rubber gloves (latex) to protect dermatitic skin is sometimes associated with the development of hypersensitivity reactions to components of the gloves. Patients can be treated with cool moist compresses followed by application of a mid- to high-potency topical glucocorticoid in a cream or ointment base. As in AD, treatment of secondary infection is essential for good control. In addition, patients with hand eczema should be examined for dermatophyte infection by KOH preparation and culture (see below).

NUMMULAR ECZEMA

Nummular eczema is characterized by circular or oval “coinlike” lesions, beginning as small edematous papules that become crusted and scaly. The etiology of nummular eczema is unknown, but dry skin is a contributing factor. Common locations are the trunk or the extensor surfaces of the extremities, particularly on the pretibial areas or dorsum of the hands. Nummular eczema occurs more frequently in men and is most common in middle age. The treatment of nummular eczema is similar to that for AD.

ASTEATOTIC ECZEMA

Asteatotic eczema, also known as *xerotic eczema* or “winter itch,” is a mildly inflammatory dermatitis that develops in areas of extremely dry skin, especially during the dry winter months. Clinically, there may be considerable overlap with nummular eczema. This form of eczema accounts for a large number of physician visits because of the associated pruritus. Fine cracks and scale, with or without erythema, characteristically develop in areas of dry skin, especially on the anterior surfaces of the lower extremities in elderly patients. Asteatotic eczema responds well to topical moisturizers and the avoidance of cutaneous irritants. Overbathing and the use of harsh soaps exacerbate asteatotic eczema.

STASIS DERMATITIS AND STASIS ULCERATION

Stasis dermatitis develops on the lower extremities secondary to venous incompetence and chronic edema. Patients may give a history of deep venous thrombosis and may have evidence of vein removal or varicose veins. Early findings in stasis dermatitis consist of mild erythema and scaling associated with pruritus. The typical initial site of involvement is the medial aspect of the ankle, often over a distended vein (Fig. 71-3).

Stasis dermatitis may become acutely inflamed, with crusting and exudate. In this state, it is easily confused with cellulitis. Chronic stasis dermatitis is often associated with dermal fibrosis that is recognized clinically as brawny edema of the skin. As the disorder progresses, the dermatitis becomes progressively pigmented due to chronic erythrocyte extravasation leading to cutaneous hemosiderin deposition. Stasis dermatitis may be complicated by secondary infection and contact dermatitis. Severe stasis dermatitis may precede the development of stasis ulcers.

TREATMENT STASIS DERMATITIS AND STASIS ULCERATION

Patients with stasis dermatitis and stasis ulceration benefit greatly from leg elevation and the routine use of compression stockings with a gradient of at least 30–40 mmHg. Stockings providing less compression, such as antiembolism hose, are poor substitutes. Use of emollients and/or mid-potency topical glucocorticoids and avoidance of irritants are also helpful in treating stasis dermatitis. Protection of the legs from injury, including scratching, and control of chronic edema are essential to prevent ulcers. Diuretics may be required to adequately control chronic edema.

Stasis ulcers are difficult to treat, and resolution is slow. It is extremely important to elevate the affected limb as much as possible. The ulcer should be kept clear of necrotic material by gentle debridement and covered with a semipermeable dressing and a compression dressing or compression stocking. Glucocorticoids



FIGURE 71-3 Stasis dermatitis. An example of stasis dermatitis showing erythematous, scaly, and oozing patches over the lower leg. Several stasis ulcers are also seen in this patient.



FIGURE 71-4 Seborrheic dermatitis. Central facial erythema with overlying greasy, yellowish scale is seen in this patient. (Courtesy of Jean Bologna, MD; with permission.)

should not be applied to ulcers, because they may retard healing; however, they may be applied to the surrounding skin to control itching, scratching, and additional trauma. Secondarily infected lesions should be treated with appropriate oral antibiotics, but it should be noted that all ulcers will become colonized with bacteria, and the purpose of antibiotic therapy should not be to clear all bacterial growth. Care must be taken to exclude treatable causes of leg ulcers (hypercoagulation, vasculitis) before beginning the chronic management outlined above.

SEBORRHEIC DERMATITIS

Seborrheic dermatitis is a common, chronic disorder characterized by greasy scales overlying erythematous patches or plaques. Induration and scale are generally less prominent than in psoriasis, but clinical overlap exists between these diseases (“sebopsoriasis”). The most common location is in the scalp, where it may be recognized as severe dandruff. On the face, seborrheic dermatitis affects the eyebrows, eyelids, glabella, and nasolabial folds (Fig. 71-4). Scaling of the external auditory canal is common in seborrheic dermatitis. In addition, the postauricular areas often become macerated and tender. Seborrheic dermatitis may also develop in the central chest, axilla, groin, submammary folds, and gluteal cleft. Rarely, it may cause widespread generalized dermatitis. Pruritus is variable.

Seborrheic dermatitis may be evident within the first few weeks of life, and within this context it typically occurs in the scalp (“cradle cap”), face, or groin. It is rarely seen in children beyond infancy but becomes evident again during adult life. Although it is frequently seen in patients with Parkinson’s disease, in those who have had cerebrovascular accidents, and in those with HIV infection, the overwhelming majority of individuals with seborrheic dermatitis have no underlying disorder.

TREATMENT SEBORRHEIC DERMATITIS

Treatment with low-potency topical glucocorticoids in conjunction with a topical antifungal agent, such as ketoconazole cream or ciclopirox cream, is often effective. The scalp and beard areas may benefit from antidandruff shampoos, which should be left in place 3–5 min before rinsing. High-potency topical glucocorticoid solutions (betamethasone or clobetasol) are effective for control of severe scalp involvement. High-potency glucocorticoids should not be used on the face because this treatment is often associated with steroid-induced rosacea or atrophy.

PAPULOSQUAMOUS DISORDERS (TABLE 71-2)

PSORIASIS

Psoriasis is one of the most common dermatologic diseases, affecting up to 2% of the world’s population. It is an immune-mediated disease clinically characterized by erythematous, sharply demarcated papules and rounded plaques covered by silvery micaceous scale. The skin lesions of psoriasis are variably pruritic. Traumatized areas often develop lesions of psoriasis (the *Koebner* or isomorphic phenomenon). In addition, other external factors may exacerbate psoriasis, including infections, stress, and medications (lithium, beta blockers, and anti-malarial drugs).

The most common variety of psoriasis is called *plaque-type*. Patients with plaque-type psoriasis have stable, slowly enlarging plaques, which remain basically unchanged for long periods of time. The most commonly involved areas are the elbows, knees, gluteal cleft, and scalp. Involvement tends to be symmetric. Plaque psoriasis generally develops slowly and runs an indolent course. It rarely remits spontaneously. *Inverse psoriasis* affects the intertriginous regions, including the axilla, groin, submammary region, and navel; it also tends to affect the scalp, palms, and soles. The individual lesions are sharply demarcated plaques (see Fig. 70-7), but they may be moist and without scale due to their locations.

Guttate psoriasis (eruptive psoriasis) is most common in children and young adults. It develops acutely in individuals without psoriasis or in those with chronic plaque psoriasis. Patients present with many small erythematous, scaling papules, frequently after upper respiratory tract infection with β -hemolytic streptococci. The differential diagnosis should include pityriasis rosea and secondary syphilis.

In *pustular psoriasis*, patients may have disease localized to the palms and soles, or the disease may be generalized. Regardless of the extent of disease, the skin is erythematous, with pustules and variable scale. Localized to the palms and soles, it is easily confused with eczema. When it is generalized, episodes are characterized by fever (39° – 40° C [102.2° – 104.0° F]) lasting several days, an accompanying generalized eruption of sterile pustules, and a background of intense erythema; patients may become erythrodermic. Episodes of fever and pustules are recurrent. Local irritants, pregnancy, medications, infections, and systemic glucocorticoid withdrawal can precipitate this form of psoriasis. Oral retinoids are the treatment of choice in nonpregnant patients.

TABLE 71-2 PAPULOSQUAMOUS DISORDERS

	Clinical Features	Other Notable Features	Histologic Features
Psoriasis	Sharply demarcated, erythematous plaques with mica-like scale; predominantly on elbows, knees, and scalp; atypical forms may localize to intertriginous areas; eruptive forms may be associated with infection	May be aggravated by certain drugs, infection; severe forms seen in association with HIV	Acanthosis, vascular proliferation
Lichen planus	Purple polygonal papules marked by severe pruritus; lacy white markings, especially associated with mucous membrane lesions	Certain drugs may induce: thiazides, antimalarial drugs	Interface dermatitis
Pityriasis rosea	Rash often preceded by herald patch; oval to round plaques with trailing scale; most often affects trunk; eruption lines up in skinfolds giving a “fir tree–like” appearance; generally spares palms and soles	Variable pruritus; self-limited, resolving in 2–8 weeks; may be imitated by secondary syphilis	Pathologic features often nonspecific
Dermatophytosis	Polymorphous appearance depending on dermatophyte, body site, and host response; sharply defined to ill-demarcated scaly plaques with or without inflammation; may be associated with hair loss	KOH preparation may show branching hyphae; culture helpful	Hyphae and neutrophils in stratum corneum

TABLE 71-3 FDA-APPROVED SYSTEMIC THERAPY FOR PSORIASIS

Agent	Medication Class	Administration		Adverse Events (Selected)
		Route	Frequency	
Methotrexate	Antimetabolite	Oral	Weekly	Hepatotoxicity, pulmonary toxicity, pancytopenia, potential for increased malignancies, ulcerative stomatitis, nausea, diarrhea, teratogenicity
Acitretin	Retinoid	Oral	Daily	Teratogenicity, hepatotoxicity, hyperostosis, hyperlipidemia/pancreatitis, depression, ophthalmologic effects, pseudotumor cerebri
Cyclosporine	Calcineurin inhibitor	Oral	Twice daily	Renal dysfunction, hypertension, hyperkalemia, hyperuricemia, hypomagnesemia, hyperlipidemia, increased risk of malignancies

Fingernail involvement, appearing as punctate pitting, onycholysis, nail thickening, or subungual hyperkeratosis, may be a clue to the diagnosis of psoriasis when the clinical presentation is not classic.

According to the National Psoriasis Foundation, up to 30% of patients with psoriasis have psoriatic arthritis (PsA). There are five subtypes of PsA: symmetric, asymmetric, distal interphalangeal predominant (DIP), spondylitis, and arthritis mutilans. Symmetric arthritis resembles rheumatoid arthritis, but is usually milder. Asymmetric arthritis can involve any joint and may present as “sausage digits.” DIP is the classic form, but occurs in only about 5% of patients with PsA. It may involve fingers and toes. Spondylitis also occurs in about 5% of patients with PsA. Arthritis mutilans is severe and deforming. It affects primarily the small joints of the hands and feet. It accounts for fewer than 5% of PsA cases.

An increased risk of metabolic syndrome, including increased morbidity and mortality from cardiovascular events, has been demonstrated in psoriasis patients. Appropriate screening tests should be performed. The etiology of psoriasis is still poorly understood, but there is clearly a genetic component to the disease. In various studies, 30–50% of patients with psoriasis report a positive family history. Psoriatic lesions contain infiltrates of activated T cells that are thought to elaborate cytokines responsible for keratinocyte hyperproliferation, which results in the characteristic clinical findings. Agents inhibiting T cell activation, clonal expansion, or release of proinflammatory cytokines are often effective for the treatment of severe psoriasis (see below).

TREATMENT PSORIASIS

Treatment of psoriasis depends on the type, location, and extent of disease. All patients should be instructed to avoid excess drying or irritation of their skin and to maintain adequate cutaneous hydration. Most cases of localized, plaque-type psoriasis can be managed with mid-potency topical glucocorticoids, although their long-term use is often accompanied by loss of effectiveness (tachyphylaxis) and atrophy of the skin. A topical vitamin D analogue (calcipotriene) and a retinoid (tazarotene) are also efficacious in the treatment of limited psoriasis and have largely replaced other topical agents such as coal tar, salicylic acid, and anthralin.

Ultraviolet (UV) light, natural or artificial, is an effective therapy for many patients with widespread psoriasis. Ultraviolet B (UVB), narrowband UVB, and ultraviolet A (UVA) light with either oral or topical psoralens (PUVA) is used clinically. UV light’s immunosuppressive properties are thought to be responsible for its therapeutic activity in psoriasis. It is also mutagenic, potentially leading to an increased incidence of nonmelanoma and melanoma skin cancer. UV-light therapy is contraindicated in patients receiving cyclosporine and should be used with great care in all immunocompromised patients due to the increased risk of skin cancer.

Various systemic agents can be used for severe, widespread psoriatic disease (Table 71-3). Oral glucocorticoids should not be used for the treatment of psoriasis due to the potential for development of life-threatening pustular psoriasis when therapy is discontinued. Methotrexate is an effective agent, especially in patients with psoriatic arthritis. The synthetic retinoid acitretin is useful, especially when immunosuppression must be avoided; however, teratogenicity limits its use.

The evidence implicating psoriasis as a T cell–mediated disorder has directed therapeutic efforts to immunoregulation. Cyclosporine and other immunosuppressive agents can be very effective in the treatment of psoriasis, and much attention is currently directed toward the development of biologic agents with more selective immunosuppressive properties and better safety profiles (Table 71-4). Experience with these biologic agents is limited, and information regarding combination therapy and adverse events continues to emerge. Use of tumor necrosis factor (TNF- α) inhibitors may worsen congestive heart failure (CHF), and they should be used with caution in patients at risk for or known to have CHF. Further, none of the immunosuppressive agents used in the treatment of psoriasis should be initiated if the patient has a severe infection; patients on such therapy should be routinely screened for tuberculosis. There have been reports of progressive multifocal leukoencephalopathy in association with treatment with the TNF- α inhibitors. Malignancies, including a risk or history of certain malignancies, may limit the use of these systemic agents.

TABLE 71-4 BIOLOGICS APPROVED FOR PSORIASIS OR PSORIATIC ARTHRITIS

Agent	Mechanism of Action	Administration			Warnings
		Indication	Route	Frequency	
Etanercept	Anti-TNF- α	Ps, PsA	SC	Once or twice weekly	Serious infections, hepatotoxicity, CHF, hematologic events, hypersensitivity reactions, neurologic events, potential for increased malignancies
Adalimumab	Anti-TNF- α	Ps, PsA	SC	Every other week	Serious infections, hepatotoxicity, CHF, hematologic events, hypersensitivity reactions, neurologic events, potential for increased malignancies
Infliximab	Anti-TNF- α	Ps, PsA	IV	Initial infusion followed by infusions at weeks 2 and 6, then every 8 weeks	Serious infections, hepatotoxicity, CHF, hematologic events, hypersensitivity reactions, neurologic events, potential for increased malignancies
Golimumab	Anti-TNF- α	PsA	SC	Monthly	Serious infections, hepatotoxicity, CHF, hypersensitivity reactions, neurologic events, potential for increased malignancies
Ustekinumab	Anti-IL-12 and anti-IL-23	Ps	SC	2 doses 4 weeks apart, then every 12 weeks	Serious infections, neurologic events, potential for increased malignancies

Abbreviations: CHF, congestive heart failure; IL, interleukin; IM, intramuscular; Ps, psoriasis; PsA, psoriatic arthritis; SC, subcutaneous; TNF, tumor necrosis factor.



FIGURE 71-5 Lichen planus. An example of lichen planus showing multiple flat-topped, violaceous papules and plaques. Nail dystrophy, as seen in this patient's thumbnail, may also be a feature. (Courtesy of Robert Swerlick, MD; with permission.)

LICHEN PLANUS

Lichen planus (LP) is a papulosquamous disorder that may affect the skin, scalp, nails, and mucous membranes. The primary cutaneous lesions are pruritic, polygonal, flat-topped, violaceous papules. Close examination of the surface of these papules often reveals a network of gray lines (*Wickham's striae*). The skin lesions may occur anywhere but have a predilection for the wrists, shins, lower back, and genitalia (Fig. 71-5). Involvement of the scalp (*lichen planopilaris*) may lead to scarring alopecia, and nail involvement may lead to permanent deformity or loss of fingernails and toenails. LP commonly involves mucous membranes, particularly the buccal mucosa, where it can present on a spectrum ranging from a mild, white, reticulate eruption of the mucosa to a severe, erosive stomatitis. Erosive stomatitis may persist for years and may be linked to an increased risk of oral squamous cell carcinoma. Cutaneous eruptions clinically resembling LP have been observed after administration of numerous drugs, including thiazide diuretics, gold, antimalarial agents, penicillamine, and phenothiazines, and in patients with skin lesions of chronic graft-versus-host disease. In addition, LP may be associated with hepatitis C infection. The course of LP is variable, but most patients have spontaneous remissions 6 months to 2 years after the onset of disease. Topical glucocorticoids are the mainstay of therapy.

PITYRIASIS ROSEA

Pityriasis rosea (PR) is a papulosquamous eruption of unknown etiology occurring more commonly in the spring and fall. Its first manifestation is the development of a 2- to 6-cm annular lesion (the herald patch). This is followed in a few days to a few weeks by the appearance of many smaller annular or papular lesions with a predilection to occur on the trunk (Fig. 71-6). The lesions are generally oval, with their long axis parallel to the skinfold lines. Individual lesions may range in color from red to brown and have a trailing scale. PR shares many clinical features with the eruption of secondary syphilis, but palm and sole lesions are extremely rare in PR and common in secondary syphilis. The eruption tends to be moderately pruritic and lasts 3–8 weeks. Treatment is directed at alleviating pruritus and consists of oral antihistamines; mid-potency topical glucocorticoids; and, in some cases, UVB phototherapy.

CUTANEOUS INFECTIONS (TABLE 71-5)

IMPETIGO, ECTHYMA, AND FURUNCULOSIS

Impetigo is a common superficial bacterial infection of skin caused most often by *S. aureus* (Chap. 172) and in some cases by group A β -hemolytic streptococci (Chap. 173). The primary lesion is a superficial pustule that ruptures and forms a characteristic yellow-brown



FIGURE 71-6 Pityriasis rosea. In this patient with pityriasis rosea, multiple round to oval erythematous patches with fine central scale are distributed along the skin tension lines on the trunk.

honey-colored crust (see Fig. 173-3). Lesions may occur on normal skin (primary infection) or in areas already affected by another skin disease (secondary infection). Lesions caused by staphylococci may be tense, clear bullae, and this less common form of the disease is called *bullous impetigo*. Blisters are caused by the production of exfoliative toxin by *S. aureus* phage type II. This is the same toxin responsible for staphylococcal scalded-skin syndrome, often resulting in dramatic loss of the superficial epidermis due to blistering. The latter syndrome is much more common in children than in adults; however, it should be considered along with toxic epidermal necrolysis and severe drug eruptions in patients with widespread blistering of the skin. *Ecthyma* is a deep non-bullous variant of impetigo that causes punched-out ulcerative lesions. It is more often caused by a primary or secondary infection with *Streptococcus pyogenes*. *Ecthyma* is a deeper infection than typical impetigo and resolves with scars. Treatment of both *ecthyma* and *impetigo* involves gentle debridement of adherent crusts, which is facilitated by the use of soaks and topical antibiotics in conjunction with appropriate oral antibiotics.

Furunculosis is also caused by *S. aureus*, and this disorder has gained prominence in the last decade because of CA-MRSA. A furuncle, or boil, is a painful, erythematous nodule that can occur on any cutaneous surface. The lesions may be solitary but are most often multiple. Patients frequently believe they have been bitten by spiders or insects. Family members or close contacts may also be affected. Furuncles can rupture and drain spontaneously or may need incision and drainage, which may be adequate therapy for small solitary furuncles without cellulitis or systemic symptoms. Whenever possible, lesional material should be sent for culture. Current recommendations for methicillin-sensitive infections are β -lactam antibiotics. Therapy for CA-MRSA was discussed previously (see "Atopic Dermatitis"). Warm compresses and nasal mupirocin are helpful therapeutic additions. Severe infections may require IV antibiotics.

ERYSIPELAS AND CELLULITIS

See Chap. 156.

DERMATOPHYTOSIS

Dermatophytes are fungi that infect skin, hair, and nails and include members of the genera *Trichophyton*, *Microsporum*, and *Epidermophyton* (Chap. 243). *Tinea corporis*, or infection of the relatively hairless skin of the body (glabrous skin), may have a variable appearance depending on the extent of the associated inflammatory reaction. Typical infections consist of erythematous, scaly plaques, with an annular appearance that accounts for the common name "ringworm." Deep inflammatory nodules or granulomas occur in

TABLE 71-5 COMMON SKIN INFECTIONS

	Clinical Features	Etiologic Agent	Treatment
Impetigo	Honey-colored crusted papules, plaques, or bullae	Group A <i>Streptococcus</i> and <i>Staphylococcus aureus</i>	Systemic or topical antistaphylococcal and antistreptococcal antibiotics
Dermatophytosis	Inflammatory or noninflammatory annular scaly plaques; may involve hair loss; groin involvement spares scrotum; hyphae on KOH preparation	<i>Trichophyton</i> , <i>Epidermophyton</i> , or <i>Microsporum</i> spp.	Topical azoles, systemic griseofulvin, terbinafine, or azoles
Candidiasis	Inflammatory papules and plaques with satellite pustules, frequently in intertriginous areas; may involve scrotum; pseudohyphae on KOH preparation	<i>Candida albicans</i> and other <i>Candida</i> spp.	Topical nystatin or azoles; systemic azoles for resistant disease
Tinea versicolor	Hyper- or hypopigmented scaly patches on trunk; characteristic mixture of hyphae and spores ("spaghetti and meatballs") on KOH preparation	<i>Malassezia furfur</i>	Topical selenium sulfide lotion or azoles

some infections, most often those inappropriately treated with mid- to high-potency topical glucocorticoids. Involvement of the groin (*tinea cruris*) is more common in males than in females. It presents as a scaling, erythematous eruption sparing the scrotum. Infection of the foot (*tinea pedis*) is the most common dermatophyte infection and is often chronic; it is characterized by variable erythema, edema, scaling, pruritus, and occasionally vesiculation. The infection may be widespread or localized but generally involves the web space between the fourth and fifth toes. Infection of the nails (*tinea unguium* or *onychomycosis*) occurs in many patients with *tinea pedis* and is characterized by opacified, thickened nails and subungual debris. The distal-lateral variant is most common. Proximal subungual onychomycosis may be a marker for HIV infection or other immunocompromised states. Dermatophyte infection of the scalp (*tinea capitis*) continues to be common, particularly affecting inner-city children but also affecting adults. The predominant organism is *Trichophyton tonsurans*, which can produce a relatively noninflammatory infection with mild scale and hair loss that is diffuse or localized. *T. tonsurans* can also cause a markedly inflammatory dermatosis with edema and nodules. This latter presentation is a *kerion*.

The diagnosis of tinea can be made from skin scrapings, nail scrapings, or hair by culture or direct microscopic examination with potassium hydroxide (KOH). Nail clippings may be sent for histologic examination with periodic acid–Schiff (PAS) stain.

TREATMENT DERMATOPHYTOSIS

Both topical and systemic therapies may be used in dermatophyte infections. Treatment depends on the site involved and the type of infection. Topical therapy is generally effective for uncomplicated tinea corporis, tinea cruris, and limited tinea pedis. Topical agents are not effective as monotherapy for tinea capitis or onychomycosis (see below). Topical imidazoles, triazoles, and allylamines may be effective therapies for dermatophyte infections, but nystatin is not active against dermatophytes. Topicals are generally applied twice daily, and treatment should continue for 1 week beyond clinical resolution of the infection. Tinea pedis often requires longer treatment courses and frequently relapses. Oral antifungal agents may be required for recalcitrant tinea pedis or tinea corporis.

Oral antifungal agents are required for dermatophyte infections involving the hair and nails and for other infections unresponsive to topical therapy. A fungal etiology should be confirmed by direct microscopic examination or by culture before oral antifungal agents are prescribed. All of the oral agents may cause hepatotoxicity. They should not be used in women who are pregnant or breast-feeding.

Griseofulvin is approved in the United States for dermatophyte infections involving the skin, hair, or nails. When griseofulvin is used, a daily dose of 500 mg microsized or 375 mg ultramicrosized, administered with a fatty meal, is adequate for most dermatophyte infections. Higher doses are required for some cases of tinea pedis and tinea capitis. Markedly inflammatory tinea capitis may result in scarring and hair loss, and systemic or topical glucocorticoids may be helpful in preventing these sequelae. The duration of griseofulvin

therapy may be 2 weeks for uncomplicated tinea corporis, 8–12 weeks for tinea capitis, or as long as 6–18 months for nail infections. Due to high relapse rates, griseofulvin is seldom used for nail infections. Common side effects of griseofulvin include gastrointestinal distress, headache, and urticaria.

Oral itraconazole is approved for onychomycosis. Itraconazole is given with food as either continuous daily therapy (200 mg/d) or pulses (200 mg bid for 1 week per month). Fingernails require 2 months of continuous therapy or two pulses. Toenails require 3 months of continuous therapy or three pulses. Itraconazole has the potential for serious interactions with other drugs requiring the P450 enzyme system for metabolism. Itraconazole should not be administered to patients with evidence of ventricular dysfunction or patients with known CHF. Terbinafine (250 mg/d) is also effective for onychomycosis, and the granule version is approved for treatment of tinea capitis. Therapy with terbinafine is continued for 6 weeks for fingernail and scalp infections and 12 weeks for toenail infections. Terbinafine has fewer interactions with other drugs than itraconazole, but caution should be used with patients who are on multiple medications. The risk/benefit ratio should be considered when an asymptomatic toenail infection is treated with systemic agents.

TINEA VERSICOLOR

Tinea versicolor is caused by a nondermatophytic, dimorphic fungus, *Malassezia furfur*, a normal inhabitant of the skin. The expression of infection is promoted by heat and humidity. The typical lesions consist of oval scaly macules, papules, and patches concentrated on the chest, shoulders, and back but only rarely on the face or distal extremities. On dark skin the lesions often appear as hypopigmented areas, while on light skin they are slightly erythematous or hyperpigmented. A KOH preparation from scaling lesions will demonstrate a confluence of short hyphae and round spores ("spaghetti and meatballs"). Lotions or shampoos containing sulfur, salicylic acid, or selenium sulfide will clear the infection if used daily for 1–2 weeks and then weekly thereafter. These preparations are irritating if left on the skin for >10 min; thus, they should be washed off completely. Treatment with some oral antifungal agents is also effective, but they do not provide lasting results and are not FDA approved for this indication. A very short course of ketoconazole has been used, as have itraconazole and fluconazole. The patient must sweat after taking the medication if it is to be effective. Griseofulvin is not effective and terbinafine is not reliably effective for tinea versicolor.

CANDIDIASIS

Candidiasis is a fungal infection caused by a related group of yeasts whose manifestations may be localized to the skin and mucous membranes or, rarely, may be systemic and life-threatening (Chap. 240). The causative organism is usually *Candida albicans*. These organisms are normal saprophytic inhabitants of the gastrointestinal tract but may overgrow due to broad-spectrum antibiotic therapy, diabetes mellitus, or immunosuppression and cause disease. Candidiasis is a

very common infection in HIV-infected individuals (Chap. 226). The oral cavity is commonly involved. Lesions may occur on the tongue or buccal mucosa (*thrush*) and appear as white plaques. Fissured, macerated lesions at the corners of the mouth (*perlèche*) are often seen in individuals with poorly fitting dentures and may also be associated with candidal infection. In addition, candidal infections have an affinity for sites that are chronically wet and macerated, including the skin around nails (onycholysis and paronychia), and in intertriginous areas. Intertriginous lesions are characteristically edematous, erythematous, and scaly, with scattered “satellite pustules.” In males, there is often involvement of the penis and scrotum as well as the inner aspect of the thighs. In contrast to dermatophyte infections, candidal infections are frequently painful and accompanied by a marked inflammatory response. Diagnosis of candidal infection is based upon the clinical pattern and demonstration of yeast on KOH preparation or culture.

TREATMENT CANDIDIASIS

Treatment involves removal of any predisposing factors such as antibiotic therapy or chronic wetness and the use of appropriate topical or systemic antifungal agents. Effective topicals include nystatin or azoles (miconazole, clotrimazole, econazole, or ketoconazole). The associated inflammatory response accompanying candidal infection on glabrous skin can be treated with a mild glucocorticoid lotion or cream (2.5% hydrocortisone). Systemic therapy is usually reserved for immunosuppressed patients or individuals with chronic or recurrent disease who fail to respond to appropriate topical therapy. Oral agents approved for the treatment of candidiasis include itraconazole and fluconazole. Oral nystatin is effective only for candidiasis of the gastrointestinal tract. Griseofulvin and terbinafine are not effective.

WARTS

Warts are cutaneous neoplasms caused by papillomaviruses. More than 100 different human papillomaviruses (HPVs) have been described. A typical wart, *verruca vulgaris*, is sessile, dome-shaped, and usually about a centimeter in diameter. Its surface is hyperkeratotic, consisting of many small filamentous projections. The HPV types that cause typical *verruca vulgaris* also cause typical plantar warts, flat warts (*verruca plana*), and filiform warts. Plantar warts are endophytic and are covered by thick keratin. Paring of the wart will generally reveal a central core of keratinized debris and punctate bleeding points. Filiform warts are most commonly seen on the face, neck, and skinfolds and present as papillomatous lesions on a narrow base. Flat warts are only slightly elevated and have a velvety, nonverrucous surface. They have a propensity for the face, arms, and legs and are often spread by shaving.

Genital warts begin as small papillomas that may grow to form large, fungating lesions. In women, they may involve the labia, perineum, or perianal skin. In addition, the mucosa of the vagina, urethra, and anus can be involved as well as the cervical epithelium. In men, the lesions often occur initially in the coronal sulcus but may be seen on the shaft of the penis, the scrotum, or the perianal skin or in the urethra.

Appreciable evidence has accumulated indicating that HPV plays a role in the development of neoplasia of the uterine cervix and anogenital skin (Chap. 117). HPV types 16 and 18 have been most intensely studied and are the major risk factors for intraepithelial neoplasia and squamous cell carcinoma of the cervix, anus, vulva, and penis. The risk is higher among patients immunosuppressed after solid organ transplantation and among those infected with HIV. Recent evidence also implicates other HPV types. Histologic examination of biopsied samples from affected sites may reveal changes associated with typical warts and/or features typical of intraepidermal carcinoma (Bowen’s disease). Squamous cell carcinomas associated with HPV infections have also been observed in extragenital skin (Chap. 105), most commonly in patients immunosuppressed after organ transplantation. Patients on long-term immunosuppression should be monitored for the development of squamous cell carcinoma and other cutaneous malignancies.

TREATMENT WARTS

Treatment of warts, other than anogenital warts, should be tempered by the observation that a majority of warts in normal individuals resolve spontaneously within 1–2 years. There are many modalities available to treat warts, but no single therapy is universally effective. Factors that influence the choice of therapy include the location of the wart, the extent of disease, the age and immunologic status of the patient, and the patient’s desire for therapy. Perhaps the most useful and convenient method for treating warts in almost any location is cryotherapy with liquid nitrogen. Equally effective for nongenital warts, but requiring much more patient compliance, is the use of keratolytic agents such as salicylic acid plasters or solutions. For genital warts, in-office application of a podophyllin solution is moderately effective but may be associated with marked local reactions. Prescription preparations of dilute, purified podophyllin are available for home use. Topical imiquimod, a potent inducer of local cytokine release, has been approved for treatment of genital warts. A new topical compound composed of green tea extracts (sin catechins) is also available. Conventional and laser surgical procedures may be required for recalcitrant warts. Recurrence of warts appears to be common to all these modalities. A highly effective vaccine for selected types of HPV has been approved by the FDA, and its use appears to reduce the incidence of anogenital and cervical carcinoma.

HERPES SIMPLEX

See Chap. 216.

HERPES ZOSTER

See Chap. 217.

ACNE

ACNE VULGARIS

Acne vulgaris is a self-limited disorder primarily of teenagers and young adults, although perhaps 10–20% of adults may continue to experience some form of the disorder. The permissive factor for the expression of the disease in adolescence is the increase in sebum production by sebaceous glands after puberty. Small cysts, called *comedones*, form in hair follicles due to blockage of the follicular orifice by retention of keratinous material and sebum. The activity of bacteria (*Propionibacterium acnes*) within the comedones releases free fatty acids from sebum, causes inflammation within the cyst, and results in rupture of the cyst wall. An inflammatory foreign-body reaction develops as result of extrusion of oily and keratinous debris from the cyst.

The clinical hallmark of acne vulgaris is the comedone, which may be closed (*whitehead*) or open (*blackhead*). Closed comedones appear as 1- to 2-mm pebbly white papules, which are accentuated when the skin is stretched. They are the precursors of inflammatory lesions of acne vulgaris. The contents of closed comedones are not easily expressed. Open comedones, which rarely result in inflammatory acne lesions, have a large dilated follicular orifice and are filled with easily expressible oxidized, darkened, oily debris. Comedones are usually accompanied by inflammatory lesions: papules, pustules, or nodules.

The earliest lesions seen in adolescence are generally mildly inflamed or noninflammatory comedones on the forehead. Subsequently, more typical inflammatory lesions develop on the cheeks, nose, and chin (Fig. 71-7). The most common location for acne is the face, but involvement of the chest and back is common. Most disease remains mild and does not lead to scarring. A small number of patients develop large inflammatory cysts and nodules, which may drain and result in significant scarring. Regardless of the severity, acne may affect a patient’s quality of life. With adequate treatment, this effect may be transient. In the case of severe, scarring acne, the effects can be permanent and profound. Early therapeutic intervention in severe acne is essential.

Exogenous and endogenous factors can alter the expression of acne vulgaris. Friction and trauma (from headbands or chin straps of athletic helmets), application of comedogenic topical agents (cosmetics



FIGURE 71-7 Acne vulgaris. An example of acne vulgaris with inflammatory papules, pustules, and comedones. (Courtesy of Kalman Watsky, MD; with permission.)

or hair preparations), or chronic topical exposure to certain industrial compounds may elicit or aggravate acne. Glucocorticoids, topical or systemic, may also elicit acne. Other systemic medications such as oral contraceptive pills, lithium, isoniazid, androgenic steroids, halogens, phenytoin, and phenobarbital may produce acneiform eruptions or aggravate preexisting acne. Genetic factors and polycystic ovary disease may also play a role.

TREATMENT ACNE VULGARIS

Treatment of acne vulgaris is directed toward elimination of comedones by normalizing follicular keratinization, decreasing sebaceous gland activity, decreasing the population of *P. acnes*, and decreasing inflammation. Minimal to moderate pauci-inflammatory disease may respond adequately to local therapy alone. Although areas affected with acne should be kept clean, overly vigorous scrubbing may aggravate acne due to mechanical rupture of comedones. Topical agents such as retinoic acid, benzoyl peroxide, or salicylic acid may alter the pattern of epidermal desquamation, preventing the formation of comedones and aiding in the resolution of preexisting cysts. Topical antibacterial agents (such as azelaic acid, erythromycin, clindamycin, or dapsone) are also useful adjuncts to therapy.

Patients with moderate to severe acne with a prominent inflammatory component will benefit from the addition of systemic therapy, such as tetracycline in doses of 250–500 mg bid or doxycycline in doses of 100 mg bid. Minocycline is also useful. Such antibiotics appear to have anti-inflammatory effects independent of their antibacterial effects. Female patients who do not respond to oral antibiotics may benefit from hormonal therapy. Several oral contraceptives are now approved by the FDA for use in the treatment of acne vulgaris.

Patients with severe nodulocystic acne unresponsive to the therapies discussed above may benefit from treatment with the synthetic retinoid isotretinoin. Its dose is based on the patient's weight, and it is given once daily for 5 months. Results are excellent in appropriately selected patients. Its use is highly regulated due to its potential for severe adverse events, primarily teratogenicity and depression. In addition, patients receiving this medication develop extremely dry skin and cheilitis and must be followed for development of hypertriglyceridemia.

At present, prescribers must enroll in a program designed to prevent pregnancy and adverse events while patients are taking isotretinoin. These measures are imposed to ensure that all prescribers are familiar with the risks of isotretinoin; that all female patients have two negative pregnancy tests prior to initiation of therapy and a negative pregnancy test prior to each refill; and that all patients have been warned about the risks associated with isotretinoin.



FIGURE 71-8 Acne rosacea. Prominent facial erythema, telangiectasia, scattered papules, and small pustules are seen in this patient with acne rosacea. (Courtesy of Robert Swerlick, MD; with permission.)

ACNE ROSACEA

Acne rosacea, commonly referred to simply as rosacea, is an inflammatory disorder predominantly affecting the central face. Persons most often affected are Caucasians of northern European background, but rosacea also occurs in patients with dark skin. Rosacea is seen almost exclusively in adults, only rarely affecting patients <30 years old. Rosacea is more common in women, but those most severely affected are men. It is characterized by the presence of erythema, telangiectases, and superficial pustules (Fig. 71-8) but is not associated with the presence of comedones. Rosacea rarely involves the chest or back.

There is a relationship between the tendency for facial flushing and the subsequent development of acne rosacea. Often, individuals with rosacea initially demonstrate a pronounced flushing reaction. This may be in response to heat, emotional stimuli, alcohol, hot drinks, or spicy foods. As the disease progresses, the flush persists longer and longer and may eventually become permanent. Papules, pustules, and telangiectases can become superimposed on the persistent flush. Rosacea of very long standing may lead to connective tissue overgrowth, particularly of the nose (*rhinophyma*). Rosacea may also be complicated by various inflammatory disorders of the eye, including keratitis, blepharitis, iritis, and recurrent chalazion. These ocular problems are potentially sight-threatening and warrant ophthalmologic evaluation.

TREATMENT ACNE ROSACEA

Acne rosacea can be treated topically or systemically. Mild disease often responds to topical metronidazole, sodium sulfacetamide, or azelaic acid. More severe disease requires oral tetracyclines: tetracycline, 250–500 mg bid; doxycycline, 100 mg bid; or minocycline, 50–100 mg bid. Residual telangiectasia may respond to laser therapy. Topical glucocorticoids, especially potent agents, should be avoided because chronic use of these preparations may elicit rosacea. Application of topical agents to the skin is not effective treatment for ocular disease.

SKIN DISEASES AND SMALLPOX VACCINATION

Although smallpox vaccinations were discontinued several decades ago for the general population, they are still required for certain military personnel and first responders. In the absence of a bioterrorism attack and a real or potential exposure to smallpox, such vaccination is contraindicated in persons with a history of skin diseases such as AD, eczema, and psoriasis, who have a higher incidence of adverse events associated with smallpox vaccination. In the case of such exposure, the risk of smallpox infection outweighs the risk of adverse events from the vaccine (Chap. 261e).

72 Skin Manifestations of Internal Disease

Jean L. Bologna, Irwin M. Braverman

It is a generally accepted concept in medicine that the skin can develop signs of internal disease. Therefore, in textbooks of medicine, one finds a chapter describing in detail the major systemic disorders that can be identified by cutaneous signs. The underlying assumption of such a chapter is that the clinician has been able to identify the specific disorder in the patient and needs only to read about it in the textbook. In reality, concise differential diagnoses and the identification of these disorders are actually difficult for the nondermatologist because he or she is not well-versed in the recognition of cutaneous lesions or their spectrum of presentations. Therefore, this chapter covers this particular topic of cutaneous medicine not by simply focusing on individual diseases, but by describing the various presenting clinical signs and symptoms that point to specific disorders. Concise differential diagnoses will be generated in which the significant diseases will be distinguished from the more common cutaneous disorders that have minimal or no significance with regard to associated internal disease. The latter disorders are reviewed in table form and always need to be excluded when considering the former. For a detailed description of individual diseases, the reader should consult a dermatologic text.

PAPULOSQUAMOUS SKIN LESIONS

(Table 72-1) When an eruption is characterized by elevated lesions, either papules (<1 cm) or plaques (>1 cm), in association with scale, it is referred to as *papulosquamous*. The most common papulosquamous diseases—*tinea*, *psoriasis*, *pityriasis rosea*, and *lichen planus*—are primary cutaneous disorders (Chap. 71). When psoriatic lesions are accompanied by arthritis, the possibility of psoriatic arthritis or reactive arthritis (formerly known as Reiter's syndrome) should be considered. A history of oral ulcers, conjunctivitis, uveitis, and/or urethritis points to the latter diagnosis. Lithium, beta blockers, HIV or streptococcal infections, and a rapid taper of systemic glucocorticoids are known to exacerbate psoriasis. Comorbidities in patients with psoriasis include cardiovascular disease and metabolic syndrome.

Whenever the diagnosis of pityriasis rosea or lichen planus is made, it is important to review the patient's medications because the eruption may resolve by simply discontinuing the offending agent. Pityriasis rosea-like drug eruptions are seen most commonly with beta blockers, angiotensin-converting enzyme (ACE) inhibitors, and metronidazole, whereas the drugs that can produce a lichenoid eruption include thiazides, antimalarials, quinidine, beta blockers, and ACE inhibitors. In some populations, there is a higher prevalence of hepatitis C viral infection in patients with lichen planus. Lichen planus-like lesions are also observed in chronic graft-versus-host disease.

In its early stages, the mycosis fungoides (MF) form of *cutaneous T cell lymphoma* (CTCL) may be confused with eczema or psoriasis, but it often fails to respond to the appropriate therapy for those inflammatory diseases. MF can develop within lesions of large-plaque parapsoriasis and is suggested by an increase in the thickness of the lesions. The diagnosis of MF is established by skin biopsy in which collections of atypical T lymphocytes are found in the epidermis and dermis. As the disease progresses, cutaneous tumors and lymph node involvement may appear.

In *secondary syphilis*, there are scattered red-brown papules with thin scale. The eruption often involves the palms and soles and can resemble pityriasis rosea. Associated findings are helpful in making the diagnosis and include annular plaques on the face, nonscarring alopecia, condyloma lata (broad-based and moist), and mucous patches as well as lymphadenopathy, malaise, fever, headache, and myalgias. The interval between the primary chancre and the secondary stage is usually 4–8 weeks, and spontaneous resolution without appropriate therapy occurs.

TABLE 72-1 SELECTED CAUSES OF PAPULOSQUAMOUS SKIN LESIONS

1. Primary cutaneous disorders
 - a. Tinea^a
 - b. Psoriasis^a
 - c. Pityriasis rosea^a
 - d. Lichen planus^a
 - e. Parapsoriasis, small plaque and large plaque
 - f. Bowen's disease (squamous cell carcinoma in situ)^a
2. Drugs
3. Systemic diseases
 - a. Lupus erythematosus, primarily subacute or chronic (discoid) lesions^a
 - b. Cutaneous T cell lymphoma, in particular, mycosis fungoides^a
 - c. Secondary syphilis
 - d. Reactive arthritis (formerly known as Reiter's syndrome)
 - e. Sarcoidosis^a

^aDiscussed in detail in Chap. 71; cardiovascular disease and the metabolic syndrome are comorbidities in psoriasis; primarily in Europe, hepatitis C virus is associated with oral lichen planus. ^bAssociated with chronic sun exposure more often than exposure to arsenic; usually one or a few lesions. ^cSee also Red Lesions in "Papulonodular Skin Lesions." ^dAlso cutaneous lesions of HTLV-1-associated adult T cell leukemia/lymphoma. ^eSee also Red-Brown Lesions in "Papulonodular Skin Lesions."

ERYTHRODERMA

(Table 72-2) *Erythroderma* is the term used when the majority of the skin surface is erythematous (red in color). There may be associated scale, erosions, or pustules as well as shedding of the hair and nails. Potential systemic manifestations include fever, chills, hypothermia, reactive lymphadenopathy, peripheral edema, hypoalbuminemia, and high-output cardiac failure. The major etiologies of erythroderma are (1) cutaneous diseases such as psoriasis and dermatitis (Table 72-3); (2) drugs; (3) systemic diseases, most commonly CTCL; and (4) idiopathic. In the first three groups, the location and description of the initial lesions, prior to the development of the erythroderma, aid in the diagnosis. For example, a history of red scaly plaques on the elbows and knees would point to psoriasis. It is also important to examine the skin carefully for a migration of the erythema and associated secondary changes such as pustules or erosions. Migratory waves of erythema studded with superficial pustules are seen in *pustular psoriasis*.

Drug-induced erythroderma (exfoliative dermatitis) may begin as an exanthematous (morbilliform) eruption (Chap. 74) or may arise as diffuse erythema. A number of drugs can produce an erythroderma, including penicillins, sulfonamides, carbamazepine, phenytoin, and allopurinol. Fever and peripheral eosinophilia often accompany the eruption, and there may also be facial swelling, hepatitis, myocarditis, thyroiditis, and allergic interstitial nephritis; this constellation is frequently referred to as *drug reaction with eosinophilia and systemic symptoms* (DRESS) or *drug-induced hypersensitivity reaction* (DIHS). In addition, these reactions, especially to aromatic anticonvulsants, can lead to a pseudolymphoma syndrome (with adenopathy and

TABLE 72-2 CAUSES OF ERYTHRODERMA

1. Primary cutaneous disorders
 - a. Psoriasis^a
 - b. Dermatitis (atopic > contact >> stasis [with autosensitization] or seborrheic)^a
 - c. Pityriasis rubra pilaris
2. Drugs
3. Systemic diseases
 - a. Cutaneous T cell lymphoma (Sézary syndrome, erythrodermic mycosis fungoides)
 - b. Other lymphomas
4. Idiopathic (usually older men)

^aDiscussed in detail in Chap. 71.

TABLE 72-3 ERYTHRODERMA (PRIMARY CUTANEOUS DISORDERS)

	Initial Lesions	Location of Initial Lesions	Other Findings	Diagnostic Aids	Treatment
Psoriasis ^a	Pink-red, silvery scale, sharply demarcated	Elbows, knees, scalp, presacral area, intergluteal fold	Nail dystrophy, arthritis, pustules; SAPHO syndrome, especially with palmoplantar pustulosis	Skin biopsy	Topical glucocorticoids, vitamin D; UV-B (narrowband) > PUVA; oral retinoid; MTX, cyclosporine, anti-TNF agents, anti-IL-12/23 Ab
Dermatitis^a					
Atopic	Acute: Erythema, fine scale, crust, indistinct borders, excoriations Chronic: Lichenification (increased skin markings), excoriations	Antecubital and popliteal fossae, neck, hands, eyelids	Pruritus Personal and/or family history of atopy, including asthma, allergic rhinitis or conjunctivitis, and atopic dermatitis Exclude secondary infection with <i>Staphylococcus aureus</i> or HSV Exclude superimposed irritant or allergic contact dermatitis	Skin biopsy	Topical glucocorticoids, tacrolimus, pimecrolimus, tar, and antipruritics; oral antihistamines; open wet dressings; UV-B ± UV-A > PUVA; oral/IM glucocorticoids (short-term); MTX; mycophenolate mofetil; azathioprine; cyclosporine Topical or oral antibiotics
Contact	Local: Erythema, crusting, vesicles, and bullae Systemic: Erythema, fine scale, crust	Depends on offending agent Generalized vs major intertriginous zones (especially groin)	Irritant—onset often within hours Allergic—delayed-type hypersensitivity; lag time of 48 h Patient has history of allergic contact dermatitis to topical agent and then receives systemic medication that is structurally related, e.g., thiuram (skin), disulfiram (oral)	Patch testing; open use test Patch testing	Remove irritant or allergen; topical glucocorticoids; oral antihistamines; oral/IM glucocorticoids (short-term) Same as local
Seborrheic (rare in adults)	Pink-red to pink-orange, greasy scale	Scalp, nasolabial folds, eyebrows, intertriginous zones	Flares with stress, HIV infection Associated with Parkinson's disease	Skin biopsy	Topical glucocorticoids and imidazoles
Stasis (with autosensitization)	Erythema, crusting, excoriations	Lower extremities	Pruritus, lower extremity edema, varicosities, hemosiderin deposits History of venous ulcers, thrombophlebitis, and/or cellulitis Exclude cellulitis Exclude superimposed contact dermatitis, e.g., topical neomycin	Skin biopsy	Topical glucocorticoids; open wet dressings; leg elevation; pressure stockings; pressure wraps if associated ulcers
Pityriasis rubra pilaris	Orange-red (salmon-colored), perifollicular papules	Generalized, but characteristic "skip" areas of normal skin	Wax-like palmoplantar keratoderma Exclude cutaneous T cell lymphoma	Skin biopsy	Isotretinoin or acitretin; MTX; perhaps anti-IL-12/23 Ab, IV anti-TNF agent

^aDiscussed in detail in [Chap. 71](#).

Abbreviations: Ab, antibody; HSV, herpes simplex virus; IL, interleukin; IM, intramuscular; IV, intravenous; MTX, methotrexate; PUVA, psoralens + ultraviolet A irradiation; SAPHO, synovitis, acne, pustulosis, hyperostosis, and osteitis (also referred to as chronic recurrent multifocal osteomyelitis); TNF, tumor necrosis factor; UV-A, ultraviolet A irradiation; UV-B, ultraviolet B irradiation.

circulating atypical lymphocytes), while reactions to allopurinol may be accompanied by gastrointestinal bleeding.

The most common malignancy that is associated with erythroderma is CTCL; in some series, up to 25% of the cases of erythroderma were due to CTCL. The patient may progress from isolated plaques and tumors, but more commonly, the erythroderma is present throughout the course of the disease (Sézary syndrome). In the Sézary syndrome, there are circulating clonal atypical T lymphocytes, pruritus, and lymphadenopathy. In cases of erythroderma where there is no apparent cause (idiopathic), longitudinal evaluation is mandatory to monitor for the possible development of CTCL. There have been isolated case reports of erythroderma secondary to some solid tumors—lung, liver, prostate, thyroid, and colon—but it is primarily during a late stage of the disease.

ALOPECIA

([Table 72-4](#)) The two major forms of alopecia are scarring and non-scarring. *Scarring alopecia* is associated with fibrosis, inflammation, and loss of hair follicles. A smooth scalp with a decreased number of follicular openings is usually observed clinically, but in some patients, the changes are seen only in biopsy specimens from affected areas. In *nonscarring alopecia*, the hair shafts are absent or miniaturized, but the hair follicles are preserved, explaining the reversible nature of nonscarring alopecia.

The most common causes of nonscarring alopecia include *androgenetic alopecia*, *telogen effluvium*, *alopecia areata*, *tinea capitis*, and the early phase of *traumatic alopecia* ([Table 72-5](#)). In women with androgenetic alopecia, an elevation in circulating levels of androgens may be seen as a result of ovarian or adrenal gland dysfunction or neoplasm. When there are signs of virilization, such as a deepened voice and enlarged clitoris, the possibility of an ovarian or adrenal gland tumor should be considered.

Exposure to various drugs can also cause diffuse hair loss, usually by inducing a telogen effluvium. An exception is the anagen effluvium observed with antimetabolic agents such as daunorubicin. Alopecia is a side effect of the following drugs: warfarin, heparin, propylthiouracil, carbimazole, isotretinoin, acitretin, lithium, beta blockers, interferons, colchicine, and amphetamines. Fortunately, spontaneous regrowth usually follows discontinuation of the offending agent.

Less commonly, nonscarring alopecia is associated with *lupus erythematosus* and *secondary syphilis*. In systemic lupus there are two forms of alopecia—one is scarring secondary to discoid lesions (see below), and the other is nonscarring. The latter form coincides with flares of systemic disease and may involve the entire scalp or just the frontal scalp, with the appearance of multiple short hairs ("lupus hairs") as a sign of initial regrowth. Scattered, poorly circumscribed patches of alopecia with a "moth-eaten" appearance are a manifestation

TABLE 72-4 CAUSES OF ALOPECIA

I. Nonscarring alopecia
A. Primary cutaneous disorders
1. Androgenetic alopecia
2. Telogen effluvium
3. Alopecia areata
4. Tinea capitis
5. Traumatic alopecia ^a
B. Drugs
C. Systemic diseases
1. Systemic lupus erythematosus
2. Secondary syphilis
3. Hypothyroidism
4. Hyperthyroidism
5. Hypopituitarism
6. Deficiencies of protein, biotin, zinc, and perhaps iron
II. Scarring alopecia
A. Primary cutaneous disorders
1. Cutaneous lupus (chronic discoid lesions) ^b
2. Lichen planus
3. Central centrifugal cicatricial alopecia
4. Folliculitis decalvans
5. Linear scleroderma (morphea) ^c
B. Systemic diseases
1. Discoid lesions in the setting of systemic lupus erythematosus ^b
2. Sarcoidosis
3. Cutaneous metastases

^aMost patients with trichotillomania, pressure-induced alopecia, or early stages of traction alopecia. ^bWhile the majority of patients with discoid lesions have only cutaneous disease, these lesions do represent one of the 11 American College of Rheumatology criteria (1982) for systemic lupus erythematosus. ^cCan involve underlying muscles and osseous structures.

of the secondary stage of syphilis. Diffuse thinning of the hair is also associated with hypothyroidism and hyperthyroidism (Table 72-4).

Scarring alopecia is more frequently the result of a primary cutaneous disorder such as *lichen planus*, *folliculitis decalvans*, *chronic cutaneous (discoid) lupus*, or *linear scleroderma (morphea)* than it is a sign of systemic disease. Although the scarring lesions of *discoid lupus* can be seen in patients with systemic lupus, in the majority of patients, the disease process is limited to the skin. Less common causes of scarring alopecia include *sarcoidosis* (see “Papulonodular Skin Lesions,” below) and *cutaneous metastases*.

In the early phases of discoid lupus, lichen planus, and folliculitis decalvans, there are circumscribed areas of alopecia. Fibrosis and subsequent loss of hair follicles are observed primarily in the center of these alopecic patches, whereas the inflammatory process is most prominent at the periphery. The areas of active inflammation in discoid lupus are erythematous with scale, whereas the areas of previous inflammation are often hypopigmented with a rim of hyperpigmentation. In lichen planus, perifollicular macules at the periphery are usually violet-colored. A complete examination of the skin and oral mucosa combined with a biopsy and direct immunofluorescence microscopy of inflamed skin will aid in distinguishing these two entities. The peripheral active lesions in folliculitis decalvans are follicular pustules; these patients can develop a reactive arthritis.

FIGURATE SKIN LESIONS

(Table 72-6) In *figurate eruptions*, the lesions form rings and arcs that are usually erythematous but can be skin-colored to brown. Most commonly, they are due to primary cutaneous diseases such as *tinea*, *urticaria*, *granuloma annulare*, and *erythema annulare centrifugum* (Chaps. 71 and 73). An underlying systemic illness is found in a second, less common group of migratory annular erythemas. It includes *erythema migrans*, *erythema gyratum repens*, *erythema marginatum*, and *necrolytic migratory erythema*.

In *erythema gyratum repens*, one sees numerous mobile concentric arcs and wavefronts that resemble the grain in wood. A search for an

TABLE 72-5 NONSCARRING ALOPECIA (PRIMARY CUTANEOUS DISORDERS)

	Clinical Characteristics	Pathogenesis	Treatment
Telogen effluvium	Diffuse shedding of normal hairs Follows major stress (high fever, severe infection) or change in hormone levels (postpartum) Reversible without treatment	Stress causes more of the asynchronous growth cycles of individual hairs to become synchronous; therefore, larger numbers of growing (anagen) hairs simultaneously enter the dying (telogen) phase	Observation; discontinue any drugs that have alopecia as a side effect; must exclude underlying metabolic causes, e.g., hypothyroidism, hyperthyroidism
Androgenetic alopecia (male pattern; female pattern)	Miniaturization of hairs along the midline of the scalp Recession of the anterior scalp line in men and some women	Increased sensitivity of affected hairs to the effects of androgens Increased levels of circulating androgens (ovarian or adrenal source in women)	If no evidence of hyperandrogenemia, then topical minoxidil; finasteride ^a ; spironolactone (women); hair transplant
Alopecia areata	Well-circumscribed, circular areas of hair loss, 2–5 cm in diameter In extensive cases, coalescence of lesions and/or involvement of other hair-bearing surfaces of the body Pitting or sandpapered appearance of the nails	The germinative zones of the hair follicles are surrounded by T lymphocytes Occasional associated diseases: hyperthyroidism, hypothyroidism, vitiligo, Down syndrome	Topical anthralin or tazarotene; intraleisional glucocorticoids; topical contact sensitizers
Tinea capitis	Varies from scaling with minimal hair loss to discrete patches with “black dots” (broken infected hairs) to boggy plaque with pustules (kerion) ^b	Invasion of hairs by dermatophytes, most commonly <i>Trichophyton tonsurans</i>	Oral griseofulvin or terbinafine plus 2.5% selenium sulfide or ketoconazole shampoo; examine family members
Traumatic alopecia ^c	Broken hairs, often of varying lengths Irregular outline	Traction with curlers, rubber bands, braiding Exposure to heat or chemicals (e.g., hair straighteners) Mechanical pulling (trichotillomania)	Discontinuation of offending hair style or chemical treatments; diagnosis of trichotillomania may require observation of shaved hairs (for growth) or biopsy, possibly followed by psychotherapy

^aTo date, Food and Drug Administration–approved for men. ^bScarring alopecia can occur at sites of kerions. ^cMay also be scarring, especially late-stage traction alopecia.

TABLE 72-6 CAUSES OF FIGURATE SKIN LESIONS

- I. Primary cutaneous disorders
 - A. Tinea
 - B. Urticaria (primary in $\geq 90\%$ of patients)
 - C. Granuloma annulare
 - D. Erythema annulare centrifugum
 - E. Psoriasis
- II. Systemic diseases
 - A. Migratory
 1. Erythema migrans (CDC case definition is ≥ 5 cm in diameter)
 2. Urticaria ($\leq 10\%$ of patients)
 3. Erythema gyratum repens
 4. Erythema marginatum
 5. Pustular psoriasis (generalized)
 6. Necrolytic migratory erythema (glucagonoma syndrome)^a
 - B. Nonmigratory
 1. Sarcoidosis
 2. Subacute cutaneous lupus erythematosus
 3. Secondary syphilis
 4. Cutaneous T cell lymphoma (especially mycosis fungoides)

^aMigratory erythema with erosions; favors lower extremities and girdle area.

Abbreviation: CDC, Centers for Disease Control and Prevention.

underlying malignancy is mandatory in a patient with this eruption. Erythema migrans is the cutaneous manifestation of Lyme disease, which is caused by the spirochete *Borrelia burgdorferi*. In the initial stage (3–30 days after tick bite), a single annular lesion is usually seen, which can expand to ≥ 10 cm in diameter. Within several days, up to half of the patients develop multiple smaller erythematous lesions at sites distant from the bite. Associated symptoms include fever, headache, photophobia, myalgias, arthralgias, and malar rash. Erythema marginatum is seen in patients with rheumatic fever, primarily on the trunk. Lesions are pink-red in color, flat to minimally elevated, and transient.

There are additional cutaneous diseases that present as annular eruptions but lack an obvious migratory component. Examples include *CTCL*, *subacute cutaneous lupus*, *secondary syphilis*, and *sarcoidosis* (see “Papulonodular Skin Lesions,” below).

ACNE

(Table 72-7) In addition to *acne vulgaris* and *acne rosacea*, the two major forms of acne (Chap. 71), there are drugs and systemic diseases that can lead to acneiform eruptions.

Patients with the *carcinoid syndrome* have episodes of flushing of the head, neck, and sometimes the trunk. Resultant skin changes of the face, in particular telangiectasias, may mimic the clinical appearance of *acne rosacea*.

TABLE 72-7 CAUSES OF ACNEIFORM ERUPTIONS

- I. Primary cutaneous disorders
 - A. Acne vulgaris
 - B. Acne rosacea
- II. Drugs, e.g., anabolic steroids, glucocorticoids, lithium, EGFR^a inhibitors, iodides
- III. Systemic diseases
 - A. Increased androgen production
 1. Adrenal origin, e.g., Cushing’s disease, 21-hydroxylase deficiency
 2. Ovarian origin, e.g., polycystic ovary syndrome, ovarian hyperthecosis
 - B. Cryptococcosis, disseminated
 - C. Dimorphic fungal infections
 - D. Behçet’s disease

^aEGFR, epidermal growth factor receptor.

PUSTULAR LESIONS

Acneiform eruptions (see “Acne,” above) and *folliculitis* represent the most common pustular dermatoses. An important consideration in the evaluation of follicular pustules is a determination of the associated pathogen, e.g., normal flora, *Staphylococcus aureus*, *Pseudomonas aeruginosa* (“hot tub” folliculitis), *Malassezia*, dermatophytes (Majocchi’s granuloma), and *Demodex* spp. Noninfectious forms of folliculitis include HIV- or immunosuppression-associated eosinophilic folliculitis and folliculitis secondary to drugs such as glucocorticoids, lithium, and epidermal growth factor receptor (EGFR) inhibitors. Administration of high-dose systemic glucocorticoids can result in a widespread eruption of follicular pustules on the trunk, characterized by lesions in the same stage of development. With regard to underlying systemic diseases, nonfollicular-based pustules are a characteristic component of pustular psoriasis (sterile) and can be seen in septic emboli of bacterial or fungal origin (see “Purpura,” below). In patients with acute generalized exanthematous pustulosis (AGEP) due primarily to medications (e.g., cephalosporins), there are large areas of erythema studded with multiple sterile pustules in addition to neutrophilia.

TELANGIECTASIAS

(Table 72-8) To distinguish the various types of telangiectasias, it is important to examine the shape and configuration of the dilated blood vessels. *Linear telangiectasias* are seen on the face of patients

TABLE 72-8 CAUSES OF TELANGIECTASIAS

- I. Primary cutaneous disorders
 - A. Linear/branching
 1. Acne rosacea
 2. Actinically damaged skin
 3. Venous hypertension
 4. Generalized essential telangiectasia
 5. Cutaneous collagenous vasculopathy
 6. Within basal cell carcinomas or cutaneous lymphoma
 - B. Poikiloderma
 1. Ionizing radiation^a
 2. Parapsoriasis, large plaque
 - C. Spider angioma
 1. Idiopathic
 2. Pregnancy
- II. Systemic diseases
 - A. Linear/branching
 1. Carcinoid
 2. Ataxia-telangiectasia
 3. Mastocytosis
 - B. Poikiloderma
 1. Dermatomyositis
 2. Mycosis fungoides
 3. Xeroderma pigmentosum
 - C. Mat
 1. Systemic sclerosis (scleroderma)
 - D. Periungual/cuticular
 1. Lupus erythematosus
 2. Systemic sclerosis (scleroderma)
 3. Dermatomyositis
 4. Hereditary hemorrhagic telangiectasia
 - E. Papular
 1. Hereditary hemorrhagic telangiectasia
 - F. Spider angioma
 1. Cirrhosis

^aBecoming less common.

with *actinically damaged skin* and *acne rosacea*, and they are found on the legs of patients with *venous hypertension* and *generalized essential telangiectasia*. Patients with an unusual form of *mastocytosis* (*telangiectasia macularis eruptiva perstans*) and the *carcinoid syndrome* (see “Acne,” above) also have linear telangiectasias. Lastly, linear telangiectasias are found in areas of cutaneous inflammation. For example, lesions of discoid lupus frequently have telangiectasias within them.

Poikiloderma is a term used to describe a patch of skin with: (1) reticulated hypo- and hyperpigmentation, (2) wrinkling secondary to epidermal atrophy, and (3) telangiectasias. Poikiloderma does not imply a single disease entity—although it is becoming less common, it is seen in skin damaged by *ionizing radiation* as well as in patients with autoimmune connective tissue diseases, primarily *dermatomyositis* (DM), and rare genodermatoses (e.g., Kindler syndrome).

In *systemic sclerosis (scleroderma)* the dilated blood vessels have a unique configuration and are known as *mat telangiectasias*. The lesions are broad macules that usually measure 2–7 mm in diameter but occasionally are larger. Mats have a polygonal or oval shape, and their erythematous color may appear uniform, but, upon closer inspection, the erythema is the result of delicate telangiectasias. The most common locations for mat telangiectasias are the face, oral mucosa, and hands—peripheral sites that are prone to intermittent ischemia. The limited form of systemic sclerosis, often referred to as the CREST (calcinosis cutis, Raynaud’s phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia) variant (Chap. 382), is associated with a chronic course and anticentromere antibodies. Mat telangiectasias are an important clue to the diagnosis of this variant as well as the diffuse form of systemic sclerosis because they may be the only cutaneous finding.

Periungual telangiectasias are pathognomonic signs of the three major autoimmune connective tissue diseases: *lupus erythematosus*, *systemic sclerosis*, and *DM*. They are easily visualized by the naked eye and occur in at least two-thirds of these patients. In both DM and lupus, there is associated nailfold erythema, and in DM, the erythema is often accompanied by “ragged” cuticles and fingertip tenderness. Under 10× magnification, the blood vessels in the nailfolds of lupus patients are tortuous and resemble “glomeruli,” whereas in systemic sclerosis and DM, there is a loss of capillary loops and those that remain are markedly dilated.

In *hereditary hemorrhagic telangiectasia* (Osler-Rendu-Weber disease), the lesions usually appear during adolescence (mucosal) and adulthood (cutaneous) and are most commonly seen on the mucous membranes (nasal, orolabial), face, and distal extremities, including under the nails. They represent arteriovenous (AV) malformations of the dermal microvasculature, are dark red in color, and are usually slightly elevated. When the skin is stretched over an individual lesion, an eccentric punctum with radiating legs is seen. Although the degree of systemic involvement varies in this autosomal dominant disease (due primarily to mutations in either the endoglin or activin receptor-like kinase gene), the major symptoms are recurrent epistaxis and gastrointestinal bleeding. The fact that these mucosal telangiectasias are actually AV communications helps to explain their tendency to bleed.

HYPOPIGMENTATION

(Table 72-9) Disorders of hypopigmentation are often classified as either diffuse or localized. The classic example of *diffuse hypopigmentation* is *oculocutaneous albinism* (OCA). The most common forms are due to mutations in the tyrosinase gene (type I) or the *P* gene (type II); patients with type IA OCA have a total lack of enzyme activity. At birth, different forms of OCA can appear similar—white hair, gray-blue eyes, and pink-white skin. However, the patients with no tyrosinase activity maintain this phenotype, whereas those with decreased activity will acquire some pigmentation of the eyes, hair, and skin as they age. The degree of pigment formation is also a function of racial background, and the pigmentary dilution is more readily apparent when patients are compared to their first-degree relatives. The ocular findings in OCA correlate with the degree of hypopigmentation and

TABLE 72-9 CAUSES OF HYPOPIGMENTATION

I. Primary cutaneous disorders
A. Diffuse
1. Generalized vitiligo ^a
B. Localized
1. Idiopathic guttate hypomelanosis
2. Postinflammatory
3. Tinea (pityriasis) versicolor
4. Vitiligo ^a
5. Chemical- or drug-induced leukoderma
6. Nevus depigmentosus
7. Piebaldism ^a
II. Systemic diseases
A. Diffuse
1. Oculocutaneous albinism ^b
2. Hermansky-Pudlak syndrome ^{b,c}
3. Chédiak-Higashi syndrome ^{b,d}
4. Phenylketonuria
B. Localized
1. Systemic sclerosis (scleroderma)
2. Melanoma-associated leukoderma, spontaneous or immunotherapy-induced
3. Vogt-Koyanagi-Harada syndrome
4. Onchocerciasis
5. Sarcoidosis
6. Cutaneous T cell lymphoma (especially mycosis fungoides)
7. Tuberculoid and indeterminate leprosy
8. Linear nevoid hypopigmentation (hypomelanosis of Ito) ^e
9. Incontinentia pigmenti (stage IV)
10. Tuberous sclerosis
11. Waardenburg syndrome and Shah-Waardenburg syndrome

^aAbsence of melanocytes in areas of leukoderma. ^bNormal number of melanocytes.

^cPlatelet storage defect and restrictive lung disease secondary to deposits of ceroid-like material or immunodeficiency; due to mutations in β subunit of adaptor protein 3 as well as subunits of biogenesis of lysosome-related organelles complex (BLOC)-1, -2, and -3.

^dGiant lysosomal granules and recurrent infections. ^eMinority of patients in a nonreferral setting have systemic abnormalities (musculoskeletal, central nervous system, ocular).

include decreased visual acuity, nystagmus, photophobia, strabismus, and a lack of normal binocular vision.

The differential diagnosis of *localized hypomelanosis* includes the following primary cutaneous disorders: *idiopathic guttate hypomelanosis*, *postinflammatory hypopigmentation*, *tinea (pityriasis) versicolor*, *vitiligo*, *chemical- or drug-induced leukoderma*, *nevus depigmentosus* (see below), and *piebaldism* (Table 72-10). In this group of diseases, the areas of involvement are macules or patches with a decrease or absence of pigmentation. Patients with vitiligo also have an increased incidence of several autoimmune disorders, including Hashimoto’s thyroiditis, Graves’ disease, pernicious anemia, Addison’s disease, uveitis, alopecia areata, chronic mucocutaneous candidiasis, and the autoimmune polyendocrine syndromes (types I and II). Diseases of the thyroid gland are the most frequently associated disorders, occurring in up to 30% of patients with vitiligo. Circulating autoantibodies are often found, and the most common ones are antithyroglobulin, antimicrosomal, and antithyroid-stimulating hormone receptor antibodies.

There are four systemic diseases that should be considered in a patient with skin findings suggestive of vitiligo—*Vogt-Koyanagi-Harada syndrome*, *systemic sclerosis*, *onchocerciasis*, and *melanoma-associated leukoderma*. A history of aseptic meningitis, nontraumatic uveitis, tinnitus, hearing loss, and/or dysacusia points to the diagnosis of the Vogt-Koyanagi-Harada syndrome. In these patients, the face and scalp are the most common locations of pigment loss. The

TABLE 72-10 HYPOPIGMENTATION (PRIMARY CUTANEOUS DISORDERS, LOCALIZED)

	Clinical Characteristics	Wood's Lamp Examination (UV-A; Peak = 365 nm)	Skin Biopsy Specimen	Pathogenesis	Treatment
Idiopathic guttate hypomelanosis	Common; acquired; 1–4 mm in diameter Shins and extensor forearms	Less enhancement than vitiligo	Abrupt decrease in epidermal melanin content	Possible somatic mutations as a reflection of aging or UV exposure	None
Postinflammatory hypopigmentation	Can develop within active lesions, as in subacute cutaneous lupus, or after the lesion fades, as in atopic dermatitis	Depends on particular disease Usually less enhancement than in vitiligo	Type of inflammatory infiltrate depends on specific disease	Block in transfer of melanin from melanocytes to keratinocytes could be secondary to edema or decrease in contact time Destruction of melanocytes if inflammatory cells attack basal layer of epidermis	Treat underlying inflammatory disease
Pityriasis (tinea) versicolor	Common disorder Upper trunk and neck (shawl-like distribution), groin Young adults Macules have fine white scale when scratched	Golden fluorescence	Hyphal forms and budding yeast in stratum corneum	Invasion of stratum corneum by the yeast <i>Malassezia</i> Yeast is lipophilic and produces C ₉ and C ₁₁ dicarboxylic acids, which in vitro inhibit tyrosinase	Selenium sulfide 2.5%; topical imidazoles; oral triazoles
Vitiligo	Acquired; progressive Symmetric areas of complete pigment loss Periorificial—around mouth, nose, eyes, nipples, umbilicus, anus Other areas—flexor wrists, extensor distal extremities Segmental form is less common—unilateral, dermatomal-like	More apparent Chalk-white	Absence of melanocytes Mild inflammation	Autoimmune phenomenon that results in destruction of melanocytes—primarily cellular (circulating skin-homing autoreactive T cells)	Topical glucocorticoids; topical calcineurin inhibitors; NBUV-B; PUVA; transplants, if stable; depigmentation (topical MBEH), if widespread
Chemical- or drug-induced leukoderma	Similar appearance to vitiligo Often begins on hands when associated with chemical exposure Satellite lesions in areas not exposed to chemicals	More apparent Chalk-white	Decreased number or absence of melanocytes	Exposure to chemicals that selectively destroy melanocytes, in particular phenols and catechols (germicides; adhesives) or ingestion of drugs such as imatinib Release of cellular antigens and activation of circulating lymphocytes may explain satellite phenomenon Possible inhibition of KIT receptor	Avoid exposure to offending agent, then treat as vitiligo Drug-induced variant may undergo repigmentation when medication is discontinued
Piebaldism	Autosomal dominant Congenital, stable White forelock Areas of amelanosis contain normally pigmented and hyperpigmented macules of various sizes Symmetric involvement of central forehead, ventral trunk, and mid regions of upper and lower extremities	Enhancement of leukoderma and hyperpigmented macules	Amelanotic areas—few to no melanocytes	Defect in migration of melanoblasts from neural crest to involved skin or failure of melanoblasts to survive or differentiate in these areas Mutations within the <i>c-kit</i> protooncogene that encodes the tyrosine kinase receptor for stem cell growth factor (kit ligand)	None; occasionally transplants

Abbreviations: MBEH, monobenzylether of hydroquinone; NBUV-B, narrow band ultraviolet B; PUVA, psoralens + ultraviolet A irradiation.

vitiligo-like leukoderma seen in patients with systemic sclerosis has a clinical resemblance to idiopathic vitiligo that has begun to repigment as a result of treatment; that is, perifollicular macules of normal pigmentation are seen within areas of depigmentation. The basis of this leukoderma is unknown; there is no evidence of inflammation in areas of involvement, but it can resolve if the underlying connective tissue disease becomes inactive. In contrast to idiopathic vitiligo, melanoma-associated leukoderma often begins on the trunk, and its appearance, if spontaneous, should prompt a search for metastatic disease. It is also seen in patients undergoing immunotherapy for melanoma, including ipilimumab, with cytotoxic T lymphocytes presumably recognizing cell surface antigens common to melanoma cells and melanocytes, and is associated with a greater likelihood of a clinical response.

There are two systemic disorders (neurocristopathies) that may have the cutaneous findings of piebaldism (Table 72-9). They are *Shah-Waardenburg syndrome* and *Waardenburg syndrome*. A possible explanation for both disorders is an abnormal embryonic migration or survival of two neural crest-derived elements, one of them being melanocytes and the other myenteric ganglion cells (leading to Hirschsprung disease in Shah-Waardenburg syndrome) or auditory nerve cells (Waardenburg syndrome). The latter syndrome is characterized by congenital sensorineural hearing loss, dystopia canthorum (lateral displacement of the inner canthi but normal interpupillary distance), heterochromic irises, and a broad nasal root, in addition to the piebaldism. The facial dysmorphism can be explained by the neural crest origin of the connective tissues of the head and neck. Patients with Waardenburg syndrome have been shown to have mutations in four genes, including *PAX-3* and *MITF*, all of which encode DNA-binding proteins, whereas patients with Hirschsprung disease plus white spotting have mutations in one of three genes—endothelin 3, endothelin B receptor, and *SOX-10*.

In *tuberous sclerosis*, the earliest cutaneous sign is macular hypomelanosis, referred to as an ash leaf spot. These lesions are often present at birth and are usually multiple; however, detection may require Wood's lamp examination, especially in fair-skinned individuals. The pigment within them is reduced, but not absent. The average size is 1–3 cm, and the common shapes are polygonal and lance-ovate. Examination of the patient for additional cutaneous signs such as multiple angiofibromas of the face (adenoma sebaceum), unguinal and gingival fibromas, fibrous plaques of the forehead, and connective tissue nevi (shagreen patches) is recommended. It is important to remember that an ash leaf spot on the scalp will result in a circumscribed patch of lightly pigmented hair. Internal manifestations include seizures, mental retardation, central nervous system (CNS) and retinal hamartomas, pulmonary lymphangioleiomyomatosis (women), renal angiomyolipomas, and cardiac rhabdomyomas. The latter can be detected in up to 60% of children (<18 years) with tuberous sclerosis by echocardiography.

Nevus depigmentosus is a stable, well-circumscribed hypomelanosis that is present at birth. There is usually a single oval or rectangular lesion, but when there are multiple lesions, the possibility of tuberous sclerosis needs to be considered. In *linear nevoid hypopigmentation*, a term that is replacing hypomelanosis of Ito and segmental or systematized nevus depigmentosus, streaks and swirls of hypopigmentation are observed. Up to a third of patients in a tertiary care setting had associated abnormalities involving the musculoskeletal system (asymmetry), the CNS (seizures and mental retardation), and the eyes (strabismus and hypertelorism). Chromosomal mosaicism has been detected in these patients, lending support to the hypothesis that the cutaneous pattern is the result of the migration of two clones of primordial melanocytes, each with a different pigment potential.

Localized areas of decreased pigmentation are commonly seen as a result of cutaneous inflammation (Table 72-10) and have been observed in the skin overlying active lesions of sarcoidosis (see “Papulonodular Skin Lesions,” below) as well as in CTCL. Cutaneous infections also present as disorders of hypopigmentation, and in *tuberculoid leprosy*, there are a few asymmetric patches of hypomelanosis that have associated anesthesia, anhidrosis, and alopecia. Biopsy specimens of the palpable border show dermal granulomas that contain rare, if any, *Mycobacterium leprae* organisms.

HYPERPIGMENTATION

(Table 72-11) Disorders of hyperpigmentation are also divided into two groups—localized and diffuse. The localized forms are due to an epidermal alteration, a proliferation of melanocytes, or an increase in pigment production. Both seborrheic keratoses and acanthosis nigricans belong to the first group. *Seborrheic keratoses* are common lesions, but in one rare clinical setting, they are a sign of systemic disease, and that setting is the sudden appearance of multiple lesions, often with an inflammatory base and in association with acrochordons (skin tags) and acanthosis nigricans. This is termed the *sign of Leser-Trélat* and alerts the clinician to search for an internal malignancy. *Acanthosis nigricans* can also be a reflection of an internal malignancy, most commonly of the gastrointestinal tract, and it appears as velvety hyperpigmentation, primarily in flexural areas. However, in the majority of patients, acanthosis nigricans is associated with obesity and insulin resistance, although it may be a reflection of an endocrinopathy such as acromegaly, Cushing's syndrome, polycystic ovary syndrome, or insulin-resistant diabetes mellitus (type A, type B, and lipodystrophic forms).

A proliferation of melanocytes results in the following pigmented lesions: *lentigo*, *melanocytic nevus*, and *melanoma* (Chap. 105). In an adult, the majority of lentigines are related to sun exposure, which explains their distribution. However, in the Peutz-Jeghers and LEOPARD (lentigines; ECG abnormalities, primarily conduction defects; ocular hypertelorism; pulmonary stenosis and subaortic valvular stenosis; abnormal genitalia [cryptorchidism, hypospadias]; retardation of growth; and deafness [sensorineural]) syndromes, lentigines do serve as a clue to systemic disease. In *LEOPARD syndrome*, hundreds of lentigines develop during childhood and are scattered over the entire surface of the body. The lentigines in patients with *Peutz-Jeghers syndrome* are located primarily around the nose and mouth, on the hands and feet, and within the oral cavity. While the pigmented macules on the face may fade with age, the oral lesions persist. However, similar intraoral lesions are also seen in Addison's disease, in Laugier-Hunziker syndrome (no internal manifestations), and as a normal finding in darkly pigmented individuals. Patients with this autosomal dominant syndrome (due to mutations in a novel serine threonine kinase gene) have multiple benign polyps of the gastrointestinal tract, testicular or ovarian tumors, and an increased risk of developing gastrointestinal (primarily colon) and pancreatic cancers.

In the *Carney complex*, numerous lentigines are also seen, but they are in association with cardiac myxomas. This autosomal dominant disorder is also known as the *LAMB* (lentigines, atrial myxomas, mucocutaneous myxomas, and blue nevi) syndrome or *NAME* (nevi, atrial myxoma, myxoid neurofibroma, and ephelides [freckles]) syndrome. These patients can also have evidence of endocrine overactivity in the form of Cushing's syndrome (pigmented nodular adrenocortical disease) and acromegaly.

The third type of localized hyperpigmentation is due to a local increase in pigment production, and it includes *ephelides* and *café au lait macules* (CALMs). While a single CALM can be seen in up to 10% of the normal population, the presence of multiple or large-sized CALMs raises the possibility of an associated genodermatosis, e.g., neurofibromatosis (NF) or McCune-Albright syndrome. CALMs are flat, uniformly brown in color (usually two shades darker than uninvolved skin), and can vary in size from 0.5–12 cm. Approximately 80–90% of adult patients with *type I NF* will have six or more CALMs measuring ≥ 1.5 cm in diameter. Additional findings are discussed in the section on neurofibromas (see “Papulonodular Skin Lesions,” below). In comparison with NF, the CALMs in patients with *McCune-Albright syndrome* (polyostotic fibrous dysplasia with precocious puberty in females due to mosaicism for an activating mutation in a G protein [G_{α}] gene) are usually larger, are more irregular in outline, and tend to respect the midline.

In incontinentia pigmenti, dyskeratosis congenita, and bleomycin pigmentation, the areas of localized hyperpigmentation form a pattern—swirled in the first, reticulated in the second, and flagellate in the third. In *dyskeratosis congenita*, atrophic reticulated

TABLE 72-11 CAUSES OF HYPERPIGMENTATION

- I. Primary cutaneous disorders
 - A. Localized
 1. Epidermal alteration
 - a. Seborrheic keratosis
 - b. Pigmented actinic keratosis
 2. Proliferation of melanocytes
 - a. Lentigo
 - b. Melanocytic nevus (mole)
 - c. Melanoma
 3. Increased pigment production
 - a. Ephelide (freckle)
 - b. Café au lait macule
 - c. Postinflammatory hyperpigmentation
 - B. Localized and diffuse
 1. Drugs (e.g., minocycline, hydroxychloroquine, bleomycin)
- II. Systemic diseases
 - A. Localized
 1. Epidermal alteration
 - a. Seborrheic keratoses (sign of Leser-Trélat)
 - b. Acanthosis nigricans (insulin resistance, other endocrine disorders, paraneoplastic)
 2. Proliferation of melanocytes
 - a. Lentiginos (Peutz-Jeghers and LEOPARD syndromes; xeroderma pigmentosum)
 - b. Melanocytic nevi (Carney complex [LAMB and NAME syndromes])^a
 3. Increased pigment production
 - a. Café au lait macules (neurofibromatosis, McCune-Albright syndrome^b)
 - b. Urticaria pigmentosa^c
 4. Dermal pigmentation
 - a. Incontinentia pigmenti (stage III)
 - b. Dyskeratosis congenita
 - B. Diffuse
 1. Endocrinopathies
 - a. Addison's disease
 - b. Nelson syndrome
 - c. Ectopic ACTH syndrome
 - d. Hyperthyroidism
 2. Metabolic
 - a. Porphyria cutanea tarda
 - b. Hemochromatosis
 - c. Vitamin B₁₂/folate deficiency
 - d. Pellagra
 - e. Malabsorption, including Whipple's disease
 3. Melanosis secondary to metastatic melanoma
 4. Autoimmune
 - a. Biliary cirrhosis
 - b. Systemic sclerosis (scleroderma)
 - c. POEMS syndrome
 - d. Eosinophilia-myalgia syndrome^d
 5. Drugs and metals (e.g., arsenic)

^aAlso lentiginos. ^bPolyostotic fibrous dysplasia. ^cSee also "Papulonodular Skin Lesions." ^dLate 1980s.

Abbreviations: LAMB, lentiginos, atrial myxomas, mucocutaneous myxomas, and blue nevi; LEOPARD, lentiginos, ECG abnormalities, ocular hypertelorism, pulmonary stenosis and subaortic valvular stenosis, abnormal genitalia, retardation of growth, and deafness (sensorineural); NAME, nevi, atrial myxoma, myxoid neurofibroma, and ephelides (freckles); POEMS, polyneuropathy, organomegaly, endocrinopathies, M-protein, and skin changes.

hyperpigmentation is seen on the neck, trunk, and thighs and is accompanied by nail dystrophy, pancytopenia, and leukoplakia of the oral and anal mucosae. The latter often develops into squamous cell carcinoma. In addition to the flagellate pigmentation (linear streaks) on the trunk, patients receiving bleomycin often have hyperpigmentation overlying the elbows, knees, and small joints of the hand.

Localized hyperpigmentation is seen as a side effect of several other *systemic medications*, including those that produce fixed drug reactions (nonsteroidal anti-inflammatory drugs [NSAIDs], sulfonamides, barbiturates, and tetracyclines) and those that can complex with melanin (antimalarials) or iron (minocycline). Fixed drug eruptions recur in the exact same location as circular areas of erythema that can become bullous and then resolve as brown macules. The eruption usually appears within hours of administration of the offending agent, and common locations include the genitalia, distal extremities, and perioral region. Chloroquine and hydroxychloroquine produce gray-brown to blue-black discoloration of the shins, hard palate, and face, while blue macules (often misdiagnosed as bruises) can be seen on the lower extremities and in sites of inflammation with prolonged minocycline administration. Estrogen in oral contraceptives can induce melasma—symmetric brown patches on the face, especially the cheeks, upper lip, and forehead. Similar changes are seen in pregnancy and in patients receiving phenytoin.

In the diffuse forms of hyperpigmentation, the darkening of the skin may be of equal intensity over the entire body or may be accentuated in sun-exposed areas. The causes of diffuse hyperpigmentation can be divided into four major groups—endocrine, metabolic, autoimmune, and drugs. The endocrinopathies that frequently have associated hyperpigmentation include *Addison's disease*, *Nelson syndrome*, and *ectopic ACTH syndrome*. In these diseases, the increased pigmentation is diffuse but is accentuated in sun-exposed areas, the palmar creases, sites of friction, and scars. An overproduction of the pituitary hormones α -MSH (melanocyte-stimulating hormone) and ACTH can lead to an increase in melanocyte activity. These peptides are products of the proopiomelanocortin gene and exhibit homology; e.g., α -MSH and ACTH share 13 amino acids. A minority of patients with Cushing's disease or hyperthyroidism have generalized hyperpigmentation.

The metabolic causes of hyperpigmentation include *porphyria cutanea tarda* (PCT), *hemochromatosis*, *vitamin B₁₂ deficiency*, *folate deficiency*, *pellagra*, and *malabsorption*, including *Whipple's disease*. In patients with PCT (see "Vesicles/Bullae," below), the skin darkening is seen in sun-exposed areas and is a reflection of the photoreactive properties of porphyrins. The increased level of iron in the skin of patients with type 1 hemochromatosis stimulates melanin pigment production and leads to the classic bronze color. Patients with pellagra have a brown discoloration of the skin, especially in sun-exposed areas, as a result of nicotinic acid (niacin) deficiency. In the areas of increased pigmentation, there is a thin, varnish-like scale. These changes are also seen in patients who are vitamin B₆ deficient, have functioning carcinoid tumors (increased consumption of niacin), or take isoniazid. Approximately 50% of the patients with Whipple's disease have an associated generalized hyperpigmentation in association with diarrhea, weight loss, arthritis, and lymphadenopathy. A diffuse, slate-blue to gray-brown color is seen in patients with *melanosis secondary to metastatic melanoma*. The color reflects widespread deposition of melanin within the dermis as a result of the high concentration of circulating melanin precursors.

Of the autoimmune diseases associated with diffuse hyperpigmentation, *biliary cirrhosis* and *systemic sclerosis* are the most common, and occasionally, both disorders are seen in the same patient. The skin is dark brown in color, especially in sun-exposed areas. In biliary cirrhosis, the hyperpigmentation is accompanied by pruritus, jaundice, and xanthomas, whereas in systemic sclerosis, it is accompanied by sclerosis of the extremities, face, and, less commonly, the trunk. Additional clues to the diagnosis of systemic sclerosis are mat and periungual telangiectasias, calcinosis cutis, Raynaud's phenomenon, and distal ulcerations (see "Telangiectasias," above). The differential diagnosis of cutaneous sclerosis with hyperpigmentation includes

the POEMS (polyneuropathy; organomegaly [liver, spleen, lymph nodes]; endocrinopathies [impotence, gynecomastia]; *M*-protein; and skin changes) syndrome. The skin changes include hyperpigmentation, induration, hypertrichosis, angiomas, clubbing, and facial lipatrophy.

Diffuse hyperpigmentation that is due to drugs or metals can result from one of several mechanisms—induction of melanin pigment formation, complexing of the drug or its metabolites to melanin, and deposits of the drug in the dermis. Busulfan, cyclophosphamide, 5-fluorouracil, and inorganic arsenic induce pigment production. Complexes containing melanin or iron plus the drug or its metabolites are seen in patients receiving minocycline, and a diffuse, blue-gray, muddy appearance within sun-exposed areas may develop, in addition to pigmentation of the mucous membranes, teeth, nails, bones, and thyroid. Administration of amiodarone can result in both a phototoxic eruption (exaggerated sunburn) and/or a slate-gray to violaceous discoloration of sun-exposed skin. Biopsy specimens of the latter show yellow-brown granules in dermal macrophages, which represent intralysosomal accumulations of lipids, amiodarone, and its metabolites. Actual deposits of a particular drug or metal in the skin are seen with silver (argyria), where the skin appears blue-gray in color; gold (chrysi-asis), where the skin has a brown to blue-gray color; and clofazimine, where the skin appears reddish brown. The associated pigmentation is accentuated in sun-exposed areas, and discoloration of the eye is seen with gold (sclerae) and clofazimine (conjunctivae).

VESICLES/BULLAE

(Table 72-12) Depending on their size, cutaneous blisters are referred to as *vesicles* (<1 cm) or *bullae* (>1 cm). The primary autoimmune blistering disorders include *pemphigus vulgaris*, *pemphigus foliaceus*, *paraneoplastic pemphigus*, *bullous pemphigoid*, *gestational pemphigoid*, *cicatricial pemphigoid*, *epidermolysis bullosa acquisita*, *linear IgA bullous dermatosis (LABD)*, and *dermatitis herpetiformis* (Chap. 73).

Vesicles and bullae are also seen in *contact dermatitis*, both allergic and irritant forms (Chap. 71). When there is a linear arrangement of vesicular lesions, an exogenous cause or herpes zoster should be suspected. Bullous disease secondary to the ingestion of drugs can take one of several forms, including phototoxic eruptions, isolated bullae, Stevens-Johnson syndrome (SJS), and toxic epidermal necrolysis (TEN) (Chap. 74). Clinically, phototoxic eruptions resemble an exaggerated sunburn with diffuse erythema and bullae in sun-exposed areas. The most commonly associated drugs are doxycycline, quinolones, thiazides, NSAIDs, voriconazole, and psoralens. The development of a phototoxic eruption is dependent on the doses of both the drug and ultraviolet (UV)-A irradiation.

Toxic epidermal necrolysis is characterized by bullae that arise on widespread areas of tender erythema and then slough. This results in large areas of denuded skin. The associated morbidity, such as sepsis, and mortality rates are relatively high and are a function of the extent of epidermal necrosis. In addition, these patients may also have involvement of the mucous membranes and respiratory and intestinal tracts. Drugs are the primary cause of TEN, and the most common offenders are aromatic anticonvulsants (phenytoin, barbiturates, carbamazepine), sulfonamides, aminopenicillins, allopurinol, and NSAIDs. Severe acute graft-versus-host disease (grade 4), vancomycin-induced LABD, and the acute syndrome of apoptotic pan-epidermolysis (ASAP) in patients with lupus can also resemble TEN.

In *erythema multiforme* (EM), the primary lesions are pink-red macules and edematous papules, the centers of which may become vesicular. In contrast to a morbilliform exanthem, the clue to the diagnosis of EM, and especially SJS, is the development of a “dusky” violet color in the center of the lesions. Target lesions are also characteristic of EM and arise as a result of active centers and borders in combination with centrifugal spread. However, target lesions need not be present to make the diagnosis of EM.

EM has been subdivided into two major groups: (1) EM minor due to herpes simplex virus (HSV) and (2) EM major due to HSV; *Mycoplasma pneumoniae*; or, occasionally, drugs. Involvement of the

TABLE 72-12 CAUSES OF VESICLES/BULLAE

I. Primary mucocutaneous diseases
A. Primary blistering diseases (autoimmune)
1. Pemphigus, foliaceus and vulgaris ^a
2. Bullous pemphigoid ^b
3. Gestational pemphigoid ^b
4. Cicatricial pemphigoid ^b
5. Dermatitis herpetiformis ^{b,c}
6. Linear IgA bullous dermatosis ^b
7. Epidermolysis bullosa acquisita ^{b,d}
B. Secondary blistering diseases
1. Contact dermatitis ^{a,b}
2. Erythema multiforme ^e
3. Stevens-Johnson syndrome ^e
4. Toxic epidermal necrolysis ^e
C. Infections
1. Varicella-zoster virus ^{a,f}
2. Herpes simplex virus ^{a,f}
3. Enteroviruses, e.g., hand-foot-and-mouth disease ^f
4. Staphylococcal scalded-skin syndrome ^{a,g}
5. Bullous impetigo ^g
II. Systemic diseases
A. Autoimmune
1. Paraneoplastic pemphigus ^g
B. Infections
1. Cutaneous emboli ^h
C. Metabolic
1. Diabetic bullae ^{a,b}
2. Porphyria cutanea tarda ^b
3. Porphyria variegata ^b
4. Pseudoporphyria ^b
5. Bullous dermatosis of hemodialysis ^b
D. Ischemia
1. Coma bullae

^aIntraepidermal. ^bSubepidermal. ^cAssociated with gluten enteropathy. ^dAssociated with inflammatory bowel disease. ^eDegeneration of cells within the basal layer of the epidermis can give impression split is subepidermal. ^fAlso systemic. ^gIn adults, associated with renal failure and immunocompromised state.

mucous membranes (ocular, nasal, oral, and genital) is seen more commonly in the latter form. Hemorrhagic crusts of the lips are characteristic of EM major and SJS as well as herpes simplex, pemphigus vulgaris, and paraneoplastic pemphigus. Fever, malaise, myalgias, sore throat, and cough may precede or accompany the eruption. The lesions of EM usually resolve over 2–4 weeks but may be recurrent, especially when due to HSV. In addition to HSV (in which lesions usually appear 7–12 days after the viral eruption), EM can also follow vaccinations, radiation therapy, and exposure to environmental toxins, including the oleoresin in poison ivy.

Induction of SJS is most often due to drugs, especially sulfonamides, phenytoin, barbiturates, lamotrigine, aminopenicillins, nonnucleoside reverse transcriptase inhibitors (e.g., nevirapine), and carbamazepine. Widespread dusky macules and significant mucosal involvement are characteristic of SJS, and the cutaneous lesions may or may not develop epidermal detachment. If the latter occurs, by definition, it is limited to <10% of the body surface area (BSA). Greater involvement leads to the diagnosis of SJS/TEN overlap (10–30% BSA) or TEN (>30% BSA).

In addition to primary blistering disorders and hypersensitivity reactions, bacterial and viral infections can lead to vesicles and bullae. The most common infectious agents are HSV (Chap. 216), varicella-zoster virus (Chap. 217), and *S. aureus* (Chap. 172).

Staphylococcal scalded-skin syndrome (SSSS) and *bullous impetigo* are two blistering disorders associated with staphylococcal (phage group II) infection. In SSSS, the initial findings are redness and tenderness of the central face, neck, trunk, and intertriginous zones. This is followed by short-lived flaccid bullae and a slough or exfoliation of the superficial epidermis. Crusted areas then develop, characteristically around the mouth in a radial pattern. SSSS is distinguished from TEN by the following features: younger age group (primarily infants), more superficial site of blister formation, no oral lesions, shorter course, lower morbidity and mortality rates, and an association with staphylococcal exfoliative toxin (“exfoliatin”), not drugs. A rapid diagnosis of SSSS versus TEN can be made by a frozen section of the blister roof or exfoliative cytology of the blister contents. In SSSS, the site of staphylococcal infection is usually extracutaneous (conjunctivitis, rhinorrhea, otitis media, pharyngitis, tonsillitis), and the cutaneous lesions are sterile, whereas in bullous impetigo, the skin lesions are the site of infection. Impetigo is more localized than SSSS and usually presents with honey-colored crusts. Occasionally, superficial purulent blisters also form. *Cutaneous emboli* from gram-negative infections may present as isolated bullae, but the base of the lesion is purpuric or necrotic, and it may develop into an ulcer (see “Purpura,” below).

Several metabolic disorders are associated with blister formation, including diabetes mellitus, renal failure, and porphyria. Local hypoxemia secondary to decreased cutaneous blood flow can also produce blisters, which explains the presence of bullae over pressure points in comatose patients (coma bullae). In *diabetes mellitus*, tense bullae with clear sterile viscous fluid arise on normal skin. The lesions can be as large as 6 cm in diameter and are located on the distal extremities. There are several types of porphyria, but the most common form with cutaneous findings is *porphyria cutanea tarda* (PCT). In sun-exposed areas (primarily the face and hands), the skin is very fragile, with trauma leading to erosions mixed with tense vesicles. These lesions then heal with scarring and formation of milia; the latter are firm, 1- to 2-mm white or yellow papules that represent epidermoid inclusion cysts. Associated findings can include hypertrichosis of the lateral malar region (men) or face (women) and, in sun-exposed areas, hyperpigmentation and firm sclerotic plaques. An elevated level of urinary uroporphyrins confirms the diagnosis and is due to a decrease in uroporphyrinogen decarboxylase activity. PCT can be exacerbated by alcohol, hemochromatosis and other forms of iron overload, chlorinated hydrocarbons, hepatitis C and HIV infections, and hepatomas.

The differential diagnosis of PCT includes (1) *porphyria variegata*—the skin signs of PCT plus the systemic findings of acute intermittent porphyria; it has a diagnostic plasma porphyrin fluorescence emission at 626 nm; (2) *drug-induced pseudoporphyria*—the clinical and histologic findings are similar to PCT, but porphyrins are normal; etiologic agents include naproxen and other NSAIDs, furosemide, tetracycline, and voriconazole; (3) *bullous dermatosis of hemodialysis*—the same appearance as PCT, but porphyrins are usually normal or occasionally borderline elevated; patients have chronic renal failure and are on hemodialysis; (4) PCT associated with hepatomas and hemodialysis; and (5) *epidermolysis bullosa acquisita* (Chap. 73).

EXANTHEMS

(Table 72-13) Exanthems are characterized by an acute generalized eruption. The most common presentation is erythematous macules and papules (morbilliform) and less often confluent blanching erythema (scarlatiniform). *Morbilliform* eruptions are usually due to either drugs or viral infections. For example, up to 5% of patients receiving penicillins, sulfonamides, phenytoin, or nevirapine will develop a maculopapular eruption. Accompanying signs may include pruritus, fever, eosinophilia, and transient lymphadenopathy. Similar maculopapular eruptions are seen in the classic childhood viral exanthems, including (1) *rubeola* (measles)—a prodrome of coryza, cough, and conjunctivitis followed by Koplik’s spots on the buccal mucosa; the eruption begins behind the ears, at the hairline, and on the forehead and then spreads down the body, often becoming confluent; (2) *rubella*—the eruption begins on the forehead and face

TABLE 72-13 CAUSES OF EXANTHEMS

I. Morbilliform
A. Drugs
B. Viral
1. Rubeola (measles)
2. Rubella
3. Erythema infectiosum (erythema of cheeks; reticulated on extremities)
4. Epstein-Barr virus, echovirus, coxsackievirus, CMV, adenovirus, HHV-6/HHV-7 ^a , dengue virus, and West Nile virus infections
5. HIV seroconversion exanthem (plus mucosal ulcerations)
C. Bacterial
1. Typhoid fever
2. Early secondary syphilis
3. Early <i>Rickettsia</i> infections
4. Early meningococemia
5. Ehrlichiosis
D. Acute graft-versus-host disease
E. Kawasaki disease
II. Scarlatiniform
A. Scarlet fever
B. Toxic shock syndrome
C. Kawasaki disease
D. Early staphylococcal scalded-skin syndrome

^aPrimary infection in infants and reactivation in the setting of immunosuppression.

Abbreviations: CMV, cytomegalovirus; HHV, human herpesvirus; HIV, human immunodeficiency virus.

and then spreads down the body; it resolves in the same order and is associated with retroauricular and suboccipital lymphadenopathy; and (3) *erythema infectiosum* (fifth disease)—erythema of the cheeks is followed by a reticulated pattern on the extremities; it is secondary to a parvovirus B19 infection, and an associated arthritis is seen in adults.

Both measles and rubella can occur in unvaccinated adults, and an atypical form of measles is seen in adults immunized with either killed measles vaccine or killed vaccine followed in time by live vaccine. In contrast to classic measles, the eruption of atypical measles begins on the palms, soles, wrists, and ankles, and the lesions may become purpuric. The patient with atypical measles can have pulmonary involvement and be quite ill. Rubelliform and roseoliform eruptions are also associated with *Epstein-Barr virus* (5–15% of patients), *echovirus*, *coxsackievirus*, *cytomegalovirus*, *adenovirus*, *dengue virus*, and *West Nile virus* infections. Detection of specific IgM antibodies or fourfold elevations in IgG antibodies allow the proper diagnosis, but polymerase chain reaction (PCR) is gradually replacing serologic assays. Occasionally, a maculopapular drug eruption is a reflection of an underlying viral infection. For example, ~95% of the patients with infectious mononucleosis who are given ampicillin will develop a rash.

Of note, early in the course of infections with *Rickettsia* and meningococcus, prior to the development of petechiae and purpura, the lesions may be erythematous macules and papules. This is also the case in chickenpox prior to the development of vesicles. Maculopapular eruptions are associated with early *HIV infection*, early secondary *syphilis*, *typhoid fever*, and *acute graft-versus-host disease*. In the last, lesions frequently begin on the dorsal hands and forearms; the macular rose spots of typhoid fever involve primarily the anterior trunk.

The prototypic *scarlatiniform* eruption is seen in *scarlet fever* and is due to an erythrogenic toxin produced by bacteriophage-containing group A β -hemolytic streptococci, most commonly in the setting of pharyngitis. This eruption is characterized by diffuse erythema, which begins on the neck and upper trunk, and red follicular puncta. Additional findings include a white strawberry tongue (white coating with red papillae) followed by a red strawberry tongue (red tongue

with red papillae); petechiae of the palate; a facial flush with circumoral pallor; linear petechiae in the antecubital fossae; and desquamation of the involved skin, palms, and soles 5–20 days after onset of the eruption. A similar desquamation of the palms and soles is seen in toxic shock syndrome (TSS), in Kawasaki disease, and after severe febrile illnesses. Certain strains of staphylococci also produce an erythrogenic toxin that leads to the same clinical findings as in streptococcal scarlet fever, except that the anti-streptolysin O or -DNase B titers are not elevated.

In *toxic shock syndrome*, staphylococcal (phage group I) infections produce an exotoxin (TSST-1) that causes the fever and rash as well as enterotoxins. Initially, the majority of cases were reported in menstruating women who were using tampons. However, other sites of infection, including wounds and nasal packing, can lead to TSS. The diagnosis of TSS is based on clinical criteria (Chap. 172), and three of these involve mucocutaneous sites (diffuse erythema of the skin, desquamation of the palms and soles 1–2 weeks after onset of illness, and involvement of the mucous membranes). The latter is characterized as hyperemia of the vagina, oropharynx, or conjunctivae. Similar systemic findings have been described in *streptococcal toxic shock syndrome* (Chap. 173), and although an exanthem is seen less often than in TSS due to a staphylococcal infection, the underlying infection is often in the soft tissue (e.g., cellulitis).

The cutaneous eruption in *Kawasaki disease* (Chap. 385) is polymorphous, but the two most common forms are morbilliform and scarlatiniform. Additional mucocutaneous findings include bilateral conjunctival injection; erythema and edema of the hands and feet followed by desquamation; and diffuse erythema of the oropharynx, red strawberry tongue, and dry fissured lips. This clinical picture can resemble TSS and scarlet fever, but clues to the diagnosis of Kawasaki disease are cervical lymphadenopathy, cheilitis, and thrombocytosis. The most serious associated systemic finding in this disease is coronary aneurysms secondary to arteritis. Scarlatiniform eruptions are also seen in the early phase of SSSS (see “Vesicles/Bullae,” above), in young adults with *Arcanobacterium haemolyticum* infection, and as reactions to drugs.

URTICARIA

(Table 72-14) *Urticaria* (hives) are transient lesions that are composed of a central wheal surrounded by an erythematous halo or flare. Individual lesions are round, oval, or figurate and are often pruritic. Acute and chronic urticaria have a wide variety of allergic etiologies and reflect edema in the dermis. Urticarial lesions can also be seen in patients with mastocytosis (urticaria pigmentosa), hypo- or hyperthyroidism, and systemic-onset juvenile idiopathic arthritis (Still's disease). In both juvenile- and adult-onset Still's disease, the lesions

coincide with the fever spike, are transient, and are due to dermal infiltrates of neutrophils.

The common *physical urticarias* include dermatographism, solar urticaria, cold urticaria, and cholinergic urticaria. Patients with *dermatographism* exhibit linear wheals following minor pressure or scratching of the skin. It is a common disorder, affecting ~5% of the population. *Solar urticaria* characteristically occurs within minutes of sun exposure and is a skin sign of one systemic disease—erythropoietic protoporphyria. In addition to the urticaria, these patients have subtle pitted scarring of the nose and hands. *Cold urticaria* is precipitated by exposure to the cold, and therefore exposed areas are usually affected. In occasional patients, the disease is associated with abnormal circulating proteins—more commonly cryoglobulins and less commonly cryofibrinogens. Additional systemic symptoms include wheezing and syncope, thus explaining the need for these patients to avoid swimming in cold water. Autosomal dominantly inherited cold urticaria is associated with dysfunction of cryopyrin. *Cholinergic urticaria* is precipitated by heat, exercise, or emotion and is characterized by small wheals with relatively large flares. It is occasionally associated with wheezing.

Whereas urticarias are the result of dermal edema, subcutaneous edema leads to the clinical picture of *angioedema*. Sites of involvement include the eyelids, lips, tongue, larynx, and gastrointestinal tract as well as the subcutaneous tissue. Angioedema occurs alone or in combination with urticaria, including urticarial vasculitis and the physical urticarias. Both acquired and hereditary (autosomal dominant) forms of angioedema occur (Chap. 376), and in the latter, urticaria is rarely, if ever, seen.

Urticarial vasculitis is an immune complex disease that may be confused with simple urticaria. In contrast to simple urticaria, individual lesions tend to last longer than 24 h and usually develop central petechiae that can be observed even after the urticarial phase has resolved. The patient may also complain of burning rather than pruritus. On biopsy, there is a leukocytoclastic vasculitis of the small dermal blood vessels. Although many cases of urticarial vasculitis are idiopathic in origin, it can be a reflection of an underlying systemic illness such as lupus erythematosus, Sjögren's syndrome, or hereditary complement deficiency. There is a spectrum of urticarial vasculitis that ranges from purely cutaneous to multisystem involvement. The most common systemic signs and symptoms are arthralgias and/or arthritis, nephritis, and crampy abdominal pain, with asthma and chronic obstructive lung disease seen less often. Hypocomplementemia occurs in one- to two-thirds of patients, even in the idiopathic cases. Urticarial vasculitis can also be seen in patients with *hepatitis B* and *hepatitis C* infections, *serum sickness*, and *serum sickness-like illnesses* (e.g., due to cefaclor, minocycline).

PAPULONODULAR SKIN LESIONS

(Table 72-15) In the *papulonodular diseases*, the lesions are elevated above the surface of the skin and may coalesce to form larger plaques. The location, consistency, and color of the lesions are the keys to their diagnosis; this section is organized on the basis of color.

WHITE LESIONS

In *calcinosis cutis*, there are firm white to white-yellow papules with an irregular surface. When the contents are expressed, a chalky white material is seen. *Dystrophic calcification* is seen at sites of previous inflammation or damage to the skin. It develops in acne scars as well as on the distal extremities of patients with systemic sclerosis and in the subcutaneous tissue and intermuscular fascial planes in DM. The latter is more extensive and is more commonly seen in children. An elevated calcium phosphate product, most commonly due to secondary hyperparathyroidism in the setting of renal failure, can lead to nodules of *metastatic calcinosis cutis*, which tend to be subcutaneous and periarticular. These patients can also develop calcification of muscular arteries and subsequent ischemic necrosis (calciphylaxis). *Osteoma cutis*, in the form of small papules, most commonly occurs on the face

TABLE 72-14 CAUSES OF URTICARIA AND ANGIOEDEMA

- I. Primary cutaneous disorders
 - A. Acute and chronic urticaria^a
 - B. Physical urticaria
 1. Dermatographism
 2. Solar urticaria^b
 3. Cold urticaria^b
 4. Cholinergic urticaria^b
 - C. Angioedema (hereditary and acquired)^{b,c}
- II. Systemic diseases
 - A. Urticarial vasculitis
 - B. Hepatitis B or C infection
 - C. Serum sickness
 - D. Angioedema (hereditary and acquired)

^aA small minority develop anaphylaxis. ^bAlso systemic. ^cAcquired angioedema can be idiopathic, associated with a lymphoproliferative disorder, or due to a drug, e.g., angiotensin-converting enzyme (ACE) inhibitors.

TABLE 72-15 PAPULONODULAR SKIN LESIONS ACCORDING TO COLOR GROUPS

I. White	3. Leukemia cutis
A. Calcinosis cutis	4. Sweet syndrome
B. Osteoma cutis (also skin-colored or blue)	C. Nodules
II. Skin-colored	1. Panniculitis
A. Rheumatoid nodules	2. Medium-sized vessel vasculitis (e.g., cutaneous polyarteritis nodosa)
B. Neurofibromas (von Recklinghausen's disease)	D. Primary cutaneous disorders
C. Angiofibromas (tuberous sclerosis, MEN syndrome, type 1)	1. Arthropod bites
D. Neuromas (MEN syndrome, type 2b)	2. Cherry hemangiomas
E. Adnexal tumors	3. Infections, e.g., streptococcal cellulitis, sporotrichosis
1. Basal cell carcinomas (nevroid basal cell carcinoma syndrome)	4. Polymorphous light eruption
2. Tricholemmomas (Cowden disease)	5. Lymphocytoma cutis (pseudolymphoma)
F. Osteomas (arise in skull and jaw in Gardner syndrome)	VI. Red-brown ^b
G. Primary cutaneous disorders	A. Sarcoidosis
1. Epidermal inclusion cysts ^a	B. Urticaria pigmentosa
2. Lipomas	C. Erythema elevatum diutinum (chronic leukocytoclastic vasculitis)
III. Pink/translucent ^b	D. Lupus vulgaris
A. Amyloidosis, primary systemic	VII. Blue ^b
B. Papular mucinosis/scleromyxedema	A. Venous malformations (e.g., blue rubber bleb syndrome)
C. Multicentric reticulohistiocytosis	B. Primary cutaneous disorders
IV. Yellow	1. Venous lake
A. Xanthomas	2. Blue nevus
B. Tophi	VIII. Violaceous
C. Necrobiosis lipoidica	A. Lupus pernio (sarcoidosis)
D. Pseudoxanthoma elasticum	B. Lymphoma cutis
E. Sebaceous adenomas (Muir-Torre syndrome)	C. Cutaneous lupus
V. Red ^b	IX. Purple
A. Papules	A. Kaposi's sarcoma
1. Angiokeratomas (Fabry disease)	B. Angiosarcoma
2. Bacillary angiomatosis (primarily in AIDS)	C. Palpable purpura (see Table 72-16)
B. Papules/plaques	X. Brown-black ^c
1. Cutaneous lupus	XI. Any color
2. Lymphoma cutis	A. Metastases

^aIf multiple with childhood onset, consider Gardner syndrome. ^bMay have darker hue in more darkly pigmented individuals. ^cSee also "Hyperpigmentation."

Abbreviation: MEN, multiple endocrine neoplasia.

of individuals with a history of acne vulgaris, whereas plate-like lesions occur in rare genetic syndromes (**Chap. 82**).

SKIN-COLORED LESIONS

There are several types of skin-colored lesions, including epidermoid inclusion cysts, lipomas, rheumatoid nodules, neurofibromas, angiofibromas, neuromas, and adnexal tumors such as tricholemmomas. Both *epidermoid inclusion cysts* and *lipomas* are very common mobile subcutaneous nodules—the former are rubbery and drain cheese-like material (sebum and keratin) if incised. Lipomas are firm and somewhat lobulated on palpation. When extensive facial epidermoid inclusion cysts develop during childhood or there is a family history of such lesions, the patient should be examined for other signs of Gardner syndrome, including osteomas and desmoid tumors. *Rheumatoid nodules* are firm 0.5- to 4-cm nodules that favor the extensor aspect of joints, especially the elbows. They are seen in ~20% of patients with rheumatoid arthritis and 6% of patients with Still's disease. Biopsies of the nodules show palisading granulomas. Similar lesions that are smaller and shorter-lived are seen in rheumatic fever.

Neurofibromas (benign Schwann cell tumors) are soft papules or nodules that exhibit the "button-hole" sign; that is, they invaginate into the skin with pressure in a manner similar to a hernia. Single lesions are seen in normal individuals, but multiple neurofibromas, usually in combination with six or more CALMs measuring >1.5 cm (see "Hyperpigmentation," above), axillary freckling, and multiple Lisch

nodules, are seen in von Recklinghausen's disease (NF type I) (**Chap. 118**). In some patients, the neurofibromas are localized and unilateral due to somatic mosaicism.

Angiofibromas are firm pink to skin-colored papules that measure from 3 mm to a few centimeters in diameter. When multiple lesions are located on the central cheeks (adenoma sebaceum), the patient has tuberous sclerosis or multiple endocrine neoplasia (MEN) syndrome, type 1. The former is an autosomal disorder due to mutations in two different genes, and the associated findings are discussed in the section on ash leaf spots as well as in **Chap. 118**.

Neuromas (benign proliferations of nerve fibers) are also firm, skin-colored papules. They are more commonly found at sites of amputation and as rudimentary supernumerary digits. However, when there are multiple neuromas on the eyelids, lips, distal tongue, and/or oral mucosa, the patient should be investigated for other signs of the MEN syndrome, type 2b. Associated findings include marfanoid habitus, protuberant lips, intestinal ganglioneuromas, and medullary thyroid carcinoma (>75% of patients; **Chap. 408**).

Adnexal tumors are derived from pluripotent cells of the epidermis that can differentiate toward hair, sebaceous, apocrine, or eccrine glands or remain undifferentiated. *Basal cell carcinomas* (BCCs) are examples of adnexal tumors that have little or no evidence of differentiation. Clinically, they are translucent papules with rolled borders, telangiectasias, and central erosion. BCCs commonly arise in sun-damaged skin of the head and neck as well as the upper trunk. When a

patient has multiple BCCs, especially prior to age 30, the possibility of the nevoid basal cell carcinoma syndrome should be raised. It is inherited as an autosomal dominant trait and is associated with jaw cysts, palmar and plantar pits, frontal bossing, medulloblastomas, and calcification of the falx cerebri and diaphragma sellae. *Tricholemmomas* are also skin-colored adnexal tumors but differentiate toward hair follicles and can have a wartlike appearance. The presence of multiple tricholemmomas on the face and cobblestoning of the oral mucosa points to the diagnosis of Cowden disease (multiple hamartoma syndrome) due to mutations in the phosphatase and tensin homolog (*PTEN*) gene. Internal organ involvement (in decreasing order of frequency) includes fibrocystic disease and carcinoma of the breast, adenomas and carcinomas of the thyroid, and gastrointestinal polyposis. Keratoses of the palms, soles, and dorsal aspect of the hands are also seen.

PINK LESIONS

The cutaneous lesions associated with primary systemic amyloidosis are often pink in color and translucent. Common locations are the face, especially the periorbital and perioral regions, and flexural areas. On biopsy, homogeneous deposits of amyloid are seen in the dermis and in the walls of blood vessels; the latter lead to an increase in vessel wall fragility. As a result, petechiae and purpura develop in clinically normal skin as well as in lesional skin following minor trauma, hence the term *pinch purpura*. Amyloid deposits are also seen in the striated muscle of the tongue and result in macroglossia.

Even though specific mucocutaneous lesions are present in only ~30% of the patients with primary systemic (AL) amyloidosis, the diagnosis can be made via histologic examination of abdominal subcutaneous fat, in conjunction with a serum free light chain assay. By special staining, amyloid deposits are seen around blood vessels or individual fat cells in 40–50% of patients. There are also three forms of amyloidosis that are limited to the skin and that should not be construed as cutaneous lesions of systemic amyloidosis. They are macular amyloidosis (upper back), lichen amyloidosis (usually lower extremities), and nodular amyloidosis. In macular and lichen amyloidosis, the deposits are composed of altered epidermal keratin. Early-onset macular and lichen amyloidosis have been associated with MEN syndrome, type 2a.

Patients with *multicentric reticulohistiocytosis* also have pink-colored papules and nodules on the face and mucous membranes as well as on the extensor surface of the hands and forearms. They have a polyarthritis that can mimic rheumatoid arthritis clinically. On histologic examination, the papules have characteristic giant cells that are not seen in biopsies of rheumatoid nodules. Pink to skin-colored papules that are firm, 2–5 mm in diameter, and often in a linear arrangement are seen in patients with *papular mucinosis*. This disease is also referred to as *generalized lichen myxedematosus* or *scleromyxedema*. The latter name comes from the induration of the face and extremities that may accompany the papular eruption. Biopsy specimens of the papules show localized mucin deposition, and serum protein electrophoresis plus immunofixation electrophoresis demonstrates a monoclonal spike of IgG, usually with a λ light chain.

YELLOW LESIONS

Several systemic disorders are characterized by yellow-colored cutaneous papules or plaques—hyperlipidemia (xanthomas), gout (tophi), diabetes (necrobiosis lipoidica), pseudoxanthoma elasticum, and Muir-Torre syndrome (sebaceous tumors). Eruptive xanthomas are the most common form of *xanthomas* and are associated with hypertriglyceridemia (primarily hyperlipoproteinemia types I, IV, and V). Crops of yellow papules with erythematous halos occur primarily on the extensor surfaces of the extremities and the buttocks, and they spontaneously involute with a fall in serum triglycerides. Types II and III result in one or more of the following types of xanthoma: xanthelasma, tendon xanthomas, and plane xanthomas. Xanthelasma are found on the eyelids, whereas tendon xanthomas are frequently associated with the Achilles and extensor finger tendons; plane xanthomas are flat and favor the palmar creases, neck, upper trunk, and flexural folds. Tuberous xanthomas are frequently associated with hypertriglyceridemia, but they

are also seen in patients with hypercholesterolemia and are found most frequently over the large joints or hand. Biopsy specimens of xanthomas show collections of lipid-containing macrophages (foam cells).

Patients with several disorders, including biliary cirrhosis, can have a secondary form of hyperlipidemia with associated tuberous and plane xanthomas. However, patients with plasma cell dyscrasias have *normolipemic plane xanthomas*. This latter form of xanthoma may be ≥ 12 cm in diameter and is most frequently seen on the upper trunk or side of the neck. It is important to note that the most common setting for eruptive xanthomas is uncontrolled diabetes mellitus. The least specific sign for hyperlipidemia is xanthelasma, because at least 50% of the patients with this finding have normal lipid profiles.

In *tophaceous gout*, there are deposits of monosodium urate in the skin around the joints, particularly those of the hands and feet. Additional sites of *tophi* formation include the helix of the ear and the olecranon and prepatellar bursae. The lesions are firm, yellow in color, and occasionally discharge a chalky material. Their size varies from 1 mm to 7 cm, and the diagnosis can be established by polarized light microscopy of the aspirated contents of a lesion. Lesions of *necrobiosis lipoidica* are found primarily on the shins (90%), and patients can have diabetes mellitus or develop it subsequently. Characteristic findings include a central yellow color, atrophy (transparency), telangiectasias, and a red to red-brown border. Ulcerations can also develop within the plaques. Biopsy specimens show necrobiosis of collagen and granulomatous inflammation.

In *pseudoxanthoma elasticum* (PXE), due to mutations in the gene *ABCC6*, there is an abnormal deposition of calcium on the elastic fibers of the skin, eye, and blood vessels. In the skin, the flexural areas such as the neck, axillae, antecubital fossae, and inguinal area are the primary sites of involvement. Yellow papules coalesce to form reticulated plaques that have an appearance similar to that of plucked chicken skin. In severely affected skin, hanging, redundant folds develop. Biopsy specimens of involved skin show swollen and irregularly clumped elastic fibers with deposits of calcium. In the eye, the calcium deposits in Bruch's membrane lead to angioid streaks and choroiditis; in the arteries of the heart, kidney, gastrointestinal tract, and extremities, the deposits lead to angina, hypertension, gastrointestinal bleeding, and claudication, respectively.

Adnexal tumors that have differentiated toward sebaceous glands include sebaceous adenoma, sebaceous carcinoma, and sebaceous hyperplasia. Except for sebaceous hyperplasia, which is commonly seen on the face, these tumors are fairly rare. Patients with Muir-Torre syndrome have one or more *sebaceous adenoma(s)*, and they can also have sebaceous carcinomas and sebaceous hyperplasia as well as keratoacanthomas. The internal manifestations of Muir-Torre syndrome include *multiple* carcinomas of the gastrointestinal tract (primarily colon) as well as cancers of the larynx, genitourinary tract, and breast.

RED LESIONS

Cutaneous lesions that are red in color have a wide variety of etiologies; in an attempt to simplify their identification, they will be subdivided into papules, papules/plaques, and subcutaneous nodules. Common red papules include *arthropod bites* and *cherry hemangiomas*; the latter are small, bright-red, dome-shaped papules that represent a benign proliferation of capillaries. In patients with AIDS (Chap. 226), the development of multiple red hemangioma-like lesions points to bacillary angiomatosis, and biopsy specimens show clusters of bacilli that stain positive with the Warthin-Starry stain; the pathogens have been identified as *Bartonella henselae* and *Bartonella quintana*. Disseminated visceral disease is seen primarily in immunocompromised hosts but can occur in immunocompetent individuals.

Multiple *angiokeratomas* are seen in Fabry disease, an X-linked recessive lysosomal storage disease that is due to a deficiency of α -galactosidase A. The lesions are red to red-blue in color and can be quite small in size (1–3 mm), with the most common location being the lower trunk. Associated findings include chronic renal disease, peripheral neuropathy, and corneal opacities (cornea verticillata). Electron photomicrographs of angiokeratomas and clinically normal skin demonstrate lamellar lipid deposits in fibroblasts, pericytes, and endothelial

cells that are diagnostic of this disease. Widespread acute eruptions of erythematous papules are discussed in the section on exanthems.

There are several infectious diseases that present as erythematous papules or nodules in a lymphocutaneous or sporotrichoid pattern, i.e., in a linear arrangement along the lymphatic channels. The two most common etiologies are *Sporothrix schenckii* (sporotrichosis) and the atypical mycobacterium *Mycobacterium marinum*. The organisms are introduced as a result of trauma, and a primary inoculation site is often seen in addition to the lymphatic nodules. Additional causes include *Nocardia*, *Leishmania*, and other atypical mycobacteria and dimorphic fungi; culture of lesional tissue will aid in the diagnosis.

The diseases that are characterized by erythematous plaques with scale are reviewed in the papulosquamous section, and the various forms of dermatitis are discussed in the section on erythroderma. Additional disorders in the differential diagnosis of red papules/plaques include *cellulitis*, *polymorphous light eruption* (PMLE), *cutaneous lymphoid hyperplasia* (lymphocytoma cutis), *cutaneous lupus*, *lymphoma cutis*, and *leukemia cutis*. The first three diseases represent primary cutaneous disorders, although cellulitis may be accompanied by a bacteremia. PMLE is characterized by erythematous papules and plaques in a primarily sun-exposed distribution—dorsum of the hand, extensor forearm, and upper trunk. Lesions follow exposure to UV-B and/or UV-A, and in higher latitudes, PMLE is most severe in the late spring and early summer. A process referred to as “hardening” occurs with continued UV exposure, and the eruption fades, but in temperate climates, it will recur in the spring. PMLE must be differentiated from cutaneous lupus, and this is accomplished by observation of the natural history, histologic examination, and direct immunofluorescence of the lesions. Cutaneous lymphoid hyperplasia (pseudolymphoma) is a *benign* polyclonal proliferation of lymphocytes in the skin that presents as infiltrated pink-red to red-purple papules and plaques; it must be distinguished from lymphoma cutis.

Several types of red plaques are seen in patients with systemic *lupus*, including (1) erythematous urticarial plaques across the cheeks and nose in the classic butterfly rash; (2) erythematous discoid lesions with fine or “carpet-tack” scale, telangiectasias, central hypopigmentation, peripheral hyperpigmentation, follicular plugging, and atrophy located on the face, scalp, external ears, arms, and upper trunk; and (3) psoriasisiform or annular lesions of subacute cutaneous lupus with hypopigmented centers located primarily on the extensor arms and upper trunk. Additional mucocutaneous findings include (1) a violaceous flush on the face and V of the neck; (2) photosensitivity; (3) urticarial vasculitis (see “Urticaria,” above); (4) lupus panniculitis (see below); (5) diffuse alopecia; (6) alopecia secondary to discoid lesions; (7) periungual telangiectasias and erythema; (8) EM-like lesions that may become bullous; (9) oral ulcers; and (10) distal ulcerations secondary to Raynaud’s phenomenon, vasculitis, or livedoid vasculopathy. Patients with only discoid lesions usually have the form of lupus that is limited to the skin. However, up to 10% of these patients eventually develop systemic lupus. Direct immunofluorescence of involved skin, in particular discoid lesions, shows deposits of IgG or IgM and C3 in a granular distribution along the dermal-epidermal junction.

In *lymphoma cutis*, there is a proliferation of malignant lymphocytes in the skin, and the clinical appearance resembles that of cutaneous lymphoid hyperplasia—infiltrated pink-red to red-purple papules and plaques. Lymphoma cutis can occur anywhere on the surface of the skin, whereas the sites of predilection for lymphocytomas include the malar ridge, tip of the nose, and earlobes. Patients with non-Hodgkin’s lymphomas have specific cutaneous lesions more often than those with Hodgkin’s disease, and, occasionally, the skin nodules precede the development of extracutaneous non-Hodgkin’s lymphoma or represent the only site of involvement (e.g., primary cutaneous B cell lymphoma). Arcuate lesions are sometimes seen in lymphoma and lymphocytoma cutis as well as in CTCL. *Adult T cell leukemia/lymphoma* that develops in association with HTLV-1 infection is characterized by cutaneous plaques, hypercalcemia, and circulating CD25+ lymphocytes. *Leukemia cutis* has the same appearance as lymphoma cutis, and specific lesions are seen more commonly in monocytic leukemias than in lymphocytic or granulocytic leukemias. Cutaneous

chloromas (granulocytic sarcomas) may precede the appearance of circulating blasts in acute myelogenous leukemia and, as such, represent a form of leukemic leukemia cutis.

Sweet syndrome is characterized by pink-red to red-brown edematous plaques that are frequently painful and occur primarily on the head, neck, and upper (and, less often, lower) extremities. The patients also have fever, neutrophilia, and a dense dermal infiltrate of neutrophils in the lesions. In ~10% of the patients, there is an associated malignancy, most commonly acute myelogenous leukemia. Sweet syndrome has also been reported with inflammatory bowel disease, systemic lupus erythematosus, and solid tumors (primarily of the genitourinary tract) as well as drugs (e.g., all-*trans*-retinoic acid, granulocyte colony-stimulating factor [G-CSF]). The differential diagnosis includes neutrophilic eccrine hidradenitis; bullous forms of pyoderma gangrenosum; and, occasionally, cellulitis. Extracutaneous sites of involvement include joints, muscles, eye, kidney (proteinuria, occasionally glomerulonephritis), and lung (neutrophilic infiltrates). The idiopathic form of Sweet syndrome is seen more often in women, following a respiratory tract infection.

Common causes of erythematous subcutaneous nodules include inflamed epidermoid inclusion cysts, acne cysts, and furuncles. *Panniculitis*, an inflammation of the fat, also presents as subcutaneous nodules and is frequently a sign of systemic disease. There are several forms of panniculitis, including erythema nodosum, erythema induratum/nodular vasculitis, lupus panniculitis, lipodermatosclerosis, α_1 -antitrypsin deficiency, factitial, and fat necrosis secondary to pancreatic disease. Except for erythema nodosum, these lesions may break down and ulcerate or heal with a scar. The shin is the most common location for the nodules of erythema nodosum, whereas the calf is the most common location for lesions of erythema induratum. In erythema nodosum, the nodules are initially red but then develop a blue color as they resolve. Patients with erythema nodosum but no underlying systemic illness can still have fever, malaise, leukocytosis, arthralgias, and/or arthritis. However, the possibility of an underlying illness should be excluded, and the most common associations are streptococcal infections, upper respiratory viral infections, sarcoidosis, and inflammatory bowel disease, in addition to drugs (oral contraceptives, sulfonamides, penicillins, bromides, iodides). Less common associations include bacterial gastroenteritis (*Yersinia*, *Salmonella*) and coccidioidomycosis followed by tuberculosis, histoplasmosis, brucellosis, and infections with *Chlamydia pneumoniae* or *Chlamydia trachomatis*, *Mycoplasma pneumoniae*, or hepatitis B virus.

Erythema induratum and nodular vasculitis have overlapping features clinically and histologically, and whether they represent two separate entities or the ends of a single disease spectrum is a point of debate; in general, the latter is usually idiopathic and the former is associated with the presence of *Mycobacterium tuberculosis* DNA by PCR within skin lesions. The lesions of lupus panniculitis are found primarily on the cheeks, upper arms, and buttocks (sites of abundant fat) and are seen in both the cutaneous and systemic forms of lupus. The overlying skin may be normal, erythematous, or have the changes of discoid lupus. The subcutaneous fat necrosis that is associated with pancreatic disease is presumably secondary to circulating lipases and is seen in patients with pancreatic carcinoma as well as in patients with acute and chronic pancreatitis. In this disorder, there may be an associated arthritis, fever, and inflammation of visceral fat. Histologic examination of deep incisional biopsy specimens will aid in the diagnosis of the particular type of panniculitis.

Subcutaneous erythematous nodules are also seen in cutaneous polyarteritis nodosa and as a manifestation of *systemic vasculitis* when there is involvement of medium-sized vessels, e.g., systemic polyarteritis nodosa, allergic granulomatosis, or granulomatosis with polyangiitis (Wegener’s) (**Chap. 385**). Cutaneous polyarteritis nodosa presents with painful subcutaneous nodules and ulcers within a red-purple, netlike pattern of livedo reticularis. The latter is due to slowed blood flow through the superficial horizontal venous plexus. The majority of lesions are found on the lower extremities, and while arthralgias and myalgias may accompany cutaneous polyarteritis nodosa, there is no evidence of systemic involvement. In both the cutaneous and systemic

forms of vasculitis, skin biopsy specimens of the associated nodules will show the changes characteristic of a necrotizing vasculitis and/or granulomatous inflammation.

RED-BROWN LESIONS

The cutaneous lesions in *sarcoidosis* (Chap. 390) are classically red to red-brown in color, and with diascopy (pressure with a glass slide), a yellow-brown residual color is observed that is secondary to the granulomatous infiltrate. The waxy papules and plaques may be found anywhere on the skin, but the face is the most common location. Usually there are no surface changes, but occasionally the lesions will have scale. Biopsy specimens of the papules show “naked” granulomas in the dermis, i.e., granulomas surrounded by a minimal number of lymphocytes. Other cutaneous findings in sarcoidosis include annular lesions with an atrophic or scaly center, papules within scars, hypopigmented papules and patches, alopecia, acquired ichthyosis, erythema nodosum, and lupus pernio (see below).

The differential diagnosis of sarcoidosis includes foreign-body granulomas produced by chemicals such as beryllium and zirconium, late secondary syphilis, and *lupus vulgaris*. *Lupus vulgaris* is a form of cutaneous tuberculosis that is seen in previously infected and sensitized individuals. There is often underlying active tuberculosis elsewhere, usually in the lungs or lymph nodes. Lesions occur primarily in the head and neck region and are red-brown plaques with a yellow-brown color on diascopy. Secondary scarring and squamous cell carcinomas can develop within the plaques. Cultures or PCR analysis of the lesions should be performed, along with an interferon γ release assay of peripheral blood, because it is rare for the acid-fast stain to show bacilli within the dermal granulomas.

A generalized distribution of red-brown macules and papules is seen in the form of mastocytosis known as *urticaria pigmentosa* (Chap. 376). Each lesion represents a collection of mast cells in the dermis, with hyperpigmentation of the overlying epidermis. Stimuli such as rubbing cause these mast cells to degranulate, and this leads to the formation of localized urticaria (Darier’s sign). Additional symptoms can result from mast cell degranulation and include headache, flushing, diarrhea, and pruritus. Mast cells also infiltrate various organs such as the liver, spleen, and gastrointestinal tract, and accumulations of mast cells in the bones may produce either osteosclerotic or osteolytic lesions on radiographs. In the majority of these patients, however, the internal involvement remains indolent. A subtype of chronic cutaneous small-vessel vasculitis, *erythema elevatum diutinum* (EED), also presents with papules that are red-brown in color. The papules coalesce into plaques on the extensor surfaces of knees, elbows, and the small joints of the hand. Flares of EED have been associated with streptococcal infections.

BLUE LESIONS

Lesions that are blue in color are the result of vascular ectasias, hyperplasias and tumors or melanin pigment within the dermis. *Venous lakes* (ectasias) are compressible dark-blue lesions that are found commonly in the head and neck region. *Venous malformations* are also compressible blue papulonodules and plaques that can occur anywhere on the body, including the oral mucosa. When there are multiple rather than single congenital lesions, the patient may have the blue rubber bleb syndrome or Maffucci’s syndrome. Patients with the blue rubber bleb syndrome also have vascular anomalies of the gastrointestinal tract that may bleed, whereas patients with Maffucci’s syndrome have associated osteochondromas. *Blue nevi* (moles) are seen when there are collections of pigment-producing nevus cells in the dermis. These benign papular lesions are dome-shaped and occur most commonly on the dorsum of the hand or foot or in the head and neck region.

VIOLACEOUS LESIONS

Violaceous papules and plaques are seen in *lupus pernio*, *lymphoma cutis*, and *cutaneous lupus*. *Lupus pernio* is a particular type of sarcoidosis that involves the tip and alar rim of the nose as well as the earlobes,

with lesions that are violaceous in color rather than red-brown. This form of sarcoidosis is associated with involvement of the upper respiratory tract. The plaques of lymphoma cutis and cutaneous lupus may be red or violaceous in color and were discussed above.

PURPLE LESIONS

Purple-colored papules and plaques are seen in vascular tumors, such as *Kaposi’s sarcoma* (Chap. 226) and *angiosarcoma*, and when there is extravasation of red blood cells into the skin in association with inflammation, as in *palpable purpura* (see “Purpura,” below). Patients with congenital or acquired AV fistulas and venous hypertension can develop purple papules on the lower extremities that can resemble Kaposi’s sarcoma clinically and histologically; this condition is referred to as pseudo-Kaposi’s sarcoma (acral angiodermatitis). Angiosarcoma is found most commonly on the scalp and face of elderly patients or within areas of chronic lymphedema and presents as purple papules and plaques. In the head and neck region, the tumor often extends beyond the clinically defined borders and may be accompanied by facial edema.

BROWN AND BLACK LESIONS

Brown- and black-colored papules are reviewed in “Hyperpigmentation,” above.

CUTANEOUS METASTASES

These are discussed last because they can have a wide range of colors. Most commonly, they present as either firm, skin-colored subcutaneous nodules or firm, red to red-brown papulonodules. The lesions of lymphoma cutis range from pink-red to plum in color, whereas metastatic melanoma can be pink, blue, or black in color. Cutaneous metastases develop from hematogenous or lymphatic spread and are most often due to the following primary carcinomas: in men, melanoma, oropharynx, lung, and colon; and in women, breast, melanoma, and ovary. These metastatic lesions may be the initial presentation of the carcinoma, especially when the primary site is the lung.

PURPURA

(Table 72-16) *Purpura* are seen when there is an extravasation of red blood cells into the dermis and, as a result, the lesions do not blanch with pressure. This is in contrast to those erythematous or violet-colored lesions that are due to localized vasodilatation—they do blanch with pressure. Purpura (≥ 3 mm) and petechiae (≤ 2 mm) are divided into two major groups: palpable and nonpalpable. The most frequent causes of *nonpalpable* petechiae and purpura are primary cutaneous disorders such as *trauma*, *solar (actinic) purpura*, and *capillaritis*. Less common causes are *steroid purpura* and *livedoid vasculopathy* (see “Ulcers,” below). Solar purpura are seen primarily on the extensor forearms, whereas steroid purpura secondary to potent topical glucocorticoids or endogenous or exogenous Cushing’s syndrome can be more widespread. In both cases, there is alteration of the supporting connective tissue that surrounds the dermal blood vessels. In contrast, the petechiae that result from capillaritis are found primarily on the lower extremities. In capillaritis, there is an extravasation of erythrocytes as a result of perivascular lymphocytic inflammation. The petechiae are bright red, 1–2 mm in size, and scattered within yellow-brown patches. The yellow-brown color is caused by hemosiderin deposits within the dermis.

Systemic causes of nonpalpable purpura fall into several categories, and those secondary to clotting disturbances and vascular fragility will be discussed first. The former group includes *thrombocytopenia* (Chap. 140), *abnormal platelet function* as is seen in uremia, and *clotting factor defects*. The initial site of presentation for thrombocytopenia-induced petechiae is the distal lower extremity. Capillary fragility leads to nonpalpable purpura in patients with systemic *amyloidosis* (see “Papulonodular Skin Lesions,” above), disorders of collagen production such as *Ehlers-Danlos syndrome*, and *scurvy*. In scurvy, there are flattened corkscrew hairs with surrounding hemorrhage on the lower extremities, in addition to gingivitis. Vitamin C is a cofactor for lysyl hydroxylase, an enzyme involved in the posttranslational modification of procollagen that is necessary for cross-link formation.

TABLE 72-16 CAUSES OF PURPURA

- I. Primary cutaneous disorders
 - A. Nonpalpable
 1. Trauma
 2. Solar (actinic, senile) purpura
 3. Steroid purpura
 4. Capillaritis
 5. Livedoid vasculopathy in the setting of venous hypertension^a
- II. Systemic diseases
 - A. Nonpalpable
 1. Clotting disturbances
 - a. Thrombocytopenia (including ITP)
 - b. Abnormal platelet function
 - c. Clotting factor defects
 2. Vascular fragility
 - a. Amyloidosis (within normal-appearing skin)
 - b. Ehlers-Danlos syndrome
 - c. Scurvy
 3. Thrombi
 - a. Disseminated intravascular coagulation
 - b. Warfarin (Coumadin)-induced necrosis
 - c. Heparin-induced thrombocytopenia and thrombosis
 - d. Antiphospholipid antibody syndrome
 - e. Monoclonal cryoglobulinemia
 - f. Vasculopathy induced by levamisole-adulterated cocaine
 - g. Thrombotic thrombocytopenic purpura
 - h. Thrombocytosis
 - i. Homozygous protein C or protein S deficiency
 4. Emboli
 - a. Cholesterol
 - b. Fat
 5. Possible immune complex
 - a. Gardner-Diamond syndrome (autoerythrocyte sensitivity)
 - b. Waldenström's hypergammaglobulinemic purpura
 - B. Palpable
 1. Vasculitis
 - a. Cutaneous small-vessel vasculitis, including in the setting of systemic vasculitides
 - b. Polyarteritis nodosa
 2. Emboli^b
 - a. Acute meningococemia
 - b. Disseminated gonococcal infection
 - c. Rocky Mountain spotted fever
 - d. Ecthyma gangrenosum

^aAlso associated with underlying disorders that lead to hypercoagulability, e.g., factor V Leiden, protein C dysfunction/deficiency. ^bBacterial (including rickettsial), fungal, or parasitic.

Abbreviation: ITP, idiopathic thrombocytopenic purpura.

In contrast to the previous group of disorders, the purpura (noninflammatory with a retiform outline) seen in the following group of diseases are associated with thrombi formation within vessels. It is important to note that these thrombi are demonstrable in skin biopsy specimens. This group of disorders includes disseminated intravascular coagulation (DIC), monoclonal cryoglobulinemia, thrombocytosis, thrombotic thrombocytopenic purpura, antiphospholipid antibody syndrome, and reactions to warfarin and heparin (heparin-induced thrombocytopenia and thrombosis). DIC is triggered by several types of infection (gram-negative, gram-positive, viral, and rickettsial) as well as by tissue injury and neoplasms. Widespread purpura and hemorrhagic infarcts of the distal extremities are seen. Similar lesions are found in purpura fulminans, which is a form of DIC associated with fever and hypotension that

occurs more commonly in children following an infectious illness such as varicella, scarlet fever, or an upper respiratory tract infection. In both disorders, hemorrhagic bullae can develop in involved skin.

Monoclonal cryoglobulinemia is associated with plasma cell dyscrasias, chronic lymphocytic leukemia, and lymphoma. Purpura, primarily of the lower extremities, and hemorrhagic infarcts of the fingers, toes, and ears are seen in these patients. Exacerbations of disease activity can follow cold exposure or an increase in serum viscosity. Biopsy specimens show precipitates of the cryoglobulin within dermal vessels. Similar deposits have been found in the lung, brain, and renal glomeruli. Patients with *thrombotic thrombocytopenic purpura* can also have hemorrhagic infarcts as a result of intravascular thromboses. Additional signs include microangiopathic hemolytic anemia and fluctuating neurologic abnormalities, especially headaches and confusion.

Administration of *warfarin* can result in painful areas of erythema that become purpuric and then necrotic with an adherent black eschar; the condition is referred to as warfarin-induced necrosis. This reaction is seen more often in women and in areas with abundant subcutaneous fat—breasts, abdomen, buttocks, thighs, and calves. The erythema and purpura develop between the third and tenth day of therapy, most likely as a result of a transient imbalance in the levels of anticoagulant and procoagulant vitamin K-dependent factors. Continued therapy does not exacerbate preexisting lesions, and patients with an inherited or acquired deficiency of protein C are at increased risk for this particular reaction as well as for purpura fulminans and calciphylaxis.

Purpura secondary to *cholesterol emboli* are usually seen on the lower extremities of patients with atherosclerotic vascular disease. They often follow anticoagulant therapy or an invasive vascular procedure such as an arteriogram but also occur spontaneously from disintegration of atheromatous plaques. Associated findings include livedo reticularis, gangrene, cyanosis, and ischemic ulcerations. Multiple step sections of the biopsy specimen may be necessary to demonstrate the cholesterol clefts within the vessels. Petechiae are also an important sign of *fat embolism* and occur primarily on the upper body 2–3 days after a major injury. By using special fixatives, the emboli can be demonstrated in biopsy specimens of the petechiae. Emboli of tumor or thrombus are seen in patients with atrial myxomas and marantic endocarditis.

In the *Gardner-Diamond syndrome* (autoerythrocyte sensitivity), female patients develop large ecchymoses within areas of painful, warm erythema. Intradermal injections of autologous erythrocytes or phosphatidyl serine derived from the red cell membrane can reproduce the lesions in some patients; however, there are instances where a reaction is seen at an injection site of the forearm but not in the midback region. The latter has led some observers to view Gardner-Diamond syndrome as a cutaneous manifestation of severe emotional stress. More recently, the possibility of platelet dysfunction (as assessed via aggregation studies) has been raised. *Waldenström's hypergammaglobulinemic purpura* is a chronic disorder characterized by petechiae on the lower extremities. There are circulating complexes of IgG–anti-IgG molecules, and exacerbations are associated with prolonged standing or walking.

Palpable purpura are further subdivided into vasculitic and embolic. In the group of vasculitic disorders, cutaneous small-vessel vasculitis, also known as *leukocytoclastic vasculitis* (LCV), is the one most commonly associated with palpable purpura (**Chap. 385**). Underlying etiologies include drugs (e.g., antibiotics), infections (e.g., hepatitis C virus), and autoimmune connective tissue diseases (e.g., rheumatoid arthritis, Sjögren's syndrome, lupus). *Henoch-Schönlein purpura* (HSP) is a subtype of acute LCV that is seen more commonly in children and adolescents following an upper respiratory infection. The majority of lesions are found on the lower extremities and buttocks. Systemic manifestations include fever, arthralgias (primarily of the knees and ankles), abdominal pain, gastrointestinal bleeding, and nephritis. Direct immunofluorescence examination shows deposits of IgA within dermal blood vessel walls. Renal disease is of particular concern in adults with HSP. In *polyarteritis nodosa*, specific cutaneous lesions result from a vasculitis of arterial vessels (arteritis), or there

may be an associated LCV. Arteritis leads to an infarct of the skin, and this explains the irregular outline of the purpura (see below).

Several types of infectious emboli can give rise to palpable purpura. These embolic lesions are usually *irregular* in outline as opposed to the lesions of LCV, which are *circular* in outline. The irregular outline is indicative of a cutaneous infarct, and the size corresponds to the area of skin that received its blood supply from that particular arteriole or artery. The palpable purpura in LCV are circular because the erythrocytes simply diffuse out evenly from the postcapillary venules as a result of inflammation. Infectious emboli are most commonly due to gram-negative cocci (meningococcus, gonococcus), gram-negative rods (Enterobacteriaceae), and gram-positive cocci (*Staphylococcus*). Additional causes include *Rickettsia* and, in immunocompromised patients, *Aspergillus* and other opportunistic fungi.

The embolic lesions in *acute meningococemia* are found primarily on the trunk, lower extremities, and sites of pressure, and a gunmetal-gray color often develops within them. Their size varies from a few millimeters to several centimeters, and the organisms can be cultured from the lesions. Associated findings include a preceding upper respiratory tract infection; fever; meningitis; DIC; and, in some patients, a deficiency of the terminal components of complement. In *disseminated gonococcal infection* (arthritis-dermatitis syndrome), a small number of inflammatory papules and vesicopustules, often with central purpura or hemorrhagic necrosis, are found on the distal extremities. Additional symptoms include arthralgias, tenosynovitis, and fever. To establish the diagnosis, a Gram stain of these lesions should be performed. *Rocky Mountain spotted fever* is a tick-borne disease that is caused by *Rickettsia rickettsii*. A several-day history of fever, chills, severe headache, and photophobia precedes the onset of the cutaneous eruption. The initial lesions are erythematous macules and papules on the wrists, ankles, palms, and soles. With time, the lesions spread centripetally and become purpuric.

Lesions of *ecthyma gangrenosum* begin as edematous, erythematous papules or plaques and then develop central purpura and necrosis. Bullae formation also occurs in these lesions, and they are frequently found in the girdle region. The organism that is classically associated with *ecthyma gangrenosum* is *Pseudomonas aeruginosa*, but other gram-negative rods such as *Klebsiella*, *Escherichia coli*, and *Serratia* can produce similar lesions. In immunocompromised hosts, the list of potential pathogens is expanded to include *Candida* and other opportunistic fungi (e.g., *Aspergillus*, *Fusarium*).

ULCERS

The approach to the patient with a cutaneous ulcer is outlined in [Table 72-17](#). Peripheral vascular diseases of the extremities are reviewed in [Chap. 302](#), as is Raynaud's phenomenon.

Livedoid vasculopathy (livedoid vasculitis; atrophie blanche) represents a combination of a vasculopathy plus intravascular thrombosis. Purpuric lesions and livedo reticularis are found in association with *painful* ulcerations of the lower extremities. These ulcers are often slow to heal, but when they do, irregularly shaped white scars form. The majority of cases are secondary to venous hypertension, but possible underlying illnesses include cryofibrinogenemia and disorders of hypercoagulability, e.g., the antiphospholipid syndrome ([Chaps. 142 and 379](#)).

In *pyoderma gangrenosum*, the border of untreated active ulcers has a characteristic appearance consisting of an undermined necrotic violaceous edge and a peripheral erythematous halo. The ulcers often begin as pustules that then expand rather rapidly to a size as large as 20 cm. Although these lesions are most commonly found on the lower extremities, they can arise anywhere on the surface of the body, including sites of trauma (pathergy). An estimated 30–50% of cases are idiopathic, and the most common associated disorders are ulcerative colitis and Crohn's disease. Less commonly, *pyoderma gangrenosum* is associated with seropositive rheumatoid arthritis, acute and chronic myelogenous leukemia, hairy cell leukemia, myelofibrosis, or a monoclonal gammopathy, usually IgA. Because the histology of *pyoderma gangrenosum* may be nonspecific

TABLE 72-17 CAUSES OF MUCOCUTANEOUS ULCERS

- I. Primary cutaneous disorders
 - A. Peripheral vascular disease ([Chap. 302](#))
 1. Venous
 2. Arterial^a
 - B. Livedoid vasculopathy in the setting of venous hypertension^b
 - C. Squamous cell carcinoma, e.g., within scars, basal cell carcinomas
 - D. Infections, e.g., ecthyma caused by *Streptococcus* ([Chap. 173](#))
 - E. Physical, e.g., trauma, pressure
 - F. Drugs, e.g., hydroxyurea
- II. Systemic diseases
 - A. Lower legs
 1. Small-vessel and medium-vessel vasculitis^c
 2. Hemoglobinopathies ([Chap. 127](#))
 3. Cryoglobulinemia,^c cryofibrinogenemia
 4. Cholesterol emboli^c
 5. Necrobiosis lipoidica^d
 6. Antiphospholipid syndrome ([Chap. 141](#))
 7. Neuropathic^e ([Chap. 417](#))
 8. Panniculitis
 9. Kaposi's sarcoma, acral angiodermatitis
 10. Diffuse dermal angiomas
 - B. Hands and feet
 1. Raynaud's phenomenon ([Chap. 302](#))
 2. Buerger disease
 - C. Generalized
 1. Pyoderma gangrenosum, but most commonly legs
 2. Calciphylaxis ([Chap. 424](#))
 3. Infections, e.g., dimorphic fungi, leishmaniasis
 4. Lymphoma
 - D. Face, especially perioral, and anogenital
 1. Chronic herpes simplex^f
- III. Mucosal
 - A. Behçet's syndrome ([Chap. 387](#))
 - B. Erythema multiforme major, Stevens-Johnson syndrome, TEN
 - C. Primary blistering disorders ([Chap. 73](#))
 - D. Lupus erythematosus, lichen planus
 - E. Inflammatory bowel disease
 - F. Acute HIV infection
 - G. Reactive arthritis (formerly known as Reiter's syndrome)

^aUnderlying atherosclerosis. ^bAlso associated with underlying disorders that lead to hypercoagulability, e.g., factor V Leiden, protein C dysfunction/deficiency, antiphospholipid antibodies. ^cReviewed in section on Purpura. ^dReviewed in section on Papulonodular Skin Lesions. ^eFavors plantar surface of the foot. ^fSign of immunosuppression.

Abbreviation: TEN, toxic epidermal necrolysis.

(dermal infiltrate of neutrophils when in untreated state), the diagnosis requires clinicopathologic correlation, in particular, the exclusion of similar-appearing ulcers such as necrotizing vasculitis, Meleney's ulcer (synergistic infection at a site of trauma or surgery), dimorphic fungi, cutaneous amebiasis, spider bites, and factitial. In the myeloproliferative disorders, the ulcers may be more superficial with a pustulobullous border, and these lesions provide a connection between classic *pyoderma gangrenosum* and acute febrile neutrophilic dermatosis (Sweet syndrome).

FEVER AND RASH

The major considerations in a patient with a fever and a rash are inflammatory diseases versus infectious diseases. In the hospital setting, the most common scenario is a patient who has a drug rash plus a fever secondary to an underlying infection. However, it should be

emphasized that a drug reaction can lead to both a cutaneous eruption and a fever (“drug fever”), especially in the setting of DRESS, AGEP, or serum sickness–like reaction. Additional inflammatory diseases that are often associated with a fever include pustular psoriasis, erythroderma, and Sweet syndrome. Lyme disease, secondary syphilis, and viral and bacterial exanthems (see “Exanthems,” above) are examples of infectious diseases that produce a rash and a fever. Lastly, it is important to determine whether or not the cutaneous lesions represent septic emboli (see “Purpura,” above). Such lesions usually have evidence of ischemia in the form of purpura, necrosis, or impending necrosis (gunmetal-gray color). In the patient with thrombocytopenia, however, purpura can be seen in inflammatory reactions such as morbilliform drug eruptions and infectious lesions.

73 Immunologically Mediated Skin Diseases

Kim B. Yancey, Thomas J. Lawley

A number of immunologically mediated skin diseases and immunologically mediated systemic disorders with cutaneous manifestations are now recognized as distinct entities with consistent clinical, histologic, and immunopathologic findings. Clinically, these disorders are characterized by morbidity (pain, pruritus, disfigurement) and, in some instances, result in death (largely due to loss of epidermal barrier function and/or secondary infection). The major features of the more common immunologically mediated skin diseases are summarized in this chapter (Table 73-1), as are the autoimmune systemic disorders with cutaneous manifestations.

AUTOIMMUNE CUTANEOUS DISEASES

PEMPHIGUS VULGARIS

Pemphigus refers to a group of autoantibody-mediated intraepidermal blistering diseases characterized by loss of cohesion between epidermal cells (a process termed *acantholysis*). Manual pressure to the skin of these patients may elicit the separation of the epidermis (*Nikolsky's sign*). This finding, while characteristic of pemphigus, is not specific to this group of disorders and is also seen in toxic epidermal necrolysis, Stevens-Johnson syndrome, and a few other skin diseases.

Pemphigus vulgaris (PV) is a mucocutaneous blistering disease that predominantly occurs in patients >40 years of age. PV typically begins on mucosal surfaces and often progresses to involve the skin. This disease is characterized by fragile, flaccid blisters that rupture to produce extensive denudation of mucous membranes and skin (Fig. 73-1). The mouth, scalp, face, neck, axilla, groin, and trunk are typically involved. PV may be associated with severe skin pain; some patients experience pruritus as well. Lesions usually heal without scarring except at sites complicated by secondary infection or mechanically induced dermal wounds. Postinflammatory hyperpigmentation is usually present for some time at sites of healed lesions.

Biopsies of early lesions demonstrate intraepidermal vesicle formation secondary to loss of cohesion between epidermal cells (i.e., acantholytic blisters). Blister cavities contain acantholytic epidermal cells, which appear as round homogeneous cells containing hyperchromatic nuclei. Basal keratinocytes remain attached to the epidermal basement membrane; hence, blister formation takes place within the suprabasal portion of the epidermis. Lesional skin may contain focal collections of intraepidermal eosinophils within blister cavities; dermal alterations are slight, often limited to an eosinophil-predominant leukocytic infiltrate. Direct immunofluorescence microscopy of lesional or intact patient skin shows deposits of IgG on the surface of keratinocytes; deposits of complement components are typically found in lesional but not in uninvolved skin. Deposits of IgG on keratinocytes are derived from circulating autoantibodies to cell-surface autoantigens. Such circulating autoantibodies

TABLE 73-1 IMMUNOLOGICALLY MEDIATED BLISTERING DISEASES

Disease	Clinical Manifestations	Histology	Immunopathology	Autoantigens ^a
Pemphigus vulgaris	Flaccid blisters, denuded skin, oromucosal lesions	Acantholytic blister formed in suprabasal layer of epidermis	Cell surface deposits of IgG on keratinocytes	Dsg3 (plus Dsg1 in patients with skin involvement)
Pemphigus foliaceus	Crusts and shallow erosions on scalp, central face, upper chest, and back	Acantholytic blister formed in superficial layer of epidermis	Cell surface deposits of IgG on keratinocytes	Dsg1
Paraneoplastic pemphigus	Painful stomatitis with papulosquamous or lichenoid eruptions that may progress to blisters	Acantholysis, keratinocyte necrosis, and vacuolar interface dermatitis	Cell surface deposits of IgG and C3 on keratinocytes and (variably) similar immunoreactants in epidermal BMZ	Plakin protein family members and desmosomal cadherins (see text for details)
Bullous pemphigoid	Large tense blisters on flexor surfaces and trunk	Subepidermal blister with eosinophil-rich infiltrate	Linear band of IgG and/or C3 in epidermal BMZ	BPAG1, BPAG2
Pemphigoid gestationis	Pruritic, urticarial plaques rimmed by vesicles and bullae on the trunk and extremities	Teardrop-shaped, subepidermal blisters in dermal papillae; eosinophil-rich infiltrate	Linear band of C3 in epidermal BMZ	BPAG2 (plus BPAG1 in some patients)
Dermatitis herpetiformis	Extremely pruritic small papules and vesicles on elbows, knees, buttocks, and posterior neck	Subepidermal blister with neutrophils in dermal papillae	Granular deposits of IgA in dermal papillae	Epidermal transglutaminase
Linear IgA disease	Pruritic small papules on extensor surfaces; occasionally larger, arciform blisters	Subepidermal blister with neutrophil-rich infiltrate	Linear band of IgA in epidermal BMZ	BPAG2 (see text for specific details)
Epidermolysis bullosa acquisita	Blisters, erosions, scars, and milia on sites exposed to trauma; widespread, inflammatory, tense blisters may be seen initially	Subepidermal blister that may or may not include a leukocytic infiltrate	Linear band of IgG and/or C3 in epidermal BMZ	Type VII collagen
Mucous membrane pemphigoid	Erosive and/or blistering lesions of mucous membranes and possibly the skin; scarring of some sites	Subepidermal blister that may or may not include a leukocytic infiltrate	Linear band of IgG, IgA, and/or C3 in epidermal BMZ	BPAG2, laminin-332, or others

^aAutoantigens bound by these patients' autoantibodies are defined as follows: Dsg1, desmoglein 1; Dsg3, desmoglein 3; BPAG1, bullous pemphigoid antigen 1; BPAG2, bullous pemphigoid antigen 2.

Abbreviation: BMZ, basement membrane zone.



A



B

FIGURE 73-1 Pemphigus vulgaris. **A.** Flaccid bullae are easily ruptured, resulting in multiple erosions and crusted plaques. **B.** Involvement of the oral mucosa, which is almost invariable, may present with erosions on the gingiva, buccal mucosa, palate, posterior pharynx, or tongue. (B, Courtesy of Robert Swerlick, MD; with permission.)

can be demonstrated in 80–90% of PV patients by indirect immunofluorescence microscopy; monkey esophagus is the optimal substrate for these studies. Patients with PV have IgG autoantibodies to *desmogleins* (Dsgs), transmembrane desmosomal glycoproteins that belong to the cadherin family of calcium-dependent adhesion molecules. Such autoantibodies can be precisely quantitated by enzyme-linked immunosorbent assay (ELISA). Patients with early PV (i.e., mucosal disease) have IgG autoantibodies to Dsg3; patients with advanced PV (i.e., mucocutaneous disease) have IgG autoantibodies to both Dsg3 and Dsg1. Experimental studies have shown that autoantibodies from patients with PV are pathogenic (i.e., responsible for blister formation) and that their titer correlates with disease activity. Recent studies have shown that the anti-Dsg autoantibody profile in these patients' sera as well as the tissue distribution of Dsg3 and Dsg1 determine the site of blister formation in patients with PV. Coexpression of Dsg3 and Dsg1 by epidermal cells protects against pathogenic IgG antibodies to either of these cadherins but not against pathogenic autoantibodies to both.

PV can be life-threatening. Prior to the availability of glucocorticoids, mortality rates ranged from 60% to 90%; the current figure is ~5%. Common causes of morbidity and death are infection and

complications of treatment with glucocorticoids. Bad prognostic factors include advanced age, widespread involvement, and the requirement for high doses of glucocorticoids (with or without other immunosuppressive agents) for control of disease. The course of PV in individual patients is variable and difficult to predict. Some patients experience remission, while others may require long-term treatment or succumb to complications of their disease or its treatment. The mainstay of treatment is systemic glucocorticoids. Patients with moderate to severe PV are usually started on prednisone at 1 mg/kg per day. If new lesions continue to appear after 1–2 weeks of treatment, the dose may need to be increased and/or prednisone may need to be combined with other immunosuppressive agents such as azathioprine (2–2.5 mg/kg per day), mycophenolate mofetil (20–35 mg/kg per day), or cyclophosphamide (1–2 mg/kg per day). Patients with severe, treatment-resistant disease may derive benefit from plasmapheresis (six high-volume exchanges [i.e., 2–3 L per exchange] over ~2 weeks), IV immunoglobulin (IVIg) (2 g/kg over 3–5 days every 6–8 weeks), or rituximab (375 mg/m² per week × 4, or 1000 mg on days 1 and 15). It is important to bring severe or progressive disease under control quickly in order to lessen the severity and/or duration of this disorder. Accordingly, some have suggested that rituximab and daily glucocorticoids should be used early in PV patients to avert the development of treatment-resistant disease.

PEMPHIGUS FOLIACEUS

Pemphigus foliaceus (PF) is distinguished from PV by several features. In PF, acantholytic blisters are located high within the epidermis, usually just beneath the stratum corneum. Hence, PF is a more superficial blistering disease than PV. The distribution of lesions in the two disorders is much the same, except that in PF mucous membranes are almost always spared. Patients with PF rarely have intact blisters but rather exhibit shallow erosions associated with erythema, scale, and crust formation. Mild cases of PF resemble severe seborrheic dermatitis; severe PF may cause extensive exfoliation. Sun exposure (ultraviolet irradiation) may be an aggravating factor.

PF has immunopathologic features in common with PV. Specifically, direct immunofluorescence microscopy of perilesional skin demonstrates IgG on the surface of keratinocytes. Similarly, patients with PF have circulating IgG autoantibodies directed against the surface of keratinocytes. In PF, autoantibodies are directed against Dsg1, a 160-kDa desmosomal cadherin. These autoantibodies can be quantitated by ELISA. As noted for PV, the autoantibody profile in patients with PF (i.e., anti-Dsg1 IgG) and the tissue distribution of this autoantigen (i.e., expression in oral mucosa that is compensated by coexpression of Dsg3) are thought to account for the distribution of lesions in this disease.

Endemic forms of PF are found in south-central rural Brazil, where the disease is known as *fogo selvagem* (FS), as well as in selected sites in Latin America and Tunisia. Endemic PF, like other forms of this disease, is mediated by IgG autoantibodies to Dsg1. Clusters of FS overlap with those of leishmaniasis, a disease transmitted by bites of the sand fly *Lutzomyia longipalis*. Recent studies have shown that sand-fly salivary antigens (specifically, the LJM11 salivary protein) are recognized by IgG autoantibodies from FS patients (as well as by monoclonal antibodies to Dsg1 derived from these patients). Moreover, mice immunized with LJM11 produce antibodies to Dsg1. Thus, these findings suggest that insect bites may deliver salivary antigens that initiate a cross-reactive humoral immune response, which may lead to FS in genetically susceptible individuals.

Although pemphigus has been associated with several autoimmune diseases, its association with thymoma and/or myasthenia gravis is particularly notable. To date, >30 cases of thymoma and/or myasthenia gravis have been reported in association with pemphigus, usually with PF. Patients may also develop pemphigus as a consequence of drug exposure; drug-induced pemphigus usually resembles PF rather than PV. Drugs containing a thiol group in their chemical structure (e.g., penicillamine, captopril, enalapril) are most commonly associated with drug-induced pemphigus. Nonthiol drugs linked to pemphigus include penicillins, cephalosporins, and piroxicam. It has

been suggested that thiol-containing and non-thiol-containing drugs induce pemphigus via biochemical and immunologic mechanisms, respectively—hence, the better prognosis upon drug withdrawal in cases of pemphigus induced by thiol-containing medications. Some cases of drug-induced pemphigus are durable and require treatment with systemic glucocorticoids and/or immunosuppressive agents.

PF is generally a less severe disease than PV and carries a better prognosis. Localized disease can sometimes be treated with topical or intralesional glucocorticoids; more active cases can usually be controlled with systemic glucocorticoids. Patients with severe, treatment-resistant disease may require more aggressive interventions, as described above for patients with PV.

PARANEOPLASTIC PEMPHIGUS

Paraneoplastic pemphigus (PNP) is an autoimmune acantholytic mucocutaneous disease associated with an occult or confirmed neoplasm. Patients with PNP typically have painful mucosal erosive lesions in association with papulosquamous and/or lichenoid eruptions that often progress to blisters. Palm and sole involvement are common in these patients and raise the possibility that prior reports of neoplasia-associated erythema multiforme actually may have represented unrecognized cases of PNP. Biopsies of lesional skin from these patients show varying combinations of acantholysis, keratinocyte necrosis, and vacuolar-interface dermatitis. Direct immunofluorescence microscopy of a patient's skin shows deposits of IgG and complement on the surface of keratinocytes and (variably) similar immunoreactants in the epidermal basement membrane zone. Patients with PNP have IgG autoantibodies to cytoplasmic proteins that are members of the plakin family (e.g., desmoplakins I and II, bullous pemphigoid antigen [BPAG]1, envoplakin, periplakin, and plectin) and to cell-surface proteins that are members of the cadherin family (e.g., Dsg1 and Dsg3). Passive transfer studies have shown that autoantibodies from patients with PNP are pathogenic in animal models.

The predominant neoplasms associated with PNP are non-Hodgkin's lymphoma, chronic lymphocytic leukemia, thymoma, spindle cell tumors, Waldenström's macroglobulinemia, and Castleman's disease; the last-mentioned neoplasm is particularly common among children with PNP. Rare cases of seronegative PNP have been reported in patients with B cell malignancies previously treated with rituximab. In addition to severe skin lesions, many patients with PNP develop life-threatening bronchiolitis obliterans. PNP is generally resistant to conventional therapies (i.e., those used to treat PV); rarely, a patient's disease may ameliorate or even remit following ablation or removal of underlying neoplasms.

BULLOUS PEMPHIGOID

Bullous pemphigoid (BP) is a polymorphic autoimmune subepidermal blistering disease usually seen in the elderly. Initial lesions may consist of urticarial plaques; most patients eventually display tense blisters on either normal-appearing or erythematous skin (Fig. 73-2). The lesions are usually distributed over the lower abdomen, groin, and flexor surface of the extremities; oral mucosal lesions are found in some patients. Pruritus may be nonexistent or severe. As lesions evolve, tense blisters tend to rupture and be replaced by erosions with or without surmounting crust. Nontraumatized blisters heal without scarring. The major histocompatibility complex class II allele HLA-DQB1*0301 is prevalent in patients with BP. Despite isolated reports, several studies have shown that patients with BP do not have a higher incidence of malignancy than appropriately age- and gender-matched controls.

Biopsies of early lesional skin demonstrate subepidermal blisters and histologic features that roughly correlate with the clinical character of the particular lesion under study. Lesions on normal-appearing skin generally contain a sparse perivascular leukocytic infiltrate with some eosinophils; conversely, biopsies of inflammatory lesions typically show an eosinophil-rich infiltrate at sites of vesicle formation and in perivascular areas. In addition to eosinophils, cell-rich lesions also contain mononuclear cells and neutrophils. It is not possible to distinguish BP from other subepidermal blistering diseases by routine histologic studies alone.



FIGURE 73-2 Bullous pemphigoid with tense vesicles and bullae on erythematous, urticarial bases. (Courtesy of the Yale Resident's Slide Collection; with permission.)

Direct immunofluorescence microscopy of normal-appearing perilesional skin from patients with BP shows linear deposits of IgG and/or C3 in the epidermal basement membrane. The sera of ~70% of these patients contain circulating IgG autoantibodies that bind the epidermal basement membrane of normal human skin in indirect immunofluorescence microscopy. IgG from an even higher percentage of patients reacts with the epidermal side of 1 M NaCl split skin (an alternative immunofluorescence microscopy test substrate used to distinguish circulating IgG autoantibodies to the basement membrane in patients with BP from those in patients with similar, yet different, subepidermal blistering diseases; see below). In BP, circulating autoantibodies recognize 230- and 180-kDa hemidesmosome-associated proteins in basal keratinocytes (i.e., BPAG1 and BPAG2, respectively). Autoantibodies to BPAG2 are thought to deposit in situ, activate complement, produce dermal mast-cell degranulation, and generate granulocyte-rich infiltrates that cause tissue damage and blister formation.

BP may persist for months to years, with exacerbations or remissions. Extensive involvement may result in widespread erosions and compromise cutaneous integrity; elderly and/or debilitated patients may die. The mainstay of treatment is systemic glucocorticoids. Local or minimal disease can sometimes be controlled with topical glucocorticoids alone; more extensive lesions generally respond to systemic glucocorticoids either alone or in combination with immunosuppressive agents. Patients usually respond to prednisone (0.75–1 mg/kg per day). In some instances, azathioprine (2–2.5 mg/kg per day), mycophenolate mofetil (20–35 mg/kg per day), or cyclophosphamide (1–2 mg/kg per day) are necessary adjuncts.

PEMPHIGOID GESTATIONIS

Pemphigoid gestationis (PG), also known as *herpes gestationis*, is a rare, nonviral, subepidermal blistering disease of pregnancy and the puerperium. PG may begin during any trimester of pregnancy or present shortly after delivery. Lesions are usually distributed over the abdomen, trunk, and extremities; mucous membrane lesions are rare. Skin lesions in these patients may be quite polymorphic and consist of erythematous urticarial papules and plaques, vesiculopapules, and/or frank bullae. Lesions are almost always extremely pruritic. Severe exacerbations of PG frequently follow delivery, typically within 24–48 h. PG tends to recur in subsequent pregnancies, often beginning earlier during such gestations. Brief flare-ups of disease may occur with resumption of menses and may develop in patients later exposed to oral contraceptives. Occasionally, infants of affected mothers have transient skin lesions.

Biopsies of early lesional skin show teardrop-shaped subepidermal vesicles forming in dermal papillae in association with an eosinophil-rich leukocytic infiltrate. Differentiation of PG from other subepidermal bullous diseases by light microscopy is difficult. However, direct immunofluorescence microscopy of perilesional skin from PG patients reveals the immunopathologic hallmark of this disorder: linear deposits of C3 in the epidermal basement membrane. These deposits develop as a consequence of complement activation produced by low-titer IgG anti-basement membrane autoantibodies directed against BPAG2, the same hemidesmosome-associated protein that is targeted by autoantibodies in patients with BP—a subepidermal bullous disease that resembles PG clinically, histologically, and immunopathologically.

The goals of therapy in patients with PG are to prevent the development of new lesions, relieve intense pruritus, and care for erosions at sites of blister formation. Many patients require treatment with moderate doses of daily glucocorticoids (i.e., 20–40 mg of prednisone) at some point in their course. Mild cases (or brief flare-ups) may be controlled by vigorous use of potent topical glucocorticoids. Infants born of mothers with PG appear to be at increased risk of being born slightly premature or “small for dates.” Current evidence suggests that there is no difference in the incidence of uncomplicated live births between PG patients treated with systemic glucocorticoids and those managed more conservatively. If systemic glucocorticoids are administered, newborns are at risk for development of reversible adrenal insufficiency.

DERMATITIS HERPETIFORMIS

Dermatitis herpetiformis (DH) is an intensely pruritic, papulovesicular skin disease characterized by lesions symmetrically distributed over extensor surfaces (i.e., elbows, knees, buttocks, back, scalp, and posterior neck) (see Fig. 70-8). Primary lesions in this disorder consist of papules, papulovesicles, or urticarial plaques. Because pruritus is prominent, patients may present with excoriations and crusted papules but no observable primary lesions. Patients sometimes report that their pruritus has a distinctive burning or stinging component; the onset of such local symptoms reliably heralds the development of distinct clinical lesions 12–24 h later. Almost all DH patients have associated, usually subclinical, gluten-sensitive enteropathy (Chap. 349), and >90% express the HLA-B8/DRw3 and HLA-DQw2 haplotypes. DH may present at any age, including in childhood; onset in the second to fourth decades is most common. The disease is typically chronic.

Biopsy of early lesional skin reveals neutrophil-rich infiltrates within dermal papillae. Neutrophils, fibrin, edema, and microvesicle formation at these sites are characteristic of early disease. Older lesions may demonstrate nonspecific features of a subepidermal bulla or an excoriating papule. Because the clinical and histologic features of this disease can be variable and resemble those of other subepidermal blistering disorders, the diagnosis is confirmed by direct immunofluorescence microscopy of normal-appearing perilesional skin. Such studies demonstrate granular deposits of IgA (with or without complement components) in the papillary dermis and along the epidermal basement membrane zone. IgA deposits in the skin are unaffected by control of disease with medication; however, these immunoreactants diminish in intensity or disappear in patients maintained for long periods on a strict gluten-free diet (see below). Patients with DH have granular deposits of IgA in their epidermal basement membrane zone and should be distinguished from individuals with linear IgA deposits at this site (see below).

Although most DH patients do not report overt gastrointestinal symptoms or have laboratory evidence of malabsorption, biopsies of the small bowel usually reveal blunting of intestinal villi and a lymphocytic infiltrate in the lamina propria. As is true for patients with celiac disease, this gastrointestinal abnormality can be reversed by a gluten-free diet. Moreover, if maintained, this diet alone may control the skin disease and eventuate in clearance of IgA deposits from these patients' epidermal basement membrane zones. Subsequent gluten exposure in such patients alters the morphology of their small bowel, elicits a flare-up of their skin disease, and is associated with the reappearance of IgA in their epidermal basement membrane zones. As in patients with

celiac disease, dietary gluten sensitivity in patients with DH is associated with IgA endomysial autoantibodies that target tissue transglutaminase. Studies indicate that patients with DH also have high-avidity IgA autoantibodies to epidermal transglutaminase 3 and that the latter is co-localized with granular deposits of IgA in the papillary dermis of DH patients. Patients with DH also have an increased incidence of thyroid abnormalities, achlorhydria, atrophic gastritis, and autoantibodies to gastric parietal cells. These associations likely relate to the high frequency of the HLA-B8/DRw3 haplotype in these patients, because this marker is commonly linked to autoimmune disorders. The mainstay of treatment of DH is dapsone, a sulfone. Patients respond rapidly (24–48 h) to dapsone (50–200 mg/d), but require careful pretreatment evaluation and close follow-up to ensure that complications are avoided or controlled. All patients taking dapsone at >100 mg/d will have some hemolysis and methemoglobinemia, which are expected pharmacologic side effects of this agent. Gluten restriction can control DH and lessen dapsone requirements; this diet must rigidly exclude gluten to be of maximal benefit. Many months of dietary restriction may be necessary before a beneficial result is achieved. Good dietary counseling by a trained dietitian is essential.

LINEAR IgA DISEASE

Linear IgA disease, once considered a variant form of DH, is actually a separate and distinct entity. Clinically, patients with linear IgA disease may resemble individuals with DH, BP, or other subepidermal blistering diseases. Lesions typically consist of papulovesicles, bullae, and/or urticarial plaques that develop predominantly on central or flexural sites. Oral mucosal involvement occurs in some patients. Severe pruritus resembles that seen in patients with DH. Patients with linear IgA disease do not have an increased frequency of the HLA-B8/DRw3 haplotype or an associated enteropathy and therefore are not candidates for treatment with a gluten-free diet.

Histologic alterations in early lesions may be virtually indistinguishable from those in DH. However, direct immunofluorescence microscopy of normal-appearing perilesional skin reveals a linear band of IgA (and often C3) in the epidermal basement membrane zone. Most patients with linear IgA disease have circulating IgA basement membrane autoantibodies directed against neopeptides in the proteolytically processed extracellular domain of BPAG2. These patients generally respond to treatment with dapsone (50–200 mg/d).

EPIDERMOLYSIS BULLOSA ACQUISITA

Epidermolysis bullosa acquisita (EBA) is a rare, noninherited, polymorphic, chronic, subepidermal blistering disease. (The inherited form is discussed in Chap. 427.) Patients with classic or noninflammatory EBA have blisters on noninflamed skin, atrophic scars, milia, nail dystrophy, and oral lesions. Because lesions generally occur at sites exposed to minor trauma, classic EBA is considered a mechanobullous disease. Other patients with EBA have widespread inflammatory scarring and bullous lesions that resemble severe BP. Inflammatory EBA may evolve into the classic, noninflammatory form of this disease. Rarely, patients present with lesions that predominate on mucous membranes. The HLA-DR2 haplotype is found with increased frequency in EBA patients. Studies suggest that EBA is sometimes associated with inflammatory bowel disease (especially Crohn's disease).

The histology of lesional skin varies with the character of the lesion being studied. Noninflammatory bullae are subepidermal, feature a sparse leukocytic infiltrate, and resemble the lesions in patients with porphyria cutanea tarda. Inflammatory lesions consist of neutrophil-rich subepidermal blisters. EBA patients have continuous deposits of IgG (and frequently C3) in a linear pattern within the epidermal basement membrane zone. Ultrastructurally, these immunoreactants are found in the sublamina densa region in association with anchoring fibrils. Approximately 50% of EBA patients have demonstrable circulating IgG basement membrane autoantibodies directed against type VII collagen—the collagen species that makes up anchoring fibrils. Such IgG autoantibodies bind the dermal side of 1 M NaCl split skin (in contrast to IgG autoantibodies in patients with BP). Studies have shown that passive transfer of experimental or clinical IgG against type

VII collagen can produce lesions in mice that clinically, histologically, and immunopathologically resemble those in patients with inflammatory EBA.

Treatment of EBA is generally unsatisfactory. Some patients with inflammatory EBA may respond to systemic glucocorticoids, either alone or in combination with immunosuppressive agents. Other patients (especially those with neutrophil-rich inflammatory lesions) may respond to dapsone. The chronic, noninflammatory form of EBA is largely resistant to treatment, although some patients may respond to cyclosporine, azathioprine, or IVIg.

MUCOUS MEMBRANE PEMPHIGOID

Mucous membrane pemphigoid (MMP) is a rare, acquired, subepithelial immunobullous disease characterized by erosive lesions of mucous membranes and skin that result in scarring of at least some sites of involvement. Common sites include the oral mucosa (especially the gingiva) and conjunctiva; other sites that may be affected include the nasopharyngeal, laryngeal, esophageal, and anogenital mucosa. Skin lesions (present in about one-third of patients) tend to predominate on the scalp, face, and upper trunk and generally consist of a few scattered erosions or tense blisters on an erythematous or urticarial base. MMP is typically a chronic and progressive disorder. Serious complications may arise as a consequence of ocular, laryngeal, esophageal, or anogenital lesions. Erosive conjunctivitis may result in shortened fornices, symblepharon, ankyloblepharon, entropion, corneal opacities, and (in severe cases) blindness. Similarly, erosive lesions of the larynx may cause hoarseness, pain, and tissue loss that, if unrecognized and untreated, may eventuate in complete destruction of the airway. Esophageal lesions may result in stenosis and/or strictures that could place patients at risk for aspiration. Strictures may also complicate anogenital involvement.

Biopsies of lesional tissue generally show subepithelial vesiculobullae and a mononuclear leukocytic infiltrate. Neutrophils and eosinophils may be seen in biopsies of early lesions; older lesions may demonstrate a scant leukocytic infiltrate and fibrosis. Direct immunofluorescence microscopy of perilesional tissue typically reveals deposits of IgG, IgA, and/or C3 in the epidermal basement membrane. Because many patients with MMP exhibit no evidence of circulating basement membrane autoantibodies, testing of perilesional skin is important diagnostically. Although MMP was once thought to be a single nosologic entity, it is now largely regarded as a disease phenotype that may develop as a consequence of an autoimmune reaction to a variety of molecules in the epidermal basement membrane (e.g., BPAG2, laminin-332, type VII collagen, and other antigens yet to be completely defined). Studies suggest that MMP patients with autoantibodies to laminin-332 have an increased relative risk for cancer. Treatment of MMP is largely dependent upon the sites of involvement. Due to potentially severe complications, patients with ocular, laryngeal, esophageal, and/or anogenital involvement require aggressive systemic treatment with dapsone, prednisone, or the latter in combination with another immunosuppressive agent (e.g., azathioprine, mycophenolate mofetil, cyclophosphamide, or rituximab) or IVIg. Less threatening forms of the disease may be managed with topical or intralesional glucocorticoids.

AUTOIMMUNE SYSTEMIC DISEASES WITH PROMINENT CUTANEOUS FEATURES

DERMATOMYOSITIS

The cutaneous manifestations of dermatomyositis ([Chap. 388](#)) are often distinctive but at times may resemble those of systemic lupus erythematosus (SLE) ([Chap. 378](#)), scleroderma ([Chap. 382](#)), or other overlapping connective tissue diseases ([Chap. 382](#)). The extent and severity of cutaneous disease may or may not correlate with the extent and severity of the myositis. The cutaneous manifestations of dermatomyositis are similar, whether the disease appears in children or in the elderly, except that calcification of subcutaneous tissue is a common late sequela in childhood dermatomyositis.

The cutaneous signs of dermatomyositis may precede or follow the development of myositis by weeks to years. Cases lacking muscle



FIGURE 73-3 Dermatomyositis. Periorbital violaceous erythema characterizes the classic heliotrope rash. (Courtesy of James Krell, MD; with permission.)

involvement (i.e., *dermatomyositis sine myositis*) have also been reported. The most common manifestation is a purple-red discoloration of the upper eyelids, sometimes associated with scaling (“heliotrope” erythema; [Fig. 73-3](#)) and periorbital edema. Erythema on the cheeks and nose in a “butterfly” distribution may resemble the malar eruption of SLE. Erythematous or violaceous scaling patches are common on the upper anterior chest, posterior neck, scalp, and the extensor surfaces of the arms, legs, and hands. Erythema and scaling may be particularly prominent over the elbows, knees, and dorsal interphalangeal joints. Approximately one-third of patients have violaceous, flat-topped papules over the dorsal interphalangeal joints that are pathognomonic of dermatomyositis (Gottron’s papules). Thin violaceous papules and plaques on the elbows and knees of patients with dermatomyositis are referred to as *Gottron’s sign* ([Fig. 73-4](#)). These lesions can be contrasted with the erythema and scaling on the dorsum of the fingers that spares the skin over the interphalangeal joints of some SLE patients. Periungual telangiectasia may be prominent in patients with dermatomyositis. Lacy or reticulated erythema may be associated with fine scaling on the extensor and lateral surfaces of the thighs and upper arms. Other patients, particularly those with long-standing disease, develop areas of hypopigmentation, hyperpigmentation, mild atrophy, and telangiectasia known as *poikiloderma*. Poikiloderma is rare in both SLE and scleroderma and thus can serve as a clinical sign that distinguishes dermatomyositis from these two diseases. Cutaneous changes may



FIGURE 73-4 Gottron’s papules. Dermatomyositis often involves the hands as erythematous flat-topped papules over the knuckles. Periungual telangiectases are also evident.

be similar in dermatomyositis and various overlap syndromes where thickening and binding down of the skin of the hands (*sclerodactyly*) as well as Raynaud's phenomenon can be seen. However, the presence of severe muscle disease, Gottron's papules, heliotrope erythema, and poikiloderma serves to distinguish patients with dermatomyositis. Skin biopsy of the erythematous, scaling lesions of dermatomyositis may reveal only mild nonspecific inflammation but sometimes may show changes indistinguishable from those found in SLE, including epidermal atrophy, hydropic degeneration of basal keratinocytes, edema of the upper dermis, and a mild mononuclear cell infiltrate. Direct immunofluorescence microscopy of lesional skin is usually negative, although granular deposits of immunoglobulin(s) and complement in the epidermal basement membrane zone have been described in some patients. Treatment should be directed at the systemic disease. Topical glucocorticoids are sometimes useful; patients should avoid exposure to ultraviolet irradiation and aggressively use photoprotective measures, including broad-spectrum sunscreens.

LUPUS ERYTHEMATOSUS

The cutaneous manifestations of lupus erythematosus (LE) (Chap. 378) can be divided into acute, subacute, and chronic or discoid types. *Acute cutaneous LE* is characterized by erythema of the nose and malar eminences in a "butterfly" distribution (Fig. 73-5A). The erythema is often sudden in onset, accompanied by edema and fine scale, and



FIGURE 73-5 Acute cutaneous lupus erythematosus (LE). **A.** Acute cutaneous LE on the face, showing prominent, scaly, malar erythema. Involvement of other sun-exposed sites is also common. **B.** Acute cutaneous LE on the upper chest, demonstrating brightly erythematous and slightly edematous papules and plaques. (B, Courtesy of Robert Swerlick, MD; with permission.)

correlated with systemic involvement. Patients may have widespread involvement of the face as well as erythema and scaling of the extensor surfaces of the extremities and upper chest (Fig. 73-5B). These acute lesions, while sometimes evanescent, usually last for days and are often associated with exacerbations of systemic disease. Skin biopsy of acute lesions may show only a sparse dermal infiltrate of mononuclear cells and dermal edema. In some instances, cellular infiltrates around blood vessels and hair follicles are notable, as is hydropic degeneration of basal cells of the epidermis. Direct immunofluorescence microscopy of lesional skin frequently reveals deposits of immunoglobulin(s) and complement in the epidermal basement membrane zone. Treatment is aimed at control of systemic disease. Photoprotection is very important in this as well as in other forms of LE.

Subacute cutaneous lupus erythematosus (SCLE) is characterized by a widespread photosensitive, nonscarring eruption. In most patients, renal and central nervous system involvement is mild or absent. SCLE may present as a papulosquamous eruption that resembles psoriasis or as annular polycyclic lesions that resemble those seen in erythema multiforme. In the papulosquamous form, discrete erythematous papules arise on the back, chest, shoulders, extensor surfaces of the arms, and dorsum of the hands; lesions are uncommon on the central face and the flexor surfaces of the arms as well as below the waist. These slightly scaling papules tend to merge into large plaques, some with a reticulate appearance. The annular form involves the same areas and presents with erythematous papules that evolve into oval, circular, or polycyclic lesions. The lesions of SCLE are more widespread but have less tendency for scarring than lesions of discoid LE. Skin biopsy reveals a dense mononuclear cell infiltrate around hair follicles and blood vessels in the superficial dermis, combined with hydropic degeneration of basal cells in the epidermis. Direct immunofluorescence microscopy of lesional skin reveals deposits of immunoglobulin(s) in the epidermal basement membrane zone in about one-half of these cases. A particulate pattern of IgG deposition throughout the epidermis has been associated with SCLE. Most SCLE patients have anti-Ro autoantibodies. Local therapy alone is usually unsuccessful. Most patients require treatment with aminoquinoline antimalarial drugs. Low-dose therapy with oral glucocorticoids is sometimes necessary. Photoprotective measures against both ultraviolet B and ultraviolet A wavelengths are very important.

Discoid lupus erythematosus (DLE, also called *chronic cutaneous LE*) is characterized by discrete lesions, most often found on the face, scalp, and/or external ears. The lesions are erythematous papules or plaques with a thick, adherent scale that occludes hair follicles (follicular plugging). When the scale is removed, its underside shows small excrescences that correlate with the openings of hair follicles (so-called "carpet tacking"), a finding relatively specific for DLE. Long-standing lesions develop central atrophy, scarring, and hypopigmentation but frequently have erythematous, sometimes raised borders (Fig. 73-6). These lesions persist for years and tend to expand slowly. Up to 15% of patients with DLE eventually meet the American College of Rheumatology criteria for SLE. However, typical discoid lesions are frequently seen in patients with SLE. Biopsy of DLE lesions shows hyperkeratosis, follicular plugging, atrophy of the epidermis, hydropic degeneration of basal keratinocytes, and a mononuclear cell infiltrate adjacent to epidermal, adnexal, and microvascular basement membranes. Direct immunofluorescence microscopy demonstrates immunoglobulin(s) and complement deposits at the basement membrane zone in ~90% of cases. Treatment is focused on control of local cutaneous disease and consists mainly of photoprotection and topical or intralesional glucocorticoids. If local therapy is ineffective, use of aminoquinoline antimalarial agents may be indicated.

SCLEDERMA AND MORPHEA

The skin changes of scleroderma (Chap. 382) usually begin on the hands, feet, and face, with episodes of recurrent nonpitting edema. Sclerosis of the skin commences distally on the fingers (*sclerodactyly*) and spreads proximally, usually accompanied by resorption of bone of the fingertips, which may have punched out ulcers, stellate scars, or areas of hemorrhage (Fig. 73-7). The fingers may actually shrink and



FIGURE 73-6 Discoid (chronic cutaneous) lupus erythematosus. Violaceous, hyperpigmented, atrophic plaques, often with evidence of follicular plugging that may result in scarring, are typical.

become sausage-shaped, and, because the fingernails are usually unaffected, they may curve over the end of the fingertips. Periungual telangiectases are usually present, but periungual erythema is rare. In advanced cases, the extremities show contractures and calcinosis cutis. Facial involvement includes a smooth, unwrinkled brow, taut skin over the nose, shrinkage of tissue around the mouth, and perioral radial furrowing (Fig. 73-8). Matlike telangiectases are often present, particularly on the face and hands. Involved skin feels indurated, smooth, and bound to underlying structures; hyper- and hypopigmentation are common as well. *Raynaud's phenomenon* (i.e., cold-induced blanching, cyanosis, and reactive hyperemia) is documented in almost all patients and can precede development of scleroderma by many years. *Linear scleroderma* is a limited form of disease that presents in a linear, bandlike distribution and tends to involve deep as well as superficial layers of skin. The combination of calcinosis cutis, Raynaud's phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia has been termed the *CREST syndrome*. Centromere antibodies have been reported in a very high percentage of patients with the CREST syndrome but in only a small minority of patients with scleroderma. Skin biopsy reveals thickening of the dermis and homogenization of collagen bundles. Direct immunofluorescence microscopy of lesional skin is usually negative.

Morphea is characterized by localized thickening and sclerosis of skin; it dominates on the trunk. This disorder may affect children or adults. Morphea begins as erythematous or flesh-colored plaques that become sclerotic, develop central hypopigmentation, and have an



FIGURE 73-7 Scleroderma showing acral sclerosis and focal digital ulcers.

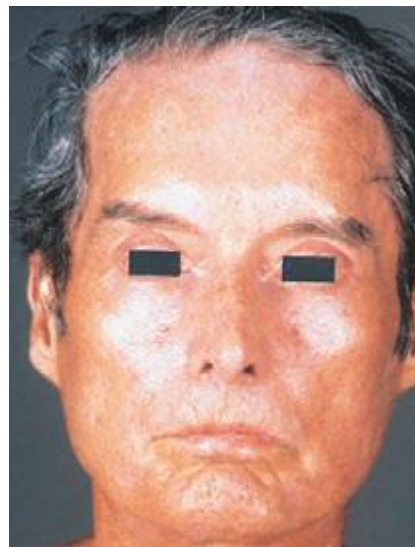


FIGURE 73-8 Scleroderma often eventuates in development of an expressionless, masklike facies.

erythematous border. In most cases, patients have one or a few lesions, and the disease is termed *localized morphea*. In some patients, widespread cutaneous lesions may occur without systemic involvement (*generalized morphea*). Many adults with generalized morphea have concomitant rheumatic or other autoimmune disorders. Skin biopsy of morphea is indistinguishable from that of scleroderma. Scleroderma and morphea are usually quite resistant to therapy. For this reason, physical therapy to prevent joint contractures and to maintain function is employed and is often helpful. Treatment options for early, rapidly progressive disease include phototherapy (UVA1 or PUVA) or methotrexate (15–20 mg/week) alone or in combination with daily glucocorticoids.

Diffuse fasciitis with eosinophilia is a clinical entity that can sometimes be confused with scleroderma. There is usually a sudden onset of swelling, induration, and erythema of the extremities, frequently following significant physical exertion. The proximal portions of the extremities (upper arms, forearms, thighs, calves) are more often involved than are the hands and feet. While the skin is indurated, it usually displays a woody, dimpled, or “pseudocellulite” appearance rather than being bound down as in scleroderma; contractures may occur early secondary to fascial involvement. The latter may also cause muscle groups to be separated and veins to appear depressed (i.e., the “groove sign”). These skin findings are accompanied by peripheral-blood eosinophilia, increased erythrocyte sedimentation rate, and sometimes hypergammaglobulinemia. Deep biopsy of affected areas of skin reveals inflammation and thickening of the deep fascia overlying muscle. An inflammatory infiltrate composed of eosinophils and mononuclear cells is usually found. Patients with eosinophilic fasciitis appear to be at increased risk for developing bone marrow failure or other hematologic abnormalities. While the ultimate course of eosinophilic fasciitis is uncertain, many patients respond favorably to treatment with prednisone in doses of 40–60 mg/d.

The *eosinophilia-myalgia syndrome*, a disorder with epidemic numbers of cases reported in 1989 and linked to ingestion of L-tryptophan manufactured by a single company in Japan, is a multisystem disorder characterized by debilitating myalgias and absolute eosinophilia in association with varying combinations of arthralgias, pulmonary symptoms, and peripheral edema. In a later phase (3–6 months after initial symptoms), these patients often develop localized sclerodermatous skin changes, weight loss, and/or neuropathy (Chap. 382). The precise cause of this syndrome, which may resemble other sclerotic skin conditions, is unknown. However, the implicated lots of L-tryptophan contained the contaminant 1,1-ethylidene bis[tryptophan]. This contaminant may be pathogenic or may be a marker for another substance that provokes the disorder.

74 Cutaneous Drug Reactions

Kanade Shinkai, Robert S. Stern, Bruce U. Wintroub

Cutaneous reactions are among the most frequent adverse reactions to drugs. Most are benign, but a few can be life threatening. Prompt recognition of severe reactions, drug withdrawal, and appropriate therapeutic interventions can minimize toxicity. This chapter focuses on adverse cutaneous reactions to systemic medications; it covers their incidence, patterns, and pathogenesis and provides some practical guidelines on treatment, assessment of causality, and future use of drugs.

USE OF PRESCRIPTION DRUGS IN THE UNITED STATES

In the United States, more than 3 billion prescriptions for over 60,000 drug products, which include more than 2000 different active agents, are dispensed annually. Hospital inpatients alone annually receive about 120 million courses of drug therapy, and half of adult Americans receive prescription drugs on a regular outpatient basis. Many patients use over-the-counter medicines that may cause adverse cutaneous reactions.

INCIDENCE OF CUTANEOUS REACTIONS

Several large cohort studies established that acute cutaneous reaction to drugs affected about 3% of hospital inpatients. Reactions usually occur a few days to 4 weeks after initiation of therapy.

Many drugs of common use are associated with a 1–2% rate of rashes during premarketing clinical trials. The risk is often higher when medications are used in general, unselected populations. The rate may reach 3–7% for amoxicillin, sulfamethoxazole, many anticonvulsants, and anti-HIV agents.

In addition to acute eruptions, a variety of skin diseases can be induced or exacerbated by prolonged use of drugs (e.g., pruritus, pigmentation, nail or hair disorders, psoriasis, bullous pemphigoid, photosensitivity, and even cutaneous neoplasms). These drug reactions are not frequent, but neither their incidence nor their impact on public health has been evaluated.

In a series of 48,005 inpatients over a 20-year period, morbilliform rash (91%) and urticaria (6%) were the most frequent skin reactions. Severe reactions are actually too rare to be detected in such cohorts. Although rare, severe cutaneous reactions to drugs have an important impact on health because of significant sequelae, including mortality. Adverse drug rashes are responsible for hospitalization, increase the duration of hospital stay, and are life threatening. Some populations are at increased risk of drug reactions, including patients with collagen vascular diseases, bone marrow graft recipients, and those with acute Epstein-Barr virus infection. The pathophysiology underlying this association is unknown, but may be related to immunocompromise or immune dysregulation. Risk of drug allergy, including severe hypersensitivity reactions, is increased with HIV infection; individuals with advanced HIV disease (e.g., CD4 T lymphocyte count <200 cells/ μ L) have a forty- to fiftyfold increased risk of adverse reactions to sulfamethoxazole ([Chap. 226](#)).

PATHOGENESIS OF DRUG REACTIONS

Adverse cutaneous responses to drugs can arise as a result of immunologic or nonimmunologic mechanisms.

NONIMMUNOLOGIC DRUG REACTIONS

Examples of responses that arise from nonimmunologic mechanisms are pigmentary changes related to dermal accumulation of medications or their metabolites; alteration of hair follicles by antimetabolites and signaling inhibitors; and lipodystrophy associated with metabolic effects of anti-HIV medications. These side effects are mostly toxic, predictable, and sometimes can be avoided in part by simple preventive measures.

IMMUNOLOGIC DRUG REACTIONS

Evidence suggests an immunologic basis for most acute drug eruptions. Drug reactions may result from immediate release of preformed mediators (e.g., urticaria, anaphylaxis), antibody-mediated reactions, immune complex deposition, and antigen-specific responses. Drug-specific T cell clones can be derived from the blood or from skin lesions of patients with a variety of drug allergies, strongly suggesting that these T cells play a role in drug allergy in an antigen-specific manner. Specific clones were obtained with penicillin G, amoxicillin, cephalosporins, sulfamethoxazole, phenobarbital, carbamazepine, and lamotrigine (medications that are frequently a cause of drug eruptions). Both CD4 and CD8 clones have been obtained; however, their specific roles in the manifestations of allergy have not been elucidated. Drug presentation to T cells was major histocompatibility complex (MHC)-restricted and may involve drug-peptide complex recognition by specific T cell receptors (TCRs).

Once a drug has induced an immune response, the final phenotype of the reaction probably depends on the nature of effectors: cytotoxic (CD8+) T cells in blistering and certain hypersensitivity reactions, chemokines for reactions mediated by neutrophils or eosinophils, and collaboration with B cells for production of specific antibodies for urticarial reaction. Immunologic reactions have recently been classified into further subtypes that provide a useful framework for designating adverse drug reactions based on involvement of specific immune pathways ([Table 74-1](#)).

Immediate Reactions Immediate reactions depend on the release of mediators of inflammation by tissue mast cells or circulating basophils. These mediators include histamine, leukotrienes, prostaglandins, bradykinins, platelet-activating factor, enzymes, and proteoglycans. Drugs can trigger mediator release either directly (“anaphylactoid” reaction) or through IgE-specific antibodies. These reactions usually manifest in the skin and gastrointestinal, respiratory, and cardiovascular systems ([Chap. 376](#)). Primary symptoms and signs include pruritus, urticaria, nausea, vomiting, abdominal cramps, bronchospasm, laryngeal edema, and, occasionally, anaphylactic shock with hypotension and death. They occur within minutes of drug exposure. Nonsteroidal anti-inflammatory drugs (NSAIDs), including aspirin, and radiocontrast media are frequent causes of direct mast cell degranulation or anaphylactoid reactions, which can occur on first exposure. Penicillins and muscle relaxants used in general anesthesia are the most frequent causes of IgE-dependent reactions to drugs, which require prior sensitization. Release of mediators is triggered when polyvalent drug protein conjugates cross-link IgE molecules fixed to sensitized cells. Certain routes of administration favor different clinical patterns (e.g., gastrointestinal effects from oral route, circulatory effects from intravenous route).

Immune Complex-Dependent Reactions Serum sickness is produced by tissue deposition of circulating immune complexes with consumption of complement. It is characterized by fever, arthritis, nephritis, neuritis, edema, and a urticarial, papular, or purpuric rash ([Chap. 385](#)). First described following administration of nonhuman sera, it currently occurs in the setting of monoclonal antibodies and other similar medications. In classic serum sickness, symptoms develop 6 days or more after exposure to a drug, the latent period representing the time needed to synthesize antibody. Cutaneous or systemic vasculitis, a relatively rare complication of drugs, may also be a result of immune complex deposition ([Chap. 385](#)). Cephalosporin and other medications, including monoclonal antibodies such as infliximab, rituximab, and omalizumab, may be associated with clinically similar “serum sickness–like” reactions. The mechanism of this reaction is unknown but is unrelated to complement activation and immune complex formation.

Delayed Hypersensitivity While not completely understood, delayed hypersensitivity directed by drug-specific T cells is an important mechanism underlying the most common drug eruptions, i.e., morbilliform eruptions, and also rare and severe forms such as drug-induced hypersensitivity syndrome (DIHS) (also known

TABLE 74-1 CLASSIFICATION OF ADVERSE DRUG REACTIONS BASED ON IMMUNE PATHWAY


Type	Key Pathway	Key Immune Mediators	Adverse Drug Reaction Type
Type I	IgE	IgE	Urticaria, angioedema, anaphylaxis
Type II	IgG-mediated cytotoxicity	IgG	Drug-induced hemolysis, thrombocytopenia (e.g., penicillin)
Type III	Immune complex	IgG + antigen	Vasculitis, serum sickness, drug-induced lupus
Type IVa	T lymphocyte-mediated macrophage inflammation	IFN- γ , TNF- α T _H 1 cells	Tuberculin skin test, contact dermatitis
Type IVb	T lymphocyte-mediated eosinophil inflammation	IL-4, IL-5, IL-13 T _H 2 cells Eosinophils	Drug-induced hypersensitivity syndrome (DIHS) Morbilliform eruption
Type IVc	T lymphocyte-mediated cytotoxic T lymphocyte inflammation	Cytotoxic T lymphocytes Granzyme Perforin Granulysin (Stevens-Johnson syndrome [SJS]/toxic epidermal necrolysis [TEN] only)	SJS/TEN Morbilliform eruption
Type IVd	T lymphocyte-mediated neutrophil inflammation	CXCL8, IL-17, GM-CSF Neutrophils	Acute generalized exanthematous pustulosis (AGEP)

Abbreviations: GM-CSF, granulocyte-macrophage colony-stimulating factor; IFN, interferon; IL, interleukin; TNF, tumor necrosis factor.

as drug rash with eosinophilia and systemic symptoms [DRESS]), acute generalized exanthematous pustulosis (AGEP), Stevens-Johnson syndrome (SJS), and toxic epidermal necrolysis (TEN) (Table 74-1). Drug-specific T cells have been detected in these types of drug eruptions. For example, drug-specific cytotoxic T cells have been detected in the skin lesions of fixed drug eruptions and of TEN. In TEN, skin lesions contain T lymphocytes reactive to autologous lymphocytes and keratinocytes in a drug-specific, HLA-restricted, and perforin/granzyme-mediated pathway.


The mechanism(s) by which medications result in T cell activation is unknown. Two hypotheses prevail: first, that the antigens driving these reactions may be the native drug itself or components of the drug covalently complexed with endogenous proteins, presented in association with HLA molecules to T cells through the classical antigen presentation pathway, or alternatively, through direct interaction of the drug/metabolite with the T cell receptor or peptide-loaded HLA (e.g., the pharmacologic interaction of drugs with immune receptors, or p-i hypothesis). Recent x-ray crystallography data characterizing binding between specific HLA molecules to particular drugs known to cause hypersensitivity reactions demonstrates unique alterations to the MHC peptide-binding groove, suggesting a molecular basis for T cell activation and the development of hypersensitivity reactions.

GENETIC FACTORS AND CUTANEOUS DRUG REACTIONS

 Genetic determinants may predispose individuals to severe drug reactions by affecting either drug metabolism or immune responses to drugs. Polymorphisms in cytochrome P450 enzymes, drug acetylation, methylation (such as thiopurine methyltransferase activity and azathioprine), and other forms of metabolism (such as glucose-6-phosphate dehydrogenase) may increase susceptibility to drug toxicity or underdosing, highlighting a role for differential pharmacokinetic or pharmacodynamic effects.

Associations between drug hypersensitivities and HLA haplotypes also suggest a key role for immune mechanisms. Hypersensitivity to the anti-HIV medication abacavir is strongly associated with HLA-B*57:01 (Chap. 226). In Taiwan, within a homogeneous Han Chinese population, a 100% association was observed between SJS/TEN (but not DIHS) related to carbamazepine and HLA-B*15:02. In the same population, another 100% association was found between SJS, TEN, or DIHS related to allopurinol and HLA-B*58:01. These associations are drug and phenotype specific; that is, HLA-specific T cell stimulation by medications leads to distinct reactions and may explain why the reaction patterns are so clinically diverse. However, the strong associations found in Taiwan have not been observed in other countries with more heterogeneous populations.

GLOBAL CONSIDERATIONS

 Recognition of the HLA associations with drug hypersensitivity has been acknowledged by recommendations to screen high-risk populations. Genetic screening for HLA-B*57:01 to prevent abacavir hypersensitivity, which carries a 100% negative predictive value when patch test confirmed and 55% positive predictive value generalizable across races, is becoming the clinical standard of care worldwide (number needed to treat = 13). The U.S. Food and Drug Administration recently mandated new labeling of carbamazepine recommending HLA-B*15:02 screening of Asian individuals prior to receiving a new prescription of the medication. The American College of Rheumatology has recommended HLA-B*58:01 screening of Han Chinese patients prescribed allopurinol. To date, screening for a single HLA (but not multiple HLA haplotypes) in specific populations has been determined to be cost-effective.

Several investigators have proposed that specific HLA haplotypes associated with drug hypersensitivity indeed play a pathogenic role; stimulation of carbamazepine-specific cytotoxic T lymphocytes (CTL) in the context of HLA-B*15:02 results in production of a putative mediator of keratinocyte necrosis in TEN. Other studies have identified CTLs reactive to carbamazepine that use highly restricted V-alpha and V-beta TCR repertoires in patients with carbamazepine hypersensitivity that are not found in carbamazepine-tolerant individuals. Although not yet clinically available, some investigators have suggested combined genetic testing for specific HLA haplotypes and functional screening for TCR repertoire to best identify patients at risk.

CLINICAL PRESENTATION OF CUTANEOUS DRUG REACTIONS

NONIMMUNE CUTANEOUS REACTIONS

Exacerbation or Induction of Dermatologic Diseases A variety of drugs can exacerbate preexisting diseases or sometimes induce a disease that may or may not disappear after withdrawal of the inducing medication. For example, NSAIDs, lithium, beta blockers, tumor necrosis factor (TNF) α cytokine antagonists, interferon (IFN) α , and angiotensin-converting enzyme (ACE) inhibitors can exacerbate plaque psoriasis, whereas antimalarials and withdrawal of systemic glucocorticoids can worsen pustular psoriasis. The situation of TNF- α inhibitors is unusual, as this class of medications is used to treat psoriasis; however, in other cases, they may induce psoriasis (especially palmar-plantar) in patients being treated for other conditions. Acne may be induced by glucocorticoids, androgens, lithium, and antidepressants. Follicular papular or pustular eruptions of the face and trunk, sometimes mimicking acne, frequently occur with epidermal growth factor (EGF) receptor antagonists. In the case of EGF-receptor antagonists, the severity of the eruption correlates with a better anticancer effect. It

may be secondarily impetiginized and often spares areas of prior or active radiation. Tetracycline antibiotics, topical corticosteroids, and topical anti-acne treatments (such as benzoyl peroxide and clindamycin lotion) are helpful.

Several medications induce or exacerbate autoimmune disease. Interleukin (IL) 2, IFN- α , and anti-TNF- α are associated with new-onset systemic lupus erythematosus (SLE). Drug-induced lupus is classically marked by antinuclear and antihistone antibodies and, in some cases, anti-double-stranded DNA (D-penicillamine, anti-TNF- α) or p-ANCA (minocycline) antibodies. Minocycline and thiazide diuretics may exacerbate subacute SLE; pemphigus can be induced by D-penicillamine and ACE inhibitors. Furosemide is associated with drug-induced bullous pemphigoid. Vancomycin is associated with linear IgA bullous dermatitis, a transient blistering disorder.

Other medications may cause highly selective cutaneous reactions. Gadolinium contrast has been associated with nephrogenic systemic fibrosis, a condition of sclerosing skin with rare internal organ involvement; advanced renal compromise may be an important risk factor. Granulocyte colony-stimulating factor may induce various neutrophilic dermatoses, including Sweet syndrome and pyoderma gangrenosum. Both systemic and topical glucocorticoids cause a variety of atrophic skin changes, including atrophy and striae, and, in sufficiently high doses, can impede wound healing.

The hypothesis that a drug may be responsible should always be considered, especially in cases with atypical clinical presentation. Resolution of the cutaneous reaction may be delayed upon discontinuation of the medication (e.g., lichenoid drug eruptions may take years to resolve).

Photosensitivity Eruptions Photosensitivity eruptions are usually most marked in sun-exposed areas but may extend to sun-protected areas. The mechanism is almost always phototoxicity. Phototoxic reactions resemble sunburn and can occur with first exposure to a drug. Blistering may occur in drug-related pseudoporphyria, most commonly with NSAIDs (Fig. 74-1). The severity of the reactions depends on the tissue level of the drug, its efficiency as a photosensitizer, and the extent of exposure to the activating wavelengths of ultraviolet (UV) light (Chap. 75).

Common orally administered photosensitizing drugs include fluoroquinolones and tetracycline antibiotics. Other drugs less frequently encountered are chlorpromazine, thiazides, and NSAIDs. Voriconazole may result in severe photosensitivity, accelerated photo-induced aging, and cutaneous carcinogenesis in organ transplant recipients.

Because UV-A and visible light, which trigger these reactions, are not easily absorbed by nonopaque sunscreens and are transmitted through window glass, photosensitivity reactions may be difficult to block. Photosensitivity reactions abate with removal of either the drug or UV radiation, use of sunscreens that block UV-A light, and treating the reaction as one would a sunburn. Rarely, individuals develop

persistent reactivity to light, necessitating long-term avoidance of sun exposure.

Pigmentation Changes Drugs, either systemic or topical, may cause a variety of pigmentary changes in the skin. Oral contraceptives may induce melasma. Long-term minocycline, pefloxacin, and amiodarone may cause blue-gray pigmentation. Phenothiazine, gold, and bismuth result in gray-brown pigmentation of sun-exposed areas. Numerous cancer chemotherapeutic agents may be associated with characteristic patterns of pigmentation (e.g., bleomycin, busulfan, daunorubicin, cyclophosphamide, hydroxyurea, and methotrexate). Clofazimine causes a drug-induced lipofuscinosis with characteristic red-brown coloration. Hyperpigmentation of the face, mucous membranes, and pretibial and subungual areas occurs with antimalarials. Quinacrine causes generalized, cutaneous yellow discoloration. Pigmentation changes may also occur in mucous membranes (busulfan, bismuth), conjunctiva (chlorpromazine, thioridazine, imipramine, clomipramine), nails (zidovudine, doxorubicin, cyclophosphamide, bleomycin, fluorouracil, hydroxyurea), hair, and teeth (tetracyclines).

Warfarin Necrosis of Skin This rare reaction (0.01–0.1%) usually occurs between the third and tenth days of therapy with warfarin, usually in women. Common sites are breasts, thighs, and buttocks (Fig. 74-2). Lesions are sharply demarcated, indurated, and erythematous or purpuric and may progress to form large, hemorrhagic bullae with eventual necrosis and slow-healing eschar formation. These lesions can be life threatening.

Development of the syndrome is unrelated to drug dose, and the course is not altered by discontinuation of the drug after onset of the eruption. Warfarin anticoagulation in heterozygous protein C deficiency causes a precipitous fall in circulating levels of protein C, permitting hypercoagulability and thrombosis in the cutaneous microvasculature, with consequent areas of necrosis. Heparin-induced necrosis may have clinically similar features but is probably due to heparin-induced platelet aggregation with subsequent occlusion of blood vessels; it can affect areas adjacent to the injection site or more distant sites if infused.

Warfarin-induced cutaneous necrosis is treated with vitamin K, heparin, surgical debridement, and intensive wound care. Treatment with protein C concentrates may also be helpful. Newer agents such as dabigatran etexilate may avoid warfarin necrosis in high-risk patients.

Drug-Induced Hair Disorders • DRUG-INDUCED HAIR LOSS Medications may affect hair follicles at two different phases of their growth cycle: anagen (growth) or telogen (resting). *Anagen effluvium* occurs within days of drug administration, especially with antimetabolite or other chemotherapeutic drugs. In contrast, *telogen effluvium*, the delay is 2 to 4 months following initiation of a new medication. Both present as diffuse nonscarring alopecia most often reversible after discontinuation



FIGURE 74-1 Pseudoporphyria due to nonsteroidal anti-inflammatory drugs.



FIGURE 74-2 Warfarin necrosis.

of the responsible agent. The prevalence and severity of alopecia depend on the drug as well as on an individual's predisposition.

A considerable number of drugs have been reported to induce hair loss. These include antineoplastic agents (alkylating agents, bleomycin, vinca alkaloids, platinum compounds), anticonvulsants (carbamazepine, valproate), antihypertensive drugs (beta blockers), antidepressants, antithyroid drugs, IFNs (especially IFN- α), oral contraceptives, and cholesterol-lowering agents.

DRUG-INDUCED HAIR GROWTH Medications may also cause hair growth. Hirsutism is an excessive growth of terminal hair with masculine hair growth pattern in a female, most often on the face and trunk, due to androgenic stimulation of hormone-sensitive hair follicles (anabolic steroids, oral contraceptives, testosterone, corticotropin). Hypertrichosis is a distinct pattern of hair growth, not in a masculine pattern, typically located on the forehead and temporal regions of the face. Drugs responsible for hypertrichosis include anti-inflammatory drugs, glucocorticoids, vasodilators (diazoxide, minoxidil), diuretics (acetazolamide), anticonvulsants (phenytoin), immunosuppressive agents (cyclosporine A), psoralens, and zidovudine.

Changes in hair color or structure are uncommon adverse effects from medications. Hair discoloration may occur with chloroquine, IFN- α , chemotherapeutic agents, and tyrosine kinase inhibitors. Changes in hair structure have been observed in patients given epidermal growth factor receptor (EGFR) inhibitors, tyrosine kinase inhibitors (Fig. 74-3), and acitretin.

Drug-Induced Nail Disorders Drug-related nail disorders usually involve all 20 nails and need months to resolve after withdrawal of the offending agent. The pathogenesis is most often toxic. Drug-induced nail changes include Beau's line (transverse depression of the nail plate), onycholysis (detachment of the distal part of the nail plate), onychomadesis (detachment of the proximal part of the nail plate), pigmentation, and paronychia (inflammation of periungual skin).

ONYCHOLYSIS Onycholysis occurs with tetracyclines, fluoroquinolones, phenothiazines, and psoralens, as well as in persons taking NSAIDs, captopril, retinoids, sodium valproate, and many chemotherapeutic agents such as anthracyclines or taxanes including paclitaxel and docetaxel. The risk of onycholysis in patients receiving cytotoxic drugs, tetracyclines, quinolones, phenothiazines, and psoralens can be increased by exposure to sunlight.

ONYCHOMADESIS Onychomadesis is caused by temporary arrest of nail matrix mitotic activity. Common drugs reported to induce onychomadesis include carbamazepine, lithium, retinoids, and chemotherapeutic agents such as cyclophosphamide and vincristine.

PARONYCHIA Paronychia and multiple pyogenic granuloma (Fig. 74-4) with progressive and painful periungual abscess of fingers and toes



FIGURE 74-3 Dysmorphic eyelashes in association with erlotinib.



FIGURE 74-4 Pyogenic granuloma in association with isotretinoin.

are side effects of systemic retinoids, lamivudine, indinavir, and anti-EGFR monoclonal antibodies (cetuximab, gefitinib).

NAIL DISCOLORATION Some drugs—including anthracyclines, taxanes, fluorouracil, psoralens, and zidovudine—may induce nail bed hyperpigmentation through melanocyte stimulation. It appears to be reversible and dose-dependent.

Toxic Erythema of Chemotherapy and Other Chemotherapy Reactions

Because many agents used in cancer chemotherapy inhibit cell division, rapidly proliferating elements of the skin, including hair, mucous membranes, and appendages, are sensitive to their effects. A broad spectrum of chemotherapy-related skin toxicities have been reported, including neutrophilic eccrine hidradenitis, sterile cellulitis, exfoliative dermatitis, and flexural erythema; although previously designated as distinct skin eruptions, recent nomenclature classifies these under the unifying diagnosis of toxic erythema of chemotherapy (TEC). Acral erythema, marked by dysesthesia and an erythematous, edematous eruption of the palms and soles, is caused by cytarabine, doxorubicin, methotrexate, hydroxyurea, and fluorouracil and may be alleviated by pyridoxine supplementation.

The recent introduction of many new monoclonal antibody and small molecular signaling inhibitors for the treatment of cancer has been accompanied by numerous reports of skin and hair toxicity; only the most common of these are mentioned here. Cetuximab and other EGFR antagonists induce follicular eruptions and nail toxicity after a mean interval of 10 days in a majority of patients. Xerosis, eczematous eruptions, acneiform eruptions, and pruritus are common. Erlotinib is associated with marked hair textural changes (Fig. 74-3). Sorafenib, a tyrosine kinase inhibitor, may result in follicular eruptions and bullous palmoplantar eruptions with dysesthesia (Fig. 74-5). BRAF inhibitors are associated with photosensitivity, dyskeratotic (Grover's-like) rash, hyperkeratotic benign cutaneous neoplasms, and keratoacanthoma-like squamous cell carcinomas. Rash, pruritus, and vitiliginous depigmentation have been reported in association with ipilimumab (anti-CTLA4) treatment.

IMMUNE CUTANEOUS REACTIONS: COMMON

Maculopapular Eruptions Morbilliform or maculopapular eruptions (Fig. 74-6) are the most common of all drug-induced reactions, often start on the trunk or intertriginous areas, and consist of erythematous macules and papules that are symmetric and confluent. Involvement of mucous membranes is unusual; the eruption may be associated with moderate to severe pruritus and fever. Diagnosis is rarely assisted by laboratory testing. Skin biopsy often shows nonspecific inflammatory changes. A viral exanthem is the principal differential diagnostic consideration, especially in children, and graft-versus-host disease is also a consideration in the proper clinical setting. Absence of enanthems;



FIGURE 74-5 Sorafenib-associated hand-foot syndrome.

absence of ear, nose, throat, and upper respiratory tract symptoms; and polymorphism of the skin lesions support a drug rather than a viral eruption. Certain medications carry very high rates of morbilliform eruption, including nevirapine and lamotrigine, even in the absence of hypersensitivity reactions. Lamotrigine morbilliform rash is associated with higher starting doses, rapid dose escalation, concomitant use of valproate (which increases lamotrigine levels and half-life) and use in children, especially with seizure disorder.

Maculopapular reactions usually develop within 1 week of initiation of therapy and last less than 2 weeks. Occasionally, these eruptions resolve despite continued use of the responsible drug. Because the eruption may also worsen, the suspect drug should be discontinued unless it is essential; it is important to note that the rash may continue to progress for a few days to up to one week following medication discontinuation. Oral antihistamines and emollients may help relieve pruritus. Short courses of potent topical glucocorticoids can reduce inflammation and symptoms. Systemic glucocorticoid treatment is rarely indicated.

Pruritus Pruritus is associated with almost all drug eruptions and, in some cases, may represent the only symptom of the adverse cutaneous reaction. It is usually alleviated by antihistamines such as hydroxyzine or diphenhydramine. Pruritus stemming from specific medications may require distinct treatment; opiate-related pruritus may require selective opiate antagonists to gain relief. Pruritus is a common complication of antimalarial therapy, occurring in up to 50% of black patients receiving chloroquine, and may be severe enough to lead to discontinuation of treatment. It is much rarer in Caucasians taking chloroquine. Intense pruritus, sometimes accompanied by an eczematous rash, may occur in 20% of patients receiving IFN and ribavirin for



FIGURE 74-6 Morbilliform drug eruption.

hepatitis C; addition of the protease inhibitor telaprevir may increase this occurrence to 50% of treated patients.

Urticaria/Angioedema/Anaphylaxis Urticaria, the second most frequent type of cutaneous reaction to drugs, is characterized by pruritic, red wheals of varying size rarely lasting more than 24 h. It has been observed in association with nearly all drugs, most frequently ACE inhibitors, aspirin, NSAIDs, penicillin, and blood products. However, medications account for no more than 10–20% of acute urticaria cases. Deep edema within dermal and subcutaneous tissues is known as angioedema and may involve respiratory and gastrointestinal mucous membranes as well. Urticaria and angioedema may be part of a life-threatening anaphylactic reaction.

Drug-induced urticaria may be caused by three mechanisms: an IgE-dependent mechanism, circulating immune complexes (serum sickness), and nonimmunologic activation of effector pathways. IgE-dependent urticarial reactions usually occur within 36 h of drug exposure but can occur within minutes. Immune complex–induced urticaria associated with serum sickness–like reactions usually occurs 6–12 days after first exposure. In this syndrome, the urticarial eruption (typically polycyclic plaques) may be accompanied by fever, hematuria, arthralgias, hepatic dysfunction, and neurologic symptoms. Certain drugs, such as NSAIDs, ACE inhibitors, angiotensin II antagonists, radiographic dye, and opiates, may induce urticarial reactions, angioedema, and anaphylaxis in the absence of drug-specific antibody through direct mast-cell degranulation.

Radiocontrast agents are a common cause of urticaria and, in rare cases, can cause anaphylaxis. High-osmolality radiocontrast media were about five times more likely to induce urticaria (1%) or anaphylaxis than were newer low-osmolality media. About one-third of those with mild reactions to previous exposure react on reexposure. Pretreatment with prednisone and diphenhydramine reduces reaction rates. Persons with a reaction to a high-osmolality contrast media may be given low-osmolality media if later contrast studies are required.

The treatment of urticaria or angioedema depends on the severity of the reaction. In severe cases with respiratory or cardiovascular compromise, epinephrine is the mainstay of therapy, but its effect is reduced in patients using beta blockers. Treatment with intravenous systemic glucocorticoids is helpful. For patients with urticaria without symptoms of angioedema or anaphylaxis, drug withdrawal and oral antihistamines are usually sufficient. Future drug avoidance is recommended; rechallenge, especially in individuals with severe reactions, should only occur in an intensive care setting.

Anaphylactoid Reactions Vancomycin is associated with red man syndrome, a histamine-related anaphylactoid reaction characterized by flushing, diffuse maculopapular eruption, and hypotension. In rare cases, cardiac arrest may be associated with rapid IV infusion of the medication.

Irritant/Allergic Contact Dermatitis Patients using topical medications may develop an irritant or allergic contact dermatitis to the medication itself or to a preservative or other component of the formulation. Reactions to chlorhexidine, neomycin sulfate, and polymyxin B are common. Allergic contact dermatitis to topical glucocorticoids may also occur and is paradoxically partially masked by the anti-inflammatory nature of the medication itself; typically this allergy is selective for one of the four classes of glucocorticoid types, as subdivided by allergenic properties. Patch testing can be useful to determine whether a patient is steroid allergic. Desoximetasone is rarely allergenic.

Fixed Drug Eruptions These less common reactions are characterized by one or more sharply demarcated, dull red to brown lesions, sometimes with central bulla (Fig. 74-7). Hyperpigmentation often results after resolution of the acute inflammation. With rechallenge, the lesion recurs in the same (e.g., fixed) location. Lesions often involve the lips, hands, legs, face, genitalia, and oral mucosa and cause a burning sensation. Most patients have multiple lesions. Fixed drug eruptions have been associated with pseudoephedrine (frequently a nonpigmented reaction), phenolphthalein (in laxatives), sulfonamides, tetracyclines, NSAIDs, and barbiturates.



FIGURE 74-7 Fixed drug eruption.

IMMUNE CUTANEOUS REACTIONS: RARE AND SEVERE

Vasculitis Cutaneous small-vessel vasculitis often presents as palpable purpuric lesions that may be generalized or limited to the lower extremities or other dependent areas (**Chap. 385**). Pustular lesions and hemorrhagic blisters also occur. Vasculitis may involve other organs, including the liver, kidney, brain, and joints. Drugs are implicated as a cause of 10–15% of all cases of small-vessel vasculitides. Infection, malignancy, and collagen vascular disease are responsible for the majority of non-drug-related cases.

Propylthiouracil induces a cutaneous vasculitis that is accompanied by leukopenia and splenomegaly. Direct immunofluorescent changes in these lesions suggest immune-complex deposition. Common drugs implicated in vasculitis include allopurinol, thiazides, sulfonamides, antimicrobials, and NSAIDs. The presence of eosinophils in the perivascular infiltrate of skin biopsy suggests a drug etiology.

Pustular Eruptions AGEP is a rare reaction pattern (3–5 cases per million per year) that is often associated with exposure to drugs (**Fig. 74-8**). Usually beginning on the face or intertriginous areas, small nonfollicular pustules overlying erythematous and edematous skin may coalesce and lead to superficial erosion. Differentiating this eruption from TEN in its initial stages may be difficult. A skin biopsy is important and shows neutrophil collections and sparse necrotic keratinocytes in the upper part of the epidermis instead of the full-thickness epidermal necrosis that characterizes TEN. Fever and leukocytosis are common, and eosinophilia occurs in one-third of cases. Acute pustular psoriasis

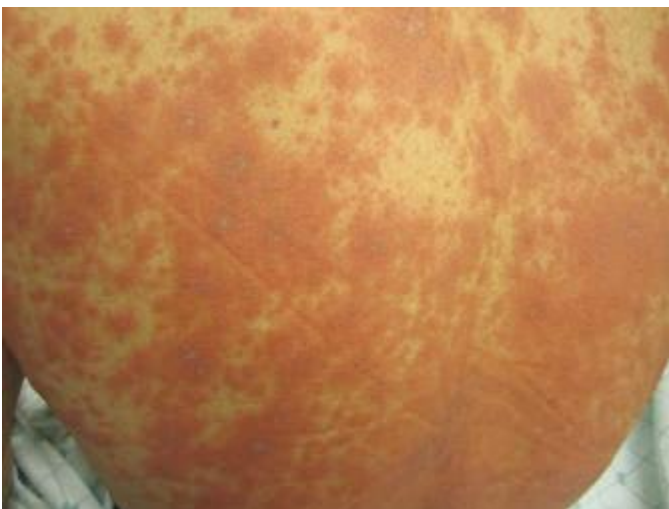


FIGURE 74-8 Acute generalized exanthematous pustulosis.

is the principal differential diagnostic consideration. DIHS with pustular features must also be clinically considered, although the timing for the onset of DIHS is distinct (much later onset). AGEP often begins within a few days of initiating drug treatment, most notably antibiotics, but may occur as late as 7–14 days after initiation of treatment. A broad range of drug classes (anticonvulsants, mercury, radiocontrast dye) and infections (viral, *Mycoplasma*) are also associated with AGEP. Patch testing with the responsible drug results in a localized pustular eruption.

Drug-induced Hypersensitivity Syndrome Drug-induced hypersensitivity syndrome (DIHS) is a multiorgan drug reaction previously known as DRESS (drug reaction with eosinophilia and systemic symptoms); since eosinophilia is not always present, the term *DIHS* is now preferred. Allopurinol is the most common cause. Although less frequently prescribed, abacavir has been reported to cause DIHS with an incidence as high as 4–8%. It presents as a widespread erythematous eruption that may become purpuric, pustular, or lichenoid and is accompanied by many of the following features: fever, facial edema, lymphadenopathy, leukocytosis (often with atypical lymphocytes and eosinophilia), hepatitis, myositis (including myocarditis), and sometimes nephritis (with proteinuria) or pneumonitis. Distinct patterns of timing of onset and organ involvement may exist; e.g., allopurinol classically induces DIHS with renal involvement. Cardiac and lung involvement is more common with minocycline; gastrointestinal involvement is almost exclusively seen with abacavir, and some medications typically lack eosinophilia (abacavir, dapsone, lamotrigine). The cutaneous reaction usually begins 2–8 weeks after the drug is started and lasts longer than mild eruptions after drug cessation. Signs and symptoms may persist for several weeks, especially those associated with hepatitis. The eruption recurs with rechallenge, and cross-reactions among aromatic anticonvulsants, including phenytoin, carbamazepine, and barbiturates, are frequent. Other drugs causing this syndrome include sulfonamides and other antibiotics. Hypersensitivity to reactive drug metabolites, hydroxylamine for sulfamethoxazole, and arene oxide for aromatic anticonvulsants may be involved in the pathogenesis of DIHS. Reactivation of herpes viruses, especially herpesvirus 6 and Epstein-Barr virus (EBV), has been frequently reported in this syndrome, although the causal role of viral infection has been debated. Recent research suggests that inciting drugs may reactivate quiescent herpes viruses, resulting in expansion of viral-specific CD8+ T lymphocytes with subsequent end-organ damage. Viral reactivation may be associated with a worse clinical prognosis. Mortality rates as high as 10% have been reported; mortality is highest in association with hepatitis. Systemic glucocorticoids (prednisone, 1–2 mg/kg daily) should be started with slow taper over 8–12 weeks. A steroid-sparing agent, such as mycophenolate mofetil, may be indicated in cases of rapid recurrence upon steroid taper. In all cases, rapid withdrawal of the suspected drug is required. Given the severe long-term complications of myocarditis, patients should undergo cardiac evaluation if heart involvement is suspected by hypotension or arrhythmia. Patients should be closely monitored for resolution of organ dysfunction and for development of late-onset autoimmune thyroiditis (up to 6 months).

Stevens-Johnson Syndrome and Toxic Epidermal Necrolysis SJS and TEN are characterized by blisters and mucosal/epidermal detachment resulting from full-thickness epidermal necrosis in the absence of substantial dermal inflammation (**Fig. 74-9**). The term *Stevens-Johnson syndrome* describes cases with blisters developing on target lesions, dusky or purpuric macules in which mucosal involvement is significant, and total body surface area blistering and eventual detachment in <10% of cases. The term *Stevens-Johnson syndrome/toxic epidermal necrolysis overlap* is used to describe cases with 10–30% detachment, and TEN is used to describe cases with >30% detachment.

Other blistering eruptions with mucositis associated with infections may be confused with SJS/TEN. Erythema multiforme (EM) associated with herpes simplex virus is characterized by mucosal involvement and target lesions often more acraly distributed and with limited skin detachment. *Mycoplasma* infection in children causes a clinically distinct presentation with prominent mucositis and limited blistering



FIGURE 74-9 Toxic epidermal necrolysis. (Photo credit: Lindy Peta Fox, MD, and Jubin Ryu, MD, PhD.)

lesions; some believe that this clinical entity is the syndrome originally described by Stevens and Johnson.

Patients with SJS, SJS/TEN, or TEN initially present with acute onset of painful skin lesions, fever $>39^{\circ}\text{C}$ (102.2°F), sore throat, and conjunctivitis resulting from mucosal lesions. Intestinal and pulmonary involvement is associated with a poor prognosis, as are a greater extent of epidermal detachment and older age. About 10% and 30% of SJS- and TEN-affected persons die from their disease, respectively. Drugs that most commonly cause SJS or TEN are sulfonamides, nevirapine (1 in 1000 risk of SJS or TEN), allopurinol, lamotrigine, aromatic anticonvulsants, and NSAIDs, specifically oxycam. Frozen-section skin biopsy may aid in rapid diagnosis. At this time, SJS and TEN have no proven effective treatment. The best results come from early diagnosis, immediate discontinuation of any suspected drug, supportive therapy, and paying close attention to ocular complications and infection. Systemic glucocorticoid therapy (prednisone 1–2 mg/kg) may be useful early in the evolution of the disease, but long-term systemic glucocorticoid use has been associated with higher mortality. Cyclosporine may be a possible therapy for SJS/TEN. After initial enthusiasm for the use of intravenous immunoglobulin (IVIG) in the treatment of SJS/TEN, some recent data questions whether IVIG benefits these patients. Randomized studies to more definitively assess the potential benefit of systemic glucocorticoids and IVIG are lacking and difficult to perform but are necessary.

Overlap Hypersensitivity Syndromes An important emerging concept in the clinical approach to severe drug eruptions is the presence of overlap syndromes, most notably DIHS with TEN-like features, DIHS with pustular eruption (AGEP-like), and AGEP with TEN-like features. In several case series of AGEP, 50% of cases had TEN-like or DRESS-like features, and 20% of cases had mucosal involvement resembling SJS/TEN. In one study, up to 20% of all severe drug eruptions had overlap features, suggesting that AGEP, DIHS, and SJS/TEN represent a clinical spectrum with common pathophysiologic mechanisms. Designation of a single diagnosis based on cutaneous and extracutaneous involvement may not always be possible in cases of hypersensitivity.

MANAGEMENT OF A PATIENT WITH A DRUG ERUPTION

There are four main questions to answer regarding an eruption:

1. Is it a drug reaction?
2. Is it a severe eruption or the onset of a form that may become severe?
3. Which drug(s) is (are) suspected, and which drug(s) should be withdrawn?
4. What is recommended for future use of drugs?

TABLE 74-2 CLINICAL AND LABORATORY FINDINGS ASSOCIATED WITH MORE SERIOUS DRUG-INDUCED CUTANEOUS CLINICAL FINDINGS

Cutaneous	
Generalized erythema	
Facial edema	
Skin pain	
Palpable purpura	
Target lesions	
Skin necrosis	
Blisters or epidermal detachment	
Positive Nikolsky's sign	
Mucous membrane erosions	
Urticaria	
Swelling of tongue	
General	
High fever (temperature $>40^{\circ}\text{C}$ [$>104^{\circ}\text{F}$])	
Enlarged lymph nodes	
Arthralgias or arthritis	
Shortness of breath, wheezing, hypotension	
Laboratory Results	
Eosinophil count $>1000/\mu\text{L}$	
Lymphocytosis with atypical lymphocytes	
Abnormal liver or kidney function tests	

Source: Adapted from JC Roujeau, RS Stern: *N Engl J Med* 331:1272, 1994.

EARLY DIAGNOSIS OF SEVERE ERUPTIONS

Rapid recognition of adverse drug reactions that may become serious or life threatening is paramount. **Table 74-2** lists clinical and laboratory features that, if present, suggest that the reaction may be serious. **Table 74-3** provides key features of the most serious adverse cutaneous reactions. Intensity of symptoms and rapid progression of signs should raise the suspicion of a severe eruption. Any doubt should lead to prompt consultation with a dermatologist and/or referral of the patient to a specialized center.

CONFIRMATION OF DRUG REACTION

The probability of drug etiology varies with the pattern of the reaction. Only fixed drug eruptions are always drug-induced. Morbilliform eruptions are usually viral in children and drug-induced in adults. Among severe reactions, drugs account for 10–20% of anaphylaxis and vasculitis and between 70–90% of AGEP, DIHS, SJS, or TEN. Skin biopsy helps in characterizing the reaction but does not indicate drug causality. Blood counts and liver and renal function tests are important for evaluating organ involvement. The association of mild elevation of liver enzymes and high eosinophil count is frequent but not specific for a drug reaction. Blood tests that could identify an alternative cause, antihistone antibody tests (to rule out drug-induced lupus), and serology or polymerase chain reaction for infections may be of great importance to determine a cause.

WHAT DRUG(S) TO SUSPECT AND WITHDRAW

Most cases of drug eruptions occur during the first course of treatment with a new medication. A notable exception is IgE-mediated urticaria and anaphylaxis that need presensitization and develop a few minutes to a few hours after rechallenge. Characteristic times of onset to drug reaction are as follows: 4–14 days for morbilliform eruptions, 2–4 days for AGEP, 5–28 days for SJS/TEN, and 14–48 days for DIHS. A drug chart, compiling information of all current and past medications/supplements and the timing of administration relative to the rash, is a key diagnostic tool to identifying the inciting drug. Medications introduced for the first time in the relevant time frame are prime suspects. Two other important elements to suspect causality at this stage are (1) previous experience with the drug in the population and (2) alternative etiologic candidates.

TABLE 74-3 CLINICAL FEATURES OF SEVERE CUTANEOUS DRUG REACTIONS

Diagnosis	Mucosal Lesions	Typical Skin Lesions	Frequent Signs and Symptoms	Alternative Causes Not Related to Drugs
Stevens-Johnson syndrome	Erosions usually at two or more sites	Small blisters on dusky purpuric macules or atypical targets; rare areas of confluence; detachment \leq 10% of body surface area	Most cases involve fever	10–20% cause not determined
Toxic epidermal necrolysis ^a	Erosions usually at two or more sites	Individual lesions like those seen in Stevens-Johnson syndrome; confluent erythema; outer layer of epidermis separates readily from basal layer with lateral pressure; large sheet of necrotic epidermis; total detachment of $>$ 30% of body surface area	Nearly all cases involve fever, "acute skin failure," leukopenia	10–20% cause not determined
Hypersensitivity syndrome	Infrequent	Severe exanthematous rash (may become purpuric), exfoliative dermatitis, facial edema	30–50% of cases involve fever, lymphadenopathy, hepatitis, nephritis, myocarditis, eosinophilia, atypical lymphocytes	Cutaneous lymphoma
Acute generalized exanthematous pustulosis	About 20% erosions (mouth, tongue)	Initially nonfollicular, small pustules overlying edematous erythema, sometimes leading to superficial erosions	Fever, burning, pruritus, facial swelling, leukocytosis, hypocalcemia	Infection
Serum sickness or reactions resembling serum sickness	Absent	Morbilloform lesions, sometimes with urticarial plaques (typically polycyclic)	Fever, arthralgias	Infection
Anticoagulant-induced necrosis	Infrequent	Erythema then purpura and necrosis, especially of fatty areas	Pain in affected areas	Disseminated intravascular coagulopathy, septicemia
Angioedema	Often involved	Urticaria or swelling of central part of face	Respiratory distress, cardiovascular collapse	Insect stings, foods

^aOverlap of Stevens-Johnson syndrome and toxic epidermal necrolysis have features of both and attachment of 10–30% of body surface area may occur.

Source: Adapted from JC Roujeau, RS Stern: *N Engl J Med* 331:1272, 1994.

The decision to continue or discontinue any medication will depend on the severity of the reaction, the severity of the primary disease, the degree of suspicion of causality, and the feasibility of an alternative safer treatment. In any potentially fatal drug reaction, elimination of all possible suspect drugs or unnecessary medications should be attempted. Some rashes may resolve when "treating through" a benign drug-related eruption. The decision to treat through an eruption should, however, remain the exception and withdrawal of every suspect drug the general rule. On the other hand, drugs that are not suspected and are important for the patient (e.g., antihypertensive agents) generally should not be quickly withdrawn. This approach prevents reluctance to future use of these agents.

RECOMMENDATION FOR FUTURE USE OF DRUGS

The aims are (1) to prevent the recurrence of the drug eruption and (2) not to compromise future treatments by contraindicating otherwise useful medications.

Begin with thorough assessment of drug causality. Drug causality is evaluated based on timing of the reaction, evaluation of other possible causes, effect of drug withdrawal or continuation, and knowledge of medications that have been associated with the observed reaction. Combination of these criteria leads to considering the causality as definite, probable, possible, or unlikely. The RegiSCAR group has proposed a useful algorithm called Algorithm of Drug Causality for Epidermal Necrolysis (ALDEN) to determine drug causality in SJS/TEN. A drug with a "definite" or "probable" causality should be contraindicated, a warning card or medical alert tag (e.g., wristband) should be given to the patient, and the drugs should be listed in the patient's medical chart as an allergy.

A drug with a "possible" causality may be submitted to further investigations depending on the expected need for future treatment.

A drug with "unlikely" causality or that has been continued when the reaction improved or was reintroduced without a reaction can be administered safely.

The usefulness of laboratory tests to determine causality is still debated. Many *in vitro* immunologic assays have been developed, but the predictive value of these tests has not been validated in any large series of affected patients; these tests exist primarily for research and not clinical purposes.

In some cases, diagnostic rechallenge may be appropriate, even for drugs with high rates of adverse reactions. Desensitization is often successful in HIV-infected patients with morbilliform eruptions to sulfonamides but is not recommended in HIV-infected patients who manifested erythroderma or a bullous reaction in response to their earlier sulfonamide exposure.

In patients with history suggesting immediate IgE-mediated reactions to penicillin, skin-prick testing with penicillins or cephalosporins has proved useful for identifying patients at risk of anaphylactic reactions to these agents. However, skin tests themselves carry a small risk of anaphylaxis. Negative skin tests do not totally rule out IgE-mediated reactivity, but the risk of anaphylaxis in response to penicillin administration in patients with negative skin tests is about 1%. In contrast, two-thirds of patients with a positive skin test experience an allergic response upon rechallenge.

For patients with delayed-type hypersensitivity, the clinical utility of skin tests is more questionable. At least one of a combination of several tests (prick, patch, and intradermal) is positive in 50–70% of patients with a reaction "definitely" attributed to a single medication. This low sensitivity corresponds to the observation that readministration of drugs with negative skin testing resulted in eruptions in 17% of cases.

CROSS-SENSITIVITY

Because of the possibility of cross-sensitivity among chemically related drugs, many physicians recommend avoidance of not only the medication that induced the reaction but also all drugs of the same pharmacologic class.

There are two types of cross-sensitivity. Reactions that depend on a pharmacologic interaction may occur with all drugs that target the same pathway, whether they are structurally similar or not. This is the case with angioedema caused by NSAIDs and ACE inhibitors. In this situation, the risk of recurrence varies from drug to drug in a particular class; however, avoidance of all drugs in the class is usually recommended. Immune recognition of structurally related drugs is the second mechanism by which cross-sensitivity occurs. A classic example is hypersensitivity to aromatic antiepileptics (barbiturates, phenytoin, carbamazepine) with up to 50% reaction to a second drug in patients who reacted to one. For other drugs, *in vitro* as well as *in vivo* data have suggested that cross-reactivity existed only between compounds with very similar chemical structures. Sulfamethoxazole-specific lymphocytes may be activated by other antibacterial sulfonamides but not diuretics, antidiabetic drugs, or anti-COX2 NSAIDs with a sulfonamide group. Approximately 10% of patients with penicillin allergies will also develop allergic reactions to cephalosporin class antibiotics.

Recent data suggest that although the risk of a drug eruption to another drug was increased in persons with a prior reaction, “cross-sensitivity” was probably not the explanation. As an example, persons with a history of an allergic-like reaction to penicillin were at higher risk to develop a reaction to antibacterial sulfonamides than to cephalosporins.

These data suggest that the list of drugs to avoid after a drug reaction should be limited to the causative one(s) and to a few very similar medications.

Because of growing evidence that some severe cutaneous reactions to drugs are associated with HLA genes, it is recommended that first-degree family members of patients with severe cutaneous reactions also should avoid these causative medications. This may be most relevant to sulfonamides and antiseizure medications.

Desensitization can be considered in those with a history of reaction to a medication that must be used again. Efficacy of such procedures has been demonstrated in cases of immediate reaction to penicillin and positive skin tests, anaphylactic reactions to platinum chemotherapy, and delayed reactions to sulfonamides in patients with AIDS. Various protocols are available, including oral and parenteral approaches. Oral desensitization appears to have a lower risk of serious anaphylactic reactions. However, desensitization carries the risk of anaphylaxis regardless of how it is performed and should be performed in monitored clinical settings such as an intensive care unit. After desensitization, many patients experience non-life-threatening reactions during therapy with the culprit drug.

REPORTING

Any severe reaction to drugs should be reported to a regulatory agency or to pharmaceutical companies (e.g., MedWatch, <http://www.fda.gov/Safety/MedWatch/default.htm>). Because severe reactions are too rare to be detected in premarketing clinical trials, spontaneous reports are of critical importance for early detection of unexpected life-threatening events. To be useful, the report should contain enough details to permit ascertainment of severity and drug causality. This enables recognition of similar cases that may be reported from several different sources.

ACKNOWLEDGMENT

We acknowledge the contribution of Dr. Jean-Claude Roujeau to this chapter in the 17th edition.

SOLAR RADIATION

Sunlight is the most visible and obvious source of comfort in the environment. The sun provides the beneficial effects of warmth and vitamin D synthesis. However, acute and chronic sun exposure also has pathologic consequences. Few effects of sun exposure beyond those affecting the skin have been identified, but cutaneous exposure to sunlight is the major cause of human skin cancer and can have immunosuppressive effects as well.

The sun’s energy reaching the earth’s surface is limited to components of the ultraviolet (UV) spectrum, the visible spectrum, and portions of the infrared spectrum. The cutoff at the short end of the UV spectrum at ~290 nm is due primarily to stratospheric ozone—formed by highly energetic ionizing radiation—that prevents penetration to the earth’s surface of the shorter, more energetic, potentially more harmful wavelengths of solar radiation. Indeed, concern about destruction of the ozone layer by chlorofluorocarbons released into the atmosphere has led to international agreements to reduce production of those chemicals.

Measurements of solar flux showed a twentyfold regional variation in the amount of energy at 300 nm that reaches the earth’s surface. This variability relates to seasonal effects, the path that sunlight traverses through ozone and air, the altitude (a 4% increase for each 300 m of elevation), the latitude (increasing intensity with decreasing latitude), and the amount of cloud cover, fog, and pollution.

The major components of the photobiologic action spectrum that are capable of affecting human skin include the UV and visible wavelengths between 290 and 700 nm. In addition, the wavelengths beyond 700 nm in the infrared spectrum primarily emit heat and in certain circumstances may exacerbate the pathologic effects of energy in the UV and visible spectra.

The UV spectrum reaching the earth represents <10% of total incident solar energy and is arbitrarily divided into two major segments, UV-B and UV-A, which constitute the wavelengths from 290 to 400 nm. UV-B consists of wavelengths between 290 and 320 nm. This portion of the photobiologic action spectrum is the most efficient in producing redness or erythema in human skin and thus is sometimes known as the “sunburn spectrum.” UV-A includes wavelengths between 320 and 400 nm and is ~1000-fold less efficient in producing skin redness than is UV-B.

The wavelengths between 400 and 700 nm are visible to the human eye. The photon energy in the visible spectrum is not capable of damaging human skin in the absence of a photosensitizing chemical. Without the absorption of energy by a molecule, there can be no photosensitivity. Thus, the *absorption spectrum* of a molecule is defined as the range of wavelengths it absorbs, whereas the *action spectrum* for an effect of incident radiation is defined as the range of wavelengths that evoke the response.

Photosensitivity occurs when a photon-absorbing chemical (*chromophore*) present in the skin absorbs incident energy, becomes excited, and transfers the absorbed energy to various structures or to molecular oxygen.

UV RADIATION (UVR) AND SKIN STRUCTURE AND FUNCTION

Human skin consists of two major compartments: the outer epidermis, which is a stratified squamous epithelium, and the underlying dermis, which is rich in matrix proteins such as collagens and elastin. Both compartments are susceptible to damage from sun exposure. The epidermis and the dermis contain several chromophores capable of absorbing incident solar energy, including nucleic acids, proteins, and lipids. The outermost epidermal layer, the stratum corneum, is a major absorber of UV-B, and <10% of incident UV-B wavelengths penetrate through the epidermis to the dermis. Approximately 3% of radiation

below 300 nm, 20% of radiation below 360 nm, and 33% of short visible radiation reach the basal cell layer in untanned human skin. In contrast, UV-A readily penetrates to the dermis and is capable of altering structural and matrix proteins that contribute to photoaging of chronically sun-exposed skin, particularly in individuals of light complexion. Thus, longer wavelengths can penetrate more deeply into the skin.

Molecular Targets for UVR-Induced Skin Effects Epidermal DNA—predominantly in keratinocytes and in Langerhans cells, which are dendritic antigen-presenting cells—absorbs UV-B and undergoes structural changes between adjacent pyrimidine bases (thymine or cytosine), including the formation of cyclobutane dimers and 6,4-photoproducts. These structural changes are potentially mutagenic and are found in most basal cell and squamous cell carcinomas (BCCs and SCCs, respectively). They can be repaired by cellular mechanisms that result in their recognition and excision and the restoration of normal base sequences. The efficient repair of these structural aberrations is crucial, since individuals with defective DNA repair are at high risk for the development of cutaneous cancer. For example, patients with xeroderma pigmentosum, an autosomal recessive disorder, have a variably deficient repair of UV-induced photoproducts. The skin of these patients often shows the dry, leathery appearance of prematurely photoaged skin, and these patients have an increased frequency of skin cancer already in the first two decades of life. Studies in transgenic mice have verified the importance of functional genes that regulate these repair pathways in preventing the development of UV-induced skin cancer. DNA damage in Langerhans cells may also contribute to the known immunosuppressive effects of UV-B (see “Photoimmunology,” below).

In addition to DNA, molecular oxygen is a target for incident solar UVR, leading to the generation of reactive oxygen species (ROS). These ROS can damage skin components, such as epidermal lipids—either free lipids in the stratum corneum or cell membrane lipids. UVR also can target proteins, leading to increased cross-linking and degradation of matrix proteins in the dermis and accumulation of abnormal dermal elastin leading to photoaging changes known as *solar elastosis*.

Cutaneous Optics and Chromophores *Chromophores* are endogenous or exogenous chemical components that can absorb physical energy. Endogenous chromophores are of two types: (1) normal components of skin, including nucleic acids, proteins, lipids, and 7-dehydrocholesterol (the precursor of vitamin D); and (2) components that are synthesized elsewhere in the body and that circulate in the bloodstream and diffuse into the skin, such as porphyrins. Normally, only trace amounts of porphyrins are present in the skin, but, in selected diseases known as the *porphyrias* (Chap. 430), porphyrins are released into the circulation in increased amounts from the bone marrow and the liver and are transported to the skin, where they absorb incident energy both in the Soret band (around 400 nm; short visible) and, to a lesser extent, in the red portion of the visible spectrum (580–660 nm). This energy absorption results in the generation of ROS that can mediate structural damage to the skin, manifested as erythema, edema, urticaria, or blister formation. It is of interest that photoexcited porphyrins are currently used in the treatment of nonmelanoma skin cancers and their precursor lesions, actinic keratoses. Known as *photodynamic therapy*, this modality generates ROS in the skin, leading to cell death. Topical photosensitizers used in photodynamic therapy are the porphyrin precursors 5-aminolevulinic acid and methyl aminolevulinic acid, which are converted to porphyrins in the skin. It is believed that photodynamic therapy targets tumor cells for destruction more selectively than it targets adjacent nonneoplastic cells. The efficacy of such therapy requires appropriate timing of the application of methyl aminolevulinic acid or 5-aminolevulinic acid to the affected skin followed by exposure to artificial sources of visible light. High-intensity blue light has been used successfully for the treatment of thin actinic keratoses. Red light has a longer wavelength, penetrates more deeply into the skin, and is more beneficial in the treatment of superficial BCCs.

Acute Effects of Sun Exposure The acute effects of skin exposure to sunlight include sunburn and vitamin D synthesis.

SUNBURN This painful skin condition is an acute inflammatory response of the skin, predominantly to UV-B. Generally, an individual’s ability to tolerate sunlight is inversely proportional to that individual’s degree of melanin pigmentation. Melanin, a complex polymer of tyrosine derivatives, is synthesized in specialized epidermal dendritic cells known as *melanocytes* and is packaged into *melanosomes* that are transferred via dendritic processes into *keratinocytes*, thereby providing photoprotection and simultaneously darkening the skin. Sun-induced melanogenesis is a consequence of increased tyrosinase activity in melanocytes. Central to the suntan response is the melanocortin-1 receptor (*MC1R*), and mutations in this gene contribute to the wide variation in human skin and hair color; individuals with red hair and fair skin typically have low *MC1R* activity. Genetic studies have revealed additional genes that influence skin color variation in humans, such as the gene for tyrosinase (*TYR*) and the genes *APBA2*[*OCA2*], *SLC45A2*, and *SLC24A5*. The human *MC1R* gene encodes a G protein-coupled receptor that binds α -melanocyte-stimulating hormone, which is secreted in the skin mainly by keratinocytes in response to UVR. The UV-induced expression of this hormone is controlled by the tumor suppressor p53, and absence of functional p53 attenuates the tanning response. Activation of the melanocortin receptor leads to increased intracellular cyclic adenosine 5'-monophosphate (cAMP) and protein kinase A activation, resulting in an increased transcription of the microphthalmia-associated transcription factor (MITF), which stimulates melanogenesis. Since the precursor of α -melanocyte-stimulating hormone, proopiomelanocortin, is also the precursor of β -endorphins, UVR may result in not only increased pigmentation but also in increased β -endorphin production, an effect that has been hypothesized to promote sun-seeking behaviors.

The Fitzpatrick classification of human skin phototypes is based on the efficiency of the epidermal-melanin unit, which usually can be ascertained by asking an individual two questions: (1) Do you burn after sun exposure? (2) Do you tan after sun exposure? The answers to these questions permit division of the population into six skin types, varying from type I (always burn, never tan) to type VI (never burn, always tan) (Table 75-1).

Sunburn erythema is due to vasodilation of dermal blood vessels. There is a lag time (usually 4–12 h) between skin exposure to sunlight and the development of visible redness. The action spectrum for sunburn erythema includes UV-B and UV-A, although UV-B is much more efficient than UV-A in evoking the response. However, UV-A may contribute to sunburn erythema at midday, when much more UV-A than UV-B is present in the solar spectrum. The erythema that accompanies the inflammatory response induced by UVR results from the orchestrated release of cytokines along with growth factors and the generation of ROS. Furthermore, UV-induced activation of nuclear factor κ B-dependent gene transcription can augment release of several proinflammatory cytokines and vasoactive mediators. These cytokines and mediators accumulate locally in sunburned skin, providing chemotactic factors that attract neutrophils, macrophages, and T lymphocytes, which promote the inflammatory response. UVR also stimulates infiltration of inflammatory cells through induced expression of adhesion molecules such as E-selectin and intercellular adhesion molecule 1 on endothelial cells and keratinocytes. UVR also has been shown to activate phospholipase A₂, resulting in increases in eicosanoids such as prostaglandin E₂, which is known to be a potent inducer of sunburn erythema. The role

TABLE 75-1 SKIN TYPE AND SUNBURN SENSITIVITY (FITZPATRICK CLASSIFICATION)

Type	Description
I	Always burn, never tan
II	Always burn, sometimes tan
III	Sometimes burn, sometimes tan
IV	Sometimes burn, always tan
V	Never burn, sometimes tan
VI	Never burn, always tan

of eicosanoids in this reaction has been verified by studies showing that nonsteroidal anti-inflammatory drugs (NSAIDs) can reduce it.

Epidermal changes in sunburn include the induction of “sunburn cells,” which are keratinocytes undergoing p53-dependent apoptosis as a defense, with elimination of cells that harbor UV-B-induced structural DNA damage.

VITAMIN D SYNTHESIS AND PHOTOCHEMISTRY Cutaneous exposure to UV-B causes photolysis of epidermal 7-dehydrocholesterol, converting it to pre-vitamin D₃, which then undergoes temperature-dependent isomerization to form the stable hormone vitamin D₃. This compound diffuses to the dermal vasculature and circulates to the liver and kidney, where it is converted to the dihydroxylated functional hormone 1,25-dihydroxyvitamin D₃. Vitamin D metabolites from the circulation and those produced in the skin itself can augment epidermal differentiation signaling and inhibit keratinocyte proliferation. These effects are exploited therapeutically in psoriasis with the topical application of synthetic vitamin D analogues. In addition, vitamin D is increasingly thought to have beneficial effects in several other inflammatory conditions, and some evidence suggests that—besides its classic physiologic effects on calcium metabolism and bone homeostasis—it is associated with a reduced risk of various internal malignancies. There is controversy regarding the risk-to-benefit ratio of sun exposure in vitamin D homeostasis. At present, it is important to emphasize that no clear-cut evidence suggests that the use of sunscreens substantially diminishes vitamin D levels. Since aging also substantially decreases the ability of human skin to photocatalytically produce vitamin D₃, the widespread use of sunscreens that filter out UV-B has led to concerns that the elderly might be unduly susceptible to vitamin D deficiency. However, the amount of sunlight needed to produce sufficient vitamin D is small and does not justify the risks of skin cancer and other types of photodamage linked to increased sun exposure or tanning behavior. Nutritional supplementation of vitamin D is a preferable strategy for patients with vitamin D deficiency.

Chronic Effects of Sun Exposure: Nonmalignant The clinical features of photoaging (*dermatoheliosis*) consist of wrinkling, blotchiness, and telangiectasia as well as a roughened, irregular, “weather-beaten” leathery appearance.

UVR is important in the pathogenesis of photoaging in human skin, and ROS are likely involved. The dermis and its connective tissue matrix are major targets for sun-associated chronic damage that manifests as solar elastosis, a massive increase in thickened irregular masses of abnormal-appearing elastic fibers. Collagen fibers are also abnormally clumped in the deeper dermis of sun-damaged skin. The chromophore(s), the action spectra, and the specific biochemical events orchestrating these changes are only partially understood, although more deeply penetrating UV-A seems to be primarily involved. Chronologically aged sun-protected skin and photoaged skin share important molecular features, including connective tissue damage and elevated levels of matrix metalloproteinases (MMPs). MMPs are enzymes involved in the degradation of the extracellular matrix. UV-A induces expression of some MMPs, including MMP-1 and MMP-3, leading to increased collagen breakdown. In addition, UV-A reduces type I procollagen mRNA expression. Thus, chronic UVR alters the structure and function of dermal collagen. On the basis of these observations, it is not surprising that high-dose UV-A phototherapy may have beneficial effects in some patients with localized fibrotic diseases of the skin, such as localized scleroderma.

Chronic Effects of Sun Exposure: Malignant One of the major known consequences of chronic excessive skin exposure to sunlight is non-melanoma skin cancer. The two most common types of non-melanoma skin cancer are BCC and SCC (**Chap. 105**). A model for skin cancer induction involves three major steps: initiation, promotion, and progression. Exposure of human skin to sunlight results in *initiation*, a step by which structural (mutagenic) changes in DNA evoke an irreversible alteration in the target cell (keratinocyte) that begins the tumorigenic process. Exposure to a tumor initiator such as UV-B is believed to be a necessary but not a sufficient step in the malignant

process, since initiated skin cells not exposed to tumor promoters generally do not develop tumors. The second stage in tumor development is *promotion*, a multistep process by which chronic exposure to sunlight evokes further changes that culminate in the clonal expansion of initiated cells and cause the development, over many years, of premalignant growths known as *actinic keratoses*, a minority of which may progress to form SCCs. As a result of extensive studies, it seems clear that UV-B is a *complete carcinogen*, meaning that it can act as both a tumor initiator and a tumor promoter. The third and final step in the malignant process is *malignant conversion* of benign precursors into malignant lesions, a process thought to require additional genetic alterations.

On a molecular level, skin carcinogenesis results from the accumulation of gene mutations that cause inactivation of tumor suppressors, activation of oncogenes, or reactivation of cellular signaling pathways that normally are expressed only during epidermal embryologic development. Accumulation of mutations in the tumor-suppressor gene *p53* secondary to UV-induced DNA damage occurs in both SCCs and BCCs and is important in promoting skin carcinogenesis. Indeed, the majority of both human and murine UV-induced skin cancers have characteristic *p53* mutations (C → T and CC → TT transitions). Studies in mice have shown that sunscreens can substantially reduce the frequency of these signature mutations in *p53* and inhibit the induction of tumors.

BCCs also harbor inactivating mutations in the tumor-suppressor gene *patched*, which result in activation of the sonic hedgehog signaling pathway and increased cell proliferation. Thus, these tumors can manifest mutations in tumor suppressors (*p53* and *patched*) or oncogenes (*smoothed*). New evidence links alterations in the Wnt/β-catenin signaling pathway, which is known to be critical for hair follicle development, to skin cancer as well. Thus interactions between this pathway and the hedgehog signaling pathway appear to be involved in both skin carcinogenesis and embryologic development of the skin and hair follicles.

Clonal analysis in mouse models of BCC revealed that tumor cells arise from long-term resident progenitor cells of the interfollicular epidermis and the upper infundibulum of the hair follicle. These BCC-initiating cells are reprogrammed to resemble embryonic hair follicle progenitors, whose tumor-initiating ability depends on activation of the Wnt/β-catenin signaling pathway.

SCC initiation occurs both in the interfollicular epidermis and in the hair follicle bulge stem cell populations. In mouse models, the combination of mutant K-Ras and *p53* is sufficient to induce invasive SCCs from these cell populations.

The transcription factor *Myc* is important for stem cell maintenance in the skin, and oncogenic activation of *Myc* has been implicated in the development of BCCs and SCCs. Thus, nonmelanoma skin cancer involves mutations and alterations in multiple genes and pathways that occur as a result of their chronic accumulation driven by exposure to environmental factors such as UVR.

Epidemiologic studies have linked excessive sun exposure to an increased risk of nonmelanoma cancers and melanoma of the skin; the evidence is far more direct for nonmelanoma skin cancers (BCCs and SCCs) than for melanoma. Approximately 80% of nonmelanoma skin cancers develop on sun-exposed body areas, including the face, neck, and hands. Major risk factors include male sex, childhood sun exposures, older age, fair skin, and residence at latitudes relatively close to the equator. Individuals with darker-pigmented skin have a lower risk of skin cancer than do fair-skinned individuals. More than 2 million individuals in the United States develop nonmelanoma skin cancer annually, and the lifetime risk that a fair-skinned individual will develop such a neoplasm is estimated at ~15%. The incidence of non-melanoma skin cancer in the population is increasing at a rate of 2–3% per year. One potential explanation is the widespread use of indoor tanning. It is estimated that 30 million people tan indoors in the United States annually, including >2 million adolescents.

The relationship of sun exposure to melanoma development is less direct, but strong evidence supports an association. Clear-cut risk

factors include a positive family or personal history of melanoma and multiple dysplastic nevi. Melanomas can occur during adolescence; the implication is that the latent period for tumor growth is shorter than that for nonmelanoma skin cancer. For reasons that are only partially understood, melanomas are among the most rapidly increasing human malignancies (**Chap. 105**). Epidemiologic studies indicate that indoor tanning is a risk factor for melanoma, which may contribute to the increasing incidence of melanoma formation. Furthermore, epidemiologic studies suggest that life in a sunny climate from birth or early childhood may increase the risk of melanoma development. In general, risk does not correlate with cumulative sun exposure but may be related to the duration and extent of exposure in childhood.

However, in contrast to nonmelanoma skin cancers, melanoma frequently develops in sun-protected skin, and oncogenic mutations in melanoma may also not be UVR-signature mutations; these observations suggest that UVR-independent factors contribute to melanomagenesis. Low MC1R activity leads to production of the red/yellow pheomelanin pigment in individuals with red hair and fair skin, while high MC1R activity results in increased production of the black/brown eumelanin. Experiments in mice suggest that high pheomelanin content in skin (as in individuals with red hair and fair skin) leads to a UVR-independent increase in the risk of melanoma through a mechanism that involves oxidative damage. Thus, both UVR-dependent and UVR-independent factors are likely to contribute to melanoma formation.

Photoimmunology Exposure to solar radiation causes both local immunosuppression (through inhibition of immune responses to antigens applied at the irradiated site) and systemic immunosuppression (through inhibition of immune responses to antigens applied at remote, unirradiated sites). For example, human skin exposure to modest doses of UV-B can deplete the epidermal antigen-presenting cells known as Langerhans cells, thereby reducing the degree of allergic sensitization to application of the potent contact allergen dinitrochlorobenzene at the irradiated skin site.

An example of the systemic immunosuppressive effects of higher doses of UVR is the diminished immunologic response to antigens introduced either epicutaneously or intracutaneously at sites distant from the irradiated site. Various immunomodulatory factors and immune cells have been implicated in UVR-induced systemic immunosuppression, including tumor necrosis factor α , interleukin 4, interleukin 10, *cis*-urocanic acid, and eicosanoids. Experimental evidence suggests that prostaglandin E_2 signaling through prostaglandin E receptor subtype 4 mediates UVR-induced systemic immunosuppression by elevating the number of regulatory T cells, and this effect can be inhibited with NSAIDs.

The major chromophores in the upper epidermis that are known to initiate UV-mediated immunosuppression include DNA, *trans*-urocanic acid, and membrane components. The action spectrum for UV-induced immunosuppression closely mimics the absorption spectrum of DNA. Pyrimidine dimers in Langerhans cells may inhibit antigen presentation. The absorption spectrum of epidermal urocanic acid closely mimics the action spectrum for UV-B-induced immunosuppression. Urocanic acid is a metabolic product of the essential amino acid histidine and accumulates in the upper epidermis through breakdown of the histidine-rich protein filaggrin due to the absence of its catabolizing enzyme in keratinocytes. Urocanic acid is synthesized as a *trans*-isomer, and UV-induced *trans-cis* isomerization of urocanic acid in the stratum corneum drives immunosuppression. *Cis*-urocanic acid may exert its immunosuppressive effects through a variety of mechanisms, including inhibition of antigen presentation by Langerhans cells.

One important consequence of chronic sun exposure and associated immunosuppression is an enhanced risk of skin cancer. In part, UV-B activates regulatory T cells that suppress antitumor immune responses via interleukin 10 expression, whereas, in the absence of high UV-B exposure, epidermal Langerhans cells present tumor-associated antigens and induce protective immunity, thereby inhibiting skin tumorigenesis. UV-induced DNA damage is a major molecular trigger of this immunosuppressive effect.

Perhaps the most graphic demonstration of the role of immunosuppression in enhancing the risk of nonmelanoma skin cancer comes from studies of organ transplant recipients who require lifelong immunosuppressive/antirejection drug regimens. More than 50% of organ transplant recipients develop BCCs and SCCs, and these cancers are the most common types of malignancies arising in these patients. Rates of BCC and SCC increase with the duration and degree of immunosuppression. These patients ideally should be screened prior to organ transplantation, be monitored closely thereafter, and adhere to rigorous photoprotection measures, including the use of sunscreens and protective clothing as well as sun avoidance. Notably, immunosuppressive drugs that target the mTOR pathway, such as sirolimus and everolimus, may reduce the risk of nonmelanoma skin cancer in organ transplant recipients from that associated with the use of calcineurin inhibitors (cyclosporine and tacrolimus), which may contribute to nonmelanoma skin cancer formation not only through their immunosuppressive effects but also through suppression of p53-dependent cancer cell senescence pathways independent of host immunity.

PHOTOSENSITIVITY DISEASES

The diagnosis of photosensitivity requires elicitation of a careful history in order to define the duration of signs and symptoms, the length of time between exposure to sunlight and the development of subjective symptoms, and visible changes in the skin. The age of onset can also be a helpful diagnostic clue; for example, the acute photosensitivity of erythropoietic protoporphyria almost always begins in childhood, whereas the chronic photosensitivity of porphyria cutanea tarda (PCT) typically begins in the fourth and fifth decades of life. A patient's history of exposure to topical and systemic drugs and chemicals may provide important diagnostic clues. Many classes of drugs can cause photosensitivity on the basis of either phototoxicity or photoallergy. Fragrances such as musk ambrette that were previously present in numerous cosmetic products are also potent photosensitizers.

Examination of the skin may offer important clues. Anatomic areas that are naturally protected from direct sunlight, such as the hairy scalp, the upper eyelids, the retroauricular areas, and the infranasal and submental regions, may be spared, whereas exposed areas show characteristic features of the pathologic process. These anatomic localization patterns are often helpful, but not infallible, in making the diagnosis. For example, airborne contact sensitizers that are blown onto the skin may produce dermatitis that can be difficult to distinguish from photosensitivity despite the fact that such material may trigger skin reactivity in areas shielded from direct sunlight.

Many dermatologic conditions may be caused or aggravated by sunlight (**Table 75-2**). The role of light in evoking these responses may be dependent on genetic abnormalities ranging from well-described defects in DNA repair that occur in xeroderma pigmentosum to the inherited abnormalities in heme synthesis that characterize the porphyrias. The chromophore has been identified in certain photosensitivity diseases, but the energy-absorbing agent remains unknown in the majority.

Polymorphous Light Eruption A common type of photosensitivity disease is *polymorphous light eruption* (PMLE). Many affected individuals never seek medical attention because the condition is often transient, becoming manifest in the spring with initial sun exposure but then subsiding spontaneously with continuing exposure, a phenomenon known as "hardening." The major manifestations of PMLE include (often intensely) pruritic erythematous papules that may coalesce into plaques in a patchy distribution on exposed areas of the trunk and forearms. The face is usually less seriously involved. Whereas the morphologic skin findings remain similar for each patient with subsequent recurrences, significant interindividual variations in skin findings are characteristic (hence the term "polymorphous").

A skin biopsy and phototest procedures in which skin is exposed to multiple erythematous doses of UV-A and UV-B may aid in the diagnosis. The action spectrum for PMLE is usually within these portions of the solar spectrum.

Whereas the treatment of an acute flare of PMLE may require topical or systemic glucocorticoids, approaches to preventing PMLE are

TABLE 75-2 CLASSIFICATION OF PHOTOSENSITIVITY DISEASES

Type	Disease
Genetic	Erythropoietic porphyria
	Erythropoietic protoporphyria
	Porphyria cutanea tarda—familial
	Variagate porphyria
	Hepatoerythropoietic porphyria
	Albinism
	Xeroderma pigmentosum
	Rothmund-Thomson syndrome
	Bloom syndrome
	Cockayne syndrome
	Kindler syndrome
	Phenylketonuria
	Metabolic
Hartnup disease	
Kwashiorkor	
Pellagra	
Carcinoid syndrome	
Phototoxic	
	Internal
External	Drugs, plants, food
Photoallergic	
	Immediate
Delayed	Drug photoallergy Persistent light reaction/chronic actinic dermatitis
Neoplastic and degenerative	Photoaging
	Actinic keratosis
	Melanoma and nonmelanoma skin cancer
Idiopathic	Polymorphous light eruption
	Hydroa aestivale
	Actinic prurigo
Photoaggravated	Lupus erythematosus
	Systemic
	Subacute cutaneous
	Discoid
	Dermatomyositis
	Herpes simplex
	Lichen planus actinicus
	Acne vulgaris (aestivale)

important and include the use of high-SPF and high UVA-protection broad-spectrum sunscreens as well as the induction of “hardening” by the cautious administration of artificial UV-B (broad-band or narrow-band) and/or UV-A radiation or the use of psoralen plus UV-A (PUVA) photochemotherapy for 2–4 weeks before initial sun exposure. Such prophylactic phototherapy or photochemotherapy at the beginning of spring may prevent the occurrence of PMLE throughout the summer.

Phototoxicity and Photoallergy These photosensitivity disorders are related to the topical or systemic administration of drugs and other chemicals. Both reactions require the absorption of energy by a drug or chemical with consequent production of an excited-state photosensitizer that can transfer its absorbed energy to a bystander molecule or to molecular oxygen, thereby generating tissue-destructive chemical species, including ROS.

Phototoxicity is a nonimmunologic reaction that can be caused by drugs and chemicals, a few of which are listed in Table 75-3. The usual clinical manifestations include erythema resembling a sunburn reaction that quickly desquamates, or “peels,” within several days. In addition, edema, vesicles, and bullae may occur.

TABLE 75-3 DRUGS THAT MAY CAUSE A PHOTOTOXIC REACTION

Drug	Topical	Systemic
Amiodarone		+
Dacarbazine		+
Fluoroquinolones		+
5-Fluorouracil	+	+
Furosemide		+
Nalidixic acid		+
Phenothiazines		+
Psoralens	+	+
Retinoids	+/-	+
Sulfonamides		+
Sulfonylureas		+
Tetracyclines		+
Thiazides		+
Vinblastine		+

Photoallergy is much less common and is distinct in that it is an immunopathologic process. The excited-state photosensitizer may create highly unstable haptenic free radicals that bind covalently to macromolecules to form a functional antigen capable of evoking a delayed-type hypersensitivity response. Some drugs and chemicals that can produce photoallergy are listed in Table 75-4. The clinical manifestations typically differ from those of phototoxicity in that an intensely pruritic eczematous dermatitis tends to predominate and evolves into lichenified, thickened, “leathery” changes in sun-exposed areas. A small subset (perhaps 5–10%) of patients with photoallergy may develop a persistent exquisite hypersensitivity to light even when the offending drug or chemical is identified and eliminated, a condition known as *persistent light reaction*.

A very uncommon type of persistent photosensitivity is known as *chronic actinic dermatitis*. The affected patients are typically elderly men with a long history of preexisting allergic contact dermatitis or photosensitivity. These individuals are usually exquisitely sensitive to UV-B, UV-A, and visible wavelengths.

Phototoxicity and photoallergy often can be diagnostically confirmed by phototest procedures. In patients with suspected phototoxicity, determining the minimal erythmal dose (MED) while the patient is exposed to a suspected agent and then repeating the MED after discontinuation of the agent may provide a clue to the causative drug or chemical. Photopatch testing can be performed to confirm the diagnosis of photoallergy. In this simple variant of ordinary patch testing, a series of known photoallergens is applied to the skin in duplicate, and one set is irradiated with a suberythmal dose of UV-A. The development of eczematous changes at sites exposed to sensitizer and light is a positive result. The characteristic abnormality in patients with persistent light reaction is a diminished threshold to erythema evoked

TABLE 75-4 DRUGS THAT MAY CAUSE A PHOTOALLERGIC REACTION

Drug	Topical	Systemic
6-Methylcoumarin	+	
Aminobenzoic acid and esters	+	
Bithionol	+	
Chlorpromazine		+
Diclofenac		+
Fluoroquinolones		+
Halogenated salicylanilides	+	
Hypericin (St. John's wort)	+	+
Musk ambrette	+	
Piroxicam		+
Promethazine		+
Sulfonamides		+
Sulfonylureas		+

by UV-B. Patients with chronic actinic dermatitis usually manifest a broad spectrum of UV hyperresponsiveness and require meticulous photoprotection, including avoidance of sun exposure, use of high-SPF (>30) sunscreens, and, in severe cases, systemic immunosuppression, such as with azathioprine.

The management of drug photosensitivity involves first and foremost the elimination of exposure to the chemical agents responsible for the reaction and the minimization of sun exposure. The acute symptoms of phototoxicity may be ameliorated by cool moist compresses, topical glucocorticoids, and systemically administered NSAIDs. In severely affected individuals, a rapidly tapered course of systemic glucocorticoids may be useful. Judicious use of analgesics may be necessary.

Photoallergic reactions require a similar management approach. Furthermore, patients with persistent light reaction and chronic actinic dermatitis must be meticulously protected against light exposure. In selected patients to whom chronic systemic high-dose glucocorticoids pose unacceptable risks, it may be necessary to employ an immunosuppressive drug such as azathioprine, cyclophosphamide, cyclosporine, or mycophenolate mofetil.

Porphyria The porphyrias (Chap. 430) are a group of diseases that have in common inherited or acquired derangements in the synthesis of heme. Heme is an iron-chelated tetrapyrrole or porphyrin, and the nonmetal chelated porphyrins are potent photosensitizers that absorb light intensely in both the short (400–410 nm) and the long (580–650 nm) portions of the visible spectrum.

Heme cannot be reutilized and must be synthesized continuously. The two body compartments with the largest capacity for its production are the bone marrow and the liver. Accordingly, the porphyrias originate in one or the other of these organs, with an end result of excessive endogenous production of potent photosensitizing porphyrins. The porphyrins circulate in the bloodstream and diffuse into the skin, where they absorb solar energy, become photoexcited, generate ROS, and evoke cutaneous photosensitivity. The mechanism of porphyrin photosensitization is known to be photodynamic, or oxygen-dependent, and is mediated by ROS such as singlet oxygen and superoxide anions.

Porphyria cutanea tarda is the most common type of porphyria and is associated with decreased activity of the enzyme uroporphyrinogen decarboxylase. There are two basic types of PCT: (1) the sporadic or acquired type, generally seen in individuals ingesting ethanol or receiving estrogens; and (2) the inherited type, in which there is autosomal dominant transmission of deficient enzyme activity. Both forms are associated with increased hepatic iron stores.

In both types of PCT, the predominant feature is chronic photosensitivity characterized by increased fragility of sun-exposed skin, particularly areas subject to repeated trauma such as the dorsa of the hands, the forearms, the face, and the ears. The predominant skin lesions are vesicles and bullae that rupture, producing moist erosions (often with a hemorrhagic base) that heal slowly, with crusting and purplish discoloration of the affected skin. Hypertrichosis, mottled pigmentary change, and scleroderma-like induration are associated features. The diagnosis can be confirmed biochemically by measurement of urinary porphyrin excretion, plasma porphyrin assay, and assay of erythrocyte and/or hepatic uroporphyrinogen decarboxylase. Multiple mutations of the uroporphyrinogen decarboxylase gene have been identified in human populations. Some patients with PCT have associated mutations in the *HFE* gene, which is linked to hemochromatosis; these mutations could contribute to the iron overload seen in PCT, although iron status as measured by serum ferritin, iron levels, and transferrin saturation is no different from that in PCT patients without *HFE* mutations. Prior hepatitis C virus infection appears to be an independent risk factor for PCT.

Treatment of PCT consists of repeated phlebotomies to diminish the excessive hepatic iron stores and/or intermittent low doses of chloroquine and hydroxychloroquine. Long-term remission of the disease can be achieved if the patient eliminates exposure to porphyrinogenic agents and prolonged exposure to sunlight.

Erythropoietic protoporphyria originates in the bone marrow and is due to a decrease in the mitochondrial enzyme ferrochelatase secondary to numerous gene mutations. The major clinical features include acute photosensitivity characterized by subjective burning and stinging of exposed skin that often develops during or just after sun exposure. There may be associated skin swelling and, after repeated episodes, a waxlike scarring.

The diagnosis is confirmed by demonstration of elevated levels of free erythrocyte protoporphyrin. Detection of increased plasma protoporphyrin helps distinguish erythropoietic protoporphyria from lead poisoning and iron-deficiency anemia, in both of which erythrocyte protoporphyrin levels are elevated in the absence of cutaneous photosensitivity and elevated plasma protoporphyrin levels.

Treatment includes reduction of sun exposure and oral administration of the carotenoid β -carotene, which is an effective scavenger of free radicals. This drug increases tolerance to sun exposure in some affected individuals, although it has no effect on deficient ferrochelatase.

An algorithm for managing patients with photosensitivity is presented in Fig. 75-1.

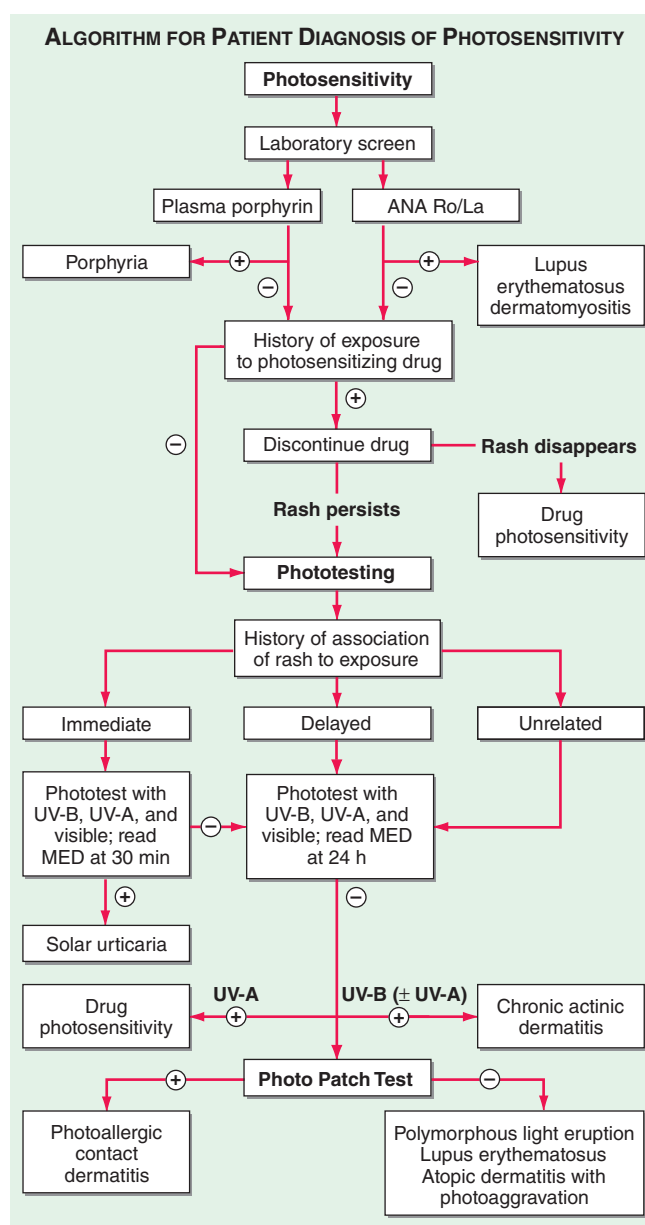


FIGURE 75-1 Algorithm for the diagnosis of a patient with photosensitivity. ANA, antinuclear antibody; MED, minimal erythral dose; UV-A and UV-B, ultraviolet spectrum segments including wavelengths of 320–400 nm and 290–320 nm, respectively.

PHOTOPROTECTION

Since photosensitivity of the skin results from exposure to sunlight, it follows that absolute avoidance of sunlight will eliminate these disorders. However, contemporary lifestyles make this approach impractical for most individuals. Thus better approaches to photoprotection have been sought.

Natural photoprotection is provided by structural proteins in the epidermis, particularly keratins and melanin. The amount of melanin and its distribution in cells are genetically regulated, and individuals of darker complexion (skin types IV–VI) are at decreased risk for the development of acute sunburn and cutaneous malignancy.

Other forms of photoprotection include clothing and sunscreens. Clothing constructed of tightly woven sun-protective fabrics, irrespective of color, affords substantial protection. Wide-brimmed hats, long sleeves, and trousers all reduce direct exposure. Sunscreens are now considered over-the-counter drugs, and a monograph from the U.S. Food and Drug Administration (FDA) has recognized category I ingredients as safe and effective. Those ingredients are listed in [Table 75-5](#). Sunscreens are rated for their photoprotective effect by their sun protection factor (SPF). The SPF is simply a ratio of the time required to produce sunburn erythema with and without sunscreen application. The SPF of most sunscreens reflects protection from UV-B but not from UV-A. The FDA monograph stipulates that sunscreens must be rated on a scale ranging from minimal (SPF ≤ 2 and <12) to moderate (SPF ≥ 12 and <30) to high (SPF ≥ 30 , labeled as 30+).

Broad-spectrum sunscreens contain both UV-B-absorbing and UV-A-absorbing chemicals, the latter including avobenzene and ecamsule (terephthalylidene dicamphor sulfonic acid). These chemicals absorb UVR and transfer the absorbed energy to surrounding cells. In contrast, physical UV blockers (zinc oxide and titanium dioxide) scatter or reflect UVR.

In addition to light absorption, a critical determinant of the sustained photoprotective effect of sunscreens is their water resistance. The FDA monograph has defined strict testing criteria for sunscreens that claim to possess a high degree of water resistance.

Some degree of photoprotection can be achieved by limiting the time of sun exposure during the day. Since a large part of an individual's total lifetime sun exposure may occur by age 18, it is important to educate parents and young children about the hazards of sunlight. Simply eliminating exposure at midday will substantially reduce lifetime UVR exposure.

PHOTOTHERAPY AND PHOTOCHEMOTHERAPY

UVR can be used therapeutically. The administration of UV-B alone or in combination with topically applied agents can induce remissions of many dermatologic diseases, including psoriasis and atopic dermatitis. In particular, narrow-band UV-B treatments (with fluorescent bulbs emitting radiation at ~ 311 nm) have enhanced efficacy over that obtained with broad-band UV-B in the treatment of psoriasis.

Photochemotherapy in which topically applied or systemically administered psoralens are combined with UV-A (PUVA) is effective in treating psoriasis and the early stages of cutaneous T cell lymphoma and vitiligo. Psoralens are tricyclic furocoumarins that, when intercalated into DNA and exposed to UV-A, form adducts with pyrimidine bases and eventually form DNA cross-links. These structural changes are thought to decrease DNA synthesis and to be related to the amelioration of psoriasis. Why PUVA photochemotherapy is effective in cutaneous T cell lymphoma is only partially understood, but it has been shown to induce apoptosis of atypical T lymphocyte populations in the skin. Consequently, direct treatment of circulating atypical lymphocytes by extracorporeal photochemotherapy (photopheresis) has been used in Sézary syndrome as well as in other severe systemic diseases with circulating atypical lymphocytes, such as graft-versus-host disease.

In addition to its effects on DNA, PUVA photochemotherapy stimulates epidermal thickening and melanin synthesis; the latter property, together with its anti-inflammatory effects, provides the rationale for use of PUVA in the depigmenting disease vitiligo. Oral 8-methoxypsoralen and UV-A appear to be most effective in this regard, but as many as 100 treatments extending over 12–18 months may be required for satisfactory repigmentation.

Not surprisingly, the major side effects of long-term UV-B phototherapy and PUVA photochemotherapy mimic those seen in individuals with chronic sun exposure and include skin dryness, actinic keratoses, and an increased risk of skin cancer. Despite these risks, the therapeutic index of these modalities continues to be excellent. It is important to choose the most appropriate phototherapeutic approach for a specific dermatologic disease. For example, narrow-band UV-B has been reported in several studies to be as effective as PUVA photochemotherapy in the treatment of psoriasis but to pose a lower risk of skin cancer development than PUVA.

TABLE 75-5 FDA CATEGORY I MONOGRAPHED SUNSCREEN INGREDIENTS^a

Ingredients	Maximum Concentration, %
<i>p</i> -Aminobenzoic acid (PABA)	15
Avobenzene	3
Cinoxate	3
Dioxybenzone (benzophenone-8)	3
Ecamsule	15
Homosalate	15
Methyl anthranilate	5
Octocrylene	10
Octyl methoxycinnamate	7.5
Octyl salicylate	5
Oxybenzone (benzophenone-3)	6
Padimate O (octyl dimethyl PABA)	8
Phenylbenzimidazole sulfonic acid	4
Sulisobenzene (benzophenone-4)	10
Titanium dioxide	25
Trolamine salicylate	12
Zinc oxide	25

^aFDA, U.S. Food and Drug Administration.

In the practice of medicine, virtually every clinician encounters patients with skin disease. Physicians of all specialties face the daily task of determining the nature and clinical implication of dermatologic disease. In patients with skin disease, the physician must confront the question of whether the cutaneous process is confined to the skin, representing a purely dermatologic event, or whether it is a manifestation of internal disease related to the patient's overall medical condition. Evaluation and accurate diagnosis of skin lesions are particularly critical given the marked rise in both melanoma and nonmelanoma skin cancer. Dermatologic conditions can be classified and categorized in many ways. In this atlas, a selected group of inflammatory skin eruptions and neoplastic conditions are grouped in the following manner: (1) common skin diseases and lesions, (2) nonmelanoma skin cancer, (3) melanoma and benign pigmented lesions, (4) infectious disease and the skin, (5) immunologically mediated skin disease, and (6) skin manifestations of internal disease.

COMMON SKIN DISEASES AND LESIONS

(Figs. 76e-1 to 76e-19) While most of these common inflammatory skin diseases and benign neoplastic and reactive lesions usually present as a predominantly dermatologic process, underlying systemic associations may be found in some settings. Atopic dermatitis is often present in patients with an atopic diathesis, including asthma or sinusitis. Psoriasis ranges from limited patches on the elbows and knees to severe erythrodermic and pustular involvement and associated psoriatic arthritis. Some patients with alopecia areata may have an underlying thyroid abnormality requiring screening. Finally, even acne vulgaris, one of the most common inflammatory dermatoses, can be associated with a systemic process such as polycystic ovarian syndrome.

NONMELANOMA SKIN CANCERS

(Figs. 76e-20 to 76e-27) In fair-skinned ethnic populations, rates of nonmelanoma skin cancer are increasing at an alarming rate. Basal cell carcinoma is the most common cancer in humans and is strongly linked to ultraviolet radiation. Squamous cell carcinoma, including keratoacanthoma, is the second most common skin cancer in most ethnic groups and is also most commonly linked to ultraviolet radiation. Less common cutaneous malignancies include cutaneous T cell lymphoma (mycosis fungoides) and carcinoma and lymphoma metastatic to skin.



FIGURE 76e-1 Acne vulgaris, with inflammatory papules, pustules, and comedones. (Courtesy of Kalman Watsky, MD; with permission.)

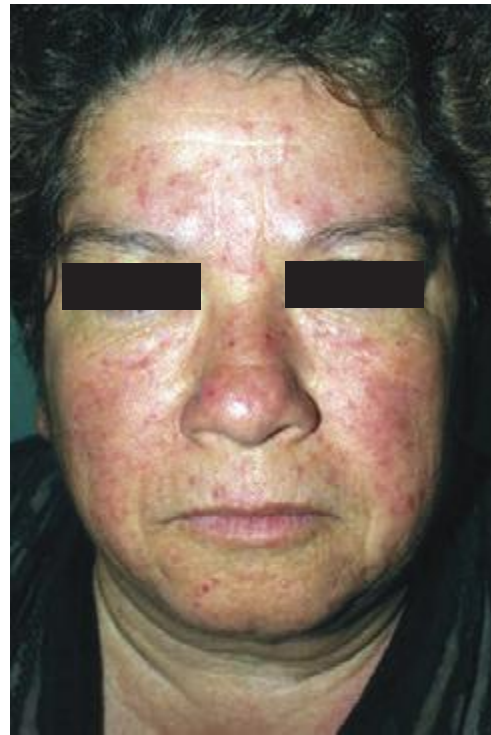


FIGURE 76e-2 Acne rosacea, with prominent facial erythema, telangiectasias, scattered papules, and small pustules. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-3 Psoriasis. **A.** Typical psoriasis is characterized by small and large erythematous plaques with adherent silvery scale. **B.** Acute inflammatory variants of psoriasis may present with widespread superficial pustules.



FIGURE 76e-4 Atopic dermatitis, with hyperpigmentation, lichenification, and scaling in the antecubital fossae. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-7 Stasis dermatitis, with erythematous, scaly, and oozing patches over the lower leg. Several stasis ulcers are also seen in this patient.



FIGURE 76e-5 Dyshidrotic eczema, characterized by deep-seated vesicles and scaling on palms and lateral fingers, is often associated with an atopic diathesis.



A



B

FIGURE 76e-8 Allergic contact dermatitis. **A.** Acute phase, with sharply demarcated, weeping, eczematous plaques in a perioral distribution. **B.** Allergic contact reaction to nickel, chronic phase, with an erythematous, lichenified, weeping plaque on skin chronically exposed to a metal snap. (B: Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-6 Seborrheic dermatitis, with erythema and scale in the nasolabial fold. (Courtesy of Robert A. Swerlick, MD; with permission.)



FIGURE 76e-9 Lichen planus, with multiple flat-topped, violaceous papules and plaques. Nail dystrophy, as seen in this patient's thumb-nail, may also be a feature. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-11 Vitiligo in a typical acral distribution, with striking cutaneous depigmentation as a result of melanocyte loss.



FIGURE 76e-10 Seborrheic keratoses are "stuck on," waxy, verrucous papules and plaques with a variety of colors ranging from light tan to black.



FIGURE 76e-12 Alopecia areata, characterized by a sharply demarcated circular patch of scalp completely devoid of hairs. Preservation of follicular orifices is indicative of nonscarring alopecia. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-13 Pityriasis rosea. Multiple round or oval erythematous patches with fine central scale are distributed along the skin tension lines on the trunk.



A



B

FIGURE 76e-14 **A.** Urticaria, with characteristic discrete and confluent, edematous, erythematous papules and plaques. **B.** Dermatographism. Erythema and whealing developed after firm stroking of the skin. (B: Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-15 **Epidermoid cysts.** Several inflamed and noninflamed firm cystic nodules are seen in this patient. Often a pathulous follicular punctum is observed on the overlying epidermal surface.



FIGURE 76e-16 **Keloids** resulting from ear piercing, with firm exophytic flesh-colored to erythematous nodules of scar tissue.



FIGURE 76e-17 **Cherry hemangiomas**—multiple erythematous to dark-purple papules, usually located on the trunk—are very common and arise in middle-aged to older adults.



FIGURE 76e-18 Frostbite of the hand, with vesiculation surrounded by edema and erythema. (Courtesy of Daniel F. Danzl, MD; with permission.)



FIGURE 76e-19 Frostbite of the foot, with vesiculation surrounded by edema and erythema. (Courtesy of Daniel F. Danzl, MD; with permission.)



FIGURE 76e-20 Kaposi's sarcoma in a patient with AIDS. Patch, plaque, and tumor stages are shown.



FIGURE 76e-21 Non-Hodgkin's lymphoma involving the skin, with typical violaceous, "plum-colored" nodules. (Courtesy of Jean Bologna, MD; with permission.)



FIGURE 76e-22 Basal cell carcinoma, with central ulceration and a pearly, rolled, telangiectatic tumor border.

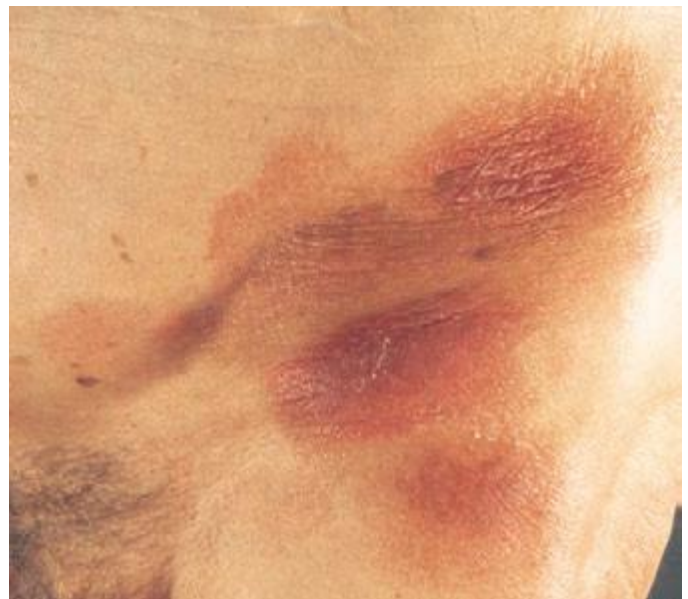


FIGURE 76e-23 Mycosis fungoides is a cutaneous T cell lymphoma. Plaque-stage lesions are seen in this patient.



FIGURE 76e-24 Metastatic carcinoma to the skin is characterized by inflammatory, often ulcerated dermal nodules.



FIGURE 76e-25 Keratoacanthoma is a low-grade squamous cell carcinoma that presents as an exophytic nodule with central keratinous debris.

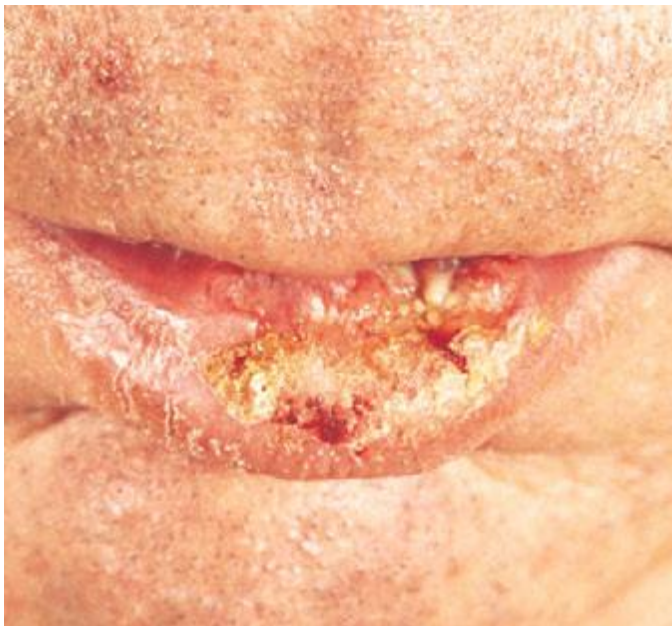


FIGURE 76e-26 Squamous cell carcinoma is seen here as a hyperkeratotic, crusted, and somewhat eroded plaque on the lower lip. Sun-exposed skin of the head, neck, hands, and arms are other typical sites of involvement.



FIGURE 76e-27 Actinic keratoses consist of hyperkeratotic erythematous papules and patches on sun-exposed skin. They arise in middle-aged to older adults and have some potential for malignant transformation. (Courtesy of Robert Swerlick, MD; with permission.)

MELANOMA AND BENIGN PIGMENTED LESIONS

(Figs. 76e-28 to 76e-33) As the prognosis of melanoma is related primarily to the microscopic depth of invasion, and as early detection with surgical treatment can be curative in a high percentage of patients, it is essential that all clinicians acquire some facility in evaluating pigmented lesions. Three clinicopathologic subtypes of melanoma—superficial spreading, lentigo maligna, and acral lentiginous melanoma—typically display features noted in the “ABCD rule”: *asymmetry* (one half of the lesion varies from the other half); *border irregularity* (the circumferential border exhibits an irregular, sometimes jagged appearance); *color* (there is uneven coloration and tone to the pigmented lesion, with various shades of brown, black, red, and white in different areas); and *diameter* (the diameter is typically >6 mm). The more uncommon subtype, nodular melanoma, may not manifest all these features but rather may present as a more symmetric, evenly pigmented, or amelanotic lesion. Dysplastic (atypical) melanocytic nevi may occur as solitary or multiple lesions as well as in the setting of familial melanoma. These nevi display some degree of asymmetry, border irregularity, and color variation. Ordinary nevi may be acquired or congenital and are quite common.



FIGURE 76e-28 Nevi are benign proliferations of nevomelanocytes characterized by regularly shaped hyperpigmented macules or papules of a uniform color.

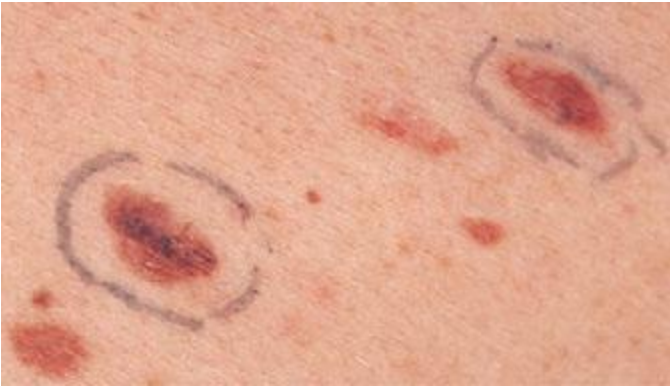


FIGURE 76e-29 Dysplastic nevi are irregularly pigmented and shaped nevocytic lesions that may be associated with familial melanoma.

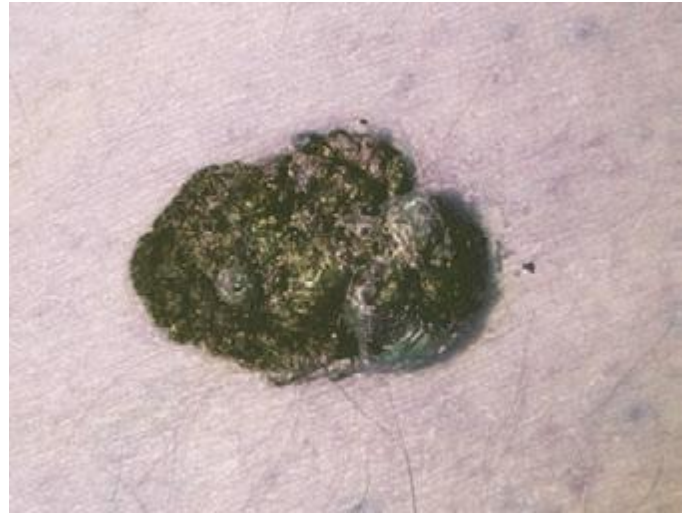


FIGURE 76e-32 Nodular melanoma most commonly manifests as a rapidly growing, often ulcerated or crusted black nodule. (Courtesy of S. Wright Caughman, MD; with permission.)



FIGURE 76e-30 Superficial spreading melanoma, the most common type of malignant melanoma, is characterized by color variegation (black, blue, brown, pink, and white) and irregular borders.

INFECTIOUS DISEASE AND THE SKIN

(Figs. 76e-34 to 76e-58) One of the roles of the skin is to function as a barrier from the outside world. In this capacity, exposure to infectious agents occurs, and bacterial, viral, fungal, and parasitic infections may result. In addition, the skin may be secondarily involved and provides diagnostic clues to systemic infections such as meningococemia, Rocky Mountain spotted fever, Lyme disease, and septic emboli. Most sexually transmitted bacterial and viral diseases exhibit cutaneous involvement; examples include primary and secondary syphilis, chancroid, genital herpes simplex, and condyloma acuminatum.

IMMUNOLOGICALLY MEDIATED SKIN DISEASE

(Figs. 76e-59 to 76e-70) Immunologically mediated skin disease may be largely localized to skin and mucous membranes and manifest with blisters and erosions such as pemphigus, pemphigoid, and dermatitis herpetiformis. In diseases such as systemic lupus erythematosus, dermatomyositis, and vasculitis, skin manifestations are often only one element of a widespread process.

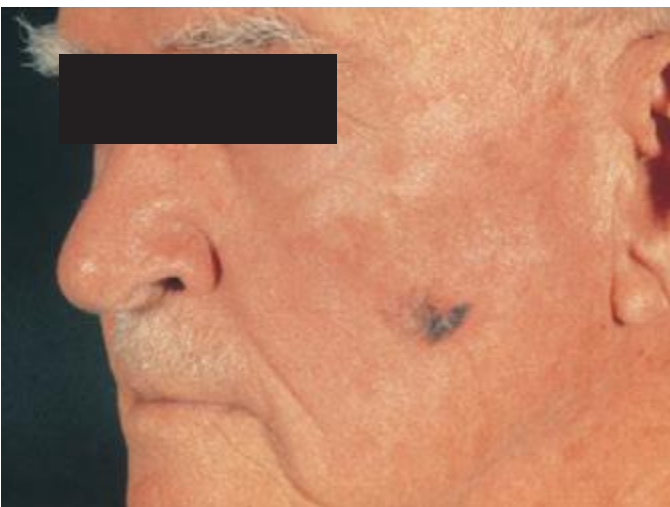


FIGURE 76e-31 Lentigo maligna melanoma occurs on sun-exposed skin as a large, hyperpigmented macule or plaque with irregular borders and variable pigmentation. (Courtesy of Alvin Solomon, MD; with permission.)



FIGURE 76e-33 Acral lentiginous melanoma is more common among blacks, Asians, and Hispanics and occurs as an enlarging hyperpigmented macule or plaque on the palms or soles. Lateral pigment diffusion is present.



FIGURE 76e-34 Erysipelas is a streptococcal infection of the superficial dermis and consists of well-demarcated, erythematous, edematous, warm plaques.



FIGURE 76e-35 Varicella, with numerous lesions in various stages of evolution: vesicles on an erythematous base, umbilicated vesicles, and crusts. (Courtesy of Robert Hartman, MD; with permission.)



FIGURE 76e-36 Herpes zoster is seen in this HIV-infected patient as hemorrhagic vesicles and pustules on an erythematous base in a dermatomal distribution. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-37 Impetigo contagiosa is a superficial streptococcal or *Staphylococcus aureus* infection consisting of honey-colored crusts and erythematous weeping erosions. Bullous lesions are occasionally seen.



FIGURE 76e-38 Tender vesicles and erosions in the mouth of a patient with hand-foot-and-mouth disease. (Courtesy of Stephen D. Gellis, MD; with permission.)



FIGURE 76e-39 Lacy reticular rash of erythema infectiosum (fifth disease).

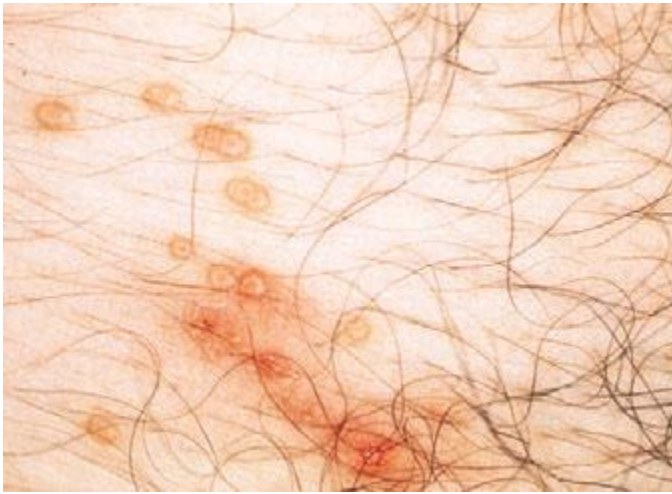


FIGURE 76e-40 *Molluscum contagiosum* is a cutaneous poxvirus infection characterized by multiple umbilicated flesh-colored or hypopigmented papules. (Courtesy of Yale Resident's Slide Collection; with permission.)



FIGURE 76e-43 Rocky Mountain spotted fever, with pinpoint petechial lesions on the palm and volar aspect of the wrist. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-41 Oral hairy leukoplakia often presents as white plaques on the lateral tongue and is associated with Epstein-Barr virus infection. (From K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005. www.accessmedicine.com.)



FIGURE 76e-44 Erythema migrans, the early cutaneous manifestation of Lyme disease, is characterized by erythematous annular patches, often with a central erythematous papule at the tick-bite site. (Courtesy of Yale Resident's Slide Collection; with permission.)

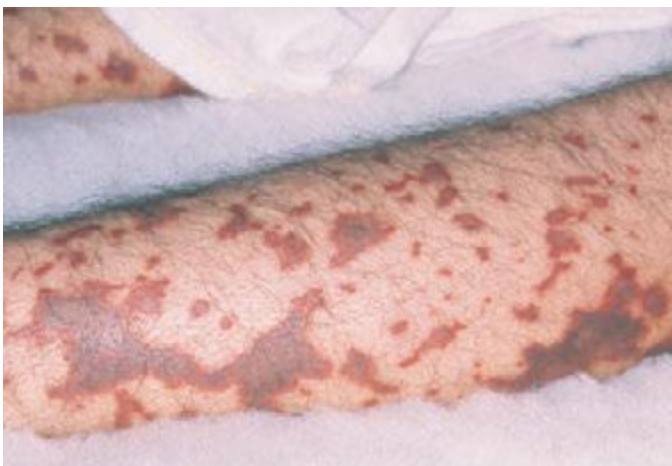


FIGURE 76e-42 Fulminant meningococemia, with extensive angular purpuric patches. (Courtesy of Stephen D. Gellis, MD; with permission.)



FIGURE 76e-45 Primary syphilis, with a firm, nontender chancre. (Courtesy of Gregory Cox, MD; with permission.)



FIGURE 76e-46 Secondary syphilis commonly affects the palms and soles, with scaling, firm, red-brown papules. (Courtesy of Alvin Solomon, MD; with permission.)



FIGURE 76e-47 Condylomata lata are moist, somewhat verrucous inter-triginous plaques seen in secondary syphilis. (Courtesy of Yale Resident's Slide Collection; with permission.)



FIGURE 76e-48 Secondary syphilis, with the characteristic papulo-squamous truncal eruption.



A



B

FIGURE 76e-49 **A.** Tinea corporis is a superficial fungal infection, seen here as an erythematous annular scaly plaque with central clearing. **B.** A common presentation of chronic dermatophyte infection involves the feet (tinea pedis), hands (tinea manum), and nails (tinea unguium).



FIGURE 76e-50 Scabies, with typical scaling erythematous papules and few linear burrows.



FIGURE 76e-51 Skin lesions caused by *Chironex fleckeri* sting. (Courtesy of V. Pranava Murthy, MD; with permission.)



FIGURE 76e-53 Condylomata acuminata are lesions induced by human papillomavirus and in this patient are seen as multiple verrucous papules coalescing into plaques. (Courtesy of S. Wright Caughman, MD; with permission.)



FIGURE 76e-52 Chancroid, with characteristic penile ulcers and associated left inguinal adenitis (bubo).



FIGURE 76e-54 A patient with features of polar lepromatous leprosy: multiple nodular skin lesions, particularly of the forehead, and loss of eyebrows. (Courtesy of Robert Gelber, MD; with permission.)



FIGURE 76e-55 Skin lesions of neutropenic patients. **A.** Hemorrhagic papules on the foot of a patient undergoing treatment for multiple myeloma. Biopsy and culture demonstrated *Aspergillus* species. **B.** Eroded nodule on the hard palate of a patient undergoing chemotherapy. Biopsy and culture demonstrated *Mucor* species. **C.** Ecthyma gangrenosum in a neutropenic patient with *Pseudomonas aeruginosa* bacteremia.



FIGURE 76e-56 Septic emboli, with hemorrhage and infarction due to acute *Staphylococcus aureus* endocarditis. (Courtesy of L. Baden, MD; with permission.)

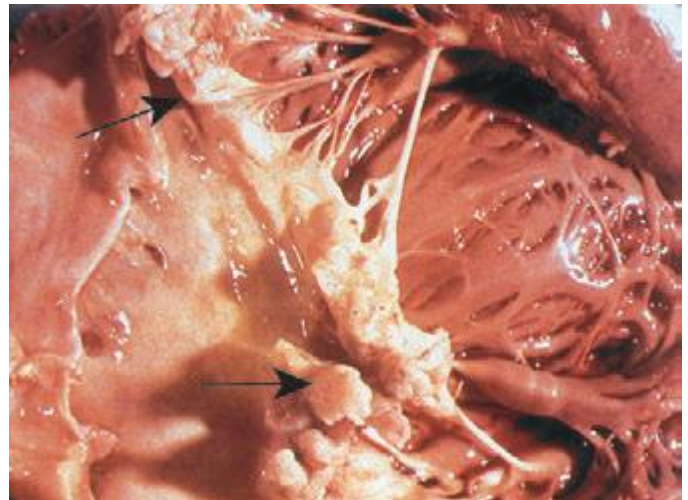


FIGURE 76e-57 Vegetations (arrows) due to viridans streptococcal endocarditis involving the mitral valve. (Courtesy of AW Karchmer, MD; with permission.)

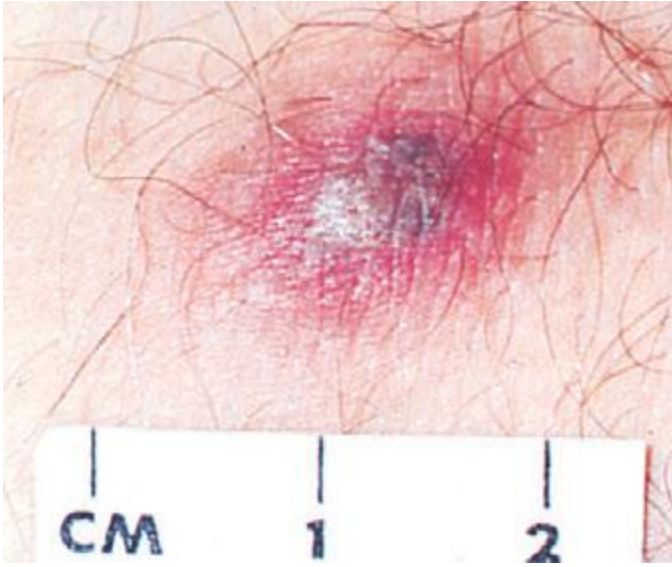


FIGURE 76e-58 Disseminated gonococemia in the skin is seen as hemorrhagic papules and pustules with purpuric centers in an acral distribution. (Courtesy of Daniel M. Musher, MD; with permission.)



FIGURE 76e-60 Discoid lupus erythematosus. Atrophic, depigmented plaques and patches surrounded by hyperpigmentation and erythema in association with scarring and alopecia are characteristic of this cutaneous form of lupus.

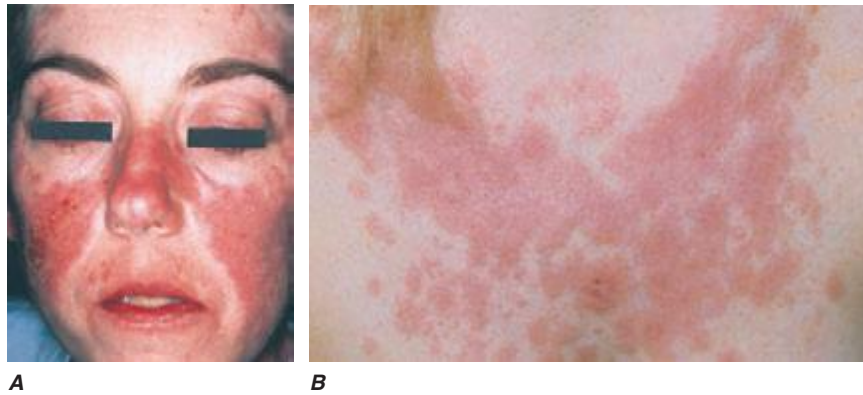


FIGURE 76e-59 Lupus erythematosus. **A.** Systemic lupus erythematosus, with prominent, scaly malar erythema. Involvement of other sun-exposed sites is also common. **B.** Acute lupus erythematosus on the upper chest, with brightly erythematous and slightly edematous coalescence of papules and plaques. (B: Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-61 Dermatomyositis. Periorbital violaceous erythema characterizes the classic heliotrope rash. (Courtesy of James Krell, MD; with permission.)



FIGURE 76e-62 Scleroderma characterized by typical expressionless, mask-like facies.



FIGURE 76e-65 Erythema multiforme is characterized by multiple erythematous plaques with a target or iris morphology and usually represents a hypersensitivity reaction to drugs or infections (especially herpes simplex virus). (Courtesy of Yale Resident's Slide Collection; with permission.)



FIGURE 76e-63 Scleroderma, with acral sclerosis and focal digital ulcers.



FIGURE 76e-66 Dermatitis herpetiformis, manifested by pruritic, grouped vesicles in a typical location. The vesicles are often excoriated and may also occur on the knees, buttocks, elbows, and posterior scalp.



FIGURE 76e-64 Dermatomyositis often involves the hands as erythematous flat-topped papules over the knuckles (Gottron's sign) and periungual telangiectasias.



A



B

FIGURE 76e-67 Pemphigus vulgaris. **A.** Eroded bullae on the back. **B.** The oral mucosa is almost invariably involved, sometimes with erosions on the gingiva, buccal mucosa, palate, posterior pharynx, or tongue. (B: Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-68 Erythema nodosum is a panniculitis characterized by tender deep-seated nodules and plaques, usually located on the lower extremities. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-69 Vasculitis. Palpable purpuric papules on the lower legs are seen in this patient with cutaneous small-vessel vasculitis. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-70 Bullous pemphigoid, with tense vesicles and bullae on an erythematous, urticarial base. (Courtesy of Yale Resident's Slide Collection; with permission.)

SKIN MANIFESTATIONS OF INTERNAL DISEASE

(Figs. 76e-71 to 76e-78) While many systemic diseases also have cutaneous manifestations, there are well-recognized dermatologic markers of internal disease, some of which are shown in this section. Many of these dermatologic markers may precede, accompany, or follow diagnosis of systemic disease. Acanthosis nigricans is a prototypical dermatologic process that often occurs in association with underlying systemic abnormalities, most commonly obesity and insulin resistance. It may also be associated with other endocrine disorders and several rare genetic syndromes. Malignant acanthosis nigricans may occur in association with several malignancies, especially adenocarcinoma of the gastrointestinal tract, lung, and breast. Other markers of internal disease in this section include pretibial myxedema, which is associated with thyroid disease, and Sweet syndrome, which may be associated with hematologic malignancies, solid tumors, infections, or inflammatory bowel disease. The skin is also involved in many systemic inflammatory diseases such as sarcoidosis, rheumatoid arthritis, and lupus erythematosus.



FIGURE 76e-71 Acanthosis nigricans, with typical hyperpigmented plaques on a velvet-like, verrucous surface on the neck.



FIGURE 76e-74 Bilateral rheumatoid nodules of the upper extremities. (Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-72 Pretibial myxedema manifesting as waxy, infiltrated plaques in a patient with Graves' disease.



FIGURE 76e-75 Neurofibromatosis, with numerous flesh-colored cutaneous neurofibromas.



FIGURE 76e-73 Erythematous, indurated plaque of Sweet syndrome, with a pseudovesicular border. (Courtesy of Robert Swerlick, MD, with permission.)



FIGURE 76e-76 Coumarin necrosis. Shown is cutaneous and subcutaneous necrosis of a breast. Other fatty areas, such as buttocks and thighs, are also common sites of involvement. (Courtesy of Kim Yancey, MD; with permission.)



A



B

FIGURE 76e-77 Sarcoid. **A.** Infiltrated papules and plaques of variable color are seen in a typical paranasal and periorbital location. **B.** Infiltrated, hyperpigmented, and slightly erythematous coalescent papules and plaques on the upper arm. (B: Courtesy of Robert Swerlick, MD; with permission.)



FIGURE 76e-78 Pyoderma gangrenosum on the dorsal aspect of both hands. Multiple necrotic ulcers are surrounded by a violaceous and undermined border. (Courtesy of Robert Swerlick, MD; with permission.)

77 Anemia and Polycythemia

John W. Adamson, Dan L. Longo

HEMATOPOIESIS AND THE PHYSIOLOGIC BASIS OF RED CELL PRODUCTION

Hematopoiesis is the process by which the formed elements of blood are produced. The process is regulated through a series of steps beginning with the hematopoietic stem cell. Stem cells are capable of producing red cells, all classes of granulocytes, monocytes, platelets, and the cells of the immune system. The precise molecular mechanism—either intrinsic to the stem cell itself or through the action of extrinsic factors—by which the stem cell becomes committed to a given lineage is not fully defined. However, experiments in mice suggest that erythroid cells come from a common erythroid/megakaryocyte progenitor that does not develop in the absence of expression of the GATA-1 and FOG-1 (friend of GATA-1) transcription factors (Chap. 89e). Following lineage commitment, hematopoietic progenitor and precursor cells come increasingly under the regulatory influence of growth factors and hormones. For red cell production, erythropoietin (EPO) is the primary regulatory hormone. EPO is required for the maintenance of committed erythroid progenitor cells that, in the absence of the hormone, undergo programmed cell death (*apoptosis*). The regulated process of red cell production is *erythropoiesis*, and its key elements are illustrated in Fig. 77-1.

In the bone marrow, the first morphologically recognizable erythroid precursor is the pronormoblast. This cell can undergo four to five cell divisions, which result in the production of 16–32 mature red cells. With increased EPO production, or the administration of EPO as a drug, early progenitor cell numbers are amplified and, in turn, give rise to increased numbers of erythrocytes. The regulation of EPO production itself is linked to tissue oxygenation.

In mammals, O₂ is transported to tissues bound to the hemoglobin contained within circulating red cells. The mature red cell is 8 μm in diameter, anucleate, discoid in shape, and extremely pliable in order to traverse the microcirculation successfully; its membrane integrity is maintained by the intracellular generation of ATP. Normal red cell production results in the daily replacement of 0.8–1% of all circulating red cells in the body, since the average red cell lives 100–120 days. The organ responsible for red cell production is called the *erythron*. The erythron is a dynamic organ made up of a rapidly proliferating pool of marrow erythroid precursor cells and a large mass of mature circulating

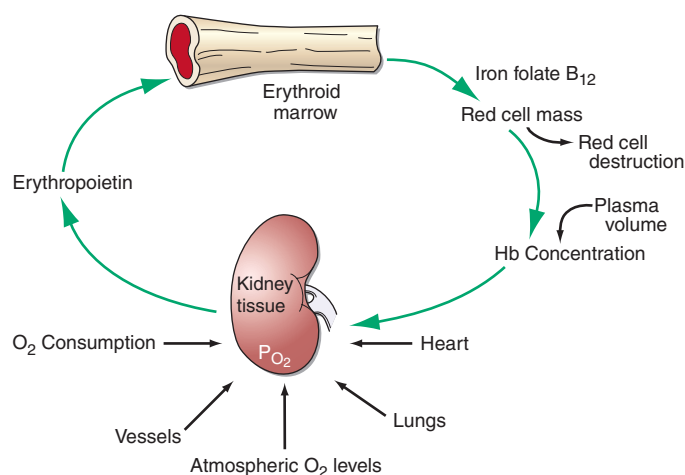


FIGURE 77-1 The physiologic regulation of red cell production by tissue oxygen tension. Hb, hemoglobin.

red blood cells. The size of the red cell mass reflects the balance of red cell production and destruction. The physiologic basis of red cell production and destruction provides an understanding of the mechanisms that can lead to anemia.

The physiologic regulator of red cell production, the glycoprotein hormone EPO, is produced and released by peritubular capillary lining cells within the kidney. These cells are highly specialized epithelial-like cells. A small amount of EPO is produced by hepatocytes. The fundamental stimulus for EPO production is the availability of O₂ for tissue metabolic needs. Key to EPO gene regulation is hypoxia-inducible factor (HIF)-1α. In the presence of O₂, HIF-1α is hydroxylated at a key proline, allowing HIF-1α to be ubiquitinated and degraded via the proteasome pathway. If O₂ becomes limiting, this critical hydroxylation step does not occur, allowing HIF-1α to partner with other proteins, translocate to the nucleus, and upregulate the expression of the EPO gene, among others.

Impaired O₂ delivery to the kidney can result from a decreased red cell mass (*anemia*), impaired O₂ loading of the hemoglobin molecule or a high O₂ affinity mutant hemoglobin (*hypoxemia*), or, rarely, impaired blood flow to the kidney (renal artery stenosis). EPO governs the day-to-day production of red cells, and ambient levels of the hormone can be measured in the plasma by sensitive immunoassays—the normal level being 10–25 U/L. When the hemoglobin concentration falls below 100–120 g/L (10–12 g/dL), plasma EPO levels increase in proportion to the severity of the anemia (Fig. 77-2). In circulation, EPO has a half-clearance time of 6–9 h. EPO acts by binding to specific receptors on the surface of marrow erythroid precursors, inducing them to proliferate and to mature. With EPO stimulation, red cell production can increase four- to fivefold within a 1- to 2-week period, but only in the presence of adequate nutrients, especially iron. The functional capacity of the erythron, therefore, requires normal renal production of EPO, a functioning erythroid marrow, and an adequate supply of substrates for hemoglobin synthesis. A defect in any of these key components can lead to anemia. Generally, anemia is recognized in the laboratory when a patient’s hemoglobin level or hematocrit is reduced below an expected value (the normal range). The likelihood and severity of anemia are defined based on the deviation of the patient’s hemoglobin/hematocrit from values expected for age- and sex-matched normal subjects. The hemoglobin concentration in adults has a Gaussian distribution. The

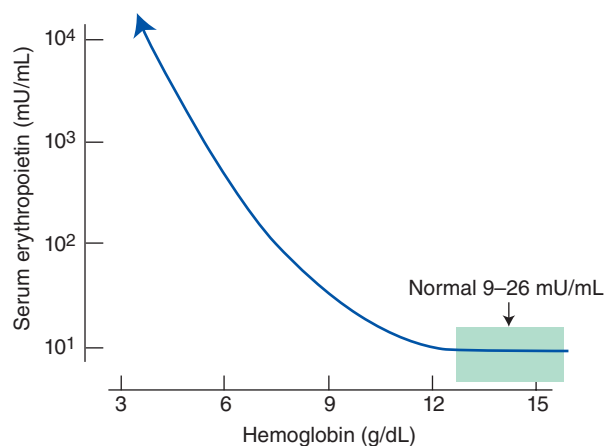


FIGURE 77-2 Erythropoietin (EPO) levels in response to anemia. When the hemoglobin level falls to 120 g/L (12 g/dL), plasma EPO levels increase logarithmically. In the presence of chronic kidney disease or chronic inflammation, EPO levels are typically lower than expected for the degree of anemia. As individuals age, the level of EPO needed to sustain normal hemoglobin levels appears to increase. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

mean hematocrit value for adult males is 47% (standard deviation, $\pm 7\%$) and that for adult females is 42% ($\pm 5\%$). Any single hematocrit or hemoglobin value carries with it a likelihood of associated anemia. Thus, a hematocrit of $<39\%$ in an adult male or $<35\%$ in an adult female has only about a 25% chance of being normal. Hematocrit levels are less useful than hemoglobin levels in assessing anemia because they are calculated rather than measured directly. Suspected low hemoglobin or hematocrit values are more easily interpreted if previous values for the same patient are known for comparison. The World Health Organization (WHO) defines anemia as a hemoglobin level <130 g/L (13 g/dL) in men and <120 g/L (12 g/dL) in women.

The critical elements of erythropoiesis—EPO production, iron availability, the proliferative capacity of the bone marrow, and effective maturation of red cell precursors—are used for the initial classification of anemia (see below).

ANEMIA

CLINICAL PRESENTATION OF ANEMIA

Signs and Symptoms Anemia is most often recognized by abnormal screening laboratory tests. Patients less commonly present with advanced anemia and its attendant signs and symptoms. Acute anemia is due to blood loss or hemolysis. If blood loss is mild, enhanced O_2 delivery is achieved through changes in the O_2 -hemoglobin dissociation curve mediated by a decreased pH or increased CO_2 (*Bohr effect*). With acute blood loss, hypovolemia dominates the clinical picture, and the hematocrit and hemoglobin levels do not reflect the volume of blood lost. Signs of vascular instability appear with acute losses of 10–15% of the total blood volume. In such patients, the issue is not anemia but hypotension and decreased organ perfusion. When $>30\%$ of the blood volume is lost suddenly, patients are unable to compensate with the usual mechanisms of vascular contraction and changes in regional blood flow. The patient prefers to remain supine and will show postural hypotension and tachycardia. If the volume of blood lost is $>40\%$ (i.e., >2 L in the average-sized adult), signs of hypovolemic shock including confusion, dyspnea, diaphoresis, hypotension, and tachycardia appear ([Chap. 129](#)). Such patients have significant deficits in vital organ perfusion and require immediate volume replacement.

With acute hemolysis, the signs and symptoms depend on the mechanism that leads to red cell destruction. Intravascular hemolysis with release of free hemoglobin may be associated with acute back pain, free hemoglobin in the plasma and urine, and renal failure. Symptoms associated with more chronic or progressive anemia depend on the age of the patient and the adequacy of blood supply to critical organs. Symptoms associated with moderate anemia include fatigue, loss of stamina, breathlessness, and tachycardia (particularly with physical exertion). However, because of the intrinsic compensatory mechanisms that govern the O_2 -hemoglobin dissociation curve, the gradual onset of anemia—particularly in young patients—may not be associated with signs or symptoms until the anemia is severe (hemoglobin <70 – 80 g/L [7 – 8 g/dL]). When anemia develops over a period of days or weeks, the total blood volume is normal to slightly increased, and changes in cardiac output and regional blood flow help compensate for the overall loss in O_2 -carrying capacity. Changes in the position of the O_2 -hemoglobin dissociation curve account for some of the compensatory response to anemia. With chronic anemia, intracellular levels of 2,3-bisphosphoglycerate rise, shifting the dissociation curve to the right and facilitating O_2 unloading. This compensatory mechanism can only maintain normal tissue O_2 delivery in the face of a 20–30 g/L (2–3 g/dL) deficit in hemoglobin concentration. Finally, further protection of O_2 delivery to vital organs is achieved by the shunting of blood away from organs that are relatively rich in blood supply, particularly the kidney, gut, and skin.

Certain disorders are commonly associated with anemia. Chronic inflammatory states (e.g., infection, rheumatoid arthritis, cancer) are associated with mild to moderate anemia, whereas lymphoproliferative disorders, such as chronic lymphocytic leukemia and certain other B cell neoplasms, may be associated with autoimmune hemolysis.

APPROACH TO THE PATIENT:

Anemia

The evaluation of the patient with anemia requires a careful history and physical examination. Nutritional history related to drugs or alcohol intake and family history of anemia should always be assessed. Certain geographic backgrounds and ethnic origins are associated with an increased likelihood of an inherited disorder of the hemoglobin molecule or intermediary metabolism. Glucose-6-phosphate dehydrogenase (G6PD) deficiency and certain hemoglobinopathies are seen more commonly in those of Middle Eastern or African origin, including African Americans who have a high frequency of G6PD deficiency. Other information that may be useful includes exposure to certain toxic agents or drugs and symptoms related to other disorders commonly associated with anemia. These include symptoms and signs such as bleeding, fatigue, malaise, fever, weight loss, night sweats, and other systemic symptoms. Clues to the mechanisms of anemia may be provided on physical examination by findings of infection, blood in the stool, lymphadenopathy, splenomegaly, or petechiae. Splenomegaly and lymphadenopathy suggest an underlying lymphoproliferative disease, whereas petechiae suggest platelet dysfunction. Past laboratory measurements are helpful to determine a time of onset.

In the anemic patient, physical examination may demonstrate a forceful heartbeat, strong peripheral pulses, and a systolic “flow” murmur. The skin and mucous membranes may be pale if the hemoglobin is <80 – 100 g/L (8–10 g/dL). This part of the physical examination should focus on areas where vessels are close to the surface such as the mucous membranes, nail beds, and palmar creases. If the palmar creases are lighter in color than the surrounding skin when the hand is hyperextended, the hemoglobin level is usually <80 g/L (8 g/dL).

LABORATORY EVALUATION

[Table 77-1](#) lists the tests used in the initial workup of anemia. A routine complete blood count (CBC) is required as part of the evaluation and includes the hemoglobin, hematocrit, and red cell indices: the mean cell volume (MCV) in femtoliters, mean cell hemoglobin (MCH) in picograms per cell, and mean concentration of hemoglobin per volume of red cells (MCHC) in grams per liter (non-SI: grams per deciliter). The red cell indices are calculated as shown in [Table 77-2](#), and the normal variations in the hemoglobin and hematocrit with age are shown in [Table 77-3](#). A number of physiologic factors affect the CBC, including age, sex, pregnancy, smoking, and altitude. High-normal hemoglobin values may be seen in men and women who live at altitude or smoke heavily. Hemoglobin elevations due to smoking reflect normal compensation due to the displacement of O_2 by CO in hemoglobin binding. Other important information is provided by the reticulocyte count and measurements of iron supply including *serum iron*, *total iron-binding capacity* (TIBC; an indirect measure of serum transferrin), and *serum ferritin*. Marked alterations in the red cell indices usually reflect disorders of maturation or iron deficiency. A careful evaluation of the peripheral blood smear is important, and clinical laboratories often provide a description of both the red and white cells, a white cell differential count, and the platelet count. In patients with severe anemia and abnormalities in red blood cell morphology and/or low reticulocyte counts, a bone marrow aspirate or biopsy can assist in the diagnosis. Other tests of value in the diagnosis of specific anemias are discussed in chapters on specific disease states.

The components of the CBC also help in the classification of anemia. *Microcytosis* is reflected by a lower than normal MCV (<80), whereas high values (>100) reflect *macrocytosis*. The MCH and MCHC reflect defects in hemoglobin synthesis (*hypochromia*). Automated cell counters describe the red cell volume distribution width (RDW). The MCV (representing the peak of the distribution curve) is insensitive to the appearance of small populations of macrocytes or microcytes. An experienced laboratory technician

TABLE 77-1 LABORATORY TESTS IN ANEMIA DIAGNOSIS

- I. Complete blood count (CBC)
 - A. Red blood cell count
 1. Hemoglobin
 2. Hematocrit
 3. Reticulocyte count
 - B. Red blood cell indices
 1. Mean cell volume (MCV)
 2. Mean cell hemoglobin (MCH)
 3. Mean cell hemoglobin concentration (MCHC)
 4. Red cell distribution width (RDW)
 - C. White blood cell count
 1. Cell differential
 2. Nuclear segmentation of neutrophils
 - D. Platelet count
 - E. Cell morphology
 1. Cell size
 2. Hemoglobin content
 3. Anisocytosis
 4. Poikilocytosis
 5. Polychromasia
- II. Iron supply studies
 - A. Serum iron
 - B. Total iron-binding capacity
 - C. Serum ferritin
- III. Marrow examination
 - A. Aspirate
 1. M/E ratio^a
 2. Cell morphology
 3. Iron stain
 - B. Biopsy
 1. Cellularity
 2. Morphology

^aM/E ratio, ratio of myeloid to erythroid precursors.

TABLE 77-2 RED BLOOD CELL INDICES

Index	Normal Value
Mean cell volume (MCV) = (hematocrit × 10)/(red cell count × 10 ⁶)	90 ± 8 fL
Mean cell hemoglobin (MCH) = (hemoglobin × 10)/(red cell count × 10 ⁶)	30 ± 3 pg
Mean cell hemoglobin concentration = (hemoglobin × 10)/hematocrit, or MCH/MCV	33 ± 2%

TABLE 77-3 CHANGES IN NORMAL HEMOGLOBIN/HEMATOCRIT VALUES WITH AGE, SEX, AND PREGNANCY

Age/Sex	Hemoglobin, g/dL	Hematocrit, %
At birth	17	52
Childhood	12	36
Adolescence	13	40
Adult man	16 (±2)	47 (±6)
Adult woman (menstruating)	13 (±2)	40 (±6)
Adult woman (postmenopausal)	14 (±2)	42 (±6)
During pregnancy	12 (±2)	37 (±6)

Source: From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.

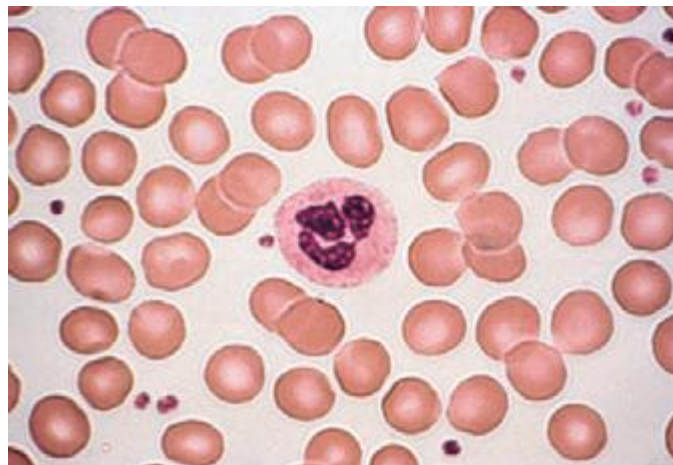


FIGURE 77-3 Normal blood smear (Wright stain). High-power field showing normal red cells, a neutrophil, and a few platelets. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

will be able to identify minor populations of large or small cells or hypochromic cells before the red cell indices change.

Peripheral Blood Smear The peripheral blood smear provides important information about defects in red cell production (**Chap. 81e**). As a complement to the red cell indices, the blood smear also reveals variations in cell size (*anisocytosis*) and shape (*poikilocytosis*). The degree of anisocytosis usually correlates with increases in the RDW or the range of cell sizes. Poikilocytosis suggests a defect in the maturation of red cell precursors in the bone marrow or fragmentation of circulating red cells. The blood smear may also reveal *polychromasia*—red cells that are slightly larger than normal and grayish blue in color on the Wright-Giemsa stain. These cells are reticulocytes that have been prematurely released from the bone marrow, and their color represents residual amounts of ribosomal RNA. These cells appear in circulation in response to EPO stimulation or to architectural damage of the bone marrow (fibrosis, infiltration of the marrow by malignant cells, etc.) that results in their disordered release from the marrow. The appearance of nucleated red cells, Howell-Jolly bodies, target cells, sickle cells, and others may provide clues to specific disorders (**Figs. 77-3 to 77-11**).

Reticulocyte Count An accurate reticulocyte count is key to the initial classification of anemia. Reticulocytes are red cells that

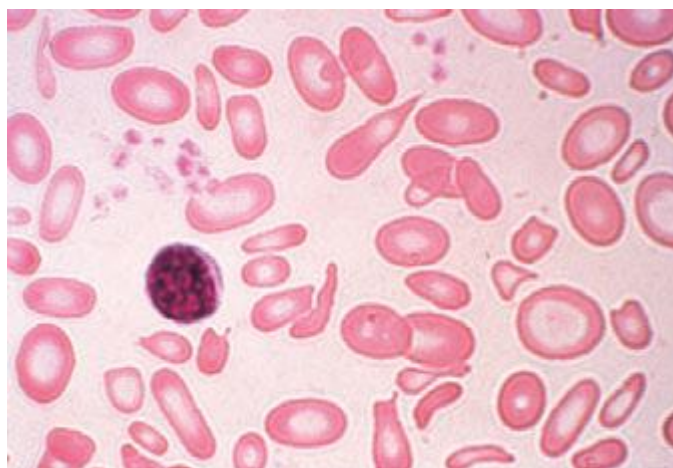


FIGURE 77-4 Severe iron-deficiency anemia. Microcytic and hypochromic red cells smaller than the nucleus of a lymphocyte associated with marked variation in size (anisocytosis) and shape (poikilocytosis). (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

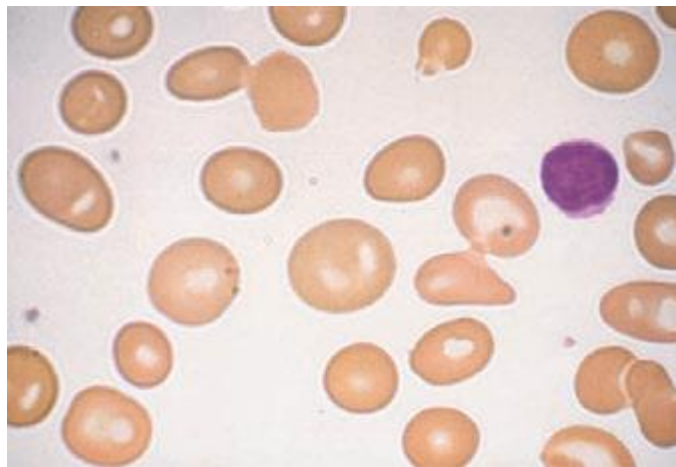


FIGURE 77-5 Macrocytosis. Red cells are larger than a small lymphocyte and well hemoglobinized. Often macrocytes are oval shaped (macro-ovalocytes).

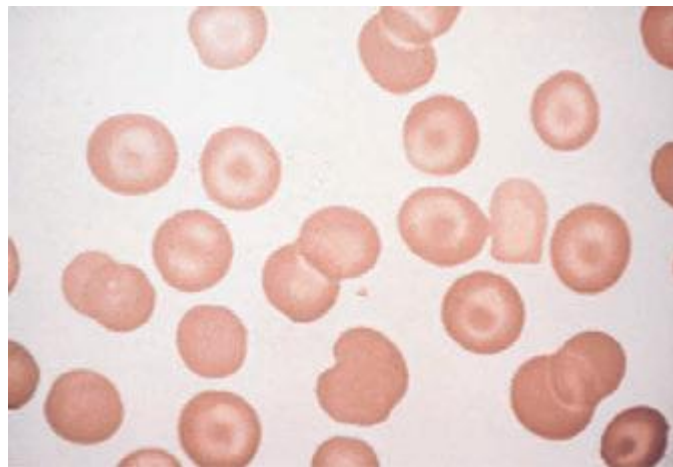


FIGURE 77-8 Target cells. Target cells have a bull's-eye appearance and are seen in thalassemia and in liver disease. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

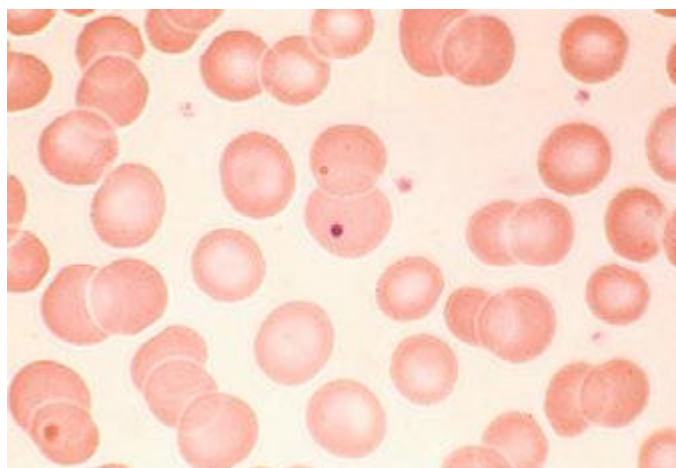


FIGURE 77-6 Howell-Jolly bodies. In the absence of a functional spleen, nuclear remnants are not culled from the red cells and remain as small homogeneously staining blue inclusions on Wright stain. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

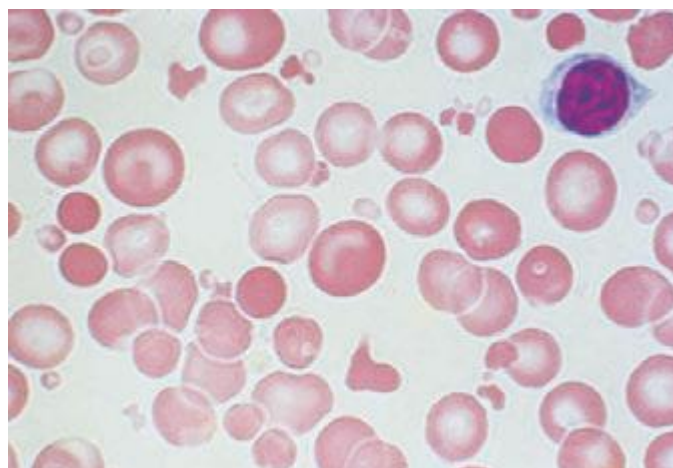


FIGURE 77-9 Red cell fragmentation. Red cells may become fragmented in the presence of foreign bodies in the circulation, such as mechanical heart valves, or in the setting of thermal injury. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

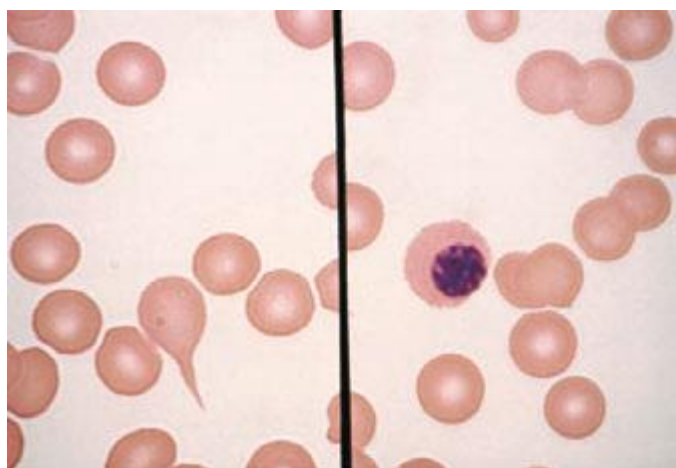


FIGURE 77-7 Red cell changes in myelofibrosis. The left panel shows a teardrop-shaped cell. The right panel shows a nucleated red cell. These forms can be seen in myelofibrosis.

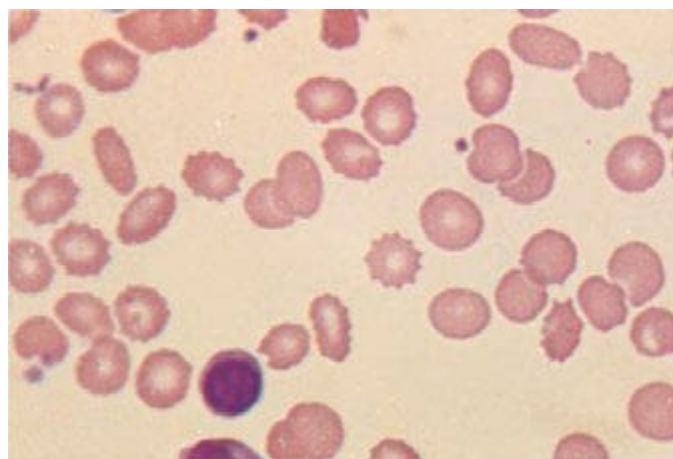


FIGURE 77-10 Uremia. The red cells in uremia may acquire numerous regularly spaced, small, spiny projections. Such cells, called burr cells or echinocytes, are readily distinguishable from irregularly spiculated acanthocytes shown in Fig. 77-11.

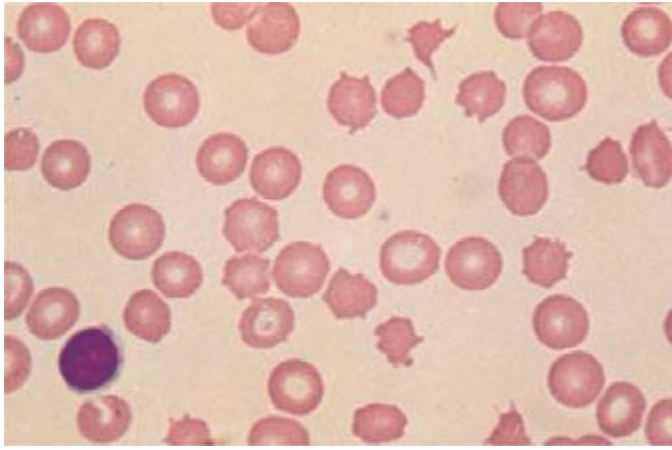


FIGURE 77-11 Spur cells. Spur cells are recognized as distorted red cells containing several irregularly distributed thornlike projections. Cells with this morphologic abnormality are also called acanthocytes. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

have been recently released from the bone marrow. They are identified by staining with a supravital dye that precipitates the ribosomal RNA (Fig. 77-12). These precipitates appear as blue or black punctate spots and can be counted manually or, currently, by fluorescent emission of dyes that bind to RNA. This residual RNA is metabolized over the first 24–36 h of the reticulocyte’s life span in circulation. Normally, the reticulocyte count ranges from 1 to 2% and reflects the daily replacement of 0.8–1.0% of the circulating red cell population. A corrected reticulocyte count provides a reliable measure of effective red cell production.

In the initial classification of anemia, the patient’s reticulocyte count is compared with the expected reticulocyte response. In general, if the EPO and erythroid marrow responses to moderate anemia [hemoglobin <100 g/L (10 g/dL)] are intact, the red cell production rate increases to two to three times normal within 10 days following the onset of anemia. In the face of established anemia, a reticulocyte response less than two to three times normal indicates an inadequate marrow response.

To use the reticulocyte count to estimate marrow response, two corrections are necessary. The first correction adjusts the reticulocyte count based on the reduced number of circulating red cells. With anemia, the percentage of reticulocytes may be increased while the absolute number is unchanged. To correct for this effect, the reticulocyte percentage is multiplied by the ratio of the patient’s hemoglobin or hematocrit to the expected hemoglobin/hematocrit

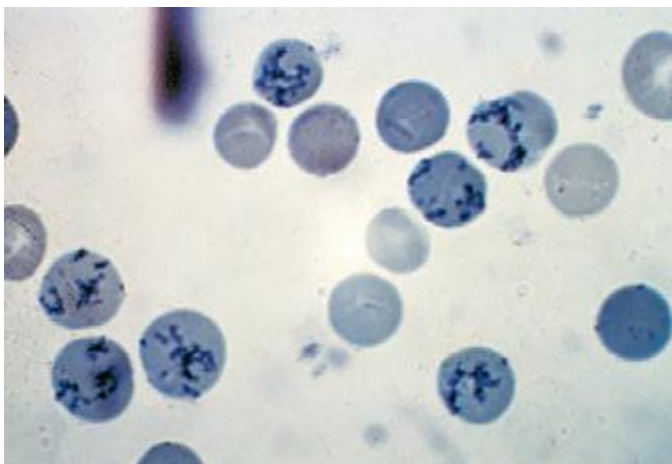


FIGURE 77-12 Reticulocytes. Methylene blue stain demonstrates residual RNA in newly made red cells. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

TABLE 77-4 CALCULATION OF RETICULOCYTE PRODUCTION INDEX

Correction #1 for Anemia:

This correction produces the corrected reticulocyte count.

In a person whose reticulocyte count is 9%, hemoglobin 7.5 g/dL, and hematocrit 23%, the absolute reticulocyte count = $9 \times (7.5/15)$ [or $\times (23/45)] = 4.5\%$

Note. This correction is not done if the reticulocyte count is reported in absolute numbers (e.g., 50,000/ μ L of blood)

Correction #2 for Longer Life of Prematurely Released Reticulocytes in the Blood:

This correction produces the reticulocyte production index.

In a person whose reticulocyte count is 9%, hemoglobin 7.5 gm/dL, and hematocrit 23%, the reticulocyte production index

$$= 9 \times \frac{(7.5/15)(\text{hemoglobin correction})}{2(\text{maturation time correction})} = 2.25$$

for the age and sex of the patient (Table 77-4). This provides an estimate of the reticulocyte count corrected for anemia. To convert the corrected reticulocyte count to an index of marrow production, a further correction is required, depending on whether some of the reticulocytes in circulation have been released from the marrow prematurely. For this second correction, the peripheral blood smear is examined to see if there are polychromatophilic macrocytes present.

These cells, representing prematurely released reticulocytes, are referred to as “shift” cells, and the relationship between the degree of shift and the necessary shift correction factor is shown in Fig. 77-13. The correction is necessary because these prematurely released cells survive as reticulocytes in circulation for >1 day, thereby providing a falsely high estimate of daily red cell production. If polychromasia is increased, the reticulocyte count, already corrected for anemia, should be divided again by 2 to account for the prolonged reticulocyte maturation time. The second correction factor varies from 1 to 3 depending on the severity of anemia. In general, a correction of 2 is simply used. An appropriate correction is shown in Table 77-4. If polychromatophilic cells are not seen on the blood smear, the second correction is not required. The now doubly

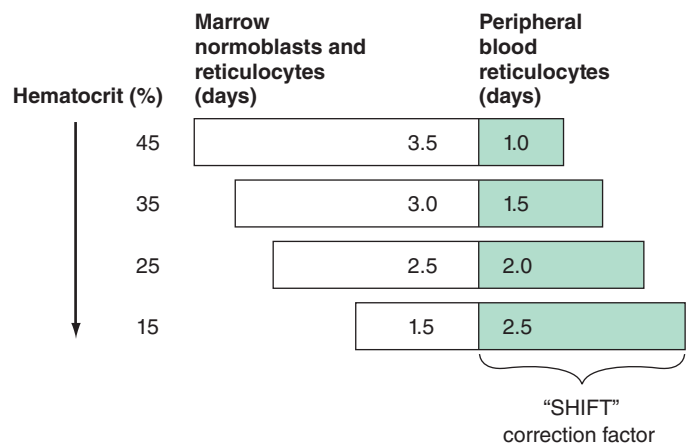


FIGURE 77-13 Correction of the reticulocyte count. To use the reticulocyte count as an indicator of effective red cell production, the reticulocyte percentage must be corrected based on the level of anemia and the circulating life span of the reticulocytes. Erythroid cells take ~4.5 days to mature. At a normal hemoglobin, reticulocytes are released to the circulation with ~1 day left as reticulocytes. However, with different levels of anemia, reticulocytes (and even earlier erythroid cells) may be released from the marrow prematurely. Most patients come to clinical attention with hematocrits in the mid-20s, and thus a correction factor of 2 is commonly used because the observed reticulocytes will live for 2 days in the circulation before losing their RNA.

TABLE 77-5 NORMAL MARROW RESPONSE TO ANEMIA

Hemoglobin	Production Index	Reticulocyte Count
15 g/dL	1	50,000/ μ L
11 g/dL	2.0–2.5	100–150,000/ μ L
8 g/dL	3.0–4.0	300–400,000/ μ L

corrected reticulocyte count is the *reticulocyte production index*, and it provides an estimate of marrow production relative to normal. In many hospital laboratories, the reticulocyte count is reported not only as a percentage but also in absolute numbers. If so, no correction for dilution is required. A summary of the appropriate marrow response to varying degrees of anemia is shown in **Table 77-5**.

Premature release of reticulocytes is normally due to increased EPO stimulation. However, if the integrity of the bone marrow release process is lost through tumor infiltration, fibrosis, or other disorders, the appearance of nucleated red cells or polychromatophilic macrocytes should still invoke the second reticulocyte correction. The shift correction should always be applied to a patient with anemia and a very high reticulocyte count to provide a true index of effective red cell production. Patients with severe chronic hemolytic anemia may increase red cell production as much as six- to sevenfold. This measure alone confirms the fact that the patient has an appropriate EPO response, a normally functioning bone marrow, and sufficient iron available to meet the demands for new red cell formation. If the reticulocyte production index is <2 in the face of established anemia, a defect in erythroid marrow proliferation or maturation must be present.

Tests of Iron Supply and Storage The laboratory measurements that reflect the availability of iron for hemoglobin synthesis include the serum iron, the TIBC, and the percent transferrin saturation. The percent transferrin saturation is derived by dividing the serum iron level ($\times 100$) by the TIBC. The normal serum iron ranges from 9 to 27 μ mol/L (50–150 μ g/dL), whereas the normal TIBC is 54–64 μ mol/L (300–360 μ g/dL); the normal transferrin saturation ranges from 25 to 50%. A diurnal variation in the serum iron leads to a variation in the percent transferrin saturation. The serum ferritin is used to evaluate total body iron stores. Adult males have serum ferritin levels that average ~ 100 μ g/L, corresponding to iron stores of ~ 1 g. Adult females have lower serum ferritin levels averaging 30 μ g/L, reflecting lower iron stores (~ 300 mg). A serum ferritin level of 10–15 μ g/L indicates depletion of body iron stores. However, ferritin is also an acute-phase reactant and, in the presence of acute or chronic inflammation, may rise several-fold above baseline levels. As a rule, a serum ferritin >200 μ g/L means there is at least some iron in tissue stores.

Bone Marrow Examination A bone marrow aspirate and smear or a needle biopsy can be useful in the evaluation of some patients with anemia. In patients with hypoproliferative anemia and normal iron status, a bone marrow is indicated. Marrow examination can diagnose primary marrow disorders such as myelofibrosis, a red cell maturation defect, or an infiltrative disease (**Figs. 77-14 to 77-16**). The increase or decrease of one cell lineage (myeloid vs erythroid) compared to another is obtained by a differential count of nucleated cells in a bone marrow smear (the myeloid/erythroid [M/E] ratio). A patient with a hypoproliferative anemia (see below) and a reticulocyte production index <2 will demonstrate an M/E ratio of 2 or 3:1. In contrast, patients with hemolytic disease and a production index >3 will have an M/E ratio of at least 1:1. Maturation disorders are identified from the discrepancy between the M/E ratio and the reticulocyte production index (see below). Either the marrow smear or biopsy can be stained for the presence of iron stores or iron in developing red cells. The storage iron is in the form of ferritin or *hemosiderin*. On carefully prepared bone marrow smears, small ferritin granules can normally be seen under oil immersion in 20–40% of developing erythroblasts. Such cells are called *sideroblasts*.

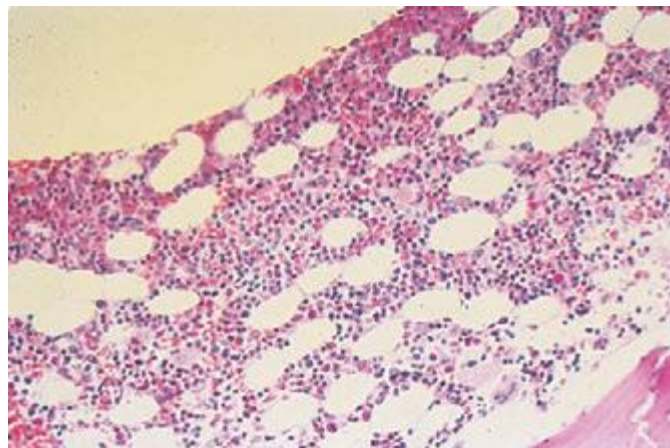


FIGURE 77-14 Normal bone marrow. This is a low-power view of a section of a normal bone marrow biopsy stained with hematoxylin and eosin (H&E). Note that the nucleated cellular elements account for ~ 40 – 50% and the fat (clear areas) accounts for ~ 50 – 60% of the area. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

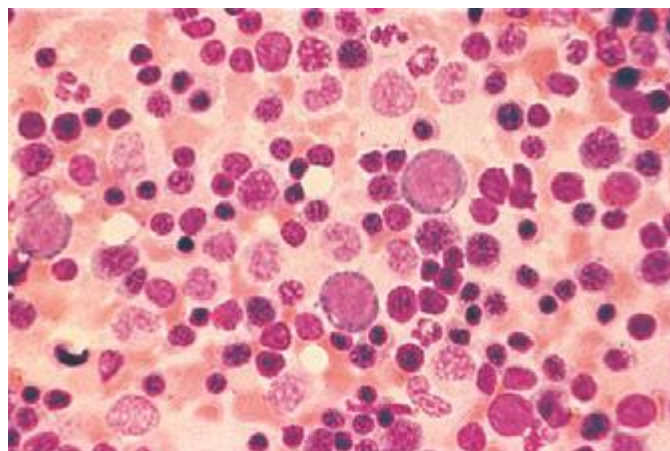


FIGURE 77-15 Erythroid hyperplasia. This marrow shows an increase in the fraction of cells in the erythroid lineage as might be seen when a normal marrow compensates for acute blood loss or hemolysis. The myeloid/erythroid (M/E) ratio is about 1:1. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

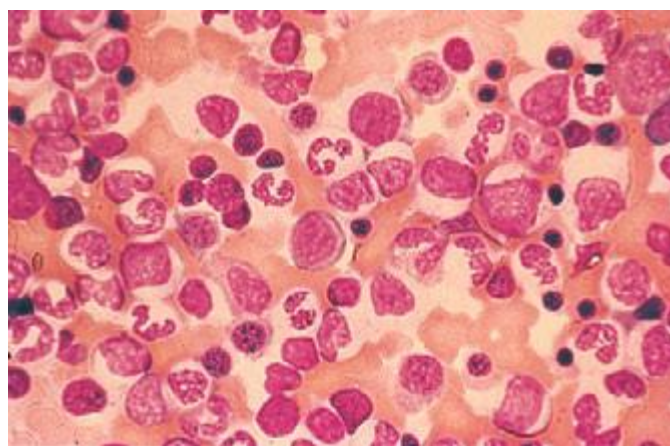


FIGURE 77-16 Myeloid hyperplasia. This marrow shows an increase in the fraction of cells in the myeloid or granulocytic lineage as might be seen in a normal marrow responding to infection. The myeloid/erythroid (M/E) ratio is $>3:1$. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2010.)

OTHER LABORATORY MEASUREMENTS

Additional laboratory tests may be of value in confirming specific diagnoses. For details of these tests and how they are applied in individual disorders, see Chaps. 126 to 130.

DEFINITION AND CLASSIFICATION OF ANEMIA

Initial Classification of Anemia The functional classification of anemia has three major categories. These are (1) marrow production defects (*hypoproliferative*), (2) red cell maturation defects (*ineffective erythropoiesis*), and (3) decreased red cell survival (*blood loss/hemolysis*). The classification is shown in Fig. 77-17. A hypoproliferative anemia is typically seen with a low reticulocyte production index together with little or no change in red cell morphology (a normocytic, normochromic anemia) (Chap. 126). Maturation disorders typically have a slight to moderately elevated reticulocyte production index that is accompanied by either macrocytic (Chap. 128) or microcytic (Chaps. 126, 127) red cell indices. Increased red blood cell destruction secondary to hemolysis results in an increase in the reticulocyte production index to at least three times normal (Chap. 129), provided sufficient iron is available. Hemorrhagic anemia does not typically result in production indices of more than 2.0–2.5 times normal because of the limitations placed on expansion of the erythroid marrow by iron availability.

In the first branch point of the classification of anemia, a reticulocyte production index >2.5 indicates that hemolysis is most likely. A reticulocyte production index <2 indicates either a hypoproliferative anemia or maturation disorder. The latter two possibilities can often be distinguished by the red cell indices, by examination of the peripheral blood smear, or by a marrow examination. If the red cell indices are normal, the anemia is almost certainly hypoproliferative in nature. Maturation disorders are characterized by ineffective red cell production and a low reticulocyte production index. Bizarre red cell shapes—macrocytes or hypochromic microcytes—are seen on the peripheral blood smear. With a hypoproliferative anemia, no erythroid

hyperplasia is noted in the marrow, whereas patients with ineffective red cell production have erythroid hyperplasia and an M/E ratio $<1:1$.

Hypoproliferative Anemias At least 75% of all cases of anemia are hypoproliferative in nature. A hypoproliferative anemia reflects absolute or relative marrow failure in which the erythroid marrow has not proliferated appropriately for the degree of anemia. The majority of hypoproliferative anemias are due to mild to moderate iron deficiency or inflammation. A hypoproliferative anemia can result from marrow damage, iron deficiency, or inadequate EPO stimulation. The last may reflect impaired renal function, suppression of EPO production by inflammatory cytokines such as interleukin 1, or reduced tissue needs for O_2 from metabolic disease such as hypothyroidism. Only occasionally is the marrow unable to produce red cells at a normal rate, and this is most prevalent in patients with renal failure. With diabetes mellitus or myeloma, the EPO deficiency may be more marked than would be predicted by the degree of renal insufficiency. In general, hypoproliferative anemias are characterized by normocytic, normochromic red cells, although microcytic, hypochromic cells may be observed with mild iron deficiency or long-standing chronic inflammatory disease. The key laboratory tests in distinguishing between the various forms of hypoproliferative anemia include the serum iron and iron-binding capacity, evaluation of renal and thyroid function, a marrow biopsy or aspirate to detect marrow damage or infiltrative disease, and serum ferritin to assess iron stores. An iron stain of the marrow will determine the pattern of iron distribution. Patients with the anemia of acute or chronic inflammation show a distinctive pattern of serum iron (low), TIBC (normal or low), percent transferrin saturation (low), and serum ferritin (normal or high). These changes in iron values are brought about by hepcidin, the iron regulatory hormone that is produced by the liver and is increased in inflammation (Chap. 126). A distinct pattern of results is noted in mild to moderate iron deficiency (low serum iron, high TIBC, low percent transferrin saturation, low serum ferritin) (Chap. 126). Marrow damage by drugs, infiltrative disease such as leukemia or lymphoma, or marrow aplasia is diagnosed from the peripheral blood and bone marrow morphology. With infiltrative disease or fibrosis, a marrow biopsy is required.

Maturation Disorders The presence of anemia with an inappropriately low reticulocyte production index, macro- or microcytosis on smear, and abnormal red cell indices suggests a maturation disorder. Maturation disorders are divided into two categories: nuclear maturation defects, associated with macrocytosis, and cytoplasmic maturation defects, associated with microcytosis and hypochromia usually from defects in hemoglobin synthesis. The inappropriately low reticulocyte production index is a reflection of the ineffective erythropoiesis that results from the destruction within the marrow of developing erythroblasts. Bone marrow examination shows erythroid hyperplasia.

Nuclear maturation defects result from vitamin B_{12} or folic acid deficiency, drug damage, or myelodysplasia. Drugs that interfere with cellular DNA synthesis, such as methotrexate or alkylating agents, can produce a nuclear maturation defect. Alcohol, alone, is also capable of producing macrocytosis and a variable degree of anemia, but this is usually associated with folic acid deficiency. Measurements of folic acid and vitamin B_{12} are critical not only in identifying the specific vitamin deficiency but also because they reflect different pathogenetic mechanisms (Chap. 128).

Cytoplasmic maturation defects result from severe iron deficiency or abnormalities in globin or heme synthesis. Iron deficiency occupies an unusual position in the classification of anemia. If the iron-deficiency anemia is mild to moderate, erythroid marrow proliferation is blunted and the anemia is classified as hypoproliferative. However, if the anemia is severe and prolonged, the erythroid marrow will become hyperplastic despite the inadequate iron supply, and the anemia will be classified as ineffective erythropoiesis with a cytoplasmic maturation defect. In either case, an inappropriately low reticulocyte production index, microcytosis, and a classic pattern of iron values make the diagnosis clear and easily distinguish iron deficiency from other cytoplasmic maturation defects such as the thalassemias. Defects in heme synthesis, in contrast to globin synthesis, are less common and

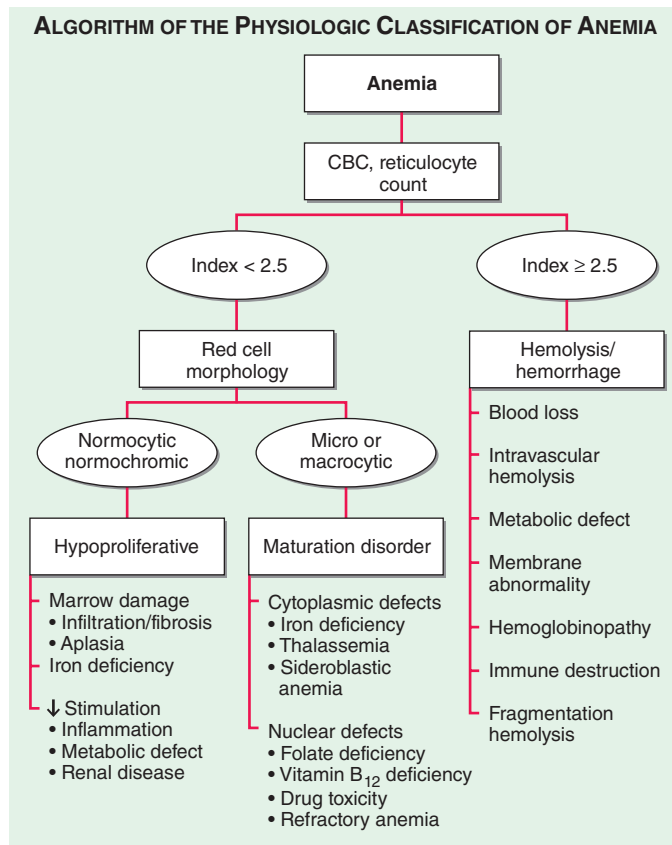


FIGURE 77-17 The physiologic classification of anemia. CBC, complete blood count.

TREATMENT ANEMIA

may be acquired or inherited (Chap. 430). Acquired abnormalities are usually associated with myelodysplasia, may lead to either a macro- or microcytic anemia, and are frequently associated with mitochondrial iron loading. In these cases, iron is taken up by the mitochondria of the developing erythroid cell but not incorporated into heme. The iron-encrusted mitochondria surround the nucleus of the erythroid cell, forming a ring. Based on the distinctive finding of so-called ringed sideroblasts on the marrow iron stain, patients are diagnosed as having a sideroblastic anemia—almost always reflecting myelodysplasia. Again, studies of iron parameters are helpful in the differential diagnosis of these patients.

Blood Loss/Hemolytic Anemia In contrast to anemias associated with an inappropriately low reticulocyte production index, hemolysis is associated with red cell production indices ≥ 2.5 times normal. The stimulated erythropoiesis is reflected in the blood smear by the appearance of increased numbers of polychromatophilic macrocytes. A marrow examination is rarely indicated if the reticulocyte production index is increased appropriately. The red cell indices are typically normocytic or slightly macrocytic, reflecting the increased number of reticulocytes. Acute blood loss is not associated with an increased reticulocyte production index because of the time required to increase EPO production and, subsequently, marrow proliferation. Subacute blood loss may be associated with modest reticulocytosis. Anemia from chronic blood loss presents more often as iron deficiency than with the picture of increased red cell production.

The evaluation of blood loss anemia is usually not difficult. Most problems arise when a patient presents with an increased red cell production index from an episode of acute blood loss that went unrecognized. The cause of the anemia and increased red cell production may not be obvious. The confirmation of a recovering state may require observations over a period of 2–3 weeks, during which the hemoglobin concentration will rise and the reticulocyte production index fall (Chap. 129).

Hemolytic disease, while dramatic, is among the least common forms of anemia. The ability to sustain a high reticulocyte production index reflects the ability of the erythroid marrow to compensate for hemolysis and, in the case of extravascular hemolysis, the efficient recycling of iron from the destroyed red cells to support red cell production. With intravascular hemolysis, such as paroxysmal nocturnal hemoglobinuria, the loss of iron may limit the marrow response. The level of response depends on the severity of the anemia and the nature of the underlying disease process.

Hemoglobinopathies, such as sickle cell disease and the thalassemias, present a mixed picture. The reticulocyte index may be high but is inappropriately low for the degree of marrow erythroid hyperplasia (Chap. 127).

Hemolytic anemias present in different ways. Some appear suddenly as an acute, self-limited episode of intravascular or extravascular hemolysis, a presentation pattern often seen in patients with autoimmune hemolysis or with inherited defects of the Embden-Meyerhof pathway or the glutathione reductase pathway. Patients with inherited disorders of the hemoglobin molecule or red cell membrane generally have a life-long clinical history typical of the disease process. Those with chronic hemolytic disease, such as hereditary spherocytosis, may actually present not with anemia but with a complication stemming from the prolonged increase in red cell destruction such as symptomatic bilirubin gallstones or splenomegaly. Patients with chronic hemolysis are also susceptible to aplastic crises if an infectious process interrupts red cell production.

The differential diagnosis of an acute or chronic hemolytic event requires the careful integration of family history, the pattern of clinical presentation, and—whether the disease is congenital or acquired—careful examination of the peripheral blood smear. Precise diagnosis may require more specialized laboratory tests, such as hemoglobin electrophoresis or a screen for red cell enzymes. Acquired defects in red cell survival are often immunologically mediated and require a direct or indirect antiglobulin test or a cold agglutinin titer to detect the presence of hemolytic antibodies or complement-mediated red cell destruction (Chap. 129).

An overriding principle is to initiate treatment of mild to moderate anemia only when a specific diagnosis is made. Rarely, in the acute setting, anemia may be so severe that red cell transfusions are required before a specific diagnosis is available. Whether the anemia is of acute or gradual onset, the selection of the appropriate treatment is determined by the documented cause(s) of the anemia. Often, the cause of the anemia is multifactorial. For example, a patient with severe rheumatoid arthritis who has been taking anti-inflammatory drugs may have a hypoproliferative anemia associated with chronic inflammation as well as chronic blood loss associated with intermittent gastrointestinal bleeding. In every circumstance, it is important to evaluate the patient's iron status fully before and during the treatment of any anemia. **Transfusion is discussed in Chap. 138e; iron therapy is discussed in Chap. 126; treatment of megaloblastic anemia is discussed in Chap. 128; treatment of other entities is discussed in their respective chapters (sickle cell anemia, Chap. 127; hemolytic anemias, Chap. 129; aplastic anemia and myelodysplasia, Chap. 130).**

Therapeutic options for the treatment of anemias have expanded dramatically during the past 30 years. Blood component therapy is available and safe. Recombinant EPO as an adjunct to anemia management has transformed the lives of patients with chronic renal failure on dialysis and reduced transfusion needs of anemic cancer patients receiving chemotherapy. Eventually, patients with inherited disorders of globin synthesis or mutations in the globin gene, such as sickle cell disease, may benefit from the successful introduction of targeted genetic therapy (Chap. 91e).

POLYCYTHEMIA

Polycythemia is defined as an increase in the hemoglobin above normal. This increase may be real or only apparent because of a decrease in plasma volume (spurious or relative polycythemia). The term *erythrocytosis* may be used interchangeably with polycythemia, but some draw a distinction between them: erythrocytosis implies documentation of increased red cell mass, whereas polycythemia refers to any increase in red cells. Often patients with polycythemia are detected through an incidental finding of elevated hemoglobin or hematocrit levels. Concern that the hemoglobin level may be abnormally high is usually triggered at 170 g/L (17 g/dL) for men and 150 g/L (15 g/dL) for women. Hematocrit levels $>50\%$ in men or $>45\%$ in women may be abnormal. Hematocrits $>60\%$ in men and $>55\%$ in women are almost invariably associated with an increased red cell mass. Given that the machine that quantitates red cell parameters actually measures hemoglobin concentrations and calculates hematocrits, hemoglobin levels may be a better index.

Features of the clinical history that are useful in the differential diagnosis include smoking history; current living at high altitude; or a history of congenital heart disease, sleep apnea, or chronic lung disease.

Patients with polycythemia may be asymptomatic or experience symptoms related to the increased red cell mass or the underlying disease process that leads to the increased red cell mass. The dominant symptoms from an increased red cell mass are related to hyperviscosity and thrombosis (both venous and arterial), because the blood viscosity increases logarithmically at hematocrits $>55\%$. Manifestations range from digital ischemia to Budd-Chiari syndrome with hepatic vein thrombosis. Abdominal vessel thromboses are particularly common. Neurologic symptoms such as vertigo, tinnitus, headache, and visual disturbances may occur. Hypertension is often present. Patients with *polycythemia vera* may have aquagenic pruritus and symptoms related to hepatosplenomegaly. Patients may have easy bruising, epistaxis, or bleeding from the gastrointestinal tract. Peptic ulcer disease is common. Patients with hypoxemia may develop cyanosis on minimal exertion or have headache, impaired mental acuity, and fatigue.

The physical examination usually reveals a ruddy complexion. Splenomegaly favors polycythemia vera as the diagnosis (Chap. 131).

The presence of cyanosis or evidence of a right-to-left shunt suggests congenital heart disease presenting in the adult, particularly tetralogy of Fallot or Eisenmenger's syndrome (**Chap. 236**). Increased blood viscosity raises pulmonary artery pressure; hypoxemia can lead to increased pulmonary vascular resistance. Together, these factors can produce cor pulmonale.

Polycythemia can be spurious (related to a decrease in plasma volume; Gaisbock's syndrome), primary, or secondary in origin. The secondary causes are all associated with increases in EPO levels: either a physiologically adapted appropriate elevation based on tissue hypoxia (lung disease, high altitude, CO poisoning, high-affinity hemoglobinopathy) or an abnormal overproduction (renal cysts, renal artery stenosis, tumors with ectopic EPO production). A rare familial form of polycythemia is associated with normal EPO levels but hyper-responsive EPO receptors due to mutations.

APPROACH TO THE PATIENT: Polycythemia

As shown in **Fig. 77-18**, the first step is to document the presence of an increased red cell mass using the principle of isotope dilution by administering ^{51}Cr -labeled autologous red blood cells to the patient and sampling blood radioactivity over a 2-h period. If the red cell mass is normal (<36 mL/kg in men, <32 mL/kg in women), the patient has spurious or relative polycythemia. If the red cell mass is increased (>36 mL/kg in men, >32 mL/kg in women), serum EPO levels should be measured. If EPO levels are low or unmeasurable, the patient most likely has polycythemia vera. A mutation in *JAK2* (Val617Phe), a key member of the cytokine intracellular signaling pathway, can be found in 90–95% of patients with polycythemia vera. Many of those without this particular *JAK2* mutation have mutations in exon 12. As a practical matter, few centers assess red

cell mass in the setting of an increased hematocrit. The short workup is to measure EPO levels, check for *JAK2* mutation, and perform an abdominal ultrasound to assess spleen size. Tests that support the diagnosis of polycythemia vera include elevated white blood cell count, increased absolute basophil count, and thrombocytosis.

If serum EPO levels are elevated, one needs to distinguish whether the elevation is a physiologic response to hypoxia or related to autonomous EPO production. Patients with low arterial O_2 saturation ($<92\%$) should be further evaluated for the presence of heart or lung disease, if they are not living at high altitude. Patients with normal O_2 saturation who are smokers may have elevated EPO levels because of CO displacement of O_2 . If carboxyhemoglobin (COHb) levels are high, the diagnosis is “smoker's polycythemia.” Such patients should be urged to stop smoking. Those who cannot stop smoking require phlebotomy to control their polycythemia. Patients with normal O_2 saturation who do not smoke either have an abnormal hemoglobin that does not deliver O_2 to the tissues (evaluated by finding elevated O_2 -hemoglobin affinity) or have a source of EPO production that is not responding to the normal feedback inhibition. Further workup is dictated by the differential diagnosis of EPO-producing neoplasms. Hepatoma, uterine leiomyoma, and renal cancer or cysts are all detectable with abdominopelvic computed tomography scans. Cerebellar hemangiomas may produce EPO, but they present with localizing neurologic signs and symptoms rather than polycythemia-related symptoms.

78 Bleeding and Thrombosis

Barbara A. Konkle

AN APPROACH TO DIAGNOSING PATIENTS WITH POLYCYTHEMIA

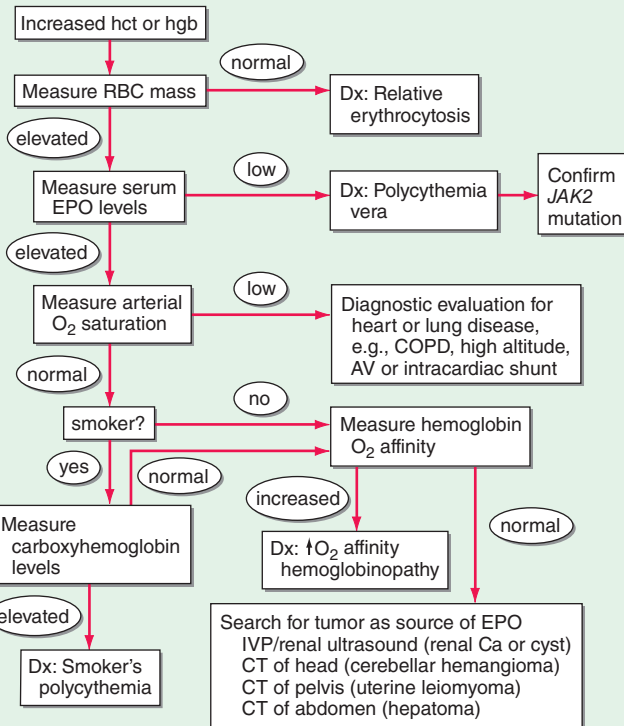


FIGURE 77-18 An approach to the differential diagnosis of patients with an elevated hemoglobin (possible polycythemia). AV, atrioventricular; COPD, chronic obstructive pulmonary disease; CT, computed tomography; EPO, erythropoietin; hct, hematocrit; hgb, hemoglobin; IVP, intravenous pyelogram; RBC, red blood cell.

The human hemostatic system provides a natural balance between procoagulant and anticoagulant forces. The procoagulant forces include platelet adhesion and aggregation and fibrin clot formation; anticoagulant forces include the natural inhibitors of coagulation and fibrinolysis. Under normal circumstances, hemostasis is regulated to promote blood flow; however, it is also prepared to clot blood rapidly to arrest blood flow and prevent exsanguination. After bleeding is successfully halted, the system remodels the damaged vessel to restore normal blood flow. The major components of the hemostatic system, which function in concert, are (1) platelets and other formed elements of blood, such as monocytes and red cells; (2) plasma proteins (the coagulation and fibrinolytic factors and inhibitors); and (3) the vessel wall.

STEPS OF NORMAL HEMOSTASIS

PLATELET PLUG FORMATION

On vascular injury, platelets adhere to the site of injury, usually the denuded vascular intimal surface. Platelet adhesion is mediated primarily by Von Willebrand factor (VWF), a large multimeric protein present in both plasma and the extracellular matrix of the subendothelial vessel wall, which serves as the primary “molecular glue,” providing sufficient strength to withstand the high levels of shear stress that would tend to detach them with the flow of blood. Platelet adhesion is also facilitated by direct binding to subendothelial collagen through specific platelet membrane collagen receptors.

Platelet adhesion results in subsequent platelet activation and aggregation. This process is enhanced and amplified by humoral mediators in plasma (e.g., epinephrine, thrombin); mediators released from activated platelets (e.g., adenosine diphosphate, serotonin); and vessel wall extracellular matrix constituents that come in contact with adherent platelets (e.g., collagen, VWF). Activated platelets undergo the release reaction, during which they secrete contents that further promote aggregation and inhibit the naturally anticoagulant endothelial cell

factors. During platelet aggregation (platelet-platelet interaction), additional platelets are recruited from the circulation to the site of vascular injury, leading to the formation of an occlusive platelet thrombus. The platelet plug is anchored and stabilized by the developing fibrin mesh.

The platelet glycoprotein (Gp) IIb/IIIa ($\alpha_{IIb}\beta_3$) complex is the most abundant receptor on the platelet surface. Platelet activation converts the normally inactive Gp IIb/IIIa receptor into an active receptor, enabling binding to fibrinogen and VWF. Because the surface of each platelet has about 50,000 Gp IIb/IIIa-binding sites, numerous activated platelets recruited to the site of vascular injury can rapidly form an occlusive aggregate by means of a dense network of intercellular fibrinogen bridges. Because this receptor is the key mediator of platelet aggregation, it has become an effective target for antiplatelet therapy.

FIBRIN CLOT FORMATION

Plasma coagulation proteins (*clotting factors*) normally circulate in plasma in their inactive forms. The sequence of coagulation protein reactions that culminate in the formation of fibrin was originally described as a *waterfall* or a *cascade*. Two pathways of blood coagulation have been described in the past: the so-called extrinsic, or tissue factor, pathway and the so-called intrinsic, or contact activation, pathway. We now know that coagulation is normally initiated through tissue factor (TF) exposure and activation through the classic *extrinsic pathway* but with critically important amplification through elements of the classic *intrinsic pathway*, as illustrated in Fig. 78-1. These reactions take place on phospholipid surfaces, usually the activated platelet surface. Coagulation testing in the laboratory can reflect other influences due to the artificial nature of the in vitro systems used (see below).

The immediate trigger for coagulation is vascular damage that exposes blood to TF that is constitutively expressed on the surfaces of subendothelial cellular components of the vessel wall, such as smooth muscle cells and fibroblasts. TF is also present in circulating microparticles, presumably shed from cells including monocytes and platelets. TF binds the serine protease factor VIIa; the complex activates factor X to factor Xa. Alternatively, the complex can indirectly activate factor X by initially converting factor IX to factor IXa, which then activates factor X. The participation of factor XI in hemostasis is not dependent on its activation by factor XIIa but rather on its positive feedback acti-

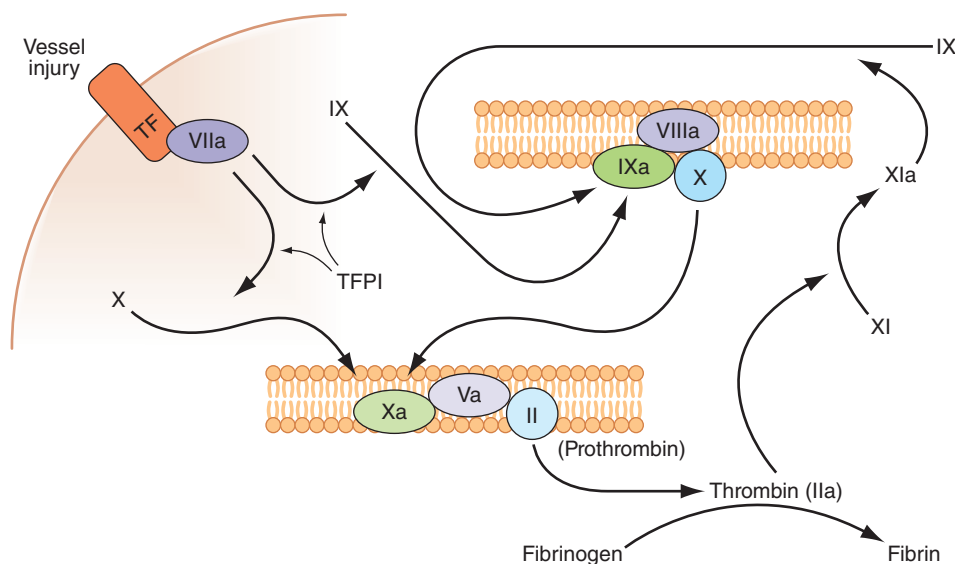


FIGURE 78-1 Coagulation is initiated by tissue factor (TF) exposure, which, with factor (F) VIIa, activates FIX and FX, which in turn, with FVIII and FV as cofactors, respectively, results in thrombin formation and subsequent conversion of fibrinogen to fibrin. Thrombin activates FXI, FVIII, and FV, amplifying the coagulation signal. Once the TF/FVIIa/FXa complex is formed, tissue factor pathway inhibitor (TFPI) inhibits the TF/FVIIa pathway, making coagulation dependent on the amplification loop through FIX/FVIII. Coagulation requires calcium (not shown) and takes place on phospholipid surfaces, usually the activated platelet membrane.

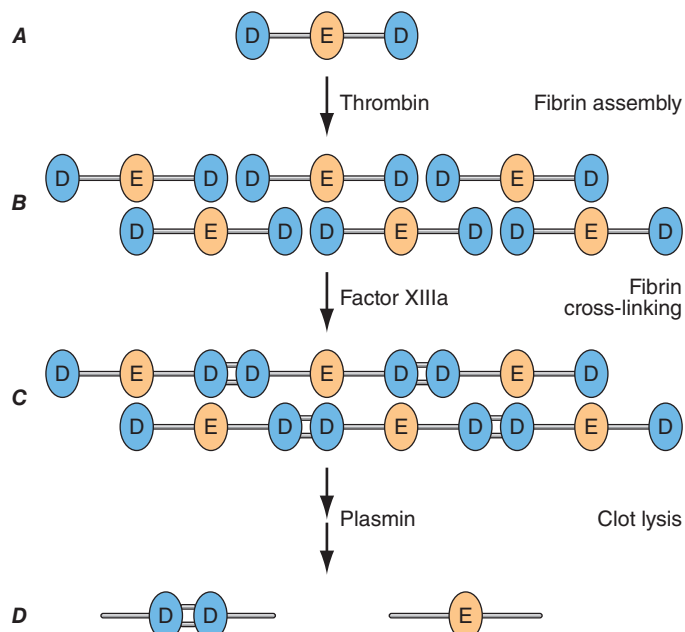


FIGURE 78-2 Fibrin formation and dissolution. (A) Fibrinogen is a trinodular structure consisting of two D domains and one E domain. Thrombin activation results in an ordered lateral assembly of protofibrils (B) with noncovalent associations. Factor XIIIa cross-links the D domains on adjacent molecules (C). Fibrin and fibrinogen (not shown) lysis by plasmin occurs at discrete sites and results in intermediary fibrin(ogen) degradation products (not shown). D-Dimers are the product of complete lysis of fibrin (D), maintaining the cross-linked D domains.

vation by thrombin. Thus, factor XIa functions in the propagation and amplification, rather than in the initiation, of the coagulation cascade.

Factor Xa can be formed through the actions of either the TF/factor VIIa complex or factor IXa (with factor VIIIa as a cofactor) and converts prothrombin to thrombin, the pivotal protease of the coagulation system. The essential cofactor for this reaction is factor Va. Like the homologous factor VIIIa, factor Va is produced by thrombin-induced limited proteolysis of factor V. Thrombin is a multifunctional enzyme that converts soluble plasma fibrinogen to an insoluble fibrin matrix. Fibrin polymerization involves an orderly process of intermolecular associations (Fig. 78-2). Thrombin also activates factor XIII (fibrin-stabilizing factor) to factor XIIIa, which covalently cross-links and thereby stabilizes the fibrin clot.

The assembly of the clotting factors on activated cell membrane surfaces greatly accelerates their reaction rates and also serves to localize blood clotting to sites of vascular injury. The critical cell membrane components, acidic phospholipids, are not normally exposed on resting cell membrane surfaces. However, when platelets, monocytes, and endothelial cells are activated by vascular injury or inflammatory stimuli, the procoagulant head groups of the membrane anionic phospholipids become translocated to the surfaces of these cells or released as part of microparticles, making them available to support and promote the plasma coagulation reactions.

Several physiologic antithrombotic mechanisms act in concert to prevent clotting under normal circumstances. These mechanisms operate to preserve blood fluidity and to limit blood clotting to specific focal sites of vascular injury. Endothelial cells have many antithrombotic effects. They produce prostacyclin, nitric oxide, and ectoADPase/CD39, which act to inhibit platelet binding, secretion, and aggregation. Endothelial cells produce anticoagulant factors including heparan proteoglycans, antithrombin, TF pathway inhibitor, and thrombomodulin. They also activate fibrinolytic mechanisms through the production of tissue plasminogen activator 1, urokinase, plasminogen activator inhibitor, and annexin-2. The sites of action of the major physiologic antithrombotic pathways are shown in Fig. 78-3.

Antithrombin (or antithrombin III) is the major plasma protease inhibitor of thrombin and the other clotting factors in coagulation. Antithrombin neutralizes thrombin and other activated coagulation factors by forming a complex between the active site of the enzyme and the reactive center of antithrombin. The rate of formation of these inactivating complexes increases by a factor of several thousand in the presence of heparin. Antithrombin inactivation of thrombin and other activated clotting factors occurs physiologically on vascular surfaces, where glycosaminoglycans, including heparan sulfates, are present to catalyze these reactions. Inherited quantitative or qualitative deficiencies of antithrombin lead to a lifelong predisposition to venous thromboembolism.

Protein C is a plasma glycoprotein that becomes an anticoagulant when it is activated by thrombin. The thrombin-induced activation of protein C occurs physiologically on thrombomodulin, a transmembrane proteoglycan-binding site for thrombin on endothelial cell surfaces. The binding of protein C to its receptor on endothelial cells places it in

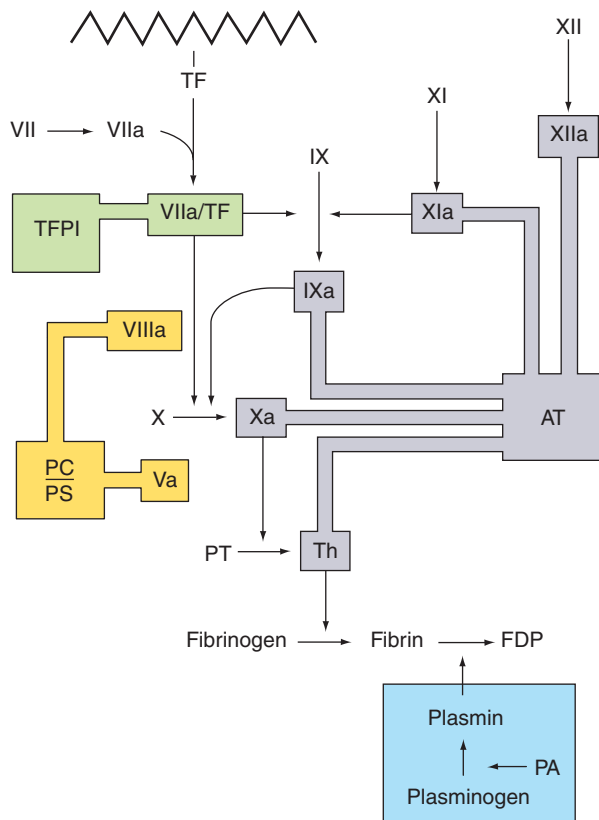


FIGURE 78-3 Sites of action of the four major physiologic antithrombotic pathways: antithrombin (AT); protein C/S (PC/PS); tissue factor pathway inhibitor (TFPI); and the fibrinolytic system, consisting of plasminogen, plasminogen activator (PA), and plasmin. PT, prothrombin; Th, thrombin; FDP, fibrin(ogen) degradation products. (Modified from BA Konkle, Al Schafer, in DP Zipes et al [eds]: Braunwald's Heart Disease, 7th ed. Philadelphia, Saunders, 2005.)

proximity to the thrombin-thrombomodulin complex, thereby enhancing its activation efficiency. Activated protein C acts as an anticoagulant by cleaving and inactivating activated factors V and VIII. This reaction is accelerated by a cofactor, protein S, which, like protein C, is a glycoprotein that undergoes vitamin K-dependent posttranslational modification. Quantitative or qualitative deficiencies of protein C or protein S, or resistance to the action of activated protein C by a specific mutation at its target cleavage site in factor Va (factor V Leiden), lead to hypercoagulable states.

Tissue factor pathway inhibitor (TFPI) is a plasma protease inhibitor that regulates the TF-induced extrinsic pathway of coagulation. TFPI inhibits the TF/factor VIIa/factor Xa complex, essentially turning off the TF/factor VIIa initiation of coagulation, which then becomes dependent on the "amplification loop" via factor XI and factor VIII activation by thrombin. TFPI is bound to lipoprotein and can also be released by heparin from endothelial cells, where it is bound to glycosaminoglycans, and from platelets. The heparin-mediated release of TFPI may play a role in the anticoagulant effects of unfractionated and low-molecular-weight heparins.

THE FIBRINOLYTIC SYSTEM

Any thrombin that escapes the inhibitory effects of the physiologic anticoagulant systems is available to convert fibrinogen to fibrin. In response, the endogenous fibrinolytic system is then activated to dispose of intravascular fibrin and thereby maintain or reestablish the patency of the circulation. Just as thrombin is the key protease enzyme of the coagulation system, plasmin is the major protease enzyme of the fibrinolytic system, acting to digest fibrin to fibrin degradation products. The general scheme of fibrinolysis and its control is shown in Fig. 78-4.

The plasminogen activators, tissue type plasminogen activator (tPA) and the urokinase-type plasminogen activator (uPA), cleave the Arg560-Val561 bond of plasminogen to generate the active enzyme plasmin. The lysine-binding sites of plasmin (and plasminogen) permit it to bind to fibrin, so that physiologic fibrinolysis is "fibrin specific." Both plasminogen (through its lysine-binding sites) and tPA possess specific affinity for fibrin and thereby bind selectively to clots. The assembly of a ternary complex, consisting of fibrin, plasminogen, and tPA, promotes the localized interaction between plasminogen and tPA and greatly accelerates the rate of plasminogen activation to plasmin. Moreover, partial degradation of fibrin by plasmin exposes new plasminogen and tPA-binding sites in carboxy-terminus lysine

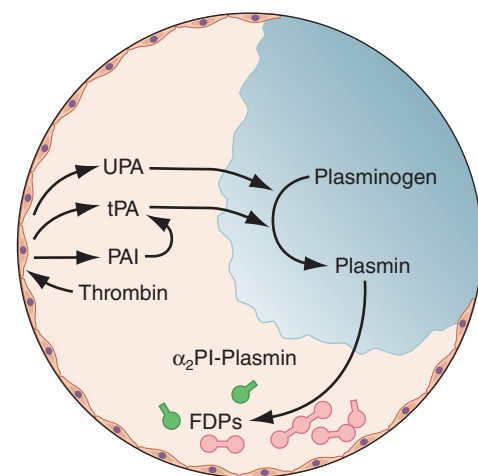


FIGURE 78-4 A schematic diagram of the fibrinolytic system.

Tissue plasminogen activator (tPA) is released from endothelial cells, binds the fibrin clot, and activates plasminogen to plasmin. Excess fibrin is degraded by plasmin to distinct degradation products (FDPs). Any free plasmin is complexed with α_2 -antiplasmin (α_2 PI). PAI, plasminogen activator inhibitor; UPA, urokinase-type plasminogen activator.

residues of fibrin fragments to enhance these reactions further. This creates a highly efficient mechanism to generate plasmin focally on the fibrin clot, which then becomes plasmin's substrate for digestion to fibrin degradation products.

Plasmin cleaves fibrin at distinct sites of the fibrin molecule, leading to the generation of characteristic fibrin fragments during the process of fibrinolysis (Fig. 78-2). The sites of plasmin cleavage of fibrin are the same as those in fibrinogen. However, when plasmin acts on covalently cross-linked fibrin, D-dimers are released; hence, D-dimers can be measured in plasma as a relatively specific test of fibrin (rather than fibrinogen) degradation. D-Dimer assays can be used as sensitive markers of blood clot formation and have been validated for clinical use to exclude the diagnosis of deep venous thrombosis (DVT) and pulmonary embolism in selected populations. In addition, D-dimer measurement can be used to stratify patients, particularly women, for risk of recurrent venous thromboembolism (VTE) when measured 1 month after discontinuation of anticoagulation given for treatment of an initial idiopathic event. D-Dimer levels may be elevated in the absence of VTE in elderly people.

Physiologic regulation of fibrinolysis occurs primarily at three levels: (1) plasminogen activator inhibitors (PAIs), specifically PAI-1 and PAI-2, inhibit the physiologic plasminogen activators; (2) the thrombin-activatable fibrinolysis inhibitor (TAFI) limits fibrinolysis; and (3) α_2 -antiplasmin inhibits plasmin. PAI-1 is the primary inhibitor of tPA and uPA in plasma. TAFI cleaves the N-terminal lysine residues of fibrin, which aid in localization of plasmin activity. α_2 -Antiplasmin is the main inhibitor of plasmin in human plasma, inactivating any nonfibrin clot-associated plasmin.

APPROACH TO THE PATIENT: Bleeding and Thrombosis

CLINICAL PRESENTATION

Disorders of hemostasis may be either inherited or acquired. A detailed personal and family history is key in determining the chronicity of symptoms and the likelihood of the disorder being inherited, as well as providing clues to underlying conditions that have contributed to the bleeding or thrombotic state. In addition, the history can give clues as to the etiology by determining (1) the bleeding (mucosal and/or joint) or thrombosis (arterial and/or venous) site and (2) whether an underlying bleeding or clotting tendency was enhanced by another medical condition or the introduction of medications or dietary supplements.

History of Bleeding A history of bleeding is the most important predictor of bleeding risk. In evaluating a patient for a bleeding disorder, a history of at-risk situations, including the response to past surgeries, should be assessed. Does the patient have a history of spontaneous or trauma/surgery-induced bleeding? Spontaneous hemarthroses are a hallmark of moderate and severe factor VIII and IX deficiency and, in rare circumstances, of other clotting factor deficiencies. Mucosal bleeding symptoms are more suggestive of underlying platelet disorders or Von Willebrand disease (VWD), termed *disorders of primary hemostasis or platelet plug formation*. Disorders affecting primary hemostasis are shown in [Table 78-1](#).

A bleeding score has been validated as a tool to predict patients more likely to have type 1 VWD (International Society on Thrombosis and Haemostasis Bleeding Assessment Tool [www.isth.org/resource/resmgr/ssc/isth-ssc_bleeding_assessment.pdf]). This is most useful tool in excluding the diagnosis of a bleeding disorder, and thus avoiding unnecessary testing. One study found that a low bleeding score (≤ 3) and a normal activated partial thromboplastin time (aPTT) had 99.6% negative predictive value for the diagnosis of VWD. Bleeding symptoms that appear to be more common in patients with bleeding disorders include prolonged bleeding with surgery, dental procedures and extractions, and/or trauma, menorrhagia or postpartum hemorrhage, and large bruises (often described with lumps).

TABLE 78-1 PRIMARY HEMOSTATIC (PLATELET PLUG) DISORDERS

Defects of Platelet Adhesion

Von Willebrand disease

Bernard-Soulier syndrome (absence or dysfunction of platelet Gp Ib-IX-V)

Defects of Platelet Aggregation

Glanzmann's thrombasthenia (absence or dysfunction of platelet glycoprotein [Gp] IIb/IIIa)

Afibrinogenemia

Defects of Platelet Secretion

Decreased cyclooxygenase activity

Drug-induced (aspirin, nonsteroidal anti-inflammatory agents, thienopyridines)

Inherited

Granule storage pool defects

Inherited

Acquired

Nonspecific inherited secretory defects

Nonspecific drug effects

Uremia

Platelet coating (e.g., paraprotein, penicillin)

Defect of Platelet Coagulant Activity

Scott's syndrome

Easy bruising and menorrhagia are common complaints in patients with and without bleeding disorders. Easy bruising can also be a sign of medical conditions in which there is no identifiable coagulopathy; instead, the conditions are caused by an abnormality of blood vessels or their supporting tissues. In Ehlers-Danlos syndrome, there may be posttraumatic bleeding and a history of joint hyperextensibility. Cushing's syndrome, chronic steroid use, and aging result in changes in skin and subcutaneous tissue, and subcutaneous bleeding occurs in response to minor trauma. The latter has been termed *senile purpura*.

Epistaxis is a common symptom, particularly in children and in dry climates, and may not reflect an underlying bleeding disorder. However, it is the most common symptom in hereditary hemorrhagic telangiectasia and in boys with VWD. Clues that epistaxis is a symptom of an underlying bleeding disorder include lack of seasonal variation and bleeding that requires medical evaluation or treatment, including cauterization. Bleeding with eruption of primary teeth is seen in children with more severe bleeding disorders, such as moderate and severe hemophilia. It is uncommon in children with mild bleeding disorders. Patients with disorders of primary hemostasis (platelet adhesion) may have increased bleeding after dental cleanings and other procedures that involve gum manipulation.

Menorrhagia is defined quantitatively as a loss of >80 mL of blood per cycle, based on the quantity of blood loss required to produce iron-deficiency anemia. A complaint of heavy menses is subjective and has a poor correlation with excessive blood loss. Predictors of menorrhagia include bleeding resulting in iron-deficiency anemia or a need for blood transfusion, passage of clots >1 inch in diameter, and changing a pad or tampon more than hourly. Menorrhagia is a common symptom in women with underlying bleeding disorders and is reported in the majority of women with VWD, women with factor XI deficiency, and symptomatic carriers of hemophilia. Women with underlying bleeding disorders are more likely to have other bleeding symptoms, including bleeding after dental extractions, postoperative bleeding, and postpartum bleeding, and are much more likely to have menorrhagia beginning at menarche than women with menorrhagia due to other causes.

Postpartum hemorrhage (PPH) is a common symptom in women with underlying bleeding disorders. In women with type 1 VWD and symptomatic carriers of hemophilia A in whom levels of VWF and factor VIII usually normalize during pregnancy, PPH

may be delayed. Women with a history of PPH have a high risk of recurrence with subsequent pregnancies. Rupture of ovarian cysts with intraabdominal hemorrhage has also been reported in women with underlying bleeding disorders.

Tonsillectomy is a major hemostatic challenge, because intact hemostatic mechanisms are essential to prevent excessive bleeding from the tonsillar bed. Bleeding may occur early after surgery or after approximately 7 days postoperatively, with loss of the eschar at the operative site. Similar delayed bleeding is seen after colonic polyp resection. Gastrointestinal (GI) bleeding and hematuria are usually due to underlying pathology, and procedures to identify and treat the bleeding site should be undertaken, even in patients with known bleeding disorders. VWD, particularly types 2 and 3, has been associated with angiodysplasia of the bowel and GI bleeding.

Hemarthroses and spontaneous muscle hematomas are characteristic of moderate or severe congenital factor VIII or IX deficiency. They can also be seen in moderate and severe deficiencies of fibrinogen, prothrombin, and factors V, VII, and X. Spontaneous hemarthroses occur rarely in other bleeding disorders except for severe VWD, with associated factor VIII levels <5%. Muscle and soft tissue bleeds are also common in acquired factor VIII deficiency. Bleeding into a joint results in severe pain and swelling, as well as loss of function, but is rarely associated with discoloration from bruising around the joint. Life-threatening sites of bleeding include bleeding into the oropharynx, where bleeding can obstruct the airway, into the central nervous system, and into the retroperitoneum. Central nervous system bleeding is the major cause of bleeding-related deaths in patients with severe congenital factor deficiencies.

Prohemorrhagic Effects of Medications and Dietary Supplements

Aspirin and other nonsteroidal anti-inflammatory drugs (NSAIDs) that inhibit cyclooxygenase 1 impair primary hemostasis and may exacerbate bleeding from another cause or even unmask a previously occult mild bleeding disorder such as VWD. All NSAIDs, however, can precipitate GI bleeding, which may be more severe in patients with underlying bleeding disorders. The aspirin effect on platelet function as assessed by aggregometry can persist for up to 7 days, although it has frequently returned to normal by 3 days after the last dose. The effect of other NSAIDs is shorter, as the inhibitor effect is reversed when the drug is removed. Thienopyridines (clopidogrel and prasugrel) inhibit ADP-mediated platelet aggregation and, like NSAIDs, can precipitate or exacerbate bleeding symptoms.

Many herbal supplements can impair hemostatic function (Table 78-2). Some are more convincingly associated with a bleeding risk than others. Fish oil or concentrated omega-3 fatty acid supplements impair platelet function. They alter platelet biochemistry to produce more PGI₃, a more potent platelet inhibitor than prostacyclin (PGI₂), and more thromboxane A₃, a less potent platelet activator than thromboxane A₂. In fact, diets naturally rich in omega-3 fatty acids can result in a prolonged bleeding time and abnormal platelet aggregation studies, but the actual associated bleeding risk is unclear. Vitamin E appears to inhibit protein kinase C-mediated platelet aggregation and nitric oxide production. In patients with unexplained bruising or bleeding, it is prudent to review any new medications or supplements and discontinue those that may be associated with bleeding.

Underlying Systemic Diseases That Cause or Exacerbate a Bleeding Tendency

Acquired bleeding disorders are commonly secondary to, or associated with, systemic disease. The clinical evaluation of a patient with a bleeding tendency must therefore include a thorough assessment for evidence of underlying disease. Bruising or mucosal bleeding may be the presenting complaint in liver disease, severe renal impairment, hypothyroidism, paraproteinemias or amyloidosis, and conditions causing bone marrow failure. All coagulation factors are synthesized in the liver, and hepatic failure results in combined factor deficiencies. This is often compounded by thrombocytopenia from splenomegaly due to portal

TABLE 78-2 HERBAL SUPPLEMENTS ASSOCIATED WITH INCREASED BLEEDING

Herbs with Potential Antiplatelet Activity

Ginkgo (*Ginkgo biloba* L.)

Garlic (*Allium sativum*)

Bilberry (*Vaccinium myrtillus*)

Ginger (*Zingiber officinale*)

Dong quai (*Angelica sinensis*)

Feverfew (*Tanacetum parthenium*)

Asian ginseng (*Panax ginseng*)

American ginseng (*Panax quinquefolius*)

Siberian ginseng/eleuthero (*Eleutherococcus senticosus*)

Turmeric (*Curcuma longa*)

Meadowsweet (*Filipendula ulmaria*)

Willow (*Salix* spp.)

Coumarin-Containing Herbs

Motherwort (*Leonurus cardiaca*)

Chamomile (*Matricaria recutita*, *Chamaemelum mobile*)

Horse chestnut (*Aesculus hippocastanum*)

Red clover (*Trifolium pratense*)

Fenugreek (*Trigonella foenum-graecum*)

hypertension. Coagulation factors II, VII, IX, and X and proteins C, S, and Z are dependent on vitamin K for posttranslational modification. Although vitamin K is required in both procoagulant and anticoagulant processes, the phenotype of vitamin K deficiency or the warfarin effect on coagulation is bleeding.

The normal blood platelet count is 150,000–450,000/μL. Thrombocytopenia results from decreased production, increased destruction, and/or sequestration. Although the bleeding risk varies somewhat by the reason for the thrombocytopenia, bleeding rarely occurs in isolated thrombocytopenia at counts <50,000/μL and usually not until <10,000–20,000/μL. Coexisting coagulopathies, as is seen in liver failure or disseminated coagulation; infection; platelet-inhibitory drugs; and underlying medical conditions can all increase the risk of bleeding in the thrombocytopenic patient. Most procedures can be performed in patients with a platelet count of 50,000/μL. The level needed for major surgery will depend on the type of surgery and the patient's underlying medical state, although a count of approximately 80,000/μL is likely sufficient.

HISTORY OF THROMBOSIS

The risk of thrombosis, like that of bleeding, is influenced by both genetic and environmental influences. The major risk factor for arterial thrombosis is atherosclerosis, whereas for venous thrombosis, the risk factors are immobility, surgery, underlying medical conditions such as malignancy, medications such as hormonal therapy, obesity, and genetic predispositions. Factors that increase risks for venous and for both venous and arterial thromboses are shown in Table 78-3.

The most important point in a history related to venous thrombosis is determining whether the thrombotic event was idiopathic (meaning there was no clear precipitating factor) or was a precipitated event. In patients without underlying malignancy, having an idiopathic event is the strongest predictor of recurrence of VTE. In patients who have a vague history of thrombosis, a history of being treated with warfarin suggests a past DVT. Age is an important risk factor for venous thrombosis—the risk of DVT increases per decade, with an approximate incidence of 1/100,000 per year in early childhood to 1/200 per year among octogenarians. Family history is helpful in determining if there is a genetic predisposition and how strong that predisposition appears to be. A genetic thrombophilia that confers a relatively small increased risk, such as being a heterozygote for the prothrombin G20210A or factor V Leiden mutation, may be a minor determinant of risk in an elderly individual

TABLE 78-3 RISK FACTORS FOR THROMBOSIS

Venous	Venous and Arterial
Inherited	Inherited
Factor V Leiden	Homocystinuria
Prothrombin G20210A	Dysfibrinogenemia
Antithrombin deficiency	Mixed (inherited and acquired)
Protein C deficiency	Hyperhomocysteinemia
Protein S deficiency	Acquired
Elevated factor VIII	Malignancy
Acquired	Antiphospholipid antibody syndrome
Age	Hormonal therapy
Previous thrombosis	Polycythemia vera
Immobilization	Essential thrombocythemia
Major surgery	Paroxysmal nocturnal hemoglobinuria
Pregnancy and puerperium	Thrombotic thrombocytopenic purpura
Hospitalization	Heparin-induced thrombocytopenia
Obesity	Disseminated intravascular coagulation
Infection	
APC resistance, nongenetic	
Smoking	
Unknown^a	
Elevated factor II, IX, XI	
Elevated TAFI levels	
Low levels of TFPI	

^aUnknown whether risk is inherited or acquired.

Abbreviations: APC, activated protein C; TAFI, thrombin-activatable fibrinolysis inhibitor; TFPI, tissue factor pathway inhibitor.

undergoing a high-risk surgical procedure. As illustrated in **Fig. 78-5**, a thrombotic event usually has more than one contributing factor. Predisposing factors must be carefully assessed to determine the risk of recurrent thrombosis and, with consideration of the patient's bleeding risk, determine the length of anticoagulation.

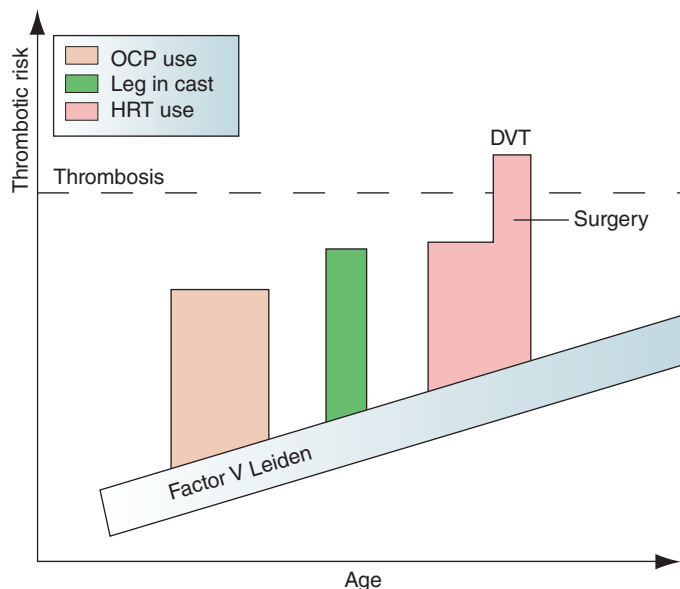


FIGURE 78-5 Thrombotic risk over time. Shown schematically is an individual's thrombotic risk over time. An underlying factor V Leiden mutation provides a "theoretically" constant increased risk. The thrombotic risk increases with age and, intermittently, with oral contraceptive (OCP) or hormone replacement therapy (HRT) use; other events may increase the risk further. At some point, the cumulative risk may increase to the threshold for thrombosis and result in deep venous thrombosis (DVT). Note: The magnitude and duration of risk portrayed in the figure are meant for example only and may not precisely reflect the relative risk determined by clinical study. (From BA Konkle, A Schafer, in DP Zipes et al [eds]: *Braunwald's Heart Disease*, 7th ed. Philadelphia, Saunders, 2005; modified with permission from FR Rosendaal: *Venous thrombosis: A multicausal disease*. *Lancet* 353:1167, 1999.)

Similar consideration should be given in determining the need, if any, to test the patient and family members for thrombophilias.

LABORATORY EVALUATION

Careful history taking and clinical examination are essential components in the assessment of bleeding and thrombotic risk. The use of laboratory tests of coagulation complement, but cannot substitute for, clinical assessment. No test exists that provides a global assessment of hemostasis. The bleeding time has been used to assess bleeding risk; however, it does not predict bleeding risk with surgery and it is not recommended for this indication. The PFA-100, an instrument that measures platelet-dependent coagulation under flow conditions, is more sensitive and specific for VWD than the bleeding time; however, it is not sensitive enough to rule out mild bleeding disorders. PFA-100 closure times are prolonged in patients with some, but not all, inherited platelet disorders. Also, its utility in predicting bleeding risk has not been determined.

For routine preoperative and preprocedure testing, an abnormal prothrombin time (PT) may detect liver disease or vitamin K deficiency that had not been previously appreciated. Studies have not confirmed the usefulness of an aPTT in preoperative evaluations in patients with a negative bleeding history. The primary use of coagulation testing should be to confirm the presence and type of bleeding disorder in a patient with a suspicious clinical history.

Because of the nature of coagulation assays, proper sample acquisition and handling is critical to obtaining valid results. In patients with abnormal coagulation assays who have no bleeding history, repeat studies with attention to these factors frequently results in normal values. Most coagulation assays are performed in sodium citrate anticoagulated plasma that is recalcified for the assay. Because the anticoagulant is in liquid solution and needs to be added to blood in proportion to the plasma volume, incorrectly filled or inadequately mixed blood collection tubes will give erroneous results. Vacutainer tubes should be filled to >90% of the recommended fill, which is usually denoted by a line on the tube. An elevated hematocrit (>55%) can result in a false value due to a decreased plasma-to-anticoagulant ratio.

Screening Assays The most commonly used screening tests are the PT, aPTT, and platelet count. The PT assesses the factors I (fibrinogen), II (prothrombin), V, VII, and X (**Fig. 78-6**). The PT measures the time for clot formation of the citrated plasma after recalcification and addition of thromboplastin, a mixture of TF and phospholipids. The sensitivity of the assay varies by the source of thromboplastin. The relationship between defects in secondary hemostasis (fibrin formation) and coagulation test abnormalities is shown in **Table 78-4**. To adjust for this variability, the overall sensitivity of different thromboplastins to reduction of the vitamin K-dependent clotting factors II, VII, IX, and X in anticoagulation patients is now expressed as the International Sensitivity Index (ISI). An inverse relationship exists between ISI and thromboplastin sensitivity. The international normalized ratio (INR) is then determined based on the formula: $INR = (PT_{patient} / PT_{normal\ mean})^{ISI}$.

The INR was developed to assess stable anticoagulation due to reduction of vitamin K-dependent coagulation factors; it is commonly used in the evaluation of patients with liver disease. Although it does allow comparison between laboratories, reagent sensitivity as used to determine the ISI is not the same in liver disease as with warfarin anticoagulation. In addition, progressive liver failure is associated with variable changes in coagulation factors; the degree of prolongation of either the PT or the INR only roughly predicts the bleeding risk. Thrombin generation has been shown to be normal in many patients with mild to moderate liver dysfunction. Because the PT only measures one aspect of hemostasis affected by liver dysfunction, we likely overestimate the bleeding risk of a mildly elevated INR in this setting.

The aPTT assesses the intrinsic and common coagulation pathways; factors XI, IX, VIII, X, V, and II; fibrinogen; prekallikrein;

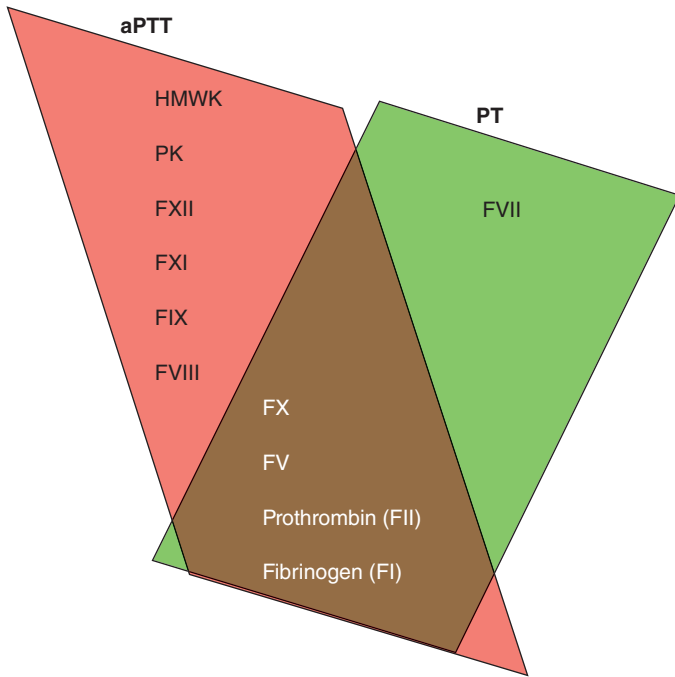


FIGURE 78-6 Coagulation factor activity tested in the activated partial thromboplastin time (aPTT) in red and prothrombin time (PT) in green, or both. F, factor; HMWK, high-molecular-weight kininogen; PK, prekallikrein.

TABLE 78-4 HEMOSTATIC DISORDERS AND COAGULATION TEST ABNORMALITIES

Prolonged Activated Partial Thromboplastin Time (aPTT)

No clinical bleeding—↓ factor XII, high-molecular-weight kininogen, prekallikrein
 Variable, but usually mild, bleeding—↓ factor XI, mild ↓ factor VIII and factor IX
 Frequent, severe bleeding—severe deficiencies of factors VIII and IX
 Heparin and direct thrombin inhibitors

Prolonged Prothrombin Time (PT)

Factor VII deficiency
 Vitamin K deficiency—early
 Warfarin anticoagulation
 Direct Xa inhibitors (rivaroxaban, apixaban)

Prolonged aPTT and PT

Factor II, V, X, or fibrinogen deficiency
 Vitamin K deficiency—late
 Direct thrombin inhibitors

Prolonged Thrombin Time

Heparin or heparin-like inhibitors
 Direct thrombin inhibitors (e.g., dabigatran, argatroban, bivalirudin)
 Mild or no bleeding—dysfibrinogenemia
 Frequent, severe bleeding—afibrinogenemia

Prolonged PT and/or aPTT Not Corrected with Mixing with Normal Plasma

Bleeding—specific factor inhibitor
 No symptoms, or clotting and/or pregnancy loss—lupus anticoagulant
 Disseminated intravascular coagulation
 Heparin or direct thrombin inhibitor

Abnormal Clot Solubility

Factor XIII deficiency
 Inhibitors or defective cross-linking

Rapid Clot Lysis

Deficiency of α_2 -antiplasmin or plasminogen activator inhibitor 1
 Treatment with fibrinolytic therapy

high-molecular-weight kininogen; and factor XII (Fig. 78-6). The aPTT reagent contains phospholipids derived from either animal or vegetable sources that function as a platelet substitute in the coagulation pathways and includes an activator of the intrinsic coagulation system, such as nonparticulate ellagic acid or the particulate activators kaolin, celite, or micronized silica.

The phospholipid composition of aPTT reagents varies, which influences the sensitivity of individual reagents to clotting factor deficiencies and to inhibitors such as heparin and lupus anticoagulants. Thus, aPTT results will vary from one laboratory to another, and the normal range in the laboratory where the testing occurs should be used in the interpretation. Local laboratories can relate their aPTT values to the therapeutic heparin anticoagulation by correlating aPTT values with direct measurements of heparin activity (anti-Xa or protamine titration assays) in samples from heparinized patients, although correlation between these assays is often poor. The aPTT reagent will vary in sensitivity to individual factor deficiencies and usually becomes prolonged with individual factor deficiencies of 30–50%.

Mixing Studies Mixing studies are used to evaluate a prolonged aPTT or, less commonly PT, to distinguish between a factor deficiency and an inhibitor. In this assay, normal plasma and patient plasma are mixed in a 1:1 ratio, and the aPTT or PT is determined immediately and after incubation at 37°C for varying times, typically 30, 60, and/or 120 min. With isolated factor deficiencies, the aPTT will correct with mixing and stay corrected with incubation. With aPTT prolongation due to a lupus anticoagulant, the mixing and incubation will show no correction. In acquired neutralizing factor antibodies, notably an acquired factor VIII inhibitor, the initial assay may or may not correct immediately after mixing but will prolong or remain prolonged with incubation at 37°C. Failure to correct with mixing can also be due to the presence of other inhibitors or interfering substances such as heparin, fibrin split products, and paraproteins.

Specific Factor Assays Decisions to proceed with specific clotting factor assays will be influenced by the clinical situation and the results of coagulation screening tests. Precise diagnosis and effective management of inherited and acquired coagulation deficiencies necessitate quantitation of the relevant factors. When bleeding is severe, specific assays are urgently required to guide appropriate therapy. Individual factor assays are usually performed as modifications of the mixing study, where the patient's plasma is mixed with plasma deficient in the factor being studied. This will correct all factor deficiencies to >50%, thus making prolongation of clot formation due to a factor deficiency dependent on the factor missing from the added plasma.

Testing for Antiphospholipid Antibodies Antibodies to phospholipids (cardiolipin) or phospholipid-binding proteins (β_2 -microglobulin and others) are detected by enzyme-linked immunosorbent assay (ELISA). When these antibodies interfere with phospholipid-dependent coagulation tests, they are termed *lupus anticoagulants*. The aPTT has variability sensitivity to lupus anticoagulants, depending in part on the aPTT reagents used. An assay using a sensitive reagent has been termed an *LA-PTT*. The dilute Russell viper venom test (dRVVT) and the tissue thromboplastin inhibition (TTI) test are modifications of standard tests with the phospholipid reagent decreased, thus increasing the sensitivity to antibodies that interfere with the phospholipid component. The tests, however, are not specific for lupus anticoagulants, because factor deficiencies or other inhibitors will also result in prolongation. Documentation of a lupus anticoagulant requires not only prolongation of a phospholipid-dependent coagulation test but also lack of correction when mixed with normal plasma and correction when the addition of activated platelet membranes or certain phospholipids (e.g., hexagonal phase).

79

Enlargement of Lymph Nodes and Spleen

Patrick H. Henry, Dan L. Longo

Other Coagulation Tests The thrombin time and the reptilase time measure fibrinogen conversion to fibrin and are prolonged when the fibrinogen level is low (usually <80–100 mg/dL) or qualitatively abnormal, as seen in inherited or acquired dysfibrinogenemias, or when fibrin/fibrinogen degradation products interfere. The thrombin time, but not the reptilase time, is prolonged in the presence of heparin. The thrombin time is markedly prolonged in the presence of the direct thrombin inhibitor, dabigatran; a dilute thrombin time can be used to assess drug activity. Measurement of anti-factor Xa plasma inhibitory activity is a test frequently used to assess low-molecular-weight heparin (LMWH) levels, as a direct measurement of unfractionated heparin (UFH) activity, or to assess activity of the new direct Xa inhibitors rivaroxaban or apixaban. Drug in the patient sample inhibits the enzymatic conversion of an Xa-specific chromogenic substrate to colored product by factor Xa. Standard curves are created using multiple concentrations of drug and are used to calculate the concentration of anti-Xa activity in the patient plasma.

Laboratory Testing for Thrombophilia Laboratory assays to detect thrombophilic states include molecular diagnostics and immunologic and functional assays. These assays vary in their sensitivity and specificity for the condition being tested. Furthermore, acute thrombosis, acute illnesses, inflammatory conditions, pregnancy, and medications affect levels of many coagulation factors and their inhibitors. Antithrombin is decreased by heparin and in the setting of acute thrombosis. Protein C and S levels may be increased in the setting of acute thrombosis and are decreased by warfarin. Antiphospholipid antibodies are frequently transiently positive in acute illness. Testing for genetic thrombophilias should, in general, only be performed when there is a strong family history of thrombosis and results would affect clinical decision making.

Because thrombophilia evaluations are usually performed to assess the need to extend anticoagulation, testing should be performed in a steady state, remote from the acute event. In most instances, warfarin anticoagulation can be stopped after the initial 3–6 months of treatment, and testing can be performed at least 3 weeks later. As a sensitive marker of coagulation activation, the quantitative D-dimer assay, drawn 4 weeks after stopping anticoagulation, can be used to stratify risk of recurrent thrombosis in patients who have an idiopathic event.

Measures of Platelet Function The bleeding time has been used to assess bleeding risk; however, it has not been found to predict bleeding risk with surgery, and it is not recommended for use for this indication. The PFA-100 and similar instruments that measure platelet-dependent coagulation under flow conditions are generally more sensitive and specific for platelet disorders and VWD than the bleeding time; however, data are insufficient to support their use to predict bleeding risk or monitor response to therapy, and they will be normal in some patients with platelet disorders or mild VWD. When they are used in the evaluation of a patient with bleeding symptoms, abnormal results, as with the bleeding time, require specific testing, such as VWF assays and/or platelet aggregation studies. Because all of these “screening” assays may miss patients with mild bleeding disorders, further studies are needed to define their role in hemostasis testing.

For classic platelet aggregometry, various agonists are added to the patient's platelet-rich plasma and platelet aggregation is measured. Tests of platelet secretion in response to agonists can also be measured. These tests are affected by many factors, including numerous medications, and the association between minor defects in aggregation or secretion in these assays and bleeding risk is not clearly established.

ACKNOWLEDGMENT

Robert I. Handin, MD, contributed this chapter in the 16th edition, and some material from that chapter has been retained here.

This chapter is intended to serve as a guide to the evaluation of patients who present with enlargement of the lymph nodes (*lymphadenopathy*) or the spleen (*splenomegaly*). Lymphadenopathy is a rather common clinical finding in primary care settings, whereas palpable splenomegaly is less so.

LYMPHADENOPATHY

Lymphadenopathy may be an incidental finding in patients being examined for various reasons, or it may be a presenting sign or symptom of the patient's illness. The physician must eventually decide whether the lymphadenopathy is a normal finding or one that requires further study, up to and including biopsy. Soft, flat, submandibular nodes (<1 cm) are often palpable in healthy children and young adults; healthy adults may have palpable inguinal nodes of up to 2 cm, which are considered normal. Further evaluation of these normal nodes is not warranted. In contrast, if the physician believes the node(s) to be abnormal, then pursuit of a more precise diagnosis is needed.

**APPROACH TO THE PATIENT:
Lymphadenopathy**

Lymphadenopathy may be a primary or secondary manifestation of numerous disorders, as shown in [Table 79-1](#). Many of these disorders are infrequent causes of lymphadenopathy. In primary care practice, more than two-thirds of patients with lymphadenopathy have nonspecific causes or upper respiratory illnesses (viral or bacterial), and <1% have a malignancy. In one study, 84% of patients referred for evaluation of lymphadenopathy had a “benign” diagnosis. The remaining 16% had a malignancy (lymphoma or metastatic adenocarcinoma). Of the patients with benign lymphadenopathy, 63% had a nonspecific or reactive etiology (no causative agent found), and the remainder had a specific cause demonstrated, most commonly infectious mononucleosis, toxoplasmosis, or tuberculosis. Thus, the vast majority of patients with lymphadenopathy will have a nonspecific etiology requiring few diagnostic tests.

CLINICAL ASSESSMENT

The physician will be aided in the pursuit of an explanation for the lymphadenopathy by a careful medical history, physical examination, selected laboratory tests, and perhaps an excisional lymph node biopsy.

The *medical history* should reveal the setting in which lymphadenopathy is occurring. Symptoms such as sore throat, cough, fever, night sweats, fatigue, weight loss, or pain in the nodes should be sought. The patient's age, sex, occupation, exposure to pets, sexual behavior, and use of drugs such as diphenylhydantoin are other important historic points. For example, children and young adults usually have benign (i.e., nonmalignant) disorders that account for the observed lymphadenopathy such as viral or bacterial upper respiratory infections; infectious mononucleosis; toxoplasmosis; and, in some countries, tuberculosis. In contrast, after age 50, the incidence of malignant disorders increases and that of benign disorders decreases.

The *physical examination* can provide useful clues such as the extent of lymphadenopathy (localized or generalized), size of nodes, texture, presence or absence of nodal tenderness, signs of inflammation over the node, skin lesions, and splenomegaly. A thorough ear, nose, and throat (ENT) examination is indicated in adult patients

TABLE 79-1 DISEASES ASSOCIATED WITH LYMPHADENOPATHY

1. Infectious diseases
 - a. Viral—*infectious mononucleosis syndromes (EBV, CMV), infectious hepatitis, herpes simplex, herpesvirus-6, varicella-zoster virus, rubella, measles, adenovirus, HIV, epidemic keratoconjunctivitis, vaccinia, herpesvirus-8*
 - b. Bacterial—*streptococci, staphylococci, cat-scratch disease, brucellosis, tularemia, plague, chancroid, melioidosis, glanders, tuberculosis, atypical mycobacterial infection, primary and secondary syphilis, diphtheria, leprosy, Bartonella*
 - c. Fungal—*histoplasmosis, coccidioidomycosis, paracoccidioidomycosis*
 - d. Chlamydial—*lymphogranuloma venereum, trachoma*
 - e. Parasitic—*toxoplasmosis, leishmaniasis, trypanosomiasis, filariasis*
 - f. Rickettsial—*scrub typhus, rickettsialpox, Q fever*
2. Immunologic diseases
 - a. Rheumatoid arthritis
 - b. Juvenile rheumatoid arthritis
 - c. Mixed connective tissue disease
 - d. Systemic lupus erythematosus
 - e. Dermatomyositis
 - f. Sjögren's syndrome
 - g. Serum sickness
 - h. Drug hypersensitivity—*diphenylhydantoin, hydralazine, allopurinol, primidone, gold, carbamazepine, etc.*
 - i. Angioimmunoblastic lymphadenopathy
 - j. Primary biliary cirrhosis
 - k. Graft-versus-host disease
 - l. Silicone-associated
 - m. Autoimmune lymphoproliferative syndrome
 - n. IgG4-related disease
 - o. Immune reconstitution inflammatory syndrome (IRIS)
3. Malignant diseases
 - a. Hematologic—*Hodgkin's disease, non-Hodgkin's lymphomas, acute or chronic lymphocytic leukemia, hairy cell leukemia, malignant histiocytosis, amyloidosis*
 - b. Metastatic—*from numerous primary sites*
4. Lipid storage diseases—*Gaucher's, Niemann-Pick, Fabry, Tangier*
5. Endocrine diseases—*hyperthyroidism*
6. Other disorders
 - a. Castleman's disease (*giant lymph node hyperplasia*)
 - b. Sarcoidosis
 - c. Dermatopathic lymphadenitis
 - d. Lymphomatoid granulomatosis
 - e. Histiocytic necrotizing lymphadenitis (*Kikuchi's disease*)
 - f. Sinus histiocytosis with massive lymphadenopathy (*Rosai-Dorfman disease*)
 - g. Mucocutaneous lymph node syndrome (*Kawasaki's disease*)
 - h. Histiocytosis X
 - i. Familial Mediterranean fever
 - g. Severe hypertriglyceridemia
 - k. Vascular transformation of sinuses
 - l. Inflammatory pseudotumor of lymph node
 - m. Congestive heart failure

Abbreviations: CMV, cytomegalovirus; EBV, Epstein-Barr virus.

with cervical adenopathy and a history of tobacco use. Localized or regional adenopathy implies involvement of a single anatomic area. Generalized adenopathy has been defined as involvement of three or more noncontiguous lymph node areas. Many of the causes of lymphadenopathy (Table 79-1) can produce localized or generalized adenopathy, so this distinction is of limited utility in the differential diagnosis. Nevertheless, generalized lymphadenopathy is frequently associated with nonmalignant disorders such as infectious

mononucleosis (Epstein-Barr virus [EBV] or cytomegalovirus [CMV]), toxoplasmosis, AIDS, other viral infections, systemic lupus erythematosus (SLE), and mixed connective tissue disease. Acute and chronic lymphocytic leukemias and malignant lymphomas also produce generalized adenopathy in adults.

The site of localized or regional adenopathy may provide a useful clue about the cause. Occipital adenopathy often reflects an infection of the scalp, and preauricular adenopathy accompanies conjunctival infections and cat-scratch disease. The most frequent site of regional adenopathy is the neck, and most of the causes are benign—upper respiratory infections, oral and dental lesions, infectious mononucleosis, or other viral illnesses. The chief malignant causes include metastatic cancer from head and neck, breast, lung, and thyroid primaries. Enlargement of supraclavicular and scalene nodes is always abnormal. Because these nodes drain regions of the lung and retroperitoneal space, they can reflect lymphomas, other cancers, or infectious processes arising in these areas. Virchow's node is an enlarged left supraclavicular node infiltrated with metastatic cancer from a gastrointestinal primary. Metastases to supraclavicular nodes also occur from lung, breast, testis, or ovarian cancers. Tuberculosis, sarcoidosis, and toxoplasmosis are nonneoplastic causes of supraclavicular adenopathy. Axillary adenopathy is usually due to injuries or localized infections of the ipsilateral upper extremity. Malignant causes include melanoma or lymphoma and, in women, breast cancer. Inguinal lymphadenopathy is usually secondary to infections or trauma of the lower extremities and may accompany sexually transmitted diseases such as lymphogranuloma venereum, primary syphilis, genital herpes, or chancroid. These nodes may also be involved by lymphomas and metastatic cancer from primary lesions of the rectum, genitalia, or lower extremities (melanoma).

The size and texture of the lymph node(s) and the presence of pain are useful parameters in evaluating a patient with lymphadenopathy. Nodes <1.0 cm² in area (1.0 cm \times 1.0 cm or less) are almost always secondary to benign, nonspecific reactive causes. In one retrospective analysis of younger patients (9–25 years) who had a lymph node biopsy, a maximum diameter of >2 cm served as one discriminant for predicting that the biopsy would reveal malignant or granulomatous disease. Another study showed that a lymph node size of 2.25 cm² (1.5 cm \times 1.5 cm) was the best size limit for distinguishing malignant or granulomatous lymphadenopathy from other causes of lymphadenopathy. Patients with node(s) ≤ 1.0 cm² should be observed after excluding infectious mononucleosis and/or toxoplasmosis unless there are symptoms and signs of an underlying systemic illness.

The texture of lymph nodes may be described as soft, firm, rubbery, hard, discrete, matted, tender, movable, or fixed. Tenderness is found when the capsule is stretched during rapid enlargement, usually secondary to an inflammatory process. Some malignant diseases such as acute leukemia may produce rapid enlargement and pain in the nodes. Nodes involved by lymphoma tend to be large, discrete, symmetric, rubbery, firm, mobile, and nontender. Nodes containing metastatic cancer are often hard, nontender, and nonmovable because of fixation to surrounding tissues. The coexistence of splenomegaly in the patient with lymphadenopathy implies a systemic illness such as infectious mononucleosis, lymphoma, acute or chronic leukemia, SLE, sarcoidosis, toxoplasmosis, cat-scratch disease, or other less common hematologic disorders. The patient's story should provide helpful clues about the underlying systemic illness.

Nonsuperficial presentations (thoracic or abdominal) of adenopathy are usually detected as the result of a symptom-directed diagnostic workup. Thoracic adenopathy may be detected by routine chest radiography or during the workup for superficial adenopathy. It may also be found because the patient complains of a cough or wheezing from airway compression; hoarseness from recurrent laryngeal nerve involvement; dysphagia from esophageal compression; or swelling of the neck, face, or arms secondary to compression of the superior vena cava or subclavian vein. The differential

diagnosis of mediastinal and hilar adenopathy includes primary lung disorders and systemic illnesses that characteristically involve mediastinal or hilar nodes. In the young, mediastinal adenopathy is associated with infectious mononucleosis and sarcoidosis. In endemic regions, histoplasmosis can cause unilateral paratracheal lymph node involvement that mimics lymphoma. Tuberculosis can also cause unilateral adenopathy. In older patients, the differential diagnosis includes primary lung cancer (especially among smokers), lymphomas, metastatic carcinoma (usually lung), tuberculosis, fungal infection, and sarcoidosis.

Enlarged intraabdominal or retroperitoneal nodes are usually malignant. Although tuberculosis may present as mesenteric lymphadenitis, these masses usually contain lymphomas or, in young men, germ cell tumors.

LABORATORY INVESTIGATION

The laboratory investigation of patients with lymphadenopathy must be tailored to elucidate the etiology suspected from the patient's history and physical findings. One study from a family practice clinic evaluated 249 younger patients with "enlarged lymph nodes, not infected" or "lymphadenitis." No laboratory studies were obtained in 51%. When studies were performed, the most common were a complete blood count (CBC) (33%), throat culture (16%), chest x-ray (12%), or monospot test (10%). Only eight patients (3%) had a node biopsy, and half of those were normal or reactive. The CBC can provide useful data for the diagnosis of acute or chronic leukemias, EBV or CMV mononucleosis, lymphoma with a leukemic component, pyogenic infections, or immune cytopenias in illnesses such as SLE. Serologic studies may demonstrate antibodies specific to components of EBV, CMV, HIV, and other viruses; *Toxoplasma gondii*; *Brucella*; and so on. If SLE is suspected, antinuclear and anti-DNA antibody studies are warranted.

The chest x-ray is usually negative, but the presence of a pulmonary infiltrate or mediastinal lymphadenopathy would suggest tuberculosis, histoplasmosis, sarcoidosis, lymphoma, primary lung cancer, or metastatic cancer and demands further investigation.

A variety of imaging techniques (computed tomography [CT], magnetic resonance imaging [MRI], ultrasound, color Doppler ultrasonography) have been used to differentiate benign from malignant lymph nodes, especially in patients with head and neck cancer. CT and MRI are comparably accurate (65–90%) in the diagnosis of metastases to cervical lymph nodes. Ultrasonography has been used to determine the long axis, short axis, and a ratio of long to short (L/S) axis in cervical nodes. An L/S ratio of <2.0 has a sensitivity and a specificity of 95% for distinguishing benign and malignant nodes in patients with head and neck cancer. This ratio has greater specificity and sensitivity than palpation or measurement of either the long or the short axis alone.

The indications for lymph node biopsy are imprecise, yet it is a valuable diagnostic tool. The decision to biopsy may be made early in a patient's evaluation or delayed for up to 2 weeks. Prompt biopsy should occur if the patient's history and physical findings suggest a malignancy; examples include a solitary, hard, nontender cervical node in an older patient who is a chronic user of tobacco; supraclavicular adenopathy; and solitary or generalized adenopathy that is firm, movable, and suggestive of lymphoma. If a primary head and neck cancer is suspected as the basis of a solitary, hard cervical node, then a careful ENT examination should be performed. Any mucosal lesion that is suspicious for a primary neoplastic process should be biopsied first. If no mucosal lesion is detected, an excisional biopsy of the largest node should be performed. Fine-needle aspiration should not be performed as the first diagnostic procedure. Most diagnoses require more tissue than such aspiration can provide, and it often delays a definitive diagnosis. Fine-needle aspiration should be reserved for thyroid nodules and for confirmation of relapse in patients whose primary diagnosis is known. If the primary physician is uncertain about whether to proceed to biopsy, consultation with a hematologist or medical oncologist should

be helpful. In primary care practices, <5% of lymphadenopathy patients will require a biopsy. That percentage will be considerably larger in referral practices, i.e., hematology, oncology, or ENT.

Two groups have reported algorithms that they claim will identify more precisely those lymphadenopathy patients who should have a biopsy. Both reports were retrospective analyses in referral practices. The first study involved patients 9–25 years of age who had a node biopsy performed. Three variables were identified that predicted those young patients with peripheral lymphadenopathy who should undergo biopsy; lymph node size >2 cm in diameter and abnormal chest x-ray had positive predictive values, whereas recent ENT symptoms had negative predictive values. The second study evaluated 220 lymphadenopathy patients in a hematology unit and identified five variables (lymph node size, location [supraclavicular or nonsupraclavicular], age [>40 years or <40 years], texture [nonhard or hard], and tenderness) that were used in a mathematical model to identify patients requiring a biopsy. Positive predictive value was found for age >40 years, supraclavicular location, node size >2.25 cm², hard texture, and lack of pain or tenderness. Negative predictive value was evident for age <40 years, node size <1.0 cm², nonhard texture, and tender or painful nodes. Ninety-one percent of those who required biopsy were correctly classified by this model. Because both of these studies were retrospective analyses and one was limited to young patients, it is not known how useful these models would be if applied prospectively in a primary care setting.

Most lymphadenopathy patients do not require a biopsy, and at least half require no laboratory studies. If the patient's history and physical findings point to a benign cause for lymphadenopathy, careful follow-up at a 2- to 4-week interval can be used. The patient should be instructed to return for reevaluation if there is an increase in the size of the nodes. Antibiotics are not indicated for lymphadenopathy unless strong evidence of a bacterial infection is present. Glucocorticoids should not be used to treat lymphadenopathy because their lympholytic effect obscures some diagnoses (lymphoma, leukemia, Castleman's disease), and they contribute to delayed healing or activation of underlying infections. An exception to this statement is the life-threatening pharyngeal obstruction by enlarged lymphoid tissue in Waldeyer's ring that is occasionally seen in infectious mononucleosis.

SPLENOMEGALY

STRUCTURE AND FUNCTION OF THE SPLEEN

The spleen is a reticuloendothelial organ that has its embryologic origin in the dorsal mesogastrium at about 5 weeks of gestation. It arises in a series of hillocks, migrates to its normal adult location in the left upper quadrant (LUQ), and is attached to the stomach via the gastrosplenic ligament and to the kidney via the lienorenal ligament. When the hillocks fail to unify into a single tissue mass, accessory spleens may develop in around 20% of persons. The function of the spleen has been elusive. Galen believed it was the source of "black bile" or melancholia, and the word *hypochondria* (literally, beneath the ribs) and the idiom "to vent one's spleen" attest to the beliefs that the spleen had an important influence on the psyche and emotions. In humans, its normal physiologic roles seem to be the following:

1. Maintenance of quality control over erythrocytes in the red pulp by removal of senescent and defective red blood cells. The spleen accomplishes this function through a unique organization of its parenchyma and vasculature (Fig. 79-1).
2. Synthesis of antibodies in the white pulp.
3. The removal of antibody-coated bacteria and antibody-coated blood cells from the circulation.

An increase in these normal functions may result in splenomegaly. The spleen is composed of *red pulp* and *white pulp*, which are Malpighi's terms for the red blood-filled sinuses and reticuloendothelial

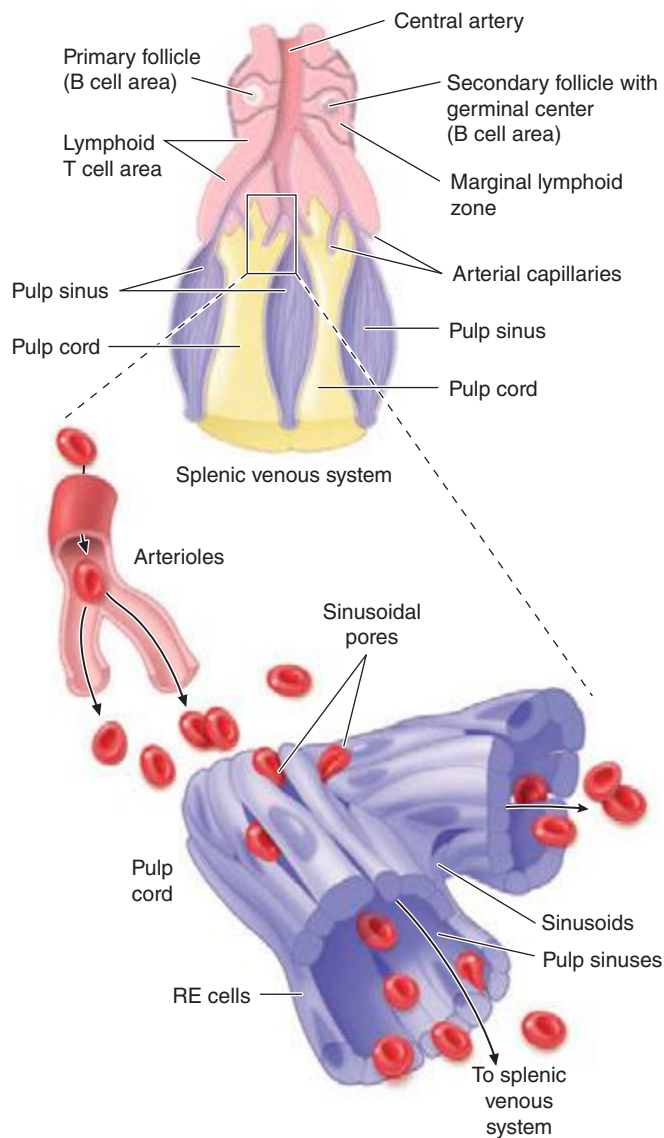


FIGURE 79-1 Schematic spleen structure. The spleen comprises many units of red and white pulp centered around small branches of the splenic artery, called *central arteries*. White pulp is lymphoid in nature and contains B cell follicles, a marginal zone around the follicles, and T cell–rich areas sheathing arterioles. The red pulp areas include pulp sinuses and pulp cords. The cords are dead ends. In order to regain access to the circulation, red blood cells must traverse tiny openings in the sinusoidal lining. Stiff, damaged, or old red cells cannot enter the sinuses. RE, reticuloendothelial. (Bottom portion of figure from RS Hillman, KA Ault: *Hematology in Clinical Practice*, 4th ed. New York, McGraw-Hill, 2005.)

cell–lined cords and the white lymphoid follicles arrayed within the red pulp matrix. The spleen is in the portal circulation. The reason for this is unknown but may relate to the fact that lower blood pressure allows less rapid flow and minimizes damage to normal erythrocytes. Blood flows into the spleen at a rate of about 150 mL/min through the splenic artery, which ultimately ramifies into central arterioles. Some blood goes from the arterioles to capillaries and then to splenic veins and out of the spleen, but the majority of blood from central arterioles flows into the macrophage-lined sinuses and cords. The blood entering the sinuses reenters the circulation through the splenic venules, but the blood entering the cords is subjected to an inspection of sorts. To return to the circulation, the blood cells in the cords must squeeze through slits in the cord lining to enter the sinuses that lead to the venules. Old and damaged erythrocytes are less deformable and are retained in the cords, where they are destroyed and their components

recycled. Red cell–inclusion bodies such as parasites (Chaps. 248 and 250e), nuclear residua (Howell–Jolly bodies, see Fig. 77-6), or denatured hemoglobin (Heinz bodies) are pinched off in the process of passing through the slits, a process called *pitting*. The culling of dead and damaged cells and the pitting of cells with inclusions appear to occur without significant delay because the blood transit time through the spleen is only slightly slower than in other organs.

The spleen is also capable of assisting the host in adapting to its hostile environment. It has at least three adaptive functions: (1) clearance of bacteria and particulates from the blood, (2) the generation of immune responses to certain pathogens, and (3) the generation of cellular components of the blood under circumstances in which the marrow is unable to meet the needs (i.e., extramedullary hematopoiesis). The latter adaptation is a recapitulation of the blood-forming function the spleen plays during gestation. In some animals, the spleen also serves a role in the vascular adaptation to stress because it stores red blood cells (often hemoconcentrated to higher hematocrits than normal) under normal circumstances and contracts under the influence of β -adrenergic stimulation to provide the animal with an autotransfusion and improved oxygen-carrying capacity. However, the normal human spleen does not sequester or store red blood cells and does not contract in response to sympathetic stimuli. The normal human spleen contains approximately one-third of the total body platelets and a significant number of margined neutrophils. These sequestered cells are available when needed to respond to bleeding or infection.

APPROACH TO THE PATIENT: Splénomegaly

CLINICAL ASSESSMENT

The most common *symptoms* produced by diseases involving the spleen are pain and a heavy sensation in the LUQ. Massive splenomegaly may cause early satiety. Pain may result from acute swelling of the spleen with stretching of the capsule, infarction, or inflammation of the capsule. For many years, it was believed that splenic infarction was clinically silent, which, at times, is true. However, Soma Weiss, in his classic 1942 report of the self-observations by a Harvard medical student on the clinical course of subacute bacterial endocarditis, documented that severe LUQ and pleuritic chest pain may accompany thromboembolic occlusion of splenic blood flow. Vascular occlusion, with infarction and pain, is commonly seen in children with sickle cell crises. Rupture of the spleen, from either trauma or infiltrative disease that breaks the capsule, may result in intraperitoneal bleeding, shock, and death. The rupture itself may be painless.

A palpable spleen is the major *physical sign* produced by diseases affecting the spleen and suggests enlargement of the organ. The normal spleen weighs <250 g, decreases in size with age, normally lies entirely within the rib cage, has a maximum cephalocaudad diameter of 13 cm by ultrasonography or maximum length of 12 cm and/or width of 7 cm by radionuclide scan, and is usually not palpable. However, a palpable spleen was found in 3% of 2200 asymptomatic, male, freshman college students. Follow-up at 3 years revealed that 30% of those students still had a palpable spleen without any increase in disease prevalence. Ten-year follow-up found no evidence for lymphoid malignancies. Furthermore, in some tropical countries (e.g., New Guinea), the incidence of splenomegaly may reach 60%. Thus, the presence of a palpable spleen does not always equate with presence of disease. Even when disease is present, splenomegaly may not reflect the primary disease but rather a reaction to it. For example, in patients with Hodgkin's disease, only two-thirds of the palpable spleens show involvement by the cancer.

Physical examination of the spleen uses primarily the techniques of palpation and percussion. Inspection may reveal fullness in the LUQ that descends on inspiration, a finding associated with a

massively enlarged spleen. Auscultation may reveal a venous hum or friction rub.

Palpation can be accomplished by bimanual palpation, ballotment, and palpation from above (Middleton maneuver). For bimanual palpation, which is at least as reliable as the other techniques, the patient is supine with flexed knees. The examiner's left hand is placed on the lower rib cage and pulls the skin toward the costal margin, allowing the fingertips of the right hand to feel the tip of the spleen as it descends while the patient inspires slowly, smoothly, and deeply. Palpation is begun with the right hand in the left lower quadrant with gradual movement toward the left costal margin, thereby identifying the lower edge of a massively enlarged spleen. When the spleen tip is felt, the finding is recorded as centimeters below the left costal margin at some arbitrary point, i.e., 10–15 cm, from the midpoint of the umbilicus or the xiphisternal junction. This allows other examiners to compare findings or the initial examiner to determine changes in size over time. Bimanual palpation in the right lateral decubitus position adds nothing to the supine examination.

Percussion for splenic dullness is accomplished with any of three techniques described by Nixon, Castell, or Barkun:

1. **Nixon's method:** The patient is placed on the right side so that the spleen lies above the colon and stomach. Percussion begins at the lower level of pulmonary resonance in the posterior axillary line and proceeds diagonally along a perpendicular line toward the lower midanterior costal margin. The upper border of dullness is normally 6–8 cm above the costal margin. Dullness >8 cm in an adult is presumed to indicate splenic enlargement.
2. **Castell's method:** With the patient supine, percussion in the lowest intercostal space in the anterior axillary line (eighth or ninth) produces a resonant note if the spleen is normal in size. This is true during expiration or full inspiration. A dull percussion note on full inspiration suggests splenomegaly.
3. **Percussion of Traube's semilunar space:** The borders of Traube's space are the sixth rib superiorly, the left midaxillary line laterally, and the left costal margin inferiorly. The patient is supine with the left arm slightly abducted. During normal breathing, this space is percussed from medial to lateral margins, yielding a normal resonant sound. A dull percussion note suggests splenomegaly.

Studies comparing methods of percussion and palpation with a standard of ultrasonography or scintigraphy have revealed sensitivity of 56–71% for palpation and 59–82% for percussion. Reproducibility among examiners is better for palpation than percussion. Both techniques are less reliable in obese patients or patients who have just eaten. Thus, the physical examination techniques of palpation and percussion are imprecise at best. It has been suggested that the examiner perform percussion first and, if positive, proceed to palpation; if the spleen is palpable, then one can be reasonably confident that splenomegaly exists. However, not all LUQ masses are enlarged spleens; gastric or colon tumors and pancreatic or renal cysts or tumors can mimic splenomegaly.

The presence of an enlarged spleen can be more precisely determined, if necessary, by liver-spleen radionuclide scan, CT, MRI, or ultrasonography. The latter technique is the current procedure of choice for routine assessment of spleen size (normal = a maximum cephalocaudal diameter of 13 cm) because it has high sensitivity and specificity and is safe, noninvasive, quick, mobile, and less costly. Nuclear medicine scans are accurate, sensitive, and reliable but are costly, require greater time to generate data, and use immobile equipment. They have the advantage of demonstrating accessory splenic tissue. CT and MRI provide accurate determination of spleen size, but the equipment is immobile and the procedures are expensive. MRI appears to offer no advantage over CT. Changes in spleen structure such as mass lesions, infarcts, inhomogeneous infiltrates, and cysts are more readily assessed by CT, MRI, or ultrasonography. None of these techniques is very reliable in the detection of patchy infiltration (e.g., Hodgkin's disease).

DIFFERENTIAL DIAGNOSIS

Many of the diseases associated with splenomegaly are listed in **Table 79-2**. They are grouped according to the presumed basic mechanisms responsible for organ enlargement:

1. Hyperplasia or hypertrophy related to a particular splenic function such as reticuloendothelial hyperplasia (work hypertrophy) in diseases such as hereditary spherocytosis or thalassemia syndromes that require removal of large numbers of defective red blood cells; immune hyperplasia in response to systemic infection (infectious mononucleosis, subacute bacterial endocarditis) or to immunologic diseases (immune thrombocytopenia, SLE, Felt's syndrome).
2. Passive congestion due to decreased blood flow from the spleen in conditions that produce portal hypertension (cirrhosis, Budd-Chiari syndrome, congestive heart failure).
3. Infiltrative diseases of the spleen (lymphomas, metastatic cancer, amyloidosis, Gaucher's disease, myeloproliferative disorders with extramedullary hematopoiesis).

The differential diagnostic possibilities are much fewer when the spleen is "massively enlarged," palpable more than 8 cm below the left costal margin or has a drained weight of ≥ 1000 g (**Table 79-3**). The vast majority of such patients will have non-Hodgkin's lymphoma, chronic lymphocytic leukemia, hairy cell leukemia, chronic myeloid leukemia, myelofibrosis with myeloid metaplasia, or polycythemia vera.

LABORATORY ASSESSMENT

The major laboratory abnormalities accompanying splenomegaly are determined by the underlying systemic illness. Erythrocyte counts may be normal, decreased (thalassemia major syndromes, SLE, cirrhosis with portal hypertension), or increased (polycythemia vera). Granulocyte counts may be normal, decreased (Felt's syndrome, congestive splenomegaly, leukemias), or increased (infections or inflammatory disease, myeloproliferative disorders). Similarly, the platelet count may be normal, decreased when there is enhanced sequestration or destruction of platelets in an enlarged spleen (congestive splenomegaly, Gaucher's disease, immune thrombocytopenia), or increased in the myeloproliferative disorders such as polycythemia vera.

The CBC may reveal cytopenia of one or more blood cell types, which should suggest *hypersplenism*. This condition is characterized by splenomegaly, cytopenia(s), normal or hyperplastic bone marrow, and a response to splenectomy. The latter characteristic is less precise because reversal of cytopenia, particularly granulocytopenia, is sometimes not sustained after splenectomy. The cytopenias result from increased destruction of the cellular elements secondary to reduced flow of blood through enlarged and congested cords (congestive splenomegaly) or to immune-mediated mechanisms. In hypersplenism, various cell types usually have normal morphology on the peripheral blood smear, although the red cells may be spherocytic due to loss of surface area during their longer transit through the enlarged spleen. The increased marrow production of red cells should be reflected as an increased reticulocyte production index, although the value may be less than expected due to increased sequestration of reticulocytes in the spleen.

The need for additional laboratory studies is dictated by the differential diagnosis of the underlying illness of which splenomegaly is a manifestation.

SPLENECTOMY

Splenectomy is infrequently performed for diagnostic purposes, especially in the absence of clinical illness or other diagnostic tests that suggest underlying disease. More often, splenectomy is performed for symptom control in patients with massive splenomegaly, for disease control in patients with traumatic splenic rupture, or for correction of cytopenias in patients with hypersplenism or immune-mediated destruction of one or more cellular blood elements. Splenectomy is

TABLE 79-2 DISEASES ASSOCIATED WITH SPLENOMEGALY GROUPED BY PATHOGENIC MECHANISM**Enlargement Due to Increased Demand for Splenic Function**

Reticuloendothelial system hyperplasia (for removal of defective erythrocytes)	Leishmaniasis
Spherocytosis	Trypanosomiasis
Early sickle cell anemia	Ehrlichiosis
Ovalocytosis	Disordered immunoregulation
Thalassemia major	Rheumatoid arthritis (Felty's syndrome)
Hemoglobinopathies	Systemic lupus erythematosus
Paroxysmal nocturnal hemoglobinuria	Collagen vascular diseases
Pernicious anemia	Serum sickness
Immune hyperplasia	Immune hemolytic anemias
Response to infection (viral, bacterial, fungal, parasitic)	Immune thrombocytopenias
Infectious mononucleosis	Immune neutropenias
AIDS	Drug reactions
Viral hepatitis	Angioimmunoblastic lymphadenopathy
Cytomegalovirus	Sarcoidosis
Subacute bacterial endocarditis	Thyrotoxicosis (benign lymphoid hypertrophy)
Bacterial septicemia	Interleukin 2 therapy
Congenital syphilis	Extramedullary hematopoiesis
Splenic abscess	Myelofibrosis
Tuberculosis	Marrow damage by toxins, radiation, strontium
Histoplasmosis	Marrow infiltration by tumors, leukemias, Gaucher's disease
Malaria	

Enlargement Due to Abnormal Splenic or Portal Blood Flow

Cirrhosis	Splenic artery aneurysm
Hepatic vein obstruction	Hepatic schistosomiasis
Portal vein obstruction, intrahepatic or extrahepatic	Congestive heart failure
Cavernous transformation of the portal vein	Hepatic echinococcosis
Splenic vein obstruction	Portal hypertension (any cause including the above): "Banti's disease"

Infiltration of the Spleen

Intracellular or extracellular depositions	Hodgkin's disease
Amyloidosis	Myeloproliferative syndromes (e.g., polycythemia vera, essential thrombocytosis)
Gaucher's disease	Angiosarcomas
Niemann-Pick disease	Metastatic tumors (melanoma is most common)
Tangier disease	Eosinophilic granuloma
Hurler's syndrome and other mucopolysaccharidoses	Histiocytosis X
Hyperlipidemias	Hamartomas
Benign and malignant cellular infiltrations	Hemangiomas, fibromas, lymphangiomas
Leukemias (acute, chronic, lymphoid, myeloid, monocytic)	Splenic cysts
Lymphomas	

Unknown Etiology

Idiopathic splenomegaly	Iron-deficiency anemia
Berylliosis	

necessary for staging of patients with Hodgkin's disease only in those with clinical stage I or II disease in whom radiation therapy alone is contemplated as the treatment. Noninvasive staging of the spleen in Hodgkin's disease is not a sufficiently reliable basis for treatment decisions because one-third of normal-sized spleens will be involved with Hodgkin's disease and one-third of enlarged spleens will be tumor-free. The widespread use of systemic therapy to test all stages of Hodgkin's disease has made staging laparotomy with splenectomy

unnecessary. Although splenectomy in chronic myeloid leukemia (CML) does not affect the natural history of disease, removal of the massive spleen usually makes patients significantly more comfortable and simplifies their management by significantly reducing transfusion requirements. The improvements in therapy of CML have reduced the need for splenectomy for symptom control. Splenectomy is an effective secondary or tertiary treatment for two chronic B cell leukemias, hairy cell leukemia and prolymphocytic leukemia, and for the very rare splenic mantle cell or marginal zone lymphoma. Splenectomy in these diseases may be associated with significant tumor regression in bone marrow and other sites of disease. Similar regressions of systemic disease have been noted after splenic irradiation in some types of lymphoid tumors, especially chronic lymphocytic leukemia and prolymphocytic leukemia. This has been termed the *abscopal effect*. Such systemic tumor responses to local therapy directed at the spleen suggest that some hormone or growth factor produced by the spleen may affect tumor cell proliferation, but this conjecture is not yet substantiated.

TABLE 79-3 DISEASES ASSOCIATED WITH MASSIVE SPLENOMEGALY*

Chronic myeloid leukemia	Gaucher's disease
Lymphomas	Chronic lymphocytic leukemia
Hairy cell leukemia	Sarcoidosis
Myelofibrosis with myeloid metaplasia	Autoimmune hemolytic anemia
Polycythemia vera	Diffuse splenic hemangiomas

*The spleen extends >8 cm below the left costal margin and/or weighs >1000 g.

A common therapeutic indication for splenectomy is traumatic or iatrogenic splenic rupture. In a fraction of patients with splenic rupture, peritoneal seeding of splenic fragments can lead to *splenosis*—the presence of multiple rests of spleen tissue not connected to the portal circulation. This ectopic spleen tissue may cause pain or gastrointestinal obstruction, as in endometriosis. A large number of hematologic, immunologic, and congestive causes of splenomegaly can lead to destruction of one or more cellular blood elements. In the majority of such cases, splenectomy can correct the cytopenias, particularly anemia and thrombocytopenia. In a large series of patients seen in two tertiary care centers, the indication for splenectomy was diagnostic in 10% of patients, therapeutic in 44%, staging for Hodgkin's disease in 20%, and incidental to another procedure in 26%. Perhaps the only contraindication to splenectomy is the presence of marrow failure, in which the enlarged spleen is the only source of hematopoietic tissue.

The absence of the spleen has minimal long-term effects on the hematologic profile. In the immediate postsplenectomy period, leukocytosis (up to 25,000/ μ L) and thrombocytosis (up to 1×10^6 / μ L) may develop, but within 2–3 weeks, blood cell counts and survival of each cell lineage are usually normal. The chronic manifestations of splenectomy are marked variation in size and shape of erythrocytes (anisocytosis, poikilocytosis) and the presence of Howell-Jolly bodies (nuclear remnants), Heinz bodies (denatured hemoglobin), basophilic stippling, and an occasional nucleated erythrocyte in the peripheral blood. When such erythrocyte abnormalities appear in a patient whose spleen has not been removed, one should suspect splenic infiltration by tumor that has interfered with its normal culling and pitting function.

The most serious consequence of splenectomy is increased susceptibility to bacterial infections, particularly those with capsules such as *Streptococcus pneumoniae*, *Haemophilus influenzae*, and some gram-negative enteric organisms. Patients under age 20 years are particularly susceptible to overwhelming sepsis with *S. pneumoniae*, and the overall actuarial risk of sepsis in patients who have had their spleens removed is about 7% in 10 years. The case-fatality rate for pneumococcal sepsis in splenectomized patients is 50–80%. About 25% of patients without spleens will develop a serious infection at some time in their life. The frequency is highest within the first 3 years after splenectomy. About 15% of the infections are polymicrobial, and lung, skin, and blood are the most common sites. No increased risk of viral infection has been noted in patients who have no spleen. The susceptibility to bacterial infections relates to the inability to remove opsonized bacteria from the bloodstream and a defect in making antibodies to T cell-independent antigens such as the polysaccharide components of bacterial capsules. Pneumococcal vaccine should be administered to all patients 2 weeks before elective splenectomy. The Advisory Committee on Immunization Practices recommends that these patients receive repeat vaccination 5 years after splenectomy. Efficacy has not been proven for this group, and the recommendation discounts the possibility that administration of the vaccine may actually lower the titer of specific pneumococcal antibodies. A more effective pneumococcal conjugate vaccine that involves T cells in the response is now available (Prevenar, 7-valent). The vaccine to *Neisseria meningitidis* should also be given to patients in whom elective splenectomy is planned. Although efficacy data for *H. influenzae* type b vaccine are not available for older children or adults, it may be given to patients who have had a splenectomy.

Splenectomized patients should be educated to consider any unexplained fever as a medical emergency. Prompt medical attention with evaluation and treatment of suspected bacteremia may be life-saving. Routine chemoprophylaxis with oral penicillin can result in the emergence of drug-resistant strains and is not recommended.

In addition to an increased susceptibility to bacterial infections, splenectomized patients are also more susceptible to the parasitic disease babesiosis. The splenectomized patient should avoid areas where the parasite *Babesia* is endemic (e.g., Cape Cod, MA).

Surgical removal of the spleen is an obvious cause of hyposplenism. Patients with sickle cell disease often suffer from autosplenectomy as a result of splenic destruction by the numerous infarcts associated with sickle cell crises during childhood. Indeed, the presence of a palpable spleen in a patient with sickle cell disease after age 5 suggests a coexisting hemoglobinopathy, e.g., thalassemia or hemoglobin C.

In addition, patients who receive splenic irradiation for a neoplastic or autoimmune disease are also functionally hyposplenic. The term *hyposplenism* is preferred to *asplenism* in referring to the physiologic consequences of splenectomy because asplenia is a rare, specific, and fatal congenital abnormality in which there is a failure of the left side of the coelomic cavity (which includes the splenic anlagen) to develop normally. Infants with asplenia have no spleens, but that is the least of their problems. The right side of the developing embryo is duplicated on the left so there is liver where the spleen should be, there are two right lungs, and the heart comprises two right atria and two right ventricles.

80

Disorders of Granulocytes and Monocytes

Steven M. Holland, John I. Gallin

Leukocytes, the major cells comprising inflammatory and immune responses, include neutrophils, T and B lymphocytes, natural killer (NK) cells, monocytes, eosinophils, and basophils. These cells have specific functions, such as antibody production by B lymphocytes or destruction of bacteria by neutrophils, but in no single infectious disease is the exact role of the cell types completely established. Thus, whereas neutrophils are classically thought to be critical to host defense against bacteria, they may also play important roles in defense against viral infections.

The blood delivers leukocytes to the various tissues from the bone marrow, where they are produced. Normal blood leukocyte counts are $4.3\text{--}10.8 \times 10^9$ /L, with neutrophils representing 45–74% of the cells, bands 0–4%, lymphocytes 16–45%, monocytes 4–10%, eosinophils 0–7%, and basophils 0–2%. Variation among individuals and among different ethnic groups can be substantial, with lower leukocyte numbers for certain African-American ethnic groups. The various leukocytes are derived from a common stem cell in the bone marrow. Three-fourths of the nucleated cells of bone marrow are committed to the production of leukocytes. Leukocyte maturation in the marrow is under the regulatory control of a number of different factors, known as colony-stimulating factors (CSFs) and interleukins (ILs). Because an alteration in the number and type of leukocytes is often associated with disease processes, total white blood cell (WBC) count (cells per μ L) and differential counts are informative. This chapter focuses on neutrophils, monocytes, and eosinophils. **Lymphocytes and basophils are discussed in Chaps. 372e and 376, respectively.**

NEUTROPHILS

MATURATION

Important events in neutrophil life are summarized in **Fig. 80-1**. In normal humans, neutrophils are produced only in the bone marrow. The minimum number of stem cells necessary to support hematopoiesis is estimated to be 400–500 at any one time. Human blood monocytes, tissue macrophages, and stromal cells produce CSFs, hormones required for the growth of monocytes and neutrophils in the bone marrow. The hematopoietic system not only produces enough neutrophils ($\sim 1.3 \times 10^{11}$ cells per 80-kg person per day) to carry out physiologic functions but also has a large reserve stored in the marrow, which can be mobilized in response to inflammation or infection. An increase in the number of blood neutrophils is called *neutrophilia*, and the presence of immature cells is termed a *shift to the left*. A decrease in the number of blood neutrophils is called *neutropenia*.

Neutrophils and monocytes evolve from pluripotent stem cells under the influence of cytokines and CSFs (**Fig. 80-2**). The proliferation phase through the metamyelocyte takes about 1 week, while the maturation phase from metamyelocyte to mature neutrophil takes

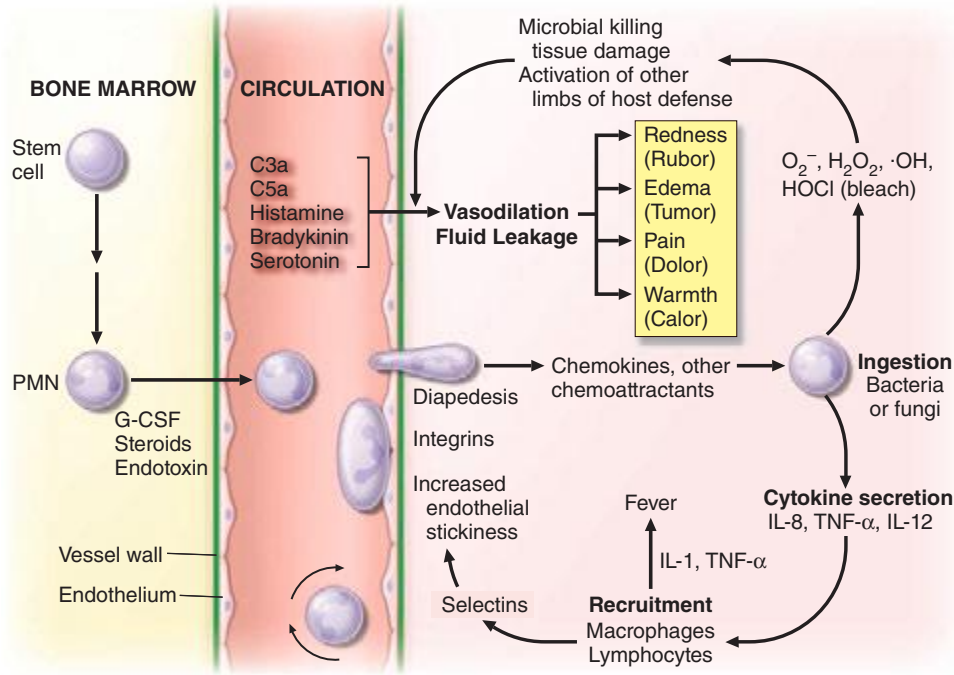


FIGURE 80-1 Schematic events in neutrophil production, recruitment, and inflammation. The four cardinal signs of inflammation (rubor, tumor, calor, dolor) are indicated, as are the interactions of neutrophils with other cells and cytokines. G-CSF, granulocyte colony-stimulating factor; IL, interleukin; PMN, polymorphonuclear leukocyte; TNF- α , tumor necrosis factor α .

Cell	Stage	Surface Markers ^a	Characteristics
	MYELOBLAST	CD33, CD13, CD15	Prominent nucleoli
	PROMYELOCYTE	CD33, CD13, CD15	Large cell Primary granules appear
	MYELOCYTE	CD33, CD13, CD15, CD14, CD11b	Secondary granules appear
	METAMYELOCYTE	CD33, CD13, CD15, CD14, CD11b	Kidney bean-shaped nucleus
	BAND FORM	CD33, CD13, CD15, CD14, CD11b, CD10, CD16	Condensed, band-shaped nucleus
	NEUTROPHIL	CD33, CD13, CD15, CD14, CD11b, CD10, CD16	Condensed, multilobed nucleus

^aCD = Cluster Determinant; ● Nucleolus; ● Primary granule; ● Secondary granule.

FIGURE 80-2 Stages of neutrophil development shown schematically. Granulocyte colony-stimulating factor (G-CSF) and granulocyte-macrophage colony-stimulating factor (GM-CSF) are critical to this process. Identifying cellular characteristics and specific cell-surface markers are listed for each maturational stage.

another week. The myeloblast is the first recognizable precursor cell and is followed by the *promyelocyte*. The promyelocyte evolves when the classic lysosomal granules, called the *primary*, or *azurophil*, *granules* are produced. The primary granules contain hydrolases, elastase, myeloperoxidase, cathepsin G, cationic proteins, and bactericidal/permeability-increasing protein, which is important for killing gram-negative bacteria. Azurophil granules also contain *defensins*, a family of cysteine-rich polypeptides with broad antimicrobial activity against bacteria, fungi, and certain enveloped viruses. The promyelocyte divides to produce the *myelocyte*, a cell responsible for the synthesis of the *specific*, or *secondary*, *granules*, which contain unique (specific) constituents such as lactoferrin, vitamin B₁₂-binding protein, membrane components of the reduced nicotinamide-adenine dinucleotide phosphate (NADPH) oxidase required for hydrogen peroxide production, histaminase, and receptors for certain chemoattractants and adherence-promoting factors (CR3) as well as receptors for the basement membrane component, laminin. The secondary granules do not contain acid hydrolases and therefore are not classic lysosomes. Packaging of secondary granule contents during myelopoiesis is controlled by CCAAT/enhancer binding protein- ϵ . Secondary granule contents are readily released extracellularly, and their mobilization is important in modulating inflammation. During the final stages of maturation, no cell division occurs, and the cell passes through the *metamyelocyte* stage and then to the *band neutrophil* with a sausage-shaped nucleus (Fig. 80-3). As the band cell matures, the nucleus assumes a lobulated configuration. The nucleus of neutrophils normally contains up to four segments (Fig. 80-4). Excessive segmentation (more than five nuclear lobes) may be a manifestation of folate or vitamin B₁₂ deficiency or the congenital neutropenia syndrome of warts, hypogammaglobulinemia, infections, and myelokathexis (WHIM) described below. The Pelger-Hüet anomaly (Fig. 80-5), an infrequent dominant benign inherited trait, results in neutrophils with distinctive bilobed nuclei that must be distinguished from band forms. Acquired bilobed nuclei, pseudo Pelger-Hüet anomaly, can occur with acute infections or in myelodysplastic syndromes. The physiologic role of the normal multilobed nucleus of neutrophils is unknown, but it may allow great deformation of neutrophils during migration into tissues at sites of inflammation.

In severe acute bacterial infection, prominent neutrophil cytoplasmic granules, called *toxic granulations*, are occasionally seen. Toxic granulations are immature or abnormally staining azurophil granules. Cytoplasmic inclusions, also called *Döhle bodies* (Fig. 80-3), can be seen during infection and are fragments of ribosome-rich endoplasmic reticulum. Large neutrophil vacuoles are often present in acute

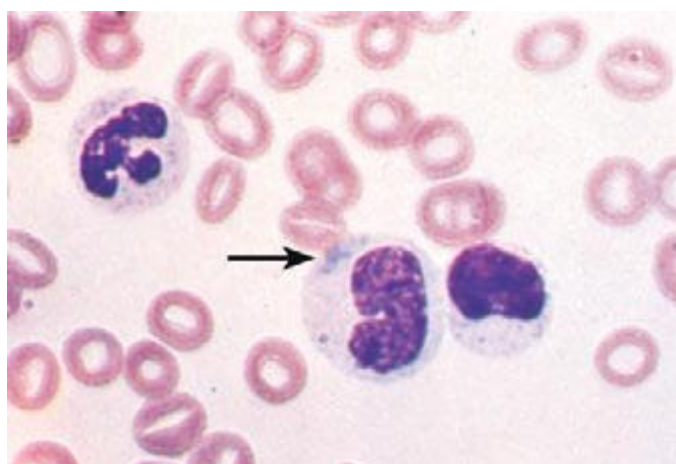


FIGURE 80-3 Neutrophil band with Döhle body. The neutrophil with a sausage-shaped nucleus in the center of the field is a band form. Döhle bodies are discrete, blue-staining, nongranular areas found in the periphery of the cytoplasm of the neutrophil in infections and other toxic states. They represent aggregates of rough endoplasmic reticulum.

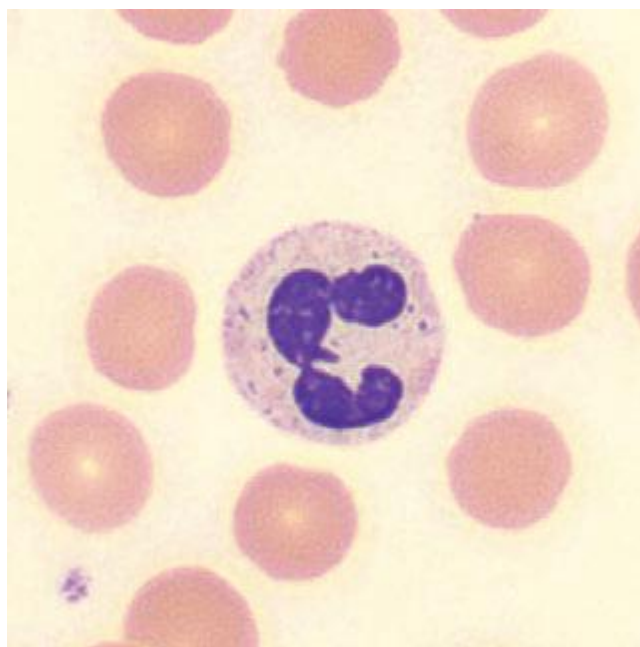


FIGURE 80-4 Normal granulocyte. The normal granulocyte has a segmented nucleus with heavy, clumped chromatin; fine neutrophilic granules are dispersed throughout the cytoplasm.

bacterial infection and probably represent pinocytosed (internalized) membrane.

Neutrophils are heterogeneous in function. Monoclonal antibodies have been developed that recognize only a subset of mature neutrophils. The meaning of neutrophil heterogeneity is not known.

The morphology of eosinophils and basophils is shown in Fig. 80-6.

MARROW RELEASE AND CIRCULATING COMPARTMENTS

Specific signals, including IL-1, tumor necrosis factor α (TNF- α), the CSFs, complement fragments, and chemokines, mobilize leukocytes from the bone marrow and deliver them to the blood in an unstimulated state. Under normal conditions, ~90% of the neutrophil pool is in the bone marrow, 2–3% in the circulation, and the remainder in the tissues (Fig. 80-7).

The circulating pool exists in two dynamic compartments: one freely flowing and one marginated. The freely flowing pool is about one-half the neutrophils in the basal state and is composed of those cells that are in the blood and not in contact with the endothelium. Marginated leukocytes are those that are in close physical contact with the

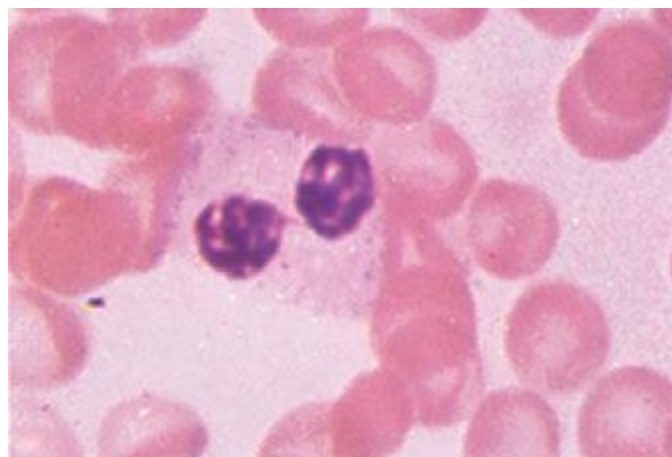


FIGURE 80-5 Pelger-Hüet anomaly. In this benign disorder, the majority of granulocytes are bilobed. The nucleus frequently has a spectacle-like, or "pince-nez," configuration.

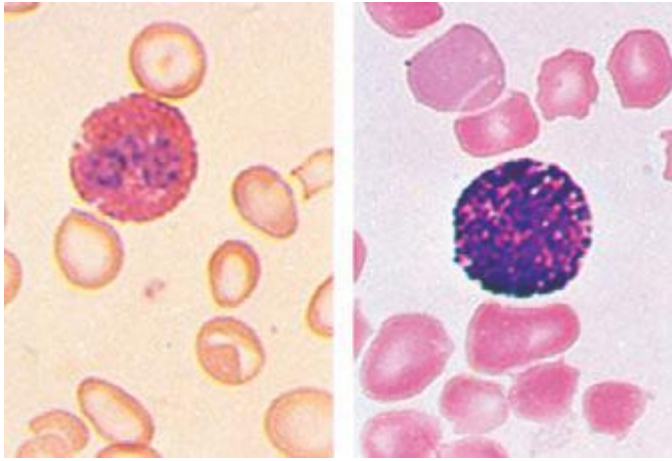


FIGURE 80-6 Normal eosinophil (left) and basophil (right). The eosinophil contains large, bright orange granules and usually a bilobed nucleus. The basophil contains large purple-black granules that fill the cell and obscure the nucleus.

endothelium (Fig. 80-8). In the pulmonary circulation, where an extensive capillary bed (~1000 capillaries per alveolus) exists, margination occurs because the capillaries are about the same size as a mature neutrophil. Therefore, neutrophil fluidity and deformability are necessary to make the transit through the pulmonary bed. Increased neutrophil rigidity and decreased deformability lead to augmented neutrophil trapping and margination in the lung. In contrast, in the systemic postcapillary venules, margination is mediated by the interaction of specific cell-surface molecules called *selectins*. Selectins are glycoproteins expressed on neutrophils and endothelial cells, among others, that cause a low-affinity interaction, resulting in “rolling” of the neutrophil along the endothelial surface. On neutrophils, the molecule L-selectin (cluster determinant [CD] 62L) binds to glycosylated proteins on endothelial cells (e.g., glycosylation-dependent cell adhesion molecule [GlyCAM1] and CD34). Glycoproteins on neutrophils, most importantly sialyl-Lewis^x (SLe^x, CD15s), are targets for binding of selectins expressed on endothelial cells (E-selectin [CD62E] and P-selectin [CD62P]) and

other leukocytes. In response to chemotactic stimuli from injured tissues (e.g., complement product C5a, leukotriene B₄, IL-8) or bacterial products (e.g., *N*-formylmethionylleucylphenylalanine [f-met-leu-phe]), neutrophil adhesiveness increases through mobilization of intracellular adhesion proteins stored in specific granules to the cell surface, and the cells “stick” to the endothelium through *integrins*. The integrins are leukocyte glycoproteins that exist as complexes of a common CD18 β chain with CD11a (LFA-1), CD11b (called Mac-1, CR3, or the C3bi receptor), and CD11c (called p150,95 or CR4). CD11a/CD18 and CD11b/CD18 bind to specific endothelial receptors (intercellular adhesion molecules [ICAM] 1 and 2).

On cell stimulation, L-selectin is shed from neutrophils, and E-selectin increases in the blood, presumably because it is shed from endothelial cells; receptors for chemoattractants and opsonins are mobilized; and the phagocytes orient toward the chemoattractant source in the extravascular space, increase their motile activity (chemokinesis), and migrate directionally (chemotaxis) into tissues. The process of migration into tissues is called *diapedesis* and involves the crawling of neutrophils between postcapillary endothelial cells that open junctions between adjacent cells to permit leukocyte passage. Diapedesis involves platelet/endothelial cell adhesion molecule (PECAM) 1 (CD31), which is expressed on both the emigrating leukocyte and the endothelial cells. The endothelial responses (increased blood flow from increased vasodilation and permeability) are mediated by anaphylatoxins (e.g., C3a and C5a) as well as vasodilators such as histamine, bradykinin, serotonin, nitric oxide, vascular endothelial growth factor (VEGF), and prostaglandins E and I. Cytokines regulate some of these processes (e.g., TNF-α induction of VEGF, interferon [IFN] γ inhibition of prostaglandin E).

In the healthy adult, most neutrophils leave the body by migration through the mucous membrane of the gastrointestinal tract. Normally, neutrophils spend a short time in the circulation (half-life, 6–7 h). Senescent neutrophils are cleared from the circulation by macrophages in the lung and spleen. Once in the tissues, neutrophils release enzymes, such as collagenase and elastase, which may help establish abscess cavities. Neutrophils ingest pathogenic materials that have been opsonized by IgG and C3b. Fibronectin and the tetrapeptide tuftsin also facilitate phagocytosis.

With phagocytosis comes a burst of oxygen consumption and activation of the hexose-monophosphate shunt. A membrane-associated NADPH oxidase, consisting of membrane and cytosolic components, is assembled and catalyzes the univalent reduction of oxygen to superoxide anion, which is then converted by superoxide dismutase to hydrogen peroxide and other toxic oxygen products (e.g., hydroxyl radical). Hydrogen peroxide + chloride + neutrophil myeloperoxidase generate hypochlorous acid (bleach), hypochlorite, and chlorine. These products oxidize and halogenate microorganisms and tumor cells and, when uncontrolled, can damage host tissue. Strongly cationic proteins, defensins, elastase, cathepsins, and probably nitric oxide also participate in microbial killing. Lactoferrin chelates iron, an important growth factor for microorganisms, especially fungi. Other enzymes, such as lysozyme and acid proteases, help digest microbial debris. After 1–4 days in tissues, neutrophils die. The apoptosis of neutrophils is also cytokine-regulated; granulocyte colony-stimulating factor (G-CSF) and IFN-γ prolong their life span. Under certain conditions, such as in delayed-type hypersensitivity, monocyte accumulation occurs within 6–12 h of initiation of inflammation. Neutrophils, monocytes, microorganisms in various states of digestion, and altered local tissue cells make up the inflammatory exudate, pus. Myeloperoxidase confers the characteristic green color to pus and may participate in turning off the inflammatory process by inactivating chemoattractants and immobilizing phagocytic cells.

Neutrophils respond to certain cytokines (IFN-γ, granulocyte-macrophage colony-stimulating factor [GM-CSF], IL-8) and produce cytokines and chemotactic signals (TNF-α, IL-8, macrophage inflammatory protein [MIP] 1) that modulate the inflammatory response. In the presence of fibrinogen, f-met-leu-phe or leukotriene B₄ induces IL-8 production by neutrophils, providing autocrine amplification of inflammation. *Chemokines* (chemoattractant cytokines) are small proteins

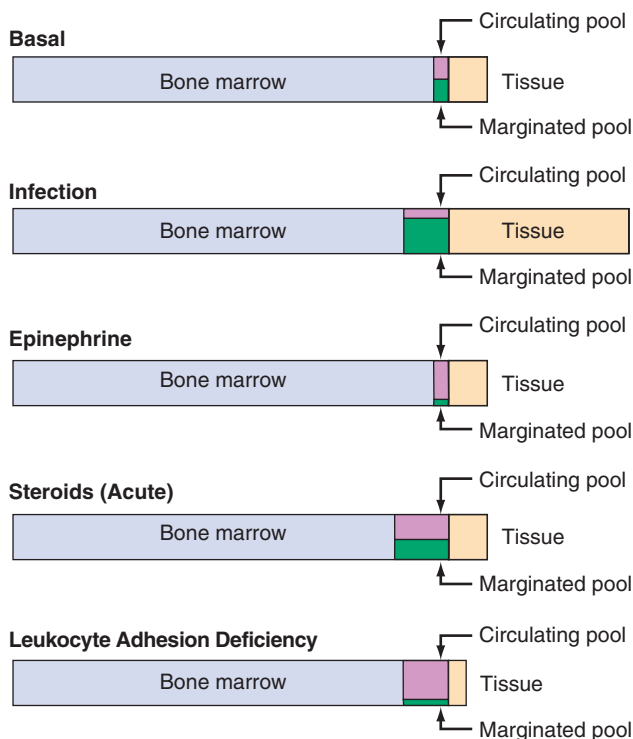


FIGURE 80-7 Schematic neutrophil distribution and kinetics between the different anatomic and functional pools.

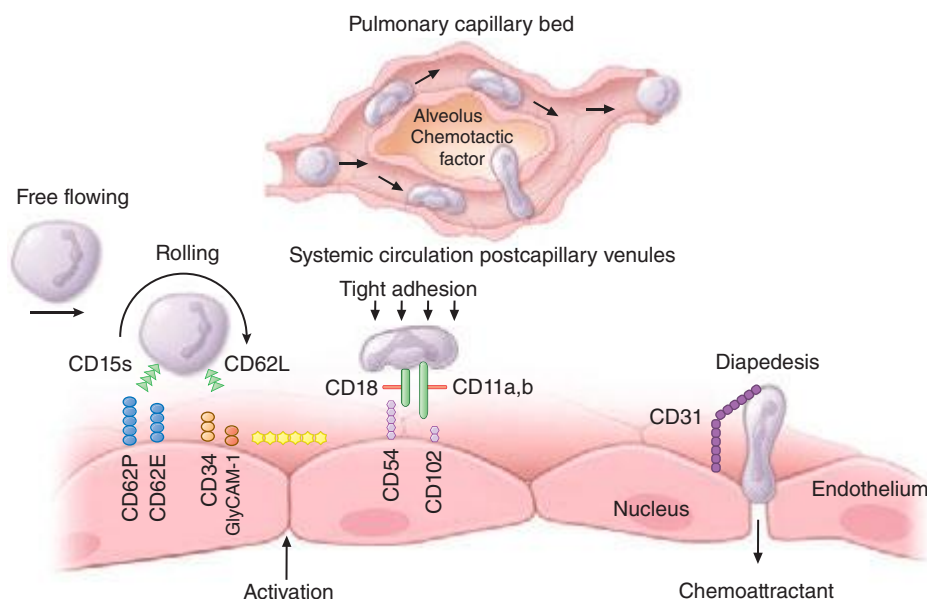


FIGURE 80-8 Neutrophil travel through the pulmonary capillaries is dependent on neutrophil deformability. Neutrophil rigidity (e.g., caused by C5a) enhances pulmonary trapping and response to pulmonary pathogens in a way that is not so dependent on cell-surface receptors. Intraalveolar chemotactic factors, such as those caused by certain bacteria (e.g., *Streptococcus pneumoniae*), lead to diapedesis of neutrophils from the pulmonary capillaries into the alveolar space. Neutrophil interaction with the endothelium of the systemic postcapillary venules is dependent on molecules of attachment. The neutrophil “rolls” along the endothelium using selectins: neutrophil CD15s (sialyl-Lewis^x) binds to CD62E (E-selectin) and CD62P (P-selectin) on endothelial cells; CD62L (L-selectin) on neutrophils binds to CD34 and other molecules (e.g., GlyCAM-1) expressed on endothelium. Chemokines or other activation factors stimulate integrin-mediated “tight adhesion”: CD11a/CD18 (LFA-1) and CD11b/CD18 (Mac-1, CR3) bind to CD54 (ICAM-1) and CD102 (ICAM-2) on the endothelium. Diapedesis occurs between endothelial cells: CD31 (PECAM-1) expressed by the emigrating neutrophil interacts with CD31 expressed at the endothelial cell-cell junction. CD, cluster determinant; GlyCAM, glycosylation-dependent cell adhesion molecule; ICAM, intercellular adhesion molecule; PECAM, platelet/endothelial cell adhesion molecule.

produced by many different cell types, including endothelial cells, fibroblasts, epithelial cells, neutrophils, and monocytes, that regulate neutrophil, monocyte, eosinophil, and lymphocyte recruitment and activation. Chemokines transduce their signals through heterotrimeric G protein-linked receptors that have seven cell membrane-spanning domains, the same type of cell-surface receptor that mediates the response to the classic chemoattractants f-met-leu-phe and C5a. Four major groups of chemokines are recognized based on the cysteine structure near the N terminus: C, CC, CXC, and CXXXX. The CXC cytokines such as IL-8 mainly attract neutrophils; CC chemokines such as MIP-1 attract lymphocytes, monocytes, eosinophils, and basophils; the C chemokine lymphotactin is T cell tropic; the CXXXX chemokine fractalkine attracts neutrophils, monocytes, and T cells. These molecules and their receptors not only regulate the trafficking and activation of inflammatory cells, but specific chemokine receptors also serve as co-receptors for HIV infection (Chap. 226) and have a role in other viral infections such as West Nile infection and atherogenesis.

NEUTROPHIL ABNORMALITIES

Defects in the neutrophil life cycle can lead to dysfunction and compromised host defenses. Inflammation is often depressed, and the clinical result is often recurrent, severe bacterial and fungal infections. Aphthous ulcers of mucous membranes (gray ulcers without pus) and gingivitis and periodontal disease suggest a phagocytic cell disorder. Patients with congenital phagocyte defects can have infections within the first few days of life. Skin, ear, upper and lower respiratory tract, and bone infections are common. Sepsis and meningitis are rare. In some disorders, the frequency of infection is variable, and patients can go for months or even years without major infection. Aggressive management of these congenital diseases has extended the life span of patients well beyond 30 years.

Neutropenia The consequences of absent neutrophils are dramatic. Susceptibility to infectious diseases increases sharply when neutrophil counts fall below 1000 cells/ μ L. When the absolute neutrophil count

(ANC; band forms and mature neutrophils combined) falls to <500 cells/ μ L, control of endogenous microbial flora (e.g., mouth, gut) is impaired; when the ANC is <200/ μ L, the local inflammatory process is absent. Neutropenia can be due to depressed production, increased peripheral destruction, or excessive peripheral pooling. A falling neutrophil count or a significant decrease in the number of neutrophils below steady-state levels, together with a failure to increase neutrophil counts in the setting of infection or other challenge, requires investigation. Acute neutropenia, such as that caused by cancer chemotherapy, is more likely to be associated with increased risk of infection than neutropenia of long duration (months to years) that reverses in response to infection or carefully controlled administration of endotoxin (see “Laboratory Diagnosis and Management,” below).

Some causes of inherited and acquired neutropenia are listed in Table 80-1. The most common neutropenias are iatrogenic, resulting from the use of cytotoxic or immunosuppressive therapies for malignancy or control of autoimmune disorders. These drugs cause neutropenia because they result in decreased production of rapidly growing progenitor (stem) cells of the marrow. Certain antibiotics such as chloramphenicol, trimethoprim-sulfamethoxazole, flucytosine, vidarabine, and the antiretroviral drug zidovudine may cause neutropenia by inhibiting proliferation of myeloid precursors. Azathioprine and 6-mercaptopurine are metabolized by the enzyme thiopurine methyltransferase (TMPT), hypofunctional polymorphisms in which are found in 11% of whites and can lead to accumulation of 6-thioguanine and profound marrow toxicity. The marrow suppression is generally dose-related and dependent on continued administration of the drug. Cessation of the offending agent and recombinant human G-CSF usually reverse these forms of neutropenia.

Another important mechanism for iatrogenic neutropenia is the effect of drugs that serve as immune haptens and sensitize neutrophils or neutrophil precursors to immune-mediated peripheral destruction. This form of drug-induced neutropenia can be seen within 7 days of exposure to the drug; with previous drug exposure, resulting in preexisting antibodies, neutropenia may occur a few hours after

TABLE 80-1 CAUSES OF NEUTROPENIA**Decreased Production**

Drug-induced—alkylating agents (nitrogen mustard, busulfan, chlorambucil, cyclophosphamide); antimetabolites (methotrexate, 6-mercaptopurine, 5-fluorouracil); noncytotoxic agents (antibiotics [chloramphenicol, penicillins, sulfonamides], phenothiazines, tranquilizers [meprobamate], anticonvulsants [carbamazepine], antipsychotics [clozapine], certain diuretics, anti-inflammatory agents, antithyroid drugs, many others)

Hematologic diseases—idiopathic, cyclic neutropenia, Chédiak-Higashi syndrome, aplastic anemia, infantile genetic disorders (see text)

Tumor invasion, myelofibrosis

Nutritional deficiency—vitamin B₁₂, folate (especially alcoholics)

Infection—tuberculosis, typhoid fever, brucellosis, tularemia, measles, infectious mononucleosis, malaria, viral hepatitis, leishmaniasis, AIDS

Peripheral Destruction

Antineutrophil antibodies and/or splenic or lung trapping

Autoimmune disorders—Felty's syndrome, rheumatoid arthritis, lupus erythematosus

Drugs as haptens—aminopyrine, α-methyl dopa, phenylbutazone, mercurial diuretics, some phenothiazines

Granulomatosis with polyangiitis (Wegener's)

Peripheral Pooling (Transient Neutropenia)

Overwhelming bacterial infection (acute endotoxemia)

Hemodialysis

Cardiopulmonary bypass

administration of the drug. Although any drug can cause this form of neutropenia, the most frequent causes are commonly used antibiotics, such as sulfa-containing compounds, penicillins, and cephalosporins. Fever and eosinophilia may also be associated with drug reactions, but often these signs are not present. Drug-induced neutropenia can be severe, but discontinuation of the sensitizing drug is sufficient for recovery, which is usually seen within 5–7 days and is complete by 10 days. Readministration of the sensitizing drug should be avoided, because abrupt neutropenia will often result. For this reason, diagnostic challenge should be avoided.

Autoimmune neutropenias caused by circulating antineutrophil antibodies are another form of acquired neutropenia that results in increased destruction of neutrophils. Acquired neutropenia may also be seen with viral infections, including infection with HIV. Acquired neutropenia may be cyclic in nature, occurring at intervals of several weeks. Acquired cyclic or stable neutropenia may be associated with an expansion of large granular lymphocytes (LGLs), which may be T cells, NK cells, or NK-like cells. Patients with large granular lymphocytosis may have moderate blood and bone marrow lymphocytosis, neutropenia, polyclonal hypergammaglobulinemia, splenomegaly, rheumatoid arthritis, and absence of lymphadenopathy. Such patients may have a chronic and relatively stable course. Recurrent bacterial infections are frequent. Benign and malignant forms of this syndrome occur. In some patients, a spontaneous regression has occurred even after 11 years, suggesting an immunoregulatory defect as the basis for at least one form of the disorder. Glucocorticoids, cyclosporine, and methotrexate are commonly used to manage these cytopenias.

Hereditary Neutropenias Hereditary neutropenias are rare and may manifest in early childhood as a profound constant neutropenia or agranulocytosis. Congenital forms of neutropenia include Kostmann's syndrome (neutrophil count <100/μL), which is often fatal and due to mutations in the antiapoptosis gene *HAX-1*; severe chronic neutropenia (neutrophil count of 300–1500/μL) due to mutations in neutrophil elastase (*ELANE*); hereditary cyclic neutropenia, or, more appropriately, cyclic hematopoiesis, also due to mutations in neutrophil elastase (*ELANE*); the cartilage-hair hypoplasia syndrome due to mutations in the mitochondrial RNA-processing endoribonuclease *RMRP*; Shwachman-Diamond syndrome associated with pancreatic insufficiency due to mutations in the Shwachman-Bodian-Diamond syndrome gene *SBDS*; the WHIM (warts, hypogammaglobulinemia,

infections, myelokathexis [retention of WBCs in the marrow]) syndrome, characterized by neutrophil hypersegmentation and bone marrow myeloid arrest due to mutations in the chemokine receptor *CXCR4*; and neutropenias associated with other immune defects, such as X-linked agammaglobulinemia, Wiskott-Aldrich syndrome, and CD40 ligand deficiency. Mutations in the G-CSF receptor can develop in severe congenital neutropenia and are linked to leukemia. Absence of both myeloid and lymphoid cells is seen in reticular dysgenesis, due to mutations in the nuclear genome-encoded mitochondrial enzyme adenylate kinase-2 (*AK2*).

Maternal factors can be associated with neutropenia in the newborn. Transplacental transfer of IgG directed against antigens on fetal neutrophils can result in peripheral destruction. Drugs (e.g., thiazides) ingested during pregnancy can cause neutropenia in the newborn by either depressed production or peripheral destruction.

In Felty's syndrome—the triad of rheumatoid arthritis, splenomegaly, and neutropenia (**Chap. 380**)—spleen-produced antibodies can shorten neutrophil life span, while large granular lymphocytes can attack marrow neutrophil precursors. Splenectomy may increase the neutrophil count in Felty's syndrome and lower serum neutrophil-binding IgG. Some Felty's syndrome patients also have neutropenia associated with an increased number of LGLs. Splenomegaly with peripheral trapping and destruction of neutrophils is also seen in lysosomal storage diseases and in portal hypertension.

Neutrophilia Neutrophilia results from increased neutrophil production, increased marrow release, or defective margination (**Table 80-2**). The most important acute cause of neutrophilia is infection. Neutrophilia from acute infection represents both increased production and increased marrow release. Increased production is also associated with chronic inflammation and certain myeloproliferative diseases. Increased marrow release and mobilization of the marginated leukocyte pool are induced by glucocorticoids. Release of epinephrine, as with vigorous exercise, excitement, or stress, will demarginate neutrophils in the spleen and lungs and double the neutrophil count in minutes. Cigarette smoking can elevate neutrophil counts above the normal range. Leukocytosis with cell counts of 10,000–25,000/μL occurs in response to infection and other forms of acute inflammation and results from both release of the marginated pool and mobilization of marrow reserves. Persistent neutrophilia with cell counts of ≥30,000–50,000/μL is called a *leukemoid reaction*, a term

TABLE 80-2 CAUSES OF NEUTROPHILIA**Increased Production**

Idiopathic

Drug-induced—glucocorticoids, G-CSF

Infection—bacterial, fungal, sometimes viral

Inflammation—thermal injury, tissue necrosis, myocardial and pulmonary infarction, hypersensitivity states, collagen vascular diseases

Myeloproliferative diseases—myelocytic leukemia, myeloid metaplasia, polycythemia vera

Increased Marrow Release

Glucocorticoids

Acute infection (endotoxin)

Inflammation—thermal injury

Decreased or Defective Margination

Drugs—epinephrine, glucocorticoids, nonsteroidal anti-inflammatory agents

Stress, excitement, vigorous exercise

Leukocyte adhesion deficiency type 1 (CD18); leukocyte adhesion deficiency type 2 (selectin ligand, CD15s); leukocyte adhesion deficiency type 3 (FERMT3)

Miscellaneous

Metabolic disorders—ketoacidosis, acute renal failure, eclampsia, acute poisoning

Drugs—lithium

Other—metastatic carcinoma, acute hemorrhage or hemolysis

Abbreviation: G-CSF, granulocyte colony-stimulating factor.

TABLE 80-3 TYPES OF GRANULOCYTE AND MONOCYTE DISORDERS

Function	Cause of Indicated Dysfunction		
	Drug-Induced	Acquired	Inherited
Adherence-aggregation	Aspirin, colchicine, alcohol, glucocorticoids, ibuprofen, piroxicam	Neonatal state, hemodialysis	Leukocyte adhesion deficiency types 1, 2, and 3
Deformability		Leukemia, neonatal state, diabetes mellitus, immature neutrophils	
Chemokinesis-chemotaxis	Glucocorticoids (high dose), auranofin, colchicine (weak effect), phenylbutazone, naproxen, indomethacin, interleukin 2	Thermal injury, malignancy, malnutrition, periodontal disease, neonatal state, systemic lupus erythematosus, rheumatoid arthritis, diabetes mellitus, sepsis, influenza virus infection, herpes simplex virus infection, acrodermatitis enteropathica, AIDS	Chédiak-Higashi syndrome, neutrophil-specific granule deficiency, hyper IgE–recurrent infection (Job’s syndrome (in some patients), Down’s syndrome, α -mannosidase deficiency, leukocyte adhesion deficiencies, Wiskott-Aldrich syndrome
Microbicidal activity	Colchicine, cyclophosphamide, glucocorticoids (high dose), TNF- α -blocking antibodies	Leukemia, aplastic anemia, certain neutropenias, tuftsin deficiency, thermal injury, sepsis, neonatal state, diabetes mellitus, malnutrition, AIDS	Chédiak-Higashi syndrome, neutrophil-specific granule deficiency, chronic granulomatous disease, defects in IFN γ /IL-12 axis

Abbreviations: IFN γ , interferon γ ; IL, interleukin; TNF- α , tumor necrosis factor alpha.

often used to distinguish this degree of neutrophilia from leukemia. In a leukemoid reaction, the circulating neutrophils are usually mature and not clonally derived.

Abnormal Neutrophil Function Inherited and acquired abnormalities of phagocyte function are listed in [Table 80-3](#). The resulting diseases are best considered in terms of the functional defects of adherence, chemotaxis, and microbicidal activity. The distinguishing features of the important inherited disorders of phagocyte function are shown in [Table 80-4](#).

DISORDERS OF ADHESION Three main types of leukocyte adhesion deficiency (LAD) have been described. All are autosomal recessive and result in the inability of neutrophils to exit the circulation to sites of infection, leading to leukocytosis and increased susceptibility to infection ([Fig. 80-8](#)). Patients with LAD 1 have mutations in *CD18*, the common component of the integrins LFA-1, Mac-1, and p150,95, leading to a defect in tight adhesion between neutrophils and the endothelium. The heterodimer formed by CD18/CD11b (Mac-1) is also the receptor for the complement-derived opsonin C3bi (CR3). The *CD18* gene is located on distal chromosome 21q. The severity of the defect determines the severity of clinical disease. Complete lack of expression of the leukocyte integrins results in a severe phenotype in which inflammatory stimuli do not increase the expression of leukocyte integrins on neutrophils or activated T and B cells. Neutrophils (and monocytes) from patients with LAD 1 adhere poorly to endothelial cells and protein-coated surfaces and exhibit defective spreading, aggregation, and chemotaxis. Patients with LAD 1 have recurrent bacterial infections involving the skin, oral and genital mucosa, and respiratory and intestinal tracts; persistent leukocytosis (resting neutrophil counts of 15,000–20,000/ μ L) because cells do not marginate; and, in severe cases, a history of delayed separation of the umbilical stump. Infections, especially of the skin, may become necrotic with progressively enlarging borders, slow healing, and development of dysplastic scars. The most common bacteria are *Staphylococcus aureus* and enteric gram-negative bacteria. LAD 2 is caused by an abnormality of fucosylation of SLe^x (CD15s), the ligand on neutrophils that interacts with selectins on endothelial cells and is responsible for neutrophil rolling along the endothelium. Infection susceptibility in LAD 2 appears to be less severe than in LAD 1. LAD 2 is also known as *congenital disorder of glycosylation IIc* (CDGIIc) due to mutation in a GDP-fucose transporter (*SLC35C1*). LAD 3 is characterized by infection susceptibility, leukocytosis, and petechial hemorrhage due to impaired integrin activation caused by mutations in the gene *FERMT3*.

DISORDERS OF NEUTROPHIL GRANULES The most common neutrophil defect is myeloperoxidase deficiency, a primary granule defect inherited as an autosomal recessive trait; the incidence is ~1 in 2000 persons. Isolated myeloperoxidase deficiency is not associated with clinically compromised defenses, presumably because other defense systems

such as hydrogen peroxide generation are amplified. Microbicidal activity of neutrophils is delayed but not absent. Myeloperoxidase deficiency may make other acquired host defense defects more serious, and patients with myeloperoxidase deficiency and diabetes are more susceptible to *Candida* infections. An acquired form of myeloperoxidase deficiency occurs in myelomonocytic leukemia and acute myeloid leukemia.

Chédiak-Higashi syndrome (CHS) is a rare disease with autosomal recessive inheritance due to defects in the lysosomal transport protein *LYST*, encoded by the gene *CHS1* at 1q42. This protein is required for normal packaging and disbursement of granules. Neutrophils (and all cells containing lysosomes) from patients with CHS characteristically have large granules ([Fig. 80-9](#)), making it a systemic disease. Patients with CHS have nystagmus, partial oculocutaneous albinism, and an increased number of infections resulting from many bacterial agents. Some CHS patients develop an “accelerated phase” in childhood with a hemophagocytic syndrome and an aggressive lymphoma requiring bone marrow transplantation. CHS neutrophils and monocytes have impaired chemotaxis and abnormal rates of microbial killing due to slow rates of fusion of the lysosomal granules with phagosomes. NK cell function is also impaired. CHS patients may develop a severe disabling peripheral neuropathy in adulthood that can lead to bed confinement.

Specific granule deficiency is a rare autosomal recessive disease in which the production of secondary granules and their contents, as well as the primary granule component defensins, is defective. The defect in killing leads to severe bacterial infections. One type of specific granule deficiency is due to a mutation in the CCAAT/enhancer binding protein- ϵ , a regulator of expression of granule components. A dominant mutation in *C/EBP- ϵ* has also been described.

CHRONIC GRANULOMATOUS DISEASE Chronic granulomatous disease (CGD) is a group of disorders of granulocyte and monocyte oxidative metabolism. Although CGD is rare, with an incidence of ~1 in 200,000 individuals, it is an important model of defective neutrophil oxidative metabolism. In about two-thirds of patients, CGD is inherited as an X-linked recessive trait; 30% of patients inherit the disease in an autosomal recessive pattern. Mutations in the genes for the five proteins that assemble at the plasma membrane account for all patients with CGD. Two proteins (a 91-kDa protein, abnormal in X-linked CGD, and a 22-kDa protein, absent in one form of autosomal recessive CGD) form the heterodimer cytochrome b-558 in the plasma membrane. Three other proteins (40, 47, and 67 kDa, abnormal in the other autosomal recessive forms of CGD) are cytoplasmic in origin and interact with the cytochrome after cell activation to form NADPH oxidase, required for hydrogen peroxide production. Leukocytes from patients with CGD have severely diminished hydrogen peroxide production. The genes involved in each of the defects have been cloned and sequenced and the chromosome locations identified. Patients with CGD characteristically have increased numbers of infections

TABLE 80-4 INHERITED DISORDERS OF PHAGOCYTE FUNCTION: DIFFERENTIAL FEATURES

Clinical Manifestations	Cellular or Molecular Defects	Diagnosis
Chronic Granulomatous Diseases (70% X-Linked, 30% Autosomal Recessive)		
Severe infections of skin, ears, lungs, liver, and bone with catalase-positive microorganisms such as <i>Staphylococcus aureus</i> , <i>Burkholderia cepacia</i> complex, <i>Aspergillus</i> spp., <i>Chromobacterium violaceum</i> ; often hard to culture organism; excessive inflammation with granulomas, frequent lymph node suppuration; granulomas can obstruct GI or GU tracts; gingivitis, aphthous ulcers, seborrheic dermatitis	No respiratory burst due to the lack of one of five NADPH oxidase subunits in neutrophils, monocytes, and eosinophils	DHR or NBT test; no superoxide and H ₂ O ₂ production by neutrophils; immunoblot for NADPH oxidase components; genetic detection
Chédiak-Higashi Syndrome (Autosomal Recessive)		
Recurrent pyogenic infections, especially with <i>S. aureus</i> ; many patients get lymphoma-like illness during adolescence; periodontal disease; partial oculocutaneous albinism, nystagmus, progressive peripheral neuropathy, mental retardation in some patients	Reduced chemotaxis and phagolysosome fusion, increased respiratory burst activity, defective egress from marrow, abnormal skin window; defect in <i>CHS1</i>	Giant primary granules in neutrophils and other granule-bearing cells (Wright's stain); genetic detection
Specific Granule Deficiency (Autosomal Recessive and Dominant)		
Recurrent infections of skin, ears, and sinopulmonary tract; delayed wound healing; decreased inflammation; bleeding diathesis	Abnormal chemotaxis, impaired respiratory burst and bacterial killing, failure to upregulate chemotactic and adhesion receptors with stimulation, defect in transcription of granule proteins; defect in <i>CEBPE</i>	Lack of secondary (specific) granules in neutrophils (Wright's stain), no neutrophil-specific granule contents (i.e., lactoferrin), no defensins, platelet α granule abnormality; genetic detection
Myeloperoxidase Deficiency (Autosomal Recessive)		
Clinically normal except in patients with underlying disease such as diabetes mellitus; then candidiasis or other fungal infections	No myeloperoxidase due to pre- and posttranslational defects in myeloperoxidase deficiency	No peroxidase in neutrophils; genetic detection
Leukocyte Adhesion Deficiency		
Type 1: Delayed separation of umbilical cord, sustained neutrophilia, recurrent infections of skin and mucosa, gingivitis, periodontal disease	Impaired phagocyte adherence, aggregation, spreading, chemotaxis, phagocytosis of C3bi-coated particles; defective production of CD18 subunit common to leukocyte integrins	Reduced phagocyte surface expression of the CD18-containing integrins with monoclonal antibodies against LFA-1 (CD18/CD11a), Mac-1 or CR3 (CD18/CD11b), p150,95 (CD18/CD11c); genetic detection
Type 2: Mental retardation, short stature, Bombay (hh) blood phenotype, recurrent infections, neutrophilia	Impaired phagocyte rolling along endothelium; due to defects in fucose transporter	Reduced phagocyte surface expression of Sialyl-Lewis ^x , with monoclonal antibodies against CD15s; genetic detection
Type 3: Petechial hemorrhage, recurrent infections	Impaired signaling for integrin activation resulting in impaired adhesion due to mutation in <i>FERMT3</i>	Reduced signaling for adhesion through integrins; genetic detection
Phagocyte Activation Defects (X-Linked and Autosomal Recessive)		
NEMO deficiency: mild hypohidrotic ectodermal dysplasia; broad-based immune defect: pyogenic and encapsulated bacteria, viruses, <i>Pneumocystis</i> , mycobacteria; X-linked	Impaired phagocyte activation by IL-1, IL-18, TLR, CD40L, TNF- α leading to problems with inflammation and antibody production	Poor in vitro response to endotoxin; impaired NF- κ B activation; genetic detection
IRAK4 and MyD88 deficiency: susceptibility to pyogenic bacteria such as staphylococci, streptococci, clostridia; resistant to <i>Candida</i> ; autosomal recessive	Impaired phagocyte activation by endotoxin through TLR and other pathways; TNF- α signaling preserved	Poor in vitro response to endotoxin; lack of NF- κ B activation by endotoxin; genetic detection
Hyper IgE–Recurrent Infection Syndrome (Autosomal Dominant) (Job's Syndrome)		
Eczematoid or pruritic dermatitis, "cold" skin abscesses, recurrent pneumonias with <i>S. aureus</i> with bronchopleural fistulae and cyst formation, mild eosinophilia, mucocutaneous candidiasis, characteristic facies, restrictive lung disease, scoliosis, delayed primary dental deciduation	Reduced chemotaxis in some patients, reduced memory T and B cells; mutation in <i>STAT3</i>	Somatic and immune features involving lungs, skeleton, and immune system; serum IgE >2000 IU/mL; genetic testing
DOCK8 deficiency (autosomal recessive), severe eczema, atopic dermatitis, cutaneous abscesses, HSV, HPV, and molluscum infections, severe allergies, cancer	Impaired T cell proliferation to mitogens; mutation in <i>DOCK8</i>	Severe allergies, viral infections, high IgE, eosinophilia, low IgM, progressive lymphopenia, genetic detection
Mycobacteria Susceptibility (Autosomal Dominant and Recessive Forms)		
Severe extrapulmonary or disseminated infections with bacille Calmette–Guérin (BCG), nontuberculous mycobacteria, salmonella, histoplasmosis, coccidioidomycosis, poor granuloma formation	Inability to kill intracellular organisms due to low IFN- γ production or response; mutations in IFN- γ receptors, IL-12 receptors, IL-12 p40, <i>STAT1</i> , <i>NEMO</i> , <i>ISG15</i> , <i>GATA2</i>	Abnormally low or very high levels of IFN- γ receptor 1; functional assays of cytokine production and response; genetic detection
GATA2 Deficiency (Autosomal Dominant)		
Persistent or disseminated warts, disseminated mycobacterial disease, low monocytes, NK cells, B cells; hypoplastic myelodysplasia, leukemia, cytogenetic abnormalities, pulmonary alveolar proteinosis	Impaired macrophage activity, cytopenias; mutations in <i>GATA2</i>	Profound circulating monocytopenia, NK and B cell cytopenias; genetic detection

Abbreviations: C/EBP ϵ , CCAAT/enhancer binding protein- ϵ ; DHR, dihydrorhodamine (oxidation test); DOCK8, dedicator of cytokinesis 8; GI, gastrointestinal; GU, genitourinary; HPV, human papilloma virus; HSV, herpes simplex virus; IFN, interferon; IL, interleukin; IRAK4, IL-1 receptor–associated kinase 4; LFA-1, leukocyte function–associated antigen 1; MyD88, myeloid differentiation primary response gene 88; NADPH, nicotinamide–adenine dinucleotide phosphate; NBT, nitroblue tetrazolium (dye test); NEMO, NF- κ B essential modulator; NF- κ B, nuclear factor- κ B; NK, natural killer; STAT1–3, signal transducer and activator of transcription 1–3; TLR, Toll-like receptor; TNF, tumor necrosis factor.

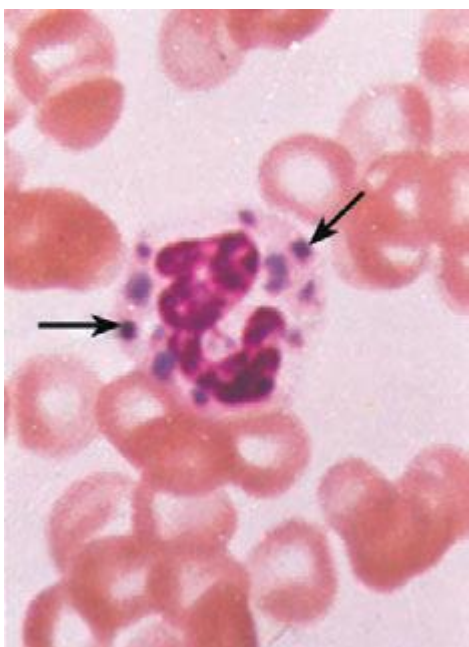


FIGURE 80-9 Chédiak-Higashi syndrome. The granulocytes contain huge cytoplasmic granules formed from aggregation and fusion of azurophilic and specific granules. Large abnormal granules are found in other granule-containing cells throughout the body.

due to catalase-positive microorganisms (organisms that destroy their own hydrogen peroxide) such as *S. aureus*, *Burkholderia cepacia*, and *Aspergillus* species. When patients with CGD become infected, they often have extensive inflammatory reactions, and lymph node suppuration is common despite the administration of appropriate antibiotics. Aphthous ulcers and chronic inflammation of the nares are often present. Granulomas are frequent and can obstruct the gastrointestinal or genitourinary tracts. The excessive inflammation is due to failure to downregulate inflammation, reflecting failure to inhibit the synthesis of, degradation of, or response to chemoattractants or residual antigens, leading to persistent neutrophil accumulation. Impaired killing of intracellular microorganisms by macrophages may lead to persistent cell-mediated immune activation and granuloma formation. Autoimmune complications such as immune thrombocytopenic purpura and juvenile rheumatoid arthritis are also increased in CGD. In addition, for unexplained reasons, discoid lupus is more common in X-linked carriers. Late complications, including nodular regenerative hyperplasia and portal hypertension, are increasingly recognized in long-term survivors of severe CGD.

DISORDERS OF PHAGOCYTE ACTIVATION Phagocytes depend on cell-surface stimulation to induce signals that evoke multiple levels of the inflammatory response, including cytokine synthesis, chemotaxis, and antigen presentation. Mutations affecting the major pathway that signals through NF- κ B have been noted in patients with a variety of infection susceptibility syndromes. If the defects are at a very late stage of signal transduction, in the protein critical for NF- κ B activation known as the NF- κ B essential modulator (NEMO), then affected males develop ectodermal dysplasia and severe immune deficiency with susceptibility to bacteria, fungi, mycobacteria, and viruses. If the defects in NF- κ B activation are closer to the cell-surface receptors, in the proteins transducing Toll-like receptor signals, IL-1 receptor–associated kinase 4 (IRAK4), and myeloid differentiation primary response gene 88 (MyD88), then children have a marked susceptibility to pyogenic infections early in life but develop resistance to infection later.

MONONUCLEAR PHAGOCYTES

The mononuclear phagocyte system is composed of monoblasts, promonocytes, and monocytes, in addition to the structurally diverse tissue macrophages that make up what was previously referred to as

the reticuloendothelial system. Macrophages are long-lived phagocytic cells capable of many of the functions of neutrophils. They are also secretory cells that participate in many immunologic and inflammatory processes distinct from neutrophils. Monocytes leave the circulation by diapedesis more slowly than neutrophils and have a half-life in the blood of 12–24 h.

After blood monocytes arrive in the tissues, they differentiate into macrophages (“big eaters”) with specialized functions suited for specific anatomic locations. Macrophages are particularly abundant in capillary walls of the lung, spleen, liver, and bone marrow, where they function to remove microorganisms and other noxious elements from the blood. Alveolar macrophages, liver Kupffer cells, splenic macrophages, peritoneal macrophages, bone marrow macrophages, lymphatic macrophages, brain microglial cells, and dendritic macrophages all have specialized functions. Macrophage-secreted products include lysozyme, neutral proteases, acid hydrolases, arginase, complement components, enzyme inhibitors (plasmin, α_2 -macroglobulin), binding proteins (transferrin, fibronectin, transcobalamin II), nucleosides, and cytokines (TNF- α ; IL-1, -8, -12, -18). IL-1 (Chaps. 23 and 372e) has many functions, including initiating fever in the hypothalamus, mobilizing leukocytes from the bone marrow, and activating lymphocytes and neutrophils. TNF- α is a pyrogen that duplicates many of the actions of IL-1 and plays an important role in the pathogenesis of gram-negative shock (Chap. 325). TNF- α stimulates production of hydrogen peroxide and related toxic oxygen species by macrophages and neutrophils. In addition, TNF- α induces catabolic changes that contribute to the profound wasting (cachexia) associated with many chronic diseases.

Other macrophage-secreted products include reactive oxygen and nitrogen metabolites, bioactive lipids (arachidonic acid metabolites and platelet-activating factors), chemokines, CSFs, and factors stimulating fibroblast and vessel proliferation. Macrophages help regulate the replication of lymphocytes and participate in the killing of tumors, viruses, and certain bacteria (*Mycobacterium tuberculosis* and *Listeria monocytogenes*). Macrophages are key effector cells in the elimination of intracellular microorganisms. Their ability to fuse to form giant cells that coalesce into granulomas in response to some inflammatory stimuli is important in the elimination of intracellular microbes and is under the control of IFN- γ . Nitric oxide induced by IFN- γ is an important effector against intracellular parasites, including tuberculosis and *Leishmania*.

Macrophages play an important role in the immune response (Chap. 372e). They process and present antigen to lymphocytes and secrete cytokines that modulate and direct lymphocyte development and function. Macrophages participate in autoimmune phenomena by removing immune complexes and other substances from the circulation. Polymorphisms in macrophage receptors for immunoglobulin (Fc γ R2) determine susceptibility to some infections and autoimmune diseases. In wound healing, they dispose of senescent cells, and they contribute to atheroma development. Macrophage elastase mediates development of emphysema from cigarette smoking.

DISORDERS OF THE MONONUCLEAR PHAGOCYTE SYSTEM

Many disorders of neutrophils extend to mononuclear phagocytes. Monocytosis is associated with tuberculosis, brucellosis, subacute bacterial endocarditis, Rocky Mountain spotted fever, malaria, and visceral leishmaniasis (kala azar). Monocytosis also occurs with malignancies, leukemias, myeloproliferative syndromes, hemolytic anemias, chronic idiopathic neutropenias, and granulomatous diseases such as sarcoidosis, regional enteritis, and some collagen vascular diseases. Patients with LAD, hyperimmunoglobulin E–recurrent infection (Job’s) syndrome, CHS, and CGD all have defects in the mononuclear phagocyte system.

Monocyte cytokine production or response is impaired in some patients with disseminated nontuberculous mycobacterial infection who are not infected with HIV. Genetic defects in the pathways regulated by IFN- γ and IL-12 lead to impaired killing of intracellular bacteria, mycobacteria, salmonellae, and certain viruses (Fig. 80-10).

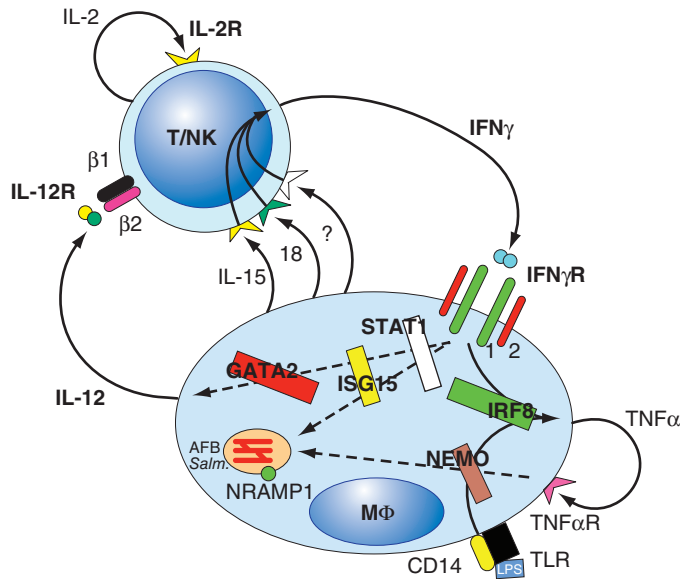


FIGURE 80-10 Lymphocyte-macrophage interactions underlying resistance to mycobacteria and other intracellular pathogens such as *Salmonella*, *Histoplasma*, and *Coccidioides*. Mycobacteria (and others) infect macrophages, leading to the production of IL-12, which activates T or NK cells through its receptor, leading to production of IL-2 and IFN- γ . IFN- γ acts through its receptor on macrophages to upregulate TNF- γ and IL-12 and kill intracellular pathogens. Other critical interacting molecules include signal transducer and activator of transcription 1 (STAT1), interferon regulatory factor 8 (IRF8), GATA2, and ISG15. Mutant forms of the cytokines and receptors shown in *bold type* have been found in severe cases of nontuberculous mycobacterial infection, salmonellosis and other intracellular pathogens. AFB, acid-fast bacilli; IFN, interferon; IL, interleukin; NEMO, nuclear factor- κ B essential modulator; NK, natural killer; TLR, Toll-like receptor; TNF, tumor necrosis factor.

Certain viral infections impair mononuclear phagocyte function. For example, influenza virus infection causes abnormal monocyte chemotaxis. Mononuclear phagocytes can be infected by HIV using CCR5, the chemokine receptor that acts as a co-receptor with CD4 for HIV. T lymphocytes produce IFN- γ , which induces FcR expression and phagocytosis and stimulates hydrogen peroxide production by mononuclear phagocytes and neutrophils. In certain diseases, such as AIDS, IFN- γ production may be deficient, whereas in other diseases, such as T cell lymphomas, excessive release of IFN- γ may be associated with erythrophagocytosis by splenic macrophages.

Autoinflammatory diseases are characterized by abnormal cytokine regulation, leading to excess inflammation in the absence of infection. These diseases can mimic infectious or immunodeficient syndromes. Gain-of-function mutations in the TNF- α receptor cause TNF- α receptor-associated periodic syndrome (TRAPS), which is characterized by recurrent fever in the absence of infection, due to persistent stimulation of the TNF- α receptor (Chap. 392). Diseases with abnormal IL-1 regulation leading to fever include familial Mediterranean fever due to mutations in *PYRIN*. Mutations in *cold-induced autoinflammatory syndrome 1* (*CIAS1*) lead to neonatal-onset multisystem autoinflammatory disease, familial cold urticaria, and Muckle-Wells syndrome. The syndrome of *pyoderma gangrenosum*, *acne*, and sterile *pyogenic arthritis* (PAPA syndrome) is caused by mutations in *PSTPIP1*. In contrast to these syndromes of overexpression of proinflammatory cytokines, blockade of TNF- α by the antagonists infliximab, adalimumab, certolizumab, golimumab, or etanercept has been associated with severe infections due to tuberculosis, nontuberculous mycobacteria, and fungi (Chap. 392).

Monocytopenia occurs with acute infections, with stress, and after treatment with glucocorticoids. Drugs that suppress neutrophil production in the bone marrow can cause monocytopenia. Persistent severe circulating monocytopenia is seen in GATA2 deficiency,

even though macrophages are found at the sites of inflammation. Monocytopenia also occurs in aplastic anemia, hairy cell leukemia, acute myeloid leukemia, and as a direct result of myelotoxic drugs.

EOSINOPHILS

Eosinophils and neutrophils share similar morphology, many lysosomal constituents, phagocytic capacity, and oxidative metabolism. Eosinophils express a specific chemoattractant receptor and respond to a specific chemokine, eotaxin, but little is known about their required role. Eosinophils are much longer lived than neutrophils, and unlike neutrophils, tissue eosinophils can recirculate. During most infections, eosinophils appear unimportant. However, in invasive helminthic infections, such as hookworm, schistosomiasis, strongyloidiasis, toxocarosis, trichinosis, filariasis, echinococcosis, and cysticercosis, the eosinophil plays a central role in host defense. Eosinophils are associated with bronchial asthma, cutaneous allergic reactions, and other hypersensitivity states.

The distinctive feature of the red-staining (Wright's stain) eosinophil granule is its crystalline core consisting of an arginine-rich protein (major basic protein) with histaminase activity, important in host defense against parasites. Eosinophil granules also contain a unique eosinophil peroxidase that catalyzes the oxidation of many substances by hydrogen peroxide and may facilitate killing of microorganisms.

Eosinophil peroxidase, in the presence of hydrogen peroxide and halide, initiates mast cell secretion in vitro and thereby promotes inflammation. Eosinophils contain cationic proteins, some of which bind to heparin and reduce its anticoagulant activity. Eosinophil-derived neurotoxin and eosinophil cationic protein are ribonucleases that can kill respiratory syncytial virus. Eosinophil cytoplasm contains Charcot-Leyden crystal protein, a hexagonal bipyramidal crystal first observed in a patient with leukemia and then in sputum of patients with asthma; this protein is lysophospholipase and may function to detoxify certain lysophospholipids.

Several factors enhance the eosinophil's function in host defense. T cell-derived factors enhance the ability of eosinophils to kill parasites. Mast cell-derived eosinophil chemotactic factor of anaphylaxis (ECF α) increases the number of eosinophil complement receptors and enhances eosinophil killing of parasites. Eosinophil CSFs (e.g., IL-5) produced by macrophages increase eosinophil production in the bone marrow and activate eosinophils to kill parasites.

EOSINOPHILIA

Eosinophilia is the presence of >500 eosinophils per μ L of blood and is common in many settings besides parasite infection. Significant tissue eosinophilia can occur without an elevated blood count. A common cause of eosinophilia is allergic reaction to drugs (iodides, aspirin, sulfonamides, nitrofurantoin, penicillins, and cephalosporins). Allergies such as hay fever, asthma, eczema, serum sickness, allergic vasculitis, and pemphigus are associated with eosinophilia. Eosinophilia also occurs in collagen vascular diseases (e.g., rheumatoid arthritis, eosinophilic fasciitis, allergic angitis, and periarteritis nodosa) and malignancies (e.g., Hodgkin's disease; mycosis fungoides; chronic myeloid leukemia; and cancer of the lung, stomach, pancreas, ovary, or uterus), as well as in Job's syndrome, DOCK8 deficiency (see below), and CGD. Eosinophilia is commonly present in helminthic infections. IL-5 is the dominant eosinophil growth factor. Therapeutic administration of the cytokines IL-2 or GM-CSF frequently leads to transient eosinophilia. The most dramatic hypereosinophilic syndromes are Loeffler's syndrome, tropical pulmonary eosinophilia, Loeffler's endocarditis, eosinophilic leukemia, and idiopathic hypereosinophilic syndrome (50,000–100,000/ μ L). IL-5 is the dominant eosinophil growth factor and can be specifically inhibited with the monoclonal antibody mepolizumab.

The idiopathic hypereosinophilic syndrome represents a heterogeneous group of disorders with the common feature of prolonged eosinophilia of unknown cause and organ system dysfunction, including the heart, central nervous system, kidneys, lungs, gastrointestinal tract, and skin. The bone marrow is involved in all affected individuals, but the

most severe complications involve the heart and central nervous system. Clinical manifestations and organ dysfunction are highly variable. Eosinophils are found in the involved tissues and likely cause tissue damage by local deposition of toxic eosinophil proteins such as eosinophil cationic protein and major basic protein. In the heart, the pathologic changes lead to thrombosis, endocardial fibrosis, and restrictive endomyocardialopathy. The damage to tissues in other organ systems is similar. Some cases are due to mutations involving the platelet-derived growth factor receptor, and these are extremely sensitive to the tyrosine kinase inhibitor imatinib. Glucocorticoids, hydroxyurea, and IFN- α each have been used successfully, as have therapeutic antibodies against IL-5. Cardiovascular complications are managed aggressively.

The *eosinophilia-myalgia syndrome* is a multisystem disease, with prominent cutaneous, hematologic, and visceral manifestations, that frequently evolves into a chronic course and can occasionally be fatal. The syndrome is characterized by eosinophilia (eosinophil count $>1000/\mu\text{L}$) and generalized disabling myalgias without other recognized causes. Eosinophilic fasciitis, pneumonitis, and myocarditis; neuropathy culminating in respiratory failure; and encephalopathy may occur. The disease is caused by ingesting contaminants in L-tryptophan-containing products. Eosinophils, lymphocytes, macrophages, and fibroblasts accumulate in the affected tissues, but their role in pathogenesis is unclear. Activation of eosinophils and fibroblasts and the deposition of eosinophil-derived toxic proteins in affected tissues may contribute. IL-5 and transforming growth factor β have been implicated as potential mediators. Treatment is withdrawal of products containing L-tryptophan and the administration of glucocorticoids. Most patients recover fully, remain stable, or show slow recovery, but the disease can be fatal in up to 5% of patients.

Eosinophilic neoplasms are discussed in Chapter 135e.

EOSINOPENIA

Eosinopenia occurs with stress, such as acute bacterial infection, and after treatment with glucocorticoids. The mechanism of eosinopenia of acute bacterial infection is unknown but is independent of endogenous glucocorticoids, because it occurs in animals after total adrenalectomy. There is no known adverse effect of eosinopenia.

HYPERIMMUNOGLOBULIN E—RECURRENT INFECTION SYNDROME

The hyperimmunoglobulin E—recurrent infection syndrome, or Job's syndrome, is a rare multisystem disease in which the immune and somatic systems are affected, including neutrophils, monocytes, T cells, B cells, and osteoclasts. Autosomal *dominant* mutations in signal transducer and activator of transcription 3 (STAT3) lead to inhibition of normal STAT signaling with broad and profound effects. Patients have characteristic facies with broad nose, kyphoscoliosis, and eczema. The primary teeth erupt normally but do not deciduate, often requiring extraction. Patients develop recurrent sinopulmonary and cutaneous infections that tend to be much less inflamed than appropriate for the degree of infection and have been referred to as “cold abscesses.” Characteristically, pneumonias cavitate, leading to pneumatoceles. Coronary artery aneurysms are common, as are cerebral demyelinated plaques that accumulate with age. Importantly, IL-17-producing T cells, which are thought responsible for protection against extracellular and mucosal infections, are profoundly reduced in Job's syndrome. Despite very high IgE levels, these patients do not have elevated levels of allergy. An important syndrome with clinical overlap with STAT3 deficiency is due to autosomal recessive defects in dedicator of cytokinesis 8 (DOCK8). In DOCK8 deficiency, IgE elevation is joined to severe allergy, viral susceptibility, and increased rates of cancer.

LABORATORY DIAGNOSIS AND MANAGEMENT

Initial studies of WBC and differential and often a bone marrow examination may be followed by assessment of bone marrow reserves (steroid challenge test), marginated circulating pool of cells (epinephrine challenge test), and marginating ability (endotoxin challenge test) (Fig. 80-7). In vivo assessment of inflammation is possible with a Rebuck

skin window test or an in vivo skin blister assay, which measures the ability of leukocytes and inflammatory mediators to accumulate locally in the skin. In vitro tests of phagocyte aggregation, adherence, chemotaxis, phagocytosis, degranulation, and microbicidal activity (for *S. aureus*) may help pinpoint cellular or humoral lesions. Deficiencies of oxidative metabolism are detected with either the nitroblue tetrazolium (NBT) dye test or the dihydrorhodamine (DHR) oxidation test. These tests are based on the ability of products of oxidative metabolism to alter the oxidation states of reporter molecules so that they can be detected microscopically (NBT) or by flow cytometry (DHR). Qualitative studies of superoxide and hydrogen peroxide production may further define neutrophil oxidative function.

Patients with leukopenias or leukocyte dysfunction often have delayed inflammatory responses. Therefore, clinical manifestations may be minimal despite overwhelming infection, and unusual infections must always be suspected. Early signs of infection demand prompt, aggressive culturing for microorganisms, use of antibiotics, and surgical drainage of abscesses. Prolonged courses of antibiotics are often required. In patients with CGD, prophylactic antibiotics (trimethoprim-sulfamethoxazole) and antifungals (itraconazole) markedly diminish the frequency of life-threatening infections. Glucocorticoids may relieve gastrointestinal or genitourinary tract obstruction by granulomas in patients with CGD. Although TNF- α -blocking agents may markedly relieve inflammatory bowel symptoms, extreme caution must be exercised in their use in CGD inflammatory bowel disease, because it profoundly increases these patients' already heightened susceptibility to infection. Recombinant human IFN- γ , which nonspecifically stimulates phagocytic cell function, reduces the frequency of infections in patients with CGD by 70% and reduces the severity of infection. This effect of IFN- γ in CGD is additive to the effect of prophylactic antibiotics. The recommended dose is 50 $\mu\text{g}/\text{m}^2$ subcutaneously three times weekly. IFN- γ has also been used successfully in the treatment of leprosy, nontuberculous mycobacteria, and visceral leishmaniasis.

Rigorous oral hygiene reduces but does not eliminate the discomfort of gingivitis, periodontal disease, and aphthous ulcers; chlorhexidine mouthwash and tooth brushing with a hydrogen peroxide-sodium bicarbonate paste help many patients. Oral antifungal agents (fluconazole, itraconazole, voriconazole, posaconazole) have reduced mucocutaneous candidiasis in patients with Job's syndrome. Androgens, glucocorticoids, lithium, and immunosuppressive therapy have been used to restore myelopoiesis in patients with neutropenia due to impaired production. Recombinant G-CSF is useful in the management of certain forms of neutropenia due to depressed neutrophil production, including those related to cancer chemotherapy. Patients with chronic neutropenia with evidence of a good bone marrow reserve need not receive prophylactic antibiotics. Patients with chronic or cyclic neutrophil counts $<500/\mu\text{L}$ may benefit from prophylactic antibiotics and G-CSF during periods of neutropenia. Oral trimethoprim-sulfamethoxazole (160/800 mg) twice daily can prevent infection. Increased numbers of fungal infections are not seen in patients with CGD on this regimen. Oral quinolones such as levofloxacin and ciprofloxacin are alternatives.

In the setting of cytotoxic chemotherapy with severe, persistent lymphocyte dysfunction, trimethoprim-sulfamethoxazole prevents *Pneumocystis jiroveci* pneumonia. These patients, and patients with phagocytic cell dysfunction, should avoid heavy exposure to airborne soil, dust, or decaying matter (mulch, manure), which are often rich in *Nocardia* and the spores of *Aspergillus* and other fungi. Restriction of activities or social contact has no proven role in reducing risk of infection for phagocyte defects.

Although aggressive medical care for many patients with phagocytic disorders can allow them to go for years without a life-threatening infection, there may still be delayed effects of prolonged antimicrobials and other inflammatory complications. Cure of most congenital phagocyte defects is possible by bone marrow transplantation, and rates of success are improving (Chap. 139e). The identification of specific gene defects in patients with LAD 1, CGD, and other immunodeficiencies has led to gene therapy trials in a number of genetic white cell disorders.

Some of the relevant findings in peripheral blood, enlarged lymph nodes, and bone marrow are illustrated in this chapter. Systematic histologic examination of the bone marrow and lymph nodes is beyond the scope of a general medicine textbook. However, every internist should know how to examine a peripheral blood smear.

The examination of a peripheral blood smear is one of the most informative exercises a physician can perform. Although advances in automated technology have made the examination of a peripheral blood smear by a physician seem less important, the technology is not a completely satisfactory replacement for a blood smear interpretation by a trained medical professional who also knows the patient's clinical history, family history, social history, and physical findings. It is useful to ask the laboratory to generate a Wright's-stained peripheral blood smear and examine it.

The best place to examine blood cell morphology is the feathered edge of the blood smear where red cells lie in a single layer, side by side, just barely touching one another but not overlapping. The author's approach is to look at the smallest cellular elements, the platelets, first and work his way up in size to red cells and then white cells.

Using an oil immersion lens that magnifies the cells 100-fold, one counts the platelets in five to six fields, averages the number per field, and multiplies by 20,000 to get a rough estimate of the platelet count. The platelets are usually 1–2 μm in diameter and have a blue granulated appearance. There is usually 1 platelet for every 20 or so red cells. Of course, the automated counter is much more accurate, but gross disparities between the automated and manual counts should be assessed. Large platelets may be a sign of rapid platelet turnover, as young platelets are often larger than old ones; alternatively, certain rare inherited syndromes can produce large platelets. Platelet clumping visible on the smear can be associated with falsely low automated platelet counts. Similarly, neutrophil fragmentation can be a source of falsely elevated automated platelet counts.

Next one examines the red blood cells. One can gauge their size by comparing the red cell to the nucleus of a small lymphocyte. Both are normally about 8 μm wide. Red cells that are smaller than the small lymphocyte nucleus may be microcytic; those larger than the small lymphocyte nucleus may be macrocytic. Macrocytic cells also tend to be more oval than spherical in shape and are sometimes called macroovalocytes. The automated mean corpuscular volume (MCV) can assist in making a classification. However, some patients may have both iron and vitamin B₁₂ deficiency, which will produce an MCV in the normal range but wide variation in red cell size. When the red cells vary greatly in size, *anisocytosis* is said to be present. When the red cells vary greatly in shape, *poikilocytosis* is said to be present. The electronic cell counter provides an independent assessment of variability in red cell size. It measures the range of red cell volumes and reports the results as "red cell distribution width" (RDW). This value is calculated from the MCV; thus, cell width is not being measured but cell volume is. The term is derived from the curve displaying the frequency of cells at each volume, also called the distribution. The width of the red cell volume distribution curve is what determines the RDW. The RDW is calculated as follows: $\text{RDW} = (\text{standard deviation of MCV} \div \text{mean MCV}) \times 100$. In the presence of morphologic anisocytosis, RDW (normally 11–14%) increases to 15–18%. The RDW is useful in at least two clinical settings. In patients with microcytic anemia, the differential diagnosis is generally between iron deficiency and thalassemia. In thalassemia, the small red cells are generally of uniform size with a normal small RDW. In iron deficiency, the size variability and the RDW are large. In addition, a large RDW can suggest a dimorphic anemia when a chronic atrophic gastritis can produce both vitamin B₁₂ malabsorption to produce macrocytic anemia and blood loss to produce iron deficiency. In such settings, RDW is also large. An elevated

RDW also has been reported as a risk factor for all-cause mortality in population-based studies (Patel KV et al: *Arch Intern Med* 169:515, 2009), a finding that is unexplained currently.

After red cell size is assessed, one examines the hemoglobin content of the cells. They are either normal in color (*normochromic*) or pale in color (*hypochromic*). They are never "hyperchromic." If more than the normal amount of hemoglobin is made, the cells get larger—they do not become darker. In addition to hemoglobin content, the red cells are examined for inclusions. Red cell inclusions are the following:

1. *Basophilic stippling*—diffuse fine or coarse blue dots in the red cell usually representing RNA residue—especially common in lead poisoning
2. *Howell-Jolly bodies*—dense blue circular inclusions that represent nuclear remnants—their presence implies defective splenic function
3. *Nuclei*—red cells may be released or pushed out of the marrow prematurely before nuclear extrusion—often implies a myelophthitic process or a vigorous narrow response to anemia, usually hemolytic anemia
4. *Parasites*—red cell parasites include malaria and babesia ([Chap. 250e](#))
5. *Polychromatophilia*—the red cell cytoplasm has a bluish hue, reflecting the persistence of ribosomes still actively making hemoglobin in a young red cell

Vital stains are necessary to see precipitated hemoglobin called *Heinz bodies*.

Red cells can take on a variety of different shapes. All abnormally shaped red cells are *poikilocytes*. Small red cells without the central pallor are *spherocytes*; they can be seen in hereditary spherocytosis, hemolytic anemias of other causes, and clostridial sepsis. *Dacrocytes* are teardrop-shaped cells that can be seen in hemolytic anemias, severe iron deficiency, thalassemias, myelofibrosis, and myelodysplastic syndromes. *Schistocytes* are helmet-shaped cells that reflect microangiopathic hemolytic anemia or fragmentation on an artificial heart valve. *Echinocytes* are spiculated red cells with the spikes evenly spaced; they can represent an artifact of abnormal drying of the blood smear or reflect changes in stored blood. They also can be seen in renal failure and malnutrition and are often reversible. *Acanthocytes* are spiculated red cells with the spikes irregularly distributed. This process tends to be irreversible and reflects underlying renal disease, abetalipoproteinemia, or splenectomy. *Elliptocytes* are elliptical-shaped red cells that can reflect an inherited defect in the red cell membrane, but they also are seen in iron deficiency, myelodysplastic syndromes, megaloblastic anemia, and thalassemias. *Stomatocytes* are red cells in which the area of central pallor takes on the morphology of a slit instead of the usual round shape. Stomatocytes can indicate an inherited red cell membrane defect and also can be seen in alcoholism. *Target cells* have an area of central pallor that contains a dense center, or bull's-eye. These cells are seen classically in thalassemia, but they are also present in iron deficiency, cholestatic liver disease, and some hemoglobinopathies. They also can be generated artifactually by improper slide making.

One last feature of the red cells to assess before moving to the white blood cells is the distribution of the red cells on the smear. In most individuals, the cells lie side by side in a single layer. Some patients have red cell clumping (called *agglutination*) in which the red cells pile upon one another; it is seen in certain paraproteinemias and autoimmune hemolytic anemias. Another abnormal distribution involves red cells lying in single cell rows on top of one another like stacks of coins. This is called *rouleaux formation* and reflects abnormal serum protein levels.

Finally, one examines the white blood cells. Three types of granulocytes are usually present: neutrophils, eosinophils, and basophils, in decreasing frequency. Neutrophils are generally the most abundant white cell. They are round, are 10–14 μm wide, and contain a lobulated nucleus with two to five lobes connected by a thin chromatin thread. Bands are immature neutrophils that have not completed nuclear condensation and have a U-shaped nucleus. Bands reflect a left shift in neutrophil maturation in an effort to make more cells more rapidly.

Neutrophils can provide clues to a variety of conditions. Vacuolated neutrophils may be a sign of bacterial sepsis. The presence of 1- to 2- μm blue cytoplasmic inclusions, called *Döhle bodies*, can reflect infections, burns, or other inflammatory states. If the neutrophil granules are larger than normal and stain a darker blue, “toxic granulations” are said to be present, and they also suggest a systemic inflammation. The presence of neutrophils with more than five nuclear lobes suggests megaloblastic anemia. Large misshapen granules may reflect the inherited Chédiak-Higashi syndrome.

Eosinophils are slightly larger than neutrophils, have bilobed nuclei, and contain large red granules. Diseases of eosinophils are associated with too many of them rather than any morphologic or qualitative change. They normally total less than one-thirtieth the number of neutrophils. Basophils are even rarer than eosinophils in the blood. They have large dark blue granules and may be increased as part of chronic myeloid leukemia.

Lymphocytes can be present in several morphologic forms. Most common in healthy individuals are small lymphocytes with a small dark nucleus and scarce cytoplasm. In the presence of viral infections, more of the lymphocytes are larger, about the size of neutrophils, with

abundant cytoplasm and a less condensed nuclear chromatin. These cells are called *reactive lymphocytes*. About 1% of lymphocytes are larger and contain blue granules in a light blue cytoplasm; they are called *large granular lymphocytes*. In chronic lymphoid leukemia, the small lymphocytes are increased in number, and many of them are ruptured in making the blood smear, leaving a smudge of nuclear material without a surrounding cytoplasm or cell membrane; they are called *smudge cells* and are rare in the absence of chronic lymphoid leukemia.

Monocytes are the largest white blood cells, ranging from 15 to 22 μm in diameter. The nucleus can take on a variety of shapes but usually appears to be folded; the cytoplasm is gray.

Abnormal cells may appear in the blood. Most often the abnormal cells originate from neoplasms of bone marrow–derived cells, including lymphoid cells, myeloid cells, and occasionally red cells. More rarely, other types of tumors can get access to the bloodstream, and rare epithelial malignant cells may be identified. The chances of seeing such abnormal cells is increased by examining blood smears made from buffy coats, the layer of cells that is visible on top of sedimenting red cells when blood is left in the test tube for an hour. Smears made from finger sticks may include rare endothelial cells.

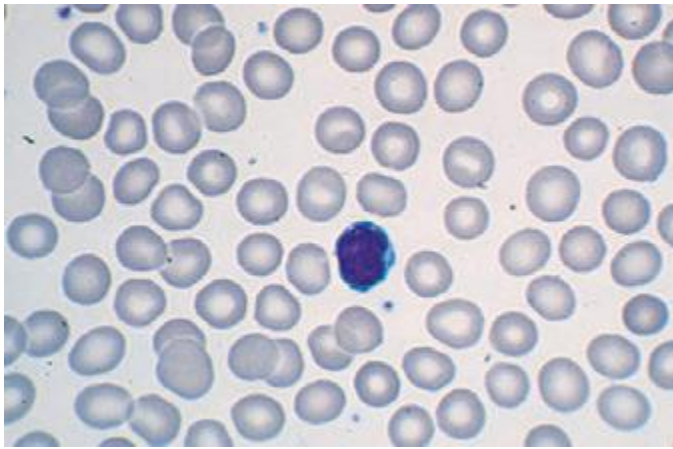


FIGURE 81e-1 Normal peripheral blood smear. Small lymphocyte in center of field. Note that the diameter of the red blood cell is similar to the diameter of the small lymphocyte nucleus.

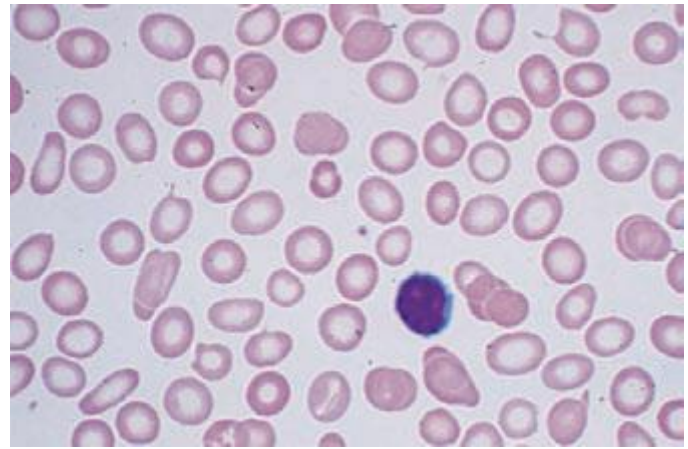


FIGURE 81e-3 Hypochromic microcytic anemia of iron deficiency. Small lymphocyte in field helps assess the red blood cell size.

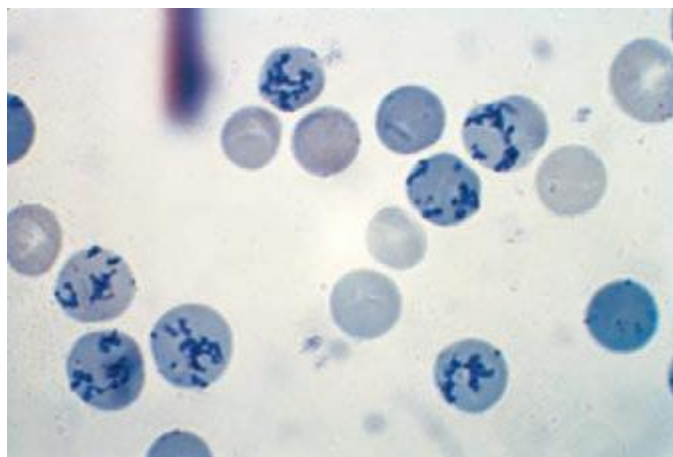


FIGURE 81e-2 Reticulocyte count preparation. This new methylene blue–stained blood smear shows large numbers of heavily stained reticulocytes (the cells containing the dark blue–staining RNA precipitates).

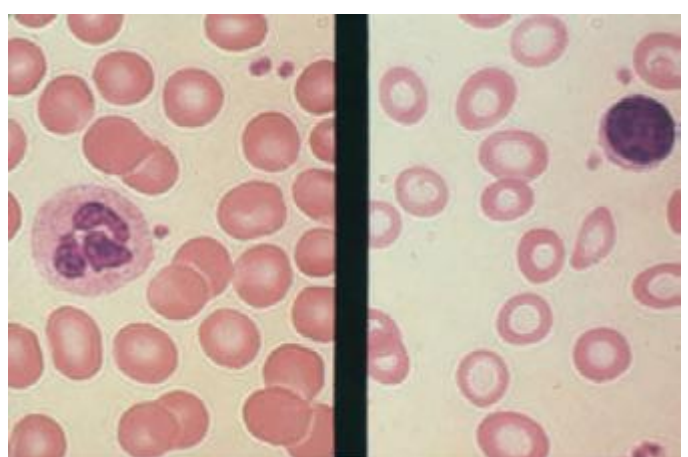


FIGURE 81e-4 Iron deficiency anemia next to normal red blood cells. Microcytes (*right panel*) are smaller than normal red blood cells (cell diameter $<7\ \mu\text{m}$) and may or may not be poorly hemoglobinized (hypochromic).

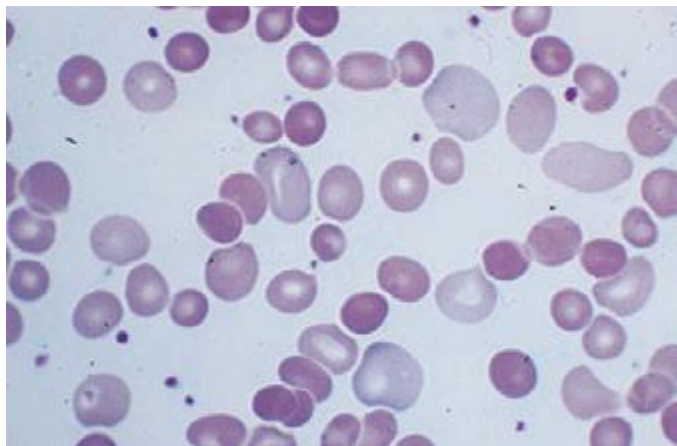


FIGURE 81e-5 Polychromatophilia. Note large red cells with light purple coloring.

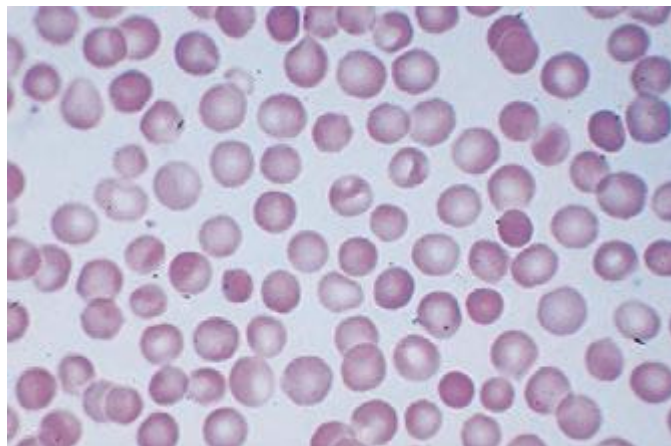


FIGURE 81e-8 Spherocytosis. Note small hyperchromatic cells without the usual clear area in the center.

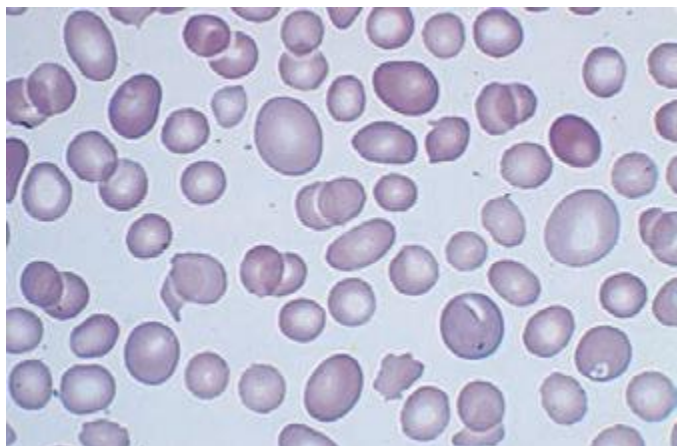


FIGURE 81e-6 Macrocytosis. These cells are both larger than normal (mean corpuscular volume >100) and somewhat oval in shape. Some morphologists call these cells macroovalocytes.

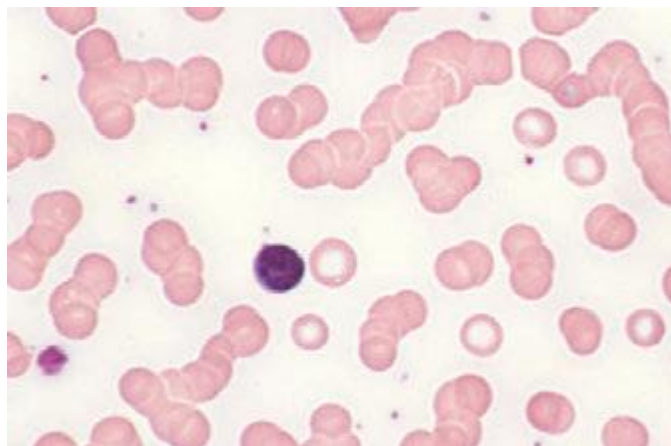


FIGURE 81e-9 Rouleaux formation. Small lymphocyte in center of field. These red cells align themselves in stacks and are related to increased serum protein levels.

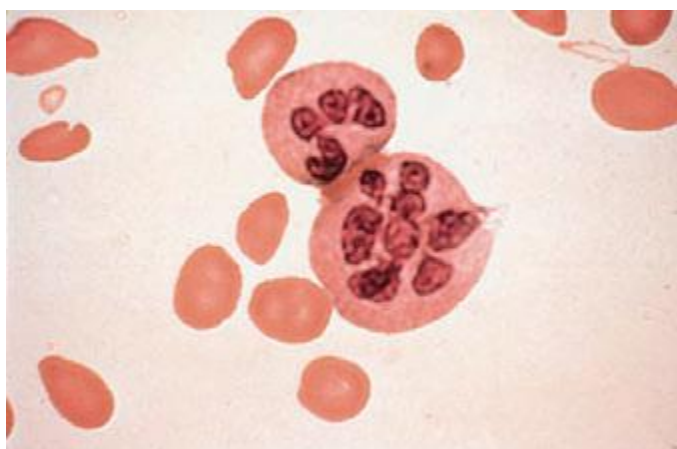


FIGURE 81e-7 Hypersegmented neutrophils. Hypersegmented neutrophils (multilobed polymorphonuclear leukocytes) are larger than normal neutrophils with five or more segmented nuclear lobes. They are commonly seen with folic acid or vitamin B₁₂ deficiency.

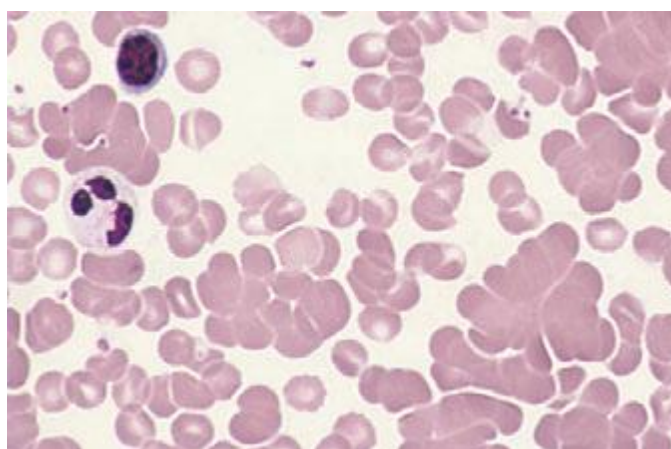


FIGURE 81e-10 Red cell agglutination. Small lymphocyte and segmented neutrophil in upper left center. Note irregular collections of aggregated red cells.

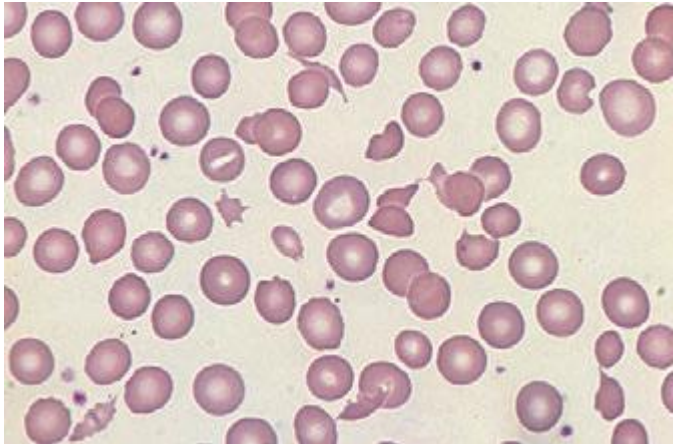


FIGURE 81e-11 Fragmented red cells. Heart valve hemolysis.

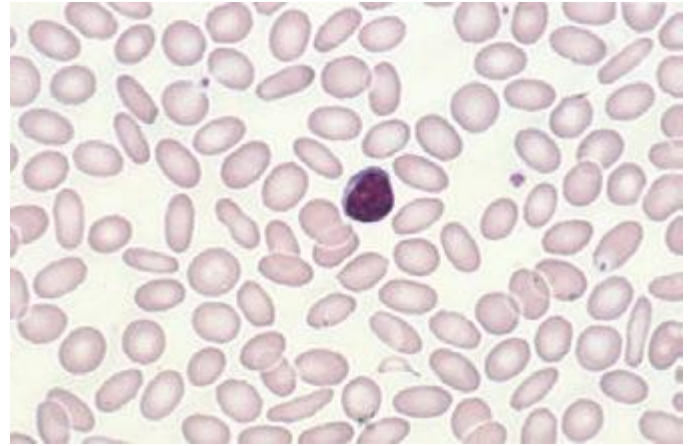


FIGURE 81e-14 Elliptocytosis. Small lymphocyte in center of field. Elliptical shape of red cells is related to weakened membrane structure, usually due to mutations in spectrin.

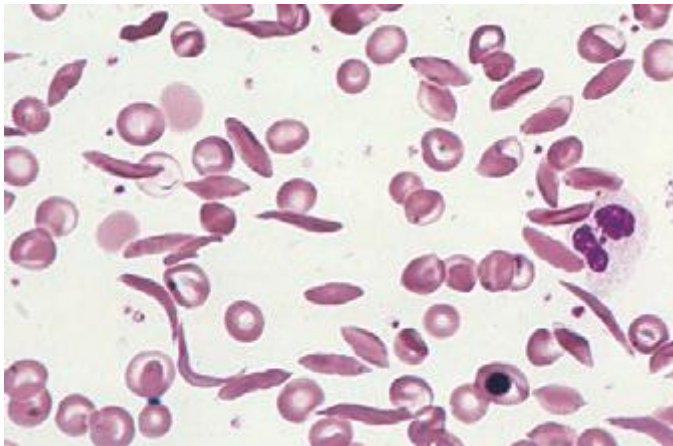


FIGURE 81e-12 Sickle cells. Homozygous sickle cell disease. A nucleated red cell and neutrophil are also in the field.

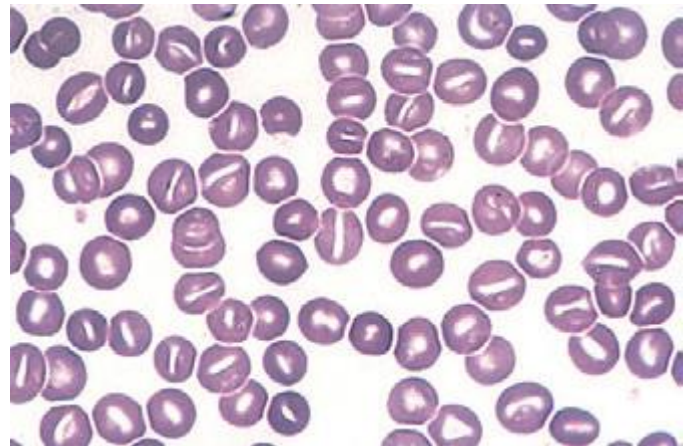


FIGURE 81e-15 Stomatocytosis. Red cells characterized by a wide transverse slit or stoma. This often is seen as an artifact in a dehydrated blood smear. These cells can be seen in hemolytic anemias and in conditions in which the red cell is overhydrated or dehydrated.

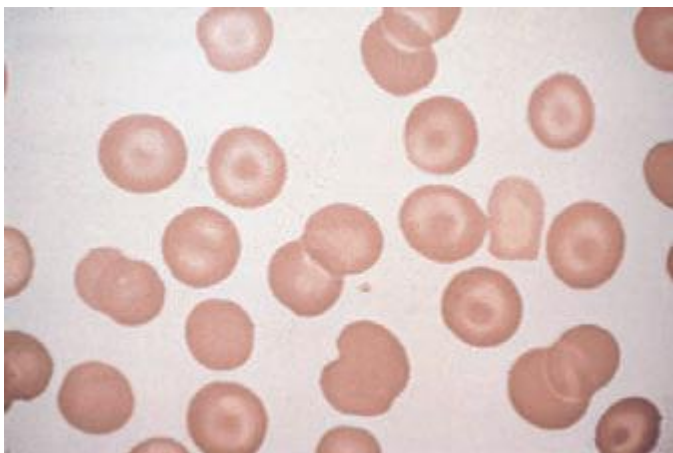


FIGURE 81e-13 Target cells. Target cells are recognized by the bull's-eye appearance of the cell. Small numbers of target cells are seen with liver disease and thalassemia. Larger numbers are typical of hemoglobin C disease.

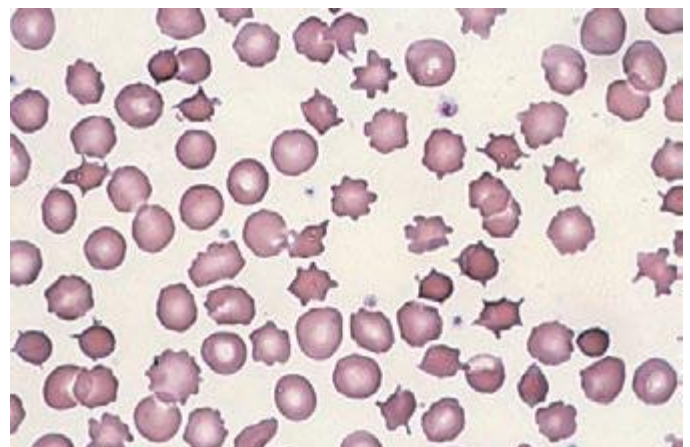


FIGURE 81e-16 Acanthocytosis. Spiculated red cells are of two types: *acanthocytes* are contracted dense cells with irregular membrane projections that vary in length and width; *echinocytes* have small, uniform, and evenly spaced membrane projections. Acanthocytes are present in severe liver disease, in patients with abetalipoproteinemia, and in rare patients with McLeod blood group. Echinocytes are found in patients with severe uremia, in glycolytic red cell enzyme defects, and in microangiopathic hemolytic anemia.

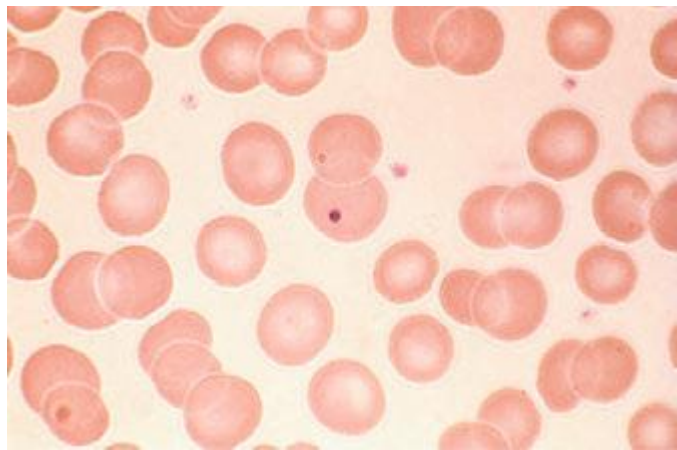


FIGURE 81e-17 Howell-Jolly bodies. Howell-Jolly bodies are tiny nuclear remnants that normally are removed by the spleen. They appear in the blood after splenectomy (defect in removal) and with maturation/dysplastic disorders (excess production).

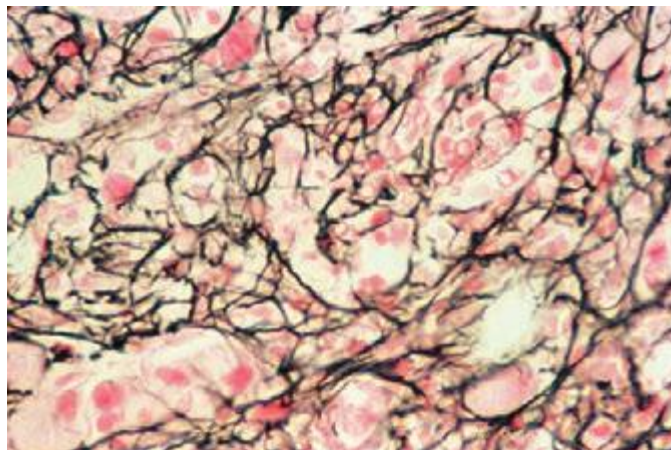


FIGURE 81e-20 Reticulin stain of marrow myelofibrosis. Silver stain of a myelofibrotic marrow showing an increase in reticulin fibers (black-staining threads).

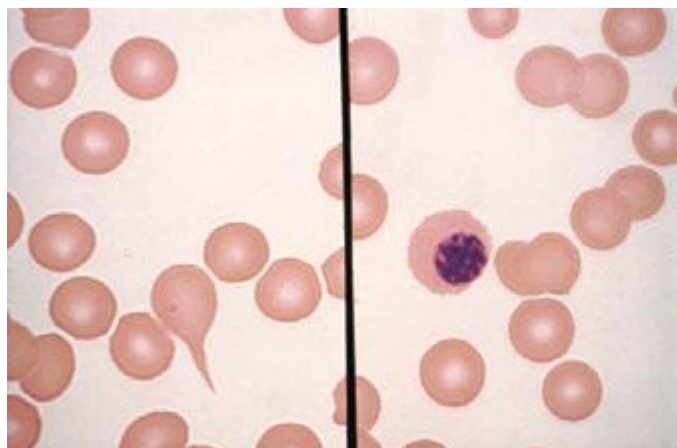


FIGURE 81e-18 Teardrop cells and nucleated red blood cells characteristic of myelofibrosis. A teardrop-shaped red blood cell (*left panel*) and a nucleated red blood cell (*right panel*) as typically seen with myelofibrosis and extramedullary hematopoiesis.

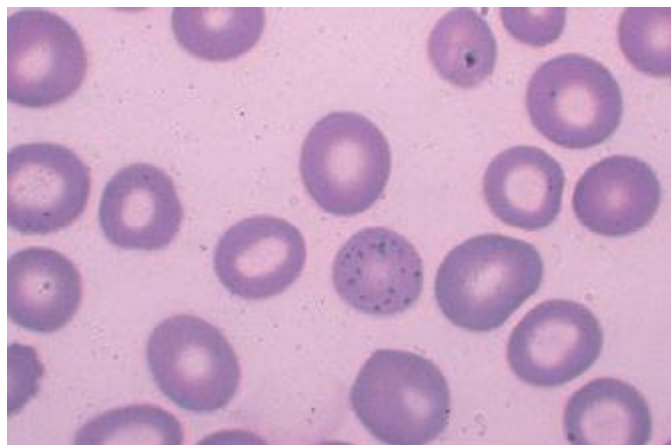


FIGURE 81e-21 Stippled red cell in lead poisoning. Mild hypochromia. Coarsely stippled red cell.

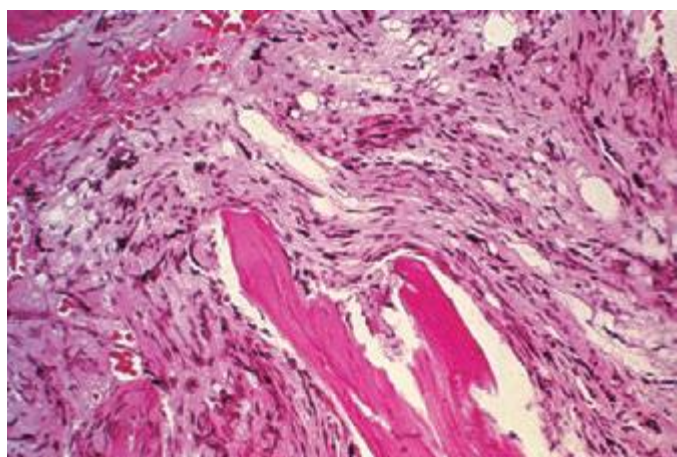


FIGURE 81e-19 Myelofibrosis of the bone marrow. Total replacement of marrow precursors and fat cells by a dense infiltrate of reticulin fibers and collagen (H&E stain).

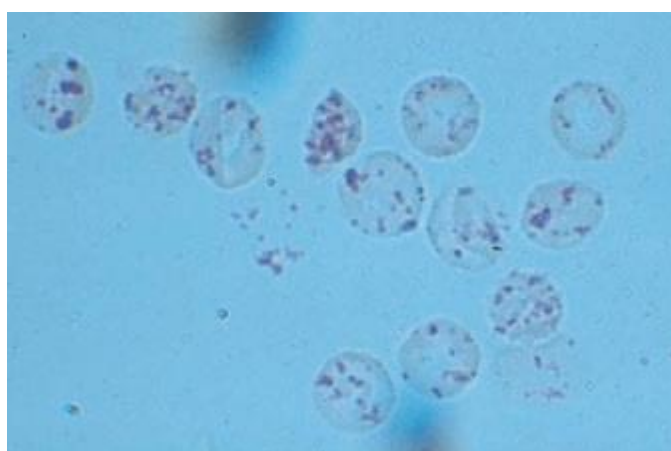


FIGURE 81e-22 Heinz bodies. Blood mixed with hypotonic solution of crystal violet. The stained material is precipitates of denatured hemoglobin within cells.

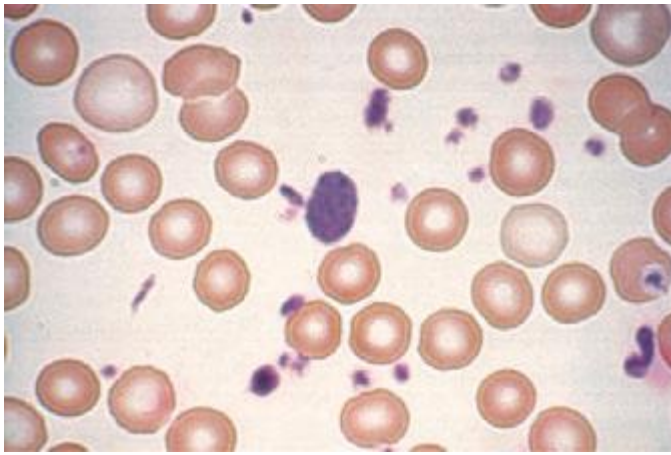


FIGURE 81e-23 Giant platelets. Giant platelets, together with a marked increase in the platelet count, are seen in myeloproliferative disorders, especially primary thrombocythemia.

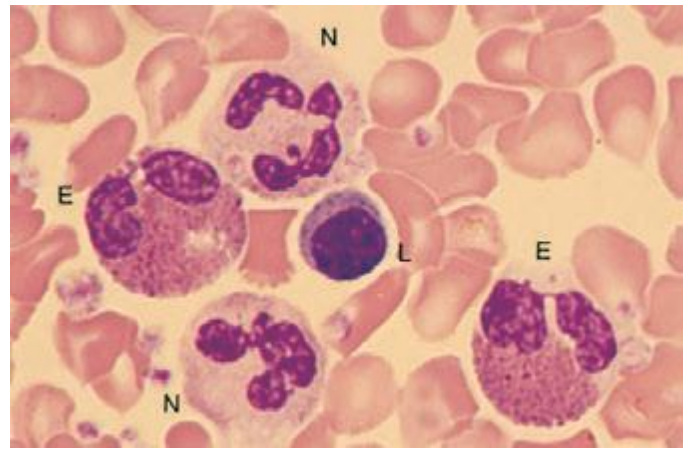


FIGURE 81e-26 Normal eosinophils. The film was prepared from the buffy coat of the blood from a normal donor. E, eosinophil; L, lymphocyte; N, neutrophil.

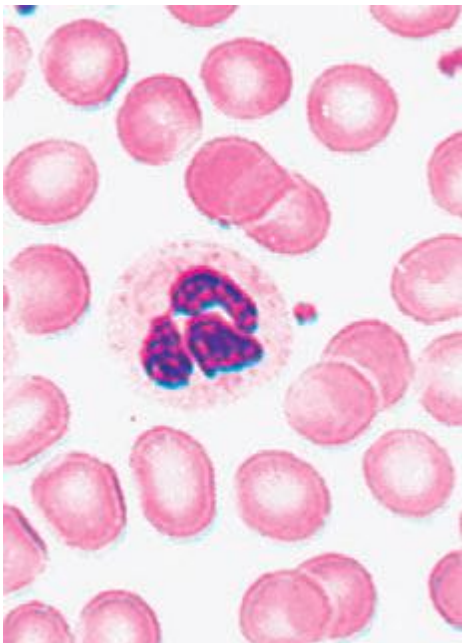


FIGURE 81e-24 Normal granulocytes. The normal granulocyte has a segmented nucleus with heavy, clumped chromatin; fine neutrophilic granules are dispersed throughout the cytoplasm.

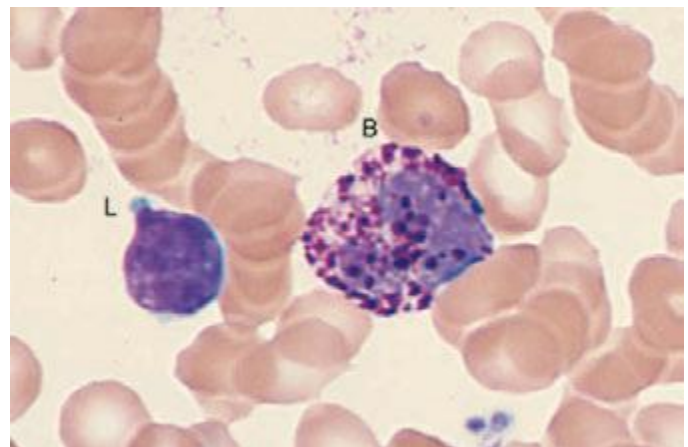


FIGURE 81e-27 Normal basophil. The film was prepared from the buffy coat of the blood from a normal donor. B, basophil; L, lymphocyte.

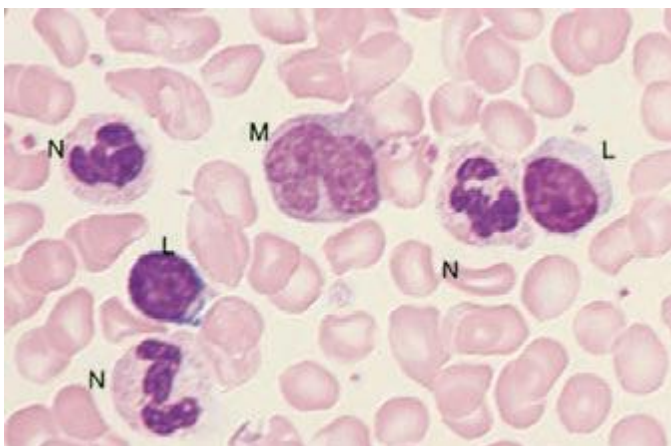


FIGURE 81e-25 Normal monocytes. The film was prepared from the buffy coat of the blood from a normal donor. L, lymphocyte; M, monocyte; N, neutrophil.

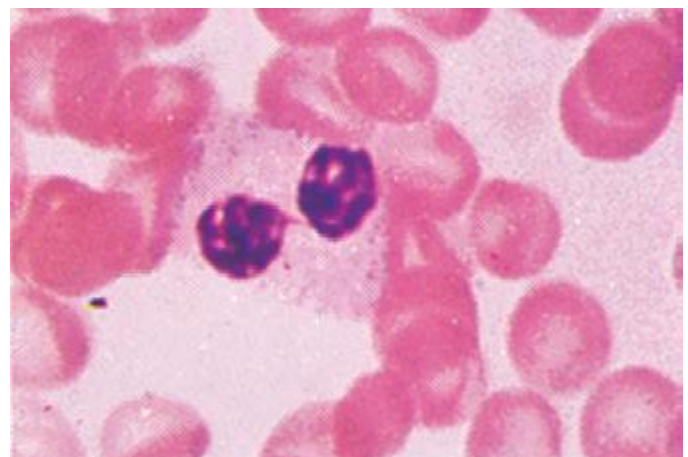


FIGURE 81e-28 Pelger-Huet anomaly. In this benign disorder, the majority of granulocytes are bilobed. The nucleus frequently has a spectacle-like, or "pince-nez," configuration.

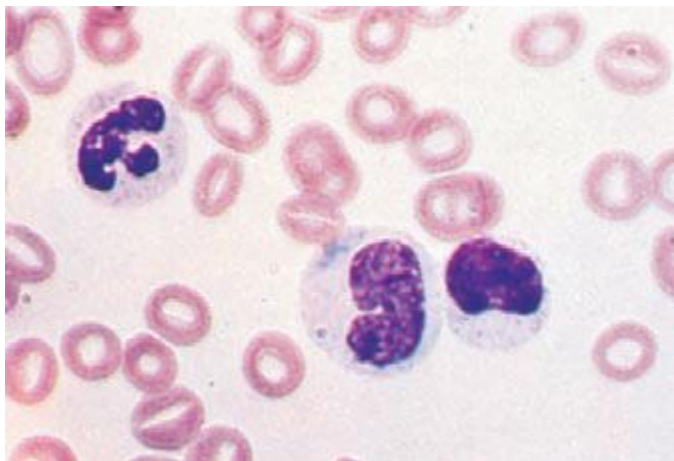


FIGURE 81e-29 Döhle body. Neutrophil band with Döhle body. The neutrophil with a sausage-shaped nucleus in the center of the field is a band form. Döhle bodies are discrete, blue-staining nongranular areas found in the periphery of the cytoplasm of the neutrophil in infections and other toxic states. They represent aggregates of rough endoplasmic reticulum.

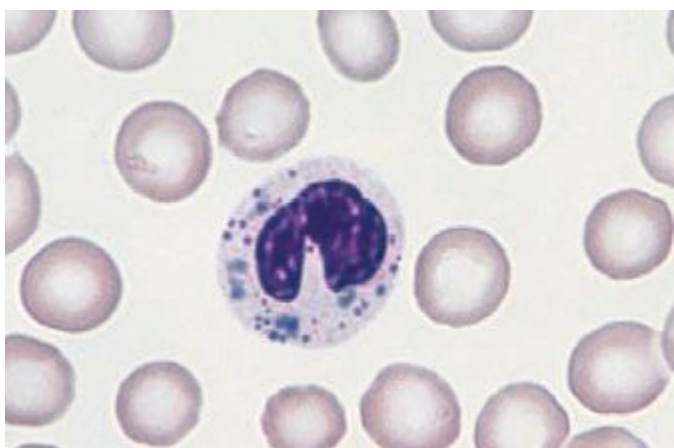


FIGURE 81e-30 Chédiak-Higashi disease. Note giant granules in neutrophil.

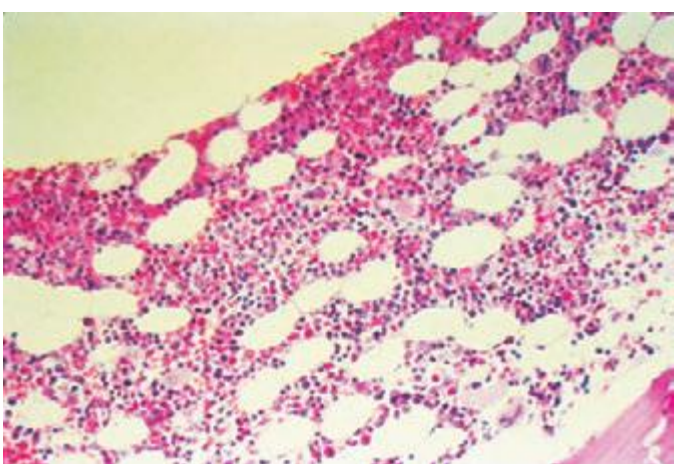


FIGURE 81e-31 Normal bone marrow. Low-power view of normal adult marrow (hematoxylin and eosin [H&E] stain), showing a mix of fat cells (clear areas) and hematopoietic cells. The percentage of the space that consists of hematopoietic cells is referred to as *marrow cellularity*. In adults, normal marrow cellularity is 35–40%. If demands for increased marrow production occur, cellularity may increase to meet the demand. As people age, the marrow cellularity decreases and the marrow fat increases. Patients >70 years old may have a 20–30% marrow cellularity.

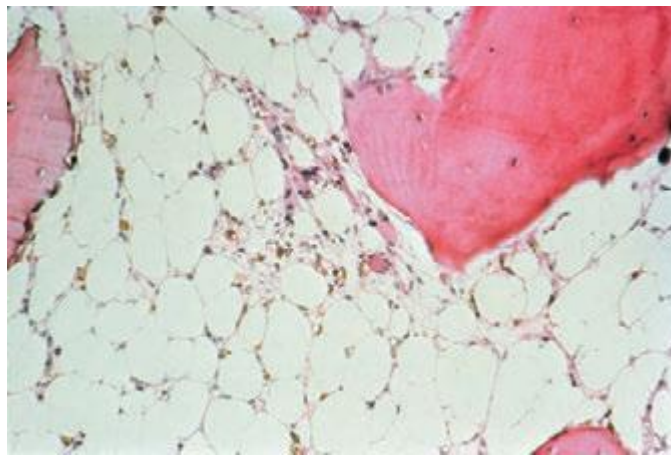


FIGURE 81e-32 Aplastic anemia bone marrow. Normal hematopoietic precursor cells are virtually absent, leaving behind fat cells, reticuloendothelial cells, and the underlying sinusoidal structure.

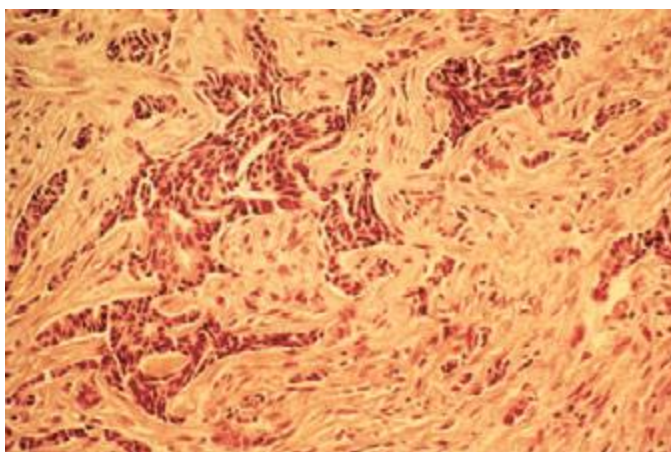


FIGURE 81e-33 Metastatic cancer in the bone marrow. Marrow biopsy specimen infiltrated with metastatic breast cancer and reactive fibrosis (H&E stain).

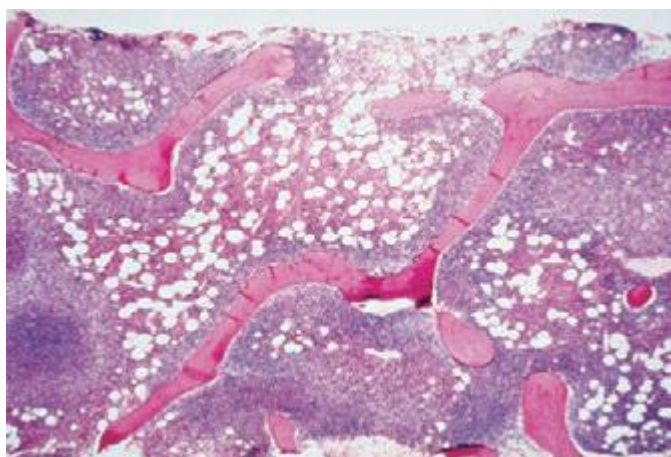


FIGURE 81e-34 Lymphoma in the bone marrow. Nodular (follicular) lymphoma infiltrate in a marrow biopsy specimen. Note the characteristic paratrabecular location of the lymphoma cells.

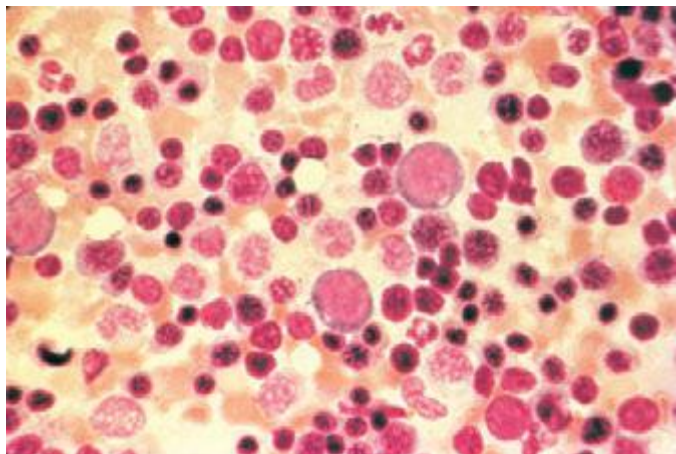


FIGURE 81e-35 Erythroid hyperplasia of the marrow. Marrow aspirate specimen with a myeloid/erythroid ratio (M/E ratio) of 1:1–2, typical for a patient with a hemolytic anemia or one recovering from blood loss.

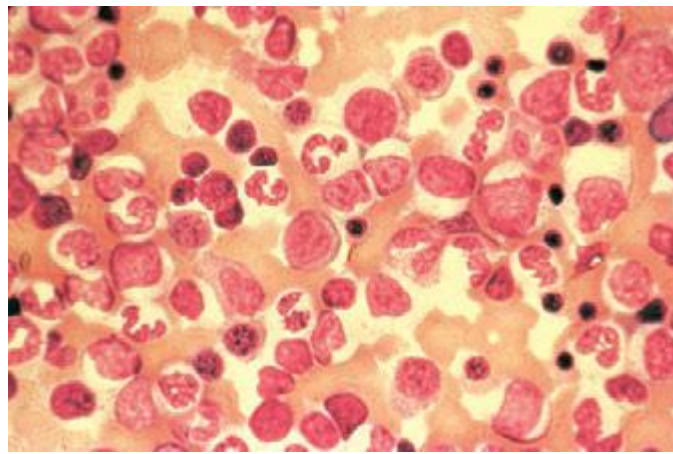


FIGURE 81e-36 Myeloid hyperplasia of the marrow. Marrow aspirate specimen showing a myeloid/erythroid ratio of $\geq 3:1$, suggesting either a loss of red blood cell precursors or an expansion of myeloid elements.

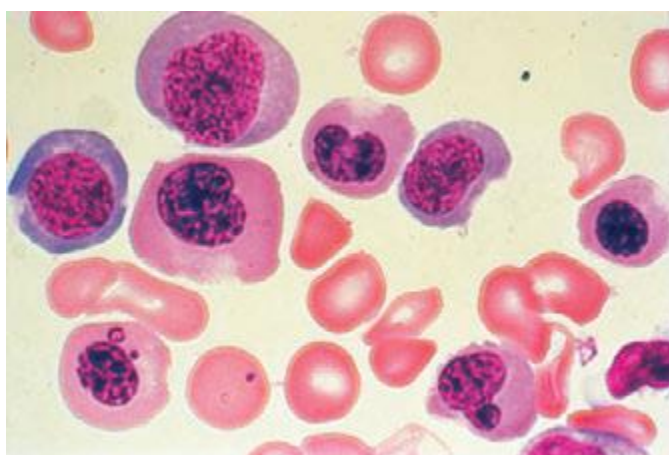


FIGURE 81e-37 Megaloblastic erythropoiesis. High-power view of megaloblastic red blood cell precursors from a patient with a macrocytic anemia. Maturation is delayed, with late normoblasts showing a more immature-appearing nucleus with a lattice-like pattern with normal cytoplasmic maturation.

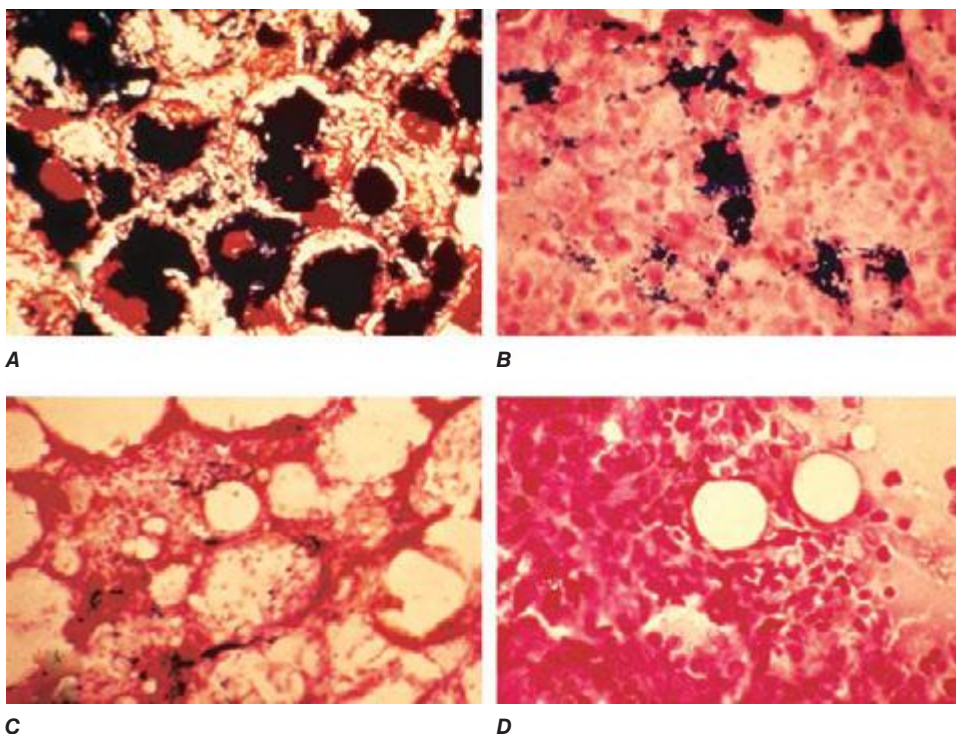


FIGURE 81e-38 Prussian blue staining of marrow iron stores. Iron stores can be graded on a scale of 0 to 4+. **A.** A marrow with excess iron stores (>4+); **B.** normal stores (2–3+); **C.** minimal stores (1+); and **D.** absent iron stores (0).

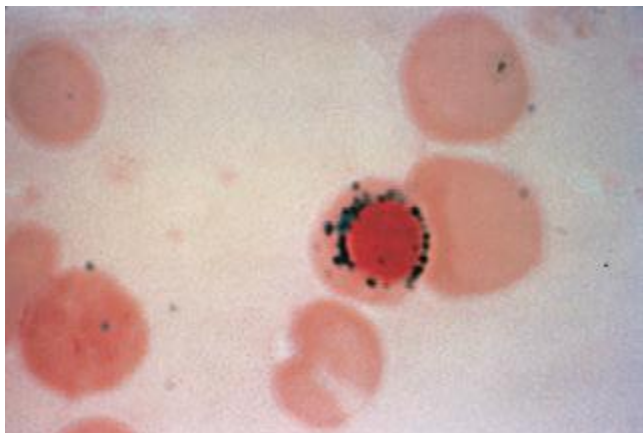


FIGURE 81e-39 Ringed sideroblast. An orthochromatic normoblast with a collar of blue granules (mitochondria encrusted with iron) surrounding the nucleus.

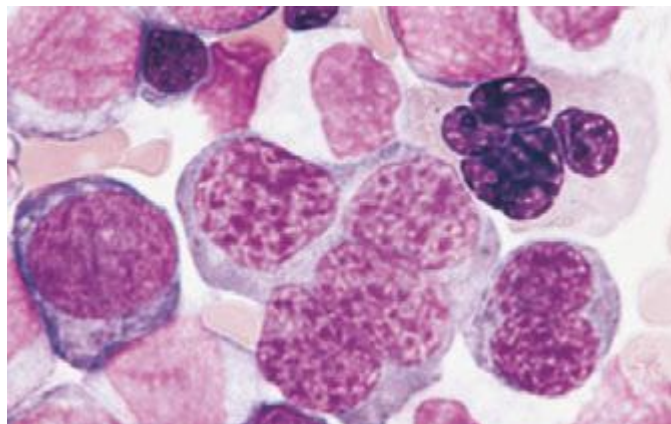


FIGURE 81e-42 Acute erythroleukemia. Note giant dysmorphic erythroblasts; two are binucleate, and one is multinucleate.

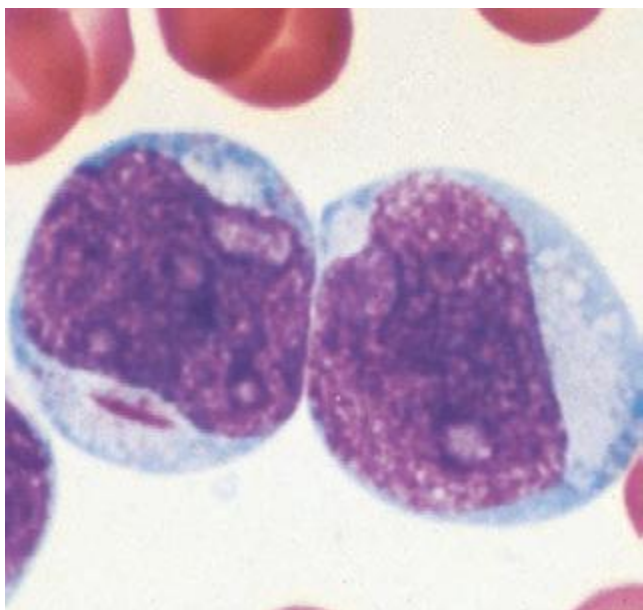


FIGURE 81e-40 Acute myeloid leukemia. Leukemic myeloblast with an Auer rod. Note two to four large, prominent nucleoli in each cell.

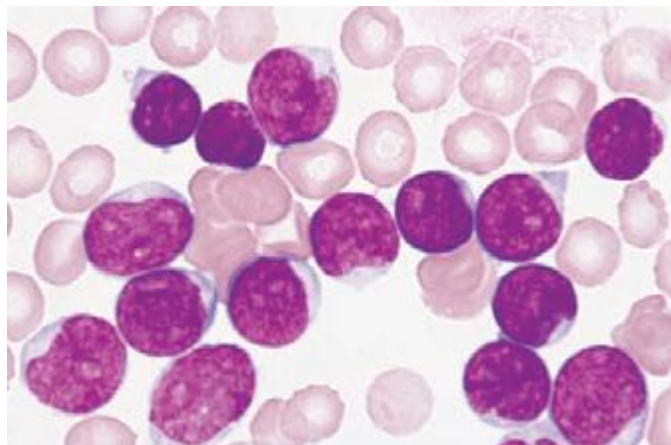


FIGURE 81e-43 Acute lymphoblastic leukemia.

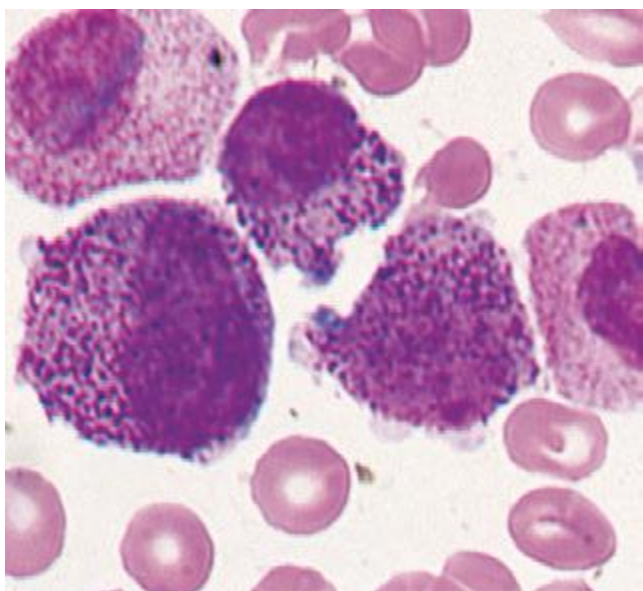


FIGURE 81e-41 Acute promyelocytic leukemia. Note prominent cytoplasmic granules in the leukemia cells.

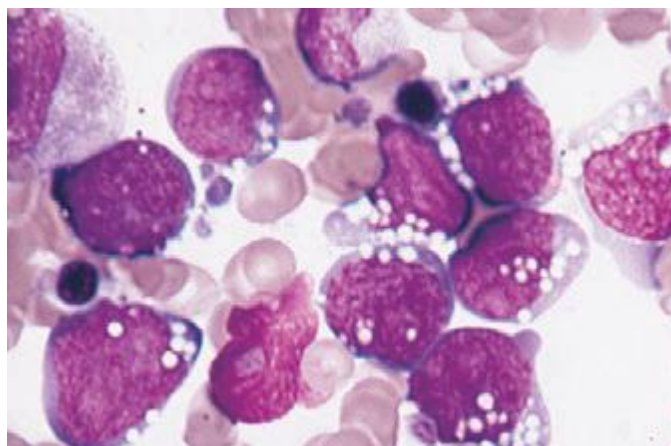


FIGURE 81e-44 Burkitt's leukemia, acute lymphoblastic leukemia.

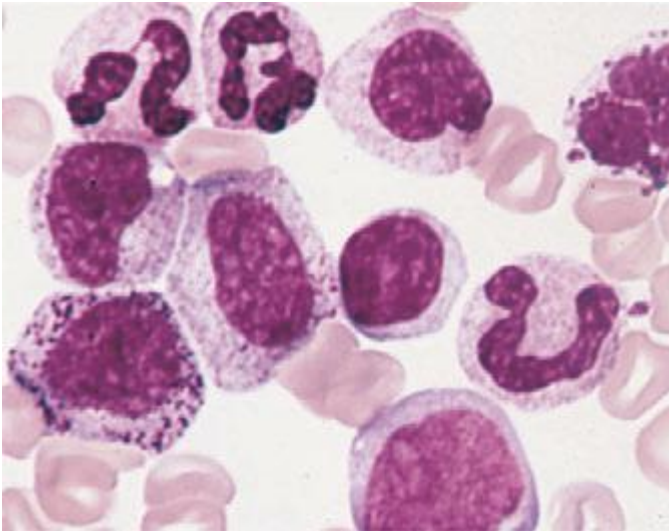


FIGURE 81e-45 Chronic myeloid leukemia in the peripheral blood.

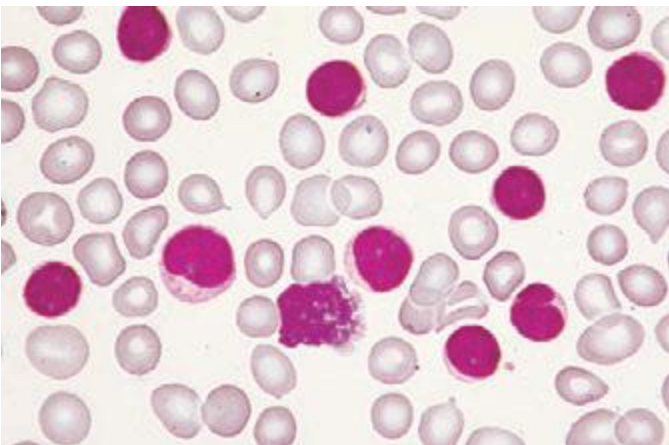


FIGURE 81e-46 Chronic lymphoid leukemia in the peripheral blood.

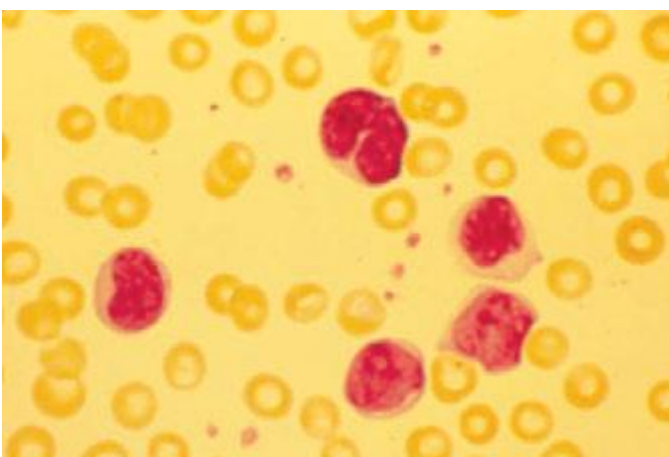


FIGURE 81e-47 Sézary's syndrome. Lymphocytes with frequently convoluted nuclei (Sézary cells) in a patient with advanced mycosis fungoides.

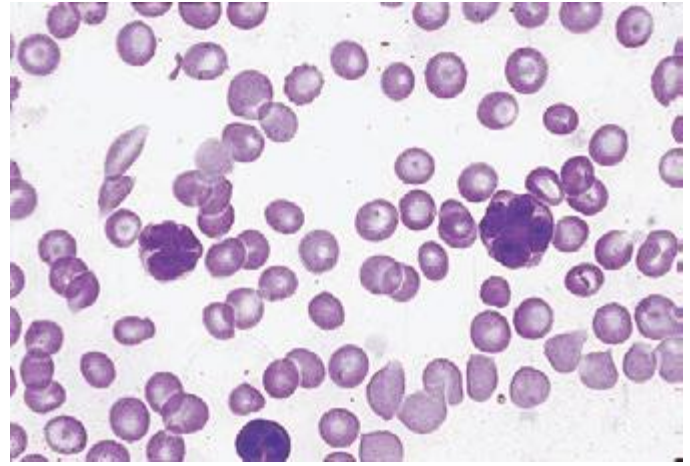


FIGURE 81e-48 Adult T cell leukemia. Peripheral blood smear showing leukemia cells with typical "flower-shaped" nucleus.

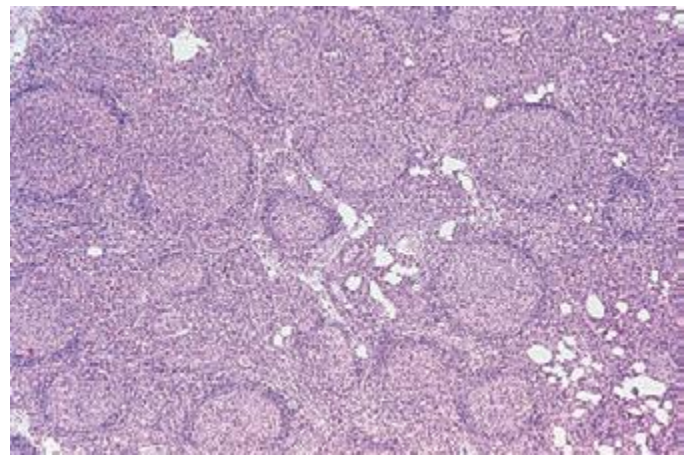


FIGURE 81e-49 Follicular lymphoma in a lymph node. The normal nodal architecture is effaced by nodular expansions of tumor cells. Nodules vary in size and contain predominantly small lymphocytes with cleaved nuclei along with variable numbers of larger cells with vesicular chromatin and prominent nucleoli.

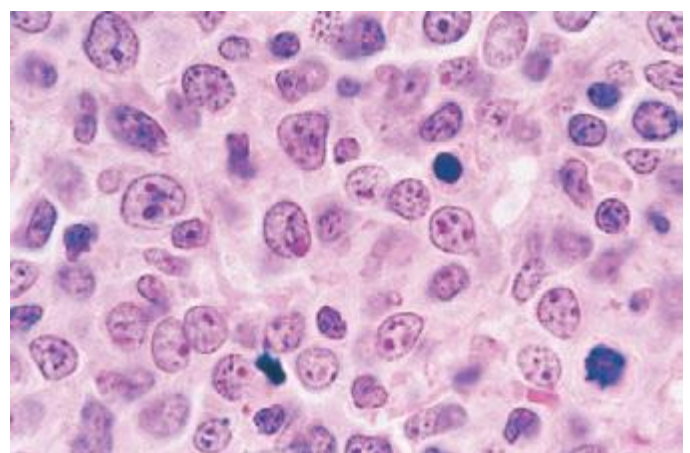


FIGURE 81e-50 Diffuse large B cell lymphoma in a lymph node. The neoplastic cells are heterogeneous but predominantly large cells with vesicular chromatin and prominent nucleoli.

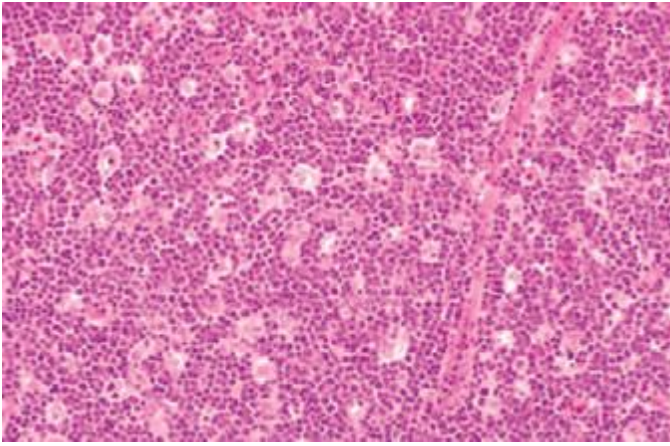


FIGURE 81e-51 Burkitt's lymphoma in a lymph node. Burkitt's lymphoma with starry-sky appearance. The lighter areas are macrophages attempting to clear dead cells.

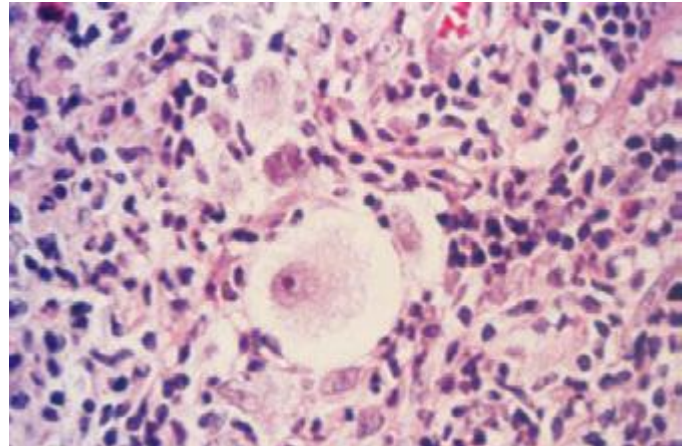


FIGURE 81e-54 Lacunar cell; Reed-Sternberg cell variant in nodular sclerosing Hodgkin's disease. High-power view of single mononuclear lacunar cell with retracted cytoplasm in a patient with nodular sclerosing Hodgkin's disease.

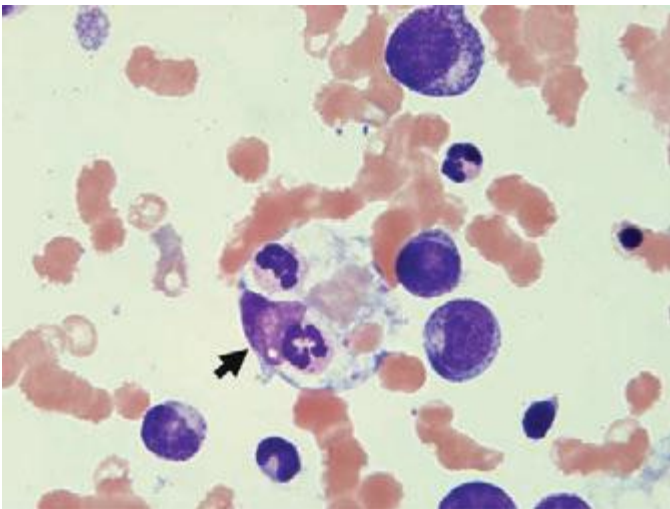


FIGURE 81e-52 Erythrophagocytosis accompanying aggressive lymphoma. The central macrophage is ingesting red cells, neutrophils, and platelets. (Courtesy of Dr. Kiyomi Tsukimori, Kyushu University, Fukuoka, Japan.)

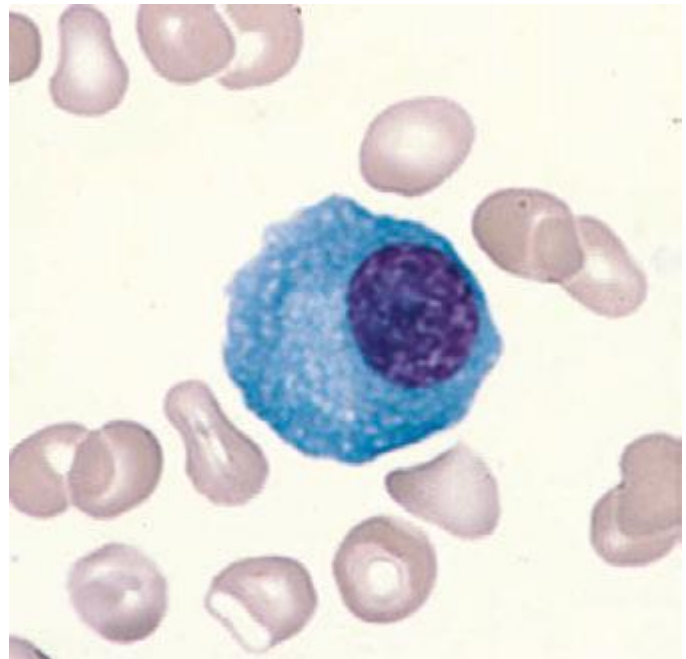


FIGURE 81e-55 Normal plasma cell.

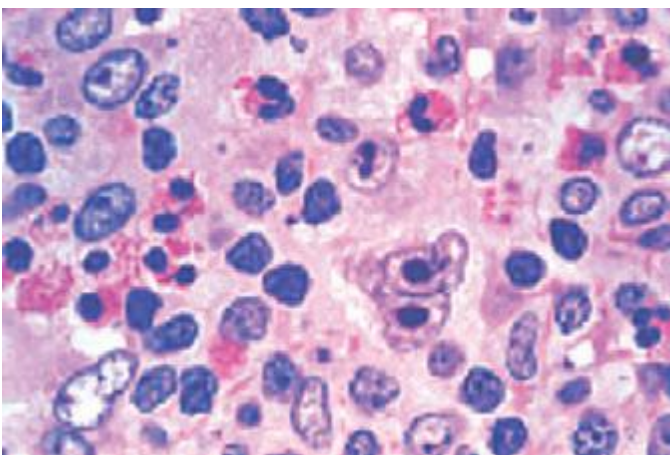


FIGURE 81e-53 Hodgkin's disease. A Reed-Sternberg cell is present near the center of the field; a large cell with a bilobed nucleus and prominent nucleoli giving an "owl's eyes" appearance. The majority of the cells are normal lymphocytes, neutrophils, and eosinophils that form a pleiomorphic cellular infiltrate.

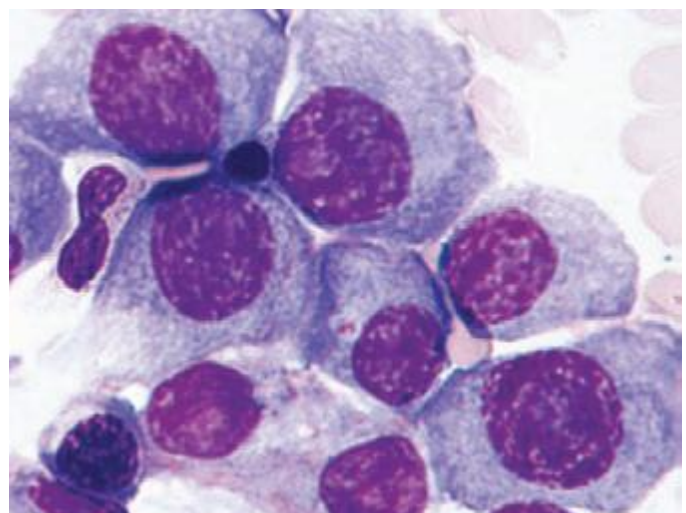


FIGURE 81e-56 Multiple myeloma.



FIGURE 81e-57 Serum color in hemoglobinemia. The distinctive red coloration of plasma (hemoglobinemia) in a spun blood sample in a patient with intravascular hemolysis.

ACKNOWLEDGMENT

Figures in this e-chapter were borrowed from Williams Hematology, 7th edition, M Lichtman et al (eds). New York, McGraw-Hill, 2005; Hematology in General Practice, 4th edition, RS Hillman, KA Ault, New York, McGraw-Hill, 2005.

This page intentionally left blank

82 Principles of Human Genetics

J. Larry Jameson, Peter Kopp

IMPACT OF GENETICS AND GENOMICS ON MEDICAL PRACTICE

The prevalence of genetic diseases, combined with their potential severity and chronic nature, imposes great human, social, and financial burdens on society. Human genetics refers to the study of individual genes, their role and function in disease, and their mode of inheritance. Genomics refers to an organism's entire genetic information, the *genome*, and the function and interaction of DNA within the genome, as well as with environmental or nongenetic factors, such as a person's lifestyle. With the characterization of the human genome, genomics complements traditional genetics in our efforts to elucidate the etiology and pathogenesis of disease and to improve therapeutic interventions and outcomes. Following impressive advances in genetics, genomics, and health care information technology, the consequences of this wealth of knowledge for the practice of medicine are profound and play an increasingly prominent role in the diagnosis, prevention, and treatment of disease (Chap. 84).

Personalized medicine, the customization of medical decisions to an individual patient, relies heavily on genetic information. For example, a patient's genetic characteristics (genotype) can be used to optimize drug therapy and predict efficacy, adverse events, and drug dosing of selected medications (*pharmacogenetics*) (Chap. 5). The mutational profile of a malignancy allows the selection of therapies that target mutated or overexpressed signaling molecules. Although still investigational, genomic risk prediction models for common diseases are beginning to emerge.

Genetics has traditionally been viewed through the window of relatively rare single-gene diseases. These disorders account for ~10% of pediatric admissions and childhood mortality. Historically, genetics has focused predominantly on chromosomal and metabolic disorders, reflecting the long-standing availability of techniques to diagnose these conditions. For example, conditions such as trisomy 21 (Down's syndrome) or monosomy X (Turner's syndrome) can be diagnosed using cytogenetics (Chap. 83e). Likewise, many metabolic disorders (e.g., phenylketonuria, familial hypercholesterolemia) are diagnosed using biochemical analyses. The advances in DNA diagnostics have extended the field of genetics to include virtually all medical specialties and have led to the elucidation of the pathogenesis of numerous monogenic disorders. In addition, it is apparent that virtually every medical condition has a genetic component. As is often evident from a patient's family history, many common disorders such as hypertension, heart disease, asthma, diabetes mellitus, and mental illnesses are significantly influenced by the genetic background. These polygenic or multifactorial (complex) disorders involve the contributions of many different genes, as well as environmental factors that can modify disease risk (Chap. 84). Genome-wide association studies (GWAS) have elucidated numerous disease-associated loci and are providing novel insights into the allelic architecture of complex traits. These studies have been facilitated by the availability of comprehensive catalogues of human single-nucleotide polymorphism (SNP) haplotypes generated through the HapMap Project. The sequencing of whole genomes or exomes (the exons within the genome) is increasingly used in the clinical realm in order to characterize individuals with complex undiagnosed conditions or to characterize the mutational profile of advanced malignancies in order to select better targeted therapies.

Cancer has a genetic basis because it results from acquired somatic mutations in genes controlling growth, apoptosis, and cellular differentiation (Chap. 101e). In addition, the development of many cancers is associated with a hereditary predisposition. Characterization

of the genome (and epigenome) in various malignancies has led to fundamental new insights into cancer biology and reveals that the genomic profile of mutations is in many cases more important in determining the appropriate chemotherapy than the organ in which the tumor originates. Hence, comprehensive mutational profiling of malignancies has increasing impact on cancer taxonomy, the choice of targeted therapies, and improved outcomes.

Genetic and genomic approaches have proven invaluable for the detection of infectious pathogens and are used clinically to identify agents that are difficult to culture such as mycobacteria, viruses, and parasites, or to track infectious agents locally or globally. In many cases, molecular genetics has improved the feasibility and accuracy of diagnostic testing and is beginning to open new avenues for therapy, including gene and cellular therapy (Chaps. 90e and 91e). Molecular genetics has also provided the opportunity to characterize the *microbiome*, a new field that characterizes the population dynamics of bacteria, viruses, and parasites that coexist with humans and other animals (Chap. 86e). Emerging data indicate that the microbiome has significant effects on normal physiology as well as various disease states.

Molecular biology has significantly changed the treatment of human disease. Peptide hormones, growth factors, cytokines, and vaccines can now be produced in large amounts using recombinant DNA technology. Targeted modifications of these peptides provide the practitioner with improved therapeutic tools, as illustrated by genetically modified insulin analogues with more favorable kinetics. Lastly, there is reason to believe that a better understanding of the genetic basis of human disease will also have an increasing impact on disease prevention.

The astounding rate at which new genetic information is being generated creates a major challenge for physicians, health care providers, and basic investigators. Although many functional aspects of the genome remain unknown, there are many clinical situations where sufficient evidence exists for the use of genetic and genomic information to optimize patient care and treatment. Much genetic information resides in databases or is being published in basic science journals. Databases provide easy access to the expanding information about the human genome, genetic disease, and genetic testing (Table 82-1). For example, several thousand monogenic disorders are summarized in a large, continuously evolving compendium, referred to as the *Online Mendelian Inheritance in Man* (OMIM) catalogue (Table 82-1). The ongoing refinement of bioinformatics is simplifying the analysis and access to this daunting amount of new information.

THE HUMAN GENOME

Structure of the Human Genome • HUMAN GENOME PROJECT The Human Genome Project was initiated in the mid-1980s as an ambitious effort to characterize the entire human genome. Although the prospect of determining the complete sequence of the human genome seemed daunting several years ago, technical advances in DNA sequencing and bioinformatics led to the completion of a draft human sequence in 2000 and the completion of the DNA sequence for the last of the human chromosomes in May 2006. Currently, facilitated by rapidly decreasing costs for comprehensive sequence analyses and improvement of bioinformatics pipelines for data analysis, the sequencing of whole genomes and exomes is used with increasing frequency in the clinical setting. The scope of a whole genome sequence analysis can be illustrated by the following analogy. Human DNA consists of ~3 billion base pairs (bp) of DNA per haploid genome, which is nearly 1000-fold greater than that of the *Escherichia coli* genome. If the human DNA sequence were printed out, it would correspond to about 120 volumes of *Harrison's Principles of Internal Medicine*.

In addition to the human genome, the genomes of numerous organisms have been sequenced completely (~4000) or partially (~10,000) (Genomes Online Database [GOLD]; Table 82-1). They include, among others, eukaryotes such as the mouse (*Mus musculus*),

TABLE 82-1 SELECTED DATABASES RELEVANT FOR GENOMICS AND GENETIC DISORDERS

Site	URL	Comment
National Center for Biotechnology Information (NCBI)	http://www.ncbi.nlm.nih.gov/	Broad access to biomedical and genomic information, literature (PubMed), sequence databases, software for analyses of nucleotides and proteins Extensive links to other databases, genome resources, and tutorials
National Human Genome Research Institute	http://www.genome.gov/	An institute of the National Institutes of Health focused on genomic and genetic research; links providing information about the human genome sequence, genomes of other organisms, and genomic research
Catalog of Published Genome-Wide Association Studies	http://www.genome.gov/GWASudies/	Published high-resolution genome-wide association studies (GWAS)
Ensembl Genome browser	http://www.ensembl.org	Maps and sequence information of eukaryotic genomes
Online Mendelian Inheritance in Man	http://www.ncbi.nlm.nih.gov/omim	Online compendium of Mendelian disorders and human genes causing genetic disorders
Office of Biotechnology Activities, National Institutes of Health	http://oba.od.nih.gov/oba	Information about recombinant DNA and gene transfer; medical, ethical, legal, and social issues raised by genetic testing; medical, ethical, legal, and social issues raised by xenotransplantation
American College of Medical Genetics and Genomics	http://www.acmg.net/	Extensive links to other databases relevant for the diagnosis, treatment, and prevention of genetic disease
American Society of Human Genetics	http://www.ashg.org	Information about advances in genetic research, professional and public education, social and scientific policies
Cancer Genome Anatomy Project (CGAP)	http://cgap.nci.nih.gov/	Information about gene expression profiles of normal, precancer, and cancer cells
GeneTests	http://www.genetests.org/	International directory of genetic testing laboratories and prenatal diagnosis clinics; reviews and educational materials
Genomes Online Database (GOLD)	http://www.genomesonline.org/	Information on published and unpublished genomes
HUGO Gene Nomenclature	http://www.genenames.org/	Gene names and symbols
MITOMAP, a human mitochondrial genome database	http://www.mitomap.org/	A compendium of polymorphisms and mutations of the human mitochondrial DNA
International HapMap Project	http://www.hapmap.org/	Catalogue of haplotypes in different ethnic groups relevant for association studies and pharmacogenomics
ENCODE	http://www.genome.gov/10005107	Encyclopedia of DNA Elements; catalogue of all functional elements in the human genome
Dolan DNA Learning Center, Cold Spring Harbor Laboratories	http://www.dnalc.org/	Educational material about selected genetic disorders, DNA, eugenics, and genetic origin
The Online Metabolic and Molecular Bases of Inherited Disease (OMMBID)	http://www.ommbid.com/	Online version of the comprehensive text on the metabolic and molecular bases of inherited disease
Online Mendelian Inheritance in Animals (OMIA)	http://omia.angis.org.au/	Online compendium of Mendelian disorders in animals
The Jackson Laboratory	http://www.jax.org/ http://www.informatics.jax.org	Information about murine models and the mouse genome Mouse genome informatics

Note: Databases are evolving constantly. Pertinent information may be found by using links listed in the few selected databases.

Saccharomyces cerevisiae, *Caenorhabditis elegans*, and *Drosophila melanogaster*; bacteria (e.g., *E. coli*); and Archaea, viruses, organelles (mitochondria, chloroplasts), and plants (e.g., *Arabidopsis thaliana*). Genomic information of infectious agents has significant impact for the characterization of infectious outbreaks and epidemics. Other ramifications arising from the availability of genomic data include, among others, (1) the comparison of entire genomes (*comparative genomics*), (2) the study of large-scale expression of RNAs (*functional genomics*) and proteins (*proteomics*) to detect differences between various tissues in health and disease, (3) the characterization of the variation among individuals by establishing catalogues of sequence variations and SNPs (HapMap Project), and (4) the identification of genes that play critical roles in the development of polygenic and multifactorial disorders.

CHROMOSOMES The human genome is divided into 23 different chromosomes, including 22 autosomes (numbered 1–22) and the X and Y sex chromosomes (Fig. 82-1). Adult cells are *diploid*, meaning they contain two homologous sets of 22 autosomes and a pair of sex chromosomes. Females have two X chromosomes (XX), whereas males have one X and one Y chromosome (XY). As a consequence of meiosis, germ cells (sperm or oocytes) are haploid and contain one set of 22 autosomes and one of the sex chromosomes. At the time of fertilization, the diploid genome is reconstituted by pairing of the homologous chromosomes from the mother and father. With each cell division

(mitosis), chromosomes are replicated, paired, segregated, and divided into two daughter cells.

STRUCTURE OF DNA DNA is a double-stranded helix composed of four different bases: adenine (A), thymidine (T), guanine (G), and cytosine (C). Adenine is paired to thymidine, and guanine is paired to cytosine, by hydrogen bond interactions that span the double helix (Fig. 82-1). DNA has several remarkable features that make it ideal for the transmission of genetic information. It is relatively stable, and the double-stranded nature of DNA and its feature of strict base-pair complementarity permit faithful replication during cell division. Complementarity also allows the transmission of genetic information from DNA → RNA → protein (Fig. 82-2). mRNA is encoded by the so-called sense or coding strand of the DNA double helix and is translated into proteins by ribosomes.

The presence of four different bases provides surprising genetic diversity. In the protein-coding regions of genes, the DNA bases are arranged into codons, a triplet of bases that specifies a particular amino acid. It is possible to arrange the four bases into 64 different triplet codons (4³). Each codon specifies 1 of the 20 different amino acids, or a regulatory signal such as initiation and stop of translation. Because there are more codons than amino acids, the genetic code is degenerate; that is, most amino acids can be specified by several different codons. By arranging the codons in different combinations and in

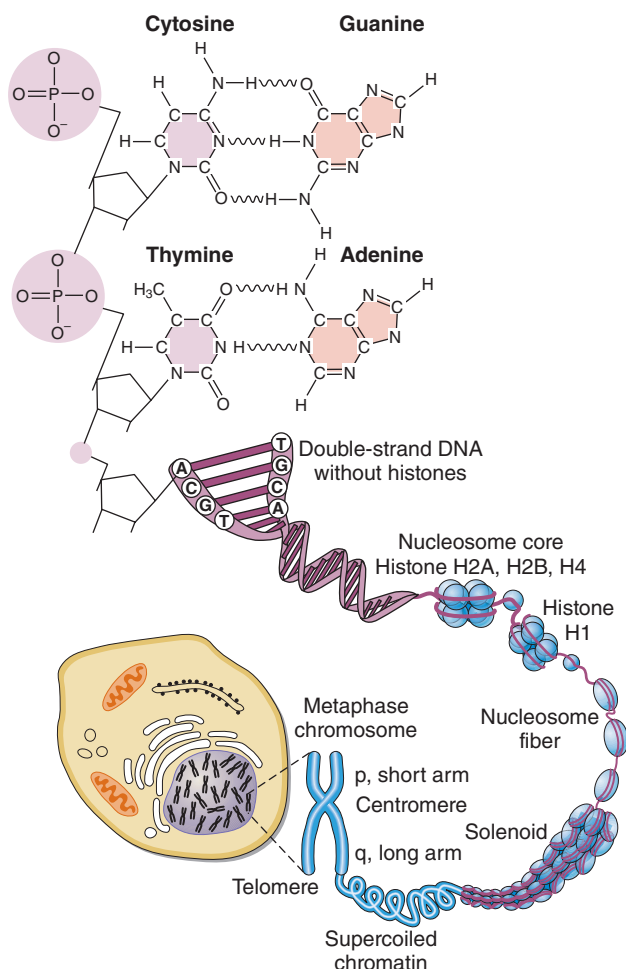


FIGURE 82-1 Structure of chromatin and chromosomes.

Chromatin is composed of double-strand DNA that is wrapped around histone and nonhistone proteins forming nucleosomes. The nucleosomes are further organized into solenoid structures. Chromosomes assume their characteristic structure, with short (p) and long (q) arms at the metaphase stage of the cell cycle.

various lengths, it is possible to generate the tremendous diversity of primary protein structure.

DNA length is normally measured in units of 1000 bp (kilobases, kb) or 1,000,000 bp (megabases, Mb). Not all DNA encodes genes. In fact, genes account for only ~10–15% of DNA. Much of the remaining DNA consists of sequences, often of highly repetitive nature, the function of which is poorly understood. These repetitive DNA regions, along with nonrepetitive sequences that do not encode genes, serve, in part, a structural role in the packaging of DNA into chromatin (i.e., DNA bound to histone proteins, and chromosomes) and exert regulatory functions (Fig. 82-1).

GENES A *gene* is a functional unit that is regulated by transcription (see below) and encodes an RNA product, which is most commonly, but not always, translated into a protein that exerts activity within or outside the cell (Fig. 82-3). Historically, genes were identified because they conferred specific traits that are transmitted from one generation to the next. Increasingly, they are characterized based on expression in various tissues (*transcriptome*). The size of genes is quite broad; some genes are only a few hundred base pairs, whereas others are extraordinarily large (2 Mb). The number of genes greatly underestimates the complexity of genetic expression, because single genes can generate multiple spliced messenger RNA (mRNA) products (*isoforms*), which are translated into proteins that are subject to complex posttranslational modification such as phosphorylation. *Exons* refer to the portion of genes that are eventually spliced together to form mRNA. *Introns* refer to the spacing regions between the exons that are spliced out of precursor RNAs during RNA processing. The gene locus also includes

regions that are necessary to control its expression (Fig. 82-2). Current estimates predict 20,687 protein-coding genes in the human genome with an average of about four different coding transcripts per gene. Remarkably, the exome only constitutes 1.14% of the genome. In addition, thousands of noncoding transcripts (RNAs of various length such as microRNAs and long noncoding RNAs), which function, at least in part, as transcriptional and posttranscriptional regulators of gene expression, have been identified. Aberrant expression of microRNAs has been found to play a pathogenic role in numerous diseases.

SINGLE-NUCLEOTIDE POLYMORPHISMS An SNP is a variation of a single base pair in the DNA. The identification of the ~10 million SNPs estimated to occur in the human genome has generated a catalogue of common genetic variants that occur in human beings from distinct ethnic backgrounds (Fig. 82-3). SNPs are the most common type of sequence variation and account for ~90% of all sequence variation. They occur on average every 100 to 300 bases and are the major source of genetic heterogeneity. Remarkably, however, the primary DNA sequence of humans has ~99.9% similarity compared to that of any other human. SNPs that are in close proximity are inherited together (e.g., they are linked) and are referred to as *haplotypes* (Fig. 82-4). The HapMap describes the nature and location of these SNP haplotypes and how they are distributed among individuals within and among populations. The haplotype map information, referred to as HapMap, is greatly facilitating GWAS designed to elucidate the complex interactions among multiple genes and lifestyle factors in multifactorial disorders (see below). Moreover, haplotype analyses are useful to assess variations in responses to medications (*pharmacogenomics*) and environmental factors, as well as the prediction of disease predisposition.

COPY NUMBER VARIATIONS Copy number variations (CNVs) are relatively large genomic regions (1 kb to several Mb) that have been duplicated or deleted on certain chromosomes (Fig. 82-5). It has been estimated that as many as 1500 CNVs, scattered throughout the genome, are present in an individual. When comparing the genomes of two individuals, approximately 0.4–0.8% of their genomes differ in terms of CNVs. Of note, *de novo* CNVs have been observed between monozygotic twins, who otherwise have identical genomes. Some CNVs have been associated with susceptibility or resistance to disease, and CNVs can be elevated in cancer cells.

Replication of DNA and Mitosis Genetic information in DNA is transmitted to daughter cells under two different circumstances: (1) somatic cells divide by *mitosis*, allowing the diploid ($2n$) genome to replicate itself completely in conjunction with cell division; and (2) germ cells (sperm and ova) undergo *meiosis*, a process that enables the reduction of the diploid ($2n$) set of chromosomes to the haploid state ($1n$).

Prior to mitosis, cells exit the resting, or G_0 state, and enter the cell cycle (Chap. 101e). After traversing a critical checkpoint in G_1 , cells undergo DNA synthesis (S phase), during which the DNA in each chromosome is replicated, yielding two pairs of sister chromatids ($2n \rightarrow 4n$). The process of DNA synthesis requires stringent fidelity in order to avoid transmitting errors to subsequent generations of cells. Genetic abnormalities of DNA mismatch/repair include xeroderma pigmentosum, Bloom's syndrome, ataxia telangiectasia, and hereditary nonpolyposis colon cancer (HNPCC), among others. Many of these disorders strongly predispose to neoplasia because of the rapid acquisition of additional mutations (Chap. 101e). After completion of DNA synthesis, cells enter G_2 and progress through a second checkpoint before entering mitosis. At this stage, the chromosomes condense and are aligned along the equatorial plate at metaphase. The two identical sister chromatids, held together at the centromere, divide and migrate to opposite poles of the cell. After formation of a nuclear membrane around the two separated sets of chromatids, the cell divides and two daughter cells are formed, thus restoring the diploid ($2n$) state.

Assortment and Segregation of Genes During Meiosis Meiosis occurs only in germ cells of the gonads. It shares certain features with mitosis but involves two distinct steps of cell division that reduce the chromosome number to the haploid state. In addition, there is active recombination that generates genetic diversity. During the first cell division, two

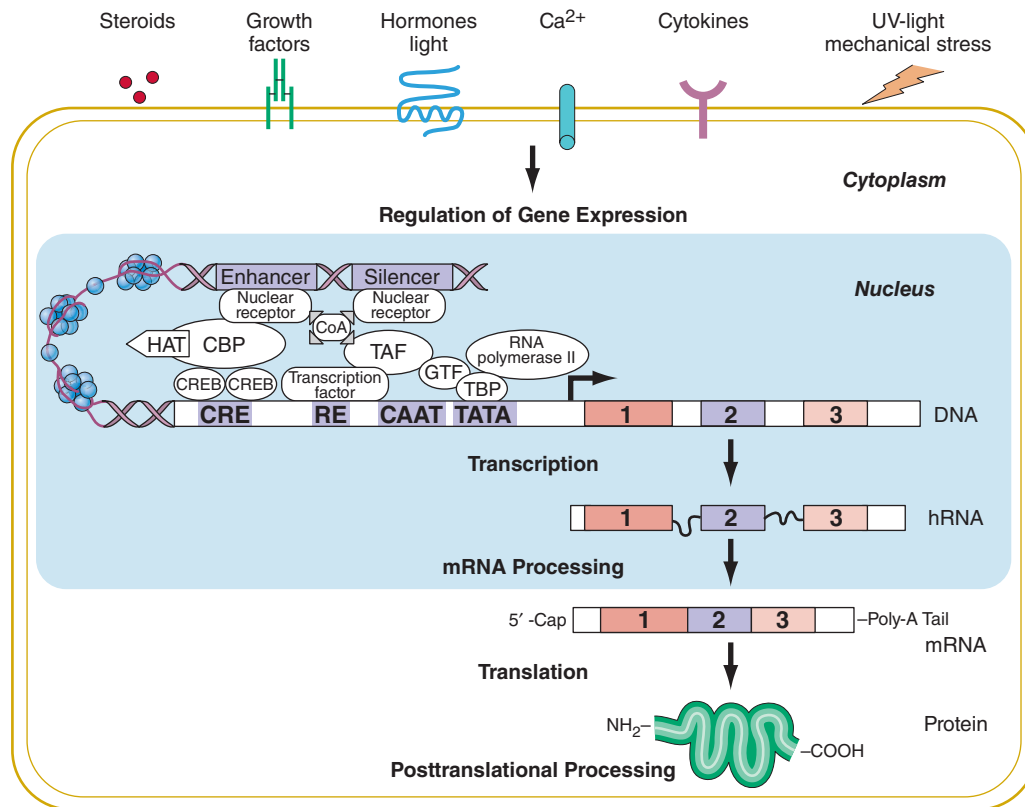


FIGURE 82-2 Flow of genetic information. Multiple extracellular signals activate intracellular signal cascades that result in altered regulation of gene expression through the interaction of transcription factors with regulatory regions of genes. RNA polymerase transcribes DNA into RNA that is processed to mRNA by excision of intronic sequences. The mRNA is translated into a polypeptide chain to form the mature protein after undergoing posttranslational processing. CBP, CREB-binding protein; CoA, co-activator; COOH, carboxyterminus; CRE, cyclic AMP responsive element; CREB, cyclic AMP response element-binding protein; GTF, general transcription factors; HAT, histone acetyl transferase; NH₂, aminotermi- nus; RE, response element; TAF, TBP-associated factors; TATA, TATA box; TBP, TATA-binding protein.

sister chromatids ($2n \rightarrow 4n$) are formed for each chromosome pair and there is an exchange of DNA between homologous paternal and maternal chromosomes. This process involves the formation of *chiasmata*, structures that correspond to the DNA segments that cross over between the maternal and paternal homologues (Fig. 82-6). Usually there is at least one crossover on each chromosomal arm; recombination occurs more frequently in female meiosis than in male meiosis. Subsequently, the chromosomes segregate randomly. Because there are 23 chromosomes, there exist 2^{23} (>8 million) possible combinations of chromosomes. Together with the genetic exchanges that occur during recombination, chromosomal segregation generates tremendous diversity, and each gamete is genetically unique. The process of recombination and the independent segregation of chromosomes provide the foundation for performing linkage analyses, whereby one attempts to correlate the inheritance of certain chromosomal regions (or linked genes) with the presence of a disease or genetic trait (see below).

After the first meiotic division, which results in two daughter cells ($2n$), the two chromatids of each chromosome separate during a second meiotic division to yield four gametes with a haploid state ($1n$). When the egg is fertilized by sperm, the two haploid sets are combined, thereby restoring the diploid state ($2n$) in the zygote.

REGULATION OF GENE EXPRESSION

Regulation by Transcription Factors The expression of genes is regulated by DNA-binding proteins that activate or repress transcription. The number of DNA sequences and transcription factors that regulate transcription is much greater than originally anticipated. Most genes contain at least 15–20 discrete regulatory elements within 300 bp of the transcription start site. This densely packed promoter region often contains binding sites for ubiquitous transcription factors such as CAAT box/enhancer binding protein (C/EBP), cyclic AMP response element-binding (CREB) protein, selective promoter factor 1 (Sp-1),

or activator protein 1 (AP-1). However, factors involved in cell-specific expression may also bind to these sequences. Key regulatory elements may also reside at a large distance from the proximal promoter. The globin and the immunoglobulin genes, for example, contain *locus control regions* that are several kilobases away from the structural sequences of the gene. Specific groups of transcription factors that bind to these promoter and enhancer sequences provide a combinatorial code for regulating transcription. In this manner, relatively ubiquitous factors interact with more restricted factors to allow each gene to be expressed and regulated in a unique manner that is dependent on developmental state, cell type, and numerous extracellular stimuli. Regulatory factors also bind within the gene itself, particularly in the intronic regions. The transcription factors that bind to DNA actually represent only the first level of regulatory control. Other proteins—*co-activators* and *co-repressors*—interact with the DNA-binding transcription factors to generate large regulatory complexes. These complexes are subject to control by numerous cell-signaling pathways and enzymes, leading to phosphorylation, acetylation, sumoylation, and ubiquitination. Ultimately, the recruited transcription factors interact with, and stabilize, components of the basal transcription complex that assembles at the site of the TATA box and initiator region. This basal transcription factor complex consists of >30 different proteins. Gene transcription occurs when RNA polymerase begins to synthesize RNA from the DNA template. A large number of identified genetic diseases involve transcription factors (Table 82-2).

The field of *functional genomics* is based on the concept that understanding alterations of gene expression under various physiologic and pathologic conditions provides insight into the underlying functional role of the gene. By revealing specific gene expression profiles, this knowledge may be of diagnostic and therapeutic relevance. The large-scale study of expression profiles, which takes advantage of microarray and bead array technologies, is also referred to as *transcriptomics*

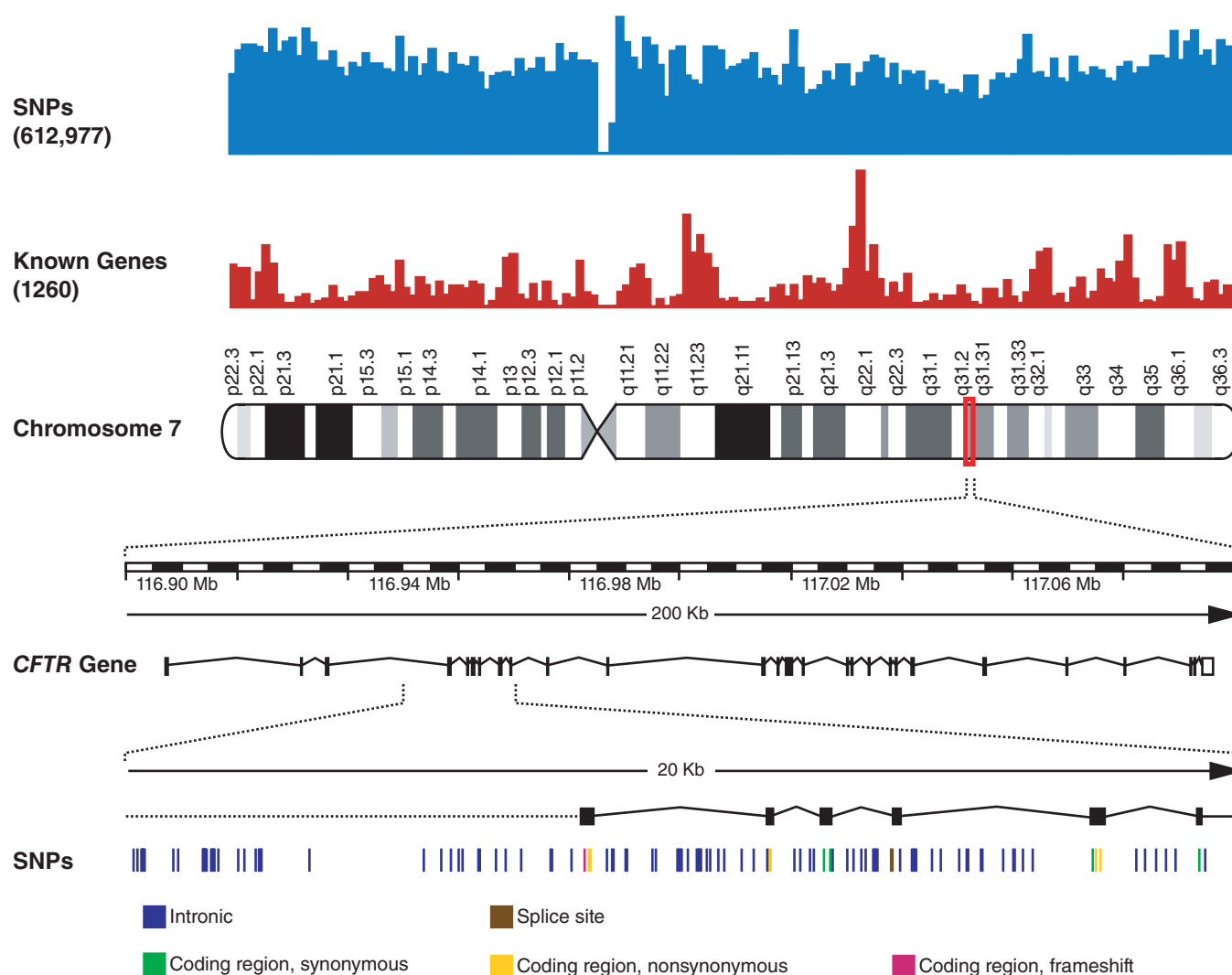


FIGURE 82-3 Chromosome 7 is shown with the density of single-nucleotide polymorphisms (SNPs) and genes above. A 200-kb region in 7q31.2 containing the *CFTR* gene is shown below. The *CFTR* gene contains 27 exons. More than 1900 mutations in this gene have been found in patients with cystic fibrosis. A 20-kb region encompassing exons 4–9 is shown further amplified to illustrate the SNPs in this region.

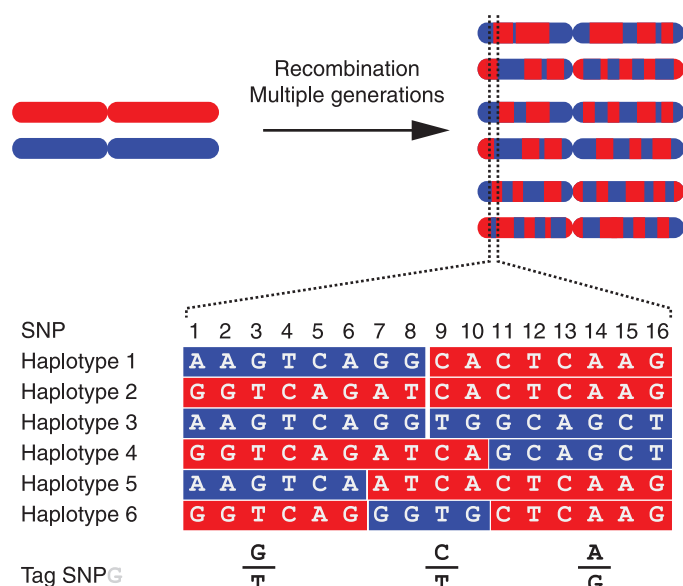


FIGURE 82-4 The origin of haplotypes is due to repeated recombination events occurring in multiple generations. Over time, this leads to distinct haplotypes. These haplotype blocks can often be characterized by genotyping selected Tag single-nucleotide polymorphisms (SNPs), an approach that facilitates performing genome-wide association studies (GWAS).

because the complement of mRNAs transcribed by the cellular genome is called the *transcriptome*.

Most studies of gene expression have focused on the regulatory DNA elements that control transcription. However, it should be emphasized that gene expression requires a series of steps, including mRNA processing, protein translation, and posttranslational modifications, all of which are actively regulated (Fig. 82-2).

Epigenetic Regulation of Gene Expression *Epigenetics* describes mechanisms and phenotypic changes that are not a result of variation in the primary DNA nucleotide sequence, but are caused by secondary modifications of DNA or histones. These modifications include heritable changes such as X-inactivation and imprinting, but they can also result from dynamic posttranslational protein modifications in response to environmental influences such as diet, age, or drugs. The epigenetic modifications result in altered expression of individual genes or chromosomal loci encompassing multiple genes. The term *epigenome* describes the constellation of covalent modifications of DNA and histones that impact chromatin structure, as well as non-coding transcripts that modulate the transcriptional activity of DNA. Although the primary DNA sequence is usually identical in all cells of an organism, tissue-specific changes in the epigenome contribute to determining the transcriptional signature of a cell (transcriptome) and hence the protein expression profile (proteome).

Mechanistically, DNA and histone modifications can result in the activation or silencing of gene expression (Fig. 82-7). DNA methylation involves the addition of a methyl group to cytosine residues. This is

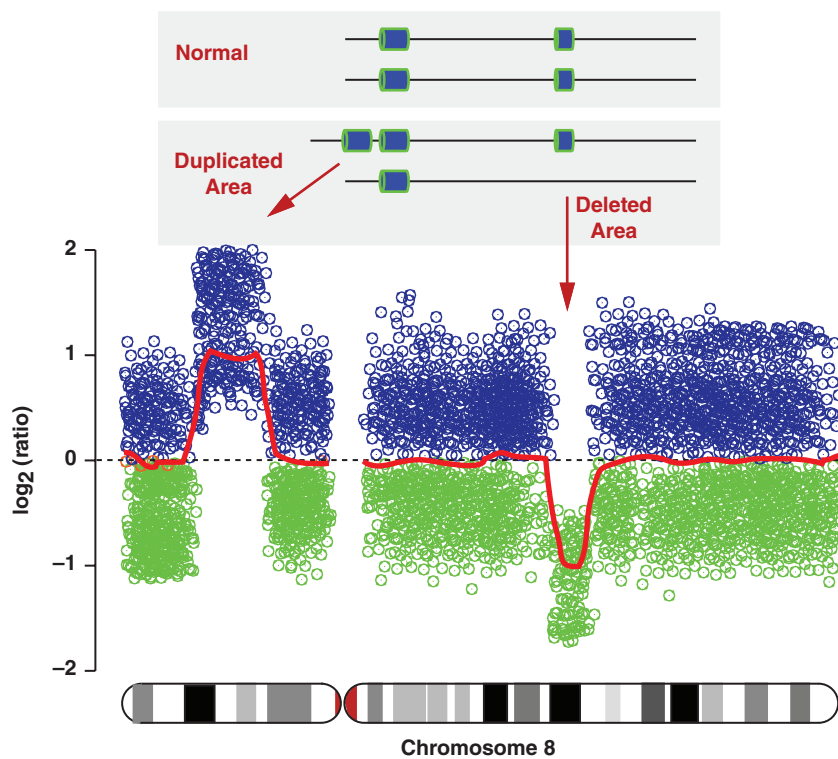


FIGURE 82-5 Copy number variations (CNV) encompass relatively large regions of the genome that have been duplicated or deleted. Chromosome 8 is shown with CNV detected by genomic hybridization. An increase in the signal strength indicates a duplication, a decrease reflects a deletion of the covered chromosomal regions.

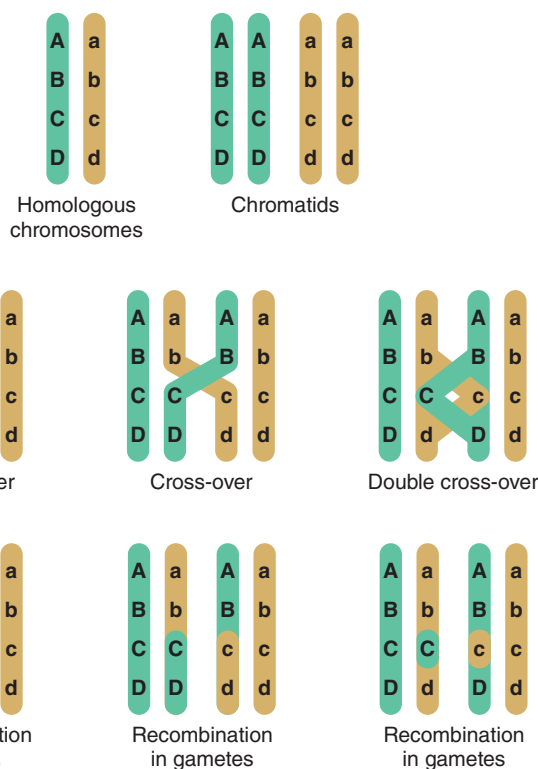


FIGURE 82-6 Crossing-over and genetic recombination. During chiasma formation, either of the two sister chromatids on one chromosome pairs with one of the chromatids of the homologous chromosome. Genetic recombination occurs through crossing-over and results in recombinant and nonrecombinant chromosome segments in the gametes. Together with the random segregation of the maternal and paternal chromosomes, recombination contributes to genetic diversity and forms the basis of the concept of linkage.

usually restricted to cytosines of CpG dinucleotides, which are abundant throughout the genome. Methylation of these dinucleotides is thought to represent a defense mechanism that minimizes the expression of sequences that have been incorporated into the genome such as retroviral sequences. CpG dinucleotides also exist in so-called *CpG islands*, stretches of DNA characterized by a high CG content, which are found in the majority of human gene promoters. CpG islands in promoter regions are typically unmethylated, and the lack of methylation facilitates transcription.

Histone methylation involves the addition of a methyl group to lysine residues in histone proteins (Fig. 82-7). Depending on the specific lysine residue being methylated, this alters chromatin configuration, either making it more open or tightly packed. Acetylation of histone proteins is another well-characterized mechanism that results in an open chromatin configuration, which favors active transcription. Acetylation is generally more dynamic than methylation, and many transcriptional activation complexes have histone acetylase activity, whereas repressor complexes often contain deacetylases and remove acetyl groups from histones. Other histone modifications, whose effects are incompletely characterized, include phosphorylation and sumoylation. Lastly, noncoding RNAs that bind to DNA can have a significant impact on transcriptional activity.

Physiologically, epigenetic mechanisms play an important role in several instances. For example, X-inactivation refers to the relative silencing of one of the two X chromosome copies present in females. The inactivation process is a form of dosage compensation such that females (XX) do not generally express twice as many X-chromosomal gene products as males (XY). In a given cell, the choice of which chromosome is inactivated occurs randomly in humans. But once the maternal or paternal X chromosome is inactivated, it will remain inactive, and this information is transmitted with each cell division. The *X-inactive specific transcript (Xist)* gene encodes a large noncoding RNA that mediates the silencing of the X chromosome from which it is transcribed by coating it with Xist RNA. The inactive X chromosome is highly methylated and has low levels of histone acetylation.

Epigenetic gene inactivation also occurs on selected chromosomal regions of autosomes, a phenomenon referred to as *genomic imprinting*. Through this mechanism, a small subset of genes is only expressed in a monoallelic fashion. Imprinting is heritable and leads to the preferential expression of one of the parental alleles, which deviates from the usual biallelic expression seen for the majority of genes. Remarkably, imprinting can be limited to a subset of tissues. Imprinting is mediated through DNA methylation of one of the alleles. The epigenetic marks on imprinted genes are maintained throughout life, but during zygote formation, they are activated or inactivated in a sex-specific manner (imprint reset) (Fig. 82-8), which allows a differential expression pattern in the fertilized egg and the subsequent mitotic divisions. Appropriate expression of imprinted genes is important for normal development and cellular functions. Imprinting defects and uniparental disomy, which is the inheritance of two chromosomes or chromosomal regions from the same parent, are the cause of several developmental disorders such as Beckwith-Wiedemann syndrome, Silver-Russell syndrome, Angelman's syndrome, and Prader-Willi syndrome (see below). Monoallelic loss-of-function mutations in the *GNAS1* gene lead to Albright's hereditary osteodystrophy (AHO). Paternal transmission of *GNAS1* mutations leads to an isolated AHO phenotype (pseudopseudohypoparathyroidism), whereas maternal transmission leads to AHO in combination with hormone resistance to parathyroid hormone, thyrotropin, and gonadotropins (pseudohypoparathyroidism type IA). These phenotypic differences are explained by tissue-specific imprinting of the *GNAS1* gene, which is expressed primarily from the maternal allele in the thyroid, gonadotropes, and the proximal renal tubule. In most other tissues, the *GNAS1* gene is expressed biallelically.

TABLE 82-2 SELECTED EXAMPLES OF DISEASES CAUSED BY MUTATIONS AND REARRANGEMENTS IN TRANSCRIPTION FACTOR CLASSES

Transcription Factor Class	Example	Associated Disorder
Nuclear receptors	Androgen receptor	Complete or partial androgen insensitivity (recessive missense mutations) Spinobulbar muscular atrophy (CAG repeat expansion)
Zinc finger proteins	WT1	WAGR syndrome: Wilms' tumor, aniridia, genitourinary malformations, mental retardation
Basic helix-loop-helix	MITF	Waardenburg's syndrome type 2A
Homeobox	IPF1	Maturity onset of diabetes mellitus type 4 (heterozygous mutation/haploinsufficiency) Pancreatic agenesis (homozygous mutation)
Leucine zipper	Retina leucine zipper (NRL)	Autosomal dominant retinitis pigmentosa
High mobility group (HMG) proteins	SRY	Sex reversal
Forkhead	HNF4 α , HNF1 α , HNF1 β	Maturity onset of diabetes mellitus types 1, 3, 5
Paired box	PAX3	Waardenburg's syndrome types 1 and 3
T-box	TBX5	Holt-Oram syndrome (thumb anomalies, atrial or ventricular septum defects, phocomelia)
Cell cycle control proteins	P53	Li-Fraumeni syndrome, other cancers
Co-activators	CREB binding protein (CBP)	Rubinstein-Taybi syndrome
General transcription factors	TATA-binding protein (TBP)	Spinocerebellar ataxia 17 (CAG expansion)
Transcription elongation factor	VHL	von Hippel-Lindau syndrome (renal cell carcinoma, pheochromocytoma, pancreatic tumors, hemangioblastomas) Autosomal dominant inheritance, somatic inactivation of second allele (Knudson two-hit model)
Runt	CBFA2	Familial thrombocytopenia with propensity to acute myelogenous leukemia
Chimeric proteins due to translocations	PML-RAR	Acute promyelocytic leukemia t(15;17)(q22;q11.2-q12) translocation

Abbreviations: CREB, cAMP responsive element-binding protein; HNF, hepatocyte nuclear factor; PML, promyelocytic leukemia; RAR, retinoic acid receptor; SRY, sex-determining region Y; VHL, von Hippel-Lindau.

In patients with isolated renal resistance to parathyroid hormone (pseudohypoparathyroidism type IB), defective imprinting of the *GNAS1* gene results in decreased Gsa expression in the proximal renal tubules. Rett's syndrome is an X-linked dominant disorder resulting in developmental regression and stereotypic hand movements in affected girls.

It is caused by mutations in the *MECP2* gene, which encodes a methyl-binding protein. The ensuing aberrant methylation results in abnormal gene expression in neurons, which are otherwise normally developed.

Remarkably, epigenetic differences also occur among monozygotic twins. Although twins are epigenetically indistinguishable during the early years of life, older monozygotic twins exhibit differences in the overall content and genomic distribution of DNA methylation and histone acetylation, which would be expected to alter gene expression in various tissues.

In cancer, the epigenome is characterized by simultaneous losses and gains of DNA methylation in different genomic regions, as well as repressive histone modifications. Hyper- and hypomethylation are associated with mutations in genes that control DNA methylation. Hypomethylation is thought to remove normal control mechanisms that prevent expression of repressed DNA regions. It is also associated with genomic instability. Hypermethylation, in contrast, results in the silencing of CpG islands in promoter regions of genes, including tumor-suppressor genes. Epigenetic alterations are considered to be more easily reversible compared to genetic changes, and modification of the epigenome with demethylating agents and histone deacetylases is being explored in clinical trials.

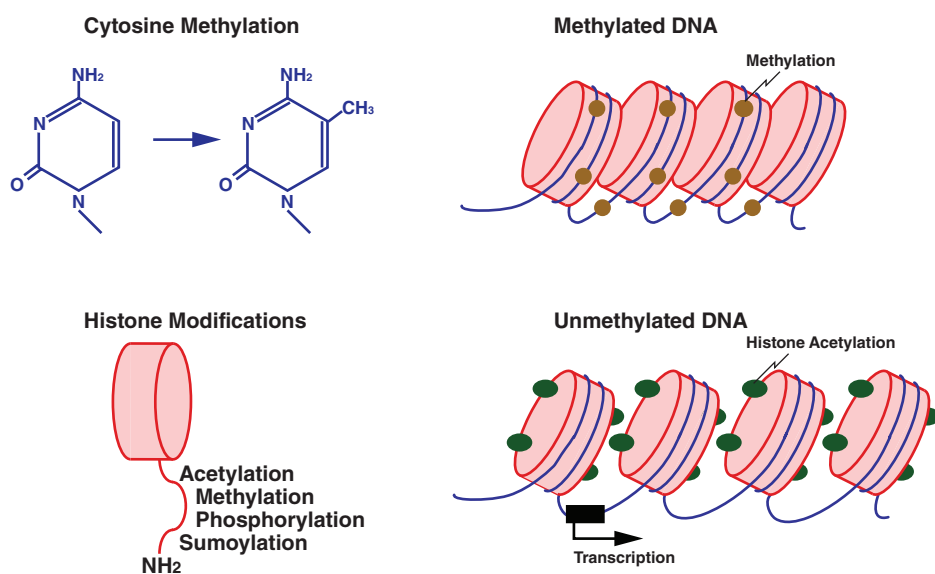


FIGURE 82-7 Epigenetic modifications of DNA and histones. Methylation of cytosine residues is associated with gene silencing. Methylation of certain genomic regions is inherited (imprinting), and it is involved in the silencing of one of the two X chromosomes in females (X-inactivation). Alterations in methylation can also be acquired, e.g., in cancer cells. Covalent posttranslational modifications of histones play an important role in altering DNA accessibility and chromatin structure and hence in regulating transcription. Histones can be reversibly modified in their amino-terminal tails, which protrude from the nucleosome core particle, by acetylation of lysine, phosphorylation of serine, methylation of lysine and arginine residues, and sumoylation. Acetylation of histones by histone acetylases (HATs), e.g., leads to unwinding of chromatin and accessibility to transcription factors. Conversely, deacetylation by histone deacetylases (HDACs) results in a compact chromatin structure and silencing of transcription.

MODELS OF GENETIC DISEASE

Several organisms have been studied extensively as genetic models, including *M. musculus* (mouse), *D. melanogaster* (fruit fly), *C. elegans* (nematode), *S. cerevisiae* (baker's yeast), and *E. coli* (colonic bacterium). The ability to use these evolutionarily distant organisms as genetic models that are relevant

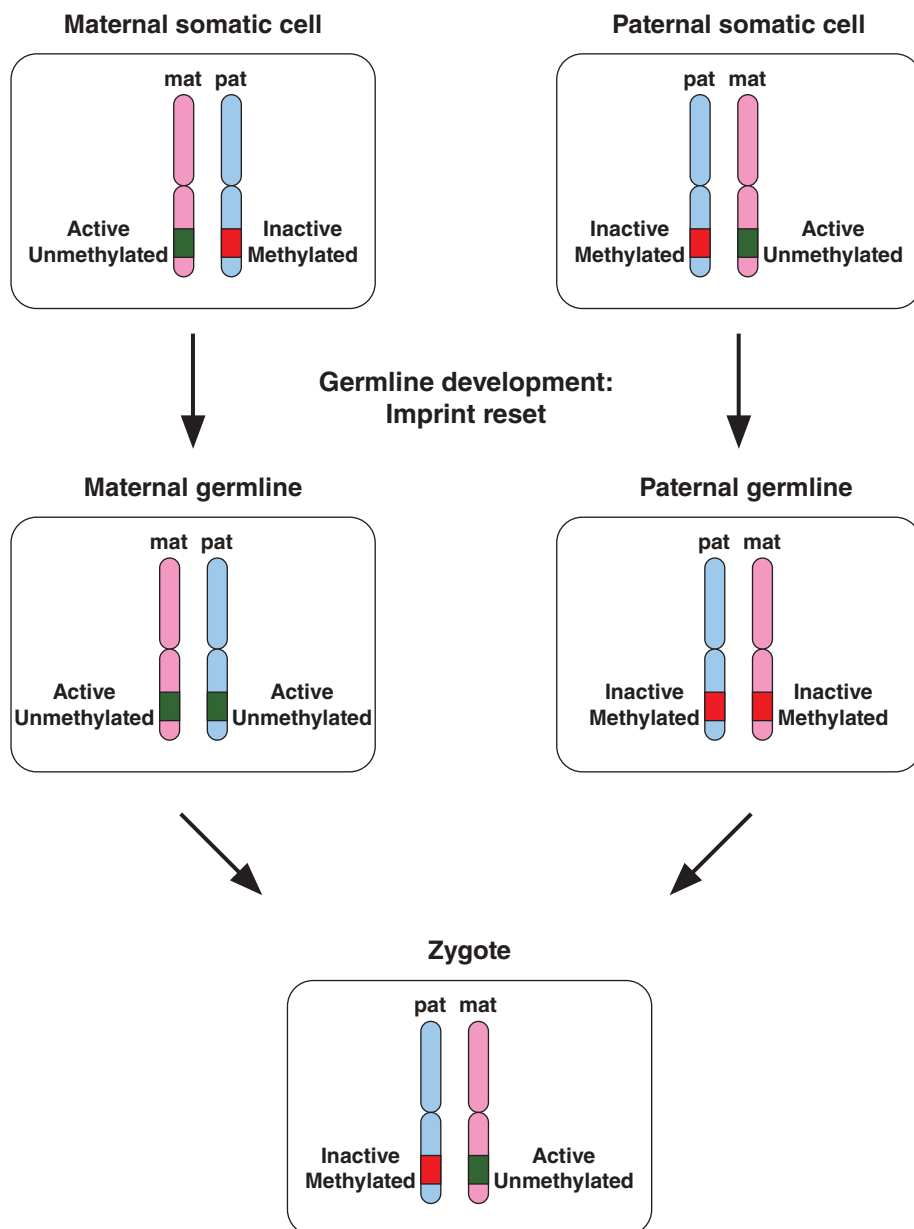


FIGURE 82-8 A few genomic regions are imprinted in a parent-specific fashion. The unmethylated chromosomal regions are actively expressed, whereas the methylated regions are silenced. In the germline, the imprint is reset in a parent-specific fashion: both chromosomes are unmethylated in the maternal (mat) germline and methylated in the paternal (pat) germline. In the zygote, the resulting imprinting pattern is identical with the pattern in the somatic cells of the parents.

to human physiology reflects a surprising conservation of genetic pathways and gene function. Transgenic mouse models have been particularly valuable, because many human and mouse genes exhibit similar structure and function and because manipulation of the mouse genome is relatively straightforward compared to that of other mammalian species. Transgenic strategies in mice can be divided into two main approaches: (1) expression of a gene by random insertion into the genome, and (2) deletion or targeted mutagenesis of a gene by homologous recombination with the native endogenous gene (knock-out, knock-in). Previous versions of this chapter provide more detail about the technical principles underlying the development of genetically modified animals. Several databases provide comprehensive information about natural and transgenic animal models, the associated phenotypes, and integrated genetic, genomic, and biologic data (Table 82-1).

TRANSMISSION OF GENETIC DISEASE

Origins and Types of Mutations A *mutation* can be defined as any change in the primary nucleotide sequence of DNA regardless of

its functional consequences. Some mutations may be lethal, others are less deleterious, and some may confer an evolutionary advantage. Mutations can occur in the germline (sperm or oocytes); these can be transmitted to progeny. Alternatively, mutations can occur during embryogenesis or in somatic tissues. Mutations that occur during development lead to *mosaicism*, a situation in which tissues are composed of cells with different genetic constitutions. If the germline is mosaic, a mutation can be transmitted to some progeny but not others, which sometimes leads to confusion in assessing the pattern of inheritance. Somatic mutations that do not affect cell survival can sometimes be detected because of variable phenotypic effects in tissues (e.g., pigmented lesions in McCune-Albright syndrome). Other somatic mutations are associated with neoplasia because they confer a growth advantage to cells. Epigenetic events may also influence gene expression or facilitate genetic damage. With the exception of triplet nucleotide repeats, which can expand (see below), mutations are usually stable.

Mutations are structurally diverse—they can involve the entire genome, as in triploidy (one extra set of chromosomes), or gross numerical or structural alterations in chromosomes or individual genes (Chap. 83e). Large deletions may affect a portion of a gene or an entire gene, or, if several genes are involved, they may lead to a *contiguous gene syndrome*. Unequal crossing-over between homologous genes can result in fusion gene mutations, as illustrated by color blindness. Mutations involving single nucleotides are referred to as *point mutations*. Substitutions are called *transitions* if a purine is replaced by another purine base ($A \leftrightarrow G$) or if a pyrimidine is replaced by another pyrimidine ($C \leftrightarrow T$). Changes from a purine to a pyrimidine, or vice versa, are referred to as *transversions*. If the DNA sequence change occurs in a coding region and alters an amino acid, it is called a *missense mutation*. Depending on the functional consequences of such a missense mutation, amino acid substitutions in different regions of the protein can lead to distinct phenotypes.

Mutations can occur in all domains of a gene (Fig. 82-9). A point mutation occurring within the coding region leads to an amino acid substitution if the codon is altered (Fig. 82-10). Point mutations that introduce a premature stop codon result in a truncated protein. Large deletions may affect a portion of a gene or an entire gene, whereas small deletions and insertions alter the reading frame if they do not represent a multiple of three bases. These “frameshift” mutations lead to an entirely altered carboxy terminus. Mutations in intronic sequences or in exon junctions may destroy or create splice donor or splice acceptor sites. Mutations may also be found in the regulatory sequences of genes, resulting in reduced or enhanced gene transcription.

Certain DNA sequences are particularly susceptible to mutagenesis. Successive pyrimidine residues (e.g., T-T or C-C) are subject to the formation of ultraviolet light-induced photoadducts. If these pyrimidine dimers are not repaired by the nucleotide excision repair pathway, mutations will be introduced after DNA synthesis. The dinucleotide C-G, or CpG, is also a hot spot for a specific type of mutation. In this case, methylation of the cytosine is associated with

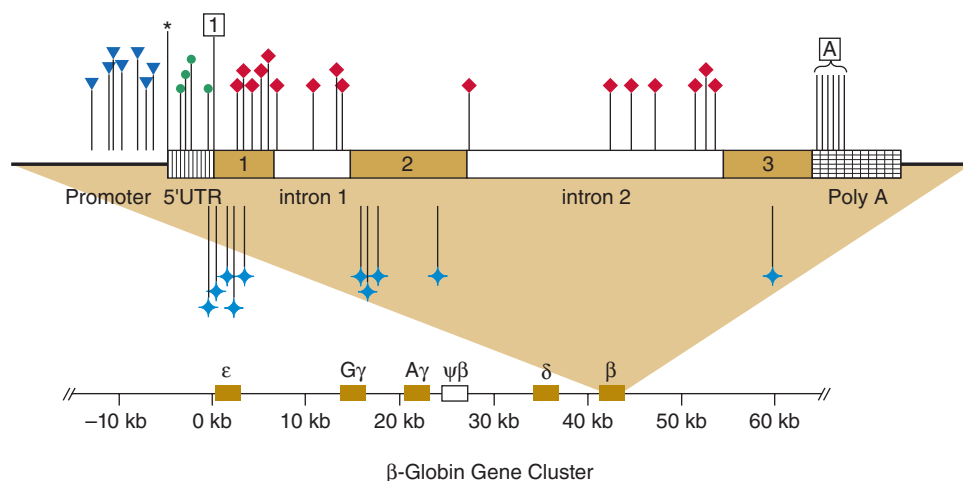


FIGURE 82-9 Point mutations causing β thalassemia as example of allelic heterogeneity. The β -globin gene is located in the globin gene cluster. Point mutations can be located in the promoter, the CAP site, the 5'-untranslated region, the initiation codon, each of the three exons, the introns, or the polyadenylation signal. Many mutations introduce missense or nonsense mutations, whereas others cause defective RNA splicing. Not shown here are deletion mutations of the β -globin gene or larger deletions of the globin locus that can also result in thalassemia. ▼, promoter mutations; *, CAP site; •, 5'UTR; □, initiation codon; ◆, defective RNA processing; ◆, missense and nonsense mutations; ▭, Poly A signal.

an enhanced rate of deamination to uracil, which is then replaced with thymine. This C \rightarrow T transition (or G \rightarrow A on the opposite strand) accounts for at least one-third of point mutations associated with polymorphisms and mutations. In addition to the fact that certain types of mutations (C \rightarrow T or G \rightarrow A) are relatively common, the nature of the genetic code also results in overrepresentation of certain amino acid substitutions.

Polymorphisms are sequence variations that have a frequency of at least 1%. Usually, they do not result in a perceptible phenotype. Often they consist of single base-pair substitutions that do not alter the protein coding sequence because of the degenerate nature of the genetic code (synonymous polymorphism), although it is possible that some might alter mRNA stability, translation, or the amino acid sequence (nonsynonymous polymorphism) (Fig. 82-10). The detection of sequence variants poses a practical problem because

mutations is much greater in the male germline than the female germline, in which rates of aneuploidy are increased (Chap. 83e). Thus, the incidence of new point mutations in spermatogonia increases with paternal age (e.g., achondroplasia, Marfan's syndrome, neurofibromatosis). It is estimated that about 1 in 10 sperm carries a new deleterious mutation. The rates for new mutations are calculated most readily for autosomal dominant and X-linked disorders and are $\sim 10^{-5}$ – 10^{-6} /locus per generation. Because most monogenic diseases are relatively rare, new mutations account for a significant fraction of cases. This is important in the context of genetic counseling, because a new mutation can be transmitted to the affected individual but does not necessarily imply that the parents are at risk to transmit the disease to other children. An exception to this is when the new mutation occurs early in germline development, leading to *gonadal mosaicism*.

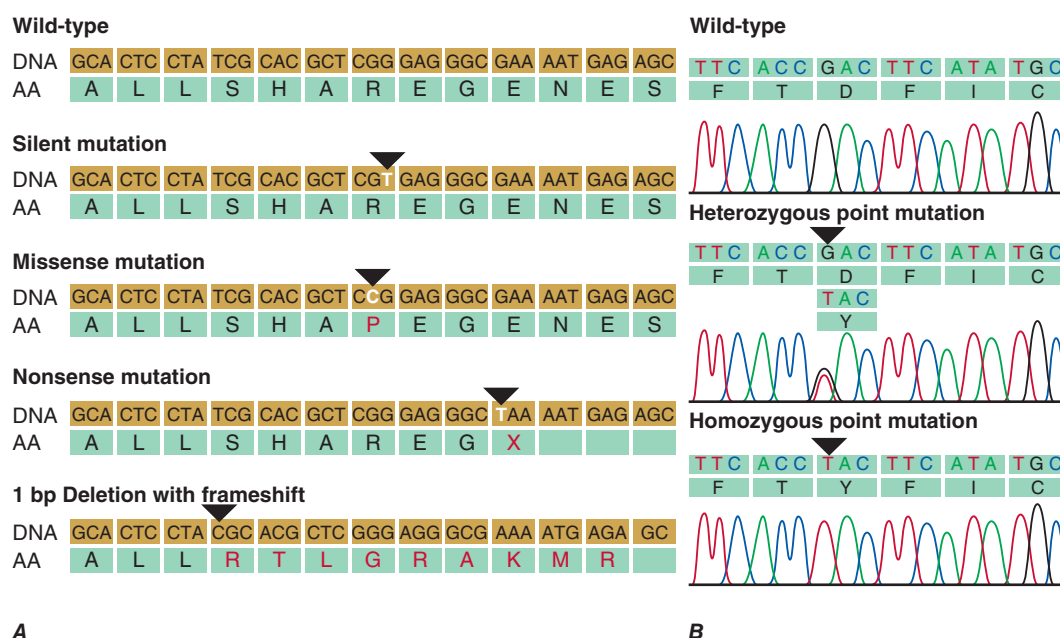


FIGURE 82-10 A. Examples of mutations. The coding strand is shown with the encoded amino acid sequence. B. Chromatograms of sequence analyses after amplification of genomic DNA by polymerase chain reaction.

UNEQUAL CROSSING-OVER Normally, DNA recombination in germ cells occurs with remarkable fidelity to maintain the precise junction sites for the exchanged DNA sequences (Fig. 82-6). However, mispairing of homologous sequences leads to unequal crossover, with gene duplication on one of the chromosomes and gene deletion on the other chromosome. A significant fraction of growth hormone (*GH*) gene deletions, for example, involve unequal crossing-over (Chap. 402). The *GH* gene is a member of a large gene cluster that includes a *GH* variant gene as well as several structurally related chorionic somatomammotropin genes and pseudogenes (highly homologous but functionally inactive relatives of a normal gene). Because such gene clusters contain multiple homologous DNA sequences arranged in tandem, they are particularly prone to undergo recombination and, consequently, gene duplication or deletion. On the other hand, duplication of the *PMP22* gene because of unequal crossing-over results in increased gene dosage and type IA Charcot-Marie-Tooth disease. Unequal crossing-over resulting in deletion of *PMP22* causes a distinct neuropathy called *hereditary liability to pressure palsy* (Chap. 459).

Glucocorticoid-remediable aldosteronism (GRA) is caused by a gene fusion or rearrangement involving the genes that encode aldosterone synthase (*CYP11B2*) and steroid 11 β -hydroxylase (*CYP11B1*), normally arranged in tandem on chromosome 8q. These two genes are 95% identical, predisposing to gene duplication and deletion by unequal crossing-over. The rearranged gene product contains the regulatory regions of 11 β -hydroxylase fused to the coding sequence of aldosterone synthetase. Consequently, the latter enzyme is expressed in the adrenocorticotrophic hormone (ACTH)-dependent zona fasciculata of the adrenal gland, resulting in overproduction of mineralocorticoids and hypertension (Chap. 406).

Gene conversion refers to a nonreciprocal exchange of homologous genetic information. It has been used to explain how an internal portion of a gene is replaced by a homologous segment copied from another allele or locus; these genetic alterations may range from a few nucleotides to a few thousand nucleotides. As a result of gene conversion, it is possible for short DNA segments of two chromosomes to be identical, even though these sequences are distinct in the parents. A practical consequence of this phenomenon is that nucleotide substitutions can occur during gene conversion between related genes, often altering the function of the gene. In disease states, gene conversion often involves intergenic exchange of DNA between a gene and a related pseudogene. For example, the 21-hydroxylase gene (*CYP21A2*) is adjacent to a non-functional pseudogene (*CYP21A1P*). Many of the nucleotide substitutions that are found in the *CYP21A2* gene in patients with congenital adrenal hyperplasia correspond to sequences that are present in the *CYP21A1P* pseudogene, suggesting gene conversion as one cause of mutagenesis. In addition, mitotic gene conversion has been suggested as a mechanism to explain revertant mosaicism in which an inherited mutation is “corrected” in certain cells. For example, patients with autosomal recessive generalized atrophic benign epidermolysis bullosa have acquired reverse mutations in one of the two mutated *COL17A1* alleles, leading to clinically unaffected patches of skin.

INSERTIONS AND DELETIONS Although many instances of insertions and deletions occur as a consequence of unequal crossing-over, there is also evidence for internal duplication, inversion, or deletion of DNA sequences. The fact that certain deletions or insertions appear to occur repeatedly as independent events indicates that specific regions within the DNA sequence predispose to these errors. For example, certain regions of the *DMD* gene, which encodes dystrophin, appear to be hot spots for deletions and result in muscular dystrophy (Chap. 462e). Some regions within the human genome are rearrangement hot spots and lead to CNVs.

ERRORS IN DNA REPAIR Because mutations caused by defects in DNA repair accumulate as somatic cells divide, these types of mutations are particularly important in the context of neoplastic disorders (Chap. 102e). Several genetic disorders involving DNA repair enzymes underscore their importance. Patients with xeroderma pigmentosum have defects in DNA damage recognition or in the nucleotide excision and repair pathway (Chap. 105). Exposed skin is dry and pigmented

and is extraordinarily sensitive to the mutagenic effects of ultraviolet irradiation. More than 10 different genes have been shown to cause the different forms of xeroderma pigmentosum. This finding is consistent with the earlier classification of this disease into different complementation groups in which normal function is rescued by the fusion of cells derived from two different forms of xeroderma pigmentosum.

Ataxia telangiectasia causes large telangiectatic lesions of the face, cerebellar ataxia, immunologic defects, and hypersensitivity to ionizing radiation (Chap. 450). The discovery of the ataxia telangiectasia mutated (*ATM*) gene reveals that it is homologous to genes involved in DNA repair and control of cell cycle checkpoints. Mutations in the *ATM* gene give rise to defects in meiosis as well as increasing susceptibility to damage from ionizing radiation. Fanconi’s anemia is also associated with an increased risk of multiple acquired genetic abnormalities. It is characterized by diverse congenital anomalies and a strong predisposition to develop aplastic anemia and acute myelogenous leukemia (Chap. 132). Cells from these patients are susceptible to chromosomal breaks caused by a defect in genetic recombination. At least 13 different complementation groups have been identified, and the loci and genes associated with Fanconi’s anemia have been cloned. HNPCC (Lynch’s syndrome) is characterized by autosomal dominant transmission of colon cancer, young age (<50 years) of presentation, predisposition to lesions in the proximal large bowel, and associated malignancies such as uterine cancer and ovarian cancer. HNPCC is predominantly caused by mutations in one of several different mismatch repair (MMR) genes including MutS homologue 2 (*MSH2*), MutL homologue 1 and 6 (*MLH1*, *MLH6*), *MSH6*, *PMS1*, and *PMS2* (Chap. 110). These proteins are involved in the detection of nucleotide mismatches and in the recognition of slipped-strand trinucleotide repeats. Germline mutations in these genes lead to microsatellite instability and a high mutation rate in colon cancer. Genetic screening tests for this disorder are now being used for families considered to be at risk (Chap. 84). Recognition of HNPCC allows early screening with colonoscopy and the implementation of prevention strategies using nonsteroidal anti-inflammatory drugs.

UNSTABLE DNA SEQUENCES *Trinucleotide repeats* may be unstable and expand beyond a critical number. Mechanistically, the expansion is thought to be caused by unequal recombination and slipped mispairing. A premutation represents a small increase in trinucleotide copy number. In subsequent generations, the expanded repeat may increase further in length and result in an increasingly severe phenotype, a process called *dynamic mutation* (see below for discussion of anticipation). Trinucleotide expansion was first recognized as a cause of the fragile X syndrome, one of the most common causes of intellectual disability. Other disorders arising from a similar mechanism include Huntington’s disease (Chap. 448), X-linked spinobulbar muscular atrophy (Chap. 452), and myotonic dystrophy (Chap. 462e). Malignant cells are also characterized by genetic instability, indicating a breakdown in mechanisms that regulate DNA repair and the cell cycle.

Functional Consequences of Mutations Functionally, mutations can be broadly classified as gain-of-function and loss-of-function mutations. Gain-of-function mutations are typically dominant (e.g., they result in phenotypic alterations when a single allele is affected). Inactivating mutations are usually recessive, and an affected individual is homozygous or compound heterozygous (e.g., carrying two different mutant alleles of the same gene) for the disease-causing mutations. Alternatively, mutation in a single allele can result in *haploinsufficiency*, a situation in which one normal allele is not sufficient to maintain a normal phenotype. Haploinsufficiency is a commonly observed mechanism in diseases associated with mutations in transcription factors (Table 82-2). Remarkably, the clinical features among patients with an identical mutation in a transcription factor often vary significantly. One mechanism underlying this variability consists in the influence of modifying genes. Haploinsufficiency can also affect the expression of rate-limiting enzymes. For example, haploinsufficiency in enzymes involved in heme synthesis can cause porphyrias (Chap. 430).

An increase in dosage of a gene product may also result in disease, as illustrated by the duplication of the *DAX1* gene in dosage-sensitive sex reversal (Chap. 410). Mutation in a single allele can also result in loss of function due to a dominant-negative effect. In this case, the mutated allele interferes with the function of the normal gene product by one of several different mechanisms: (1) a mutant protein may interfere with the function of a multimeric protein complex, as illustrated by mutations in type 1 collagen (*COL1A1*, *COL1A2*) genes in osteogenesis imperfecta (Chap. 427); (2) a mutant protein may occupy binding sites on proteins or promoter response elements, as illustrated by thyroid hormone resistance, a disorder in which inactivated thyroid hormone receptor β binds to target genes and functions as an antagonist of normal receptors (Chap. 405); or (3) a mutant protein can be cytotoxic as in α_1 antitrypsin deficiency (Chap. 314) or autosomal dominant neurohypophyseal diabetes insipidus (Chap. 404), in which the abnormally folded proteins are trapped within the endoplasmic reticulum and ultimately cause cellular damage.

Genotype and Phenotype • ALLELES, GENOTYPES, AND HAPLOTYPES An observed trait is referred to as a *phenotype*; the genetic information defining the phenotype is called the *genotype*. Alternative forms of a gene or a genetic marker are referred to as *alleles*. Alleles may be polymorphic variants of nucleic acids that have no apparent effect on gene expression or function. In other instances, these variants may have subtle effects on gene expression, thereby conferring adaptive advantages associated with genetic diversity. On the other hand, allelic variants may reflect mutations that clearly alter the function of a gene product. The common Glu6Val (E6V) sickle cell mutation in the β -globin gene and the Δ F508 deletion of phenylalanine (F) in the *CFTR* gene are examples of allelic variants of these genes that result in disease. Because each individual has two copies of each chromosome (one inherited from the mother and one inherited from the father), he or she can have only two alleles at a given locus. However, there can be many different alleles in the population. The normal or common allele is usually referred to as *wild type*. When alleles at a given locus are identical, the individual is *homozygous*. Inheriting identical copies of a mutant allele occurs in many autosomal recessive disorders, particularly in circumstances of consanguinity or isolated populations. If the alleles are different on the maternal and the paternal copy of the gene, the individual is *heterozygous* at this locus (Fig. 82-10). If two different mutant alleles are inherited at a given locus, the individual is said to be a *compound heterozygote*. *Hemizygous* is used to describe males with a mutation in an X chromosomal gene or a female with a loss of one X chromosomal locus.

Genotypes describe the specific alleles at a particular locus. For example, there are three common alleles (E2, E3, E4) of the apolipoprotein E (*APOE*) gene. The genotype of an individual can therefore be described as *APOE3/4* or *APOE4/4* or any other variant. These designations indicate which alleles are present on the two chromosomes in the *APOE* gene at locus 19q13.2. In other cases, the genotype might be assigned arbitrary numbers (e.g., 1/2) or letters (e.g., B/b) to distinguish different alleles.

A *haplotype* refers to a group of alleles that are closely linked together at a genomic locus (Fig. 82-4). Haplotypes are useful for tracking the transmission of genomic segments within families and for detecting evidence of genetic recombination, if the crossover event occurs between the alleles (Fig. 82-6). As an example, various alleles at the histocompatibility locus antigen (HLA) on chromosome 6p are used to establish haplotypes associated with certain disease states. For example, 21-hydroxylase deficiency, complement deficiency, and hemochromatosis are each associated with specific HLA haplotypes. It is now recognized that these genes lie in close proximity to the HLA locus, which explains why HLA associations were identified even before the disease genes were cloned and localized. In other cases, specific HLA associations with diseases such as ankylosing spondylitis (HLA-B27) or type 1 diabetes mellitus (HLA-DR4) reflect the role of specific HLA allelic variants in susceptibility to these autoimmune diseases. The characterization of common SNP haplotypes in numerous populations from different parts of the world through the HapMap Project is providing a novel tool for association studies designed

to detect genes involved in the pathogenesis of complex disorders (Table 82-1). The presence or absence of certain haplotypes may also become relevant for the customized choice of medical therapies (pharmacogenomics) or for preventive strategies.

Genotype-phenotype correlation describes the association of a specific mutation and the resulting phenotype. The phenotype may differ depending on the location or type of the mutation in some genes. For example, in von Hippel-Lindau disease, an autosomal dominant multisystem disease that can include renal cell carcinoma, hemangioblastomas, and pheochromocytomas, among others, the phenotype varies greatly and the identification of the specific mutation can be clinically useful in order to predict the phenotypic spectrum.

ALLELIC HETEROGENEITY *Allelic heterogeneity* refers to the fact that different mutations in the same genetic locus can cause an identical or similar phenotype. For example, many different mutations of the β -globin locus can cause β thalassemia (Table 82-3) (Fig. 82-9). In essence, allelic heterogeneity reflects the fact that many different mutations are capable of altering protein structure and function. For this reason, maps of inactivating mutations in genes usually show a near-random distribution. Exceptions include (1) a founder effect, in which a particular mutation that does not affect reproductive capacity can be traced to a single individual; (2) “hot spots” for mutations, in which the nature of the DNA sequence predisposes to a recurring mutation; and (3) localization of mutations to certain domains that are particularly critical for protein function. Allelic heterogeneity creates a practical problem for genetic testing because one must often examine the entire genetic locus for mutations, because these can differ in each patient. For example, there are currently 1963 reported mutations in the *CFTR* gene (Fig. 82-3). Mutational analysis may initially focus on a panel of mutations that are particularly frequent (often taking the ethnic background of the patient into account), but a negative result does not exclude the presence of a mutation elsewhere in the gene. One should also be aware that mutational analyses generally focus on the coding region of a gene without considering regulatory and intronic regions. Because disease-causing mutations may be located outside the coding regions, negative results need to be interpreted with caution. The advent of more comprehensive sequencing technologies greatly facilitates concomitant mutational analyses of several genes after targeted enrichment, or even mutational analysis of the whole exome or genome. However, comprehensive sequencing can result in significant diagnostic challenges because the detection of a sequence alteration alone is not always sufficient to establish that it has a causal role.

PHENOTYPIC HETEROGENEITY *Phenotypic heterogeneity* occurs when more than one phenotype is caused by allelic mutations (e.g., different mutations in the same gene) (Table 82-3). For example, laminopathies are monogenic multisystem disorders that result from mutations in the *LMNA* gene, which encodes the nuclear lamins A and C. Twelve autosomal dominant and four autosomal recessive disorders are caused by mutations in the *LMNA* gene. They include several forms of lipodystrophies, Emery-Dreifuss muscular dystrophy, progeria syndromes, a form of neuronal Charcot-Marie-Tooth disease (type 2B1), and a group of overlapping syndromes. Remarkably, hierarchical cluster analysis has revealed that the phenotypes vary depending on the position of the mutation (*genotype-phenotype correlation*). Similarly, identical mutations in the *FGFR2* gene can result in very distinct phenotypes: Crouzon’s syndrome (craniofacial synostosis) or Pfeiffer’s syndrome (acrocephalopolysyndactyly).

LOCUS OR NONALLELIC HETEROGENEITY AND PHENOCOPIES *Nonallelic or locus heterogeneity* refers to the situation in which a similar disease phenotype results from mutations at different genetic loci (Table 82-3). This often occurs when more than one gene product produces different subunits of an interacting complex or when different genes are involved in the same genetic cascade or physiologic pathway. For example, osteogenesis imperfecta can arise from mutations in two different procollagen genes (*COL1A1* or *COL1A2*) that are located on different chromosomes, and at least eight other genes (Chap. 427). The effects of inactivating mutations in these two genes are similar because the protein products comprise different subunits

TABLE 82-3 SELECTED EXAMPLES OF LOCUS HETEROGENEITY AND PHENOTYPIC HETEROGENEITY

Phenotypic Heterogeneity			
Gene, Protein	Phenotype	Inheritance	OMIM
LMNA, Lamin A/C	Emery-Dreifuss muscular dystrophy (AD)	AD	181350
	Familial partial lipodystrophy Dunnigan	AD	151660
	Hutchinson-Gilford progeria	AD	176670
	Atypical Werner's syndrome	AD	150330
	Dilated cardiomyopathy	AD	115200
	Early-onset atrial fibrillation	AD	607554
	Emery-Dreifuss muscular dystrophy (AR)	AR	604929
	Limb-girdle muscular dystrophy type 1B	AR	159001
	Charcot-Marie-Tooth type 2B1	AR	605588
KRAS	Noonan's syndrome	AD	163950
	Cardio-facio-cutaneous syndrome	AD	115150
Locus Heterogeneity			
Phenotype	Gene	Chromosomal Location	Protein
Familial hypertrophic cardiomyopathy Genes encoding sarcomeric proteins	MYH7	14q12	Myosin heavy chain beta
	TNNT2	1q2	Troponin-T2
	TPM1	15q22.1	Tropomyosin alpha
			Myosin-binding protein C
	MYBPC3	11p11q	protein C
	TNNI3	19q13.4	Troponin 1
	MYL2	12q23-24.3	Myosin light chain 2
	MYL3	3p	Myosin light chain 3
	TTN	2q24.3	Cardiac titin
	ACTC	15q11	Cardiac alpha actin
	MYH6	14q1	Myosin heavy chain alpha
	MYLK2	20q13.3	Myosin light-peptide kinase
	CAV3	3p25	Caveolin 3
Genes encoding nonsarcomeric proteins	MTT1	Mitochondrial	tRNA isoleucine
	MTTG	Mitochondrial	tRNA glycine
	PRKAG2	7q35-q36	AMP-activated protein kinase γ 2 subunit
	DMPK	19q13.2-13.3	Myotonic protein kinase (myotonic dystrophy)
	FRDA	9q13	Frataxin (Friedreich's ataxia)
Polycystic kidney disease	PKD1	16p13.3-13.12	Polycystin 1 (AD)
	PKD2	4q21.-23	Polycystin 2 (AD)
	PKHD1	6p21.1-p12	Fibrocystin (AR)
Noonan's syndrome	PTPN11	12q24.1	Protein-tyrosine phosphatase 2c
	KRAS	12p12.1	KRAS

of the helical collagen fiber. Similarly, muscular dystrophy syndromes can be caused by mutations in various genes, consistent with the fact that it can be transmitted in an X-linked (Duchenne or Becker), autosomal dominant (limb-girdle muscular dystrophy type 1), or autosomal recessive (limb-girdle muscular dystrophy type 2) manner (Chap. 462e). Mutations in the X-linked *DMD* gene, which encodes dystrophin, are the most common cause of muscular dystrophy. This feature reflects the large size of the gene as well as the fact that the phenotype is expressed in hemizygous males because they have only a single copy of the X chromosome. Dystrophin is associated with a large protein complex linked to the membrane-associated cytoskeleton in muscle. Mutations in several different components of this protein complex can also cause muscular dystrophy syndromes. Although the phenotypic features of some of these disorders are distinct, the phenotypic spectrum caused by mutations in different genes overlaps, thereby leading to nonallelic heterogeneity. It should be noted that mutations in dystrophin also cause allelic heterogeneity. For example, mutations in the *DMD* gene can cause either Duchenne's or the less severe Becker's muscular dystrophy, depending on the severity of the protein defect.

Recognition of nonallelic heterogeneity is important for several reasons: (1) the ability to identify disease loci in linkage studies is reduced

by including patients with similar phenotypes but different genetic disorders; (2) genetic testing is more complex because several different genes need to be considered along with the possibility of different mutations in each of the candidate genes; and (3) novel information is gained about how genes or proteins interact, providing unique insights into molecular physiology.

Phenocopies refer to circumstances in which nongenetic conditions mimic a genetic disorder. For example, features of toxin- or drug-induced neurologic syndromes can resemble those seen in Huntington's disease, and vascular causes of dementia share phenotypic features with familial forms of Alzheimer's dementia (Chap. 448). As in nonallelic heterogeneity, the presence of phenocopies has the potential to confound linkage studies and genetic testing. Patient history and subtle differences in phenotype can often provide clues that distinguish these disorders from related genetic conditions.

VARIABLE EXPRESSIVITY AND INCOMPLETE PENETRANCE The same genetic mutation may be associated with a phenotypic spectrum in different affected individuals, thereby illustrating the phenomenon of *variable expressivity*. This may include different manifestations of a disorder variably involving different organs

(e.g., multiple endocrine neoplasia [MEN]), the severity of the disorder (e.g., cystic fibrosis), or the age of disease onset (e.g., Alzheimer's dementia). MEN 1 illustrates several of these features. In this autosomal dominant tumor syndrome, affected individuals carry an inactivating germline mutation that is inherited in an autosomal dominant fashion. After somatic inactivation of the alternate allele, they can develop tumors of the parathyroid gland, endocrine pancreas, and the pituitary gland (Chap. 408). However, the pattern of tumors in the different glands, the age at which tumors develop, and the types of hormones produced vary among affected individuals, even within a given family. In this example, the phenotypic variability arises, in part, because of the requirement for a second somatic mutation in the normal copy of the *MEN1* gene, as well as the large array of different cell types that are susceptible to the effects of *MEN1* gene mutations. In part, variable expression reflects the influence of modifier genes, or genetic background, on the effects of a particular mutation. Even in identical twins, in whom the genetic constitution is essentially the same, one can occasionally see variable expression of a genetic disease.

Interactions with the environment can also influence the course of a disease. For example, the manifestations and severity of hemochromatosis can be influenced by iron intake (Chap. 428), and the course of phenylketonuria is affected by exposure to phenylalanine in the diet (Chap. 434e). Other metabolic disorders, such as hyperlipidemias and porphyria, also fall into this category. Many mechanisms, including genetic effects and environmental influences, can therefore lead to variable expressivity. In genetic counseling, it is particularly important to recognize this variability, because one cannot always predict the course of disease, even when the mutation is known.

Penetrance refers to the proportion of individuals with a mutant genotype that express the phenotype. If all carriers of a mutant express the phenotype, penetrance is complete, whereas it is said to be *incomplete* or *reduced* if some individuals do not exhibit features of the phenotype. Dominant conditions with incomplete penetrance are characterized by skipping of generations with unaffected carriers transmitting the mutant gene. For example, hypertrophic obstructive cardiomyopathy (HCM) caused by mutations in the *myosin-binding protein C* gene is a dominant disorder with clinical features in only a subset of patients who carry the mutation (Chap. 283). Patients who have the mutation but no evidence of the disease can still transmit the disorder to subsequent generations. In many conditions with postnatal onset, the proportion of gene carriers who are affected varies with age. Thus, when describing penetrance, one has to specify age. For example, for disorders such as Huntington's disease or familial amyotrophic lateral sclerosis, which present later in life, the rate of penetrance is influenced by the age at which the clinical assessment is performed. **Imprinting** can also modify the penetrance of a disease. For example, in patients with Albright's hereditary osteodystrophy, mutations in the $G\alpha$ subunit (*GNAS1* gene) are expressed clinically only in individuals who inherit the mutation from their mother (Chap. 424).

SEX-INFLUENCED PHENOTYPES Certain mutations affect males and females quite differently. In some instances, this is because the gene resides on the X or Y sex chromosomes (X-linked disorders and Y-linked disorders). As a result, the phenotype of mutated X-linked genes will be expressed fully in males but variably in heterozygous females, depending on the degree of X-inactivation and the function of the gene. For example, most heterozygous female carriers of factor VIII deficiency (hemophilia A) are asymptomatic because sufficient factor VIII is produced to prevent a defect in coagulation (Chap. 141). On the other hand, some females heterozygous for the X-linked lipid storage defect caused by α -galactosidase A deficiency (Fabry's disease) experience mild manifestations of painful neuropathy, as well as other features of the disease (Chap. 432e). Because only males have a Y chromosome, mutations in genes such as *SRY*, which causes male-to-female sex reversal, or *DAZ* (deleted in azoospermia), which causes abnormalities of spermatogenesis, are unique to males (Chap. 410).

Other diseases are expressed in a sex-limited manner because of the differential function of the gene product in males and females. Activating mutations in the luteinizing hormone receptor cause

dominant male-limited precocious puberty in boys (Chap. 411). The phenotype is unique to males because activation of the receptor induces testosterone production in the testis, whereas it is functionally silent in the immature ovary. Biallelic inactivating mutations of the follicle-stimulating hormone (FSH) receptor cause primary ovarian failure in females because the follicles do not develop in the absence of FSH action. In contrast, affected males have a more subtle phenotype, because testosterone production is preserved (allowing sexual maturation) and spermatogenesis is only partially impaired (Chap. 411). In congenital adrenal hyperplasia, most commonly caused by 21-hydroxylase deficiency, cortisol production is impaired and ACTH stimulation of the adrenal gland leads to increased production of androgenic precursors (Chap. 406). In females, the increased androgen level causes ambiguous genitalia, which can be recognized at the time of birth. In males, the diagnosis may be made on the basis of adrenal insufficiency at birth, because the increased adrenal androgen level does not alter sexual differentiation, or later in childhood, because of the development of precocious puberty. Hemochromatosis is more common in males than in females, presumably because of differences in dietary iron intake and losses associated with menstruation and pregnancy in females (Chap. 428).

Chromosomal Disorders Chromosomal or cytogenetic disorders are caused by numerical or structural aberrations in chromosomes. For a detailed discussion of disorders of chromosome number and structure, see Chap. 83e. Deviations in chromosome number are common causes of abortions, developmental disorders, and malformations. **Contiguous gene syndromes** (e.g., large deletions affecting several genes) have been useful for identifying the location of new disease-causing genes. Because of the variable size of gene deletions in different patients, a systematic comparison of phenotypes and locations of deletion breakpoints allows positions of particular genes to be mapped within the critical genomic region.

Monogenic Mendelian Disorders Monogenic human diseases are frequently referred to as *Mendelian disorders* because they obey the principles of genetic transmission originally set forth in Gregor Mendel's classic work. The continuously updated OMIM catalogue lists several thousand of these disorders and provides information about the clinical phenotype, molecular basis, allelic variants, and pertinent animal models (Table 82-1). The mode of inheritance for a given phenotypic trait or disease is determined by pedigree analysis. All affected and unaffected individuals in the family are recorded in a pedigree using standard symbols (Fig. 82-11). The principles of allelic segregation, and the transmission of alleles from parents to children, are illustrated in Fig. 82-12. One dominant (A) allele and one recessive (a) allele can display three Mendelian modes of inheritance: autosomal dominant, autosomal recessive, and X-linked. About 65% of human monogenic disorders are autosomal dominant, 25% are autosomal recessive, and 5% are X-linked. Genetic testing is now available for many of these disorders and plays an increasingly important role in clinical medicine (Chap. 84).

AUTOSOMAL DOMINANT DISORDERS These disorders assume particular relevance because mutations in a single allele are sufficient to cause the disease. In contrast to recessive disorders, in which disease pathogenesis is relatively straightforward because there is loss of gene function, dominant disorders can be caused by various disease mechanisms, many of which are unique to the function of the genetic pathway involved.

In autosomal dominant disorders, individuals are affected in successive generations; the disease does not occur in the offspring of unaffected individuals. Males and females are affected with equal frequency because the defective gene resides on one of the 22 autosomes (Fig. 82-13A). Autosomal dominant mutations alter one of the two alleles at a given locus. Because the alleles segregate randomly at meiosis, the probability that an offspring will be affected is 50%. Unless there is a new germline mutation, an affected individual has an affected parent. Children with a normal genotype do not transmit the disorder. Due to differences in penetrance or expressivity (see above), the

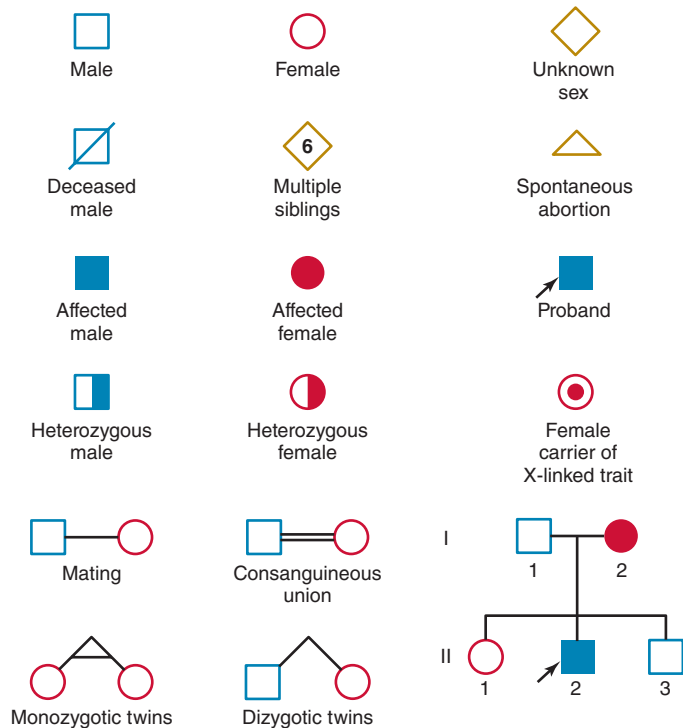


FIGURE 82-11 Standard pedigree symbols.

clinical manifestations of autosomal dominant disorders may be variable. Because of these variations, it is sometimes challenging to determine the pattern of inheritance.

It should be recognized, however, that some individuals acquire a mutated gene from an unaffected parent. *De novo* germline mutations occur more frequently during later cell divisions in gametogenesis, which explains why siblings are rarely affected. As noted before, new germline mutations occur more frequently in fathers of advanced age. For example, the average age of fathers with new germline mutations that cause Marfan's syndrome is ~37 years, whereas fathers who transmit the disease by inheritance have an average age of ~30 years.

AUTOSOMAL RECESSIVE DISORDERS In recessive disorders, the mutated alleles result in a complete or partial loss of function. They frequently involve enzymes in metabolic pathways, receptors, or proteins in signaling cascades. In an autosomal recessive disease, the affected individual, who can be of either sex, is a homozygote or compound heterozygote for a single-gene defect. With a few important exceptions, autosomal recessive diseases are rare and often occur in the context of parental consanguinity. The relatively high frequency of certain recessive disorders such as sickle cell anemia, cystic fibrosis, and thalassemia, is partially explained by a selective biologic advantage for the heterozygous state (see below). Although heterozygous carriers of a defective allele are usually clinically normal, they may display subtle differences in phenotype that only become apparent with more precise testing or in the context of certain environmental influences. In sickle

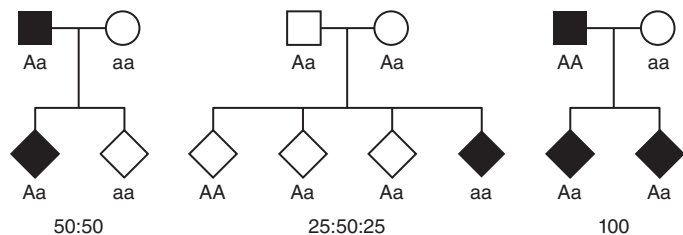


FIGURE 82-12 Segregation of alleles. Segregation of genotypes in the offspring of parents with one dominant (A) and one recessive (a) allele. The distribution of the parental alleles to their offspring depends on the combination present in the parents. Filled symbols = affected individuals.

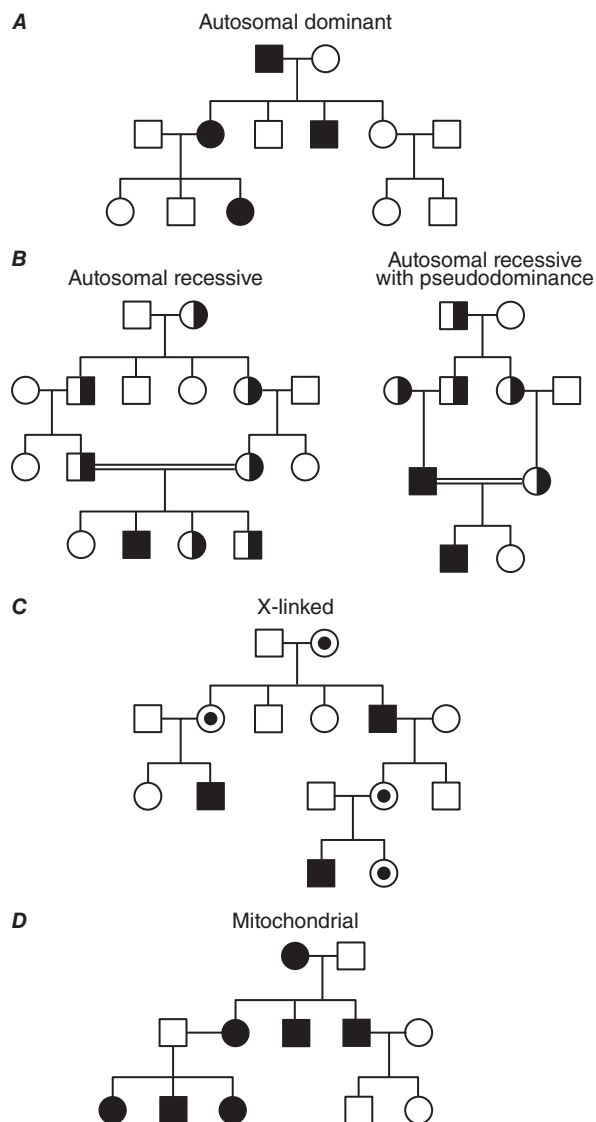


FIGURE 82-13 (A) Dominant, (B) recessive, (C) X-linked, and (D) mitochondrial (matrilinear) inheritance.

cell anemia, for example, heterozygotes are normally asymptomatic. However, in situations of dehydration or diminished oxygen pressure, sickle cell crises can also occur in heterozygotes (**Chap. 127**).

In most instances, an affected individual is the offspring of heterozygous parents. In this situation, there is a 25% chance that the offspring will have a normal genotype, a 50% probability of a heterozygous state, and a 25% risk of homozygosity for the recessive alleles (**Figs. 82-10, 82-13B**). In the case of one unaffected heterozygous and one affected homozygous parent, the probability of disease increases to 50% for each child. In this instance, the pedigree analysis mimics an autosomal dominant mode of inheritance (*pseudodominance*). In contrast to autosomal dominant disorders, new mutations in recessive alleles are rarely manifest because they usually result in an asymptomatic carrier state.

X-LINKED DISORDERS Males have only one X chromosome; consequently, a daughter always inherits her father's X chromosome in addition to one of her mother's two X chromosomes. A son inherits the Y chromosome from his father and one maternal X chromosome. Thus, the characteristic features of X-linked inheritance are (1) the absence of father-to-son transmission, and (2) the fact that all daughters of an affected male are obligate carriers of the mutant allele (**Fig. 82-13C**). The risk of developing disease due to a mutant X-chromosomal gene differs in the two sexes. Because males have only one X chromosome, they are hemizygous for the mutant allele; thus, they are more likely

to develop the mutant phenotype, regardless of whether the mutation is dominant or recessive. A female may be either heterozygous or homozygous for the mutant allele, which may be dominant or recessive. The terms *X-linked dominant* or *X-linked recessive* are therefore only applicable to expression of the mutant phenotype in women. In addition, the expression of X-chromosomal genes is influenced by X chromosome inactivation.

Y-LINKED DISORDERS The Y chromosome has a relatively small number of genes. One such gene, the sex-region determining Y factor (*SRY*), which encodes the testis-determining factor (*TDF*), is crucial for normal male development. Normally there is infrequent exchange of sequences on the Y chromosome with the X chromosome. The *SRY* region is adjacent to the pseudoautosomal region, a chromosomal segment on the X and Y chromosomes with a high degree of homology. A crossing-over event occasionally involves the *SRY* region with the distal tip of the X chromosome during meiosis in the male. Translocations can result in XY females with the Y chromosome lacking the *SRY* gene or XX males harboring the *SRY* gene on one of the X chromosomes (Chap. 410). Point mutations in the *SRY* gene may also result in individuals with an XY genotype and an incomplete female phenotype. Most of these mutations occur *de novo*. Men with oligospermia/azoospermia frequently have microdeletions on the long arm of the Y chromosome that involve one or more of the azoospermia factor (*AZF*) genes.

Exceptions to Simple Mendelian Inheritance Patterns • MITOCHONDRIAL DISORDERS Mendelian inheritance refers to the transmission of genes encoded by DNA contained in the nuclear chromosomes. In addition, each mitochondrion contains several copies of a small circular chromosome (Chap. 85e). The mitochondrial DNA (mtDNA) is ~16.5 kb and encodes transfer and ribosomal RNAs and 13 core proteins that are components of the respiratory chain involved in oxidative phosphorylation and ATP generation. The mitochondrial genome does not recombine and is inherited through the maternal line because sperm does not contribute significant cytoplasmic components to the zygote. A noncoding region of the mitochondrial chromosome, referred to as D-loop, is highly polymorphic. This property, together with the absence of mtDNA recombination, makes it a valuable tool for studies tracing human migration and evolution, and it is also used for specific forensic applications.

Inherited mitochondrial disorders are transmitted in a matrilineal fashion; all children from an affected mother will inherit the disease, but it will not be transmitted from an affected father to his children (Fig. 82-13D). Alterations in the mtDNA that involves enzymes required for oxidative phosphorylation lead to reduction of ATP supply, generation of free radicals, and induction of apoptosis. Several syndromic disorders arising from mutations in the mitochondrial genome are known in humans and they affect both protein-coding and tRNA genes (Chap. 85e). The broad clinical spectrum often involves (cardio) myopathies and encephalopathies because of the high dependence of these tissues on oxidative phosphorylation. The age of onset and the clinical course are highly variable because of the unusual mechanisms of mtDNA transmission, which replicates independently from nuclear DNA. During cell replication, the proportion of wild-type and mutant mitochondria can drift among different cells and tissues. The resulting heterogeneity in the proportion of mitochondria with and without a mutation is referred to as *heteroplasmy* and underlies the phenotypic variability that is characteristic of mitochondrial diseases.

Acquired somatic mutations in mitochondria are thought to be involved in several age-dependent degenerative disorders affecting predominantly muscle and the peripheral and central nervous system (e.g., Alzheimer's and Parkinson's diseases). Establishing that an mtDNA alteration is causal for a clinical phenotype is challenging because of the high degree of polymorphism in mtDNA and the phenotypic variability characteristic of these disorders. Certain pharmacologic treatments may have an impact on mitochondria and/or their function. For example, treatment with the antiretroviral compound azidothymidine (AZT) causes an acquired mitochondrial myopathy through depletion of muscular mtDNA.

MOSAICISM Mosaicism refers to the presence of two or more genetically distinct cell lines in the tissues of an individual. It results from a mutation that occurs during embryonic, fetal, or extrauterine development. The developmental stage at which the mutation arises will determine whether germ cells and/or somatic cells are involved. Chromosomal mosaicism results from nondisjunction at an early embryonic mitotic division, leading to the persistence of more than one cell line, as exemplified by some patients with Turner's syndrome (Chap. 410). Somatic mosaicism is characterized by a patchy distribution of genetically altered somatic cells. The McCune-Albright syndrome, for example, is caused by activating mutations in the stimulatory G protein α (*Gsa*) that occur early in development (Chap. 424). The clinical phenotype varies depending on the tissue distribution of the mutation; manifestations include ovarian cysts that secrete sex steroids and cause precocious puberty, polyostotic fibrous dysplasia, café-au-lait skin pigmentation, growth hormone-secreting pituitary adenomas, and hypersecreting autonomous thyroid nodules (Chap. 412).

X-INACTIVATION, IMPRINTING, AND UNIPARENTAL DISOMY According to traditional Mendelian principles, the parental origin of a mutant gene is irrelevant for the expression of the phenotype. There are, however, important exceptions to this rule. *X-inactivation* prevents the expression of most genes on one of the two X chromosomes in every cell of a female. Gene inactivation through genomic imprinting occurs on selected chromosomal regions of autosomes and leads to inheritable preferential expression of one of the parental alleles. It is of pathophysiologic importance in disorders where the transmission of disease is dependent on the sex of the transmitting parent and, thus, plays an important role in the expression of certain genetic disorders. Two classic examples are the Prader-Willi syndrome and Angelman's syndrome (Chap. 83e). Prader-Willi syndrome is characterized by diminished fetal activity, obesity, hypotonia, mental retardation, short stature, and hypogonadotropic hypogonadism. Deletions of the paternal copy of the Prader-Willi locus located on the short arm of chromosome 15 result in a contiguous gene syndrome involving missing paternal copies of the *necdin* and *SNRPN* genes, among others. In contrast, patients with Angelman's syndrome, characterized by mental retardation, seizures, ataxia, and hypotonia, have deletions involving the maternal copy of this region on chromosome 15. These two syndromes may also result from *uniparental disomy*. In this case, the syndromes are not caused by deletions on chromosome 15 but by the inheritance of either two maternal chromosomes (Prader-Willi syndrome) or two paternal chromosomes (Angelman's syndrome). Lastly, the two distinct phenotypes can also be caused by an imprinting defect that impairs the resetting of the imprint during zygote development (defect in the father leads to Prader-Willi syndrome; defect in the mother leads to Angelman's syndrome).

Imprinting and the related phenomenon of allelic exclusion may be more common than currently documented, because it is difficult to examine levels of mRNA expression from the maternal and paternal alleles in specific tissues or in individual cells. Genomic imprinting, or uniparental disomy, is involved in the pathogenesis of several other disorders and malignancies (Chap. 83e). For example, hydatidiform moles contain a normal number of diploid chromosomes, but they are all of paternal origin. The opposite situation occurs in ovarian teratomata, with 46 chromosomes of maternal origin. Expression of the imprinted gene for insulin-like growth factor II (*IGF-II*) is involved in the pathogenesis of the cancer-predisposing Beckwith-Wiedemann syndrome (BWS) (Chap. 101e). These children show somatic overgrowth with organomegalies and hemihypertrophy, and they have an increased risk of embryonal malignancies such as Wilms' tumor. Normally, only the paternally derived copy of the *IGF-II* gene is active and the maternal copy is inactive. Imprinting of the *IGF-II* gene is regulated by *H19*, which encodes an RNA transcript that is not translated into protein. Disruption or lack of *H19* methylation leads to a relaxation of *IGF-II* imprinting and expression of both alleles. Alterations of the epigenome through gain and loss of DNA methylation, as well as altered histone modifications, play an important role in the pathogenesis of malignancies.

SOMATIC MUTATIONS Cancer can be considered a genetic disease at the cellular level (**Chap. 101e**). Cancers are monoclonal in origin, indicating that they have arisen from a single precursor cell with one or several mutations in genes controlling growth (proliferation or apoptosis) and/or differentiation. These acquired somatic mutations are restricted to the tumor and its metastases and are not found in the surrounding normal tissue. The molecular alterations include dominant gain-of-function mutations in oncogenes, recessive loss-of-function mutations in tumor-suppressor genes and DNA repair genes, gene amplification, and chromosome rearrangements. Rarely, a single mutation in certain genes may be sufficient to transform a normal cell into a malignant cell. In most cancers, however, the development of a malignant phenotype requires several genetic alterations for the gradual progression from a normal cell to a cancerous cell, a phenomenon termed *multistep carcinogenesis* (**Chaps. 101e and 102e**). Genome-wide analyses of cancers using deep sequencing often reveal somatic rearrangements resulting in fusion genes and mutations in multiple genes. Comprehensive sequence analyses provide further insight into genetic heterogeneity within malignancies; these include intratumoral heterogeneity among the cells of the primary tumor, intermetastatic and intrametastatic heterogeneity, and interpatient differences. These analyses further support the notion of cancer as an ongoing process of clonal evolution, in which successive rounds of clonal selection within the primary tumor and metastatic lesions result in diverse genetic and epigenetic alterations that require targeted (personalized) therapies. The heterogeneity of mutations within a tumor can also lead to resistance to target therapies because cells with mutations that are resistant to the therapy, even if they are a minor part of the tumor population, will be selected as the more sensitive cells are killed. Most human tumors express telomerase, an enzyme formed of a protein and an RNA component, which adds telomere repeats at the ends of chromosomes during replication. This mechanism impedes shortening of the telomeres, which is associated with senescence in normal cells and is associated with enhanced replicative capacity in cancer cells. Telomerase inhibitors provide a novel strategy for treating advanced human cancers.

In many cancer syndromes, there is an inherited *predisposition* to tumor formation. In these instances, a germline mutation is inherited in an autosomal dominant fashion inactivating one allele of an autosomal tumor-suppressor gene. If the second allele is inactivated by a somatic mutation or by epigenetic silencing in a given cell, this will lead to neoplastic growth (Knudson two-hit model). Thus, the defective allele in the germline is transmitted in a dominant mode, although tumorigenesis results from a biallelic loss of the tumor-suppressor gene

in an affected tissue. The classic example to illustrate this phenomenon is retinoblastoma, which can occur as a sporadic or hereditary tumor. In sporadic retinoblastoma, both copies of the retinoblastoma (*RB*) gene are inactivated through two somatic events. In hereditary retinoblastoma, one mutated or deleted *RB* allele is inherited in an autosomal dominant manner and the second allele is inactivated by a subsequent somatic mutation. This two-hit model applies to other inherited cancer syndromes such as MEN 1 (**Chap. 408**) and neurofibromatosis type 2 (**Chap. 118**).

NUCLEOTIDE REPEAT EXPANSION DISORDERS Several diseases are associated with an increase in the number of nucleotide repeats above a certain threshold (**Table 82-4**). The repeats are sometimes located within the coding region of the genes, as in Huntington's disease or the X-linked form of spinal and bulbar muscular atrophy (SBMA; Kennedy's syndrome). In other instances, the repeats probably alter gene regulatory sequences. If an expansion is present, the DNA fragment is unstable and tends to expand further during cell division. The length of the nucleotide repeat often correlates with the severity of the disease. When repeat length increases from one generation to the next, disease manifestations may worsen or be observed at an earlier age; this phenomenon is referred to as *anticipation*. In Huntington's disease, for example, there is a correlation between age of onset and length of the triplet codon expansion (**Chap. 444e**). Anticipation has also been documented in other diseases caused by dynamic mutations in trinucleotide repeats (**Table 82-4**). The repeat number may also vary in a tissue-specific manner. In myotonic dystrophy, the CTG repeat may be tenfold greater in muscle tissue than in lymphocytes (**Chap. 462e**).

Complex Genetic Disorders The expression of many common diseases such as cardiovascular disease, hypertension, diabetes, asthma, psychiatric disorders, and certain cancers is determined by a combination of genetic background, environmental factors, and lifestyle. A trait is called *polygenic* if multiple genes contribute to the phenotype or *multifactorial* if multiple genes are assumed to interact with environmental factors. Genetic models for these complex traits need to account for genetic heterogeneity and interactions with other genes and the environment. Complex genetic traits may be influenced by modifier genes that are not linked to the main gene involved in the pathogenesis of the trait. This type of gene-gene interaction, or *epistasis*, plays an important role in polygenic traits that require the simultaneous presence of variations in multiple genes to result in a pathologic phenotype.

Type 2 diabetes mellitus provides a paradigm for considering a multifactorial disorder, because genetic, nutritional, and lifestyle factors are intimately interrelated in disease pathogenesis (**Table 82-5**) (**Chap. 417**).

TABLE 82-4 SELECTED TRINUCLEOTIDE REPEAT DISORDERS

Disease	Locus	Repeat	Triplet Length (Normal/Disease)	Inheritance	Gene Product
X-chromosomal spinobulbar muscular atrophy (SBMA)	Xq11-q12	CAG	11–34/40–62	XR	Androgen receptor
Fragile X syndrome (FRAXA)	Xq27.3	CGG	6–50/200–300	XR	FMR-1 protein
Fragile X syndrome (FRAXE)	Xq28	GCC	6–25/>200	XR	FMR-2 protein
Dystrophia myotonica (DM)	19q13.2-q13.3	CTG	5–30/200–1000	AD, variable penetrance	Myotonin protein kinase
Huntington's disease (HD)	4p16.3	CAG	6–34/37–180	AD	Huntingtin
Spinocerebellar ataxia type 1 (SCA1)	6p21.3-21.2	CAG	6–39/40–88	AD	Ataxin 1
Spinocerebellar ataxia type 2 (SCA2)	12q24.1	CAG	15–31/34–400	AD	Ataxin 2
Spinocerebellar ataxia type 3 (SCA3); Machado-Joseph disease (MD)	14q21	CAG	13–36/55–86	AD	Ataxin 3
Spinocerebellar ataxia type 6 (SCA6, CACNA1A)	19p13.1-13.2	CAG	4–16/20–33	AD	Alpha 1A voltage-dependent L-type calcium channel
Spinocerebellar ataxia type 7 (SCA7)	3p21.1-p12	CAG	4–19/37 to >300	AD	Ataxin 7
Spinocerebellar ataxia type 12 (SCA12)	5q31	CAG	6–26/66–78	AD	Protein phosphatase 2A
Dentatorubral pallidolusian atrophy (DRPLA)	12p	CAG	7–23/49–75	AD	Atrophin 1
Friedreich's ataxia (FRDA1)	9q13-21	GAA	7–22/200–900	AR	Frataxin

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; XR, X-linked recessive.

TABLE 82-5 GENES AND LOCI INVOLVED IN MONO- AND POLYGENIC FORMS OF DIABETES

Disorder	Genes or Susceptibility Locus	Chromosomal Location	Other Factors	
Monogenic permanent neonatal diabetes mellitus	<i>KCNJ11</i> (inwardly rectifying potassium channel Kir6.2)	11p15.1	AD	
	<i>GCK</i> (glucokinase)	7p15-p13	AR	
	<i>INS</i> (insulin)	11p15.5	AR, hyperproinsulinemia	
	<i>ABCC8</i> (ATP-binding cassette, subfamily c, member 8; sulfonylurea receptor)	11p15.1	AD or AR	
Maturity-onset diabetes of the young (MODY): Monogenic forms of diabetes mellitus	<i>GLIS3</i> (GLIS family zinc finger protein 3)	9p24.2	AR, diabetes, congenital hypothyroidism	
	MODY 1	<i>HNF4a</i> (hepatocyte nuclear factor 4a)	20q12-q13.1	AD inheritance
	MODY 2	<i>GCK</i> (glucokinase)	7p15-p13	
	MODY 3	<i>HNF1a</i> (hepatocyte nuclear factor 1a)	12q24.2	
	MODY 4	<i>IPF1</i> (insulin receptor substrate)	13q12.1	
	MODY 5 (renal cysts, diabetes)	<i>HNF1β</i> (hepatocyte nuclear factor 1β)	17cen-q21.3	
	MODY 6	<i>NeuroD1</i> (neurogenic differentiation factor 1)	2q32	
	MODY 7	<i>KLF1</i> (Kruppel-like factor 1)	19p13.13-p13.12	
	MODY 8	<i>CEL</i> (carboxyl ester lipase)	9q34.3	
	MODY 9	<i>PAX4</i> (paired box transcription factor 4)	7q32	
	MODY 10	<i>INS</i> (insulin)	11p15.5	
MODY 11	<i>BLK</i> (B-lymphocyte-specific tyrosine kinase)	8p23-p22		
Diabetes mellitus type 2; loci and genes linked and/or associated with susceptibility for diabetes mellitus type 2	Genes and loci identified by linkage/association studies		Heavily influenced by diet, energy expenditure, obesity	
	<i>PPARG, KCNJ11/ABCC8, TCF7L2, IGF2BP2, CDKAL1, SLC30A8, CDKN2A/B, HHEX, FTO, HNF1B, NOTCH2, THADA, ADAMSTS9, JAZF1, CDC122/CAMK1D, KCNQ1, TSPAN8/LGR5, IRS1, DUSP9, PROX1, BCK11A, G6PC2, GCKR, ADCY5, SLC2A2, WFS1, ZBED3, DGKB/TMEM195, GCK, KLF14, TP53INP1, GLIS3, TLE4, ADRA2A, CENTD2, CRY2, FADS1, MADD, MTNR1B, HMGA1, HNF1A, IGF1A, IGF1, C2CD4B, PRC1, VPS13C, ZFAND6, GIPR</i>			

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; MODY, maturity onset diabetes of the young.

The identification of genetic variations and environmental factors that either predispose to or protect against disease is essential for predicting disease risk, designing preventive strategies, and developing novel therapeutic approaches. The study of rare monogenic diseases may provide insight into some of the genetic and molecular mechanisms important in the pathogenesis of complex diseases. For example, the identification of the genes causing monogenic forms of permanent neonatal diabetes mellitus or maturity-onset diabetes defined them as *candidate genes* in the pathogenesis of diabetes mellitus type 2 (Tables 82-2 and 82-5). Genome scans have identified numerous genes and loci that may be associated with susceptibility to development of diabetes mellitus in certain populations. Efforts to identify susceptibility genes require very large sample sizes, and positive results may depend on ethnicity, ascertainment criteria, and statistical analysis. Association studies analyzing the potential influence of (biologically functional) SNPs and SNP haplotypes on a particular phenotype are providing new insights into the genes involved in the pathogenesis of these common disorders. Large variants ([micro]deletions, duplications, and inversions) present in the human population also contribute to the pathogenesis of complex disorders, but their contributions remain poorly understood.

Linkage and Association Studies There are two primary strategies for mapping genes that cause or increase susceptibility to human disease: (1) classic linkage can be performed based on a known genetic model or, when the model is unknown, by studying pairs of affected relatives; or (2) disease genes can be mapped using allelic association studies (Table 82-6).

GENETIC LINKAGE *Genetic linkage* refers to the fact that genes are physically connected, or linked, to one another along the chromosomes.

Two fundamental principles are essential for understanding the concept of linkage: (1) when two genes are close together on a chromosome, they are usually transmitted together, unless a recombination event separates them (Figs. 82-6); and (2) the odds of a crossover, or recombination event, between two linked genes is proportional to the distance that separates them. Thus, genes that are farther apart are more likely to undergo a recombination event than genes that are very close together. The detection of chromosomal loci that segregate with a disease by linkage can be used to identify the gene responsible for the disease (*positional cloning*) and to predict the odds of disease gene transmission in genetic counseling.

Polymorphisms are essential for linkage studies because they provide a means to distinguish the maternal and paternal chromosomes in an individual. On average, 1 out of every 1000 bp varies from one person to the next. Although this degree of variation seems low (99.9% identical), it means that >3 million sequence differences exist between any two unrelated individuals and the probability that the sequence at such loci will differ on the two homologous chromosomes is high (often >70–90%). These sequence variations include variable number of tandem repeats (VNTRs), short tandem repeats (STRs), and SNPs. Most STRs, also called *polymorphic microsatellite markers*, consist of di-, tri-, or tetranucleotide repeats that can be characterized readily using the polymerase chain reaction (PCR). Characterization of SNPs, using DNA chips or beads, permits comprehensive analyses of genetic variation, linkage, and association studies. Although these sequence variations often have no apparent functional consequences, they provide much of the basis for variation in genetic traits.

In order to identify a chromosomal locus that segregates with a disease, it is necessary to characterize polymorphic DNA markers from affected and unaffected individuals of one or several pedigrees. One can

TABLE 82-6 GENETIC APPROACHES FOR IDENTIFYING DISEASE GENES

Method	Indications and Advantages	Limitations
Linkage Studies		
Classical linkage analysis (parametric methods)	Analysis of monogenic traits	Difficult to collect large informative pedigrees
	Suitable for genome scan	Difficult to obtain sufficient statistical power for complex traits
	Control population not required	
	Useful for multifactorial disorders in isolated populations	
Allele-sharing methods (nonparametric methods)	Suitable for identification of susceptibility genes in polygenic and multifactorial disorders	Difficult to collect sufficient number of subjects
Affected sib and relative pair analyses	Suitable for genome scan	Difficult to obtain sufficient statistical power for complex traits
Sib pair analysis	Control population not required if allele frequencies are known	Reduced power compared to classical linkage, but not sensitive to specification of genetic mode
	Statistical power can be increased by including parents and relatives	
Association Studies		
Case-control studies	Suitable for identification of susceptibility genes in polygenic and multifactorial disorders	Requires large sample size and matched control population
Linkage disequilibrium	Suitable for testing specific allelic variants of known candidate loci	False-positive results in the absence of suitable control population
Transmission disequilibrium test (TDT)	Facilitated by HapMap data, making GWAS more feasible	Candidate gene approach does not permit detection of novel genes and pathways
Whole-genome association studies	Does not necessarily need relatives	Susceptibility genes can vary among different populations

Abbreviation: GWAS, genome-wide association study.

then assess whether certain marker alleles cosegregate with the disease. Markers that are closest to the disease gene are less likely to undergo recombination events and therefore receive a higher linkage score. Linkage is expressed as a lod (logarithm of odds) score—the ratio of the probability that the disease and marker loci are linked rather than unlinked. Lod scores of +3 (1000:1) are generally accepted as supporting linkage, whereas a score of –2 is consistent with the absence of linkage.

ALLELIC ASSOCIATION, LINKAGE DISEQUILIBRIUM, AND HAPLOTYPES *Allelic association* refers to a situation in which the frequency of an allele is significantly increased or decreased in individuals affected by a particular disease in comparison to controls. Linkage and association differ in several aspects. Genetic linkage is demonstrable in families or sibships. Association studies, on the other hand, compare a population of affected individuals with a control population. Association studies can be performed as case-control studies that include unrelated affected individuals and matched controls or as family-based studies that compare the frequencies of alleles transmitted or not transmitted to affected children.

Allelic association studies are particularly useful for identifying susceptibility genes in complex diseases. When alleles at two loci occur more frequently *in combination* than would be predicted (based on known allele frequencies and recombination fractions), they are said to be in *linkage disequilibrium*. Evidence for linkage disequilibrium can be helpful in mapping disease genes because it suggests that the two loci are tightly linked.

Detecting the genetic factors contributing to the pathogenesis of common complex disorders remains a great challenge. In many instances, these are low-penetrance alleles (e.g., variations that individually have a subtle effect on disease development, and they can only be identified by unbiased GWAS) (Catalog of Published Genome-Wide Association Studies; Table 82-1) (Fig. 82-14). Most variants occur in noncoding or regulatory sequences but do not alter protein structure. The analysis of complex disorders is further complicated by ethnic differences in disease prevalence, differences in allele frequencies in known susceptibility genes among different populations, locus and allelic heterogeneity, gene-gene and gene-environment interactions, and the possibility of phenocopies. The data generated by the HapMap Project are greatly facilitating GWAS for the characterization of complex disorders. Adjacent SNPs are inherited together

as blocks, and these blocks can be identified by genotyping selected marker SNPs, so-called *Tag SNPs*, thereby reducing cost and workload (Fig. 82-4). The availability of this information permits the characterization of a limited number of SNPs to identify the set of haplotypes present in an individual (e.g., in cases and controls). This, in turn, permits performing GWAS by searching for associations of certain haplotypes with a disease phenotype of interest, an essential step for unraveling the genetic factors contributing to complex disorders.

POPULATION GENETICS In population genetics, the focus changes from alterations in an individual's genome to the distribution pattern of different genotypes in the population. In a case where there are only two alleles, A and a, the frequency of the genotypes will be $p^2 + 2pq + q^2 = 1$, with p^2 corresponding to the frequency of AA, $2pq$ to the frequency of Aa, and q^2 to aa. When the frequency of an allele is known, the frequency of the genotype can be calculated. Alternatively, one can determine an allele frequency if the genotype frequency has been determined.

Allele frequencies vary among ethnic groups and geographic regions. For example, heterozygous mutations in the *CFTR* gene are relatively common in populations of European origin but are rare in the African population. Allele frequencies may vary because certain allelic variants confer a selective advantage. For example, heterozygotes for the sickle cell mutation, which is particularly common in West Africa, are more resistant to malarial infection because the erythrocytes of heterozygotes provide a less favorable environment for *Plasmodium* parasites. Although homozygosity for the sickle cell mutation is associated with severe anemia and sickle crises (Chap. 127), heterozygotes have a higher probability of survival because of the reduced morbidity and mortality from malaria; this phenomenon has led to an increased frequency of the mutant allele. Recessive conditions are more prevalent in geographically isolated populations because of the more restricted gene pool.

APPROACH TO THE PATIENT: Inherited Disorders

For the practicing clinician, the family history remains an essential step in recognizing the possibility of a hereditary predisposition to disease. When taking the history, it is useful to draw a detailed

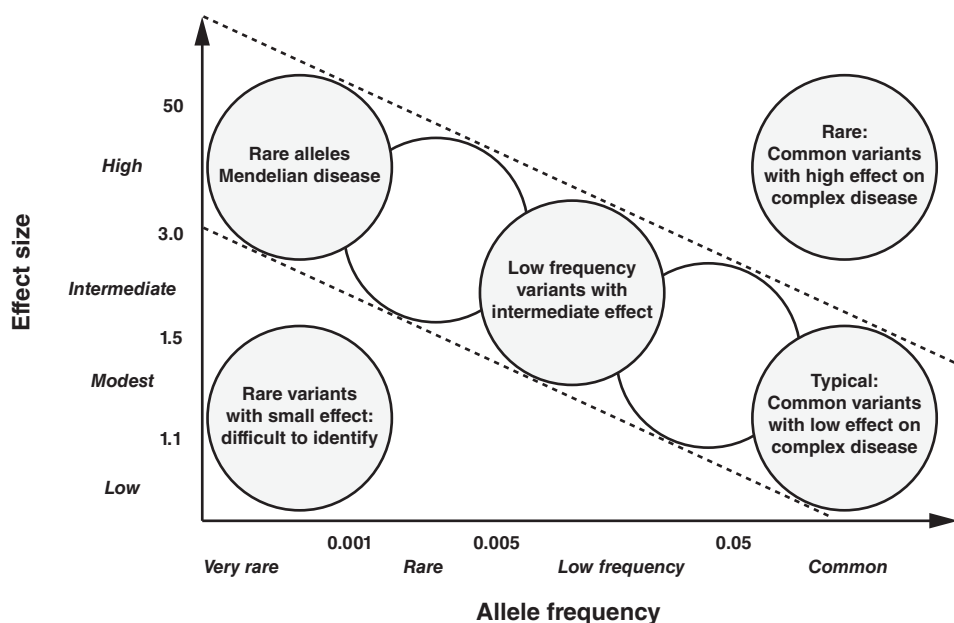


FIGURE 82-14 Relationship between allele frequency and effect size in monogenic and polygenic disorders. In classic Mendelian disorders, the allele frequency is typically low but has a high impact (single gene disorder). This contrasts with polygenic disorders that require the combination of multiple low impact alleles that are frequently quite common in the general population.

pedigree of the first-degree relatives (e.g., parents, siblings, and children), because they share 50% of genes with the patient. Standard symbols for pedigrees are depicted in Fig. 82-11. The family history should include information about ethnic background, age, health status, and deaths, including infants. Next, the physician should explore whether there is a family history of the same or related illnesses to the current problem. An inquiry focused on commonly occurring disorders such as cancers, heart disease, and diabetes mellitus should follow. Because of the possibility of age-dependent expressivity and penetrance, the family history will need intermittent updating. If the findings suggest a genetic disorder, the clinician should assess whether some of the patient's relatives may be at risk of carrying or transmitting the disease. In this circumstance, it is useful to confirm and extend the pedigree based on input from several family members. This information may form the basis for genetic counseling, carrier detection, early intervention, and disease prevention in relatives of the index patient (Chap. 84).

In instances where a diagnosis at the molecular level may be relevant, it is important to identify an appropriate laboratory that can perform the appropriate test. Genetic testing is available for a rapidly growing number of monogenic disorders through commercial laboratories. For uncommon disorders, the test may only be performed in a specialized research laboratory. Approved laboratories offering testing for inherited disorders can be identified in continuously updated online resources (e.g., GeneTests; Table 82-1). If genetic testing is considered, the patient and the family should be counseled about the potential implications of positive results, including psychological distress and the possibility of discrimination. The patient or caretakers should be informed about the meaning of a negative result, technical limitations, and the possibility of false-negative and inconclusive results. For these reasons, genetic testing should only be performed after obtaining *informed consent*. Published ethical guidelines address the specific aspects that should be considered when testing children and adolescents. Genetic testing should usually be limited to situations in which the results may have an impact on medical management.

IDENTIFYING THE DISEASE-CAUSING GENE

Genomic medicine aims to enhance the quality of medical care through the use of genotypic analysis (DNA testing) to identify genetic predisposition to disease, to select more specific pharmacotherapy, and to design individualized medical care based on genotype.

Genotype can be deduced by analysis of protein (e.g., hemoglobin, apoprotein E), mRNA, or DNA. However, technologic advances have made DNA analysis particularly useful because it can be readily applied.

DNA testing is performed by mutational analysis or linkage studies in individuals at risk for a genetic disorder known to be present in a family. Mass screening programs require tests of high sensitivity and specificity to be cost-effective. Prerequisites for the success of genetic screening programs include the following: that the disorder is potentially serious; that it can be influenced at a presymptomatic stage by changes in behavior, diet, and/or pharmaceutical manipulations; and that the screening does not result in any harm or discrimination. Screening in Jewish populations for the autosomal recessive neurodegenerative storage disease Tay-Sachs has reduced the number of affected individuals. In contrast, screening for sickle cell trait/disease in African Americans has led to unanticipated problems of discrimination by health insurers and employers. Mass screening programs harbor additional potential problems. For example, screening for the most common genetic alteration in cystic fibrosis, the $\Delta F508$ mutation with a frequency of ~70% in northern Europe, is feasible and seems to be effective. One has to keep in mind, however, that there is pronounced allelic heterogeneity and that the disease can be caused by about 2000 other mutations. The search for these less common mutations would substantially increase costs but not the effectiveness of the screening program as a whole. Next-generation genome sequencing permits comprehensive and cost-effective mutational analyses after selective enrichment of candidate genes. For example, tests that sequence all the common genes causing hereditary deafness are already commercially available. Occupational screening programs aim to detect individuals with increased risk for certain professional activities (e.g., α_1 antitrypsin deficiency and smoke or dust exposure). Integrating genomic data into electronic medical records is evolving and may provide significant decision support at the point of care, for example, by providing the clinician with genomic data and decision algorithms for the prescription of drugs that are subject to pharmacogenetic influences.

Mutational Analyses DNA sequence analysis is now widely used as a diagnostic tool and has significantly enhanced diagnostic accuracy. It is used for determining carrier status and for prenatal testing in monogenic disorders (Chap. 84). Numerous techniques,

discussed in previous versions of this chapter, are available for the detection of mutations. In a very broad sense, one can distinguish between techniques that allow for screening of known mutations (screening mode) or techniques that definitively characterize mutations. Analyses of large alterations in the genome are possible using classic methods such as cytogenetics, fluorescent in situ hybridization (FISH), and Southern blotting (Chap. 83e), as well as more sensitive novel techniques that search for multiple single exon deletions or duplications.

More discrete sequence alterations rely heavily on the use of PCR, which allows rapid gene amplification and analysis. Moreover, PCR makes it possible to perform genetic testing and mutational analysis with small amounts of DNA extracted from leukocytes or even from single cells, buccal cells, or hair roots. DNA sequencing can be performed directly on PCR products or on fragments cloned into plasmid vectors amplified in bacterial host cells. Sequencing of all exons of the genome or selected chromosomes, or sequencing of numerous candidate genes in a single run, is now possible with next-generation sequencing platforms.

The majority of traditional diagnostic methods were gel-based. Novel technologies for the analysis of mutations, genotyping, large-scale sequencing, and mRNA expression profiles are undergoing rapid evolution. DNA chip technologies allow hybridization of DNA or RNA to hundreds of thousands of probes simultaneously. Microarrays are being used clinically for mutational analysis of several human disease genes, as well as for the identification of viral or bacterial sequence variations. With advances in high-throughput DNA sequencing technology, complete sequencing of the genome or an exome has entered the clinical realm. Although comprehensive sequencing of large genomic regions or multiple genes is already a reality, the subsequent bioinformatics analysis, assembly of sequence fragments, and comparative alignments remains a significant and commonly underestimated challenge. The discovery of incidental (or secondary) findings that are unrelated to the indication for the sequencing analysis but indicators of other disorders of potential relevance for patient care can pose a difficult ethical dilemma. It can lead to the detection of undiagnosed medically actionable genetic conditions, but can also reveal deleterious mutations that cannot be influenced, as numerous sequence variants are of unknown significance.

A general algorithm for the approach to mutational analysis is outlined in Fig. 82-15. The importance of a detailed clinical phenotype cannot be overemphasized. This is the step where one should also consider the possibility of genetic heterogeneity and phenocopies. If obvious candidate genes are suggested by the phenotype, they

can be analyzed directly. After identification of a mutation, it is essential to demonstrate that it segregates with the phenotype. The functional characterization of novel mutations is labor intensive and may require analyses in vitro or in transgenic models in order to document the relevance of the genetic alteration.

Prenatal diagnosis of numerous genetic diseases in instances with a high risk for certain disorders is now possible by direct DNA analysis. *Amniocentesis* involves the removal of a small amount of amniotic fluid, usually at 16 weeks of gestation. Cells can be collected and submitted for karyotype analyses, FISH, and mutational analysis of selected genes. The main indications for amniocentesis include advanced maternal age (>35 years), an abnormal serum triple marker test (α -fetoprotein, β human chorionic gonadotropin, pregnancy-associated plasma protein A, or unconjugated estriol), a family history of chromosomal abnormalities, or a Mendelian disorder amenable to genetic testing. Prenatal diagnosis can also be performed by *chorionic villus sampling* (CVS), in which a small amount of the chorion is removed by a transcervical or transabdominal biopsy. Chromosomes and DNA obtained from these cells can be submitted for cytogenetic and mutational analyses. CVS can be performed earlier in gestation (weeks 9–12) than amniocentesis, an aspect that may be of relevance when termination of pregnancy is a consideration. Later in pregnancy, beginning at about 18 weeks of gestation, percutaneous umbilical blood sampling (PUBS) permits collection of fetal blood for lymphocyte culture and analysis. Recently, the entire fetal genome has been determined prenatally from cells taken from the mother's plasma through deep sequencing and the counting of parental haplotypes, or by inferring it from DNA sequences obtained from blood samples from the mother, father, and umbilical cord. These approaches enable screening for clinically relevant and deleterious alleles inherited from the parents, as well as for de novo germline mutations, and they may have the potential to change the diagnosis of genetic disorders in the prenatal setting.

In combination with in vitro fertilization (IVF) techniques, it is even possible to perform genetic diagnoses in a single cell removed from the four- to eight-cell embryo or to analyze the first polar body from an oocyte. Preconceptual diagnosis thereby avoids therapeutic abortions but is costly and labor intensive. It should be emphasized that excluding a specific disorder by any of these approaches is never equivalent to the assurance of having a normal child.

Mutations in certain cancer susceptibility genes such as *BRCA1* and *BRCA2* may identify individuals with an increased risk for the development of malignancies and result in risk-reducing interventions. The detection of mutations is an important diagnostic and

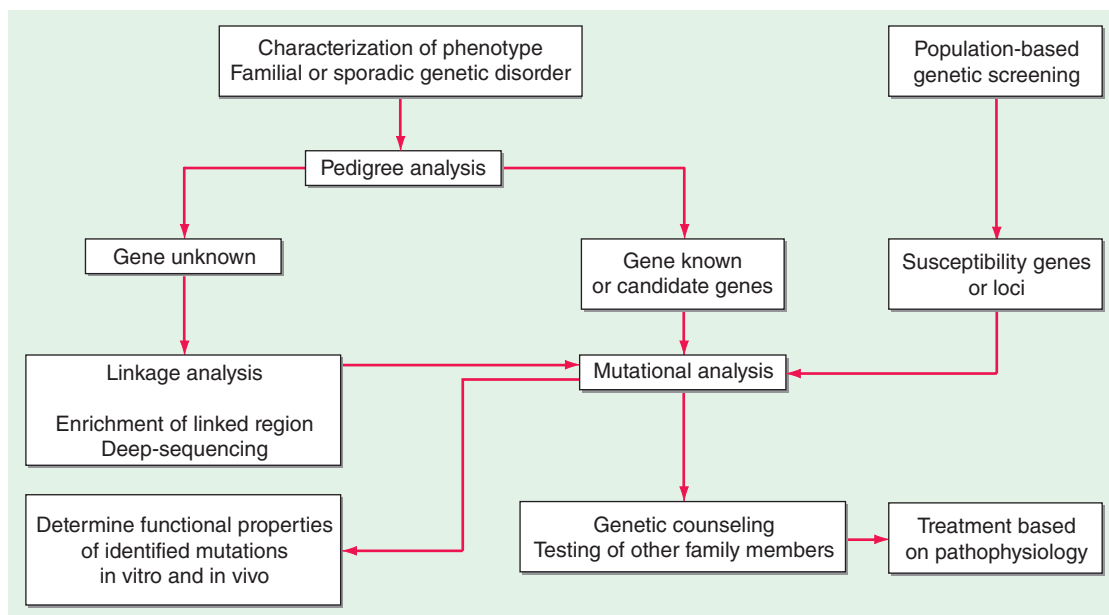


FIGURE 82-15 Approach to genetic disease.

prognostic tool in leukemias and lymphomas. The demonstration of the presence or absence of mutations and polymorphisms is also relevant for the rapidly evolving field of pharmacogenomics, including the identification of differences in drug treatment response or metabolism as a function of genetic background. For example, the thiopurine drugs 6-mercaptopurine and azathioprine are commonly used cytotoxic and immunosuppressive agents. They are metabolized by thiopurine methyltransferase (TPMT), an enzyme with variable activity associated with genetic polymorphisms in 10% of whites and complete deficiency in about 1 in 300 individuals. Patients with intermediate or deficient TPMT activity are at risk for excessive toxicity, including fatal myelosuppression. Characterization of these polymorphisms allows mercaptopurine doses to be modified based on TPMT genotype. Pharmacogenomics may increasingly permit individualized drug therapy, improve drug effectiveness, reduce adverse side effects, and provide cost-effective pharmaceutical care (Chap. 5).

ETHICAL ISSUES

Determination of the association of genetic defects with disease, comprehensive data of an individual's genome, and studies of genetic variation raise many ethical and legal issues. Genetic information is generally regarded as sensitive information that should not be readily accessible without explicit consent (*genetic privacy*). The disclosure of genetic information may risk possible discrimination by insurers or employers. The scientific components of the Human Genome Project have been paralleled by efforts to examine ethical, social, and legal implications. An important milestone emerging from these endeavors consists in the Genetic Information Nondiscrimination Act (GINA), signed into law in 2008, which aims to protect asymptomatic individuals against the misuse of genetic information for health insurance and employment. It does not, however, protect the symptomatic individual. Provisions of the U.S. Patient Protection and Affordable Care Act, effective in 2014, will fill this gap and prohibit exclusion from, or termination of, health insurance based on personal health status. Potential threats to the maintenance of genetic privacy consist in the emerging integration of genomic data into electronic medical records, compelled disclosures of health records, and direct-to-consumer genetic testing.

It is widely accepted that identifying disease-causing genes can lead to improvements in diagnosis, treatment, and prevention. However, the information gleaned from genotypic results can have quite different impacts, depending on the availability of strategies to modify the course of disease (Chap. 84). For example, the identification of mutations that cause MEN 2 or hemochromatosis allows specific interventions for affected family members. On the other hand, at present, the identification of an Alzheimer's or Huntington's disease gene does not currently alter therapy and outcomes. Most genetic disorders are likely to fall into an intermediate category where the opportunity for prevention or treatment is significant but limited (Chap. 84). However, the progress in this area is unpredictable, as underscored by the finding that angiotensin II receptor blockers may slow disease progression in Marfan's syndrome. Genetic test results can generate anxiety in affected individuals and family members. Comprehensive sequence analyses are particularly challenging because most individuals can be expected to harbor several serious recessive gene mutations.

The impact of genetic testing on health care costs is currently unclear. It is likely to vary among disorders and depend on the availability of effective therapeutic modalities. A significant problem arises from the marketing of genetic testing directly to consumers by commercial companies. The validity of these tests has not been defined, and there are numerous concerns about the lack of appropriate regulatory oversight, the accuracy and confidentiality of genetic information, the availability of counseling, and the handling of these results.

Many issues raised by the genome project are familiar, in principle, to medical practitioners. For example, an asymptomatic

patient with increased low-density lipoprotein (LDL) cholesterol, high blood pressure, or a strong family history of early myocardial infarction is known to be at increased risk of coronary heart disease. In such cases, it is clear that the identification of risk factors and an appropriate intervention are beneficial. Likewise, patients with phenylketonuria, cystic fibrosis, or sickle cell anemia are often identified as having a genetic disease early in life. These precedents can be helpful for adapting policies that relate to genetic information. We can anticipate similar efforts, whether based on genotypes or other markers of genetic predisposition, to be applied to many disorders. One confounding aspect of the rapid expansion of information is that our ability to make clinical decisions often lags behind initial insights into genetic mechanisms of disease. For example, when genes that predispose to breast cancer such as *BRCA1* are described, they generate tremendous public interest in the potential to predict disease, but many years of clinical research are still required to rigorously establish genotype and phenotype correlations.

Genomics may contribute to improvements in global health by providing a better understanding of pathogens and diagnostics, and through contributions to drug development. There is, however, concern about the development of a "genomics divide" because of the costs associated with these developments and uncertainty as to whether these advances will be accessible to the populations of developing countries. The World Health Organization has summarized the current issues and inequities surrounding genomic medicine in a detailed report titled "Genomics and World Health."

Whether related to informed consent, participation in research, or the management of a genetic disorder that affects an individual or his or her family, there is a great need for more information about fundamental principles of genetics. The pervasive nature of the role of genetics in medicine makes it important for physicians and other health care professionals to become more informed about genetics and to provide advice and counseling in conjunction with trained genetic counselors (Chap. 84). The application of screening and prevention strategies will therefore require intensive patient and physician education, changes in health care financing, and legislation to protect patient's rights.

CHROMOSOME DISORDERS

Alterations of the chromosomes (numerical and structural) occur in about 1% of the general population, in 8% of stillbirths, and in close to 50% of spontaneously aborted fetuses. The 3×10^9 base pairs that encode the human genome are packaged into 23 pairs of chromosomes, which consist of discrete portions of DNA, bound to several classes of regulatory proteins. Technical advances that led to the ability to analyze human chromosomes immediately translated into the revelation that human disorders can be caused by an abnormality of chromosome number. In 1959, the clinically recognizable disorder, Down syndrome, was demonstrated to result from having three copies of chromosome 21 (trisomy 21). Very soon thereafter, in 1960, a small, structurally abnormal chromosome was recognized in the cells of some patients with chronic myelogenous leukemia (CML), and this abnormal chromosome is now known as the Philadelphia chromosome.

Since these early discoveries, the techniques for analysis of human chromosomes, and DNA in general, have gone through several revolutions, and with each technical advancement, our understanding of the role of chromosomal abnormalities in human disease has expanded. While early studies in the 1950s and 1960s easily identified abnormalities of chromosome number (aneuploidy) and large structural alterations such as deletions (chromosomes with missing regions), duplications (extra copies of chromosome regions), or translocations (where portions of the chromosomes are rearranged), many other types of structural alterations could only be identified as techniques improved. The first important technical advance was the introduction of chromosome banding in the late 1960s, a technique that allowed for the staining of the chromosomes, so that each chromosome could be recognized by its pattern of alternating dark and light (or fluorescent and nonfluorescent) bands. Other technical innovations ranged from the introduction of fluorescence *in situ* hybridization in the 1980s to use of array-based and sequencing technologies in the early 2000s. Currently, we can appreciate that many types of chromosome abnormalities contribute to human disease including aneuploidy; structural alterations such as deletions and duplications, translocations, or inversions; uniparental disomy, where two copies of one chromosome (or a portion of a chromosome) are inherited from one parent; complex alterations such as isochromosomes, markers, and rings; and mosaicism for all of the aforementioned abnormalities. The first chromosome disorders identified had very striking and generally severe phenotypes, because the abnormalities involved large regions of the genome, but as methods have become more sensitive, it is now possible to recognize many more subtle phenotypes, often involving smaller genomic regions.

METHODS FOR CHROMOSOME ANALYSIS

STANDARD CYTOGENETIC ANALYSIS

Standard cytogenetic analysis refers to the examination of banded human chromosomes. Banded chromosome analysis allows for both the determination of the number and identity of chromosomes in the cell and recognition of abnormal banding patterns associated with a structural rearrangement. A stained band is defined as the part of a chromosome that is clearly distinguishable from its adjacent segments by appearing darker or lighter with one or more banding techniques. Cytogenetic analysis is most commonly carried out on cells in mitosis, requiring dividing cells. Actively growing cells are most often obtained from peripheral blood; however, it is only a small subset of the blood cells that are actually used for cytogenetic analysis. Often, chemicals, like phytohemagglutinin (PHA), are used to specially stimulate growth of T cells in a blood sample. Other sources of dividing cells include skin-derived fibroblasts, amniotic fluid or placental tissue (for prenatal diagnosis), or tumor tissue (for cancer diagnosis). After culturing, cells are treated with a mitotic spindle inhibitor, which prevents

the separation of the chromatids during metaphase. Halting mitosis in metaphase is essential, because chromosomes are at their most condensed state during this stage of mitosis. The banding pattern of a metaphase chromosome is easily recognizable and is ideal for karyotyping. There are several different types of chromosome staining techniques, including R-banding, C-banding, and quinacrine staining, but the most commonly used is G-banding. G-banding is accomplished by treatment of the chromosomes with a proteolytic enzyme, such as trypsin, which digests some of the proteins holding DNA in a three-dimensional structure, followed by staining with a dye (Giemsa) that binds DNA. The resulting patterns have both dark and light bands; in general, the light bands occur in regions on the chromosome in which genes are actively being transcribed, and dark bands are in regions of less active transcription.

The banded human karyotype has now been standardized based on an internationally agreed upon system for designating not only individual chromosomes but also chromosome regions, providing a way in which structural rearrangements and variants can be described in terms of their composition. The normal human female karyotype is referred to as 46,XX (46 chromosomes, with 22 pairs of autosomes and two of the same type of sex chromosomes [two Xs], indicating this is a female); and the normal human male karyotype is referred to as 46,XY (46 chromosomes, with 22 pairs of autosomes and one of each type of sex chromosome [one X and one Y], indicating this is a male). The anatomy of a chromosome includes the central constriction, known as the centromere, which is critical for movement of the chromosomes during mitosis and meiosis; the two chromosome arms (p for the smaller or petite arm, and q for the longer arm); and the chromosome ends, which contain the telomeres. The telomeres are made up of a hexanucleotide repeat (TTAGGG)_n, and unlike the centromere, they are not visible at the light microscope level. Telomeres are functionally important because they confer stability to the end of the chromosome. Broken chromosomes tend to fuse end to end, whereas a normal chromosome with an intact telomere structure is stable. To create the standard chromosome-banding map, each chromosome is divided into segments that are numbered, and then further subdivided. The precise band names are recorded in an international document so that each band has a distinct number. **Figure 83e-1** shows an ideogram (chromosome map with bands) of the X chromosome and a G-banded X chromosome. This system provides a way for a chromosome abnormality to be written, with an indication of which band is deleted, duplicated, or rearranged.

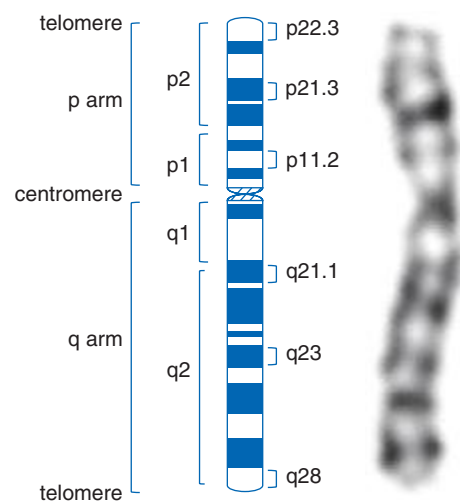


FIGURE 83e-1 Ideogram of the X chromosome and a G-banded X chromosome. The labeling of the X ideogram shows the positioning of the p and q arms, the centromere, and the telomeres. The numbering of the bands is also demonstrated, indicating the broadest subbands (p1, p2, q1, q2) and the further subdivisions to the right. Numbering begins at the centromere and moves out along each arm toward the telomeres.

Molecular cytogenetics provides a link between chromosome and molecular analysis and overcomes some of the limitations of standard cytogenetics. Deletions smaller than several million base pairs are not routinely detectable by standard G-banding techniques, and chromosomal abnormalities with indistinct or novel banding patterns can be difficult or impossible to interpret. To carry out cytogenetic analysis, cells must be dividing, which is not always possible to obtain (e.g., in autopsy or tumor material that has already been fixed). Finally, growth selection or bias may occasionally cause the results of cytogenetic studies to be misleading because cells that proliferate *in vitro* may not be representative of the original population, as is often the case with tumor specimens.

Fluorescence *in situ* hybridization (FISH) is a combined cytogenetic-molecular technique that solves many of the aforementioned problems. FISH permits determination of the number and location of specific DNA sequences in human cells. FISH can be performed on metaphase chromosomes, as with G-banding, but can also be performed on cells not actively progressing through mitosis. FISH performed on nondividing cells is referred to as interphase or nuclear FISH (Fig. 83e-2). The FISH procedure relies on the complementarity between the two strands of the DNA double helix and uses a molecular probe, which can be a pool of sequences across an entire chromosome, a DNA sequence for a repetitive part of the genome (e.g., centromeres or telomeres), or a specific DNA sequence found only once in the genome (e.g., a disease-associated gene). The choice of probes for FISH studies is important and will vary with the information needed for the diagnosis of a particular disorder. The most common type of probes are locus-specific probes, which are used to determine if a critical gene or region is absent (indicating a deletion), or present in the normal

number of copies, or if an additional copy of the region is present. FISH on metaphase chromosomes will give the additional information of the location of the additional copy, which is necessary information to determine whether a structural rearrangement, such as a translocation, is present. FISH can also be performed with probes that bind to repeated sequences, such as DNA found in centromeres or telomeres, or with probes that bind to an entire chromosome ("painting" probes), to determine the chromosome composition of an abnormal chromosome. Interphase FISH studies can also help to identify structural alterations when probes are used that map to both sides of a translocation breakpoint. Each side of the breakpoint is labeled in a different color, and when no translocation is present the two probes appear to be overlapping. When a translocation is present, the two probes appear separate from one another. These set of probes, called "break-apart" probes, are commonly used to detect recurrent translocations in cancer cells.

ARRAY-BASED METHODOLOGIES (CYTOGENOMICS)

Array-based methods were introduced into the clinical lab beginning in 2003 and quickly revolutionized the field of cytogenetics. These techniques used arrays (collections of DNA segments from the entire genome) which could be interrogated with respect to copy number. With standard cytogenetics, the missing or extra pieces of DNA have to be big enough to see in the microscope on banded chromosomes (usually larger than 5 Mb). FISH requires a preselection of an informative molecular probe prior to analysis. In contrast, array-based techniques permit analysis of many regions of the genome in a single analysis, with greatly increased resolution over standard cytogenetics. Array-based techniques allow for scanning of the genome for small deletions or duplications quickly and accurately. The resolution of the

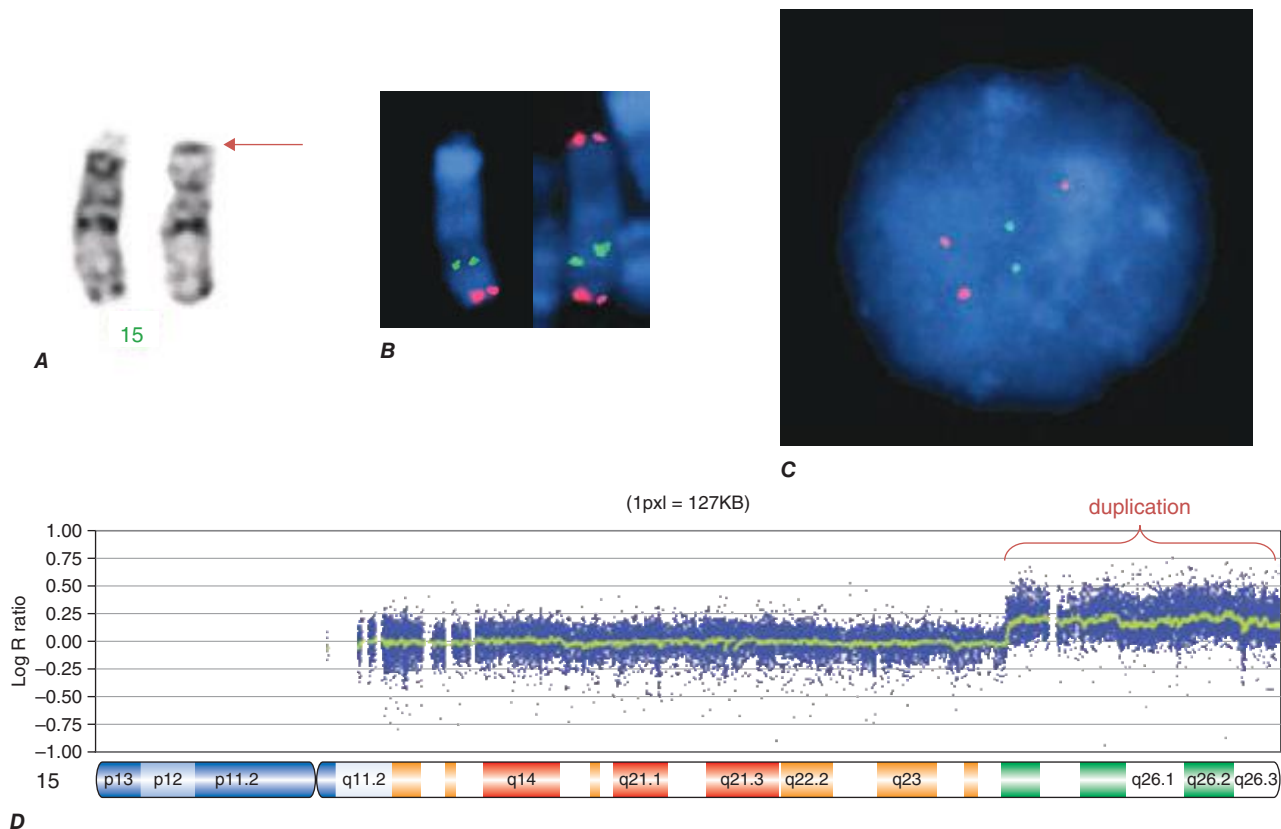


FIGURE 83e-2 G-banding, fluorescence *in situ* hybridization (FISH), and single nucleotide polymorphism (SNP) array demonstrate an abnormal chromosome 15. **A.** G-banding shows an abnormal chromosome 15, with unrecognizable material in place of the p arm in the chromosome on the right (*top arrow*). **B.** Metaphase FISH (only chromosome 15s are shown) using a probe from the 15q telomere region (*red*) and a control probe that maps outside of the duplicated region (*green*). **C.** Interphase FISH demonstrates three copies of the 15q tel probe in *red*, and two copies of the 15q control probe (*green*). **D.** Genome-wide SNP array demonstrates the increased copy number for a portion of 15q. Note that the G-banding alone indicates the abnormal chromosome 15, but the origin of the extra material can only be demonstrated by FISH or array. The FISH analysis requires additional information about possible genetic causes to select the correct probe. The array can exactly identify the origin of the extra material, but by itself would not provide positional information.

TABLE 83e-1 COMPARISON OF CYTOGENETIC AND CYTOGENOMIC TECHNIQUES

Method	Requires Growing Cells	Detects Deletions and Duplications	Detects Balanced Structural Rearrangements	Detects Uniparental Disomy	Detection Limits (Lower Limit)
G-banding	Yes	Yes	Yes	No	5–10 million bases
Metaphase FISH	Yes	Yes	Yes	No	40–250 thousand bases
Interphase FISH	No	Yes	Some	No	40–250 thousand bases
CGH Array	No	Yes	No	No	Single exon or single gene
SNP Array	No	Yes	No	Some	Single exon or single gene

test is a function of the number of probes or DNA sequences present on the array. Arrays may use probes of different sizes (ranging from 50 to 200,000 base pairs of DNA) and different probe densities depending on the requirements of the application. Low-resolution platforms can have hundreds of probes, targeted to known disease regions, whereas high-resolution platforms can have millions of probes spread across the entire genome. Depending on the size of the probes and the probe placement across the genome, array-based testing may be able to detect single exon deletions or duplications.

Comparative Genomic Hybridization (CGH) and Single Nucleotide Polymorphism (SNP) Analysis CGH and SNP-based genotyping arrays can both be used for the analysis of genomic deletions and duplications. For both techniques, oligonucleotide probes are placed onto a slide or chip in a grid format. Each of these probes is specific for a particular genomic region. In array CGH, the amount of DNA from a patient is compared to that in a clinically normal control, or pool of controls, for each of the probes present on the array. DNA from a patient is fluorescently labeled with a dye of one color, and DNA from a control individual is labeled with another color. These DNA samples are then hybridized at the same time to the array. The resulting fluorescent signal will vary depending on whether both the control and patient DNA are present in equal amounts or if one has a different copy number than the other. SNP platforms use arrays targeting SNPs that are distributed across the genome. SNP arrays vary in density of markers and in the technology used for genotyping, depending on the manufacturer of the array. SNP arrays were initially designed to determine genotypes at a biallelic, polymorphic base (e.g., CC, CT, or TT) and have been increasingly used in genome-wide association studies to identify disease susceptibility genes. SNP arrays were subsequently adapted to identify genomic deletions and duplications (Fig. 83e-2). SNP arrays, in addition to identifying copy number changes, can also detect regions of the genome that have an excess of homozygous genotypes and absence of heterozygous genotypes (e.g., CC and TT genotypes only, with no CT genotypes). Absence of heterozygosity is sometimes associated with uniparental disomy (discussed later in this chapter) but is also observed when an individual's parents are related to one another (identity by descent). Regions of homozygosity have been used to help identify genes in which homozygous mutations result in disease phenotypes in families with known consanguinity.

Array-based techniques (which we will now refer to as cytogenomic analysis) have proven superior to chromosome analysis in the identification of clinically significant deletions or duplications. It is estimated that for a deletion or duplication to be visualized by standard cytogenetics it must be minimally between 5 and 10 million base pairs in size. In almost all cases, deletions and duplications of this size contain multiple genes, and these deletions and duplications are disease causing. However, utilization of array-based cytogenomic testing, which can routinely identify deletions and duplications smaller than 50,000 base pairs, reveals that clinically normal individuals all have some deletions and duplications. This presents a dilemma for the analyst to discern which smaller copy number variations (CNVs) are disease causing (pathogenic) and which are likely benign polymorphisms. Although initially burdensome, the cytogenomics community has been curating these CNVs for almost a decade, and databases have been created reporting CNVs routinely seen in clinically normal individuals and those routinely seen in individuals with clinical abnormalities. Nevertheless, each copy number variant that is identified in an

individual undergoing genomic testing must be evaluated for gene content and overlap with CNVs in other patients and in controls.

Array technologies are DNA based, unlike cytogenetic technologies, which are cell based. Although resolution of gains and losses are greatly increased with array technology, this technique cannot identify structural changes. When DNA is extracted for array studies, chromosomal structure is lost because the DNA is fragmented for better hybridization to the slides. As an example, the array may be able to detect a duplication of a small region of a chromosome, but no information on the location of this extra material can be determined from this test. The location of this extra copy in the genome may be critical, as the chromosomal material may be involved in a translocation, insertion, marker, or other complex rearrangement. Depending on the chromosomal position of this extra material, the patient may have different clinical outcomes, and recurrence risks for the family can be significantly different. Often, combinations of array-based and cytogenetic-based techniques are required to fully characterize chromosomal abnormalities (see [Table 83e-1](#) for comparison of these technologies).

NEXT-GENERATION SEQUENCING—BASED METHODOLOGIES

Recent advances in genomic sequencing, known as next-generation sequencing (NGS), have vastly increased the speed and throughput of DNA sequence analysis. NGS is rapidly finding its way into the diagnostic lab for detection of clinically relevant intragenic mutations, and new bioinformatic tools for analysis of genomic deletions and duplications are being developed. It is anticipated that NGS will soon allow the complete analysis of a patient's genome, with identification of intragenic mutations as well as chromosome abnormalities resulting in gain or loss of genetic material. Identification of completely balanced translocations is the most challenging for NGS, but recent reports of successes in this area suggest that in a matter of time, sequencing will be used for all types of genomic analysis.

INDICATIONS FOR CHROMOSOME/CYTOGENOMIC ANALYSIS

Cytogenetic analysis is most commonly used for (1) examination of the fetal chromosomes or genome during pregnancy (prenatal diagnosis) or in the event of a spontaneous miscarriage; (2) examination of chromosomes in the neonatal or pediatric population to look for an underlying diagnosis in the case of congenital or developmental anomalies, including short stature and abnormalities of sexual differentiation or progression; (3) chromosome analysis in adults who are facing fertility problems; or (4) examination of cancer cells to look for alterations that aid in establishing a diagnosis or contributing to the prognosis of a tumor ([Table 83e-2](#)).

PRENATAL DIAGNOSIS

Prenatal diagnosis is carried out by analysis of samples obtained by four techniques: amniocentesis, chorionic villous sampling, fetal blood sampling, and analysis of cell free DNA from maternal serum. Amniocentesis, which has been the most commonly used test to date, is usually performed between 15 and 17 weeks of gestational age and carries a small but significant risk for miscarriage. Amniocentesis can be performed as early as 12 weeks, but because there is a lower volume of fluid, the risks for fetal injury or miscarriage are greater. Chorionic villous sampling (CVS) or placental biopsy is routinely carried out earlier than amniocentesis, between 10 and 12 weeks, but a reported increase in limb defects when the procedure is carried out earlier than

TABLE 83e-2 INDICATIONS FOR CYTOGENETIC AND CYTOGENOMIC ANALYSIS ACROSS THE LIFESPAN

Timing of Testing	Indications for Testing
Prenatal	Advanced maternal age Abnormalities on ultrasound Increased risk for genetic disorder on maternal serum screen
Neonatal and Childhood	Multiple congenital anomalies Intellectual disability Autism Developmental delay Failure to thrive Short stature Disorders of sexual development History of familial chromosomal alteration Cancer
Adult	Infertility Recurrent miscarriage Cancer

10 weeks has resulted in reduced use of this test in some centers. Fetal blood sampling (percutaneous umbilical blood sampling [PUBS]) is a riskier procedure that is carried out in the second or third trimester of pregnancy, usually to follow up on an unclear finding from an amniocentesis (such as mosaicism) or an ultrasound abnormality that was detected later in pregnancy. One of the far-reaching recent advances in prenatal diagnosis of chromosome and other genetic disorders is the utilization of cell free fetal DNA that can be identified in maternal serum. The obvious advantages of using fetal DNA obtained from maternal serum is that the DNA can be obtained at minimal risk to the pregnancy, because it requires a maternal blood sample, rather than amniotic fluid which is obtained by puncturing the uterine membranes and carries a risk of miscarriage or infection. Although cell free fetal DNA screening, also called noninvasive prenatal screening, has started to be offered clinically, it requires further confirmation of fetal tissues when an abnormal result is identified. Furthermore, ethical concerns have been raised, because it is feared that the ease of doing this test may encourage testing for individuals who are not truly prepared to deal with the choices that accompany diagnosis of a genetic disease and this testing may change the ethical implications of prenatal testing. Nevertheless, this is an active area of research, both in terms of the technology and the utilization and implications.

Common Indications Common indications for prenatal diagnosis by cytogenetic or cytogenomic analysis are (1) advanced maternal age, (2) presence of an abnormality of the fetus on ultrasound examination, and (3) abnormalities in maternal serum screening that reveal an increased risk for chromosome abnormality.

Maternal age is well known to be an important risk factor for having a fetus with trisomy. At a maternal age less than 25 years, 2% of all clinically recognized pregnancies are trisomic, but by a maternal age of 36 years, this figure increases to 10%, and by the maternal age of 42 years, the figure increases to >33%. Based on the risk of having a chromosomally abnormal fetus in comparison to the risk for an adverse event from amniocentesis or CVS, the recommendation is that women over the age of 35 consider prenatal testing if they want to know the chromosomal status of their fetus. The precise mechanism for the maternal age effect is not known, but it is believed that it involves a breakdown in the process of chromosome segregation. A similar effect is not seen for trisomy and paternal age. This difference may reflect the fact that oocytes are generated early in ovary development in the female, whereas spermatogonia are generated continuously after puberty in the male.

Abnormalities on prenatal ultrasound are the second most frequent indication for prenatal genetic screening. Ultrasound screening can

reveal structural or functional anomalies in the fetus, which might be associated with chromosome or genomic disorders. Follow-up chromosome studies may therefore be recommended.

Maternal serum screening results are the third most frequent indication for prenatal chromosome analysis. There have been several versions of maternal serum screening offered over the past few decades. Currently, the “quad” screen analyzes levels of a fetoprotein (AFP), human chorionic gonadotropin (hCG), estriol, and inhibin-A. The values of these analytes are used to adjust the maternal age–predicted risk of a trisomy 21 or trisomy 18 fetus.

POSTNATAL INDICATIONS

Postnatal indications for cytogenetic or cytogenomic analysis in neonates or children are varied, and the list has been growing with the increasing ability to diagnose smaller genomic alterations via array-based techniques. Common indications include multiple congenital anomalies, suspicion of a known cytogenetic or cytogenomic syndrome, intellectual disability or developmental delay both with and without accompanying dysmorphic features, autism, failure to thrive in infancy or short stature during childhood, and disorders of sexual development. The ability to detect smaller genomic alterations with involvement of fewer genes, sometimes as few as a single gene, suggests that a wider range of phenotypes could be investigated by cytogenomic analysis. Reasons for chromosome testing in adults include recurrent miscarriages or infertility, where balanced chromosome rearrangements such as reciprocal translocations may occur. Additionally, some adults with anomalies who were not diagnosed when they were children are referred for cytogenetic analysis, often when other members of their family want to understand any potential genetic implications, as they plan their own families.

TYPES OF CHROMOSOME ABNORMALITIES

NUMERICAL CHROMOSOME ABNORMALITIES

Aneuploidy (extra or missing chromosomes) is the most common type of abnormality, occurring in 3/1000 newborns and at much higher frequency (about 35%) in spontaneously aborted fetuses. The only autosomal trisomies that are compatible with being live born in humans are trisomies 13, 18, and 21, although there are several chromosomes that can be trisomic in mosaic form. Trisomy 21 is associated with the relatively common disorder Down syndrome. Down syndrome has characteristic features including recognizable facial features, along with intellectual disability and abnormalities of multiple other organ systems including the heart. Both trisomy 13 and trisomy 18 are much more severe disorders than Down syndrome, with low frequency of patients surviving past 1 year of age. Trisomy 13 is characterized by low birth weight, postaxial polydactyly, microcephaly, ocular malformations such as anophthalmia or microphthalmia, cleft lip and palate, cardiac defects, and renal malformations. Trisomy 18 neonates have distinct facial characteristics at birth accompanied by an abnormal neurologic exam, underdeveloped genitalia, general lack of responsiveness, and structural birth defects such as congenital heart disease, esophageal atresia, and omphalocele.

Mosaicism refers to the presence of two or more populations of cells with distinct chromosome constitutions: for example, an individual with a normal female karyotype in some cells (46,XX) and trisomy 21 in other cells (47,XX,+21). In general, individuals who are mosaic for a chromosomal abnormality have less severe phenotypes than individuals with that same finding in every cell. The severity and presentation of phenotypes are related to the mosaic levels and the tissue distribution of the abnormal cells. There are a number of trisomies that have been reported in mosaic form including mosaic trisomies for chromosomes 8, 9, 14, 17, and 22. A number of trisomies have also been reported in spontaneous abortions (SABs) that have not been seen in live-born individuals, including trisomy 16, which is the most common trisomy in SABs. Monosomy for human chromosomes is very rare, with the single exception being monosomy for the X chromosome, associated with Turner syndrome (45,X). Monosomy for the X chromosome occurs in 1% of all conceptions, yet 98% of these conceptions do not go to term and result in SABs. Trisomies for

the sex chromosomes also occur, with 47,XXX (trisomy X or triple X syndrome), 47,XXY (Klinefelter syndrome), and 47,XYY all reported in individuals with relatively mild phenotypes (Chap. 410). Klinefelter syndrome is the most common clinically recognized sex chromosome abnormality, and clinical features include gynecomastia, azoospermia, small testes, and hypogonadism. The 47,XYY karyotype is most often found in boys with developmental delay and/or behavioral difficulties, but population-based studies have shown that intelligence for individuals with this karyotype is generally within the normal range, although slightly lower than that found in siblings.

STRUCTURAL CHROMOSOME ABNORMALITIES

Structural chromosome abnormalities include deletions, duplications, translocations, inversions, as well as other types of abnormalities, each relatively rare, but nonetheless contributing to clinical disease resulting from chromosome anomalies. These rare alterations include isochromosomes, ring chromosomes, dicentric chromosomes, and marker chromosomes (structurally abnormal chromosomes that cannot be identified based on cytogenetics alone). Both translocations and inversions can be completely balanced in some cases, such that there is no disruption of coding regions of the genome, with a completely normal clinical phenotype; however, carriers are at risk for unbalanced forms of these rearrangements in their offspring.

Reciprocal translocations are found in approximately 1/500–1/600 individuals in the general population and result from the exchange of chromosomal segments between at least two chromosomes. These usually occur between nonhomologous chromosomes and can be identified based on an altered banding pattern on G-banding. Balanced translocation carriers are at risk for abnormal chromosome segregation during meiosis and therefore have a higher risk for infertility, SAB, and live-born offspring with multiple congenital malformations. These phenotypes are observed when only one of the pairs of chromosomes involved in a translocation is inherited from a parent, resulting in an unbalanced genotype (Fig. 83e-3). Sometimes the exchanged segments are so small that they cannot be appreciated by

banding (cryptic translocation), and these are sometimes recognized when a phenotypically affected child with an unbalanced form is born. Parental chromosomes can then be studied by FISH to determine if the rearrangement is inherited from a parent with a balanced form of the translocation. The majority of reciprocal, apparently balanced translocations occur in phenotypically normal individuals. The risk for a clinical abnormality when a new reciprocal translocation is identified (usually during prenatal diagnostic studies) is about 7%. Analysis of cytogenetically reciprocal translocations using arrays has demonstrated that translocations in clinically normal individuals are more likely to show no deletions or duplications at the breakpoint, whereas translocations in clinically affected individuals are more likely to have breakpoint-associated deletions or duplications. Most reciprocal translocations occur uniquely, at apparently random positions throughout the genome; however, there are a few exceptions with multiple cases of recurrent translocations occurring. These recurrent translocations include t(11;22), which results in Emanuel syndrome in the unbalanced form, and several translocations involving a region on 4p, 8p, and 12p. These recurrent translocations occur in regions of the genome that contain specific types of AT-rich repeats, or other repeat sequences, that are prone to rearrangement. A special category of translocations is the Robertsonian translocations, which involve the acrocentric chromosomes. An acrocentric chromosome has unique genetic material only on the long arm of the chromosomes, whereas the short arm contains repetitive DNA. The acrocentric chromosomes are 13, 14, 15, 21, and 22. Robertsonian translocations occur when an entire long arm of an acrocentric chromosome is translocated onto the short arm of another acrocentric chromosome. Balanced carriers of a Robertsonian translocation contain only 45 chromosomes, with one chromosome consisting of two long arms of an acrocentric chromosome. Technically, this is an unbalanced translocation, as two short arms of the acrocentric chromosomes are missing; however, because the short arms are repetitive, there is no phenotypic consequence. Unbalanced Robertsonian carriers have 46 chromosomes, but have three copies of the long arm of an acrocentric chromosome. The most

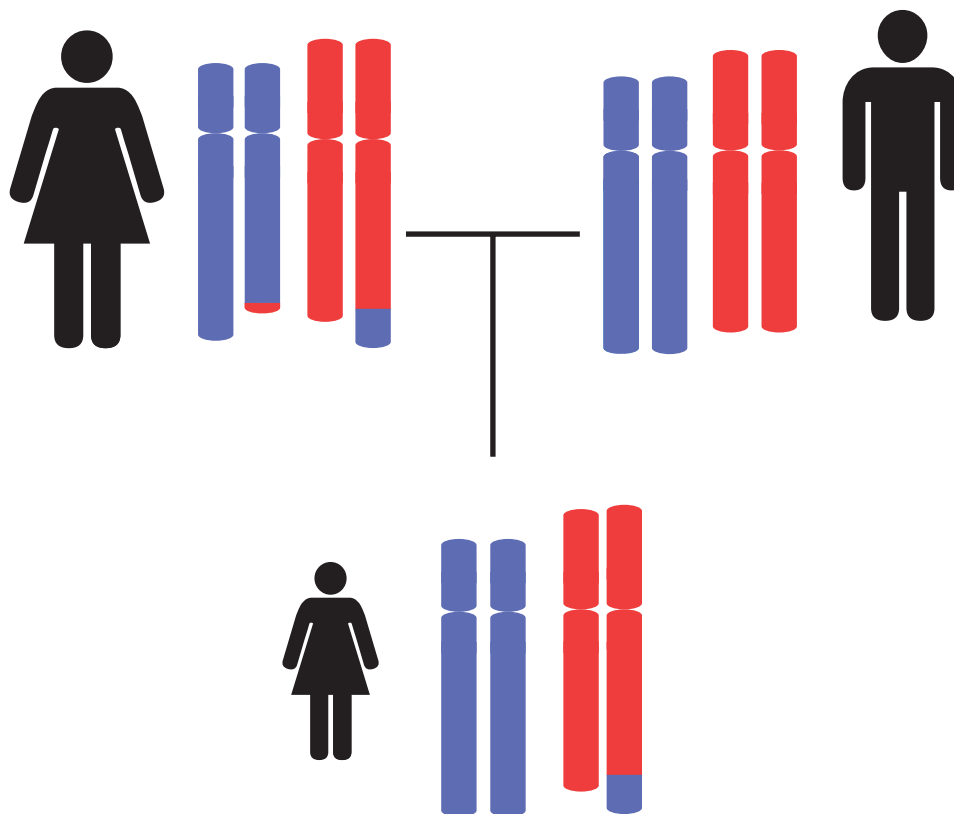


FIGURE 83e-3 Segregation of a balanced translocation in a mother, with inheritance of an unbalanced form in her child. Note that the mother has two rearranged chromosomes, but her child only received one of these, resulting in extra copies of a region of the blue chromosome, with loss of some material from the red chromosome.

common Robertsonian translocation involves chromosomes 13 and 14. Unbalanced Robertsonian translocations involving chromosomes 13 and 21 result in trisomy 13 and Down syndrome, respectively. Approximately 4% of patients with Down syndrome have a translocation, and because recurrence risks are different for families of these individuals, all patients with clinically identified Down syndrome should have a karyotype to look for translocations.

Inversions are another type of chromosome abnormality involving rearranged segments, where there are two breaks within a chromosome, with the intervening chromosomal material inserted in an inverted orientation. As with reciprocal translocations, if a break occurs within a gene or control region for a gene, a clinical phenotype may result, but often there are no consequences for the inversion carrier; however, there is risk for abnormalities in the offspring of carriers, as recombinant chromosomes may result after crossing over between a normal chromosome and an inverted chromosome during meiosis.

Deletion refers to the loss of a chromosomal segment, which results in the presence of only a single copy of that region in an individual's genome. A deletion can be at the end of a chromosome (terminal), or it can be within the chromosome (interstitial). Deletions that are visible at the microscopic level in standard cytogenetic analysis are generally greater than 5 Mb in size. Smaller deletions have been identified by FISH and by chromosomal microarray. The clinical consequences of a deletion depend on the number and function of genes in the deleted region. Genes that cause a phenotype when a single copy is deleted are known as haploinsufficient genes (one copy is not sufficient), and it is estimated that less than 10% of genes are haploinsufficient. Genes associated with disease that are not haploinsufficient include genes for known recessive disorders, such as cystic fibrosis or Tay-Sachs disease.

The first chromosome deletion syndromes were diagnosed clinically and were subsequently demonstrated to be caused by a chromosome deletion on cytogenetic analysis. Examples of these disorders include the Wolf-Hirschhorn syndrome, which is associated with deletions of a small region of the short arm of chromosome 4 (4p); the cri-du-chat syndrome, associated with deletion of a small region of the short arm of chromosome 5 (5p); Williams syndrome, which is associated with interstitial deletions of the long arm of chromosome 7 (7q11.23); and the DiGeorge/velocardiofacial syndromes, associated with interstitial deletions of the long arm of chromosome 22 (22q11.2). Initial cytogenetic studies were able to provide a rough localization of the deletions in different patients, but with the increased usage of arrays, precise mapping of the extent and gene content of these deletions has become much easier. In many cases, one or two genes that are critical for the phenotype associated with these deletions have been identified. In other cases, the phenotype stems from the deletion of multiple genes. The increased utilization of genomic testing by array, which can identify deletions that are much smaller than those detectable by standard cytogenetic analysis, has resulted in the discovery of several new cytogenomic disorders. These include the 1q21.1, 15q13.3, 16p11.2, and 17q21.31 microdeletion syndromes.

Duplication of genomic regions is better tolerated than deletion, as evidenced by the viability of several autosomal trisomies (whole chromosome duplications) but no autosomal monosomies (whole chromosome deletions). There are several duplication syndromes where the duplicated region of the genome is present as a supernumerary chromosome. Utilization of chromosome microarray analysis has made analysis of the origins of duplicated chromosome material straightforward (Fig. 83e-2). Recurrent syndromes associated with supernumerary chromosomes include the inverted duplication 15 (inv dup 15) syndrome, caused by the presence of a marker chromosome derived from chromosome 15, with two copies of proximal 15q resulting in tetrasomy (four copies) of this region. The inv dup 15 syndrome has a distinct phenotype and is associated with hypotonia, developmental delay, intellectual disability, epilepsy, and autistic behavior. Another syndrome is the cat eye syndrome, named for the “cat-eye-like” appearance of the pupil, resulting from a coloboma of the iris. This syndrome results from a supernumerary chromosome derived from a portion of chromosome 22, and the marker chromosomes can vary in size and are often mosaic. Consistent with expectations of a

mosaic disorder, the phenotype of this syndrome is highly variable and includes renal malformations, urinary tract anomalies, congenital heart defects, anal atresia with fistula, imperforate anus, and mild to moderate intellectual disability. Another rare duplication syndrome is the Pallister-Killian syndrome (PKS), which illustrates the principle of tissue-specific mosaicism. Individuals with PKS have coarse facial features with pigmentary skin anomalies, localized alopecia, profound intellectual disability, and seizures. The disorder is caused by a supernumerary isochromosome for the short arm of chromosome 12 (isochromosome 12p). Isochromosomes consist of two copies of one chromosome arm (p or q), rather than one copy of each arm. This isochromosome is not generally seen in peripheral blood lymphocytes when they are analyzed by G-banding, but it is detected in fibroblasts. Array technology has been reported to detect the isochromosome in uncultured peripheral blood in some patients, and it has been hypothesized that a growth bias against cells with the isochromosome prevents their identification in cytogenetic studies.

Numerical abnormalities, translocations, and deletions are the most common chromosome alterations observed in the diagnostic laboratory, but in addition to inversions and duplications, several other types of abnormal chromosomes have been reported, including ring chromosomes, where the two ends of the chromosome fuse to form a circle, and insertions, where a piece of one chromosome is inserted into another chromosome or elsewhere into the same chromosome.

Uniparental disomy (UPD) is the inheritance of a pair of chromosomes (or part of a chromosome) from only one parent. This usually occurs as a result of nondisjunction during meiosis, with a gamete missing or having an extra copy of a chromosome. A resulting fertilized egg would then have only one parental contribution for a given chromosome pair, or a trisomy for a given chromosome. If the monosomy or trisomy is not compatible with life, the embryo may undergo a “rescue” to normal copy number. If a monosomy is rescued, the single chromosome may be duplicated, resulting in a cell with two identical chromosomes (monosomy rescue) (Fig. 83e-4). In the case of trisomies, a subsequent nondisjunction can result in cells where one of the extra chromosomes is lost (trisomy rescue) (Fig. 83e-4). For trisomy rescue, there is a one in three chance that the lost

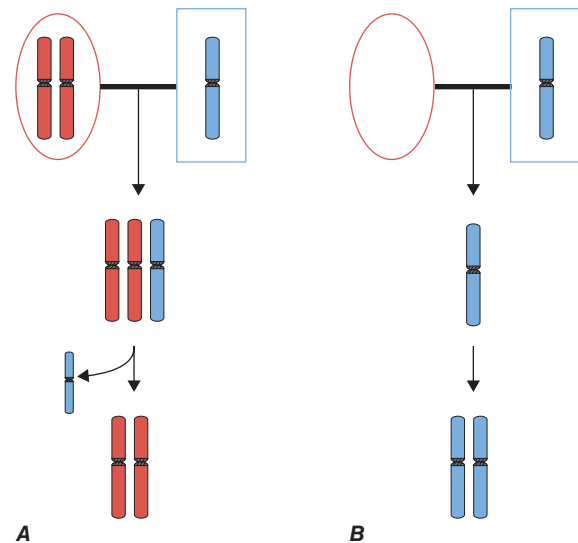


FIGURE 83e-4 Mechanisms of formation of uniparental disomy.

Panel A demonstrates nondisjunction in one parent (mother, represented in red), with trisomy in the zygote. A subsequent nondisjunction, with loss of the paternal chromosome (represented in blue), restores the diploid karyotype but leaves two copies of the maternal chromosome (maternal uniparental disomy [UPD]). **Panel B** demonstrates nondisjunction in one parent (mother, indicated by red oval), resulting in only one copy of this chromosome in the zygote. Subsequent nondisjunction duplicates the single chromosome, rescuing the monosomy, but resulting in two copies of the paternal chromosome (represented in blue; paternal UPD).

chromosome will be the sole chromosome from one parent, resulting in a cell with two chromosomes from the same parent. UPD is sometimes associated with clinical abnormalities, and this can occur by two mechanisms. UPD can cause disease when there is an imprinted gene on the involved chromosome, resulting in altered gene expression. Imprinting is the chemical marking of the parental origin of a chromosome, and genes that are imprinted are only expressed from either the maternal or paternal chromosome ([Chap. 82](#)). Imprinting therefore results in the differential expression of affected genes, based on parent of origin. Imprinting usually occurs through differential modification of the chromosome from one of the parents, and methylation is one of several epigenetic mechanisms (others include histone acetylation, ubiquitylation, and phosphorylation). Imprinted chromosomes that are associated with phenotypes include paternal UPD6 (associated with neonatal diabetes), maternal UPD7 and UPD11 (associated with Russell-Silver syndrome), paternal UPD11 (associated with Beckwith-Wiedemann syndrome), paternal UPD14, maternal UPD15 (Angelman syndrome), and paternal UPD15 (Prader-Willi syndrome). UPD can also result in disease if the two copies from the same parent are the same chromosome (uniparental isodisomy), and the chromosome contains an allele involving a pathogenic mutation associated with a recessive disorder. Two copies of the deleterious allele would result in the associated disease, even though only one parent is a disease carrier.

ACQUIRED CHROMOSOME ABNORMALITIES IN CANCER

Chromosome changes can occur during meiosis or mitosis and can occur at any point across the lifespan. Mosaicism for a developmental disorder is one consequence of mitotic chromosome abnormalities,

and another consequence is cancer, when the chromosome change confers a growth or proliferation advantage on the cell. The types of chromosome abnormalities seen in cancer are similar to those seen in the developmental disorders (e.g., aneuploidy, deletion, duplication, translocation, isochromosomes, rings, inversion). Tumor cells often have multiple chromosome changes, some of which happen early in the development of a tumor, and may contribute to its selective advantage, whereas others are secondary effects of the deregulation that characterizes many tumors. Chromosome changes in cancer have been studied extensively and have been shown to provide important diagnostic, classification, and prognostic information. The identification of cancer type-specific translocation breakpoints has led to the identification of a number of cancer-associated genes. For example, the small abnormal chromosome found to be associated with chronic myelogenous leukemia (CML) in 1960 was shown to be the result of translocation between chromosomes 9 and 22 once techniques for analysis of banded chromosomes were introduced, and subsequently, the translocation breakpoint was cloned to reveal the *c-abl* oncogene on chromosome 9. This translocation produces a fusion protein, which has been targeted for treatment of CML. [For detailed discussion of cancer genetics, see Chap. 101e.](#)

84 The Practice of Genetics in Clinical Medicine

Susan M. Domchek, J. Larry Jameson, Susan Miesfeldt

APPLICATIONS OF MOLECULAR GENETICS IN CLINICAL MEDICINE

Genetic testing for inherited abnormalities associated with disease risk is increasingly used in the practice of clinical medicine. Germline alterations include chromosomal abnormalities (Chap. 83e), specific gene mutations with autosomal dominant or recessive patterns of transmission (Chap. 82), and single nucleotide polymorphisms with small relative risks associated with disease. Germline alterations are responsible for disorders beyond classic Mendelian conditions with genetic susceptibility to common adult-onset diseases such as asthma, hypertension, diabetes mellitus, macular degeneration, and many forms of cancer. For many of these diseases, there is a complex interplay of genes (often multiple) and environmental factors that affect lifetime risk, age of onset, disease severity, and treatment options.

The expansion of knowledge related to genetics is changing our understanding of pathophysiology and influencing our classification of diseases. Awareness of genetic etiology can have an impact on clinical management, including prevention and screening for or treatment of a range of diseases. Primary care physicians are relied upon to help patients navigate testing and treatment options. Consequently, they must understand the genetic basis for a large number of genetically influenced diseases, incorporate personal and family history to determine the risk for a specific mutation, and be positioned to provide counseling. Even if patients are seen by genetic specialists who assess genetic risk and coordinate testing, primary care providers should provide information to their patients regarding the indications, limitations, risks, and benefits of genetic counseling and testing. They must also be prepared to offer risk-based management following genetic risk assessment. Given the pace of genetics, this is an increasingly difficult task. The field of clinical genetics is rapidly moving from single gene testing to multigene panel testing, with techniques such as whole-exome and -genome sequencing on the horizon, increasing the complexity of test selection and interpretation, as well as patient education and medical decision making.

COMMON ADULT-ONSET GENETIC DISORDERS

INHERITANCE PATTERNS

Adult-onset hereditary diseases follow multiple patterns of inheritance. Some are autosomal dominant conditions. These include many common cancer susceptibility syndromes such as hereditary breast and ovarian cancer (due to germline *BRCA1* and *BRCA2* mutations) and Lynch syndrome (caused by germline mutations in the mismatch repair genes *MLH1*, *MSH2*, *MSH6*, and *PMS2*). In both of these examples, inherited mutations are associated with a high penetrance (lifetime risk) of cancer, although risk is not 100%. In other conditions, although there is autosomal dominant transmission, there is lower penetrance, thereby making the disorders more difficult to recognize. For example, germline mutations in *CHEK2* increase the risk of breast cancer, but with a moderate lifetime risk in the range of 20–40%, as opposed to 50–70% for mutations in *BRCA1* or *BRCA2*. Other adult-onset hereditary diseases are transmitted in an autosomal recessive fashion where two mutant alleles are necessary to cause disease. Examples include hemochromatosis and *MYH*-associated colon cancer. There are more pediatric-onset autosomal recessive disorders, such as lysosomal storage diseases and cystic fibrosis.

The genetic risk for many adult-onset disorders is multifactorial. Risk can be conferred by genetic factors at a number of loci, which individually have very small effects (usually with relative risks of <1.5). These risk loci (generally single nucleotide polymorphisms [SNPs]) combine with other genes and environmental factors in ways that are not well understood. SNP panels are available to assess risk of disease,

but the optimal way of using this information in the clinical setting remains uncertain.

Many diseases have multiple patterns of inheritance, adding to the complexity of evaluating patients and families for these conditions. For example, colon cancer can be associated with a single germline mutation in a mismatch repair gene (Lynch syndrome, autosomal dominant), biallelic mutations in *MYH* (autosomal recessive), or multiple SNPs (polygenic). Many more individuals will have SNP risk alleles than germline mutations in high-penetrance genes, but cumulative lifetime risk of colon cancer related to the former is modest, whereas the risk related to the latter is significant. Personal and family histories provide important insights into the possible mode of inheritance.

FAMILY HISTORY

When two or more first-degree relatives are affected with asthma, cardiovascular disease, type 2 diabetes, breast cancer, colon cancer, or melanoma, the relative risk for disease among close relatives ranges from two- to fivefold, underscoring the importance of family history for these prevalent disorders. In most situations, the key to assessing the inherited risk for common adult-onset diseases is the collection and interpretation of a detailed personal and family medical history in conjunction with a directed physical examination.

Family history should be recorded in the form of a pedigree. Pedigrees should convey health-related data on first- and second-degree relatives. When such pedigrees suggest inherited disease, they should be expanded to include additional family members. The determination of risk for an asymptomatic individual will vary depending on the size of the pedigree, the number of unaffected relatives, the types of diagnoses, and the ages of disease onset. For example, a woman with two first-degree relatives with breast cancer is at greater risk for a specific Mendelian disorder if she has a total of 3 female first-degree relatives (with only 1 unaffected) than if she has a total of 10 female first-degree relatives (with 7 unaffected). Factors such as adoption and limited family structure (few women in a family) should be taken into consideration in the interpretation of a pedigree. Additional considerations include young age of disease onset (e.g., a 30-year nonsmoking woman with a myocardial infarction), unusual diseases (e.g., male breast cancer or medullary thyroid cancer), and the finding of multiple potentially related diseases in an individual (e.g., a woman with a history of both colon and endometrial cancer). Some adult-onset diseases are more prevalent in certain ethnic groups. For instance, 2.5% of individuals of Ashkenazi Jewish ancestry carry one of three founder mutations in *BRCA1* and *BRCA2*. Factor V Leiden mutations are much more common in Caucasians than in Africans or Asians.

Additional variables that should be documented are nonhereditary risk factors among those with disease (such as cigarette smoking and myocardial infarction; asbestos exposure and lung disease; and mantle radiation and breast cancer). Significant associated environmental exposures or lifestyle factors decrease the likelihood of a specific genetic disorder. In contrast, the absence of nonhereditary risk factors typically associated with a disease raises concern about a genetic association. A personal or family history of deep vein thrombosis in the absence of known environmental or medical risk factors suggests a hereditary thrombotic disorder. The physical examination may also provide important clues about the risk for a specific inherited disorder. A patient presenting with xanthomas at a young age should prompt consideration of familial hypercholesterolemia. The presence of trichilemmomas in a woman with breast cancer raises concern for Cowden syndrome, associated with *PTEN* mutations.

Recall of family history is often inaccurate. This is especially so when the history is remote and families lose contact or separate geographically. It can be helpful to ask patients to fill out family history forms before or after their visits, because this provides them with an opportunity to contact relatives. Ideally, this information should be embedded in electronic health records and updated intermittently. Attempts should be made to confirm the illnesses reported in the family history before making important and, in certain circumstances, irreversible management decisions. This process is often labor

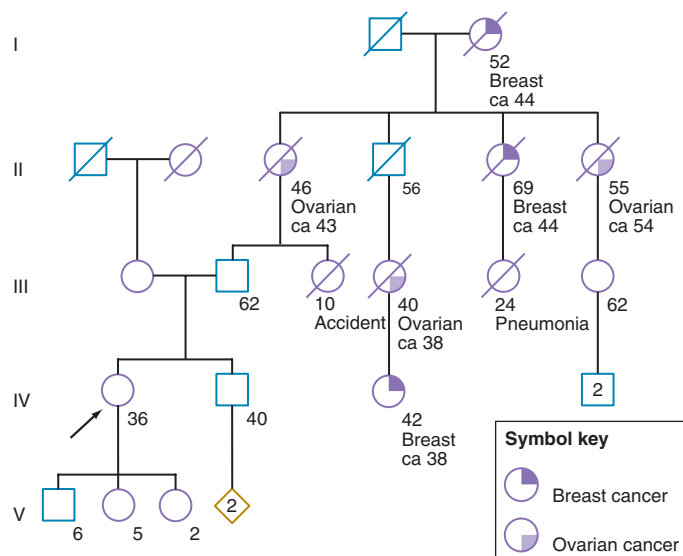


FIGURE 84-1 A 36-year-old woman (arrow) seeks consultation because of her family history of cancer. The patient expresses concern that the multiple cancers in her relatives imply an inherited predisposition to develop cancer. The family history is recorded, and records of the patient's relatives confirm the reported diagnoses.

intensive and ideally involves interviews of additional family members or reviewing medical records, autopsy reports, and death certificates.

Although many inherited disorders will be suggested by the clustering of relatives with the same or related conditions, it is important to note that disease penetrance is incomplete for most genetic disorders. As a result, the pedigree obtained in such families may not exhibit a clear Mendelian inheritance pattern, because not all family members carrying the disease-associated alleles will manifest clinical evidence of the condition. Furthermore, genes associated with some of these disorders often exhibit variable disease expression. For example, the breast cancer-associated gene *BRCA2* can predispose to several different malignancies in the same family, including cancers of the breast, ovary, pancreas, skin, and prostate. For common diseases such as breast cancer, some family members without the susceptibility allele (or genotype) may develop breast cancer (or phenotype) sporadically. Such phenocopies represent another confounding variable in the pedigree analysis.

Some of the aforementioned features of the family history are illustrated in Fig. 84-1. In this example, the proband, a 36-year-old woman (IV-1), has a strong history of breast and ovarian cancer on the paternal side of her family. The early age of onset and the co-occurrence of breast and ovarian cancer in this family suggest the possibility of an inherited mutation in *BRCA1* or *BRCA2*. It is unclear however, without genetic testing, whether her father harbors such a mutation and transmitted it to her. After appropriate genetic counseling of the proband and her family, the most informative and cost-effective approach to DNA analysis in this family is to test the cancer-affected 42-year-old living cousin for the presence of a *BRCA1* or *BRCA2* mutation. If a mutation is found, then it is possible to test for this particular alteration in other family members, if they so desire. In the example shown, if the proband's father has a *BRCA1* mutation, there is a 50:50 probability that the mutation was transmitted to her, and genetic testing can be used to establish the absence or presence of this alteration. In this same example, if a mutation is not detected in the cancer-affected cousin, testing would not be indicated for cancer-unaffected relatives.

GENETIC TESTING FOR ADULT-ONSET DISORDERS

A critical first step before initiating genetic testing is to ensure that the correct clinical diagnosis has been made, whether it is based on family history, characteristic physical findings, pathology, or biochemical testing. Such careful clinical assessment can define the phenotype. In the traditional model of genetic testing, testing is directed initially

toward the most probable genes (determined by the phenotype), which prevents unnecessary testing. Many disorders exhibit the feature of locus heterogeneity, which refers to the fact that mutations in different genes can cause phenotypically similar disorders. For example, osteogenesis imperfecta (Chap. 427), long QT syndrome (Chap. 277), muscular dystrophy (Chap. 462e), and hereditary predisposition to breast (Chap. 108) or colon (Chap. 110) cancer can each be caused by mutations in a number of distinct genes. The patterns of disease transmission, disease risk, clinical course, and treatment may differ significantly depending on the specific gene affected. Historically, the choice of which gene to test has been determined by unique clinical and family history features and the relative prevalence of candidate genetic disorders. However, rapid changes in genetic testing techniques, as discussed below, may impact this paradigm. It is now technically and financially feasible to sequence many genes (or even the whole exome) at one time. The incorporation of multiplex testing for germline mutations is rapidly evolving.

METHODOLOGIC APPROACHES TO GENETIC TESTING

Genetic testing is regulated and performed in much the same way as other specialized laboratory tests. In the United States, genetic testing laboratories are Clinical Laboratory Improvement Amendments (CLIA) approved to ensure that they meet quality and proficiency standards. A useful information source for various genetic tests is www.genetests.org. It should be noted that many tests need to be ordered through specialized laboratories.

Genetic testing is performed largely by DNA sequence analysis for mutations, although genotype can also be deduced through the study of RNA or protein (e.g., apolipoprotein E, hemoglobin S, and immunohistochemistry). For example, universal screening for Lynch syndrome via immunohistochemical analysis of colorectal cancers for absence of expression of mismatch repair proteins is under way at multiple hospitals throughout the United States. The determination of DNA sequence alterations relies heavily on the use of polymerase chain reaction (PCR), which allows rapid amplification and analysis of the gene of interest. In addition, PCR enables genetic testing on minimal amounts of DNA extracted from a wide range of tissue sources including leukocytes, mucosal epithelial cells (obtained via saliva or buccal swabs), and archival tissues. Amplified DNA can be analyzed directly by DNA sequencing, or it can be hybridized to DNA chips or blots to detect the presence of normal and altered DNA sequences. Direct DNA sequencing is frequently used for determination of hereditary disease susceptibility and prenatal diagnosis. Analyses of large alterations of the genome are possible using cytogenetics, fluorescent in situ hybridization (FISH), Southern blotting, or multiplex ligation-dependent probe amplification (MLPA) (Chap. 83e).

Massively parallel sequencing (also called *next-generation sequencing*) is significantly altering the approach to genetic testing for adult-onset hereditary susceptibility disorder. This technology encompasses several high-throughput approaches to DNA sequencing, all of which can reliably sequence many genes at one time. Technically, this involves the use of amplified DNA templates in a flow cell, a very different process than traditional Sanger sequencing which is time-consuming and expensive.

Multiplex panels for inherited susceptibility are commercially available and include testing of a number of genes that have been associated with the condition of interest. For example, panels are available for Brugada syndrome, hypertrophic cardiomyopathy, and Charcot-Marie-Tooth neuropathy. For many syndromes, this type of panel testing may make sense. However, in other situations, the utility of panel testing is less certain. Currently available breast cancer susceptibility panels contain six genes or more. Many of the genes included in the larger panels are associated with only a modest risk of breast cancer, and the clinical application is uncertain. An additional problem of sequencing many genes (rather than the genes for which there is most suspicion) is the identification of one or more variants of uncertain significance (VUS), discussed below.

Whole-exome sequencing (WES) is also now commercially available, although largely used in individuals with syndromes unexplained by

traditional genetic testing. As cost declines, WES may be more widely used. Whole-genome sequencing is also commercially available. Although it may be quite feasible to sequence the entire genome, there are many issues in doing so, including the daunting task of analyzing the vast amount of data generated. Other issues include: (1) the optimal way in which to obtain informed consent, (2) interpretation of frequent sequence variation of uncertain significance, (3) interpretation of alterations in genes with unclear relevance to specific human pathology, and (4) management of unexpected but clinically significant genetic findings.

Testing strategies are evolving as a result of these new genetic testing platforms. As the cost of multiple gene panels and WES continue to fall, and as interpretation of such test results improve, there may be a shift from sequential single-gene (or a few genes) testing to multigene testing. For example, presently, a 30-year-old woman with breast cancer but no family history of cancer and no syndromic features would undergo *BRCA1/2* testing. If negative, she would subsequently be offered *TP53* testing. Notably, a reasonable number of individuals offered *TP53* testing for Li-Fraumeni syndrome decline because mutations are associated with extremely high cancer risks (including childhood cancers) in multiple organs and there are no proven interventions to mitigate risk. Without features consistent with Cowden syndrome, the woman would not be routinely offered *PTEN* testing or testing for *CHEK2*, *ATM*, *BRIP1*, *BARD*, *NBN*, and *PALB2*. However, it is now possible to synchronously analyze all of the aforementioned genes, for a nominally higher cost than *BRCA1/2* testing alone. Concerns about such panels include appropriate consent strategies related to unexpected findings, VUS, and unclear clinical utility of testing moderate-penetrance genes. Thus, changes from the traditional model of single-gene genetic testing should be done with caution (Fig. 84-2).

Limitations to the accuracy and interpretation of genetic testing exist. In addition to technical errors, genetics tests are sometimes designed to detect only the most common mutations. In addition, genetic testing has evolved over time. For example, it was not possible to obtain commercially available comprehensive large genomic rearrangement testing for *BRCA1* and *BRCA2* until 2006. Therefore, a

negative result must be qualified by the possibility that the individual may have a mutation that was not included in the test. In addition, a negative result does not mean that there is not a mutation in some other gene that causes a similar inherited disorder. A negative result, unless there is known mutation in the family, is typically classified as uninformative.

VUS are another limitation to genetic testing. A VUS (also termed *unclassified variant*) is a sequence variation in a gene where the effect of the alteration on the function of the protein is not known. Many of these variants are single nucleotide substitutions (also called missense mutations) that result in a single amino acid change. Although many VUSs will ultimately be reclassified as benign polymorphisms, some will prove to be functionally important. As more genes are sequenced (for example, in a multiplex panel or through WES), the percentage of individuals found to have a VUS increases significantly. The finding of a VUS is difficult for patients and providers alike and complicates decisions regarding medical management.

Clinical utility is an important consideration because genetic testing for susceptibility to chronic diseases is increasingly integrated into the practice of medicine. In some situations, there is clear clinical utility to genetic testing with significant evidence-based changes in medical management decisions based on results. However, in many cases, the discovery of disease-associated genes has outpaced studies that assess how such information should be used in the clinical management of the patient and family. This is particularly true for moderate- and low-penetrance gene mutations. Therefore, predictive genetic testing should be approached with caution and only offered to patients who have been adequately counseled and have provided informed consent.

Predictive genetic testing falls into two distinct categories. Presymptomatic testing applies to diseases where a specific genetic alteration is associated with a near 100% likelihood of developing disease. In contrast, predisposition testing predicts a risk for disease that is less than 100%. For example, presymptomatic testing is available for those at risk for Huntington's disease; whereas, predisposition testing is considered for those at risk for hereditary colon cancer. It is important to note that for the majority of adult-onset disorders, testing is only predictive. Test results cannot reveal with confidence whether,

when, or how the disease will manifest itself. For example, not everyone with the apolipoprotein E4 allele will develop Alzheimer's disease, and individuals without this genetic marker can still develop the disorder.

The optimal testing strategy for a family is to initiate testing in an affected family member first. Identification of a mutation can direct the testing of other at-risk family members (whether symptomatic or not). In the absence of additional familial or environmental risk factors, individuals who test negative for the mutation found in the affected family member can be informed that they are at general population risk for that particular disease. Furthermore, they can be reassured that they are not at risk for passing the mutation on to their children. On the other hand, asymptomatic family members who test positive for the known mutation must be informed that they are at increased risk for disease development and for transmitting the alteration to their children.

Pretest counseling and education are important, as is an assessment of the patient's ability to understand and cope with test results. Genetic testing has implications for entire families, and thus individuals interested in pursuing genetic testing must consider how test results might impact their relationships with relatives, partners, spouses, and children. In families with a known genetic mutation, those who test positive must consider the impact of their carrier

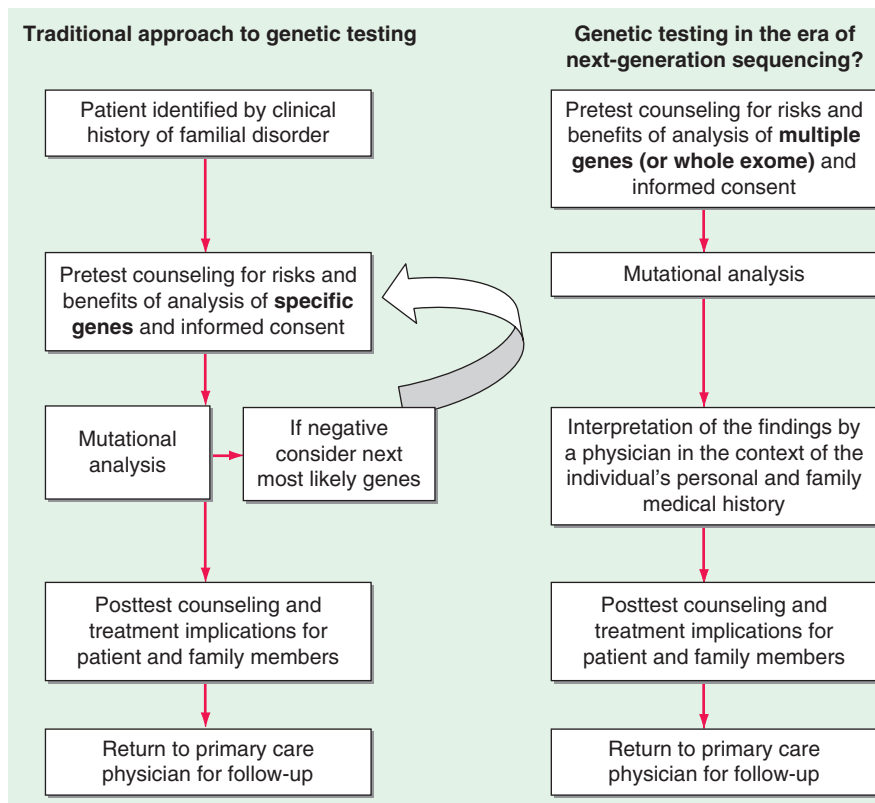


FIGURE 84-2 Approach to genetic testing.

status on their present and future lifestyles; those who test negative may manifest survivor guilt. Parents who are found to have a disease-associated mutation often express considerable anxiety and despair as they address the issue of risk to their children. In addition, some individuals consider options such as preimplantation genetic diagnosis in their reproductive decision making.

When a condition does not manifest until adulthood, clinicians and parents are faced with the question of whether at-risk children should be offered genetic testing and, if so, at what age. Although the matter is debated, several professional organizations have cautioned that genetic testing for adult-onset disorders should not be offered to children. Many of these conditions have no known interventions in childhood to prevent disease; consequently, such information can pose significant psychosocial risk to the child. In addition, there is concern that testing during childhood violates a child's right to make an informed decision regarding testing upon reaching adulthood. On the other hand, testing should be offered in childhood for disorders that may manifest early in life, especially when management options are available. For example, children with multiple endocrine neoplasia 2 (MEN 2) may develop medullary thyroid cancer early in life and should be considered for prophylactic thyroidectomy (Chap. 408). Similarly, children with familial adenomatous polyposis (FAP) due to a mutation in *APC* may develop polyps in their teens with progression to invasive cancer in the twenties, and therefore, colonoscopy screening is started between the ages of 10 and 15 years (Chap. 110).

INFORMED CONSENT

Informed consent for genetic testing begins with education and counseling. The patient should understand the risks, benefits, and limitations of genetic testing, as well as the potential implications of test results. Informed consent should include a written document, drafted clearly and concisely in a language and format that is understandable to the patient. Because molecular genetic testing of an asymptomatic individual often allows prediction of future risk, the patient should understand all potential long-term medical, psychological, and social implications of testing. There have long been concerns about the potential for genetic discrimination. The Genetic Information Nondiscrimination Act (GINA) was passed in 2008 and provides some protections related to job and health insurance discrimination. It is important to explore with patients the potential impact of genetic test results on future health as well as disability and life insurance coverage. Patients should understand that alternatives remain available if they decide not to pursue genetic testing, including the option of delaying testing to a later date. The option of DNA banking should be presented so that samples are readily available for future use by family members, if needed.

FOLLOW-UP CARE AFTER TESTING

Depending on the nature of the genetic disorder, posttest interventions may include: (1) cautious surveillance and awareness; (2) specific medical interventions such as enhanced screening, chemoprevention, or risk-reducing surgery; (3) risk avoidance; and (4) referral to support services. For example, patients with known deleterious mutations in *BRCA1* or *BRCA2* are strongly encouraged to undergo risk-reducing salpingo-oophorectomy and are offered intensive breast cancer screening as well as the option of risk-reducing mastectomy. In addition, such women may wish to take chemoprevention with tamoxifen, raloxifene, or exemestane. Those with more limited medical management and prevention options, such as patients with Huntington's disease, should be offered continued follow-up and supportive services, including physical and occupational therapy and social services or support groups as indicated. Specific interventions will change as research continues to enhance our understanding of the medical management of these genetic conditions and more is learned about the functions of the gene products involved.

Individuals who test negative for a mutation in a disease-associated gene identified in an affected family member must be reminded that they may still be at risk for the disease. This is of particular importance for common diseases such as diabetes mellitus, cancer, and coronary

artery disease. For example, a woman who finds that she does not carry the disease-associated mutation in *BRCA2* previously discovered in the family should be reminded that she still requires the same breast cancer screening recommended for the general population.

GENETIC COUNSELING AND EDUCATION

Genetic counseling should be distinguished from genetic testing and screening, although genetic counselors are often involved in issues related to testing. Genetic counseling refers to a communication process that deals with human problems associated with the occurrence of risk of a genetic disorder in a family. Genetic risk assessment is complex and often involves elements of uncertainty. Counseling, therefore, includes genetic education as well as psychosocial counseling. Genetic counseling can be useful in a wide range of situations (Table 84-1). The role of the genetic counselor includes the following:

1. Gather and document a detailed family history.
2. Educate patients about general genetic principles related to disease risk, both for themselves and for others in the family.
3. Assess and enhance the patient's ability to cope with the genetic information offered.
4. Discuss how nongenetic factors may relate to the ultimate expression of disease.
5. Address medical management issues.
6. Assist in determining the role of genetic testing for the individual and the family.
7. Ensure the patient is aware of the indications, process, risks, benefits, and limitations of the various genetic testing options.
8. Assist the patient, family, and referring physician in the interpretation of the test results.
9. Refer the patient and other at-risk family members for additional medical and support services, if necessary.

Genetic counseling is generally offered in a nondirective manner, wherein patients learn to understand how their values factor into a particular medical decision. Nondirective counseling is particularly appropriate when there are no data demonstrating a clear benefit associated with a particular intervention or when an intervention is considered experimental. For example, nondirective genetic counseling is used when a person is deciding whether to undergo genetic testing for Huntington's disease. At this time, there is no clear benefit (in terms of medical outcome) to an at-risk individual undergoing genetic testing for this disease because its course cannot be altered by therapeutic interventions. However, testing can have an important impact on the individual's perception of advanced care planning and his or her interpersonal relationships and plans for childbearing. Therefore, the decision to pursue testing rests on the individual's belief system and values. On the other hand, a more directive approach is appropriate when a condition can be treated. In a family with FAP, colon cancer screening and prophylactic colectomy should be recommended for known *APC* mutation carriers. The counselor and clinician following this family must ensure that the at-risk family members have access to the resources necessary to adhere to these recommendations.

Genetic education is central to an individual's ability to make an informed decision regarding testing options and treatment. An adequate knowledge of patterns of inheritance will allow patients to understand the probability of disease risk for themselves and other family members. It is also important to impart the concepts of disease penetrance and expression. For most complex adult-onset genetic

TABLE 84-1 INDICATIONS FOR GENETIC COUNSELING

Advanced maternal age (>35 years)
Consanguinity
Previous history of a child with birth defects or a genetic disorder
Personal or family history suggestive of a genetic disorder
High-risk ethnic groups
Documented genetic alteration in a family member
Ultrasound or prenatal testing suggesting a genetic disorder

disorders, asymptomatic patients should be advised that a positive test result does not always translate into future disease development. In addition, the role of nongenetic factors, such as environmental exposures and lifestyle, must be discussed in the context of multifactorial disease risk and disease prevention. Finally, patients should understand the natural history of the disease as well as the potential options for intervention, including screening, prevention, and in certain circumstances, pharmacologic treatment or prophylactic surgery.

Therapeutic Interventions Based on Genetic Risk for Disease

Specific treatments are available for a number of genetic disorders. Strategies for the development of therapeutic interventions have a long history in childhood metabolic diseases; however, these principles have been applied in the diagnosis and management of adult-onset diseases as well (Table 84-2). Hereditary hemochromatosis is usually caused

by mutations in *HFE* (although other genes have been less commonly associated) and manifests as a syndrome of iron overload, which can lead to liver disease, skin pigmentation, diabetes mellitus, arthropathy, impotence in males, and cardiac issues (Chap. 428). When identified early, the disorder can be managed effectively with therapeutic phlebotomy. Therefore, when the diagnosis of hemochromatosis has been made in a proband, it is important to counsel and offer testing to other family members in order to minimize the impact of the disorder.

Preventative measures and therapeutic interventions are not restricted to metabolic disorders. Identification of familial forms of long QT syndrome, associated with ventricular arrhythmias, allows early electrocardiographic testing and the use of prophylactic antiarrhythmic therapy, overdrive pacemakers, or defibrillators. Individuals with familial hypertrophic cardiomyopathy can be screened by ultrasound, treated with beta blockers or other drugs, and counseled about the importance of avoiding strenuous exercise and dehydration. Those with Marfan's syndrome can be treated with beta blockers or

TABLE 84-2 EXAMPLE OF GENETIC TESTING AND POSSIBLE INTERVENTIONS

Genetic Disorder	Inheritance	Genes	Interventions
Oncologic			
Lynch syndrome (HNPCC)	AD	<i>MLH1, MSH2, MSH6, PMS2</i>	Early endoscopic screening; risk-reducing surgery
Familial adenomatous polyposis	AD	<i>APC</i>	Early and frequent endoscopy; prophylactic colectomy
Hereditary breast and ovarian cancer	AD	<i>BRCA1, BRCA2</i>	Risk reducing salpingo-oophorectomy; intensified breast surveillance including breast MRI; risk-reducing mastectomy
Hereditary diffuse gastric cancer	AD	<i>CDH1</i>	Prophylactic gastrectomy; enhanced breast cancer surveillance
Hematologic			
Factor V Leiden	AD	<i>F5</i>	Avoidance of thrombogenic risk factors
Hemophilia A	XL	<i>F8</i>	Factor VIII replacement
Hemophilia B	XL	<i>F9</i>	Factor IX replacement
Glucose 6-phosphate dehydrogenase deficiency	XL	<i>G6PD</i>	Avoidance of oxidant drugs and certain foods
Cardiovascular			
Hypertrophic cardiomyopathy	AD	>10 genes including <i>MYBPC3, MYH7, TNNT2, TPM1</i>	Echocardiographic screening; pharmacologic intervention; myomectomy
Long QT syndrome	AD, AR	>10 genes including <i>KCNQ1, SCN5A, KCNE1, KCNE2</i>	Electrocardiographic screening; pharmacologic intervention; implantable cardiac defibrillator devices
Marfan's syndrome	AD	<i>FBN1</i>	Echocardiographic screening; prophylactic beta blockers; aortic valve replacement as indicated
Gastrointestinal			
Familial Mediterranean fever	AR	<i>MEFV</i>	Colchicine
Hemochromatosis	AR	<i>HFE</i>	Phlebotomy
Pulmonary			
α_1 Antitrypsin deficiency	AR	<i>SERPINA1</i>	Avoidance of smoking and occupational and environmental toxins
Cystic fibrosis	AR	<i>CFTR</i>	Chest physiotherapy; agents to promote airway secretion clearance; <i>CFTR</i> modulators (G551D mutations); lung transplantation
Endocrine			
Neurohypophyseal diabetes insipidus	AD	<i>AVP</i>	Replace vasopressin
Familial hypocalciuric hypercalcemia	AD	<i>CASR</i>	Avoidance of parathyroidectomy
Multiple endocrine neoplasia type 2	AD	<i>RET</i>	Prophylactic thyroidectomy; screening for pheochromocytoma and hyperparathyroidism
Renal			
Polycystic kidney disease	AD, AR	<i>PKD1, PKD2, PKHD1</i>	Prevention of hypertension; prevention of urinary tract infections; kidney transplantation
Nephrogenic diabetes insipidus	XL, AR	<i>AVPR2, AQP2</i>	Fluid replacement; thiazides with or without amiloride
Neurologic			
Malignant hyperthermia	AD	<i>RYR1, CACNA1S</i>	Avoidance of precipitating anesthetics
Hyperkalemic periodic paralysis	AD	<i>SCN4A</i>	Diet rich in calcium and low in potassium; thiazides or acetazolamide
Duchenne's and Becker's muscular dystrophy	XL	<i>DMD</i>	Corticosteroids; physical therapy
Wilson's disease	AR	<i>ATP7B</i>	Zinc, trientine

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; HNPCC, hereditary nonpolyposis colorectal cancer; MRI, magnetic resonance imaging; XL, X-linked.

angiotensin II receptor blockers and monitored for the development of aortic aneurysms.

The field of pharmacogenetics identifies genes that alter drug metabolism or confer susceptibility to toxic drug reactions. Pharmacogenetics seeks to individualize drug therapy in an attempt to improve treatment outcomes and reduce toxicity. Examples include thiopurine methyltransferase (TPMT) deficiency, dihydropyrimidine dehydrogenase deficiency, malignant hyperthermia, and glucose-6-phosphate deficiency. Despite successes in this area, it is not always clear how to incorporate pharmacogenetics into clinical care. For example, although there is an association with *CYP2C6* and *VKORC1* genotypes and warfarin dosing, there is no evidence that incorporating genotyping into clinical practice improves patient outcomes.

The identification of germline abnormalities that increase the risk of specific types of cancer is rapidly changing clinical management. Identifying family members with mutations that predispose to FAP or Lynch syndrome leads to recommendations of early cancer screening and prophylactic surgery, as well as consideration of chemoprevention and attention to healthy lifestyle habits. Similar principles apply to familial forms of melanoma as well as cancers of the breast, ovary, and thyroid. In addition to increased screening and prophylactic surgery, the identification of germline mutations associated with cancer may also lead to the development of targeted therapeutics, for example, the ongoing development of PARP inhibitors in those with *BRCA*-associated cancers.

Although the role of genetic testing in the clinical setting continues to evolve, such testing holds the promise of allowing early and more targeted interventions that can reduce morbidity and mortality. Rapid technologic advances are changing the ways in which genetic testing is performed. As genetic testing becomes less expensive and technically easier to perform, it is anticipated that there will be an expansion of its use. This will present challenges, but also opportunities. It is critical that physicians and other health care professionals keep current with advances in genetic medicine in order to facilitate appropriate referral for genetic counseling and judicious use of genetic testing, as well as to provide state-of-the-art, evidence-based care for affected or at-risk patients and their relatives.

85e Mitochondrial DNA and Heritable Traits and Diseases

Karl Skorecki, Doron Behar

Mitochondria are cytoplasmic organelles whose major function is to generate ATP by the process of oxidative phosphorylation under aerobic conditions. This process is mediated by the respiratory electron transport chain (ETC) multiprotein enzyme complexes I–V and the two electron carriers, coenzyme Q (CoQ) and cytochrome c. Other cellular processes to which mitochondria make a major contribution include apoptosis (programmed cell death) and additional cell type-specific functions (Table 85e-1). The efficiency of the mitochondrial ETC in ATP production is a major determinant of overall body energy balance and thermogenesis. In addition, mitochondria are the predominant source of reactive oxygen species (ROS), whose rate of production also relates to the coupling of ATP production to oxygen consumption. Given the centrality of oxidative phosphorylation to the normal activities of almost all cells, it is not surprising that mitochondrial dysfunction can affect almost any organ system (Fig. 85e-1). Thus, physicians in many disciplines might encounter patients with mitochondrial diseases and should be aware of their existence and characteristics.

The integrated activity of an estimated 1500 gene products is required for normal mitochondrial biogenesis, function, and integrity. Almost all of these are encoded by nuclear genes and thus follow the rules and patterns of nuclear genomic inheritance (Chap. 84). These nuclear-encoded proteins are synthesized in the cell cytoplasm and imported to their location of activity within the mitochondria through a complex biochemical process. In addition, the mitochondria contain their own small genome consisting of numerous copies (polyploidy) per mitochondrion of a circular, double-strand mitochondrial DNA (mtDNA) molecule comprising 16,569 nucleotides. This mtDNA sequence (also known as the “mitogenome”) might represent the remnants of endosymbiotic prokaryotes from which mitochondria are thought to have originated. The mtDNA sequence contains a total of 37 genes, of which 13 encode mitochondrial protein components of the ETC (Fig. 85e-2). The remaining 22 tRNA- and 2 rRNA-encoding genes are dedicated to the process of translating the 13 mtDNA-encoded proteins. This dual nuclear and mitochondrial genetic control of mitochondrial function results in unique and diagnostically challenging patterns of inheritance. The current chapter focuses on heritable traits and diseases related to the mtDNA component of the dual genetic control of mitochondrial function. The reader is referred to Chaps. 84 and 462e for consideration of mitochondrial disease originating from mutations in the nuclear genome. The latter include (1) disorders due to mutations in nuclear genes directly encoding structural components or assembly factors of the oxidative phosphorylation complexes, (2) disorders due to mutations in nuclear genes encoding proteins indirectly related to oxidative phosphorylation, and (3) mtDNA depletion syndromes (MDS) characterized by a reduction

of mtDNA copy number in affected tissues without mutations or rearrangements in the mtDNA.

MITOCHONDRIAL DNA STRUCTURE AND FUNCTION

As a result of its circular structure and extranuclear location, the replication and transcription mechanisms of mtDNA differ from the corresponding mechanisms in the nuclear genome, whose nucleosomal packaging and structure are more complex. Because each cell contains many copies of mtDNA, and because the number of mitochondria can vary during the lifetime of each cell, mtDNA copy number is not directly coordinated with the cell cycle. Thus, vast differences in mtDNA copy number are observed between different cell types and tissues and during the lifetime of a cell. Another important feature of the mtDNA replication process is a reduced stringency of proofreading and replication error correction, leading to a greater degree of sequence variation compared to the nuclear genome. Some of these sequence variants are silent polymorphisms that do not have the potential for a phenotypic or pathogenic effect, whereas others may be considered pathogenic mutations.

With respect to transcription, initiation can occur on both strands and proceeds through the production of an intronless polycistronic precursor RNA, which is then processed to produce the 13 individual mRNA and 24 individual tRNA and rRNA products. The 37 mtDNA genes comprise fully 93% of the 16,569 nucleotides of the mtDNA in what is known as the *coding region*. The *control region* consists of ~1.1 kilobases (kb) of noncoding DNA, which is thought to have an important role in replication and transcription initiation.

MATERNAL INHERITANCE AND LACK OF RECOMBINATION

In contrast to homologous pair recombination that takes place in the nucleus, mtDNA molecules do not undergo recombination, such that mutational events represent the only source of mtDNA genetic diversification. Moreover, with very rare exceptions, it is only the maternal DNA that is transmitted to the offspring. The fertilized oocyte degrades mtDNA carried from the sperm in a complex process involving the ubiquitin proteasome system. Thus, although mothers transmit their mtDNA to both their sons and daughters, only the daughters are able to transmit the inherited mtDNA to future generations. Accordingly, mtDNA sequence variation and associated phenotypic traits and diseases are inherited exclusively along maternal lines.

As noted below, because of the complex relationship between mtDNA mutations and disease expression, sometimes this maternal inheritance is difficult to recognize at the clinical or pedigree level. However, evidence of paternal transmission can almost certainly rule out an mtDNA genetic origin of phenotypic variation or disease; conversely, a disease affecting both sexes without evidence of paternal transmission strongly suggests a heritable mtDNA disorder (Fig. 85e-2).

MULTIPLE COPY NUMBER (POLYPOIDY), HIGH MUTATION RATE, HETEROPLASMY, AND MITOTIC SEGREGATION

Each aerobic cell in the body has multiple mitochondria, often numbering many hundreds or more in cells with extensive energy production requirements. Furthermore, the number of copies of mtDNA within each mitochondrion varies from several to hundreds; this is true of both somatic as well as germ cells, including oocytes in females. In the case of somatic cells, this means that the impact of most newly acquired somatic mutations is likely to be very small in terms of total cellular or organ system function; however, because of the manyfold higher mutation rate during mtDNA replication, numerous different mutations may accumulate with aging of the organism. It has been proposed that the total cumulative burden of acquired somatic mtDNA mutations with age may result in an overall perturbation of mitochondrial function, contributing to age-related reduction in the efficiency of oxidative phosphorylation and increased production of damaging ROS. The accumulation of such acquired somatic mtDNA mutations with aging may contribute to age-related diseases, such as metabolic syndrome and diabetes, cancer, and neurodegenerative and cardiovascular disease in any given individual. However, somatic mutations are not carried forward to the next generation, and the

TABLE 85e-1 FUNCTIONS OF MITOCHONDRIA

All Cells and Tissues

Oxidative phosphorylation
Apoptosis (programmed cell death)

Tissue- or Cell-Specific

Cholesterol metabolism
Amino and organic acid metabolism
Fatty acid beta oxidation
Sex steroid synthesis
Heme synthesis
Hepatic ammonia detoxification
Neurotransmitter metabolism

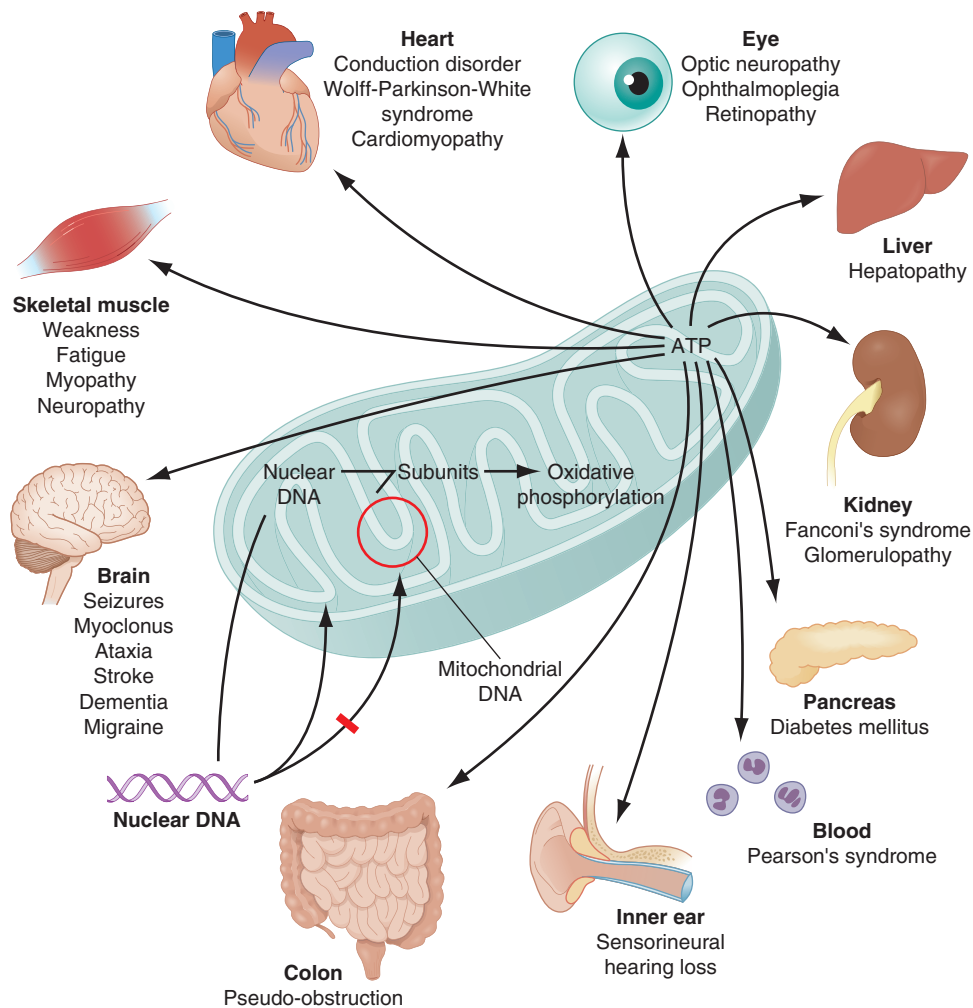


FIGURE 85e-1 Dual genetic control and multiple organ system manifestations of mitochondrial disease. (Reproduced with permission from DR Johns: Mitochondrial DNA and disease. *N Engl J Med* 333:638, 1995.)

hereditary impact of mtDNA mutagenesis requires separate consideration of events in the female germline.

The multiple mtDNA copy number within each cell, including the maternal germ cells, results in the phenomenon of heteroplasmy, in contrast to much greater uniformity (homoplasmy) of somatic nuclear DNA sequence. Heteroplasmy for a given mtDNA sequence variant or mutation arises as a result of the coexistence within a cell, tissue, or individual of mtDNA molecules bearing more than one version of the sequence variant (Fig. 85e-3). The importance of the heteroplasmy phenomena to the understanding of mtDNA-related mitochondrial diseases is critical. The coexistence of mutant and nonmutant mtDNA and the variation of the mutant load among individuals from the same maternal sibship, and across organs and tissues within the same

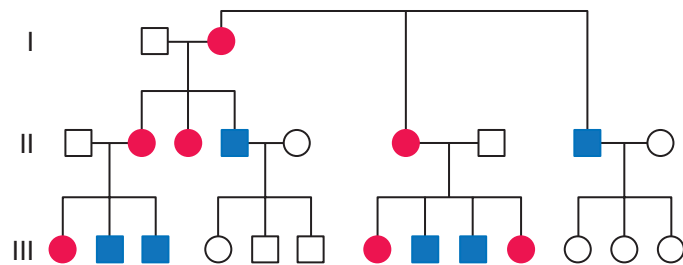


FIGURE 85e-2 Maternal inheritance of mitochondrial DNA (mtDNA) disorders and heritable traits. Affected women (filled circles) transmit the trait to their children. Affected men (filled squares) do not transmit the trait to any of their offspring.

individual, play a pivotal role in the manifestation and severity of disease and are crucial to understanding the complexity of inheritance of mtDNA disorders. At the level of the oocyte, the percentage of mtDNA molecules bearing each version of the polymorphic sequence variant or mutation depends on stochastic events related to partitioning of mtDNA molecules during the process of oogenesis itself. Thus, oocytes differ from each other in the degree of heteroplasmy for that sequence variant or mutation. In turn, the heteroplasmic state is carried forward to the zygote and to the organism as a whole, to varying degrees, depending on mitotic segregation of mtDNA molecules during organ system development and maintenance. For this reason, in vitro fertilization, followed by preimplantation genetic diagnosis (PGD), is not as predictive of the genetic health of the offspring in the case of mtDNA mutations as in the case of the nuclear genome. Similarly, the impact of somatic mtDNA mutations acquired during development and subsequently also shows an enormous spectrum of variability.

Mitotic segregation refers to the unequal distribution of wild-type and mutant versions of mtDNA molecules during all cell divisions that occur during prenatal development and subsequently throughout the lifetime of an individual. The phenotypic effect or disease impact will, thus, be a function not only of the inherent disruptive effect (pathogenicity) on the mtDNA-encoded gene (coding region mutations) or integrity of the mtDNA molecule

(control region mutations), but also of its distribution among the multiple copies of mtDNA in the various mitochondria, cells, and tissues of the affected individual. Thus, one consequence can be the generation of a bottleneck due to the marked decline in given sets of mtDNA variants, consequent to such mitotic segregation. Heterogeneity arises from differences in the degree of heteroplasmy among oocytes of the affected female, together with subsequent mitotic segregation of the pathogenic mutation during tissue and organ development, and throughout the lifetime of the individual offspring. The actual expression of disease might then depend on a threshold percentage of mitochondria whose function is disrupted by mtDNA mutations. This in turn confounds hereditary transmission patterns and hence genetic diagnosis of pathogenic heteroplasmic mutations. Generally, if the proportion of mutant mtDNA is less than 60%, the individual is unlikely to be affected, whereas proportions exceeding 90% cause clinical disease.

HOMOPLASMIC VARIANTS AND HUMAN MTDNA PHYLOGENY

In contrast to classic mtDNA diseases, most of which begin in childhood and are the result of heteroplasmic mutations as noted above, during the course of human evolution, certain mtDNA sequence variants have drifted to a state of homoplasmy, wherein all of the mtDNA molecules in the organism contain the new sequence variant. This arises due to a "bottleneck" effect followed by genetic drift during the very process of oogenesis itself (Fig. 85e-3). In other words, during certain stages of oogenesis, the mtDNA copy number becomes so substantially reduced that the particular mtDNA species bearing the novel or derived sequence variant may become the increasingly predominant,

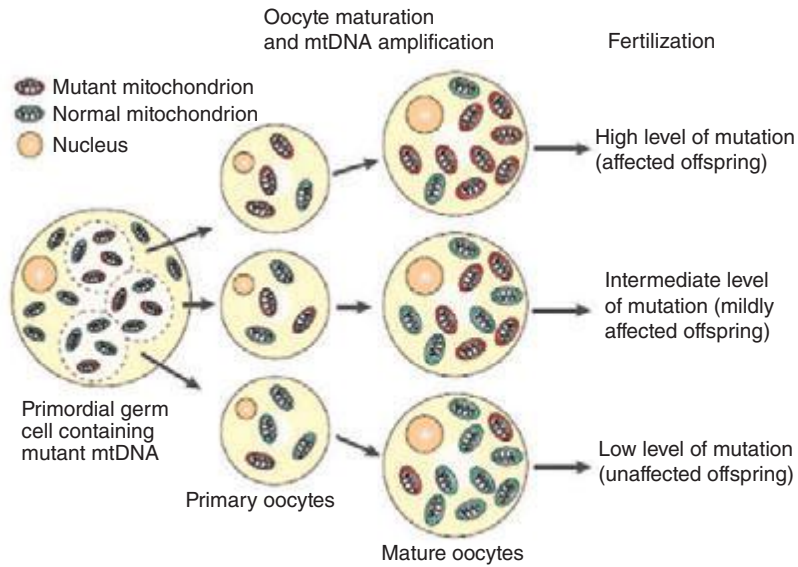


FIGURE 85e-3 Heteroplasmy and the mitochondrial genetic bottleneck.

During the production of primary oocytes, a selected number of mitochondrial DNA (mtDNA) molecules are transferred into each oocyte. Oocyte maturation is associated with the rapid replication of this mtDNA population. This restriction-amplification event can lead to a random shift of mtDNA mutational load between generations and is responsible for the variable levels of mutated mtDNA observed in affected offspring from mothers with pathogenic mtDNA mutations. Mitochondria that contain mutated mtDNA are shown in red, and those with normal mtDNA are shown in green. (Reproduced with permission from R Taylor, D Turnbull: *Mitochondrial DNA mutations in human disease*. *Nat Rev Genetics* 6:389, 2005.)

and eventually exclusive, version of the mtDNA for that particular nucleotide site. All of the offspring of a woman bearing an mtDNA sequence variant or mutation that has become homoplasmic will also be homoplasmic for that variant and will transmit the sequence variant forward in subsequent generations.

Considerations of reproductive fitness limit the evolutionary or population emergence of homoplasmic mutations that are lethal or cause severe disease in infancy or childhood. Thus, with a number of notable exceptions (e.g., mtDNA mutations causing Leber's hereditary optic neuropathy; see below), most homoplasmic mutations are considered to be neutral markers of human evolution, which are useful and interesting in the population genetics analysis of shared maternal ancestry but which have little significance in human phenotypic variation or disease predisposition.

More importantly is the understanding that this accumulation of homoplasmic mutations occurs at a genetic locus that is transmitted only through the female germline and that lacks recombination. In turn, this enables reconstruction of the sequential topology and radiating phylogeny of mutations accumulated through the course of human evolution since the time of the most recent common mtDNA ancestor of all contemporary mtDNA sequences, some 200,000 years ago. The term haplogroup is usually used to define major branching points in the human mtDNA phylogeny, nested one within the other, which often demonstrate striking continental geographic ancestral partitioning. At the level of the complete mtDNA sequence, the term haplotype is usually used to describe the sum of mutations observed for a given mtDNA sequence and as compared to a reference sequence, such that all haplotypes falling within a given haplogroup share the total sum of mutations that have accumulated since the most recent common ancestor and the bifurcation point they mark. The remaining observed variants are private to each haplotype. Consequentially, human mtDNA sequence is an almost perfect molecular prototype for a nonrecombining locus, and its variation has been extensively used in phylogenetic studies. Moreover, the mtDNA mutation rate is considerably higher than the rate observed for the nuclear genome, especially in the control region, which contains the displacement, or

D-loop, in turn comprising two adjacent hypervariable regions (HVR-I and HVR-II). Together with the absence of recombination, this amplifies drift to high frequencies of novel haplotypes. As a result, mtDNA haplotypes are more highly partitioned across geographically defined populations than sequence variants in other parts of the genome. Despite extensive research, it has not been well established that such haplotype-based partitioning has a significant influence on human health conditions. However, mtDNA-based phylogenetic analysis can be used both as a quality assurance tool and as a filter in distinguishing neutral mtDNA variants comprising human mtDNA phylogeny from potentially deleterious mutations.

MITOCHONDRIAL DNA DISEASE

The true prevalence of mtDNA disease is difficult to estimate because of the phenotypic heterogeneity that occurs as a function of heteroplasmy, the challenge of detecting and assessing heteroplasmy in different affected tissues, and the other unique features of mtDNA function and inheritance described above. It is estimated that at least 1 in 200 healthy humans harbors a pathogenic mtDNA mutation with the potential to cause disease, but that heteroplasmic germline pathogenic mtDNA mutations actually affect up to approximately 1 in 8500 individuals.

The true disease burden relating to mtDNA sequence variation will only be known when the following capabilities become available: (1) ability to distinguish a completely neutral sequence variant from a true phenotype-modifying or pathogenic mutation, (2) accurate assessment of heteroplasmy that can be determined with fidelity, and (3) a systems biology approach (Chap. 87e)

to determine the network of epistatic interactions of mtDNA sequence variations with mutations in the nuclear genome.

OVERVIEW OF CLINICAL AND PATHOLOGIC FEATURES OF HUMAN MTDNA DISEASE

Given the vital roles of mitochondria in all nucleated cells, it is not surprising that mtDNA mutations can affect numerous tissues with pleiotropic effects. More than 200 different disease-causing, mostly heteroplasmic mtDNA mutations have been described affecting ETC function. Figure 85e-4 provides a partial mtDNA map of some of the better characterized of these disorders. A number of clinical clues can increase the index of suspicion for a heteroplasmic mtDNA mutation as an etiology of a heritable trait or disease, including (1) familial clustering with absence of paternal transmission; (2) adherence to one of the classic syndromes (see below) or paradigmatic combinations of disease phenotypes involving several organ systems that normally do not fit together within a single nuclear genomic mutation category; (3) a complex of laboratory and pathologic abnormalities that reflect disruption in cellular energetics (e.g., lactic acidosis and neurodegenerative and myodegenerative symptoms with the finding of ragged red fibers, reflecting the accumulation of abnormal mitochondria under the muscle sarcolemmal membrane); or (4) a mosaic pattern reflecting a heteroplasmic state.

Heteroplasmy can sometimes be elegantly demonstrated at the tissue level using histochemical staining for enzymes in the oxidative phosphorylation pathway, with a mosaic pattern indicating heterogeneity of the genotype for the coding region for the mtDNA-encoded enzyme. Complex II, CoQ, and cytochrome c are exclusively encoded by nuclear DNA. In contrast, complexes I, III, IV, and V contain at least some subunits encoded by mtDNA. Just 3 of the 13 subunits of the ETC complex IV enzyme, cytochrome c oxidase, are encoded by mtDNA, and, therefore, this enzyme has the lowest threshold for dysfunction when a threshold level of mutated mtDNA is reached. Histochemical staining for cytochrome c oxidase activity in tissues of patients affected with heteroplasmic inherited mtDNA mutations

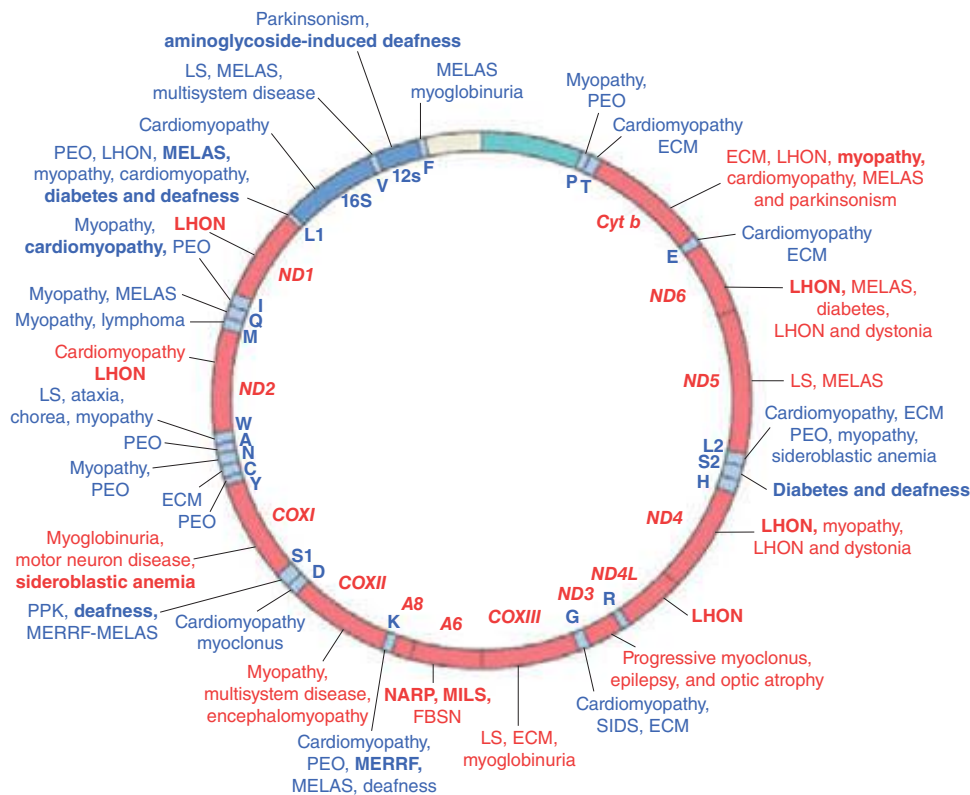


FIGURE 85e-4 Mutations in the human mitochondrial genome known to cause disease.

Disorders that are frequently or prominently associated with mutations in a particular gene are shown in **boldface**. Diseases due to mutations that impair mitochondrial protein synthesis are shown in **blue**. Diseases due to mutations in protein-coding genes are shown in **red**. ECM, encephalomyopathy; FBSN, familial bilateral striatal necrosis; LHON, Leber's hereditary optic neuropathy; LS, Leigh syndrome; MELAS, mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes; MERRF, myoclonic epilepsy with ragged red fibers; MILS, maternally inherited Leigh syndrome; NARP, neuropathy, ataxia, and retinitis pigmentosa; PEO, progressive external ophthalmoplegia; PPK, palmoplantar keratoderma; SIDS, sudden infant death syndrome. (Reproduced with permission from S DiMauro, E Schon: *Mitochondrial respiratory-chain diseases*. *N Engl J Med* 348:2656, 2003.)

(or with the somatic accumulation of mtDNA mutations, see below) can show a mosaic pattern of reduced histochemical staining in comparison with histochemical staining for the complex II enzyme, succinate dehydrogenase (Fig. 85e-5). Heteroplasmy can also be detected at the genetic level through direct Sanger-type mtDNA genotyping under special conditions, although clinically significant low levels of heteroplasmy can escape detection in genomic samples extracted from whole blood using conventional genotyping and sequencing techniques.

The emerging next-generation sequencing (NGS) techniques and their rapid penetration and recognition as useful clinical diagnostic tools are expected to also dramatically improve the clinical genetic diagnostic evaluation of mitochondrial diseases at the level of both the nuclear genome and mtDNA. In the context of the larger nuclear genome, the ability of NGS techniques to dramatically increase the speed at which DNA can be sequenced at a fraction of the cost of conventional Sanger-type sequencing technology is particularly beneficial. Low sequencing costs and short turnaround time expedite "first-tier" screening of panels of hundreds of previously known or suspected mitochondrial disease genes or screening for the entire exome or genome in an attempt to identify novel genes and mutations affecting different patients or families. In the context of the mtDNA, NGS approaches hold the particular promise for rapid and reliable detection of heteroplasmy in different affected tissues. Although Sanger sequencing allows for complete coverage of the mtDNA, it is limited by the lack of deep coverage and low sensitivity for heteroplasmy detection when it is much less than 50%. In contrast, NGS technology

is an excellent tool for rapidly and accurately obtaining a patient's predominant mtDNA sequence and also lower frequency heteroplasmic variants. This is enabled by deep coverage of the genome reads. Accordingly, recent studies making use of NGS techniques have demonstrated sequence accuracy equivalent to Sanger-type sequencing, but also have uncovered heretofore unappreciated heteroplasmy rates ranging between 10 and 50% and detection of single-nucleotide heteroplasmy down to levels of <10%.

Clinically, the most striking overall characteristic of mitochondrial genetic disease is the phenotypic heterogeneity associated with mtDNA mutations. This extends to intrafamilial phenotypic heterogeneity for the same mtDNA pathogenic mutation and, conversely, to the overlap of phenotypic disease manifestations with distinct mutations. Thus, although fairly consistent and well-defined "classic" syndromes have been attributed to specific mutations, frequently "nonclassic" combinations of disease phenotypes ranging from isolated myopathy to extensive multi-system disease are often encountered, rendering genotype-phenotype correlation challenging. In both classical and nonclassical mtDNA disorders, there is often a clustering of some combination of abnormalities affecting the neurologic system (including optic nerve atrophy, pigment retinopathy, and sensorineural hearing loss), cardiac and skeletal muscle (including extraocular muscles), and endocrine and metabolic systems (including diabetes mellitus).

Additional organ systems that may be affected include the hematopoietic, renal, hepatic, and gastrointestinal systems, although these are more frequently involved in infants and children. Disease-causing mtDNA coding region mutations can affect either one of the 13 protein encoding genes or one of the 24 protein synthetic genes. Clinical manifestations do not readily distinguish these two categories, although lactic acidosis and muscle pathologic findings tend to be more prominent in the latter. In all cases, either defective ATP production due to disturbances in the ETC or enhanced generation of ROS has been invoked as the mediating biochemical mechanism between mtDNA mutation and disease manifestation.

mtDNA DISEASE PRESENTATIONS

The clinical presentation of adult patients with mtDNA disease can be divided into three categories: (1) clinical features suggestive of mitochondrial disease (Table 85e-2), but not a well-defined classic syndrome; (2) classic mtDNA syndromes; and (3) clinical presentation confined to one organ system (e.g., isolated sensorineural deafness, cardiomyopathy, or diabetes mellitus).

Table 85e-3 provides a summary of eight illustrative classic mtDNA syndromes or disorders that affect adult patients and highlights some of the most interesting features of mtDNA disease in terms of molecular pathogenesis, inheritance, and clinical presentation. The first five of these syndromes result from heritable point mutations in either protein-encoding or protein synthetic mtDNA genes; the other three result from rearrangements or deletions that usually do not involve the germline.

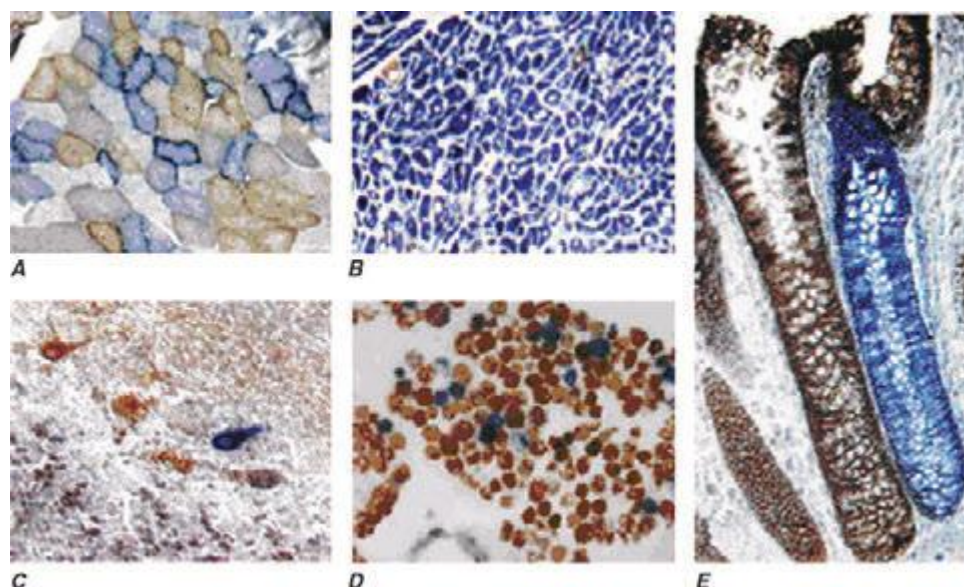


FIGURE 85e-5 Cytochrome c oxidase (COX) deficiency in mitochondrial DNA (mtDNA)-associated disease. Transverse tissue sections that have been stained for COX and succinate dehydrogenase (SDH) activities sequentially, with COX-positive cells shown in *brown* and COX-deficient cells shown in *blue*. **A.** Skeletal muscle from a patient with a heteroplasmic mitochondrial tRNA point mutation. The section shows a typical “mosaic” pattern of COX activity, with many muscle fibers harboring levels of mutated mtDNA that are above the crucial threshold to produce a functional enzyme complex. **B.** Cardiac tissue (left ventricle) from a patient with a homoplasmic tRNA mutation that causes hypertrophic cardiomyopathy, which demonstrates an absence of COX in most cells. **C.** A section of cerebellum from a patient with mtDNA rearrangement that highlights the presence of COX-deficient neurons. **D, E.** Tissues that show COX deficiency due to clonal expansion of somatic mtDNA mutations within single cells—a phenomenon that is seen in both post-mitotic cells (**D**; extraocular muscles) and rapidly dividing cells (**E**; colonic crypt) in aging humans. (Reproduced with permission from R Taylor, D Turnbull: *Mitochondrial DNA mutations in human disease*. *Nat Rev Genetics* 6:389, 2005.)

Leber’s hereditary optic neuropathy (LHON) is a common cause of maternally inherited visual failure. LHON typically presents during young adulthood with subacute painless loss of vision in one eye, with symptoms developing in the other eye 6–12 weeks after the initial onset. In some instances, cerebellar ataxia, peripheral neuropathy, and cardiac conduction defects are observed. In >95% of cases, LHON is due to one of three homoplasmic point mutations of mtDNA that affect genes encoding different subunits of complex I of the mitochondrial ETC; however, not all individuals who inherit a primary LHON mtDNA mutation develop optic neuropathy, and males are four to five times more likely than females to be affected, indicating that additional environmental (e.g., tobacco exposure) or genetic factors are important in the etiology of the disorder. Both the nuclear and mitochondrial genomic backgrounds modify disease penetrance. Indeed, a region of the X chromosome containing a high-risk haplotype for LHON was recently identified, supporting the formulation that nuclear genes act as modifiers and affording an explanation for the male prevalence of LHON. This haplotype can be used in predictive genomic testing and prenatal screening for this disease. In contrast to the other classic mtDNA disorders, it is of interest that patients with this syndrome

are often homoplasmic for the disease-causing mutation. The somewhat later onset in young adulthood and modifying effect of protective background nuclear genomic haplotypes may have enabled homoplasmic pathogenic mutations to have escaped evolutionary censoring.

Mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (MELAS) is a multisystem disorder with a typical onset between 2 to 10 years of age. Following normal early psychomotor development, the most common initial symptoms are seizures, recurrent headaches, anorexia, and recurrent vomiting. Exercise intolerance or proximal limb weakness can be the initial manifestation, followed by generalized tonic-clonic seizures. Short stature is common. Seizures are often associated with stroke-like episodes of transient hemiparesis or cortical blindness that may produce altered consciousness and may recur. The cumulative residual effects of the stroke-like episodes gradually impair motor abilities, vision, and cognition, often by adolescence or young adulthood. Sensorineural hearing loss adds to the progressive decline of these individuals. A plethora of less common symptoms have been described including myoclonus, ataxia, episodic coma, optic atrophy, cardiomyopathy, pigmentary retinopathy, ophthalmoplegia, diabetes mellitus, hirsutism, gastrointestinal dysmotility, and nephropathy. The typical age of death

ranges from 10 to 35 years, but some individuals live into their sixth decade. Intercurrent infections or intestinal obstructions are often the terminal events. Laboratory investigation commonly demonstrates elevated lactate concentrations at rest with excessive increase after moderate exercise. Brain imaging during stroke-like episodes shows areas of increased T2 signal, typically involving the posterior cerebrum and not conforming to the distribution of major arteries. Electrocardiogram (ECG) may show evidence of cardiomyopathy, preexcitation, or incomplete heart block. Electromyography and nerve conduction studies are consistent with a myopathic process, but axonal and sensory neuropathy may coexist. Muscle biopsy typically shows ragged red fibers with the modified Gomori trichrome stain or “ragged blue fibers” resulting from the hyperintense reaction with the histochemical staining for succinate dehydrogenase. The diagnosis of MELAS is based on a combination of clinical findings and molecular genetic testing. Mutations in the mtDNA gene *MT-TL1* encoding tRNA^{leu} are causative. The most common mutation, present in approximately 80% of individuals with typical clinical findings, is an A-to-G transition at nucleotide 3243 (m.3243A>G). Mutations can usually be detected in mtDNA from leukocytes in individuals with typical MELAS; however, the occurrence of heteroplasmy can result in varying tissue distribution of mutated mtDNA. In the absence of specific treatment, various manifestations of MELAS are treated according to standard modalities for prevention, surveillance, and treatment.

Myoclonic epilepsy with ragged red fibers (MERRF) is a multisystem disorder characterized by myoclonus, seizures, ataxia, and myopathy with ragged red fibers. Hearing loss, exercise intolerance, neuropathy, and short stature are often present. Almost all MERRF patients have mutation in the mtDNA tRNA^{lys} gene, and the m.8344A>G mutation in the mtDNA gene encoding the lysine amino acid tRNA is responsible for 80–90% of MERRF cases.

TABLE 85e-2 COMMON FEATURES OF mtDNA-ASSOCIATED DISEASES IN ADULTS

Neurologic: stroke, epilepsy, migraine headache, peripheral neuropathy, cranial neuropathy (optic atrophy, sensorineural deafness, dysphagia, dysphasia)
Skeletal myopathy: ophthalmoplegia, exercise intolerance, myalgia
Cardiac: conduction block, cardiomyopathy
Respiratory: hypoventilation, aspiration pneumonitis
Endocrine: diabetes mellitus, premature ovarian failure, hypothyroidism, hypoparathyroidism
Ophthalmologic: cataracts, pigment retinopathy, neurologic and myopathic (optic atrophy, ophthalmoplegia)

TABLE 85e-3 MITOCHONDRIAL DISEASES DUE TO mtDNA POINT MUTATIONS AND LARGE-SCALE REARRANGEMENTS

Disease	Phenotype	Most Frequent mtDNA Mutations	Homoplasmic (usually)	Maternal
NARP, Leigh disease	Loss of central vision leading to blindness in young adult life	m.1778G>A, m.14484T>C, m.3460G>A	Heteroplasmic	Maternal
MELAS	Mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes; may manifest only as diabetes mellitus	Point mutation in tRNA ^{Leu}	Heteroplasmic	Maternal
MERRF	Myoclonic epilepsy, ragged red fibers in muscle, ataxia, increased CSF protein, sensorineural deafness, dementia	Point mutation in tRNA ^{lys}	Heteroplasmic	Maternal
Deafness	Progressive sensorineural deafness, often induced by aminoglycoside antibiotics	m.1555A>G mutation in 12S rRNA	Homoplasmic	Maternal
	Nonsyndromic sensorineural deafness	m.7445A>G mutation in 12S rRNA	Homoplasmic	Maternal
Chronic progressive external ophthalmoplegia (PEO)	Late-onset bilateral ptosis and ophthalmoplegia, proximal muscle weakness, and exercise intolerance	Single deletions or duplications	Heteroplasmic	Mostly sporadic, somatic mutations
Pearson syndrome	Pancreatic insufficiency, pancytopenia, lactic acidosis	Large deletion	Heteroplasmic	Sporadic, somatic mutations
Kearns-Sayre syndrome (KSS)	External ophthalmoplegia, heart block, retinal pigmentation, ataxia	The 5-kb “common deletion”	Heteroplasmic	Sporadic, somatic mutations

Abbreviations: CSF, cerebrospinal fluid; NARP, neuropathy, ataxia, and retinitis pigmentosa.

Neuropathy, ataxia, and retinitis pigmentosa (NARP) is characterized by moderate diffuse cerebral and cerebellar atrophy and symmetric lesions of the basal ganglia on magnetic resonance imaging (MRI). A heteroplasmic m.8993T>G mutation in the ATPase 6 subunit gene has been identified as causative. Ragged red fibers are not observed in muscle biopsy. When >95% of mtDNA molecules are mutant, a more severe clinical, neuroradiologic, and neuropathologic picture (Leigh syndrome) emerges. Point mutations in the mtDNA gene encoding the 12S rRNA result in heritable nonsyndromic hearing loss. One such mutation causes heritable ototoxic susceptibility to aminoglycoside antibiotics, which opens a pathway for a simple pharmacogenetic test in the appropriate clinical settings.

Kearns-Sayre syndrome (KSS), sporadic progressive external ophthalmoplegia (PEO), and Pearson syndrome are three disease phenotypes caused by large-scale mtDNA rearrangements including partial deletions or partial duplication. The majority of single large-scale rearrangements of mtDNA are thought to result from clonal amplification of a single sporadic mutational event, occurring in the maternal oocyte or during early embryonic development. Because germline involvement is rare, most cases are sporadic rather than inherited. KSS is characterized by the triad of onset before age 20, chronic progressive external ophthalmoplegia, and pigmentary retinopathy. Cerebellar syndrome, heart block, increased cerebrospinal fluid protein content, diabetes mellitus, and short stature are also part of the syndrome. Single deletions/duplication can also result in milder phenotypes such as PEO, characterized by late-onset progressive external ophthalmoplegia, proximal myopathy, and exercise intolerance. In both KSS and PEO, diabetes mellitus and hearing loss are frequent accompaniments. Pearson syndrome is also characterized by diabetes mellitus from pancreatic insufficiency, together with pancytopenia and lactic acidosis, caused by the large-scale sporadic deletion of several mtDNA genes.

Two important dilemmas in classic mtDNA disease have benefited from recent important research insights. The first relates to the greater involvement of neuronal, muscular, renal, hepatic, and pancreatic manifestations in mtDNA disease in these syndromes. This observation has appropriately been mostly attributed to the high energy utilization of the involved tissues and organ systems and, hence, greater dependency on mitochondrial ETC integrity and health. However, because mutations are stochastic events, mitochondrial mutations should occur in any organ during embryogenesis and development. Recently, additional explanations have been suggested based on studies of the common m.3243A>G transition. The proportion of this mutation in peripheral blood cells was shown to decrease exponentially with age.

A selective process acting at the stem cell level with a strong bias against the mutated form would have its greatest effect to reduce the mutant mtDNA only in highly proliferating cells, such as those derived from the hematopoietic system. Tissues and organs with lower cell turnover, such as those involved with mtDNA mutations, would not benefit from this effect and, thus, would be the most affected.

The other dilemma arises from the observation that only a subset of mtDNA mutations accounts for the majority of the familial mtDNA diseases. The random occurrence of mutations in the mtDNA sequence should yield a more uniform distribution of disease-causing mutations. However, recent studies using the introduction of one severe and one mild point mutation into the female germline of experimental animals demonstrated selective elimination during oogenesis of the severe mutation and selective retention of the milder mutation, with the emergence of mitochondrial disease in offspring after multiple generations. Thus, oogenesis itself can act as an “evolutionary” filter for mtDNA disease.

THE INVESTIGATION OF SUSPECTED mtDNA DISEASE

The clinical presentations of classic syndromes, groupings of disease manifestations in multiple organ systems, or unexplained isolated presentations of one of the disease features of a classic mtDNA syndrome should prompt a systematic clinical investigation as outlined in Fig. 85e-6. Indeed, mitochondrial disease should be considered in the differential diagnosis of any progressive multisystem disorder. Despite the centrality of disruptive oxidative phosphorylation, an elevated blood lactate level is neither specific nor sensitive because there are many causes of blood lactic acidosis, and many patients with mtDNA defects presenting in adulthood have normal blood lactate. An elevated cerebrospinal fluid lactate is a more specific test for mitochondrial disease if there is central nervous system involvement. The serum creatine kinase may be elevated but is often normal, even in the presence of a proximal myopathy. Urinary organic and amino acids may also be abnormal, reflecting metabolic and kidney proximal tubule dysfunction. Every patient with seizures or cognitive decline should have an electroencephalogram. A brain computed tomography (CT) scan may show calcified basal ganglia or bilateral hypodense regions with cortical atrophy. MRI is indicated in patients with brainstem signs or stroke-like episodes.

For some mitochondrial diseases, it is possible to obtain an accurate diagnosis with a simple molecular genetic screen. For examples, 95% of patients with LHON harbor one of three mtDNA point mutations (m.11778A>G, m.A3460A>G, or m.14484T>C). These patients have very high levels of mutated mtDNA in peripheral blood cells, and

CLINICAL AND LABORATORY INVESTIGATION OF SUSPECTED mtDNA DISORDER

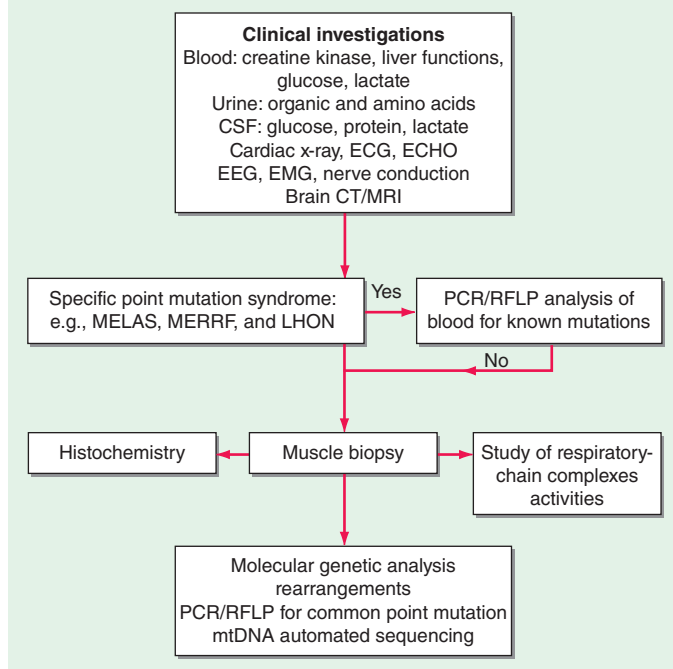


FIGURE 85e-6 Clinical and laboratory investigation of a suspected mitochondrial DNA (mtDNA) disorder. CSF, cerebrospinal fluid; CT, computed tomography; ECG, electrocardiogram; ECHO, echocardiography; EEG, electroencephalogram; EMG, electromyogram; LHON, Leber's hereditary optic neuropathy; MELAS, mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes; MERRF, myoclonic epilepsy with ragged red fibers; MRI, magnetic resonance imaging; PCR, polymerase chain reaction; RFLP, restriction fragment length polymorphism.

therefore, it is appropriate to send a blood sample for molecular genetic analysis by polymerase chain reaction (PCR) or restriction fragment length polymorphism. The same is true for most MERRF patients who harbor a point mutation in the lysine tRNA gene at position 8344. In contrast, patients with the m.3243A>G MELAS mutation often have low levels of mutated mtDNA in blood. If clinical suspicion is strong enough to warrant peripheral blood testing, then patients with a negative result should be investigated further by performing a skeletal muscle biopsy.

Muscle biopsy histochemical analysis is the cornerstone for investigation of patients with suspected mitochondrial disease. Histochemical analysis may show subsarcolemmal accumulation of mitochondria with the appearance of ragged red fibers. Electron microscopy might show abnormal mitochondria with paracrystalline inclusions. Muscle histochemistry may show cytochrome c oxidase (COX)-deficient fibers, which indicate mitochondrial dysfunction (Fig. 85e-5). Respiratory chain complex assays may also show reduced enzyme function. Either of these two abnormalities confirms the presence of a mitochondrial disease, to be followed by an in-depth molecular genetic analysis.

Recent evidence has provided important insights into the importance of nuclear-mtDNA genomic cross-talk and has provided a descriptive framework for classifying and understanding disorders that emanate from perturbations in this cross-talk. Although not strictly considered as mtDNA genetic disorders, manifestations do overlap those highlighted above (Fig. 85e-7).

IMPACT OF HOMOPLASMIC SEQUENCE VARIATION ON HERITABLE TRAITS AND DISEASE

The relationship among the degree of heteroplasmy, tissue distribution of the mutant mtDNA, and disease phenotype simplifies inference of a clear causative relationship between heteroplasmic mutation and disease. With the exception of certain mutations (e.g., those causing most cases of LHON), drift to homoplasmy of such mutations would be precluded normally by the severity of impaired oxidative phosphorylation and the consequent reduction in reproductive fitness. Therefore, sequence variants that have reached homoplasmy should be neutral in terms of human evolution and, hence, useful only for tracing human evolution, demography, and migration, as described above. One important exception is in the case of one or more of the homoplasmic population-level variants, which designate the mtDNA haplogroup J, and the interaction with the mtDNA mutations causing LHON. Reduced disease predilection suggests that one or more of the ancient sequence variants designating mtDNA haplogroup J appears to attenuate predisposition to degenerative disease, in the face of other risk factors. Whether or not additional epistatic interactions between population-level mtDNA haplotypes and common health conditions will be found remains to be determined. If such influences do exist, then they are more likely to be relevant to health conditions in the postreproductive age groups, wherein evolutionary filters would not have had the opportunity to censor deleterious effects and interactions and wherein the effects of oxidative stress may play a role. Although much has been written about the possible associations of population-level common mtDNA variants and human health and disease phenotypes or adaptation to different environmental influences (e.g., climate), a word of caution is in order.

Many studies that purport to show such associations with phenotypes such as longevity, athletic performance, and metabolic and neurodegenerative disease are limited by small sample sizes, possible genotyping inaccuracies, and the possibility of population stratification or ethnic ancestry bias. Because mtDNA haplogroups are so prominently partitioned along phylogeographic lines, it is difficult to rule out the possibility that a haplogroup for which an association has been found is simply a marker for differences in

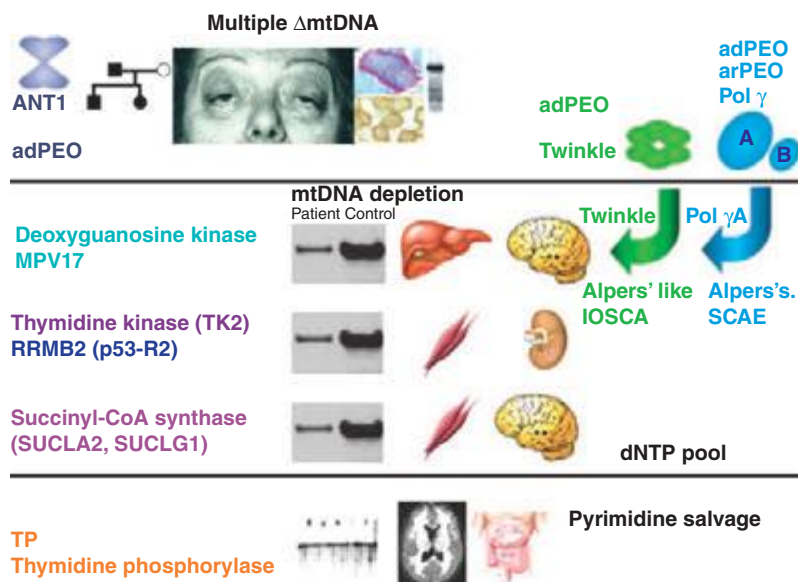


FIGURE 85e-7 Disorders associated with perturbations in nuclear-mitochondrial genomic cross-talk. Clinical features and genes associated with multiple mitochondrial DNA (mtDNA) deletions, mtDNA depletion, and mitochondrial neurogastrointestinal encephalomyopathy syndromes. ANT, adenine nucleotide translocators; adPEO, autosomal dominant progressive external ophthalmoplegia; arPEO, autosomal recessive progressive external ophthalmoplegia; IOSCA, infantile-onset spinocerebellar ataxia; SCAE, spinocerebellar ataxia and epilepsy. (Reproduced with permission from A Spinazzola, M Zeviani: Disorders from perturbations of nuclear-mitochondrial intergenomic cross-talk. *J Intern Med* 265:174, 2009.)

populations with a societal or environmental difference or with different allele frequencies at other genomic loci, which are actually causally related to the heritable trait or disease of interest. The difficulty in generating cellular or animal models to test the functional influence of homoplasmic sequence variants (as a result of mtDNA polyploidy) further compounds the challenge. The most likely formulation is that the risk conferred by different mtDNA haplogroup-defining homoplasmic mutations for common diseases depends on the concomitant nuclear genomic background, together with environmental influences. Progress in minimizing potentially misleading associations in mtDNA heritable trait and disease studies should include ensuring adequate sample size taken from a large sample recruitment base, using carefully matched controls and population structure determination, and performing analysis that takes into account epistatic interactions with other genomic loci and environmental factors.

IMPACT OF ACQUIRED SOMATIC mtDNA MUTATION ON HUMAN HEALTH AND DISEASE

Studies on aging humans and animals have shown a potentially important correlation of age with the accumulation of heterogeneous mtDNA mutations, especially in those organ systems that undergo the most prominent age-related degenerative tissue phenotype. Sequencing of PCR-amplified single mtDNA molecules has demonstrated an average of two to three point mutations per molecule in elderly subjects when compared with younger ones. Point mutations observed include those responsible for known heritable heteroplasmic mtDNA disorders, such as the m.3344A>G and m.3243A>G mutations responsible for the MERRF and MELAS syndromes, respectively. However, the cumulative burden of these acquired somatic point mutations with age was observed to remain well below the threshold expected for phenotypic expression (<2%). Point mutations at other sites not normally involved in inherited mtDNA disorders have also been shown to accumulate to much higher levels in some tissues of elderly individuals, with the description of tissue-specific “hot spots” for mtDNA point mutations. Along the same lines, an age-associated and tissue-specific accumulation of mtDNA deletions has been observed, including deletions involved in known heritable mtDNA disorders, as well as others. The accumulation of functional mtDNA deletions in a given tissue is expected to be associated with mitochondrial dysfunction, as reflected in an age-associated patchy and reduced COX activity on histochemical staining, especially in skeletal and cardiac muscle and brain. A particularly well-studied and potentially important example is the accumulation of mtDNA deletions and COX deficiency observed in neurons of the substantia nigra in Parkinson’s disease patients.

The progressive accumulation of ROS has been proposed as the key factor connecting mtDNA mutations with aging and age-related disease pathogenesis (Fig. 85e-8). As noted above, ROS are a by-product of oxidative phosphorylation and are removed by detoxifying antioxidants into less harmful moieties; however, exaggerated production of ROS or impaired removal results in their accumulation. One of the main targets for ROS-mediated injury is DNA, and mtDNA is particularly vulnerable because of its lack of protective histones and less efficient injury repair systems compared with nuclear DNA. In turn, accumulation of mtDNA mutations results in inefficient oxidative phosphorylation, with the potential for excessive production of ROS, generating a “vicious cycle” of cumulative mtDNA damage. Indeed, measurement of the oxidative stress biomarker 8-hydroxy-2-deoxyguanosine has been used to measure age-dependent increases in mtDNA oxidative damage at a rate exceeding that of nuclear DNA. It should be noted that mtDNA mutation can potentially occur in postmitotic cells as well, because mtDNA replication is not synchronized with the cell cycle. Two other proposed links between mtDNA mutation and aging, besides ROS-mediated tissue injury, are the perturbations in efficiency of oxidative phosphorylation with disturbed cellular aerobic function

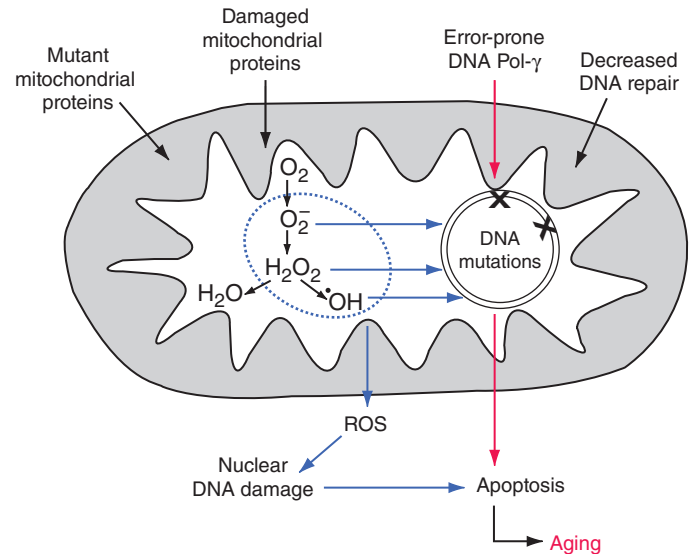


FIGURE 85e-8 Multiple pathways of mitochondrial DNA (mtDNA) damage and aging. Multiple factors may impinge on the integrity of mitochondria that lead to loss of cell function, apoptosis, and aging. The classic pathway is indicated with *blue arrows*; the generation of reactive oxygen species (ROS; superoxide anion, hydrogen peroxide, and hydroxyl radicals), as a by-product of mitochondrial oxidative phosphorylation, results in damage to mitochondrial macromolecules, including the mtDNA, with the latter leading to deleterious mutations. When these factors damage the mitochondrial energy-generating apparatus beyond a functional threshold, proteins are released from the mitochondria that activate the caspase pathway, leading to apoptosis, cell death, and aging. (Reproduced with permission from L Loeb et al: *The mitochondrial theory of aging and its relationship to reactive oxygen species damage and somatic mtDNA mutations. Proc Natl Acad Sci USA 102:18769, 2005.*)

and perturbations in apoptotic pathways, whose execution steps involve mitochondrial activity.

Genetic intervention studies in animal models have sought to clarify the potential causative relationship between acquired somatic mtDNA mutation and the aging phenotype, and the role of ROS in particular. Replication of the mitochondrial genome is mediated by the activity of the nuclear-encoded polymerase gamma gene. A transgenic homozygous mouse knock-in mutation of this gene renders the polymerase enzyme deficient in proofreading and results in a three- to fivefold increase in mtDNA mutation rate. Such mice develop a premature aging phenotype, which includes subcutaneous lipoatrophy, alopecia, kyphonia, and weight loss with premature death. Although the finding of increased mtDNA mutation and mitochondrial dysfunction with age has been solidly established, the causative role and specific contribution of mitochondrial ROS to aging and age-related disease in humans has yet to be proved. Similarly, although many tumors display higher levels of heterogeneous mtDNA mutations, a causal relationship to tumorigenesis has not been proved.

Besides the age-dependent acquired accumulation in somatic cells of heterogeneous point mutations and deletions, a quite different effect of nonheritable and acquired mtDNA mutation has been described affecting tissue stem cells. In particular, disease phenotypes attributed to acquired mtDNA mutation have been observed in sporadic and apparently nonfamilial cases involving a single individual or even tissue, usually skeletal muscle. The presentation consists of decreased exercise tolerance and myalgias, sometimes progressing to rhabdomyolysis. As in the case of the sporadic, heteroplasmic, large-scale deletion, classic syndromes of chronic PEO, Pearson syndrome, and KSS, the absence of a maternal inheritance pattern, together with the finding of limited tissue distribution, suggests a molecular pathogenic mechanism emanating from mutations arising *de novo* in muscle stem cells after germline differentiation (somatic mutations that are

not sporadic and occur in tissue-specific stem cells during fetal development or in the postnatal maintenance or postinjury repair stage). Such mutations would be expected to be propagated only within the progeny of that stem cell and affect a particular tissue within a given individual, without evidence of heritability.

PROSPECTS FOR CLINICAL MANAGEMENT OF mtDNA DISEASE

TREATMENT OF mtDNA DISORDERS

No specific curative treatment for mtDNA disorders is currently available; therefore, the management of mitochondrial disease is largely supportive. Management issues may include early diagnosis and treatment of diabetes mellitus, cardiac pacing, ptosis correction, and intraocular lens replacement for cataracts. Less specific interventions in the case of other disorders involve combined treatment strategies including dietary intervention and removal of toxic metabolites. Cofactors and vitamin supplements are widely used in the treatment of diseases of mitochondrial oxidative phosphorylation, although there is little evidence, apart from anecdotal reports, to support their use. This includes administration of artificial electron acceptors, including vitamin K₃, vitamin C, and ubiquinone (coenzyme Q10); administration of cofactors (coenzymes) including riboflavin, carnitine, and creatine; and use of oxygen radical scavengers, such as vitamin E, copper, selenium, ubiquinone, and idebenone. Drugs that could interfere with mitochondrial function, such as the anesthetic agent propofol, barbiturates, and high doses of valproate, should be avoided. Supplementation with the nitric oxide synthase substrate, L-arginine, has been advocated as a vasodilator treatment during stroke-like episodes. The physician should also be familiar with environmental interactions, such as the strong and consistent association between visual loss in LHON and smoking. A clinical penetrance of 93% was found in men who smoked. Asymptomatic carriers of an LHON mtDNA mutation should, therefore, be strongly advised not to smoke and to moderate their alcohol intake. Although not a cure, these interventions might stave off the devastating clinical manifestations of the LHON mutation. Another example is strict avoidance of aminoglycosides in the familial syndrome of ototoxic susceptibility to aminoglycosides in the presence of the mtDNA m.1555A>G mutation of the 12SrRNA encoding gene.

GENETIC COUNSELING, PRENATAL DIAGNOSIS, AND PREIMPLANTATION GENETIC DIAGNOSIS IN mtDNA DISORDERS

The provision of accurate genetic counseling and reproductive options to families with mtDNA mutations is challenging due to the unique genetic features of mtDNA inheritance that distinguish it from Mendelian genetics. mtDNA defects are transmitted by maternal inheritance. mtDNA de novo mutations are often large deletions, affect one family member, and usually represent no significant risk to other members of the family. In contrast, mtDNA point mutations or duplications can be transmitted down the maternal line. Accordingly, the father of an affected individual has no risk of harboring the disease-causing mutation, and a male cannot transmit the mtDNA mutation to his offspring. In contrast, the mother of an affected individual usually harbors the same mutation but might be completely asymptomatic. This wide phenotypic variability is primarily related to the phenomena of heteroplasmy and the mutation load carried by different members of the same family. Consequently, a symptomatic or asymptomatic female harboring a disease-causing mutation in a heteroplasmic state will transmit to her offspring variable amounts of the mutant mtDNA molecules. The offspring will be symptomatic or asymptomatic primarily according to the mutant load transmitted via the oocyte and, to some extent, subsequent mitotic segregation during development. Interactions with the mtDNA haplotype background or nuclear human genome (as in the case of LHON) serve as an additional important determinant of disease penetrance. Because the severity of the disease phenotype associated with the heteroplasmic mutation load

is a function of the stochastic differential segregation and copy number of mutant mtDNA during the oogenesis bottleneck and, subsequently, following tissue and organ development in the offspring, it is rarely predictable with any degree of accuracy. For this reason, prenatal diagnosis (PND) and PGD techniques that have evolved into integral and well-accepted standards of practice are severely hampered in the case of mtDNA-related diseases.

The value of PND and PGD is limited, partly due to the absence of data on the rules that govern the segregation of wild-type and mutant mtDNA species (heteroplasmy) among tissue in the developing embryo. Three factors are required to ensure the reliability of PND and PGD: (1) a close correlation between the mutant load and the disease severity, (2) a uniform distribution of mutant load among tissues, and (3) no major change in mutant load with time. These criteria are suggested to be fulfilled for the NARP m.8993T>G mutation but do not seem to apply to other mtDNA disorders. In fact, the level of mutant mtDNA in a chorionic villous or amniotic fluid sample may be very different from the level in the fetus, and it would be difficult to deduce whether the mutational load in the prenatal samples provides clinically useful information regarding the postnatal and adult state.

PREVENTION OF MITOCHONDRIAL DISEASE INHERITANCE BY ASSISTED REPRODUCTIVE TECHNOLOGIES

Because the treatment options for patients with mitochondrial disease are rather limited, preventive interventions that eliminate the likelihood of transmission of affected mtDNA into offspring are desirable. The lack of utility of PND and PGD techniques to reliably diagnose and predict mitochondrial disorders at preimplantation-stage products of conception has resulted in the search for alternative preventive approaches for the same problem. One possible approach to “diluting” or even entirely eliminating the mutant mtDNA is applicable only in the earliest embryonic state and in effect represents a form of germline preventive therapy (Fig. 85e-9). This possibility has been explored by using alternative assisted reproduction techniques such as ooplasmic transfer (OT), metaphase chromosome transfer (CT), pronuclear transfer (PNT), and germinal vesicle transfer (GVT) in animal models and, to an extent, in humans. OT is a technique wherein a certain volume (5–15%) of healthy donor oocyte cytoplasm with normal mitochondria is injected into the patient oocyte containing mutated mitochondria. The reasoning behind OT is to supplement the patient’s oocyte with uncompromised cytoplasmic factors such as mtDNA, mRNA, proteins, and other molecules by injecting cytoplasm from healthy oocytes. In PNT, following fertilization, pronuclei of a patient’s zygote are removed with a cytoplasm (“karyoplast”). The karyoplast is transferred to the perivitelline space of a donated zygote, which has been already enucleated. The karyoplast is then fused with enucleated zygote by electric pulses or inactivated Sendai viruses (HVJ). The reconstructed zygote contains a nucleus from the patient (patient nuclear DNA) and cytoplasm from the donor. Thus, the majority of the patient mtDNA is replaced with mtDNA from the donor oocyte. In CT, meiosis II stage of oocyte maturation provides an opportunity for the reconstruction of oocytes with different nuclear and cytoplasmic components before fertilization takes place. Reconstructed oocytes by metaphase chromosome transfer are then fertilized to produce embryos with desired mtDNA haplotypes. In GVT, replacement of compromised cytoplasm with healthy cytoplasm through germinal vesicle transfer before the start of chromosome segregation is carried out.

These approaches have not yet met with widely reported clinical success, yet there is room for optimism. As noted above, analysis of heteroplasmy and inheritance patterns indicates that even a small increase in copies of nonmutant mtDNA can exceed the threshold required to ameliorate serious clinical disease. All of the approaches described above show promise in achieving this goal and thus reducing the burden of clinical mtDNA disease in the future.

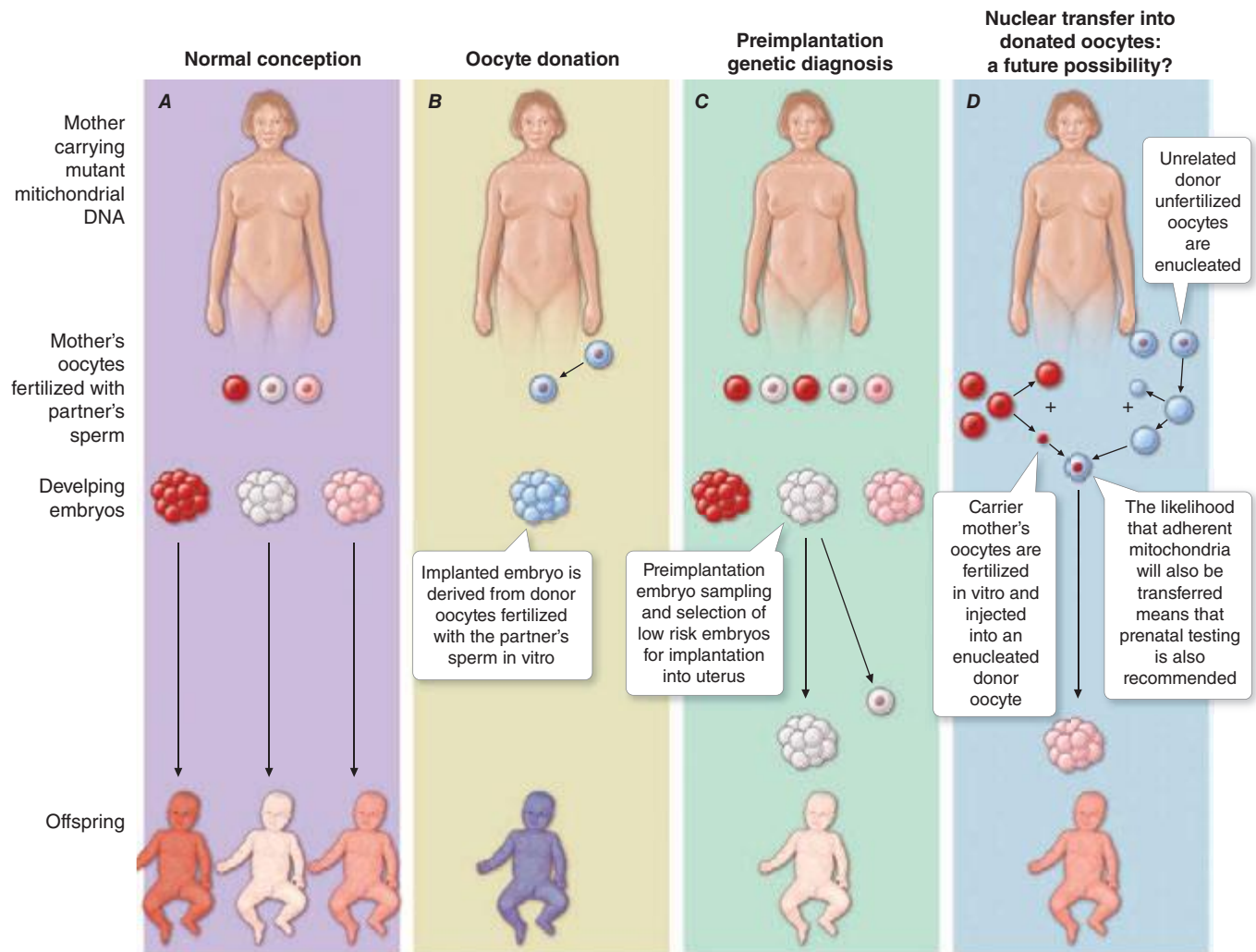


FIGURE 85e-9 Possible approaches for prevention of mitochondrial DNA (mtDNA) disease. **A.** No intervention: offspring's mutant mtDNA load will vary greatly. **B.** Oocyte donation: currently permitted in some constituencies but limited by the availability of oocyte donors. **C.** Preimplantation genetic diagnosis: available for some mtDNA diseases (reliable in determining background nuclear genomic haplotype risk). **D.** Nuclear transfer: research stage, including initial studies in nonhuman primates. Red represents mutant mtDNA, pink and white represent successively higher proportions of normal mtDNA. Blue represents genetic material from an unrelated donor. (Adapted with permission from J Poulton et al: Preventing transmission of maternally inherited mitochondrial DNA diseases. *Br Med J* 338:b94, 2009.)

The technologies that allowed us to decipher the human genome have revolutionized our ability to delineate the composition and functions of the microbial communities that colonize our bodies and make up our *microbiota*. Each body habitat, including the skin, nose, mouth, airways, gastrointestinal tract, and vagina, harbors a distinctive community of microbes. Efforts to understand our microbiota and its collection of microbial genes (our *microbiome*) are changing our views of “self” and deepening our understanding of many normal physiologic, metabolic, and immunologic features and their interpersonal and intrapersonal variations. In addition, this area of research is beginning to provide new insights into diseases not previously known to have microbial “contributors” and is suggesting new strategies for treatment

and prevention. Key terms used in the discussion of the human microbiome are defined in [Table 86e-1](#).

We are holobionts—collections of human and microbial cells that function together in an elaborate symbiosis. The aggregate number of microbial cells in our microbiota exceeds the number of human cells in our adult bodies by up to 10-fold, and each healthy adult is estimated to harbor 10^5 – 10^6 microbial genes, in contrast to ~20,000 *Homo sapiens* genes. Members of our microbiota can function as mutualists (i.e., both host and microbe benefit from each other’s presence), as commensals (one partner benefits; the other is seemingly unaffected), and as potential or overt pathogens (one partner benefits; the other is harmed). Many clinicians view pathogens as individual microbial species or strains that can elicit disease in susceptible hosts. An emerging, more ecologic view is that pathogens do not function in isolation; rather, their invasion, emergence, and effects on the host reflect interactions with other members of a microbiota. An even more expansive view is that multiple organisms in a community conspire to

TABLE 86e-1 GLOSSARY OF TERMS USED IN DISCUSSION OF THE HUMAN MICROBIOME

Term	Definition
Culture-independent analysis	A type of analysis in which the culture of microbes is not required but rather information is extracted directly from environmental samples
Diversity (alpha and beta)	<i>Alpha</i> diversity measures the effective number of species (kinds of organisms) at the level of individual habitats, sites, or samples. <i>Beta</i> diversity measures differences in the number of kinds of organisms across habitats, sites, or samples.
Domains of life	The three major branches of life on Earth: the Eukarya (including humans), the Bacteria, and the Archaea
Dysbiosis	Any deleterious condition arising from a structural and/or functional aberration in one or more of the host organism’s microbial communities
Gnotobiotics	The rearing of animals under sterile (germ-free) conditions. These animals can subsequently be colonized at various stages of the life cycle with defined collections of microbes.
Holobiont	The biologic entity consisting of a host and all its internal and external symbionts, their gene repertoires, and their functions
Human microbiome	In ecology, <i>biome</i> refers to a habitat and the organisms in it. In this sense, the human <i>microbiome</i> would be defined as the collection of microorganisms associated with the human body. However, the term <i>microbiome</i> is also used to refer to the collective genomes and genes present in members of a given microbiota (see “Microbiota,” below), and the human <i>metagenome</i> is the sum of the human genome and microbial genes (microbiome). A <i>core human microbiome</i> is defined as everything shared in a given body habitat among all or the vast majority of human microbiomes. A core microbiome may include a common set of genomes and genes encoding various protein families and/or metabolic capabilities. Microbial genes that are variably represented in different humans may contribute to distinctive physiologic/metabolic phenotypes.
Metagenomics	An emerging field encompassing culture-independent studies of the structures and functions of microbial communities as well as the interactions of these communities with the habitats they occupy. Metagenomics includes (1) shotgun sequencing of microbial DNA isolated directly from a given environment and (2) high-throughput screening of expression libraries constructed from cloned community DNA to identify specific functions such as antibiotic resistance (<i>functional metagenomics</i>). DNA-level analyses provide the foundation for profiling of mRNAs and proteins produced by a microbiome (<i>metatranscriptomics</i> and <i>metaproteomics</i>) and for identification of a community’s metabolic network (<i>metametabolomics</i>).
Microbial source tracking	A collection of methods for assessing the environments of origin for microbes. One method, SourceTracker, uses a Bayesian approach to identify each bacterial taxon’s origins and estimates the proportions of each community made up by bacteria originating from different environments.
Microbiota	A microbial community—including Bacteria, Archaea, Eukarya, and viruses—that occupies a given habitat
Pan-genome	The group of genes found in genomes that make up a given microbial phylotype, including both <i>core</i> genes found in all genomes and variably represented genes found in a subset of genomes within the phylotype
Phylogenetic analysis	Characterization of the evolutionary relationships between organisms and their gene products
Phylogenetic tree	A “tree” in which organisms are shown according to their relationships to hypothetical common ancestors. When built from molecular sequences, the branch lengths are proportional to the amount of evolutionary change separating each ancestor–descendant pair.
Phylotype	A phylogenetic group of microbes, currently defined by a threshold percentage identity shared among their small-subunit rRNA genes (e.g., $\geq 97\%$ for a species-level phylotype)
Principal coordinates analysis	An ordination method for visualizing multivariate data based on the similarity/dissimilarity of the measured entities (e.g., visualization of bacterial communities based on their UniFrac distances; see “UniFrac,” below)
Random Forests analysis/machine learning	<i>Machine learning</i> is a collection of approaches that allow a computer to learn without being explicitly programmed. <i>Random Forests</i> is a machine-learning method for classification and regression that uses multiple decision trees during a training step.
Rarefaction	A procedure in which subsampling is used to assess whether all the diversity present in a given sample or set of samples has been observed at a given sampling depth and to extrapolate how much additional sampling would be needed to observe all the diversity
Resilience	A community’s ability to return to its initial state after a perturbation
Shotgun sequencing	A method for sequencing large DNA regions or collections of regions by fragmenting DNA and sequencing the resulting smaller sections
Succession (primary and secondary)	<i>Succession</i> (in an ecologic context) refers to changes in the structure of a community through time. <i>Primary</i> succession describes the sequence of colonizations and extinctions that occur in a new habitat. <i>Secondary</i> succession refers to changes in community structure after a disturbance.
UniFrac	A measure of the phylogenetic dissimilarity between two communities, calculated as the unshared proportion of the phylogenetic tree containing all the organisms present in either community

produce pathogenic effects in certain host and environmental contexts (a pathologic community).

The ability to characterize microbial communities without culturing their component members has spawned the field of *metagenomics* (Table 86e-1). Metagenomics reflects a confluence of experimental and computational advances in the genome sciences as well as a more ecologic understanding of medical microbiology, according to which the functions of a given microbe and its impact on human biology depend on the context of other microbes in the same community. Traditional microbiology relies on culturing individual microbes, but metagenomics skips this step, instead sequencing DNA isolated directly from a given microbial community. The resulting datasets facilitate follow-up functional studies, such as the profiling of RNA and protein products expressed from the microbiome or the characterization of a microbial community's metabolic activities.

Metagenomics provides insight into how microbial communities vary in several situations critical to human health. One such situation is how microbial communities are assembled following birth and how they operate over time, including responses of established communities to various perturbations. Another is how microbial communities normally vary between different anatomic sites within an individual and between different groups of people representing different ages, physiologic states, lifestyles, geographies, and gender. Yet another is how microbial communities vary in disease; whether such variations are consistent among individuals grouped according to current criteria for a disease or its subtypes; whether the microbiota or microbiome provides new ways of classifying disease states; and, importantly, whether the structural and functional configurations of microbial communities are a cause or a consequence of disease.

Analysis of our microbiomes also addresses one of the most fundamental questions in genetics: How does environment select our genes and directly influence their function? Each human encounters a unique environment during the course of his or her lifetime. Part of this personally experienced environment is incorporated into the genes and capabilities of our microbial communities. The microbiome therefore expands our conceptualization of “human” genetic potential from a single set of genes “fixed” at birth to a microbiome with additional genes and capabilities acquired via a process influenced by our family and life experiences, including modifiable lifestyle choices such as diet. This view recognizes a previously underappreciated dimension of human evolution that occurs at the level of our microbiomes and inspires us to determine how—and how fast—this microbial evolution effects changes in our human biology. For example, Westernization is associated with loss of bacterial species diversity (richness) in the microbiota, and this loss may be associated with the suite of Western diseases. The study of our microbiomes also raises important questions about personal identity, how we define the origins of health disparities, and privacy. Further, it offers the possibility of entirely new approaches to disease prevention and treatment, including *regenerative* medicine, which involves administration of microbial species (*probiotics*) to individuals harboring communities that have not developed into a mature, fully functional state or that have been perturbed in ways that can be restored by the addition of species that fill unoccupied “jobs” (niches).

This chapter provides a general overview of how human microbial communities are analyzed; reviews ecologic principles that guide our understanding of microbial communities in health and disease; summarizes recent studies that establish correlations and, in some cases, causal relationships between our microbiota/microbiomes and various diseases; and discusses challenges faced in the translation of these findings to new therapeutic interventions.

A TOOLBOX FOR METAGENOMIC ANALYSES OF HUMAN MICROBIAL COMMUNITIES

Life on Earth has been classified into three domains: Bacteria, Archaea, and Eukarya. The habitats of the surface-exposed human body harbor members of each domain plus their viruses. In large part, microbial diversity has not been characterized by culture-based approaches, partly because we do not know how to re-create the metabolic milieu fashioned by these communities in their native habitats and

partly because a few organisms tend to outgrow the others. Culture-independent methods readily identify which organisms are present in a microbiota and their relative abundance. The gene widely used to identify microbes and their evolutionary relationships encodes the major RNA component of the small subunit (SSU) of ribosomes. Within each domain of life, the SSU gene is highly conserved, allowing the SSU gene sequences present in different organisms in that domain to be accurately aligned and regions of nucleotide sequence variation to be identified. Pairwise comparisons of SSU ribosomal RNA (rRNA) genes from different microbes allow construction of a phylogenetic tree that represents an evolutionary map on which previously unknown organisms can be assigned a position. This approach, known as *molecular phylogenetics*, permits characterization of each organism on the basis of its evolutionary distance from other organisms. Different phylogenetic types (*phylotypes*) can be viewed as comprising branches on an evolutionary tree.

Characterization of Bacteria Because members of the Bacteria dominate our microbiota, most studies defining our various body habitat-associated microbial communities have sequenced the bacterial SSU gene that encodes 16S rRNA. This gene has a mosaic structure, with highly conserved domains flanking more variable regions. The most straightforward way to identify bacterial taxonomic groups (*taxa*) in a given community is to sequence polymerase chain reaction (PCR) products (*amplicons*) generated from the 16S rRNA genes present in that community. PCR primers directed at the conserved regions of the gene yield PCR amplicons encompassing one or more of that gene's nine variable regions. PCR primer design is critical: differential annealing with primer pairs designed to amplify different variable regions can lead to over- or underrepresentation of specific taxa, and different regions within the 16S rRNA gene can have different patterns of evolution. Therefore, caution must be exercised in comparisons of the relative abundance of taxa in samples characterized in different studies, as methodologic differences can lead to larger perceived differences in the inferred taxonomy than actually exist.

A key innovation is *multiplex sequencing*. Amplicons from each microbial-community DNA sample are tagged by incorporation of a unique oligonucleotide barcode into the PCR primer. Amplicons harboring these sample-specific barcodes can then be pooled together so that multiple samples representing multiple communities can be sequenced simultaneously (Fig. 86e-1). One important choice is the tradeoff between the number of samples that can be processed simultaneously and the number of sequences generated per sample. Interpersonal differences in the bacterial components of the microbiota are typically large, as are differences between communities occupying different body habitats in the same individual (see below); thus fewer than 1000 16S rRNA reads are characteristically required to discriminate community type. However, the identification of systematic differences in microbiota composition that correlate with physiologic status or disease state is confounded by the substantial interpersonal variation that occurs normally.

Sequencing of bacterial 16S rRNA genes creates a challenge for medical microbiology: how to define the taxonomic groups present in a community in a systematic and informative manner, so that one community can be compared with and contrasted to another. Within each domain of life, microbes are classified in a hierarchy beginning with phylum (the broadest group) followed by class, order, family, genus, and species. To determine taxonomy, 16S rRNA sequences are aligned on the basis of their sequence similarity—a process known as *picking operational taxonomic units* (OTUs). Grouping of 16S rRNA sequences from a given variable region into “bins” that share $\geq 97\%$ nucleotide sequence identity (97%ID OTUs) is a commonly accepted, albeit arbitrary, way to define a species.

Looking beyond the 16S rRNA gene, we find that different isolates (strains) of a given bacterial species have overlapping but not identical sets of genes in their genomes. The aggregate set of genes identified in all isolates (strains) of a given species-level phylotype represents its *pan-genome*. Most species are represented by multiple strains, sometimes with markedly different functions (for example,

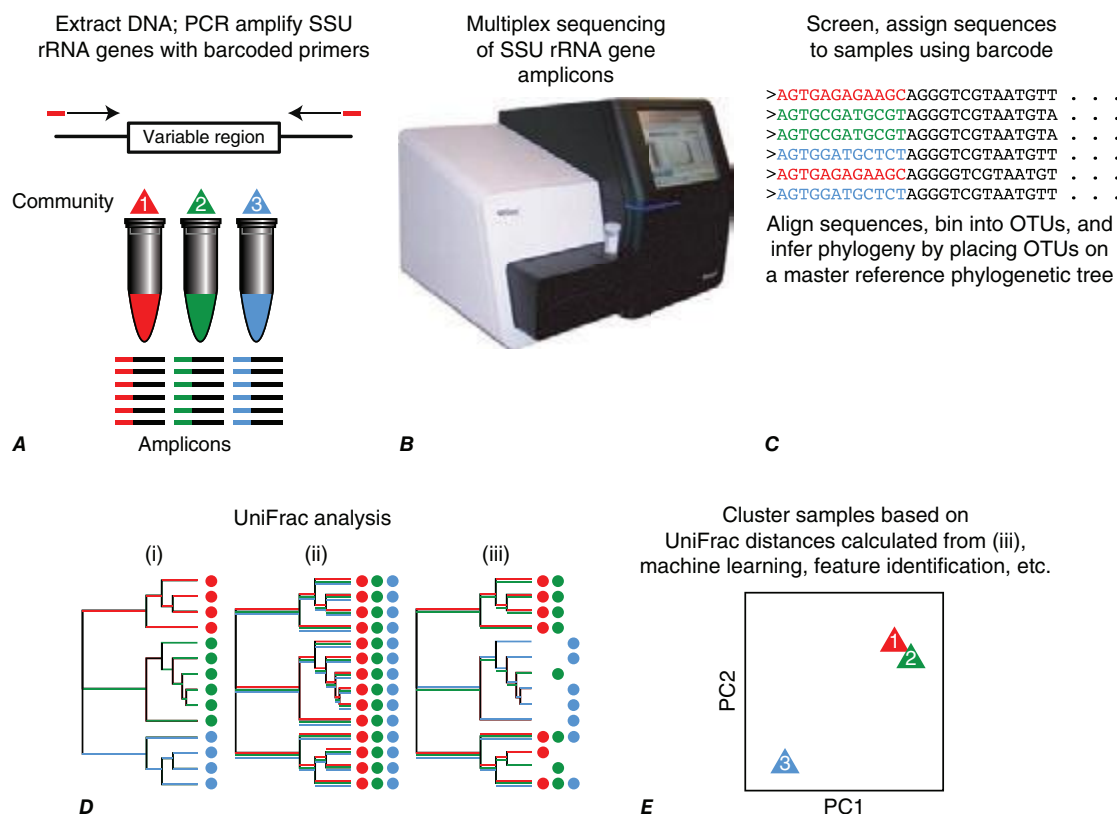


FIGURE 86e-1 Pipeline for culture-independent studies of a microbiota. **(A)** DNA is extracted directly from a sampled human body habitat-associated microbial community. The precise location of the community and relevant patient clinical data are collected. Polymerase chain reaction (PCR) is used to amplify portions of small-subunit (SSU) rRNA genes (e.g., the genes encoding bacterial 16S rRNA) containing one or more variable regions. Primers with sample-specific, error-correcting barcodes are designed to recognize the more conserved regions of the 16S rRNA gene that flank the targeted variable region(s). **(B)** Barcoded amplicons from multiple samples (communities 1–3) are pooled and sequenced in batch in a highly parallel next-generation DNA sequencer. **(C)** The resulting reads are then processed, with barcodes denoting which sample the sequence came from. After barcode sequences are removed in silico, reads are aligned and grouped according to a specified level of shared identity; e.g., sequences that share $\geq 97\%$ nucleotide sequence identity are regarded as representing a species. Once reads are binned into operational taxonomic units (OTUs) in this fashion, they are placed on a phylogenetic tree of all known bacteria and their phylogeny is inferred. **(D)** Communities can be compared to one another by either taxon-based methods, in which phylogeny is not considered and the number of shared taxa are simply scored, or phylogenetic methods, in which community similarity is considered in light of the evolutionary relationships of community members. The UniFrac metric is commonly used for phylogeny-based comparisons. In stylized examples (i), (ii), and (iii), communities with varying degrees of similarity are shown. Each circle represents an OTU colored on the basis of its community of origin and placed on a master phylogenetic tree that includes all lineages from all communities. Branches (horizontal lines) are colored with each community that contains members from that branch. The three examples vary in the amount of branch length shared between the OTUs from each community. In (i), there is no shared branch length, and thus the three communities have a similarity score of 0. In (ii), the communities are identical, and a similarity score of 1 is assigned. In (iii), there is an intermediate level of similarity: communities represented in red and green share more branch length and thus have a higher similarity score than red vs. blue or green vs. blue. The amount of shared branch length in each pairwise community comparison provides a distance matrix. **(E)** The results of taxon- or phylogeny-based distance matrices can be displayed by principal coordinates analysis (PCoA), which plots each community spatially such that the largest component of variance is captured on the x-axis (PC1) and the second largest component of variance is displayed on the y-axis (PC2). In the example shown, the three communities in example (iii) from panel D are compared. Note that for shotgun sequencing of whole-community DNA (microbiome analysis), reads are compared with genes that are present in the genomes of sequenced cultured microbes and/or with genes that have been annotated by hierarchical functional classification schemes in various databases, such as the Kyoto Encyclopedia of Genes and Genomes (KEGG). Communities can then be compared on the basis of the distribution of functional groups in their microbiomes—an approach analogous to taxon-based methods for 16S rRNA-based comparisons—and the results plotted with PCoA.

enteropathogenic versus commensal *Escherichia coli*). Differences in genome content among strains of a given species reflect differences in community membership as well as differences in the selective pressures these strains experience within and between habitats. Horizontal gene transfer among members of a microbiota—mediated by phage, plasmids, and other mechanisms—is a major contributor to this strain-level variation.

Strain-level diversity can be important in any consideration of how microbial communities differ between individuals and how these communities accommodate perturbations. For example, the great bacterial strain-level diversity that exists in the gut is thought to be one of the features that allows this microbiota, which occupies a constantly

perfused ecosystem exposed to the complex and varying set of substances we ingest, to adapt to changing circumstances rather than depending on one strain to occupy a given niche important for proper community functioning. In ecologic studies of different environments, such as grasslands, forests, and reefs, increased diversity within a community increases its capacity to respond to disturbances and to restore itself (i.e., its *resilience*); the same is likely true of microbial ecosystems. When characterizing the mechanisms by which a given species produces an effect or effects on humans, it is important to consider the strain being tested; strain-level diversity has an impact on discovery and development efforts aimed at identifying next-generation probiotics that can be used therapeutically to promote health or treat disease.

Identification of Archaeal and Eukaryotic Members Surveys based on SSU rRNA gene sequencing have largely focused on Bacteria, yet the census of “who’s there” in human body habitat-associated communities must also include the other two domains of life: Archaea and Eukarya. Differences in the sequences of archaeal and bacterial 16S rRNA genes, first recognized by Carl Woese in 1977, allowed these two domains of life to be distinguished. The representation of Archaea in human microbial communities is less well defined than that of Bacteria, in part due to the difficulty in optimizing the design of PCR primers that specifically target conserved regions of archaeal (versus bacterial) 16S rRNA genes. Identifying archaeal members is important to our understanding of the functional properties of the microbiota. For example, a major challenge faced by microbial communities when breaking down polysaccharides (the most abundant biologic polymers on Earth) is the maintenance of redox balance in the setting of maximal energy production. Many microbial species have branched fermentation pathways that allow them to dispose of reducing equivalents (e.g., by the production of H_2 , which is energetically efficient). However, there is a caveat: the hydrogen must be removed or it will inhibit reoxidation of pyridine nucleotides. Therefore, hydrogen-consuming (*hydrogenotrophic*) species are key to maximizing the energy-extracting capacity of primary fermenters.

In the human gut, hydrogenotrophs include a phylogenetically diverse group of bacterial acetogens, a more limited group of sulfate-reducing bacteria that generate hydrogen sulfide, and methane-producing archaeal organisms (*methanogens*) that can represent up to 10% of the anaerobes present in the feces of some humans. However, the degree of archaeal diversity in the gut microbiota of healthy individuals appears to be low.

Culture-independent surveys of eukaryotic diversity are also confounded by challenges related to the design of PCR primers that target the eukaryotic SSU gene (18S rRNA) as well as the internal transcribed spacer regions of rRNA operons. Metagenomic studies of healthy human adults living in countries with distinct cultural traditions and disparate geographic features and locations have revealed that the degree of eukaryotic diversity is lower than that of bacterial diversity. In the gut, which contains far more microbes than any other body habitat, the representation of fungi is significantly lower in individuals living in Westernized societies than in those living in non-Western societies. The most abundant fungal sequences belong to the phylum-level taxa *Ascomycota* and *Microsporidia*. The phyla *Ascomycota* and *Basidiomycota* appear to be mutually exclusive, and the presence of *Candida* in particular correlates with recent consumption of carbohydrates.

Elucidation of Viral Dynamics Viruses are the most abundant biologic entity on Earth. Viral particles outnumber microbial cells by 10:1 in most environments. Humans are no exception in terms of viral colonization; our feces alone contain 10^8 – 10^9 viral particles per gram. Despite this abundance, many eukaryotic viral communities remain incompletely characterized, in part because the identification of viruses within metagenomic sequencing datasets is itself very challenging. Characterizing viral diversity requires different approaches: because no single gene is found in all viruses, no universal phylogenetic “barcode of life” equivalent to the SSU rRNA gene exists. One approach has been to selectively purify virus-like particles from community biospecimens, amplify the small amounts of DNA that are recovered, and randomly fragment the DNA and sequence the fragments (*shotgun sequencing*). The resulting sequences can be assembled into larger *contigs* whose function can be computationally predicted from homology to known genes, and the information obtained can be used to populate/expand nonredundant viral databases. These annotated nonredundant databases can then be used for more targeted mining of the rapidly expanding number of shotgun sequencing datasets generated from total-community DNA for known or putative DNA viruses.

Given the dominance of bacteria in the gut microbiota, it is not surprising that *phages* (viruses that infect bacteria) dominate the identifiable components of the gut’s DNA virome. *Prophages* are a manifestation of a so-called temperate viral–bacterial host dynamic, in which

a phage is integrated into its host bacterium’s genome. This temperate dynamic provides a way to constantly refashion the genomes of bacterial species through horizontal gene transfer. Genes encoded by a prophage genome may expand the niche and fitness of their bacterial host, for example, by enabling the metabolism of previously inaccessible nutrient sources. Prophage integration can also protect the host strain from superinfection, “immunizing” the strain against infection by closely related phages. A temperate prophage life cycle allows the virus to expand in a 1:1 ratio with its bacterial host. If the integrated virus conveys increased fitness, the prevalence of the bacterial host and its phage will increase in the microbiota. Induction of a lytic cycle, where the prophage replicates and kills the host, may follow. Lytic cycles can cause high bacterial turnover. Lysis debris (e.g., components of capsules) can be used as nutrient sources by surviving bacteria; this change in the energy dynamic in a community is referred to as a *phage shunt*. A subpopulation of bacteria that undergoes lytic induction may sweep away other sensitive species present in the community, thus increasing the niche space available for survivors (i.e., those bacteria that already have an integrated prophage). Periodic induction of prophages leads to a “constant diversity dynamic” that helps maintain community structure and function.

Interest in viral communities has expanded in recent years, especially given a potentially therapeutic role for phages as an alternative or adjunct to antibiotics. Virome members have evolved elegant survival mechanisms that allow them to evade host defenses, diversify, and establish elaborate and mutually beneficial symbioses with their hosts. A number of recent studies have tried to adapt these mechanisms for therapeutic purposes (e.g., the use of synthetic phages to treat *Pseudomonas aeruginosa* infections in burn patients or in other settings). Phage therapy is not a new idea: Félix d’Herelle, co-discoverer of phages, recognized their potential medical applications nearly a century ago. However, only recently have our technologic capabilities and our knowledge of the human microbiota made phage therapy realistically attainable within our lifetimes.

ECOLOGIC PRINCIPLES AND PARAMETERS FOR COMPARING MICROBIAL COMMUNITIES

At many levels, different people are very much alike: our genomes are >99% identical, and we have similar collections of human cells. However, our microbial communities differ drastically, both between people and between habitats within a single human body. The greatest variation (beta diversity, described below) is between body sites. For example, the difference between the microbial communities residing in a person’s mouth versus the same person’s gut is comparable to the difference in communities residing in soil versus seawater. Even within a body site, the differences among people are not subtle: gut, skin, and oral communities can all differ by 80–90%, even from the broad, bacterial species-level view. The English poet John Donne said that “no man is an island”; however, from a microbial perspective, each of us consists of not just one isolated island but rather a whole archipelago of distinct habitats that exchange microbes with one another and with the outside environment at some as yet undetermined level. Before we can discuss these differences and understand their relevance to human disease, it is important to understand some basic terms and ecologic principles.

Alpha Diversity *Alpha diversity* is defined as the effective number of species present in a given sample. Communities that are compositionally more diverse (i.e., have more OTUs) or that are phylogenetically more diverse are defined as having greater alpha diversity. Alpha diversity can be measured by plotting the number of different types of SSU rRNA sequences identified at a given phylogenetic level (species, genera, etc.) in a sample as a function of the number of SSU rRNA gene reads collected. The most commonly used metrics of alpha diversity are S_{obs} (the number of species observed in a given number of sequences), Chao1 (a measure based on the number of species observed only once), the Shannon index (a measure of the number of bits of information gained by revealing the identity of a randomly chosen member of the community), and phylogenetic diversity

(a measure of the total branch length of a phylogenetic tree encompassing a sample). Diversity estimators are particularly sensitive to errors introduced during PCR and sequencing.

Beta Diversity *Beta diversity* refers to the differences between communities and can be defined with phylogenetic or nonphylogenetic distance measurements. UniFrac is a commonly used phylogenetic metric that compares the evolutionary history of different microbial communities, noting the degree to which any two communities share branch length on a tree of microbial life: the more similar communities are to each other, the more branch length they share (Fig. 86e-1). UniFrac-based measurements of distances between communities can be visually represented with principal coordinates analysis or other geometric techniques that project a high-dimensional dataset down onto a small number of dimensions for a more approachable analysis (Fig. 86e-1). Principal coordinates analysis can also be applied to non-phylogenetic methods for comparing communities, such as Euclidean distance, Jensen-Shannon divergence, or Bray-Curtis dissimilarity, which operate independent of evolutionary tree data but can make biologic patterns more difficult to identify. The taxonomic data or distance matrices can also be used as input into a range of machine-learning algorithms (such as Random Forests) that employ supervised classification to identify differences between labeled groups of samples. Supervised classification is useful for identifying differences between cases and controls but can obscure important patterns intrinsic to the data, including confounding variables such as different sequencing runs or patient populations.

As noted above, the greatest beta diversity is that among body sites. This fact underscores the need to specify body habitat in microbiota analyses of any type, including microbial surveillance studies examining the flow of normal and pathogenic organisms into and out of different body sites in patients and their health care providers. Several other key points have emerged from beta diversity studies of human-associated microbial communities—notably, that (1) there is a high level of *interpersonal* variability in every body habitat studied to date, (2) *intrapersonal* variation in a given body habitat is less pronounced, and (3) family members have more similar communities than unrelated individuals living in separate households. Thus, a person is his/her own best control, and examination of an individual over time as a function of disease state or treatment intervention is desirable. Similarly, family members serve as logical reference controls, although age is a major covariate that affects microbiota structure.

Studies of fecal samples obtained from twins over time have shown that the overall degree of phylogenetic similarity of bacterial communities does not differ significantly between monozygotic and dizygotic twin pairs, although monozygotic twin pairs may be more similar in some populations at earlier ages. These results, together with intervention studies in mice and epidemiologic observations in humans, emphasize that early environmental exposures are a very important determinant of adult-gut microbial ecology. In humans, the initial exposures depend on delivery mode: babies sampled within 20 min of birth have relatively undifferentiated microbial communities in the mouth, the skin, and the gut. For vaginally delivered babies, these communities resemble the specific microbial communities found in the mother's vagina. For babies delivered by cesarean section, the communities resemble skin communities. Although studies of older children and of adults stratified by delivery mode are still rare in the literature, these differences have been shown to persist until at least 4 months of age and perhaps until age 7 years. The infant gut microbiota changes to resemble the adult gut community over the first 3 years of life; comparable studies have not been done in other body habitats to date.

Exposures to environmental microbial reservoirs can continue to influence community structure. For example, unrelated cohabiting adults have more similar microbiotas in all of their body habitats than do non-cohabiting adults, and humans resemble the dogs they live with, at least in terms of skin microbiota. Gender and sexual maturation may also affect the microbiota structure, although efforts to isolate these variables are complicated by many confounding factors; any gender effect must be small compared with the effects of other variables such as

diet (except in the case of the female urinary tract, which is influenced by the vaginal microbiota).

The vaginal microbiota illustrates another intriguing aspect of the contributions made by various factors to interpersonal differences in microbial community structure within a given body habitat. Bacterial 16S rRNA-based studies of the midvaginal microbiota in sexually active women have documented significant differences in community configurations between four self-reported ethnic groups: Caucasian, black, Hispanic, and Asian. Unlike most other body habitats that have been surveyed, this ecosystem is dominated by a single genus, *Lactobacillus*. Four species of this genus together account for more than half of the bacteria in most vaginal communities. Five community categories have been defined: four are dominated by *L. iners*, *L. crispatus*, *L. gasseri*, and *L. jensenii*, respectively, and the fifth has proportionally fewer lactobacilli and more anaerobes. The representation of these community categories is distinct within each of the four ethnic groups and correlates with vaginal pH and Nugent score (the latter being a biomarker for bacterial vaginosis). Longitudinal studies of individuals are being conducted to identify factors that determine the assembly of these distinct communities—both within and among ethnic groups—as well as their resistance to or resilience after various physiologic and pathologic disturbances. For example, the menstrual cycle and pregnancy turn out to be surprisingly significant factors (cause larger changes) compared with sexual activity.

Yet another factor affecting beta diversity is spatial location within a habitat. Several surveys show that the skin harbors bacterial communities with predictable, albeit complex, biogeographic features. To determine whether these differences are due to differences in local environmental factors, to the history of a given site's exposure to microbes, or to a combination of the two, reciprocal microbiota transplantation has been performed. Microbial communities from one region of the skin were depleted by treatment with germicidal agents, and the region (*plot*) was inoculated with a “foreign” microbiota harvested from different regions of the skin or from different body habitats from the same or another individual. Community assembly at the site of transplantation was then tracked over time. Remarkably, assembly proceeded differently at different sites: forearm plots receiving a tongue microbiota remained more similar to tongue communities than to native forearm communities in terms of their composition and diversity, while forehead plots inoculated with tongue bacteria changed to become more similar to native forehead communities. Thus, in addition to the history of exposure to tongue bacteria, environmental factors operating at the forehead plot likely shape community assembly. Intriguingly, the factors that shape fungal skin communities appear to be entirely different from those that shape bacterial skin communities. The palm and forearm have high bacterial and low fungal diversity, whereas the feet have the opposite diversity pattern. Moreover, fungal communities are generally shaped by location (foot, torso, head), whereas bacterial communities are generally shaped by moisture phenotype (dry, moist, or sebaceous).

Co-Occurrence Analysis *Co-occurrence analysis* seeks to identify which phylotypes are co-distributed across individuals in a given body habitat and/or between habitats and to determine the factors that explain the observed patterns of co-distribution. Positive correlations tend to reflect shared preferences for certain environmental features, while negative correlations typically reflect divergent preferences or a competitive relationship. *Syntrophic* (cross-feeding) relationships reflect interdependent interactions based on nutrient-sharing strategies. For example, in *food webs*, the products of one organism's metabolism can be used by the other for its own unique metabolic capabilities (e.g., the interactions between fermentative organisms and methanogens).

Enterotype Analysis *Enterotype analysis* seeks to classify individuals into discrete groups based on the configuration of their microbiotas, essentially drawing boundaries on a map defined by principal coordinates analysis or other ordination techniques. The first enterotype analysis used supervised clustering to define three major types of human-gut microbial configurations across three distinct human studies and provided a view that presupposed the existence of three

clusters. Subsequent work has shown that the range of variability in the gut microbiota of children and of non-Western populations greatly exceeds the variability captured in the populations used to define the original enterotypes; in addition, even in Western populations, the variability follows more of a continuum dominated by a gradient in the abundance of the genera *Bacteroides* and *Prevotella*. Another consideration in enterotype analysis is whether location on a map defined by healthy human variation is relevant to predisposition to disease or whether instead rare species with particular functions are more important discriminants.

Functional Redundancy *Functional redundancy* arises when functions are performed by many bacterial taxa. Thus interpersonal differences in microbial bacterial diversity (i.e., which bacteria are present) are not necessarily accompanied by comparable degrees of difference in functional diversity (i.e., what these bacteria can do). Characterization of a microbiome by shotgun sequencing is important because, unlike SSU rRNA analyses, shotgun sequencing provides a direct readout of the genes (and, via comparative genomics, their functions) in a given community. One fundamental question is the degree to which variations in the species occupying a given body habitat correlate with variations in a community's functional capabilities. For example, the *neutral theory of community assembly* developed by macroecologists suggests that species are added to the community without respect to function, automatically endowing the community with functional redundancy. If applicable to the microbial world, neutral community assembly would predict a high level of variation in the types of microbial lineages that occupy a given body habitat in different individuals, although the broad functions encoded in the microbiomes of these communities could be quite similar.

Shotgun sequencing of the fecal microbiome has revealed that different microbial communities converge on the same functional state: in other words, there is a group of microbial genes represented in the guts of unrelated as well as related individuals. The same principle holds true at other body sites (Fig. 86e-2). The “core” gut microbiome is enriched in functions related to microbial survival (e.g., translation; metabolism of nucleotides, carbohydrates, and amino acids) and in functions that benefit the host (nutrient and energy partitioning from the diet to microbes and host). The latter functions encompass the food webs mentioned above, in which products of one type of microbe become the substrates for other microbes. These webs, which can be incredibly elaborate, change as microbes adjust their patterns of gene expression and metabolism in response to alterations in nutrient availability. Thus the sum of all the activities of the members of a microbial community can be viewed as an emergent rather than a fixed property.

It is important to note that pairwise comparisons have shown that family members have functionally more similar gut microbiomes than do unrelated individuals. Thus, intrafamilial transmission of a gut microbiome within a given generation and across multiple generations could shape the biologic features of humans belonging to a kinship and modulate/mediate risks for a variety of diseases.

Stability Like other ecosystems, human body habitat-associated microbial communities vary over time, and an understanding of this variation is essential for a functional understanding of our microbiota. Few high-resolution time series of individual healthy adults have been published to date, but one available daily time series suggests that individuals tend to resemble themselves microbially day to day over a span of 6–15 months, retaining their separate identities during cohabitation. The development of low-error amplicon sequencing methods has provided a much more reliable

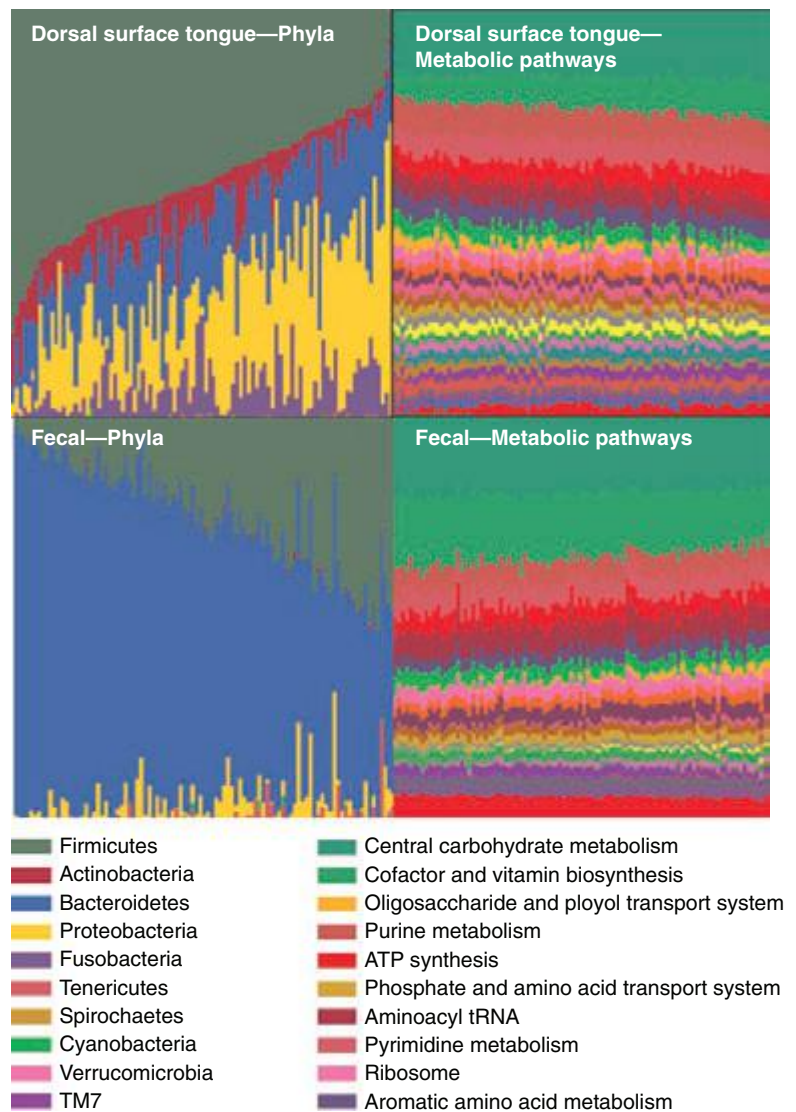


FIGURE 86e-2 Interpersonal variation in organismal representation in body habitat-associated communities is more extensive than interpersonal variation in gene functional features. Bacterial taxonomy and metabolic function are compared in 107 oral microbiota and microbiome samples (top) and in 139 fecal microbiota and microbiome samples (bottom). Samples represent an arbitrarily chosen subset from 242 healthy young adults living in the United States, with equal numbers of men and women. The same DNA extracts from the same samples were used for both taxonomic and functional classifications; each sample was analyzed by bacterial 16S rRNA amplicon sequencing (mean, 5400 sequences per sample) and by shotgun sequencing of community DNA (mean, 2.9 billion bases per sample). Taxonomic groups vary dramatically in their representation among different samples, with different characteristic bacterial phyla in the oral versus the fecal microbiota; e.g., members of the Actinobacteria and Fusobacteria are far more common in the mouth than in the gut, while members of Bacteroidetes are far more common in fecal samples. In contrast, metabolic pathways are far more consistently represented in different samples, even when the species that contribute to these pathways are completely different. These results suggest a high degree of functional redundancy in microbial ecosystems—similar to that observed in macroecosystems, in which many fundamentally different lineages of organisms can play the same ecologic roles (e.g., pollinator or top predator). (Adapted from Human Microbiome Project Consortium: *Nature* 486:207, 2012; and CA Lozupone et al: *Nature* 489:220, 2012.)

way for defining stability at the strain level than was available in the past. Application of these methods to the guts of healthy individuals sampled over time has disclosed that a healthy adult gut harbors a persistent collection of ~100 bacterial species and several hundred strains. The stability of the bacterial components follows a *power law*: bacterial strains acquired early in life can persist in the gut for decades, although

their proportional representation changes as a function of numerous factors, including diet. Whole-genome sequencing of culturable components of the microbiota of study participants has confirmed that strains are retained in individuals for prolonged periods and are shared among family members.

Resilience The ability of a microbiota or microbiome to rebound from a short-term perturbation, such as antibiotic administration or an infection, is defined as its *resilience*. This capacity can be visualized as a ball rolling over a landscape of local minima; essentially, the community moves into a new state and, to recover, must move through another, unstable state. In some cases, recovery will lead to the original stable state; in others, it will lead to a new stable state, which may be either healthy or unhealthy. Changes in, for example, diet or host physiologic status may introduce alterations into the landscape itself, making it easier to move from the initial state to any one of a number of other states, potentially with different health consequences. Microbial communities in our body habitats differ widely in resilience. For example, hand washing leads to profound changes in the microbial community, greatly increasing diversity (presumably because of the preferential removal of high-abundance, dominant phylotypes such as *Propionibacterium*). Within 6 h, the hand microbiota rebounds to resemble the original hand communities. The effects of repeated hand washing still need to be defined; for example, the surface microbiota on the skin (as measured by scrape biopsies) consists of ~50,000 microbial cells/cm², whereas the subsurface microbiota (as measured by punch biopsies) consists of ~1,000,000 microbial cells/cm².

In a study of three healthy adult volunteers given a short course of ciprofloxacin (500 mg by mouth twice a day for 5 days—a regimen commonly used against uncomplicated urinary tract infections), overall gut-community configuration came to resemble baseline within 6 months after treatment cessation, although some taxa failed to recover. However, the effects of the antibiotic perturbation were highly individualized. Administration of a second course of treatment months later led to altered-community states, relative to baseline, in all three volunteers; again, the extent of the alteration differed with the individual. Crucially, as shown in this and other studies, a given bacterial taxon can respond differently to the same antibiotic in different individuals; this observation suggests that the rest of the microbial community plays an important role in determining the effects of antibiotics on a per-individual basis.

In any body habitat, the microbial-community state after disturbance may be degraded. However, this degraded state may itself be resilient, and it may therefore be difficult to restore a more functional state. For example, *Clostridium difficile* infection can persist for years. The development and resilience of a degraded state may be driven by positive feedback loops, such as reactive oxygen species cascades involving host macrophages that promote the further growth of proinflammatory Proteobacteria, as well as negative-feedback loops such as depletion of the butyrate needed for promotion of a healthy gut epithelial barrier and further establishment of beneficial members of the microbiota. Consequently, microbiota-based therapies may require either (1) the elimination of a feedback loop that prevents establishment of a new community or (2) identification of a direction for change and a stimulus of sufficient magnitude (e.g., invasion and establishment of microbes from a fecal transplant or from a defined consortium of cultured, sequenced members of the human gut microbiota; see below) to overcome the resilience mechanisms inherent in the degraded state. A critical unresolved question that especially affects infants, whose microbiota is changing rapidly, is whether intervention during periods of rapid change or during periods of relative stability is generally more effective.

ESTABLISHING CAUSAL RELATIONSHIPS BETWEEN THE GUT MICROBIOTA AND NORMAL PHYSIOLOGIC, METABOLIC, AND IMMUNOLOGIC PHENOTYPES AS WELL AS DISEASE STATES

Gnotobiotic animals are raised in germ-free environments—with no exposure to microbes—and then colonized at specific stages of life with specified microbial communities. Gnotobiotic mice provide an excellent system for controlling host genotype, microbial community composition, diet, and housing conditions. Microbial communities

harvested from donor mice with defined genotypes and phenotypes can be used to determine how the donors' microbial communities affect the properties of formerly germ-free recipients. The recipients may also affect the transplanted microbiota and its microbiome. Thus gnotobiotic mice afford investigators an opportunity to marry comparative studies of donor communities to functional assays of community properties and to determine how (and for how long) these functions influence host biology.

The Cardiovascular System The gut microbiota affects the elaborate microvasculature underlying the small-intestinal epithelium: capillary network density is markedly reduced in adult germ-free animals but can be restored to normal levels within 2 weeks after gut microbiota transplantation. Mechanistic studies have shown that the microbiota promotes vascular remodeling in the gut through effects on a novel extravascular tissue factor–protease-activated receptor (PAR1) signaling pathway. Heart weight measured echocardiographically or as wet mass and normalized to tibial length or lean body weight is significantly reduced in germ-free mice; this difference is eliminated within 2 weeks after colonization with a gut microbiota. During fasting, a gut microbiota–dependent increase in hepatic ketogenesis (regulated by peroxisome proliferator–activated receptor α) occurs, and myocardial metabolism is directed to ketone body utilization. Analyses of isolated, perfused working hearts from germ-free and colonized animals, together with *in vivo* assessments, have shown that myocardial performance in germ-free mice is maintained by increasing glucose utilization. However, heart weight is significantly reduced in both fasted and fed mice; this heart-mass phenotype is completely reversed in germ-free mice fed a ketogenic diet. These findings illustrate how the gut microbiota benefits the host during periods of nutrient deprivation and represent one link between gut microbes and cardiovascular metabolism and health.

Conventionally raised *apoE*-deficient mice develop a less severe form of atherosclerosis than their germ-free counterparts when fed a high-fiber diet. This protective effect of the microbiota is obviated when animals are fed a diet low in fiber and high in simple sugars and fat. A number of the beneficial effects attributed to diets with high proportional representation of whole grains, fruits, and vegetables are thought to be mediated by end products of microbial metabolism of dietary compounds, including short-chain fatty acids and metabolites derived from flavonoids. Conversely, microbes can convert otherwise harmless dietary compounds into metabolites that increase risk for cardiovascular disease. Studies of mice and human volunteers have revealed that gut microbiota metabolism of dietary L-carnitine, which is present in large amounts in red meat, yields trimethylamine-N-oxide, which can accelerate atherosclerosis in mice by suppressing reverse cholesterol transport.

Yet another facet of microbial influence on cardiovascular physiology was revealed in a study of mice deficient in *Olfir78* (a G protein–coupled receptor expressed in the juxtaglomerular apparatus, where it regulates renin secretion in response to short-chain fatty acids) or *Gpr41* (another short-chain fatty acid receptor that, together with *Olfir78*, is expressed in the smooth muscle cells present in small resistance vessels). This study demonstrated that the microbiota can modulate host blood pressure via short-chain fatty acids produced by microbial fermentation.

Bone Adult germ-free mice have greater bone mass than their conventionally raised counterparts. This increase in bone mass is associated with reduced numbers of osteoclasts per unit bone surface area, reduced numbers of CD11b+/GR1 osteoclast precursors in bone marrow, decreased numbers of CD4+ T cells, and reduced levels of expression of the osteolytic cytokine tumor necrosis factor α . Colonization with a normal gut microbiota resolves these observed differences between germ-free and conventionally raised animals.

Brain Adult germ-free and conventionally raised mice differ significantly in levels of 38 out of 196 identified cerebral metabolites, 10 of which have known roles in brain function; included in the latter group are *N*-acetylaspartic acid (a marker of neuronal health and attenuation),

pipecolic acid (a presynaptic modulator of γ -aminobutyric acid levels), and serine (an obligatory co-agonist at the glycine site of the N-methyl-D-aspartate receptor). Propionate, a short-chain fatty acid product of gut microbial-community metabolism of dietary fiber, affects expression of genes involved in intestinal gluconeogenesis via a gut-brain neural circuit involving free fatty-acid receptor 3; this effect provides a mechanistic explanation for the documented beneficial impact of dietary fiber in enhancing insulin sensitivity and reducing body mass and adiposity.

Studies of a mouse model (maternal immune activation) with stereotyped/repetitive and anxiety-like behaviors indicate that treatment with a member of the human gut microbiota, *Bacteroides fragilis*, corrects gut barrier (permeability) defects; reduces elevated levels of 4-ethylphenylsulfate, a metabolite seen in the maternal immune activation model that has been causally associated with the animals' behavioral phenotypes; and ameliorates some behavioral effects. These observations highlight the importance of further exploration of potentially co-evolved relationships between the microbiota and host behavior.

Immune Function Many foundational studies have shown that the gut microbiota plays a key role in the maturation of the innate as well as the adaptive components of the immune system. The intestinal epithelium, which is composed of four principal cell lineages (enterocytes plus goblet, Paneth, and enteroendocrine cells), acts as a physical and functional barrier to microbial penetration. Goblet cells produce mucus that overlies the epithelium, where it forms two layers: an outer (luminal-facing) looser layer that harbors microbes and a denser lower layer that normally excludes microbes. Members of the Paneth cell lineage reside at the base of crypts of Lieberkühn and secrete antimicrobial peptides. Studies in mice have demonstrated that Paneth cells directly sense the presence of a microbiota through expression of the signaling adaptor protein MyD88, which helps transduce signals to host cells upon recognition of microbial products through Toll-like receptors (TLRs). This recognition drives expression of antibacterial products (e.g., the lectin RegIIIy) that act to prevent microbial translocation across the gut mucosal barrier.

The intestine is enriched for B cells that produce IgA, which is secreted into the lumen; there it functions to exclude microbes from crossing the mucosal barrier and to restrict dissemination of food antigens. The microbiota plays a key role in development of an IgA response: germ-free mice display a marked reduction in IgA+ B cells. The absence of a normal IgA response can lead to a massive increase in bacterial load. B cell-derived IgA that targets specific members of the gut microbiota plays an important role in preventing activation of microbiota-specific T cells.

Gut bacterial species elicit development of protective T_H17 and T_H1 responses that help ward off pathogen attack. Members of the microbiota also promote the development of a specialized population of CD4+ T cells that prevent unwarranted inflammatory responses. These regulatory T cells (T_{regs}) are characterized by expression of the transcription factor forkhead box P3 (FOXP3) and by expression of other cell-surface markers. There is a paucity of T_{regs} in the colonic lamina propria of germ-free mice. Specific members of the microbiota—including a consortium of *Clostridium* strains isolated from the mouse and human gut as well as several human-gut *Bacteroides* species—expand the T_{reg} compartment and enhance immunosuppressive functions.

The microbiota is a key trigger in the development of inflammatory bowel disease (IBD) in mice that harbor mutations in genes associated with IBD risk in humans. Moreover, components of the gut microbiota can modify the activity of the immune system to ameliorate or prevent IBD. Mice containing a mutant ATG16L1 allele linked to Crohn's disease are particularly susceptible to IBD. Upon infection with mouse norovirus and treatment with dextran sodium sulfate, expression of a hypomorphic ATG16L1 allele leads to defects in small-intestinal Paneth cells and renders mice significantly more susceptible to ileitis than are wild-type control animals. This process is dependent on the gut microbiota and highlights how the intersection of host genetics, infectious agents, and the microbiota can lead to severe immune

pathology; i.e., the pathogenic potential of a microbiota may be context-dependent, requiring a confluence of factors. An important observation is that members of the gut microbiota, including *B. fragilis* or members of *Clostridium*, prevent the severe inflammation that develops in mouse models mimicking various aspects of human IBD.

The gut microbiota has been implicated in promoting immunopathology outside of the intestine. Multiple sclerosis develops in conventionally raised mice whose CD4+ T cell compartment is reactive to myelin oligodendrocyte protein; their germ-free counterparts are completely protected from development of multiple sclerosis-like symptoms. This protection is reversed by colonization with a gut microbiota from conventionally raised animals.

Inflammasomes are cytoplasmic multiprotein complexes that sense stress and damage-associated patterns. Mice deficient in NLRP6, a component of the inflammasome, are more susceptible to colitis induced by administration of dextran sodium sulfate. This enhanced susceptibility is associated with alterations in the gut microbiota of these animals relative to that of wild-type controls. Mice are coprophagic, and co-housing of NLRP6-deficient mice with wild-type mice is sufficient to transfer the enhanced susceptibility to colitis induced by dextran sodium sulfate. Similar findings have been reported for mice deficient in the inflammasome adaptor ASC (apoptosis-associated speck-like protein containing a caspase recruitment domain). ASC-deficient mice are more susceptible to the development of a model of nonalcoholic steatohepatitis. This susceptibility is associated with alterations in gut microbiota structure and can be transferred to wild-type animals by co-housing.

Obesity and Diabetes Germ-free mice are resistant to diet-induced obesity. Genetically obese *ob/ob* mice have gut microbial-community structures that are profoundly altered from those in their lean wild-type (+/+) and heterozygous *+/ob* littermates. Transplantation of the *ob/ob* mouse microbiota into wild-type germ-free animals transmits an increased-adiposity phenotype not seen in mice receiving microbiota transplants from +/+ and *+/ob* littermates. These differences are not attributable to differences in food consumption but rather are associated with differences in microbial community metabolism. Roux-en-Y gastric bypass produces pronounced decreases in weight and adiposity as well as improved glucose metabolism—changes that are not ascribable simply to decreased caloric intake or reduced nutrient absorption. 16S rRNA analyses have documented that changes in the gut microbiota after this surgery are conserved among mice, rats, and humans; animal studies have demonstrated these changes along the length of the gut but most prominently downstream of the site of surgical manipulation of the bowel. Notably, transplantation of the gut microbiota from mice that have undergone Roux-en-Y gastric bypass to germ-free mice that have not had this surgery produces reductions in weight and adiposity not seen in recipients of microbiotas from mice that underwent sham surgery.

The gut microbiota confers protection against the development of type 1 diabetes mellitus in the non-obese diabetic (NOD) mouse model. Disease incidence is significantly lower in conventionally raised male NOD mice than in their female counterparts, while germ-free males are as susceptible as their female counterparts. Castration of males increases disease incidence, while androgen treatment of females provides protection. Transfer of the gut microbiota from adult male NOD mice to female NOD weanlings is sufficient to reduce the severity of disease relative to that among females receiving a microbiota from an adult female or an unmanipulated female. The blocking of protection by treatment with flutamide highlights a functional role for testosterone signaling in this microbiota-mediated protection against type 1 diabetes.

NOD mice deficient in MyD88, a key component of the TLR signaling pathway, do not develop diabetes and exhibit increased relative abundance of members of the family-level taxon Lactobacillaceae. Consistent with these findings, investigators have documented lower levels of representation of members of the genus *Lactobacillus* in children with type 1 diabetes than in healthy controls. Components of lactobacilli have been shown to promote gut barrier integrity. Studies

in various animal models indicate that translocation of bacterial components, including bacterial lipopolysaccharides, across a leaky gut barrier triggers low-grade inflammation, which contributes to insulin resistance. Mice deficient in TLR5 exhibit alterations in the gut microbiota and hyperphagia, and they develop features of metabolic syndrome, including hypertension, hyperlipidemia, insulin resistance, and increased adiposity.

The gut microbiota regulates biosynthesis as well as metabolism of host-derived products; these products can signal through host receptors to shape host physiology. An example of this symbiosis is provided by bile acids, which direct metabolic effects that are largely mediated through the farnesoid X receptor (FXR, also known as NR1H4). In leptin-deficient mice, FXR deficiency protects against obesity and improves insulin sensitivity. In mice with diet-induced obesity that are subjected to vertical sleeve gastrectomy, the surgical procedure results in elevated levels of circulating bile acids, changes in the gut microbiota, weight loss, and improved glucose homeostasis. However, weight reduction and improved insulin sensitivity are mitigated in animals with engineered FXR-deficiency.

Xenobiotic Metabolism Evidence is accumulating that pharmacogenomic studies need to consider the gene repertoire present in our *H. sapiens* genome as well as that in our microbiomes. For example, digoxin is inactivated by the human gut bacterium *Eggerthella lenta*, but only by strains with a cytochrome-containing operon. Expression of this operon is induced by digoxin and inhibited by arginine. Studies in gnotobiotic mice established that dietary protein affects (reduces) microbial metabolism of digoxin, with corresponding alterations in levels of the drug in both serum and urine. These findings reinforce the need to consider strain-level diversity in the gut microbiota when examining interpersonal variations in the metabolism of orally administered drugs.

Characterizing the Effects of the Human Microbiota on Host Biology in Mice and Humans Questions about the relationship between human microbial communities and health status can be posed in the following general format: Is there a consistent configuration of the microbiota definable in the study population that is associated with a given disease state? How is the configuration affected by remission/relapse or by treatment? If a reconfiguration does occur with treatment, is it durable? How is host biology related to the configuration or reconfiguration? What is the effect size? Are correlations robust to individuals from different families and communities representing different ages, geographic locales, and lifestyles?

As in all studies involving human microbial ecology, the issue of what constitutes a suitable reference control is extremely important. Should we choose the person himself or herself, family members, or age- or gender-matched individuals living in the same locale and representing similar cultural traditions? Critically, are the relationships observed between microbial community structure and expressed functions a response to disease state (i.e., side effects of other processes), or are they a contributing cause? In this sense, we are challenged to evolve a set of Koch's postulates that can be applied to whole microbial communities or components of communities rather than just to a single purified organism. As in other circumstances in which experiments to determine causality of human disease are difficult or unethical, Hill's criteria, which examine the strength, consistency, and biologic plausibility of epidemiologic data, can be useful.

Sets of mono- and dizygotic twins and their family members represent a valuable resource for initially teasing out relationships between environmental exposures, genotypes, and our own microbial ecology. Similarly, monozygotic twins discordant for various disease states enhance the ability to determine whether various diseases can be linked to a person's microbiota and microbiome. A twin-pair sampling design rather than a conventional unrelated case-control design has advantages owing to the pronounced between-family variability in microbiota/microbiome composition and the potential for multiple states of a community associated with disease. Transplantation of a microbiota from suitable human donor controls representing different disease states and communities (e.g., twins discordant for a disease)

to germ-free mice is helpful in establishing a causal role for the community in pathogenesis and for providing insights relevant to underlying mechanisms. In addition, transplantation provides a preclinical platform for identifying next-generation probiotics, prebiotics, or combinations of the two (*synbiotics*). Obesity and obesity-associated metabolic dysfunction illustrate these points.

The gut microbiotas (and microbiomes) of obese individuals are significantly less diverse than those of lean individuals; the implication is that there may be unfilled niches (unexpressed functions) that contribute to obesity and its associated metabolic abnormalities. Le Chatelier and colleagues observed a bimodal distribution of gene abundance in their analysis of 292 fecal microbiomes: low-gene-count (LGC) individuals averaged 380,000 microbial genes per gut microbiome, while high-gene-count (HGC) individuals averaged 640,000 genes. LGC individuals had an increased risk for type 2 diabetes and other metabolic abnormalities, whereas the HGC group was metabolically healthy. When gene content was used to identify taxa that discriminated HGC and LGC individuals, the results revealed associations between anti-inflammatory bacterial species such as *Faecalibacterium prausnitzii* and the HGC group and between proinflammatory species such as *Ruminococcus gnavus* and the LGC group. LGC microbiomes had significantly greater representation of genes assigned to tricarboxylic acid cycle modules, peroxidases, and catalases—an observation suggesting a greater capacity to handle oxygen exposure and oxidative stress; HGC microbiomes were enriched in genes involved in the production of organic acids, including lactate, propionate, and butyrate—a result suggesting increased fermentative capacity.

Transplantation of an uncultured fecal microbiota from twins stably discordant for obesity or of bacterial culture collections generated from their microbiota transmits their discordant adiposity phenotypes as well as obesity-associated metabolic abnormalities to recipient germ-free mice. Co-housing of the recipient coprophagic gnotobiotic mice results in invasion of specific bacterial species from the transplanted lean twin's culture collection into the guts of cage mates harboring the obese twin's culture collection (but not vice versa), thereby preventing the latter animals from developing obesity and its associated metabolic abnormalities. It is noteworthy that invasion and prevention of obesity and metabolic phenotypes are dependent on the type of human diets fed to animals: prevention is associated with a diet low in saturated fats and high in fruit and vegetable content, but not with a diet high in saturated fats and low in fruit and vegetable content.

This approach provides evidence for a causal role for the microbiota in obesity and its attendant metabolic abnormalities. It also provides a method for defining unoccupied niches in disease-associated microbial communities, the role of dietary components in determining how these niches can be filled by human gut-derived bacterial taxa, and the effects of such occupancy on microbial and host metabolism. It also provides a way to identify health-promoting diets and next-generation probiotics representing naturally occurring members of our indigenous microbial communities that are well adapted to persist in a given body habitat.

A key to this approach is the ability to harvest a microbial community from a donor representing a physiology, disease state, lifestyle, or geography of interest; to preserve the donor's community by freezing it; and then to resurrect and replicate it in multiple recipient gnotobiotic animals that can be reared under conditions where environmental and host variables can be controlled and manipulated to a degree not achievable in clinical studies. Since these mice can be followed as a function of time prior to and after transplantation, in essence, a snapshot of a donor's community can be converted into a movie. Transplantation of intact uncultured human (fecal) microbiota samples from multiple donors representing the phenotype of interest, with administration of the donors' diets (or derivatives of those diets) to different groups of mice, is one way to assess whether transmissible responses are shared features of the microbiota or are highly donor specific. A second step is to determine whether the culturable component of a representative microbiota sample can transmit the phenotype(s) observed with the intact uncultured sample. Possession of a collection of cultured organisms that have co-evolved in a given donor's body habitat sets the stage

for the selection of subsets of the collection for testing in gnotobiotic mice, the determination of which members are responsible for effecting the phenotype, and the elucidation of the mechanisms underlying these effects. The models used may inform the design and interpretation of clinical studies of the very individuals and populations whose microbiota are selected for creating these models.

Human-to-human fecal microbiota transplantation (FMT) is currently the most direct way to establish proof-of-concept for a causal role for the microbiota in disease pathogenesis. A human donor's feces are provided to a recipient via nasogastric tube or another technique. Numerous small trials have documented the effects of FMT from healthy donors to recipients with diseases ranging from *C. difficile* infection to Crohn's disease, ulcerative colitis, and type 2 diabetes. Only a few of these studies have used a double-blind, placebo-controlled design.

In a double-blind, controlled trial involving men 21–65 years old with a body mass index of >30 kg/m² and documented insulin resistance, FMT was performed using a microbiota from metabolically healthy lean donors or from the study participants themselves. A microbiota from lean donors significantly improved peripheral insulin sensitivity over that in controls. This change was associated with an increase in the relative abundance of the butyrate-producing bacteria related to *Roseburia intestinalis* (in the feces) and *Eubacterium hallii* (in the small intestine).

The efficacy of FMT for the treatment of recurrent *C. difficile* infection has been assessed in a number of small trials. One unblinded, placebo-controlled trial assessed the use of FMT in 42 patients with recurrent *C. difficile* infection (defined as at least one relapse after treatment with vancomycin or metronidazole for ≥10 d). Patients were pretreated with oral vancomycin. The experimental group then received FMT via nasoduodenal tube from healthy volunteer donors (<60 years of age) selected from the community. Controls underwent sterile lavage or received oral vancomycin alone. In 10 weeks of follow-up, infection was cured (with cure defined as three negative fecal tests for *C. difficile* toxin) in 81% of patients in the FMT group (13 of 16) but in only 23% (3 of 13) in the bowel-lavage control arm and 31% (4 of 13) in the vancomycin-only group. Metagenomic analysis of microbiota samples collected before and after treatment revealed an increased representation of Bacteroidetes and *Clostridium* clusters IV and XIVa, along with a 100-fold decrease in the relative abundance of Proteobacteria, in the FMT group.

A meta-analysis of FMT in *C. difficile* infection examined 20 case-series publications, 15 case reports, and the one unblinded study described above. All but one of these studies used fresh (not frozen) fecal samples. Donor selection varied, although most donors were family members or relatives and most studies excluded donors who had recently received antibiotics. It is noteworthy that the concentrations of infused donor feces varied widely (i.e., from 5 g to 200 g, resuspended in 10–500 mL); these fecal suspensions were introduced at different sites along the gastrointestinal tract, including the stomach and points throughout the small intestine and colon. Resolution of infection, which was frequently assessed on the basis of symptom resolution (with *C. difficile* toxin testing rarely performed), was documented in 87% (467) of 536 treated patients. The most common adverse events reported were diarrhea (94% of cases) and abdominal cramps (31%) on the day of infusion. The meta-analysis was limited to clinical outcomes and did not specifically address the role of the microbiota in disease resolution (e.g., the extent of invasion of donor taxa; their persistence; or the long-term effects of transplantation on various facets of host biology, which generally have not been evaluated).

Sober and thoughtful consideration needs to be applied to the therapeutic use of FMT, which represents an early and rudimentary approach to microbiota manipulation that very likely will be replaced by administration of defined collections of sequenced, cultured members of the human microbiota (*probiotic consortia*). A number of published reports on FMT have garnered significant public attention. This attention, coupled with an increasing public appreciation of the beneficial nature of our interactions with microbes, demands that the precautionary principle be honored and that risks versus benefits of such interventions be carefully evaluated.

To date, most FMT trials have failed to define (or have differed in) significant confounders, including (1) the criteria used for donor sample selection; (2) the methods used for donor sample preparation and characterization as well as the decision about whether or not to create a repository for donor and recipient samples that will permit retrospective analyses (and meta-analyses for given disease states); (3) the development of minimal standards for assessing the invasion of recipient gut communities by taxa from donor microbiota (using microbial source-tracking methods) as well as the timing, duration, nature, and breadth of sampling of the recipient as a function of transplantation; (4) the adoption of minimal standards for collection of patients' clinical data (e.g., age, diet, antibiotic use) and the establishment of databases for entering these data (including use of a defined vocabulary for annotating the clinical data); and (5) the development of standards for informed consent in lieu of knowledge of the long-term effects of the procedure. The regulatory landscape is evolving. The U. S. Food and Drug Administration recently issued an enforcement policy specifically addressing the use of FMT for the treatment of recurrent *C. difficile* infection; this policy indicates that the agency intends to “exercise enforcement discretion regarding the investigational new drug (IND) requirements for the use of FMT to treat *C. difficile* infection not responding to standard therapies,” but it does not waive IND requirements for other FMT studies.

MOVING FORWARD

The design of human microbiome studies is rapidly evolving, in part because the data are highly multivariate, are compositional, and do not meet distributional assumptions of standard statistical tests such as analysis of variance. Consequently, the proper number of subjects to enroll and the proper populations to target remain to be established. One useful approach is to review published studies and ask whether the reported conclusion could be obtained with fewer subjects (*sample rarefaction*) and/or fewer sequencing reads from SSU rRNA genes, whole-community DNA (microbiomes), or expressed community mRNA (metatranscriptomes) per subject (*sequence rarefaction*). A common yet critical problem to avoid is under-sampling of the types of objects under study. For example, if the goal is to compare factors applying to individuals (e.g., individual diet), then dozens of individuals in each clinical category may be needed. If the goal is to compare factors applying to populations (e.g., demographic properties), then many populations may be needed.

Another key issue is whether the effect size to be studied, especially in meta-analysis, is greater than or less than technical effects. As noted above, different PCR primers will lead to different readouts of the taxonomy of a microbial community; these differences are, for example, greater than the differences between lean and obese subjects' fecal microbiota but less than the difference between fecal communities in newborns and adults.

A central challenge in human microbiome research is establishing the extent to which diagnostic tests and therapeutic approaches are generalizable. This challenge is illustrated by studies of the capacities of gut microbiomes to metabolize orally administered drugs. The results could be very informative for the pharmaceutical industry as it seeks new and more accurate ways to predict bioavailability and toxicity. However, these studies should prompt consideration of the fact that many clinical trials are outsourced to countries where trial participants have diets and microbial community structures that differ from those of the intended initial recipients of the (marketed) drug. Capture and preservation of the wide range of microbial diversity present in different human populations—and thus of the capacity of our microbial communities to catalyze elaborate and in many respects uncharacterized biotransformations—represent potentially fertile ground for the discovery of new drugs (and new industrial processes of societal value). The chemical entities that our microbial communities have evolved to synthesize in order to support their mutually beneficial relationships and the human genes that these chemotypes influence may become new classes of drugs and new targets for drug discovery, respectively. Therefore, characterization of groups of individuals living in countries that are undergoing rapid transformations in cultural traditions

and socioeconomic conditions and are witnessing the emergence of a variety of diseases associated with increasingly Western lifestyles (globalization) is a timely challenge. Birth cohort studies (including studies of twins) initiated every 10 years in these countries may be able to capture the impact of globalization, including changing diets, on human microbial ecology.

Although microbiome-associated diagnostics and therapeutics provide new and exciting dimensions for personalized medicine, attention must be paid to the potentially broad societal impact of this work. For example, studies of the human gut microbiome are likely to have a disruptive effect on current views of human nutrition, enhancing appreciation of how food and the metabolic output of interactions of dietary components with the microbiota are intimately connected to myriad features of human biology. Underlying the efforts to elucidate the relations among food, the microbiome, and human nutrition is a need to proactively develop materials for educational outreach with a narrative and vocabulary that is understandable to broad and varied consumer populations representing different cultural traditions and widely ranging degrees of scientific literacy. The results have the potential to catalyze efforts to integrate agricultural policies and practice, food production, and nutritional recommendations for consumer populations representing different ages, geographic locales, and states of health.

Defining our *metagenome* (the genes embedded in our *H. sapiens* genome plus those in our microbiome) will likely lead to an entirely

new level of refinement in our description of self, our genetic evolution, our postnatal development, the microbial legacy of our connection to family, and the consequences of personal lifestyle choices. While this information can help us understand the origins of certain yet unexplained health disparities, care must be taken to avoid stigmatization of individuals or groups of individuals having different cultural norms, belief systems, or behaviors. In partnership with human microbiome researchers, anthropologists need to examine the impact of studies of the human microbiome on the participants, assessing how this field and participants' cultural traditions interact to affect these individuals' perceptions about the natural world, the forces that affect their lives, and their connections to one another within the context of family and community.

SUMMARY

Studies of human microbial ecology are an important manifestation of progress in the genome sciences, represent a timely step in our quest to achieve a better understanding of our place in the natural world, and reflect the evolving focus of twenty-first-century medicine on disease prevention, new definitions of health, new ways to determine the origins of individual biologic differences, and new approaches to evaluating the impact of changes in our lifestyles and biosphere on our biology. As microbiome-directed diagnostics and therapeutics emerge, we must be sensitive to the societal impact of this work.

The field of human biology has progressed over the last three centuries largely as a result of the reductionist approach to the scientific problems that challenge the discipline. Biologists study the experimental response of a variable of interest in a cell or organism while holding all other variables constant. In this way, it is possible to dissect the individual components of a biologic system and assume that a thorough understanding of a specific component (e.g., an enzyme or a transcription factor) will provide sufficient insight to explain the global behavior of that system (e.g., a metabolic pathway or a gene network, respectively). Biologic systems are, however, much more complex than this approach assumes and manifest behaviors that frequently (if not invariably) cannot be predicted from knowledge of their component parts characterized in isolation. Growing recognition of this shortcoming of conventional biologic research has led to the development of a new discipline, *systems biology*, which is defined as the holistic study of living organisms or their cellular or molecular network components to predict their response to perturbations. Concepts of systems biology can be applied readily to human disease and therapy and define the field of *systems pathobiology*, in which genetic or environmental perturbations produce disease and drug perturbations restore normal system behavior.

Systems biology evolved from the field of systems engineering in which a linked collection of component parts constitute a network whose output the engineer wishes to predict. The simple example of an electronic circuit can be used to illustrate some basic systems engineering concepts. All the individual elements of the circuit—resistors, capacitors, transistors—have well-defined properties that can be characterized precisely. However, they can be linked (wired or configured) in a variety of ways, each of which yields a circuit whose response to voltage applied across it is different from the response of every other configuration. To predict the circuit's (i.e., system's) behavior, the engineer must study its response to perturbation (e.g., voltage applied across it) holistically rather than its individual components' responses to that perturbation. Viewed another way, the resulting behavior of the system is greater than (or different from) the simple sum of its parts, and systems engineering utilizes rigorous mathematical approaches to predict these complex, often nonlinear, responses. By analogy to biologic systems, one can reason that detailed knowledge of a single enzyme in a metabolic pathway or of a single transcription factor in a gene network will not provide sufficient detail to predict the output of that metabolic pathway or transcriptional network, respectively. Only a systems-based approach will suffice.

It has taken biologists a long time to appreciate the importance of systems approaches to biomedical problems. Reductionism has reigned supreme for many decades, largely because it is experimentally and analytically simpler than holism, and because it has provided insights into biologic mechanisms and disease pathogenesis that have led to successful therapies. However, reductionism cannot solve all biomedical problems. For example, the so-called off-target effects of new drugs that frequently limit their adoption likely reflect the failure of a drug to be studied in holistic context, i.e., the failure to explore all possible actions aside from the principal target action for which it was developed. Other approaches to understanding biology therefore are clearly needed. With the growing body of genomic, proteomic, and metabolomic data sets in which dynamic changes in the expression of many genes and many metabolites are recorded after a perturbation and with the growth of rigorous mathematical approaches to analyzing those changes, the stage has been set for applying systems engineering principles to modern biology.

Physiologists historically have had more of a (bio)engineering perspective on the conduct of their studies and have been among the first systems biologists. Yet, with few exceptions, they, too, have focused

on comparatively simple physiologic systems that are tractable using conventional reductionist approaches. Efforts at integrative modeling of human physiologic systems, as first attempted by Guyton for blood pressure regulation, represent one application of systems engineering to human biology. These dynamic physiologic models often focus on the acute response of a measurable physiologic parameter to a system perturbation, and do so from a classic analytic perspective in which all the conventional physiologic determinants of the output parameter are known and can be modeled quantitatively.

Until recently, molecular systems analysis has been limited owing to inadequate knowledge of the molecular determinants of a biologic system of interest. Although biochemists have approached metabolic pathways from a systems perspective for over 50 years, their efforts have been limited by the inadequacy of key information for each enzyme (K_M , k_{cat} , and concentration) and substrate (concentration) in the pathway. With increasingly rich molecular data sets available for systems-based analyses, including genomic, transcriptomic, proteomic, and metabolomic data, biochemists are now poised to use systems biology approaches to explore biologic and pathobiologic phenomena.

PROPERTIES OF COMPLEX BIOLOGIC SYSTEMS

To understand how best to apply the principles of systems biology to human biomedicine, it is necessary to review briefly the building blocks of any biologic system and the determinants of system complexity. All systems can be analyzed by defining their static topology (architecture) and their dynamic (i.e., time-dependent) response to perturbation. In the discussion that follows, system properties are described that derive from the consequences of topology (form) or dynamic response (function). Any system of interacting elements can be represented schematically as a *network* in which the individual elements are depicted as nodes and their connections are depicted as links. The nature of the links among nodes reflects the degree of complexity of the system. *Simple systems* are those in which the nodes are linearly linked with occasional feedback or feedforward loops modulating system throughput in highly predictable ways. By contrast, *complex systems* are nodes that are linked in more complicated, nonlinear networks; the behavior of these systems by definition is inherently more difficult to predict owing to the nature of the interacting links, the dependence of the system's behavior on its initial conditions, and the inability to measure the overall state of the system at any specific time with great precision. Complex systems can be depicted as a network of lower-complexity interacting components or modules, each of which can be reduced further to simpler analyzable canonical motifs (such as feedback and feedforward loops, or negative and positive autoregulation); however, a central property of complex systems is that simplifying their structures by identifying and characterizing the individual nodes and links or even simpler substructures does not necessarily yield a predictable understanding of a system's behavior. Thus, the functioning system is greater than (or different from) the sum of its individual, tractable parts.

Defined in this way, most biologic systems are complex systems that can be represented as networks whose behaviors are not readily predictable from simple reductionist principles. The nodes, for example, can be metabolites that are linked by the enzymes that cause their transformations, transcription factors that are linked by the genes whose expression they influence, or proteins in an interaction network that are linked by cofactors that facilitate interactions or by thermodynamic forces that facilitate their physical association. Biologic systems typically are organized as *scale-free*, rather than stochastic, networks of nodes. Scale-free networks are those in which a few nodes have many links to other nodes (highly linked nodes, or hubs) but most nodes have only a few links (weakly linked nodes). The term *scale-free* refers to the fact that the connectivity of nodes in the network is invariant with respect to the size of the network. This is quite different from two other common network architectures: random (Poisson) and exponential distributions. Scale-free networks can be mathematically described by a power law that defines the probability of the number of

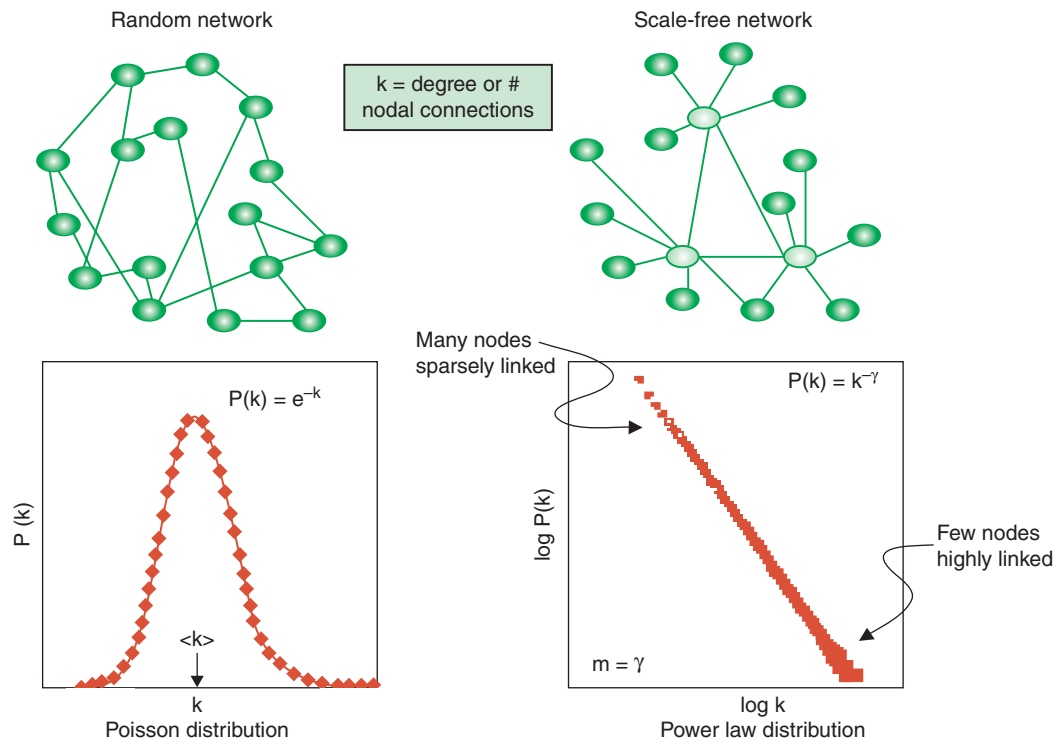


FIGURE 87e-1 Network representations and their distributions. A random network is depicted on the left, and its Poisson distribution of the number of nodal connections (k) is shown in the graph below it. A scale-free network is depicted on the right, and its power law distribution of the number of nodal connections (k) is shown in the graph below it. Highly connected nodes (hubs) are lightly shaded.

links per node ($P[k] = k^{-\gamma}$, where k is the number of links per node and γ is the slope of the $\log P[k]$ versus $\log[k]$ plot); this unique property of most biologic networks is a reflection of their self-similarity or fractal nature (Fig. 87e-1).

There are unique properties of scale-free biologic systems that reflect their evolution and promote their adaptability and survival. Biologic networks likely evolved one node at a time in a process in which new nodes are more likely to link to a highly connected node than to a sparsely connected node. Furthermore, scale-free networks can become sparsely linked to one another, yielding more complex, *modular scale-free topologies*. This evolutionary growth of biologic networks has three important properties that affect system function and survival. First, this scale-free addition of new nodes promotes *system redundancy*, which minimizes the consequences of errors and accommodates adverse perturbations to the system robustly with minimal effects on critical functions (unless the highly connected nodes are the focus of the perturbation). Second, this resulting network redundancy provides a survival advantage to the system. In complex gene networks, for example, mutations or polymorphisms in weakly linked genes account for biodiversity and biologic variability without disrupting the critical functions of the system; only mutations in highly linked (*essential*) genes (hubs) can shut down the system and cause embryonic lethality. Third, scale-free biologic systems facilitate the flow of information (e.g., metabolite flux) across the system compared with randomly organized biologic systems; this so-called “small-world” property of the system (in which the clustered nature of the highly linked hubs defines a local neighborhood within the network that communicates through weaker, less frequent links to other clusters) minimizes the energy cost for the dynamic action of the system (e.g., minimizes the transition time between states in a metabolic network).

These basic organizing principles of complex biologic systems lead to three unique properties that require emphasis. First, biologic systems are *robust*, which means that they are quite stable in response to most changes in external conditions or internal modification. Second, a corollary to the property of robustness is that complex biologic systems are *sloppy*, which means that they are insensitive to changes in external conditions or internal modification except under certain

uncommon conditions (i.e., when a hub is involved in the change). Third, complex biologic systems exhibit *emergent properties*, which means that they manifest behaviors that cannot be predicted from the reductionist principles used to characterize their component parts. Examples of emergent behavior in biologic systems include spontaneous, self-sustained oscillations in glycolysis; spiral and scroll waves of depolarization in cardiac tissue that cause reentrant arrhythmias; and self-organizing patterns in biochemical systems governed by diffusion and chemical reaction.

APPLICATIONS OF SYSTEMS BIOLOGY TO PATHOBIOLOGY

The principles of systems biology have been applied to complex pathologic processes with some early successes. The key to these applications is the identification of emergent properties of the system under study in order to define novel, otherwise unpredictable (i.e., from the reductionist perspective) methods for regulating the system’s response. Systems biology approaches have been used to characterize epidemics and ways to control them, taking advantage of the scale-free properties of the network of infected individuals that constitute the epidemic. Through the use of a systems analysis of a neural protein-protein interaction network, unique disease-modifying proteins have been identified that are common to a wide range of cerebellar neurodegenerative disorders causing inherited ataxias. Systems analysis and disease network construction of a pulmonary arterial hypertension network led to the identification of a unique disease module involving a pathway governed by microRNA21. Systems biology models have been used to dissect the dynamics of the inflammatory response using oscillatory changes in the transcription factor nuclear factor (NF) κ B as the system output. Systems biology principles also have been used to predict the development of an idiotypic–anti-idiotypic antibody network, describe the dynamics of species growth in microbial biofilms, and analyze the innate immune response. In each of these examples, a systems (patho)biology approach provided insights into the behavior of these complex systems that could not have been recognized with conventional scientific reductionism.

A unique application of systems biology to biomedicine is in the area of drug development. Conventional drug development involves

identifying a potential target protein and then designing or screening compounds to identify those that inhibit the function of that target. This reductionist analysis has identified many potential drug targets and drugs, yet only when a drug is tested in animal models or humans are the systems consequences of the drug's action revealed; not uncommonly, so-called off-target effects may become apparent and be sufficiently adverse for researchers to cease development of the agent. A good example of this problem is the unexpected outcomes of the vitamin B–based regimens for lowering homocysteine levels. In these trials, plasma homocysteine levels were reduced effectively; however, there was no effect of this reduction on clinical vascular endpoints. One explanation for this outcome is that one of the B vitamins in the regimen, folate, has a panoply of effects on cell proliferation and metabolism that probably offset its homocysteine-lowering benefits, promoting progressive atherosclerotic plaque growth and its consequences for clinical events. In addition to these types of unexpected outcomes exerted through pathways that were not considered *ab initio*, conventional approaches to drug development typically do not take into consideration the possibility of emergent behaviors of the organism or the metabolic pathway or the transcriptional network of interest. Thus, a systems-based analysis of potential drugs (drug-target network analysis) can benefit the development paradigm both by enhancing the likelihood that a compound of interest will not manifest unforeseen adverse effects and by promoting novel analytic methods for identifying unique control points or pathways in metabolic or genetic networks that would benefit from drug-based modulation.

SYSTEMS PATHOBIOLOGY AND HUMAN DISEASE CLASSIFICATION: NETWORK MEDICINE

Perhaps most important, systems pathobiology can be used to revise and refine the definition of human disease. The classification of human disease used in this and all medical textbooks derives from the correlation between pathologic analysis and clinical syndromes that began in the nineteenth century. Although this approach has been very successful, serving as the basis for the development of many effective therapies, it has major shortcomings. Those shortcomings include a lack of sensitivity in defining preclinical disease, a primary focus on overtly manifest disease, failure to recognize different and potentially differentiable causes of common late-stage pathophenotypes, and a limited ability to incorporate the growing body of molecular and genetic determinants of pathophenotype into the conventional classification scheme.

Two examples will illustrate the weakness of simple correlation analyses grounded in the reductionist principle of simplification (Occam's razor) in defining human disease. Sickle cell anemia, the "classic" Mendelian disorder, is caused by a Val6Gln substitution in the β chain of hemoglobin. If conventional genetic teaching holds, this single mutation should lead to a single phenotype in patients who harbor it (genotype-phenotype correlation). This assumption is, however, false, as patients with sickle cell disease manifest a variety of pathophenotypes, including hemolytic anemia, stroke, acute chest syndrome, bony infarction, and painful crisis, as well as an overtly normal phenotype. The reasons for these different phenotypic presentations include the presence of disease-modifying genes or gene products (e.g., hemoglobin F, hemoglobin C, glucose-6-phosphate dehydrogenase), exposure to adverse environmental factors (e.g., hypoxia, dehydration), and the genetic and environmental determinants of common intermediate pathophenotypes (i.e., variations in those generic pathologic mechanisms underlying all human disease—*inflammation, thrombosis/hemorrhage, fibrosis, cell proliferation, apoptosis/necrosis, immune response*).

A second example of note is familial pulmonary arterial hypertension. This disorder is associated with over 100 different mutations in three members of the transforming growth factor β (TGF- β) superfamily: bone morphogenetic protein receptor-2 (BMPR-2), activin receptor-like kinase-1 (Alk-1), and endoglin. All these different genotypes are associated with a common pathophenotype, and each leads to that pathophenotype by molecular mechanisms that range from haploinsufficiency to dominant negative effects. As only approximately one-fourth of individuals in families that harbor these mutations

manifest the pathophenotype, other disease-modifying genes (e.g., the serotonin receptor 5-HT_{2B}, the serotonin transporter 5-HTT), genomic and environmental determinants of common intermediate pathophenotypes, and environmental exposures (e.g., hypoxia, infective agents [HIV], anorexigens) probably account for the incomplete penetrance of the disorder.

On the basis of these and many other related examples, one can approach human disease from a systems pathobiology perspective in which each "disease" can be depicted as a network that includes the following modules: the primary disease-determining elements of the genome (or proteome, if posttranslationally modified), the disease-modifying elements of the genome or proteome, environmental determinants, and genomic and environmental determinants of the generic intermediate pathophenotypes. **Figure 87e-2** graphically depicts these genotype-phenotype relationships as modules for the six common disease types with specific examples for each type. **Figure 87e-3** shows a network-based depiction of sickle cell disease using this kind of modular approach.

Goh and colleagues developed the concept of a human disease network (**Fig. 87e-4**) in which they used a systems approach to characterize the disease-gene associations listed in the Online Mendelian Inheritance in Man database. Their analysis showed that genes linked to similar disorders are more likely to have products that physically associate and greater similarity between their transcription profiles than do genes not associated with similar disorders. In addition, proteins associated with the same pathophenotype are significantly more likely to interact with one another than with other proteins not associated with the pathophenotype. Finally, these authors showed that the great majority of disease-associated genes are not highly connected genes (i.e., not hubs) and are typically weakly linked nodes within the functional periphery of the network in which they operate.

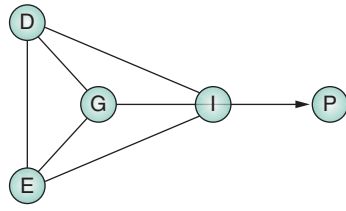
This type of analysis validates the potential importance of defining disease on the basis of its systems pathobiologic determinants. Clearly, doing this will require a more careful dissection of the molecular elements in the relevant pathways (i.e., more precise molecular pathophenotyping), less reliance on overt manifestations of disease for their classification, and an understanding of the dynamics (not just the static architecture) of the pathobiologic networks that underlie pathophenotypes defined in this way. **Figure 87e-5** illustrates the elements of a molecular network within which a disease module is contained. This network is first identified by determining the interactions (physical or regulatory) among the proteins or genes that comprise it (the "interactome"). These interactions then define a topologic module within which exists functional modules (pathways) and disease modules. One approach to constructing this module is illustrated in **Fig. 87e-6**. Examples of the use of this approach in defining novel determinants of disease are given in **Table 87e-1**.

TABLE 87e-1 EXAMPLES OF SYSTEMS BIOLOGY APPLICATION TO DISEASE

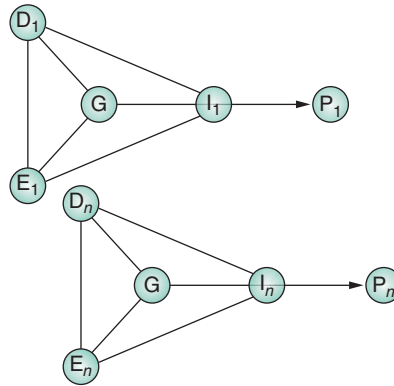
Disease	Analysis	Reference
Hereditary ataxias	Many ataxia-causing proteins share interacting partners that affect neurodegeneration	Lim et al: Cell 125:801-814, 2006
Diabetes mellitus	Metabolite-protein network analysis links three unique metabolite abnormalities in prediabetics to seven type 2 diabetes genes through four enzymes	Wang-Sattler et al: Mol Syst Biol 8:615, 2012
Ebstein-Barr virus infection	Viral proteome exerts its effects through linking to host interactome	Gulbahce et al: PLoS One 8:e1002531, 2012
Pulmonary arterial hypertension	Network analysis indicates adaptive role for microRNA 21 in suppressing rho kinase pathway	Parikh et al: Circulation 125:1520-1532, 2012

**Classic mendelian disorder:
Single phenotype**

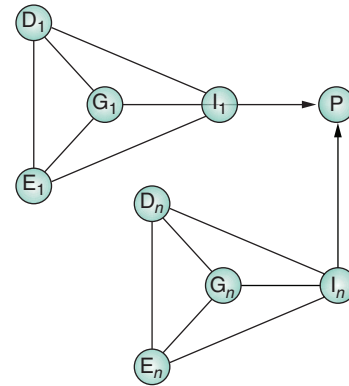
Example:?

**Classic mendelian disorder:
Multiple phenotypes**

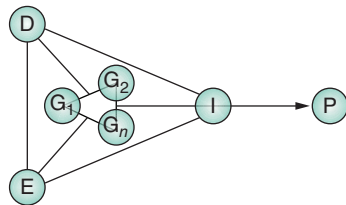
Example: Sickle cell disease

**Classic mendelian disorder:
Multiple mutations, single phenotype**

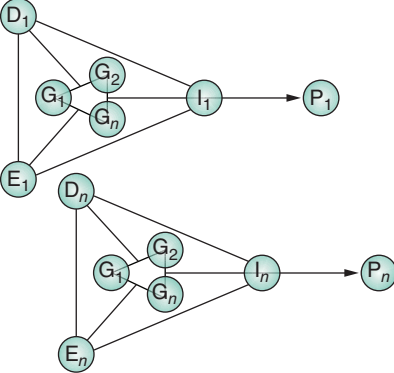
Example: Hypertrophic cardiomyopathy

**Polygenic disorder:
Single phenotype**

Example: Essential hypertension

**Polygenic disorder:
Multiple phenotypes**

Example: Ischemic heart disease

**Environmental disorder**

Example: Subacute bacterial endocarditis

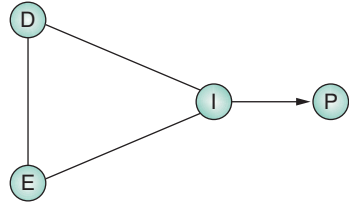


FIGURE 87e-2 Examples of modular representations of human disease. D, secondary human disease genome or proteome; E, environmental determinants; G, primary human disease genome or proteome; I, intermediate phenotype; P, pathophenotype. (Reproduced with permission from J Loscalzo et al: *Molec Syst Biol* 3:124, 2007.)

As yet another potential consideration, one can argue that disease reflects the later-stage consequences of the predilection of an organ system to manifest a particular intermediate pathophenotype in response to injury. This paradigm reflects a reverse causality view in which a disease is defined as a tendency to heightened inflammation, thrombosis, or fibrosis after an injurious perturbation. Where the process is manifest (i.e., the organ in which it occurs) is less important than that it occurs (with the exception of the organ-specific pathophysiologic consequences that may require acute attention). For example, from this perspective, acute myocardial infarction (AMI) and its consequences are a reflection of thrombosis (in the coronary artery), inflammation (in the acutely injured myocardium), and fibrosis (at the site or sites of cardiomyocyte death). In effect, the major therapies for AMI address these intermediate pathophenotypes (e.g., antithrombotics, statins) rather than any organ-specific disease-determining process. This paradigm would argue for a systems-based analysis that would first identify the intermediate pathophenotypes to which a person is predisposed, then determine how and when to intervene to attenuate that adverse predisposition, and finally limit the likelihood that a major organ-specific event will occur. Evidence

for the validity of this approach is found in the work of Rzhetsky and colleagues, who reviewed 1.5 million patient records and 161 diseases and found that these disease phenotypes form a network of strong pairwise correlations. This result is consistent with the notion that underlying genetic predispositions to intermediate pathophenotypes form the predicate basis for conventionally defined end organ diseases.

Regardless of the specific nature of the systems pathobiologic approach used, these analyses will lead to a drastic revision of the way human disease is defined and treated, establishing the discipline of network medicine. This will be a lengthy and complicated process but ultimately will lead to better disease prevention and therapy and probably do so from an increasingly personalized perspective. The analysis of pathobiology from a systems-based perspective is likely to help define specific subsets of patients more likely to respond to particular interventions based on shared disease mechanisms. Although it is unlikely that the extreme of “individualized medicine” will ever be practical (or even desirable), complex diseases can be mechanistically subclassified and interventions may be tailored to those settings in which they are more likely to work.

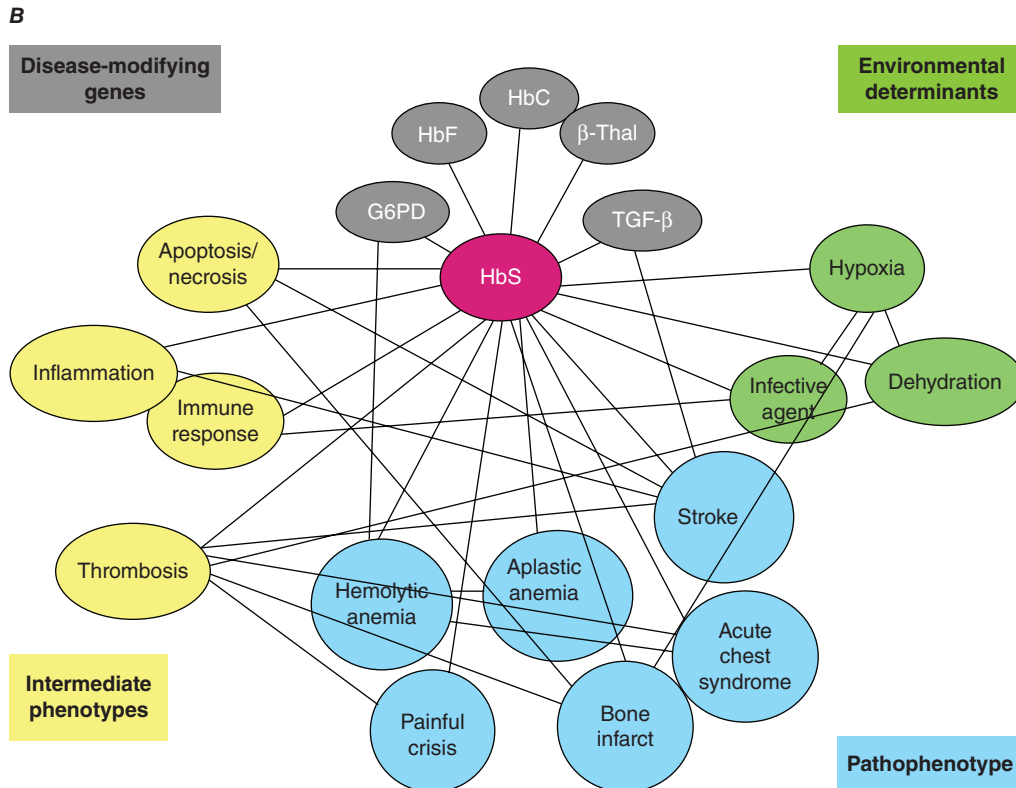
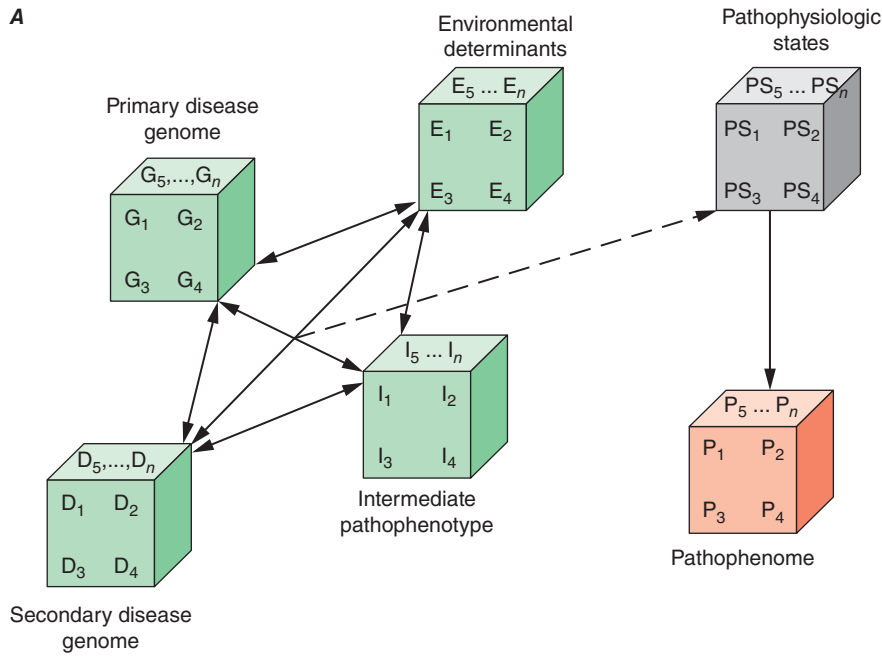


FIGURE 87e-3 **A.** Theoretical human disease network illustrating the relationships among genetic and environmental determinants of the pathophenotypes. Key: D, secondary disease genome or proteome; E, environmental determinants; G, primary disease genome or proteome; I, intermediate phenotype; PS, pathophysiologic states leading to P, pathophenotype. **B.** Example of this theoretical construct applied to sickle cell disease. Key: Red, primary molecular abnormality; gray, disease-modifying genes; yellow, intermediate phenotypes; green, environmental determinants; blue, pathophenotypes. (Reproduced with permission from J Loscalzo et al: *Molec Syst Biol* 3:124, 2007.)

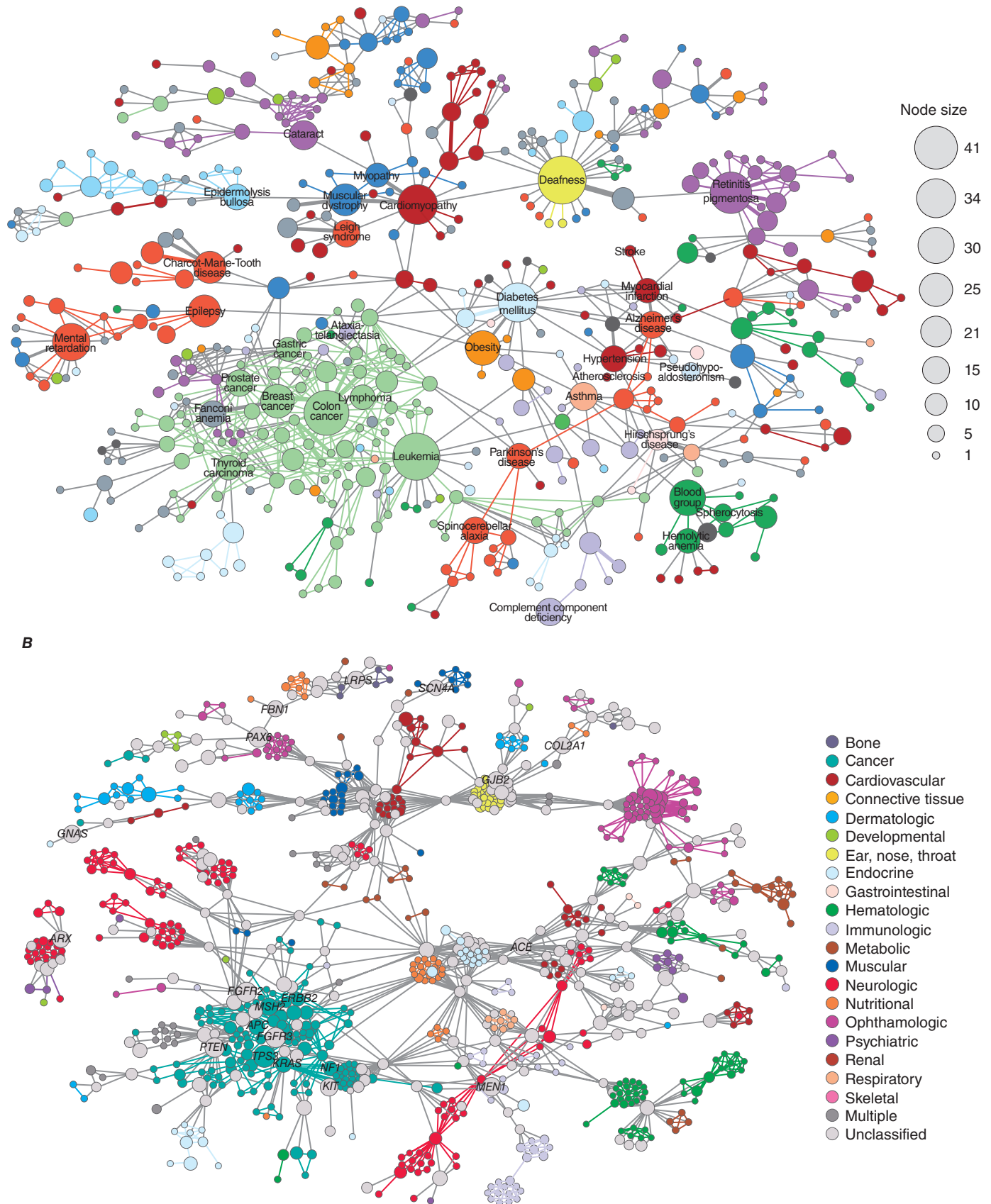


FIGURE 87e-4 **A.** Human disease network. Each node corresponds to a specific disorder colored by class (22 classes, shown in the key to **B**). The size of each node is proportional to the number of genes contributing to the disorder. Edges between disorders in the same disorder class are colored with the same (lighter) color, and edges connecting different disorder classes are colored gray, with the thickness of the edge proportional to the number of genes shared by the disorders connected by it. **B.** Disease gene network. Each node is a single gene, and any two genes are connected if implicated in the same disorder. In this network map, the size of each node is proportional to the number of specific disorders in which the gene is implicated. (Reproduced with permission from KI Goh et al: *Proc Natl Acad Sci USA* 104:8685, 2007.)

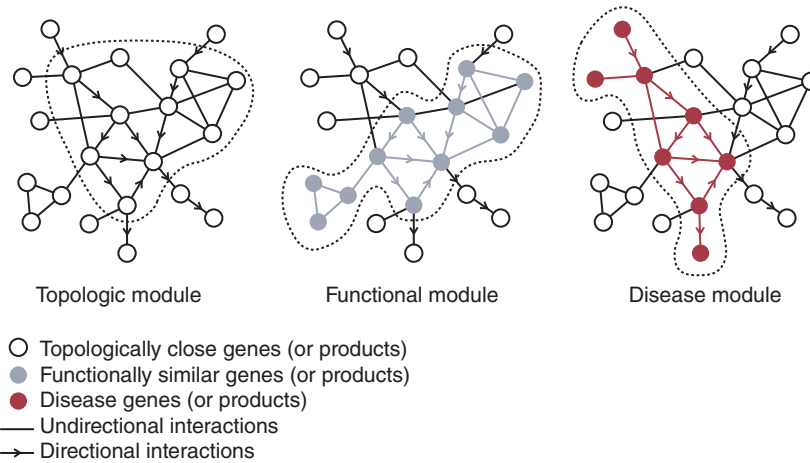


FIGURE 87e-5. The elements of the interactome. The interactome includes topologic modules (genes or gene products that are closely associated with one another through direct interactions), functional modules (genes or gene products that work together to define a pathway), and disease modules (genes or gene products that interact to yield a pathophenotype). (Reproduced with permission from AL Barabasi et al: *Nat Rev Genet* 12:56, 2011.)

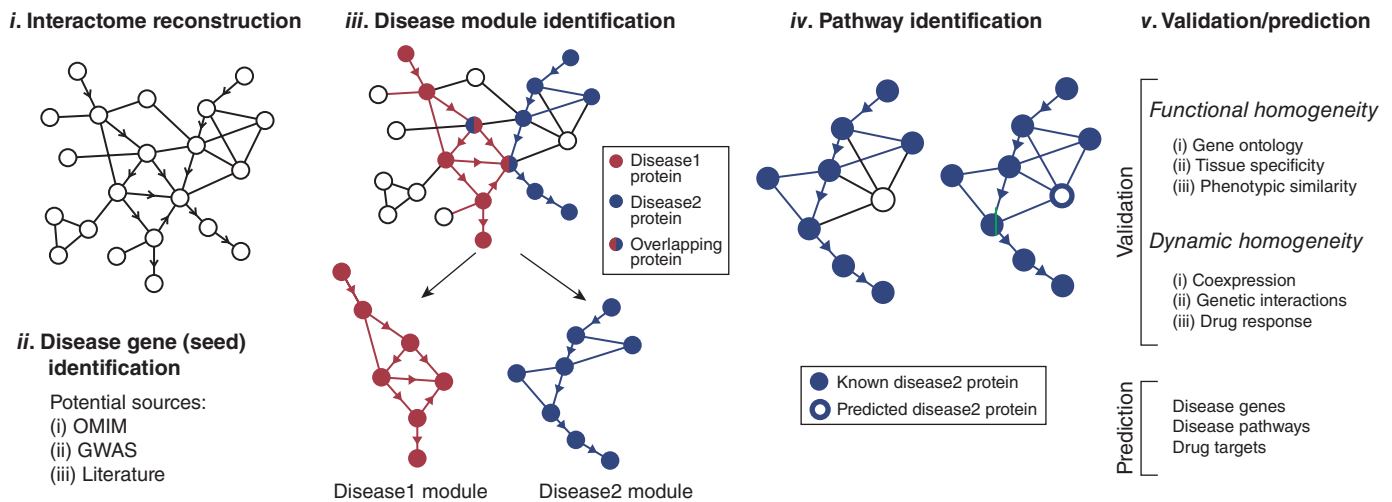


FIGURE 87e-6. Approaches to identifying disease modules in molecular networks. A strategy for defining disease modules involves (i) reconstructing the interactome; (ii) ascertaining potential seed (disease) genes from the curated literature, the Online Mendelian Inheritance in Man (OMIM) database, or genomic analyses (genome-wide association studies [GWAS] or transcriptional profiling); (iii) identifying the disease module using different modeling or statistical approaches; (iv) identifying pathways and the role of disease genes or modules in those pathways; and (v) disease module validation and prediction. (Reproduced with permission from AL Barabasi et al: *Nat Rev Genet* 12:56, 2011.)

This page intentionally left blank

88

Stem Cell Biology

Minoru S. H. Ko

Stem cell biology is a rapidly expanding field that explores the characteristics and possible clinical applications of a variety of stem cells that serve as the progenitors of more differentiated cell types. In addition to potential therapeutic applications (Chap. 90e), patient-derived stem cells can also be used as disease models and as a means of testing drug efficacy. Stem cells and their niche are a major focus of medical research because they play central roles in tissue and organ homeostasis and repair, which are important aspects of aging and disease.

IDENTIFICATION, ISOLATION, AND DERIVATION OF STEM CELLS

Resident Stem Cells The definition of stem cells remains elusive. Stem cells were originally postulated as *unspecified* or *undifferentiated* cells that provide a source of renewal of skin, intestine, and blood cells throughout life. These *resident stem cells* have been identified in a variety of organs (e.g., epithelia of the skin and digestive system, bone marrow, blood vessels, brain, skeletal muscle, liver, testis, and pancreas) based on their specific locations, morphology, and biochemical markers.

Isolated Stem Cells Unequivocal identification of stem cells requires their separation and purification, usually based on a combination of specific cell-surface markers. These *isolated stem cells* (e.g., hematopoietic stem [HS] cells) can be studied in detail and used in clinical applications, such as bone marrow transplantation (Chap. 89e). However, the lack of specific cell-surface markers for other types of stem cells has made it difficult to isolate them in large quantities. This challenge has been partially addressed in animal models by genetically marking different cell types with green-fluorescent protein driven by cell-specific promoters. Alternatively, putative stem cells have been isolated from a variety of tissues as side population (SP) cells using fluorescence-activated cell sorting after staining with the Hoechst 33342 dye.

Cultured Stem Cells It is desirable to culture and expand stem cells *in vitro* to obtain a sufficient quantity for analysis and potential therapeutic use. Although the derivation of stem cells *in vitro* has been a major obstacle in stem cell biology, the number and types of *cultured stem cells* have increased progressively (Table 88-1). Cultured stem cells derived from resident stem cells are often called *adult stem cells* or *somatic stem cells* to distinguish them from *embryonic stem* (ES) and *embryonic germ* (EG) cells. However, considering the existence of embryo-derived, tissue-specific stem cells (e.g., trophoblast stem [TS] cells) and the possible derivation of similar cells from an embryo/fetus (e.g., neural stem [NS] cells), it is more appropriate to use the term, *tissue stem cells*.

Successful derivation of cultured stem cells (both embryonic and tissue stem cells) often requires the identification of necessary growth factors and culture conditions, mimicking the microenvironment or *niche* of the resident stem cells. Recently, long-term maintenance of tissue stem cells *in vitro* is increasingly possible by growing them as three-dimensional (3D) organoids, which contain both stem cells and niche cells (Chap. 92e). For example, intestinal stem cells can now be cultured as “epithelial mini-guts” in the presence of R-spondin, epidermal growth factor (EGF), and noggin on Matrigel. Similarly, lung stem cells can be cultured as self-renewing “alveolospheres.” A growing list of cultured stem cells, although not comprehensive, is shown in Table 88-1. Please note that the establishment of cultured stem cells is often under dispute due to the difficulties in assessing the characteristics of these cells.

SELF-RENEWAL AND PROLIFERATION OF STEM CELLS

Symmetric and Asymmetric Cell Division The most widely accepted stem cell definition is a cell with a unique capacity to produce unaltered daughter cells (*self-renewal*) and to generate specialized cell types (*potency*). Self-renewal can be achieved in two ways. *Asymmetric cell division* produces one daughter cell that is identical to the parental cell and one daughter cell that is different from the parental cell and is a progenitor or differentiated cell. Asymmetric cell division does not increase the number of stem cells. *Symmetric cell division* produces two

TABLE 88-1 EXAMPLES OF CULTURED STEM CELLS

Name	Source
Embryonic stem cells (ES, ESC)	Blastocysts or immunosurgically isolated inner cell mass (ICM) from blastocysts
Embryonic germ cells (EG, EGC)	Primordial germ cells (PGCs) from embryos at E8.5–E12.5 (m); gonadal tissues from 5–11 week postfertilization embryo/fetus (h)
Trophoblast stem cells (TS, TSC)	Trophectoderm of E3.5 blastocysts, extraembryonic ectoderm of E6.5 embryos, and chorionic ectoderm of E7.5 embryos (m)
Extraembryonic endoderm cells (XEN)	ICM from blastocysts (m)
Embryonal carcinoma cells (EC)	Teratocarcinoma—a type of cancer that develops in the testes and ovaries (m, h)
Mesenchymal stem cells (MS, MSC)	Bone marrow, muscle, adipose tissue, peripheral blood, and umbilical cord blood (m, h)
Multipotent adult stem cells (MAPC)	Bone marrow mononuclear cells (m, h); postnatal muscle and brain (m)
Spermatogonial stem cells (SS, SSC)	Newborn testis (m)
Germline stem cells (GS, GSC)	Neonatal testis (m)
Multipotent adult germline stem cells (maGSC)	Adult testis (m)
Neural stem cells (NS, NSC)	Fetal and adult brain (subventricular zone, ventricular zone, and hippocampus) (m, h)
Unrestricted somatic stem cells (USSC)	Mononuclear fraction of cord blood (h)
Epistem cells (EpiSC)	Early postimplantation epiblast (m)
Induced pluripotent stem cells (iPS, iPSC)	Variety of terminally differentiated cells and tissue stem cells (m, h)
Multilineage-differentiating stress-enduring (MUSE) cells	Adult human bone marrow stromal cells and dermal fibroblasts
Urine-derived stem cells (USC)	Urine (h)
Lung stem cells	Lung (m, h)
Amniotic fluid-derived stem (AFS) cells	Amniotic fluid (m, h)
Umbilical cord blood stem cells	Umbilical cord (h)
Adipose stem cells (AST)	Fat (m, h)
Cardiac stem cells	Heart (m, h)
Renal stem cells	Renal papilla (m, h)
Crypt stem cells	Intestine (m, h)
Colon stem cells (CoSC)	Colon (m, h)
Hepatic stem cells	Liver (m, h)
Dental pulp stem cells (DPSC)	Dental pulp (m, h)
Hair follicle stem cells	Hair (m, h)

Abbreviations: h, human; m, mouse.

identical daughter cells. For stem cells to proliferate in vitro, they must divide symmetrically.

Unlimited Expansion In Vitro Resident stem cells are often quiescent and divide infrequently. However, once the stem cells are successfully cultured in vitro, they often acquire the capacity to divide continuously and the ability to proliferate beyond the normal passage limit typical of primary cultured cells (sometimes called *immortality*). These features are primarily seen in ES cells but have also been demonstrated for tissue stem cells, such as NS cells and mesenchymal stem (MS) cells, thereby enhancing the potential of these cells for therapeutic use (Table 88-1).

Stability of Genotype and Phenotype The capacity to actively proliferate is often associated with the accumulation of chromosomal abnormalities and mutations. Mouse ES cells appear to be an exception to this rule and tend to maintain their euploid karyotype and genome integrity. By contrast, human ES cells appear to be more susceptible to mutations after long-term culture. However, it is also important to note that even euploid mouse ES cells can form teratomas when injected into immunosuppressed animals, raising concerns about the possible formation of tumors after transplanting actively dividing stem cells.

POTENCY AND DIFFERENTIATION OF STEM CELLS

Developmental Potency The term *potency* is used to indicate a cell's ability to differentiate into multiple specialized cell types. The current lack of knowledge about the molecular nature of potency requires the experimental manipulation of stem cells to demonstrate their potency. For example, in vivo testing can be done by injecting stem cells into mouse blastocysts or immunosuppressed adult mice and determining how many different cell types are formed from the injected cells. However, these in vivo assays are not applicable to human stem cells. In vitro testing can be performed by differentiating cells in various culture conditions to determine how many different cell types are formed from the cells. The formal test of self-renewal and potency is performed by demonstrating that a single cell possesses such abilities in vitro (*clonality*). Cultured stem cells are tentatively grouped according to their potency (Fig. 88-1). Only some examples are shown, because many cultured stem cells, especially human cells, lack definitive information about their developmental potency.

From Totipotency to Unipotency *Totipotent cells* can form an entire organism autonomously. Only a fertilized egg (zygote) possesses this feature. *Pluripotent cells* (e.g., ES cells) can form almost all of the body's cell lineages (endoderm, mesoderm, and ectoderm), including germ cells. *Multipotent cells* (e.g., HS cells) can form multiple cell lineages

but cannot form all of the body's cell lineages. *Oligopotent cells* (e.g., NS cells) can form more than one cell lineage but are more restricted than multipotent cells. Oligopotent cells are sometimes called *progenitor cells* or *precursor cells*; however, these terms are often more strictly used to define partially differentiated or lineage-committed cells (e.g., myeloid progenitor cells) that can divide into different cell types but lack self-renewing capacity. *Unipotent cells* or *monopotent cells* (e.g., spermatogonial stem [SS] cells) can form a single differentiated cell lineage.

Nuclear Reprogramming Development naturally progresses from totipotent fertilized eggs to pluripotent epiblast cells to multipotent cells and, finally, to terminally differentiated cells. According to Waddington's epigenetic landscape, this is analogous to a ball moving down a slope. The reversal of the terminally differentiated cells to totipotent or pluripotent cells (called *nuclear reprogramming*) can thus be seen as an uphill gradient. Nuclear reprogramming has been achieved using *nuclear transplantation*, or *nuclear transfer* (NT), procedures (often called "cloning"), where the nucleus of a differentiated cell is transferred into an enucleated oocyte. Although this is an error-prone procedure with a very low success rate, live animals have been produced using adult somatic cells as donors in sheep, mice, and other mammals. In mice, it has been demonstrated that ES cells derived from blastocysts made by somatic cell NT are indistinguishable from normal ES cells. NT can potentially be used to produce *patient-specific ES cells* carrying a genome identical to that of the patient, although such strategies have not been pursued due to ethical issues and technical challenges. Recent success in generating human ES cells by NT has rekindled an interest in this area; however, the limited supply of human oocytes will still be a major problem for clinical applications of NT.

An alternative approach that has become a method of choice is the direct conversion of terminally differentiated cells into ES-like cells (called induced pluripotent stem [iPS] cells) by overexpressing a combination of key transcription factors (TFs). The original method was to infect mouse embryonic fibroblast cells with retrovirus vectors carrying four TFs [*Pou5f1* (*Oct4*), *Sox2*, *Klf4*, and *Myc*] and to identify rare ES-like cells in culture. This approach was soon adapted to human cells, followed by a more refined procedure (e.g., the use of fewer TFs, different cell types, and different gene-delivery methods). Because a clinical trial using iPS cells is imminent, the safety of iPS-based therapy is a major concern and a variety of measures are being taken to ensure the safety. For example, it has now become a standard to use footprint-free methods such as an episomal vector, Sendai virus vector, and synthetic mRNAs to deliver reprogramming factors into cells, resulting in the production of patient-specific iPS cells with minimal alteration of their genetic makeup. In addition to cell replacement therapy, disease-specific iPS cells are expected to play a role in modeling human disease in vitro and in screening drugs for personalized medicine.

It has also become possible to convert one type of terminally differentiated cell (e.g., fibroblast cell) into another type of terminally differentiated cell (e.g., cardiac muscle, neuron, or hepatocyte) by overexpressing specific sets of TFs (called direct reprogramming). Direct reprogramming can bypass the step of making iPS cells, possibly providing the safer route to desired cell types for therapy; however, the technology is currently limited by its low efficiency.

Stem Cell Plasticity, Transdifferentiation, and Facultative Stem Cells The prevailing paradigm in developmental biology is that once cells are differentiated, their phenotypes are stable. However, more recent studies show that tissue stem cells, which have traditionally been thought to be lineage-committed multipotent cells, possess the capacity to differentiate into cell types outside their lineage restrictions (called *transdifferentiation*). For example, HS cells may be converted into neurons as well as germ cells. This feature may provide a means to use tissue stem cells derived directly from a patient for therapeutic purposes, thereby eliminating the need to use embryonic stem cells or elaborate procedures such as nuclear reprogramming of a patient's

Source Potency	Preimplantation	Embryonic, fetal	Postnatal	Adult
Totipotent	Zygote ^{m,h}			
Pluripotent	ES ^{m,h}	EG ^{m,h} EpiSC ^m	GS ^m iPS ^{m,h} USSC ^h	EC ^{m,h} iPS ^{m,h} maGSC ^m
Multipotent				MS ^{m,h}
Oligopotent	TS ^m			NS ^{m,h}
Unipotent	XEN ^m			SSC ^m
Terminally differentiated cells				

FIGURE 88-1 Potency and source developmental stage of cultured stem cells. For abbreviations of stem cells, see Table 88-1. Note that stem cells are often abbreviated with or without "cells," e.g., ES cells or ESCs for embryonic stem cells. h, human; m, mouse.

somatic cells. However, more strict criteria and rigorous validation are required to establish tissue stem cell plasticity. For example, observations of transdifferentiation may reflect cell fusion, contamination with progenitor cells from other cell lineages, or persistence of pluripotent embryonic cells in adult organs. Therefore, the assignment of potency to each cultured stem cell in Fig. 88-1 should be considered with caution. Whether transdifferentiation exists and can be used for therapeutic purposes remains to be determined conclusively. A similar, but distinct, concept is the facultative stem cell, which is defined as a unipotent cell or a terminally differentiated cell that can function as a stem cell upon tissue injury. The presence of such cells has been proposed for some organs such as liver, intestine, pancreas, and testis, but is still debated.

Directed Differentiation of Stem Cells Pluripotent stem cells (e.g., ES and iPS cells) can differentiate into multiple cell types, but in culture, they normally differentiate into heterogeneous cell populations in a stochastic manner. However, for therapeutic uses, it is desirable to direct stem cells into specific cell types (e.g., insulin-secreting beta cells). This is an active area of stem cell research, and protocols are being developed to achieve this goal. In any of these directed cell differentiation systems, the cell phenotype must be evaluated critically. Alternatively, the heterogeneity of the cell population derived from pluripotent stem cells can be actively exploited, as different types of cells interact with each other in culture and further enhance their own differentiation. In some instances, e.g., optic cup, self-organizing tissue morphogenesis has been demonstrated in 3D culture.

MOLECULAR CHARACTERIZATION OF STEM CELLS

Genomics and Proteomics In addition to standard molecular biological approaches, high-throughput genomics and proteomics have been extensively applied to the analysis of stem cells. For example, DNA microarray analyses have revealed the expression levels of essentially all genes and identified specific markers for some stem cells. Chromatin immunoprecipitation coupled with next-generation sequencing technologies, capable of producing billions of sequence reads in a single run, has revealed chromatin modifications (“epigenetic marks”) relevant to stem cell properties. Similarly, the protein profiles of stem cells have been assessed by using mass spectrometry. These methods are beginning to provide a novel means to characterize and classify various stem cells and the molecular mechanisms that give them their unique characteristics.

ES Cell Regulation It is important to identify genes involved in the regulation of stem cell function and to examine the effects of altered gene expression on ES and other stem cells. For example, core networks of TFs such as *Pou5f1* (*Oct4*), *Nanog*, and *Sox2*, govern key gene regulatory pathways/networks for the maintenance of self-renewal and pluripotency of mouse and human ES cells. These TF networks are modulated by specific external factors through signal transduction pathways, such as leukemia inhibitory factor (*Lif*)/*Stat3*, mitogen-activated protein kinase 1/3 (*Mapk1/3*), the transforming growth factor β (TGF β) superfamily, and *Wnt*/glycogen synthase kinase 3 beta (*Gsk3b*). Inhibitors of *Mapk1/3* and *Gsk3b* signaling enhance the derivation of ES cells and help maintain ES cells in full pluripotency (“ground” or “naive state”). Recent data also indicate that 20–25 nucleotide RNAs, called microRNAs (miRNAs), play an important role in regulating stem cell function by repressing the translation of their target genes. For example, it has been shown that miR-21 regulates cell cycle progression in ES cells and miR-128 prevents the differentiation of hematopoietic progenitor cells. These types of analyses should provide molecular clues about the function of stem cells and lead to a more effective means to manipulate stem cells for future therapeutic use.

All of the cell types in the peripheral blood and some cells in every tissue of the body are derived from hematopoietic (*hemo*: blood; *poiesis*: creation) stem cells. If the hematopoietic stem cell is damaged and can no longer function (e.g., due to a nuclear accident), a person would survive 2–4 weeks in the absence of extraordinary support measures. With the clinical use of hematopoietic stem cells, tens of thousands of lives are saved each year (Chap. 139e). Stem cells produce hundreds of billions of blood cells daily from a stem cell pool that is estimated to be only in the tens of thousands. How stem cells do this, how they persist for many decades despite the production demands, and how they may be better used in clinical care are important issues in medicine.

The study of blood cell production has become a paradigm for how other tissues may be organized and regulated. Basic research in hematopoiesis includes defining stepwise molecular changes accompanying functional changes in maturing cells, aggregating cells into functional subgroups, and demonstrating hematopoietic stem cell regulation by a specialized microenvironment; these concepts are worked out in hematology, but they offer models for other tissues. Moreover, these concepts may not be restricted to normal tissue function but extend to malignancy. Stem cells are rare cells among a heterogeneous population of cell types, and their behavior is assessed mainly in experimental animal models involving reconstitution of hematopoiesis. Thus, much of what we know about stem cells is imprecise and based on inferences from genetically manipulated animals.

CARDINAL FUNCTIONS OF HEMATOPOIETIC STEM CELLS

All stem cell types have two cardinal functions: self-renewal and differentiation (Fig. 89e-1). Stem cells exist to generate, maintain, and repair tissues. They function successfully if they can replace a wide variety of shorter-lived mature cells over prolonged periods. The process of self-renewal (see below) assures that a stem cell population can be sustained over time. Without self-renewal, the stem cell pool would become exhausted and tissue maintenance would not be possible. The process of differentiation leads to production of the effectors of tissue function: mature cells. Without proper differentiation, the integrity of

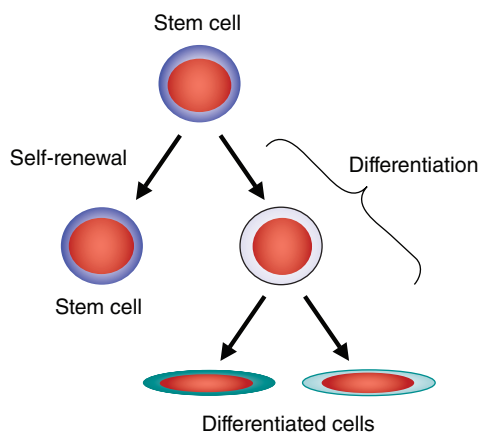


FIGURE 89e-1 Signature characteristics of the stem cell. Stem cells have two essential features: the capacity to differentiate into a variety of mature cell types and the capacity for self-renewal. Intrinsic factors associated with self-renewal include expression of *Bmi-1*, *Gfi-1*, *PTEN*, *STAT5*, *Tel/Atv6*, *p21*, *p18*, *MCL-1*, *Mel-18*, *RAE28*, and *HoxB4*. Extrinsic signals for self-renewal include Notch, Wnt, SHH, and Tie2/Ang-1. Based mainly on murine studies, hematopoietic stem cells express the following cell surface molecules: CD34, Thy-1 (CD90), c-Kit receptor (CD117), CD133, CD164, and c-Mpl (CD110, also known as the thrombopoietin receptor).

tissue function would be compromised and organ failure or neoplasia would ensue.

In the blood, mature cells have variable average life spans, ranging from 7 h for mature neutrophils to a few months for red blood cells to many years for memory lymphocytes. However, the stem cell pool is the central, durable source of all blood and immune cells, maintaining a capacity to produce a broad range of cells from a single cell source, yet keeping itself vigorous over decades of life. As an individual stem cell divides, it has the capacity to accomplish one of three division outcomes: two stem cells, two cells destined for differentiation, or one stem cell and one differentiating cell. The former two outcomes are the result of symmetric cell division, whereas the latter indicates a different outcome for the two daughter cells—an event termed *asymmetric cell division*. The relative balance for these types of outcomes may change during development and under particular kinds of demands on the stem cell pool.

DEVELOPMENTAL BIOLOGY OF HEMATOPOIETIC STEM CELLS

During development, blood cells are produced at different sites. Initially, the yolk sac provides oxygen-carrying red blood cells, and then the placenta and several sites of intraembryonic blood cell production become involved. These intraembryonic sites engage in sequential order, moving from the genital ridge at a site where the aorta, gonadal tissue, and mesonephros are emerging to the fetal liver and then, in the second trimester, to the bone marrow and spleen. As the location of stem cells changes, the cells they produce also change. The yolk sac provides red cells expressing embryonic hemoglobins while intraembryonic sites of hematopoiesis generate red cells, platelets, and the cells of innate immunity. The production of the cells of adaptive immunity occurs when the bone marrow is colonized and the thymus forms. Stem cell proliferation remains high, even in the bone marrow, until shortly after birth, when it appears to dramatically decline. The cells in the bone marrow are thought to arrive by the bloodborne transit of cells from the fetal liver after calcification of the long bones has begun. The presence of stem cells in the circulation is not unique to a time window in development; however, hematopoietic stem cells appear to circulate throughout life. The time that cells spend freely circulating appears to be brief (measured in minutes in the mouse), but the cells that do circulate are functional and can be used for transplantation. The number of stem cells that circulate can be increased in a number of ways to facilitate harvest and transfer to the same or a different host.

MOBILITY OF HEMATOPOIETIC STEM CELLS

Cells entering and exiting the bone marrow do so through a series of molecular interactions. Circulating stem cells (through CD162 and CD44) engage the lectins (carbohydrate binding proteins) P- and E-selectin on the endothelial surface to slow the movement of the cells to a rolling phenotype. Stem cell integrins are then activated and accomplish firm adhesion between the stem cell and vessel wall, with a particularly important role for stem cell VCAM-1 engaging endothelial VLA-4. The chemokine CXCL12 (SDF1) interacting with stem cell CXCR4 receptors and ionic calcium interacting with the calcium sensing receptor appear to be important in the process of stem cells getting from the circulation to where they engraft in the bone marrow. This is particularly true in the developmental move from fetal liver to bone marrow.

However, the role for CXCR4 in adults appears to be more related to retention of stem cells in the bone marrow rather than the process of getting them there. Interrupting that retention process through either specific molecular blockers of the CXCR4/CXCL12 interaction, cleavage of CXCL12, or downregulation of the CXCR4 receptor can all result in the release of stem cells into the circulation. This process is an increasingly important aspect of recovering stem cells for therapeutic use as it has permitted the harvesting process to be done by leukapheresis rather than bone marrow punctures in the operating room. Granulocyte colony-stimulating factor and plerixafor, a macrocyclic compound that can block CXCR4, are both used clinically to mobilize marrow hematopoietic stem cells for transplant. Refining our

knowledge of how stem cells get into and out of the bone marrow may improve our ability to obtain stem cells and make them more efficient at finding their way to the specific sites for blood cell production, the so-called stem cell niche.

HEMATOPOIETIC STEM CELL MICROENVIRONMENT

The concept of a specialized microenvironment, or stem cell niche, was first proposed to explain why cells derived from the bone marrow of one animal could be used in transplantation and again be found in the bone marrow of the recipient. This niche is more than just a housing site for stem cells, however. It is an anatomic location where regulatory signals are provided that allow the stem cells to thrive, to expand if needed, and to provide varying amounts of descendant daughter cells. In addition, unregulated growth of stem cells may be problematic based on their undifferentiated state and self-renewal capacity. Thus, the niche must also regulate the number of stem cells produced. In this manner, the niche has the dual function of serving as a site of nurture but imposing limits for stem cells: in effect, acting as both a nutritive and constraining home.

The niche for blood stem cells changes with each of the sites of blood production during development, but for most of human life it is located in the bone marrow. Within the bone marrow, the perivascular space particularly in regions of trabecular bone serves as a niche. The mesenchymal and endothelial cells of the marrow microvessels produce kit ligand and CXCL12, both known to be important for hematopoietic stem cells. Other cell types, such as sympathetic neurons, nonmyelinating Schwann cells, macrophages, osteoclasts, and osteoblasts, have been shown to regulate stem cells, but it is unclear whether their effects are direct or indirect. Extracellular matrix proteins like osteopontin also affect stem cell function. The endosteal region is particularly important for transplanted cells, suggesting that there may be distinctive features of that region that are yet to be defined that are important mediators of stem cell engraftment. The functioning of the niche as a supportive context for stem cells is of obvious importance for maintaining hematopoiesis and in transplantation. An active area of study involves determining whether the niche is altered in disease and whether drugs can modify niche function to improve transplantation or normal stem cell function in hematologic disease.

EXCESS CAPACITY OF HEMATOPOIETIC STEM CELLS

In the absence of disease, one never runs out of hematopoietic stem cells. Indeed, serial transplantation studies in mice suggest that sufficient stem cells are present to reconstitute several animals in succession, with each animal having normal blood cell production. The fact that allogeneic stem cell transplant recipients also never run out of blood cells in their life span, which can extend for decades, argues that even the limiting numbers of stem cells provided to them are sufficient. How stem cells respond to different conditions to increase or decrease their mature cell production remains poorly understood. Clearly, negative feedback mechanisms affect the level of production of most of the cells, leading to the normal tightly regulated blood cell counts. However, many of the regulatory mechanisms that govern production of more mature progenitor cells do not apply or apply differently to stem cells. Similarly, most of the molecules shown to be able to change the size of the stem cell pool have little effect on more mature blood cells. For example, the growth factor erythropoietin, which stimulates red blood cell production from more mature precursor cells, has no effect on stem cells. Similarly, granulocyte colony-stimulating factor drives the rapid proliferation of granulocyte precursors but has little or no effect on the cell cycling of stem cells. Rather, it changes the location of stem cells by indirect means, altering molecules such as CXCL12 that tether stem cells to their niche. Molecules shown to be important for altering the proliferation, self-renewal, or survival of stem cells, such as cyclin-dependent kinase inhibitors, transcription factors like *Bmi-1*, or microRNA-processing enzymes like Dicer, have little or different effects on progenitor cells. Hematopoietic stem cells have governing mechanisms that are distinct from the cells they generate.

HEMATOPOIETIC STEM CELL DIFFERENTIATION

Hematopoietic stem cells sit at the base of a branching hierarchy of cells culminating in the many mature cell types that compose the blood and immune system (Fig. 89e-2). The maturation steps leading to terminally differentiated and functional blood cells take place both as a consequence of intrinsic changes in gene expression and niche-directed and cytokine-directed changes in the cells. Our knowledge of the details remains incomplete. As stem cells mature to progenitors, precursors, and, finally, mature effector cells, they undergo a series of functional changes. These include the obvious acquisition of functions defining mature blood cells, such as phagocytic capacity or hemoglobin synthesis. They also include the progressive loss of plasticity (i.e., the ability to become other cell types). For example, the myeloid progenitor can make all cells in the myeloid series but none in the lymphoid series. As common myeloid progenitors mature, they become precursors for either monocytes and granulocytes or erythrocytes and megakaryocytes, but not both. Some amount of reversibility of this process may exist early in the differentiation cascade, but that is lost beyond a distinct stage in normal physiologic conditions. With genetic interventions, however, blood cells, like other somatic cells, can be reprogrammed to become a variety of cell types.

As cells differentiate, they may also lose proliferative capacity (Fig. 89e-3). Mature granulocytes are incapable of proliferation and only increase in number by increased production from precursors. The exceptions to the rule are some resident macrophages, which appear capable of proliferation, and lymphoid cells. Lymphoid cells retain the capacity to proliferate but have linked their proliferation to the recognition of particular proteins or peptides by specific antigen receptors on their surface. Like many tissues with short-lived mature cells such as the skin and intestine, blood cell proliferation is largely accomplished by a more immature progenitor population. In general, cells within the highly proliferative progenitor cell compartment are also relatively short-lived, making their way through the differentiation process in a defined molecular program involving the sequential activation of particular sets of genes. For any particular cell type, the differentiation program is difficult to speed up. The time it takes for hematopoietic progenitors to become mature cells is ~10–14 days in humans, evident clinically by the interval between cytotoxic chemotherapy and blood count recovery in patients.

Although hematopoietic stem cells are generally thought to have the capacity to form all cells of the blood, it is becoming clear that individual stem cells may not be equal in their differentiation potential. That is, some stem cells are “biased” to become mature cells of a particular type. In addition, the general concept of cells having a binary choice of lymphoid or myeloid differentiation is not entirely accurate. A cell population with limited myeloid (monocyte and granulocyte) and lymphoid potential is now added to the commitment steps stem cells may undergo.

SELF-RENEWAL

The hematopoietic stem cell must balance its three potential fates: apoptosis, self-renewal, and differentiation. The proliferation of cells is generally not associated with the ability to undergo a self-renewing division except among memory T and B cells and among stem cells. Self-renewal capacity gives way to differentiation as the only option after cell division when cells leave the stem cell compartment, until they have the opportunity to become memory lymphocytes. In addition to this self-renewing capacity, stem cells have an additional feature characterizing their proliferation machinery. Stem cells in many mature adult tissues may be heterogeneous with some being deeply quiescent, serving as a deep reserve, whereas others are more proliferative and replenish the short-lived progenitor population. In the hematopoietic system, stem cells are generally cytokine-resistant, remaining dormant even when cytokines drive bone marrow progenitors to proliferation rates measured in hours. Stem cells, in contrast, are thought to divide at far longer intervals, measured in months to years, for the most quiescent cells. This quiescence is difficult to overcome in vitro, limiting the ability to effectively expand human hematopoietic stem

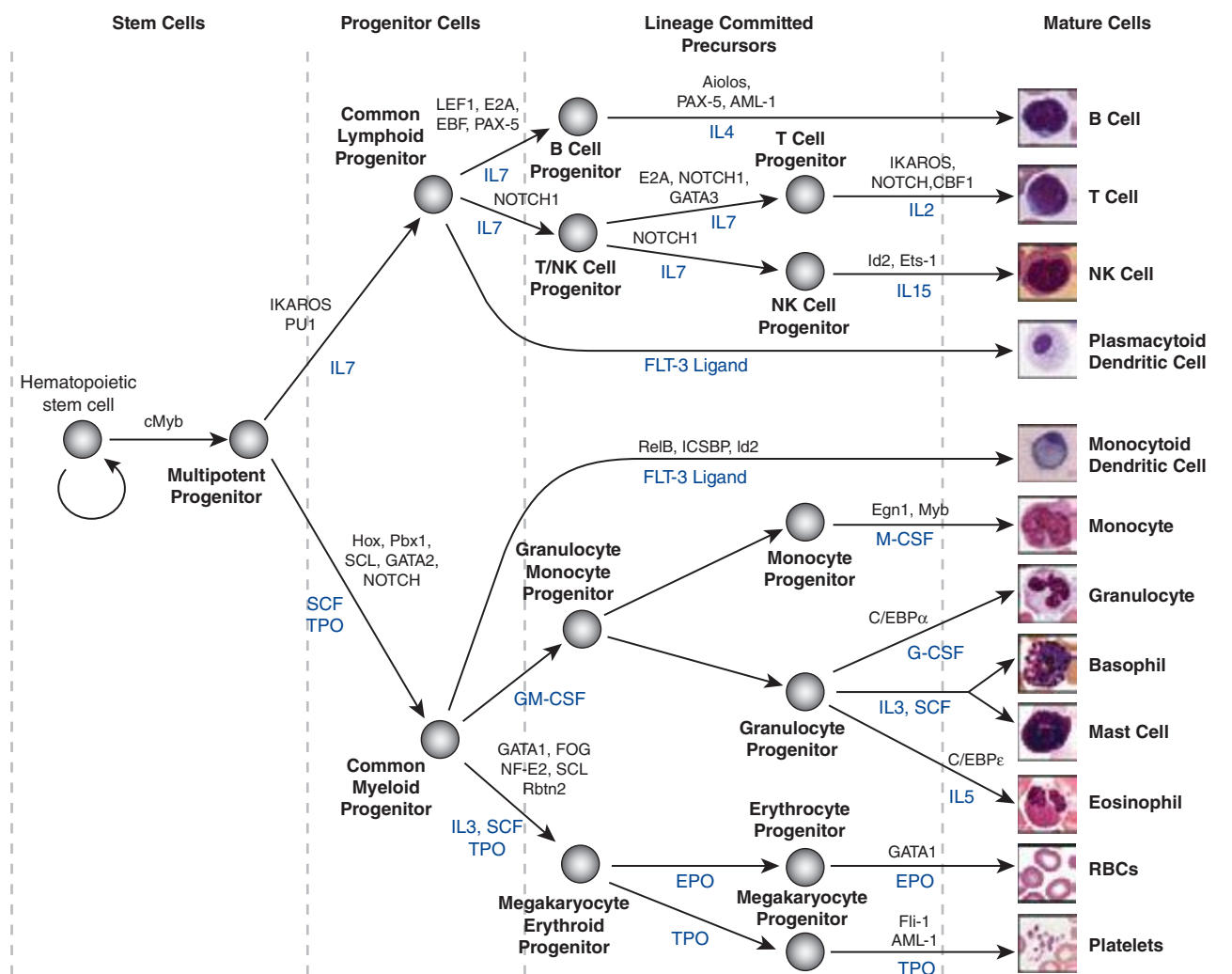


FIGURE 89e-2 Hierarchy of hematopoietic differentiation. *Stem cells* are multipotent cells that are the source of all descendant cells and have the capacity to provide either long-term (measured in years) or short-term (measured in months) cell production. *Progenitor cells* have a more limited spectrum of cells they can produce and are generally a short-lived, highly proliferative population also known as transient amplifying cells. *Precursor cells* are cells committed to a single blood cell lineage but with a continued ability to proliferate; they do not have all the features of a fully mature cell. *Mature cells* are the terminally differentiated product of the differentiation process and are the effector cells of specific activities of the blood and immune system. Progress through the pathways is mediated by alterations in gene expression. The regulation of the differentiation by soluble factors and cell-cell communications within the bone marrow niche are still being defined. The transcription factors that characterize particular cell transitions are illustrated on the *arrows*; the soluble factors that contribute to the differentiation process are in *blue*. This picture is a simplification of the process. Active research is revealing multiple discrete cell types in the maturation of B cells and T cells and has identified cells that are biased toward one lineage or another (rather than uncommitted) in their differentiation. EPO, erythropoietin; RBC, red blood cell; SCF, stem cell factor; TPO, thrombopoietin.

cells. The process may be controlled by particularly high levels of cyclin-dependent kinase inhibitors like p57 or CDKN1c that restrict entry of stem cells into the cell cycle, blocking the G_1 -S transition. Exogenous signals from the niche also appear to enforce quiescence, including the activation of the tyrosine kinase receptor Tie2 on stem cells by angiopoietin 1 on niche cells.

The regulation of stem cell proliferation also appears to change with age. In mice, the cyclin-dependent kinase inhibitor p16INK4a accumulates in stem cells in older animals and is associated with a change in five different stem cell functions, including cell cycling. Lowering expression of p16INK4a in older animals improves stem cell cycling and capacity to reconstitute hematopoiesis in adoptive hosts, making them similar to younger animals. Mature cell numbers are unaffected. Therefore, molecular events governing the specific functions of stem cells are being gradually made clear and offer the potential of new approaches to changing stem cell function for therapy. One critical stem cell function that remains poorly defined is the molecular regulation of self-renewal.

For medicine, self-renewal is perhaps the most important function of stem cells because it is critical in regulating the number of stem cells. Stem cell number is a key limiting parameter for both autologous and allogeneic stem cell transplantation. Were we to have the ability to use fewer stem cells or expand limited numbers of stem cells *ex vivo*, it might be possible to reduce the morbidity and expense of stem cell harvests and enable use of other stem cell sources. Specifically, umbilical cord blood is a rich source of stem cells. However, the volume of cord blood units is extremely small, and therefore, the total number of hematopoietic stem cells that can be obtained in any single cord blood unit is generally only sufficient to transplant an individual of <40 kg. This limitation restricts what would otherwise be an extremely promising source of stem cells. Two features of cord blood stem cells are particularly important. (1) They are derived from a diversity of individuals that far exceeds the adult donor pool and therefore can overcome the majority of immunologic cross-matching obstacles. (2) Cord blood stem cells have a large number of T cells associated with them, but (paradoxically) they appear to be associated with a lower

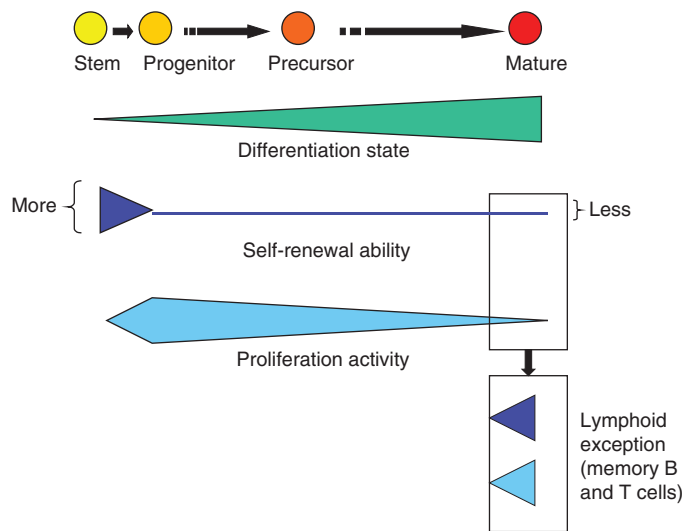


FIGURE 89e-3 Relative function of cells in the hematopoietic hierarchy. The boxes represent distinct functional features of cells in the myeloid (*upper box*) versus lymphoid (*lower box*) lineages.

incidence of graft-versus-host disease when compared with similarly mismatched stem cells from other sources. If stem cell expansion by self-renewal could be achieved, the number of cells available might be sufficient for use in larger adults. An alternative approach to this problem is to improve the efficiency of engraftment of donor stem cells. Graft engineering is exploring methods of adding cell components that may enhance engraftment. Furthermore, at least some data suggest that depletion of host NK (natural killer) cells may lower the number of stem cells necessary to reconstitute hematopoiesis.

Some limited understanding of self-renewal exists and, intriguingly, implicates gene products that are associated with the chromatin state, a high-order organization of chromosomal DNA that influences transcription. These include members of the polycomb family, a group of zinc finger-containing transcriptional regulators that interact with the chromatin structure, contributing to the accessibility of groups of genes for transcription. One member, *Bmi-1*, is important in enabling hematopoietic stem cell self-renewal through modification of cell cycle regulators such as the cyclin-dependent kinase inhibitors. In the absence of *Bmi-1* or of the transcriptional regulator, *Gfi-1*, hematopoietic stem cells decline in number and function. In contrast, dysregulation of *Bmi-1* has been associated with leukemia; it may promote leukemic stem cell self-renewal when it is overexpressed. Other transcription regulators have also been associated with self-renewal, particularly homeobox, or “hox,” genes. These transcription factors are named for their ability to govern large numbers of genes, including those determining body patterning in invertebrates. *HoxB4* is capable of inducing extensive self-renewal of stem cells through its DNA-binding motif. Other members of the hox family of genes have been noted to affect normal stem cells, but they are also associated with leukemia. External signals that may influence the relative self-renewal versus differentiation outcomes of stem cell cycling include specific Wnt ligands. Intracellular signal transducing intermediates are also implicated in regulating self-renewal. They include PTEN, an inhibitor of the AKT pathway, and STAT5, both of which are downstream of activated growth factor receptors and necessary for normal stem cell functions including self-renewal, at least in mouse models. The connections between these molecules remain to be defined, and their

role in physiologic regulation of stem cell self-renewal is still poorly understood.

CANCER IS SIMILAR TO AN ORGAN WITH SELF-RENEWING CAPACITY

The relationship of stem cells to cancer is an important evolving dimension of adult stem cell biology. Cancer may share principles of organization with normal tissues. Cancer cells are heterogeneous even within a given patient and may have a hierarchical organization of cells with a base of stem-like cells capable of the signature stem cell features: self-renewal and differentiation. These stem-like cells might be the basis for perpetuation of the tumor and represent a slowly dividing, rare population with distinct regulatory mechanisms, including a relationship with a specialized microenvironment. A subpopulation of self-renewing cells has been defined for some, but not all, cancers. A more sophisticated understanding of the stem cell organization of cancers may lead to improved strategies for developing new therapies for the many common and difficult-to-treat types of malignancies that have been relatively refractory to interventions aimed at dividing cells.

Does the concept of cancer stem cells provide insight into the cellular origin of cancer? The fact that some cells within a cancer have stem cell-like properties does not necessarily mean that the cancer arose in the stem cell itself. Rather, more mature cells could have acquired the self-renewal characteristics of stem cells. Any single genetic event is unlikely to be sufficient to enable full transformation of a normal cell to a frankly malignant one. Rather, cancer is a multistep process, and for the multiple steps to accumulate, the cell of origin must be able to persist for prolonged periods. It must also be able to generate large numbers of daughter cells. The normal stem cell has these properties and, by virtue of its having intrinsic self-renewal capability, may be more readily converted to a malignant phenotype. This hypothesis has been tested experimentally in the hematopoietic system. Taking advantage of the cell-surface markers that distinguish hematopoietic cells of varying maturity, stem cells, progenitors, precursors, and mature cells can be isolated. Powerful transforming gene constructs were placed in these cells, and it was found that the cell with the greatest potential to produce a malignancy was dependent on the transforming gene. In some cases, it was the stem cell, but in others, the progenitor cell functioned to initiate and perpetuate the cancer. This shows that cells can acquire stem cell-like properties in malignancy.

WHAT ELSE CAN HEMATOPOIETIC STEM CELLS DO?

Some experimental data have suggested that hematopoietic stem cells or other cells mobilized into the circulation by the same factors that mobilize hematopoietic stem cells are capable of playing a role in healing the vascular and tissue damage associated with stroke and myocardial infarction. These data are controversial, and the applicability of a stem cell approach to nonhematopoietic conditions remains experimental. However, reprogramming technology offers the potential for using the readily obtained hematopoietic stem cell as a source for cells with other capabilities.

The stem cell, therefore, represents a true dual-edged sword. It has tremendous healing capacity and is essential for life. Uncontrolled, it can threaten the life it maintains. Understanding how stem cells function, the signals that modify their behavior, and the tissue niches that modulate stem cell responses to injury and disease are critical for more effectively developing stem cell-based medicine. That aspect of medicine will include the use of the stem cells and the use of drugs to target stem cells to enhance repair of damaged tissues. It will also include the careful balance of interventions to control stem cells where they may be dysfunctional or malignant.

90e Applications of Stem Cell Biology in Clinical Medicine

John A. Kessler

Damage to an organ initiates a series of events that lead to the reconstruction of the damaged tissue, including proliferation, differentiation, and migration of various cell types; release of cytokines and chemokines; and remodeling of the extracellular matrix. Endogenous stem and progenitor cells are among the cell populations that are involved in these injury responses. In normal steady-state conditions, an equilibrium is maintained in which endogenous stem cells intrinsic to the tissue replenish dying cells. After tissue injury, stem cells in organs such as the liver and skin have a remarkable ability to regenerate the organ, whereas other stem cell populations, such as those in the heart and brain, have a much more limited capability for self-repair. In rare circumstances, circulating stem cells may contribute to regenerative responses by migrating into a tissue and differentiating into organ-specific cell types. The goal of stem cell therapies is to promote cell replacement in organs that are damaged beyond their ability to self-repair.

GENERAL STRATEGIES FOR STEM CELL REPLACEMENT

At least three different therapeutic concepts for cell replacement can be envisaged (Fig. 90e-1). One therapeutic approach involves direct administration of stem cells. The cells may be injected directly into the damaged organ, where they can differentiate into the desired cell type. Alternatively, stem cells may be injected systemically since they have the capacity to home in on damaged tissues by following gradients of cytokines and chemokines released by the diseased organ. A second approach involves transplantation of differentiated cells derived from stem cells. For example, pancreatic islet cells can be

generated from stem cells before transplantation into diabetic patients, and cardiomyocytes can be generated to treat ischemic heart disease. A third approach involves stimulation of endogenous stem cells to facilitate repair. This goal might be accomplished by administration of appropriate growth factors and drugs that amplify the number of endogenous stem/progenitor cells and/or direct them to differentiate into the desired cell types. Therapeutic stimulation of precursor cells is already a clinical reality in the hematopoietic system, where factors such as erythropoietin, granulocyte colony-stimulating factor, and granulocyte-macrophage colony-stimulating factor are used to increase production of specific blood elements. In addition to these strategies for cell replacement, a number of other approaches could involve stem cells for ex vivo or in situ generation of tissues, a process termed *tissue engineering* (Chap. 92e). Stem cells are also excellent candidates as vehicles for cellular gene therapy (Chap. 91e). Finally, transplanted stem cells may exert paracrine effects on damaged tissues without the differentiation and replacement of lost cells.

Stem cell transplantation is not a new concept but rather is already part of established medical practice. Hematopoietic stem cells (Chap. 89e) are responsible for the long-term repopulation of all blood elements in recipients of bone marrow transplants, and hematopoietic stem cell transplantation is the gold standard against which other stem cell transplantation therapies will be measured. Transplantation of differentiated cells is also a clinical reality, and donated organs and tissues are often used to replace damaged tissues. However, the need for transplantable tissues and organs far outweighs the available supply, and organ transplantation has limited potential for some tissues, such as the brain. Stem cells offer the possibility of a renewable source of replacement cells for virtually all organs.

SOURCES OF STEM CELLS FOR TISSUE REPAIR

A variety of different types of stem cells (Chap. 88) could be used in regenerative strategies, including embryonic stem (ES) cells, induced pluripotent stem (iPS) cells, umbilical-cord blood stem cells (USCs), organ-specific somatic stem cells (e.g., neural stem cells for treatment of the brain), and somatic stem cells that generate cell types specific for the target organ rather than the donor organ (e.g., bone marrow mesenchymal stem cells or CD34+ hematopoietic stem cells for cardiac repair). Although each cell type has potential advantages and disadvantages, there are a number of generic problems in developing any of these cell types into a useful and reliable clinical tool.

Embryonic Stem Cells Embryonic stem cells have the potential to generate all the cell types in the body; thus, in theory, there are no restrictions on the organs that could be regenerated. ES cells can self-renew endlessly, so that a single cell line with carefully characterized traits potentially could generate almost limitless numbers of cells. In the absence of moral or ethical constraints (see “Ethical Issues,” below), unused human blastocysts from fertility clinics could be used to derive new ES cell lines that are matched immunologically with potential transplant recipients. Alternatively, somatic cell nuclear transfer (“therapeutic cloning”) could be used to create ES cell lines that are genetically identical to those of the patient, although this endeavor has been technically refractory for human cells. However, human ES cells are difficult to culture and grow slowly. Techniques for differentiating them into specific cell types are just beginning

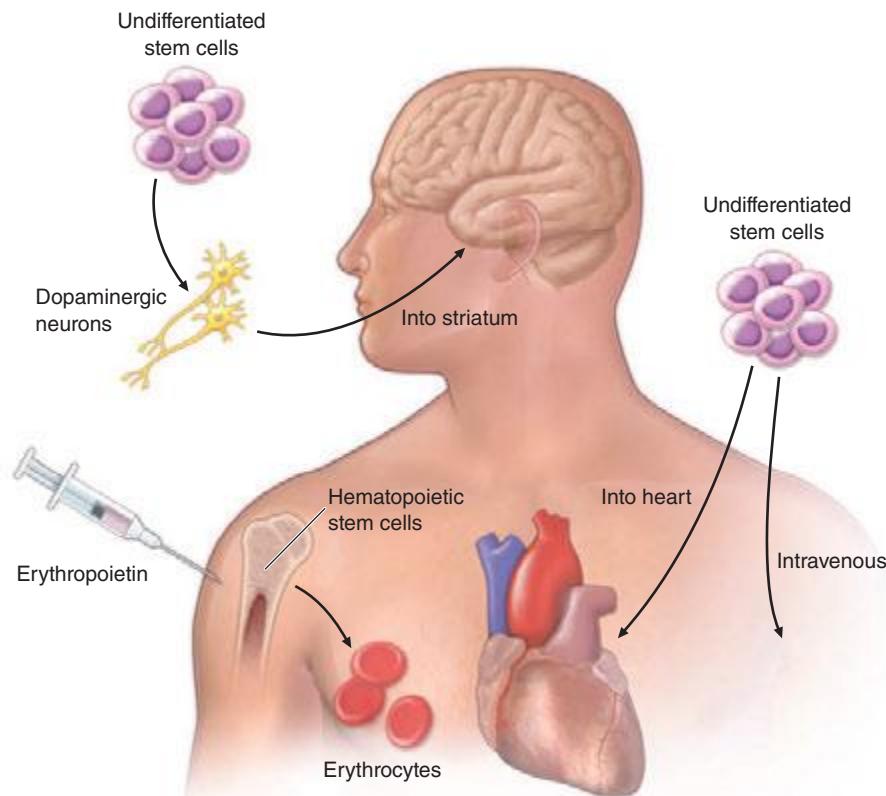


FIGURE 90e-1 Strategies for transplantation of stem cells. 1. Undifferentiated or partially differentiated stem cells may be injected directly into the target organ or intravenously. 2. Stem cells may be differentiated ex vivo before injection into the target organ. 3. Growth factors or other drugs may be injected to stimulate endogenous stem cell populations.

to be developed. Cells tend to develop abnormal karyotypes and other abnormalities with increased time in culture, and ES cells have the potential to form teratomas if all cells are not committed to the desired cell types before transplantation. Further, human ES cells are ethically controversial and, on these grounds, would be unacceptable to some patients and physicians despite their therapeutic potential. Nevertheless, there have been limited clinical trials of ES-derived cells in a number of disorders, including macular degeneration, myopia, and spinal cord injury.

Induced Pluripotent Stem Cells The field of stem cell biology was transformed by the discovery that adult somatic cells can be converted (“reprogrammed”) into pluripotent cells through the overexpression of four transcription factors normally expressed in pluripotent cells (Chap. 88). These iPS cells share most properties with ES cells, although there are distinct differences in gene expression between ES and iPS cells. The initial use of viruses to insert the transcription factors into somatic cells made the resulting cells unsuitable for clinical use. However, a number of strategies have since been developed to circumvent this problem, including the insertion of modified mRNAs, proteins, or microRNAs rather than cDNAs; the use of non-integrating viruses such as Sendai virus; the insertion of transposons with the programming factors, followed by their subsequent removal; and the use of floxed viral constructs, followed by treatment with Cre recombinase to excise those constructs. The safety of iPS cells in humans remains to be demonstrated, but clinical trials in macular degeneration and other disorders are planned. Potential advantages of iPS cells are that somatic cells from patients would generate pluripotent cells genetically identical to those of the patient and that these cells are not subject to the same ethical constraints as ES cells. It is not clear whether the differences in gene expression between ES and iPS cells will have any impact on their potential clinical utility, and studies of both cell types will be essential to resolve this issue.

Umbilical-Cord Stem Cells Umbilical-cord blood stem/progenitor cells (USCs) are widely and readily available. These cells appear to be associated with less graft-versus-host disease than are some other cell types, such as marrow stem cells. They have less human leukocyte antigen restriction than adult marrow stem cells and are less likely to be contaminated with herpesvirus. However, it is unclear how many different cell types can be generated from USCs, and methods for differentiating these cells into nonhematopoietic phenotypes are largely lacking. Nevertheless, there are ongoing clinical trials of these cells in dozens of disorders, including cirrhosis, cardiopathies, multiple sclerosis, burns, stroke, autism, and critical limb ischemia.

Organ-Specific Multipotent Stem Cells Organ-specific multipotent stem cells have the advantage of already being somewhat specialized so that the inducement of desired cell types may be easier. Cells potentially could be obtained from the patient and amplified in cell culture, circumventing the problems associated with immune rejection. Stem cells are relatively easy to harvest from some tissues, such as bone marrow and blood, but are difficult to harvest from other tissues, such as heart and brain. Moreover, these populations of cells are more limited in potentiality than are pluripotent ES or iPS cells, and they may be difficult to obtain in large quantities from many organs. Therefore, substantial efforts have been devoted to developing techniques for using more easily obtainable stem cell populations, such as bone marrow mesenchymal stem cells (MSCs), CD34+ hematopoietic stem cells (HSCs), cardiac mesenchymal cells, and adipose-derived stem cells (ASCs), for use in regenerative strategies. Tissue culture evidence suggests that these stem cell populations may be able to generate differentiated cell types unrelated to their organ source (including myocytes, chondrocytes, tendon cells, osteoblasts, cardiomyocytes, adipocytes, hepatocytes, and neurons) in a process known as *transdifferentiation*. However, it is still unclear whether these stem cells are capable of generating differentiated cell types that integrate into organs, survive, and function after transplantation in vivo. A number of early studies of MSCs transplanted into heart, liver, and other organs suggested that the cells had differentiated into organ-specific cell types with beneficial

effects in animal models of disease. Unfortunately, subsequent studies revealed that the stem cells had simply fused with cells resident in the organs and that the observed beneficial effects were due to paracrine release of trophic and anti-inflammatory cytokines. Further studies will be necessary to determine whether transdifferentiation of MSCs, ASCs, or other stem cell populations occurs at a high enough frequency to make these cells useful for stem cell replacement therapy. Despite the remaining issues, clinical trials of MSCs, autologous HSCs, USCs, and ASCs are being performed in many disorders, including ischemic cardiac disease, cardiomyopathy, diabetes, stroke, cirrhosis, and muscular dystrophy.

Regardless of the source of the stem cells used in regenerative strategies, a number of generic problems must be overcome for the development of successful clinical applications. These problems include the devising of methods to reliably generate large numbers of specific cell types, to minimize the risk of tumor formation or proliferation of inappropriate cell types, to ensure the viability and function of the engrafted cells, to overcome immune rejection when autografts are not used, and to facilitate revascularization of regenerated tissue. Each organ system will also pose tissue-specific problems for stem cell therapies.

DISEASE-SPECIFIC APPLICATIONS OF STEM CELLS

Ischemic Heart Disease and Cardiomyocyte Regeneration Because of the high prevalence of ischemic heart disease, extensive efforts have been devoted to the development of strategies for stem cell replacement of cardiomyocytes. Historically, the adult heart has been viewed as a terminally differentiated organ without the capacity for regeneration. However, recent studies have demonstrated that the heart has the capacity for low levels of cardiomyocyte regeneration (Chap. 265e). This regeneration appears to be accomplished by cardiac stem cells resident in the heart and possibly also by cells originating in the bone marrow. The heart might be an ideal source of stem cells for therapeutic use, but techniques for isolating, characterizing, and amplifying large numbers of these cells have not yet been perfected. For successful myocardial repair, stem cell therapy must deliver cells either systemically or locally, and the cells must survive, engraft, and differentiate into functional cardiomyocytes that couple mechanically and electrically with the recipient myocardium. The optimal method for cell delivery is not clear, and various experimental and clinical studies have successfully employed intramyocardial, transendocardial, intravenous, intracoronary, and retrograde coronary venous injections. In experimental myocardial infarction, functional improvements have been achieved after transplantation of a variety of different cell types, including ES cells, HSCs, MSCs, USCs, and ASCs. Early studies suggested that each of these cell types might have the potential to engraft and generate cardiomyocytes. However, most investigators have found that the generation of new cardiomyocytes by these cells is at best a rare event and that graft survival over long periods is poor. The preponderance of evidence suggests that the observed beneficial effects of most experimental therapies were not derived from direct stem cell generation of cardiomyocytes but rather from indirect effects of the stem cells on resident cells. It is not clear whether these effects reflect the release of soluble trophic factors, the induction of angiogenesis, the release of anti-inflammatory cytokines, or another mechanism. A wide variety of cell delivery methods, cell types, and cell doses have been used in a progressively enlarging series of clinical trials, but the fate of the cells and the mechanisms by which they alter cardiac function are still open questions. In aggregate, however, these studies have shown a small but measurable improvement in cardiac function and, in some cases, reduction in infarct size. In short, the available evidence suggests that the beneficial clinical impact reflects an indirect effect of the transplanted cells rather than genuine cell replacement.

Diabetes Successes with islet cell and pancreas transplantation have provided proof of concept for cell-based therapies for type 1 diabetes. However, the demand for donor pancreases far exceeds the number available, and maintenance of long-term graft survival is a problem. The search for a renewable source of stem cells capable of regenerating pancreatic islets has therefore been intensive. Pancreatic beta cell

turnover occurs even in the normal pancreas, although the source of the new beta cells remains controversial. This persistent turnover suggests that, in principle, it should be possible to develop strategies for reconstituting the beta cell population in diabetics. Attempts to devise techniques for promoting endogenous regenerative processes by using combinations of growth factors, drugs, and gene therapy have failed thus far, but this remains a potentially viable approach. A number of different cell types are candidates for use in stem cell replacement strategies, including iPS cells, ES cells, hepatic progenitor cells, pancreatic ductal progenitor cells, and MSCs. Successful therapy will depend on the development of a source of cells that can be amplified to produce large numbers of progeny with the ability to synthesize, store, and release insulin when it is required, primarily in response to changes in the ambient level of glucose. The proliferative capacity of the replacement cells must be tightly regulated to avoid excessive expansion of beta cell numbers and the consequent development of hyperinsulinemia/hypoglycemia; moreover, the cells must withstand immune rejection. Although it has been reported that ES and iPS cells can be differentiated into cells that produce insulin, these cells have a low content of insulin and a high rate of apoptosis and generally lack the capacity to normalize blood glucose levels in diabetic animals. Thus, ES and iPS cells have not yet been useful for the large-scale production of differentiated islet cells. During embryogenesis, the pancreas, liver, and gastrointestinal tract are all derived from the anterior endoderm, and transdifferentiation of pancreas to liver and vice versa has been observed in a number of pathologic conditions. There is also substantial evidence that multipotential stem cells reside within gastric glands and intestinal crypts. These observations suggest that hepatic, pancreatic, and/or gastrointestinal precursor cells may be reasonable candidates for cell-based therapy for diabetes, although it is unclear whether insulin-producing cells derived from pancreatic stem cells or liver progenitors can be expanded *in vitro* to clinically useful numbers. MSCs and neural stem cells both reportedly have the capacity to generate insulin-producing cells, but there is no convincing evidence that either cell type will be clinically useful. Clinical trials of MSCs, USCs, HSCs, and ASCs in both type 1 and type 2 diabetes are ongoing.

Nervous System Substantial progress has been made in the development of methodologies for generating neural cells from different stem cell populations. Human ES or iPS cells can be induced to generate cells with the properties of neural stem cells, and these cells in turn give rise to neurons, oligodendroglia, and astrocytes. Reasonably large numbers of these cells can be transplanted into the rodent brain with formation of appropriate cell types and no tumor formation. Multipotent stem cells present in the adult brain also can be easily amplified in number and used to generate all the major neural cell types, but the need for invasive procedures to obtain autologous cells is a major limitation. Fetal neural stem cells derived from miscarriages or abortions are an alternative but raise ethical concerns. Nevertheless, clinical trials of fetal neural stem cells have commenced in amyotrophic lateral sclerosis (ALS), stroke, and several other disorders. Transdifferentiation of MSCs and ASCs into neural stem cells, and vice versa, has been reported by numerous investigators, and clinical trials of such cells have begun for a number of neurologic diseases. Clinical trials of a conditionally immortalized human cell line and of USCs in stroke are also in progress. Because of the incapacitating nature of neural disorders and the limited endogenous repair capacity of the nervous system, clinical trials of stem cells in neurologic disorders have been particularly numerous, including trials in spinal cord injury, multiple sclerosis, epilepsy, Alzheimer's disease, ALS, acute and chronic stroke, numerous genetic disorders, traumatic brain injury, Parkinson's disease, and others. In diseases such as ALS, possible benefits are more likely to be due to indirect trophic effects than to neuron replacement. In Parkinson's disease, the major motor features of the disorder result from the loss of a single cell population: dopaminergic neurons within the substantia nigra; this circumstance suggests that cell replacement should be relatively straightforward. However, two clinical trials of fetal nigral transplantation failed to meet their primary endpoint and were complicated by the development of dyskinesia. Transplantation

of stem cell-derived dopamine-producing cells offers a number of potential advantages over the fetal transplants, including the ability of stem cells to migrate and disperse within tissue, the potential for engineering regulatable release of dopamine, and the ability to engineer cells to produce factors that will enhance cell survival. Nevertheless, the experience with fetal transplants points out the difficulties that may be encountered.

At least some of the neurologic dysfunction after spinal cord injury reflects demyelination, and both ES cells and MSCs can facilitate remyelination after experimental spinal cord injury (SCI). Clinical trials of MSCs in this disorder have commenced in a number of countries, and SCI was the first disorder targeted for the clinical use of ES cells. However, the ES cell trial in SCI was terminated early for nonmedical reasons. At present, no population of transplanted stem cells has been shown to have the capacity to generate neurons that extend axons over long distances to form synaptic connections (as would be necessary for replacement of upper motor neurons in ALS, stroke, or other disorders). For many injuries, including SCI, the balance between scar formation and tissue repair/regeneration may prove to be an important consideration. For example, it may ultimately prove necessary to limit scar formation so that axons can reestablish connections.

Liver Liver transplantation is currently the only successful treatment for end-stage liver diseases, but the shortage of liver grafts limits its application. Clinical trials of hepatocyte transplantation demonstrate its potential as a substitute for organ transplantation, but this approach is limited by the paucity of available cells. Potential sources of stem cells for regenerative strategies include endogenous liver stem cells (such as oval cells), ES cells, MSCs, and USCs. Although a series of studies in humans as well as animals suggested that transplanted MSCs and HSCs can generate hepatocytes, fusion of the transplanted cells with endogenous liver cells, giving the erroneous appearance of new hepatocytes, appears to be the underlying event in most circumstances. The available evidence suggests that transplanted HSCs and MSCs can generate hepatocyte-like cells in the liver only at a very low frequency, but there are beneficial consequences presumably related to indirect paracrine effects. ES cells can be differentiated into hepatocytes and transplanted in animal models of liver failure without the formation of teratomas. Clinical trials are in progress in cirrhosis with numerous cell types, including MSCs, USCs, HSCs, and ASCs.

Other Organ Systems and the Future The use of stem cells in regenerative strategies has been studied for many other organ systems and cell types, including skin, eye, cartilage, bone, kidney, lung, endometrium, vascular endothelium, smooth muscle, and striated muscle, and clinical trials in these and other organs are ongoing. In fact, the potential for stem cell regeneration of damaged organs and tissues is virtually limitless. However, numerous obstacles must be overcome before stem cell therapies can become a widespread clinical reality. Only HSCs have been adequately characterized by surface markers so that they can be unambiguously identified, a prerequisite for reliable clinical applications. The pathways for differentiating stem cells into specific cellular phenotypes are largely unknown, and the ability to control the migration of transplanted cells or predict the response of the cells to the environment of diseased organs is presently limited. Some strategies may employ the coadministration of scaffolding, artificial extracellular matrix, and/or growth factors to orchestrate differentiation of stem cells and their organization into appropriate constituents of the organ. There is currently no way to image stem cells *in vivo* after transplantation into humans, and it will be necessary to develop techniques to do so. Fortunately, stem cells can be engineered before transplantation to contain a contrast agent that may make their *in vivo* imaging feasible. The potential for tumor formation and the problems associated with immune rejection are impediments, and it will also be necessary to develop techniques for ensuring vascularization of regenerated tissues. There already are many strategies for supporting cell replacement, including coadministration of vasoactive endothelial growth factor to foster vascularization of the transplant. Some strategies also include the genetic engineering of stem cells with an inducible suicide gene so that the cells can be easily eradicated in

the event of tumor formation or another complication. The potential for stem cell therapies to revolutionize medical care is extraordinary, and disorders such as myocardial infarction, diabetes, and Parkinson's disease, among many others, are potentially curable by such therapies. However, stem cell-based therapies are still at a very early stage of development, and perfection of techniques for clinical transplantation of predictable, well-characterized cells is going to be a difficult and lengthy undertaking.

ETHICAL ISSUES

Stem cell therapies raise ethical and socially contentious issues that must be addressed in parallel with the scientific and medical opportunities. Society has great diversity with respect to religious beliefs, concepts of individual rights, tolerance for uncertainty and risk, and boundaries for how scientific interventions should be used to alter the outcome of disease. In the United States, the federal government has authorized research using existing human ES cell lines but still restricts the use of federal funds for developing new human ES cell lines. Ongoing studies of existing lines have indicated that they develop abnormalities with time in culture and that they may be contaminated with mouse proteins. These findings highlight the need to develop new human ES cell lines. The development of iPS cell technology may lessen the need for deriving new ES cell lines, but it is still not clear whether the differences in gene expression by ES and iPS cells are important for potential clinical use.

In considering ethical issues associated with the use of stem cells, it is helpful to draw from experience with other scientific advances, such as organ transplantation, recombinant DNA technology, implantation of mechanical devices, neuroscience and cognitive research, in vitro fertilization, and prenatal genetic testing. These and other precedents have pointed to the importance of understanding and testing fundamental biology in the laboratory setting and in animal models before applying new techniques in carefully controlled clinical trials. When these trials occur, they must include full informed consent and careful oversight by external review groups.

Ultimately, there will be medical interventions that are scientifically feasible but ethically or socially unacceptable to some members of a society. Stem cell research raises fundamentally difficult questions about the definition of human life, and it has raised deep fears about the ability to balance issues of justice and safety with the needs of critically ill patients. Health care providers and experts with backgrounds in ethics, law, and sociology must help guard against the premature or inappropriate application of stem cell therapies and the inappropriate involvement of vulnerable population groups. However, these therapies offer important new strategies for the treatment of otherwise irreversible disorders. An open dialogue among the scientific community, physicians, patients and their advocates, lawmakers, and the lay population is critically important to raise and address important ethical issues and balance the benefits and risks associated with stem cell transfer.

Gene transfer is a novel area of therapeutics in which the active agent is a nucleic acid sequence rather than a protein or small molecule. Because delivery of naked DNA or RNA to a cell is an inefficient process, most gene transfer is carried out using a vector, or gene delivery vehicle. These vehicles have generally been engineered from viruses by deleting some or all of the viral genome and replacing it with the therapeutic gene of interest under the control of a suitable promoter (Table 91e-1). Gene transfer strategies can thus be described in terms of three essential elements: (1) a vector; (2) a gene to be delivered, sometimes called the transgene; and (3) a physiologically relevant target cell to which the DNA or RNA is delivered. The series of steps in which the vector and donated DNA enter the target cell and express the transgene is referred to as *transduction*. Gene delivery can take place *in vivo*, in which the vector is directly injected into the patient, or, in the case of hematopoietic and some other target cells, *ex vivo*, with removal of the target cells from the patient, followed by return of the gene-modified autologous cells to the patient after manipulation in the laboratory. The latter approach effectively combines gene transfer techniques with cellular therapies (Chap. 90e).

Gene transfer is one of the most powerful concepts in modern molecular medicine and has the potential to address a host of diseases for which there are currently no available treatments. Clinical trials of gene therapy have been under way since 1990; a recent landmark in the field was the licensing, in 2012, of the first gene therapy product approved in Europe or the United States (see below). Given that vector-mediated gene therapy is arguably one of the most complex therapeutics yet developed, consisting of both a nucleic acid and a protein component, this time course from first clinical trial to licensed product is noteworthy for being similar to that seen with other novel classes of therapeutics, including monoclonal antibodies or bone marrow transplantation. Over 5000 subjects have been enrolled in gene transfer studies, and serious adverse events have been rare. Some of the initial trials were characterized by an overabundance of optimism and a failure to be appropriately critical of preclinical studies in animals; in addition, it was in some contexts not fully appreciated that animal studies are only a partial guide to safety profiles of products in humans (e.g., insertional mutagenesis). Initial exuberance was driven

by many factors, including an intense desire to develop therapies for hitherto untreatable diseases, lack of understanding of risks, and, in some cases, undisclosed financial conflicts of interest. After a teenager died of complications related to vector infusion, the field underwent a retrenchment; continued efforts led to a more nuanced understanding of the risks and benefits of these new therapies and more sophisticated selection of disease targets. Currently, gene therapies are being developed for a wide variety of disease entities (Fig. 91e-1).

GENE TRANSFER FOR GENETIC DISEASE

Gene transfer strategies for genetic disease generally involve gene addition therapy, an approach characterized by transfer of the missing gene to a physiologically relevant target cell. However, other strategies are possible, including supplying a gene that achieves a similar biologic effect through an alternative pathway (e.g., factor VIIa for hemophilia A); supplying an antisense oligonucleotide to splice out a mutant exon if the sequence is not critical to the function of the protein (as has been done with the dystrophin gene in Duchenne's muscular dystrophy); or downregulating a harmful effect through a small interfering RNA (siRNA). Two distinct strategies are used to achieve long-term gene expression: one is to transduce stem cells with an integrating vector, so that all progeny cells will carry the donated gene; and the other is to transduce long-lived cells, such as skeletal muscle or neurons. In the case of long-lived cells, integration into the target cell genome is unnecessary. Instead, because the cells are nondividing, the donated DNA, if stabilized in an episomal form, will give rise to expression for the life of the cell. This approach thus avoids problems related to integration and insertional mutagenesis.

IMMUNODEFICIENCY DISORDERS: PROOF OF PRINCIPLE

Early attempts to effect gene replacement into hematopoietic stem cells (HSCs) were stymied by the relatively low transduction efficiency of retroviral vectors, which require dividing target cells for integration. Because HSCs are normally quiescent, they are a formidable transduction target. However, identification of cytokines that induced cell division without promoting differentiation of stem cells, along with technical improvements in the isolation and transduction of HSCs, led to modest but real gains in transduction efficiency.

The first convincing therapeutic effect from gene transfer occurred with X-linked severe combined immunodeficiency disease (SCID), which results from mutations in the gene (*IL2RG*) encoding the γ c subunit of cytokine receptors required for normal development of

TABLE 91e-1 CHARACTERISTICS OF GENE DELIVERY VEHICLES

Features	Viral Vectors						
	Retroviral	Lentiviral	Adenoviral	AAV	Human Foamy Virus	HSV-1	Alpha Viruses
Viral genome	RNA	RNA	DNA	DNA	RNA	DNA	RNA
Cell division requirement	Yes	G ₁ phase	No	No	No	No	No
Packaging limitation	8 kb	8 kb	8–30 kb	5 kb	8.5 kb	40–150 kb	5 kb
Immune responses to vector	Few	Few	Extensive	Few	Few	Few in recombinant virus	Few
Genome integration	Yes	Yes	Poor	Poor	Yes	No	No
Long-term expression	Yes	Yes	No	Yes	Yes	No	No
Main advantages	Persistent gene transfer in dividing cells	Persistent gene transfer in transduced tissues	Highly effective in transducing various tissues	Elicits few inflammatory responses, nonpathogenic	Persistent gene expression in both dividing and nondividing cells	Large packaging capacity with persistent gene transfer	Limited immune responses against the vector
Main disadvantages	Theoretical risk of insertional mutagenesis (occurred in multiple cases)	Might induce oncogenesis in some cases (not yet observed)	Viral capsid elicits strong immune responses	Limited packaging capacity	In need of a stable packaging system	Residual cytotoxicity with neuron specificity	Transduced gene expression is transient

Abbreviations: AAV, adeno-associated virus; HSV, herpes simplex virus; SV, sarcoma virus.

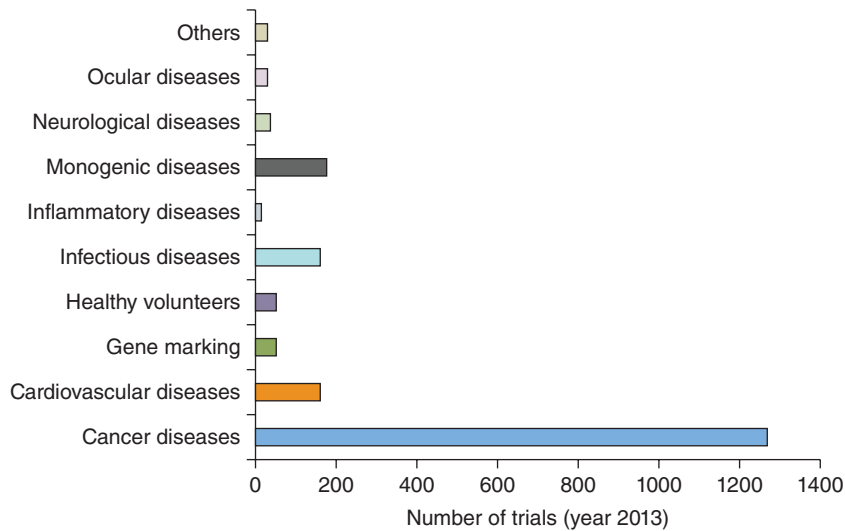


FIGURE 91e-1 Indications in gene therapy clinical trials. The bar graph classifies clinical gene transfer studies by disease. A majority of trials have addressed cancer, with monogenic disorders, infectious diseases, and cardiovascular diseases the next largest categories. (Adapted from SL Ginn et al: *J Gene Med* 15:65-77, 2013. Published online in Wiley Online.)

T and natural killer (NK) cells (Chap. 374). Affected infants present in the first few months of life with overwhelming infections and/or failure to thrive. In this disorder, it was recognized that the transduced cells, even if few in number, would have a proliferative advantage compared to the nontransduced cells, which lack receptors for the cytokines required for lymphocyte development and maturation. Complete reconstitution of the immune system, including documented responses to standard childhood vaccinations, clearing of infections, and remarkable gains in growth occurred in most of the treated children. However, among 20 children treated in two separate trials, five eventually developed a syndrome similar to T cell acute lymphocytic leukemia, with splenomegaly, rising white counts, and the emergence of a single clone of T cells. Molecular studies revealed that, in most of these children, the retroviral vector had integrated within a gene, *LMO-2* (*LIM* only-2), which encodes a component of a transcription factor complex involved in hematopoietic development. The retroviral long terminal repeat increases the expression of *LMO-2*, resulting in T cell leukemia.

The X-linked SCID studies were a watershed event in the evolution of gene therapy. They demonstrated conclusively that gene therapy could cure disease; of the 20 children eventually treated in these trials, 18 achieved correction of the immunodeficiency disorder. Unfortunately, 5 of the 20 patients later developed a leukemia-like disorder, and one died of this complication; the rest are alive and free of complications at time periods ranging up to 14 years after initial treatment. These studies demonstrated that insertional mutagenesis leading to cancer was more than a theoretical possibility (Table 91e-2). As a result of the experience in these trials, all protocols using integrating vectors in hematopoietic cells must include a plan for monitoring sites of insertion and clonal proliferation. Strategies to overcome this complication have included using a “suicide” gene cassette in the

TABLE 91e-2 POTENTIAL COMPLICATIONS OF GENE THERAPY

Gene silencing – repression of promoter
Genotoxicity – complications arising from insertional mutagenesis
Phenotoxicity – complications arising from overexpression or ectopic expression of the transgene
Immunotoxicity – harmful immune response to either the vector or transgene
Risks of horizontal transmission – shedding of infectious vector into environment
Risks of vertical transmission – germline transmission of donated DNA

vector, so that errant clones can be quickly ablated, or using “insulator” elements in the cassette, which can limit the activation of genes surrounding the insertion site. Lentiviral vectors, which can efficiently transduce nondividing target cells, are also likely to be safer than retroviral vectors, based on patterns of integration; the field is thus gradually moving toward these to replace retroviral vectors.

More clear-cut success has been achieved in a gene therapy trial for another form of SCID, adenosine deaminase (ADA) deficiency (Chap. 374). ADA-SCID is clinically similar to X-linked SCID, although it can be treated by enzyme replacement therapy with a pegylated form of the enzyme (PEG-ADA), which leads to immune reconstitution but not always to normal T cell counts. Enzyme replacement therapy is expensive (annual costs: \$200,000–\$300,000 in U.S. dollars). The initial trials of gene therapy for ADA-SCID were unsuccessful, but modifications of this protocol to include the use of HSCs rather than T cells as the target for transduction; discontinuation of PEG-ADA at the time of vector infusion, so that the transduced cells have a proliferative advantage over the nontransduced; and the use of a mild conditioning regimen to facilitate engraftment of the transduced cells have led to success without the complications

seen in the X-linked SCID trials. There have been no complications in the 10 children treated on the Milan protocol, with a median follow-up of >8 years. ADA-SCID, then, is an example where gene therapy has changed therapeutic options for patients. For those with a human leukocyte antigen (HLA)-identical sibling, bone marrow transplantation is still the best treatment option, but this includes only a minority of those affected. For those without an HLA-identical match, gene therapy has comparable efficacy to PEG-ADA, does not require repetitive injections, and does not run the risk of neutralizing antibodies to the bovine enzyme.

NEURODEGENERATIVE DISEASES: EXTENSION OF PRINCIPLE

The SCID trials gave support to the hypothesis that gene transfer into HSCs could be used to treat any disease for which allogeneic bone marrow transplantation was therapeutic. Moreover, the use of genetically modified autologous cells carried several advantages including no risk of graft-versus-host disease, guaranteed availability of a “donor” (unless the disease itself damages the stem cell population of the patient), and low likelihood of failure of engraftment. Cartier and Aubourg capitalized on this realization to conduct the first trial of lentiviral vector transduction of HSCs for a neurodegenerative disorder, X-linked adrenoleukodystrophy (ALD). X-linked ALD is a fatal demyelinating disease of the central nervous system caused by mutations in the gene encoding an adenosine triphosphate-binding cassette transporter. Deficiency of this protein leads to accumulation of very-long-chain fatty acids in oligodendrocytes and microglia, disrupting myelin maintenance by these cells. Affected boys present with clinical and neuroradiographic evidence of disease at age 6–8 and usually die before adolescence. Following lentiviral transduction of autologous HSCs in young boys with the disease, dramatic stabilization of disease occurred, demonstrating that stem cell transduction could work for neurodegenerative as well as immunologic disorders. Investigators in Milan carried this observation one step further to develop a treatment for another neurodegenerative disorder that has previously responded poorly to bone marrow transplantation. Metachromatic leukodystrophy is a lysosomal storage disorder caused by mutations in the gene encoding arylsulfatase A (ARSA). The late infantile form of the disease is characterized by progressive motor and cognitive impairment, and death within a few years of onset, due to accumulation of the ARSA substrate sulfatide in oligodendrocytes, microglia, and some neurons. Recognizing that endogenous levels of production of ARSA were too low to provide cross-correction by allogeneic transplant, Naldini and colleagues engineered a lentiviral vector that directed supraphysiologic

levels of ARSA expression in transduced cells. Transduction of autologous HSCs from children born with the disease, at a point when they were still presymptomatic, led to preservation and continued acquisition of motor and cognitive milestones at time periods as long as 32 months after affected siblings had begun to lose milestones. These results illustrate that the ability to engineer levels of expression can allow gene therapy approaches to succeed where allogeneic bone marrow transplantation cannot. It is likely that a similar approach will be used in other neurodegenerative conditions.

Transduction of HSCs to treat the hemoglobinopathies is an obvious extension of studies already conducted but represents a higher hurdle in terms of the extent of transduction required to achieve a therapeutic effect. Trials are now under way for thalassemia and for a number of other hematologic disorders, including Wiskott-Aldrich syndrome, and chronic granulomatous disease.

LONG-TERM EXPRESSION IN GENETIC DISEASE: IN VIVO GENE TRANSFER WITH RECOMBINANT ADENO-ASSOCIATED VIRAL VECTORS

Recombinant adeno-associated viral (AAV) vectors have emerged as attractive gene delivery vehicles for genetic disease. Engineered from a small replication-defective DNA virus, they are devoid of viral coding sequences and trigger very little immune response in experimental animals. They are capable of transducing nondividing target cells, and the donated DNA is stabilized primarily in an episomal form, thus minimizing risks arising from insertional mutagenesis. Because the vector has a tropism for certain long-lived cell types, such as skeletal muscle, the central nervous system (CNS), and hepatocytes, long-term expression can be achieved even in the absence of integration.

FIRST LICENSED PRODUCT

These features of AAV were used to develop the first licensed gene therapy product in Europe, an AAV vector for treatment of the autosomal recessive disorder lipoprotein lipase (LPL) deficiency. This rare disorder (1–2/million) is due to loss-of-function mutations in the gene encoding LPL, an enzyme normally produced in skeletal muscle and required for the catabolism of triglyceride-rich lipoproteins and chylomicrons. Affected individuals have lipemic serum and may have eruptive xanthomas, hepatosplenomegaly, and in some cases, recurrent bouts of acute pancreatitis. Clinical trials demonstrated the safety of intramuscular injection of AAV-LPL and its efficacy in reducing frequency of pancreatitis episodes in affected individuals, leading to drug approval in Europe. Additional clinical trials currently under way that use AAV vectors in the setting of genetic disease include those for muscular dystrophies, α_1 antitrypsin deficiency, Parkinson's disease, Batten's disease, hemophilia B, and several forms of congenital blindness.

HEMOPHILIA

Hemophilia (Chap. 78) has long been considered a promising disease model for gene transfer, because the gene product does not require precise regulation of expression and biologically active clotting factors can be synthesized in a variety of tissue types, permitting latitude in the choice of target tissue. Moreover, raising circulating factor levels from <1% (levels seen in those severely affected) into the range of 5% greatly improves the phenotype of the disease. Preclinical studies with recombinant AAV vectors infused into skeletal muscle or liver have resulted in long-term (>5 years) expression of factor VIII or factor IX in the hemophilic dog model. Administration to skeletal muscle of an AAV vector expressing factor IX in patients with hemophilia B was safe and resulted in long-term expression as measured on muscle biopsy, but circulating levels never rose to >1% for sustained periods, and a large number of IM injections (>80–100) was required to access a large muscle mass. Intravascular vector delivery has been used to access large areas of skeletal muscle in animal models of hemophilia and will likely be tested for this and other disorders in upcoming trials.

The first trial of an AAV vector expressing factor IX delivered to the liver in humans with hemophilia B resulted in therapeutic circulating levels at the highest dose tested, but expression at these levels (>5%) lasted for only 6–10 weeks before declining to baseline (<1%).

A memory T cell response to viral capsid, present in humans but not in other animal species (which are not natural hosts for the virus), likely led to the loss of expression (Table 91e-2). In response to these findings, a second trial included a short course of prednisolone, to be administered if factor IX levels began to decline. This approach resulted in long-term expression of factor IX, in the range of 2–5%, in men with severe hemophilia B. Current efforts are focused on expanding these trials, and extending the approach to hemophilia A.

RETINAL GENE THERAPY

A logical conclusion from the early experience with AAV in liver in the hemophilia trial was that avoidance of immune responses was key to long-term expression. Thus immunoprivileged sites such as the retina began to attract substantial interest as therapeutic targets. This inference has been elegantly confirmed in the setting of the retinal degenerative disease Leber's congenital amaurosis (LCA). Characterized by early-onset blindness, LCA is not currently treatable and is caused by mutations in several different genes; ~15% of cases of LCA are due to a mutation in a gene, *RPE65*, encoding a retinal pigment epithelial-associated 65-kDa protein. In dogs with a null mutation in *RPE65*, sight was restored after subretinal injection of an AAV vector expressing *RPE65*. Transgene expression appears to be stable, with the first animals treated >10 years ago continuing to manifest electroretinal and behavioral evidence of visual function. As is the case for X-linked SCID, gene transfer must occur relatively early in life to achieve optimal correction of the genetic disease, although the exact limitations imposed by age have not yet been defined. AAV-RPE65 trials carried out in both the United States and the United Kingdom have shown restoration of visual and retinal function in over 30 subjects, with the most marked improvement occurring in the younger subjects. Trials for other inherited retinal degenerative disorders such as choroideremia are under way, as are studies for certain complex acquired disorders such as age-related macular degeneration, which affects several million people worldwide. The neovascularization that occurs in age-related macular degeneration can be inhibited by expression of vascular endothelial growth factor (VEGF) inhibitors such as angiostatin or through the use of RNA interference (RNAi)-mediated knockdown of VEGF. Early-phase trials of siRNAs that target VEGF RNA are under way, but these require repeated intravitreal injection of the siRNAs; an AAV vector-mediated approach, which would allow long-term inhibition of the biological effects of VEGF through a soluble VEGF receptor, is now in clinical testing.

GENE THERAPY FOR CANCER

The majority of clinical gene transfer experience has been in subjects with cancer (Fig. 91e-1). As a general rule, a feature that distinguishes gene therapies from conventional cancer therapeutics is that the former are less toxic, in some cases because they are delivered locally (e.g., intratumoral injections), and in other cases because they are targeted specifically to elements of the tumor (immunotherapies, antiangiogenic approaches). Because cancer is a disease of aging, and many elderly are frail, the development of therapeutics with milder side effects is an important goal.

LOCAL APPROACHES

Cancer gene therapies can be divided into local and systemic approaches (Table 91e-3). Some of the earliest cancer gene therapy trials focused

TABLE 91e-3 GENE THERAPY STRATEGIES IN CANCER

Local/regional approaches
Suicide gene/prodrug
Suppressor oncogene
Oncolytic virus
Systemic approaches
Chemoprotection
Immunomodulation
Antiangiogenesis

on local delivery of a prodrug or a suicide gene that would increase sensitivity of tumor cells to cytotoxic drugs. A frequently used strategy has been intratumoral injection of an adenoviral vector expressing the thymidine kinase (*TK*) gene. Cells that take up and express the *TK* gene can be killed after the administration of ganciclovir, which is phosphorylated to a toxic nucleoside by *TK*. Because cell division is required for the toxic nucleoside to affect cell viability, this strategy was initially used in aggressive brain tumors (glioblastoma multiforme) where the cycling tumor cells were affected but the nondividing normal neurons were not. More recently, this approach has been explored for locally recurrent prostate, breast, and colon tumors, among others.

Another local approach uses adenoviral-mediated expression of the tumor suppressor p53, which is mutated in a wide variety of cancers. This strategy has resulted in complete and partial responses in squamous cell carcinoma of the head and neck, esophageal cancer, and non-small-cell lung cancer after direct intratumoral injection of the vector. Response rates (~15%) are comparable to those of other single agents. The use of oncolytic viruses that selectively replicate in tumor cells but not in normal cells has also shown promise in squamous cell carcinoma of the head and neck and in other solid tumors. This approach is based on the observation that deletion of certain viral genes abolishes their ability to replicate in normal cells but not in tumor cells. An advantage of this strategy is that the replicating vector can proliferate and spread within the tumor, facilitating eventual tumor clearance. However, physical limitations to viral spread, including fibrosis, intermixed normal cells, basement membranes, and necrotic areas within the tumor, may limit clinical efficacy. Oncolytic viruses are licensed and available in some countries but not in the United States.

SYSTEMIC APPROACHES

Because metastatic disease rather than uncontrolled growth of the primary tumor is the source of mortality for most cancers, there has been considerable interest in developing systemic gene therapy approaches. One strategy has been to promote more efficient recognition of tumor cells by the immune system. Approaches have included transduction of tumor cells with immune-enhancing genes encoding cytokines, chemokines, or co-stimulatory molecules; and *ex vivo* manipulation of dendritic cells to enhance the presentation of tumor antigens. Recently, considerable success has been achieved using lentiviral transduction of autologous lymphocytes with a cDNA encoding a chimeric antigen receptor (CAR). The CAR moiety consists of a tumor antigen-binding domain (e.g., an antibody to the B cell antigen CD19) fused to an intracellular signaling domain that allows T cell activation. The transduced lymphocytes can then recognize and destroy cells bearing the antigen. This CAR-T cell approach has proven extraordinarily successful in the setting of refractory chronic lymphocytic leukemia and pre-B-cell acute lymphoblastic leukemia. Infusion of gene-modified T cells engineered to recognize the B cell antigen CD19 has resulted in >1000-fold expansion *in vivo*, trafficking of the T cells to the bone marrow, and complete remission in a subset of patients who had failed multiple chemotherapy regimens. The cells persist as memory CAR+ T cells, providing ongoing antitumor functionality. Some patients experience a delayed tumor lysis syndrome requiring intensive medical management. This approach also causes an on-target toxicity, leading to B cell aplasia that necessitates lifelong IgG infusions. Current results indicate that long-lasting remissions can be achieved and the strategy can theoretically be extended to other tumor types if a tumor antigen can be identified.

Gene transfer strategies have also been developed for inhibiting tumor angiogenesis. These have included constitutive expression of angiogenesis inhibitors such as angiostatin and endostatin; use of siRNA to reduce levels of VEGF or VEGF receptor; and combined approaches in which autologous T cells are genetically modified to recognize antigens specific to tumor vasculature. These studies are still in early-phase testing.

Another novel systemic approach is the use of gene transfer to protect normal cells from the toxicities of chemotherapy. The most

extensively studied of these approaches has been transduction of hematopoietic cells with genes encoding resistance to chemotherapeutic agents, including the multidrug resistance gene *MDR1* or the gene encoding O⁶-methylguanine DNA methyltransferase (*MGMT*). *Ex vivo* transduction of hematopoietic cells, followed by autologous transplantation, is being investigated as a strategy for allowing administration of higher doses of chemotherapy than would otherwise be tolerated.

GENE THERAPY FOR VASCULAR DISEASE

The third major category addressed by gene transfer studies is cardiovascular disease. Initial experience was in trials designed to increase blood flow to either skeletal (critical limb ischemia) or cardiac muscle (angina/myocardial ischemia). First-line treatment for both of these groups includes mechanical revascularization or medical management, but a subset of patients are not candidates for or fail these approaches. These patients formed the first cohorts for evaluation of gene transfer to achieve therapeutic angiogenesis. The major transgene used has been VEGF, attractive because of its specificity for endothelial cells; other transgenes have included fibroblast growth factor (FGF) and hypoxia-inducible factor 1, α subunit (HIF-1 α). The design of most of the trials has included direct IM (or myocardial) injection of either a plasmid or an adenoviral vector expressing the transgene. Both of these vectors are likely to result in only short-term expression of VEGF, which may be adequate because there is no need for continued transgene expression once the new vessels have formed. Direct injection favors local expression, which should help to avoid systemic effects such as retinal neovascularization or new vessel formation in a nascent tumor. Initial trials of adeno-VEGF or plasmid-VEGF injection resulted in improvement over baseline in angiographically detectable vasculature, but no change in amputation frequency or cardiovascular mortality. Studies using different routes of administration or different transgenes are currently under way.

More recent studies have used AAV vectors to develop a therapeutic approach for individuals with refractory congestive heart failure. In preclinical studies, a vector encoding sarcoplasmic reticulum Ca²⁺ ATPase (SERCA2a) demonstrated positive left ventricular inotropic effects in a swine model of volume-overloaded heart failure. Results of a phase II study in which vector was infused via the coronary arteries in patients with congestive heart failure demonstrated safety and some indications of efficacy; larger studies are now planned.

OTHER APPROACHES

This chapter has focused on gene addition therapy, in which a normal gene is transferred to a target tissue to drive expression of a gene product with therapeutic effects. Another powerful technique under development is genome editing, in which a mutation is corrected *in situ*, generating a wild-type copy under the control of the endogenous regulatory signals. This approach makes use of novel reagents including zinc finger nucleases, TALENs and CRISPR, which introduce double-stranded breaks into the DNA near the site of the mutation and then rely on a donated repair sequence and cellular mechanisms for repair of double-strand breaks to reconstitute a functioning gene. Another strategy recently introduced into clinical trials is the use of siRNAs or short hairpin RNAs as transgenes to knock down expression of deleterious genes (e.g., mutant huntingtin in Huntington's disease or genes of the hepatitis C genome in infected individuals).

SUMMARY

The power and versatility of gene transfer approaches are such that there are few serious disease entities for which gene transfer therapies are *not* under development. The development of new classes of therapeutics typically takes two to three decades; monoclonal antibodies and recombinant proteins are recent examples. Gene therapeutics, which entered clinical testing in the early 1990s, traversed the same time course. Examples of clinical success are now abundant, and gene therapy approaches are likely to become increasingly important as a

TABLE 91e-4 TAKING HISTORY FROM SUBJECTS ENROLLED IN GENE TRANSFER STUDIES**Elements of History for Subjects Enrolled in Gene Transfer Trials**

1. What vector was administered? Is it predominantly integrating (retroviral, lentiviral, herpesvirus [latency and reactivation]) or nonintegrating (plasmid, adenoviral, adeno-associated viral)?
2. What was the route of administration of the vector?
3. What was the target tissue?
4. What gene was transferred in? A disease-related gene? A marker?
5. Were there any adverse events noted after gene transfer?

Screening Questions for Long-Term Follow-Up in Gene Transfer Subjects^a

1. Has a new malignancy been diagnosed?
2. Has a new neurologic/ophthalmologic disorder, or exacerbation of a preexisting disorder, been diagnosed?
3. Has a new autoimmune or rheumatologic disorder been diagnosed?
4. Has a new hematologic disorder been diagnosed?

^aFactors influencing long-term risk include: integration of the vector into the genome, vector persistence without integration, and transgene-specific effects.

therapeutic modality in the twenty-first century. A central question to be addressed is the long-term safety of gene transfer, and regulatory agencies have mandated a 15-year follow-up for subjects enrolled in gene therapy trials (Table 91e-4). Realization of the therapeutic benefits of modern molecular medicine will depend on continued progress in gene transfer technology.

Tissue engineering is a field that applies principles of regenerative medicine to restore the function of various organs by combining cells with biomaterials. It is multidisciplinary, often combining the skills of physicians, cell biologists, bioengineers, and material scientists, to recapitulate the native three-dimensional architecture of an organ, the appropriate cell types, and the supportive nutrients and growth factors that allow normal cell growth, differentiation, and function. Tissue engineering is a relatively new field, originating in the late 1970s. Early studies focused on efforts to create skin substitutes using biomaterials and epithelial skin cells with a goal of providing barrier protection for patients with burns. The early strategies employed a tissue biopsy, followed by *ex vivo* expansion of cells seeded on scaffolds. The cell-scaffold composite was later implanted back into the same patient, where the new tissue would mature. However, there were many hurdles to overcome. The three major challenges in the field of tissue engineering involved: (1) the ability to grow and expand normal primary human cells in large quantities; (2) the identification of appropriate biomaterials; and (3) the requirement for adequate vascularization and innervation of the engineered constructs.

ISOLATION AND GROWTH OF CELLS

The original model for tissue engineering focused largely on the isolation of tissue from the organ of interest, the growth and expansion of the tissue-specific cells, and the seeding of these cells onto three-dimensional scaffolds. Just a few decades ago, most primary cultures of human cells could not be grown and expanded in large quantities, representing a major impediment to the engineering of human tissues. However, the identification of specific tissue progenitor cells in the 1990s allowed expansion of multiple cell types, and progress has occurred steadily since then. Some cell types are more amenable to expansion than others, reflecting in part their native regenerative capacity but also varying requirements for nutrients, growth factors, and cell-cell contacts. As an example of progress, after years of effort, protocols for the growth and expansion of human cardiomyocytes are now available. However, there are still many tissue-specific cell types that cannot be expanded from tissue sources, including the pancreas, liver, and nerves. The discovery of pluripotent or highly multipotent stem cells (Chap. 88) may ultimately allow most human cell types to be used for tissue engineering. The stem cell characteristics depend on their origin and their degree of plasticity, with cells from the earliest developmental stages, such as embryonic stem cells, having the greatest plasticity. Induced pluripotent stem cells have the advantage that they can be derived from individual patients, allowing autologous transplants. They can also be differentiated, *in vitro*, along cell-specific lineages, although these protocols are still at an early stage of development. Human embryonic and induced pluripotent stem cells have a very high replicative potential, but they also have the potential for rejection and tumor formation (e.g., teratomas). The more recently described amniotic fluid and placental stem cells have a high replicative potential but without an apparent propensity for tumor formation. Moreover, they have the potential to be used in an autologous manner without rejection. Adult stem cells, such as those derived from bone marrow, also have less propensity for tumor formation and, if used in an autologous manner, will not be rejected, but their replicative potential is limited, especially for endoderm and ectoderm cells.

Stem cells can be derived from autologous or heterologous sources. Heterologous cells can be used when only temporary coverage is needed, such as replacing skin after a burn or wound. However, if a more permanent construct is required, autologous cells are preferred to avoid rejection. There are also practical issues related to tissue sources. For example, if a patient presents with end-stage heart disease, obtaining a cardiac tissue biopsy for cell expansion is unlikely to be feasible, and bone marrow-derived mesenchymal cells may provide an alternative.

BIOMATERIALS AS SCAFFOLDS FOR TISSUE ENGINEERING

The biomaterials used to create the scaffolds for tissue engineering require specific properties to enhance the long-term success of the implanted constructs. Ideally, the biomaterials should be biocompatible; elicit minimal inflammatory responses; have appropriate biomechanical properties; and promote cell attachment, viability, proliferation, and differentiated function. Ideally, the scaffolds should replicate the biomechanical and structural properties of the tissue being replaced. In addition, biodegradation should be controlled such that the scaffold retains its structural integrity until the cells deposit their own matrix. If the scaffolds degrade too quickly, the constructs may collapse. If the scaffolds degrade too slowly, fibrotic tissue may form. Also, the degradation of the scaffolds should not alter the local environment unfavorably, because this can impair the function of cells or newly formed tissue.

The first scaffolds designed for tissue regeneration were naturally derived materials, such as collagen. The first artificially derived material for tissue engineering used a biodegradable scaffold made of polyglycolic acid. Naturally derived scaffolds have properties very similar to the native matrix, but there is an inherent batch-to-batch variability, whereas the production of artificially derived biomaterials can be better controlled, allowing for more uniform results. More recently, combination scaffolds, made of both naturally and artificially derived biomaterials, have been used for tissue engineering.

An emerging area is the use of peptide nanostructures to facilitate tissue engineering. Some of these are self-assembling peptide amphiphiles that allow scaffolds to form *in vivo*, for example at sites of spinal cord injury where they have been used experimentally to prevent scar formation and facilitate nerve and blood vessel regeneration. Peptide nanostructures can be combined with other biomaterials, and they can be linked to growth factors, antibodies, and various signaling molecules that can modulate cell behavior during organ regeneration.

VASCULARIZATION AND INNERVATION

Implanted tissue-engineered constructs require adequate vascularity and innervation. Judah Folkman, a pioneer in the field of angiogenesis, made the observation that cells could survive in volumes up to 3 mm³ via nutrient diffusion alone, but larger cell volumes required vascularization for survival. Adequate vascularity was also essential for normal innervation to occur. This was a major challenge in the field of tissue engineering, which largely depended on the patient's native angiogenesis and innervation. Even if sufficient cell quantities are available, there is a theoretical limit on the types of tissue constructs that could be created. In response to this challenge, material scientists designed scaffolds with much greater porosity and architecture. Scaffold designs included the creation of thin, porous sponges comprised of 95% air, markedly increasing the surface area for the resident cells. These properties promoted increased vascularity and innervation. The addition of growth factors, such as vascular endothelial growth factor and nerve growth factor, has been used to enhance angiogenesis and innervation.

LEVEL OF COMPLEXITY FOR THE ENGINEERING OF TISSUES AND ORGANS

All human tissues are complex. However, from an architectural aspect, tissues can be categorized under four levels. Flat tissue structures, such as skin, are the least complex (level 1), comprised predominantly of a single epithelial cell type. Tubular structures, such as blood vessels and the trachea, are more complex architecturally (level 2) and must be constructed to ensure that the structure does not collapse over time. These tissues typically have two major cell types. They are designed to act as a conduit for air or fluid at a steady state within a defined physiologic range. Hollow nontubular organs, such as the stomach, bladder, or uterus, are more complex architecturally (level 3). The cells are functionally more complex, and these cell types often have a functional interdependence. By far, the most complex are the solid organs (level 4), because the amount of cells per cm² are exponentially greater than any of the other tissue types.

For the first three tissue levels (1–3), when the constructs are initially implanted, the cell layering on the scaffolds is thin, not unlike that seen in tissue culture matrices. The cell layering continues to

mature, in concert with the recipient's native angiogenesis and neoinnervation. For level 4 (solid) organs, the vascularity requirements are substantial, and native tissue angiogenesis is not sufficient. The engineering strategies for tissues vary according to their complexity level.

STRATEGIES FOR TISSUE ENGINEERING

The basic principles of tissue engineering involve the use of the relevant cell populations, where the cell biology is well understood and the cells can be reproducibly retrieved and expanded, and the use of optimized biomaterials and scaffold designs. Cell seeding can be performed using various techniques, including static or flow-based systems that use bioreactors.

Most techniques for the engineering of tissues fall under one of five strategies (Fig. 92e-1):

1. Scaffolds can be used alone, without cells, and implanted, where they depend on native cell migration onto the scaffold from the adjacent tissue for regeneration. The first use of decellularized scaffolds for tissue regeneration was for urethral reconstruction. These techniques are most optimal when the size of the defect is relatively small, usually <0.5 cm from each tissue edge. Larger defects tend to heal by scarring, due to the deposition of fibroblasts, and eventual fibrosis. Scaffolds alone have also been used for other applications, including for wound coverage, soft tissue coverage after joint surgery, urogynecologic applications for sling surgery, and as materials for hernia repair.
2. A more recent strategy in tissue engineering involves the use of proteins, cytokines, genes, or small molecules that induce in situ

tissue regeneration, either alone or with the use of scaffolds. For example, gene transcription factors used in the mouse pancreas led to tissue regeneration. Surgically implanted decellularized heart valve scaffolds, coated with proteins that attract vascular stem cells, led to the creation of in situ cell-seeded functional heart valves in sheep. Drugs that induce muscle regeneration are being tested clinically. Small molecules that induce tissue regeneration are currently under investigation for multiple applications, including growth of skin and hair and for musculoskeletal applications.

3. The most common strategy for the engineering of tissues uses scaffolds seeded with cells. The most direct and established type of tissue engineering uses flat scaffolds, either artificial or naturally derived, that are seeded with cells and used for the replacement or repair of flat tissue structures. The flat scaffolds can also be sized and molded at the time of surgical implantation, or they can be shaped prior to cell seeding, for example, for tubular organs such as blood vessels or nontubular hollow tissues such as bladders. Bioreactors are often used to expose the cell-scaffold construct to mechanical forces, such as, stress, strain, and pulsatile flow that aid in the normal development of the cells into tissues (Video 92e-1, engineered heart valve in a pulsatile bioreactor showing the valves opening and closing). This strategy is the most common method used for tissue regeneration to date, and tissues and organs, such as skin, blood vessels, urethras, tracheas, vaginas, and bladders, have been engineered and implanted in patients using these techniques.
4. The fourth strategy in tissue engineering is applicable for solid organs, where discarded organs are exposed to mild detergents and are decellularized, leaving behind a three-dimensional scaffold that preserves its vascular tree. The scaffold can then be reseeded with the patient's own expanded vascular and tissue-specific cells. This strategy was used initially to create solid phallic structures in rabbits that were functional and able to produce offspring. Similar strategies were also used to recellularize miniature heart, liver, and kidney structures, with limited functionality to date, but with an established proof of concept (Video 92e-2, a dye is injected through the portal artery of a decellularized liver showing an intact vascular tree). These techniques are currently under investigation and have not been used clinically to date.

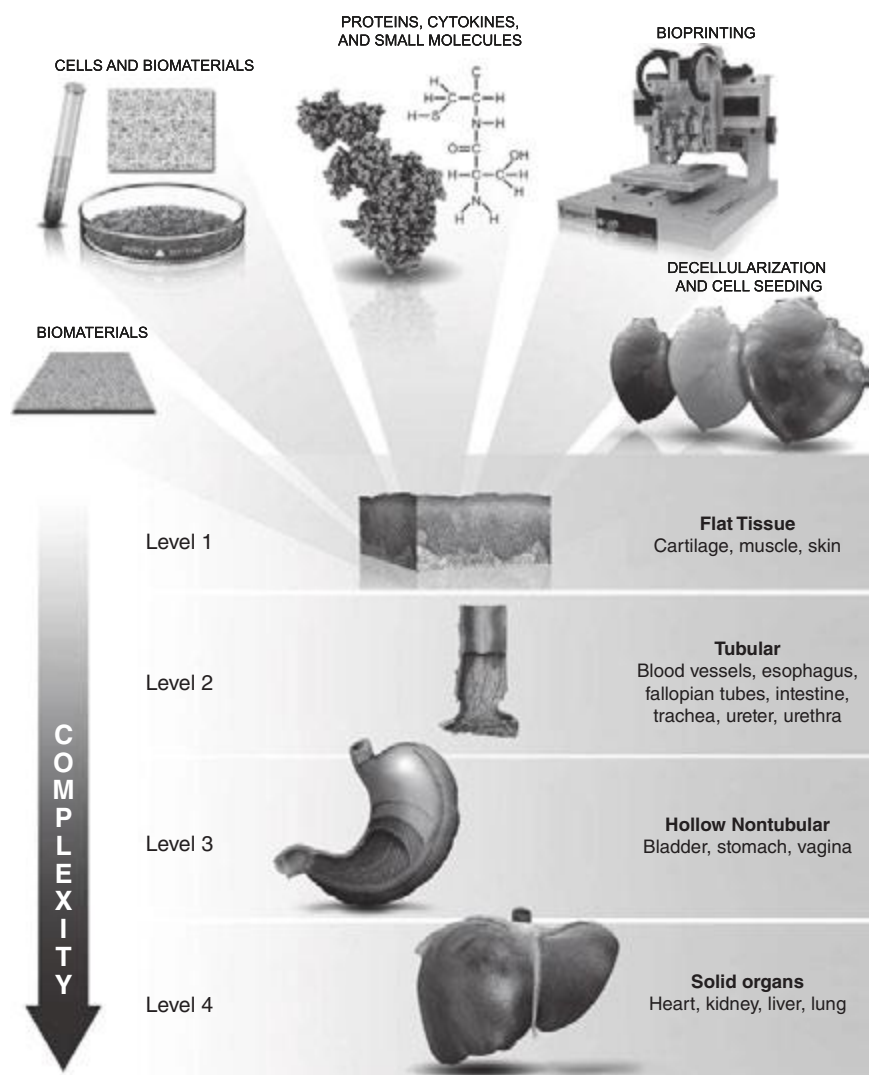


FIGURE 92e-1 Strategies for tissue and organ engineering.

5. The fifth strategy for tissue engineering involves the use of bioprinting. These technologies arose through the use of modified desktop inkjet printers over a decade ago. The inkjet cartridges were filled with a cell-hydrogel combination instead of ink. A rudimentary three-dimensional elevator was lowered each time the cartridge deposited the cells and hydrogel, thus building miniature solid structures, such as two-chambered heart organoids, one layer at a time. More sophisticated bioprinters have now been built that have additional computer-aided design (CAD) and three-dimensional printing technologies. The information to print the organ can be personalized using the patient's own imaging studies that help to define the size and shape of the particular tissue (Video 92e-3, a modified inkjet printer shows the three-dimensional construction of a two-chambered heart and how the structure beats with the cardiomyocytes in synchrony). Bioprinting is a tool that allows a scale-up option for the production of engineered tissues. Its use is still experimental and has not been applied clinically to date.

FUTURE DIRECTIONS

A number of engineered tissues, including architecturally flat, tubular, and hollow nontubular organs,

TABLE 92e-1 ENGINEERED TISSUES IN PATIENTS

Engineered Tissues in Patients	Year	Clinical Indication	Cell Source
Skin	1989	Burns	Autologous cultured epithelial cells
Cartilage	1997	Knee articular cartilage injuries	Autologous cultured chondrocytes
Skin	1998	Venous leg ulcers	Allogeneic neonatal dermal keratinocytes and fibroblasts
Bladder	1998	Myelomeningocele with poorly compliant bladders	Autologous smooth-muscle and urothelial cells on biodegradable scaffolds
Blood vessel	2001	Single ventricle physiology (pulmonary artery)	Autologous mixed population of endothelial, muscle, and fibroblast cells on biodegradable scaffolds
Blood vessel	2001	Single ventricle physiology (pulmonary artery)	Autologous bone marrow mononuclear cells on biodegradable scaffolds
Skin	2003	Diabetic foot ulcers	Allogeneic neonatal dermal fibroblasts on biodegradable scaffolds
Urethra	2004	Urethral strictures	Autologous smooth-muscle and epithelial cells seeded on biodegradable scaffolds
Skin	2004	Deep partial-thickness burns	Autologous keratinocytes, melanocytes, Langerhans cells, and fibroblasts
Blood vessel	2004	Kidney dialysis arteriovenous shunts	Autologous tubularized fibroblasts
Trachea	2008	End-stage airway disease	Autologous epithelial cells and mesenchymal stem cell–derived chondrocytes seeded on biodegradable scaffolds
Skin	2013	Acute and chronic wound repair	Allogeneic mesenchymal stem cells, fibroblasts, and epithelial cells from placental membrane
Blood vessel	2013	Kidney dialysis arteriovenous shunts	Acellular collagenous extracellular matrix secreted by smooth muscle

have been implanted in patients dating back to the 1990s. These include bladders, blood vessels, urethras, vaginal organs, tracheas, and skin for permanent replacement (**Table 92e-1**). Various types of skin substitutes, which were used as temporary “living wound dressings” to cover burn areas until skin grafts could be obtained from the same patient, were implanted starting in the 1990s. However, the use of engineered skin as a permanent replacement occurred only recently. Many engineered tissues are still being used in patients under regulatory guidelines for clinical trials. To date, solid organs have not yet been engineered for clinical use.

Tissue engineering is a rapidly evolving field where new technologies are continuously being applied to achieve success. The field still has many challenges ahead, including the long regulatory timelines required for the approval of widespread use, the need for improved scale-up production technologies, and the cost of the technologies, which include multiple processes involving biologics. Nonetheless, the list of tissues and organs being implanted in patients keeps growing, and the ability of these technologies to improve health has been demonstrated. More patients should be able to benefit from these technologies in the coming years.

VIDEO 92e-1 Engineered heart valve in a pulsatile bioreactor showing the valves opening and closing.

VIDEO 92e-2 A dye is injected through the portal artery of a decellularized liver showing an intact vascular tree.

VIDEO 92e-3 A modified inkjet printer shows the three-dimensional construction of a two-chambered heart and how the structure beats with the cardiomyocytes in synchrony.

93e World Demography of Aging

Richard M. Suzman, John G. Haaga

Population aging is transforming the world in dramatic and fundamental ways. The age distributions of populations have changed and will continue to change radically, due to long-term declines in fertility rates and improvements in mortality rates (Table 93e-1). This transformation, known as the Demographic Transition, is also accompanied by an epidemiologic transition, in which noncommunicable chronic diseases are becoming the major causes of death and contributors to the burden of disease and disability. A concomitant of population aging is the change in key ratios expressing “dependency” of one form or another—the ratio of adults in the workforce to those typically out of the workforce, such as infants, children, retired “young old” (those still active but in ways other than paid work), and the oldest old. Global aging will affect economic growth, migration, patterns of work and retirement, family structures, pension and health systems, and even trade and the relative standing of nations. Both absolute numbers (the size of an age group) and ratios (the ratio of those in working ages to dependents such as the young or retired, or the ratio of children to older people) are important. The size of age groups might affect the number of hospital beds needed, whereas the ratio of children to older people could affect the relative demand for pediatricians and geriatricians.

Although the increase in life expectancy, resulting from a series of social, economic, public health, and medical victories over disease, might very well be considered the crowning achievement of the past century and a half, the increased length of life coupled with the shifts in dependency ratios present formidable long-term challenges.

The pace of the change is accelerating. In countries where the Demographic Transition began earlier, the process was slower: it took France 115 years for the proportion of the age group 65 and older to

increase from 7 to 14% of the total population, and the United States will soon have completed this same increase in 69 years. But in countries that started the transition later, the process is occurring much more rapidly: Japan took 26 years to go from 7 to 14% age 65 and older, while China and Brazil are projected to require just 24 years.

Sometime around the year 2020, for the first time ever, the number of people age 65 and older in the world is expected to exceed that of children under the age of 5. Around the middle of the twentieth century, the under-5 age group constituted almost 15% of the total population and the over-65 age group 5%. It took about 70 years for these two to reach equal proportions. But demographers predict it will take only another 25–30 years for the 65 and older age group to equal about 15% and be about double the number of children under age 5. By the middle of their careers, medical students in most countries should expect to be practicing in far older populations. Preparations for these changes need to begin decades in advance, and the costs and penalties for delay can be very high. Although some governments have started planning for the long term, many, if not most, have yet to begin.

HISTORICAL

Population aging around the world in recent decades has followed a broadly similar pattern, starting with a decline in infant and childhood mortality that precedes a decline in fertility; at later stages, mortality at older ages declines as well. Declining fertility began as early as the beginning of the nineteenth century in the United States and France and extended to the rest of Europe and North America and parts of East Asia by the middle of the twentieth century. Since World War II, fertility declines have started in all other world regions. In fact, more than half the world’s population now lives in countries or provinces with fertility rates below the replacement level of just over two live births per woman. Mortality rates also began to change, relatively slowly at first, in Western Europe and North America during the nineteenth century. At first, changes were most evident at the youngest ages.

TABLE 93e-1 SELECTED INDICATORS OF POPULATION AGING, ESTIMATES FOR 2009, AND PROJECTIONS TO 2050; SELECTED REGIONS AND COUNTRIES

	Population Age 60+ (in millions)		Percentage of Population 60+		Life Expectancy at Birth		Life Expectancy at 60		Old Age Support Ratio ^a	
	2012	2050	2012	2050	Male	Female	Male	Female	2012	2050
World	809	2031	11	22	65.4	69.8	18.1	21.2	8	4
More developed regions	279	418	21	32	73.6	80.5	19.6	23.7	4	2
Less developed regions	530	1613	9	20	63.9	67.4	17.3	19.6	11	4
Least developed countries	46	181	5	11	54.7	57.2	15.3	16.8	16	9
Africa	60	215	6	10	52.9	55.3	15.2	17.1	16	10
Asia	446	1252	11	24	67.1	75.7	17.6	20.3	10	4
China	181	439	13	34	71.3	74.8	18.2	20.7	8	2
Japan	40	45	32	41	79	86.2	22.1	27.8	3	1
Republic of Korea	8	18	17	39	75.9	82.5	19.7	24.6	6	2
India	100	323	8	19	62.1	65	16	17.9	13	5
Europe	166	242	22	34	71.1 ^b	79.1 ^b	18.3	22.6	4	2
Russian Federation	26	39	19	31	60.3	73.1	14.3	19.2	6	3
United Kingdom	14	22	23	29	77.2	81.6	20.4	24	4	3
Italy	15.8	22.3	26	39	78.1	84.1	21.5	25.9	3	2
Germany	21.1	27.9	26	40	77.1	82.4	20.4	24.6	3	2
France	14.1	22.0	23	33	77.6	84.7	21.6	26.6	4	2
Americas										
United States	56.2	110.5	18	27	76.9	81.4	21.2	24.6	5	3
Canada	6.5	14.1	20	32	78.3	82.9	21.7	25.2	5	2

^aUN Population Division defines Old Age Support Ratio as the number of people age 15 to 64 years for every person age 65 or older. ^bThe UN includes all European regions in its overall statistics; life expectancy at birth for males ranges from 63.8 years in Eastern Europe to 77.4 years in Western Europe. For women it ranges from 74.8 to 83.1 years in Western Europe.

Improvements in water supply and sewage handling, as well as in nutrition and housing, accounted for most of the improvement before the 1940s, when antibiotics and vaccines and increasing education of mothers began to make a major impact. Since the middle of the twentieth century, the “Child Survival Revolution” has spread to all parts of the world. Children almost everywhere in the world are much more likely to reach late middle age now than in previous generations.

Especially since around 1960, mortality at older ages has improved steadily. This improvement has been primarily due to advances in care of heart disease and stroke and in control of conditions like hypertension and hypercholesterolemia that lead to circulatory diseases. In some parts of the world, smoking rates have declined, and these declines have led to lower incidence of many cancers, heart disease, and stroke.

The initial decline in fertility resulted in older age groups becoming a larger fraction of the total population. Declines in adult and old age mortality contributed to population aging in the later stages of the process. Life expectancy at birth—the average age to which someone is expected to live, under prevailing mortality conditions—has been calculated at around 28 years in ancient Greece, perhaps 30 years in medieval Britain, and less than 25 years in the colony of Virginia in North America. In the United States, life expectancy climbed slowly during the nineteenth century, reaching 49 years for white women by 1900. White men had a life expectancy 2 years lower than that for white women, and black Americans had a life expectancy 14 years lower than did white Americans in 1900. By the early twenty-first century, life expectancy in the United States had improved dramatically for all, with the sex gap wider and the racial gaps narrower than at the beginning of the century: 76 years for white men in 2006; 81 years for white women; and 70 and 76 years for black men and women, respectively. However, although the United States had a relatively high life expectancy compared to other high-income countries around 1980, almost all such countries have in the interim exceeded the United States in life expectancy. Female life expectancy, especially for whites in the United States, has done particularly poorly, and this has been attributed to relatively high rates of lifetime smoking.

At later stages of the demographic transition, mortality declines at the oldest ages, leading to increases in the 65 and older population, and the oldest old, those older than age 85 years. Migration can also affect population aging. An influx of young migrants with high birth rates can slow (though not stop) the process, as it has in the United States and Canada; or the out-migration of the young leaving older people behind can accelerate aging at the population level, as it has in many rural areas of the world.

REGIONAL AGING—NUMBERS AND PERCENTAGES OLDER THAN AGE 60 YEARS

Regions of the world are at very different stages of the demographic transition (Fig. 93e-1). Of a world population of 6.8 billion in 2012, approximately 11% were older than age 60 years, with Japan (32%) and Europe (22%) being the oldest regions (Germany and Italy 27% each) and the United States having 19%. The percentage of the population older than age 60 years in the United States has remained lower than in Europe, due both to modestly higher fertility rates and to higher rates of immigration. Asia has about 10% older than age 60 years, with the population giants close to the average—China (12%), Indonesia (9%), and India (7%). Middle Eastern and African countries have the lowest proportions of older people (5% or lower).

Based on estimates from the United Nations Population Division, 809 million people were age 60 years or older in 2012, of whom 279 million lived in more developed countries and 530 million in less developed countries (as classified by the United Nations). The countries with the largest populations of those age 60 and older were China (181 million), India (100 million), and the United States (60 million).

NUMBERS—POPULATION SIZE PROJECTIONS

Population projections make use of expected fertility, mortality, and migration rates and should be regarded as uncertain when applied 40 or more years in the future. However, the population that will be age 60 and older in 2050 have all been born and survived childhood in 2014, so uncertainty about their numbers (as distinct from their proportion of the total population) is not great. Comparing the maps of the world in 2010 (Fig. 93e-1) and 2050 (Fig. 93e-2), it is apparent that the middle- and low-income countries in Latin America, Asia, and much of Africa will soon be joining the “oldest” category. In less than four decades between 2012 and 2050, the United Nations Population Division projects that the world population age 60 and older will more than double to 2.03 billion, with the least developed regions more than quadrupling. China’s 60+ population is projected to reach 439 million, India’s 323 million, and the United States’s 107 million. Over the same period, the median age of the world’s population is expected to increase by 10 years.

Current global life expectancy at birth is estimated to be 65.4 for men and 69.8 for women, with the comparable figures for the more developed region being 73.6 and 80.5 years. Life expectancy in the least developed countries averaged only 57.2 for women and 54.7 for men. Life expectancy at birth is heavily influenced by infant and child mortality, which is considerably higher in poor countries. At older ages, the gap between rich and poor nations is narrower; so while women who have reached age 60 in wealthy countries can expect 23.7 more years of life on average, women at age 60 in poor countries live 16.8 years on average—a significant difference but not so stark as the difference in life expectancy at birth. At the lowest levels of per capita gross national product (GNP), life expectancy shows a powerful positive association with this measure of economic development, but then the slope of the relationship flattens out; for countries with average incomes above about \$20,000 per year, life expectancy is not closely related to income. At each level of economic development, there is significant variation in life expectancy, indicating that many other factors influence life expectancy.

Japan, France, Italy, and Australia currently have some of the highest life expectancies in the world, while the United States has lagged behind other high-income countries since about 1980, especially in the case of white women. The causes of this lag are being explored, but the cumulative number of years that people have smoked tobacco by the time they reach older ages and the prevalence of obesity appear to play important roles.

GROWTH OF THE OLDEST OLD POPULATION—THOSE OVER AGE 85

A modern feature of population aging has been the almost explosive growth of the age group known as the oldest old, variously defined as those over age 80 or age 85. This is the age group with the highest burden of noncommunicable degenerative disease and related disability. Thirty years ago, this group attracted little attention because they were hidden within the overall older population in most statistical reports; for example, the U.S. Census Bureau merged them into a 65+ category. The reduction of mortality at older ages coupled with larger birth



FIGURE 93e-1 Percentages of national populations age 60+, in 2010. (From the U.S. Census Bureau, International Database. StatPlanet Mapping Software.)

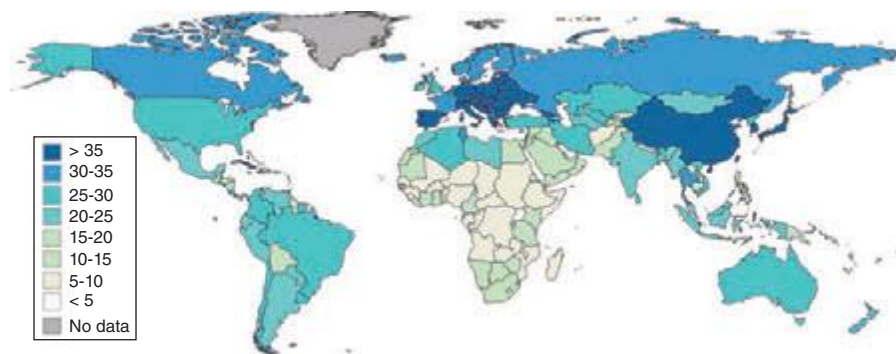


FIGURE 93e-2 Percentages of national populations age 60+, in 2050 (projections).

(From the U.S. Census Bureau, International Database. StatPlanet Mapping Software.)

cohorts surviving into old age led to the rapid growth of the oldest old. This age group is predicted to grow at a significantly higher rate than the 60+ population, and one estimate has the current 102 million age 80+ increasing to almost 400 million by 2050 (Table 93e-2). Projected increases are astounding: China's 80+ population might increase from 20 to 96 million, India from 8 to 43 million, the United States from 12 to 32 million, and Japan from 9 to 16 million. The numbers of centenarians are increasing at an even faster rate.

THE FUTURE OF LIFE EXPECTANCY

The members of the population who could potentially become age 80 and older in 2050 are already alive today. The actual numbers of people who will be age 80 and older in 2050 will therefore depend almost solely on adult and old age mortality rates over the next 35 years. The history of the decline of mortality suggests that improvements in the standard of living, including increased and improved education and improved nutrition, coupled with improvements in public health stemming from an understanding of the germ theory of disease initially led to the decline in mortality, with medical achievements such as antibiotics and improved understanding of risk factors for cardiovascular and circulatory diseases becoming factors only in the post-World War II period; the largest strides in cardiovascular disease came only in more recent decades. The improvements in educational attainment of succeeding generations have been credited in large part for improvements in child mortality during the past century, because educated mothers

TABLE 93e-2 ESTIMATES (2012) AND PROJECTIONS (2050) FOR THE POPULATION AGED 80 YEARS AND OLDER: SELECTED REGIONS AND COUNTRIES

	Population Age 80 and Over (in millions)	
	2012	2050
World	113	4,062
More developed regions	56	121
Less developed regions	58	274
Africa	5	22
Asia	54	238
China	20	97
Japan	9	16
India	9	45
Europe	33	68
Russian Federation	4	7
Italy	4	8
Germany	4	10
Americas		
United States	12	32
Canada	14	
Mexico	2	8

Source: United Nations Population Division, *World Population Ageing* 2012.

are especially likely to understand and take advantage of measures to reduce infection. The effects of continuing progress will likely be seen in coming decades as well, because educational attainment is associated with improved health and survival at older ages. Countries vary in the extent to which the "future elderly" cohorts will be more educated. China in particular will have a much more educated elderly population in 2050 (with more than two-thirds of the 65+ population having completed secondary school) than it did in 2000 (when only 10% of older people had a secondary education). In the United States and other rich nations, this change has largely taken place already; future changes in educational attainment of the elderly population will be less dramatic.

Holding aside the possibility of new infectious diseases ravaging populations as AIDS did in some African countries, debates about future life expectancy revolve around the balance and influence of risk factors such as obesity; the possibility of reducing the deaths from current killers such as cancer, heart disease, and diabetes; whether there is some natural limit to life expectancy; and the distant though nonzero possibility that science will find a way to slow the basic processes of aging.

While some have posited natural limits to human life expectancy, the limits have been surpassed with some regularity, and at the very oldest ages in the leading countries with the highest life expectancy, there appears to be little evidence of any approaching asymptote. Indeed a surprising discovery was that life expectancy in the leading country over the last century and a half, with different countries taking the lead in different epochs, could be represented almost perfectly by a straight line, with the increase for females showing a steady and astonishing increase of three months per year or 2.5 years per decade (Fig. 93e-3). No single country kept that pace of improvement the entire time, but this trend calls into question the notion that improvement must slow down, at least in the near future.

There remains a great deal of diversity in health conditions both among and within national populations. There is nothing inevitable

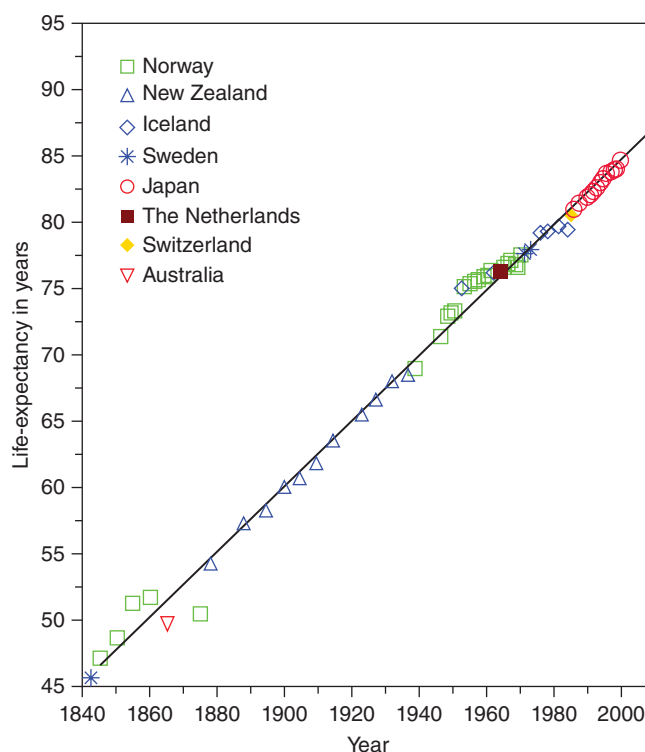


FIGURE 93e-3 Life expectancy in most advanced nations, 1800–2000, females. (From J Oeppen, JW Vaupel: *Science* 296:1029, 2002.)

about the mortality transition—in several African countries, the prevalence of AIDS has been high enough to cause life expectancy to fall below the levels of 1980. Though none has so far reached a scale to rival the AIDS epidemic, periodic outbreaks of new influenza viruses or “emerging infectious” agents remind us that infectious diseases could again come to the fore. Progress against chronic disease is also reversible: In Russia and some other countries that formed part of the Soviet Union before 1992, life expectancy for men has been declining, now reaching levels below those of men in South Asia. Much of the gap between Russian and Western European men is explainable by much greater heart disease and injuries among the former.

DEPENDENCY AND CAREGIVING RATIOS

Ratios of different age groups provide useful though crude indicators of potential demands on resources and resource availability. One set of ratios, known variously as dependency or support ratios, compare the age groups who are most likely to be in the labor force with the age groups typically dependent on the productive capacity of those working—the young and the old, or just the old. A commonly used ratio is the number of persons age 15–64 per persons age 65 and older. Even though many in some countries do not enter the labor force until significantly older than age 15, retire before age 65, or work past age 65, the ratios do summarize important facts, especially in countries where financial support for the retired comes partially or mainly from those currently in the labor force through either a formal pension system or through informal support from the family. While many countries still have very basic pension systems with incomplete coverage, in Europe public pensions are quite generous, and these countries face dramatic changes in their ratios of working age to older populations. Over the next 40 years, Western Europe faces a drop in the ratio from 4 to 2. In other words, while in crude terms there are today 4 workers supporting the pensions and other costs of each older person, by 2050 there will only be 2. China faces an even steeper drop from 9 persons of working age to only 3, while Japan declines from 3 to just 1. Even in India, projected to become the most populous country, the decline is quite steep from 13 to 5.

The dramatically declining number of workers per older person (however determined) is at the crux of the economic challenge of population aging. The extra years of life that can be considered the crowning achievement in medicine and public health of the last 150 years have to be financed. The economic model of the life cycle assumes that people are economically productive for a limited number of years and that the proceeds of their work during those years have to be smoothed over to finance consumption during less economically productive ages, either within families or by institutions such as the state in order to provide for the young, the old, and the infirm. There are only so many ways to meet the challenge of an extended period of dependency, including increasing the productivity of those in the labor force, saving more, reducing consumption, increasing the number of years worked by increasing the age of retirement, increasing the voluntary nonmonetary productive contributions of the retired, and immigration of very large numbers of young workers into the “old” countries. Pressures to increase retirement ages in industrialized countries and to reduce benefits are increasing. But no single one of these measures can bear the full load of adaptation to population aging, since the changes would have to be so severe and disruptive as to be politically impossible. More likely, there will be some combination of these measures.

Population health and the ability to function at work and in everyday life interact with these population ratios in significant ways. The physical and cognitive capacity to continue to work at older ages is crucial if the age of retirement is raised. Similarly, caregiving often requires significant physical and emotional stamina. Further, healthier older populations require less caregiving and medical services. Just two decades ago, the prevalent view of aging was highly pessimistic. Epidemiologists held that while modern medicine could keep older people alive, nothing much could be done to prevent, delay, or significantly treat the degenerative chronic diseases of aging. The result would be that more and more older people with chronic diseases

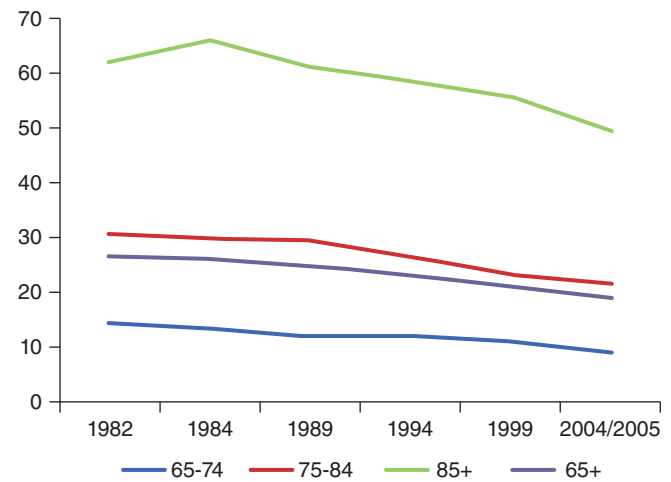


FIGURE 93e-4 Disability prevalence, various years 1982–2005, by age group over 65, United States. (Adapted from KG Manton et al: *Proc Natl Acad Sci U S A.* 103:18374, 2006.)

would be kept from dying, with the consequent piling up of the older people disabled by chronic disease. Surprisingly, between 1984 and about 2000, the prevalence of disability in the 65+ population in the United States declined by about 25%, suggesting that in this respect, aging was more plastic than had been previously believed (Fig. 93e-4). All the causes of this significant shift in disability are not yet understood, but rising levels of education, improved treatment of cardiovascular diseases and cataracts, greater availability of assistive devices, and less physically demanding occupations have been found to contribute. One calculation showed that if the rate of improvement could be maintained until 2050, that the numbers of disabled in the older population could be kept constant in the United States despite the aging of the baby boomers and the older population itself growing older. Unfortunately, the rapid increase in obesity rates could slow and perhaps even reverse this most positive trend. Because of the absence of comparable data in other countries, it is less certain whether the same pattern of improvement in disability rates (with recent deceleration) is occurring outside of the United States. Using estimates and projections of disease prevalence from the Global Burden of Disease Study, the global population of those “dependent and in need of care” is projected to rise from about 350 million in 2010 to over 600 million in 2050. Worldwide, about half of the older persons in need of care (two-thirds of the dependent population age 90 and above) suffer from dementia or cognitive impairment. A global network of longitudinal studies on aging, health, and retirement is now providing comparable data that may allow more definitive projections on disease and disability trends in the future. One estimate (World Alzheimer’s Report 2010) projected that the 36 million people with dementia worldwide in 2010 would increase to 115 million by 2050. The largest increases would occur in low- and middle-income countries where about two-thirds already live. The estimated costs were \$604 billion in 2010 with 70% occurring in North America and Western Europe. A 2013 study using a nationally representative U.S. sample found that annual dementia costs could be as high as \$215 billion. Direct costs of dementia care exceeded the direct costs for either heart disease or cancer. Given the age-associated prevalence of dementia and the expected increase in the older population, coupled with the associated decline in family members able to provide care, countries need to plan for a pandemic of individuals requiring long-term care.

Population aging, and related demographic changes including changes in family structure, could affect the “supply side” of long-term care as well as the demand for care and health care. In every country, long-term care of the disabled and the chronically ill relies heavily on informal, typically unpaid caregivers—usually spouses or children; and increasingly in more developed countries, caregivers for the oldest old are in their 60s and early 70s. Although there are many men who provide care, on the population level, informal caregiving is still

mainly done by women. Because women live longer than men, lack of a spousal caregiver is especially likely to be a problem for older women. Both men and women have fewer children on whom they can call for informal caregiving, because of the worldwide decline in fertility rates. An increasing proportion of older men in Europe and North America have spent much or all of their adult lives apart from their biological children. Lower fertility rates, delayed marriage, and increasing divorce rates mean that people approaching old age may be less likely to have close ties with daughters and daughters-in-law—the adults who have in the past been the most common caregivers apart from spouses. Adult women who in the past have provided uncompensated care (and much other essential volunteer work) are now more likely than in the past to be working for pay and thus have fewer hours to devote to the unpaid roles.

These broad demographic and economic trends do not dictate particular social adaptations or policy responses, of course. One can imagine many different responses to the challenges of caring for the disabled—increased reliance on home health agencies and assisted living communities, “naturally occurring retirement communities” in which neighbors fulfill many of the roles once reserved for close kin; private or even publicly financed direct payments to compensate formerly unpaid family caregivers (a reform that has proved very popular in Germany). These and other responses to the challenge of long-term care are being tested in aging countries, and continued experimentation will no doubt be needed.

THE EPIDEMIOLOGIC TRANSITION—CHANGES IN THE BURDEN OF DISEASE AND RISK FACTORS

The secular improvements in ages at death have been accompanied by changes in causes of death. In the broadest terms, the proportion of deaths due to infectious disease and conditions associated with pregnancy and delivery has fallen, and the proportion due to chronic, noncommunicable diseases, such as heart and cerebrovascular diseases, diabetes, cancers, and age-related neurodegenerative diseases such as Alzheimer’s and Parkinson’s diseases, has increased steadily and is expected to continue to increase. [Figure 93e-5](#) shows results from an international comparative project that drew on a wide variety

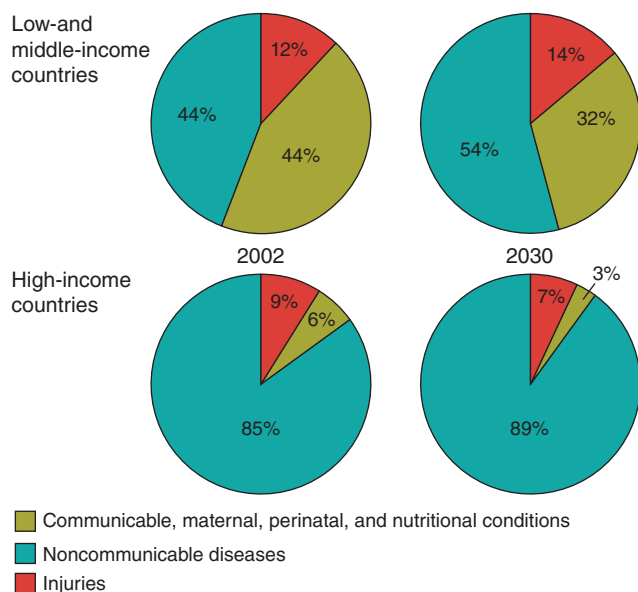


FIGURE 93e-5 Leading causes of burden of illness in world regions 2002 and projected for 2030. (Adapted from CD Mathers, D Loncar. *PLoS Med* 3:e442, 2006.)

TABLE 93e-3 RANKING OF DISEASES AND CONDITIONS CAUSING THE GREATEST BURDEN OF DISEASE WORLDWIDE IN 1990 AND 2010

	1990	2010
1	Lower respiratory infections	Ischemic heart disease
2	Diarrheal diseases	Lower respiratory infections
3	Preterm birth complications	Cerebrovascular disease
4	Ischemic heart disease	Diarrheal diseases
5	Cerebrovascular disease	HIV/AIDS
6	COPD	Low back pain
7	Malaria	Malaria
8	Tuberculosis	Preterm birth complications
9	Protein energy malnutrition	COPD
10	Neonatal encephalopathy	Road injury

Note: “Burden of disease” takes into account years of life lost due to death from this cause and also a weighted estimate for years spent with disability, pain, or impairments due to the condition. These estimates were aggregated from many different national reporting systems and special surveys or surveillance systems, with adjustments for incomplete coverage and different reporting schemes as part of the Global Burden of Disease Study 2010 updating previous global estimates.

Abbreviation: COPD, chronic obstructive pulmonary disease.

Source: CJL Murray et al. *Lancet* 380:2197, 2013, Fig. 5.

of data sources to provide estimates of the global burden of disease at the beginning of this century, with projections to future years based on recent trends in disease prevalence and demographic rates. Burden of disease in these pie charts is a composite measure, one that takes into account both the number of deaths due to a particular disease or condition and the timing of such deaths—an infant death represents a loss of more potential life-years lived than does the death of a very old person. Nor is death the only outcome that matters; most diseases or conditions cause significant disability and suffering even when nonfatal, so this measure of burden captures nonfatal outcomes using statistical weighting. As [Table 93e-3](#) shows, the “modern plagues” of chronic noncommunicable diseases are already among the leading causes of premature death and disability even in low-income countries. This is due to a mix of factors—lower fertility rates mean fewer infants and children at prime ages of susceptibility to infections; more people reaching older ages when chronic disease incidence is high; and often changing incidence rates due to increased exposure to tobacco, Western diets, and inactivity. Noncommunicable diseases, once thought of as “diseases of affluence,” are projected to account for more than half of the disease burden even in low- and middle-income countries by the year 2030 ([Fig. 93e-5](#)).

SUMMARY

Population aging is a global phenomenon with profound short- and long-term implications for health and long-term care needs, and indeed for the economic and social well-being of nations. The timing and context of aging vary across and within world regions and countries; the industrialized nations became wealthy before they aged significantly, while many of the low-resource regions will age before they reach high-income levels. The variation at both the population level and individual levels indicates that there is much flexibility in successful aging, but meeting the challenges will require advance planning and preparation. The extent to which research can find solutions that reduce physical and cognitive disability at older ages will determine how countries cope with this fundamental transformation.

94e The Biology of Aging

Rafael de Cabo, David G. Le Couteur

THE IMPACT OF AGING ON MEDICINE

Aging and old age are among the most significant challenges facing medicine this century. The aging process is the major risk factor underlying disease and disability in developed nations, and older people respond differently to therapies developed for younger adults (usually with less effectiveness and more adverse reactions). Modern medicine and healthier lifestyles have increased the likelihood that younger adults will now achieve old age. However, this has led to rapidly increasing numbers of older people, often encumbered with age-related disorders that are predicted to overwhelm health care systems. Improved health in old age and further extension of the human healthspan are now likely to result primarily from increased understanding of the biology of aging, age-related susceptibility to disease, and modifiable factors that influence the aging process.

Definitions of Aging Aging is easy to recognize but difficult to define. Most definitions of aging indicate that it is a progressive process associated with declines in structure and function, impaired maintenance and repair systems, increased susceptibility to disease and death, and reduced reproductive capacity. There are both statistical and phenotypic components to aging. As recognized by Gompertz in the nineteenth century, aging in humans is associated with an exponential risk of mortality with time (Fig. 94e-1), although it is now realized that this plateaus in extreme old age because of healthy survivor bias. The phenotypic components of aging include structural and functional changes that are separated, somewhat artificially, into either primary aging changes (e.g., sarcopenia, gray hair, oxidative stress, increased peripheral vascular resistance) or age-related disease (e.g., dementia, osteoporosis, arthritis, insulin resistance, hypertension).

Definitions of aging rarely acknowledge the possibility that some of those biological and functional changes with aging might be adaptive or even reflect improvement and gain. Nor do they emphasize the effect of aging on responses to medical treatments. Old age is associated with increased vulnerability to many perturbations, including therapeutic interventions. This is a critical issue for clinicians; the problem with aging would be more limited if our disease-specific therapies retained their balance of risk to benefit into old age.

Aging and Disease Susceptibility Old age is the major independent risk factor for chronic diseases (and associated mortality) that are most prevalent in developed countries such as cardiovascular disease, cancers, and neurodegenerative disorders (Fig. 94e-2). Consequently,

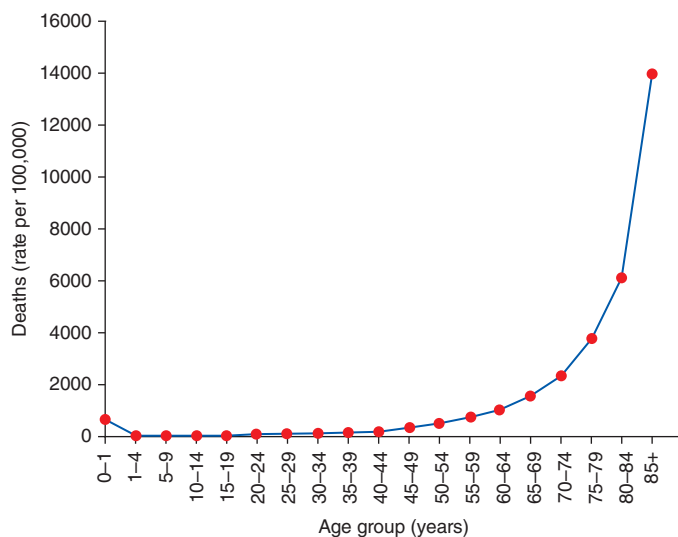


FIGURE 94e-1 The rates of death in the United States (2010) showing exponential increase in mortality risk with chronologic age.

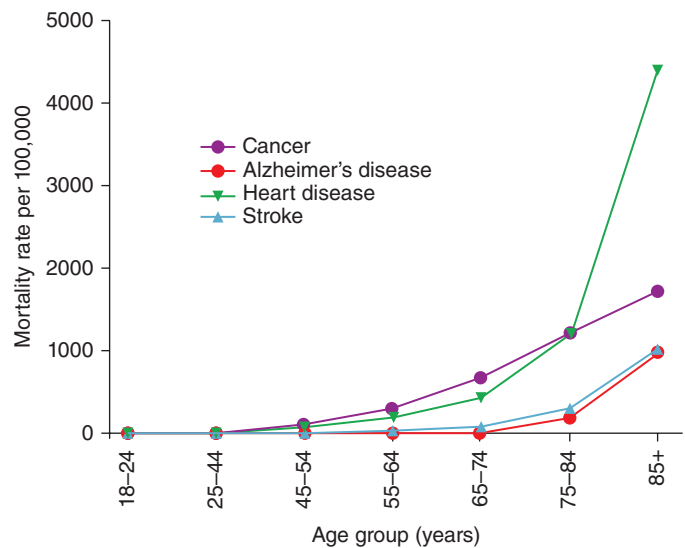


FIGURE 94e-2 The rates of most common chronic diseases and related mortality increase with old age. (Data from USA 2008–2010 CDC.)

older people have multiple comorbidities, usually in the range of 5 to 10 illnesses per person.

Disease in older people is typically multifactorial with a strong component related to the underlying aging process. For example, in younger patients with dementia, Alzheimer's disease is a single disorder confirmed by examining brain tissue for plaques and tangles containing amyloid and tau proteins. However, the vast majority of people with dementia are elderly, and here the association between typical Alzheimer's neuropathology and dementia becomes less definitive. In the oldest-old, the prevalence of Alzheimer's-type brain pathology is similar in people with and without clinical features of dementia. On the other hand, brains of older people with dementia usually show mixed pathology, with evidence of Alzheimer's pathology along with features of other dementias such as vascular lesions, Lewy bodies, and non-Alzheimer's tauopathy. Typical aging changes, such as microvascular dysfunction, oxidative injury, and mitochondrial impairment, underlie many of the pathologic changes.

The Longevity Dividend *Compression of morbidity* refers to the concept that the burden of lifetime illness might be compressed by medical interventions into a shorter period before death without necessarily increasing longevity. However, continuing development of successful therapeutic and preventative interventions focusing on individual diseases is less effective in older people because of multiple comorbidities, complications of overtreatment, and competing causes of death. Therefore, it has been proposed that further gains in healthspan and life expectancy will be achieved by a single intervention that delays aging and age-related disease susceptibility, rather than multiple treatments each targeting different individual age-related illnesses. This is called the *longevity dividend* and is driving an explosion of research into aging biology and, more importantly, interventions (genetic, pharmaceutical, and nutritional) that influence the rate of aging and delay age-related disease.

EVOLUTIONARY MECHANISMS FOR AGING

At the most basic level, living things have only two approaches to maintain their existence: immortality or reproduction. In a changing environment, reproduction combined with a finite lifespan has proved to be the successful strategy. Of course, finite lifespan is not the same as aging, although aging, by definition, contributes to a finite lifespan.

Many evolutionary theories related to aging are linked by their attempts to explain this interaction between reproduction and longevity (Fig. 94e-3). Most mainstream aging theories stem from the fact that evolution is driven by early reproductive success, whereas there is minimal selection pressure for late-life reproduction or postreproductive

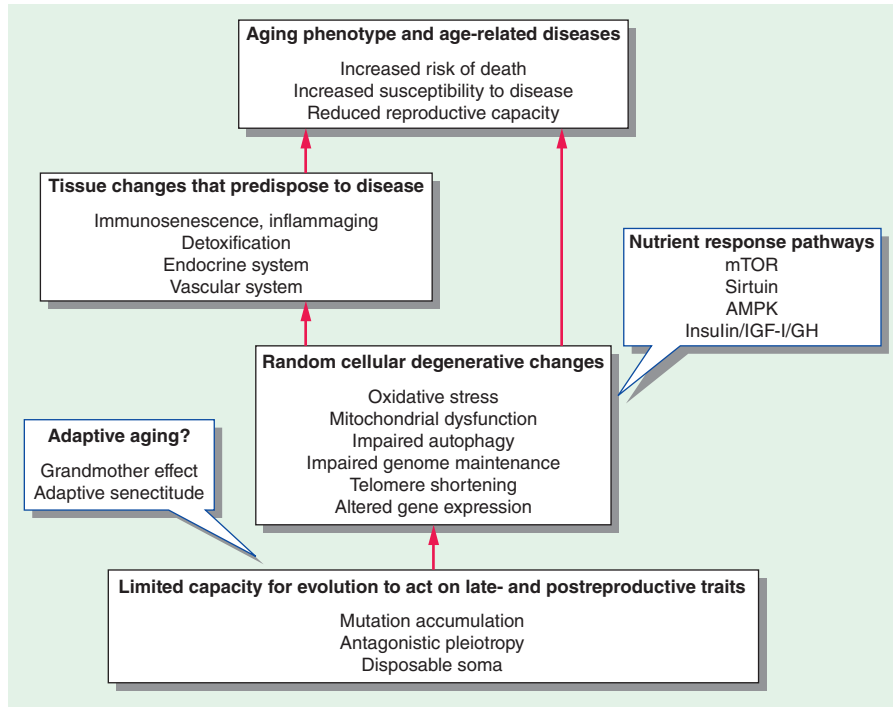


FIGURE 94e-3 Schema linking evolution and cellular and tissue changes with aging. The call-out blue boxes indicate factors that might delay the aging process including nutrient response pathways and, possibly, adaptive evolutionary effects.

survival. Aging is seen as the random degeneration resulting from the inability of evolution to prevent it, i.e., the nonadaptive consequence of evolutionary “neglect.” This conclusion is supported by studies that restricted reproduction to later life in the fruit fly, *Drosophila melanogaster*, thus permitting natural selection to operate on later life traits and leading to an increase in longevity. There are some species of plants and animals that do not appear to age, or at least they undergo an extremely slow aging process, termed “negligible senescence.” The mortality rates of these species are relatively constant with time, and they do not display any obvious phenotypic changes of aging. Conversely, there are some living things that undergo programmed death immediately after reproduction, such as annual plants and semelparous animals (Fig. 94e-4). However, many other living things from yeast to humans undergo a gradual aging process leading to death

that is surprisingly similar at the cellular and biochemical level across taxa. Some of the major classical evolutionary theories of aging include the following:

- *Programmed death.* The first evolutionary theory of aging was proposed by Weissman in 1882. This theory states that aging and death are programmed and have evolved to remove older animals from the population so that environmental resources such as food and water are freed up for younger members of the species.
- *Mutation accumulation.* This theory was proposed by Medawar in 1952. Natural selection is most powerful for those traits that influence reproduction in early life, and therefore, the ability of evolution to shape our biology declines with age. Germline mutations that are deleterious in later life can accumulate simply because natural selection cannot act to prevent them.
- *Antagonistic pleiotropy.* George C. Williams extended Medawar’s theory when he proposed that evolution can allow for the selection of genes that are pleiotropic, i.e., beneficial for survival and reproduction in early life, but harmful in old age. For example, genes for sex hormones are necessary for reproduction in early life but contribute to the risk of cancer in old age.
- *Life history theory.* Evolution is influenced by the way that limited resources are allocated to all aspects of life including development, sexual maturation, reproduction, number of offspring, and senescence and death. Therefore, “trade-offs” occur between these phases of life. For example, in a hostile environment, survival is highest for those species that have large numbers of offspring and short lifespan, whereas in a safe and abundant environment, survival is highest for those species that invest resources in a smaller number of offspring and a longer life.
- *Disposable soma theory.* Kirkwood and Holliday in 1979 combined many of these ideas in the disposable soma theory of aging. There are finite resources available for the maintenance and repair of both germ and soma cells, so there must be a trade-off between germ cells (i.e., reproduction) and soma cells (i.e., longevity and aging). The soma cells are disposable from an evolutionary perspective, so they accumulate damage that causes aging while resources are preferentially diverted to the maintenance and repair of the germ cells. For example, the longevity of the nematode worm, *Caenorhabditis elegans*, is increased when its germ cells are ablated early in life.





Negligible senescence	Rapid death postreproduction (semelparous animals, annual plants)
 Rougheye rockfish  Bristlecone pine	 Pacific salmon  Sunflower

FIGURE 94e-4 The typical features of aging (aging phenotype and exponential increase in risk of death) are not universal findings in living things. Some living things (e.g., rockfish and the bristlecone pine, sometimes called the Methuselah tree) undergo negligible senescence, whereas others die almost immediately after reproduction is completed (e.g., semelparous animals and annual plants).

All of these theories assume that natural selection has negligible or negative influences on aging. Some postmodern ideas propose that aspects of aging might be adaptive and raise the possibility that evolution can act on the aging process in a positive way. These include the following:

- **Grandmother hypothesis.** The grandmother hypothesis proposed by Hamilton in 1966 describes how evolution can enhance old age. In some animals, including humans, the survival of multiple, dependent offspring is beyond the capacity and resources of a single parent. In this situation, the presence of a long-lived grandmother who shares in the care of her grandchildren can have a major impact on their survival. These children share some of the genes of their grandmother including those that promoted their grandmother's longevity.
- **Mother's curse.** Mitochondrial dysfunction is a key component of the aging process. Mitochondria contain their own DNA and are only passed on from mother to child because sperm cells contain almost no mitochondria. Therefore, natural selection can only act on the evolution of mitochondrial DNA in females. The "mother's curse" of the maternal inheritance of mitochondrial DNA might explain why females live longer and age more slowly than males.
- **Adaptive senectitude.** Many traits that are harmful in younger humans such as obesity, hypertension, and oxidative stress paradoxically appear to be associated with greater survival and function in very old people. Perhaps driven by the grandmother effect, this might represent "adaptive senectitude" or "reverse antagonistic pleiotropy," whereby some traits that are harmful in young people become beneficial in older people.

CELLULAR PROCESSES THAT ACCOMPANY AGING

There are many cellular processes that change with aging. These are generally considered to be degenerative and stochastic or random changes that reflect some sort of time-dependent damage (Fig. 94e-3). Whether any of these is the root cause of aging is unknown, but they all contribute to the aging phenotype and disease susceptibility.

Oxidative Stress and the Free Radical Theory of Aging Free radicals are chemical species that are highly reactive because they contain unpaired electrons. Oxidants are oxygen-based free radicals that include the hydroxyl free radical, superoxide, and hydrogen peroxide. Most cellular oxidants are waste products generated by mitochondria during the production of ATP from oxygen. More recently, the role of oxidants in cellular signaling and inflammatory responses has been recognized. Unchecked, oxidants can generate chain reactions leading to widespread damage to biological molecules. Cells contain numerous antioxidant defense mechanisms to prevent such oxidative stress including enzymes (superoxide dismutase, catalase, glutathione peroxidase) and chemicals (uric acid, ascorbate). In 1956, Harman proposed the "free radical theory of aging," whereby oxidants generated by metabolism or irradiation are responsible for age-related damage. It is now well established that old age in most species is associated with increased oxidative stress, for example to DNA (8-hydroxyguanosine derivatives), proteins (carbonyls), lipids (lipoperoxides, malondialdehydes), and prostaglandins (isoprostanes). Conversely, many of the cellular antioxidant defense mechanisms, including the antioxidant enzymes, decline in old age. The free radical theory of aging has spawned numerous studies of supplementation with antioxidants such as vitamin E to delay aging in animals and humans. Unfortunately, meta-analyses of human clinical trials performed to treat and prevent various diseases with antioxidant supplements indicate that they have no effect on, or may even increase, mortality.

Mitochondrial Dysfunction Aging is characterized by altered mitochondrial production of ATP and oxygen-derived free radicals. This leads to a vicious cycle mediated by accumulation of oxidative injury to mitochondrial proteins and DNA. With age, the number of mitochondria in cells decreases, and there is an increase in their size (megamitochondria) associated with other structural changes including vacuolization and disrupted cristae. These morphologic aging changes are linked with decreased activity of mitochondrial complexes I, II, and

IV and decreased ATP production. Of all of the complexes involved in ATP production, the activity of complex IV (COX) is usually reported to be most impaired in old age. Reduced energy production is linked with generation of hydrogen peroxide and superoxide radicals leading to oxidative injury to mitochondrial DNA and accumulation of carbonylated mitochondrial proteins and mitochondrial lipoperoxides. As well as being implicated in the aging process, common geriatric syndromes including sarcopenia, frailty, and cognitive impairment are associated with mitochondrial dysfunction.

Telomere Shortening and Replicative Senescence Cells that are isolated from animal tissue and grown in culture only divide for a certain number of times before entering a senescent phase. This number of divisions is known as the Hayflick limit and tends to be less in cells isolated from older animals compared to younger animals. It has been suggested that aging in vivo might in part be secondary to some cells ceasing to divide because they have reached their Hayflick limit. One mechanism for replicative senescence relates to telomeres. Telomeres are repeat sequences of DNA at the end of linear chromosomes that shorten by around 50–200 base pairs during each cell division by mitosis. Once telomeres become too short, cell division can no longer occur. This mechanism contributes to the Hayflick limit and has been called the cellular clock. There are some studies that suggest that the length of telomeres in circulating leukocytes (leukocyte telomere length [LTL]) decreases with age in humans. However, the aging process also occurs in tissues that do not undergo repeated cell division such as neurons.

Altered Gene Expression, Epigenetics, and microRNA There are changes in the expression of many genes and proteins during the aging process. These changes are complicated and vary between species and tissue. Such heterogeneity reflects increasing dysregulation of gene expression with age while appearing to exclude a programmed and/or uniform response. With old age, there are often reductions in the expression of genes and proteins associated with mitochondrial function and increased expression of those involved with inflammation, genome repair, and oxidative stress. There are several factors controlling the regulation of gene and protein expression that change with aging. These include the epigenetic state of the chromosomes (e.g., DNA methylation and histone acetylation) and microRNAs (miRNA). DNA methylation correlates with age, although the pattern of change is complex. Histone acetylation is regulated by many enzymes including SIRT1, a protein that has marked effects on aging and the response to dietary restriction in many species. miRNAs are a very large group of noncoding lengths of RNA (18–25 nucleotides) that inhibit translation of multiple different mRNAs through binding their 3' untranslated regions (UTRs). The expression of miRNAs usually decreases with aging and is altered in some age-related diseases. Specific miRNAs linked with aging pathways include miR-21 (associated with target of rapamycin pathway) and miR-1 (associated with insulin/insulin-like growth factor 1 pathway).

Impaired Autophagy There are a number of ways that cells can remove damaged macromolecules and organelles, often generating cellular energy as a byproduct. Intracellular degradation is undertaken by the lysosomal system and the ubiquitin proteasomal system. Both are impaired with aging, leading to the accumulation of waste products that alter cellular functions. Such waste products include lipofuscin, a brown autofluorescent pigment found within lysosomes of most cells in old age and often considered to be one of the most characteristic histologic features of aging cells. They also include aggregated proteins characteristic of age-related neurodegenerative diseases (e.g., tau, β -amyloid, α -synuclein). Lysosomes are organelles that contain proteases, lipases, glycases, and nucleotidases that degrade intracellular macromolecules, membrane components, organelles, and some pathogens through a process called autophagy. The lysosomal process most impaired with aging is macroautophagy, which is regulated by numerous autophagy-related genes (ATGs). Old age is associated with some impairment in chaperone-mediated autophagy, whereas the effect of aging on the third lysosomal process, microautophagy, is unclear.

AGING CHANGES IN SPECIFIC TISSUES THAT PREDISPOSE TO DISEASE

Aging changes in some tissues increase susceptibility to age-related disease as a secondary or downstream phenomenon (Fig. 94e-3). In humans, this includes, but is not limited to, the immune system (leading to increased infections and autoimmunity), hepatic detoxification (leading to increased exposure to disease-inducing endobiotics and xenobiotics), the endocrine system (leading to hypogonadism and bone disease), and the vascular system (leading to segmental or global ischemic changes in many tissues).

Inflammaging and Immunosenescence Old age is associated with increased background levels of inflammation including blood measurements of C-reactive protein (CRP), erythrocyte sedimentation rate (ESR), and cytokines such as interleukin 6 (IL-6) and tumor necrosis factor α (TNF- α). This has been termed *inflammaging*. T cells (particularly naïve T cells) are less numerous because of age-related atrophy of the thymus, whereas B cells overproduce autoantibodies, leading to the age-related increase in autoimmune diseases and gammopathies. Thus older people are generally considered to be immunocompromised and have reduced responses to infection (fever, leukocytosis) with increased mortality.

Detoxification and the Liver Old age is associated with impaired detoxification of various disease-causing endobiotics (e.g., lipoproteins) and xenobiotics (e.g., neurotoxins, carcinogens), leading to increased systemic exposure. In humans, the liver is the major organ for the clearance of such toxins. Hepatic clearance of many substrates is reduced in old age as a consequence of reduced hepatic blood flow, impaired hepatic microcirculation, and in some cases, reduced expression of xenobiotic metabolizing enzymes. These changes in hepatic detoxification also increase the likelihood of increased blood levels of, and adverse reactions to, medications.

Endocrine System Hormonal changes with aging have been a focus of aging research for over a century, partly because of the erroneous belief that supplementation with sex hormones will delay aging and rejuvenate older people. There are age-related reductions in sex steroids secondary to hypogonadism and, in females, menopause. Age-related declines in growth hormone and dehydroepiandrosterone (DHEA) are well established, as is the increase in insulin levels and associated insulin resistance. These hormonal changes contribute to some features of aging such as sarcopenia and osteoporosis, which may be delayed by hormonal supplementation. However, adverse effects of long-term hormonal supplementation outweigh any potential beneficial effects on lifespan.

Vascular Changes There is a continuum from vascular aging through to atherosclerotic disease, present in many, but not all, older people. Vascular aging changes overlap with the early stages of hypertension and atherosclerosis, with increasing arterial stiffness and vascular resistance. This contributes to myocardial ischemia and strokes but also appears to be associated with geriatric conditions such as dementia, sarcopenia, and osteoporosis. In these conditions, impaired exchange between blood and tissues is a common pathogenic factor. For example, the risk of Alzheimer's disease and dementia is increased in patients with risk factors for vascular disease, and there is pathologic evidence for microvascular changes in postmortem studies of brains of people with established Alzheimer's disease. Similarly, strong epidemiologic links have been found between osteoporosis and standard vascular risk factors, whereas there are significant age-associated changes in the microcirculation of osteoporotic bone. Sarcopenia might also be related to the effects of age on the muscle vasculature, which is altered in old age. The sinusoidal microcirculation of the liver becomes markedly altered during aging (pseudocapillarization), which influences hepatic uptake of lipoproteins and other substrates. In fact, it has often been overlooked that in his original exposition of the free radical theory of aging, Harman proposed that the primary target of oxidative stress was the vasculature and that many aging changes were secondary to impaired exchange across the damaged blood vessels.

GENETIC INFLUENCES ON AGING

There is variability in aging and lifespan in populations of genetically identical species such as mice that are housed in the same environment. Moreover, the heritability of lifespan in human twin studies is estimated to be only 25% (although there is stronger hereditary contribution to extreme longevity). These two observations indicate that the cause of aging is unlikely to lie only within the DNA code. On the other hand, genetic studies initially undertaken in the nematode worm *C. elegans* and, more recently, in models from yeast to mice have shown that manipulating genes can have profound effects on the rate of aging. Perhaps surprisingly, this can often be generated by variability in *single* genes, and for some genetic mechanisms, there is very strong evolutionary conservation.

Genetic Progeroid Syndromes There are a few very rare, genetic premature aging conditions that are called progeroid syndromes. These conditions recapitulate some, but not all, age-related diseases and senescent phenotypes. They are mostly caused by impairment of genome and nuclear maintenance. These syndromes include the following:

- *Werner's syndrome*. This is an autosomal recessive condition caused by a mutation in the *WRN* gene. This gene codes for a RecQ helicase, which unwinds DNA for both repair and replication. It is typically diagnosed in teen years, and there is premature onset of atherosclerosis, osteoporosis, cancers, and diabetes, with death by age of 50 years.
- *Hutchinson-Gilford progeria syndrome (HGPS)*. This usually occurs as a de novo, noninherited mutation in the lamin A gene (*LMNA*), leading to an abnormal protein called progerin. *LMNA* is required for the nuclear lamina, which provides structural support to the nucleus. There are marked development changes obvious in infancy with subsequent onset of atherosclerosis, kidney failure, and scleroderma-like features and death during the teen years.
- *Cockayne syndrome*. This includes a number of autosomal recessive disorders with features such as impaired neurologic growth, photosensitivity (xeroderma pigmentosa), and death during childhood years. These disorders are caused by mutations in the genes for DNA excision repair proteins, *ERCC-6* and *ERCC-8*.

Gene Studies in Long-Lived Humans The main genes that have been consistently associated with increased longevity in human candidate gene studies are *APOE* and *FOXO3A*. ApoE is an apoprotein found in chylomicrons, whereas the ApoE4 isoform is a risk factor for Alzheimer's disease and cardiovascular disease, which might explain its association with reduced lifespan. *FOXO3A* is a transcription factor involved in the insulin/IGF-I pathway, and its homolog in *C. elegans*, *daf16*, has a substantial impact on aging in these nematodes. Genome-wide association studies (GWAS) of centenarians have confirmed the association of longevity with *APOE*. GWAS have been used to identify a range of other single nucleotide polymorphisms (SNPs) that might be associated with longevity including SNPs in the sirtuin genes and the progeroid syndrome genes, *LMNA* and *WRN*. Gene set analysis of GWAS studies has shown that both the insulin/IGF-I signaling pathway and the telomere maintenance pathway are associated with longevity.

Of particular interest are people with Laron-type dwarfism. These people have mutations in the growth hormone receptor, which causes severe growth hormone resistance. In mice, similar knockout of the growth hormone receptor (GHRKO mice, Methuselah mice) is associated with extremely long life. Therefore, subjects with Laron's syndrome have been carefully studied, and it was found that they have very low rates of cancer and diabetes mellitus and, possibly, longer lives.

Nutrient-Sensing Pathways Many living things have evolved to respond to periods of nutritional shortage and famine by increasing cellular resilience and delaying reproduction until food supply becomes abundant once again. This increases the chances of reproductive success and survival of offspring. Lifelong food shortage, often termed *caloric*

restriction (or dietary restriction) increases lifespan and delays aging in many animals, probably as a nonadaptive side effect of this famine response. Many of the genes and pathways that regulate the way that cells respond to nutritional undersupply have been identified, initially in yeast and *C. elegans*. In general, manipulation of these pathways (through genetic knockout or overexpression or pharmacologic agonists and antagonists) alters the aging benefits of caloric restriction and, in some cases, the lifespan of animals on normal diets. These pathways are all very influential cellular “switches” that control a wide range of key functions including protein translation, autophagy, mitochondrial function and bioenergetics, and the cellular metabolism of fats, proteins, and carbohydrates. The discovery of these nutrient-sensing pathways has led to targets for pharmacologic extension of lifespan. The main nutrient-sensing pathways that influence aging and responses to caloric restriction include the following:

- **SIRT1.** The sirtuins are a class of histone deacetylases that inhibit gene expression. The key nutrient-sensing member of this class in mammals is SIRT1. The activity of SIRT1 is regulated by levels of reduced nicotinamide adenine dinucleotide (NAD⁺), which are increased when cellular energy stores are depleted. Important downstream targets include PGC1 α and NRF2, which act on mitochondrial biogenesis.
- **Target of rapamycin (TOR, or mTOR in mammals).** mTOR is activated by branched-chain amino acids, providing a link to dietary protein intake. It is a complex that orchestrates two pathways (TORC1 and TORC2). Key downstream targets of mTOR of relevance to aging include the tuberous sclerosis protein (TSC) and 4EBP1, which influence protein production.
- **5' Adenosine monophosphate-activated protein kinase (AMPK).** AMPK is activated by increased levels of AMP, which reflect cellular energy status.
- **Insulin signaling and IGF-I/growth hormone.** These two pathways are usually considered together because they are the same in lower animals and have diverged only in higher animals. Insulin responds to carbohydrate intake. An important downstream target for this pathway is a transcription factor called daf16 in worms and FOXO in mammals and the fruit fly.

Mitochondrial Genes Mitochondrial function is influenced by genes located both in the mitochondria (mtDNA) and the nucleus. mtDNA is considered to have a prokaryotic origin and is highly conserved across taxa. It forms a circular loop of 16,569 nucleotides in humans. Aging is associated with increased frequency of mutations in mtDNA as a consequence of its high exposure to oxygen-derived free radicals and relatively inefficient DNA repair machinery. Nuclear DNA encodes approximately 1000–1500 genes for mitochondrial function, including genes involved with oxidative phosphorylation, mitochondrial metabolic pathways, and enzymes required for biogenesis. These genes are thought to have originated in mtDNA but subsequently translocated to the nucleus, and unlike mtDNA genes, their sequence is stable with aging.

Genetic manipulation of mitochondrial genes in animals influences aging and lifespan. In *C. elegans*, many mutants with defective electron transfer chain function have increased lifespan. The mtDNA “mutator” mice, which lack the mtDNA proofreading enzyme, have increased mtDNA mutations and premature aging, whereas overexpression of mitochondrial uncoupling proteins leads to longer lifespan. In humans, hereditary variability in mtDNA is associated with diseases (mitochondriopathies such as Leigh’s disease) and aging. For example, in Europeans, mitochondrial DNA haplogroup J (haplogroups are combinations of genetic variants that exist in specific populations) is associated with longevity, and haplogroup D is over-represented in Asian centenarians.

STRATEGIES THAT INCREASE HEALTHSPAN AND DELAY AGING

Aging is an intrinsic feature of human life whose manipulation has fascinated humans ever since becoming conscious of their own existence. Recent reports and scientific literature are shaping a picture

where different dietary restriction regimes and exercise interventions may improve healthy aging in laboratory animals. Several long-term experimental interventions (e.g., resveratrol, rapamycin, spermidine, metformin) may open doors for corresponding pharmacologic strategies. Surprisingly, most of the effective aging interventions proposed converge on only a few molecular pathways: nutrient signaling, mitochondrial proteostasis, and the autophagic machinery.

Lifespan is inevitably accompanied by functional decline, a steady increase in a plethora of chronic diseases, and ultimately death. For millennia, it has been a dream of mankind to prolong both lifespan and healthspan. Developed countries have profited from the medical improvements and their transfer to public health care systems, as well as from better living conditions derived from their socioeconomic power, to achieve remarkable increases in life expectancy during the last century. In the United States, the percentage of the population age 65 years or older is projected to increase from 13% in 2010 to 19.3% in 2030. However, old age remains the main risk factor for major life-threatening disorders, and the number of people suffering from age-related diseases is anticipated to almost double over the next two decades. The prevalence of age-related pathologies represents a major threat as well as an economic burden that urgently needs effective interventions.

Molecules, drugs, and other interventions that might decelerate aging processes continue to raise interest among both the general public and scientists of all biologic and medical fields. Over the past two decades, this interest has taken root in the fact that many of the molecular mechanisms underlying aging are interconnected and linked with pathways that cause disease, including cancer and cardiovascular and neurodegenerative disorders. Unfortunately, among the many proposed aging interventions, only a few have reached a certain age themselves. Results often lack reproducibility because of a simple inherent problem: interventions in aging research take a lifetime. Experiments lasting the lifetime of animal models are prone to develop artifacts, increasing the possibilities and time windows for experimental discrepancies. Some inconsistencies in the field arise from overinterpreting lifespan-shortening models and scenarios as being related to accelerated aging.

Many substances and interventions have been claimed to be antiaging throughout history and into the present. In the following sections, interventions will be restricted to those that meet the following highly selective criteria: (1) promotion of lifespan and/or healthspan, (2) validation in at least three model organisms, and (3) confirmation by at least three different laboratories. These interventions include (1) caloric restriction and fasting regimens, (2) some pharmacotherapies (resveratrol, rapamycin, spermidine, metformin), and (3) exercise.

Caloric Restriction One of the most important and robust interventions that delays aging is caloric restriction. This outcome has been recorded in rodents, dogs, worms, flies, yeasts, monkeys, and prokaryotes. Calorie restriction is defined as a reduction in the total caloric intake, usually of about 30% and without malnutrition. Caloric restriction reduces the release of growth factors such as growth hormone, insulin, and IGF-I, which are activated by nutrients and have been shown to accelerate aging and enhance the probability for mortality in many organisms. Yet the effects of caloric restriction on aging were first discovered by McCay in 1935 long before the effects of such hormones and growth factors on aging were recognized. The cellular pathways that mediate this remarkable response have been explored in many experimental models. These include the nutrient-sensing pathways (TOR, AMPK, insulin/IGF-I, sirtuins) and transcription factors (FOXO in *D. melanogaster* and daf16 in *C. elegans*). The transcription factor Nrf2 appears to confer most of the anticancer properties of caloric restriction in mice, even though it is dispensable for lifespan extension.

Two studies have reported the effects of caloric restriction in monkeys with different outcomes: one study observed prolonged life, while the other did not. However, both studies confirmed that caloric restriction increases healthspan by reducing the risk for diabetes, cardiovascular

disease, and cancer. In humans, caloric restriction is associated with increased lifespan and healthspan. This is most convincingly demonstrated in Okinawa, Japan, where one of the most long-lived human populations resides. In comparison to the rest of the Japanese population, Okinawan people usually combine an above-average amount of daily exercise with a below-average food intake. However, when Okinawan families move to Brazil, they adopt a Western lifestyle that affects both exercise and nutrition, causing a rise in weight and a reduction in life expectancy by nearly two decades. In the Biosphere II project, where volunteers lived together for 24 months undergoing an unforeseen severe caloric restriction, there were improvements in insulin, blood sugar, glycosylated hemoglobin, cholesterol levels, and blood pressure—all outcomes that would be expected to benefit lifespan. Caloric restriction changes many aspects of human aging that might influence lifespan such as the transcriptome, hormonal status (especially IGF-I and thyroid hormones), oxidative stress, inflammation, mitochondrial function, glucose homeostasis, and cardiometabolic risk factors. Epigenetic modifications are an emerging target for caloric restriction.

It must be noted that maintaining caloric restriction and avoiding malnutrition is not only arduous in humans but is also linked with substantial side effects. For instance, prolonged reduction of calorie intake may decrease fertility and libido, impair wound healing, reduce the potential to combat infections, and lead to amenorrhea and osteoporosis.

Although extreme obesity (body mass index [BMI] >35) leads to a 29% increased risk of dying, people with BMI in the overweight range seem to have reduced mortality, at least in population studies of middle-aged and older subjects. People with a BMI in the overweight range seem more able to counteract and respond to disease, trauma, and infection, whereas caloric restriction impairs healing and immune responses. On the other hand, BMI is an insufficient denominator of body and body fat composition. A well-trained athlete may have a similar BMI as an overweight person because of the higher muscle mass density. The waist-to-hip ratio is a much better indicator for body fat and an excellent and stringent predictor of the risk of dying from cardiovascular disease: the lower the waist-to-hip ratio, the lower is the risk.

PERIODIC FASTING How can caloric restriction be translated to humans in a socially and medically feasible way? A whole series of periodic fasting regimens are asserting themselves as suitable strategies, among them the alternate-day fasting diet, the “five:two” intermittent fasting diet, and a 48-h fast once or twice each month. Periodic fasting is psychologically more viable, lacks some of the negative side effects, and is only accompanied by minimal weight loss.

It is striking that many cultures implement periodic fasting rituals, for example Buddhists, Christians, Hindus, Jews, Muslims, and some African animistic religions. It could be speculated that a selective advantage of fasting versus nonfasting populations is conferred by health-promoting attributes of religious routines that periodically limit caloric intake. Indeed, several lines of evidence indicate that intermittent fasting regimens exert antiaging effects. For example improved morbidity and longevity were observed among Spanish home nursing residents who underwent alternate-day fasting. Even rats subjected to alternate-day fasting live up to 83% longer than normally fed control animals, and one 24-h fasting period every 4 days is sufficient to generate lifespan extension.

Repeated fasting and eating cycles may circumvent the negative side effects of sustained caloric restriction. This strategy may even yield effects despite extreme overeating during the nonfasting periods. In a spectacular experiment, mice fed a high-fat diet in a time-restricted manner, i.e., with regular fasting breaks, showed reduced inflammation markers and no fatty liver and were slim in comparison to mice with equivalent total calorie consumption but ad libitum. From an evolutionary point of view, this kind of feeding pattern may reflect mammalian adaptation to food availability: overeating in times of nutrient availability (e.g., after a hunting success) and starvation in between. This is how some indigenous peoples who have avoided

Western lifestyles live today; those who have been investigated show limited signs of age-induced diseases such as cancer, neurodegeneration, diabetes, cardiovascular disease, and hypertension.

Fasting exerts beneficial effects on healthspan by minimizing the risk of developing age-related diseases including hypertension, neurodegeneration, cancer, and cardiovascular diseases. The most effective and rapid repercussion of fasting is reduction in hypertension. Two weeks of water-only fasting resulted in a blood pressure below 120/80 mmHg in 82% of subjects with borderline hypertension. Ten days of fasting cured all hypertensive patients who had been taking antihypertensive medication previously.

Periodic fasting dampens the consequences of many age-related neurodegenerative diseases (Alzheimer’s disease, Parkinson’s disease, Huntington’s disease, and frontotemporal dementia, but not amyotrophic lateral sclerosis in mouse models). Fasting cycles are as effective as chemotherapy against certain tumors in mice. In combination with chemotherapy, fasting protected mice against the negative side effects of chemotherapeutic drugs, while it enhanced their efficacy against tumors. Combining fasting and chemotherapy rendered 20–60% of mice cancer-free when inoculated with highly aggressive tumors like glioblastoma or pancreatic tumors, which have 100% mortality even with chemotherapy. This approach has been attempted in people with some indication that toxicities of chemotherapy are reduced.

Pharmacologic Interventions to Delay Aging and Increase Lifespan

Virtually all obese people know that stable weight reduction will reduce their elevated risk of cardiometabolic disease and enhance their overall survival, yet only 20% of overweight individuals are able to lose 10% of their weight for a period of at least 1 year. Even in the most motivated people (such as the “Cronies” who deliberately attempt long-term caloric restriction in order to extend their lives), long-term caloric restriction is extremely difficult. Thus, focus has been directed at the possibility of developing medicines that replicate the beneficial effects of caloric restriction without the need for reducing food intake (“CR-mimetics,” Fig. 94e-5):

- **Resveratrol.** Resveratrol, an agonist of SIRT1, is a polyphenol that is found in grapes and in red wine. The potential of resveratrol to promote lifespan was first identified in yeast, and it has gathered fame since, at least in part because it might be responsible for the so-called French paradox whereby wine reduces some of the cardiometabolic risks of a high-fat diet. Resveratrol has been reported to increase lifespan in many lower order species such as yeast, fruit flies, worms, and mice on high-fat diets. In monkeys fed a diet high in sugar and fat, resveratrol had beneficial outcomes related to inflammation and cardiometabolic parameters. Some studies in humans have also shown improvements in cardiometabolic function, whereas

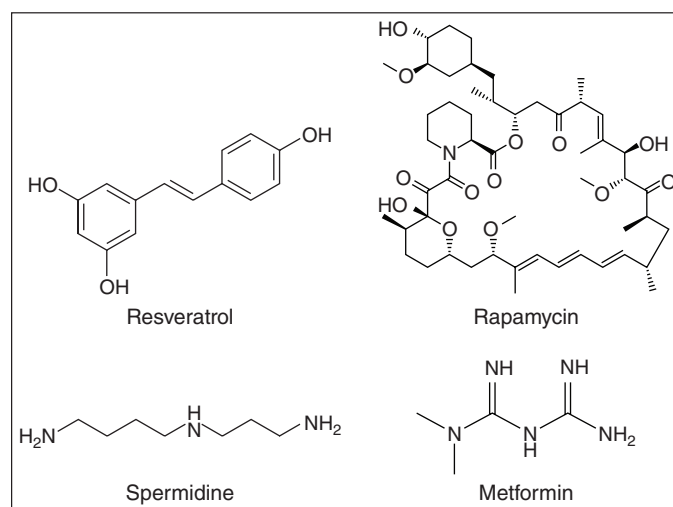


FIGURE 94e-5 Chemical structures of four agents (resveratrol, rapamycin, spermidine, and metformin) that have been shown to delay aging in experimental animal models.

others have been negative. Gene expression studies in animals and humans reveal that resveratrol mimics some of the metabolic and gene expression changes of caloric restriction.

- **Rapamycin.** Rapamycin, an inhibitor of mTOR, was originally discovered on Easter Island (Rapa Nui; hence its name) as a bacterial secretion with antibiotic properties. Before its immersion in the antiaging field, rapamycin already had a longstanding career as an immunosuppressant and cancer chemotherapeutic in humans. Rapamycin extends lifespan in all organisms tested so far, including yeast, flies, worms, and mice. However, the potential utility of rapamycin for human lifespan extension is likely to be limited by adverse effects related to immunosuppression, wound healing, proteinuria, and hypercholesterolemia, among others. An alternative strategy may be intermittent rapamycin feeding, which was found to increase mouse lifespan.
- **Spermidine.** Spermidine is a physiologic polyamine that induces autophagy-mediated lifespan extension in yeast, flies, and worms. Spermidine levels decrease during the life of virtually all organisms including humans, with the stunning exception of centenarians. Oral administration of spermidine and upregulation of bacterial polyamine production in the gut both lead to lifespan extension in short-lived mouse models. Spermidine has also been found to have beneficial effects on neurodegeneration probably by increasing transcription of genes involved in autophagy.
- **Metformin.** Metformin, an activator of AMPK, is a biguanide first isolated from the French lilac that is widely used for the treatment of type 2 diabetes mellitus. Metformin decreases hepatic gluconeogenesis and increases insulin sensitivity. Metformin has other actions including inhibition of mTOR and mitochondrial complex I and activation of the transcription factor SKN-1/Nrf2. Metformin increases lifespan in different mouse strains including female mouse strains predisposed to high incidence of mammary tumors. At a biochemical level, metformin supplementation is associated with reduced oxidative damage and inflammation and mimics some of the gene expression changes seen with caloric restriction.

Exercise and Physical Activity In humans and animals, regular exercise reduces the risk of morbidity and mortality. Given that cardiovascular diseases are the dominant cause of aging in humans but not in mice, the effects on human health may be even stronger than those seen in mouse experiments. An increase in aerobic exercise capacity, which declines during aging, is associated with favorable effects on blood pressure, lipids, glucose tolerance, bone density, and depression in older people. Likewise, exercise training protects against aging disorders such as cardiovascular diseases, diabetes mellitus, and osteoporosis. Exercise is the only treatment that can prevent or even reverse

sarcopenia (age-related muscle wasting). Even moderate or low levels of exercise (30 min walking per day) have significant protective effects in obese subjects. In older people, regular physical activity has been found to increase the duration of independent living.

While clearly promoting health and thus quality of life, regular exercise does not extend lifespan. Furthermore, the combination of exercise with caloric restriction has no additive effect on maximal lifespan in rodents. On the other hand, alternate-day fasting with exercise is more beneficial for the muscle mass than single treatments alone. In nonobese humans, exercise combined with caloric restriction has synergistic effects on insulin sensitivity and inflammation. From the evolutionary perspective, the responses to hunger and exercise are linked: when food is scarce, increased activity is required to hunt and gather.

Hormesis The term *hormesis* describes the, at first sight paradoxical, protective effects conferred by the exposure to low doses of stressors or toxins (or as Nietzsche stated, “What does not kill him makes him stronger”). Adaptive stress responses elicited by noxious agents (chemical, thermal, or radioactive) precondition an organism, rendering it resistant to subsequent higher and otherwise lethal doses of the same trigger. Hormetic stressors have been found to influence aging and lifespan, presumably by increasing cellular resilience to factors that might contribute to aging such as oxidative stress.

Yeast cells that have been exposed to low doses of oxidative stress exhibit a marked antistress response that inhibits death following exposure to lethal doses of oxidants. During ischemic preconditioning in humans, short periods of ischemia protect the brain and the heart against a more severe deprivation of oxygen and subsequent reperfusion-induced oxidative stress. Similarly, the lifelong and periodic exposure to various stressors can inhibit or retard the aging process. Consistent with this concept, heat or mild doses of oxidative stress can lead to lifespan extension in *C. elegans*. Caloric restriction can also be considered to be a type of hormetic stress that results in the activation of antistress transcription factors (Rim15, Gis1, and Msn2/Msn4 in yeast and FOXO in mammals) that enhance the expression of free radical-scavenging factors and heat shock proteins.

CONCLUSIONS

Clinicians need to understand aging biology in order to better manage people who are elderly now. Moreover there is an urgent need to develop strategies based on aging biology that delay aging, reduce or postpone the onset of age-related disorders, and increase functional life and healthspan for future generations. Interventions related to nutritional interventions and drugs that act on nutrient-sensing pathways are being developed and, in some cases, are already being studied in humans. Whether these interventions are universally effective or species/individual specific needs to be determined.

95e Nutrient Requirements and Dietary Assessment

Johanna Dwyer

Nutrients are substances that are not synthesized in sufficient amounts in the body and therefore must be supplied by the diet. Nutrient requirements for groups of healthy persons have been determined experimentally. The absence of essential nutrients leads to growth impairment, organ dysfunction, and failure to maintain nitrogen balance or adequate status of other nutrients. For good health, we require energy-providing nutrients (protein, fat, and carbohydrate), vitamins, minerals, and water. Requirements for organic nutrients include 9 essential amino acids, several fatty acids, glucose, 4 fat-soluble vitamins, 10 water-soluble vitamins, dietary fiber, and choline. Several inorganic substances, including 4 minerals, 7 trace minerals, 3 electrolytes, and the ultratrace elements, must also be supplied by diet.

The amounts of the essential nutrients that are required by individuals differ by age and physiologic state. Conditionally essential nutrients are not required in the diet but must be supplied to individuals who do not synthesize them in adequate amounts, such as those with genetic defects, those with pathologic conditions such as infection or trauma with nutritional implications, and developmentally immature infants. For example, inositol, taurine, arginine, and glutamine may be needed by premature infants. Many other organic and inorganic compounds that are present in foods, such as pesticides and lead, also have health effects.

ESSENTIAL NUTRIENT REQUIREMENTS

Energy For weight to remain stable, energy intake must match energy output. The major components of energy output are resting energy expenditure (REE) and physical activity; minor components include the energy cost of metabolizing food (thermic effect of food, or specific dynamic action) and shivering thermogenesis (e.g., cold-induced thermogenesis). The average energy intake is ~2600 kcal/d for American men and ~1800 kcal/d for American women, though these estimates vary with body size and activity level. Formulas for roughly estimating REE are useful in assessing the energy needs of an individual whose weight is stable. Thus, for males, $REE = 900 + 10m$, and for females, $REE = 700 + 7m$, where m is mass in kilograms. The calculated REE is then adjusted for physical activity level by multiplying by 1.2 for sedentary, 1.4 for moderately active, or 1.8 for very active individuals. The final figure, the estimated energy requirement (EER), provides an approximation of total caloric needs in a state of energy balance for a person of a certain age, sex, weight, height, and physical activity level.

For further discussion of energy balance in health and disease, see Chap. 97.

Protein Dietary protein consists of both essential and nonessential amino acids that are required for protein synthesis. The nine essential amino acids are histidine, isoleucine, leucine, lysine, methionine/cystine, phenylalanine/tyrosine, threonine, tryptophan, and valine. Certain amino acids, such as alanine, can also be used for energy and gluconeogenesis. When energy intake is inadequate, protein intake must be increased, because ingested amino acids are diverted into pathways of glucose synthesis and oxidation. In extreme energy deprivation, protein-calorie malnutrition may ensue (Chap. 97).

For adults, the recommended dietary allowance (RDA) for protein is ~0.6 g/kg desirable body mass per day, assuming that energy needs are met and that the protein is of relatively high biologic value. Current recommendations for a healthy diet call for at least 10–14% of calories from protein. Most American diets provide at least those amounts. Biologic value tends to be highest for animal proteins, followed by proteins from legumes (beans), cereals (rice, wheat, corn), and roots.

Combinations of plant proteins that complement one another in biologic value or combinations of animal and plant proteins can increase biologic value and lower total protein requirements. In healthy people with adequate diets, the timing of protein intake over the course of the day has little effect.

Protein needs increase during growth, pregnancy, lactation, and rehabilitation after injury or malnutrition. Tolerance to dietary protein is decreased in renal insufficiency (with consequent uremia) and in liver failure. Normal protein intake can precipitate encephalopathy in patients with cirrhosis of the liver.

Fat and Carbohydrate Fats are a concentrated source of energy and constitute, on average, 34% of calories in U.S. diets. However, for optimal health, fat intake should total no more than 30% of calories. Saturated fat and trans fat should be limited to <10% of calories and polyunsaturated fats to <10% of calories, with monounsaturated fats accounting for the remainder of fat intake. At least 45–55% of total calories should be derived from carbohydrates. The brain requires ~100 g of glucose per day for fuel; other tissues use about 50 g/d. Some tissues (e.g., brain and red blood cells) rely on glucose supplied either exogenously or from muscle proteolysis. Over time, adaptations in carbohydrate needs are possible during hypocaloric states. Like fat (9 kcal/g), carbohydrate (4 kcal/g), and protein (4 kcal/g), alcohol (ethanol) provides energy (7 kcal/g). However, it is not a nutrient.

Water For adults, 1–1.5 mL of water per kilocalorie of energy expenditure is sufficient under usual conditions to allow for normal variations in physical activity, sweating, and solute load of the diet. Water losses include 50–100 mL/d in the feces; 500–1000 mL/d by evaporation or exhalation; and, depending on the renal solute load, ≥1000 mL/d in the urine. If external losses increase, intakes must increase accordingly to avoid underhydration. Fever increases water losses by ~200 mL/d per °C; diarrheal losses vary but may be as great as 5 L/d in severe diarrhea. Heavy sweating, vigorous exercise, and vomiting also increase water losses. When renal function is normal and solute intakes are adequate, the kidneys can adjust to increased water intake by excreting up to 18 L of excess water per day (Chap. 404). However, obligatory urine outputs can compromise hydration status when there is inadequate water intake or when losses increase in disease or kidney damage.

Infants have high requirements for water because of their large ratio of surface area to volume, their inability to communicate their thirst, and the limited capacity of the immature kidney to handle high renal solute loads. Increased water needs during pregnancy are ~30 mL/d. During lactation, milk production increases daily water requirements so that ~1000 mL of additional water is needed, or 1 mL for each milliliter of milk produced. Special attention must be paid to the water needs of the elderly, who have reduced total body water and blunted thirst sensation and are more likely to be taking medications such as diuretics.

Other Nutrients See Chap. 96e for detailed descriptions of vitamins and trace minerals.

DIETARY REFERENCE INTAKES AND RECOMMENDED DIETARY ALLOWANCES

Fortunately, human life and well-being can be maintained within a fairly wide range for most nutrients. However, the capacity for adaptation is not infinite—too much, as well as too little, intake of a nutrient can have adverse effects or alter the health benefits conferred by another nutrient. Therefore, benchmark recommendations regarding nutrient intakes have been developed to guide clinical practice. These quantitative estimates of nutrient intakes are collectively referred to as the *dietary reference intakes* (DRIs). The DRIs have supplanted the RDAs—the single reference values used in the United States until the early 1990s. DRIs include the *estimated average requirement* (EAR) for nutrients as well as other reference values used for dietary planning for individuals: the RDA, the *adequate intake* (AI), and the *tolerable upper*

TABLE 95e-1 DIETARY REFERENCE INTAKES (DRIs): RECOMMENDED DIETARY ALLOWANCES AND ADEQUATE INTAKES FOR VITAMINS

Life-Stage Group	Vitamin A (μg/d) ^a	Vitamin C (mg/d)	Vitamin D (μg/d) ^{b,c}	Vitamin E (mg/d) ^d	Vitamin K (μg/d)	Thiamin (mg/d)	Riboflavin (mg/d)	Niacin (mg/d) ^e	Vitamin B ₆ (mg/d)	Folate (μg/d) ^f	Vitamin B ₁₂ (μg/d)	Pantothenic Acid (mg/d)	Biotin (μg/d)	Choline (mg/d) ^g
Infants														
Birth to 6 mo	400*	40*	10	4*	2.0*	0.2*	0.3*	2*	0.1*	65*	0.4*	1.7*	5*	125*
6–12 mo	500*	50*	10	5*	2.5*	0.3*	0.4*	4*	0.3*	80*	0.5*	1.8*	6*	150*
Children														
1–3 y	300	15	15	6	30*	0.5	0.5	6	0.5	150	0.9	2*	8*	200*
4–8 y	400	25	15	7	55*	0.6	0.6	8	0.6	200	1.2	3*	12*	250*
Males														
9–13 y	600	45	15	11	60*	0.9	0.9	12	1.0	300	1.8	4*	20*	375*
14–18 y	900	75	15	15	75*	1.2	1.3	16	1.3	400	2.4	5*	25*	550*
19–30 y	900	90	15	15	120*	1.2	1.3	16	1.3	400	2.4	5*	30*	550*
31–50 y	900	90	15	15	120*	1.2	1.3	16	1.3	400	2.4	5*	30*	550*
51–70 y	900	90	15	15	120*	1.2	1.3	16	1.7	400	2.4^h	5*	30*	550*
>70 y	900	90	20	15	120*	1.2	1.3	16	1.7	400	2.4^h	5*	30*	550*
Females														
9–13 y	600	45	15	11	60*	0.9	0.9	12	1.0	300	1.8	4*	20*	375*
14–18 y	700	65	15	15	75*	1.0	1.0	14	1.2	400ⁱ	2.4	5*	25*	400*
19–30 y	700	75	15	15	90*	1.1	1.1	14	1.3	400ⁱ	2.4	5*	30*	425*
31–50 y	700	75	15	15	90*	1.1	1.1	14	1.3	400ⁱ	2.4	5*	30*	425*
51–70 y	700	75	15	15	90*	1.1	1.1	14	1.5	400	2.4^h	5*	30*	425*
>70 y	700	75	20	15	90*	1.1	1.1	14	1.5	400	2.4^h	5*	30*	425*
Pregnant women														
14–18 y	750	80	15	15	75*	1.4	1.4	18	1.9	600^j	2.6	6*	30*	450*
19–30 y	770	85	15	15	90*	1.4	1.4	18	1.9	600^j	2.6	6*	30*	450*
31–50 y	770	85	15	15	90*	1.4	1.4	18	1.9	600^j	2.6	6*	30*	450*
Lactating women														
14–18 y	1200	115	15	19	75*	1.4	1.6	17	2.0	500	2.8	7*	35*	550*
19–30 y	1300	120	15	19	90*	1.4	1.6	17	2.0	500	2.8	7*	35*	550*
31–50 y	1300	120	15	19	90*	1.4	1.6	17	2.0	500	2.8	7*	35*	550*

Note: This table (taken from the DRI reports; see www.nap.edu) presents recommended dietary allowances (RDAs) in **bold type** and adequate intakes (AIs) in ordinary type followed by an asterisk (*). An RDA is the average daily dietary intake level sufficient to meet the nutrient requirements of nearly all healthy individuals (97–98%) in a group. The RDA is calculated from an estimated average requirement (EAR). If sufficient scientific evidence is not available to establish an EAR and thus to calculate an RDA, an AI is usually developed. For healthy breast-fed infants, an AI is the mean intake. The AI for other life-stage and sex-specific groups is believed to cover the needs of all healthy individuals in those groups, but lack of data or uncertainty in the data makes it impossible to specify with confidence the percentage of individuals covered by this intake.

^aAs retinol activity equivalents (RAEs). 1 RAE = 1 μg retinol, 12 μg β-carotene, 24 μg α-carotene, or 24 μg β-cryptoxanthin. The RAE for dietary provitamin A carotenoids is twofold greater than the retinol equivalent (RE), whereas the RAE for preformed vitamin A is the same as the RE. ^bAs cholecalciferol. 1 μg cholecalciferol = 40 IU vitamin D. ^cUnder the assumption of minimal sunlight. ^dAs α-tocopherol. α-Tocopherol includes *RRR*-α-tocopherol, the only form of α-tocopherol that occurs naturally in foods, and the *2R*-stereoisomeric forms of α-tocopherol (*RRR*-, *RSR*-, *RRS*-, and *RSS*-α-tocopherol) that occur in fortified foods and supplements. It does not include the *2S*-stereoisomeric forms of α-tocopherol (*SRR*-, *SSR*-, *SRS*-, and *SSS*-α-tocopherol) also found in fortified foods and supplements. ^eAs niacin equivalents (NEs). 1 mg of niacin = 60 mg of tryptophan; 0–6 months = preformed niacin (not NE). ^fAs dietary folate equivalents (DFEs). 1 DFE = 1 μg food folate = 0.6 μg of folic acid from fortified food or as a supplement consumed with food = 0.5 μg of a supplement taken on an empty stomach. ^gAlthough AIs have been set for choline, there are few data to assess whether a dietary supply of choline is needed at all stages of the life cycle, and it may be that the choline requirement can be met by endogenous synthesis at some of these stages. ^hBecause 10–30% of older people may malabsorb food-bound B₁₂, it is advisable for those >50 years of age to meet their RDA mainly by consuming foods fortified with B₁₂ or a supplement containing B₁₂. ⁱIn view of evidence linking inadequate folate intake with neural tube defects in the fetus, it is recommended that all women capable of becoming pregnant consume 400 μg of folate from supplements or fortified foods in addition to intake of food folate from a varied diet. ^jIt is assumed that women will continue consuming 400 μg from supplements or fortified food until their pregnancy is confirmed and they enter prenatal care, which ordinarily occurs after the end of the periconceptional period—the critical time for formation of the neural tube.

Source: Food and Nutrition Board, Institute of Medicine, National Academies (<http://www.iom.edu/Activities/Nutrition/SummaryDRIs/DRI-Tables.aspx>).

level (UL). The DRIs also include acceptable macronutrient distribution ranges (AMDRs) for protein, fat, and carbohydrate. The current DRIs for vitamins and elements are provided in **Tables 95e-1 and 95e-2**, respectively. **Table 95e-3** provides DRIs for water and macronutrients. **EERs are discussed in Chap. 97 on energy balance in health and disease.**

Estimated Average Requirement When florid manifestations of the classic dietary-deficiency diseases such as rickets (deficiency of vitamin D and calcium), scurvy (deficiency of vitamin C), xerophthalmia (deficiency of vitamin A), and protein-calorie malnutrition were common, nutrient adequacy was inferred from the absence of their clinical signs. Later, biochemical and other changes were found to be evident long before the deficiency became clinically apparent. Consequently, criteria of adequacy are now based on biologic markers when they are available. Priority is given to sensitive biochemical, physiologic, or behavioral tests that reflect early changes in regulatory processes;

maintenance of body stores of nutrients; or, if available, the amount of a nutrient that minimizes the risk of chronic degenerative disease. Current efforts focus on this last variable, but relevant markers often are not available.

The EAR is the amount of a nutrient estimated to be adequate for half of the healthy individuals of a specific age and sex. The types of evidence and criteria used to establish nutrient requirements vary by nutrient, age, and physiologic group. The EAR is not an effective estimate of nutrient adequacy in individuals because it is a median requirement for a group; 50% of individuals in a group fall below the requirement and 50% fall above it. Thus, a person with a usual intake at the EAR has a 50% risk of inadequate intake. For these reasons, other standards, described below, are more useful for clinical purposes.

Recommended Dietary Allowances The RDA is the average daily dietary intake level that meets the nutrient requirements of nearly all healthy persons of a specific sex, age, life stage, or physiologic condition

TABLE 95e-2 DIETARY REFERENCE INTAKES (DRIs): RECOMMENDED DIETARY ALLOWANCES AND ADEQUATE INTAKES FOR ELEMENTS

Life-Stage Group	Calcium (mg/d)	Chromium (µg/d)	Copper (µg/d)	Fluoride (mg/d)	Iodine (µg/d)	Iron (mg/d)	Magnesium (mg/d)	Manganese (mg/d)	Molybdenum (µg/d)	Phosphorus (mg/d)	Selenium (µg/d)	Zinc (mg/d)	Potassium (g/d)	Sodium (g/d)	Chloride (g/d)
Infants															
Birth to 6 mo	200*	0.2*	200*	0.01*	110*	0.27*	30*	0.003*	2*	100*	15*	2*	0.4*	0.12*	0.18*
6–12 mo	260*	5.5*	220*	0.5*	130*	11	75*	0.6*	3*	275*	20*	3	0.7*	0.37*	0.57*
Children															
1–3 y	700	11*	340	0.7*	90	7	80	1.2*	17	460	20	3	3.0*	1.0*	1.5*
4–8 y	1000	15*	440	1*	90	10	130	1.5*	22	500	30	5	3.8*	1.2*	1.9*
Males															
9–13 y	1300	25*	700	2*	120	8	240	1.9*	34	1250	40	8	4.5*	1.5*	2.3*
14–18 y	1300	35*	890	3*	150	11	410	2.2*	43	1250	55	11	4.7*	1.5*	2.3*
19–30 y	1000	35*	900	4*	150	8	400	2.3*	45	700	55	11	4.7*	1.5*	2.3*
31–50 y	1000	35*	900	4*	150	8	420	2.3*	45	700	55	11	4.7*	1.5*	2.3*
51–70 y	1000	30*	900	4*	150	8	420	2.3*	45	700	55	11	4.7*	1.3*	2.0*
>70 y	1200	30*	900	4*	150	8	420	2.3*	45	700	55	11	4.7*	1.2*	1.8*
Females															
9–13 y	1300	21*	700	2*	120	8	240	1.6*	34	1250	40	8	4.5*	1.5*	2.3*
14–18 y	1300	24*	890	3*	150	15	360	1.6*	43	1250	55	9	4.7*	1.5*	2.3*
19–30 y	1000	25*	900	3*	150	18	310	1.8*	45	700	55	8	4.7*	1.5*	2.3*
31–50 y	1000	25*	900	3*	150	18	320	1.8*	45	700	55	8	4.7*	1.5*	2.3*
51–70 y	1200	20*	900	3*	150	8	320	1.8*	45	700	55	8	4.7*	1.3*	2.0*
>70 y	1200	20*	900	3*	150	8	320	1.8*	45	700	55	8	4.7*	1.2*	1.8*
Pregnant women															
14–18 y	1300	29*	1000	3*	220	27	400	2.0*	50	1250	60	12	4.7*	1.5*	2.3*
19–30 y	1000	30*	1000	3*	220	27	350	2.0*	50	700	60	11	4.7*	1.5*	2.3*
31–50 y	1000	30*	1000	3*	220	27	360	2.0*	50	700	60	11	4.7*	1.5*	2.3*
Lactating women															
14–18 y	1300	44*	1300	3*	290	10	360	2.6*	50	1250	70	13	5.1*	1.5*	2.3*
19–30 y	1000	45*	1300	3*	290	9	310	2.6*	50	700	70	12	5.1*	1.5*	2.3*
31–50 y	1000	45*	1300	3*	290	9	320	2.6*	50	700	70	12	5.1*	1.5*	2.3*

Note: This table (taken from the DRI reports; see www.nap.edu) presents recommended dietary allowances (RDAs) in **bold type** and adequate intakes (AIs) in ordinary type followed by an asterisk (*). An RDA is the average daily dietary intake level sufficient to meet the nutrient requirements of nearly all healthy individuals (97–98%) in a group. The RDA is calculated from an estimated average requirement (EAR). If sufficient scientific evidence is not available to establish an EAR and thus to calculate an RDA, an AI is usually developed. For healthy breast-fed infants, an AI is the mean intake. The AI for other life-stage and sex-specific groups is believed to cover the needs of all healthy individuals in those groups, but lack of data or uncertainty in the data makes it impossible to specify with confidence the percentage of individuals covered by this intake.

Sources: Food and Nutrition Board, Institute of Medicine, National Academies (<http://www.iom.edu/Activities/Nutrition/SummaryDRIs/DRI-Tables.aspx>), based on: *Dietary Reference Intakes for Calcium, Phosphorus, Magnesium, Vitamin D, and Fluoride* (1997); *Dietary Reference Intakes for Thiamin, Riboflavin, Niacin, Vitamin B₆, Folate, Vitamin B₁₂, Pantothenic Acid, Biotin, and Choline* (1998); *Dietary Reference Intakes for Vitamin C, Vitamin E, Selenium, and Carotenoids* (2000); and *Dietary Reference Intakes for Vitamin A, Vitamin K, Arsenic, Boron, Chromium, Copper, Iodine, Iron, Manganese, Molybdenum, Nickel, Silicon, Vanadium, and Zinc* (2001); *Dietary Reference Intakes for Water, Potassium, Sodium, Chloride, and Sulfate* (2005); and *Dietary Reference Intakes for Calcium and Vitamin D* (2011). These reports can be accessed via www.nap.edu.

(e.g., pregnancy or lactation). The RDA, which is the nutrient-intake goal for planning diets of individuals, is defined statistically as two standard deviations above the EAR to ensure that the needs of any given individual are met. The online tool at <http://fnic.nal.usda.gov/interactiveDRI/> allows health professionals to calculate individualized daily nutrient recommendations for dietary planning based on the DRIs for persons of a given age, sex, and weight. The RDAs are used to formulate food guides such as the U.S. Department of Agriculture (USDA) MyPlate Food Guide for individuals (www.supertracker.usda.gov/default.aspx), to create food-exchange lists for therapeutic diet planning, and as a standard for describing the nutritional content of foods and nutrient-containing dietary supplements.

The risk of dietary inadequacy increases as intake falls below the RDA. However, the RDA is an overly generous criterion for evaluating nutrient adequacy. For example, by definition, the RDA exceeds the actual requirements of all but ~2–3% of the population. Therefore, many people whose intake falls below the RDA may still be getting enough of the nutrient. On food labels, the nutrient content in a food is stated by weight or as a percent of the daily value (DV), a variant of the RDA used on the nutrition facts panel that, for an adult, represents the highest RDA for an adult consuming 2000 kcal.

Adequate Intake It is not possible to set an RDA for some nutrients that do not have an established EAR. In this circumstance, the AI is

based on observed or experimentally determined approximations of nutrient intakes in healthy people. In the DRIs, AIs rather than RDAs are proposed for nutrients consumed by infants (up to age 1 year) as well as for chromium, fluoride, manganese, sodium, potassium, pantothenic acid, biotin, choline, and water consumed by persons of all ages. Vitamin D and calcium recommendations were recently revised, and more precise estimates are now available.

Tolerable Upper Levels of Nutrient Intake Healthy individuals derive no established benefit from consuming nutrient levels above the RDA or AI. In fact, excessive nutrient intake can disturb body functions and cause acute, progressive, or permanent disabilities. The tolerable UL is the highest level of chronic nutrient intake (usually daily) that is unlikely to pose a risk of adverse health effects for most of the population. Data on the adverse effects of large amounts of many nutrients are unavailable or too limited to establish a UL. Therefore, the lack of a UL does *not* mean that the risk of adverse effects from high intake is nonexistent. Nutrients in commonly eaten foods rarely exceed the UL. However, highly fortified foods and dietary supplements provide more concentrated amounts of nutrients per serving and thus pose a potential risk of toxicity. Nutrient supplements are labeled with supplement facts that express the amount of nutrient in absolute units or as the percentage of the DV provided per recommended serving size. Total nutrient consumption, including that in foods, supplements,

TABLE 95e-3 DIETARY REFERENCE INTAKES (DRIs): RECOMMENDED DIETARY ALLOWANCES AND ADEQUATE INTAKES FOR TOTAL WATER AND MACRONUTRIENTS

Life-Stage Group	Total Water ^a (L/d)	Carbohydrate (g/d)	Total Fiber (g/d)	Fat (g/d)	Linoleic Acid (g/d)	α -Linolenic Acid (g/d)	Protein ^b (g/d)
Infants							
Birth to 6 mo	0.7*	60*	ND ^c	31*	4.4*	0.5*	9.1*
6–12 mo	0.8*	95*	ND	30*	4.6*	0.5*	11.0
Children							
1–3 y	1.3*	130	19*	ND	7*	0.7*	13
4–8 y	1.7*	130	25*	ND	10*	0.9*	19
Males							
9–13 y	2.4*	130	31*	ND	12*	1.2*	34
14–18 y	3.3*	130	38*	ND	16*	1.6*	52
19–30 y	3.7*	130	38*	ND	17*	1.6*	56
31–50 y	3.7*	130	38*	ND	17*	1.6*	56
51–70 y	3.7*	130	30*	ND	14*	1.6*	56
>70 y	3.7*	130	30*	ND	14*	1.6*	56
Females							
9–13 y	2.1*	130	26*	ND	10*	1.0*	34
14–18 y	2.3*	130	26*	ND	11*	1.1*	46
19–30 y	2.7*	130	25*	ND	12*	1.1*	46
31–50 y	2.7*	130	25*	ND	12*	1.1*	46
51–70 y	2.7*	130	21*	ND	11*	1.1*	46
>70 y	2.7*	130	21*	ND	11*	1.1*	46
Pregnant women							
14–18 y	3.0*	175	28*	ND	13*	1.4*	71
19–30 y	3.0*	175	28*	ND	13*	1.4*	71
31–50 y	3.0*	175	28*	ND	13*	1.4*	71
Lactating women							
14–18	3.8*	210	29*	ND	13*	1.3*	71
19–30 y	3.8*	210	29*	ND	13*	1.3*	71
31–50 y	3.8*	210	29*	ND	13*	1.3*	71

Note: This table (taken from the DRI reports; see www.nap.edu) presents recommended dietary allowances (RDAs) in **bold type** and adequate intakes (AIs) in ordinary type followed by an asterisk (*). An RDA is the average daily dietary intake level sufficient to meet the nutrient requirements of nearly all healthy individuals (97–98%) in a group. The RDA is calculated from an estimated average requirement (EAR). If sufficient scientific evidence is not available to establish an EAR and thus to calculate an RDA, an AI is usually developed. For healthy breast-fed infants, an AI is the mean intake. The AI for other life-stage and sex-specific groups is believed to cover the needs of all healthy individuals in those groups, but lack of data or uncertainty in the data make it impossible to specify with confidence the percentage of individuals covered by this intake.

^aTotal water includes all water contained in food, beverages, and drinking water. ^bBased on grams of protein per kilogram of body weight for the reference body weight (e.g., for adults: 0.8 g/kg body weight for the reference body weight). ^cNot determined.

Source: Food and Nutrition Board, Institute of Medicine, National Academies (<http://www.iom.edu/Activities/Nutrition/SummaryDRIs/DRI-Tables.aspx>), based on: *Dietary Reference Intakes for Energy, Carbohydrate, Fiber, Fat, Fatty Acids, Cholesterol, Protein, and Amino Acids* (2002/2005) and *Dietary Reference Intakes for Water, Potassium, Sodium, Chloride, and Sulfate* (2005). These reports can be accessed via www.nap.edu.

and over-the-counter medications (e.g., antacids), should not exceed RDA levels.

Acceptable Macronutrient Distribution Ranges The AMDRs are not experimentally determined but are rough ranges for energy-providing macronutrient intakes (protein, carbohydrate, and fat) that the Institute of Medicine's Food and Nutrition Board considers to be healthful. These ranges are 10–35% of calories for protein, 20–35% of calories for fat, and 45–65% of calories for carbohydrate. Alcohol, which also provides energy, is not a nutrient; therefore, no recommendations are not provided.

FACTORS ALTERING NUTRIENT NEEDS

The DRIs are affected by age, sex, rate of growth, pregnancy, lactation, physical activity level, concomitant diseases, drugs, and dietary composition. If requirements for nutrient sufficiency are close to levels indicating excess of a nutrient, dietary planning is difficult.

Physiologic Factors Growth, strenuous physical activity, pregnancy, and lactation all increase needs for energy and several essential nutrients. Energy needs rise during pregnancy due to the demands of fetal growth and during lactation because of the increased energy required for milk production. Energy needs decrease with loss of lean body mass, the major determinant of REE. Because lean tissue, physical activity,

and health often decline with age, energy needs of older persons, especially those over 70, tend to be lower than those of younger persons.

Dietary Composition Dietary composition affects the biologic availability and use of nutrients. For example, the absorption of iron may be impaired by large amounts of calcium or lead; likewise, non-heme iron uptake may be impaired by a lack of ascorbic acid and amino acids in the meal. Protein use by the body may be decreased when essential amino acids are not present in sufficient amounts—a rare scenario in U.S. diets. Animal foods, such as milk, eggs, and meat, have high biologic values, with most of the needed amino acids present in adequate amounts. Plant proteins in corn (maize), soy, rice, and wheat have lower biologic values and must be combined with other plant or animal proteins or fortified with the amino acids that are deficient to achieve optimal use by the body.

Route of Intake The RDAs apply only to oral intakes. When nutrients are administered parenterally, similar values can sometimes be used for amino acids, glucose (carbohydrate), fats, sodium, chloride, potassium, and most vitamins because their intestinal absorption rate is nearly 100%. However, the oral bioavailability of most mineral elements may be only half that obtained by parenteral administration. For some nutrients that are not readily stored in the body or that cannot be stored in large amounts, timing of administration may also

be important. For example, amino acids cannot be used for protein synthesis if they are not supplied together; instead, they will be used for energy production, although in healthy individuals eating adequate diets, the distribution of protein intake over the course of the day has little effect on health.

Disease Dietary deficiency diseases include protein-calorie malnutrition, iron-deficiency anemia, goiter (due to iodine deficiency), rickets and osteomalacia (vitamin D deficiency), and xerophthalmia (vitamin A deficiency), megaloblastic anemia (vitamin B₁₂ or folic acid deficiency), scurvy (vitamin C/ascorbic acid deficiency), beriberi (thiamin deficiency), and pellagra (niacin and tryptophan deficiency) (**Chaps. 96e and 97**). Each deficiency disease is characterized by imbalances at the cellular level between the supply of nutrients or energy and the body's nutritional needs for growth, maintenance, and other functions. Imbalances and excesses in nutrient intakes are recognized as risk factors for certain chronic degenerative diseases, such as saturated fat and cholesterol in coronary artery disease; sodium in hypertension; obesity in hormone-dependent endometrial and breast cancers; and ethanol in alcoholism. Because the etiology and pathogenesis of these disorders are multifactorial, diet is only one of many risk factors. Osteoporosis, for example, is associated with calcium deficiency, sometimes secondary to vitamin D deficiency, as well as with risk factors related to environment (e.g., smoking, sedentary lifestyle), physiology (e.g., estrogen deficiency), genetic determinants (e.g., defects in collagen metabolism), and drug use (chronic steroid and aromatase inhibitors) (**Chap. 425**).

DIETARY ASSESSMENT

In clinical situations, nutritional assessment is an iterative process that involves: (1) screening for malnutrition, (2) assessing the diet and other data to establish either the absence or the presence of malnutrition and its possible causes, (3) planning and implementing the most appropriate nutritional therapy, and (4) reassessing intakes to make sure that they have been consumed. Some disease states affect the bioavailability, requirements, use, or excretion of specific nutrients. In these circumstances, specific measurements of various nutrients or their biomarkers may be required to ensure adequate replacement (**Chap. 96e**).

Most health care facilities have nutrition-screening processes in place for identifying possible malnutrition after hospital admission. Nutritional screening is required by the Joint Commission, which accredits and certifies health care organizations in the United States. However, there are no universally recognized or validated standards. The factors that are usually assessed include abnormal weight for height or body mass index (e.g., BMI <19 or >25); reported weight change (involuntary loss or gain of >5 kg in the past 6 months) (**Chap. 56**); diagnoses with known nutritional implications (e.g., metabolic disease, any disease affecting the gastrointestinal tract, alcoholism); present therapeutic dietary prescription; chronic poor appetite; presence of chewing and swallowing problems or major food intolerances; need for assistance with preparing or shopping for food, eating, or other aspects of self-care; and social isolation. The nutritional status of hospitalized patients should be reassessed periodically—at least once every week.

A more complete dietary assessment is indicated for patients who exhibit a high risk of or frank malnutrition on nutritional screening. The type of assessment varies with the clinical setting, the severity of the patient's illness, and the stability of the patient's condition.

Acute-Care Settings In acute-care settings, anorexia, various other diseases, test procedures, and medications can compromise dietary intake. Under such circumstances, the goal is to identify and avoid inadequate intake and to assure appropriate alimentation. Dietary assessment focuses on what patients are currently eating, whether or not they are able and willing to eat, and whether or not they experience any problems with eating. Dietary intake assessment is based on information from observed intakes; medical records; history; clinical examination; and anthropometric, biochemical, and functional status

evaluations. The objective is to gather enough information to establish the likelihood of malnutrition due to poor dietary intake or other causes in order to assess whether nutritional therapy is indicated (**Chap. 98e**).

Simple observations may suffice to suggest inadequate oral intake. These include dietitians' and nurses' notes; observation of a patient's frequent refusal to eat or the amount of food eaten on trays; the frequent performance of tests and procedures that are likely to cause meals to be skipped; adherence to nutritionally inadequate diet orders (e.g., clear liquids or full liquids) for more than a few days; the occurrence of fever, gastrointestinal distress, vomiting, diarrhea, or a comatose state; and the presence of diseases or use of treatments that involve any part of the alimentary tract. Acutely ill patients with diet-related diseases such as diabetes need assessment because an inappropriate diet may exacerbate these conditions and adversely affect other therapies. Abnormal biochemical values (serum albumin levels <35 g/L [<3.5 mg/dL]; serum cholesterol levels <3.9 mmol/L [<150 mg/dL]) are nonspecific but may indicate a need for further nutritional assessment.

Most therapeutic diets offered in hospitals are calculated to meet individual nutrient requirements and the RDA *if they are eaten*. Exceptions include clear liquids, some full-liquid diets, and test diets (such as those adhered to in preparation for gastrointestinal procedures), which are inadequate for several nutrients and should not be used, if possible, for more than 24 h. However, because as much as half of the food served to hospitalized patients is not eaten, it cannot be assumed that the intakes of hospitalized patients are adequate. Dietary assessment should compare how much and what kinds of food the patient has consumed with the diet that has been provided. Major deviations in intakes of energy, protein, fluids, or other nutrients of special concern for the patient's illness should be noted and corrected.

Nutritional monitoring is especially important for patients who are very ill and who have extended lengths of hospital stay. Patients who are fed by enteral and parenteral routes also require special nutritional assessment and monitoring by physicians and/or dietitians with certification in nutritional support (**Chap. 98e**).

Ambulatory Settings The aim of dietary assessment in the outpatient setting is to determine whether or not the patient's usual diet is a health risk in itself or if it contributes to existing chronic disease-related problems. Dietary assessment also provides the basis for planning a diet that fulfills therapeutic goals while ensuring patient adherence. The outpatient dietary assessment should review the adequacy of present and usual food intakes, including vitamin and mineral supplements, oral nutritional supplements, medical foods, other dietary supplements, medications, and alcohol, because all of these may affect the patient's nutritional status. The assessment should focus on the dietary constituents that are most likely to be involved or compromised by a specific diagnosis as well as on any comorbidities that are present. More than one day's intake should be reviewed to provide a better representation of the usual diet.

There are many ways to assess the adequacy of a patient's habitual diet. These include use of a food guide, a food-exchange list, a diet history, or a food-frequency questionnaire. A commonly used food guide for healthy persons is the USDA's Choose My Plate, which is useful as a rough guide for avoiding inadequate intakes of essential nutrients as well as likely excesses in the amounts of fat (especially saturated and trans fats), sodium, sugar, and alcohol consumed (**Table 95e-4**). The Choose My Plate graphic emphasizes a balance between calories and nutritional needs, encouraging increased intake of fruits and vegetables, whole grains, and low-fat milk in conjunction with reduced intake of sodium and high-calorie sugary drinks. The Web version of the guide provides a calculator that tailors the number of servings suggested for healthy patients of different weights, sexes, ages, and life-cycle stages to help them to meet their needs while avoiding excess (<http://www.supertracker.usda.gov/default.aspx> and www.ChooseMyPlate.gov). Patients who follow ethnic or unusual dietary patterns may need extra instruction on how foods should be categorized and on the appropriate portion sizes that constitute a serving. The process of reviewing the guide with patients helps them transition

TABLE 95e-4 CHOOSE MY PLATE: A GUIDE TO INDIVIDUALIZED DIETARY PLANNING

Dietary Factor, Unit of Measure (Advice)	Examples of Standard Portion Sizes at Indicated Energy Level		
	Lower: 1600 kcal	Moderate: 2200 kcal	Higher: 2800 kcal
Fruits, cups (Focus on fruits.)	1.5	2	2.5
Vegetables, cups (Vary vegetables.)	2	3	3.5
Grains, oz eq (Make at least half of grains whole.) ^a	5	7	10
Protein foods, oz eq (Go lean with protein.) ^b	5	6	7
Dairy, cups or oz ^c (Choose calcium-rich foods.)	3	3	3
"Empty" calories, kcal ^d	120	260	400
Sodium, mg	<2300 at all energy levels		
Physical activity, min	At least 150 min vigorous physical activity per week at all energy levels		

Note: Oils (formerly listed with portions of 5, 6, and 8 teaspoons for the lower, moderate, and higher energy levels, respectively) are no longer singled out in Choose My Plate, but rather are included in the empty calories/added sugar category with SOFAS (calories from solid fats and added sugars). The limit is the remaining number of calories in each food pattern above after intake of the recommended amounts of the nutrient-dense foods.

^aFor example, 1 serving equals 1 slice bread, 1 cup ready-to-eat cereal, or 0.5 cup cooked rice, pasta, or cooked cereal. ^bFor example, 1 serving equals 1 oz lean meat, poultry, or fish; 1 egg; 1 tablespoon peanut butter; 0.25 cup cooked dry beans; or 0.5 oz nuts or seeds. ^cFor example, 1 serving equals 1 cup milk or yogurt, 1.5 oz natural cheese, or 2 oz processed cheese. ^dFormerly called "discretionary calorie allowance." Portions are calculated as the number of calories remaining after all of the above allotments are accounted for.

Abbreviation: oz eq, ounce equivalent.

Source: Data from U.S. Department of Agriculture (<http://www.Choosemyplate.gov>).

to healthier dietary patterns and identifies food groups eaten in excess of recommendations or in insufficient quantities. For persons on therapeutic diets, assessment against food-exchange lists may be useful. These include, for example, American Diabetes Association food-

exchange lists for diabetes and the Academy of Nutrition and Dietetics food-exchange lists for renal disease.

NUTRITIONAL STATUS ASSESSMENT

Full nutritional status assessment is reserved for seriously ill patients and those at very high nutritional risk when the cause of malnutrition is still uncertain after the initial clinical evaluation and dietary assessment. It involves multiple dimensions, including documentation of dietary intake, anthropometric measurements, biochemical measurements of blood and urine, clinical examination, health history elicitation, and functional status evaluation. Therapeutic dietary prescriptions and menu plans for most diseases are available from most hospitals and from the Academy of Nutrition and Dietetics. **For further discussion of nutritional assessment, see Chap. 97.**

GLOBAL CONSIDERATIONS



The DRIs (e.g., the EAR, the UL, and energy needs) are estimates of physiologic requirements based on experimental evidence. Assuming that appropriate adjustments are made for age, sex, body size, and physical activity level, these estimates should be applicable to individuals in most parts of the world. However, the AIs are based on customary and adequate intakes in U.S. and Canadian populations, which appear to be compatible with good health, rather than on a large body of direct experimental evidence. Similarly, the AMDRs represent expert opinion regarding the approximate intakes of energy-providing nutrients that are healthful in these North American populations. Thus these measures should be used with caution in other settings. Nutrient-based standards like the DRIs have also been developed by the World Health Organization/Food and Agricultural Organization of the United Nations and are available on the Web (<http://www.who.int/nutrition/topics/nutrecomm/en/index.html>). The European Food Safety Authority (EFSA) Panel on Dietetic Products, Nutrition and Allergies periodically publishes its recommendations in the *EFSA Journal*. Other countries have promulgated similar recommendations. The different standards have many similarities in their basic concepts, definitions, and nutrient recommendation levels, but there are some differences from the DRIs as a result of the functional criteria chosen, environmental differences, the timeliness of the evidence reviewed, and expert judgment.

Vitamins are required constituents of the human diet since they are synthesized inadequately or not at all in the human body. Only small amounts of these substances are needed to carry out essential biochemical reactions (e.g., by acting as coenzymes or prosthetic groups). Overt vitamin or trace mineral deficiencies are rare in Western countries because of a plentiful, varied, and inexpensive food supply; food fortification; and use of supplements. However, multiple nutrient deficiencies may appear together in persons who are chronically ill or alcoholic. After gastric bypass surgery, patients are at high risk for multiple nutrient deficiencies. Moreover, subclinical vitamin and trace mineral deficiencies, as diagnosed by laboratory testing, are quite common in the normal population, especially in the geriatric age group. Conversely, because of the widespread use of nutrient supplements, nutrient toxicities are gaining pathophysiologic and clinical importance.



Victims of famine, emergency-affected and displaced populations, and refugees are at increased risk for protein-energy malnutrition and classic micronutrient deficiencies (vitamin A, iron, iodine) as well as for overt deficiencies in thiamine (beriberi), riboflavin, vitamin C (scurvy), and niacin (pellagra).

Body stores of vitamins and minerals vary tremendously. For example, stores of vitamin B₁₂ and vitamin A are large, and an adult may not become deficient until ≥ 1 year after beginning to eat a deficient diet. However, folate and thiamine may become depleted within weeks among those eating a deficient diet. Therapeutic modalities can deplete essential nutrients from the body; for example, hemodialysis removes water-soluble vitamins, which must be replaced by supplementation.

Vitamins and trace minerals play several roles in diseases: (1) Deficiencies of vitamins and minerals may be caused by disease states such as malabsorption. (2) Either deficiency or excess of vitamins and minerals can cause disease in and of itself (e.g., vitamin A intoxication and liver disease). (3) Vitamins and minerals in high doses may be

used as drugs (e.g., niacin for hypercholesterolemia). Since they are covered elsewhere, the hematologic-related vitamins and minerals (Chaps. 126 and 128) either are not considered or are considered only briefly in this chapter, as are the bone-related vitamins and minerals (vitamin D, calcium, phosphorus, magnesium; Chap. 423).

VITAMINS

See also Table 96e-1 and Fig. 96e-1.

THIAMINE (VITAMIN B₁)

Thiamine was the first B vitamin to be identified and therefore is referred to as vitamin B₁. Thiamine functions in the decarboxylation of α -ketoacids (e.g., pyruvate α -ketoglutarate) and branched-chain amino acids and thus is essential for energy generation. In addition, thiamine pyrophosphate acts as a coenzyme for a transketolase reaction that mediates the conversion of hexose and pentose phosphates. It has been postulated that thiamine plays a role in peripheral nerve conduction, although the exact chemical reactions underlying this function are not known.

Food Sources The median intake of thiamine in the United States from food alone is 2 mg/d. Primary food sources for thiamine include yeast, organ meat, pork, legumes, beef, whole grains, and nuts. Milled rice and grains contain little thiamine. Thiamine deficiency is therefore more common in cultures that rely heavily on a rice-based diet. Tea, coffee (regular and decaffeinated), raw fish, and shellfish contain thiaminases, which can destroy the vitamin. Thus, drinking large amounts of tea or coffee can theoretically lower thiamine body stores.

Deficiency Most dietary deficiency of thiamine worldwide is the result of poor dietary intake. In Western countries, the primary causes of thiamine deficiency are alcoholism and chronic illnesses such as cancer. Alcohol interferes directly with the absorption of thiamine and with the synthesis of thiamine pyrophosphate, and it increases urinary excretion. Thiamine should always be replenished when a patient with alcoholism is being refeed, as carbohydrate repletion without adequate thiamine can precipitate acute thiamine deficiency with lactic acidosis. Other at-risk populations are women with prolonged hyperemesis gravidarum and anorexia, patients with overall poor nutritional status who are receiving parenteral glucose, patients who have had

TABLE 96e-1 PRINCIPAL CLINICAL FINDINGS OF VITAMIN MALNUTRITION

Nutrient	Clinical Finding	Dietary Level per Day Associated with Overt Deficiency in Adults	Contributing Factors to Deficiency
Thiamine	Beriberi: neuropathy, muscle weakness and wasting, cardiomegaly, edema, ophthalmoplegia, confabulation	<0.3 mg/1000 kcal	Alcoholism, chronic diuretic use, hyperemesis, thiaminases in food
Riboflavin	Magenta tongue, angular stomatitis, seborrhea, cheilosis	<0.6 mg	Alcoholism
Niacin	Pellagra: pigmented rash of sun-exposed areas, bright red tongue, diarrhea, apathy, memory loss, disorientation	<9.0 niacin equivalents	Alcoholism, vitamin B ₆ deficiency, riboflavin deficiency, tryptophan deficiency
Vitamin B ₆	Seborrhea, glossitis, convulsions, neuropathy, depression, confusion, microcytic anemia	<0.2 mg	Alcoholism, isoniazid
Folate	Megaloblastic anemia, atrophic glossitis, depression, \uparrow homocysteine	<100 μ g/d	Alcoholism, sulfasalazine, pyrimethamine, triamterene
Vitamin B ₁₂	Megaloblastic anemia, loss of vibratory and position sense, abnormal gait, dementia, impotence, loss of bladder and bowel control, \uparrow homocysteine, \uparrow methylmalonic acid	<1.0 μ g/d	Gastric atrophy (pernicious anemia), terminal ileal disease, strict vegetarianism, acid-reducing drugs (e.g., H ₂ blockers), metformin
Vitamin C	Scurvy: petechiae, ecchymosis, coiled hairs, inflamed and bleeding gums, joint effusion, poor wound healing, fatigue	<10 mg/d	Smoking, alcoholism
Vitamin A	Xerophthalmia, night blindness, Bitot's spots, follicular hyperkeratosis, impaired embryonic development, immune dysfunction	<300 μ g/d	Fat malabsorption, infection, measles, alcoholism, protein-energy malnutrition
Vitamin D	Rickets: skeletal deformation, rachitic rosary, bowed legs; osteomalacia	<2.0 μ g/d	Aging, lack of sunlight exposure, fat malabsorption, deeply pigmented skin
Vitamin E	Peripheral neuropathy, spinocerebellar ataxia, skeletal muscle atrophy, retinopathy	Not described unless underlying contributing factor is present	Occurs only with fat malabsorption or genetic abnormalities of vitamin E metabolism/transport
Vitamin K	Elevated prothrombin time, bleeding	<10 μ g/d	Fat malabsorption, liver disease, antibiotic use

bariatric bypass surgery (*bariatric Wernicke*), and patients receiving chronic diuretic therapy (e.g., in hypertension or heart failure) due to increased urinary thiamine losses. Maternal thiamine deficiency can lead to infantile beriberi in breast-fed children. Thiamine deficiency could be an underlying factor in motor vehicle accidents and could be overlooked in the setting of head injury.

Thiamine deficiency in its early stage induces anorexia and non-specific symptoms (e.g., irritability, decrease in short-term memory).

Prolonged thiamine deficiency causes *beriberi*, which is classically categorized as wet or dry although there is considerable overlap between the two categories. In either form of beriberi, patients may complain of pain and paresthesia. *Wet beriberi* presents primarily with cardiovascular symptoms that are due to impaired myocardial energy metabolism and dysautonomia; it can occur after 3 months of a thiamine-deficient diet. Patients present with an enlarged heart, tachycardia, high-output congestive heart failure, peripheral edema,

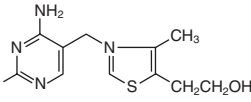
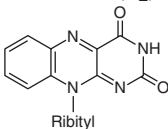
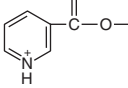
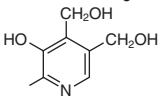
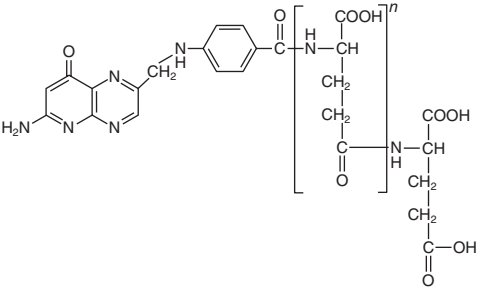
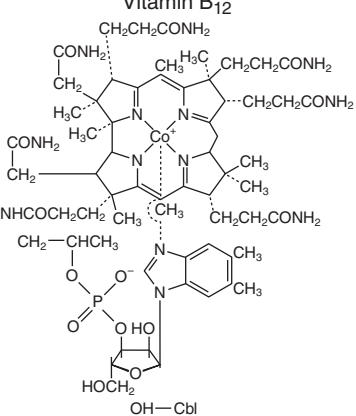
Vitamin	Active derivative or cofactor form	Principal function
<p>Thiamine (B₁)</p> 	Thiamine pyrophosphate	Coenzyme for cleavage of carbon-carbon bonds; amino acid and carbohydrate metabolism
<p>Riboflavin (B₂)</p> 	Flavin mononucleotide (FMN) and flavin adenine dinucleotide (FAD)	Cofactor for oxidation, reduction reactions, and covalently attached prosthetic groups for some enzymes
<p>Niacin</p> 	Nicotinamide adenine dinucleotide phosphate (NADP) and nicotinamide adenine dinucleotide (NAD)	Coenzymes for oxidation and reduction reactions
<p>Vitamin B₆</p> 	Pyridoxal phosphate	Cofactor for enzymes of amino acid metabolism
<p>Folate</p> 	Polyglutamate forms of (5, 6, 7, 8) tetrahydrofolate with carbon unit attachments	Coenzyme for one carbon transfer in nucleic acid and amino acid metabolism
<p>Vitamin B₁₂</p> 	Methylcobalamine Adenosylcobalamine	Coenzyme for methionine synthase and L-methylmalonyl-CoA mutase

FIGURE 96e-1 Structures and principal functions of vitamins associated with human disorders.

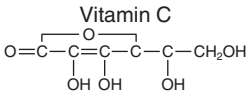
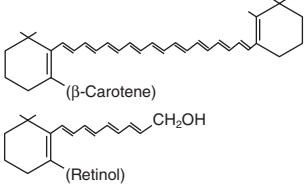
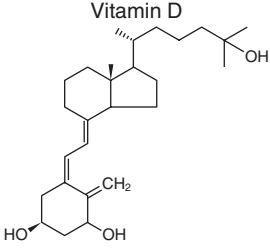
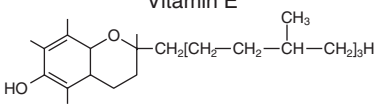
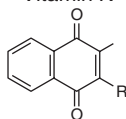
Vitamin	Active derivative or cofactor form	Principal function
<p>Vitamin C</p> 	Ascorbic acid and dehydroascorbic acid	Participation as a redox ion in many biologic oxidation and hydrogen transfer reactions
<p>Vitamin A</p> 	Retinol, retinaldehyde, and retinoic acid	Formation of rhodopsin (vision) and glycoproteins (epithelial cell function); also regulates gene transcription
<p>Vitamin D</p> 	1,25-Dihydroxyvitamin D	Maintenance of blood calcium and phosphorus levels; antiproliferative hormone
<p>Vitamin E</p> 	Tocopherols and tocotrienols	Antioxidants
<p>Vitamin K</p> 	Vitamin K hydroquinone	Cofactor for posttranslation carboxylation of many proteins including essential clotting factors

FIGURE 96e-1 (Continued)

and peripheral neuritis. Patients with *dry beriberi* present with a symmetric peripheral neuropathy of the motor and sensory systems, with diminished reflexes. The neuropathy affects the legs most markedly, and patients have difficulty rising from a squatting position.

Alcoholic patients with chronic thiamine deficiency also may have central nervous system (CNS) manifestations known as *Wernicke's encephalopathy*, which consists of horizontal nystagmus, ophthalmoplegia (due to weakness of one or more extraocular muscles), cerebellar ataxia, and mental impairment (Chap. 467). When there is an additional loss of memory and a confabulatory psychosis, the syndrome is known as *Wernicke-Korsakoff syndrome*. Despite the typical clinical picture and history, Wernicke-Korsakoff syndrome is underdiagnosed.

The laboratory diagnosis of thiamine deficiency usually is made by a functional enzymatic assay of transketolase activity measured before and after the addition of thiamine pyrophosphate. A >25% stimulation in response to the addition of thiamine pyrophosphate (i.e., an activity coefficient of 1.25) is interpreted as abnormal. Thiamine or the phosphorylated esters of thiamine in serum or blood also can be measured by high-performance liquid chromatography to detect deficiency.

TREATMENT THIAMINE DEFICIENCY

In acute thiamine deficiency with either cardiovascular or neurologic signs, 200 mg of thiamine three times daily should be given intravenously until there is no further improvement in acute symptoms; oral thiamine (10 mg/d) should subsequently be given until recovery

is complete. Cardiovascular and ophthalmoplegic improvement occurs within 24 h. Other manifestations gradually clear, although psychosis in Wernicke-Korsakoff syndrome may be permanent or may persist for several months. Other nutrient deficiencies should be corrected concomitantly.

Toxicity Although anaphylaxis has been reported after high intravenous doses of thiamine, no adverse effects have been recorded from either food or supplements at high doses. Thiamine supplements may be bought over the counter in doses of up to 50 mg/d.

RIBOFLAVIN (VITAMIN B₂)

Riboflavin is important for the metabolism of fat, carbohydrate, and protein, acting as a respiratory coenzyme and an electron donor. Enzymes that contain flavin adenine dinucleotide (FAD) or flavin mononucleotide (FMN) as prosthetic groups are known as *flavoenzymes* (e.g., succinic acid dehydrogenase, monoamine oxidase, glutathione reductase). FAD is a cofactor for methyltetrahydrofolate reductase and therefore modulates homocysteine metabolism. The vitamin also plays a role in drug and steroid metabolism, including detoxification reactions.

Although much is known about the chemical and enzymatic reactions of riboflavin, the clinical manifestations of riboflavin deficiency are nonspecific and are similar to those of other deficiencies of B vitamins. Riboflavin deficiency is manifested principally by lesions of the mucocutaneous surfaces of the mouth and skin. In addition, corneal

vascularization, anemia, and personality changes have been described with riboflavin deficiency.

Deficiency and Excess Riboflavin deficiency almost always is due to dietary deficiency. Milk, other dairy products, and enriched breads and cereals are the most important dietary sources of riboflavin in the United States, although lean meat, fish, eggs, broccoli, and legumes are also good sources. Riboflavin is extremely sensitive to light, and milk should be stored in containers that protect against photodegradation. Laboratory diagnosis of riboflavin deficiency can be made by determination of red blood cell or urinary riboflavin concentrations or by measurement of erythrocyte glutathione reductase activity, with and without added FAD. Because the capacity of the gastrointestinal tract to absorb riboflavin is limited (~20 mg after one oral dose), riboflavin toxicity has not been described.

NIACIN (VITAMIN B₃)

The term *niacin* refers to nicotinic acid and nicotinamide and their biologically active derivatives. Nicotinic acid and nicotinamide serve as precursors of two coenzymes, nicotinamide adenine dinucleotide (NAD) and NAD phosphate (NADP), which are important in numerous oxidation and reduction reactions in the body. In addition, NAD and NADP are active in adenine diphosphate–ribose transfer reactions involved in DNA repair and calcium mobilization.

Metabolism and Requirements Nicotinic acid and nicotinamide are absorbed well from the stomach and small intestine. The bioavailability of niacin from beans, milk, meat, and eggs is high; bioavailability from cereal grains is lower. Since flour is enriched with “free” niacin (i.e., the non-coenzyme form), bioavailability is excellent. Median intakes of niacin in the United States considerably exceed the recommended dietary allowance (RDA).

The amino acid tryptophan can be converted to niacin with an efficiency of 60:1 by weight. Thus, the RDA for niacin is expressed in niacin equivalents. A lower-level conversion of tryptophan to niacin occurs in vitamin B₆ and/or riboflavin deficiencies and in the presence of isoniazid. The urinary excretion products of niacin include 2-pyridone and 2-methyl nicotinamide, measurements of which are used in the diagnosis of niacin deficiency.



Deficiency Niacin deficiency causes *pellagra*, which is found mostly among people eating corn-based diets in parts of China, Africa, and India. Pellagra in North America is found mainly among alcoholics; among patients with congenital defects of intestinal and kidney absorption of tryptophan (Hartnup disease; [Chap. 434e](#)); and among patients with carcinoid syndrome ([Chap. 113](#)), in which there is increased conversion of tryptophan to serotonin. The antituberculosis drug isoniazid is a structural analog of niacin and can precipitate pellagra. In the setting of famine or population displacement, pellagra results from the absolute lack of niacin but also from the deficiency of micronutrients required for the conversion of tryptophan to niacin (e.g., iron, riboflavin, and pyridoxine). The early symptoms of pellagra include loss of appetite, generalized weakness and irritability, abdominal pain, and vomiting. Bright red glossitis then ensues and is followed by a characteristic skin rash that is pigmented and scaling, particularly in skin areas exposed to sunlight. This rash is known as *Casal's necklace* because it forms a ring around the neck; it is seen in advanced cases. Vaginitis and esophagitis also may occur. Diarrhea (due in part to proctitis and in part to malabsorption), depression, seizures, and dementia are also part of the pellagra syndrome. The primary manifestations of this syndrome are sometimes referred to as “the four D’s”: dermatitis, diarrhea, and dementia leading to death.

TREATMENT PELLAGRA

Treatment of pellagra consists of oral supplementation with 100–200 mg of nicotinamide or nicotinic acid three times daily for 5 days. High doses of nicotinic acid (2 g/d in a time-release form) are used for the treatment of elevated cholesterol and triglyceride levels and/or low high-density lipoprotein cholesterol levels ([Chap. 421](#)).

Toxicity Prostaglandin-mediated flushing due to binding of the vitamin to a G protein–coupled receptor has been observed at daily nicotinic acid doses as low as 30 mg taken as a supplement or as therapy for dyslipidemia. There is no evidence of toxicity from niacin that is derived from food sources. Flushing always starts in the face and may be accompanied by skin dryness, itching, paresthesia, and headache. Pharmaceutical preparations of nicotinic acid combined with laropiprant, a selective prostaglandin D₂ receptor 1 antagonist, or premedication with aspirin may alleviate these symptoms. Flushing is subject to tachyphylaxis and often improves with time. Nausea, vomiting, and abdominal pain also occur at similar doses of niacin. Hepatic toxicity is the most serious toxic reaction caused by sustained-release niacin and may present as jaundice with elevated aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels. A few cases of fulminant hepatitis requiring liver transplantation have been reported at doses of 3–9 g/d. Other toxic reactions include glucose intolerance, hyperuricemia, macular edema, and macular cysts. The combination of nicotinic acid preparations for dyslipidemia with 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase inhibitors may increase the risk of rhabdomyolysis. The upper limit for daily niacin intake has been set at 35 mg. However, this upper limit does not pertain to the therapeutic use of niacin.

PYRIDOXINE (VITAMIN B₆)

Vitamin B₆ refers to a family of compounds that includes pyridoxine, pyridoxal, pyridoxamine, and their 5′-phosphate derivatives. 5′-Pyridoxal phosphate (PLP) is a cofactor for more than 100 enzymes involved in amino acid metabolism. Vitamin B₆ also is involved in heme and neurotransmitter synthesis and in the metabolism of glycogen, lipids, steroids, sphingoid bases, and several vitamins, including the conversion of tryptophan to niacin.

Dietary Sources Plants contain vitamin B₆ in the form of pyridoxine, whereas animal tissues contain PLP and pyridoxamine phosphate. The vitamin B₆ contained in plants is less bioavailable than that in animal tissues. Rich food sources of vitamin B₆ include legumes, nuts, wheat bran, and meat, although it is present in all food groups.

Deficiency Symptoms of vitamin B₆ deficiency include epithelial changes, as seen frequently with other B vitamin deficiencies. In addition, severe vitamin B₆ deficiency can lead to peripheral neuropathy, abnormal electroencephalograms, and personality changes that include depression and confusion. In infants, diarrhea, seizures, and anemia have been reported. Microcytic hypochromic anemia is due to diminished hemoglobin synthesis, since the first enzyme involved in heme biosynthesis (aminolevulinic synthase) requires PLP as a cofactor ([Chap. 126](#)). In some case reports, platelet dysfunction has been reported. Since vitamin B₆ is necessary for the conversion of homocysteine to cystathionine, it is possible that chronic low-grade vitamin B₆ deficiency may result in hyperhomocysteinemia and increased risk of cardiovascular disease ([Chaps. 291e and 434e](#)). Independent of homocysteine, low levels of circulating vitamin B₆ have been associated with inflammation and elevated levels of C-reactive protein.

Certain medications, such as isoniazid, L-dopa, penicillamine, and cycloserine, interact with PLP due to a reaction with carbonyl groups. Pyridoxine should be given concurrently with isoniazid to avoid neuropathy. The increased ratio of AST to ALT seen in alcoholic liver disease reflects the relative vitamin B₆ dependence of ALT. Vitamin B₆ dependency syndromes that require pharmacologic doses of vitamin B₆ are rare; they include cystathionine β-synthase deficiency, pyridoxine-responsive (primarily sideroblastic) anemias, and gyrate atrophy with chorioretinal degeneration due to decreased activity of the mitochondrial enzyme ornithine aminotransferase. In these situations, 100–200 mg/d of oral vitamin B₆ is required for treatment.

High doses of vitamin B₆ have been used to treat carpal tunnel syndrome, premenstrual syndrome, schizophrenia, autism, and diabetic neuropathy but have not been found to be effective.

The laboratory diagnosis of vitamin B₆ deficiency is generally based on low plasma PLP values (<20 nmol/L). Vitamin B₆ deficiency is treated with 50 mg/d; higher doses of 100–200 mg/d are given if the

deficiency is related to medication use. Vitamin B₆ should not be given with L-dopa, since the vitamin interferes with the action of this drug.

Toxicity The safe upper limit for vitamin B₆ has been set at 100 mg/d, although no adverse effects have been associated with high intakes of vitamin B₆ from food sources only. When toxicity occurs, it causes severe sensory neuropathy, leaving patients unable to walk. Some cases of photosensitivity and dermatitis have been reported.

FOLATE (VITAMIN B₁₂)

See Chap. 128.

VITAMIN C

Both ascorbic acid and its oxidized product dehydroascorbic acid are biologically active. Actions of vitamin C include antioxidant activity, promotion of nonheme iron absorption, carnitine biosynthesis, conversion of dopamine to norepinephrine, and synthesis of many peptide hormones. Vitamin C is also important for connective tissue metabolism and cross-linking (proline hydroxylation), and it is a component of many drug-metabolizing enzyme systems, particularly the mixed-function oxidase systems.

Absorption and Dietary Sources Vitamin C is almost completely absorbed if <100 mg is administered in a single dose; however, only 50% or less is absorbed at doses >1 g. Enhanced degradation and fecal and urinary excretion of vitamin C occur at higher intake levels.

Good dietary sources of vitamin C include citrus fruits, green vegetables (especially broccoli), tomatoes, and potatoes. Consumption of five servings of fruits and vegetables a day provides vitamin C in excess of the RDA of 90 mg/d for men and 75 mg/d for women. In addition, ~40% of the U.S. population consumes vitamin C as a dietary supplement in which “natural forms” of the vitamin are no more bioavailable than synthetic forms. Smoking, hemodialysis, pregnancy, and stress (e.g., infection, trauma) appear to increase vitamin C requirements.

Deficiency Vitamin C deficiency causes scurvy. In the United States, this condition is seen primarily among the poor and the elderly, in alcoholics who consume <10 mg/d of vitamin C, and in individuals consuming macrobiotic diets. Vitamin C deficiency also can occur in young adults who eat severely unbalanced diets. In addition to generalized fatigue, symptoms of scurvy primarily reflect impaired formation of mature connective tissue and include bleeding into the skin (petechiae, ecchymoses, perifollicular hemorrhages); inflamed and bleeding gums; and manifestations of bleeding into joints, the peritoneal cavity, the pericardium, and the adrenal glands. In children, vitamin C deficiency may cause impaired bone growth. Laboratory diagnosis of vitamin C deficiency is based on low plasma or leukocyte levels.

Administration of vitamin C (200 mg/d) improves the symptoms of scurvy within several days. High-dose vitamin C supplementation (e.g., 1–2 g/d) may slightly decrease the symptoms and duration of upper respiratory tract infections. Vitamin C supplementation has also been reported to be useful in Chédiak-Higashi syndrome (Chap. 80) and osteogenesis imperfecta (Chap. 427). Diets high in vitamin C have been claimed to lower the incidence of certain cancers, particularly esophageal and gastric cancers. If proved, this effect may be due to the fact that vitamin C can prevent the conversion of nitrites and secondary amines to carcinogenic nitrosamines. However, an intervention study from China did not show vitamin C to be protective. A potential role for parenteral ascorbic acid in the treatment of advanced cancers has been suggested.

Toxicity Taking >2 g of vitamin C in a single dose may result in abdominal pain, diarrhea, and nausea. Since vitamin C may be metabolized to oxalate, it is feared that chronic high-dose vitamin C supplementation could result in an increased prevalence of kidney stones. However, except in patients with preexisting renal disease, this association has not been borne out in several trials. Nevertheless, it is reasonable to advise patients with a history of kidney stones not to take large doses of vitamin C. There is also an unproven but possible risk that chronic high doses of vitamin C could promote iron overload and iron toxicity. High doses of vitamin C can induce hemolysis in patients

with glucose-6-phosphate dehydrogenase deficiency, and doses >1 g/d can cause false-negative guaiac reactions and interfere with tests for urinary glucose. High doses may interfere with the activity of certain drugs (e.g., bortezomib in myeloma patients).

BIOTIN

Biotin is a water-soluble vitamin that plays a role in gene expression, gluconeogenesis, and fatty acid synthesis and serves as a CO₂ carrier on the surface of both cytosolic and mitochondrial carboxylase enzymes. The vitamin also functions in the catabolism of specific amino acids (e.g., leucine) and in gene regulation by histone biotinylation. Excellent food sources of biotin include organ meat such as liver or kidney, soy and other beans, yeast, and egg yolks; however, egg white contains the protein avidin, which strongly binds the vitamin and reduces its bioavailability.

Biotin deficiency due to low dietary intake is rare; rather, deficiency is due to inborn errors of metabolism. Biotin deficiency has been induced by experimental feeding of egg white diets and by biotin-free parenteral nutrition in patients with short bowels. In adults, biotin deficiency results in mental changes (depression, hallucinations), paresthesia, anorexia, and nausea. A scaling, seborrheic, and erythematous rash may occur around the eyes, nose, and mouth as well as on the extremities. In infants, biotin deficiency presents as hypotonia, lethargy, and apathy. In addition, infants may develop alopecia and a characteristic rash that includes the ears. The laboratory diagnosis of biotin deficiency can be established on the basis of a decreased concentration of urinary biotin (or its major metabolites), increased urinary excretion of 3-hydroxyisovaleric acid after a leucine challenge, or decreased activity of biotin-dependent enzymes in lymphocytes (e.g., propionyl-CoA carboxylase). Treatment requires pharmacologic doses of biotin—i.e., up to 10 mg/d. No toxicity is known.

PANTOTHENIC ACID (VITAMIN B₅)

Pantothenic acid is a component of coenzyme A and phosphopantetheine, which are involved in fatty acid metabolism and the synthesis of cholesterol, steroid hormones, and all compounds formed from isoprenoid units. In addition, pantothenic acid is involved in the acetylation of proteins. The vitamin is excreted in the urine, and the laboratory diagnosis of deficiency is based on low urinary vitamin levels.

The vitamin is ubiquitous in the food supply. Liver, yeast, egg yolks, whole grains, and vegetables are particularly good sources. Human pantothenic acid deficiency has been demonstrated only by experimental feeding of diets low in pantothenic acid or by administration of a specific pantothenic acid antagonist. The symptoms of pantothenic acid deficiency are nonspecific and include gastrointestinal disturbance, depression, muscle cramps, paresthesia, ataxia, and hypoglycemia. Pantothenic acid deficiency is believed to have caused the “burning feet syndrome” seen in prisoners of war during World War II. No toxicity of this vitamin has been reported.

CHOLINE

Choline is a precursor for acetylcholine, phospholipids, and betaine. Choline is necessary for the structural integrity of cell membranes, cholinergic neurotransmission, lipid and cholesterol metabolism, methyl-group metabolism, and transmembrane signaling. Recently, a recommended adequate intake was set at 550 mg/d for men and 425 mg/d for women, although certain genetic polymorphisms can increase an individual’s requirement. Choline is thought to be a “conditionally essential” nutrient in that its de novo synthesis occurs in the liver and results in lesser-than-used amounts only under certain stress conditions (e.g., alcoholic liver disease). The dietary requirement for choline depends on the status of other nutrients involved in methyl-group metabolism (folate, vitamin B₁₂, vitamin B₆, and methionine) and thus varies widely. Choline is widely distributed in food (e.g., egg yolks, wheat germ, organ meat, milk) in the form of lecithin (phosphatidylcholine). Choline deficiency has occurred in patients receiving parenteral nutrition devoid of choline. Deficiency results in fatty liver, elevated aminotransferase levels, and skeletal muscle damage with high creatine phosphokinase values. The diagnosis of choline deficiency is

currently based on low plasma levels, although nonspecific conditions (e.g., heavy exercise) may also suppress plasma levels.

Toxicity from choline results in hypotension, cholinergic sweating, diarrhea, salivation, and a fishy body odor. The upper limit for choline intake has been set at 3.5 g/d. Because of its ability to lower cholesterol and homocysteine levels, choline treatment has been suggested for patients with dementia and patients at high risk of cardiovascular disease. However, the benefits of such treatment have not been firmly documented. Choline- and betaine-restricted diets are of therapeutic value in trimethylaminuria (“fish odor syndrome”).

FLAVONOIDS

Flavonoids constitute a large family of polyphenols that contribute to the aroma, taste, and color of fruits and vegetables. Major groups of dietary flavonoids include anthocyanidins in berries; catechins in green tea and chocolate; flavonols (e.g., quercetin) in broccoli, kale, leeks, onions, and the skins of grapes and apples; and isoflavones (e.g., genistein) in legumes. Isoflavones have a low bioavailability and are partially metabolized by the intestinal flora. The dietary intake of flavonoids is estimated at 10–100 mg/d; this figure is almost certainly an underestimate attributable to a lack of information on their concentrations in many foods. Several flavonoids have antioxidant activity and affect cell signaling. From observational epidemiologic studies and limited clinical (human and animal) studies, flavonoids have been postulated to play a role in the prevention of several chronic diseases, including neurodegenerative disease, diabetes, and osteoporosis. The ultimate importance and usefulness of these compounds against human disease have not been demonstrated.

VITAMIN A

Vitamin A, in the strictest sense, refers to retinol. However, the oxidized metabolites retinaldehyde and retinoic acid are also biologically active compounds. The term *retinoids* includes all molecules (including synthetic molecules) that are chemically related to retinol. Retinaldehyde (11-*cis*) is the essential form of vitamin A that is required for normal vision, whereas retinoic acid is necessary for normal morphogenesis, growth, and cell differentiation. Retinoic acid does not function in vision and, in contrast to retinol, is not involved in reproduction. Vitamin A also plays a role in iron utilization, humoral immunity, T cell-mediated immunity, natural killer cell activity, and phagocytosis. Vitamin A is commercially available in esterified forms (e.g., acetate, palmitate), which are more stable than other forms.

There are more than 600 carotenoids in nature, ~50 of which can be metabolized to vitamin A. β -Carotene is the most prevalent carotenoid with provitamin A activity in the food supply. In humans, significant fractions of carotenoids are absorbed intact and are stored in liver and fat. It is estimated that ≥ 12 μg (range, 4–27 μg) of dietary all-*trans* β -carotene is equivalent to 1 μg of retinol activity, whereas the figure is ≥ 24 μg for other dietary provitamin A carotenoids (e.g., cryptoxanthin, α -carotene). The vitamin A equivalency for a β -carotene supplement in an oily solution is 2:1.

Metabolism The liver contains ~90% of the vitamin A reserves and secretes vitamin A in the form of retinol, which is bound to retinol-binding protein. Once binding has occurred, the retinol-binding protein complex interacts with a second protein, transthyretin. This trimolecular complex functions to prevent vitamin A from being filtered by the kidney glomerulus, thus protecting the body against the toxicity of retinol and allowing retinol to be taken up by specific cell-surface receptors that recognize retinol-binding protein. A certain amount of vitamin A enters peripheral cells even if it is not bound to retinol-binding protein. After retinol is internalized by the cell, it becomes bound to a series of cellular retinol-binding proteins, which function as sequestering and transporting agents as well as co-ligands for enzymatic reactions. Certain cells also contain retinoic acid-binding proteins, which have sequestering functions but also shuttle retinoic acid to the nucleus and enable its metabolism.

Retinoic acid is a ligand for certain nuclear receptors that act as transcription factors. Two families of receptors (retinoic acid receptors

[RARs] and retinoid X receptors [RXRs]) are active in retinoid-mediated gene transcription. Retinoid receptors regulate transcription by binding as dimeric complexes to specific DNA sites—the retinoic acid response elements—in target genes (Chap. 400e). The receptors can either stimulate or repress gene expression in response to their ligands. RARs bind all-*trans* retinoic acid and 9-*cis*-retinoic acid, whereas RXRs bind only 9-*cis*-retinoic acid.

The retinoid receptors play an important role in controlling cell proliferation and differentiation. Retinoic acid is useful in the treatment of promyelocytic leukemia (Chap. 132) and also is used in the treatment of cystic acne because it inhibits keratinization, decreases sebum secretion, and possibly alters the inflammatory reaction (Chap. 71). RXRs dimerize with other nuclear receptors to function as coregulators of genes responsive to retinoids, thyroid hormone, and calcitriol. RXR agonists induce insulin sensitivity experimentally, perhaps because RXRs are cofactors for the peroxisome proliferator-activated receptors, which are targets for thiazolidinedione drugs such as rosiglitazone and troglitazone (Chap. 418).

Dietary Sources The retinol activity equivalent (RAE) is used to express the vitamin A value of food: 1 RAE is defined as 1 μg of retinol (0.003491 mmol), 12 μg of β -carotene, and 24 μg of other provitamin A carotenoids. In older literature, vitamin A often was expressed in international units (IU), with 1 μg of retinol equal to 3.33 IU of retinol and 20 IU of β -carotene, but these units are no longer in scientific use.

Liver, fish, and eggs are excellent food sources for preformed vitamin A; vegetable sources of provitamin A carotenoids include dark green and deeply colored fruits and vegetables. Moderate cooking of vegetables enhances carotenoid release for uptake in the gut. Carotenoid absorption is also aided by some fat in a meal. Infants are particularly susceptible to vitamin A deficiency because neither breast nor cow's milk supplies enough vitamin A to prevent deficiency. In developing countries, chronic dietary deficiency is the main cause of vitamin A deficiency and is exacerbated by infection. In early childhood, low vitamin A status results from inadequate intakes of animal food sources and edible oils, both of which are expensive, coupled with seasonal unavailability of vegetables and fruits and lack of marketed fortified food products. Concurrent zinc deficiency can interfere with the mobilization of vitamin A from liver stores. Alcohol interferes with the conversion of retinol to retinaldehyde in the eye by competing for alcohol (retinol) dehydrogenase. Drugs that interfere with the absorption of vitamin A include mineral oil, neomycin, and cholestyramine.



Deficiency Vitamin A deficiency is endemic in areas where diets are chronically poor, especially in southern Asia, sub-Saharan Africa, some parts of Latin America, and the western Pacific, including parts of China. Vitamin A status is usually assessed by measuring serum retinol (normal range, 1.05–3.50 $\mu\text{mol/L}$ [30–100 $\mu\text{g/dL}$]) or blood-spot retinol or by tests of dark adaptation. Stable isotopic or invasive liver biopsy methods are available to estimate total body stores of vitamin A. As judged by deficient serum retinol (< 0.70 $\mu\text{mol/L}$ [20 $\mu\text{g/dL}$]), vitamin A deficiency worldwide is present in > 90 million preschool-age children, among whom > 4 million have an ocular manifestation of deficiency termed *xerophthalmia*. This condition includes milder stages of night blindness and conjunctival *xerosis* (dryness) with *Bitot's spots* (white patches of keratinized epithelium appearing on the sclera) as well as rare, potentially blinding corneal ulceration and necrosis. *Keratomalacia* (softening of the cornea) leads to corneal scarring that blinds at least a quarter of a million children each year and is associated with fatality rates of 4–25%. However, vitamin A deficiency at any stage poses an increased risk of death from diarrhea, dysentery, measles, malaria, or respiratory disease. Vitamin A deficiency can compromise barrier, innate, and acquired immune defenses to infection. In areas where deficiency is widely prevalent, vitamin A supplementation can markedly reduce the risk of childhood mortality (by 23–34%, on average). About 10% of pregnant women in undernourished settings also develop night blindness (assessed by history) during the latter half of pregnancy, and this moderate vitamin A deficiency is associated with an increased risk of maternal infection and death.

TREATMENT VITAMIN A DEFICIENCY

Any stage of xerophthalmia should be treated with 60 mg (or RAE) of vitamin A in oily solution, usually contained in a soft-gel capsule. The same dose is repeated 1 and 14 days later. Doses should be reduced by half for patients 6–11 months of age. Mothers with night blindness or Bitot's spots should be given vitamin A orally—either 3 mg daily or 7.5 mg twice a week for 3 months. These regimens are efficacious, and they are less expensive and more widely available than injectable water-miscible vitamin A. A common approach to prevention is to provide vitamin A supplementation every 4–6 months to young children and infants (both HIV-positive and HIV-negative) in high-risk areas. Infants 6–11 months of age should receive 30 mg vitamin A; children 12–59 months of age, 60 mg. For reasons that are not clear, vitamin A supplementation has not proven useful in high-risk settings for preventing morbidity or death among infants 1–5 months of age.

Uncomplicated vitamin A deficiency is rare in industrialized countries. One high-risk group—extremely low-birth-weight (<1000-g) infants—is likely to be vitamin A–deficient and should receive a supplement of 1500 µg (or RAE) three times a week for 4 weeks. Severe measles in any society can lead to secondary vitamin A deficiency. Children hospitalized with measles should receive two 60-mg doses of vitamin A on two consecutive days. Vitamin A deficiency most often occurs in patients with malabsorptive diseases (e.g., celiac sprue, short-bowel syndrome) who have abnormal dark adaptation or symptoms of night blindness without other ocular changes. Typically, such patients are treated for 1 month with 15 mg/d of a water-miscible preparation of vitamin A. This treatment is followed by a lower maintenance dose, with the exact amount determined by monitoring serum retinol.

No specific signs or symptoms result from carotenoid deficiency. It was postulated that β-carotene would be an effective chemopreventive agent for cancer because numerous epidemiologic studies had shown that diets high in β-carotene were associated with lower incidences of cancers of the respiratory and digestive systems. However, intervention studies in smokers found that treatment with high doses of β-carotene actually resulted in more lung cancers than did treatment with placebo. Non-provitamin A carotenoids such as lutein and zeaxanthin have been suggested to confer protection against macular degeneration, and one large-scale intervention study did not show a beneficial effect except in those with a low lutein status. The use of the non-provitamin A carotenoid lycopene to protect against prostate cancer has been proposed. Again, however, the effectiveness of these agents has not been proved by intervention studies, and the mechanisms underlying these purported biologic actions are unknown.

Selective plant-breeding techniques that lead to a higher provitamin A content in staple foods may decrease vitamin A malnutrition in low-income countries. Moreover, a recently developed genetically modified food (Golden Rice) has an improved β-carotene-to-vitamin A conversion ratio of ~3:1.

Toxicity The acute toxicity of vitamin A was first noted in Arctic explorers who ate polar bear liver and has also been seen after administration of 150 mg to adults or 100 mg to children. Acute toxicity is manifested by increased intracranial pressure, vertigo, diplopia, bulging fontanels (in children), seizures, and exfoliative dermatitis; it may result in death. Among children being treated for vitamin A deficiency according to the protocols outlined above, transient bulging of fontanels occurs in 2% of infants, and transient nausea, vomiting, and headache occur in 5% of preschoolers. Chronic vitamin A intoxication is largely a concern in industrialized countries and has been seen in otherwise healthy adults who ingest 15 mg/d and children who ingest 6 mg/d over a period of several months. Manifestations include dry skin, cheilosis, glossitis, vomiting, alopecia, bone demineralization and pain, hypercalcemia, lymph node enlargement, hyperlipidemia, amenorrhea, and features of pseudotumor cerebri with increased intracranial

pressure and papilledema. Liver fibrosis with portal hypertension and bone demineralization may result from chronic vitamin A intoxication. Provision of vitamin A in excess to pregnant women has resulted in spontaneous abortion and in congenital malformations, including craniofacial abnormalities and valvular heart disease. In pregnancy, the daily dose of vitamin A should not exceed 3 mg. Commercially available retinoid derivatives are also toxic, including 13-*cis*-retinoic acid, which has been associated with birth defects. Thus contraception should be continued for at least 1 year and possibly longer in women who have taken 13-*cis*-retinoic acid.

In malnourished children, vitamin A supplements (30–60 mg), in amounts calculated as a function of age and given in several rounds over 2 years, are considered to amplify nonspecific effects of vaccines. However, for unclear reasons, there may be a negative effect on mortality rates in incompletely vaccinated girls.

High doses of carotenoids do not result in toxic symptoms but should be avoided in smokers due to an increased risk of lung cancer. Very high doses of β-carotene (~200 mg/d) have been used to treat or prevent the skin rashes of erythropoietic protoporphyria. Carotenemia, which is characterized by a yellowing of the skin (in creases of the palms and soles) but not the sclerae, may follow ingestion of >30 mg of β-carotene daily. Hypothyroid patients are particularly susceptible to the development of carotenemia due to impaired breakdown of carotene to vitamin A. Reduction of carotenes in the diet results in the disappearance of skin yellowing and carotenemia over a period of 30–60 days.

VITAMIN D

The metabolism of the fat-soluble vitamin D is described in detail in [Chap. 423](#). The biologic effects of this vitamin are mediated by vitamin D receptors, which are found in most tissues; binding with these receptors potentially expands vitamin D actions on nearly all cell systems and organs (e.g., immune cells, brain, breast, colon, and prostate) as well as exerting classic endocrine effects on calcium metabolism and bone health. Vitamin D is thought to be important for maintaining normal function of many nonskeletal tissues such as muscle (including heart muscle), for immune function, and for inflammation as well as for cell proliferation and differentiation. Studies have shown that vitamin D may be useful as adjunctive treatment for tuberculosis, psoriasis, and multiple sclerosis or for the prevention of certain cancers. Vitamin D insufficiency may increase the risk of type 1 diabetes mellitus, cardiovascular disease (insulin resistance, hypertension, or low-grade inflammation), or brain dysfunction (e.g., depression). However, the exact physiologic roles of vitamin D in these nonskeletal diseases and the importance of these roles have not been clarified.

The skin is a major source of vitamin D, which is synthesized upon skin exposure to ultraviolet B radiation (UV-B; wavelength, 290–320 nm). Except for fish, food (unless fortified) contains only limited amounts of vitamin D. Vitamin D₂ (ergocalciferol) is obtained from plant sources and is the chemical form found in some supplements.

Deficiency Vitamin D status has been assessed by measuring serum levels of 25-dihydroxyvitamin D (25[OH]₂ vitamin D); however, there is no consensus on a uniform assay or on optimal serum levels. The optimal level might, in fact, differ according to the targeted disease entity. Epidemiologic and experimental data indicate that a 25(OH)₂ vitamin D level of >20 ng/mL (≥50 nmol/L; to convert ng/mL to nmol/L, multiply by 2.496) is sufficient for good bone health. Some experts advocate higher serum levels (e.g., >30 ng/mL) for other desirable endpoints of vitamin D action. There is insufficient evidence to recommend combined vitamin D and calcium supplementation as a primary preventive strategy for reduction of the incidence of fractures in healthy men and premenopausal women.

Risk factors for vitamin D deficiency are old age, lack of sun exposure, dark skin (especially among residents of northern latitudes), fat malabsorption, and obesity. *Rickets* represents the classic disease of vitamin D deficiency. Signs of deficiency are muscle soreness, weakness, and bone pain. Some of these effects are independent of calcium intake.

The U.S. National Academy of Sciences recently concluded that the majority of North Americans are receiving adequate amounts of vitamin D (RDA = 15 µg/d or 600 IU/d; **Chap. 95e**). However, for people older than 70 years, the RDA is set at 20 µg/d (800 IU/d). The consumption of fortified or enriched foods as well as suberythral sun exposure should be encouraged for people at risk for vitamin D deficiency. If adequate intake is impossible, vitamin D supplements should be taken, especially during the winter months. Vitamin D deficiency can be treated by the oral administration of 50,000 IU/week for 6–8 weeks followed by a maintenance dose of 800 IU/d (100 µg/d) from food and supplements once normal plasma levels have been attained. The physiologic effects of vitamin D₂ and vitamin D₃ are identical when these vitamins are ingested over long periods.

Toxicity The upper limit of intake has been set at 4000 IU/d. Contrary to earlier beliefs, acute vitamin D intoxication is rare and usually is caused by the uncontrolled and excessive ingestion of supplements or by faulty food fortification practices. High plasma levels of 1,25(OH)₂ vitamin D and calcium are central features of toxicity and mandate discontinuation of vitamin D and calcium supplements; in addition, treatment of hypercalcemia may be required.

VITAMIN E

Vitamin E is the collective designation for all stereoisomers of tocopherols and tocotrienols, although only the RR tocopherols meet human requirements. Vitamin E acts as a chain-breaking antioxidant and is an efficient pyroxyl radical scavenger that protects low-density lipoproteins and polyunsaturated fats in membranes from oxidation. A network of other antioxidants (e.g., vitamin C, glutathione) and enzymes maintains vitamin E in a reduced state. Vitamin E also inhibits prostaglandin synthesis and the activities of protein kinase C and phospholipase A₂.

Absorption and Metabolism After absorption, vitamin E is taken up from chylomicrons by the liver, and a hepatic α-tocopherol transport protein mediates intracellular vitamin E transport and incorporation into very low density lipoprotein. The transport protein has a particular affinity for the RRR isomeric form of α-tocopherol; thus, this natural isomer has the most biologic activity.

Requirement Vitamin E is widely distributed in the food supply, with particularly high levels in sunflower oil, safflower oil, and wheat germ oil; γ-tocotrienols are notably present in soybean and corn oils. Vitamin E is also found in meats, nuts, and cereal grains, and small amounts are present in fruits and vegetables. Vitamin E pills containing doses of 50–1000 mg are ingested by ~10% of the U.S. population. The RDA for vitamin E is 15 mg/d (34.9 µmol or 22.5 IU) for all adults. Diets high in polyunsaturated fats may necessitate a slightly higher intake of vitamin E.

Dietary deficiency of vitamin E does not exist. Vitamin E deficiency is seen only in severe and prolonged malabsorptive diseases, such as celiac disease, or after small-intestinal resection or bariatric surgery. Children with cystic fibrosis or prolonged cholestasis may develop vitamin E deficiency characterized by areflexia and hemolytic anemia. Children with abetalipoproteinemia cannot absorb or transport vitamin E and become deficient quite rapidly. A familial form of isolated vitamin E deficiency also exists; it is due to a defect in the α-tocopherol transport protein. Vitamin E deficiency causes axonal degeneration of the large myelinated axons and results in posterior column and spinocerebellar symptoms. Peripheral neuropathy is initially characterized by areflexia, with progression to an ataxic gait, and by decreased vibration and position sensations. Ophthalmoplegia, skeletal myopathy, and pigmented retinopathy may also be features of vitamin E deficiency. A deficiency of either vitamin E or selenium in the host has been shown to increase certain viral mutations and, therefore, virulence. The laboratory diagnosis of vitamin E deficiency is based on low blood levels of α-tocopherol (<5 µg/mL, or <0.8 mg of α-tocopherol per gram of total lipids).

TREATMENT VITAMIN E DEFICIENCY

Symptomatic vitamin E deficiency should be treated with 800–1200 mg of α-tocopherol per day. Patients with abetalipoproteinemia may need as much as 5000–7000 mg/d. Children with symptomatic vitamin E deficiency should be treated orally with water-miscible esters (400 mg/d); alternatively, 2 mg/kg per day may be administered intramuscularly. Vitamin E in high doses may protect against oxygen-induced retrolental fibroplasia and bronchopulmonary dysplasia as well as intraventricular hemorrhage of prematurity. Vitamin E has been suggested to increase sexual performance, treat intermittent claudication, and slow the aging process, but evidence for these properties is lacking. When given in combination with other antioxidants, vitamin E may help prevent macular degeneration. High doses (60–800 mg/d) of vitamin E have been shown in controlled trials to improve parameters of immune function and reduce colds in nursing home residents, but intervention studies using vitamin E to prevent cardiovascular disease or cancer have not shown efficacy, and, at doses >400 mg/d, vitamin E may even increase all-cause mortality rates.

Toxicity All forms of vitamin E are absorbed and could contribute to toxicity; however, the toxicity risk seems to be rather low as long as liver function is normal. High doses of vitamin E (>800 mg/d) may reduce platelet aggregation and interfere with vitamin K metabolism and are therefore contraindicated in patients taking warfarin and antiplatelet agents (such as aspirin or clopidogrel). Nausea, flatulence, and diarrhea have been reported at doses >1 g/d.

VITAMIN K

There are two natural forms of vitamin K: vitamin K₁, also known as *phylloquinone*, from vegetable and animal sources, and vitamin K₂, or *menaquinone*, which is synthesized by bacterial flora and found in hepatic tissue. Phylloquinone can be converted to menaquinone in some organs.

Vitamin K is required for the posttranslational carboxylation of glutamic acid, which is necessary for calcium binding to γ-carboxylated proteins such as prothrombin (factor II); factors VII, IX, and X; protein C; protein S; and proteins found in bone (osteocalcin) and vascular smooth muscle (e.g., matrix Gla protein). However, the importance of vitamin K for bone mineralization and prevention of vascular calcification is not known. Warfarin-type drugs inhibit γ-carboxylation by preventing the conversion of vitamin K to its active hydroquinone form.

Dietary Sources Vitamin K is found in green leafy vegetables such as kale and spinach, and appreciable amounts are also present in margarine and liver. Vitamin K is present in vegetable oils; olive, canola, and soybean oils are particularly rich sources. The average daily intake by Americans is estimated to be ~100 µg/d.

Deficiency The symptoms of vitamin K deficiency are due to hemorrhage; newborns are particularly susceptible because of low fat stores, low breast milk levels of vitamin K, relative sterility of the infantile intestinal tract, liver immaturity, and poor placental transport. Intracranial bleeding as well as gastrointestinal and skin bleeding can occur in vitamin K-deficient infants 1–7 days after birth. Thus, vitamin K (0.5–1 mg IM) is given prophylactically at delivery.

Vitamin K deficiency in adults may be seen in patients with chronic small-intestinal disease (e.g., celiac disease, Crohn's disease), in those with obstructed biliary tracts, or after small-bowel resection. Broad-spectrum antibiotic treatment can precipitate vitamin K deficiency by reducing numbers of gut bacteria, which synthesize menaquinones, and by inhibiting the metabolism of vitamin K. In patients with warfarin therapy, the anti-obesity drug orlistat can lead to international normalized ratio changes due to vitamin K malabsorption. Vitamin K deficiency usually is diagnosed on the basis of an elevated prothrombin time or reduced clotting factors, although vitamin K may also be measured directly by high-pressure liquid chromatography. Vitamin K deficiency

is treated with a parenteral dose of 10 mg. For patients with chronic malabsorption, 1–2 mg/d should be given orally or 1–2 mg per week can be taken parenterally. Patients with liver disease may have an elevated prothrombin time because of liver cell destruction as well as vitamin K deficiency. If an elevated prothrombin time does not improve during vitamin K therapy, it can be deduced that this abnormality is not the result of vitamin K deficiency.

Toxicity Toxicity from dietary phyloquinones and menaquinones has not been described. High doses of vitamin K can impair the actions of oral anticoagulants.

MINERALS

See also Table 96e-2.

CALCIUM

See Chap. 423.

ZINC

Zinc is an integral component of many metalloenzymes in the body; it is involved in the synthesis and stabilization of proteins, DNA, and RNA and plays a structural role in ribosomes and membranes. Zinc is necessary for the binding of steroid hormone receptors and several other transcription factors to DNA. Zinc is absolutely required for normal spermatogenesis, fetal growth, and embryonic development.

Absorption The absorption of zinc from the diet is inhibited by dietary phytate, fiber, oxalate, iron, and copper as well as by certain drugs, including penicillamine, sodium valproate, and ethambutol. Meat, shellfish, nuts, and legumes are good sources of bioavailable zinc, whereas zinc in grains and legumes is less available for absorption.



Deficiency Mild zinc deficiency has been described in many diseases, including diabetes mellitus, HIV/AIDS, cirrhosis, alcoholism, inflammatory bowel disease, malabsorption syndromes, and sickle cell disease. In these diseases, mild chronic zinc deficiency can cause stunted growth in children, decreased taste sensation (*hypogeusia*), and impaired immune function. Severe chronic zinc deficiency has been described as a cause of hypogonadism and dwarfism in several Middle Eastern countries. In these children, hypopigmented hair is also part of the syndrome. Acrodermatitis enteropathica is a rare autosomal recessive disorder characterized by abnormalities in zinc absorption. Clinical manifestations include diarrhea, alopecia, muscle wasting, depression, irritability, and a rash involving the extremities, face, and perineum. The rash is characterized by vesicular and pustular crusting with scaling and erythema. Occasional patients with Wilson's disease have developed zinc deficiency as a consequence of penicillamine therapy (Chap. 429).

Zinc deficiency is prevalent in many developing countries and usually coexists with other micronutrient deficiencies (especially iron deficiency). Zinc (20 mg/d until recovery) may be an effective adjunctive therapeutic strategy for diarrheal disease and pneumonia in children ≥ 6 months of age.

The diagnosis of zinc deficiency is usually based on a serum zinc level $<12 \mu\text{mol/L}$ ($<70 \mu\text{g/dL}$). Pregnancy and birth control pills may cause a slight depression in serum zinc levels, and hypoalbuminemia from any cause can result in hypozincemia. In acute stress situations, zinc may be redistributed from serum into tissues. Zinc deficiency may be treated with 60 mg of elemental zinc taken by mouth twice a day. Zinc gluconate lozenges (13 mg of elemental zinc every 2 h while awake) have been reported to reduce the duration and symptoms of the common cold in adults, but study results are conflicting.

TABLE 96e-2 DEFICIENCIES AND TOXICITIES OF METALS

Element	Deficiency	Toxicity	Tolerable Upper (Dietary) Intake Level
Boron	No biologic function determined	Developmental defects, male sterility, testicular atrophy	20 mg/d (extrapolated from animal data)
Calcium	Reduced bone mass, osteoporosis	Renal insufficiency (milk-alkali syndrome), nephrolithiasis, impaired iron absorption, thiazide diuretics	2500 mg/d (milk-alkali)
Copper	Anemia, growth retardation, defective keratinization and pigmentation of hair, hypothermia, degenerative changes in aortic elastin, osteopenia, mental deterioration	Nausea, vomiting, diarrhea, hepatic failure, tremor, mental deterioration, hemolytic anemia, renal dysfunction	10 mg/d (liver toxicity)
Chromium	Impaired glucose tolerance	<i>Occupational:</i> Renal failure, dermatitis, pulmonary cancer	Not determined
Fluoride	\uparrow Dental caries	Dental and skeletal fluorosis, osteosclerosis	10 mg/d (fluorosis)
Iodine	Thyroid enlargement, $\downarrow T_4$, cretinism	Thyroid dysfunction, acne-like eruptions	1100 $\mu\text{g/d}$ (thyroid dysfunction)
Iron	Muscle abnormalities, koilonychia, pica, anemia, \downarrow work performance, impaired cognitive development, premature labor, \uparrow perinatal maternal death	Gastrointestinal effects (nausea, vomiting, diarrhea, constipation), iron overload with organ damage, acute and chronic systemic toxicity, increased susceptibility to malaria, increased risk association with certain chronic diseases (e.g., diabetes)	45 mg/d of elemental iron (gastrointestinal side effects)
Manganese	Impaired growth and skeletal development, reproduction, lipid and carbohydrate metabolism; upper body rash	<i>General:</i> Neurotoxicity, Parkinson-like symptoms <i>Occupational:</i> Encephalitis-like syndrome, Parkinson-like syndrome, psychosis, pneumoconiosis	11 mg/d (neurotoxicity)
Molybdenum	Severe neurologic abnormalities	Reproductive and fetal abnormalities	2 mg/d (extrapolated from animal data)
Selenium	Cardiomyopathy, heart failure, striated muscle degeneration	<i>General:</i> Alopecia, nausea, vomiting, abnormal nails, emotional lability, peripheral neuropathy, lassitude, garlic odor to breath, dermatitis <i>Occupational:</i> Lung and nasal carcinomas, liver necrosis, pulmonary inflammation	400 $\mu\text{g/d}$ (hair, nail changes)
Phosphorus	Rickets (osteomalacia), proximal muscle weakness, rhabdomyolysis, paresthesia, ataxia, seizure, confusion, heart failure, hemolysis, acidosis	Hyperphosphatemia	4000 mg/d
Zinc	Growth retardation, \downarrow taste and smell, alopecia, dermatitis, diarrhea, immune dysfunction, failure to thrive, gonadal atrophy, congenital malformations	<i>General:</i> Reduced copper absorption, gastritis, sweating, fever, nausea, vomiting <i>Occupational:</i> Respiratory distress, pulmonary fibrosis	40 mg/d (impaired copper metabolism)

96e-10 Toxicity Acute zinc toxicity after oral ingestion causes nausea, vomiting, and fever. Zinc fumes from welding may also be toxic and cause fever, respiratory distress, excessive salivation, sweating, and headache. Chronic large doses of zinc may depress immune function and cause hypochromic anemia as a result of copper deficiency. Intranasal zinc preparations should be avoided because they may lead to irreversible damage of the nasal mucosa and anosmia.

COPPER

Copper is an integral part of numerous enzyme systems, including amine oxidases, ferroxidase (ceruloplasmin), cytochrome c oxidase, superoxide dismutase, and dopamine hydroxylase. Copper is also a component of ferroprotein, a transport protein involved in the basolateral transfer of iron during absorption from the enterocyte. As such, copper plays a role in iron metabolism, melanin synthesis, energy production, neurotransmitter synthesis, and CNS function; the synthesis and cross-linking of elastin and collagen; and the scavenging of superoxide radicals. Dietary sources of copper include shellfish, liver, nuts, legumes, bran, and organ meats.

Deficiency Dietary copper deficiency is relatively rare, although it has been described in premature infants who are fed milk diets and in infants with malabsorption (Table 96e-2). Copper-deficiency anemia (refractory to therapeutic iron) has been reported in patients with malabsorptive diseases and nephrotic syndrome and in patients treated for Wilson's disease with chronic high doses of oral zinc, which can interfere with copper absorption. *Menkes kinky hair syndrome* is an X-linked metabolic disturbance of copper metabolism characterized by mental retardation, hypocupremia, and decreased circulating ceruloplasmin (Chap. 427). This syndrome is caused by mutations in the copper-transporting *ATP7A* gene. Children with this disease often die within 5 years because of dissecting aneurysms or cardiac rupture. Aceruloplasminemia is a rare autosomal recessive disease characterized by tissue iron overload, mental deterioration, microcytic anemia, and low serum iron and copper concentrations.

The diagnosis of copper deficiency is usually based on low serum levels of copper (<65 µg/dL) and low ceruloplasmin levels (<20 mg/dL). Serum levels of copper may be elevated in pregnancy or stress conditions since ceruloplasmin is an acute-phase reactant and 90% of circulating copper is bound to ceruloplasmin.

Toxicity Copper toxicity is usually accidental (Table 96e-2). In severe cases, kidney failure, liver failure, and coma may ensue. In Wilson's disease, mutations in the copper-transporting *ATP7B* gene lead to accumulation of copper in the liver and brain, with low blood levels due to decreased ceruloplasmin (Chap. 429).

SELENIUM



Selenium, in the form of selenocysteine, is a component of the enzyme glutathione peroxidase, which serves to protect proteins, cell membranes, lipids, and nucleic acids from oxidant

molecules. As such, selenium is being actively studied as a chemopreventive agent against certain cancers, such as prostate cancer. Selenocysteine is also found in the deiodinase enzymes, which mediate the deiodination of thyroxine to triiodothyronine (Chap. 405). Rich dietary sources of selenium include seafood, muscle meat, and cereals, although the selenium content of cereal is determined by the soil concentration. Countries with low soil concentrations include parts of Scandinavia, China, and New Zealand. *Keshan disease* is an endemic cardiomyopathy found in children and young women residing in regions of China where dietary intake of selenium is low (<20 µg/d). Concomitant deficiencies of iodine and selenium may worsen the clinical manifestations of cretinism. Chronic ingestion of large amounts of selenium leads to selenosis, characterized by hair and nail brittleness and loss, garlic breath odor, skin rash, myopathy, irritability, and other abnormalities of the nervous system.

CHROMIUM

Chromium potentiates the action of insulin in patients with impaired glucose tolerance, presumably by increasing insulin receptor-mediated signaling, although its usefulness in treating type 2 diabetes is uncertain. In addition, improvement in blood lipid profiles has been reported in some patients. The usefulness of chromium supplements in muscle building has not been substantiated. Rich food sources of chromium include yeast, meat, and grain products. Chromium in the trivalent state is found in supplements and is largely nontoxic; however, chromium-6 is a product of stainless steel welding and is a known pulmonary carcinogen as well as a cause of liver, kidney, and CNS damage.

MAGNESIUM

See Chap. 423.

FLUORIDE, MANGANESE, AND ULTRATRACE ELEMENTS

An essential function for fluoride in humans has not been described, although it is useful for the maintenance of structure in teeth and bones. Adult fluorosis results in mottled and pitted defects in tooth enamel as well as brittle bone (skeletal fluorosis).

Manganese and molybdenum deficiencies have been reported in patients with rare genetic abnormalities and in a few patients receiving prolonged total parenteral nutrition. Several manganese-specific enzymes have been identified (e.g., manganese superoxide dismutase). Deficiencies of manganese have been reported to result in bone demineralization, poor growth, ataxia, disturbances in carbohydrate and lipid metabolism, and convulsions.

Ultratrace elements are defined as those needed in amounts <1 mg/d. Essentiality has not been established for most ultratrace elements, although selenium, chromium, and iodine are clearly essential (Chap. 405). Molybdenum is necessary for the activity of sulfite and xanthine oxidase, and molybdenum deficiency may result in skeletal and brain lesions.

This page intentionally left blank

This page intentionally left blank

This page intentionally left blank

97 Malnutrition and Nutritional Assessment

Douglas C. Heimburger

Malnutrition can arise from primary or secondary causes, resulting in the former case from inadequate or poor-quality food intake and in the latter case from diseases that alter food intake or nutrient requirements, metabolism, or absorption. *Primary* malnutrition occurs mainly in developing countries and under conditions of political unrest, war, or famine. *Secondary* malnutrition, the main form encountered in industrialized countries, was largely unrecognized until the early 1970s, when it was appreciated that persons with adequate food supplies can become malnourished as a result of acute or chronic diseases that alter nutrient intake or metabolism, particularly diseases that cause acute or chronic inflammation. Various studies have shown that protein-energy malnutrition (PEM) affects one-third to one-half of patients on general medical and surgical wards in teaching hospitals. The consistent finding that nutritional status influences patient prognosis underscores the importance of preventing, detecting, and treating malnutrition.

Definitions for forms of PEM are in flux. Traditionally, the two major types of PEM have been *marasmus* and *kwashiorkor*. These conditions are compared in [Table 97-1](#). Marasmus is the end result of a long-term deficit of dietary energy, whereas kwashiorkor has been understood to result from a protein-poor diet. Although the former concept remains essentially correct, evidence is accumulating that PEM syndromes are distinguished by two main features: insufficient dietary intake and underlying inflammatory processes. Energy-poor diets with minimal inflammation cause gradual erosion of body mass, resulting in classic marasmus. By contrast, inflammation from acute illnesses such as injury or sepsis or from chronic illnesses such as cancer, lung or heart disease, or HIV infection can erode lean body mass even in the presence of relatively sufficient dietary intake, leading to a kwashiorkor-like state. Quite often, inflammatory illnesses impair appetite and dietary intake, producing combinations of the two conditions.

Consensus committees have proposed the following revised definitions. *Starvation-related malnutrition* is suggested for instances of chronic starvation without inflammation, *chronic disease-related malnutrition* when inflammation is chronic and of mild to moderate degree, and *acute disease- or injury-related malnutrition* when inflammation is acute and of a severe degree. However, because distinguishing diagnostic criteria for these conditions have not been universally adopted, this chapter integrates the older and newer terms.

MARASMUS (STARVATION-RELATED MALNUTRITION) AND CACHEXIA (CHRONIC DISEASE-RELATED MALNUTRITION)

Marasmus (starvation-related malnutrition) is a state in which virtually all available body fat stores have been exhausted due to starvation without systemic inflammation. *Cachexia* (chronic disease-related malnutrition) is a state that involves substantial loss of lean body mass in the presence of chronic systemic inflammation. Conditions that produce cachexia tend to be chronic and indolent, such as cancer and chronic pulmonary disease, whereas, in high-income countries, the classic setting for marasmus is in patients with anorexia nervosa. These conditions are relatively easy to detect because of the patient's starved

TABLE 97-1 COMPARISON OF MARASMUS/CACHEXIA AND KWASHIORKOR/ACUTE MALNUTRITION

Feature	Marasmus (Starvation-Related Malnutrition) and Cachexia (Chronic Disease-Related Malnutrition)	Kwashiorkor (Acute Disease- or Injury-Related Malnutrition) ^a
Clinical setting	Prolonged ↓ energy and protein intake with or without systemic inflammation	Acute ↓ energy and protein intake with substantial systemic inflammation
Time course to develop	Months or years	Weeks
Clinical features	Starved appearance (body mass index <18.5) Reduced triceps skinfold Reduced midarm muscle circumference	Normal body mass index (although loss of body mass may be masked by edema) Easy hair pluckability ^b Edema
Laboratory findings	Serum albumin normal (marasmus, no inflammation) or decreased (cachexia, with inflammation)	Serum albumin <2.8 g/dL Total iron-binding capacity <200 μg/dL Lymphocytes <1500/μL Anergy
Clinical course	Reasonably preserved responsiveness to short-term stress	Infections Poor wound healing, decubitus ulcers, skin breakdown
Mortality risk	Low unless related to underlying disease	High
Diagnostic criteria	Triceps skinfold <3 mm Midarm muscle circumference <15 cm	Serum albumin <2.8 g/dL At least one of the following: Poor wound healing, decubitus ulcers, or skin breakdown Easy hair pluckability ^b Edema

^aThe findings used to diagnose kwashiorkor/acute malnutrition must be unexplained by other causes. ^bTested by *firmly* pulling a lock of hair from the top (not the sides or back), grasping with the thumb and forefinger. An average of three or more hairs removed easily and painlessly is considered abnormal hair pluckability.

appearance. The diagnosis is based on fat and muscle wastage resulting from prolonged calorie deficiency and/or inflammation. Diminished skinfold thickness reflects the loss of fat reserves; reduced arm muscle circumference with temporal and interosseous muscle wasting reflects the catabolism of protein throughout the body, including in vital organs such as the heart, liver, and kidneys.

Routine laboratory findings in cachexia/marasmus are relatively unremarkable. The creatinine-height index (24-h urinary creatinine excretion compared with normal values based on height) is low, reflecting the loss of muscle mass. Occasionally, the serum albumin level is reduced, but it remains above 2.8 g/dL when systemic inflammation is absent. Despite a morbid appearance, immunocompetence, wound healing, and the ability to handle short-term stress are reasonably well preserved in most patients.

Pure starvation-related malnutrition is a chronic, fairly well adapted form of starvation rather than an acute illness; it should be treated cautiously in an attempt to reverse the downward trend gradually. Although nutritional support is necessary, overly aggressive repletion can result in severe, even life-threatening metabolic imbalances such as hypophosphatemia and cardiorespiratory failure (*refeeding syndrome*). When possible, oral or enteral nutritional support is preferred; treatment started slowly allows readaptation of metabolic and intestinal functions (**Chap. 98e**).

KWASHIORKOR (ACUTE DISEASE- OR INJURY-RELATED MALNUTRITION)

By contrast, *kwashiorkor* (acute disease- or injury-related malnutrition) in developed countries occurs mainly in connection with acute,

life-threatening conditions such as trauma and sepsis. The physiologic stress produced by these illnesses increases protein and energy requirements at a time when intake is often limited. A classic scenario is an acutely stressed patient who receives only 5% dextrose solutions for periods as brief as 2 weeks. Although the etiologic mechanisms are not fully known, the protein-sparing response normally seen in starvation is blocked by the stressed state and by carbohydrate infusion.

In its early stages, the physical findings of kwashiorkor/acute malnutrition are few and subtle. Initially unaffected fat reserves and muscle mass give the deceptive appearance of adequate nutrition. Signs that support the diagnosis include easy hair pluckability, edema, skin breakdown, and poor wound healing. The major sine qua non is severe reduction of levels of serum proteins such as albumin (<2.8 g/dL) and transferrin (<150 mg/dL) or of iron-binding capacity (<200 μg/dL). Cellular immune function is depressed, as reflected by lymphopenia (<1500 lymphocytes/μL in adults and older children) and lack of response to skin test antigens (*anergy*).

The prognosis of adult patients with full-blown kwashiorkor/acute malnutrition is not good even with aggressive nutritional support. Surgical wounds often dehisce (fail to heal), pressure sores develop, gastroparesis and diarrhea can occur with enteral feeding, the risk of gastrointestinal bleeding from stress ulcers is increased, host defenses are compromised, and death from overwhelming infection may occur despite antibiotic therapy. Unlike treatment of marasmus, therapy for kwashiorkor entails aggressive nutritional support to restore better metabolic balance rapidly (**Chap. 98e**).

PHYSIOLOGIC CHARACTERISTICS OF HYPOMETABOLIC AND HYPERMETABOLIC STATES

The metabolic characteristics and nutritional needs of hypermetabolic patients who are stressed from injury, infection, or chronic inflammatory illness differ from those of hypometabolic patients who are unstressed but chronically starved. In both cases, nutritional support is important, but misjudgments in selecting the appropriate approach may have serious adverse consequences.

The *hypometabolic* patient is typified by the relatively less stressed but mildly catabolic and chronically starved individual who, with time, will develop cachexia/marasmus. The *hypermetabolic* patient stressed from injury or infection is catabolic (experiencing rapid breakdown of body mass) and is at high risk for developing acute malnutrition/kwashiorkor if nutritional needs are not met and/or the illness does not resolve quickly. As summarized in **Table 97-2**, the two states are distinguished by differing perturbations of metabolic rate, rates of protein breakdown (*proteolysis*), and rates of gluconeogenesis. These differences are mediated by proinflammatory cytokines and counter-regulatory hormones—tumor necrosis factor, interleukins 1 and 6, C-reactive protein, catecholamines (epinephrine and norepinephrine), glucagon, and cortisol—whose levels are relatively reduced in hypometabolic patients and increased in hypermetabolic patients. Although insulin levels are also elevated in stressed patients, insulin resistance in the target tissues blocks insulin-mediated anabolic effects. Physiologic

TABLE 97-2 PHYSIOLOGIC CHARACTERISTICS OF HYPOMETABOLIC AND HYPERMETABOLIC STATES

Physiologic Characteristics	Hypometabolic, Nonstressed Patient (Risk for Starvation/ Marasmus)	Hypermetabolic, Stressed Patient (Risk for Kwashiorkor/ Acute Malnutrition)
Cytokines, catecholamines, glucagon, cortisol, insulin	↓	↑
Metabolic rate, O ₂ consumption	↓	↑
Proteolysis, gluconeogenesis	↓	↑
Ureagenesis, urea excretion	↓	↑
Fat catabolism, fatty acid utilization	Relative ↑	Absolute ↑
Adaptation to starvation	Normal	Abnormal

characteristics of patients at risk for chronic disease-related malnutrition are less predictable and likely represent a mixture of the two extremes depicted in Table 97-2.

Metabolic Rate In starvation and semistarvation, the resting metabolic rate falls between 10% and 30% as an adaptive response to energy restriction, slowing the rate of weight loss. By contrast, the resting metabolic rate rises in the presence of physiologic stress in proportion to the degree of the insult. The rate may increase by ~10% after elective surgery, 20–30% after bone fractures, 30–60% with severe infections such as peritonitis or gram-negative septicemia, and as much as 110% after major burns.

If the metabolic rate (energy requirement) is not matched by energy intake, weight loss results—slowly in hypometabolism and quickly in hypermetabolism. Losses of up to 10% of body mass are unlikely to be detrimental; however, greater losses in acutely ill hypermetabolic patients may be associated with rapid deterioration in body functions.

Protein Catabolism The rate of endogenous protein breakdown (*catabolism*) to supply energy needs normally falls during uncomplicated energy deprivation. After ~10 days of total starvation, an unstressed individual loses about 12–18 g of protein per day (equivalent to ~60 g of muscle tissue or ~2–3 g of nitrogen). In contrast, in injury and sepsis, protein breakdown accelerates in proportion to the degree of stress, reaching 30–60 g/d after elective surgery, 60–90 g/d with infection, 100–130 g/d with severe sepsis or skeletal trauma, and >175 g/d with major burns or head injuries. These losses are reflected by proportional increases in the excretion of urea nitrogen, the major by-product of protein breakdown.

Gluconeogenesis The major aim of protein catabolism during a state of starvation is to provide the glucogenic amino acids (especially alanine and glutamine) that serve as substrates for endogenous glucose production (*gluconeogenesis*) in the liver. In the hypometabolic/starved state, protein breakdown for gluconeogenesis is minimized, especially as ketones derived from fatty acids become the substrate preferred by certain tissues. In the hypermetabolic/stress state, gluconeogenesis increases dramatically and in proportion to the degree of the insult to increase the supply of glucose (the major fuel of reparation). Glucose is the only fuel that can be utilized by hypoxic tissues (*anaerobic glycolysis*), white blood cells, and newly generated fibroblasts. Infusions of glucose partially offset a negative energy balance but do not significantly suppress the high rates of gluconeogenesis in catabolic patients. Hence, adequate supplies of protein are needed to replace the amino acids used for this metabolic response.

In summary, a hypometabolic patient is adapted to starvation and conserves body mass through reduction of the metabolic rate and use of fat as the primary fuel (rather than glucose and its precursor amino acids). A hypermetabolic patient also uses fat as a fuel but rapidly breaks down body protein to produce glucose, with consequent loss of muscle and organ tissue and danger to vital body functions.

MICRONUTRIENT MALNUTRITION

The same illnesses and reductions in nutrient intake that lead to PEM often produce deficiencies of vitamins and minerals as well (**Chap. 96e**). Deficiencies of nutrients that are stored in small amounts (such as the water-soluble vitamins) occur because of loss through external secretions, such as zinc in diarrhea fluid or burn exudate, and are probably more common than is generally recognized.


Deficiencies of vitamin C, folic acid, and zinc are relatively common in sick patients. Signs of scurvy, such as corkscrew hairs on the lower extremities, are found frequently in chronically ill and/or alcoholic patients. The diagnosis can be confirmed by determination of plasma vitamin C levels. Folic acid intakes and blood levels are often less than optimal, even among healthy persons; with illness, alcoholism, poverty, or poor dentition, these deficiencies are common. Low blood zinc levels are prevalent in patients with malabsorption syndromes such as inflammatory bowel disease. Patients with zinc deficiency often exhibit poor wound healing, pressure ulcer formation, and impaired

immunity. Thiamine deficiency is a common complication of alcoholism but may be prevented by therapeutic doses of thiamine in patients treated for alcohol abuse.

Patients with low plasma vitamin C levels usually respond to the doses in multivitamin preparations, but patients with deficiencies should be supplemented with 250–500 mg/d. Folic acid is absent from some oral multivitamin preparations; patients with deficiencies should be supplemented with ~1 mg/d. Patients with zinc deficiencies resulting from large external losses sometimes require oral supplementation with 220 mg of zinc sulfate one to three times daily. For these reasons, laboratory assessments of the micronutrient status of patients at high risk are desirable.

Hypophosphatemia develops in hospitalized patients with remarkable frequency and generally results from rapid intracellular shifts of phosphate in underweight or alcoholic patients receiving intravenous glucose (**Chap. 63**). The adverse clinical sequelae are numerous; some, such as acute cardiopulmonary failure, are collectively called *refeeding syndrome* and can be life-threatening.

GLOBAL CONSIDERATIONS

 Many developing countries are still faced with high prevalences of the classic forms of PEM: marasmus and kwashiorkor. *Food insecurity*, which characterizes many poor countries, prevents consistent dietary sufficiency and/or quality and leads to endemic or cyclic malnutrition. Factors threatening food security include marked seasonal variations in agricultural productivity (rainy season–dry season cycles), periodic droughts, political unrest or injustice, and disease epidemics (especially of HIV/AIDS). The coexistence of malnutrition and disease epidemics exacerbates the latter and increases complications and mortality rates, creating vicious cycles of malnutrition and disease.

As economic prosperity improves, developing countries have been observed to undergo an epidemiologic transition, a component of which has been termed the *nutrition transition*. As improved economic resources make greater dietary diversity possible, middle-income populations (e.g., in southern Asia, China, and Latin America) typically begin to adopt lifestyle habits of industrialized nations, with increased consumption of energy and fat and decreased levels of physical activity. These changes lead to rising levels of obesity, metabolic syndrome, diabetes, cardiovascular disease, and cancer, sometimes coexisting in populations with persistent undernutrition.

Micronutrient deficiencies also remain prevalent in many countries of the world, impairing functional status and productivity and increasing mortality rates. Vitamin A deficiency impairs vision and increases morbidity and mortality rates from infections such as measles. Mild to moderate iron deficiency may be prevalent in up to 50% of the world, resulting from poor dietary diversity coupled with periodic blood loss and pregnancies. Iodine deficiency remains prevalent, causing goiter, hypothyroidism, and cretinism. Zinc deficiency is endemic in many populations, producing growth retardation, hypogonadism, and dermatoses and impairing wound healing. Fortunately, public health supplementation programs have substantially improved vitamin A and zinc status in developing countries during the past two decades, reducing mortality rates from measles, diarrheal diseases, and other manifestations. However, with the advancing nutrition transition and a shift toward nutritionally related chronic noncommunicable conditions, it is estimated that nutrition remains one of the three greatest contributors of risk for morbidity and mortality worldwide.

NUTRITIONAL ASSESSMENT

Because interactions between illness and nutrition are complex, many physical and laboratory findings reflect both underlying disease and nutritional status. Therefore, the nutritional evaluation of a patient requires an integration of history, physical examination, anthropometrics, and laboratory studies. This approach helps both to detect nutritional problems and to prevent the conclusion that isolated findings indicate nutritional problems when they do not. For example, hypoalbuminemia caused by an inflammatory illness does not necessarily indicate malnutrition.

TABLE 97-3 NUTRITIONAL DEFICIENCY: THE HIGH-RISK PATIENT

Underweight (body mass index <18.5) and/or recent loss of $\geq 10\%$ of usual body mass
Poor intake: anorexia, food avoidance (e.g., psychiatric condition), or NPO ^a status for more than ~ 5 days
Protracted nutrient losses: malabsorption, enteric fistulas, draining abscesses or wounds, renal dialysis
Hypermetabolic states: sepsis, protracted fever, extensive trauma or burns
Alcohol abuse or use of drugs with antinutrient or catabolic properties: glucocorticoids, antimetabolites (e.g., methotrexate), immunosuppressants, antitumor agents
Impoverishment, isolation, advanced age

^aNil per os (nothing by mouth).

Nutritional History Elicitation of a nutritional history is directed toward the identification of underlying mechanisms that put patients at risk for nutritional depletion or excess. These mechanisms include inadequate intake, impaired absorption, decreased utilization, increased losses, and increased requirements for nutrients.

Individuals with the characteristics listed in **Table 97-3** are at particular risk for nutritional deficiencies.

Physical Examination Physical findings that suggest vitamin, mineral, and protein-energy deficiencies and excesses are outlined in **Table 97-4**. Most of the physical findings are not specific for individual nutrient deficiencies and must be integrated with historic, anthropometric, and laboratory findings. For example, follicular hyperkeratosis on the back of the arms is a fairly common, normal finding. However, if it is widespread in a person who consumes few fruits and vegetables and smokes regularly (increasing ascorbic acid requirements), vitamin C deficiency is likely. Similarly, easily pluckable hair may be a consequence of chemotherapy but suggests acute malnutrition/kwashiorkor in a hospitalized patient who has poorly healing surgical wounds and hypoalbuminemia.

Anthropometric Measurements Anthropometric measurements provide information on body muscle mass and fat reserves. The most practical and commonly used measurements are body weight, height, triceps skinfold (TSF), and midarm muscle circumference (MAMC). Body weight is one of the most useful nutritional parameters to follow in patients who are acutely or chronically ill. Unintentional weight loss during illness often reflects loss of lean body mass (muscle and organ tissue), especially if it is rapid and is not caused by diuresis. Such weight loss can be an ominous sign since it indicates use of vital body protein stores for metabolic fuel. The reference standard for normal body weight, body mass index (BMI: weight in kilograms divided by height, in meters, squared), is discussed in **Chap. 416**. BMI values <18.5 are considered underweight; <17, significantly underweight; and <16, severely wasted. Values of 18.5–24.9 are normal; 25–29.9, overweight; and ≥ 30 , obese.

Measurement of skinfold thickness is useful for estimating body fat stores, because $\sim 50\%$ of body fat is normally located in the subcutaneous region. This measurement can also permit discrimination of fat mass from muscle mass. The triceps is a convenient site that is generally representative of the body's overall fat level. A thickness <3 mm suggests virtually complete exhaustion of fat stores. The MAMC can be used to estimate skeletal muscle mass, calculated as follows:

$$\text{MAMC (cm)} = \text{upper arm circumference (cm)} - [0.314 \times \text{TSF (mm)}]$$

Laboratory Studies A number of laboratory tests used routinely in clinical medicine can yield valuable information about a patient's nutritional status if a slightly different approach to their interpretation is used. For example, abnormally low serum albumin levels, low total iron-binding capacity, and anergy may have a distinct explanation, but collectively they may represent kwashiorkor. In the clinical setting of a hypermetabolic, acutely ill patient who is edematous and has easily pluckable hair and inadequate protein intake, the diagnosis of acute malnutrition/kwashiorkor is clear-cut. Commonly used laboratory

tests for assessing nutritional status are outlined in **Table 97-5**. The table also provides tips to avoid the assignment of nutritional significance to tests that may be abnormal for nonnutritional reasons.

ASSESSMENT OF CIRCULATING (VISCERAL) PROTEINS The serum proteins most commonly used to assess nutritional status include albumin, total iron-binding capacity (or transferrin), thyroxine-binding prealbumin (or transthyretin), and retinol-binding protein. Because they have different synthesis rates and half-lives (the half-life of serum albumin is ~ 21 days, whereas those of prealbumin and retinol-binding protein are ~ 2 days and ~ 12 h, respectively), some of these proteins reflect changes in nutritional status more quickly than do others. However, rapid fluctuations can also make shorter-half-life proteins less reliable.

Levels of circulating proteins are influenced by their rates of synthesis and catabolism, "third spacing" (loss into interstitial spaces), and, in some cases, external loss. Although an adequate intake of calories and protein is necessary for optimal circulating protein levels, serum protein levels generally do not reflect protein intake. For example, a drop in the serum level of albumin or transferrin often accompanies significant physiologic stress (e.g., from infection or injury) and is not necessarily an indication of malnutrition or poor intake. A low serum albumin level in a burned patient with both hypermetabolism and increased dermal losses of protein may not indicate malnutrition. However, adequate nutritional support of the patient's calorie and protein needs is critical for returning circulating proteins to normal levels as stress resolves. Thus low values by themselves do not define malnutrition, but they often point to increased risk of malnutrition because of the hypermetabolic stress state. As long as significant physiologic stress persists, serum protein levels remain low, even with aggressive nutritional support. However, if the levels do not rise after the underlying illness improves, the patient's protein and calorie needs should be reassessed to ensure that intake is sufficient.

ASSESSMENT OF VITAMIN AND MINERAL STATUS The use of laboratory tests to confirm suspected micronutrient deficiencies is desirable because the physical findings for those deficiencies are often equivocal or nonspecific. Low blood micronutrient levels can predate more serious clinical manifestations and also may indicate drug-nutrient interactions.

ESTIMATING PROTEIN AND ENERGY REQUIREMENTS

A patient's basal energy expenditure (BEE, measured in kilocalories per day) can be estimated from height, weight, age, and sex with the Harris-Benedict equations:

$$\begin{aligned} \text{Men: BEE} &= 66.47 + 13.75W + 5.00H - 6.76A \\ \text{Women: BEE} &= 655.10 + 9.56W + 1.85H - 4.68A \end{aligned}$$

In these equations, W is weight in kilograms, H is height in centimeters, and A is age in years. After these equations are solved, total energy requirements are estimated by multiplying BEE by a factor that accounts for the stress of illness. Multiplying by 1.1–1.4 yields a range 10–40% above basal that estimates the 24-h energy expenditure of the majority of patients. The lower value (1.1) is used for patients without evidence of significant physiologic stress; the higher value (1.4) is appropriate for patients with marked stress such as sepsis or trauma. The result is used as a 24-h energy goal for feeding.

When it is important to have a more accurate assessment, energy expenditure can be measured at the bedside by indirect calorimetry. This technique is useful in patients who are thought to be hypermetabolic from sepsis or trauma and whose body weight cannot be ascertained accurately. Indirect calorimetry can also be useful in patients who have difficulty weaning from a ventilator and whose energy needs therefore should not be exceeded to avoid excessive CO_2 production. Patients at the extremes of weight (e.g., obese persons) and/or age are good candidates as well, because the Harris-Benedict equations were developed from measurements in adults with roughly normal body weights.

Because urea is a major by-product of protein catabolism, the amount of urea nitrogen excreted each day can be used to estimate the rate of protein catabolism and determine whether protein intake is adequate to

TABLE 97-4 PHYSICAL FINDINGS OF NUTRITIONAL DEFICIENCIES

Clinical Findings	Possible Deficiency or Form of Malnutrition	Possible Excess
Hair, nails		
Corkscrew hairs and unemerged coiled hairs	Vitamin C	
Easily pluckable hair	Acute malnutrition	
Flag sign (transverse depigmentation of hair)	Acute malnutrition	
Sparse hair	Biotin, zinc; acute malnutrition	Vitamin A
Transverse ridging of nails	Acute malnutrition	
Skin		
Cellophane appearance	Acute malnutrition	
Cracking ("flaky-paint" or "crazy-pavement" dermatosis)	Acute malnutrition	
Follicular hyperkeratosis	Vitamins A, C	
Petechiae (especially perifollicular)	Vitamin C	
Purpura	Vitamins C, K	
Pigmentation, scaling of sun-exposed areas	Niacin	
Poor wound healing, decubitus ulcers	Vitamin C, zinc; acute malnutrition	
Scaling	Vitamin A, essential fatty acids, biotin Zinc (hyperpigmented)	Vitamin A
Yellow pigmentation sparing sclerae (benign)		Carotene
Eyes		
Night blindness	Vitamin A	
Papilledema		Vitamin A
Perioral		
Angular stomatitis	Riboflavin, pyridoxine, niacin	
Cheilosis (dry, cracking, ulcerated lips)	Riboflavin, pyridoxine, niacin	
Oral		
Atrophic lingual papillae (slick tongue)	Riboflavin, niacin, folate, vitamin B ₁₂ , iron; acute malnutrition	
Glossitis (scarlet, raw tongue)	Riboflavin, niacin, pyridoxine, folate, vitamin B ₁₂	
Hypogeusesthesia, hyposmia	Zinc	
Swollen, retracted, bleeding gums (if teeth present)	Vitamin C	
Bones, joints		
Beading of ribs, epiphyseal swelling, bowlegs	Vitamin D	
Tenderness, subperiosteal hemorrhage in children	Vitamin C	
Neurologic		
Confabulation, disorientation	Thiamine (Korsakoff's psychosis)	
Drowsiness, lethargy, vomiting		Vitamin A
Dementia	Niacin, vitamin B ₁₂ , folate	
Headache		Vitamin A
Ophthalmoplegia	Thiamine, phosphorus	
Peripheral neuropathy (e.g., weakness, paresthesias, ataxia, footdrop, and decreased tendon reflexes, fine tactile sense, vibratory sense, and position sense)	Thiamine, pyridoxine, vitamin B ₁₂	Pyridoxine
Tetany	Calcium, magnesium	
Other		
Edema	Thiamine; acute malnutrition	
Heart failure	Thiamine ("wet" beriberi), phosphorus	
Hepatomegaly	Acute malnutrition	Vitamin A
Parotid enlargement	Acute malnutrition (consider also bulimia)	
Sudden heart failure, death	Vitamin C	

offset it. Total protein loss and protein balance can be calculated from urinary urea nitrogen (UUN) as follows:

$$\text{Protein catabolic rate (g/d)} = [24\text{-h UUN (g)} + 4] \times 6.25 \text{ (g protein/g nitrogen)}$$

The value of 4 g added to the UUN represents a liberal estimate of the unmeasured nitrogen lost in the urine (e.g., creatinine and uric acid), sweat, hair, skin, and feces. When protein intake is low (e.g., less than

~20 g/d), the equation indicates both the patient's protein requirement and the severity of the catabolic state (Table 97-5). More substantial protein intakes can raise the UUN because some of the ingested (or intravenously infused) protein is catabolized and converted to UUN. Thus, at lower protein intakes, the equation is useful for estimating requirements, and at higher protein intakes it is useful for assessing protein balance.

$$\text{Protein balance (g/d)} = \text{protein intake} - \text{protein catabolic rate}$$

TABLE 97-5 LABORATORY TESTS FOR NUTRITIONAL ASSESSMENT

Test (Normal Values)	Nutritional Use	Causes of Normal Value Despite Malnutrition	Other Causes of Abnormal Value
Serum albumin (3.5–5.5 g/dL)	2.8–3.5 g/dL: Protein depletion or systemic inflammation <2.8 g/dL: Possible acute malnutrition or severe inflammation Increasing value reflects positive protein balance	Dehydration Infusion of albumin, fresh-frozen plasma, or whole blood	Low <i>Common:</i> Infection and other stress, especially with poor protein intake Burns, trauma Congestive heart failure Fluid overload Severe liver disease <i>Uncommon:</i> Nephrotic syndrome Zinc deficiency Bacterial stasis/overgrowth of small intestine
Serum prealbumin, also called transthyretin (20–40 mg/dL; lower in prepubertal children)	10–15 mg/dL: Mild protein depletion or inflammation 5–10 mg/dL: Moderate protein depletion or inflammation <5 mg/dL: Severe protein depletion or inflammation Increasing value reflects positive protein balance	Chronic renal failure	Similar to serum albumin
Serum total iron-binding capacity (240–450 µg/dL)	<200 µg/dL: Protein depletion or inflammatory state; possible acute malnutrition Increasing value reflects positive protein balance More labile than albumin	Iron deficiency	Low Similar to serum albumin High Iron deficiency
Prothrombin time (2.0–15.5 s)	Prolongation: vitamin K deficiency		Prolonged Anticoagulant therapy (warfarin) Severe liver disease
Serum creatinine (0.6–1.6 mg/dL)	<0.6 mg/dL: Muscle wasting due to prolonged energy deficit Reflects muscle mass		High Despite muscle wasting: Renal failure Severe dehydration
24-h urinary creatinine (500–1200 mg/d, standardized for height and sex)	Low value: muscle wasting due to prolonged energy deficit	>24-h collection Decreasing serum creatinine	Low Incomplete urine collection Increasing serum creatinine Neuromuscular wasting
24-h urinary urea nitrogen (UUN; <5 g/d; depends on level of protein intake)	<i>Determine level of catabolism</i> (as long as protein intake is ≥10 g below calculated protein loss or <20 g total, and as long as carbohydrate intake has been at least 100 g) 5–10 g/d: Mild catabolism or normal fed state 10–15 g/d: Moderate catabolism >15 g/d: Severe catabolism <i>Estimate protein balance</i> Protein balance = protein intake – protein loss where protein loss (protein catabolic rate) = [24-h UUN (g) + 4] × 6.25. Adjustments required in burn patients and others with large nonurinary nitrogen losses and in patients with fluctuating levels of blood urea nitrogen (e.g., in renal failure)		
Blood urea nitrogen (8–23 mg/dL)	<8 mg/dL: Possibly inadequate protein intake 12–23 mg/dL: Possibly adequate protein intake >23 mg/dL: Possibly excessive protein intake If serum creatinine is normal, use BUN. If serum creatinine is elevated, use BUN/creatinine ratio. (Normal range is essentially the same as for BUN.)		Low Severe liver disease Anabolic state Syndrome of inappropriate antidiuretic hormone High Despite poor protein intake: Renal failure (Use BUN/creatinine ratio.) Congestive heart failure Gastrointestinal hemorrhage

When correctly implemented, specialized nutritional support (SNS) plays a major and often life-saving role in medicine. SNS is used for two main purposes: (1) to provide an appropriate nutritional substrate in order to maintain or replenish the nutritional status of patients unable to voluntarily ingest or absorb sufficient amounts of food, and (2) to maintain the nutritional and metabolic status of adequately nourished patients who are experiencing systemic hypercatabolic effects of severe inflammation, injury, or infection in the course of persistent critical illness. Patients with permanent major loss of intestinal length or function often require lifelong SNS. Many patients who require treatment in chronic-care facilities receive enteral SNS, most often because their voluntary food intake is deemed insufficient or because impaired chewing and swallowing create a high risk of aspiration pneumonia.

Enteral SNS is the provision of liquid formula meals through a tube placed into the gut. Parenteral SNS is the direct infusion of complete mixtures of crystalline amino acids, dextrose, triglyceride emulsions, and micronutrients into the bloodstream through a central venous catheter or (rarely in adults) via a peripheral vein. The enteral route is almost always preferred because of its relative simplicity and safety, its low cost, and the benefits of maintaining digestive, absorptive, and immunologic barrier functions of the gastrointestinal tract. Pliable, small-bore feeding tubes make placement relatively easy and acceptable to patients. Constant-rate infusion pumps increase the reliability of nutrient delivery. The chief disadvantage of enteral SNS is that many days may be required to meet the patient's nutrient requirements.

For short-term use, the feeding tube can be placed via the nose into the stomach, duodenum, or jejunum. For long-term use, these sites may be accessed through the abdominal wall by endoscopic or surgical procedures. The chief disadvantage of tube feeding in acute illness is intolerance due to gastric retention, risk of vomiting, or diarrhea. The presence of severe coagulopathy is a relative contraindication to the insertion of a feeding tube. In adults, parenteral nutrition (PN) almost always requires aseptic insertion of a central venous catheter with a dedicated port. Many circumstances can delay or slow the progression of enteral SNS, whereas parenteral SNS can provide a complete substrate mix easily and promptly. This practical advantage is mitigated by the need to infuse relatively large fluid volumes and the real risk of inadvertent toxic overfeeding.

APPROACH TO THE PATIENT:

Requirements for Specialized Nutritional Support

INDICATIONS FOR SPECIALIZED NUTRITIONAL SUPPORT

Approximately one-fifth to one-quarter of patients in acute-care hospitals suffer from at least moderate protein-energy malnutrition (PEM), the defining features of which are malnutrition-induced weight loss and skeletal muscle atrophy. Usually, but not always, other features further compromise clinical responses; these features include a subnormal adipose tissue mass, with the accompanying adverse consequences of weakness, skin thinning, and breakdown; reduced ventilatory drive; ineffective cough; immunodeficiency; and impaired thermoregulation. Commonly, PEM is already present at the time of hospital admission and remains unimproved or worsens during the ensuing hospital stay. Common reasons for PEM worsening during hospitalization are refusal of food (because of anorexia, nausea, pain, or delirium), communication barriers, an unmet need for hand-feeding of patients with physical or sensory impairment, disordered or ineffective chewing or swallowing, and prolonged periods of physician-ordered fasting—all potentially

TABLE 98e-1 BODY MASS INDEX (BMI), MUSCLE MASS, AND PROTEIN ENERGY MALNUTRITION (PEM)

BMI (kg/m ²)	Skeletal Muscle Mass	Nutritional Status
>30	Normal	Obese
25–29.9	Normal	Overweight
20–24.9	Normal	Normal
>18.5	Decreased	PEM despite adequate or excessive adipose tissue store
>18.5	Decreased	Moderate PEM
<16	Decreased	Severe PEM
<13	Decreased	Lethal in men
<11	Decreased	Lethal in women

taking place in a context of caregiver unawareness and inattention. Most patients who are suffering from in-hospital PEM do not, or ought not, to require SNS. A large proportion of these patients can be expected to improve with appropriate management of their primary disease. Others have a terminal disease whose downward course will not be altered by SNS. In yet other cases, the PEM is sufficiently mild that the benefits of SNS are exceeded by its risks. For patients who fall into this last category, the correct approach is to intensify and/or modify the patient's oral nutrition as directed by the unit dietitian.

PEM is often classified as minimal, moderate, or severe on the basis of weight for height (body mass index, BMI) and percentage of body weight recently lost. As shown in [Table 98e-1](#), the BMI (when corrected for abnormal extracellular fluid accumulation) is a crude but useful indicator of PEM severity. Note, however, that obesity does not preclude moderate or severe PEM, especially in older or bedridden patients; indeed, obesity can mask the presence of PEM if the patient's muscle mass is not specifically examined.

The decision to implement SNS must be based on the determinations (1) that intensified or modified oral nutrition has failed or is impossible, impractical, or undesirable; and (2) that SNS will increase the patient's rate and likelihood of recovery, reduce the risk of infection, improve healing, or otherwise shorten the hospital stay. In chronic-care situations, the decision to institute SNS is based on the likelihood that the intervention will extend the duration or quality of the patient's life. An algorithm for determining when to use SNS is depicted in [Fig. 98e-1](#).

The decision to enhance oral nutrition or—that attempt failing—to resort to SNS is based on the anticipated consequences of non-intervention. The mnemonic “in-in-in” (for inanition-inflammation-inactivity) can serve as a reminder of the three main factors that come into play when deciding whether or not it is acceptable to withhold SNS from a patient with PEM.

Inanition Key issues include whether normal food intake is likely to be impossible for a prolonged period and whether the patient can tolerate prolonged starvation. A previously well-nourished person can tolerate ~7 days of starvation without harm, even in the presence of a moderate systemic response to inflammation (SRI), whereas the degree of tolerance to prolonged starvation is much less in patients whose skeletal muscle mass is already reduced, whether from PEM, from the muscle atrophy of old age (sarcopenia), or from muscle atrophy due to neuromuscular disease. Excess body fat does not exclude the possibility of coexisting muscle atrophy from any of these causes. In general, unintentional weight loss of >10% during the previous 6 months or a weight-to-height ratio that is <90% of standard, when associated with physiologic impairment, crudely predicts that the patient has moderate PEM. Weight loss >20% of usual or <80% of standard makes severe PEM more likely.

Inflammation The anorexia that invariably accompanies the SRI reduces the likelihood that a patient's nutritional goals will be achieved by intensifying or modifying the diet, by providing counseling, or by hand-feeding. Furthermore, the protein-catabolic

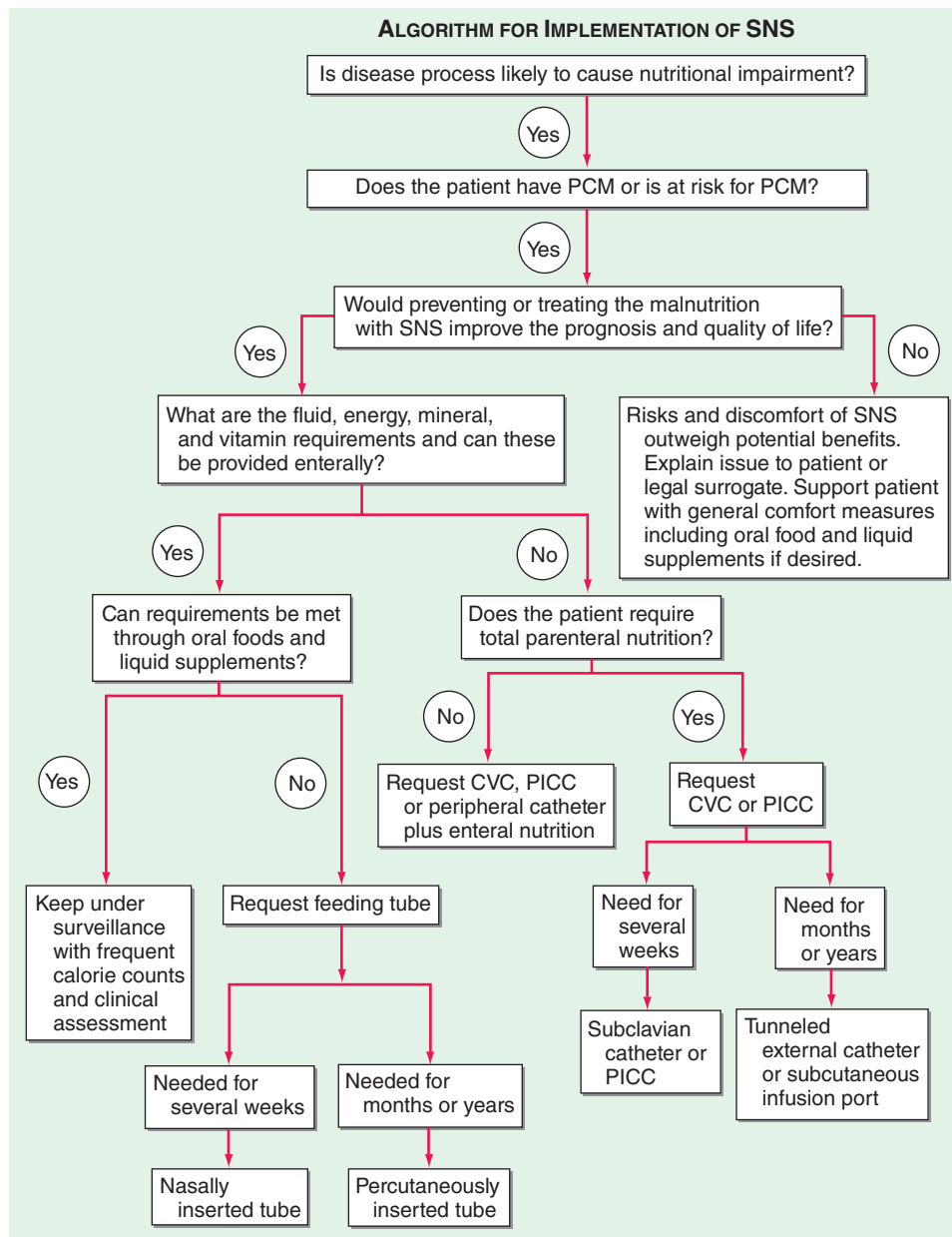


FIGURE 98e-1 Decision-making for the implementation of specialized nutritional support (SNS). CVC, central venous catheter; PICC, peripherally inserted central catheter. (Adapted from the chapter on this topic in *Harrison's Principles of Internal Medicine, 16e*, by Lyn Howard, MD.)

effects of the SRI accelerate skeletal muscle wasting and substantially block normal protein-sparing adaptation to protein and energy starvation.

Inactivity A nutritional red flag should be raised over every acutely ill patient who remains bedridden or inactive for a prolonged period. Such patients commonly manifest muscle atrophy (due to nutritional deficiencies and disuse) and anorexia with inadequate voluntary food intake.

Once it has been determined that a patient has significant—and, in particular, *progressive*—PEM despite meaningful efforts to reverse it by modifying the diet or the way food is provided, the next step is to decide whether SNS will have a net positive effect on the patient's clinical outcome. The pathway to the end stage of most severe chronic diseases leads through PEM. In most patients with end-stage untreatable cancer or certain end-organ diseases, SNS will neither reverse PEM nor improve the quality of life. Provision of food and water is commonly regarded as an aspect of basic humane care; in contrast, enteral and parenteral SNS is a therapeutic intervention that can cause discomfort and pose risks. As with other

life-support interventions, the discontinuation of enteral or parenteral SNS can be psychologically difficult for patients, their families, and their caregivers. Indeed, the difficulty can be greater with SNS than with other life-support interventions because the provision of food and water is often considered equivalent to comfort care. In such difficult, near end-of-life situations, it is prudent to explicitly state the treatment goals at the outset of a course of SNS therapy. Such clarity can smooth the way for subsequent appropriate discontinuation in those patients whose prognosis has become hopeless.

After the decision has been made that SNS is indeed appropriate, the next determinations are the route of delivery (enteral versus parenteral), timing, and calculation of the patient's nutritional goals. Although enteral SNS is the default option, the choice of optimal route depends on the degree of gut function as well as on available technical resources.

Both the choice of route and the timing of SNS require an evaluation of the patient's current nutritional status, the presence and extent of the SRI, and the anticipated clinical course. Severe SRI is identified on the basis of the standard clinical signs of leukocytosis, tachycardia, tachypnea, and temperature elevation or depression.

Serum albumin is a negative acute-phase protein and hence a marker of the SRI. More severe hypoalbuminemia is a crude indicator of greater SRI severity, but this condition is almost certainly worsened by concurrent dietary protein deficiency. Despite the importance of adequate protein provision to patients with the SRI, no amount of SNS will raise serum albumin levels into the normal range as long as the SRI persists.

The SRI can be graded as mild, moderate, or severe. Examples of a severe SRI include (1) sepsis or other major inflammatory diseases (e.g., pancreatitis) that require care in the intensive care unit; (2) multiple trauma with an Injury Severity Score >20–25 or an Acute Physiology and Chronic Health Evaluation II (APACHE II) score >25; (3) closed head injury with a Glasgow Coma Scale <8; and (4) major third-degree burns of >40% of the body surface area. A moderate SRI occurs with less severe infections, injuries, or inflammatory conditions like pneumonia, uncomplicated major surgery, acute hepatic or renal injury, and exacerbations of ulcerative colitis or regional enteritis requiring hospitalization.

Patients with a severe SRI require the initiation of SNS within the first several days of care, for they are highly unlikely to consume an adequate amount of food voluntarily over the next 7 days. On the other hand, a moderate SRI, as is common during the period following major uncomplicated surgery without oral intake, may be tolerated for 5–7 days as long as the patient is initially well nourished. Patients awaiting elective major surgery benefit from preoperative nutritional repletion for 5–10 days but only in the presence of significant PEM. When adequate preoperative nutrition or SNS is impractical, early postoperative SNS is usually indicated. Furthermore, patients with a combination of a moderate SRI and moderate PEM are likely to benefit from early postoperative SNS.

RISKS AND BENEFITS OF SPECIALIZED NUTRITIONAL SUPPORT

The risks of enteral SNS are determined primarily by the patient's state of alertness and swallowing competence, the anatomy and function of the gastrointestinal tract, and the experience of the supervising clinical team. The safest and least costly approach is to avoid SNS by close attention to oral food intake; personal encouragement; dietary modifications; hand-feeding, when possible; and, often, the addition of an oral liquid supplement. For this reason, all patients at nutritional risk should be assessed and followed by a nutritionist. There is increasing interest in the use, under selected circumstances and when not contraindicated, of pharmacologic doses of anabolic steroids to stimulate appetite and promote muscle anabolism.

Nasogastric tube insertion is a bedside procedure, but many critically ill patients have impaired gastric emptying and a high risk of aspiration pneumonia. This risk can be reduced by placing the tip of the feeding tube in the jejunum beyond the ligament of Treitz, a procedure that usually requires fluoroscopic or endoscopic guidance. When a laparotomy is planned for a patient who has other surgical conditions likely to necessitate prolonged SNS, it is advantageous to place a jejunal feeding tube at the time of surgery.

A major disadvantage of enteral SNS is that the amounts of protein and calories provided to critically ill patients commonly fail to reach target goals within the first 7–14 days after SNS is initiated. This problem is compounded by the lack of enteral products that allow the provision of the recommended protein target of 1.5–2.0 g/kg without simultaneously inducing potentially harmful caloric overfeeding.

Enteral SNS is often required in patients with anorexia, impaired swallowing, or small-intestinal disease. The bowel and its associated digestive organs derive 70% of their required nutrients directly from nutritional substrates absorbed from the intestinal lumen. Enteral feeding also supports gut function by stimulating splanchnic blood flow, neuronal activity, IgA antibody release, and secretion of gastrointestinal hormones that stimulate gut trophic activity. These factors support the gut as an immunologic barrier against enteric pathogens. For these reasons, current evidence indicates that some luminal nutrition should be provided, even when PN is required to

provide most of the nutritional support. The nonessential amino acids arginine and glutamine, short-chain fatty acids, long-chain omega 3 fatty acids, and nucleotides are available in some specialty enteral formulas and appear to have an important role in maintaining immune function. The addition of supplemental PN to enteral feeding (either by mouth or as SNS by enteral tube) may hasten the transition to full enteral feeding, which is usually successful when >50% of requirements can be met enterally. As long as protein and other essential nutrient requirements are met, substantial nutritional benefit can be achieved by providing ~50% of energy needs for periods of up to 10 days. As a rule of thumb, dietary protein provision should be increased by ~25–50% when energy intake is reduced by this amount, since negative energy balance reduces the efficiency of dietary protein retention. For longer periods and in patients who have a normal or increased body fat content, it may be preferable to provide only 75–80% of energy needs (together with increased protein), as the mild energy deficit improves gastrointestinal tolerance, makes glycemic control far easier, and avoids excess fluid administration.

The main risks of PN are related to the placement of a central venous catheter, with its complications of thrombosis and infection, and the relatively large intravenous volumes infused. Less often appreciated are the risks associated with the ease of inadvertently infusing excessive carbohydrate and lipid directly into the bloodstream. These risks include hyperglycemia, inadequate lipid clearance from the circulation, hepatic steatosis and inflammation, and even respiratory failure in patients with borderline pulmonary function. On the other hand, renal dysfunction does not reduce a patient's requirement for protein or amino acids. In cases in which renal function is a limiting factor, appropriate renal replacement therapy must be provided along with SNS.

In the past, bowel rest through PN was the cornerstone of treatment for many severe gastrointestinal disorders. However, the value of providing even minimal amounts of enteral nutrition (EN) is now widely accepted. Protocols to facilitate more widespread use of EN include initiation within 24 h of ICU admission; aggressive use of the head-upright position; use of postpyloric and nasojejunal feeding tubes; use of prokinetic agents; more rapid increases in feeding rates; tolerance of higher gastric residuals; and adherence to nurse-directed algorithms for feeding progression. Parenteral SNS alone is generally necessary only for severe gut dysfunction due to prolonged ileus, intestinal obstruction, or severe hemorrhagic pancreatitis.

In critically ill patients, parenteral SNS can be commenced within the first 24 h of care, with the anticipation of a better clinical outcome and a lower mortality risk than those following delayed or inadequate enteral SNS; however, this point remains controversial. Some evidence suggests that early SNS is associated with a reduced risk of death but also with an increased risk of serious infection. More recent data, obtained in studies of moderately critically ill patients, suggest that early hypocaloric parenteral SNS lessens morbidity and mitigates muscle atrophy without an increased risk of infection, but also without a detectable reduction in mortality risk. Unfortunately, the current clinical-trial evidence fails to address several important unknowns. It is important to note that the level of protein substrate provided in published clinical trials generally falls well below the current recommendation, even in trials of supplemental parenteral SNS. Much of the increase in morbidity associated with parenteral and enteral SNS can be ascribed to hyperglycemia, which can be prevented by appropriately intensive insulin therapy. The level of glycemia necessary to prevent complications, whether <110 mg/dL or <150 mg/dL, remains unclear. Adequately fed surgical patients may benefit from the lower glucose range, but studies of intensive insulin therapy alone, without full feeding, suggest improved morbidity and mortality outcomes with looser control of glucose at <180 mg/dL.

In the early years of its use, PN was relatively expensive, but its components now are often less costly than specialty enteral formulas. Percutaneous placement of a central venous catheter

into the subclavian vein or (less desirably) the internal jugular vein with advancement into the superior vena cava can be accomplished at the bedside by trained personnel using sterile techniques. Peripherally inserted central catheters (PICCs) can also be used, although they are usually more appropriate for non-ICU patients. Subclavian or internal jugular catheters carry the risks of pneumothorax or serious vascular damage but are generally well tolerated and, rather than requiring reinsertion, can be exchanged over a wire when catheter infection is suspected.

Although most SNS is delivered in hospitals, some patients require it on a long-term basis. At-home SNS requires a safe home environment, a stable clinical condition, and the patient's ability and willingness to learn appropriate self-care techniques. Other important considerations in determining the appropriateness of at-home parenteral or enteral SNS are that the patient's prognosis indicates survival for longer than several months and that the therapy enhances the patient's quality of life.

DISEASE-SPECIFIC NUTRITIONAL SUPPORT

The purpose of SNS is to correct and prevent malnutrition. Certain conditions require special modification of the SNS regimen. Protein intake may need to be limited in many stable patients with renal insufficiency or borderline liver function. In renal disease, except for brief periods, protein intakes should approach the required level for normal adults of at least 0.8 g/kg and should aim for 1.2 g/kg as long as severe azotemia does not occur. Patients with severe renal failure who require SNS need concurrent renal replacement therapy. In hepatic failure, protein intakes of 1.2–1.4 g/kg (up to 1.5 g/kg) should be provided as long as encephalopathy due to protein intolerance does not occur. In the presence of protein intolerance, formulas containing 33–50% branched-chain amino acids are available and can be provided at the 1.2- to 1.4-g/kg level. Cardiac patients and many other severely stressed patients often benefit from fluid and sodium restriction to 1000 mL of PN formula and 5–20 meq of sodium per day. In patients with severe chronic PEM characterized by severe weight loss, it is important to initiate PN gradually because of the profound antinatriuresis, antidiuresis, and intracellular accumulation of potassium, magnesium, and phosphorus that develop as a consequence of the resulting high insulin levels. This modification of parenteral SNS is usually accomplished by limiting daily fluid intake initially to ~1000 mL; limiting carbohydrate intake to 10–20% dextrose; limiting sodium intake; and providing ample potassium, magnesium, and phosphorus, with careful daily assessment of fluid and electrolyte status. Protein need not be restricted.

THE DESIGN OF INDIVIDUAL REGIMENS

FLUID REQUIREMENTS

Normal adults require ~30 mL of fluid/kg of body weight from all sources each day as well as the replacement of abnormal losses such as those caused by diuretic therapy, nasogastric tube drainage, wound output, high rates of perspiration (which can be several liters per day during periods of extreme heat), and diarrhea/ostomy losses. Electrolyte and mineral losses can be estimated or measured and need to be replaced (Table 98e-2). Fluid restriction may be necessary in patients with fluid overload. Total fluid input can usually be limited to 1200 mL/d as long as urine is the only significant source of fluid output. In severe fluid overload, a 1-L central vein PN solution of 7% crystalline amino acids (70 g) and 21% dextrose (210 g) can temporarily provide an acceptable amount of glucose and protein substrate in the absence of significant catabolic stress.

Patients who require PN or EN in the acute-care setting generally have associated hormonal adaptations to their underlying disease (e.g., increased secretion of antidiuretic hormone, aldosterone, insulin, glucagon, or cortisol), and these signals promote fluid retention and hyperglycemia. In critical illness, body weight is invariably increased due to fluid resuscitation and fluid retention. Lean-tissue accretion is minimal in the acute phase of critical illness, no matter how much

TABLE 98e-2 APPROXIMATE VOLUME AND COMPOSITION OF FLUID SECRETIONS^a

Secretion	Volume (L/d)	Na	K	Cl	HCO ₃	pH
Saliva	1–2	15	30	15	30	6–8
Gastric juice	1–2	50–90	5–30	90–130	0	1.5–3.5
Hepatic bile	0.5–1	130–150	5–10	80–120	30–50	7–9
Pancreas	0.5–1	130–150	5–10	70–100	90–110	8–9
Small intestine	1–3	120–140	10–20	80–120	20–40	7–9
Diarrhea	Varies	30–50	20–30		Varies	

^aConcentration values are in mmol/L.

protein and or how many calories are provided. Because excess fluid removal can be difficult, limiting fluid intake to allow for balanced intake and output is more effective.

ENERGY REQUIREMENTS

Total energy expenditure comprises resting energy expenditure, activity energy expenditure, and the thermal effect of feeding (Chap. 97). Resting energy expenditure accounts for two-thirds of total energy expenditure, activity energy expenditure for one-fourth to one-third, and the thermal effect of feeding for ~10%. For normally nourished, healthy individuals, the total energy expenditure is ~30–35 kcal/kg. Critical illness increases resting energy expenditure, but this increase is significant only in initially well-nourished individuals with a robust SRI who experience, for example, severe multiple trauma, extensive burns, sepsis, sustained high fever, or closed head injury. In these situations, total energy expenditure can reach 40–45 kcal/kg. The chronically starved patient with adapted PEM has a reduced energy expenditure and is inactive, with a usual total energy expenditure of ~20–25 kcal/kg. Very few patients with adapted PEM require as much as 30 kcal/kg for energy balance. Because providing ~50% of measured energy expenditure as SNS is at least as effective as 100% for the first 10 days of critical illness, actual measurement of energy expenditure generally is not necessary in the early period of SNS. However, in patients who remain critically ill beyond several weeks, in patients with severe PEM for whom estimates of energy expenditure are unreliable, and in patients who are difficult to wean from ventilators, it is reasonable to measure energy expenditure directly when the technique is available, targeting an energy intake of 100–120% of the measured energy expenditure.

Insulin resistance due to the SRI is associated with increased gluconeogenesis and reduced peripheral glucose utilization, with resulting hyperglycemia. Hyperglycemia is aggravated by excessive exogenous carbohydrate administration from SNS. In critically ill patients receiving SNS, normalization of blood glucose levels by insulin infusion reduces morbidity and mortality risk. In mildly or moderately malnourished patients, it is reasonable to provide metabolic support in order to improve protein synthesis and maintain metabolic homeostasis. Hypocaloric nutrition, with provision of ~1000 kcal and 70 g protein per day for up to 10 days, requires less fluid and reduces the likelihood of poor glycemic control, although a higher protein intake would be optimal. During the second week of SNS, energy and protein provision can be advanced to 20–25 kcal/kg and 1.5 g/kg per day, respectively, as metabolic conditions permit. As mentioned above, patients with multiple trauma, closed head injury, and severe burns often have greatly elevated energy expenditures, but there is little evidence that providing >30 kcal/kg daily confers further benefit, and such high caloric intake may well be harmful as it substantially increases the risk of hyperglycemia.

As a rule, amino acids and glucose are provided in an increasing dose until energy provision matches estimated resting energy expenditure. At this point, it becomes beneficial to add fat. A surfeit of glucose merely stimulates de novo lipogenesis—an energy-inefficient process. Polyunsaturated long-chain triglycerides (e.g., in soybean oil) are the chief ingredient in most parenteral fat emulsions and provide the majority of the fat in enteral feeding formulas. These vegetable oil-based emulsions provide essential fatty acids. The fat content of

enteral feeding formulas ranges from 3% to 50% of energy. Parenteral fat is provided in separate containers as 20% and 30% emulsions that can be infused separately or mixed in the sterile pharmacy as an all-in-one or total nutrient admixture of amino acids, glucose, lipid, electrolytes, vitamins, and minerals. Although parenteral fat needs to make up only ~3% of the energy requirement in order to meet essential fatty acid requirements, when provided daily as an all-in-one mixture of carbohydrate, fat, and protein, the complete admixture has a fat content of 2–3 g/dL and provides 20–30% of the total energy requirement—an acceptable level that offers the advantage of ensuring emulsion stability. When given as a separate infusion, parenteral fat should not be provided at rates exceeding 0.11 g/kg of body mass or 100 g over 12 h—equivalent to 500 mL of 20% parenteral fat.

Medium-chain triglycerides containing saturated fatty acids with chain lengths of 6, 8, 10, or 12 carbons (>95% of which are C8 and C10) are included in a number of enteral feeding formulas because they are absorbed preferentially. Fish oil contains polyunsaturated fatty acids of the omega 3 family, which improve immune function and reduce the inflammatory response. At this time, fish oil injectable emulsions are available in the United States as an investigational new drug.

PN formulations provide carbohydrate as hydrous glucose (3.4 kcal/g). In enteral formulas, glucose is the carbohydrate source for so-called monomeric diets. These diets provide protein as amino acids and fat in minimal amounts (3%) to meet essential fatty acid requirements. Monomeric formulas are designed to optimize absorption in the seriously compromised gut. These formulas, like immune-enhancing diets, are expensive. In polymeric diets, the carbohydrate source is usually an osmotically less active polysaccharide, the protein is usually soy or casein protein, and fat is present at concentrations of 25–50%. Such formulas are usually well tolerated by patients with normal intestinal length, and some are acceptable for oral consumption.

PROTEIN OR AMINO ACID REQUIREMENTS

The daily protein recommendation for healthy adults is 0.8 g/kg, but body proteins are replenished faster with 1.5 g/kg in patients with PEM, and net protein catabolism is reduced in critically ill patients when 1.5–2.0 g/kg is provided. In patients who are not critically ill but who require SNS in the acute-care setting, at least 1 g of protein/kg is recommended, and larger amounts up to 1.5 g/kg are appropriate when volume, renal, and hepatic tolerances allow. The standard parenteral and enteral formulas contain protein of high biologic value and meet the requirements for the eight essential amino acids when nitrogen needs are met. Parenteral amino acid mixtures and elemental enteral mixtures consist of hydrated individual amino acids. Because of their hydrated status, elemental amino acid solutions deliver 17% less protein substrate than intact proteins. In protein-intolerant conditions such as renal and hepatic failure, modified amino acid formulas may be considered. In hepatic failure, higher branched-chain, amino acid-enriched formulas appear to improve outcomes. Conditionally essential amino acids like arginine and glutamine may also have some benefit in supplemental amounts.

Protein (nitrogen) balance provides a measure of the efficacy of parenteral or enteral SNS. This balance is calculated as protein intake/6.25 (because proteins are, on average, 16% nitrogen) minus the 24-h urine urea nitrogen plus 4 g of nitrogen (the latter reflecting other nitrogen losses). In critical illness, a mild negative nitrogen balance of 2–4 g/d is often achievable. A similarly mild positive nitrogen balance is observed in the nonstressed recuperating patient. Each gram of nitrogen lost or gained represents ~30 g of lean tissue.

MINERAL AND VITAMIN REQUIREMENTS

Parenteral electrolyte, vitamin, and trace mineral requirements are summarized in [Tables 98e-3, 98e-4, and 98e-5](#), respectively. Electrolyte modifications are necessary with substantial gastrointestinal losses from nasogastric drainage or intestinal losses from fistulas, diarrhea, or ostomy outputs. Such losses also imply extra calcium, magnesium, and zinc losses. Zinc losses are high in secretory diarrhea. Secretory diarrhea contains ~12 mg of zinc/L, and patients with intestinal fistulas or chronic diarrhea require an average of ~12 mg of

TABLE 98e-3 USUAL DAILY ELECTROLYTE ADDITIONS TO PARENTERAL NUTRITION

Electrolyte	Parenteral Equivalent	Usual Intake
Sodium		1–2 meq/kg + replacement, but can be as low as 5–40 meq/d
Potassium		40–100 meq/d + replacement of unusual losses
Chloride		As needed for acid-base balance, but usually 2:1 to 1:1 with acetate
Acetate		As needed for acid-base balance
Calcium	10 meq	10–20 meq/d
Magnesium	10 meq	8–16 meq/d
Phosphorus	30 mmol	20–40 mmol

parenteral zinc/d (equivalent to 30 mg of oral elemental zinc) to maintain zinc balance. Excessive urinary potassium losses with amphotericin or magnesium losses with cisplatin or in renal failure necessitate adjustments in sodium, potassium, magnesium, phosphorus, and acid-base balance. Vitamin and trace element requirements are met by the daily provision of a complete parenteral vitamin supplement and trace elements via PN and by the provision of adequate amounts of enteral feeding formulas that contain these micronutrients.

Iron is a highly reactive catalyst of oxidative reactions and thus is not included in PN mixtures. The parenteral iron requirement is normally only ~1 mg/d. Iron deficiency occurs with considerable frequency in acutely ill hospitalized patients, especially those with PEM and gastrointestinal tract disease, and in patients subjected to frequent blood withdrawals. Iron deficiency is sometimes inadequately considered in hospitalized patients because there are commoner causes: the inflammation-mediated anemia of chronic disease (with an associated increase in serum ferritin, an acute-phase protein) and redistribution of the intravascular fluid volume during prolonged bed rest. Iron deficiency should be considered in every patient receiving SNS. A falling mean red cell volume, even if still in the low-normal range, together with an intermediate serum ferritin concentration is suggestive of iron deficiency. Intravenous iron infusions follow standard guidelines, always with a termination order and never as a standing order because of the risk of inadvertent iron overdosing. Major iron replacement during critical illness is of some concern because of the possibility that a substantial rise in the serum iron concentration may increase susceptibility to some bacterial infections.

TABLE 98e-4 PARENTERAL MULTIVITAMIN REQUIREMENTS FOR ADULTS

Vitamin	Revised Values
Vitamin A	3300 IU
Thiamin (B ₁)	6 mg
Riboflavin (B ₂)	3.6 mg
Niacin (B ₃)	40 mg
Folic acid	600 µg
Pantothenic acid	15 mg
Pyridoxine (B ₆)	6 mg
Cyanocobalamin (B ₁₂)	5 µg
Biotin	60 µg
Ascorbic acid (C)	200 mg
Vitamin D	200 IU ^a
Vitamin E	10 IU
Vitamin K ^b	150 µg

^aThe current vitamin D requirement—a minimum of 600 IU/day—cannot be met with available injectable vitamin formulations. Calcitriol is not equivalent to vitamin D and is not a suitable replacement for it, since it is not a substrate for 25-hydroxyvitamin D biosynthesis. ^bA product is available without vitamin K. Vitamin K supplementation is recommended at 2–4 mg/week in patients not receiving oral anticoagulation therapy when the vitamin K-free product is used.

TABLE 98e-5 PARENTERAL TRACE METAL SUPPLEMENTATION FOR ADULTS^a

Trace Mineral	Intake
Zinc ^b	2.5–4 mg/d; an additional 10–15 mg/d per L of stool or ileostomy output
Copper	0.5–1.5 mg/d; possibility of retention in biliary tract obstruction
Manganese	0.1–0.3 mg/d; possibility of retention in biliary tract obstruction
Chromium	10–15 µg/d
Selenium	20–100 µg/d; necessary for long-term PN, optional for short term
Molybdenum	20–120 µg/d; necessary for long-term PN, optional for short term
Iodine	75–150 µg/d; necessary for long-term PN, optional for short term

^aCommercial products are available with the first four, the first five, and all seven of these metals in recommended amounts. ^bThe basal IV zinc requirement is approximately one-third of the oral requirement, because only approximately one-third of orally ingested zinc is absorbed.

Abbreviation: PN, parenteral nutrition.

PARENTERAL NUTRITION

INFUSION TECHNIQUE AND PATIENT MONITORING

Parenteral feeding through a peripheral vein is limited by osmolarity and volume constraints. Solutions with an osmolarity >900 mOsm/L (e.g., those which contain >3% amino acids and 5% glucose [290 kcal/L]) are poorly tolerated peripherally. Parenteral lipid emulsions (20%) can be given to increase the calories delivered. The total volume required for a marginal amino acid provision rate of 60 g (equivalent to 50 g of protein) and a total of 1680 kcal is 2.5 L. Moreover, the risk of significant morbidity and mortality from incompatibilities of calcium and phosphate salts is greatest in these low-osmolarity, low-glucose regimens. For short-term infusions, calcium may be temporarily limited or even omitted from the mixture. Parenteral feeding via a peripheral vein is generally intended as a supplement to oral feeding; it is not suitable for the critically ill. Peripheral PN may be enhanced by small amounts of heparin (1000 U/L) and co-infusion with parenteral fat to reduce osmolarity, but volume constraints still limit the value of this therapy, especially in critical illness.

PICCs may be used to infuse solutions of 20–25% dextrose and 4–7% amino acids, thus avoiding the traumatic complications of percutaneous central vein catheter placement. With PICC lines, however, flow can be position-related, and the lines cannot be exchanged over a wire for infection monitoring. It is important to withdraw blood samples carefully and appropriately from a dual-port PICC because intermixing of the blood sample with even tiny volumes of nutrient infusate will falsely indicate hyperglycemia and hyperkalemia. For all these reasons, centrally placed catheters are preferred in critical illness. The subclavian approach is best tolerated by the patient and is the easiest to dress. The jugular approach is less likely to cause a pneumothorax. Femoral vein catheterization is strongly discouraged because of the risk of catheter infection. For long-term feeding at home, tunneled catheters and implanted ports are used to reduce infection risk and are more acceptable to patients. Tunneled catheters require placement in the operating room.

Catheters are made of Silastic[®], polyurethane, or polyvinyl chloride. Silastic catheters are less thrombogenic and are best for tunneled catheters. Polyurethane is best for temporary catheters. To avoid infection, dressing changes with dry gauze should be performed at regular intervals by nurses skilled in catheter care. Chlorhexidine solution is more effective than alcohol or iodine compounds. Appropriate monitoring for patients receiving PN is summarized in [Table 98e-6](#).

STANDARD VERSUS INDIVIDUALIZED NUTRIENT PROVISION

Even though premixed solutions of crystalline amino acids and dextrose are in common use, the future of evidence-based PN lies in computer-controlled sterile compounders that rapidly and inexpensively

TABLE 98e-6 MONITORING THE PATIENT RECEIVING PARENTERAL NUTRITION^a

Clinical Data	
General sense of well-being	
Strength, as evidenced by getting out of bed, walking, and resistance exercise as appropriate	
Vital signs, including temperature, blood pressure, pulse, and respiratory rate	
Fluid balance: weight (recorded at least several times weekly); fluid intake (parenteral and enteral) vs. fluid output (urine, stool, gastric drainage, wound, ostomy)	
Parenteral nutrition delivery equipment: tubing, pump, filter, catheter, dressing	
Nutrient solution composition	
Laboratory Data	
Finger-stick glucose	Three times daily until stable
Blood glucose, Na, K, Cl, HCO ₃ , BUN	Daily until stable and fully advanced; then twice weekly
Serum creatinine, albumin, PO ₄ , Ca, Mg, Hb/Hct, WBC count	Baseline; then twice weekly
INR	Baseline; then weekly
Micronutrient tests	As indicated

^aParameters are assessed daily unless otherwise specified.

Abbreviations: BUN, blood urea nitrogen; Hb, hemoglobin; Hct, hematocrit; INR, international normalized ratio; WBC, white blood cell.

Source: Adapted from the chapter on this topic in *Harrison's Principles of Internal Medicine*, 16e, by Lyn Howard, MD.

generate personalized solutions that meet the specific protein and calorie goals for different patients in different clinical situations. For example, 1 L of a standard mixture of 5% amino acids/25% dextrose solution provides 50 g of amino acids (41.5 g of protein substrate) and 1000 kcal; the use of this solution to meet the 1.5–to 2.0-g/kg protein requirement of an acutely ill 70-kg patient requires the infusion of 2.5–3.4 L of fluid and a potentially excessively high energy dose of 2500–3300 kcal. When the body fat store is adequate, clinical evidence increasingly supports the greater safety and efficacy of high-protein, moderately hypocaloric SNS in such patients. A sterile compounder can accurately generate an appropriate recipe for such a patient. For example, 1 L of a solution including 600 mL of 15% amino acids, 300 mL of 50% dextrose, and 100 mL of electrolyte/micronutrient mix contains 75 g of protein substrate and 800 kcal; thus it is feasible to meet the patient's protein requirement with only 1.4–1.9 L of solution and a more appropriate 1100–1520 kcal; any mild gap in energy provision is easily filled by use of intravenous lipid.

COMPLICATIONS

Mechanical The insertion of a central venous catheter should be performed by trained and experienced personnel using aseptic techniques to limit the major common complications of pneumothorax and inadvertent arterial puncture or injury. The catheter's position should be radiographically confirmed to be in the superior vena cava distal to the junction with the jugular or subclavian vein and not directly against the vessel wall. Thrombosis related to the catheter may occur at the site of entry into the vein and extend to encase the catheter. Catheter infection predisposes to thrombosis, as does the SRI. The addition of 6000 U of heparin to the daily parenteral formula for hospitalized patients with temporary catheters reduces the risk of fibrin sheath formation and catheter infection. Temporary catheters that develop a thrombus should be removed and, according to clinical findings, treated with anticoagulants. Thrombolytic therapy can be considered for patients with permanent catheters, depending on the ease of replacement and the presence of alternative, reasonably acceptable venous access sites. Low-dose warfarin therapy (1 mg/d) reduces the risk of thrombosis in permanent catheters used for at-home parenteral SNS, but full anticoagulation may be required for patients who have recurrent thrombosis related to permanent catheters. A recent U.S. Food and Drug Administration mandate to reformulate parenteral multivitamins to include vitamin K at a dose of 150 µg/d may affect

the efficacy of low-dose warfarin therapy. A “no vitamin K” version is available for patients receiving this therapy. Catheters can become occluded due to mechanical factors; by fibrin at the tip; or by fat, minerals, or drugs intraluminally. These occlusions can be managed with low-dose alteplase for fibrin, with indwelling 70% alcohol for fat, with 0.1 N hydrochloric acid for mineral precipitates, and with either 0.1 N hydrochloric acid or 0.1 N sodium hydroxide for drugs, depending on the pH of the drug.

Metabolic The most common problems caused by parenteral SNS are fluid overload and hyperglycemia (Table 98e-7). Hypertonic dextrose stimulates a much higher insulin level than meal feeding. Because insulin is a potent antinatriuretic and antidiuretic hormone,

hyperinsulinemia leads to sodium and fluid retention. Consequently, in the absence of gastrointestinal losses or renal dysfunction, net fluid retention is likely when total fluid intake exceeds 2000 mL/d. Close monitoring of body mass as well as of fluid intake and output is necessary to prevent this complication. In the absence of significant renal impairment, the sodium content of the urine is likely to be <10 meq/L. Provision of sodium in limited amounts (40 meq/d) and the use of both glucose and fat in the PN mixture will reduce serum glucose levels and help reduce fluid retention. The elevated insulin level also increases the intracellular transport of potassium, magnesium, and phosphorus, which can precipitate a dangerous re-feeding syndrome if the total glucose content of the PN solution is advanced too quickly in severely malnourished patients. To assess glucose tolerance, it is generally best to start PN with <200 g of glucose/d. Regular insulin can be added to the PN formula to establish glycemic control, and the insulin doses can be increased proportionately as the glucose content is advanced. As a general rule, patients with insulin-dependent diabetes require about twice their usual at-home insulin dose when receiving PN at 20–25 kcal/kg, largely as a consequence of parenteral glucose administration and some loss of insulin to the formula’s container. As a rough estimate, the amount of insulin provided can be proportionately similar to the number of calories provided as total parenteral nutrition (TPN) relative to full feeding, and the insulin can be placed in the TPN formula. Subcutaneous regular insulin can be provided to improve glucose control as assessed by measurements of blood glucose every 6 h. About two-thirds of the total 24-h amount can be added to the next day’s order, with SC insulin supplements as needed. Advances in the TPN glucose concentration should be made when reasonable glucose control is established, and the insulin dose can be adjusted proportionately to the calories added as glucose and amino acids. These are general rules, and they are conservative. Given the adverse clinical impact of hyperglycemia, it may be necessary to use intensive insulin therapy as a separate infusion with a standard protocol to initially establish control. Once control is established, this insulin dose can be added to the PN formula. Acid-base imbalance is also common during parenteral SNS. Amino acid formulas are buffered, but critically ill patients are prone to metabolic acidosis, often due to renal tubular impairment. The use of sodium and potassium acetate salts in the PN formula may address this problem. Bicarbonate salts should not be used because they are incompatible with TPN formulations. Nasogastric drainage produces hypochloremic alkalosis that can be managed by attention to chloride balance. Occasionally, hydrochloric acid may be required for a more rapid response or when diuretic therapy limits the ability to provide substantial sodium chloride. Up to 100 meq/L and up to 150 meq of hydrochloric acid per day may be placed in a fat-free TPN formula.

Infectious Infections of the central access catheter rarely occur in the first 72 h. Fever during this period is usually attributable to infection elsewhere or another cause. Fever that develops during parenteral SNS can be addressed by checking the catheter site and, if the site looks clean, exchanging the catheter over a wire, with cultures taken through the catheter and at the catheter tip. If these cultures are negative, as they usually are, the new catheter can continue to be used. If a culture is positive for a relatively nonpathogenic bacterium like *Staphylococcus epidermidis*, a second exchange over a wire with repeat cultures or replacement of the catheter can be considered in light of the clinical circumstances. If cultures are positive for more pathogenic bacteria or for fungi like *Candida albicans*, it is generally best to replace the catheter at a new site. Whether antibiotic treatment is required is a clinical decision, but *C. albicans* grown from the blood culture in a patient receiving PN should always be treated with an antifungal drug because the consequences of failure to treat can be dire.

Catheter infections can be minimized by dedicating the feeding catheter to TPN, without blood sampling or medication administration. Central catheter infections are a serious complication, with an attributed mortality rate of 12–25%. Fewer than three infections per 1000 catheter-days should occur in central venous catheters dedicated to feeding. At-home TPN catheter infections may be treated through

TABLE 98e-7 SELECTED METABOLIC DISTURBANCES CAUSED BY PN AND THEIR CORRECTION

Disturbance	Cause	Corrective Action with PN
Hyponatremia	Increased total body water or decreased total body sodium	Decrease free water or increase sodium.
Hypernatremia	Occurs commonly with excessive isotonic or hypertonic fluid followed by diuretic administration with free water clearance; can also occur with dehydration and normal total body sodium	Increase free water to produce net positive fluid balance, maintaining sodium and chloride balance.
Hypokalemia	Inadequate intake relative to need	Use supplements.
	Excessive diuresis, tubular dysfunction	Use supplements.
	Magnesium deficiency	Increase PN magnesium.
	Metabolic alkalosis	Correct alkalosis.
Hyperkalemia	Excessive provision	Reduce supplements.
	Metabolic acidosis	Evaluate acidosis. Treat with PN acetate salt, and decrease potassium.
	Renal deterioration	Evaluate patient and adjust PN as indicated.
Hypocalcemia	Reciprocal response to phosphorus repletion	Increase calcium.
	Critical illness effect	Increase calcium.
	Severe malabsorption	Supplement calcium.
Hypercalcemia	Excessive administration or pathology (cancer, hyperparathyroidism)	Reduce or eliminate calcium.
Hypomagnesemia	Increased requirements due to diuretic use, alcoholism, malabsorption, malnutrition	Supplement magnesium.
	Critical illness	Supplement magnesium.
Hypophosphatemia	Inadequate intake relative to needs related to malnutrition, alcohol use	Supplement phosphorus.
	Increased calcium intake	Use supplements.
Hyperphosphatemia	Excessive administration or worsening renal function	Reduce phosphorus.
Azotemia	Excessive amino acid infusion or worsening renal function	Reduce amino acid level if feasible, but use renal replacement therapy if 1 g of protein/kg cannot be provided for prolonged periods.

Abbreviation: PN, parenteral nutrition.

the catheter without its removal, particularly if the offending organism is *S. epidermidis*. Clearing of the biofilm and fibrin sheath by local treatment of the catheter with indwelling alteplase may increase the likelihood of eradication. Antibiotic lock therapy with high concentrations of antibiotic, with or without heparin in addition to systemic therapy, may improve efficacy. Sepsis with hypotension should precipitate catheter removal in either the temporary or the permanent TPN setting.

ENTERAL NUTRITION

TUBE PLACEMENT AND PATIENT MONITORING

The types of enteral feeding tubes, methods of insertion, their clinical uses, and potential complications are outlined in [Table 98e-8](#). The different types of enteral formulas are listed in [Table 98e-9](#). Patients receiving EN are at risk for many of the same metabolic complications as those who receive PN and should be monitored in the same manner. EN can be a source of similar problems, but not to the same degree, because the insulin response to EN is about half of that to PN. Enteral feeding formulas have fixed electrolyte compositions that are generally modest in sodium and somewhat higher in potassium. Acid-base disturbances can be addressed to a more limited extent with EN.

TABLE 98e-8 ENTERAL FEEDING TUBES

Type/Insertion Technique	Clinical Uses	Potential Complications
Nasogastric Tube		
External measurement: nostril, ear, xiphisternum; tube stiffened by ice water or stylet; position verified by air injection and auscultation or by x-ray	Short-term clinical situation (weeks) or longer periods with intermittent insertion; bolus feeding is simpler, but continuous drip with pump is better tolerated	Aspiration; ulceration of nasal and esophageal tissues, leading to stricture
Nasoduodenal Tube		
External measurement: nostril, ear, anterior superior iliac spine; tube stiffened by stylet and passed through pylorus under fluoroscopy or with endoscopic loop	Short-term clinical situations where gastric emptying is impaired or proximal leak is suspected; requires continuous drip with pump	Spontaneous pulling back into stomach (position verified by aspirating content, pH >6); diarrhea common, fiber-containing formulas may help
Gastrostomy Tube		
Percutaneous placement endoscopically, radiologically, or surgically; after track is established, can be converted to a gastric "button"	Long-term clinical situations, swallowing disorders, or impaired small-bowel absorption requiring continuous drip	Aspiration; irritation around tube exit site; peritoneal leak; balloon migration and obstruction of pylorus
Jejunostomy Tube		
Percutaneous placement endoscopically or radiologically via pylorus or endoscopically or surgically directly into the jejunum	Long-term clinical situations where gastric emptying is impaired; requires continuous drip with pump; direct endoscopic placement (PEJ) is most comfortable for patient	Clogging or displacement of tube; jejunal fistula if large-bore tube is used; diarrhea from dumping; irritation of surgical anchoring suture
Combined Gastrojejunostomy Tube		
Percutaneous placement endoscopically, radiologically, or surgically; intragastric arm for continuous or intermittent gastric suction; jejunal arm for enteral feeding	Used for patients with impaired gastric emptying and high risk for aspiration or patients with acute pancreatitis or proximal leaks	Clogging, especially of small-bore jejunal tube

Abbreviation: PEJ, percutaneous endoscopic jejunostomy.

Note: All small tubes are at risk for clogging, especially if used for crushed medications. In long-term enteral nutrition patients, gastrostomy and jejunostomy tubes can be exchanged for a low-profile "button" once the track is established.

Source: Adapted from the chapter on this topic in *Harrison's Principles of Internal Medicine*, 16e, by Lyn Howard, MD.

TABLE 98e-9 ENTERAL FORMULAS

Composition Characteristics	Clinical Indications
Standard Enteral Formula	
Complete dietary products (+) ^a	Suitable for most patients requiring tube feeding; some standard formulas can be used orally
1. Caloric density: 1 kcal/mL	
2. Protein: ~14% cals (caseinates, soy, lactalbumin)	
3. Carbohydrate: ~60% cals (hydrolyzed corn starch, maltodextrin, sucrose)	
4. Fat: ~30% cals (corn, soy, safflower oils)	
5. Recommended daily intake of all minerals and vitamins in >1500 kcal/d	
6. Osmolality: ~300 mosmol/kg	
Modified Enteral Formulas	
1. Caloric density: 1.5–2 kcal/mL (+)	Fluid-restricted patients
2. Protein	
a. High protein (~20–25% protein) (+)	Critically ill patients
b. Hydrolyzed protein to small peptides (+)	Impaired absorption
c. ↑ Arginine, glutamine, nucleotides, ω3 fat (+++)	Immune-enhancing diets
d. ↑ Branched-chain amino acids, ↓ aromatic amino acids (+++)	Liver failure patients intolerant of 0.8 g of protein/kg
e. Low protein of high biologic value	Renal failure patients for brief periods if critically ill
3. Fat	
a. Low-fat partial MCT substitution (+)	Fat malabsorption
b. ↑ Fat (>40% cals) (++)	Pulmonary failure with CO ₂ retention on standard formula, limited utility
c. ↑ Fat from MUFA (++)	Improvement in glycemic index control in diabetes
d. ↑ Fat from ω3 and ↓ fat from ω6 linoleic acid (+++)	Improved ventilation in ARDS
4. Fiber: provided as soy polysaccharide (+)	Improved laxation

^aCost: +, inexpensive; ++, moderately expensive; +++, very expensive.

Note: ARDS, acute respiratory distress syndrome; MCT, medium-chain triglyceride; MUFA, monounsaturated fatty acids; ω3 or ω6, polyunsaturated fat with first double bond at carbon 3 (fish oils) or carbon 6 (vegetable oils).

Source: Adapted from the chapter on this topic in *Harrison's Principles of Internal Medicine*, 16e, by Lyn Howard, MD.

Acetate salts can be added to the formula to treat chronic metabolic acidosis. Calcium chloride can be added to treat mild chronic metabolic alkalosis. Medications and other additives to enteral feeding formulas can clog the tubes (e.g., calcium chloride may interact with casein-based formulas to form insoluble calcium caseinate products) and may reduce the efficacy of some drugs (e.g., phenytoin). Since small-bore tubes are easily displaced, tube position should be checked at intervals by aspirating and measuring the pH of the gut fluid (normal: <4 in the stomach, >6 in the jejunum).

COMPLICATIONS

Aspiration The debilitated patient with poor gastric emptying and impairment of swallowing and cough is at risk for aspiration; this complication is particularly common among patients who are mechanically ventilated. Tracheal suctioning induces coughing and gastric regurgitation, and cuffs on endotracheal or tracheostomy tubes seldom protect against aspiration. Preventive measures include elevating the head of the bed to 30°, using nurse-directed algorithms for formula advancement, combining enteral with parenteral feeding, and using

post-ligament of Treitz feeding. Tube feeding should not be discontinued for gastric residuals of <300 mL unless there are other signs of gastrointestinal intolerance, such as nausea, vomiting, or abdominal distention. Continuous feeding using pumps is better tolerated intragastrically than bolus feeding and is essential for feeding into the jejunum. For small-bowel feeding, residuals are not assessed, but abdominal pain and distention should be monitored.

Diarrhea Enteral feeding often leads to diarrhea, especially if bowel function is compromised by disease or drugs (most often, broad-spectrum antibiotics). Sorbitol used to flavor some medications can also cause diarrhea. Diarrhea may be controlled by the use of a continuous drip, with a fiber-containing formula, or by the addition of an antidiarrheal agent to the formula. However, *Clostridium difficile*, which is a common cause of diarrhea in patients being tube-fed, should be ruled out as the etiology before antidiarrheal agents are used. H₂ blockers may help reduce the net volume of fluid presented to the colon. Diarrhea associated with enteral feeding does not necessarily imply inadequate absorption of nutrients other than water and electrolytes. Amino acids and glucose are particularly well absorbed in the upper small bowel except in the most diseased or shortest bowel. Since luminal nutrients exert trophic effects on the gut mucosa, it is often appropriate to persist with tube feeding despite diarrhea, even when this course necessitates supplemental parenteral fluid support.

Apart from conditions with drastically diminished small-intestinal absorptive function, there are no established indications for short peptide-based or elemental formulas.

GLOBAL CONSIDERATIONS



In the United States, the only parenteral lipid emulsion available is made with soybean oil, whose constituent fatty acids have been suggested to be immunosuppressive under certain circumstances. In Europe and Japan, a number of other lipid emulsions are available, including those containing fish oil only; mixtures of fish oil, medium-chain triglycerides, and long-chain triglycerides as olive oil and/or soybean oil; mixtures of medium-chain triglycerides and long-chain triglycerides as soybean oil; and long-chain triglyceride mixtures as olive oil and soybean oil, which may be more beneficial in terms of metabolism and hepatic and immune function. Furthermore, a glutamine-containing dipeptide for inclusion in TPN formulas is available in Europe and may be helpful in terms of immune function and resistance to infection, although a recent study using a larger-than-recommended dose was associated with net harm.

ACKNOWLEDGMENT

The authors acknowledge the contributions of Lyn Howard, MD, the author in earlier editions of HPIM, to material in this chapter.

This page intentionally left blank

This page intentionally left blank

SECTION 1 NEOPLASTIC DISORDERS

99 Approach to the Patient with Cancer

Dan L. Longo

The application of current treatment techniques (surgery, radiation therapy, chemotherapy, and biologic therapy) results in the cure of nearly two of three patients diagnosed with cancer. Nevertheless, patients experience the diagnosis of cancer as one of the most traumatic and revolutionary events that has ever happened to them. Independent of prognosis, the diagnosis brings with it a change in a person's self-image and in his or her role in the home and workplace. The prognosis of a person who has just been found to have pancreatic cancer is the same as the prognosis of the person with aortic stenosis who develops the first symptoms of congestive heart failure (median survival, ~8 months). However, the patient with heart disease may remain functional and maintain a self-image as a fully intact person with just a malfunctioning part, a diseased organ ("a bum ticker"). By contrast, the patient with pancreatic cancer has a completely altered self-image and is viewed differently by family and anyone who knows the diagnosis. He or she is being attacked and invaded by a disease that could be anywhere in the body. Every ache or pain takes on desperate significance. Cancer is an exception to the coordinated interaction among cells and organs. In general, the cells of a multicellular organism are programmed for collaboration. Many diseases occur because the specialized cells fail to perform their assigned task. Cancer takes this malfunction one step further. Not only is there a failure of the cancer cell to maintain its specialized function, but it also strikes out on its own; the cancer cell competes to survive using natural mutability and natural selection to seek advantage over normal cells in a recapitulation of evolution. One consequence of the traitorous behavior of cancer cells is that the patient feels betrayed by his or her body. The cancer patient feels that he or she, and not just a body part, is diseased.

THE MAGNITUDE OF THE PROBLEM

No nationwide cancer registry exists; therefore, the incidence of cancer is estimated on the basis of the National Cancer Institute's Surveillance, Epidemiology, and End Results (SEER) database, which tabulates cancer incidence and death figures from 13 sites, accounting for about 10% of the U.S. population, and from population data from the U.S. Census Bureau. In 2014, 1.665 million new cases of invasive cancer (855,220 men, 810,320 women) were diagnosed, and 585,720 persons (310,010 men, 275,710 women) died from cancer. The percent distribution of new cancer cases and cancer deaths by site for men and women is shown in [Table 99-1](#). Cancer incidence has been declining by about 2% each year since 1992. Cancer is the cause of one in four deaths in the United States.

The most significant risk factor for cancer overall is age; two-thirds of all cases were in those older than age 65 years. Cancer incidence increases as the third, fourth, or fifth power of age in different sites. For the interval between birth and age 49 years, 1 in 29 men and 1 in 19 women will develop cancer; for the interval between ages 50 and 59 years, 1 in 15 men and 1 in 17 women will develop cancer; for the interval between ages 60 and 69 years, 1 in 6 men and 1 in 10 women will develop cancer; and for people age 70 and older, 1 in 3 men and 1 in 4 women will develop cancer. Overall, men have a 44% risk of developing cancer at some time during their lives; women have a 38% lifetime risk.


TABLE 99-1 DISTRIBUTION OF CANCER INCIDENCE AND DEATHS FOR 2014

Sites	Male		Female		
	%	Number	%	Number	
Cancer Incidence					
Prostate	27	233,000	Breast	29	232,670
Lung	14	116,000	Lung	13	108,210
Colorectal	8	71,830	Colorectal	8	65,000
Bladder	7	56,390	Endometrial	6	52,630
Melanoma	5	43,890	Thyroid	6	47,790
Kidney	4	39,140	Lymphoma	4	32,530
Lymphoma	4	38,270	Melanoma	4	32,210
Oral cavity	4	30,220	Kidney	3	24,780
Leukemia	4	30,100	Pancreas	3	22,890
Liver	3	24,600	Leukemia	3	22,280
All others	20	171,780	All others	21	169,330
All sites	100	855,220	All sites	100	810,320
Cancer Deaths					
Lung	28	86,930	Lung	26	72,330
Prostate	10	29,480	Breast	15	40,000
Colorectal	8	26,270	Colorectal	9	24,040
Pancreas	7	20,170	Pancreas	7	19,420
Liver	5	15,870	Ovary	5	14,270
Leukemia	5	14,040	Leukemia	4	10,050
Esophagus	4	12,450	Endometrial	3	8,590
Bladder	4	11,170	Lymphoma	3	8,520
Lymphoma	3	10,470	Liver	3	7,130
Kidney	3	8,900	CNS	2	6,230
All others	23	74,260	All others	23	65,130
All sites	100	310,010	All sites	100	275,710

Source: From R Siegel et al: Cancer statistics, 2014. *CA Cancer J Clin* 64:9, 2014.

Cancer is the second leading cause of death behind heart disease. Deaths from heart disease have declined 45% in the United States since 1950 and continue to decline. Cancer has overtaken heart disease as the number one cause of death in persons younger than age 85 years. Incidence trends over time are shown in [Fig. 99-1](#). After a 70-year period of increase, cancer deaths began to decline in 1990–1991 ([Fig. 99-2](#)). Between 1990 and 2010, cancer deaths decreased by 21% among men and 12.3% among women. The magnitude of the decline is illustrated in [Fig. 99-3](#). The five leading causes of cancer deaths are shown for various populations in [Table 99-2](#). The 5-year survival for white patients was 39% in 1960–1963 and 69% in 2003–2009. Cancers are more often deadly in blacks; the 5-year survival was 61% for the 2003–2009 interval; however, the racial differences are narrowing over time. Incidence and mortality vary among racial and ethnic groups ([Table 99-3](#)). The basis for these differences is unclear.

CANCER AROUND THE WORLD

 In 2008, 12.7 million new cancer cases and 7.6 million cancer deaths were estimated worldwide, according to estimates of GLOBOCAN 2008, developed by the International Agency for Research on Cancer (IARC). When broken down by region of the world, ~45% of cases were in Asia, 26% in Europe, 14.5% in North America, 7.1% in Central/South America, 6% in Africa, and 1% in Australia/New Zealand ([Fig. 99-4](#)). Lung cancer is the most common cancer and the most common cause of cancer death in the world. Its

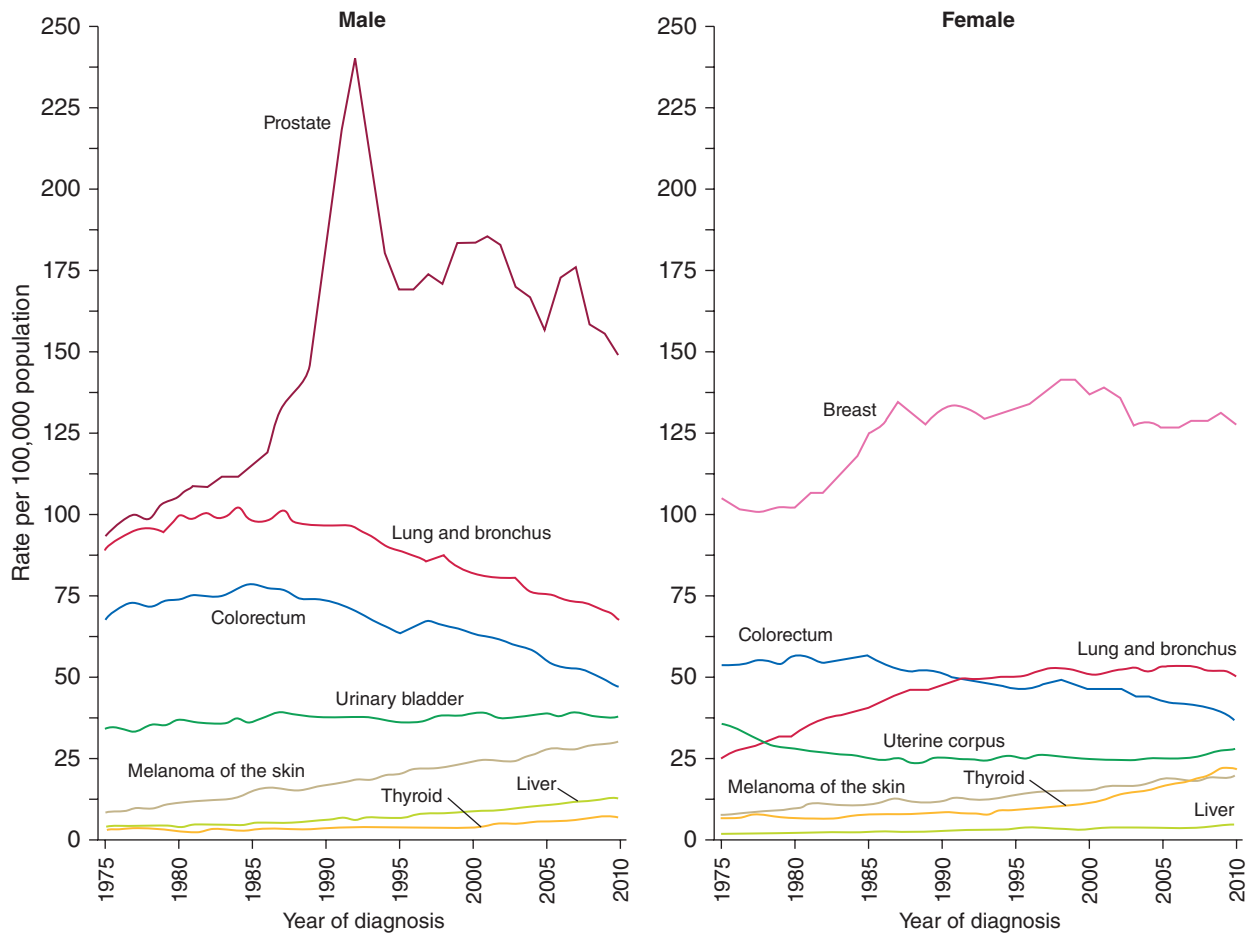


FIGURE 99-1 Incidence rates for particular types of cancer over the last 35 years in men (A) and women (B). (From R Siegel et al: *CA Cancer J Clin* 64:9, 2014.)

incidence is highly variable, affecting only 2 per 100,000 African women but as many as 61 per 100,000 North American men. Breast cancer is the second most common cancer worldwide; however, it ranks fifth as a cause of death behind lung, stomach, liver, and colorectal cancer. Among the eight most common forms of cancer, lung (2-fold), breast (3-fold), prostate (2.5-fold), and colorectal (3-fold) cancers are more common in more developed countries than in less developed countries. By contrast, liver (2-fold), cervical (2-fold), and esophageal (2- to 3-fold) cancers are more common in less developed countries. Stomach cancer incidence is similar in more and less developed countries but is much more common in Asia than North America or Africa. The most common cancers in Africa are cervical, breast, and liver cancers. It has been estimated that nine modifiable risk factors are responsible for more than one-third of cancers worldwide. These include smoking, alcohol consumption, obesity, physical inactivity, low fruit and vegetable consumption, unsafe sex, air pollution, indoor smoke from household fuels, and contaminated injections.

PATIENT MANAGEMENT

Important information is obtained from every portion of the routine history and physical examination. The duration of symptoms may reveal the chronicity of disease. The past medical history may alert the physician to the presence of underlying diseases that may affect the choice of therapy or the side effects of treatment. The social history may reveal occupational exposure to carcinogens or habits, such as smoking or alcohol consumption, that may influence the course of disease and its treatment. The family history may suggest an underlying familial cancer predisposition and point out the need to begin surveillance or other preventive therapy for unaffected siblings of the

patient. The review of systems may suggest early symptoms of metastatic disease or a paraneoplastic syndrome.

DIAGNOSIS

The diagnosis of cancer relies most heavily on invasive tissue biopsy and should never be made without obtaining tissue; no noninvasive diagnostic test is sufficient to define a disease process as cancer. Although in rare clinical settings (e.g., thyroid nodules), fine-needle aspiration is an acceptable diagnostic procedure, the diagnosis generally depends on obtaining adequate tissue to permit careful evaluation of the histology of the tumor, its grade, and its invasiveness and to yield further molecular diagnostic information, such as the expression of cell-surface markers or intracellular proteins that typify a particular cancer, or the presence of a molecular marker, such as the t(8;14) translocation of Burkitt's lymphoma. Increasing evidence links the expression of certain genes with the prognosis and response to therapy (Chaps. 101e and 102e).

Occasionally a patient will present with a metastatic disease process that is defined as cancer on biopsy but has no apparent primary site of disease. Efforts should be made to define the primary site based on age, sex, sites of involvement, histology and tumor markers, and personal and family history. Particular attention should be focused on ruling out the most treatable causes (Chap. 120e).

Once the diagnosis of cancer is made, the management of the patient is best undertaken as a multidisciplinary collaboration among the primary care physician, medical oncologists, surgical oncologists, radiation oncologists, oncology nurse specialists, pharmacists, social workers, rehabilitation medicine specialists, and a number of other consulting professionals working closely with each other and with the patient and family.

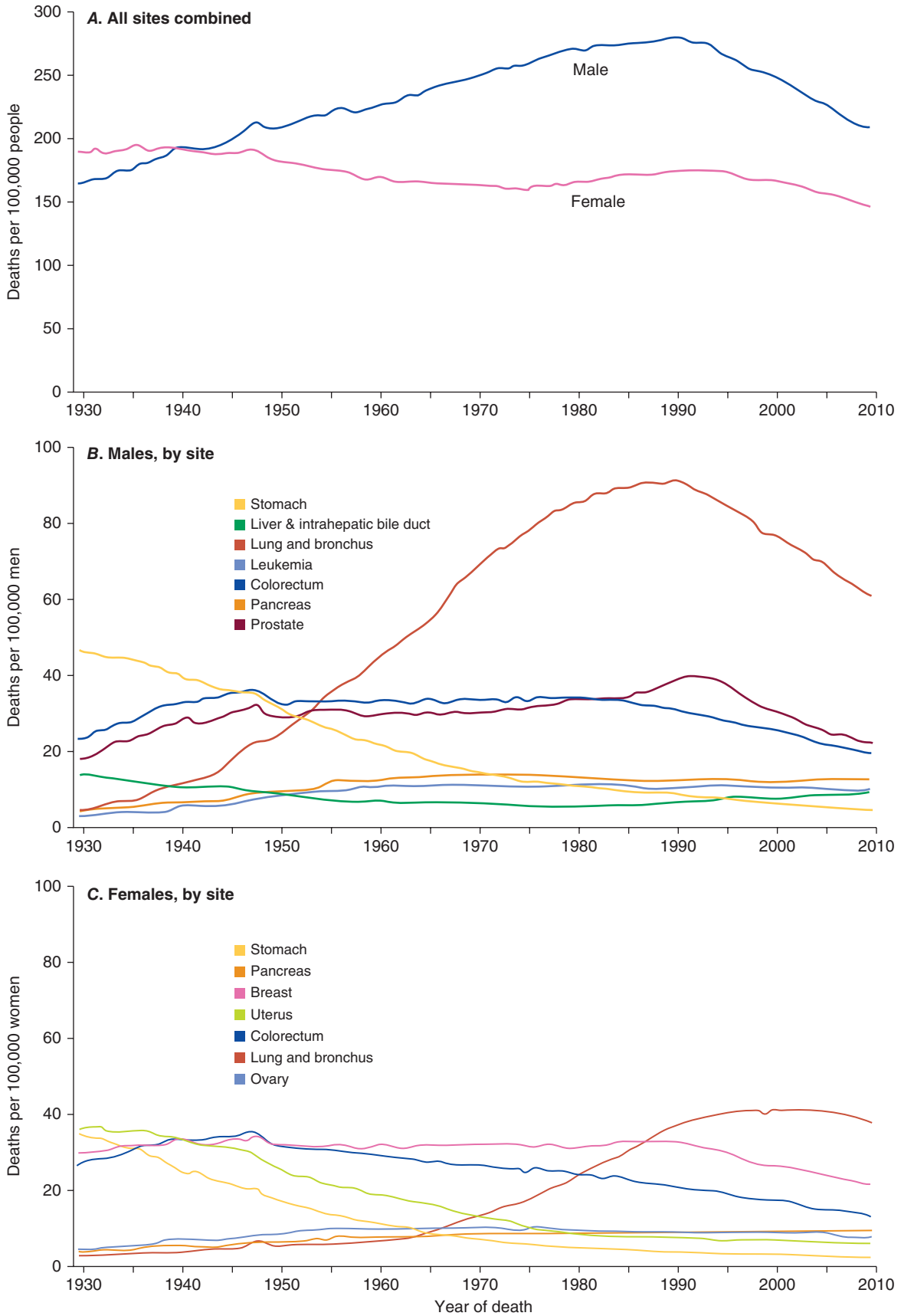


FIGURE 99-2 Eighty-year trend in cancer death rates for (A) women and (B) men by site in the United States, 1930–2010. Rates are per 100,000 age-adjusted to the 2000 U.S. standard population. All sites combined (A), individual sites in men (B) and individual sites in women (C) are shown. (From R Siegel et al: *CA Cancer J Clin* 64:9, 2014.)

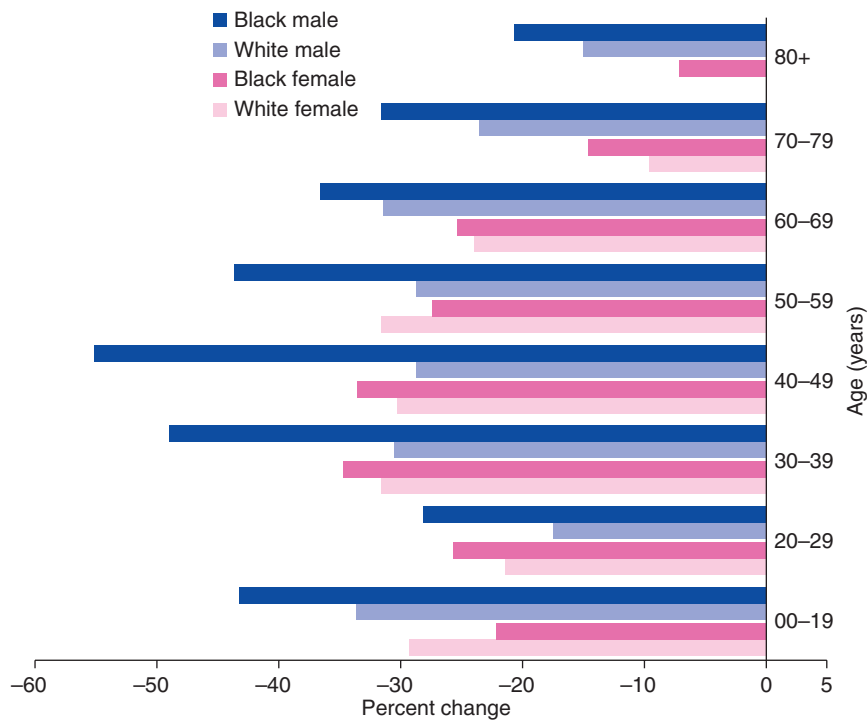


FIGURE 99-3 The decline in death rates from cancer is shown for different age ranges by sex and race for the 20-year period between 1991 and 2010 expressed as a percentage of the 1991 rate. (From R Siegel et al: *CA Cancer J Clin* 64:9, 2014.)

DEFINING THE EXTENT OF DISEASE AND THE PROGNOSIS

The first priority in patient management after the diagnosis of cancer is established and shared with the patient is to determine the extent of disease. The curability of a tumor usually is inversely proportional to the tumor burden. Ideally, the tumor will be diagnosed before symptoms develop or as a consequence of screening efforts (**Chap. 100**). A very high proportion of such patients can be cured. However, most patients with cancer present with symptoms related to the cancer, caused either by mass effects of the tumor or by alterations associated with the production of cytokines or hormones by the tumor.

For most cancers, the extent of disease is evaluated by a variety of noninvasive and invasive diagnostic tests and procedures. This process is called *staging*. There are two types. *Clinical staging* is based on physical examination, radiographs, isotopic scans, computed tomography (CT) scans, and other imaging procedures; *pathologic staging* takes into account information obtained during a surgical procedure, which might include intraoperative palpation, resection of regional lymph nodes and/or tissue adjacent to the tumor, and inspection and biopsy of organs commonly involved in disease spread. Pathologic staging includes

histologic examination of all tissues removed during the surgical procedure. Surgical procedures performed may include a simple lymph node biopsy or more extensive procedures such as thoracotomy, mediastinoscopy, or laparotomy. Surgical staging may occur in a separate procedure or may be done at the time of definitive surgical resection of the primary tumor.

Knowledge of the predilection of particular tumors for spreading to adjacent or distant organs helps direct the staging evaluation.

Information obtained from staging is used to define the extent of disease as localized, as exhibiting spread outside of the organ of origin to regional but not distant sites, or as metastatic to distant sites. The most widely used system of staging is the TNM (tumor, node, metastasis) system codified by the International Union Against Cancer and the American Joint Committee on Cancer. The TNM classification is an anatomically based system that categorizes the tumor on the basis of the size of the primary tumor lesion (T1-4, where a higher number indicates a tumor of larger size), the presence of nodal involvement (usually N0 and N1 for the absence and presence, respectively, of involved nodes, although some tumors have more elaborate systems of nodal grading), and the presence of metastatic disease (M0 and M1 for the absence and presence, respectively, of metastases). The various permutations of T, N, and M scores (sometimes including tumor histologic grade [G]) are then broken into stages, usually designated by the roman numerals I through IV. Tumor

burden increases and curability decreases with increasing stage. Other anatomic staging systems are used for some tumors, e.g., the Dukes classification for colorectal cancers, the International Federation of Gynecologists and Obstetricians classification for gynecologic cancers, and the Ann Arbor classification for Hodgkin's disease.

Certain tumors cannot be grouped on the basis of anatomic considerations. For example, hematopoietic tumors such as leukemia, myeloma, and lymphoma are often disseminated at presentation and do not spread like solid tumors. For these tumors, other prognostic factors have been identified (**Chaps. 132-136**).

In addition to tumor burden, a second major determinant of treatment outcome is the physiologic reserve of the patient. Patients who are bedridden before developing cancer are likely to fare worse, stage for stage, than fully active patients. Physiologic reserve is a determinant of how a patient is likely to cope with the physiologic stresses imposed by the cancer and its treatment. This factor is difficult to assess directly. Instead, surrogate markers for physiologic reserve are used, such as the patient's age or Karnofsky performance status (**Table 99-4**) or Eastern Cooperative Oncology Group (ECOG) performance status

TABLE 99-2 THE FIVE LEADING PRIMARY TUMOR SITES FOR PATIENTS DYING OF CANCER BASED ON AGE AND SEX IN 2010

Rank	Sex	All Ages	Age, years				
			Under 20	20-39	40-59	60-79	>80
1	M	Lung	Leukemia	Leukemia	Lung	Lung	Lung
	F	Lung	Leukemia	Breast	Breast	Lung	Lung
2	M	Prostate	CNS	CNS	Colorectal	Colorectal	Prostate
	F	Breast	CNS	Cervix	Lung	Breast	Breast
3	M	Colorectal	Bone sarcoma	Colorectal	Liver	Prostate	Colorectal
	F	Colorectal	Bone sarcoma	Leukemia	Colorectal	Colorectal	Colorectal
4	M	Pancreas	Soft tissue sarcoma	Lymphoma	Pancreas	Pancreas	Bladder
	F	Pancreas	Soft tissue sarcoma	Colorectal	Ovary	Pancreas	Pancreas
5	M	Liver	Lymphoma	Lung	Esophagus	Liver	Pancreas
	F	Ovary	Liver	CNS	Pancreas	Ovary	Lymphoma

Abbreviations: CNS, central nervous system; F, female; M, male.

Source: From R Siegel et al: Cancer statistics, 2014. *CA Cancer J Clin* 64:9, 2014.

TABLE 99-3 CANCER INCIDENCE AND MORTALITY IN RACIAL AND ETHNIC GROUPS, UNITED STATES, 2006–2010

Site	Sex	White	Black	Asian/Pacific Islander	American Indian ^a	Hispanic
Incidence per 100,000 Population						
All	M	548.1	601.0	326.1	441.1	426.8
	F	436.2	395.9	282.6	372.0	330.8
Breast		127.3	118.4	84.7	90.3	91.1
Colorectal	M	50.9	62.5	40.8	51.7	47.3
	F	38.6	46.7	31.0	42.7	32.6
Kidney	M	21.6	23.0	10.6	30.6	20.5
	F	11.2	12.2	5.1	17.5	11.5
Liver	M	8.7	14.9	21.3	17.8	11.5
	F	2.9	4.4	8.0	8.0	6.9
Lung	M	82.9	94.7	48.8	70.2	45.9
	F	57.1	50.7	27.6	41.3	26.5
Prostate		138.6	220.0	75.0	104.1	124.2
Cervix		7.2	10.3	6.7	9.7	10.9
Deaths per 100,000 Population						
All	M	217.3	276.6	132.4	191.0	152.2
	F	153.6	171.2	92.1	139.0	101.3
Breast		22.7	30.8	11.5	15.5	14.8
Colorectal	M	19.2	28.7	13.1	18.7	16.1
	F	13.6	19.0	9.7	15.4	10.2
Kidney	M	5.9	5.7	3.0	9.5	5.1
	F	2.6	2.6	1.2	4.4	2.3
Liver	M	7.1	11.8	14.4	13.2	12.3
	F	2.9	4.1	6.0	6.1	5.4
Lung	M	65.7	78.5	35.5	49.6	31.3
	F	42.7	37.2	18.4	33.1	14.1
Prostate		21.3	50.9	10.1	20.7	19.2
Cervix		2.1	4.2	1.9	3.5	2.9

^aBased on Indian Health Service delivery areas.

Abbreviations: F, female; M, male.

Source: From R Siegel R et al: Cancer statistics, 2014. *CA Cancer J Clin* 64:9, 2014.

(Table 99-5). Older patients and those with a Karnofsky performance status <70 or ECOG performance status ≥ 3 have a poor prognosis unless the poor performance is a reversible consequence of the tumor.

Increasingly, biologic features of the tumor are being related to prognosis. The expression of particular oncogenes, drug-resistance genes, apoptosis-related genes, and genes involved in metastasis is being found to influence response to therapy and prognosis. The presence of selected cytogenetic abnormalities may influence survival. Tumors with higher growth fractions, as assessed by expression of proliferation-related markers such as proliferating cell nuclear antigen, behave more aggressively than tumors with lower growth fractions. Information obtained from studying the tumor itself will increasingly be used to influence treatment decisions. Host genes involved in drug metabolism can influence the safety and efficacy of particular treatments.

Enormous heterogeneity has been noted by studying tumors; we have learned that morphology is not capable of discerning certain distinct subsets of patients whose tumors have different sets of abnormalities. Tumors that look the same by light microscopy can be very different. Similarly, tumors that look quite different from one another histologically can share genetic lesions that predict responses to treatments. Furthermore, tumor cells vary enormously within a single patient even though the cells share a common origin.

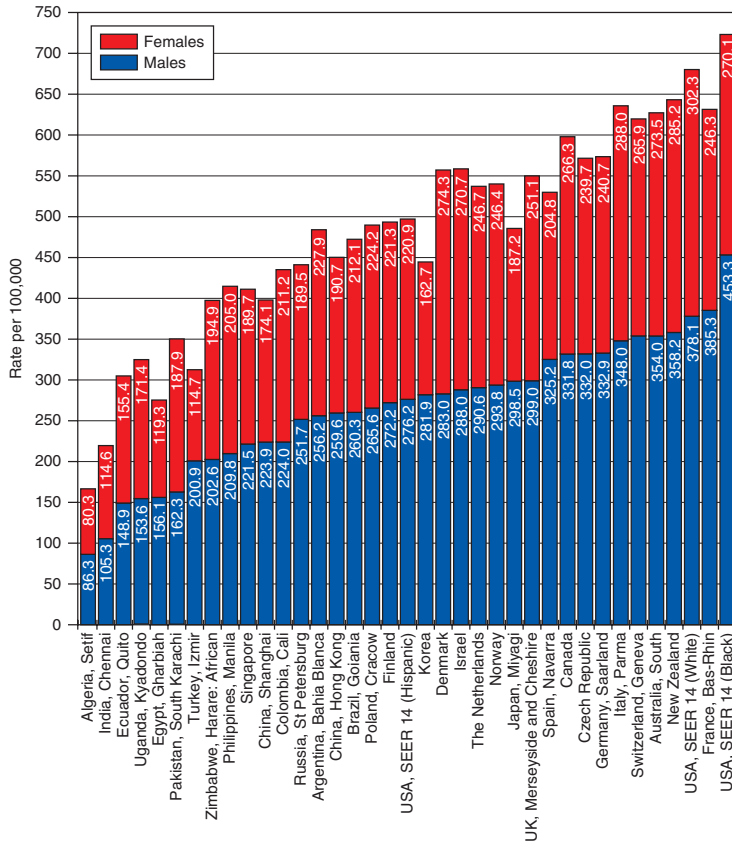
MAKING A TREATMENT PLAN

From information on the extent of disease and the prognosis and in conjunction with the patient's wishes, it is determined whether the treatment approach should be curative or palliative in intent. Cooperation among the various professionals involved in cancer

treatment is of the utmost importance in treatment planning. For some cancers, chemotherapy or chemotherapy plus radiation therapy delivered before the use of definitive surgical treatment (so-called neoadjuvant therapy) may improve the outcome, as seems to be the case for locally advanced breast cancer and head and neck cancers. In certain settings in which combined-modality therapy is intended, coordination among the medical oncologist, radiation oncologist, and surgeon is crucial to achieving optimal results. Sometimes the chemotherapy and radiation therapy need to be delivered sequentially, and other times concurrently. Surgical procedures may precede or follow other treatment approaches. It is best for the treatment plan either to follow a standard protocol precisely or else to be part of an ongoing clinical research protocol evaluating new treatments. Ad hoc modifications of standard protocols are likely to compromise treatment results.

The choice of treatment approaches was formerly dominated by the local culture in both the university and the practice settings. However, it is now possible to gain access electronically to standard treatment protocols and to every approved clinical research study in North America through a personal computer interface with the Internet.¹

¹The National Cancer Institute maintains a database called PDQ (Physician Data Query) that is accessible on the Internet under the name CancerNet at www.cancer.gov/cancertopics/pdq/cancerdatabase. Information can be obtained through a facsimile machine using CancerFax by dialing 301-402-5874. Patient information is also provided by the National Cancer Institute in at least three formats: on the Internet via CancerNet at www.cancer.gov, through the CancerFax number listed above, or by calling 1-800-4-CANCER. The quality control for the information provided through these services is rigorous.



Incidence (n = 10,864,499) Mortality (n = 6,724,931) Prevalence (n = 24,576,453)

FIGURE 99-4 Worldwide overall annual cancer incidence, mortality, and 5-year prevalence for the period of 1993–2001. (Adapted from A Jemal et al: *Cancer Epidemiol Biomarkers Prev* 19:1893, 2010.)

The skilled physician also has much to offer the patient for whom curative therapy is no longer an option. Often a combination of guilt and frustration over the inability to cure the patient and the pressure of a busy schedule greatly limit the time a physician spends with a patient who is receiving only palliative care. Resist these forces. In addition to the medicines administered to alleviate symptoms (see below), it is important to remember the comfort that is provided by holding the patient's hand, continuing regular examinations, and taking time to talk.

TABLE 99-4 KARNOFSKY PERFORMANCE INDEX	
Performance Status	Functional Capability of the Patient
100	Normal; no complaints; no evidence of disease
90	Able to carry on normal activity; minor signs or symptoms of disease
80	Normal activity with effort; some signs or symptoms of disease
70	Cares for self; unable to carry on normal activity or do active work
60	Requires occasional assistance but is able to care for most needs
50	Requires considerable assistance and frequent medical care
40	Disabled; requires special care and assistance
30	Severely disabled; hospitalization is indicated, although death is not imminent
20	Very sick; hospitalization is necessary; active supportive treatment is necessary
10	Moribund, fatal processes progressing rapidly
0	Dead

MANAGEMENT OF DISEASE AND TREATMENT COMPLICATIONS

Because cancer therapies are toxic (Chap. 103e), patient management involves addressing complications of both the disease and its treatment as well as the complex psychosocial problems associated with cancer. In the short term during a course of curative therapy, the patient's functional status may decline. Treatment-induced toxicity is less acceptable if the goal of therapy is palliation. The most common side effects of treatment are nausea and vomiting (see below), febrile neutropenia (Chap. 104), and myelosuppression (Chap. 103e). Tools are now available to minimize the acute toxicity of cancer treatment.

New symptoms developing in the course of cancer treatment should always be assumed to be reversible until proven otherwise. The fatalistic attribution of anorexia, weight loss, and jaundice to recurrent or progressive tumor could result in a patient dying from a reversible intercurrent cholecystitis. Intestinal obstruction may be due to reversible adhesions rather than progressive tumor. Systemic infections, sometimes with unusual pathogens, may be a consequence of the immunosuppression associated with cancer therapy. Some drugs used to treat cancer or its complications (e.g., nausea) may produce central nervous system symptoms that look like metastatic disease or may mimic paraneoplastic syndromes such as the syndrome of inappropriate antidiuretic hormone. A definitive diagnosis should be pursued and may even require a repeat biopsy.

A critical component of cancer management is assessing the response to treatment. In addition to a careful physical examination in which all sites of disease are physically measured and recorded in a flow chart by date, response assessment usually requires periodic repeating of imaging tests that were abnormal at the time of staging. If imaging tests have become normal, repeat biopsy of previously involved tissue is performed to document complete response by pathologic criteria. Biopsies are not usually required if there is macroscopic residual disease. A complete response is defined as disappearance of all evidence of disease, and a partial response as >50% reduction in the sum of the products of the perpendicular diameters of all measurable lesions.

The determination of partial response may also be based on a 30% decrease in the sums of the longest diameters of lesions (Response Evaluation Criteria in Solid Tumors [RECIST]). Progressive disease is defined as the appearance of any new lesion or an increase of >25% in the sum of the products of the perpendicular diameters of all measurable lesions (or an increase of 20% in the sums of the longest diameters by RECIST). Tumor shrinkage or growth that does not meet any of these criteria is considered stable disease. Some sites of involvement (e.g., bone) or patterns of involvement (e.g., lymphangitic lung or diffuse pulmonary infiltrates) are considered unmeasurable. No response is complete without biopsy documentation of their resolution, but partial responses may exclude their assessment unless clear objective progression has occurred.

TABLE 99-5 THE EASTERN COOPERATIVE ONCOLOGY GROUP (ECOG) PERFORMANCE SCALE	
ECOG Grade 0:	Fully active, able to carry on all predisease performance without restriction
ECOG Grade 1:	Restricted in physically strenuous activity but ambulatory and able to carry out work of a light or sedentary nature, e.g., light housework, office work
ECOG Grade 2:	Ambulatory and capable of all self-care but unable to carry out any work activities. Up and about more than 50% of waking hours
ECOG Grade 3:	Capable of only limited self-care, confined to bed or chair more than 50% of waking hours
ECOG Grade 4:	Completely disabled. Cannot carry on any self-care. Totally confined to bed or chair
ECOG Grade 5:	Dead

Source: From MM Oken et al: *Am J Clin Oncol* 5:649, 1982.

TABLE 99-6 TUMOR MARKERS

Tumor Markers	Cancer	Nonneoplastic Conditions
Hormones		
Human chorionic gonadotropin	Gestational trophoblastic disease, gonadal germ cell tumor	Pregnancy
Calcitonin	Medullary cancer of the thyroid	
Catecholamines	Pheochromocytoma	
Oncofetal Antigens		
α Fetoprotein	Hepatocellular carcinoma, gonadal germ cell tumor	Cirrhosis, hepatitis
Carcinoembryonic antigen	Adenocarcinomas of the colon, pancreas, lung, breast, ovary	Pancreatitis, hepatitis, inflammatory bowel disease, smoking
Enzymes		
Prostatic acid phosphatase	Prostate cancer	Prostatitis, prostatic hypertrophy
Neuron-specific enolase	Small-cell cancer of the lung, neuroblastoma	
Lactate dehydrogenase	Lymphoma, Ewing's sarcoma	Hepatitis, hemolytic anemia, many others
Tumor-Associated Proteins		
Prostate-specific antigen	Prostate cancer	Prostatitis, prostatic hypertrophy
Monoclonal immunoglobulin	Myeloma	Infection, MGUS
CA-125	Ovarian cancer, some lymphomas	Menstruation, peritonitis, pregnancy
CA 19-9	Colon, pancreatic, breast cancer	Pancreatitis, ulcerative colitis
CD30	Hodgkin's disease, anaplastic large-cell lymphoma	—
CD25	Hairy cell leukemia, adult T cell leukemia/lymphoma	—

Abbreviation: MGUS, monoclonal gammopathy of uncertain significance.

Tumor markers may be useful in patient management in certain tumors. Response to therapy may be difficult to gauge with certainty. However, some tumors produce or elicit the production of markers that can be measured in the serum or urine, and in a particular patient, rising and falling levels of the marker are usually associated with increasing or decreasing tumor burden, respectively. Some clinically useful tumor markers are shown in Table 99-6. Tumor markers are not in themselves specific enough to permit a diagnosis of malignancy to be made, but once a malignancy has been diagnosed and shown to be associated with elevated levels of a tumor marker, the marker can be used to assess response to treatment.

The recognition and treatment of depression are important components of management. The incidence of depression in cancer patients is ~25% overall and may be greater in patients with greater debility. This diagnosis is likely in a patient with a depressed mood (dysphoria) and/or a loss of interest in pleasure (anhedonia) for at least 2 weeks. In addition, three or more of the following symptoms are usually present: appetite change, sleep problems, psychomotor retardation or agitation, fatigue, feelings of guilt or worthlessness, inability to concentrate, and suicidal ideation. Patients with these symptoms should receive therapy. Medical therapy with a serotonin reuptake inhibitor such as fluoxetine (10–20 mg/d), sertraline (50–150 mg/d), or paroxetine (10–20 mg/d) or a tricyclic antidepressant such as amitriptyline (50–100 mg/d) or desipramine (75–150 mg/d) should be tried, allowing 4–6 weeks for response. Effective therapy should be continued at least 6 months after resolution of symptoms. If therapy is unsuccessful, other classes of antidepressants may be used. In addition to medication, psychosocial

interventions such as support groups, psychotherapy, and guided imagery may be of benefit.

Many patients opt for unproven or unsound approaches to treatment when it appears that conventional medicine is unlikely to be curative. Those seeking such alternatives are often well educated and may be early in the course of their disease. Unsound approaches are usually hawked on the basis of unsubstantiated anecdotes and not only cannot help the patient but may be harmful. Physicians should strive to keep communications open and nonjudgmental, so that patients are more likely to discuss with the physician what they are actually doing. The appearance of unexpected toxicity may be an indication that a supplemental therapy is being taken.²

LONG-TERM FOLLOW-UP/LATE COMPLICATIONS

At the completion of treatment, sites originally involved with tumor are reassessed, usually by radiography or imaging techniques, and any persistent abnormality is biopsied. If disease persists, the multidisciplinary team discusses a new salvage treatment plan. If the patient has been rendered disease-free by the original treatment, the patient is followed regularly for disease recurrence. The optimal guidelines for follow-up care are not known. For many years, a routine practice has been to follow the patient monthly for 6–12 months, then every other month for a year, every 3 months for a year, every 4 months for a year, every 6 months for a year, and then annually. At each visit, a battery of laboratory and radiographic and imaging tests were obtained on the assumption that it is best to detect recurrent disease before it becomes symptomatic. However, where follow-up procedures have been examined, this assumption has been found to be untrue. Studies of breast cancer, melanoma, lung cancer, colon cancer, and lymphoma have all failed to support the notion that asymptomatic relapses are more readily cured by salvage therapy than symptomatic relapses. In view of the enormous cost of a full battery of diagnostic tests and their manifest lack of impact on survival, new guidelines are emerging for less frequent follow-up visits, during which the history and physical examination are the major investigations performed.

As time passes, the likelihood of recurrence of the primary cancer diminishes. For many types of cancer, survival for 5 years without recurrence is tantamount to cure. However, important medical problems can occur in patients treated for cancer and must be examined (Chap. 125). Some problems emerge as a consequence of the disease and some as a consequence of the treatment. An understanding of these disease- and treatment-related problems may help in their detection and management.

Despite these concerns, most patients who are cured of cancer return to normal lives.

SUPPORTIVE CARE

In many ways, the success of cancer therapy depends on the success of the supportive care. Failure to control the symptoms of cancer and its treatment may lead patients to abandon curative therapy. Of equal importance, supportive care is a major determinant of quality of life. Even when life cannot be prolonged, the physician must strive to preserve its quality. Quality-of-life measurements have become common endpoints of clinical research studies. Furthermore, palliative care has been shown to be cost-effective when approached in an organized fashion. A credo for oncology could be to cure sometimes, to extend life often, and to comfort always.

Pain Pain occurs with variable frequency in the cancer patient: 25–50% of patients present with pain at diagnosis, 33% have pain associated with treatment, and 75% have pain with progressive disease. The pain may have several causes. In ~70% of cases, pain is caused by the tumor itself—by invasion of bone, nerves, blood vessels, or mucous

²Information about unsound methods may be obtained from the National Council Against Health Fraud, Box 1276, Loma Linda, CA 92354, or from the Center for Medical Consumers and Health Care Information, 237 Thompson Street, New York, NY 10012.

474 membranes or obstruction of a hollow viscus or duct. In ~20% of cases, pain is related to a surgical or invasive medical procedure, to radiation injury (mucositis, enteritis, or plexus or spinal cord injury), or to chemotherapy injury (mucositis, peripheral neuropathy, phlebitis, steroid-induced aseptic necrosis of the femoral head). In 10% of cases, pain is unrelated to cancer or its treatment.

Assessment of pain requires the methodical investigation of the history of the pain, its location, character, temporal features, provocative and palliative factors, and intensity (**Chap. 18**); a review of the oncologic history and past medical history as well as personal and social history; and a thorough physical examination. The patient should be given a 10-division visual analogue scale on which to indicate the severity of the pain. The clinical condition is often dynamic, making it necessary to reassess the patient frequently. Pain therapy should not be withheld while the cause of pain is being sought.

A variety of tools are available with which to address cancer pain. About 85% of patients will have pain relief from pharmacologic intervention. However, other modalities, including antitumor therapy (such as surgical relief of obstruction, radiation therapy, and strontium-89 or samarium-153 treatment for bone pain), neurostimulatory techniques, regional analgesia, or neuroablative procedures, are effective in an additional 12% or so. Thus, very few patients will have inadequate pain relief if appropriate measures are taken. **A specific approach to pain relief is detailed in Chap. 10.**

Nausea Emesis in the cancer patient is usually caused by chemotherapy (**Chap. 103e**). Its severity can be predicted from the drugs used to treat the cancer. Three forms of emesis are recognized on the basis of their timing with regard to the noxious insult. *Acute emesis*, the most common variety, occurs within 24 h of treatment. *Delayed emesis* occurs 1–7 days after treatment; it is rare, but, when present, usually follows cisplatin administration. *Anticipatory emesis* occurs before administration of chemotherapy and represents a conditioned response to visual and olfactory stimuli previously associated with chemotherapy delivery.

Acute emesis is the best understood form. Stimuli that activate signals in the chemoreceptor trigger zone in the medulla, the cerebral cortex, and peripherally in the intestinal tract lead to stimulation of the vomiting center in the medulla, the motor center responsible for coordinating the secretory and muscle contraction activity that leads to emesis. Diverse receptor types participate in the process, including dopamine, serotonin, histamine, opioid, and acetylcholine receptors. The serotonin receptor antagonists ondansetron and granisetron are the most effective drugs against highly emetogenic agents, but they are expensive.

As with the analgesia ladder, emesis therapy should be tailored to the situation. For mildly and moderately emetogenic agents, prochlorperazine, 5–10 mg PO or 25 mg PR, is effective. Its efficacy may be enhanced by administering the drug before the chemotherapy is delivered. Dexamethasone, 10–20 mg IV, is also effective and may enhance the efficacy of prochlorperazine. For highly emetogenic agents such as cisplatin, mechlorethamine, dacarbazine, and streptozocin, combinations of agents work best and administration should begin 6–24 h before treatment. Ondansetron, 8 mg PO every 6 h the day before therapy and IV on the day of therapy, plus dexamethasone, 20 mg IV before treatment, is an effective regimen. Addition of oral aprepitant (a substance P/neurokinin 1 receptor antagonist) to this regimen (125 mg on day 1, 80 mg on days 2 and 3) further decreases the risk of both acute and delayed vomiting. Like pain, emesis is easier to prevent than to alleviate.

Delayed emesis may be related to bowel inflammation from the therapy and can be controlled with oral dexamethasone and oral metoclopramide, a dopamine receptor antagonist that also blocks serotonin receptors at high dosages. The best strategy for preventing anticipatory emesis is to control emesis in the early cycles of therapy to prevent the conditioning from taking place. If this is unsuccessful, prophylactic antiemetics the day before treatment may help. Experimental studies are evaluating behavior modification.

Effusions Fluid may accumulate abnormally in the pleural cavity, pericardium, or peritoneum. Asymptomatic malignant effusions may

not require treatment. Symptomatic effusions occurring in tumors responsive to systemic therapy usually do not require local treatment but respond to the treatment for the underlying tumor. Symptomatic effusions occurring in tumors unresponsive to systemic therapy may require local treatment in patients with a life expectancy of at least 6 months.

Pleural effusions due to tumors may or may not contain malignant cells. Lung cancer, breast cancer, and lymphomas account for ~75% of malignant pleural effusions. Their exudative nature is usually gauged by an effusion/serum protein ratio of ≥ 0.5 or an effusion/serum lactate dehydrogenase ratio of ≥ 0.6 . When the condition is symptomatic, thoracentesis is usually performed first. In most cases, symptomatic improvement occurs for < 1 month. Chest tube drainage is required if symptoms recur within 2 weeks. Fluid is aspirated until the flow rate is < 100 mL in 24 h. Then either 60 units of bleomycin or 1 g of doxycycline is infused into the chest tube in 50 mL of 5% dextrose in water; the tube is clamped; the patient is rotated on four sides, spending 15 min in each position; and, after 1–2 h, the tube is again attached to suction for another 24 h. The tube is then disconnected from suction and allowed to drain by gravity. If < 100 mL drains over the next 24 h, the chest tube is pulled, and a radiograph is taken 24 h later. If the chest tube continues to drain fluid at an unacceptably high rate, sclerosis can be repeated. Bleomycin may be somewhat more effective than doxycycline but is very expensive. Doxycycline is usually the drug of first choice. If neither doxycycline nor bleomycin is effective, talc can be used.

Symptomatic pericardial effusions are usually treated by creating a pericardial window or by stripping the pericardium. If the patient's condition does not permit a surgical procedure, sclerosis can be attempted with doxycycline and/or bleomycin.

Malignant ascites is usually treated with repeated paracentesis of small volumes of fluid. If the underlying malignancy is unresponsive to systemic therapy, peritoneovenous shunts may be inserted. Despite the fear of disseminating tumor cells into the circulation, widespread metastases are an unusual complication. The major complications are occlusion, leakage, and fluid overload. Patients with severe liver disease may develop disseminated intravascular coagulation.

Nutrition Cancer and its treatment may lead to a decrease in nutrient intake of sufficient magnitude to cause weight loss and alteration of intermediary metabolism. The prevalence of this problem is difficult to estimate because of variations in the definition of cancer cachexia, but most patients with advanced cancer experience weight loss and decreased appetite. A variety of both tumor-derived factors (e.g., bombesin, adrenocorticotrophic hormone) and host-derived factors (e.g., tumor necrosis factor, interleukins 1 and 6, growth hormone) contribute to the altered metabolism, and a vicious cycle is established in which protein catabolism, glucose intolerance, and lipolysis cannot be reversed by the provision of calories.

It remains controversial how to assess nutritional status and when and how to intervene. Efforts to make the assessment objective have included the use of a prognostic nutritional index based on albumin levels, triceps skinfold thickness, transferrin levels, and delayed-type hypersensitivity skin testing. However, a simpler approach has been to define the threshold for nutritional intervention as $< 10\%$ unexplained body weight loss, serum transferrin level < 1500 mg/L (150 mg/dL), and serum albumin < 34 g/L (3.4 g/dL).

The decision is important, because it appears that cancer therapy is substantially more toxic and less effective in the face of malnutrition. Nevertheless, it remains unclear whether nutritional intervention can alter the natural history. Unless some pathology is affecting the absorptive function of the gastrointestinal tract, enteral nutrition provided orally or by tube feeding is preferred over parenteral supplementation. However, the risks associated with the tube may outweigh the benefits. Megestrol acetate, a progestational agent, has been advocated as a pharmacologic intervention to improve nutritional status. Research in this area may provide more tools in the future as cytokine-mediated mechanisms are further elucidated.

Psychosocial Support The psychosocial needs of patients vary with their situation. Patients undergoing treatment experience fear, anxiety,

and depression. Self-image is often seriously compromised by deforming surgery and loss of hair. Women who receive cosmetic advice that enables them to look better also feel better. Loss of control over how one spends time can contribute to the sense of vulnerability. Juggling the demands of work and family with the demands of treatment may create enormous stresses. Sexual dysfunction is highly prevalent and needs to be discussed openly with the patient. An empathetic health care team is sensitive to the individual patient's needs and permits negotiation where such flexibility will not adversely affect the course of treatment.

Cancer survivors have other sets of difficulties. Patients may have fears associated with the termination of a treatment they associate with their continued survival. Adjustments are required to physical losses and handicaps, real and perceived. Patients may be preoccupied with minor physical problems. They perceive a decline in their job mobility and view themselves as less desirable workers. They may be victims of job and/or insurance discrimination. Patients may experience difficulty reentering their normal past life. They may feel guilty for having survived and may carry a sense of vulnerability to colds and other illnesses. Perhaps the most pervasive and threatening concern is the ever-present fear of relapse (the Damocles syndrome).

Patients in whom therapy has been unsuccessful have other problems related to the end of life.

Death and Dying The most common causes of death in patients with cancer are infection (leading to circulatory failure), respiratory failure, hepatic failure, and renal failure. Intestinal blockage may lead to inanition and starvation. Central nervous system disease may lead to seizures, coma, and central hypoventilation. About 70% of patients develop dyspnea preterminally. However, many months usually pass between the diagnosis of cancer and the occurrence of these complications, and during this period, the patient is severely affected by the possibility of death. The path of unsuccessful cancer treatment usually occurs in three phases. First, there is optimism at the hope of cure; when the tumor recurs, there is the acknowledgment of an incurable disease, and the goal of palliative therapy is embraced in the hope of being able to live with disease; finally, at the disclosure of imminent death, another adjustment in outlook takes place. The patient imagines the worst in preparation for the end of life and may go through stages of adjustment to the diagnosis. These stages include denial, isolation, anger, bargaining, depression, acceptance, and hope. Of course, patients do not all progress through all the stages or proceed through them in the same order or at the same rate. Nevertheless, developing an understanding of how the patient has been affected by the diagnosis and is coping with it is an important goal of patient management.

It is best to speak frankly with the patient and the family regarding the likely course of disease. These discussions can be difficult for the physician as well as for the patient and family. The critical features of the interaction are to reassure the patient and family that everything that can be done to provide comfort will be done. They will not be abandoned. Many patients prefer to be cared for in their homes or in a hospice setting rather than a hospital. The American College of Physicians has published a book called *Home Care Guide for Cancer: How to Care for Family and Friends at Home* that teaches an approach to successful problem-solving in home care. With appropriate planning, it should be possible to provide the patient with the necessary medical care as well as the psychological and spiritual support that will prevent the isolation and depersonalization that can attend in-hospital death.

The care of dying patients may take a toll on the physician. A “burn-out” syndrome has been described that is characterized by fatigue, disengagement from patients and colleagues, and a loss of self-fulfillment. Efforts at stress reduction, maintenance of a balanced life, and setting realistic goals may combat this disorder.

End-of-Life Decisions Unfortunately, a smooth transition in treatment goals from curative to palliative may not be possible in all cases because of the occurrence of serious treatment-related complications or rapid disease progression. Vigorous and invasive medical support for a reversible disease or treatment complication is assumed to be

justified. However, if the reversibility of the condition is in doubt, the patient's wishes determine the level of medical care. These wishes should be elicited before the terminal phase of illness and reviewed periodically. Information about advance directives can be obtained from the American Association of Retired Persons, 601 E Street, NW, Washington, DC 20049, 202-434-2277, or Choice in Dying, 250 West 57th Street, New York, NY 10107, 212-366-5540. Some states allow physicians to assist patients who choose to end their lives. This subject is challenging from an ethical and a medical point of view. Discussions of end-of-life decisions should be candid and involve clear informed consent, waiting periods, second opinions, and documentation. **A full discussion of end-of-life management is in Chap. 10.**

100 Prevention and Early Detection of Cancer

Jennifer M. Croswell, Otis W. Brawley, Barnett S. Kramer

Improved understanding of carcinogenesis has allowed cancer prevention and early detection (also known as cancer control) to expand beyond the identification and avoidance of carcinogens. Specific interventions to prevent cancer in those at risk, and effective screening for early detection of cancer, are the goals.

Carcinogenesis is not an event but a process, a continuum of discrete tissue and cellular changes over time resulting in aberrant physiologic processes. Prevention concerns the identification and manipulation of the biologic, environmental, social, and genetic factors in the causal pathway of cancer.

EDUCATION AND HEALTHFUL HABITS

Public education on the avoidance of identified risk factors for cancer and encouraging healthy habits contributes to cancer prevention and control. The clinician is a powerful messenger in this process. The patient-provider encounter provides an opportunity to teach patients about the hazards of smoking, the features of a healthy lifestyle, use of proven cancer screening methods, and avoidance of excessive sun exposure.

SMOKING CESSATION

Tobacco smoking is a strong, modifiable risk factor for cardiovascular disease, pulmonary disease, and cancer. Smokers have an approximately 1 in 3 lifetime risk of dying prematurely from a tobacco-related cancer, cardiovascular, or pulmonary disease. Tobacco use causes more deaths from cardiovascular disease than from cancer. Lung cancer and cancers of the larynx, oropharynx, esophagus, kidney, bladder, pancreas, and stomach are all tobacco-related.

The number of cigarettes smoked per day and the level of inhalation of cigarette smoke are correlated with risk of lung cancer mortality. Light- and low-tar cigarettes are not safer, because smokers tend to inhale them more frequently and deeply.

Those who stop smoking have a 30–50% lower 10-year lung cancer mortality rate compared to those who continue smoking, despite the fact that some carcinogen-induced gene mutations persist for years after smoking cessation. Smoking cessation and avoidance would save more lives than any other public health activity.

The risk of tobacco smoke is not limited to the smoker. Environmental tobacco smoke, known as secondhand or passive smoke, causes lung cancer and other cardiopulmonary diseases in nonsmokers.

Tobacco use prevention is a pediatric issue. More than 80% of adult American smokers began smoking before the age of 18 years. Approximately 20% of Americans in grades 9 through 12 have smoked a cigarette in the past month. Counseling of adolescents and young adults is critical to prevent smoking. A clinician's simple advice can be of benefit. Providers should query patients on tobacco use and offer smokers assistance in quitting.

Current approaches to smoking cessation recognize smoking as an addiction (Chap. 470). The smoker who is quitting goes through identifiable stages that include contemplation of quitting, an action phase in which the smoker quits, and a maintenance phase. Smokers who quit completely are more likely to be successful than those who gradually reduce the number of cigarettes smoked or change to lower- or lower-nicotine cigarettes. More than 90% of the Americans who have successfully quit smoking did so on their own, without participation in an organized cessation program, but cessation programs are helpful for some smokers. The Community Intervention Trial for Smoking Cessation (COMMIT) was a 4-year program showing that light smokers (<25 cigarettes per day) were more likely to benefit from simple cessation messages and cessation programs than those who did not receive an intervention. Quit rates were 30.6% in the intervention group and 27.5% in the control group. The COMMIT interventions were unsuccessful in heavy smokers (<25 cigarettes per day). Heavy smokers may need an intensive broad-based cessation program that includes counseling, behavioral strategies, and pharmacologic adjuncts, such as nicotine replacement (gum, patches, sprays, lozenges, and inhalers), bupropion, and/or varenicline.

The health risks of cigars are similar to those of cigarettes. Smoking one or two cigars daily doubles the risk for oral and esophageal cancers; smoking three or four cigars daily increases the risk of oral cancers more than eightfold and esophageal cancer fourfold. The risks of occasional use are unknown.

Smokeless tobacco also represents a substantial health risk. Chewing tobacco is a carcinogen linked to dental caries, gingivitis, oral leukoplakia, and oral cancer. The systemic effects of smokeless tobacco (including snuff) may increase risks for other cancers. Esophageal cancer is linked to carcinogens in tobacco dissolved in saliva and swallowed. The net effects of e-cigarettes on health are poorly studied. Whether they aid in smoking cessation or serve as a “gateway” for nonsmoking children to acquire a smoking habit is debated.

PHYSICAL ACTIVITY

Physical activity is associated with a decreased risk of colon and breast cancer. A variety of mechanisms have been proposed. However, such studies are prone to confounding factors such as recall bias, association of exercise with other health-related practices, and effects of preclinical cancers on exercise habits (reverse causality).

DIET MODIFICATION

International epidemiologic studies suggest that diets high in fat are associated with increased risk for cancers of the breast, colon, prostate, and endometrium. These cancers have their highest incidence and mortalities in Western cultures, where fat composes an average of one-third of the total calories consumed.

Despite correlations, dietary fat has not been proven to cause cancer. Case-control and cohort epidemiologic studies give conflicting results. In addition, diet is a highly complex exposure to many nutrients and chemicals. Low-fat diets are associated with many dietary changes beyond simple subtraction of fat. Other lifestyle changes are also associated with adherence to a low-fat diet.

In observational studies, dietary fiber is associated with a reduced risk of colonic polyps and invasive cancer of the colon. However, cancer-protective effects of increasing fiber and lowering dietary fat have not been proven in the context of a prospective clinical trial. The putative protective mechanisms are complex and speculative. Fiber binds oxidized bile acids and generates soluble fiber products, such as butyrate, that may have differentiating properties. Fiber does not increase bowel transit times. Two large prospective cohort studies of >100,000 health professionals showed no association between fruit and vegetable intake and risk of cancer.

The Polyp Prevention Trial randomly assigned 2000 elderly persons, who had polyps removed, to a low-fat, high-fiber diet versus routine diet for 4 years. No differences were noted in polyp formation.

The U.S. National Institutes of Health Women’s Health Initiative, launched in 1994, was a long-term clinical trial enrolling >100,000 women age 45–69 years. It placed women in 22 intervention groups.

Participants received calcium/vitamin D supplementation; hormone replacement therapy; and counseling to increase exercise, eat a low-fat diet with increased consumption of fruits, vegetables, and fiber, and cease smoking. The study showed that although dietary fat intake was lower in the diet intervention group, invasive breast cancers were not reduced over an 8-year follow-up period compared to the control group. No reduction was seen in the incidence of colorectal cancer in the dietary intervention arm. The difference in dietary fat averaged ~10% between the two groups. Evidence does not currently establish the anticarcinogenic value of vitamin, mineral, or nutritional supplements in amounts greater than those provided by a balanced diet.

ENERGY BALANCE

Risk of cancer appears to increase as body mass index increases beyond 25 kg/m². Obesity is associated with increased risk for cancers of the colon, breast (female postmenopausal), endometrium, kidney (renal cell), and esophagus, although causality has not been established.

In observational studies, relative risks of colon cancer are increased in obesity by 1.5–2 for men and 1.2–1.5 for women. Obese postmenopausal women have a 30–50% increased relative risk of breast cancer. An unproven hypothesis for the association is that adipose tissue serves as a depot for aromatase that facilitates estrogen production.

SUN AVOIDANCE

Nonmelanoma skin cancers (basal cell and squamous cell) are induced by cumulative exposure to ultraviolet (UV) radiation. Intermittent acute sun exposure and sun damage have been linked to melanoma, but the evidence is inconsistent. Sunburns, especially in childhood and adolescence, may be associated with an increased risk of melanoma in adulthood. Reduction of sun exposure through use of protective clothing and changing patterns of outdoor activities can reduce skin cancer risk. Sunscreens decrease the risk of actinic keratoses, the precursor to squamous cell skin cancer, but melanoma risk may not be reduced. Sunscreens prevent burning, but they may encourage more prolonged exposure to the sun and may not filter out wavelengths of energy that cause melanoma.

Educational interventions to help individuals assess their risk of developing skin cancer have some impact. In particular, appearance-focused behavioral interventions in young women can decrease indoor tanning use and other UV exposures. Self-examination for skin pigment characteristics associated with skin cancer, such as freckling, may be useful in identifying people at high risk. Those who recognize themselves as being at risk tend to be more compliant with sun-avoidance recommendations. Risk factors for melanoma include a propensity to sunburn, a large number of benign melanocytic nevi, and atypical nevi.

CANCER CHEMOPREVENTION

Chemoprevention involves the use of specific natural or synthetic chemical agents to reverse, suppress, or prevent carcinogenesis before the development of invasive malignancy.

Cancer develops through an accumulation of tissue abnormalities associated with genetic and epigenetic changes, and growth regulatory pathways that are potential points of intervention to prevent cancer. The initial changes are termed *initiation*. The alteration can be inherited or acquired through the action of physical, infectious, or chemical carcinogens. Like most human diseases, cancer arises from an interaction between genetics and environmental exposures (Table 100-1). Influences that cause the initiated cell and its surrounding tissue microenvironment to progress through the carcinogenic process and change phenotypically are termed *promoters*. Promoters include hormones such as androgens, linked to prostate cancer, and estrogen, linked to breast and endometrial cancer. The distinction between an initiator and promoter is indistinct; some components of cigarette smoke are “complete carcinogens,” acting as both initiators and promoters. Cancer can be prevented or controlled through interference with the factors that cause cancer initiation, promotion, or progression. Compounds of interest in chemoprevention often have antimutagenic, hormone modulation, anti-inflammatory, antiproliferative, or proapoptotic activity (or a combination).

TABLE 100-1 SUSPECTED CARCINOGENS

Carcinogens ^a	Associated Cancer or Neoplasm
Alkylating agents	Acute myeloid leukemia, bladder cancer
Androgens	Prostate cancer
Aromatic amines (dyes)	Bladder cancer
Arsenic	Cancer of the lung, skin
Asbestos	Cancer of the lung, pleura, peritoneum
Benzene	Acute myelocytic leukemia
Chromium	Lung cancer
Diethylstilbestrol (prenatal)	Vaginal cancer (clear cell)
Epstein-Barr virus	Burkitt's lymphoma, nasal T cell lymphoma
Estrogens	Cancer of the endometrium, liver, breast
Ethyl alcohol	Cancer of the breast, liver, esophagus, head and neck
<i>Helicobacter pylori</i>	Gastric cancer, gastric MALT lymphoma
Hepatitis B or C virus	Liver cancer
Human immunodeficiency virus	Non-Hodgkin's lymphoma, Kaposi's sarcoma, squamous cell carcinomas (especially of the urogenital tract)
Human papilloma virus	Cancers of the cervix, anus, oropharynx
Human T cell lymphotropic virus type 1 (HTLV-1)	Adult T cell leukemia/lymphoma
Immunosuppressive agents (azathioprine, cyclosporine, glucocorticoids)	Non-Hodgkin's lymphoma
Ionizing radiation (therapeutic or diagnostic)	Breast, bladder, thyroid, soft tissue, bone, hematopoietic, and many more
Nitrogen mustard gas	Cancer of the lung, head and neck, nasal sinuses
Nickel dust	Cancer of the lung, nasal sinuses
Diesel exhaust	Lung cancer (miners)
Phenacetin	Cancer of the renal pelvis and bladder
Polycyclic hydrocarbons	Cancer of the lung, skin (especially squamous cell carcinoma of scrotal skin)
Radon gas	Lung cancer
Schistosomiasis	Bladder cancer (squamous cell)
Sunlight (ultraviolet)	Skin cancer (squamous cell and melanoma)
Tobacco (including smokeless)	Cancer of the upper aerodigestive tract, bladder
Vinyl chloride	Liver cancer (angiosarcoma)

^aAgents that are thought to act as cancer initiators and/or promoters.

CHEMOPREVENTION OF CANCERS OF THE UPPER AERODIGESTIVE TRACT

Smoking causes diffuse epithelial injury in the oral cavity, neck, esophagus, and lung. Patients cured of squamous cell cancers of the lung, esophagus, oral cavity, and neck are at risk (as high as 5% per year) of developing second cancers of the upper aerodigestive tract. Cessation of cigarette smoking does not markedly decrease the cured cancer patient's risk of second malignancy, even though it does lower the cancer risk in those who have never developed a malignancy. Smoking cessation may halt the early stages of the carcinogenic process (such as metaplasia), but it may have no effect on late stages of carcinogenesis. This "field carcinogenesis" hypothesis for upper aerodigestive tract cancer has made "cured" patients an important population for chemoprevention of second malignancies.

Oral human papilloma virus (HPV) infection, particularly HPV-16, increases the risk for cancers of the oropharynx. This association exists even in the absence of other risk factors such as smoking or alcohol use (although the magnitude of increased risk appears greater than additive when HPV infection and smoking are both present). Oral HPV infection is believed to be largely sexually acquired. Although no direct evidence currently exists to confirm the hypothesis, the introduction of the HPV vaccine may eventually reduce oropharyngeal cancer rates.

Oral leukoplakia, a premalignant lesion commonly found in smokers, has been used as an intermediate marker of chemopreventive

activity in smaller shorter-duration, randomized, placebo-controlled trials. Response was associated with upregulation of retinoic acid receptor- β (RAR- β). Therapy with high, relatively toxic doses of isotretinoin (13-*cis*-retinoic acid) causes regression of oral leukoplakia. However, the lesions recur when the therapy is withdrawn, suggesting the need for long-term administration. More tolerable doses of isotretinoin have not shown benefit in the prevention of head and neck cancer. Isotretinoin also failed to prevent second malignancies in patients cured of early-stage non-small cell lung cancer; mortality rates were actually increased in current smokers.

Several large-scale trials have assessed agents in the chemoprevention of lung cancer in patients at high risk. In the α -tocopherol/ β -carotene (ATBC) Lung Cancer Prevention Trial, participants were male smokers, age 50–69 years at entry. Participants had smoked an average of one pack of cigarettes per day for 35.9 years. Participants received α -tocopherol, β -carotene, and/or placebo in a randomized, two-by-two factorial design. After median follow-up of 6.1 years, lung cancer incidence and mortality were statistically significantly increased in those receiving β -carotene. α -Tocopherol had no effect on lung cancer mortality, and no evidence suggested interaction between the two drugs. Patients receiving α -tocopherol had a higher incidence of hemorrhagic stroke.

The β -Carotene and Retinol Efficacy Trial (CARET) involved 17,000 American smokers and workers with asbestos exposure. Entrants were randomly assigned to one of four arms and received β -carotene, retinol, and/or placebo in a two-by-two factorial design. This trial also demonstrated harm from β -carotene: a lung cancer rate of 5 per 1000 subjects per year for those taking placebo and of 6 per 1000 subjects per year for those taking β -carotene.

The ATBC and CARET results demonstrate the importance of testing chemoprevention hypotheses thoroughly before their widespread implementation because the results contradict a number of observational studies. The Physicians' Health Trial showed no change in the risk of lung cancer for those taking β -carotene; however, fewer of its participants were smokers than those in the ATBC and CARET studies.

CHEMOPREVENTION OF COLON CANCER

Many colon cancer prevention trials are based on the premise that most colorectal cancers develop from adenomatous polyps. These trials use adenoma recurrence or disappearance as a surrogate endpoint (not yet validated) for colon cancer prevention. Early clinical trial results suggest that nonsteroidal anti-inflammatory drugs (NSAIDs), such as piroxicam, sulindac, and aspirin, may prevent adenoma formation or cause regression of adenomatous polyps. The mechanism of action of NSAIDs is unknown, but they are presumed to work through the cyclooxygenase pathway. Although two randomized controlled trials (the Physicians' Health Study and the Women's Health Study) did not show an effect of aspirin on colon cancer or adenoma incidence in persons with no previous history of colonic lesions after 10 years of therapy, these trials did show an approximately 18% relative risk reduction for colonic adenoma incidence in persons with a previous history of adenomas after 1 year. Pooled findings from observational cohort studies do demonstrate a 22% and 28% relative reduction in colorectal cancer and adenoma incidence, respectively, with regular aspirin use, and a well-conducted meta-analysis of four randomized controlled trials (albeit primarily designed to examine aspirin's effects on cardiovascular events) found that aspirin at doses of at least 75 mg resulted in a 24% relative reduction in colorectal cancer incidence after 20 years, with no clear increase in efficacy at higher doses. Cyclooxygenase-2 (COX-2) inhibitors have also been considered for colorectal cancer and polyp prevention. Trials with COX-2 inhibitors were initiated, but an increased risk of cardiovascular events in those taking the COX-2 inhibitors was noted, suggesting that these agents are not suitable for chemoprevention in the general population.

Epidemiologic studies suggest that diets high in calcium lower colon cancer risk. Calcium binds bile and fatty acids, which cause proliferation of colonic epithelium. It is hypothesized that calcium reduces intraluminal exposure to these compounds. The randomized controlled Calcium Polyp Prevention Study found that calcium supplementation

478 decreased the absolute risk of adenomatous polyp recurrence by 7% at 4 years; extended observational follow-up demonstrated a 12% absolute risk reduction 5 years after cessation of treatment. However, in the Women's Health Initiative, combined use of calcium carbonate and vitamin D twice daily did not reduce the incidence of invasive colorectal cancer compared with placebo after 7 years.

The Women's Health Initiative demonstrated that postmenopausal women taking estrogen plus progestin have a 44% lower relative risk of colorectal cancer compared to women taking placebo. Of >16,600 women randomized and followed for a median of 5.6 years, 43 invasive colorectal cancers occurred in the hormone group and 72 in the placebo group. The positive effect on colon cancer is mitigated by the modest increase in cardiovascular and breast cancer risks associated with combined estrogen plus progestin therapy.

A case-control study suggested that statins decrease the incidence of colorectal cancer; however, several subsequent case-control and cohort studies have not demonstrated an association between regular statin use and a reduced risk of colorectal cancer. No randomized controlled trials have addressed this hypothesis. A meta-analysis of statin use showed no protective effect of statins on overall cancer incidence or death.

CHEMOPREVENTION OF BREAST CANCER

Tamoxifen is an antiestrogen with partial estrogen agonistic activity in some tissues, such as endometrium and bone. One of its actions is to upregulate transforming growth factor β , which decreases breast cell proliferation. In randomized placebo-controlled trials to assess tamoxifen as adjuvant therapy for breast cancer, tamoxifen reduced the number of new breast cancers in the opposite breast by more than a third. In a randomized placebo-controlled prevention trial involving >13,000 pre- and postmenopausal women at high risk, tamoxifen decreased the risk of developing breast cancer by 49% (from 43.4 to 22 per 1000 women) after a median follow-up of nearly 6 years. Tamoxifen also reduced bone fractures; a small increase in risk of endometrial cancer, stroke, pulmonary emboli, and deep vein thrombosis was noted. The International Breast Cancer Intervention Study (IBIS-1) and the Italian Randomized Tamoxifen Prevention Trial also demonstrated a reduction in breast cancer incidence with tamoxifen use. A trial comparing tamoxifen with another selective estrogen receptor modulator, raloxifene, in postmenopausal women showed that raloxifene is comparable to tamoxifen in cancer prevention. This trial only included postmenopausal women. Raloxifene was associated with more invasive breast cancers and a trend toward more noninvasive breast cancers, but fewer thromboembolic events than tamoxifen; the drugs are similar in risks of other cancers, fractures, ischemic heart disease, and stroke. Both tamoxifen and raloxifene (the latter for postmenopausal women only) have been approved by the U.S. Food and Drug Administration (FDA) for reduction of breast cancer in women at high risk for the disease (1.66% risk at 5 years based on the Gail risk model: <http://www.cancer.gov/bcrisktool/>).

Because the aromatase inhibitors are even more effective than tamoxifen in adjuvant breast cancer therapy, it has been hypothesized that they would be more effective in breast cancer prevention. A randomized, placebo-controlled trial of exemestane reported a 65% relative reduction (from 5.5 to 1.9 per 1000 women) in the incidence of invasive breast cancer in women at elevated risk after a median follow-up of about 3 years. Common adverse effects included arthralgias, hot flashes, fatigue, and insomnia. No trial has directly compared aromatase inhibitors with selective estrogen receptor modulators for breast cancer chemoprevention.

CHEMOPREVENTION OF PROSTATE CANCER

Finasteride and dutasteride are 5- α -reductase inhibitors. They inhibit conversion of testosterone to dihydrotestosterone (DHT), a potent stimulator of prostate cell proliferation. The Prostate Cancer Prevention Trial (PCPT) randomly assigned men age 55 years or older at average risk of prostate cancer to finasteride or placebo. All men in the trial were being regularly screened with prostate-specific antigen (PSA) levels and digital rectal examination. After 7 years of therapy, the incidence of

prostate cancer was 18.4% in the finasteride arm, compared with 24.4% in the placebo arm, a statistically significant difference. However, the finasteride group had more patients with tumors of Gleason score 7 and higher compared with the placebo arm (6.4 vs 5.1%). Reassuringly, long-term (10–15 years) follow-up did not reveal any statistically significant differences in overall mortality between all men in the finasteride and placebo arms or in men diagnosed with prostate cancer; differences in prostate cancer in favor of finasteride persisted.

Dutasteride has also been evaluated as a preventive agent for prostate cancer. The Reduction by Dutasteride of Prostate Cancer Events (REDUCE) trial was a randomized double-blind trial in which approximately 8200 men with an elevated PSA (2.5–10 ng/mL for men age 50–60 years and 3–10 ng/mL for men age 60 years or older) and negative prostate biopsy on enrollment received daily 0.5 mg of dutasteride or placebo. The trial found a statistically significant 23% relative risk reduction in the incidence of biopsy-detected prostate cancer in the dutasteride arm at 4 years of treatment (659 cases vs 858 cases, respectively). Overall, across years 1 through 4, there was no difference between the arms in the number of tumors with a Gleason score of 7 to 10; however, during years 3 and 4, there was a statistically significant difference in tumors with Gleason score of 8 to 10 in the dutasteride arm (12 tumors vs 1 tumor, respectively).

The clinical importance of the apparent increased incidence of higher-grade tumors in the 5- α -reductase inhibitor arms of these trials is controversial. It may likely represent an increased sensitivity of PSA and digital rectal exam for high-grade tumors in men receiving these agents. The FDA has analyzed both trials, and it determined that the use of a 5- α -reductase inhibitor for prostate cancer chemoprevention would result in one additional high-grade (Gleason score 8 to 10) prostate cancer for every three to four lower-grade (Gleason score <6) tumors averted. Although it acknowledged that detection bias may have accounted for the finding, it stated that it could not conclusively dismiss a causative role for 5- α -reductase inhibitors. These agents are therefore not FDA-approved for prostate cancer prevention.

Because all men in both the PCPT and REDUCE trials were being screened and because screening approximately doubles the rate of prostate cancer, it is not known if finasteride or dutasteride decreases the risk of prostate cancer in men who are not being screened.

Several favorable laboratory and observational studies led to the formal evaluation of selenium and α -tocopherol (vitamin E) as potential prostate cancer preventives. The Selenium and Vitamin E Cancer Prevention Trial (SELECT) assigned 35,533 men to receive 200 μ g/d selenium, 400 IU/d α -tocopherol, selenium plus vitamin E, or placebo. After a median follow-up of 7 years, a trend toward an increased risk of developing prostate cancer was observed for those men taking vitamin E alone as compared to the placebo arm (hazard ratio 1.17; 95% confidence interval, 1.004–1.36).

VACCINES AND CANCER PREVENTION

A number of infectious agents cause cancer. Hepatitis B and C are linked to liver cancer; some HPV strains are linked to cervical, anal, and head and neck cancer; and *Helicobacter pylori* is associated with gastric adenocarcinoma and gastric lymphoma. Vaccines to protect against these agents may reduce the risk of their associated cancers.

The hepatitis B vaccine is effective in preventing hepatitis and hepatomas due to chronic hepatitis B infection.

A quadrivalent HPV vaccine (covering HPV strains 6, 11, 16, and 18) and a bivalent vaccine (covering HPV strains 16 and 18) are available for use in the United States. HPV types 16 and 18 cause cervical and anal cancer; reduction in these HPV types could prevent >70% of cervical cancers worldwide. HPV types 6 and 11 cause genital papillomas. For individuals not previously infected with these HPV strains, the vaccines demonstrate high efficacy in preventing persistent strain-specific HPV infections; however, the trials and substudies that evaluated the vaccines' ability to prevent cervical and anal cancer relied on surrogate outcome measures (cervical or anal intraepithelial neoplasia [CIN/AIN] I, II, and III), and the degree of durability of the immune response beyond 5 years is not currently known. The vaccines do not appear to impact preexisting infections and the efficacy appears to

be markedly lower for populations that had previously been exposed to vaccine-specific HPV strains. The vaccine is recommended in the United States for females and males age 9–26 years.

SURGICAL PREVENTION OF CANCER

Some organs in some individuals are at such high risk of developing cancer that surgical removal of the organ at risk may be considered. Women with severe cervical dysplasia are treated with laser or loop electrosurgical excision or conization and occasionally even hysterectomy. Colectomy is used to prevent colon cancer in patients with familial polyposis or ulcerative colitis.

Prophylactic bilateral mastectomy may be chosen for breast cancer prevention among women with genetic predisposition to breast cancer. In a prospective series of 139 women with *BRCA1* and *BRCA2* mutations, 76 chose to undergo prophylactic mastectomy and 63 chose close surveillance. At 3 years, no cases of breast cancer had been diagnosed in those opting for surgery, but eight patients in the surveillance group had developed breast cancer. A larger (n = 639) retrospective cohort study reported that three patients developed breast cancer after prophylactic mastectomy compared with an expected incidence of 30–53 cases: a 90–94% reduction in breast cancer risk. Postmastectomy breast cancer–related deaths were reduced by 81–94% for high-risk women compared with sister controls and by 100% for moderate-risk women when compared with expected rates.

Prophylactic oophorectomy may also be employed for the prevention of ovarian and breast cancers among high-risk women. A prospective cohort study evaluating the outcomes of *BRCA* mutation carriers demonstrated a statistically significant association between prophylactic oophorectomy and a reduced incidence of ovarian or primary peritoneal cancer (36% relative risk reduction, or a 4.5% absolute difference). Studies of prophylactic oophorectomy for prevention of breast cancer in women with genetic mutations have shown relative risk reductions of approximately 50%; the risk reduction may be greatest for women having the procedure at younger (i.e., <50 years) ages.

All of the evidence concerning the use of prophylactic mastectomy and oophorectomy for prevention of breast and ovarian cancer in high-risk women has been observational in nature; such studies are prone to a variety of biases, including case selection bias, family relationships between patients and controls, and inadequate information about hormone use. Thus, they may give an overestimate of the magnitude of benefit.

CANCER SCREENING

Screening is a means of detecting disease early in asymptomatic individuals, with the goal of decreasing morbidity and mortality. While screening can potentially reduce disease-specific deaths and has been shown to do so in cervical, colon, lung, and breast cancer, it is also subject to a number of biases that can suggest a benefit when actually there is none. Biases can even mask net harm. Early detection does not in itself confer benefit. Cause-specific mortality, rather than survival after diagnosis, is the preferred endpoint (see below).

Because screening is done on asymptomatic, healthy persons, it should offer substantial likelihood of benefit that outweighs harm. Screening tests and their appropriate use should be carefully evaluated before their use is widely encouraged in screening programs, as a matter of public policy.

A large and increasing number of genetic mutations and nucleotide polymorphisms have been associated with an increased risk of cancer. Testing for these genetic mutations could in theory define a high-risk population. However, most of the identified mutations have very low penetrance and individually provide minimal predictive accuracy. The ability to predict the development of a particular cancer may some day present therapeutic options as well as ethical dilemmas. It may eventually allow for early intervention to prevent a cancer or limit its severity. People at high risk may be ideal candidates for chemoprevention and screening; however, efficacy of these interventions in the high-risk population should be investigated. Currently, persons at high risk for a particular cancer can engage in intensive screening. While this course is clinically reasonable, it is not known if it reduces mortality in these populations.

TABLE 100-2 ASSESSMENT OF THE VALUE OF A DIAGNOSTIC TEST^a

	Condition Present	Condition Absent
Positive test	a	b
Negative test	c	d
a = true positive		
b = false positive		
c = false negative		
d = true negative		
Sensitivity	The proportion of persons with the condition who test positive: $a/(a + c)$	
Specificity	The proportion of persons without the condition who test negative: $d/(b + d)$	
Positive predictive value (PPV)	The proportion of persons with a positive test who have the condition: $a/(a + b)$	
Negative predictive value	The proportion of persons with a negative test who do not have the condition: $d/(c + d)$	
Prevalence, sensitivity, and specificity determine PPV		
$PPV = \frac{\text{prevalence} \times \text{sensitivity}}{(\text{prevalence} \times \text{sensitivity}) + (1 - \text{prevalence})(1 - \text{specificity})}$		

^aFor diseases of low prevalence, such as cancer, poor specificity has a dramatic adverse effect on PPV such that only a small fraction of positive tests are true positives.

The Accuracy of Screening A screening test's accuracy or ability to discriminate disease is described by four indices: sensitivity, specificity, positive predictive value, and negative predictive value (Table 100-2). *Sensitivity*, also called the true-positive rate, is the proportion of persons with the disease who test positive in the screen (i.e., the ability of the test to detect disease when it is present). *Specificity*, or 1 minus the false-positive rate, is the proportion of persons who do not have the disease that test negative in the screening test (i.e., the ability of a test to correctly identify that the disease is not present). The *positive predictive value* is the proportion of persons who test positive that actually have the disease. Similarly, *negative predictive value* is the proportion testing negative that do not have the disease. The sensitivity and specificity of a test are independent of the underlying prevalence (or risk) of the disease in the population screened, but the predictive values depend strongly on the prevalence of the disease.

Screening is most beneficial, efficient, and economical when the target disease is common in the population being screened. Specificity is at least as important to the ultimate feasibility and success of a screening test as sensitivity.

Potential Biases of Screening Tests Common biases of screening are lead time, length-biased sampling, and selection. These biases can make a screening test seem beneficial when actually it is not (or even causes net harm). Whether beneficial or not, screening can create the false impression of an epidemic by increasing the number of cancers diagnosed. It can also produce a shift in the proportion of patients diagnosed at an early stage and inflate survival statistics without reducing mortality (i.e., the number of deaths from a given cancer relative to the number of those at risk for the cancer). In such a case, the *apparent* duration of survival (measured from date of diagnosis) increases without lives being saved or life expectancy changed.

Lead-time bias occurs whether or not a test influences the natural history of the disease; the patient is merely diagnosed at an earlier date. Survival *appears* increased even if life is not really prolonged. The screening test only prolongs the time the subject is aware of the disease and spends as a patient.

Length-biased sampling occurs because screening tests generally can more easily detect slow-growing, less aggressive cancers than fast-growing cancers. Cancers diagnosed due to the onset of symptoms between scheduled screenings are on average more aggressive, and treatment outcomes are not as favorable. An extreme form of length bias sampling is termed *overdiagnosis*, the detection of "pseudo disease." The reservoir of some undetected slow-growing tumors is large. Many of these tumors fulfill the histologic criteria of cancer but will

480 never become clinically significant or cause death. This problem is compounded by the fact that the most common cancers appear most frequently at ages when competing causes of death are more frequent.

Selection bias must be considered in assessing the results of any screening effort. The population most likely to seek screening may differ from the general population to which the screening test might be applied. In general, volunteers for studies are more health conscious and likely to have a better prognosis or lower mortality rate, irrespective of the screening result. This is termed the *healthy volunteer effect*.

Potential Drawbacks of Screening Risks associated with screening include harm caused by the screening intervention itself, harm due to the further investigation of persons with positive tests (both true and false positives), and harm from the treatment of persons with a true-positive result, whether or not life is extended by treatment (e.g., even if a screening test reduces relative cause-specific mortality by 20–30%, 70–80% of those diagnosed still go on to die of the target cancer). The diagnosis and treatment of cancers that would never have caused medical problems can lead to the harm of unnecessary treatment and give patients the anxiety of a cancer diagnosis. The psychosocial impact of cancer screening can also be substantial when applied to the entire population.

Assessment of Screening Tests Good clinical trial design can offset some biases of screening and demonstrate the relative risks and benefits of a screening test. A randomized controlled screening trial with cause-specific mortality as the endpoint provides the strongest support for a screening intervention. Overall mortality should also be reported to detect an adverse effect of screening and treatment on other disease outcomes (e.g., cardiovascular disease). In a randomized trial, two like populations are randomly established. One is given the usual standard of care (which may be no screening at all) and the other receives the screening intervention being assessed. The two populations are compared over time. Efficacy for the population studied is established when the group receiving the screening test has a better cause-specific mortality rate than the control group. Studies showing a reduction in the incidence of advanced-stage disease, improved survival, or a stage shift are weaker (and possibly misleading) evidence of benefit. These latter criteria are early indicators but not sufficient to establish the value of a screening test.

Although a randomized, controlled screening trial provides the strongest evidence to support a screening test, it is not perfect. Unless the trial is population-based, it does not remove the question of generalizability to the target population. Screening trials generally involve thousands of persons and last for years. Less definitive study designs are therefore often used to estimate the effectiveness of screening practices. However, every nonrandomized study design is subject to strong confounders. In descending order of strength, evidence may also be derived from the findings of internally controlled trials using intervention allocation methods other than randomization (e.g., allocation by birth date, date of clinic visit); the findings of analytic observational studies; or the results of multiple time series studies with or without the intervention.

Screening for Specific Cancers Screening for cervical, colon, and breast cancer is beneficial for certain age groups. Depending on age and smoking history, lung cancer screening can also be beneficial in specific settings. Special surveillance of those at high risk for a specific cancer because of a family history or a genetic risk factor may be prudent, but few studies have assessed the influence on mortality. A number of organizations have considered whether or not to endorse routine use of certain screening tests. Because these groups have not used the same criteria to judge whether a screening test should be endorsed, they have arrived at different recommendations. The American Cancer Society (ACS) and the U.S. Preventive Services Task Force (USPSTF) publish screening guidelines (Table 100-3); the American Academy of Family Practitioners (AAFP) generally follow/endorse the USPSTF recommendations; and the American College of Physicians (ACP) develops recommendations based on structured reviews of other organizations' guidelines.

BREAST CANCER Breast self-examination, clinical breast examination by a caregiver, mammography, and magnetic resonance imaging (MRI) have all been variably advocated as useful screening tools.

A number of trials have suggested that annual or biennial screening with mammography or mammography plus clinical breast examination in normal-risk women older than age 50 years decreases breast cancer mortality. Each trial has been criticized for design flaws. In most trials, breast cancer mortality rate is decreased by 15–30%. Experts disagree on whether average-risk women age 40–49 years should receive regular screening (Table 100-3). The U.K. Age Trial, the only randomized trial of breast cancer screening to specifically evaluate the impact of mammography in women age 40–49 years, found no statistically significant difference in breast cancer mortality for screened women versus controls after about 11 years of follow-up (relative risk 0.83; 95% confidence interval 0.66–1.04); however, <70% of women received screening in the intervention arm, potentially diluting the observed effect. A meta-analysis of eight large randomized trials showed a 15% relative reduction in mortality (relative risk 0.85; 95% confidence interval 0.75–0.96) from mammography screening for women age 39–49 years after 11–20 years of follow-up. This is equivalent to a number needed to invite to screening of 1904 over 10 years to prevent one breast cancer death. At the same time, nearly half of women age 40–49 years screened annually will have false-positive mammograms necessitating further evaluation, often including biopsy. Estimates of overdiagnosis range from 10 to 40% of diagnosed invasive cancers. In the United States, widespread screening over the last several decades has not been accompanied by a reduction in incidence of metastatic breast cancer despite a large increase in early-stage disease, suggesting a substantial amount of overdiagnosis at the population level.

No study of breast self-examination has shown it to decrease mortality. A randomized controlled trial of approximately 266,000 women in China demonstrated no difference in breast cancer mortality between a group that received intensive breast self-exam instruction and reinforcement/reminders and controls at 10 years of follow-up. However, more benign breast lesions were discovered and more breast biopsies were performed in the self-examination arm.

Genetic screening for *BRCA1* and *BRCA2* mutations and other markers of breast cancer risk has identified a group of women at high risk for breast cancer. Unfortunately, when to begin and the optimal frequency of screening have not been defined. Mammography is less sensitive at detecting breast cancers in women carrying *BRCA1* and *BRCA2* mutations, possibly because such cancers occur in younger women, in whom mammography is known to be less sensitive. MRI screening may be more sensitive than mammography in women at high risk due to genetic predisposition or in women with very dense breast tissue, but specificity may be lower. An increase in overdiagnosis may accompany the higher sensitivity. The impact of MRI on breast cancer mortality with or without concomitant use of mammography has not been evaluated in a randomized controlled trial.

CERVICAL CANCER Screening with Papanicolaou (Pap) smears decreases cervical cancer mortality. The cervical cancer mortality rate has fallen substantially since the widespread use of the Pap smear. With the onset of sexual activity comes the risk of sexual transmission of HPV, the fundamental etiologic factor for cervical cancer. Screening guidelines recommend regular Pap testing for all women who have reached the age of 21 (prior to this age, even in individuals that have begun sexual activity, screening may cause more harm than benefit). The recommended interval for Pap screening is 3 years. Screening more frequently adds little benefit but leads to important harms, including unnecessary procedures and overtreatment of transient lesions. Beginning at age 30, guidelines also offer the alternative of combined Pap smear and HPV testing for women. The screening interval for women who test normal using this approach may be lengthened to 5 years.

An upper age limit at which screening ceases to be effective is not known, but women age 65 years with no abnormal results in the previous 10 years may choose to stop screening. Screening should be

TABLE 100-3 SCREENING RECOMMENDATIONS FOR ASYMPTOMATIC SUBJECTS NOT KNOWN TO BE AT INCREASED RISK FOR THE TARGET CONDITION^a

Cancer Type	Test or Procedure	USPSTF	ACS
Breast	Self-examination	"D"	Women ≥20 years: Breast self-exam is an option
	Clinical examination	Women ≥40 years: "I" (as a stand-alone without mammography)	Women 20–39 years: Perform every 3 years Women ≥40 years: Perform annually
	Mammography	Women 40–49 years: The decision should be an individual one, and take patient context/values into account ("C") Women 50–74 years: Every 2 years ("B") Women ≥75 years: "I"	Women ≥40 years: Screen annually for as long as the woman is in good health
	Magnetic resonance imaging (MRI)	"I"	Women with >20% lifetime risk of breast cancer: Screen with MRI plus mammography annually Women with 15–20% lifetime risk of breast cancer: Discuss option of MRI plus mammography annually Women with <15% lifetime risk of breast cancer: Do not screen annually with MRI
Cervical	Pap test (cytology)	Women 21–65 years: Screen every 3 years ("A")	Women 21–29 years: Screen every 3 years
		Women <21 years: "D"	Women 30–65 years: Acceptable approach to screen with cytology every 3 years (see HPV test below)
		Women >65 years, with adequate, normal prior Pap screenings: "D"	Women <21 years: No screening Women >65 years: No screening following adequate negative prior screening
	HPV test	Women after total hysterectomy for noncancerous causes: "D"	Women after total hysterectomy for noncancerous causes: Do not screen
		Women 30–65 years: Screen in combination with cytology every 5 years if woman desires to lengthen the screening interval (see Pap test, above) ("A")	Women 30–65 years: Preferred approach to screen with HPV and cytology co-testing every 5 years (see Pap test above)
		Women <30 years: "D"	Women <30 years: Do not use HPV testing
	Women >65 years, with adequate, normal prior Pap screenings: "D"	Women >65 years: No screening following adequate negative prior screening	
	Women after total hysterectomy for noncancerous causes: "D"	Women after total hysterectomy for noncancerous causes: Do not screen	
Colorectal	Sigmoidoscopy	Adults 50–75 years: every 5 years in combination with high-sensitivity FOBT every 3 years ("A") ^b	Adults ≥50 years: Screen every 5 years
		Adults 76–85 years: "C"	
		Adults ≥85 years: "D"	
	Fecal occult blood testing (FOBT)	Adults 50–75 years: Annually, for high-sensitivity FOBT ("A")	Adults ≥50 years: Screen every year
		Adults 76–85 years: "C"	
	Colonoscopy	Adults 50–75 years: every 10 years ("A") Adults 76–85 years: "C" Adults ≥85 years: "D"	Adults ≥50 years: Screen every 10 years
Fecal DNA testing	"I"	Adults ≥50 years: Screen, but interval uncertain	
	Fecal immunochemical testing (FIT)	"I"	Adults ≥50 years: Screen every year
CT colonography	"I"	Adults ≥50 years: Screen every 5 years	
Lung	Low-dose computed tomography (CT) scan	Adults 55–80 years, with a ≥30 pack-year smoking history, still smoking or have quit within past 15 years. Discontinue once a person has not smoked for 15 years or develops a health problem that substantially limits life expectancy or the ability to have curative lung surgery: "B"	Men and women, 55–74 years, with ≥30 pack-year smoking history, still smoking or have quit within past 15 years: Discuss benefits, limitations, and potential harms of screening; only perform screening in facilities with the right type of CT scanner and with high expertise/specialists
Ovarian	CA-125	"D"	There is no sufficiently accurate test proven effective in the early detection of ovarian cancer. For women at high risk of ovarian cancer and/or who have unexplained, persistent symptoms, the combination of CA-125 and transvaginal ultrasound with pelvic exam may be offered.
	Transvaginal ultrasound	"D"	

(Continued)

Cancer Type	Test or Procedure	USPSTF	ACS
Prostate	Prostate-specific antigen (PSA)	Men, all ages: "D"	Starting at age 50, men should talk to a doctor about the pros and cons of testing so they can decide if testing is the right choice for them. If African American or have a father or brother who had prostate cancer before age 65, men should have this talk starting at age 45. How often they are tested will depend on their PSA level.
	Digital rectal examination (DRE)	No individual recommendation	As for PSA; if men decide to be tested, they should have the PSA blood test with or without a rectal exam
Skin	Complete skin examination by clinician or patient	"I"	Self-examination monthly; clinical exam as part of routine cancer-related checkup

^aSummary of the screening procedures recommended for the general population by the USPSTF and the ACS. These recommendations refer to asymptomatic persons who are not known to have risk factors, other than age or gender, for the targeted condition. ^bUSPSTF lettered recommendations are defined as follows: "A": The USPSTF recommends the service, because there is high certainty that the net benefit is substantial; "B": The USPSTF recommends the service, because there is high certainty that the net benefit is moderate or moderate certainty that the net benefit is moderate to substantial; "C": The USPSTF recommends selectively offering or providing this service to individual patients based on professional judgment and patient preferences; there is at least moderate certainty that the net benefit is small; "D": The USPSTF recommends against the service because there is moderate or high certainty that the service has no net benefit or that the harms outweigh the benefits; "I": The USPSTF concludes that the current evidence is insufficient to assess the balance of benefits and harms of the service.

Abbreviations: ACS, American Cancer Society; USPSTF, U.S. Preventive Services Task Force.

discontinued in women who have undergone a hysterectomy for non-cancerous reasons.

Although the efficacy of the Pap smear in reducing cervical cancer mortality has never been directly confirmed in a randomized, controlled setting, a clustered randomized trial in India evaluated the impact of one-time cervical visual inspection and immediate colposcopy, biopsy, and/or cryotherapy (where indicated) versus counseling on cervical cancer deaths in women age 30–59 years. After 7 years of follow-up, the age-standardized rate of death due to cervical cancer was 39.6 per 100,000 person-years in the intervention group versus 56.7 per 100,000 person-years in controls.

COLORECTAL CANCER Fecal occult blood testing (FOBT), digital rectal examination (DRE), rigid and flexible sigmoidoscopy, colonoscopy, and computed tomography (CT) colonography have been considered for colorectal cancer screening. A meta-analysis of four randomized controlled trials demonstrated a 15% relative reduction in colorectal cancer mortality with FOBT. The sensitivity for FOBT is increased if specimens are rehydrated before testing, but at the cost of lower specificity. The false-positive rate for rehydrated FOBT is high; 1–5% of persons tested have a positive test. Only 2–10% of those with occult blood in the stool have cancer. The high false-positive rate of FOBT dramatically increases the number of colonoscopies performed.

Fecal immunochemical tests appear to have higher sensitivity for colorectal cancer than nonrehydrated FOBT tests. Fecal DNA testing is an emerging testing modality; it may have increased sensitivity and comparable specificity to FOBT and could potentially reduce harms associated with follow-up of false-positive tests. The body of evidence on the operating characteristics and effectiveness of fecal DNA tests in reducing colorectal cancer mortality is limited.

Two meta-analyses of five randomized controlled trials of sigmoidoscopy (i.e., the NORCCAP, SCORE, PLCO, Telemark, and U.K. trials) found an 18% relative reduction in colorectal cancer incidence and a 28% relative reduction in colorectal cancer mortality. Participant ages ranged from 50 to 74 years, with follow-up ranging from 6 to 13 years. Diagnosis of adenomatous polyps by sigmoidoscopy should lead to evaluation of the entire colon with colonoscopy. The most efficient interval for screening sigmoidoscopy is unknown, but an interval of 5 years is often recommended. Case-control studies suggest that intervals of up to 15 years may confer benefit; the U.K. trial demonstrated benefit with one-time screening.

One-time colonoscopy detects ~25% more advanced lesions (polyps >10 mm, villous adenomas, adenomatous polyps with high-grade dysplasia, invasive cancer) than one-time FOBT with sigmoidoscopy; comparative *programmatic* performance of the two modalities over time is not known. Perforation rates are about 3/1000 for colonoscopy and 1/1000 for sigmoidoscopy. Debate continues on whether colonoscopy is too expensive and invasive and whether sufficient provider capacity exists to be recommended as the preferred screening tool in

standard-risk populations. Some observational studies suggest that efficacy of colonoscopy to decrease colorectal cancer mortality is primarily limited to the left side of the colon.

CT colonography, if done at expert centers, appears to have a sensitivity for polyps ≥6 mm comparable to colonoscopy. However, the rate of extracolonic findings of abnormalities of uncertain significance that must nevertheless be worked up is high (~15–30%); the long-term cumulative radiation risk of repeated colonography screenings is also a concern.

LUNG CANCER Chest x-ray and sputum cytology have been evaluated in several randomized lung cancer screening trials. The most recent and largest (n = 154,901) of these, one substudy of the Prostate, Lung, Colorectal, and Ovarian (PLCO) cancer screening trial, found that, compared with usual care, annual chest x-ray did not reduce the risk of dying from lung cancer (relative risk 0.99; 95% confidence interval 0.87–1.22) after 13 years. Low-dose CT has also been evaluated in several randomized trials. The largest and longest of these, the National Lung Screening Trial (NLST), was a randomized controlled trial of screening for lung cancer in approximately 53,000 persons age 55–74 years with a 30+ pack-year smoking history. It demonstrated a statistically significant relative reduction of about 15–20% in lung cancer mortality in the CT arm compared to the chest x-ray arm (or about 3 fewer deaths per 1000 people screened with CT). However, the harms include the potential radiation risks associated with multiple scans, the discovery of incidental findings of unclear significance, and a high rate of false-positive test results. Both incidental findings and false-positive tests can lead to invasive diagnostic procedures associated with anxiety, expense, and complications (e.g., pneumo- or hemothorax after lung biopsy). The NLST was performed at experienced screening centers, and the balance of benefits and harms may differ in the community setting at less experienced centers.

OVARIAN CANCER Adnexal palpation, transvaginal ultrasound (TVUS), and serum CA-125 assay have been considered for ovarian cancer screening. A large randomized controlled trial has shown that an annual screening program of TVUS and CA-125 in average-risk women does not reduce deaths from ovarian cancer (relative risk 1.21; 95% confidence interval 0.99–1.48). Adnexal palpation was dropped early in the study because it did not detect any ovarian cancers that were not detected by either TVUS or CA-125. The risks and costs associated with the high number of false-positive results are impediments to routine use of these modalities for screening. In the PLCO trial, 10% of participants had a false-positive result from TVUS or CA-125, and one-third of these women underwent a major surgical procedure; the ratio of surgeries to screen-detected ovarian cancer was approximately 20:1.

PROSTATE CANCER The most common prostate cancer screening modalities are DRE and serum PSA assay. An emphasis on PSA screening has caused prostate cancer to become the most common nonskin cancer diagnosed in American males. This disease is prone to lead-time bias,

length bias, and overdiagnosis, and substantial debate continues among experts as to whether screening should be offered unless the patient specifically asks to be screened. Virtually all organizations stress the importance of informing men about the uncertainty regarding screening efficacy and the harms associated with screening. Prostate cancer screening clearly detects many asymptomatic cancers, but the ability to distinguish tumors that are lethal but still curable from those that pose little or no threat to health is limited, and randomized trials indicate that the effect of PSA screening on prostate cancer mortality across a population is, at best, small. Men older than age 50 years have a high prevalence of indolent, clinically insignificant prostate cancers (about 30–50% of men, increasing further as men age).

Two major randomized controlled trials of the impact of PSA screening on prostate cancer mortality have been published. The PLCO Cancer Screening Trial was a multicenter U.S. trial that randomized almost 77,000 men age 55–74 years to receive either annual PSA testing for 6 years or usual care. At 13 years of follow-up, no statistically significant difference in the number of prostate cancer deaths were noted between the arms (rate ratio 1.09; 95% confidence interval 0.87–1.36). Approximately 50% of men in the control arm received at least one PSA test during the trial, which may have potentially diluted a small effect.

The European Randomized Study of Screening for Prostate Cancer (ERSPC) was a multinational study that randomized approximately 182,000 men between age 50 and 74 years (with a predefined “core” screening group of men age 55–69 years) to receive PSA testing or no screening. Recruitment and randomization procedures, as well as actual frequency of PSA testing, varied by country. After a median follow-up of 11 years, a 20% relative reduction in the risk of prostate cancer death in the screened arm was noted in the “core” screening group. The trial found that 1055 men would need to be invited to screening, and 37 cases of prostate cancer detected, to avert 1 death from prostate cancer. Of the seven countries included in the mortality analysis, two demonstrated statistically significant reductions in prostate cancer deaths, whereas five did not. There was also an imbalance in treatment between the two study arms, with a higher proportion of men with clinically localized cancer receiving radical prostatectomy in the screening arm and receiving it at experienced referral centers.

Treatments for low-stage prostate cancer, such as surgery and radiation therapy, can cause significant morbidity, including impotence and urinary incontinence. In a trial conducted in the United States after the initiation of widespread PSA testing, random assignment to radical prostatectomy compared with “watchful waiting” did not result in a statistically significant decrease in prostate cancer deaths (absolute risk reduction 2.7%; 95% confidence interval –1.3 to 6.2%).

SKIN CANCER Visual examination of all skin surfaces by the patient or by a health care provider is used in screening for basal and squamous cell cancers and melanoma. No prospective randomized study has been performed to look for a mortality decrease. Unfortunately, screening is associated with a substantial rate of overdiagnosis.

CANCER IS A GENETIC DISEASE

Cancer arises through a series of somatic alterations in DNA that result in unrestrained cellular proliferation. Most of these alterations involve actual sequence changes in DNA (i.e., mutations). They may originate as a consequence of random replication errors, exposure to carcinogens (e.g., radiation), or faulty DNA repair processes. While most cancers arise sporadically, familial clustering of cancers occurs in certain families that carry a germline mutation in a cancer gene.

HISTORICAL PERSPECTIVE

The idea that cancer progression is driven by sequential somatic mutations in specific genes has only gained general acceptance in the past 25 years. Before the advent of the microscope, cancer was believed to be composed of aggregates of mucus or other noncellular matter. By the middle of the nineteenth century, it became clear that tumors were masses of cells and that these cells arose from the normal cells of the tissue from which the cancer originated. However, the molecular basis for the uncontrolled proliferation of cancer cells was to remain a mystery for another century. During that time, a number of theories for the origin of cancer were postulated. The great biochemist Otto Warburg proposed the combustion theory of cancer, which stipulated that cancer was due to abnormal oxygen metabolism. In addition, some believed that all cancers were caused by viruses, and that cancer was in fact a contagious disease.

In the end, observations of cancer occurring in chimney sweeps, studies of x-rays, and the overwhelming data demonstrating cigarette smoke as a causative agent in lung cancer, together with Ames's work on chemical mutagenesis, provided convincing evidence that cancer originated through changes in DNA. Although the viral theory of cancer did not prove to be generally accurate (with the exception of human papillomaviruses, which can cause cervical and other cancers in human), the study of retroviruses led to the discovery of the first human *oncogenes* in the late 1970s. Soon after, the study of families with genetic predisposition to cancer was instrumental in the discovery of *tumor-suppressor genes*. The field that studies the type of mutations, as well as the consequence of these mutations in tumor cells, is now known as *cancer genetics*.

THE CLONAL ORIGIN AND MULTISTEP NATURE OF CANCER

Nearly all cancers originate from a single cell; this clonal origin is a critical discriminating feature between neoplasia and hyperplasia.

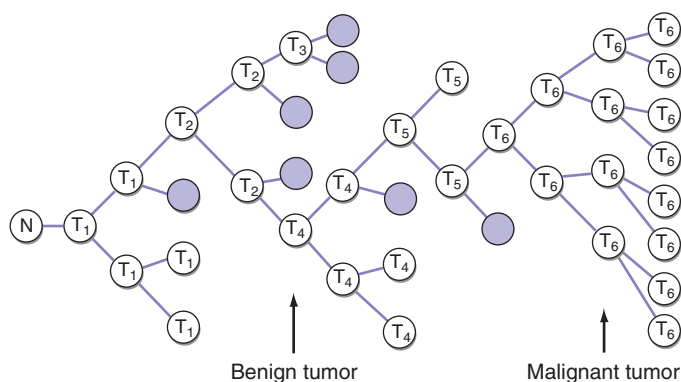


FIGURE 101e-1 Multistep clonal development of malignancy. In this diagram a series of five cumulative mutations (T_1, T_2, T_4, T_5, T_6), each with a modest growth advantage acting alone, eventually results in a malignant tumor. Note that not all such alterations result in progression; for example, the T_3 clone is a dead end. The actual number of cumulative mutations necessary to transform from the normal to the malignant state is unknown in most tumors. (After P Nowell: *Science* 194:23, 1976, with permission.)

Multiple cumulative mutational events are invariably required for the progression of a tumor from normal to fully malignant phenotype. The process can be seen as Darwinian microevolution in which, at each successive step, the mutated cells gain a growth advantage resulting in an increased representation relative to their neighbors (Fig. 101e-1). Based on observations of cancer frequency increases during aging, as well as molecular genetics work, it is believed that 5 to 10 accumulated mutations are necessary for a cell to progress from the normal to the fully malignant phenotype.

We are beginning to understand the precise nature of the genetic alterations responsible for some malignancies and to get a sense of the order in which they occur. The best-studied example is colon cancer, in which analyses of DNA from tissues extending from normal colon epithelium through adenoma to carcinoma have identified some of the genes mutated in the process (Fig. 101e-2). Other malignancies are believed to progress in a similar stepwise fashion, although the order and identity of genes affected may be different.

TWO TYPES OF CANCER GENES: ONCOGENES AND TUMOR-SUPPRESSOR GENES

There are two major types of cancer genes. The first type comprises genes that positively influence tumor formation and are known as *oncogenes*. The second type of cancer genes negatively impact tumor growth and have been named *tumor-suppressor genes*. Both oncogenes and tumor-suppressor genes exert their effects on tumor growth

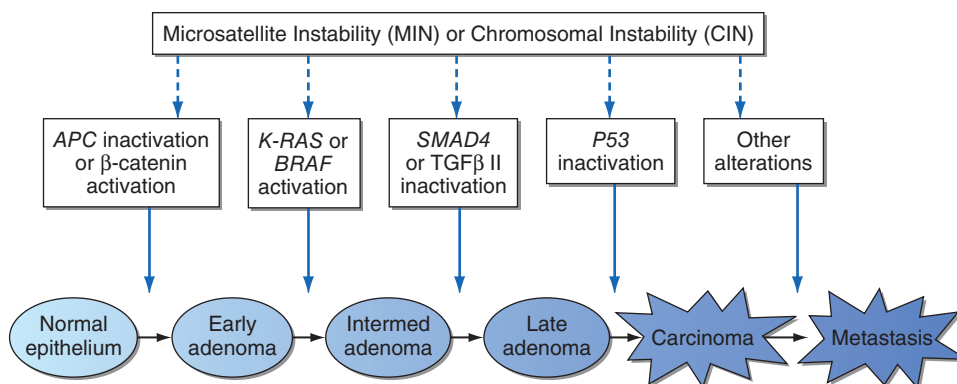


FIGURE 101e-2 Progressive somatic mutational steps in the development of colon carcinoma. The accumulation of alterations in a number of different genes results in the progression from normal epithelium through adenoma to full-blown carcinoma. Genetic instability (microsatellite or chromosomal) accelerates the progression by increasing the likelihood of mutation at each step. Patients with familial polyposis are already one step into this pathway, because they inherit a germline alteration of the *APC* gene. TGF, transforming growth factor.

TABLE 101e-1 COMMON ONCOGENES ALTERED IN HUMAN CANCERS

Oncogene	Function	Alteration in Cancer	Neoplasm
<i>AKT1</i>	Serine/threonine kinase	Amplification	Stomach
<i>AKT2</i>	Serine/threonine kinase	Amplification	Ovarian, breast, pancreatic
<i>BRAF</i>	Serine/threonine kinase	Point mutation	Melanoma, lung, colorectal
<i>CDK4</i>	Cyclin-dependent kinase	Point mutation, amplification	Breast, melanoma, myeloma, others
<i>CTNNB1</i>	Signal transduction	Point mutation	Colon, prostate, melanoma, skin, others
<i>FOS</i>	Transcription factor	Overexpression	Osteosarcomas
<i>ERBB2</i>	Receptor tyrosine kinase	Point mutation, amplification	Breast, ovary, stomach, neuroblastoma
<i>JUN</i>	Transcription factor	Overexpression	Lung
<i>MET</i>	Receptor tyrosine kinase	Point mutation, rearrangement	Osteocarcinoma, kidney, glioma
<i>MYB</i>	Transcription factor	Amplification	AML, CML, colorectal, melanoma
<i>C-MYC</i>	Transcription factor	Amplification	Breast, colon, gastric, lung
<i>L-MYC</i>	Transcription factor	Amplification	Lung, bladder
<i>N-MYC</i>	Transcription factor	Amplification	Neuroblastoma, lung
<i>PIK3A</i>	Phosphoinositol-3-kinase	Point Mutation	Multiple cancers
<i>HRAS</i>	GTPase	Point mutation	Colon, lung, pancreas
<i>KRAS</i>	GTPase	Point mutation	Melanoma, colorectal, AML
<i>NRAS</i>	GTPase	Point mutation	Various carcinomas, melanoma
<i>REL</i>	Transcription factor	Rearrangement, amplification	Lymphomas
<i>WNT1</i>	Growth factor	Amplification	Retinoblastoma

Abbreviations: AML, acute myeloid leukemia; CML, chronic myeloid leukemia.

through their ability to control cell division (cell birth) or cell death (apoptosis), although the mechanisms can be extremely complex. While tightly regulated in normal cells, oncogenes acquire mutations in cancer cells, and the mutations typically relieve this control and lead to increased activity of the gene products. This mutational event typically occurs in a single allele of the oncogene and acts in a dominant fashion. In contrast, the normal function of tumor-suppressor genes is usually to restrain cell growth, and this function is lost in cancer. Because of the diploid nature of mammalian cells, both alleles must be inactivated for a cell to completely lose the function of a tumor-suppressor gene, leading to a recessive mechanism at the cellular level. From these ideas and studies on the inherited form of retinoblastoma, Knudson and others formulated the *two-hit hypothesis*, which in its modern version states that both copies of a tumor-suppressor gene must be inactivated in cancer.

There is a subset of tumor-suppressor genes, the *caretaker genes*, that do not affect cell growth directly, but rather control the ability of the cell to maintain the integrity of its genome. Cells with a deficiency in these genes have an increased rate of mutations throughout their genomes, including in oncogenes and tumor-suppressor genes. This “mutator” phenotype was first hypothesized by Loeb to explain how the multiple mutational events required for tumorigenesis can occur in the lifetime of an individual. A mutator phenotype has now been observed in some forms of cancer, such as those associated with deficiencies in DNA mismatch repair. The great majority of cancers, however, do not harbor repair deficiencies, and their rate of mutation is similar to that observed in normal cells. Many of these cancers, however, appear to harbor a different kind of genetic instability, affecting the loss or gains of whole chromosomes or large parts thereof (as explained in more detail below).

ONCOGENES IN HUMAN CANCER

Work by Peyton Rous in the early 1900s revealed that a chicken sarcoma could be transmitted from animal to animal in cell-free extracts, suggesting that cancer could be induced by an agent acting positively to promote tumor formation. The agent responsible for the transmission of the cancer was a retrovirus (Rous sarcoma virus, RSV) and the oncogene responsible was identified 75 years later as *v-src*. Other oncogenes were also discovered through their presence in the genomes of retroviruses that are capable of causing cancers in chickens, mice, and rats. The cellular homologues of these viral genes are called

protooncogenes and are often targets of mutation or aberrant regulation in human cancer. Whereas many oncogenes were discovered because of their presence in retroviruses, other oncogenes, particularly those involved in translocations characteristic of particular leukemias and lymphomas, were isolated through genomic approaches. Investigators cloned the sequences surrounding the chromosomal translocations observed cytogenetically and then deduced the nature of the genes that were the targets of these translocations (see below). Some of these were oncogenes known from retroviruses (like *ABL*, involved in chronic myeloid leukemia [CML]), whereas others were new (like *BCL2*, involved in B cell lymphoma). In the normal cellular environment, protooncogenes have crucial roles in cell proliferation and differentiation. **Table 101e-1** is a partial list of oncogenes known to be involved in human cancer.

The normal growth and differentiation of cells is controlled by growth factors that bind to receptors on the surface of the cell. The signals generated by the membrane receptors are transmitted inside the cells through signaling cascades involving kinases, G proteins, and other regulatory proteins. Ultimately, these signals affect the activity of transcription factors in the nucleus, which regulate the expression of genes crucial in cell proliferation, cell differentiation, and cell death. Oncogene products have been found to function at critical steps in these pathways (**Chap. 102e**), and inappropriate activation of these pathways can lead to tumorigenesis.

MECHANISMS OF ONCOGENE ACTIVATION

POINT MUTATION

Point mutation is a common mechanism of oncogene activation. For example, mutations in one of the *RAS* genes (*HRAS*, *KRAS*, or *NRAS*) are present in up to 85% of pancreatic cancers and 45% of colon cancers but are less common in other cancer types, although they can occur at significant frequencies in leukemia, lung, and thyroid cancers. Remarkably—and in contrast to the diversity of mutations found in tumor-suppressor genes (see below)—most of the activated *RAS* genes contain point mutations in codons 12, 13, or 61 (these mutations reduce *RAS* GTPase activity, leading to constitutive activation of the mutant *RAS* protein). The restricted pattern of mutations observed in oncogenes compared to that of tumor-suppressor genes reflects the fact that gain-of-function mutations are less likely to occur than mutations that simply lead to loss of activity. Indeed, inactivation of a gene can in theory be accomplished through the introduction of a stop

codon anywhere in the coding sequence, whereas activations require precise substitutions at residues that can somehow lead to an increase in the activity of the encoded protein. Importantly, the specificity of oncogene mutations provides diagnostic opportunities, as tests that identify mutations at defined positions are easier to design than tests aimed at detecting random changes in a gene.

DNA AMPLIFICATION

The second mechanism for activation of oncogenes is DNA sequence amplification, leading to overexpression of the gene product. This increase in DNA copy number may cause cytologically recognizable chromosome alterations referred to as *homogeneous staining regions* (HSRs) if integrated within chromosomes, or *double minutes* (dmins) if extrachromosomal. The recognition of DNA amplification is accomplished through various cytogenetic techniques such as comparative genomic hybridization (CGH) or fluorescence in situ hybridization (FISH), which allow the visualization of chromosomal aberrations using fluorescent dyes. In addition, noncytogenetic, microarray-based approaches are now available for identifying changes in copy number at high resolution. Newer short-tag-based sequencing approaches have been used to evaluate amplifications. When paired with next-generation sequencing, this approach offers the highest degree of resolution and quantification available. With both microarray and sequencing technologies, the entire genome can be surveyed for gains and losses of DNA sequences, thus pinpointing chromosomal regions likely to contain genes important in the development or progression of cancer.

Numerous genes have been reported to be amplified in cancer. Several of these genes, including *NMYC* and *LMYC*, were identified through their presence within the amplified DNA sequences of a tumor and had homology to known oncogenes. Because the region amplified often includes hundreds of thousands of base pairs, multiple oncogenes may be amplified in a single amplicon in some cancers (particularly in sarcomas). Indeed, *MDM2*, *GLI*, *CDK4*, and *SAS* at chromosomal location 12q13-15 have been shown to be simultaneously amplified in several types of sarcomas and other tumors. Amplification of a cellular gene is often a predictor of poor prognosis; for example, *ERBB2/HER2* and *NMYC* are often amplified in aggressive breast cancers and neuroblastoma, respectively.

CHROMOSOMAL REARRANGEMENT

Chromosomal alterations provide important clues to the genetic changes in cancer. The chromosomal alterations in human solid tumors such as carcinomas are heterogeneous and complex and occur as a result of the frequent chromosomal instability (CIN) observed in these tumors (see below). In contrast, the chromosome alterations in myeloid and lymphoid tumors are often simple translocations, i.e., reciprocal transfers of chromosome arms from one chromosome to another. Consequently, many detailed and informative chromosome analyses have been performed on hematopoietic cancers. The breakpoints of recurring chromosome abnormalities usually occur at the site of cellular oncogenes. **Table 101e-2** lists representative examples of recurring chromosome alterations in malignancy and the associated gene(s) rearranged or deregulated by the chromosomal rearrangement. Translocations are particularly common in lymphoid tumors, probably because these cell types have the capability to rearrange their DNA to generate antigen receptors. Indeed, antigen receptor genes are commonly involved in the translocations, implying that an imperfect regulation of receptor gene rearrangement may be involved in the pathogenesis. An interesting example is Burkitt's lymphoma, a B cell tumor characterized by a reciprocal translocation between chromosomes 8 and 14. Molecular analysis of Burkitt's lymphomas demonstrated that the breakpoints occurred within or near the *MYC* locus on chromosome 8 and within the immunoglobulin heavy chain locus on chromosome 14, resulting in the transcriptional activation of *MYC*. Enhancer activation by translocation, although not universal, appears to play an important role in malignant progression. In addition to transcription factors and signal transduction molecules, translocation may result in the overexpression of cell cycle

TABLE 101e-2 REPRESENTATIVE ONCOGENES AT CHROMOSOMAL TRANSLOCATIONS

Gene (Chromosome)	Translocation	Malignancy
<i>ABL</i> (9q34.1)– <i>BCR</i> (22q11)	(9;22)(q34;q11)	Chronic myeloid leukemia
<i>ATF1</i> (12q13)– <i>EWS</i> (22q12)	(12;22)(q13;q12)	Malignant melanoma of soft parts
<i>BCL1</i> (11q13.3)– <i>IgH</i> (14q32)	(11;14)(q13;q32)	Mantle cell lymphoma
<i>BCL2</i> (18q21.3)– <i>IgH</i> (14q32)	(14;18)(q32;q21)	Follicular lymphoma
<i>FLI1</i> (11q24)– <i>EWS</i> (22q12)	(11;22)(q24;q12)	Ewing's sarcoma
<i>LCK</i> (1p34)– <i>TCRB</i> (7q35)	(1;7)(p34;q35)	T cell acute lymphocytic leukemia
<i>MYC</i> (8q24)– <i>IgH</i> (14q32)	(8;14)(q24;q32)	Burkitt's lymphoma, B cell acute lymphocytic leukemia
<i>PAX3</i> (2q35)– <i>FKHR/ALV</i> (13q14)	(2;13)(q35;q14)	Alveolar rhabdomyosarcoma
<i>PAX7</i> (1p36)– <i>KHR/ALV</i> (13q14)	(1;13)(p36;q14)	Alveolar rhabdomyosarcoma
<i>REL</i> (2p13)– <i>NRG</i> (2p11.2-14)	Inv(2)(p13;p11.2-14)	Non-Hodgkin's lymphoma
<i>RET</i> (10q11.2)– <i>PKAR1A</i> (17q23)	(10;17)(q11.2;q23)	Thyroid carcinoma
<i>TAL1</i> (1p32)– <i>TCTA</i> (3p21)	(1;3)(p34;p21)	Acute T cell leukemia
<i>TRK</i> (1q23-1q24)– <i>TPM3</i> (1q31)	Inv1(q23;q31)	Colon carcinoma
<i>WT1</i> (11p13)– <i>EWS</i> (22q12)	(11;22)(p13;q12)	Desmoplastic small round cell tumor

Source: From R Hesketh: *The Oncogene and Tumour Suppressor Gene Facts Book*, 2nd ed. San Diego, Academic Press, 1997; with permission.

regulatory proteins or proteins such as cyclins and of proteins that regulate cell death.

The first reproducible chromosome abnormality detected in human malignancy was the Philadelphia chromosome detected in CML. This cytogenetic abnormality is generated by reciprocal translocation involving the *ABL* oncogene on chromosome 9, encoding a tyrosine kinase, being placed in proximity to the *BCR* (breakpoint cluster region) gene on chromosome 22. **Figure 101e-3** illustrates the generation of the translocation and its protein product. The consequence of expression of the *BCR-ABL* gene product is the activation of signal transduction pathways leading to cell growth independent of normal external signals. Imatinib (marketed as Gleevec), a drug that specifically blocks the activity of Abl tyrosine kinase, has shown remarkable efficacy with little toxicity in patients with CML. It is hoped that knowledge of genetic alterations in other cancers will likewise lead to mechanism-based design and development of a new generation of chemotherapeutic agents.

CHROMOSOMAL INSTABILITY IN SOLID TUMORS

Solid tumors are generally highly aneuploid, containing an abnormal number of chromosomes; these chromosomes also exhibit structural alterations such as translocations, deletions, and amplifications. These abnormalities are collectively referred to as chromosomal instability (CIN). Normal cells possess several cell cycle checkpoints, essentially quality-control requirements that have to be met before subsequent events are allowed to take place. The mitotic checkpoint, which ensures proper chromosome attachment to the mitotic spindle before allowing the sister chromatids to separate, is altered in certain cancers. The molecular basis of CIN remains unclear, although a number of mitotic checkpoint genes are found mutated or abnormally expressed in various tumors. The exact effects of these changes on the mitotic checkpoint are unknown, and both weakening and overactivation of the checkpoint have been proposed. The identification of the cause of CIN

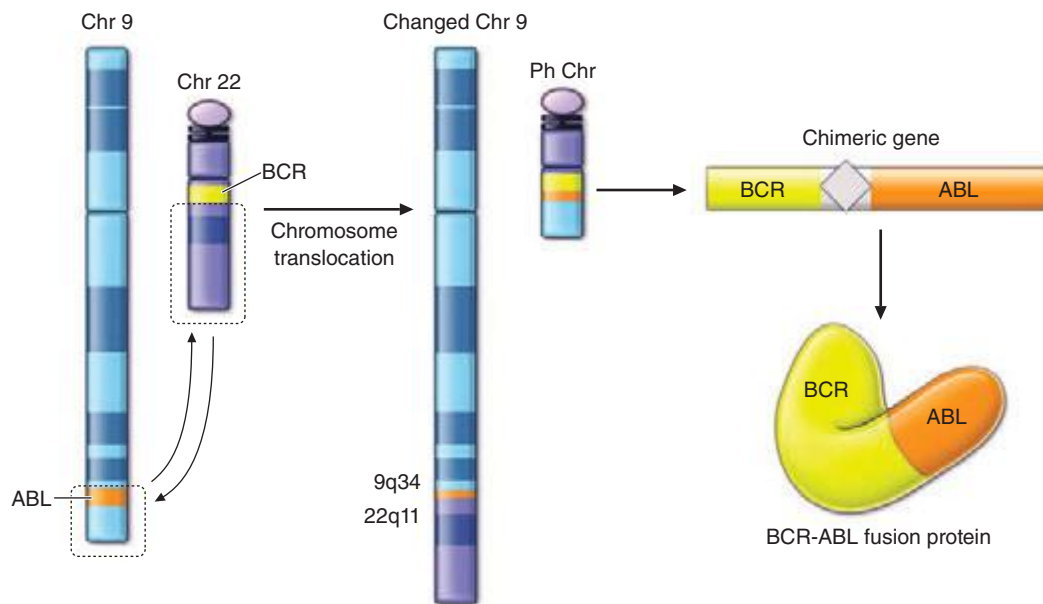


FIGURE 101e-3 Specific translocation seen in chronic myeloid leukemia (CML). The Philadelphia chromosome (Ph) is derived from a reciprocal translocation between chromosomes 9 and 22 with the breakpoint joining the sequences of the *ABL* oncogene with the *BCR* gene. The fusion of these DNA sequences allows the generation of an entirely novel fusion protein with modified function.

in tumors will likely be a formidable task, considering that several hundred genes are thought to control the mitotic checkpoint and other cellular processes ensuring proper chromosome segregation. Regardless of the mechanisms underlying CIN, the measurement of the number of chromosomal alterations present in tumors is now possible with both cytogenetic and molecular techniques, and several studies have shown that this information can be useful for prognostic purposes. In addition, because the mitotic checkpoint is essential for cellular viability, it may become a target for novel therapeutic approaches.

TUMOR-SUPPRESSOR GENE INACTIVATION IN CANCER

The first indication of the existence of tumor-suppressor genes came from experiments showing that fusion of mouse cancer cells with normal mouse fibroblasts led to a nonmalignant phenotype in the fused cells. The normal role of tumor-suppressor genes is to restrain cell growth, and the function of these genes is inactivated in cancer. The two major types of somatic lesions observed in tumor-suppressor genes during tumor development are *point mutations* and *large deletions*. Point mutations in the coding region of tumor-suppressor genes will frequently lead to truncated protein products or otherwise nonfunctional proteins. Similarly, deletions lead to the loss of a functional product and sometimes encompass the entire gene or even the entire chromosome arm, leading to loss of heterozygosity (LOH) in the tumor DNA compared to the corresponding normal tissue DNA (Fig. 101e-4). LOH in tumor DNA is considered a hallmark for the presence of a tumor-suppressor gene at a particular chromosomal location, and LOH studies have been useful in the positional cloning of many tumor-suppressor genes.

Gene silencing, an epigenetic change that leads to the loss of gene expression and occurs in conjunction with hypermethylation of the promoter and histone deacetylation, is another mechanism of tumor-suppressor gene inactivation. (An *epigenetic modification* refers to a change in the genome, heritable by cell progeny, that does not involve a change in the DNA sequence. The inactivation of the second X chromosome in female cells is an example of an epigenetic silencing that prevents gene expression from the inactivated chromosome.) During embryologic development, regions of chromosomes from one parent are silenced and gene expression proceeds from the chromosome of the other parent. For most genes, expression occurs from both alleles or randomly from one allele or the other. The preferential expression of a particular gene exclusively from the allele contributed by one

parent is called *parental imprinting* and is thought to be regulated by covalent modifications of chromatin protein and DNA (often methylation) of the silenced allele.

The role of epigenetic control mechanisms in the development of human cancer is unclear. However, a general decrease in the level of DNA methylation has been noted as a common change in cancer. In addition, numerous genes, including some tumor-suppressor genes, appear to become hypermethylated and silenced during tumorigenesis. *VHL* and *p16INK4* are well-studied examples of such tumor-suppressor genes. Overall, epigenetic mechanisms may be responsible for reprogramming the expression of a large number of genes in cancer and, together with the mutation of specific genes, are likely to be crucial in the development of human malignancies. The use of drugs that can reverse epigenetic changes in cancer cells may represent a novel therapeutic option in certain cancers or premalignant conditions. For example, demethylating agents (azacitidine or decitabine) are now approved by the U.S. Food and Drug Administration (FDA) for the treatment of patients with high-risk myelodysplastic syndrome (MDS).

FAMILIAL CANCER SYNDROMES

A small fraction of cancers occur in patients with a genetic predisposition. In these families, the affected individuals have a predisposing loss-of-function mutation in one allele of a tumor-suppressor gene. The tumors in these patients show a loss of the remaining normal allele as a result of somatic events (point mutations or deletions), in agreement with the two-hit hypothesis (Fig. 101e-4). Thus, most cells of an individual with an inherited loss-of-function mutation in a tumor-suppressor gene are functionally normal, and only the rare cells that develop a mutation in the remaining normal allele will exhibit uncontrolled regulation.

Roughly 100 syndromes of familial cancer have been reported, although many are rare. The majority are inherited as autosomal dominant traits, although some of those associated with DNA repair abnormalities (xeroderma pigmentosum, Fanconi's anemia, ataxia telangiectasia) are autosomal recessive. Table 101e-3 shows a number of cancer predisposition syndromes and the responsible genes. The current paradigm states that the genes mutated in familial syndromes can also be targets for somatic mutations in sporadic (noninherited) tumors. The study of cancer syndromes has thus provided invaluable insights into the mechanisms of progression for many tumor types. This section examines the case of inherited colon cancer in detail, but

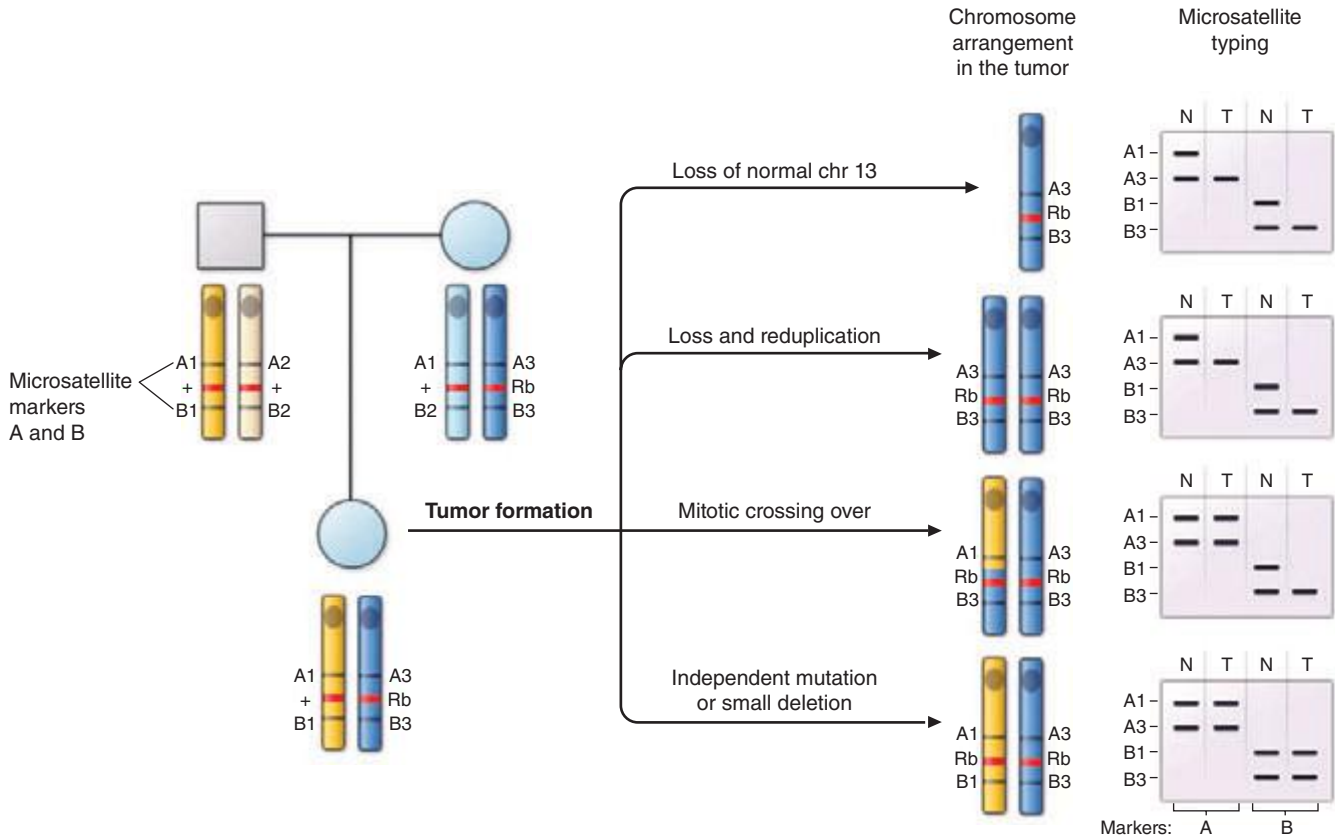


FIGURE 101e-4 Diagram of possible mechanisms for tumor formation in an individual with hereditary (familial) retinoblastoma. On the left is shown the pedigree of an affected individual who has inherited the abnormal (*Rb*) allele from her affected mother. The normal allele is shown as a (+). The four chromosomes of her two parents are drawn to indicate their origin. Flanking the retinoblastoma locus are microsatellite markers (A and B) also analyzed in this family. Markers A3 and B3 are on the chromosome carrying the retinoblastoma disease gene. Tumor formation results when the normal allele, which this patient inherited from her father, is inactivated. On the right are shown four possible ways in which this could occur. In each case, the resulting chromosome 13 arrangement is shown, as well as the results of PCR typing using the microsatellite markers comparing normal tissue (N) with tumor tissue (T). Note that in the first three situations, the normal allele (B1) has been lost in the tumor tissue, which is referred to as loss of heterozygosity (LOH) at this locus.

similar lessons can be applied to many of the cancer syndromes listed in Table 101e-3. In particular, the study of inherited colon cancer will clearly illustrate the difference between two types of tumor-suppressor genes: the *gatekeepers*, which directly regulate the growth of tumors, and the *caretakers*, which, when mutated, lead to genetic instability and therefore act indirectly on tumor growth.

Familial adenomatous polyposis (FAP) is a dominantly inherited colon cancer syndrome due to germline mutations in the adenomatous polyposis coli (*APC*) tumor-suppressor gene on chromosome 5. Patients with this syndrome develop hundreds to thousands of adenomas in the colon. Each of these adenomas has lost the normal remaining allele of *APC* but has not yet accumulated the required additional mutations to generate fully malignant cells (Fig. 101e-2). The loss of the second functional *APC* allele in tumors from FAP families often occurs through loss of heterozygosity. However, out of these thousands of benign adenomas, several will invariably acquire further abnormalities and a subset will even develop into fully malignant cancers. *APC* is thus considered to be a gatekeeper for colon tumorigenesis: in the absence of mutation of this gatekeeper (or a gene acting within the same pathway), a colorectal tumor simply cannot form. **Figure 101e-5** shows germline and somatic mutations found in the *APC* gene. The function of the *APC* protein is still not completely understood, but it likely provides differentiation and apoptotic cues to colonic cells as they migrate up the crypts. Defects in this process may lead to abnormal accumulation of cells that should normally undergo apoptosis.

In contrast to patients with FAP, patients with hereditary nonpolyposis colon cancer (HNPCC, or Lynch's syndrome) do not develop multiple polyposis, but instead develop only one or a small number

of adenomas that rapidly progress to cancer. Most HNPCC cases are due to mutations in one of four DNA mismatch repair genes (Table 101e-3), which are components of a repair system that is normally responsible for correcting errors in freshly replicated DNA. Germline mutations in *MSH2* and *MLH1* account for more than 90% of HNPCC cases, whereas mutations in *MSH6* and *PMS2* are much less frequent. When a somatic mutation inactivates the remaining wild-type allele of a mismatch repair gene, the cell develops a hypermutable phenotype characterized by profound genomic instability, especially for the short repeated sequences called *microsatellites*. This microsatellite instability (MSI) favors the development of cancer by increasing the rate of mutations in many genes, including oncogenes and tumor-suppressor genes (Fig. 101e-2). These genes can thus be considered caretakers. Interestingly, CIN can also be found in colon cancer, but MSI and CIN appear to be mutually exclusive, suggesting that they represent alternative mechanisms for the generation of a mutator phenotype in this cancer (Fig. 101e-2). Other cancer types rarely exhibit MSI, but most exhibit CIN.

Although most autosomal dominant inherited cancer syndromes are due to mutations in tumor-suppressor genes (Table 101e-3), there are a few interesting exceptions. Multiple endocrine neoplasia type 2, a dominant disorder characterized by pituitary adenomas, medullary carcinoma of the thyroid, and (in some pedigrees) pheochromocytoma, is due to gain-of-function mutations in the protooncogene *RET* on chromosome 10. Similarly, gain-of-function mutations in the tyrosine kinase domain of the *MET* oncogene lead to hereditary papillary renal carcinoma. Interestingly, loss-of-function mutations in the *RET* gene cause a completely different disease, Hirschsprung's disease (aganglionic megacolon [**Chaps. 353 and 408**]).

TABLE 101e-3 CANCER PREDISPOSITION SYNDROMES AND ASSOCIATED GENES

Syndrome	Gene	Chromosome	Inheritance	Tumors
Ataxia telangiectasia	<i>ATM</i>	11q22-q23	AR	Breast
Autoimmune lymphoproliferative syndrome	<i>FAS</i> <i>FASL</i>	10q24 1q23	AD	Lymphomas
Bloom syndrome	<i>BLM</i>	15q26.1	AR	Several types
Cowden syndrome	<i>PTEN</i>	10q23	AD	Breast, thyroid
Familial adenomatous polyposis	<i>APC</i>	5q21	AD	Intestinal adenoma, colorectal
Familial melanoma	<i>p16INK4</i>	9p21	AD	Melanoma, pancreatic
Familial Wilms' tumor	<i>WT1</i>	11p13	AD	Kidney (pediatric)
Hereditary breast/ovarian cancer	<i>BRCA1</i> <i>BRCA2</i>	17q21 13q12.3	AD	Breast, ovarian, colon, prostate
Hereditary diffuse gastric cancer	<i>CDH1</i>	16q22	AD	Stomach
Hereditary multiple exostoses	<i>EXT1</i> <i>EXT2</i>	8q24 11p11-12	AD	Exostoses, chondrosarcoma
Hereditary prostate cancer	<i>HPC1</i>	1q24-25	AD	Prostate
Hereditary retinoblastoma	<i>RB1</i>	13q14.2	AD	Retinoblastoma, osteosarcoma
Hereditary nonpolyposis colon cancer (HNPCC)	<i>MSH2</i> <i>MLH1</i> <i>MSH6</i> <i>PMS2</i>	2p16 3p21.3 2p16 7p22	AD	Colon, endometrial, ovarian, stomach, small bowel, ureter carcinoma
Hereditary papillary renal carcinoma	<i>MET</i>	7q31	AD	Papillary kidney
Juvenile polyposis	<i>SMAD4</i>	18q21	AD	Gastrointestinal, pancreatic
Li-Fraumeni	<i>TP53</i>	17p13.1	AD	Sarcoma, breast
Multiple endocrine neoplasia type 1	<i>MEN1</i>	11q13	AD	Parathyroid, endocrine, pancreas, and pituitary
Multiple endocrine neoplasia type 2a	<i>RET</i>	10q11.2	AD	Medullary thyroid carcinoma, pheochromocytoma
Neurofibromatosis type 1	<i>NF1</i>	17q11.2	AD	Neurofibroma, neurofibrosarcoma, brain
Neurofibromatosis type 2	<i>NF2</i>	22q12.2	AD	Vestibular schwannoma, meningioma, spine
Nevoid basal cell carcinoma syndrome (Gorlin's syndrome)	<i>PTCH</i>	9q22.3	AD	Basal cell carcinoma, medulloblastoma, jaw cysts
Tuberous sclerosis	<i>TSC1</i> <i>TSC2</i>	9q34 16p13.3	AD	Angiofibroma, renal angiomyolipoma
von Hippel-Lindau	<i>VHL</i>	3p25-26	AD	Kidney, cerebellum, pheochromocytoma

Abbreviations: AD, autosomal dominant; AR, autosomal recessive

Although the Mendelian forms of cancer have taught us much about the mechanisms of growth control, most forms of cancer do not follow simple patterns of inheritance. In many instances (e.g., lung cancer), a strong environmental contribution is at work. Even in such circumstances, however, some individuals may be more genetically susceptible to developing cancer, given the appropriate exposure, due to the presence of modifier alleles.

GENETIC TESTING FOR FAMILIAL CANCER

The discovery of cancer susceptibility genes raises the possibility of DNA testing to predict the risk of cancer in individuals of affected families. An algorithm for cancer risk assessment and decision making in high-risk families using genetic testing is shown in Fig. 101e-6. Once a mutation is discovered in a family, subsequent testing of asymptomatic family members can be crucial in patient management. A negative gene test in these individuals can prevent years of anxiety in the knowledge that their cancer risk is no higher than that of the general population. On the other hand, a positive test may lead to alteration of clinical management, such as increased frequency of cancer screening and, when feasible and appropriate, prophylactic surgery. Potential negative consequences of a positive test result include psychological distress (anxiety, depression) and discrimination, although the Genetic Information Nondiscrimination Act (GINA) makes it illegal for predictive genetic information to be used to discriminate in health insurance or employment. Testing should therefore not be conducted without counseling before and after disclosure of the test result. In

addition, the decision to test should depend on whether effective interventions exist for the particular type of cancer to be tested. Despite these caveats, genetic cancer testing for some cancer syndromes already appears to have greater benefits than risks. Companies offer genetic testing for many of the cancer syndromes listed in Table 83-3, including FAP (*APC* gene), hereditary breast and ovarian cancer syndrome (*BRCA1* and *BRCA2* genes), Lynch's syndrome (mismatch repair genes), Li-Fraumeni syndrome (*TP53* gene), Cowden syndrome (*PTEN* gene), hereditary retinoblastoma (*RB1* gene), and others.

Because of the inherent problems of genetic testing such as cost, specificity, and sensitivity, it is not yet appropriate to offer these tests to the general population. However, testing may be appropriate in some subpopulations with a known increased risk, even without a defined family history. For example, two mutations in the breast cancer susceptibility gene *BRCA1*, 185delAG and 5382insC, exhibit a sufficiently high frequency in the Ashkenazi Jewish population that genetic testing of an individual of this ethnic group may be warranted.

As noted above, it is important that genetic test results be communicated to families by trained genetic counselors, especially for high-risk high-penetrance conditions such as the hereditary breast and ovarian cancer syndrome (*BRCA1/BRCA2*). To ensure that the families clearly understand its advantages and disadvantages and the impact it may have on disease management and psyche, genetic testing should never be done before counseling. Significant expertise is needed to communicate the results of genetic testing to families. For example, one common mistake is to misinterpret the result of negative genetic tests. For many cancer predisposition genes, the sensitivity of genetic

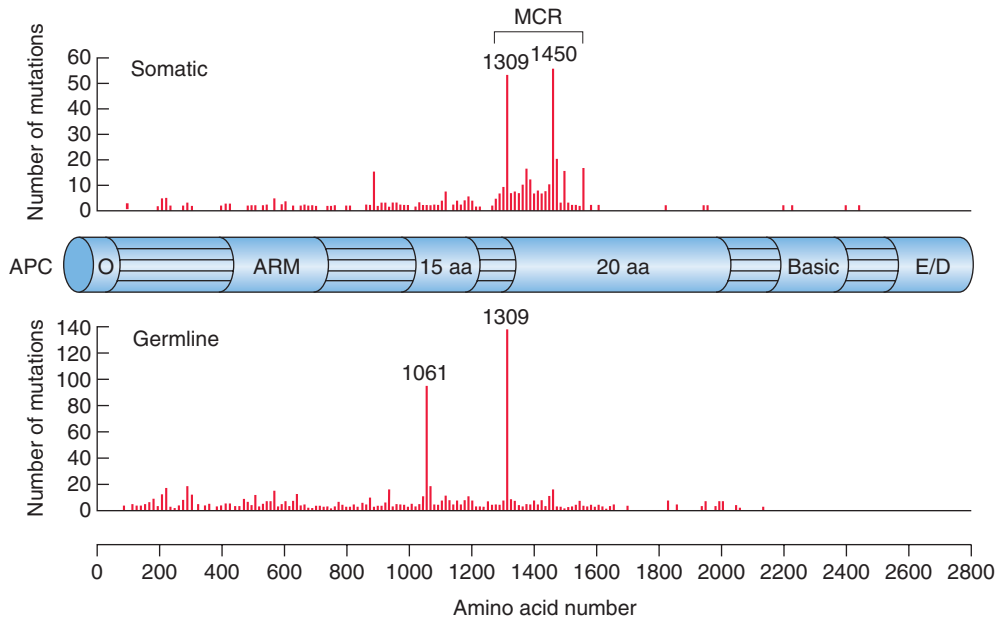


FIGURE 101e-5 Germline and somatic mutations in the tumor-suppressor gene *APC*. *APC* encodes a 2843-amino-acid protein with six major domains: an oligomerization region (O), armadillo repeats (ARM), 15-amino-acid repeats (15 aa), 20-amino-acid repeats (20 aa), a basic region, and a domain involved in binding EB1 and the *Drosophila* discs large homologue (E/D). Shown are the positions within the *APC* gene of a total of 650 somatic and 826 germline mutations (from the *APC* database at <http://www.umd.be/APC>). The vast majority of these mutations result in the truncation of the *APC* protein. Germline mutations are found to be relatively evenly distributed up to codon 1600 except for two mutation hotspots at amino acids 1061 and 1309, which together account for one-third of the mutations found in familial adenomatous polyposis (FAP) families. Somatic *APC* mutations in colon tumors cluster in an area of the gene known as the *mutation cluster region* (MCR). The location of the MCR suggests that the 20-amino-acid domain plays a crucial role in tumor suppression.

testing is less than 70% (i.e., of 100 kindreds tested, disease-causing mutations can be identified in 70 at most). Therefore, such testing should in general begin with an affected member of the kindred (the youngest family member still alive who has had the cancer of interest). If a mutation is not identified in this individual, then the test should be reported as noninformative (Fig. 101e-6) rather than negative (because it is possible that, for technical reasons, the mutation in this individual is not detectable by standard genetic assays). On the other hand, if a mutation can be identified in this individual, then testing of other family members can be performed, and the sensitivity of such subsequent tests will be 100% (because the mutation in the family is in this case known to be detectable by the method used).

MICRORNAs AND CANCER

MicroRNAs (miRNAs) are small noncoding RNAs 20–22 nucleotides in length that are involved in posttranscriptional gene regulation. Studies in chronic lymphocytic leukemia first suggested a link between miRNAs and cancer when *miR-15* and *miR-16* were found to be deleted or downregulated in the vast majority of tumors. Various miRNAs have since been found abnormally expressed in several human malignancies. Aberrant expression of miRNAs in cancer has been attributed to several mechanisms, such as chromosomal rearrangements, genomic copy number change, epigenetic modifications, defects in miRNA biogenesis pathway, and regulation by transcriptional factors. Somatic mutations of miRNAs have been identified in many cancers, but the exact functional consequences of these changes on cancer development remain to be determined. The SomaMir database (<http://compbio.uthsc.edu/SomamiR>) catalogs somatic and germline miRNA mutations that have been identified in cancer.

Functionally, miRNAs have been suggested to contribute to tumorigenesis through their ability to regulate oncogenic signaling pathways. For example, *miR-15* and *miR-16* have been shown to target the *BCL2*

oncogene, leading to its downregulation in leukemic cells and apoptosis. As another example of miRNAs' involvement in oncogenic pathways, the p53 tumor suppressor can transcriptionally induce *miR-34* following genotoxic stress, and this induction is important in mediating p53 function. The expression of miRNAs is extremely specific, and there is evidence that miRNA expression patterns may be useful in distinguishing lineage and differentiation state, as well as cancer diagnosis and outcome prediction.

VIRUSES IN HUMAN CANCER

Certain human malignancies are associated with viruses. Examples include Burkitt's lymphoma (Epstein-Barr virus; Chap. 218), hepatocellular carcinoma (hepatitis viruses), cervical cancer (human papillomavirus [HPV]; Chap. 222), and T cell leukemia (retroviruses; Chap. 225e). The mechanisms of action of these viruses are varied but always involve activation of growth-promoting pathways or inhibition of tumor-suppressor products in the infected cells. For example, HPV proteins E6 and E7 bind and inactivate cellular tumor suppressors p53 and pRB, respectively. There are several HPV types, and some of these types have been associated with the development of several malignancies,

including cervical, vulvar, vaginal, penile, anal, and oropharyngeal cancer. Viruses are not sufficient for cancer development, but constitute one alteration in the multistep process of cancer progression.

GENE EXPRESSION IN CANCER

The tumorigenesis process, driven by alterations in tumor suppressors, oncogenes, and epigenetic regulation, is accompanied by changes in gene expression. The advent of powerful techniques for high-throughput gene expression profiling, based on sequencing or microarrays, has allowed the comprehensive study of gene expression in neoplastic cells. It is indeed possible to identify the expression levels of thousands of genes expressed in normal and cancer tissues. Figure 101e-7 shows a typical microarray experiment examining gene expression in cancer. This global knowledge of gene expression allows the identification of differentially expressed genes and, in principle, the understanding of the complex molecular circuitry regulating normal and neoplastic behaviors. Such studies have led to molecular profiling of tumors, which has suggested general methods for distinguishing tumors of various biologic behaviors (molecular classification), elucidating pathways relevant to the development of tumors, and identifying molecular targets for the detection and therapy of cancer. The first practical applications of this technology have suggested that global gene expression profiling can provide prognostic information not evident from other clinical or laboratory tests. The Gene Expression Omnibus (GEO, <http://www.ncbi.nlm.nih.gov/geo/>) is a searchable online repository for expression profiling data.

GENOMEWIDE MUTATIONAL PROFILING IN CANCER

With the completion of the Human Genome Project and advances in sequencing technologies, systematic mutational analysis of the cancer genome has become possible. In fact, whole genome sequencing of cancer cells is now possible, and this technology has the potential to

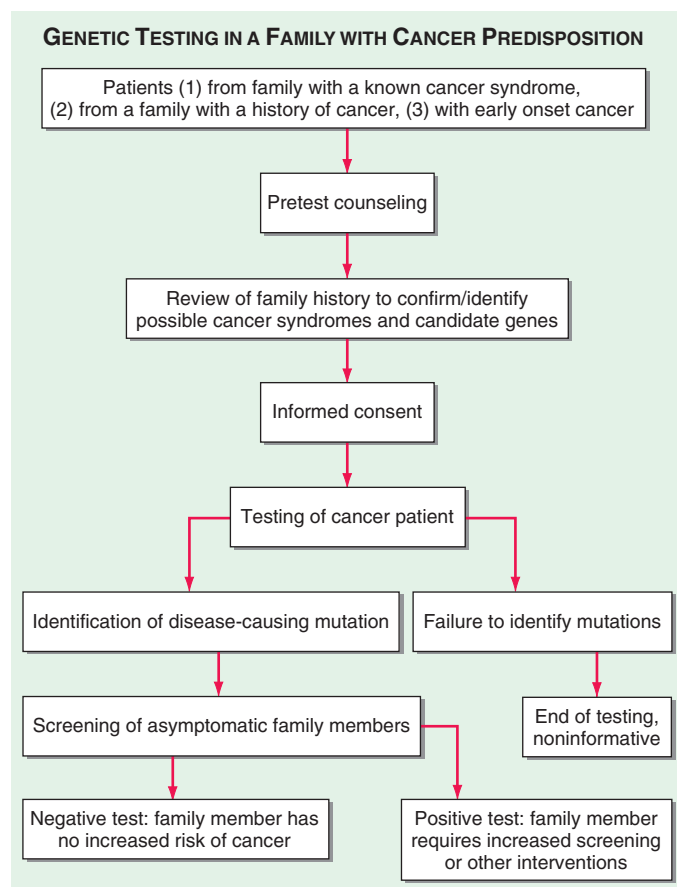


FIGURE 101e-6 Algorithm for genetic testing in a family with cancer predisposition. The key step is the identification of a mutation in a cancer patient, which allows testing of asymptomatic family members. Asymptomatic family members who test positive may require increased screening or surgery, whereas others are at no greater risk for cancer than the general population.

revolutionize our approach to cancer prevention, diagnosis, and treatment. The International Cancer Genome Consortium (<http://icgc.org/>) was developed by leading cancer agencies worldwide, genome and cancer scientists, and statisticians with the goal to launch and coordinate cancer genomics research projects worldwide and to disseminate the data. Hundreds of cancer genomes from at least 25 cancer types have been sequenced through various collaborative efforts. In addition, exome sequencing (sequencing all the coding regions of the genome) has also been performed on a large number of tumors. These sequencing data have been used to elucidate the mutational profile of cancer, including the identification of driver mutations that are functionally involved in tumor development. There are generally 40 to 100 genetic alterations that affect protein sequence in a typical cancer, although statistical analyses suggest that only 8–15 are functionally involved in tumorigenesis. The picture that emerges from these studies is that most genes found mutated in tumors are actually mutated at relatively low frequencies (<5%), whereas a small number of genes (such as *p53*, *KRAS*) are mutated in a large proportion of tumors (Fig. 101e-8). In the past, the focus of research has been on the frequently mutated genes, but it appears that the large number of genes that are infrequently mutated in cancer are major contributors to the cancer phenotype. Understanding the signaling pathways altered by mutations in these genes, as well as the functional relevance of these different mutations, represents the next challenge in the field. Moreover, a detailed knowledge of the genes altered in a particular tumor may allow for a new era of personalized treatment in cancer medicine (see below). A major effort in the United States, The Cancer Genome Atlas (<http://cancergenome.nih.gov>) is a coordinated effort from the National Cancer Institute and the National Human Genome

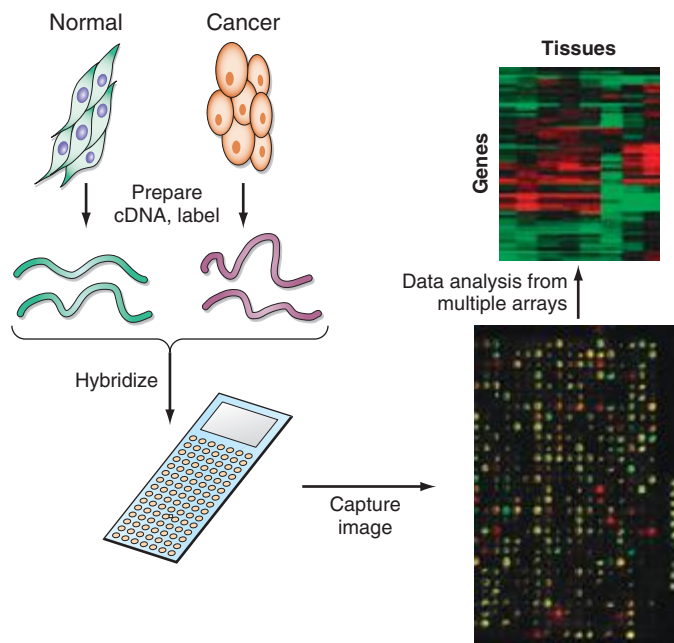


FIGURE 101e-7 A microarray experiment. RNA is prepared from cells, reverse transcribed to cDNA, and labeled with fluorescent dyes (typically green for normal cells and red for cancer cells). The fluorescent probes are mixed and hybridized to a cDNA array. Each spot on the array is an oligonucleotide (or cDNA fragment) that represents a different gene. The image is then captured with a fluorescence camera; red spots indicate higher expression in tumor cells compared with reference, while green spots represent the lower expression in tumor cells. Yellow signals indicate equal expression levels in normal and tumor specimens. After clustering analysis of multiple arrays, the results are typically represented graphically using a visualization software, which shows, for each sample, a color-coded representation of gene expression for every gene on the array.

Research Institute to systematically characterize the entire spectrum of genomic changes involved in human cancers. Similarly, COSMIC (Catalogue of Somatic Mutations in Cancer) is an initiative from the Wellcome Trust Sanger Institute to store and display somatic mutation information and related details regarding human cancers (<http://cancer.sanger.ac.uk/>).

PERSONALIZED CANCER TREATMENT BASED ON MOLECULAR PROFILES: PRECISION THERAPY

Gene expression profiling and genomewide sequencing approaches have allowed for an unprecedented understanding of cancer at the molecular level. It has been suggested that individualized knowledge of pathways or genes deregulated in a given tumor (personalized genomics) may provide a guide for therapeutic options on the tumor, thus leading to personalized therapy (also called precision medicine). Because tumor behavior is highly heterogeneous, even within a tumor type, personalized information-based medicine will likely supplement or perhaps one day supplant the current histology-based therapy, especially in the case of tumors resistant to conventional therapeutic approaches. Molecular nosology has revealed similarities in tumors of diverse histotype. The success of this approach will be dependent on the identification of sufficient actionable changes (mutations or pathways that can be targeted with a specific drug). Examples of currently actionable changes include mutations in *BRAF* (targeted by the drug vemurafenib) and *RET* (targeted by sunitinib and sorafenib), and *ALK* rearrangements (targeted by crizotinib). Interestingly, studies have reported that 20% of triple-negative breast cancers and 60% of lung cancers have potentially actionable genetic changes. Gene expression also offers the potential to predict drug sensitivities as well as provide prognostic information. Commercial diagnostic tests, such as Mammprint and Oncotype DX for breast cancer, are available

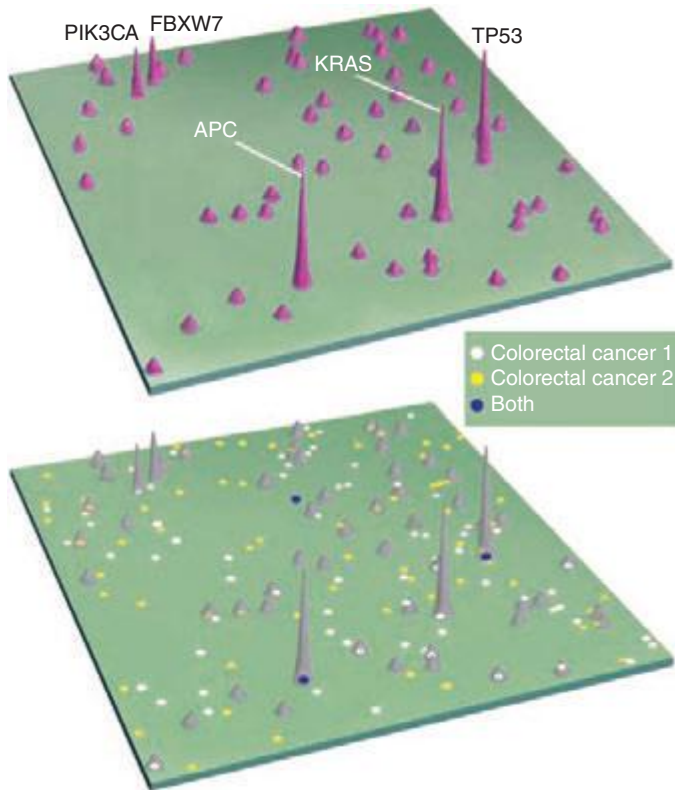


FIGURE 101e-8 A two-dimensional map of genes mutated in colorectal cancer. The two-dimensional landscape represents the positions of the RefSeq genes along the chromosomes and the height of the peaks represents the mutation frequency. On the top map, the taller peaks represent the genes that are commonly mutated in colon cancer, while the large number of smaller hills indicates the genes that are mutated at lower frequency. On the lower map, the mutations of two individual tumors are indicated. Note that there is little overlap between the mutated genes of the two colorectal tumors shown. These differences may represent the basis for the heterogeneity in terms of behavior and responsiveness to therapy observed in human cancer. (From LD Wood et al: *Science* 318:1108, 2007, with permission.)

to help the patients and their physicians make treatment decisions. Personalized medicine is an exciting new avenue for cancer treatment based on matching the unique features of a tumor to an effective therapy, and this concept is in the process of changing our approach to cancer therapy in fundamental ways. On a cautionary note, gene expression can vary enormously within a single person's cancer and at different anatomic sites in the patient. We have not yet determined whether such clonal variation within an individual tumor will interfere with the goal tailoring therapy to a particular patient's tumor.

THE FUTURE

A revolution in cancer genetics has occurred in the past 25 years. Identification of cancer genes has led to a deep understanding of the tumorigenesis process and has had important repercussions on all fields of cancer biology. In particular, the advancement of powerful techniques for genomewide expression profiling and mutation analyses has provided a detailed picture of the molecular defects present in individual tumors. Individualized treatment based on the specific genetic alterations within a given tumor has already become possible. Although these advances have not yet translated into overall changes in cancer prevention, prognosis, or treatment, it is expected that breakthroughs in these areas will continue to emerge and be applicable to an ever-increasing number of cancers.

Cancers are characterized by unregulated cell division, avoidance of cell death, tissue invasion, and the ability to metastasize. A neoplasm is *benign* when it grows in an unregulated fashion without tissue invasion. The presence of unregulated growth and tissue invasion is characteristic of *malignant* neoplasms. Cancers are named based on their origin: those derived from epithelial tissue are called *carcinomas*, those derived from mesenchymal tissues are *sarcomas*, and those derived from hematopoietic tissue are *leukemias*, *lymphomas*, and *plasma cell dyscrasias* (including *multiple myeloma*).

Cancers nearly always arise as a consequence of genetic alterations, the vast majority of which begin in a single cell and therefore are monoclonal in origin. However, because a wide variety of genetic and epigenetic changes can occur in different cells within malignant tumors over time, most cancers are characterized by marked heterogeneity in the populations of cells. This heterogeneity significantly complicates the treatment of most cancers because it is likely that there are subsets of cells that will be resistant to therapy and will therefore survive and proliferate even if the majority of cells are killed.

A few cancers appear to, at least initially, be primarily driven by an alteration in a dominant gene that produces uncontrolled cell proliferation. Examples include chronic myeloid leukemia (*abl*), about half of melanomas (*braf*), Burkitt's lymphoma (*c-myc*), and subsets of lung adenocarcinomas (*egfr*, *alk*, *ros1*, and *ret*). The genes that can promote cell growth when altered are often called *oncogenes*. They were first identified as critical elements of viruses that cause animal tumors; it was subsequently found that the viral genes had normal counterparts with important functions in the cell and had been captured and mutated by viruses as they passed from host to host.

However, the vast majority of human cancers are characterized by a multiple-step process involving many genetic abnormalities, each of which contributes to the loss of control of cell proliferation and differentiation and the acquisition of capabilities, such as tissue invasion, the ability to metastasize, and angiogenesis. These properties are not found in the normal adult cell from which the tumor is derived. Indeed, normal cells have a large number of safeguards against uncontrolled proliferation and invasion. Many cancers go through recognizable steps of progressively more abnormal phenotypes: hyperplasia, to adenoma, to dysplasia, to carcinoma in situ, to invasive cancer with the ability to metastasize (Table 102e-1). For most cancers, these changes occur over a prolonged period of time, usually many years.

In most organs, only primitive undifferentiated cells are capable of proliferating and the cells lose the capacity to proliferate as they differentiate and acquire functional capability. The expansion of the primitive cells is linked to some functional need in the host through receptors that receive signals from the local environment or through hormonal and other influences delivered by the vascular supply. In the absence of such signals, the cells are at rest. The signals that keep the primitive cells at rest remain incompletely understood. These signals must be environmental, based on the observations that a regenerating liver stops growing when it has replaced the portion that has been surgically removed after partial hepatectomy and regenerating bone marrow stops growing when the peripheral blood counts return to normal. Cancer cells clearly have lost responsiveness to such controls and do not recognize when they have overgrown the niche normally occupied by the organ from which they are derived. A better understanding of the mechanisms of growth regulation is evolving.

CELL CYCLE CHECKPOINTS

Normal cells have a number of control mechanisms that are targeted by specific genetic alterations in cancer. Critical proteins in these control processes that are frequently mutated or otherwise inactivated in cancers are called tumor-suppressor genes. Examples include p53

TABLE 102e-1 PHENOTYPIC CHARACTERISTICS OF MALIGNANT CELLS

Deregulated cell proliferation: Loss of function of negative growth regulators (tumor-suppressor genes, i.e., *Rb*, *p53*), and increased action of positive growth regulators (oncogenes, i.e., *Ras*, *Myc*). Leads to aberrant cell cycle control and includes loss of normal checkpoint responses.

Failure to differentiate: Arrest at a stage before terminal differentiation. May retain stem cell properties. (Frequently observed in leukemias due to transcriptional repression of developmental programs by the gene products of chromosomal translocations.)

Loss of normal apoptosis pathways: Inactivation of p53, increases in Bcl-2 family members. This defect enhances the survival of cells with oncogenic mutations and genetic instability and allows clonal expansion and diversification within the tumor without activation of physiologic cell death pathways.

Genetic instability: Defects in DNA repair pathways leading to either single-nucleotide or oligonucleotide mutations (as in microsatellite instability, MIN) or more commonly chromosomal instability (CIN) leading to aneuploidy. Caused by loss of function of *p53*, *BRCA1/2*, mismatch repair genes, DNA repair enzymes, and the spindle checkpoint. Leads to accumulation of a variety of mutations in different cells within the tumor and heterogeneity.

Loss of replicative senescence: Normal cells stop dividing in vitro after 25–50 population doublings. Arrest is mediated by the *Rb*, *p16^{INK4a}*, and *p53* pathways. Further replication leads to telomere loss, with crisis. Surviving cells often harbor gross chromosomal abnormalities. Relevance to human in vivo cancer remains uncertain. Many human cancers express telomerase.

Nonresponsiveness to external growth-inhibiting signals: Cancer cells have lost responsiveness to signals normally present to stop proliferating when they have overgrown the niche normally occupied by the organ from which they are derived. We know very little about this mechanism of growth regulation.

Increased angiogenesis: Due to increased gene expression of proangiogenic factors (VEGF, FGF, IL-8) by tumor or stromal cells, or loss of negative regulators (endostatin, tumstatin, thrombospondin).

Invasion: Loss of cell-cell contacts (gap junctions, cadherins) and increased production of matrix metalloproteinases (MMPs). Often takes the form of epithelial-to-mesenchymal transition (EMT), with anchored epithelial cells becoming more like motile fibroblasts.

Metastasis: Spread of tumor cells to lymph nodes or distant tissue sites. Limited by the ability of tumor cells to survive in a foreign environment.

Evasion of the immune system: Downregulation of MHC class I and II molecules; induction of T cell tolerance; inhibition of normal dendritic cell and/or T cell function; antigenic loss variants and clonal heterogeneity; increase in regulatory T cells.

Shift in cell metabolism: Energy generation shifts to aerobic glycolysis.

Abbreviations: FGF, fibroblast growth factor; IL, interleukin; MHC, major histocompatibility complex; VEGF, vascular endothelial growth factor.

and *Rb* (discussed below). The progression of a cell through the cell division cycle is regulated at a number of checkpoints by a wide array of genes. In the first phase, G_1 , preparations are made to replicate the genetic material. The cell stops before entering the DNA synthesis phase, or S phase, to take inventory. Are we ready to replicate our DNA? Is the DNA repair machinery in place to fix any mutations that are detected? Are the DNA replicating enzymes available? Is there an adequate supply of nucleotides? Is there sufficient energy? The main brake on the process is the retinoblastoma protein, *Rb*. When the cell determines that it is prepared to move ahead, sequential activation of cyclin-dependent kinases (CDKs) results in the inactivation of the brake, *Rb*, by phosphorylation. Phosphorylated *Rb* releases the S phase-regulating transcription factor, E2F/DP1, and genes required for S phase progression are expressed. If the cell determines that it is unready to move ahead with DNA replication, a number of inhibitors are capable of blocking the action of the CDKs, including *p21^{Cip2/Waf1}*, *p16^{INK4a}*, and *p27^{Kip1}*. *Nearly every cancer has one or more genetic lesions in the G_1 checkpoint that permits progression to S phase.*

At the end of S phase, when the cell has exactly duplicated its DNA content, a second inventory is taken at the S checkpoint. Have all of the chromosomes been fully duplicated? Were any segments of DNA copied more than once? Do we have the right number of chromosomes and the right amount of DNA? If so, the cell proceeds to G_2 , in which the cell prepares for division by synthesizing mitotic spindle and other proteins needed to produce two daughter cells. When DNA damage is

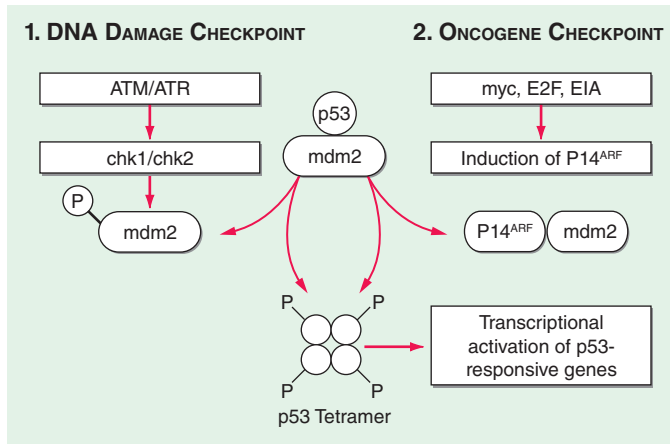


FIGURE 102e-1 Induction of p53 by the DNA damage and oncogene checkpoints. In response to noxious stimuli, p53 and mdm2 are phosphorylated by the ataxia-telangiectasia mutated (ATM) and related (ATR) serine/threonine kinases, as well as the immediate downstream checkpoint kinases, Chk1 and Chk2. This causes dissociation of p53 from mdm2, leading to increased p53 protein levels and transcription of genes leading to cell cycle arrest (p21^{Cip1/Waf1}) or apoptosis (e.g., the proapoptotic Bcl-2 family members Noxa and Puma). Inducers of p53 include hypoxemia, DNA damage (caused by ultraviolet radiation, gamma irradiation, or chemotherapy), ribonucleotide depletion, and telomere shortening. A second mechanism of p53 induction is activated by oncogenes such as *Myc*, which promote aberrant G₁/S transition. This pathway is regulated by a second product of the *Ink4a* locus, p14^{ARF} (p19 in mice), which is encoded by an alternative reading frame of the same stretch of DNA that codes for p16^{Ink4a}. Levels of ARF are upregulated by *Myc* and E2F, and ARF binds to mdm2 and rescues p53 from its inhibitory effect. This *oncogene checkpoint* leads to the death or senescence (an irreversible arrest in G₁ of the cell cycle) of renegade cells that attempt to enter S phase without appropriate physiologic signals. Senescent cells have been identified in patients whose premalignant lesions harbor activated oncogenes, for instance, dysplastic nevi that encode an activated form of BRAF (see below), demonstrating that induction of senescence is a protective mechanism that operates in humans to prevent the outgrowth of neoplastic cells.

detected, the p53 pathway is normally activated. Called the guardian of the genome, p53 is a transcription factor that is normally present in the cell in very low levels. Its level is generally regulated through its rapid turnover. Normally, p53 is bound to mdm2, a ubiquitin ligase, that both inhibits p53 transcriptional activation and also targets p53 for degradation in the proteasome. When damage is sensed, the ATM (ataxia-telangiectasia mutated) pathway is activated; ATM phosphorylates mdm2, which no longer binds to p53, and p53 then stops cell cycle progression, directs the synthesis of repair enzymes, or if the damage is too great, initiates apoptosis of the cell to prevent the propagation of a damaged cell (Fig. 102e-1).

A second method of activating p53 involves the induction of p14^{ARF} by hyperproliferative signals from oncogenes. p14^{ARF} competes with p53 for binding to mdm2, allowing p53 to escape the effects of mdm2 and accumulate in the cell. Then p53 stops cell cycle progression by activating CDK inhibitors such as p21 and/or initiating the apoptosis pathway. Not surprisingly given its critical role in controlling cell cycle progression, mutations in the gene for p53 on chromosome 17p are found in more than 50% of human cancers. Most commonly these mutations are acquired in the malignant tissue in one allele and the second allele is deleted, leaving the cell unprotected from DNA-damaging agents or oncogenes. Some environmental exposures produce signature mutations in p53; for example, aflatoxin exposure leads to mutation of arginine to serine at codon 249 and leads to hepatocellular carcinoma. In rare instances, p53 mutations are in the germline (Li-Fraumeni syndrome) and produce a familial cancer syndrome.

The absence of p53 leads to chromosome instability and the accumulation of DNA damage including the acquisition of properties that give the abnormal cell a proliferative and survival advantage. Like *Rb dysfunction*, most cancers have mutations that disable the p53 pathway. Indeed, the importance of p53 and Rb in the development of cancer is underscored by the neoplastic transformation mechanism of human papillomavirus. This virus has two main oncogenes, E6 and E7. E6 acts to increase the rapid turnover of p53, and E7 acts to inhibit Rb function; inhibition of these two targets is required for transformation of epithelial cells.

Another cell cycle checkpoint exists when the cell is undergoing division, the spindle checkpoint. The details of this checkpoint are still being discovered; however, it appears that if the spindle apparatus does not properly align the chromosomes for division, if the chromosome number is abnormal (i.e., greater or less than 4*n*), or if the centromeres are not properly paired with their duplicated partners, then the cell initiates a cell death pathway to prevent the production of aneuploid progeny (having an altered number of chromosomes). Abnormalities in the spindle checkpoint facilitate the development of aneuploidy. In some tumors, aneuploidy is a predominant genetic feature. In others, a defect in the cells' ability to repair errors in the DNA due to mutations in genes coding for the proteins critical for mismatched DNA repair is the primary genetic lesion. This is usually detected by finding alterations in repeat sequences of DNA (called microsatellites) or microsatellite instability in malignant cells. In general, tumors either have defects in chromosome number or microsatellite instability, but not both. Defects that lead to cancer include abnormal cell cycle checkpoints, inadequate DNA repair, and failure to preserve genome integrity.

Efforts are under way to therapeutically restore the defects in cell cycle regulation that characterize cancer, although this remains a challenging problem because it is much more difficult to restore normal biologic function than to inhibit abnormal function of proteins driving cell proliferation, such as oncogenes.

CANCER AS AN ORGAN THAT IGNORES ITS NICHE

The fundamental cellular defects that create a malignant neoplasm act at the cellular level. However, that is not the entire story. Cancers behave as organs that have lost their specialized function and stopped responding to signals that normally limit their growth. Human cancers usually become clinically detectable when a primary mass is at least 1 cm in diameter—such a mass consists of about 10⁹ cells. More commonly, patients present with tumors that are 10¹⁰ cells or greater. A lethal tumor burden is about 10¹² to 10¹³ cells. If all tumor cells were dividing at the time of diagnosis, patients would reach a lethal tumor burden in a very short time. However, human tumors grow by Gompertzian kinetics—this means that not every daughter cell produced by a cell division is itself capable of dividing. The growth fraction of a tumor declines exponentially with time. The growth fraction of the first malignant cell is 100%, and by the time a patient presents for medical care, the growth fraction is 2–3% or less. This fraction is similar to the growth fraction of normal bone marrow and normal intestinal epithelium, the most highly proliferative normal tissues in the human body, a fact that may explain the dose-limiting toxicities of agents that target dividing cells.

The implication of these data is that the tumor is slowing its own growth over time. How does it do this? The tumor cells have multiple genetic lesions that tend to promote proliferation, yet by the time the tumor is clinically detectable, its capacity for proliferation has declined. We need to better understand how a tumor slows its own growth. A number of factors are known to contribute to the failure of tumor cells to proliferate in vivo. Some cells are hypoxic and have inadequate supply of nutrients and energy. Some have sustained too much genetic damage to complete the cell cycle but have lost the capacity to undergo apoptosis and therefore survive but do not proliferate. However, an important subset is not actively dividing but retains the capacity to divide and can start dividing again under certain conditions such as when the tumor mass is reduced by treatments. Just as the bone marrow increases its rate of proliferation in response to bone

marrow-damaging agents, the tumor also seems to sense when tumor cell numbers have been reduced and can respond by increasing growth rate. However, the critical difference is that the marrow stops growing when it has reached its production goals, whereas tumors do not.

Additional tumor cell vulnerabilities are likely to be detected when we learn more about how normal cells respond to “stop” signals from their environment and why and how tumor cells fail to heed such signals.

IS IN VITRO SENESCENCE RELEVANT TO CARCINOGENESIS?

When normal cells are placed in culture *in vitro*, most are not capable of sustained growth. Fibroblasts are an exception to this rule. When they are cultured, fibroblasts may divide 30–50 times and then they undergo what has been termed a “crisis” during which the majority of cells stop dividing (usually due to an increase in p21 expression, a CDK inhibitor), many die, and a small fraction emerge that have acquired genetic changes that permit their uncontrolled growth. The cessation of growth of normal cells in culture has been termed “senescence,” and whether this phenomenon is relevant to any physiologic event *in vivo* is debated.

Among the cellular changes during *in vitro* propagation is telomere shortening. DNA polymerase is unable to replicate the tips of chromosomes, resulting in the loss of DNA at the specialized ends of chromosomes (called *telomeres*) with each replication cycle. At birth, human telomeres are 15- to 20-kb pairs long and are composed of tandem repeats of a six-nucleotide sequence (TTAGGG) that associates with specialized telomere-binding proteins to form a T-loop structure that protects the ends of chromosomes from being mistakenly recognized as damaged. The loss of telomeric repeats with each cell division cycle causes gradual telomere shortening, leading to growth arrest (called *senescence*) when one or more critically short telomeres trigger a p53-regulated DNA-damage checkpoint response. Cells can bypass this growth arrest if pRb and p53 are nonfunctional, but cell death usually ensues when the unprotected ends of chromosomes lead to chromosome fusions or other catastrophic DNA rearrangements. *The ability to bypass telomere-based growth limitations is thought to be a critical step in the evolution of most malignancies.* This occurs by the reactivation of telomerase expression in cancer cells. Telomerase is an enzyme that adds TTAGGG repeats onto the 3′ ends of chromosomes. It contains a catalytic subunit with reverse transcriptase activity (hTERT) and an RNA component that provides the template for telomere extension. Most normal somatic cells do not express sufficient telomerase to prevent telomere attrition with each cell division. Exceptions include stem cells (such as those found in hematopoietic tissues, gut and skin epithelium, and germ cells) that require extensive cell division to maintain tissue homeostasis. More than 90% of human cancers express high levels of telomerase that prevent telomere shortening to critical levels and allow indefinite cell proliferation. *In vitro* experiments indicate that inhibition of telomerase activity leads to tumor cell apoptosis. Major efforts are under way to develop methods to inhibit telomerase activity in cancer cells. For example, the protein component of telomerase (hTERT) may act as one of the most widely expressed tumor-associated antigens and be targeted by vaccine approaches.

Although most of the functions of telomerase relate to cell division, it also has several other effects including interfering with the differentiated functions of at least certain stem cells, although the impact on differentiated function of normal non-stem cells is less clear. Nevertheless, a major growth industry in medical research has been discovering an association between short telomeres and human diseases ranging from diabetes and coronary artery disease to Alzheimer’s disease. The picture is further complicated by the fact that rare genetic defects in the telomerase enzyme seem to cause pulmonary fibrosis, aplastic anemia, or dyskeratosis congenita (characterized by abnormalities in skin, nails, and oral mucosa with increased risk for certain malignancies) but not defects in nutrient absorption in the gut, a site that might be presumed to be highly sensitive to defective cell proliferation. Much remains to be learned about how telomere shortening and telomere maintenance are related to human illness in general and cancer in particular.

SIGNAL TRANSDUCTION PATHWAYS IN CANCER CELLS

Signals that affect cell behavior come from adjacent cells, the stroma in which the cells are located, hormonal signals that originate remotely, and from the cells themselves (autocrine signaling). These signals generally exert their influence on the receiving cell through activation of signal transduction pathways that have as their end result the induction of activated transcription factors that mediate a change in cell behavior or function or the acquisition of effector machinery to accomplish a new task. Although signal transduction pathways can lead to a wide variety of outcomes, many such pathways rely on cascades of signals that sequentially activate different proteins or glycoproteins and lipids or glycolipids, and the activation steps often involve the addition or removal of one or more phosphate groups on a downstream target. Other chemical changes can result from signal transduction pathways, but phosphorylation and dephosphorylation play a major role. The proteins that add phosphate groups to proteins are called kinases. There are two major distinct classes of kinases; one class acts on tyrosine residues, and the other acts on serine/threonine residues. The tyrosine kinases often play critical roles in signal transduction pathways; they may be receptor tyrosine kinases, or they may be linked to other cell-surface receptors through associated docking proteins (Fig. 102e-2).

Normally, tyrosine kinase activity is short-lived and reversed by protein tyrosine phosphatases (PTPs). However, in many human cancers, tyrosine kinases or components of their downstream pathways are activated by mutation, gene amplification, or chromosomal translocations. Because these pathways regulate proliferation, survival, migration, and angiogenesis, they have been identified as important targets for cancer therapeutics.

Inhibition of kinase activity is effective in the treatment of a number of neoplasms. Lung cancers with mutations in the epidermal growth factor receptor are highly responsive to erlotinib and gefitinib (Table 102e-2). Lung cancers with activation of anaplastic lymphoma kinase (ALK) or ROS1 by translocations respond to crizotinib, an ALK and ROS1 inhibitor. A BRAF inhibitor is highly effective in melanomas and thyroid cancers in which BRAF is mutated. Targeting a protein (MEK) downstream of BRAF also has activity against BRAF mutant melanomas. Janus kinase inhibitors are active in myeloproliferative syndromes in which JAK2 activation is a pathogenetic event. Imatinib (which targets a number of tyrosine kinases) is an effective agent in tumors that have translocations of the *c-Abl* and BCR gene (such as chronic myeloid leukemia), mutant *c-Kit* (gastrointestinal stromal cell tumors), or mutant platelet-derived growth factor receptor (PDGFR; chronic myelomonocytic leukemia); second-generation inhibitors of BCR-Abl, dasatinib, and nilotinib are even more effective. The third-generation agent bosutinib has activity in some patients who have progressed on other inhibitors, whereas the third-generation agent ponatinib has activity against the T315I mutation, which is resistant to the other agents. Sorafenib and sunitinib, agents that inhibit a large number of kinases, have shown antitumor activity in a number of malignancies, including renal cell cancer (RCC) (both), hepatocellular carcinoma (sorafenib), thyroid cancer (sorafenib), gastrointestinal stromal tumor (GIST) (sunitinib), and pancreatic neuroendocrine tumors (sunitinib). Inhibitors of the mammalian target of rapamycin (mTOR) are active in RCC, pancreatic neuroendocrine tumors, and breast cancer. The list of active agents and treatment indications is growing rapidly. These new agents have ushered in a new era of personalized therapy. It is becoming more routine for resected tumors to be assessed for specific molecular changes that predict response and to have clinical decision-making guided by those results.

However, none of these therapies has yet been curative by themselves for any malignancy, although prolonged periods of disease control lasting many years frequently occur in chronic myeloid leukemia. The reasons for the failure to cure are not completely defined, although resistance to the treatment ultimately develops in most patients. In some tumors, resistance to kinase inhibitors is related to an acquired mutation in the target kinase that inhibits drug binding. Many of these kinase inhibitors act as competitive inhibitors of the ATP-binding pocket. ATP is the phosphate donor in these phosphorylation

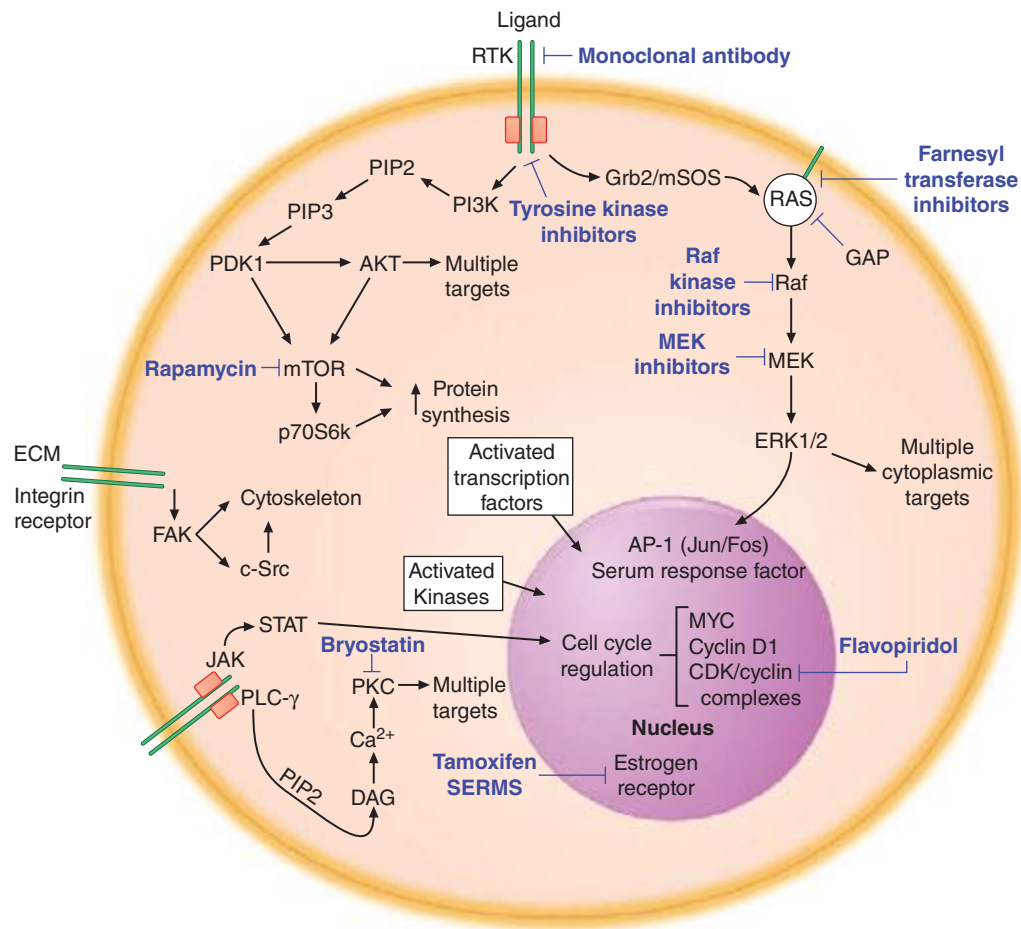


FIGURE 102e-2 Therapeutic targeting of signal transduction pathways in cancer cells. Three major signal transduction pathways are activated by receptor tyrosine kinases (RTK). **1.** The protooncogene Ras is activated by the Grb2/mSOS guanine nucleotide exchange factor, which induces an association with Raf and activation of downstream kinases (MEK and ERK1/2). **2.** Activated PI3K phosphorylates the membrane lipid PIP₂ to generate PIP₃, which acts as a membrane-docking site for a number of cellular proteins including the serine/threonine kinases PDK1 and Akt. PDK1 has numerous cellular targets, including Akt and mTOR. Akt phosphorylates target proteins that promote resistance to apoptosis and enhance cell cycle progression, whereas mTOR and its target p70S6k upregulate protein synthesis to potentiate cell growth. **3.** Activation of PLC- γ leads to the formation of diacylglycerol (DAG) and increased intracellular calcium, with activation of multiple isoforms of PKC and other enzymes regulated by the calcium/calmodulin system. Other important signaling pathways involve non-RTKs that are activated by cytokine or integrin receptors. Janus kinases (JAK) phosphorylate STAT (signal transducer and activator of transcription) transcription factors, which translocate to the nucleus and activate target genes. Integrin receptors mediate cellular interactions with the extracellular matrix (ECM), inducing activation of FAK (focal adhesion kinase) and c-Src, which activate multiple downstream pathways, including modulation of the cell cytoskeleton. Many activated kinases and transcription factors migrate into the nucleus, where they regulate gene transcription, thus completing the path from extracellular signals, such as growth factors, to a change in cell phenotype, such as induction of differentiation or cell proliferation. The nuclear targets of these processes include transcription factors (e.g., Myc, AP-1, and serum response factor) and the cell cycle machinery (CDKs and cyclins). Inhibitors of many of these pathways have been developed for the treatment of human cancers. Examples of inhibitors that are currently being evaluated in clinical trials are shown in purple type.

reactions. Mutation in the BCR-ABL kinase in the ATP-binding pocket (such as the threonine to isoleucine change at codon 315 [T315I]) can prevent imatinib binding. Other resistance mechanisms include altering other signal transduction pathways to bypass the inhibited pathway. As resistance mechanisms become better defined, rational strategies to overcome resistance will emerge. In addition, many kinase inhibitors are less specific for an oncogenic target than was hoped, and toxicities related to off-target inhibition of kinases limit the use of the agent at a dose that would optimally inhibit the cancer-relevant kinase.

Targeted agents can also be used to deliver highly toxic compounds. An important component of the technology for developing effective conjugates is the design of the linker between the two, which needs to be stable. Currently approved antibody drug conjugates include brentuximab vedotin, which links the microtubule toxin monomethyl auristatin E (MMAE) to an antibody targeting the cell surface antigen CD30, which is expressed on a number of malignant cells but especially in Hodgkin's disease and anaplastic lymphoma. The linker in this

case is cleavable, which allows diffusion of the drug out of the cell after delivery. The second approved conjugate is ado-trastuzumab emtansine, which links the microtubule formation inhibitor mertansine and the monoclonal antibody trastuzumab targeted against human epidermal growth factor receptor 2 (HER2) on breast cancer cells. In this case, the linker is noncleavable, thus trapping the chemotherapeutic agent within the cells. There are theoretical pluses and minuses to having either cleavable or noncleavable linkers, and it is likely that both will be used in future developments of antibody-drug conjugates.

Another strategy to enhance the antitumor effects of targeted agents is to use them in rational combinations with each other and in empiric combinations with chemotherapy agents that kill cells in ways distinct from targeted agents. Combinations of trastuzumab (a monoclonal antibody that targets the HER2 receptor [member of the epidermal growth factor receptor (EGFR) family]) with chemotherapy have significant activity against breast and stomach cancers that have high levels of expression of the HER2 protein. The activity of trastuzumab

TABLE 102e-2 SOME FDA-APPROVED MOLECULARLY TARGETED AGENTS FOR THE TREATMENT OF CANCER

Drug	Molecular Target	Disease	Mechanism of Action
All- <i>trans</i> retinoic acid	PML-RAR α oncogene	Acute promyelocytic leukemia M3 AML; t(15;17)	Inhibits transcriptional repression by PML-RAR α
Imatinib	Bcr-Abl, c-Abl, c-Kit, PDGFR- α/β	Chronic myeloid leukemia; GIST	Blocks ATP binding to tyrosine kinase active site
Dasatinib, nilotinib, ponatinib, bosutinib	Bcr-Abl (primarily)	Chronic myeloid leukemia	Blocks ATP binding to tyrosine kinase active site
Sunitinib	c-Kit, VEGFR-2, PDGFR- β , Flt-3	GIST; renal cell cancer	Inhibits activated c-Kit and PDGFR in GIST; inhibits VEGFR in RCC
Sorafenib	RAF, VEGFR-2, PDGFR- α/β , Flt-3, c-Kit	RCC; hepatocellular carcinoma; TC	Targets VEGFR pathways in RCC. Possible activity against BRAF in thyroid cancer
Regorafenib	VEGFR-1 to -3, TIE-2, FGFR1, KIT, RET, PDGFR	Colorectal cancer; GIST	Competitive inhibitor of ATP binding site of tyrosine kinase domain multiple kinases
Axitinib	VEGFR-1 to -3	RCC	Competitive inhibitor of ATP binding site of tyrosine kinase domain VEGF receptors
Erlotinib	EGFR	Non-small-cell lung cancer; pancreatic cancer	Competitive inhibitor of the ATP-binding site of the EGFR
Afatinib	EGFR (and other HER family)	Non-small-cell lung cancer	Irreversible inhibitor of ATP-binding site of HER family members
Lapatinib	HER2/neu	Breast cancer	Competitive inhibitor of the ATP binding site of HER2
Crizotinib (Xalkori)	ALK	Non-small-cell lung cancer	Inhibitor of ALK tyrosine kinase
Bortezomib, carfilzomib	Proteasome	Multiple myeloma	Inhibits proteolytic degradation of multiple cellular proteins
Vemurafenib, dabrafenib	BRAF	Melanoma	Inhibitor of serine-threonine kinase domain of V600E mutant of BRAF
Trametinib	MEK	Melanoma	Inhibitor of serine-threonine kinase domain of V600E mutant of MEK
Cabozantinib	RET, MET, VEGFR	MTC	Competitive inhibitor of ATP binding site of tyrosine kinase domain multiple kinases
Vandetanib	RET, VEGFR, EGFR	MTC	Competitive inhibitor of ATP binding site of tyrosine kinase domain multiple kinases
Temsirolimus	mTOR	RCC	Competitive inhibitor of mTOR serine-threonine kinase
Everolimus	mTOR	RCC; breast cancer	Binds to immunophilin FK binding protein-12, which forms complex that inhibits mTOR kinase
Vorinostat, romidepsin	HDAC	CTCL	HDAC inhibitor
Ruxolitinib	JAK-1, -2	Myelofibrosis	Competitive inhibitor of tyrosine kinase
Vismodegib	Hedgehog pathway	Basal cell cancer (skin)	Inhibits smoothed in hedgehog pathway
Monoclonal Antibodies Alone			
Trastuzumab	HER2/neu (ERBB2)	Breast cancer	Binds HER2 on tumor cell surface and induces receptor internalization
Pertuzumab	HER2/neu (ERBB2)	Breast cancer	Binds HER2 on tumor cell surface at distinct site from trastuzumab and prevents binding to other receptors
Cetuximab	EGFR	Colon cancer, squamous cell carcinoma of the head and neck	Binds extracellular domain of EGFR and blocks binding of EGF and TGF- α ; induces receptor internalization; potentiates the efficacy of chemotherapy and radiotherapy
Panitumumab	EGFR	Colon cancer	Similar to cetuximab but fully humanized rather than chimeric
Rituximab	CD20	B cell lymphomas and leukemias that express CD20	Multiple potential mechanisms, including direct induction of tumor cell apoptosis and immune mechanisms
Alemtuzumab	CD52	Chronic lymphocytic leukemia and CD52-expressing lymphoid tumors	Immune mechanisms
Bevacizumab	VEGF	Colorectal, lung cancers, RCC, glioblastoma, cervical cancer	Inhibits angiogenesis by high-affinity binding to VEGF
Ziv-aflibercept	VEGF-A, VEGF-B, PLGF	Colorectal cancers	Inhibits angiogenesis by high-affinity binding to VEGF-A, VEGF-B, and PLGF
Ipilimumab	CTLA-4	Melanoma	Blocks CTLA-4, preventing interaction with CD80/86 and T cell inhibition
Denosumab	RANK ligand	Breast, prostate cancer	Inhibits RANK ligand, the primary signal for bone removal
Pembrolizumab	PD-1	Melanoma	Blocks PD-1 preventing interaction with PD-L1 T cell inhibition
Antibody-Chemotherapy Conjugates			
Brentuximab vedotin	CD30	Hodgkin's disease, anaplastic lymphoma	Delivery of chemotherapeutic agent (MMAE) to CD30-expressing tumor cells
Ado-trastuzumab emtansine	HER2	Breast cancer	Delivery of chemotherapeutic agent emtansine to HER2-expressing breast cancer cells

Abbreviations: AML, acute myeloid leukemia; CTCL, cutaneous T cell lymphoma; EGFR, epidermal growth factor receptor; FDA, Food and Drug Administration; Flt-3, fms-like tyrosine kinase-3; GIST, gastrointestinal stromal tumor; MTC, medullary thyroid cancer; mTOR, mammalian target of rapamycin; PDGFR, platelet-derived growth factor receptor; PLGF, placental growth factor; PML-RAR α , promyelocytic leukemia-retinoic acid receptor-alpha; RCC, renal cell cancer; t(15;17), translocation between chromosomes 15 and 17; TC, thyroid cancer; TGF- α , transforming growth factor-alpha; VEGFR, vascular endothelial growth factor receptor.

and chemotherapy can be enhanced further by combinations with another targeted monoclonal antibody (pertuzumab), which prevents dimerization of the HER2 receptor with other HER family members including HER3.

Although targeted therapies have not yet resulted in cures when used alone, their use in the adjuvant setting and when combined with other effective treatments has substantially increased the fraction of patients cured. For example, the addition of rituximab, an anti-CD20 antibody, to combination chemotherapy in patients with diffuse large B cell lymphoma improves cure rates by 15–20%. The addition of trastuzumab, antibody to HER2, to combination chemotherapy in the adjuvant treatment of HER2-positive breast cancer reduces relapse rates by 50%.

A major effort is under way to develop targeted therapies for mutations in the *ras* family of genes, which are the most common mutations in oncogenes in cancers (especially *kras*) but have proved to be very difficult targets for a number of reasons related to how RAS proteins are activated and inactivated. Targeted therapies against proteins downstream of RAS (including mitogen-activated protein [MAP] kinase and ERK) are currently being studied, both individually and in combination. A large number of inhibitors of phospholipid signaling pathways such as the phosphatidylinositol-3-kinase (PI3K) and phospholipase C-gamma pathways, which are involved in a large number of cellular processes that are important in cancer development and progression, are being evaluated. The targeting of a variety of other pathways that are activated in malignant cells, such as the MET pathway, hedgehog pathway, and various angiogenesis pathways, is also being explored.

One of the strategies for new drug development is to take advantage of so-called oncogene addiction. This situation (Fig. 102e-3) is created

when a tumor cell develops an activating mutation in an oncogene that becomes a dominant pathway for survival and growth with reduced contributions from other pathways, even when there may be abnormalities in those pathways. This dependency on a single pathway creates a cell that is vulnerable to inhibitors of that oncogene pathway. For example, cells harboring mutations in *BRAF* are very sensitive to MEK inhibitors that inhibit downstream signaling in the BRAF pathway.

Targeting proteins critical for transcription of proteins vital for malignant cell survival or proliferation provides another potential target for treating cancers. The transcription factor nuclear factor- κ B (NF- κ B) is a heterodimer composed of p65 and p50 subunits that associate with an inhibitor, I κ B, in the cell cytoplasm. In response to growth factor or cytokine signaling, a multi-subunit kinase called IKK (I κ B kinase) phosphorylates I κ B and directs its degradation by the ubiquitin/proteasome system. NF- κ B, free of its inhibitor, translocates to the nucleus and activates target genes, many of which promote the survival of tumor cells. Novel drugs called *proteasome inhibitors* block the proteolysis of I κ B, thereby preventing NF- κ B activation. For unexplained reasons, this is selectively toxic to tumor cells. The antitumor effects of proteasome inhibitors are more complicated and involve the inhibition of the degradation of multiple cellular proteins. Proteasome inhibitors (e.g., bortezomib [Velcade]) have activity in patients with multiple myeloma, including partial and complete remissions. Inhibitors of IKK are also in development, with the hope of more selectively blocking the degradation of I κ B, thus “locking” NF- κ B in an inhibitory complex and rendering the cancer cell more susceptible to apoptosis-inducing agents. Many other transcription factors are activated by phosphorylation, which can be prevented by tyrosine kinase inhibitors or serine/threonine kinase inhibitors, a number of which are currently in clinical trials.

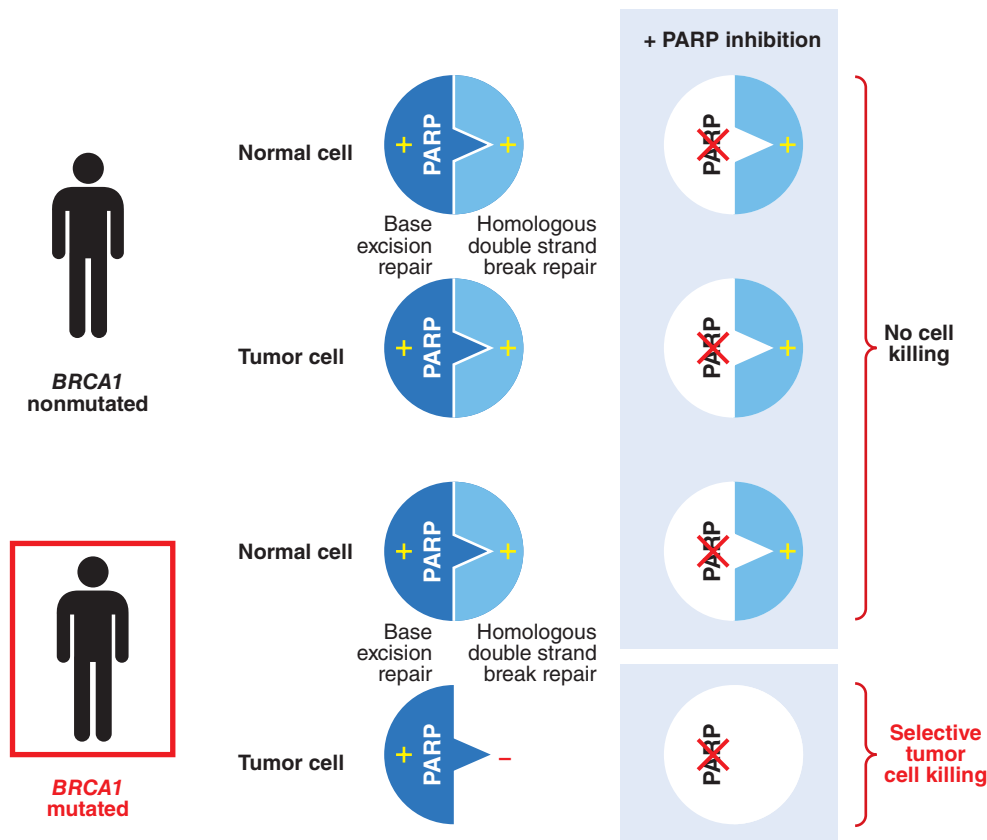


FIGURE 102e-3 Synthetic lethality. Genes are said to have a synthetic lethal relationship when mutation of either gene alone is tolerated by the cell but mutation of both genes leads to lethality, as originally noted by Bridges and later named by Dobzhansky. Thus, mutant *gene a* and *gene b* have a synthetic lethal relationship, implying that the loss of one gene makes the cell dependent on the function of the other gene. In cancer cells, loss of function of a DNA repair gene like *BRCA1*, which repairs double-strand breaks, makes the cell dependent on base excision repair mediated in part by *PARP*. If the *PARP* gene product is inhibited, the cell attempts to repair the break using the error-prone nonhomologous end-joining method, which results in tumor cell death. High-throughput screens can now be performed using isogenic cell line pairs in which one cell line has a defined defect in a DNA repair pathway. Compounds can be identified that selectively kill the mutant cell line; targets of these compounds have a synthetic lethal relationship to the repair pathway and are potentially important targets for future therapeutics.

Estrogen receptors (ERs) and androgen receptors (ARs), members of the steroid hormone family of nuclear receptors, are targets of inhibition by drugs used to treat breast and prostate cancers, respectively. Tamoxifen, a partial agonist and antagonist of ER function, can mediate tumor regression in metastatic breast cancer and can prevent disease recurrence in the adjuvant setting. Tamoxifen binds to the ER and modulates its transcriptional activity, inhibiting activity in the breast but promoting activity in bone and uterine epithelium. Selective ER modulators (SERMs) have been developed with the hope of a more beneficial modulation of ER activity, i.e., antiestrogenic activity in the breast, uterus, and ovary, but estrogenic for bone, brain, and cardiovascular tissues. Aromatase inhibitors, which block the conversion of androgens to estrogens in breast and subcutaneous fat tissues, have demonstrated improved clinical efficacy compared with tamoxifen and are often used as first-line therapy in patients with ER-positive disease. A number of approaches have been developed for blocking androgen stimulation of prostate cancer, including decreasing production (e.g., orchiectomy, luteinizing hormone-releasing hormone agonists or antagonists, estrogens, ketoconazole, and inhibitors of enzymes such as CYP17 involved in androgen production) and AR blockers (Chap. 108).

ONCOGENE ADDICTION AND SYNTHETIC LETHALITY

The concepts of oncogene addiction and synthetic lethality have spurred new drug development targeting oncogene- and tumor-suppressor pathways. As discussed earlier in this chapter and outlined in Fig. 102e-3, cancer cells can become dependent on signaling pathways containing activated oncogenes; this can effect proliferation (i.e., mutated *Kras*, *Braf*, overexpressed *Myc*, or activated tyrosine kinases), DNA repair (loss of *BRCA1* or *BRCA2* gene function), survival (overexpression of *Bcl-2* or *NF-κB*), cell metabolism (as occurs when mutant *Kras* enhances glucose uptake and aerobic glycolysis), and perhaps angiogenesis (production of VEGF in response to HIF-2α in RCC). In such cases, targeted inhibition of the pathway can lead to specific killing of the cancer cells. However, targeting defects in tumor-suppressor genes has been much more difficult, both because the target of mutation is often deleted and because it is much more difficult to restore normal function than to inhibit abnormal function of a protein. Synthetic lethality occurs when loss of function in either of two genes alone has limited effects on cell survival but loss of function in both genes leads to cell death. Identifying genes that have a synthetic lethal relationship to tumor-suppressor pathways that have been mutated in tumor cells may allow targeting of proteins required uniquely by those cells (Fig. 102e-3). Several examples of this have been identified. For instance, cells with mutations in the *BRCA1* or *BRCA2* tumor-suppressor genes (e.g., a subset of breast and ovarian cancers) are unable to repair DNA damage by homologous recombination. PARP are a family of proteins important for single-strand break (SSB) DNA repair. PARP inhibition results in selective killing of cancer cells with *BRCA1* or *BRCA2* loss. Preliminary trials have suggested some effectiveness of PARP inhibition, especially in combination with chemotherapy; clinical trials are ongoing. The concept of synthetic lethality provides a framework for genetic screens to identify other synthetic lethal combinations involving known tumor-suppressor genes and development of novel therapeutic agents to target dependent pathways.

EPIGENETIC INFLUENCES ON CANCER GENE TRANSCRIPTION

Chromatin structure regulates the hierarchical order of sequential gene transcription that governs differentiation and tissue homeostasis. Disruption of chromatin remodeling (the process of modifying chromatin structure to control exposure of specific genes to transcriptional proteins, thereby controlling the expression of those genes) leads to aberrant gene expression and can induce proliferation of undifferentiated cells. *Epigenetics* is defined as changes that alter the pattern of gene expression that persist across at least one cell division but are not caused by changes in the DNA code. Epigenetic changes include alterations of chromatin structure mediated by methylation of cytosine residues in CpG dinucleotides, modification of histones by acetylation or methylation, or changes in higher-order chromosome structure

(Fig. 102e-4). The transcriptional regulatory regions of active genes often contain a high frequency of CpG dinucleotides (referred to as *CpG islands*), which are normally unmethylated. Expression of these genes is controlled by transient association with repressor or activator proteins that regulate transcriptional activation. However, hypermethylation of promoter regions is a common mechanism by which tumor-suppressor loci are epigenetically silenced in cancer cells. Thus one allele may be inactivated by mutation or deletion (as occurs in loss of heterozygosity), while expression of the other allele is epigenetically silenced, usually by methylation.

Acetylation of the amino terminus of the core histones H3 and H4 induces an open chromatin conformation that promotes transcription initiation. Histone acetylases are components of coactivator complexes recruited to promoter/enhancer regions by sequence-specific transcription factors during the activation of genes (Fig. 102e-4). Histone deacetylases (HDACs; at least 17 are encoded in the human genome) are recruited to genes by transcriptional repressors and prevent the initiation of gene transcription. Methylated cytosine residues in promoter regions become associated with methyl cytosine-binding proteins that recruit protein complexes with HDAC activity. The balance between permissive and inhibitory chromatin structure is therefore largely determined by the activity of transcription factors in modulating the “histone code” and the methylation status of the genetic regulatory elements of genes.

The pattern of gene transcription is aberrant in all human cancers, and in many cases, epigenetic events are responsible. Unlike genetic events that alter DNA primary structure (e.g., deletions), epigenetic changes are potentially reversible and appear amenable to therapeutic intervention. In certain human cancers, including pancreatic cancer and multiple myeloma, the p16^{ink4a} promoter is inactivated by methylation, thus permitting the unchecked activity of CDK4/cyclin D and rendering pRb nonfunctional. In sporadic forms of renal, breast, and colon cancer, the von Hippel-Lindau (*VHL*), breast cancer 1 (*BRCA1*), and serine/threonine kinase 11 (*STK11*) genes, respectively, are epigenetically silenced. Other targeted genes include the p15^{ink4b} CDK inhibitor, glutathione-S-transferase (which detoxifies reactive oxygen species), and the E-cadherin molecule (important for junction formation between epithelial cells). Epigenetic silencing can occur in pre-malignant lesions and can affect genes involved in DNA repair, thus predisposing to further genetic damage. Examples include MLH1 (mut L homologue) in hereditary nonpolyposis colon cancer (HNPCC, also called Lynch’s syndrome), which is critical for repair of mismatched bases that occur during DNA synthesis, and O⁶-methylguanine-DNA methyltransferase, which removes alkylated guanine adducts from DNA and is often silenced in colon, lung, and lymphoid tumors.

Human leukemias often have chromosomal translocations that code for novel fusion proteins with enzymatic activities that alter chromatin structure. The promyelocytic leukemia-retinoic acid receptor (PML-RAR) fusion protein, generated by the t(15;17) observed in most cases of acute promyelocytic leukemia (APL), binds to promoters containing retinoic acid response elements and recruits HDAC to these promoters, effectively inhibiting gene expression. This arrests differentiation at the promyelocyte stage and promotes tumor cell proliferation and survival. Treatment with pharmacologic doses of all-*trans* retinoic acid (ATRA), the ligand for RARα, results in the release of HDAC activity and the recruitment of coactivators, which overcome the differentiation block. This induced differentiation of APL cells has improved treatment of these patients but also has led to a novel treatment toxicity when newly differentiated tumor cells infiltrate the lungs. However, ATRA represents a treatment paradigm for the reversal of epigenetic changes in cancer. For other leukemia-associated fusion proteins, such as acute myeloid leukemia (AML)-eight-twenty-one (ETO) and the MLL fusion proteins seen in AML and acute lymphocytic leukemia, no ligand is known. Therefore, efforts are ongoing to determine the structural basis for interactions between translocation fusion proteins and chromatin-remodeling proteins and to use this information to rationally design small molecules that will disrupt specific protein-protein associations, although this has proven to be technically difficult. Drugs that block the enzymatic activity of HDAC are

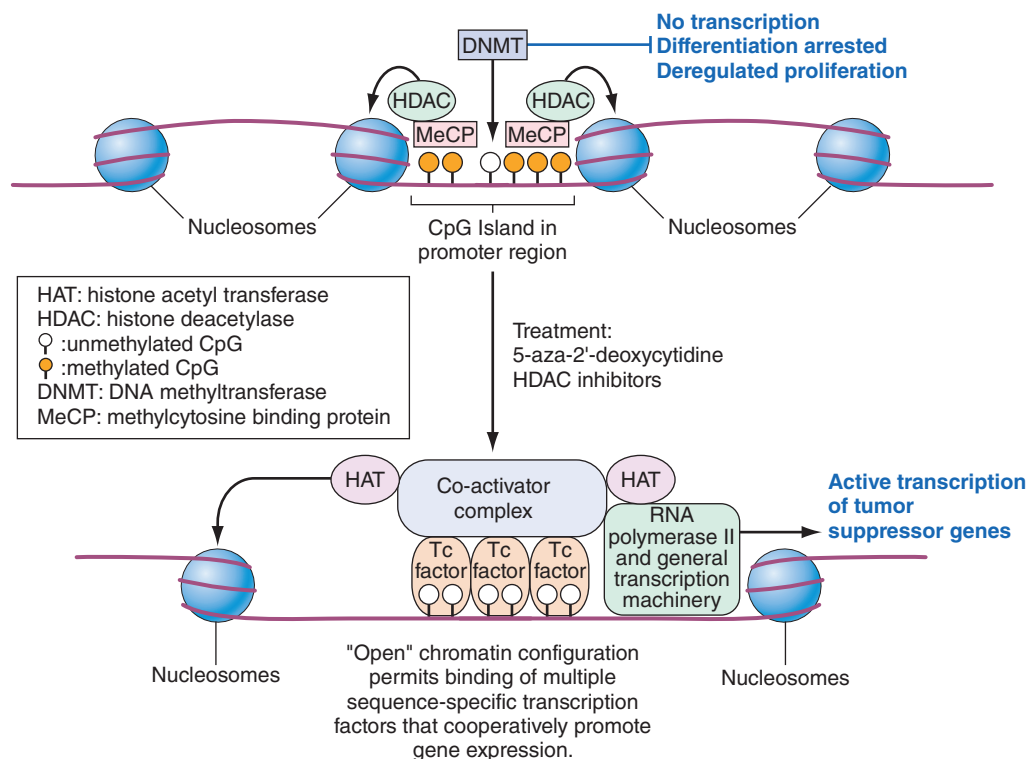


FIGURE 102e-4 Epigenetic regulation of gene expression in cancer cells. Tumor-suppressor genes are often epigenetically silenced in cancer cells. In the upper portion, a CpG island within the promoter and enhancer regions of the gene has been methylated, resulting in the recruitment of methyl-cytosine binding proteins (MeCP) and complexes with histone deacetylase (HDAC) activity. Chromatin is in a condensed, non-permissive conformation that inhibits transcription. Clinical trials are under way using the combination of demethylating agents such as 5-aza-2'-deoxycytidine plus HDAC inhibitors, which together confer an open, permissive chromatin structure (*lower portion*). Transcription factors bind to specific DNA sequences in promoter regions and, through protein-protein interactions, recruit coactivator complexes containing histone acetyl transferase (HAT) activity. This enhances transcription initiation by RNA polymerase II and associated general transcription factors. The expression of the tumor-suppressor gene commences, with phenotypic changes that may include growth arrest, differentiation, or apoptosis.

being tested. HDAC inhibitors have demonstrated antitumor activity in clinical studies against cutaneous T cell lymphoma (e.g., vorinostat) and some solid tumors. HDAC inhibitors may target cancer cells via a number of mechanisms, including upregulation of death receptors (DR4/5, FAS, and their ligands) and p21^{Cip1/Waf1}, as well as inhibition of cell cycle checkpoints.

Efforts are also under way to reverse the hypermethylation of CpG islands that characterizes many malignancies. Drugs that induce DNA demethylation, such as 5-aza-2'-deoxycytidine, can lead to reexpression of silenced genes in cancer cells with restoration of function, and 5-aza-2'-deoxycytidine is approved for use in myelodysplastic syndrome (MDS). However, 5-aza-2'-deoxycytidine has limited aqueous solubility and is myelosuppressive. Other inhibitors of DNA methyltransferases are in development. In ongoing clinical trials, inhibitors of DNA methylation are being combined with HDAC inhibitors. The hope is that by reversing coexisting epigenetic changes, the deregulated patterns of gene transcription in cancer cells will be at least partially reversed.

Epigenetic gene regulation can also occur via microRNAs or long non-coding RNAs (lncRNAs). MicroRNAs are short (average 22 nucleotides in length) RNA molecules that silence gene expression after transcription by binding and inhibiting the translation or promoting the degradation of mRNA transcripts. It is estimated that more than 1000 microRNAs are encoded in the human genome. Each tissue has a distinctive repertoire of microRNA expression, and this pattern is altered in specific ways in cancers. However, specific correlations between microRNA expression and tumor biology and clinical behavior are just now emerging. Therapies targeting microRNAs are not currently at hand but represent a novel area of treatment development. lncRNAs are longer than 200 nucleotides and compose the largest group of noncoding RNAs. Some of them have been shown

to play important roles in gene regulation. The potential for altering these RNAs for therapeutic benefit is an area of active investigation, although much more needs to be learned before this will be feasible.

APOPTOSIS AND OTHER MECHANISMS OF CELL DEATH

Tissue homeostasis requires a balance between the death of aged, terminally differentiated cells or severely damaged cells and their renewal by proliferation of committed progenitors. Genetic damage to growth-regulating genes of stem cells could lead to catastrophic results for the host as a whole. Thus, genetic events causing activation of oncogenes or loss of tumor suppressors, which would be predicted to lead to unregulated cell proliferation unless corrected, usually activate signal transduction pathways that block aberrant cell proliferation. These pathways can lead to a form of programmed cell death (*apoptosis*) or irreversible growth arrest (*senescence*). Much as a panoply of intra- and extracellular signals impinge upon the core cell cycle machinery to regulate cell division, so too are these signals transmitted to a core enzymatic machinery that regulates cell death and survival.

Apoptosis is induced by two main pathways (Fig. 102e-5). The extrinsic pathway of apoptosis is activated by cross-linking members of the tumor necrosis factor (TNF) receptor superfamily, such as CD95 (Fas) and death receptors DR4 and DR5, by their ligands, Fas ligand or TRAIL (TNF-related apoptosis-inducing ligand), respectively. This induces the association of FADD (Fas-associated death domain) and procaspase-8 to death domain motifs of the receptors. Caspase-8 is activated and then cleaves and activates effector caspases-3 and -7, which then target cellular constituents (including caspase-activated DNase, cytoskeletal proteins, and a number of regulatory proteins), inducing the morphologic appearance characteristic of apoptosis, which pathologists term "karyorrhexis." The intrinsic pathway of apoptosis is initiated by the release of cytochrome c and SMAC (second

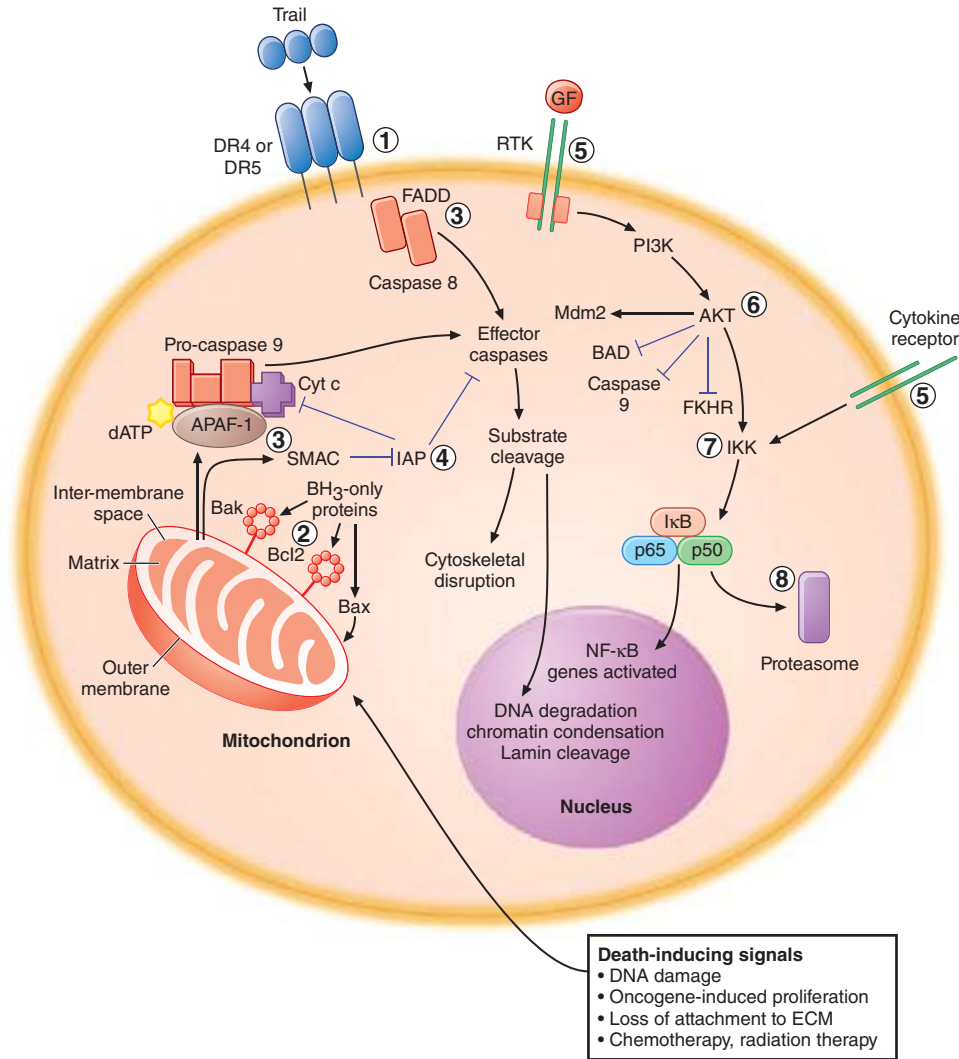


FIGURE 102e-5 Therapeutic strategies to overcome aberrant survival pathways in cancer cells. **1.** The extrinsic pathway of apoptosis can be selectively induced in cancer cells by TRAIL (the ligand for death receptors 4 and 5) or by agonistic monoclonal antibodies. **2.** Inhibition of antiapoptotic Bcl-2 family members with antisense oligonucleotides or inhibitors of the BH₃-binding pocket will promote formation of Bak- or Bax-induced pores in the mitochondrial outer membrane. **3.** Epigenetic silencing of APAF-1, caspase-8, and other proteins can be overcome using demethylating agents and inhibitors of histone deacetylases. **4.** Inhibitor of apoptosis proteins (IAP) blocks activation of caspases; small-molecule inhibitors of IAP function (mimicking SMAC action) should lower the threshold for apoptosis. **5.** Signal transduction pathways originating with activation of receptor tyrosine kinase receptors (RTKs) or cytokine receptors promote survival of cancer cells by a number of mechanisms. Inhibiting receptor function with monoclonal antibodies, such as trastuzumab or cetuximab, or inhibiting kinase activity with small-molecule inhibitors can block the pathway. **6.** The Akt kinase phosphorylates many regulators of apoptosis to promote cell survival; inhibitors of Akt may render tumor cells more sensitive to apoptosis-inducing signals; however, the possibility of toxicity to normal cells may limit the therapeutic value of these agents. **7** and **8.** Activation of the transcription factor NF- κ B (composed of p65 and p50 subunits) occurs when its inhibitor, I κ B, is phosphorylated by I κ B kinase (IKK), with subsequent degradation of I κ B by the proteasome. Inhibition of IKK activity should selectively block the activation of NF- κ B target genes, many of which promote cell survival. Inhibitors of proteasome function are Food and Drug Administration approved and may work in part by preventing destruction of I κ B, thus blocking NF- κ B nuclear localization. NF- κ B is unlikely to be the only target for proteasome inhibitors.

mitochondrial activator of caspases) from the mitochondrial inter-membrane space in response to a variety of noxious stimuli, including DNA damage, loss of adherence to the extracellular matrix (ECM), oncogene-induced proliferation, and growth factor deprivation. Upon release into the cytoplasm, cytochrome *c* associates with dATP, procaspase-9, and the adaptor protein APAF-1, leading to the sequential activation of caspase-9 and effector caspases. SMAC binds to and blocks the function of inhibitor of apoptosis proteins (IAP), negative regulators of caspase activation.

The release of apoptosis-inducing proteins from the mitochondria is regulated by pro- and antiapoptotic members of the Bcl-2 family. Antiapoptotic members (e.g., Bcl-2, Bcl-XL, and Mcl-1) associate with the mitochondrial outer membrane via their carboxyl termini, exposing to the cytoplasm a hydrophobic binding pocket composed

of Bcl-2 homology (BH) domains 1, 2, and 3 that is crucial for their activity. Perturbations of normal physiologic processes in specific cellular compartments lead to the activation of BH3-only proapoptotic family members (such as Bad, Bim, Bid, Puma, Noxa, and others) that can alter the conformation of the outer-membrane proteins Bax and Bak, which then oligomerize to form pores in the mitochondrial outer membrane resulting in cytochrome *c* release. If proteins composed only of BH3 domains are sequestered by Bcl-2, Bcl-XL, or Mcl-1, pores do not form and apoptosis-inducing proteins are not released from the mitochondria. The ratio of levels of antiapoptotic Bcl-2 family members and the levels of proapoptotic BH3-only proteins at the mitochondrial membrane determines the activation state of the intrinsic pathway. The mitochondrion must therefore be recognized not only as an organelle with vital roles in intermediary metabolism and

102e-10 oxidative phosphorylation but also as a central regulatory structure of the apoptotic process.

The evolution of tumor cells to a more malignant phenotype requires the acquisition of genetic changes that subvert apoptosis pathways and promote cancer cell survival and resistance to anticancer therapies. However, cancer cells may be more vulnerable than normal cells to therapeutic interventions that target the apoptosis pathways that cancer cells depend on. For instance, overexpression of Bcl-2 as a result of the t(14;18) translocation contributes to follicular lymphoma. Upregulation of Bcl-2 expression is also observed in prostate, breast, and lung cancers and melanoma. Targeting of antiapoptotic Bcl-2 family members has been accomplished by the identification of several low-molecular-weight compounds that bind to the hydrophobic pockets of either Bcl-2 or Bcl-XL and block their ability to associate with death-inducing BH3-only proteins. These compounds inhibit the antiapoptotic activities of Bcl-2 and Bcl-XL at nanomolar concentrations in the laboratory and are entering clinical trials.

Preclinical studies targeting death receptors DR4 and DR5 have demonstrated that recombinant, soluble, human TRAIL or humanized monoclonal antibodies with agonist activity against DR4 or DR5 can induce apoptosis of tumor cells while sparing normal cells. The mechanisms for this selectivity may include expression of decoy receptors or elevated levels of intracellular inhibitors (such as FLIP, which competes with caspase-8 for FADD) by normal cells but not tumor cells. Synergy has been shown between TRAIL-induced apoptosis and chemotherapeutic agents. For instance, some colon cancers encode mutated Bax protein as a result of mismatch repair (MMR) defects and are resistant to TRAIL. However, upregulation of Bak by chemotherapy restores the ability of TRAIL to activate the mitochondrial pathway of apoptosis. However, clinical studies have not yet shown significant activity of approaches targeting the TRAIL pathway.

Many of the signal transduction pathways perturbed in cancer promote tumor cell survival (Fig. 102e-5). These include activation of the PI3K/Akt pathway, increased levels of the NF- κ B transcription factor, and epigenetic silencing of genes such as *APAF-1* and *caspase-8*. Each of these pathways is a target for therapeutic agents that, in addition to affecting cancer cell proliferation or gene expression, may render cancer cells more susceptible to apoptosis, thus promoting synergy when combined with other chemotherapeutic agents.

Some tumor cells resist drug-induced apoptosis by expression of one or more members of the ABC family of ATP-dependent efflux pumps that mediate the multidrug-resistance (MDR) phenotype. The prototype, P-glycoprotein (PGP), spans the plasma membrane 12 times and has two ATP-binding sites. Hydrophobic drugs (e.g., anthracyclines and vinca alkaloids) are recognized by PGP as they enter the cell and are pumped out. Numerous clinical studies have failed to demonstrate that drug resistance can be overcome using inhibitors of PGP. However, ABC transporters have different substrate specificities, and inhibition of a single family member may not be sufficient to overcome the MDR phenotype. Efforts to reverse PGP-mediated drug resistance continue.

Cells, including cancer cells, can also undergo other mechanisms of cell death including *autophagy* (degradation of proteins and organelles by lysosomal proteases) and *necrosis* (digestion of cellular components and rupturing of the cell membrane). Necrosis usually occurs in response to external forces resulting in release of cellular components, which leads to inflammation and damage to surrounding tissues. Although necrosis was thought to be unprogrammed, evidence now suggests that at least some aspects may be programmed. The exact role of necrosis in cancer cell death in various settings is still being determined. In addition to its role in cell death, autophagy can serve as a homeostatic mechanism to promote survival for the cell by recycling cellular components to provide necessary energy. The mechanisms that control the balance between enhancing survival versus leading to cell death are still not fully understood. Autophagy appears to play conflicting roles in the development and survival of cancer. Early in the carcinogenic process, it can act as a tumor suppressor by preventing the cell from accumulating abnormal proteins and organelles. However, in established tumors, it may serve as a mechanism of survival for cancer

cells when they are stressed by damage such as from chemotherapy. Inhibition of this process can enhance the sensitivity of cancer cells to chemotherapy. Better understanding of the factors that control the survival-promoting versus death-inducing aspects of autophagy is required in order to know how to best manipulate it for therapeutic benefit.

METASTASIS

The metastatic process accounts for the vast majority of deaths from solid tumors, and therefore, an understanding of this process is critical. The biology of metastasis is complex and requires multiple steps. The three major features of tissue invasion are cell adhesion to the basement membrane, local proteolysis of the membrane, and movement of the cell through the rent in the membrane and the ECM. Cells that lose contact with the ECM normally undergo programmed cell death (anoikis), and this process has to be suppressed in cells that metastasize. Another process important for metastasizing epithelial cancer cells is epithelial-mesenchymal transition (EMT). This is a process by which cells lose their epithelial properties and gain mesenchymal properties. This normally occurs during the developmental process in embryos, allowing cells to migrate to their appropriate destinations in the embryo. It also occurs in wound healing, tissue regeneration, and fibrotic reactions, but in all of these processes, cells stop proliferating when the process is complete. Malignant cells that metastasize undergo EMT as an important step in that process but retain the capacity for unregulated proliferation. Malignant cells that gain access to the circulation must then repeat those steps at a remote site, find a hospitable niche in a foreign tissue, avoid detection by host defenses, and induce the growth of new blood vessels. The rate-limiting step for metastasis is the ability for tumor cells to survive and expand in the novel microenvironment of the metastatic site, and multiple host-tumor interactions determine the ultimate outcome (Fig. 102e-6). Few drugs have been developed to attempt to directly target the process of metastasis, in part because the specifics of the critical steps in the process that would be potentially good targets for drugs are still being identified. However, a number of potential targets are known. HER2 can enhance the metastatic potential of breast cancer cells, and as discussed above, the monoclonal antibody trastuzumab, which targets HER2, improves survival in the adjuvant setting for HER2-positive breast cancer patients. Other potential targets that increase metastatic potential of cells in preclinical studies include HIF-1 and -2, transcription factors induced by hypoxia within tumors; growth factors (e.g., cMET and VEGFR); oncogenes (e.g., *SRC*); adhesion molecules (e.g., focal adhesion kinase [FAK]); ECM proteins (e.g., matrix metalloproteinases-1 and -2); and inflammatory molecules (e.g., COX-2).

The metastatic phenotype is likely restricted to a small fraction of tumor cells (Fig. 102e-6). A number of genetic and epigenetic changes are required for tumor cells to be able to metastasize, including activation of metastasis-promoting genes and inhibition of genes that suppress the metastatic ability. Cells with metastatic capability frequently express chemokine receptors that are likely important in the metastatic process. A number of candidate metastasis-suppressor genes have been identified, including genes coding for proteins that enhance apoptosis, suppress cell division, are involved in the interactions of cells with each other or the ECM, or suppress cell migration. The loss of function of these genes enhances metastasis. Gene expression profiling is being used to study the metastatic process and other properties of tumor cells that may predict susceptibilities.

An example of the ability of malignant cells to survive and grow in a novel microenvironment is bone metastasis. Bone metastases are extremely painful, cause fractures of weight-bearing bones, can lead to hypercalcemia, and are a major cause of morbidity for cancer patients. Osteoclasts and their monocyte-derived precursors express the surface receptor RANK (receptor activator of NF- κ B), which is required for terminal differentiation and activation of osteoclasts. Osteoblasts and other stromal cells express RANK ligand (RANKL), as both a membrane-bound and soluble cytokine. Osteoprotegerin (OPG), a soluble receptor for RANKL produced by stromal cells, acts as a decoy receptor to inhibit RANK activation. The relative balance of RANKL

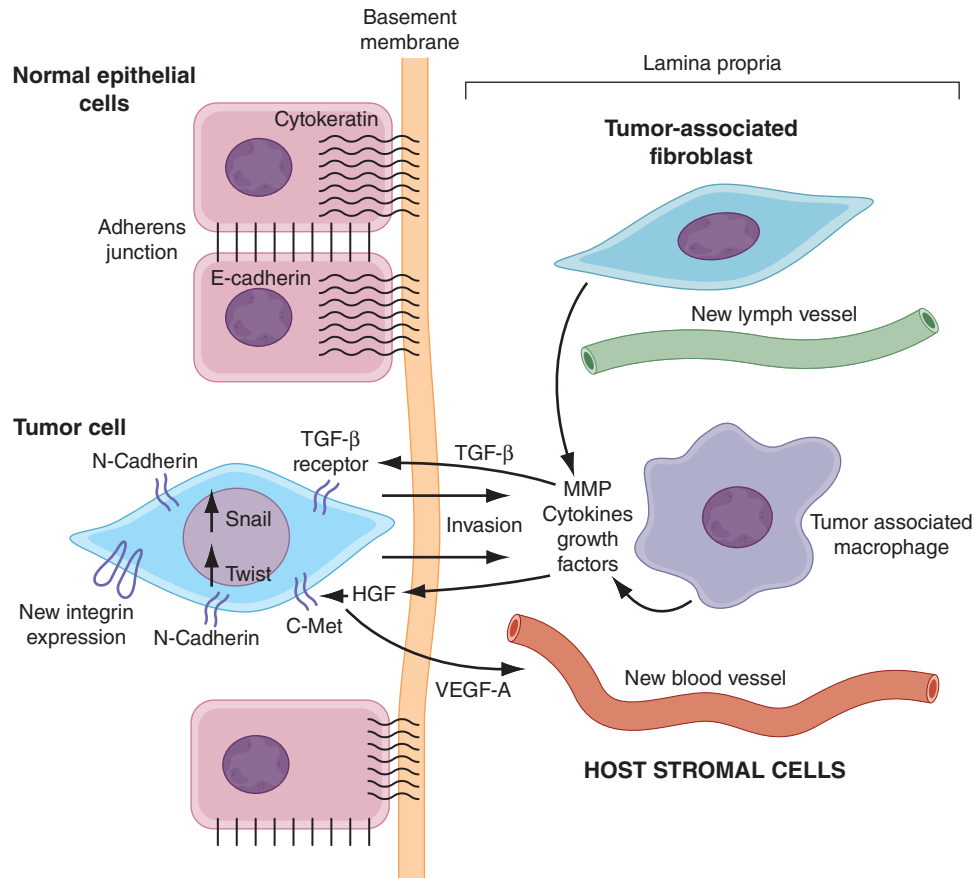


FIGURE 102e-6 Oncogene signaling pathways are activated during tumor progression and promote metastatic potential. This figure shows a cancer cell that has undergone epithelial to mesenchymal transition (EMT) under the influence of several environmental signals. Critical components include activated transforming growth factor β (TGF- β) and the hepatocyte growth factor (HGF)/c-Met pathways, as well as changes in the expression of adhesion molecules that mediate cell-cell and cell-extracellular matrix interactions. Important changes in gene expression are mediated by the Snail and Twist family of transcriptional repressors (whose expression is induced by the oncogenic pathways), leading to reduced expression of E-cadherin, a key component of adherens junctions between epithelial cells. This, in conjunction with upregulation of N-cadherin, a change in the pattern of expression of integrins (which mediate cell-extracellular matrix associations that are important for cell motility), and a switch in intermediate filament expression from cytokeratin to vimentin, results in the phenotypic change from adherent highly organized epithelial cells to motile and invasive cells with a fibroblast or mesenchymal morphology. EMT is thought to be an important step leading to metastasis in some human cancers. Host stromal cells, including tumor-associated fibroblasts and macrophages, play an important role in modulating tumor cell behavior through secretion of growth factors and proangiogenic cytokines, and matrix metalloproteinases that degrade the basement membrane. VEGF-A, -C, and -D are produced by tumor cells and stromal cells in response to hypoxemia or oncogenic signals and induce production of new blood vessels and lymphatic channels through which tumor cells metastasize to lymph nodes or tissues.

and OPG determines the activation state of RANK on osteoclasts. Many tumors increase osteoclast activity by secretion of substances such as parathyroid hormone (PTH), PTH-related peptide, interleukin (IL)-1, or Mip1 that perturb the homeostatic balance of bone remodeling by increasing RANK signaling. One example is multiple myeloma, where tumor cell-stromal cell interactions activate osteoclasts and inhibit osteoblasts, leading to the development of multiple lytic bone lesions. Inhibition of RANKL by an antibody (denosumab) can prevent further bone destruction. Bisphosphonates are also effective inhibitors of osteoclast function that are used in the treatment of cancer patients with bone metastases.

CANCER STEM CELLS

Only a small proportion of the cells within a tumor are capable of initiating colonies *in vitro* or forming tumors at high efficiency when injected into immunocompromised NOD/SCID mice. Acute and chronic myeloid leukemias (AML and CML) have a small population of cells (<1%) that have properties of stem cells, such as unlimited self-renewal and the capacity to cause leukemia when serially transplanted in mice. These cells have an undifferentiated phenotype (Thy1-CD34+CD38- and do not express other differentiation markers) and resemble normal stem cells in many ways, but are no longer under

homeostatic control (Fig. 102e-7). Solid tumors may also contain a population of stem cells. Cancer stem cells, like their normal counterparts, have unlimited proliferative capacity and paradoxically traverse the cell cycle at a very slow rate; cancer growth occurs largely due to expansion of the stem cell pool, the unregulated proliferation of an amplifying population, and failure of apoptosis pathways (Fig. 102e-7). Slow cell cycle progression and high levels of expression of antiapoptotic Bcl-2 family members and drug efflux pumps of the MDR family render cancer stem cells less vulnerable to cancer chemotherapy or radiation therapy. Implicit in the cancer stem cell hypothesis is the idea that failure to cure most human cancers is due to the fact that current therapeutic agents do not kill the stem cells. If cancer stem cells can be identified and isolated, then aberrant signaling pathways that distinguish these cells from normal tissue stem cells can be identified and targeted. Evidence that cells with stem cell properties can arise from other epithelial cells within the cancer by processes such as epithelial mesenchymal transition also implies that it is essential to treat all of the cancer cells, and not just those with current stem cell-like properties, in order to eliminate the self-renewing cancer cell population. The exact nature of cancer stem cells remains an area of investigation. One of the unanswered questions is the exact origin of cancer stem cells for the different cancers.

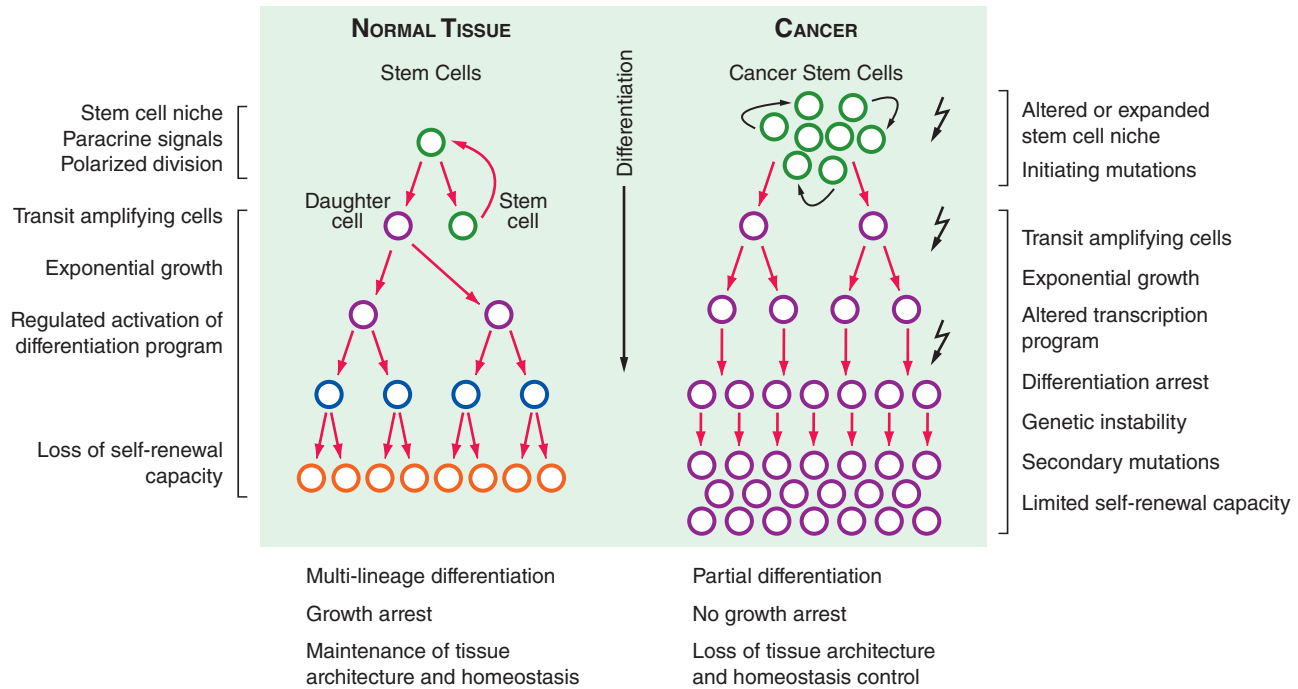


FIGURE 102e-7 Cancer stem cells play a critical role in the initiation, progression, and resistance to therapy of malignant neoplasms.

In normal tissues (*left*), homeostasis is maintained by asymmetric division of stem cells, leading to one progeny cell that will differentiate and one cell that will maintain the stem cell pool. This occurs within highly specific niches unique to each tissue, such as in close apposition to osteoblasts in bone marrow, or at the base of crypts in the colon. Here, paracrine signals from stromal cells, such as sonic hedgehog or Notch ligands, as well as upregulation of β -catenin and telomerase, help to maintain stem cell features of unlimited self-renewal while preventing differentiation or cell death. This occurs in part through upregulation of the transcriptional repressor Bmi-1 and inhibition of the p16^{Ink4a}/Arf and p53 pathways. Daughter cells leave the stem cells niche and enter a proliferative phase (referred to as *transit-amplifying*) for a specified number of cell divisions, during which time a developmental program is activated, eventually giving rise to fully differentiated cells that have lost proliferative potential. Cell renewal equals cell death, and homeostasis is maintained. In this hierarchical system, only stem cells are long-lived. The hypothesis is that cancers harbor stem cells that make up a small fraction (i.e., 0.001–1%) of all cancer cells. These cells share several features with normal stem cells, including an undifferentiated phenotype, unlimited self-renewal potential, and a capacity for some degree of differentiation; however, due to initiating mutations (mutations are indicated by lightning bolts), they are no longer regulated by environmental cues. The cancer stem cell pool is expanded, and rapidly proliferating progeny, through additional mutations, may attain stem cell properties, although most of this population is thought to have a limited proliferative capacity. Differentiation programs are dysfunctional due to reprogramming of the pattern of gene transcription by oncogenic signaling pathways. Within the cancer transit-amplifying population, genomic instability generates aneuploidy and clonal heterogeneity as cells attain a fully malignant phenotype with metastatic potential. The cancer stem cell hypothesis has led to the idea that current cancer therapies may be effective at killing the bulk of tumor cells but do not kill tumor stem cells, leading to a regrowth of tumors that is manifested as tumor recurrence or disease progression. Research is in progress to identify unique molecular features of cancer stem cells that can lead to their direct targeting by novel therapeutic agents.

PLASTICITY AND RESISTANCE

Cancer cells, and especially stem cells, have the capacity for significant plasticity, allowing them to alter multiple aspects of cell biology in response to external factors (e.g., chemotherapy, inflammation, immune response). Thus, a major problem in cancer therapy is that malignancies have a wide spectrum of mechanisms for both initial and adaptive resistance to treatments. These include inhibiting drug delivery to the cancer cells, blocking drug uptake and retention, increasing drug metabolism, altering levels of target proteins, acquiring mutations in target proteins, modifying metabolism and cell signaling pathways, using alternate signaling pathways, adjusting the cell replication process including mechanisms by which the cell deals with DNA damage, inhibiting apoptosis, and evading the immune system. Thus, most metastatic cancers (except those curable with chemotherapy such as germ cell tumors) eventually become resistant to the therapy being used. Overcoming resistance is a major area of research.

CANCER METABOLISM

One of the distinguishing characteristics of cancer cells is that they have altered metabolism as compared with normal cells in supporting survival and their high rates of proliferation. These cells must focus a significant fraction of their energy resources on synthesis of proteins and other molecules while still maintaining sufficient ATP production

to survive and grow. Although normal proliferating cells also have similar needs, there are differences in how cancer cells metabolize glucose and a number of other compounds, including glutamine, as compared to normal cells. Many cancer cells use aerobic glycolysis (the Warburg effect) (Fig. 102e-8) to metabolize glucose, leading to increased lactic acid production, whereas normal cells use oxidative phosphorylation in mitochondria under aerobic conditions, a much more efficient process. One consequence is increased glucose uptake by cancer cells, a fact used in fluorodeoxyglucose (FDG) positron emission tomography (PET) scanning to detect tumors. A number of proteins in cancer cells, including CMYC, HIF1, RAS, p53, pRB, and AKT, are all involved in modulating glycolytic processes and controlling the Warburg effect. Although these pathways remain difficult to target therapeutically, both the PI3 kinase pathway with signaling through mTOR and the AMP-activated kinase (AMPK) pathway, which inhibits mTOR complex 1 (mTORC1; a protein complex that includes mTOR), are important in controlling the glycolytic process and thus provide potential targets for inhibiting this process. The inefficient utilization of glucose also leads to a need for alternative metabolic pathways for other compounds as well, one of which is glutamine. Similar to glucose, this provides both a source for structural molecules as well as energy production. Glutamine is also inefficiently used by cancer cells.

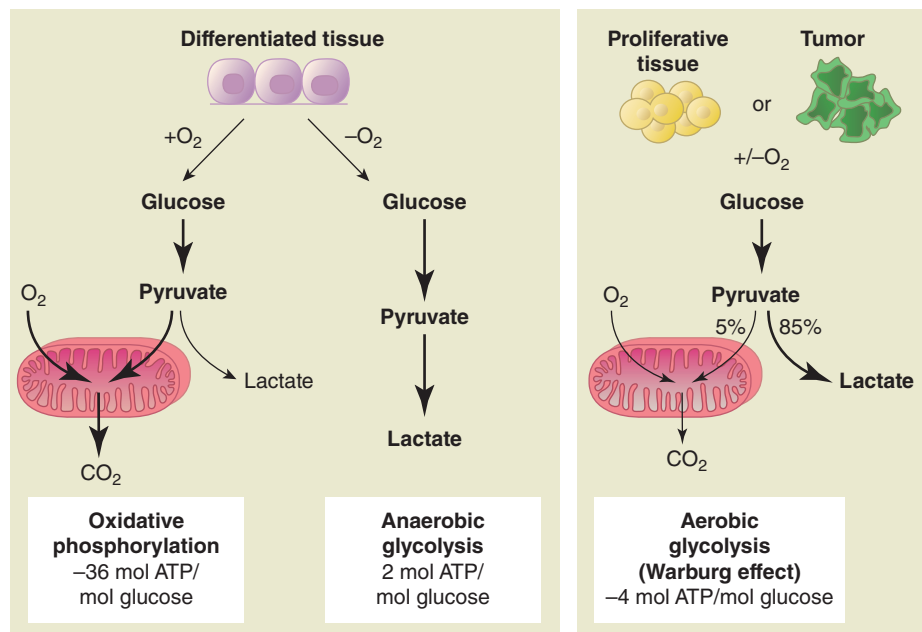


FIGURE 102e-8 Warburg effect versus oxidative phosphorylation. In most normal tissues, the vast majority of cells are differentiated and dedicated to a particular function within the organ in which they reside. The metabolic needs are mainly for energy and not for building blocks for new cells. In these tissues, ATP is generated by oxidative phosphorylation that efficiently generates about 36 molecules of ATP for each molecule of glucose metabolized. By contrast, proliferative tumor tissues, especially in the setting of hypoxia, a typical condition within tumors, use aerobic glycolysis to generate energy for cell survival and generation of building blocks for new cells.

Mutations in genes involved in the metastatic process occur in a number of cancers. Among the most frequently found to date are mutations in isocitrate dehydrogenases 1 and 2 (*IDH1* and *IDH2*). These have been most commonly seen in gliomas, AML, and intrahepatic cholangiocarcinomas. These mutations lead to the production of an oncometabolite (2-hydroxyglutarate [2HG]) instead of the normal product α -ketoglutarate. Although the exact mechanisms of oncogenesis by 2HG are still being elucidated, α -ketoglutarate is a key cofactor for a number of dioxygenases involved in controlling DNA methylation. 2HG can act as a competitive inhibitor for α -ketoglutarate, leading to alterations in methylation status (primarily hypermethylation) of genes (epigenetic changes) that can have profound effects on a number of cellular processes including differentiation. Inhibitors of mutant *IDH1* and *IDH2* are being developed.

Much needs to be learned about the specific differences in metabolism between cancer cells and normal cells; however, modulators of metabolism are being tested clinically. The first of these is the antidiabetic agent metformin, both alone and in combination with chemotherapeutic agents. Metformin inhibits gluconeogenesis and may have direct effects on tumor cells by activating the 5'-adenosine monophosphate-activated kinase (AMPK), a serine/threonine protein kinase that is downstream of the LKB1 tumor suppressor, and thus inhibiting mTORC1. This leads to decreased protein synthesis and proliferation. A second approach being tested involves dichloroacetate (DCA), an inhibitor of pyruvate dehydrogenase kinase (PDK). PDK inhibits pyruvate dehydrogenase in cancer cells, leading to a switch from mitochondrial oxidative phosphorylation of glucose to cytoplasmic glycolysis (the Warburg effect). By blocking PDK, DCA inhibits glycolysis. Additional approaches targeting tumor metabolism will likely emerge.

TUMOR MICROENVIRONMENT, ANGIOGENESIS, AND IMMUNE EVASION

Tumors consist not only of malignant cells but also of a complex microenvironment including many other types of cells (e.g., inflammatory cells), ECM, secreted factors (e.g., growth factors), reactive oxygen and nitrogen species, mechanical factors, blood vessels, and lymphatics. This microenvironment is not static but rather is dynamic and

continually evolving. Both the complexity and dynamic nature of the microenvironment enhance the difficulty of treating tumors. There are also a number of mechanisms by which the microenvironment can contribute to resistance to anticancer therapies.

One of the critical elements of tumor cell proliferation is delivery of oxygen, nutrients, and circulating factors important for growth and survival. The diffusion limit for oxygen in tissues is ~ 100 – 200 μm , and thus, a critical aspect in the growth of tumors is the development of new blood vessels, or angiogenesis. The growth of primary and metastatic tumors to larger than a few millimeters requires the recruitment of blood vessels and vascular endothelial cells to support their metabolic requirements. Thus, a critical element in growth of primary tumors and formation of metastatic sites is the *angiogenic switch*: the ability of the tumor to promote the formation of new capillaries from pre-existing host vessels. The angiogenic switch is a phase in tumor development when the dynamic balance of pro- and antiangiogenic factors is tipped in favor of vessel formation by the effects of the tumor on its immediate environment. Stimuli for tumor angiogenesis include hypoxemia, inflammation, and genetic lesions in oncogenes or tumor suppressors that alter tumor cell gene expression. Angiogenesis consists of several steps,

including the stimulation of endothelial cells (ECs) by growth factors, degradation of the ECM by proteases, proliferation and migration of ECs into the tumor, and the eventual formation of new capillary tubes.

Tumor blood vessels are not normal; they have chaotic architecture and blood flow. Due to an imbalance of angiogenic regulators such as VEGF and angiopoietins (see below), tumor vessels are tortuous and dilated with an uneven diameter, excessive branching, and shunting. Tumor blood flow is variable, with areas of hypoxemia and acidosis leading to the selection of variants that are resistant to hypoxemia-induced apoptosis (often due to the loss of p53 expression). Tumor vessel walls have numerous openings, widened interendothelial junctions, and discontinuous or absent basement membrane; this contributes to the high vascular permeability of these vessels and, together with lack of functional intratumoral lymphatics, causes increased interstitial pressure within the tumor (which also interferes with the delivery of therapeutics to the tumor; Figs. 102e-9, 102e-10, and 102e-11). Tumor blood vessels lack perivascular cells such as pericytes and smooth-muscle cells that normally regulate flow in response to tissue metabolic needs.

Unlike normal blood vessels, the vascular lining of tumor vessels is not a homogeneous layer of ECs but often consists of a mosaic of ECs and tumor cells with upregulated genes seen in ECs and vessel formation that can occur in hypoxic conditions because of their plasticity; the concept of cancer cell-derived vascular channels, which may be lined by ECM secreted by the tumor cells, is referred to as *vascular mimicry*. During tumor angiogenesis, ECs are highly proliferative and express a number of plasma membrane proteins that are characteristic of activated endothelium, including growth factor receptors and adhesion molecules such as integrins.

MECHANISMS OF TUMOR VESSEL FORMATION

Tumors use a number of mechanisms to promote vascularization, subverting normal angiogenic processes for this purpose (Fig. 102e-9). Primary or metastatic tumor cells sometimes arise in proximity to host blood vessels and grow around these vessels, parasitizing nutrients by co-opting the local blood supply. However, most tumor blood vessels arise by the process of *sprouting*, in which tumors secrete trophic

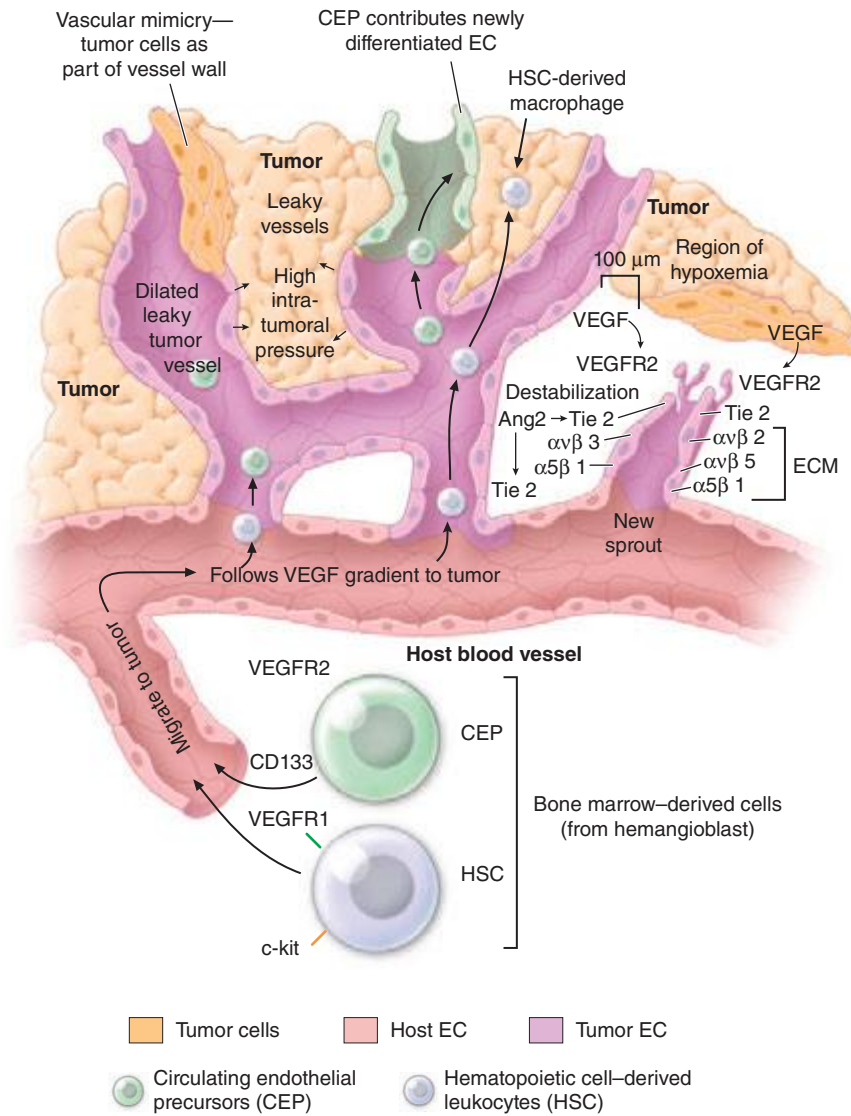


FIGURE 102e-9 Tumor angiogenesis is a complex process involving many different cell types that must proliferate, migrate, invade, and differentiate in response to signals from the tumor microenvironment. Endothelial cells (ECs) sprout from host vessels in response to VEGF, bFGF, Ang2, and other proangiogenic stimuli. Sprouting is stimulated by VEGF/VEGFR2, Ang2/Tie2, and integrin/extracellular matrix (ECM) interactions. Bone marrow-derived circulating endothelial precursors (CEPs) migrate to the tumor in response to VEGF and differentiate into ECs, while hematopoietic stem cells differentiate into leukocytes, including tumor-associated macrophages that secrete angiogenic growth factors and produce matrix metalloproteinases (MMPs) that remodel the ECM and release bound growth factors. Tumor cells themselves may directly form parts of vascular channels within tumors. The pattern of vessel formation is haphazard: vessels are tortuous, dilated, and leaky and branch in random ways. This leads to uneven blood flow within the tumor, with areas of acidosis and hypoxemia (which stimulate release of angiogenic factors) and high intratumoral pressures that inhibit delivery of therapeutic agents.

angiogenic molecules, the most potent being vascular endothelial growth factors (VEGF), that induce the proliferation and migration of host ECs into the tumor. Sprouting in normal and pathogenic angiogenesis is regulated by three families of transmembrane receptor tyrosine kinases (RTKs) expressed on ECs and their ligands (VEGFs, angiopoietins, ephrins; Fig. 102e-10), which are produced by tumor cells, inflammatory cells, or stromal cells in the tumor microenvironment.

When tumor cells arise in or metastasize to an avascular area, they grow to a size limited by hypoxemia and nutrient deprivation. Hypoxemia, a key regulator of tumor angiogenesis, causes the transcriptional induction of the gene encoding VEGF. VEGF and its receptors are required for embryonic *vasculogenesis* (development of new blood vessels when none preexist) and normal (wound healing, corpus luteum formation) and pathologic angiogenesis (tumor angiogenesis, inflammatory conditions such as rheumatoid arthritis). VEGF-A is a heparin-binding glycoprotein with at least four isoforms (splice variants) that

regulates blood vessel formation by binding to the RTKs VEGFR1 and VEGFR2, which are expressed on all ECs in addition to a subset of hematopoietic cells (Fig. 102e-9). VEGFR2 regulates EC proliferation, migration, and survival, whereas VEGFR1 may act as an antagonist of R2 in ECs but is probably also important for angioblast differentiation during embryogenesis. Tumor vessels may be more dependent on VEGFR signaling for growth and survival than normal ECs. Although VEGF signaling is a critical initiator of angiogenesis, this is a complex process regulated by additional signaling pathways (Fig. 102e-10). The angiopoietin, Ang1, produced by stromal cells, binds to the EC RTK Tie2 and promotes the interaction of ECs with the ECM and perivascular cells, such as pericytes and smooth-muscle cells, to form tight, nonleaky vessels. Platelet-derived growth factor (PDGF) and basic fibroblast growth factor (bFGF) help to recruit these perivascular cells. Ang1 is required for maintaining the quiescence and stability of mature blood vessels and prevents the vascular permeability normally induced by VEGF and inflammatory cytokines.

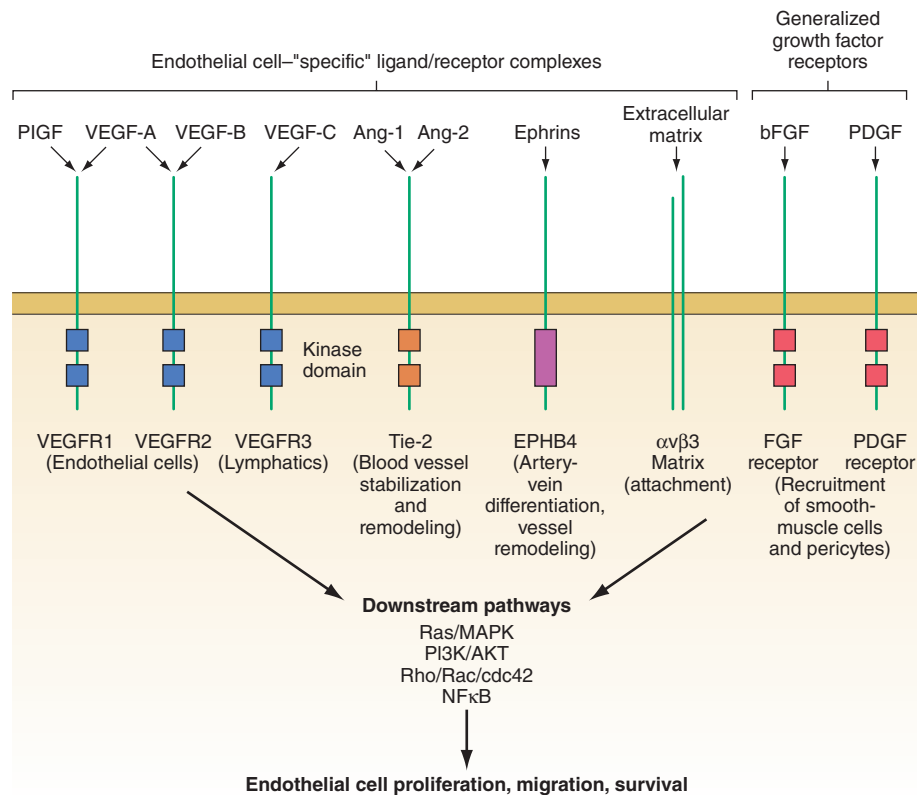


FIGURE 102e-10 Critical molecular determinants of endothelial cell biology. Angiogenic endothelium expresses a number of receptors not found on resting endothelium. These include receptor tyrosine kinases (RTKs) and integrins that bind to the extracellular matrix and mediate endothelial cell (EC) adhesion, migration, and invasion. ECs also express RTK (i.e., the FGF and PDGF receptors) that are found on many other cell types. Critical functions mediated by activated RTK include proliferation, migration, and enhanced survival of endothelial cells, as well as regulation of the recruitment of perivascular cells and bloodborne circulating endothelial precursors and hematopoietic stem cells to the tumor. Intracellular signaling via EC-specific RTK uses molecular pathways that may be targets for future antiangiogenic therapies.

For tumor cell-derived VEGF to initiate sprouting from host vessels, the stability conferred by the Ang1/Tie2 pathway must be perturbed; this occurs by the secretion of Ang2 by ECs that are undergoing active remodeling. Ang2 binds to Tie2 and is a competitive inhibitor of Ang1 action: under the influence of Ang2, preexisting blood vessels become more responsive to remodeling signals, with less adherence of ECs to stroma and associated perivascular cells and more responsiveness to VEGF. Therefore, Ang2 is required at early stages of tumor angiogenesis for destabilizing the vasculature by making host ECs more sensitive to angiogenic signals. Because tumor ECs are blocked by Ang2, there is no stabilization by the Ang1/Tie2 interaction, and tumor blood vessels are leaky, hemorrhagic, and have poor association of ECs with underlying stroma. Sprouting tumor ECs express high levels of the transmembrane protein ephrin-B2 and its receptor, the RTK EPH, whose signaling appears to work with the angiopoietins during vessel remodeling. During embryogenesis, EPH receptors are expressed on the endothelium of primordial venous vessels while the transmembrane ligand ephrin-B2 is expressed by cells of primordial arteries; the reciprocal expression may regulate differentiation and patterning of the vasculature.

A number of ubiquitously expressed host molecules play critical roles in normal and pathologic angiogenesis. Proangiogenic cytokines, chemokines, and growth factors secreted by stromal cells or inflammatory cells make important contributions to neovascularization, including bFGF, transforming growth factor α (TGF- α), TNF- α , and IL-8. In contrast to normal endothelium, angiogenic endothelium overexpresses specific members of the integrin family of ECM-binding proteins that mediate EC adhesion, migration, and survival. Specifically, expression of integrins $\alpha_v\beta_3$, $\alpha_v\beta_5$, and $\alpha_5\beta_1$ mediates spreading and migration of ECs and is required for angiogenesis induced by VEGF and bFGF, which in turn can upregulate EC integrin expression. The $\alpha_v\beta_3$ integrin physically associates with VEGFR2 in the plasma membrane and promotes signal

transduction from each receptor to promote EC proliferation (via focal adhesion kinase, src, PI3K, and other pathways) and survival (by inhibition of p53 and increasing the Bcl-2/Bax expression ratio). In addition, $\alpha_v\beta_3$ forms cell-surface complexes with matrix metalloproteinases (MMPs), zinc-requiring proteases that cleave ECM proteins, leading to enhanced EC migration and the release of heparin-binding growth factors, including VEGF and bFGF. EC adhesion molecules can be upregulated (i.e., by VEGF, TNF- α) or downregulated (by TGF- β); this, together with chaotic blood flow, explains poor leukocyte-endothelial interactions in tumor blood vessels and may help tumor cells avoid immune surveillance.

Lymphatic vessels also exist within tumors. Development of tumor lymphatics is associated with expression of VEGFR3 and its ligands VEGF-C and VEGF-D. The role of these vessels in tumor cell metastasis to regional lymph nodes remains to be determined. However, VEGF-C levels correlate significantly with metastasis to regional lymph nodes in lung, prostate, and colorectal cancers.

ANTIANGIOGENIC THERAPY

Angiogenesis inhibitors function by targeting the critical molecular pathways involved in EC proliferation, migration, and/or survival, many of which are unique to the activated endothelium in tumors. Inhibition of growth factor and adhesion-dependent signaling pathways can induce EC apoptosis with concomitant inhibition of tumor growth. Different types of tumors can use distinct combinations of molecular mechanisms to activate the angiogenic switch. Therefore, it is doubtful that a single antiangiogenic strategy will suffice for all human cancers; rather, a number of agents or combinations of agents will be needed, depending on distinct programs of angiogenesis used by different human cancers. Despite this, experimental data indicate that for some tumor types, blockade of a single growth factor (e.g., VEGF) may inhibit tumor-induced vascular growth.

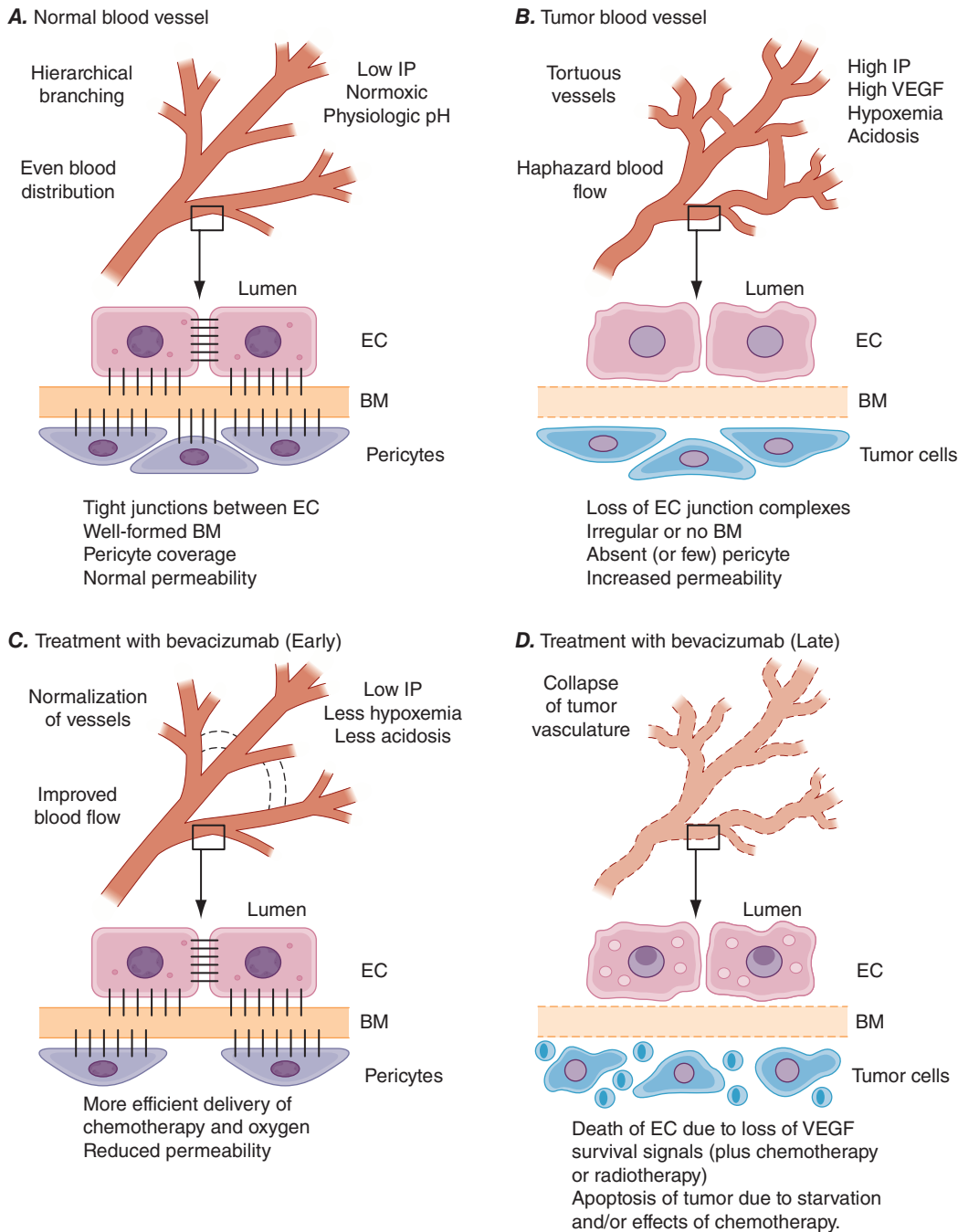


FIGURE 102e-11 Normalization of tumor blood vessels due to inhibition of VEGF signaling. **A.** Blood vessels in normal tissues exhibit a regular hierarchical branching pattern that delivers blood to tissues in a spatially and temporally efficient manner to meet the metabolic needs of the tissue (*top*). At the microscopic level, tight junctions are maintained between endothelial cells (ECs), which are adherent to a thick and evenly distributed basement membrane (BM). Pericytes form a surrounding layer that provides trophic signals to the EC and helps maintain proper vessel tone. Vascular permeability is regulated, interstitial fluid pressure is low, and oxygen tension and pH are physiologic. **B.** Tumors have abnormal vessels with tortuous branching and dilated, irregular interconnecting branches, causing uneven blood flow with areas of hypoxemia and acidosis. This harsh environment selects genetic events that result in resistant tumor variants, such as the loss of p53. High levels of VEGF (secreted by tumor cells) disrupt gap junction communication, tight junctions, and adherens junctions between EC via src-mediated phosphorylation of proteins such as connexin 43, zonula occludens-1, VE-cadherin, and α/β -catenins. Tumor vessels have thin, irregular BM, and pericytes are sparse or absent. Together, these molecular abnormalities result in a vasculature that is permeable to serum macromolecules, leading to high tumor interstitial pressure, which can prevent the delivery of drugs to the tumor cells. This is made worse by the binding and activation of platelets at sites of exposed BM, with release of stored VEGF and microvessel clot formation, creating more abnormal blood flow and regions of hypoxemia. **C.** In experimental systems, treatment with bevacizumab or blocking antibodies to VEGFR2 leads to changes in the tumor vasculature that has been termed *vessel normalization*. During the first week of treatment, abnormal vessels are eliminated or pruned (*dotted lines*), leaving a more normal branching pattern. ECs partially regain features such as cell-cell junctions, adherence to a more normal BM, and pericyte coverage. These changes lead to a decrease in vascular permeability, reduced interstitial pressure, and a transient increase in blood flow within the tumor. Note that in murine models, this normalization period lasts only for ~5–6 days. **D.** After continued anti-VEGF/VEGFR therapy (which is often combined with chemo- or radiotherapy), ECs die, leading to tumor cell death (either due to direct effects of the chemotherapy or lack of blood flow).

Bevacizumab, an antibody that binds VEGF, appears to potentiate the effects of a number of different types of active chemotherapeutic regimens used to treat a variety of different tumor types including colon cancer, lung cancer, cervical cancer, and RCC.

Bevacizumab is administered IV every 2–3 weeks (its half-life is nearly 20 days) and is generally well tolerated. Hypertension is the most common side effect of inhibitors of VEGF (or its receptors), but can be treated with antihypertensive agents and rarely requires discontinuation of therapy. Rare but serious potential risks include arterial thromboembolic events, including stroke and myocardial infarction, and hemorrhage. Another serious complication is bowel perforation, which has been observed in 1–3% of patients (mainly those with colon and ovarian cancers). Inhibition of wound healing is also seen.

Several small-molecule inhibitors (SMIs) that target VEGFR tyrosine kinase activity but are also inhibitory to other kinases have also been approved to treat certain cancers. Sunitinib (see above and Table 102e-2) has activity directed against mutant c-Kit receptors (approved for GIST), but also targets VEGFR and PDGFR, and has shown significant antitumor activity against metastatic RCC, presumably on the basis of its antiangiogenic activity. Similarly, sorafenib, originally developed as a Raf kinase inhibitor but with potent activity against VEGFR and PDGFR, has activity against RCC, thyroid cancer, and hepatocellular cancer. Other inhibitors of VEGFR approved for the treatment of RCC include axitinib and pazopanib.

The success in targeting tumor angiogenesis has led to enhanced enthusiasm for the development of drugs that target other aspects of the angiogenic process; some of these therapeutic approaches are outlined in Fig. 102e-12.

EVASION OF THE IMMUNE SYSTEM BY CANCERS

Cancers have a number of mechanisms that allow them to evade detection and elimination by the immune system. These include downregulation of cell surface proteins involved in immune recognition (including MHC proteins and tumor-specific antigens), expression of other cell surface proteins that inhibit immune function (including members of the B7 family of proteins such as PD-L1), secretion of proteins and other molecules that are immunosuppressive, recruitment and expansion of immunosuppressive cells such as regulatory T cells, and induction of T cell tolerance. In addition, the inflammatory effects of some of the immune mediator cells in the tumor microenvironment (especially tissue-associated macrophages and myeloid-derived suppressor cells) can suppress T cell responses to the tumor as well as stimulate inflammation that can enhance tumor growth.

Immunotherapy approaches to treat cancer aimed at activating the immune response against tumors using immunostimulatory molecules such as interferons, IL-2, and monoclonal antibodies have had some successes. Another approach that has shown particular clinical promise is the targeting of proteins or cells (such as regulatory T cells) involved in normal homeostatic control to prevent autoimmune damage to the host but that malignant cells and their stroma can also use to inhibit the immune response directed against them. The approach that is furthest along clinically has involved targeting CTLA-4, PD-1, and PDL-1, co-inhibitory molecules that are expressed on the surface of cancer cells, cells of the immune system, and/or stromal cells and are involved in inhibiting the immune response against cancer (Fig. 102e-13). Monoclonal antibodies directed against CTLA-4 and PD-1 are approved for the treatment of melanoma, and additional antibodies

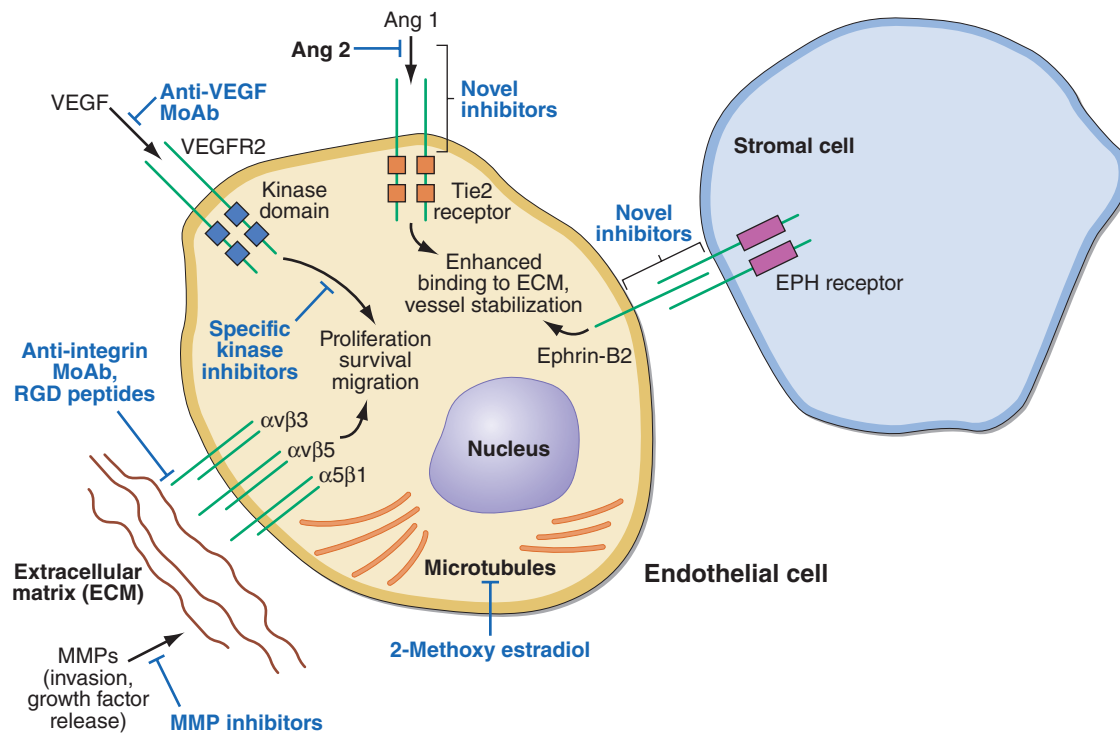


FIGURE 102e-12 Knowledge of the molecular events governing tumor angiogenesis has led to a number of therapeutic strategies to block tumor blood vessel formation. The successful therapeutic targeting of VEGF is described in the text. Other endothelial cell-specific receptor tyrosine kinase pathways (e.g., angiopoietin/Tie2 and ephrin/EPH) are likely targets for the future. Ligand of the $\alpha_v\beta_3$ integrin is required for endothelial cell (EC) survival. Integrins are also required for EC migration and are important regulators of matrix metalloproteinase (MMP) activity, which modulates EC movement through the extracellular matrix (ECM) as well as release of bound growth factors. Targeting of integrins includes development of blocking antibodies, small peptide inhibitors of integrin signaling, and arg-gly-asp-containing peptides that prevent integrin:ECM binding. Peptides derived from normal proteins by proteolytic cleavage, including endostatin and tumstatin, inhibit angiogenesis by mechanisms that include interfering with integrin function. Signal transduction pathways that are dysregulated in tumor cells indirectly regulate EC function. Inhibition of EGF-family receptors, whose signaling activity is upregulated in a number of human cancers (e.g., breast, colon, and lung cancers), results in downregulation of VEGF and IL-8, while increasing expression of the antiangiogenic protein thrombospondin-1. The Ras/MAPK, PI3K/Akt, and Src kinase pathways constitute important antitumor targets that also regulate the proliferation and survival of tumor-derived EC. The discovery that ECs from normal tissues express tissue-specific “vascular addressins” on their cell surface suggests that targeting specific EC subsets may be possible.

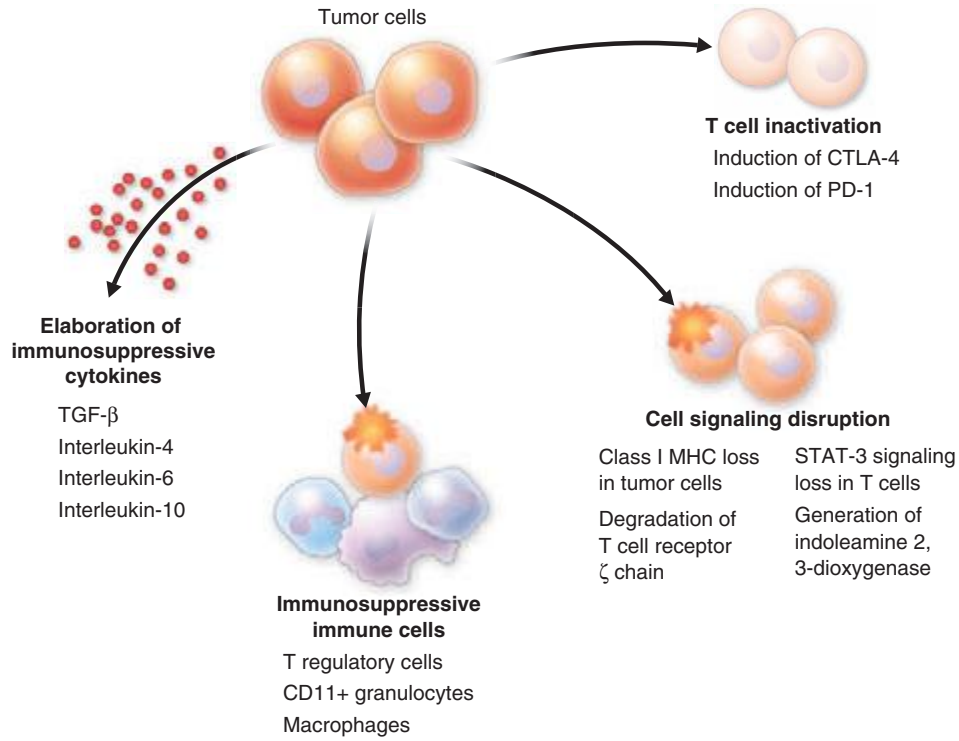


FIGURE 102e-13 Tumor-host interactions that suppress the immune response to the tumor.

targeting PD-1 or PDL-1 have shown activity against melanoma, RCC, and lung cancer and continue to be evaluated against other malignancies as well. Combination approaches targeting more than one protein or involving other anticancer approaches (targeted agents, chemotherapy, radiation therapy) are also being explored and have shown promise in early studies. An important aspect of these approaches is balancing sufficient release of the negative control of the immune response to allow immune-mediated attack on the tumors while not allowing too much release and inducing severe autoimmune effects (such as against skin, thyroid, pituitary gland, or the gastrointestinal tract).

SUMMARY

The explosion of information on tumor cell biology, metastasis, and tumor-host interactions (including angiogenesis and immune evasion by tumors) has ushered in a new era of rational targeted therapy

for cancer. Furthermore, it has become clear that specific molecular factors detected in individual tumors (specific gene mutations, gene-expression profiles, microRNA expression, overexpression of specific proteins) can be used to tailor therapy and maximize antitumor effects.

ACKNOWLEDGMENT

Robert G. Fenton contributed to this chapter in prior editions, and important material from those prior chapters has been included here.

CANCER PRESENTATION

Cancer in a localized or systemic state is a frequent item in the differential diagnosis of a variety of common complaints. Although not all forms of cancer are curable at diagnosis, affording patients the greatest opportunity for cure or meaningful prolongation of life is greatly aided by diagnosing cancer at the earliest point possible in its natural history and defining treatments that prevent or retard its systemic spread. Indeed, certain forms of cancer, notably breast, colon, and possibly lung cancers in certain patients, can be prevented by screening appropriately selected asymptomatic patients; screening is arguably the earliest point in the spectrum of possible cancer-related interventions where cure is possible (Table 103e-1).

DETECTION OF A CANCER

The term *cancer*, as used here, is synonymous with the term *tumor*, whose original derivation from Latin simply meant “swelling,” not otherwise specified. We now understand that the swelling that is a common physical manifestation of a tumor derives from increased interstitial fluid pressure and increased cellular and stromal mass per volume, compared to normal tissue. Tumors historically were referred to as *carcinomas*, or “crab-like” infiltrating tumors, or *sarcomas*, or “fleshy tumors,” derived from the Greek terms for “crab” and “flesh,” respectively. Leukemias are a special case of a cancer of the blood-forming tissues presenting in a disseminated form frequently without definable tumor masses. In addition to localized swelling, tumors present by altered function of the organ they afflict, such as dyspnea on exertion from the anemia caused by leukemia replacing normal hematopoietic cells, cough from lung cancers, jaundice from tumors disrupting the hepatobiliary tree, or seizures and neurologic signs from brain tumors. Hemorrhage is also a frequent presenting sign of tumors involving hollow viscera, as are decreases in the number of platelets and inappropriate inhibition of blood coagulation. Thus, although statistically the fraction of patients with cancer underlying a particular presenting sign or symptom may be low, the implications for a patient with cancer of missing an early-stage tumor call for vigilance; therefore, persistent signs or symptoms should be evaluated as possibly coming from an early-stage tumor.

Evidence of a tumor’s existence can objectively be established by careful physical examination, such as enlarged lymph nodes in lymphomas or a palpable mass in a breast or soft tissue site. A mass

TABLE 103e-1 SPECTRUM OF CANCER-RELATED INTERVENTIONS

Screening for cancer in an asymptomatic patient
Consideration of cancer in a differential diagnosis
Physical examination, imaging, or endoscopy to define a possible tumor
Diagnosis of cancer by biopsy or removal:
Routine histology
Specialized histology: immunohistochemistry
Molecular studies
Cytogenetic studies
Staging the cancer: Where has it spread?
Treatment
Localized
Systemic
Supportive care
During treatment: related to tumor effects on patient
During treatment to counteract side effects of treatment
Palliative and end of life
When useful treatments are not feasible or desired

may also be detected or confirmed by an imaging modality, such as plain x-ray, computed tomography (CT) scan, ultrasound, positron emission tomography (PET) imaging, or nuclear magnetic resonance approaches. Sensitivity of these technologies varies considerably, and the index of suspicion for a tumor should match the technology chosen. For example, low-dose helical CT scans are superior to plain chest radiographs in detecting lung cancers. Another way of initially establishing the existence of a possible tumor is through direct visualization of an afflicted organ by endoscopy.

ESTABLISHING A CANCER DIAGNOSIS

Once the existence of a likely tumor is defined, unequivocally establishing the diagnosis is the next step in the spectrum of correctly addressing a patient’s needs. This is usually accomplished by a biopsy procedure and the emergence after pathologic examination of an unequivocal statement that cancer is present. The underlying principle in cancer diagnosis is to obtain as much tissue as safely as possible. Due to tumor heterogeneity, pathologists are better able to make the diagnosis when they have more tissue to examine. In addition to light microscopic inspection of a tumor for pattern of growth, degree of cellular atypia, invasiveness, and morphologic features that aid in the differential diagnosis, sufficient tissue is of value in searching for genetic abnormalities and protein expression patterns, such as hormone receptor expression in breast cancers, that may aid in the differential diagnosis or provide information about prognosis or likely response to treatment. Efforts to define “personalized” information from the biology of each patient’s tumor and pertinent to each patient’s treatment plan are becoming increasingly important in selecting treatment options. The general internist should make sure that a patient’s cancer biopsy is appropriately referred from the surgical suite for important molecular studies that can advise the best treatment (Table 103e-2).

Similar-appearing tumors by microscopic morphology may have very different gene expression patterns when assessed by such techniques as microarray analysis for gene expression patterns using gene

TABLE 103e-2 DIAGNOSTIC BIOPSY: STANDARD OF CARE MOLECULAR AND SPECIAL STUDIES

Breast cancer: primary and suspected metastatic
Hormone receptors: estrogen, progesterone
HER2/neu oncoprotein
Lung cancer: primary and suspected metastatic
If nonsquamous non-small cell: epidermal growth factor receptor mutation; alk oncoprotein gene fusion
Colon cancer: suspected metastatic
Ki-ras mutation
Gastrointestinal stromal tumor
c-kit oncoprotein mutation
Melanoma
B-raf oncoprotein mutation
c-kit expression and mutation
Leukemia (peripheral blood mononuclear cells and/or bone marrow)
Cytogenetics
Flow cytometry
Treatment-defining chromosomal translocations
Bcr-Abl fusion protein
t(15,17)
inversion 16
t(8,21)
Lymphoma
Immunohistochemistry for CD20, CD30, T cell markers
Treatment defining chromosomal translocations:
t(14,18)
t(8,14)

chips, with important differences in biology and response to treatment. Such testing requires that the tissue be handled properly (e.g., immunologic detection of proteins is more effective in fresh-frozen tissue rather than in formalin-fixed tissue). Coordination among the surgeon, pathologist, and primary care physician is essential to ensure that the amount of information learned from the biopsy material is maximized. These goals are best met by an *excisional biopsy* in which the entire tumor mass is removed with a small margin of normal tissue surrounding it. If an excisional biopsy cannot be performed, *incisional biopsy* is the procedure of second choice. A wedge of tissue is removed, and an effort is made to include the majority of the cross-sectional diameter of the tumor in the biopsy to minimize sampling error. Biopsy techniques that involve cutting into tumor carry with them a risk of facilitating the spread of the tumor, and consideration of whether the biopsy might be the prelude to a curative surgery if certain diagnoses are established should inform the actual approach taken. *Core-needle biopsy* usually obtains considerably less tissue, but this procedure often provides enough information to plan a definitive surgical procedure. *Fine-needle aspiration* generally obtains only a suspension of cells from within a mass. This procedure is minimally invasive, and if positive for cancer, it may allow inception of systemic treatment when metastatic disease is evident, or it can provide a basis for planning a more meticulous and extensive surgical procedure. However, a negative fine-needle aspiration for a neoplastic diagnosis cannot be taken as definitive evidence that a tumor is absent or make a definitive diagnosis in someone not known to have a cancer.

CANCER STAGING

An essential component of correct patient management in many cancer types is defining the extent of disease, because this information critically informs whether localized treatments, “combined-modality” approaches, or systemic treatments should initially be considered. Radiographic and other imaging tests can be helpful in defining the clinical stage; however, pathologic staging requires defining the extent of involvement by documenting the histologic presence of tumor in tissue biopsies obtained through a surgical procedure. Axillary lymph node sampling in breast cancer and lymph node sampling at laparotomy for testicular, colon, and other intraabdominal cancers may provide crucial information for treatment planning and may determine the extent and nature of primary cancer treatment.

For tumors associated with a potential “primary site,” staging systems have evolved to define a “T” component related to the size of the tumor or its invasion into local structures, an “N” component related to the number and nature of lymph node groups adjacent to the tumor with evidence of tumor spread, and an “M” component, based on the presence of local or distant metastatic sites. The various “TNM” components are then aggregated to stages, usually stage I to III or IV, depending on the anatomic site. The numerical stages reflect similar long-term survival outcomes of the aggregated TNM groupings in a numeric stage after treatment tailored to the stage. In general, stage I tumors are T1 (reflecting small size), N0 or N1 (reflecting no or minimal node spread), and M0 (no metastases). Such early-stage tumors are amenable to curative approaches with local treatments. On the other hand, stage IV tumors usually have metastasized to distant sites or locally invaded viscera in a nonresectable way and are dealt with using techniques that have palliative intent, except for those diseases with exceptional sensitivity to systemic treatments such as chemotherapy or immunotherapy. Also, the TNM staging system is not useful in diseases such as leukemia, where bone marrow infiltration is never really localized, or central nervous system tumors, where tumor histology and the extent of anatomic feasible resection are more important in driving prognosis.

CANCER TREATMENT

The goal of cancer treatment is first to eradicate the cancer. If this primary goal cannot be accomplished, the goal of cancer treatment shifts to palliation, the amelioration of symptoms, and preservation of quality of life while striving to extend life. The dictum *primum non nocere*

may not always be the guiding principle of cancer therapy. When cure of cancer is possible, cancer treatments may be considered despite the certainty of severe and perhaps life-threatening toxicities. Every cancer treatment has the potential to cause harm, and treatment may be given that produces toxicity with no benefit. The therapeutic index of many interventions may be quite narrow, with treatments given to the point of toxicity. Conversely, when the clinical goal is palliation, careful attention to minimizing the toxicity of potentially toxic treatments becomes a significant goal.

Cancer treatments are divided into two main types: local and systemic. Local treatments include surgery, radiation therapy (including photodynamic therapy), and ablative approaches, including radiofrequency and cryosurgical approaches. Systemic treatments include chemotherapy (including hormonal therapy and molecularly targeted therapy) and biologic therapy (including immunotherapy). The modalities are often used in combination, and agents in one category can act by several mechanisms. For example, cancer chemotherapy agents can induce differentiation, and antibodies (a form of immunotherapy) can be used to deliver radiation therapy. *Oncology*, the study of tumors including treatment approaches, is a multidisciplinary effort with surgical, radiation, and internal medicine–related areas of oncologic expertise. Treatments for patients with hematologic malignancies are often shared by hematologists and medical oncologists.

In many ways, cancer mimics an organ attempting to regulate its own growth. However, cancers have not set an appropriate limit on how much growth should be permitted. Normal organs and cancers share the property of having (1) a population of cells actively progressing through the cell cycle with their division providing a basis for tumor growth, and (2) a population of cells not in cycle. In cancers, cells that are not dividing are heterogeneous; some have sustained too much genetic damage to replicate but have defects in their death pathways that permit their survival, some are starving for nutrients and oxygen, and some are out of cycle but poised to be recruited back into cycle and expand if needed (i.e., reversibly growth-arrested). Severely damaged and starving cells are unlikely to kill the patient. The problem is that the cells that are reversibly not in cycle are capable of replenishing tumor cells physically removed or damaged by radiation and chemotherapy. These include *cancer stem cells*, whose properties are being elucidated, as they may serve as a basis for giving rise to tumor initiating or repopulating cells. The stem cell fraction may define new targets for therapies that will retard their ability to reenter the cell cycle.

Tumors follow a Gompertzian growth curve (Fig. 103e-1), with the apparent growth fraction of a neoplasm being high with small tumor burdens and declining until, at the time of diagnosis, with a tumor burden of $1-5 \times 10^9$ tumor cells, the growth fraction is usually 1–4% for many solid tumors. By this view, the most rapid growth rate occurs before the tumor is detectable. An alternative explanation for such growth properties may also emerge from the ability of tumors at metastatic sites to recruit circulating tumor cells from the primary tumor or other metastases. An additional key feature of a successful tumor is the ability to stimulate the development of a new supporting stroma through angiogenesis and production of proteases to allow invasion through basement membranes and normal tissue barriers (Chap. 102e). Specific cellular mechanisms promote entry or withdrawal of tumor cells from the cell cycle. For example, when a tumor recurs after surgery or chemotherapy, frequently its growth is accelerated and the growth fraction of the tumor is increased. This pattern is similar to that seen in regenerating organs. Partial resection of the liver results in the recruitment of cells into the cell cycle, and the resected liver volume is replaced. Similarly, chemotherapy-damaged bone marrow increases its growth to replace cells killed by chemotherapy. However, cancers do not recognize a limit on their expansion. Monoclonal gammopathy of uncertain significance may be an example of a clonal neoplasm with intrinsic features that stop its growth before a lethal tumor burden is reached. A fraction of patients with this disorder go on to develop fatal multiple myeloma, but probably this occurs because of the accumulation of additional genetic lesions. Elucidation of the mechanisms that regulate this “organ-like” behavior of tumors may provide additional clues to cancer control and treatment.

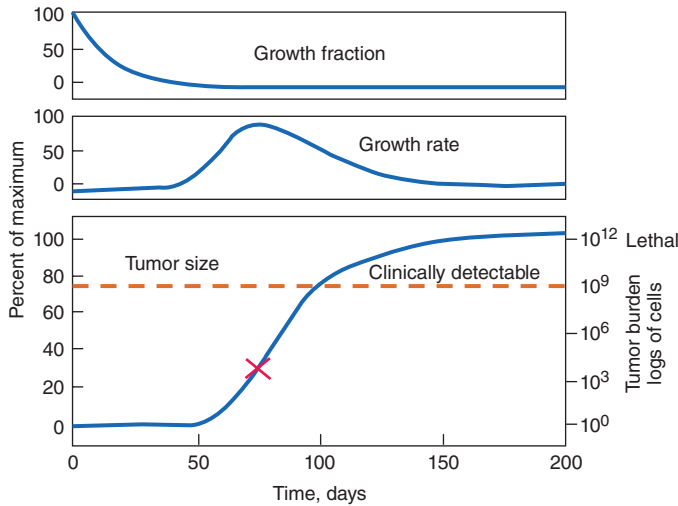


FIGURE 103e-1 Gompertzian tumor growth. The growth fraction of a tumor declines exponentially over time (*top*). The growth rate of a tumor peaks before it is clinically detectable (*middle*). Tumor size increases slowly, goes through an exponential phase, and slows again as the tumor reaches the size at which limitation of nutrients or autoregulatory or host regulatory influences can occur. The maximum growth rate occurs at $1/e$, the point at which the tumor is about 37% of its maximum size (*marked with an X*). Tumor becomes detectable at a burden of about 10^9 (1 cm^3) cells and kills the patient at a tumor cell burden of about 10^{12} (1 kg). Efforts to treat the tumor and reduce its size can result in an increase in the growth fraction and an increase in growth rate.

LOCALIZED CANCER TREATMENTS

SURGERY

Surgery is unquestionably the most effective means of treating cancer. Today at least 40% of cancer patients are cured by surgery. Unfortunately, a large fraction of patients with solid tumors (perhaps 60%) have metastatic disease that is not accessible for removal. However, even when the disease is not curable by surgery alone, the removal of tumor can obtain important benefits, including local control of tumor, preservation of organ function, debulking that permits subsequent therapy to work better, and staging information on extent of involvement. Cancer surgery aiming for cure is usually planned to excise the tumor completely with an adequate margin of normal tissue (the margin varies with the tumor and the anatomy), touching the tumor as little as possible to prevent vascular and lymphatic spread, and minimizing operative risk. Such a resection is defined as an R0 resection. R1 and R2 resections, in contrast, are imprecisely defined pathologically as having microscopic or macroscopic tumor at resection margins. Such outcomes may be necessitated by proximity of the tumor to vital structures or recognition only in the resected specimen of the extent of tumor involvement, and may be the basis for reoperation to obtain optimal margins if feasible. Extending the procedure to resect draining lymph nodes obtains prognostic information and may, in some anatomic locations, improve survival.

Increasingly, laparoscopic approaches are being used to address primary abdominal and pelvic tumors. Lymph node spread may be assessed using the sentinel node approach, in which the first draining lymph node a spreading tumor would encounter is defined by injecting a dye or radioisotope into the tumor site at operation and then resecting the first node to turn blue or collect label. The sentinel node assessment is continuing to undergo clinical evaluation but appears to provide reliable information without the risks (lymphedema, lymphangiosarcoma) associated with resection of all the regional nodes. Advances in adjuvant chemotherapy (chemotherapy given systemically after removal of all disease by operation and without evidence of active metastatic disease) and radiation therapy following surgery have permitted a substantial decrease in the extent of primary surgery

necessary to obtain the best outcomes. Thus, lumpectomy with radiation therapy is as effective as modified radical mastectomy for breast cancer, and limb-sparing surgery followed by adjuvant radiation therapy and chemotherapy has replaced radical primary surgical procedures involving amputation and disarticulation for childhood rhabdomyosarcomas and osteosarcomas. More limited surgery is also being used to spare organ function, as in larynx and bladder cancer. The magnitude of operations necessary to optimally control and cure cancer has also been diminished by technical advances; for example, the circular anastomotic stapler has allowed narrower ($<2 \text{ cm}$) margins in colon cancer without compromise of local control rates, and many patients who would have had colostomies are able to maintain normal anatomy.

In some settings (e.g., bulky testicular cancer or stage III breast cancer), surgery is not the first treatment modality used. After an initial diagnostic biopsy, chemotherapy and/or radiation therapy is delivered to reduce the size of the tumor and clinically control undetected metastatic disease. Such therapy is followed by a surgical procedure to remove residual masses; this is called *neoadjuvant therapy*. Because the sequence of treatment is critical to success and is different from the standard surgery-first approach, coordination among the surgical oncologist, radiation oncologist, and medical oncologist is crucial.

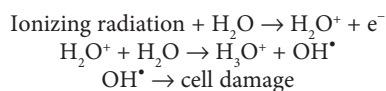
Surgery may be curative in a subset of patients with metastatic disease. Patients with lung metastases from osteosarcoma may be cured by resection of the lung lesions. In patients with colon cancer who have fewer than five liver metastases restricted to one lobe and no extrahepatic metastases, hepatic lobectomy may produce long-term disease-free survival in 25% of selected patients. Surgery can also be associated with systemic antitumor effects. In the setting of hormonally responsive tumors, oophorectomy and/or adrenalectomy may eliminate estrogen production, and orchiectomy may reduce androgen production, hormones that drive certain breast and all prostate cancers, respectively; both procedures can have useful effects on metastatic tumor growth. If resection of the primary lesion takes place in the presence of metastases, acceleration of metastatic growth has also been described in certain cases, perhaps based on the removal of a source of angiogenesis inhibitors and mass-related growth regulators in the tumor.

In selecting a surgeon or center for primary cancer treatment, consideration must be given to the volume of cancer surgeries undertaken by the site. Studies in a variety of cancers have shown that increased annual procedure volume appears to correlate with outcome. In addition, facilities with extensive support systems—e.g., for joint thoracic and abdominal surgical teams with cardiopulmonary bypass, if needed—may allow resection of certain tumors that would otherwise not be possible.

Surgery is used in a number of ways for palliative or supportive care of the cancer patient, not related to the goal of curing the cancer. These include insertion and care of central venous catheters, control of pleural and pericardial effusions and ascites, caval interruption for recurrent pulmonary emboli, stabilization of cancer-weakened weight-bearing bones, and control of hemorrhage, among others. Surgical bypass of gastrointestinal, urinary tract, or biliary tree obstruction can alleviate symptoms and prolong survival. Surgical procedures may provide relief of otherwise intractable pain or reverse neurologic dysfunction (cord decompression). Splenectomy may relieve symptoms and reverse hypersplenism. Intrathecal or intrahepatic therapy relies on surgical placement of appropriate infusion portals. Surgery may correct other treatment-related toxicities such as adhesions or strictures. Surgical procedures are also valuable in rehabilitative efforts to restore health or function. Orthopedic procedures may be necessary to ensure proper ambulation. Breast reconstruction can make an enormous impact on the patient's perception of successful therapy. Plastic and reconstructive surgery can correct the effects of disfiguring primary treatment.

Surgery is also a tool valuable in the prevention of cancers in high-risk populations. Prophylactic mastectomy, colectomy, oophorectomy, and thyroidectomy are mainstays of prevention of genetic cancer syndromes. Resection of premalignant skin and uterine cervix lesions and colonic polyps prevents progression to frank malignancy.

Radiation Biology and Medicine Therapeutic radiation is ionizing; it damages any tissue in its path. The selectivity of radiation for causing cancer cell death may be due to defects in a cancer cell's ability to repair sublethal DNA and other damage. Ionizing radiation causes breaks in DNA and generates free radicals from cell water that may damage cell membranes, proteins, and organelles. Radiation damage is augmented by oxygen; hypoxic cells are more resistant. Augmentation of oxygen presence is one basis for radiation sensitization. Sulfhydryl compounds interfere with free radical generation and may act as radiation protectors. X-rays and gamma rays are the forms of ionizing radiation most commonly used to treat cancer. They are both electromagnetic, nonparticulate waves that cause the ejection of an orbital electron when absorbed. This orbital electron ejection is called *ionization*. X-rays are generated by linear accelerators; gamma rays are generated from decay of atomic nuclei in radioisotopes such as cobalt and radium. These waves behave biologically as packets of energy, called *photons*. Particulate ionizing radiation using protons has also become available. Most radiation-induced cell damage is due to the formation of hydroxyl radicals from tissue water:



Radiation is quantitated based on the amount of radiation absorbed by the tumor in the patient; it is not based on the amount of radiation generated by the machine. The International System (SI) unit for radiation absorbed is the Gray (Gy): 1 Gy refers to 1 J/kg of tissue; 1 Gy equals 100 centigrays (cGy) of absorbed dose. A historically used unit appearing in the oncology literature, the *rad* (radiation absorbed dose), is defined as 100 ergs of energy absorbed per gram of tissue and is equivalent to 1 cGy. Radiation dosage is defined by the energy absorbed per mass of tissue. Radiation dose is measured by placing detectors at the body surface or based on radiating phantoms that resemble human form and substance, containing internal detectors. The features that make a particular cell more sensitive or more resistant to the biologic effects of radiation are not completely defined and critically involve DNA repair proteins that, in their physiologic role, protect against environmentally related DNA damage.

Localized Radiation Therapy Radiation effect is influenced by three determinants: total absorbed dose, number of fractions, and time of treatment. A frequent error is to omit the number of fractions and the duration of treatment. This is analogous to saying that a runner completed a race in 20 s; without knowing how far he or she ran, the result is difficult to interpret. The time could be very good for a 200-m race or very poor for a 100-m race. Thus, a typical course of radiation therapy should be described as 4500 cGy delivered to a particular target (e.g., mediastinum) over 5 weeks in 180-cGy fractions. Most curative radiation treatment programs are delivered once a day, 5 days a week, in 150- to 200-cGy fractions.

A number of parameters influence the damage done to tissue (normal and tumor) by radiation. Hypoxic cells are relatively resistant. Nondividing cells are more resistant than dividing cells, and this is one rationale for delivering radiation in repeated fractions, to ultimately expose a larger number of tumor cells that have entered the division cycle. In addition to these biologic parameters, physical parameters of the radiation are also crucial. The energy of the radiation determines its ability to penetrate tissue. Low-energy orthovoltage beams (150–400 kV) scatter when they strike the body, much like light diffuses when it strikes particles in the air. Such beams result in more damage to adjacent normal tissues and less radiation delivered to the tumor. Megavoltage radiation (>1 MeV) has very low lateral scatter; this produces a skin-sparing effect, more homogeneous distribution of the radiation energy, and greater deposit of the energy in the tumor, or *target volume*. The tissues that the beam passes through to get to the tumor are called the *transit volume*. The maximum dose in the target volume is often the cause of complications to tissues in the transit volume, and the minimum dose in the target volume influences

the likelihood of tumor recurrence. Dose homogeneity in the target volume is the goal. Computational approaches and delivery of many beams to converge on a target lesion are the basis for “gamma knife” and related approaches to deliver high doses to small volumes of tumor, sparing normal tissue.

Therapeutic radiation is delivered in three ways: (1) *teletherapy*, with focused beams of radiation generated at a distance and aimed at the tumor within the patient; (2) *brachytherapy*, with encapsulated sources of radiation implanted directly into or adjacent to tumor tissues; and (3) *systemic therapy*, with radionuclides administered, for example, intravenously but targeted by some means to a tumor site. Teletherapy with x-ray or gamma-ray photons is the most commonly used form of radiation therapy. Particulate forms of radiation are also used in certain circumstances, such as the use of proton beams. The difference between photons and protons relates to the volume in which the greatest delivery of energy occurs. Typically protons have a much narrower range of energy deposition, theoretically resulting in more precise delivery of radiation with improvement in the degree to which adjacent structures may be affected, in comparison to photons. Electron beams are a particulate form of radiation that, in contrast to photons and protons, have a very low tissue penetrance and are used to treat cutaneous tumors. Apart from sparing adjacent structures, particulate forms of radiation are in most applications not superior to x-rays or gamma rays in clinical studies reported thus far, but this is an active area of investigation.

Certain drugs used in cancer treatment may also act as radiation sensitizers. For example, compounds that incorporate into DNA and alter its stereochemistry (e.g., halogenated pyrimidines, cisplatin) augment radiation effects at local sites, as does hydroxyurea, another DNA synthesis inhibitor. These are important adjuncts to the local treatment of certain tumors, such as squamous head and neck, uterine cervix, and rectal cancers.

Toxicity of Radiation Therapy Although radiation therapy is most often administered to a local region, systemic effects, including fatigue, anorexia, nausea, and vomiting, may develop that are related in part to the volume of tissue irradiated, dose fractionation, radiation fields, and individual susceptibility. Injured tissues release cytokines that act systemically to produce these effects. Bone is among the most radio-resistant organs, with radiation effects being manifested mainly in children through premature fusion of the epiphyseal growth plate. By contrast, the male testis, female ovary, and bone marrow are the most sensitive organs. Any bone marrow in a radiation field will be eradicated by therapeutic irradiation. Organs with less need for cell renewal, such as heart, skeletal muscle, and nerves, are more resistant to radiation effects. In radiation-resistant organs, the vascular endothelium is the most sensitive component. Organs with more self-renewal as a part of normal homeostasis, such as the hematopoietic system and mucosal lining of the intestinal tract, are more sensitive. Acute toxicities include mucositis, skin erythema (ulceration in severe cases), and bone marrow toxicity. Often these can be alleviated by interruption of treatment.

Chronic toxicities are more serious. Radiation of the head and neck region often produces thyroid failure. Cataracts and retinal damage can lead to blindness. Salivary glands stop making saliva, which leads to dental caries and poor dentition. Taste and smell can be affected. Mediastinal irradiation leads to a threefold increased risk of fatal myocardial infarction. Other late vascular effects include chronic constrictive pericarditis, lung fibrosis, viscus stricture, spinal cord transection, and radiation enteritis. A serious late toxicity is the development of second solid tumors in or adjacent to the radiation fields. Such tumors can develop in any organ or tissue and occur at a rate of about 1% per year beginning in the second decade after treatment. Some organs vary in susceptibility to radiation carcinogenesis. A woman who receives mantle field radiation therapy for Hodgkin's disease at age 25 years has a 30% risk of developing breast cancer by age 55 years. This is comparable in magnitude to genetic breast cancer syndromes. Women treated after age 30 years have little or no increased risk of breast cancer. No data suggest that a threshold dose of therapeutic radiation exists below which the incidence of second

cancers is decreased. High rates of second tumors occur in people who receive as little as 1000 cGy.

OTHER LOCALIZED CANCER TREATMENTS

Endoscopy techniques may allow the placement of stents to unblock viscera by mechanical means, palliating, for example, gastrointestinal or biliary obstructions. Radiofrequency ablation (RFA) refers to the use of focused microwave radiation to induce thermal injury within a volume of tissue. RFA can be useful in the control of metastatic lesions, particularly in liver, that may threaten biliary drainage (as one example) and threaten quality and duration of useful life in patients with otherwise unresectable disease. Cryosurgery uses extreme cold to sterilize lesions in certain sites, such as prostate and kidney, when at a very early stage, eliminating the need for modalities with more side effects such as surgery or radiation.

Some chemicals (porphyrins, phthalocyanines) are preferentially taken up by cancer cells by mechanisms not fully defined. When light, usually delivered by a laser, is shone on cells containing these compounds, free radicals are generated and the cells die. Hematoporphyrins and light (phototherapy) are being used with increasing frequency to treat skin cancer; ovarian cancer; and cancers of the lung, colon, rectum, and esophagus. Palliation of recurrent locally advanced disease can sometimes be dramatic and last many months.

Infusion of chemotherapeutic or biologic agents or radiation-bearing delivery devices such as isotope-coated glass spheres into local sites through catheters inserted into specific vascular sites such as liver or an extremity have been used in an effort to control disease limited to that site; in selected cases, prolonged control of truly localized disease has been possible.

SYSTEMIC CANCER TREATMENTS

The concept that systemically administered agents may have a useful effect on cancers was historically derived from three sets of observations. Paul Ehrlich in the nineteenth century observed that different dyes reacted with different cell and tissue components. He hypothesized the existence of compounds that would be “magic bullets” that might bind to tumors, owing to the affinity of the agent for the tumor. A second observation was the toxic effects of certain mustard gas derivatives on the bone marrow during World War I, leading to the idea that smaller doses of these agents might be used to treat tumors of marrow-derived cells. Finally, the observation that certain tumors from hormone-responsive tissues, e.g., breast tumors, could shrink after oophorectomy led to the idea that endogenous substances promoting the growth of a tumor might be antagonized. Chemicals achieving each of the goals are actually or intellectually the forerunners of the currently used cancer chemotherapy agents.

Systemic cancer treatments are of four broad types. *Conventional “cytotoxic” chemotherapy agents* were historically derived by the empirical observation that these “small molecules” (generally with molecular mass <1500 Da) could cause major regression of experimental tumors growing in animals. These agents mainly target DNA structure or segregation of DNA as chromosomes in mitosis. *Targeted agents* refer to small molecules or “biologics” (generally macromolecules such as antibodies or cytokines) designed and developed to interact with a defined molecular target important in maintaining the malignant state or expressed by the tumor cells. As described in [Chap. 102e](#), successful tumors have activated biochemical pathways that lead to uncontrolled proliferation through the action of, e.g., oncogene products, loss of cell cycle inhibitors, or loss of cell death regulation, and have acquired the capacity to replicate chromosomes indefinitely, invade, metastasize, and evade the immune system. Targeted therapies seek to capitalize on the biology behind the aberrant cellular behavior as a basis for therapeutic effects. *Hormonal therapies* (the first form of targeted therapy) capitalize on the biochemical pathways underlying estrogen and androgen function and action as a therapeutic basis for approaching patients with tumors of breast, prostate, uterus, and ovarian origin. *Biologic therapies* are often macromolecules that have a particular target (e.g., antiproliferation factor

or cytokine antibodies) or may have the capacity to regulate growth of tumor cells or induce a host immune response to kill tumor cells. Thus, biologic therapies include not only antibodies but also cytokines and gene therapies.

CANCER CHEMOTHERAPY

Principles The usefulness of any drug is governed by the extent to which a given dose causes a useful result (therapeutic effect; in the case of anticancer agents, toxicity to tumor cells) as opposed to a toxic effect to the host. The *therapeutic index* is the degree of separation between toxic and therapeutic doses. Really useful drugs have large therapeutic indices, and this usually occurs when the drug target is expressed in the disease-causing compartment as opposed to the normal compartment. Classically, selective toxicity of an agent for a tissue or cell type is governed by the differential expression of a drug’s target in the “sensitive” cell type or by differential drug accumulation into or elimination from compartments where greater or lesser toxicity is experienced, respectively. Currently used chemotherapeutic agents have the unfortunate property that their targets are present in both normal and tumor tissues. Therefore, they have relatively narrow therapeutic indices.

[Figure 103e-2](#) illustrates steps in cancer drug development. Following demonstration of antitumor activity in animal models, potentially useful anticancer agents are further evaluated to define an optimal schedule of administration and arrive at a drug formulation designed for a given route of administration and schedule. Safety testing in two species on an analogous schedule of administration defines the starting dose for a phase 1 trial in humans, usually but not always in patients with cancer who have exhausted “standard” (already approved) treatments. The initial dose is usually one-sixth to one-tenth of the dose just causing easily reversible toxicity in the more sensitive animal species. Escalating doses of the drug are then given during the human phase 1 trial until reversible toxicity is observed. Dose-limiting toxicity (DLT) defines a dose that conveys greater toxicity than would be acceptable in routine practice, allowing definition of a lower maximum-tolerated dose (MTD). The occurrence of toxicity is, if possible, correlated with plasma drug concentrations. The MTD or a dose just lower than the MTD is usually the dose suitable for phase 2 trials, where a fixed dose is administered to a relatively homogeneous set of patients with a particular tumor type in an effort to define whether the drug causes regression of tumors. In a phase 3 trial, evidence of improved overall survival or improvement in the time to progression of disease on the part of the new drug is sought in comparison to an appropriate control population, which is usually receiving an acceptable “standard of care” approach. A favorable outcome of a phase 3 trial is the basis for application to a regulatory agency for approval of the new agent for commercial marketing as safe and possessing a measure of clinical effectiveness.

Response, defined as tumor shrinkage, is the most immediate indicator of drug effect. To be clinically valuable, responses must translate into clinical benefit. This is conventionally established by a beneficial effect on overall survival, or at least an increased time to further progression of disease. Karnofsky was among the first to champion the evaluation of a chemotherapeutic agent’s benefit by carefully quantitating its effect on tumor size and using these measurements to objectively decide the basis for further treatment of a particular patient or further clinical evaluation of a drug’s potential. A partial response (PR) is defined conventionally as a decrease by at least 50% in a tumor’s bidimensional area; a complete response (CR) connotes disappearance of all tumor; progression of disease signifies an increase in size of existing lesions by >25% from baseline or best response or development of new lesions; and stable disease fits into none of the above categories. Newer evaluation systems, such as Response Evaluation Criteria in Solid Tumors (RECIST), use unidimensional measurement, but the intent is similar in rigorously defining evidence for the activity of the agent in assessing its value to the patient. An active chemotherapy agent conventionally has PR rates of at least 20–25% with reversible non-life-threatening side effects, and it may then be suitable for study in phase 3 trials to assess efficacy in comparison to standard or no therapy. Active efforts are being made to quantitate effects of anticancer

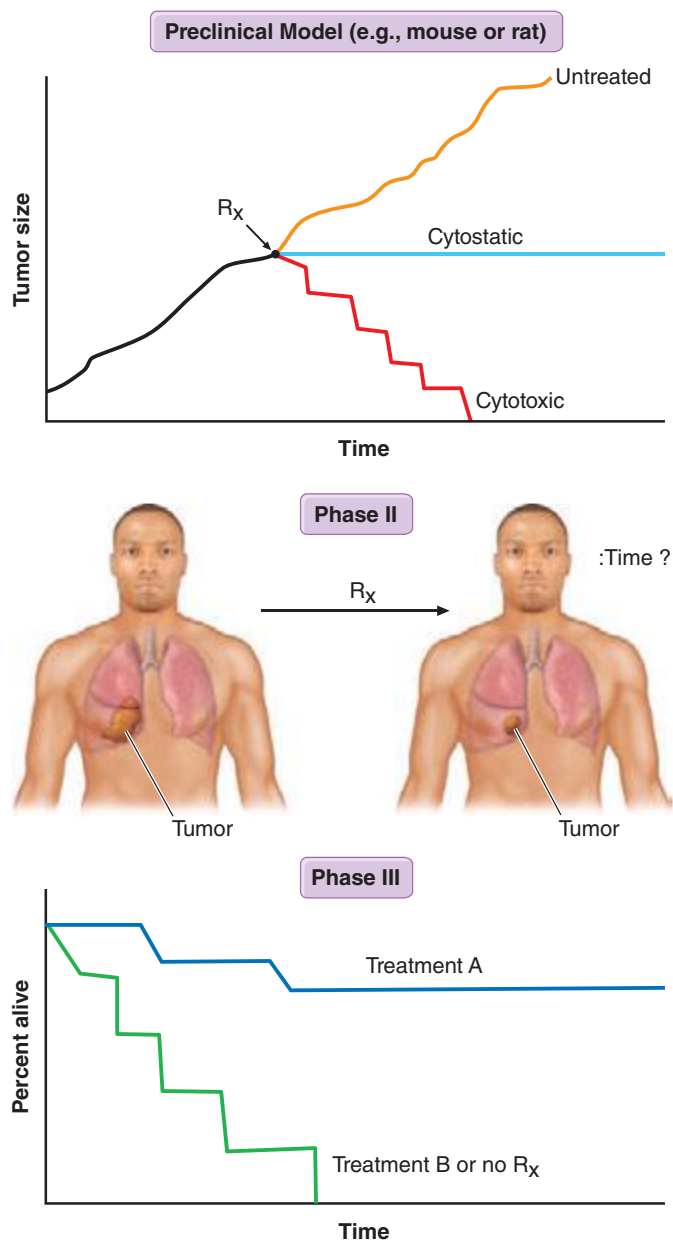


FIGURE 103e-2 Steps in cancer drug discovery and development.

Preclinical activity (*top*) in animal models of cancers may be used as evidence to support the entry of the drug candidate into phase 1 trials in humans to define a correct dose and observe any clinical antitumor effect that may occur. The drug may then be advanced to phase 2 trials directed against specific cancer types, with rigorous quantitation of antitumor effects (*middle*). Phase 3 trials then may reveal activity superior to standard or no treatment (*bottom*).

agents on quality of life. Cancer drug clinical trials conventionally use a toxicity grading scale where grade 1 toxicities do not require treatment, grade 2 toxicities may require symptomatic treatment but are not life-threatening, grade 3 toxicities are potentially life-threatening if untreated, grade 4 toxicities are actually life-threatening, and grade 5 toxicities are those that result in the patient's death.

Development of targeted agents may proceed quite differently. While phase 1–3 trials are still conducted, molecular analysis of human tumors may allow the precise definition of target expression in a patient's tumor that is necessary for or relevant to the drug's action. This information might then allow selection of patients expressing the drug target for participation in all trial phases. These patients may then have a greater chance of developing a useful response to the drug by virtue of expressing the target in the tumor. Clinical trials may be

designed to incorporate an assessment of the behavior of the target in relation to the drug (pharmacodynamic studies). Ideally, the plasma concentration that affects the drug target is known, so escalation to MTD may not be necessary. Rather, the correlation of host toxicity while achieving an “optimal biologic dose” becomes a more relevant endpoint for phase 1 and early phase 2 trials with targeted agents.

Useful cancer drug treatment strategies using conventional chemotherapy agents, targeted agents, hormonal treatments, or biologics have one of two valuable outcomes. They can induce cancer cell death, resulting in tumor shrinkage with corresponding improvement in patient survival, or increase the time until the disease progresses. Another potential outcome is to induce cancer cell *differentiation* or *dormancy* with loss of tumor cell replicative potential and reacquisition of phenotypic properties resembling normal cells. A blocking in normal cellular differentiation may be a key feature in the pathogenesis of certain leukemias.

Cell death is a closely regulated process. *Necrosis* refers to cell death induced, for example, by physical damage with the hallmarks of cell swelling and membrane disruption. *Apoptosis*, or programmed cell death, refers to a highly ordered process whereby cells respond to defined stimuli by dying, and it recapitulates the necessary cell death observed during the ontogeny of the organism. Cancer chemotherapeutic agents can cause both necrosis and apoptosis. Apoptosis is characterized by chromatin condensation (giving rise to “apoptotic bodies”), cell shrinkage, and, in living animals, phagocytosis by surrounding stromal cells without evidence of inflammation. This process is regulated either by signal transduction systems that promote a cell's demise after a certain level of insult is achieved or in response to specific cell-surface receptors that mediate physiologic cell death responses, such as occurs in the developing organism or in the normal function of immune cells. Influencing apoptosis by manipulation of signal transduction pathways has emerged as a basis for understanding the actions of drugs and designing new strategies to improve their use. *Autophagy* is a cellular response to injury where the cell does not initially die but catabolizes itself in a way that can lead to loss of replicative potential. A general view of how cancer treatments work is that the interaction of a chemotherapeutic drug with its target induces a “cascade” of further signaling steps. These signals ultimately lead to cell death by triggering an “execution phase” where proteases, nucleases, and endogenous regulators of the cell death pathway are activated ([Fig. 103e-3](#)).

Targeted agents differ from chemotherapy agents in that they do not indiscriminately cause macromolecular lesions but regulate the action of particular pathways. For example, the p210^{bcr-abl} fusion protein tyrosine kinase drives chronic myeloid leukemia (CML), and HER2/neu stimulates the proliferation of certain breast cancers. The tumor has been described as “addicted” to the function of these molecules in the sense that without the pathway's continued action, the tumor cell cannot survive. In this way, targeted agents directed at p210^{bcr-abl} or HER2/neu may alter the “threshold” tumors driven by these molecules may have for undergoing apoptosis without actually creating any molecular lesions such as direct DNA strand breakage or altered membrane function.

While apoptotic mechanisms are important in regulating cellular proliferation and the behavior of tumor cells *in vitro*, *in vivo* it is unclear whether all of the actions of chemotherapeutic agents to cause cell death can be attributed to apoptotic mechanisms. However, changes in molecules that regulate apoptosis are correlated with clinical outcomes (e.g., *bcl2* overexpression in certain lymphomas conveys poor prognosis; proapoptotic *bax* expression is associated with a better outcome after chemotherapy for ovarian carcinoma). A better understanding of the relationship of cell death and cell survival mechanisms is needed.

Chemotherapy agents may be used for the treatment of active, clinically apparent cancer. The goal of such treatment in some cases is cure of the cancer, that is, elimination of all clinical and pathologic evidence of cancer and return of the patient to an expected survival no different than the general population. [Table 103e-3, A](#) lists those tumors considered curable by conventionally available chemotherapeutic agents

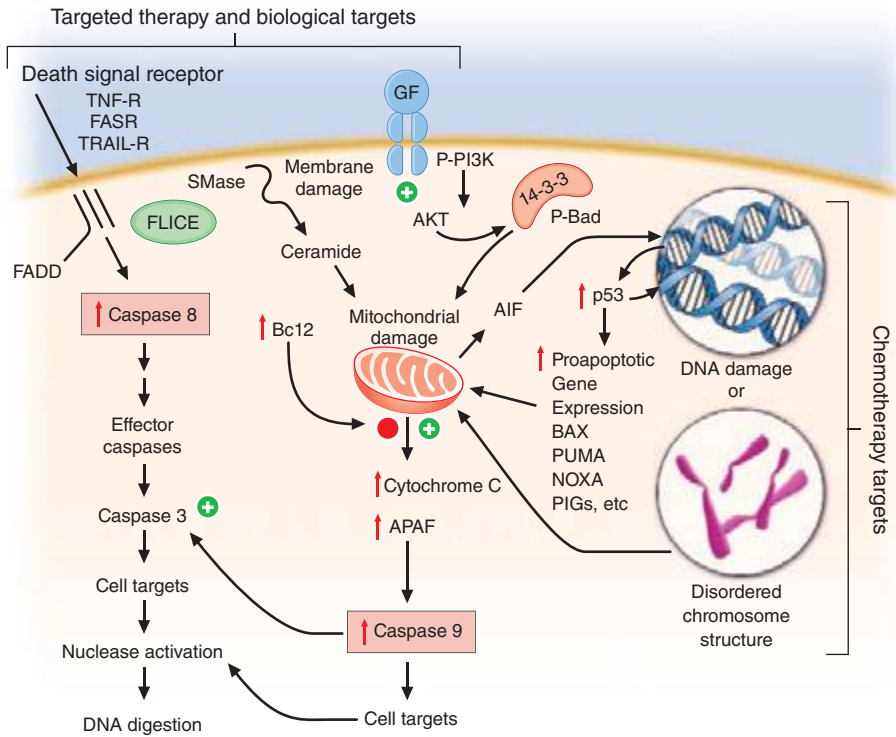


FIGURE 103e-3 Integration of cell death responses. Cell death through an apoptotic mechanism requires active participation of the cell. In response to interruption of growth factor (GF) or propagation of certain cytokine death signals (e.g., tumor necrosis factor receptor [TNF-R]), there is activation of “upstream” cysteine aspartyl proteases (caspases), which then directly digest cytoplasmic and nuclear proteins, resulting in activation of “downstream” caspases; these cause activation of nucleases, resulting in the characteristic DNA fragmentation that is a hallmark of apoptosis. Chemotherapy agents that create lesions in DNA or alter mitotic spindle function seem to activate aspects of this process by damage ultimately conveyed to the mitochondria, perhaps by activating the transcription of genes whose products can produce or modulate the toxicity of free radicals. In addition, membrane damage with activation of sphingomyelinases results in the production of ceramides that can have a direct action at mitochondria. The anti-apoptotic protein bcl2 attenuates mitochondrial toxicity, while proapoptotic gene products such as bax antagonize the action of bcl2. Damaged mitochondria release cytochrome C and apoptosis-activating factor (APAF), which can directly activate caspase 9, resulting in propagation of a direct signal to other downstream caspases through protease activation. Apoptosis-inducing factor (AIF) is also released from the mitochondrion and then can translocate to the nucleus, bind to DNA, and generate free radicals to further damage DNA. An additional proapoptotic stimulus is the bad protein, which can heterodimerize with bcl2 gene family members to antagonize apoptosis. Importantly, though, bad protein function can be retarded by its sequestration as phospho-bad through the 14-3-3 adapter proteins. The phosphorylation of bad is mediated by the action of the AKT kinase in a way that defines how growth factors that activate this kinase can retard apoptosis and promote cell survival.

when used to address disseminated or metastatic cancers. If a tumor is localized to a single site, serious consideration of surgery or primary radiation therapy should be given, because these treatment modalities may be curative as local treatments. Chemotherapy may then be used after the failure of these modalities to eradicate a local tumor or as part of multimodality approaches to offer primary treatment to a clinically localized tumor. In this event, it can allow organ preservation when given with radiation, as in the larynx or other upper airway sites, or sensitize tumors to radiation when given, e.g., to patients concurrently receiving radiation for lung or cervix cancer (Table 103e-3, B). Chemotherapy can be administered as an *adjuvant*, i.e., in addition to surgery or radiation (Table 103e-3, C), even after all clinically apparent disease has been removed. This use of chemotherapy has curative potential in breast and colorectal neoplasms, as it attempts to eliminate clinically unapparent tumor that may have already disseminated. As noted above, small tumors frequently have high growth fractions and therefore may be intrinsically more susceptible to the action of antiproliferative agents. *Neoadjuvant* chemotherapy refers to administration of

chemotherapy prior to any surgery or radiation to a local tumor in an effort to enhance the effect of the local treatment.

Chemotherapy is routinely used in “conventional” dose regimens. In general, these doses produce reversible acute side effects, primarily consisting of transient myelosuppression with or without gastrointestinal toxicity (usually nausea), which are readily managed. “High-dose” chemotherapy regimens are predicated on the observation that the dose-response curve for many anticancer agents is rather steep, and increased dose can produce markedly increased therapeutic effect, although at the cost of potentially life-threatening complications that require intensive support, usually in the form of hematopoietic stem cell support from the patient (*autologous*) or from donors matched for histocompatibility loci (*allogeneic*), or pharmacologic “rescue” strategies to repair the effect of the high-dose chemotherapy on normal tissues. High-dose regimens have definite curative potential in defined clinical settings (Table 103e-3, D).

If cure is not possible, chemotherapy may be undertaken with the goal of palliating some aspect of the tumor’s effect on the host. In this usage, value is perceived by the demonstration of improved symptom relief, progression-free survival, or overall survival at a certain time from the inception of treatment in the treated population, compared to a relevant control population established as the result of clinical research protocol or other organized comparative study. Such clinical research protocols are the basis for U.S. Food and Drug Administration (FDA) approval of a particular cancer treatment as safe and effective and are the benchmark for an evidence-based approach to the use of chemotherapeutic agents. Common tumors that may be meaningfully addressed by chemotherapy with palliative intent are listed in Table 103e-3, E.

Usually, tumor-related symptoms manifest as pain, weight loss, or some local symptom related to the tumor’s effect on normal structures. Patients treated with palliative intent should be aware of their diagnosis and the limitations of the proposed treatments, have access to supportive care, and have suitable “performance status,” according to assessment algorithms such as the one developed by Karnofsky (see Table 99-4) or by the Eastern Cooperative Oncology Group (ECOG) (see Table 99-5). ECOG performance status 0 (PS0) patients are without symptoms; PS1 patients are ambulatory but restricted in strenuous physical activity; PS2 patients are ambulatory but unable to work and are up and about 50% or more of the time; PS3 patients are capable of limited self-care and are up <50% of the time; and PS4 patients are totally confined to bed or chair and incapable of self-care. Only PS0, PS1, and PS2 patients are generally considered suitable for palliative (noncurative) treatment. If there is curative potential, even poor-performance status patients may be treated, but their prognosis is usually inferior to that of good-performance status patients treated with similar regimens.

An important perspective the primary care provider may bring to patients and their families facing incurable cancer is that, given the limited value of chemotherapeutic approaches at some point in the

TABLE 103e-3 CURABILITY OF CANCERS WITH CHEMOTHERAPY

<p>A. Advanced Cancers with Possible Cure</p> <p>Acute lymphoid and acute myeloid leukemia (pediatric/adult)</p> <p>Hodgkin's disease (pediatric/adult)</p> <p>Lymphomas—certain types (pediatric/adult)</p> <p>Germ cell neoplasms</p> <ul style="list-style-type: none"> Embryonal carcinoma Teratocarcinoma Seminoma or dysgerminoma Choriocarcinoma <p>Gestational trophoblastic neoplasia</p> <p>Pediatric neoplasms</p> <ul style="list-style-type: none"> Wilms' tumor Embryonal rhabdomyosarcoma Ewing's sarcoma Peripheral neuroepithelioma Neuroblastoma <p>Small-cell lung carcinoma</p> <p>Ovarian carcinoma</p> <p>B. Advanced Cancers Possibly Cured by Chemotherapy and Radiation</p> <p>Squamous carcinoma (head and neck)</p> <p>Squamous carcinoma (anus)</p> <p>Breast carcinoma</p> <p>Carcinoma of the uterine cervix</p> <p>Non-small-cell lung carcinoma (stage III)</p> <p>Small-cell lung carcinoma</p> <p>C. Cancers Possibly Cured with Chemotherapy as Adjuvant to Surgery</p> <p>Breast carcinoma</p> <p>Colorectal carcinoma^a</p> <p>Osteogenic sarcoma</p> <p>Soft tissue sarcoma</p>	<p>D. Cancers Possibly Cured with "High-Dose" Chemotherapy with Stem Cell Support</p> <p>Relapsed leukemias, lymphoid and myeloid</p> <p>Relapsed lymphomas, Hodgkin's and non-Hodgkin's</p> <p>Chronic myeloid leukemia</p> <p>Multiple myeloma</p> <p>E. Cancers Responsive with Useful Palliation, But Not Cure, by Chemotherapy</p> <p>Bladder carcinoma</p> <p>Chronic myeloid leukemia</p> <p>Hairy cell leukemia</p> <p>Chronic lymphocytic leukemia</p> <p>Lymphoma—certain types</p> <p>Multiple myeloma</p> <p>Gastric carcinoma</p> <p>Cervix carcinoma</p> <p>Endometrial carcinoma</p> <p>Soft tissue sarcoma</p> <p>Head and neck cancer</p> <p>Adrenocortical carcinoma</p> <p>Islet cell neoplasms</p> <p>Breast carcinoma</p> <p>Colorectal carcinoma</p> <p>Renal carcinoma</p> <p>F. Tumors Poorly Responsive in Advanced Stages to Chemotherapy</p> <p>Pancreatic carcinoma</p> <p>Biliary tract neoplasms</p> <p>Thyroid carcinoma</p> <p>Carcinoma of the vulva</p> <p>Non-small-cell lung carcinoma</p> <p>Prostate carcinoma</p> <p>Melanoma (subsets)</p> <p>Hepatocellular carcinoma</p> <p>Salivary gland cancer</p>
---	--

^aRectum also receives radiation therapy.

natural history of most metastatic cancers, *palliative care* or *hospice-based* approaches, with meticulous and ongoing attention to symptom relief and with family, psychological, and spiritual support, should receive prominent attention as a valuable therapeutic plan (**Chaps. 10 and 99**). Optimizing the quality of life rather than attempting to extend it becomes a valued intervention. Patients facing the impending progression of disease in a life-threatening way frequently choose to undertake toxic treatments of little to no potential value, and support provided by the primary caregiver in accessing palliative and hospice-based options in contrast to receiving toxic and ineffective regimen can be critical in providing a basis for patients to make sensible choices.

Cytotoxic Chemotherapy Agents Table 103e-4 lists commonly used cytotoxic cancer chemotherapy agents and pertinent clinical aspects of their use, with particular reference to adverse effects that might be encountered by the generalist in the care of patients. The drugs listed may be usefully grouped into two general categories: those affecting DNA and those affecting microtubules.

DIRECT DNA-INTERACTIVE AGENTS DNA replication occurs during the synthesis or S-phase of the cell cycle, with chromosome segregation of

the replicated DNA occurring in the M, or mitosis, phase. The G₁ and G₂ "gap phases" precede S and M, respectively. Historically, chemotherapeutic agents have been divided into "phase-nonspecific" agents, which can act in any phase of the cell cycle, and "phase-specific" agents, which require the cell to be at a particular cell cycle phase to cause greatest effect. Once the agent has acted, cells may progress to "checkpoints" in the cell cycle where the drug-related damage may be assessed and either repaired or allowed to initiate apoptosis. An important function of certain tumor-suppressor genes such as *p53* may be to modulate checkpoint function.

Alkylating agents as a class are cell cycle phase-nonspecific agents. They break down, either spontaneously or after normal organ or tumor cell metabolism, to reactive intermediates that covalently modify bases in DNA. This leads to cross-linkage of DNA strands or the appearance of breaks in DNA as a result of repair efforts. "Broken" or cross-linked DNA is intrinsically unable to complete normal replication or cell division; in addition, it is a potent activator of cell cycle checkpoints and further activates cell-signaling pathways that can precipitate apoptosis. As a class, alkylating agents share similar toxicities: myelosuppression, alopecia, gonadal dysfunction, mucositis, and pulmonary fibrosis. They differ greatly in a spectrum of normal organ toxicities. As a class, they share the capacity to cause "second" neoplasms, particularly leukemia, many years after use, particularly when used in low doses for protracted periods.

Cyclophosphamide is inactive unless metabolized by the liver to 4-hydroxy-cyclophosphamide, which decomposes into an alkylating species, as well as to chloroacetaldehyde and acrolein. The latter causes chemical cystitis; therefore, excellent hydration must be maintained while using cyclophosphamide. If severe, the cystitis may be prevented from progressing or prevented altogether (if expected from the dose of cyclophosphamide to be used) by mesna (2-mercaptoethanesulfonate). Liver disease impairs cyclophosphamide activation. Sporadic interstitial pneumonitis leading to pulmonary fibrosis can accompany the use of cyclophosphamide, and high doses used in conditioning regimens for bone marrow transplant can cause cardiac dysfunction. Ifosfamide is a cyclophosphamide analogue also activated in the liver, but more slowly, and it requires coadministration of mesna to prevent bladder injury. Central nervous system (CNS) effects, including somnolence, confusion, and psychosis, can follow ifosfamide use; the incidence appears related to low body surface area or decreased creatinine clearance.

Several alkylating agents are less commonly used. Nitrogen mustard (mechlorethamine) is the prototypic agent of this class, decomposing rapidly in aqueous solution to potentially yield a bifunctional carbonium ion. It must be administered shortly after preparation into a rapidly flowing intravenous line. It is a powerful vesicant, and infiltration may be symptomatically ameliorated by infiltration of the affected site with 1/6 M thiosulfate. Even without infiltration, aseptic thrombophlebitis is frequent. It can be used topically as a dilute solution or ointment in cutaneous lymphomas, with a notable incidence of hypersensitivity reactions. It causes moderate nausea after intravenous administration. Bendamustine is a nitrogen mustard derivative with evidence of activity in chronic lymphocytic leukemia and certain lymphomas.

Chlorambucil causes predictable myelosuppression, azoospermia, nausea, and pulmonary side effects. Busulfan can cause profound myelosuppression, alopecia, and pulmonary toxicity but is relatively "lymphocyte sparing." Its routine use in treatment of CML has been curtailed in favor of imatinib (Gleevec) or dasatinib, but it is still used in transplant preparation regimens. Melphalan shows variable oral bioavailability and undergoes extensive binding to albumin and α_2 -acidic glycoprotein. Mucositis appears more prominently; however, it has prominent activity in multiple myeloma.

Nitrosoureas break down to carbamylating species that not only cause a distinct pattern of DNA base pair-directed toxicity but also can covalently modify proteins. They share the feature of causing relatively delayed bone marrow toxicity, which can be cumulative and long-lasting. Procarbazine is metabolized in the liver and possibly in tumor cells to yield a variety of free radical and alkylating species. In addition to

TABLE 103e-4 CYTOTOXIC CHEMOTHERAPY AGENTS

Drug	Toxicity	Interactions, Issues
Direct DNA-Interacting Agents		
Alkylator		
Cyclophosphamide	Marrow (relative platelet sparing) Cystitis Common alkylator ^a Cardiac (high dose)	Liver metabolism required to activate to phosphoramidate mustard + acrolein Mesna protects against "high-dose" bladder damage
Mechlorethamine	Marrow Vesicant Nausea	Topical use in cutaneous lymphoma
Chlorambucil	Marrow Common alkylator ^a	
Melphalan	Marrow (delayed nadir) GI (high dose)	Decreased renal function delays clearance
Carmustine (BCNU)	Marrow (delayed nadir) GI, liver (high dose) Renal	
Lomustine (CCNU)	Marrow (delayed nadir)	
Ifosfamide	Myelosuppressive Bladder Neurologic Metabolic acidosis	Analogue of cyclophosphamide Must use mesna Greater activity vs testicular neoplasms and sarcomas
Procarbazine	Marrow Nausea Neurologic Common alkylator ^a	Liver and tissue metabolism required Disulfiram-like effect with ethanol Acts as MAOI HBP after tyrosinase-rich foods
Dacarbazine (DTIC)	Marrow Nausea Flulike	Metabolic activation
Temozolomide	Nausea/vomiting Headache/fatigue Constipation	Infrequent myelosuppression
Altretamine (formerly hexamethyl-melamine)	Nausea Neurologic (mood swing) Neuropathy Marrow (less)	Liver activation Barbiturates enhance/cimetidine diminishes
Cisplatin	Nausea Neuropathy Auditory Marrow platelets > WBCs Renal Mg ²⁺ , Ca ²⁺	Maintain high urine flow; osmotic diuresis, monitor intake/output K ⁺ , Mg ²⁺ Emetogenic—prophylaxis needed Full dose if CrCl >60 mL/min and tolerate fluid push
Carboplatin	Marrow platelets > WBCs Nausea Renal (high dose)	Reduce dose according to CrCl: to AUC of 5–7 mg/mL per min [AUC = dose/(CrCl + 25)]
Oxaliplatin	Nausea Anemia	Acute reversible neurotoxicity; chronic sensory neurotoxicity cumulative with dose; reversible laryngopharyngeal spasm
Antitumor Antibiotics and Topoisomerase Poisons		
Bleomycin	Pulmonary Skin effects Raynaud's Hypersensitivity	Inactivate by bleomycin hydrolase (decreased in lung/skin) O ₂ enhances pulmonary toxicity Cisplatin-induced decrease in CrCl may increase skin/lung toxicity Reduce dose if CrCl <60 mL/min
Dactinomycin	Marrow Nausea Mucositis Vesicant Alopecia	Radiation recall

(Continued)

TABLE 103e-4 CYTOTOXIC CHEMOTHERAPY AGENTS (CONTINUED)

Drug	Toxicity	Interactions, Issues
Etoposide (VP16-213)	Marrow (WBCs > platelet) Alopecia Hypotension Hypersensitivity (rapid IV) Nausea Mucositis (high dose)	Hepatic metabolism—renal 30% Reduce doses with renal failure Schedule-dependent (5-day schedule better than 1-day) Late leukemogenic Accentuate antimetabolite action
Topotecan	Marrow Mucositis Nausea Mild alopecia	Reduce dose with renal failure No liver toxicity
Irinotecan	Diarrhea: "early onset" with cramping, flushing, vomiting; "late onset" after several doses Marrow Alopecia Nausea Vomiting Pulmonary	Prodrug requires enzymatic clearance to active drug "SN 38" Early diarrhea likely due to biliary excretion Late diarrhea, use "high-dose" loperamide (2 mg q2–4 h)
Doxorubicin and daunorubicin	Marrow Mucositis Alopecia Cardiovascular acute/chronic Vesicant	Heparin aggregate; coadministration increases clearance Acetaminophen, BCNU increase liver toxicity Radiation recall
Idarubicin	Marrow Cardiac (less than doxorubicin)	None established
Epirubicin	Marrow Cardiac	None established
Mitoxantrone	Marrow Cardiac (less than doxorubicin) Vesicant (mild) Blue urine, sclerae, nails	Interacts with heparin Less alopecia, nausea than doxorubicin Radiation recall Less alopecia, nausea than doxorubicin
Indirectly DNA-Interacting Agents		
Antimetabolites		
Deoxycoformycin	Nausea Immunosuppression Neurologic Renal	Excretes in urine Reduce dose for renal failure Inhibits adenosine deaminase Reduce dose for renal failure
6-Mercaptopurine (6-MP)	Marrow Liver Nausea	Variable bioavailability Metabolize by xanthine oxidase Decrease dose with allopurinol Increased toxicity with thiopurine methyltransferase deficiency
6-Thioguanine	Marrow Liver Nausea	Variable bioavailability Increased toxicity with thiopurine methyltransferase deficiency
Azathioprine	Marrow Nausea Liver	Metabolizes to 6-MP; therefore, reduce dose with allopurinol Increase toxicity with thiopurine methyltransferase deficiency
2-Chlorodeoxyadenosine	Marrow Renal Fever	Notable use in hairy cell leukemia
Hydroxyurea	Marrow Nausea Mucositis Skin changes Rare renal, liver, lung, CNS	Decrease dose with renal failure Augments antimetabolite effect

(Continued)

TABLE 103e-4 CYTOTOXIC CHEMOTHERAPY AGENTS (CONTINUED)

Drug	Toxicity	Interactions, Issues
Methotrexate	Marrow Liver/lung Renal tubular Mucositis	Toxicity lessened by “rescue” with leucovorin Excreted in urine Decrease dose in renal failure; NSAIDs increase renal toxicity
5-Fluorouracil (5FU)	Marrow Mucositis Neurologic Skin changes	Toxicity enhanced by leucovorin by increasing “ternary complex” with thymidylate synthase; dihydropyrimidine dehydrogenase deficiency increases toxicity; metabolism in tissue
Capecitabine	Diarrhea Hand-foot syndrome	Prodrug of 5FU due to intratumoral metabolism
Cytosine arabinoside	Marrow Mucositis Neurologic (high dose) Conjunctivitis (high dose) Noncardiogenic pulmonary edema	Enhances activity of alkylating agents Metabolizes in tissues by deamination but renal excretion prominent at doses >500 mg; therefore, dose reduce in “high-dose” regimens in patients with decreased CrCl
Azacitidine	Marrow Nausea Liver Neurologic Myalgia	Use limited to leukemia Altered methylation of DNA alters gene expression
Gemcitabine	Marrow Nausea Hepatic Fever/“flu syndrome”	
Fludarabine phosphate	Marrow Neurologic Lung	Dose reduction with renal failure Metabolized to F-ara converted to F-ara ATP in cells by deoxycytidine kinase
Clofarabine	Myelosuppression Mucositis Rare cardiac/inflammatory	
Nelarabine	Myelosuppression Neurologic	T cell ALL; T cell lymphoblastic lymphoma
Asparaginase	Protein synthesis; indirect inhibition of DNA synthesis by decreased histone synthesis Clotting factors Glucose Albumin Hypersensitivity CNS Pancreatitis Hepatic	Blocks methotrexate action
Pemetrexed	Anemia Neutropenia Thrombocytopenia	Supplement folate/B ₁₂ Caution in renal failure
Pralatrexate	Myelosuppression Mucositis	Active in peripheral T cell lymphoma
Antimitotic Agents		
Vincristine	Vesicant Marrow Neurologic GI: ileus/constipation; bladder hypotoxicity; SIADH Cardiovascular	Hepatic clearance Dose reduction for bilirubin >1.5 mg/dL Prophylactic bowel regimen

(Continued)

TABLE 103e-4 CYTOTOXIC CHEMOTHERAPY AGENTS (CONTINUED)

Drug	Toxicity	Interactions, Issues
Vinblastine	Vesicant Marrow Neurologic (less common but similar spectrum to other vincas) Hypertension Raynaud's	Hepatic clearance Dose reduction as with vincristine
Vinorelbine	Vesicant Marrow Allergic/bronchospasm (immediate) Dyspnea/cough (subacute) Neurologic (less prominent but similar spectrum to other vincas)	Hepatic clearance
Paclitaxel	Hypersensitivity Marrow Mucositis Alopecia Sensory neuropathy CV conduction disturbance Nausea—infrequent	Premedicate with steroids, H ₁ and H ₂ blockers Hepatic clearance Dose reduction as with vincas
Docetaxel	Hypersensitivity Fluid retention syndrome Marrow Dermatologic Sensory neuropathy Nausea infrequent Some stomatitis	Premedicate with steroids, H ₁ and H ₂ blockers
Estramustine phosphate	Nausea Vomiting Diarrhea CHF Thrombosis Gynecomastia	
Nab-paclitaxel (protein bound)	Neuropathy Anemia Neutropenia Thrombocytopenia	Caution in hepatic insufficiency
Ixabepilone	Myelosuppression Neuropathy	

*Common alkylator: alopecia, pulmonary, infertility, plus teratogenesis.

Abbreviations: ALL, acute lymphocytic leukemia; AUC, area under the curve; CHF, congestive heart failure; CNS, central nervous system; CrCl, creatinine clearance; CV, cardiovascular; GI, gastrointestinal; HBP, high blood pressure; MAOI, monoamine oxidase inhibitor; NSAID, nonsteroidal anti-inflammatory drug; SIADH, syndrome of inappropriate antidiuretic hormone secretion.

myelosuppression, it causes hypnotic and other CNS effects, including vivid nightmares. It can cause a disulfiram-like syndrome on ingestion of ethanol. Altretamine (formerly hexa-methylmelamine) and thiotepa can chemically give rise to alkylating species, although the nature of the DNA damage has not been well characterized in either case. Dacarbazine (DTIC) is activated in the liver to yield the highly reactive methyl diazonium cation. It causes only modest myelosuppression 21–25 days after a dose but causes prominent nausea on day 1. Temozolomide is structurally related to dacarbazine but was designed to be activated by nonenzymatic hydrolysis in tumors and is bioavailable orally.

Cisplatin was discovered fortuitously by observing that bacteria present in electrolysis solutions with platinum electrodes could not divide. Only the *cis* diamine configuration is active as an antitumor agent. It is hypothesized that in the intracellular environment, a chloride is lost from each position, being replaced by a water molecule. The resulting positively charged species is an efficient bifunctional interactor with DNA, forming Pt-based cross-links.

Cisplatin requires administration with adequate hydration, including forced diuresis with mannitol to prevent kidney damage; even with the use of hydration, gradual decrease in kidney function is common, along with noteworthy anemia. Hypomagnesemia frequently attends cisplatin use and can lead to hypocalcemia and tetany. Other common toxicities include neurotoxicity with stocking-and-glove sensorimotor neuropathy. Hearing loss occurs in 50% of patients treated with conventional doses. Cisplatin is intensely emetogenic, requiring prophylactic antiemetics. Myelosuppression is less evident than with other alkylating agents. Chronic vascular toxicity (Raynaud's phenomenon, coronary artery disease) is a more unusual toxicity. Carboplatin displays less nephro-, oto-, and neurotoxicity. However, myelosuppression is more frequent, and because the drug is exclusively cleared through the kidney, adjustment of dose for creatinine clearance must be accomplished through use of various dosing nomograms. Oxaliplatin is a platinum analogue with noteworthy activity in colon cancers refractory to other treatments. It is prominently neurotoxic.

ANTITUMOR ANTIBIOTICS AND TOPOISOMERASE POISONS Antitumor antibiotics are substances produced by bacteria that in nature appear to provide a chemical defense against other hostile microorganisms. As a class, they bind to DNA directly and can frequently undergo electron transfer reactions to generate free radicals in close proximity to DNA, leading to DNA damage in the form of single-strand breaks or cross-links. Topoisomerase poisons include natural products or semisynthetic species derived ultimately from plants, and they modify enzymes that regulate the capacity of DNA to unwind to allow normal replication or transcription. These include topoisomerase I, which creates single-strand breaks that then rejoin following the passage of the other DNA strand through the break. Topoisomerase II creates double-strand breaks through which another segment of DNA duplex passes before rejoining. DNA damage from these agents can occur in any cell cycle phase, but cells tend to arrest in S-phase or G₂ of the cell cycle in cells with p53 and Rb pathway lesions as the result of defective checkpoint mechanisms in cancer cells. Owing to the role of topoisomerase I in the procession of the replication fork, topoisomerase I poisons cause lethality if the topoisomerase I-induced lesions are made in S-phase.

Doxorubicin can intercalate into DNA, thereby altering DNA structure, replication, and topoisomerase II function. It can also undergo reduction reactions by accepting electrons into its quinone ring system, with the capacity to undergo reoxidation to form reactive oxygen radicals after reoxidation. It causes predictable myelosuppression, alopecia, nausea, and mucositis. In addition, it causes acute cardiotoxicity in the form of atrial and ventricular dysrhythmias, but these are rarely of clinical significance. In contrast, cumulative doses >550 mg/m² are associated with a 10% incidence of chronic cardiomyopathy. The incidence of cardiomyopathy appears to be related to schedule (peak serum concentration), with low-dose, frequent treatment or continuous infusions better tolerated than intermittent higher-dose exposures. Cardiotoxicity has been related to iron-catalyzed oxidation and reduction of doxorubicin, and not to topoisomerase action. Cardiotoxicity is related to peak plasma dose; thus, lower doses and continuous infusions are less likely to cause heart damage. Doxorubicin's cardiotoxicity is increased when given together with trastuzumab (Herceptin), the anti-HER2/neu antibody. Radiation recall or interaction with concomitantly administered radiation to cause local site complications is frequent. The drug is a powerful vesicant, with necrosis of tissue apparent 4–7 days after an extravasation; therefore, it should be administered into a rapidly flowing intravenous line. Dexrazoxane is an antidote to doxorubicin-induced extravasation. Doxorubicin is metabolized by the liver, so doses must be reduced by 50–75% in the presence of liver dysfunction. Daunorubicin is closely related to doxorubicin and was actually introduced first into leukemia treatment, where it remains part of curative regimens and has been shown preferable to doxorubicin owing to less mucositis and colonic damage. Idarubicin is also used in acute myeloid leukemia treatment and may be preferable to daunorubicin in activity. Encapsulation of daunorubicin into a liposomal formulation has attenuated cardiac toxicity and antitumor activity in Kaposi's sarcoma, other sarcomas, multiple myeloma, and ovarian cancer.

Bleomycin refers to a mixture of glycopeptides that have the unique feature of forming complexes with Fe²⁺ while also bound to DNA. It remains an important component of curative regimens for Hodgkin's disease and germ cell neoplasms. Oxidation of Fe²⁺ gives rise to superoxide and hydroxyl radicals. The drug causes little, if any, myelosuppression. The drug is cleared rapidly, but augmented skin and pulmonary toxicity in the presence of renal failure has led to the recommendation that doses be reduced by 50–75% in the face of a creatinine clearance <25 mL/min. Bleomycin is not a vesicant and can be administered intravenously, intramuscularly, or subcutaneously. Common side effects include fever and chills, facial flush, and Raynaud's phenomenon. Hypertension can follow rapid intravenous administration, and the incidence of anaphylaxis with early preparations of the drug has led to the practice of administering a test dose of 0.5–1 unit before the rest of the dose. The most feared complication of bleomycin treatment is pulmonary fibrosis, which increases in incidence at >300 cumulative units administered and is minimally responsive to treatment

(e.g., glucocorticoids). The earliest indicator of an adverse effect is usually a decline in the carbon monoxide diffusing capacity (DL_{CO}) or coughing, although cessation of drug immediately upon documentation of a decrease in DL_{CO} may not prevent further decline in pulmonary function. Bleomycin is inactivated by a bleomycin hydrolase, whose concentration is diminished in skin and lung. Because bleomycin-dependent electron transport is dependent on O₂, bleomycin toxicity may become apparent after exposure to transient very high fraction of inspired oxygen (FIO₂). Thus, during surgical procedures, patients with prior exposure to bleomycin should be maintained on the lowest FIO₂ consistent with maintaining adequate tissue oxygenation.

Mitoxantrone is a synthetic compound that was designed to recapitulate features of doxorubicin but with less cardiotoxicity. It is quantitatively less cardiotoxic (comparing the ratio of cardiotoxic to therapeutically effective doses) but is still associated with a 10% incidence of cardiotoxicity at cumulative doses of >150 mg/m². It also causes alopecia. Cases of acute promyelocytic leukemia (APL) have arisen shortly after exposure of patients to mitoxantrone, particularly in the adjuvant treatment of breast cancer. Although chemotherapy-associated leukemia is generally of the acute myeloid type, APL arising in the setting of prior mitoxantrone treatment had the typical t(15;17) chromosome translocation associated with APL, but the breakpoints of the translocation appeared to be at topoisomerase II sites that would be preferred sites of mitoxantrone action, clearly linking the action of the drug to the generation of the leukemia.

Etoposide was synthetically derived from the plant product podophyllotoxin; it binds directly to topoisomerase II and DNA in a reversible ternary complex. It stabilizes the covalent intermediate in the enzyme's action where the enzyme is covalently linked to DNA. This "alkali-labile" DNA bond was historically a first hint that an enzyme such as a topoisomerase might exist. The drug therefore causes a prominent G₂ arrest, reflecting the action of a DNA damage checkpoint. Prominent clinical effects include myelosuppression, nausea, and transient hypotension related to the speed of administration of the agent. Etoposide is a mild vesicant but is relatively free from other large-organ toxicities. When given at high doses or very frequently, topoisomerase II inhibitors may cause acute leukemia associated with chromosome 11q23 abnormalities in up to 1% of exposed patients.

Camptothecin was isolated from extracts of a Chinese tree and had notable antileukemia activity in preclinical mouse models. Early human clinical studies with the sodium salt of the hydrolyzed camptothecin lactone showed evidence of toxicity with little antitumor activity. Identification of topoisomerase I as the target of camptothecins and the need to preserve lactone structure allowed additional efforts to identify active members of this series. Topoisomerase I is responsible for unwinding the DNA strand by introducing single-strand breaks and allowing rotation of one strand about the other. In S-phase, topoisomerase I-induced breaks that are not promptly resealed lead to progress of the replication fork off the end of a DNA strand. The DNA damage is a potent signal for induction of apoptosis. Camptothecins promote the stabilization of the DNA linked to the enzyme in a so-called cleavable complex, analogous to the action of etoposide with topoisomerase II. Topotecan is a camptothecin derivative approved for use in gynecologic tumors and small-cell lung cancer. Toxicity is limited to myelosuppression and mucositis. CPT-11, or irinotecan, is a camptothecin with evidence of activity in colon carcinoma. In addition to myelosuppression, it causes a secretory diarrhea related to the toxicity of a metabolite called SN-38. Levels of SN-38 are particularly high in the setting of Gilbert's disease, characterized by defective glucuronyl transferase and indirect hyperbilirubinemia, a condition that affects about 10% of the white population in the United States. The diarrhea can be treated effectively with loperamide or octreotide.

INDIRECT MODULATORS OF NUCLEIC ACID FUNCTION: ANTIMETABOLITES A broad definition of antimetabolites would include compounds with structural similarity to precursors of purines or pyrimidines, or compounds that interfere with purine or pyrimidine synthesis. Some antimetabolites can cause DNA damage indirectly, through misincorporation into DNA, abnormal timing or progression through DNA synthesis, or

103e-14 altered function of pyrimidine and purine biosynthetic enzymes. They tend to convey greatest toxicity to cells in S-phase, and the degree of toxicity increases with duration of exposure. Common toxic manifestations include stomatitis, diarrhea, and myelosuppression. Second malignancies are not associated with their use.

Methotrexate inhibits dihydrofolate reductase, which regenerates reduced folates from the oxidized folates produced when thymidine monophosphate is formed from deoxyuridine monophosphate. Without reduced folates, cells die a “thymine-less” death. N5-Tetrahydrofolate or N5-formyltetrahydrofolate (leucovorin) can bypass this block and rescue cells from methotrexate, which is maintained in cells by polyglutamylated. The drug and other reduced folates are transported into cells by the folate carrier, and high concentrations of drug can bypass this carrier and allow diffusion of drug directly into cells. These properties have suggested the design of “high-dose” methotrexate regimens with leucovorin rescue of normal marrow and mucosa as part of curative approaches to osteosarcoma in the adjuvant setting and hematopoietic neoplasms of children and adults. Methotrexate is cleared by the kidney via both glomerular filtration and tubular secretion, and toxicity is augmented by renal dysfunction and drugs such as salicylates, probenecid, and nonsteroidal anti-inflammatory agents that undergo tubular secretion. With normal renal function, 15 mg/m² leucovorin will rescue 10⁻⁸ to 10⁻⁶ M methotrexate in three to four doses. However, with decreased creatinine clearance, doses of 50–100 mg/m² are continued until methotrexate levels are <5 × 10⁻⁸ M. In addition to bone marrow suppression and mucosal irritation, methotrexate can cause renal failure itself at high doses owing to crystallization in renal tubules; therefore, high-dose regimens require alkalization of urine with increased flow by hydration. Methotrexate can be sequestered in third-space collections and diffuse back into the general circulation, causing prolonged myelosuppression. Less frequent adverse effects include reversible increases in transaminases and hypersensitivity-like pulmonary syndrome. Chronic low-dose methotrexate can cause hepatic fibrosis. When administered to the intrathecal space, methotrexate can cause chemical arachnoiditis and CNS dysfunction.

Pemetrexed is a novel folate-directed antimetabolite. It is “multi-targeted” in that it inhibits the activity of several enzymes, including thymidylate synthetase, dihydrofolate reductase, and glycinamide ribonucleotide formyltransferase, thereby affecting the synthesis of both purine and pyrimidine nucleic acid precursors. To avoid significant toxicity to the normal tissues, patients receiving pemetrexed should also receive low-dose folate and vitamin B₁₂ supplementation. Pemetrexed has notable activity against certain lung cancers and, in combination with cisplatin, also against mesotheliomas. Pralatrexate is an antifolate approved for use in T cell lymphoma that is very efficiently transported into cancer cells.

5-Fluorouracil (5FU) represents an early example of “rational” drug design in that it originated from the observation that tumor cells incorporate radiolabeled uracil more efficiently into DNA than normal cells, especially gut. 5FU is metabolized in cells to 5’FdUMP, which inhibits thymidylate synthetase (TS). In addition, misincorporation can lead to single-strand breaks, and RNA can aberrantly incorporate FUMP. 5FU is metabolized by dihydropyrimidine dehydrogenase, and deficiency of this enzyme can lead to excessive toxicity from 5FU. Oral bioavailability varies unreliably, but orally administered analogues of 5FU such as capecitabine have been developed that allow at least equivalent activity to many parenteral 5FU-based approaches. Intravenous administration of 5FU leads to bone marrow suppression after short infusions but to stomatitis after prolonged infusions. Leucovorin augments the activity of 5FU by promoting formation of the ternary covalent complex of 5FU, the reduced folate, and TS. Less frequent toxicities include CNS dysfunction, with prominent cerebellar signs, and endothelial toxicity manifested by thrombosis, including pulmonary embolus and myocardial infarction.

Cytosine arabinoside (ara-C) is incorporated into DNA after formation of ara-CTP, resulting in S-phase-related toxicity. Continuous infusion schedules allow maximal efficiency, with uptake maximal at 5–7 μM. Ara-C can be administered intrathecally. Adverse effects

include nausea, diarrhea, stomatitis, chemical conjunctivitis, and cerebellar ataxia. Gemcitabine is a cytosine derivative that is similar to ara-C in that it is incorporated into DNA after anabolism to the triphosphate, rendering DNA susceptible to breakage and repair synthesis, which differs from that in ara-C in that gemcitabine-induced lesions are very inefficiently removed. In contrast to ara-C, gemcitabine appears to have useful activity in a variety of solid tumors, with limited nonmyelosuppressive toxicities.

6-Thioguanine and 6-mercaptopurine (6MP) are used in the treatment of acute lymphoid leukemia. Although administered orally, they display variable bioavailability. 6MP is metabolized by xanthine oxidase and therefore requires dose reduction when used with allopurinol. 6MP is also metabolized by thiopurine methyltransferase; genetic deficiency of thiopurine methyltransferase results in excessive toxicity.

Fludarabine phosphate is a prodrug of F-adenine arabinoside (F-ara-A), which in turn was designed to diminish the susceptibility of ara-A to adenosine deaminase. F-ara-A is incorporated into DNA and can cause delayed cytotoxicity even in cells with low growth fraction, including chronic lymphocytic leukemia and follicular B cell lymphoma. CNS and peripheral nerve dysfunction and T cell depletion leading to opportunistic infections can occur in addition to myelosuppression. 2-Chlorodeoxyadenosine is a similar compound with activity in hairy cell leukemia. 2-Deoxycytidine inhibits adenosine deaminase, with resulting increase in dATP levels. This causes inhibition of ribonucleotide reductase as well as augmented susceptibility to apoptosis, particularly in T cells. Renal failure and CNS dysfunction are notable toxicities in addition to immunosuppression. Hydroxyurea inhibits ribonucleotide reductase, resulting in S-phase block. It is orally bioavailable and useful for the acute management of myeloproliferative states.

Asparaginase is a bacterial enzyme that causes breakdown of extracellular asparagine required for protein synthesis in certain leukemic cells. This effectively stops tumor cell DNA synthesis, as DNA synthesis requires concurrent protein synthesis. The outcome of asparaginase action is therefore very similar to the result of the small-molecule antimetabolites. Because asparaginase is a foreign protein, hypersensitivity reactions are common, as are effects on organs such as pancreas and liver that normally require continuing protein synthesis. This may result in decreased insulin secretion with hyperglycemia, with or without hyperamylasemia and clotting function abnormalities. Close monitoring of clotting functions should accompany use of asparaginase. Paradoxically, owing to depletion of rapidly turning over anticoagulant factors, thromboses particularly affecting the CNS may also be seen with asparaginase.

MITOTIC SPINDLE INHIBITORS Microtubules are cellular structures that form the mitotic spindle, and in interphase cells, they are responsible for the cellular “scaffolding” along which various motile and secretory processes occur. Microtubules are composed of repeating noncovalent multimers of a heterodimer of α and β isoform of the protein tubulin. Vincristine binds to the tubulin dimer with the result that microtubules are disaggregated. This results in the block of growing cells in M-phase; however, toxic effects in G₁ and S-phase are also evident, reflecting effects on normal cellular activities of microtubules. Vincristine is metabolized by the liver, and dose adjustment in the presence of hepatic dysfunction is required. It is a powerful vesicant, and infiltration can be treated by local heat and infiltration of hyaluronidase. At clinically used intravenous doses, neurotoxicity in the form of glove-and-stocking neuropathy is frequent. Acute neuropathic effects include jaw pain, paralytic ileus, urinary retention, and the syndrome of inappropriate antidiuretic hormone secretion. Myelosuppression is not seen. Vinblastine is similar to vincristine, except that it tends to be more myelotoxic, with more frequent thrombocytopenia and also mucositis and stomatitis. Vinorelbine is a vinca alkaloid that appears to have differences in resistance patterns in comparison to vincristine and vinblastine; it may be administered orally.

The taxanes include paclitaxel and docetaxel. These agents differ from the vinca alkaloids in that the taxanes stabilize microtubules against depolymerization. The “stabilized” microtubules

function abnormally and are not able to undergo the normal dynamic changes of microtubule structure and function necessary for cell cycle completion. Taxanes are among the most broadly active antineoplastic agents for use in solid tumors, with evidence of activity in ovarian cancer, breast cancer, Kaposi's sarcoma, and lung tumors. They are administered intravenously, and paclitaxel requires use of a Cremophor-containing vehicle that can cause hypersensitivity reactions. Premedication with dexamethasone (8–16 mg orally or intravenously 12 and 6 h before treatment) and diphenhydramine (50 mg) and cimetidine (300 mg), both 30 min before treatment, decreases but does not eliminate the risk of hypersensitivity reactions to the paclitaxel vehicle. Docetaxel uses a polysorbate 80 formulation, which can cause fluid retention in addition to hypersensitivity reactions, and dexamethasone premedication with or without antihistamines is frequently used. A protein-bound formulation of paclitaxel (called *nab-paclitaxel*) has at least equivalent antineoplastic activity and decreased risk of hypersensitivity reactions. Paclitaxel may also cause hypersensitivity reactions, myelosuppression, neurotoxicity in the form of glove-and-stocking numbness, and paresthesia. Cardiac rhythm disturbances were observed in phase 1 and 2 trials, most commonly asymptomatic bradycardia but also, much more rarely, varying degrees of heart block. These have not emerged as clinically significant in the majority of patients. Docetaxel causes comparable degrees of myelosuppression and neuropathy. Hypersensitivity reactions, including bronchospasm, dyspnea, and hypotension, are less frequent but occur to some degree in up to 25% of patients. Fluid retention appears to result from a vascular leak syndrome that can aggravate preexisting effusions. Rash can complicate docetaxel administration, appearing prominently as a pruritic maculopapular rash affecting the forearms, but it has also been associated with fingernail ridging, breakdown, and skin discoloration. Stomatitis appears to be somewhat more frequent than with paclitaxel. Cabazitaxel is a taxane with somewhat better activity in prostate cancers than earlier generations of taxanes, perhaps due to superior delivery to sites of disease.

Resistance to taxanes has been related to the emergence of efficient efflux of taxanes from tumor cells through the p170 P-glycoprotein (*mdr* gene product) or the presence of variant or mutant forms of tubulin. Etoposides represent a class of novel microtubule-stabilizing agents that have been conscientiously optimized for activity in taxane-resistant tumors. Ixabepilone has clear evidence of activity in breast cancers resistant to taxanes and anthracyclines such as doxorubicin. It retains acceptable expected side effects, including myelosuppression, and can also cause peripheral sensory neuropathy. Eribulin is a microtubule-directed agent with activity in patients who have had progression of disease on taxanes and is more similar to vinca alkaloids in its action but has similar side effects as vinca alkaloids and taxanes.

Estramustine was originally synthesized as a mustard derivative that might be useful in neoplasms that possessed estrogen receptors. However, no evidence of interaction with DNA was observed. Surprisingly, the drug caused metaphase arrest, and subsequent study revealed that it binds to microtubule-associated proteins, resulting in abnormal microtubule function. Estramustine binds to estramustine-binding proteins (EMBPs), which are notably present in prostate tumor tissue, where the drug is used. Gastrointestinal and cardiovascular adverse effects related to the estrogen moiety occur in up to 10% of patients, including worsened heart failure and thromboembolic phenomena. Gynecomastia and nipple tenderness can also occur.

Targeted Chemotherapy • HORMONE RECEPTOR-DIRECTED THERAPY Steroid hormone receptor-related molecules have emerged as prominent targets for small molecules useful in cancer treatment. When bound to their cognate ligands, these receptors can alter gene transcription and, in certain tissues, induce apoptosis. The pharmacologic effect is a mirror or parody of the normal effects of the agents acting on nontransformed normal tissues, although the effects on tumors are mediated by indirect effects in some cases. While in some cases, such as breast cancer, demonstration of the target hormone receptor is necessary, in other cases such prostate cancer (androgen receptor) and

lymphoid neoplasms (glucocorticoid receptor), the relevant receptor is always present in the tumor.

Glucocorticoids are generally given in “pulsed” high doses in leukemias and lymphomas, where they induce apoptosis in tumor cells. Cushing's syndrome and inadvertent adrenal suppression on withdrawal from high-dose glucocorticoids can be significant complications, along with infections common in immunosuppressed patients, in particular *Pneumocystis pneumonia*, which classically appears a few days after completing a course of high-dose glucocorticoids.

Tamoxifen is a partial estrogen receptor antagonist; it has a 10-fold greater antitumor activity in breast cancer patients whose tumors express estrogen receptors than in those who have low or no levels of expression. It might be considered the prototypic “molecularly targeted” agent. Owing to its agonistic activities in vascular and uterine tissue, side effects include a somewhat increased risk of cardiovascular complications, such as thromboembolic phenomena, and a small increased incidence of endometrial carcinoma, which appears after chronic use (usually >5 years). Progestational agents—including medroxyprogesterone acetate, androgens including fluoxymesterone (Halotestin), and, paradoxically, estrogens—have approximately the same degree of activity in primary hormonal treatment of breast cancers that have elevated expression of estrogen receptor protein. Estrogen itself is not used often owing to prominent cardiovascular and uterotrophic activity.

Aromatase refers to a family of enzymes that catalyze the formation of estrogen in various tissues, including the ovary and peripheral adipose tissue and some tumor cells. Aromatase inhibitors are of two types, the irreversible steroid analogues such as exemestane and the reversible inhibitors such as anastrozole or letrozole. Anastrozole is superior to tamoxifen in the adjuvant treatment of breast cancer in postmenopausal patients with estrogen receptor-positive tumors. Letrozole treatment affords benefit following tamoxifen treatment. Adverse effects of aromatase inhibitors may include an increased risk of osteoporosis.

Prostate cancer is classically treated by androgen deprivation. Diethylstilbestrol (DES) acting as an estrogen at the level of the hypothalamus to downregulate hypothalamic luteinizing hormone (LH) production results in decreased elaboration of testosterone by the testicle. For this reason, orchiectomy is equally as effective as moderate-dose DES, inducing responses in 80% of previously untreated patients with prostate cancer but without the prominent cardiovascular side effects of DES, including thrombosis and exacerbation of coronary artery disease. In the event that orchiectomy is not accepted by the patient, testicular androgen suppression can also be effected by luteinizing hormone-releasing hormone (LHRH) agonists such as leuprolide and goserelin. These agents cause tonic stimulation of the LHRH receptor, with the loss of its normal pulsatile activation resulting in decreased output of LH by the anterior pituitary. Therefore, as primary hormonal manipulation in prostate cancer, one can choose orchiectomy or leuprolide, but not both. The addition of androgen receptor blockers, including flutamide or bicalutamide, is of uncertain additional benefit in extending overall response duration; the combined use of orchiectomy or leuprolide plus flutamide is referred to as *total androgen blockade*. Enzalutamide also binds to the androgen receptor and antagonizes androgen action in a mechanistically distinct way. Somewhat analogous to inhibitors of aromatase, agents have been derived that inhibit testosterone and other androgen synthesis in the testis, adrenal gland, and prostate tissue. Abiraterone inhibits 17 α -hydroxylase/C17,20 lyase (CYP 17A1) and has been shown to be active in prostate cancer patients experiencing progression despite androgen blockade.

Tumors that respond to a primary hormonal manipulation may frequently respond to second and third hormonal manipulations. Thus, breast tumors that had previously responded to tamoxifen have, on relapse, notable response rates to withdrawal of tamoxifen itself or to subsequent addition of an aromatase inhibitor or progestin. Likewise, initial treatment of prostate cancers with leuprolide plus flutamide may be followed after disease progression by response to withdrawal of flutamide. These responses may result from the removal

103e-16 of antagonists from mutant steroid hormone receptors that have come to depend on the presence of the antagonist as a growth-promoting influence.

Additional strategies to treat refractory breast and prostate cancers that possess steroid hormone receptors may also address adrenal capacity to produce androgens and estrogens, even after orchiectomy or oophorectomy, respectively. Thus, aminoglutethimide or ketoconazole can be used to block adrenal synthesis by interfering with the enzymes of steroid hormone metabolism. Administration of these agents requires concomitant hydrocortisone replacement and additional glucocorticoid doses administered in the event of physiologic stress.

Humoral mechanisms can also result in complications from an underlying malignancy producing the hormone. Adrenocortical carcinomas can cause Cushing's syndrome as well as syndromes of androgen or estrogen excess. Mitotane can counteract these by decreasing synthesis of steroid hormones. Islet cell neoplasms can cause debilitating diarrhea, treated with the somatostatin analogue octreotide. Prolactin-secreting tumors can be effectively managed by the dopaminergic agonist bromocriptine.

DIAGNOSTICALLY GUIDED THERAPY The basis for discovery of drugs of this type was the prior knowledge of the importance of the drugs' molecular target to drive tumors in different contexts. **Figure 103e-4** summarizes how FDA-approved targeted agents act. In the case of diagnostically guided targeted chemotherapy, prior demonstration of a specific target is necessary to guide the rational use of the agent, while in the case of targeted agents directed at oncogenic pathways, specific diagnosis of pathway activation is not yet necessary or in some cases feasible, although this is an area of ongoing clinical research. **Table 103e-5** lists currently approved targeted chemotherapy agents, with features of their use.

In hematologic tumors, the prototypic agent of this type is imatinib, which targets the ATP binding site of the p210^{bcr-abl} protein tyrosine kinase that is formed as the result of the chromosome 9;22 translocation producing the Philadelphia chromosome in CML. Imatinib is superior to interferon plus chemotherapy in the initial treatment of the chronic phase of this disorder. It has lesser activity in the blast phase of CML, where the cells may have acquired additional mutations in p210^{bcr-abl} itself or other genetic lesions. Its side effects are relatively tolerable in most patients and include hepatic dysfunction, diarrhea, and fluid retention. Rarely, patients receiving imatinib have decreased cardiac function, which may persist after discontinuation of the drug. The quality of response to imatinib enters into the decision about when to refer patients with CML for consideration of transplant approaches. Nilotinib is a tyrosine protein kinase inhibitor with a similar spectrum of activity to imatinib, but with increased potency and perhaps better tolerance by certain patients. Dasatinib, another inhibitor of the p210^{bcr-abl} oncoproteins, is active in certain mutant variants of p210^{bcr-abl} that are refractory to imatinib and arise during therapy with imatinib or are present de novo. Dasatinib also has inhibitory action against kinases belonging to the src tyrosine protein kinase family; this activity may contribute to its effects in hematopoietic tumors and suggest a role in solid tumors where src kinases are active. The T3151 mutant of p210^{bcr-abl} is resistant to imatinib, nilotinib, bosutinib, and dasatinib; ponatinib has activity in patients with this p210^{bcr-abl} variant, but ponatinib has noteworthy associated thromboembolic toxicity. Use of this class of targeted agents is thus critically guided not only by the presence of the p210^{bcr-abl} tyrosine kinase, but also by the presence of different mutations in the ATP binding site.

All-trans-retinoic acid (ATRA) targets the PML-retinoic acid receptor (RAR) a fusion protein, which is the result of the chromosome 15;17 translocation pathogenic for most forms of APL. Administered orally, it causes differentiation of the neoplastic promyelocytes to mature granulocytes and attenuates the rate of hemorrhagic complications. Adverse effects include headache with or without pseudotumor cerebri and gastrointestinal and cutaneous toxicities.

In epithelial solid tumors, the small-molecule epidermal growth factor (EGF) antagonists act at the ATP binding site of the EGF receptor

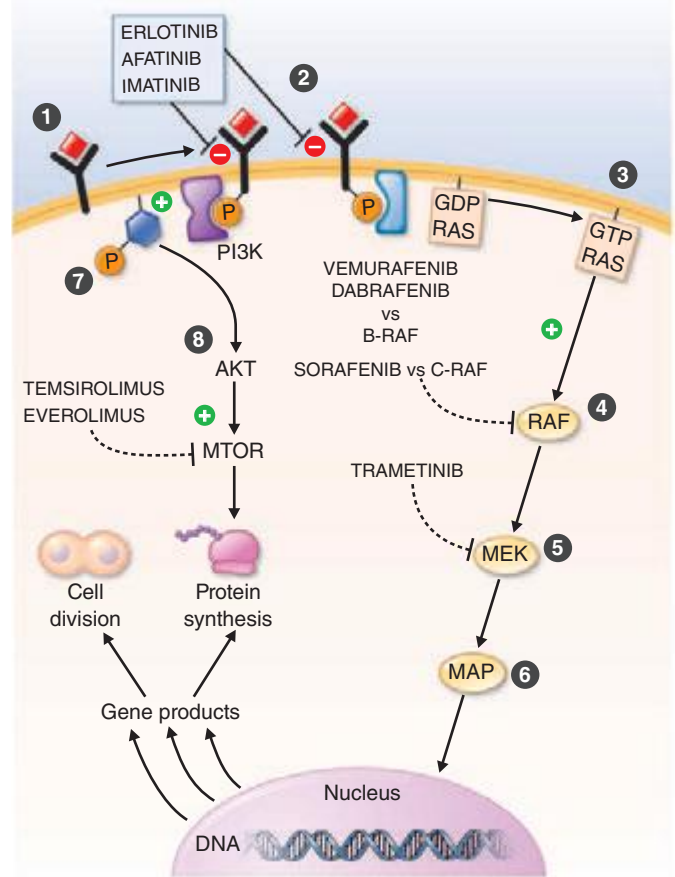


FIGURE 103e-4 Targeted chemotherapeutic agents act in most instances by interrupting cell growth factor-mediated signaling pathways. After a growth factor binds to its cognate receptor (1), in many cases there is activation of tyrosine kinase activity particularly after dimerization of the receptors (2). This leads to autophosphorylation of the receptor and docking of “adaptor” proteins. One important pathway activated occurs after exchange of GDP for GTP in the RAS family of proto-oncogene products (3). GTP-RAS activates the RAF proto-oncogene kinase (4), leading to a phosphorylation cascade of kinases (5, 6) that ultimately impart signals to regulators of gene function to produce transcripts which activate cell cycle progression and increase protein synthesis. In parallel, tyrosine phosphorylated receptors can activate the phosphatidylinositol-3-kinase to produce the phosphorylated lipid phosphatidylinositol-3-phosphate (7). This leads to the activation of the AKT kinase (8) which in turn stimulates the mammalian “Target of Rapamycin” kinase (mTOR), which directly increases the translation of key mRNAs for gene products regulating cell growth. Erlotinib and afatinib, are examples of Epidermal Growth Factor receptor tyrosine kinase inhibitors; imatinib can act on the nonreceptor tyrosine kinase bcr-abl or c-KIT membrane bound tyrosine kinase. Vemurafenib and Dabrafenib act on the B isoform of RAF uniquely in melanoma, and c-RAF is inhibited by sorafenib. Trametinib acts on MEK. Temsirolimus and everolimus inhibit mTOR kinase to downregulate translation of oncogenic mRNAs.

tyrosine kinase. In early clinical trials, gefitinib showed evidence of responses in a small fraction of patients with non-small-cell lung cancer (NSCLC). Side effects were generally acceptable, consisting mostly of rash and diarrhea. Subsequent analysis of responding patients revealed a high frequency of activating mutations in the EGF receptor. Patients with such activating mutations who initially responded to gefitinib but who then had progression of the disease then acquired additional mutations in the enzyme, analogous functionally to mutational variants responsible for imatinib resistance in CML. Erlotinib is another EGF receptor tyrosine kinase antagonist with a superior outcome in clinical

TABLE 103e-5 MOLECULARLY TARGETED AGENTS

Drug	Target	Adverse Events	Notes
Diagnostically Guided Protein Kinase Antagonists			
Imatinib	Bcr-Abl fusion protein (CML/ALL); c-kit mutants, PDGFR variants (GI stromal tumor; eosinophilic syndromes)	Nausea Periorbital edema Rare CHF QTc prolongation	Myelosuppression not frequent in solid tumor indications
Nilotinib	Bcr-Abl fusion protein (CML) and some imatinib-resistant variants	Interaction with CYP3A4-metabolized drugs CHF Hepatotoxicity Hypothyroidism	Chronic phase and in patients resistant to imatinib
Dasatinib	Bcr-Abl fusion protein (CML/ALL); wild-type and imatinib-resistant mutants	Myelosuppression (bleeding, infection) Pulmonary hypertension CHF Fluid retention QTc prolongation	Chronic phase and imatinib or nilotinib resistant
Bosutinib	Bcr-Abl fusion protein (CML); wild-type and imatinib-resistant mutants	Myelosuppression Hepatic QTc prolongation	Chronic phase and imatinib or nilotinib resistant
Ponatinib	T315I mutation of Bcr-Abl fusion protein (CML)	Clotting Hepatic CHF Pancreatitis Neuropathy Rash	
Gefitinib	First-line treatment of NSCLC with ATP site mutation of EGFR	Diarrhea Interstitial pneumonitis	In United States, only with prior documented benefit in second-line treatment of NSCLC
Erlotinib	First-line treatment of NSCLC with ATP site mutation of EGFR; second-line treatment of wild-type EGFR NSCLC	Rash Diarrhea Rare interstitial pneumonitis	1 h before, 2 h after meals
Afatinib	First-line treatment of NSCLC with ATP site mutation of EGFR	Diarrhea Cutaneous	Interacts with Pgp inhibitors
Crizotinib	EML4-Alk fusion protein	Interstitial pneumonitis Hepatic QTc prolongation Bradycardia	
Vemurafenib	<i>BRAF</i> V600E in melanoma	Nausea Rash Cutaneous Second cutaneous neoplasms	
Dabrafenib	<i>BRAF</i> V600E in melanoma	Cutaneous Second cutaneous neoplasms	
Trametinib	<i>BRAF</i> V600E in melanoma (both as single agent and in combination with dabrafenib)	Rash Diarrhea Lymphedema	In combination with dabrafenib, second neoplasms, hemorrhage, venous thrombosis, CHF, ocular, hyperglycemia
Drug	Indication	Adverse Events	Notes
Diagnostically Guided Retinoid			
Tretinoin	APL t(15,17)	Teratogenic Cutaneous	APL differentiation syndrome: pulmonary dysfunction/infiltrate, pleural/pericardial effusion, fever
Nondiagnostically Guided Agents			
Retinoid			
Bexarotene	Cutaneous T cell lymphoma	Hypercholesterolemia Hypertriglyceridemia Cutaneous Teratogenic	Central hypothyroidism

(Continued)

TABLE 103e-5 MOLECULARLY TARGETED AGENTS (CONTINUED)

Drug	Target	Adverse Events	Notes
Multikinase inhibitors			
Sorafenib	Renal cell, hepatocellular, differentiated thyroid carcinoma	Diarrhea Hand-foot syndrome Other rash Hypertension CHF	Targets c-raf, VEGFR
Pazopanib	Renal cell carcinoma, soft tissue sarcoma	Fatigue Diarrhea/GI Hypertension Thromboses QTc	Target VEGFR, c-kit, PDGFR
Regorafenib	Second-line colorectal cancer; GI stromal tumor	Hypertension Hand-foot syndrome Thromboses Perforations	VEGFR/TIE2
Sunitinib	Renal cell carcinoma, pancreatic neuroendocrine tumor, GI stromal tumor	Fatigue Diarrhea Neutropenia	Target VEGFR
Vandetanib	Medullary thyroid cancer	Diarrhea Rash Hypertension Prolonged QTc Thromboses	Target VEGFR, ret, EGFR
Cabozantinib	Medullary thyroid cancer	Hypertension Wound healing Fistulas Osteonecrosis Proteinuria	Target VEGFR, c-met
Axitinib	Renal cell carcinoma, second line	Diarrhea/other GI Fatigue Hand-foot syndrome	Target VEGFR, PDGFR, c-kit
Proteasome Inhibitors			
Bortezomib	Multiple myeloma, mantle cell lymphoma	Neuropathy Thrombocytopenia GI	
Carfilzomib	Multiple myeloma, second line	Infusion reaction CHF Thrombocytopenia Pulmonary Tumor lysis	
Histone Deacetylase Inhibitors			
Vorinostat	Cutaneous T cell lymphoma, second line	Fatigue Diarrhea Thrombocytopenia Embolism	
Romidepsin	Cutaneous T cell lymphoma, second line	Nausea Vomiting Cytopenias Cardiac conduction	
mTOR Inhibitors			
Temsirolimus	Renal cell carcinoma, second line or poor prognosis	Stomatitis Thrombocytopenia Nausea Anorexia, fatigue Metabolic (glucose, lipid)	

(Continued)

TABLE 103e-5 MOLECULARLY TARGETED AGENTS (CONTINUED)

Drug	Target	Adverse Events	Notes
Everolimus	Renal cell carcinoma, advanced; subependymal giant-cell astrocytoma; breast cancer, hormone receptor positive, resistant to antiestrogen; pancreatic neuroendocrine	Stomatitis Fatigue	
Miscellaneous			
Arsenic trioxide	APL	↑ QTc	APL differentiation syndrome (see under tretinoin)
Vismodegib	Metastatic basal cell carcinoma	GI Hair loss Fatigue Muscle spasm Dysgeusia	Target smoothed receptor in hedgehog pathway

Abbreviations: APL, acute promyelocytic leukemia; ALL, acute lymphocytic leukemia; CHF, congestive heart failure; CML, chronic myeloid leukemia; EGFR, epidermal growth factor receptor; GI, gastrointestinal; mTOR, mammalian target of rapamycin kinase; NSCLC, non-small-cell lung cancer; PDGFR, platelet-derived growth factor receptor; Pgp, P-glycoprotein; VEGFR, vascular endothelial growth factor receptor.

trials in NSCLC; an overall survival advantage was demonstrated in subsets of patients who were treated after demonstrating progression of disease and who also had not been preselected for the presence of activating mutations. Thus, although even patients with wild-type EGF receptors may benefit from erlotinib treatment, the presence of EGF receptor tyrosine kinase mutations has recently been shown to be a basis for recommending erlotinib and afatinib for first-line treatment of advanced NSCLC. Likewise, crizotinib targeting the *alk* protooncogene fusion protein has value in the initial treatment of *alk*-positive NSCLC. Lapatinib is a tyrosine kinase inhibitor with both EGF receptor and HER2/neu antagonist activity, which is important in the treatment of breast cancers expressing the HER2/neu oncoprotein.

In addition to the p210^{bcr-abl} kinase, imatinib also has activity against the c-kit tyrosine kinase (the receptor for the *steel* growth factor, also called stem cell factor) and the platelet-derived growth factor receptor (PDGFR), both of which can be expressed in gastrointestinal stromal sarcoma (GIST). Imatinib has found clinical utility in GIST, a tumor previously notable for its refractoriness to chemotherapeutic approaches. Imatinib's degree of activity varies with the specific mutational variant of kit or PDGFR present in a particular patient's tumor.

The *BRAF* V600E mutation has been detected in a notable fraction of melanomas, thyroid tumors, and hairy cell leukemia, and preclinical models supported the concept that *BRAF* V600E drives oncogenic signaling in these tumors. Vemurafenib and dabrafenib, with selective capacity to inhibit the *BRAF* V600E serine kinase activity, were each shown to cause noteworthy responses in patients with *BRAF* V600E-mutated melanomas, although early relapse occurred in many patients treated with the drugs as single agents. Trametinib, acting downstream of *BRAF* V600E by directly inhibiting the MEK serine kinase by a non-ATP binding site mechanism, also displayed noteworthy responses in *BRAF* V600E-mutated melanomas, and the combination of trametinib and dabrafenib is even more active, by targeting the *BRAF* V600E-driven pathway at two points in the pathway leading to gene activation.

ONCOGENICALLY ACTIVATED PATHWAYS This group of agents also targets specific regulatory molecules in promoting the viability of tumor cells, but they do not require the diagnostically verified presence of a particular target or target variant at this time.

"Multitargeted" kinase antagonists are small-molecule ATP site-directed antagonists that inhibit more than one protein kinase and have value in the treatment of several solid tumors. Drugs of this type with prominent activity against the vascular endothelial growth factor receptor (VEGFR) tyrosine kinase have activity in renal cell carcinoma. Sorafenib is a VEGFR antagonist with activity against the *raf* serine-threonine protein kinase, and regorafenib is a closely related drug with value in relapsed advanced colon cancer. Pazopanib also prominently targets VEGFR and has activity in renal carcinoma and soft tissue sarcomas. Sunitinib has anti-VEGFR, anti-PDGFR, and

anti-c-kit activity. It causes prominent responses and stabilization of disease in renal cell cancers and GISTs. Side effects for agents with anti-VEGFR activity prominently include hypertension, proteinuria, and, more rarely, bleeding and clotting disorders and perforation of scarred gastrointestinal lesions. Also encountered are fatigue, diarrhea, and the hand-foot syndrome, with erythema and desquamation of the distal extremities, in some cases requiring dose modification, particularly with sorafenib.

Temsirolimus and everolimus are mammalian target of rapamycin (mTOR) inhibitors with activity in renal cancers. They produce stomatitis, fatigue, and some hyperlipidemia (10%), myelosuppression (10%), and rare lung toxicity. Everolimus is also useful in patients with hormone receptor-positive breast cancers displaying resistance to hormonal inhibition and in certain neuroendocrine and brain tumors, the latter arising in patients with sporadic or inherited mutations in the pathway activating mTOR.

In hematologic neoplasms, bortezomib is an inhibitor of the proteasome, the multisubunit assembly of protease activities responsible for the selective degradation of proteins important in regulating activation of transcription factors, including nuclear factor- κ B (NF- κ B) and proteins regulating cell cycle progression. It has activity in multiple myeloma and certain lymphomas. Adverse effects include neuropathy, orthostatic hypotension with or without hyponatremia, and reversible thrombocytopenia. Carfilzomib is a proteasome inhibitor chemically unrelated to bortezomib without prominent neuropathy, but with evidence of a cytokine release syndrome, which can be a cardiopulmonary stress. Other agents active in multiple myeloma and certain other hematologic neoplasms include the immunomodulatory agents related to thalidomide, including lenalidomide and pomalidomide. All these agents collectively inhibit aberrant angiogenesis in the bone marrow microenvironment, as well as influence stromal cell immune functions to alter the cytokine milieu supporting the growth of myeloma cells. Thalidomide, although clinically active, has prominent cytopenic, neuropathic, procoagulant, and CNS toxicities that have been somewhat attenuated in the other drugs of the class, although use of these agents frequently entails concomitant anticoagulant prophylaxis.

Ibrutinib is representative a novel class of inhibitors directed at Bruton's tyrosine kinase, which is important in the function of B cells. Initially approved for use in mantle cell lymphoma, it is potentially applicable to a number of B cell neoplasms that depend on signals through the B cell antigen receptor. Janus kinases likewise function downstream of a variety of cytokine receptors to amplify cytokine signals, and Janus kinase inhibitors including ruxolitinib have approved activity in myelofibrosis to ameliorate splenomegaly and systemic symptoms.

Vorinostat is an inhibitor of histone deacetylases, which are responsible for maintaining the proper orientation of histones on DNA, with resulting capacity for transcriptional readiness. Acetylated histones allow access of transcription factors to target genes and therefore

increase expression of genes that are selectively repressed in tumors. The result can be differentiation with the emergence of a more normal cellular phenotype, or cell cycle arrest with expression of endogenous regulators of cell cycle progression. Vorinostat is approved for clinical use in cutaneous T cell lymphoma, with dramatic skin clearing and very few side effects. Romidepsin is a distinct molecular class of histone deacetylase inhibitor also active in cutaneous T cell lymphoma. An active retinoid in cutaneous T cell lymphoma is the synthetic retinoid X receptor ligand bexarotene.

DNA methyltransferase inhibitors, including 5-aza-cytidine and 2'-deoxy-5-azacytidine (decitabine), can also increase transcription of genes "silenced" during the pathogenesis of a tumor by causing demethylation of the methylated cytosines that are acquired as an "epigenetic" (i.e., after the DNA is replicated) modification of DNA. These drugs were originally considered antimetabolites but have clinical value in myelodysplastic syndromes and certain leukemias when administered at low doses.

CANCER BIOLOGIC THERAPY

Principles The goal of biologic therapy is to manipulate the host-tumor interaction in favor of the host, potentially at an optimum biologic dose that might be different than the MTD. As a class, biologic therapies may be distinguished from molecularly targeted agents in that many biologic therapies require an active response (e.g., reexpression of silenced genes or antigen expression) on the part of the tumor cell or on the part of the host (e.g., immunologic effects) to allow therapeutic effect. This may be contrasted with the more narrowly defined antiproliferative or apoptotic response that is the ultimate goal of molecularly targeted agents discussed above. However, there is much commonality in the strategies to evaluate and use molecularly targeted and biologic therapies.

Immune Cell-Mediated Therapies Tumors have a variety of means of avoiding the immune system: (1) they are often only subtly different from their normal counterparts; (2) they are capable of downregulating their major histocompatibility complex antigens, effectively masking them from recognition by T cells; (3) they are inefficient at presenting antigens to the immune system; (4) they can cloak themselves in a protective shell of fibrin to minimize contact with surveillance mechanisms; and (5) they can produce a range of soluble molecules, including potential immune targets, that can distract the immune system from recognizing the tumor cell or can kill or inactivate the immune effector cells. Some of the cell products initially polarize the immune response away from cellular immunity (shifting from T_H1 to T_H2 responses; [Chap. 372e](#)) and ultimately lead to defects in T cells that prevent their activation and cytotoxic activity. Cancer treatment further suppresses host immunity. A variety of strategies are being tested to overcome these barriers.

Cell-Mediated Immunity The strongest evidence that the immune system can exert clinically meaningful antitumor effects comes from allogeneic bone marrow transplantation. Adoptively transferred T cells from the donor expand in the tumor-bearing host, recognize the tumor as being foreign, and can mediate impressive antitumor effects (graft-versus-tumor effects). Three types of experimental interventions are being developed to take advantage of the ability of T cells to kill tumor cells.

1. *Transfer of allogeneic T cells.* This occurs in three major settings: in allogeneic bone marrow transplantation; as purified lymphocyte transfusions following bone marrow recovery after allogeneic bone marrow transplantation; and as pure lymphocyte transfusions following immunosuppressive (nonmyeloablative) therapy (also called minitransplants). In each of these settings, the effector cells are donor T cells that recognize the tumor as being foreign, probably through minor histocompatibility differences. The main risk of such therapy is the development of graft-versus-host disease because of the minimal difference between the cancer and the normal host cells. This approach has been highly effective in certain hematologic cancers.

2. *Transfer of autologous T cells.* In this approach, the patient's own T cells are removed from the tumor-bearing host, manipulated in several ways in vitro, and given back to the patient. There are three major classes of autologous T-cell manipulation. First, tumor antigen-specific T cells can be developed and expanded to large numbers over many weeks ex vivo before administration. Second, the patient's T cells can be activated by exposure to polyclonal stimulators such as anti-CD3 and anti-CD28 after a short period ex vivo, and then amplified in the host after transfer by stimulation with IL-2, for example. Short periods removed from the patient permit the cells to overcome the tumor-induced T cell defects, and such cells traffic and home to sites of disease better than cells that have been in culture for many weeks. In a third approach, genes that encode for a T cell receptor specific for an antigen expressed by the tumor along with genes that facilitate T cell activation can be introduced into subsets of a patient's T cells, which, after transfer back into the patient, allow homing of cytotoxic T cells to tumor cells expressing the antigen.

3. *Tumor vaccines aimed at boosting T cell immunity.* The finding that mutant oncogenes that are expressed only intracellularly can be recognized as targets of T cell killing greatly expanded the possibilities for tumor vaccine development. No longer is it difficult to find something different about tumor cells. However, major difficulties remain in getting the tumor-specific peptides presented in a fashion to prime the T cells. Tumors themselves are very poor at presenting their own antigens to T cells at the first antigen exposure (*priming*). Priming is best accomplished by professional antigen-presenting cells (dendritic cells). Thus, a number of experimental strategies are aimed at priming host T cells against tumor-associated peptides. Vaccine adjuvants such as granulocyte-macrophage colony-stimulating factor (GM-CSF) appear capable of attracting antigen-presenting cells to a skin site containing a tumor antigen. Such an approach has been documented to eradicate microscopic residual disease in follicular lymphoma and give rise to tumor-specific T cells. Purified antigen-presenting cells can be pulsed with tumor, its membranes, or particular tumor antigens and delivered as a vaccine. One such vaccine, Sipuleucel-T, is approved for use in patients with hormone-independent prostate cancer. In this approach, the patient undergoes leukapheresis, wherein mononuclear cells (that include antigen-presenting cells) are removed from the patient's blood. The cells are pulsed in a laboratory with an antigenic fusion protein comprising a protein frequently expressed by prostate cancer cells, prostate acid phosphatase, fused to GM-CSF, and matured to increase their capacity to present the antigen to immune effector cells. The cells are then returned to the patient in a well-tolerated treatment. Although no objective tumor response was documented in clinical trials, median survival was increased by about 4 months. Tumor cells can also be transfected with genes that attract antigen-presenting cells.

Another important vaccine strategy is directed at infectious agents whose action ultimately is tied to the development of human cancer. Hepatitis B vaccine in an epidemiologic sense prevents hepatocellular carcinoma, and a tetravalent human papillomavirus vaccine prevents infection by virus types currently accounting for 70% of cervical cancer. Unfortunately, these vaccines are ineffective at treating patients who have developed a virus-induced cancer.

Antibody-Mediated Therapeutic Approaches In general, antibodies are not very effective at killing cancer cells. Because the tumor seems to influence the host toward making antibodies rather than generating cellular immunity, it is inferred that antibodies are easier for the tumor to fend off. Many patients can be shown to have serum antibodies directed at their tumors, but these do not appear to influence disease progression. However, the ability to grow very large quantities of high-affinity antibody directed at a tumor by the hybridoma technique has led to the application of antibodies in the treatment of cancer. In this approach, antibodies are derived where the antigen-combining regions are grafted onto human immunoglobulin gene products (chimerized or humanized) or derived de novo from mice bearing

TABLE 103e-6 ANTIBODIES USED IN CANCER TREATMENT

Drug	Target	Indications and Features of Use
Tumor Regulatory Antibodies		
Rituximab	CD20	B cell neoplasms (also emerging role in autoimmune disease); chimeric antibody with frequent mouse-derived sequences; frequent infusion reactions, particularly on initial doses; reactivation of infections, particularly hepatitis; progressive multifocal leukoencephalopathy; tumor lysis syndrome
Ofatumumab	CD20	active in CLL; fully human antibody with distinct binding site compared to rituximab; decreased intensity infusion reactions;
Trastuzumab	HER2/neu	Active in breast cancer and GI cancers expressing HER2/neu; cardiotoxicity, particularly in setting of prior anthracyclines, requires monitoring; infusion reactions
Pertuzumab	HER2/neu	Breast cancer; targets distinct binding site from trastuzumab, inhibiting dimerization of HER2 family members; infusion reactions; cardiac toxicity
Cetuximab	EGFR	Colorectal cancers with wild-type Ki-ras oncoprotein; head and neck cancers with radiation; rash, diarrhea, infusion reactions
Panitumumab	EGFR	Colorectal cancers with wild-type Ki-ras oncoprotein; fully humanized; decreased infusion reactions; different IgG subtype than cetuximab
Bevacizumab	VEGF	Metastatic colorectal cancer and non-small-cell lung cancer (nonsquamous) with chemotherapy; renal cancer and glioblastoma as single agents; prominent HBP, proteinuria, GI perforations, hemorrhage, thrombosis (venous and arterial)
Immunoregulatory Antibodies		
Alemtuzumab	CD52	CLL, T cell lymphomas; activates complement after binding to cell surface; infusion reactions, hypersensitivity, tumor lysis, activation of infections, cytopenias
Ipilimumab	CTLA4	Melanoma; inhibits the negative proliferative signal to T cells acting through CTLA4, resulting in prominent T cell activation; side effects include immune-mediated toxicity to liver, skin, pituitary, gut, which if severe calls for steroids, which inhibit antineoplastic effect
Pembrolizumab	PD-1	Melanoma unresectable or metastatic and if B-RAF V600 mutated, refractory to a B-RAF inhibitor; also can cause immune related colitis, hepatitis, hypophysitis, nephritis, and altered thyroid function; also consider steroids for treatment of severe adverse events

Abbreviations: CLL, chronic lymphocytic leukemia; EGFR, epidermal growth factor receptor; GI, gastrointestinal; HBP, high blood pressure; VEGF, vascular endothelial growth factor.

human immunoglobulin gene loci. Three general strategies have emerged using antibodies. *Tumor-regulatory antibodies* target tumor cells directly or indirectly to modulate intracellular functions or attract immune or stromal cells. *Immunoregulatory antibodies* target antigens expressed on the tumor cells or host immune cells to modulate primarily the host's immune responsiveness to the tumor. Finally, *antibody conjugates* can be made with the antibody linked to drugs, toxins, or radioisotopes to target these "warheads" for delivery to the tumor. **Table 103e-6** lists features of currently used or promising antibodies for cancer treatment.

TUMOR-REGULATORY ANTIBODIES Humanized antibodies against the CD20 molecule expressed on B cell lymphomas (rituximab and ofatumumab) are exemplary of antibodies that affect both signaling events driving lymphomagenesis as well as activating immune responses against B cell neoplasms. They are used as single agents and in combination with chemotherapy and radiation in the treatment of B cell neoplasms. Obinutuzumab is an antibody with an altered glycosylation that enhances its ability to fix complement; it is also directed against CD20 and is of value in chronic lymphocytic leukemia. It seems to be more effective in this setting than rituximab.

The HER2/neu receptor overexpressed on epithelial cancers, especially breast cancer, was initially targeted by trastuzumab, with noteworthy activity in potentiating the action of chemotherapy in breast cancer as well as some evidence of single-agent activity. Trastuzumab also appears to interrupt intracellular signals derived from HER2/neu and to stimulate immune mechanisms. The anti-HER2 antibody pertuzumab, specifically targeting the domain of HER2/neu responsible for dimerization with other HER2 family members, is more specifically directed against HER2 signaling function and augments the action of trastuzumab.

EGF receptor (EGFR)-directed antibodies (such as cetuximab and panitumumab) have activity in colorectal cancer refractory to chemotherapy, particularly when used to augment the activity of an additional chemotherapy program, and in the primary treatment of head and neck cancers treated with radiation therapy. The mechanism of action is unclear. Direct effects on the tumor may mediate an antiproliferative effect as well as stimulate the participation of host mechanisms involving immune cell or complement-mediated response to tumor cell-bound antibody. Alternatively, the antibody may alter the release of paracrine factors promoting tumor cell survival.

The anti-VEGF antibody bevacizumab shows little evidence of antitumor effect when used alone, but when combined with chemotherapeutic agents, it improves the magnitude of tumor shrinkage and time to disease progression in colorectal and nonsquamous lung cancers. The mechanism for the effect is unclear and may relate to the capacity of the antibody to alter delivery and tumor uptake of the active chemotherapeutic agent. Ziv-aflibercept is not an antibody, but a solubilized VEGF receptor VEGF binding domain, and therefore may have a distinct mechanism of action with comparable side effects.

Unintended side effects of any antibody use include infusion-related hypersensitivity reactions, usually limited to the first infusion, which can be managed with glucocorticoid and/or antihistamine prophylaxis. In addition, distinct syndromes have emerged with different antibodies. Anti-EGFR antibodies produce an acneiform rash that poorly responds to glucocorticoid cream treatment. Trastuzumab (anti-HER2) can inhibit cardiac function, particularly in patients with prior exposure to anthracyclines. Bevacizumab has a number of side effects of medical significance, including hypertension, thrombosis, proteinuria, hemorrhage, and gastrointestinal perforations with or without prior surgeries; these adverse events also occur with small-molecule drugs modulating VEGFR function.

IMMUNOREGULATORY ANTIBODIES Purely immunoregulatory antibodies stimulate immune responses to mediate tumor-directed cytotoxicity. First-generation approaches sought to activate complement and are exemplified by antibodies to CD52; these are active in chronic lymphoid leukemia and T cell malignancies. A more refined understanding of the tumor-host interface has defined that cytotoxic tumor-directed T cells are frequently inhibited by ligands upregulated in the tumor cells. The programmed death ligand 1 (PD-L1; also known as B7-homolog 1) was initially recognized as an entity that induced T cell death through a receptor present on T cells, termed the PD receptor (**Fig 103e-5**), which physiologically exists to regulate the intensity of the immune response. The PD family of ligands and receptors also regulates macrophage function, present in tumor stroma. These actions raised the hypothesis that antibodies directed against the PD signaling axis (both anti-PD-L1 and anti-PD) might be useful in cancer treatment by allowing reactivation of the immune response against tumors. Indeed, nivolumab and lambrolizumab, both anti-PD antibodies, have shown evidence of important immune-mediated actions against certain solid tumors, including melanoma and lung cancers.

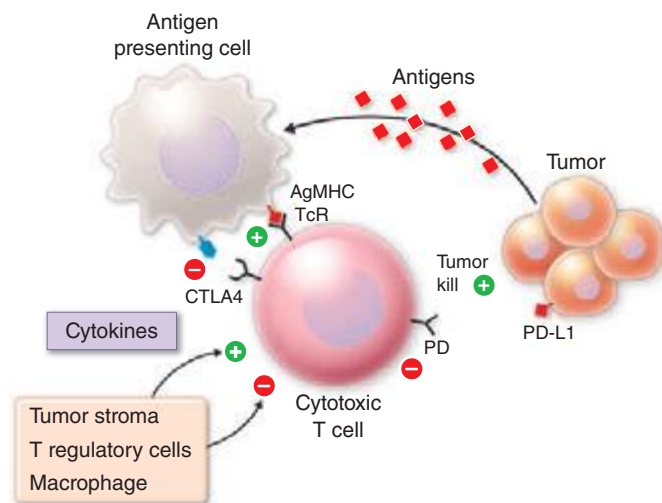


FIGURE 103e-5 Tumors possess a microenvironment (tumor stroma) with immune cells including both helper T cells, suppressor T cells (both “regulatory” of other immune cell function), macrophages, and cytotoxic T cells. Cytokines found in the stroma and deriving from macrophages and regulatory T cells modulate the activities of cytotoxic T cells, which have the potential to kill tumor cells. Antigens released by tumor cells are taken up by Antigen Presenting Cells (APCs), also in the stroma. Antigens are processed by the APCs to peptides presented by the Major Histocompatibility Complex to T-cell antigen receptors, thus providing an (+) activation signal for the cytotoxic tumor cells to kill tumor cells bearing that antigen. Negative (–) signals inhibiting cytotoxic T cell action include the CTLA4 receptor (on T cells), interacting with the B7 family of negative regulatory signals from APCs, and the PD receptor (on T cells), interacting with the PD-L1 (–) signal coming from tumor cells expressing the PD-1 ligand (PD-L1). As both CTLA4 and PD1 signals attenuate the anti-tumor T cell response, strategies which inhibit CTLA4 and PD1 function are a means of stimulating cytotoxic T cell activity to kill tumor cells. Cytokines from other immune cells and macrophages can provide both (+) and (–) signals for T cell action, and are under investigation as novel immunoregulatory therapeutics.

Already approved for clinical use in melanoma is ipilimumab, an antibody directed against the anti-CTLA4 (cytotoxic T lymphocyte antigen 4), which is expressed on T cells (not tumor cells), responds to signals from antigen-presenting cells (Fig. 103e-5), and also down-regulates the intensity of the T cell proliferative response to antigens derived from tumor cells. Indeed, manipulation of the CTLA4 axis was the first demonstration that purely immunoregulatory antibody strategies directed at T cell physiology could be safe and effective in the treatment of cancer, although it acts at a very early stage in T cell activation and can be considered somewhat nonspecific in its basis for T cell stimulation. Pembrolizumab, an anti-PD ligand blocking agent was also approved for melanoma, with a similar spectrum of potential adverse events, but acting in the tumor microenvironment. Indeed, prominent activation of autoimmune hepatic, endocrine, cutaneous, neurologic, and gastrointestinal responses is a basis for adverse events with the use of ipilimumab; the emergent use of glucocorticoids may be required to attenuate severe toxicities, which unfortunately can cause potential attenuation of antitumor effect. Importantly for the general internist, these events may occur late after exposure to ipilimumab while the patient may otherwise be enjoying sustained control of tumor growth owing to the beneficial actions of ipilimumab.

Another class of immunoregulatory antibody is the “bispecific” antibody blinatumomab, which was constructed to have an anti-CD19 antigen combining site as one valency of an antibody with anti-CD3 binding site as the other valency. This antibody thus can bring T cells (with its anti-CD3 activity) close to B cells bearing the CD19 determinant. Blinatumomab is active in B cell neoplasms such as acute

lymphocytic leukemia, which may not have prominent expression of the CD20 targeted by rituximab.

ANTIBODY CONJUGATES Conjugates of antibodies with drugs and isotopes have also been shown to be effective in the treatment of cancer and have the intent of increasing the therapeutic index of the drug or isotope by delivering the toxic “warhead” directly to the tumor cell or tumor microenvironment. Ado-trastuzumab is a conjugate of the HER2/neu-directed trastuzumab and a highly toxic microtubule targeted drug (emtansine), which by itself is too toxic for human use; the antibody-drug conjugate shows valuable activity in patients with breast cancer who have developed resistance to the “naked” antibody. Brentuximab vedotin is an anti-CD30 antibody drug conjugate with a distinct microtubule poison with activity in neoplasms such as Hodgkin’s lymphoma where the tumor cells frequently express CD30. Radioconjugates targeting CD20 on lymphomas have been approved for use (ibratumomab tiuxetan [Zevalin], using yttrium-90 or ¹³¹I-tositumomab). Toxicity concerns have limited their use.

Cytokines There are >70 separate proteins and glycoproteins with biologic effects in humans: interferon (IFN) α , β , and γ ; interleukin (IL) 1 through 29 (so far); the tumor necrosis factor (TNF) family (including lymphotoxin, TNF-related apoptosis-inducing ligand [TRAIL], CD40 ligand, and others); and the chemokine family. Only a fraction of these has been tested against cancer; only IFN- α and IL-2 are in routine clinical use.

About 20 different genes encode IFN- α , and their biologic effects are indistinguishable. IFN induces the expression of many genes, inhibits protein synthesis, and exerts a number of different effects on diverse cellular processes. The two recombinant forms that are commercially available are IFN- α 2a and - α 2b. Interferon is not curative for any tumor but can induce partial responses in follicular lymphoma, hairy cell leukemia, CML, melanoma, and Kaposi’s sarcoma. It has been used in the adjuvant setting in stage II melanoma, multiple myeloma, and follicular lymphoma, with uncertain effects on survival. It produces fever, fatigue, a flulike syndrome, malaise, myelosuppression, and depression and can induce clinically significant autoimmune disease. IFN- α is not generally the treatment of choice for any cancer.

IL-2 exerts its antitumor effects indirectly through augmentation of immune function. Its biologic activity is to promote the growth and activity of T cells and natural killer (NK) cells. High doses of IL-2 can produce tumor regression in certain patients with metastatic melanoma and renal cell cancer. About 2–5% of patients may experience complete remissions that are durable, unlike any other treatment for these tumors. IL-2 is associated with myriad clinical side effects, including intravascular volume depletion, capillary leak syndrome, adult respiratory distress syndrome, hypotension, fever, chills, skin rash, and impaired renal and liver function. Patients may require blood pressure support and intensive care to manage the toxicity. However, once the agent is stopped, most of the toxicities reverse completely within 3–6 days.

Ligand Receptor-Directed Constructs High-affinity receptors for cytokines have led to the design of cytokine-toxin recombinant fusion proteins, such as IL-2 expressed in frame with a fragment of diphtheria toxin. A commercially available construct has activity against certain T cell lymphomas. Likewise, the high-affinity folate receptor is the target for folate conjugated to chemotherapeutic agents. In both cases, the drug’s utility derives from the internalization of the targeted receptor and cleavage of the active drug or toxin moiety.

SYSTEMIC RADIATION THERAPY

Although total-body irradiation has a role in preparing a patient to received allogeneic stem cells, and antibodies as described above can specifically target radioisotopes, systemically administered isotopes of iodide salts have an important role in the treatment of thyroid neoplasms, owing to the selective upregulation of the iodide transporter in the tumor cell compartment. Likewise, isotopes of samarium and radium have been found useful in the palliation of symptoms from

advanced bony metastases of prostate cancer owing to their selective deposition at the tumor–bone matrix interface, thereby potentially affecting the function of both tumor and stromal cells in the progressive growth of the metastatic deposit.

RESISTANCE TO CANCER TREATMENTS

Resistance mechanisms to the conventional cytotoxic agents were initially characterized in the late twentieth century as defects in drug uptake, metabolism, or export by tumor cells. The *multidrug resistance* (*mdr*) gene defined in vitro in cell lines exposed to increasing concentrations of drugs led to the definition of a family of transport proteins that, when overexpressed, result in the facile transport of a variety of hydrophobic drugs out of the cancer cell. Although efforts to manipulate this transporter to promote drug residence in tumor cells have been pursued, none are clinically useful at this time. Drug-metabolizing enzymes such as cytidine deaminase are upregulated in resistant tumor cells, and this is the basis for so-called “high-dose cytarabine” regimens in the treatment of leukemia. Another resistance mechanism defined during this era involved increased expression of a drug’s target, exemplified by amplification of the dihydrofolate reductase gene, in patients who had lost responsiveness to methotrexate, or mutation of topoisomerase II in tumors that relapsed after topoisomerase II modulator treatment.

A second class of resistance mechanisms involves loss of the cellular apoptotic mechanism activated after the engagement of a drug’s target by the drug. This occurs in a way that is heavily influenced by the biology of the particular tumor type. For example, decreased alkylguanine alkyltransferase defines a subset of glioblastoma patients with the prospect of greatest benefit from treatment with temozolomide, but has no predictive value for benefit from temozolomide in epithelial neoplasms. Likewise, ovarian cancers resistant to platinating agents have decreased expression of the proapoptotic gene *bax*. These types of findings have prompted the idea that responsive tumors to chemotherapeutic agents are populated by cells that express drug-related cell death controlling genes, creating in effect a state of “synthetic lethality” of the drug (Chap. 102e) with the genes expressed in responsive tumors, analogous to the existence in yeast of mutations that are well tolerated in the absence of a physiologic stressor but become lethal in the presence of that stressor. In the case of tumors, the chemotherapy inducing the cell death response is the analogous physiologic stressor.

A third class of resistance mechanisms emerged from sequencing of the targets of agents directed at oncogenic kinases. Thus, patients with CML resistant to imatinib have acquired mutations in the ATP binding domain of p210^{bcr-abl} in some cases, leading to the screening and design of agents with activity against the mutant proteins. Entirely analogous resistance mechanisms have emerged in patients with lung cancer treated with the EGFR antagonists gefitinib and erlotinib.

A final category of tumor resistance mechanisms to targeted agents includes the upregulation of alternate means of activating the pathway targeted by the agent. Thus melanomas initially responsive to *BRAF* V600E antagonists such as vemurafenib may reactivate raf signaling by upregulating isoforms that can bypass the variant blocked by the drug. Likewise, inhibition of HER2/neu signaling in breast cancer cells can lead to the emergence of variants with distinct oncogenic signaling pathways such as PI3 kinase. Analogously in NSCLC, EGFR inhibitor treatment leads to the emergence of cells with a predominance of c-met protooncogene–dependent signaling in the resistant tumors.

The susceptibility of a tumor to different treatments as a function of its expression of potential drug targets or their mutational profile has led to efforts to define the dominant pathways driving a patient’s tumor by genomic techniques including whole exome sequencing. The difficulty with applying such data to patient treatment is recognizing that these pathways may change during the natural history of a tumor and that different sites in a single patient may have tumors with different patterns of gene mutation.

MYELOSUPPRESSION

The common cytotoxic chemotherapeutic agents almost invariably affect bone marrow function. Titration of this effect determines the MTD of the agent on a given schedule. The normal kinetics of blood cell turnover influences the sequence and sensitivity of each of the formed elements. Polymorphonuclear leukocytes (PMNs; $t_{1/2}$ = 6–8 h), platelets ($t_{1/2}$ = 5–7 days), and red blood cells (RBCs; $t_{1/2}$ = 120 days) have most, less, and least susceptibility, respectively, to usually administered cytotoxic agents. The nadir count of each cell type in response to classes of agents is characteristic. Maximal neutropenia occurs 6–14 days after conventional doses of anthracyclines, antifolates, and antimetabolites. Alkylating agents differ from each other in the timing of cytopenias. Nitrosoureas, DTIC, and procarbazine can display delayed marrow toxicity, first appearing 6 weeks after dosing.

Complications of myelosuppression result from the predictable sequelae of the missing cells’ function. *Febrile neutropenia* refers to the clinical presentation of fever (one temperature $\geq 38.5^\circ\text{C}$ or three readings $\geq 38^\circ\text{C}$ but $\leq 38.5^\circ\text{C}$ per 24 h) in a neutropenic patient with an uncontrolled neoplasm involving the bone marrow or, more usually, in a patient undergoing treatment with cytotoxic agents. Mortality from uncontrolled infection varies inversely with the neutrophil count. If the nadir neutrophil count is $>1000/\mu\text{L}$, there is little risk; if $<500/\mu\text{L}$, risk of death is markedly increased. Management of febrile neutropenia has conventionally included empirical coverage with antibiotics for the duration of neutropenia (Chap. 104). Selection of antibiotics is governed by the expected association of infections with certain underlying neoplasms; careful physical examination (with scrutiny of catheter sites, dentition, mucosal surfaces, and perirectal and genital orifices by gentle palpation); chest x-ray; and Gram stain and culture of blood, urine, and sputum (if any) to define a putative site of infection. In the absence of any originating site, a broadly acting β -lactam with anti-*Pseudomonas* activity, such as ceftazidime, is begun empirically. The addition of vancomycin to cover potential cutaneous sites of origin (until these are ruled out or shown to originate from methicillin-sensitive organisms) or metronidazole or imipenem for abdominal or other sites favoring anaerobes reflects modifications tailored to individual patient presentations. The coexistence of pulmonary compromise raises a distinct set of potential pathogens, including *Legionella*, *Pneumocystis*, and fungal agents that may require further diagnostic evaluations, such as bronchoscopy with bronchoalveolar lavage. Febrile neutropenic patients can be stratified broadly into two prognostic groups. The first, with expected short duration of neutropenia and no evidence of hypotension or abdominal or other localizing symptoms, may be expected to do well even with oral regimens, e.g., ciprofloxacin or moxifloxacin, or amoxicillin plus clavulanic acid. A less favorable prognostic group is patients with expected prolonged neutropenia, evidence of sepsis, and end organ compromise, particularly pneumonia. These patients require tailoring of their antibiotic regimen to their underlying presentation, with frequent empirical addition of antifungal agents if fever and neutropenia persists for 7 days without identification of an adequately treated organism or site.

Transfusion of granulocytes has no role in the management of febrile neutropenia, owing to their exceedingly short half-life, mechanical fragility, and clinical syndromes of pulmonary compromise with leukostasis after their use. Instead, colony-stimulating factors (CSFs) are used to augment bone marrow production of PMNs. Early-acting factors such as IL-1, IL-3, and stem cell factor have not been as useful clinically as late-acting, lineage-specific factors such as granulocyte colony-stimulating factor (G-CSF) or GM-CSF, erythropoietin (EPO), thrombopoietin, IL-6, and IL-11. CSFs may easily become overused in oncology practice. The settings in which their use has been proved effective are limited. G-CSF, GM-CSF, EPO, and IL-11 are currently approved for use. The American Society of Clinical Oncology has developed practice guidelines for the use of G-CSF and GM-CSF (Table 103e-7).

Primary prophylaxis (i.e., shortly after completing chemotherapy to reduce the nadir) administers G-CSF to patients receiving cytotoxic

TABLE 103e-7 INDICATIONS FOR THE CLINICAL USE OF G-CSF OR GM-CSF**Preventive Uses**

With the first cycle of chemotherapy (so-called *primary CSF administration*)

- Not needed on a routine basis
- Use if the probability of febrile neutropenia is $\geq 20\%$
- Use if patient has preexisting neutropenia or active infection
- Age >65 years treated for lymphoma with curative intent or other tumor treated by similar regimens
- Poor performance status
- Extensive prior chemotherapy
- Dose-dense regimens in a clinical trial or with strong evidence of benefit

With subsequent cycles if febrile neutropenia has previously occurred (so-called *secondary CSF administration*)

- Not needed after short-duration neutropenia without fever
- Use if patient had febrile neutropenia in previous cycle
- Use if prolonged neutropenia (even without fever) delays therapy

Therapeutic Uses

Afebrile neutropenic patients

- No evidence of benefit

Febrile neutropenic patients

- No evidence of benefit
- May feel compelled to use in the face of clinical deterioration from sepsis, pneumonia, or fungal infection, but benefit unclear

In bone marrow or peripheral blood stem cell transplantation

- Use to mobilize stem cells from marrow
- Use to hasten myeloid recovery

In acute myeloid leukemia

- G-CSF of minor or no benefit
- GM-CSF of no benefit and may be harmful

In myelodysplastic syndromes

- Not routinely beneficial
- Use intermittently in subset with neutropenia and recurrent infection

What Dose and Schedule Should Be Used?

G-CSF: 5 mg/kg per day subcutaneously

GM-CSF: 250 mg/m² per day subcutaneously

Pegfilgrastim: one dose of 6 mg 24 h after chemotherapy

When Should Therapy Begin and End?

When indicated, start 24–72 h after chemotherapy

Continue until absolute neutrophil count is 10,000/ μ L

Do not use concurrently with chemotherapy or radiation therapy

Abbreviations: CSF, cerebrospinal fluid; G-CSF, granulocyte colony-stimulating factor; GM-CSF, granulocyte-macrophage colony-stimulating factor.

Source: From the American Society of Clinical Oncology: *J Clin Oncol* 24:3187, 2006.

regimens associated with a 20% incidence of febrile neutropenia. “Dose-dense” regimens, where cycling of chemotherapy is intended to be completed without delay of administered doses, may also benefit, but such patients should be on a clinical trial. Administration of G-CSF in these circumstances has reduced the incidence of febrile neutropenia in several studies by about 50%. Most patients, however, receive regimens that do not have such a high risk of expected febrile neutropenia, and therefore most patients initially should not receive G-CSF or GM-CSF. Special circumstances—such as a documented history of febrile neutropenia with the regimen in a particular patient or categories of patients at increased risk, such as patients older than age 65 years with aggressive lymphoma treated with curative chemotherapy regimens; extensive compromise of marrow by prior radiation or chemotherapy; or active, open wounds or deep-seated infection—may support primary treatment with G-CSF or GM-CSF. Administration of G-CSF or GM-CSF to afebrile neutropenic patients or to patients with low-risk febrile neutropenia is not recommended, and patients receiving concomitant chemoradiation treatment, particularly those with thoracic neoplasms, likewise are not generally recommended for

treatment. In contrast, administration of G-CSF to high-risk patients with febrile neutropenia and evidence of organ compromise including sepsis syndrome, invasive fungal infection, concurrent hospitalization at the time fever develops, pneumonia, profound neutropenia ($<0.1 \times 10^9/L$), or age >65 years is reasonable.

Secondary prophylaxis refers to the administration of CSFs in patients who have experienced a neutropenic complication from a prior cycle of chemotherapy; dose reduction or delay may be a reasonably considered alternative. G-CSF or GM-CSF is conventionally started 24–72 h after completion of chemotherapy and continued until a PMN count of 10,000/ μ L is achieved, unless a “depot” preparation of G-CSF such as pegfilgrastim is used, where one dose is administered at least 14 days before the next scheduled administration of chemotherapy. Also, patients with myeloid leukemias undergoing induction therapy may have a slight reduction in the duration of neutropenia if G-CSF is commenced after completion of therapy and may be of particular value in elderly patients, but the influence on long-term outcome has not been defined. GM-CSF probably has a more restricted utility than G-CSF, with its use currently limited to patients after autologous bone marrow transplants, although proper head-to-head comparisons with G-CSF have not been conducted in most instances. GM-CSF may be associated with more systemic side effects.

Dangerous degrees of thrombocytopenia do not frequently complicate the management of patients with solid tumors receiving cytotoxic chemotherapy (with the possible exception of certain carboplatin-containing regimens), but they are frequent in patients with certain hematologic neoplasms where marrow is infiltrated with tumor. Severe bleeding related to thrombocytopenia occurs with increased frequency at platelet counts $<20,000/\mu$ L and is very prevalent at counts $<5000/\mu$ L.

The precise “trigger” point at which to transfuse patients has been defined as a platelet count of 10,000/ μ L or less in patients without medical comorbidities that may increase the risk of bleeding. This issue is important not only because of the costs of frequent transfusion, but unnecessary platelet transfusions expose the patient to the risks of allosensitization and loss of value from subsequent transfusion owing to rapid platelet clearance, as well as the infectious and hypersensitivity risks inherent in any transfusion. Prophylactic transfusions to keep platelets $>20,000/\mu$ L are reasonable in patients with leukemia who are stressed by fever or concomitant medical conditions (the threshold for transfusion is 10,000/ μ L in patients with solid tumors and no other bleeding diathesis or physiologic stressors such as fever or hypotension, a level that might also be reasonably considered for leukemia patients who are thrombocytopenic but not stressed or bleeding). In contrast, patients with myeloproliferative states may have functionally altered platelets despite normal platelet counts, and transfusion with normal donor platelets should be considered for evidence of bleeding in these patients. Careful review of medication lists to prevent exposure to nonsteroidal anti-inflammatory agents and maintenance of clotting factor levels adequate to support near-normal prothrombin and partial thromboplastin time tests are important in minimizing the risk of bleeding in the thrombocytopenic patient.

Certain cytokines in clinical investigation have shown an ability to increase platelets (e.g., IL-6, IL-1, thrombopoietin), but clinical benefit and safety are not yet proven. IL-11 (oprelvekin) is approved for use in the setting of expected thrombocytopenia, but its effects on platelet counts are small, and it is associated with side effects such as headache, fever, malaise, syncope, cardiac arrhythmias, and fluid retention. Eltrombopag and romiplostim are thrombopoietin agonists with demonstrated efficacy in certain thrombocytopenic states, but they have not been systematically studied in chemotherapy-induced thrombocytopenia.

Anemia associated with chemotherapy can be managed by transfusion of packed RBCs. Transfusion is not undertaken until the hemoglobin falls to <80 g/L (8 g/dL), compromise of end organ function occurs, or an underlying condition (e.g., coronary artery disease) calls for maintenance of hemoglobin >90 g/L (9 g/dL). Patients who are to receive therapy for >2 months on a “stable” regimen and who are likely to require continuing transfusions are also candidates for erythropoietin (EPO). Randomized trials in certain tumors have raised the pos-

sibility that EPO use may promote tumor-related adverse events. This information should be considered in the care of individual patients. In the event EPO treatment is undertaken, maintenance of hemoglobin of 90–100 g/L (9–10 g/dL) should be the target. In the setting of adequate iron stores and serum EPO levels <100 ng/mL, EPO, 150 U three times a week, can produce a slow increase in hemoglobin over about 2 months of administration. Depot formulations can be administered less frequently. It is unclear whether higher hemoglobin levels, up to 110–120 g/L (11–12 g/dL), are associated with improved quality of life to a degree that justifies the more intensive EPO use. Efforts to achieve levels at or above 120 g/L (12 g/dL) have been associated with increased thromboses and mortality rates. EPO may rescue hypoxic cells from death and contribute to tumor radioresistance.

NAUSEA AND VOMITING

The most common side effect of chemotherapy administration is nausea, with or without vomiting. Nausea may be acute (within 24 h of chemotherapy), delayed (>24 h), or anticipatory of the receipt of chemotherapy. Patients may be likewise stratified for their risk of susceptibility to nausea and vomiting, with increased risk in young, female, heavily pretreated patients without a history of alcohol or drug use but with a history of motion or morning sickness. Antineoplastic agents vary in their capacity to cause nausea and vomiting. Highly emetogenic drugs (>90%) include mechlorethamine, streptozotocin, DTIC, cyclophosphamide at >1500 mg/m², and cisplatin; moderately emetogenic drugs (30–90% risk) include carboplatin, cytosine arabinoside (>1 mg/m²), ifosfamide, conventional-dose cyclophosphamide, and anthracyclines; low-risk (10–30%) agents include 5FU, taxanes, etoposide, and bortezomib, with minimal risk (<10%) afforded by treatment with antibodies, bleomycin, busulfan, fludarabine, and vinca alkaloids. *Emesis* is a reflex caused by stimulation of the vomiting center in the medulla. Input to the vomiting center comes from the chemoreceptor trigger zone (CTZ) and afferents from the peripheral gastrointestinal tract, cerebral cortex, and heart. The different emesis “syndromes” require distinct management approaches. In addition, a conditioned reflex may contribute to anticipatory nausea arising after repeated cycles of chemotherapy. Accordingly, antiemetic agents differ in their locus and timing of action. Combining agents from different classes or the sequential use of different classes of agent is the cornerstone of successful management of chemotherapy-induced nausea and vomiting. Of great importance are the prophylactic administration of agents and such psychological techniques as the maintenance of a supportive milieu, counseling, and relaxation to augment the action of antiemetic agents.

Serotonin antagonists (5-HT₃) and neurokinin 1 (NK1) receptor antagonists are useful in “high-risk” chemotherapy regimens. The combination acts at both peripheral gastrointestinal and CNS sites that control nausea and vomiting. For example, the 5-HT₃ blocker dolasetron, 100 mg intravenously or orally; dexamethasone, 12 mg; and the NK1 antagonist aprepitant, 125 mg orally, are combined on the day of administration of severely emetogenic regimens, with repetition of dexamethasone (8 mg) and aprepitant (80 mg) on days 2 and 3 for delayed nausea. Alternate 5-HT₃ antagonists include ondansetron, given as 0.15 mg/kg intravenously for three doses just before and at 4 and 8 h after chemotherapy; palonosetron at 0.25 mg over 30 s, 30 min before chemotherapy; and granisetron, given as a single dose of 0.01 mg/kg just before chemotherapy. Emesis from moderately emetic chemotherapy regimens may be prevented with a 5-HT₃ antagonist and dexamethasone alone for patients not receiving doxorubicin and cyclophosphamide combinations; the latter combination requires the 5-HT₃/dexamethasone/aprepitant on day 1 but aprepitant alone on days 2 and 3. Emesis from low-emetic-risk regimens may be prevented with 8 mg of dexamethasone alone or with non-5-HT₃, non-NK1 antagonist approaches including the following.

Antidopaminergic phenothiazines act directly at the CTZ and include prochlorperazine (Compazine), 10 mg intramuscularly or intravenously, 10–25 mg orally, or 25 mg per rectum every 4–6 h for up to four doses; and thiethylperazine, 10 mg by potentially all of the above routes every 6 h. Haloperidol is a butyrophenone

dopamine antagonist given at 1 mg intramuscularly or orally every 8 h. Antihistamines such as diphenhydramine have little intrinsic antiemetic capacity but are frequently given to prevent or treat dystonic reactions that can complicate use of the antidopaminergic agents. Lorazepam is a short-acting benzodiazepine that provides an anxiolytic effect to augment the effectiveness of a variety of agents when used at 1–2 mg intramuscularly, intravenously, or orally every 4–6 h. Metoclopramide acts on peripheral dopamine receptors to augment gastric emptying and is used in high doses for highly emetogenic regimens (1–2 mg/kg intravenously 30 min before chemotherapy and every 2 h for up to three additional doses as needed); intravenous doses of 10–20 mg every 4–6 h as needed or 50 mg orally 4 h before and 8 and 12 h after chemotherapy are used for moderately emetogenic regimens. 5-9-Tetrahydrocannabinol (Marinol) is a rather weak antiemetic compared to other available agents, but it may be useful for persisting nausea and is used orally at 10 mg every 3–4 h as needed.

DIARRHEA

Regimens that include 5FU infusions and/or irinotecan may produce severe diarrhea. Similar to the vomiting syndromes, chemotherapy-induced diarrhea may be immediate or occur in a delayed fashion up to 48–72 h after the drugs. Careful attention to maintained hydration and electrolyte repletion, intravenously if necessary, along with antimotility treatments such as “high-dose” loperamide, commenced with 4 mg at the first occurrence of diarrhea, with 2 mg repeated every 2 h until 12 h without loose stools, not to exceed a total daily dose of 16 mg. Octreotide (100–150 µg), a somatostatin analogue, or opiate-based preparations may be considered for patients not responding to loperamide.

MUCOSITIS

Irritation and inflammation of the mucous membranes particularly afflicting the oral and anal mucosa, but potentially involving the gastrointestinal tract, may accompany cytotoxic chemotherapy. Mucositis is due to damage to the proliferating cells at the base of the mucosal squamous epithelia or in the intestinal crypts. Topical therapies, including anesthetics and barrier-creating preparations, may provide symptomatic relief in mild cases. Palifermin or keratinocyte growth factor, a member of the fibroblast growth factor family, is effective in preventing severe mucositis in the setting of high-dose chemotherapy with stem cell transplantation for hematologic malignancies. It may also prevent or ameliorate mucositis from radiation.

ALOPECIA

Chemotherapeutic agents vary widely in causing alopecia, with anthracyclines, alkylating agents, and topoisomerase inhibitors reliably causing near-total alopecia when given at therapeutic doses. Antimetabolites are more variably associated with alopecia. Psychological support and the use of cosmetic resources are to be encouraged, and “chemo caps” that reduce scalp temperature to decrease the degree of alopecia should be discouraged, particularly during treatment with curative intent of neoplasms, such as leukemia or lymphoma, or in adjuvant breast cancer therapy. The richly vascularized scalp can certainly harbor micrometastatic or disseminated disease.

GONADAL DYSFUNCTION AND PREGNANCY

Cessation of ovulation and azoospermia reliably result from alkylating agent- and topoisomerase poison-containing regimens. The duration of these effects varies with age and sex. Males treated for Hodgkin's disease with mechlorethamine- and procarbazine-containing regimens are effectively sterile, whereas fertility usually returns after regimens that include cisplatin, vinblastine, or etoposide and after bleomycin for testicular cancer. Sperm banking before treatment may be considered to support patients likely to be sterilized by treatment. Females experience amenorrhea with anovulation after alkylating agent therapy; they are likely to recover normal menses if treatment is completed before age 30 but unlikely to recover menses after age 35. Even those who regain menses usually experience premature

103e-26 menopause. Because the magnitude and extent of decreased fertility can be difficult to predict, patients should be counseled to maintain effective contraception, preferably by barrier means, during and after therapy. Resumption of efforts to conceive should be considered in the context of the patient's likely prognosis. Hormone replacement therapy should be undertaken in women who do not have a hormonally responsive tumor. For patients who have had a hormone-sensitive tumor primarily treated by a local modality, conventional practice would counsel against hormone replacement, but this issue is under investigation.

Chemotherapy agents have variable effects on the success of pregnancy. All agents tend to have increased risk of adverse outcomes when administered during the first trimester, and strategies to delay chemotherapy, if possible, until after this milestone should be considered if the pregnancy is to continue to term. Patients in their second or third trimester can be treated with most regimens for the common neoplasms afflicting women in their childbearing years, with the exception of antimetabolites, particularly antifolates, which have notable teratogenic or fetotoxic effects throughout pregnancy. The need for anticancer chemotherapy per se is infrequently a clear basis to recommend termination of a concurrent pregnancy, although each treatment strategy in this circumstance must be tailored to the individual needs of the patient.

SPECIAL ISSUES WITH TARGETED TREATMENTS

Treatment with EGFR-directed small molecules (e.g., erlotinib, afatinib, lapatinib), antibodies (e.g., cetuximab, panitumumab), and mTOR antagonists (e.g., everolimus, temsirolimus) reliably produces an acneiform rash that can be a source of distress to patients and can be ameliorated with topically applied clindamycin gels and low-potency corticosteroid creams. Diarrhea frequently accompanies tyrosine

kinase inhibitor administration and may respond to antimotility agents such as loperamide or stool-bulking agents.

Anti-VEGFR-directed treatments, including the specific antibody bevacizumab, and the "multikinase" inhibitors with anti VEGFR activity, such as sorafenib, sunitinib, and pazopanib, reliably produce hypertension in a significant fraction of patients that typically can be treated with lisinopril, amlodipine, or clonidine alone or in combination. More difficult to treat is proteinuria with resultant azotemia; this can be a basis for discontinuing treatment depending on the clinical context. Thyroid function is prominently affected by chronic exposure to this group of multikinase inhibitors including sorafenib and pazopanib, and periodic surveillance of thyroid-stimulating hormone and thyroxine (T_4) levels during treatment is reasonable. Gastrointestinal perforations, arterial thromboses, and hemorrhage likewise have no specific treatments and may be a basis to avoid this class of agents. Palmar-plantar dysesthesia ("hand-foot syndrome") can be seen after administration of these agents (as well as some cytotoxic agents including gemcitabine and liposomal preparations of doxorubicin) and is a basis for considering dose reduction if not responsive to topical emollients and analgesics.

Protein kinase antagonists as a class have been associated with poorly predicted hepatic and cardiac toxicities (imatinib, dasatinib, sorafenib, pazopanib) or cardiac conduction deficits including prolonged QT interval (pazopanib). The occurrence of new cardiac or liver abnormalities in a patient receiving treatment with a protein kinase antagonist should lead to a consideration of the risk versus benefit and the possible relation of the agent to the new adverse event. The existence of prior cardiac dysfunction is a relative contraindication to the use of certain targeted therapies (e.g., trastuzumab), although each patient's needs should be individualized. **Chronic effects of cancer treatment are reviewed in Chap. 125.**

104 Infections in Patients with Cancer

Robert W. Finberg

Infections are a common cause of death and an even more common cause of morbidity in patients with a wide variety of neoplasms. Autopsy studies show that most deaths from acute leukemia and half of deaths from lymphoma are caused directly by infection. With more intensive chemotherapy, patients with solid tumors have also become more likely to die of infection. Fortunately, an evolving approach to prevention and treatment of infectious complications of cancer has decreased infection-associated mortality rates and will probably continue to do so. This accomplishment has resulted from three major steps:

1. The practice of using “early empirical” antibiotics reduced mortality rates among patients with leukemia and bacteremia from 84% in 1965 to 44% in 1972. Recent studies suggest that the mortality rate due to infection in febrile neutropenic patients dropped to <10% by 2013. This dramatic improvement is attributed to early intervention with appropriate antimicrobial therapy.
2. “Empirical” antifungal therapy has also lowered the incidence of disseminated fungal infection, with dramatic decreases in mortality rates. An antifungal agent is administered—on the basis of likely fungal infection—to neutropenic patients who, after 4–7 days of antibiotic therapy, remain febrile but have no positive cultures.
3. Use of antibiotics for afebrile neutropenic patients as broad-spectrum prophylaxis against infections has decreased both mortality and morbidity even further. The current approach to treatment of severely neutropenic patients (e.g., those receiving high-dose chemotherapy for leukemia or high-grade lymphoma) is based on initial prophylactic therapy at the onset of neutropenia, subsequent “empirical” antibacterial therapy targeting the organisms whose involvement is likely in light of physical findings (most often fever alone), and finally “empirical” antifungal therapy based on the known likelihood that fungal infection will become a serious issue after 4–7 days of broad-spectrum antibacterial therapy.

A physical predisposition to infection in patients with cancer (Table 104-1) can be a result of the neoplasm’s production of a break in the skin. For example, a squamous cell carcinoma may cause local invasion

of the epidermis, which allows bacteria to gain access to subcutaneous tissue and permits the development of cellulitis. The artificial closing of a normally patent orifice can also predispose to infection; for example, obstruction of a ureter by a tumor can cause urinary tract infection, and obstruction of the bile duct can cause cholangitis. Part of the host’s normal defense against infection depends on the continuous emptying of a viscus; without emptying, a few bacteria that are present as a result of bacteremia or local transit can multiply and cause disease.

A similar problem can affect patients whose lymph node integrity has been disrupted by radical surgery, particularly patients who have had radical node dissections. A common clinical problem following radical mastectomy is the development of cellulitis (usually caused by streptococci or staphylococci) because of lymphedema and/or inadequate lymph drainage. In most cases, this problem can be addressed by local measures designed to prevent fluid accumulation and breaks in the skin, but antibiotic prophylaxis has been necessary in refractory cases.

A life-threatening problem common to many cancer patients is the loss of the reticuloendothelial capacity to clear microorganisms after splenectomy, which may be performed as part of the management of hairy cell leukemia, chronic lymphocytic leukemia (CLL), and chronic myelogenous leukemia (CML) and in Hodgkin’s disease. Even after curative therapy for the underlying disease, the lack of a spleen predisposes such patients to rapidly fatal infections. The loss of the spleen through trauma similarly predisposes the normal host to overwhelming infection throughout life. The splenectomized patient should be counseled about the risks of infection with certain organisms, such as the protozoan *Babesia* (Chap. 249) and *Capnocytophaga canimorsus*, a bacterium carried in the mouths of animals (Chaps. 167e and 183e). Because encapsulated bacteria (*Streptococcus pneumoniae*, *Haemophilus influenzae*, and *Neisseria meningitidis*) are the organisms most commonly associated with postsplenectomy sepsis, splenectomized persons should be vaccinated (and revaccinated; Table 104-2 and Chap. 148) against the capsular polysaccharides of these organisms. Many clinicians recommend giving splenectomized patients a small supply of antibiotics effective against *S. pneumoniae*, *N. meningitidis*, and *H. influenzae* to avert rapid, overwhelming sepsis in the event that they cannot present for medical attention immediately after the onset of fever or other signs or symptoms of bacterial infection. A few tablets of amoxicillin/clavulanic acid (or levofloxacin if resistant strains of *S. pneumoniae* are prevalent locally) are a reasonable choice for this purpose.

TABLE 104-1 DISRUPTION OF NORMAL BARRIERS THAT MAY PREDISPOSE TO INFECTIONS IN PATIENTS WITH CANCER

Type of Defense	Specific Lesion	Cells Involved	Organism	Cancer Association	Disease
Physical barrier	Breaks in skin	Skin epithelial cells	Staphylococci, streptococci	Head and neck, squamous cell carcinoma	Cellulitis, extensive skin infection
Emptying of fluid collections	Occlusion of orifices: ureters, bile duct, colon	Luminal epithelial cells	Gram-negative bacilli	Renal, ovarian, biliary tree, metastatic diseases of many cancers	Rapid, overwhelming bacteremia; urinary tract infection
Lymphatic function	Node dissection	Lymph nodes	Staphylococci, streptococci	Breast cancer surgery	Cellulitis
Splenic clearance of microorganisms	Splenectomy	Splenic reticuloendothelial cells	<i>Streptococcus pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Neisseria meningitidis</i> , <i>Babesia</i> , <i>Capnocytophaga canimorsus</i>	Hodgkin’s disease, leukemia	Rapid, overwhelming sepsis
Phagocytosis	Lack of granulocytes	Granulocytes (neutrophils)	Staphylococci, streptococci, enteric organisms, fungi	Acute myeloid and acute lymphocytic leukemias, hairy cell leukemia	Bacteremia
Humoral immunity	Lack of antibody	B cells	<i>S. pneumoniae</i> , <i>H. influenzae</i> , <i>N. meningitidis</i>	Chronic lymphocytic leukemia, multiple myeloma	Infections with encapsulated organisms, sinusitis, pneumonia
Cellular immunity	Lack of T cells	T cells and macrophages	<i>Mycobacterium tuberculosis</i> , <i>Listeria</i> , herpesviruses, fungi, intracellular parasites	Hodgkin’s disease, leukemia, T cell lymphoma	Infections with intracellular bacteria, fungi, parasites; virus reactivation

TABLE 104-2 VACCINATION OF CANCER PATIENTS RECEIVING CHEMOTHERAPY^a

Vaccine	Use in Indicated Patients		
	Intensive Chemotherapy	Hodgkin's Disease	Hematopoietic Stem Cell Transplantation
Diphtheria-tetanus ^b	Primary series and boosters as necessary	No special recommendation	3 doses given 6–12 months after transplantation
Poliomyelitis ^c	Complete primary series and boosters	No special recommendation	3 doses given 6–12 months after transplantation
<i>Haemophilus influenzae</i> type b conjugate	Primary series and booster for children	Single dose for adults	3 doses given 6–12 months after transplantation (separated by 1 month)
Human papillomavirus (HPV)	Quadrivalent HPV vaccine is approved for males and females 9–26 years of age. Check Centers for Disease Control and Prevention (CDC) website (www.cdc.gov/vaccines) for updated recommendations.	Quadrivalent HPV vaccine is approved for males and females 9–26 years of age. Check CDC website (www.cdc.gov/vaccines) for updated recommendations.	Quadrivalent HPV vaccine is approved for males and females 9–26 years of age. Check CDC website (www.cdc.gov/vaccines) for updated recommendations.
Hepatitis A	As indicated for normal hosts on the basis of occupation and lifestyle	As indicated for normal hosts on the basis of occupation and lifestyle	As indicated for normal hosts on the basis of occupation and lifestyle
Hepatitis B	Same as for normal hosts	As indicated for normal hosts on the basis of occupation and lifestyle	3 doses given 6–12 months after transplantation
Pneumococcal conjugate vaccine (PCV13) Pneumococcal polysaccharide vaccine (PPSV23) ^d	Finish series prior to chemotherapy if possible	Patients with splenectomy should receive PPSV23.	Three doses of PCV13, beginning 3–6 months after transplantation, are followed by a dose of PPSV23 at least 8 weeks later. A second PPSV23 dose can be given 5 years later.
Quadrivalent meningococcal vaccine ^e	Should be administered to splenectomized patients and to patients living in endemic areas, including college students in dormitories	Should be administered to splenectomized patients and to patients living in endemic areas, including college students in dormitories. An additional dose can be given after 5 years.	Should be administered to splenectomized patients and to patients living in endemic areas, including college students in dormitories. An additional dose can be given after 5 years.
Influenza	Seasonal immunization	Seasonal immunization	Seasonal immunization (A seasonal dose is recommended and can be given as early as 4 months after transplantation; if given <6 months after transplantation, an additional dose is recommended.)
Measles/mumps/rubella	Contraindicated	Contraindicated during chemotherapy	After 24 months in patients without graft-versus-host disease
Varicella-zoster virus ^f	Contraindicated ^g	Contraindicated	Contraindicated (CDC recommends use on a case-by-case basis following reevaluation.)

^aThe latest recommendations by the Advisory Committee on Immunization Practices and the CDC guidelines can be found at <http://www.cdc.gov/vaccines>. ^bA single dose of TdAp (tetanus–diphtheria–acellular pertussis), followed by a booster dose of Td (tetanus–diphtheria) every 10 years, is recommended for adults. ^cLive-virus vaccine is contraindicated; inactivated vaccine should be used. ^dTwo types of vaccine are used to prevent pneumococcal disease. A conjugate vaccine active against 13 serotypes (13-valent pneumococcal conjugate vaccine, or PCV13) is currently administered in three separate doses to all children. A polysaccharide vaccine active against 23 serotypes (23-valent pneumococcal polysaccharide vaccine, or PPSV23) elicits titers of antibody lower than those achieved with the conjugate vaccine, and immunity may wane more rapidly. Because the ablative chemotherapy given to recipients of hematopoietic stem cell transplants (HSCTs) eradicates immunologic memory, revaccination is recommended for all such patients. Vaccination is much more effective once immunologic reconstitution has occurred; however, because of the need to prevent serious disease, pneumococcal vaccine should be administered 6–12 months after transplantation in most cases. Because PPSV23 includes serotypes not present in PCV13, HSCT recipients should receive a dose of PPSV23 at least 8 weeks after the last dose of PCV13. Although antibody titers from PPSV23 clearly decay, experience with multiple doses of PPSV23 is limited, as are data on the safety, toxicity, or efficacy of such a regimen. For this reason, the CDC currently recommends the administration of one additional dose of PPSV23 at least 5 years after the last dose to immunocompromised patients, including transplant recipients, as well as patients with Hodgkin's disease, multiple myeloma, lymphoma, or generalized malignancies. Beyond this single additional dose, further doses are not recommended at this time.

^eMeningococcal conjugate vaccine MenACWY is recommended for adults ≤55 years old, and meningococcal polysaccharide vaccine (MPSV4) is recommended for those ≥56 years old.

^fIncludes both varicella vaccine for children and zoster vaccine for adults. ^gContact the manufacturer for more information on use in children with acute lymphocytic leukemia.

The level of suspicion of infections with certain organisms should depend on the type of cancer diagnosed (Table 104-3). Diagnosis of multiple myeloma or CLL should alert the clinician to the possibility of hypogammaglobulinemia. While immunoglobulin replacement therapy can be effective, in most cases prophylactic antibiotics are a cheaper, more convenient method of eliminating bacterial infections in CLL patients with hypogammaglobulinemia. Patients with acute lymphocytic leukemia (ALL), patients with non-Hodgkin's lymphoma, and all cancer patients treated with high-dose glucocorticoids (or glucocorticoid-containing chemotherapy regimens) should receive antibiotic prophylaxis for *Pneumocystis* infection (Table 104-3) for the duration of their chemotherapy. In addition to exhibiting susceptibility to certain infectious organisms, patients with cancer are likely to manifest their infections in characteristic ways. For example, fever—generally a sign of infection in normal hosts—continues to be a reliable indicator in neutropenic patients. In contrast, patients receiving glucocorticoids and agents that impair T cell function and cytokine secretion may have serious infections in the absence of fever. Similarly, neutropenic

patients commonly present with cellulitis without purulence and with pneumonia without sputum or even x-ray findings (see below).

The use of monoclonal antibodies that target B and T cells as well as drugs that interfere with lymphocyte signal transduction events is associated with reactivation of latent infections. The use of rituximab, the antibody to CD20 (a B cell surface protein), is associated with the development of reactivation tuberculosis as well as other latent viral infections, including hepatitis B and cytomegalovirus (CMV) infection. Like organ transplant recipients (Chap. 169), patients with latent bacterial disease (like tuberculosis) and latent viral disease (like herpes simplex or zoster) should be carefully monitored for reactivation disease.

SYSTEM-SPECIFIC SYNDROMES

SKIN-SPECIFIC SYNDROMES

Skin lesions are common in cancer patients, and the appearance of these lesions may permit the diagnosis of systemic bacterial or fungal infection. While cellulitis caused by skin organisms such as

TABLE 104-3 INFECTIONS ASSOCIATED WITH SPECIFIC TYPES OF CANCER

Cancer	Underlying Immune Abnormality	Organisms Causing Infection
Multiple myeloma	Hypogammaglobulinemia	<i>Streptococcus pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Neisseria meningitidis</i>
Chronic lymphocytic leukemia	Hypogammaglobulinemia	<i>S. pneumoniae</i> , <i>H. influenzae</i> , <i>N. meningitidis</i>
Acute myeloid or lymphocytic leukemia	Granulocytopenia, skin and mucous membrane lesions	Extracellular gram-positive and gram-negative bacteria, fungi
Hodgkin's disease	Abnormal T cell function	Intracellular pathogens (<i>Mycobacterium tuberculosis</i> , <i>Listeria</i> , <i>Salmonella</i> , <i>Cryptococcus</i> , <i>Mycobacterium avium</i>); herpesviruses
Non-Hodgkin's lymphoma and acute lymphocytic leukemia	Glucocorticoid chemotherapy, T and B cell dysfunction	<i>Pneumocystis</i>
Colon and rectal tumors	Local abnormalities ^a	<i>Streptococcus bovis</i> biotype 1 (bacteremia)
Hairy cell leukemia	Abnormal T cell function	Intracellular pathogens (<i>M. tuberculosis</i> , <i>Listeria</i> , <i>Cryptococcus</i> , <i>M. avium</i>)

^aThe reason for this association is not well defined.

Streptococcus or *Staphylococcus* is common, neutropenic patients—i.e., those with <500 functional polymorphonuclear leukocytes (PMNs)/ μ L—and patients with impaired blood or lymphatic drainage may develop infections with unusual organisms. Innocent-looking macules or papules may be the first sign of bacterial or fungal sepsis in immunocompromised patients (Fig. 104-1). In the neutropenic host, a macule progresses rapidly to ecthyma gangrenosum (see Fig. 25e-35), a usually painless, round, necrotic lesion consisting of a central black or gray-black eschar with surrounding erythema. Ecthyma gangrenosum, which is located in nonpressure areas (as distinguished from necrotic lesions associated with lack of circulation), is often associated with *Pseudomonas aeruginosa* bacteremia (Chap. 189) but may be caused by other bacteria.

Candidemia (Chap. 240) is also associated with a variety of skin conditions (see Fig. 25e-38) and commonly presents as a maculopapular rash. Punch biopsy of the skin may be the best method for diagnosis.

TABLE 104-4 ORGANISMS LIKELY TO CAUSE INFECTIONS IN GRANULOCYTOPENIC PATIENTS

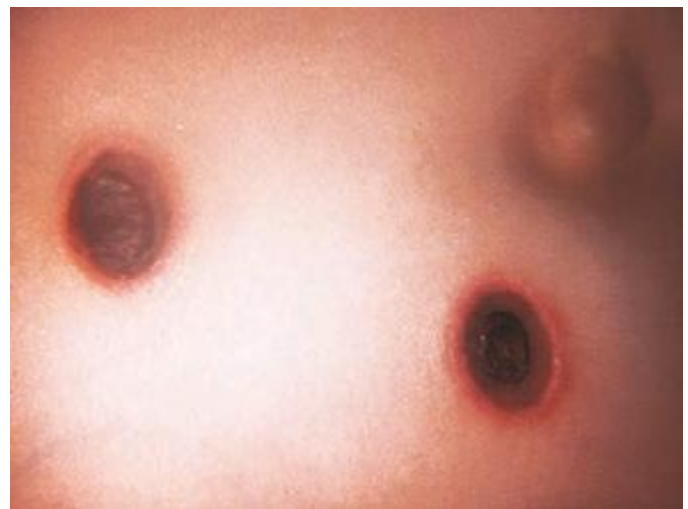
Gram-Positive Cocci	
<i>Staphylococcus epidermidis</i>	<i>Staphylococcus aureus</i>
Viridans <i>Streptococcus</i>	<i>Enterococcus faecalis</i>
<i>Streptococcus pneumoniae</i>	
Gram-Negative Bacilli	
<i>Escherichia coli</i>	<i>Serratia</i> spp.
<i>Klebsiella</i> spp.	<i>Acinetobacter</i> spp. ^a
<i>Pseudomonas aeruginosa</i>	<i>Stenotrophomonas</i> spp.
<i>Enterobacter</i> spp.	<i>Citrobacter</i> spp.
Non-aeruginosa <i>Pseudomonas</i> spp. ^a	
Gram-Positive Bacilli	
Diphtheroids	JK bacillus ^a
Fungi	
<i>Candida</i> spp.	<i>Mucor/Rhizopus</i>
<i>Aspergillus</i> spp.	

^aOften associated with intravenous catheters.

Cellulitis, an acute spreading inflammation of the skin, is most often caused by infection with group A *Streptococcus* or *Staphylococcus aureus*, virulent organisms normally found on the skin (Chap. 156). Although cellulitis tends to be circumscribed in normal hosts, it may spread rapidly in neutropenic patients. A tiny break in the skin may lead to spreading cellulitis, which is characterized by pain and erythema; in the affected patients, signs of infection (e.g., purulence) are often lacking. What might be a furuncle in a normal host may require amputation because of uncontrolled infection in a patient presenting with leukemia. A dramatic response to an infection that might be trivial in a normal host can mark the first sign of leukemia. Fortunately, granulocytopenic patients are likely to be infected with certain types of organisms (Table 104-4); thus the selection of an antibiotic regimen is somewhat easier than it might otherwise be (see “Antibacterial Therapy,” below). It is essential to recognize cellulitis early and to treat it aggressively. Patients who are neutropenic or who have previously received antibiotics for other reasons may develop cellulitis with unusual organisms (e.g., *Escherichia coli*, *Pseudomonas*, or fungi). Early treatment, even of innocent-looking lesions, is essential to prevent necrosis and loss of tissue. Debridement to prevent spread may sometimes be necessary early in the course of disease, but it can often be performed after chemotherapy, when the PMN count increases.



A



B

FIGURE 104-1 A. Papules related to *Escherichia coli* bacteremia in a patient with acute lymphocytic leukemia. B. The same lesions on the following day.

Sweet syndrome, or *febrile neutrophilic dermatosis*, was originally described in women with elevated white blood cell (WBC) counts. The disease is characterized by the presence of leukocytes in the lower dermis, with edema of the papillary body. Ironically, this disease now is usually seen in neutropenic patients with cancer, most often in association with acute myeloid leukemia (AML) but also in association with a variety of other malignancies. Sweet syndrome usually presents as red or bluish-red papules or nodules that may coalesce and form sharply bordered plaques (see Fig. 25e-41). The edema may suggest vesicles, but on palpation the lesions are solid, and vesicles probably never arise in this disease. The lesions are most common on the face, neck, and arms. On the legs, they may be confused with erythema nodosum (see Fig. 25e-40). The development of lesions is often accompanied by high fevers and an elevated erythrocyte sedimentation rate. Both the lesions and the temperature elevation respond dramatically to glucocorticoid administration. Treatment begins with high doses of glucocorticoids (prednisone, 60 mg/d) followed by tapered doses over the next 2–3 weeks.

Data indicate that *erythema multiforme* (see Fig. 25e-25) with mucous membrane involvement is often associated with herpes simplex virus (HSV) infection and is distinct from Stevens-Johnson syndrome, which is associated with drugs and tends to have a more widespread distribution. Because cancer patients are both immunosuppressed (and therefore susceptible to herpes infections) and heavily treated with drugs (and therefore subject to Stevens-Johnson syndrome [see Fig. 46e-4]), both of these conditions are common in this population.

Cytokines, which are used as adjuvants or primary treatments for cancer, can themselves cause characteristic rashes, further complicating the differential diagnosis. This phenomenon is a particular problem in bone marrow transplant recipients (Chap. 169), who, in addition to having the usual chemotherapy-, antibiotic-, and cytokine-induced rashes, are plagued by graft-versus-host disease.

CATHETER-RELATED INFECTIONS

Because IV catheters are commonly used in cancer chemotherapy and are prone to cause infection (Chap. 168), they pose a major problem in the care of patients with cancer. Some catheter-associated infections can be treated with antibiotics, whereas in others the catheter must be removed (Table 104-5). If the patient has a “tunneled” catheter (which consists of an entrance site, a subcutaneous tunnel, and an exit site), a red streak over the subcutaneous part of the line (the tunnel) is grounds

for immediate device removal. Failure to remove catheters under these circumstances may result in extensive cellulitis and tissue necrosis.

More common than tunnel infections are exit-site infections, often with erythema around the area where the line penetrates the skin. Most authorities (Chap. 172) recommend treatment (usually with vancomycin) for an exit-site infection caused by coagulase-negative *Staphylococcus*. Treatment of coagulase-positive staphylococcal infection is associated with a poorer outcome, and it is advisable to remove the catheter if possible. Similarly, most clinicians remove catheters associated with infections due to *P. aeruginosa* and *Candida* species, because such infections are difficult to treat and bloodstream infections with these organisms are likely to be deadly. Catheter infections caused by *Burkholderia cepacia*, *Stenotrophomonas* species, *Agrobacterium* species, *Acinetobacter baumannii*, *Pseudomonas* species other than *aeruginosa*, and carbapenem-resistant Enterobacteriaceae are likely to be very difficult to eradicate with antibiotics alone. Similarly, isolation of *Bacillus*, *Corynebacterium*, and *Mycobacterium* species should prompt removal of the catheter.

GASTROINTESTINAL TRACT–SPECIFIC SYNDROMES

Upper Gastrointestinal Tract Disease

INFECTIONS OF THE MOUTH The oral cavity is rich in aerobic and anaerobic bacteria (Chap. 201) that normally live in a commensal relationship with the host. The antimetabolic effects of chemotherapy cause a breakdown of mucosal host defenses, leading to ulceration of the mouth and the potential for invasion by resident bacteria. Mouth ulcerations afflict most patients receiving cytotoxic chemotherapy and have been associated with viridans streptococcal bacteremia. *Candida* infections of the mouth are very common. Fluconazole is clearly effective in the treatment of both local infections (thrush) and systemic infections (esophagitis) due to *Candida albicans*. Other azoles (e.g., voriconazole) as well as echinocandins offer similar efficacy as well as activity against the fluconazole-resistant organisms that are associated with chronic fluconazole treatment (Chap. 240).

Noma (cancrum oris), commonly seen in malnourished children, is a penetrating disease of the soft and hard tissues of the mouth and adjacent sites, with resulting necrosis and gangrene. It has a counterpart in immunocompromised patients and is thought to be due to invasion of the tissues by *Bacteroides*, *Fusobacterium*, and other normal inhabitants of the mouth. Noma is associated with debility, poor oral hygiene, and immunosuppression.

TABLE 104-5 APPROACH TO CATHETER INFECTIONS IN IMMUNOCOMPROMISED PATIENTS

Clinical Presentation or Isolated Pathogen	Catheter Removal	Antibiotics	Comments
Evidence of Infection, Negative Blood Cultures			
Exit-site erythema	Not necessary if infection responds to treatment	Usually, begin treatment for gram-positive cocci.	Coagulase-negative staphylococci are most common.
Tunnel-site erythema	Required	Treat for gram-positive cocci pending culture results.	Failure to remove the catheter may lead to necrosis of the involved area requiring skin grafts in the future.
Blood Culture–Positive Infections			
Coagulase-negative staphylococci	Line removal optimal but may be unnecessary if patient is clinically stable and responds to antibiotics	Usually, start with vancomycin. Linezolid, quinupristin/dalfopristin, and daptomycin are alternative agents.	If there are no contraindications to line removal, this course of action is optimal. If the line is removed, antibiotics may not be necessary.
Other gram-positive cocci (e.g., <i>Staphylococcus aureus</i> , <i>Enterococcus</i>); gram-positive rods (<i>Bacillus</i> , <i>Corynebacterium</i> spp.)	Recommended	Treat with antibiotics to which the organism is sensitive, with duration based on the clinical setting.	The incidence of metastatic infections following <i>S. aureus</i> infection and the difficulty of treating enterococcal infection make line removal the recommended course of action. In addition, gram-positive rods do not respond readily to antibiotics alone.
Gram-negative bacteria	Recommended	Use an agent to which the organism is shown to be sensitive.	Organisms like <i>Stenotrophomonas</i> , <i>Pseudomonas</i> , and <i>Burkholderia</i> are notoriously hard to treat, as are carbapenem-resistant organisms.
Fungi	Recommended	—	Fungal infections of catheters are extremely difficult to treat.

488 Viruses, particularly HSV, are a prominent cause of morbidity in immunocompromised patients, in whom they are associated with severe mucositis. The use of acyclovir, either prophylactically or therapeutically, is of value.

ESOPHAGEAL INFECTIONS The differential diagnosis of esophagitis (usually presenting as substernal chest pain upon swallowing) includes herpes simplex and candidiasis, both of which are readily treatable.

Lower Gastrointestinal Tract Disease Hepatic candidiasis (Chap. 240) results from seeding of the liver (usually from a gastrointestinal source) in neutropenic patients. It is most common among patients being treated for AML and usually presents symptomatically around the time the neutropenia resolves. The characteristic picture is that of persistent fever unresponsive to antibiotics, abdominal pain and tenderness or nausea, and elevated serum levels of alkaline phosphatase in a patient with hematologic malignancy who has recently recovered from neutropenia. The diagnosis of this disease (which may present in an indolent manner and persist for several months) is based on the finding of yeasts or pseudohyphae in granulomatous lesions. Hepatic ultrasound or CT may reveal bull's-eye lesions. MRI scans reveal small lesions not visible by other imaging modalities. The pathology (a granulomatous response) and the timing (with resolution of neutropenia and an elevation in granulocyte count) suggest that the host response to *Candida* is an important component of the manifestations of disease. In many cases, although organisms are visible, cultures of biopsied material may be negative. The designation *hepatosplenic candidiasis* or *hepatic candidiasis* is a misnomer because the disease often involves the kidneys and other tissues; the term *chronic disseminated candidiasis* may be more appropriate. Because of the risk of bleeding with liver biopsy, diagnosis is often based on imaging studies (MRI, CT). Treatment should be directed to the causative agent (usually *C. albicans* but sometimes *Candida tropicalis* or other less common *Candida* species).

Typhlitis *Typhlitis* (also referred to as necrotizing colitis, neutropenic colitis, necrotizing enteropathy, ileocecal syndrome, and cecitis) is a clinical syndrome of fever and right-lower-quadrant (or generalized abdominal) tenderness in an immunosuppressed host. This syndrome is classically seen in neutropenic patients after chemotherapy with cytotoxic drugs. It may be more common among children than among adults and appears to be much more common among patients with AML or ALL than among those with other types of cancer. Physical examination reveals right-lower-quadrant tenderness, with or without rebound tenderness. Associated diarrhea (often bloody) is common, and the diagnosis can be confirmed by the finding of a thickened cecal wall on CT, MRI, or ultrasonography. Plain films may reveal a right-lower-quadrant mass, but CT with contrast or MRI is a much more sensitive means of diagnosis. Although surgery is sometimes attempted to avoid perforation from ischemia, most cases resolve with medical therapy alone. The disease is sometimes associated with positive blood cultures (which usually yield aerobic gram-negative bacilli), and therapy is recommended for a broad spectrum of bacteria (particularly gram-negative bacilli, which are likely to be found in the bowel flora). Surgery is indicated in the case of perforation.

Clostridium difficile–Induced Diarrhea Patients with cancer are predisposed to the development of *C. difficile* diarrhea (Chap. 161) as a consequence of chemotherapy alone. Thus, they may test positive for *C. difficile* even without receiving antibiotics. Obviously, such patients are also subject to *C. difficile*–induced diarrhea as a result of antibiotic pressure. *C. difficile* should always be considered as a possible cause of diarrhea in cancer patients who have received either chemotherapy or antibiotics.

CENTRAL NERVOUS SYSTEM–SPECIFIC SYNDROMES

Meningitis The presentation of meningitis in patients with lymphoma or CLL and in patients receiving chemotherapy (particularly with glucocorticoids) for solid tumors suggests a diagnosis of cryptococcal or listerial infection. As noted previously, splenectomized patients are susceptible to rapid, overwhelming infection with encapsulated bacteria (including *S. pneumoniae*, *H. influenzae*, and *N. meningitidis*).

Similarly, patients who are antibody-deficient (e.g., those with CLL, those who have received intensive chemotherapy, or those who have undergone bone marrow transplantation) are likely to have infections caused by these bacteria. Other cancer patients, however, because of their defective cellular immunity, are likely to be infected with other pathogens (Table 104-3). Central nervous system (CNS) tuberculosis should be considered, especially in patients from countries where tuberculosis is highly prevalent in the population.

Encephalitis The spectrum of disease resulting from viral encephalitis is expanded in immunocompromised patients. A predisposition to infections with intracellular organisms similar to those encountered in patients with AIDS (Chap. 226) is seen in cancer patients receiving (1) high-dose cytotoxic chemotherapy, (2) chemotherapy affecting T cell function (e.g., fludarabine), or (3) antibodies that eliminate T cells (e.g., anti-CD3, alemtuzumab, anti-CD52) or cytokine activity (anti-tumor necrosis factor agents or interleukin 1 receptor antagonists). Infection with varicella-zoster virus (VZV) has been associated with encephalitis that may be caused by VZV-related vasculitis. Chronic viral infections may also be associated with dementia and encephalitic presentations. A diagnosis of progressive multifocal leukoencephalopathy (Chap. 164) should be considered when a patient who has received chemotherapy (rituximab in particular) presents with dementia (Table 104-6). Other abnormalities of the CNS that may be confused with infection include normal-pressure hydrocephalus and vasculitis resulting from CNS irradiation. It may be possible to differentiate these conditions by MRI.

Brain Masses Mass lesions of the brain most often present as headache with or without fever or neurologic abnormalities. Infections associated with mass lesions may be caused by bacteria (particularly *Nocardia*), fungi (particularly *Cryptococcus* or *Aspergillus*), or parasites (*Toxoplasma*). Epstein-Barr virus (EBV)–associated lymphoma may also present as single—or sometimes multiple—mass lesions of the brain. A biopsy may be required for a definitive diagnosis.

PULMONARY INFECTIONS

Pneumonia (Chap. 153) in immunocompromised patients may be difficult to diagnose because conventional methods of diagnosis depend on the presence of neutrophils. Bacterial pneumonia in neutropenic patients may present without purulent sputum—or, in fact, without any sputum at all—and may not produce physical findings suggestive of chest consolidation (rales or egophony).

In granulocytopenic patients with persistent or recurrent fever, the chest x-ray pattern may help to localize an infection and thus to determine which investigative tests and procedures should be undertaken and which therapeutic options should be considered (Table 104-7). In this setting, a simple chest x-ray is a screening tool; because the impaired host response results in less evidence of consolidation or infiltration, high-resolution CT is recommended for the diagnosis of pulmonary infections. The difficulties encountered in the management of pulmonary infiltrates relate in part to the difficulties of performing diagnostic procedures on the patients involved. When platelet counts can be increased to adequate levels by transfusion,

TABLE 104-6 DIFFERENTIAL DIAGNOSIS OF CENTRAL NERVOUS SYSTEM INFECTIONS IN PATIENTS WITH CANCER

Findings on CT or MRI	Underlying Predisposition	
	Prolonged Neutropenia	Defects in Cellular Immunity ^a
Mass lesions	<i>Aspergillus</i> , <i>Nocardia</i> , or <i>Cryptococcus</i> brain abscess	Toxoplasmosis, Epstein-Barr virus lymphoma (rare)
Diffuse encephalitis	Progressive multifocal leukoencephalopathy (JC virus)	Infection with varicella-zoster virus, cytomegalovirus, herpes simplex virus, human herpesvirus type 6, JC virus, <i>Listeria</i>

^aHigh-dose glucocorticoid therapy, cytotoxic chemotherapy.

TABLE 104-7 DIFFERENTIAL DIAGNOSIS OF CHEST INFILTRATES IN IMMUNOCOMPROMISED PATIENTS

Infiltrate	Cause of Pneumonia	
	Infectious	Noninfectious
Localized	Bacteria (including <i>Legionella</i> , mycobacteria)	Local hemorrhage or embolism, tumor
Nodular	Fungi (e.g., <i>Aspergillus</i> or <i>Mucor</i>), <i>Nocardia</i>	Recurrent tumor
Diffuse	Viruses (especially cytomegalovirus), <i>Chlamydia</i> , <i>Pneumocystis</i> , <i>Toxoplasma gondii</i> , mycobacteria	Congestive heart failure, radiation pneumonitis, drug-induced lung injury, lymphangitic spread of cancer

microscopic and microbiologic evaluation of the fluid obtained by endoscopic bronchial lavage is often diagnostic. Lavage fluid should be cultured for *Mycoplasma*, *Chlamydia*, *Legionella*, *Nocardia*, more common bacterial pathogens, fungi, and viruses. In addition, the possibility of *Pneumocystis* pneumonia should be considered, especially in patients with ALL or lymphoma who have not received prophylactic trimethoprim-sulfamethoxazole (TMP-SMX). The characteristics of the infiltrate may be helpful in decisions about further diagnostic and therapeutic maneuvers. Nodular infiltrates suggest fungal pneumonia (e.g., that caused by *Aspergillus* or *Mucor*). Such lesions may best be approached by visualized biopsy procedures. It is worth noting that while bacterial pneumonias classically present as lobar infiltrates in normal hosts, bacterial pneumonias in granulocytopenic hosts present with a paucity of signs, symptoms, or radiographic abnormalities; thus, the diagnosis is difficult.

Aspergillus species (Chap. 241) can colonize the skin and respiratory tract or cause fatal systemic illness. Although this fungus may cause aspergillomas in a previously existing cavity or may produce allergic bronchopulmonary disease in some patients, the major problem posed by this genus in neutropenic patients is invasive disease, primarily due to *Aspergillus fumigatus* or *Aspergillus flavus*. The organisms enter the host following colonization of the respiratory tract, with subsequent invasion of blood vessels. The disease is likely to present as a thrombotic or embolic event because of this ability of the fungi to invade blood vessels. The risk of infection with *Aspergillus* correlates directly with the duration of neutropenia. In prolonged neutropenia, positive surveillance cultures for nasopharyngeal colonization with *Aspergillus* may predict the development of disease.

Patients with *Aspergillus* infection often present with pleuritic chest pain and fever, which are sometimes accompanied by cough. Hemoptysis may be an ominous sign. Chest x-rays may reveal new focal infiltrates or nodules. Chest CT may reveal a characteristic halo consisting of a mass-like infiltrate surrounded by an area of low attenuation. The presence of a “crescent sign” on chest x-ray or chest CT, in which the mass progresses to central cavitation, is characteristic of invasive *Aspergillus* infection but may develop as the lesions are resolving.

In addition to causing pulmonary disease, *Aspergillus* may invade through the nose or palate, with deep sinus penetration. The appearance of a discolored area in the nasal passages or on the hard palate should prompt a search for invasive *Aspergillus*. This situation is likely to require surgical debridement. Catheter infections with *Aspergillus* usually require both removal of the catheter and antifungal therapy.

Diffuse interstitial infiltrates suggest viral, parasitic, or *Pneumocystis* pneumonia. If the patient has a diffuse interstitial pattern on chest x-ray, it may be reasonable, while considering invasive diagnostic procedures, to institute empirical treatment for *Pneumocystis* with TMP-SMX and for *Chlamydia*, *Mycoplasma*, and *Legionella* with a quinolone or azithromycin. Noninvasive procedures, such as staining of induced sputum smears for *Pneumocystis*, serum cryptococcal antigen tests, and urine testing for *Legionella* antigen, may be helpful. Serum galactomannan and β -D-glucan tests may be of value in diagnosing *Aspergillus* infection, but their utility is limited by their lack of sensitivity and specificity. The presence of an elevated level of β -D-glucan in the serum of a patient being treated for cancer who is not receiving prophylaxis against *Pneumocystis* suggests the diagnosis

of *Pneumocystis* pneumonia. Infections with viruses that cause only upper respiratory symptoms in immunocompetent hosts, such as respiratory syncytial virus (RSV), influenza viruses, and parainfluenza viruses, may be associated with fatal pneumonitis in immunocompromised hosts. CMV reactivation occurs in cancer patients receiving chemotherapy, but CMV pneumonia is most common among HSCT recipients (Chap. 169). Polymerase chain reaction testing now allows rapid diagnosis of viral pneumonia, which can lead to treatment in some cases (e.g., influenza). Multiplex studies that can detect a wide array of viruses in the lung and upper respiratory tract are now available and will lead to specific diagnoses of viral pneumonias.

Bleomycin is the most common cause of chemotherapy-induced lung disease. Other causes include alkylating agents (such as cyclophosphamide, chlorambucil, and melphalan), nitrosoureas (carmustine [BCNU], lomustine [CCNU], and methyl-CCNU), busulfan, procarbazine, methotrexate, and hydroxyurea. Both infectious and noninfectious (drug- and/or radiation-induced) pneumonitis can cause fever and abnormalities on chest x-ray; thus, the differential diagnosis of an infiltrate in a patient receiving chemotherapy encompasses a broad range of conditions (Table 104-7). The treatment of radiation pneumonitis (which may respond dramatically to glucocorticoids) or drug-induced pneumonitis is different from that of infectious pneumonia, and a biopsy may be important in the diagnosis. Unfortunately, no definitive diagnosis can be made in ~30% of cases, even after bronchoscopy.

Open-lung biopsy is the gold standard of diagnostic techniques. Biopsy via a visualized thoracostomy can replace an open procedure in many cases. When a biopsy cannot be performed, empirical treatment can be undertaken; a quinolone or an erythromycin derivative (azithromycin) and TMP-SMX are used in the case of diffuse infiltrates, and an antifungal agent is administered in the case of nodular infiltrates. The risks should be weighed carefully in these cases. If inappropriate drugs are administered, empirical treatment may prove toxic or ineffective; either of these outcomes may be riskier than biopsy.

CARDIOVASCULAR INFECTIONS

Patients with Hodgkin's disease are prone to persistent infections by *Salmonella*, sometimes (and particularly often in elderly patients) affecting a vascular site. The use of IV catheters deliberately lodged in the right atrium is associated with a high incidence of bacterial endocarditis, presumably related to valve damage followed by bacteremia. Nonbacterial thrombotic endocarditis (marantic endocarditis) has been described in association with a variety of malignancies (most often solid tumors) and may follow bone marrow transplantation as well. The presentation of an embolic event with a new cardiac murmur suggests this diagnosis. Blood cultures are negative in this disease of unknown pathogenesis.

ENDOCRINE SYNDROMES

Infections of the endocrine system have been described in immunocompromised patients. *Candida* infection of the thyroid may be difficult to diagnose during the neutropenic period. It can be defined by indium-labeled WBC scans or gallium scans after neutrophil counts increase. CMV infection can cause adrenalitis with or without resulting adrenal insufficiency. The presentation of a sudden endocrine anomaly in an immunocompromised patient can be a sign of infection in the involved end organ.

MUSCULOSKELETAL INFECTIONS

Infection that is a consequence of vascular compromise, resulting in gangrene, can occur when a tumor restricts the blood supply to muscles, bones, or joints. The process of diagnosis and treatment of such infection is similar to that in normal hosts, with the following caveats:

1. *In terms of diagnosis*, a lack of physical findings resulting from a lack of granulocytes in the granulocytopenic patient should make the clinician more aggressive in obtaining tissue rather than more willing to rely on physical signs.
2. *In terms of therapy*, aggressive debridement of infected tissues may be required. However, it is usually difficult to operate on patients

who have recently received chemotherapy, both because of a lack of platelets (which results in bleeding complications) and because of a lack of WBCs (which may lead to secondary infection). A blood culture positive for *Clostridium perfringens*—an organism commonly associated with gas gangrene—can have a number of meanings (Chap. 179). *Clostridium septicum* bacteremia is associated with the presence of an underlying malignancy. Bloodstream infections with intestinal organisms such as *Streptococcus bovis* biotype 1 and *C. perfringens* may arise spontaneously from lower gastrointestinal lesions (tumor or polyps); alternatively, these lesions may be harbingers of invasive disease. The clinical setting must be considered in order to define the appropriate treatment for each case.

RENAL AND URETERAL INFECTIONS

Infections of the urinary tract are common among patients whose ureteral excretion is compromised (Table 104-1). *Candida*, which has a predilection for the kidney, can invade either from the bloodstream or in a retrograde manner (via the ureters or bladder) in immunocompromised patients. The presence of “fungus balls” or persistent candiduria suggests invasive disease. Persistent funguria (with *Aspergillus* as well as *Candida*) should prompt a search for a nidus of infection in the kidney.

Certain viruses are typically seen only in immunosuppressed patients. BK virus (polyomavirus hominis 1) has been documented in the urine of bone marrow transplant recipients and, like adenovirus, may be associated with hemorrhagic cystitis.

ABNORMALITIES THAT PREDISPOSE TO INFECTION

(Table 104-1)

THE LYMPHOID SYSTEM

It is beyond the scope of this chapter to detail how all the immunologic abnormalities that result from cancer or from chemotherapy for cancer lead to infections. Disorders of the immune system are discussed in other sections of this book. As has been noted, patients with antibody deficiency are predisposed to overwhelming infection with encapsulated bacteria (including *S. pneumoniae*, *H. influenzae*, and *N. meningitidis*). Infections that result from the lack of a functional cellular immune system are described in Chap. 226. It is worth mentioning, however, that patients undergoing intensive chemotherapy for any form of cancer will have not only defects due to granulocytopenia but also lymphocyte dysfunction, which may be profound. Thus, these patients—especially those receiving glucocorticoid-containing regimens or drugs that inhibit either T cell activation (calcineurin inhibitors or drugs like fludarabine, which affect lymphocyte function) or cytokine induction—should be given prophylaxis for *Pneumocystis pneumonia*.

Patients receiving treatment that eliminates B cells (e.g., with anti-CD20 antibodies or rituximab) are especially vulnerable to intercurrent viral infections. The incidence of progressive multifocal leukoencephalopathy (caused by JC virus) is elevated in these patients.

THE HEMATOPOIETIC SYSTEM

Initial studies in the 1960s revealed a dramatic increase in the incidence of infections (fatal and nonfatal) among cancer patients with a granulocyte count of $<500/\mu\text{L}$. The use of prophylactic antibacterial agents has reduced the number of bacterial infections, but 35–78% of febrile neutropenic patients being treated for hematologic malignancies develop infections at some time during chemotherapy. Aerobic pathogens (both gram-positive and gram-negative) predominate in all series, but the exact organisms isolated vary from center to center. Infections with anaerobic organisms are uncommon. Geographic patterns affect the types of fungi isolated. Tuberculosis and malaria are common causes of fever in the developing world and may present in this setting as well.

Neutropenic patients are unusually susceptible to infection with a wide variety of bacteria; thus, antibiotic therapy should be initiated

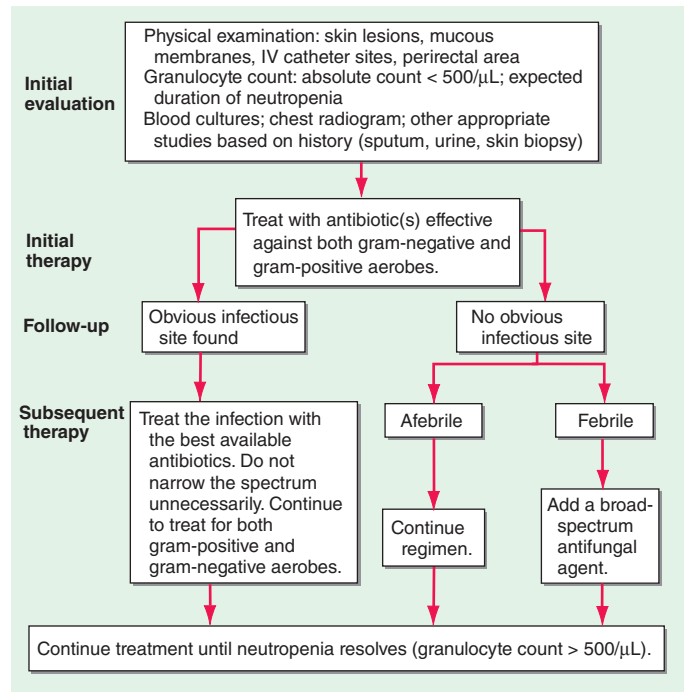


FIGURE 104-2 Algorithm for the diagnosis and treatment of fever and neutropenia.

promptly to cover likely pathogens if infection is suspected. Indeed, early initiation of antibacterial agents is mandatory to prevent deaths. Like most immunocompromised patients, neutropenic patients are threatened by their own microbial flora, including gram-positive and gram-negative organisms found commonly on the skin and mucous membranes and in the bowel (Table 104-4). Because treatment with narrow-spectrum agents leads to infection with organisms not covered by the antibiotics used, the initial regimen should target all pathogens likely to be the initial causes of bacterial infection in neutropenic hosts. As noted in the algorithm shown in Fig. 104-2, administration of antimicrobial agents is routinely continued until neutropenia resolves—i.e., the granulocyte count is sustained above $500/\mu\text{L}$ for at least 2 days. In some cases, patients remain febrile after resolution of neutropenia. In these instances, the risk of sudden death from overwhelming bacteremia is greatly reduced, and the following diagnoses should be seriously considered: (1) fungal infection, (2) bacterial abscesses or undrained foci of infection, and (3) drug fever (including reactions to antimicrobial agents as well as to chemotherapy or cytokines). In the proper setting, viral infection or graft-versus-host disease should be considered. In clinical practice, antibacterial therapy is usually discontinued when the patient is no longer neutropenic and all evidence of bacterial disease has been eliminated. Antifungal agents are then discontinued if there is no evidence of fungal disease. If the patient remains febrile, a search for viral diseases or unusual pathogens is conducted while unnecessary cytokines and other drugs are systematically eliminated from the regimen.

TREATMENT INFECTIONS IN CANCER PATIENTS

ANTIBACTERIAL THERAPY

Hundreds of antibacterial regimens have been tested for use in patients with cancer. The major risk of infection is related to the degree of neutropenia seen as a consequence of either the disease or the therapy. Many of the relevant studies have involved small populations in which the outcomes have generally been good, and most have lacked the statistical power to detect differences among the regimens studied. Each febrile neutropenic patient should be approached as a unique problem, with particular attention given to previous infections and recent antibiotic exposures. Several general

guidelines are useful in the initial treatment of neutropenic patients with fever (Fig. 104-2):

1. In the initial regimen, it is necessary to use antibiotics active against both gram-negative and gram-positive bacteria (Table 104-4).
2. Monotherapy with an aminoglycoside or an antibiotic lacking good activity against gram-positive organisms (e.g., ciprofloxacin or aztreonam) is not adequate in this setting.
3. The agents used should reflect both the epidemiology and the antibiotic resistance pattern of the hospital.
4. If the pattern of resistance justifies its use, a single third-generation cephalosporin constitutes an appropriate initial regimen in many hospitals.
5. Most standard regimens are designed for patients who have not previously received prophylactic antibiotics. The development of fever in a patient who has received antibiotics affects the choice of subsequent therapy, which should target resistant organisms and organisms known to cause infections in patients being treated with the antibiotics already administered.
6. Randomized trials have indicated the safety of oral antibiotic regimens in the treatment of “low-risk” patients with fever and neutropenia. Outpatients who are expected to remain neutropenic for <10 days and who have no concurrent medical problems (such as hypotension, pulmonary compromise, or abdominal pain) can be classified as low risk and treated with a broad-spectrum oral regimen.
7. Several large-scale studies indicate that prophylaxis with a fluoroquinolone (ciprofloxacin or levofloxacin) decreases morbidity and mortality rates among afebrile patients who are anticipated to have neutropenia of long duration.

Commonly used antibiotic regimens for the treatment of febrile patients in whom prolonged neutropenia (>7 days) is anticipated include (1) ceftazidime or cefepime, (2) piperacillin/tazobactam, or (3) imipenem/cilastatin or meropenem. All three regimens have shown equal efficacy in large trials. All three are active against *P. aeruginosa* and a broad spectrum of aerobic gram-positive and gram-negative organisms. Imipenem/cilastatin has been associated with an elevated rate of *C. difficile* diarrhea, and many centers reserve carbapenem antibiotics for treatment of gram-negative bacteria that produce extended-spectrum β -lactamases; these limitations make carbapenems less attractive as an initial regimen. Despite the frequent involvement of coagulase-negative staphylococci, the initial use of vancomycin or its automatic addition to the initial regimen has not resulted in improved outcomes, and the antibiotic does exert toxic effects. For these reasons, only judicious use of vancomycin is recommended—for example, when there is good reason to suspect the involvement of coagulase-negative staphylococci (e.g., the appearance of erythema at the exit site of a catheter or a positive culture for methicillin-resistant *S. aureus* or coagulase-negative staphylococci). Because the sensitivities of bacteria vary from hospital to hospital, clinicians are advised to check their local sensitivities and to be aware that resistance patterns can change quickly, necessitating a change in approach to patients with fever and neutropenia. Similarly, infection control services should monitor for basic antibiotic resistance and for fungal infections. The appearance of a large number of *Aspergillus* infections, in particular, suggests the possibility of an environmental source that requires further investigation and remediation.

The initial antibacterial regimen should be refined on the basis of culture results (Fig. 104-2). Blood cultures are the most relevant basis for selection of therapy; surface cultures of skin and mucous membranes may be misleading. In the case of gram-positive bacteremia or another gram-positive infection, it is important that the antibiotic be optimal for the organism isolated. Once treatment with broad-spectrum antibiotics has begun, it is not desirable to discontinue all antibiotics because of the risk of failing to treat a potentially fatal bacterial infection; the addition of more and more

antibacterial agents to the regimen is not appropriate unless there is a clinical or microbiologic reason to do so. Planned progressive therapy (the serial, empirical addition of one drug after another without culture data) is not efficacious in most settings and may have unfortunate consequences. Simply adding another antibiotic for fear that a gram-negative infection is present is a dubious practice. The synergy exhibited by β -lactams and aminoglycosides against certain gram-negative organisms (especially *P. aeruginosa*) provides the rationale for using two antibiotics in this setting, but recent analyses suggest that efficacy is not enhanced by the addition of aminoglycosides, while toxicity may be increased. Mere “double coverage,” with the addition of a quinolone or another antibiotic that is not likely to exhibit synergy, has not been shown to be of benefit and may cause additional toxicities and side effects. Cephalosporins can cause bone marrow suppression, and vancomycin is associated with neutropenia in some healthy individuals. Furthermore, the addition of multiple cephalosporins may induce β -lactamase production by some organisms; cephalosporins and double β -lactam combinations should probably be avoided altogether in *Enterobacter* infections.

ANTIFUNGAL THERAPY

Fungal infections in cancer patients are most often associated with neutropenia. Neutropenic patients are predisposed to the development of invasive fungal infections, most commonly those due to *Candida* and *Aspergillus* species and occasionally those caused by *Mucor*, *Rhizopus*, *Fusarium*, *Trichosporon*, *Bipolaris*, and others. Cryptococcal infection, which is common among patients taking immunosuppressive agents, is uncommon among neutropenic patients receiving chemotherapy for AML. Invasive candidal disease is usually caused by *C. albicans* or *C. tropicalis* but can be caused by *C. krusei*, *C. parapsilosis*, and *C. glabrata*.

For decades, it has been common clinical practice to add amphotericin B to antibacterial regimens if a neutropenic patient remains febrile despite 4–7 days of treatment with antibacterial agents. The rationale for this empirical addition is that it is difficult to culture fungi before they cause disseminated disease and that mortality rates from disseminated fungal infections in granulocytopenic patients are high. Before the introduction of newer azoles into clinical practice, amphotericin B was the mainstay of antifungal therapy. The insolubility of amphotericin B has resulted in the marketing of several lipid formulations that are less toxic than the amphotericin B deoxycholate complex. Echinocandins (e.g., caspofungin) are useful in the treatment of infections caused by azole-resistant *Candida* strains as well as in therapy for aspergillosis and have been shown to be equivalent to liposomal amphotericin B for the empirical treatment of patients with prolonged fever and neutropenia. Newer azoles have also been demonstrated to be effective in this setting. Although fluconazole is efficacious in the treatment of infections due to many *Candida* species, its use against serious fungal infections in immunocompromised patients is limited by its narrow spectrum: it has no activity against *Aspergillus* or against several non-*albicans Candida* species. The broad-spectrum azoles (e.g., voriconazole and posaconazole) provide another option for the treatment of *Aspergillus* infections (Chap. 241), including CNS infection. Clinicians should be aware that the spectrum of each azole is somewhat different and that no drug can be assumed to be efficacious against all fungi. *Aspergillus terreus* is resistant to amphotericin B. Although voriconazole is active against *Pseudallescheria boydii*, amphotericin B is not; however, voriconazole has no activity against *Mucor*. Posaconazole, which is administered orally, is useful as a prophylactic agent in patients with prolonged neutropenia. Studies in progress are assessing the use of these agents in combinations. **For a full discussion of antifungal therapy, see Chap. 235.**

ANTIVIRAL THERAPY

The availability of a variety of agents active against herpes-group viruses, including some new agents with a broader spectrum of activity, has heightened focus on the treatment of viral infections,

492 which pose a major problem in cancer patients. Viral diseases caused by the herpes group are prominent. Serious (and sometimes fatal) infections due to HSV and VZV are well documented in patients receiving chemotherapy. CMV may also cause serious disease, but fatalities from CMV infection are more common in HSCT recipients. The roles of human herpesvirus (HHV)-6, HHV-7, and HHV-8 (Kaposi's sarcoma-associated herpesvirus) in cancer patients are still being defined (Chap. 219). EBV lymphoproliferative disease (LPD) can occur in patients receiving chemotherapy but is much more common among transplant recipients (Chap. 169). While clinical experience is most extensive with acyclovir, which can be used therapeutically or prophylactically, a number of derivative drugs offer advantages over this agent (Chap. 215e).

In addition to the herpes group, several respiratory viruses (especially RSV) may cause serious disease in cancer patients. Although influenza vaccination is recommended (see below), it may be ineffective in this patient population. The availability of antiviral drugs with activity against influenza viruses gives the clinician additional options for the prophylaxis and treatment of these patients (Chaps. 215e and 224).

OTHER THERAPEUTIC MODALITIES

Another way to address the problems posed by the febrile neutropenic patient is to replenish the neutrophil population. Although granulocyte transfusions may be effective in the treatment of refractory gram-negative bacteremia, they do not have a documented role in prophylaxis. Because of the expense, the risk of leukoagglutinin reactions (which has probably been decreased by improved cell-separation procedures), and the risk of transmission of CMV from unscreened donors (which has been reduced by the use of filters), granulocyte transfusion is reserved for patients whose condition is unresponsive to antibiotics. This modality is efficacious for documented gram-negative bacteremia refractory to antibiotics, particularly in situations where granulocyte numbers will be depressed for only a short period. The demonstrated usefulness of granulocyte colony-stimulating factor in mobilizing neutrophils and advances in preservation techniques may make this option more useful than in the past.

A variety of cytokines, including granulocyte colony-stimulating factor and granulocyte-macrophage colony-stimulating factor, enhance granulocyte recovery after chemotherapy and consequently shorten the period of maximal vulnerability to fatal infections. The role of these cytokines in routine practice is still a matter of some debate. Most authorities recommend their use only when neutropenia is both severe and prolonged. The cytokines themselves may have adverse effects, including fever, hypoxemia, and pleural effusions or serositis in other areas (Chap. 372e).

Once neutropenia has resolved, the risk of infection decreases dramatically. However, depending on what drugs they receive, patients who continue on chemotherapeutic protocols remain at high risk for certain diseases. Any patient receiving more than a maintenance dose of glucocorticoids (e.g., in many treatment regimens for diffuse lymphoma) should also receive prophylactic TMP-SMX because of the risk of *Pneumocystis* infection; those with ALL should receive such prophylaxis for the duration of chemotherapy.

PREVENTION OF INFECTION IN CANCER PATIENTS

EFFECT OF THE ENVIRONMENT

Outbreaks of fatal *Aspergillus* infection have been associated with construction projects and materials in several hospitals. The association between spore counts and risk of infection suggests the need for a high-efficiency air-handling system in hospitals that care for large numbers of neutropenic patients. The use of laminar-flow rooms and prophylactic antibiotics has decreased the number of infectious episodes in severely neutropenic patients. However, because of the expense of such a program and the failure to show that it dramatically affects mortality rates, most centers do not routinely use laminar flow to care for neutropenic patients. Some centers use "reverse isolation," in which health care providers and visitors to a patient who is

neutropenic wear gowns and gloves. Since most of the infections these patients develop are due to organisms that colonize the patients' own skin and bowel, the validity of such schemes is dubious, and limited clinical data do not support their use. Hand washing by all staff caring for neutropenic patients should be required to prevent the spread of resistant organisms.

The presence of large numbers of bacteria (particularly *P. aeruginosa*) in certain foods, especially fresh vegetables, has led some authorities to recommend a special "low-bacteria" diet. A diet consisting of cooked and canned food is satisfactory to most neutropenic patients and does not involve elaborate disinfection or sterilization protocols. However, there are no studies to support even this type of dietary restriction. Counseling of patients to avoid leftovers, deli foods, undercooked meat, and unpasteurized dairy products is recommended.

PHYSICAL MEASURES

Although few studies address this issue, patients with cancer are predisposed to infections resulting from anatomic compromise (e.g., lymphedema resulting from node dissections after radical mastectomy). Surgeons who specialize in cancer surgery can provide specific guidelines for the care of such patients, and patients benefit from common-sense advice about how to prevent infections in vulnerable areas.

IMMUNOGLOBULIN REPLACEMENT

Many patients with multiple myeloma or CLL have immunoglobulin deficiencies as a result of their disease, and all allogeneic bone marrow transplant recipients are hypogammaglobulinemic for a period after transplantation. However, current recommendations reserve intravenous immunoglobulin replacement therapy for those patients with severe (<400 mg of total IgG/dL), prolonged hypogammaglobulinemia and a history of repeated infections. Antibiotic prophylaxis has been shown to be cheaper and is efficacious in preventing infections in most CLL patients with hypogammaglobulinemia. Routine use of immunoglobulin replacement is not recommended.

SEXUAL PRACTICES

The use of condoms is recommended for severely immunocompromised patients. Any sexual practice that results in oral exposure to feces is not recommended. Neutropenic patients should be advised to avoid any practice that results in trauma, as even microscopic cuts may result in bacterial invasion and fatal sepsis.

ANTIBIOTIC PROPHYLAXIS

Several studies indicate that the use of oral fluoroquinolones prevents infection and decreases mortality rates among severely neutropenic patients. Prophylaxis for *Pneumocystis* is mandatory for patients with ALL and for all cancer patients receiving glucocorticoid-containing chemotherapy regimens.

VACCINATION OF CANCER PATIENTS

In general, patients undergoing chemotherapy respond less well to vaccines than do normal hosts. Their greater need for vaccines thus leads to a dilemma in their management. Purified proteins and inactivated vaccines are almost never contraindicated and should be given to patients even during chemotherapy. For example, all adults should receive diphtheria-tetanus toxoid boosters at the indicated times as well as seasonal influenza vaccine. However, if possible, vaccination should not be undertaken concurrent with cytotoxic chemotherapy. If patients are expected to be receiving chemotherapy for several months and vaccination is indicated (e.g., influenza vaccination in the fall), the vaccine should be given midcycle—as far apart in time as possible from the antimetabolic agents that will prevent an immune response. The meningococcal and pneumococcal polysaccharide vaccines should be given to patients before splenectomy, if possible. The *H. influenzae* type b conjugate vaccine should be administered to all splenectomized patients.

In general, live virus (or live bacterial) vaccines should not be given to patients during intensive chemotherapy because of the risk of disseminated infection. Recommendations on vaccination are summarized in Table 104-2 (see www.cdc.gov/vaccine for updated recommendations).

MELANOMA

Pigmented lesions are among the most common findings on skin examination. The challenge is to distinguish cutaneous melanomas, which account for the overwhelming majority of deaths resulting from skin cancer, from the remainder, which are usually benign. Cutaneous melanoma can occur in adults of all ages, even young individuals, and people of all colors; its location on the skin and its distinct clinical features make it detectable at a time when complete surgical excision is possible. Examples of malignant and benign pigmented lesions are shown in [Fig. 105-1](#).

EPIDEMIOLOGY

Melanoma is an aggressive malignancy of melanocytes, pigment-producing cells that originate from the neural crest and migrate to

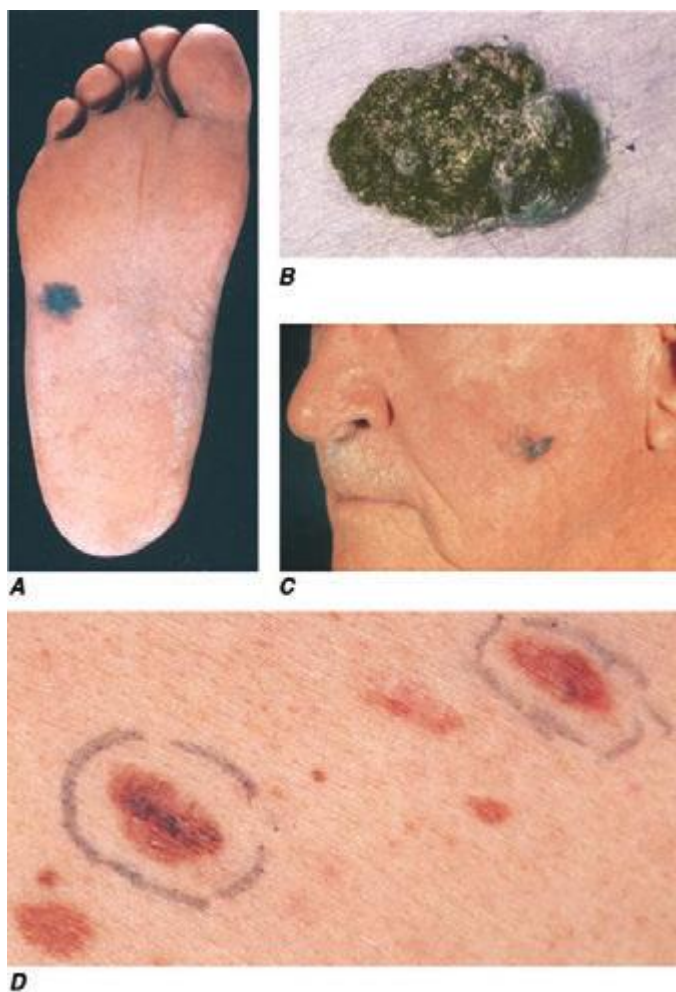


FIGURE 105-1 Atypical and malignant pigmented lesions. The most common melanoma is superficial spreading melanoma (not pictured). **A.** Acral lentiginous melanoma is the most common melanoma in blacks, Asians, and Hispanics and occurs as an enlarging hyperpigmented macule or plaque on the palms and soles. Lateral pigment diffusion is present. **B.** Nodular melanoma most commonly manifests as a rapidly growing, often ulcerated or crusted black nodule. **C.** Lentigo maligna melanoma occurs on sun-exposed skin as a large, hyperpigmented macule or plaque with irregular borders and variable pigmentation. **D.** Dysplastic nevi are irregularly pigmented and shaped nevomelanocytic lesions that may be associated with familial melanoma.

the skin, meninges, mucous membranes, upper esophagus, and eyes. Melanocytes in each of these locations have the potential for malignant transformation. Cutaneous melanoma is predominantly a malignancy of white-skinned people (98% of cases), and the incidence correlates with latitude of residence, providing strong evidence for the role of sun exposure. Men are affected slightly more than women (1.3:1), and the median age at diagnosis is the late fifties. Dark-skinned populations (such as those of India and Puerto Rico), blacks, and East Asians also develop melanoma, albeit at rates 10–20 times lower than those in whites. Cutaneous melanomas in these populations are diagnosed more often at a higher stage, and patients tend to have worse outcomes. Furthermore, in nonwhite populations, there is a much higher frequency of acral (subungual, plantar, palmar) and mucosal melanomas. In 2014, more than 76,000 individuals in the United States were expected to develop melanoma, and approximately 9700 were expected to die. There will be nearly 50,000 annual deaths worldwide as a result of melanoma. Data from the Connecticut Tumor Registry support an unremitting increase in the incidence and mortality of melanoma. In the past 60 years, there have been 17-fold and 9-fold increases in incidence for men and women, respectively. In the same six decades, there has been a tripling of mortality rates for men and doubling for women. Mortality rates begin to rise at age 55, with the greatest increase in men age >65 years. Of particular concern is the increase in rates among women <40 years of age. Much of this increase is believed to be associated with a greater emphasis on tanned skin as a marker of beauty, the increased availability and use of indoor tanning beds, and exposure to intense ultraviolet (UV) light in childhood. These statistics highlight the need to promote prevention and early detection.

RISK FACTORS

Presence of Nevi The risk of developing melanoma is related to genetic, environmental, and host factors ([Table 105-1](#)). The strongest risk factors for melanoma are the presence of multiple benign or atypical nevi and a family or personal history of melanoma. The presence of melanocytic nevi, common or dysplastic, is a marker for increased risk of melanoma. Nevi have been referred to as precursor lesions because they can transform into melanomas; however, the actual risk for any specific nevus is exceedingly low. About one-quarter of melanomas are histologically associated with nevi, but the majority arise *de novo*. The number of clinically atypical moles may vary from one to several hundred, and they usually differ from one another in appearance. The borders are often hazy and indistinct, and the pigment pattern is more highly varied than that in benign acquired nevi. Individuals with clinically atypical moles and a strong family history of melanoma have been reported to have a >50% lifetime risk for developing melanoma and warrant close follow-up with a dermatologist. Of the 90% of patients whose disease is sporadic (i.e., who lack a family history of melanoma), ~40% have clinically atypical moles, compared with an estimated 5–10% of the population at large.

Congenital melanocytic nevi, which are classified as small (≤ 1.5 cm), medium (1.5–20 cm), and giant (> 20 cm), can be precursors for melanoma. The risk is highest for the giant melanocytic nevus, also called the bathing trunk nevus, a rare malformation that affects 1 in 30,000–100,000 individuals. Since the lifetime risk of melanoma development is estimated to be as high as 6%, prophylactic excision early in life is prudent. This usually requires staged removal with coverage

TABLE 105-1 FACTORS ASSOCIATED WITH INCREASED RISK OF MELANOMA

Total body nevi (higher number = higher risk)
Dysplastic nevi (10-fold increased risk)
Family or personal history
Ultraviolet exposure/sunburns/tanning booths
Light skin/hair/eye color
Poor tanning ability
Freckling
<i>CDKN2A</i> , <i>CDK4</i> , <i>MITF</i> mutations
<i>MCT1R</i> variants

494 by split-thickness skin grafts. Surgery cannot remove all at-risk nevus cells, as some may penetrate into the muscles or central nervous system (CNS) below the nevus. Small- to medium-size congenital melanocytic nevi affect approximately 1% of persons; the risk of melanoma developing in these lesions is not known but appears to be relatively low. The management of small- to medium-size congenital melanocytic nevi remains controversial.

Personal and Family History Once diagnosed, patients with melanoma require a lifetime of surveillance because their risk of developing another melanoma is 10 times that of the general population. First-degree relatives have a higher risk of developing melanoma than do individuals without a family history, but only 5–10% of all melanomas are truly familial. In familial melanoma, patients tend to be younger at first diagnosis, lesions are thinner, survival is improved, and multiple primary melanomas are common.



Genetic Susceptibility Approximately 20–40% of cases of hereditary melanoma (0.2–2% of all melanomas) are due to germline mutations in the cell cycle regulatory gene cyclin-dependent kinase inhibitor 2A (*CDKN2A*). In fact, 70% of all cutaneous melanomas have mutations or deletions affecting the *CDKN2A* locus on chromosome 9p21. This locus encodes two distinct tumor-suppressor proteins from alternate reading frames: p16 and ARF (p14^{ARF}). The p16 protein inhibits CDK4/6-mediated phosphorylation and inactivation of the retinoblastoma (RB) protein, whereas ARF inhibits MDM2 ubiquitin-mediated degradation of p53. The end result of the loss of *CDKN2A* is inactivation of two critical tumor-suppressor pathways, RB and p53, which control entry of cells into the cell cycle. Several studies have shown an increased risk of pancreatic cancer among melanoma-prone families with *CDKN2A* mutations. A second high-risk locus for melanoma susceptibility, *CDK4*, is located on chromosome 12q13 and encodes the kinase inhibited by p16. *CDK4* mutations, which also inactivate the RB pathway, are much rarer than *CDKN2A* mutations. Germline mutations in the melanoma lineage-specific oncogene microphthalmia-associated transcription factor (*MITF*) predispose to both familial and sporadic melanomas.

The melanocortin-1 receptor (*MC1R*) gene is a moderate-risk inherited melanoma susceptibility factor. Solar radiation stimulates the production of melanocortin (α -melanocyte-stimulating hormone [α -MSH]), the ligand for *MC1R*, which is a G-protein-coupled receptor that signals via cyclic AMP and regulates the amount and type of pigment produced. *MC1R* is highly polymorphic, and among its 80 variants are those that result in partial loss of signaling and lead to the production of red/yellow pheomelanins, which are not sun-protective and produce red hair, rather than brown/black eumelanins that are photoprotective. This red hair color (RHC) phenotype is associated with fair skin, red hair, freckles, increased sun sensitivity, and increased risk of melanoma. In addition to its weak UV shielding capacity relative to eumelanin, increased pheomelanin production in patients with inactivating polymorphisms of *MC1R* also provides a UV-independent carcinogenic contribution to melanomagenesis via oxidative damage.

A number of other more common, low-penetrance polymorphisms that have small effects on melanoma susceptibility include other genes related to pigmentation, nevus count, immune responses, DNA repair, metabolism, and the vitamin D receptor.

PREVENTION AND EARLY DETECTION

Primary prevention of melanoma and nonmelanoma skin cancer (NMSC) is based on protection from the sun. Public health initiatives, such as the SunSmart program that started in Australia and now is operative in Europe and the United States, have demonstrated that behavioral change can decrease the incidence of NMSC and melanoma. Preventive measures should start early in life because damage from UV light begins early despite the fact that cancers develop years later. Biological factors are increasingly being understood, such as tanning addiction, which is postulated to involve stimulation of reward centers in the brain involving dopamine pathways, and cutaneous secretion of β -endorphins after UV exposure, and may represent

another area for preventive intervention. Regular use of broad-spectrum sunscreens that block UVA and UVB with a sun protection factor (SPF) of at least 30 and protective clothing should be encouraged. Avoidance of tanning beds and midday (10:00 A.M. to 2:00 P.M.) sun exposure is recommended.

Secondary prevention comprises education, screening, and early detection. Patients should be educated in the clinical features of melanoma (ABCDEs; see following “Diagnosis” section) and advised to report any growth or other change in a pigmented lesion. Brochures are available from the American Cancer Society, the American Academy of Dermatology, the National Cancer Institute, and the Skin Cancer Foundation. Self-examination at 6- to 8-week intervals may enhance the likelihood of detecting change. Although the U.S. Preventive Services Task Force states that evidence is insufficient to recommend for or against skin cancer screening, a full-body skin exam seems to be a simple, practical way to approach reducing the mortality rate for skin cancer. Depending on the presence or absence of risk factors, strategies for early detection can be individualized. This is particularly true for patients with clinically atypical moles (dysplastic nevi) and those with a personal history of melanoma. For these individuals, surveillance should be performed by the dermatologist and include total-body photography and dermoscopy where appropriate. Individuals with three or more primary melanomas and families with at least one invasive melanoma and two or more cases of melanoma and/or pancreatic cancer among first- or second-degree relatives on the same side of the family may benefit from genetic testing. Precancerous and in situ lesions should be treated early. Early detection of small tumors allows the use of simpler treatment modalities with higher cure rates and lower morbidity.

DIAGNOSIS

The main goal is to identify a melanoma before tumor invasion and life-threatening metastases have occurred. Early detection may be facilitated by applying the ABCDEs: *a*symmetry (benign lesions are usually symmetric); *b*order irregularity (most nevi have clear-cut borders); *c*olor variegation (benign lesions usually have uniform light or dark pigment); *d*iameter >6 mm (the size of a pencil eraser); and *e*volving (any change in size, shape, color, or elevation or new symptoms such as bleeding, itching, and crusting). Benign nevi usually appear on sun-exposed skin above the waist, rarely involving the scalp, breasts, or buttocks; atypical moles usually appear on sun-exposed skin, most often on the back, but can involve the scalp, breasts, or buttocks. Benign nevi are present in 85% of adults, with 10–40 moles scattered over the body; atypical nevi can be present in the hundreds.

The entire skin surface, including the scalp and mucous membranes, as well as the nails should be examined in each patient. Bright room illumination is important, and a hand lens is helpful for evaluating variation in pigment pattern. Any suspicious lesions should be biopsied, evaluated by a specialist, or recorded by chart and/or photography for follow-up. A focused method for examining individual lesions, dermoscopy, employs low-level magnification of the epidermis and may allow a more precise visualization of patterns of pigmentation than is possible with the naked eye. Complete physical examination with attention to the regional lymph nodes is part of the initial evaluation in a patient with suspected melanoma. The patient should be advised to have other family members screened if either melanoma or clinically atypical moles (dysplastic nevi) are present. Patients who fit into high-risk groups should be instructed to perform monthly self-examinations.

Biopsy Any pigmented cutaneous lesion that has changed in size or shape or has other features suggestive of malignant melanoma is a candidate for biopsy. An excisional biopsy with 1- to 3-mm margins is suggested. This facilitates pathologic assessment of the lesion, permits accurate measurement of thickness if the lesion is melanoma, and constitutes definitive treatment if the lesion is benign. For lesions that are large or on anatomic sites where excisional biopsy may not be feasible (such as the face, hands, and feet), an incisional biopsy through the most nodular or darkest area of the lesion is acceptable; this should

include the vertical growth phase of the primary tumor, if present. Incisional biopsy does not appear to facilitate the spread of melanoma. For suspicious lesions, every attempt should be made to preserve the ability to assess the deep and peripheral margins and to perform immunohistochemistry. Shave biopsies are an acceptable alternative, particularly if the suspicion of malignancy is low, but they should be deep and include underlying fat; cauterization should be avoided. The biopsy should be read by a pathologist experienced in pigmented lesions, and the report should include Breslow thickness, mitoses per square millimeter for lesions ≤ 1 mm, presence or absence of ulceration, and peripheral and deep margin status. Breslow thickness is the greatest thickness of a primary cutaneous melanoma measured on the slide from the top of the epidermal granular layer, or from the ulcer base, to the bottom of the tumor. To distinguish melanomas from benign nevi in cases with challenging histology, fluorescence in situ hybridization (FISH) with multiple probes and comparative genome hybridization (CGH) can be helpful.

CLINICAL CLASSIFICATION

Four major types of cutaneous melanoma have been recognized (Table 105-2). In three of these types—*superficial spreading melanoma*, *lentigo maligna melanoma*, and *acral lentiginous melanoma*—the lesion has a period of superficial (so-called radial) growth during which it increases in size but does not penetrate deeply. It is during this period that the melanoma is most capable of being cured by surgical excision. The fourth type—*nodular melanoma*—does not have a recognizable radial growth phase and usually presents as a deeply invasive lesion that is capable of early metastasis. When tumors begin to penetrate deeply into the skin, they are in the so-called vertical growth phase. Melanomas with a radial growth phase are characterized by irregular and sometimes notched borders, variation in pigment pattern, and variation in color. An increase in size or change in color is noted by the patient in 70% of early lesions. Bleeding, ulceration, and pain are late signs and are of little help in early recognition. Superficial spreading melanoma is the most common variant observed in the white population. The back is the most common site for melanoma in men. In women, the back and the lower leg (from knee to ankle) are common sites. Nodular melanomas are dark brown-black to blue-black nodules. Lentigo maligna melanoma usually is confined to chronically sun-damaged sites in older individuals. Acral lentiginous melanoma occurs on the palms, soles, nail beds, and mucous membranes. Although this type occurs in whites, it occurs most frequently (along with nodular melanoma) in blacks and East Asians. A fifth type of melanoma, *desmoplastic melanoma*, is associated with a fibrotic response, neural invasion, and a greater tendency for local recurrence. Occasionally, melanomas appear clinically to be amelanotic, in which case the diagnosis is established microscopically after biopsy of a new or a changing skin nodule. Melanomas can also arise in the mucosa of the head and neck (nasal cavity, paranasal sinuses and oral cavity), the gastrointestinal tract, the CNS, the female genital tract (vulva, vagina), and the uveal tract of the eye.

Although cutaneous melanoma subtypes are clinically and histopathologically distinct, this classification does not have independent prognostic value. Histologic subtype is not part of American Joint Committee on Cancer (AJCC) staging, although the College of American Pathologists (CAP) recommends inclusion in the pathology report. Newer classifications will increasingly emphasize molecular features of each melanoma (see below). The molecular analysis of individual melanomas will provide a basis for distinguishing benign nevi from melanomas, and determination of the mutational status of the tumor will help elucidate the molecular mechanisms of tumorigenesis and be used to identify targets that will guide therapy.

PATHOGENESIS AND MOLECULAR CLASSIFICATION

Considerable evidence from epidemiologic and molecular studies suggests that cutaneous melanomas arise via multiple causal pathways. There are both environmental and genetic components. UV solar radiation causes genetic changes in the skin, impairs cutaneous immune function, increases the production of growth factors, and induces the formation of DNA-damaging reactive oxygen species that affect keratinocytes and melanocytes. A comprehensive catalog of somatic mutations from a human melanoma revealed more than 33,000 base mutations with damage to almost 300 protein-coding segments compared with normal cells from the same patient. The dominant mutational signature reflected DNA damage due to UV light exposure. The melanoma also contained previously described driver mutations (i.e., mutations that confer selective clonal growth advantage and are implicated in oncogenesis). These driver mutations affect pathways that promote cell proliferation and inhibit normal pathways of apoptosis in response to DNA repair (see below). The altered melanocytes accumulate DNA damage, and selection occurs for all the attributes that constitute the malignant phenotype: invasion, metastasis, and angiogenesis.

An understanding of the molecular changes that occur during the transformation of normal melanocytes into malignant melanoma would not only help classify patients but also would contribute to the understanding of etiology and aid the development of new therapeutic options. A genome-wide assessment of melanomas classified into four groups based on their location and degree of exposure to the sun has confirmed that there are distinct genetic pathways in the development of melanoma. The four groups were cutaneous melanomas on skin without chronic sun-induced damage, cutaneous melanomas with chronic sun-induced damage, mucosal melanomas, and acral melanomas. Distinct patterns of DNA alterations were noted that varied with the site of origin and were independent of the histologic subtype of the tumor. Thus, although the genetic changes are diverse, the overall pattern of mutation, amplification, and loss of cancer genes indicates they have convergent effects on key biochemical pathways involved in proliferation, senescence, and apoptosis. The *p16* mutation that affects cell cycle arrest and the *ARF* mutation that results in defective apoptotic responses to genotoxic damage were described earlier. The proliferative pathways affected were the mitogen-activated protein (MAP) kinase and phosphatidylinositol 3' kinase/AKT pathways (Fig. 105-2).

TABLE 105-2 HISTOLOGIC SUBTYPES OF MALIGNANT MELANOMA

Type	Site	Average Age at Diagnosis, Years	Duration of Known Existence, Years	Color
Lentigo maligna melanoma	Sun-exposed surfaces, particularly malar region of cheek and temple	70	5–20 or longer ^a	In flat portions, shades of brown and tan predominate, but whitish gray occasionally present; in nodules, shades of reddish brown, bluish gray, bluish black
Superficial spreading melanoma	Any site (more common on upper back and, in women, lower legs)	40–50	1–7	Shades of brown mixed with bluish red (violaceous), bluish black, reddish brown, and often whitish pink, and the border of lesion is at least in part visibly and/or palpably elevated
Nodular melanoma	Any	40–50	Months–<5 years	Reddish blue (purple) or bluish black; either uniform in color or mixed with brown or black
Acral lentiginous melanoma	Palm, sole, nail bed, mucous membrane	60	1–10	In flat portions, dark brown predominantly; in raised lesions (plaques), brown-black or blue-black predominantly

^aDuring much of this time, the precursor stage, lentigo maligna, is confined to the epidermis.

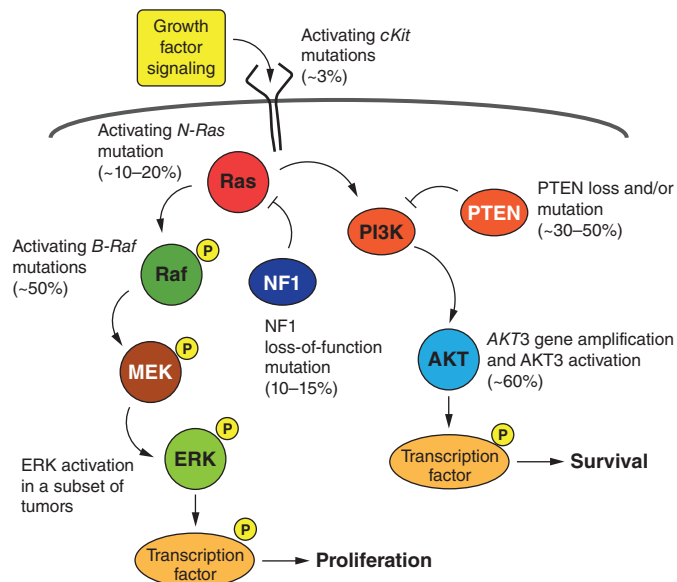


FIGURE 105-2 Major pathways involved in melanoma. The MAP kinase and PI3K/AKT pathways, which promote proliferation and inhibit apoptosis, respectively, are subject to mutations in melanoma. ERK, extracellular signal-regulated kinase; MEK, mitogen-activated protein kinase kinase; NF-1; neurofibromatosis type 1 gene; PTEN, phosphatase and tensin homolog.

RAS and BRAF, members of the MAP kinase pathway, which classically mediates the transcription of genes involved in cell proliferation and survival, undergo somatic mutation in melanoma and thereby generate potential therapeutic targets. N-RAS is mutated in approximately 20% of melanomas, and somatic activating BRAF mutations are found in most benign nevi and 40–60% of melanomas. Neither mutation by itself appears to be sufficient to cause melanoma; thus, they often are accompanied by other mutations. The BRAF mutation is most commonly a point mutation (T→A nucleotide change) that results in a valine-to-glutamate amino acid substitution (V600E). V600E BRAF mutations do not have the standard UV signature mutation (pyrimidine dimer); they are more common in younger patients and are present in most melanomas that arise on sites with intermittent sun exposure and are less common in melanomas from chronically sun-damaged skin.

Melanomas also harbor mutations in AKT (primarily in AKT3) and PTEN (phosphatase and tensin homolog). AKT can be amplified, and PTEN may be deleted or undergo epigenetic silencing that leads to constitutive activation of the PI3K/AKT pathway and enhanced cell survival by antagonizing the intrinsic pathway of apoptosis. Loss of PTEN, which dysregulates AKT activity, and mutation of AKT3 both prolong cell survival through inactivation of BAD, Bcl2-antagonist of cell death, and activation of the forkhead transcription factor FOXO1, which leads to synthesis of prosurvival genes. A loss-of-function mutation in NF1, which can affect both MAP kinase and PI3K/AKT pathways, has been described in 10–15% of melanomas. In melanoma, these two signaling pathways (MAP kinase and PI3K/AKT) enhance tumorigenesis, chemoresistance, migration, and cell cycle dysregulation. Targeted agents that inhibit these pathways have been developed, and some are available for clinical use (see below). Optimal treatment of patients with melanoma may require simultaneous inhibition of both MAPK and PI3K pathways as well as promotion of immune eradication of malignancy.

PROGNOSTIC FACTORS

The prognostic factors of greatest importance to a newly diagnosed patient are included in the staging classification (Table 105-3). The best predictor of metastatic risk is the lesion's Breslow thickness. The Clark level, which defines melanomas on the basis of the layer

of skin to which a melanoma has invaded, does not add significant prognostic information and has minimal influence on treatment decisions. The anatomic site of the primary is also prognostic; favorable sites are the forearm and leg (excluding the feet), and unfavorable sites include the scalp, hands, feet, and mucous membranes. In general, women with stage I or II disease have better survival than men, perhaps in part because of earlier diagnosis; women frequently have melanomas on the lower leg, where self-recognition is more likely and the prognosis is better. The effect of age is not straightforward. Older individuals, especially men over 60, have worse prognoses, a finding that has been explained in part by a tendency toward later diagnosis (and thus thicker tumors) and in part by a higher proportion of acral melanomas in men. However, there is a greater risk of lymph node metastasis in young patients. Other important adverse factors recognized via the staging classification include high mitotic rate, presence of ulceration, microsatellite lesions and/or in-transit metastases, evidence of nodal involvement, elevated serum lactate dehydrogenase (LDH), and presence and site of distant metastases.

STAGING

Once the diagnosis of melanoma has been made, the tumor must be staged to determine the prognosis and treatment. Staging helps determine prognosis and aids in treatment selection. The current melanoma staging criteria and estimated 15-year survival by stage are depicted in Table 105-3. The clinical stage of the patient is determined after the pathologic evaluation of the melanoma skin lesion and clinical/radiologic assessment for metastatic disease. Pathologic staging also includes the microscopic evaluation of the regional lymph nodes obtained at sentinel lymph node biopsy or completion lymphadenectomy as indicated. All patients should have a complete history, with attention to symptoms that may represent metastatic disease such as malaise, weight loss, headaches, visual changes, and pain, and physical examination directed to the site of the primary melanoma, looking for persistent disease or for dermal or subcutaneous nodules that could represent satellite or in-transit metastases, and to the regional draining lymph nodes, CNS, liver, and lungs. A complete blood count (CBC), complete metabolic panel, and LDH should be performed. Although these are low-yield tests for uncovering occult metastatic disease, a microcytic anemia would raise the possibility of bowel metastases, particularly in the small bowel, and an unexplained elevated LDH should prompt a more extensive evaluation, including computed tomography (CT) scan or possibly a positron emission tomography (PET) (or CT/PET combined) scan. If signs or symptoms of metastatic disease are present, appropriate diagnostic imaging should be performed. At initial presentation, more than 80% of patients will have disease confined to the skin and a negative history and physical exam, in which case imaging is not indicated.

TREATMENT MELANOMA

MANAGEMENT OF CLINICALLY LOCALIZED MELANOMA (STAGE I, II)

For a newly diagnosed cutaneous melanoma, wide surgical excision of the lesion with a margin of normal skin is necessary to remove all malignant cells and minimize possible local recurrence. The following margins are recommended for a primary melanoma: in situ, 0.5–1.0 cm; invasive up to 1 mm thick, 1 cm; >1.01–2 mm, 1–2 cm; and >2 mm, 2 cm. For lesions on the face, hands, and feet, strict adherence to these margins must give way to individual considerations about the constraints of surgery and minimization of morbidity. In all instances, however, inclusion of subcutaneous fat in the surgical specimen facilitates adequate thickness measurement and assessment of surgical margins by the pathologist. Topical imiquimod also has been used, particularly for lentigo maligna, in cosmetically sensitive locations.

Sentinel lymph node biopsy (SLNB) is a valuable staging tool that has replaced elective regional nodal dissection for the evaluation of regional nodal status. SLNB provides prognostic information and

TABLE 105-3 STAGING CRITERIA FOR MELANOMA

Pathologic and TNM Stage	Thickness, mm	Ulceration	No. of Involved Lymph Nodes	Nodal Involvement	15-Year Survival Estimate (%)
0					98
Tis	In situ	No	0	None	
IA					92
T1a	<1	No, mitosis <1/mm	0	None	
IB					80
T1b	<1	Yes or mitosis >1/mm	0	None	
T2a	1.01–2	No	0	None	
IIA					62
T2b	1.01–2	Yes	0	None	
T3a	2.01–4	No	0	None	
IIB					51
T3b	2.01–4	Yes	0	None	
T4a	>4	No	0	None	
IIC					37
T4b	>4	Yes	0	None	
IIIA					68
N1a	T1–4a	No	1	Microscopic	
N2a	T1–4a	No	2 or 3	Microscopic	
IIIB					38
N1a	Any	Yes	1	Microscopic	
N2a	Any	Yes	2 or 3	Microscopic	
N1b	Any	Yes or no	1	Macroscopic	
N2b	Any	Yes or no	2 or 3	Macroscopic	
N2c	Any	Yes or no	In-transit metastases/satellites, no nodal involvement		
IIIC					22
N1b	Any	Yes or no	1	Macroscopic	
N2b	Any	Yes or no	2 or 3	Macroscopic	
N2c	Any	Yes or no	In-transit metastases/satellites, no nodal involvement		
N3	Any	Yes or no	4+ metastatic nodes, matted nodes or in-transit metastases/satellites, with metastatic nodes		
IV		Distant metastasis			<10
M1a		Skin, subcutaneous			
M1b		Lung			
M1c		Other visceral site			
		Elevated lactate dehydrogenase			

helps identify patients at high risk for relapse who may be candidates for adjuvant therapy. The initial (sentinel) draining node(s) from the primary site is (are) identified by injecting a blue dye and a radioisotope around the primary site. The sentinel node(s) then is (are) identified by inspection of the nodal basin for the blue-stained node and/or the node with high uptake of the radioisotope. The identified nodes are removed and subjected to careful histopathologic analysis with serial section using hematoxylin and eosin stains as well as immunohistochemical stains (e.g., S100, HMB45, and MelanA) to identify melanocytes.

Not every patient requires a SLNB. Patients whose melanomas are ≤0.75 mm thick have <5% risk of sentinel lymph node (SLN) disease and do not require a SLNB. Patients with tumors >1 mm thick generally undergo SLNB. For melanomas 0.76–1.0 mm thick, SLNB may be considered for lesions with high-risk features such as ulceration, high mitotic index, or lymphovascular invasion, but wide excision alone is the usual definitive therapy. Most other patients with clinically negative lymph nodes should undergo a SLNB. Patients

whose SLNB is negative are spared a complete node dissection and its attendant morbidities, and can simply be followed or, based on the features of the primary melanoma, be considered for adjuvant therapy or a clinical trial. The current standard of care for all patients with a positive SLN is to perform a complete lymphadenectomy; however, ongoing clinical studies will determine whether patients with small-volume SLN metastases can be managed safely without additional surgery. Patients with microscopically positive lymph nodes should be considered for adjuvant therapy with interferon or enrollment in a clinical trial.

MANAGEMENT OF REGIONALLY METASTATIC MELANOMA (STAGE III)

Melanomas may recur at the edge of the scar or graft, as satellite metastases, which are separate from but within 2 cm of the scar; as in-transit metastases, which are recurrences >2 cm from the primary lesion but not beyond the regional nodal basin; or, most commonly, as metastasis to a draining lymph node basin. Each of these presentations is managed surgically, following which there

498 is the possibility of long-term disease-free survival. Isolated limb perfusion or infusion with melphalan and hyperthermia are options for patients with extensive cutaneous regional recurrences in an extremity. High complete response rates have been reported and significant palliation of symptoms can be achieved, but there is no change in overall survival.

Patients rendered free of disease after surgery may be at high risk for a local or distant recurrence and should be considered for adjuvant therapy. Radiotherapy can reduce the risk of local recurrence after lymphadenectomy, but does not affect overall survival. Patients with large nodes (>3–4 cm), four or more involved lymph nodes, or extranodal spread on microscopic examination should be considered for radiation. Systemic adjuvant therapy is indicated primarily for patients with stage III disease, but high-risk, node-negative patients (>4 mm thick or ulcerated lesions) and patients with completely resected stage IV disease also may benefit. Either interferon $\alpha 2b$ (IFN- $\alpha 2b$), which is given at 20 million units/m² IV 5 days a week for 4 weeks followed by 10 million units/m² SC three times a week for 11 months (1 year total), or subcutaneous peginterferon $\alpha 2b$ (6 μ g/kg per week for 8 weeks followed by 3 μ g/kg per week for a total of 5 years) is acceptable adjuvant therapy. Treatment is accompanied by significant toxicity, including a flu-like illness, decline in performance status, and the development of depression. Side effects can be managed in most patients by appropriate treatment of symptoms, dose reduction, and treatment interruption. Sometimes IFN must be permanently discontinued before all of the planned doses are administered because of unacceptable toxicity. The high-dose regimen is significantly more toxic than peginterferon, but the latter requires 4 additional years of therapy. Adjuvant treatment with IFN improves disease-free survival, but its impact on overall survival remains controversial. Enrollment in a clinical trial is appropriate for these patients, many of whom will otherwise be observed without treatment either because they are poor candidates for IFN or because the patient (or their oncologist) does not believe the beneficial effects of IFN outweigh the toxicity. The recently approved immunotherapy and targeted agents are being evaluated in the adjuvant setting.

TREATMENT METASTATIC DISEASE

At diagnosis, most patients with melanoma will have early-stage disease; however, some will present with metastases, and others will develop metastases after initial therapy. Patients with a history of melanoma who develop signs or symptoms suggesting recurrent disease should undergo restaging that includes physical examination, CBC, complete metabolic panel, LDH, and appropriate diagnostic imaging that may include a magnetic resonance image (MRI) of the brain and total-body PET/CT or CT scans of the chest, abdomen, and pelvis. Distant metastases (stage IV), which may involve any organ, commonly involve the skin and lymph nodes as well as viscera, bone, or the brain. Historically, metastatic melanoma was considered incurable; median survival ranges from 6 to 15 months, depending on the organs involved. The prognosis is better for patients with skin and subcutaneous metastases (M1a) than for lung (M1b) and worst for those with metastases to liver, bone, and brain (M1c). An elevated serum LDH is a poor prognostic factor and places the patient in stage M1c regardless of the site of the metastases (Table 105-3). Although historical data suggest that the 15-year survival of patients with M1a, M1b, and M1c disease is less than 10%, there is optimism that newer therapies will increase the number of melanoma patients with long-term survival, especially patients with M1a and M1b disease.

The treatment for patients with stage IV melanoma has changed dramatically in the past 2 years. Two new classes of therapeutic agents for melanoma have been approved by the U.S. Food and Drug Administration (FDA). The immune T cell checkpoint inhibitor, ipilimumab, and three new oral agents that target the MAP kinase pathway: the BRAF inhibitors, vemurafenib and dabrafenib, and the

TABLE 105-4 TREATMENT OPTIONS FOR METASTATIC MELANOMA

Surgery: Metastasectomy for small number of lesions
Immunotherapy:
Interleukin 2
Immune checkpoint blockade
- FDA approved
▪ Anti-CTLA-4: ipilimumab
- Experimental
▪ Anti-PD-1: nivolumab, lambrolizumab
▪ Anti-PD-L1
Molecular targeted therapy:
BRAF inhibitor: vemurafenib, dabrafenib
MEK inhibitor: trametinib
Chemotherapy: dacarbazine, temozolomide, paclitaxel, albumin-bound paclitaxel (Abraxane), carboplatin

MEK inhibitor, trametinib, are now available, so patients with stage IV disease now have multiple therapeutic options (Table 105-4).

Patients with oligometastatic disease should be referred to a surgical oncologist for consideration of metastasectomy, because they may experience long-term disease-free survival after surgery. Patients with solitary metastases are the best candidates, but surgery increasingly is being used even for patients with metastases at more than one site. Patients rendered free of disease can be considered for IFN therapy or a clinical trial because their risk of developing additional metastases is very high. Surgery can also be used as an adjunct to immunotherapy when only a few of many metastatic lesions prove resistant to systemic therapy.

IMMUNOTHERAPY

The cytokine interleukin 2 (IL-2 or aldesleukin) has been approved to treat patients with melanoma since 1995. IL-2 is used to treat stage IV patients who have a good performance status and is administered at centers with experience managing IL-2-related toxicity. Patients require hospitalization in an intensive care unit-like setting to receive high-dose IL-2 600,000 or 720,000 IU every 8 h for up to 14 doses (one cycle). Patients continue treatment until they achieve maximal benefit, usually 4–6 cycles. Treatment is associated with long-term disease-free survival (probable cures) in 5% of treated patients. The mechanism by which IL-2 causes tumor regression has not been identified, but it is presumed that IL-2 induces melanoma-specific T cells that eliminate tumor cells by recognizing specific antigens. Rosenberg and his colleagues at the National Cancer Institute (NCI) have combined adoptive transfer of in vitro-expanded tumor-infiltrating lymphocytes with high-dose IL-2 in patients who were preconditioned with nonmyeloablative chemotherapy (sometimes combined with total-body irradiation). Tumor regression was observed in more than 50% of patients with IL-2-refractory metastatic melanoma.

Immune checkpoint blockade with monoclonal antibodies to the inhibitory immune receptors CTLA-4 and PD-1 has shown promising clinical efficacy. An array of inhibitory receptors are upregulated during an immune response. An absolute requirement to ensure proper regulation of a normal immune response, the continued expression of inhibitory receptors during chronic infection (hepatitis, HIV) and in cancer patients denotes exhausted T cells with limited potential for proliferation, cytokine production, or cytotoxicity (Fig. 105-3). Checkpoint blockade with a monoclonal antibody results in improved T cell function with eradication of tumor cells in preclinical animal models. Ipilimumab, a fully human IgG antibody that binds CTLA-4 and blocks inhibitory signals, was the first treatment of any kind to improve survival in patients with metastatic melanoma. A full course of therapy is four IV outpatient infusions of ipilimumab 3 mg/kg every 3 weeks. Although response rates were low (~10%) in randomized clinical trials, survival of both previously treated and untreated patients was improved, and ipilimumab was approved by the FDA in March 2011.

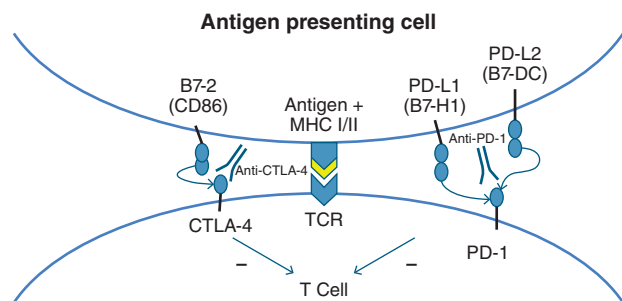


FIGURE 105-3 Inhibitory regulatory pathways that influence T cell function, memory, and lifespan after engagement of the T cell receptor by antigen presented by antigen-presenting cells in the context of MHC I/II. CTLA-4 and PD-1 are part of the CD28 family and have inhibitory effects that can be mitigated by antagonistic antibodies to the receptors or ligand, resulting in enhanced T cell function and antitumor effects. CTLA-4, cytotoxic T lymphocyte antigen-4; MHC, major histocompatibility complex; PD-1, programmed death-1; PD-L1, programmed death ligand-1; PD-L2, programmed death ligand-2; TCR, T cell receptor.

In addition to its antitumor effects, ipilimumab's interference with normal regulatory mechanisms produced a novel spectrum of side effects that resembled autoimmunity. The most common immune-related adverse events were skin rash and diarrhea (sometimes severe, life-threatening colitis), but toxicity could involve most any organ (e.g., hypophysitis, hepatitis, nephritis, pneumonitis, myocarditis, neuritis). Vigilance and early treatment with steroids that do not appear to interfere with the antitumor effects are required to manage these patients safely. Widespread use of ipilimumab has not been completely embraced by the oncology community because of the low objective response rate, significant toxicity (including death), and high cost (drug cost alone for a course of therapy is approximately \$120,000 in 2013). Despite these reservations, ipilimumab's overall survival benefit (17% of patients alive at 7 years) indicates that treatment should be strongly considered for all eligible patients.

Chronic T cell activation also leads to induction of PD-1 on the surface of T cells. Expression of one of its ligands, PD-L1, on tumor cells can protect them from immune destruction (Fig. 105-3). Early trials attempting to block the PD-1:PD-L1 axis by IV administration of anti-PD-1 or anti-PD-L1 have shown substantial clinical activity in patients with advanced melanoma (and lung cancer) with significantly less toxicity than ipilimumab. Anti-PD-1 therapy looks promising, but is not currently available except by participation in clinical trials. Intriguingly, preliminary results from a clinical trial indicate that blocking both inhibitory pathways with ipilimumab and anti-PD-1 leads to superior antitumor activity than treatment with either agent alone. The main benefit to patients from immune-based therapy (IL-2, ipilimumab, and anti-PD-1) is the durability of the responses achieved. Although the percentage of patients whose tumors regress following immunotherapy is lower than the response rate after targeted therapy (see below), the durability of immunotherapy-induced responses (>10 years in some cases) appears to be superior to responses after targeted therapy and suggests that many of these patients have been cured.

TARGETED THERAPY

RAF and MEK inhibitors of the MAP kinase pathway are a new and exciting approach for patients whose melanomas harbor a *BRAF* mutation. The high frequency of oncogenic mutations in the RAS-RAF-MEK-ERK pathway, which delivers proliferation and survival signals from the cell surface to the cytoplasm and nucleus, has led to the development of inhibitors to BRAF and MEK. Two BRAF inhibitors, vemurafenib and dabrafenib, have been approved for the treatment of stage IV patients whose melanomas harbor a mutation

at position 600 in the gene for *BRAF*. The oral BRAF inhibitors cause tumor regression in approximately 50% of patients, and overall survival is improved compared to treatment with chemotherapy. Treatment is accompanied by manageable side effects that differ from those following immunotherapy or chemotherapy. A class-specific complication of BRAF inhibition is the development of numerous skin lesions, some of which are well-differentiated squamous cell skin cancers (seen in up to a quarter of patients). Patients should be co-managed with a dermatologist as these skin cancers will need excision. Metastases have not been reported, and treatment can be continued safely following simple excision. Long-term results following treatment with BRAF inhibitors are not yet available, but the current concern is that over time the vast majority of patients will relapse and eventually die from drug-resistant disease. There are a number of mechanisms by which resistance develops, usually via maintenance of MAP kinase signaling; however, mutations in the *BRAF* gene that affect binding of the inhibitor are not among them. The MEK inhibitor trametinib has activity as a single agent, but appears to be less effective than either of the BRAF inhibitors. Combined therapy with the BRAF inhibitor and MEK inhibitor showed improved progression-free survival compared to BRAF inhibitor therapy alone; and, interestingly, the neoplastic skin lesions that were so troubling with BRAF inhibition alone did not occur. Although the durability of responses following combined therapy remains to be determined, its use in metastatic melanoma is FDA approved. Activating mutations in the *c-kit* receptor tyrosine kinase are found in a minority of cutaneous melanomas with chronic sun damage, but more commonly in mucosal and acral lentiginous subtypes. Overall, the number of patients with *c-kit* mutations is exceedingly small, but when present, they are largely identical to mutations found in gastrointestinal stromal tumors (GISTs); melanomas with activating *c-kit* mutations can have clinically meaningful responses to imatinib.

CHEMOTHERAPY

No chemotherapy regimen has ever been shown to improve survival in metastatic melanoma, and the advances in immunotherapy and targeted therapy have relegated chemotherapy to the palliation of symptoms. Drugs with antitumor activity include dacarbazine (DTIC) or its orally administered analog temozolomide (TMZ), cisplatin and carboplatin, the taxanes (paclitaxel alone or albumin-bound and docetaxel), and carmustine (BCNU), which have reported response rates of 12–20%.

INITIAL APPROACH TO PATIENT WITH METASTATIC DISEASE

Upon diagnosis of stage IV disease, whether by biopsy or diagnostic imaging, a sample of the patient's tumor needs to undergo molecular testing to determine whether a druggable mutation (e.g., *BRAF*) is present. Analysis of a metastatic lesion is preferred, but any biopsy will suffice because there is little discordance between primary and metastatic lesions. Treatment algorithms start with the tumor's *BRAF* status. For *BRAF* "wild-type" tumors, immunotherapy is recommended. For patients whose tumors harbor a *BRAF* mutation, initial therapy with either a BRAF inhibitor or immunotherapy is acceptable. Molecular testing may also include *N-RAS* and *c-kit* in appropriate tumors.

The majority of patients still die from their melanoma, despite improvements in therapy. Therefore, enrollment in a clinical trial is always an important consideration, even for previously untreated patients. Most patients with stage IV disease will eventually progress despite advances in therapy, and many, because of disease burden, poor performance status, or concomitant illness, will be unsuitable for therapy. Therefore, a major focus of care should be the timely integration of palliative care and hospice.

FOLLOW-UP

Skin examination and surveillance at least once a year are recommended for all patients with melanoma. The National Comprehensive Cancer Network (NCCN) guidelines for patients with stage IA–IIA

500 melanoma recommend a comprehensive history and physical examination every 6–12 months for 5 years, and then annually as clinically indicated. Particular attention should be paid to the draining lymph nodes in stage I–III patients as resection of lymph node recurrences may still be curative. A CBC, LDH, and chest x-ray are recommended at the physician's discretion, but are ineffective tools for the detection of occult metastases. Routine imaging for metastatic disease is not recommended at this time. For patients with higher stage disease (IIB–IV), imaging (chest x-ray, CT, and/or PET/CT scans) every 4–12 months can be considered. Because no discernible survival benefit has been demonstrated for routine surveillance, it is reasonable to perform scans only if clinically indicated.

NONMELANOMA SKIN CANCER

Nonmelanoma skin cancer (NMSC) is the most common cancer in the United States. Although tumor registries do not routinely gather data on the incidence of basal cell and squamous cell skin cancers, it is estimated that the annual incidence is 1.5–2 million cases in the United States. Basal cell carcinomas (BCCs) account for 70–80% of NMSCs. Squamous cell carcinomas (SCCs), which comprise ~20% of NMSCs, are more significant because of their ability to metastasize and account for 2400 NMSC deaths annually. There has also been an increase in the incidence of nonepithelial skin cancer, especially Merkel cell carcinoma, with nearly 5000 new diagnoses and 3000 deaths annually.

PATHOPHYSIOLOGY AND ETIOLOGY

The most significant cause of BCC and SCC is UV exposure, whether through direct exposure to sunlight or by artificial UV light sources (tanning beds). Both UVA and UVB can induce DNA damage through free radical formation (UVA) or induction of pyrimidine dimers (UVB). The sun emits energy across the UV spectrum, whereas tanning bed equipment typically emits 97% UVA and 3% UVB. DNA damage induced by UV irradiation can result in cell death or repair of damaged DNA by nucleotide excision repair (NER). Inherited disorders of NER, such as xeroderma pigmentosum, are associated with a greatly increased incidence of skin cancer and help to establish the link between UV-induced DNA damage, inadequate DNA repair, and skin cancer. The genes damaged most commonly by UV in BCC involve the Hedgehog pathway (Hh). In SCC, *p53* and *N-RAS* are commonly affected. There is a dose-response relationship between tanning bed use and the incidence of skin cancer. As few as four tanning bed visits per year confers a 15% increase in BCC and an 11% increase in SCC and melanoma. Tanning bed use as a teenager or young adult confers greater risk than comparable exposure in older individuals. Other associations include blond or red hair, blue or green eyes, a tendency to sunburn easily, and an outdoor occupation. The incidence of NMSC increases with decreasing latitude. Most tumors develop on sun-exposed areas of the head and neck. The risk of lip or oral SCC is increased with cigarette smoking. Human papillomaviruses and UV radiation may act as cocarcinogens.

Solid organ transplant recipients on chronic immunosuppression have a 65-fold increase in SCC and a 10-fold increase in BCC. The frequency of skin cancer is proportional to the level and duration of immunosuppression and the extent of sun exposure before and after transplantation. SCCs in this population also demonstrate higher rates of local recurrence, metastasis, and mortality. There is increasing use of tumor necrosis factor (TNF) antagonists to treat inflammatory bowel disease and autoimmune disorders such as rheumatoid and psoriatic arthritis. TNF antagonists may also confer an increased risk of NMSC. BRAF-targeted therapy can induce SCCs including keratoacanthoma-type SCCs in keratinocytes, with preexisting *H-RAS* overexpression present in approximately 60% of patients.

Other risk factors include HIV infection, ionizing radiation, thermal burn scars, and chronic ulcerations. Albinism, xeroderma pigmentosum, Muir-Torre syndrome, Rombo's syndrome, Bazex-Dupré-Christol syndrome, dyskeratosis congenita, and basal cell nevus syndrome (Gorlin syndrome) also increase the incidence of NMSC. Mutations in Hh genes encoding the tumor-suppressor patched

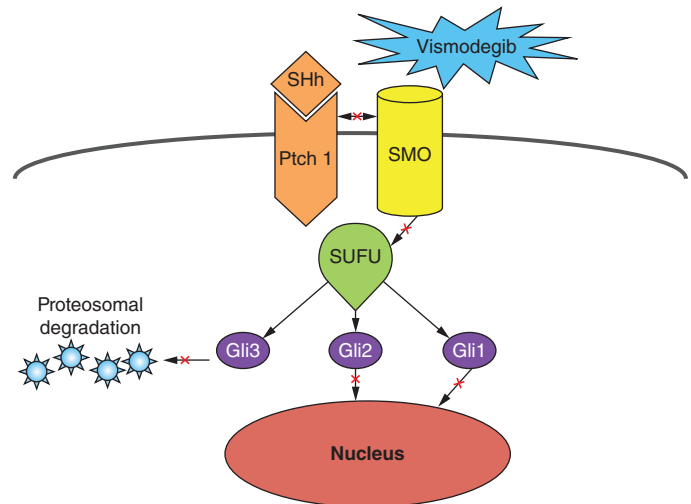


FIGURE 105-4 Influence of vismodegib on the hedgehog (Hh) pathway. Normally, one of three Hh ligands (sonic [SHh], Indian, or desert) binds to patched homolog 1 (PTCH1), causing its degradation and release of smoothed homolog (SMO). The downstream events of SMO release are the activation of Gli1, Gli2, and Gli3 through the transcriptional regulator known as SUFU. Gli1 and Gli2 translocate to the nucleus and promote gene transcription. Vismodegib is an SMO antagonist that decreases the interaction between SMO and PTCH1, resulting in decreased Hh pathway signaling, gene transcription, and cell division. The downstream Hh pathway events inhibited by vismodegib are indicated in red.

homolog 1 (*PTCH1*) and smoothed homolog (*SMO*) occur in BCC. Aberrant *PTCH1* signaling is propagated by the nuclear transcription factors Gli1 and Gli2, which are salient in the development of BCC and have led to the FDA approval of an oral SMO inhibitor, vismodegib, to treat advanced inoperable or metastatic BCC (Fig. 105-4). Vismodegib also reduces the incidence of BCC in patients with basal cell nevus syndrome who have *PTCH1* mutations, affirming the importance of Hh in the onset of BCC.

CLINICAL PRESENTATION

Basal Cell Carcinoma BCC arises from epidermal basal cells. The least invasive of BCC subtypes, superficial BCC, consists of often subtle, erythematous scaling plaques that slowly enlarge and are most commonly seen on the trunk and proximal extremities (Fig. 105-5). This BCC subtype may be confused with benign inflammatory dermatoses, especially nummular eczema and psoriasis. BCC also can present as a small, slowly growing pearly nodule, often with tortuous telangiectatic vessels on its surface, rolled borders, and a central crust (nodular BCC). The occasional presence of melanin in this variant of nodular BCC (pigmented BCC) may lead to confusion with melanoma. Morpheaform (fibrosing), infiltrative, and micronodular BCC, the most invasive and potentially aggressive subtypes, manifest as solitary, flat or slightly depressed, indurated whitish, yellowish, or pink scar-like plaques. Borders are typically indistinct, and lesions can be subtle; thus, delay in treatment is common, and tumors can be more extensive than expected clinically.

Squamous Cell Carcinoma Primary *cutaneous* SCC is a malignant neoplasm of keratinizing epidermal cells. SCC has a variable clinical course, ranging from indolent to rapid growth kinetics, with the potential for metastasis to regional and distant sites. Commonly, SCC appears as an ulcerated erythematous nodule or superficial erosion on sun-exposed skin of the head, neck, trunk, and extremities (Fig. 105-5). It may also appear as a banal, firm, dome-shaped papule or rough-textured plaque. It is commonly mistaken for a wart or callous when the inflammatory response to the lesion is minimal. Clinically visible overlying telangiectasias are uncommon, although dotted or coiled vessels are a hallmark of SCC when viewed through a dermatoscope.

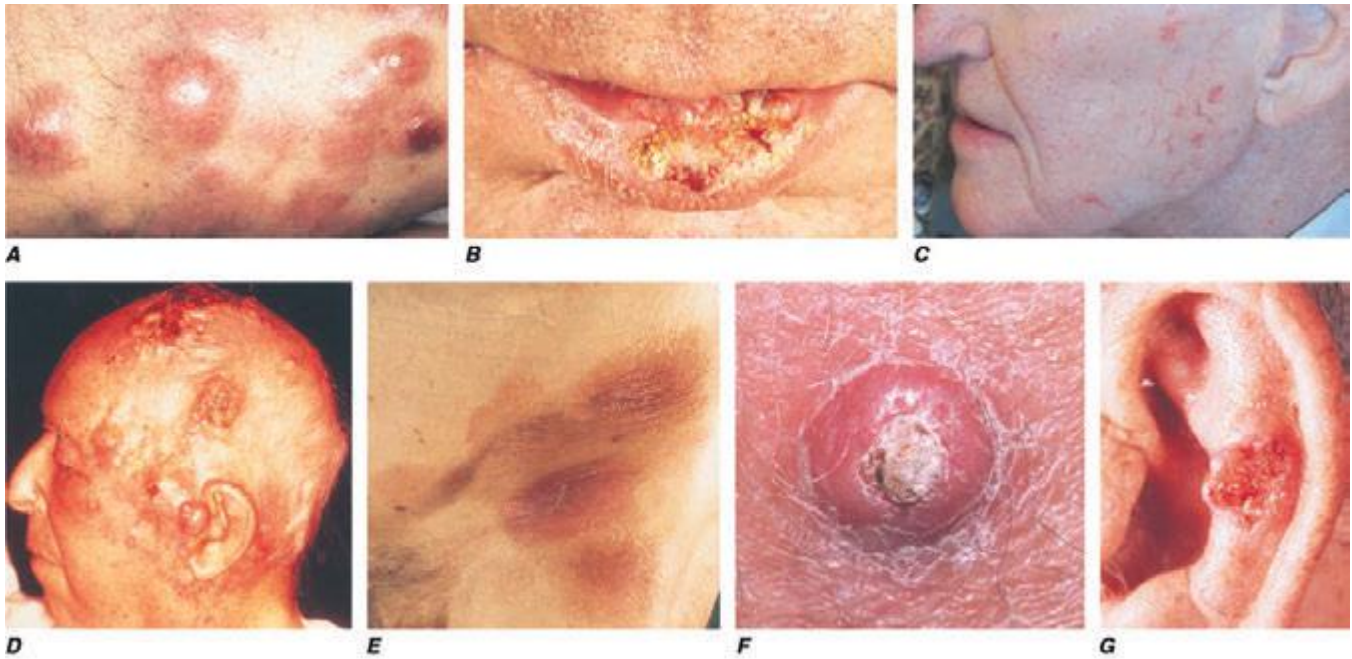


FIGURE 105-5 Cutaneous neoplasms. **A.** Non-Hodgkin's lymphoma involves the skin with typical violaceous, "plum-colored" nodules. **B.** Squamous cell carcinoma is seen here as a hyperkeratotic crusted and somewhat eroded plaque on the lower lip. Sun-exposed skin in areas such as the head, neck, hands, and arms represent other typical sites of involvement. **C.** Actinic keratoses consist of hyperkeratotic erythematous papules and patches on sun-exposed skin. They arise in middle-aged to older adults and have some potential for malignant transformation. **D.** Metastatic carcinoma to the skin is characterized by inflammatory, often ulcerated dermal nodules. **E.** Mycosis fungoides is a cutaneous T cell lymphoma, and plaque-stage lesions are seen in this patient. **F.** Keratoacanthoma is a low-grade squamous cell carcinoma that presents as an exophytic nodule with central keratinous debris. **G.** This basal cell carcinoma shows central ulceration and a pearly, rolled telangiectatic tumor border.

The margins of this tumor may be ill defined, and fixation to underlying structures may occur ("tethering").

A very rapidly growing but low-grade form of SCC, called keratoacanthoma (KA), typically appears as a large dome-shaped papule with a central keratotic crater. Some KAs regress spontaneously without therapy, but because progression to metastatic SCC has been documented, KAs should be treated in the same manner as other types of cutaneous SCC. KAs are also associated with medications that target *BRAF* mutations and occur in 15–25% of patients receiving these medications.

Actinic keratoses and *cheilitis* (actinic keratoses occurring on the lip), both premalignant forms of SCC, present as hyperkeratotic papules on sun-exposed areas. The potential for malignant degeneration in untreated lesions ranges from 0.25 to 20%. SCC in situ, also called *Bowen's disease*, is the intraepidermal form of SCC and usually presents as a scaling, erythematous plaque. As with invasive SCC, SCC in situ most commonly arises on sun-damaged skin, but can occur anywhere on the body. Bowen's disease forming secondary to infection with human papillomavirus (HPV) can arise on skin with minimal or no prior sun exposure, such as the buttock or posterior thigh. Treatment of premalignant and in situ lesions reduces the subsequent risk of invasive disease.

NATURAL HISTORY

Basal Cell Carcinoma The natural history of BCC is that of a slowly enlarging, locally invasive neoplasm. The degree of local destruction and risk of recurrence vary with the size, duration, location, and histologic subtype of the tumor. Location on the central face, ears, or scalp may portend a higher risk. Small nodular, pigmented, cystic, or superficial BCCs respond well to most treatments. Large lesions and micronodular, infiltrative, and morpheaform subtypes may be more aggressive. The metastatic potential of BCC is low (0.0028–0.1% in immunocompetent patients), but the risk of recurrence or a new primary NMSC is about 40% over 5 years.

Squamous Cell Carcinoma The natural history of SCC depends on tumor and host characteristics. Tumors arising on sun-damaged

skin have a lower metastatic potential than do those on non-sun-exposed areas. Cutaneous SCC metastasizes in 0.3–5.2% of individuals, most frequently to regional lymph nodes. Tumors occurring on the lower lip and ear develop regional metastases in 13 and 11% of patients, respectively, whereas the metastatic potential of SCC arising in scars, chronic ulcerations, and genital or mucosal surfaces is higher. Recurrent SCC has a much higher potential for metastatic disease, approaching 30%. Large, poorly differentiated, deep tumors with perineural or lymphatic invasion, multifocal tumors, and those arising in immunosuppressed patients often behave aggressively.

TREATMENT BASAL CELL CARCINOMA

Treatments used for BCC include electrodesiccation and curettage (ED&C), excision, cryosurgery, radiation therapy (RT), laser therapy, Mohs micrographic surgery (MMS), topical 5-fluorouracil, photodynamic therapy (PDT), and topical immunomodulators such as imiquimod. The therapy chosen depends on tumor characteristics including depth and location, patient age, medical status, and patient preference. ED&C remains the most commonly employed method for superficial, minimally invasive nodular BCCs and low-risk tumors (e.g., a small tumor of a less aggressive subtype in a favorable location). Wide local excision with standard margins is usually selected for invasive, ill-defined, and more aggressive subtypes of tumors, or for cosmetic reasons. MMS, a specialized type of surgical excision that provides the best method for tumor removal while preserving uninvolved tissue, is associated with cure rates >98%. It is the preferred modality for lesions that are recurrent, in high-risk or cosmetically sensitive locations (including recurrent tumors in these locations), and in which maximal tissue conservation is critical (e.g., the eyelids, lips, ears, nose, and digits). RT can cure patients not considered surgical candidates and can be used as a surgical adjunct in high-risk tumors. Younger patients may not be good candidates

502 for RT because of the risks of long-term carcinogenesis and radio-dermatitis. Imiquimod can be used to treat superficial and smaller nodular BCCs, although it is not FDA-approved for nodular BCC. Topical 5-fluorouracil therapy should be limited to superficial BCC. PDT, which uses selective activation of a photoactive drug by visible light, has been used in patients with numerous tumors. Intravesical chemotherapy (5-fluorouracil and IFN) for NMSC has existed since the mid-twentieth century, but is used so infrequently that recent consensus guidelines for the treatment of BCC and SCC do not include it. Like RT, it remains an option for well-selected patients who cannot or will not undergo surgery.

SQUAMOUS CELL CARCINOMA

Therapy for cutaneous SCC should be based on the size, location, histologic differentiation, patient age, and functional status. Surgical excision and MMS are standard treatments. Cryosurgery and ED&C have been used for premalignant lesions and small, superficial, in situ primary tumors. Lymph node metastases are treated with surgical resection, RT, or both. Systemic chemotherapy combinations that include cisplatin can palliate patients with advanced disease. SCC and keratoacanthomas that develop in patients receiving BRAF-targeted therapy should be excised, but their development should not deter the continued use of BRAF therapy. Retinoid prophylaxis can also be considered for patients receiving BRAF-targeted therapy, although no prospective studies have been completed thus far.

PREVENTION

The general principles for prevention are those described for melanoma earlier. Unique strategies for NMSC include active surveillance for patients on immunosuppressive medications or BRAF-targeted therapy. Chemoprophylaxis using synthetic retinoids and immunosuppression reduction when possible may be useful in controlling new lesions and managing patients with multiple tumors.

OTHER NONMELANOMA CUTANEOUS MALIGNANCIES

Neoplasms of cutaneous adnexae and sarcomas of fibrous, mesenchymal, fatty, and vascular tissues make up the remaining 1–2% of NMSCs.

Merkel cell carcinoma (MCC) is a neural crest–derived highly aggressive malignancy with mortality rates approaching 33% at 3 years. An oncogenic Merkel cell polyomavirus is present in 80% of tumors. Many patients have detectable cellular or humoral immune responses to polyoma viral proteins, although this immune response is insufficient to eradicate the malignancy. Survival depends on extent of disease: 90% survive with local disease, 52% with nodal involvement, and only 10% with distant disease at 3 years. MCC incidence tripled over the last 20 years with an estimated 1600 cases per year in the United States. Immunosuppression can increase incidence and diminish prognosis. MCC lesions typically present as an asymptomatic rapidly expanding bluish-red/violaceous tumor on sun-exposed skin of older white patients. Treatment is surgical excision with sentinel lymph node biopsy for accurate staging in patients with localized disease, often followed by adjuvant RT. Patients with extensive disease can be offered systemic chemotherapy; however, there is no convincing survival benefit. Whenever possible a clinical trial should be considered for this rare but aggressive NMSC, especially in light of the potential for new treatments directed at the oncogenic virus that causes this malignancy.

Extramammary Paget's disease is an uncommon apocrine malignancy arising from stem cells of the epidermis that are characterized histologically by the presence of Paget cells. These tumors present as moist erythematous patches on anogenital or axillary skin of the elderly. Outcomes are generally good with site-directed surgery, and 5-year disease specific survival is approximately 95% with localized disease. Advanced age and extensive disease at presentation are factors that confer diminished prognosis. RT or topical imiquimod can be considered for more extensive disease. Local management may be challenging because these tumors often extend far beyond clinical margins; surgical excision with MMS has the highest cure rates. Similarly,

MMS is the treatment of choice in other rare cutaneous tumors with extensive subclinical extension such as *dermatofibrosarcoma protuberans*.

Kaposi's sarcoma (KS) is a soft tissue sarcoma of vascular origin that is induced by the human herpesvirus 8. The incidence of KS increased dramatically during the AIDS epidemic, but has now decreased tenfold with the institution of highly active antiretroviral therapy.

ACKNOWLEDGMENT


Carl V. Washington, MD, and Hari Nadiminti, MD, contributed to this chapter in the 18th edition, and material from that chapter is included here. Claudia Taylor, MD, and Steven Kolker, MD, provided valued feedback and suggested many improvements to this chapter.

106 Head and Neck Cancer

Everett E. Vokes

Epithelial carcinomas of the head and neck arise from the mucosal surfaces in the head and neck and typically are squamous cell in origin. This category includes tumors of the paranasal sinuses, the oral cavity, and the nasopharynx, oropharynx, hypopharynx, and larynx. Tumors of the salivary glands differ from the more common carcinomas of the head and neck in etiology, histopathology, clinical presentation, and therapy. They are rare and histologically highly heterogeneous. **Thyroid malignancies are described in Chap. 405.**

INCIDENCE AND EPIDEMIOLOGY

 The number of new cases of head and neck cancers (oral cavity, pharynx, and larynx) in the United States was 53,640 in 2013, accounting for about 3% of adult malignancies; 11,520 people died from the disease. The worldwide incidence exceeds half a million cases annually. In North America and Europe, the tumors usually arise from the oral cavity, oropharynx, or larynx. The incidence of oropharyngeal cancers is increasing in recent years. Nasopharyngeal cancer is more commonly seen in the Mediterranean countries and in the Far East, where it is endemic in some areas.

ETIOLOGY AND GENETICS

Alcohol and tobacco use are the most significant risk factors for head and neck cancer, and when used together, they act synergistically. Smokeless tobacco is an etiologic agent for oral cancers. Other potential carcinogens include marijuana and occupational exposures such as nickel refining, exposure to textile fibers, and woodworking.

Some head and neck cancers have a viral etiology. Epstein-Barr virus (EBV) infection is frequently associated with nasopharyngeal cancer, especially in endemic areas of the Mediterranean and Far East. EBV antibody titers can be measured to screen high-risk populations. Nasopharyngeal cancer has also been associated with consumption of salted fish and in-door pollution.

In Western countries, the human papilloma virus (HPV) is associated with a rising incidence of tumors arising from the oropharynx, i.e., the tonsillar bed and base of tongue. Over 50% of oropharyngeal tumors are caused by HPV in the United States. HPV 16 is the dominant viral subtype, although HPV 18 and other oncogenic subtypes are seen as well. Alcohol- and tobacco-related cancers, on the other hand, have decreased in incidence. HPV-related oropharyngeal cancer occurs in a younger patient population and is associated with increased numbers of sexual partners and oral sexual practices. It is associated with a better prognosis, especially for nonsmokers.

Dietary factors may contribute. The incidence of head and neck cancer is higher in people with the lowest consumption of fruits and vegetables. Certain vitamins, including carotenoids, may be protective if included in a balanced diet. Supplements of retinoids, such as *cis*-retinoic acid, have not been shown to prevent head and neck cancers

(or lung cancer) and may increase the risk in active smokers. No specific risk factors or environmental carcinogens have been identified for salivary gland tumors.

HISTOPATHOLOGY, CARCINOGENESIS, AND MOLECULAR BIOLOGY

Squamous cell head and neck cancers are divided into well-differentiated, moderately well-differentiated, and poorly differentiated categories. Poorly differentiated tumors have a worse prognosis than well-differentiated tumors. For nasopharyngeal cancers, the less common differentiated squamous cell carcinoma is distinguished from nonkeratinizing and undifferentiated carcinoma (lymphoepithelioma) that contains infiltrating lymphocytes and is commonly associated with EBV.

Salivary gland tumors can arise from the major (parotid, submandibular, sublingual) or minor salivary glands (located in the submucosa of the upper aerodigestive tract). Most parotid tumors are benign, but half of submandibular and sublingual gland tumors and most minor salivary gland tumors are malignant. Malignant tumors include mucoepidermoid and adenoid cystic carcinomas and adenocarcinomas.

The mucosal surface of the entire pharynx is exposed to alcohol- and tobacco-related carcinogens and is at risk for the development of a premalignant or malignant lesion. Erythroplakia (a red patch) or leukoplakia (a white patch) can be histopathologically classified as hyperplasia, dysplasia, carcinoma in situ, or carcinoma. However, most head and neck cancer patients do not present with a history of premalignant lesions. Multiple synchronous or metachronous cancers can also be observed. In fact, over time, patients with early-stage head and neck cancer are at greater risk of dying from a second malignancy than from a recurrence of the primary disease.

Second head and neck malignancies are usually not therapy-induced; they reflect the exposure of the upper aerodigestive mucosa to the same carcinogens that caused the first cancer. These second primaries develop in the head and neck area, the lung, or the esophagus. Thus, computed tomography (CT) screening for lung cancer in heavy smokers who have already developed a head and neck cancer should be considered. Rarely, patients can develop a radiation therapy-induced sarcoma after having undergone prior radiotherapy for a head and neck cancer.

Much progress has been made in describing the molecular features of head and neck cancer. These features have allowed investigators to describe the genetic and epigenetic alterations and the mutational spectrum of these tumors. Early reports demonstrated frequent overexpression of the epidermal growth factor receptor (EGFR). Overexpression was shown to correlate with poor prognosis. However, it has not proved to be a good predictor of tumor response to EGFR inhibitors, which are successful in only about 10–15% of patients. *p53* mutations are also found frequently with other major affected oncogenic driver pathways including the mitotic signaling and Notch pathways and cell cycle regulation. The PI3K pathway is frequently altered, especially in HPV-positive tumors, where it is the only mutated cancer gene identified to date. Overall, these alterations affect mitogenic signaling, genetic stability, cellular proliferation, and differentiation. HPV is known to act through inhibition of the *p53* and *RB* tumor-suppressor genes, thereby initiating the carcinogenic process, and has a mutational spectrum distinct from alcohol- and tobacco-related tumors.

CLINICAL PRESENTATION AND DIFFERENTIAL DIAGNOSIS

Most tobacco-related head and neck cancers occur in patients older than age 60 years. HPV-related malignancies are frequently diagnosed in younger patients, usually in their forties or fifties, whereas EBV-related nasopharyngeal cancer can occur in all ages, including teenagers. The manifestations vary according to the stage and primary site of the tumor. Patients with nonspecific signs and symptoms in the head and neck area should be evaluated with a thorough otolaryngologic exam, particularly if symptoms persist longer than 2–4 weeks. Males are more frequently affected than women by head and neck cancers, including HPV-positive tumors.

Cancer of the nasopharynx typically does not cause early symptoms. However, it may cause unilateral serous otitis media due to obstruction

of the eustachian tube, unilateral or bilateral nasal obstruction, or epistaxis. Advanced nasopharyngeal carcinoma causes neuropathies of the cranial nerves due to skull base involvement.

Carcinomas of the oral cavity present as nonhealing ulcers, changes in the fit of dentures, or painful lesions. Tumors of the tongue base or oropharynx can cause decreased tongue mobility and alterations in speech. Cancers of the oropharynx or hypopharynx rarely cause early symptoms, but they may cause sore throat and/or otalgia. HPV-related tumors frequently present with neck lymphadenopathy as the first sign.

Hoarseness may be an early symptom of laryngeal cancer, and persistent hoarseness requires referral to a specialist for indirect laryngoscopy and/or radiographic studies. If a head and neck lesion treated initially with antibiotics does not resolve in a short period, further workup is indicated; to simply continue the antibiotic treatment may be to lose the chance of early diagnosis of a malignancy.

Advanced head and neck cancers in any location can cause severe pain, otalgia, airway obstruction, cranial neuropathies, trismus, odynophagia, dysphagia, decreased tongue mobility, fistulas, skin involvement, and massive cervical lymphadenopathy, which may be unilateral or bilateral. Some patients have enlarged lymph nodes even though no primary lesion can be detected by endoscopy or biopsy; these patients are considered to have carcinoma of unknown primary (Fig. 106-1). If the enlarged nodes are located in the upper neck and the tumor cells are of squamous cell histology, the malignancy probably arose from a mucosal surface in the head or neck. Tumor cells in supraclavicular lymph nodes may also arise from a primary site in the chest or abdomen.

The physical examination should include inspection of all visible mucosal surfaces and palpation of the floor of the mouth and of the tongue and neck. In addition to tumors themselves, leukoplakia (a white mucosal patch) or erythroplakia (a red mucosal patch) may be observed; these “pre-malignant” lesions can represent hyperplasia, dysplasia, or carcinoma in situ and require biopsy. Further examination should be performed by a specialist. Additional staging procedures include CT of the head and neck to identify the extent of the disease. Patients with lymph node involvement should have CT scan of the chest and upper abdomen to screen for distant metastases. In heavy smokers, the CT scan of the chest can also serve as a screening tool to rule out a second lung primary tumor. A positron emission tomography (PET) scan may also be administered and can help to identify or exclude distant metastases. The definitive staging procedure is an

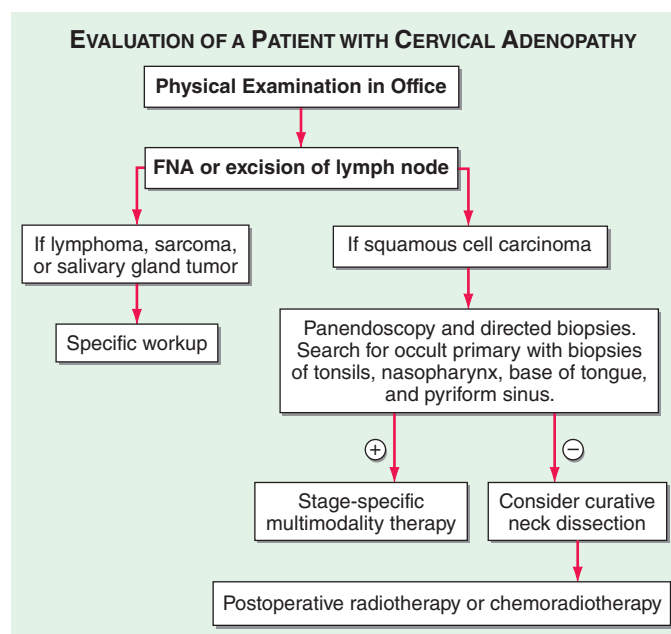


FIGURE 106-1 Evaluation of a patient with cervical adenopathy without a primary mucosal lesion; a diagnostic workup. FNA, fine-needle aspiration.

504 endoscopic examination under anesthesia, which may include laryngoscopy, esophagoscopy, and bronchoscopy; during this procedure, multiple biopsy samples are obtained to establish a primary diagnosis, define the extent of primary disease, and identify any additional premalignant lesions or second primaries.

Head and neck tumors are classified according to the tumor-node-metastasis (TNM) system of the American Joint Committee on Cancer (Fig. 106-2). This classification varies according to the specific anatomic subsite. In general, primary tumors are classified as T1 to T3 by increasing size, whereas T4 usually represents invasion of another

structure such as bone, muscle, or root of tongue. Lymph nodes are staged by size, number, and location (ipsilateral vs contralateral to the primary). Distant metastases are found in <10% of patients at initial diagnosis and are more common in patients with advanced lymph node stage; microscopic involvement of the lungs, bones, or liver is more common, particularly in patients with advanced neck lymph node disease. Modern imaging techniques may increase the number of patients with clinically detectable distant metastases in the future.

In patients with lymph node involvement and no visible primary, the diagnosis should be made by lymph node excision (Fig. 106-1). If






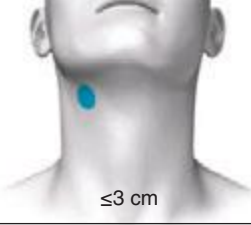
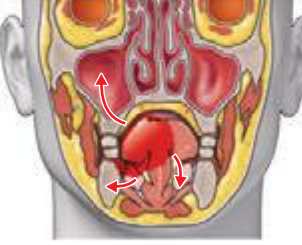
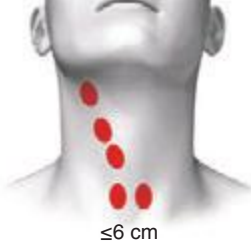
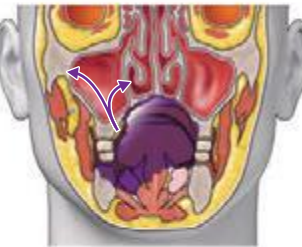
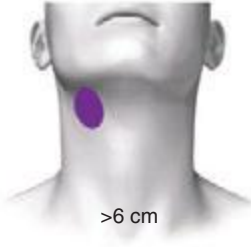
Definition of TNM			Stage groupings		
Stage I T1  Tumor ≤ 2 cm in greatest dimension without extraparenchymal extension	N0  N0- No regional lymph node metastasis	T1 N0 M0			
Stage II T2  Tumor ≥ 2 cm but not more than 4 cm in greatest dimension without extraparenchymal extension	N0  N0- No regional lymph node metastasis	T2 N0 M0			
Stage III T3  Tumor ≥ 4 cm and/or tumor having extraparenchymal extension	N1  N1- Metastasis in a single ipsilateral lymph node, ≤ 3 cm in greatest dimension	T3 N0 M0 T1 N1 M0 T2 N1 M0 T3 N1 M0			
Stage IVA T4a  Tumor invades skin, mandible, ear canal, and/or fascial nerve	N2  N2a- Metastasis in a single ipsilateral lymph node, >3 cm but ≤6 cm N2b- Metastasis in a multiple ipsilateral lymph node, none >6 cm N2c- Metastasis in a bilateral or contralateral lymph nodes, none >6 cm	T4a N0 M0 T4a N1 M0 T1 N2 M0 T2 N2 M0 T3 N2 M0 T4a N2 M0			
Stage IVB T4b  Tumor invades skull base and/or pterygoid plates and/or encases carotid artery	N3  N3- Metastasis in a lymph node >6 cm in greatest dimension	T4b Any N M0 Any T N3 M0			
Stage IVC	M1	Any T Any N M1			

FIGURE 106-2 Tumor-node-metastasis (TNM) staging system.

the results indicate squamous cell carcinoma, a panendoscopy should be performed, with biopsy of all suspicious-appearing areas and directed biopsies of common primary sites, such as the nasopharynx, tonsil, tongue base, and pyriform sinus. HPV-positive tumors especially can have small primary tumors that spread early to locoregional lymph nodes.

TREATMENT HEAD AND NECK CANCER

Patients with head and neck cancer can be grossly categorized into three clinical groups: those with localized disease, those with locally or regionally advanced disease (lymph node positive), and those with recurrent and/or metastatic disease. Comorbidities associated with tobacco and alcohol abuse can affect treatment outcome and define long-term risks for patients who are cured of their disease.

LOCALIZED DISEASE

Nearly one-third of patients have localized disease, that is, T1 or T2 (stage I or stage II) lesions without detectable lymph node involvement or distant metastases. These patients are treated with curative intent by either surgery or radiation therapy. The choice of modality differs according to anatomic location and institutional expertise. Radiation therapy is often preferred for laryngeal cancer to preserve voice function, and surgery is preferred for small lesions in the oral cavity to avoid the long-term complications of radiation, such as xerostomia and dental decay. Overall 5-year survival is 60–90%. Most recurrences occur within the first 2 years following diagnosis and are usually local.

LOCALLY OR REGIONALLY ADVANCED DISEASE

Locally or regionally advanced disease—disease with a large primary tumor and/or lymph node metastases—is the stage of presentation for >50% of patients. Such patients can also be treated with curative intent, but not with surgery or radiation therapy alone. Combined-modality therapy including surgery, radiation therapy, and chemotherapy is most successful. It can be administered as induction chemotherapy (chemotherapy before surgery and/or radiotherapy) or as concomitant (simultaneous) chemotherapy and radiation therapy. The latter is currently most commonly used and supported by the best evidence. Five-year survival rates exceed 50% in many trials, but part of this increased survival may be due to an increasing fraction of study populations with HPV-related tumors who carry a better prognosis. HPV testing of newly diagnosed tumors is now performed for most patients at the time of diagnosis, and clinical trials for HPV-related tumors are focused on exploring reductions in treatment intensity, especially radiation dose, in order to ameliorate long-term toxicities (fibrosis, swallowing dysfunction).

In patients with intermediate-stage tumors (stage III and early stage IV), concomitant chemoradiotherapy can be administered either as a primary treatment for patients with unresectable disease, to pursue an organ-preserving approach, or in the postoperative setting for intermediate-stage resectable tumors.

Induction Chemotherapy In this strategy, patients receive chemotherapy (current standard is a three-drug regimen of docetaxel, cisplatin, and fluorouracil [5-FU]) before surgery and radiation therapy. Most patients who receive three cycles show tumor reduction, and the response is clinically “complete” in up to half of patients. This “sequential” multimodality therapy allows for organ preservation (omission of surgery) in patients with laryngeal and hypopharyngeal cancer, and it has been shown to result in higher cure rates compared with radiotherapy alone.

Concomitant Chemoradiotherapy With the concomitant strategy, chemotherapy and radiation therapy are given simultaneously rather than in sequence. Tumor recurrences from head and neck cancer develop most commonly locoregionally (in the head and neck area of the primary and draining lymph nodes). The concomitant approach is aimed at enhancing tumor cell killing by radiation therapy in the presence of chemotherapy (radiation enhancement)

and is a conceptually attractive approach for bulky tumors. Toxicity (especially mucositis, grade 3 or 4 in 70–80%) is increased with concomitant chemoradiotherapy. However, meta-analyses of randomized trials document an improvement in 5-year survival of 8% with concomitant chemotherapy and radiation therapy. Results seem more favorable in recent trials as more active drugs or more intensive radiotherapy schedules are used. In addition, concomitant chemoradiotherapy produces better laryngectomy-free survival (organ preservation) than radiation therapy alone in patients with advanced larynx cancer. The use of radiation therapy together with cisplatin has also produced improved survival in patients with advanced nasopharyngeal cancer. The outcome of HPV-related cancers seems to be especially favorable following cisplatin-based chemoradiotherapy.

The success of concomitant chemoradiotherapy in patients with unresectable disease has led to the testing of a similar approach in patients with resected intermediate-stage disease as a postoperative therapy. Concomitant chemoradiotherapy produces a significant improvement over postoperative radiation therapy alone for patients whose tumors demonstrate higher risk features, such as extracapsular spread beyond involved lymph nodes, involvement of multiple lymph nodes, or positive margins at the primary site following surgery.

A monoclonal antibody to EGFR (cetuximab) increases survival rates when administered during radiotherapy. EGFR blockade results in radiation sensitization and has milder systemic side effects than traditional chemotherapy agents, although an acneiform skin rash is commonly observed. Nevertheless, the integration of cetuximab into current standard chemoradiotherapy regimens has failed to show additional improvement in survival and is not recommended.

RECURRENT AND/OR METASTATIC DISEASE

Five to ten percent of patients present with metastatic disease, and 30–50% of patients with locoregionally advanced disease experience recurrence, frequently outside the head and neck region. Patients with recurrent and/or metastatic disease are, with few exceptions, treated with palliative intent. Some patients may require local or regional radiation therapy for pain control, but most are given chemotherapy. Response rates to chemotherapy average only 30–50%; the durations of response are short, and the median survival time is 8–10 months. Therefore, chemotherapy provides transient symptomatic benefit. Drugs with single-agent activity in this setting include methotrexate, 5-FU, cisplatin, paclitaxel, and docetaxel. Combinations of cisplatin with 5-FU, carboplatin with 5-FU, and cisplatin or carboplatin with paclitaxel or docetaxel are frequently used.

EGFR-directed therapies, including monoclonal antibodies (e.g., cetuximab) and tyrosine kinase inhibitors (TKIs) of the EGFR signaling pathway (e.g., erlotinib or gefitinib), have single-agent activity of approximately 10%. Side effects are usually limited to an acneiform rash and diarrhea (for the TKIs). The addition of cetuximab to standard combination chemotherapy with cisplatin or carboplatin and 5-FU was shown to result in a significant increase in median survival. Drugs targeting specific mutations are under investigation, but no such strategy has yet been shown to be feasible in head and neck cancer.

COMPLICATIONS

Complications from treatment of head and neck cancer are usually correlated to the extent of surgery and exposure of normal tissue structures to radiation. Currently, the extent of surgery has been limited or completely replaced by chemotherapy and radiation therapy as the primary approach. Acute complications of radiation include mucositis and dysphagia. Long-term complications include xerostomia, loss of taste, decreased tongue mobility, second malignancies, dysphagia, and neck fibrosis. The complications of chemotherapy vary with the regimen used but usually include myelosuppression, mucositis, nausea and vomiting, and nephrotoxicity (with cisplatin).

506 The mucosal side effects of therapy can lead to malnutrition and dehydration. Many centers address issues of dentition before starting treatment, and some place feeding tubes to ensure control of hydration and nutrition intake. About 50% of patients develop hypothyroidism from the treatment; thus, thyroid function should be monitored.

SALIVARY GLAND TUMORS

Most benign salivary gland tumors are treated with surgical excision, and patients with invasive salivary gland tumors are treated with surgery and radiation therapy. These tumors may recur regionally; adenoid cystic carcinoma has a tendency to recur along the nerve tracks. Distant metastases may occur as late as 10–20 years after the initial diagnosis. For metastatic disease, therapy is given with palliative intent, usually chemotherapy with doxorubicin and/or cisplatin. Identification of novel agents with activity in these tumors is a high priority.

107 Neoplasms of the Lung

Leora Horn, Christine M. Lovly, David H. Johnson

Lung cancer, which was rare prior to 1900 with fewer than 400 cases described in the medical literature, is considered a disease of modern man. By the mid-twentieth century, lung cancer had become epidemic and firmly established as the leading cause of cancer-related death in North America and Europe, killing over three times as many men as prostate cancer and nearly twice as many women as breast cancer. This fact is particularly troubling because lung cancer is one of the most preventable of all of the major malignancies. Tobacco consumption is the primary cause of lung cancer, a reality firmly established in the mid-twentieth century and codified with the release of the U.S. Surgeon General's 1964 report on the health effects of tobacco smoking. Following the report, cigarette use started to decline in North America and parts of Europe, and with it, so did the incidence of lung cancer. To date, the decline in lung cancer is seen most clearly in men; only recently has the decline become apparent among women in the United States. Unfortunately, in many parts of the world, especially in countries with developing economies, cigarette use continues to increase, and along with it, the incidence of lung cancers is also rising. Although tobacco smoking remains the primary cause of lung cancer worldwide, approximately 60% of new lung cancers in the United States occur in former smokers (smoked ≥ 100 cigarettes per lifetime, quit ≥ 1 year), many of whom quit decades ago, or never smokers (smoked < 100 cigarettes per lifetime). Moreover, one in five women and one in 12 men diagnosed with lung cancer have never smoked. Given the magnitude of the problem, it is incumbent that every internist has a general knowledge of lung cancer and its management.

EPIDEMIOLOGY

Lung cancer is the most common cause of cancer death among American men and women. More than 225,000 individuals will be diagnosed with lung cancer in the United States in 2013, and over 150,000 individuals will die from the disease. The incidence of lung cancer peaked among men in the late 1980s and has plateaued in women. Lung cancer is rare below age 40, with rates increasing until age 80, after which the rate tapers off. The projected lifetime probability of developing lung cancer is estimated to be approximately 8% among males and approximately 6% among females. The incidence of lung cancer varies by racial and ethnic group, with the highest age-adjusted incidence rates among African Americans. The excess in age-adjusted rates among African Americans occurs only among men, but examinations of age-specific rates show that below age 50, mortality from lung cancer

is more than 25% higher among African American than Caucasian women. Incidence and mortality rates among Hispanics and Native and Asian Americans are approximately 40–50% those of whites.

RISK FACTORS

Cigarette smokers have a 10-fold or greater increased risk of developing lung cancer compared to those who have never smoked. A deep sequencing study suggested that one genetic mutation is induced for every 15 cigarettes smoked. The risk of lung cancer is lower among persons who quit smoking than among those who continue smoking; former smokers have a ninefold increased risk of developing lung cancer compared to men who have never smoked versus the 20-fold excess in those who continue to smoke. The size of the risk reduction increases with the length of time the person has quit smoking, although generally even long-term former smokers have higher risks of lung cancer than those who never smoked. Cigarette smoking has been shown to increase the risk of all the major lung cancer cell types. Environmental tobacco smoke (ETS) or second-hand smoke is also an established cause of lung cancer. The risk from ETS is less than from active smoking, with about a 20–30% increase in lung cancer observed among never smokers married for many years to smokers, in comparison to the 2000% increase among continuing active smokers.

Although cigarette smoking is the cause of the majority of lung cancers, several other risk factors have been identified, including occupational exposures to asbestos, arsenic, bischloromethyl ether, hexavalent chromium, mustard gas, nickel (as in certain nickel-refining processes), and polycyclic aromatic hydrocarbons. Occupational observations also have provided insight into possible mechanisms of lung cancer induction. For example, the risk of lung cancer among asbestos-exposed workers is increased primarily among those with underlying asbestosis, raising the possibility that the scarring and inflammation produced by this fibrotic nonmalignant lung disease may in many cases (although likely not in all) be the trigger for asbestos-induced lung cancer. Several other occupational exposures have been associated with increased rates of lung cancer, but the causal nature of the association is not as clear.

The risk of lung cancer appears to be higher among individuals with low fruit and vegetable intake during adulthood. This observation led to hypotheses that specific nutrients, in particular retinoids and carotenoids, might have chemopreventative effects for lung cancer. However, randomized trials failed to validate this hypothesis. In fact, studies found the incidence of lung cancer was increased among smokers with supplementation. Ionizing radiation is also an established lung carcinogen, most convincingly demonstrated from studies showing increased rates of lung cancer among survivors of the atom bombs dropped on Hiroshima and Nagasaki and large excesses among workers exposed to alpha irradiation from radon in underground uranium mining. Prolonged exposure to low-level radon in homes might impart a risk of lung cancer equal or greater than that of ETS. Prior lung diseases such as chronic bronchitis, emphysema, and tuberculosis have been linked to increased risks of lung cancer as well.

Smoking Cessation Given the undeniable link between cigarette smoking and lung cancer (not even addressing other tobacco-related illnesses), physicians must promote tobacco abstinence. Physicians also must help their patients who smoke to stop smoking. Smoking cessation, even well into middle age, can minimize an individual's subsequent risk of lung cancer. Stopping tobacco use before middle age avoids more than 90% of the lung cancer risk attributable to tobacco. However, there is little health benefit derived from just "cutting back." Importantly, smoking cessation can even be beneficial in individuals with an established diagnosis of lung cancer, as it is associated with improved survival, fewer side effects from therapy, and an overall improvement in quality of life. Moreover, smoking can alter the metabolism of many chemotherapy drugs, potentially adversely altering the toxicities and therapeutic benefits of the agents. Consequently, it is important to promote smoking cessation even *after* the diagnosis of lung cancer is established.

Physicians need to understand the essential elements of smoking cessation therapy. The individual must want to stop smoking and must be

willing to work hard to achieve the goal of smoking abstinence. Self-help strategies alone only marginally affect quit rates, whereas individual and combined pharmacotherapies in combination with counseling can significantly increase rates of cessation. Therapy with an antidepressant (e.g., bupropion) and nicotine replacement therapy (varenicline, a $\alpha_4\beta_2$ nicotinic acetylcholine receptor partial agonist) are approved by the U.S. Food and Drug Administration (FDA) as first-line treatments for nicotine dependence. However, both drugs have been reported to increase suicidal ideation and must be used with caution. In a randomized trial, varenicline was shown to be more efficacious than bupropion or placebo. Prolonged use of varenicline beyond the initial induction phase proved useful in maintaining smoking abstinence. Clonidine and nortriptyline are recommended as second-line treatments. Of note, reducing cigarettes smoked before quit day and quitting abruptly, with no prior reduction, yield comparable quit rates. Therefore, patients can be given the choice to quit in either of these ways (Chap. 470).

Inherited Predisposition to Lung Cancer Exposure to environmental carcinogens, such as those found in tobacco smoke, induce or facilitate the transformation from bronchoepithelial cells to the malignant phenotype. The contribution of carcinogens on transformation is modulated by polymorphic variations in genes that affect aspects of carcinogen metabolism. Certain genetic polymorphisms of the P450 enzyme system, specifically CYP1A1, and chromosome fragility are associated with the development of lung cancer. These genetic variations occur at relatively high frequency in the population, but their contribution to an individual's lung cancer risk is generally low. However, because of their population frequency, the overall impact on lung cancer risk could be high. In addition, environmental factors, as modified by inherited modulators, likely affect specific genes by deregulating important pathways to enable the cancer phenotype.

First-degree relatives of lung cancer probands have a two- to three-fold excess risk of lung cancer and other cancers, many of which are not smoking-related. These data suggest that specific genes and/or genetic variants may contribute to susceptibility to lung cancer. However, very few such genes have yet been identified. Individuals with inherited mutations in *RB* (patients with retinoblastoma living to adulthood) and *p53* (Li-Fraumeni syndrome) genes may develop lung cancer. Common gene variants involved in lung cancer have been recently identified through large, collaborative, genome-wide association studies. These studies identified three separate loci that are associated with lung cancer (5p15, 6p21, and 15q25) and include genes that regulate acetylcholine nicotinic receptors and telomerase production. A rare germline mutation (T790M) involving the epidermal growth factor receptor (EGFR) maybe be linked to lung cancer susceptibility in never smokers. Likewise, a susceptibility locus on chromosome 6q greatly increases risk lung cancer risk among light and never smokers. Although progress has been made, there is a significant amount of work that remains to be done in identifying heritable risk factors for lung cancer. Currently no molecular criteria are suitable to select patients for more intense screening programs or for specific chemopreventative strategies.

PATHOLOGY

The World Health Organization (WHO) defines lung cancer as tumors arising from the respiratory epithelium (bronchi, bronchioles, and alveoli). The WHO classification system divides epithelial lung cancers into four major cell types: small-cell lung cancer (SCLC), adenocarcinoma, squamous cell carcinoma, and large-cell carcinoma; the latter three types are collectively known as non-small-cell carcinomas (NSCLCs) (Fig. 107-1). Small-cell carcinomas consist of small cells with scant cytoplasm, ill-defined cell borders, finely granular nuclear chromatin, absent or inconspicuous nucleoli, and a high mitotic count. SCLC may be distinguished from NSCLC by the presence of neuroendocrine markers including CD56, neural cell adhesion molecule (NCAM), synaptophysin, and chromogranin. In North America, adenocarcinoma is the most common histologic type of lung cancer. Adenocarcinomas possess glandular differentiation or mucin production and may show acinar, papillary, lepidic, or solid features or a mixture of these patterns. Squamous cell carcinomas of the

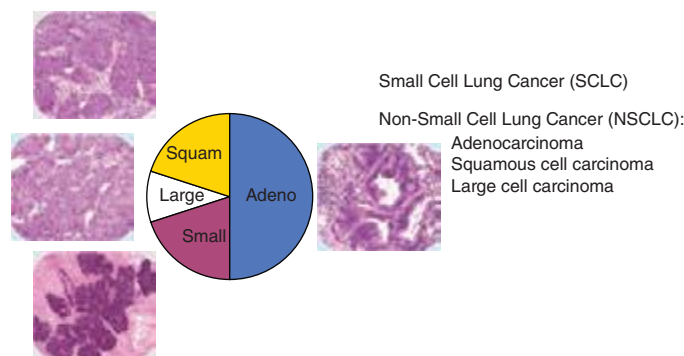


FIGURE 107-1 Traditional histologic view of lung cancer.

lung are morphologically identical to extrapulmonary squamous cell carcinomas and cannot be distinguished by immunohistochemistry alone. Squamous cell tumors show keratinization and/or intercellular bridges that arise from bronchial epithelium. The tumor tends to consist of sheets of cells rather than the three-dimensional groups of cells characteristic of adenocarcinomas. Large-cell carcinomas comprise less than 10% of lung carcinomas. These tumors lack the cytologic and architectural features of small-cell carcinoma and glandular or squamous differentiation. Together these four histologic types account for approximately 90% of all epithelial lung cancers.

All histologic types of lung cancer can develop in current and former smokers, although squamous and small-cell carcinomas are most commonly associated with heavy tobacco use. Through the first half of the twentieth century, squamous carcinoma was the most common subtype of NSCLC diagnosed in the United States. However, with the decline in cigarette consumption over the past four decades, adenocarcinoma has become the most frequent histologic subtype of lung cancer in the United States as both squamous carcinoma and small-cell carcinoma are on the decline. In lifetime never smokers or former light smokers (<10 pack-year history), women, and younger adults (<60 years), adenocarcinoma tends to be the most common form of lung cancer.

Historically, the major pathologic distinction was simply between SCLC and NSCLC, because these tumors have quite different natural histories and therapeutic approaches (see below). Likewise, until fairly recently, there was no apparent need to distinguish among the various subtypes of NSCLC because there were no clear differences in therapeutic outcome based on histology alone. However, this perspective radically changed in 2004 with the recognition that a small percentage of lung adenocarcinomas harbored mutation in *EGFR* that rendered those tumors exquisitely sensitive to inhibitors of the EGFR tyrosine kinases (e.g., gefitinib and erlotinib). This observation, coupled with the subsequent identification of other “actionable” molecular alterations (Table 107-1) and the recognition that some active

TABLE 107-1 DRIVER MUTATIONS IN NON-SMALL-CELL LUNG CANCER (NSCLC)

Gene	Alteration	Frequency in NSCLC	Typical Histology
<i>AKT1</i>	Mutation	1%	Adenocarcinoma, squamous
<i>ALK</i>	Rearrangement	3–7%	Adenocarcinoma
<i>BRAF</i>	Mutation	1–3%	Adenocarcinoma
<i>DDR2</i>	Mutation	~4%	Squamous
<i>EGFR</i>	Mutation	10–35%	Adenocarcinoma
<i>FGFR1</i>	Amplification	~20%	Squamous
<i>HER2</i>	Mutation	2–4%	Adenocarcinoma
<i>KRAS</i>	Mutation	15–25%	Adenocarcinoma
<i>MEK1</i>	Mutation	1%	Adenocarcinoma
<i>MET</i>	Amplification	2–4%	Adenocarcinoma
<i>NRAS</i>	Mutation	1%	Adenocarcinoma
<i>PIK3CA</i>	Mutation	1–3%	Squamous
<i>PTEN</i>	Mutation	4–8%	Squamous

508 chemotherapy agents performed quite differently in squamous carcinomas versus adenocarcinomas, firmly established the need for modifications in the then-existing 2004 WHO lung cancer classification system. The revised 2011 classification system, developed jointly by the International Association for the Study of Lung Cancer, the American Thoracic Society, and the European Respiratory Society, provides an integrated approach to the classification of lung adenocarcinomas that includes clinical, molecular, radiographic, and pathologic information. It also recognizes that most lung cancers present in an advanced stage and are often diagnosed based on small biopsies or cytologic specimens, rendering clear histologic distinctions difficult if not impossible.

Previously, in the 2004 classification system, tumors *failing* to show definite glandular or squamous morphology in a small biopsy or cytologic specimen were simply classified as *non-small-cell carcinoma, not otherwise specified*. However, because the distinction between adenocarcinoma and squamous carcinoma is now viewed as critical to optimal therapeutic decision making, the modified classification approach recommends these lesions be further characterized using a limited special stain workup. This distinction can be achieved using a single marker for adenocarcinoma (thyroid transcription factor-1 or napsin-A) plus a squamous marker (p40 or p63) and/or mucin stains. The modified classification system also recommends preservation of sufficient specimen material for appropriate molecular testing necessary to help guide therapeutic decision making (see below).

Another significant modification to the WHO classification system is the discontinuation of the terms *bronchioloalveolar carcinoma* and *mixed-subtype adenocarcinoma*. The term *bronchioloalveolar carcinoma* was dropped due to its inconsistent use and because it caused confusion in routine clinical care and research. As formerly used, the term encompassed at least five different entities with diverse clinical and molecular properties. The terms *adenocarcinoma in situ* and *minimally invasive adenocarcinoma* are now recommended for small solitary adenocarcinomas (≤ 3 cm) with either pure lepidic growth (term used to describe single-layered growth of atypical cuboidal cells coating the alveolar walls) or predominant lepidic growth with ≤ 5 mm invasion. Individuals with these entities experience 100% or near 100% 5-year disease-free survival with complete tumor resection. *Invasive adenocarcinomas*, representing more than 70–90% of surgically resected lung adenocarcinomas, are now classified by their predominant pattern: lepidic, acinar, papillary, and solid patterns. Lepidic-predominant subtype has a favorable prognosis, acinar and papillary have an intermediate prognosis, and solid-predominant has a poor prognosis. The terms *signet ring* and *clear cell adenocarcinoma* have been eliminated from the variants of invasive lung adenocarcinoma, whereas the term *micropapillary*, a subtype with a particularly poor prognosis, has been added. Although *EGFR* mutations are encountered most frequently in nonmucinous adenocarcinomas with a lepidic- or papillary-predominant pattern, most adenocarcinoma subtypes can harbor *EGFR* or *KRAS* mutations. The same is true of *ALK*, *RET*, and *ROS1* rearrangements. What was previously termed *mucinous bronchioloalveolar carcinoma* is now called *invasive mucinous adenocarcinoma*. These tumors generally lack *EGFR* mutations and show a strong correlation with *KRAS* mutations. Overall, the revised WHO reclassification of lung cancer addresses important advances in diagnosis and treatment, most importantly, the critical advances in understanding the specific genes and molecular pathways that initiate and sustain lung tumorigenesis resulting in new “targeted” therapies with improved specificity and better antitumor efficacy.

IMMUNOHISTOCHEMISTRY

The diagnosis of lung cancer most often rests on the morphologic or cytologic features correlated with clinical and radiographic findings. Immunohistochemistry may be used to verify neuroendocrine differentiation within a tumor, with markers such as neuron-specific enolase (NSE), CD56 or NCAM, synaptophysin, chromogranin, and Leu7. Immunohistochemistry is also helpful in differentiating primary from metastatic adenocarcinomas; thyroid transcription factor-1 (TTF-1), identified in tumors of thyroid and pulmonary origin, is positive in

over 70% of pulmonary adenocarcinomas and is a reliable indicator of primary lung cancer, provided a thyroid primary has been excluded. A negative TTF-1, however, does not exclude the possibility of a lung primary. TTF-1 is also positive in neuroendocrine tumors of pulmonary and extrapulmonary origin. Napsin-A (Nap-A) is an aspartic protease that plays an important role in maturation of surfactant B7 and is expressed in cytoplasm of type II pneumocytes. In several studies, Nap-A has been reported in $>90\%$ of primary lung adenocarcinomas. Notably, a combination of Nap-A and TTF-1 is useful in distinguishing primary lung adenocarcinoma (Nap-A positive, TTF-1 positive) from primary lung squamous cell carcinoma (Nap-A negative, TTF-1 negative) and primary SCLC (Nap-A negative, TTF-1 positive). Cytokeratins 7 and 20 used in combination can help narrow the differential diagnosis; nonsquamous NSCLC, SCLC, and mesothelioma may stain positive for CK7 and negative for CK20, whereas squamous cell lung cancer often will be both CK7 and CK20 negative. p63 is a useful marker for the detection of NSCLCs with squamous differentiation when used in cytologic pulmonary samples. Mesothelioma can be easily identified ultrastructurally, but it has historically been difficult to differentiate from adenocarcinoma through morphology and immunohistochemical staining. Several markers in the last few years have proven to be more helpful including CK5/6, calretinin, and Wilms tumor gene-1 (*WT-1*), all of which show positivity in mesothelioma.

MOLECULAR PATHOGENESIS

Cancer is a disease involving dynamic changes in the genome. As proposed by Hanahan and Weinberg, virtually all cancer cells acquire six hallmark capabilities: self-sufficiency in growth signals, insensitivity to antigrowth signals, evading apoptosis, limitless replicative potential, sustained angiogenesis, and tissue invasion and metastasis. The order in which these hallmark capabilities are acquired appears quite variable and can differ from tumor to tumor. Events leading to acquisition of these hallmarks can vary widely, although broadly, cancers arise as a result from accumulations of gain-of-function mutations in oncogenes and loss-of-function mutations in tumor-suppressor genes. Further complicating the study of lung cancer, the sequence of events that lead to disease is clearly different for the various histopathologic entities.

The exact cell of origin for lung cancers is not clearly defined. Whether one cell of origin leads to all histologic forms of lung cancer is unclear. However, for lung adenocarcinoma, evidence suggests that type II epithelial cells (or alveolar epithelial cells) have the capacity to give rise to tumors. For SCLC, cells of neuroendocrine origin have been implicated as precursors.

For cancers in general, one theory holds that a small subset of the cells within a tumor (i.e., “stem cells”) are responsible for the full malignant behavior of the tumor. As part of this concept, the large bulk of the cells in a cancer are “offspring” of these cancer stem cells. While clonally related to the cancer stem cell subpopulation, most cells by themselves cannot regenerate the full malignant phenotype. The stem cell concept may explain the failure of standard medical therapies to eradicate lung cancers, even when there is a clinical complete response. Disease recurs because therapies do not eliminate the stem cell component, which may be more resistant to chemotherapy. Precise human lung cancer stem cells have yet to be identified.

Lung cancer cells harbor multiple chromosomal abnormalities, including mutations, amplifications, insertions, deletions, and translocations. One of the earliest sets of oncogenes found to be aberrant was the *MYC* family of transcription factors (*MYC*, *MYCN*, and *MYCL*). *MYC* is most frequently activated via gene amplification or transcriptional dysregulation in both SCLC and NSCLC. Currently, there are no *MYC*-specific drugs.

Among lung cancer histologies, adenocarcinomas have been the most extensively catalogued for recurrent genomic gains and losses as well as for somatic mutations (Fig. 107-2). While multiple different kinds of aberrations have been found, a major class involves “driver mutations,” which are mutations that occur in genes encoding signaling proteins that when aberrant, drive initiation and maintenance of tumor cells. Importantly, driver mutations can serve as potential Achilles’ heels for tumors, if their gene products can be targeted

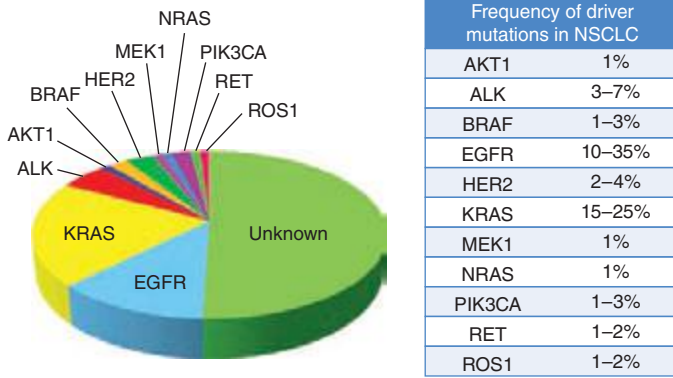


FIGURE 107-2 Driver mutations in adenocarcinomas.

appropriately. For example, one set of mutations involves EGFR, which belongs to the ERBB (HER) family of protooncogenes, including *EGFR* (ERBB1), *HER2/neu* (ERBB2), *HER3* (ERBB3), and *HER4* (ERBB4). These genes encode cell-surface receptors consisting of an extracellular ligand-binding domain, a transmembrane structure, and an intracellular tyrosine kinase (TK) domain. The binding of ligand to receptor activates receptor dimerization and TK autophosphorylation, initiating a cascade of intracellular events, and leading to increased cell proliferation, angiogenesis, metastasis, and a decrease in apoptosis. Lung adenocarcinomas can arise when tumors express mutant *EGFR*. These same tumors display high sensitivity to small-molecule EGFR TK inhibitors (TKIs). Additional examples of driver mutations in lung adenocarcinoma include the GTPase *KRAS*, the serine-threonine kinase *BRAF*, and the lipid kinase *PIK3CA*. More recently, more subsets of lung adenocarcinoma have been identified as defined by the presence of specific chromosomal rearrangements resulting in the aberrant activation of the TKs ALK, ROS1, and RET. Notably, most driver mutations in lung cancer appear to be mutually exclusive, suggesting that acquisition of one of these mutations is sufficient to drive tumorigenesis. Although driver mutations have mostly been found in adenocarcinomas, three potential molecular targets recently have been identified in squamous cell lung carcinomas: *FGFR1* amplification, *DDR2* mutations, and *PIK3CA* mutations/*PTEN* loss (Table 107-1). Together, these potentially “actionable” defects occur in up to 50% of squamous carcinomas.

A large number of tumor-suppressor genes have also been identified that are inactivated during the pathogenesis of lung cancer. These include *TP53*, *RBI*, *RASSF1A*, *CDKN2A/B*, *LKB1* (*STK11*), and *FHIT*. Nearly 90% of SCLCs harbor mutations in *TP53* and *RBI*. Several tumor-suppressor genes on chromosome 3p appear to be involved in nearly all lung cancers. Allelic loss for this region occurs very early in lung cancer pathogenesis, including in histologically normal smoking-damaged lung epithelium.

EARLY DETECTION AND SCREENING

In lung cancer, clinical outcome is related to the stage at diagnosis, and hence, it is generally assumed that early detection of occult tumors will lead to improved survival. Early detection is a process that involves screening tests, surveillance, diagnosis, and early treatment. Screening refers to the use of simple tests across a healthy population in order to identify individuals who harbor asymptomatic disease. For a screening program to be successful, there must be a high burden of disease within the target population; the test must be sensitive, specific, accessible, and cost effective; and there must be effective treatment that can reduce mortality. With any screening procedure, it is important to consider the possible influence of *lead-time bias* (detecting the cancer earlier without an effect on survival), *length-time bias* (indolent cancers are detected on screening and may not affect survival, whereas aggressive cancers are likely to cause symptoms earlier in patients and are less likely to be detected), and *overdiagnosis* (diagnosing cancers so slow growing that they are unlikely to cause the death of the patient) (Chap. 100).

Because a majority of lung cancer patients present with advanced disease beyond the scope of surgical resection, there is understandable skepticism about the value of screening in this condition. Indeed, randomized controlled trials conducted in the 1960s to 1980s using screening chest x-rays (CXR), with or without sputum cytology, reported no impact on lung cancer-specific mortality in patients characterized as high risk (males age ≥ 45 years with a smoking history). These studies have been criticized for their design, statistical analyses, and outdated imaging modalities. The results of the more recently conducted Prostate, Lung, Colorectal and Ovarian Cancer Screening Trial (PLCO) are consistent with these earlier reports. Initiated in 1993, participants in the PLCO lung cancer screening trial received annual CXR screening for 4 years, whereas participants in the usual care group received no interventions other than their customary medical care. The diagnostic follow-up of positive screening results was determined by participants and their physicians. The PLCO trial differed from previous lung cancer screening studies in that women and never smokers were eligible. The study was designed to detect a 10% reduction in lung cancer mortality in the interventional group. A total of 154,901 individuals between 55 and 74 years of age were enrolled (77,445 assigned to annual CXR screenings; 77,456 assigned to usual care). Participant demographics and tumor characteristics were well balanced between the two groups. Through 13 years of follow-up, cumulative lung cancer incidence rates (20.1 vs 19.2 per 10,000 person-years; rate ratio [RR], 1.05; 95% confidence interval [CI], 0.98–1.12) and lung cancer mortality ($n = 1213$ vs $n = 1230$) were identical between the two groups. The stage and histology of detected cancers in the two groups also were similar. These data corroborate previous recommendations *against* CXR screening for lung cancer.

In contrast to CXR, low-dose, noncontrast, thin-slice spiral chest computed tomography (LDCT) has emerged as an effective tool to screen for lung cancer. In nonrandomized studies conducted in the 1990s, LDCT scans were shown to detect more lung nodules and cancers than standard CXR in selected high-risk populations (e.g., age ≥ 60 years and a smoking history of ≥ 10 pack-years). Notably, up to 85% of the lung cancers discovered in these trials were classified as stage I disease and therefore considered potentially curable with surgical resection.

These data prompted the National Cancer Institute (NCI) to initiate the National Lung Screening Trial (NLST), a randomized study designed to determine if LDCT screening could reduce mortality from lung cancer in high-risk populations as compared with standard posterior anterior CXR. High-risk patients were defined as individuals between 55 and 74 years of age, with a ≥ 30 pack-year history of cigarette smoking; former smokers must have quit within the previous 15 years. Excluded from the trial were individuals with a previous lung cancer diagnosis, a history of hemoptysis, an unexplained weight loss of >15 lb in the preceding year, or a chest CT within 18 months of enrollment. A total of 53,454 persons were enrolled and randomized to annual screening yearly for three years (LDCT screening, $n = 26,722$; CXR screening, $n = 26,732$). Any noncalcified nodule measuring ≥ 4 mm in any diameter found on LDCT and CXR images with any noncalcified nodule or mass were classified as “positive.” Participating radiologists had the option of not calling a final screen positive if a noncalcified nodule had been stable on the three screening exams. Overall, 39.1% of participants in the LDCT group and 16% in the CXR group had at least one positive screening result. Of those who screened positive, the false-positive rate was 96.4% in the LDCT group and 94.5% in the CXR group. This was consistent across all three rounds. In the LDCT group, 1060 cancers were identified compared with 941 cancers in the CXR group (645 vs 572 per 100,000 person-years; RR, 1.13; 95% CI, 1.03 to 1.23). Nearly twice as many early-stage IA cancers were detected in the LDCT group compared with the CXR group (40% vs 21%). The overall rates of lung cancer death were 247 and 309 deaths per 100,000 participants in the LDCT and CXR groups, respectively, representing a 20% reduction in lung cancer mortality in the LDCT-screened population (95% CI, 6.8–26.7%; $p = .004$). Compared with the CXR group, the rate of death in the LDCT group from *any* cause was reduced by 6.7% (95% CI, 1.2–13.6; $p = .02$) (Table 107-2). The

	Event Number		Rates of Events per 100,000 Person-Years		Relative Risk (95% CI)	p Value
	LDCT (n = 26,772)	CXR (n = 26,732)	LDCT	CXR	RR	
Lung cancer mortality	356	443	247	309	0.80 (0.73–0.93)	.004
All-cause mortality	1877	2000	1303	1395	0.93 (0.86–0.99)	.02
Mortality not due to lung cancer	1521	1557	1056	1086	0.99 (0.95–1.02)	.51

Abbreviations: CI, confidence interval; CXR, chest x-ray; LDCT, low-dose computed tomography; RR, rate ratio.

Source: Modified from PB Bach et al: JAMA 307:2418, 2012.

number needed to screen (NNTS) to prevent one lung cancer death was calculated to be 320.

LDCT screening for lung cancer comes with known risks including a high rate of false-positive results, false-negative results, potential for unnecessary follow-up testing, radiation exposure, overdiagnosis, changes in anxiety and quality of life, and substantial financial costs. By far the biggest challenge confronting the use of CT screening is the high false-positive rate. False positives can have a substantial impact on patients through the expense and risk of unneeded further evaluation and emotional stress. The management of these patients usually consists of serial CT scans over time to see if the nodules grow, attempted fine-needle aspirates, or surgical resection. At \$300 per scan (NCI estimated cost), the outlay for initial LDCT alone could run into the billions of dollars annually, an expense that only further escalates when factoring in various downstream expenditures an individual might incur in the assessment of positive findings. A formal cost-effectiveness analysis of the NLST is expected soon that should help resolve this crucial concern.

Despite the aforementioned caveats, screening of individuals who meet the NLST criteria for lung cancer risk (or in some cases, modified versions of these criteria) seems warranted, provided comprehensive multidisciplinary coordinated care and follow-up similar to those provided to NLST participants are available. Algorithms to improve candidate selection are under development. When discussing the option of LDCT screening, use of absolute risks rather than relative risks is helpful because studies indicate the public can process absolute terminology more effectively than relative risk projections. A useful guide has been developed by the NCI to help patients and physicians assess the benefits and harms of LDCT screening for lung cancer (Table 107-3). Finally, even a small negative effect of screening on smoking behavior (either lower quit rates or higher recidivism) could easily offset the potential gains in a population. Fortunately no such impact has been reported to date. Nonetheless, smoking cessation must be included as an indispensable component of any screening program.

CLINICAL MANIFESTATIONS

Over half of all patients diagnosed with lung cancer present with locally advanced or metastatic disease at the time of diagnosis. The majority of patients present with signs, symptoms, or laboratory abnormalities that can be attributed to the primary lesion, local tumor growth, invasion or obstruction of adjacent structures, growth at distant metastatic sites, or a paraneoplastic syndrome (Tables 107-4 and 107-5). The prototypical lung cancer patient is a current or former smoker of either sex, usually in the seventh decade of life. A history of chronic cough with or without hemoptysis in a current or former smoker with chronic obstructive pulmonary disease (COPD) age 40 years or older should prompt a thorough investigation for lung cancer even in the face of a normal CXR. A persistent pneumonia without constitutional symptoms and unresponsive to repeated courses of antibiotics also should prompt an evaluation for the underlying cause. Lung cancer arising in a lifetime never smoker is more common in women and East Asians. Such patients also tend to be younger than their smoking counterparts at the time of diagnosis. The clinical presentation of lung cancer in never smokers tends to mirror that of current and former smokers.

Patients with central or endobronchial growth of the primary tumor may present with cough, hemoptysis, wheeze, stridor, dyspnea, or postobstructive pneumonitis. Peripheral growth of the primary tumor may cause pain from pleural or chest wall involvement, dyspnea on a restrictive basis, and symptoms of a lung abscess resulting from tumor cavitation. Regional spread of tumor in the thorax (by contiguous growth or by metastasis to regional lymph nodes) may cause tracheal obstruction, esophageal compression with dysphagia, recurrent laryngeal paralysis with hoarseness, phrenic nerve palsy with elevation of the hemidiaphragm and dyspnea, and sympathetic nerve paralysis with Horner's syndrome (enophthalmos, ptosis, miosis, and anhidrosis). Malignant pleural effusions can cause pain, dyspnea, or cough. Pancoast (or superior sulcus tumor) syndromes result from local extension of a tumor growing in the apex of the lung with involvement of the eighth cervical and first and second thoracic nerves, and present with shoulder pain that characteristically radiates in the ulnar

TABLE 107-3 THE BENEFITS AND HARMS OF LDCT SCREENING FOR LUNG CANCER BASED ON NLST DATA

	LDCT	CXR
Benefits: How Did CT Scans Help Compared To CXR?		
4 in 1000 fewer died from lung cancer	13 in 1000	17 in 1000
5 in 1000 fewer died from all causes	70 in 1000	75 in 1000
Harms: What Problems Did CT Scans Cause Compared to CXR?		
223 in 1000 had at least 1 false alarm	365 in 1000	142 in 1000
18 in 1000 had a false alarm leading to an invasive procedure	25 in 1000	7 in 1000
2 in 1000 had a major complication from an invasive procedure	3 in 1000	1 in 1000

Abbreviations: CXR, chest x-ray; LDCT, low-dose computed tomography; NLST, National Lung Screening Trial.

Source: Modified from S Woloshin et al: N Engl J Med 367:1677, 2012.

TABLE 107-4 PRESENTING SIGNS AND SYMPTOMS OF LUNG CANCER

Symptom and Signs	Range of Frequency
Cough	8–75%
Weight loss	0–68%
Dyspnea	3–60%
Chest pain	20–49%
Hemoptysis	6–35%
Bone pain	6–25%
Clubbing	0–20%
Fever	0–20%
Weakness	0–10%
Superior vena cava obstruction	0–4%
Dysphagia	0–2%
Wheezing and stridor	0–2%

Source: Reproduced with permission from MA Beckles: Chest 123:97-104, 2003.

TABLE 107-5 CLINICAL FINDINGS SUGGESTIVE OF METASTATIC DISEASE

Symptoms elicited in history	<ul style="list-style-type: none"> • Constitutional: weight loss >10 lb • Musculoskeletal: focal skeletal pain • Neurologic: headaches, syncope, seizures, extremity weakness, recent change in mental status
Signs found on physical examination	<ul style="list-style-type: none"> • Lymphadenopathy (>1 cm) • Hoarseness, superior vena cava syndrome • Bone tenderness • Hepatomegaly (>13 cm span) • Focal neurologic signs, papilledema • Soft-tissue mass
Routine laboratory tests	<ul style="list-style-type: none"> • Hematocrit, <40% in men; <35% in women • Elevated alkaline phosphatase, GGT, SGOT, and calcium levels

Abbreviations: GGT, gamma-glutamyltransferase; SGOT, serum glutamic-oxaloacetic transaminase.

Source: Reproduced with permission from GA Silvestri et al: *Chest* 123(1 Suppl):147S, 2003.

distribution of the arm, often with radiologic destruction of the first and second ribs. Often Horner's syndrome and Pancoast syndrome coexist. Other problems of regional spread include superior vena cava syndrome from vascular obstruction; pericardial and cardiac extension with resultant tamponade, arrhythmia, or cardiac failure; lymphatic obstruction with resultant pleural effusion; and lymphangitic spread through the lungs with hypoxemia and dyspnea. In addition, lung cancer can spread transbronchially, producing tumor growth along multiple alveolar surfaces with impairment of gas exchange, respiratory insufficiency, dyspnea, hypoxemia, and sputum production. Constitutional symptoms may include anorexia, weight loss, weakness, fever, and night sweats. Apart from the brevity of symptom duration, these parameters fail to clearly distinguish SCLC from NSCLC or even from neoplasms metastatic to lungs.

Extrathoracic metastatic disease is found at autopsy in more than 50% of patients with squamous carcinoma, 80% of patients with adenocarcinoma and large-cell carcinoma, and greater than 95% of patients with SCLC. Approximately one-third of patients present with symptoms as a result of distant metastases. Lung cancer metastases may occur in virtually every organ system, and the site of metastatic involvement largely determines other symptoms. Patients with brain metastases may present with headache, nausea and vomiting, seizures, or neurologic deficits. Patients with bone metastases may present with pain, pathologic fractures, or cord compression. The latter may also occur with epidural metastases. Individuals with bone marrow invasion may present with cytopenias or leukoerythroblastosis. Those with liver metastases may present with hepatomegaly, right upper quadrant pain, fever, anorexia, and weight loss. Liver dysfunction and biliary obstructions are rare. Adrenal metastases are common but rarely cause pain or adrenal insufficiency unless they are large.

Paraneoplastic syndromes are common in patients with lung cancer, especially those with SCLC, and may be the presenting finding or the first sign of recurrence. In addition, paraneoplastic syndromes may mimic metastatic disease and, unless detected, lead to inappropriate palliative rather than curative treatment. Often the paraneoplastic syndrome may be relieved with successful treatment of the tumor. In some cases, the pathophysiology of the paraneoplastic syndrome is known, particularly when a hormone with biological activity is secreted by a tumor. However, in many cases, the pathophysiology is unknown. Systemic symptoms of anorexia, cachexia, weight loss (seen in 30% of patients), fever, and suppressed immunity are paraneoplastic syndromes of unknown etiology or at least not well defined. Weight loss greater than 10% of total body weight is considered a bad prognostic sign. Endocrine syndromes are seen in 12% of patients; hypercalcemia resulting from ectopic production of parathyroid hormone (PTH), or more commonly, PTH-related peptide, is the most common life-threatening metabolic complication of malignancy, primarily occurring with squamous cell carcinomas of the lung. Clinical symptoms

include nausea, vomiting, abdominal pain, constipation, polyuria, thirst, and altered mental status.

Hyponatremia may be caused by the syndrome of inappropriate secretion of antidiuretic hormone (SIADH) or possibly atrial natriuretic peptide (ANP). SIADH resolves within 1-4 weeks of initiating chemotherapy in the vast majority of cases. During this period, serum sodium can usually be managed and maintained above 128 mEq/L via fluid restriction. Demeclocycline can be a useful adjunctive measure when fluid restriction alone is insufficient. Vasopressin receptor antagonists like tolvaptan also have been used in the management of SIADH. However, there are significant limitations to the use of tolvaptan including liver injury and overly rapid correction of the hyponatremia, which can lead to irreversible neurologic injury. Likewise, the cost of tolvaptan may be prohibitive (as high as \$300 per tablet in some areas). Of note, patients with ectopic ANP may have worsening hyponatremia if sodium intake is not concomitantly increased. Accordingly, if hyponatremia fails to improve or worsens after 3-4 days of adequate fluid restriction, plasma levels of ANP should be measured to determine the causative syndrome.

Ectopic secretion of ACTH by SCLC and pulmonary carcinoids usually results in additional electrolyte disturbances, especially hypokalemia, rather than the changes in body habitus that occur in Cushing's syndrome from a pituitary adenoma. Treatment with standard medications, such as metyrapone and ketoconazole, is largely ineffective due to extremely high cortisol levels. The most effective strategy for management of the Cushing's syndrome is effective treatment of the underlying SCLC. Bilateral adrenalectomy may be considered in extreme cases.

Skeletal-connective tissue syndromes include clubbing in 30% of cases (usually NSCLCs) and hypertrophic primary osteoarthropathy in 1-10% of cases (usually adenocarcinomas). Patients may develop periostitis, causing pain, tenderness, and swelling over the affected bones and a positive bone scan. Neurologic-myopathic syndromes are seen in only 1% of patients but are dramatic and include the myasthenic Eaton-Lambert syndrome and retinal blindness with SCLC, whereas peripheral neuropathies, subacute cerebellar degeneration, cortical degeneration, and polymyositis are seen with all lung cancer types. Many of these are caused by autoimmune responses such as the development of anti-voltage-gated calcium channel antibodies in Eaton-Lambert syndrome. Patients with this disorder present with proximal muscle weakness, usually in the lower extremities, occasional autonomic dysfunction, and rarely, cranial nerve symptoms or involvement of the bulbar or respiratory muscles. Depressed deep tendon reflexes are frequently present. In contrast to patients with myasthenia gravis, strength improves with serial effort. Some patients who respond to chemotherapy will have resolution of the neurologic abnormalities. Thus, chemotherapy is the initial treatment of choice. Paraneoplastic encephalomyelitis and sensory neuropathies, cerebellar degeneration, limbic encephalitis, and brainstem encephalitis occur in SCLC in association with a variety of antineuronal antibodies such as anti-Hu, anti-CRMP5, and ANNA-3. Paraneoplastic cerebellar degeneration may be associated with anti-Hu, anti-Yo, or P/Q calcium channel autoantibodies. Coagulation or thrombotic or other hematologic manifestations occur in 1-8% of patients and include migratory venous thrombophlebitis (Trousseau's syndrome), nonbacterial thrombotic (marantic) endocarditis with arterial emboli, and disseminated intravascular coagulation with hemorrhage, anemia, granulocytosis, and leukoerythroblastosis. Thrombotic disease complicating cancer is usually a poor prognostic sign. Cutaneous manifestations such as dermatomyositis and acanthosis nigricans are uncommon (1%), as are the renal manifestations of nephrotic syndrome and glomerulonephritis ($\leq 1\%$).

DIAGNOSING LUNG CANCER

Tissue sampling is required to confirm a diagnosis in all patients with suspected lung cancer. In patients with suspected metastatic disease, a biopsy of the most distant site of disease is preferred for tissue confirmation. Given the greater emphasis placed on molecular testing for NSCLC patients, a core biopsy is preferred to ensure adequate

512 tissue for analysis. Tumor tissue may be obtained via minimally invasive techniques such as bronchial or transbronchial biopsy during fiberoptic bronchoscopy, by fine-needle aspiration or percutaneous biopsy using image guidance, or via endobronchial ultrasound (EBUS) guided biopsy. Depending on the location, lymph node sampling may occur via transesophageal endoscopic ultrasound-guided biopsy (EUS), EBUS, or blind biopsy. In patients with clinically palpable disease such as a lymph node or skin metastasis, a biopsy may be obtained. In patients with suspected metastatic disease, a diagnosis may be confirmed by percutaneous biopsy of a soft tissue mass, lytic bone lesion, bone marrow, pleural or liver lesion, or an adequate cell block obtained from a malignant pleural effusion. In patients with a suspected malignant pleural effusion, if the initial thoracentesis is negative, a repeat thoracentesis is warranted. Although the majority of pleural effusions are due to malignant disease, particularly if they are exudative or bloody, some may be parapneumonic. In the absence of distant disease, such patients should be considered for possible curative treatment.

The diagnostic yield of any biopsy depends on several factors including location (accessibility) of the tumor, tumor size, tumor type, and technical aspects of the diagnostic procedure including the experience level of the bronchoscopist and pathologist. In general, central lesions such as squamous cell carcinomas, small-cell carcinomas, or endobronchial lesions such as carcinoid tumors are more readily diagnosed by bronchoscopic examination, whereas peripheral lesions such as adenocarcinomas and large-cell carcinomas are more amenable to transthoracic biopsy. Diagnostic accuracy for SCLC versus NSCLC for most specimens is excellent, with lesser accuracy for subtypes of NSCLC.

Bronchoscopic specimens include bronchial brush, bronchial wash, bronchioloalveolar lavage, transbronchial fine-needle aspiration (FNA), and core biopsy. For more accurate histologic classification, mutation analysis, or investigational purposes, reasonable efforts (e.g., a core needle biopsy) should be made to obtain more tissue than what is contained in a routine cytology specimen obtained by FNA. Overall sensitivity for combined use of bronchoscopic methods is approximately 80%, and together with tissue biopsy, the yield increases to

85–90%. Like transbronchial core biopsy specimens, transthoracic core biopsy specimens are also preferred. Sensitivity is highest for larger lesions and peripheral tumors. In general, core biopsy specimens, whether transbronchial, transthoracic, or EUS-guided, are superior to other specimen types. This is primarily due to the higher percentage of tumor cells with fewer confounding factors such as obscuring inflammation and reactive nonneoplastic cells.

Sputum cytology is inexpensive and noninvasive but has a lower yield than other specimen types due to poor preservation of the cells and more variability in acquiring a good-quality specimen. The yield for sputum cytology is highest for larger and centrally located tumors such as squamous cell carcinoma and small-cell carcinoma histology. The specificity for sputum cytology averages close to 100%, although sensitivity is generally <70%. The accuracy of sputum cytology improves with increased numbers of specimens analyzed. Consequently, analysis of at least three sputum specimens is recommended.

STAGING LUNG CANCER

Lung cancer staging consists of two parts: first, a determination of the location of the tumor and possible metastatic sites (anatomic staging), and second, an assessment of a patient's ability to withstand various antitumor treatments (physiologic staging). All patients with lung cancer should have a complete history and physical examination, with evaluation of all other medical problems, determination of performance status, and history of weight loss. The most significant dividing line is between those patients who are candidates for surgical resection and those who are inoperable but will benefit from chemotherapy, radiation therapy, or both. Staging with regard to a patient's potential for surgical resection is principally applicable to NSCLC.

ANATOMIC STAGING OF PATIENTS WITH LUNG CANCER

The accurate staging of patients with NSCLC is essential for determining the appropriate treatment in patients with resectable disease and avoiding unnecessary surgical procedures in patients with advanced disease (Fig. 107-3). All patients with NSCLC should undergo initial

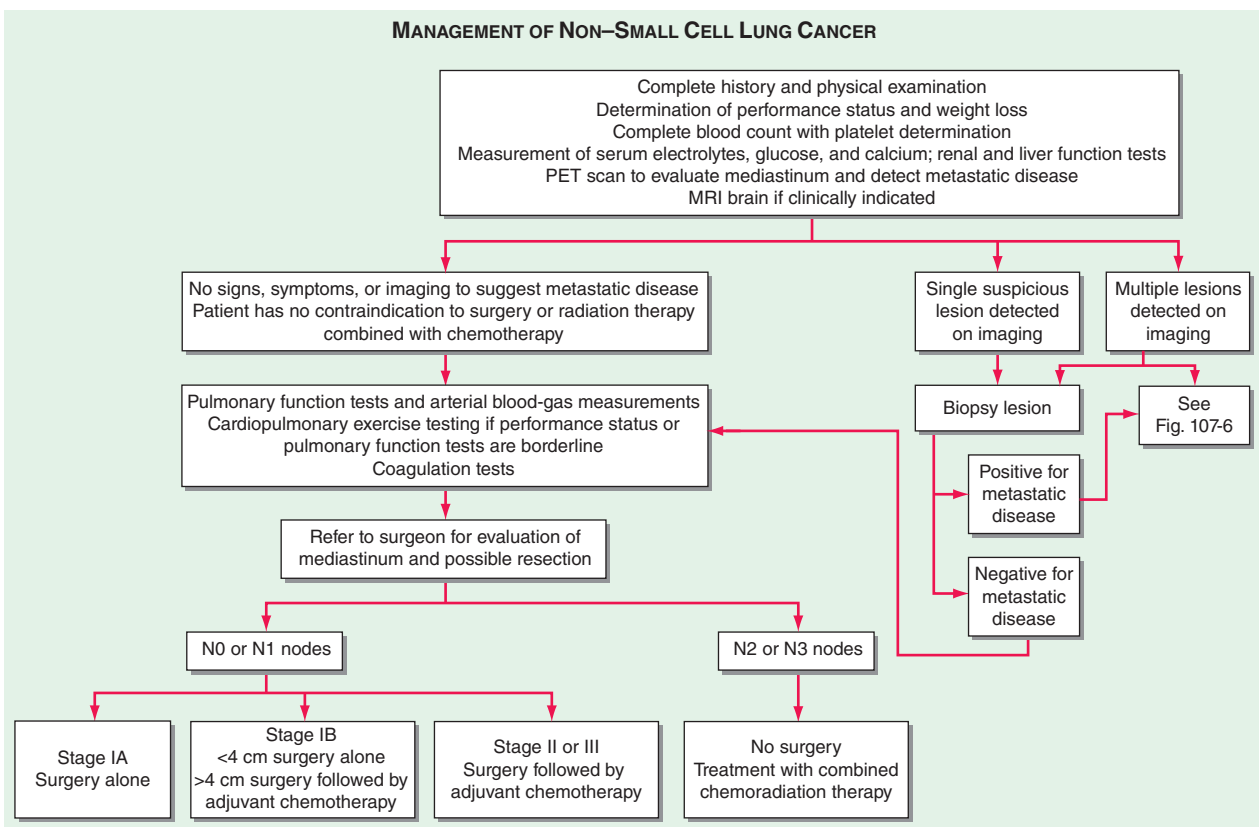


FIGURE 107-3 Algorithm for management of non-small-cell lung cancer. MRI, magnetic resonance imaging; PET, positron emission tomography.

radiographic imaging with CT scan, positron emission tomography (PET), or preferably CT-PET. PET scanning attempts to identify sites of malignancy based on glucose metabolism by measuring the uptake of ^{18}F -fluorodeoxyglucose (FDG). Rapidly dividing cells, presumably in the lung tumors, will preferentially take up ^{18}F -FDG and appear as a “hot spot.” To date, PET has been mostly used for staging and detection of metastases in lung cancer and in the detection of nodules >15 mm in diameter. Combined ^{18}F -FDG PET-CT imaging has been shown to improve the accuracy of staging in NSCLC compared to visual correlation of PET and CT or either study alone. CT-PET has been found to be superior in identifying pathologically enlarged mediastinal lymph nodes and extrathoracic metastases. A standardized uptake value (SUV) of >2.5 on PET is highly suspicious for malignancy. False negatives can be seen in diabetes, in lesions <8 mm, and in slow-growing tumors (e.g., carcinoid tumors or well-differentiated adenocarcinoma). False positives can be seen in certain infections and granulomatous disease (e.g., tuberculosis). Thus, PET should never be used alone to diagnose lung cancer, mediastinal involvement, or metastases. Confirmation with tissue biopsy is required. For brain metastases, magnetic resonance imaging (MRI) is the most effective method. MRI can also be useful in selected circumstances, such as superior sulcus tumors to rule out brachial plexus involvement, but in general, MRI does not play a major role in NSCLC staging.

In patients with NSCLC, the following are contraindications to potential curative resection: extrathoracic metastases, superior vena cava syndrome, vocal cord and, in most cases, phrenic nerve paralysis, malignant pleural effusion, cardiac tamponade, tumor within 2 cm of the carina (potentially curable with combined chemoradiotherapy), metastasis to the contralateral lung, metastases to supraclavicular lymph nodes, contralateral mediastinal node metastases (potentially curable with combined chemoradiotherapy), and involvement of the main pulmonary artery. In situations where it will make a difference in treatment, abnormal scan findings require tissue confirmation of malignancy so that patients are not precluded from having potentially curative therapy.

The best predictor of metastatic disease remains a careful history and physical examination. If signs, symptoms, or findings from the physical examination suggest the presence of malignancy, then sequential imaging starting with the most appropriate study should be performed. If the findings from the clinical evaluation are negative, then imaging studies beyond CT-PET are unnecessary and the search for metastatic disease is complete. More controversial is how one should assess patients with known stage III disease. Because these patients are more likely to have asymptomatic occult metastatic disease, current guidelines recommend a more extensive imaging evaluation including imaging of the brain with either CT scan or MRI. In patients in whom distant metastatic disease has been ruled out, lymph node status needs to be assessed via a combination of radiographic imaging and/or minimally invasive techniques such as those mentioned above and/or invasive techniques such as mediastinoscopy, mediastinotomy, thoracoscopy, or thoracotomy. Approximately one-quarter to one-half of patients diagnosed with NSCLC will have mediastinal lymph node metastases at the time of diagnosis. Lymph node sampling is recommended in all patients with enlarged nodes detected by CT or PET scan and in patients with large tumors or tumors occupying the inner third of the lung. The extent of mediastinal lymph node involvement is important in determining the appropriate treatment strategy: surgical resection followed by adjuvant chemotherapy versus combined chemoradiation alone (see below). A standard nomenclature for referring to the location of lymph nodes involved with lung cancer has evolved (Fig. 107-4).

In SCLC patients, current staging recommendations include a CT scan of the chest and abdomen (because of the high frequency of hepatic and adrenal involvement), MRI of the brain (positive in 10% of asymptomatic patients), and radionuclide bone scan if symptoms or signs suggest disease involvement in these areas (Fig. 107-5). Although there are less data on the use of CT-PET in SCLC, the most recent American College of Chest Physicians Evidence-Based Clinical Practice Guidelines recommend PET scans in patients with clinical

stage I SCLC who are being considered for curative intent surgical resection. In addition, invasive mediastinal staging and extrathoracic imaging (head MRI/CT and PET or abdominal CT plus bone scan) is also recommended for patients with clinical stage I SCLC if curative intent surgical resection is contemplated. Some practice guidelines also recommend the use of PET scanning in the staging of SCLC patients who are potential candidates for the addition of thoracic radiotherapy to chemotherapy. Bone marrow biopsies and aspirations are rarely performed now given the low incidence of isolated bone marrow metastases. Confirmation of metastatic disease, ipsilateral or contralateral lung nodules, or metastases beyond the mediastinum may be achieved by the same modalities recommended earlier for patients with NSCLC.

If a patient has signs or symptoms of spinal cord compression (pain, weakness, paralysis, urinary retention), a spinal CT or MRI scan and examination of the cerebrospinal fluid cytology should be performed. If metastases are evident on imaging, a neurosurgeon should be consulted for possible palliative surgical resection and/or a radiation oncologist should be consulted for palliative radiotherapy to the site of compression. If signs or symptoms of leptomeningitis develop at any time in a patient with lung cancer, an MRI of the brain and spinal cord should be performed, as well as a spinal tap, for detection of malignant cells. If the spinal tap is negative, a repeat spinal tap should be considered. There is currently no approved therapy for the treatment of leptomeningeal disease.

STAGING SYSTEM FOR NON-SMALL-CELL LUNG CANCER

The tumor-node-metastasis (TNM) international staging system provides useful prognostic information and is used to stage all patients with NSCLC. The various T (tumor size), N (regional node involvement), and M (presence or absence of distant metastasis) are combined to form different stage groups (Tables 107-6 and 107-7). The previous edition of the TNM staging system for lung cancer was developed based on a relatively small database of patients from a single institution. The latest seventh edition of the TNM staging system went into effect in 2010 and developed using a much more robust database of more than 100,000 patients with lung cancer who were treated in multiple countries between 1990 and 2000. Data from 67,725 patients with NSCLC were then used to reevaluate the prognostic value of the TNM descriptors (Table 107-8). The major distinction between the sixth and seventh editions of the international staging systems is within the T classification; T1 tumors are divided into tumors ≤ 2 cm in size, as these patients were found to have a better prognosis compared to patients with tumors >2 cm but ≤ 3 cm. T2 tumors are divided into those that are >3 cm but ≤ 5 cm and those that are >5 cm but ≤ 7 cm. Tumors that are >7 cm are considered T3 tumors. T3 tumors also include tumors with invasion into local structures such as chest wall and diaphragm and additional nodules in the same lobe. T4 tumors include tumors of any size with invasion into mediastinum, heart, great vessels, trachea, or esophagus or multiple nodules in the ipsilateral lung. No changes have been made to the current classification of lymph node involvement (N). Patients with metastasis may be classified as M1a (malignant pleural or pericardial effusion, pleural nodules, or nodules in the contralateral lung) or M1b (distant metastasis; e.g., bone, liver, adrenal, or brain metastasis). Based on these data, approximately one-third of patients have localized disease that can be treated with curative attempt (surgery or radiotherapy), one-third have local or regional disease that may or may not be amenable to a curative attempt, and one-third have metastatic disease at the time of diagnosis.

STAGING SYSTEM FOR SMALL-CELL LUNG CANCER

In patients with SCLC, it is now recommended that both the Veterans Administration system and the American Joint Committee on Cancer/International Union Against Cancer seventh edition system (TNM) be used to classify the tumor stage. The Veterans Administration system is a distinct two-stage system dividing patients into those with limited- or extensive-stage disease. Patients with limited-stage disease (LD) have cancer that is confined to the ipsilateral hemithorax and can be encompassed within a tolerable radiation port. Thus, contralateral

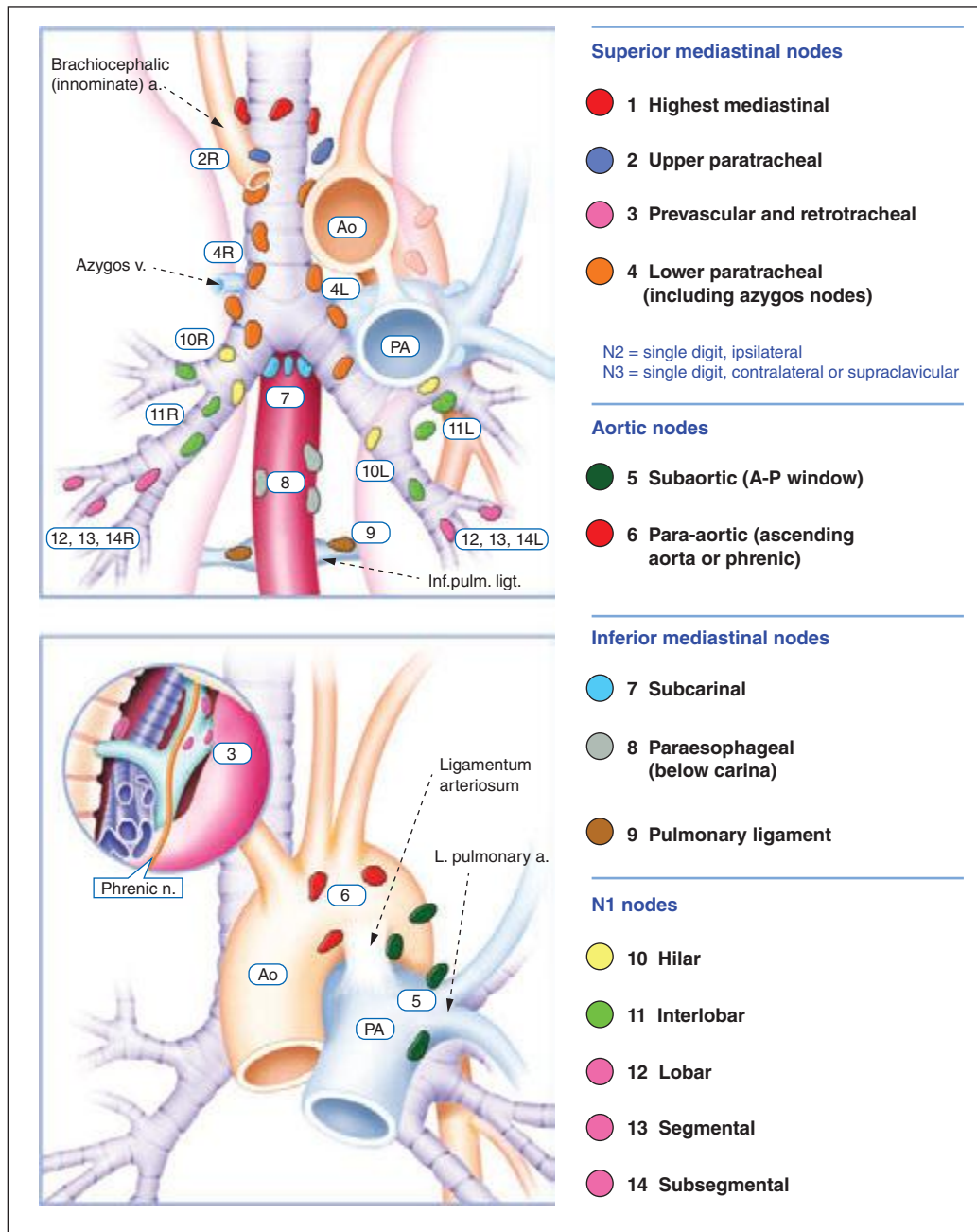


FIGURE 107-4 Lymph node stations in staging non-small-cell lung cancer. The International Association for the Study of Lung Cancer (IASLC) lymph node map, including the proposed grouping of lymph node stations into “zones” for the purposes of prognostic analyses. a., artery; Ao, aorta; Inf. pulm. lig., inferior pulmonary ligament; n., nerve; PA, pulmonary artery; v., vein.

supraclavicular nodes, recurrent laryngeal nerve involvement, and superior vena caval obstruction can all be part of LD. Patients with extensive-stage disease (ED) have overt metastatic disease by imaging or physical examination. Cardiac tamponade, malignant pleural effusion, and bilateral pulmonary parenchymal involvement generally qualify disease as ED, because the involved organs cannot be encompassed safely or effectively within a single radiation therapy port. Sixty to 70% of patients are diagnosed with ED at presentation. The TNM staging system is preferred in the rare SCLC patient presenting with what appears to be clinical stage I disease (see above).

PHYSIOLOGIC STAGING

Patients with lung cancer often have other comorbid conditions related to smoking including cardiovascular disease and COPD. To improve their preoperative condition, correctable problems (e.g., anemia, electrolyte and fluid disorders, infections, cardiac disease, and arrhythmias) should be addressed, appropriate chest physical

therapy should be instituted, and patients should be encouraged to stop smoking. Because it is not always possible to predict whether a lobectomy or pneumonectomy will be required until the time of operation, a conservative approach is to restrict surgical resection to patients who could potentially tolerate a pneumonectomy. Patients with a forced expiratory volume in 1 s (FEV_1) of greater than 2 L or greater than 80% of predicted can tolerate a pneumonectomy, and those with an FEV_1 greater than 1.5 L have adequate reserve for a lobectomy. In patients with borderline lung function but a resectable tumor, cardiopulmonary exercise testing could be performed as part of the physiologic evaluation. This test allows an estimate of the maximal oxygen consumption (VO_{2max}). A $VO_{2max} < 15$ mL/(kg·min) predicts for a higher risk of postoperative complications. Patients deemed unable to tolerate lobectomy or pneumonectomy from a pulmonary functional standpoint may be candidates for more limited resections, such as wedge or anatomic segmental resection, although such procedures are associated with significantly higher rates of local

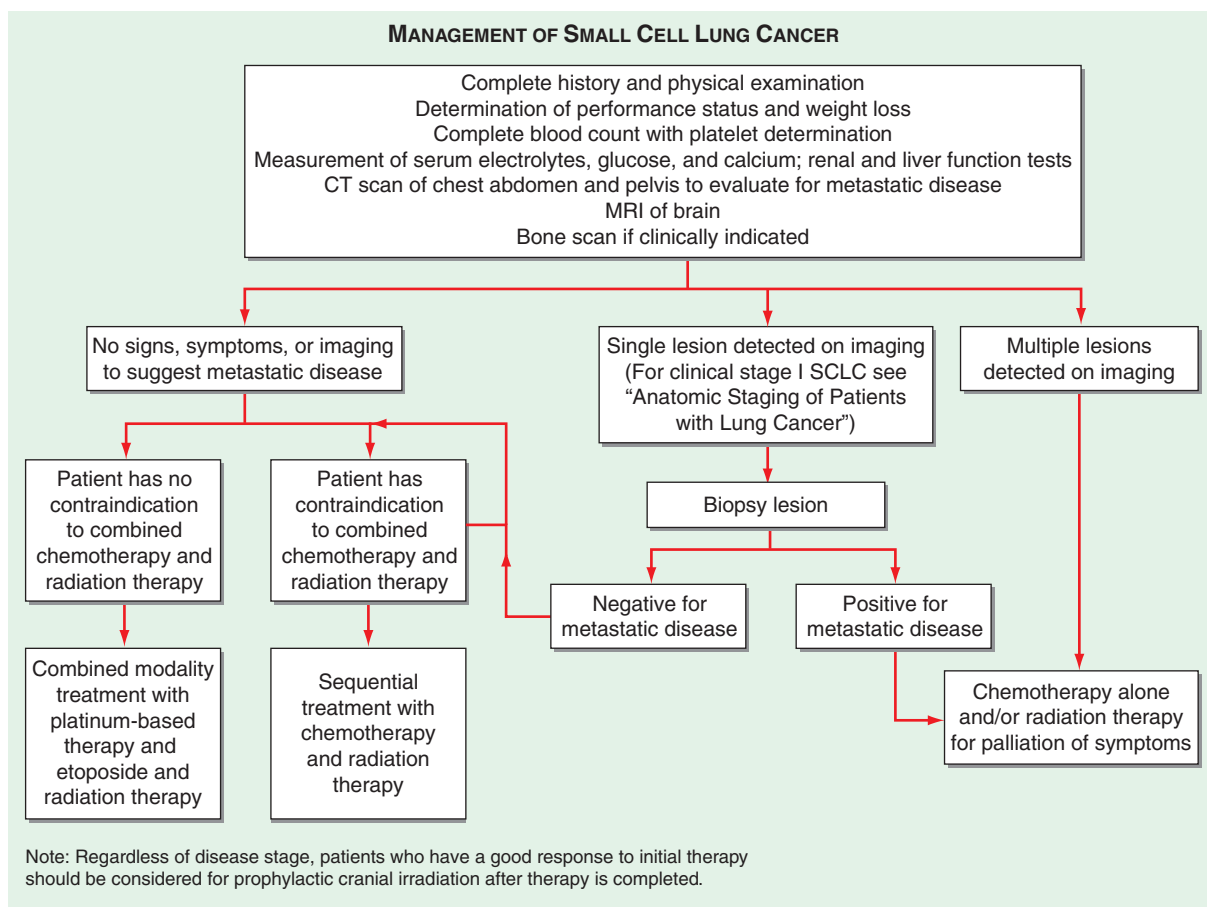


FIGURE 107-5 Algorithm for management of small-cell lung cancer. CT, computed tomography; MRI, magnetic resonance imaging.

recurrence and a trend toward decreased overall survival. All patients should be assessed for cardiovascular risk using American College of Cardiology and American Heart Association guidelines. A myocardial infarction within the past 3 months is a contraindication to thoracic surgery because 20% of patients will die of reinfarction. An infarction in the past 6 months is a relative contraindication. Other major contraindications include uncontrolled arrhythmias, an FEV₁ of less than 1 L, CO₂ retention (resting Pco₂ >45 mmHg), DL_{co} <40%, and severe pulmonary hypertension.

TREATMENT NON-SMALL-CELL LUNG CANCER

The overall treatment approach to patients with NSCLC is shown in Fig. 107-3.

OCCULT AND STAGE 0 CARCINOMAS

Patients with severe atypia on sputum cytology have an increased risk of developing lung cancer compared to those without atypia. In the uncommon circumstance where malignant cells are identified in a sputum or bronchial washing specimen but the chest imaging appears normal (TX tumor stage), the lesion must be localized. More than 90% of tumors can be localized by meticulous examination of the bronchial tree with a fiberoptic bronchoscope under general anesthesia and collection of a series of differential brushings and biopsies. Surgical resection following bronchoscopic localization has been shown to improve survival compared to no treatment. Close follow-up of these patients is indicated because of the high incidence of second primary lung cancers (5% per patient per year).

SOLITARY PULMONARY NODULE AND "GROUND-GLASS" OPACITIES

A solitary pulmonary nodule is defined as an x-ray density completely surrounded by normal aerated lung with circumscribed

margins, of any shape, usually 1–6 cm in greatest diameter. The approach to a patient with a solitary pulmonary nodule is based on an estimate of the probability of cancer, determined according to the patient's smoking history, age, and characteristics on imaging (Table 107-9). Prior CXRs and CT scans should be obtained if available for comparison. A PET scan may be useful if the lesion is greater than 7–8 mm in diameter. If no diagnosis is apparent, Mayo investigators reported that clinical characteristics (age, cigarette smoking status, and prior cancer diagnosis) and three radiologic characteristics (nodule diameter, spiculation, and upper lobe location) were independent predictors of malignancy. At present, only two radiographic criteria are thought to predict the benign nature of a solitary pulmonary nodule: lack of growth over a period >2 years and certain characteristic patterns of calcification. Calcification alone, however, does not exclude malignancy; a dense central nidus, multiple punctuate foci, and "bull's eye" (granuloma) and "popcorn ball" (hamartoma) calcifications are highly suggestive of a benign lesion. In contrast, a relatively large lesion, lack of or asymmetric calcification, chest symptoms, associated atelectasis, pneumonitis, or growth of the lesion revealed by comparison with an old x-ray or CT scan or a positive PET scan may be suggestive of a malignant process and warrant further attempts to establish a histologic diagnosis. An algorithm for assessing these lesions is shown in Fig. 107-6.

Since the advent of screening CTs, small "ground-glass" opacities (GGOs) have often been observed, particularly as the increased sensitivity of CTs enables detection of smaller lesions. Many of these GGOs, when biopsied, are found to be atypical adenomatous hyperplasia (AAH), adenocarcinoma in situ (AIS), or minimally invasive adenocarcinoma (MIA). AAH is usually a nodule of <5 mm and is minimally hazy, also called nonsolid or ground glass (i.e., hazy slightly increased attenuation, no solid component, and preservation of bronchial and vascular margins). On thin-section CT, AIS is

TABLE 107-6 SEVENTH EDITION TNM STAGING SYSTEMS FOR NON-SMALL-CELL LUNG CANCER

TNM Staging System for Lung Cancer (7th Edition)	
Primary Tumor (T)	
T1	Tumor ≤3 cm diameter, surrounded by lung or visceral pleura, without invasion more proximal than lobar bronchus
T1a	Tumor ≤2 cm in diameter
T1b	Tumor >2 cm but ≤3 cm in diameter
T2	Tumor >3 cm but ≤7 cm, or tumor with any of the following features: Involves main bronchus, ≥2 cm distal to carina Invades visceral pleura Associated with atelectasis or obstructive pneumonitis that extends to the hilar region but does not involve the entire lung
T2a	Tumor >3 cm but ≤5 cm
T2b	Tumor >5 cm but ≤7 cm
T3	Tumor >7 cm or any of the following: Directly invades any of the following: chest wall, diaphragm, phrenic nerve, mediastinal pleura, parietal pericardium, main bronchus <2 cm from carina (without involvement of carina) Atelectasis or obstructive pneumonitis of the entire lung Separate tumor nodules in the same lobe
T4	Tumor of any size that invades the mediastinum, heart, great vessels, trachea, recurrent laryngeal nerve, esophagus, vertebral body, carina, or with separate tumor nodules in a different ipsilateral lobe
Regional Lymph Nodes (N)	
N0	No regional lymph node metastases
N1	Metastasis in ipsilateral peribronchial and/or ipsilateral hilar lymph nodes and intrapulmonary nodes, including involvement by direct extension
N2	Metastasis in ipsilateral mediastinal and/or subcarinal lymph node(s)
N3	Metastasis in contralateral mediastinal, contralateral hilar, ipsilateral or contralateral scalene, or supraclavicular lymph node(s)
Distant Metastasis (M)	
M0	No distant metastasis
M1	Distant metastasis
M1a	Separate tumor nodule(s) in a contralateral lobe; tumor with pleural nodules or malignant pleural or pericardial effusion
M1b	Distant metastasis (in extrathoracic organs)

Abbreviation: TNM, tumor-node-metastasis.

Source: Reproduced with permission from P Goldstraw et al: J Thorac Oncol 2:706, 2007.

TABLE 107-7 SEVENTH EDITION TNM STAGING SYSTEMS FOR NON-SMALL-CELL LUNG CANCER

Stage groupings			
Stage IA	T1a-T1b	N0	M0
Stage IB	T2a	N0	M0
Stage IIA	T1a,T1b,T2a	N1	M0
	T2b	N0	M0
Stage IIB	T2b	N1	M0
	T3	N0	M0
Stage IIIA	T1a,T1b,T2a,T2b	N2	M0
	T3	N1,N2	M0
	T4	N0,N1	M0
Stage IIIB	T4	N2	M0
	Any T	N3	M0
Stage IV	Any T	Any N	M1a or M1b

Abbreviation: TNM, tumor-node-metastasis.

Source: Reproduced with permission from P Goldstraw P et al: J Thorac Oncol 2:706, 2007.

TABLE 107-8 FIVE-YEAR SURVIVAL BY STAGE AND TNM CLASSIFICATION OF NON-SMALL-CELL LUNG CANCER (SEVENTH EDITION)

Stage	TNM Seventh Edition	5-Year Survival (%)
IA	T1a-T1bN0M0	73%
IB	T2aN0M0	58%
IIA	T1a-T2aN1M0	46%
	T2bN0M0	
IIB	T2bN1M0	36%
	T3N0M0	
IIIA	T1a-T3N2M0	24%
	T3N1M0	
	T4N0-1M0	
IIIB	T4N2M0	9%
	T1a-T4N3M0	
IV	Any T Any N plus M1a or M1b	13%

Abbreviation: TNM, tumor-node-metastasis.

usually a nonsolid nodule and tends to be slightly more opaque than AAH. MIA is mainly solid, usually with a small (<5 mm) central solid component. However, overlap exists among the imaging features of the preinvasive and minimally invasive lesions in the lung adenocarcinoma spectrum. Lepidic adenocarcinomas are usually solid but may be nonsolid. Likewise, the small invasive adenocarcinomas also are usually solid but may exhibit a small nonsolid component.

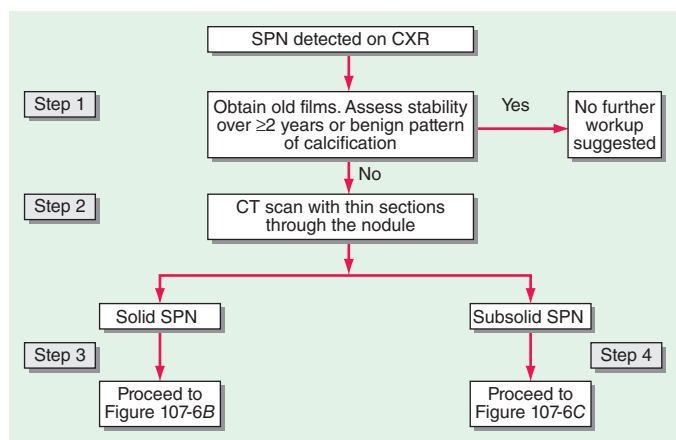
MANAGEMENT OF STAGES I AND II NSCLC

Surgical Resection of Stage I and II NSCLC Surgical resection, ideally by an experienced thoracic surgeon, is the treatment of choice for patients with clinical stage I and II NSCLC who are able to tolerate the procedure. Operative mortality rates for patients resected by thoracic or cardiothoracic surgeons are lower compared to general surgeons. Moreover, survival rates are higher in patients who undergo resection in facilities with a high surgical volume compared to those performing fewer than 70 procedures per year, even though the higher-volume facilities often serve older and less socio-economic advantaged populations. The improvement in survival is most evident in the immediate postoperative period. The extent of resection is a matter of surgical judgment based on findings at exploration. In patients with stage IA NSCLC, lobectomy is superior to wedge resection with respect to rates of local recurrence. There is also a trend toward improvement in overall survival. In patients with comorbidities, compromised pulmonary reserve, and small peripheral lesions, a limited resection, wedge resection, and segmentectomy (potentially by video-assisted thoracoscopic surgery) may be reasonable surgical option. Pneumonectomy is reserved for patients with central tumors and should be performed only in

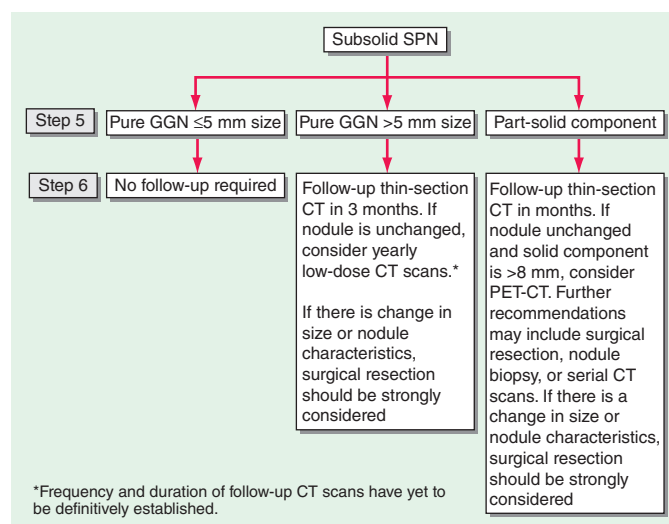
TABLE 107-9 ASSESSMENT OF RISK OF CANCER IN PATIENTS WITH SOLITARY PULMONARY NODULES

Variable	Risk		
	Low	Intermediate	High
Diameter (cm)	<1.5	1.5–2.2	≥2.3
Age (years)	<45	45–60	>60
Smoking status	Never smoker	Current smoker (<20 cigarettes/d)	Current smoker (>20 cigarettes/d)
Smoking cessation status	Quit ≥7 years ago or quit	Quit <7 years ago	Never quit
Characteristics of nodule margins	Smooth	Scalloped	Corona radiata or spiculated

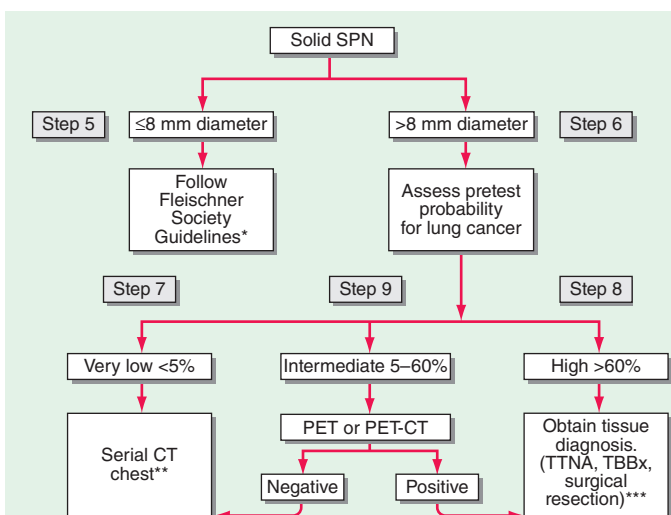
Source: Reproduced with permission from D Ost et al: N Engl J Med 348:2535, 2003.



A



C

*Fleischner society guidelines; modified from: H. MacMahon, et al: *Radiology* 2005; 237:395–400

Nodule size (a):	Low-risk patient (b):	High-risk patient (c):
≤ 4 mm	No follow-up needed (d)	Follow-up at 12 months; if unchanged, no further follow-up
> 4 –6 mm	Follow-up CT at 12 months; if unchanged, no further follow-up	Follow-up CT at 6–12 months; then 18–24 months if no change
> 6 –8 mm	Follow-up CT at 6–12 months; then 18–24 months if no change	Follow-up CT at 3–6 months; then 9–12 and 24 months if no change
> 8 mm	Follow-up CT at 3, 9, and 24 months; dynamic contrast-enhanced CT, PET, and/or biopsy	Same as low-risk patient

(a) Average of largest and smallest axial diameters of the nodule

(b) No smoking history and absence of other risk factors

(c) Previous or current smoking history or other risk factors

(d) Risk of malignancy ($< 0.1\%$) is substantially lower than for an asymptomatic smoker**ACCP guidelines (see MK Gould et al: *Chest* 2007;132(suppl 3):108s-130S.

***Consider patient preference, severity of medical comorbidities, center specific expertise prior to tissue diagnosis.

B

FIGURE 107-6 A. Algorithm for evaluation of solitary pulmonary nodule (SPN). B. Algorithm for evaluation of solid SPN. C. Algorithm for evaluation of semisolid SPN. CT, computed tomography; CXR, chest radiograph; GGN, ground-glass nodule; PET, positron emission tomography; TTBx, transbronchial biopsy; TTNA, transthoracic needle biopsy. (Adapted from VK Patel et al: *Chest* 143:840, 2013.)

patients with excellent pulmonary reserve. The 5-year survival rates are 60–80% for patients with stage I NSCLC and 40–50% for patients with stage II NSCLC.

Accurate pathologic staging requires adequate segmental, hilar, and mediastinal lymph node sampling. Ideally this includes a mediastinal lymph node dissection. On the right side, mediastinal stations 2R, 4R, 7, 8R, and 9R should be dissected; on the left side, stations 5, 6, 7, 8L, and 9L should be dissected. Hilar lymph nodes are typically resected and sent for pathologic review, although it is helpful to specifically dissect and label level 10 lymph nodes when possible. On the left side, level 2 and sometimes level 4 lymph nodes are generally obscured by the aorta. Although the therapeutic benefit of nodal dissection versus nodal sampling is controversial, a pooled analysis of three trials involving patients with stages I to IIIA NSCLC demonstrated a superior 4-year survival in patients undergoing resection and a complete mediastinal lymph node dissection compared with lymph node sampling. Moreover, complete mediastinal lymphadenectomy added little morbidity to a pulmonary resection for lung cancer when carried out by an experienced thoracic surgeon.

Radiation Therapy in Stages I and II NSCLC There is currently no role for postoperative radiation therapy in patients following resection of stage I or II NSCLC. However, patients with stage I and II disease

who either refuse or are not suitable candidates for surgery should be considered for radiation therapy with curative intent. Stereotactic body radiation therapy (SBRT) is a relatively new technique used to treat patients with isolated pulmonary nodules (≤ 5 cm) who are not candidates for or refuse surgical resection. Treatment is typically administered in three to five fractions delivered over 1–2 weeks. In uncontrolled studies, disease control rates are $> 90\%$, and 5-year survival rates of up to 60% have been reported with SBRT. By comparison, survival rates typically range from 13 to 39% in patients with stage I or II NSCLC treated with standard external-beam radiotherapy. Cryoablation is another technique occasionally used to treat small, isolated tumors (i.e., ≤ 3 cm). However, very little data exist on long-term outcomes with this technique.

Chemotherapy in Stages I and II NSCLC Although a landmark meta-analysis of cisplatin-based adjuvant chemotherapy trials in patients with resected stages I to IIIA NSCLC (the Lung Adjuvant Cisplatin Evaluation [LACE] Study) demonstrated a 5.4% improvement in 5-year survival for adjuvant chemotherapy compared to surgery alone, the survival benefit was seemingly confined to patients with stage II or III disease (Table 107-10). By contrast, survival was actually worsened in stage IA patients with the application of adjuvant therapy. In stage IB, there was a modest improvement in survival of questionable clinical significance. Adjuvant chemotherapy was

TABLE 107-10 ADJUVANT CHEMOTHERAPY TRIALS IN NON-SMALL-CELL LUNG CANCER

Trial	Stage	Treatment	No. of Patients	5-Year Survival (%)	p
IALT	I–III	Cisplatin-based	932	44.5	< .03
		Control	835	40.4	
BR10	IB–II	Cisplatin + vinorelbine	242	69	.03
		Control	240	54	
ANITA	IB–IIIA	Cisplatin + vinorelbine	407	60	.017
		Control	433	58	
ALPI	I–III	MVP	548	50	.49
		Control	540	45	
BLT	I–III	Cisplatin-based	192	60	.90
		Control	189	58	
CALGB	IB	Carboplatin + paclitaxel	173	59	.10
			171	57	

Abbreviations: ALPI, Adjuvant Lung Cancer Project Italy; ANITA, Adjuvant Navelbine International Trialist Association; BLT, Big Lung Trial; CALGB, Cancer and Lung Cancer Group B; IALT, International Adjuvant Lung Cancer Trial; MVP, mitomycin, vindesine, and cisplatin.

also detrimental in patients with poor performance status (Eastern Cooperative Oncology Group [ECOG] performance status = 2). These data suggest that adjuvant chemotherapy is best applied in patients with resected stage II or III NSCLC. There is no apparent role for adjuvant chemotherapy in patients with resected stage IA or IB NSCLC. A possible exception to the prohibition of adjuvant therapy in this setting is the stage IB patient with a resected lesion ≥ 4 cm.

As with any treatment recommendation, the risks and benefits of adjuvant chemotherapy should be considered on an individual patient basis. If a decision is made to proceed with adjuvant chemotherapy, in general, treatment should be initiated 6–12 weeks after surgery, assuming the patient has fully recovered, and should be administered for no more than four cycles. Although a cisplatin-based chemotherapy is the preferred treatment regimen, carboplatin can be substituted for cisplatin in patients who are unlikely to tolerate cisplatin for reasons such as reduced renal function, presence of neuropathy, or hearing impairment. No specific chemotherapy regimen is considered optimal in this setting, although platinum plus vinorelbine is most commonly used.

Neoadjuvant chemotherapy, which is the application of chemotherapy administered *before* an attempted surgical resection, has been advocated by some experts on the assumption that such an approach will more effectively extinguish occult micrometastases compared to postoperative chemotherapy. In addition, it is thought that preoperative chemotherapy might render an inoperable lesion resectable. With the exception of superior sulcus tumors, however, the role of neoadjuvant chemotherapy in stage I to III disease is not well defined. However, a meta-analysis of 15 randomized controlled trials involving more than 2300 patients with stage I to III NSCLC suggested there may be a modest 5-year survival benefit (i.e., ~5%) that is virtually identical to the survival benefit achieved with postoperative chemotherapy. Accordingly, neoadjuvant therapy may prove useful in selected cases (see below). A decision to use neoadjuvant chemotherapy should always be made in consultation with an experienced surgeon.

It should be noted that all patients with resected NSCLC are at high risk of recurrence, most of which occurs within 18–24 months of surgery, or developing a second primary lung cancer. Thus, it is reasonable to follow these patients with periodic imaging studies. Given the results of the NLST, periodic CT scans appear to be the most appropriate screening modality. Based on the timing of most recurrences, some guidelines recommend a contrasted chest CT scan every 6 months for the first 3 years after surgery, followed by yearly CT scans of the chest without contrast thereafter.

MANAGEMENT OF STAGE III NSCLC

Management of patients with stage III NSCLC usually requires a combined-modality approach. Patients with stage IIIA disease

commonly are stratified into those with “nonbulky” or “bulky” mediastinal lymph node (N2) disease. Although the definition of “bulky” N2 disease varies somewhat in the literature, the usual criteria include the size of a dominant lymph node (i.e., >2 – 3 cm in short-axis diameter as measured by CT), groupings of multiple smaller lymph nodes, evidence of extracapsular nodal involvement, or involvement of more than two lymph node stations. The distinction between nonbulky and bulky stage IIIA disease is mainly used to select potential candidates for *upfront* surgical resection or for resection after neoadjuvant therapy. Many aspects of therapy of patients with stage III NSCLC remain controversial, and the optimal treatment strategy has not been clearly defined. Moreover, although there are many potential treatment options, none yields a very high probability of cure. Furthermore, because stage III disease is highly heterogeneous, no single treatment approach can be recommended for all patients. Key factors guiding treatment choices include the particular combination of tumor (T) and nodal (N) disease, the ability to achieve a complete surgical resection if indicated, and the patient’s overall physical condition and preferences. For example, in carefully selected patients with limited stage IIIA disease where involved mediastinal lymph nodes can be completely resected, initial surgery followed by postoperative chemotherapy (with or without radiation therapy) may be indicated. By contrast, for patients with clinically evident bulky mediastinal lymph node involvement, the standard approach to treatment is concurrent chemoradiotherapy. Nevertheless, in some cases, the latter group of patients may be candidates for surgery following chemoradiotherapy.

Absent and Nonbulky Mediastinal (N2, N3) Lymph Node Disease For the subset of stage IIIA patients initially thought to have clinical stage I or II disease (i.e., pathologic involvement of mediastinal [N2] lymph nodes is *not* detected preoperatively), surgical resection is often the treatment of choice. This is followed by adjuvant chemotherapy in patients with microscopic lymph node involvement in a resection specimen. Postoperative radiation therapy (PORT) may also have a role for those with close or positive surgical margins. Patients with tumors involving the chest wall or proximal airways within 2 cm of the carina with hilar lymph node involvement (but not N2 disease) are classified as having T3N1 stage IIIA disease. They too are best managed with surgical resection, if technically feasible, followed by adjuvant chemotherapy if completely resected. Patients with tumors exceeding 7 cm in size also are now classified as T3 and are considered stage IIIA if tumor has spread to N1 nodes. The appropriate initial management of these patients involves surgical resection when feasible, provided the mediastinal staging is negative, followed by adjuvant chemotherapy for those who achieve complete tumor resection. Patients with T3N0 or T3N1 disease due to the presence of satellite nodules within the same lobe as the primary tumor also are candidates for surgery, as are patients with ipsilateral

nodules in another lobe and negative mediastinal nodes (IIIA, T4N0 or T4N1). Although data regarding adjuvant chemotherapy in the latter subsets of patients are limited, it is often recommended.

Patients with T4N0-1 were reclassified as having stage IIIA tumors in the seventh edition of the TNM system. These patients may have involvement of the carina, superior vena cava, or a vertebral body and yet still be candidates for surgical resection in selected circumstances. The decision to proceed with an attempted resection must be made in consultation with an experienced thoracic surgeon often in association with a vascular or cardiac surgeon and an orthopedic surgeon depending on tumor location. However, if an incomplete resection is inevitable or if there is evidence of N2 involvement (stage IIIB), surgery for T4 disease is contraindicated. Most T4 lesions are best treated with chemoradiotherapy.

The role of PORT in patients with completely resected stage III NSCLC is controversial. To a large extent, the use of PORT is dictated by the presence or absence of N2 involvement and, to a lesser degree, by the biases of the treating physician. Using the Surveillance, Epidemiology, and End Results (SEER) database, a recent meta-analysis of PORT identified a significant increase in survival in patients with N2 disease but not in patients with N0 or N1 disease. An earlier analysis by the PORT Meta-analysis Trialist Group using an older database produced similar results.

Known Mediastinal (N2, N3) Lymph Node Disease When pathologic involvement of mediastinal lymph nodes is documented preoperatively, a combined-modality approach is recommended assuming the patient is a candidate for treatment with curative intent. These patients are at high risk for both local and distant recurrence if managed with resection alone. For patients with stage III disease who are not candidates for initial surgical resection, *concurrent* chemoradiotherapy is most commonly used as the initial treatment. Concurrent chemoradiotherapy has been shown to produce superior survival compared to *sequential* chemoradiotherapy; however, it also is associated with greater host toxicities (including fatigue, esophagitis, and neutropenia). Therefore, for patients with a good performance status, concurrent chemoradiotherapy is the preferred treatment approach, whereas sequential chemoradiotherapy may be more appropriate for patients with a performance status that is not as good. For patients who are *not* candidates for a combined-modality treatment approach, typically due to a poor performance status or a comorbidity that makes chemotherapy untenable, radiotherapy alone may provide a modest survival benefit in addition to symptom palliation.

For patients with potentially resectable N2 disease, it remains uncertain whether surgery after neoadjuvant chemoradiotherapy improves survival. In an NCI-sponsored Intergroup randomized trial comparing concurrent chemoradiotherapy alone to concurrent chemoradiotherapy followed by attempted surgical resection, no survival benefit was observed in the trimodality arm compared to the bimodality therapy. In fact, patients subjected to a pneumonectomy had a worse survival outcome. By contrast, those treated with a lobectomy appeared to have a survival advantage based on a retrospective subset analysis. Thus, in carefully selected, otherwise healthy patients with nonbulky mediastinal lymph node involvement, surgery may be a reasonable option if the primary tumor can be fully resected with a lobectomy. This is not the case if a pneumonectomy is required to achieve complete resection.

Superior Sulcus Tumors (Pancoast Tumors) Superior sulcus tumors represent a distinctive subset of stage III disease. These tumors arise in the apex of the lung and may invade the second and third ribs, the brachial plexus, the subclavian vessels, the stellate ganglion, and adjacent vertebral bodies. They also may be associated with Pancoast syndrome, characterized by pain that may arise in the shoulder or chest wall or radiate to the neck. Pain characteristically radiates to the ulnar surface of the hand. Horner's syndrome (enophthalmos, ptosis, miosis, and anhidrosis) due to invasion of the paravertebral sympathetic chain may be present as well. Patients with these tumors should undergo the same staging procedures as all

patients with stage II and III NSCLC. Neoadjuvant chemotherapy or combined chemoradiotherapy followed by surgery is reserved for those without N2 involvement. This approach yields excellent survival outcomes (>50% 5-year survival in patients with an R0 resection). Patients with N2 disease are less likely to benefit from surgery and can be managed with chemoradiotherapy alone. Patients presenting with metastatic disease can be treated with radiation therapy (with or without chemotherapy) for symptom palliation.

MANAGEMENT OF METASTATIC NSCLC

Approximately 40% of NSCLC patients present with advanced, stage IV disease at the time of diagnosis. These patients have a poor median survival (4–6 months) and a 1-year survival of 10% when managed with best supportive care alone. In addition, a significant number of patients who first presented with early-stage NSCLC will eventually relapse with distant disease. Patients who have recurrent disease have a better prognosis than those presenting with metastatic disease at the time of diagnosis. Standard medical management, the judicious use of pain medications, and the appropriate use of radiotherapy and chemotherapy form the cornerstone of management. Chemotherapy palliates symptoms, improves the quality of life, and improves survival in patients with stage IV NSCLC, particularly in patients with good performance status. In addition, economic analysis has found chemotherapy to be cost-effective palliation for stage IV NSCLC. However, the use of chemotherapy for NSCLC requires clinical experience and careful judgment to balance potential benefits and toxicities. Of note, the early application of palliative care in conjunction with chemotherapy is associated with improved survival and a better quality of life.

First-Line Chemotherapy for Metastatic or Recurrent NSCLC A landmark meta-analysis published in 1995 provided the earliest meaningful indication that chemotherapy could provide a survival benefit in metastatic NSCLC as opposed to supportive care alone. However, the survival benefit was seemingly confined to cisplatin-based chemotherapy regimens (hazard ratio 0.73; 27% reduction in the risk of death; 10% improvement in survival at 1 year). These data launched two decades of clinical research aimed at detecting the optimal chemotherapy regimen for advanced NSCLC. For the most part, however, these efforts proved unsuccessful because the overwhelming majority of randomized trials showed no major survival improvement with any one regimen versus another (Table 107-11). On the other hand, differences in progression-free survival, cost, side effects, and schedule were frequently observed. These first-line studies were later extended to elderly patients, where doublet chemotherapy was found to improve overall survival compared to single agents in the “fit” elderly (e.g., elderly patients with no major comorbidities) and in patients with an ECOG performance status of 2. An ongoing debate in the treatment of patients with advanced NSCLC is the appropriate duration of platinum-based chemotherapy. Several large phase III randomized trials have failed to show a meaningful benefit for increasing the duration of platinum-based doublet chemotherapy beyond four to six cycles. In fact, longer duration of chemotherapy has been associated with increased toxicities and impaired quality of life. Therefore, prolonged front-line therapy (beyond four to six cycles) with platinum-based regimens is not recommended. Maintenance therapy following initial platinum-based therapy is discussed below.

Although specific tumor histology was once considered irrelevant to treatment choice in NSCLC, with the recent recognition that selected chemotherapy agents perform quite differently in squamous versus adenocarcinomas, accurate determination of histology has become essential. Specifically, in a landmark randomized phase III trial, patients with nonsquamous NSCLC were found to have an improved survival when treated with cisplatin and pemetrexed compared to cisplatin and gemcitabine. By contrast, patients with squamous carcinoma had an improved survival when treated with cisplatin and gemcitabine. This survival difference is thought to be related to the differential expression of thymidylate synthase (TS),

TABLE 107-11 FIRST-LINE CHEMOTHERAPY TRIALS FOR METASTATIC NON-SMALL-CELL LUNG CANCER

Trial	Regimen	No. of Patients	RR (%)	Median Survival (months)
ECOG1594	Cisplatin + paclitaxel	288	21	7.8
	Cisplatin + gemcitabine	288	22	8.1
	Cisplatin + docetaxel	289	17	7.4
	Carboplatin + paclitaxel	290	17	8.1
TAX-326	Cisplatin + docetaxel	406	32	11.3
	Cisplatin + vinorelbine	394	25	10.1
	Carboplatin + docetaxel	404	24	9.4
EORTC	Cisplatin + paclitaxel	159	32	8.1
	Cisplatin + gemcitabine	160	37	8.9
	Paclitaxel + gemcitabine	161	28	6.7
ILCP	Cisplatin + gemcitabine	205	30	9.8
	Carboplatin + paclitaxel	204	32	9.9
	Cisplatin + vinorelbine	203	30	9.5
SWOG	Cisplatin + vinorelbine	202	28	8.0
	Carboplatin + paclitaxel	206	25	8.0
FACS	Cisplatin + irinotecan	145	31	13.9
	Carboplatin + paclitaxel	145	32	12.3
	Cisplatin + gemcitabine	146	30	14.0
	Cisplatin + vinorelbine	145	33	11.4
Scagliotti	Cisplatin + gemcitabine	863	28	10.3
	Cisplatin + pemetrexed	862	31	10.3
iPASS ^a	Carboplatin + paclitaxel	608	32	17.3
	Gefitinib	609	43%	18.6

^aEnrolled selected patients: 18 years of age or older, had histologic or cytologically confirmed stage IIIb or IV non-small-cell lung cancer with histologic features of adenocarcinoma (including bronchioloalveolar carcinoma), were nonsmokers (defined as patients who had smoked <100 cigarettes in their lifetime) or former light smokers (those who had stopped smoking at least 15 years previously and had a total of ≤10 pack-years of smoking), and had had no previous chemotherapy or biologic or immunologic therapy.

Abbreviations: ECOG, Eastern Cooperative Oncology Group; EORTC, European Organization for Research and Treatment of Cancer; ILCP, Italian Lung Cancer Project; SWOG, Southwest Oncology Group; FACS, Follow-up After Colorectal Surgery; iPASS, Iressa Pan-Asian Study.

one of the targets of pemetrexed, between tumor types. Squamous cancers have a much higher expression of TS compared to adenocarcinomas, accounting for their lower responsiveness to pemetrexed. By contrast, the activity of gemcitabine is not impacted by the levels of TS. Bevacizumab, a monoclonal antibody against VEGF, has been shown to improve response rate, progression-free survival, and overall survival in patients with advanced disease when combined with chemotherapy (see below). However, bevacizumab cannot be given to patients with squamous cell histology NSCLC because of their tendency to experience serious hemorrhagic effects.

Agents That Inhibit Angiogenesis Bevacizumab, a monoclonal antibody directed against VEGF, was the first antiangiogenic agent approved for the treatment of patients with advanced NSCLC in the United States. This drug primarily acts by blocking the growth of new blood vessels, which are required for tumor viability. Two randomized phase III trials of chemotherapy with or without bevacizumab had conflicting results. The first trial, conducted in North America, compared carboplatin plus paclitaxel with or without bevacizumab in patients with recurrent or advanced nonsquamous NSCLC and reported a significant improvement in response rate, progression-free survival, and overall survival in patients treated with chemotherapy plus bevacizumab versus chemotherapy alone. Bevacizumab-treated patients had a significantly higher incidence of toxicities. The second trial, conducted in Europe, compared cisplatin/gemcitabine with or without bevacizumab in patients with recurrent or advanced nonsquamous NSCLC and reported a significant improvement in progression-free survival but no improvement

in overall survival for bevacizumab-treated patients. A randomized phase III trial compared carboplatin/pemetrexed and bevacizumab to carboplatin/paclitaxel and bevacizumab as first-line therapy in patients with recurrent or advanced nonsquamous NSCLC and reported no significant difference in progression-free survival or overall survival between treatment groups. Therefore, currently carboplatin/paclitaxel and bevacizumab or carboplatin/pemetrexed and bevacizumab are appropriate regimens for first-line treatment for stage IV nonsquamous NSCLC patients. Of note, there are many small-molecule inhibitors of VEGFR; however, these VEGFR TKIs have not proven to be effective in the treatment of NSCLC.

Maintenance Therapy for Metastatic NSCLC Maintenance chemotherapy in nonprogressing patients (patients with a complete response, partial response, or stable disease) is a controversial topic in the treatment of NSCLC. Conceptually, there are two types of maintenance strategies: (1) switch maintenance therapy, where patients receive four to six cycles of platinum-based chemotherapy and are switched to an entirely different regimen; and (2) continuation maintenance therapy, where patients receive four to six cycles of platinum-based chemotherapy and then the platinum agent is discontinued but the agent it is paired with is continued (Table 107-12). Two studies investigated switch maintenance single-agent chemotherapy with docetaxel or pemetrexed in nonprogressing patients following treatment with first-line platinum-based chemotherapy. Both trials randomized patients to immediate single-agent therapy versus observation and reported improvements in progression-free and overall survival. In both trials, a significant portion of patients in the observation arm did not receive therapy with the agent under investigation upon disease progression; 37% of study patients never received docetaxel in the docetaxel study and 81% of patients never received pemetrexed in the pemetrexed study. In the trial of maintenance docetaxel versus observation, survival was identical to the treatment group in the subset of patients who received docetaxel on progression, indicating this is an active agent in NSCLC. These data are not available for the pemetrexed study. Two additional trials evaluated switch maintenance therapy with erlotinib after platinum-based chemotherapy in patients with advanced

TABLE 107-12 MAINTENANCE THERAPY TRIALS

Group	CT	No. of Patients	Survival	
			OS (months)	PFS (months)
Switch Maintenance				
Fidias	Immediate docetaxel	153	12.3	5.7
	Delayed docetaxel	156	9.7	2.7
Ciuleanu	Pemetrexed	444	13.4	4.3
	BSC	222	10.6	2.6
Paramount	Pemetrexed	472	13.9	4.1
	BSC	297	11.0	2.8
ATLAS	Bev + erlotinib	384	15.9	4.8
	Bev + placebo	384	13.9	3.8
SATURN	Erlotinib	437	12.3	2.9
	Placebo	447	11.1	2.6
Continuation Maintenance				
ECOG4599	Bev 15 mg/kg	444	12.3	6.2
	BSC	434	10.3	4.5
AVAiL	Bev 15 mg/kg	351	13.4	6.5
	Bev 7.5 mg/kg	345	13.6	6.7
	Placebo	347	13.1	6.1
	Bev 15 mg/kg	347	13.1	6.1
POINTBREAK	Pemetrexed + Bev 15 mg/kg			8.6
	Bev 15 mg/kg			6.9

Abbreviations: Bev, bevacizumab; BSC, best supportive care; CT, chemotherapy; OS, overall survival; PFS, progression-free survival.

NSCLC and reported an improvement in progression-free survival and overall survival in the erlotinib treatment group. Currently, maintenance pemetrexed or erlotinib following platinum-based chemotherapy in patients with advanced NSCLC are approved by the U.S. FDA. However, maintenance therapy is not without toxicity and, at this time, should be considered on an individual patient basis.

Targeted Therapies for Select Molecular Cohorts of NSCLC As the efficacy of traditional cytotoxic chemotherapeutic agents plateaued in NSCLC, there was a critical need to define novel therapeutic treatment strategies. These novel strategies have largely been based on the identification of somatic driver mutations within the tumor. These driver mutations occur in genes encoding signaling proteins that, when aberrant, drive initiation and maintenance of tumor cells. Importantly, driver mutations can serve as Achilles' heels for tumors, if their gene products can be targeted therapeutically with small-molecule inhibitors. For example, *EGFR* mutations have been detected in 10–15% of North American patients diagnosed with NSCLC. *EGFR* mutations are associated with younger age, light (<10 pack-year) and nonsmokers, and adenocarcinoma histology. Approximately 90% of these mutations are exon 19 deletions or exon 21 L858R point mutations within the *EGFR* TK domain, resulting in hyperactivation of both *EGFR* kinase activity and downstream signaling. Lung tumors that harbor activating mutations within the *EGFR* kinase domain display high sensitivity to small-molecule *EGFR* TKIs. Erlotinib and afatinib are FDA-approved oral small-molecule TKIs that inhibit *EGFR*. Outside the United States, gefitinib also is available. Several large, international, phase III studies have demonstrated improved response rates, progression-free survival, and overall survival in patients with *EGFR* mutation-positive NSCLC patients treated with an *EGFR* TKI as compared with standard first-line chemotherapy regimens (Table 107-13).

Although response rates with *EGFR* TKI therapy are clearly superior in patients with lung tumors harboring activating *EGFR* kinase domain mutations, the *EGFR* TKI erlotinib is also FDA approved for second- and third-line therapy in patients with advanced NSCLC irrespective of tumor genotype. The reason for this apparent discrepancy is that erlotinib was initially evaluated in lung cancer before the discovery of *EGFR* activating mutations. In fact, *EGFR* mutations were initially identified in lung cancer by studying the tumors of patients who had dramatic responses to this agent. With the rapid pace of scientific discovery, additional driver mutations in lung cancer have been identified and targeted therapeutically with impressive clinical results. For example, chromosomal rearrangements involving the anaplastic lymphoma kinase (*ALK*) gene on chromosome 2 have been found in ~3–7% of NSCLC. The result of these *ALK* rearrangements is hyperactivation of the *ALK* TK domain. Similar to

EGFR, *ALK* rearrangements are typically (but not exclusively) associated with younger age, light (<10 pack-year) and nonsmokers, and adenocarcinoma histology. Remarkably, *ALK* rearrangements were initially described in lung cancer in 2007, and by 2011, the first *ALK* inhibitor, crizotinib, received FDA approval for patients with lung tumors harboring *ALK* rearrangements.

In addition to *EGFR* and *ALK*, other driver mutations have been discovered with varying frequencies in NSCLC, including *KRAS*, *BRAF*, *PIK3CA*, *NRAS*, *AKT1*, *MET*, *MEK1* (*MAP2K1*), *ROS1*, and *RET*. Mutations within the *KRAS* GTPase are found in approximately 20% of lung adenocarcinomas. To date, however, no small-molecule inhibitors are available to specifically target mutant *KRAS*. Each of the other driver mutations occurs in less than 1–3% of lung adenocarcinomas. The great majority of the driver mutations are mutually exclusive, and there are ongoing clinical studies for their specific inhibitors. For example, the *BRAF* inhibitor vemurafenib and the *RET* inhibitor cabozantinib have already demonstrated efficacy in patients with lung cancer harboring *BRAF* mutations or *RET* gene fusions, respectively. Most of these mutations are present in adenocarcinoma; however, mutations that may be linked to future targeted therapies in squamous cell carcinomas are emerging. In addition, there are active research efforts aimed at defining novel targetable mutations in lung cancer as well as defining mechanisms of acquired resistance to small-molecule inhibitors used in the treatment of patients with NSCLC.

Second-Line Chemotherapy and Beyond Second-line therapy for advanced NSCLC was almost never recommended until a seminal study in 2000 showed that docetaxel improved survival compared to supportive care alone. As first-line chemotherapy regimens improve, a substantial number of patients will maintain a good performance status and a desire for further therapy when they develop recurrent disease. Currently, several agents are FDA approved for second-line use in NSCLC including docetaxel, pemetrexed, erlotinib (approved for second-line therapy regardless of tumor genotype), and crizotinib (for patients with *ALK*-mutant lung cancer only). Most of the survival benefit for any of these agents is realized in patients who maintain a good performance status.

Immunotherapy For more than 30 years, the investigation of vaccines and immunotherapies in lung cancer has yielded little in the way of meaningful benefit. Recently, however, this perception has changed based on preliminary results of studies using monoclonal antibodies that activate antitumor immunity through blockade of immune checkpoints. For example, ipilimumab, a monoclonal antibody directed at cytotoxic T lymphocyte antigen-4 (CTLA-4), was studied in combination with paclitaxel plus carboplatin in patients with both SCLC and NSCLC. There appeared to be a small but not statistically significant advantage to the combination when ipilimumab was instituted after several cycles of chemotherapy. A randomized phase III trial in SCLC is under way to validate these data. Antibodies to the T cell programmed cell death receptor 1 (PD-1), nivolumab and pembrolizumab, have been shown to produce responses in lung cancer, renal cell cancer, and melanoma. Many of these responses have had very long duration (i.e., >1 year). Monoclonal antibodies to the PD-1 ligand (anti-PDL-1), which may be expressed on the tumor cell, have also been shown to produce responses in patients with melanoma and lung cancer. Preliminary studies in melanoma suggest that the combination of ipilimumab and nivolumab could produce higher response rates compared to either agent alone. A similar strategy is being investigated in SCLC patients. Further evaluation of these agents in both NSCLC and SCLC is ongoing in combination with already approved chemotherapy and targeted agents.

Supportive Care No discussion of the treatment strategies for patients with advanced lung cancer would be complete without a mention of supportive care. Coincident with advances in chemotherapy and targeted therapy was a pivotal study that demonstrated that the early integration of palliative care with standard treatment strategies improved both quality of life and mood for patients with advanced lung cancer. Aggressive pain and symptom control is an important component for optimal treatment of these patients.

TABLE 107-13 RESULTS OF PHASE III TRIALS COMPARING CHEMOTHERAPY AND FIRST-LINE *EGFR* TKI IN *EGFR* MUTATION-POSITIVE PATIENTS

Study	Therapy	No. of Patients	ORR (%)	PFS (months)
IPASS	CbP	129	47	6.3
	Gefitinib	132	71	9.3
EURTAC	CG	87	15	5.2
	Erlotinib	86	58	9.7
OPTIMAL	CG	72	36	4.6
	Erlotinib	82	83	13.1
NEJ002	CG	114	31	5.4
	Gefitinib	114	74	10.8
WJTOG3405	CD	89	31	6.3
	Gefitinib	88	62	9.2
LUX LUNG 3	CP	115	23	6.9
	Afatinib	230	56	11.1

Abbreviations: CbP, carboplatin and paclitaxel; CD, cisplatin and docetaxel; CG, cisplatin and gemcitabine; CP, cisplatin and paclitaxel; ORR, overall response rate; PFS, progression-free survival.

SURGERY FOR LIMITED-DISEASE SMALL-CELL LUNG CANCER

SCLC is a highly aggressive disease characterized by its rapid doubling time, high growth fraction, early development of disseminated disease, and dramatic response to first-line chemotherapy and radiation. In general, surgical resection is *not* routinely recommended for patients because even patients with LD-SCLC still have occult micrometastases. However, the most recent American College of Chest Physicians Evidence-Based Clinical Practice Guidelines recommend surgical resection over nonsurgical treatment in SCLC patients with clinical stage I disease after a thorough evaluation for distant metastases and invasive mediastinal stage evaluation (grade 2C). After resection, these patients should receive platinum-based adjuvant chemotherapy (grade 1C). If the histologic diagnosis of SCLC is made in patients on review of a resected surgical specimen, such patients should receive standard SCLC chemotherapy as well.

CHEMOTHERAPY

Chemotherapy significantly prolongs survival in patients with SCLC. Four to six cycles of platinum-based chemotherapy with either cisplatin or carboplatin plus either etoposide or irinotecan has been the mainstay of treatment for nearly three decades and is recommended over other chemotherapy regimens irrespective of initial stage. Cyclophosphamide, doxorubicin (Adriamycin), and vincristine (CAV) may be an alternative for patients who are unable to tolerate a platinum-based regimen. Despite response rates to first-line therapy as high as 80%, the median survival ranges from 12 to 20 months for patients with LD and from 7 to 11 months for patients with ED. Regardless of disease extent, the majority of patients relapse and develop chemotherapy-resistant disease. Only 6–12% of patients with LD-SCLC and 2% of patients with ED-SCLC live beyond 5 years. The prognosis is especially poor for patients who relapse within the first 3 months of therapy; these patients are said to have *chemotherapy-resistant disease*. Patients are said to have *sensitive disease* if they relapse more than 3 months after their initial therapy and are thought to have a somewhat better overall survival. These patients also are thought to have the greatest potential benefit from second-line chemotherapy (Fig. 107-7). Topotecan is the

only FDA-approved agent for second-line therapy in patients with SCLC. Topotecan has only modest activity and can be given either intravenously or orally. In one randomized trial, 141 patients who were not considered candidates for further IV chemotherapy were randomized to receive either oral topotecan or best supportive care. Although the response rate to oral topotecan was only 7%, overall survival was significantly better in patients receiving chemotherapy (median survival time, 26 weeks vs 14 weeks; $p = .01$). Moreover, patients given topotecan had a slower decline in quality of life than did those not receiving chemotherapy. Other agents with similar low levels of activity in the second-line setting include irinotecan, paclitaxel, docetaxel, vinorelbine, oral etoposide, and gemcitabine. Clearly novel treatments for this all too common disease are desperately needed.

THORACIC RADIATION THERAPY

Thoracic radiation therapy (TRT) is a standard component of induction therapy for good performance status and limited-stage SCLC patients. Meta-analyses indicate that chemotherapy combined with chest irradiation improves 3-year survival by approximately 5% as compared with chemotherapy alone. The 5-year survival rate, however, remains disappointingly low at ~10–15%. Most commonly, TRT is combined with cisplatin and etoposide chemotherapy due to a superior toxicity profile as compared to anthracycline-containing chemotherapy regimens. As observed in locally advanced NSCLC, *concurrent* chemoradiotherapy is more effective than *sequential* chemoradiation but is associated with significantly more esophagitis and hematologic toxicity. Ideally TRT should be administered with the first two cycles of chemotherapy because later application appears slightly less effective. If for reasons of fitness or availability, this regimen cannot be offered, TRT should follow induction chemotherapy. With respect to fractionation of TRT, twice-daily 1.5-Gy fractionated radiation therapy has been shown to improve survival in LD-SCLC patients but is associated with higher rates of grade 3 esophagitis and pulmonary toxicity. Although it is feasible to deliver once-daily radiation therapy doses up to 70 Gy concurrently with cisplatin-based chemotherapy, there are no data to support equivalency of this approach compared with the 45-Gy twice-daily radiotherapy dose. Therefore, the current standard regimen of a 45-Gy dose administered in 1.5-Gy fractions twice daily for 30 days is being compared with higher-dose regimens in two phase III trials, one in the United States and one in Europe. Patients should be carefully selected for concurrent chemoradiation therapy based on good performance status and adequate pulmonary reserve. The role of radiotherapy in ED-SCLC is largely restricted to palliation of tumor-related symptoms such as bone pain and bronchial obstruction.

PROPHYLACTIC CRANIAL IRRADIATION

Prophylactic cranial irradiation (PCI) should be considered in all patients with either LD-SCLC or ED-SCLC who have responded well to initial therapy. A meta-analysis including seven trials and 987 patients with LD-SCLC who had achieved a complete remission after upfront chemotherapy yielded a 5.4% improvement in overall survival for patients treated with PCI. In patients with ED-SCLC who have responded to first-line chemotherapy, a prospective randomized phase III trial showed that PCI reduced the occurrence of symptomatic brain metastases and prolonged disease-free and overall survival compared to no radiation therapy. Long-term toxicities, including deficits in cognition, have been reported after PCI but are difficult to sort out from the effects of chemotherapy or normal aging.

SUMMARY

The management of NSCLC has undergone major change in the past decade. To a lesser extent, the same is true for SCLC. For patients with early-stage disease, advances in radiotherapy and surgical procedures as well as new systemic therapies have greatly improved prognosis in both diseases. For patients with advanced disease, major progress in understanding tumor genetics has led to the development of

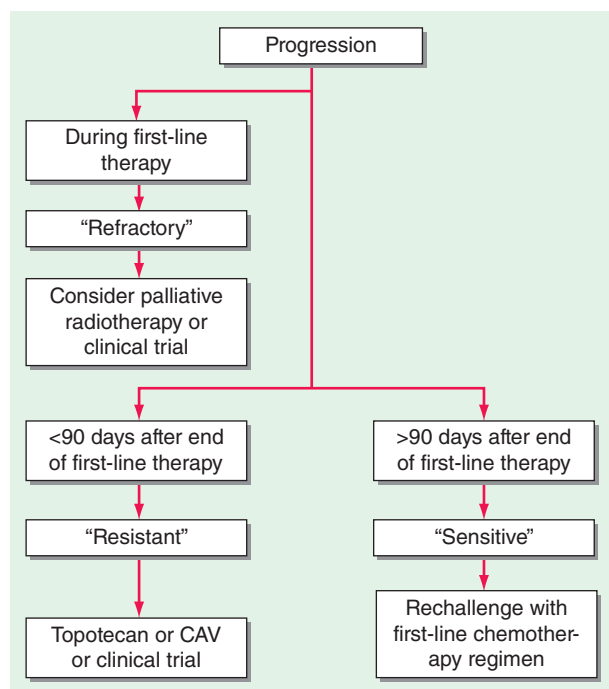


FIGURE 107-7 Management of recurrent small-cell lung cancer (SCLC). CAV, cyclophosphamide, doxorubicin, and vincristine. (Adapted with permission from JP van Meerbeeck et al: *Lancet* 378:1741, 2011.)

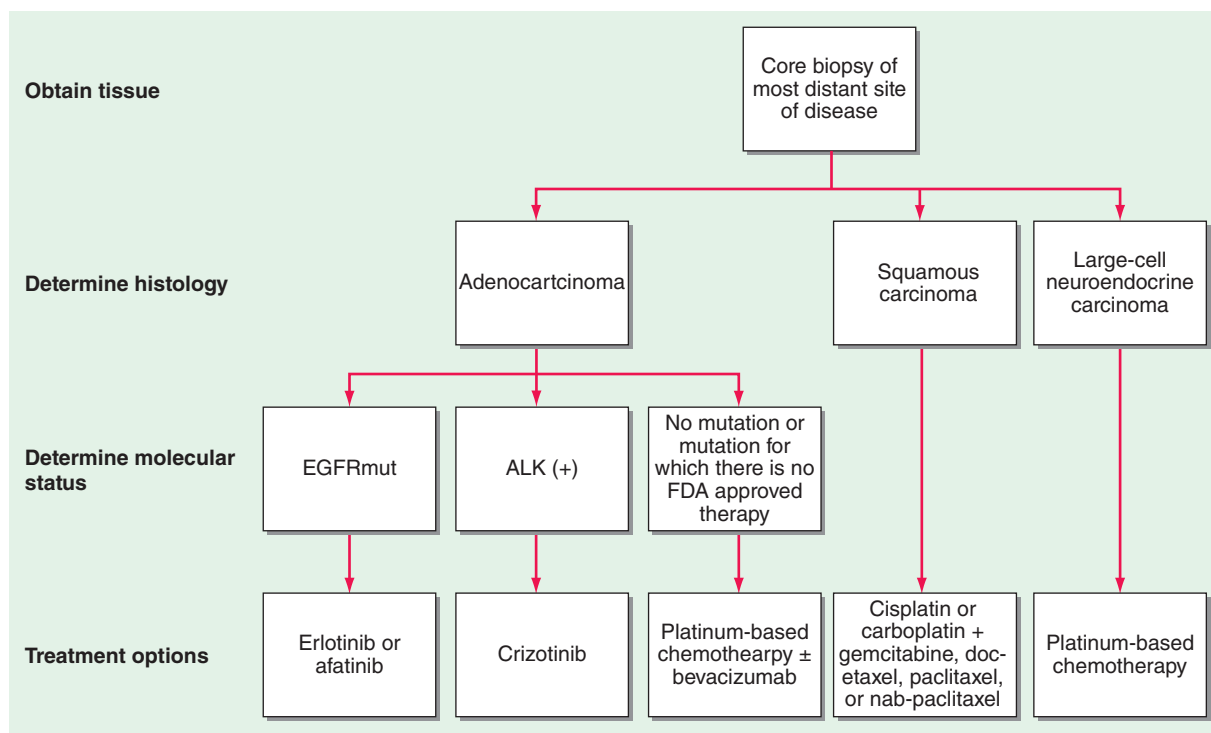


FIGURE 107-8 Approach to first-line therapy in a patient with stage IV non-small-cell lung cancer (NSCLC). EGFRmut, *EGFR* mutation; FDA, Food and Drug Administration.

targeted inhibitors based specifically on the tumor's molecular profile. Furthermore, increased understanding of how to activate the immune system to drive antitumor immunity is proving to be a promising therapeutic strategy for some patients with advanced lung cancer. In Fig. 107-8, we propose an algorithm of the treatment approach for patient with stage IV NSCLC. However, the reality is that the majority of patients treated with targeted therapies or chemotherapy eventually develop resistance, which provides strong motivation for further research and enrollment of patients onto clinical trials in this rapidly evolving area.

Thus, breast cancer may exist for a long period as either a noninvasive disease or an invasive but nonmetastatic disease. These facts have significant clinical ramifications.

Not more than 10% of human breast cancers can be linked directly to germline mutations. Several genes have been implicated in familial cases. The Li-Fraumeni syndrome is characterized by inherited mutations in the *p53* tumor-suppressor gene, which lead to an increased incidence of breast cancer, osteogenic sarcomas, and other malignancies. Inherited mutations in *PTEN* have also been reported in breast cancer.

Another tumor-suppressor gene, *BRCA1*, has been identified at the chromosomal locus 17q21; this gene encodes a zinc finger protein, and the protein product functions as a transcription factor and is involved in gene repair. Women who inherit a mutated allele of this gene from either parent have at least a 60–80% lifetime chance of developing breast cancer and about a 33% chance of developing ovarian cancer. The risk is higher among women born after 1940, presumably due to promotional effects of hormonal factors. Men who carry a mutant allele of the gene have an increased incidence of prostate cancer and breast cancer. A fourth gene, termed *BRCA2*, which has been localized to chromosome 13q12, is also associated with an increased incidence of breast cancer in men and women.

Germline mutations in *BRCA1* and *BRCA2* can be readily detected; patients with these mutations should be counseled appropriately. All women with strong family histories for breast cancer should be referred to genetic screening programs, particularly women of Ashkenazi Jewish descent who have a high likelihood of a specific founder *BRCA1* mutation (substitution of adenine for guanine at position 185).

Even more important than the role these genes play in inherited forms of breast cancer may be their role in sporadic breast cancer. A *p53* mutation is present in nearly 40% of human breast cancers as an acquired defect. Acquired mutations in *PTEN* occur in about 10% of the cases. *BRCA1* mutation in sporadic primary breast cancer has not been reported. However, decreased expression of *BRCA1* mRNA (possibly via gene methylation) and abnormal cellular location of the *BRCA1* protein have been found in some breast cancers. Loss of heterozygosity of *BRCA1* and *BRCA2* suggests that tumor-suppressor

108 Breast Cancer

Marc E. Lippman

Breast cancer is a malignant proliferation of epithelial cells lining the ducts or lobules of the breast. In the year 2014, about 180,000 cases of invasive breast cancer and 40,000 deaths will occur in the United States. In addition, about 2000 men will be diagnosed with breast cancer. Epithelial malignancies of the breast are the most common cause of cancer in women (excluding skin cancer), accounting for about one-third of all cancer in women. As a result of improved treatment and earlier detection, the mortality rate from breast cancer has begun to decrease very substantially in the United States. This Chapter will not consider rare malignancies presenting in the breast, such as sarcomas and lymphomas, but will focus on the epithelial cancers.

GENETIC CONSIDERATIONS



Human breast cancer is a clonal disease; a single transformed cell—the product of a series of somatic (acquired) or germline mutations—is eventually able to express full malignant potential.

524 activity may be inactivated in sporadic cases of human breast cancer. Finally, increased expression of a dominant oncogene plays a role in about a quarter of human breast cancer cases. The product of this gene, a member of the epidermal growth factor receptor superfamily, is called *erbB2* (HER/2 neu) and is overexpressed in these breast cancers due to gene amplification; this overexpression can contribute to transformation of human breast epithelium and is the target of effective systemic therapy in adjuvant and metastatic disease settings. A series of acquired “driver” mutations have been identified in sporadic breast cancer by major sequencing consortia. Unfortunately, most occur in no more than 5% of cases and generally do not have effective agents to target them, so “personalized medicine” is for now more of a dream than a reality.

EPIDEMIOLOGY

Breast cancer is a hormone-dependent disease. Women without functioning ovaries who never receive estrogen replacement therapy do not develop breast cancer. The female-to-male ratio is about 150:1. For most epithelial malignancies, a log-log plot of incidence versus age shows a single-component straight-line increase with every year of life. A similar plot for breast cancer shows two components: a straight-line increase with age but with a decrease in slope beginning at the age of menopause. The three dates in a woman’s life that have a major impact on breast cancer incidence are age at menarche, age at first full-term pregnancy, and age at menopause. Women who experience menarche at age 16 years have only 50–60% of the breast cancer risk of a woman having menarche at age 12 years; the lower risk persists throughout life. Similarly, menopause occurring 10 years before the median age of menopause (52 years), whether natural or surgically induced, reduces lifetime breast cancer risk by about 35%. Women who have a first full-term pregnancy by age 18 years have a 30–40% lower risk of breast cancer compared with nulliparous women. Thus, length of menstrual life—particularly the fraction occurring before first full-term pregnancy—is a substantial component of the total risk of breast cancer. These three factors (menarche, age of first full-term pregnancy, and menopause) can account for 70–80% of the variation in breast cancer frequency in different countries. Also, duration of maternal nursing correlates with substantial risk reduction independent of either parity or age at first full-term pregnancy.

International variation in incidence has provided some of the most important clues on hormonal carcinogenesis. A woman living to age 80 years in North America has one chance in nine of developing invasive breast cancer. Asian women have one-fifth to one-tenth the risk of breast cancer of women in North America or Western Europe. Asian women have substantially lower concentrations of estrogens and progesterone. These differences cannot be explained on a genetic basis because Asian women living in a Western environment have sex steroid hormone concentrations and risks identical to those of their Western counterparts. These migrant women, and more notably their daughters, also differ markedly in height and weight from Asian women in Asia; height and weight are critical regulators of age of menarche and have substantial effects on plasma concentrations of estrogens.

The role of diet in breast cancer etiology is controversial. While there are associative links between total caloric and fat intake and breast cancer risk, the exact role of fat in the diet is unproven. Increased caloric intake contributes to breast cancer risk in multiple ways: earlier menarche, later age at menopause, and increased postmenopausal estrogen concentrations reflecting enhanced aromatase activities in fatty tissues. On the other hand, central obesity is both a risk factor for occurrence and recurrence of breast cancer. Moderate alcohol intake also increases the risk by an unknown mechanism. Folic acid supplementation appears to modify risk in women who use alcohol but is not additionally protective in abstainers. Recommendations favoring abstinence from alcohol must be weighed against other social pressures and the possible cardioprotective effect of moderate alcohol intake. Chronic low-dose aspirin use is associated with a decreased incidence of breast cancer. Depression is also associated with both occurrence and recurrence of breast cancer.

Understanding the potential role of exogenous hormones in breast cancer is of extraordinary importance because millions of American women regularly use oral contraceptives and postmenopausal hormone replacement therapy. The most credible meta-analyses of oral contraceptive use suggest that these agents cause a small increased risk of breast cancer. By contrast, oral contraceptives offer a substantial protective effect against ovarian epithelial tumors and endometrial cancers. Hormone replacement therapy (HRT) has a powerful effect on breast cancer risk. Data from the Women’s Health Initiative (WHI) trial showed that conjugated equine estrogens plus progestins increased the risk of breast cancer and adverse cardiovascular events but decreased the risk of bone fractures and colorectal cancer. On balance, there were more negative events with HRT; 6–7 years of HRT nearly doubled the risk of breast cancer. A parallel WHI trial with >12,000 women enrolled testing conjugated estrogens alone (estrogen replacement therapy in women who have had hysterectomies) showed no significant increase in breast cancer incidence. Thus, there are serious concerns about long-term HRT use in terms of cardiovascular disease and breast cancer. The WHI trial of conjugated equine estrogen alone demonstrated few adverse effects for women age <70; however, no comparable safety data are available for other more potent forms of estrogen replacement, and they should not be routinely used as substitutes. HRT in women previously diagnosed with breast cancer increases recurrence rates. Rapid decrease in the number of women on HRT has already led to a coincident decrease in breast cancer incidence.

In addition to the other factors, radiation is a risk factor in younger women. Women who have been exposed before age 30 years to radiation in the form of multiple fluoroscopies (200–300 cGy) or treatment for Hodgkin’s disease (>3600 cGy) have a substantial increase in risk of breast cancer, whereas radiation exposure after age 30 years appears to have a minimal carcinogenic effect on the breast.

EVALUATION OF BREAST MASSES IN MEN AND WOMEN

Because the breasts are a common site of potentially fatal malignancy in women, examination of the breast is an essential part of the physical examination. Unfortunately, internists frequently do not examine breasts in men, and in women, they are apt to defer this evaluation to gynecologists. Because of the plausible association between early detection and improved outcome, it is the duty of every physician to identify breast abnormalities at the earliest possible stage and to institute a diagnostic workup. Women should be trained in breast self-examination (BSE). Although breast cancer in men is unusual, unilateral lesions should be evaluated in the same manner as in women, with the recognition that gynecomastia in men can sometimes begin unilaterally and is often asymmetric.

Virtually all breast cancer is diagnosed by biopsy of a nodule detected either on a mammogram or by palpation. Algorithms have been developed to enhance the likelihood of diagnosing breast cancer and reduce the frequency of unnecessary biopsy (Fig. 108-1).

THE PALPABLE BREAST MASS

Women should be strongly encouraged to examine their breasts monthly. A potentially flawed study from China has suggested that BSE does not alter survival, but given its safety, the procedure should still be encouraged. At worst, this practice increases the likelihood of detecting a mass at a smaller size when it can be treated with more limited surgery. Breast examination by the physician should be performed in good light so as to see retractions and other skin changes. The nipple and areolae should be inspected, and an attempt should be made to elicit nipple discharge. All regional lymph node groups should be examined, and any lesions should be measured. Physical examination alone cannot exclude malignancy. Lesions with certain features are more likely to be cancerous (hard, irregular, tethered or fixed, or painless lesions). A negative mammogram in the presence of a persistent lump in the breast does not exclude malignancy. Palpable lesions require additional diagnostic procedures, including biopsy.

In premenopausal women, lesions that are either equivocal or nonsuspicious on physical examination should be reexamined in 2–4

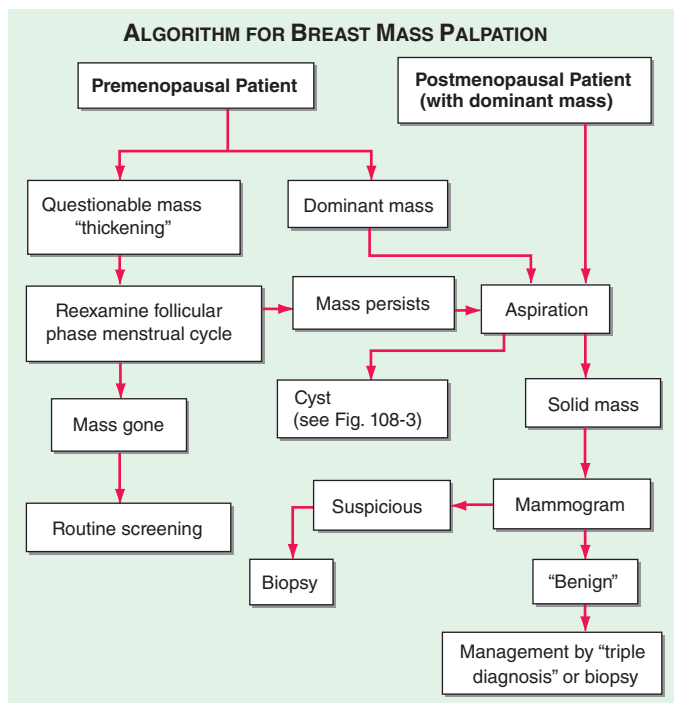


FIGURE 108-1 Approach to a palpable breast mass.

weeks, during the follicular phase of the menstrual cycle. Days 5–7 of the cycle are the best time for breast examination. A dominant mass in a postmenopausal woman or a dominant mass that persists through a menstrual cycle in a premenopausal woman should be aspirated by fine-needle biopsy or referred to a surgeon. If nonbloody fluid is aspirated, the diagnosis (cyst) and therapy have been accomplished together. Solid lesions that are persistent, recurrent, complex, or bloody cysts require mammography and biopsy, although in selected patients the so-called triple diagnostic technique (palpation, mammography, aspiration) can be used to avoid biopsy (Figs. 108-1, 108-2, and 108-3). Ultrasound can be used in place of fine-needle aspiration to distinguish cysts from solid lesions. Not all solid masses are detected by ultrasound; thus, a palpable mass that is not visualized on ultrasound must be presumed to be solid.

Several points are essential in pursuing these management decision trees. First, risk-factor analysis is not part of the decision structure. No constellation of risk factors, by their presence or absence, can be used to exclude biopsy. Second, fine-needle aspiration should be used only in centers that have proven skill in obtaining such specimens and analyzing them. The likelihood of cancer is low in the setting of a “triple negative” (benign-feeling lump, negative mammogram, and negative fine-needle aspiration), but it is not zero. The patient and physician

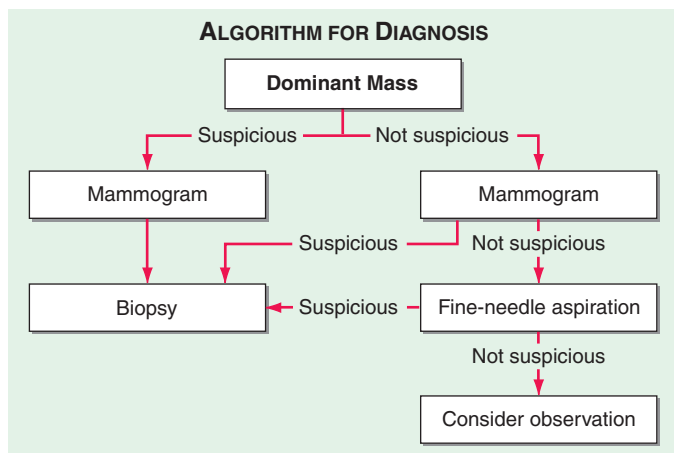


FIGURE 108-2 The “triple diagnosis” technique.

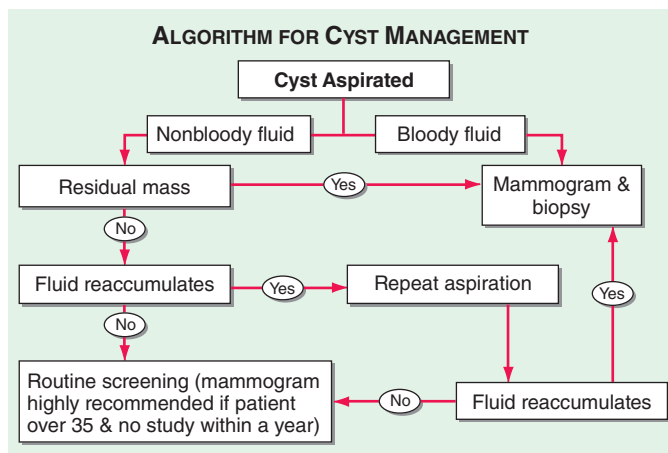


FIGURE 108-3 Management of a breast cyst.

must be aware of a 1% risk of false negatives. Third, additional technologies such as magnetic resonance imaging (MRI), ultrasound, and sestamibi imaging cannot be used to exclude the need for biopsy, although in unusual circumstances, they may provoke a biopsy.

THE ABNORMAL MAMMOGRAM

Diagnostic mammography should not be confused with *screening mammography*, which is performed after a palpable abnormality has been detected. Diagnostic mammography is aimed at evaluating the rest of the breast before biopsy is performed or occasionally is part of the triple-test strategy to exclude immediate biopsy.

Subtle abnormalities that are first detected by screening mammography should be evaluated carefully by compression or magnified views. These abnormalities include clustered microcalcifications, densities (especially if spiculated), and new or enlarging architectural distortion. For some nonpalpable lesions, ultrasound may be helpful either to identify cysts or to guide biopsy. If there is no palpable lesion and detailed mammographic studies are unequivocally benign, the patient should have routine follow-up appropriate to the patient’s age. It cannot be stressed too strongly that in the presence of a breast lump a negative mammogram does not rule out cancer.

If a nonpalpable mammographic lesion has a low index of suspicion, mammographic follow-up in 3–6 months is reasonable. Workup of indeterminate and suspicious lesions has been rendered more complex by the advent of stereotactic biopsies. Morrow and colleagues have suggested that these procedures are indicated for lesions that require biopsy but are likely to be benign—that is, for cases in which the procedure probably will eliminate additional surgery. When a lesion is more probably malignant, open biopsy should be performed with a needle localization technique. Others have proposed more widespread use of stereotactic core biopsies for nonpalpable lesions on economic grounds and because diagnosis leads to earlier treatment planning. However, stereotactic diagnosis of a malignant lesion does not eliminate the need for definitive surgical procedures, particularly if breast conservation is attempted. For example, after a breast biopsy with needle localization (i.e., local excision) of a stereotactically diagnosed malignancy, reexcision may still be necessary to achieve negative margins. To some extent, these issues are decided on the basis of referral pattern and the availability of the resources for stereotactic core biopsies. A reasonable approach is shown in Fig. 108-4.

BREAST MASSES IN THE PREGNANT OR LACTATING WOMAN

During pregnancy, the breast grows under the influence of estrogen, progesterone, prolactin, and human placental lactogen. Lactation is suppressed by progesterone, which blocks the effects of prolactin. After delivery, lactation is promoted by the fall in progesterone levels, which leaves the effects of prolactin unopposed. The development of a dominant mass during pregnancy or lactation should never be attributed to hormonal changes. A dominant mass must be treated with the same concern in a pregnant woman as any other. Breast

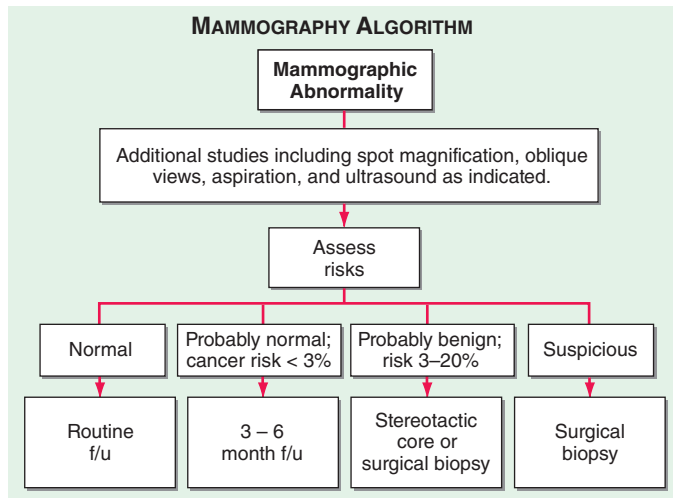


FIGURE 108-4 Approaches to abnormalities detected by mammogram.

cancer develops in 1 in every 3000–4000 pregnancies. Stage for stage, breast cancer in pregnant patients is no different from premenopausal breast cancer in nonpregnant patients. However, pregnant women often have more advanced disease because the significance of a breast mass was not fully considered and/or because of endogenous hormone stimulation. Persistent lumps in the breast of pregnant or lactating women *cannot* be attributed to benign changes based on physical findings; such patients should be promptly referred for diagnostic evaluation.

BENIGN BREAST MASSES

Only about 1 in every 5–10 breast biopsies leads to a diagnosis of cancer, although the rate of positive biopsies varies in different countries and clinical settings. (These differences may be related to interpretation, medico-legal considerations, and availability of mammograms.) The vast majority of benign breast masses are due to “fibrocystic” disease, a descriptive term for small fluid-filled cysts and modest epithelial cell and fibrous tissue hyperplasia. However, fibrocystic disease is a histologic, not a clinical, diagnosis, and women who have had a biopsy with benign findings are at greater risk of developing breast cancer than those who have not had a biopsy. The subset of women with ductal or lobular cell proliferation (about 30% of patients), particularly the small fraction (3%) with atypical hyperplasia, have a fourfold greater risk of developing breast cancer than those women who have not had a biopsy, and the increase in the risk is about ninefold for women in this category who also have an affected first-degree relative. Thus, careful follow-up of these patients is required. By contrast, patients with a benign biopsy without atypical hyperplasia are at little risk and may be followed routinely.

SCREENING

Breast cancer is virtually unique among the epithelial tumors in adults in that screening (in the form of annual mammography) improves survival. Meta-analysis examining outcomes from every randomized trial of mammography conclusively shows a 25–30% reduction in the chance of dying from breast cancer with annual screening after age 50 years; the data for women between ages 40 and 50 years are almost as positive; however, since the incidence is much lower in younger women, there are more false positives. While controversy continues to surround the assessment of screening mammography, the preponderance of data strongly supports the benefits of screening mammography. New analyses of older randomized studies have occasionally suggested that screening may not work. While the design defects in some older studies cannot be retrospectively corrected, most experts, including panels of the American Society of Clinical Oncology and the American Cancer Society (ACS), continue to believe that screening conveys substantial benefit. Furthermore, the profound drop in breast

cancer mortality rate seen over the past decade is unlikely to be solely attributable to improvements in therapy. It seems prudent to recommend annual or biannual mammography for women past the age of 40 years. Although no randomized study of BSE has ever shown any improvement in survival, its major benefit is identification of tumors appropriate for conservative local therapy. Better mammographic technology, including digitized mammography, routine use of magnified views, and greater skill in mammographic interpretation, combined with newer diagnostic techniques (MRI, magnetic resonance spectroscopy, positron emission tomography, etc.) may make it possible to identify breast cancers even more reliably and earlier. Screening by any technique other than mammography is not indicated. However, the ACS suggests that younger women who are *BRCA1* or *BRCA2* carriers or untested first-degree relatives of women with cancer; women with a history of radiation therapy to the chest between ages 10 and 30 years; women with a lifetime risk of breast cancer of at least 20%; and women with a history of Li-Fraumeni, Cowden, or Bannayan-Riley-Ruvalcaba syndromes may benefit from MRI screening, where the higher sensitivity may outweigh the loss of specificity.

STAGING

Correct staging of breast cancer patients is of extraordinary importance. Not only does it permit an accurate prognosis, but in many cases, therapeutic decision-making is based largely on the TNM (primary tumor, regional nodes, metastasis) classification (Table 108-1). Comparison with historic series should be undertaken with caution, as the staging has changed several times in the past 20 years. The current staging is complex and results in significant changes in outcome by stage as compared with prior staging systems.

TREATMENT BREAST CANCER

One of the most exciting aspects of breast cancer biology has been its subdivision into at least five subtypes based on gene expression profiling.

- 1. Luminal A:** The luminal tumors express cytokeratins 8 and 18, have the highest levels of estrogen receptor expression, tend to be low grade, are most likely to respond to endocrine therapy, and have a favorable prognosis. They tend to be less responsive to chemotherapy.
- 2. Luminal B:** Tumor cells are also of luminal epithelial origin, but with a gene expression pattern distinct from luminal A. Prognosis is somewhat worse than luminal A.
- 3. Normal breast-like:** These tumors have a gene expression profile reminiscent of nonmalignant “normal” breast epithelium. Prognosis is similar to the luminal B group. This subtype is somewhat controversial and may represent contamination of the sample by normal mammary epithelium.
- 4. HER2 amplified:** These tumors have amplification of the *HER2* gene on chromosome 17q and frequently exhibit coamplification and overexpression of other genes adjacent to *HER2*. Historically the clinical prognosis of such tumors was poor. However, with the advent of trastuzumab and other targeted therapies, the clinical outcome of *HER2*-positive patients is markedly improving.
- 5. Basal:** These estrogen receptor/progesterone receptor–negative and *HER2*-negative tumors (so-called triple negative) are characterized by markers of basal/myoepithelial cells. They tend to be high grade, and express cytokeratins 5/6 and 17 as well as vimentin, p63, CD10, α -smooth muscle actin, and epidermal growth factor receptor (EGFR). Patients with *BRCA* mutations also fall within this molecular subtype. They also have stem cell characteristics.

PRIMARY BREAST CANCER

Breast-conserving treatments, consisting of the removal of the primary tumor by some form of lumpectomy with or without irra-

TABLE 108-1 STAGING OF BREAST CANCER

Primary Tumor (T)			
T0	No evidence of primary tumor		
TIS	Carcinoma in situ		
T1	Tumor ≤2 cm		
T1a	Tumor >0.1 cm but ≤0.5 cm		
T1b	Tumor >0.5 but ≤1 cm		
T1c	Tumor >1 cm but ≤2 cm		
T2	Tumor >2 cm but ≤5 cm		
T3	Tumor >5 cm		
T4	Extension to chest wall, inflammation, satellite lesions, ulcerations		
Regional Lymph Nodes (N)			
PN0(i-)	No regional lymph node metastasis histologically, negative IHC		
PN0(i+)	No regional lymph node metastasis histologically, positive IHC, no IHC cluster greater than 0.2 mm		
PN0(mol-)	No regional lymph node metastasis histologically, negative molecular findings (RT-PCR)		
PN0(mol+)	No regional lymph node metastasis histologically, positive molecular findings (RT-PCR)		
PN1	Metastasis in one to three axillary lymph nodes, or in internal mammary nodes with microscopic disease detected by sentinel lymph node dissection but not clinically apparent		
PN1mi	Micrometastasis (>0.2 mm, none >2 mm)		
PN1a	Metastasis in one to three axillary lymph nodes		
PN1b	Metastasis in internal mammary nodes with microscopic disease detected by sentinel lymph node dissection but not <i>clinically apparent</i> ^a		
PN1c	Metastasis in one to three axillary lymph nodes and in internal mammary lymph nodes with microscopic disease detected by sentinel lymph node dissection but not clinically apparent. ^a (If associated with greater than three positive axillary lymph nodes, the internal mammary nodes are classified as pN3b to reflect increased tumor burden.)		
pN2	Metastasis in four to nine axillary lymph nodes, or in clinically apparent internal mammary lymph nodes in the <i>absence</i> of axillary lymph node metastasis		
pN3	Metastasis in 10 or more axillary lymph nodes, or in infraclavicular lymph nodes, or in clinically apparent ^a ipsilateral internal mammary lymph nodes in the <i>presence</i> of 1 or more positive axillary lymph nodes; or in more than 3 axillary lymph nodes with clinically negative microscopic metastasis in internal mammary lymph nodes; or in ipsilateral subcarinal lymph nodes		
Distant Metastasis (M)			
M0	No distant metastasis		
M1	Distant metastasis (includes spread to ipsilateral supraclavicular nodes)		
Stage Grouping			
Stage 0	TIS	N0	M0
Stage I	T1	N0	M0
Stage IIA	T0	N1	M0
	T1	N1	M0
	T2	N0	M0
Stage IIB	T2	N1	M0
	T3	N0	M0
	T3	N1, N2	M0
Stage IIIA	T0	N2	M0
	T1	N2	M0
	T2	N2	M0
	T3	N1, N2	M0
Stage IIIB	T4	N0, N1, N2	M0
Stage IIIC	Any T	N3	M0
Stage IV	Any T	Any N	M1

^aClinically apparent is defined as detected by imaging studies (excluding lymphoscintigraphy) or by clinical examination.

Abbreviations: IHC, immunohistochemistry; RT-PCR, reverse transcriptase polymerase chain reaction.

Source: Used with permission of the American Joint Committee on Cancer (AJCC), Chicago, Illinois. The original source for this material is the *AJCC Cancer Staging Manual*, 7th ed. New York, Springer, 2010; www.springeronline.com.

diating the breast, result in a survival that is as good as (or slightly superior to) that after extensive surgical procedures, such as mastectomy or modified radical mastectomy, with or without further irradiation. Postlumpectomy breast irradiation greatly reduces the risk of recurrence in the breast. While breast conservation is associated with a possibility of recurrence in the breast, 10-year survival is at least as good as that after more extensive surgery. Postoperative radiation to regional nodes following mastectomy is also associated with an improvement in survival. Because radiation therapy can also reduce the rate of local or regional recurrence, it should be strongly considered following mastectomy for women with high-risk primary

tumors (i.e., T2 in size, positive margins, positive nodes). At present, nearly one-third of women in the United States are managed by lumpectomy. Breast-conserving surgery is not suitable for all patients: it is not generally suitable for tumors >5 cm (or for smaller tumors if the breast is small), for tumors involving the nipple-areola complex, for tumors with extensive intraductal disease involving multiple quadrants of the breast, for women with a history of collagen-vascular disease, and for women who either do not have the motivation for breast conservation or do not have convenient access to radiation therapy. However, these groups probably do not account for more than one-third of patients who are treated with

528 mastectomy. Thus, a great many women still undergo mastectomy who could safely avoid this procedure and probably would if appropriately counseled.

Sentinel lymph node biopsy (SLNB) is generally the standard of care for women with localized breast cancer and clinically negative axilla. If SLNB is negative, more extensive axillary surgery is not required, avoiding much of the risk of lymphedema following more extensive axillary dissections. In the presence of minimal involvement of a sentinel lymph node, further axillary surgery is not required.

An extensive intraductal component is a predictor of recurrence in the breast, and so are several clinical variables. Both axillary lymph node involvement and involvement of vascular or lymphatic channels by metastatic tumor in the breast are associated with a higher risk of relapse in the breast but are not contraindications to breast-conserving treatment. When these patients are excluded, and when lumpectomy with negative tumor margins is achieved, breast conservation is associated with a recurrence rate in the breast of 5% or less. The survival of patients who have recurrence in the breast is somewhat worse than that of women who do not. Thus, recurrence in the breast is a negative prognostic variable for long-term survival. However, recurrence in the breast is not the cause of distant metastasis. If recurrence in the breast caused metastatic disease, then women treated with lumpectomy, who have a higher rate of recurrence in the breast, should have poorer survival than women treated with mastectomy, and they do not. Most patients should consult with a radiation oncologist before making a final decision concerning local therapy. However, a multimodality clinic in which the surgeon, radiation oncologist, medical oncologist, and other caregivers cooperate to evaluate the patient and develop a treatment plan is usually considered a major advantage by patients.

Adjuvant Therapy The use of systemic therapy after local management of breast cancer substantially improves survival. More than half of the women who would otherwise die of metastatic breast cancer remain disease-free when treated with the appropriate systemic regimen. These data have grown more and more impressive with longer follow-up and more effective regimens.

PROGNOSTIC VARIABLES The most important prognostic variables are provided by *tumor staging*. The size of the tumor and the status of the axillary lymph nodes provide reasonably accurate information on the likelihood of tumor relapse. The relation of pathologic stage to 5-year survival is shown in **Table 108-2**. For most women, the need for adjuvant therapy can be readily defined on this basis alone. In the absence of lymph node involvement, involvement of microvessels (either capillaries or lymphatic channels) in tumors is nearly equivalent to lymph node involvement. The greatest controversy concerns women with intermediate prognoses. *There is rarely justification for adjuvant chemotherapy in most women with tumors <1 cm in size whose axillary lymph nodes are negative. HER2-positive tumors are a potential exception.* Detection of breast cancer cells either in the circulation or bone marrow is associated with an increased relapse rate. The most exciting development in this area is the use of gene expression arrays to analyze patterns of tumor gene expression. Several groups have independently defined gene sets that reliably predict disease-free and overall survival far more accu-

rately than any single prognostic variable including the Oncotype DX[®] analysis of 21 genes. Also, the use of such standardized risk assessment tools such as Adjuvant! Online (www.adjuvantonline.com) is very helpful. These tools are highly recommended in otherwise ambiguous circumstances.

Estrogen receptor status and progesterone receptor status are of prognostic significance. Tumors that lack either or both of these receptors are more likely to recur than tumors that have them.

Several *measures of tumor growth rate* correlate with early relapse. S-phase analysis using flow cytometry is the most accurate measure. Indirect S-phase assessments using antigens associated with the cell cycle, such as PCNA (Ki67), are also valuable. Tumors with a high proportion (more than the median) of cells in S-phase pose a greater risk of relapse; chemotherapy offers the greatest survival benefit for these tumors. Assessment of DNA content in the form of ploidy is of modest value, with nondiploid tumors having a somewhat worse prognosis.

Histologic classification of the tumor has also been used as a prognostic factor. Tumors with a poor nuclear grade have a higher risk of recurrence than tumors with a good nuclear grade. Semiquantitative measures such as the Elston score improve the reproducibility of this measurement.

Molecular changes in the tumor are also useful. Tumors that overexpress *erbB2* (HER2/neu) or have a mutated *p53* gene have a worse prognosis. Particular interest has centered on *erbB2* overexpression as measured by immunohistochemistry or fluorescence in situ hybridization. Tumors that overexpress *erbB2* are more likely to respond to doxorubicin-containing regimens; *erbB2* overexpression also predicts those tumors that will respond to HER2/neu antibodies (trastuzumab) (Herceptin) and HER2/neu kinase inhibitors.

Other variables that have also been used to evaluate prognosis include proteins associated with invasiveness, such as type IV collagenase, cathepsin D, plasminogen activator, plasminogen activator receptor, and the metastasis-suppressor gene *nm23*. None of these has been widely accepted as a prognostic variable for therapeutic decision-making. One problem in interpreting these prognostic variables is that most of them have not been examined in a study using a large cohort of patients.

ADJUVANT REGIMENS Adjuvant therapy is the use of systemic therapies in patients whose known disease has received local therapy but who are at risk of relapse. Selection of appropriate adjuvant chemotherapy or hormone therapy is highly controversial in some situations. Meta-analyses have helped to define broad limits for therapy but do not help in choosing optimal regimens or in choosing a regimen for certain subgroups of patients. A summary of recommendations is shown in **Table 108-3**. In general, premenopausal women for whom any form of adjuvant systemic therapy is indicated should receive multidrug chemotherapy. Antihormone therapy improves survival in premenopausal patients who are estrogen receptor positive and should be added following completion of chemotherapy. Prophylactic surgical or medically induced castration may also be associated with a substantial survival benefit (primarily in estrogen receptor-positive patients) but is not widely used in this country.

Data on postmenopausal women are also controversial. The impact of adjuvant chemotherapy is quantitatively less clear-cut than in premenopausal patients, particularly in estrogen receptor-positive cases, although survival advantages have been shown. The first decision is whether chemotherapy or endocrine therapy should be used. While adjuvant endocrine therapy (aromatase inhibitors and tamoxifen) improves survival regardless of axillary lymph node status, the improvement in survival is modest for patients in whom multiple lymph nodes are involved. For this reason, it has been usual to give chemotherapy to postmenopausal patients who have no medical contraindications and who have more than one positive lymph node; hormone therapy is commonly given subsequently. For postmenopausal women for whom systemic therapy is warranted but who have a more favorable prognosis (based more commonly on analysis such as the Oncotype DX methodology), hormone therapy may be used alone. Large clinical trials have

TABLE 108-2 5-YEAR SURVIVAL RATE FOR BREAST CANCER BY STAGE

Stage	5-Year Survival, %
0	99
I	92
IIA	82
IIB	65
IIIA	47
IIIB	44
IV	14

Source: Modified from data of the National Cancer Institute: Surveillance, Epidemiology, and End Results (SEER).

TABLE 108-3 SUGGESTED APPROACHES TO ADJUVANT THERAPY

Age Group	Lymph Node Status ^a	Estrogen Receptor (ER) Status	Tumor	Recommendation
Premenopausal	Positive	Any	Any	Multidrug chemotherapy + tamoxifen if ER-positive + trastuzumab in HER2/neu-positive tumors
Premenopausal	Negative	Any	>2 cm, or 1–2 cm with other poor prognostic variables	Multidrug chemotherapy + tamoxifen if ER-positive + trastuzumab in HER2/neu-positive tumors. Consider Oncotype or similar testing.
Postmenopausal	Positive	Negative	Any	Multidrug chemotherapy + trastuzumab in HER2/neu-positive tumors
Postmenopausal	Positive	Positive	Any	Aromatase inhibitors and tamoxifen with or without chemotherapy + trastuzumab in HER2/neu-positive tumors
Postmenopausal	Negative	Positive	>2 cm, or 1–2 cm with other poor prognostic variables	Aromatase inhibitors and tamoxifen + trastuzumab in HER2/neu-positive tumors
Postmenopausal	Negative	Negative	>2 cm, or 1–2 cm with other poor prognostic variables	Consider multidrug chemotherapy + trastuzumab in HER2/neu-positive tumors

^aAs determined by pathologic examination.

shown superiority for aromatase inhibitors over tamoxifen alone in the adjuvant setting, although tamoxifen appears essentially equivalent in women who are obese and therefore presumably have higher endogenous concentrations of estrogen. Unfortunately the optimal plan is unclear. Tamoxifen for 5 years followed by an aromatase inhibitor, the reverse strategy, or even switching to an aromatase inhibitor after 2–3 years of tamoxifen has been shown to be better than tamoxifen alone. Continuation of tamoxifen for 10 years yields further benefit and is a reasonable decision for women with less favorable prognoses. Unfortunately, multiple studies have revealed very suboptimal adherence to long-term adjuvant endocrine regimens, and every effort should be made to encourage their continuous use. No valid information currently permits selection among the three clinically approved aromatase inhibitors. Concomitant use of bisphosphonates is almost always warranted; however, it is not finally settled as to whether their prophylactic use increases survival in addition to just decreasing recurrences in bone.

Most comparisons of adjuvant chemotherapy regimens show little difference among them, although small advantages for doxorubicin-containing regimens and “dose-dense” regimens are usually seen.

One approach—so-called neoadjuvant chemotherapy—involves the administration of adjuvant therapy before definitive surgery and radiation therapy. Because the objective response rates of patients with breast cancer to systemic therapy in this setting exceed 75%, many patients will be “downstaged” and may become candidates for breast-conserving therapy. However, overall survival has not been improved using this approach as compared with the same drugs given postoperatively. Patients who achieve a pathologic complete remission after neoadjuvant chemotherapy not unexpectedly have a substantially improved survival. The neoadjuvant setting also provides a wonderful opportunity for the evaluation of new agents. For example, a second HER2 targeting antibody, pertuzumab, has been shown to provide additional benefit when combined with trastuzumab in the neoadjuvant setting.

Other adjuvant treatments under investigation include the use of taxanes, such as paclitaxel and docetaxel, and therapy based on alternative kinetic and biologic models. In such approaches, high doses of single agents are used separately in relatively dose-intensive cycling regimens. Node-positive patients treated with doxorubicin-cyclophosphamide for four cycles followed by four cycles of a taxane have a substantial improvement in survival compared with women receiving doxorubicin-cyclophosphamide alone, particularly in women with estrogen receptor–negative tumors. In addition, administration of the same drug combinations at the same dose but at more frequent intervals (every 2 weeks with cytokine support as compared with the standard every 3 weeks) is even more effective. Among the 25% of women whose tumors overexpress HER2/neu, addition of trastuzumab given concurrently with a taxane and then for a year after chemotherapy produces significant improvement in survival. Although longer follow-up will be important, this is now

the standard care for most women with HER2/neu-positive breast cancers. Cardiotoxicity, immediate and long-term, remains a concern, and further efforts to exploit non-anthracycline-containing regimens are being pursued. Very-high-dose therapy with stem cell transplantation in the adjuvant setting has not proved superior to standard-dose therapy and should not be routinely used.

A variety of exciting approaches are close to adoption, and the literature needs to be followed attentively. Tyrosine kinase inhibitors such as lapatinib and additional HER2-targeting antibodies such as pertuzumab are very promising. Finally, as described in the next section, a novel class of agents targeting DNA repair—the so-called poly-ADP ribose polymerase (PARP) inhibitors—is likely to have a major effect on breast cancers either caused by *BRCA1* or *BRCA2* mutations or sharing similar defects in DNA repair in their etiology.

SYSTEMIC THERAPY OF METASTATIC DISEASE

About one-third of patients treated for apparently localized breast cancer develop metastatic disease. Although a small number of these patients enjoy long remissions when treated with combinations of systemic and local therapy, most eventually succumb to metastatic disease. The median survival for all patients diagnosed with metastatic breast cancer is less than 3 years. Soft tissue, bony, and visceral (lung and liver) metastases each account for approximately one-third of sites of initial relapses. However, by the time of death, most patients will have bony involvement. Recurrences can appear at any time after primary therapy. A very cruel fact about breast cancer recurrences is that at least half of all breast cancer recurrences occur >5 years after initial therapy. It is now clear that a variety of host factors can influence recurrence rates, including depression and central obesity, and these diseases should be managed as aggressively as possible.

Because the diagnosis of metastatic disease alters the outlook for the patient so drastically, it should rarely be made without a confirmatory biopsy. Every oncologist has seen patients with tuberculosis, gallstones, sarcoidosis, or other nonmalignant diseases misdiagnosed and treated as though they had metastatic breast cancer or even second malignancies such as multiple myeloma thought to be recurrent breast cancer. This is a catastrophic mistake and justifies biopsy for virtually every patient at the time of initial suspicion of metastatic disease. Furthermore, there are well-documented changes in hormone receptor status that can occur and substantially alter treatment decisions.

The choice of therapy requires consideration of local therapy needs, the overall medical condition of the patient, and the hormone receptor status of the tumor, as well as clinical judgment. Because therapy of systemic disease is palliative, the potential toxicities of therapies should be balanced against the response rates. Several variables influence the response to systemic therapy. For example, the presence of estrogen and progesterone receptors is a strong indication for endocrine therapy. On the other hand, patients

530 with short disease-free intervals, rapidly progressive visceral disease, lymphangitic pulmonary disease, or intracranial disease are unlikely to respond to endocrine therapy.

In many cases, systemic therapy can be withheld while the patient is managed with appropriate local therapy. Radiation therapy and occasionally surgery are effective at relieving the symptoms of metastatic disease, particularly when bony sites are involved. Many patients with bone-only or bone-dominant disease have a relatively indolent course. Under such circumstances, systemic chemotherapy has a modest effect, whereas radiation therapy may be effective for long periods. Other systemic treatments, such as strontium-89 and/or bisphosphonates, may provide a palliative benefit without inducing objective responses. Most patients with metastatic disease, and certainly all who have bone involvement, should receive concurrent bisphosphonates. Because the goal of therapy is to maintain well-being for as long as possible, emphasis should be placed on avoiding the most hazardous complications of metastatic disease, including pathologic fracture of the axial skeleton and spinal cord compression. New back pain in patients with cancer should be explored aggressively on an emergent basis; to wait for neurologic symptoms is a potentially catastrophic error. Metastatic involvement of endocrine organs can cause profound dysfunction, including adrenal insufficiency and hypopituitarism. Similarly, obstruction of the biliary tree or other impaired organ function may be better managed with a local therapy than with a systemic approach.

Many patients are inappropriately treated with toxic regimens into their last days of life. Often oncologists are unwilling to have the difficult conversations that are required with patients nearing the end of life, and not uncommonly, patients and families can pressure physicians into treatments with very little survival value. Palliative care consultation and realistic assessment of treatment expectations need to be reviewed with patients and families. We urge consideration of palliative care consultations for patients who have received at least two lines of therapy for metastatic disease.

Endocrine Therapy Normal breast tissue is estrogen dependent. Both primary and metastatic breast cancer may retain this phenotype. The best means of ascertaining whether a breast cancer is hormone dependent is through analysis of estrogen and progesterone receptor levels on the tumor. Tumors that are positive for the estrogen receptor and negative for the progesterone receptor have a response rate of ~30%. Tumors that are positive for both receptors have a response rate approaching 70%. If neither receptor is present, the objective response rates are <5%. Receptor analyses provide information as to the correct ordering of endocrine therapies as opposed to chemotherapy. Because of their lack of toxicity and because some patients whose receptor analyses are reported as negative respond to endocrine therapy, an endocrine treatment should be attempted in virtually every patient with metastatic breast cancer. Potential endocrine therapies are summarized in **Table 108-4**. The choice of endocrine therapy is usually determined by toxicity profile and availability. In most postmenopausal patients, the initial endocrine therapy should be an aromatase inhibitor rather than tamoxifen. For the subset of postmenopausal women who are estrogen receptor-positive but also HER2/neu-positive, response rates to aromatase inhibitors are substantially higher than to tamoxifen. Aromatase inhibitors are not used in premenopausal women because their hypothalamus can respond to estrogen deprivation by producing gonadotropins that promote estrogen synthesis. Newer “pure” antiestrogens that are free of agonistic effects are also effective. Cases in which tumors shrink in response to tamoxifen withdrawal (as well as withdrawal of pharmacologic doses of estrogens) have been reported. A series of studies with aromatase inhibitors, tamoxifen, and fulvestrant have all shown that the addition of everolimus to the hormonal treatment can lead to significant benefit after progression on the endocrine agent alone. Everolimus (an mTOR inhibitor) in coordination with endocrine agents is now being explored as front-line therapy and in the adjuvant setting. Endogenous estrogen formation may be blocked by analogues of

TABLE 108-4 ENDOCRINE THERAPIES FOR BREAST CANCER

Therapy	Comments
Castration	For premenopausal women
Surgical	
LHRH agonists	
Antiestrogens	
Tamoxifen	Useful in pre- and postmenopausal women ^a
“Pure” antiestrogens	Responses in tamoxifen-resistant and aromatase inhibitor-resistant patients ^a
Surgical adrenalectomy	Rarely used second-line choice
Aromatase inhibitors	Low toxicity; now first choice for metastatic disease ^a
High-dose progestogens	Common fourth-line choice after aromatase inhibitors, tamoxifen, and fulvestrant
Hypophysectomy	Rarely used
Additive androgens or estrogens	Plausible fourth-line therapies; potentially toxic

^aConsider retreatment with Everolimus in combination for disease progression

Abbreviation: LHRH, luteinizing hormone-releasing hormone.

luteinizing hormone-releasing hormone in premenopausal women. Additive endocrine therapies, including treatment with progestogens, estrogens, and androgens, may also be tried in patients who respond to initial endocrine therapy; the mechanism of action of these latter therapies is unknown. Patients who respond to one endocrine therapy have at least a 50% chance of responding to a second endocrine therapy. It is not uncommon for patients to respond to two or three sequential endocrine therapies; however, combination endocrine therapies do not appear to be superior to individual agents, and combinations of chemotherapy with endocrine therapy are not useful. The median survival of patients with metastatic disease is approximately 2 years, although many patients, particularly older persons and those with hormone-dependent disease, may respond to endocrine therapy for 3–5 years or longer.

Chemotherapy Unlike many other epithelial malignancies, breast cancer responds to multiple chemotherapeutic agents, including anthracyclines, alkylating agents, taxanes, and antimetabolites. Multiple combinations of these agents have been found to improve response rates somewhat, but they have had little effect on duration of response or survival. The choice among multidrug combinations frequently depends on whether adjuvant chemotherapy was administered and, if so, what type. Although patients treated with adjuvant regimens such as cyclophosphamide, methotrexate, and fluorouracil (CMF regimens) may subsequently respond to the same combination in the metastatic disease setting, most oncologists use drugs to which the patients have not been previously exposed. Once patients have progressed after combination drug therapy, it is most common to treat them with single agents. Given the significant toxicity of most drugs, the use of a single effective agent will minimize toxicity by sparing the patient exposure to drugs that would be of little value. No method to select the drugs most efficacious for a given patient has been demonstrated to be useful.

Most oncologists use either an anthracycline or paclitaxel following failure with the initial regimen. However, the choice has to be balanced with individual needs. One randomized study has suggested that docetaxel may be superior to paclitaxel. A nanoparticle formulation of paclitaxel (Abraxane) is also effective.

The use of a humanized antibody to *erbB2* (trastuzumab [Herceptin]) combined with paclitaxel can improve response rate and survival for women whose metastatic tumors overexpress *erbB2*. A novel antibody conjugate (ADC) that links trastuzumab to a cytotoxic agent has been approved for management of HER2-positive breast cancer. The magnitude of the survival extension is modest in patients with metastatic disease. Similarly, the use of bevacizumab

(Avastin) has improved the response rate and response duration to paclitaxel. Objective responses in previously treated patients may also be seen with gemcitabine, vinca alkaloids, capecitabine, vinorelbine, and oral etoposide, as well as a new class of agents, epothilones. There are few comparative trials of one agent versus another in metastatic disease. It is a sad fact that choices are often influenced by aggressive marketing of new very expensive agents that have not been shown to be superior to other generic agents. Platinum-based agents have become far more widely used in both the adjuvant and advanced disease settings for some breast cancers, particularly those of the “triple-negative” subtype.

HIGH-DOSE CHEMOTHERAPY INCLUDING AUTOLOGOUS BONE MARROW TRANSPLANTATION Autologous bone marrow transplantation combined with high doses of single agents can produce objective responses even in heavily pretreated patients. However, such responses are rarely durable and do not alter the clinical course for most patients with advanced metastatic disease.

STAGE III BREAST CANCER

Between 10 and 25% of patients present with so-called locally advanced, or stage III, breast cancer at diagnosis. Many of these cancers are technically operable, whereas others, particularly cancers with chest wall involvement, inflammatory breast cancers, or cancers with large matted axillary lymph nodes, cannot be managed with surgery initially. Although no randomized trials have shown any survival benefit for neoadjuvant regimens as compared to adjuvant therapy, this approach has gained widespread use. More than 90% of patients with locally advanced breast cancer show a partial or better response to multidrug chemotherapy regimens that include an anthracycline. Early administration of this treatment reduces the bulk of the disease and frequently makes the patient a suitable candidate for salvage surgery and/or radiation therapy. These patients should be managed in multimodality clinics to coordinate surgery, radiation therapy, and systemic chemotherapy. Such approaches produce long-term disease-free survival in about 30–50% of patients. The neoadjuvant setting is also an ideal time to evaluate the efficacy of novel treatments because the effect on the tumor can be directly assessed.

BREAST CANCER PREVENTION

Women who have one breast cancer are at risk of developing a contralateral breast cancer at a rate of approximately 0.5% per year. When adjuvant tamoxifen or an aromatase inhibitor is administered to these patients, the rate of development of contralateral breast cancers is reduced. In other tissues of the body, tamoxifen has estrogen-like effects that are beneficial, including preservation of bone mineral density and long-term lowering of cholesterol. However, tamoxifen has estrogen-like effects on the uterus, leading to an increased risk of uterine cancer (0.75% incidence after 5 years on tamoxifen). Tamoxifen also increases the risk of cataract formation. The Breast Cancer Prevention Trial (BCPT) revealed a >49% reduction in breast cancer among women with a risk of at least 1.66% taking the drug for 5 years. Raloxifene has shown similar breast cancer prevention potency but may have different effects on bone and heart. The two agents have been compared in a prospective randomized prevention trial (the Study of Tamoxifen and Raloxifene [STAR] trial). The agents are approximately equivalent in preventing breast cancer with fewer thromboembolic events and endometrial cancers with raloxifene; however, raloxifene did not reduce noninvasive cancers as effectively as tamoxifen, so no clear winner has emerged. A newer selective estrogen receptor modulator (SERM), lasofoxifene, has been shown to reduce cardiovascular events in addition to breast cancer and fractures, and further studies of this agent should be watched with interest. It should be recalled that prevention of contralateral breast cancers in women diagnosed with one cancer is a reasonable surrogate for breast cancer prevention because these are second primaries not recurrences. In this regard, the aromatase inhibitors are all considerably more effective than tamoxifen; however, they are not approved for primary breast

cancer prevention. It remains puzzling that agents with the safety profile of raloxifene, which can reduce breast cancer risk by 50% with additional benefits in preventing osteoporotic fracture, are still so infrequently prescribed. They should be far more commonly offered to women than they are.

NONINVASIVE BREAST CANCER

Breast cancer develops as a series of molecular changes in the epithelial cells that lead to ever more malignant behavior. Increased use of mammography has led to more frequent diagnoses of noninvasive breast cancer. These lesions fall into two groups: ductal carcinoma in situ (DCIS) and lobular carcinoma in situ (lobular neoplasia). The management of both entities is controversial.

Ductal Carcinoma In Situ Proliferation of cytologically malignant breast epithelial cells within the ducts is termed *DCIS*. Atypical hyperplasia may be difficult to differentiate from DCIS. At least one-third of patients with untreated DCIS develop invasive breast cancer within 5 years. However, many low-grade DCIS lesions do not appear to progress over many years; therefore, many patients are overtreated. Unfortunately there is no reliable means of distinguishing patients who require treatment from those who may be safely observed. For many years, the standard treatment for this disease was mastectomy. However, treatment of this condition by lumpectomy and radiation therapy gives survival that is as good as the survival for invasive breast cancer treated by mastectomy. In one randomized trial, the combination of wide excision plus irradiation for DCIS caused a substantial reduction in the local recurrence rate as compared with wide excision alone with negative margins, although survival was identical in the two arms. No studies have compared either of these regimens to mastectomy. Addition of tamoxifen to any DCIS surgical/radiation therapy regimen further improves local control. Data for aromatase inhibitors in this setting are not available.

Several prognostic features may help to identify patients at high risk for local recurrence after either lumpectomy alone or lumpectomy with radiation therapy. These include extensive disease; age <40; and cytologic features such as necrosis, poor nuclear grade, and comedo subtype with overexpression of *erbB2*. Some data suggest that adequate excision with careful determination of pathologically clear margins is associated with a low recurrence rate. When surgery is combined with radiation therapy, recurrence (which is usually in the same quadrant) occurs with a frequency of ≤10%. Given the fact that half of these recurrences will be invasive, about 5% of the initial cohort will eventually develop invasive breast cancer. A reasonable expectation of mortality for these patients is about 1%, a figure that approximates the mortality rate for DCIS managed by mastectomy. Although this train of reasoning has not formally been proved valid, it is reasonable to recommend that patients who desire breast preservation, and in whom DCIS appears to be reasonably localized, be managed by adequate surgery with meticulous pathologic evaluation, followed by breast irradiation and tamoxifen. For patients with localized DCIS, axillary lymph node dissection is unnecessary. More controversial is the question of what management is optimal when there is any degree of invasion. Because of a significant likelihood (10–15%) of axillary lymph node involvement even when the primary lesion shows only microscopic invasion, it is prudent to do at least a sentinel lymph node sampling for all patients with any degree of invasion. Further management is dictated by the presence of nodal spread.

Lobular Neoplasia Proliferation of cytologically malignant cells within the lobules is termed *lobular neoplasia*. Nearly 30% of patients who have had adequate local excision of the lesion develop breast cancer (usually infiltrating ductal carcinoma) over the next 15–20 years. Ipsilateral and contralateral cancers are equally common. Therefore, lobular neoplasia may be a premalignant lesion that suggests an elevated risk of subsequent breast cancer, rather than a form of malignancy itself, and aggressive local management seems unreasonable. Most patients should be treated with an SERM or an aromatase inhibitor (for postmenopausal women) for 5 years and


109 Upper Gastrointestinal Tract Cancers

Robert J. Mayer

Upper gastrointestinal cancers include malignancies arising in the esophagus, stomach, and small intestine.

ESOPHAGEAL CANCER

INCIDENCE AND ETIOLOGY

 Cancer of the esophagus is an increasingly common and extremely lethal malignancy. The diagnosis was made in 18,170 Americans in 2014 and led to 15,450 deaths. Almost all esophageal cancers are either squamous cell carcinomas or adenocarcinomas; the two histologic subtypes have a similar clinical presentation but different causative factors.

Worldwide, squamous cell carcinoma is the more common cell type, having an incidence that rises strikingly in association with geographic location. It occurs frequently within a region extending from the southern shore of the Caspian Sea on the west to northern China on the east, encompassing parts of Iran, central Asia, Afghanistan, Siberia, and Mongolia. Familial increased risk has been observed in regions with high incidence, although gene associations are not yet defined. High-incidence “pockets” of the disease are also present in such disparate locations as Finland, Iceland, Curaçao, southeastern Africa, and northwestern France. In North America and western Europe, the disease is more common in blacks than whites and in males than females; it appears most often after age 50 and seems to be associated with a lower socioeconomic status. Such cancers generally arise in the cervical and thoracic portions of the esophagus.

A variety of causative factors have been implicated in the development of squamous cell cancers of the esophagus (Table 109-1). In the United States, the etiology of such cancers is primarily related to excess alcohol consumption and/or cigarette smoking. The relative risk increases with the amount of tobacco smoked or alcohol consumed, with these factors acting synergistically. The consumption of whiskey is linked to a higher incidence than the consumption of wine or beer. Squamous cell esophageal carcinoma has also been associated with the ingestion of nitrates, smoked opiates, and fungal toxins in pickled vegetables, as well as mucosal damage caused by such physical insults as long-term exposure to extremely hot tea, the ingestion of lye, radiation-induced strictures, and chronic achalasia. The presence of an esophageal web in association with glossitis and iron deficiency (i.e., Plummer-Vinson or Paterson-Kelly syndrome) and congenital

TABLE 109-1 SOME ETIOLOGIC FACTORS ASSOCIATED WITH SQUAMOUS CELL CANCER OF THE ESOPHAGUS

Excess alcohol consumption
Cigarette smoking
Other ingested carcinogens
Nitrates (converted to nitrites)
Smoked opiates
Fungal toxins in pickled vegetables
Mucosal damage from physical agents
Hot tea
Lye ingestion
Radiation-induced strictures
Chronic achalasia
Host susceptibility
Esophageal web with glossitis and iron deficiency (i.e., Plummer-Vinson or Paterson-Kelly syndrome)
Congenital hyperkeratosis and pitting of the palms and soles (i.e., tylosis palmaris et plantaris)
? Dietary deficiencies of selenium, molybdenum, zinc, and vitamin A

532 followed with careful annual mammography and semiannual physical examinations. Additional molecular analysis of these lesions may make it possible to discriminate between patients who are at risk of further progression and require additional therapy and those in whom simple follow-up is adequate.

MALE BREAST CANCER

Breast cancer is about 1/150th as frequent in men as in women; 1720 men developed breast cancer in 2006. It usually presents as a unilateral lump in the breast and is frequently not diagnosed promptly. Given the small amount of soft tissue and the unexpected nature of the problem, locally advanced presentations are somewhat more common. When male breast cancer is matched to female breast cancer by age and stage, its overall prognosis is identical. Although gynecomastia may initially be unilateral or asymmetric, any unilateral mass in a man older than age 40 years should receive a careful workup including biopsy. On the other hand, bilateral symmetric breast development rarely represents breast cancer and is almost invariably due to endocrine disease or a drug effect. It should be kept in mind, nevertheless, that the risk of cancer is much greater in men with gynecomastia; in such men, gross asymmetry of the breasts should arouse suspicion of cancer. Male breast cancer is best managed by mastectomy and axillary lymph node dissection or SLNB. Patients with locally advanced disease or positive nodes should also be treated with irradiation. Approximately 90% of male breast cancers contain estrogen receptors, and approximately 60% of cases with metastatic disease respond to endocrine therapy. No randomized studies have evaluated adjuvant therapy for male breast cancer. Two historic experiences suggest that the disease responds well to adjuvant systemic therapy, and, if not medically contraindicated, the same criteria for the use of adjuvant therapy in women should be applied to men.

The sites of relapse and spectrum of response to chemotherapeutic drugs are virtually identical for breast cancers in either sex.

FOLLOW-UP OF BREAST CANCER PATIENTS

Despite the availability of sophisticated and expensive imaging techniques and a wide range of serum tumor marker tests, survival is not influenced by early diagnosis of relapse. Surveillance guidelines are given in Table 108-5. Despite pressure from patients and their families, routine computed tomography scans (or other imaging) are not recommended.

TABLE 108-5 BREAST CANCER SURVEILLANCE GUIDELINES

Test	Frequency
Recommended	
History; eliciting symptoms; physical examination	q3–6 months × 3 years; q6–12 months × 2 years; then annually
Breast self-examination	Monthly
Mammography	Annually
Pelvic examination	Annually (particularly for patients on SERMs)
Patient education about symptoms of recurrence	Ongoing
Coordination of care	Ongoing
Not Recommended	
Complete blood count	
Serum chemistry studies	
Chest radiographs	
Bone scans	
Ultrasound examination of the liver	
Computed tomography of chest, abdomen, or pelvis	
Tumor markers CA 15-3, CA 27-29, CEA	

Abbreviations: CEA, carcinoembryonic antigen; SERM, selective estrogen receptor modulator.

Source: Recommended Breast Cancer Surveillance Guidelines, ASCO Education Book, Fall, 1997.

TABLE 109-2 SOME ETIOLOGIC FACTORS ASSOCIATED WITH ADENOCARCINOMA OF THE ESOPHAGUS

Chronic gastroesophageal reflux
Obesity
Barrett's esophagus
Male sex
Cigarette smoking

hyperkeratosis and pitting of the palms and soles (i.e., tylosis palmaris et plantaris) have each been linked with squamous cell esophageal cancer, as have dietary deficiencies of molybdenum, zinc, selenium, and vitamin A. Patients with head and neck cancer are at increased risk of squamous cell cancer of the esophagus.

For unclear reasons, the incidence of squamous cell esophageal cancer has decreased somewhat in both the black and white populations in the United States over the past 40 years, whereas the rate of adenocarcinoma has risen sevenfold, particularly in white males (male-to-female ratio of 6:1). Whereas squamous cell cancers comprised the vast majority of esophageal cancers in the United States as recently as 40–50 years ago, more than 75% of esophageal tumors are now adenocarcinomas, with the incidence of this histologic subtype continuing to increase rapidly. Understanding the cause for this increase is the focus of current investigation.

Several strong etiologic associations have been observed to account for the development of adenocarcinoma of the esophagus (Table 109-2). Such tumors arise in the distal esophagus in association with chronic gastric reflux, often in the presence of Barrett's esophagus (replacement of the normal squamous epithelium of the distal esophagus by columnar mucosa), which occurs more commonly in obese individuals. Adenocarcinomas arise within dysplastic columnar epithelium in the distal esophagus. Even before frank neoplasia is detectable, aneuploidy and *p53* mutations are found in the dysplastic epithelium. These adenocarcinomas behave clinically like gastric adenocarcinomas, although they are not associated with *Helicobacter pylori* infections. Approximately 15% of esophageal adenocarcinomas overexpress the *HER2/neu* gene.

CLINICAL FEATURES

About 5% of esophageal cancers occur in the upper third of the esophagus (cervical esophagus), 20% in the middle third, and 75% in the lower third. Squamous cell carcinomas and adenocarcinomas cannot be distinguished radiographically or endoscopically.

Progressive dysphagia and weight loss of short duration are the initial symptoms in the vast majority of patients. Dysphagia initially occurs with solid foods and gradually progresses to include semisolids and liquids. By the time these symptoms develop, the disease is already very advanced, because difficulty in swallowing does not occur until >60% of the esophageal circumference is infiltrated with cancer. Dysphagia may be associated with pain on swallowing (odynophagia), pain radiating to the chest and/or back, regurgitation or vomiting, and aspiration pneumonia. The disease most commonly spreads to adjacent and supraclavicular lymph nodes, liver, lungs, pleura, and bone. Tracheoesophageal fistulas may develop, primarily in patients with upper and mid-esophageal tumors. As with other squamous cell carcinomas, hypercalcemia may occur in the absence of osseous metastases, probably from parathormone-related peptide secreted by tumor cells (Chap. 121).

DIAGNOSIS

Attempts at endoscopic and cytologic screening for carcinoma in patients with Barrett's esophagus, while effective as a means of detecting high-grade dysplasia, have not yet been shown to reduce the likelihood of death from esophageal adenocarcinoma. Esophagoscopy should be performed in all patients suspected of having an esophageal abnormality, to both visualize and identify a tumor and also to obtain histopathologic confirmation of the diagnosis. Because the population of persons at risk for squamous cell carcinoma of the esophagus (i.e., smokers and drinkers) also has a high rate of cancers of the lung and

the head and neck region, endoscopic inspection of the larynx, trachea, and bronchi should also be carried out. A thorough examination of the fundus of the stomach (by retroflexing the endoscope) is imperative as well. The extent of tumor spread to the mediastinum and para-aortic lymph nodes should be assessed by computed tomography (CT) scans of the chest and abdomen and by endoscopic ultrasound. Positron emission tomography scanning provides a useful assessment of the presence of distant metastatic disease, offering accurate information regarding spread to mediastinal lymph nodes, which can be helpful in defining radiation therapy fields. Such scans, when performed sequentially, appear to provide a means of making an early assessment of responsiveness to preoperative chemotherapy.

TREATMENT ESOPHAGEAL CANCER

The prognosis for patients with esophageal carcinoma is poor. Approximately 10% of patients survive 5 years after the diagnosis; thus, management focuses on symptom control. Surgical resection of all gross tumor (i.e., total resection) is feasible in only 45% of cases, with residual tumor cells frequently present at the resection margins. Such esophagectomies have been associated with a postoperative mortality rate of approximately 5% due to anastomotic fistulas, subphrenic abscesses, and cardiopulmonary complications. Although debate regarding the comparative benefits of trans-thoracic versus transhiatal resections has continued, experienced thoracic surgeons are now favoring minimally invasive transthoracic esophagectomies. Endoscopic resections of superficial squamous cell cancers or adenocarcinomas are being examined but have not yet been shown to result in a similar likelihood of survival as observed with conventional surgical procedures. Similarly, the value of endoscopic ablation of dysplastic lesions in an area of Barrett's esophagus on reducing subsequent mortality from esophageal carcinoma is uncertain. Some experts have advocated fundoplication surgery (i.e., the removal of the gastroesophageal junction) as a means of cancer prevention in patients with Barrett's esophagus; again, objective data are not yet available to fully assess the risks versus benefits of this invasive procedure. About 20% of patients who survive a total surgical resection live for 5 years. The evaluation of chemotherapeutic agents in patients with esophageal carcinoma has been hampered by ambiguity in the definition of "response" and the debilitated physical condition of many treated individuals, particularly those with squamous cell cancers. Nonetheless, significant reductions in the size of measurable tumor masses have been reported in 15–25% of patients given single-agent treatment and in 30–60% of patients treated with drug combinations that include cisplatin. In the small subset of patients whose tumors overexpress the *HER2/neu* gene, the addition of the monoclonal antibody trastuzumab (Herceptin) appears to further enhance the likelihood of benefit, particularly in patients with gastroesophageal lesions. The use of the antiangiogenic agent bevacizumab (Avastin) seems to be of limited value in the setting of esophageal cancer. Combination chemotherapy and radiation therapy as the initial therapeutic approach, either alone or followed by an attempt at operative resection, seems to be beneficial. When administered along with radiation therapy, chemotherapy produces a better survival outcome than radiation therapy alone. The use of preoperative chemotherapy and radiation therapy followed by esophageal resection appears to prolong survival compared with surgery alone according to several randomized trials and a meta-analysis; some reports suggest that no additional benefit accrues when surgery is added if significant shrinkage of tumor has been achieved by the chemoradiation combination.

For the incurable, surgically unresectable patient with esophageal cancer, dysphagia, malnutrition, and the management of tracheoesophageal fistulas are major issues. Approaches to palliation include repeated endoscopic dilatation, the surgical placement of a gastrostomy or jejunostomy for hydration and feeding, endoscopic placement of an expansive metal stent to bypass the tumor, and radiation therapy.

GASTRIC ADENOCARCINOMA



Incidence and Epidemiology For unclear reasons, the incidence and mortality rates for gastric cancer have decreased in the United States during the past 80 years, although the disease remains the second most frequent cause of worldwide cancer-related death. The mortality rate from gastric cancer in the United States has dropped in men from 28 to 5.8 per 100,000 persons, whereas in women, the rate has decreased from 27 to 2.8 per 100,000. Nonetheless, in 2014, 22,220 new cases of stomach cancer were diagnosed in the United States, and 10,990 Americans died of the disease. Although the incidence of gastric cancer has decreased worldwide, it remains high in such disparate geographic regions as Japan, China, Chile, and Ireland.

The risk of gastric cancer is greater among lower socioeconomic classes. Migrants from high- to low-incidence nations maintain their susceptibility to gastric cancer, whereas the risk for their offspring approximates that of the new homeland. These findings suggest that an environmental exposure, probably beginning early in life, is related to the development of gastric cancer, with dietary carcinogens considered the most likely factor(s).

Pathology About 85% of stomach cancers are adenocarcinomas, with 15% due to lymphomas, gastrointestinal stromal tumors (GISTs), and leiomyosarcomas. Gastric adenocarcinomas may be subdivided into two categories: a *diffuse type*, in which cell cohesion is absent, so that individual cells infiltrate and thicken the stomach wall without forming a discrete mass; and an *intestinal type*, characterized by cohesive neoplastic cells that form glandlike tubular structures. The diffuse carcinomas occur more often in younger patients, develop throughout the stomach (including the cardia), result in a loss of distensibility of the gastric wall (so-called *limitis plastica*, or “leather bottle” appearance), and carry a poorer prognosis. Diffuse cancers have defective intercellular adhesion, mainly as a consequence of loss of expression of E-cadherin. Intestinal-type lesions are frequently ulcerative, more commonly appear in the antrum and lesser curvature of the stomach, and are often preceded by a prolonged precancerous process, often initiated by *H. pylori* infection. Although the incidence of diffuse carcinomas is similar in most populations, the intestinal type tends to predominate in the high-risk geographic regions and is less likely to be found in areas where the frequency of gastric cancer is declining. Thus, different etiologic factor(s) are likely involved in these two subtypes. In the United States, ~30% of gastric cancers originate in the distal stomach, ~20% arise in the midportion of the stomach, and ~40% originate in the proximal third of the stomach. The remaining 10% involve the entire stomach.

Etiology The long-term ingestion of high concentrations of nitrates found in dried, smoked, and salted foods appears to be associated with a higher risk. The nitrates are thought to be converted to carcinogenic nitrites by bacteria (Table 109-3). Such bacteria may be introduced exogenously through the ingestion of partially decayed

foods, which are consumed in abundance worldwide by the lower socioeconomic classes. Bacteria such as *H. pylori* may also contribute to this effect by causing chronic inflammatory atrophic gastritis, loss of gastric acidity, and bacterial growth in the stomach. Although the risk for developing gastric cancer is thought to be sixfold higher in people infected with *H. pylori*, it remains uncertain whether eradicating the bacteria after infection has already occurred actually reduces this risk. Loss of acidity may occur when acid-producing cells of the gastric antrum have been removed surgically to control benign peptic ulcer disease or when achlorhydria, atrophic gastritis, and even pernicious anemia develop in the elderly. Serial endoscopic examinations of the stomach in patients with atrophic gastritis have documented replacement of the usual gastric mucosa by intestinal-type cells. This process of intestinal metaplasia may lead to cellular atypia and eventual neoplasia. Because the declining incidence of gastric cancer in the United States primarily reflects a decline in distal, ulcerating, intestinal-type lesions, it is conceivable that better food preservation and the availability of refrigeration for all socioeconomic classes have decreased the dietary ingestion of exogenous bacteria. *H. pylori* has not been associated with the diffuse, more proximal form of gastric carcinoma or with cancers arising at the gastroesophageal junction or in the distal esophagus. Approximately 10–15% of adenocarcinomas appearing in the proximal stomach, the gastroesophageal junction, and the distal esophagus overexpress the *HER2/neu* gene; individuals whose tumors demonstrate this overexpression benefit from treatment directed against this target (i.e., trastuzumab [Herceptin]).

Several additional etiologic factors have been associated with gastric carcinoma. Gastric ulcers and adenomatous polyps have occasionally been linked, but data on a cause-and-effect relationship are unconvincing. The inadequate clinical distinction between benign gastric ulcers and small ulcerating carcinomas may, in part, account for this presumed association. The presence of extreme hypertrophy of gastric rugal folds (i.e., Ménétrier’s disease), giving the impression of polypoid lesions, has been associated with a striking frequency of malignant transformation; such hypertrophy, however, does not represent the presence of true adenomatous polyps. Individuals with blood group A have a higher incidence of gastric cancer than persons with blood group O; this observation may be related to differences in the mucous secretion, leading to altered mucosal protection from carcinogens. A germline mutation in the E-cadherin gene (*CDH1*), inherited in an autosomal dominant pattern and coding for a cell adhesion protein, has been linked to a high incidence of occult diffuse-type gastric cancers in young asymptomatic carriers. Duodenal ulcers are not associated with gastric cancer.

Clinical Features Gastric cancers, when superficial and surgically curable, usually produce no symptoms. As the tumor becomes more extensive, patients may complain of an insidious upper abdominal discomfort varying in intensity from a vague, postprandial fullness to a severe, steady pain. Anorexia, often with slight nausea, is very common but is not the usual presenting complaint. Weight loss may eventually be observed, and nausea and vomiting are particularly prominent in patients whose tumors involve the pylorus; dysphagia and early satiety may be the major symptoms caused by diffuse lesions originating in the cardia. There may be no early physical signs. A palpable abdominal mass indicates long-standing growth and predicts regional extension.

Gastric carcinomas spread by direct extension through the gastric wall to the perigastric tissues, occasionally adhering to adjacent organs such as the pancreas, colon, or liver. The disease also spreads via lymphatics or by seeding of peritoneal surfaces. Metastases to intraabdominal and supraclavicular lymph nodes occur frequently, as do metastatic nodules to the ovary (Krukenberg’s tumor), periumbilical region (“Sister Mary Joseph node”), or peritoneal cul-de-sac (Blumer’s shelf palpable on rectal or vaginal examination); malignant ascites may also develop. The liver is the most common site for hematogenous spread of tumor.

TABLE 109-3 NITRATE-CONVERTING BACTERIA AS A FACTOR IN THE CAUSATION OF GASTRIC CARCINOMA^a

Exogenous sources of nitrate-converting bacteria:

Bacterially contaminated food (common in lower socioeconomic classes, who have a higher incidence of the disease; diminished by improved food preservation and refrigeration)

Helicobacter pylori infection

Endogenous factors favoring growth of nitrate-converting bacteria in the stomach:

Decreased gastric acidity

Prior gastric surgery (antrectomy) (15- to 20-year latency period)

Atrophic gastritis and/or pernicious anemia

? Prolonged exposure to histamine H₂-receptor antagonists

^aHypothesis: Dietary nitrates are converted to carcinogenic nitrites by bacteria.

The presence of iron-deficiency anemia in men and of occult blood in the stool in both sexes mandates a search for an occult gastrointestinal tract lesion. A careful assessment is of particular importance in patients with atrophic gastritis or pernicious anemia. Unusual clinical features associated with gastric adenocarcinomas include migratory thrombophlebitis, microangiopathic hemolytic anemia, diffuse seborrheic keratoses (so-called Leser-Trélat sign), and acanthosis nigricans.

Diagnosis The use of double-contrast radiographic examinations has been supplanted by esophagogastrosopy and CT scanning for the evaluation of patients with epigastric complaints.

Gastric ulcers identified at the time of such endoscopic procedure may appear benign but merit biopsy in order to exclude a malignancy. Malignant gastric ulcers must be recognized before they penetrate into surrounding tissues, because the rate of cure of early lesions limited to the mucosa or submucosa is >80%. Because gastric carcinomas are difficult to distinguish clinically or endoscopically from gastric lymphomas, endoscopic biopsies should be made as deeply as possible, due to the submucosal location of lymphoid tumors.

The staging system for gastric carcinoma is shown in [Table 109-4](#).

Stage	TNM	Features	Data from ACS in the United States	
			No. of Cases, %	5-Year Survival, %
0	T _b N0M0	Node negative; limited to mucosa	1	90
IA	T1N0M0	Node negative; invasion of lamina propria or submucosa	7	59
IB	T2N0M0 T1N1M0	Node negative; invasion of muscularis propria	10	44
II	T1N2M0 T2N1M0 T3N0M0	Node positive; invasion beyond mucosa but within wall or Node negative; extension through wall	17	29
IIIA	T2N2M0 T3N1-2M0	Node positive; invasion of muscularis propria or through wall	21	15
IIIB	T4N0-1M0	Node negative; adherence to surrounding tissue	14	9
IIIC	T4N2-3M0 T3N3M0	>3 nodes positive; invasion of serosa or adjacent structures 7 or more positive nodes; penetrates wall without invading serosa or adjacent structures		
IV	T4N2M0 T1-4N0-2-M1	Node positive; adherence to surrounding tissue or Distant metastases	30	3

Abbreviations: ACS, American Cancer Society; TNM, tumor, node, metastasis.

Complete surgical removal of the tumor with resection of adjacent lymph nodes offers the only chance for cure. However, this is possible in less than a third of patients. A subtotal gastrectomy is the treatment of choice for patients with distal carcinomas, whereas total or near-total gastrectomies are required for more proximal tumors. The inclusion of extended lymph node dissection in these procedures appears to confer an added risk for complications without providing a meaningful enhancement in survival. The prognosis following complete surgical resection depends on the degree of tumor penetration into the stomach wall and is adversely influenced by regional lymph node involvement and vascular invasion, characteristics found in the vast majority of American patients. As a result, the probability of survival after 5 years for the 25–30% of patients able to undergo complete resection is ~20% for distal tumors and <10% for proximal tumors, with recurrences continuing for at least 8 years after surgery. In the absence of ascites or extensive hepatic or peritoneal metastases, even patients whose disease is believed to be incurable by surgery should be offered resection of the primary lesion. Reduction of tumor bulk is the best form of palliation and may enhance the probability of benefit from subsequent therapy. In high-incidence regions such as Japan and Korea, where the use of endoscopic screening programs has identified patients with superficial tumors, the use of laparoscopic gastrectomy has gained popularity. In the United States and western Europe, the use of this less invasive surgical approach remains investigational.

Gastric adenocarcinoma is a relatively radioresistant tumor, and the adequate control of the primary tumor requires doses of external-beam irradiation that exceed the tolerance of surrounding structures, such as bowel mucosa and spinal cord. As a result, the major role of radiation therapy in patients has been palliation of pain. Radiation therapy alone after a complete resection does not prolong survival. In the setting of surgically unresectable disease limited to the epigastrium, patients treated with 3500–4000 cGy did not live longer than similar patients not receiving radiotherapy; however, survival was prolonged slightly when 5-fluorouracil (5-FU) plus leucovorin was given in combination with radiation therapy (3-year survival 50% vs 41% for radiation therapy alone). In this clinical setting, the 5-FU likely functions as a radiosensitizer.

The administration of combinations of cytotoxic drugs to patients with advanced gastric carcinoma has been associated with partial responses in 30–50% of cases; responders appear to benefit from treatment. Such drug combinations have generally included cisplatin combined with epirubicin or docetaxel and infusional 5-FU or capecitabine, or with irinotecan. Despite the encouraging response rates, complete remissions are uncommon, the partial responses are transient, and the overall impact of multidrug therapy on survival has been limited; the median survival time for patients treated in this manner remains less than 12 months. As with adenocarcinomas arising in the esophagus, the addition of bevacizumab (Avastin) to chemotherapy regimens in treating gastric cancer appears to provide limited benefit. However, preliminary results utilizing another antiangiogenic compound—ramucirumab (Cyranza)—in the treatment of gastric cancer are encouraging. The use of adjuvant chemotherapy alone following the complete resection of a gastric cancer has only minimally improved survival. However, combination chemotherapy administered before and after surgery (*perioperative treatment*) as well as postoperative chemotherapy combined with radiation therapy reduces the recurrence rate and prolongs survival.

PRIMARY GASTRIC LYMPHOMA

Primary lymphoma of the stomach is relatively uncommon, accounting for <15% of gastric malignancies and ~2% of all lymphomas. The stomach is, however, the most frequent extranodal site for lymphoma, and gastric lymphoma has increased in frequency during the past

536 35 years. The disease is difficult to distinguish clinically from gastric adenocarcinoma; both tumors are most often detected during the sixth decade of life; present with epigastric pain, early satiety, and generalized fatigue; and are usually characterized by ulcerations with a ragged, thickened mucosal pattern demonstrated by contrast radiographs or endoscopic appearance. The diagnosis of lymphoma of the stomach may occasionally be made through cytologic brushings of the gastric mucosa but usually requires a biopsy at gastroscopy or laparotomy. Failure of gastroscopic biopsies to detect lymphoma in a given case should not be interpreted as being conclusive, because superficial biopsies may miss the deeper lymphoid infiltrate. The macroscopic pathology of gastric lymphoma may also mimic adenocarcinoma, consisting of either a bulky ulcerated lesion localized in the corpus or antrum or a diffuse process spreading throughout the entire gastric submucosa and even extending into the duodenum. Microscopically, the vast majority of gastric lymphoid tumors are lymphomas of B-cell origin. Histologically, these tumors may range from well-differentiated, superficial processes (mucosa-associated lymphoid tissue [MALT]) to high-grade, large-cell lymphomas. Like gastric adenocarcinoma, infection with *H. pylori* increases the risk for gastric lymphoma in general and MALT lymphomas in particular. Large-cell lymphomas of the stomach spread initially to regional lymph nodes (often to Waldeyer's ring) and may then disseminate.

TREATMENT PRIMARY GASTRIC LYMPHOMA

Primary gastric lymphoma is a far more treatable disease than adenocarcinoma of the stomach, a fact that underscores the need for making the correct diagnosis. Antibiotic treatment to eradicate *H. pylori* infection has led to regression of about 75% of gastric MALT lymphomas and should be considered before surgery, radiation therapy, or chemotherapy is undertaken in patients having such tumors. A lack of response to such antimicrobial treatment has been linked to a specific chromosomal abnormality, i.e., t(11;18). Responding patients should undergo periodic endoscopic surveillance because it remains unclear whether the neoplastic clone is eliminated or merely suppressed, although the response to antimicrobial treatment is quite durable. Subtotal gastrectomy, usually followed by combination chemotherapy, has led to 5-year survival rates of 40–60% in patients with localized high-grade lymphomas. The need for a major surgical procedure has been questioned, particularly in patients with preoperative radiographic evidence of nodal involvement, for whom chemotherapy (CHOP [cyclophosphamide, doxorubicin, vincristine, and prednisone]) plus rituximab is highly effective therapy. A role for radiation therapy is not defined because most recurrences develop at distant sites.

GASTRIC (NONLYMPHOID) SARCOMA

Leiomyosarcomas and GISTs make up 1–3% of gastric neoplasms. They most frequently involve the anterior and posterior walls of the gastric fundus and often ulcerate and bleed. Even those lesions that appear benign on histologic examination may behave in a malignant fashion. These tumors rarely invade adjacent viscera and characteristically do not metastasize to lymph nodes, but they may spread to the liver and lungs. The treatment of choice is surgical resection. Combination chemotherapy should be reserved for patients with metastatic disease. All such tumors should be analyzed for a mutation in the *c-kit* receptor. GISTs are unresponsive to conventional chemotherapy; yet ~50% of patients experience objective response and prolonged survival when treated with imatinib mesylate (Gleevec) (400–800 mg PO daily), a selective inhibitor of the *c-kit* tyrosine kinase. Many patients with GIST whose tumors have become refractory to imatinib subsequently benefit from sunitinib (Sutent) or regorafenib (Stivarga), other inhibitors of the *c-kit* tyrosine kinase.

TUMORS OF THE SMALL INTESTINE

Small-bowel tumors comprise <3% of gastrointestinal neoplasms. Because of their rarity and inaccessibility, a correct diagnosis is often

delayed. Abdominal symptoms are usually vague and poorly defined, and conventional radiographic studies of the upper and lower intestinal tract often appear normal. Small-bowel tumors should be considered in the differential diagnosis in the following situations: (1) recurrent, unexplained episodes of crampy abdominal pain; (2) intermittent bouts of intestinal obstruction, especially in the absence of inflammatory bowel disease (IBD) or prior abdominal surgery; (3) intussusception in the adult; and (4) evidence of chronic intestinal bleeding in the presence of negative conventional and endoscopic examination. A careful small-bowel barium study should be considered in such a circumstance; the diagnostic accuracy may be improved by infusing barium through a nasogastric tube placed into the duodenum (enteroclysis). Alternatively, capsule endoscopic procedures have been used.

BENIGN TUMORS

The histology of benign small-bowel tumors is difficult to predict on clinical and radiologic grounds alone. The symptomatology of benign tumors is not distinctive, with pain, obstruction, and hemorrhage being the most frequent symptoms. These tumors are usually discovered during the fifth and sixth decades of life, more often in the distal rather than the proximal small intestine. The most common benign tumors are adenomas, leiomyomas, lipomas, and angiomas.

Adenomas These tumors include those of the islet cells and Brunner's glands as well as polypoid adenomas. *Islet cell adenomas* are occasionally located outside the pancreas; the associated syndromes are discussed in [Chap. 113](#). *Brunner's gland adenomas* are not truly neoplastic but represent a hypertrophy or hyperplasia of submucosal duodenal glands. These appear as small nodules in the duodenal mucosa that secrete a highly viscous alkaline mucus. Most often, this is an incidental radiographic finding not associated with any specific clinical disorder.

Polypoid Adenomas About 25% of benign small-bowel tumors are polypoid adenomas (see [Table 110-2](#)). They may present as single polypoid lesions or, less commonly, as papillary villous adenomas. As in the colon, the sessile or papillary form of the tumor is sometimes associated with a coexisting carcinoma. Occasionally, patients with Gardner's syndrome develop premalignant adenomas in the small bowel; such lesions are generally in the duodenum. Multiple polypoid tumors may occur throughout the small bowel (and occasionally the stomach and colorectum) in the Peutz-Jeghers syndrome. The polyps are usually hamartomas (juvenile polyps) having a low potential for malignant degeneration. Mucocutaneous melanin deposits as well as tumors of the ovary, breast, pancreas, and endometrium are also associated with this autosomal dominant condition.

Leiomyomas These neoplasms arise from smooth-muscle components of the intestine and are usually intramural, affecting the overlying mucosa. Ulceration of the mucosa may cause gastrointestinal hemorrhage of varying severity. Cramping or intermittent abdominal pain is frequently encountered.

Lipomas These tumors occur with greatest frequency in the distal ileum and at the ileocecal valve. They have a characteristic radiolucent appearance and are usually intramural and asymptomatic, but on occasion cause bleeding.

Angiomas While not true neoplasms, these lesions are important because they frequently cause intestinal bleeding. They may take the form of telangiectasia or hemangiomas. Multiple intestinal telangiectasias occur in a nonhereditary form confined to the gastrointestinal tract or as part of the hereditary Osler-Rendu-Weber syndrome. Vascular tumors may also take the form of isolated hemangiomas, most commonly in the jejunum. Angiography, especially during bleeding, is the best procedure for evaluating these lesions.

MALIGNANT TUMORS

While rare, small-bowel malignancies occur in patients with longstanding regional enteritis and celiac sprue as well as in individuals with AIDS. Malignant tumors of the small bowel are frequently associated with fever, weight loss, anorexia, bleeding, and a palpable

abdominal mass. After ampullary carcinomas (many of which arise from biliary or pancreatic ducts), the most frequently occurring small-bowel malignancies are adenocarcinomas, lymphomas, carcinoid tumors, and leiomyosarcomas.

ADENOCARCINOMAS

The most common primary cancers of the small bowel are adenocarcinomas, accounting for ~50% of malignant tumors. These cancers occur most often in the distal duodenum and proximal jejunum, where they tend to ulcerate and cause hemorrhage or obstruction. Radiologically, they may be confused with chronic duodenal ulcer disease or with Crohn's disease if the patient has long-standing regional enteritis. The diagnosis is best made by endoscopy and biopsy under direct vision. Surgical resection is the treatment of choice with suggested postoperative adjuvant chemotherapy options generally following treatment patterns used in the management of colon cancer.

LYMPHOMAS

Lymphoma in the small bowel may be primary or secondary. A diagnosis of a primary intestinal lymphoma requires histologic confirmation in a clinical setting in which palpable adenopathy and hepatosplenomegaly are absent and no evidence of lymphoma is seen on chest radiograph, CT scan, or peripheral blood smear or on bone marrow aspiration and biopsy. Symptoms referable to the small bowel are present, usually accompanied by an anatomically discernible lesion. Secondary lymphoma of the small bowel consists of involvement of the intestine by a lymphoid malignancy extending from involved retroperitoneal or mesenteric lymph nodes (**Chap. 134**).

Primary intestinal lymphoma accounts for ~20% of malignancies of the small bowel. These neoplasms are non-Hodgkin's lymphomas; they usually have a diffuse, large-cell histology and are of T cell origin. Intestinal lymphoma involves the ileum, jejunum, and duodenum, in decreasing frequency—a pattern that mirrors the relative amount of normal lymphoid cells in these anatomic areas. The risk of small-bowel lymphoma is increased in patients with a prior history of malabsorptive conditions (e.g., celiac sprue), regional enteritis, and depressed immune function due to congenital immunodeficiency syndromes, prior organ transplantation, autoimmune disorders, or AIDS.

The development of localized or nodular masses that narrow the lumen results in periumbilical pain (made worse by eating) as well as weight loss, vomiting, and occasional intestinal obstruction. The diagnosis of small-bowel lymphoma may be suspected from the appearance on contrast radiographs of patterns such as infiltration and thickening of mucosal folds, mucosal nodules, areas of irregular ulceration, or stasis of contrast material. The diagnosis can be confirmed by surgical exploration and resection of involved segments. Intestinal lymphoma can occasionally be diagnosed by peroral intestinal mucosal biopsy, but because the disease mainly involves the lamina propria, full-thickness surgical biopsies are usually required.

Resection of the tumor constitutes the initial treatment modality. While postoperative radiation therapy has been given to some patients following a total resection, most authorities favor short-term (three cycles) systemic treatment with combination chemotherapy. The frequent presence of widespread intraabdominal disease at the time of diagnosis and the occasional multicentricity of the tumor often make a total resection impossible. The probability of sustained remission or cure is ~75% in patients with localized disease but is ~25% in individuals with unresectable lymphoma. In patients whose tumors are not resected, chemotherapy may lead to bowel perforation.

A unique form of small-bowel lymphoma, diffusely involving the entire intestine, was first described in oriental Jews and Arabs and is referred to as *immunoproliferative small intestinal disease* (IPSID), *Mediterranean lymphoma*, or *α heavy chain disease*. This is a B cell tumor. The typical presentation includes chronic diarrhea and steatorrhea associated with vomiting and abdominal cramps; clubbing of the digits may be observed. A curious feature in many patients with IPSID is the presence in the blood and intestinal secretions of an abnormal IgA that contains a shortened α heavy chain and is devoid of light chains. It is suspected that the abnormal α chains are produced

by plasma cells infiltrating the small bowel. The clinical course of patients with IPSID is generally one of exacerbations and remissions, with death frequently resulting from either progressive malnutrition and wasting or the development of an aggressive lymphoma. The use of oral antibiotics such as tetracycline appears to be beneficial in the early phases of the disorder, suggesting a possible infectious etiology. Combination chemotherapy has been administered during later stages of the disease, with variable results. Results are better when antibiotics and chemotherapy are combined.

CARCINOID TUMORS

Carcinoid tumors arise from argentaffin cells of the crypts of Lieberkühn and are found from the distal duodenum to the ascending colon, areas embryologically derived from the midgut. More than 50% of intestinal carcinoids are found in the distal ileum, with most congregating close to the ileocecal valve. Most intestinal carcinoids are asymptomatic and of low malignant potential, but invasion and metastases may occur, leading to the carcinoid syndrome (**Chap. 113**).

LEIOMYOSARCOMAS

Leiomyosarcomas often are >5 cm in diameter and may be palpable on abdominal examination. Bleeding, obstruction, and perforation are common. Such tumors should be analyzed for the expression of mutant *c-kit* receptor (defining GIST), and in the presence of metastatic disease, justifying treatment with imatinib mesylate (Gleevec) or, in imatinib-refractory patients, sunitinib (Sutent) or regorafenib (Stivarga).


110 Lower Gastrointestinal Cancers

Robert J. Mayer

Lower gastrointestinal cancers include malignant tumors of the colon, rectum, and anus.

COLORECTAL CANCER

INCIDENCE

 Cancer of the large bowel is second only to lung cancer as a cause of cancer death in the United States: 136,830 new cases occurred in 2014, and 50,310 deaths were due to colorectal cancer. The incidence rate has decreased significantly during the past 25 years, likely due to enhanced and more compliantly followed screening practices. Similarly, mortality rates in the United States have decreased by approximately 25%, resulting largely from earlier detection and improved treatment.

POLYPS AND MOLECULAR PATHOGENESIS

Most colorectal cancers, regardless of etiology, arise from adenomatous polyps. A polyp is a grossly visible protrusion from the mucosal surface and may be classified pathologically as a nonneoplastic hamartoma (e.g., *juvenile polyp*), a hyperplastic mucosal proliferation (*hyperplastic polyp*), or an adenomatous polyp. Only adenomas are clearly premalignant, and only a minority of adenomatous polyps evolve into cancer. Adenomatous polyps may be found in the colons of ~30% of middle-aged and ~50% of elderly people; however, <1% of polyps ever become malignant. Most polyps produce no symptoms and remain clinically undetected. Occult blood in the stool is found in <5% of patients with polyps.

A number of molecular changes are noted in adenomatous polyps and colorectal cancers that are thought to reflect a multistep process in the evolution of normal colonic mucosa to life-threatening invasive carcinoma. These developmental steps toward carcinogenesis include, but are not restricted to, point mutations in the *K-ras* protooncogene; hypomethylation of DNA, leading to gene activation; loss of DNA

538 (*allelic loss*) at the site of a tumor-suppressor gene (the adenomatous polyposis coli [APC] gene) on the long arm of chromosome 5 (5q21); allelic loss at the site of a tumor-suppressor gene located on chromosome 18q (the deleted in colorectal cancer [DCC] gene); and allelic loss at chromosome 17p, associated with mutations in the *p53* tumor-suppressor gene (see Fig. 101e-2). Thus, the altered proliferative pattern of the colonic mucosa, which results in progression to a polyp and then to carcinoma, may involve the mutational activation of an oncogene followed by and coupled with the loss of genes that normally suppress tumorigenesis. It remains uncertain whether the genetic aberrations always occur in a defined order. Based on this model, however, cancer is believed to develop only in those polyps in which most (if not all) of these mutational events take place.

Clinically, the probability of an adenomatous polyp becoming a cancer depends on the gross appearance of the lesion, its histologic features, and its size. Adenomatous polyps may be pedunculated (stalked) or sessile (flat-based). Invasive cancers develop more frequently in sessile polyps. Histologically, adenomatous polyps may be tubular, villous (i.e., papillary), or tubulovillous. Villous adenomas, most of which are sessile, become malignant more than three times as often as tubular adenomas. The likelihood that any polypoid lesion in the large bowel contains invasive cancer is related to the size of the polyp, being negligible (<2%) in lesions <1.5 cm, intermediate (2–10%) in lesions 1.5–2.5 cm, and substantial (10%) in lesions >2.5 cm in size.

Following the detection of an adenomatous polyp, the entire large bowel should be visualized endoscopically because synchronous lesions are noted in about one-third of cases. Colonoscopy should then be repeated periodically, even in the absence of a previously documented malignancy, because such patients have a 30–50% probability of developing another adenoma and are at a higher-than-average risk for developing a colorectal carcinoma. Adenomatous polyps are thought to require >5 years of growth before becoming clinically significant; colonoscopy need not be carried out more frequently than every 3 years for the vast majority of patients.

ETIOLOGY AND RISK FACTORS

 Risk factors for the development of colorectal cancer are listed in Table 110-1.

Diet The etiology for most cases of large-bowel cancer appears to be related to environmental factors. The disease occurs more often in upper socioeconomic populations who live in urban areas. Mortality from colorectal cancer is directly correlated with per capita consumption of calories, meat protein, and dietary fat and oil as well as elevations in the serum cholesterol concentration and mortality from coronary artery disease. Geographic variations in incidence largely are unrelated to genetic differences, since migrant groups tend to assume the large-bowel cancer incidence rates of their adopted countries. Furthermore, population groups such as Mormons and Seventh Day Adventists, whose lifestyle and dietary habits differ somewhat

TABLE 110-1 RISK FACTORS FOR THE DEVELOPMENT OF COLORECTAL CANCER

Diet: Animal fat
Hereditary syndromes
Polyposis coli
MYH-associated polyposis
Nonpolyposis syndrome (Lynch's syndrome)
Inflammatory bowel disease
<i>Streptococcus bovis</i> bacteremia
? Tobacco use

from those of their neighbors, have significantly lower-than-expected incidence and mortality rates for colorectal cancer. The incidence of colorectal cancer has increased in Japan since that nation has adopted a more “Western” diet. At least three hypotheses have been proposed to explain the relationship to diet, none of which is fully satisfactory.

ANIMAL FATS One hypothesis is that the ingestion of animal fats found in red meats and processed meat leads to an increased proportion of anaerobes in the gut microflora, resulting in the conversion of normal bile acids into carcinogens. This provocative hypothesis is supported by several reports of increased amounts of fecal anaerobes in the stools of patients with colorectal cancer. Diets high in animal (but not vegetable) fats are also associated with high serum cholesterol, which is also associated with enhanced risk for the development of colorectal adenomas and carcinomas.

INSULIN RESISTANCE The large number of calories in Western diets coupled with physical inactivity has been associated with a higher prevalence of obesity. Obese persons develop insulin resistance with increased circulating levels of insulin, leading to higher circulating concentrations of insulin-like growth factor type I (IGF-I). This growth factor appears to stimulate proliferation of the intestinal mucosa.

FIBER Contrary to prior beliefs, the results of randomized trials and case-controlled studies have *failed* to show any value for dietary fiber or diets high in fruits and vegetables in preventing the recurrence of colorectal adenomas or the development of colorectal cancer.

The weight of epidemiologic evidence, however, implicates diet as being the major etiologic factor for colorectal cancer, particularly diets high in animal fat and in calories.

HEREDITARY FACTORS AND SYNDROMES

Up to 25% of patients with colorectal cancer have a family history of the disease, suggesting a hereditary predisposition. Inherited large-bowel cancers can be divided into two main groups: the well-studied but uncommon polyposis syndromes and the more common nonpolyposis syndromes (Table 110-2).

Polyposis Coli Polyposis coli (familial polyposis of the colon) is a rare condition characterized by the appearance of thousands of adenomatous polyps throughout the large bowel. It is transmitted as an autosomal

TABLE 110-2 HEREDITABLE (AUTOSOMAL DOMINANT) GASTROINTESTINAL POLYPOSIS SYNDROMES

Syndrome	Distribution of Polyps	Histologic Type	Malignant Potential	Associated Lesions
Familial adenomatous polyposis	Large intestine	Adenoma	Common	None
Gardner's syndrome	Large and small intestines	Adenoma	Common	Osteomas, fibromas, lipomas, epidermoid cysts, ampullary cancers, congenital hypertrophy of retinal pigment epithelium
Turcot's syndrome	Large intestine	Adenoma	Common	Brain tumors
MYH-associated polyposis	Large intestine	Adenoma	Common	None
Nonpolyposis syndrome (Lynch's syndrome)	Large intestine (often proximal)	Adenoma	Common	Endometrial and ovarian tumors (most frequently) gastric, genitourinary, pancreatic, biliary cancers (less frequently)
Peutz-Jeghers syndrome	Small and large intestines, stomach	Hamartoma	Rare	Mucocutaneous pigmentation; tumors of the ovary, breast, pancreas, endometrium
Juvenile polyposis	Large and small intestines, stomach	Hamartoma, rarely progressing to adenoma	Rare	Various congenital abnormalities

dominant trait; the occasional patient with no family history probably developed the condition due to a spontaneous mutation. Polyposis coli is associated with a deletion in the long arm of chromosome 5 (including the *APC* gene) in both neoplastic (somatic mutation) and normal (germline mutation) cells. The loss of this genetic material (i.e., allelic loss) results in the absence of tumor-suppressor genes whose protein products would normally inhibit neoplastic growth. The presence of soft tissue and bony tumors, congenital hypertrophy of the retinal pigment epithelium, mesenteric desmoid tumors, and ampullary cancers in addition to the colonic polyps characterizes a subset of polyposis coli known as *Gardner's syndrome*. The appearance of malignant tumors of the central nervous system accompanying polyposis coli defines *Turcot's syndrome*. The colonic polyps in all these conditions are rarely present before puberty but are generally evident in affected individuals by age 25. If the polyposis is not treated surgically, colorectal cancer will develop in almost all patients before age 40. Polyposis coli results from a defect in the colonic mucosa, leading to an abnormal proliferative pattern and impaired DNA repair mechanisms. Once the multiple polyps are detected, patients should undergo a total colectomy. Medical therapy with nonsteroidal anti-inflammatory drugs (NSAIDs) such as sulindac and selective cyclooxygenase-2 inhibitors such as celecoxib can decrease the number and size of polyps in patients with polyposis coli; however, this effect on polyps is only temporary, and the use of NSAIDs has not been shown to reduce the risk of cancer. Colectomy remains the primary therapy/prevention. The offspring of patients with polyposis coli, who often are prepubertal when the diagnosis is made in the parent, have a 50% risk for developing this premalignant disorder and should be carefully screened by annual flexible sigmoidoscopy until age 35. Proctosigmoidoscopy is a sufficient screening procedure because polyps tend to be evenly distributed from cecum to anus, making more invasive and expensive techniques such as colonoscopy or barium enema unnecessary. Testing for occult blood in the stool is an inadequate screening maneuver. If a causative germline *APC* mutation has been identified in an affected family member, an alternative method for identifying carriers is testing DNA from peripheral blood mononuclear cells for the presence of the specific *APC* mutation. The detection of such a germline mutation can lead to a definitive diagnosis before the development of polyps.

MYH-Associated Polyposis MYH-associated polyposis (MAP) is a rare autosomal recessive syndrome caused by a biallelic mutation in the *MUT4H* gene. This hereditary condition may have a variable clinical presentation, resembling polyposis coli or colorectal cancer occurring in younger individuals without polyposis. Screening and colectomy guidelines for this syndrome are less clear than for polyposis coli, but annual to biennial colonoscopic surveillance is generally recommended starting at age 25–30.

Hereditary Nonpolyposis Colon Cancer Hereditary nonpolyposis colon cancer (HNPCC), also known as *Lynch's syndrome*, is another autosomal dominant trait. It is characterized by the presence of three or more relatives with histologically documented colorectal cancer, one of whom is a first-degree relative of the other two; one or more cases of colorectal cancer diagnosed before age 50 in the family; and colorectal cancer involving at least two generations. In contrast to polyposis coli, HNPCC is associated with an unusually high frequency of cancer arising in the proximal large bowel. The median age for the appearance of an adenocarcinoma is <50 years, 10–15 years younger than the median age for the general population. Despite having a poorly differentiated, mucinous histologic appearance, the proximal colon tumors that characterize HNPCC have a better prognosis than sporadic tumors from patients of similar age. Families with HNPCC often include individuals with multiple primary cancers; the association of colorectal cancer with either ovarian or endometrial carcinomas is especially strong in women, and an increased appearance of gastric, small-bowel, genitourinary, pancreaticobiliary, and sebaceous skin tumors has been reported as well. It has been recommended that members of such families undergo annual or biennial colonoscopy beginning at age 25 years, with intermittent pelvic ultrasonography and endometrial biopsy for afflicted women; such a screening strategy

has not yet been validated. HNPCC is associated with germline mutations of several genes, particularly *hMSH2* on chromosome 2 and *hMLH1* on chromosome 3. These mutations lead to errors in DNA replication and are thought to result in DNA instability because of defective repair of DNA mismatches resulting in abnormal cell growth and tumor development. Testing tumor cells through molecular analysis of DNA or immunohistochemical staining of paraffin-fixed tissue for “microsatellite instability” (sequence changes reflecting defective mismatch repair) in patients with colorectal cancer and a positive family history for colorectal or endometrial cancer may identify probands with HNPCC.

INFLAMMATORY BOWEL DISEASE

(Chap. 351) Large-bowel cancer is increased in incidence in patients with long-standing inflammatory bowel disease (IBD). Cancers develop more commonly in patients with ulcerative colitis than in those with granulomatous (i.e., Crohn's) colitis, but this impression may result in part from the occasional difficulty of differentiating these two conditions. The risk of colorectal cancer in a patient with IBD is relatively small during the initial 10 years of the disease, but then appears to increase at a rate of ~0.5–1% per year. Cancer may develop in 8–30% of patients after 25 years. The risk is higher in younger patients with pancolitis.

Cancer surveillance strategies in patients with IBD are unsatisfactory. Symptoms such as bloody diarrhea, abdominal cramping, and obstruction, which may signal the appearance of a tumor, are similar to the complaints caused by a flare-up of the underlying disease. In patients with a history of IBD lasting ≥15 years who continue to experience exacerbations, the surgical removal of the colon can significantly reduce the risk for cancer and also eliminate the target organ for the underlying chronic gastrointestinal disorder. The value of such surveillance techniques as colonoscopy with mucosal biopsies and brushings for less symptomatic individuals with chronic IBD is uncertain. The lack of uniformity regarding the pathologic criteria that characterize dysplasia and the absence of data that such surveillance reduces the development of lethal cancers have made this costly practice an area of controversy.

OTHER HIGH-RISK CONDITIONS

Streptococcus bovis Bacteremia For unknown reasons, individuals who develop endocarditis or septicemia from this fecal bacterium have a high incidence of occult colorectal tumors and, possibly, upper gastrointestinal cancers as well. Endoscopic or radiographic screening appears advisable.

Tobacco Use Cigarette smoking is linked to the development of colorectal adenomas, particularly after >35 years of tobacco use. No biologic explanation for this association has yet been proposed.

PRIMARY PREVENTION

Several orally administered compounds have been assessed as possible inhibitors of colon cancer. The most effective class of chemopreventive agents is aspirin and other NSAIDs, which are thought to suppress cell proliferation by inhibiting prostaglandin synthesis. Regular aspirin use reduces the risk of colon adenomas and carcinomas as well as death from large-bowel cancer; such use also appears to diminish the likelihood for developing additional premalignant adenomas following successful treatment for a prior colon carcinoma. This effect of aspirin on colon carcinogenesis increases with the duration and dosage of drug use. Oral folic acid supplements and oral calcium supplements appear to reduce the risk of adenomatous polyps and colorectal cancers in case-controlled studies. The value of vitamin D as a form of chemoprevention is under study. Antioxidant vitamins such as ascorbic acid, tocopherols, and β-carotene are ineffective at reducing the incidence of subsequent adenomas in patients who have undergone the removal of a colon adenoma. Estrogen replacement therapy has been associated with a reduction in the risk of colorectal cancer in women, conceivably by an effect on bile acid synthesis and composition or by decreasing synthesis of IGF-I.

The rationale for colorectal cancer screening programs is that the removal of adenomatous polyps will prevent colorectal cancer, and that earlier detection of localized, superficial cancers in asymptomatic individuals will increase the surgical cure rate. Such screening programs are particularly important for individuals with a family history of the disease in first-degree relatives. The relative risk for developing colorectal cancer increases to 1.75 in such individuals and may be even higher if the relative was afflicted before age 60. The prior use of proctosigmoidoscopy as a screening tool was based on the observation that 60% of early lesions are located in the rectosigmoid. For unexplained reasons, however, the proportion of large-bowel cancers arising in the rectum has been decreasing during the past several decades, with a corresponding increase in the proportion of cancers in the more proximal descending colon. As such, the potential for proctosigmoidoscopy to detect a sufficient number of occult neoplasms to make the procedure cost-effective has been questioned.

Screening strategies for colorectal cancer that have been examined during the past several decades are listed in [Table 110-3](#).

Many programs directed at the early detection of colorectal cancers have focused on digital rectal examinations and fecal occult blood (i.e., stool guaiac) testing. The digital examination should be part of any routine physical evaluation in adults older than age 40 years, serving as a screening test for prostate cancer in men, a component of the pelvic examination in women, and an inexpensive maneuver for the detection of masses in the rectum. However, because of the proximal migration of colorectal tumors, its value as an overall screening modality for colorectal cancer has become limited. The development of the fecal occult blood test has greatly facilitated the detection of occult fecal blood. Unfortunately, even when performed optimally, the fecal occult blood test has major limitations as a screening technique. About 50% of patients with documented colorectal cancers have a negative fecal occult blood test, consistent with the intermittent bleeding pattern of these tumors. When random cohorts of asymptomatic persons have been tested, 2–4% have fecal occult blood-positive stools. Colorectal cancers have been found in <10% of these “test-positive” cases, with benign polyps being detected in an additional 20–30%. Thus, a colorectal neoplasm will not be found in most asymptomatic individuals with occult blood in their stool. Nonetheless, persons found to have fecal occult blood-positive stool routinely undergo further medical evaluation, including sigmoidoscopy and/or colonoscopy—procedures that are not only uncomfortable and expensive but also associated with a small risk for significant complications. The added cost of these studies would appear justifiable if the small number of patients found to have occult neoplasms because of fecal occult blood screening could be shown to have an improved prognosis and prolonged survival. Prospectively controlled trials have shown a statistically significant reduction in mortality rate from colorectal cancer for individuals undergoing annual stool guaiac screening. However, this benefit only emerged after >13 years of follow-up and was extremely expensive to achieve, because all positive tests (most of which were falsely positive) were followed by colonoscopy. Moreover, these colonoscopic examinations quite likely provided the opportunity for cancer prevention through the removal of potentially premalignant adenomatous polyps

because the eventual development of cancer was reduced by 20% in the cohort undergoing annual screening.

With the appreciation that the carcinogenic process leading to the progression of the normal bowel mucosa to an adenomatous polyp and then to a cancer is the result of a series of molecular changes, investigators have examined fecal DNA for evidence of mutations associated with such molecular changes as evidence of the occult presence of precancerous lesions or actual malignancies. Such a strategy has been tested in more than 4000 asymptomatic individuals whose stool was assessed for occult blood and for 21 possible mutations in fecal DNA; these study subjects also underwent colonoscopy. Although the fecal DNA strategy suggested the presence of more advanced adenomas and cancers than did the fecal occult blood testing approach, the overall sensitivity, using colonoscopic findings as the standard, was less than 50%, diminishing enthusiasm for further pursuit of the fecal DNA screening strategy.

The use of imaging studies to screen for colorectal cancers has also been explored. Air contrast barium enemas had been used to identify sources of occult blood in the stool prior to the advent of fiberoptic endoscopy; the cumbersome nature of the procedure and inconvenience to patients limited its widespread adoption. The introduction of computed tomography (CT) scanning led to the development of virtual (i.e., CT) colonography as an alternative to the growing use of endoscopic screening techniques. Virtual colonography was proposed as being equivalent in sensitivity to colonoscopy and being available in a more widespread manner because it did not require the same degree of operator expertise as fiberoptic endoscopy. However, virtual colonography requires the same cathartic preparation that has limited widespread acceptance of endoscopic colonoscopy, is diagnostic but not therapeutic (i.e., patients with suspicious findings must undergo a subsequent endoscopic procedure for polypectomy or biopsy), and, in the setting of general radiology practices, appears to be less sensitive as a screening technique when compared with endoscopic procedures.

With the appreciation of the inadequacy of fecal occult blood testing alone, concerns about the practicality of imaging approaches, and the wider adoption of endoscopic examinations by the primary care community, screening strategies in asymptomatic persons have changed. At present, both the American Cancer Society and the National Comprehensive Cancer Network suggest either fecal occult blood testing annually coupled with flexible sigmoidoscopy every 5 years or colonoscopy every 10 years beginning at age 50 in asymptomatic individuals with no personal or family history of polyps or colorectal cancer. The recommendation for the inclusion of flexible sigmoidoscopy is strongly supported by the recently published results of three randomized trials performed in the United States, the United Kingdom, and Italy, involving more than 350,000 individuals, which consistently showed that periodic (even single) sigmoidoscopic examinations, after more than a decade of median follow-up, lead to an approximate 21% reduction in the development of colorectal cancer and a more than 25% reduction in mortality from the malignant disease. Less than 20% of participants in these studies underwent a subsequent colonoscopy. In contrast to the cathartic preparation required before colonoscopic procedures, which is only performed by highly trained specialists, flexible sigmoidoscopy requires only an enema as preparation and can be accurately performed by nonspecialty physicians or physician-extenders. The randomized screening studies using flexible sigmoidoscopy led to the estimate that approximately 650 individuals needed to be screened to prevent one colorectal cancer death; this contrasts with the data for mammography where the number of women needing to be screened to prevent one breast cancer death is 2500, reinforcing the efficacy of endoscopic surveillance for colorectal cancer screening. Presumably the benefit from the sigmoidoscopic screening is the result of the identification and removal of adenomatous polyps; it is intriguing that this benefit has been achieved using a technique that leaves the proximal half of the large bowel unvisualized.

It remains to be seen whether surveillance colonoscopy, which has gained increasing popularity in the United States for colorectal cancer screening, will prove to be more effective than flexible sigmoidoscopy. Ongoing randomized trials being conducted in Europe are addressing

TABLE 110-3 SCREENING STRATEGIES FOR COLORECTAL CANCER

Digital rectal examination
Stool testing
• Occult blood
• Fecal DNA
Imaging
• Contrast barium enema
• Virtual (i.e., computed tomography colonography)
Endoscopy
• Flexible sigmoidoscopy
• Colonoscopy

this issue. Although flexible sigmoidoscopy only visualizes the distal half of the large bowel, leading to the assumption that colonoscopy represents a more informative approach, colonoscopy has been reported as being less accurate for screening the proximal rather than the distal colon, perhaps due to technical considerations but also possibly because of a greater frequency of serrated (i.e., “flat”) polyps in the right colon, which are more difficult to identify. At present, colonoscopy performed every 10 years has been offered as an alternative to annual fecal occult blood testing with periodic (every 5 years) flexible sigmoidoscopy. Colonoscopy has been shown to be superior to double-contrast barium enema and also to have a higher sensitivity for detecting villous or dysplastic adenomas or cancers than the strategy using occult fecal blood testing and flexible sigmoidoscopy. Whether colonoscopy performed every 10 years beginning at age 50 is medically superior and economically equivalent to flexible sigmoidoscopy remains to be determined.

CLINICAL FEATURES

Presenting Symptoms Symptoms vary with the anatomic location of the tumor. Because stool is relatively liquid as it passes through the ileocecal valve into the right colon, cancers arising in the cecum and ascending colon may become quite large without resulting in any obstructive symptoms or noticeable alterations in bowel habits. Lesions of the right colon commonly ulcerate, leading to chronic, insidious blood loss without a change in the appearance of the stool. Consequently, patients with tumors of the ascending colon often present with symptoms such as fatigue, palpitations, and even angina pectoris and are found to have a hypochromic, microcytic anemia indicative of iron deficiency. Because the cancers may bleed intermittently, a random fecal occult blood test may be negative. As a result, the unexplained presence of iron-deficiency anemia in any adult (with the possible exception of a premenopausal, multiparous woman) mandates a thorough endoscopic and/or radiographic visualization of the entire large bowel (Fig. 110-1).

Because stool becomes more formed as it passes into the transverse and descending colon, tumors arising there tend to impede the passage of stool, resulting in the development of abdominal cramping, occasional obstruction, and even perforation. Radiographs of the abdomen often reveal characteristic annular, constricting lesions (“apple-core” or “napkin-ring”) (Fig. 110-2).

Cancers arising in the rectosigmoid are often associated with hematochezia, tenesmus, and narrowing of the caliber of stool; anemia is an infrequent finding. While these symptoms may lead patients and their physicians to suspect the presence of hemorrhoids, the development of rectal bleeding and/or altered bowel habits demands a prompt digital rectal examination and proctosigmoidoscopy.

Staging, Prognostic Factors, and Patterns of Spread The prognosis for individuals having colorectal cancer is related to the depth of tumor penetration into the bowel wall and the presence of both regional lymph node involvement and distant metastases. These variables are incorporated into the staging system introduced by Dukes and subsequently applied to a TNM classification method, in which T represents the depth of tumor penetration, N the presence of lymph node involvement, and M the presence or absence of distant metastases (Fig. 110-3). Superficial lesions that do not involve regional lymph nodes and do not penetrate through the submucosa (T1) or the muscularis (T2) are designated as *stage I* (T1–2N0M0) disease; tumors that penetrate through the muscularis but have not spread to lymph nodes are *stage II* disease (T3–4N0M0); regional lymph node involvement defines *stage III* (TXN1–2M0) disease; and metastatic spread to sites such as liver, lung, or bone indicates *stage IV* (TXNXM1) disease. Unless gross evidence of metastatic disease is present, disease stage cannot be determined accurately before surgical resection and pathologic analysis of the operative specimens. It is not clear whether the detection of nodal metastases by special immunohistochemical molecular techniques has the same prognostic implications as disease detected by routine light microscopy.

Most recurrences after a surgical resection of a large-bowel cancer occur within the first 4 years, making 5-year survival a fairly reliable



FIGURE 110-1 Double-contrast air-barium enema revealing a sessile tumor of the cecum in a patient with iron-deficiency anemia and guaiac-positive stool. The lesion at surgery was a stage II adenocarcinoma.

indicator of cure. The likelihood for 5-year survival in patients with colorectal cancer is stage-related (Fig. 110-3). That likelihood has improved during the past several decades when similar surgical stages have been compared. The most plausible explanation for this improvement is more thorough intraoperative and pathologic staging. In particular, more exacting attention to pathologic detail has revealed that the prognosis following the resection of a colorectal cancer is not related merely to the presence or absence of regional lymph node involvement; rather, prognosis may be more precisely gauged by the number of involved lymph nodes (one to three lymph nodes [“N1”] vs four or more lymph nodes [“N2”]) and the number of nodes examined. A minimum of 12 sampled lymph nodes is thought necessary to



FIGURE 110-2 Annular, constricting adenocarcinoma of the descending colon. This radiographic appearance is referred to as an “apple-core” lesion and is always highly suggestive of malignancy.

Stage	I		II	III		IV
	T1	T2	T3	N1	N2	M
Extent of tumor	No deeper than submucosa	Not through muscularis	Through muscularis	1–3 lymph node metastases	≥4 lymph node metastases	Distant metastases
5-year survival	>95%	>90%	70–85%	50–70%	25–60%	<5%
Stage at presentation	Colon	23%	31%	26%		20%
	Rectal	34%	25%	26%		15%

FIGURE 110-3 Staging and prognosis for patients with colorectal cancer.

accurately define tumor stage, and the more nodes examined, the better. Other predictors of a poor prognosis after a total surgical resection include tumor penetration through the bowel wall into pericolonic fat, poorly differentiated histology, perforation and/or tumor adherence to adjacent organs (increasing the risk for an anatomically adjacent recurrence), and venous invasion by tumor (Table 110-4). Regardless of the clinicopathologic stage, a preoperative elevation of the plasma carcinoembryonic antigen (CEA) level predicts eventual tumor recurrence. The presence of aneuploidy and specific chromosomal deletions, such as a mutation in the *b-raf* gene in tumor cells, appears to predict for a higher risk for metastatic spread. Conversely, the detection of microsatellite instability in tumor tissue indicates a more favorable outcome. In contrast to most other cancers, the prognosis in colorectal cancer is not influenced by the size of the primary lesion when adjusted for nodal involvement and histologic differentiation.

Cancers of the large bowel generally spread to regional lymph nodes or to the liver via the portal venous circulation. The liver represents the most frequent visceral site of metastasis; it is the initial site of distant spread in one-third of recurring colorectal cancers and is involved in more than two-thirds of such patients at the time of death. In general, colorectal cancer rarely spreads to the lungs, supraclavicular lymph nodes, bone, or brain without prior spread to the liver. A major exception to this rule occurs in patients having primary tumors in the distal rectum, from which tumor cells may spread through the paravertebral

venous plexus, escaping the portal venous system and thereby reaching the lungs or supraclavicular lymph nodes without hepatic involvement. The median survival after the detection of distant metastases has ranged in the past from 6–9 months (hepatomegaly, abnormal liver chemistries) to 24–30 months (small liver nodule initially identified by elevated CEA level and subsequent CT scan), but effective systemic therapy is significantly improving this prognosis.

Efforts to use gene expression profiles to identify patients at risk of recurrence or those particularly likely to benefit from adjuvant therapy have not yet yielded practice-changing results. Despite a burgeoning literature examining a host of prognostic factors, pathologic stage at diagnosis remains the best predictor of long-term prognosis. Patients with lymphovascular invasion and high preoperative CEA levels are likely to have a more aggressive clinical course.

TREATMENT COLORECTAL CANCER

Total resection of tumor is the optimal treatment when a malignant lesion is detected in the large bowel. An evaluation for the presence of metastatic disease, including a thorough physical examination, biochemical assessment of liver function, measurement of the plasma CEA level, and a CT scan of the chest, abdomen, and pelvis, should be performed before surgery. When possible, a colonoscopy of the entire large bowel should be performed to identify synchronous neoplasms and/or polyps. The detection of metastases should not preclude surgery in patients with tumor-related symptoms such as gastrointestinal bleeding or obstruction, but it often prompts the use of a less radical operative procedure. The necessity for a primary tumor resection in asymptomatic individuals with metastatic disease is an area of controversy. At the time of laparotomy, the entire peritoneal cavity should be examined, with thorough inspection of the liver, pelvis, and hemidiaphragm and careful palpation of the full length of the large bowel. Following recovery from a complete resection, patients should be observed carefully for 5 years by semi-annual physical examinations and blood chemistry measurements. If a complete colonoscopy was not performed preoperatively, it should be carried out within the first several postoperative months. Some authorities favor measuring plasma CEA levels at 3-month intervals because of the sensitivity of this test as a marker for otherwise undetectable tumor recurrence. Subsequent endoscopic

TABLE 110-4 PREDICTORS OF POOR OUTCOME FOLLOWING TOTAL SURGICAL RESECTION OF COLORECTAL CANCER

Tumor spread to regional lymph nodes
Number of regional lymph nodes involved
Tumor penetration through the bowel wall
Poorly differentiated histology
Perforation
Tumor adherence to adjacent organs
Venous invasion
Preoperative elevation of CEA titer (>5 ng/mL)
Aneuploidy
Specific chromosomal deletion (e.g., mutation in the <i>b-raf</i> gene)

Abbreviation: CEA, carcinoembryonic antigen.

surveillance of the large bowel, probably at triennial intervals, is indicated, because patients who have been cured of one colorectal cancer have a 3–5% probability of developing an additional bowel cancer during their lifetime and a >15% risk for the development of adenomatous polyps. Anastomotic (“suture-line”) recurrences are infrequent in colorectal cancer patients, provided the surgical resection margins are adequate and free of tumor. The value of periodic CT scans of the abdomen, assessing for an early, asymptomatic indication of tumor recurrence, is an area of uncertainty, with some experts recommending the test be performed annually for the first 3 postoperative years.

Radiation therapy to the pelvis is recommended for patients with rectal cancer because it reduces the 20–25% probability of regional recurrences following complete surgical resection of stage II or III tumors, especially if they have penetrated through the serosa. This alarmingly high rate of local disease recurrence is believed to be due to the fact that the contained anatomic space within the pelvis limits the extent of the resection and because the rich lymphatic network of the pelvic side wall immediately adjacent to the rectum facilitates the early spread of malignant cells into surgically inaccessible tissue. The use of sharp rather than blunt dissection of rectal cancers (*total mesorectal excision*) appears to reduce the likelihood of local disease recurrence to ~10%. Radiation therapy, either pre- or postoperatively, further reduces the likelihood of pelvic recurrences but does not appear to prolong survival. Combining radiation therapy with 5-fluorouracil (5-FU)-based chemotherapy, preferably prior to surgical resection, lowers local recurrence rates and improves overall survival. Preoperative radiotherapy is indicated for patients with large, potentially unresectable rectal cancers; such lesions may shrink enough to permit subsequent surgical removal. Radiation therapy is not effective as the primary treatment of colon cancer.

Systemic therapy for patients with colorectal cancer has become more effective. 5-FU remains the backbone of treatment for this disease. Partial responses are obtained in 15–20% of patients. The probability of tumor response appears to be somewhat greater for patients with liver metastases when chemotherapy is infused directly into the hepatic artery, but intraarterial treatment is costly and toxic and does not appear to appreciably prolong survival. The concomitant administration of folinic acid (leucovorin) improves the efficacy of 5-FU in patients with advanced colorectal cancer, presumably by enhancing the binding of 5-FU to its target enzyme, thymidylate synthase. A threefold improvement in the partial response rate is noted when folinic acid is combined with 5-FU; however, the effect on survival is marginal, and the optimal dose schedule remains to be defined. 5-FU is generally administered intravenously but may also be given orally in the form of capecitabine (Xeloda) with seemingly similar efficacy.

Irinotecan (CPT-11), a topoisomerase 1 inhibitor, prolongs survival when compared to supportive care in patients whose disease has progressed on 5-FU. Furthermore, the addition of irinotecan to 5-FU and leucovorin (LV) (e.g., FOLFIRI) improves response rates and survival of patients with metastatic disease. The *FOLFIRI regimen* is as follows: irinotecan, 180 mg/m² as a 90-min infusion on day 1; LV, 400 mg/m² as a 2-h infusion during irinotecan administration; immediately followed by 5-FU bolus, 400 mg/m², and 46-h continuous infusion of 2.4–3 g/m² every 2 weeks. Diarrhea is the major side effect from irinotecan. Oxaliplatin, a platinum analogue, also improves the response rate when added to 5-FU and LV (FOLFOX) as initial treatment of patients with metastatic disease. The *FOLFOX regimen* is as follows: 2-h infusion of LV (400 mg/m² per day) followed by a 5-FU bolus (400 mg/m² per day) and 22-h infusion (1200 mg/m²) every 2 weeks, together with oxaliplatin, 85 mg/m² as a 2-h infusion on day 1. Oxaliplatin frequently causes a dose-dependent sensory neuropathy that often but not always resolves following the cessation of therapy. FOLFIRI and FOLFOX are equal in efficacy. In metastatic disease, these regimens may produce median survivals of 2 years.

Monoclonal antibodies are also effective in patients with advanced colorectal cancer. Cetuximab (Erbix) and panitumumab

(Vectibix) are directed against the epidermal growth factor receptor (EGFR), a transmembrane glycoprotein involved in signaling pathways affecting growth and proliferation of tumor cells. Both cetuximab and panitumumab, when given alone, have been shown to benefit a small proportion of previously treated patients, and cetuximab appears to have therapeutic synergy with such chemotherapeutic agents as irinotecan, even in patients previously resistant to this drug; this suggests that cetuximab can reverse cellular resistance to cytotoxic chemotherapy. The antibodies are not effective in the approximate 40% subset of colon tumors that contain mutated *K-ras*. The use of both cetuximab and panitumumab can lead to an acne-like rash, with the development and severity of the rash being correlated with the likelihood of antitumor efficacy. Inhibitors of the EGFR tyrosine kinase such as erlotinib (Tarceva) or sunitinib (Sutent) do not appear to be effective in colorectal cancer.

Bevacizumab (Avastin) is a monoclonal antibody directed against the vascular endothelial growth factor (VEGF) and is thought to act as an antiangiogenesis agent. The addition of bevacizumab to irinotecan-containing combinations and to FOLFOX initially appeared to significantly improve the outcome observed with chemotherapy alone, but subsequent studies have suggested a lesser degree of benefit. The use of bevacizumab can lead to hypertension, proteinuria, and an increased likelihood of thromboembolic events.

Patients with solitary hepatic metastases without clinical or radiographic evidence of additional tumor involvement should be considered for partial liver resection, because such procedures are associated with 5-year survival rates of 25–30% when performed on selected individuals by experienced surgeons.

The administration of 5-FU and LV for 6 months after resection of tumor in patients with stage III disease leads to a 40% decrease in recurrence rates and 30% improvement in survival. The likelihood of recurrence has been further reduced when oxaliplatin has been combined with 5-FU and LV (e.g., FOLFOX); unexpectedly, the addition of irinotecan to 5-FU and LV as well as the addition of either bevacizumab or cetuximab to FOLFOX did not significantly enhance outcome. Patients with stage II tumors do not appear to benefit appreciably from adjuvant therapy, with the use of such treatment generally restricted to those patients having biologic characteristics (e.g., perforated tumors, T4 lesions, lymphovascular invasion) that place them at higher likelihood for recurrence. The addition of oxaliplatin to adjuvant treatment for patients older than age 70 and those with stage II disease does not appear to provide any therapeutic benefit.

In rectal cancer, the delivery of preoperative or postoperative combined-modality therapy (5-FU plus radiation therapy) reduces the risk of recurrence and increases the chance of cure for patients with stage II and III tumors, with the preoperative approach being better tolerated. The 5-FU acts as a radiosensitizer when delivered together with radiation therapy. Life-extending adjuvant therapy is used in only about half of patients older than age 65 years. This age bias is unfortunate because the benefits and likely the tolerance of adjuvant therapy in patients age ≥65 years appear similar to those seen in younger individuals.

CANCERS OF THE ANUS

Cancers of the anus account for 1–2% of the malignant tumors of the large bowel. Most such lesions arise in the anal canal, the anatomic area extending from the anorectal ring to a zone approximately halfway between the pectinate (or dentate) line and the anal verge. Carcinomas arising proximal to the pectinate line (i.e., in the transitional zone between the glandular mucosa of the rectum and the squamous epithelium of the distal anus) are known as *basaloid*, *cuboidal*, or *cloacogenic* tumors; about one-third of anal cancers have this histologic pattern. Malignancies arising distal to the pectinate line have squamous histology, ulcerate more frequently, and constitute ~55% of anal cancers. The prognosis for patients with basaloid and squamous cell cancers of

544 the anus is identical when corrected for tumor size and the presence or absence of nodal spread.

The development of anal cancer is associated with infection by human papillomavirus, the same organism etiologically linked to cervical cancer. The virus is sexually transmitted. The infection may lead to anal warts (condyloma acuminata), which may progress to anal intraepithelial neoplasia and on to squamous cell carcinoma. The risk for anal cancer is increased among homosexual males, presumably related to anal intercourse. Anal cancer risk is increased in both men and women with AIDS, possibly because their immunosuppressed state permits more severe papillomavirus infection. Vaccination against human papilloma viruses may reduce the eventual risk for anal cancer. Anal cancers occur most commonly in middle-aged persons and are more frequent in women than men. At diagnosis, patients may experience bleeding, pain, sensation of a perianal mass, and pruritus.

Radical surgery (abdominal-perineal resection with lymph node sampling and a permanent colostomy) was once the treatment of choice for this tumor type. The 5-year survival rate after such a procedure was 55–70% in the absence of spread to regional lymph nodes and <20% if nodal involvement was present. An alternative therapeutic approach combining external beam radiation therapy with concomitant chemotherapy (5-FU and mitomycin C) has resulted in biopsy-proven disappearance of all tumor in >80% of patients whose initial lesion was <3 cm in size. Tumor recurrences develop in <10% of these patients, meaning that ~70% of patients with anal cancers can be cured with nonoperative treatment and without the need for a colostomy. Surgery should be reserved for the minority of individuals who are found to have residual tumor after being managed initially with radiation therapy combined with chemotherapy.

TABLE 111-1 AGE-ADJUSTED INCIDENCE RATES FOR HEPATOCELLULAR CARCINOMA

Country	Persons per 100,000 per Year	
	Male	Female
Argentina	6.0	2.5
Brazil, Recife	9.2	8.3
Brazil, Sao Paulo	3.8	2.6
Mozambique	112.9	30.8
South Africa, Cape: Black	26.3	8.4
South Africa, Cape: White	1.2	0.6
Senegal	25.6	9.0
Nigeria	15.4	3.2
Gambia	33.1	12.6
Burma	25.5	8.8
Japan	7.2	2.2
Korea	13.8	3.2
China, Shanghai	34.4	11.6
India, Bombay	4.9	2.5
India, Madras	2.1	0.7
Great Britain	1.6	0.8
France	6.9	1.2
Italy, Varese	7.1	2.7
Norway	1.8	1.1
Spain, Navarra	7.9	4.7

EPIDEMIOLOGY

There are two general types of epidemiologic studies of HCC—those of country-based incidence rates (Table 111-1) and those of migrants. Endemic hot spots occur in areas of China and sub-Saharan Africa, which are associated both with high endemic hepatitis B carrier rates as well as mycotoxin contamination of foodstuffs (aflatoxin B₁), stored grains, drinking water, and soil. Environmental factors are important, for example, Japanese in Japan have a higher incidence than Japanese living in Hawaii, who in turn have a higher incidence than those living in California.

ETIOLOGIC FACTORS

Chemical Carcinogens Causative agents for HCC have been studied along two general lines. First are agents identified as carcinogenic in experimental animals (particularly rodents) that are thought to be present in the human environment (Table 111-2). Second is the association of HCC with various other clinical conditions. Probably the best-studied and most potent ubiquitous natural chemical carcinogen is a product of the *Aspergillus* fungus, called aflatoxin B₁. This mold and aflatoxin product can be found in a variety of stored grains in hot, humid places, where peanuts and rice are stored in unrefrigerated conditions. Aflatoxin contamination of foodstuffs correlates well with incidence rates in Africa and to some extent in China. In endemic areas of China, even farm animals such as ducks have HCC. The most potent carcinogens appear to be natural products of plants, fungi, and bacteria, such as bush trees containing pyrrolizidine alkaloids as well as tannic acid and safrole. Pollutants such as pesticides and insecticides are known rodent carcinogens.

Hepatitis Both case-control and cohort studies have shown a strong association between chronic hepatitis B carrier rates and increased incidence of HCC. In Taiwanese male postal carriers who were hepatitis B surface antigen (HBsAg)-positive, a 98-fold greater risk for HCC was found compared to HBsAg-negative individuals. The incidence of HCC in Alaskan natives is markedly increased related to a high prevalence of HBV infection. HBV-based HCC may involve rounds of hepatic destruction with subsequent proliferation and not necessarily frank cirrhosis. The increase in Japanese

111 Tumors of the Liver and Biliary Tree

Brian I. Carr

HEPATOCELLULAR CARCINOMA

INCIDENCE

Hepatocellular carcinoma (HCC) is one of the most common malignancies worldwide. The annual global incidence is approximately 1 million cases, with a male-to-female ratio of approximately 4:1 (1:1 without cirrhosis to 9:1 in many high-incidence countries). The incidence rate equals the death rate. In the United States, approximately 22,000 new cases are diagnosed annually, with 18,000 deaths. The death rates in males in low-incidence countries such as the United States are 1.9 per 100,000 per year; in intermediate areas such as Austria and South Africa, they range from 5.1–20; and in high-incidence areas such as in the Orient (China and Korea), they are as high as 23.1–150 per 100,000 per year (Table 111-1). The incidence of HCC in the United States is approximately 3 per 100,000 persons, with significant gender, ethnic, and geographic variations. These numbers are rapidly increasing and may be an underestimate. Approximately 4 million chronic hepatitis C virus (HCV) carriers are in the United States alone. Approximately 10% of them, or 400,000, are likely to develop cirrhosis. Approximately 5%, or 20,000, of these patients may develop HCC annually. Add to this the two other common predisposing factors—hepatitis B virus (HBV) and chronic alcohol consumption—and 60,000 new HCC cases annually seem possible. Future advances in HCC survival will likely depend in part on immunization strategies for HBV (and HCV) and earlier diagnosis by screening of patients at risk of HCC development.

Current Directions With the U.S. HCV epidemic, HCC is increasing in most states, and obesity-associated liver disease (nonalcoholic steatohepatitis [NASH]) is increasingly recognized as a cause.

TABLE 111-2 FACTORS ASSOCIATED WITH AN INCREASED RISK OF DEVELOPING HEPATOCELLULAR CARCINOMA

Common	Unusual
Cirrhosis from any cause	Primary biliary cirrhosis
Hepatitis B or C chronic infection	Hemochromatosis
Ethanol chronic consumption	α_1 Antitrypsin deficiency
NASH/NAFL	Glycogen storage diseases
Aflatoxin B ₁ or other mycotoxins	Citrullinemia
	Porphyria cutanea tarda
	Hereditary tyrosinemia
	Wilson's disease

Abbreviations: NAFL, nonalcoholic fatty liver; NASH, nonalcoholic steatohepatitis.

HCC incidence rates in the last three decades is thought to be from hepatitis C. A large-scale World Health Organization (WHO)-sponsored intervention study is currently under way in Asia involving HBV vaccination of the newborn. HCC in African blacks is not associated with severe cirrhosis but is poorly differentiated and very aggressive. Despite uniform HBV carrier rates among the South African Bantu, there is a ninefold difference in HCC incidence between Mozambicans living along the coast and inland. These differences are attributed to the additional exposure to dietary aflatoxin B₁ and other carcinogenic mycotoxins. A typical interval between HCV-associated transfusion and subsequent HCC is approximately 30 years. HCV-associated HCC patients tend to have more frequent and advanced cirrhosis, but in HBV-associated HCC, only half the patients have cirrhosis, with the remainder having chronic active hepatitis (**Chap. 362**).



Other Etiologic Conditions The 75–85% association of HCC with underlying cirrhosis has long been recognized, more typically with macronodular cirrhosis in Southeast Asia, but also with micronodular cirrhosis (alcohol) in Europe and the United States (**Chap. 365**). It is still not clear whether cirrhosis itself is a predisposing factor to the development of HCC or whether the underlying causes of the cirrhosis are actually the carcinogenic factors. However, ~20% of U.S. patients with HCC do not have underlying cirrhosis. Several underlying conditions are associated with an increased risk for cirrhosis-associated HCC (Table 111-2), including hepatitis, alcohol, autoimmune chronic active hepatitis, cryptogenic cirrhosis, and NASH. A less common association is with primary biliary cirrhosis and several metabolic diseases including hemochromatosis, Wilson's disease, α_1 antitrypsin deficiency, tyrosinemia, porphyria cutanea tarda, glycogenesis types 1 and 3, citrullinemia, and orotic aciduria. The etiology of HCC in those 20% of patients who have no cirrhosis is currently unclear, and their HCC natural history is not well-defined.

Current Directions Many patients have multiple etiologies, and the interactions of HBV, HCV, alcohol, smoking, and aflatoxins are just beginning to be explored.

CLINICAL FEATURES



Symptoms These include abdominal pain, weight loss, weakness, abdominal fullness and swelling, jaundice, and nausea (**Table 111-3**). Presenting signs and symptoms differ somewhat between high- and low-incidence areas. In high-risk areas, especially in South African blacks, the most common symptom is abdominal pain; by contrast, only 40–50% of Chinese and Japanese patients present with abdominal pain. Abdominal swelling may occur as a consequence of ascites due to the underlying chronic liver disease or may be due to a rapidly expanding tumor. Occasionally, central necrosis or acute hemorrhage into the peritoneal cavity leads to death. In countries with an active surveillance program, HCC tends to be identified at an earlier stage, when symptoms may be due only to the underlying disease. Jaundice is usually due to obstruction of the intrahepatic ducts from underlying liver disease. Hematemesis may occur due to

TABLE 111-3 HEPATOCELLULAR CARCINOMA CLINICAL PRESENTATION (N = 547)

Symptom	No. of Patients (%)
No symptom	129 (24)
Abdominal pain	219 (40)
Other (workup of anemia and various diseases)	64 (12)
Routine physical exam finding, elevated LFTs	129 (24)
Weight loss	112 (20)
Appetite loss	59 (11)
Weakness/malaise	83 (15)
Jaundice	30 (5)
Routine CT scan screening of known cirrhosis	92 (17)
Cirrhosis symptoms (ankle swelling, abdominal bloating, increased girth, pruritus, GI bleed)	98 (18)
Diarrhea	7 (1)
Tumor rupture	1
Patient Characteristics	
Mean age (yr)	56 ± 13
Male:Female	3:1
Ethnicity	
White	72%
Middle Eastern	10%
Asian	13%
African American	5%
Cirrhosis	81%
No cirrhosis	19%
Tumor Characteristics	
Hepatic tumor numbers	
1	20%
2	25%
3 or more	65%
Portal vein invasion	75%
Unilobar	25%
Bilobar	75%

Abbreviations: CT, computed tomography; GI, gastrointestinal; LFT, liver function test.

esophageal varices from the underlying portal hypertension. Bone pain is seen in 3–12% of patients, but necropsies show pathologic bone metastases in ~20% of patients. However, 25% of patients may be asymptomatic.

Physical Signs Hepatomegaly is the most common physical sign, occurring in 50–90% of the patients. Abdominal bruits are noted in 6–25%, and ascites occurs in 30–60% of patients. Ascites should be examined by cytology. Splenomegaly is mainly due to portal hypertension. Weight loss and muscle wasting are common, particularly with rapidly growing or large tumors. Fever is found in 10–50% of patients, from unclear cause. The signs of chronic liver disease may often be present, including jaundice, dilated abdominal veins, palmar erythema, gynecomastia, testicular atrophy, and peripheral edema. Budd-Chiari syndrome can occur due to HCC invasion of the hepatic veins, with tense ascites and a large tender liver (**Chap. 365**).

Paraneoplastic Syndromes Most paraneoplastic syndromes in HCC are biochemical abnormalities without associated clinical consequences. They include hypoglycemia (also caused by end-stage liver failure), erythrocytosis, hypercalcemia, hypercholesterolemia, dysfibrinogenemia, carcinoid syndrome, increased thyroxin-binding globulin, changes in secondary sex characteristics (gynecomastia, testicular atrophy, and precocious puberty), and porphyria cutanea tarda. Mild hypoglycemia occurs in rapidly growing HCC as part of

546 terminal illness, and profound hypoglycemia may occur, although the cause is unclear. Erythrocytosis occurs in 3–12% of patients and hypercholesterolemia in 10–40%. A high percentage of patients have thrombocytopenia associated with their fibrosis or leukopenia, resulting from portal hypertension, and not from cancer infiltration of bone marrow, as in other tumor types. Furthermore, large HCCs have normal or high platelet levels (thrombocytosis), as in ovarian and other gastrointestinal cancers, probably related to elevated interleukin 6 (IL-6) levels.

STAGING

Multiple clinical staging systems for HCC have been described. A widely used one has been the American Joint Committee on Cancer (AJCC) tumor-node-metastasis (TNM) classification. However, the Cancer of the Liver Italian Program (CLIP) system is now popular because it takes cirrhosis into account, based on the original Okuda system (Table 111-4). Patients with Okuda stage III disease have a dire prognosis because they usually cannot be curatively resected, and the condition of their liver typically precludes chemotherapy. Other staging systems have been proposed, and a consensus is needed. They are all based on combining the prognostic features of liver damage with those of tumor aggressiveness and include the Barcelona Clinic Liver Cancer (BCLC) system from Spain (Fig. 111-1), which is externally validated and incorporates baseline survival estimates; the Chinese University Prognostic Index (CUPI); the important and simple Japan Integrated Staging Score (JIS); and SLiDe, which stands for *s*tage, *l*iver damage, and *de* s- γ -carboxy prothrombin. CLIP and BCLC appear most popular in the West, whereas JIS is favored in Japan. Each system

TABLE 111-4 CLIP AND OKUDA STAGING SYSTEMS FOR HEPATOCELLULAR CARCINOMA

CLIP Classification			
Variables	Points		
	0	1	2
i. Tumor number	Single	Multiple	–
Hepatic replacement by tumor (%)	<50	<50	>50
ii. Child-Pugh score	A	B	C
iii. α Fetoprotein level (ng/mL)	<400	\geq 400	–
iv. Portal vein thrombosis (CT)	No	Yes	–

CLIP stages (score = sum of points): CLIP 0, 0 points; CLIP 1, 1 point; CLIP 2, 2 points; CLIP 3, 3 points.

Okuda Classification							
Tumor Extent ^a	Ascites	Albumin (g/L)	Bilirubin (mg/dL)				
\geq 50%	<50	+	–	\leq 3	>3	\geq 3	<3
(+)	(–)	(+)	(–)	(+)	(–)	(+)	(–)

Okuda stages: stage 1, all (–); stage 2, 1 or 2 (+); stage 3, 3 or 4 (+).

^aExtent of liver occupied by tumor.

Abbreviation: CLIP, Cancer of the Liver Italian Program.

has its champions. The best prognosis is for stage I, solitary tumors less than 2 cm in diameter without vascular invasion. Adverse prognostic features include ascites, jaundice, vascular invasion, and elevated α fetoprotein (AFP). Vascular invasion in particular has profound effects

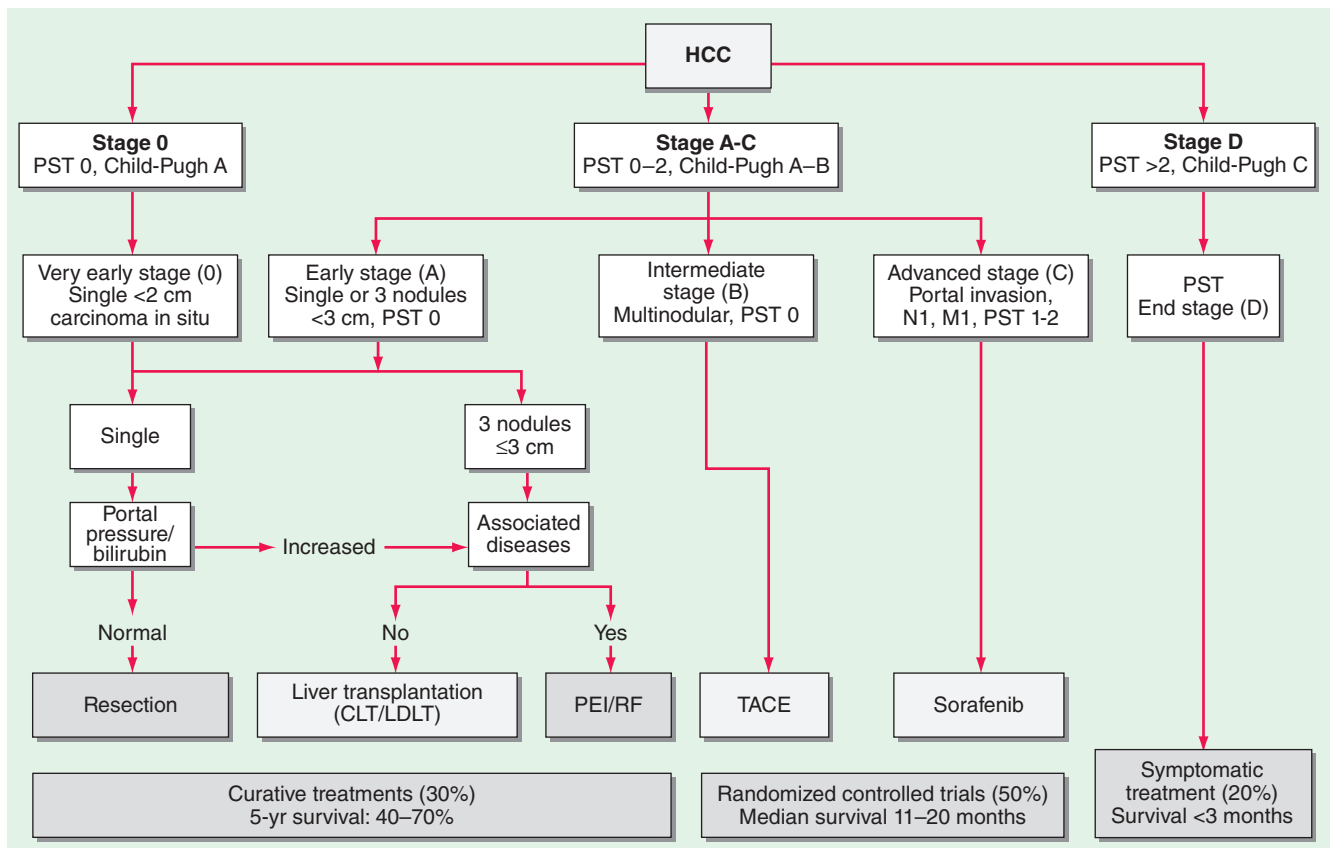


FIGURE 111-1 Barcelona Clinic Liver Cancer (BCLC) staging classification and treatment schedule. Patients with very early hepatocellular carcinoma (HCC) (stage 0) are optimal candidates for resection. Patients with early HCC (stage A) are candidates for radical therapy (resection, liver transplantation [LT], or local ablation via percutaneous ethanol injection [PEI] or radiofrequency [RF] ablation). Patients with intermediate HCC (stage B) benefit from transcatheter arterial chemoembolization (TACE). Patients with advanced HCC, defined as presence of macroscopic vascular invasion, extrahepatic spread, or cancer-related symptoms (Eastern Cooperative Oncology Group performance status 1 or 2) (stage C), benefit from sorafenib. Patients with end-stage disease (stage D) will receive symptomatic treatment. Treatment strategy will transition from one stage to another on treatment failure or contraindications for the procedures. CLT, cadaveric liver transplantation; LDLT, living donor liver transplantation; PST, Performance Status Test. (Modified from JM Llovet et al: *JNCI* 100:698, 2008.)

on prognosis and may be microscopic or macroscopic (visible on computed tomography [CT] scans). Most large tumors have microscopic vascular invasion, so full staging can usually be made only after surgical resection. Stage III disease contains a mixture of lymph node-positive and -negative tumors. Stage III patients with positive lymph node disease have a poor prognosis, and few patients survive 1 year. The prognosis of stage IV is poor after either resection or transplantation, and 1-year survival is rare.

New Directions Consensus is needed on staging. These systems will soon be refined or upended by proteomics.

APPROACH TO THE PATIENT: Hepatocellular Carcinoma

HISTORY AND PHYSICAL

The history is important in evaluating putative predisposing factors, including a history of hepatitis or jaundice, blood transfusion, or use of intravenous drugs. A family history of HCC or hepatitis should be sought and a detailed social history taken to include job descriptions for industrial exposure to possible carcinogenic drugs as well as contraceptive hormones. Physical examination should include assessing stigmata of underlying liver disease such as jaundice, ascites, peripheral edema, spider nevi, palmar erythema, and weight loss. Evaluation of the abdomen for hepatic size, masses or ascites, hepatic nodularity and tenderness, and splenomegaly is needed, as is assessment of overall performance status and psychosocial evaluation.

SEROLOGIC ASSAYS

AFP is a serum tumor marker for HCC; however, it is only increased in approximately one-half of U.S. patients. The lens culinaris agglutinin-reactive fraction of AFP (AFP-L3) assay is thought to be more specific. The other widely used assay is that for des- γ -carboxy prothrombin (DCP), a protein induced by vitamin K absence (PIVKA-2). This protein is increased in as many as 80% of HCC patients but may also be elevated in patients with vitamin K deficiency; it is always elevated after warfarin use. It may also predict for portal vein invasion. Both AFP-L3 and DCP are U.S. Food and Drug Administration (FDA) approved. Many other assays have been developed, such as glypican-3, but none have greater aggregate sensitivity and specificity. In a patient presenting with either a new hepatic mass or other indications of recent hepatic decompensation, carcinoembryonic antigen (CEA), vitamin B₁₂, AFP, ferritin, PIVKA-2, and antimitochondrial antibody should be measured, and standard liver function tests should be performed, including prothrombin time (PT), partial thromboplastin time (PTT), albumin, transaminases, γ -glutamyl transpeptidase, and alkaline phosphatase. γ -Glutamyl transpeptidase and alkaline phosphatase may be particularly important in the 50% of HCC patients who have low AFP levels. Decreases in platelet count and white blood cell count may reflect portal hypertension and associated hypersplenism. Hepatitis A, B, and C serology should be measured. If HBV or HCV serology is positive, quantitative measurements of HBV DNA or HCV RNA are needed.

New Directions Newer biomarkers are being evaluated, especially tissue- and serum-based genomics profiling. Newer plasma biomarkers include glypican-3, osteopontin, insulin-like growth factor I, and vascular endothelial growth factor. However, they are still in process of validation. Furthermore, the commercial availability of kits for isolating circulating tumor cells is permitting the molecular profiling of HCCs without the need for further tissue biopsy.

RADIOLOGY

An ultrasound examination of the liver is an excellent screening tool. The two characteristic vascular abnormalities are hypervascularity of the tumor mass (neovascularization or abnormal tumor-feeding arterial vessels) and thrombosis by tumor invasion

of otherwise normal portal veins. To determine tumor size and extent and the presence of portal vein invasion accurately, a helical/triphasic CT scan of the abdomen and pelvis, with fast-contrast bolus technique, should be performed to detect the vascular lesions typical of HCC. Portal vein invasion is normally detected as an obstruction and expansion of the vessel. A chest CT is used to exclude metastases. Magnetic resonance imaging (MRI) can also provide detailed information, especially with the newer contrast agents. Ethiodol (Lipiodol) is an ethiodized oil emulsion retained by liver tumors that can be delivered by hepatic artery injection (5–15 mL) for CT imaging 1 week later. For small tumors, Ethiodol injection is very helpful before biopsy because the histologic presence of the dye constitutes proof that the needle biopsied the mass under suspicion. A prospective comparison of triphasic CT, gadolinium-enhanced MRI, ultrasound, and fluorodeoxyglucose positron emission tomography (FDG-PET) showed similar results for CT, MRI, and ultrasound; PET imaging appears to be positive in only a subset of HCC patients. Abdominal CT versus MRI/CT uses a faster single breath-hold, is less complex, and is less dependent on patient cooperation. MRI requires a longer examination, and ascites can cause artifacts, but MRI is better able to distinguish dysplastic or regenerative nodules from HCC. Imaging criteria have been developed for HCC that do not require biopsy proof, as they have >90% specificity. The criteria include nodules >1 cm with arterial enhancement and portal venous washout and, for small tumors, specified growth rates on two scans performed less than 6 months apart (Organ Procurement and Transplant Network). Nevertheless, explant pathology after liver transplant for HCC has shown that ~20% of patients diagnosed without biopsy did not actually have a tumor.

New Directions The altered tumor vascularity that is a consequence of molecularly targeted therapies is the basis for newer imaging techniques including contrast-enhanced ultrasound (CEUS) and dynamic MRI.

PATHOLOGIC DIAGNOSIS

Histologic proof of the presence of HCC is obtained through a core liver biopsy of the liver mass under ultrasound guidance, as well as random biopsy of the underlying liver. Bleeding risk is increased compared to other cancers because (1) the tumors are hypervascular and (2) patients often have thrombocytopenia and decreased liver-dependent clotting factors. Bleeding risk is further increased in the presence of ascites. Tracking of tumor has an uncommon problem. Fine-needle aspirates can provide sufficient material for diagnosis of cancer, but core biopsies are preferred. Tissue architecture allows the distinction between HCC and adenocarcinoma. Laparoscopic approaches can also be used. For patients suspected of having portal vein involvement, a core biopsy of the portal vein may be performed safely. If positive, this is regarded as an exclusion criterion for transplantation for HCC.

New Directions Immunohistochemistry has become mainstream. Prognostic subgroupings are being defined based on growth signaling pathway proteins and genotyping strategies, including a prognostically significant five-gene profile score. Furthermore, molecular profiling of the underlying liver has provided evidence for a “field-effect” of cirrhosis in generating recurrent or new HCCs after primary resection. In addition, characteristics of HCC stem cells have been identified and include EpCAM, CD44, and CD90 expression, which may form the basis of stem cell therapeutic targeting strategies.

SCREENING HIGH-RISK POPULATIONS

There are two goals of screening, both in patients at increased risk for developing HCC, such as those with cirrhosis. The first goal is to detect smaller tumors that are potentially curable by ablation. The second goal is to enhance survival, compared with patients who were not diagnosed by surveillance. Evidence from Taiwan has shown a survival advantage to population screening in HBV-positive patients, and other evidence has shown its efficacy in diagnosis for HCV. Prospective studies in

548 high-risk populations showed that ultrasound was more sensitive than AFP elevations alone, although most practitioners request both tests at 6-month intervals for HBV and HCV carriers, especially in the presence of cirrhosis or worsening of liver function tests. However, an Italian study in patients with cirrhosis identified a yearly HCC incidence of 3% but showed no increase in the rate of detection of potentially curable tumors with aggressive screening. Prevention strategies including universal vaccination against hepatitis are more likely to be effective than screening efforts. Despite absence of formal guidelines, most practitioners obtain 6-month AFP and ultrasound (cheap and ubiquitous, even in poor countries) or CT (more sensitive, especially in overweight patients, but more costly) studies when following high-risk patients (HBV carriers, HCV cirrhosis, family history of HCC).

Current Directions Cost-benefit analysis is not yet convincing, even though screening is intuitively sound. However, studies from areas with high HBV carrier rates have shown a survival benefit for screening as a result of earlier stage at diagnosis. A definitive clinical trial on screening is unlikely, due to difficulties in obtaining informed consent for patients who are not to be screened. γ -Glutamyl transpeptidase appears useful for detecting small tumors.

PREVENTION

Prevention strategies can only be planned when the causes of a cancer are known or strongly suspected. This is true of few human cancers, with significant exceptions being smoking and lung cancer, papilloma virus and cancer of the cervix uteri, and cirrhosis of any cause or dietary contamination by aflatoxin B₁ for HCC. Aflatoxin B₁ is one of the most potent known chemical carcinogens and is a product of the *Aspergillus* mold that grows on peanuts and rice when stored in hot and humid climates. The obvious strategy is to refrigerate these foodstuffs when stored and to conduct surveillance programs for elevated aflatoxin B₁ levels, as happens in the United States, but not usually in Asia. HBV is commonly transmitted from mother to fetus in Asia (except Japan), and neonatal HBV vaccination programs have resulted in a big decrease in adolescent HBV and, thus, in predicted HCC rates. There are millions of HBV and HCV carriers (4 million with HCV in the United States) who are already infected. Nucleoside analogue-based chemoprevention

(entecavir) of HBV-mediated HCC in Japan resulted in a fivefold decrease in HCC incidence over 5 years in cirrhotic but not in non-cirrhotic HBV patients. More powerful and effective HCV therapies promise the possibility of prevention of HCV-based HCC in the future.

TREATMENT HEPATOCELLULAR CARCINOMA

Most HCC patients have two liver diseases, cirrhosis and HCC, each of which is an independent cause of death. The presence of cirrhosis usually places constraints on resection surgery, ablative therapies, and chemotherapy. Thus patient assessment and treatment planning have to take the severity of the nonmalignant liver disease into account. The clinical management choices for HCC can be complex (Fig. 111-2, Tables 111-5 and 111-6). The natural history of HCC is highly variable. Patients presenting with advanced tumors (vascular invasion, symptoms, extrahepatic spread) have a median survival of ~4 months, with or without treatment. Treatment results from the literature are difficult to interpret. Survival is not always a measure of the efficacy of therapy because of the adverse effects on survival of the underlying liver disease. A multidisciplinary team, including a hepatologist, interventional radiologist, surgical oncologist, resection surgeon, transplant surgeon, and medical oncologist, is important for the comprehensive management of HCC patients.

TNM STAGES I AND II HCC

Early-stage tumors are successfully treated using various techniques, including surgical resection, local ablation (thermal, radiofrequency [RFA], or microwave ablation [MWA]), and local injection therapies (Table 111-6). Because the majority of patients with HCC suffer from a field defect in the cirrhotic liver, they are at risk for subsequent multiple primary liver tumors. Many will also have significant underlying liver disease and may not tolerate major surgical loss of hepatic parenchyma, and they may be eligible for orthotopic liver transplant (OLT). Living related donor transplants have increased in popularity, resulting in absence of waiting for a transplant. An important principle in treating early-stage HCC in the nontransplant

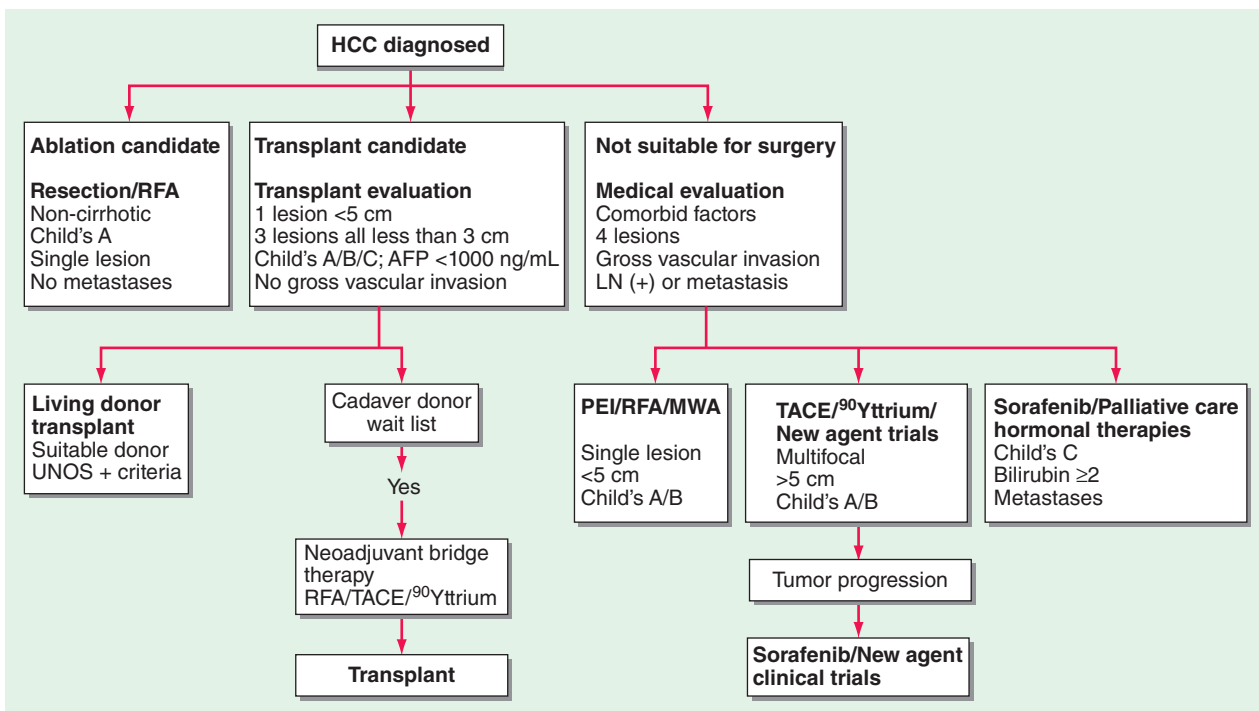


FIGURE 111-2 Hepatocellular carcinoma (HCC) treatment algorithm. The initial clinical evaluation is aimed at assessing the extent of the tumor and the underlying functional compromise of the liver by cirrhosis. Patients are classified as having resectable disease or unresectable disease or as being candidates for transplantation. AFP, α fetoprotein; LN, lymph node; MWA, microwave ablation; OLT, orthotopic liver transplantation; PEI, percutaneous ethanol injection; RFA, radiofrequency ablation; TACE, transcatheter arterial chemoembolization; UNOS, United Network for Organ Sharing. Child's A/B/C refers to the Child-Pugh classification of liver failure.

TABLE 111-5 TREATMENT OPTIONS FOR HEPATOCELLULAR CARCINOMA

Surgery	
Resection	
Liver transplantation	
Local Ablative Therapies	
Radiofrequency ablation (RFA)	
Microwave ablation (MWA)	
Cryosurgery	
Percutaneous ethanol injection (PEI)	
Regional Therapies: Hepatic Artery Transcatheter Treatments	
Transarterial chemotherapy	
Transarterial embolization	
Transarterial chemoembolization	
Transarterial drug-eluting beads	
Transarterial radiotherapies:	
⁹⁰ Yttrium microspheres	
¹³¹ Iodine–Ethiodol	
Proton beam radiation	
Conformal External-Beam Radiation and Intensity-Modulated Radiation Therapy	
Systemic therapies	
Molecularly targeted therapies (sorafenib, etc.)	
Chemotherapy	
Immunotherapy	
Hormonal therapy + growth control	
Supportive Therapies	

setting is to use liver-sparing treatments and to focus on treatment of both the tumor and the cirrhosis.

Surgical Excision The risk of major hepatectomy is high (5–10% mortality rate) due to the underlying liver disease and the potential for liver failure, but acceptable in selected cases and highly dependent on surgical experience. The risk is lower in high-volume centers. Preoperative portal vein occlusion can sometimes be performed to cause atrophy of the HCC-involved lobe and compensatory hypertrophy of the noninvolved liver, permitting safer resection. Intraoperative ultrasound is useful for planning the surgical approach. The ultrasound can image the proximity of major vascular structures that may be encountered during the dissection. In cirrhotic patients, any major liver surgery can result in liver failure. The Child-Pugh classification of liver failure is still a reliable prognosticator for tolerance of hepatic surgery, and only Child A patients should be considered for surgical resection. Child B and C

patients with stages I and II HCC should be referred for OLTX if appropriate, as well as patients with ascites or a recent history of variceal bleeding. Although open surgical excision is the most reliable, the patient may be better served with a laparoscopic approach to resection, using RFA, MWA, or percutaneous ethanol injection (PEI). No adequate comparisons of these different techniques have been undertaken, and the choice of treatment is usually based on physician skill. However, RFA has been shown to be superior to PEI in necrosis induction for tumors <3 cm in diameter and is thought to be equivalent to open resection and, thus, is the treatment of first choice for these small tumors. As tumors get larger than 3 cm, especially ≥5 cm, the effectiveness of RFA-induced necrosis diminishes. The combination of transcatheter arterial chemoembolization (TACE) with RFA has shown superior results to TACE alone in a prospective, randomized trial. Although vascular invasion is a preeminent negative prognostic factor, microvascular invasion in small tumors appears not to be a negative factor.

Local Ablation Strategies RFA uses heat to ablate tumors. The maximum size of the probe arrays allows for a 7-cm zone of necrosis, which would be adequate for a 3- to 4-cm tumor. The heat reliably kills cells within the zone of necrosis. Treatment of tumors close to the main portal pedicles can lead to bile duct injury and obstruction. This limits the location of tumors that are anatomically suited for this technique. RFA can be performed percutaneously with CT or ultrasound guidance, or at the time of laparoscopy with ultrasound guidance.

Local Injection Therapy Numerous agents have been used for local injection into tumors, most commonly ethanol (PEI). The relatively soft HCC within the hard background cirrhotic liver allows for injection of large volumes of ethanol into the tumor without diffusion into the hepatic parenchyma or leakage out of the liver. PEI causes direct destruction of cancer cells, but it is not selective for cancer and will destroy normal cells in the vicinity. However, it usually requires multiple injections (average three), in contrast to one for RFA. The maximum size of tumor reliably treated is 3 cm, even with multiple injections.

CURRENT DIRECTIONS Resection and RFA each obtain similar results. However, a distinction has been made between the causes and prevention strategies needed to prevent early versus late tumor recurrences after resection. Early recurrence has been linked to tumor invasion factors, especially microvascular tumor invasion with elevated transaminases, whereas late recurrence has been associated with cirrhosis and virus hepatitis factors and, thus, the development of new tumors. See the section on virus-directed adjuvant therapy below.

Liver Transplantation (OLT) A viable option for stages I and II tumors in the setting of cirrhosis is OLT, with survival approaching that for noncancer cases. OLT for patients with a single lesion ≤5 cm or three or fewer nodules, each ≤3 cm (Milan criteria), resulted in excellent tumor-free survival (≥70% at 5 years). For advanced HCC, OLT has been abandoned due to high tumor recurrence rates. Priority scoring for OLT previously led to HCC patients waiting too long for their OLT, resulting in some tumors becoming too advanced during the patient's wait for a donated liver. A variety of therapies were used as a “bridge” to OLT, including RFA, TACE, and hepatic arterial ⁹⁰Y-radioembolization. These pre-transplant treatments allow patients to remain on the waiting list longer, giving them greater opportunities to be transplanted, because they can stabilize the tumor and prevent it from

TABLE 111-6 SOME RANDOMIZED CLINICAL TRIALS INVOLVING TRANSHEPATIC ARTERY CHEMOEMBOLIZATION (TACE) FOR HEPATOCELLULAR CARCINOMA

Author	Year	Agents 1	Agents 2	Survival Effect
Kawaii	1992	Doxorubicin + Embo	Embo	No
Chang	1994	Cisplatin + Embo	Embo	No
Hatanaka	1995	Cisplatin, doxorubicin, + Embo	Same + Lipiodol	No
Uchino	1993	Cisplatin, doxorubicin, + oral FU	Same + Tamoxifen	No
Lin	1988	Embo	Embo + IV FU	No
Yoshikawa	1994	Epirubicin + Ethiodol	Epirubicin	No
Pelletier	1990	Doxorubicin + Gelfoam	None	No
Trinchet	1995	Cisplatin + Gelfoam	None	No
Bruix	1998	Coils + Gelfoam	None	No
Pelletier	1998	Cisplatin + Ethiodol	None	No
Trinchet	1995	Cisplatin + Gelfoam	None	No
Lo	2002	Cisplatin + Ethiodol	None	Yes
Llovet	2002	Doxorubicin + Ethiodol	None	Yes

Abbreviations: Embo, embolization; FU, 5-fluorouracil.

550 growing in the months until a donor liver becomes available. What remains unclear, however, is whether this translates into prolonged survival after transplant. Further, it is not known whether patients who have had their tumor(s) treated preoperatively follow the recurrence pattern predicted by their tumor status at the time of transplant (i.e., post-local ablative therapy), or if they follow the course set by their tumor parameters present before such treatment. The United Network for Organ Sharing (UNOS) point system for priority scoring of OLTX recipients now includes additional points for patients with HCC. The success of living related donor liver transplantation programs has also led to patients receiving transplantation earlier for HCC and often with greater than minimal tumors.

CURRENT DIRECTIONS Expanded criteria for larger HCCs beyond the Milan criteria (one lesion <5 cm or three lesions, each <3 cm), such as the University of California, San Francisco (UCSF) criteria (single lesion ≤6.5 cm or two lesions ≤4.5 cm with a total diameter ≤8 cm; 1- and 5-year survival rates of 90 and 75%, respectively), are being increasingly accepted by various UNOS areas for OLTX with satisfactory longer-term survival comparable to Milan criteria results. Furthermore, downstaging of HCCs that are too large for the Milan criteria by medical therapy (TACE) is increasingly recognized as acceptable treatment before OLTX with equivalent outcomes to patients who originally were within Milan criteria. Within-criteria patients with AFP levels >1000 ng/mL have exceptionally high post-OLTX recurrence rates. Also, the use of “salvage” OLTX after recurrent HCC after resection has produced conflicting outcomes. Shortages of organs combined with advances in resection safety have led to increasing use of resection for patients with good liver function.

Adjuvant Therapy The role of adjuvant chemotherapy for patients after resection or OLTX remains unclear. Both adjuvant and neoadjuvant approaches have been studied, but no clear advantage in disease-free or overall survival has been found. However, a meta-analysis of several trials revealed a significant improvement in disease-free and overall survival. Although analysis of postoperative adjuvant systemic chemotherapy trials demonstrated no disease-free or overall survival advantage, single studies of TACE and neoadjuvant ¹³¹I-Ethiodol showed enhanced survival after resection.

Antiviral therapy, instead of anticancer therapy, has been successful in decreasing postresection tumor recurrences in the postresection adjuvant setting. Nucleoside analogues in HBV-based HCC and peg-interferon plus ribavirin for HCV-based HCC have both been effective in reducing recurrence rates.

CURRENT DIRECTIONS A large adjuvant trial examining resection and transplantation, with or without sorafenib (see below) is in progress. The success of viral therapies in decreasing HCC recurrence after resection is part of a broader focus on the tumor microenvironment (stroma, blood vessels, inflammatory cells, and cytokines) as mediators of HCC progression and as targets for new therapies.

TNM STAGES III AND IV HCC

Fewer surgical options exist for stage III tumors involving major vascular structures. In patients without cirrhosis, a major hepatectomy is feasible, although prognosis is poor. Patients with Child A cirrhosis may be resected, but a lobectomy is associated with significant morbidity and mortality rates, and long-term prognosis is poor. Nevertheless, a small percentage of patients will achieve long-term survival, justifying an attempt at resection when feasible. Because of the advanced nature of these tumors, even successful resection can be followed by rapid recurrence. These patients are not considered candidates for transplantation because of the high tumor recurrence rates, unless their tumors can first be downstaged with neoadjuvant therapy. Decreasing the size of the primary tumor allows for less surgery, and the delay in surgery allows for extrahepatic disease to manifest on imaging studies and avoid unhelpful OLTX. The prognosis is poor for stage IV tumors, and no surgical treatment is recommended.

Systemic Chemotherapy A large number of controlled and uncontrolled clinical studies have been performed with most of the major

classes of cancer chemotherapy. No single agent or combination of agents given systemically reproducibly leads to even a 25% response rate or has any effect on survival.

Regional Chemotherapy In contrast to the dismal results of systemic chemotherapy, a variety of agents given via the hepatic artery have activity for HCC confined to the liver (Table 111-6). Two randomized controlled trials have shown a survival advantage for TACE in a selected subset of patients. One used doxorubicin, and the other used cisplatin. Despite the fact that increased hepatic extraction of chemotherapy has been shown for very few drugs, some drugs such as cisplatin, doxorubicin, mitomycin C, and possibly neocarzinostatin, produce substantial objective responses when administered regionally. Few data are available on continuous hepatic arterial infusion for HCC, although pilot studies with cisplatin have shown encouraging responses. Because the reports have not usually stratified responses or survival based on TNM staging, it is difficult to know long-term prognosis in relation to tumor extent. Most of the studies on regional hepatic arterial chemotherapy also use an embolizing agent such as Ethiodol, gelatin sponge particles (Gelfoam), starch (Spherex), or microspheres. Two products are composed of microspheres of defined size ranges—Embospheres (Biospheres) and Contour SE—using particles of 40–120, 100–300, 300–500, and 500–1000 μm in size. The optimal diameter of the particles for TACE has yet to be defined. Consistently higher objective response rates are reported for arterial administration of drugs together with some form of hepatic artery occlusion compared with any form of systemic chemotherapy to date. The widespread use of some form of embolization in addition to chemotherapy has added to its toxicities. These include a frequent but transient fever, abdominal pain, and anorexia (all in >60% of patients). In addition, >20% of patients have increased ascites or transient elevation of transaminases. Cystic artery spasm and cholecystitis are also not uncommon. However, higher responses have also been obtained. The hepatic toxicities associated with embolization may be ameliorated by the use of degradable starch microspheres, with 50–60% response rates. Two randomized studies of TACE versus placebo showed a survival advantage for treatment (Table 111-6). In addition, it is not clear that formal oncologic CT response criteria are adequate for HCC. A loss of vascularity on CT without size change may be an index of loss of viability and thus of response to TACE. A major problem that TACE trials have had in showing a survival advantage is that many HCC patients die of their underlying cirrhosis, not the tumor. Nevertheless, two randomized controlled trials, one using doxorubicin and the other using cisplatin, showed a survival advantage for TACE versus placebo (Table 111-6). However, improving quality of life is a legitimate goal of regional therapy. Drug-eluting beads using doxorubicin (DEB-TACE) have been claimed to produce equivalent survival with less toxicity, but this strategy has not been tested in a randomized trial.

Kinase Inhibitors A survival advantage has been observed for the oral multikinase inhibitor, sorafenib (Nexavar), versus placebo in two randomized trials. It targets both the Raf mitogenic pathway and the vascular endothelial growth factor receptor (VEGFR) endothelial vasculogenesis pathway. However, tumor responses were negligible, and the survival in the treatment arm in Asians was less than the placebo arm in the Western trial (Table 111-7). Sorafenib has considerable toxicity, with 30–40% of patients requiring “drug holidays,” dose reductions, or cessation of therapy. The most common toxicities include fatigue, hypertension, diarrhea, mucositis, and skin changes, such as the painful hand-foot syndrome, hair loss, and itching, each in 20–40% of patients. Several “look-alike” new agents that also target angiogenesis have either proved to be inferior or more toxic. These include sunitinib, brivanib, linifanib, everolimus, and bevacizumab (Table 111-8). The idea of angiogenesis alone as a major HCC therapeutic target may need revision.

New Therapies Although prolonged survival has been reported in phase II trials using newer agents, such as bevacizumab plus

TABLE 111-7 TARGETED THERAPIES IN HEPATOCELLULAR CARCINOMA: TRIALS

Phase III	Target	Survival (mo)
Sorafenib vs placebo	Raf, VEGFR, PDGFR	10.7 vs 7.9
Sorafenib vs placebo (Asians)	Raf, VEGFR, PDGFR	6.5 vs 4.2

Abbreviations: PDGFR, platelet-derived growth factor receptor; Raf, rapidly accelerated fibrosarcoma; VEGFR vascular endothelial growth factor receptor.

erlotinib, the data from a phase III trial were disappointing. Several forms of *radiation therapy* have been used in the treatment of HCC, including external-beam radiation and conformal radiation therapy. Radiation hepatitis remains a dose-limiting problem. The pure beta emitter ⁹⁰Yttrium attached to either glass (TheraSphere) or resin (SIR-Spheres) microspheres injected into a major branch hepatic artery has been assessed in phase II trials of HCC and has encouraging tumor control and survival effects with minimal toxicities. Randomized phase III trials comparing it to TACE have yet to be completed. The main attractiveness of ⁹⁰Yttrium therapy is its safety in the presence of major branch portal vein thrombosis, where TACE is dangerous or contraindicated. Furthermore, external-beam radiation has been reported to be safe and useful in the control of major branch portal or hepatic vein invasion (thrombosis) by tumors. The studies have all been small. Vitamin K has been assessed in clinical trials at high dosage for its HCC-inhibitory actions. This idea is based on the characteristic biochemical defect in HCC of elevated plasma levels of immature prothrombin (DCP or PIVKA-2), due to a defect in the activity of prothrombin carboxylase, a vitamin K-dependent enzyme. Two vitamin K randomized controlled trials from Japan show decreased tumor occurrence, but a major phase III trial aimed at limiting postresection recurrence was not successful.

CURRENT DIRECTIONS A number of new kinase inhibitors are being evaluated for HCC (Tables 111-9 and 111-10). These include the biologicals, such as Raf kinase and vascular endothelial growth factor (VEGF) inhibitors, and agents that target various steps of the cell growth pathway. Current hopes focus particularly on the Met pathway inhibitors such as tivantinib and several IGF receptor antagonists. ⁹⁰Yttrium looks promising and without chemotherapy toxicities. It is particularly attractive because, unlike TACE, it seems safe in the presence of portal vein thrombosis, a pathognomonic feature of HCC aggressiveness. The bottleneck of liver donors for OLTX is at last widening with increasing use of living donors, and criteria for OLTX for larger HCCs are slowly expanding. Patient participation in clinical trials assessing new therapies is encouraged (www.clinicaltrials.gov).

The main effort now is the evaluation of combinations of the compounds listed in Tables 111-7 to 111-9 that target different pathways, as well as the combination of any of these targeted therapies, but especially sorafenib, with TACE or ⁹⁰Yttrium radioembolization. Combining TACE with sorafenib appears to be safe in phase II studies with promising survival data, but randomized studies are still in progress. The same is true for intra-arterial ⁹⁰Yttrium plus sorafenib as therapy for HCC and as bridge to transplant therapy.

TABLE 111-8 PROMISING TARGETED THERAPIES THAT FAILED THEIR CLINICAL TRIAL GOALS

Sunitinib
Brivanib
Linifanib
Everolimus
Erlotinib
ThermaDox
Oncolytic virus JX-594
Bevacizumab
Bevacizumab plus erlotinib vs sorafenib
Sorafenib plus erlotinib vs sorafenib

TABLE 111-9 NEW TARGETED AGENTS AND THEIR TARGETS IN CURRENT CLINICAL TRIALS

Targets	Inhibitors
EGF receptor	Erlotinib
	Gefitinib
	Cetuximab
	Panitumumab
cMET	Tivantinib (ARQ197)
	EMD1204831
	Cabozantinib
VEGF receptor	Bevacizumab
	Regorafenib
	Brivanib
	Cediranib
	Sunitinib
FGF1 receptor	AEW54
	R1507 (MAb)
	Linsitinib (OSI-906)
	Brivanib
TRAIL-R1 (proapoptosis)	Mapatumumab
PDGF receptor	Sorafenib
	Dovitinib
	Linifanib
IGF-I receptor	IMC-A12
	B11B022
	Cixutumumab
Ubiquitin-proteasome	Bortezomib

Abbreviations: EGF, epidermal growth factor; FGF1, fibroblast growth factor 1; IGF-I, insulin-like growth factor I; PDGF, platelet-derived growth factor; VEGF, vascular endothelial growth factor.

SIGNIFICANCE AND EVALUATION OF RESPONSES TO NONSURGICAL THERAPIES

Tumor growth or spread is considered a poor prognostic sign and evidence of treatment failure. By contrast, patients receiving chemotherapy are judged to have a response if there is shrinkage of tumor size. Lack of response/size decrease has been thought of as treatment failure. Three considerations in HCC management have completely changed the views concerning nonshrinkage after therapy. First, the correlation between response to chemotherapy and survival is poor in various tumors; in some tumors, such as ovarian cancer and small-cell lung cancer, substantial tumor shrinkage on chemotherapy is followed by rapid

TABLE 111-10 SOME NOVEL MEDICAL TREATMENTS FOR HEPATOCELLULAR CARCINOMA

EGF receptor antagonists: erlotinib, gefitinib, lapatinib, cetuximab, brivanib
Multikinase antagonists: sorafenib, sunitinib
VEGF antagonist: bevacizumab
VEGFR antagonist: ABT-869 (linifanib)
mTOR antagonists: sirolimus, temsirolimus, everolimus
Proteasome inhibitors: bortezomib
Vitamin K
¹³¹ I-Ethiodol (lipiodol)
¹³¹ I-Ferritin
⁹⁰ Yttrium microspheres (TheraSphere, SIR-Spheres)
¹⁶⁶ Holmium, ¹⁸⁸ Rhenium
Three-dimensional conformal radiation
Proton beam high-dose radiotherapy
Gamma knife, CyberKnife
New targets: inhibitors of cyclin dependent kinases (Cdk), TRAIL induction caspases, and stem cells

Abbreviations: EGF, epidermal growth factor; mTOR, mammalian target of rapamycin; VEGF, vascular endothelial growth factor; VEGFR, vascular endothelial growth factor receptor.

552 tumor regrowth. Second, the Sorafenib HCC Assessment Randomized Protocol (SHARP) phase III trial of sorafenib versus placebo for unresectable HCC showed that survival could be significantly enhanced in the treatment arm with only 2% of the patients having tumor response but 70% of patients having disease stabilization. This observation has led to a reconsideration of the usefulness of response and the significance of disease stability. Third, HCC is a typically highly vascular tumor, and the vascularity is considered to be a measure of tumor viability. As a result, the Response Evaluation Criteria in Solid Tumors (RECIST) have been modified to mRECIST, which requires measurement of vascular/viable tumor on the CT or MRI scan. A partial response is defined as a 30% decrease in the sum of diameters of viable (arterially enhancing) target tumors. The need for semiquantitation of tumor vascularity on scans has led to the introduction of diffusion-weighted MRI imaging. Tissue-specific imaging agents such as gadoxetic acid (Primovist or Eovist) and the move to functional and genetic imaging mark a shift in approaches. Furthermore, plasma AFP response may be a biologic marker of radiologic response.

TREATMENT SUMMARY

Long-term survival is associated with resection or ablation or transplantation, all of which can yield >70% 5-year survival. Liver transplant is the only therapy that can treat the tumor and the underlying liver disease simultaneously and may be the most important advance in HCC therapy in 50 years. Unfortunately, it benefits only patients with limited size tumors without macrovascular portal vein invasion. Untreated patients with multinodular asymptomatic tumors without vascular invasion or extrahepatic spread have a median survival of approximately 16 months. Chemoembolization (TACE) improves their median survival to 19–20 months and is considered standard therapy for these patients, who represent the majority of HCC patients, although ⁹⁰Yttrium therapy may provide similar results with less toxicity. Patients with advanced-stage disease, vascular invasion, or metastases have a median survival of around 6 months. Among this group, outcomes may vary according to their underlying liver disease. It is this group at which kinase inhibitors are directed.

SUMMARY (TABLE 111-5)

The Most Common Modes of Patient Presentation

1. A patient with known history of hepatitis, jaundice, or cirrhosis, with an abnormality on ultrasound or CT scan, or rising AFP or DCP (PIVKA-2)
2. A patient with an abnormal liver function test as part of a routine examination
3. Radiologic workup for liver transplant for cirrhosis
4. Symptoms of HCC including cachexia, abdominal pain, or fever

History and Physical Examination

1. Clinical jaundice, asthenia, itching (scratches), tremors, or disorientation
2. Hepatomegaly, splenomegaly, ascites, peripheral edema, skin signs of liver failure

Clinical Evaluation

1. Blood tests: full blood count (splenomegaly), liver function tests, ammonia levels, electrolytes, AFP and DCP (PIVKA-2), Ca²⁺ and Mg²⁺; hepatitis B, C, and D serology (and quantitative HBV DNA or HCV RNA, if either is positive); neurotensin (specific for fibrolamellar HCC)
2. Triphasic dynamic helical (spiral) CT scan of liver (if inadequate, then follow with an MRI); chest CT scan; upper and lower gastrointestinal endoscopy (for varices, bleeding, ulcers); and brain scan (only if symptoms suggest)
3. Core biopsy: of the tumor and separate biopsy of the underlying liver

Therapy (Tables 111-5 and 111-6)

1. HCC <2 cm: RFA, PEI, or resection
2. HCC >2 cm, no vascular invasion: liver resection, RFA, or OLTX

3. Multiple unilobar tumors or tumor with vascular invasion: TACE or sorafenib
4. Bilobar tumors, no vascular invasion: TACE with OLTX for patients with tumor response
5. Extrahepatic HCC or elevated bilirubin: sorafenib or bevacizumab plus erlotinib (combination agent trials are in progress)

OTHER PRIMARY LIVER TUMORS

FIBROLAMELLAR HCC (FL-HCC)

This rarer variant of HCC has a quite different biology than adult-type HCC. None of the known HCC causative factors seem important here. It is typically a disease of younger adults, often teenagers and predominantly females. It is AFP-negative, but patients typically have elevated blood neurotensin levels, normal liver function tests, and no cirrhosis. Radiology is similar for HCC, except that characteristic adult-type portal vein invasion is less common. Although it is often multifocal in the liver, and therefore not resectable, metastases are common, especially to lungs and locoregional lymph nodes, but survival is often much better than with adult-type HCC. Resectable tumors are associated with 5-year survival $\geq 50\%$. Patients often present with a huge liver or unexplained weight loss, fever, or elevated liver function tests on routine evaluations. These huge masses suggest quite slow growth for many tumors. Surgical resection is the best management option, even for metastases, as these tumors respond much less well to chemotherapy than adult-type HCC. Although several series of OLTX for FL-HCC have been reported, the patients seem to die from tumor recurrences, with a 2- to 5-year lag compared with OLTX for adult-type HCC. Anecdotal responses to gemcitabine plus cisplatin-TACE are reported.

Epithelioid Hemangioendothelioma (EHE) This rare vascular tumor of adults is also usually multifocal and can also be associated with prolonged survival, even in the presence of metastases, which are commonly in the lung. There is usually no underlying cirrhosis. Histologically, these tumors are usually of borderline malignancy and express factor VIII, confirming their endothelial origin. OLTX may produce prolonged survival.

Cholangiocarcinoma (CCC) CCC typically refers to mucin-producing adenocarcinomas (different from HCC) that arise from the biliary tract and have features of cholangiocyte differentiation. They are grouped by their anatomic site of origin, as intrahepatic (IHC), perihilar (central, ~65% of CCCs), and peripheral (or distal, ~30% of CCCs). IHC is the second most common primary liver tumor. Depending on the site of origin, they have different features and require different treatments. They arise on the basis of cirrhosis less frequently than HCC, but may complicate primary biliary cirrhosis. However, cirrhosis and both primary biliary cirrhosis and HCV predispose to IHC. Nodular tumors arising at the bifurcation of the common bile duct are called *Klatskin* tumors and are often associated with a collapsed gallbladder, a finding that mandates visualization of the entire biliary tree. The approach to management of central and peripheral CCC is quite different. Incidence is increasing. Although most CCCs have no obvious cause (etiology unknown), a number of predisposing factors have been identified. Predisposing diseases include primary sclerosing cholangitis (10–20% of primary sclerosing cholangitis [PSC] patients), an autoimmune disease, and liver fluke in Asians, especially *Opisthorchis viverrini* and *Clonorchis sinensis*. CCC seems also to be associated with any cause of chronic biliary inflammation and injury, with alcoholic liver disease, choledocholithiasis, choledochal cysts (10%), and Caroli's disease (a rare inherited form of bile duct ectasia). CCC most typically presents as painless jaundice, often with pruritus or weight loss. Diagnosis is made by biopsy, percutaneously for peripheral liver lesions, or more commonly via endoscopic retrograde cholangiopancreatography (ERCP) under direct vision for central lesions. The tumors often stain positively for cytokeratins 7, 8, and 19 and negatively for cytokeratin 20. However, histology alone cannot usually distinguish CCC from metastases from colon or pancreas primary tumors. Serologic tumor markers appear to be nonspecific, but CEA, CA 19-9, and CA-125 are often elevated in CCC patients and are useful for following response to

therapy. Radiologic evaluation typically starts with ultrasound, which is very useful in visualizing dilated bile ducts, and then proceeds with either MRI or magnetic resonance cholangiopancreatography (MRCP) or helical CT scans. Invasive cholangiopancreatography (ERCP) is then needed to define the biliary tree and obtain a biopsy or is needed therapeutically to decompress an obstructed biliary tree with internal stent placement. If that fails, then percutaneous biliary drainage will be needed, with the biliary drainage flowing into an external bag. Central tumors often invade the porta hepatis, and locoregional lymph node involvement by tumor is frequent. Incidence has been increasing in recent decades; few patients survive 5 years. The usual treatment is surgical, but combination systemic chemotherapy may be effective. After complete surgical resection for IHC, 5-year survival is 25–30%. Combination radiation therapy with liver transplant has produced a 5-year recurrence-free survival rate of 65%.

TREATMENT CHOLANGIOCARCINOMA

Hilar CCC is resectable in ~30% of patients and usually involves bile duct resection and lymphadenectomy for prognostication. Typical survival is approximately 24 months, with recurrences being mainly in the operative bed but with ~30% in the lungs and liver. Distal CCC, which involves the main ducts, is normally treated by resection of the extrahepatic bile ducts, often with pancreaticoduodenectomy. Survival is similar. Due to the high rates of locoregional recurrences or positive surgical margins, many patients receive postoperative adjuvant radiotherapy. Its effect on survival has not been assessed. Intraluminal brachyradiotherapy has also shown some promise. However, photodynamic therapy enhanced survival in one study. In this technique, sodium porfimer is injected intravenously and then subjected to intraluminal red light laser photoactivation. OLTX has been assessed for treatment of unresectable CCC. Five-year survival was ~20%, so enthusiasm waned. However, neoadjuvant radiotherapy with sensitizing chemotherapy has shown better survival rates for CCC treated by OLTX and is currently used by UNOS for perihilar CCC <3 cm with neither intrahepatic or extrahepatic metastases. A 12-center data collection study of 287 patients with perihilar CCC confirmed the benefit of this approach in a subset of patients, with a 53% 5-year survival rate but with 10% patient dropout before transplantation. The patients had neoadjuvant external radiation with radiosensitizing therapy. Patients with tumors >3 cm had significantly shorter survival. Multiple chemotherapeutic agents have been assessed for activity and survival in unresectable CCC. Most have been inactive. However, both systemic and hepatic arterial gemcitabine have shown promising results. The combination of cisplatin plus gemcitabine has produced a survival advantage compared with gemcitabine alone in a 410-patient randomized controlled phase III trial for patients with locally advanced or metastatic CCC and is now considered standard therapy for unresectable CCC. Median overall survival in the combination arm was 11.7 months versus 8.1 months for gemcitabine alone. Significant responses were seen mainly in patients with IHC and gallbladder cancer. However, neither surgery for lymph node–positive disease nor regional chemotherapy in nonsurgical patients has shown any survival advantage thus far. Several case series have shown safety and some responses for hepatic arterial chemotherapy with gemcitabine, drug-eluting beads, and ⁹⁰Yttrium microspheres, but no convincing clinical trials are available. Clinical trials are under way with targeted therapies. Bevacizumab plus erlotinib gave a 10% partial response rate with median overall survival of 9.9 months. A sorafenib trial yielded an overall survival of 4.4 months, but 50% of the patients had received previous chemotherapy. Patients with unresectable tumors should be treated in clinical trials.

GALLBLADDER CANCER

Gallbladder (GB) cancer has an even worse prognosis than CCC, with a typical survival of ~6 months or less. Women are affected much more commonly than men (4:1), unlike HCC or CCC, and GB cancer occurs

more frequently than CCC. Most patients have a history of antecedent gallstones, but very few patients with gallstones develop GB cancer (~0.2%). GB cancer presents similarly to CCC and is often diagnosed unexpectedly during gallstone or cholecystitis surgery. Presentation is typically that of chronic cholecystitis, chronic right upper quadrant pain, and weight loss. Useful but nonspecific serum markers include CEA and CA 19-9. CT scans or MRCP typically reveal a GB mass. The mainstay of treatment is surgical, either simple or radical cholecystectomy for stage I or II disease, respectively. Survival rates are near 100% at 5 years for stage I, and range from 60–90% at 5 years for stage II. More advanced GB cancer has worse survival, and many patients are unresectable. Adjuvant radiotherapy, used in the presence of local lymph node disease, has not been shown to enhance survival. Chemotherapy is not useful in advanced or metastatic GB cancer.

CARCINOMA OF THE AMPULLA OF VATER

This tumor arises within 2 cm of the distal end of the common bile duct and is mainly (90%) an adenocarcinoma. Locoregional lymph nodes are commonly involved (50%), and the liver is the most frequent site for metastases. The most common clinical presentation is jaundice, and many patients also have pruritus, weight loss, and epigastric pain. Initial evaluation is performed with an abdominal ultrasound to assess vascular involvement, biliary dilation, and liver lesions. This is followed by a CT scan or MRI and especially MRCP. The most effective therapy is resection by pylorus-sparing pancreaticoduodenectomy, an aggressive procedure resulting in better survival rates than with local resection. Survival rates are ~25% at 5 years in operable patients with involved lymph nodes and ~50% in patients without involved nodes. Unlike CCC, approximately 80% of patients are thought to be resectable at diagnosis. Adjuvant chemotherapy or radiotherapy has not been shown to enhance survival. For metastatic tumors, chemotherapy is currently experimental.

TUMORS METASTATIC TO THE LIVER

These are predominantly from colon, pancreas, and breast primary tumors but can originate from any organ primary. Ocular melanomas are prone to liver metastasis. Tumor spread to the liver normally carries a poor prognosis for that tumor type. Colorectal and breast hepatic metastases were previously treated with continuous hepatic arterial infusion chemotherapy. However, more effective systemic drugs for each of these two cancers, especially the addition of oxaliplatin to colorectal cancer regimens, have reduced the use of hepatic artery infusion therapy. In a large randomized study of systemic versus infusional plus systemic chemotherapy for resected colorectal metastases to the liver, the patients receiving infusional therapy had no survival advantage, mainly due to extrahepatic tumor spread. ⁹⁰Yttrium resin beads are approved in the United States for treatment of colorectal hepatic metastases. The role of this modality, either alone or in combination with chemotherapy, is being evaluated in many centers. Palliation may be obtained from chemoembolization, PEI, or RFA.

BENIGN LIVER TUMORS

Three common benign tumors occur and all are found predominantly in women. They are *hemangiomas*, *adenomas*, and *focal nodular hyperplasia* (FNH). FNH is typically benign, and usually no treatment is needed. Hemangiomas are the most common and are entirely benign. Treatment is unnecessary unless their expansion causes symptoms. Adenomas are associated with contraceptive hormone use. They can cause pain and can bleed or rupture, causing acute problems. Their main interest for the physician is a low potential for malignant change and a 30% risk of bleeding. For this reason, considerable effort has gone into differentiating these three entities radiologically. On discovery of a liver mass, patients are usually advised to stop taking sex steroids, because adenoma regression may then occasionally occur. Adenomas can often be large masses ranging from 8–15 cm. Due to their size and definite, but low, malignant potential and potential for bleeding, adenomas are typically resected. The most useful diagnostic differentiating tool is a triphasic CT scan performed with HCC fast bolus protocol for arterial-phase imaging, together with subsequent delayed venous-phase imaging. Adenomas usually do not appear on

554 the basis of cirrhosis, although both adenomas and HCCs are intensely vascular on the CT arterial phase and both can exhibit hemorrhage (40% of adenomas). However, adenomas have smooth, well-defined edges, and enhance homogeneously, especially in the portal venous phase on delayed images, when HCCs no longer enhance. FNHs exhibit a characteristic central scar that is hypovascular on the arterial-phase and hypervascular on the delayed-phase CT images. MRI is even more sensitive in depicting the characteristic central scar of FNH.

112 Pancreatic Cancer

Elizabeth Smyth, David Cunningham

Pancreatic cancer is the fourth leading cause of cancer death in the United States and is associated with a poor prognosis. Endocrine tumors affecting the pancreas are discussed in [Chap. 113](#). Infiltrating ductal adenocarcinomas, the subject of this Chapter, account for the vast majority of cases and arise most frequently in the head of pancreas. At the time of diagnosis, 85–90% of patients have inoperable or metastatic disease, which is reflected in the 5-year survival rate of only 6% for all stages combined. An improved 5-year survival of up to 24% may be achieved when the tumor is detected at an early stage and when complete surgical resection is accomplished.

EPIDEMIOLOGY

Pancreatic cancer represents 3% of all newly diagnosed malignancies in the United States. The most common age group at diagnosis is 65–84 years for both sexes. Pancreatic cancer was estimated to have been diagnosed in approximately 45,220 patients and accounted for approximately 38,460 deaths in 2013. Although survival rates have almost doubled over the past 35 years for this disease, overall survival remains low.

GLOBAL CONSIDERATIONS



An estimated 278,684 cases of pancreatic cancer occur annually worldwide (the thirteenth most common cancer globally), with up to 60% of these cases diagnosed in more developed countries. It remains the eighth most common cause of cancer death in men and the ninth most common in women. The incidence is highest in the United States and western Europe and lowest in parts of Africa and South Central Asia. However, increasing rates of obesity, diabetes, and tobacco use in addition to access to diagnostic radiology in the developing world are likely to increase incidence rates in these countries. In this situation, consideration of the cost implications of adoption of current treatment paradigms in resource-constrained environments will be necessary. Primary prevention such as limiting tobacco use and avoiding obesity may be more cost effective than improvements in treatment of preexisting disease.

RISK FACTORS

Cigarette smoking may be the cause of up to 20–25% of all pancreatic cancers and is the most common environmental risk factor for this disease. A longstanding history of type 1 or type 2 diabetes also appears to be a risk factor; however, diabetes may also occur in association with pancreatic cancer, possibly confounding this interpretation. Other risk factors may include obesity, chronic pancreatitis, and ABO blood group status. Alcohol does not appear to be a risk factor unless excess consumption gives rise to chronic pancreatitis.

GENETIC AND MOLECULAR CONSIDERATIONS



Pancreatic cancer is associated with a number of well-defined molecular hallmarks. The four genes most commonly mutated or inactivated in pancreatic cancer are *KRAS* (predominantly codon 12, in 60–75% of pancreatic cancers), the tumor-suppressor genes *p16* (deleted in 95% of tumors), *p53* (inactivated or mutated in 50–70% of tumors), and *SMAD4* (deleted in 55% of tumors). The pancreatic

cancer precursor lesion pancreatic intraepithelial neoplasia (PanIN) acquires these genetic abnormalities in a progressive manner associated with increasing dysplasia; initial *KRAS* mutations are followed by *p16* loss and finally *p53* and *SMAD4* alterations. *SMAD4* gene inactivation is associated with a pattern of widespread metastatic disease in advanced-stage patients and poorer survival in patients with surgically resected pancreatic adenocarcinoma.

Up to 16% of pancreatic cancers may be inherited. Germline mutations in the following genes are associated with a significantly increased risk of pancreatic cancer and other cancers: (1) *STK11* gene (Peutz-Jeghers syndrome), which carries a 132-fold increased lifetime risk of pancreatic cancer above the general population; (2) *BRCA2* (increased risk of breast, ovarian, and pancreatic cancer); (3) *p16/CDKN2A* (familial atypical multiple mole melanoma), which carries an increased risk of melanoma and pancreatic cancer; (4) *PALB2*, which confers an increased risk of breast and pancreatic cancer; (5) *hMLH1* and *MSH2* (Lynch syndrome), which carries an increased risk of colon and pancreatic cancer; and (6) *ATM* (ataxia-telangiectasia), which carries an increased risk of breast cancer, lymphoma, and pancreatic cancer. Familial pancreatitis and an increased risk of pancreatic cancer are associated with mutations of the *PRSS1* (serine protease 1) gene. However, for most familial pancreatic syndromes, the underlying genetic cause remains unexplained. The absolute number of affected first-degree relatives is also correlated with increased cancer risk, and patients with at least two first-degree relatives with pancreatic cancer should be considered to have familial pancreatic cancer until proven otherwise.

The desmoplastic stroma surrounding pancreatic adenocarcinoma functions as a mechanical barrier to chemotherapy and secretes compounds essential for tumor progression and metastasis. Key mediators of these functions include the activated pancreatic stellate cell and the glycoprotein SPARC (secreted protein acidic and rich in cysteine), which is expressed in 80% of pancreatic ductal adenocarcinomas. Targeting this extracellular environment has become increasingly important in the treatment of advanced disease.

SCREENING AND PRECURSOR LESIONS

Screening is not routinely recommended because the incidence of pancreatic cancer in the general population is low (lifetime risk 1.3%), putative tumor markers such as carbohydrate antigen 19-9 (CA19-9) and carcinoembryonic antigen (CEA) have insufficient sensitivity, and computed tomography (CT) has inadequate resolution to detect pancreatic dysplasia. Endoscopic ultrasound (EUS) is a more promising screening tool, and preclinical efforts are focused on identifying biomarkers that may detect pancreatic cancer at an early stage. Consensus practice recommendations based largely on expert opinion have chosen a threshold of greater than fivefold increased risk for developing pancreatic cancer to select individuals who may benefit from screening. This includes people with two or more first-degree relatives with pancreatic cancer, patients with Peutz-Jeghers syndrome, and *BRCA 2*, *p16*, and hereditary nonpolyposis colorectal cancer (HNPCC) mutation carriers with one or more affected first-degree relatives.

PanIN represents a spectrum of small (<5 mm) neoplastic but non-invasive precursor lesions of the pancreatic ductal epithelium demonstrating mild, moderate, or severe dysplasia (PanIN 1–3, respectively); however, not all PanIN lesions will progress to frank invasive malignancy. Cystic pancreatic tumors such as intraductal mucinous papillary neoplasms (IPMNs) and mucinous cystic neoplasms (MCNs) are increasingly detected radiologically and are frequently asymptomatic. Main duct IPMNs are more likely to occur in older persons and have higher malignant potential than branched duct IPMNs (invasive cancer in 45% vs 18% of resected lesions, respectively). In contrast, MCNs are solitary lesions of the distal pancreas that do not communicate with the duct system. MCNs have an almost exclusive female distribution (95%). The rate of invasive cancer in resected MCNs is lower (<18%) with increased rates associated with larger tumors or the presence of nodules.

CLINICAL FEATURES

Clinical Presentation Obstructive jaundice occurs frequently when the cancer is located in the head of the pancreas. This may be accompanied

by symptoms of abdominal discomfort, pruritus, lethargy, and weight loss. Less common presenting features include epigastric pain, backache, new-onset diabetes mellitus, and acute pancreatitis caused by pressure effects on the pancreatic duct. Nausea and vomiting, resulting from gastroduodenal obstruction, may also be a symptom of this disease.

Physical Signs Patients can present with jaundice and cachexia, and scratch marks may be present. Of patients with operable tumors, 25% have a palpable gallbladder (Courvoisier's sign). Physical signs related to the development of distant metastases include hepatomegaly, ascites, left supraclavicular lymphadenopathy (Virchow's node), and periumbilical nodules (Sister Mary Joseph's nodes).

DIAGNOSIS

Diagnostic Imaging Patients who present with clinical features suggestive of pancreatic cancer undergo imaging to confirm the presence of a tumor and to establish whether the mass is likely to be inflammatory or malignant in nature. Other imaging objectives include the local and distant staging of the tumor, which will determine resectability and provide prognostic information. Dual-phase, contrast-enhanced spiral CT is the imaging modality of choice (Fig. 112-1). It provides accurate visualization of surrounding viscera, vessels, and lymph nodes, thus determining tumor resectability. Intestinal infiltration and liver and lung metastases are also reliably depicted on CT. There is no advantage of magnetic resonance imaging (MRI) over CT in predicting tumor resectability, but selected cases may benefit from MRI to characterize the nature of small indeterminate liver lesions and to evaluate the cause of biliary dilatation when no obvious mass is seen on CT. Endoscopic retrograde cholangiopancreatography (ERCP) is useful for revealing small pancreatic lesions, identifying stricture or obstruction in pancreatic or common bile ducts, and facilitating stent placement; however, it is associated with a risk of pancreatitis (Fig. 112-2). Magnetic resonance cholangiopancreatography (MRCP) is a noninvasive method for accurately depicting the level and degree of bile and pancreatic duct dilatation. EUS is highly sensitive in detecting lesions less than 3 cm in size (more sensitive than CT for lesions <2 cm) and is useful as a local staging tool for assessing vascular invasion and lymph node involvement. Fluorodeoxyglucose positron emission tomography (FDG-PET) should be considered before surgery or radical chemoradiotherapy (CRT), because it is superior to conventional imaging in detecting distant metastases.

Tissue Diagnosis and Cytology Preoperative confirmation of malignancy is not always necessary in patients with radiologic appearances consistent with operable pancreatic cancer. However, EUS-guided fine-needle aspiration is the technique of choice when there is any doubt, and also for use in patients who require neoadjuvant treatment. It has an accuracy of approximately 90% and has a smaller risk of intraperitoneal dissemination compared with the percutaneous route.

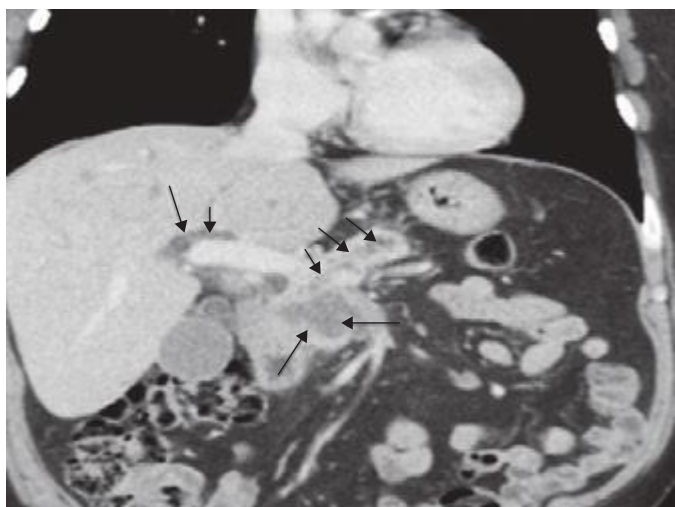


FIGURE 112-1 Coronal computed tomography showing pancreatic cancer and dilated intrahepatic and pancreatic ducts (arrows).

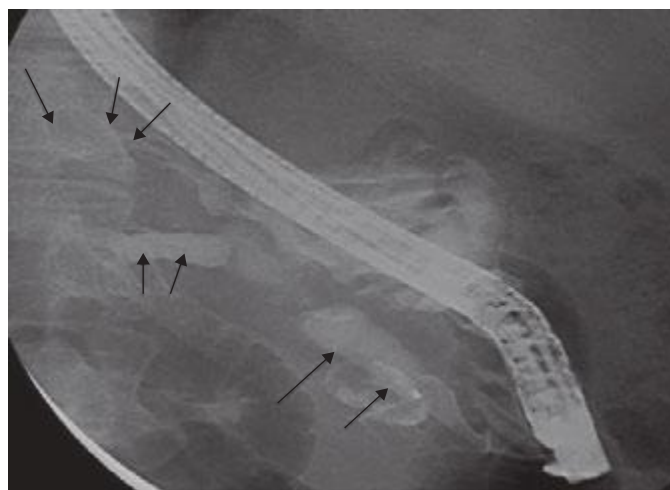


FIGURE 112-2 Endoscopic retrograde cholangiopancreatography showing contrast in dilated pancreatic duct (arrows).

Percutaneous biopsy of the pancreatic primary or liver metastases is only acceptable in patients with inoperable or metastatic disease. ERCP is a useful method for obtaining ductal brushings, but the sensitivity of ERCP for diagnosis ranges from 35 to 70%.

Serum Markers Tumor-associated CA19-9 is elevated in approximately 70–80% of patients with pancreatic carcinoma but is not recommended as a routine diagnostic or screening test because its sensitivity and specificity are inadequate for accurate diagnosis. Preoperative CA19-9 levels correlate with tumor stage, and postresection CA19-9 level has prognostic value. It is an indicator of asymptomatic recurrence in patients with completely resected tumors and is used as a biomarker of response in patients with advanced disease undergoing chemotherapy. A number of studies have established a high pretreatment CA19-9 level as an independent prognostic factor.

STAGING

The American Joint Committee on Cancer (AJCC) tumor-node-metastasis (TNM) staging of pancreatic cancer takes into account the location and size of the tumor, the involvement of lymph nodes, and distant metastasis. This information is then combined to assign a stage (Fig. 112-3). From a practical standpoint, patients are grouped according to whether the cancer is resectable, locally advanced (unresectable, but without distant spread), or metastatic.

TREATMENT PANCREATIC CANCER

RESECTABLE DISEASE

Approximately 10% of patients present with localized nonmetastatic disease that is potentially suitable for surgical resection. Approximately 30% of patients have R1 resection (microscopic residual disease) following surgery. Those who undergo R0 resection (no microscopic or macroscopic residual tumor) and who receive adjuvant treatment have the best chance of cure, with an estimated median survival of 20–23 months and a 5-year survival of approximately 20%. Outcomes are more favorable in patients with small (<3 cm), well-differentiated tumors and lymph node–negative disease.

Patients should have surgery in dedicated pancreatic centers that have lower postoperative morbidity and mortality rates. The standard surgical procedure for patients with tumors of the pancreatic head or uncinate process is a pylorus-preserving pancreaticoduodenectomy (modified Whipple's procedure). The procedure of choice for tumors of the pancreatic body and tail is a distal pancreatectomy, which routinely includes splenectomy.

Postoperative treatment improves long-term outcomes in this group of patients. Adjuvant chemotherapy comprising six cycles of gemcitabine is common practice worldwide based on data from three randomized controlled trials (Table 112-1). The Charité

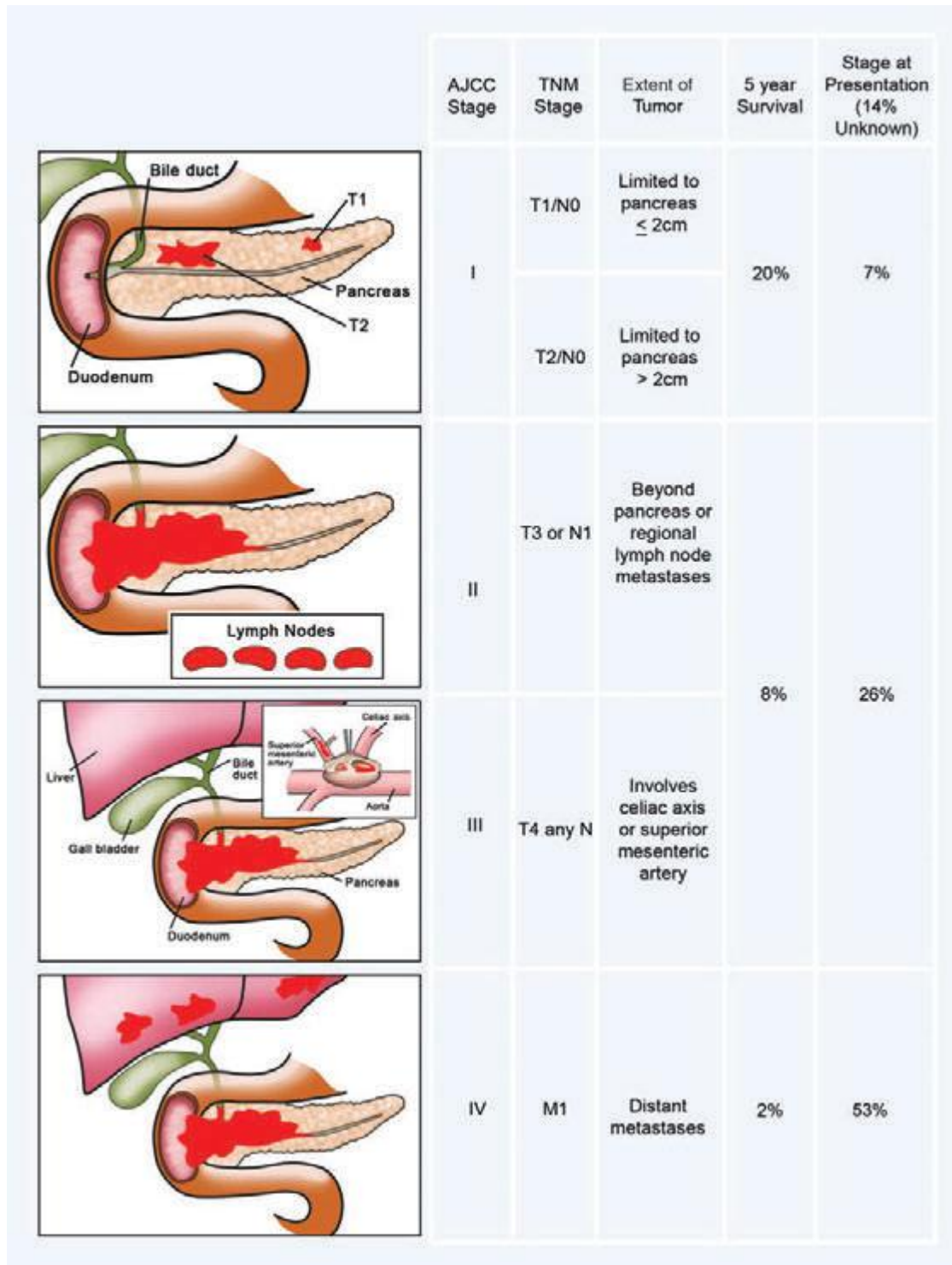


FIGURE 112-3 Staging of pancreatic cancer, and survival according to stage. AJCC, American Joint Committee on Cancer. (Illustration by Stephen Millward.)

Study	Comparator Arm	No. of Patients	Survival	
			PFS/DFS (months)	Median Survival (months)
ESPAC-1, Neoptolemos et al: N Engl J Med 350:1200, 2004	Chemotherapy (folinic acid + bolus 5-FU) vs no chemotherapy	289	PFS 15.3 vs 9.4. ($p = .02$)	20.1 vs 15.5 (HR 0.71; 95% CI 0.55–0.92; $p = .009$)
CONKO 001, Oettle et al: JAMA 297:267, 2007	Gemcitabine vs observation	368	Median DFS 13.4 vs 6.9 ($p < .001$)	22.1 vs 20.2 ($p = .06$)
ESPAC-3, Neoptolemos et al: JAMA 304:1073, 2010	5-FU/LV vs gemcitabine	1088		23 vs 23.6 (HR 0.94; 95% CI 0.81–1.08, $p = .39$)

Abbreviations: CI, confidence interval; CONKO, Charité Onkologie; DFS, disease-free survival; ESPAC, European Study Group for Pancreatic Cancer; 5-FU, 5-fluorouracil; HR, hazard ratio; LV, leucovorin; PFS, progression-free survival.

TABLE 112-2 SELECTED PHASE III STUDIES EVALUATING CHEMOTHERAPY TREATMENT IN ADVANCED PANCREATIC CANCER

Study	Comparator Arm	No. of Patients	Survival	
			PFS (months)	Median Survival (months)
Moore et al: J Clin Oncol 26:1960, 2007	Gemcitabine vs gemcitabine + erlotinib	569	3.55 vs 3.75 (HR 0.77; 95% CI 0.64–0.92; $p = .004$)	5.91 vs 6.24 (HR 0.82; 95% CI 0.69–0.99; $p = .038$)
Cunningham et al: J Clin Oncol 27:5513, 2009	Gemcitabine vs gemcitabine + capecitabine (GEM-CAP)	533	3.8 vs 5.3 (HR 0.78; 95% CI 0.66–0.93; $p = .004$)	6.2 vs 7.1 (HR 0.86; 95% CI 0.72–1.02; $p = .08$)
Von Hoff et al: N Engl J Med 369:1691, 2013	Gemcitabine vs gemcitabine + nab-paclitaxel	861	3.7 vs 5.5 (HR 0.69; 95% CI 0.58–0.82; $p < .001$)	6.7 vs 8.5 (HR 0.72; 95% CI 0.62–0.83; $p < .001$)
Conroy et al: N Engl J Med 364:1817, 2011	Gemcitabine vs FOLFIRINOX	342	3.3 vs 6.4 (HR 0.47; 95% CI 0.37–0.59; $p < .001$)	6.8 vs 11.1 (HR 0.57; 95% CI 0.45–0.73; $p < .001$)

Onkologie trial (CONKO 001) found that the use of gemcitabine after complete resection significantly delayed the development of recurrent disease compared with surgery alone. The European Study Group for Pancreatic Cancer 3 (ESPAC-3) trial, which investigated the benefit of adjuvant 5-fluorouracil/folinic acid (5-FU/FA) versus gemcitabine, revealed no survival difference between the two drugs. However, the toxicity profile of adjuvant gemcitabine was superior to 5-FU/FA by virtue of its lower incidence of stomatitis and diarrhea. Adjuvant radiotherapy is not commonly used in Europe based on the negative results of the ESPAC-1 study. Adjuvant 5-FU-based CRT with gemcitabine before and after radiotherapy as used in the Radiation Therapy Oncology Group (RTOG) 97-04 trial is preferred in the United States. This approach may be most beneficial in patients with bulky tumors involving the pancreatic head.

INOPERABLE LOCALLY ADVANCED DISEASE

Approximately 30% of patients present with locally advanced, unresectable, but nonmetastatic pancreatic carcinoma. The median survival with gemcitabine is 9 months. Patients who respond to chemotherapy or who achieve stable disease after 3–6 months of gemcitabine have frequently been offered consolidation radiotherapy. However, a large, phase III, randomized controlled trial, LAP-07, did not demonstrate any improvement in survival for patients treated with CRT after 4 months of disease control on either gemcitabine or a gemcitabine/erlotinib combination.

METASTATIC DISEASE

Approximately 60% of patients with pancreatic cancer present with metastatic disease. Patients with poor performance status do not usually benefit from chemotherapy. Gemcitabine was the standard

treatment with a median survival of 6 months and a 1-year survival rate of only 20%. The addition of nab-paclitaxel (an albumin bound nanoparticle formulation of paclitaxel) to gemcitabine results in significantly improved 1-year survival compared to gemcitabine alone (35% vs 22%, $p < .001$). Capecitabine, an oral fluoropyrimidine, has also been combined with gemcitabine (GEM-CAP) in a phase III trial that showed an improvement in response rate and progression-free survival over single-agent gemcitabine, but no overall survival benefit. However, pooling of two other randomized controlled trials with this trial in a meta-analysis resulted in a survival advantage with GEM-CAP. Addition of erlotinib, a small-molecule epidermal growth factor receptor inhibitor, produced a statistically significant but clinically marginal benefit when added to gemcitabine in the advanced disease setting. A phase III trial limited to good performance status patients with metastatic pancreatic cancer showed improved survival with the combination of 5-FU/FA, irinotecan, and oxaliplatin (FOLFIRINOX) compared with gemcitabine, but with increased toxicity (Table 112-2).

FUTURE DIRECTIONS

The early detection and future treatment of pancreatic cancer relies on an improved understanding of molecular pathways involved in the development of this disease. This will ultimately lead to the discovery of novel agents and the identification of patient groups who are likely to benefit most from targeted therapy.

ACKNOWLEDGMENT

Dr. Irene Chong is acknowledged for her work on this chapter in the 18th edition.

113 Endocrine Tumors of the Gastrointestinal Tract and Pancreas

Robert T. Jensen

GENERAL FEATURES OF GASTROINTESTINAL NEUROENDOCRINE TUMORS

Gastrointestinal (GI) neuroendocrine tumors (NETs) are tumors derived from the diffuse neuroendocrine system of the GI tract; that system is composed of amine- and acid-producing cells with different hormonal profiles, depending on the site of origin. The tumors historically are divided into GI-NETs (in the GI tract) (also frequently called *carcinoid tumors*) and pancreatic neuroendocrine tumors (pNETs), although newer pathologic classifications have proposed that they all be classified as GI-NETs. The term *GI-NET* has been proposed to replace the term *carcinoid*; however, the term *carcinoid* is widely used, and many are not familiar with this change.

Accordingly, this chapter will use the term *GI-NETs* (carcinoids). These tumors originally were classified as APUDomas (for *amine precursor uptake and decarboxylation*), as were pheochromocytomas, melanomas, and medullary thyroid carcinomas, because they share certain cytochemical features as well as various pathologic, biologic, and molecular features (Table 113-1). It was originally proposed that APUDomas had a similar embryonic origin from neural crest cells, but it is now known the peptide-secreting cells are not of neuroectodermal origin. Nevertheless, the concept of APUDomas is useful because these tumors have important similarities as well as some differences (Table 113-1). In this section, the areas of similarity between pNETs and GI-NETs (carcinoids) will be discussed together, and areas in which there are important differences will be discussed separately.

CLASSIFICATION/PATHOLOGY/TUMOR BIOLOGY OF NETs

NETs generally are composed of monotonous sheets of small round cells with uniform nuclei, and mitoses are uncommon. They can be identified tentatively on routine histology; however, these tumors are now recognized principally by their histologic staining patterns due to shared cellular proteins. Historically, silver staining was used, and

TABLE 113-1 GENERAL CHARACTERISTICS OF GASTROINTESTINAL NEUROENDOCRINE TUMORS (GI-NETs [CARCINOIDS], PANCREATIC NEUROENDOCRINE TUMORS [pNETs])

A.	Share general neuroendocrine cell markers (identification used for diagnosis)
	<ol style="list-style-type: none"> 1. Chromogranins (A, B, C) are acidic monomeric soluble proteins found in the large secretory granules. Chromogranin A is the most widely used. 2. Neuron-specific enolase (NSE) is the γ-γ dimer of the enzyme enolase and is a cytosolic marker of neuroendocrine differentiation. 3. Synaptophysin is an integral membrane glycoprotein of 38,000 molecular weight found in small vesicles of neurons and neuroendocrine tumors.
B.	Pathologic similarities
	<ol style="list-style-type: none"> 1. All are APUDomas showing amine precursor uptake and decarboxylation. 2. Ultrastructurally, they have dense-core secretory granules (>80 nm). 3. Histologically, they generally appear similar with few mitoses and uniform nuclei. 4. Frequently synthesize multiple peptides/amines, which can be detected immunocytochemically but may not be secreted. 5. Presence or absence of clinical syndrome or type cannot be predicted by immunocytochemical studies. 6. Histologic classifications (grading, TNM classification) have prognostic significance. Only invasion or metastases establish malignancy.
C.	Similarities of biologic behavior
	<ol style="list-style-type: none"> 1. Generally slow growing, but some are aggressive. 2. Most are well-differentiated tumors having low proliferative indices. 3. Secrete biologically active peptides/amines, which can cause clinical symptoms. 4. Generally have high densities of somatostatin receptors, which are used for both localization and treatment. 5. Most (>70%) secrete chromogranin A, which is frequently used as a tumor marker.
D.	Similarities/differences in molecular abnormalities
	<ol style="list-style-type: none"> 1. Similarities <ol style="list-style-type: none"> a. Uncommon—mutations in common oncogenes (<i>ras</i>, <i>jun</i>, <i>fos</i>, etc). b. Uncommon—mutations in common tumor-suppressor genes (<i>p53</i>, retinoblastoma). c. Alterations at MEN 1 locus (11q13) (frequently foregut, less commonly mid/hindgut NETs) and $p16^{\text{INK4a}}$ (9p21) occur in a proportion (10–45%). d. Methylation of various genes occurs in 40–87% (<i>ras</i>-associated domain family I, <i>p14</i>, <i>p16</i>, <i>O</i>⁶-methylguanine methyltransferases, retinoic acid receptor β). 2. Differences <ol style="list-style-type: none"> a. pNETs—loss of 1p (21%), 3p (8–47%), 3q (8–41%), 11q (21–62%), 6q (18–68%), Y (45%). Gains at 17q (10–55%), 7q (16–68%), 4q (33%), 18 (up to 45%). b. GI-NETs (carcinoids)—loss of 18q (38–88%), >18p (33–43%), >9p, 16q21 (21–23%). Gains at 17q, 19p (57%), 4q (33%), 14q (20%), 5 (up to 36%). c. pNETs: <i>ATRX/DAXX</i> mutations in 43%, MEN 1 mutations in 44%, mTOR mutations (14%); uncommon in midgut GI-NETs (0–2%).

Abbreviations: *ATRX*, alpha-thalassemia X-lined mental retardation protein; *DAXX*, death domain associated protein; *MEN 1*, multiple endocrine neoplasia type 1; TNM, tumor, node, metastasis.

tumors were classified as showing an argentaffin reaction if they took up and reduced silver or as being argyrophilic if they did not reduce it. Currently, immunocytochemical localization of chromogranins (A, B, C), neuron-specific enolase, and synaptophysin, which are all neuroendocrine cell markers, is used (Table 113-1). Chromogranin A is the most widely used.

Ultrastructurally, these tumors possess electron-dense neurosecretory granules and frequently contain small clear vesicles that correspond to synaptic vesicles of neurons. NETs synthesize numerous peptides, growth factors, and bioactive amines that may be ectopically secreted, giving rise to a specific clinical syndrome (Table 113-2). The diagnosis of the specific syndrome requires the clinical features of the disease (Table 113-2) and cannot be made from the immunocytochemistry results alone. The presence or absence of a specific clinical syndrome also cannot be predicted from the immunocytochemistry alone (Table 113-1). Furthermore, pathologists cannot distinguish between benign and malignant NETs unless metastasis or invasion is present.

GI-NETs (carcinoids) frequently are classified according to their anatomic area of origin (i.e., foregut, midgut, hindgut) because tumors with similar areas of origin share functional manifestations, histochemistry, and secretory products (Table 113-3). Foregut tumors generally have a low serotonin (5-HT) content; are argentaffin-negative but argyrophilic; occasionally secrete adrenocorticotropic hormone (ACTH) or 5-hydroxytryptophan (5-HTP), causing an atypical carcinoid syndrome (Fig. 113-1); are often multihormonal; and may metastasize to bone. They uncommonly produce a clinical syndrome due to the secreted products. Midgut carcinoids are argentaffin-positive, have a high serotonin content, most frequently cause the typical carcinoid syndrome when they metastasize (Table 113-3, Fig. 113-1), release serotonin and tachykinins (substance P, neuropeptide K, substance K), rarely secrete 5-HTP or ACTH, and less commonly metastasize to

bone. Hindgut carcinoids (rectum, transverse and descending colon) are argentaffin-negative, are often argyrophilic, rarely contain serotonin or cause the carcinoid syndrome (Fig. 113-1, Table 113-3), rarely secrete 5-HTP or ACTH, contain numerous peptides, and may metastasize to bone.

pNETs can be classified into nine well-established specific functional syndromes (Table 113-2), six additional very rare specific functional syndromes (less than five cases described), five possible specific functional syndromes (pNETs secreting calcitonin, neurotensin, pancreatic polypeptide, ghrelin) (Table 113-2), and nonfunctional pNETs. Other functional hormonal syndromes due to nonpancreatic tumors (usually intraabdominal in location) have been described only rarely and are not included in (Table 113-2). These include secretion by intestinal and ovarian tumors of peptide tyrosine tyrosine (PYY), which results in altered motility and constipation, and ovarian tumors secreting renin or aldosterone causing alterations in blood pressure or somatostatin causing diabetes or reactive hypoglycemia. Each of the functional syndromes listed in Table 113-2 is associated with symptoms due to the specific hormone released. In contrast, nonfunctional pNETs release no products that cause a specific clinical syndrome. “Nonfunctional” is a misnomer in the strict sense because those tumors frequently ectopically secrete a number of peptides (pancreatic polypeptide [PP], chromogranin A, ghrelin, neurotensin, α subunits of human chorionic gonadotropin, and neuron-specific enolase); however, they cause no specific clinical syndrome. The symptoms caused by nonfunctional pNETs are entirely due to the tumor per se. pNETs frequently ectopically secrete PP (60–85%), neurotensin (30–67%), calcitonin (30–42%), and to a lesser degree, ghrelin (5–65%). Whereas a few studies have proposed their secretion can cause a specific functional syndrome, most studies support the conclusion that their ectopic secretion is not associated with a specific clinical syndrome, and thus they are listed in Table 113-2 as possible clinical syndromes.

TABLE 113-2 GASTROINTESTINAL NEUROENDOCRINE TUMOR SYNDROMES

Name	Biologically Active Peptide(s) Secreted	Incidence (New Cases/10 ⁶ Population/Year)	Tumor Location	Malignant, %	Associated with MEN 1, %	Main Symptoms/Signs
I. Established Specific Functional Syndromes						
A. Carcinoid syndrome due to GI-NET						
Carcinoid syndrome	Serotonin, possibly tachykinins, motilin, prostaglandins	0.5–2	Midgut (75–87%) Foregut (2–33%) Hindgut (1–8%) Unknown (2–15%)	95–100	Rare	Diarrhea (32–84%) Flushing (63–75%) Pain (10–34%) Asthma (4–18%) Heart disease (11–41%)
B. Well-established functional pNET syndromes						
Zollinger-Ellison syndrome	Gastrin	0.5–1.5	Duodenum (70%) Pancreas (25%) Other sites (5%)	60–90	20–25	Pain (79–100%) Diarrhea (30–75%) Esophageal symptoms (31–56%)
Insulinoma	Insulin	1–2	Pancreas (>99%)	<10	4–5	Hypoglycemic symptoms (100%)
VPoma (Verner-Morrison syndrome, pancreatic cholera, WDHA)	Vasoactive intestinal peptide	0.05–0.2	Pancreas (90%, adult) Other (10%, neural, adrenal, periganglionic)	40–70	6	Diarrhea (90–100%) Hypokalemia (80–100%) Dehydration (83%)
Glucagonoma	Glucagon	0.01–0.1	Pancreas (100%)	50–80	1–20	Rash (67–90%) Glucose intolerance (38–87%) Weight loss (66–96%)
Somatostatinoma	Somatostatin	Rare	Pancreas (55%) Duodenum/jejunum (44%)	>70	45	Diabetes mellitus (63–90%) Cholelithiasis (65–90%) Diarrhea (35–90%)
GRFoma	Growth hormone–releasing hormone	Unknown	Pancreas (30%) Lung (54%) Jejunum (7%) Other (13%)	>60	16	Acromegaly (100%)
ACTHoma	ACTH	Rare	Pancreas (4–16% all ectopic Cushing's)	>95	Rare	Cushing's syndrome (100%)
pNET causing carcinoid syndrome	Serotonin, ?tachykinins	Rare (43 cases)	Pancreas (<1% all carcinoids)	60–88	Rare	Same as carcinoid syndrome above
pNET causing hypercalcemia	PTHrP Others unknown	Rare	Pancreas (rare cause of hypercalcemia)	84	Rare	Abdominal pain due to hepatic metastases
II. Rare Specific Functional Syndromes						
pNET secreting renin	Renin	Rare	Pancreas	Unknown	No	Hypertension
pNET secreting luteinizing hormone	Luteinizing hormone	Rare	Pancreas	Unknown	No	Anovulation, virilization (female); reduced libido (male)
pNET secreting erythropoietin	Erythropoietin	Rare	Pancreas	100	No	Polycythemia
pNET secreting IGF-II	Insulin-like growth factor II	Rare	Pancreas	Unknown	No	Hypoglycemia
pNET secreting GLP-1	Glucagon-like peptide-1	Rare	Pancreas	Unknown	No	Hypoglycemia, diabetes
pNET secreting enteroglucagon	Enteroglucagon	Rare	Pancreas, small intestine	Unknown	Rare	Small intestinal hypertrophy, intestinal stasis, malabsorption
III. Possible Specific Functional pNET Syndromes						
pNET secreting calcitonin	Calcitonin	Rare	Pancreas (rare cause of hypercalcitonemia)	>80	16	Diarrhea (50%)
pNET secreting neurotensin	Neurotensin	Rare	Pancreas (100%)	Unknown	No	Motility disturbances, vascular symptoms
pNET secreting pancreatic polypeptide (PPoma)	Pancreatic polypeptide	1–2	Pancreas	>60	18–44	Watery diarrhea
pNET secreting ghrelin	Ghrelin	Rare	Pancreas	Unknown	No	Effects on appetite, body weight
IV. Nonfunctional Syndrome pNET						
PPoma/nonfunctional ^a	None	1–2	Pancreas (100%)	>60	18–44	Weight loss (30–90%) Abdominal mass (10–30%) Pain (30–95%)

Abbreviations: ACTH, adrenocorticotropic hormone; GRFoma, growth hormone–releasing factor secreting pancreatic endocrine tumor; IGF-II, insulin-like growth factor II; MEN, multiple endocrine neoplasia; pNET, pancreatic neuroendocrine tumor; PPoma, tumor secreting pancreatic polypeptide; PTHrP, parathyroid hormone–related peptide; VPoma, tumor secreting vasoactive intestinal peptide; WDHA, watery diarrhea, hypokalemia, and achlorhydria syndrome.

^aPancreatic polypeptide–secreting tumors (PPomas) are listed in two places because most authorities classify these as not associated with a specific hormonal syndrome (nonfunctional); however, rare cases of watery diarrhea proposed to be due to PPomas have been reported.

TABLE 113-3 GI-NET (CARCINOID) LOCATION, FREQUENCY OF METASTASES, AND ASSOCIATION WITH THE CARCINOID SYNDROME

	Location (% of Total)	Incidence of Metastases	Incidence of Carcinoid Syndrome
Foregut			
Esophagus	<0.1	—	—
Stomach	4.6	10	9.5
Duodenum	2.0	—	3.4
Pancreas	0.7	71.9	20
Gallbladder	0.3	17.8	5
Bronchus, lung, trachea	27.9	5.7	13
Midgut			
Jejunum	1.8	{58.4	9
Ileum	14.9	—	9
Meckel's diverticulum	0.5	—	13
Appendix	4.8	38.8	<1
Colon	8.6	51	5
Liver	0.4	32.	—
Ovary	1.0	2 32	50
Testis	<0.1	—	50
Hindgut			
Rectum	13.6	3.9	—

Abbreviation: GI-NET, gastrointestinal neuroendocrine tumor.

Source: Location is from the PAN-SEER data (1973–1999), and incidence of metastases is from the SEER data (1992–1999), reported by IM Modlin et al: Cancer 97:934, 2003. Incidence of carcinoid syndrome is from 4349 cases studied from 1950–1971, reported by JD Godwin: Cancer 36:560, 1975.

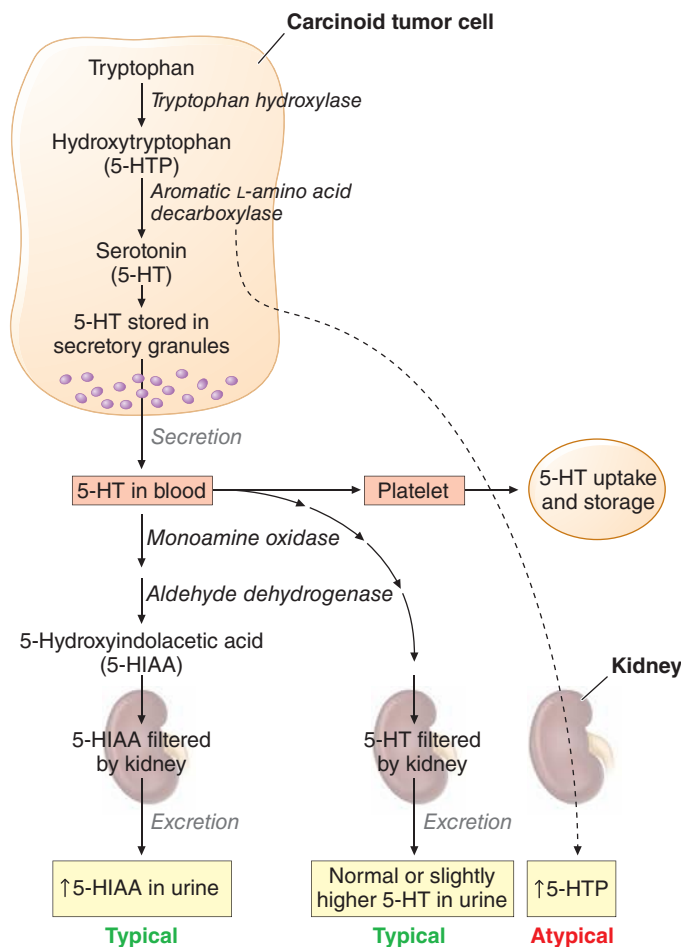


FIGURE 113-1 Synthesis, secretion, and metabolism of serotonin (5-HT) in patients with typical and atypical carcinoid syndromes. 5-HIAA, 5-hydroxyindolacetic acid.

Because a large proportion of nonfunctional pNETs (60–90%) secrete PP, these tumors are often referred to as PPomas (Table 113-2).

GI-NETs (carcinoids) can occur in almost any GI tissue (Table 113-3); however, at present, most (70%) have their origin in one of three sites: bronchus, jejunoleum, or colon/rectum. In the past, GI-NET (carcinoids) most frequently were reported in the appendix (i.e., 40%); however, the bronchus/lung, rectum, and small intestine are now the most common sites. Overall, the GI tract is the most common site for these tumors, accounting for 64%, with the respiratory tract a distant second at 28%. Both race and sex can affect the frequency as well as the distribution of GI-NETs (carcinoids). African Americans have a higher incidence of carcinoids. Race is particularly important for rectal carcinoids, which are found in 41% of Asians/Pacific Islanders with NETs compared to 32% of American Indians/Alaskan natives, 26% of African Americans, and 12% of white Americans. Females have a lower incidence of small intestinal and pancreatic carcinoids.

The term *pancreatic neuroendocrine* or *endocrine tumor*, although widely used and therefore retained here, is also a misnomer, strictly speaking, because these tumors can occur either almost entirely in the pancreas (insulinomas, glucagonomas, nonfunctional pNETs, pNETs causing hypercalcemia) or at both pancreatic and extrapancreatic sites (gastrinomas, VIPomas [vasoactive intestinal peptide], somatostatinomas, GRFomas [growth hormone-releasing factor]). pNETs are also called islet cell tumors; however, the use of this term is discouraged because it is not established that they originate from the islets, and many can occur at extrapancreatic sites.

Whereas the classification of GI neuroendocrine tumors into foregut, midgut, or hindgut is widely used and generally useful because the NETs within these areas have many similarities, they also have marked differences, particularly in biologic behavior, and it has not proved useful for prognostic purposes. More general classifications have been developed that allow NETs with similar features in different locations to be compared, have proven prognostic value, and are widely used. New classification systems have been developed for both GI-NETs (carcinoids) and pNETs by the World Health Organization (WHO), European Neuroendocrine Tumor Society (ENETS), and the American Joint Committee on Cancer/International Union Against Cancer (AJCC/UICC). Although there are some differences between these different classification systems, each uses similar information, and it is now recommended that the basic data underlying the classification be included in all standard pathology reports. These classification systems divide NETs from all sites into those that are well differentiated (low grade [G1] or intermediate grade [G2]) and those that are poorly differentiated (high grade [G3] divided into either small-cell carcinoma or large-cell neuroendocrine carcinoma). In these classification systems, both pNETs and GI-NETs (carcinoids) are classified as neuroendocrine tumors, and the old term of carcinoid is equivalent to well-differentiated neuroendocrine tumors of the GI tract. These classification systems are based on not only the differentiation of the NET, but also a grading system assessing proliferative indices (Ki-67 and the mitotic count). NETs are considered low grade (ENETS G1) if the Ki-67 is <3% and the mitotic count is <2 mitoses/high-power field (HPF), intermediate grade (ENETS G2) if the Ki-67 is 3–20% and the mitotic count is 2–20 mitoses/HPF, and high grade (ENETS G3) if the Ki-67 is >20% and the mitotic count is >20 mitoses/HPF. In addition to the grading system, a TNM classification has been proposed that is based on the level of tumor invasion, tumor size, and tumor extent (see Table 113-4 for an example with pNETs and appendiceal GI-NETs [carcinoids]). Because of the proven prognostic value of these classification and grading systems, as well as the fact that NETs with different classifications/grades respond differently to treatments, the systems are now essential for the management of all NETs.

In addition to these classification/grading systems, a number of other factors have been identified that provide important prognostic information that can guide treatment (Table 113-5).

The exact incidence of GI-NETs (carcinoids) or pNETs varies according to whether only symptomatic tumors or all tumors are considered. The incidence of clinically significant carcinoids is 7–13 cases/million population per year, whereas any malignant carcinoids

TABLE 113-4 COMPARISON OF THE CRITERIA FOR THE TUMOR CATEGORY IN THE ENETS AND SEVENTH EDITION AJCC TNM CLASSIFICATIONS OF PANCREATIC AND APPENDICEAL NETS

	ENETS TNM	AJCC/UICC TNM
pNETs		
T1	Confined to pancreas, <2 cm	Confined to pancreas, <2 cm
T2	Confined to pancreas, 2–4 cm	Confined to pancreas, >2 cm
T3	Confined to pancreas, >4 cm, or invasion of duodenum or bile duct	Peripancreatic spread, but without major vascular invasion (truncus coeliacus, superior mesenteric artery)
T4	Invasion of adjacent organs or major vessels	Major vascular invasion
Appendiceal NETs		
T1	≤1 cm; invasion of muscularis propria	T1a, ≤1 cm; T1b, >1–2 cm
T2	≤2 cm and <3 mm invasion of subserosa/mesoappendix	>2–4 cm or invasion of cecum
T3	>2 cm or >3 mm invasion of subserosa/mesoappendix	>4 cm or invasion of ileum
T4	Invasion of peritoneum/other organs	Invasion of peritoneum/other organs

Abbreviations: AJCC, American Joint Committee on Cancer; ENETS, European Neuroendocrine Tumor Society; NET, neuroendocrine tumor; pNET, pancreatic neuroendocrine tumor; TNM, tumor, node, metastasis; UICC, International Union Against Cancer.

Source: Modified from DS Klimstra: *Semin Oncol* 40:23, 2013 and G Kloppel et al: *Virchow Arch* 456:595, 2010.

at autopsy are reported in 21–84 cases/million population per year. The incidence of GI-NETs (carcinoids) is approximately 25–50 cases per million in the United States, which makes them less common than adenocarcinomas of the GI tract. However, their incidence has increased sixfold in the last 30 years. In an analysis of 35,825 GI-NETs (carcinoids) (2004) from the U.S. Surveillance, Epidemiology, and End Results (SEER) database, their incidence was 5.25/100,000 per year, and the 29-year prevalence was 35/100,000. Clinically significant pNETs have a prevalence of 10 cases/million population, with insulinomas, gastrinomas, and nonfunctional pNETs having an incidence of 0.5–2 cases/million population per year (Table 113-2). pNETs account for 1–10% of all tumors arising in the pancreas and 1.3% of tumors in the SEER database, which consists primarily of malignant tumors. VIPomas are 2–8 times less common, glucagonomas are 17–30 times less common, and somatostatinomas are the least common. In autopsy studies, 0.5–1.5% of all cases have a pNET; however, in less than 1 in 1000 cases was a functional tumor thought to occur.

Both GI-NETs (carcinoids) and pNETs commonly show malignant behavior (Tables 113-2 and 113-3). With pNETs, except for insulinomas in which <10% are malignant, 50–100% in different series are malignant. With GI-NETs (carcinoids), the percentage showing malignant behavior varies in different locations (Table 113-3). For the three most common sites of occurrence, the incidence of metastases varies greatly from the jejunioileum (58%), lung/bronchus (6%), and rectum (4%) (Table 113-3). With both GI-NETs (carcinoids) and pNETs, a number of factors (Table 113-5) are important prognostic factors in determining survival and the aggressiveness of the tumor. Patients with pNETs (excluding insulinomas) generally have a poorer prognosis than do patients with GI-NETs (carcinoids). The presence of liver metastases is the single most important prognostic factor in single and multivariate analyses for both GI-NETs (carcinoids) and pNETs. Particularly important in the development of liver metastases is the size of the primary tumor. For example, with small intestinal carcinoids, which are the most common cause of the carcinoid syndrome due to metastatic disease in the liver (Table 113-2), metastases occur in 15–25% if the tumor is <1 cm in diameter, 58–80% if it is 1–2 cm in diameter, and >75% if it is >2 cm in diameter. Similar data exist for gastrinomas and other pNETs; the size of the primary tumor is an independent predictor of the development of liver metastases.

TABLE 113-5 PROGNOSTIC FACTORS IN NEUROENDOCRINE TUMORS**I. Both GI-NETs (carcinoids) and pNETs**

Symptomatic presentation ($p < .05$)
 Presence of liver metastases ($p < .001$)
 Extent of liver metastases ($p < .001$)
 Presence of lymph node metastases ($p < .001$)
 Development of bone or extrahepatic metastases ($p < .01$)
 Depth of invasion ($p < .001$)
 Rapid rate of tumor growth
 Elevated serum alkaline phosphatase levels ($p = .003$)
 Primary tumor site ($p < .001$)
 Primary tumor size ($p < .005$)
 High serum chromogranin A level ($p < .01$)
 Presence of one or more circulating tumor cells ($p < .001$)
 Various histologic features
 Tumor differentiation ($p < .001$)
 High growth indices (high Ki-67 index, PCNA expression)
 High mitotic counts ($p < .001$)
 Necrosis present
 Presence of cytokeratin 19 ($p < .02$)
 Vascular or perineural invasion
 Vessel density (low microvessel density, increased lymphatic density)
 High CD10 metalloproteinase expression (in series with all grades of NETs)
 Flow cytometric features (i.e., aneuploidy)
 High VEGF expression (in low-grade or well-differentiated NETs only)
 WHO, ENETS, AJCC/UICC, and grading classification
 Presence of a pNET rather than GI-NET associated with poorer prognosis ($p = .0001$)
 Older age ($p < .01$)

II. GI-NETs (Carcinoids)

Location of primary: appendix < lung, rectum < small intestine < pancreas
 Presence of carcinoid syndrome
 Laboratory results (urinary 5-HIAA levels [$p < .01$], plasma neuropeptide K [$p < .05$], serum chromogranin A [$p < .01$])
 Presence of a second malignancy
 Male sex ($p < .001$)
 Molecular findings (TGF- α expression [$p < .05$], chr 16q LOH or gain chr 4p [$p < .05$])
 WHO, ENETS, AJCC/UICC, and grading classification
 Molecular findings (gain in chr 14, loss of 3p13 [ileal carcinoid], upregulation of Hoxc6)

III. pNETs

Location of primary: duodenal (gastrinoma) better than pancreatic
 Ha-ras oncogene or p53 overexpression
 Female gender
 MEN 1 syndrome absent
 Presence of nonfunctional tumor (some studies, not all)
 WHO, ENETS, AJCC/UICC, and grading classification
 Various histologic features: IHC positivity for c-KIT, low cyclin B1 expression ($p < .01$), loss of PTEN or of tuberous sclerosis-2 IHC, expression of fibroblast growth factor-13
 Laboratory findings (increased chromogranin A in some studies; gastrinomas—increased gastrin level)
 Molecular findings (increased HER2/*neu* expression [$p = .032$], chr 1q, 3p, 3q, or 6q LOH [$p = .0004$], EGF receptor overexpression [$p = .034$], gains in chr 7q, 17q, 17p, 20q; alterations in the VHL gene [deletion, methylation]; presence of FGFR4-G388R single-nucleotide polymorphism)

Abbreviations: 5-HIAA, 5-hydroxyindoleacetic acid; AJCC, American Joint Committee on Cancer; chr, chromosome; EGF, epidermal growth factor; FGFR, fibroblast growth factor receptor; GI-NET, gastrointestinal neuroendocrine tumor; IHC, immunohistochemistry; Ki-67, proliferation-associated nuclear antigen recognized by Ki-67 monoclonal antibody; LOH, loss of heterozygosity; MEN, multiple endocrine neoplasia; NET, neuroendocrine tumor; PCNA, proliferating cell nuclear antigen; pNET, pancreatic neuroendocrine tumor; PTEN, phosphatase and tensin homologue deleted from chromosome 10; TGF- α , transforming growth factor α ; TNM, tumor, node, metastasis; UICC, International Union Against Cancer; VEGF, vascular endothelial growth factor; WHO, World Health Organization.

562 The presence of lymph node metastases or extrahepatic metastases; the depth of invasion; the rapid rate of growth; various histologic features (differentiation, mitotic rates, growth indices, vessel density, vascular endothelial growth factor [VEGF], and CD10 metalloproteinase expression); necrosis; presence of cytokeratin; elevated serum alkaline phosphatase levels; older age; presence of circulating tumor cells; and flow cytometric results, such as the presence of aneuploidy, are all important prognostic factors for the development of metastatic disease (Table 113-5). For patients with GI-NETs (carcinoids), additional associations with a worse prognosis include the development of the carcinoid syndrome (especially the development of carcinoid heart disease), male sex, the presence of a symptomatic tumor or greater increases in a number of tumor markers (5-hydroxyindolacetic acid [5-HIAA], neuropeptide K, chromogranin A), and the presence of various molecular features. With pNETs or gastrinomas, a worse prognosis is associated with female sex, overexpression of the *Ha-ras* oncogene or p53, the absence of multiple endocrine neoplasia type 1 (MEN 1), higher levels of various tumor markers (i.e., chromogranin A, gastrin), and presence of various histologic features (immunohistochemistry for c-KIT, low cyclin B1, loss of PTEN/TSC-2, expression of fibroblast growth factor-13) and various molecular features (Table 113-5). The TNM classification systems and the grading systems (G1–G3) have important prognostic value.

A number of diseases due to various genetic disorders are associated with an increased incidence of NETs (Table 113-6). Each one is caused by a loss of a possible tumor-suppressor gene. The most important is MEN 1, which is an autosomal dominant disorder due to a defect in a 10-exon gene on 11q13, which encodes for a 610-amino-acid nuclear protein, menin (Chap. 408). Patients with MEN 1 develop hyperparathyroidism due to parathyroid hyperplasia in 95–100% of cases, pNETs in 80–100%, pituitary adenomas in 54–80%, adrenal adenomas in 27–36%, bronchial carcinoids in 8%, thymic carcinoids in 8%, gastric carcinoids in 13–30% of patients with Zollinger-Ellison syndrome, skin tumors (angiofibromas [88%], collagenomas [72%]), central nervous system (CNS) tumors (meningiomas [$<8\%$]), and smooth-muscle tumors (leiomyomas, leiomyosarcomas [1–7%]). Among patients with MEN 1, 80–100% develop nonfunctional pNETs (most are microscopic with 0–13% large/symptomatic), and functional pNETs occur in 20–80% in different series, with a mean of 54% developing Zollinger-Ellison syndrome, 18% insulinomas, 3% glucagonomas, 3%

VIPomas, and $<1\%$ GRFomas or somatostatinomas. MEN 1 is present in 20–25% of all patients with Zollinger-Ellison syndrome, 4% of patients with insulinomas, and a low percentage ($<5\%$) of patients with other pNETs.

Three phacomatoses associated with NETs are von Hippel–Lindau disease (VHL), von Recklinghausen’s disease (neurofibromatosis type 1 [NF-1]), and tuberous sclerosis (Bourneville’s disease) (Table 113-6). VHL is an autosomal dominant disorder due to defects on chromosome 3p25, which encodes for a 213-amino-acid protein that interacts with the elongin family of proteins as a transcriptional regulator (Chaps. 118, 339, 407, and 408). In addition to cerebellar hemangioblastomas, renal cancer, and pheochromocytomas, 10–17% develop a pNET. Most are nonfunctional, although insulinomas and VIPomas have been reported. Patients with NF-1 (von Recklinghausen’s disease) have defects in a gene on chromosome 17q11.2 that encodes for a 2845-amino-acid protein, neurofibromin, which functions in normal cells as a suppressor of the *ras* signaling cascade (Chap. 118). Up to 10% of these patients develop an upper GI-NET (carcinoid), characteristically in the periampullary region (54%). Many are classified as somatostatinomas because they contain somatostatin immunocytochemically; however, they uncommonly secrete somatostatin and rarely produce a clinical somatostatinoma syndrome. NF-1 has rarely been associated with insulinomas and Zollinger-Ellison syndrome. NF-1 accounts for 48% of all duodenal somatostatinomas and 23% of all ampullary GI-NETs (carcinoids). Tuberous sclerosis is caused by mutations that alter either the 1164-amino-acid protein hamartin (TSC1) or the 1807-amino-acid protein tuberlin (TSC2) (Chap. 118). Both hamartin and tuberlin interact in a pathway related to phosphatidylinositol 3-kinases and mammalian target of rapamycin (mTOR) signaling cascades. A few cases including nonfunctional and functional pNETs (insulinomas and gastrinomas) have been reported in these patients (Table 113-6). Mahvash disease is associated with the development of α -cell hyperplasia, hyperglucagonemia, and the development of NF pNETs and is due to a homozygous P86S mutation of the human glucagon receptor.

Mutations in common oncogenes (*ras*, *myc*, *fos*, *src*, *jun*) or common tumor-suppressor genes (*p53*, retinoblastoma susceptibility gene) are not commonly found in either pNETs or GI-NETs (carcinoids) (Table 113-1). However, frequent (70%) gene amplifications in *MDM2*, *MDM4*, and *WIP1* inactivating the p53 pathway are noted in well-differentiated pNETs, and the retinoblastoma pathway is altered in the majority of pNETs. In addition to these genes, additional alterations that may be important in their pathogenesis include changes in the *MEN1* gene, *p16/MTS1* tumor-suppressor gene, and *DPC4/Smad4* gene; amplification of the *HER-2/neu* protooncogene; alterations in transcription factors (Hoxc6 [GI carcinoids]), growth factors, and their receptors; methylation of a number of genes that probably results in their inactivation; and deletions of unknown tumor-suppressor genes as well as gains in other unknown genes (Table 113-1). The clinical antitumor activity of everolimus, an mTOR inhibitor, and sunitinib, a tyrosine kinase inhibitor (PDGFR, VEGFR1, VEGFR2, c-KIT, FLT-3), support the importance of the mTOR-AKT pathway and tyrosine kinase receptors in mediating growth of malignant NETs (especially pNETs). The importance of the mTOR pathway in pNET growth is further supported by the finding that a single-nucleotide polymorphism (FGFR4-G388R, in fibroblast growth factor receptor 4) affects selectivity to the mTOR inhibitor and can result in significantly higher risk of advanced pNET stage and liver metastases (Table 113-5). Comparative genomic hybridization, genome-wide allelotyping studies, and genome-wide single-nucleotide polymorphism analyses have shown that chromosomal losses and gains are common in pNETs and GI-NETs (carcinoids), but they differ between these two NETs, and some have prognostic significance (Table 113-5). Mutations in the *MEN1* gene are probably particularly important. Loss of heterozygosity at the MEN 1 locus on chromosome 11q13 is noted in 93% of sporadic pNETs (i.e., in patients without MEN 1) and in 26–75% of sporadic GI-NETs (carcinoids). Mutations in the *MEN1* gene are reported in 31–34% of sporadic gastrinomas. Exomic sequencing of sporadic pNETs found that the most frequently altered gene was

TABLE 113-6 GENETIC SYNDROMES ASSOCIATED WITH AN INCREASED INCIDENCE OF NEUROENDOCRINE TUMORS (NETS) (GI-NETS [CARCINOIDS] OR PNETS)

Syndrome	Location of Gene Mutation and Gene Product	NETs Seen/Frequency
Multiple endocrine neoplasia type 1 (MEN 1)	11q13 (encodes 610-amino-acid protein, menin)	80–100% develop pNETs (microscopic), 20–80% (clinical): (nonfunctional > gastrinoma > insulinoma) GI-NETs (Carcinoids): gastric (13–30%), bronchial/thymic (8%)
von Hippel–Lindau disease	3q25 (encodes 213-amino-acid protein)	12–17% develop pNETs (almost always nonfunctional)
von Recklinghausen’s disease (neurofibromatosis 1 [NF-1])	17q11.2 (encodes 2485-amino-acid protein, neurofibromin)	0–10% develop pNETs, primarily duodenal somatostatinomas (usually nonfunctional) Rarely insulinoma, gastrinoma
Tuberous sclerosis	9q34 (TSC1) (encodes 1164-amino-acid protein, hamartin), 16p13 (TSC2) (encodes 1807-amino-acid protein, tuberlin)	Uncommonly develop pNETs (nonfunctional and functional [insulinoma, gastrinoma])

Abbreviations: GI, gastrointestinal; pNETs, pancreatic neuroendocrine tumors.

MEN1, occurring in 44% of patients, followed by mutations in 43% of patients in genes encoding for two subunits of a transcription/chromatin remodeling complex consisting of DAXX (death-domain-associated protein) and ATRX (α -thalassemia/mental retardation syndrome X-linked) and in 15% of patients in the mTOR pathway. The presence of a number of these molecular alterations in pNETs or GI-NETs (carcinoids) correlates with tumor growth, tumor size, and disease extent or invasiveness and may have prognostic significance (Table 113-5).

GI-NETs (CARCINOIDS) AND CARCINOID SYNDROME

CHARACTERISTICS OF THE MOST COMMON GI-NETs (CARCINOIDS)

Appendiceal NETs (Carcinoids) Appendiceal NETs (carcinoids) occur in 1 in every 200–300 appendectomies, usually in the appendiceal tip, have an incidence of 0.15/100,000 per year, comprise 2–5% of all GI-NETs (carcinoids), and comprise 32–80% of all appendiceal tumors. Most (i.e., >90%) are <1 cm in diameter without metastases in older studies, but more recently, 2–35% have had metastases (Table 113-3). In the SEER data of 1570 appendiceal carcinoids, 62% were localized, 27% had regional metastases, and 8% had distant metastases. The risk of metastases increases with size, with those <1 cm having a 0 to <10% risk of metastases and those >2 cm having a 25–44% risk. Besides tumor size, other important prognostic factors for metastases include basal location, invasion of mesoappendix, poor differentiation, advanced stage or WHO/ENETS classification, older age, and positive resection margins. The 5-year survival is 88–100% for patients with localized disease, 78–100% for patients with regional involvement, and 12–28% for patients with distal metastases. In patients with tumors <1 cm in diameter, the 5-year survival is 95–100%, whereas it is 29% if tumors are >2 cm in diameter. Most tumors are well-differentiated G1 tumors (87%) (Table 113-4), with the remainder primarily well-differentiated G2 tumors (13%); poorly differentiated G3 tumors are uncommon (<1%). Their percentage of the total number of carcinoids decreased from 43.9% (1950–1969) to 2.4% (1992–1999). Appendiceal goblet cell (GC) NETs (carcinoids)/carcinomas are a rare subtype (<5%) that are mixed adeno-neuroendocrine carcinomas. They are malignant and are thought to comprise a distinct entity; they frequently present with advanced disease and are recommended to be treated as adenocarcinomas, not carcinoid tumors.

SMALL INTESTINAL NETs (CARCINOIDS)

Small intestinal (SI) NETs (carcinoids) have a reported incidence of 0.67/100,000 in the United States, 0.32/100,000 in England, and 1.12/100,000 in Sweden and comprise >50% of all SI tumors. There is a male predominance (1.5:1), and race affects frequency, with a lower frequency in Asians and greater frequency in African Americans. The mean age of presentation is 52–63 years, with a wide range (1–93 years). Familial SI carcinoid families exist but are very uncommon. These are frequently multiple; 9–18% occur in the jejunum, 70–80% are present in the ileum, and 70% occur within 6 cm (2.4 in.) of the ileocecal valve. Forty percent are <1 cm in diameter, 32% are 1–2 cm, and 29% are >2 cm. They are characteristically well differentiated; however, they are generally invasive, with 1.2% being intramucosal in location, 27% penetrating the submucosa, and 20% invading the muscularis propria. Metastases occur in a mean of 47–58% (range 20–100%). Liver metastases occur in 38%, to lymph nodes in 37% and more distant in 20–25%. They characteristically cause a marked fibrotic reaction, which can lead to intestinal obstruction. Tumor size is an important variable in the frequency of metastases. However, even small NETs (carcinoids) of the small intestine (<1 cm) have metastases in 15–25% of cases, whereas the proportion increases to 58–100% for tumors 1–2 cm in diameter. Carcinoids also occur in the duodenum, with 31% having metastases. Duodenal tumors <1 cm virtually never metastasize, whereas 33% of those >2 cm had metastases. SI NETs (carcinoids) are the most common cause (60–87%) of the carcinoid syndrome and are discussed in a later section (Table 113-7). Important prognostic factors are listed in (Table 113-5), and particularly important are the

TABLE 113-7 CLINICAL CHARACTERISTICS IN PATIENTS WITH CARCINOID SYNDROME

	Percentage (Range)	
	At Presentation	During Course of Disease
Symptoms/signs		
Diarrhea	32–93%	68–100%
Flushing	23–100%	45–96%
Pain	10%	34%
Asthma/wheezing	4–14%	3–18%
Pellagra	0–7%	0–5%
None	12%	22%
Carcinoid heart disease present	11–40%	14–41%
Demographics		
Male	46–59%	46–61%
Age		
Mean	57 yrs	59.2 yrs
Range	25–79 yrs	18–91 yrs
Tumor location		
Foregut	5–14%	0–33%
Midgut	57–87%	60–100%
Hindgut	1–7%	0–8%
Unknown	2–21%	0–26%

tumor extent, proliferative index by grading, and stage (Table 113-4). The overall survival at 5 years is 55–75%; however, it varies markedly with disease extent, being 65–90% with localized disease, 66–72% with regional involvement, and 36–43% with distant disease.

Rectal NETs (Carcinoids) Rectal NETs (carcinoids) comprise 27% of all GI-NETs (carcinoids) and 16% of all NETs and are increasing in frequency. In the U.S. SEER data, they currently have an incidence of 0.86/100,000 per year (up from 0.2/100,000 per year in 1973) and represent 1–2% of all rectal tumors. They are found in approximately 1 in every 1500/2500 proctoscopies/colonoscopies or 0.05–0.07% of individuals undergoing these procedures. Nearly all occur between 4 and 13 cm above the dentate line. Most are small, with 66–80% being <1 cm in diameter, and rarely metastasize (5%). Tumors between 1 and 2 cm can metastasize in 5–30%, and those >2 cm, which are uncommon, in >70%. Most invade only to the submucosa (75%), with 2.1% confined to the mucosa, 10% to the muscular layer, and 5% to adjacent structures. Histologically, most are well differentiated (98%) with 72% ENETS/WHO grade G1 and 28% grade G2 (Table 113-4). Overall survival is 88%; however, it is very much dependent of the stage, with 5-year survival of 91% for localized disease, 36–49% for regional disease, and 20–32% for distant disease. Risk factors are listed in Table 113-5 and particularly include tumor size, depth of invasion, presence of metastases, differentiation, and recent TNM classification and grade.

Bronchial NETs (Carcinoids) Bronchial NETs (carcinoids) comprise 25–33% of all well-differentiated NETs and 90% of all the poorly differentiated NETs found, likely due to a strong association with smoking. Their incidence ranges from 0.2 to 2/100,000 per year in the United States and European countries and is increasing at a rate of 6% per year. They are slightly more frequent in females and in whites compared with those of Hispanic/Asian/African descent, and are most commonly seen in the sixth decade of life, with a younger age of presentation for typical carcinoids (45 years) compared to atypical carcinoids (55 years).

A number of different classifications of bronchial GI-NETs (carcinoids) have been proposed. In some studies, they are classified into four categories: typical carcinoid (also called bronchial carcinoid tumor, Kulchitsky cell carcinoma I [KCC-I]), atypical carcinoid (also called well-differentiated neuroendocrine carcinoma [KC-II]), intermediate small-cell neuroendocrine carcinoma, and small-cell neuroendocrine carcinoma

564 (KC-III). Another proposed classification includes three categories of lung NETs: benign or low-grade malignant (typical carcinoid), low-grade malignant (atypical carcinoid), and high-grade malignant (poorly differentiated carcinoma of the large-cell or small-cell type). The WHO classification includes four general categories: typical carcinoid, atypical carcinoid, large-cell neuroendocrine carcinoma, and small-cell carcinoma. The ratio of typical to atypical carcinoids is 8–10:1, with the typical carcinoids comprising 1–2% of lung tumors, atypical 0.1–0.2%, large-cell neuroendocrine tumors 0.3%, and small-cell lung cancer 9.8% of all lung tumors. These different categories of lung NETs have different prognoses, varying from excellent for typical carcinoid to poor for small-cell neuroendocrine carcinomas. The occurrence of large-cell and small-cell lung carcinoids, but not typical or atypical lung carcinoids, is related to tobacco use. The 5-year survival is very much influenced by the classification of the tumor, with survival of 92–100% for patients with a typical carcinoid, 61–88% with an atypical carcinoid, 13–57% with a large-cell neuroendocrine tumor, and 5% with a small-cell lung cancer.

Gastric NET (Carcinoids) Gastric NETs (carcinoids) account for 3 of every 1000 gastric neoplasms and 1.3–2% of all carcinoids, and their relative frequency has increased three- to fourfold over the last five decades (2.2% in 1950 to 9.6% in 2000–2007, SEER data). At present, it is unclear whether this increase is due to better detection with the increased use of upper GI endoscopy or to a true increase in incidence. Gastric NETs (carcinoids) are classified into three different categories, and this has important implications for pathogenesis, prognosis, and treatment. Each originates from gastric enterochromaffin-like (ECL) cells, one of the six types of gastric neuroendocrine cells, in the gastric mucosa. Two subtypes are associated with hypergastrinemic states, either chronic atrophic gastritis (type I) (80% of all gastric NETs [carcinoids]) or Zollinger-Ellison syndrome, which is almost always a part of the MEN 1 syndrome (type II) (6% of all cases). These tumors generally pursue a benign course, with type I uncommonly (<10%) associated with metastases, whereas type II tumors are slightly more aggressive, with 10–30% associated with metastases. They are usually multiple, small, and infiltrate only to the submucosa. The third subtype of gastric NETs (carcinoids) (type III) (sporadic) occurs without hypergastrinemia (14–25% of all gastric carcinoids) and has an aggressive course, with 54–66% developing metastases. Sporadic carcinoids are usually single, large tumors; 50% have atypical histology, and they can be a cause of the carcinoid syndrome. Five-year survival is 99–100% in patients with type I, 60–90% in patients with type II, and 50% in patients with type III gastric NETs (carcinoids).

CLINICAL PRESENTATION OF NETS (CARCINOIDS)

GI/Lung NET (Carcinoid) Without the Carcinoid Syndrome The age of patients at diagnosis ranges from 10 to 93 years, with a mean age of 63 years for the small intestine and 66 years for the rectum. The presentation is diverse and is related to the site of origin and the extent of malignant spread. In the appendix, NETs (carcinoids) usually are found incidentally during surgery for suspected appendicitis. SI NETs (carcinoids) in the jejunioileum present with periodic abdominal pain (51%), intestinal obstruction with ileus/invagination (31%), an abdominal tumor (17%), or GI bleeding (11%). Because of the vagueness of the symptoms, the diagnosis usually is delayed approximately 2 years from onset of the symptoms, with a range up to 20 years. Duodenal, gastric, and rectal NETs (carcinoids) are most frequently found by chance at endoscopy. The most common symptoms of rectal carcinoids are melena/bleeding (39%), constipation (17%), and diarrhea (12%). Bronchial NETs (carcinoids) frequently are discovered as a lesion on a chest radiograph, and 31% of the patients are asymptomatic. Thymic NETs (carcinoids) present as anterior mediastinal masses, usually on chest radiograph or computed tomography (CT) scan. Ovarian and testicular NETs (carcinoids) usually present as masses discovered on physical examination or ultrasound. Metastatic NETs (carcinoids) in the liver frequently presents as hepatomegaly in a patient who may have minimal symptoms and nearly normal liver function test results.

GI-NETS (CARCINOIDS) WITH SYSTEMIC SYMPTOMS DUE TO SECRETED PRODUCTS

GI/lung NETs (carcinoids) immunocytochemically can contain numerous GI peptides: gastrin, insulin, somatostatin, motilin, neurotensin, tachykinins (substance K, substance P, neuropeptide K), glucagon, gastrin-releasing peptide, vasoactive intestinal peptide (VIP), PP, ghrelin, other biologically active peptides (ACTH, calcitonin, growth hormone), prostaglandins, and bioactive amines (serotonin). These substances may or may not be released in sufficient amounts to cause symptoms. In various studies of patients with GI-NETs (carcinoids), elevated serum levels of PP were found in 43%, motilin in 14%, gastrin in 15%, and VIP in 6%. Foregut NETs (carcinoids) are more likely to produce various GI peptides than are midgut NETs (carcinoids). Ectopic ACTH production causing Cushing's syndrome is seen increasingly with foregut carcinoids (respiratory tract primarily) and, in some series, has been the most common cause of the ectopic ACTH syndrome, accounting for 64% of all cases. Acromegaly due to growth hormone-releasing factor release occurs with foregut NETs (carcinoids), as does the somatostatinoma syndrome, but rarely occurs with duodenal NETs (carcinoids). The most common systemic syndrome with GI-NETs (carcinoids) is the carcinoid syndrome, which is discussed in detail in the next section.

CARCINOID SYNDROME

Clinical Features The cardinal features from a number of series at presentation as well as during the disease course are shown in Table 113-7. Flushing and diarrhea are the two most common symptoms, occurring in a mean of 69–70% of patients initially and in up to 78% of patients during the course of the disease. The characteristic flush is of sudden onset; it is a deep red or violaceous erythema of the upper body, especially the neck and face, often associated with a feeling of warmth and occasionally associated with pruritus, lacrimation, diarrhea, or facial edema. Flushes may be precipitated by stress; alcohol; exercise; certain foods, such as cheese; or certain agents, such as catecholamines, pentagastrin, and serotonin reuptake inhibitors. Flushing episodes may be brief, lasting 2–5 min, especially initially, or may last hours, especially later in the disease course. Flushing usually is associated with metastatic midgut NETs (carcinoids) but can also occur with foregut NETs (carcinoids). With bronchial NETs (carcinoids), the flushes frequently are prolonged for hours to days, reddish in color, and associated with salivation, lacrimation, diaphoresis, diarrhea, and hypotension. The flush associated with gastric NETs (carcinoids) can also be reddish in color, but with a patchy distribution over the face and neck, although the classic flush seen with midgut NETs (carcinoids) can also be seen with gastric NETs (carcinoids). It may be provoked by food and have accompanying pruritus.

Diarrhea usually occurs with flushing (85% of cases). The diarrhea usually is described as watery, with 60% of patients having <1 L/d of diarrhea. Steatorrhea is present in 67%, and in 46%, it is >15 g/d (normal <7 g). Abdominal pain may be present with the diarrhea or independently in 10–34% of cases.

Cardiac manifestations occur initially in 11–40% (mean 26%) of patients with carcinoid syndrome and in 14–41% (mean 30%) at some time in the disease course. The cardiac disease is due to the formation of fibrotic plaques (composed of smooth-muscle cells, myofibroblasts, and elastic tissue) involving the endocardium, primarily on the right side, although lesions on the left side also occur occasionally, especially if a patent foramen ovale exists. The dense fibrous deposits are most commonly on the ventricular aspect of the tricuspid valve and less commonly on the pulmonary valve cusps. They can result in constriction of the valves, and pulmonic stenosis is usually predominant, whereas the tricuspid valve is often fixed open, resulting in regurgitation predominating. Overall, in patients with carcinoid heart disease, 90–100% have tricuspid insufficiency, 43–59% have tricuspid stenosis, 50–81% have pulmonary insufficiency, 25–59% have pulmonary stenosis, and 11% (0–25%) left-side lesions. Up to 80% of patients with cardiac lesions develop heart failure. Lesions on the left side are much less extensive, occur in 30% at autopsy, and most frequently affect the mitral valve. Up to 80% of patients with cardiac lesions have evidence of heart failure. At diagnosis in various series, 27–43% of patients are

in New York Heart Association class I, 30–40% are in class II, 13–31% are in class III, and 3–12% are in class IV. At present, carcinoid heart disease is reported to be decreasing in frequency and severity, with mean occurrence in 20% of patients and occurrence in as few as 3–4% in some reports. Whether this decrease is due to the widespread use of somatostatin analogues, which control the release of bioactive agents thought involved in mediating the heart disease, is unclear.

Other clinical manifestations include wheezing or asthma-like symptoms (8–18%), pellagra-like skin lesions (2–25%), and impaired cognitive function. A variety of noncardiac problems due to increased fibrous tissue have been reported, including retroperitoneal fibrosis causing urethral obstruction, Peyronie's disease of the penis, intraabdominal fibrosis, and occlusion of the mesenteric arteries or veins.

Pathobiology Carcinoid syndrome occurred in 8% of 8876 patients with GI-NETs (carcinoids), with a rate of 1.7–18.4% in different studies. It occurs only when sufficient concentrations of products secreted by the tumor reach the systemic circulation. In 91–100% of cases, this occurs after distant metastases to the liver. Rarely, primary GI-NETs (carcinoids) with nodal metastases with extensive retroperitoneal invasion, pNETs (carcinoids) with retroperitoneal lymph nodes, or NETs (carcinoids) of the lung or ovary with direct access to the systemic circulation can cause the carcinoid syndrome without hepatic metastases. All GI-NETs (carcinoids) do not have the same propensity to metastasize and cause the carcinoid syndrome (Table 113-3). Midgut NETs (carcinoids) account for 57–67% of cases of carcinoid syndrome, foregut NETs (carcinoids) for 0–33%, hindgut for 0–8%, and an unknown primary location for 2–26% (Tables 113-3 and 113-7).

One of the main secretory products of GI-NETs (carcinoids) involved in the carcinoid syndrome is serotonin (5-HT) (Fig. 113-1), which is synthesized from tryptophan. Up to 50% of dietary tryptophan can be used in this synthetic pathway by tumor cells, and this can result in inadequate supplies for conversion to niacin; hence, some patients (2.5%) develop pellagra-like lesions. Serotonin has numerous biologic effects, including stimulating intestinal secretion with inhibition of absorption, stimulating increases in intestinal motility, and stimulating fibrogenesis. In various studies, 56–88% of all GI-NETs (carcinoids) were associated with serotonin overproduction; however, 12–26% of the patients did not have the carcinoid syndrome. In one study, platelet serotonin was elevated in 96% of patients with midgut NETs (carcinoids), 43% with foregut tumors, and 0% with hindgut tumors. In 90–100% of patients with the carcinoid syndrome, there is evidence of serotonin overproduction. Serotonin is thought to be predominantly responsible for the diarrhea. Patients with the carcinoid syndrome have increased colonic motility with a shortened transit time and possibly a secretory/absorptive alteration that is compatible with the known actions of serotonin in the gut mediated primarily through 5-HT₃ and, to a lesser degree, 5-HT₄ receptors. Serotonin receptor antagonists (especially 5-HT₃ antagonists) relieve the diarrhea in many, but not all, patients. A tryptophan 5-hydroxylase inhibitor, LX-1031, which inhibits serotonin synthesis in peripheral tissues, is reported to cause a 44% decrease in bowel movement frequency and a 20% improvement in stool form in patients with the carcinoid syndrome. Additional studies suggest that tachykinins may be important mediators of diarrhea in some patients. In one study, plasma tachykinin levels correlated with symptoms of diarrhea. Serotonin does not appear to be involved in the flushing because serotonin receptor antagonists do not relieve flushing. In patients with gastric carcinoids, the characteristic red, patchy pruritic flush is thought due to histamine release because H₁ and H₂ receptor antagonists can prevent it. Numerous studies have shown that tachykinins (substance P, neuropeptide K) are stored in GI-NETs (carcinoids) and released during flushing. However, some studies have demonstrated that octreotide can relieve the flushing induced by pentagastrin in these patients without altering the stimulated increase in plasma substance P, suggesting that other mediators must be involved in the flushing. A correlation between plasma tachykinin levels (but not substance P levels) and flushing has been reported. Prostaglandin release could be involved in mediating either the diarrhea or flush, but conflicting data exist.

Both histamine and serotonin may be responsible for the wheezing as well as the fibrotic reactions involving the heart, causing Peyronie's disease and intraabdominal fibrosis.

The exact mechanism of the heart disease remains unclear, although increasing evidence supports a central role for serotonin. Patients with heart disease have higher plasma levels of neurokinin A, substance P, plasma atrial natriuretic peptide (ANP), pro-brain natriuretic peptide, chromogranin A, and activin A as well as higher urinary 5-HIAA excretion.

The valvular heart disease caused by the appetite-suppressant drug dexfenfluramine is histologically indistinguishable from that observed in carcinoid disease. Furthermore, ergot-containing dopamine receptor agonists used for Parkinson's disease (pergolide, cabergoline) cause valvular heart disease that closely resembles that seen in the carcinoid syndrome. Furthermore, in animal studies, the formation of valvular plaques/fibrosis occurs after prolonged treatment with serotonin as well as in animals with a deficiency of the 5-HIAA transporter gene, which results in an inability to inactivate serotonin. Metabolites of fenfluramine, as well as the dopamine receptor agonists, have high affinity for serotonin receptor subtype 5-HT_{2B} receptors, whose activation is known to cause fibroblast mitogenesis. Serotonin receptor subtypes 5-HT_{1B,1D,2A,2B} normally are expressed in human heart valve interstitial cells. High levels of 5-HT_{2B} receptors are known to occur in heart valves and occur in cardiac fibroblasts and cardiomyocytes. Studies of cultured interstitial cells from human cardiac valves have demonstrated that these valvulopathic drugs induce mitogenesis by activating 5-HT_{2B} receptors and stimulating upregulation of transforming growth factor β and collagen biosynthesis. These observations support the conclusion that serotonin overproduction by GI-NETs (carcinoids) is important in mediating the valvular changes, possibly by activating 5-HT_{2B} receptors in the endocardium. Both the magnitude of serotonin overproduction and prior chemotherapy are important predictors of progression of the heart disease, whereas patients with high plasma levels of ANP have a worse prognosis. Plasma connective tissue growth factor levels are elevated in many fibrotic conditions; elevated levels occur in patients with carcinoid heart disease and correlate with the presence of right ventricular dysfunction and the extent of valvular regurgitation in patients with GI-NETs (carcinoids).

Patients may develop either a typical or, rarely, an atypical carcinoid syndrome (Fig. 113-1). In patients with the typical form, which characteristically is caused by midgut NETs (carcinoids), the conversion of tryptophan to 5-HTP is the rate-limiting step (Fig. 113-1). Once 5-HTP is formed, it is rapidly converted to 5-HT and stored in secretory granules of the tumor or in platelets. A small amount remains in plasma and is converted to 5-HIAA, which appears in large amounts in the urine. These patients have an expanded serotonin pool size, increased blood and platelet serotonin, and increased urinary 5-HIAA. Some GI-NETs (carcinoids) cause an atypical carcinoid syndrome that is thought to be due to a deficiency in the enzyme dopa decarboxylase; thus, 5-HTP cannot be converted to 5-HT (serotonin), and 5-HTP is secreted into the bloodstream (Fig. 113-1). In these patients, plasma serotonin levels are normal but urinary levels may be increased because some 5-HTP is converted to 5-HT in the kidney. Characteristically, urinary 5-HTP and 5-HT are increased, but urinary 5-HIAA levels are only slightly elevated. Foregut carcinoids are the most likely to cause an atypical carcinoid syndrome; however, they also can cause a typical carcinoid syndrome.

One of the most immediate life-threatening complications of the carcinoid syndrome is the development of a carcinoid crisis. This is more common in patients who have intense symptoms or have greatly increased urinary 5-HIAA levels (i.e., >200 mg/d). The crisis may occur spontaneously; however, it is usually provoked by procedures such as anesthesia, chemotherapy, surgery, biopsy, endoscopy, or radiologic examinations such as during biopsies, hepatic artery embolization, and vessel catheterization. It can be provoked by stress or procedures as mild as repeated palpation of the tumor during physical examination. Patients develop intense flushing, diarrhea, abdominal pain, cardiac abnormalities including tachycardia, hypertension, or hypotension, and confusion or stupor. If not adequately treated, this can be a terminal event.

The diagnosis of carcinoid syndrome relies on measurement of urinary or plasma serotonin or its metabolites in the urine. The measurement of 5-HIAA is used most frequently. False-positive elevations may occur if the patient is eating serotonin-rich foods such as bananas, pineapples, walnuts, pecans, avocados, or hickory nuts or is taking certain medications (cough syrup containing guaifenesin, acetaminophen, salicylates, serotonin reuptake inhibitors, or L-dopa). The normal range for daily urinary 5-HIAA excretion is 2–8 mg/d. Serotonin overproduction was noted in 92% of patients with carcinoid syndrome in one study, and in another study, 5-HIAA had 73% sensitivity and 100% specificity for carcinoid syndrome. Serotonin overproduction is *not* synonymous with the presence of clinical carcinoid syndrome because 12–26% of patients with serotonin overproduction do not have clinical evidence of the carcinoid syndrome.

Most physicians use only the urinary 5-HIAA excretion rate; however, plasma and platelet serotonin levels, if available, may provide additional information. Platelet serotonin levels are more sensitive than urinary 5-HIAA but are not generally available. A single plasma 5-HIAA determination was found to correlate with the 24-h urinary values, raising the possibility that this could replace the standard urinary collection because of its greater convenience and avoidance of incomplete or improper collections. Because patients with foregut NETs (carcinoids) may produce an atypical carcinoid syndrome, if this syndrome is suspected and the urinary 5-HIAA is minimally elevated or normal, other urinary metabolites of tryptophan, such as 5-HTP and 5-HT, should be measured (Fig. 113-1).

Flushing occurs in a number of other diseases, including systemic mastocytosis, chronic myeloid leukemia with increased histamine release, menopause, reactions to alcohol or glutamate, and side effects of chlorpromazine, calcium channel blockers, and nicotinic acid. None of these conditions cause increased urinary 5-HIAA.

The diagnosis of carcinoid tumor can be suggested by the carcinoid syndrome, recurrent abdominal symptoms in a healthy-appearing individual, or the discovery of hepatomegaly or hepatic metastases associated with minimal symptoms. Ileal NETs (carcinoids), which make up 25% of all clinically detected carcinoids, should be suspected in patients with bowel obstruction, abdominal pain, flushing, or diarrhea.

Serum chromogranin A levels are elevated in 56–100% of patients with GI-NETs (carcinoids), and the level correlates with tumor bulk. Serum chromogranin A levels are not specific for GI-NETs (carcinoids) because they are also elevated in patients with pNETs and other NETs. Furthermore, a major problem is caused by potent acid antisecretory drugs such as proton pump inhibitors (omeprazole and related drugs) because they almost invariably cause elevation of plasma chromogranin A levels; the elevation occurs rapidly (3–5 days) with continued use, and the elevated levels overlap with the levels seen in many patients with NETs. Plasma neuron-specific enolase levels are also used as a marker of GI-NETs (carcinoids) but are less sensitive than chromogranin A, being increased in only 17–47% of patients. Newer markers have been proposed including pancreastatin (a chromogranin A breakdown product) and activin A. The former is not affected by proton pump inhibitors; however, its sensitivity and specificity are not established. Plasma activin elevations are reported to correlate with the presence of cardiac disease with a sensitivity of 87% and specificity of 57%.

TREATMENT CARCINOID SYNDROME AND NONMETASTATIC GASTROINTESTINAL NEUROENDOCRINE TUMORS (CARCINOIDS)

CARCINOID SYNDROME

Treatment includes avoiding conditions that precipitate flushing, dietary supplementation with nicotinamide, treatment of heart failure with diuretics, treatment of wheezing with oral bronchodilators, and control of the diarrhea with antidiarrheal agents such as

loperamide and diphenoxylate. If patients still have symptoms, serotonin receptor antagonists or somatostatin analogues (Fig. 113-2) are the drugs of choice.

There are 14 subclasses of serotonin receptors, and antagonists for many are not available. The 5-HT₁ and 5-HT₂ receptor antagonists methysergide, cyproheptadine, and ketanserin have all been used to control the diarrhea but usually do not decrease flushing. The use of methysergide is limited because it can cause or enhance retroperitoneal fibrosis. Ketanserin diminishes diarrhea in 30–100% of patients. 5-HT₃ receptor antagonists (ondansetron, tropisetron, alosetron) can control diarrhea and nausea in up to 100% of patients and occasionally ameliorate the flushing. A combination of histamine H₁ and H₂ receptor antagonists (i.e., diphenhydramine and cimetidine or ranitidine) may control flushing in patients with foregut carcinoids. The tryptophan 5-hydroxylase inhibitor telotristat etiprate decreased bowel frequency in 44% and improved stool consistency in 20%.

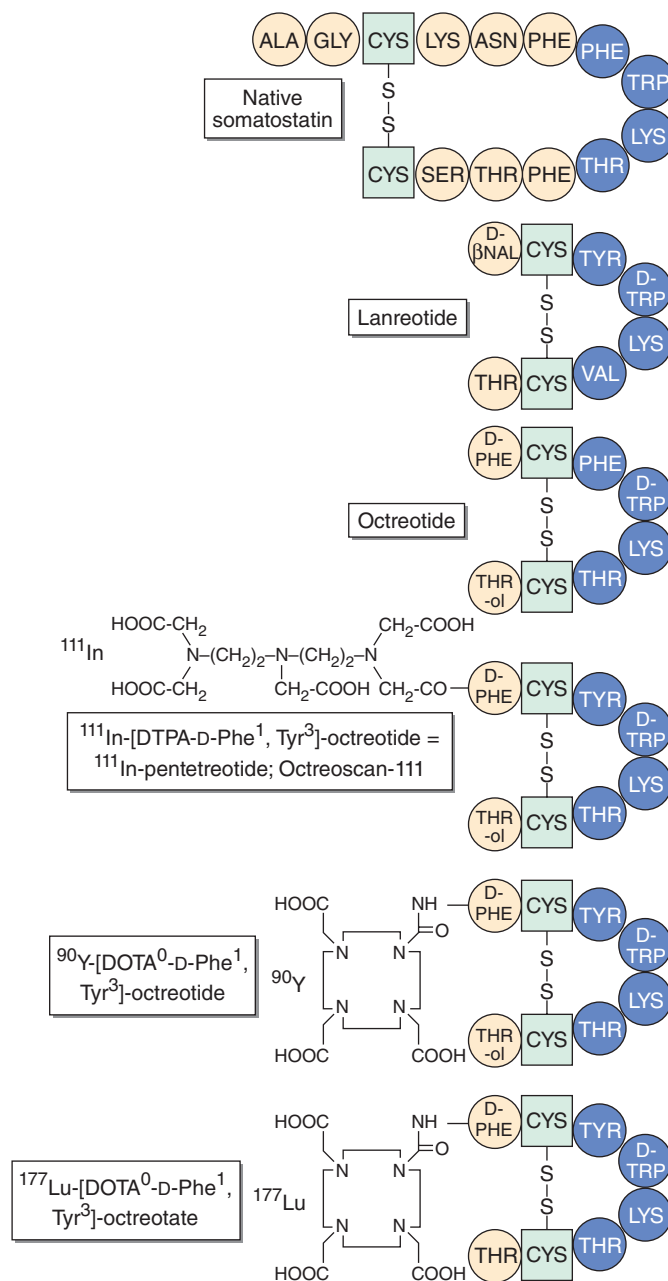


FIGURE 113-2 Structure of somatostatin and synthetic analogues used for diagnostic or therapeutic indications.

Synthetic analogues of somatostatin (octreotide, lanreotide) are now the most widely used agents to control the symptoms of patients with carcinoid syndrome (Fig. 113-2). These drugs are effective at relieving symptoms and decreasing urinary 5-HIAA levels in patients with this syndrome. Octreotide-LAR and lanreotide-SR/autogel (Somatuline) (sustained-release formulations allowing monthly injections) control symptoms in 74% and 68% of patients, respectively, with carcinoid syndrome and show a biochemical response in 51% and 64%, respectively. Patients with mild to moderate symptoms usually are treated initially with octreotide 100 µg SC every 8 h and then begun on the long-acting monthly depot forms (octreotide-LAR or lanreotide-autogel). Forty percent of patients escape control after a median time of 4 months, and the depot dosage may have to be increased as well as supplemented with the shorter-acting formulation, SC octreotide. Pasireotide (SOM230) is a somatostatin analogue with broader selectivity (high-affinity somatostatin receptors [$ss_{1,2,3,5}$] than octreotide/lanreotide ($ss_{2,5}$). In a phase II study of patients with refractory carcinoid syndrome, pasireotide controlled symptoms in 27%.

Carcinoid heart disease is associated with a decreased mean survival (3.8 years), and therefore, it should be sought for and carefully assessed in all patients with carcinoid syndrome. Transthoracic echocardiography remains a key element in establishing the diagnosis of carcinoid heart disease and determining the extent and type of cardiac abnormalities. Treatment with diuretics and somatostatin analogues can reduce the negative hemodynamic effects and secondary heart failure. It remains unclear whether long-term treatment with these drugs will decrease the progression of carcinoid heart disease. Balloon valvuloplasty for stenotic valves or cardiac valve surgery may be required.

In patients with carcinoid crises, somatostatin analogues are effective at both treating the condition and preventing their development during known precipitating events such as surgery, anesthesia, chemotherapy, and stress. It is recommended that octreotide 150–250 µg SC every 6 to 8 h be used 24–48 h before anesthesia and then continued throughout the procedure.

Currently, sustained-release preparations of both octreotide (octreotide-LAR [long-acting release], 10, 20, 30 mg) and lanreotide (lanreotide-PR [prolonged release, lanreotide-autogel], 60, 90, 120 mg) are available and widely used because their use greatly facilitates long-term treatment. Octreotide-LAR (30 mg/month) gives a plasma level ≥ 1 ng/mL for 25 days, whereas this requires three to six injections a day of the non-sustained-release form. Lanreotide-autogel (Somatuline) is given every 4–6 weeks.

Short-term side effects occur in up to one-half of patients. Pain at the injection site and side effects related to the GI tract (59% discomfort, 15% nausea, diarrhea) are the most common. They are usually short-lived and do not interrupt treatment. Important long-term side effects include gallstone formation, steatorrhea, and deterioration in glucose tolerance. The overall incidence of gallstones/biliary sludge in one study was 52%, with 7% having symptomatic disease that required surgical treatment.

Interferon α is reported to be effective in controlling symptoms of the carcinoid syndrome either alone or combined with hepatic artery embolization. With interferon α alone, the clinical response rate is 30–70%, and with interferon α with hepatic artery embolization, diarrhea was controlled for 1 year in 43% and flushing was controlled in 86%. Side effects develop in almost all patients, with the most frequent being a flu-like syndrome (80–100%), followed by anorexia and fatigue, even though these frequently improve with continued treatment. Other more severe side effects include bone marrow toxicity, hepatotoxicity, autoimmune disorders, and rarely CNS side effects (depression, mental disorders, visual problems).

Hepatic artery embolization alone or with chemotherapy (chemoembolization) has been used to control the symptoms of carcinoid syndrome. Embolization alone is reported to control symptoms in up to 76% of patients, and chemoembolization (5-fluorouracil, doxorubicin, cisplatin, mitomycin) controls symptoms in 60–75% of

patients. Hepatic artery embolization can have major side effects, including nausea, vomiting, pain, and fever. In two studies, 5–7% of patients died from complications of hepatic artery occlusion.

Other drugs have been used successfully in small numbers of patients to control the symptoms of carcinoid syndrome. Parachlorophenylalanine can inhibit tryptophan hydroxylase and therefore the conversion of tryptophan to 5-HTP. However, its severe side effects, including psychiatric disturbances, make it intolerable for long-term use. α -Methyldopa inhibits the conversion of 5-HTP to 5-HT, but its effects are only partial.

Peptide radioreceptor therapy (using radiotherapy with radiolabeled somatostatin analogues), the use of radiolabeled microspheres, and other methods for treatment of advanced metastatic disease may facilitate control of the carcinoid syndrome and are discussed in a later section dealing with treatment of advanced disease.

GI-NETS (CARCINOIDS) (NONMETASTATIC)

Surgery is the only potentially curative therapy. Because with most GI-NETs (carcinoids), the probability of metastatic disease increases with increasing size, the extent of surgical resection is determined accordingly. With appendiceal NETs (carcinoids) < 1 cm, simple appendectomy was curative in 103 patients followed for up to 35 years. With rectal NETs (carcinoids) < 1 cm, local resection is curative. With SI NETs (carcinoids) < 1 cm, there is not complete agreement. Because 15–69% of SI NETs (carcinoids) this size have metastases in different studies, some recommend a wide resection with en bloc resection of the adjacent lymph-bearing mesentery. If the tumor is > 2 cm for rectal, appendiceal, or SI NETs (carcinoids), a full cancer operation should be done. This includes a right hemicolectomy for appendiceal NETs (carcinoids), an abdominoperineal resection or low anterior resection for rectal NETs (carcinoids), and an en bloc resection of adjacent lymph nodes for SI NETs (carcinoids). For appendiceal NETs (carcinoids) 1–2 cm in diameter, a simple appendectomy is proposed by some, whereas others favor a formal right hemicolectomy. For 1–2 cm rectal NETs (carcinoids), it is recommended that a wide, local, full-thickness excision be performed.

With type I or II gastric NETs (carcinoids), which are usually < 1 cm, endoscopic removal is recommended. In type I or II gastric carcinoids, if the tumor is > 2 cm or if there is local invasion, some recommend total gastrectomy, whereas others recommend antrectomy in type I to reduce the hypergastrinemia, which has led to regression of the carcinoids in a number of studies. For types I and II gastric NETs (carcinoids) of 1–2 cm, there is no agreement, with some recommending endoscopic treatment followed by chronic somatostatin treatment and careful follow-up and others recommending surgical treatment. With type III gastric NETs (carcinoids) > 2 cm, excision and regional lymph node clearance are recommended. Most tumors < 1 cm are treated endoscopically.

Resection of isolated or limited hepatic metastases may be beneficial and will be discussed in a later section on treatment of advanced disease.

PANCREATIC NEUROENDOCRINE TUMORS

Functional pNETs usually present clinically with symptoms due to the hormone-excess state (Table 113-2). Only late in the course of the disease does the tumor per se cause prominent symptoms such as abdominal pain. In contrast, all the symptoms due to nonfunctional pNETs are due to the tumor per se. The overall result of this is that some functional pNETs may present with severe symptoms with a small or undetectable primary tumor, whereas nonfunctional tumors usually present late in the disease course with large tumors, which are frequently metastatic. The mean delay between onset of continuous symptoms and diagnosis of a functional pNET syndrome is 4–7 years. Therefore, the diagnoses frequently are missed for extended periods.

TREATMENT PANCREATIC NEUROENDOCRINE TUMOR (GENERAL POINTS)

Treatment of pNETs requires two different strategies. First, treatment must be directed at the hormone-excess state such as the gastric acid hypersecretion in gastrinomas or the hypoglycemia in insulinomas. Ectopic hormone secretion usually causes the presenting symptoms and can cause life-threatening complications. Second, with all the tumors except insulinomas, >50% are malignant (Table 113-2); therefore, treatment must also be directed against the tumor per se. Because in many patients these tumors are not surgically curable due to the presence of advanced disease at diagnosis, surgical resection for cure, which addresses both treatment aspects, is often not possible.

GASTRINOMA (ZOLLINGER-ELLISON SYNDROME)

A gastrinoma is an NET that secretes gastrin; the resultant hypergastrinemia causes gastric acid hypersecretion (Zollinger-Ellison syndrome [ZES]). The chronic hypergastrinemia results in marked gastric acid hypersecretion and growth of the gastric mucosa with increased numbers of parietal cells and proliferation of gastric ECL cells. The gastric acid hypersecretion characteristically causes peptic ulcer disease (PUD), often refractory and severe, as well as diarrhea. The most common presenting symptoms are abdominal pain (70–100%), diarrhea (37–73%), and gastroesophageal reflux disease (GERD) (30–35%); 10–20% of patients have diarrhea only. Although peptic ulcers may occur in unusual locations, most patients have a typical duodenal ulcer. Important observations that should suggest this diagnosis include PUD with diarrhea; PUD in an unusual location or with multiple ulcers; PUD refractory to treatment or persistent; PUD associated with prominent gastric folds; PUD associated with findings suggestive of MEN 1 (endocrinopathy, family history of ulcer or endocrinopathy, nephrolithiasis); and PUD without *Helicobacter pylori* present. *H. pylori* is present in >90% of idiopathic peptic ulcers but is present in <50% of patients with gastrinomas. Chronic unexplained diarrhea also should suggest ZES.

Approximately 20–25% of patients with ZES have MEN 1 (MEN1/ZES), and in most cases, hyperparathyroidism is present before the ZES develops. These patients are treated differently from those without MEN 1 (sporadic ZES); therefore, MEN 1 should be sought in all patients with ZES by family history and by measuring plasma ionized calcium and prolactin levels and plasma hormone levels (parathormone, growth hormone).

Most gastrinomas (50–90%) in sporadic ZES are present in the duodenum, followed by the pancreas (10–40%) and other intraabdominal sites (mesentery, lymph nodes, biliary tract, liver, stomach, ovary). Rarely, the tumor may involve extraabdominal sites (heart, lung cancer). In MEN 1/ZES the gastrinomas are also usually in the duodenum (70–90%), followed by the pancreas (10–30%), and are almost always multiple. About 60–90% of gastrinomas are malignant (Table 113-2) with metastatic spread to lymph nodes and liver. Distant metastases to bone occur in 12–30% of patients with liver metastases.

Diagnosis The diagnosis of ZES requires the demonstration of inappropriate fasting hypergastrinemia, usually by demonstrating hypergastrinemia occurring with an increased basal gastric acid output (BAO) (hyperchlorhydria). More than 98% of patients with ZES have fasting hypergastrinemia, although in 40–60% the level may be elevated less than tenfold. Therefore, when the diagnosis is suspected, a fasting gastrin is usually the initial test performed. It is important to remember that potent gastric acid suppressant drugs such as proton pump inhibitors (PPIs) (omeprazole, esomeprazole, pantoprazole, lansoprazole, rabeprazole) can suppress acid secretion sufficiently to cause hypergastrinemia; because of their prolonged duration of action, these drugs have to be tapered or frequently discontinued for a week before the gastrin determination. Withdrawal of PPIs should be performed carefully because PUD complications can rapidly develop in some patients and is best done in consultation with GI units with

experience in this area. The widespread use of PPIs can confound the diagnosis of ZES by raising a false-positive diagnosis by causing hypergastrinemia in a patient being treated with idiopathic PUD (without ZES) and lead to a false-negative diagnosis because at routine doses used to treat patients with idiopathic PUD, PPIs control symptoms in most ZES patients and thus mask the diagnosis. If ZES is suspected and the gastrin level is elevated, it is important to show that it is increased when gastric pH is ≤ 2.0 because physiologically hypergastrinemia secondary to achlorhydria (atrophic gastritis, pernicious anemia) is one of the most common causes of hypergastrinemia. Nearly all ZES patients have a fasting pH ≤ 2 when off antisecretory drugs. If the fasting gastrin is >1000 pg/mL (increased tenfold) and the pH is ≤ 2.0 , which occurs in 40–60% of patients with ZES, the diagnosis of ZES is established after the possibility of retained antrum syndrome has been ruled out by history. In patients with hypergastrinemia with fasting gastrins <1000 pg/mL (<10-fold increased) and gastric pH ≤ 2.0 , other conditions, such as *H. pylori* infections, antral G-cell hyperplasia/hyperfunction, gastric outlet obstruction, and, rarely, renal failure, can masquerade as ZES. To establish the diagnosis in this group, a determination of BAO and a secretin provocative test should be done. In patients with ZES without previous gastric acid-reducing surgery, the BAO is usually (>90%) elevated (i.e., >15 mEq/h). The secretin provocative test is usually positive, with the criterion of a >120-pg/mL increase over the basal level having the highest sensitivity (94%) and specificity (100%). Unfortunately the diagnosis of ZES is becoming increasing more difficult. This is due not only to the widespread use of PPIs (leading to false-positive results as well as masking ZES presentation), but also recent studies demonstrate that many of the commercial gastrin kits that are used by most laboratories to measure fasting serum gastrin levels are not reliable. In one study, 7 of the 12 tested commercial gastrin kits inaccurately assessed the true serum concentration of gastrin primarily because the antibodies used had inappropriate specificity for the different circulating forms of gastrin and were not adequately validated. Both underestimation and overestimation of fasting serum gastrin levels occurred using these commercial kits. To circumvent this problem, it is either necessary to use one of the five reliable kits identified or, alternatively, to refer the patient to a center with expertise in making the diagnosis in your area, or if this is not possible, to contact such a center and use the gastrin assay they recommend. An accurate gastrin assay is essential for accurate measurement of fasting serum gastrin level as well as for assessing gastrin levels during the secretin provocative test, and thus, the diagnosis of ZES cannot reliably be made without one.

TREATMENT ZOLLINGER-ELLISON SYNDROME

Gastric acid hypersecretion in patients with ZES can be controlled in almost every case by oral gastric antisecretory drugs. Because of their long duration of action and potency, which allows dosing once or twice a day, the PPIs (H^+ , K^+ -ATPase inhibitors) are the drugs of choice. Histamine H_2 -receptor antagonists are also effective, although more frequent dosing (q 4–8 h) and high doses are required. In patients with MEN 1/ZES with hyperparathyroidism, correction of the hyperparathyroidism increases the sensitivity to gastric antisecretory drugs and decreases the basal acid output. Long-term treatment with PPIs (>15 years) has proved to be safe and effective, without development of tachyphylaxis. Although patients with ZES, especially those with MEN 1/ZES, more frequently develop gastric NETs (carcinoids), no data suggest that the long-term use of PPIs increases this risk in these patients. With long-term PPI use in ZES patients, vitamin B_{12} deficiency can develop; thus, vitamin B_{12} levels should be assessed during follow-up. Epidemiologic studies suggest that long-term PPI use may be associated with an increased incidence of bone fractures; however, at present, there is no such report in ZES patients.

With the increased ability to control acid hypersecretion, more than 50% of patients who are not cured (>60% of patients) will die from tumor-related causes. At presentation, careful imaging studies are essential to localize the extent of the tumor to determine the appropriate treatment. A third of patients present with hepatic

metastases, and in <15% of those patients, the disease is limited, so that surgical resection may be possible. Surgical short-term cure is possible in 60% of all patients without MEN 1/ZES or liver metastases (40% of all patients) and in 30% of patients long term. In patients with MEN 1/ZES, long-term surgical cure is rare because the tumors are multiple, frequently with lymph node metastases. Surgical studies demonstrate that successful resection of the gastrinoma not only decreases the chances of developing liver metastases but also increases the disease-related survival rate. Therefore, all patients with gastrinomas without MEN 1/ZES or a medical condition that limits life expectancy should undergo surgery by a surgeon experienced in the treatment of these disorders.

INSULINOMAS

An insulinoma is an NET of the pancreas that is thought to be derived from beta cells that ectopically secrete insulin, which results in hypoglycemia. The average age of occurrence is 40–50 years old. The most common clinical symptoms are due to the effect of the hypoglycemia on the CNS (neuroglycemic symptoms) and include confusion, headache, disorientation, visual difficulties, irrational behavior, and even coma. Also, most patients have symptoms due to excess catecholamine release secondary to the hypoglycemia, including sweating, tremor, and palpitations. Characteristically, these attacks are associated with fasting.

Insulinomas are generally small (>90% are <2 cm) and usually not multiple (90%); only 5–15% are malignant, and they almost invariably occur only in the pancreas, distributed equally in the pancreatic head, body, and tail.

Insulinomas should be suspected in all patients with hypoglycemia, especially when there is a history suggesting that attacks are provoked by fasting, or with a family history of MEN 1. Insulin is synthesized as proinsulin, which consists of a 21-amino-acid α chain and a 30-amino-acid β chain connected by a 33-amino-acid connecting peptide (C peptide). In insulinomas, in addition to elevated plasma insulin levels, elevated plasma proinsulin levels are found, and C-peptide levels are elevated.

Diagnosis The diagnosis of insulinoma requires the demonstration of an elevated plasma insulin level at the time of hypoglycemia. A number of other conditions may cause fasting hypoglycemia, such as the inadvertent or surreptitious use of insulin or oral hypoglycemic agents, severe liver disease, alcoholism, poor nutrition, and other extrapancreatic tumors. Furthermore, postprandial hypoglycemia can be caused by a number of conditions that confuse the diagnosis of insulinoma. Particularly important here is the increased occurrence of hypoglycemia after gastric bypass surgery for obesity, which is now widely performed. A new entity, insulinomatosis, was described that can cause hypoglycemia and mimic insulinomas. It occurs in 10% of patients with persistent hyperinsulinemic hypoglycemia and is characterized by the occurrence of multiple macro-/microadenomas expressing insulin, and it is not clear how to distinguish this entity from insulinoma preoperatively. The most reliable test to diagnose insulinoma is a fast up to 72 h with serum glucose, C-peptide, proinsulin, and insulin measurements every 4–8 h. If at any point the patient becomes symptomatic or glucose levels are persistently below <2.2 mmol/L (40 mg/dL), the test should be terminated, and repeat samples for the above studies should be obtained before glucose is given. Some 70–80% of patients will develop hypoglycemia during the first 24 h, and 98% by 48 h. In nonobese normal subjects, serum insulin levels should decrease to <43 pmol/L (<6 μ U/mL) when blood glucose decreases to <2.2 mmol/L (<40 mg/dL) and the ratio of insulin to glucose is <0.3 (in mg/dL). In addition to having an insulin level >6 μ U/mL when blood glucose is <40 mg/dL, some investigators also require an elevated C-peptide and serum proinsulin level, an insulin/glucose ratio >0.3, and a decreased plasma β -hydroxybutyrate level for the diagnosis of insulinomas. Surreptitious use of insulin or hypoglycemic agents may be difficult to distinguish from insulinomas. The combination of proinsulin levels (normal in exogenous insulin/hypoglycemic agent users), C-peptide levels (low in exogenous insulin users), antibodies to insulin (positive in exogenous insulin users), and measurement of sulfonylurea levels in serum or plasma will allow the correct diagnosis to be made. The diagnosis of

insulinoma has been complicated by the introduction of specific insulin assays that do not also interact with proinsulin, as do many of the older radioimmunoassays (RIAs), and therefore give lower plasma insulin levels. The increased use of these specific insulin assays has resulted in increased numbers of patients with insulinomas having lower plasma insulin values (<6 μ U/mL) than levels proposed to be characteristic of insulinomas by RIA. In these patients, the assessment of proinsulin and C-peptide levels at the time of hypoglycemia is particularly helpful for establishing the correct diagnosis. An elevated proinsulin level when the fasting glucose level is <45 mg/dL is sensitive and specific.

TREATMENT INSULINOMAS

Only 5–15% of insulinomas are malignant; therefore, after appropriate imaging (see below), surgery should be performed. In different studies, 75–100% of patients are cured by surgery. Before surgery, the hypoglycemia can be controlled by frequent small meals and the use of diazoxide (150–800 mg/d). Diazoxide is a benzothiadiazide whose hyperglycemic effect is attributed to inhibition of insulin release. Its side effects are sodium retention and GI symptoms such as nausea. Approximately 50–60% of patients respond to diazoxide. Other agents effective in some patients to control the hypoglycemia include verapamil and diphenylhydantoin. Long-acting somatostatin analogues such as octreotide and lanreotide are acutely effective in 40% of patients. However, octreotide must be used with care because it inhibits growth hormone secretion and can alter plasma glucagon levels; therefore, in some patients, it can worsen the hypoglycemia.

For the 5–15% of patients with malignant insulinomas, these drugs or somatostatin analogues are used initially. In a small number of patients with insulinomas, some with malignant tumors, mammalian target of rapamycin (mTOR) inhibitors (everolimus, rapamycin) are reported to control the hypoglycemia. If they are not effective, various antitumor treatments such as hepatic arterial embolization, chemoembolization, chemotherapy, and peptide receptor radiotherapy have been used (see below).

Insulinomas, which are usually benign (>90%) and intrapancreatic in location, are increasingly resected using a laparoscopic approach, which has lower morbidity rates. This approach requires that the insulinoma be localized on preoperative imaging studies.

GLUCAGONOMAS

A glucagonoma is NET of the pancreas that secretes excessive amounts of glucagon, which causes a distinct syndrome characterized by dermatitis, glucose intolerance or diabetes, and weight loss. Glucagonomas principally occur between 45 and 70 years of age. The tumor is clinically heralded by a characteristic dermatitis (migratory necrolytic erythema) (67–90%), accompanied by glucose intolerance (40–90%), weight loss (66–96%), anemia (33–85%), diarrhea (15–29%), and thromboembolism (11–24%). The characteristic rash usually starts as an annular erythema at intertriginous and periorificial sites, especially in the groin or buttock. It subsequently becomes raised, and bullae form; when the bullae rupture, eroded areas form. The lesions can wax and wane. The development of a similar rash in patients receiving glucagon therapy suggests that the rash is a direct effect of the hyperglucagonemia. A characteristic laboratory finding is hypoaminoacidemia, which occurs in 26–100% of patients.

Glucagonomas are generally large tumors at diagnosis (5–10 cm). Some 50–80% occur in the pancreatic tail. From 50 to 82% have evidence of metastatic spread at presentation, usually to the liver. Glucagonomas are rarely extrapancreatic and usually occur singly.

Two new entities have been described that can also cause hyperglucagonemia and may mimic glucagonomas. Mahvah disease is due to a homozygous P86S mutation of the human glucagon receptor. It is associated with the development of α -cell hyperplasia, hyperglucagonemia, and the development of nonfunctioning pNETs. A second disease called *glucagon cell adenomatosis* can mimic glucagonoma syndrome clinically and is characterized by the presence of hyperplastic islets staining positive for glucagon instead of a single glucagonoma.

570 Diagnosis The diagnosis is confirmed by demonstrating an increased plasma glucagon level. Characteristically, plasma glucagon levels exceed 1000 pg/mL (normal is <150 pg/mL) in 90%; 7% are between 500 and 1000 pg/mL, and 3% are <500 pg/mL. A trend toward lower levels at diagnosis has been noted in the last decade. A plasma glucagon level >1000 pg/mL is considered diagnostic of glucagonoma. Other diseases causing increased plasma glucagon levels include cirrhosis, diabetic ketoacidosis, celiac disease, renal insufficiency, acute pancreatitis, hypercorticism, hepatic insufficiency, severe stress, and prolonged fasting or familial hyperglucagonemia, as well as danazol treatment. With the exception of cirrhosis, these disorders do not increase plasma glucagon >500 pg/mL.

Necrolytic migratory erythema is not pathognomonic for glucagonoma and occurs in myeloproliferative disorders, hepatitis B infection, malnutrition, short-bowel syndrome, inflammatory bowel disease, zinc deficiency, and malabsorption disorders.

TREATMENT GLUCAGONOMAS

In 50–80% of patients, hepatic metastases are present, and so curative surgical resection is not possible. Surgical debulking in patients with advanced disease or other antitumor treatments may be beneficial (see below). Long-acting somatostatin analogues such as octreotide and lanreotide improve the skin rash in 75% of patients and may improve the weight loss, pain, and diarrhea, but usually do not improve the glucose intolerance.

SOMATOSTATINOMA SYNDROME

The somatostatinoma syndrome is due to an NET that secretes excessive amounts of somatostatin, which causes a distinct syndrome characterized by diabetes mellitus, gallbladder disease, diarrhea, and steatorrhea. There is no general distinction in the literature between a tumor that contains somatostatin-like immunoreactivity (somatostatinoma) and does (11–45%) or does not (55–90%) produce a clinical syndrome (somatostatinoma syndrome) by secreting somatostatin. In a review of 173 cases of somatostatinomas, only 11% were associated with the somatostatinoma syndrome. The mean age is 51 years. Somatostatinomas occur primarily in the pancreas and small intestine, and the frequency of the symptoms and occurrence of the somatostatinoma syndrome differ in each. Each of the usual symptoms is more common in pancreatic than in intestinal somatostatinomas: diabetes mellitus (95% vs 21%), gallbladder disease (94% vs 43%), diarrhea (92% vs 38%), steatorrhea (83% vs 12%), hypochlorhydria (86% vs 12%), and weight loss (90% vs 69%). The somatostatinoma syndrome occurs in 30–90% of pancreatic and 0–5% of SI somatostatinomas. In various series, 43% of all duodenal NETs contain somatostatin; however, the somatostatinoma syndrome is rarely present (<2%). Somatostatinomas occur in the pancreas in 56–74% of cases, with the primary location being the pancreatic head. The tumors are usually solitary (90%) and large (mean size 4.5 cm). Liver metastases are common, being present in 69–84% of patients. Somatostatinomas are rare in patients with MEN 1, occurring in only 0.65%.

Somatostatin is a tetradecapeptide that is widely distributed in the CNS and GI tract, where it functions as a neurotransmitter or has paracrine and autocrine actions. It is a potent inhibitor of many processes, including release of almost all hormones, acid secretion, intestinal and pancreatic secretion, and intestinal absorption. Most of the clinical manifestations are directly related to these inhibitory actions.

Diagnosis In most cases, somatostatinomas have been found by accident either at the time of cholecystectomy or during endoscopy. The presence of psammoma bodies in a duodenal tumor should particularly raise suspicion. Duodenal somatostatin-containing tumors are increasingly associated with von Recklinghausen's disease (NF-1) (Table 113-6). Most of these tumors (>98%) do not cause the somatostatinoma syndrome. The diagnosis of the somatostatinoma syndrome requires the demonstration of elevated plasma somatostatin levels.

TREATMENT SOMATOSTATINOMAS

Pancreatic tumors are frequently (70–92%) metastatic at presentation, whereas 30–69% of SI somatostatinomas have metastases. Surgery is the treatment of choice for those without widespread hepatic metastases. Symptoms in patients with the somatostatinoma syndrome are also improved by octreotide treatment.

VIPOMAS

VIPomas are NETs that secrete excessive amounts of vasoactive intestinal peptide (VIP), which causes a distinct syndrome characterized by large-volume diarrhea, hypokalemia, and dehydration. This syndrome also is called Verner-Morrison syndrome, pancreatic cholera, and WDHA syndrome for watery diarrhea, hypokalemia, and achlorhydria, which some patients develop. The mean age of patients with this syndrome is 49 years; however, it can occur in children, and when it does, it is usually caused by a ganglioneuroma or ganglioneuroblastoma.

The principal symptoms are large-volume diarrhea (100%) severe enough to cause hypokalemia (80–100%), dehydration (83%), hypochlorhydria (54–76%), and flushing (20%). The diarrhea is secretory in nature, persisting during fasting, and is almost always >1 L/d and in 70% is >3 L/d. In a number of studies, the diarrhea was intermittent initially in up to half the patients. Most patients do not have accompanying steatorrhea (16%), and the increased stool volume is due to increased excretion of sodium and potassium, which, with the anions, accounts for the osmolality of the stool. Patients frequently have hyperglycemia (25–50%) and hypercalcemia (25–50%).

VIP is a 28-amino-acid peptide that is an important neurotransmitter, ubiquitously present in the CNS and GI tract. Its known actions include stimulation of SI chloride secretion as well as effects on smooth-muscle contractility, inhibition of acid secretion, and vasodilatory effects, which explain most features of the clinical syndrome.

In adults, 80–90% of VIPomas are pancreatic in location, with the rest due to VIP-secreting pheochromocytomas, intestinal carcinoids, and rarely ganglioneuromas. These tumors are usually solitary, 50–75% are in the pancreatic tail, and 37–68% have hepatic metastases at diagnosis. In children <10 years old, the syndrome is usually due to ganglioneuromas or ganglioblastomas and is less often malignant (10%).

Diagnosis The diagnosis requires the demonstration of an elevated plasma VIP level and the presence of large-volume diarrhea. A stool volume <700 mL/d is proposed to exclude the diagnosis of VIPoma. When the patient fasts, a number of diseases can be excluded that can cause marked diarrhea because the high volume of diarrhea is not sustained during the fast. Other diseases that can produce a secretory large-volume diarrhea include gastrinomas, chronic laxative abuse, carcinoid syndrome, systemic mastocytosis, rarely medullary thyroid cancer, diabetic diarrhea, sprue, and AIDS. Among these conditions, only VIPomas caused a marked increase in plasma VIP. Chronic surreptitious use of laxatives/diuretics can be particularly difficult to detect clinically. Hence, in a patient with unexplained chronic diarrhea, screens for laxatives should be performed; they will detect many, but not all, laxative abusers. Elevated plasma levels of VIP should not be the only basis of the diagnosis of VIPomas because they can occur with some diarrheal states including inflammatory bowel disease, post small bowel resection, and radiation enteritis. Furthermore, nesidioblastosis can mimic VIPomas by causing elevated plasma VIP levels, diarrhea, and even false-positive location in the pancreatic region on somatostatin receptor scintigraphy.

TREATMENT VIPOMAS

The most important initial treatment in these patients is to correct their dehydration, hypokalemia, and electrolyte losses with fluid and electrolyte replacement. These patients may require 5 L/d of fluid and >350 mEq/d of potassium. Because 37–68% of adults with VIPomas have metastatic disease in the liver at presentation, a

significant number of patients cannot be cured surgically. In these patients, long-acting somatostatin analogues such as octreotide and lanreotide are the drugs of choice.

Octreotide/lanreotide will control the diarrhea short- and long-term in 75–100% of patients. In nonresponsive patients, the combination of glucocorticoids and octreotide/lanreotide has proved helpful in a small number of patients. Other drugs reported to be helpful in small numbers of patients include prednisone (60–100 mg/d), clonidine, indomethacin, phenothiazines, loperamide, lidamide, lithium, propranolol, and metoclopramide. Treatment of advanced disease with cytoreductive surgery, embolization, chemoembolization, chemotherapy, radiotherapy, radiofrequency ablation, and peptide receptor radiotherapy may be helpful (see below).

NONFUNCTIONAL PANCREATIC NEUROENDOCRINE TUMORS (NF-pNETs)

NF-pNETs are NETs that originate in the pancreas and either secrete no products or their products do not cause a specific clinical syndrome. Their symptoms are due entirely to the tumor per se. NF-pNETs secrete chromogranin A (90–100%), chromogranin B (90–100%), α -HCG (human chorionic gonadotropin) (40%), neuron-specific enolase (31%), and β -HCG (20%), and because 40–90% secrete PP, they are also often called PPomas. Because the symptoms are due to the tumor mass, patients with NF-pNETs usually present late in the disease course with invasive tumors and hepatic metastases (64–92%), and the tumors are usually large (72% >5 cm). NF-pNETs are usually solitary except in patients with MEN 1, in which case they are multiple. They occur primarily in the pancreatic head. Even though these tumors do not cause a functional syndrome, immunocytochemical studies show that they synthesize numerous peptides and cannot be distinguished from functional pNETs by immunocytochemistry. In MEN 1, 80–100% of patients have microscopic NF-pNETs, but they become large or symptomatic in a minority (0–13%) of cases. In VHL, 12–17% develop NF-pNETs, and in 4%, they are ≥ 3 cm in diameter.

The most common symptoms are abdominal pain (30–80%), jaundice (20–35%), and weight loss, fatigue, or bleeding; 10–35% are found incidentally. The average time from the beginning of symptoms to diagnosis is 5 years.

Diagnosis The diagnosis is established by histologic confirmation in a patient without either the clinical symptoms or the elevated plasma hormone levels of one of the established syndromes. The principal difficulty in diagnosis is to distinguish an NF-pNET from a nonendocrine pancreatic tumor, which is more common, as well as from a functional pNET. Even though chromogranin A levels are elevated in almost every patient, this is not specific for this disease as it can be found in functional pNETs, GI-NETs (carcinoids), and other neuroendocrine disorders. Plasma PP elevations should strongly suggest the diagnosis in a patient with a pancreatic mass because it is usually normal in patients with pancreatic adenocarcinomas. Elevated plasma PP is not diagnostic of this tumor because it is elevated in a number of other conditions, such as chronic renal failure, old age, inflammatory conditions, alcohol abuse, pancreatitis, hypoglycemia, postprandially, and diabetes. A positive somatostatin receptor scan in a patient with a pancreatic mass should suggest the presence of pNET/NF-pNET rather than a nonendocrine tumor.

TREATMENT NONFUNCTIONAL PANCREATIC NEUROENDOCRINE TUMORS (NF-pNETs)

Overall survival in patients with sporadic NF-pNET is 30–63% at 5 years, with a median survival of 6 years. Unfortunately, surgical curative resection can be considered only in a minority of these patients because 64–92% present with diffuse metastatic disease. Treatment needs to be directed against the tumor per se using the various modalities discussed below for advanced disease. The treatment of NF-pNETs in either MEN 1 patients or patients with VHL is controversial. Most recommend surgical resection for any tumor

>2–3 cm in diameter; however, there is no consensus on smaller NF-pNETs in these inherited disorders, with most recommending careful surveillance of these patients. The treatment of small sporadic, asymptomatic NF-pNETs (≤ 2 cm) is also controversial. Most of these are low- or intermediate-grade lesions, and <7% are malignant. Some advocate a nonoperative approach with careful, regular follow-up, whereas other recommend an operative approach with special consideration for a laparoscopic surgical approach.

GRFOMAS

GRFomas are NETs that secrete excessive amounts of growth hormone-releasing factor (GRF) that cause acromegaly. GRF is a 44-amino-acid peptide, and 25–44% of pNETs have GRF immunoreactivity, although it is uncommonly secreted. GRFomas are lung tumors in 47–54% of cases, pNETs in 29–30%, and SI carcinoids in 8–10%; up to 12% occur at other sites. Patients have a mean age of 38 years, and the symptoms usually are due to either acromegaly or the tumor per se. The acromegaly caused by GRFomas is indistinguishable from classic acromegaly. The pancreatic tumors are usually large (>6 cm), and liver metastases are present in 39%. They should be suspected in any patient with acromegaly and an abdominal tumor, a patient with MEN 1 with acromegaly, or a patient without a pituitary adenoma with acromegaly or associated with hyperprolactinemia, which occurs in 70% of GRFomas. GRFomas are an uncommon cause of acromegaly. GRFomas occur in <1% of MEN 1 patients. The diagnosis is established by performing plasma assays for GRF and growth hormone. Most GRFomas have a plasma GRF level >300 pg/mL (normal <5 pg/mL men, <10 pg/mL women). Patients with GRFomas also have increased plasma levels of insulin-like growth factor type I (IGF-I) similar to those in classic acromegaly. Surgery is the treatment of choice if diffuse metastases are not present. Long-acting somatostatin analogues such as octreotide and lanreotide are the agents of choice, with 75–100% of patients responding.

OTHER RARE PANCREATIC NEUROENDOCRINE TUMOR SYNDROMES

Cushing's syndrome (ACTHoma) due to a pNET occurs in 4–16% of all ectopic Cushing's syndrome cases. It occurs in 5% of cases of sporadic gastrinomas, almost invariably in patients with hepatic metastases, and is an independent poor prognostic factor. Paraneoplastic hypercalcemia due to pNETs releasing parathyroid hormone-related peptide (PTHrP), a PTH-like material, or unknown factor, is rarely reported. The tumors are usually large, and liver metastases are usually present. Most (88%) appear to be due to release of PTHrP. pNETs occasionally can cause the carcinoid syndrome. A number of very rare pNET syndromes involving a few cases (less than five) have been described; these include a renin-producing pNET in a patient presenting with hypertension; pNETs secreting luteinizing hormone, resulting in masculinization or decreased libido; a pNET secreting erythropoietin, resulting in polycythemia; pNETs secreting IGF-II, causing hypoglycemia; and pNETs secreting enteroglucagon, causing small intestinal hypertrophy, colonic/SI stasis, and malabsorption (Table 113-2). A number of other possible functional pNETs have been proposed, but most authorities classify these as unclear or as a nonfunctional pNET because in each case numerous patients have been described with similar plasma hormone elevations that do not cause any symptoms. These include pNETs secreting calcitonin, neurotensin (neurotensinoma), PP (PPoma), and ghrelin (Table 113-2).

TUMOR LOCALIZATION

Localization of the primary tumor and knowledge of the extent of the disease are essential to the proper management of all GI-NETs (carcinoids) and pNETs. Without proper localization studies, it is not possible to determine whether the patient is a candidate for surgical resection (curative or cytoreductive) or requires antitumor treatment, to determine whether the patient is responding to antitumor therapies, or to appropriately classify/stage the patient's disease to assess prognosis.

Numerous tumor localization methods are used in both types of NETs, including cross-sectional imaging studies (CT, magnetic resonance imaging [MRI], transabdominal ultrasound), selective angiography, somatostatin receptor scintigraphy (SRS), and positron emission

572 tomography. In pNETs, endoscopic ultrasound (EUS) and functional localization by measuring venous hormonal gradients are also reported to be useful. Bronchial carcinoids are usually detected by standard chest radiography and assessed by CT. Rectal, duodenal, colonic, and gastric carcinoids are usually detected by GI endoscopy. Because of their wide availability, CT and MRI are generally initially used to determine the location of the primary NETs and the extent of disease. NETs are hypervascular tumors, and with both MRI and CT, contrast enhancement is essential for maximal sensitivity, and it is recommended that generally triple-phase scanning be used. The ability of cross-sectional imaging and, to a lesser extent, SRS to detect NETs is a function of NET size. With CT and MRI, <10% of tumors <1 cm in diameter are detected, 30–40% of tumors 1–3 cm are detected, and >50% of tumors >3 cm are detected. Many primary GI-NETs (carcinoids) are small, as are insulinomas and duodenal gastrinomas, and are frequently not detected by cross-sectional imaging, whereas most other pNETs present late in the course of their disease and are large (>4 cm). Selective angiography is more sensitive, localizing 60–90% of all NETs; however, it is now used infrequently. For detecting liver metastases, CT and MRI are more sensitive than ultrasound, and with recent improvements, 5–25% of patients with liver metastases will be missed by CT and/or MRI.

pNETs, as well as GI-NETs (carcinoids), frequently (>80%) overexpress high-affinity somatostatin receptors in both the primary tumors and the metastases. Of the five types of somatostatin receptors (sst_{1-5}), radiolabeled octreotide binds with high affinity to sst_2 and sst_3 , has a lower affinity for sst_3 , and has a very low affinity for sst_1 and sst_4 . Between 80 and 100% of GI-NETs (carcinoids) and pNETs possess sst_2 , and many also have the other four sst subtypes. Interaction with these receptors can be used to treat these tumors as well as to localize NETs by using radiolabeled somatostatin analogues (SRS). In the United States, [^{111}In -DTPA-D-Phe 1]octreotide (octreoscan) is generally used with gamma camera detection using single-photon emission computed tomography (SPECT) imaging. Numerous studies, primarily in Europe, using gallium-68-labeled somatostatin analogues and positron emission tomography (PET) detection, demonstrate even greater sensitivity than with SRS with ^{111}In -labeled somatostatin analogues. Although not yet approved in the United States, there are a number of centers starting to use this approach. Because of its sensitivity and ability to localize tumor throughout the body, SRS is the initial imaging modality of choice for localizing both the primary tumor and metastatic NETs. SRS localizes tumor in 73–95% of patients with GI-NETs (carcinoids) and in 56–100% of patients with pNETs, except insulinomas. Insulinomas are usually small and have low densities of sst receptors, resulting in SRS being positive in only 12–50% of patients with insulinomas. SRS identifies >90–95% of patients with liver metastases due to NETs. **Figure 113-3** shows an example of the increased sensitivity of SRS in a patient with a GI-NET (carcinoid) tumor. The CT scan showed a single liver metastasis, whereas the SRS demonstrated three metastases in the liver in multiple locations. Occasional false-positive responses with SRS can occur (12% in one study) because numerous other normal tissues as well as diseases can have high densities of sst receptors, including granulomas (sarcoid, tuberculosis, etc.), thyroid diseases (goiter, thyroiditis), and activated lymphocytes (lymphomas, wound infections). If liver metastases are identified by SRS, to plan the proper treatment, either a CT or an MRI (with contrast enhancement) is recommended to assess the size and exact location of the metastases because SRS does not provide information on tumor size. For pNETs in the pancreas, EUS is highly sensitive, localizing 77–100% of insulinomas, which occur almost exclusively within the pancreas. Endoscopic ultrasound is less sensitive for extrapancreatic tumors. It is increasingly used in patients with MEN 1, and to a lesser extent VHL, to detect small pNETs not seen with other modalities or for serial pNET assessments to determine size changes or rapid growth in patients in whom surgery is deferred. EUS with cytologic evaluation also is used frequently to distinguish an NF-pNET from a pancreatic adenocarcinoma or another nonendocrine pancreatic tumor. Not infrequently patients present with liver metastases due to an NET and the primary site is unclear. Occult small intestinal NETs (carcinoids) are increasingly detected by double-balloon enteroscopy or capsule endoscopy.

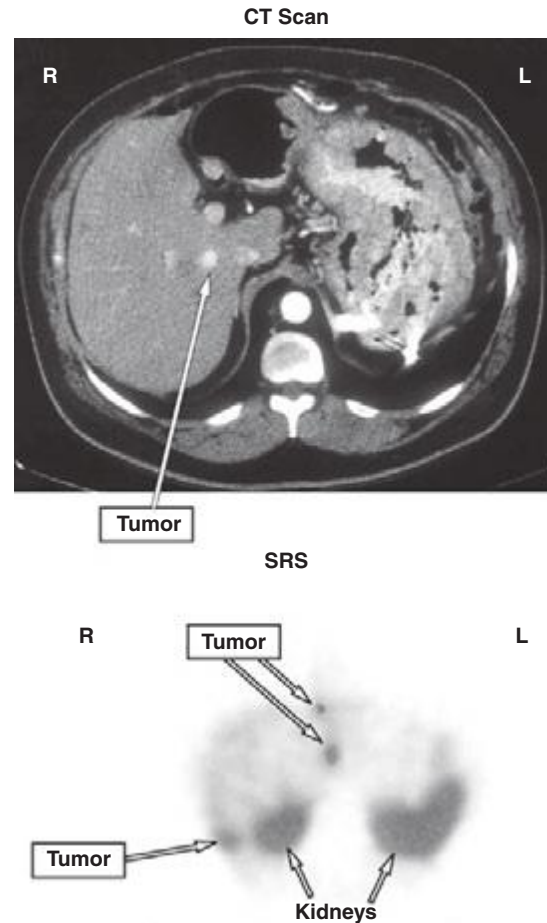


FIGURE 113-3 Ability of computed tomography (CT) scanning (*top*) or somatostatin receptor scintigraphy (SRS) (*bottom*) to localize metastatic carcinoid in the liver.

Insulinomas frequently overexpress receptors for glucagon-like peptide-1 (GLP-1), and radiolabeled GLP-1 analogues have been developed that can detect occult insulinomas not localized by other imaging modalities. Functional localization by measuring hormonal gradients is now uncommonly used with gastrinomas (after intra-arterial secretin injections) but is still frequently used in insulinoma patients in whom other imaging studies are negative (assessing hepatic vein insulin concentrations post-intra-arterial calcium injections). Functional localization measuring hormone gradients in insulinomas or gastrin gradients in gastrinoma is a sensitive method, being positive in 80–100% of patients. The intra-arterial calcium test may also allow differentiation of the cause of the hypoglycemia and indicate whether it is due to an insulinoma or a nesidioblastosis. The latter entity is becoming increasingly important because hypoglycemia after gastric bypass surgery for obesity is increasing in frequency, and it is primarily due to nesidioblastosis, although it can occasionally be due to an insulinoma.

PET and use of hybrid scanners such as CT and SRS may have increased sensitivity. PET scanning with ^{18}F -fluoro-DOPA in patients with carcinoids or with ^{11}C -5-HTP in patients with pNETs or GI-NETs (carcinoids) has greater sensitivity than cross-sectional imaging studies and may be used increasingly in the future. PET scanning for GI-NETs is not currently approved in the United States.

TREATMENT ADVANCED DISEASE (DIFFUSE METASTATIC DISEASE)

The single most important prognostic factor for survival is the presence of liver metastases (**Fig. 113-4**). For patients with foregut carcinoids without hepatic metastases, the 5-year survival in one

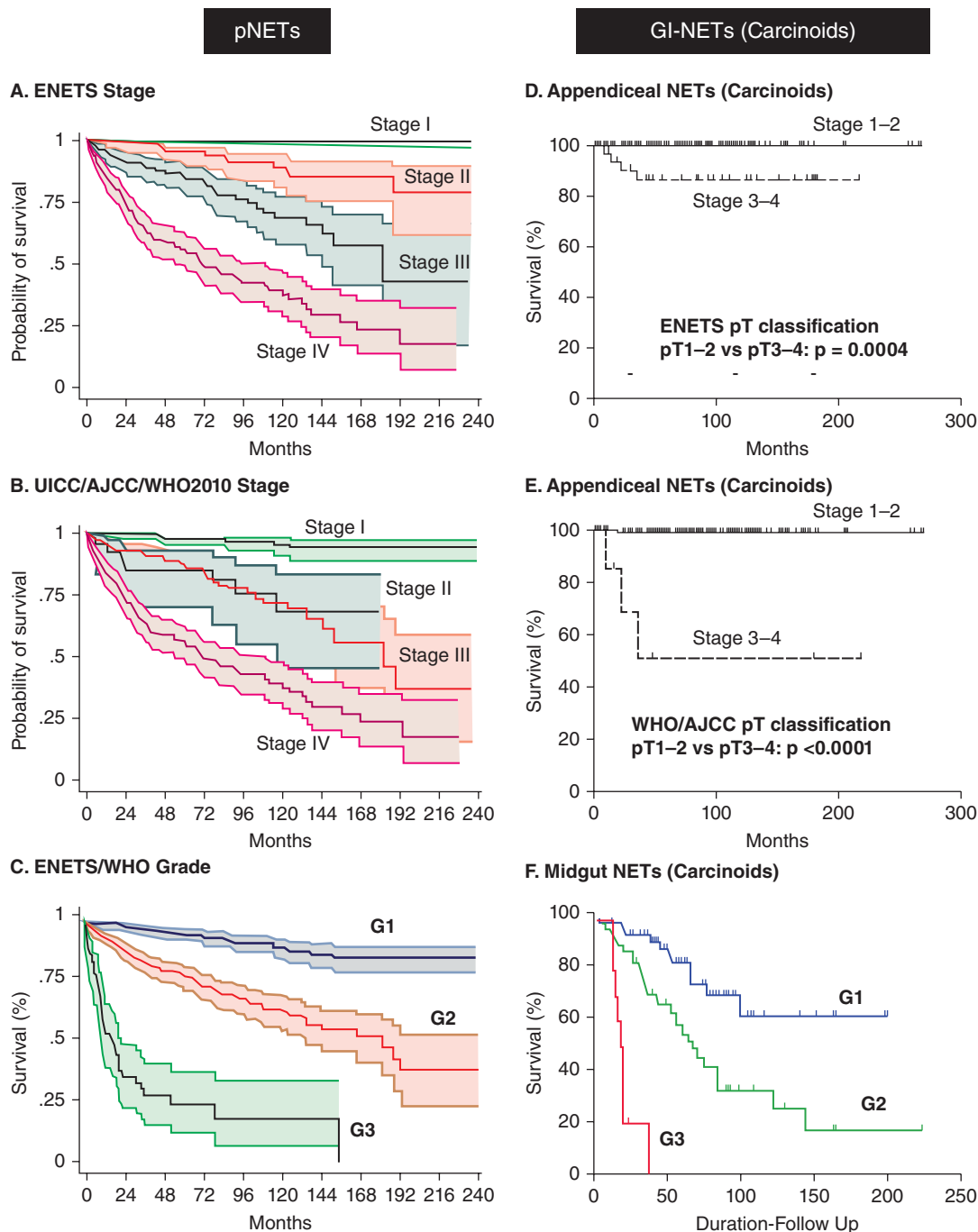


FIGURE 113-4 Survival (Kaplan-Meier plots) of patients with pancreatic neuroendocrine tumors (pNETs; $n = 1072$) (A–C) or gastrointestinal neuroendocrine tumors (GI-NETs; carcinoids) (appendix, $n = 138$; midgut, $n = 238$) (D–F) stratified according to recent proposed classification and grading systems. (Panels A–C are drawn from data in G Rindi et al: *J Natl Cancer Inst* 104:764, 2012; panels D and E are drawn from data in M Volante et al: *Am J Surg Pathol* 37:606, 2013; and panel F is drawn from data in MS Khan: *Br J Cancer* 108:1838, 2013.)

study was 95%, and with distant metastases, it was 20% (Fig. 113-4). With gastrinomas, the 5-year survival without liver metastases is 98%; with limited metastases in one hepatic lobe, it is 78%; and with diffuse metastases, 16% (Fig. 113-4). In a large study of 156 patients (67 pNETs, rest carcinoids), the overall 5-year survival rate was 77%; it was 96% without liver metastases, 73% with liver metastases, and 50% with distant disease. Another very important prognostic factor is whether the NET is well-differentiated (G1/G2) or poorly differentiated (<1% of all NETs) (G3). Well-differentiated NETs have a 5-year survival of 50–80%, whereas poorly differentiated NETs have a 5-year survival of only 0–15%.

Therefore, treatment for advanced metastatic disease is an important challenge. A number of different modalities are reported to be effective, including cytoreductive surgery (surgically or by

radiofrequency ablation [RFA]), treatment with chemotherapy, somatostatin analogues, interferon α , hepatic embolization alone or with chemotherapy (chemoembolization), molecular targeted therapy, radiotherapy with radiolabeled beads/microspheres, peptide radioreceptor therapy (PRRT), and liver transplantation.

SPECIFIC ANTITUMOR TREATMENTS

Cytoreductive surgery is considered if either all of the visible metastatic disease or at least 90% is thought resectable; however, unfortunately, this is possible in only the 9–22% of patients who present with limited hepatic metastases. Although no randomized studies have proven that it extends life, results from a number of studies suggest that it may increase survival; therefore, it is recommended, if possible. RFA can be applied to NET liver metastases if they are

574 limited in number (usually less than five) and size (usually <3.5 cm in diameter). It can be used at the time of surgery (either general or laparoscopic) or using radiologic guidance.

Response rates are >80%, the responses can last up to 3 years, the morbidity rate is low, and this procedure may be particularly helpful in patients with functional pNETs that are difficult to control medically. Although RFA has not been established in a controlled trial, both the European and North American Neuroendocrine Tumor Society guidelines (ENETS, NANETS) state it can be an effective antitumor treatment for both refractory functional syndromes and for palliative treatment.

Chemotherapy plays a different role in the treatment of patients with pNETs and GI-NETs (carcinoids). Chemotherapy continues to be widely used in the treatment of patients with advanced pNETs with moderate success (response rates 20–70%); however, in general, its results in patients with metastatic GI-NETs (carcinoids) has been disappointing, with response rates of 0–30% with various two- and three-drug combinations, and thus, it is infrequently used in these patients. An important distinction in patients with pNETs is whether the tumor is well differentiated (G1/G2) or poorly differentiated (G3). The chemotherapeutic approach is different for these two groups. The current regimen of choice for patients with well-differentiated pNETs is the combination of streptozotocin and doxorubicin with or without 5-fluorouracil. Streptozotocin is a glucosamine nitrourea compound originally found to have cytotoxic effects on pancreatic islets, and later in studies with doxorubicin with or without 5-fluorouracil, it produced response rates of 20–45% in advanced pNETs. Streptozotocin causes considerable morbidity, with 70–100% of patients developing side effects (most prominent being nausea/vomiting in 60–100% or leukopenia/thrombocytopenia) and 15–40% of patients developing some degree of renal dysfunction (proteinuria in 40–50%, decreased creatine clearance). The combination of temozolomide (TMZ) with capecitabine produces partial response rates as high as 70% in patients with advanced pNETs and a 2-year survival of 92%. The use of TMZ or another alkylating agent in advanced pNETs is supported by studies that show low levels of the DNA repair enzyme O⁶-methylguanine DNA methyltransferase in pNETs, but not in GI-NETs (carcinoids), which increases the sensitivity of pNETs to TMZ. In poorly differentiated NETs (G3), chemotherapy with a cisplatin-based regimen with etoposide or other agents (vincristine, paclitaxel) is the recommended treatment, with response rates of 40–70%; however, responses are generally short-lived (<12 months). This chemotherapy regimen can be associated with significant toxicity including GI toxicities (nausea, vomiting), myelosuppression, and renal toxicity.

In addition to the effectiveness in controlling the functional hormonal state, long-acting somatostatin analogues such as octreotide and lanreotide are increasingly used for their antiproliferative effects. Whereas somatostatin analogues rarely decrease tumor size (i.e., 0–17%), these drugs have tumoristatic effects, stopping additional growth in 26–95% of patients with NETs. In a randomized, double-blind study in patients with metastatic midgut carcinoids (PROMID study) octreotide-LAR demonstrated a marked lengthening of time to progression (14.3 vs 6 months, $p = .000072$). This improvement was seen in patients with limited liver involvement. This study did not assess whether such treatment will extend survival. A double-blind, randomized, placebo-controlled, phase III study in patients with well-differentiated, metastatic, inoperable pNETs (45%) or GI-NETs (carcinoids) (55%) (CLARINET study) showed that monthly treatment with lanreotide-autogel reduced tumor progression or death by 53%. Somatostatin analogues can induce apoptosis in GI-NETs (carcinoids), which probably contributes to their tumoristatic effects. Treatment with somatostatin analogues is generally well-tolerated, with most side effects being mild and uncommonly leading to stopping the drug. Potential long-term side effects include diabetes/glucose intolerance, steatorrhea, and the development of gallbladder sludge/gallstones (10–80%), although only 1% of patients develop symptomatic gallbladder

disease. Because of these phase III studies, somatostatin analogues are generally recommended as first-line treatment for patients with well-differentiated metastatic NETs.

Interferon α , similar to somatostatin analogues, is effective at controlling the hormonal excess symptoms of NETs and has antiproliferative effects in NETs, which primarily result in disease stabilization (30–80%), with a decrease in tumor size in <15% of patients. Interferon can inhibit DNA synthesis, block cell cycle progression in the G₁ phase, inhibit protein synthesis, inhibit angiogenesis, and induce apoptosis. Interferon α treatment results in side effects in the majority of patients, with the most frequent being a flu-like syndrome (80–100%), anorexia with weight loss, and fatigue. These side effects frequently decrease in severity with continued treatment. In addition, patients become accommodated to the symptoms. More serious side effects include hepatotoxicity (31%), hyperlipidemia (31%), bone marrow toxicity, thyroid disease (19%), and rarely CNS side effects (depression, mental/visual disorders). ENETS 2012 guidelines conclude that in patients with well-differentiated NETs that are slowly progressive, interferon α treatment should be considered if the tumor is somatostatin receptor negative or if somatostatin treatment fails.

Selective internal radiation therapy (SIRT) using yttrium-90 (⁹⁰Y) glass or resin microspheres is a relatively newer approach being evaluated in patients with unresectable NET liver metastases, with approximately 500 NET patients treated. The treatment requires careful evaluation for vascular shunting before treatment and a pretreatment angiogram to evaluate placement of the catheter and is generally reserved for patients without extrahepatic metastatic disease and with adequate hepatic reserve. One of two types of ⁹⁰Y microspheres are used: either microspheres with a 20- to 60- μ m diameter and 50 Bq/sphere (SIR-Spheres) or glass microspheres (TheraSpheres) with a 20- to 30- μ m diameter and 2500 Bq/sphere. The ⁹⁰Y-microspheres are delivered to the liver by intra-arterial injection from percutaneously placed catheters. In four studies involving metastatic NETs, the response rate varied from 50–61% (partial or complete), tumor stabilization occurred in 22–41%, 60–100% had symptomatic improvement, and overall survival varied from 25–70 months. Side effects include postembolization syndrome (pain, fever, nausea/vomiting [frequent]), which is usually mild, although grade 2 (43%) or grade 3 (1%) symptoms can occur; radiation-induced liver disease (<1%); and radiation pneumonitis (<1%). Contraindications to use include excess shunting to the GI tract or lung, inability to isolate the liver arterial supply, and inadequate liver reserve. Because of the limited data available in the ENETS 2012 guidelines, treatment with SIRT is considered experimental.

Molecular targeted medical treatment with either an mTOR inhibitor (everolimus) or a tyrosine kinase inhibitor (sunitinib) is now approved treatment in the United States and Europe for patients with metastatic unresectable pNET, each supported by a phase III, double-blind, prospective, placebo-controlled trial. mTOR is a serine-threonine kinase that plays an important role in proliferation, cell growth, and apoptosis in both normal and neoplastic cells. Activation of the mTOR cascade is important in mediating NET cell growth, especially in pNETs. A number of mTOR inhibitors have shown promising antitumor activity in NETs including everolimus and temsirolimus, with the former undergoing a phase III trial (RADIANT-3) involving 410 patients with advanced progressive pNETs. Everolimus caused significant improvement in progression-free survival (11 vs 4.6 months, $p < .001$) and increased by a factor of 3.7 the proportion of patients progression-free at 18 months (37% vs 9%). Everolimus treatment was associated with frequent side effects, causing a twofold increase in adverse events, with the most frequent being grade 1 or 2. Grade 3 or 4 side effects included hematologic, GI (diarrhea), stomatitis, or hypoglycemia occurring in 3–7% of patients. Most grade 3 or 4 side effects were controlled by dose reduction or drug interruption. The ENETS 2012 guidelines conclude that everolimus, similar to sunitinib (below), should be considered as a first-line treatment in selected cases of

114 Bladder and Renal Cell Carcinomas

Howard I. Scher, Jonathan E. Rosenberg,
Robert J. Motzer

well-differentiated pNETs that are unresectable. NETs, like other normal and neoplastic cells, frequently possess multiple types of the 20 different tyrosine kinase (TK) receptors that are known and mediate the action of different growth factors. Numerous studies demonstrate that TK receptors in normal and neoplastic tissues as well as NETs are especially important in mediating cell growth, angiogenesis, differentiation, and apoptosis. Whereas a number of TK inhibitors show antiproliferative activity in NETs only sunitinib has undergone a phase III controlled trial. Sunitinib is an orally active small-molecule inhibitor of TK receptors (PDGFRs, VEGFR-1, VEGFR-2, c-KIT, FLT-3). In a phase III study in which 171 patients with progressive, metastatic, nonresectable pNETs were treated with sunitinib (37.5 mg/d) or placebo, sunitinib treatment caused a doubling of progression-free survival (11.4 vs 4.5 months, $p < .001$), an increase in objective tumor response rate (9% vs 0%, $p = .007$), and an increase in overall survival. Sunitinib treatment was associated with an overall threefold increase in side effects, although most were grade 1 or 2. The most frequent grade 3 or 4 side effects were neutropenia (12%) and hypertension (9.6%), which were controlled by dose reduction or temporary interruption. There is no consensus regarding the order of sunitinib or everolimus use in patients with advanced, well-differentiated, progressive pNETs.

PRRT for NETs involves treatment with radiolabeled somatostatin analogues. The success of this approach is based on the finding that somatostatin receptors (sst) are overexpressed or ectopically expressed by 60–100% of all NETs, which allows the targeting of cytotoxic, radiolabeled somatostatin receptor ligands.

Three different radionuclides are being used. High doses of [^{111}In -DTPA-D-Phe 1]octreotide, which emits γ -rays, internal conversion, and Auger electrons; ^{90}Y trium, which emits high-energy β -particles coupled by a DOTA chelating group to octreotide or octreotate; and ^{177}Lu tetium-coupled analogues, which emit both, are all in clinical studies. At present, the ^{177}Lu tetium-coupled analogues are the most widely used. ^{111}In dium-, ^{90}Y trium-, and ^{177}Lu tetium-labeled compounds caused tumor stabilization in 41–81%, 44–88%, and 23–40%, respectively, and a decrease in tumor size in 8–30%, 6–37%, and 38%, respectively, of patients with advanced metastatic NETs. In one large study involving 504 patients with malignant NETs, ^{177}Lu tetium-labeled analogues produced a reduction of tumor size of >50% in 30% of patients (2% complete) and tumor stabilization in 51% of patients. An effect on survival has not been established. At present, PRRT is not approved for use in either the United States or Europe, but because of the above promising results, a large phase III study is now being conducted in both the United States and Europe. The ENETS 2012, NANETS 2010, Nordic 2010, and European Society for Medical Oncology (ESMO) guidelines list PRRT as an experimental or investigational treatment at present.

The use of liver transplantation has been abandoned for treatment of most metastatic tumors to the liver. However, for metastatic NETs, it is still a consideration. Among 213 European patients with NETs (50% functional NETs) who had liver transplantation from 1982 to 2009, the overall 5-year survival was 52% and disease free-survival was 30%. In various studies, the postoperative mortality rate is 10–14%. These results are similar to the United Network for Organ Sharing data in the United States in which 150 NET patients had liver transplants and the 5-year survival was 49%. In various studies, important prognostic factors for a poor outcome include a major resection performed in addition at the time of the liver transplant; poor tumor differentiation; hepatomegaly; age >45 years; a primary NET in the duodenum or pancreas; the presence of extrahepatic metastatic disease or extensive liver involvement (>50%); Ki-67 proliferative index >10%; and abnormal E-cadherin staining. The ENETS 2012 guidelines conclude that liver transplantation should be viewed as providing palliative care, with cure an exception, and recommend it be reserved for patients with life-threatening hormonal disturbances refractory to other treatments or for selected patients with a nonfunctional tumor with diffuse liver metastatic disease refractory to all other treatments.

BLADDER CANCER

Transitional cell epithelium lines the urinary tract from the renal pelvis to the ureter, urinary bladder, and the proximal two-thirds of the urethra. Cancers can occur at any point: 90% of malignancies develop in the bladder, 8% in the renal pelvis, and 2% in the ureter or urethra. Bladder cancer is the fourth most common cancer in men and the thirteenth in women, with an estimated 72,570 new cases and 15,210 deaths in the United States predicted for the year 2013. The almost 5:1 ratio of incidence to mortality reflects the higher frequency of the less lethal superficial variants compared to the more lethal invasive and metastatic variants. The incidence is roughly four times higher in men than in women and twofold higher in white men than in black men, with a median age of 65 years.

Once diagnosed, urothelial tumors exhibit polychronotropism, which is the tendency to recur over time in new locations in the urothelial tract. As long as urothelium is present, continuous monitoring is required.

EPIDEMIOLOGY

Cigarette smoking is believed to contribute to up to 50% of urothelial cancers in men and nearly 40% in women. The risk of developing a urothelial cancer in male smokers is increased two- to fourfold relative to nonsmokers and continues for 10 years or longer after cessation. Other implicated agents include aniline dyes, the drugs phenacetin and chlor-naphazine, and external beam radiation. Chronic cyclophosphamide exposure also increases risk, whereas vitamin A supplements appear to be protective. Exposure to *Schistosoma haematobium*, a parasite found in many developing countries, is associated with an increase in both squamous and transitional cell carcinomas of the bladder.

PATHOLOGY

Clinical subtypes are grouped into three categories: 75% are superficial, 20% invade muscle, and 5% are metastatic at presentation. Staging of the tumor within the bladder is based on the pattern of growth and depth of invasion. The revised tumor, node, metastasis (TNM) staging system is illustrated in Fig. 114-1. About half of invasive tumors presented originally as superficial lesions that later progressed. Tumors are also rated by grade. Low-grade (highly differentiated) tumors rarely progress to a higher stage, whereas high-grade tumors do.

More than 95% of urothelial tumors in the United States are transitional cell in origin. Pure squamous cancers with keratinization constitute 3%, adenocarcinomas 2%, and small cell tumors (often with paraneoplastic syndromes) <1%. Adenocarcinomas develop primarily in the urachal remnant in the dome of the bladder or in the periurethral tissues. Paragangliomas, lymphomas, and melanomas are rare. Of the transitional cell tumors, low-grade papillary lesions that grow on a central stalk are most common. These tumors are very friable, have a tendency to bleed, and have a high risk for recurrence, yet they rarely progress to the more lethal invasive variety. In contrast, carcinoma in situ (CIS) is a high-grade tumor that is considered a precursor of the more lethal muscle-invasive disease.

PATHOGENESIS

The multicentric nature of the disease and high recurrence suggests a field effect in the urothelium that results in a predisposition to develop cancer. Molecular genetic analyses suggest that the superficial and invasive lesions develop along distinct molecular pathways. Low-grade noninvasive papillary tumors harbor constitutive activation of the receptor tyrosine kinase-Ras signal transduction pathway and high frequencies of fibroblast growth factor receptor 3 and phosphoinositide-3

576 kinase α subunit mutations. In contrast, CIS and invasive tumors have a higher frequency of *TP53* and *RB* gene alterations. Within all clinical stages, including Tis, T1, and T2 or greater lesions, tumors with alterations in *p53*, *p21*, and/or *RB* have a higher probability of recurrence, metastasis, and death from disease.

CLINICAL PRESENTATION, DIAGNOSIS, AND STAGING

Hematuria occurs in 80–90% of patients and often reflects exophytic tumors. The bladder is the most common source of gross hematuria (40%), but benign cystitis (22%) is a more common cause than bladder cancer (15%) (Chap. 61). Microscopic hematuria is more commonly of prostate origin (25%); only 2% of bladder cancers produce microscopic hematuria. Once hematuria is documented, a urinary cytology, visualization of the urothelial tract by computed tomography (CT) or magnetic resonance urogram or intravenous pyelogram, and cystoscopy are recommended if no other etiology is found. Screening asymptomatic individuals for hematuria increases the diagnosis of tumors at an early stage but has not been shown to prolong life. After hematuria, irritative symptoms are the next most common presentation. Ureteral obstruction may cause flank pain. Symptoms of metastatic disease are rarely the first presenting sign.

The endoscopic evaluation includes an examination under anesthesia to determine whether a palpable mass is present. A flexible endoscope is inserted into the bladder, and bladder barbotage for cytology is performed. Visual inspection includes mapping the location, size, and number of lesions, as well as a description of the growth pattern (solid vs papillary). All visible tumors should be resected, and a sample of the muscle underlying the tumor should be obtained to assess the depth of invasion. Normal-appearing areas are biopsied at random to ensure no CIS is present. A notation is made as to whether a tumor

was completely or incompletely resected. Selective catheterization and visualization of the upper tracts should be performed if the cytology is positive and no disease is visible in the bladder. Ultrasonography, CT, and/or magnetic resonance imaging (MRI) are used to determine whether a tumor extends to perivesical fat (T3) and to document nodal spread. Distant metastases are assessed by CT of the chest and abdomen, MRI, or radionuclide imaging of the skeleton.

TREATMENT BLADDER CANCER

Management depends on whether the tumor invades muscle and whether it has spread to the regional lymph nodes and beyond. The probability of spread increases with increasing T stage.

NON-MUSCLE-INVASIVE DISEASE

At a minimum, the management is complete endoscopic resection with or without intravesical therapy. The decision to recommend intravesical therapy depends on the histologic subtype, number of lesions, depth of invasion, presence or absence of CIS, and antecedent history. Recurrences develop in upward of 50% of cases, of which 5–20% progress to a more advanced stage. In general, solitary papillary lesions are managed by transurethral surgery alone. CIS and recurrent disease are treated by transurethral surgery followed by intravesical therapy.

Intravesical therapies are used in two general contexts: as an adjuvant to a complete endoscopic resection to prevent recurrence or to eliminate disease that cannot be controlled by endoscopic resection alone. Intravesical treatments are advised for patients with diffuse CIS, recurrent disease, >40% involvement of the bladder surface by tumor, or T1 disease. The standard therapy, based on randomized comparisons, is *Bacillus Calmette-Guérin* (BCG) in six weekly instillations, often followed by maintenance administrations for ≥ 1 year. Other agents with activity include mitomycin C, interferon, and gemcitabine. The side effects of intravesical therapies include dysuria, urinary frequency, and, depending on the drug, myelosuppression or contact dermatitis. Rarely, intravesical BCG may produce a systemic illness associated with granulomatous infections in multiple sites requiring antituberculin therapy.

Following the endoscopic resection, patients are monitored for recurrence at 3-month intervals during the first year. Recurrence may develop anywhere along the urothelial tract, including the renal pelvis, ureter, or urethra. Persistent disease in the bladder and new tumors are treated with a second course of BCG or intravesical chemotherapy with valrubicin or gemcitabine. In some cases, cystectomy is recommended. Tumors in the ureter or renal pelvis are typically managed by resection during retrograde examination or, in some cases, by instillation through the renal pelvis. Prostatic urethral tumors may require cystoprostatectomy if the tumor cannot be resected completely.

MUSCLE-INVASIVE DISEASE

The treatment of a tumor that has invaded muscle can be separated into control of the primary tumor and systemic chemotherapy to treat micrometastatic disease. Radical cystectomy is the standard treatment in the United States, although in selected cases, a bladder-sparing approach is used. This approach includes complete endoscopic resection; partial cystectomy; or a combination of resection, systemic chemotherapy, and external beam radiation therapy. In some countries, external beam radiation therapy is considered standard. In the United States, it is generally limited to those patients deemed unfit for cystectomy, those with unresectable local disease, or as part of an experimental bladder-sparing approach.

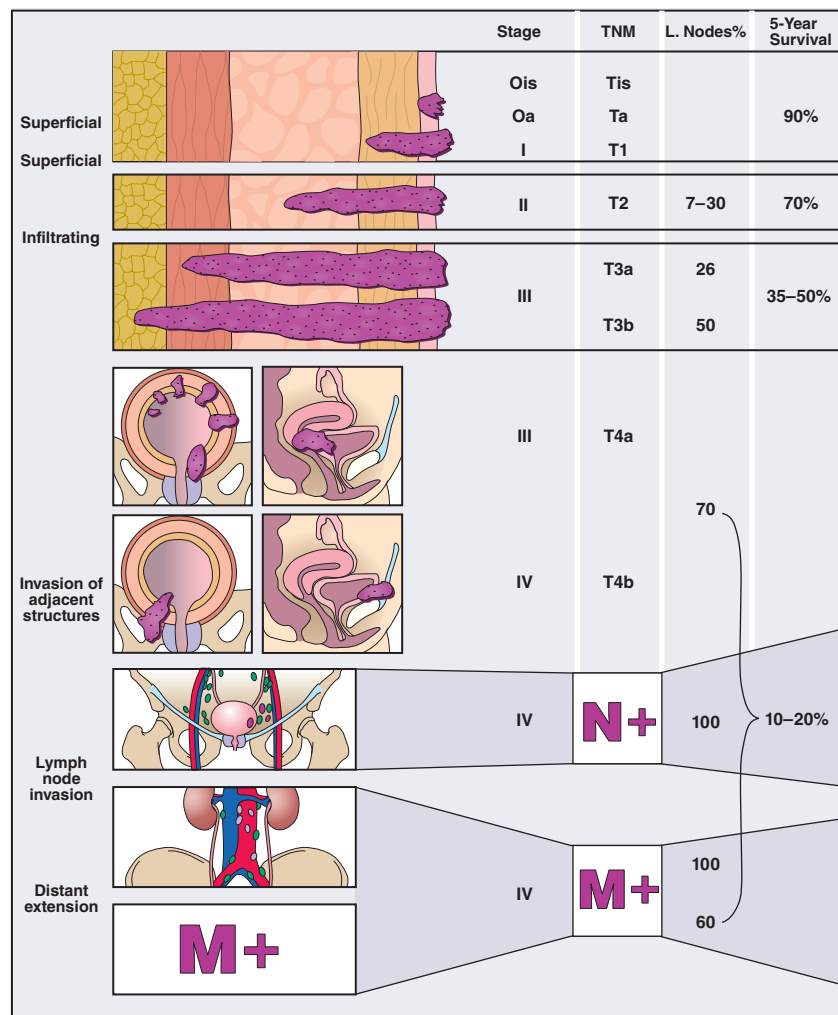


FIGURE 114-1 Bladder staging. TNM, tumor, node, metastasis.

Indications for cystectomy include muscle-invasive tumors not suitable for segmental resection; non-muscle-invasive tumors unsuitable for conservative management (e.g., due to multicentric and frequent recurrences resistant to intravesical instillations); high-grade T1 tumors especially if associated with CIS; and bladder symptoms (e.g., frequency or hemorrhage) that impair quality of life.

Radical cystectomy is major surgery that requires appropriate preoperative evaluation and management. It involves removal of the bladder and pelvic lymph nodes and creation of a conduit or reservoir for urinary flow. Grossly abnormal lymph nodes are evaluated by frozen section. If metastases are confirmed, the procedure is often aborted. In males, radical cystectomy includes the removal of the prostate, seminal vesicles, and proximal urethra. Impotence is universal unless the nerves responsible for erectile function are preserved. In females, the procedure includes removal of the bladder, urethra, uterus, fallopian tubes, ovaries, anterior vaginal wall, and surrounding fascia.

Several options are frequently used for urinary diversion. Ileal conduits bring urine directly from the ureter to the abdominal wall. Some patients receive either a continent cutaneous reservoir constructed from detubularized bowel or an orthotopic neobladder. Approximately 25% of men receive a neobladder, leading to 85–90% continence during the day. Cutaneous reservoirs are drained by intermittent catheterization. Contraindications to a neobladder include renal insufficiency, an inability to self-catheterize, or CIS or an exophytic tumor in the urethra. Diffuse CIS in the bladder is a relative contraindication based on the risk of a urethral recurrence. Concurrent ulcerative colitis or Crohn's disease may hinder the use of bowel.

A partial cystectomy may be considered when the disease is limited to the dome of the bladder, a ≥ 2 cm margin can be achieved, there is no associated CIS, and the bladder capacity is adequate after resection. This occurs in 5–10% of cases. Carcinomas in the ureter or in the renal pelvis are treated with nephroureterectomy with a bladder cuff to remove the tumor.

The probability of recurrence following surgery is based on pathologic stage, presence or absence of lymphatic or vascular invasion, and nodal spread. Among those whose cancers recur, the recurrence develops in a median of 1 year. Long-term outcomes vary by pathologic stage and histology (Table 114-1). The number of lymph nodes removed is also prognostic, whether or not the nodes contained tumor.

Chemotherapy (described below) has been shown to prolong the survival of patients with muscle-invasive disease when combined with definitive treatment of the bladder by radical cystectomy or radiation therapy. Presurgical (or neoadjuvant) chemotherapy has been the most thoroughly explored, and increases the cure rate by 5–15%, whereas postsurgical (adjuvant) chemotherapy has not been proven definitively beneficial. For the majority of patients, chemotherapy alone is inadequate to eradicate the disease. Use of neoadjuvant chemotherapy is increasing, although it still remains underused. Experimental studies are evaluating bladder preservation strategies by combining chemotherapy and radiation therapy in patients whose tumors were endoscopically removed.

METASTATIC DISEASE

The primary goal of metastatic disease treatment is to achieve complete remission with chemotherapy alone or with a combined-modality approach of chemotherapy followed by surgical resection of residual disease. One can define a goal in terms of cure or palliation

on the basis of the probability of achieving a complete response to chemotherapy using prognostic factors, such as Karnofsky performance status (KPS) (<80%) and whether the pattern of spread is nodal or visceral (liver, lung, or bone). For those with zero, one, or two risk factors, the probability of complete remission is 38, 25, and 5%, respectively, and median survival is 33, 13.4, and 9.3 months, respectively. Patients who have low KPS or who have visceral disease or bone metastases rarely achieve long-term survival. The toxicities also vary as a function of risk, and treatment-related mortality rates are as high as 3–4% using some combinations in these poor-risk patient groups. For most patients, treatment is palliative, aimed at delaying or relieving cancer-related symptoms, because few patients experience durable complete remissions.

CHEMOTHERAPY

A number of chemotherapeutic drugs have activity as single agents; cisplatin, paclitaxel, and gemcitabine are considered most active. Standard therapy consists of two-, three-, or four-drug combinations. Overall response rates of >50% have been reported using combinations such as methotrexate, vinblastine, doxorubicin, and cisplatin (MVAC); gemcitabine and cisplatin (GC); or gemcitabine, paclitaxel, and cisplatin (GPC). MVAC was considered standard, but the toxicities of neutropenia and fever, mucositis, diminished renal and auditory function, and peripheral neuropathy led to the development of alternative regimens. At present, GC is used more commonly than MVAC based on the results of a comparative trial of MVAC versus GC that showed less neutropenia and fever and less mucositis for the GC regimen with similar response rates and median overall survival. Anemia and thrombocytopenia were more common with GC. GPC is not more effective than GC.

Chemotherapy has also been tested in the neoadjuvant and adjuvant settings. In a randomized trial, patients receiving three cycles of neoadjuvant MVAC followed by cystectomy had a significantly better median (6.2 years) and 5-year survival (57%) compared to cystectomy alone (median survival 3.8 years; 5-year survival 42%). Similar results were obtained in an international study of three cycles of cisplatin, methotrexate, and vinblastine (CMV) followed by either radical cystectomy or radiation therapy. The decision to administer adjuvant therapy is based on recurrence risk after cystectomy. Studies of adjuvant chemotherapy have been underpowered, and most closed for lack of accrual. One underpowered study using the GPC regimen suggested that adjuvant treatment improved survival, although many patients never received chemotherapy for metastases. Another underpowered study did not show a benefit for GC chemotherapy. Therefore, preoperative chemotherapy is preferred when medically appropriate. Indications for adjuvant chemotherapy in patients who did not receive neoadjuvant treatment include nodal disease, extravesical tumor extension, or vascular invasion in the resected specimen.

The management of bladder cancer is summarized in Table 114-2.

CARCINOMA OF THE RENAL PELVIS AND URETER

About 5000 cases of renal pelvis and ureter cancer occur each year; nearly all are transitional cell carcinomas similar to bladder cancer in biology and appearance. This tumor is associated with chronic phenacetin abuse and aristolochic acid consumption in Chinese herbal preparations; aristolochic acid also seems to be associated with Balkan nephropathy, a chronic interstitial nephritis endemic in Bulgaria, Greece, Bosnia-Herzegovina,

TABLE 114-1 SURVIVAL FOLLOWING SURGERY FOR BLADDER CANCER

Pathologic Stage	5-Year Survival, %	10-Year Survival, %
T2,N0	89	87
T3a,N0	78	76
T3b,N0	62	61
T4,N0	50	45
Any T,N1	35	34

TABLE 114-2 MANAGEMENT OF BLADDER CANCER

Nature of Lesion	Management Approach
Non-muscle-invasive disease	Endoscopic removal, usually with intravesical therapy
Muscle-invasive disease	Cystectomy \pm systemic chemotherapy (before or after surgery)
Metastatic disease	Curative or palliative chemotherapy (based on prognostic factors) \pm surgery

578 and Romania. In addition, upper tract urothelial carcinoma is linked to hereditary nonpolyposis colorectal cancer.

The most common symptom is painless gross hematuria, and the disease is usually detected on imaging during the workup for hematuria. Patterns of spread are like bladder cancer. For low-grade disease localized to the renal pelvis and ureter, nephroureterectomy (including excision of the distal ureter with a portion of the bladder) is associated with 5-year survival of 80–90%. More invasive or poorly differentiated tumors are more likely to recur locally and to metastasize. Metastatic disease is treated with the chemotherapy used in bladder cancer, and the outcome is similar to that of metastatic bladder cancer.

RENAL CELL CARCINOMA

Renal cell carcinomas account for 90–95% of malignant neoplasms arising from the kidney. Notable features include resistance to cytotoxic agents, infrequent responses to biologic response modifiers such as interleukin (IL) 2, robust activity to antiangiogenesis targeted agents, and a variable clinical course for patients with metastatic disease, including anecdotal reports of spontaneous regression.

EPIDEMIOLOGY

The incidence of renal cell carcinoma continues to rise and is now nearly 65,000 cases annually in the United States, resulting in 13,700 deaths. The male-to-female ratio is 2:1. Incidence peaks between the ages of 50 and 70 years, although this malignancy may be diagnosed at any age. Many environmental factors have been investigated as possible contributing causes; the strongest association is with cigarette smoking. Risk is also increased for patients who have acquired cystic disease of the kidney associated with end-stage renal disease and for those with tuberous sclerosis. Most cases are sporadic, although familial forms have been reported. One is associated with von Hippel-Lindau (VHL) syndrome. VHL syndrome is an autosomal dominant disorder. Genetic studies identified the *VHL* gene on the short arm of chromosome 3. Approximately 35% of individuals with VHL disease develop clear cell renal cell carcinoma. Other associated neoplasms include retinal hemangioma, hemangioblastoma of the spinal cord and cerebellum, pheochromocytoma, neuroendocrine tumors and cysts, and cysts in the epididymis of the testis in men and the broad ligament in women.

PATHOLOGY AND GENETICS

Renal cell neoplasia represents a heterogeneous group of tumors with distinct histopathologic, genetic, and clinical features ranging from benign to high-grade malignant (Table 114-3). They are classified on the basis of morphology and histology. Categories include clear cell carcinoma (60% of cases), papillary tumors (5–15%), chromophobe tumors (5–10%), oncocytomas (5–10%), and collecting or Bellini duct tumors (<1%). Papillary tumors tend to be bilateral and multifocal. Chromophobe tumors have a more indolent clinical course, and oncocytomas are considered benign neoplasms. In contrast, Bellini duct carcinomas, which are thought to arise from the collecting ducts within the renal medulla, are rare but often very aggressive. Clear cell tumors, the predominant histology, are found in >80% of patients who

develop metastases. Clear cell tumors arise from the epithelial cells of the proximal tubules and usually show chromosome 3p deletions. Deletions of 3p21–26 (where the *VHL* gene maps) are identified in patients with familial as well as sporadic tumors. *VHL* encodes a tumor suppressor protein that is involved in regulating the transcription of vascular endothelial growth factor (VEGF), platelet-derived growth factor (PDGF), and a number of other hypoxia-inducible proteins. Inactivation of *VHL* leads to overexpression of these agonists of the VEGF and PDGF receptors, which promote tumor angiogenesis and tumor growth. Agents that inhibit proangiogenic growth factor activity show antitumor effects. Enormous genetic variability has been documented in tumors from individual patients. Although the tumors have a clear clonal origin and often contain *VHL* mutations in common, different portions of the primary tumor and different metastatic sites may have wide variation in genetic lesions they contain. This tumor heterogeneity may underlie the emergence of treatment resistance.

CLINICAL PRESENTATION

The presenting signs and symptoms include hematuria, abdominal pain, and a flank or abdominal mass. Other symptoms are fever, weight loss, anemia, and a varicocele. The tumor is most commonly detected as an incidental finding on a radiograph. Widespread use of radiologic cross-sectional imaging procedures (CT, ultrasound, MRI) contributes to earlier detection, including incidental renal masses detected during evaluation for other medical conditions. The increasing number of incidentally discovered low-stage tumors has contributed to an improved 5-year survival for patients with renal cell carcinoma and increased use of nephron-sparing surgery (partial nephrectomy). A spectrum of paraneoplastic syndromes has been associated with these malignancies, including erythrocytosis, hypercalcemia, nonmetastatic hepatic dysfunction (Stauffer's syndrome), and acquired dysfibrinogenemia. Erythrocytosis is noted at presentation in only about 3% of patients. Anemia, a sign of advanced disease, is more common.

The standard evaluation of patients with suspected renal cell tumors includes a CT scan of the abdomen and pelvis, chest radiograph, urine analysis, and urine cytology. If metastatic disease is suspected from the chest radiograph, a CT of the chest is warranted. MRI is useful in evaluating the inferior vena cava in cases of suspected tumor involvement or invasion by thrombus. In clinical practice, any solid renal masses should be considered malignant until proven otherwise; a definitive diagnosis is required. If no metastases are demonstrated, surgery is indicated, even if the renal vein is invaded. The differential diagnosis of a renal mass includes cysts, benign neoplasms (adenoma, angiomyolipoma, oncocytoma), inflammatory lesions (pyelonephritis or abscesses), and other primary or metastatic cancers. Other malignancies that may involve the kidney include transitional cell carcinoma of the renal pelvis, sarcoma, lymphoma, and Wilms' tumor. All of these are less common causes of renal masses than is renal cell cancer.

STAGING AND PROGNOSIS

Staging is based on the American Joint Committee on Cancer (AJCC) staging system (Fig. 114-2). Stage I tumors are <7 cm in greatest diameter and confined to the kidney, stage II tumors are ≥7 cm and confined to the kidney, stage III tumors extend through the renal capsule but are confined to Gerota's fascia (IIIa) or involve a single hilar lymph node (N1), and stage IV disease includes tumors that have invaded adjacent organs (excluding the adrenal gland) or involve multiple lymph nodes or distant metastases. The 5-year survival rate varies by stage: >90% for stage I, 85% for stage II, 60% for stage III, and 10% for stage IV.

TREATMENT RENAL CELL CARCINOMA

LOCALIZED TUMOR

The standard management for stage I or II tumors and selected cases of stage III disease is radical or partial nephrectomy. A radical nephrectomy involves en bloc removal of Gerota's fascia and

TABLE 114-3 CLASSIFICATION OF EPITHELIAL NEOPLASMS ARISING FROM THE KIDNEY

Carcinoma Type	Growth Pattern	Cell of Origin	Cytogenetics
Clear cell	Acinar or sarcomatoid	Proximal tubule	3p-, 5q+, 14q-
Papillary	Papillary or sarcomatoid	Proximal tubule	+7, +17, -Y
Chromophobe	Solid, tubular, or sarcomatoid	Distal tubules/cortical collecting duct	Whole arm losses (1, 2, 6, 10, 13, 17, and 21)
Oncocytic	Tumor nests	Cortical collecting duct	Undetermined
Collecting duct	Papillary or sarcomatoid	Medullary collecting duct	Undetermined

TNM	Involvement	Extent of Disease	Anatomic Stage / Prognostic Groups			
T0	Primary not involved		I	T1	N0	M0
T1	≤ 7 cm	limited to kidney	II	T2	N0	M0
T1a	≤ 4 cm		III	T1 or T2	N1	M0
T1b	> 4 cm		III	T3	N0 or N1	M0
T2	> 7 cm	limited to kidney	IV	T4	Any N	M0
T2a	> 7 cm to ≤ 10 cm		IV	Any T	Any N	M1
T2b	> 10 cm					
T3	into major veins or perinephric tissues	not beyond Gerota's fascia				
T3a	in renal vein or renal sinus fat	not beyond Gerota's fascia				
T3b	into vena cava	below diaphragm				
T3c	into vena cava	above diaphragm				
T4	invasion beyond Gerota's fascia	including contiguous extensions & into ipsilateral adrenal gland				
Regional						
NX	Regional lymph nodes not assessed					
N0	No lymph node involvement					
N1	Regional lymph node involvement					
Distant Metastases						
M0	No distant metastases					
M1	Distant metastases					

FIGURE 114-2 Renal cell carcinoma staging. TNM, tumor, node, metastasis.

its contents, including the kidney, the ipsilateral adrenal gland in some cases, and adjacent hilar lymph nodes. The role of a regional lymphadenectomy is controversial. Extension into the renal vein or inferior vena cava (stage III disease) does not preclude resection even if cardiopulmonary bypass is required. If the tumor is resected, half of these patients have prolonged survival.

Nephron-sparing approaches via open or laparoscopic surgery may be appropriate for patients who have only one kidney, depending on the size and location of the lesion. A nephron-sparing approach can also be used for patients with bilateral tumors. Partial nephrectomy techniques are applied electively to resect small masses for patients with a normal contralateral kidney. Adjuvant therapy following this surgery does not improve outcome, even in cases with a poor prognosis.

ADVANCED DISEASE

Surgery has a limited role for patients with metastatic disease. Long-term survival may occur in patients who relapse after nephrectomy in a solitary site that is removed. One indication for nephrectomy with metastases at initial presentation is to alleviate pain or hemorrhage of a primary tumor. Also, a cytoreductive nephrectomy before systemic treatment improves survival for carefully selected patients with stage IV tumors.

Metastatic renal cell carcinoma is refractory to chemotherapy. Cytokine therapy with IL-2 or interferon α (IFN- α) produces regression in 10–20% of patients. IL-2 produces durable complete remission in a small proportion of cases. In general, cytokine therapy is considered unsatisfactory for most patients.

The situation changed dramatically when two large-scale randomized trials established a role for antiangiogenic therapy, as predicted by the genetic studies. These trials separately evaluated two orally administered antiangiogenic agents, sorafenib and sunitinib, that inhibited receptor tyrosine kinase signaling through the VEGF and PDGF receptors. Both showed efficacy as second-line treatment following progression during cytokine treatment, resulting in approval by regulatory authorities for the treatment of advanced renal cell carcinoma. A randomized phase III trial comparing sunitinib to IFN- α showed superior efficacy for sunitinib with an acceptable safety profile. The trial resulted in a change in the standard first-line treatment from IFN to sunitinib. Sunitinib is usually given orally at a dose of 50 mg/d for 4 out of 6 weeks. Pazopanib and axitinib are newer agents of the same class. Pazopanib was compared to sunitinib in a randomized first-line phase III trial. Efficacy was similar, and there was less fatigue and skin toxicity, resulting in better quality of life scores for pazopanib compared with sunitinib. Temeosolimus and everolimus, inhibitors of the mammalian target

of rapamycin (mTOR), show activity in patients with untreated poor-prognosis tumors and in sunitinib/sorafenib-refractory tumors. Patients benefit from the sequential use of axitinib and everolimus following progression to sunitinib or pazopanib first-line therapy.

The prognosis of metastatic renal cell carcinoma is variable. In one analysis, no prior nephrectomy, a KPS <80, low hemoglobin, high corrected calcium, and abnormal lactate dehydrogenase were poor prognostic factors. Patients with zero, one or two, and three or more factors had a median survival of 24, 12, and 5 months, respectively. These tumors may follow an unpredictable and protracted clinical course. It may be best to document progression before considering systemic treatment.

115 Benign and Malignant Diseases of the Prostate

Howard I. Scher, James A. Eastham

Benign and malignant changes in the prostate increase with age. Autopsies of men in the eighth decade of life show hyperplastic changes in >90% and malignant changes in >70% of individuals. The high prevalence of these diseases among the elderly, who often have competing causes of morbidity and mortality, mandates a risk-adapted approach to diagnosis and treatment. This can be achieved by considering these diseases as a series of states. Each state represents a distinct clinical milestone for which therapy(ies) may be recommended based on current symptoms, the risk of developing symptoms, or death from disease in relation to death from other causes within a given time frame. For benign proliferative disorders, symptoms of urinary frequency, infection, and potential for obstruction are weighed against the side effects and complications of medical or surgical intervention. For prostate malignancies, the risks of developing the disease, symptoms, or death from cancer are balanced against the morbidities of the recommended treatments and preexisting comorbidities.

ANATOMY

The prostate is located in the pelvis and is surrounded by the rectum, the bladder, the periprostatic and dorsal vein complexes and neurovascular bundles that are responsible for erectile function, and the urinary sphincter that is responsible for passive urinary control. The

580 prostate is composed of branching tubuloalveolar glands arranged in lobules surrounded by fibromuscular stroma. The acinar unit includes an epithelial compartment made up of epithelial, basal, and neuroendocrine cells and separated by a basement membrane, and a stromal compartment that includes fibroblasts and smooth-muscle cells. Prostate-specific antigen (PSA) and prostatic acid phosphatase (PAP) are produced in the epithelial cells. Both prostate epithelial cells and stromal cells express androgen receptors (ARs) and depend on androgens for growth. Testosterone, the major circulating androgen, is converted by the enzyme 5 α -reductase to dihydrotestosterone in the gland.

The periurethral portion of the gland increases in size during puberty and after the age of 55 years due to the growth of nonmalignant cells in the transition zone of the prostate that surrounds the urethra. Most cancers develop in the peripheral zone, and cancers in this location may be palpated during a digital rectal examination (DRE).

PROSTATE CANCER

In 2013, approximately 238,590 prostate cancer cases were diagnosed, and 29,720 men died from prostate cancer in the United States. The absolute number of prostate cancer deaths has decreased in the past 5 years, which has been attributed by some to the widespread use of PSA-based detection strategies. However, the benefit of screening on survival is unclear. The paradox of management is that although 1 in 6 men will eventually be diagnosed with the disease, and the disease remains the second leading cause of cancer deaths in men, only 1 man in 30 with prostate cancer will die of his disease.

EPIDEMIOLOGY

Epidemiologic studies show that the risk of being diagnosed with prostate cancer increases by a factor of two if one first-degree relative is affected and by four if two or more are affected. Current estimates are that 40% of early-onset and 5–10% of all prostate cancers are hereditary. Prostate cancer affects ethnic groups differently. Matched for age, African-American males have both a higher incidence of prostate cancer and larger tumors and more worrisome histologic features than white males. Polymorphic variants of the AR, the cytochrome P450 C17, and the steroid 5 α -reductase type II (SRD5A2) genes have been implicated in the variations in incidence.

The prevalence of autopsy-detected cancers is similar around the world, while the incidence of clinical disease varies. Thus, environmental and dietary factors may play a role in prostate cancer growth and progression. High consumption of dietary fats, such as α -linolenic acid or the polycyclic aromatic hydrocarbons that form when red meats are cooked, is believed to increase risk. Similar to breast cancer in Asian women, the risk of prostate cancer in Asian men increases when they move to Western environments. Protective factors include consumption of the isoflavonoid genistein (which inhibits 5 α -reductase) found in many legumes, cruciferous vegetables that contain the isothiocyanate sulforaphane, retinoids such as lycopene found in tomatoes, and inhibitors of cholesterol biosynthesis (e.g., statin drugs). The development of prostate cancer is a multistep process. One early change is hypermethylation of the *GSTP1* gene promoter, which leads to loss of function of a gene that detoxifies carcinogens. The finding that many prostate cancers develop adjacent to a lesion termed proliferative inflammatory atrophy (PIA) suggests a role for inflammation.

PREVENTION

Currently no drugs or dietary supplements are approved by the U.S. Food and Drug Administration (FDA) for prevention of prostate cancer, nor are any recommended by the major clinical guidelines. Although statins may have some protective effect, the potential risks outweigh the benefits given the small number of men who die of prostate cancer. The results from several large, double-blind, randomized chemoprevention trials established 5 α -reductase inhibitors (5ARI) as the most likely therapy to reduce the future risk of a prostate cancer diagnosis. The Prostate Cancer Prevention Trial (PCPT), in which men older than age 55 years received placebo or the 5ARI finasteride,

which inhibits the type 1 isoform, showed a 25% (95% confidence interval 19–31%) reduction in the period prevalence of prostate cancer across all age groups in favor of finasteride (18.4%) over placebo (24.4%). In the Reduction by Dutasteride of Prostate Cancer Events (REDUCE) trial, a similar 23% reduction in the 4-year period prevalence was observed in favor of dutasteride ($p = .001$). Dutasteride inhibits both the type 1 and type 2 5ARI isoforms. While both studies met their endpoint, there was concern that most of the cancers that were prevented were low risk and that there was a slightly higher rate of clinically significant cancers (those with higher Gleason score) in the treatment arm. Neither drug was FDA-approved for prostate cancer prevention. In comparison, the Selenium and Vitamin E Cancer Prevention Trial (SELECT), which enrolled African-American men age ≥ 50 years and others age ≥ 55 years, showed no difference in cancer incidence in patients receiving vitamin E (4.6%) or selenium (4.9%) alone or in combination (4.6%) relative to placebo (4.4%). A similar lack of benefit for vitamin E, vitamin C, and selenium was seen in the Physicians Health Study II.

THE CLINICAL STATES MODEL

The prostate cancer continuum—from the appearance of a preneoplastic and invasive lesion localized to the prostate, to a metastatic lesion that results in symptoms and, ultimately, mortality—can span decades. To facilitate disease management, competing risks are considered in the context of a series of clinical states (Fig. 115-1). The states are defined operationally on the basis of whether or not a cancer diagnosis has been established and, for those with a diagnosis, whether or not metastases are detectable on imaging studies and the measured level of testosterone in the blood. With this approach, an individual resides in only one state and remains in that state until he has progressed. At each assessment, the decision to offer treatment and the specific form of treatment are based on the risk posed by the cancer relative to competing causes of mortality that may be present in that individual. It follows that the more advanced the disease, the greater is the need for treatment.

For those without a cancer diagnosis, the decision to undergo testing to detect a cancer is based on the individual's estimated life expectancy and, separately, the probability that a clinically significant cancer may be present. For those with a prostate cancer diagnosis, the clinical states model considers the probability of developing symptoms or dying from prostate cancer. Thus, a patient with localized prostate cancer who has had all cancer removed surgically remains in the state of localized disease as long as the PSA remains undetectable. The time within a state becomes a measure of the efficacy of an intervention, although the effect may not be assessable for years. Because many men with active cancer are not at risk for metastases, symptoms, or death, the clinical states model allows a distinction between *cure*—the elimination of all cancer cells, the primary therapeutic objective when treating most cancers—and *cancer control*, in which the tempo of the illness is altered and symptoms are controlled until the patient dies of other causes. These can be equivalent therapeutically from a patient standpoint if the patient has not experienced symptoms of the disease or the treatment needed to control it. Even when a recurrence is documented, immediate therapy is not always necessary. Rather, as at the time of diagnosis, the need for intervention is based on the tempo of the illness as it unfolds in the individual, relative to the risk-to-benefit ratio of the therapy being considered.

SCREENING AND DIAGNOSIS

Physical Examination The need to pursue a diagnosis of prostate cancer is based on symptoms, an abnormal DRE, or, more typically, a change in or an elevated serum PSA. The urologic history should focus on symptoms of outlet obstruction, continence, potency, or change in ejaculatory pattern.

The DRE focuses on prostate size and consistency and abnormalities within or beyond the gland. Many cancers occur in the peripheral zone and may be palpated on DRE. Carcinomas are characteristically hard, nodular, and irregular, while induration may also be due to benign prostatic hypertrophy (BPH) or calculi. Overall, 20–25% of men with an abnormal DRE have cancer.

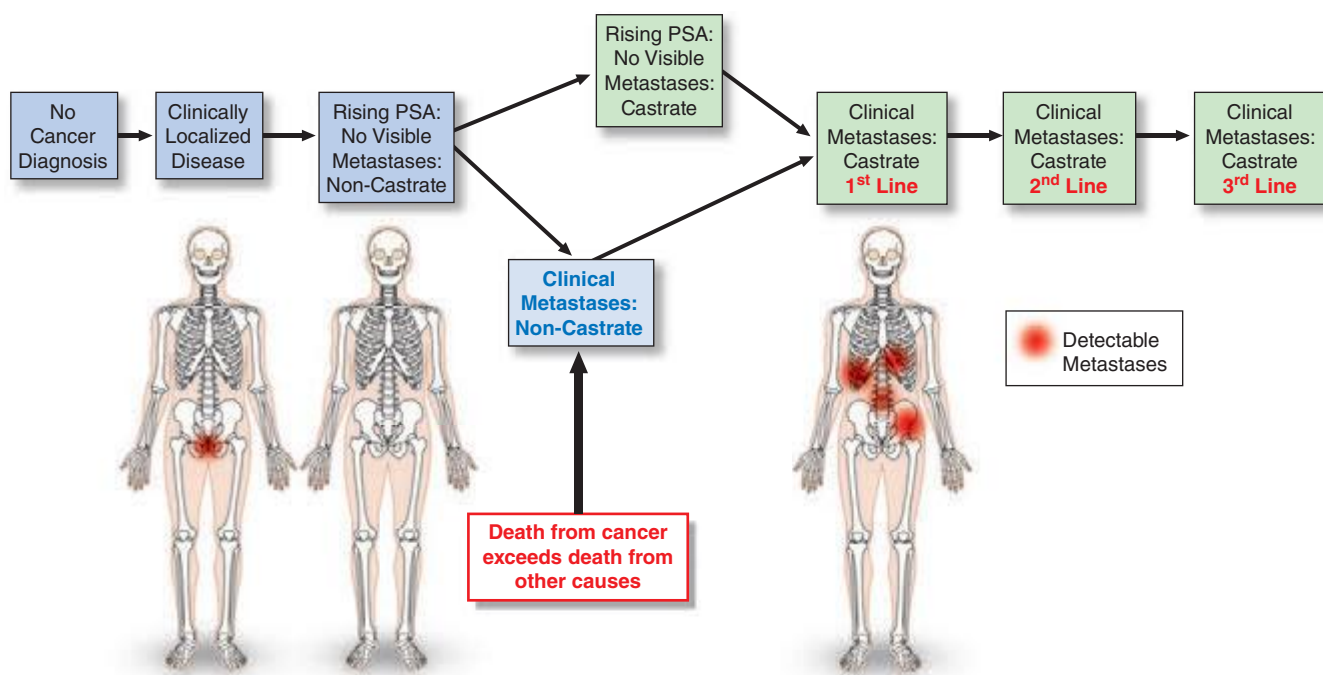


FIGURE 115-1 Clinical states of prostate cancer. PSA, prostate-specific antigen.

Prostate-Specific Antigen PSA (kallikrein-related peptidase 3; *KLK3*) is a kallikrein-related serine protease that causes liquefaction of seminal coagulum. It is produced by both nonmalignant and malignant epithelial cells and, as such, is prostate-specific, not prostate cancer-specific. Serum levels may also increase from prostatitis and BPH. Serum levels are not significantly affected by DRE, but the performance of a prostate biopsy can increase PSA levels up to tenfold for 8–10 weeks. PSA circulating in the blood is inactive and mainly occurs as a complex with the protease inhibitor α_1 -antichymotrypsin and as free (unbound) PSA forms. The formation of complexes between PSA, α_2 -macroglobulin, or other protease inhibitors is less significant. Free PSA is rapidly eliminated from the blood by glomerular filtration with an estimated half-life of 12–18 h. Elimination of PSA bound to α_1 -antichymotrypsin is slow (estimated half-life of 1–2 weeks) because it too is largely cleared by the kidneys. Levels should be undetectable after about 6 weeks if the prostate has been removed. Immunohistochemical staining for PSA can be used to establish a prostate cancer diagnosis.

PSA-BASED SCREENING AND EARLY DETECTION PSA testing was approved by the U.S. FDA in 1994 for early detection of prostate cancer, and the widespread use of the test has played a significant role in the proportion of men diagnosed with early-stage cancers: more than 70–80% of newly diagnosed cancers are clinically organ-confined. The level of PSA in blood is strongly associated with the risk and outcome of prostate cancer. A single PSA measured at age 60 is associated (area under the curve [AUC] of 0.90) with lifetime risk of death from prostate cancer. Most prostate cancer deaths (90%) occur among men with PSA levels in the top quartile (>2 ng/mL), although only a minority of men with PSA >2 ng/mL will develop lethal prostate cancer. Despite this and mortality rate reductions reported from large randomized prostate cancer screening trials, routine use of the test remains controversial.

The U.S. Preventive Services Task Force (USPSTF) reviewed the evidence for screening for prostate cancer and made a clear recommendation against screening. By giving a grade of “D” in the recommendation statement that was based on this review, the USPSTF concluded that “there is moderate or high certainty that this service has no net benefit or that the harms outweigh the benefits.” Whether the harms of screening, overdiagnosis, and overtreatment are justified by the benefits in terms of reduced prostate cancer mortality is open to reasonable doubt. In response to the USPSTF, the American Urological Association (AUA) updated their consensus statement regarding prostate cancer screening.

They concluded that the quality of evidence for the benefits of screening was moderate, and evidence for harm was high for men age 55–69 years. For men outside this age range, evidence was lacking for benefit, but the harms of screening, including overdiagnosis and overtreatment, remained. The AUA recommends shared decision making considering PSA-based screening for men age 55–69, a target age group for whom benefits may outweigh harms. Outside this age range, PSA-based screening as a routine test was not recommended based on the available evidence. The entire guideline is available at www.AUAnet.org/education/guidelines/prostate-cancer-detection.cfm.

The PSA criteria used to recommend a diagnostic prostate biopsy have evolved over time. However, based on the commonly used cut point for prostate biopsy (a total PSA ≥ 4 ng/mL), most men with a PSA elevation do not have histologic evidence of prostate cancer at biopsy. In addition, many men with PSA levels below this cut point harbor cancer cells in their prostate. Information from the PCPT demonstrates that there is no PSA below which the risk of prostate cancer is zero. Thus, the PSA level establishes the likelihood that a man will harbor cancer if he undergoes a prostate biopsy. The goal is to increase the sensitivity of the test for younger men more likely to die of the disease and to reduce the frequency of detecting cancers of low malignant potential in elderly men more likely to die of other causes. Patients with symptomatic prostatitis should have a course of antibiotics before biopsy. However, the routine use of antibiotics in an asymptomatic man with an elevated PSA level is strongly discouraged.

Prostate Biopsy A diagnosis of cancer is established by an image-guided needle biopsy. Direct visualization by transrectal ultrasound (TRUS) or magnetic resonance imaging (MRI) assures that all areas of the gland are sampled. Contemporary schemas advise an extended-pattern 12-core biopsy that includes sampling from the peripheral zone as well as a lesion-directed palpable nodule or suspicious image-guided sampling. Men with an abnormal PSA and negative biopsy are advised to undergo a repeat biopsy.

BIOPSY PATHOLOGY Each core of the biopsy is examined for the presence of cancer, and the amount of cancer is quantified based on the length of the cancer within the core and the percentage of the core involved. Of the cancers identified, $>95\%$ are adenocarcinomas; the rest are squamous or transitional cell tumors or, rarely, carcinosarcomas. Metastases to the prostate are rare, but in some cases colon cancers or transitional cell tumors of the bladder invade the gland by direct extension.

TABLE 115-1 TNM CLASSIFICATION

TNM Staging System for Prostate Cancer ^a	
Tx	Primary tumor cannot be assessed
T0	No evidence of primary tumor
Localized Disease	
T1	Clinically inapparent tumor, neither palpable nor visible by imaging
T1a	Tumor incidental histologic finding in ≤5% of resected tissue; not palpable
T1b	Tumor incidental histologic finding in >5% of resected tissue
T1c	Tumor identified by needle biopsy (e.g., because of elevated PSA)
T2	Tumor confined within prostate ^b
T2a	Tumor involves half of one lobe or less
T2b	Tumor involves more than one half of one lobe, not both lobes
T2c	Tumor involves both lobes
Local Extension	
T3	Tumor extends through the prostate capsule ^c
T3a	Extracapsular extension (unilateral or bilateral)
T3b	Tumor invades seminal vesicle(s)
T4	Tumor is fixed or invades adjacent structures other than seminal vesicles such as external sphincter, rectum, bladder, levator muscles, and/or pelvic wall.
Metastatic Disease	
N1	Positive regional lymph nodes
M1	Distant metastases

^aRevised from SB Edge et al (eds): *AJCC Cancer Staging Manual*, 7th ed. New York, Springer, 2010. ^bTumor found in one or both lobes by needle biopsy, but not palpable or reliably visible by imaging, is classified as T1c. ^cInvasion into the prostatic apex or into (but not beyond) the prostatic capsule is classified not as T3 but as T2.

Abbreviations: PSA, prostate-specific antigen; TNM, tumor, node, metastasis.

When prostate cancer is diagnosed, a measure of histologic aggressiveness is assigned using the *Gleason grading system*, in which the dominant and secondary glandular histologic patterns are scored from 1 (well-differentiated) to 5 (undifferentiated) and summed to give a total score of 2–10 for each tumor. The most poorly differentiated area of tumor (i.e., the area with the highest histologic grade) often determines biologic behavior. The presence or absence of perineural invasion and extracapsular spread is also recorded.

Prostate Cancer Staging The tumor, node, metastasis (TNM) staging system includes categories for cancers identified solely on the basis of an abnormal PSA (T1c), those that are palpable but clinically confined to the gland (T2), and those that have extended outside the gland (T3 and T4) (Table 115-1, Fig. 115-2). DRE alone is inaccurate in determining the extent of disease within the gland, the presence or absence of capsular

invasion, involvement of seminal vesicles, and extension of disease to lymph nodes. Because of the inadequacy of DRE for staging, the TNM staging system was modified to include the results of imaging. Unfortunately, no single test has proven to accurately indicate the stage or the presence of organ-confined disease, seminal vesicle involvement, or lymph node spread.

TRUS is the imaging technique most frequently used to assess the primary tumor, but its chief use is directing prostate biopsies, not staging. No TRUS finding consistently indicates cancer with certainty. Computed tomography (CT) lacks sensitivity and specificity to detect extraprostatic extension and is inferior to MRI in visualization of lymph nodes. In general, MRI performed with an endorectal coil is superior to CT to detect cancer in the prostate and to assess local disease extent. T1-weighted MRI produces a high signal in the periprostatic fat, periprostatic venous plexus, perivesicular tissues, lymph nodes, and bone marrow. T2-weighted MRI demonstrates the internal architecture of the prostate and seminal vesicles. Most cancers have a low signal, while the normal peripheral zone has a high signal, although the technique lacks sensitivity and specificity. MRI is also useful for the planning of surgery and radiation therapy.

Radionuclide bone scans (bone scintigraphy) are used to evaluate spread to osseous sites. This test is sensitive but relatively nonspecific because areas of increased uptake are not always related to metastatic disease. Healing fractures, arthritis, Paget's disease, and other conditions will also cause abnormal uptake. True-positive bone scans are uncommon when the PSA is <10 ng/mL unless the tumor is high grade.

TREATMENT TREATMENT OF PROSTATE CANCER BY CLINICAL STATE

CLINICALLY LOCALIZED PROSTATE CANCER

Clinically localized prostate cancers are those that appear to be nonmetastatic after staging studies are performed. Patients with clinically localized disease are managed by radical prostatectomy, radiation therapy, or active surveillance. Choice of therapy requires the consideration of several factors: the presence of symptoms, the probability that the untreated tumor will adversely affect the quality or duration of survival and thus require treatment, and the probability that the tumor can be cured by single-modality therapy directed at the prostate or that it will require both local and systemic therapy to achieve cure.

Data from the literature do not provide clear evidence for the superiority of any one treatment relative to another. Comparison of outcomes of various forms of therapy is limited by the lack of prospective trials, referral bias, the experience of the treating teams, and differences in endpoints and cancer control definitions. Often, PSA relapse-free survival is used because an effect on metastatic progression

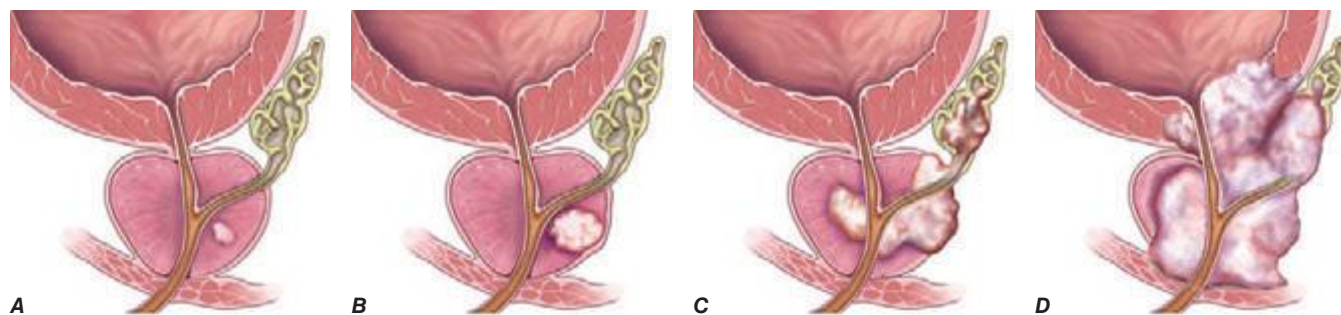


FIGURE 115-2 T stages of prostate cancer. (A) T1—Clinically inapparent tumor, neither palpable nor visible by imaging; (B) T2—Tumor confined within prostate; (C) T3—Tumor extends through prostate capsule and may invade the seminal vesicles; (D) T4—Tumor is fixed or invades adjacent structures. Eighty-one percent of patients present with local disease (T1 and T2), which is associated with a 5-year survival rate of 100%. An additional 12% of patients present with regional disease (T3 and T4 without metastases), which is also associated with a 100% survival rate after 5 years. Four percent of patients present with distant disease (T4 with metastases), which is associated with a 28% 5-year survival rate. (Three percent of patients are ungraded, and this group is associated with a 73% 5-year survival rate.) (Data from AJCC, <http://seer.cancer.gov/statfacts/html/prost.html>. Figure © 2014 Memorial Sloan-Kettering Cancer Center; used with permission.)

or survival may not be apparent for years. After radical surgery to remove all prostate tissue, PSA should become undetectable in the blood within 6 weeks. If PSA remains or becomes detectable after radical prostatectomy, the patient is considered to have persistent disease. After radiation therapy, in contrast, PSA does not become undetectable because the remaining nonmalignant elements of the gland continue to produce PSA even if all cancer cells have been eliminated. Similarly, cancer control is not well defined for a patient managed by active surveillance because PSA levels will continue to rise in the absence of therapy. Other outcomes are time to objective progression (local or systemic), cancer-specific survival, and overall survival; however, these outcomes may take years to assess.

The more advanced the disease, the lower the probability of local control and the higher the probability of systemic relapse. More important is that within the categories of T1, T2, and T3 disease are cancers with a range of prognoses. Some T3 tumors are curable with therapy directed solely at the prostate, and some T1 lesions have a high probability of systemic relapse that requires the integration of local and systemic therapy to achieve cure. For T1c cancers in particular, stage alone is inadequate to predict outcome and select treatment; other factors must be considered.

Nomograms To better assess risk and guide treatment selection, many groups have developed prognostic models or nomograms that use a combination of the initial clinical T stage, biopsy Gleason score, and baseline PSA. Some use discrete cut points (PSA <10 or ≥ 10 ng/mL; Gleason score of ≤ 6 , 7, or ≥ 8); others employ nomograms that use PSA and Gleason score as continuous variables. More than 100 nomograms have been reported to predict the probability that a clinically significant prostate cancer is present, disease extent (organ-confined vs non-organ-confined, node-negative or -positive), or the probability of success of treatment for specific local therapies using pretreatment variables. Considerable controversy exists over what constitutes “high risk” based on a predicted probability of success or failure. In these situations, nomograms and predictive models can only go so far. Exactly what probability of success or failure would lead a physician to recommend and a patient to seek alternative approaches is controversial. As an example, it may be appropriate to recommend radical surgery for a younger patient with a low probability of cure. Nomograms are being refined continually to incorporate additional clinical parameters, biologic determinants, and year of treatment, which can also affect outcomes, making treatment decisions a dynamic process.

Treatment-Related Adverse Events The frequency of adverse events varies by treatment modality and the experience of the treating team. For example, following radical prostatectomy, incontinence rates range from 2–47% and impotence rates range from 25–89%. Part of the variability relates to how the complication is defined and whether the patient or physician is reporting the event. The time of the assessment is also important. After surgery, impotence is immediate but may reverse over time, while with radiation therapy impotence is not immediate but may develop over time. Of greatest concern to patients are the effects on continence, sexual potency, and bowel function.

Radical Prostatectomy The goal of radical prostatectomy is to excise the cancer completely with a clear margin, to maintain continence by preserving the external sphincter, and to preserve potency by sparing the autonomic nerves in the neurovascular bundle. The procedure is advised for patients with a life expectancy of 10 years or more and is performed via a retropubic or perineal approach or via a minimally invasive robotic-assisted or hand-held laparoscopic approach. Outcomes can be predicted using postoperative nomograms that consider pretreatment factors and the pathologic findings at surgery. PSA failure is usually defined as a value greater than 0.1 or 0.2 ng/mL. Specific criteria to guide the choice of one approach over another are lacking. Minimally invasive approaches offer the advantage of a shorter hospital stay and reduced blood loss. Rates of cancer control, recovery of continence, and recovery of erectile function are comparable between open and minimally

invasive approaches. The individual surgeon rather than the surgical approach used is most important in determining outcomes after surgery.

Neoadjuvant hormonal therapy has also been explored in an attempt to improve the outcomes of surgery for high-risk patients, using a variety of definitions. The results of several large trials testing 3 or 8 months of androgen depletion before surgery showed that serum PSA levels decreased by 96%, prostate volumes decreased by 34%, and margin positivity rates decreased from 41% to 17%. Unfortunately, hormones did not produce an improvement in PSA relapse-free survival. Thus, neoadjuvant hormonal therapy is not recommended.

Factors associated with incontinence following radical prostatectomy include older age and urethral length, which impacts the ability to preserve the urethra beyond the apex and the distal sphincter. The skill and experience of the surgeon are also factors. Recovery of erectile function is associated with younger age, quality of erections before surgery, and the absence of damage to the neurovascular bundles. In general, erectile function begins to return about 6 months after surgery if both neurovascular bundles are preserved. Potency is reduced by half if at least one neurovascular bundle is sacrificed. Overall, with the availability of drugs such as phosphodiesterase-5 (PDE5) inhibitors, intraurethral inserts of alprostadil, and intracavernosal injections of vasodilators, many patients recover satisfactory sexual function.

Radiation Therapy Radiation therapy is given by external beam, by radioactive sources implanted into the gland, or by a combination of the two techniques.

EXTERNAL-BEAM RADIATION THERAPY Contemporary external-beam radiation therapy requires three-dimensional conformal treatment plans to maximize the dose to the prostate and to minimize the exposure of the surrounding normal tissue. Intensity-modulated radiation therapy (IMRT) permits shaping of the dose and allows the delivery of higher doses to the prostate and a further reduction in normal tissue exposure than three-dimensional conformal treatment alone. These advances have enabled the safe administration of doses >80 Gy and resulted in higher local control rates and fewer side effects.

Cancer control after radiation therapy has been defined by various criteria, including a decline in PSA to <0.5 or 1 ng/mL, “nonrising” PSA values, and a negative biopsy of the prostate 2 years after completion of treatment. The current standard definition of biochemical failure (the Phoenix definition) is a rise in PSA by ≥ 2 ng/mL higher than the lowest PSA achieved. The date of failure is “at call” and not backdated.

Radiation dose is critical to the eradication of prostate cancer. In a representative study, a PSA nadir of <1.0 ng/mL was achieved in 90% of patients receiving 75.6 or 81.0 Gy versus 76% and 56% of those receiving 70.2 and 64.8 Gy, respectively. Positive biopsy rates at 2.5 years were 4% for those treated with 81 Gy versus 27% and 36% for those receiving 75.6 and 70.2 Gy, respectively.

Overall, radiation therapy is associated with a higher frequency of bowel complications (mainly diarrhea and proctitis) than surgery. The frequency relates directly to the volume of the anterior rectal wall receiving full-dose treatment. In one series, grade 3 rectal or urinary toxicities were seen in 2.1% of patients who received a median dose of 75.6 Gy, whereas grade 3 urethral strictures requiring dilatation developed in 1% of cases, all of whom had undergone a transurethral resection of the prostate (TURP). Pooled data show that the frequency of grade 3 and 4 toxicities is 6.9% and 3.5%, respectively, for patients who received >70 Gy. The frequency of erectile dysfunction is related to the age of the patient, the quality of erections pretreatment, the dose administered, and the time of assessment. Postradiation erectile dysfunction is related to a disruption of the vascular supply and not the nerve fibers.

Neoadjuvant hormone therapy before radiation therapy has the aim of decreasing the size of the prostate and, consequently, reducing the exposure of normal tissues to full-dose radiation, increasing local control rates, and decreasing the rate of systemic failure.

584 Short-term hormone therapy can reduce toxicities and improve local control rates, but long-term treatment (2–3 years) is needed to prolong the time to PSA failure and lower the risk of metastatic disease in men with high-risk cancers. The impact on survival has been less clear.

BRACHYTHERAPY Brachytherapy is the direct implantation of radioactive sources (seeds) into the prostate. It is based on the principle that the deposition of radiation energy in tissues decreases as a function of the square of the distance from the source (**Chap. 103e**). The goal is to deliver intensive irradiation to the prostate, minimizing the exposure of the surrounding tissues. The current standard technique achieves a more homogeneous dose distribution by placing seeds according to a customized template based on imaging assessment of the cancer and computer-optimized dosimetry. The implantation is performed transperineally as an outpatient procedure with real-time imaging.

Improvements in brachytherapy techniques have resulted in fewer complications and a marked reduction in local failure rates. In a series of 197 patients followed for a median of 3 years, 5-year actuarial PSA relapse-free survival for patients with pretherapy PSA levels of 0–4, 4–10, and >10 ng/mL were 98%, 90%, and 89%, respectively. In a separate report of 201 patients who underwent posttreatment biopsies, 80% were negative, 17% were indeterminate, and 3% were positive. The results did not change with longer follow-up. Nevertheless, many physicians feel that implantation is best reserved for patients with good or intermediate prognostic features.

Brachytherapy is well tolerated, although most patients experience urinary frequency and urgency that can persist for several months. Incontinence has been seen in 2–4% of cases. Higher complication rates are observed in patients who have undergone a prior TURP, whereas those with obstructive symptoms at baseline are at a higher risk for retention and persistent voiding symptoms. Proctitis has been reported in <2% of patients.

Active Surveillance Although prostate cancer is the most common form of cancer affecting men in the United States, patients are being diagnosed earlier and more frequently present with early-stage disease. Active surveillance, described previously as *watchful waiting* or *deferred therapy*, is the policy of monitoring the illness at fixed intervals with DREs, PSA measurements, and repeat prostate biopsies as indicated until histopathologic or serologic changes correlative of progression warrant treatment with curative intent. It evolved from studies that evaluated predominantly elderly men with well-differentiated tumors who demonstrated no clinically significant progression for protracted periods, recognition of the contrast between incidence and disease-specific mortality, the high prevalence of autopsy cancers, and an effort to reduce overtreatment. A recent screening study estimated that between 50–100 men with low-risk disease would need to be treated to prevent one prostate cancer-specific death.

Arguing against active surveillance are the results of a Swedish randomized trial of radical prostatectomy versus active surveillance. With a median follow-up of 6.2 years, men treated by radical surgery had a lower risk of prostate cancer death relative to active surveillance patients (4.6% vs 8.9%) and a lower risk of metastatic progression (hazard ratio 0.63). Case selection is critical, and determining clinical parameters predictive of cancer aggressiveness that can be used to reliably select men most likely to benefit from active surveillance is an area of intense study. In one prostatectomy series, it was estimated that 10–15% of those treated had “insignificant” disease. One set of criteria includes men with clinical T1c tumors that are biopsy Gleason grade 6 or less involving three or fewer cores, each of them having less than 50% involvement by tumor and a PSA density of less than 0.15.

Concerns about active surveillance include the limited ability to predict pathologic findings by needle biopsy even when multiple cores are obtained, the recognized multifocality of the disease, and the possibility of a missed opportunity to cure the disease. Nomograms to help predict which patients can safely be managed

by active surveillance continue to be refined, and as their predictive accuracy improves, it can be anticipated that more patients will be candidates.

RISING PSA AFTER DEFINITIVE LOCAL THERAPY

This term is applied to a group of patients in whom the sole manifestation of disease is a rising PSA after surgery and/or radiation therapy. By definition, there is no evidence of disease on an imaging study. For these patients, the central issue is whether the rise in PSA results from persistent disease in the primary site, systemic disease, or both. In theory, disease in the primary site may still be curable by additional local treatment.

The decision to recommend radiation therapy after prostatectomy is guided by the pathologic findings at surgery, because imaging studies such as CT and bone scan are typically uninformative. Some recommend a choline-11 positron emission tomography (PET) scan, but availability in the United States is limited. Others recommend that a biopsy of the urethrovesical anastomosis be obtained before considering radiation, whereas others treat empirically based on risk. Factors that predict for response to salvage radiation therapy are a positive surgical margin, lower Gleason score in the radical prostatectomy specimen, long interval from surgery to PSA failure, slow PSA doubling time, absence of disease in the lymph nodes, and a low (<0.5–1 ng/mL) PSA value at the time of radiation treatment. Radiation therapy is generally not recommended if the PSA was persistently elevated after surgery, which usually indicates that the disease has spread outside of the area of the prostate bed and is unlikely to be controlled with radiation therapy. As is the case for other disease states, nomograms to predict the likelihood of success are available.

For patients with a rising PSA after radiation therapy, salvage local therapy can be considered if the disease was “curable” at the time of diagnosis, if persistent disease has been documented by a biopsy of the prostate, and if no metastatic disease is seen on imaging studies. Unfortunately, case selection is poorly defined in most series, and morbidities are significant. Options include salvage radical prostatectomy, salvage cryotherapy, salvage radiation therapy, and salvage irreversible electroporation.

The rise in PSA after surgery or radiation therapy may indicate subclinical or micrometastatic disease with or without local recurrence. In these cases, the need for treatment depends, in part, on the estimated probability that the patient will develop clinically detectable metastatic disease on a scan and in what time frame. That immediate therapy is not always required was shown in a series where patients who developed a biochemical recurrence after radical prostatectomy received no systemic therapy until metastatic disease was documented. Overall, the median time to metastatic progression by imaging was 8 years, and 63% of the patients with rising PSA values remained free of metastases at 5 years. Factors associated with progression included the Gleason score of the radical prostatectomy specimen, time to recurrence, and PSA doubling time. For those with Gleason grade ≥ 8 , the probability of metastatic progression was 37%, 51%, and 71% at 3, 5, and 7 years, respectively. If the time to recurrence was <2 years and PSA doubling time was long (>10 months), the proportions with metastatic disease at the same time intervals were 23%, 32%, and 53%, versus 47%, 69%, and 79% if the doubling time was short (<10 months). PSA doubling times are also prognostic for survival. In one series, all patients who succumbed to disease had PSA doubling times of 3 months or less.

Most physicians advise treatment if the PSA doubling time is 12 months or less. A difficulty with predicting the risk of metastatic spread, symptoms, or death from disease in the rising PSA state is that most patients receive some form of therapy before the development of metastases. Nevertheless, predictive models continue to be refined.

METASTATIC DISEASE: NONCASTRATE

The state of *noncastrate metastatic prostate cancer* includes men with metastases visible on an imaging study and noncastrate levels of testosterone (>150 ng/dL). The patient may be newly diagnosed

or have a recurrence after treatment for localized disease. Symptoms of metastatic disease include pain from osseous spread, although many patients are asymptomatic despite extensive spread. Less common are symptoms related to marrow compromise (myelophthisis), spinal cord compression, or a coagulopathy.

Standard treatment is to deplete/lower androgens by medical or surgical means and/or to block androgen binding to the AR with antiandrogens. More than 90% of male hormones originate in the testes; <10% are synthesized in the adrenal gland. Surgical orchiectomy is the “gold standard” but is rarely used due to the availability of effective medical therapies and the more widespread use of hormones on an intermittent basis by which patients are treated for defined periods of time, following which the treatments are intentionally discontinued (discussed further below) (Fig. 115-3).

Testosterone-Lowering Agents Medical therapies that lower testosterone levels include the gonadotropin-releasing hormone (GnRH) agonists/antagonists, 17,20-lyase inhibitors, CYP17 inhibitors, estrogens, and progestational agents. Of these, GnRH analogues such as leuprolide acetate and goserelin acetate initially produce a rise in luteinizing hormone and follicle-stimulating hormone, followed by a downregulation of receptors in the pituitary gland, which effects a chemical castration. They were approved on the basis of randomized comparisons showing an improved safety profile (specifically, reduced cardiovascular toxicities) relative to diethylstilbestrol (DES), with equivalent potency. The initial rise in testosterone may result in a clinical flare of the disease. As such, these agents are relatively contraindicated in men with significant obstructive symptoms, cancer-related pain, or spinal cord compromise. GnRH antagonists such as degarelix achieve castrate levels of testosterone within 48 h without the initial rise in serum testosterone and do not cause a flare in the disease. Estrogens such as DES are rarely used due to the risk of vascular complications such as fluid retention, phlebitis, embolic events, and stroke. Progestational agents alone are less efficacious.

Agents that lower testosterone are associated with an androgen-depletion syndrome that includes hot flashes, weakness, fatigue, loss of libido, impotence, sarcopenia, anemia, change in personality, and depression. Changes in lipids, obesity, and insulin resistance, along with an increased risk of diabetes and cardiovascular disease, can also occur, mimicking the metabolic syndrome. A decrease in bone density may also result that worsens over time and results in an increased risk of clinical fractures. This is a particular concern, often underappreciated, for men with preexisting osteopenia secondary to hypogonadism or glucocorticoid or alcohol use. Baseline fracture risk can be assessed using the Fracture Risk Assessment Scale (FRAX), and to minimize fracture risk, patients are advised to take calcium and vitamin D supplementation, along with a bisphosphonate or the RANK ligand inhibitor, denosumab.

Antiandrogens First-generation nonsteroidal antiandrogens such as flutamide, bicalutamide, and nilutamide block ligand binding to the AR and were initially approved to block the disease flare that may occur with the rise in serum testosterone associated with GnRH agonist therapy. When antiandrogens are given alone, testosterone levels typically increase above baseline, but relative to testosterone-lowering therapies, they cause fewer hot flashes, less of an effect on libido, less muscle wasting, fewer personality changes, and less bone loss. Gynecomastia remains a significant problem but can be alleviated in part by tamoxifen.

Most reported randomized trials suggest that the cancer-specific outcomes are inferior when antiandrogens are used alone. Bicalutamide, even at 150 mg (three times the recommended dose), was associated with a shorter time to progression and inferior survival compared to surgical castration for patients with established metastatic disease. Nevertheless, some men may accept the trade-off of a potentially inferior cancer outcome for an improved quality of life.

Combined androgen blockade, the administration of an antiandrogen plus a GnRH analogue or surgical orchiectomy, and triple androgen blockade, which includes the addition of a 5ARI, have not

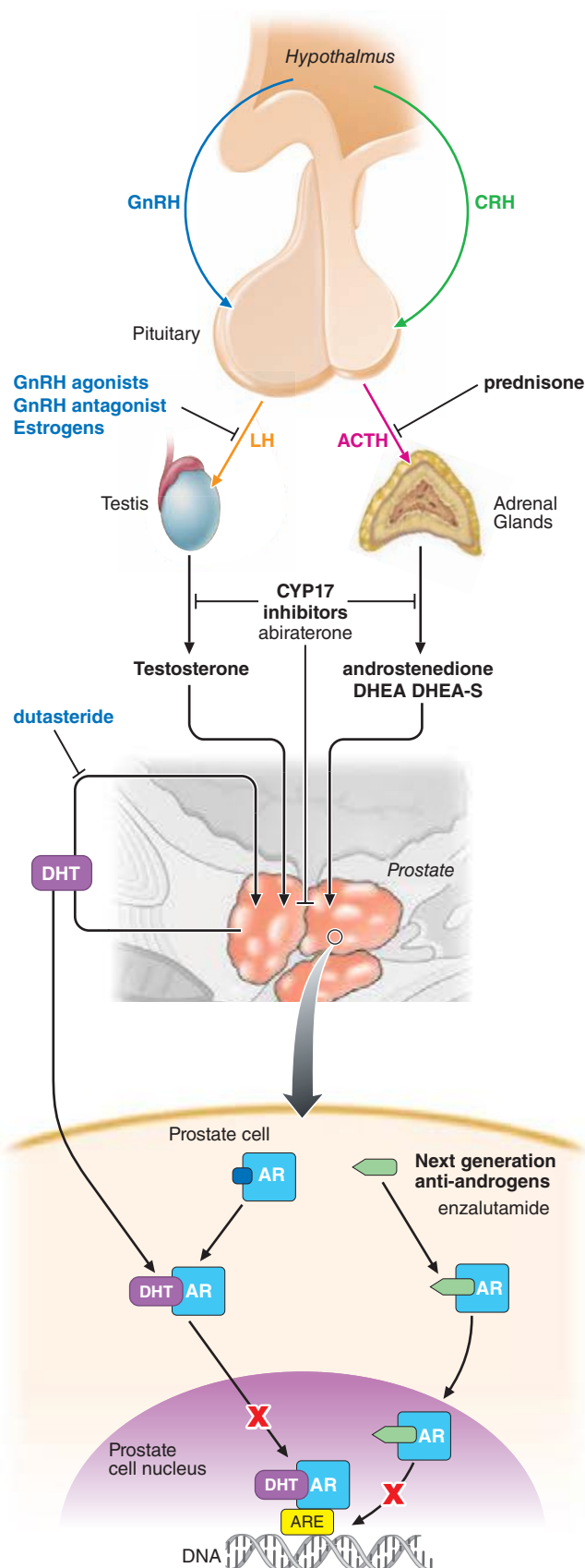


FIGURE 115-3 Sites of action of different hormone therapies.

ACTH, adrenocorticotropic hormone; AR, androgen receptor; ARE, androgen-response element; CRH, corticotropin-releasing hormone; DHEA, dehydroepiandrosterone; DHEA-S, dehydroepiandrosterone sulphate; DHT, dihydrotestosterone; GnRH, gonadotropin-releasing hormone; LH, luteinizing hormone.

586 been shown to be superior to androgen depletion monotherapies and are no longer recommended. In practice, most patients who are treated with a GnRH agonist receive an antiandrogen for the first 2–4 weeks of treatment to protect against the flare.

Intermittent Androgen Deprivation Therapy (IADT) The use of hormones in an “on-and-off” approach was initially proposed as a way to prevent the selection of cells that are resistant to androgen depletion and to reduce side effects. The hypothesis is that by allowing endogenous testosterone levels to rise, the cells that survive androgen depletion will induce a normal differentiation pathway. It is postulated that by allowing the surviving cells to proliferate in the presence of androgen, sensitivity to subsequent androgen depletion will be retained and the chance of developing a castration-resistant state will be reduced. Applied in the clinic, androgen depletion is continued for 2–6 months beyond the point of maximal response. Once treatment is stopped, endogenous testosterone levels increase, and the symptoms associated with hormone treatment abate. PSA levels also begin to rise, and at some level, treatment is restarted. With this approach, multiple cycles of regression and proliferation have been documented in individual patients. It is unknown whether the intermittent approach increases, decreases, or does not change the overall duration of sensitivity to androgen depletion. The approach is safe, but long-term data are needed to assess the course in men with low PSA levels. A randomized trial showed similar survival time between patients treated with intermittent versus continuous treatment, with a slightly higher risk of prostate cancer-specific mortality in the intermittent group, and higher cardiovascular mortality in patients on continuous therapy. The intermittent therapy was better tolerated.

Outcomes of Androgen Depletion The anti-prostate cancer effects of the various androgen depletion/blockade strategies are similar, and the outcomes predictable: an initial response, then a period of stability in which tumor cells are dormant and nonproliferative, followed after a variable period of time by a rise in PSA and tumor regrowth as a castration-resistant lesion that for most men is invariably lethal. Androgen depletion is not curative because cells that survive castration are present when the disease is first diagnosed. Considered by disease manifestation, PSA levels return to normal in 60–70% of cases, and measurable lesions regress in about 50%; improvements in bone scan occur in 25% of cases, but the majority of cases remain stable. The duration of response and survival is inversely proportional to disease extent at the time androgen depletion is first started, whereas the degree of PSA decline at 6 months has been shown to be prognostic. In a large-scale trial, PSA nadir proved prognostic.

An active question is whether hormones should be given in the adjuvant setting after surgery or radiation treatment of the primary tumor or whether to wait until PSA recurrence, metastatic disease, or symptoms are documented. Trials in support of early therapy have often been underpowered relative to the reported benefit or have been criticized on methodologic grounds. One trial showing a survival benefit for patients treated with radiation therapy and 3 years of androgen depletion, relative to radiation alone, was criticized for the poor outcomes of the control group. Another showing a survival benefit for patients with positive lymph nodes who were randomized to immediate medical or surgical castration compared to observation ($p = .02$) was criticized because the confidence intervals around the 5- and 8-year survival distributions for the two groups overlapped. A large randomized study comparing early to late hormone treatment (orchiectomy or GnRH analogue) in patients with locally advanced or asymptomatic metastatic disease showed that patients treated early were less likely to progress from M0 to M1 disease, to develop pain, and to die of prostate cancer. This trial was criticized because therapy was delayed “too long” in the late-treatment group. Noteworthy is that the American Society of Clinical Oncology Guidelines recommend deferring treatment until the disease has recurred and the prognosis has been reassessed. These guidelines do not support immediate therapy.

METASTATIC DISEASE: CASTRATE

Castration-resistant prostate cancer (CRPC) is defined as disease that progresses despite androgen suppression by medical or surgical therapies where the measured levels of testosterone are 50 ng/mL or lower. The rise in PSA indicates continued signaling through the AR signaling axis, the result of a series of oncogenic changes that include overexpression of androgen biosynthetic enzymes that can lead to increased intratumoral androgens, and overexpression of the receptor itself that enables signaling to occur even in the setting of low levels of androgen. The majority of CRPC cases are not “hormone-refractory,” and considering them as such can deny patients safe and effective treatment. CRPC can manifest in many ways. For some, it is a rise in PSA with no change in radiographs and no new symptoms. In others, it is a rising PSA and progression in bone with or without symptoms of disease. Still others will show soft tissue disease with or without osseous metastases, and others have visceral spread.

For the individual patient, it is first essential to ensure that a castrate status be documented. Patients receiving an antiandrogen alone, whose serum testosterone levels are elevated, should be treated first with a GnRH analogue or orchiectomy and observed for response. Patients on an antiandrogen in combination with a GnRH analogue should have the antiandrogen discontinued, because approximately 20% will respond to the selective discontinuation of the antiandrogen.

Chemotherapy and New Agents Through 2009, docetaxel was the only systemic therapy proven to prolong life. As a single agent, the drug produced PSA declines in 50% of patients, measurable disease regression in 25%, and improvement in both preexisting pain and prevention of future cancer-related pain. Since then, six agents with diverse mechanisms of action that target the tumor itself or other aspects of the metastatic process have been proven to prolong life and were FDA approved. The first was sipuleucel-T, the first biologic approach shown to prolong life in which antigen-presenting cells are activated *ex vivo*, pulsed with antigen, and reinfused. The second, cabazitaxel, a non-cross-resistant taxane, was shown to be superior to mitoxantrone in the post-docetaxel setting. This was followed by the CYP17 inhibitor abiraterone acetate, which lowers androgen levels in the tumor, adrenal glands, and testis, and the next-generation antiandrogen enzalutamide, which not only has a higher binding affinity to the AR relative to first-generation compounds, but uniquely inhibits nuclear location and DNA binding of the receptor complex. Both abiraterone acetate and enzalutamide were first approved for postchemotherapy treated patients on the basis of placebo-controlled phase III trials—a further indication that these tumors are not uniformly hormone-refractory. The indication for abiraterone acetate was later expanded to the prechemotherapy setting, based on a second trial using a co-primary endpoint of radiographic progression-free survival and overall survival. Similar results were seen with enzalutamide, for which an expanded indication is also anticipated. Alpharadin (radium-223 chloride), an alpha-emitting bone-seeking radioisotope, has been shown to prolong life in patients with symptoms related to osseous disease. The alpharadin result validated the bone microenvironment as a therapeutic target independent of direct effects on the tumor itself, as no declines in PSA were observed in the trial. Notable is that in addition to a survival benefit, the drug also reduced the development of significant skeletal events.

Other bone-targeted agents, such as the bisphosphonates and the RANK ligand inhibitor denosumab, protect against bone loss associated with androgen depletion and also reduce skeletal-related events by targeting bone osteoclasts. In one trial, denosumab was shown to be superior to zoledronic acid with respect to skeletal-related events, but had a slightly higher frequency of osteonecrosis of the jaw.

In clinical practice, most men seek to avoid chemotherapy and are first treated with a biologic agent and/or newer hormonal agent approved for this indication. It is crucial to the management of the individual patient to define therapeutic objectives before initiating treatment, as there are defined standards of care for different

disease manifestations. For example, sipuleucel-T is not indicated for patients with symptoms or visceral disease because the effects on the disease occur late. Similarly, alpharadin is not indicated for patients with disease that is predominantly in soft tissue or who have osseous disease that is not causing symptoms.

Pain Management Management of pain secondary to osseous metastatic disease is a critical part of therapy. Optimal palliation requires assessing whether the symptoms are from metastases that threaten or that are already affecting the spinal cord, the cauda equina, or the base of the skull, which are best treated with external-beam radiation, as are single sites of pain. Neurologic symptoms require emergency evaluation because loss of function may be permanent if not addressed quickly. Because the disease is often diffuse, palliation at one site is often followed by the emergence of symptoms in a separate site that had not received radiation. In these cases, bone-seeking radioisotopes such as alpharadin or the beta emitter ¹⁵³Sm-EDTMP (Quadramet) can be considered in addition to abiraterone acetate, docetaxel, and mitoxantrone, each of which is formally approved for the palliation of pain due to prostate cancer metastases.

BENIGN DISEASE

BENIGN PROSTATIC HYPERTROPHY

BPH is a pathologic process that contributes to the development of lower urinary tract symptoms in men. Such symptoms, arising from lower urinary tract dysfunction, are further subdivided into obstructive symptoms (urinary hesitancy, straining, weak stream, terminal dribbling, prolonged voiding, incomplete emptying) and irritative symptoms (urinary frequency, urgency, nocturia, urge incontinence, small voided volumes). Lower urinary tract symptoms and other sequelae of BPH are not just due to a mass effect, but are also likely due to a combination of the prostatic enlargement and age-related detrusor dysfunction.

TREATMENT BENIGN PROSTATIC HYPERTROPHY

The symptoms are generally measured using a validated, reproducible index that is designed to determine disease severity and response to therapy—the AUA's Symptom Index (AUASI), also adopted as the International Prostate Symptom Score (IPSS) (Table 115-2). Serial AUASI is particularly useful in following patients as they are treated with various forms of therapy. Asymptomatic patients do not require treatment regardless of the size of the gland, whereas patients with an inability to urinate, gross hematuria, recurrent

infection, or bladder stones may require surgery. In patients with symptoms, uroflowmetry can identify those with normal flow rates who are unlikely to benefit from treatment, and bladder ultrasound can identify those with high postvoid residuals who may need intervention. Pressure-flow (urodynamic) studies detect primary bladder dysfunction. Cystoscopy is recommended if hematuria is documented and to assess the urinary outflow tract before surgery. Imaging of the upper tracts is advised for patients with hematuria, a history of calculi, or prior urinary tract problems.

Symptomatic relief is the most common reason men seek treatment for BPH, and therefore the goal of therapy for BPH is usually relief of these symptoms. Alpha-adrenergic receptor antagonists are thought to treat the dynamic aspect of BPH by reducing sympathetic tone of the bladder outlet, thereby decreasing resistance and improving urinary flow. 5ARIs are thought to treat the static aspect of BPH by reducing prostate volume and having a similar, albeit delayed effect. They have also proven to be beneficial in the prevention of BPH progression, as measured by prostate volume, the risk of developing acute urinary retention, and the risk of having BPH-related surgery. The use of an alpha-adrenergic receptor antagonist and a 5ARI as combination therapy seeks to provide symptomatic relief while preventing progression of BPH.

Another class of medications that has shown improvement in lower urinary tract symptoms secondary to BPH is PDE5 inhibitors, used currently in the treatment of erectile dysfunction. All three of the PDE5 inhibitors available in the United States, sildenafil, vardenafil, and tadalafil, appear to be effective in the treatment of symptoms secondary to BPH. The use of PDE5 inhibitors is not without controversy, however, given the fact that short-acting phosphodiesterase inhibitors such as sildenafil need to be dosed separately from alpha blockers such as tamsulosin because of potential hypotensive effects. Newer classes of pharmacologic agents have been used to treat symptoms secondary to BPH. Symptoms due to BPH often coexist with symptoms due to overactive bladder, and the most common pharmacologic agents for the treatment of overactive bladder symptoms are anticholinergics. This has led to multiple studies evaluating the efficacy of anticholinergics for the treatment of lower urinary tract symptoms secondary to BPH. Surgical therapy is now considered second-line therapy and is usually reserved for patients after a trial of medical therapy. The goal of surgical therapy is to reduce the size of the prostate, effectively reducing resistance to urine flow.

Surgical approaches include TURP, transurethral incision, or removal of the gland via a retropubic, suprapubic, or perineal approach. Also used are transurethral ultrasound-guided laser-induced prostatectomy (TULIP), stents, and hyperthermia.

TABLE 115-2 AUA SYMPTOM INDEX

Questions to Be Answered	AUA Symptom Score (Circle 1 Number on Each Line)					
	Not at All	Less Than 1 Time in 5	Less Than Half the Time	About Half the Time	More Than Half the Time	Almost Always
Over the past month, how often have you had a sensation of not emptying your bladder completely after you finished urinating?	0	1	2	3	4	5
Over the past month, how often have you had to urinate again less than 2 h after you finished urinating?	0	1	2	3	4	5
Over the past month, how often have you found you stopped and started again several times when you urinated?	0	1	2	3	4	5
Over the past month, how often have you found it difficult to postpone urination?	0	1	2	3	4	5
Over the past month, how often have you had a weak urinary stream?	0	1	2	3	4	5
Over the past month, how often have you had to push or strain to begin urination?	0	1	2	3	4	5
Over the past month, how many times did you most typically get up to urinate from the time you went to bed at night until the time you got up in the morning?	(None)	(1 time)	(2 times)	(3 times)	(4 times)	(5 times)
Sum of 7 circled numbers (AUA Symptom Score): _____						

Abbreviation: AUA, American Urological Association.

Source: MJ Barry et al: J Urol 148:1549, 1992. Used with permission.

Primary germ cell tumors (GCTs) of the testis arising by the malignant transformation of primordial germ cells constitute 95% of all testicular neoplasms. Infrequently, GCTs arise from an extragonadal site, including the mediastinum, retroperitoneum, and, very rarely, the pineal gland. This disease is notable for the young age of the afflicted patients, the totipotent capacity for differentiation of the tumor cells, and its curability; approximately 95% of newly diagnosed patients are cured. Experience in the management of GCTs leads to improved outcome.

INCIDENCE AND EPIDEMIOLOGY



The incidence of testicular GCT is now approximately 8000 cases annually in the United States, resulting in nearly 400 deaths. The tumor occurs most frequently in men between the ages of 20 and 40 years. A testicular mass in a male ≥ 50 years should be regarded as a lymphoma until proved otherwise. GCT is at least four to five times more common in white than in African-American males, and a higher incidence has been observed in Scandinavia and New Zealand than in the United States.

ETIOLOGY AND GENETICS

Cryptorchidism is associated with a several-fold higher risk of GCT. Abdominal cryptorchid testes are at a higher risk than inguinal cryptorchid testes. Orchiopexy should be performed before puberty, if possible. Early orchiopexy reduces the risk of GCT and improves the ability to save the testis. An abdominal cryptorchid testis that cannot be brought into the scrotum should be removed. Approximately 2% of men with GCTs of one testis will develop a primary tumor in the other testis. Testicular feminization syndromes and family history increase the risk of testicular GCT, and Klinefelter's syndrome is associated with mediastinal GCT.

An isochromosome of the short arm of chromosome 12 [i(12p)] is pathognomonic for GCT. Excess 12p copy number, either in the form of i(12p) or as increased 12p on aberrantly banded marker chromosomes, occurs in nearly all GCTs, but the gene(s) on 12p involved in the pathogenesis are not yet defined.

CLINICAL PRESENTATION

A painless testicular mass is pathognomonic for a testicular malignancy. More commonly, patients present with testicular discomfort or swelling suggestive of epididymitis and/or orchitis. In this circumstance, a trial of antibiotics is reasonable. However, if symptoms persist or a residual abnormality remains, then testicular ultrasound examination is indicated.

Ultrasound of the testis is indicated whenever a testicular malignancy is considered and for persistent or painful testicular swelling. If a testicular mass is detected, a radical inguinal orchiectomy should be performed. Because the testis develops from the gonadal ridge, its blood supply and lymphatic drainage originate in the abdomen and descend with the testis into the scrotum. An inguinal approach is taken to avoid breaching anatomic barriers and permitting additional pathways of spread.

Back pain from retroperitoneal metastases is common and must be distinguished from musculoskeletal pain. Dyspnea from pulmonary metastases occurs infrequently. Patients with increased serum levels of human chorionic gonadotropin (hCG) may present with gynecomastia. A delay in diagnosis is associated with a more advanced stage and possibly worse survival.

The staging evaluation for GCT includes a determination of serum levels of a fetoprotein (AFP), hCG, and lactate dehydrogenase (LDH). After orchiectomy, a computed tomography (CT) scan of the chest, abdomen, and pelvis is generally performed. Stage I disease is limited to the testis, epididymis, or spermatic cord. Stage II disease is limited

to retroperitoneal (regional) lymph nodes. Stage III disease is disease outside the retroperitoneum, involving supradiaphragmatic nodal sites or viscera. The staging may be "clinical"—defined solely by physical examination, blood marker evaluation, and radiographs—or "pathologic"—defined by an operative procedure.

The regional draining lymph nodes for the testis are in the retroperitoneum, and the vascular supply originates from the great vessels (for the right testis) or the renal vessels (for the left testis). As a result, the lymph nodes that are involved first by a right testicular tumor are the interaortocaval lymph nodes just below the renal vessels. For a left testicular tumor, the first involved lymph nodes are lateral to the aorta (para-aortic) and below the left renal vessels. In both cases, further retroperitoneal nodal spread is inferior, contralateral, and, less commonly, above the renal hilum. Lymphatic involvement can extend cephalad to the retrocrural, posterior mediastinal, and supraclavicular lymph nodes. Treatment is determined by tumor histology (seminoma versus nonseminoma) and clinical stage (Fig. 116-1).

PATHOLOGY

GCTs are divided into nonseminoma and seminoma subtypes. Nonseminomatous GCTs are most frequent in the third decade of life and can display the full spectrum of embryonic and adult cellular differentiation. This entity comprises four histologies: embryonal carcinoma, teratoma, choriocarcinoma, and endodermal sinus (yolk sac) tumor. Choriocarcinoma, consisting of both cytotrophoblasts and syncytiotrophoblasts, represents malignant trophoblastic differentiation and is invariably associated with secretion of hCG. Endodermal sinus tumor is the malignant counterpart of the fetal yolk sac and is associated with secretion of AFP. Pure embryonal carcinoma may secrete AFP or hCG, or both; this pattern is biochemical evidence of differentiation. Teratoma is composed of somatic cell types derived from two or more germ layers (ectoderm, mesoderm, or endoderm). Each of these histologies may be present alone or in combination with others. Nonseminomatous GCTs tend to metastasize early to sites such as the retroperitoneal lymph nodes and lung parenchyma. Sixty percent of patients present with disease limited to the testis (stage I), 20% with retroperitoneal metastases (stage II), and 20% with more extensive supradiaphragmatic nodal or visceral metastases (stage III).

Seminoma represents approximately 50% of all GCTs, has a median age in the fourth decade, and generally follows a more indolent clinical course. Eighty percent of patients present with stage I disease, approximately 10% with stage II disease, and 10% with stage III disease; lung or other visceral metastases are rare. When a tumor contains both seminoma and nonseminoma components, patient management is directed by the more aggressive nonseminoma component.

TUMOR MARKERS

Careful monitoring of the serum tumor markers AFP and hCG is essential in the management of patients with GCT, because these markers are important for diagnosis, as prognostic indicators, in monitoring treatment response, and in the early detection of relapse. Approximately 70% of patients presenting with disseminated nonseminomatous GCT have increased serum concentrations of AFP and/or hCG. Although hCG concentrations may be increased in patients with either nonseminoma or seminoma histology, the AFP concentration is increased only in patients with nonseminoma. The presence of an increased AFP level in a patient whose tumor shows only seminoma indicates that an occult nonseminomatous component exists, and the patient should be treated for nonseminomatous GCT. LDH levels are less specific than AFP or hCG but are increased in 50–60% patients with metastatic nonseminoma and in up to 80% of patients with advanced seminoma.

AFP, hCG, and LDH levels should be determined before and after orchiectomy. Increased serum AFP and hCG concentrations decay according to first-order kinetics; the half-life is 24–36 h for hCG and 5–7 days for AFP. AFP and hCG should be assayed serially during and after treatment. The reappearance of hCG and/or AFP or the failure of these markers to decline according to the predicted half-life is an indicator of persistent or recurrent tumor.

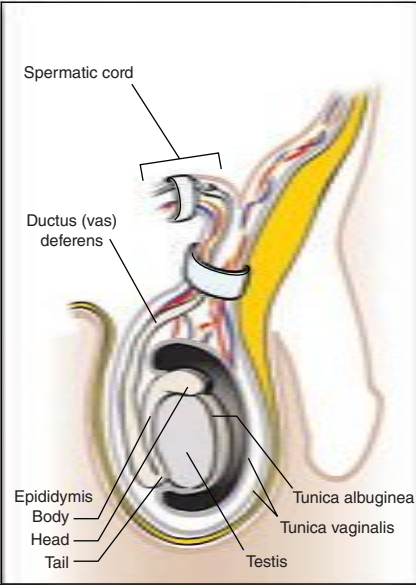
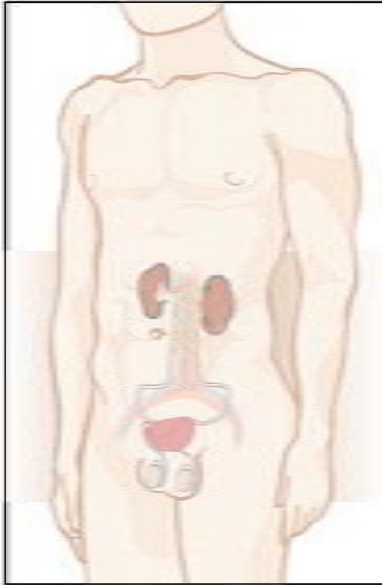
				Treatment Option	
Stage	Extent of Disease	Stage	Extent of Disease	Seminoma	Nonseminoma
	<p>pT1</p> <p>Tumor limited to the testis and epididymis without vascular/lymphatic invasion; tumor may invade into the tunica albuginea but not the tunica vaginalis</p>		<p>IA</p> <p>Testis only, no vascular/lymphatic invasion (T1)</p>	Observation Chemotherapy or RT	Observation
	<p>pT2</p> <p>Tumor limited to the testis and epididymis with vascular/lymphatic invasion; tumor extending through the tunica albuginea with involvement of the tunica vaginalis</p>		<p>IB</p> <p>Testis only, with vascular/lymphatic invasion (T2), or extension through tunica albuginea (T2), or involvement of spermatic cord (T3) or scrotum (T4)</p>	Observation Chemotherapy or RT	RPLND or Chemotherapy
	<p>pT3</p> <p>Tumor invades the spermatic cord with or without vascular/lymphatic invasion</p>		<p>IIA</p> <p>Retroperitoneal Nodes < 2 cm</p>	RT	RPLND +/- adjuvant Chemotherapy or Chemotherapy, often followed by RPLND
	<p>pT4</p> <p>Tumor invades the scrotum with or without vascular/lymphatic invasion</p>		<p>IIB</p> <p>Retroperitoneal Nodes 2-5 cm</p>	RT or Chemotherapy	Chemotherapy, often followed by RPLND
		<p>IIC</p> <p>Retroperitoneal Nodes > 5 cm</p>		Chemotherapy	Chemotherapy, often followed by RPLND
		<p>Distant Metastases</p>			
		<p>III</p> <p>Common sites include distant (or extra-abdominal) lymph nodes, lung, liver, bone, and brain</p>	Chemotherapy	Chemotherapy, often followed by surgery (biopsy or resection)	

FIGURE 116-1 Germ cell tumor staging and treatment. RPLND, retroperitoneal lymph node dissection; RT, radiotherapy.

STAGE I NONSEMINOMA

Patients with radiographs and physical examination showing no evidence of disease and serum AFP and hCG concentrations that are either normal or declining to normal according to the known half-life have clinical stage I disease. Approximately 20–50% of such patients will have retroperitoneal lymph node metastases (pathologic stage II) but will still be cured in over 95% of cases. Depending on risk of relapse, which is determined by the pathology (see below), surveillance, a nerve-sparing retroperitoneal lymph node dissection (RPLND), or adjuvant chemotherapy (one to two cycles of bleomycin, etoposide, and cisplatin [BEP]) may be appropriate choices depending on *the availability of surgical expertise and patient and physician preference*. If the primary tumor shows no evidence for lymphatic or vascular invasion and is limited to the testis (T1, clinical stage IA), then the risk of relapse is only 10–20%. Because over 80% of patients with clinical stage IA nonseminoma are cured with orchiectomy alone and there is no survival advantage to RPLND (or adjuvant chemotherapy), surveillance is the preferred treatment option. This avoids overtreatment with the potential for both acute and long-term toxicities (see below). Surveillance requires patients to be carefully followed with periodic chest radiography, physical examination, CT scan of the abdomen, and serum tumor marker determinations. The median time to relapse is approximately 7 months, and late relapses (>2 years) are rare. Noncompliant patients can be considered for RPLND or adjuvant BEP.

If lymphatic or vascular invasion is present or the tumor extends through the tunica, spermatic cord, or scrotum (T2 through T4, clinical stage IB), then the risk of relapse is approximately 50%, and RPLND and adjuvant chemotherapy can be considered. Relapse rates are reduced to 3–5% after one to two cycles of adjuvant BEP. All three approaches (surveillance, RPLND, and adjuvant BEP) should cure >95% of patients with clinical stage IB disease.

RPLND is the standard operation for removal of the regional lymph nodes of the testis (retroperitoneal nodes). The operation removes the lymph nodes draining the primary site and the nodal groups adjacent to the primary landing zone. The standard (modified bilateral) RPLND removes all node-bearing tissue down to the bifurcation of the great vessels, including the ipsilateral iliac nodes. The major long-term effect of this operation is retrograde ejaculation with resultant infertility. Nerve-sparing RPLND can preserve anterograde ejaculation in ~90% of patients. Patients with pathologic stage I disease are observed, and only the <10% who relapse require additional therapy. If nodes are found to be involved at RPLND, then a decision regarding adjuvant chemotherapy is made on the basis of the extent of retroperitoneal disease (see “Stage II Nonseminoma” below). Hence, because less than 20% of patients require chemotherapy, of the three approaches, RPLND results in the lowest number of patients at risk for the late toxicities of chemotherapy.

STAGE II NONSEMINOMA

Patients with limited, ipsilateral retroperitoneal adenopathy ≤ 2 cm in largest diameter and normal levels of AFP and hCG can be treated with either a modified bilateral nerve-sparing RPLND or chemotherapy. The local recurrence rate after a properly performed RPLND is very low. Depending on the extent of disease, the postoperative management options include either surveillance or two cycles of adjuvant chemotherapy. Surveillance is the preferred approach for patients with resected “low-volume” metastases (tumor nodes ≤ 2 cm in diameter and <6 nodes involved) because the probability of relapse is one-third or less. For those who relapse, risk-directed chemotherapy is indicated (see section on advanced GCT below). Because relapse occurs in $\geq 50\%$ of patients with “high-volume” metastases (>6 nodes involved, or any involved node >2 cm in largest diameter, or extranodal tumor extension), two cycles of adjuvant chemotherapy should be considered, as it results in a cure in $\geq 98\%$ of patients. Regimens consisting of etoposide plus

cisplatin (EP) with or without bleomycin every 3 weeks are effective and well tolerated.

Increased levels of either AFP or hCG imply metastatic disease outside the retroperitoneum; full-dose (not adjuvant) chemotherapy is used in this setting. Primary management with chemotherapy is also favored for patients with larger (>2 cm) or bilateral retroperitoneal nodes (see section on advanced GCT below).

STAGES I AND II SEMINOMA

Inguinal orchiectomy followed by immediate retroperitoneal radiation therapy or surveillance with treatment at relapse both result in cure in nearly 100% of patients with stage I seminoma. Historically, radiation was the mainstay of treatment, but the reported association between radiation and secondary malignancies and the absence of a survival advantage of radiation over surveillance has led many to favor surveillance for compliant patients. Approximately 15% of patients relapse, which is usually treated with chemotherapy. Long-term follow-up is essential, because approximately 30% of relapses occur after 2 years and 5% occur after 5 years. A single dose of carboplatin has also been investigated as an alternative to radiation therapy; the outcome was similar, but long-term safety data are lacking, and the retroperitoneum remained the most frequent site of relapse.

Generally, nonbulky retroperitoneal disease (stage IIA and small IIB) is treated with retroperitoneal radiation therapy. Approximately 90% of patients achieve relapse-free survival with retroperitoneal masses <3 cm in diameter. Due to higher relapse rates after radiation for bulkier disease, initial chemotherapy is preferred for all stage IIC and some stage IIB patients. Chemotherapy has been studied as an alternative to radiation for stage IIA and small stage IIB seminoma with lower recurrence rates compared with historical controls. These results, combined with studies demonstrating a threefold increase in the incidence of secondary malignancies and cardiovascular disease among patients who receive both radiation and chemotherapy (patients relapsing after radiation fall into this category), have led some experts to prefer chemotherapy for all stage II seminomas.

CHEMOTHERAPY FOR ADVANCED GCT

Regardless of histology, all patients with stage IIC and stage III and most with stage IIB GCT are treated with chemotherapy. Combination chemotherapy programs based on cisplatin at doses of 100 mg/m² plus etoposide at doses of 500 mg/m² per cycle cure 70–80% of such patients, with or without bleomycin, depending on risk stratification (see below). A complete response (the complete disappearance of all clinical evidence of tumor on physical examination and radiography plus normal serum levels of AFP and hCG for ≥ 1 month) occurs after chemotherapy alone in ~60% of patients, and another 10–20% become disease free with surgical resection of residual masses containing viable GCT. Lower doses of cisplatin result in inferior survival rates.

The toxicity of four cycles of the BEP is substantial. Nausea, vomiting, and hair loss occur in most patients, although nausea and vomiting have been markedly ameliorated by modern antiemetic regimens. Myelosuppression is frequent, and symptomatic bleomycin pulmonary toxicity occurs in ~5% of patients. Treatment-induced mortality due to neutropenia with septicemia or bleomycin-induced pulmonary failure occurs in 1–3% of patients. Dose reductions for myelosuppression are rarely indicated. Long-term permanent toxicities include nephrotoxicity (reduced glomerular filtration and persistent magnesium wasting), ototoxicity, peripheral neuropathy, and infertility. When bleomycin is administered by weekly bolus injection, Raynaud’s phenomenon appears in 5–10% of patients. Other evidence of small blood vessel damage, such as transient ischemic attacks and myocardial infarction, is seen less often.

RISK-DIRECTED CHEMOTHERAPY

Because not all patients are cured and treatment may cause significant toxicities, patients are stratified into “good-risk,” “intermediate-risk,” and “poor-risk” groups according to pretreatment clinical features established by the International Germ Cell Cancer Consensus Group (Table 116-1). For good-risk patients, the goal is to achieve

TABLE 116-1 INTERNATIONAL GERM CELL CANCER CONSENSUS GROUP RISK CLASSIFICATION FOR ADVANCED GERM CELL TUMORS

Risk	Nonseminoma	Seminoma
Good	Gonadal or retroperitoneal primary site Absent nonpulmonary visceral metastases AFP <1000 ng/mL β -hCG <5000 mIU/mL LDH <1.5 \times upper limit or normal (ULN)	Any primary site Absent nonpulmonary visceral metastases Any LDH, hCG
Intermediate	Gonadal or retroperitoneal primary site Absent nonpulmonary visceral metastases AFP 1000–10,000 ng/mL β -hCG 5000–50,000 mIU/mL LDH 1.5–10 \times ULN	Any primary site Presence of nonpulmonary visceral metastases Any LDH, hCG
Poor	Mediastinal primary site Presence of nonpulmonary visceral metastases AFP >10,000 ng/mL β -hCG >50,000 mIU/mL LDH >10 \times ULN	No patients classified as poor prognosis

Abbreviations: AFP, a fetoprotein; hCG, human chorionic gonadotropin; LDH, lactate dehydrogenase.

Source: From International Germ Cell Cancer Consensus Group.

maximum efficacy with minimal toxicity. For intermediate- and poor-risk patients, the goal is to identify more effective therapy with tolerable toxicity.

The marker cut offs are included in the TNM (primary tumor, regional nodes, metastasis) staging of GCT. Hence, TNM stage groupings are based on both anatomy (site and extent of disease) and biology (marker status and histology). Seminoma is either good- or intermediate-risk, based on the absence or presence of nonpulmonary visceral metastases. No poor-risk category exists for seminoma. Marker levels and primary site play no role in defining risk for seminoma. Nonseminomas have good-, intermediate-, and poor-risk categories based on the primary site of the tumor, the presence or absence of nonpulmonary visceral metastases, and marker levels.

For ~90% of patients with good-risk GCTs, four cycles of EP or three cycles of BEP produce durable complete responses, with minimal acute and chronic toxicity, and a low relapse rate. Pulmonary toxicity is absent when bleomycin is not used and is rare when therapy is limited to 9 weeks; myelosuppression with neutropenic fever is less frequent; and the treatment mortality rate is negligible. Approximately 75% of intermediate-risk patients and 50% of poor-risk patients achieve durable complete remission with four cycles of BEP, and no regimen has proved superior.

POSTCHEMOTHERAPY SURGERY Resection of residual metastases after the completion of chemotherapy is an integral part of therapy. If the initial histology is nonseminoma and the marker values have normalized, all sites of residual disease should be resected. In general, residual retroperitoneal disease requires a modified bilateral RPLND. Thoracotomy (unilateral or bilateral) and neck dissection are less frequently required to remove residual mediastinal, pulmonary parenchymal, or cervical nodal disease. Viable tumor (seminoma, embryonal carcinoma, yolk sac tumor, or choriocarcinoma) will be present in 15%, mature teratoma in 40%, and necrotic debris and fibrosis in 45% of resected specimens. The frequency of teratoma or viable disease is highest in residual mediastinal tumors. If necrotic debris or mature teratoma is present, no further chemotherapy is

necessary. If viable tumor is present but is completely excised, two additional cycles of chemotherapy are given.

If the initial histology is pure seminoma, mature teratoma is rarely present, and the most frequent finding is necrotic debris. For residual retroperitoneal disease, a complete RPLND is technically difficult due to extensive postchemotherapy fibrosis. Observation is recommended when no radiographic abnormality exists on CT scan. Positive findings on a positron emission tomography (PET) scan correlate with viable seminoma in residua and mandate surgical excision or biopsy.

SALVAGE CHEMOTHERAPY

Of patients with advanced GCT, 20–30% fail to achieve a durable complete response to first-line chemotherapy. A combination of vinblastine, ifosfamide, and cisplatin (VeIP) will cure approximately 25% of patients as a second-line therapy. Patients are more likely to achieve a durable complete response if they had a testicular primary tumor and relapsed from a prior complete remission to first-line cisplatin-containing chemotherapy. Substitution of paclitaxel for vinblastine (TIP) in this setting was associated with durable remission in nearly two-thirds of patients. In contrast, for patients with a primary mediastinal nonseminoma or who did not achieve a complete response with first-line chemotherapy, then VeIP standard-dose salvage therapy is rarely beneficial. Such patients are usually managed with high-dose chemotherapy and/or surgical resection.

Chemotherapy consisting of dose-intensive, high-dose carboplatin plus high-dose etoposide, with peripheral blood stem cell support, induces a complete response in 25–40% of patients who have progressed after ifosfamide-containing salvage chemotherapy. Approximately one-half of the complete responses will be durable. High-dose therapy is standard of care for this patient population and has been suggested as the treatment of choice for all patients with relapsed or refractory disease. Paclitaxel is active when incorporated into high-dose combination programs. Cure is still possible in some relapsed patients.

EXTRAGONADAL GCT

The prognosis and management of patients with extragonadal GCT depends on the tumor histology and site of origin. All patients with a diagnosis of extragonadal GCT should have a testicular ultrasound examination. Nearly all patients with retroperitoneal or mediastinal seminoma achieve a durable complete response to BEP or EP. The clinical features of patients with primary retroperitoneal nonseminoma GCT are similar to those of patients with a primary tumor of testis origin, and careful evaluation will find evidence of a primary testicular GCT in about two-thirds of cases. In contrast, a primary mediastinal nonseminomatous GCT is associated with a poor prognosis; one-third of patients are cured with standard therapy (four cycles of BEP). Patients with newly diagnosed mediastinal nonseminoma are considered to have poor-risk disease and should be considered for clinical trials testing regimens of possibly greater efficacy. In addition, mediastinal nonseminoma is associated with hematologic disorders, including acute myelogenous leukemia, myelodysplastic syndrome, and essential thrombocytosis unrelated to previous chemotherapy. These hematologic disorders are very refractory to treatment. Nonseminoma of any primary site may change into other malignant histologies such as embryonal rhabdomyosarcoma or adenocarcinoma. This is called *malignant transformation*. *i(12p)* has been identified in the transformed cell type, indicating GCT clonal origin.

A group of patients with poorly differentiated tumors of unknown histogenesis, midline in distribution, and not associated with secretion of AFP or hCG has been described; a few (10–20%) are cured by standard cisplatin-containing chemotherapy. An *i(12p)* is present in ~25% of such tumors (the fraction that are cisplatin-responsive), confirming their origin from primitive germ cells. This finding is also predictive of the response to cisplatin-based chemotherapy and resulting long-term survival. These tumors are heterogeneous; neuroepithelial tumors and lymphoma may also present in this fashion.

Infertility is an important consequence of the treatment of GCTs. Preexisting infertility or impaired fertility is often present. Azoospermia and/or oligospermia are present at diagnosis in at least 50% of patients with testicular GCTs. Ejaculatory dysfunction is associated with RPLND, and germ cell damage may result from cisplatin-containing chemotherapy. Nerve-sparing techniques to preserve the retroperitoneal sympathetic nerves have made retrograde ejaculation less likely in the subgroups of patients who are candidates for this operation. Spermatogenesis does recur in some patients after chemotherapy. However, because of the significant risk of impaired reproductive capacity, semen analysis and cryopreservation of sperm in a sperm bank should be recommended to all patients before treatment.

117 Gynecologic Malignancies

Michael V. Seiden

OVARIAN CANCER

INCIDENCE AND PATHOLOGY

Cancer arising in or near the ovary is actually a collection of diverse malignancies. This collection of malignancies, often referred to as “ovary cancer,” is the most lethal gynecologic malignancy in the United States and other countries that routinely screen women for cervical neoplasia. In 2014, it was estimated that there were 21,980 cases of ovarian cancer with 14,270 deaths in the United States. The ovary is a complex and dynamic organ and, between the ages of approximately 11 and 50 years, is responsible for follicle maturation associated with egg maturation, ovulation, and cyclical sex steroid hormone production. These complex and linked biologic functions are coordinated through a variety of cells within the ovary, each of which possesses neoplastic potential. By far the most common and most lethal of the ovarian neoplasms arise from the ovarian epithelium or, alternatively, the neighboring specialized epithelium of the fallopian tube, uterine corpus, or cervix. Epithelial tumors may be benign (50%), malignant (33%), or of borderline malignancy (16%). Age influences risk of malignancy; tumors in younger women are more likely benign. The most common of the ovarian epithelial malignancies are serous tumors (50%); tumors of mucinous (25%), endometrioid (15%), clear cell (5%), and transitional cell histology or Brenner tumor (1%) represent smaller proportions of epithelial ovarian tumors. In contrast, stromal tumors arise from the steroid hormone-producing cells and likewise have different phenotypes and clinical presentations largely dependent on the type and quantity of hormone production. Tumors arising in the germ cell are most similar in biology and behavior to testicular tumors in males (Chap. 116).

Tumors may also metastasize to the ovary from breast, colon, appendiceal, gastric, and pancreatic primaries. Bilateral ovarian masses from metastatic mucin-secreting gastrointestinal cancers are termed *Krukenberg tumors*.

OVARIAN CANCER OF EPITHELIAL ORIGIN

Epidemiology and Pathogenesis A female has approximately a 1 in 72 lifetime risk (1.6%) of developing ovarian cancer, with the majority of affected women developing epithelial tumors. Each of the histologic variants of epithelial tumors is distinct with unique molecular features. As a group of malignancies, epithelial tumors of the ovary have a peak incidence in women in their sixties, although age at presentation can range across the extremes of adult life, with cases being reported in women in their twenties to nineties. Each histologic subtype of ovarian cancer likely has its own associated risk factors. Serous cancer, the

most common type of epithelial ovarian cancer, is seen with increased frequency in women who are nulliparous or have a history of use of talc agents applied to the perineum; other risk factors include obesity and probably hormone replacement therapy. Protective factors include the use of oral contraceptives, multiparity, and breast-feeding. These protective factors are thought to work through suppression of ovulation and perhaps the associated reduction of ovulation associated inflammation of the ovarian epithelium or, alternatively, the serous epithelium located within the fimbriae of the fallopian tube. Other protective factors, such as fallopian tube ligation, are thought to protect the ovarian epithelium (or perhaps the distal fallopian tube fimbriae) from carcinogens that migrate from the vagina to the tubes and ovarian surface epithelium. Mucinous tumors are more frequent in women with a history of cigarette smoking, whereas endometrioid and clear cell tumors are more frequent in women with a history of endometriosis.

Considerable evidence now suggests that the precursor cell to serous carcinoma of the ovary might actually arise in the fimbria of the fallopian tube with extension or metastasis to the ovarian surface or capture of preneoplastic or neoplastic exfoliating tubal cells into an involuting ovarian follicle around the time of ovulation. Careful histologic and molecular analysis of tubal epithelium demonstrates molecular and histologic abnormalities, termed serous tubular intraepithelial carcinoma (STIC) lesions, in a high proportion of women undergoing risk-reducing salpingo-oophorectomies in the context of high-risk germline mutations in *BRCA1* and *BRCA2*, as well as a modest proportion of women with ovarian cancer in the absence of such mutations.



Genetic Risk Factors A variety of genetic syndromes substantially increase a woman’s risk of developing ovarian cancer. Approximately 10% of women with ovarian cancer have a germline mutation in one of two DNA repair genes: *BRCA1* (chromosome 17q12-21) or *BRCA2* (chromosome 13q12-13). Individuals inheriting a single copy of a mutant allele have a very high incidence of breast and ovarian cancer. Most of these women have a family history that is notable for multiple cases of breast and/or ovarian cancer, although inheritance through male members of the family can camouflage this genotype through several generations. The most common malignancy in these women is breast carcinoma, although women harboring germline *BRCA1* mutations have a marked increased risk of developing ovarian malignancies in their forties and fifties with a 30–50% lifetime risk of developing ovarian cancer. Women harboring a mutation in *BRCA2* have a lower penetrance of ovarian cancer with perhaps a 20–40% chance of developing this malignancy, with onset typically in their fifties or sixties. Women with a *BRCA2* mutation also are at slightly increased risk of pancreatic cancer. Likewise women with mutations in the DNA mismatch repair genes associated with Lynch syndrome, type 2 (*MSH2*, *MLH1*, *MLH6*, *PMS1*, *PMS2*) may have a risk of ovarian cancer as high as 1% per year in their forties and fifties. Finally, a small group of women with familial ovarian cancer may have mutations in other *BRCA*-associated genes such as *RAD51*, *CHK2*, and others. Screening studies in this select population suggest that current screening techniques, including serial evaluation of the CA-125 tumor marker and ultrasound, are insufficient at detecting early-stage and curable disease, so women with these germline mutations are advised to undergo prophylactic removal of ovaries and fallopian tubes typically after completing childbearing and ideally before age 35–40 years. Early prophylactic oophorectomy also protects these women from subsequent breast cancer with a reduction of breast cancer risk of approximately 50%.

Presentation Neoplasms of the ovary tend to be painless unless they undergo torsion. Symptoms are therefore typically related to compression of local organs or due to symptoms from metastatic disease. Women with tumors localized to the ovary do have an increased incidence of symptoms including pelvic discomfort, bloating, and perhaps changes in a woman’s typical urinary or bowel pattern. Unfortunately, these symptoms are frequently dismissed by either the woman or her health care team. It is believed that high-grade tumors metastasize early in the neoplastic process. Unlike other epithelial malignancies,

TABLE 117-1 STAGING AND SURVIVAL IN GYNECOLOGIC MALIGNANCIES

Stage	Ovarian	5-Year Survival, %	Endometrial	5-Year Survival, %	Cervix	5-Year Survival, %
0	—	—	—	—	Carcinoma in situ	100
I	Confined to ovary	90–95	Confined to corpus	89	Confined to uterus	85
II	Confined to pelvis	70–80	Involves corpus and cervix	73	Invades beyond uterus but not to pelvic wall	65
III	Intraabdominal spread	20–50	Extends outside the uterus but not outside the true pelvis or to lymph nodes	52	Extends to pelvic wall and/or lower third of vagina, or hydronephrosis	35
IV	Spread outside abdomen	1–5	Extends outside the true pelvis or involves the bladder or rectum	17	Invades mucosa of bladder or rectum or extends beyond the true pelvis	7

these tumors tend to exfoliate throughout the peritoneal cavity and thus present with symptoms associated with disseminated intraperitoneal tumors. The most common symptoms at presentation include a multimonth period of progressive complaints that typically include some combination of heartburn, nausea, early satiety, indigestion, constipation, and abdominal pain. Signs include the rapid increase in abdominal girth due to the accumulation of ascites that typically alerts the patient and her physician that the concurrent gastrointestinal symptoms are likely associated with serious pathology. Radiologic evaluation typically demonstrates a complex adnexal mass and ascites. Laboratory evaluation usually demonstrates a markedly elevated CA-125, a shed mucin (Muc 16) associated with, but not specific for, ovarian cancer. Hematogenous and lymphatic spread are seen but are not the typical presentation. Ovarian cancers are divided into four stages, with stage I tumors confined to the ovary, stage II malignancies confined to the pelvis, and stage III tumors confined to the peritoneal cavity (Table 117-1). These three stages are subdivided, with the most common presentation, stage IIIC, defined as tumors with bulky intraperitoneal disease. About 60% of women present with stage IIIC disease. Stage IV disease includes women with parenchymal metastases (liver, lung, spleen) or, alternatively, abdominal wall or pleural disease. The 40% not presenting with stage IIIC disease are roughly evenly distributed among the other stages, although mucinous and clear cell tumors are overrepresented in stage I tumors.

Screening Ovarian cancer is the fifth most lethal malignancy in women in the United States. It is curable in early stages, but seldom curable in advanced stages; hence, the development of effective screening strategies is of considerable interest. Furthermore, the ovary is well visualized with a variety of imaging techniques, most notably transvaginal ultrasound. Early-stage tumors often produce proteins that can be measured in the blood such as CA-125 and HE-4. Nevertheless, the incidence of ovarian cancer in the middle-aged female population is low, with only approximately 1 in 2000 women between the ages of 50 and 60 carrying an asymptomatic and undetected tumor. Thus effective screening techniques must be sensitive but, more importantly, highly specific to minimize the number of false-positive results. Even a screening test with 98% specificity and 50% sensitivity would have a positive predictive value of only about 1%. A large randomized study of active screening versus usual standard care demonstrated that a screening program consisting of six annual CA-125 measurements and four annual transvaginal ultrasounds in a population of women age 55–74 was not effective at reducing death from ovarian cancer and was associated with significant morbidity in the screened arm due to complications associated with diagnostic testing in the screened group. Although ongoing studies are evaluating the utility of alternative screening strategies, currently screening of normal-risk women is not recommended outside of a clinical trial.

TREATMENT OVARIAN CANCER

In women presenting with a localized ovarian mass, the principal diagnostic and therapeutic maneuver is to determine if the tumor is benign or malignant and, in the event that the tumor is malignant,

whether the tumor arises in the ovary or is a site of metastatic disease. Metastatic disease to the ovary can be seen from primary tumors of the colon, appendix, stomach (Krukenberg tumors), and breast. Typically women undergo a unilateral salpingo-oophorectomy, and if pathology reveals a primary ovarian malignancy, then the procedure is followed by a hysterectomy, removal of the remaining tube and ovary, omentectomy, and pelvic node sampling along with some random biopsies of the peritoneal cavity. This extensive surgical procedure is performed because approximately 30% of tumors that by visual inspection appear to be confined to the ovary have already disseminated to the peritoneal cavity and/or surrounding lymph nodes.

If there is evidence of bulky intraabdominal disease, a comprehensive attempt at maximal tumor cytoreduction is attempted even if it involves partial bowel resection, splenectomy, and in certain cases more extensive upper abdominal surgery. The ability to debulk metastatic ovarian cancer to minimal visible disease is associated with an improved prognosis compared with women left with visible disease. Patients without gross residual disease after resection have a median survival of 39 months, compared with 17 months for those left with macroscopic tumor. Once tumors have been surgically debulked, women receive therapy with a platinum agent, typically a taxane. Debate continues as to whether this therapy should be delivered intravenously or, alternatively, whether some of the therapy should be delivered directly into the peritoneal cavity via a catheter. Three randomized studies have demonstrated improved survival with intraperitoneal therapy, but this approach is still not widely accepted due to technical challenges associated with this delivery route and increased toxicity. In women who present with bulky intraabdominal disease, an alternative approach is to treat with platinum plus a taxane for several cycles before attempting a surgical debulking procedure (neoadjuvant therapy). Subsequent surgical procedures are more effective at leaving the patient without gross residual tumor and appear to be less morbid. Two studies have demonstrated that the neoadjuvant approach is associated with an overall survival that is comparable to the traditional approach of primary surgery followed by chemotherapy.

With optimal debulking surgery and platinum-based chemotherapy (usually carboplatin dosed to an area under the curve [AUC] of 6 plus paclitaxel 175 mg/m² by 3-h infusion in 21-day cycles), 70% of women who present with advanced-stage tumors respond, and 40–50% experience a complete remission with normalization of their CA-125, computed tomography (CT) scans, and physical examination. Unfortunately, a small proportion of women who obtain a complete response to therapy will remain in remission. Disease recurs within 1–4 years from the completion of their primary therapy in 75% of the complete responders. CA-125 levels often increase as a first sign of relapse; however, data are not clear that early intervention in relapsing patients influences survival. Recurrent disease is effectively managed, but not cured, with a variety of chemotherapeutic agents. Eventually all women with recurrent disease develop chemotherapy-refractory disease at which point refractory ascites, poor bowel motility, and obstruction or pseudoobstruction due to a

594 tumor-infiltrated aperistaltic bowel are common. Limited surgery to relieve intestinal obstruction, localized radiation therapy to relieve pressure or pain from masses, or palliative chemotherapy may be helpful. Agents with >15% response rates include gemcitabine, topotecan, liposomal doxorubicin, pemetrexed, and bevacizumab. Approximately 10% of ovarian cancers are HER2/neu positive, and trastuzumab may induce responses in this subset.

Five-year survival correlates with the stage of disease: stage I, 85–90%; stage II, 70–80%; stage III, 20–50%; and stage IV, 1–5% (Table 117-1). Low-grade serous tumors are molecularly distinct from high-grade serous tumors and are, in general, poorly responsive to chemotherapy. Targeted therapies focused on inhibiting kinases downstream of *RAS* and *BRAF* are being tested. Patients with tumors of low malignant potential are managed by surgery; chemotherapy and radiation therapy do not improve survival.

OVARIAN SEX CORD AND STROMAL TUMORS

Epidemiology, Presentation, and Predisposing Syndromes Approximately 7% of ovarian neoplasms are stromal or sex cord tumors, with approximately 1800 cases expected each year in the United States. Ovarian stromal tumors or sex cord tumors are most common in women in their fifties or sixties, but tumors can present in the extremes of age, including the pediatric population. These tumors arise from the mesenchymal components of the ovary, including steroid-producing cells as well as fibroblasts. Essentially all of these tumors are of low malignant potential and present as unilateral solid masses. Three clinical presentations are common: the detection of an abdominal mass; abdominal pain due to ovarian torsion, intratumoral hemorrhage, or rupture; or signs and symptoms due to hormonal production by these tumors.

The most common hormone-producing tumors include thecomas, granulosa cell tumor, or juvenile granulosa tumors in children. These estrogen-producing tumors often present with breast tenderness as well as isosexual precocious pseudopuberty in children, menometrorrhagia, oligomenorrhea, or amenorrhea in premenopausal women, or alternatively as postmenopausal bleeding in older women. In some women, estrogen-associated secondary malignancies, such as endometrial or breast cancer, may present as synchronous malignancies. Alternatively, endometrial cancer may serve as the presenting malignancy with evaluation subsequently identifying a unilateral solid ovarian neoplasm that proves to be an occult granulosa cell tumor. Sertoli-Leydig tumors often present with hirsutism, virilization, and occasionally Cushing's syndrome due to increased production of testosterone, androstenedione, or other 17-ketosteroids. Hormonally inert tumors include fibroma that presents as a solitary mass often in association with ascites and occasionally hydrothorax also known as Meigs' syndrome. A subset of these tumors present in individuals with a variety of inherited disorders that predispose them to mesenchymal neoplasia. Associations include juvenile granulosa cell tumors and perhaps Sertoli-Leydig tumors with Ollier's disease (multiple enchondromatosis) or Maffucci's syndrome, ovarian sex cord tumors with annular tubules with Peutz-Jeghers syndrome, and fibromas with Gorlin's disease. Essentially all granulosa tumors and a minority of juvenile granulosa cell tumors and thecomas have a defined somatic point mutation in the *FOXO2* gene at C134W generated by replacement of cysteine with a guanine at position 402. About 30% of Sertoli-Leydig tumors harbor a mutation in the RNA-processing gene *DICER* in the RNAPIIIb domain.

TREATMENT SEX CORD TUMORS

The mainstay of treatment for sex cord tumors is surgical resection. Most women present with tumors confined to the ovary. For the small subset of women who present with metastatic disease or develop evidence of tumor recurrence after primary resection, survival is still typically long, often in excess of a decade. Because these tumors are slow growing and relatively refractory to chemotherapy,

women with metastatic disease are often debulked because disease is usually peritoneal-based (as with epithelial ovarian cancer). Definitive data that surgical debulking of metastatic or recurrent disease prolongs survival are lacking, but ample data document women who have survived years or, in some cases, decades after resection of recurrent disease. In addition, large peritoneal-based metastases also have a proclivity for hemorrhage, sometimes with catastrophic complications. Chemotherapy is occasionally effective, and women tend to receive regimens designed to treat epithelial or germ cell tumors. Bevacizumab has some activity in clinical trials but is not approved for this specific indication. These tumors often produce high levels of müllerian inhibiting substance (MIS), inhibin, and, in the case of Sertoli-Leydig tumors, a fetoprotein (AFP). These proteins are detectable in serum and can be used as tumor markers to monitor women for recurrent disease because the increase or decrease of these proteins in the serum tends to reflect the changing bulk of systemic tumor.

GERM CELL TUMORS OF THE OVARY

Germ cell tumors, like their counterparts in the testis, are cancers of germ cells. These totipotent cells contain the programming for differentiation to essentially all tissue types, and hence the germ cell tumors include a histologic menagerie of bizarre tumors, including benign teratomas and a variety of malignant tumors, such as immature teratomas, dysgerminomas, yolk sac malignancies, and choriocarcinomas. Benign teratoma (or dermoid cyst) is the most common germ cell neoplasm of the ovary and often presents in young woman. These tumors include a complex mixture of differentiated tissue including tissues from all three germ layers. In older women, these differentiated tumors can develop malignant transformation, most commonly squamous cell carcinomas. Malignant germ cell tumors include dysgerminomas, yolk sac tumors, immature teratomas, and embryonal carcinoma and choriocarcinomas. There are no known genetic abnormalities that unify these tumors. A subset of dysgerminomas harbor mutations in *c-Kit* oncogenes (as seen in gastrointestinal stromal tumors [GIST]), whereas a subset of germ cell tumors have isochromosome 12 abnormalities, as seen in testicular malignancies. In addition, a subset of dysgerminomas is associated with dysgenetic ovaries. Identification of a dysgerminoma arising in genotypic XY gonads is important in that it highlights the need to identify and remove the contralateral gonad due to risk of gonadoblastoma.

Presentation Germ cell tumors can present at all ages, but the peak age of presentation tends to be in females in their late teens or early twenties. Typically these tumors will become large ovarian masses, which eventually present as palpable low abdominal or pelvic masses. Like sex cord tumors, torsion or hemorrhage may present urgently or emergently as acute abdominal pain. Some of these tumors produce elevated levels of human chorionic gonadotropin (hCG), which can lead to isosexual precocious puberty when tumors present in younger girls. Unlike epithelial ovarian cancer, these tumors have a higher proclivity for nodal or hematogenous metastases. As with testicular tumors, some of these tumors tend to produce AFP (yolk sac tumors) or hCG (embryonal carcinoma, choriocarcinomas, and some dysgerminomas) that are reliable tumor markers.

TREATMENT GERM CELL TUMORS

Germ cell tumors typically present in women who are still of child-bearing age, and because bilateral tumors are uncommon (except in dysgerminoma, 10–15%), the typical treatment is unilateral oophorectomy or salpingo-oophorectomy. Because nodal metastases to pelvic and para-aortic nodes are common and may affect treatment choices, these nodes should be carefully inspected and, if enlarged, should be resected if possible. Women with malignant germ cell tumors typically receive bleomycin, etoposide, and cisplatin (BEP) chemotherapy. In the majority of women, even those with

advanced-stage disease, cure is expected. Close follow-up without adjuvant therapy of women with stage I tumors is reasonable if there is high confidence that the patient and health care team are committed to compulsive and careful follow-up, as chemotherapy at the time of tumor recurrence is likely to be curative.

Dysgerminoma is the ovarian counterpart of testicular seminoma. The 5-year disease-free survival is 100% in early-stage patients and 61% in stage III disease. Although the tumor is highly radiation-sensitive, radiation produces infertility in many patients. BEP chemotherapy is as effective or more so without causing infertility. The use of BEP following incomplete resection is associated with a 2-year disease-free survival rate of 95%. This chemotherapy is now the treatment of choice for dysgerminoma.

FALLOPIAN TUBE CANCER

Transport of the egg to the uterus occurs via transit through the fallopian tube, with the distal ends of these tubes composed of fimbriae that drape about the ovarian surface and capture the egg as it erupts from the ovarian cortex. Fallopian tube malignancies are typically serous tumors. Previous teaching was that these malignancies were rare, but more careful histologic examination suggests that many “ovarian malignancies” might actually arise in the distal fimbria of the fallopian tube (see above). These women often present with adnexal masses, and like ovarian cancer, these tumors spread relatively early throughout the peritoneal cavity and respond to platinum and taxane therapy and have a natural history that is essentially identical to ovarian cancer (Table 117-1).

CERVICAL CANCER

GLOBAL CONSIDERATIONS



Cervical cancer is the second most common and most lethal malignancy in women worldwide likely due to the widespread infection with high-risk strains of human papillomavirus (HPV) and limited utilization of or access to Pap smear screening in many nations throughout the world. Nearly 500,000 cases of cervical cancer are expected worldwide, with approximately 240,000 deaths annually. Cancer incidence is particularly high in women residing in Central and South America, the Caribbean, and southern and eastern Africa. Mortality rate is disproportionately high in Africa. In the United States, 12,360 women were diagnosed with cervical cancer and 4020 women died in 2014. Developed countries have looked at high-technology screening techniques for HPV involving automated polymerase chain reaction in thin preps that identify dysplastic cytology as well as high-risk HPV genetic material. Visual inspection of the cervix coated with acetic acid has demonstrated the ability to reduce mortality from cervical cancer with potential broad applicability in low-resource environments. The development of effective vaccines for high-risk HPV types makes it imperative to determine economical, socially acceptable, and logistically feasible strategies to deliver and distribute this vaccine to girls and boys before their engagement in sexual activity.

HPV INFECTION AND PREVENTIVE VACCINATION

HPV is the primary neoplastic-initiating event in the vast majority of women with invasive cervical cancer. This double-strand DNA virus infects epithelium near the transformation zone of the cervix. More than 60 types of HPV are known, with approximately 20 types having the ability to generate high-grade dysplasia and malignancy. HPV-16 and -18 are the types most frequently associated with high-grade dysplasia and targeted by both U.S. Food and Drug Administration–approved vaccines. The large majority of sexually active adults are exposed to HPV, and most women clear the infection without specific intervention. The 8-kilobase HPV genome encodes seven early genes, most notably *E6* and *E7*, which can bind to *RB* and *p53*, respectively. High-risk types of HPV encode *E6* and *E7* molecules that are particularly effective at inhibiting the normal cell cycle checkpoint functions of these regulatory proteins, leading to immortalization but not full

transformation of cervical epithelium. A minority of woman will fail to clear the infection with subsequent HPV integration into the host genome. Over the course of as short as months but more typically years, some of these women develop high-grade dysplasia. The time from dysplasia to carcinoma is likely years to more than a decade and almost certainly requires the acquisition of other poorly defined genetic mutations within the infected and immortalized epithelium.

Risk factors for HPV infection and, in particular, dysplasia include a high number of sexual partners, early age of first intercourse, and history of venereal disease. Smoking is a cofactor; heavy smokers have a higher risk of dysplasia with HPV infection. HIV infection, especially when associated with low CD4+ T cell counts, is associated with a higher rate of high-grade dysplasia and likely a shorter latency period between infection and invasive disease. The administration of highly active antiretroviral therapy reduces the risk of high-grade dysplasia associated with HPV infection.

Currently approved vaccines include the recombinant proteins to the late proteins, L1 and L2, of HPV-16 and -18. Vaccination of women before the initiation of sexual activity dramatically reduces the rate of HPV-16 and -18 infection and subsequent dysplasia. There is also partial protection against other HPV types, although vaccinated women are still at risk for HPV infection and still require standard Pap smear screening. Although no randomized trial data demonstrate the utility of Pap smears, the dramatic drop in cervical cancer incidence and death in developed countries employing wide-scale screening provides strong evidence for its effectiveness. In addition, even visual inspection of the cervix with preapplication of acetic acid using a “see and treat” strategy has demonstrated a 30% reduction in cervical cancer death. The incorporation of HPV testing by polymerase chain reaction or other molecular techniques increases the sensitivity of detecting cervical pathology but at the cost of identifying many women with transient infections who require no specific medical intervention.

CLINICAL PRESENTATIONS

The majority of cervical malignancies are squamous cell carcinomas associated with HPV. Adenocarcinomas are also HPV-related and arise deep in the endocervical canal; they are typically not seen by visual inspection of the cervix and thus are often missed by Pap smear screening. A variety of rarer malignancies including atypical epithelial tumors, carcinoids, small cell carcinomas, sarcomas, and lymphomas have also been reported.

The principal role of Pap smear testing is the detection of asymptomatic preinvasive cervical dysplasia of squamous epithelial lining. Invasive carcinomas often have symptoms or signs including post-coital spotting or intermenstrual cycle bleeding or menometrorrhagia. Foul-smelling or persistent yellow discharge may also be seen. Presentations that include pelvic or sacral pain suggest lateral extension of the tumor into pelvic nerve plexus by either the primary tumor or a pelvic node and are signs of advanced-stage disease. Likewise, flank pain from hydronephrosis from ureteral compression or deep venous thrombosis from iliac vessel compression suggests either extensive nodal disease or direct extension of the primary tumor to the pelvic sidewall. The most common finding of physical exam is a visible tumor on the cervix.

TREATMENT CERVICAL CANCER

Scans are not part of the formal clinical staging of cervical cancer yet are very useful in planning appropriate therapy. CT can detect hydronephrosis indicative of pelvic sidewall disease but is not accurate at evaluating other pelvic structures. Magnetic resonance imaging (MRI) is more accurate at estimating uterine extension and paracervical extension of disease into soft tissues typically bordered by broad and cardinal ligaments that support the uterus in the central pelvis. Positron emission tomography (PET) scan is the most accurate technique for evaluating the pelvis and more importantly nodal (pelvic, para-aortic, and scalene) sites for disease.

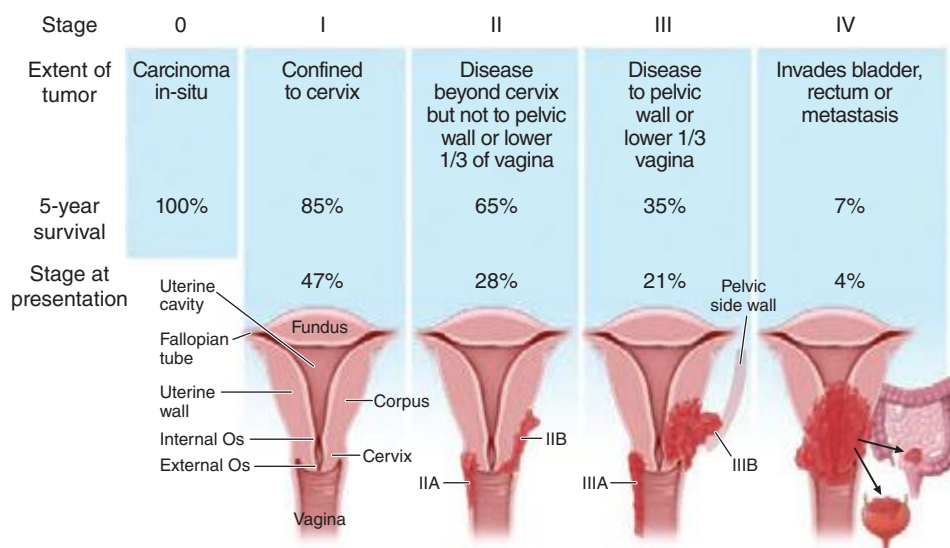


FIGURE 117-1 Anatomic display of the stages of cervix cancer defined by location, extent of tumor, frequency of presentation, and 5-year survival.

This technique seems more prognostic and accurate than CT, MRI, or lymphangiogram, especially in the para-aortic region.

Stage I cervical tumors are confined to the cervix, whereas stage II tumors extend into the upper vagina or paracervical soft tissue (Fig. 117-1). Stage III tumors extend to the lower vagina or the pelvic sidewalls, whereas stage IV tumors invade the bladder or rectum or have spread to distant sites. Very small stage I cervical tumors can be treated with a variety of surgical procedures. In young women desiring to maintain fertility, radical trachelectomy removes the cervix with subsequent anastomosis of the upper vagina to the uterine corpus. Larger cervical tumors confined to the cervix can be treated with either surgical resection or radiation therapy in combination with cisplatin-based chemotherapy with a high chance of cure. Larger tumors that extend regionally down the vagina or into the paracervical soft tissues or the pelvic sidewalls are treated with combination chemotherapy and radiation therapy. The treatment of recurrent or metastatic disease is unsatisfactory due to the relative resistance of these tumors to chemotherapy and currently available biological agents, although bevacizumab, a monoclonal antibody that is said to inhibit tumor-associated angiogenesis, has demonstrated clinically meaningful activity in the management of metastatic disease.

UTERINE CANCER

EPIDEMIOLOGY

Several different tumor types arise in uterine corpus. Most tumors arise in the glandular lining and are endometrial adenocarcinomas. Tumors can also arise in the smooth muscle; most are benign (uterine leiomyoma), with a small minority of tumors being sarcomas. The endometrioid histologic subtype of endometrial cancer is the most common gynecologic malignancy in the United States. In 2014, an estimated 52,630 women were diagnosed with cancer of the uterine corpus, with 8590 deaths from the disease. Development of these tumors is a multistep process, with estrogen playing an important early role in driving endometrial gland proliferation. Relative overexposure to this class of hormones is a risk factor for the subsequent development of endometrioid tumors. In contrast, progestins drive glandular maturation and are protective. Hence, women with high endogenous or pharmacologic exposure to estrogens, especially if unopposed by progesterone, are at high risk for endometrial cancer.

Obese women, women treated with unopposed estrogens, or women with estrogen-producing tumors (such as granulosa cell tumors of the ovary) are at higher risk for endometrial cancer. In addition, treatment with tamoxifen, which has antiestrogenic effects in breast tissue but estrogenic effects in uterine epithelium, is associated with an increased risk of endometrial cancer. Events such as the loss of the *PTEN* tumor suppressor gene with activation and often additional mutations in the *PIK-3CA/AKT* pathways likely serve as secondary events in carcinogenesis. The Cancer Genome Atlas Research Network has demonstrated that endometrioid tumors can be divided into four subgroups: ultramutated, microsatellite instability hypermutated, copy number low, and copy number high subgroups. These groups have different natural histories; therapy for these subgroups may eventually be individualized. Serous tumors of the uterine corpus represent approximately 5–10% of epithelial tumors of the uterine corpus and possess distinct molecular characteristics that are most similar to those seen in serous tumors arising in the ovary or fallopian tube.

Women with a mutation in one of a series of DNA mismatch repair genes associated with the Lynch syndrome, also known as hereditary nonpolyposis colon cancer (HNPCC), are at increased risk for endometrioid endometrial carcinoma. These individuals have germline mutations in *MSH2*, *MLH1*, and in rare cases *PMS1* and *PMS2*, with resulting microsatellite instability and hypermutation. Individuals who carry these mutations typically have a family history of cancer and are at markedly increased risk for colon cancer and modestly increased risk for ovarian cancer and a variety of other tumors. Middle-aged women with HNPCC carry a 4% annual risk of endometrial cancer and a relative overall risk of approximately 200-fold as compared to age-matched women without HNPCC.

PRESENTATIONS

The majority of women with tumors of the uterine corpus present with postmenopausal vaginal bleeding due to shedding of the malignant endometrial lining. Premenopausal women often will present with atypical bleeding between typical menstrual cycles. These signs typically bring a woman to the attention of a health care professional, and hence the majority of women present with early-stage disease with the tumor confined to the uterine corpus. Diagnosis is typically established by endometrial biopsy. Epithelial tumors may spread to pelvic or para-aortic lymph nodes. Pulmonary metastases can appear later in the natural history of this disease but are very uncommon at initial presentation. Serous tumors tend to have patterns of spread much more

reminiscent of ovarian cancer with many patients presenting with disseminated peritoneal disease and sometimes ascites. Some women presenting with uterine sarcomas will present with pelvic pain. Nodal metastases are uncommon with sarcomas, which are more likely to present with either intraabdominal disease or pulmonary metastases.

TREATMENT UTERINE CANCER

Most women with endometrial cancer have disease that is localized to the uterus (75% are stage I, Table 117-1), and definitive treatment typically involves a hysterectomy with removal of the ovaries and fallopian tubes. The resection of lymph nodes does not improve outcome but does provide prognostic information. Node involvement defines stage III disease, which is present in 13% of patients. Tumor grade and depth of invasion are the two key prognostic variables in early-stage tumors, and women with low-grade and/or minimally invasive tumors are typically observed after definitive surgical therapy. Patients with high-grade tumors or tumors that are deeply invasive (stage IB, 13%) are at higher risk for pelvic recurrence or recurrence at the vaginal cuff, which is typically prevented by vaginal vault brachytherapy.

Women with regional metastases or metastatic disease (3% of patients) with low-grade tumors can be treated with progesterone. Poorly differentiated tumors are typically resistant to hormonal manipulation and thus are treated with chemotherapy. The role of chemotherapy in the adjuvant setting is currently under investigation. Chemotherapy for metastatic disease is delivered with palliative intent. Drugs that effectively target and inhibit signaling of the AKT-mTOR pathway are currently under investigation.

Five-year survival is 89% for stage I, 73% for stage II, 52% for stage III, and 17% for stage IV disease (Table 117-1).

GESTATIONAL TROPHOBLASTIC TUMORS

GLOBAL CONSIDERATIONS



Gestational trophoblastic diseases represent a spectrum of neoplasia from benign hydatidiform mole to choriocarcinoma due to persistent trophoblastic disease associated most commonly with molar pregnancy but occasionally seen after normal gestation. The most common presentations of trophoblastic tumors are partial and complete molar pregnancies. These represent approximately 1 in 1500 conceptions in developed Western countries. The incidence widely varies globally, with areas in Southeast Asia having a much higher incidence of molar pregnancy. Regions with high molar pregnancy rates are often associated with diets low in carotene and animal fats.

RISK FACTORS

Trophoblastic tumors result from the outgrowth or persistence of placental tissue. They arise most commonly in the uterus but can also arise in other sites such as the fallopian tubes due to ectopic pregnancy. Risk factors include poorly defined dietary and environmental factors as well as conceptions at the extremes of reproductive age, with the incidence particularly high in females conceiving younger than age 16 or older than age 50. In older women, the incidence of molar pregnancy might be as high as one in three, likely due to increased risk of abnormal fertilization of the aged ova. Most trophoblastic neoplasms are associated with complete moles, diploid tumors with all genetic material from the paternal donor (known as parental disomy). This is thought to occur when a single sperm fertilizes an enucleate egg that subsequently duplicates the paternal DNA. Trophoblastic proliferation occurs with exuberant villous stroma. If pseudopregnancy extends out past the 12th week, fluid progressively accumulates within the stroma, leading to “hydropic changes.” There is no fetal development in complete moles.

Partial moles arise from the fertilization of an egg with two sperm; hence two-thirds of genetic material is paternal in these triploid tumors. Hydropic changes are less dramatic, and fetal development can often occur through late first trimester or early second trimester at which point spontaneous abortion is common. Laboratory findings will include excessively high hCG and high AFP. The risk of persistent gestational trophoblastic disease after partial mole is approximately 5%. Complete and partial moles can be noninvasive or invasive. Myometrial invasion occurs in no more than one in six complete moles and a lower portion of partial moles.

PRESENTATION OF INVASIVE TROPHOBLASTIC DISEASE

The clinical presentation of molar pregnancy is changing in developed countries due to the early detection of pregnancy with home pregnancy kits and the very early use of Doppler and ultrasound to evaluate the early fetus and uterine cavity for evidence of a viable fetus. Thus, in these countries, the majority of women presenting with trophoblastic disease have their moles detected early and have typical symptoms of early pregnancy including nausea, amenorrhea, and breast tenderness. With uterine evacuation of early complete and partial moles, most women experience spontaneous remission of their disease as monitored by serial hCG levels. These women require no chemotherapy. Patients with persistent elevation of hCG or rising hCG after evacuation have persistent or actively growing gestational trophoblastic disease and require therapy. Most series suggest that between 15 and 25% of women will have evidence of persistent gestational trophoblastic disease after molar evacuation.

In women who lack access to prenatal care, presenting symptoms can be life threatening including the development of preeclampsia or even eclampsia. Hyperthyroidism can also be seen. Evacuation of large moles can be associated with life-threatening complications including uterine perforation, volume loss, high-output cardiac failure, and adult respiratory distress syndrome (ARDS).

For women with evidence of rising hCG or radiologic confirmation of metastatic or persistent regional disease, prognosis can be estimated through a variety of scoring algorithms that identify those women at low, intermediate, and high risk for requiring multiagent chemotherapy. In general, women with widely metastatic nonpulmonary disease, very elevated hCG, and prior normal antecedent term pregnancy are considered at high risk and typically require multiagent chemotherapy for cure.

TREATMENT INVASIVE TROPHOBLASTIC DISEASE

The management for a persistent and rising hCG after evacuation of a molar conception is typically chemotherapy, although surgery can play an important role for disease that is persistently isolated in the uterus (especially if childbearing is complete) or to control hemorrhage. For women wishing to maintain fertility or with metastatic disease, the preferred treatment is chemotherapy. Chemotherapy is guided by the hCG level, which typically drops to undetectable levels with effective therapy. Single-agent treatment with methotrexate or dactinomycin cures 90% of women with low-risk disease. Patients with high-risk disease (high hCG levels, presentation 4 or more months after pregnancy, brain or liver metastases, failure of methotrexate therapy) are typically treated with multiagent chemotherapy (e.g., etoposide, methotrexate, and dactinomycin alternating with cyclophosphamide and vincristine [EMA-CO]), which is typically curative even in women with extensive metastatic disease. Cisplatin, bleomycin, and either etoposide or vinblastine are also active combinations. Survival in high-risk disease exceeds 80%. Cured women may get pregnant again without evidence of increased fetal or maternal complications.

118 Primary and Metastatic Tumors of the Nervous System

Lisa M. DeAngelis, Patrick Y. Wen

Primary brain tumors are diagnosed in approximately 52,000 people each year in the United States. At least one-half of these tumors are malignant and associated with a high mortality. Glial tumors account for about 30% of all primary brain tumors, and 80% of those are malignant. Meningiomas account for 35%, vestibular schwannomas 10%, and central nervous system (CNS) lymphomas about 2%. Brain metastases are three times more common than all primary brain tumors combined and are diagnosed in approximately 150,000 people each year. Metastases to the leptomeninges and epidural space of the spinal cord each occur in approximately 3–5% of patients with systemic cancer and are also a major cause of neurologic disability.

APPROACH TO THE PATIENT: Primary and Metastatic Tumors of the Nervous System

CLINICAL FEATURES

Brain tumors of any type can present with a variety of symptoms and signs that fall into two categories: general and focal; patients often have a combination of the two (**Table 118-1**). General or nonspecific symptoms include headache, with or without nausea or vomiting, cognitive difficulties, personality change, and gait disorder. Generalized symptoms arise when the enlarging tumor and its surrounding edema cause an increase in intracranial pressure or direct compression of cerebrospinal fluid (CSF) circulation leading to hydrocephalus. The classic headache associated with a brain tumor is most evident in the morning and improves during the day, but this particular pattern is actually seen in a minority of patients. Headaches are often holocephalic but can be ipsilateral to the side of a tumor. Occasionally, headaches have features of a typical migraine with unilateral throbbing pain associated with visual scotoma. Personality changes may include apathy and withdrawal from social circumstances, mimicking depression. Focal or lateralizing findings include hemiparesis, aphasia, or visual field defect. Lateralizing symptoms are typically subacute and progressive. A visual field defect is often unnoticed by the patient; its presence may only be revealed after it leads to an injury such as an automobile accident occurring in the blind visual field. Language difficulties may be mistaken for confusion. Seizures are a common presentation of brain tumors, occurring in about 25% of patients with brain metastases or malignant gliomas but can be the presenting symptom in up to 90% of patients with a low-grade glioma. All seizures that arise from a brain tumor will have a focal onset whether or not it is apparent clinically.

TABLE 118-1 SYMPTOMS AND SIGNS AT PRESENTATION OF BRAIN TUMORS

	High-Grade Glioma (%)	Low-Grade Glioma (%)	Meningioma (%)	Metastases (%)
Generalized				
Impaired cognitive function	50	10	30	60
Hemiparesis	40	10	36	60
Headache	50	40	37	50
Lateralizing				
Seizures	20	70+	17	18
Aphasia	20	<5	—	18
Visual field deficit	—	—	—	7

NEUROIMAGING

Cranial MRI is the preferred diagnostic test for any patient suspected of having a brain tumor and should be performed with gadolinium contrast administration. Computed tomography (CT) scan should be reserved for those patients unable to undergo magnetic resonance imaging (MRI; e.g., pacemaker). Malignant brain tumors—whether primary or metastatic—typically enhance with gadolinium and may have central areas of necrosis; they are characteristically surrounded by edema of the neighboring white matter. Low-grade gliomas usually do not enhance with gadolinium and are best appreciated on fluid-attenuated inversion recovery (FLAIR) MRIs. Meningiomas have a characteristic appearance on MRI because they are dural-based with a dural tail and compress but do not invade the brain. Dural metastases or a dural lymphoma can have a similar appearance. Imaging is characteristic for many primary and metastatic tumors and sometimes will suffice to establish a diagnosis when the location precludes surgical intervention (e.g., brainstem glioma). Functional MRI is useful in presurgical planning to define eloquent sensory, motor, or language cortex. Positron emission tomography (PET) is useful in determining the metabolic activity of the lesions seen on MRI; MR perfusion and spectroscopy can provide information on blood flow or tissue composition. These techniques may help distinguish tumor progression from necrotic tissue as a consequence of treatment with radiation and chemotherapy or identify foci of high-grade tumor in an otherwise low-grade-appearing glioma.

Neuroimaging is the only test necessary to diagnose a brain tumor. Laboratory tests are rarely useful, although patients with metastatic disease may have elevation of a tumor marker in their serum that reflects the presence of brain metastases (e.g., β human chorionic gonadotropin [β -hCG] from testicular cancer). Additional testing such as cerebral angiogram, electroencephalogram (EEG), or lumbar puncture is rarely indicated or helpful.

TREATMENT BRAIN TUMORS

Therapy of any intracranial malignancy requires both symptomatic and definitive treatments. Definitive treatment is based on the specific tumor type and includes surgery, radiotherapy, and chemotherapy. However, symptomatic treatments apply to brain tumors of any type. Most high-grade malignancies are accompanied by substantial surrounding edema, which contributes to neurologic disability and raised intracranial pressure. Glucocorticoids are highly effective at reducing perilesional edema and improving neurologic function, often within hours of administration. Dexamethasone has been the glucocorticoid of choice because of its relatively low mineralocorticoid activity. Initial doses are typically 12–16 mg/d in divided doses given orally or IV (both are equivalent). Although glucocorticoids rapidly ameliorate symptoms and signs, their long-term use causes substantial toxicity including insomnia, weight gain, diabetes mellitus, steroid myopathy, and personality changes. Consequently, a taper is indicated as definitive treatment is administered and the patient improves.

Patients with brain tumors who present with seizures require antiepileptic drug therapy. There is no role for prophylactic antiepileptic drugs in patients who have not had a seizure. The agents of choice are those drugs that do not induce the hepatic microsomal enzyme system. These include levetiracetam, topiramate, lamotrigine, valproic acid, and lacosamide (**Chap. 445**). Other drugs, such as phenytoin and carbamazepine, are used less frequently because they are potent enzyme inducers that can interfere with both glucocorticoid metabolism and the metabolism of chemotherapeutic agents needed to treat the underlying systemic malignancy or the primary brain tumor. Venous thromboembolic disease occurs in 20–30% of patients with high-grade gliomas and brain metastases. Therefore, prophylactic anticoagulants should be used during hospitalization and in nonambulatory patients. Those who have had either a deep vein thrombosis or pulmonary embolus can receive therapeutic doses of anticoagulation safely and without increasing

the risk for hemorrhage into the tumor. Inferior vena cava filters are reserved for patients with absolute contraindications to anticoagulation such as recent craniotomy.

PRIMARY BRAIN TUMORS

PATHOGENESIS

No underlying cause has been identified for the majority of primary brain tumors. The only established risk factors are exposure to ionizing radiation (meningiomas, gliomas, and schwannomas) and immunosuppression (primary CNS lymphoma). Evidence for an association with exposure to electromagnetic fields including cellular telephones, head injury, foods containing *N*-nitroso compounds, or occupational risk factors are unproven. A small minority of patients have a family history of brain tumors. Some of these familial cases are associated with genetic syndromes (Table 118-2).

As with other neoplasms, brain tumors arise as a result of a multistep process driven by the sequential acquisition of genetic alterations. These include loss of tumor-suppressor genes (e.g., *p53* and phosphatase and tensin homolog on chromosome 10 [*PTEN*]) and amplification and overexpression of protooncogenes such as the epidermal growth factor receptor (*EGFR*) and the platelet-derived growth factor receptors (*PDGFR*). The accumulation of these genetic abnormalities results in uncontrolled cell growth and tumor formation.

Important progress has been made in understanding the molecular pathogenesis of several types of brain tumors, including glioblastoma and medulloblastoma. Morphologically indistinguishable glioblastomas can be separated into four subtypes defined by molecular profiling: (1) classical, characterized by overactivation of the *EGFR* pathway; (2) proneural, characterized by overexpression of *PDGFRA*, mutations

of the isocitrate dehydrogenase (*IDH*) 1 and 2 genes, and expression of neural markers; (3) mesenchymal, defined by expression of mesenchymal markers and loss of *NF1*; and (4) neural, characterized by overactivity of *EGFR* and expression of neural markers. The clinical implications of these subtypes are under study. Medulloblastoma is the other primary brain tumor that has been highly analyzed, and four molecular subtypes have also been identified: (1) the Wnt subtype is defined by a mutation in β -catenin and has an excellent prognosis; (2) the SHH subtype has mutations in *PTCH1*, *SMO*, *GLI2*, or *SUFU* and has an intermediate prognosis; (3) group 3 has elevated *MYC* expression and has the worst prognosis; and (4) group 4 is characterized by isochromosome 17q. Targeted therapeutics are under development for some of the medulloblastoma subtypes, especially the SHH group.

INTRINSIC "MALIGNANT" TUMORS

ASTROCYTOMAS

These are infiltrative tumors with a presumptive glial cell of origin. The World Health Organization (WHO) classifies astrocytomas into four prognostic grades based on histologic features: grade I (pilocytic astrocytoma, subependymal giant cell astrocytoma); grade II (diffuse astrocytoma); grade III (anaplastic astrocytoma); and grade IV (glioblastoma). Grades I and II are considered low-grade astrocytomas, and grades III and IV are considered high-grade astrocytomas.

Low-Grade Astrocytoma These tumors occur predominantly in children and young adults.

GRADE I ASTROCYTOMAS Pilocytic astrocytomas (WHO grade I) are the most common tumor of childhood. They occur typically in the cerebellum but may also be found elsewhere in the neuraxis, including the optic nerves and brainstem. Frequently they appear as cystic lesions

TABLE 118-2 GENETIC SYNDROMES ASSOCIATED WITH PRIMARY BRAIN TUMORS

Syndrome	Inheritance	Gene/Protein	Associated Tumors
Cowden's syndrome	AD	Mutations of <i>PTEN</i> (ch10p23)	Dysplastic cerebellar gangliocytoma (Lhermitte-Duclos disease), meningioma, astrocytoma Breast, endometrial, thyroid cancer, trichilemmomas
Familial schwannomatosis	Sporadic Hereditary	Mutations in <i>INI1/SNF5</i> (ch22q11)	Schwannomas, gliomas
Gardner's syndrome	AD	Mutations in <i>APC</i> (ch5q21)	Medulloblastoma, glioblastoma, craniopharyngioma Familial polyposis, multiple osteomas, skin and soft tissue tumors
Gorlin syndrome (basal cell nevus syndrome)	AD	Mutations in <i>Patched 1</i> gene (ch9q22.3)	Medulloblastomas Basal cell carcinoma
Li-Fraumeni syndrome	AD	Mutations in <i>p53</i> (ch17p13.1)	Gliomas, medulloblastomas Sarcomas, breast cancer, leukemias, others
Multiple endocrine neoplasia 1 (Werner's syndrome)	AD	Mutations in <i>Menin</i> (ch11q13)	Pituitary adenoma, malignant schwannomas Parathyroid and pancreatic islet cell tumors
Neurofibromatosis type 1 (NF1)	AD	Mutations in <i>NF1</i> /neurofibromin (ch17q12-22)	Schwannomas, astrocytomas, optic nerve gliomas, meningiomas Neurofibromas, neurofibrosarcomas, others
Neurofibromatosis type 2 (NF2)	AD	Mutations in <i>NF2</i> /merlin (ch22q12)	Bilateral vestibular schwannomas, astrocytomas, multiple meningiomas, ependymomas
Tuberous sclerosis (TSC) (Bourneville disease)	AD	Mutations in <i>TSC1/TSC2</i> (ch9q34/16)	Subependymal giant-cell astrocytoma, ependymomas, glioma, ganglioglioma, hamartoma
Turcot syndrome	AD AR	Mutations in <i>APC</i> ^a (ch5) hMLH1 (ch3p21)	Gliomas, medulloblastomas Adenomatous colon polyps, adenocarcinoma
von Hippel-Lindau (VHL)	AD	Mutations in <i>VHL</i> gene (ch3p25)	Hemangioblastomas Retinal angiomas, renal cell carcinoma, pheochromocytoma, pancreatic tumors and cysts, endolymphatic sac tumors of the middle ear

^aVarious DNA mismatch repair gene mutations may cause a similar clinical phenotype, also referred to as Turcot syndrome, in which there is a predisposition to nonpolyposis colon cancer and brain tumors.

Abbreviations: AD, autosomal dominant; APC, adenomatous polyposis coli; AR, autosomal recessive; ch, chromosome; PTEN, phosphatase and tensin homologue; TSC, tuberous sclerosis complex.

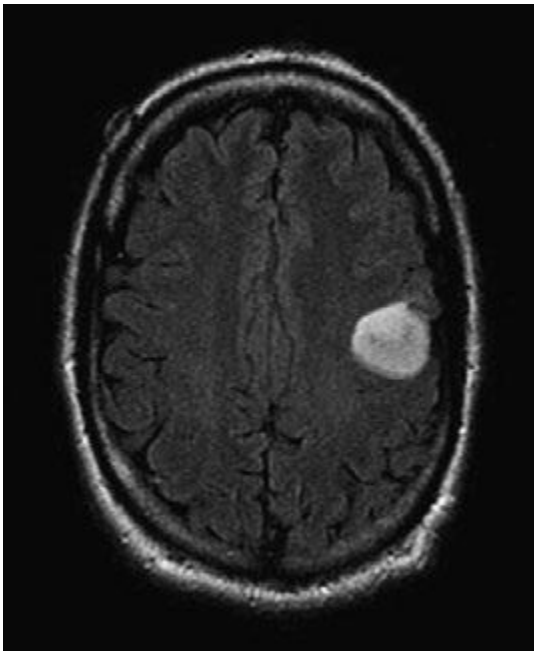


FIGURE 118-1 Fluid-attenuated inversion recovery (FLAIR) MRI of a left frontal low-grade astrocytoma. This lesion did not enhance.

with an enhancing mural nodule. These are well-demarcated lesions that are potentially curable if they can be resected completely. Giant-cell subependymal astrocytomas are usually found in the ventricular wall of patients with tuberous sclerosis. They often do not require intervention but can be treated surgically or with inhibitors of the mammalian target of rapamycin (mTOR).

GRADE II ASTROCYTOMAS These are infiltrative tumors that usually present with seizures in young adults. They appear as nonenhancing tumors with increased T2/FLAIR signal (Fig. 118-1). If feasible, patients should undergo maximal surgical resection, although complete resection is rarely possible because of the invasive nature of the tumor. Radiation therapy (RT) is helpful, but there is no difference in overall survival between RT administered postoperatively or delayed until the time of tumor progression. There is increasing evidence that chemotherapeutic agents such as temozolomide, an oral alkylating agent, can be helpful in some patients. The tumor transforms to a malignant astrocytoma in the majority of patients, leading to variable survival with a median of about 5 years.

High-Grade Astrocytoma

GRADE III (ANAPLASTIC) ASTROCYTOMA These account for approximately 15–20% of high-grade astrocytomas. They generally present in the fourth and fifth decades of life as variably enhancing tumors. Treatment is the same as for glioblastoma, consisting of maximal safe surgical resection followed by RT with concurrent and adjuvant temozolomide or by RT and adjuvant temozolomide alone.

GRADE IV ASTROCYTOMA (GLIOBLASTOMA) Glioblastoma accounts for the majority of high-grade astrocytomas. They are the most common malignant primary brain tumor, with over 10,000 cases diagnosed each year in the United States. Patients usually present in the sixth and seventh decades of life with headache, seizures, or focal neurologic deficits. The tumors appear as ring-enhancing masses with central necrosis and surrounding edema (Fig. 118-2). These are highly infiltrative tumors, and the areas of increased T2/FLAIR signal surrounding the main tumor mass contain invading tumor cells. Treatment involves maximal surgical resection followed by partial-field external-beam RT (6000 cGy in thirty 200-cGy fractions) with concomitant temozolomide, followed by 6–12 months of adjuvant temozolomide. With this regimen, median survival is increased to 14.6 months compared to only 12 months with RT alone, and 2-year survival is increased to 27%, compared to 10% with RT alone. Patients whose tumor contains

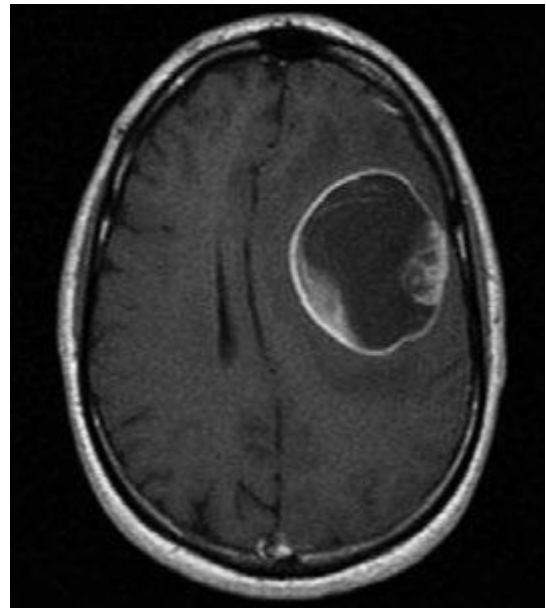


FIGURE 118-2 Postgadolinium T1 MRI of a large cystic left frontal glioblastoma.

the DNA repair enzyme O⁶-methylguanine-DNA methyltransferase (MGMT) are relatively resistant to temozolomide and have a worse prognosis compared to those whose tumors contain low levels of MGMT as a result of silencing of the MGMT gene by promoter hypermethylation. Implantation of biodegradable polymers containing the chemotherapeutic agent carmustine into the tumor bed after resection of the tumor also produces a modest improvement in survival.

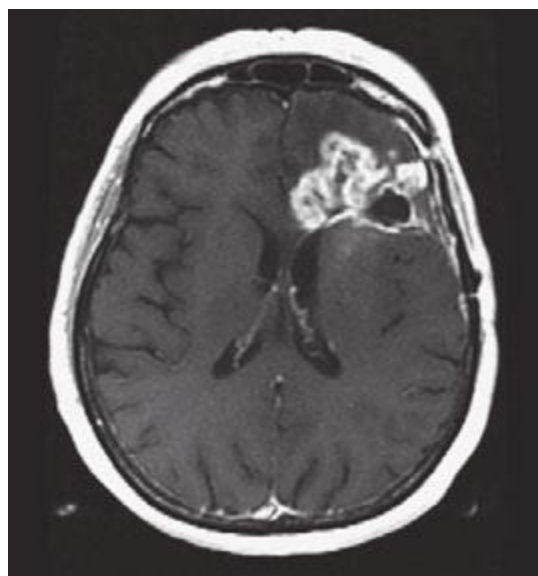
Despite optimal therapy, glioblastomas invariably recur. Treatment options for recurrent disease may include reoperation, carmustine wafers, and alternate chemotherapeutic regimens. Reirradiation is rarely helpful. Bevacizumab, a humanized vascular endothelial growth factor (VEGF) monoclonal antibody, has activity in recurrent glioblastoma, increasing progression-free survival and reducing peritumoral edema and glucocorticoid use (Fig. 118-3). Treatment decisions for patients with recurrent glioblastoma must be made on an individual basis, taking into consideration such factors as previous therapy, time to relapse, performance status, and quality of life. Whenever feasible, patients with recurrent disease should be enrolled in clinical trials. Novel therapies undergoing evaluation in patients with glioblastoma include targeted molecular agents directed at receptor tyrosine kinases and signal transduction pathways; antiangiogenic agents, especially those directed at the VEGF receptors; chemotherapeutic agents that cross the blood-brain barrier more effectively than currently available drugs; gene therapy; immunotherapy; and infusion of radiolabeled drugs and targeted toxins into the tumor and surrounding brain by means of convection-enhanced delivery.

The most important adverse prognostic factors in patients with high-grade astrocytomas are older age, histologic features of glioblastoma, poor Karnofsky performance status, and unresectable tumor. Patients whose tumor contains an unmethylated MGMT promoter resulting in the presence of the repair enzyme in tumor cells and resistance to temozolomide also have a worse prognosis.

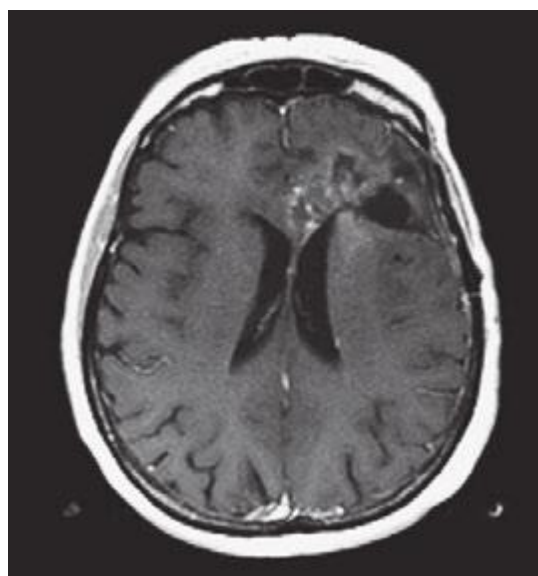
Gliomatosis Cerebri Rarely, patients may present with a highly infiltrating, nonenhancing tumor of variable histologic grade involving more than two lobes of the brain. These tumors may be indolent initially, but will eventually behave aggressively and have a poor outcome. Treatment involves RT and temozolomide chemotherapy.

OLIGODENDROGLIOMA

Oligodendrogliomas account for approximately 15–20% of gliomas. They are classified by the WHO into well-differentiated oligodendrogliomas (grade II) or anaplastic oligodendrogliomas (AOs) (grade III). Tumors with oligodendroglial components have distinctive pathologic



A



B

FIGURE 118-3 Postgadolinium T1 MRI of a recurrent glioblastoma before (A) and after (B) administration of bevacizumab. Note the decreased enhancement and mass effect.

features such as perinuclear clearing—giving rise to a “fried-egg” appearance—and a reticular pattern of blood vessel growth. Some tumors have both an oligodendroglial as well as an astrocytic component. These mixed tumors, or oligoastrocytomas (OAs), are also classified into well-differentiated OA (grade II) or anaplastic oligoastrocytomas (AOAs) (grade III).

Grade II oligodendrogliomas and OAs are generally more responsive to therapy and have a better prognosis than pure astrocytic tumors. These tumors present similarly to grade II astrocytomas in young adults. The tumors are nonenhancing and often partially calcified. They should be treated with surgery and, if necessary, RT and chemotherapy. Patients with oligodendrogliomas have a median survival in excess of 10 years.

AOs and AOAs present in the fourth and fifth decades as variably enhancing tumors. They are more responsive to therapy than grade III astrocytomas. Co-deletion of chromosomes 1p and 19q, mediated by an unbalanced translocation of 19p to 1q, occurs in 61–89% of patients with AO and 14–20% of patients with AOA. Tumors with the 1p and 19q co-deletion are particularly sensitive to chemotherapy with procarbazine, lomustine (cyclohexylchloroethylnitrosourea [CCNU]), and vincristine (PCV) or temozolomide, as well as to RT. Median

survival of patients with AO or AOA is approximately 3–6 years, but those with co-deleted tumors can have a median survival of 10–14 years if treated with RT and chemotherapy.

EPENDYMOMAS

Ependymomas are tumors derived from ependymal cells that line the ventricular surface. They account for approximately 5% of childhood tumors and frequently arise from the wall of the fourth ventricle in the posterior fossa. Although adults can have intracranial ependymomas, they occur more commonly in the spine, especially in the filum terminale of the spinal cord where they have a myxopapillary histology. Ependymomas that can be completely resected are potentially curable. Partially resected ependymomas will recur and require irradiation. The less common anaplastic ependymoma is more aggressive and is treated with resection and RT; chemotherapy has limited efficacy. Subependymomas are slow-growing benign lesions arising in the wall of ventricles that often do not require treatment.

OTHER LESS COMMON GLIOMAS

Gangliogliomas and pleomorphic xanthoastrocytomas occur in young adults. They behave as more indolent forms of grade II gliomas and are treated in the same way. Brainstem gliomas usually occur in children or young adults. Despite treatment with RT and chemotherapy, the prognosis is poor, with a median survival of only 1 year. Gliosarcomas contain both an astrocytic as well as a sarcomatous component and are treated in the same way as glioblastomas.

PRIMARY CENTRAL NERVOUS SYSTEM LYMPHOMA

Primary central nervous system lymphoma (PCNSL) is a rare non-Hodgkin lymphoma accounting for less than 3% of primary brain tumors. For unclear reasons, its incidence is increasing, particularly in immunocompetent individuals.

PCNSL in immunocompetent patients usually consists of a diffuse large B cell lymphoma. PCNSL may also occur in immunocompromised patients, usually those infected with the human immunodeficiency virus (HIV) or organ transplant recipients on immunosuppressive therapy. PCNSL in immunocompromised patients is typically large cell with immunoblastic and more aggressive features. These patients are usually severely immunocompromised, with CD4 counts of less than 50/mL. The Epstein-Barr virus (EBV) frequently plays an important role in the pathogenesis of HIV-related PCNSL.

Immunocompetent patients with PCNSL are older (median 60 years) compared to patients with HIV-related PCNSL (median 31 years). PCNSL usually presents as a mass lesion, with neuropsychiatric symptoms, symptoms of increased intracranial pressure, lateralizing signs, or seizures.

On contrast-enhanced MRI, PCNSL usually appears as a densely enhancing tumor (Fig. 118-4). Immunocompetent patients have solitary lesions more often than immunosuppressed patients. Frequently there is involvement of the basal ganglia, corpus callosum, or periventricular region. Although the imaging features are often characteristic, PCNSL can sometimes be difficult to differentiate from high-grade gliomas, infections, or demyelination. Stereotactic biopsy is necessary to obtain a histologic diagnosis. Whenever possible, glucocorticoids should be withheld until after the biopsy has been obtained because they have a cytolytic effect on lymphoma cells and may lead to nondiagnostic tissue. In addition, patients should be tested for HIV and the extent of disease should be assessed by performing PET or CT of the body, MRI of the spine, CSF analysis, and slit-lamp examination of the eye. Bone marrow biopsy and testicular ultrasound are occasionally performed.

TREATMENT

PRIMARY CENTRAL NERVOUS SYSTEM LYMPHOMA

PCNSL is more sensitive to glucocorticoids, chemotherapy, and RT than other primary brain tumors. Durable complete responses and long-term survival are possible with these treatments. High-dose methotrexate, a folate antagonist that interrupts DNA synthesis, produces response rates ranging from 35–80% and median

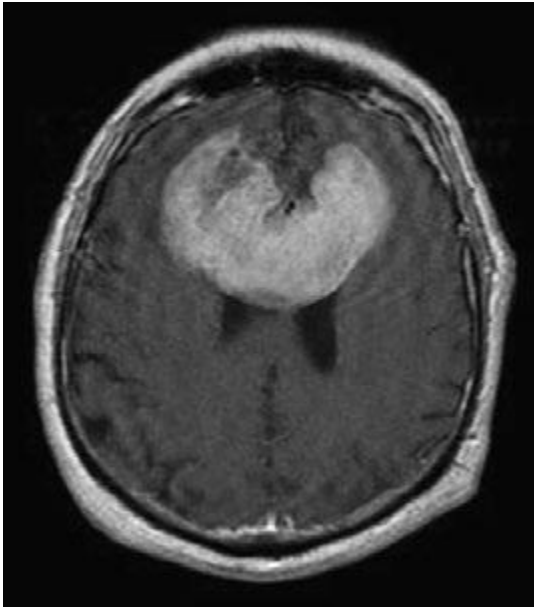


FIGURE 118-4 Postgadolinium T1 MRI demonstrating a large bifrontal primary central nervous system lymphoma (PCNSL). The periventricular location and diffuse enhancement pattern are characteristic of lymphoma.

survival of up to 50 months. The combination of methotrexate with other chemotherapeutic agents such as cytarabine increases the response rate to 70–100%. The addition of whole-brain RT to methotrexate-based chemotherapy prolongs progression-free survival but not overall survival. Furthermore, RT is associated with delayed neurotoxicity, especially in patients over the age of 60 years. As a result, full-dose RT is frequently omitted, but there may be a role for reduced-dose RT. The anti-CD20 monoclonal antibody rituximab has activity in PCNSL and is often incorporated into the chemotherapy regimen. For some patients, high-dose chemotherapy with autologous stem cell rescue may offer the best chance of preventing relapse. At least 50% of patients will eventually develop recurrent disease. Treatment options include RT for patients who have not had prior irradiation, re-treatment with methotrexate, as well as other agents such as temozolomide, rituximab, procarbazine, topotecan, and pemetrexed. High-dose chemotherapy with autologous stem cell rescue may have a role in selected patients with relapsed disease.

PCNSL IN IMMUNOCOMPROMISED PATIENTS

PCNSL in immunocompromised patients often produces multiple ring-enhancing lesions that can be difficult to differentiate from metastases and infections such as toxoplasmosis. The diagnosis is usually established by examination of the CSF for cytology and EBV DNA, toxoplasmosis serologic testing, brain PET imaging for hypermetabolism of the lesions consistent with tumor instead of infection, and, if necessary, brain biopsy. Since the advent of highly active antiretroviral drugs, the incidence of HIV-related PCNSL has declined. These patients may be treated with whole-brain RT, high-dose methotrexate, and initiation of highly active antiretroviral therapy. In organ transplant recipients, reduction of immunosuppression may improve outcome.

MEDULLOBLASTOMAS

Medulloblastomas are the most common malignant brain tumor of childhood, accounting for approximately 20% of all primary CNS tumors among children. They arise from granule cell progenitors or from multipotent progenitors from the ventricular zone. Approximately 5% of children have inherited disorders with germline mutations of genes that predispose to the development of medulloblastoma. Gorlin syndrome, the most common of these inherited disorders, is due to mutations in the patched-1 (*PTCH-1*) gene, a key

component in the sonic hedgehog pathway. Turcot syndrome, caused by mutations in the adenomatous polyposis coli (*APC*) gene and familial adenomatous polyposis, has also been associated with an increased incidence of medulloblastoma. Histologically, medulloblastomas are highly cellular tumors with abundant dark staining, round nuclei, and rosette formation (Homer-Wright rosettes). They present with headache, ataxia, and signs of brainstem involvement. On MRI they appear as densely enhancing tumors in the posterior fossa, sometimes associated with hydrocephalus. Seeding of the CSF is common. Treatment involves maximal surgical resection, craniospinal irradiation, and chemotherapy with agents such as cisplatin, lomustine, cyclophosphamide, and vincristine. Approximately 70% of patients have long-term survival but usually at the cost of significant neurocognitive impairment. A major goal of current research is to improve survival while minimizing long-term complications.

PINEAL REGION TUMORS

A large number of tumors can arise in the region of the pineal gland. These typically present with headache, visual symptoms, and hydrocephalus. Patients may have Parinaud syndrome characterized by impaired upgaze and accommodation. Some pineal tumors such as pineocytomas and benign teratomas can be treated simply by surgical resection. Germinomas respond to irradiation, whereas pineoblastomas and malignant germ cell tumors require craniospinal radiation and chemotherapy.

EXTRINSIC “BENIGN” TUMORS

MENINGIOMAS

Meningiomas are diagnosed with increasing frequency as more people undergo neuroimaging for various indications. They are now the most common primary brain tumor, accounting for approximately 35% of the total. Their incidence increases with age. They tend to be more common in women and in patients with neurofibromatosis type 2. They also occur more commonly in patients with a past history of cranial irradiation.

Meningiomas arise from the dura mater and are composed of neoplastic meningothelial (arachnoidal cap) cells. They are most commonly located over the cerebral convexities, especially adjacent to the sagittal sinus, but can also occur in the skull base and along the dorsum of the spinal cord. Meningiomas are classified by the WHO into three histologic grades of increasing aggressiveness: grade I (benign), grade II (atypical), and grade III (malignant).

Many meningiomas are found incidentally following neuroimaging for unrelated reasons. They can also present with headaches, seizures, or focal neurologic deficits. On imaging studies they have a characteristic appearance usually consisting of a partially calcified, densely enhancing extraaxial tumor arising from the dura (Fig. 118-5). Occasionally they may have a dural tail, consisting of thickened, enhanced dura extending like a tail from the mass. The main differential diagnosis of meningioma is a dural metastasis.

If the meningioma is small and asymptomatic, no intervention is necessary and the lesion can be observed with serial MRI studies. Larger, symptomatic lesions should be resected. If complete resection is achieved, the patient is cured. Incompletely resected tumors tend to recur, although the rate of recurrence can be very slow with grade I tumors. Tumors that cannot be resected, or can only be partially removed, may benefit from treatment with external-beam RT or stereotactic radiosurgery (SRS). These treatments may also be helpful in patients whose tumor has recurred after surgery. Hormonal therapy and chemotherapy are currently unproven.

Rarer tumors that resemble meningiomas include hemangiopericytomas and solitary fibrous tumors. These are treated with surgery and RT but have a higher propensity to recur locally or metastasize systemically.

SCHWANNOMAS

These are generally benign tumors arising from the Schwann cells of cranial and spinal nerve roots. The most common schwannomas, termed *vestibular schwannomas* or *acoustic neuromas*, arise from the

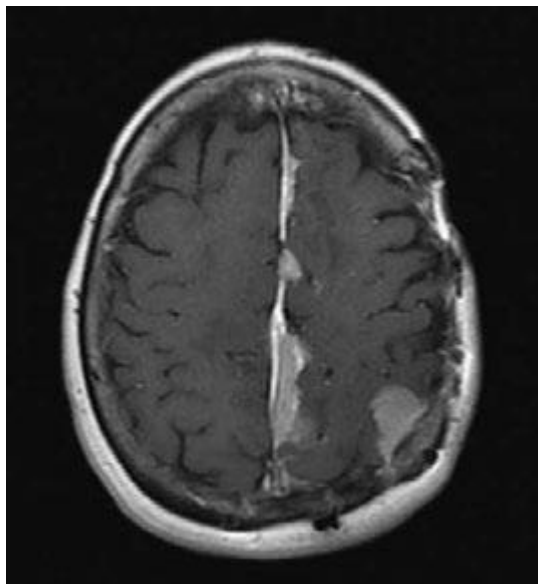


FIGURE 118-5 Postgadolinium T1 MRI demonstrating multiple meningiomas along the falx and left parietal cortex.

vestibular portion of the eighth cranial nerve and account for approximately 9% of primary brain tumors. Patients with neurofibromatosis type 2 have a high incidence of vestibular schwannomas that are frequently bilateral. Schwannomas arising from other cranial nerves, such as the trigeminal nerve (cranial nerve V), occur with much lower frequency. Neurofibromatosis type 1 is associated with an increased incidence of schwannomas of the spinal nerve roots.

Vestibular schwannomas may be found incidentally on neuroimaging or present with progressive unilateral hearing loss, dizziness, tinnitus, or less commonly, symptoms resulting from compression of the brainstem and cerebellum. On MRI they appear as densely enhancing lesions, enlarging the internal auditory canal and often extending into the cerebellopontine angle (Fig. 118-6). The differential diagnosis includes meningioma. Very small, asymptomatic lesions can be observed with serial MRIs. Larger lesions should be treated with surgery or SRS. The optimal treatment will depend on the size of the tumor, symptoms, and the patient's preference. In patients with small

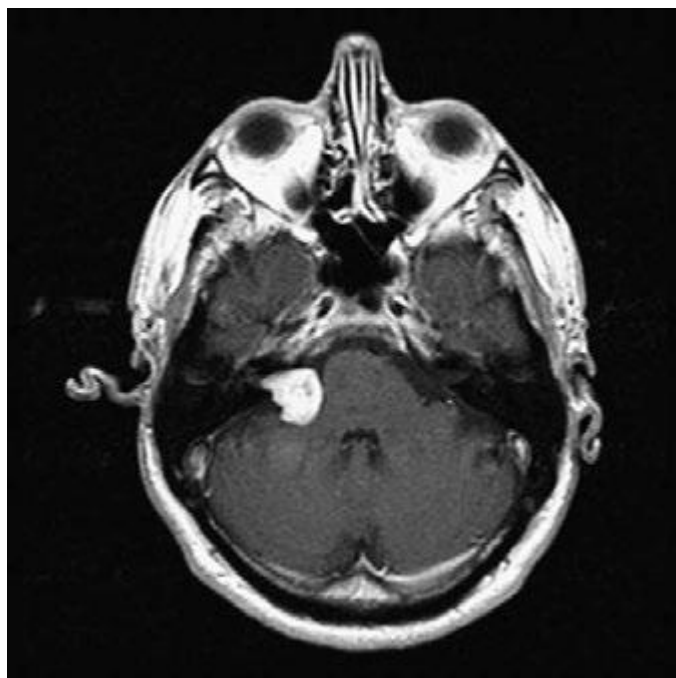


FIGURE 118-6 Postgadolinium MRI of a right vestibular schwannoma. The tumor can be seen to involve the internal auditory canal.

vestibular schwannomas and relatively intact hearing, early surgical intervention increases the chance of preserving hearing.

PITUITARY TUMORS (CHAP. 401e)

These account for approximately 9% of primary brain tumors. They can be divided into functioning and nonfunctioning tumors. Functioning tumors are usually microadenomas (<1 cm in diameter) that secrete hormones and produce specific endocrine syndromes (e.g., acromegaly for growth hormone-secreting tumors, Cushing syndrome for adrenocorticotropic hormone [ACTH]-secreting tumors, and galactorrhea, amenorrhea, and infertility for prolactin-secreting tumors). Nonfunctioning pituitary tumors tend to be macroadenomas (>1 cm) that produce symptoms by mass effect, giving rise to headaches, visual impairment (such as bitemporal hemianopia), and hypopituitarism. Prolactin-secreting tumors respond well to dopamine agonists such as bromocriptine and cabergoline. Other pituitary tumors usually require treatment with surgery and sometimes RT or radiosurgery and hormonal therapy.

CRANIOPHARYNGIOMAS

Craniopharyngiomas are rare, usually suprasellar, partially calcified, solid, or mixed solid-cystic benign tumors that arise from remnants of Rathke's pouch. They have a bimodal distribution, occurring predominantly in children but also between the ages of 55 and 65 years. They present with headaches, visual impairment, and impaired growth in children and hypopituitarism in adults. Treatment involves surgery, RT, or a combination of the two.

OTHER BENIGN TUMORS

Dysembryoplastic Neuroepithelial Tumors (DNTs) These are benign, supratentorial tumors, usually in the temporal lobe. They typically occur in children and young adults with a long-standing history of seizures. Surgical resection is curative.

Epidermoid Cysts These consist of squamous epithelium surrounding a keratin-filled cyst. They are usually found in the cerebellopontine angle and the intrasellar and suprasellar regions. They may present with headaches, cranial nerve abnormalities, seizures, or hydrocephalus. Imaging studies demonstrate extraaxial lesions with characteristics that are similar to CSF but have restricted diffusion. Treatment involves surgical resection.

Dermoid Cysts Like epidermoid cysts, dermoid cysts arise from epithelial cells that are retained during closure of the neural tube. They contain both epidermal and dermal structures such as hair follicles, sweat glands, and sebaceous glands. Unlike epidermoid cysts, these tumors usually have a midline location. They occur most frequently in the posterior fossa, especially the vermis, fourth ventricle, and suprasellar cistern. Radiographically, dermoid cysts resemble lipomas, demonstrating T1 hyperintensity and variable signal on T2. Symptomatic dermoid cysts can be treated with surgery.

Colloid Cysts These usually arise in the anterior third ventricle and may present with headaches, hydrocephalus, and, very rarely, sudden death. Surgical resection is curative, or a third ventriculostomy may relieve the obstructive hydrocephalus and be sufficient therapy.

NEUROCUTANEOUS SYNDROMES (PHAKOMATOSSES)

A number of genetic disorders are characterized by cutaneous lesions and an increased risk of brain tumors. Most of these disorders have an autosomal dominant inheritance with variable penetrance.

NEUROFIBROMATOSIS TYPE 1 (NF1) (von RECKLINGHAUSEN'S DISEASE)

NF1 is an autosomal dominant disorder with an incidence of approximately 1 in 2600–3000. Approximately one-half the cases are familial; the remainder are caused by new mutations arising in patients with unaffected parents. The *NF1* gene on chromosome 17q11.2 encodes a protein, neurofibromin, a guanosine triphosphatase (GTPase)-activating protein (GAP) that modulates signaling through the ras pathway. Mutations of *NF1* result in a large number of nervous system tumors

604 including neurofibromas, plexiform neurofibromas, optic nerve gliomas, astrocytomas, and meningiomas. In addition to neurofibromas, which appear as multiple, soft, rubbery cutaneous tumors, other cutaneous manifestations of NF1 include café-au-lait spots and axillary freckling. NF1 is also associated with hamartomas of the iris termed Lisch nodules, pheochromocytomas, pseudoarthrosis of the tibia, scoliosis, epilepsy, and mental retardation.

NEUROFIBROMATOSIS TYPE 2 (NF2)

NF2 is less common than NF1, with an incidence of 1 in 25,000–40,000. It is an autosomal dominant disorder with full penetrance. As with NF1, approximately one-half the cases arise from new mutations. The *NF2* gene on 22q encodes a cytoskeletal protein, merlin (moesin, ezrin, radixin-like protein) that functions as a tumor suppressor. NF2 is characterized by bilateral vestibular schwannomas in over 90% of patients, multiple meningiomas, and spinal ependymomas and astrocytomas. Treatment of bilateral vestibular schwannomas can be challenging because the goal is to preserve hearing for as long as possible. These patients may also have diffuse schwannomatosis that may affect the cranial, spinal, or peripheral nerves; posterior subcapsular lens opacities; and retinal hamartomas.

TUBEROUS SCLEROSIS (BOURNEVILLE DISEASE)

This is an autosomal dominant disorder with an incidence of approximately 1 in 5000–10,000 live births. It is caused by mutations in either the *TSC1* gene, which maps to chromosome 9q34 and encodes a protein termed hamartin, or the *TSC2* gene, which maps to chromosome 16p13.3 and encodes the protein tuberlin. Hamartin forms a complex with tuberlin, which inhibits cellular signaling through the mTOR, and acts as a negative regulator of the cell cycle. Patients with tuberous sclerosis may have seizures, mental retardation, adenoma sebaceum (facial angiofibromas), shagreen patch, hypomelanotic macules, periungual fibromas, renal angiomyolipomas, and cardiac rhabdomyomas. These patients have an increased incidence of subependymal nodules, cortical tubers, and subependymal giant-cell astrocytomas (SEGA). Patients frequently require anticonvulsants for seizures. SEGAs do not always require therapeutic intervention, but the most effective therapy is with the mTOR inhibitors sirolimus or everolimus, which often decrease seizures as well as SEGA size.

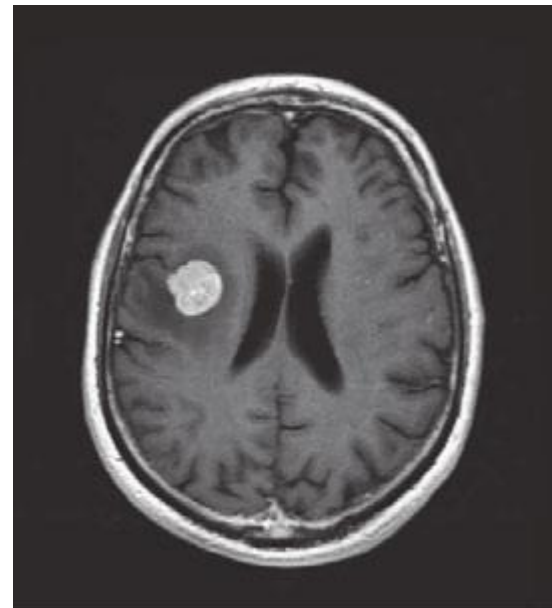
TUMORS METASTATIC TO THE BRAIN

Brain metastases arise from hematogenous spread and frequently either arise from a lung primary or are associated with pulmonary metastases. Most metastases develop at the gray matter–white matter junction in the watershed distribution of the brain where intravascular tumor cells lodge in terminal arterioles. The distribution of metastases in the brain approximates the proportion of blood flow such that about 85% of all metastases are supratentorial and 15% occur in the posterior fossa. The most common sources of brain metastases are lung and breast carcinomas; melanoma has the greatest propensity to metastasize to the brain, being found in 80% of patients at autopsy (Table 118-3). Other tumor

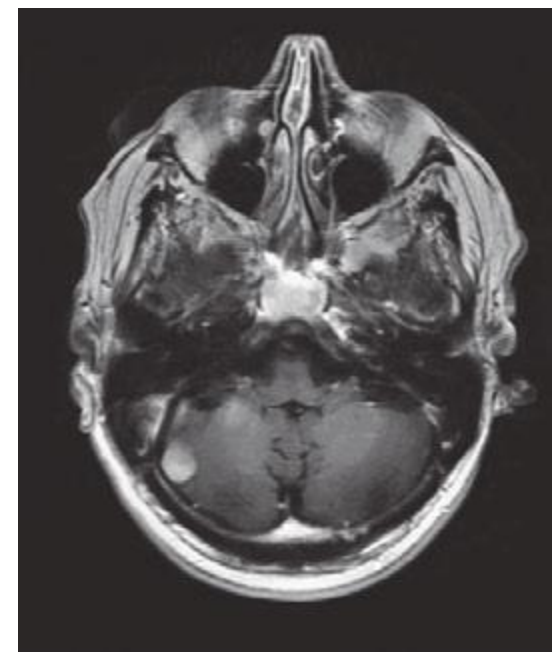
types such as ovarian and esophageal carcinoma rarely metastasize to the brain. Prostate and breast cancer also have a propensity to metastasize to the dura and can mimic meningioma. Leptomeningeal metastases are common from hematologic malignancies and also breast and lung cancers. Spinal cord compression primarily arises in patients with prostate and breast cancer, tumors with a strong propensity to metastasize to the axial skeleton.

DIAGNOSIS OF METASTASES

Brain metastases are best visualized on MRI, where they usually appear as well-circumscribed lesions (Fig. 118-7). The amount of perilesional edema can be highly variable, with large lesions causing minimal edema and sometimes very small lesions causing extensive edema. Enhancement may be in a ring pattern or diffuse. Occasionally,



A



B

FIGURE 118-7 Postgadolinium T1 MRI of multiple brain metastases from non-small-cell lung cancer involving the right frontal (A) and right cerebellar (B) hemispheres. Note the diffuse enhancement pattern and absence of central necrosis.

TABLE 118-3 FREQUENCY OF NERVOUS SYSTEM METASTASES BY COMMON PRIMARY TUMORS			
	Brain %	LM %	ESCC %
Lung	41	17	15
Breast	19	57	22
Melanoma	10	12	4
Prostate	1	1	10
GIT	7	—	5
Renal	3	2	7
Lymphoma	<1	10	10
Sarcoma	7	1	9
Other	11	—	18

Abbreviations: ESCC, epidural spinal cord compression; GIT, gastrointestinal tract; LM, leptomeningeal metastases.

intracranial metastases will hemorrhage; although melanoma, thyroid, and kidney cancer have the greatest propensity to hemorrhage, the most common cause of a hemorrhagic metastasis is lung cancer because it accounts for the majority of brain metastases. The radiographic appearance of brain metastasis is nonspecific, and similar-appearing lesions can occur with infection including brain abscesses and also with demyelinating lesions, sarcoidosis, radiation necrosis in a previously treated patient, or a primary brain tumor that may be a second malignancy in a patient with systemic cancer. However, biopsy is rarely necessary for diagnosis in most patients because imaging alone in the appropriate clinical situation usually suffices. This is straightforward for the majority of patients with brain metastases because they have a known systemic cancer. However, in approximately 10% of patients, a systemic cancer may present with a brain metastasis, and if there is not an easily accessible systemic site to biopsy, then a brain lesion must be removed for diagnostic purposes.

TREATMENT TUMORS METASTATIC TO THE BRAIN

DEFINITIVE TREATMENT

The number and location of brain metastases often determine the therapeutic options. The patient's overall condition and the current or potential control of the systemic disease are also major determinants. Brain metastases are single in approximately one-half of patients and multiple in the other half.

RADIATION THERAPY

The standard treatment for brain metastases has been whole-brain radiotherapy (WBRT) usually administered to a total dose of 3000 cGy in 10 fractions. This affords rapid palliation, and approximately 80% of patients improve with glucocorticoids and RT. However, it is not curative. Median survival is only 4–6 months. More recently, SRS delivered through a variety of techniques including the gamma knife, linear accelerator, proton beam, and CyberKnife all can deliver highly focused doses of RT, usually in a single fraction. SRS can effectively sterilize the visible lesions and afford local disease control in 80–90% of patients. In addition, there are some patients who have clearly been cured of their brain metastases using SRS, whereas this is distinctly rare with WBRT. However, SRS can be used only for lesions 3 cm or less in diameter and should be confined to patients with only one to three metastases. The addition of WBRT to SRS improves disease control in the nervous system but does not prolong survival.

SURGERY

Randomized controlled trials have demonstrated that surgical extirpation of a single brain metastasis followed by WBRT is superior to WBRT alone. Removal of two lesions or a single symptomatic mass, particularly if compressing the ventricular system, can also be useful. This is particularly useful in patients who have highly radioresistant lesions such as renal carcinoma. Surgical resection can afford rapid symptomatic improvement and prolonged survival. WBRT administered after complete resection of a brain metastasis improves disease control but does not prolong survival.

CHEMOTHERAPY

Chemotherapy is rarely useful for brain metastases. Metastases from certain tumor types that are highly chemosensitive, such as germ cell tumors or small-cell lung cancer, may respond to chemotherapeutic regimens chosen according to the underlying malignancy. Increasingly, there are data demonstrating responsiveness of brain metastases to chemotherapy including small molecule–targeted therapy when the lesion possesses the target. This has been best illustrated in patients with lung cancer harboring EGFR mutations that sensitize them to EGFR inhibitors. Antiangiogenic agents such as bevacizumab may also prove efficacious in the treatment of CNS metastases.

LEPTOMENINGEAL METASTASES

Leptomeningeal metastases are also identified as carcinomatous meningitis, meningeal carcinomatosis, or in the case of specific tumors, leukemic or lymphomatous meningitis. Among the hematologic malignancies, acute leukemia is the most common to metastasize to the subarachnoid space, and in lymphomas the aggressive diffuse lymphomas can metastasize to the subarachnoid space frequently as well. Among solid tumors, breast and lung carcinomas and melanoma most frequently spread in this fashion. Tumor cells reach the subarachnoid space via the arterial circulation or occasionally through retrograde flow in venous systems that drain metastases along the bony spine or cranium. In addition, leptomeningeal metastases may develop as a direct consequence of prior brain metastases and can develop in almost 40% of patients who have a metastasis resected from the cerebellum.

CLINICAL FEATURES

Leptomeningeal metastases are characterized clinically by multilevel symptoms and signs along the neuraxis. Combinations of lumbar and cervical radiculopathies, cranial neuropathies, seizures, confusion, and encephalopathy from hydrocephalus or raised intracranial pressure can be present. Focal deficits such as hemiparesis or aphasia are rarely due to leptomeningeal metastases unless there is direct brain infiltration, and they are more often associated with coexisting brain lesions. New-onset limb pain in patients with breast cancer, lung cancer, or melanoma should prompt consideration of leptomeningeal spread.

LABORATORY AND IMAGING DIAGNOSIS

Leptomeningeal metastases are particularly challenging to diagnose because identification of tumor cells in the subarachnoid compartment may be elusive. MRI can be definitive in patients when there are clear tumor nodules adherent to the cauda equina or spinal cord, enhancing cranial nerves, or subarachnoid enhancement on brain imaging (Fig. 118-8). Imaging is diagnostic in approximately 75% of patients and is more often positive in patients with solid tumors. Demonstration of tumor cells in the CSF is definitive and often considered the gold standard. However, CSF cytologic examination is positive in only 50% of patients on the first lumbar puncture and still misses 10% after three CSF samples. CSF cytologic examination is most useful in hematologic malignancies. Accompanying CSF abnormalities include an elevated protein concentration and an elevated white count. Hypoglycorrhachia is noted in less than 25% of patients but is useful when present. Identification of tumor markers or molecular confirmation of clonal proliferation with techniques such as flow cytometry within the CSF can also be definitive when present. Tumor markers are usually specific to solid tumors, and chromosomal or molecular markers are most useful in patients with hematologic malignancies. New technologies, such as rare cell capture, may enhance identification of tumor cells in the CSF.

TREATMENT LEPTOMENINGEAL METASTASES

The treatment of leptomeningeal metastasis is palliative because there is no curative therapy. RT to the symptomatically involved areas, such as skull base for cranial neuropathy, can relieve pain and sometimes improve function. Whole-neuraxis RT has extensive toxicity with myelosuppression and gastrointestinal irritation as well as limited effectiveness. Systemic chemotherapy with agents that can penetrate the blood-CSF barrier may be helpful. Alternatively, intrathecal chemotherapy can be effective, particularly in hematologic malignancies. This is optimally delivered through an intraventricular cannula (Ommaya reservoir) rather than by lumbar puncture. Few drugs can be delivered safely into the subarachnoid space, and they have a limited spectrum of antitumor activity, perhaps accounting for the relatively poor response to this approach. In addition, impaired CSF flow dynamics can compromise intrathecal drug delivery. Surgery has a limited role in the treatment of leptomeningeal metastasis, but placement of a ventriculoperitoneal shunt can relieve raised intracranial pressure. However, it compromises delivery of chemotherapy into the CSF.



A



B

FIGURE 118-8 Postgadolinium MRI images of extensive leptomeningeal metastases from breast cancer. Nodules along the dorsal surface of the spinal cord (A) and cauda equina (B) are seen.

EPIDURAL METASTASIS

Epidural metastasis occurs in 3–5% of patients with a systemic malignancy and causes neurologic compromise by compressing the spinal cord or cauda equina. The most common cancers that metastasize to the epidural space are those malignancies that spread to bone, such as breast and prostate. Lymphoma can cause bone involvement and compression, but it can also invade the intervertebral foramina and cause spinal cord compression without bone destruction. The thoracic spine is affected most commonly, followed by the lumbar and then cervical spine.

CLINICAL FEATURES

Back pain is the presenting symptom of epidural metastasis in virtually all patients; the pain may precede neurologic findings by weeks or months. The pain is usually exacerbated by lying down; by contrast,

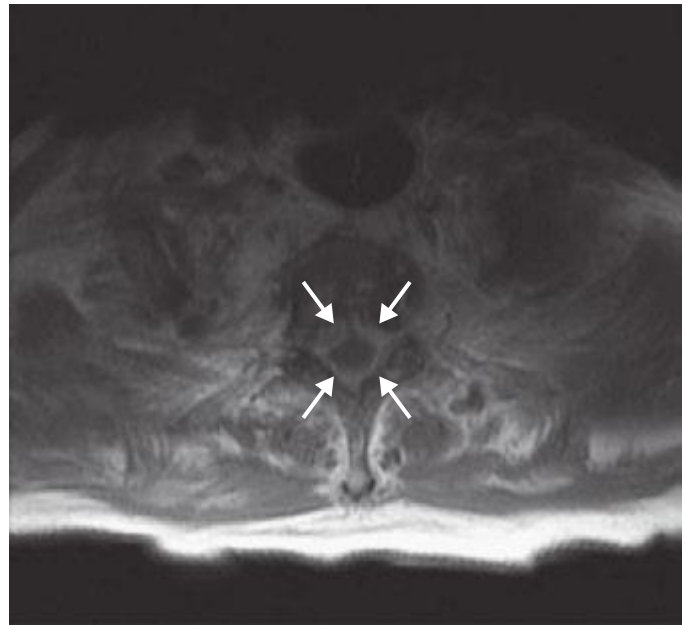


FIGURE 118-9 Postgadolinium T1 MRI showing circumferential epidural tumor around the thoracic spinal cord from esophageal cancer.

arthritic pain is often relieved by recumbency. Leg weakness is seen in about 50% of patients, as is sensory dysfunction. Sphincter problems are present in about 25% of patients at diagnosis.

DIAGNOSIS

Diagnosis is established by imaging, with MRI of the complete spine being the best test (Fig. 118-9). Contrast is not needed to identify spinal or epidural lesions. Any patient with cancer who has severe back pain should undergo an MRI. Plain films, bone scans, or even CT scans may show bone metastases, but only MRI can reliably delineate epidural tumor. For patients unable to have an MRI, CT myelography should be performed to outline the epidural space. The differential diagnosis of epidural tumor includes epidural abscess, acute or chronic hematomas, and rarely, extramedullary hematopoiesis.

TREATMENT EPIDURAL METASTASIS

Epidural metastasis requires immediate treatment. A randomized controlled trial demonstrated the superiority of surgical resection followed by RT compared to RT alone. However, patients must be able to tolerate surgery, and the surgical procedure of choice is a complete removal of the mass, which is typically anterior to the spinal canal, necessitating an extensive approach and resection. Otherwise, RT is the mainstay of treatment and can be used for patients with radiosensitive tumors, such as lymphoma, or for those unable to undergo surgery. Chemotherapy is rarely used for epidural metastasis unless the patient has minimal to no neurologic deficit and a highly chemosensitive tumor such as lymphoma or germinoma. Patients generally fare well if treated before there is severe neurologic deficit. Recovery after paraparesis is better after surgery than with RT alone, but survival is often short due to widespread metastatic tumor.

NEUROLOGIC TOXICITY OF THERAPY

TOXICITY FROM RADIOTHERAPY

RT can cause a variety of toxicities in the CNS. These are usually described based on their relationship in time to the administration of RT: acute (occurring within days of RT), early delayed (months), or late delayed (years). In general, the acute and early delayed syndromes

resolve and do not result in persistent deficits, whereas the late delayed toxicities are usually permanent and sometimes progressive.

Acute Toxicity Acute cerebral toxicity usually occurs during RT to the brain. RT can cause a transient disruption of the blood-brain barrier, resulting in increased edema and elevated intracranial pressure. This is usually manifest as headache, lethargy, nausea, and vomiting and can be both prevented and treated with the administration of glucocorticoids. There is no acute RT toxicity that affects the spinal cord.

Early Delayed Toxicity Early delayed toxicity is usually apparent weeks to months after completion of cranial irradiation and is likely due to focal demyelination. Clinically it may be asymptomatic or take the form of worsening or reappearance of a preexisting neurologic deficit. At times a contrast-enhancing lesion can be seen on MRI/CT that can mimic the tumor for which the patient received the RT. For patients with a malignant glioma, this has been described as “pseudoprogression” because it mimics tumor recurrence on MRI but actually represents inflammation and necrotic debris engendered by effective therapy. This is seen with increased frequency when chemotherapy, particularly temozolomide, is given concurrently with RT. Pseudoprogression can resolve on its own or, if very symptomatic, may require resection. A rare form of early delayed toxicity is the somnolence syndrome that occurs primarily in children and is characterized by marked sleepiness.

In the spinal cord, early delayed RT toxicity is manifest as a Lhermitte symptom with paresthesias of the limbs or along the spine when the patient flexes the neck. Although frightening, it is benign, resolves on its own, and does not portend more serious problems.

Late Delayed Toxicity Late delayed toxicities are the most serious because they are often irreversible and cause severe neurologic deficits. In the brain, late toxicities can take several forms, the most common of which include radiation necrosis and leukoencephalopathy. Radiation necrosis is a focal mass of necrotic tissue that is contrast enhancing on CT/MRI and may be associated with significant edema. This may appear identical to pseudoprogression but is seen months to years after RT and is always symptomatic. Clinical symptoms and signs include seizure and lateralizing findings referable to the location of the necrotic mass. The necrosis is caused by the effect of RT on cerebral vasculature with resultant fibrinoid necrosis and occlusion of the blood vessels. It can mimic tumor radiographically, but unlike tumor, it is typically hypometabolic on a PET scan and has reduced perfusion on perfusion MR sequences. It may require resection for diagnosis and treatment unless it can be managed with glucocorticoids. There are rare reports of improvement with hyperbaric oxygen or anticoagulation, but the usefulness of these approaches is questionable.

Leukoencephalopathy is seen most commonly after WBRT as opposed to focal RT. On T2 or FLAIR MR sequences, there is diffuse increased signal seen throughout the hemispheric white matter, often bilaterally and symmetrically. There tends to be a periventricular predominance that may be associated with atrophy and ventricular enlargement. Clinically, patients develop cognitive impairment, gait disorder, and later urinary incontinence, all of which can progress over time. These symptoms mimic those of normal pressure hydrocephalus, and placement of a ventriculoperitoneal shunt can improve function in some patients but does not reverse the deficits completely. Increased age is a risk factor for leukoencephalopathy but not for radiation necrosis. Necrosis appears to depend on an as yet unidentified predisposition.

Other late neurologic toxicities include endocrine dysfunction if the pituitary or hypothalamus was included in the RT port. An RT-induced neoplasm can occur many years after therapeutic RT for either a prior CNS tumor or a head and neck cancer; accurate diagnosis requires surgical resection or biopsy. In addition, RT causes accelerated atherosclerosis, which can cause stroke either from intracranial vascular disease or carotid plaque from neck irradiation.

The peripheral nervous system is relatively resistant to RT toxicities. Peripheral nerves are rarely affected by RT, but the plexus is more

TABLE 118-4 NEUROLOGIC SIGNS CAUSED BY AGENTS COMMONLY USED IN PATIENTS WITH CANCER

Acute encephalopathy (delirium)	Seizures
Methotrexate (high-dose IV, IT)	Methotrexate
Cisplatin	Etoposide (high-dose)
Vincristine	Cisplatin
Asparaginase	Vincristine
Procarbazine	Asparaginase
5-Fluorouracil (± levamisole)	Nitrogen mustard
Cytarabine (high-dose)	Carmustine
Nitrosoureas (high-dose or arterial)	Dacarbazine (intraarterial or high-dose)
Ifosfamide	Busulfan (high-dose)
Etoposide (high-dose)	Myelopathy (intrathecal drugs)
Bevacizumab (PRES)	Methotrexate
Chronic encephalopathy (dementia)	Cytarabine
Methotrexate	Thiotepa
Carmustine	Peripheral neuropathy
Cytarabine	Vinca alkaloids
Fludarabine	Cisplatin
Visual loss	Procarbazine
Tamoxifen	Etoposide
Gallium nitrate	Teniposide
Cisplatin	Cytarabine
Fludarabine	Taxanes
Cerebellar dysfunction/ataxia	Suramin
5-Fluorouracil (± levamisole)	Bortezomib
Cytarabine	
Procarbazine	

Abbreviations: IT, intrathecal; IV, intravenous; PRES, posterior reversible encephalopathy syndrome.

vulnerable. Plexopathy develops more commonly in the brachial distribution than in the lumbosacral distribution. It must be differentiated from tumor progression in the plexus, which is usually accomplished with CT/MR imaging of the area or PET scan demonstrating tumor infiltrating the region. Clinically, tumor progression is usually painful, whereas RT-induced plexopathy is painless. Radiation plexopathy is also more commonly associated with lymphedema of the affected limb. Sensory loss and weakness are seen in both.

TOXICITY FROM CHEMOTHERAPY

Neurotoxicity is second to myelosuppression as the dose-limiting toxicity of chemotherapeutic agents (Table 118-4). Chemotherapy causes peripheral neuropathy from a number of commonly used agents, and the type of neuropathy can differ, depending on the drug. Vincristine causes paresthesias but little sensory loss and is associated with motor dysfunction, autonomic impairment (frequently ileus), and rarely cranial nerve compromise. Cisplatin causes large fiber sensory loss resulting in sensory ataxia but little cutaneous sensory loss and no weakness. The taxanes also cause a predominately sensory neuropathy. Agents such as bortezomib and thalidomide also cause neuropathy.

Encephalopathy and seizures are common toxicities from chemotherapeutic drugs. Ifosfamide can cause a severe encephalopathy, which is reversible with discontinuation of the drug and the use of methylene blue for severely affected patients. Fludarabine also causes a severe global encephalopathy that may be permanent. Bevacizumab and other anti-VEGF agents can cause posterior reversible encephalopathy syndrome. Cisplatin can cause hearing loss and less frequently vestibular dysfunction. Immunotherapy with anti-CTLA-4 monoclonal antibodies, such as ipilimumab, can cause an autoimmune hypophysitis.

Sarcomas are rare (<1% of all malignancies) mesenchymal neoplasms that arise in bone and soft tissues. These tumors are usually of mesodermal origin, although a few are derived from neuroectoderm, and they are biologically distinct from the more common epithelial malignancies. Sarcomas affect all age groups; 15% are found in children <15 years of age, and 40% occur after age 55 years. Sarcomas are one of the most common solid tumors of childhood and are the fifth most common cause of cancer deaths in children. Sarcomas may be divided into two groups, those derived from bone and those derived from soft tissues.

SOFT TISSUE SARCOMAS

Soft tissues include muscles, tendons, fat, fibrous tissue, synovial tissue, vessels, and nerves. Approximately 60% of soft tissue sarcomas arise in the extremities, with the lower extremities involved three times as often as the upper extremities. Thirty percent arise in the trunk, the retroperitoneum accounting for 40% of all trunk lesions. The remaining 10% arise in the head and neck.

INCIDENCE

Approximately 11,410 new cases of soft tissue sarcomas occurred in the United States in 2013. The annual age-adjusted incidence is 3 per 100,000 population, but the incidence varies with age. Soft tissue sarcomas constitute 0.7% of all cancers in the general population and 6.5% of all cancers in children.

EPIDEMIOLOGY

Malignant transformation of a benign soft tissue tumor is extremely rare, with the exception that malignant peripheral nerve sheath tumors (neurofibrosarcoma, malignant schwannoma) can arise from neurofibromas in patients with neurofibromatosis. Several etiologic factors have been implicated in soft tissue sarcomas.

Environmental Factors Trauma or previous injury is rarely involved, but sarcomas can arise in scar tissue resulting from a prior operation, burn, fracture, or foreign body implantation. Chemical carcinogens such as polycyclic hydrocarbons, asbestos, and dioxin may be involved in the pathogenesis.

Iatrogenic Factors Sarcomas in bone or soft tissues occur in patients who are treated with radiation therapy. The tumor nearly always arises in the irradiated field. The risk increases with time.

Viruses Kaposi's sarcoma (KS) in patients with HIV type 1, classic KS, and KS in HIV-negative homosexual men is caused by human herpesvirus (HHV) 8 (Chap. 219). No other sarcomas are associated with viruses.

Immunologic Factors Congenital or acquired immunodeficiency, including therapeutic immunosuppression, increases the risk of sarcoma.

GENETIC CONSIDERATIONS



Li-Fraumeni syndrome is a familial cancer syndrome in which affected individuals have germline abnormalities of the tumor-suppressor gene *p53* and an increased incidence of soft tissue sarcomas and other malignancies, including breast cancer, osteosarcoma, brain tumor, leukemia, and adrenal carcinoma (Chap. 101e). Neurofibromatosis 1 (*NF-1*, peripheral form, von Recklinghausen's disease) is characterized by multiple neurofibromas and café-au-lait spots. Neurofibromas occasionally undergo malignant degeneration to become malignant peripheral nerve sheath tumors. The gene for *NF-1* is located in the pericentromeric region of chromosome 17 and encodes neurofibromin, a tumor-suppressor protein with guanosine

5'-triphosphate (GTP)ase-activating activity that inhibits Ras function (Chap. 118). Germline mutation of the *Rb-1* locus (chromosome 13q14) in patients with inherited retinoblastoma is associated with the development of osteosarcoma in those who survive the retinoblastoma and of soft tissue sarcomas unrelated to radiation therapy. Other soft tissue tumors, including desmoid tumors, lipomas, leiomyomas, neuroblastomas, and paragangliomas, occasionally show a familial predisposition.

Ninety percent of synovial sarcomas contain a characteristic chromosomal translocation t(X;18)(p11;q11) involving a nuclear transcription factor on chromosome 18 called *SYT* and two breakpoints on X. Patients with translocations to the second X breakpoint (*SSX2*) may have longer survival than those with translocations involving *SSX1*.

Insulin-like growth factor (IGF) type II is produced by some sarcomas and may act as an autocrine growth factor and as a motility factor that promotes metastatic spread. IGF-II stimulates growth through IGF-I receptors, but its effects on motility are through different receptors. If secreted in large amounts, IGF-II may produce hypoglycemia (Chaps. 121 and 420).

CLASSIFICATION

Approximately 20 different groups of sarcomas are recognized on the basis of the pattern of differentiation toward normal tissue. For example, rhabdomyosarcoma shows evidence of skeletal muscle fibers with cross-striations; leiomyosarcomas contain interlacing fascicles of spindle cells resembling smooth muscle; and liposarcomas contain adipocytes. When precise characterization of the group is not possible, the tumors are called *unclassified sarcomas*. All of the primary bone sarcomas can also arise from soft tissues (e.g., extraskeletal osteosarcoma). The entity *malignant fibrous histiocytoma (MFH)* includes many tumors previously classified as fibrosarcomas or as pleomorphic variants of other sarcomas and is characterized by a mixture of spindle (fibrous) cells and round (histiocytic) cells arranged in a storiform pattern with frequent giant cells and areas of pleomorphism. As immunohistochemical suggestion of differentiation, particularly myogenic differentiation, may be found in a significant fraction of these patients, many are now characterized as poorly differentiated leiomyosarcomas, and the terms *undifferentiated pleomorphic sarcoma (UPS)* and *myxofibrosarcoma* are replacing MFH and myxoid MFH.

For purposes of treatment, most soft tissue sarcomas can be considered together. However, some specific tumors have distinct features. For example, *liposarcoma* can have a spectrum of behaviors. Pleomorphic liposarcomas and dedifferentiated liposarcomas behave like other high-grade sarcomas; in contrast, well-differentiated liposarcomas (better termed *atypical lipomatous tumors*) lack metastatic potential, and myxoid liposarcomas metastasize infrequently, but, when they do, they have a predilection for unusual metastatic sites containing fat, such as the retroperitoneum, mediastinum, and subcutaneous tissue. Rhabdomyosarcomas, Ewing's sarcoma, and other small-cell sarcomas tend to be more aggressive and are more responsive to chemotherapy than other soft tissue sarcomas.

Gastrointestinal stromal cell tumors (GISTs), previously classified as gastrointestinal leiomyosarcomas, are now recognized as a distinct entity within soft tissue sarcomas. Its cell of origin resembles the interstitial cell of Cajal, which controls peristalsis. The majority of malignant GISTs have activating mutations of the *c-kit* gene that result in ligand-independent phosphorylation and activation of the KIT receptor tyrosine kinase, leading to tumorigenesis. Approximately 5–10% of tumors will have a mutation in the platelet-derived growth factor receptor α (*PDGFRA*). GISTs that are wild type for both *KIT* and *PDGFRA* mutations may show mutations in *SDH* B, C, or D and may be driven by the IGF-I pathway.

DIAGNOSIS

The most common presentation is an asymptomatic mass. Mechanical symptoms referable to pressure, traction, or entrapment of nerves or muscles may be present. All new and persistent or growing masses should be biopsied, either by a cutting needle (core-needle biopsy) or by a small incision, placed so that it can be encompassed in the

TABLE 119e-1 AMERICAN JOINT COMMITTEE ON CANCER STAGING SYSTEM FOR SARCOMAS

Histologic Grade (G)	Tumor Size (T)	Node Status (N)	Metastases (M)
Well differentiated (G1)	≤5 cm (T1)	Not involved (N0)	Absent (M0)
Moderately differentiated (G2)	>5 cm (T2)	Involved (N1)	Present (M1)
Poorly differentiated (G3)	Superficial fascial involvement (Ta)		
Undifferentiated (G4)	Deep fascial involvement (Tb)		
Disease Stage	5-Year Survival, %		
Stage I	98.8		
A: G1,2; T1a,b; N0; M0			
B: G1,2; T2a; N0; M0			
Stage II	81.8		
A: G1,2; T2b; N0; M0			
B: G3,4; T1; N0; M0			
C: G3,4; T2a; N0; M0			
Stage III: G3,4; T2b; N0; M0	51.7		
Stage IV	<20		
A: any G; any T; N1; M0			
B: any G; any T; any N; M1			

subsequent excision without compromising a definitive resection. Lymph node metastases occur in 5%, except in synovial and epithelioid sarcomas, clear-cell sarcoma (melanoma of the soft parts), angiosarcoma, and rhabdomyosarcoma, where nodal spread may be seen in 17%. The pulmonary parenchyma is the most common site of metastases. Exceptions are GISTs, which metastasize to the liver; myxoid liposarcomas, which seek fatty tissue; and clear-cell sarcomas, which may metastasize to bones. Central nervous system metastases are rare, except in alveolar soft part sarcoma.

Radiographic Evaluation Imaging of the primary tumor is best with plain radiographs and magnetic resonance imaging (MRI) for tumors of the extremities or head and neck and by computed tomography (CT) for tumors of the chest, abdomen, or retroperitoneal cavity. A radiograph and CT scan of the chest are important for the detection of lung metastases. Other imaging studies may be indicated, depending on the symptoms, signs, or histology.

STAGING AND PROGNOSIS

The histologic grade, relationship to fascial planes, and size of the primary tumor are the most important prognostic factors. The current American Joint Committee on Cancer (AJCC) staging system is shown in [Table 119e-1](#). Prognosis is related to the stage. Cure is common in the absence of metastatic disease, but a small number of patients with metastases can also be cured. Most patients with stage IV disease die within 12 months, but some patients may live with slowly progressive disease for many years.

TREATMENT SOFT TISSUE SARCOMAS

AJCC stage I patients are adequately treated with surgery alone. Stage II patients are considered for adjuvant radiation therapy. Stage III patients may benefit from adjuvant chemotherapy. Stage IV patients are managed primarily with chemotherapy, with or without other modalities.

SURGERY

Soft tissue sarcomas tend to grow along fascial planes, with the surrounding soft tissues compressed to form a pseudocapsule that gives the sarcoma the appearance of a well-encapsulated lesion. This is invariably deceptive because “shelling out,” or marginal excision, of such lesions results in a 50–90% probability of local recurrence. Wide excision with a negative margin, incorporating the biopsy site, is the standard surgical procedure for local disease. The adjuvant use of radiation therapy and/or chemotherapy

improves the local control rate and permits the use of limb-sparing surgery with a local control rate (85–90%) comparable to that achieved by radical excisions and amputations. Limb-sparing approaches are indicated except when negative margins are not obtainable, when the risks of radiation are prohibitive, or when neurovascular structures are involved so that resection will result in serious functional consequences to the limb.

RADIATION THERAPY

External-beam radiation therapy is an adjuvant to limb-sparing surgery for improved local control. Preoperative radiation therapy allows the use of smaller fields and smaller doses but results in a higher rate of wound complications. Postoperative radiation therapy must be given to larger fields, because the entire surgical bed must be encompassed, and in higher doses to compensate for hypoxia in the operated field. This results

in a higher rate of late complications. Brachytherapy or interstitial therapy, in which the radiation source is inserted into the tumor bed, is comparable in efficacy (except in low-grade lesions), less time-consuming, and less expensive.

ADJUVANT CHEMOTHERAPY

Chemotherapy is the mainstay of treatment for Ewing's primitive neuroectodermal tumors (PNET) and rhabdomyosarcomas. Meta-analysis of 14 randomized trials revealed a significant improvement in local control and disease-free survival in favor of doxorubicin-based chemotherapy. Overall survival improvement was 4% for all sites and 7% for the extremity site. An updated meta-analysis including four additional trials with doxorubicin and ifosfamide combination has reported a statistically significant 6% survival advantage in favor of chemotherapy. A chemotherapy regimen including an anthracycline and ifosfamide with growth factor support improved overall survival by 19% for high-risk (high-grade, ≥5 cm primary, or locally recurrent) extremity soft tissue sarcomas.

ADVANCED DISEASE

Metastatic soft tissue sarcomas are largely incurable, but up to 20% of patients who achieve a complete response become long-term survivors. The therapeutic intent, therefore, is to produce a complete remission with chemotherapy (<10%) and/or surgery (30–40%). Surgical resection of metastases, whenever possible, is an integral part of the management. Some patients benefit from repeated surgical excision of metastases. The two most active chemotherapeutic agents are doxorubicin and ifosfamide. These drugs show a steep dose-response relationship in sarcomas. Gemcitabine with or without docetaxel has become an established second-line regimen and is particularly active in patients with undifferentiated pleomorphic sarcoma (UPS) and leiomyosarcomas. Dacarbazine also has some modest activity. Taxanes have selective activity in angiosarcomas, and vincristine, etoposide, and irinotecan are effective in rhabdomyosarcomas and Ewing's sarcomas. Pazopanib, an inhibitor of the vascular endothelial growth factor, platelet-derived growth factor (PDGF), and c-kit is now approved for patients with advanced soft tissue sarcomas excluding liposarcomas after failure of chemotherapy. Imatinib targets the KIT and PDGF tyrosine kinase activity and is standard therapy for advanced/metastatic GISTs and dermatofibrosarcoma protuberans. Imatinib is now also indicated as adjuvant therapy for completely resected primary GISTs. Three years of adjuvant imatinib appears to be superior to 1 year of therapy for high-risk GISTs, although the optimal treatment duration remains unknown.

BONE SARCOMAS

INCIDENCE AND EPIDEMIOLOGY

Bone sarcomas are rarer than soft tissue sarcomas; they accounted for only 0.2% of all new malignancies and 2890 new cases in the United States in 2013. Several benign bone lesions have the potential for malignant transformation. Enchondromas and osteochondromas can transform into chondrosarcoma; fibrous dysplasia, bone infarcts, and Paget's disease of bone can transform into either UPS or osteosarcoma.

CLASSIFICATION

Benign Tumors The common benign bone tumors include enchondroma, osteochondroma, chondroblastoma, and chondromyxoid fibroma, of cartilage origin; osteoid osteoma and osteoblastoma, of bone origin; fibroma and desmoplastic fibroma, of fibrous tissue origin; hemangioma, of vascular origin; and giant-cell tumor, of unknown origin.

Malignant Tumors The most common malignant tumors of bone are plasma cell tumors (Chap. 136). The four most common malignant nonhematopoietic bone tumors are osteosarcoma, chondrosarcoma, Ewing's sarcoma, and UPS. Rare malignant tumors include chordoma (of notochordal origin), malignant giant-cell tumor and adamantinoma (of unknown origin), and hemangioendothelioma (of vascular origin).

Musculoskeletal Tumor Society Staging System Sarcomas of bone are staged according to the Musculoskeletal Tumor Society staging system based on grade and compartmental localization. A Roman numeral reflects the tumor grade: stage I is low grade, stage II is high grade, and stage III includes tumors of any grade that have lymph node or distant metastases. In addition, the tumor is given a letter reflecting its compartmental localization. Tumors designated A are intracompartmental (i.e., confined to the same soft tissue compartment as the initial tumor), and tumors designated B are extracompartmental (i.e., extending into the adjacent soft tissue compartment or into bone). The tumor-node-metastasis (TNM) staging system is shown in Table 119e-2.

OSTEOSARCOMA

Osteosarcoma, accounting for almost 45% of all bone sarcomas, is a spindle cell neoplasm that produces osteoid (unmineralized bone) or bone. Approximately 60% of all osteosarcomas occur in children and adolescents in the second decade of life, and approximately 10% occur in the third decade of life. Osteosarcomas in the fifth and sixth decades of life are frequently secondary to either radiation therapy or transformation in a preexisting benign condition, such as Paget's disease. Males are affected 1.5–2 times as often as females. Osteosarcoma has a predilection for metaphyses of long bones; the most common sites of involvement are the distal femur, proximal tibia, and proximal humerus. The classification of osteosarcoma is complex, but 75% of osteosarcomas fall into the "classic" category, which include osteoblastic, chondroblastic, and fibroblastic osteosarcomas. The remaining 25% are classified as "variants" on the basis of (1) clinical characteristics, as in the case of osteosarcoma of the jaw, postradiation osteosarcoma, or Paget's osteosarcoma; (2) morphologic characteristics, as in the case of telangiectatic osteosarcoma, small-cell osteosarcoma, or epithelioid osteosarcoma; or (3) location, as in parosteal or periosteal osteosarcoma. Diagnosis usually requires a synthesis of clinical, radiologic, and pathologic features. Patients typically present with pain and swelling of the affected area. A plain radiograph reveals a destructive lesion with a moth-eaten appearance, a spiculated periosteal reaction (sunburst appearance), and a cuff of periosteal new bone formation at the margin of the soft tissue mass (Codman's triangle). A CT scan of the primary tumor is best for defining bone destruction and the pattern of calcification, whereas MRI is better for defining intramedullary and soft tissue extension. A chest radiograph and CT scan are used to detect lung metastases. Metastases to the bony skeleton should be imaged by a bone scan or by fluorodeoxyglucose positron emission tomography (FDG-PET). Almost all osteosarcomas are hypervascular. Angiography is not helpful for diagnosis, but it is the most sensitive test for assessing the response to preoperative

TABLE 119e-2 STAGING SYSTEM FOR BONE SARCOMAS

Primary tumor (T)	TX	Primary tumor cannot be assessed
	T0	No evidence of primary tumor
	T1	Tumor ≤8 cm in greatest dimension
	T2	Tumor >8 cm in greatest dimension
	T3	Discontinuous tumors in the primary bone site
Regional lymph nodes (N)	NX	Regional lymph nodes cannot be assessed
	N0	No regional lymph node metastasis
	N1	Regional lymph node metastasis
Distant metastasis (M)	MX	Distant metastasis cannot be assessed
	M0	No distant metastasis
	M1	Distant metastasis
	M1a	Lung
	M1b	Other distant sites
Histologic grade (G)	GX	Grade cannot be assessed
	G1	Well differentiated—low grade
	G2	Moderately differentiated—low grade
	G3	Poorly differentiated—high grade
	G4	Undifferentiated—high grade (Ewing's is always classed G4)

Stage Grouping

Stage IA	T1	N0	M0	G1,2 low grade
Stage IB	T2	N0	M0	G1,2 low grade
Stage IIA	T1	N0	M0	G3,4 high grade
Stage IIB	T2	N0	M0	G3,4 high grade
Stage III	T3	N0	M0	Any G
Stage IVA	Any T	N0	M1a	Any G
Stage IVB	Any T	N1	Any M	Any G
	Any T	Any N	M1b	Any G

chemotherapy. Pathologic diagnosis is established either with a core-needle biopsy, where feasible, or with an open biopsy with an appropriately placed incision that does not compromise future limb-sparing resection. Most osteosarcomas are high-grade. The most important prognostic factor for long-term survival is response to chemotherapy. Preoperative chemotherapy followed by limb-sparing surgery (which can be accomplished in >80% of patients) followed by postoperative chemotherapy is standard management. The effective drugs are doxorubicin, ifosfamide, cisplatin, and high-dose methotrexate with leucovorin rescue. The various combinations of these agents that have been used have all been about equally successful. Long-term survival rates in extremity osteosarcoma range from 60 to 80%. Osteosarcoma is radioresistant; radiation therapy has no role in the routine management. UPS is considered a part of the spectrum of osteosarcoma and is managed similarly.

CHONDROSARCOMA

Chondrosarcoma, which constitutes ~20–25% of all bone sarcomas, is a tumor of adulthood and old age with a peak incidence in the fourth to sixth decades of life. It has a predilection for the flat bones, especially the shoulder and pelvic girdles, but can also affect the diaphyseal portions of long bones. Chondrosarcomas can arise de novo or as a malignant transformation of an enchondroma or, rarely, of the cartilaginous cap of an osteochondroma. Chondrosarcomas have an indolent natural history and typically present as pain and swelling. Radiographically, the lesion may have a lobular appearance with mottled or punctate or annular calcification of the cartilaginous matrix. It is difficult to distinguish low-grade chondrosarcoma from benign lesions by x-ray or histologic examination. The diagnosis is therefore influenced by clinical history and physical examination. A new onset of pain, signs of inflammation, and progressive increase in the size of the mass suggest malignancy. The histologic classification is complex,

but most tumors fall within the classic category. Like other bone sarcomas, high-grade chondrosarcomas spread to the lungs. Most chondrosarcomas are resistant to chemotherapy, and surgical resection of primary or recurrent tumors, including pulmonary metastases, is the mainstay of therapy. This rule does not hold for two histologic variants. Dedifferentiated chondrosarcoma has a high-grade osteosarcoma or a malignant fibrous histiocytoma component that responds to chemotherapy. Mesenchymal chondrosarcoma, a rare variant composed of a small-cell element, also is responsive to systemic chemotherapy and is treated like Ewing's sarcoma.

EWING'S SARCOMA

Ewing's sarcoma, which constitutes ~10–15% of all bone sarcomas, is common in adolescence and has a peak incidence in the second decade of life. It typically involves the diaphyseal region of long bones and also has an affinity for flat bones. The plain radiograph may show a characteristic “onion peel” periosteal reaction with a generous soft tissue mass, which is better demonstrated by CT or MRI. This mass is composed of sheets of monotonous, small, round, blue cells and can be confused with lymphoma, embryonal rhabdomyosarcoma, and small-cell carcinoma. The presence of p30/32, the product of the *mic-2* gene (which maps to the pseudoautosomal region of the X and Y chromosomes), is a cell-surface marker for Ewing's sarcoma (and other members of the Ewing's family of tumors, sometimes called PNETs). Most PNETs arise in soft tissues; they include peripheral neuroepithelioma, Askin's tumor (chest wall), and esthesioneuroblastoma. Glycogen-filled cytoplasm detected by staining with periodic acid–Schiff is also characteristic of Ewing's sarcoma cells. The classic cytogenetic abnormality associated with this disease (and other PNETs) is a reciprocal translocation of the long arms of chromosomes 11 and 22, t(11;22), which creates a chimeric gene product of unknown function with components from the *fli-1* gene on chromosome 11 and *ews* on 22. This disease is very aggressive, and it is therefore considered a systemic disease. Common sites of metastases are lung, bones, and bone marrow. Systemic chemotherapy is the mainstay of therapy, often being used before surgery. Doxorubicin, cyclophosphamide or ifosfamide, etoposide, vincristine, and dactinomycin are active drugs. Topotecan or irinotecan in combination with an alkylating agent is often used in relapsed patients. Targeted therapy with an anti-IGF-I receptor antibody in combination with an inhibitor of mammalian target of rapamycin (mTOR) appears to have promising activity in refractory cases. Local treatment for the primary tumor includes surgical resection, usually with limb salvage or radiation therapy. Patients with lesions below the elbow and below the mid-calf have a 5-year survival rate of 80% with effective treatment. Ewing's sarcoma at first presentation is a curable tumor, even in the presence of obvious metastatic disease, especially in children <11 years old.

TUMORS METASTATIC TO BONE

Bone is a common site of metastasis for carcinomas of the prostate, breast, lung, kidney, bladder, and thyroid and for lymphomas and sarcomas. Prostate, breast, and lung primaries account for 80% of all bone metastases. Metastatic tumors of bone are more common than primary bone tumors. Tumors usually spread to bone hematogenously, but local invasion from soft tissue masses also occurs. In descending order of frequency, the sites most often involved are the vertebrae, proximal femur, pelvis, ribs, sternum, proximal humerus, and skull. Bone metastases may be asymptomatic or may produce pain, swelling, nerve root or spinal cord compression, pathologic fracture, or myelophthisis

(replacement of the marrow). Symptoms of hypercalcemia may be noted in cases of bony destruction.

Pain is the most frequent symptom. It usually develops gradually over weeks, is usually localized, and often is more severe at night. When patients with back pain develop neurologic signs or symptoms, emergency evaluation for spinal cord compression is indicated ([Chap. 331](#)). Bone metastases exert a major adverse effect on quality of life in cancer patients.

Cancer in the bone may produce osteolysis, osteogenesis, or both. Osteolytic lesions result when the tumor produces substances that can directly elicit bone resorption (vitamin D–like steroids, prostaglandins, or parathyroid hormone–related peptide) or cytokines that can induce the formation of osteoclasts (interleukin 1 and tumor necrosis factor). Osteoblastic lesions result when the tumor produces cytokines that activate osteoblasts. In general, purely osteolytic lesions are best detected by plain radiography, but they may not be apparent until they are >1 cm. These lesions are more commonly associated with hypercalcemia and with the excretion of hydroxyproline-containing peptides indicative of matrix destruction. When osteoblastic activity is prominent, the lesions may be readily detected using radionuclide bone scanning (which is sensitive to new bone formation), and the radiographic appearance may show increased bone density or sclerosis. Osteoblastic lesions are associated with higher serum levels of alkaline phosphatase and, if extensive, may produce hypocalcemia. Although some tumors may produce mainly osteolytic lesions (e.g., kidney cancer) and others mainly osteoblastic lesions (e.g., prostate cancer), most metastatic lesions produce both types of lesion and may go through stages where one or the other predominates.

In older patients, particularly women, it may be necessary to distinguish metastatic disease of the spine from osteoporosis. In osteoporosis, the cortical bone may be preserved, whereas cortical bone destruction is usually noted with metastatic cancer.

TREATMENT METASTATIC BONE DISEASE

Treatment of metastatic bone disease depends on the underlying malignancy and the symptoms. Some metastatic bone tumors are curable (lymphoma, Hodgkin's disease), and others are treated with palliative intent. Pain may be relieved by local radiation therapy. Hormonally responsive tumors are responsive to hormone inhibition (antiandrogens for prostate cancer, antiestrogens for breast cancer). Strontium-89, samarium-153, and radium-223 are bone-seeking radionuclides that can exert antitumor effects and relieve symptoms. Denosumab, a monoclonal antibody that binds to RANK ligand, inhibits osteoclastic activity and increases bone mineral density. Bisphosphonates such as pamidronate may relieve pain and inhibit bone resorption, thereby maintaining bone mineral density and reducing risk of fractures in patients with osteolytic metastases from breast cancer and multiple myeloma. Careful monitoring of serum electrolytes and creatinine is recommended. Monthly administration prevents bone-related clinical events and may reduce the incidence of bone metastases in women with breast cancer. When the integrity of a weight-bearing bone is threatened by an expanding metastatic lesion that is refractory to radiation therapy, prophylactic internal fixation is indicated. Overall survival is related to the prognosis of the underlying tumor. Bone pain at the end of life is particularly common; an adequate pain relief regimen including sufficient amounts of narcotic analgesics is required. The management of hypercalcemia is discussed in [Chap. 424](#).

Carcinoma of unknown primary (CUP) is a biopsy-proven malignancy for which the anatomic site of origin remains unidentified after an intensive search. CUP is one of the 10 most frequently diagnosed cancers worldwide, accounting for 3–5% of all cancers. Most investigators limit CUP to epithelial cancers and do not include lymphomas, metastatic melanomas, and metastatic sarcomas because these cancers have specific histology- and stage-based treatments that guide management.

The emergence of sophisticated imaging, robust immunohistochemistry (IHC), and genomic and proteomic tools has challenged the “unknown” designation. Additionally, effective targeted therapies in several cancers have moved the paradigm from empiricism to considering a personalized approach to CUP management. The reasons cancers present as CUP remain unclear. One hypothesis is that the primary tumor either regresses after seeding the metastasis or remains so small that it is not detected. It is possible that CUP falls on the continuum of cancer presentation where the primary has been contained or eliminated by the natural body defenses. Alternatively, CUP may represent a specific malignant event that results in an increase in metastatic spread or survival relative to the primary. Whether the CUP metastases truly define a clone that is genetically and phenotypically unique to this diagnosis remains to be determined.

CUP BIOLOGY

Studies looking for unique signature abnormalities in CUP tumors have not been positive. Abnormalities in chromosomes 1 and 12 and other complex cytogenetic abnormalities have been reported. Aneuploidy has been described in 70% of CUP patients with metastatic adenocarcinoma or undifferentiated carcinoma. The overexpression of various genes, including *Ras*, *bcl-2* (40%), *her-2* (11%), and *p53* (26–53%), has been studied in CUP samples, but they have no effect on response to therapy or survival. The extent of angiogenesis in CUP relative to that in metastases from known primaries has also been evaluated, but no consistent findings have emerged. Using the Sequenom (SQM) Massarray platform, a study in consecutive CUP patients showed that the overall mutational rate was surprisingly low (18%). No “new” low-frequency mutations were found using a panel of mutations involving the P13K/AKT pathway, MEK pathway, receptors, and downstream effectors. Nevertheless, it is possible that newer “deep sequencing” techniques in select patients may yield consistent abnormalities.

CLINICAL EVALUATION

Initial CUP evaluation has two goals: search for the primary tumor based on pathologic evaluation of the metastases and determine the extent of disease. Obtaining a thorough medical history from CUP patients is essential, including paying particular attention to previous surgeries, removed lesions, and family medical history to assess potential hereditary cancers. Adequate physical examination, including a digital rectal examination in men and breast and pelvic examinations in women, should be performed based on clinical presentation.

Role of Serum Tumor Markers and Cytogenetics Most tumor markers, including CEA, CA-125, CA 19-9, and CA 15-3, when elevated, are nonspecific and not helpful in determining the primary tumor site. Men who present with adenocarcinoma and osteoblastic metastasis should undergo a prostate-specific antigen (PSA) test. In patients with undifferentiated or poorly differentiated carcinoma (especially with a midline tumor), elevated β -human chorionic gonadotropin (β -hCG) and α fetoprotein (AFP) levels suggest the possibility of an extragonadal germ cell (testicular) tumor. With the availability of IHC, cytogenetic studies are rarely needed.

Role of Imaging Studies In the absence of contraindications, a baseline IV contrast computed tomography (CT) scan of the chest, abdomen, and pelvis is the standard of care. This helps to search for the primary tumor, evaluate the extent of disease, and select the most accessible biopsy site. Older studies suggested that the primary tumor site is detected in 20–35% of patients who undergo a CT scan of the abdomen and pelvis, although by current definition, these patients do not have CUP. These studies also suggest a latent primary tumor prevalence of 20%; with more sophisticated imaging, this has decreased to $\leq 5\%$ today.

Mammography should be performed in all women who present with metastatic adenocarcinoma, especially in those with adenocarcinoma and isolated axillary lymphadenopathy. Magnetic resonance imaging (MRI) of the breast is a follow-up modality in patients with axillary adenopathy and suspected occult primary breast carcinoma following a negative mammography and ultrasound. The results of these imaging modalities can influence surgical management; a negative breast MRI result predicts a low tumor yield at mastectomy.

A conventional workup for a squamous cell carcinoma and cervical CUP (neck lymphadenopathy with no known primary tumor) includes a CT scan or MRI and invasive studies, including indirect and direct laryngoscopy, bronchoscopy, and upper endoscopy. Ipsilateral (or bilateral) staging tonsillectomy has been recommended for these patients. 18-Fluorodeoxyglucose positron emission tomography (18-FDG-PET) scans are useful in this patient population and may help guide the biopsy; determine the extent of disease; facilitate the appropriate treatment, including planning radiation fields; and help with disease surveillance. A smaller radiation field encompassing the primary (when found) and metastatic adenopathy decreases the risk of chronic xerostomia. Several studies have evaluated the utility of PET in patients with squamous cervical CUP, and head and neck primary tumors were identified in ~21–30%.

The diagnostic contribution of PET to the evaluation of other CUP (outside of the neck adenopathy indication) remains controversial and is not routinely recommended. PET-CT can be helpful for patients who are candidates for surgical intervention for solitary metastatic disease because the presence of disease outside the primary site may affect surgical planning.

Invasive studies, including upper endoscopy, colonoscopy, and bronchoscopy, should be limited to symptomatic patients or those with laboratory, imaging, or pathologic abnormalities that suggest that these techniques will result in a high yield in finding a primary cancer.

Role of Pathologic Studies A detailed pathologic examination of the most accessible biopsied tissue specimen is mandatory in CUP patients. Pathologic evaluation typically consists of hematoxylin and eosin stains and immunohistochemical tests.

LIGHT MICROSCOPY EVALUATION Adequate tissue obtained preferably by excisional biopsy or core-needle biopsy (instead of only a fine-needle aspiration) is stained with hematoxylin and eosin and subjected to light microscopic examination. On light microscopy, 60–65% of CUP is adenocarcinoma, and 5% is squamous cell carcinoma. The remaining 30–35% is poorly differentiated adenocarcinoma, poorly differentiated carcinoma, or poorly differentiated neoplasm. A small percentage of lesions are diagnosed as neuroendocrine cancers (2%), mixed tumors (adenosquamous or sarcomatoid carcinomas), or undifferentiated neoplasms (Table 120e-1).

TABLE 120e-1 MAJOR HISTOLOGIES IN CARCINOMA OF UNKNOWN PRIMARY

Histology	Proportion, %
Well to moderately differentiated adenocarcinoma	60
Squamous cell cancer	5
Poorly differentiated adenocarcinoma, poorly differentiated carcinoma	30
Neuroendocrine	2
Undifferentiated malignancy	3

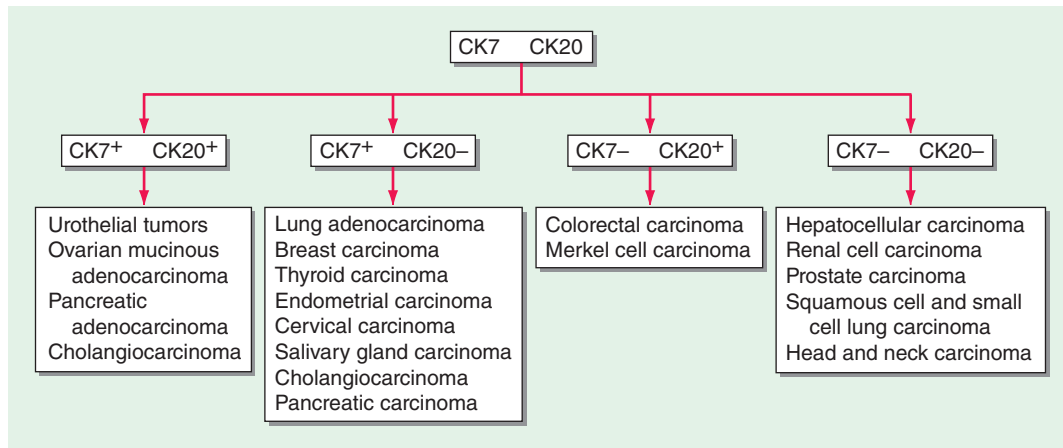


FIGURE 120e-1 Approach to cytokeratin (CK7 and CK20) markers used in adenocarcinoma of unknown primary.

ROLE OF IMMUNOHISTOCHEMICAL ANALYSIS Immunohistochemical stains are peroxidase-labeled antibodies against specific tumor antigens that are used to define tumor lineage. The number of available immunohistochemical stains is ever-increasing. However, in CUP cases, more is not necessarily better, and immunohistochemical stains should be used in conjunction with the patient's clinical presentation and imaging studies to select the best therapy. Communication between the clinician and pathologist is essential. No stain is 100% specific, and overinterpretation should be avoided. PSA and thyroglobulin tissue markers, which are positive in prostate and thyroid cancer, respectively, are the most specific of the current marker panel. However, these cancers rarely present as CUP, so the yield of these tests may be low. Fig. 120e-1 delineates a simple algorithm for immunohistochemical staining in CUP cases. Table 120e-2 lists additional tests that may be useful to further define the tumor lineage. A more comprehensive algorithm may improve the diagnostic accuracy but can make the process complex. With the use of immunohistochemical markers, electron microscopic analysis, which is time-consuming and expensive, is rarely needed.

There are >20 subtypes of cytokeratin (CK) intermediate filaments with different molecular weights and differential expression in various cell types and cancers. Monoclonal antibodies to specific CK subtypes have been used to help classify tumors according to their site of origin; commonly used CK stains in adenocarcinoma CUP are CK7 and CK20. CK7 is found in tumors of the lung, ovary, endometrium, breast, and upper gastrointestinal tract including pancreaticobiliary cancers, whereas CK20 is normally expressed in the gastrointestinal epithelium, urothelium, and Merkel cells. The nuclear CDX-2 transcription factor, which is the product of a homeobox gene necessary for intestinal organogenesis, is often used to aid in the diagnosis of gastrointestinal adenocarcinomas.

Thyroid transcription factor 1 (TTF-1) nuclear staining is typically positive in lung and thyroid cancers. Approximately 68% of adenocarcinomas and 25% of squamous cell lung cancers stain positive for TTF-1, which helps differentiate a lung primary tumor from metastatic adenocarcinoma in a pleural effusion, the mediastinum, or the lung parenchyma.

Gross cystic disease fibrous protein-15, a 15-kDa monomer protein, is a marker of apocrine differentiation that is detected in 62–72% of breast carcinomas. UROIII, high-molecular-weight cytokeratin, thrombomodulin, and CK20 are the markers used to diagnose lesions of urothelial origin.

IHC performs the best when used in groups that give rise to patterns that are strongly indicative of certain profiles. For example, the TTF-1/CK7+ and CK20+/CDX-2+/CK7- phenotypes have been reported as very suggestive of lung and lower gastrointestinal cancer profiles, respectively, although these patterns have not been validated prospectively in the absence of a primary cancer. IHC is not without its limitations; several factors affect tissue antigenicity (antigen retrieval, specimen processing, and fixation), interpretation of stains in tumor

(nuclear, cytoplasmic, membrane) versus normal tissue, inter- and intraobserver variability, and tissue heterogeneity and inadequacy (given small biopsy sizes). Communication with the pathologist is critical to determine if additional tissue will be beneficial in the pathologic evaluation.

ROLE OF TISSUE OF ORIGIN MOLECULAR PROFILING In the absence of a known primary, developing therapeutic strategies for CUP is challenging. The current diagnostic yield with imaging and immunochemistry is ~20–30% for CUP patients. The use of gene expression studies holds the

TABLE 120e-2 SELECT IMMUNOHISTOCHEMICAL STAINS USEFUL IN THE DIAGNOSIS OF CARCINOMA OF UNKNOWN PRIMARY (CUP)

Likely Primary Profile	Commonly Considered IHC to Assist in Differential Diagnosis of CUP ^a
Breast	Estrogen receptor (ER), gross cystic disease fibrous protein-15 (GCDFFP-15), mammaglobin, Her-2/neu
Ovarian/mullerian	Estrogen receptor (ER), Wilms' tumor gene (WT-1), CK7, PAX8, PAX2
Lung adenocarcinoma	Thyroid transcription factor (TTF-1; nuclear staining), napsin A, surfactant protein A precursor (SP-A1)
Germ cell	β-hCG, AFP, OCT3/4, CKIT, CD30 (embryonal), SALL4
Prostate	PSA, α-methylacyl CoA racemase/P504S (AMACR/P504S), P501S (prostein), and prostate-specific membrane antigen (PSMA)
Intestinal	CK7, CK20, CDX-2, carcinoembryonic antigen (CEA)
Neuroendocrine	Chromogranin, synaptophysin, CD56
Sarcoma	Desmin (desmoid tumors), factor VIII (angiosarcomas), CD31, smooth muscle actin (leiomyosarcoma), MyoD1 (rhabdomyosarcoma)
Renal	RCC, CD10, PAX8
Hepatocellular carcinoma	Hep par-1, arginase-1 (Arg-1), TTF-1 (granular cytoplasmic staining)
Melanoma	S100, vimentin, HMB-45, tyrosinase and melan-A
Urothelial	CK7, CK20, thrombomodulin
Mesothelioma	Calretinin, WT-1
Lymphoma	Leukocyte common antigen (LCA), CD3, CD4, CD5, CD20, CD45
Squamous cell carcinoma (SCC)	p63, p40 (lung SCC), CK5/6

^aPatterns emerging from coexpression of stains are better than individual stains to suggest putative primary site. Even with optimization, no IHC panel is 100% sensitive or specific (e.g., ovarian mucinous carcinoma can exhibit positivity with intestinal markers).

Abbreviations: AFP, α fetoprotein; β-hCG, β human chorionic gonadotropin; CUP, carcinoma of unknown primary; IHC, immunohistochemistry; PSA, prostate-specific antigen.

promise of substantially increasing this yield. Gene expression profiles are most commonly generated using quantitative reverse transcriptase polymerase chain reaction (RT-PCR) or DNA microarray.

Neural network programs have been used to develop predictive algorithms from the gene expression profiles. Typically, a training set of gene profiles from known cancers (preferably from metastatic sites) is used to train the software. The program can then be used to predict the putative origin of a test tumor and presumably of true CUP. Comprehensive gene expression databases that have become available for common malignancies may also be useful in CUP. These approaches have been effective in testing against known primary cancers and their metastases.

mRNA- or microRNA-based tissue of origin molecular profiling assays have been studied in prospective and retrospective CUP trials. Most of the CUP studies have evaluated assay *performance*, although the challenge with validating the accuracy of an assay for CUP is that, by definition, the primary cancer diagnosis cannot be verified. Thus, current estimates of tissue of origin test accuracy have relied on indirect metrics including comparison with IHC, clinical presentation, and appearance of latent primaries. Using these measures, the assays suggest a plausible primary in ~70% of patients studied. The only outcomes-based study is a single-arm study reporting a median survival of 12.5 months for patients who received assay-directed site-specific therapy. Firm conclusions of therapeutic impact cannot be drawn from this study given the nonrandomized design, statistical biases, confounding variables including use of subsequent lines of (empiric) therapy, and the heterogeneity of the CUP cancers. Additional studies are needed to better understand the clinical influence of tissue of origin profiling tools and how these assays complement IHC and help guide therapy.

TREATMENT CARCINOMA OF UNKNOWN PRIMARY

GENERAL CONSIDERATIONS

The treatment of CUP continues to evolve, albeit slowly. The median survival duration of most patients with disseminated CUP is ~6–10 months. Systemic chemotherapy is the primary treatment modality

in most patients with disseminated disease, but the careful integration of surgery, radiation therapy, and even periods of observation is important in the overall management of this condition (Figs. 120e-2 and 120e-3). Prognostic factors include performance status, site and number of metastases, response to chemotherapy, and serum lactate dehydrogenase (LDH) levels. Culine and colleagues developed a prognostic model using performance status and serum LDH levels, which allowed the assignment of patients into two subgroups with divergent outcomes. Future prospective trials using this prognostic model are warranted. Clinically, some CUP diagnoses fall into a favorable prognostic subset. Others, including those with disseminated CUP, do not and have a more unfavorable prognosis.

TREATMENT OF FAVORABLE CUP SUBSETS

Women with Isolated Axillary Adenopathy Women with isolated axillary adenopathy with adenocarcinoma or carcinoma are usually treated for stage II or III breast cancer based on pathologic findings. These patients should undergo a breast MRI if mammogram and ultrasound are negative. Radiation therapy to the ipsilateral breast is indicated if the breast MRI is positive. Chemotherapy and/or hormonal therapy are indicated based on patient's age (premenopausal or postmenopausal), nodal disease bulk, and hormone receptor status (Chap. 108). It is important to verify that the pathology suggests a breast cancer profile (morphology, immunohistochemical breast markers including estrogen receptor, mammoglobin, GCDFP-15, HER-2 gene expression) before embarking on a breast cancer therapeutic program.

WOMEN WITH PERITONEAL CARCINOMATOSIS

The term *primary peritoneal papillary serous carcinoma* (PPSC) has been used to describe CUP with carcinomatosis with the pathologic and laboratory (elevated CA-125 antigen) characteristics of ovarian cancer but no ovarian primary tumor identified on transvaginal sonography or laparotomy. Studies suggest that ovarian cancer and PPSC, which are both of müllerian origin, have similar gene expression profiles. Similar to patients with ovarian cancer, patients with PPSC are candidates for cytoreductive surgery, followed by adjuvant taxane- and platinum-based chemotherapy. In one

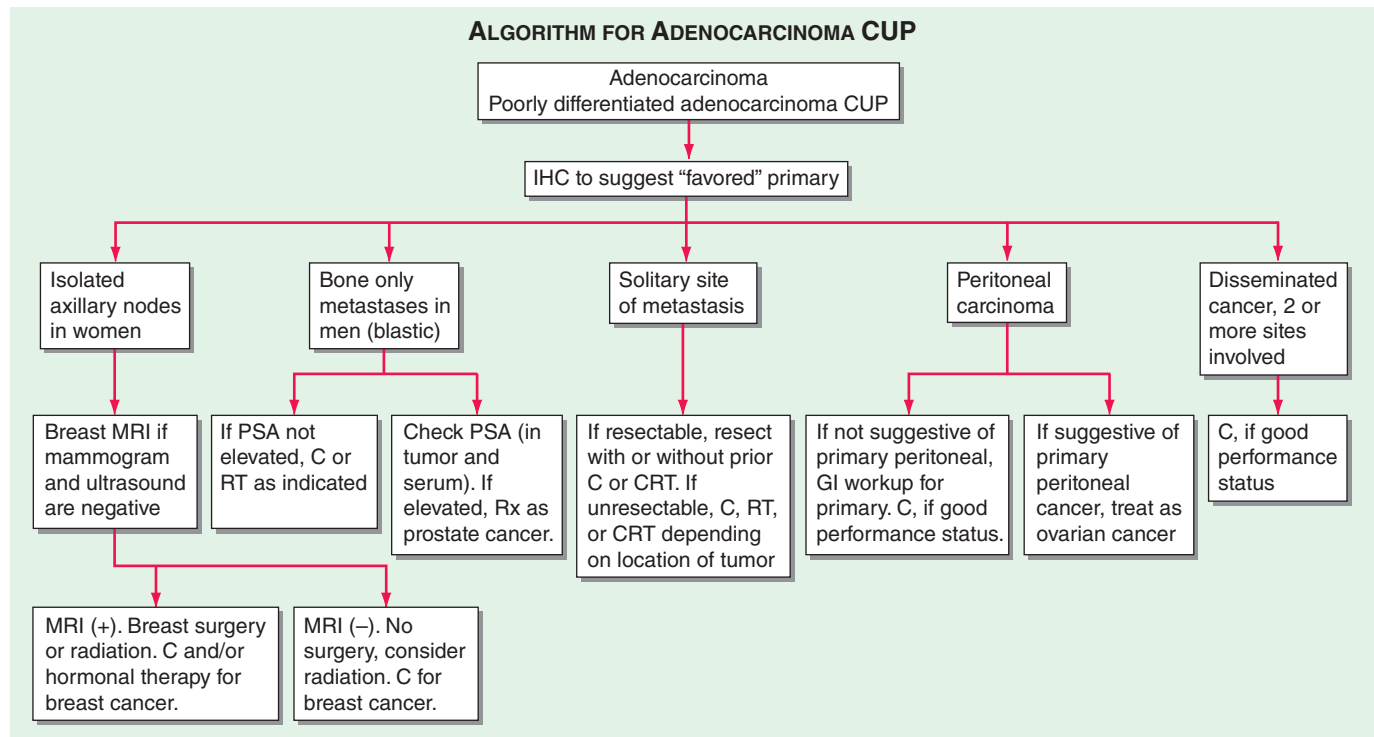


FIGURE 120e-2 Treatment algorithm for adenocarcinoma and poorly differentiated adenocarcinoma of unknown primary (CUP).

C, chemotherapy; CRT, chemoradiation; GI, gastrointestinal; IHC, immunohistochemistry; MRI, magnetic resonance imaging; PSA, prostate-specific antigen; RT, radiation.

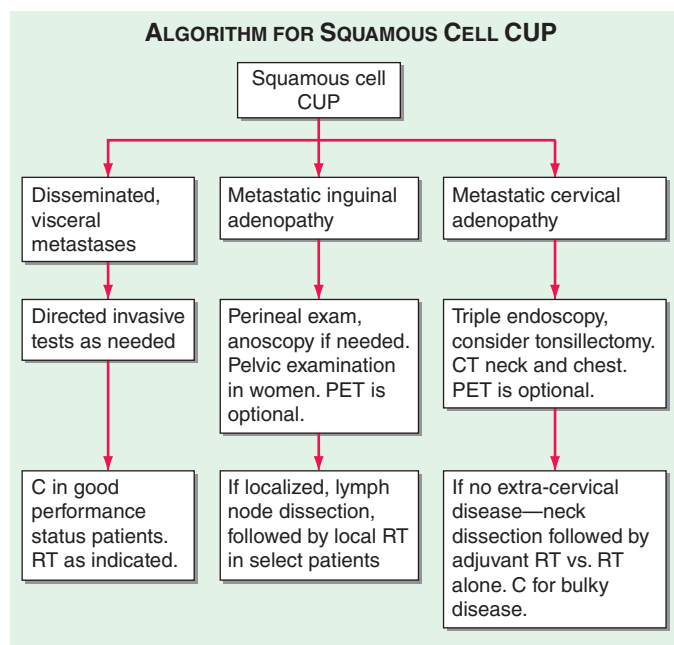


FIGURE 120e-3 Treatment algorithm for squamous cell carcinoma of unknown primary (CUP). C, chemotherapy; CT, computed tomography; PET, positron emission tomography; RT, radiation.

retrospective study of 258 women with peritoneal carcinomatosis who had undergone cytoreductive surgery and chemotherapy, 22% of patients had a complete response to chemotherapy; the median survival duration was 18 months (range 11–24 months). However, not all peritoneal carcinomatosis in women is PPSC. Careful pathological evaluation can help diagnose a colon cancer profile (CDX-2+, CK-20+, CK7–) or a pancreaticobiliary cancer or even a mislabeled peritoneal mesothelioma (calretinin positive).

POORLY DIFFERENTIATED CARCINOMA WITH MIDLINE ADENOPATHY

Men with poorly differentiated or undifferentiated carcinoma that presents as a midline adenopathy should be evaluated for extragonadal germ cell malignancy. If diagnosed and treated as such, they often experience a good response to treatment with platinum-based combination chemotherapy. Response rates of >50% have been noted, and long-term survival rates of 10–15% long have been reported. Older patients (especially smokers) who present with mediastinal adenopathy are more likely to have a lung or head-and-neck cancer profile.

NEUROENDOCRINE CARCINOMA

Low-grade neuroendocrine carcinoma often has an indolent course, and treatment decisions are based on symptoms and tumor bulk. Urine 5-HIAA and serum chromogranin may be elevated and can be followed as markers. Often the patient is treated with somatostatin analogues alone for hormone-related symptoms (diarrhea, flushing, nausea). Specific local therapies or systemic therapy would only be indicated if the patient is symptomatic with local pain secondary to significant growth of the metastasis or the hormone-related symptoms are not controlled with endocrine therapy. Patients with high-grade neuroendocrine carcinoma are treated as having small-cell lung cancer and are responsive to chemotherapy; 20–25% show a complete response, and up to 10% patients survive more than 5 years.

SQUAMOUS CELL CARCINOMA PRESENTING AS NECK ADENOPATHY

Patients with early-stage squamous cell carcinoma involving the cervical lymph nodes are candidates for node dissection and radiation therapy, which can result in long-term survival. The role of chemotherapy in these patients is undefined, although chemoradiation therapy or induction chemotherapy is often used and is beneficial in bulky N2/N3 lymph node disease.

SOLITARY METASTATIC SITE

Patients with solitary metastases can also experience good treatment outcomes. Some patients who present with locoregional disease are candidates for aggressive trimodality management; both prolonged disease-free interval and occasionally cure are possible.

MEN WITH BLASTIC SKELETAL METASTASES AND ELEVATED PSA

Blastic bone-only metastasis is a rare presentation, and elevated serum PSA or tumor staining with PSA may provide confirmatory evidence of prostate cancer in these patients. Those with elevated levels are candidates for hormonal therapy for prostate cancer, although it is important to rule out other primary tumors (lung most common).

MANAGEMENT OF DISSEMINATED CUP

Patients who present with liver, brain, and adrenal metastatic disease usually have a poor prognosis. Patients with nonserous papillary primary peritoneal carcinomatosis can have a large differential diagnosis, which is mainly of gastrointestinal profile and includes gastric, appendiceal, colon, and pancreaticobiliary profiles.

Traditionally, platinum-based combination chemotherapy regimens have been used to treat CUP. Several broadly used regimens have been studied in the last two decades; these include paclitaxel-carboplatin, gemcitabine-cisplatin, gemcitabine-oxaliplatin, and irinotecan and fluoropyrimidine-based therapies. These chemotherapeutic agents used as empiric regimens have shown a response rate of 25–40%, and their use obtains median survival times of 6–13 months.

Outside of favorable subsets, there is a small group of patients with a “definitive” IHC. These patients usually have a single diagnosis based on their clinicopathologic presentation and are often treated for the putative primary tumor. This does not guarantee a response, although it increases the probability of response when select drugs are chosen from a class of drugs known to work in that cancer type. Patients who do not fall into those categories are candidates for broad-spectrum platinum-based regimens, clinical trials, and additional trial-based genomic and proteomic tests. Today, we do not have effective drugs for several CUP cancer profiles, and treatments overlap for some cancers. However, as novel therapies are developed for additional known cancers, tissue of origin and assessment of molecular features of the tumor will be important and might direct more selective treatment.

SUMMARY

Patients with CUP should undergo a directed diagnostic search for the primary tumor on the basis of clinical and pathologic data. Subsets of patients have prognostically favorable disease, as defined by clinical or histologic criteria, and may substantially benefit from aggressive treatment and expect prolonged survival. However, for most patients who present with advanced CUP, the prognosis remains poor with early resistance to available cytotoxic therapy. The current focus has shifted away from empirical chemotherapeutic trials to understanding the metastatic phenotype, tissue of origin profiling, and evaluating molecular targets in CUP patients.

121 Paraneoplastic Syndromes: Endocrinologic/Hematologic

J. Larry Jameson, Dan L. Longo

Neoplastic cells can produce a variety of products that can stimulate hormonal, hematologic, dermatologic, and neurologic responses. *Paraneoplastic syndromes* is the term used to refer to the disorders that accompany benign or malignant tumors but are not directly related to mass effects or invasion. Tumors of neuroendocrine origin, such as small-cell lung carcinoma (SCLC) and carcinoids, produce a wide array of peptide hormones and are common causes of paraneoplastic syndromes. However, almost every type of tumor has the potential to produce hormones or to induce cytokine and immunologic responses. Careful studies of the prevalence of paraneoplastic syndromes indicate that they are more common than is generally appreciated. The signs, symptoms, and metabolic alterations associated with paraneoplastic disorders may be overlooked in the context of a malignancy and its treatment. Consequently, atypical clinical manifestations in a patient with cancer should prompt consideration of a paraneoplastic syndrome. The most common endocrinologic and hematologic syndromes associated with underlying neoplasia will be discussed here.

ENDOCRINE PARANEOPLASTIC SYNDROMES

Etiology Hormones can be produced from eutopic or ectopic sources. *Eutopic* refers to the expression of a hormone from its normal tissue of origin, whereas *ectopic* refers to hormone production from an atypical tissue source. For example, adrenocorticotropic hormone (ACTH) is expressed eutopically by the corticotrope cells of the anterior pituitary, but it can be expressed ectopically in SCLC. Many hormones are produced at low levels from a wide array of tissues in addition to the classic endocrine source. Thus, ectopic expression is often a quantitative change rather than an absolute change in tissue expression. Nevertheless, the term *ectopic expression* is firmly entrenched and conveys the abnormal physiology associated with hormone production by neoplastic cells. In addition to high levels of hormones, ectopic expression typically is characterized by abnormal regulation of hormone production (e.g., defective feedback control) and peptide processing (resulting in large, unprocessed precursors).

A diverse array of molecular mechanisms has been suggested to cause ectopic hormone production. In rare instances, genetic rearrangements explain aberrant hormone expression. For example, translocation of the parathyroid hormone (*PTH*) gene can result in high levels of *PTH* expression in tissues other than the parathyroid gland because the genetic rearrangement brings the *PTH* gene under the control of atypical regulatory elements. A related phenomenon is well documented in many forms of leukemia and lymphoma, in which somatic genetic rearrangements confer a growth advantage and alter cellular differentiation and function (Chap. 134). Although genetic rearrangements cause selected cases of ectopic hormone production, this mechanism is rare, as many tumors are associated with excessive production of numerous peptides. Cellular dedifferentiation probably underlies most cases of ectopic hormone production. Many cancers are poorly differentiated, and certain tumor products, such as human chorionic gonadotropin (hCG), parathyroid hormone–related protein (PTHrP), and α fetoprotein, are characteristic of gene expression at earlier developmental stages. In contrast, the propensity of certain cancers to produce particular hormones (e.g., squamous cell carcinomas produce PTHrP) suggests that dedifferentiation is partial or that selective pathways are derepressed. These expression profiles probably reflect epigenetic modifications that alter transcriptional repression, microRNA expression, and other pathways that govern cell differentiation.

In SCLC, the pathway of differentiation has been relatively well defined. The neuroendocrine phenotype is dictated in part by the basic-helix-loop-helix (bHLH) transcription factor human achaete-scute homologue 1 (hASH-1), which is expressed at abnormally high

TABLE 121-1 PARANEOPLASTIC SYNDROMES CAUSED BY ECTOPIC HORMONE PRODUCTION

Paraneoplastic Syndrome	Ectopic Hormone	Typical Tumor Types ^a
Common		
Hypercalcemia of malignancy	Parathyroid hormone–related protein (PTHrP)	Squamous cell (head and neck, lung, skin), breast, genitourinary, gastrointestinal
	1,25-dihydroxyvitamin D	Lymphomas
	Parathyroid hormone (PTH) (rare)	Lung, ovary
	Prostaglandin E ₂ (PGE ₂) (rare)	Renal, lung
Syndrome of inappropriate antidiuretic hormone secretion (SIADH)	Vasopressin	Lung (squamous, small cell), gastrointestinal, genitourinary, ovary
Cushing's syndrome	Adrenocorticotrophic hormone (ACTH)	Lung (small cell, bronchial carcinoid, adenocarcinoma, squamous), thymus, pancreatic islet, medullary thyroid carcinoma
	Corticotropin-releasing hormone (CRH) (rare)	Pancreatic islet, carcinoid, lung, prostate
	Ectopic expression of gastric inhibitory peptide (GIP), luteinizing hormone (LH)/human chorionic gonadotropin (hCG), other G protein–coupled receptors (rare)	Macronodular adrenal hyperplasia
Less Common		
Non–islet cell hypoglycemia	Insulin-like growth factor type II (IGF-II)	Mesenchymal tumors, sarcomas, adrenal, hepatic, gastrointestinal, kidney, prostate
	Insulin (rare)	Cervix (small-cell carcinoma)
Male feminization	hCG ^b	Testis (embryonal, seminomas), germinomas, choriocarcinoma, lung, hepatic, pancreatic islet
Diarrhea or intestinal hypermotility	Calcitonin ^c	Lung, colon, breast, medullary thyroid carcinoma
	Vasoactive intestinal peptide (VIP)	Pancreas, pheochromocytoma, esophagus
Rare		
Oncogenic osteomalacia	Phosphatonin (fibroblast growth factor 23 [FGF23])	Hemangiopericytomas, osteoblastomas, fibromas, sarcomas, giant cell tumors, prostate, lung
Acromegaly	Growth hormone–releasing hormone (GHRH)	Pancreatic islet, bronchial, and other carcinoids
	Growth hormone (GH)	Lung, pancreatic islet
Hyperthyroidism	Thyroid-stimulating hormone (TSH)	Hydatidiform mole, embryonal tumors, struma ovarii
Hypertension	Renin	Juxtaglomerular tumors, kidney, lung, pancreas, ovary

^aOnly the most common tumor types are listed. For most ectopic hormone syndromes, an extensive list of tumors has been reported to produce one or more hormones. ^bhCG is produced ectopically by trophoblastic tumors. Certain tumors produce disproportionate amounts of the hCG α or hCG β subunit. High levels of hCG rarely cause hyperthyroidism because of weak binding to the TSH receptor. ^cCalcitonin is produced ectopically by medullary thyroid carcinoma and is used as a tumor marker.

levels in SCLC associated with ectopic ACTH. The activity of hASH-1 is inhibited by hairy enhancer of split 1 (HES-1) and by Notch proteins, which also are capable of inducing growth arrest. Thus, abnormal expression of these developmental transcription factors appears to provide a link between cell proliferation and differentiation.

Ectopic hormone production would be merely an epiphenomenon associated with cancer if it did not result in clinical manifestations. Excessive and unregulated production of hormones such as ACTH, PTHrP, and vasopressin can lead to substantial morbidity and complicate the cancer treatment plan. Moreover, the paraneoplastic endocrinopathies may be a presenting clinical feature of underlying malignancy and prompt the search for an unrecognized tumor.

A large number of paraneoplastic endocrine syndromes have been described, linking overproduction of particular hormones with specific types of tumors. However, certain recurring syndromes emerge from this group (Table 121-1). The most common paraneoplastic endocrine syndromes include hypercalcemia from overproduction of PTHrP and other factors, hyponatremia from excess vasopressin, and Cushing's syndrome from ectopic ACTH.

HYPERCALCEMIA CAUSED BY ECTOPIC PRODUCTION OF PTHrP

(See also Chap. 424)

Etiology Humoral hypercalcemia of malignancy (HHM) occurs in up to 20% of patients with cancer. HHM is most common in cancers of the lung, head and neck, skin, esophagus, breast, and genitourinary tract and in multiple myeloma and lymphomas. There are several distinct humoral causes of HHM, but it is caused most commonly by overproduction of PTHrP. In addition to acting as a circulating

humoral factor, bone metastases (e.g., breast, multiple myeloma) may produce PTHrP, leading to local osteolysis and hypercalcemia. PTHrP may also affect the initiation and progression of tumors by acting through pro-survival and chemokine pathways.

PTHrP is structurally related to PTH and binds to the PTH receptor, explaining the similar biochemical features of HHM and hyperparathyroidism. PTHrP plays a key role in skeletal development and regulates cellular proliferation and differentiation in other tissues, including skin, bone marrow, breast, and hair follicles. The mechanism of PTHrP induction in malignancy is incompletely understood; however, tumor-bearing tissues commonly associated with HHM normally produce PTHrP during development or cell renewal. PTHrP expression is stimulated by hedgehog pathways and Gli transcription factors that are active in many malignancies. Transforming growth factor β (TGF- β), which is produced by many tumors, also stimulates PTHrP, in part by activating the Gli pathway. Mutations in certain oncogenes, such as *Ras*, also can activate PTHrP expression. In adult T cell lymphoma, the transactivating Tax protein produced by human T cell lymphotropic virus 1 (HTLV-1) stimulates PTHrP promoter activity. Metastatic lesions to bone are more likely to produce PTHrP than are metastases in other tissues, suggesting that bone produces factors (e.g., TGF- β) that enhance PTHrP production or that PTHrP-producing metastases have a selective growth advantage in bone. PTHrP activates the pro-survival AKT pathway and the chemokine receptor CXCR4. Thus, PTHrP production can be stimulated by mutations in oncogenes, altered expression of viral or cellular transcription factors, and local growth factors. In addition to its role in HHM, the PTHrP pathway may also provide a potential target for therapeutic intervention to impede cancer growth.

Another relatively common cause of HHM is excess production of 1,25-dihydroxyvitamin D. Like granulomatous disorders associated with hypercalcemia, lymphomas can produce an enzyme that converts 25-hydroxyvitamin D to the more active 1,25-dihydroxyvitamin D, leading to enhanced gastrointestinal calcium absorption. Other causes of HHM include tumor-mediated production of osteolytic cytokines and inflammatory mediators.

Clinical Manifestations The typical presentation of HHM is a patient with a known malignancy who is found to be hypercalcemic on routine laboratory tests. Less often, hypercalcemia is the initial presenting feature of malignancy. Particularly when calcium levels are markedly increased (>3.5 mmol/L [>14 mg/dL]), patients may experience fatigue, mental status changes, dehydration, or symptoms of nephrolithiasis.

Diagnosis Features that favor HHM, as opposed to primary hyperparathyroidism, include known malignancy, recent onset of hypercalcemia, and very high serum calcium levels. Like hyperparathyroidism, hypercalcemia caused by PTHrP is accompanied by hypercalciuria and hypophosphatemia. Patients with HHM typically have metabolic alkalosis rather than hyperchloremic acidosis, as is seen in hyperparathyroidism. Measurement of PTH is useful to exclude primary hyperparathyroidism; the PTH level should be suppressed in HHM. An elevated PTHrP level confirms the diagnosis, and it is increased in ~80% of hypercalcemic patients with cancer. 1,25-Dihydroxyvitamin D levels may be increased in patients with lymphoma.

TREATMENT HUMORAL HYPERCALCEMIA OF MALIGNANCY

The management of HHM begins with removal of excess calcium in the diet, medications, or IV solutions. Saline rehydration (typically 200–500 mL/h) is used to dilute serum calcium and promote calciuresis; exercise caution in patients with cardiac, hepatic, or renal insufficiency. Forced diuresis with furosemide (20–80 mg IV in escalating doses) or other loop diuretics can enhance calcium excretion but provides relatively little value except in life-threatening hypercalcemia. When used, loop diuretics should be administered only after complete rehydration and with careful monitoring of fluid balance. Oral phosphorus (e.g., 250 mg Neutra-Phos 3–4 times daily) should be given until serum phosphorus is >1 mmol/L (>3 mg/dL). Bisphosphonates such as pamidronate (60–90 mg IV), zoledronate (4–8 mg IV), and etidronate (7.5 mg/kg per day PO for 3–7 consecutive days) can reduce serum calcium within 1–2 days and suppress calcium release for several weeks. Bisphosphonate infusions can be repeated, or oral bisphosphonates can be used for chronic treatment. Dialysis should be considered in severe hypercalcemia when saline hydration and bisphosphonate treatments are not possible or are too slow in onset. Previously used agents such as calcitonin and mithramycin have little utility now that bisphosphonates are available. Calcitonin (2–8 U/kg SC every 6–12 h) should be considered when rapid correction of severe hypercalcemia is needed. Hypercalcemia associated with lymphomas, multiple myeloma, or leukemia may respond to glucocorticoid treatment (e.g., prednisone 40–100 mg PO in four divided doses).

ECTOPIC VASOPRESSIN: TUMOR-ASSOCIATED SIADH (See also Chap. 63)

Etiology Vasopressin is an antidiuretic hormone normally produced by the posterior pituitary gland. Ectopic vasopressin production by tumors is a common cause of the syndrome of inappropriate antidiuretic hormone (SIADH), occurring in at least half of patients with SCLC. SIADH also can be caused by a number of nonneoplastic conditions, including central nervous system (CNS) trauma, infections, and medications (Chap. 404). Compensatory responses to SIADH, such as decreased thirst, may mitigate the development of hyponatremia. However, with prolonged production of excessive vasopressin, the osmostat controlling thirst and hypothalamic vasopressin secretion may

become reset. In addition, intake of free water, orally or intravenously, can quickly worsen hyponatremia because of reduced renal diuresis.

Tumors with neuroendocrine features, such as SCLC and carcinoids, are the most common sources of ectopic vasopressin production, but it also occurs in other forms of lung cancer and with CNS lesions, head and neck cancer, and genitourinary, gastrointestinal, and ovarian cancers. The mechanism of activation of the vasopressin gene in these tumors is unknown but often involves concomitant expression of the adjacent oxytocin gene, suggesting derepression of this locus.

Clinical Manifestations Most patients with ectopic vasopressin secretion are asymptomatic and are identified because of the presence of hyponatremia on routine chemistry testing. Symptoms may include weakness, lethargy, nausea, confusion, depressed mental status, and seizures. The severity of symptoms reflects the rapidity of onset as well as the severity of hyponatremia. Hyponatremia usually develops slowly but may be exacerbated by the administration of IV fluids or the institution of new medications.

Diagnosis The diagnostic features of ectopic vasopressin production are the same as those of other causes of SIADH (Chaps. 63 and 404). Hyponatremia and reduced serum osmolality occur in the setting of an inappropriately normal or increased urine osmolality. Urine sodium excretion is normal or increased unless volume depletion is present. Other causes of hyponatremia should be excluded, including renal, adrenal, or thyroid insufficiency. Physiologic sources of vasopressin stimulation (CNS lesions, pulmonary disease, nausea), adaptive circulatory mechanisms (hypotension, heart failure, hepatic cirrhosis), and medications, including many chemotherapeutic agents, also should be considered as possible causes of hyponatremia. Vasopressin measurements are not usually necessary to make the diagnosis.

TREATMENT ECTOPIC VASOPRESSIN: TUMOR-ASSOCIATED SIADH

Most patients with ectopic vasopressin production develop hyponatremia over several weeks or months. The disorder should be corrected gradually unless mental status is altered or there is risk of seizures. Treatment of the underlying malignancy may reduce ectopic vasopressin production, but this response is slow if it occurs at all. Fluid restriction to less than urine output, plus insensible losses, is often sufficient to correct hyponatremia partially. However, strict monitoring of the amount and types of liquids consumed or administered intravenously is required for fluid restriction to be effective. Salt tablets and saline are not helpful unless volume depletion is also present. Demeclocycline (150–300 mg orally three to four times daily) can be used to inhibit vasopressin action on the renal distal tubule, but its onset of action is relatively slow (1–2 weeks). Conivaptan, a nonpeptide V_2 -receptor antagonist, can be administered either PO (20–120 mg bid) or IV (10–40 mg) and is particularly effective when used in combination with fluid restriction in euolemic hyponatremia. Tolvaptan (15 mg PO daily) is another vasopressin antagonist. The dose can be increased to 30–60 mg/d based on response. Severe hyponatremia ($\text{Na} < 115$ meq/L) or mental status changes may require treatment with hypertonic (3%) or normal saline infusion together with furosemide to enhance free water clearance. The rate of sodium correction should be slow (0.5–1 meq/L per hour) to prevent rapid fluid shifts and the possible development of central pontine myelinolysis.

CUSHING'S SYNDROME CAUSED BY ECTOPIC ACTH PRODUCTION (See also Chap. 406)

Etiology Ectopic ACTH production accounts for 10–20% of cases of Cushing's syndrome. The syndrome is particularly common in neuroendocrine tumors. SCLC is the most common cause of ectopic ACTH, followed by bronchial and thymic carcinoids, islet cell tumors, other carcinoids, and pheochromocytomas. Ectopic ACTH production is caused by increased expression of the proopiomelanocortin (POMC)

gene, which encodes ACTH, along with melanocyte-stimulating hormone (MSH), β lipotropin, and several other peptides. In many tumors, there is abundant but aberrant expression of the *POMC* gene from an internal promoter, proximal to the third exon, which encodes ACTH. However, because this product lacks the signal sequence necessary for protein processing, it is not secreted. Increased production of ACTH arises instead from less abundant, but unregulated, *POMC* expression from the same promoter site used in the pituitary. However, because the tumors lack many of the enzymes needed to process the *POMC* polypeptide, it is typically released as multiple large, biologically inactive fragments along with relatively small amounts of fully processed, active ACTH.

Rarely, corticotropin-releasing hormone (CRH) is produced by pancreatic islet cell tumors, SCLC, medullary thyroid cancer, carcinoids, or prostate cancer. When levels are high enough, CRH can cause pituitary corticotrope hyperplasia and Cushing's syndrome. Tumors that produce CRH sometimes also produce ACTH, raising the possibility of a paracrine mechanism for ACTH production.

A distinct mechanism for ACTH-independent Cushing's syndrome involves ectopic expression of various G protein-coupled receptors in the adrenal nodules. Ectopic expression of the gastric inhibitory peptide (GIP) receptor is the best-characterized example of this mechanism. In this case, meals induce GIP secretion, which inappropriately stimulates adrenal growth and glucocorticoid production.

Clinical Manifestations The clinical features of hypercortisolemia are detected in only a small fraction of patients with documented ectopic ACTH production. Patients with ectopic ACTH syndrome generally exhibit less marked weight gain and centripetal fat redistribution, probably because the exposure to excess glucocorticoids is relatively brief and because cachexia reduces the propensity for weight gain and fat deposition. The ectopic ACTH syndrome is associated with several clinical features that distinguish it from other causes of Cushing's syndrome (e.g., pituitary adenomas, adrenal adenomas, iatrogenic glucocorticoid excess). The metabolic manifestations of ectopic ACTH syndrome are dominated by fluid retention and hypertension, hypokalemia, metabolic alkalosis, glucose intolerance, and occasionally steroid psychosis. The very high ACTH levels often cause increased pigmentation, and melanocyte-stimulating hormone (MSH) activity derived from the *POMC* precursor peptide is also increased. The extraordinarily high glucocorticoid levels in patients with ectopic sources of ACTH can lead to marked skin fragility and easy bruising. In addition, the high cortisol levels often overwhelm the renal 11β -hydroxysteroid dehydrogenase type II enzyme, which normally inactivates cortisol and prevents it from binding to renal mineralocorticoid receptors. Consequently, in addition to the excess mineralocorticoids produced by ACTH stimulation of the adrenal gland, high levels of cortisol exert activity through the mineralocorticoid receptor, leading to severe hypokalemia.

Diagnosis The diagnosis of ectopic ACTH syndrome is usually not difficult in the setting of a known malignancy. Urine free cortisol levels fluctuate but are typically greater than two to four times normal, and the plasma ACTH level is usually >22 pmol/L (>100 pg/mL). A suppressed ACTH level excludes this diagnosis and indicates an ACTH-independent cause of Cushing's syndrome (e.g., adrenal or exogenous glucocorticoid). In contrast to pituitary sources of ACTH, most ectopic sources of ACTH do not respond to glucocorticoid suppression. Therefore, high-dose dexamethasone (8 mg PO) suppresses 8:00 A.M. serum cortisol (50% decrease from baseline) in $\sim 80\%$ of pituitary ACTH-producing adenomas but fails to suppress ectopic ACTH in $\sim 90\%$ of cases. Bronchial and other carcinoids are well-documented exceptions to these general guidelines, as these ectopic sources of ACTH may exhibit feedback regulation indistinguishable from pituitary adenomas, including suppression by high-dose dexamethasone, and ACTH responsiveness to adrenal blockade with metyrapone. If necessary, petrosal sinus catheterization can be used to evaluate a patient with ACTH-dependent Cushing's syndrome when the source of ACTH is unclear. After CRH stimulation, a 3:1 petrosal sinus:peripheral ACTH ratio strongly suggests a pituitary ACTH

source. Imaging studies (computed tomography or magnetic resonance imaging) are also useful in the evaluation of suspected carcinoid lesions, allowing biopsy and characterization of hormone production using special stains. If available, positron emission tomography or octreotide scanning may identify some sources of ACTH production.

TREATMENT CUSHING'S SYNDROME CAUSED BY ECTOPIC ACTH PRODUCTION

The morbidity associated with the ectopic ACTH syndrome can be substantial. Patients may experience depression or personality changes because of extreme cortisol excess. Metabolic derangements, including diabetes mellitus and hypokalemia, can worsen fatigue. Poor wound healing and predisposition to infections can complicate the surgical management of tumors, and opportunistic infections caused by organisms such as *Pneumocystis carinii* and mycoses are often the cause of death in patients with ectopic ACTH production. These patients likely have increased risk of venous thromboembolism reflecting the combination of malignancy and altered coagulation factor profiles. Depending on prognosis and treatment plans for the underlying malignancy, measures to reduce cortisol levels are often indicated. Treatment of the underlying malignancy may reduce ACTH levels but is rarely sufficient to reduce cortisol levels to normal. Adrenalectomy is not practical for most of these patients but should be considered during surgery for the malignancy or if the underlying tumor is not resectable and the prognosis is otherwise favorable (e.g., carcinoid). Medical therapy with ketoconazole (300–600 mg PO bid), metyrapone (250–500 mg PO every 6 h), mitotane (3–6 g PO in four divided doses, tapered to maintain low cortisol production), or other agents that block steroid synthesis or action is often the most practical strategy for managing the hypercortisolism associated with ectopic ACTH production. Glucocorticoid replacement should be provided to prevent adrenal insufficiency (**Chap. 406**). Unfortunately, many patients eventually progress despite medical blockade.

TUMOR-INDUCED HYPOGLYCEMIA CAUSED BY EXCESS PRODUCTION OF IGF-II

(See also **Chap. 420**) Mesenchymal tumors, hemangiopericytomas, hepatocellular tumors, adrenal carcinomas, and a variety of other large tumors have been reported to produce excessive amounts of insulin-like growth factor type II (IGF-II) precursor, which binds weakly to insulin receptors and more strongly to IGF-I receptors, leading to insulin-like actions. The gene encoding IGF-II resides on a chromosome 11p15 locus that is normally imprinted (that is, expression is exclusively from a single parental allele). Biallelic expression of the IGF-II gene occurs in a subset of tumors, suggesting loss of methylation and loss of imprinting as a mechanism for gene induction. In addition to increased IGF-II production, IGF-II bioavailability is increased due to complex alterations in circulating binding proteins. Increased IGF-II suppresses growth hormone (GH) and insulin, resulting in reduced IGF binding protein 3 (IGFBP-3), IGF-I, and acid-labile subunit (ALS). The reduction in ALS and IGFBP-3, which normally sequester IGF-II, causes it to be displaced to a small circulating complex that has greater access to insulin target tissues. For this reason, circulating IGF-II levels may not be markedly increased despite causing hypoglycemia. In addition to IGF-II-mediated hypoglycemia, tumors may occupy enough of the liver to impair gluconeogenesis.

In most cases, a tumor causing hypoglycemia is clinically apparent (usually >10 cm in size) and hypoglycemia develops in association with fasting. The diagnosis is made by documenting low serum glucose and suppressed insulin levels in association with symptoms of hypoglycemia. Serum IGF-II levels may not be increased (IGF-II assays may not detect IGF-II precursors). Increased IGF-II mRNA expression is found in most of these tumors. Any medications associated with hypoglycemia should be eliminated. Treatment of the underlying malignancy, if possible, may reduce the predisposition to hypoglycemia. Frequent meals and IV glucose, especially during sleep or fasting, are often

612 necessary to prevent hypoglycemia. Glucagon and glucocorticoids have also been used to enhance glucose production.

HUMAN CHORIONIC GONADOTROPIN

hCG is composed of α and β subunits and can be produced as intact hormone, which is biologically active, or as uncombined biologically inert subunits. Ectopic production of intact hCG occurs most often in association with testicular embryonal tumors, germ cell tumors, extragonadal germinomas, lung cancer, hepatoma, and pancreatic islet tumors. Eutopic production of hCG occurs with trophoblastic malignancies. hCG α subunit production is particularly common in lung cancer and pancreatic islet cancer. In men, high hCG levels stimulate steroidogenesis and aromatase activity in testicular Leydig cells, resulting in increased estrogen production and the development of gynecomastia. Precocious puberty in boys or gynecomastia in men should prompt measurement of hCG and consideration of a testicular tumor or another source of ectopic hCG production. Most women are asymptomatic. hCG is easily measured. Treatment should be directed at the underlying malignancy.

ONCOGENIC OSTEOMALACIA

Hypophosphatemic oncogenic osteomalacia, also called tumor-induced osteomalacia (TIO), is characterized by markedly reduced serum phosphorus and renal phosphate wasting, leading to muscle weakness, bone pain, and osteomalacia. Serum calcium and PTH levels are normal, and 1,25-dihydroxyvitamin D is low. Oncogenic osteomalacia is usually caused by benign mesenchymal tumors, such as hemangiopericytomas, fibromas, and giant cell tumors, often of the skeletal extremities or head. It has also been described in sarcomas and in patients with prostate and lung cancer. Resection of the tumor reverses the disorder, confirming its humoral basis. The circulating phosphaturic factor is called *phosphatonin*—a factor that inhibits renal tubular reabsorption of phosphate and renal conversion of 25-hydroxyvitamin D to 1,25-dihydroxyvitamin D. Phosphatonin has been identified as fibroblast growth factor 23 (FGF23). FGF23 levels are increased in some, but not all, patients with osteogenic osteomalacia. FGF23 forms a ternary complex with the klotho protein and renal FGF receptors to reduce renal phosphate reabsorption. Treatment involves removal of the tumor, if possible, and supplementation with phosphate and vitamin D. Octreotide treatment reduces phosphate wasting in some patients with tumors that express somatostatin receptor subtype 2. Octreotide scans may also be useful in detecting these tumors.

HEMATOLOGIC SYNDROMES

The elevation of granulocyte, platelet, and eosinophil counts in most patients with myeloproliferative disorders is caused by the proliferation of the myeloid elements due to the underlying disease rather than to a paraneoplastic syndrome. The paraneoplastic hematologic syndromes in patients with solid tumors are less well characterized than are the endocrine syndromes because the ectopic hormone(s) or cytokines responsible have not been identified in most of these tumors (Table 121-2). The extent of the paraneoplastic syndromes parallels the course of the cancer.

ERYTHROCYTOSIS

Ectopic production of erythropoietin by cancer cells causes most paraneoplastic erythrocytosis. The ectopically produced erythropoietin stimulates the production of red blood cells (RBCs) in the bone marrow and raises the hematocrit. Other lymphokines and hormones produced by cancer cells may stimulate erythropoietin release but have not been proved to cause erythrocytosis.

Most patients with erythrocytosis have an elevated hematocrit (>52% in men, >48% in women) that is detected on a routine blood count. Approximately 3% of patients with renal cell cancer, 10% of patients with hepatoma, and 15% of patients with cerebellar hemangioblastomas have erythrocytosis. In most cases, the erythrocytosis is asymptomatic.

Patients with erythrocytosis due to a renal cell cancer, hepatoma, or CNS cancer should have measurement of red cell mass. If the red cell

TABLE 121-2 PARANEOPLASTIC HEMATOLOGIC SYNDROMES

Syndrome	Proteins	Cancers Typically Associated with Syndrome
Erythrocytosis	Erythropoietin	Renal cancers, hepatocarcinoma, cerebellar hemangioblastomas
Granulocytosis	G-CSF, GM-CSF, IL-6	Lung cancer, gastrointestinal cancer, ovarian cancer, genitourinary cancer, Hodgkin's disease
Thrombocytosis	IL-6	Lung cancer, gastrointestinal cancer, breast cancer, ovarian cancer, lymphoma
Eosinophilia	IL-5	Lymphoma, leukemia, lung cancer
Thrombophlebitis	Unknown	Lung cancer, pancreatic cancer, gastrointestinal cancer, breast cancer, genitourinary cancer, ovarian cancer, prostate cancer, lymphoma

Abbreviations: G-CSF, granulocyte colony-stimulating factor; GM-CSF, granulocyte-macrophage colony-stimulating factor; IL, interleukin.

mass is elevated, the serum erythropoietin level should be measured. Patients with an appropriate cancer, elevated erythropoietin levels, and no other explanation for erythrocytosis (e.g., hemoglobinopathy that causes increased O₂ affinity; Chap. 77) have the paraneoplastic syndrome.

TREATMENT ERYTHROCYTOSIS

Successful resection of the cancer usually resolves the erythrocytosis. If the tumor cannot be resected or treated effectively with radiation therapy or chemotherapy, phlebotomy may control any symptoms related to erythrocytosis.

GRANULOCYTOSIS

Approximately 30% of patients with solid tumors have granulocytosis (granulocyte count >8000/ μ L). In about half of patients with granulocytosis and cancer, the granulocytosis has an identifiable nonparaneoplastic etiology (infection, tumor necrosis, glucocorticoid administration, etc.). The other patients have proteins in urine and serum that stimulate the growth of bone marrow cells. Tumors and tumor cell lines from patients with lung, ovarian, and bladder cancers have been documented to produce granulocyte colony-stimulating factor (G-CSF), granulocyte-macrophage colony-stimulating factor (GM-CSF), and/or interleukin 6 (IL-6). However, the etiology of granulocytosis has not been characterized in most patients.

Patients with granulocytosis are nearly all asymptomatic, and the differential white blood cell count does not have a shift to immature forms of neutrophils. Granulocytosis occurs in 40% of patients with lung and gastrointestinal cancers, 20% of patients with breast cancer, 30% of patients with brain tumors and ovarian cancers, 20% of patients with Hodgkin's disease, and 10% of patients with renal cell carcinoma. Patients with advanced-stage disease are more likely to have granulocytosis than are those with early-stage disease.

Paraneoplastic granulocytosis does not require treatment. The granulocytosis resolves when the underlying cancer is treated.

THROMBOCYTOSIS

Some 35% of patients with thrombocytosis (platelet count >400,000/ μ L) have an underlying diagnosis of cancer. IL-6, a candidate molecule for the etiology of paraneoplastic thrombocytosis, stimulates the production of platelets in vitro and in vivo. Some patients with cancer and thrombocytosis have elevated levels of IL-6 in plasma. Another candidate molecule is thrombopoietin, a peptide hormone that stimulates megakaryocyte proliferation and platelet production. The etiology of thrombocytosis has not been established in most cases.

Patients with thrombocytosis are nearly all asymptomatic. Thrombocytosis is not clearly linked to thrombosis in patients with

cancer. Thrombocytosis is present in 40% of patients with lung and gastrointestinal cancers; 20% of patients with breast, endometrial, and ovarian cancers; and 10% of patients with lymphoma. Patients with thrombocytosis are more likely to have advanced-stage disease and have a poorer prognosis than do patients without thrombocytosis. In ovarian cancer, IL-6 has been shown to directly promote tumor growth. Paraneoplastic thrombocytosis does not require treatment other than treatment of the underlying tumor.

EOSINOPHILIA

Eosinophilia is present in ~1% of patients with cancer. Tumors and tumor cell lines from patients with lymphomas or leukemia may produce IL-5, which stimulates eosinophil growth. Activation of IL-5 transcription in lymphomas and leukemias may involve translocation of the long arm of chromosome 5, to which the genes for IL-5 and other cytokines map.

Patients with eosinophilia are typically asymptomatic. Eosinophilia is present in 10% of patients with lymphoma, 3% of patients with lung cancer, and occasional patients with cervical, gastrointestinal, renal, and breast cancer. Patients with markedly elevated eosinophil counts (>5000/ μ L) can develop shortness of breath and wheezing. A chest radiograph may reveal diffuse pulmonary infiltrates from eosinophil infiltration and activation in the lungs.

TREATMENT EOSINOPHILIA

Definitive treatment is directed at the underlying malignancy: Tumors should be resected or treated with radiation or chemotherapy. In most patients who develop shortness of breath related to eosinophilia, symptoms resolve with the use of oral or inhaled glucocorticoids. IL-5 antagonists exist but have not been evaluated in this clinical setting.

THROMBOPHLEBITIS

Deep venous thrombosis and pulmonary embolism are the most common thrombotic conditions in patients with cancer. Migratory or recurrent thrombophlebitis may be the initial manifestation of cancer. Nearly 15% of patients who develop deep venous thrombosis or pulmonary embolism have a diagnosis of cancer (Chap. 142). The coexistence of peripheral venous thrombosis with visceral carcinoma, particularly pancreatic cancer, is called *Trousseau's syndrome*.

Pathogenesis Patients with cancer are predisposed to thromboembolism because they are often at bed rest or immobilized, and tumors may obstruct or slow blood flow. Postoperative deep venous thrombosis is twice as common in cancer patients who undergo surgery. Chronic IV catheters also predispose to clotting. In addition, clotting may be promoted by release of procoagulants or cytokines from tumor cells or associated inflammatory cells or by platelet adhesion or aggregation. The specific molecules that promote thromboembolism have not been identified.

Chemotherapeutic agents, particularly those associated with endothelial damage, can induce venous thrombosis. The annual risk of venous thrombosis in patients with cancer receiving chemotherapy is about 11%, sixfold higher than the risk in the general population. Bleomycin, L-asparaginase, thalidomide analogues, cisplatin-based regimens, and high doses of busulfan and carmustine are all associated with an increased risk.

In addition to cancer and its treatment causing secondary thrombosis, primary thrombophilic diseases may be associated with cancer. For example, the antiphospholipid antibody syndrome is associated with a wide range of pathologic manifestations (Chap. 379). About 20% of patients with this syndrome have cancers. Among patients with cancer and antiphospholipid antibodies, 35–45% develop thrombosis.

Clinical Manifestations Patients with cancer who develop deep venous thrombosis usually develop swelling or pain in the leg, and physical examination reveals tenderness, warmth, and redness. Patients who present with pulmonary embolism develop dyspnea, chest pain, and

syncope, and physical examination shows tachycardia, cyanosis, and hypotension. Some 5% of patients with no history of cancer who have a diagnosis of deep venous thrombosis or pulmonary embolism will have a diagnosis of cancer within 1 year. The most common cancers associated with thromboembolic episodes include lung, pancreatic, gastrointestinal, breast, ovarian, and genitourinary cancers; lymphomas; and brain tumors. Patients with cancer who undergo surgical procedures requiring general anesthesia have a 20–30% risk of deep venous thrombosis.

Diagnosis The diagnosis of deep venous thrombosis in patients with cancer is made by impedance plethysmography or bilateral compression ultrasonography of the leg veins. Patients with a noncompressible venous segment have deep venous thrombosis. If compression ultrasonography is normal and there is a high clinical suspicion for deep venous thrombosis, venography should be done to look for a luminal filling defect. Elevation of D-dimer is not as predictive of deep venous thrombosis in patients with cancer as it is in patients without cancer; elevations are seen in people over age 65 years without concomitant evidence of thrombosis, probably as a consequence of increased thrombin deposition and turnover in aging.

Patients with symptoms and signs suggesting a pulmonary embolism should be evaluated with a chest radiograph, electrocardiogram, arterial blood gas analysis, and ventilation-perfusion scan. Patients with mismatched segmental perfusion defects have a pulmonary embolus. Patients with equivocal ventilation-perfusion findings should be evaluated as described above for deep venous thrombosis in their legs. If deep venous thrombosis is detected, they should be anticoagulated. If deep venous thrombosis is not detected, they should be considered for a pulmonary angiogram.

Patients without a diagnosis of cancer who present with an initial episode of thrombophlebitis or pulmonary embolus need no additional tests for cancer other than a careful history and physical examination. In light of the many possible primary sites, diagnostic testing in asymptomatic patients is wasteful. However, if the clot is refractory to standard treatment or is in an unusual site or if the thrombophlebitis is migratory or recurrent, efforts to find an underlying cancer are indicated.

TREATMENT THROMBOPHLEBITIS

Patients with cancer and a diagnosis of deep venous thrombosis or pulmonary embolism should be treated initially with IV unfractionated heparin or low-molecular-weight heparin for at least 5 days, and warfarin should be started within 1 or 2 days. The warfarin dose should be adjusted so that the international normalized ratio (INR) is 2–3. Patients with proximal deep venous thrombosis and a relative contraindication to heparin anticoagulation (hemorrhagic brain metastases or pericardial effusion) should be considered for placement of a filter in the inferior vena cava (Greenfield filter) to prevent pulmonary embolism. Warfarin should be administered for 3–6 months. An alternative approach is to use low-molecular-weight heparin for 6 months. Patients with cancer who undergo a major surgical procedure should be considered for heparin prophylaxis or pneumatic boots. Breast cancer patients undergoing chemotherapy and patients with implanted catheters should be considered for prophylaxis. Guidelines recommend that hospitalized patients with cancer and patients receiving a thalidomide analogue receive prophylaxis with low-molecular-weight heparin or low-dose aspirin. Use of prophylaxis routinely during chemotherapy is controversial and not recommended by the American Society of Clinical Oncology.

Cutaneous paraneoplastic syndromes are discussed in Chap. 72. Neurologic paraneoplastic syndromes are discussed in Chap. 122.

ACKNOWLEDGMENT

The authors acknowledge the contributions of Bruce E. Johnson to prior versions of this chapter.

122 Paraneoplastic Neurologic Syndromes and Autoimmune Encephalitis

Josep Dalmau, Myrna R. Rosenfeld

Paraneoplastic neurologic disorders (PNDs) are cancer-related syndromes that can affect any part of the nervous system (Table 122-1). They are caused by mechanisms other than metastasis or by any of the complications of cancer such as coagulopathy, stroke, metabolic and nutritional conditions, infections, and side effects of cancer therapy. In 60% of patients, the neurologic symptoms precede the cancer diagnosis. Clinically disabling PNDs occur in 0.5–1% of all cancer patients, but they affect 2–3% of patients with neuroblastoma or small-cell lung cancer (SCLC) and 30–50% of patients with thymoma or sclerotic myeloma.

PATHOGENESIS

Most PNDs are mediated by immune responses triggered by neuronal proteins (onconeural antigens) expressed by tumors. In PNDs of the central nervous system (CNS), many antibody-associated immune responses have been identified (Table 122-2). These antibodies react with the patient's tumor, and their detection in serum or cerebrospinal fluid (CSF) usually predicts the presence of cancer. When the antigens are intracellular, most syndromes are associated with extensive infiltrates of CD4+ and CD8+ T cells, microglial activation, gliosis, and variable neuronal loss. The infiltrating T cells are often in close contact with neurons undergoing degeneration, suggesting a primary pathogenic role. T cell-mediated cytotoxicity may contribute directly to cell death in these PNDs. Thus both humoral and cellular immune mechanisms participate in the pathogenesis of many PNDs. This complex immunopathogenesis may underlie the resistance of many of these conditions to therapy.

In contrast to the disorders associated with immune responses against intracellular antigens, those associated with antibodies to antigens expressed on the neuronal cell surface of the CNS or at the neuromuscular junction are more responsive to immunotherapy (Table 122-3, Fig. 122-1). These disorders occur with and without a cancer association and may affect children and young adults, and there is increasing evidence that they are mediated by the antibodies.

Other PNDs are likely immune-mediated, although their antigens are unknown. These include several syndromes of inflammatory neuropathies and myopathies. In addition, many patients with typical PND syndromes are antibody-negative.

TABLE 122-1 PARANEOPLASTIC SYNDROMES OF THE NERVOUS SYSTEM

Classic Syndromes: Usually Occur with Cancer Association	Nonclassic Syndromes: May Occur with and Without Cancer Association
Encephalomyelitis	Brainstem encephalitis
Limbic encephalitis	Stiff-person syndrome
Cerebellar degeneration (adults)	Necrotizing myelopathy
Opsoclonus-myoclonus	Motor neuron disease
Subacute sensory neuropathy	Guillain-Barré syndrome
Gastrointestinal paresis or pseudo-obstruction	Subacute and chronic mixed sensory-motor neuropathies
Dermatomyositis (adults)	Neuropathy associated with plasma cell dyscrasias and lymphoma
Lambert-Eaton myasthenic syndrome	Vasculitis of nerve
Cancer- or melanoma-associated retinopathy	Pure autonomic neuropathy
	Acute necrotizing myopathy
	Polymyositis
	Vasculitis of muscle
	Optic neuropathy
	BDUMP

Abbreviation: BDUMP, bilateral diffuse uveal melanocytic proliferation.

TABLE 122-2 ANTIBODIES TO INTRACELLULAR ANTIGENS, SYNDROMES, AND ASSOCIATED CANCERS

Antibody	Associated Neurologic Syndrome(s)	Tumors
Anti-Hu (ANNA1)	Encephalomyelitis, subacute sensory neuropathy	SCLC
Anti-Yo (PCA1)	Cerebellar degeneration	Ovary, breast
Anti-Ri (ANNA2)	Cerebellar degeneration, opsoclonus, brainstem encephalitis	Breast, gynecologic, SCLC
Anti-Tr	Cerebellar degeneration	Hodgkin's lymphoma
Anti-CRMP5 (CV2)	Encephalomyelitis, chorea, optic neuritis, uveitis, peripheral neuropathy	SCLC, thymoma, other
Anti-Ma proteins	Limbic, hypothalamic, brainstem encephalitis	Testicular (Ma2), other (Ma)
Anti-amphiphysin	Stiff-person syndrome, encephalomyelitis	Breast, SCLC
Recoverin, bipolar cell antibodies, others ^a	Cancer-associated retinopathy (CAR) Melanoma-associated retinopathy (MAR)	SCLC (CAR), melanoma (MAR)
Anti-GAD	Stiff-person, cerebellar syndromes, limbic encephalitis	Infrequent tumor association (thymoma)

^aA variety of target antigens have been identified.

Abbreviations: CRMP, collapsing response-mediator protein; SCLC, small-cell lung cancer.

TABLE 122-3 ANTIBODIES TO CELL SURFACE OR SYNAPTIC ANTIGENS, SYNDROMES, AND ASSOCIATED TUMORS

Antibody	Neurologic Syndrome	Tumor Type When Associated
Anti-AChR (muscle) ^a	Myasthenia gravis	Thymoma
Anti-AChR (neuronal) ^a	Autonomic ganglionopathy	SCLC
Anti-VGCC ^b	LEMS, cerebellar degeneration	SCLC
Anti-NMDAR ^c	Anti-NMDAR encephalitis	Teratoma in young women (children and men rarely have tumors)
Anti-LGI1 ^c	Limbic encephalitis, hyponatremia, faciobrachial tonic or dystonic seizures	Rarely thymoma
Anti-Caspr2 ^c	Morvan's syndrome, neuro-myotonia	Thymoma, prostate cancer
Anti-GABA _B R ^d	Limbic encephalitis, seizures	SCLC, neuroendocrine
Anti-GABA _A R ^d	Encephalitis with prominent seizures and status epilepticus; less often opsoclonus and stiff-person syndrome	Rarely thymoma
Anti-AMPA ^d	Limbic encephalitis with relapses	SCLC, thymoma, breast
Glycine receptor ^d	Encephalomyelitis with rigidity, stiff-person syndrome	Rarely, thymoma, lung cancer
Anti-DPPX ^d	Agitation, myoclonus, tremor, seizures, hyperekplexia, encephalomyelitis with rigidity	No cancer, but frequent diarrhea or cachexia suggesting paraneoplasia

^aA direct pathogenic role of these antibodies has been demonstrated. ^bAnti-VGCC antibodies are pathogenic for LEMS. ^cPreviously named voltage-gated potassium channel antibodies (VGKC); currently included under the term VGKC-complex proteins. Of note, the significance of antibodies to VGKC-complex proteins other than LGI1 and Caspr2 is uncertain (the antigens are unknown, and the response to immunotherapy is variable). ^dThese antibodies are strongly suspected to be pathogenic.

Abbreviations: AChR, acetylcholine receptor; AMPAR, α-amino-3-hydroxy-5-methylisoxazole-4-propionic acid receptor; Caspr2, contactin-associated protein-like 2; DPPX, dipeptidyl-peptidase-like protein-6; GABA_BR, γ-aminobutyric acid B receptor; GAD, glutamic acid decarboxylase; LEMS, Lambert-Eaton myasthenic syndrome; LGI1, leucine-rich glioma-inactivated 1; NMDAR, N-methyl-D-aspartate receptor; SCLC, small-cell lung cancer; VGCC, voltage-gated calcium channel.

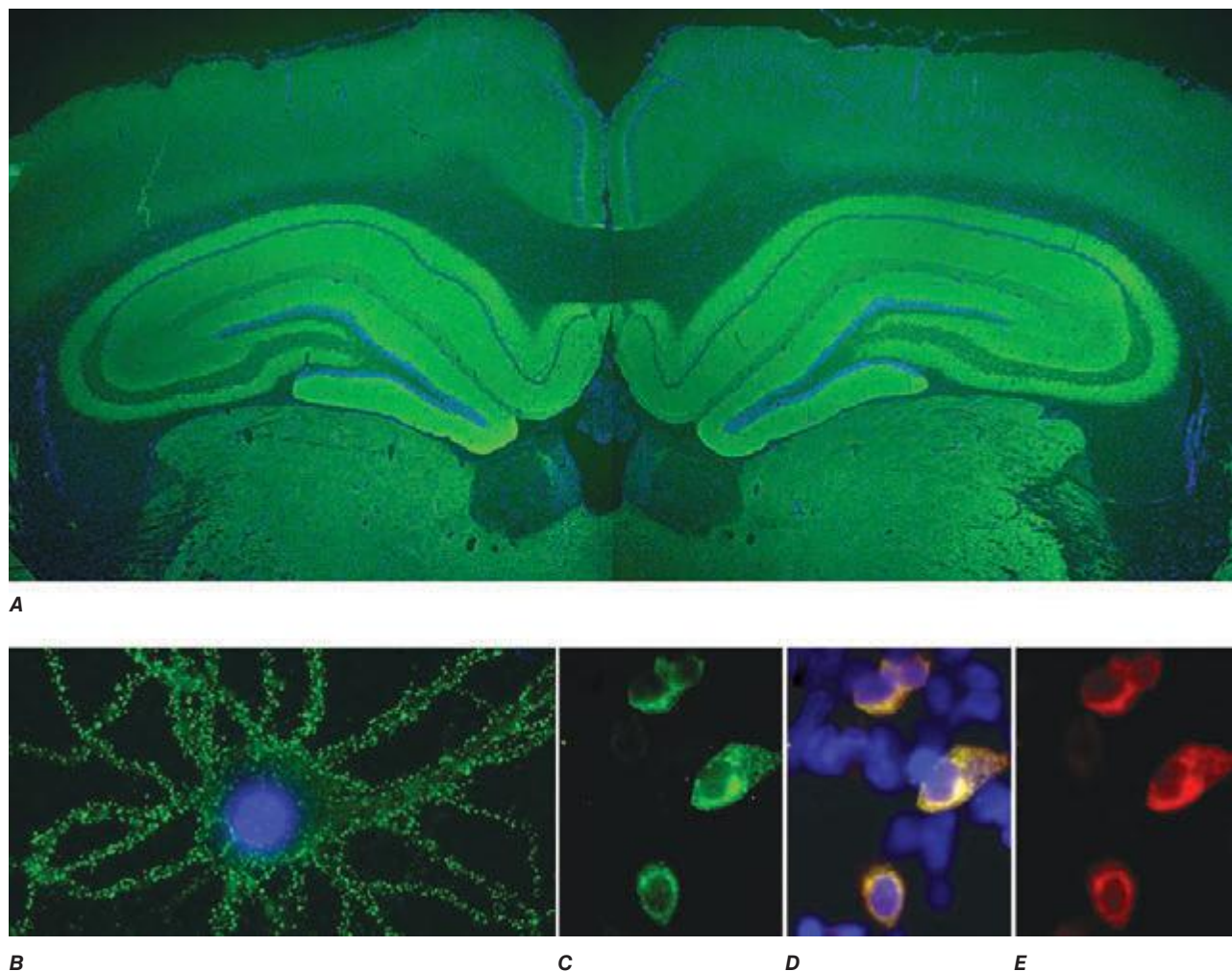


FIGURE 122-1 Antibodies to the GluN1 subunit of the *N*-methyl-D-aspartate (NMDA) receptor in a patient with anti-NMDA receptor encephalitis and ovarian teratoma. (A) Coronal section of rat brain immunolabeled (green fluorescence) with the patient's antibodies. The reactivity predominates in the hippocampus, which is highly enriched in NMDA receptors. (B) This image shows the antibody reactivity with cultures of rat hippocampal neurons; the intense green immunolabeling is due to the antibodies against the GluN1 subunit of NMDA receptors. (C–E) Images of HEK cells (a human kidney cell line) transfected to express NMDA receptors, showing reactivity with patient's antibodies (C) and with a commercial monoclonal antibody against NMDA receptors (E); the patient's antibody reactivity co-labels only the cells that express NMDA receptors (D). (From J Dalmau et al: *Lancet Neurol* 7:1091, 2008; with permission.)

For still other PNDs, the cause remains quite obscure. These include, among others, several neuropathies that occur in the terminal stages of cancer and a number of neuropathies associated with plasma cell dyscrasias or lymphoma without evidence of inflammatory infiltrates or deposits of immunoglobulin, cryoglobulin, or amyloid.

APPROACH TO THE PATIENT: Paraneoplastic Neurologic Disorders

Three key concepts are important for the diagnosis and management of PNDs. First, it is common for symptoms to appear before the presence of a tumor is known; second, the neurologic syndrome usually develops rapidly, producing severe deficits in a short period of time; and third, there is evidence that prompt tumor control improves the neurologic outcome. Therefore, the major concern of the physician is to recognize a disorder promptly as paraneoplastic and to identify and treat the tumor.

PND OF THE CENTRAL NERVOUS SYSTEM AND DORSAL ROOT GANGLIA

When symptoms involve brain, spinal cord, or dorsal root ganglia, the suspicion of PND is usually based on a combination of clinical, radiologic, and CSF findings. Presence of antineuronal antibodies (Tables 122-2 and 122-3) may help in the diagnosis, but only

60–70% of PNDs of the CNS and less than 20% of those involving the peripheral nervous system have neuronal or neuromuscular junction antibodies that can be used as diagnostic tests.

Magnetic resonance imaging (MRI) and CSF studies are important to rule out neurologic complications due to the direct spread of cancer, particularly metastatic and leptomeningeal disease. In most PNDs, the MRI findings are nonspecific. Paraneoplastic limbic encephalitis is usually associated with characteristic MRI abnormalities in the mesial temporal lobes (see below), but similar findings can occur with other disorders (e.g., nonparaneoplastic autoimmune limbic encephalitis and human herpesvirus type 6 [HHV-6] encephalitis) (Fig. 122-2). The CSF profile of patients with PND of the CNS or dorsal root ganglia typically consists of mild to moderate pleocytosis (<200 mononuclear cells, predominantly lymphocytes), an increase in the protein concentration, and a variable presence of oligoclonal bands. There are no specific electrophysiologic tests that are diagnostic of PND. Moreover, a biopsy of the affected tissue is often difficult to obtain, and although useful to rule out other disorders (e.g., metastasis) the pathologic findings are not specific for PND.

PND OF NERVE AND MUSCLE

If symptoms involve peripheral nerve, neuromuscular junction, or muscle, the diagnosis of a specific PND is usually established on clinical, electrophysiologic, and pathologic grounds. The clinical

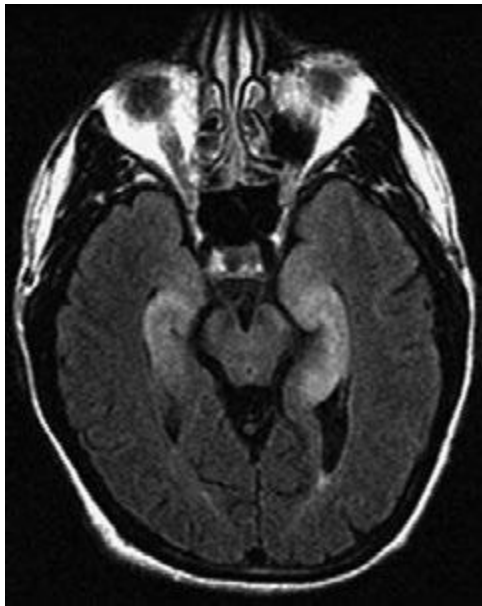


FIGURE 122-2 Fluid-attenuated inversion recovery sequence magnetic resonance imaging of a patient with limbic encephalitis and LGI1 antibodies. Note the abnormal hyperintensity involving the medial aspect of the temporal lobes.

history, accompanying symptoms (e.g., anorexia, weight loss), and type of syndrome dictate the studies and degree of effort needed to demonstrate a neoplasm. For example, the frequent association of Lambert-Eaton myasthenic syndrome (LEMS) with SCLC should lead to a chest and abdomen computed tomography (CT) or body positron emission tomography (PET) scan and, if negative, periodic tumor screening for at least 3 years after the neurologic diagnosis. In contrast, the weak association of polymyositis with cancer calls into question the need for repeated cancer screenings in this situation. Serum and urine immunofixation studies should be considered in patients with peripheral neuropathy of unknown cause; detection of a monoclonal gammopathy suggests the need for additional studies to uncover a B cell or plasma cell malignancy. In paraneoplastic neuropathies, diagnostically useful antineuronal antibodies are limited to anti-CV₂/CRMP5 and anti-Hu.

For any type of PND, if antineuronal antibodies are negative, the diagnosis relies on the demonstration of cancer and the exclusion of other cancer-related or independent neurologic disorders. Combined CT and PET scans often uncover tumors undetected by other tests. For germ cell tumors of the testis and teratomas of the ovary, ultrasound and CT or MRI of the abdomen and pelvis may reveal tumors undetectable by PET.

SPECIFIC PARANEOPLASTIC NEUROLOGIC SYNDROMES

PARANEOPLASTIC ENCEPHALOMYELITIS AND FOCAL ENCEPHALITIS

The term *encephalomyelitis* describes an inflammatory process with multifocal involvement of the nervous system, including brain, brainstem, cerebellum, and spinal cord. It is often associated with dorsal root ganglia and autonomic dysfunction. For any given patient, the clinical manifestations are determined by the areas predominantly involved, but pathologic studies almost always reveal abnormalities beyond the symptomatic regions. Several clinicopathologic syndromes may occur alone or in combination: (1) *cortical encephalitis*, which may present as “epilepsia partialis continua”; (2) *limbic encephalitis*, characterized by confusion, depression, agitation, anxiety, severe short-term memory deficits, partial complex seizures, and sometimes dementia (the MRI usually shows unilateral or bilateral medial temporal lobe abnormalities, best seen with T2 and fluid-attenuated inversion recovery sequences); (3) *brainstem encephalitis*, resulting in eye movement disorders (nystagmus, opsoclonus, supranuclear or nuclear paresis), cranial nerve paresis, dysarthria, dysphagia, and central autonomic dysfunction; (4) *cerebellar gait and limb ataxia*; (5) *myelitis*, which may cause lower or upper motor neuron symptoms, myoclonus, muscle rigidity, and spasms; and (6) *autonomic dysfunction* as a result of involvement of the neuraxis at multiple levels, including hypothalamus, brainstem, and autonomic nerves (see Paraneoplastic Peripheral Neuropathies, below). Cardiac arrhythmias, postural hypotension, and central hypoventilation are frequent causes of death in patients with encephalomyelitis.

Paraneoplastic encephalomyelitis and focal encephalitis are usually associated with SCLC, but many other cancers have been implicated. Patients with SCLC and these syndromes usually have anti-Hu antibodies in serum and CSF. Anti-CRMP5 antibodies occur less frequently; some of these patients may develop chorea, uveitis, or optic neuritis. Antibodies to Ma proteins are associated with limbic, hypothalamic, and brainstem encephalitis and occasionally with cerebellar symptoms (Fig. 122-3); some patients develop hypersomnia, cataplexy, and severe hypokinesia. MRI abnormalities are frequent, including those described with limbic encephalitis and variable involvement of the hypothalamus, basal ganglia, or upper brainstem. The oncologic associations of these antibodies are shown in Table 122-2.

TREATMENT ENCEPHALOMYELITIS AND FOCAL ENCEPHALITIS

Most types of paraneoplastic encephalitis and encephalomyelitis respond poorly to treatment. Stabilization of symptoms or partial neurologic improvement may occasionally occur, particularly if there is a satisfactory response of the tumor to treatment. Controlled trials of therapy are lacking, but many experts recommend treatment initially with glucocorticoids. If there is



FIGURE 122-3 Magnetic resonance imaging (MRI) and tumor of a patient with anti-Ma2-associated encephalitis. (A and B) Fluid-attenuated inversion recovery MRI sequences showing abnormal hyperintensities in the medial temporal lobes, hypothalamus, and upper brainstem. (C) This image corresponds to a section of the patient’s orchietomy incubated with a specific marker (Oct4) of germ cell tumors. The positive (brown) cells correspond to an intratubular germ cell neoplasm.

no response within several days, one can advance to intravenous immunoglobulin (IVIg) or plasma exchange, and then to immunosuppression with either rituximab or cyclophosphamide. Approximately 30% of patients with anti-Ma2-associated encephalitis respond to treatment of the tumor (usually a germ cell neoplasm of the testis) and immunotherapy.

ENCEPHALITIDES WITH ANTIBODIES TO CELL-SURFACE OR SYNAPTIC PROTEINS (TABLE 122-3)

These disorders are important for three reasons: (1) they can occur with and without tumor association, (2) some syndromes predominate in young individuals and children, and (3) despite the severity of the symptoms patients usually respond to treatment of the tumor, if found, and immunotherapy (e.g., glucocorticoids, IVIg, plasma exchange, rituximab, or cyclophosphamide).

Encephalitis with N-methyl-D-aspartate (NMDA) receptor antibodies (Fig. 122-1) usually occurs in young women and children, but men and older patients of both sexes can be affected. The disorder has a characteristic pattern of symptom progression that includes a prodrome resembling a viral process, followed in a few days by the onset of severe psychiatric symptoms, memory loss, seizures, decreased level of consciousness, abnormal movements (orofacial, limb, and trunk dyskinesias, dystonic postures), autonomic instability, and frequent hypoventilation. Monosymptomatic episodes, such as pure psychosis, occur in 4% of the patients. Clinical relapses occur in 12–24% of patients (12% during the first 2 years after initial presentation). Most patients have intrathecal synthesis of antibodies, likely by infiltrating plasma cells in brain and meninges (Fig. 122-4A). The syndrome is often misdiagnosed as a viral or idiopathic encephalitis, neuroleptic malignant syndrome, or encephalitis lethargica, and many patients are initially evaluated by psychiatrists with the suspicion of acute psychosis. The detection of an associated teratoma is dependent on age and gender: 46% of female patients 12 years or older have uni- or bilateral ovarian teratomas, whereas less than 7% of girls younger than 12 have a teratoma (Fig. 122-4B). In male patients, the detection of a tumor is rare. Patients older than 45 years are more frequently male; about 20% of these patients have tumors (e.g., cancer of the breast, ovary, or lung).

Encephalitis with leucine-rich glioma-inactivated 1 (LGII) antibodies predominates in patients older than 50 years (65% male) and frequently presents with memory loss and seizures (limbic encephalopathy), along with hyponatremia and sleep dysfunction. In a small number of patients, the encephalitis is preceded by or occurs with myoclonic-like movements called faciobrachial dystonic or tonic seizures. Less than 10% of patients have thymoma.

Encephalitis with contactin-associated protein-like 2 (Caspr2) antibodies predominates in patients older than 50 years and is associated

with Morvan's syndrome (encephalitis, insomnia, confusion, hallucinations, autonomic dysfunction, and neuromyotonia) and, less frequently, with limbic encephalitis, neuromyotonia, and neuropathic pain. About 30–40% of patients have thymoma.

Encephalitis with γ -aminobutyric acid type B (GABA_B) receptor antibodies is usually associated with limbic encephalitis and seizures. In rare instances, patients develop cerebellar symptoms and opsoclonus. Fifty percent of patients have SCLC or a neuroendocrine tumor of the lung. Patients may have additional antibodies to glutamic acid decarboxylase (GAD), which are of unclear significance. Other antibodies to nonneuronal proteins are often found in these patients as well as in patients with α -amino-3-hydroxy-5-methylisoxazole-4-propionic acid (AMPA) receptor antibodies, indicating a general tendency to autoimmunity.

Encephalitis with GABA_A receptor antibodies may affect children and adults. When antibodies are present at high titer in serum and CSF, the disorder associates with prominent seizures and status epilepticus, often requiring pharmacologically induced coma. Low titer antibodies in serum are often associated with other autoimmune conditions, and the spectrum of symptoms is wider, including encephalitis, seizures, opsoclonus, or stiff-person syndrome. Most patients do not have an underlying tumor, but some may have thymoma.

Encephalitis with AMPA receptor antibodies affects middle-aged women, who develop acute limbic dysfunction or, less frequently, prominent psychiatric symptoms; 70% of the patients have an underlying tumor in the lung, breast, or thymus. Neurologic relapses may occur; these also respond to immunotherapy and are not necessarily associated with tumor recurrence.

Encephalitis with glycine receptor (GlyR) antibodies has been described in adults with progressive encephalomyelitis with rigidity and myoclonus (PERM) and stiff-person spectrum of symptoms (with or without GAD antibodies). The disorder usually occurs without tumor association, although some patients have lung cancer, thymoma, or Hodgkin's lymphoma.

Encephalitis with dipeptidyl-peptidase-like protein-6 (or DPPX) antibodies results in symptoms of CNS hyperexcitability including agitation, hallucinations, paranoid delusions, tremor, myoclonus, nystagmus, seizures, and sometimes hyperekplexia. Some patients develop progressive encephalomyelitis with rigidity and myoclonus. Diarrhea, other gastrointestinal symptoms, and substantial loss of weight often suggest the presence of an underlying tumor, but no tumor association has been identified. The disorder responds to immunotherapy.

PARANEOPLASTIC CEREBELLAR DEGENERATION

This disorder is often preceded by a prodrome that may include dizziness, oscillopsia, blurry or double vision, nausea, and vomiting. A few

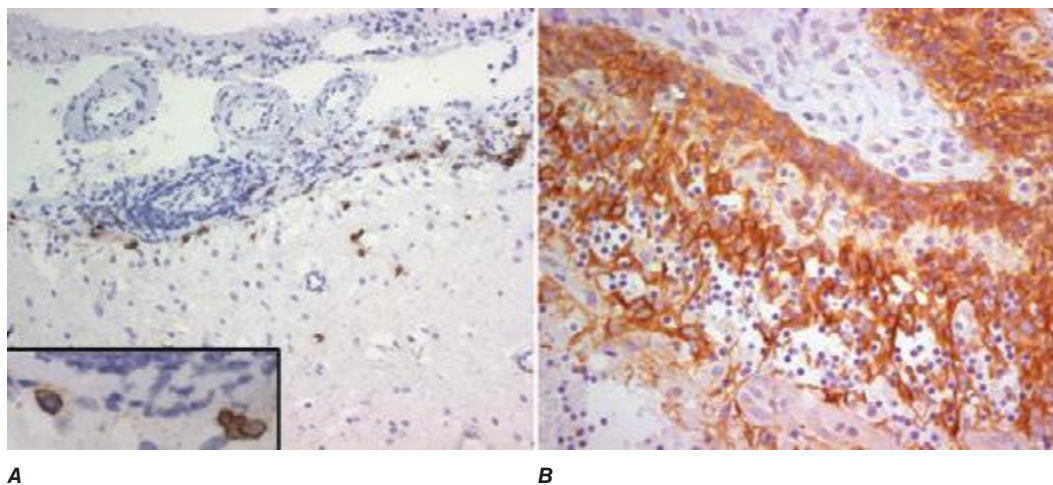


FIGURE 122-4 Pathologic findings in anti-N-methyl-D-aspartate (NMDA) receptor encephalitis. Infiltrates of plasma cells (brown cells; stained for CD138) in the meninges and brain of a patient (A); the inset is a magnification of some plasma cells. (B) Neurons and neuronal processes in the teratoma of a patient (brown cells; stained with MAP2); these neurons express NMDA receptors (not shown). (From E Martinez-Hernandez et al: *Neurology* 77:589, 2011, with permission.)

618 days or weeks later, patients develop dysarthria, gait and limb ataxia, and variable dysphagia. The examination usually shows downbeating nystagmus and, rarely, opsoclonus. Brainstem dysfunction, upgoing toes, or a mild neuropathy may occur. Early in the course, MRI studies are usually normal; later, the MRI reveals cerebellar atrophy. The disorder results from extensive degeneration of Purkinje cells, with variable involvement of other cerebellar cortical neurons, deep cerebellar nuclei, and spinocerebellar tracts. The tumors more frequently involved are SCLC, cancer of the breast and ovary, and Hodgkin's lymphoma.

Anti-Yo antibodies in patients with breast and gynecologic cancers and anti-Tr antibodies in patients with Hodgkin's lymphoma are the two immune responses typically associated with prominent or pure cerebellar degeneration. Antibodies to P/Q-type voltage-gated calcium channels (VGCC) occur in some patients with SCLC and cerebellar dysfunction; only some of these patients develop LEMS. A variable degree of cerebellar dysfunction can be associated with virtually any of the antibodies and PND of the CNS shown in Table 122-2.

A number of single case reports have described neurologic improvement after tumor removal, plasma exchange, IVIg, cyclophosphamide, rituximab, or glucocorticoids. However, most patients with paraneoplastic cerebellar degeneration do not improve with treatment.

PARANEOPLASTIC OPSOCLONUS-MYOCLONUS SYNDROME

Opsoclonus is a disorder of eye movement characterized by involuntary, chaotic saccades that occur in all directions of gaze; it is frequently associated with myoclonus and ataxia. Opsoclonus-myoclonus may be cancer-related or idiopathic. When the cause is paraneoplastic, the tumors involved are usually cancer of the lung and breast in adults, neuroblastoma in children, and ovarian teratoma in adolescents and young women. The pathologic substrate of opsoclonus-myoclonus is unclear, but studies suggest that disinhibition of the fastigial nucleus of the cerebellum is involved. Most patients do not have antineuronal antibodies. A small subset of patients with ataxia, opsoclonus, and other eye-movement disorders develop anti-Ri antibodies; in rare instances, muscle rigidity, laryngeal spasms, autonomic dysfunction, and dementia also occur. The tumors most frequently involved in anti-Ri-associated syndromes are breast and ovarian cancer. If the tumor is not successfully treated, the syndrome in adults often progresses to encephalopathy, coma, and death. In addition to treating the tumor, symptoms may respond to immunotherapy (glucocorticoids, plasma exchange, and/or IVIg).

At least 50% of children with opsoclonus-myoclonus have an underlying neuroblastoma. Hypotonia, ataxia, behavioral changes, and irritability are frequent accompanying symptoms. Neurologic symptoms often improve with treatment of the tumor and glucocorticoids, adrenocorticotropic hormone (ACTH), plasma exchange, IVIg, and rituximab. Many patients are left with psychomotor retardation and behavioral and sleep problems.

PARANEOPLASTIC SYNDROMES OF THE SPINAL CORD

The number of reports of paraneoplastic spinal cord syndromes, such as *subacute motor neuronopathy* and *acute necrotizing myelopathy*, has decreased in recent years. This may represent a true decrease in incidence, due to improved and prompt oncologic interventions, or the identification of nonparaneoplastic etiologies. Some patients with cancer develop *upper or lower motor neuron dysfunction* or both, resembling amyotrophic lateral sclerosis. It is unclear whether these disorders have a paraneoplastic etiology or simply coincide with the presence of cancer. There are isolated case reports of cancer patients with motor neuron dysfunction who had neurologic improvement after tumor treatment. A search for lymphoma should be undertaken in patients with a rapidly progressive motor neuron syndrome and a monoclonal protein in serum or CSF.

Paraneoplastic myelitis may present with upper or lower motor neuron symptoms, segmental myoclonus, and rigidity, and can be the first manifestation of encephalomyelitis. *Neuromyelitis optica (NMO) with aquaporin 4 antibodies* may occur in rare instances as a paraneoplastic manifestation of a cancer.

PARANEOPLASTIC STIFF-PERSON SYNDROME

This disorder is characterized by progressive muscle rigidity, stiffness, and painful spasms triggered by auditory, sensory, or emotional stimuli. Rigidity mainly involves the lower trunk and legs, but it can affect the upper extremities and neck. Sometimes, only one extremity is affected (*stiff-limb syndrome*). Symptoms improve with sleep and general anesthetics. Electrophysiologic studies demonstrate continuous motor unit activity. The associated antibodies target proteins (GAD, amphiphysin) involved in the function of inhibitory synapses using γ -aminobutyric acid (GABA) or glycine as neurotransmitters. The presence of amphiphysin antibodies usually indicates a paraneoplastic etiology related to SCLC and breast cancer. By contrast, GAD antibodies may occur in some cancer patients but are much more frequently present in the nonparaneoplastic disorder. GlyR antibodies may occur in some patients with stiff-person syndrome; these antibodies are also detectable in patients with PERM.

Optimal treatment of stiff-person syndrome requires therapy of the underlying tumor, glucocorticoids, and symptomatic use of drugs that enhance GABA-ergic transmission (diazepam, baclofen, sodium valproate, tiagabine, vigabatrin). IVIg and plasma exchange are transiently effective in some patients.

PARANEOPLASTIC SENSORY NEURONOPATHY OR DORSAL ROOT GANGLIONOPATHY

This syndrome is characterized by sensory deficits that may be symmetric or asymmetric, painful dysesthesias, radicular pain, and decreased or absent reflexes. All modalities of sensation and any part of the body including face and trunk can be involved. Specialized sensations such as taste and hearing can also be affected. Electrophysiologic studies show decreased or absent sensory nerve potentials with normal or near-normal motor conduction velocities. Symptoms result from an inflammatory, likely immune-mediated, process that targets the dorsal root ganglia, causing neuronal loss and secondary degeneration of the posterior columns of the spinal cord. The dorsal and, less frequently, the anterior nerve roots and peripheral nerves may also be involved. This disorder often precedes or is associated with encephalomyelitis and autonomic dysfunction and has the same immunologic and oncologic associations (Hu antibodies, SCLC).

As with anti-Hu-associated encephalomyelitis, the therapeutic approach focuses on prompt treatment of the tumor. Glucocorticoids occasionally produce clinical stabilization or improvement. The benefit of IVIg and plasma exchange is not proven.

PARANEOPLASTIC PERIPHERAL NEUROPATHIES

These disorders may develop any time during the course of the neoplastic disease. Neuropathies occurring at late stages of cancer or lymphoma usually cause mild to moderate sensorimotor deficits due to axonal degeneration of unclear etiology. These neuropathies are often masked by concurrent neurotoxicity from chemotherapy and other cancer therapies. In contrast, the neuropathies that develop in the early stages of cancer frequently show a rapid progression, sometimes with a relapsing and remitting course, and evidence of inflammatory infiltrates and axonal loss or demyelination. If demyelinating features predominate (**Chaps. 459 and 460**), IVIg, plasma exchange, or glucocorticoids may improve symptoms. Occasionally anti-CRMP5 antibodies are present; detection of anti-Hu suggests concurrent dorsal root ganglionitis.

Guillain-Barré syndrome and *brachial plexitis* have occasionally been reported in patients with lymphoma, but there is no clear evidence of a paraneoplastic association (**Chap. 460**).

Malignant monoclonal gammopathies include: (1) multiple myeloma and sclerotic myeloma associated with IgG or IgA monoclonal proteins; and (2) Waldenström's macroglobulinemia, B cell lymphoma, and chronic B cell lymphocytic leukemia associated with IgM monoclonal proteins. These disorders may cause neuropathy by a variety of mechanisms, including compression of roots and plexuses by metastasis to vertebral bodies and pelvis, deposits of amyloid in peripheral nerves, and paraneoplastic mechanisms. The paraneoplastic variety has several distinctive features. Approximately half of patients with sclerotic myeloma develop a sensorimotor neuropathy with predominantly motor deficits, resembling a chronic inflammatory demyelinating

neuropathy (**Chap. 460**); some patients develop elements of the POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, M protein, skin changes). Treatment of the plasmacytoma or sclerotic lesions usually improves the neuropathy. In contrast, the sensorimotor or sensory neuropathy associated with multiple myeloma is more refractory to treatment. Between 5 and 10% of patients with Waldenström's macroglobulinemia develop a distal symmetric sensorimotor neuropathy with predominant involvement of large sensory fibers. These patients may have IgM antibodies in their serum against myelin-associated glycoprotein and various gangliosides (**Chap. 460**). In addition to treating the Waldenström's macroglobulinemia, other therapies may improve the neuropathy, including plasma exchange, IVIg, chlorambucil, cyclophosphamide, fludarabine, or rituximab.

Vasculitis of the nerve and muscle causes a painful symmetric or asymmetric distal axonal sensorimotor neuropathy with variable proximal weakness. It predominantly affects elderly men and is associated with an elevated erythrocyte sedimentation rate and increased CSF protein concentration. SCLC and lymphoma are the primary tumors involved. Glucocorticoids and cyclophosphamide often result in neurologic improvement.

Peripheral nerve hyperexcitability (neuromyotonia, or Isaacs' syndrome) is characterized by spontaneous and continuous muscle fiber activity of peripheral nerve origin. Clinical features include cramps, muscle twitching (fasciculations or myokymia), stiffness, delayed muscle relaxation (pseudomyotonia), and spontaneous or evoked carpal or pedal spasms. The involved muscles may be hypertrophic, and some patients develop paresthesias and hyperhidrosis. CNS dysfunction, including mood changes, sleep disorder, hallucinations, and autonomic symptoms may occur. The electromyogram (EMG) shows fibrillations; fasciculations; and doublet, triplet, or multiplet single-unit (myokymic) discharges that have a high intraburst frequency. Some patients have Caspr2 antibodies in the context of Morvan's syndrome, but most cases of isolated neuromyotonia are antibody negative. The disorder often occurs without cancer; if paraneoplastic, benign and malignant thymomas and SCLC are the usual tumors. Phenytoin, carbamazepine, and plasma exchange improve symptoms.

Paraneoplastic autonomic neuropathy usually develops as a component of other disorders, such as LEMS and encephalomyelitis. It may rarely occur as a pure or predominantly autonomic neuropathy with cholinergic or adrenergic dysfunction at the pre- or postganglionic levels. Patients can develop several life-threatening complications, such as gastrointestinal paresis with pseudo-obstruction, cardiac dysrhythmias, and postural hypotension. Other clinical features include abnormal pupillary responses, dry mouth, anhidrosis, erectile dysfunction, and problems in sphincter control. The disorder occurs in association with several tumors, including SCLC, cancer of the pancreas or testis, carcinoid tumors, and lymphoma. Because autonomic symptoms can be the presenting feature of encephalomyelitis, serum anti-Hu and anti-CRMP5 antibodies should be sought. Antibodies to ganglionic (alpha3-type) neuronal acetylcholine receptors are the cause of autoimmune autonomic ganglionopathy, a disorder that frequently occurs without cancer association (**Chap. 454**).

LAMBERT-EATON MYASTHENIC SYNDROME

LEMS is discussed in **Chap. 461**.

MYASTHENIA GRAVIS

Myasthenia gravis is discussed in **Chap. 461**.

POLYMYOSITIS-DERMATOMYOSITIS

Polymyositis and dermatomyositis are discussed in detail in **Chap. 388**.

ACUTE NECROTIZING MYOPATHY

Patients with this syndrome develop myalgias and rapid progression of weakness involving the extremities and the pharyngeal and respiratory muscles, often resulting in death. Serum muscle enzymes are elevated, and muscle biopsy shows extensive necrosis with minimal or absent inflammation and sometimes deposits of complement. The disorder occurs as a paraneoplastic manifestation of a variety of cancers including SCLC and cancer of the gastrointestinal tract, breast, kidney, and

prostate, among others. Glucocorticoids and treatment of the underlying tumor rarely control the disorder.

PARANEOPLASTIC VISUAL SYNDROMES

This group of disorders involves the retina and, less frequently, the uvea and optic nerves. The term *cancer-associated retinopathy* is used to describe paraneoplastic cone and rod dysfunction characterized by photosensitivity, progressive loss of vision and color perception, central or ring scotomas, night blindness, and attenuation of photopic and scotopic responses in the electroretinogram (ERG). The most commonly associated tumor is SCLC. Melanoma-associated retinopathy affects patients with metastatic cutaneous melanoma. Patients develop acute onset of night blindness and shimmering, flickering, or pulsating photopsias that often progress to visual loss. The ERG shows reduced b waves with normal dark adapted a waves. Paraneoplastic optic neuritis and uveitis are very uncommon and can develop in association with encephalomyelitis. Some patients with paraneoplastic uveitis and optic neuritis have anti-CRMP5 antibodies.

Some paraneoplastic retinopathies are associated with serum antibodies that specifically react with the subset of retinal cells undergoing degeneration, supporting an immune-mediated pathogenesis (**Table 122-2**). Paraneoplastic retinopathies usually fail to improve with treatment, although rare responses to glucocorticoids, plasma exchange, and IVIg have been reported.

The thymus is derived from the third and fourth pharyngeal pouches and is located in the anterior mediastinum. It is composed of epithelial and stromal cells derived from the pharyngeal pouch and lymphoid precursors derived from mesodermal cells. It is the site to which bone marrow precursors that are committed to differentiate into T cells migrate to complete their differentiation. Like many organs, it is organized into functional regions, in this case the cortex and the medulla. The cortex of the thymus contains ~85% of the lymphoid cells, and the medulla contains ~15%. It appears that the primitive bone marrow progenitors enter the thymus at the corticomedullary junction and migrate first through the cortex toward the periphery of the gland and then toward the medulla as they mature. Medullary thymocytes have a phenotype that cannot be distinguished readily from that of mature peripheral blood and lymph node T cells.

Several things can go wrong with the thymus, but thymic abnormalities are very rare. If the thymus does not develop properly, serious deficiencies in T-cell development ensue and severe immunodeficiency is seen (e.g., DiGeorge syndrome, [Chap. 374](#)). If a lymphoid cell within the thymus becomes neoplastic, the disease that develops is a lymphoma. The majority of lymphoid tumors that develop in the thymus are derived from the precursor T cells, and the tumor is a precursor T-cell lymphoblastic lymphoma ([Chap. 134](#)). Rare B cells exist in the thymus, and when they become neoplastic, the tumor is a mediastinal (thymic) B cell lymphoma ([Chap. 134](#)). Hodgkin's disease, particularly the nodular sclerosing subtype, often involves the anterior mediastinum. Extranodal marginal zone (mucosa-associated lymphoid tissue [MALT]) lymphomas have been reported to involve the thymus in the setting of Sjögren's syndrome or other autoimmune disorders, and the lymphoma cells often express IgA instead of IgM on their surface. Castleman's disease can involve the thymus. Germ cell tumors and carcinoid tumors occasionally may arise in the thymus. If the epithelial cells of the thymus become neoplastic, the tumor that develops is a *thymoma*.

CLINICAL PRESENTATION AND DIFFERENTIAL DIAGNOSIS

Thymoma, although rare (0.1–0.15 cases per 100,000 person-years), is the most common cause of an anterior mediastinal mass in adults, accounting for ~40% of all mediastinal masses. The other major causes of anterior mediastinal masses are lymphomas, germ cell tumors, and substernal thyroid tumors. Carcinoid tumors, lipomas, and thymic cysts also may produce radiographic masses. After combination chemotherapy for another malignancy, teenagers and young adults may develop a rebound thymic hyperplasia in the first few months after treatment. Granulomatous inflammatory diseases (tuberculosis, sarcoidosis) can produce thymic enlargement. Thymomas are most common in the fifth and sixth decades, are uncommon in children, and are distributed evenly between men and women.

About 40–50% of patients are asymptomatic; masses are detected incidentally on routine chest radiographs. When symptomatic, patients may have cough, chest pain, dyspnea, fever, wheezing, fatigue, weight loss, night sweats, or anorexia. Occasionally, thymomas may obstruct the superior vena cava. Pericardial effusion may be present. About 40% of patients with thymoma have another systemic autoimmune illness related to the thymoma. About 30% of patients with thymoma have myasthenia gravis, 5–8% have pure red cell aplasia, and ~5% have hypogammaglobulinemia. Thymoma with hypogammaglobulinemia also is called Good's syndrome. Among patients with myasthenia gravis, ~10–15% have a thymoma. Thymoma more rarely may be associated with polymyositis, systemic lupus erythematosus, thyroiditis, Sjögren's syndrome, ulcerative colitis, pernicious anemia, Addison's disease, stiff person syndrome, scleroderma, and panhypopituitarism. In one series, 70% of patients with thymoma were found to have another systemic illness.

DIAGNOSIS AND STAGING

Once a mediastinal mass is detected, a surgical procedure is required for definitive diagnosis. An initial mediastinoscopy or limited thoracotomy can be undertaken to get sufficient tissue to make an accurate diagnosis. Fine-needle aspiration is poor at distinguishing between lymphomas and thymomas but is more reliable in diagnosing germ cell tumors and metastatic carcinoma. Thymomas and lymphomas require sufficient tissue to examine the tumor architecture to assure an accurate diagnosis and obtain prognostic information.

Once a diagnosis of thymoma is defined, subsequent staging generally occurs at surgery. However, chest computed tomography (CT) scans can assess local invasiveness in some instances. Magnetic resonance imaging (MRI) has a defined role in the staging of posterior mediastinal tumors, but it is not clear that it adds important information to the CT scan in anterior mediastinal tumors. Somatostatin receptor imaging with indium-labeled somatostatin analogues may be of value. If invasion is not distinguished by noninvasive testing, an effort to resect the entire tumor should be undertaken. If invasion is present, neoadjuvant chemotherapy may be warranted before surgery (see "Treatment" section below).

Some 90% of thymomas are in the anterior mediastinum, but some may be in other mediastinal sites or even the neck, based on aberrant migration of the developing thymic enlage.

The staging system for thymoma was developed by Masaoka and colleagues ([Table 123e-1](#)). It is an anatomic system in which the stage is increased on the basis of the degree of invasiveness. The 5-year survival of patients in the various stages is as follows: stage I, 96%; stage II, 86%; stage III, 69%; and stage IV, 50%. The French Study Group on Thymic Tumors (GETT) has proposed modifications to the Masaoka scheme based on the degree of surgical removal because the extent of surgery has been noted to be a prognostic indicator. In their system, stage I tumors are divided into A and B on the basis of whether the surgeon suspects adhesions to adjacent structures; stage III tumors are divided into A and B based on whether disease was subtotally resected or only biopsied. The concurrence between the two systems is high.

PATHOLOGY AND ETIOLOGY

Thymomas are epithelial tumors, and all of them have malignant potential. It is not worthwhile to try to divide them into benign and malignant forms; the key prognostic feature is whether they are noninvasive or invasive. About 65% of thymomas are encapsulated

TABLE 123e-1 MASAOKA STAGING SYSTEM FOR THYOMAS

Stage	Diagnostic Criteria	Stage Distribution, %	5-Year Survival, %	10-Year Survival, %
I	Macroscopically and microscopically completely encapsulated; no invasion through capsule	36	95–100	86–100
II	IIA Microscopic invasion outside the capsule IIB Macroscopic invasion into surrounding fat or grossly adherent to pleura or pericardium	26	70–100	50–100
III	IIIA Macroscopic invasion into neighboring organs, pericardium, or pleura but not great vessels IIIB Macroscopic invasion into neighboring organs that includes great vessels	22	68–89	47–60
IV	IVA Pleural or pericardial dissemination IVB Lymphatic or hematogenous metastases	10	47–69	0–11

Source: From A Masaoka et al: *Cancer* 48:2485, 1981. Updated from S Tomaszek et al: *Ann Thorac Surg* 87:1973, 2009, and CB Falkson et al: *J Thorac Oncol* 4:911, 2009.

TABLE 123e-2 WORLD HEALTH ORGANIZATION (WHO) HISTOLOGIC CLASSIFICATION OF THYMUS TUMORS

Type	Histologic Description	
A	Medullary thymoma	
AB	Mixed thymoma	
B1	Predominantly cortical thymoma	
B2	Cortical thymoma	
B3	Well-differentiated thymic carcinoma	
C	Thymic carcinoma	
Type	Distribution, %	Prognosis (10-year disease-free survival), %
A	8	100
AB	26	90–100
B1	15	78–94
B2	28	83
B3	15	36
C	8	0–35

Source: From S Tomaszek et al: Ann Thorac Surg 87:1973, 2009.

and noninvasive, and about 35% are invasive. They may have a variable percentage of lymphocytes within the tumor, but genetic studies suggest that the lymphocytes are benign polyclonal cells. The epithelial component of the tumor may consist primarily of round or oval cells derived mainly from the cortex or spindle-shaped cells derived mainly from the medulla or combinations of the two types (World Health Organization classification; Table 123e-2). Cytologic features are not reliable predictors of biologic behavior. In part, this unreliability may be related to the moderate reproducibility of the system. About 90% of A, AB, and B1 tumors are localized. A very small number of patients have aggressive histology features characteristic of carcinomas. Thymic carcinomas are invasive and have a poor prognosis.

Genetic lesions are common in thymomas. The most common abnormalities affect chromosome 6p21.3 (the MHC locus) and 6p25.2–25.3 (usually loss of heterozygosity). Abnormalities affecting a number of other genes altered in other types of tumors are also seen, including *p53*, *RB*, *FHIT*, and *APC*. Thymic carcinomas may overexpress *c-kit*, *HER2*, or growth factor receptor genes (epidermal growth factor receptor and insulin-like growth factor receptor). Some data suggest that Epstein-Barr virus may be associated with thymomas. Some tumors overexpress the p21 *ras* gene product. However, molecular pathogenesis remains undefined. A thymoma susceptibility locus has been defined on *rat* chromosome 7, but the relationship between this gene locus, termed *Tsr1*, and human thymoma has not been examined.

INFLUENCE OF THYMECTOMY ON THE COURSE OF ACCOMPANYING DISEASES

Patients with myasthenia gravis have a high incidence of thymic abnormalities (~80%), but overt thymoma is present in only ~10–15% of patients with myasthenia gravis. It is thought that the thymus plays

a role in breaking self-tolerance and generating T cells that recognize the acetylcholine receptor as a foreign antigen. Although patients with thymoma and myasthenia gravis are less likely to have a remission in the myasthenia as a consequence of thymectomy than are patients with thymic abnormalities other than thymoma, the course of myasthenia gravis is not significantly different in patients with or without thymoma. Thymectomy produces at least some symptomatic improvement in ~65% of patients with myasthenia gravis. In one large series, thymoma patients with myasthenia gravis had a better long-term survival from thymoma resection than did those without myasthenia gravis.

About 30–50% of patients with pure red cell aplasia have a thymoma. Thymectomy results in the resolution of pure red cell aplasia in ~30% of patients. About 10% of patients with hypogammaglobulinemia have a thymoma, but hypogammaglobulinemia rarely responds to thymectomy.

TREATMENT THYMOMA

Treatment is determined by the stage of disease. For patients with encapsulated tumors and stage I disease, complete resection is sufficient to cure 96% of patients. For patients with stage II disease, complete resection may be followed by 30–60 Gy of postoperative radiation therapy to the site of the primary tumor. However, the value of radiation therapy in this setting has not been established. The main predictors of long-term survival are Masaoka stage and completeness of resection. For patients with stage III and IV disease, the use of neoadjuvant chemotherapy followed by radical surgery, with or without additional radiation therapy, and additional consolidation chemotherapy has been associated with excellent survival. Chemotherapy regimens that are most effective generally include a platinum compound (either cisplatin or carboplatin) and an anthracycline. Addition of cyclophosphamide, vincristine, and prednisone seems to improve response rates. Response rates of 50–93% have been reported in series of patients each of which involved fewer than 40 patients. A single most effective regimen has not been defined. No randomized controlled phase III studies have been reported. If surgery after neoadjuvant chemotherapy fails to produce a complete resection of residual disease, radiation therapy (50–60 Gy) may help reduce recurrence rates.

This multimodality approach appears to be superior to the use of surgery followed by radiation therapy alone, which produces a 5-year survival of ≤50% in patients with advanced-stage disease.

Some thymic carcinomas express *c-kit*, and one patient whose *c-kit* locus was mutated responded dramatically to imatinib. Many thymomas express epidermal growth factor receptors, but the antibodies to the receptor and the kinase inhibitors that block its action have not been evaluated systematically. Octreotide plus prednisone produces responses in about one-third of patients.

Cancer complicates ~1 in every 1000 pregnancies. Of all the cancers that occur in women, less than 1% complicate pregnancies. The four cancers that most commonly complicate pregnancies are cervical cancer, breast cancer, melanoma, and lymphomas (particularly Hodgkin's lymphoma); however, virtually every form of cancer has been reported in pregnant women (Table 124e-1). In addition to cancers developing in other organs of the mother, gestational trophoblastic tumors can arise from the placenta. The problem of cancer in a pregnant woman is complex. One must take into account (1) the possible influence of the pregnancy on the natural history of the cancer, (2) effects on the mother and fetus of complications from the malignancy (e.g., anorexia, nausea, vomiting, malnutrition), (3) potential effects of diagnostic and staging procedures, and (4) potential effects of cancer treatments on both the mother and the developing fetus. Generally, the management that optimizes maternal physiology is also best for the fetus. However, the dilemma occasionally arises that what is best for the mother may be harmful to the fetus, and what is best for the fetus may compromise the ultimate prognosis for the mother. The best way to approach management of a pregnant woman with cancer is to ask, "What would we do for this woman in this clinical situation if she was not pregnant? Now, which, if any, of those plans need to be modified because she is pregnant?"

Pregnancy is associated with a number of physiologic changes that frequently result in symptoms that may make it difficult to recognize symptoms or physical findings suggestive of a neoplasm. Increased sensitivity of central chemoreceptors to PCO_2 drives an increase in minute ventilation that many women perceive as dyspnea at rest or with minimal exertion. The combination of increased total body water, decreased colloid oncotic pressure, and some obstruction of venous return from the lower extremities causes demonstrable dependent edema in more than 50% of pregnant women. Decreased gastrointestinal motility due to high serum progesterone levels and mechanical compression from an enlarging uterus cause early satiety, gastroesophageal reflux, nausea, vomiting, and constipation. Hemorrhoids develop and often bleed. Breasts enlarge and increase in density and "lumpiness." These changes may result in delayed recognition and more advanced disease at diagnosis.

Physiologic changes in the maternal immune system necessary to facilitate retention of the fetal semi-allograft raise concerns that the relationship of a cancer with its host may be altered to the detriment of the maternal host. One half of all the genes necessary to create a new individual by sexual reproduction come from each parent. This provides the opportunity for many antigenic differences between the conceptus and the mother. Mammalian placentation has been a very successful method of reproduction, but it has necessitated some combination of both fetal and maternal evolutionary immune adaptations. These mechanisms are incompletely understood and remain an area of active investigation. It does seem likely, however, that this has been accomplished without a general, nonspecific blunting of the maternal

immune response, which would be maladaptive to the mother. The multiple mechanisms likely include some "masking" of fetal antigens from recognition by the maternal immune system, blunting the maternal inflammatory response locally at the placental-maternal interface and induction of fetal-specific maternal immune tolerance to avoid rejection. Attention has turned to a subset of CD4+ induced, peripherally produced regulatory T cells that express the X chromosome encoded transcription factor *Foxp3* (so-called *Tregs*). When these *Foxp3* cells develop centrally in the thymus, they are termed "Tregs." When *Foxp3*-expressing cells develop peripherally, they are called "Pregs." These regulatory cells suppress the immune response against "self" and foreign antigens. They seem to be capable of suppressing the maternal response to paternal antigens expressed by the fetus and creating memory cells that retain tolerance to the same paternal antigens in subsequent pregnancies. Unfortunately, in a mouse model, the interleukin (IL) 10 produced by these cells enhanced susceptibility to infection by *Listeria* and *Salmonella*, while ironically not proving essential for retaining the fetal graft. Undoubtedly much remains to be learned about this critical immune balance.

RADIATION IN PREGNANCY

Exposure of developing fetuses to ionizing radiation may cause adverse fetal effects; awareness among physicians of this potential toxicity has resulted in a disproportionate aversion to diagnostic imaging in pregnancy. First, it must be stated that there are very useful imaging modalities (i.e., ultrasound and magnetic resonance imaging [MRI]) that do not use any ionizing radiation and are not associated with any demonstrable adverse fetal effects. There are three potential adverse fetal effects of ionizing radiation: teratogenesis (induction of anatomic birth defects), mutagenesis, and carcinogenesis. The fetus is most sensitive to teratogenesis during organogenesis in the first trimester. The dose of ionizing radiation necessary to induce birth defects in human fetuses is derived from studies of the survivors of the atomic bomb explosions and by extrapolation from controlled experiments in nonhuman mammals. From these data sources, it is clear that a minimum of 5 rem and more likely greater than 10 rem exposure is needed to induce birth defects in the first trimester. The fetal doses of radiation associated with some common diagnostic radiologic procedures are displayed in Table 124e-2. The data in Table 124e-2 show that no single procedure or selective combination of diagnostic procedures will exceed the very conservative 5 rem teratogenic threshold. Teratogenic effects later in pregnancy are largely limited to microcephaly and require exposures exceeding 25 rem. The reason for the disproportionate concern about radiation exposure and birth defects is that 2.5% of all fetuses are affected with birth defects without radiation exposure and, therefore, 2.5% of women undergoing any diagnostic imaging procedure will deliver malformed fetuses. Spontaneous mutations occur relatively infrequently, and high doses of radiation (>150 rem) are required to cause a demonstrable increase in that rate. The magnitude of the risk of carcinogenesis in offspring exposed as fetuses to diagnostic doses of radiation has been very difficult to

TABLE 124e-1 INCIDENCE OF MALIGNANT TUMORS DURING GESTATION

Tumor Type	Incidence per 10,000 Pregnancies ^a	% of Cases ^b
Breast cancer	1–3	25%
Cervical cancer	1.2–4.5	25%
Thyroid cancer	1.2	15%
Hodgkin's disease	1.6	10%
Melanoma	1–2.6	8%
Ovarian cancer	0.8	2%
All sites	10	100%

^aThese are estimates based on extrapolations from a review of more than 3 million pregnancies (LH Smith et al: *Am J Obstet Gynecol* 184:1504, 2001). ^bBased on accumulating case reports from the literature; the precision of these data is not high.

TABLE 124e-2 ESTIMATED FETAL EXPOSURE FROM SOME COMMON RADIOLOGIC PROCEDURES

Procedure	Fetal Exposure
Chest x-ray (2 views)	0.02–0.07 mrad
Abdominal film (single view)	100 mrad
Intravenous pyelography	≥1 rad ^a
Hip film (single view)	7–20 mrad
Barium enema or small bowel series	2–4 rad
CT scan of head or chest	<1 rad
CT scan of abdomen and lumbar spine	3.5 rad
CT pelvimetry	250 mrad

^aExposure depends on the number of films.

Abbreviation: CT, computed tomography.

Source: Data from FG Cunningham et al: General considerations and maternal evaluation. In *Williams Obstetrics*, 21st ed. New York: McGraw-Hill; 2001, pp. 1143–1158.

measure due to the relative rarity of cancer in children and the long duration of follow-up that might credibly be needed to see the effect. The inconsistent results and small effect sizes observed from diagnostic exposures make it likely that, if there is an effect, it is very small and, if there is not a significant effect, it will be impossible to prove that fact to everyone's satisfaction. No imaging using ionizing radiation should be done without a compelling reason and due consideration to obtaining the necessary information by other imaging modalities. Exposure to diagnostic and therapeutic radionuclides, especially radioactive iodine, poses unique risks, but a full discussion of these is beyond the scope of this chapter. Radiation therapy uses radiation doses three orders of magnitude greater than diagnostic procedures, entails substantial risks if the fetus is in the radiation field, and is rarely appropriate in pregnancy. Finally, although difficult to prove, it is likely that more harm has come to pregnant women from failing to perform appropriate diagnostic procedures than has been done to their offspring from performing appropriate diagnostic procedures.

CHEMOTHERAPY IN PREGNANCY

There are a number of reasons why it is impossible to make many definitive statements regarding the safety and efficacy of chemotherapy in pregnancy. All of the available data in the literature are published as case reports or case series. The quality and completeness of the data are inconsistent and often poor. Reports may come from medical oncologists, obstetricians, pediatricians, or other treating physicians familiar with the information important to the report from their own perspective but missing information important for other specialty areas. Reports frequently lack critical details of drug administration, such as dose, duration, cumulative dose, and timing of exposure in gestation, and outcomes, including birth weight and gestational age at delivery, indication for or cause of premature delivery, and follow-up of offspring beyond the immediate neonatal period. There are a wide variety of agents available to treat cancer, and they are usually used in combinations. This results in the fact that every patient is almost unique (an experiment of one) in the combination of agents, doses, durations, and gestational ages of administration, making it very difficult to attribute what benefit or toxicity accrues to which agent. Fortunately, cancer in pregnant women is sufficiently rare that it takes quite a while to accumulate enough information for any one agent or combination of agents to be confident about what toxicities (including congenital malformations) are truly associated with which agents. There is such rapid progress in cancer chemotherapy that by the time there may seem to be enough information about the agents currently in use to use them intelligently and counsel patients meaningfully, the cancer community has moved on to newer, more efficacious, and hopefully less toxic agents for which there is little or no experience in pregnancy. Finally, for obvious reasons, there are no untreated controls for comparison. It may be very difficult to sort out the maternal consequences (nausea, vomiting, fever, weight loss, dehydration) that might result directly from the malignancy and cause adverse pregnancy outcomes from some of the toxicities of the chemotherapeutic agents used to treat the malignancy.

Generally, toxic chemotherapy should be avoided during pregnancy, if at all possible. It should virtually never be given in the first trimester. However, a variety of single agents and combinations have been given in the second and third trimesters, without a high frequency of toxic effects to the pregnancy or the fetus, but data on safety are sparse. Maternal factors that may influence the pharmacology of chemotherapeutic agents include the 50% increase in plasma volume, altered absorption and protein binding, increased glomerular filtration rate, increased hepatic mixed function oxidase activity, and third space created by amniotic fluid. The fetus is protected from some agents by placental expression of drug efflux pumps, but decreased fetal hepatic mixed function oxidase and glucuronidation activity may prolong the half-life of agents that do cross the placenta. A database on the risks associated with individual chemotherapy agents is available on the Internet (http://ntp.niehs.nih.gov/ntp/ohat/cancer_chemo_preg/chemopregnancy_monofinal_508.pdf).

Optimal management strategies have not been developed based on prospective clinical trials. Management of a malignancy complicating pregnancy will be critically determined by the gestational age when the malignancy is diagnosed and the anticipated natural history of the lesion, if left untreated. On one extreme, if the malignancy is slowly progressive, the patient is near her delivery date, and waiting until delivery to begin treatment would not be anticipated to compromise maternal prognosis, then treatment could be delayed until after delivery to avoid fetal exposure to chemotherapy. If there is a greater sense of urgency to begin definitive treatment to avoid compromising maternal prognosis, and the patient is beyond 24 weeks of gestation but remote from her delivery date, then treatment (surgical, medical, or both) might be initiated during pregnancy and plans made to deliver the fetus early to avoid exposure to more chemotherapy than absolutely necessary. Finally, if the patient is in her first trimester and toxic chemotherapy must be initiated promptly to avoid a very poor maternal outcome, then it may be necessary to consider therapeutic abortion to avoid maternal disaster and fetal survival with injury resulting in long-term morbid sequelae. No two cases are precisely alike, and inevitably, decision making must be individualized, preferably with consultation with a multidisciplinary team including medical oncology, surgical oncology if appropriate, maternal-fetal medicine, neonatology, and anesthesia. Pregnancy appears to have little or no impact on the natural history of malignancies, despite the hormonal influences. Spread of the mother's cancer to the fetus (so-called *vertical transmission*) is exceedingly rare.

CERVICAL CANCER DURING PREGNANCY

The incidence of cervical cancer in pregnant women is roughly comparable to that of age-matched controls who are not pregnant. Invasive cervical cancer complicates about 0.45 in 1000 live births, and carcinoma in situ is seen in 1 in 750 pregnancies. About 1% of women diagnosed with cervical cancer are pregnant at the time of diagnosis. Early signs of cervical cancer include vaginal spotting or discharge, pain, and postcoital bleeding, which are also common features of pregnancy. Early visual changes in the cervix related to invasive cancer can be mistaken for cervical decidualization or ectropion (columnar epithelium on the cervix) due to pregnancy. Women diagnosed with cervical cancer during pregnancy report having had symptoms for 4.5 months on average.

Approximately 95% of all cervical cancer is caused by human papillomavirus (HPV) infections, with types 16 and 18 accounting for about 70% of cervical cancer. The rate of carriage of these serotypes is highest among women in their early twenties and can be reduced with the use of vaccination before exposure. Women generally tend to clear the infection by age 30, with the risk of cervical cancer being highest among those who fail to clear the infection. Screening is recommended at the first prenatal visit and 6 weeks postpartum. The rate of cytologic abnormalities on Pap smear in pregnant women is about 5–8% and is not much different than the rate in nonpregnant women of the same age.

In 2012, several sets of recommendations were published for screening for cervical cancer: one by the American Cancer Society (ACS), the American Society for Colposcopy and Cervical Pathology (ASCCP), and the American Society for Clinical Pathology (ASCP); a second by the U.S. Preventive Services Task Force (USPSTF); and a third by the American College of Obstetricians and Gynecologists (ACOG). Although the details of the recommendations for screening and management of abnormal results differ slightly among the three sets of guidelines, there is general consensus that cytology screening should start at age 21 and continue every 3 years through age 29. After age 30, cytology screening frequency may be reduced to every 5 years if accompanied by co-testing for HPV. Recommendations for management of abnormal cytology findings are complex and determined by the degree of abnormality of the cytology finding (e.g., atypical squamous cells of undetermined significance; atypical squamous cells, cannot exclude high-grade squamous intraepithelial lesion; low-grade squamous intraepithelial lesion; or high-grade squamous intraepithelial

lesion), the HPV status of the patient, the age of the patient, and whether this is the first abnormal finding or a persistent abnormality. A full discussion of all the treatment recommendations based on these factors is beyond the scope of this chapter. Some of the diagnostic procedures recommended for evaluation of nonpregnant women are contraindicated in pregnancy, and the indications for some procedures are modified in the setting of pregnancy. Suffice it to say that the evaluation of women with abnormal cervical cytology in pregnancy should be referred to knowledgeable and experienced gynecologists or gynecologic oncologists.

Cervical intraepithelial neoplasia is a slowly progressive lesion and has a low risk of progression to invasive cancer during pregnancy (~0.4%), and many low-grade lesions (36–70%) regress spontaneously. Accordingly, some physicians defer definitive diagnostic procedures in pregnant women until 6 weeks postpartum unless they are at high risk for invasive disease. If invasive disease is suspected and the pregnancy is between 16 and 20 weeks, a cone biopsy may be performed to make the diagnosis and may be curative for some lesions; however, the procedure may cause heavy bleeding due to the increased vasculature in the gravid cervix and increases the risk of premature rupture of membranes and preterm labor two- to threefold. Cone biopsy should not be done within 4 weeks of delivery. The only indication for therapy of cervical neoplasia in pregnant women is the documentation of invasive cancer.

Management of invasive disease is guided by the stage of disease, the gestational age of the fetus, and the desire of the mother to have the baby. If the disease is in early stage and the pregnancy is desired, it is safe to delay treatment regardless of gestational age until fetal maturity allows for safe delivery. Abortion followed by definitive therapy is recommended for women with advanced, but potentially curable, cancer in the first or second trimester (Chap. 117). If the disease is in an advanced stage in early pregnancy and the patient declines pregnancy termination to permit prompt definitive therapy, she must be informed of the fact that the maternal safety of delaying therapy is unproven. In women in the third trimester with advanced disease, the mother should be treated with betamethasone to accelerate fetal lung maturation and the baby should be delivered at the earliest possible gestational age followed immediately by stage-appropriate therapy. Most women with invasive cancer have early-stage disease. If the disease is microinvasive, vaginal delivery can take place and be followed by definitive treatment, usually conization. If a lesion is visible on the cervix, delivery is best done by caesarian section and followed by radical hysterectomy.

BREAST CANCER DURING PREGNANCY

Breast cancer complicates approximately 1 in 3000 to 10,000 live births. About 5% of all breast cancers occur in women age 40 years or younger. Among all premenopausal women with breast cancer, 25–30% were pregnant at the time of diagnosis. It has been recognized for some time that breast cancer associated with pregnancy generally seems to have a poorer prognosis for both overall survival and progression-free survival. The definition of *pregnancy-associated breast cancer* (PABC) has differed in various publications, but a generally accepted definition is breast cancer diagnosed during pregnancy or within 1 year of delivery. There are likely several reasons for the observation of the poorer prognosis. Breast cancers diagnosed during pregnancy are often diagnosed at a later stage of disease and so have a poorer outcome. The late diagnosis is often due to the fact that early physical signs of the disease are missed or attributed to the changes that occur in the breast normally as a function of pregnancy. However, a discreet breast mass in a pregnant woman should never be assumed to be normal. Another reason is the more aggressive behavior of the cancer possibly related to the hormonal milieu (estrogen increases 100-fold; progesterone increases 1000-fold) of the pregnancy. However, about 70% of the breast cancers found in pregnancy are estrogen receptor-negative. About 28–58% of the tumors express HER2, a biologically more aggressive breast cancer subset. Another factor is that aggressive, definitive chemotherapy and radiation therapy are often delayed due

TABLE 124e-3 DIFFERENCES IN BREAST CANCERS IN PREGNANT AND NONPREGNANT WOMEN

	Pregnant	Nonpregnant
Tumor size	3.5 cm	2 cm
Estrogen receptor +	30% ^a	67%
HER2 +	Up to 58%	10–25%
Stage II, III	65–90%	45–66%
Lymph node +	56–89%	38–54%

^aLower measured levels could be in part artifactual due to the increased levels of estrogen in the milieu.

to concerns about the consequences of those treatments for the fetus. Younger women with breast cancer have a higher likelihood of having mutations in *BRCA1* or *BRCA2*.

Differences in presentation between PABC and breast cancers diagnosed in nonpregnant women are shown in Table 124e-3. About 20% of breast cancers are detected in the first trimester, 45% in the second trimester, and 35% in the third trimester. Some argue that stage for stage, the outcome is the same for breast cancer diagnosed in pregnant and nonpregnant women. Primary tumors in pregnant women are 3.5 cm on average, compared to <2 cm in nonpregnant women. A dominant mass and a nipple discharge are the most common presenting signs, and they should prompt ultrasonography and breast MRI exam (if available) followed by lumpectomy if the mass is solid and aspiration if the mass is cystic. Mammography is less reliable in pregnancy due to the increased breast density. Needle aspirates of breast masses in pregnant women are often nondiagnostic or falsely positive. Even in pregnancy, most breast masses are benign (~80% are adenoma, lobular hyperplasia, milk retention cyst, fibrocystic disease, fibroadenoma, or other rarer entities).

Many studies comparing outcomes among women with PABC to those of nonpregnant women have small sample sizes, and there is considerable heterogeneity among the study results, but a formal meta-analysis including multiple adjustments and sensitivity analyses confirms the clinical impression of poorer outcomes for women with PABC. The hazard ratios were 1.44 for poorer overall survival and 1.60 for poorer disease-free survival.

Although having had a pregnancy is a protective factor against breast cancer in women in general, it is questionable as to whether it retains its protective effect in carriers of *BRCA1* and *BRCA2* mutations. Cullinane et al. (Int J Cancer 117:988, 2005) found a statistically insignificant difference (odds ratio 0.94) in breast cancer risk among *BRCA1* carriers who had ever been pregnant versus those who never had a pregnancy. Stratifying the risk of breast cancer according to the number of prior pregnancies versus no pregnancies, no statistically significant protective trend was observed. For *BRCA2* carriers, there was a marginally statistically significant increased risk of breast cancer among women with prior pregnancies. In an international study with more than 65,000 person-years of observation (Andrieu J: Natl Cancer Inst 98:535, 2006), there was no significant effect in either direction of pregnancy on breast cancer risk for carriers of either mutation. Staging the axillary lymph nodes is currently somewhat controversial. Sentinel lymph node sampling is not straightforward in pregnant women. Blue dye has been carcinogenic in rats, and fetuses cannot be shielded from administered radionuclides. For this reason, many surgeons favor axillary node dissection to stage the nodes. Largely due to the typical delay in diagnosis, axillary nodes are more often positive in pregnant than in nonpregnant women.

As with other types of cancer in pregnant women, counseling following diagnosis in the first trimester should include a discussion of pregnancy termination to allow definitive therapeutic intervention at the earliest possible time without the potential for permanent injury to a surviving fetus. While definitive local surgery can safely be performed in the first trimester, radiation therapy and chemotherapy are considerably more risky. Delay in administration of systemic therapy can increase the risk of axillary spread. In the second and third trimesters,

chemotherapy (particularly anthracycline-based combinations) is both safe and effective (**Chap. 108**). Lumpectomy followed by adjuvant chemotherapy is frequently used; fluorouracil and cyclophosphamide with either doxorubicin or epirubicin have been given without major risk to the fetus. Taxanes and gemcitabine are also beginning to be used; however, safety data are sparse. Methotrexate and other folate antagonists are to be avoided because of effects on the fetal nervous system. Myelotoxic therapy is generally not administered after 33 or 34 weeks of gestation to allow 3 weeks off therapy before delivery for recovery of blood counts. Endocrine therapy and trastuzumab are unsafe during pregnancy. Experience with lapatinib is anecdotal, but no fetal malformations have been reported. Antiemetics and colony-stimulating factors are also considered safe. Women being treated into the postpartum period should not breast-feed their babies because of excretion of cancer chemotherapy agents, particularly alkylating agents, in milk.

Subsequent pregnancies following gestational breast cancer do not appear to influence relapse rate or overall survival. A meta-analysis found that pregnancy in breast cancer survivors was associated with a reduced the risk of dying from breast cancer by as much as 42%. This finding, however, is heavily confounded by the “healthy survivor effect”; women with more extensive or advanced disease are more likely to avoid pregnancy.

MELANOMA DURING PREGNANCY

Speculation about melanoma occurring during pregnancy based largely on anecdotal evidence and small case series concluded that it occurred with increased frequency, was more aggressive in its natural history, and was caused in part by the hormonal changes that also produced hyperpigmentation (so-called *melasma*) during pregnancy. However, more complete epidemiologic data suggest that melanoma is no more frequent in pregnant women than in nonpregnant women in the same age group, that melanoma is not more aggressive during pregnancy, and that hormones seem to have little or nothing to do with the etiology. Pregnant and nonpregnant women do not differ in the location of primary tumor, depth of primary tumor, tumor ulceration, or vascular invasion.

Suspicious lesions should be looked for and managed definitively with excisional biopsy during pregnancy. Wide excision with sampling of regional lymph nodes is warranted. If lymph nodes are involved, the course of action is less clear. Several agents have demonstrated some activity in melanoma, but none have been used during pregnancy. Adjuvant interferon α is toxic, and its safety in pregnancy has not been documented. Agents active in advanced disease include dacarbazine, IL-2, ipilimumab (antibody to CTLA-4), and in those with *BRAF* mutation V600E, a *BRAF* kinase inhibitor. In the setting of metastatic disease, abortion may be indicated so that systemic therapy can be initiated as soon as possible (**Chap. 105**).

Melanoma is one of the very few cancers that are well documented to metastasize transplacentally to the fetus, where it seems to have a predilection for the head and neck. It has a very grave prognosis in the offspring. Fortunately, transplacental spread is rare.

Pregnancy subsequent to the diagnosis and treatment of melanoma is not associated with an increased risk of melanoma recurrence.

HODGKIN'S DISEASE AND NON-HODGKIN'S LYMPHOMA DURING PREGNANCY

(**See Chap. 134**) Hodgkin's disease occurs mainly in the age range that coincides with child-bearing. However, Hodgkin's disease is not more common in pregnant than nonpregnant women. Hodgkin's disease is diagnosed in approximately 1 in 6000 pregnancies. It generally presents as a nontender lymph node swelling, most often in the left supraclavicular region. It may be accompanied by B symptoms (fever, night sweats, unexplained weight loss). Excisional biopsy is the preferred diagnostic procedure because fine-needle aspiration cannot reveal the architectural framework that is an essential component of Hodgkin's disease diagnosis. The stage at presentation appears to be unaffected by pregnancy. Women diagnosed in the second and third

trimester can be treated safely with combination chemotherapy, usually doxorubicin, bleomycin, vinblastine, and dacarbazine (ABVD). In general, the patient in the first trimester is asymptomatic, and a woman with a desired pregnancy can be followed until the second or third trimester when definitive multiagent chemotherapy can be safely given. Radiation therapy is not given during pregnancy and is not necessary for optimal management of the pregnant patient. If symptoms requiring treatment appear during the first trimester, anecdotal evidence suggests that Hodgkin's disease symptoms can be controlled with weekly low-dose vinblastine. Such an approach has been safely used to avoid termination of pregnancy. Pregnancy does not have an adverse effect on treatment outcome.

Non-Hodgkin's lymphomas are more unusual in pregnancy (approximately 0.8 per 100,000 pregnancies), but are usually tumors with an aggressive natural history, such as diffuse large B cell lymphoma, Burkitt's lymphoma, or peripheral T cell lymphoma. Diagnosis relies on an excisional biopsy of a tumor mass, not fine-needle aspiration. Staging evaluation is generally limited to ultrasound or MRI examinations. Diagnosis in the first trimester should prompt termination of the pregnancy followed by definitive treatment with combination chemotherapy, because aggressive lymphomas are not likely to be held at bay with single-agent chemotherapy. Women diagnosed in the second or third trimesters can be treated with standard chemotherapy, such as with cyclophosphamide, doxorubicin, vincristine, and prednisone (CHOP). The experience with rituximab in this setting is anecdotal. However, infants born of mothers who have received rituximab may have transient delay in B cell development that typically normalizes by 6 months. The treatment outcome is similar in lymphomas diagnosed in pregnant and nonpregnant women of the same clinical stage.

THYROID CANCER DURING PREGNANCY

(**See Chap. 405**) Thyroid cancer, along with melanomas, brain tumors, and lymphomas, are cancers that are increasing in incidence in the general population. Thyroid cancers are rising faster among women in North America than the other increasing tumor types. The Endocrine Society has developed practice guidelines to inform the management of patients with thyroid disease during pregnancy (<http://www.endocrine.org/~media/endosociety/Files/Publications/Clinical%20Practice%20Guidelines/Thyroid-Exec-Summ.pdf>). Thyroid nodules 1 cm or larger are approached by fine-needle aspiration. If a malignancy is diagnosed, surgery is generally recommended in the second and third trimesters. However, surgical complications appear to be twice as common when the patient is pregnant. Because the growth of thyroid tumors is often indolent, surgery can safely be postponed until after the first trimester. Patients with follicular cancer or early papillary cancer can be observed until the postpartum period. The fetal thyroid begins trapping iodine by 12 weeks of gestation and does so with very high avidity. Even small doses of radioactive iodine given during pregnancy can completely ablate the fetal thyroid with serious consequences for the fetus and should be avoided throughout pregnancy. Radioactive iodine can be safely administered after delivery. Patients with a history of thyroid cancer who become pregnant should be maintained on thyroid hormone replacement during pregnancy because of the adverse impact of maternal hypothyroidism on the fetus. Women who are breast-feeding should not be treated with radioactive iodine, and women treated with radioactive iodine should not become pregnant for 6–12 months after treatment.

The assessment of thyroid function during pregnancy is challenging because of the physiologic changes that occur during pregnancy. Women who have previously been treated for thyroid cancer are at risk of hypothyroidism. The demand for thyroid hormone increases during pregnancy, and doses to maintain normal function may increase by 30–50%. Total T_4 levels are higher during pregnancy, but target therapeutic levels also increase (**Table 124e-4**). It is recommended that the upper and lower limits of the laboratory range be multiplied by 1.5 in the second and third trimester to establish a pregnancy-specific normal range. The target thyroid-stimulating hormone (TSH) level is <2.5 mIU/L.

TABLE 124e-4 THYROID FUNCTION TEST DURING PREGNANCY (MEAN LEVELS)

	Nonpregnant	First Trimester	Second Trimester	Third Trimester
Thyroid-stimulating hormone (mIU/L)	1.38	0.91	1.03	1.32
Total thyroxine (µg/dL)	7.35	10.98	11.88	11.08

Source: Based on the National Health and Nutrition Examination Survey III (NHANES III) (OP Soldin et al: *Ther Drug Monit* 17:303, 2007).

GESTATIONAL TROPHOBLASTIC DISEASE

(See Chap. 117) Gestational trophoblastic disease encompasses hydatidiform mole, choriocarcinoma, placental site trophoblastic tumor, and assorted miscellaneous and unclassifiable trophoblastic tumors. Moles are the most common, occurring in 1 in 1500 pregnancies in the United States. The incidence is higher in Asia. In general, if the serum level of β -human chorionic gonadotropin (β -hCG) returns to normal after surgical removal (evacuation) of the mole, the illness is considered gestational trophoblastic disease. By contrast, if the β -hCG level remains elevated after mole evacuation, the patient is considered to have gestational trophoblastic neoplasia. Choriocarcinoma occurs in 1 in 25,000 pregnancies. Maternal age >45 years and prior history of molar pregnancy are risk factors. A previous molar pregnancy makes choriocarcinoma about 1000 times more likely to occur (incidence 1–2%).

Hydatidiform moles are characterized by clusters of villi with hydropic changes, trophoblastic hyperplasia, and absence of fetal blood vessels. Invasive moles are distinguished by invasion of the myometrium. Placental site trophoblastic tumors are composed mainly of cytotrophoblast cells arising at the site of placental implantation. Choriocarcinomas contain anaplastic trophoblastic tissue with both cytotrophoblast and syncytiotrophoblast features and no identifiable villi.

Moles can be partial, typically associated with fetal tissue, or complete, typically not associated with any fetal or embryonic tissue. Partial moles have a distinct molecular origin and usually are smaller tumors with less hydropic villi and considerably less potential for persistent or malignant disease. Partial moles result from fertilization of an egg by two sperm, resulting in diandric triploidy. Complete moles usually have a 46,XX genotype; 95% develop by a single male sperm fertilizing an empty egg and undergoing gene duplication (diandric diploidy); 5% develop from dispermic fertilization of an empty egg (diandric dispermy).

Women with molar gestations often present with first-trimester bleeding, disproportionately high serum β -hCG levels for menstrual age, unusually large uterine size for menstrual age, hyperemesis gravidarum, theca lutein cysts in the ovaries (due to β -hCG stimulation), and hyperthyroidism (due to cross-reactivity of β -hCG and TSH) and may develop preeclampsia before 20 weeks of menstrual age. Pelvic ultrasound imaging of complete moles shows absence of fetal parts, an enlarged echo-bright, hydropic placenta in an enlarged uterus, and enlarged multicystic ovaries. If the diagnosis is uncertain at the initial examination and the pregnancy is desired, then a serum β -hCG level should be obtained and the examination repeated in a week. If no

embryo is seen within 7–10 days and the serum β -hCG is elevated, then this is a nonviable pregnancy that should be evacuated. Diagnosis of partial molar pregnancies can be more difficult because an embryo or fetus with visible heart motion is usually present, and the hydropic changes in the placenta, uterine enlargement, and elevations of β -hCG are not usually as dramatic. Although an embryo or fetus is present, it rarely grows normally with normal anatomy, and repeated ultrasound examinations usually make the diagnosis. Amniocentesis will also make the diagnosis by demonstration of triploidy.

Patients with molar pregnancies require prompt uterine evacuation with suction curettage, which may be complicated by very heavy bleeding. Following evacuation of complete moles, approximately 20% will result in persistent, invasive, or metastatic disease. Partial moles are considerably less likely (<5%) to result in persistent disease. Patients should be monitored with serial determinations of serum β -hCG until the values fall below the lower limit of the assay and remain low for at least 6 months. Patients should be advised not to become pregnant for at least 12 months.

A variety of criteria have been used to make the diagnosis of post-molar gestational trophoblastic disease, but current consensus guidelines as adopted by the International Federation of Gynecology and Obstetrics are listed below:

1. A β -hCG level plateau of four values plus or minus 10% recorded over a 3-week duration (days 1, 7, 14, and 21)
2. A β -hCG level increase of more than 10% in three values recorded over a 2-week duration (days 1, 7, and 14)
3. Persistence of detectable β -hCG for more than 6 months after molar evacuation

About half of choriocarcinomas develop after a molar pregnancy, and half develop after ectopic pregnancy or, rarely, after a normal full-term pregnancy. Disease is classified as stage I if it is confined to the uterus, stage II if disease is limited to genital structures (~30% have vaginal involvement), stage III if disease has spread to the lungs but no other organs, and stage IV if disease has spread to liver, brain, or other organs.

Patients without widely metastatic disease are generally managed with single-agent methotrexate (either 30 mg/m² IM weekly until β -hCG normalizes or 1 mg/kg IM every other day for four doses followed by leucovorin 0.1 mg/kg IV 24 h after methotrexate), which cures >90% of patients. Patients with very high β -hCG levels, presenting >4 months after a pregnancy, with brain or liver metastases, or failing to be cured by single-agent methotrexate are treated with combination chemotherapy. Etoposide, methotrexate, and dactinomycin alternating with cyclophosphamide and vincristine (EMA-CO) is the most commonly used regimen, producing long-term survival in >80% of patients. Brain metastases can usually be controlled with brain radiation therapy. The vast majority of choriocarcinomas can be cured with chemotherapy alone. Hysterectomy is reserved for women who have completed their child-bearing, women with chemotherapy-resistant disease in the uterus, and women with rare placental site trophoblastic tumors confined to the uterus because these tumors are less reliably sensitive to chemotherapy. Women cured of trophoblastic disease who have not undergone hysterectomy do not appear to have increased risk of fetal abnormalities or maternal complications with subsequent pregnancies.

125 Late Consequences of Cancer and Its Treatment

Carl E. Freter, Dan L. Longo

There are over 10 million American cancer survivors. The vast majority of these will bear some mark of their cancer and its treatment, and a large proportion will experience long-term consequences including medical problems, psychosocial dysfunction, economic hardship, sexual dysfunction, and discrimination regarding employment and insurance. Many of these problems are directly related to cancer treatment. As patients survive longer from more types of malignancies, we are increasingly recognizing the biologic toll our very imperfect therapies take in terms of morbidity and mortality. The human face of these consequences of therapy confronts the cancer specialist who treats them every day. Although long-term survivors of childhood leukemias, Hodgkin's lymphoma, and testicular cancer, as examples, have taught us much about the consequences of cancer treatment, we keep learning more as patients survive longer with newer therapies. Newer "targeted" chemotherapy drugs have their own, often unique, long-term toxicities about which we remain in a learning process. Cancer "survivorship" clinics are increasing to expressly follow patients for long-term toxicities of cancer treatment.

The pace of developing therapies that mitigate treatment-related consequences has been slow, partly due to an understandable aversion to alter regimens that work and partly due to a lack of new, effective, less toxic therapeutic agents with less "collateral damage" to replace known agents with known toxicities. The types of damage from

cancer treatment vary. Often, a final common pathway is irreparable damage to DNA. Surgery can create dysfunction, including blind gut loops with absorption problems and loss of function of removed body parts. Radiation may damage end-organ function, for example, loss of potency in prostate cancer patients, pulmonary fibrosis, and neurocognitive impairment, and may act as a direct carcinogen. Cancer chemotherapy can be a direct carcinogen and has a kaleidoscope of other toxicities discussed in this chapter. **Table 125-1** lists the late effects of cancer treatment.

TABLE 125-1 LATE EFFECTS OF CANCER THERAPY

Surgical Procedure		Effect
Amputation		Functional loss
Lymph node dissection		Risk of lymphedema
Ostomy		Psychosocial impact
Splenectomy		Risk of sepsis
Adhesions		Risk of obstruction
Bowel anastomoses		Malabsorption syndromes
Radiation Therapy		Effect
Organ		
Bone		Premature termination of growth, osteonecrosis
Soft tissues		Atrophy, fibrosis
Brain		Neuropsychiatric deficits, cognitive dysfunction
Thyroid		Hypothyroidism, Graves' disease, cancer
Salivary glands		Dry mouth, caries, dysgeusia
Eyes		Cataracts
Heart		Pericarditis, myocarditis, coronary artery disease
Lung		Pulmonary fibrosis
Kidney		Decreased function, hypertension
Liver		Decreased function
Intestine		Malabsorption, stricture
Gonads		Infertility, premature menopause
Any		Secondary neoplasia
Chemotherapy		Effect
Organ	Drug	
Bone	Glucocorticoids	Osteoporosis, avascular necrosis
Brain	Methotrexate, cytarabine, others	Neuropsychiatric deficits, cognitive decline?
Peripheral nerves	Vincristine, platinum, taxanes	Neuropathy, hearing loss
Eyes	Glucocorticoids	Cataracts
Heart	Anthracyclines, trastuzumab	Cardiomyopathy
Lung	Bleomycin	Pulmonary fibrosis
	Methotrexate	Pulmonary hypersensitivity
Kidney	Platinum, others	Decreased function, hypomagnesemia
Liver	Various	Altered function
Gonads	Alkylating agents, others	Infertility, premature menopause
Bone marrow	Various	Aplasia, myelodysplasia, secondary leukemia

The first goal of therapy is to eradicate or control the malignancy. Late treatment consequences are, indeed, testimony to the increasing success of such treatment. Their occurrence sharply underlines the necessity to develop more effective therapies with less long-term morbidity and mortality. At the same time, a sense of perspective and relative risk is necessary; fear of long-term complications should not prevent the application of effective, particularly curative, cancer treatment.

CARDIOVASCULAR DYSFUNCTION

CHEMOTHERAPEUTIC AGENTS

Cardiovascular toxicity of cancer chemotherapeutic agents includes dysrhythmias, cardiac ischemia, cardiomyopathic congestive heart failure (CHF), pericardial disease, and peripheral vascular disease. Because these cardiac toxicities are difficult to distinguish from disease that is not associated with cancer treatment, clear etiologic implication of cancer chemotherapeutic agents may be difficult. Cardiovascular complications occurring in an unexpected clinical setting in patients who have undergone cancer therapy are often important in raising suspicion. Dose-dependent myocardial toxicity of anthracyclines with characteristic myofibrillar dropout is pathologically pathognomonic on endomyocardial biopsy. Anthracycline cardiotoxicity occurs through a root mechanism of chemical free radical damage. Fe³⁺-doxorubicin complexes damage DNA, nuclear and cytoplasmic membranes, and mitochondria. About 5% of patients receiving >450–550 mg/m² of doxorubicin will develop CHF. Cardiotoxicity in relation to the dose of anthracycline is clearly not a step function, but rather a continuous function, and occasional patients are seen with CHF at substantially lower doses. Advanced age, other concomitant cardiac disease, hypertension, diabetes, and thoracic radiation therapy are all important cofactors in promoting anthracycline-associated CHF. The risk of cardiac failure appears to be substantially lower when doxorubicin is administered by continuous infusion. Anthracycline-related CHF is difficult to reverse and has a mortality rate as high as 50%, making prevention crucial. Some anthracyclines such as mitoxantrone are associated with less cardiotoxicity, and continuous-infusion regimens and liposomally encapsulated doxorubicin are associated with less cardiotoxicity. Dexrazoxane, an intracellular iron chelator, may limit anthracycline toxicity, but the concern of limiting chemotherapeutic efficacy has somewhat limited its use. Monitoring patients for cardiac toxicity typically involves periodic gated nuclear cardiac blood pool ejection fraction testing (multigated acquisition scan [MUGA]) or cardiac ultrasonography. More recently, cardiac magnetic resonance imaging (MRI) has been used, but MRI is not standard or widespread. Testing is performed more frequently at higher cumulative doses, with additional risk factors, and certainly for any newly developing CHF or other symptoms of cardiac dysfunction.

After anthracyclines, trastuzumab is the next most frequent cardiotoxic drug currently in use. Trastuzumab is frequently used as adjuvant breast cancer therapy, sometimes in conjunction with anthracyclines, which is believed to result in additive or possibly synergistic toxicity. In contrast to anthracyclines, cardiotoxicity is not dose-related, is usually reversible, is not associated with pathologic changes of anthracyclines on cardiac myofibrils, and has a different biochemical mechanism inhibiting intrinsic cardiac repair mechanisms. Toxicity is typically routinely monitored every three to four doses using functional cardiac testing as mentioned earlier for anthracyclines.

Other cardiotoxic drugs include lapatinib, phosphoramidate mustards (cyclophosphamide), ifosfamide, interleukin 2, ponatinib, imatinib, and sunitinib.

RADIATION THERAPY

Radiation therapy that includes the heart can cause interstitial myocardial fibrosis, acute and chronic pericarditis, valvular disease, and accelerated premature atherosclerotic coronary artery disease. Repeated or high (>6000 cGy) radiation doses are associated with

greater risk, as is concomitant or distant cardiotoxic cancer chemotherapy exposure. Symptoms of acute pericarditis, which peaks about 9 months after treatment, include dyspnea, chest pain, and fever. Chronic constrictive pericarditis may develop 5–10 years following radiation therapy. Cardiac valvular disease includes aortic insufficiency from fibrosis or papillary muscle dysfunction resulting in mitral regurgitation. A threefold increased risk of fatal myocardial infarction is associated with mantle field radiation with accelerated coronary artery disease. Carotid radiation similarly increases the risk of embolic stroke.

TREATMENT CHEMOTHERAPEUTIC/RADIATION-INDUCED CARDIOVASCULAR DISEASE

Therapy for chemotherapeutic/radiation-induced cardiovascular disease is essentially the same as therapy for disease not associated with cancer treatment. Discontinuation of the offending agent is the first step. Diuretics, fluid and sodium restriction, and antiarrhythmic agents are often useful for acute symptoms. Afterload reduction with angiotensin-converting enzyme (ACE) inhibitors or, in some cases, β -adrenergic blockers (carvedilol) often is of significant benefit, and digitalis may be helpful as well.

A hybrid discipline of “cardio-oncology” has been developing in clinics to expressly follow chemotherapy-treated patients for cardiotoxicity. The goals are early intervention using more sensitive techniques, management of cardiotoxicity before it becomes symptomatic, and using clinical trials to identify cardioprotective strategies.

PULMONARY DYSFUNCTION

CHEMOTHERAPEUTIC AGENTS

Bleomycin generates activated free radical oxygen species and causes pneumonitis associated with a radiographic or interstitial ground-glass appearance diffusely throughout both lungs, often worse in the lower lobes. A nonproductive cough with or without fever may be an early sign. This toxicity is dose-related and dose-limiting. The diffusion capacity of the lungs for carbon dioxide (DL_{CO}) is a sensitive measure of toxicity and recovery, and a baseline value is generally obtained for future comparison prior to bleomycin therapy. Additive or synergistic risk factors include age, prior lung disease, and concomitant use of other chemotherapy, lung irradiation, and high concentrations of inspired oxygen. Other chemotherapeutic agents notable for pulmonary toxicity include mitomycin, nitrosoureas, doxorubicin with radiation, gemcitabine combined with weekly docetaxel, methotrexate, and fludarabine. High-dose alkylating agents, cyclophosphamide, ifosfamide, and melphalan are frequently used in the hematopoietic stem cell transplant setting, often with whole-body radiation. This therapy may result in severe pulmonary fibrosis and/or pulmonary venoocclusive disease.

RADIATION THERAPY

Risk factors for radiation pneumonitis include advanced age, poor performance status, preexisting compromised pulmonary function, and radiation volume and dose. The dose “threshold” is thought to be in the range of 5 to 20 Gy. Hypoxemia and dyspnea on exertion are characteristic. Fine, high-pitched “Velcro rales” may be an accompanying physical finding, and fever, cough, and pleuritic chest pain are common symptoms. The DL_{CO} is the most sensitive measure of pulmonary functional impairment, and ground-glass infiltrates often correspond with relatively sharp edges to the irradiated volume, although the pneumonitis may progress beyond the field and even occasionally involve the contralateral unirradiated lung.

Chemotherapy- and radiation-induced pneumonitis is generally very corticosteroid responsive, except in the case of nitrosoureas. Prednisone 1 mg/kg is often used to control acute symptoms and pulmonary dysfunction with a generally slow taper. Prolonged glucocorticoid therapy requires gastrointestinal protection with proton pump inhibitors, management of hyperglycemia, heightened infection management, and treatment of steroid-induced osteoporosis. Antibiotics, bronchodilators, oxygen in only necessary doses, and diuretics may all play an important role in management of pneumonitis, and consultation with a pulmonologist should be routinely undertaken. Amifostine has been studied as a pulmonary radioprotectant, with inconclusive results, and is associated with skin rash, fatigue, and nausea; hence, it is not considered standard therapy at this time. Transforming growth factor β (TGF- β) is believed to be a major inducer of radiation fibrosis and represents a therapeutic target for development of anti-TGF- β therapies.

NEUROLOGIC DYSFUNCTION

CHEMOTHERAPEUTIC AGENTS

Chemotherapy- and radiation-induced neurologic dysfunction is unfortunately increasing in both incidence and severity as a result of improved supportive care leading to more aggressive regimens and longer cancer survivorship allowing the development of late toxicity. Direct effects on myelin, glial cells, and neurons have all been implicated, with alterations in cellular cytoskeleton, axonal transport, and cellular metabolism as mechanisms.

Vinca alkaloids produce a characteristic “stocking-glove” neuropathy with numbness and tingling advancing to loss of motor function, which is highly dose related. Distal sensorimotor polyneuropathy prominently involves loss of deep tendon reflexes with initially loss of pain and temperature sensation, followed by proprioceptive and vibratory loss. This requires careful patient history and physical examination by experienced oncologists to decide when the drug must be stopped due to toxicity. Milder toxicity often slowly completely resolves. Vinca alkaloids may sometimes be associated with jaw claudication, autonomic neuropathy, ileus, cranial nerve palsies, and, in severe cases, encephalopathy, seizures, and coma.

Cisplatin is associated with sensorimotor neuropathy and hearing loss, especially at doses >400 mg/m², requiring audiometry in patients with preexisting hearing compromise. Carboplatin is often substituted in such cases given its lesser effect on hearing.

Many of the agents that target kinase enzymes in tumor cells and 5-fluorouracil congeners produce dysesthesias and painful hands and feet known as hand-foot syndrome or palmar-plantar erythrodysesthesia. Symptoms usually abate when the agent is stopped.

Neurocognitive dysfunction has been well described in childhood survivors of acute lymphoblastic leukemia (ALL) treatment, including intrathecal methotrexate or cytosine arabinoside in conjunction with prophylactic cranial irradiation. Methotrexate alone may cause acute leukoencephalopathy characterized by somnolence and confusion that is often reversible. Acute toxicity is dose related, especially at doses >3 g/m², with younger patients being at greater risk. Subacute methotrexate toxicity occurs weeks after therapy and is often ameliorated with glucocorticoid therapy. Chronic methotrexate toxicity (leukoencephalopathy) develops months or years after treatment and is characterized clinically as progressive loss of cognitive function and focal neurologic signs, which are irreversible, promoted by synchronous or metachronous radiation therapy, and more pronounced at a younger age.

Neurocognitive decline following chemotherapy alone occurs notably in breast cancer patients receiving adjuvant chemotherapy; this has been referred to as “chemo brain.” It is clinically associated with impaired memory, learning, attention, and speed of information

processing. There is no clear mechanistic explanation for its cause and no clearly effective therapy. This entity is justifiably attracting more attention and clearly needs to be studied to develop effective therapy or prophylaxis.

Many cancer patients experience intrusive or debilitating concerns about cancer recurrence following successful therapy. In addition, these patients may experience job, insurance, stress, relationship, financial, and sexual difficulties. Oncologists need to ask about and address these issues explicitly with patients and provide appropriate counseling or support systems. Suicidal ideation and suicide have an increased incidence in cancer patients and survivors.

RADIATION THERAPY

Acute radiation central nervous system (CNS) toxicity occurs within weeks; is characterized by nausea, drowsiness, hypersomnia, and ataxia; and is most often associated with recovery. Early delayed toxicity occurring weeks to 3 months following therapy is associated with similar symptoms as acute toxicity and is pathologically associated with reversible demyelination. Chronic, late radiation injury occurs 9 months to up to 10 years following therapy. Focal necrosis is a common pathologic finding, and glucocorticoid therapy may be helpful. Diffuse radiation injury is associated with global CNS neurologic dysfunction and diffuse white matter changes on computed tomography (CT) or MRI. Pathologically, small vessel changes are prominent. Glucocorticoids may be symptomatically useful but do not alter the course. Necrotizing encephalopathy is the most severe form of radiation injury and almost always is associated with chemotherapy, notably methotrexate.

Cranial radiation may also be associated with an array of endocrine abnormalities with disruption of normal pituitary/hypothalamic axis function, and a high index of suspicion needs to be maintained to identify and treat this toxicity.

Radiation-associated spinal cord injury (myelopathy) is highly dose-dependent and rarely occurs with modern radiation therapy. An early, self-limited form involving electric sensations down the spine on neck flexion (Lhermitte’s sign) is seen 6–12 weeks after treatment and generally resolves over weeks. Peripheral nerve toxicity is quite rare owing to relative radiation resistance.

HEPATIC DYSFUNCTION

CHEMOTHERAPEUTIC AGENTS

Long-term hepatic damage from standard chemotherapy regimens is rare. Long-term methotrexate or high-dose chemotherapy alone or with radiation therapy, for example, in preparative regimens for bone marrow transplantation, may result in venoocclusive disease of the liver. This potentially lethal complication classically presents with anicteric ascites, elevated alkaline phosphatase, and hepatosplenomegaly. Pathologically, there is venous congestion, epithelial cell proliferation, and hepatocyte atrophy progressing to frank fibrosis. Frequent monitoring of liver function tests during any chemotherapy is necessary to avoid both idiosyncratic and expected toxicities.

Certain nucleoside drugs have been associated with hepatic dysfunction; however, this complication is rare in oncology.

RADIATION THERAPY

Hepatic radiation damage depends on dose, volume, fractionation, preexisting liver disease, and synchronous or metachronous chemotherapy. In general, radiation doses to the liver >1500 cGy can produce hepatic dysfunction with a steep dose-injury curve. Radiation-induced liver disease closely mimics hepatic venoocclusive disease.

RENAL/BLADDER DYSFUNCTION

Cisplatin produces reversible decrements in renal function, but may also produce severe irreversible toxicity in the presence of renal disease and may predispose to accentuated damage with subsequent renal insults. Cyclophosphamide and ifosfamide, as prodrugs primarily

activated in the liver, have cleavage products (acrolein) that can produce hemorrhagic cystitis. This can be prevented with the free radical scavenger MESNA (mercaptoethane sulfonate), which is required for ifosfamide administration. Hemorrhagic cystitis caused by these agents may predispose to bladder cancer.

REPRODUCTIVE AND ENDOCRINE DYSFUNCTION

CHEMOTHERAPEUTIC AGENTS

Alkylating agents are associated with the highest rates of male and female infertility, which is directly dependent on age, dose, and duration of treatment. The age at treatment is an important determinant of fertility outcome, with prepubertal patients having the highest tolerance. Ovarian failure is age related, and females who resume menses after treatment are still at increased risk for premature menopause. Males generally have reversible azoospermia during lower intensity alkylator chemotherapy, and long-term infertility is associated with doses of cyclophosphamide >9 g/m² and with high-intensity therapy, such as that used in hematopoietic stem cell transplantation. Males undergoing potentially sterilizing chemotherapy should be offered sperm banking. Gonadotropin-releasing hormone (GnRH) analogs remain experimental to preserve ovarian function. Assisted reproductive technologies can be helpful to couples with chemotherapy-induced infertility.

RADIATION THERAPY

Testicles and ovaries in prepubertal patients are less sensitive to radiation damage; spermatogenesis is affected by low doses of radiation, and complete azoospermia occurs at 600–700 cGy. Leydig cell dysfunction, in contrast, occurs at <2000 cGy, and hence, endocrine function is lost at much higher radiation doses than spermatogenesis. Erectile dysfunction occurs in up to 80% of men treated with external-beam radiation therapy for prostate cancer. Sildenafil may be useful in reversing erectile dysfunction. Ovarian function damage with radiation is age related and occurs at doses of 150–500 cGy. Premature induction of menopause can have serious medical and psychological sequelae. Hormone replacement therapy is often contraindicated (as in estrogen receptor–positive breast cancer). Attention must be paid to maintenance of bone mass with calcium and vitamin D supplements and oral bisphosphonates, and bone mass should be monitored using bone density determinations. Paroxetine, clonidine, pregabalin, and other drugs may be useful in symptomatically controlling hot flashes.

Long-term survivors of childhood cancer (e.g., ALL) who have received cranial radiation may have altered leptin biology and growth hormone deficiency, leading to obesity and reduced strength, exercise tolerance, and bone density.

Radiation therapy to the neck (e.g., in Hodgkin's lymphoma) may lead to hypothyroidism, Graves' disease, thyroiditis, and thyroid malignancies. Thyroid-stimulating hormone (TSH) is followed routinely in such patients to prevent hypothyroidism, and to suppress persistently elevated levels of TSH which may cause or drive thyroid cancer.

OCULAR COMPLICATIONS

Cataracts may be caused by glucocorticoids, depending on duration and dose; radiation therapy; and uncommonly tamoxifen. Orbital radiation therapy may cause blindness.

ORAL COMPLICATIONS

Radiation therapy can produce xerostomia (dry mouth), with an attendant increase in caries and poor dentition. Taste and appetite may be suppressed. Bisphosphonate use may result in osteonecrosis of the jaw.

RAYNAUD'S PHENOMENON

Up to 40% of patients treated with bleomycin may develop Raynaud's phenomenon as a result of an unknown mechanism.

SECOND MALIGNANCIES

Second malignancies in patients cured of cancer are a major cause of death, and treated cancer patients must be monitored for their occurrence. The induction of second malignancies is governed by the complex interplay of a number of factors including age, gender, environmental exposures, genetic susceptibility, and cancer treatment itself. In a number of settings, the events leading to the primary cancer themselves increase the risk of second malignancies. Patients with lung cancer are at increased risk of esophageal and head and neck cancers, and vice versa, due to shared risk factors including alcohol and tobacco abuse. Indeed, the risk of developing a second primary head and neck, esophageal, or lung cancer is also increased in these patients. Patients with breast cancer are at increased risk of breast cancer in the opposite breast. Patients with Hodgkin's lymphoma are at risk for non-Hodgkin's lymphomas. Genetic cancer syndromes (e.g., multiple endocrine neoplasia or Li-Fraumeni, Lynch's, Cowden's, and Gardner's syndromes) are examples of genetically based second malignancies of specific types. Cancer treatment itself does not appear to be responsible for the risk of these secondary malignancies. Deficient DNA repair can greatly increase the risk of cancers from DNA-damaging agents, as in ataxia-telangiectasia. Importantly, the risk of treatment-related second malignancies is at least additive and often synergistic with combined chemotherapy and radiation therapy, and hence for such combined-therapy treatment approaches, it is important to establish the necessity of each in the treatment program. All of these patients require special surveillance or, in some cases, prophylactic surgery as part of appropriate treatment and follow-up.

CHEMOTHERAPEUTIC AGENTS

Chemotherapy is significantly associated with two fatal second malignancies, acute leukemia and myelodysplastic syndromes. Two types of leukemia have been described; in patients treated with alkylating agents, acute myeloid leukemia is associated with deletions in chromosome 5 or 7. The lifetime risk is about 1–5%, is increased by radiation therapy, and increases with age. The incidence of these leukemias peaks at 4–6 years, with risk returning close to baseline at 10 years. The other type of acute myeloid leukemia is related to therapy with topoisomerase inhibitors, is associated with chromosome 10q23 translocations, has an incidence of $<1\%$, and generally occurs 1.5–3 years after treatment. Both of these acute myeloid leukemias are refractory to treatment and have a high mortality. The development of myelodysplastic syndromes is increased following chemotherapy, and these are often associated with leukemic progression and a dismal prognosis.

RADIATION THERAPY

Patients receiving radiation have an increasing and lifelong risk of second malignancies that is 1–2% in the second decade following treatment but increases to $>25\%$ after 25 years. These malignancies include cancers of the thyroid and breast, sarcomas, and CNS cancers, which often tend to be aggressive and have a poor prognosis. An example of organ-, age-, and sex-dependent radiation-induced secondary malignancy is breast cancer, in which the risk is small with radiation in women under age 30 but increases about 20-fold over baseline in women over 30. A 25-year-old woman treated with mantle radiation for Hodgkin's lymphoma has a 29% actuarial risk of developing breast cancer by age 55.

HORMONAL THERAPY

Treatment of breast cancer with tamoxifen for 5 years or longer is associated with a 1–2% risk of endometrial cancer. Surveillance is generally effective at finding these cancers at an early stage. The risk of mortality from tamoxifen-induced endometrial cancer is low compared to the benefit of tamoxifen as adjuvant therapy for breast cancer.

IMMUNOSUPPRESSIVE THERAPY

Immunosuppressive therapy, as used in allogeneic bone marrow transplantation, particularly with T cell depletion using antithymocyte globulin or other means, increases the risk of Epstein Barr virus-associated

TABLE 125-2 LONG-TERM TREATMENT EFFECTS BY CANCER TYPE

Cancer Type	Late Effects
Pediatric cancers	Majority have at least one late effect 30% with moderate/severe problems Cardiovascular: radiation, anthracyclines Lungs: radiation Skeletal abnormalities: radiation Psychological, cognitive, and sexual problems Second neoplasms significant cause of death
Hodgkin's lymphoma	Thyroid dysfunction: radiation Premature coronary artery disease: radiation Gonadal dysfunction: chemotherapy Postsplenectomy sepsis Myelodysplasia Acute myeloid leukemia Non-Hodgkin's lymphomas Breast cancer, lung cancer, and melanoma Fatigue, psychological and sexual problems Peripheral neuropathy
Non-Hodgkin's lymphoma	Myelodysplasia Acute leukemia Bladder cancer Peripheral neuropathy
Acute leukemia	Second malignancies: hematologic, solid tumors Neuropsychiatric dysfunction Subnormal growth Thyroid abnormalities Infertility
Bone marrow stem cell transplantation	Infertility Graft-versus-host disease (allogeneic transplant) Psychosexual dysfunction.
Head and neck cancer	Poor dentition, dry mouth, poor nutrition: radiation
Breast cancer	Tamoxifen: endometrial cancer, blood clots Aromatase inhibitors: osteoporosis, arthritis Cardiomyopathy: anthracycline ± radiation, trastuzumab Acute leukemia Hormone deficiency symptoms: hot flashes, vaginal dryness, dyspareunia Psychosocial dysfunction "Chemo brain"
Testicular cancer	Raynaud's phenomenon Renal dysfunction Pulmonary dysfunction Retrograde ejaculation: surgery 15% sexual dysfunction
Colon cancer	Major risk is second colon cancer. Quality of life high in survivors
Prostate cancer	Impotence Urinary incontinence (0–15%) Chronic proctitis, prostatitis/cystitis: radiation

B cell lymphoproliferative disorder. The incidence at 10 years after T cell depletion is 9–12%. Discontinuing immunosuppressive therapy, if possible, is often associated with complete disease regression.

RECOMMENDATIONS FOR FOLLOW-UP

All former cancer patients should be followed indefinitely. This is most often done by oncologists, but demographic changes suggest that more primary care physicians will need to be trained in the follow-up of treated cancer patients in remission. Cancer patients need to be educated about signs and symptoms of recurrence and potentially adverse effects related to therapy. Localized pain or palpable abnormality in a previously radiated field should prompt radiographic evaluation. Screening tests, when available and validated, should be used on a routine and regular basis (e.g., mammography and Pap smear), particularly in patients receiving radiation to specific organs. Annual mammography should start no later than 10 years after breast radiation. Patients receiving radiation fields encompassing thyroid tissue should have regular thyroid exams and TSH testing. Patients treated with alkylating agents or topoisomerase inhibitors should have a complete blood count every 6–12 months, and cytopenias, abnormal cells on peripheral smear, or macrocytosis should be evaluated with bone marrow biopsy and aspirate, to include cytogenetics, flow cytometry, or fluorescence in situ hybridization (FISH) studies as appropriate.

As the population of cancer survivors lives longer and grows, cancer survivorship has become an increasingly recognized subject, and the Institute of Medicine and National Research Council have published a monograph entitled *From Cancer Patient to Cancer Survivor: Lost in Transition*. The monograph proposes a plan that would inform clinicians caring for cancer survivors in complete detail of their previous treatments, complications thereof, signs and symptoms of late effects, and recommended screening and follow-up procedures. [Table 125-2](#) lists long-term treatment effects by cancer type.

OUTLOOK

Clearly, the challenge for the future is to combine chemotherapy, targeted agents, biologic therapies, radiation, and surgery to produce better outcomes with less toxicity, including late effects of therapy. This is easily said but less easily accomplished. As treatment becomes more effective in new patient populations (ovarian, bladder, anal, and laryngeal cancers, for example), we will expect to discover new populations at risk for late effects. These populations will need to be followed carefully, so that such effects are recognized and treated. Cancer survivors represent an underused resource for prevention studies. Childhood cancer survivors, especially, suffer multiple chronic health impairments. The incidence of these late treatment consequences appears to have no plateau with age, throwing in stark relief the necessity of close monitoring and therapies with fewer late consequences of treatment.

126 Iron Deficiency and Other Hypoproliferative Anemias

John W. Adamson

Anemias associated with normocytic and normochromic red cells and an inappropriately low reticulocyte response (reticulocyte index <2–2.5) are *hypoproliferative anemias*. This category includes early iron deficiency (before hypochromic microcytic red cells develop), acute and chronic inflammation (including many malignancies), renal disease, hypometabolic states such as protein malnutrition and endocrine deficiencies, and anemias from marrow damage. **Marrow damage states are discussed in Chap. 130.**

Hypoproliferative anemias are the most common anemias, and in the clinic, iron deficiency anemia is the most common of these followed by the anemia of inflammation. The anemia of inflammation, similar to iron deficiency, is related in part to abnormal iron metabolism. The anemias associated with renal disease, inflammation, cancer, and hypometabolic states are characterized by a suboptimal erythropoietin response to the anemia.

IRON METABOLISM

Iron is a critical element in the function of all cells, although the amount of iron required by individual tissues varies during development. At the same time, the body must protect itself from free iron, which is highly toxic in that it participates in chemical reactions that generate free radicals such as singlet O₂ or OH⁻. Consequently, elaborate mechanisms have evolved that allow iron to be made available for physiologic functions while at the same time conserving this element and handling it in such a way that toxicity is avoided.

The major role of iron in mammals is to carry O₂ as part of hemoglobin. O₂ is also bound by myoglobin in muscle. Iron is a critical element in iron-containing enzymes, including the cytochrome system in mitochondria. Iron distribution in the body is shown in **Table 126-1**. Without iron, cells lose their capacity for electron transport and energy metabolism. In erythroid cells, hemoglobin synthesis is impaired, resulting in anemia and reduced O₂ delivery to tissue.

THE IRON CYCLE IN HUMANS

Figure 126-1 outlines the major pathways of internal iron exchange in humans. Iron absorbed from the diet or released from stores circulates in the plasma bound to *transferrin*, the iron transport protein. Transferrin is a bilobed glycoprotein with two iron binding sites. Transferrin that carries iron exists in two forms—*monoferric* (one iron atom) or *diferric* (two iron atoms). The turnover (half-clearance time) of transferrin-bound iron is very rapid—typically 60–90 min. Because almost all of the iron transported by transferrin is delivered to the erythroid marrow, the clearance time of transferrin-bound iron from the circulation is affected most by the plasma iron level and the erythroid marrow activity. When erythropoiesis is markedly stimulated, the pool of erythroid cells requiring iron increases and the clearance time of iron from the circulation decreases. The half-clearance time of iron in

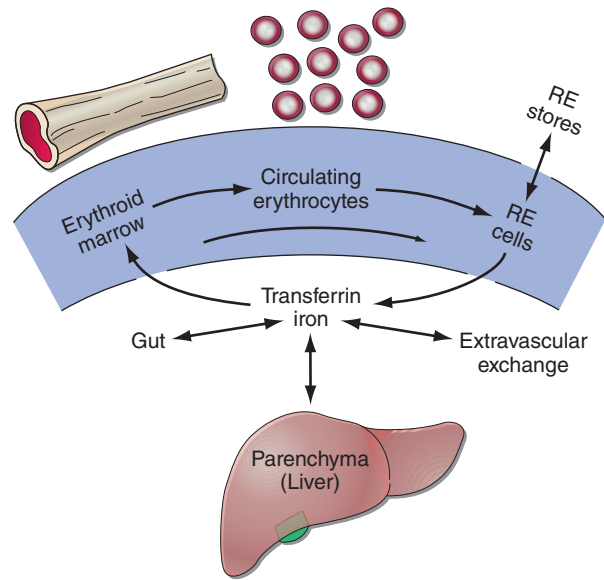


FIGURE 126-1 Internal iron exchange. Normally 80% of iron passing through the plasma transferrin pool is recycled from senescent red cells. Absorption of approximately 1 mg/d is required from the diet in men, and 1.4 mg/d in women to maintain homeostasis. As long as transferrin saturation is maintained between 20 and 60% and erythropoiesis is not increased, use of iron stores is not required. However, in the event of blood loss, dietary iron deficiency, or inadequate iron absorption, up to 40 mg/d of iron can be mobilized from stores. RE, reticuloendothelial.

the presence of iron deficiency is as short as 10–15 min. With suppression of erythropoiesis, the plasma iron level typically increases and the half-clearance time may be prolonged to several hours. Normally, the iron bound to transferrin turns over 6–8 times per day. Assuming a normal plasma iron level of 80–100 µg/dL, the amount of iron passing through the transferrin pool is 20–24 mg/d.

The iron-transferrin complex circulates in the plasma until it interacts with specific *transferrin receptors* on the surface of marrow erythroid cells. Diferric transferrin has the highest affinity for transferrin receptors; apotransferrin (not carrying iron) has very little affinity. Although transferrin receptors are found on cells in many tissues within the body—and all cells at some time during development will display transferrin receptors—the cell having the greatest number of receptors (300,000–400,000/cell) is the developing erythroblast.

Once the iron-bearing transferrin interacts with its receptor, the complex is internalized via clathrin-coated pits and transported to an acidic endosome, where the iron is released at the low pH. The iron is then made available for heme synthesis while the transferrin-receptor complex is recycled to the surface of the cell, where the bulk of the transferrin is released back into circulation and the transferrin receptor reanchors into the cell membrane. At this point a certain amount of the transferrin receptor protein may be released into circulation and can be measured as soluble transferrin receptor protein. Within the erythroid cell, iron in excess of the amount needed for hemoglobin synthesis binds to a storage protein, *apoferritin*, forming *ferritin*. This mechanism of iron exchange also takes place in other cells of the body expressing transferrin receptors, especially liver parenchymal cells where the iron can be incorporated into heme-containing enzymes or stored. The iron incorporated into hemoglobin subsequently enters the circulation as new red cells are released from the bone marrow. The iron is then part of the red cell mass and will not become available for reutilization until the red cell dies.

TABLE 126-1 BODY IRON DISTRIBUTION

	Iron Content, mg	
	Adult Male, 80 kg	Adult Female, 60 kg
Hemoglobin	2500	1700
Myoglobin/enzymes	500	300
Transferrin iron	3	3
Iron stores	600–1000	0–300

In a normal individual, the average red cell life span is 120 days. Thus, 0.8–1% of red cells are replaced each day. At the end of its life span, the red cell is recognized as senescent by the cells of the *reticulo-endothelial (RE) system*, and the red cell undergoes phagocytosis. Once within the RE cell, the ingested hemoglobin is broken down, the globin and other proteins are returned to the amino acid pool, and the iron is shuttled back to the surface of the RE cell, where it is presented to circulating transferrin. It is the efficient and highly conserved recycling of iron from senescent red cells that supports steady-state (and even mildly accelerated) erythropoiesis.

Because each milliliter of red cells contains 1 mg of elemental iron, the amount of iron needed to replace those red cells lost through senescence amounts to 20 mg/d (assuming an adult with a red cell mass of 2 L). Any additional iron required for daily red cell production comes from the diet. Normally, an adult male will need to absorb at least 1 mg of elemental iron daily to meet needs, while females in the child-bearing years will need to absorb an average of 1.4 mg/d. However, to achieve a maximum proliferative erythroid marrow response to anemia, additional iron must be available. With markedly stimulated erythropoiesis, demands for iron are increased by as much as six- to eightfold. With extravascular hemolytic anemia, the rate of red cell destruction is increased, but the iron recovered from the red cells is efficiently reutilized for hemoglobin synthesis. In contrast, with intravascular hemolysis or blood loss anemia, the rate of red cell production is limited by the amount of iron that can be mobilized from stores. Typically, the rate of mobilization under these circumstances will not support red cell production more than 2.5 times normal. If the delivery of iron to the stimulated marrow is suboptimal, the marrow's proliferative response is blunted, and hemoglobin synthesis is impaired. The result is a hypoproliferative marrow accompanied by microcytic, hypochromic anemia.

Whereas blood loss or hemolysis places a demand on the iron supply, inflammatory conditions interfere with iron release from stores and can result in a rapid decrease in the serum iron (see below).

NUTRITIONAL IRON BALANCE

The balance of iron in humans is tightly controlled and designed to conserve iron for reutilization. There is no regulated excretory pathway for iron, and the only mechanisms by which iron is lost are blood loss (via gastrointestinal bleeding, menses, or other forms of bleeding) and the loss of epithelial cells from the skin, gut, and genitourinary tract. Normally, the only route by which iron comes into the body is via absorption from food or from medicinal iron taken orally. Iron may also enter the body through red cell transfusions or injection of iron complexes. The margin between the amount of iron available for absorption and the requirement for iron in growing infants and the adult female is narrow; this accounts for the great prevalence of iron deficiency worldwide—currently estimated at one-half billion people.

The amount of iron required from the diet to replace losses averages approximately 10% of body iron content a year in men and 15% in women of childbearing age. Dietary iron content is closely related to total caloric intake (approximately 6 mg of elemental iron per 1000 calories). Iron bioavailability is affected by the nature of the foodstuff, with heme iron (e.g., red meat) being most readily absorbed. In the United States, the average iron intake in an adult male is 15 mg/d with 6% absorption; for the average female, the daily intake is 11 mg/d with 12% absorption. An individual with iron deficiency can increase iron absorption to approximately 20% of the iron present in a meat-containing diet but only 5–10% of the iron in a vegetarian diet. As a result, one-third of the female population in the United States has virtually no iron stores. Vegetarians are at an additional disadvantage because certain foodstuffs that include phytates and phosphates reduce iron absorption by approximately 50%. When ionizable iron salts are given together with food, the amount of iron absorbed is reduced. When the percentage of iron absorbed from individual food items is compared with the percentage for an equivalent amount of ferrous salt, iron in vegetables is only about one-twentieth as available, egg iron one-eighth, liver iron one-half, and heme iron one-half to two-thirds.

Infants, children, and adolescents may be unable to maintain normal iron balance because of the demands of body growth and lower dietary intake of iron. During the last two trimesters of pregnancy, daily iron requirements increase to 5–6 mg, and iron supplements are strongly recommended for pregnant women in developed countries.

Iron absorption takes place largely in the proximal small intestine and is a carefully regulated process. For absorption, iron must be taken up by the luminal cell. That process is facilitated by the acidic contents of the stomach, which maintains the iron in solution. At the brush border of the absorptive cell, the ferric iron is converted to the ferrous form by a ferrireductase. Transport across the membrane is accomplished by divalent metal transporter type 1 (DMT-1, also known as natural resistance macrophage-associated protein type 2 [Nramp 2] or DCT-1). DMT-1 is a general cation transporter. Once inside the gut cell, iron may be stored as ferritin or transported through the cell to be released at the basolateral surface to plasma transferrin through the membrane-embedded iron exporter, ferroportin. The function of ferroportin is negatively regulated by hepcidin, the principal iron regulatory hormone. In the process of release, iron interacts with another ferroxidase, hephaestin, which oxidizes the iron to the ferric form for transferrin binding. Hephaestin is similar to ceruloplasmin, the copper-carrying protein.

Iron absorption is influenced by a number of physiologic states. Erythroid hyperplasia stimulates iron absorption even in the face of normal or increased iron stores, and hepcidin levels are inappropriately low. Thus, patients with anemias associated with high levels of ineffective erythropoiesis absorb excess amounts of dietary iron. The molecular mechanism underlying this relationship is not known. Over time, this may lead to iron overload and tissue damage. In iron deficiency, hepcidin levels are also low and iron is much more efficiently absorbed; the contrary is true in states of secondary iron overload. The normal individual can reduce iron absorption in situations of excessive intake or medicinal iron intake; however, while the percentage of iron absorbed goes down, the absolute amount goes up. This accounts for the acute iron toxicity occasionally seen when children ingest large numbers of iron tablets. Under these circumstances, the amount of iron absorbed exceeds the transferrin binding capacity of the plasma, resulting in free iron that affects critical organs such as cardiac muscle cells.

IRON-DEFICIENCY ANEMIA



Iron deficiency is one of the most prevalent forms of malnutrition. Globally, 50% of anemia is attributable to iron deficiency and accounts for approximately 841,000 deaths annually worldwide. Africa and parts of Asia bear 71% of the global mortality burden; North America represents only 1.4% of the total morbidity and mortality associated with iron deficiency.

STAGES OF IRON DEFICIENCY

The progression to iron deficiency can be divided into three stages (Fig. 126-2). The first stage is *negative iron balance*, in which the demands for (or losses of) iron exceed the body's ability to absorb iron from the diet. This stage results from a number of physiologic mechanisms, including blood loss, pregnancy (in which the demands for red cell production by the fetus outstrip the mother's ability to provide iron), rapid growth spurts in the adolescent, or inadequate dietary iron intake. Blood loss in excess of 10–20 mL of red cells per day is greater than the amount of iron that the gut can absorb from a normal diet. Under these circumstances, the iron deficit must be made up by mobilization of iron from RE storage sites. During this period, iron stores—reflected by the serum ferritin level or the appearance of stainable iron on bone marrow aspirations—decrease. As long as iron stores are present and can be mobilized, the serum iron, total iron-binding capacity (TIBC), and red cell protoporphyrin levels remain within normal limits. At this stage, red cell morphology and indices are normal.

When iron stores become depleted, the serum iron begins to fall. Gradually, the TIBC increases, as do red cell protoporphyrin levels. By definition, marrow iron stores are absent when the serum ferritin level is <15 µg/L. As long as the serum iron remains within the normal

	Normal	Negative iron balance	Iron-deficient erythropoiesis	Iron-deficiency anemia
Iron stores				
Erythron iron				
Marrow iron stores	1-3+	0-1+	0	0
Serum ferritin (µg/L)	50-200	<20	<15	<15
TIBC (µg/dL)	300-360	>360	>380	>400
SI (µg/dL)	50-150	NL	<50	<30
Saturation (%)	30-50	NL	<20	<10
Marrow sideroblasts (%)	40-60	NL	<10	<10
RBC protoporphyrin (µg/dL)	30-50	NL	>100	>200
RBC morphology	NL	NL	NL	Microcytic/hypochromic

FIGURE 126-2 Laboratory studies in the evolution of iron deficiency. Measurements of marrow iron stores, serum ferritin, and total iron-binding capacity (TIBC) are sensitive to early iron-store depletion. Iron-deficient erythropoiesis is recognized from additional abnormalities in the serum iron (SI), percent transferrin saturation, the pattern of marrow sideroblasts, and the red blood cell (RBC) protoporphyrin level. Patients with iron-deficiency anemia demonstrate all the same abnormalities plus hypochromic microcytic anemia. (From RS Hillman, CA Finch: *The Red Cell Manual*, 7th ed. Philadelphia, F.A. Davis and Co., 1996, with permission.)

range, hemoglobin synthesis is unaffected despite the dwindling iron stores. Once the transferrin saturation falls to 15–20%, hemoglobin synthesis becomes impaired. This is a period of *iron-deficient erythropoiesis*. Careful evaluation of the peripheral blood smear reveals the first appearance of microcytic cells, and if the laboratory technology is available, one finds hypochromic reticulocytes in circulation. Gradually, the hemoglobin and hematocrit begin to fall, reflecting *iron-deficiency anemia*. The transferrin saturation at this point is 10–15%.

When moderate anemia is present (hemoglobin 10–13 g/dL), the bone marrow remains hypoproliferative. With more severe anemia (hemoglobin 7–8 g/dL), hypochromia and microcytosis become more prominent, target cells and misshapen red cells (poikilocytes) appear on the blood smear as cigar- or pencil-shaped forms, and the erythroid marrow becomes increasingly ineffective. Consequently, with severe prolonged iron-deficiency anemia, erythroid hyperplasia of the marrow develops, rather than hypoproliferation.

CAUSES OF IRON DEFICIENCY

Conditions that increase demand for iron, increase iron loss, or decrease iron intake or absorption can produce iron deficiency (Table 126-2).

CLINICAL PRESENTATION OF IRON DEFICIENCY

Certain clinical conditions carry an increased likelihood of iron deficiency. Pregnancy, adolescence, periods of rapid growth, and an intermittent history of blood loss of any kind should alert the clinician to possible iron deficiency. A cardinal rule is that the appearance of iron deficiency in an adult male means gastrointestinal blood loss until proven otherwise. Signs related to iron deficiency depend on the severity and chronicity of the anemia in addition to the usual signs of anemia—fatigue, pallor, and reduced exercise capacity. *Cheilosis* (fissures at the corners of the mouth) and *koilonychia* (spooning of the fingernails) are signs of advanced tissue iron deficiency. The diagnosis of iron deficiency is typically based on laboratory results.

TABLE 126-2 CAUSES OF IRON DEFICIENCY

Increased Demand for Iron
Rapid growth in infancy or adolescence
Pregnancy
Erythropoietin therapy
Increased Iron Loss
Chronic blood loss
Menses
Acute blood loss
Blood donation
Phlebotomy as treatment for polycythemia vera
Decreased Iron Intake or Absorption
Inadequate diet
Malabsorption from disease (sprue, Crohn's disease)
Malabsorption from surgery (gastrectomy and some forms of bariatric surgery)
Acute or chronic inflammation

LABORATORY IRON STUDIES

Serum Iron and Total Iron-Binding Capacity The serum iron level represents the amount of circulating iron bound to transferrin. The TIBC is an indirect measure of the circulating transferrin. The normal range for the serum iron is 50–150 µg/dL; the normal range for TIBC is 300–360 µg/dL. Transferrin saturation, which is normally 25–50%, is obtained by the following formula: $\text{serum iron} \times 100 \div \text{TIBC}$. Iron-deficiency states are associated with saturation levels below 20%. There is a diurnal variation in the serum iron. A transferrin saturation % >50% indicates that a disproportionate amount of the iron bound to transferrin is being delivered to nonerythroid tissues. If this persists for an extended time, tissue iron overload may occur.

Serum Ferritin Free iron is toxic to cells, and the body has established an elaborate set of protective mechanisms to bind iron in various tissue compartments. Within cells, iron is stored complexed to protein as ferritin or hemosiderin. Apoferritin binds to free ferrous iron and stores it in the ferric state. As ferritin accumulates within cells of the RE system, protein aggregates are formed as hemosiderin. Iron in ferritin or hemosiderin can be extracted for release by the RE cells, although hemosiderin is less readily available. Under steady-state conditions, the serum ferritin level correlates with total body iron stores; thus, the serum ferritin level is the most convenient laboratory test to estimate iron stores. The normal value for ferritin varies according to the age and gender of the individual (Fig. 126-3). Adult males have serum

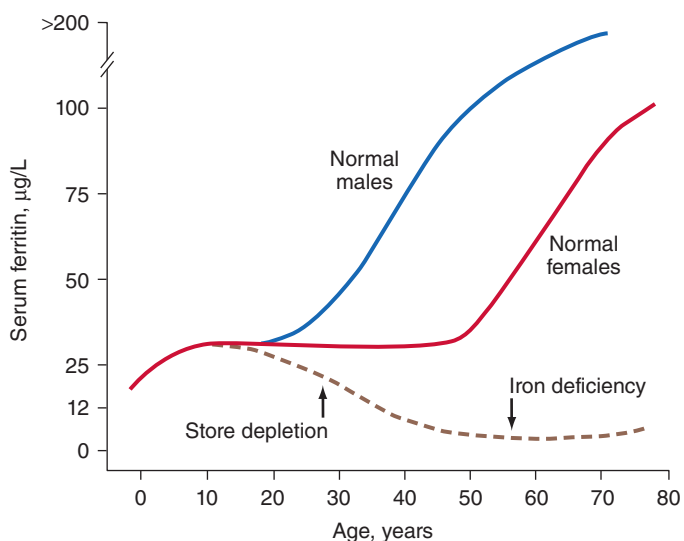


FIGURE 126-3 Serum ferritin levels as a function of sex and age. Iron store depletion and iron deficiency are accompanied by a decrease in serum ferritin level below 20 µg/L. (From RS Hillman et al: *Hematology in Clinical Practice*, 5th ed. New York, McGraw-Hill, 2011, with permission.)

TABLE 126-3 IRON STORE MEASUREMENTS

Iron Stores	Marrow Iron Stain, 0–4+	Serum Ferritin, µg/L
0	0	<15
1–300 mg	Trace to 1+	15–30
300–800 mg	2+	30–60
800–1000 mg	3+	60–150
1–2 g	4+	>150
Iron overload	—	>500–1000

ferritin values averaging 100 µg/L, while adult females have levels averaging 30 µg/L. As iron stores are depleted, the serum ferritin falls to <15 µg/L. Such levels are diagnostic of absent body iron stores.

Evaluation of Bone Marrow Iron Stores Although RE iron stores can be estimated from the iron stain of a bone marrow aspirate or biopsy, the measurement of serum ferritin has largely supplanted these procedures for determination of storage iron (Table 126-3). The serum ferritin level is a better indicator of iron overload than the marrow iron stain. However, in addition to storage iron, the marrow iron stain provides information about the effective delivery of iron to developing erythroblasts. Normally, when the marrow smear is stained for iron, 20–40% of developing erythroblasts—called *sideroblasts*—will have visible ferritin granules in their cytoplasm. This represents iron in excess of that needed for hemoglobin synthesis. In states in which release of iron from storage sites is blocked, RE iron will be detectable, and there will be few or no sideroblasts. In the myelodysplastic syndromes, mitochondrial dysfunction can occur, and accumulation of iron in mitochondria appears in a necklace fashion around the nucleus of the erythroblast. Such cells are referred to as *ringed sideroblasts*.

Red Cell Protoporphyrin Levels Protoporphyrin is an intermediate in the pathway to heme synthesis. Under conditions in which heme synthesis is impaired, protoporphyrin accumulates within the red cell. This reflects an inadequate iron supply to erythroid precursors to support hemoglobin synthesis. Normal values are <30 µg/dL of red cells. In iron deficiency, values in excess of 100 µg/dL are seen. The most common causes of increased red cell protoporphyrin levels are absolute or relative iron deficiency and lead poisoning.

Serum Levels of Transferrin Receptor Protein Because erythroid cells have the highest numbers of transferrin receptors of any cell in the body, and because transferrin receptor protein (TRP) is released by cells into the circulation, serum levels of TRP reflect the total erythroid marrow mass. Another condition in which TRP levels are elevated is absolute iron deficiency. Normal values are 4–9 µg/L determined by immunoassay. This laboratory test is becoming increasingly available and, along with the serum ferritin, has been proposed to distinguish between iron deficiency and the anemia of inflammation (see below).

DIFFERENTIAL DIAGNOSIS

Other than iron deficiency, only three conditions need to be considered in the differential diagnosis of a hypochromic microcytic anemia (Table 126-4). The first is an inherited defect in globin chain synthesis:

the thalassemias. These are differentiated from iron deficiency most readily by serum iron values; normal or increased serum iron levels and transferrin saturation are characteristic of the thalassemias. In addition, the red blood cell distribution width (RDW) index is generally normal in thalassemia and elevated in iron deficiency.

The second condition is the anemia of inflammation (AI; also referred to as the anemia of chronic disease) with inadequate iron supply to the erythroid marrow. The distinction between true iron-deficiency anemia and AI is among the most common diagnostic problems encountered by clinicians (see below). Usually, AI is normocytic and normochromic. The iron values usually make the differential diagnosis clear, as the ferritin level is normal or increased and the percent transferrin saturation and TIBC are typically below normal.

Finally, the myelodysplastic syndromes represent the third and least common condition. Occasionally, patients with myelodysplasia have impaired hemoglobin synthesis with mitochondrial dysfunction, resulting in impaired iron incorporation into heme. The iron values again reveal normal stores and more than an adequate supply to the marrow, despite the microcytosis and hypochromia.

TREATMENT IRON-DEFICIENCY ANEMIA

The severity and cause of iron-deficiency anemia will determine the appropriate approach to treatment. As an example, symptomatic elderly patients with severe iron-deficiency anemia and cardiovascular instability may require red cell transfusions. Younger individuals who have compensated for their anemia can be treated more conservatively with iron replacement. The foremost issue for the latter patient is the precise identification of the cause of the iron deficiency.

For the majority of cases of iron deficiency (pregnant women, growing children and adolescents, patients with infrequent episodes of bleeding, and those with inadequate dietary intake of iron), oral iron therapy will suffice. For patients with unusual blood loss or malabsorption, specific diagnostic tests and appropriate therapy take priority. Once the diagnosis of iron-deficiency anemia and its cause is made, there are three major therapeutic approaches.

RED CELL TRANSFUSION

Transfusion therapy is reserved for individuals who have symptoms of anemia, cardiovascular instability, and continued and excessive blood loss from whatever source and who require immediate intervention. The management of these patients is less related to the iron deficiency than it is to the consequences of the severe anemia. Not only do transfusions correct the anemia acutely, but the transfused red cells provide a source of iron for reutilization, assuming they are not lost through continued bleeding. Transfusion therapy will stabilize the patient while other options are reviewed.

ORAL IRON THERAPY

In the asymptomatic patient with established iron-deficiency anemia, treatment with oral iron is usually adequate. Multiple preparations are available, ranging from simple iron salts to complex iron compounds designed for sustained release throughout the small intestine (Table 126-5). Although the various preparations

TABLE 126-4 DIAGNOSIS OF MICROCYTIC ANEMIA

Tests	Iron Deficiency	Inflammation	Thalassemia	Sideroblastic Anemia
Smear	Micro/hypo	Normal micro/hypo	Micro/hypo with targeting	Variable
Serum iron (µg/dL)	<30	<50	Normal to high	Normal to high
TIBC (µg/dL)	>360	<300	Normal	Normal
Percent saturation	<10	10–20	30–80	30–80
Ferritin (µg/L)	<15	30–200	50–300	50–300
Hemoglobin pattern on electrophoresis	Normal	Normal	Abnormal with β thalassemia; can be normal with α thalassemia	Normal

Abbreviation: TIBC, total iron-binding capacity.

TABLE 126-5 ORAL IRON PREPARATIONS

Generic Name	Tablet (Iron Content), mg	Elixir (Iron Content), mg in 5 mL
Ferrous sulfate	325 (65)	300 (60)
	195 (39)	90 (18)
Extended release	525 (105)	
Ferrous fumarate	325 (107)	
	195 (64)	100 (33)
Ferrous gluconate	325 (39)	300 (35)
Polysaccharide iron	150 (150)	100 (100)
	50 (50)	

contain different amounts of iron, they are generally all absorbed well and are effective in treatment. Some come with other compounds designed to enhance iron absorption, such as ascorbic acid. It is not clear whether the benefits of such compounds justify their costs. Typically, for iron replacement therapy, up to 200 mg of elemental iron per day is given, usually as three or four iron tablets (each containing 50–65 mg elemental iron) given over the course of the day. Ideally, oral iron preparations should be taken on an empty stomach, since food may inhibit iron absorption. Some patients with gastric disease or prior gastric surgery require special treatment with iron solutions, because the retention capacity of the stomach may be reduced. The retention capacity is necessary for dissolving the shell of the iron tablet before the release of iron. A dose of 200 mg of elemental iron per day should result in the absorption of iron up to 50 mg/d. This supports a red cell production level of two to three times normal in an individual with a normally functioning marrow and appropriate erythropoietin stimulus. However, as the hemoglobin level rises, erythropoietin stimulation decreases, and the amount of iron absorbed is reduced. The goal of therapy in individuals with iron-deficiency anemia is not only to repair the anemia, but also to provide stores of at least 0.5–1 g of iron. Sustained treatment for a period of 6–12 months after correction of the anemia will be necessary to achieve this.

Of the complications of oral iron therapy, gastrointestinal distress is the most prominent and is seen in 15–20% of patients. Abdominal pain, nausea, vomiting, or constipation may lead to noncompliance. Although small doses of iron or iron preparations with delayed release may help somewhat, the gastrointestinal side effects are a major impediment to the effective treatment of a number of patients.

The response to iron therapy varies, depending on the erythropoietin stimulus and the rate of absorption. Typically, the reticulocyte count should begin to increase within 4–7 days after initiation of therapy and peak at 1–1½ weeks. The absence of a response may be due to poor absorption, noncompliance (which is common), or a confounding diagnosis. A useful test in the clinic to determine the patient's ability to absorb iron is the *iron tolerance test*. Two iron tablets are given to the patient on an empty stomach, and the serum iron is measured serially over the subsequent 2 h. Normal absorption will result in an increase in the serum iron of at least 100 µg/dL. If iron deficiency persists despite adequate treatment, it may be necessary to switch to parenteral iron therapy.

PARENTERAL IRON THERAPY

Intravenous iron can be given to patients who are unable to tolerate oral iron; whose needs are relatively acute; or who need iron on an ongoing basis, usually due to persistent gastrointestinal blood loss. Parenteral iron use has been increasing rapidly in the last several years with the recognition that recombinant erythropoietin (EPO) therapy induces a large demand for iron—a demand that frequently cannot be met through the physiologic release of iron from RE sources or oral iron absorption. The safety of parenteral iron—particularly iron dextran—has been a concern. The serious adverse reaction rate to intravenous high-molecular-weight iron dextran is 0.7%. Fortunately, newer iron complexes are available in the United States, such as ferumoxytol (Feraheme), sodium ferric

gluconate (Ferrlecit), iron sucrose (Venofer), and ferric carboxymaltose (Injectafer), that have much lower rates of adverse effects. Ferumoxytol delivers 510 mg of iron per injection; ferric gluconate 125 mg per injection, ferric carboxymaltose 750 mg per injection, and iron sucrose 200 mg per injection.

Parenteral iron is used in two ways: one is to administer the total dose of iron required to correct the hemoglobin deficit and provide the patient with at least 500 mg of iron stores; the second is to give repeated small doses of parenteral iron over a protracted period. The latter approach is common in dialysis centers, where it is not unusual for 100 mg of elemental iron to be given weekly for 10 weeks to augment the response to recombinant EPO therapy. The amount of iron needed by an individual patient is calculated by the following formula:

$$\text{Body weight (kg)} \times 2.3 \times (15 - \text{patient's hemoglobin, g/dL}) + 500 \text{ or } 1000 \text{ mg (for stores).}$$

In administering intravenous iron dextran, anaphylaxis is a concern. Anaphylaxis is much rarer with the newer preparations. The factors that have correlated with an anaphylactic-like reaction include a history of multiple allergies or a prior allergic reaction to dextran (in the case of iron dextran). Generalized symptoms appearing several days after the infusion of a large dose of iron can include arthralgias, skin rash, and low-grade fever. These may be dose-related, but they do not preclude the further use of parenteral iron in the patient. To date, patients with sensitivity to iron dextran have been safely treated with other parenteral iron preparations. If a large dose of iron dextran is to be given (>100 mg), the iron preparation should be diluted in 5% dextrose in water or 0.9% NaCl solution. The iron solution can then be infused over a 60- to 90-min period (for larger doses) or at a rate convenient for the attending nurse or physician. Although a test dose (25 mg) of parenteral iron dextran is recommended, in reality a slow infusion of a larger dose of parenteral iron solution will afford the same kind of early warning as a separately injected test dose. Early in the infusion of iron, if chest pain, wheezing, a fall in blood pressure, or other systemic symptoms occur, the infusion of iron should be stopped immediately.

OTHER HYPOPROLIFERATIVE ANEMIAS

In addition to mild to moderate iron-deficiency anemia, the hypoproliferative anemias can be divided into four categories: (1) chronic inflammation, (2) renal disease, (3) endocrine and nutritional deficiencies (hypometabolic states), and (4) marrow damage (Chap. 130). With chronic inflammation, renal disease, or hypometabolism, endogenous EPO production is inadequate for the degree of anemia observed. For the anemia of chronic inflammation, the erythroid marrow also responds inadequately to stimulation, due in part to defective *iron reutilization*. As a result of the lack of adequate EPO stimulation, an examination of the peripheral blood smear will disclose only an occasional polychromatophilic (“shift”) reticulocyte. In cases of iron deficiency or marrow damage, appropriate elevations in endogenous EPO levels are typically found, and shift reticulocytes will be present on the blood smear.

ANEMIA OF ACUTE AND CHRONIC INFLAMMATION/INFECTION (AI)

AI—which encompasses inflammation, infection, tissue injury, and conditions (such as cancer) associated with the release of proinflammatory cytokines—is one of the most common forms of anemia seen clinically. It is the most important anemia in the differential diagnosis of iron deficiency, because many of the features of the anemia are brought about by inadequate iron delivery to the marrow, despite the presence of normal or increased iron stores. This is reflected by a low serum iron, increased red cell protoporphyrin, a hypoproliferative marrow, transferrin saturation in the range of 15–20%, and a normal or increased serum ferritin. The serum ferritin values are often the most distinguishing features between true iron-deficiency anemia and the iron-restricted erythropoiesis associated with inflammation. Typically, serum ferritin values increase threefold over basal levels

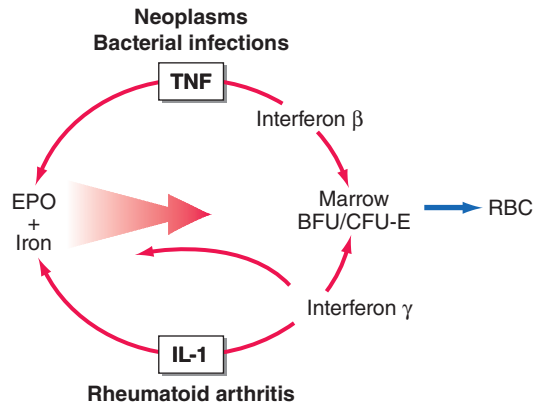


FIGURE 126-4 Suppression of erythropoiesis by inflammatory cytokines. Through the release of tumor necrosis factor (TNF) and interferon γ (IFN- γ), neoplasms and bacterial infections suppress erythropoietin (EPO) production and the proliferation of erythroid progenitors (erythroid burst-forming units and erythroid colony-forming units [BFU/CFU-E]). The mediators in patients with vasculitis and rheumatoid arthritis include interleukin 1 (IL-1) and IFN- γ . The red arrows indicate sites of inflammatory cytokine inhibitory effects. RBC, red blood cell.

in the face of inflammation. These changes are due to the effects of inflammatory cytokines and hepcidin, the key iron regulatory hormone, acting at several levels of erythropoiesis (Fig. 126-4).

Interleukin 1 (IL-1) directly decreases EPO production in response to anemia. IL-1, acting through accessory cell release of interferon γ (IFN- γ), suppresses the response of the erythroid marrow to EPO—an effect that can be overcome by EPO administration in vitro and in vivo. In addition, tumor necrosis factor (TNF), acting through the release of IFN- γ by marrow stromal cells, also suppresses the response to EPO. Hepcidin, made by the liver, is increased in inflammation via an IL-6 mediated pathway, and acts to suppress iron absorption and iron release from storage sites. The overall result is a chronic hypoproliferative anemia with classic changes in iron metabolism. The anemia is further compounded by a mild to moderate shortening in red cell survival.

With chronic inflammation, the primary disease will determine the severity and characteristics of the anemia. For example, many patients with cancer also have anemia that is typically normocytic and normochromic. In contrast, patients with long-standing active rheumatoid arthritis or chronic infections such as tuberculosis will have a microcytic, hypochromic anemia. In both cases, the bone marrow is hypoproliferative, but the differences in red cell indices reflect differences in the availability of iron for hemoglobin synthesis. Occasionally, conditions associated with chronic inflammation are also associated with chronic blood loss. Under these circumstances, a bone marrow aspirate stained for iron may be necessary to rule out absolute iron deficiency. However, the administration of iron in this case will correct the iron deficiency component of the anemia and leave the inflammatory component unaffected.

The anemia associated with acute infection or inflammation is typically mild but becomes more pronounced over time. Acute infection can produce a decrease in hemoglobin levels of 2–3 g/dL within 1 or 2 days; this is largely related to the hemolysis of red cells near the end of their natural life span. The fever and cytokines released exert a selective pressure against cells with more limited capacity to maintain the red cell membrane. In most individuals, the mild anemia is reasonably well tolerated, and symptoms, if present, are associated with the underlying disease. Occasionally, in patients with preexisting cardiac disease, moderate anemia (hemoglobin 10–11 g/dL) may be associated

with angina, exercise intolerance, and shortness of breath. The erythropoietic profile that distinguishes the anemia of inflammation from the other causes of hypoproliferative anemias is shown in Table 126-6.

ANEMIA OF CHRONIC KIDNEY DISEASE (CKD)

Progressive CKD is usually associated with a moderate to severe hypoproliferative anemia; the level of the anemia correlates with the stage of CKD. Red cells are typically normocytic and normochromic, and reticulocytes are decreased. The anemia is primarily due to a failure of EPO production by the diseased kidney and a reduction in red cell survival. In certain forms of acute renal failure, the correlation between the anemia and renal function is weaker. Patients with the hemolytic-uremic syndrome increase erythropoiesis in response to the hemolysis, despite renal failure requiring dialysis. Polycystic kidney disease also shows a smaller degree of EPO deficiency for a given level of renal failure. By contrast, patients with diabetes or myeloma have more severe EPO deficiency for a given level of renal failure.

Assessment of iron status provides information to distinguish the anemia of CKD from the other forms of hypoproliferative anemia (Table 126-6) and to guide management. Patients with the anemia of CKD usually present with normal serum iron, TIBC, and ferritin levels. However, those maintained on chronic hemodialysis may develop iron deficiency from blood loss through the dialysis procedure. Iron must be replenished in these patients to ensure an adequate response to EPO therapy (see below).

ANEMIA IN HYPOMETABOLIC STATES

Patients who are starving, particularly for protein, and those with a variety of endocrine disorders that produce lower metabolic rates, may develop a mild to moderate hypoproliferative anemia. The release of EPO from the kidney is sensitive to the need for O_2 , not just O_2 levels. Thus, EPO production is triggered at lower levels of blood O_2 content in disease states (such as hypothyroidism and starvation) where metabolic activity, and thus O_2 demand, is decreased.

Endocrine Deficiency States The difference in the levels of hemoglobin between men and women is related to the effects of androgen and estrogen on erythropoiesis. Testosterone and anabolic steroids augment erythropoiesis; castration and estrogen administration to males decrease erythropoiesis. Patients who are hypothyroid or have deficits in pituitary hormones also may develop a mild anemia. Pathogenesis may be complicated by other nutritional deficiencies because iron and folic acid absorption can be affected by these disorders. Usually, correction of the hormone deficiency reverses the anemia.

Anemia may be more severe in Addison's disease, depending on the level of thyroid and androgen hormone dysfunction; however, anemia may be masked by decreases in plasma volume. Once such patients are given cortisol and volume replacement, the hemoglobin level may fall rapidly. Mild anemia complicating hyperparathyroidism may be due to decreased EPO production as a consequence of the renal effects of hypercalcemia or to impaired proliferation of erythroid progenitors.

Protein Starvation Decreased dietary intake of protein may lead to mild to moderate hypoproliferative anemia; this form of anemia may be prevalent in the elderly. The anemia can be more severe in patients with a greater degree of starvation. In marasmus, where patients are

TABLE 126-6 DIAGNOSIS OF HYPOPROLIFERATIVE ANEMIAS

Tests	Iron Deficiency	Inflammation	Renal Disease	Hypometabolic States
Anemia	Mild to severe	Mild	Mild to severe	Mild
MCV (fL)	60–90	80–90	90	90
Morphology	Normo-microcytic	Normocytic	Normocytic	Normocytic
SI (μ g/dL)	<30	<50	Normal	Normal
TIBC (μ g/dL)	>360	<300	Normal	Normal
Saturation (%)	<10	10–20	Normal	Normal
Serum ferritin (μ g/L)	<15	30–200	115–150	Normal
Iron stores	0	2–4+	1–4+	Normal

Abbreviations: MCV, mean corpuscular volume; SI, serum iron; TIBC, total iron-binding capacity.

127 Disorders of Hemoglobin

Edward J. Benz, Jr.

both protein and calorie deficient, the release of EPO is impaired in proportion to the reduction in metabolic rate; however, the degree of anemia may be masked by volume depletion and becomes apparent after refeeding. Deficiencies in other nutrients (iron, folate) may also complicate the clinical picture but may not be apparent at diagnosis. Changes in the erythrocyte indices on refeeding should prompt evaluation of iron, folate, and B₁₂ status.

Anemia in Liver Disease A mild hypoproliferative anemia may develop in patients with chronic liver disease from nearly any cause. The peripheral blood smear may show spur cells and stomatocytes from the accumulation of excess cholesterol in the membrane from a deficiency of lecithin-cholesterol acyltransferase. Red cell survival is shortened, and the production of EPO is inadequate to compensate. In alcoholic liver disease, nutritional deficiencies are common and complicate the management. Folate deficiency from inadequate intake, as well as iron deficiency from blood loss and inadequate intake, can alter the red cell indices.

TREATMENT HYPOPROLIFERATIVE ANEMIAS

Many patients with hypoproliferative anemias experience recovery of normal hemoglobin levels when the underlying disease is appropriately treated. For those in whom such reversals are not possible—such as patients with end-stage kidney disease, cancer, and chronic inflammatory diseases—symptomatic anemia requires treatment. The two major forms of treatment are transfusions and EPO.

TRANSFUSIONS

Thresholds for transfusion should be determined based on the patient's symptoms. In general, patients without serious underlying cardiovascular or pulmonary disease can tolerate hemoglobin levels above 7–8 g/dL and do not require intervention until the hemoglobin falls below that level. Patients with more physiologic compromise may need to have their hemoglobin levels kept above 11 g/dL. Usually, a unit of packed red cells increases the hemoglobin level by 1 g/dL. Transfusions are associated with certain infectious risks (Chap. 138e), and chronic transfusions can produce iron overload. Importantly, the liberal use of blood has been associated with increased morbidity and mortality, particularly in the intensive care setting. Therefore, in the absence of documented tissue hypoxia, a conservative approach to the use of red cell transfusions is preferable.

ERYTHROPOIETIN (EPO)

EPO is particularly useful in anemias in which endogenous EPO levels are inappropriately low, such as CKD or AI. Iron status must be evaluated and iron replaced to obtain optimal effects from EPO. In patients with CKD, the usual dose of EPO is 50–150 U/kg three times a week intravenously. Hemoglobin levels of 10–12 g/dL are usually reached within 4–6 weeks if iron levels are adequate; 90% of these patients respond. Once a target hemoglobin level is achieved, the EPO dose can be decreased. A decrease in hemoglobin level occurring in the face of EPO therapy usually signifies the development of an infection or iron depletion. Aluminum toxicity and hyperparathyroidism can also compromise the response to EPO. When an infection intervenes, it is best to interrupt the EPO therapy and rely on transfusions to correct the anemia until the infection is adequately treated. The dose of EPO needed to correct chemotherapy-induced anemia in patients with cancer is higher, up to 300 U/kg three times a week, and only approximately 60% of patients respond. Because of evidence that there is an increased risk of thromboembolic complications and tumor progression with EPO administration, the risks and benefits of using EPO in such patients must be weighed carefully, and the target hemoglobin should be that necessary to avoid transfusions.

Longer-acting preparations of EPO can reduce the frequency of injections. Darbepoetin alfa, a molecularly modified EPO with additional carbohydrate, has a half-life in the circulation that is three to four times longer than recombinant human EPO, permitting weekly or every other week dosing.

Hemoglobin is critical for normal oxygen delivery to tissues; it is also present in erythrocytes in such high concentrations that it can alter red cell shape, deformability, and viscosity. Hemoglobinopathies are disorders affecting the structure, function, or production of hemoglobin. These conditions are usually inherited and range in severity from asymptomatic laboratory abnormalities to death in utero. Different forms may present as hemolytic anemia, erythrocytosis, cyanosis, or vasoocclusive stigmata.

PROPERTIES OF THE HUMAN HEMOGLOBINS

HEMOGLOBIN STRUCTURE

Different hemoglobins are produced during embryonic, fetal, and adult life (Fig. 127-1). Each consists of a tetramer of globin polypeptide chains: a pair of α -like chains 141 amino acids long and a pair of β -like chains 146 amino acids long. The major adult hemoglobin, HbA, has the structure $\alpha_2\beta_2$. HbF ($\alpha_2\gamma_2$) predominates during most of gestation, and HbA₂ ($\alpha_2\delta_2$) is minor adult hemoglobin. Embryonic hemoglobins need not be considered here.

Each globin chain enfolds a single heme moiety, consisting of a protoporphyrin IX ring complexed with a single iron atom in the ferrous state (Fe²⁺). Each heme moiety can bind a single oxygen molecule; a molecule of hemoglobin can transport up to four oxygen molecules.

The amino acid sequences of the various globins are highly homologous to one another. Each has a highly helical *secondary structure*. Their globular *tertiary structures* cause the exterior surfaces to be rich in polar (hydrophilic) amino acids that enhance solubility, and the interior to be lined with nonpolar groups, forming a hydrophobic pocket into which heme is inserted. The tetrameric *quaternary structure* of HbA contains two $\alpha\beta$ dimers. Numerous tight interactions (i.e., $\alpha_1\beta_1$ contacts) hold the α and β chains together. The complete tetramer is held together by interfaces (i.e., $\alpha_1\beta_2$ contacts) between the α -like chain of one dimer and the non- α chain of the other dimer.

The hemoglobin tetramer is highly soluble, but individual globin chains are insoluble. Unpaired globin precipitates, forming inclusions that damage the cell and can trigger apoptosis. Normal globin chain synthesis is balanced so that each newly synthesized α or non- α globin chain will have an available partner with which to pair.

Solubility and reversible oxygen binding are the key properties deranged in hemoglobinopathies. Both depend most on the hydrophilic surface amino acids, the hydrophobic amino acids lining the heme pocket, a key histidine in the F helix, and the amino acids forming the $\alpha_1\beta_1$ and $\alpha_1\beta_2$ contact points. Mutations in these strategic regions tend to be the ones that alter oxygen affinity or solubility.

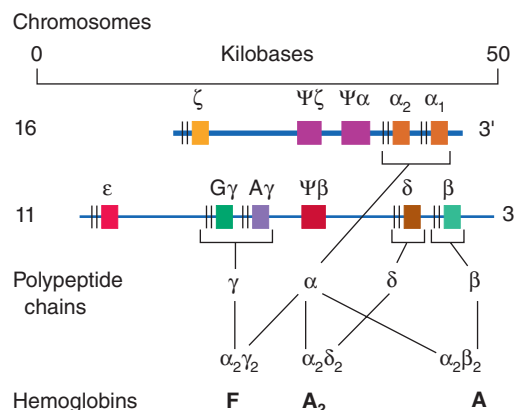


FIGURE 127-1 The globin genes. The α -like genes (α , ζ) are encoded on chromosome 16; the β -like genes (β , γ , δ , ϵ) are encoded on chromosome 11. The ζ and ϵ genes encode embryonic globins.

To support oxygen transport, hemoglobin must bind O_2 efficiently at the partial pressure of oxygen (PO_2) of the alveolus, retain it in the circulation, and release it to tissues at the PO_2 of tissue capillary beds. Oxygen acquisition and delivery over a relatively narrow range of oxygen tensions depend on a property inherent in the tetrameric arrangement of heme and globin subunits within the hemoglobin molecule called *cooperativity* or *heme-heme interaction*.

At low oxygen tensions, the hemoglobin tetramer is fully deoxygenated (Fig. 127-2). Oxygen binding begins slowly as O_2 tension rises. However, as soon as some oxygen has been bound by the tetramer, an abrupt increase occurs in the slope of the curve. Thus, hemoglobin molecules that have bound some oxygen develop a higher oxygen affinity, greatly accelerating their ability to combine with more oxygen. This S-shaped oxygen equilibrium curve (Fig. 127-2), along which substantial amounts of oxygen loading and unloading can occur over a narrow range of oxygen tensions, is physiologically more useful than the high-affinity hyperbolic curve of individual monomers.

Oxygen affinity is modulated by several factors. The Bohr effect is the ability of hemoglobin to deliver more oxygen to tissues at low pH. It arises from the stabilizing action of protons on deoxyhemoglobin, which binds protons more readily than oxyhemoglobin because the latter is a weaker acid (Fig. 127-2). Thus, hemoglobin has a lower oxygen affinity at low pH. The major small molecule that alters oxygen affinity in humans is 2,3-bisphosphoglycerate (2,3-BPG; formerly 2,3-DPG), which lowers oxygen affinity when bound to hemoglobin. HbA has a reasonably high affinity for 2,3-BPG. HbF does not bind 2,3-BPG, so it tends to have a higher oxygen affinity in vivo. Hemoglobin also binds nitric oxide reversibly; this interaction influences vascular tone, but its clinical relevance remains incompletely understood.

Proper oxygen transport depends on the tetrameric structure of the proteins, the proper arrangement of hydrophilic and hydrophobic amino acids, and interaction with protons or 2,3-BPG.

DEVELOPMENTAL BIOLOGY OF HUMAN HEMOGLOBINS

Red cells first appearing at about 6 weeks after conception contain the embryonic hemoglobins Hb Portland ($\zeta_2\gamma_2$), Hb Gower I ($\zeta_2\varepsilon_2$), and Hb Gower II ($\alpha_2\varepsilon_2$). At 10–11 weeks, fetal hemoglobin (HbF; $\alpha_2\gamma_2$) becomes predominant. The switch to nearly exclusive synthesis of adult hemoglobin (HbA; $\alpha_2\beta_2$) occurs at about 38 weeks (Fig. 127-1).

Fetuses and newborns therefore require α -globin but not β -globin for normal gestation. A major advance in understanding the HbF to HbA transition has been the demonstration that the transcription factor Bcl11a plays a pivotal role in its regulation. Small amounts of HbF are produced during postnatal life. A few red cell clones called *F cells* are progeny of a small pool of immature committed erythroid precursors (BFU-e) that retain the ability to produce HbF. Profound erythroid stresses, such as severe hemolytic anemias, bone marrow transplantation, or cancer chemotherapy, cause more of the F-potent BFU-e to be recruited. HbF levels thus tend to rise in some patients with sickle cell anemia or thalassemia. This phenomenon probably explains the ability of hydroxyurea to increase levels of HbF in adults. Agents such as butyrate and histone deacetylase inhibitors can also activate fetal globin genes partially after birth.

GENETICS AND BIOSYNTHESIS OF HUMAN HEMOGLOBIN

The human hemoglobins are encoded in two tightly linked gene clusters; the α -like globin genes are clustered on chromosome 16 and the β -like genes on chromosome 11 (Fig. 127-1). The α -like cluster consists of two α -globin genes and a single copy of the ζ gene. The non- α gene cluster consists of a single ε gene, the γ and δ fetal globin genes, and the adult β gene.

Important regulatory sequences flank each gene. Immediately upstream are typical promoter elements needed for the assembly of the transcription initiation complex. Sequences in the 5' flanking region of the γ and the β genes appear to be crucial for the correct developmental regulation of these genes, whereas elements that function like classic enhancers and silencers are in the 3' flanking regions. The locus control region (LCR) elements located far upstream appear to control the overall level of expression of each cluster. These elements achieve their regulatory effects by interacting with *trans*-acting transcription factors. Some of these factors are ubiquitous (e.g., Sp1 and YY1), while others are more or less limited to erythroid cells or hematopoietic cells (e.g., GATA-1, NFE-2, and EKLF). The LCR controlling the α -globin gene cluster is modulated by a SWI/SNF-like protein called ATRX; this protein appears to influence chromatin remodeling and DNA methylation. The association of α thalassemia with mental retardation and myelodysplasia in some families appears to be related to mutations in the ATRX pathway. This pathway also modulates genes specifically expressed during erythropoiesis, such

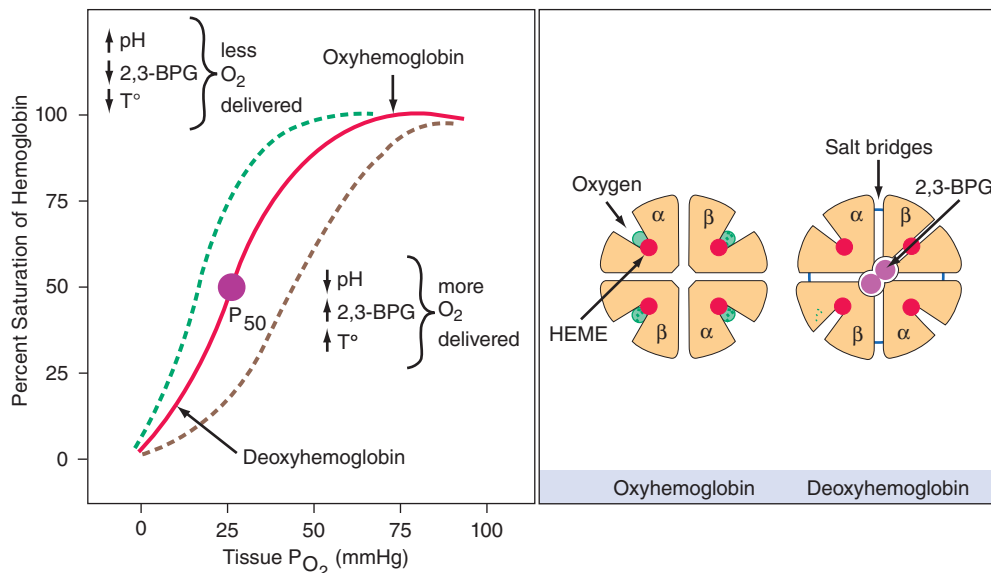


FIGURE 127-2 Hemoglobin-oxygen dissociation curve. The hemoglobin tetramer can bind up to four molecules of oxygen in the iron-containing sites of the heme molecules. As oxygen is bound, 2,3-bisphosphoglycerate (2,3-BPG) and carbon dioxide (CO_2) are expelled. Salt bridges are broken, and each of the globin molecules changes its conformation to facilitate oxygen binding. Oxygen release to the tissues is the reverse process, with salt bridges being formed and 2,3-BPG and CO_2 bound. Deoxyhemoglobin does not bind oxygen efficiently until the cell returns to conditions of higher pH, the most important modulator of O_2 affinity (Bohr effect). When acid is produced in the tissues, the dissociation curve shifts to the right, facilitating oxygen release and CO_2 binding. Alkalosis has the opposite effect, reducing oxygen delivery.

as those that encode the enzymes for heme biosynthesis. Normal red blood cell (RBC) differentiation requires the coordinated expression of the globin genes with the genes responsible for heme and iron metabolism. RBC precursors contain a protein, α -hemoglobin-stabilizing protein (AHSP), that enhances the folding and solubility of a globin, which is otherwise easily denatured, leading to insoluble precipitates. These precipitates play an important role in the thalassemia syndromes and certain unstable hemoglobin disorders. Polymorphic variation in the amounts and/or functional capacity of AHSP might explain some of the clinical variability seen in patients inheriting identical thalassemia mutations.

CLASSIFICATION OF HEMOGLOBINOPATHIES

There are five major classes of hemoglobinopathies (Table 127-1). *Structural hemoglobinopathies* occur when mutations alter the amino acid sequence of a globin chain, altering the physiologic properties of the variant hemoglobins and producing the characteristic clinical abnormalities. The most clinically relevant variant hemoglobins polymerize abnormally, as in sickle cell anemia, or exhibit altered solubility or oxygen-binding affinity. *Thalassemia syndromes* arise from mutations that impair production or translation of globin mRNA, leading to deficient globin chain biosynthesis. Clinical abnormalities are attributable to the inadequate supply of hemoglobin and the imbalances in the production of individual globin chains, leading to premature destruction of erythroblasts and RBC. *Thalassemic hemoglobin variants* combine features of thalassemia (e.g., abnormal globin biosynthesis) and of structural hemoglobinopathies (e.g., an abnormal amino acid sequence). *Hereditary persistence of fetal hemoglobin* (HPFH) is characterized by synthesis of high levels of fetal hemoglobin in adult life. *Acquired hemoglobinopathies* include modifications of the hemoglobin molecule by toxins (e.g., acquired methemoglobinemia) and clonal abnormalities of hemoglobin synthesis (e.g., high levels of HbF production in preleukemia and α thalassemia in myeloproliferative disorders).

TABLE 127-1 CLASSIFICATION OF HEMOGLOBINOPATHIES

- I. Structural hemoglobinopathies—hemoglobins with altered amino acid sequences that result in deranged function or altered physical or chemical properties
 - A. Abnormal hemoglobin polymerization—HbS, hemoglobin sickling
 - B. Altered O₂ affinity
 1. High affinity—polycythemia
 2. Low affinity—cyanosis, pseudoanemia
 - C. Hemoglobins that oxidize readily
 1. Unstable hemoglobins—hemolytic anemia, jaundice
 2. M hemoglobins—methemoglobinemia, cyanosis
- II. Thalassemias—defective biosynthesis of globin chains
 - A. α Thalassemias
 - B. β Thalassemias
 - C. $\delta\beta$, $\gamma\delta\beta$, $\alpha\beta$ Thalassemias
- III. Thalassemic hemoglobin variants—structurally abnormal Hb associated with coinherited thalassemic phenotype
 - A. HbE
 - B. Hb Constant Spring
 - C. Hb Lepore
- IV. Hereditary persistence of fetal hemoglobin—persistence of high levels of HbF into adult life
- V. Acquired hemoglobinopathies
 - A. Methemoglobin due to toxic exposures
 - B. Sulfhemoglobin due to toxic exposures
 - C. Carboxyhemoglobin
 - D. HbH in erythroleukemia
 - E. Elevated HbF in states of erythroid stress and bone marrow dysplasia

EPIDEMIOLOGY



Hemoglobinopathies are especially common in areas in which malaria is endemic. This clustering of hemoglobinopathies is assumed to reflect a selective survival advantage for the abnormal RBC, which presumably provides a less hospitable environment during the obligate RBC stages of the parasitic life cycle. Very young children with a thalassemia are *more* susceptible to infection with the nonlethal *Plasmodium vivax*. Thalassemia might then favor a natural protection against infection with the more lethal *Plasmodium falciparum*.

Thalassemias are the most common genetic disorders in the world, affecting nearly 200 million people worldwide. About 15% of African Americans are silent carriers for a thalassemia; a thalassemia trait (minor) occurs in 3% of African American and in 1–15% of persons of Mediterranean origin. β Thalassemia has a 10–15% incidence in individuals from the Mediterranean and Southeast Asia and 0.8% in African Americans. The number of severe cases of thalassemia in the United States is about 1000. Sickle cell disease is the most common structural hemoglobinopathy, occurring in heterozygous form in ~8% of African Americans and in homozygous form in 1 in 400. Between 2 and 3% of African Americans carry a hemoglobin C allele.

INHERITANCE AND ONTOGENY

Hemoglobinopathies are autosomal codominant traits—thus, compound heterozygotes who inherit a different abnormal mutant allele from each parent exhibit composite features of each. For example, patients inheriting sickle β thalassemia exhibit features of β thalassemia and sickle cell anemia. The α chain is present in HbA, HbA₂, and HbF; α -chain mutations thus cause abnormalities in all three. The α -globin hemoglobinopathies are symptomatic in utero and after birth because normal function of the α -globin gene is required throughout gestation and adult life. In contrast, infants with β -globin hemoglobinopathies tend to be asymptomatic until 3–9 months of age, when HbA has largely replaced HbF. Prevention or partial reversion of the switch should thus be an effective therapeutic strategy for β -chain hemoglobinopathies.

DETECTION AND CHARACTERIZATION OF HEMOGLOBINOPATHIES—GENERAL METHODS

Electrophoretic techniques are still widely used for hemoglobin analysis. Electrophoresis at pH 8.6 on cellulose acetate membranes is especially simple, inexpensive, and reliable for initial screening. Agar gel electrophoresis at pH 6.1 in citrate buffer is often used as a complementary method because each method detects different variants. Some important variants are electrophoretically silent. These mutant hemoglobins can usually be characterized by more specialized techniques such as mass spectroscopy, which is rapidly replacing electrophoresis for initial analysis.

Quantitation of the hemoglobin profile is often desirable. HbA₂ is frequently elevated in β thalassemia trait and depressed in iron deficiency. HbF is elevated in HPFH, some β thalassemia syndromes, and occasional periods of erythroid stress or marrow dysplasia. For characterization of sickle cell trait, sickle thalassemia syndromes, or HbSC disease, and for monitoring the progress of exchange transfusion therapy to lower the percentage of circulating HbS, quantitation of individual hemoglobins is also required. In most laboratories, quantitation is performed only if the test is specifically ordered. Complete characterization, including amino acid sequencing or gene cloning and sequencing, is readily available from several reference laboratories.

Because some variants can comigrate with HbA or HbS (sickle hemoglobin), electrophoretic assessment should always be regarded as incomplete unless functional assays for hemoglobin sickling, solubility, or oxygen affinity are also performed, as dictated by the clinical presentation. The best sickling assays involve measurement of the degree to which the hemoglobin sample becomes insoluble, or gelled, as it is deoxygenated (i.e., sickle solubility test). Unstable hemoglobins are detected by their precipitation in isopropanol or after heating to 50°C. High-O₂ affinity and low-O₂ affinity variants are detected

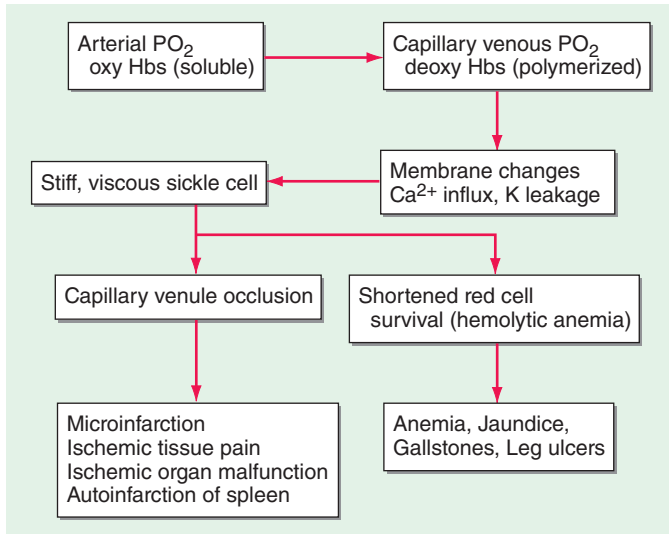


FIGURE 127-3 Pathophysiology of sickle cell crisis.

by quantitating the P_{50} , the partial pressure of oxygen at which the hemoglobin sample becomes 50% saturated with oxygen. Direct tests for the percent carboxyhemoglobin and methemoglobin, using spectrophotometric techniques, can readily be obtained from most clinical laboratories on an urgent basis.

Laboratory evaluation remains an adjunct, rather than the sole diagnostic aid. Diagnosis is best established by recognition of a characteristic history, physical findings, peripheral blood smear morphology, and abnormalities of the complete blood cell count (e.g., profound microcytosis with minimal anemia in thalassemia trait).

STRUCTURALLY ABNORMAL HEMOGLOBINS

SICKLE CELL SYNDROMES

The sickle cell syndromes are caused by a mutation in the β -globin gene that changes the sixth amino acid from glutamic acid to valine. HbS ($\alpha_2\beta_2^{Glu\rightarrow Val}$) polymerizes reversibly when deoxygenated to form a gelatinous network of fibrous polymers that stiffen the RBC membrane, increase viscosity, and cause dehydration due to potassium leakage and calcium influx (Fig. 127-3). These changes also produce the sickle shape. Sickled cells lose the pliability needed to traverse small capillaries. They possess altered “sticky” membranes that are abnormally adherent to the endothelium of small venules. These abnormalities provoke unpredictable episodes of microvascular vasoocclusion and premature RBC destruction (hemolytic anemia). Hemolysis occurs because the spleen destroys the abnormal RBC. The rigid adherent cells clog small capillaries and venules, causing tissue ischemia, acute pain, and gradual end-organ damage. This venoocclusive component usually dominates the clinical course. Prominent manifestations include episodes of ischemic pain (i.e., painful crises) and ischemic malfunction or frank infarction in the spleen, central nervous system, bones, joints, liver, kidneys, and lungs (Fig. 127-3).

Several sickle syndromes occur as the result of inheritance of HbS from one parent and another hemoglobinopathy, such as β thalassemia or HbC ($\alpha_2\beta_2^{Glu\rightarrow Lys}$), from the other parent. The prototype disease, sickle cell anemia, is the homozygous state for HbS (Table 127-2).

Clinical Manifestations of Sickle Cell Anemia Most patients with sickling syndromes suffer from hemolytic anemia, with hematocrits from 15 to 30%, and significant reticulocytosis. Anemia was once thought to exert protective effects against vasoocclusion by reduc-

ing blood viscosity. However, natural history and drug therapy trials suggest that an increase in the hematocrit and feedback inhibition of reticulocytosis might be beneficial, even at the expense of increased blood viscosity. The role of adhesive reticulocytes in vasoocclusion might account for these paradoxical effects.

Granulocytosis is common. The white count can fluctuate substantially and unpredictably during and between painful crises, infectious episodes, and other intercurrent illnesses.

Vasoocclusion causes protean manifestations. Intermittent episodes of vasoocclusion in connective and musculoskeletal structures produce ischemia manifested by acute pain and tenderness, fever, tachycardia, and anxiety. These recurrent episodes, called *painful crises*, are the most common clinical manifestation. Their frequency and severity vary greatly. Pain can develop almost anywhere in the body and may last from a few hours to 2 weeks. Repeated crises requiring hospitalization (>3 episodes per year) correlate with reduced survival in adult life, suggesting that these episodes are associated with accumulation of chronic end-organ damage. Provocative factors include infection, fever, excessive exercise, anxiety, abrupt changes in temperature, hypoxia, or hypertonic dyes.

Repeated microinfarction can destroy tissues having microvascular beds prone to sickling. Thus, splenic function is frequently lost within the first 18–36 months of life, causing susceptibility to infection, particularly by pneumococci. Acute venous obstruction of the spleen (*splenic sequestration crisis*), a rare occurrence in early childhood, may require emergency transfusion and/or splenectomy to prevent trapping of the entire arterial output in the obstructed spleen. Occlusion of retinal vessels can produce hemorrhage, neovascularization, and eventual detachments. Renal papillary necrosis invariably produces isosthenuria. More widespread renal necrosis leads to renal failure in adults, a common late cause of death. Bone and joint ischemia can lead to aseptic necrosis, especially of the femoral or humeral heads; chronic arthropathy; and unusual susceptibility to osteomyelitis, which may be caused by organisms, such as *Salmonella*, rarely encountered in other settings. The *hand-foot syndrome* is caused by painful infarcts of the digits and dactylitis. Stroke is especially common in children; a small subset tends to suffer repeated episodes. Stroke is less common in adults and is often hemorrhagic. A particularly painful complication in males is priapism, due to infarction of the penile venous outflow tracts; permanent impotence is a frequent consequence. Chronic lower leg ulcers probably arise from ischemia and superinfection in the distal circulation.

Acute chest syndrome is a distinctive manifestation characterized by chest pain, tachypnea, fever, cough, and arterial oxygen desaturation. It can mimic pneumonia, pulmonary emboli, bone marrow infarction and embolism, myocardial ischemia, or in situ lung infarction. Acute chest syndrome is thought to reflect in situ sickling within the lung, producing pain and temporary pulmonary dysfunction. Often it is difficult or impossible to distinguish among other possibilities. Pulmonary infarction and pneumonia are the most frequent underlying or concomitant conditions in patients with this syndrome. Repeated episodes of acute chest pain correlate with reduced survival. Acutely, reduction in arterial oxygen saturation is especially ominous because it promotes sickling on a massive

TABLE 127-2 CLINICAL FEATURES OF SICKLE HEMOGLOBINOPATHIES

Condition	Clinical Abnormalities	Hemoglobin Level, g/L (g/dL)	MCV, fL	Hemoglobin Electrophoresis
Sickle cell trait	None; rare painless hematuria	Normal	Normal	HbS/A: 40/60
Sickle cell anemia	Vasoocclusive crises with infarction of spleen, brain, marrow, kidney, lung; aseptic necrosis of bone; gallstones; priapism; ankle ulcers	70–100 (7–10)	80–100	HbS/A: 100/0 HbF: 2–25%
S/ β^0 thalassemia	Vasoocclusive crises; aseptic necrosis of bone	70–100 (7–10)	60–80	HbS/A: 100/0 HbF: 1–10%
S/ β^+ thalassemia	Rare crises and aseptic necrosis	100–140 (10–14)	70–80	HbS/A: 60/40
Hemoglobin SC	Rare crises and aseptic necrosis; painless hematuria	100–140 (10–14)	80–100	HbS/A: 50/0 HbC: 50%

scale. Chronic acute or subacute pulmonary crises lead to pulmonary hypertension and cor pulmonale, an increasingly common cause of death as patients survive longer. Considerable controversy exists about the possible role played by free plasma HbS in scavenging nitrogen dioxide (NO₂), thus raising pulmonary vascular tone. Trials of sildenafil to restore NO₂ levels were terminated because of adverse effects.

Chronic subacute central nervous system damage in the absence of an overt stroke is a distressingly common phenomenon beginning in early childhood. Modern functional imaging techniques have pinpointed circulatory dysfunction due to a likely CNS sickle vasculopathy; these changes correlate with an array of cognitive and behavioral abnormalities in children and young adults. It is important to be aware of these often subtle changes because they can complicate clinical management or be misinterpreted as “difficult patient” behaviors.

Sickle cell syndromes are remarkable for their clinical heterogeneity. Some patients remain virtually asymptomatic into or even through adult life, while others suffer repeated crises requiring hospitalization from early childhood. Patients with sickle thalassemia and sickle-HbE tend to have similar, slightly milder symptoms, perhaps because of the ameliorating effects of production of other hemoglobins within the RBC. Hemoglobin SC disease, one of the more common variants of sickle cell anemia, is frequently marked by lesser degrees of hemolytic anemia and a greater propensity for the development of retinopathy and aseptic necrosis of bones. In most respects, however, the clinical manifestations resemble sickle cell anemia. Some rare hemoglobin variants actually aggravate the sickling phenomenon.

The clinical variability in different patients inheriting the same disease-causing mutation (sickle hemoglobin) has made sickle cell disease the focus of efforts to identify modifying genetic polymorphisms in other genes that might account for the heterogeneity. The complexity of the data obtained thus far has dampened the expectation that genome-wide analysis will yield individualized profiles that predict a patient's clinical course. Nevertheless, a number of interesting patterns have emerged from these modifying gene analyses. For example, genes affecting the inflammatory response or cytokine expression appear to be modifying candidates. Genes that affect transcriptional regulation of lymphocytes may also be involved.

Clinical Manifestations of Sickle Cell Trait Sickle cell trait is often asymptomatic. Anemia and painful crises are rare. An uncommon but highly distinctive symptom is painless hematuria often occurring in adolescent males, probably due to papillary necrosis. Isosthenuria is a more common manifestation of the same process. Sloughing of papillae with urethral obstruction has been reported, as have isolated cases of massive sickling or sudden death due to exposure to high altitudes or extremes of exercise and dehydration. Avoidance of dehydration or extreme physical stress should be advised.

Diagnosis Sickle cell syndromes are suspected on the basis of hemolytic anemia, RBC morphology (Fig. 127-4), and intermittent episodes of ischemic pain. Diagnosis is confirmed by hemoglobin electrophoresis, mass spectroscopy, and the sickling tests already discussed. Thorough characterization of the exact hemoglobin profile of the patient is important, because sickle thalassemia and hemoglobin SC disease have distinct prognoses or clinical features. Diagnosis is usually established in childhood, but occasional patients, often with compound heterozygous states, do not develop symptoms until the onset of puberty, pregnancy, or early adult life. Genotyping of family members and potential parental partners is critical for genetic counseling. Details of the childhood history establish prognosis and need for aggressive or experimental therapies. Factors associated with increased morbidity and reduced survival include more than three crises requiring hospitalization per year, chronic neutrophilia, a history of splenic sequestration or hand-foot syndrome, and second episodes of acute chest syndrome. Patients with a history of cerebrovascular accidents are at higher risk for repeated episodes and require partial exchange transfusion and especially close monitoring using Doppler carotid flow measurements. Patients with severe or repeated episodes of acute

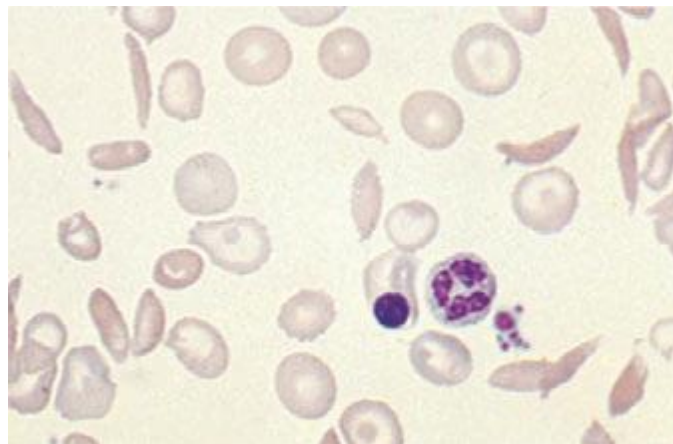


FIGURE 127-4 Sickle cell anemia. The elongated and crescent-shaped red blood cells seen on this smear represent circulating irreversibly sickled cells. Target cells and a nucleated red blood cell are also seen.

chest syndrome may need lifelong transfusion support, using partial exchange transfusion, if possible.

TREATMENT SICKLE CELL SYNDROMES

Patients with sickle cell syndromes require ongoing continuity of care. Familiarity with the pattern of symptoms provides the best safeguard against excessive use of the emergency room, hospitalization, and habituation to addictive narcotics. Additional preventive measures include regular slit-lamp examinations to monitor development of retinopathy; antibiotic prophylaxis appropriate for splenectomized patients during dental or other invasive procedures; and vigorous oral hydration during or in anticipation of periods of extreme exercise, exposure to heat or cold, emotional stress, or infection. Pneumococcal and *Haemophilus influenzae* vaccines are less effective in splenectomized individuals. Thus, patients with sickle cell anemia should be vaccinated early in life.

The management of an acute painful crisis includes vigorous hydration, thorough evaluation for underlying causes (such as infection), and aggressive analgesia administered by a standing order and/or patient-controlled analgesia (PCA) pump. Morphine (0.1–0.15 mg/kg every 3–4 h) should be used to control severe pain. Bone pain may respond as well to ketorolac (30–60 mg initial dose, then 15–30 mg every 6–8 h). Inhalation of nitrous oxide can provide short-term pain relief, but great care must be exercised to avoid hypoxia and respiratory depression. Nitrous oxide also elevates O₂ affinity, reducing O₂ delivery to tissues. Its use should be restricted to experts. Many crises can be managed at home with oral hydration and oral analgesia. Use of the emergency room should be reserved for especially severe symptoms or circumstances in which other processes, e.g., infection, are strongly suspected. Nasal oxygen should be used as appropriate to protect arterial saturation. Most crises resolve in 1–7 days. Use of blood transfusion should be reserved for extreme cases: transfusions do not shorten the duration of the crisis.

No tests are definitive to diagnose acute painful crisis. Critical to good management is an approach that recognizes that most patients reporting crisis symptoms do indeed have crisis or another significant medical problem. Diligent diagnostic evaluation for underlying causes is imperative, even though these are found infrequently. In adults, the possibility of aseptic necrosis or sickle arthropathy must be considered, especially if pain and immobility become repeated or chronic at a single site. Nonsteroidal anti-inflammatory agents are often effective for sickle cell arthropathy.

Acute chest syndrome is a medical emergency that may require management in an intensive care unit. Hydration should be monitored carefully to avoid the development of pulmonary edema, and oxygen therapy should be especially vigorous for protection of

636 arterial saturation. Diagnostic evaluation for pneumonia and pulmonary embolism should be especially thorough, since these may occur with atypical symptoms. Critical interventions are transfusion to maintain a hematocrit >30, and emergency exchange transfusion if arterial saturation drops to <90%. As patients with sickle cell syndrome increasingly survive into their fifth and sixth decades, end-stage renal failure and pulmonary hypertension are becoming increasingly prominent causes of end-stage morbidity. A sickle cell cardiomyopathy and/or premature coronary artery disease may compromise cardiac function in later years. Sickle cell patients have received kidney transplants, but they often experience an increase in the frequency and severity of crises, possibly due to increased infection as a consequence of immunosuppression.

The most significant advance in the therapy of sickle cell anemia has been the introduction of hydroxyurea as a mainstay of therapy for patients with severe symptoms. Hydroxyurea (10–30 mg/kg per day) increases fetal hemoglobin and may also exert beneficial effects on RBC hydration, vascular wall adherence, and suppression of the granulocyte and reticulocyte counts; dosage is titrated to maintain a white cell count between 5000 and 8000/μL. White cells and reticulocytes may play a major role in the pathogenesis of sickle cell crisis, and their suppression may be an important side benefit of hydroxyurea therapy.

Hydroxyurea should be considered in patients experiencing repeated episodes of acute chest syndrome or with more than three crises per year requiring hospitalization. The utility of this agent for reducing the incidence of other complications (priapism, retinopathy) is under evaluation, as are the long-term side effects. To date, however, minimal risk of bone marrow dyscrasias or other neoplasms has been documented. Hydroxyurea offers broad benefits to most patients whose disease is severe enough to impair their functional status, and it may improve survival. HbF levels increase in most patients within a few months.

The antitumor drug 5-azacytidine was the first agent found to elevate HbF. It never achieved widespread use because of concerns about acute toxicity and carcinogenesis. However, low doses of the related agent 5-deoxyazacytidine (decitabine) can elevate HbF with more acceptable toxicity.

Bone marrow transplantation can provide definitive cures but is known to be effective and safe only in children. Clinical trials studying partially myeloablative conditioning regimens (“mini” transplants) are likely to support more widespread use of this modality in older patients. Prognostic features justifying bone marrow transplant are the presence of repeated crises early in life, a high neutrophil count, or the development of hand-foot syndrome. Children at risk for stroke can now be identified through the use of Doppler ultrasound techniques. Prophylactic exchange transfusion appears to substantially reduce the risk of stroke in this population. Children who do suffer a cerebrovascular accident should be maintained for at least 3–5 years on a program of vigorous exchange transfusion, as the risk of second strokes is extremely high.

Gene therapy for sickle cell anemia is being intensively pursued, but no safe measures are currently available. The development of newer methods of direct gene correction in situ (e.g., zinc finger nucleases, or “CRISPR” [clustered regularly interspaced short palindromic repeats] technology) could well find clinical use in these patients. Experimental methods of derepressing HbF by interfering with Bcl11a are also being explored.

UNSTABLE HEMOGLOBINS

Amino acid substitutions that reduce solubility or increase susceptibility to oxidation result in unstable hemoglobins that precipitate, forming inclusion bodies injurious to the RBC membrane. Representative mutations are those that interfere with contact points between the α and β subunits (e.g., Hb Philly [$\beta^{35Tyr\rightarrow Phe}$]), alter the helical segments (e.g., Hb Genova [$\beta^{28Leu\rightarrow Pro}$]), or disrupt interactions of the hydrophobic pockets of the globin subunits with heme (e.g., Hb Köln [$\beta^{98Val\rightarrow Met}$]) (Table 127-3). The inclusions, called *Heinz bodies*, are clinically

TABLE 127-3 REPRESENTATIVE ABNORMAL HEMOGLOBINS WITH ALTERED SYNTHESIS OR FUNCTION

Designation	Mutation	Population	Main Clinical Effects ^a
Sickle or S	$\beta^{6Glu\rightarrow Val}$	African	Anemia, ischemic infarcts
C	$\beta^{6Glu\rightarrow Lys}$	African	Mild anemia; interacts with HbS
E	$\beta^{26Glu\rightarrow Lys}$	Southeast Asian	Microcytic anemia, splenomegaly, thalassemic phenotype
Köln	$\beta^{98Val\rightarrow Met}$	Sporadic	Hemolytic anemia, Heinz bodies when splenectomized
Yakima	$\beta^{99Asp\rightarrow His}$	Sporadic	Polycythemia
Kansas	$\beta^{102Asn\rightarrow Lys}$	Sporadic	Mild anemia
M Iwata	$\beta^{87His\rightarrow Tyr}$	Sporadic	Methemoglobinemia

^aSee text for details.

detectable by staining with supravital dyes such as crystal violet. Removal of these inclusions by the spleen generates pitted, rigid cells that have shortened life spans, producing hemolytic anemia of variable severity, sometimes requiring chronic transfusion support. Splenectomy may be needed to correct the anemia. Leg ulcers and premature gallbladder disease due to bilirubin loading are frequent stigmata.

Unstable hemoglobins occur sporadically, often by spontaneous new mutations. Heterozygotes are often symptomatic because a significant Heinz body burden can develop even when the unstable variant accounts for only a portion of the total hemoglobin. Symptomatic unstable hemoglobins tend to be β -globin variants, because sporadic mutations affecting only one of the four α globins alleles would generate only 20–30% abnormal hemoglobin.

HEMOGLOBINS WITH ALTERED OXYGEN AFFINITY

High-affinity hemoglobins (e.g., Hb Yakima [$\beta^{99Asp\rightarrow His}$]) bind oxygen more readily but deliver less O₂ to tissues at normal capillary Po₂ levels (Fig. 127-2). Mild tissue hypoxia ensues, stimulating RBC production and erythrocytosis (Table 127-3). In extreme cases, the hematocrits can rise to 60–65%, increasing blood viscosity and producing typical symptoms (headache, somnolence, or dizziness). Phlebotomy may be required. Typical mutations alter interactions within the heme pocket or disrupt the Bohr effect or salt-bond site. Mutations that impair the interaction of HbA with 2,3-BPG can increase O₂ affinity because 2,3-BPG binding lowers O₂ affinity.

Low-affinity hemoglobins (e.g., Hb Kansas [$\beta^{102Asn\rightarrow Lys}$]) bind sufficient oxygen in the lungs, despite their lower oxygen affinity, to achieve nearly full saturation. At capillary oxygen tensions, they lose sufficient amounts of oxygen to maintain homeostasis at a low hematocrit (Fig. 127-2) (*pseudoanemia*). Capillary hemoglobin desaturation can also be sufficient to produce clinically apparent cyanosis. Despite these findings, patients usually require no specific treatment.

METHEMOGLOBINEMIAS

Methemoglobin is generated by oxidation of the heme iron moieties to the ferric state, causing a characteristic bluish-brown muddy color resembling cyanosis. Methemoglobin has such high oxygen affinity that virtually no oxygen is delivered. Levels >50–60% are often fatal.

Congenital methemoglobinemia arises from globin mutations that stabilize iron in the ferric state (e.g., HbM Iwata [$\alpha^{87His\rightarrow Tyr}$], Table 127-3) or from mutations that impair the enzymes that reduce methemoglobin to hemoglobin (e.g., methemoglobin reductase, NADP diaphorase). Acquired methemoglobinemia is caused by toxins that oxidize heme iron, notably nitrate and nitrite-containing compounds, including drugs commonly used in cardiology and anesthesiology.

DIAGNOSIS AND MANAGEMENT OF PATIENTS WITH UNSTABLE HEMOGLOBINS, HIGH-AFFINITY HEMOGLOBINS, AND METHEMOGLOBINEMIA

Unstable hemoglobin variants should be suspected in patients with nonimmune hemolytic anemia, jaundice, splenomegaly, or premature biliary tract disease. Severe hemolysis usually presents during infancy as neonatal jaundice or anemia. Milder cases may present in adult life with anemia or only as unexplained reticulocytosis, hepatosplenomegaly, premature biliary tract disease, or leg ulcers. Because spontaneous

mutation is common, family history of anemia may be absent. The peripheral blood smear often shows anisocytosis, abundant cells with punctate inclusions, and irregular shapes (i.e., poikilocytosis).

The two best tests for diagnosing unstable hemoglobins are the Heinz body preparation and the isopropanol or heat stability test. Many unstable Hb variants are electrophoretically silent. A normal electrophoresis does not rule out the diagnosis. Mass spectroscopy or direct gene analysis will provide a definitive diagnosis.

Severely affected patients may require transfusion support for the first 3 years of life, because splenectomy before age 3 is associated with a significantly higher immune deficit. Splenectomy is usually effective thereafter, but occasional patients may require lifelong transfusion support. After splenectomy, patients can develop cholelithiasis and leg ulcers, hypercoagulable states, and susceptibility to overwhelming sepsis. Splenectomy should thus be avoided or delayed unless it is the only alternative. Precipitation of unstable hemoglobins is aggravated by oxidative stress, e.g., infection and antimalarial drugs, which should be avoided where possible.

High-O₂ affinity hemoglobin variants should be suspected in patients with erythrocytosis. The best test for confirmation is measurement of the P₅₀. A high-O₂ affinity hemoglobin causes a significant left shift (i.e., lower numeric value of the P₅₀); confounding conditions, e.g., tobacco smoking or carbon monoxide exposure, can also lower the P₅₀.

High-affinity hemoglobins are often asymptomatic; rubor or plethora may be telltale signs. When the hematocrit approaches 60%, symptoms of high blood viscosity and sluggish flow (headache, lethargy, dizziness, etc.) may be present. These persons may benefit from judicious phlebotomy. Erythrocytosis represents an appropriate attempt to compensate for the impaired oxygen delivery by the abnormal variant. Overzealous phlebotomy may stimulate increased erythropoiesis or aggravate symptoms by thwarting this compensatory mechanism. The guiding principle of phlebotomy should be to improve oxygen delivery by reducing blood viscosity and increasing blood flow rather than restoration of a normal hematocrit. Phlebotomy-induced modest iron deficiency may aid in control.

Low-affinity hemoglobins should be considered in patients with cyanosis or a low hematocrit with no other reason apparent after thorough evaluation. The P₅₀ test confirms the diagnosis. Counseling and reassurance are the interventions of choice.

Methemoglobin should be suspected in patients with hypoxic symptoms who appear cyanotic but have a Pao₂ sufficiently high that hemoglobin should be fully saturated with oxygen. A history of nitrite or other oxidant ingestions may not always be available; some exposures may be inapparent to the patient, and others may result from suicide attempts. The characteristic muddy appearance of freshly drawn blood can be a critical clue. The best diagnostic test is methemoglobin assay, which is usually available on an emergency basis.

Methemoglobinemia often causes symptoms of cerebral ischemia at levels >15%; levels >60% are usually lethal. Intravenous injection of 1 mg/kg of methylene blue is effective emergency therapy. Milder cases and follow-up of severe cases can be treated orally with methylene blue (60 mg three to four times each day) or ascorbic acid (300–600 mg/d).

THALASSEMIA SYNDROMES

The thalassemia syndromes are inherited disorders of α - or β -globin biosynthesis. The reduced supply of globin diminishes production of hemoglobin tetramers, causing hypochromia and microcytosis. Unbalanced accumulation of α and β subunits occurs because the synthesis of the unaffected globins proceeds at a normal rate. Unbalanced chain accumulation dominates the clinical phenotype. Clinical severity varies widely, depending on the degree to which the synthesis of the affected globin is impaired, altered synthesis of other globin chains, and coinheritance of other abnormal globin alleles.

CLINICAL MANIFESTATIONS OF β THALASSEMIA SYNDROMES

Mutations causing thalassemia can affect any step in the pathway of globin gene expression: transcription, processing of the mRNA precursor, translation, and posttranslational metabolism of the β -globin

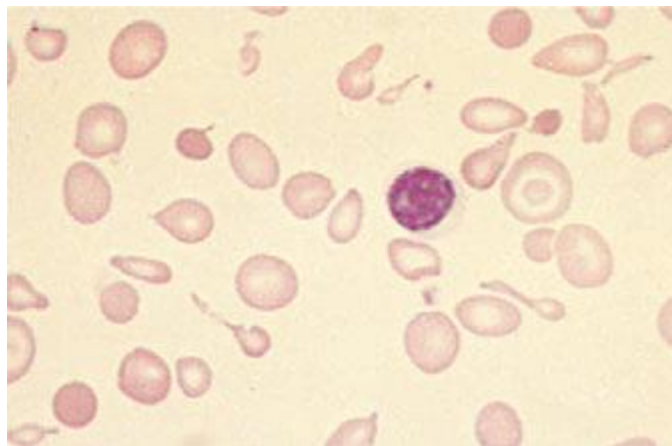


FIGURE 127-5 β Thalassemia intermedia. Microcytic and hypochromic red blood cells are seen that resemble the red blood cells of severe iron-deficiency anemia. Many elliptical and teardrop-shaped red blood cells are noted.

polypeptide chain. The most common forms arise from mutations that derange splicing of the mRNA precursor or prematurely terminate translation of the mRNA.

Hypochromia and microcytosis characterize all forms of β thalassemia because of the reduced amounts of hemoglobin tetramers (Fig. 127-5). In heterozygotes (β thalassemia trait), this is the only abnormality seen. Anemia is minimal. In more severe homozygous states, unbalanced α - and β -globin accumulation causes accumulation of highly insoluble unpaired α chains. They form toxic inclusion bodies that kill developing erythroblasts in the marrow. Few of the proerythroblasts beginning erythroid maturation survive. The surviving RBCs bear a burden of inclusion bodies that are detected in the spleen, shortening the RBC life span and producing severe hemolytic anemia. The resulting profound anemia stimulates erythropoietin release and compensatory erythroid hyperplasia, but the marrow response is sabotaged by the ineffective erythropoiesis. Anemia persists. Erythroid hyperplasia can become exuberant and produce masses of extramedullary erythropoietic tissue in the liver and spleen.

Massive bone marrow expansion deranges growth and development. Children develop characteristic “chipmunk” facies due to maxillary marrow hyperplasia and frontal bossing. Thinning and pathologic fracture of long bones and vertebrae may occur due to cortical invasion by erythroid elements and profound growth retardation. Hemolytic anemia causes hepatosplenomegaly, leg ulcers, gallstones, and high-output congestive heart failure. The conscription of caloric resources to support erythropoiesis leads to inanition, susceptibility to infection, endocrine dysfunction, and in the most severe cases, death during the first decade of life. Chronic transfusions with RBCs improve oxygen delivery, suppress the excessive ineffective erythropoiesis, and prolong life, but the inevitable side effects, notably iron overload, often prove fatal by age 30 years.

Severity is highly variable. Known modulating factors are those that ameliorate the burden of unpaired α -globin inclusions. Alleles associated with milder synthetic defects and coinheritance of a thalassemia trait reduce clinical severity by reducing accumulation of excess α globin. HbF persists to various degrees in β thalassemias. γ -Globin gene chains can substitute for β chains, generating more hemoglobin and reducing the burden of α -globin inclusions. The terms *β thalassemia major* and *β thalassemia intermedia* are used to reflect the clinical heterogeneity. Patients with β thalassemia major require intensive transfusion support to survive. Patients with β thalassemia intermedia have a somewhat milder phenotype and can survive without transfusion. The terms *β thalassemia minor* and *β thalassemia trait* describe asymptomatic heterozygotes for β thalassemia.

THALASSEMIA SYNDROMES

The four classic α thalassemias, most common in Asians, are a thalassemia-2 trait, in which one of the four α -globin loci is deleted;

TABLE 127-4 THE α THALASSEMIAS

Condition	Hemoglobin A, %	Hemoglobin H (β_4), %	Hemoglobin Level, g/L (g/dL)	MCV, fL
Normal	97	0	150 (15)	90
Silent thalassemia: $-\alpha/\alpha$	98–100	0	150 (15)	90
Thalassemia trait: $-\alpha/-\alpha$ homozygous α -thal-2 ^a or $-\alpha/\alpha$ heterozygous α -thal-1 ^a	85–95	Rare red blood cell inclusions	120–130 (12–13)	70–80
Hemoglobin H disease: $---/\alpha$ heterozygous α -thal-1/ α -thal-2	70–95	5–30	60–100 (6–10)	60–70
Hydrops fetalis: $---/---$ homozygous α -thal-1	0	5–10 ^b	Fatal in utero or at birth	

^aWhen both α alleles on one chromosome are deleted, the locus is called α -thal-1; when only a single α allele on one chromosome is deleted, the locus is called α -thal-2. ^b90–95% of the hemoglobin is hemoglobin Barts (tetramers of γ chains).

α thalassemia-1 trait, with two deleted loci; HbH disease, with three loci deleted; and hydrops fetalis with Hb Barts, with all four loci deleted (Table 127-4). Nondeletion forms of α thalassemia also exist.

α Thalassemia-2 trait is an asymptomatic, silent carrier state. α Thalassemia-1 trait resembles β thalassemia minor. Offspring doubly heterozygous for α thalassemia-2 and α thalassemia-1 exhibit a more severe phenotype called *HbH disease*. Heterozygosity for a deletion that removes both genes from the same chromosome (*cis* deletion) is common in Asians and in those from the Mediterranean region, as is homozygosity for α thalassemia-2 (*trans* deletion). Both produce asymptomatic hypochromia and microcytosis.

In *HbH disease*, HbA production is only 25–30% normal. Fetuses accumulate some unpaired γ chains (Hb Barts; γ -chain tetramers). In adults, unpaired β chains accumulate and are soluble enough to form β_4 tetramers called HbH. HbH forms few inclusions in erythroblasts and precipitates in circulating RBC. Patients with HbH disease have thalassemia intermedia characterized by moderately severe hemolytic anemia but milder ineffective erythropoiesis. Survival into midadult life without transfusions is common.

The homozygous state for the α thalassemia-1 *cis* deletion (hydrops fetalis) causes total absence of α -globin synthesis. No physiologically useful hemoglobin is produced beyond the embryonic stage. Excess γ globin forms tetramers called *Hb Barts* (γ_4), which has a very high oxygen affinity. It delivers almost no O_2 to fetal tissues, causing tissue asphyxia, edema (hydrops fetalis), congestive heart failure, and death in utero. α Thalassemia-2 trait is common (15–20%) among people of African descent. The *cis* α thalassemia-1 deletion is almost never seen, however. Thus, α thalassemia-2 and the *trans* form of α thalassemia-1 are very common, but HbH disease and hydrops fetalis are rare.

It has been known for some time that some patients with myelodysplasia or erythroleukemia produce RBC clones containing HbH. This phenomenon is due to mutations in the ATRX pathway that affect the LCR of the α -globin gene cluster.

DIAGNOSIS AND MANAGEMENT OF THALASSEMIAS

The diagnosis of β -thalassemia major is readily made during childhood on the basis of severe anemia accompanied by the characteristic signs of massive ineffective erythropoiesis: hepatosplenomegaly, profound microcytosis, a characteristic blood smear (Fig. 127-5), and elevated levels of HbF, HbA₂, or both. Many patients require chronic hypertransfusion therapy designed to maintain a hematocrit of at least 27–30% so that erythropoiesis is suppressed. Splenectomy is required if the annual transfusion requirement (volume of RBCs per kilogram of body weight per year) increases by >50%. Folic acid supplements may be useful. Vaccination with Pneumovax in anticipation of eventual splenectomy is advised, as is close monitoring for infection, leg ulcers, and biliary tract disease. Many patients develop endocrine deficiencies as a result of iron overload. Early endocrine evaluation is required for glucose intolerance, thyroid dysfunction, and delayed onset of puberty or secondary sexual characteristics.

Patients with β thalassemia intermedia exhibit similar stigmata but can survive without chronic hypertransfusion. Management is particularly challenging because a number of factors can aggravate the anemia, including infection, onset of puberty, and development of splenomegaly and hypersplenism. Some patients may eventually benefit from splenectomy. The expanded erythron can cause absorption of excessive dietary iron and hemosiderosis, even without transfusion. Some patients eventually become transfusion dependent.

β Thalassemia minor (i.e., thalassemia trait) usually presents as profound microcytosis and hypochromia with target cells, but only minimal or mild anemia. The mean corpuscular volume is

rarely >75 fL; the hematocrit is rarely <30–33%. Hemoglobin analysis classically reveals an elevated HbA₂ (3.5–7.5%), but some forms are associated with normal HbA₂ and/or elevated HbF. Genetic counseling and patient education are essential. Patients with β thalassemia trait should be warned that their blood picture resembles iron deficiency and can be misdiagnosed. They should eschew empirical use of iron, yet iron deficiency requiring replacement therapy can develop during pregnancy or from chronic bleeding.

Persons with α thalassemia trait may exhibit mild hypochromia and microcytosis usually without anemia. HbA₂ and HbF levels are normal. Affected individuals usually require only genetic counseling. HbH disease resembles β thalassemia intermedia, with the added complication that the HbH molecule behaves like moderately unstable hemoglobin. Patients with HbH disease should undergo splenectomy if excessive anemia or a transfusion requirement develops. Oxidative drugs should be avoided. Iron overload leading to death can occur in more severely affected patients.

PREVENTION

Antenatal diagnosis of thalassemia syndromes is now widely available. DNA diagnosis is based on polymerase chain reaction (PCR) amplification of fetal DNA, obtained by amniocentesis or chorionic villus biopsy followed by hybridization to allele-specific oligonucleotide probes or direct DNA sequencing.

THALASSEMIC STRUCTURAL VARIANTS

Thalassemic structural variants are characterized by both defective synthesis and abnormal structure.

HEMOGLOBIN LEPORE

Hb Lepore [$\alpha_2(\delta\beta)_2$] arises by an unequal crossover and recombination event that fuses the proximal end of the δ -gene with the distal end of the closely linked β -gene. It is common in the Mediterranean basin. The resulting chromosome contains only the fused $\delta\beta$ gene. The Lepore ($\delta\beta$) globin is synthesized poorly because the fused gene is under the control of the weak δ -globin promoter. Hb Lepore alleles have a phenotype like β thalassemia, except for the added presence of 2–20% Hb Lepore. Compound heterozygotes for Hb Lepore and a classic β thalassemia allele may also have severe thalassemia.

HEMOGLOBIN E

HbE (i.e., $\alpha_2\beta_2^{26\text{Glu}\rightarrow\text{Lys}}$) is extremely common in Cambodia, Thailand, and Vietnam. The gene has become far more prevalent in the United States as a result of immigration of Asian persons, especially in California, where HbE is the most common variant detected. HbE is mildly unstable but not enough to affect RBC life span significantly. Heterozygotes resemble individuals with a mild β -thalassemia trait. Homozygotes have somewhat more marked abnormalities but are asymptomatic. Compound heterozygotes for



HbE and a β thalassemia gene can have β thalassemia intermedia or β thalassemia major, depending on the severity of the coinherited thalassaemic gene.

The β^E allele contains a single base change in codon 26 that causes the amino acid substitution. This mutation also activates a cryptic RNA splice site, generating a structurally abnormal globin mRNA that cannot be translated, from about 50% of the initial pre-mRNA molecules. The remaining 40–50% are normally spliced and generate functional mRNA that is translated into β^E -globin because the mature mRNA carries the base change that alters codon 26.

Genetic counseling of the persons at risk for HbE should focus especially on the interaction of HbE with β thalassemia, because HbE homozygosity is a condition associated with mildly asymptomatic microcytosis, hypochromia, and hemoglobin levels rarely <100 g/L (<10 g/dL).

HEREDITARY PERSISTENCE OF FETAL HEMOGLOBIN

HPFH is characterized by continued synthesis of high levels of HbF in adult life. No deleterious effects are apparent, even when all of the hemoglobin produced is HbF. These rare patients demonstrate convincingly that prevention or reversal of the fetal to adult hemoglobin switch would provide effective therapy for sickle cell anemia and β thalassemia.

ACQUIRED HEMOGLOBINOPATHIES

The two most important acquired hemoglobinopathies are carbon monoxide poisoning and methemoglobinemia (see above). Carbon monoxide has a higher affinity for hemoglobin than does oxygen; it can replace oxygen and diminish O_2 delivery. Chronic elevation of carboxyhemoglobin levels to 10 or 15%, as occurs in smokers, can lead to secondary polycythemia. Carboxyhemoglobin is cherry red in color and masks the development of cyanosis usually associated with poor O_2 delivery to tissues.

Abnormalities of hemoglobin biosynthesis have also been described in blood dyscrasias. In some patients with myelodysplasia, erythroleukemia, or myeloproliferative disorders, elevated HbF or a mild form of HbH disease may also be seen. The abnormalities are not severe enough to alter the course of the underlying disease.

TREATMENT TRANSFUSIONAL HEMOSIDEROSIS

Chronic blood transfusion can lead to bloodborne infection, alloimmunization, febrile reactions, and lethal iron overload (Chap. 138e). A unit of packed RBCs contains 250–300 mg iron (1 mg/mL). The iron assimilated by a single transfusion of 2 units of packed RBCs is thus equal to a 1- to 2-year oral intake of iron. Iron accumulates in chronically transfused patients because no mechanisms exist for increasing iron excretion: an expanded erythron causes especially rapid development of iron overload because accelerated erythropoiesis promotes excessive absorption of dietary iron. Vitamin C should not be supplemented because it generates free radicals in iron excess states.

Patients who receive >100 units of packed RBCs usually develop hemosiderosis. The ferritin level rises, followed by early endocrine dysfunction (glucose intolerance and delayed puberty), cirrhosis, and cardiomyopathy. Liver biopsy shows both parenchymal and reticuloendothelial iron. The superconducting quantum-interference device (SQUID) is accurate at measuring hepatic iron but not widely available. Cardiac toxicity is often insidious. Early development of pericarditis is followed by dysrhythmia and pump failure. The onset of heart failure is ominous, often presaging death within a year (Chap. 428).

The decision to start long-term transfusion support should also prompt one to institute therapy with iron-chelating agents. Deferoxamine (Desferal) is for parenteral use. Its iron-binding kinetics require chronic slow infusion via a metering pump. The constant

presence of the drug improves the efficiency of chelation and protects tissues from occasional releases of the most toxic fraction of iron—low-molecular-weight iron—which may not be sequestered by protective proteins.

Deferoxamine is relatively nontoxic. Occasional cataracts, deafness, and local skin reactions, including urticaria, occur. Skin reactions can usually be managed with antihistamines. Negative iron balance can be achieved, even in the face of a high transfusion requirement, but this alone does not prevent long-term morbidity and mortality in chronically transfused patients. Irreversible end-organ deterioration develops at relatively modest levels of iron overload, even if symptoms do not appear for many years thereafter. To enjoy a significant survival advantage, chelation must begin before 5–8 years of age in β thalassemia major.

Deferasirox is an oral iron-chelating agent. Single daily doses of 20–30 mg/kg deferasirox produced reductions in liver iron concentration comparable to deferoxamine in long-term transfused adult and pediatric patients. Deferasirox produces some elevations in liver enzymes and slight but persistent increases in serum creatinine, without apparent clinical consequence. Other toxicities are similar to those of deferoxamine. Its toxicity profile is acceptable, although long-term effects are still being evaluated.

EXPERIMENTAL THERAPIES

BONE MARROW TRANSPLANTATION, GENE THERAPY, AND MANIPULATION OF HbF

Bone marrow transplantation provides stem cells able to express normal hemoglobin; it has been used in a large number of patients with β thalassemia and a smaller number of patients with sickle cell anemia. Early in the course of disease, before end-organ damage occurs, transplantation is curative in 80–90% of patients. In highly experienced centers, the treatment-related mortality is $<10\%$. Because survival into adult life is possible with conventional therapy, the decision to transplant is best made in consultation with specialized centers.

Gene therapy of thalassemia and sickle cell disease has proved to be an elusive goal, but experimental advances are raising expectations.

Reestablishing high levels of fetal hemoglobin synthesis should ameliorate the symptoms of β -chain hemoglobinopathies. Cytotoxic agents such as hydroxyurea and cytarabine promote high levels of HbF synthesis, probably by stimulating proliferation of the primitive HbF-producing progenitor cell population (i.e., F cell progenitors). Unfortunately, this regimen has not yet been effective in β thalassemia. Butyrates stimulate HbF production, but only transiently. Pulsed or intermittent administration has been found to sustain HbF induction in the majority of patients with sickle cell disease. It is unclear whether butyrates will have similar activity in patients with β thalassemia.

APLASTIC AND HYPOPLASTIC CRISIS IN PATIENTS WITH HEMOGLOBINOPATHIES

Patients with hemolytic anemias sometimes exhibit an alarming decline in hematocrit during and immediately after acute illnesses. Bone marrow suppression occurs in almost everyone during acute and chronic inflammatory illnesses. In patients with short RBC life spans, suppression can affect RBC counts more dramatically. These hypoplastic crises are usually transient and self-correcting before intervention is required.

Aplastic crisis refers to a profound cessation of erythroid activity in patients with chronic hemolytic anemias. It is associated with a rapidly falling hematocrit. Episodes are usually self-limited. Aplastic crises are caused by infection with a particular strain of parvovirus, B19A. Children infected with this virus usually develop permanent immunity. Aplastic crises do not often recur and are rarely seen in adults. Management requires close monitoring of the hematocrit and reticulocyte count. If anemia becomes symptomatic, transfusion support is indicated. Most crises resolve spontaneously within 1–2 weeks.

128 Megaloblastic Anemias

A. Victor Hoffbrand

The megaloblastic anemias are a group of disorders characterized by the presence of distinctive morphologic appearances of the developing red cells in the bone marrow. The marrow is usually hypercellular and the anemia is based on ineffective erythropoiesis. The cause is usually a deficiency of either cobalamin (vitamin B₁₂) or folate, but megaloblastic anemia may occur because of genetic or acquired abnormalities that affect the metabolism of these vitamins or because of defects in DNA synthesis not related to cobalamin or folate (Table 128-1). Cobalamin and folate absorption and metabolism are described next, followed by the biochemical basis, clinical and laboratory features, causes, and treatment of megaloblastic anemia.

COBALAMIN

Cobalamin (vitamin B₁₂) exists in a number of different chemical forms. All have a cobalt atom at the center of a corrin ring. In nature, the vitamin is mainly in the 2-deoxyadenosyl (ado) form, which is located in mitochondria. It is the cofactor for the enzyme methylmalonyl coenzyme A (CoA) mutase. The other major natural cobalamin is methylcobalamin, the form in human plasma and in cell cytoplasm. It is the cofactor for methionine synthase. There are also minor amounts of hydroxocobalamin to which methyl- and adocobalamin are converted rapidly by exposure to light.

DIETARY SOURCES AND REQUIREMENTS

Cobalamin is synthesized solely by microorganisms. Ruminants obtain cobalamin from the foregut, but the only source for humans is food of animal origin, e.g., meat, fish, and dairy products. Vegetables, fruits, and other foods of nonanimal origin are free from cobalamin unless they are contaminated by bacteria. A normal Western diet contains 5–30 µg of cobalamin daily. Adult daily losses (mainly in the urine and feces) are 1–3 µg (~0.1% of body stores), and because the body does not have the ability to degrade cobalamin, daily requirements are also about 1–3 µg. Body stores are of the order of 2–3 mg, sufficient for 3–4 years if supplies are completely cut off.

ABSORPTION

Two mechanisms exist for cobalamin absorption. One is passive, occurring equally through buccal, duodenal, and ileal mucosa; it is rapid but extremely inefficient, with <1% of an oral dose being absorbed by this process. The normal physiologic mechanism is active; it occurs through the ileum and is efficient for small (a few micrograms) oral doses of cobalamin, and it is mediated by gastric intrinsic factor (IF). Dietary cobalamin is released from protein complexes by enzymes in the stomach, duodenum, and jejunum; it combines rapidly with a salivary glycoprotein that belongs to the family of cobalamin-binding proteins known as haptocorrins (HCs). In the intestine, the haptocorrin is digested by pancreatic trypsin and the cobalamin is transferred to IF.

TABLE 128-1 CAUSES OF MEGALOBLASTIC ANEMIA

Cobalamin deficiency or abnormalities of cobalamin metabolism (see Tables 128-3, 128-4)
Folate deficiency or abnormalities of folate metabolism (see Table 128-5)
Therapy with antifolate drugs (e.g., methotrexate)
Independent of either cobalamin or folate deficiency and refractory to cobalamin and folate therapy:
Some cases of acute myeloid leukemia, myelodysplasia
Therapy with drugs interfering with synthesis of DNA (e.g., cytosine arabinoside, hydroxyurea, 6-mercaptopurine, azidothymidine [AZT])
Orotic aciduria (responds to uridine)
Thiamine-responsive

IF (gene at chromosome 11q13) is produced in the gastric parietal cells of the fundus and body of the stomach, and its secretion parallels that of hydrochloric acid. Normally, there is a vast excess of IF. The IF-cobalamin complex passes to the ileum, where IF attaches to a specific receptor (cubilin) on the microvillus membrane of the enterocytes. Cubilin also is present in yolk sac and renal proximal tubular epithelium. Cubilin appears to traffic by means of amnionless (AMN), an endocytic receptor protein that directs sublocalization and endocytosis of cubilin with its ligand IF-cobalamin complex. The cobalamin-IF complex enters the ileal cell, where IF is destroyed. After a delay of about 6 h, the cobalamin appears in portal blood attached to transcobalamin (TC) II.

Between 0.5 and 5 µg of cobalamin enter the bile each day. This binds to IF, and a major portion of biliary cobalamin normally is reabsorbed together with cobalamin derived from sloughed intestinal cells. Because of the appreciable amount of cobalamin undergoing enterohepatic circulation, cobalamin deficiency develops more rapidly in individuals who malabsorb cobalamin than it does in vegans, in whom reabsorption of biliary cobalamin is intact.

TRANSPORT

Two main cobalamin transport proteins exist in human plasma; they both bind cobalamin—one molecule for one molecule. One HC, also known as TC I, is closely related to other cobalamin-binding HCs in milk, gastric juice, bile, saliva, and other fluids. The gene *TCN1* is at chromosome 11q11-q12.3. These HCs differ from each other only in the carbohydrate moiety of the molecule. TC I is derived primarily from the specific granules in neutrophils. Normally, it is about two-thirds saturated with cobalamin, which it binds tightly. TC I does not enhance cobalamin entry into tissues. Glycoprotein receptors on liver cells are involved in the removal of TC I from plasma, and TC I may play a role in the transport of cobalamin analogues (which it binds more effectively than IF) to the liver for excretion in bile.

The other major cobalamin transport protein in plasma is transcobalamin, also known as TC II. The gene is on chromosome 22q11-q13.1. As for IF and HC, there are nine exons. The three proteins are likely to have a common ancestral origin. TC II is synthesized by liver and by other tissues, including macrophages, ileum, and vascular endothelium. It normally carries only 20–60 ng of cobalamin per liter of plasma and readily gives up cobalamin to marrow, placenta, and other tissues, which it enters by receptor-mediated endocytosis involving the TC II receptor and megalin (encoded by the *LRP-2* gene). The TC II cobalamin is internalized by endocytosis via clathrin-coated pits; the complex is degraded, but the receptor probably is recycled to the cell membrane as is the case for transferrin. Export of “free” cobalamin is via the ATP-binding cassette drug transporter alias multidrug resistance protein 1.

FOLATE

DIETARY FOLATE

Folic (pteroylglutamic) acid is a yellow, crystalline, water-soluble substance. It is the parent compound of a large family of natural folate compounds, which differ from it in three respects: (1) they are partly or completely reduced to di- or tetrahydrofolate (THF) derivatives, (2) they usually contain a single carbon unit (Table 128-2), and (3) 70–90% of natural folates are folate-polyglutamates.

Most foods contain some folate. The highest concentrations are found in liver, yeast, spinach, other greens, and nuts (>100 µg/100 g). The total folate content of an average Western diet is ~250 µg daily, but the amount varies widely according to the type of food eaten and the method of cooking. Folate is easily destroyed by heating, particularly in large volumes of water. Total body folate in the adult is ~10 mg, with the liver containing the largest store. Daily adult requirements are ~100 µg, and so stores are sufficient for only 3–4 months in normal adults and severe folate deficiency may develop rapidly.

ABSORPTION

Folates are absorbed rapidly from the upper small intestine. The absorption of folate polyglutamates is less efficient than that of monoglutamates; on average, ~50% of food folate is absorbed. Polyglutamate

TABLE 128-2 BIOCHEMICAL REACTIONS OF FOLATE COENZYMES

Reaction	Coenzyme Form of Folate Involved	Single Carbon Unit Transferred	Importance
<i>Formate activation</i>	THF	–CHO	Generation of 10-formyl-THF
<i>Purine synthesis</i>			
Formation of glycinamide ribonucleotide	5,10-Methylene-THF	–CHO	Formation of purines needed for DNA, RNA synthesis, but reactions probably not rate-limiting
Formylation of aminoimidazole carboxamide ribonucleotide (AICAR)	10-Formyl (CHO)THF		
<i>Pyrimidine synthesis</i>			
Methylation of deoxyuridine monophosphate (dUMP) to thymidine monophosphate (dTMP)	5,10-Methylene-THF	–CH ₃	Rate limiting in DNA synthesis Oxidizes THF to DHF Some breakdown of folate at the C-9–N-10 bond
<i>Amino acid interconversion</i>			
Serine–glycine interconversion	THF	=CH ₂	Entry of single carbon units into active pool
Homocysteine to methionine	5-Methyl(M)THF	–CH ₃	Demethylation of 5-MTHF to THF; also requires cobalamin, flavine adenine dinucleotide, ATP, and adenosylmethionine
Forminoglutamic acid to glutamic acid in histidine catabolism	THF	–HN–CH=	

Abbreviations: DHF, dihydrofolate; THF, tetrahydrofolate.

forms are hydrolyzed to the monoglutamate derivatives either in the lumen of the intestine or within the mucosa. All dietary folates are converted to 5-methylTHF (5-MTHF) within the small intestinal mucosa before entering portal plasma. The monoglutamates are actively transported across the enterocyte by a proton-coupled folate transporter (PCFT, SCL46A1). This is situated at the apical brush border and is most active at pH 5.5, which is about the pH of the duodenal and jejunal surface. Genetic mutations of this protein underlie hereditary malabsorption of folate (see below). Pteroylglutamic acid at doses >400 µg is absorbed largely unchanged and converted to natural folates in the liver. Lower doses are converted to 5-MTHF during absorption through the intestine.

About 60–90 µg of folate enters the bile each day and is excreted into the small intestine. Loss of this folate, together with the folate of sloughed intestinal cells, accelerates the speed with which folate deficiency develops in malabsorption conditions.

TRANSPORT

Folate is transported in plasma; about one-third is loosely bound to albumin, and two-thirds is unbound. In all body fluids (plasma, cerebrospinal fluid, milk, bile), folate is largely, if not entirely, 5-MTHF in the monoglutamate form. Three types of folate-binding protein are involved. A reduced folate transporter (RFC, SLC19A1) is the major route of delivery of plasma folate (5-MTHF) to cells. Two folate receptors, FR2 and FR3 embedded in the cell membrane by a glycosyl phosphatidylinositol anchor, transport folate into the cell via receptor-mediated endocytosis. The third protein, PCFT, transports folate at low pH from the vesicle to the cell cytoplasm. The reduced folate transporter also mediates uptake of methotrexate by cells.

BIOCHEMICAL FUNCTIONS

Folates (as the intracellular polyglutamate derivatives) act as coenzymes in the transfer of single-carbon units (Fig. 128-1 and Table 128-2). Two of these reactions are involved in purine synthesis and one in pyrimidine synthesis necessary for DNA and RNA replication. Folate is also a coenzyme for methionine synthesis, in which methylcobalamin is also involved and in which THF is regenerated. THF is the acceptor of single carbon units newly entering the active pool via conversion of serine to

glycine. Methionine, the other product of the methionine synthase reaction, is the precursor for S-adenosylmethionine (SAM), the universal methyl donor involved in >100 methyltransferase reactions (Fig. 128-1).

During thymidylate synthesis, 5,10-methylene-THF is oxidized to DHF (dihydrofolate). The enzyme DHF reductase converts this to THF. The drugs methotrexate, pyrimethamine, and (mainly in bacteria) trimethoprim inhibit DHF reductase and so prevent formation of active THF coenzymes from DHF. A small fraction of the folate coenzyme is not recycled during thymidylate synthesis but is degraded at the C9–N10 bond.

BIOCHEMICAL BASIS OF MEGALOBlastic ANEMIA

The common feature of all megaloblastic anemias is a defect in DNA synthesis that affects rapidly dividing cells in the bone marrow. All conditions that give rise to megaloblastic changes have in common a disparity in the rate of synthesis or availability of the four immediate precursors of DNA: the deoxyribonucleoside triphosphates (dNTPs)—dA(adenine)TP and dG(guanine)TP (purines), dT(thymine)TP and dC(cytosine)TP (pyrimidines). In deficiencies of either folate or cobalamin, there is failure to convert deoxyuridine monophosphate (dUMP) to deoxythymidine monophosphate (dTMP), the precursor of dTTP (Fig 128-1). This is the case because folate is needed as the coenzyme 5,10-methylene-THF polyglutamate for conversion of dUMP to dTMP; the availability of 5,10-methylene-THF is reduced in either cobalamin or folate deficiency. An alternative theory for megaloblastic anemia in cobalamin or folate deficiency is misincorporation of uracil into DNA because of a buildup of deoxyuridine triphosphate (dUTP) at the DNA replication fork as a consequence of the block in conversion of dUMP to dTMP.

COBALAMIN-FOLATE RELATIONS

Folate is required for many reactions in mammalian tissues. Only two reactions in the body are known to require cobalamin. Methylmalonyl CoA isomerization requires adocobalamin, and the methylation of homocysteine to methionine requires both methylcobalamin and 5-MTHF (Fig. 128-1). This reaction is the first step in the pathway by which 5-MTHF, which enters bone marrow and other cells from plasma, is converted into all the intracellular folate coenzymes. The coenzymes are all polyglutamated (the larger size aiding retention in the cell), but the enzyme folate polyglutamate synthase can use only THF, not MTHF, as substrate. In cobalamin deficiency, MTHF accumulates in plasma, and intracellular folate concentrations fall due to failure of formation of THF, the substrate on which folate polyglutamates are built. This has been termed *THF starvation*, or the *methylfolate trap*.

This theory explains the abnormalities of folate metabolism that occur in cobalamin deficiency (high serum folate, low cell folate, positive purine precursor aminoimidazole carboxamide ribonucleotide [AICAR] excretion; Table 128-2) and also why the anemia of cobalamin deficiency responds to folic acid in large doses.

CLINICAL FEATURES

Many symptomless patients are detected through the finding of a raised mean corpuscular volume (MCV) on a routine blood count. The main clinical features in more severe cases are those of anemia. Anorexia is

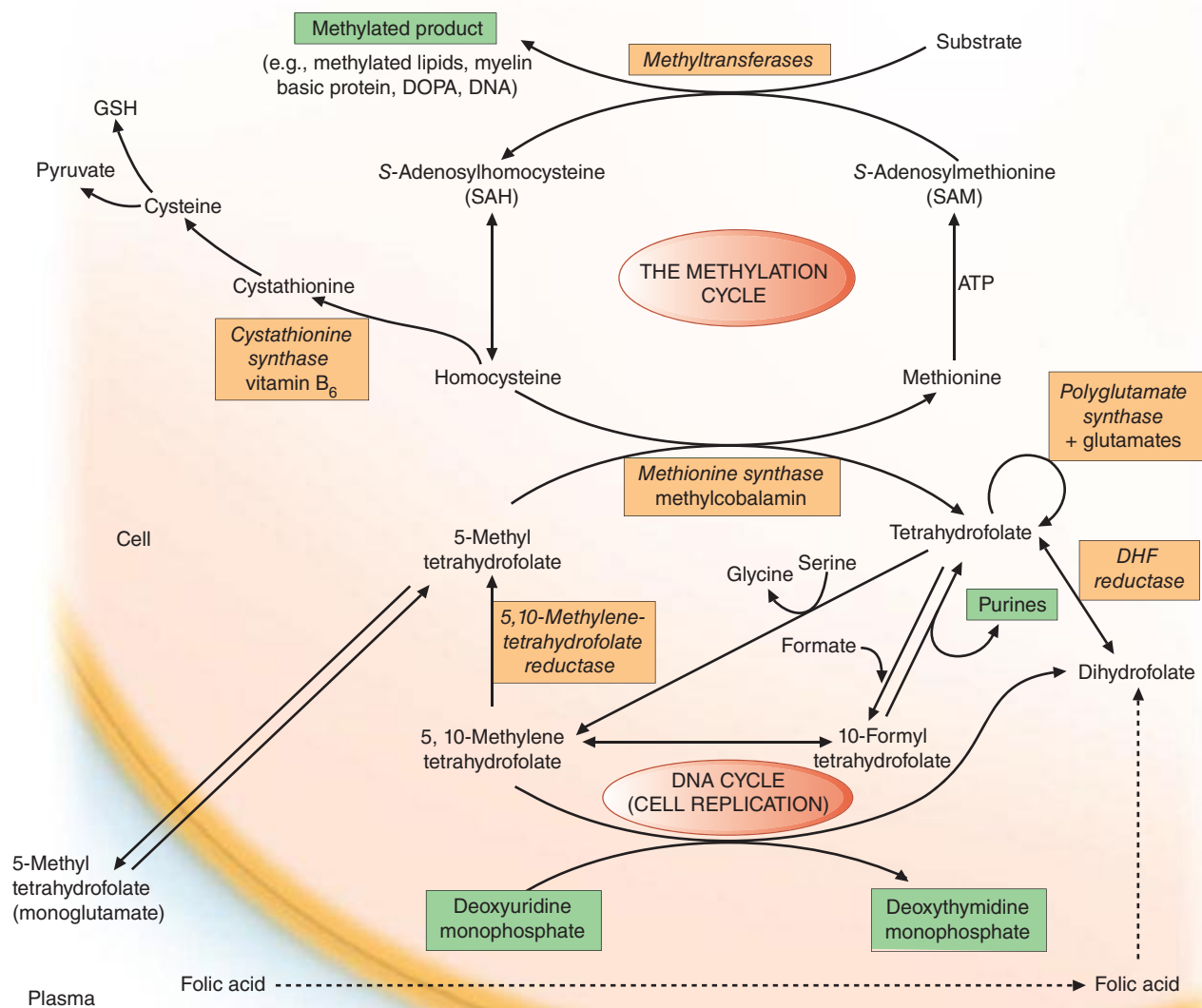


FIGURE 128-1 The role of folates in DNA synthesis and in formation of S-adenosylmethionine (SAM), which is involved in numerous methylation reactions. DHF, dihydrofolate; GSH, glutathione. (Reprinted from AV Hoffbrand et al [eds]: Postgraduate Haematology, 5th ed. Oxford, UK, Blackwell Publishing, 2005; with permission.)

usually marked, and there may be weight loss, diarrhea, or constipation. Glossitis, angular cheilosis, a mild fever in more severely anemic patients, jaundice (unconjugated), and reversible melanin skin hyperpigmentation also may occur with a deficiency of either folate or cobalamin. Thrombocytopenia sometimes leads to bruising, and this may be aggravated by vitamin C deficiency or alcohol in malnourished patients. The anemia and low leukocyte count may predispose to infections, particularly of the respiratory and urinary tracts. Cobalamin deficiency has also been associated with impaired bactericidal function of phagocytes.

GENERAL TISSUE EFFECTS OF COBALAMIN AND FOLATE DEFICIENCIES

Epithelial Surfaces After the marrow, the next most frequently affected tissues are the epithelial cell surfaces of the mouth, stomach, and small intestine and the respiratory, urinary, and female genital tracts. The cells show macrocytosis, with increased numbers of multinucleate and dying cells. The deficiencies may cause cervical smear abnormalities.

Complications of Pregnancy The gonads are also affected, and infertility is common in both men and women with either deficiency. Maternal folate deficiency has been implicated as a cause of prematurity, and both folate deficiency and cobalamin deficiency have been implicated in recurrent fetal loss and neural tube defects, as discussed below.

Neural Tube Defects Folic acid supplements at the time of conception and in the first 12 weeks of pregnancy reduce by ~70% the incidence of neural tube defects (NTDs) (anencephaly, meningomyelocele, encephalocele, and spina bifida) in the fetus. Most of this protective effect can be achieved by taking folic acid, 0.4 mg daily, at the time of conception.

The incidence of cleft palate and harelip also can be reduced by prophylactic folic acid. There is no clear simple relationship between maternal folate status and these fetal abnormalities, although overall the lower the maternal folate, the greater the risk to the fetus. NTDs also can be caused by antifolate and antiepileptic drugs.

An underlying maternal folate metabolic abnormality has also been postulated. One abnormality has been identified: reduced activity of the enzyme 5,10-methylene-THF reductase (MTHFR) (Fig. 128-1) caused by a common C677T polymorphism in the *MTHFR* gene. In one study, the prevalence of this polymorphism was found to be higher than in controls in the parents of NTD fetuses and in the fetuses themselves: homozygosity for the TT mutation was found in 13% of cases compared with 5% of control subjects. The polymorphism codes for a thermolabile form of MTHFR. The homozygous state results in a lower mean serum and red cell folate level compared with control subjects, as well as significantly higher serum homocysteine levels. Tests for mutations in other enzymes possibly associated with NTDs,

e.g., methionine synthase and serine-glycine hydroxymethylase, have been negative. Serum vitamin B₁₂ levels are also lower in the sera of mothers of NTD infants than in controls. In addition, maternal TC II receptor polymorphisms are associated with increased risk of NTD births. There are, however, no studies showing dietary fortification with vitamin B₁₂ reduces the incidence of NTDs.

Cardiovascular Disease Children with severe homocystinuria (blood levels ≥ 100 $\mu\text{mol/L}$) due to deficiency of one of three enzymes, methionine synthase, MTHFR, or cystathionine synthase (Fig. 128-1), have vascular disease, e.g., ischemic heart disease, cerebrovascular disease, or pulmonary embolus, as teenagers or in young adulthood. Lesser degrees of raised serum homocysteine and low levels of serum folate and homozygous inherited mutations of MTHFR have been found to be associated with cerebrovascular, peripheral vascular, and coronary heart disease and with deep vein thrombosis. Prospective randomized trials of lowering homocysteine levels with supplements of folic acid, vitamin B₁₂, and vitamin B₆ against placebo over a 5-year period in patients with vascular disease or diabetes have not, however, shown a reduction of first event fatal or nonfatal myocardial infarction, nor have these supplements reduced the risk of recurrent cardiovascular disease after an acute myocardial infarct. Meta-analysis showed an 18% reduction in strokes but no significant prevention of death from any cause. Venous thrombosis has been reported to be more frequent in vitamin B₁₂-deficient subjects than in controls. This was ascribed to raised plasma homocysteine levels in vitamin B₁₂ deficiency.

Malignancy Prophylactic folic acid in pregnancy has been found in some but not all studies to reduce the subsequent incidence of acute lymphoblastic leukemia (ALL) in childhood. A significant negative association has also been found with the MTHFR C677T polymorphism and leukemias with mixed lineage leukemia (MLL) translocations, but a positive association with hyperdiploidy in infants with ALL or acute myeloid leukemia or with childhood ALL. A second polymorphism in the MTHFR gene, A1298C, is also strongly associated with hyperdiploid leukemia. There are various positive and negative associations between polymorphisms in folate-dependent enzymes and the incidence of adult ALL. The C677T polymorphism is thought to lead to increased thymidine pools and “better quality” of DNA synthesis by shunting one-carbon groups toward thymidine and purine synthesis. This may explain its reported association with a lower risk for colorectal cancer. Most but not all studies suggest that prophylactic folic acid also protects against colon adenomas. Other tumors that have been associated with folate polymorphisms or status include follicular lymphoma, breast cancer, and gastric cancer. A meta-analysis of 50,000 individuals given folic acid or placebo in cardiovascular or colon adenoma prevention trials found that folic acid supplementation did not substantially increase or decrease the incidence of site-specific cancer during the first 5 years of treatment. Because folic acid may “feed” tumors, it probably should be avoided in those with established tumors unless there is severe megaloblastic anemia due to folate deficiency.

Neurologic Manifestations Cobalamin deficiency may cause a bilateral peripheral neuropathy or degeneration (demyelination) of the posterior and pyramidal tracts of the spinal cord and, less frequently, optic atrophy or cerebral symptoms.

The patient, more frequently male, presents with paresthesias, muscle weakness, or difficulty in walking and sometimes dementia, psychotic disturbances, or visual impairment. Long-term nutritional cobalamin deficiency in infancy leads to poor brain development and impaired intellectual development. Folate deficiency has been suggested to cause organic nervous disease, but this is uncertain, although methotrexate injected into the cerebrospinal fluid may cause brain or spinal cord damage.

An important clinical problem is the nonanemic patient with neurologic or psychiatric abnormalities and a low or borderline serum cobalamin level. In such patients, it is necessary to try to establish whether there is significant cobalamin deficiency, e.g., by careful examination of the blood film, tests for serum gastrin level and for antibodies to IF or parietal cells, along with serum methylmalonic acid

(MMA) measurement if available. A trial of cobalamin therapy for at least 3 months will usually also be needed to determine whether the symptoms improve.

The biochemical basis for cobalamin neuropathy remains obscure. Its occurrence in the absence of methylmalonic aciduria in TC II deficiency suggests that the neuropathy is related to the defect in homocysteine-methionine conversion. Accumulation of S-adenosylhomocysteine in the brain, resulting in inhibition of transmethylation reactions, has been suggested.

Psychiatric disturbance is common in both folate and cobalamin deficiencies. This, like the neuropathy, has been attributed to a failure of the synthesis of SAM, which is needed in methylation of biogenic amines (e.g., dopamine) as well as that of proteins, phospholipids, and neurotransmitters in the brain (Fig. 128-1). Associations between lower serum folate or cobalamin levels and higher homocysteine levels and the development of decreased cognitive function and dementia in Alzheimer’s disease have been reported. A meta-analysis of randomized, placebo-controlled trials of homocysteine-lowering B-vitamin supplementation of individuals with and without cognitive impairment, however, showed that supplementation with vitamin B₁₂, vitamin B₆, and folic acid alone or in combination did not improve cognitive function. It is unknown whether prolonged treatment with these B vitamins can reduce the risk of dementia in later life.

HEMATOLOGIC FINDINGS

PERIPHERAL BLOOD

Oval macrocytes, usually with considerable anisocytosis and poikilocytosis, are the main feature (Fig. 128-2A). The MCV is usually >100 fL unless a cause of microcytosis (e.g., iron deficiency or thalassemia trait) is present. Some of the neutrophils are hypersegmented (more than five nuclear lobes). There may be leukopenia due to a reduction in granulocytes and lymphocytes, but this is usually $>1.5 \times 10^9/\text{L}$; the platelet count may be moderately reduced, rarely to $<40 \times 10^9/\text{L}$. The severity of all these changes parallels the degree of anemia. In a non-anemic patient, the presence of a few macrocytes and hypersegmented neutrophils in the peripheral blood may be the only indication of the underlying disorder.

BONE MARROW

In a severely anemic patient, the marrow is hypercellular with an accumulation of primitive cells due to selective death by apoptosis of more mature forms. The erythroblast nucleus maintains a primitive appearance despite maturation and hemoglobinization of the cytoplasm. The cells are larger than normoblasts, and an increased number of cells with eccentric lobulated nuclei or nuclear fragments may be present (Fig. 128-2B). Giant and abnormally shaped metamyelocytes and enlarged hyperpolyploid megakaryocytes are characteristic. In severe cases, the accumulation of primitive cells may mimic acute myeloid leukemia, whereas in less anemic patients, the changes in the marrow may be difficult to recognize. The terms *intermediate*, *mild*, and *early* have been used. The term *megaloblastoid* does not mean mildly megaloblastic. It is used to describe cells with both immature-appearing nuclei and defective hemoglobinization and is usually seen in myelodysplasia.

CHROMOSOMES

Bone marrow cells, transformed lymphocytes, and other proliferating cells in the body show a variety of changes, including random breaks, reduced contraction, spreading of the centromere, and exaggeration of secondary chromosomal constrictions and overprominent satellites. Similar abnormalities may be produced by antimetabolite drugs (e.g., cytosine arabinoside, hydroxyurea, and methotrexate) that interfere with either DNA replication or folate metabolism and that also cause megaloblastic appearances.

INEFFECTIVE HEMATOPOIESIS

There is an accumulation of unconjugated bilirubin in plasma due to the death of nucleated red cells in the marrow (ineffective erythropoiesis). Other evidence for this includes raised urine urobilinogen, reduced haptoglobins and positive urine hemosiderin, and a raised

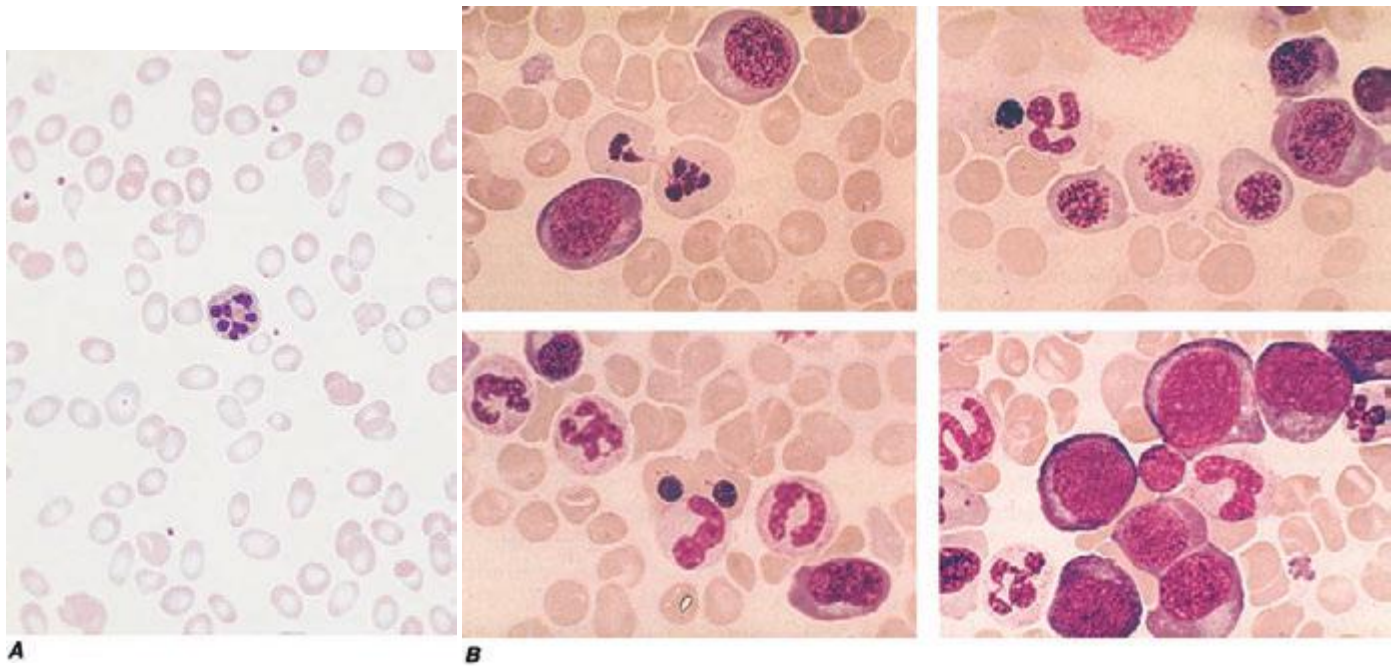


FIGURE 128-2 **A.** The peripheral blood in severe megaloblastic anemia. **B.** The bone marrow in severe megaloblastic anemia. (Reprinted from AV Hoffbrand et al [eds]: *Postgraduate Haematology*, 5th ed. Oxford, UK, Blackwell Publishing, 2005; with permission.)

serum lactate dehydrogenase. A weakly positive direct antiglobulin test due to complement can lead to a false diagnosis of autoimmune hemolytic anemia.

CAUSES OF COBALAMIN DEFICIENCY

Cobalamin deficiency is usually due to malabsorption. The only other cause is inadequate dietary intake.

INADEQUATE DIETARY INTAKE

Adults Dietary cobalamin deficiency arises in vegans who omit meat, fish, eggs, cheese, and other animal products from their diet. The largest group in the world consists of Hindus, and it is likely that many millions of Indians are at risk of deficiency of cobalamin on a nutritional basis. Subnormal serum cobalamin levels are found in up to 50% of randomly selected, young, adult Indian vegans, but the deficiency usually does not progress to megaloblastic anemia since the diet of most vegans is not totally lacking in cobalamin and the enterohepatic circulation of cobalamin is intact. Dietary cobalamin deficiency may also arise rarely in nonvegetarian individuals who exist on grossly inadequate diets because of poverty or psychiatric disturbance.

Infants Cobalamin deficiency has been described in infants born to severely cobalamin-deficient mothers. These infants develop megaloblastic anemia at about 3–6 months of age, presumably because they are born with low stores of cobalamin and because they are fed breast milk with low cobalamin content. The babies have also shown growth retardation, impaired psychomotor development, and other neurologic sequelae.

GASTRIC CAUSES OF COBALAMIN MALABSORPTION

See Tables 128-3 and 128-4.

Pernicious Anemia Pernicious anemia (PA) may be defined as a severe lack of IF due to gastric atrophy. It is a common disease in north Europeans but occurs in all countries and ethnic groups. The overall incidence is about 120 per 100,000 population in the United Kingdom (UK). The ratio of incidence in men and women among whites is ~1:1.6, and the peak age of onset is 60 years, with only 10% of patients being <40 years of age. However, in some ethnic groups, notably black individuals and Latin Americans, the age at onset of PA is generally lower. The disease occurs more commonly than by chance

TABLE 128-3 CAUSES OF COBALAMIN DEFICIENCY SUFFICIENTLY SEVERE TO CAUSE MEGALOBLASTIC ANEMIA

Nutritional	Vegans
Malabsorption	Pernicious anemia
Gastric causes	Congenital absence of intrinsic factor or functional abnormality Total or partial gastrectomy
Intestinal causes	Intestinal stagnant loop syndrome: jejunal diverticulosis, ileocolic fistula, anatomic blind loop, intestinal stricture, etc. Ileal resection and Crohn's disease Selective malabsorption with proteinuria Tropical sprue Transcobalamin II deficiency Fish tapeworm

TABLE 128-4 MALABSORPTION OF COBALAMIN MAY OCCUR IN THE FOLLOWING CONDITIONS BUT IS NOT USUALLY SUFFICIENTLY SEVERE AND PROLONGED TO CAUSE MEGALOBLASTIC ANEMIA

Gastric causes	Simple atrophic gastritis (food cobalamin malabsorption) Zollinger-Ellison syndrome Gastric bypass surgery Use of proton pump inhibitors
Intestinal causes	Gluten-induced enteropathy Severe pancreatitis HIV infection Radiotherapy Graft-versus-host disease
Deficiencies of cobalamin, folate, protein, ?riboflavin, ?nicotinic acid	
Therapy with colchicine, para-aminosalicylate, neomycin, slow-release potassium chloride, anticonvulsant drugs, metformin, phenformin, cytotoxic drugs	
Alcohol	

in close relatives and in persons with other organ-specific autoimmune diseases, e.g., thyroid diseases, vitiligo, hypoparathyroidism, and Addison's disease. It is also associated with hypogammaglobulinemia, with premature graying or blue eyes, and persons of blood group A. An association with human leukocyte antigen (HLA) 3 has been reported in some but not all series and, in those with endocrine disease, with HLA-B8, -B12, and -BW15. Life expectancy is normal in women once regular treatment has begun. Men have a slightly subnormal life expectancy as a result of a higher incidence of carcinoma of the stomach than in control subjects. Gastric output of hydrochloric acid, pepsin, and IF is severely reduced. The serum gastrin level is raised, and serum pepsinogen I levels are low.

Gastric Biopsy A single endoscopic examination is recommended if PA is diagnosed. Gastric biopsy usually shows atrophy of all layers of the body and fundus, with loss of glandular elements, an absence of parietal and chief cells and replacement by mucous cells, a mixed inflammatory cell infiltrate, and perhaps intestinal metaplasia. The infiltrate of plasma cells and lymphocytes contains an excess of CD4 cells. These are directed against gastric H/K-ATPase. The antral mucosa is usually well preserved. *Helicobacter pylori* infection occurs infrequently in PA, but it has been suggested that *H. pylori* gastritis occurs at an early phase of atrophic gastritis and presents in younger patients as iron-deficiency anemia but in older patients as PA. *H. pylori* is suggested to stimulate an autoimmune process directed against parietal cells, with the *H. pylori* infection then being gradually replaced, in some individuals, by an autoimmune process.

Serum Antibodies Two types of IF immunoglobulin G antibody may be found in the sera of patients with PA. One, the "blocking," or type I, antibody, prevents the combination of IF and cobalamin, whereas the "binding," or type II, antibody prevents attachment of IF to ileal mucosa. Type I occurs in the sera of ~55% of patients, and type II in 35%. IF antibodies cross the placenta and may cause temporary IF deficiency in a newborn infant. Patients with PA also show cell-mediated immunity to IF. Type I antibody has been detected rarely in the sera of patients without PA but with thyrotoxicosis, myxedema, Hashimoto's disease, or diabetes mellitus and in relatives of PA patients. IF antibodies also have been detected in gastric juice in ~80% of PA patients. These gastric antibodies may reduce absorption of dietary cobalamin by combining with small amounts of remaining IF.

Parietal cell antibody is present in the sera of almost 90% of adult patients with PA but is frequently present in other subjects. Thus, it occurs in as many as 16% of randomly selected female subjects age >60 years. The parietal cell antibody is directed against the α and β subunits of the gastric proton pump (H^+,K^+ -ATPase).

JUVENILE PERNICIOUS ANEMIA

This usually occurs in older children and resembles PA of adults. Gastric atrophy, achlorhydria, and serum IF antibodies are all present, although parietal cell antibodies are usually absent. About one-half of these patients show an associated endocrinopathy such as autoimmune thyroiditis, Addison's disease, or hypoparathyroidism; in some, mucocutaneous candidiasis occurs.

CONGENITAL INTRINSIC FACTOR DEFICIENCY OR FUNCTIONAL ABNORMALITY

An affected child usually presents with megaloblastic anemia in the first to third year of life; a few have presented as late as the second decade. The child usually has no demonstrable IF but has a normal gastric mucosa and normal secretion of acid. The inheritance is autosomal recessive. Parietal cell and IF antibodies are absent. Variants have been described in which the child is born with IF that can be detected immunologically but is unstable or functionally inactive, unable to bind cobalamin or to facilitate its uptake by ileal receptors.

GASTRECTOMY

After total gastrectomy, cobalamin deficiency is inevitable, and prophylactic cobalamin therapy should be commenced immediately after the operation. After partial gastrectomy, 10–15% of patients also develop this deficiency. The exact incidence and time of onset are

most influenced by the size of the resection and the preexisting size of cobalamin body stores.

FOOD COBALAMIN MALABSORPTION

Failure of release of cobalamin from binding proteins in food is believed to be responsible for this condition, which is more common in the elderly. It is associated with low serum cobalamin levels, with or without raised serum levels of MMA and homocysteine. Typically, these patients have normal cobalamin absorption, as measured with crystalline cobalamin, but show malabsorption when a modified test using food-bound cobalamin is used. The frequency of progression to severe cobalamin deficiency and the reasons for this progression are not clear.

INTESTINAL CAUSES OF COBALAMIN MALABSORPTION

Intestinal Stagnant Loop Syndrome Malabsorption of cobalamin occurs in a variety of intestinal lesions in which there is colonization of the upper small intestine by fecal organisms. This may occur in patients with jejunal diverticulosis, enteroanastomosis, or an intestinal stricture or fistula or with an anatomic blind loop due to Crohn's disease, tuberculosis, or an operative procedure.

Ileal Resection Removal of ≥ 1.2 m of terminal ileum causes malabsorption of cobalamin. In some patients after ileal resection, particularly if the ileocecal valve is incompetent, colonic bacteria may contribute further to the onset of cobalamin deficiency.

Selective Malabsorption of Cobalamin with Proteinuria (Imlerslund's Syndrome; Imlerslund-Gräsbeck Syndrome; Congenital Cobalamin Malabsorption; Autosomal Recessive Megaloblastic Anemia; MGA1) This autosomally recessive disease is the most common cause of megaloblastic anemia due to cobalamin deficiency in infancy in Western countries. More than 200 cases have been reported, with familial clusters in Finland, Norway, the Middle East, and North Africa. The patients secrete normal amounts of IF and gastric acid but are unable to absorb cobalamin. In Finland, impaired synthesis, processing, or ligand binding of cubilin due to inherited mutations is found. In Norway, mutation of the gene for AMN has been reported. Other tests of intestinal absorption are normal. Over 90% of these patients show nonspecific proteinuria, but renal function is otherwise normal and renal biopsy has not shown any consistent renal defect. A few have shown aminoaciduria and congenital renal abnormalities, such as duplication of the renal pelvis.

Tropical Sprue Nearly all patients with acute and subacute tropical sprue show malabsorption of cobalamin; this may persist as the principal abnormality in the chronic form of the disease, when the patient may present with megaloblastic anemia or neuropathy due to cobalamin deficiency. Absorption of cobalamin usually improves after antibiotic therapy and, in the early stages, folic acid therapy.

Fish Tapeworm Infestation The fish tapeworm (*Diphyllobothrium latum*) lives in the small intestine of humans and accumulates cobalamin from food, rendering the cobalamin unavailable for absorption. Individuals acquire the worm by eating raw or partly cooked fish. Infestation is common around the lakes of Scandinavia, Germany, Japan, North America, and Russia. Megaloblastic anemia or cobalamin neuropathy occurs only in those with a heavy infestation.

Gluten-Induced Enteropathy Malabsorption of cobalamin occurs in ~30% of untreated patients (presumably those in whom the disease extends to the ileum). Cobalamin deficiency is not severe in these patients and is corrected with a gluten-free diet.

Severe Chronic Pancreatitis In this condition, lack of trypsin is thought to cause dietary cobalamin attached to gastric non-IF (R) binder to be unavailable for absorption. It also has been proposed that in pancreatitis, the concentration of calcium ions in the ileum falls below the level needed to maintain normal cobalamin absorption.

HIV Infection Serum cobalamin levels tend to fall in patients with HIV infection and are subnormal in 10–35% of those with AIDS.

646 Malabsorption of cobalamin not corrected by IF has been shown in some, but not all, patients with subnormal serum cobalamin levels. Cobalamin deficiency sufficiently severe to cause megaloblastic anemia or neuropathy is rare.

Zollinger-Ellison Syndrome Malabsorption of cobalamin has been reported in the Zollinger-Ellison syndrome. It is thought that there is a failure to release cobalamin from R-binding protein due to inactivation of pancreatic trypsin by high acidity, as well as interference with IF binding of cobalamin.

Radiotherapy Both total-body irradiation and local radiotherapy to the ileum (e.g., as a complication of radiotherapy for carcinoma of the cervix) may cause malabsorption of cobalamin.

Graft-versus-Host Disease This commonly affects the small intestine. Malabsorption of cobalamin due to abnormal gut flora, as well as damage to ileal mucosa, is common.

Drugs The drugs that have been reported to cause malabsorption of cobalamin are listed in Table 105-4. However, megaloblastic anemia due to these drugs is rare.

ABNORMALITIES OF COBALAMIN METABOLISM

Congenital Transcobalamin II Deficiency or Abnormality Infants with TC II deficiency usually present with megaloblastic anemia within a few weeks of birth. Serum cobalamin and folate levels are normal, but the anemia responds to massive (e.g., 1 mg three times weekly) injections of cobalamin. Some cases show neurologic complications. The protein may be present but functionally inert. Genetic abnormalities found include mutations of an intra-exonic cryptic splice site, extensive deletion, single nucleotide deletion, nonsense mutation, and an RNA editing defect. Malabsorption of cobalamin occurs in all cases, and serum immunoglobulins are usually reduced. Failure to institute adequate cobalamin therapy or treatment with folic acid may lead to neurologic damage.

Congenital Methylmalonic Acidemia and Aciduria Infants with this abnormality are ill from birth with vomiting, failure to thrive, severe metabolic acidosis, ketosis, and mental retardation. Anemia, if present, is normocytic and normoblastic. The condition may be due to a functional defect in either mitochondrial methylmalonyl CoA mutase or its cofactor adocobalamin. Mutations in the methylmalonyl CoA mutase are not responsive, or only poorly responsive, to treatment with cobalamin. A proportion of infants with failure of adocobalamin synthesis respond to cobalamin in large doses. Some children have combined methylmalonic aciduria and homocystinuria due to defective formation of both cobalamin coenzymes. This usually presents in the first year of life with feeding difficulties, developmental delay, microcephaly, seizures, hypotonia, and megaloblastic anemia.

Acquired Abnormality of Cobalamin Metabolism: Nitrous Oxide Inhalation Nitrous oxide (N_2O) irreversibly oxidizes methylcobalamin to an inactive precursor; this inactivates methionine synthase. Megaloblastic anemia has occurred in patients undergoing prolonged N_2O anesthesia (e.g., in intensive care units). A neuropathy resembling cobalamin neuropathy has been described in dentists and anesthetists who are exposed repeatedly to N_2O . Methylmalonic aciduria does not occur as adocobalamin is not inactivated by N_2O .

CAUSES OF FOLATE DEFICIENCY

(Table 128-5)

NUTRITIONAL

Dietary folate deficiency is common. Indeed, in most patients with folate deficiency a nutritional element is present. Certain individuals are particularly prone to have diets containing inadequate amounts of folate (Table 128-5). In the United States and other countries where fortification of the diet with folic acid has been adopted, the prevalence of folate deficiency has dropped dramatically and is now almost restricted to high-risk groups with increased folate needs. Nutritional

TABLE 128-5 CAUSES OF FOLATE DEFICIENCY

Dietary ^a
Particularly in: old age, infancy, poverty, alcoholism, chronic invalids, and the psychiatrically disturbed; may be associated with scurvy or kwashiorkor
Malabsorption
Major causes of deficiency
Tropical sprue, gluten-induced enteropathy in children and adults, and in association with dermatitis herpetiformis, specific malabsorption of folate, intestinal megaloblastosis caused by severe cobalamin or folate deficiency
Minor causes of deficiency
Extensive jejunal resection, Crohn's disease, partial gastrectomy, congestive heart failure, Whipple's disease, scleroderma, amyloid, diabetic enteropathy, systemic bacterial infection, lymphoma, sulfasalazine (Salazopyrin)
Excess utilization or loss
Physiologic
Pregnancy and lactation, prematurity
Pathologic
Hematologic diseases: chronic hemolytic anemias, sickle cell anemia, thalassemia major, myelofibrosis
Malignant diseases: carcinoma, lymphoma, leukemia, myeloma
Inflammatory diseases: tuberculosis, Crohn's disease, psoriasis, exfoliative dermatitis, malaria
Metabolic disease: homocystinuria
Excess urinary loss: congestive heart failure, active liver disease
Hemodialysis, peritoneal dialysis
Antifolate drugs ^b
Anticonvulsant drugs (phenytoin, primidone, barbiturates), sulfasalazine
Nitrofurantoin, tetracycline, antituberculosis (less well documented)
Mixed causes
Liver diseases, alcoholism, intensive care units

^aIn severely folate-deficient patients with causes other than those listed under Dietary, poor dietary intake is often present. ^bDrugs inhibiting dihydrofolate reductase are discussed in the text.

folate deficiency occurs in kwashiorkor and scurvy and in infants with repeated infections or those who are fed solely on goats' milk, which has a low folate content.

MALABSORPTION

Malabsorption of dietary folate occurs in tropical sprue and in gluten-induced enteropathy. In the rare congenital recessive syndrome of selective malabsorption of folate due to mutation of the proton-coupled folate transporter (PCFT), there is an associated defect of folate transport into the cerebrospinal fluid, and these patients show megaloblastic anemia, which responds to physiologic doses of folic acid given parenterally but not orally. They also show mental retardation, convulsions, and other central nervous system abnormalities. Minor degrees of malabsorption may also occur after jejunal resection or partial gastrectomy, in Crohn's disease, and in systemic infections, but in these conditions, if severe deficiency occurs, it is usually largely due to poor nutrition. Malabsorption of folate has been described in patients receiving sulfasalazine (Salazopyrin), cholestyramine, and triamterene.

EXCESS UTILIZATION OR LOSS

Pregnancy Folate requirements are increased by 200–300 μg to ~400 μg daily in a normal pregnancy, partly because of transfer of the vitamin to the fetus but mainly because of increased folate catabolism due to cleavage of folate coenzymes in rapidly proliferating tissues. Megaloblastic anemia due to this deficiency is prevented by prophylactic folic acid therapy. It occurred in 0.5% of pregnancies in the UK and other Western countries before prophylaxis with folic acid, but the incidence is much higher in countries where the general nutritional status is poor.

Prematurity A newborn infant, whether full term or premature, has higher serum and red cell folate concentrations than does an adult. However, a newborn infant's demand for folate has been estimated to be up to 10 times that of adults on a weight basis, and the neonatal folate level falls rapidly to the lowest values at about 6 weeks of age. The falls are steepest and are liable to reach subnormal levels in premature babies, a number of whom develop megaloblastic anemia responsive to folic acid at about 4–6 weeks of age. This occurs particularly in the smallest babies (<1500 g birth weight) and those who have feeding difficulties or infections or have undergone multiple exchange transfusions. In these babies, prophylactic folic acid should be given.

Hematologic Disorders Folate deficiency frequently occurs in chronic hemolytic anemia, particularly in sickle cell disease, autoimmune hemolytic anemia, and congenital spherocytosis. In these and other conditions of increased cell turnover (e.g., myelofibrosis, malignancies), folate deficiency arises because it is not completely reutilized after performing coenzyme functions.

Inflammatory Conditions Chronic inflammatory diseases such as tuberculosis, rheumatoid arthritis, Crohn's disease, psoriasis, exfoliative dermatitis, bacterial endocarditis, and chronic bacterial infections cause deficiency by reducing the appetite and increasing the demand for folate. Systemic infections also may cause malabsorption of folate. Severe deficiency is virtually confined to the patients with the most active disease and the poorest diet.

Homocystinuria This is a rare metabolic defect in the conversion of homocysteine to cystathionine. Folate deficiency occurring in most of these patients may be due to excessive utilization because of compensatory increased conversion of homocysteine to methionine.

Long-Term Dialysis Because folate is only loosely bound to plasma proteins, it is easily removed from plasma by dialysis. In patients with anorexia, vomiting, infections, and hemolysis, folate stores are particularly likely to become depleted. Routine folate prophylaxis is now given.

Congestive Heart Failure, Liver Disease Excess urinary folate losses of >100 µg per day may occur in some of these patients. The explanation appears to be release of folate from damaged liver cells.

ANTIFOLATE DRUGS

A large number of epileptics who are receiving long-term therapy with phenytoin or primidone, with or without barbiturates, develop low serum and red cell folate levels. The exact mechanism is unclear. Alcohol may also be a folate antagonist, as patients who are drinking spirits may develop megaloblastic anemia that will respond to normal quantities of dietary folate or to physiologic doses of folic acid only if alcohol is withdrawn. Macrocytosis of red cells is associated with chronic alcohol intake even when folate levels are normal. Inadequate folate intake is the major factor in the development of deficiency in spirit-drinking alcoholics. Beer is relatively folate-rich in some countries, depending on the technique used for brewing.

The drugs that inhibit DHF reductase include methotrexate, pyrimethamine, and trimethoprim. Methotrexate has the most powerful action against the human enzyme, whereas trimethoprim is most active against the bacterial enzyme and is likely to cause megaloblastic anemia only when used in conjunction with sulfamethoxazole in patients with preexisting folate or cobalamin deficiency. The activity of pyrimethamine is intermediate. The antidote to these drugs is folic acid (5-formyl-THF).

CONGENITAL ABNORMALITIES OF FOLATE METABOLISM

Some infants with congenital defects of folate enzymes (e.g., cyclohydrolase or methionine synthase) have had megaloblastic anemia.

DIAGNOSIS OF COBALAMIN AND FOLATE DEFICIENCIES

The diagnosis of cobalamin or folate deficiency has traditionally depended on the recognition of the relevant abnormalities in the peripheral blood and analysis of the blood levels of the vitamins.

COBALAMIN DEFICIENCY

Serum Cobalamin This is measured by an automated enzyme-linked immunosorbent assay (ELISA) or competitive-binding luminescence assay (CBLA). Normal serum levels range from 118–148 pmol/L (160–200 ng/L) to ~738 pmol/L (1000 ng/L). In patients with megaloblastic anemia due to cobalamin deficiency, the level is usually <74 pmol/L (100 ng/L). In general, the more severe the deficiency, the lower is the serum cobalamin level. In patients with spinal cord damage due to the deficiency, levels are very low even in the absence of anemia. Values between 74 and 148 pmol/L (100 and 200 ng/L) are regarded as borderline. They may occur, for instance, in pregnancy, in patients with megaloblastic anemia due to folate deficiency. They may also be due to heterozygous, homozygous, or compound heterozygous mutations of the gene *TCNI* that codes for haptocorrin (transcobalamin I). There is no clinical or hematologic abnormality. The serum cobalamin level is sufficiently robust, cost-effective, and most convenient to rule out cobalamin deficiency in the vast majority of patients suspected of having this problem. However, problems have arisen with commercial CBLA assays involving intrinsic factor in PA patients with intrinsic antibodies in serum. These antibodies may cause false normal serum vitamin B₁₂ levels in up to 50% of cases tested. Where clinical indications of PA are strong, a normal serum vitamin B₁₂ does not rule out the diagnosis. Serum MMA levels will be elevated in PA (see below).

Serum Methylmalonate and Homocysteine In patients with cobalamin deficiency sufficient to cause anemia or neuropathy, the serum MMA level is raised. Sensitive methods for measuring MMA and homocysteine in serum have been introduced and recommended for the early diagnosis of cobalamin deficiency, even in the absence of hematologic abnormalities or subnormal levels of serum cobalamin. Serum MMA levels fluctuate, however, in patients with renal failure. Mildly elevated serum MMA and/or homocysteine levels occur in up to 30% of apparently healthy volunteers, with serum cobalamin levels up to 258 pmol/L (350 ng/L) and normal serum folate levels; 15% of elderly subjects, even with cobalamin levels >258 pmol/L (>350 ng/L), have this pattern of raised metabolite levels. These findings bring into question the exact cutoff points for normal MMA and homocysteine levels. It is also unclear at present whether these mildly raised metabolite levels have clinical consequences.

Serum homocysteine is raised in both early cobalamin and folate deficiency but may be raised in other conditions, e.g., chronic renal disease, alcoholism, smoking, pyridoxine deficiency, hypothyroidism, and therapy with steroids, cyclosporine, and other drugs. Levels are also higher in serum than in plasma, in men than in premenopausal women, in women taking hormone replacement therapy or in oral contraceptive users, and in elderly persons and patients with several inborn errors of metabolism affecting enzymes in trans-sulfuration pathways of homocysteine metabolism. Thus, homocysteine levels must be carefully interpreted for diagnosis of cobalamin or folate deficiency.

Tests for the Cause of Cobalamin Deficiency Only vegans, strict vegetarians, or people living on a totally inadequate diet will become vitamin B₁₂ deficient because of inadequate intake. Studies of cobalamin absorption once were widely used, but difficulty in obtaining radioactive cobalamin and ensuring that IF preparations are free of viruses has made these tests obsolete. Tests to diagnose PA include serum gastrin, which is raised; serum pepsinogen I, which is low in PA (90–92%) but also in other conditions; and gastric endoscopy. Tests for IF and parietal cell antibodies are also used, as well as tests for individual intestinal diseases.

FOLATE DEFICIENCY

Serum Folate This is also measured by an ELISA technique. In most laboratories, the normal range is from 11 nmol/L (2 µg/L) to ~82 nmol/L (15 µg/L). The serum folate level is low in all folate-deficient patients. It also reflects recent diet. Because of this, serum folate may be low before there is hematologic or biochemical evidence of deficiency. Serum folate rises in severe cobalamin deficiency because of the block in conversion of MTHF to THF inside cells; raised levels have also

648 been reported in the intestinal stagnant loop syndrome due to absorption of bacterially synthesized folate.

Red Cell Folate The red cell folate assay is a valuable test of body folate stores. It is less affected than the serum assay by recent diet and traces of hemolysis. In normal adults, concentrations range from 880–3520 $\mu\text{mol/L}$ (160–640 $\mu\text{g/L}$) of packed red cells. Subnormal levels occur in patients with megaloblastic anemia due to folate deficiency but also in nearly two-thirds of patients with severe cobalamin deficiency. False-normal results may occur if a folate-deficient patient has received a recent blood transfusion or if a patient has a raised reticulocyte count. Serum homocysteine assay is discussed earlier.

Tests for the Cause of Folate Deficiency The diet history is important. Tests for transglutaminase antibodies are performed to confirm or exclude celiac disease. If positive, duodenal biopsy is needed. An underlying disease causing increased folate breakdown should also be excluded.

TREATMENT COBALAMIN AND FOLATE DEFICIENCY

It is usually possible to establish which of the two deficiencies, folate or cobalamin, is the cause of the anemia and to treat only with the appropriate vitamin. In patients who enter the hospital severely ill, however, it may be necessary to treat with both vitamins in large doses once blood samples have been taken for cobalamin and folate assays and a bone marrow biopsy has been performed (if deemed necessary). Transfusion is usually unnecessary and inadvisable. If it is essential, packed red cells should be given slowly, one or two units only, with the usual treatment for heart failure if present. Potassium supplements have been recommended to obviate the danger of the hypokalemia but are not necessary. Occasionally, an excessive rise in platelets occurs after 1–2 weeks of therapy. Antiplatelet therapy, e.g., aspirin, should be considered if the platelet count rises to $>800 \times 10^9/\text{L}$.

COBALAMIN DEFICIENCY

It is usually necessary to treat patients who have developed cobalamin deficiency with lifelong regular cobalamin injections. In the UK, the form used is hydroxocobalamin; in the United States, cyanocobalamin. In a few instances, the underlying cause of cobalamin deficiency can be permanently corrected, e.g., fish tapeworm, tropical sprue, or an intestinal stagnant loop that is amenable to surgery. The indications for starting cobalamin therapy are a well-documented megaloblastic anemia or other hematologic abnormalities and neuropathy due to the deficiency. Patients with borderline serum cobalamin levels but no hematologic or other abnormality may be followed to make sure that the cobalamin deficiency does not progress (see below). If malabsorption of cobalamin or rises in serum MMA levels have been demonstrated, however, these patients also should be given regular maintenance cobalamin therapy. Cobalamin should be given routinely to all patients who have had a total gastrectomy or ileal resection. Patients who have undergone gastric reduction for control of obesity or who are receiving long-term treatment with proton pump inhibitors should be screened and, if necessary, given cobalamin replacement.

Replenishment of body stores should be complete with six 1000- μg IM injections of hydroxocobalamin given at 3- to 7-day intervals. More frequent doses are usually used in patients with cobalamin neuropathy, but there is no evidence that they produce a better response. Allergic reactions are rare and may require desensitization or antihistamine or glucocorticoid cover. For maintenance therapy, 1000 μg hydroxocobalamin IM once every 3 months is satisfactory. Because of the poorer retention of cyanocobalamin, protocols generally use higher and more frequent doses, e.g., 1000 μg IM, monthly, for maintenance treatment.

Because a small fraction of cobalamin can be absorbed passively through mucous membranes even when there is complete failure of physiologic IF-dependent absorption, large daily oral doses (1000–2000 μg) of cyanocobalamin have been used in PA for replacement

and maintenance of normal cobalamin status in, e.g., food malabsorption of cobalamin. Sublingual therapy has also been proposed for those in whom injections are difficult because of a bleeding tendency and who may not tolerate oral therapy. If oral therapy is used, it is important to monitor compliance, particularly with elderly, forgetful patients. This author prefers parenteral therapy for initial treatment, particularly in severe anemia or if a neuropathy is present, and for maintenance.

For treatment of patients with subnormal serum vitamin B₁₂ (B₁₂) levels with a normal MCV and no hypersegmentation of neutrophils, a negative IF antibody test in the absence of tests of B₁₂ absorption is problematic. Some (perhaps 15%) cases may be due to TC I (HC) deficiency. Homocysteine and/or MMA measurements may help, but in the absence of these tests and with otherwise normal gastrointestinal function, repeat serum B₁₂ assay after 6–12 months may help one decide whether to start cobalamin therapy.

Vitamin B₁₂ injections are used in a wide variety of diseases, often neurologic, despite normal serum B₁₂ and folate levels and a normal blood count and in the absence of randomized, double-blind, controlled trials. These conditions include multiple sclerosis and chronic fatigue syndrome/myalgic encephalomyelitis (ME). It seems probable that any benefit is due to the placebo effect of a usually painless, pink injection. In ME, oral B₁₂ therapy, despite providing equally large amounts of B₁₂, has not been beneficial, supporting the view of the effect of the injections being placebo only.

FOLATE DEFICIENCY

Oral doses of 5–15 mg folic acid daily are satisfactory, as sufficient folate is absorbed from these extremely large doses even in patients with severe malabsorption. The length of time therapy must be continued depends on the underlying disease. It is customary to continue therapy for about 4 months, when all folate-deficient red cells will have been eliminated and replaced by new folate-replete populations.

Before large doses of folic acid are given, cobalamin deficiency must be excluded and, if present, corrected; otherwise cobalamin neuropathy may develop despite a response of the anemia of cobalamin deficiency to folate therapy. Studies in the United States, however, suggest that there is no increase in the proportion of individuals with low serum cobalamin levels and no anemia since food fortification with folic acid, but it is unknown if there has been a change in incidence of cobalamin neuropathy.

Long-term folic acid therapy is required when the underlying cause of the deficiency cannot be corrected and the deficiency is likely to recur, e.g., in chronic dialysis or hemolytic anemias. It may also be necessary in gluten-induced enteropathy that does not respond to a gluten-free diet. Where mild but chronic folate deficiency occurs, it is preferable to encourage improvement in the diet after correcting the deficiency with a short course of folic acid. In any patient receiving long-term folic acid therapy, it is important to measure the serum cobalamin level at regular (e.g., once-yearly) intervals to exclude the coincidental development of cobalamin deficiency.

Folinic Acid (5-Formyl-THF) This is a stable form of fully reduced folate. It is given orally or parenterally to overcome the toxic effects of methotrexate or other DHF reductase inhibitors, e.g., trimethoprim or cotrimoxazole.

PROPHYLACTIC FOLIC ACID

Prophylactic folic acid is used in chronic dialysis patients and in parenteral feeds. Prophylactic folic acid has been used to reduce homocysteine levels to prevent cardiovascular disease and for cognitive function in the elderly, but there are no firm data to show any benefit.

Pregnancy In over 70 countries (but none in Europe), food is fortified with folic acid (in grain or flour) to reduce the risk of NTDs. Nevertheless, folic acid, 400 μg daily, should be given as a supplement before and throughout pregnancy to prevent megaloblastic

anemia and reduce the incidence of NTDs, even in countries with fortification of the diet. The levels of fortification provide up to 400 µg daily on average in Chile, but in most countries, it is nearer to 200 µg, so periconceptual folic acid is still needed. Studies in early pregnancy show significant lack of compliance with the folic acid supplements, emphasizing the benefit of food fortification. Supplemental folic acid reduces the incidence of birth defects in babies born to diabetic mothers. In women who have had a previous fetus with an NTD, 5 mg daily is recommended when pregnancy is contemplated and throughout the subsequent pregnancy.

Infancy and Childhood The incidence of folate deficiency is so high in the smallest premature babies during the first 6 weeks of life that folic acid (e.g., 1 mg daily) should be given routinely to those weighing <1500 g at birth and to larger premature babies who require exchange transfusions or develop feeding difficulties, infections, or vomiting and diarrhea.

The World Health Organization currently recommends routine supplementation with iron and folic acid in children in countries where iron deficiency is common and child mortality, largely due to infectious diseases, is high. However, some studies suggest that in areas where malaria rates are high, this approach may increase the incidence of severe illness and death. Even where malaria is rare, there appears to be no survival benefit.

MEGALOBlastic ANEMIA NOT DUE TO COBALAMIN OR FOLATE DEFICIENCY OR ALTERED METABOLISM

This may occur with many antimetabolic drugs (e.g., hydroxyurea, cytosine arabinoside, 6-mercaptopurine) that inhibit DNA replication. Antiviral nucleoside analogues used in treatment of HIV infection may also cause macrocytosis and megaloblastic marrow changes. In the rare disease orotic aciduria, two consecutive enzymes in purine synthesis are defective. The condition responds to therapy with uridine, which bypasses the block. In thiamine-responsive megaloblastic anemia, there is a genetic defect in the high-affinity thiamine transport (*SLC19A2*) gene. This causes defective RNA ribose synthesis through impaired activity of transketolase, a thiamine-dependent enzyme in the pentose cycle. This leads to reduced nucleic acid production. It may be associated with diabetes mellitus and deafness and the presence of many ringed sideroblasts in the marrow. The explanation is unclear for megaloblastic changes in the marrow in some patients with acute myeloid leukemia and myelodysplasia.

129 Hemolytic Anemias and Anemia Due to Acute Blood Loss

Lucio Luzzatto

DEFINITIONS

A finite life span is a distinct characteristic of red cells. Hence, a logical, time-honored classification of anemias is in three groups: (1) decreased production of red cells, (2) increased destruction of red cells, and (3) acute blood loss. Decreased production is covered in [Chaps. 126, 128, and 130](#); increased destruction and acute blood loss are covered in this chapter.

All patients who are anemic as a result of either increased destruction of red cells or acute blood loss have one important element in common: the anemia results from overconsumption of red cells from the peripheral blood, whereas the supply of cells from the bone marrow is normal (indeed, it is usually increased). On the other hand, these two groups differ in that physical loss of red cells from the bloodstream or from the body itself, as in acute hemorrhage, is fundamentally different from destruction of red cells *within* the body, as in hemolytic anemias.

TABLE 129-1 CLASSIFICATION OF HEMOLYTIC ANEMIAS^a

	Intracorporeal Defects	Extracorporeal Factors
Hereditary	Hemoglobinopathies Enzymopathies Membrane-cytoskeletal defects	Familial (atypical) hemolytic-uremic syndrome
Acquired	Paroxysmal nocturnal hemoglobinuria (PNH)	Mechanical destruction (microangiopathic) Toxic agents Drugs Infectious Autoimmune

^aHereditary causes correlate with intracorporeal defects, because these defects are due to inherited mutations; the one exception is PNH, because the defect is due to an acquired somatic mutation. Similarly, acquired causes correlate with extracorporeal factors, because mostly these factors are exogenous; the one exception is familial hemolytic-uremic syndrome (HUS; often referred to as atypical HUS), because here an inherited abnormality allows complement activation to be excessive, with bouts of production of membrane attack complex capable of destroying normal red cells.

Therefore, the clinical aspects and pathophysiology of anemia in these two groups of patients are quite different, and they will be considered separately.

HEMOLYTIC ANEMIAS

With respect to primary etiology, anemias due to increased destruction of red cells, which we know as hemolytic anemias (HAs), may be *inherited* or *acquired*; from a clinical point of view, they may be more *acute* or more *chronic*, and they may vary from mild to very severe; the site of hemolysis may be predominantly *intravascular* or *extravascular*. With respect to mechanisms, HAs may be due to *intracorporeal* causes or to *extracorporeal* causes ([Table 129-1](#)). But before reviewing the individual types of HA, it is appropriate to consider what they have in common.

GENERAL CLINICAL AND LABORATORY FEATURES

The clinical presentation of a patient with anemia is greatly influenced in the first place by whether the onset is abrupt or gradual, and HAs are no exception. A patient with autoimmune HA or with favism may be a medical emergency, whereas a patient with mild hereditary spherocytosis or with cold agglutinin disease may be diagnosed after years. This is due in large measure to the remarkable ability of the body to adapt to anemia when it is slowly progressing ([Chap. 77](#)).

What differentiates HAs from other anemias is that the patient has signs and symptoms arising directly from hemolysis ([Table 129-2](#)). At the clinical level, the main sign is *jaundice*; in addition, the patient may report discoloration of the urine. In many cases of HA, the spleen is enlarged, because it is a preferential site of hemolysis; and in some cases, the liver may be enlarged as well. In all severe congenital forms of HA, there may also be skeletal changes due to overactivity of the bone marrow (although they are never as severe as they are in thalassemia).

TABLE 129-2 FEATURES COMMON TO MOST PATIENTS WITH A HEMOLYTIC DISORDER

General examination	Jaundice, pallor
Other physical findings	Spleen may be enlarged; bossing of skull in severe congenital cases
Hemoglobin level	From normal to severely reduced
MCV, MCH	Usually increased
Reticulocytes	Increased
Bilirubin	Increased (mostly unconjugated)
LDH	Increased (up to 10 time normal with intravascular hemolysis)
Haptoglobin	Reduced to absent (if hemolysis in part intravascular)

Abbreviations: LDH, lactate dehydrogenase; MCH, mean corpuscular hemoglobin; MCV, mean corpuscular volume.

The laboratory features of HA are related to hemolysis per se and the erythropoietic response of the bone marrow. Hemolysis regularly produces an increase in unconjugated bilirubin and aspartate aminotransferase (AST) in the serum; urobilinogen will be increased in both urine and stool. If hemolysis is mainly intravascular, the telltale sign is hemoglobinuria (often associated with hemosiderinuria); in the serum, there is hemoglobin, lactate dehydrogenase (LDH) is increased, and haptoglobin is reduced. In contrast, the bilirubin level may be normal or only mildly elevated. The main sign of the erythropoietic response by the bone marrow is an increase in reticulocytes (a test all too often neglected in the initial workup of a patient with anemia). Usually the increase will be reflected in both the percentage of reticulocytes (the more commonly quoted figure) and the absolute reticulocyte count (the more definitive parameter). The increased number of reticulocytes is associated with an increased mean corpuscular volume (MCV) in the blood count. On the blood smear, this is reflected in the presence of macrocytes; there is also polychromasia, and sometimes one sees nucleated red cells. In most cases, a bone marrow aspirate is not necessary in the diagnostic workup; if it is done, it will show erythroid hyperplasia. In practice, once an HA is suspected, specific tests will usually be required for a definitive diagnosis of a specific type of HA.

GENERAL PATHOPHYSIOLOGY

The mature red cell is the product of a developmental pathway that brings the phenomenon of differentiation to an extreme. An orderly sequence of events produces synchronous changes, whereby the gradual accumulation of a huge amount of hemoglobin in the cytoplasm (to a final level of 340 g/L, i.e., about 5 mM) goes hand in hand with the gradual loss of cellular organelles and of biosynthetic abilities. In the end, the erythroid cell undergoes a process that has features of apoptosis, including nuclear pyknosis and actual loss of the nucleus. However, the final result is more altruistic than suicidal; the cytoplasmic body, instead of disintegrating, is now able to provide oxygen to all cells in the human organism for some remaining 120 days of the red cell life span.

As a result of this unique process of differentiation and maturation, intermediary metabolism is drastically curtailed in mature red cells (Fig. 129-1); for instance, cytochrome-mediated oxidative phosphorylation has been lost with the loss of mitochondria (through a process of physiologic autophagy); therefore, there is no backup to anaerobic glycolysis, which in the red cell is the only provider of adenosine triphosphate (ATP). Also the capacity of making protein has been lost with the loss of ribosomes. This places the cell's limited metabolic apparatus at risk, because if any protein component deteriorates, it cannot be replaced, as it would be in most other cells; and in fact the activity of most enzymes gradually decreases as red cells age. At the same time, during their long time in circulation, various red cell components inevitably accumulate damage; in senescent red cells, the membrane protein band 3 molecules (see below and Fig. 129-1), having bound hemichromes on their intracellular domains, tend to cluster. Now they bind anti-band 3 IgG antibodies (present in most people) and C3 complement fragments; thus they become opsonized and are eventually removed by phagocytosis in the reticuloendothelial system.

Another consequence of the relative simplicity of red cells is that they have a very limited range of ways to manifest distress under hardship; in essence, any sort of metabolic failure will eventually lead either to structural damage to the membrane or to failure of the cation pump. In either case, the life span of the red cell is reduced, which is the definition of a *hemolytic disorder*. If the rate of red cell destruction exceeds the capacity of the bone marrow to produce more red cells, the hemolytic disorder will manifest as HA.

Thus, the essential pathophysiologic process common to all HAs is an increased red cell turnover; and in many HAs, this is due at least in part to an acceleration of the senescence process described above. The gold standard for proving that the life span of red cells is reduced (compared to the normal value of about 120 days) is a *red cell survival study*, which can be carried out by labeling the red cells with ^{51}Cr and measuring residual radioactivity over several days or weeks; however, this classic test is now available in very few centers, and it is rarely necessary.

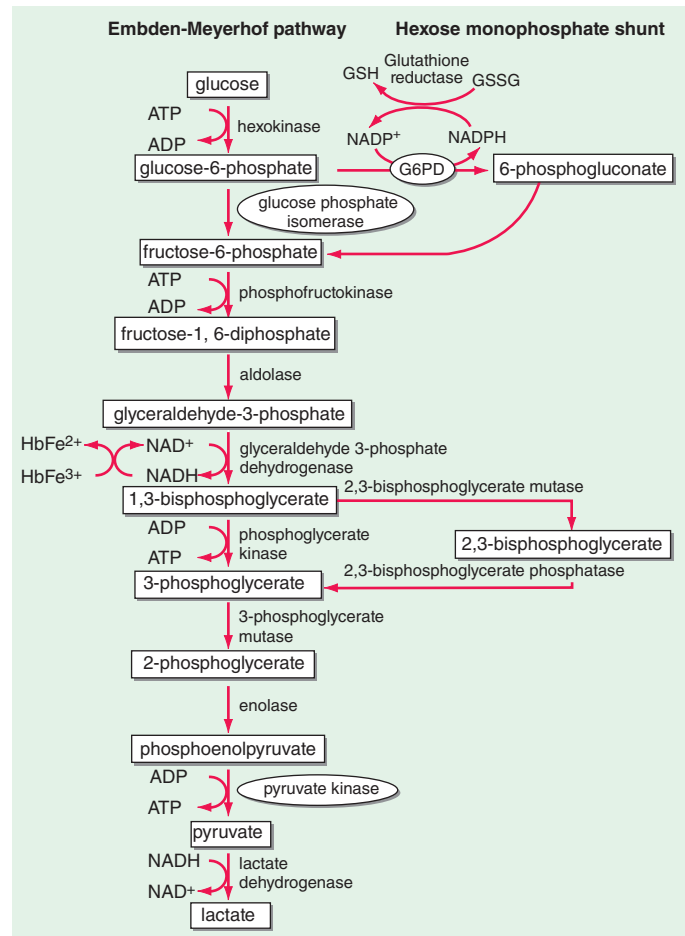


FIGURE 129-1 Red blood cell (RBC) metabolism. The Embden-Meyerhof pathway (glycolysis) generates ATP for energy and membrane maintenance. The generation of NADPH maintains hemoglobin in a reduced state. The hexose monophosphate shunt generates NADPH that is used to reduce glutathione, which protects the red cell against oxidant stress. Regulation of 2,3-bisphosphoglycerate levels is a critical determinant of oxygen affinity of hemoglobin. Enzyme deficiency states in order of prevalence: glucose 6-phosphate dehydrogenase (G6PD) > pyruvate kinase > glucose-6-phosphate isomerase > rare deficiencies of other enzymes in the pathway. The more common enzyme deficiencies are encircled.

If the hemolytic event is transient, it does not usually cause any long-term consequences, except for an increased requirement for erythropoietic factors, particularly folic acid. However, if hemolysis is recurrent or persistent, the increased bilirubin production favors the formation of gallstones. If a considerable proportion of hemolysis takes place in the spleen, as is often the case, splenomegaly may become increasingly a feature, and hypersplenism may develop, with consequent neutropenia and/or thrombocytopenia.

The increased red cell turnover also has metabolic consequences. In normal subjects, the iron from effete red cells is very efficiently recycled by the body; however, with chronic intravascular hemolysis, the persistent hemoglobinuria will cause considerable iron loss, needing replacement. With chronic extravascular hemolysis, the opposite problem, iron overload, is more common, especially if the patient needs frequent blood transfusions. Chronic iron overload will cause secondary hemochromatosis; this will cause damage particularly to the liver, eventually leading to cirrhosis, and to the heart muscle, eventually causing heart failure.

Compensated Hemolysis Versus Hemolytic Anemia Red cell destruction is a potent stimulus for erythropoiesis, which is mediated by erythropoietin (EPO) produced by the kidney. This mechanism is so effective that

in many cases the increased output of red cells from the bone marrow can fully balance an increased destruction of red cells. In such cases, we say that hemolysis is *compensated*. The pathophysiology of compensated hemolysis is similar to what we have just described, except there is no anemia. This notion is important from the diagnostic point of view, because a patient with a hemolytic condition, even an inherited one, may present without anemia; and it is also important from the point of view of management, because compensated hemolysis may become “decompensated,” i.e., anemia may suddenly appear, in certain circumstances, for instance in pregnancy, folate deficiency, or renal failure interfering with adequate EPO production. Another general feature of chronic HAs is seen when any intercurrent condition, such as an acute infection, depresses erythropoiesis. When this happens, in view of the increased rate of red cell turnover, the effect will be predictably much more marked than in a person who does not have hemolysis. The most dramatic example is infection by parvovirus B19, which may cause a rather precipitous fall in hemoglobin—an occurrence sometimes referred to as *aplastic crisis*.

INHERITED HEMOLYTIC ANEMIAS

There are three essential components in the red cell: (1) hemoglobin, (2) the membrane-cytoskeleton complex, and (3) the metabolic machinery necessary to keep hemoglobin and the membrane-cytoskeleton complex in working order. Diseases caused by abnormalities of hemoglobin, or hemoglobinopathies, are covered in **Chap. 127**. Here we will deal with diseases of the other two components.

Hemolytic Anemias due to Abnormalities of the Membrane-Cytoskeleton Complex

The detailed architecture of the red cell membrane is complex, but its basic design is relatively simple (**Fig. 129-2**). The lipid bilayer incorporates phospholipids and cholesterol, and it is spanned by a number of proteins that have their hydrophobic transmembrane domain(s) embedded in the membrane; most of these proteins also extend to both the outside (extracellular domains) and the inside of the cell (cytoplasmic domains). Other proteins are tethered to the membrane through a glycosylphosphatidylinositol (GPI) anchor; these have only an extracellular domain, and they include ion channels, receptors for complement components, and receptors for other ligands. The most abundant red cell membrane proteins are glycophorins and the so-called *band 3*, an anion transporter. The extracellular domains of many of these proteins are heavily glycosylated, and they carry antigenic

determinants that correspond to blood groups. Underneath the membrane, and tangential to it, is a network of other proteins that make up the cytoskeleton. The main cytoskeletal protein is spectrin, the basic unit of which is a dimer of α -spectrin and β -spectrin. The membrane is physically linked to the cytoskeleton by a third set of proteins (including ankyrin and the so-called *band 4.1* and *band 4.2*), which thus make these two structures intimately connected to each other.

The membrane-cytoskeleton complex is so integrated that, not surprisingly, an abnormality of almost any of its components will be disturbing or disruptive, causing structural failure, which results ultimately in hemolysis. These abnormalities are almost invariably inherited mutations; thus, diseases of the membrane-cytoskeleton complex belong to the category of inherited HAs. Before the red cells lyse, they often exhibit more or less specific morphologic changes that alter the normal biconcave disk shape. Thus, the majority of the diseases in this group have been known for over a century as *hereditary spherocytosis* and *hereditary elliptocytosis*. Over the past 20 years, their molecular basis has been elucidated; it has emerged that both conditions can arise from mutations in several genes with considerable overlap (**Fig. 129-3**).

HEREDITARY SPHEROCYTOSIS (HS) This is a relatively common type of genetically determined HA, with an estimated frequency of at least 1 in 5000. Its identification is credited to Minkowsky and Chauffard, who, at the end of the nineteenth century, reported families who had the presence of numerous spherocytes in the peripheral blood (**Fig 129-4A**). In vitro studies revealed that the red cells were abnormally susceptible to lysis in hypotonic media; indeed, the presence of *osmotic fragility* became the main diagnostic test for HS. Today we know that HS, thus defined, is genetically heterogeneous; i.e., it can arise from a variety of mutations in one of several genes (**Table 129-3**). It has been also recognized that the inheritance of HS is not always autosomal dominant (with the patient being heterozygous); indeed, some of the most severe forms are instead autosomal recessive (with the patient being homozygous).

Clinical presentation and diagnosis The spectrum of clinical severity of HS is broad. Severe cases may present in infancy with severe anemia, whereas mild cases may present in young adults or even later in life. The main clinical findings are jaundice, an enlarged spleen, and often gallstones; indeed, it may be the finding of gallstones in a young person that triggers diagnostic investigations.

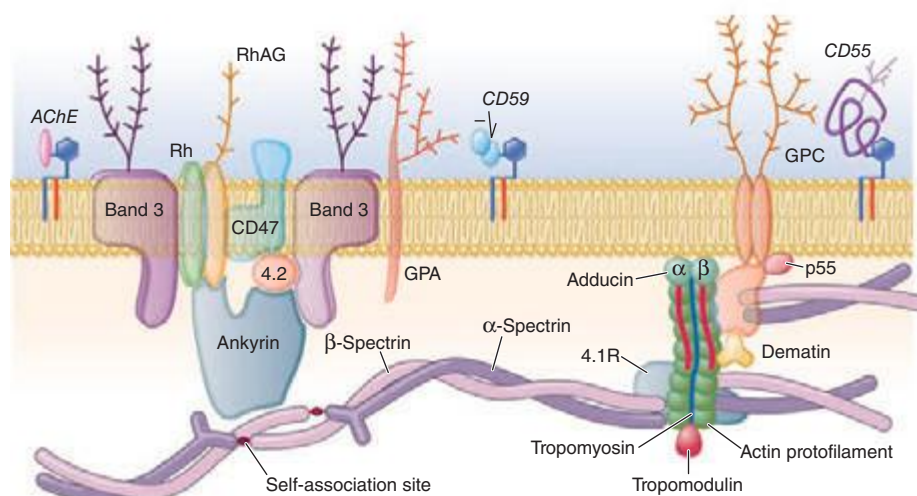


FIGURE 129-2 The red cell membrane. In this figure, one sees, within the lipid bilayer, several membrane proteins, of which band 3 (anion exchanger 1 [AE1]) is the most abundant; the α - β spectrin dimers that associate to form most of the cytoskeleton; and several proteins (e.g., ankyrin) that connect the membrane to the cytoskeleton. In addition, as examples of glycosylphosphatidylinositol (GPI)-linked proteins, one sees acetylcholinesterase (AChE) and the two complement-regulatory proteins CD59 and CD55. The (nonrealistic) shapes of the protein moieties of the GPI-linked proteins are meant to indicate that they can be very different from each other and that, unlike with the other membrane proteins shown, the entire polypeptide chain is extracellular. Branched lines symbolize carbohydrate moiety of proteins. The molecules are obviously not drawn to the same scale. Additional explanations can be found in the text. (From N Young et al: *Clinical Hematology*. Copyright Elsevier, 2006; with permission.)

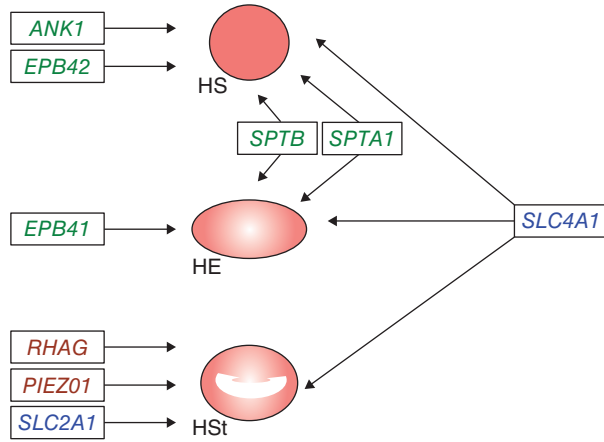


FIGURE 129-3 Hereditary spherocytosis (HS), hereditary elliptocytosis (HE), and hereditary stomatocytosis (HSst) are three morphologically distinct forms of congenital hemolytic anemia. It has emerged that each one can arise from mutation of one of several genes and that different mutations of the same gene can give one or another form. (See also Table 129-3.)

The variability in clinical manifestations that is observed among patients with HS is largely due to the different underlying molecular lesions (Table 129-3). Not only are mutations of several genes involved, but also individual mutations of the same gene can also give very different clinical manifestations. In milder cases, hemolysis is often compensated (see above), and this may cause variation in time even in the same patient, due to the fact that intercurrent conditions (e.g., pregnancy, infection) may cause decompensation. The anemia is usually normocytic, with the characteristic morphology that gives the disease its name. An increased mean corpuscular hemoglobin concentration (MCHC) on an ordinary blood count report should raise the suspicion of HS, because HS is almost the only condition in which this abnormality occurs. It has been apparent for a long time that the spleen plays a special role in HS through a dual mechanism. On one hand, like in many other HAs, the spleen itself is a major site of destruction; on the other hand, transit through the splenic circulation makes the defective red cells more spherocytic and, therefore, accelerates their demise, even though that may take place elsewhere.

When there is a family history, it is usually easy to make a diagnosis based on features of HA and typical red cell morphology. However, there may be no family history for at least two reasons. First, the patient may have a *de novo* mutation, i.e., a mutation that has taken place in a germ cell of one of his parents or early after zygote formation. Second, the patient may have a recessive form of HS (Table 129-3). In such cases, more extensive laboratory investigations are required, including osmotic fragility, the acid glycerol lysis test, the eosin-5'-maleimide (EMA)-binding test, and SDS-gel electrophoresis of membrane proteins; these tests are usually carried out in laboratories with special expertise in this area. Sometimes a definitive diagnosis can be obtained only by molecular studies demonstrating a mutation in one of the genes underlying HS (Table 129-3).

TREATMENT HEREDITARY SPHEROCYTOSIS

We do not have a causal treatment for HS; i.e., no way has yet been found to correct the basic defect in the membrane-cytoskeleton structure. Given the special role of the spleen in HS (see above), it has long been thought that an almost obligatory therapeutic measure was splenectomy. Because this operation may have more than trivial consequences, today we have more articulate recommendations, based on disease severity (having found out, whenever possible, about the outcome of splenectomy in the patient's relatives with HS), as follows. In mild cases, avoid splenectomy. Delay splenectomy until puberty in moderate cases or until 4–6 years of age in severe cases. Antipneumococcal vaccination before splenectomy

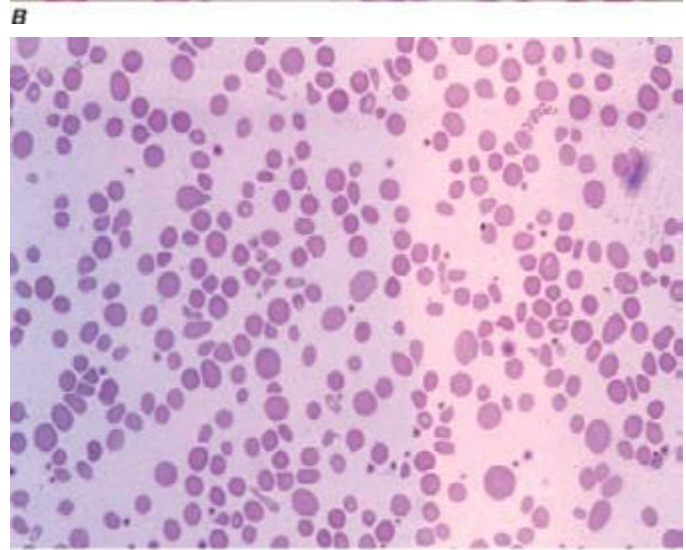
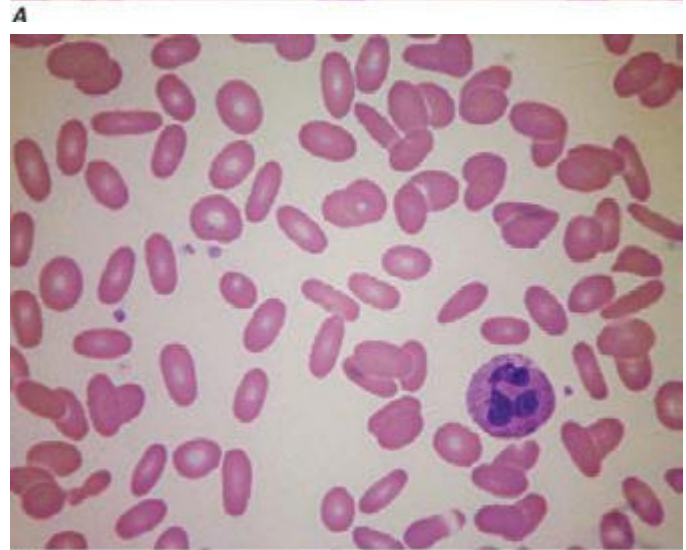
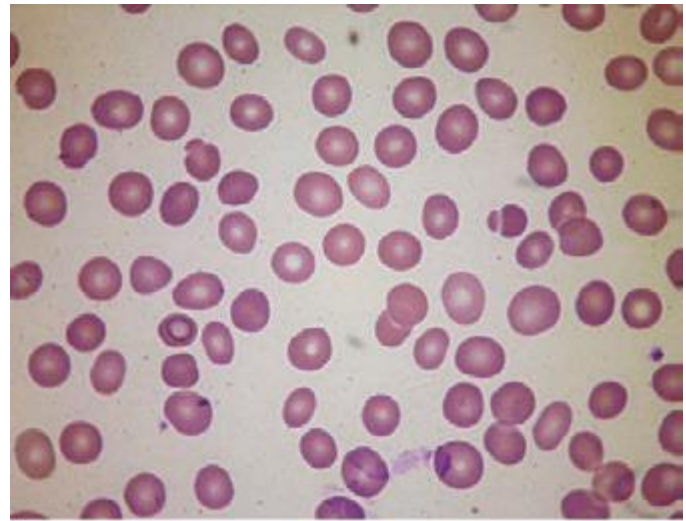


FIGURE 129-4 Peripheral blood smear from patients with membrane-cytoskeleton abnormalities. **A.** Hereditary spherocytosis. **B.** Hereditary elliptocytosis, heterozygote. **C.** Elliptocytosis, with both alleles of the α -spectrin gene mutated.

is imperative, whereas penicillin prophylaxis after splenectomy is controversial. Along with splenectomy, cholecystectomy should not be regarded as automatic; it should be carried out, usually by the laparoscopic approach, when clinically indicated.

TABLE 129-3 INHERITED DISEASES OF THE RED CELL MEMBRANE-CYTOSKELETON COMPLEX

Gene	Chromosomal Location	Protein Produced	Disease(s) with Certain Mutations (Inheritance)	Comments
<i>SPTA1</i>	1q22-q23	α -Spectrin	HS (recessive) HE (dominant)	Rare Mutations of this gene account for about 65% of HE. More severe forms may be due to coexistence of an otherwise silent mutant allele.
<i>SPTB</i>	14q23-q24.1	β -Spectrin	HS (dominant) HE (dominant)	Rare Mutations of this gene account for about 30% of HE, including some severe forms.
<i>ANK1</i>	8p11.2	Ankyrin	HS (dominant)	May account for majority of HS.
<i>SLC4A1</i>	17q21	Band 3; also known as AE (anion exchanger) or AE1	HS (dominant) Southeast Asia ovalocytosis (dominant) Stomatocytosis	Mutations of this gene may account for about 25% of HS. Polymorphic mutation (deletion of 9 amino acids); clinically asymptomatic; protective against <i>Plasmodium falciparum</i> . Certain specific missense mutations shift protein function from anion exchanger to cation conductance.
<i>EPB41</i>	1p33-p34.2	Band 4.1	HE (dominant)	Mutations of this gene account for about 5% of HE, mostly with prominent morphology but no hemolysis in heterozygotes; severe hemolysis in homozygotes.
<i>EPB42</i>	15q15-q21	Band 4.2	HS (recessive)	Mutations of this gene account for about 3% of HS.
<i>RHAG</i>	6p21.1-p11	Rhesus antigen	Chronic nonspherocytic hemolytic anemia (recessive)	Very rare; associated with total loss of all Rh antigens. A specific mutation causes overhydrated stomatocytosis.
<i>PIEZO1</i>	16q23-q24	PIEZO1	Dehydrated hereditary stomatocytosis (dominant)	Also known as xerocytosis with pseudohyperkalemia. Patients may present with perinatal edema. PIEZO1 is a mechanosensitive cation channel.

Abbreviations: HE, hereditary elliptocytosis; HS, hereditary spherocytosis.

HEREDITARY ELLIPTOCYTOSIS (HE) HE is at least as heterogeneous as HS, both from the genetic point of view (Table 129-3, Fig. 129-3) and from the clinical point of view. Again, it is the shape of the red cells (Fig. 129-4B) that gives the name to the condition, but there is no direct correlation between the elliptocytic morphology and clinical severity. In fact, some mild or even asymptomatic cases may have nearly 100% elliptocytes, whereas in severe cases, all kinds of bizarre poikilocytes can predominate. Clinical features and recommended management are similar to those outlined above for HS. Although the spleen may not have the specific role it has in HS, in severe cases, splenectomy may be beneficial. The prevalence of HE causing clinical disease is similar to that of HS. However, an in-frame deletion of nine amino acids in the *SLC4A1* gene encoding band 3, causing the so-called *Southeast Asia ovalocytosis*, has a frequency of up to 7% in certain populations, presumably as a result of malaria selection; it is asymptomatic in heterozygotes and probably lethal in homozygotes.

Disorders of Cation Transport These rare conditions with autosomal dominant inheritance are characterized by increased intracellular sodium in red cells, with concomitant loss of potassium; indeed, they are sometimes discovered through the incidental finding, in a blood test, of a high serum K^+ (*pseudohyperkalemia*). In patients from some families, the cation transport disturbance is associated with gain of water; as a result, the red cells are overhydrated (low MCHC), and on a blood smear, the normally round-shaped central pallor is replaced by a linear-shaped central pallor, which has earned this disorder the name *stomatocytosis* (Fig. 129-3). In patients from other families, instead, the red cells are dehydrated (high MCHC), and their consequent rigidity has earned this disorder the name *xerocytosis*. One would surmise that in these disorders the primary defect may be in a cation transporter; indeed, xerocytosis results from mutations in *PIEZO1*. In other patients with stomatocytosis, mutations are found in other genes also related to solute transport (Table 129-3), including *SLC4A1* (encoding band 3), the Rhesus gene *RHAG*, and the glucose transporter gene *SLC2A1* responsible for a special form called *cryohydrocytosis*. Hemolysis can vary from relatively mild to quite severe. From the practical point of view, it is important to know that in stomatocytosis, splenectomy is strongly contraindicated because it has been followed in a significant proportion of cases by severe thromboembolic complications.

Enzyme Abnormalities When there is an important defect in the membrane or in the cytoskeleton, hemolysis is a direct consequence of the fact that the very structure of the red cell is abnormal. Instead, when one of the enzymes is defective, the consequences will depend on the precise role of that enzyme in the metabolic machinery of the red cell, which, in first approximation, has two important functions: (1) to provide energy in the form of ATP and (2) to prevent oxidative damage to hemoglobin and to other proteins by providing sufficient reductive potential; the key molecule for this is NADPH.

ABNORMALITIES OF THE GLYCOLYTIC PATHWAY Because red cells, in the course of their differentiation, have sacrificed not only their nucleus and their ribosomes, but also their mitochondria, they rely exclusively on the anaerobic portion of the glycolytic pathway for producing energy in the form of ATP. Most of the ATP is required by the red cell for cation transport against a concentration gradient across the membrane. If this fails, due to a defect of any of the enzymes of the glycolytic pathway (Table 129-4), the result will be hemolytic disease.

Pyruvate kinase deficiency Abnormalities of the glycolytic pathway are all inherited and all rare. Among them, deficiency of pyruvate kinase (PK) is the least rare, with an estimated prevalence in most populations of the order of 1:10,000. However, very recently, a polymorphic PK mutation (E277K) was found in some African populations, with heterozygote frequencies of 1–7%, suggesting that this may be another malaria-related polymorphism. The clinical picture of homozygous (or compound biallelic) PK deficiency is that of an HA that often presents in the newborn with neonatal jaundice; the jaundice persists, and it is usually associated with a very high reticulocytosis. The anemia is of variable severity; sometimes it is so severe as to require regular blood transfusion treatment, whereas sometimes it is mild, bordering on a nearly compensated hemolytic disorder. As a result, the diagnosis may be delayed, and in some cases, it is made, for instance, in a young woman during her first pregnancy, when the anemia may get worse. The delay in diagnosis may be also helped by the fact that the anemia is remarkably well tolerated, because the metabolic block at the last step in glycolysis causes an increase in bisphosphoglycerate (or DPG; Fig. 129-1), a major effector of the hemoglobin-oxygen dissociation curve; thus, the oxygen delivery to the tissues is enhanced, a remarkable compensatory feat.

Enzyme (Acronym)	Chromosomal Location	Prevalence of Enzyme Deficiency (Rank)	Clinical Manifestations Extra-Red Cell	Comments
Glycolytic Pathway				
Hexokinase (HK)	10q22	Very rare		Other isoenzymes known
Glucose 6-phosphate isomerase (G6PI)	19q31.1	Rare (4) ^a	NM, CNS	
Phosphofructokinase (PFK)	12q13	Very rare	Myopathy	
Aldolase	16q22-24	Very rare		
Triose phosphate isomerase (TPI)	12p13	Very rare	CNS (severe), NM	
Glyceraldehyde 3-phosphate dehydrogenase (GAPD)	12p13.31-p13.1	Very rare	Myopathy	
Diphosphoglycerate mutase (DPGM)	7q31-q34	Very rare		Erythrocytosis rather than hemolysis
Phosphoglycerate kinase (PGK)	Xq13	Very rare	CNS, NM	May benefit from splenectomy
Pyruvate kinase (PK)	1q21	Rare (2) ^a		May benefit from splenectomy
Redox				
Glucose 6-phosphate dehydrogenase (G6PD)	Xq28	Common (1) ^a	Very rarely granulocytes	In almost all cases, only AHA from exogenous trigger
Glutathione synthase	20q11.2	Very rare	CNS	
γ-Glutamylcysteine synthase	6p12	Very rare	CNS	
Cytochrome b5 reductase	22q13.31-qter	Rare	CNS	Methemoglobinemia rather than hemolysis
Nucleotide Metabolism				
Adenylate kinase (AK)	9q34.1	Very rare	CNS	
Pyrimidine 5'-nucleotidase (P5N)	3q11-q12	Rare (3) ^a		May benefit from splenectomy

^aThe numbers from (1) to (4) indicate the ranking order of these enzymopathies in terms of frequency.

Abbreviations: AHA, acquired hemolytic anemia; CNS, central nervous system; NM, neuromuscular.

TREATMENT PYRUVATE KINASE DEFICIENCY

The management of PK deficiency is mainly supportive. In view of the marked increase in red cell turnover, oral folic acid supplements should be given constantly. Blood transfusion should be used as necessary, and iron chelation may have to be added if the blood transfusion requirement is high enough to cause iron overload. In these patients, who have more severe disease, splenectomy may be beneficial. There is a single case report of curative treatment of PK deficiency by bone marrow transplantation from an HLA-identical PK-normal sibling. This seems a viable option for severe cases when a sibling donor is available. Rescue of inherited PK deficiency through lentiviral-mediated human PK gene transfer has been successful in mice. Prenatal diagnosis has been carried out in a mother who had already had an affected child.

Other glycolytic enzyme abnormalities All of these defects are rare to very rare (Table 129-4), and all cause hemolytic anemia with varying degrees of severity. It is not unusual for the presentation to be in the guise of severe neonatal jaundice, which may require exchange transfusion; if the anemia is less severe, it may present later in life, or it may even remain asymptomatic and be detected incidentally when a blood count is done for unrelated reasons. The spleen is often enlarged. When other systemic manifestations occur, they can involve the central nervous system (sometimes entailing severe mental retardation, particularly in the case of triose phosphate isomerase deficiency), the neuromuscular system, or both. This is not altogether surprising, if we consider that these are housekeeping genes. The *diagnosis* of hemolytic anemia is usually not difficult, thanks to the triad of normomacrocytic anemia, reticulocytosis, and hyperbilirubinemia. Enzymopathies should be considered in the differential diagnosis of any chronic Coombs-negative hemolytic anemia. Unlike with membrane disorders where the red cells show characteristic morphologic abnormalities, in most cases of glycolytic enzymopathies, these are conspicuous by their absence. A definitive diagnosis can be made only by demonstrating the deficiency of an individual enzyme by quantitative assays; these

are carried out in only a few specialized laboratories. If a particular molecular abnormality is already known in the family, then one could test directly for that defect at the DNA level, thus bypassing the need for enzyme assays. Of course the time may be getting nearer when a patient will present with her or his exome already sequenced, and we will need to concentrate on which genes to look up within the file. The principles for the management of these conditions are similar as for PK deficiency. In one case of phosphoglycerate kinase deficiency, allogeneic bone marrow transplantation (BMT) effectively controlled the hematologic manifestations but did not reverse neurologic damage.

ABNORMALITIES OF REDOX METABOLISM

Glucose 6-phosphate dehydrogenase (G6PD) deficiency G6PD is a housekeeping enzyme critical in the redox metabolism of all aerobic cells (Fig. 129-1). In red cells, its role is even more critical, because it is the only source of NADPH, which directly and via glutathione (GSH) defends these cells against oxidative stress (Fig. 129-5). G6PD deficiency is a prime example of an HA due to interaction between

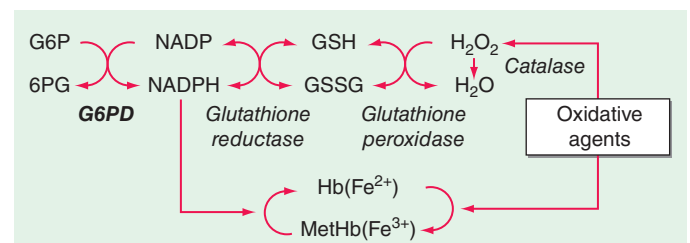


FIGURE 129-5 Diagram of redox metabolism in the red cell. 6PG, 6-phosphogluconate; G6P, glucose 6-phosphate; G6PD, glucose 6-phosphate dehydrogenase; GSH, reduced glutathione; GSSG, oxidized glutathione; Hb, hemoglobin; MetHb, methemoglobin; NADP, nicotinamide adenine dinucleotide phosphate; NADPH, reduced nicotinamide adenine dinucleotide phosphate.

an intracorporeal cause and an extracorporeal cause, because in the majority of cases hemolysis is triggered by an exogenous agent. Although a decrease in G6PD activity is present in most tissues of G6PD-deficient subjects, in other cells, the decrease is much less marked than in red cells, and it does not seem to impact on clinical expression.

GENETIC CONSIDERATIONS



The *G6PD* gene is X-linked, and this has important implications. First, because males have only one *G6PD* gene (i.e., they are hemizygous for this gene), they must be either normal or G6PD deficient. By contrast, females, who have two *G6PD* genes, can be either normal or deficient (homozygous) or intermediate (heterozygous). As a result of the phenomenon of X chromosome inactivation, heterozygous females are genetic mosaics, with a highly variable ratio of G6PD-normal to G6PD-deficient cells and an equally variable degree of clinical expression; some heterozygotes can be just as affected as hemizygous males. The enzymatically active form of G6PD is either a dimer or a tetramer of a single protein subunit of 514 amino acids. G6PD-deficient subjects have been found invariably to have mutations in the coding region of the *G6PD* gene (Fig. 129-5). Almost all of the approximately 180 different mutations known are single missense point mutations, entailing single amino acid replacements in the G6PD protein. In most cases, these mutations cause G6PD deficiency by decreasing the *in vivo* stability of the protein; thus, the physiologic decrease in G6PD activity that takes place with red cell aging is greatly accelerated. In some cases, an amino acid replacement can also affect the catalytic function of the enzyme.

Among these mutations, those underlying chronic nonspherocytic hemolytic anemia (CNSHA; see below) are a discrete subset. This much more severe clinical phenotype can be ascribed in some cases to adverse qualitative changes (for instance, a decreased affinity for the substrate, glucose 6-phosphate) or simply to the fact that the enzyme deficit is more extreme, because of a more severe instability of the enzyme. For instance, a cluster of mutations map at or near the dimer interface, and clearly they compromise severely the formation of the dimer.



Epidemiology G6PD deficiency is widely distributed in tropical and subtropical parts of the world (Africa, Southern Europe, the Middle East, Southeast Asia, and Oceania) (Fig. 129-6) and wherever people from those areas have migrated. A conservative estimate is that at least 400 million people have a *G6PD* deficiency gene. In several of these areas, the frequency of a *G6PD* deficiency gene may be as high as 20% or more. It would be quite extraordinary for a trait that causes significant pathology to spread widely and reach high frequencies in many populations without conferring some biologic advantage. Indeed, G6PD is one of the best-characterized examples of genetic polymorphisms in the human species. Clinical field studies and

in vitro experiments strongly support the view that G6PD deficiency has been selected by *Plasmodium falciparum* malaria, by virtue of the fact that it confers a relative resistance against this highly lethal infection. Different G6PD variants underlie G6PD deficiency in different parts of the world. Some of the more widespread variants are G6PD Mediterranean on the shores of that sea, in the Middle East, and in India; G6PD A⁻ in Africa and in Southern Europe; G6PD Vianchan and G6PD Mahidol in Southeast Asia; G6PD Canton in China; and G6PD Union worldwide. The heterogeneity of polymorphic G6PD variants is proof of their independent origin, and it supports the notion that they have been selected by a common environmental agent, in keeping with the concept of convergent evolution (Fig. 129-6).

Clinical manifestations The vast majority of people with G6PD deficiency remain clinically asymptomatic throughout their lifetime; however, all of them have an increased risk of developing neonatal jaundice (NNJ) and a risk of developing acute HA (AHA) when challenged by a number of oxidative agents. NNJ related to G6PD deficiency is very rarely present at birth; the peak incidence of clinical onset is between day 2 and day 3, and in most cases, the anemia is not severe. However, NNJ can be very severe in some G6PD-deficient babies, especially in association with prematurity, infection, and/or environmental factors (such as naphthalene-camphor balls, which are used in babies' bedding and clothing), and the risk of severe NNJ is also increased by the coexistence of a monoallelic or biallelic mutation in the uridyl transferase gene (*UGT1A1*; the same mutations are associated with Gilbert's syndrome). If inadequately managed, NNJ associated with G6PD deficiency can produce kernicterus and permanent neurologic damage.

AHA can develop as a result of three types of triggers: (1) fava beans, (2) infections, and (3) drugs (Table 129-5). Typically, a hemolytic attack starts with malaise, weakness, and abdominal or lumbar pain. After an interval of several hours to 2–3 days, the patient develops jaundice and often dark urine. The onset can be extremely abrupt, especially with favism in children. The anemia is moderate to extremely severe, usually normocytic and normochromic, and due partly to intravascular hemolysis; hence, it is associated with hemoglobinemia, hemoglobinuria, high LDH, and low or absent plasma haptoglobin. The blood film shows anisocytosis, polychromasia, and spherocytes typical of hemolytic anemias. The most typical feature of G6PD deficiency is the presence of bizarre poikilocytes, with red cells that appear to have unevenly distributed hemoglobin ("hemighosts") and red cells that appear to have had parts of them bitten away ("bite cells" or "blister cells") (Fig. 129-7). A classical test, now rarely carried out, is supravital staining with methyl violet, which, if done promptly, reveals the presence of Heinz bodies (consisting of precipitates of denatured hemoglobin and hemichromes), which are regarded as a signature of oxidative damage to red cells (they are also seen with unstable hemoglobins). LDH is high, and so is the unconjugated

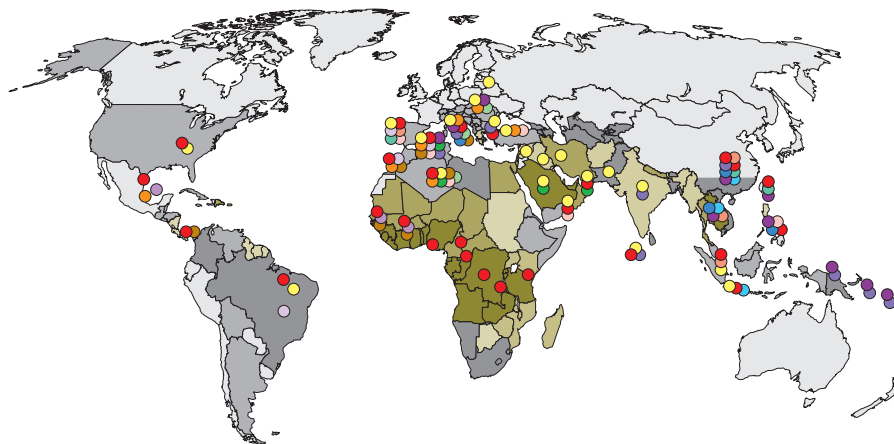


FIGURE 129-6 Epidemiology of glucose 6-phosphate dehydrogenase (G6PD) deficiency throughout the world. The different shadings indicate increasingly high levels of prevalence, up to about 20%; the different colored symbols indicate individual genetic variants of G6PD, each one having a different mutation. (From L Luzzatto et al, in C Scriver et al [eds]: *The Metabolic & Molecular Bases of Inherited Disease*, 8th ed. New York, McGraw-Hill, 2001.)

	Definite Risk	Possible Risk	Doubtful Risk
Antimalarials	Primaquine Dapsone/chlorproguanil ^a	Chloroquine	Quinine
Sulphonamides/sulphones	Sulfamethoxazole Others Dapsone	Sulfasalazine Sulfadimidine	Sulfisoxazole Sulfadiazine
Antibacterial/antibiotics	Cotrimoxazole Nalidixic acid Nitrofurantoin Niridazole	Ciprofloxacin Norfloxacin	Chloramphenicol <i>p</i> -Aminosalicylic acid
Antipyretic/analgesics	Acetanilide Phenazopyridine	Acetylsalicylic acid high dose (>3 g/d)	Acetylsalicylic acid (<3 g/d) Acetaminophen Phenacetin
Other	Naphthalene Methylene blue Rasburicase	Vitamin K analogues Ascorbic acid (>1 g)	Doxorubicin Probenecid

^aMarketed as Lapdap from 2003 to 2008.

bilirubin, indicating that there is also extravascular hemolysis. The most serious threat from AHA in adults is the development of acute renal failure (this is exceedingly rare in children). Once the threat of acute anemia is over and in the absence of comorbidity, full recovery from AHA associated with G6PD deficiency is the rule.

Although it was primaquine (PQ) that led to the discovery of G6PD deficiency, this drug has not been very prominent subsequently, because it is not necessary for the treatment of life-threatening *P. falciparum* malaria. Today there is a revival of interest in PQ because it is the only effective agent for eliminating the gametocytes of *P. falciparum* (thus preventing further transmission) and eliminating the hypnozoites of *Plasmodium vivax* (thus preventing endogenous relapse). In countries aiming to eliminate malaria, there may be a call for mass administration of PQ; this ought to be associated with G6PD testing. At the other end of the historic spectrum, the latest addition to the list of potentially hemolytic drugs (Table 129-5) is rasburicase; again G6PD testing ought to be made mandatory before giving this drug because fatal cases have been reported in newborns with kidney injury and in adults with tumor lysis syndrome.

A very small minority of subjects with G6PD deficiency have *chronic nonspherocytic hemolytic anemia* (CNSHA) of variable severity. The patient is nearly always a male, usually with a history of NNJ, who may present with anemia, unexplained jaundice, or gallstones later in life. The spleen may be enlarged. The severity of anemia ranges in different patients from borderline to transfusion dependent. The anemia is usually normomacrocytic, with reticulocytosis. Bilirubin and LDH are increased. Although hemolysis is, by definition, chronic in these patients, they are also vulnerable to acute oxidative damage, and therefore the same agents that can cause AHA in people with the ordinary type of G6PD deficiency will cause severe exacerbations in people with CNSHA associated with G6PD deficiency. In some cases of CNSHA, the deficiency of G6PD is so severe in granulocytes that it becomes rate-limiting for their oxidative burst, with consequent increased susceptibility to some bacterial infections.

Laboratory diagnosis The suspicion of G6PD deficiency can be confirmed by semiquantitative methods often referred to as screening tests, which are suitable for population studies and can correctly classify male subjects, in the steady state, as G6PD normal or G6PD deficient. However, in clinical practice, a diagnostic test is usually needed when the patient has had a hemolytic attack; this implies that the oldest, most G6PD-deficient red cells have been selectively destroyed, and young red cells, having higher G6PD activity, are being released into the circulation. Under these conditions, only a quantitative test can give a definitive result. In males, this test will identify normal hemizygotes and G6PD-deficient hemizygotes; among females, some heterozygotes will be missed, but those who are at most risk of hemolysis will be identified. Of course, G6PD deficiency also can be diagnosed by DNA testing.

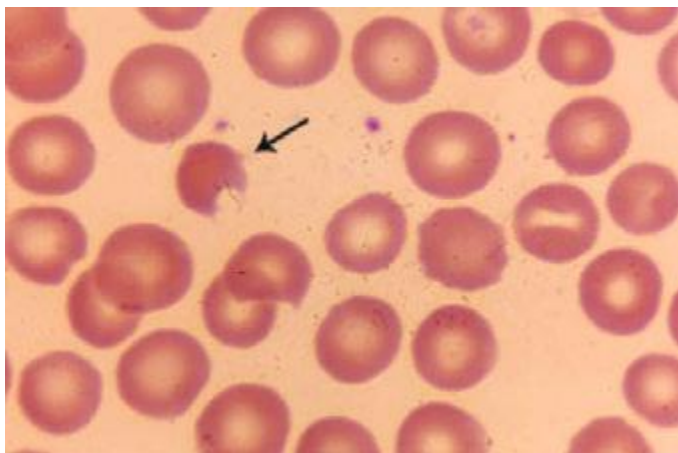


FIGURE 129-7 Peripheral blood smear from a glucose 6-phosphate dehydrogenase (G6PD)-deficient boy experiencing hemolysis. Note the red cells that are misshapen and called “bite” cells. (From MA Lichtman et al: *Lichtman's Atlas of Hematology*; <http://www.accessmedicine.com>. Copyright © The McGraw-Hill Companies, Inc. All rights reserved.)

TREATMENT G6PD DEFICIENCY

The AHA of G6PD deficiency is largely preventable by avoiding exposure to triggering factors of previously screened subjects. Of course, the practicability and cost-effectiveness of screening depend on the prevalence of G6PD deficiency in each individual community. Favism is entirely preventable in G6PD-deficient subjects by not eating fava beans. Drug-induced hemolysis can be prevented by testing for G6PD deficiency before prescribing; in most cases, one can use alternative drugs. When AHA develops and once its cause

is recognized, in most cases, no specific treatment is needed. However, if the anemia is severe, it may be a medical emergency, especially in children, requiring immediate action, including blood transfusion. This has been the case with an antimalarial drug combination containing dapsone (called Lapdap, introduced in 2003) that has caused severe acute hemolytic episodes in children with malaria in several African countries; after a few years, the drug was taken off the market. If there is acute renal failure, hemodialysis may be necessary, but if there is no previous kidney disease, recovery is the rule. The management of NNJ associated with G6PD deficiency is no different from that of NNJ due to other causes.

In cases with CNSHA, if the anemia is not severe, regular folic acid supplements and regular hematologic surveillance will suffice. It will be important to avoid exposure to potentially hemolytic drugs, and blood transfusion may be indicated when exacerbations occur, mostly in concomitance with intercurrent infection. In rare patients, regular blood transfusions may be required, in which case appropriate iron chelation should be instituted. Unlike in HS, there is no evidence of selective red cell destruction in the spleen; however, in practice, splenectomy has proven beneficial in severe cases.

Other abnormalities of the redox system As mentioned above, GSH is a key player in the defense against oxidative stress. Inherited defects of GSH metabolism are exceedingly rare, but each one can give rise to chronic HA (Table 129-4). A rare, peculiar, usually self-limited severe HA of the first month of life, called *infantile poikilocytosis*, may be associated with deficiency of glutathione peroxidase (GSHPX) due not to an inherited abnormality, but to transient nutritional deficiency of selenium, an element essential for the activity of GSHPX.

PYRIMIDINE 5'-NUCLEOTIDASE (P5N) DEFICIENCY P5N is a key enzyme in the catabolism of nucleotides arising from the degradation of nucleic acids that takes place in the final stages of erythroid cell maturation. How exactly its deficiency causes HA is not well understood, but a highly distinctive feature of this condition is a morphologic abnormality of the red cells known as *basophilic stippling*. The condition is rare, but it probably ranks third in frequency among red cell enzyme defects (after G6PD deficiency and PK deficiency). The anemia is lifelong, of variable severity, and may benefit from splenectomy.

Familial (Atypical) Hemolytic-Uremic Syndrome The term *familial (atypical) hemolytic-uremic syndrome* is used to designate a group of rare disorders, mostly affecting children, characterized by microangiopathic HA with presence of fragmented erythrocytes in the peripheral blood smear, thrombocytopenia (usually mild), and acute renal failure. (The word *atypical* is part of the phrase for historical reasons; it was hemolytic-uremic syndrome [HUS] caused by infection with *Escherichia coli* producing the Shiga toxin that was regarded as *typical*). The

genetic basis of atypical HUS (aHUS) has been elucidated. Studies of >100 families have revealed that those family members who developed HUS had mutations in any one of several genes encoding complement regulatory proteins: complement factor H (CFH), CD46 or membrane cofactor protein (MCP), complement factor I (CFI), complement component C3, complement factor B (CFB), and thrombomodulin. Thus, whereas all other inherited HAs are due to intrinsic red cell abnormalities, this group is unique in that hemolysis results from an inherited defect external to red cells (Table 129-1). Because the regulation of the complement cascade has considerable redundancy, in the steady state, any of the above abnormalities can be tolerated. However, when an intercurrent infection or some other trigger activates complement through the alternative pathway, the deficiency of one of the complement regulators becomes critical. Endothelial cells get damaged, especially in the kidney; at the same time, and partly as a result of this, there will be brisk hemolysis (thus, the more common Shiga toxin-related HUS [Chap. 149] can be regarded as a phenocopy of aHUS). aHUS is a severe disease, with up to 15% mortality in the acute phase and up to 50% of cases progressing to end-stage renal disease. Not infrequently, aHUS undergoes spontaneous remission; but because its basis is an inherited abnormality, it is not surprising that, given renewed exposure to a trigger, the syndrome will tend to recur; when it does, the prognosis is always serious. The standard treatment has been plasma exchange, which will supply the deficient complement regulator. The anti-C5 complement inhibitor eculizumab (see below) was found to greatly ameliorate the microangiopathic picture, with improvement in platelet counts and in renal function, thus abrogating the need for plasma exchange. It remains to be seen for how long eculizumab treatment will have to be continued in individual patients and whether it will influence the controversial issue of kidney (and liver) transplantation.

ACQUIRED HEMOLYTIC ANEMIA

Mechanical Destruction of Red Cells Although red cells are characterized by the remarkable deformability that enables them to squeeze through capillaries narrower than themselves for thousands of times in their lifetime, there are at least two situations in which they succumb to shear, if not to wear and tear; the result is intravascular hemolysis, resulting in hemoglobinuria (Table 129-6). One situation is acute and self-inflicted, *march hemoglobinuria*. Why sometimes a marathon runner may develop this complication, whereas on another occasion, this does not happen, we do not know (perhaps her or his footwear needs attention). A similar syndrome may develop after prolonged barefoot ritual dancing or intense playing of bongo drums. The other situation is chronic and iatrogenic (it has been called *microangiopathic hemolytic anemia*). It takes place in patients with prosthetic heart valves, especially when paraprosthetic regurgitation is present. If the hemolysis consequent on mechanical trauma to the red cells is mild, and if the supply of iron is adequate, the loss may be largely compensated; if more

TABLE 129-6 DISEASES AND CLINICAL SITUATIONS WITH PREDOMINANTLY INTRAVASCULAR HEMOLYSIS

	Onset/Time Course	Main Mechanism	Appropriate Diagnostic Procedure	Comments
Mismatched blood transfusion	Abrupt	Nearly always ABO incompatibility	Repeat cross-match	
Paroxysmal nocturnal hemoglobinuria (PNH)	Chronic with acute exacerbations	Complement (C)-mediated destruction of CD59(−) red cells	Flow cytometry to display a CD59(−) red cell population	Exacerbations due to C activation through any pathway
Paroxysmal cold hemoglobinuria (PCH)	Acute	Immune lysis of normal red cells	Test for Donath-Landsteiner antibody	Often triggered by viral infection
Septicemia	Very acute	Exotoxins produced by <i>Clostridium perfringens</i>	Blood cultures	Other organisms may be responsible
Microangiopathic	Acute or chronic	Red cell fragmentation	Red cell morphology on blood smear	Different causes ranging from endothelial damage to hemangioma to leaky prosthetic heart valve
March hemoglobinuria	Abrupt	Mechanical destruction	Targeted history taking	
Favism	Acute	Destruction of older fraction of G6PD-deficient red cells	G6PD assay	Triggered by ingestion of large dish of fava beans, but trigger can be infection or drug instead

Abbreviation: G6PD, glucose 6-phosphate dehydrogenase.

658 than mild anemia develops, reintervention to correct regurgitation may be required.

Infection By far the most frequent infectious cause of HA, in endemic areas, is malaria (Chap. 248). In other parts of the world, the most frequent direct cause is probably Shiga toxin-producing *E. coli* O157:H7, now recognized as the main etiologic agent of HUS, which is more common in children than in adults (Chap. 149). Life-threatening intravascular hemolysis, due to a toxin with lecithinase activity, occurs with *Clostridium perfringens* sepsis, particularly following open wounds, septic abortion, or as a disastrous accident due to a contaminated blood unit. Rarely, and if at all in children, HA is seen with sepsis or endocarditis from a variety of organisms. In addition, bacterial and viral infections can cause HA by indirect mechanisms (see above section on G6PD deficiency and Table 129-6).

Immune Hemolytic Anemias These can arise through at least two distinct mechanisms. (1) There is a true autoantibody directed against a red cell antigen, i.e., a molecule present on the surface of red cells. (2) When an antibody directed against a certain molecule (e.g., a drug) reacts with that molecule, red cells may get caught in the reaction, whereby they are damaged or destroyed. Because the antibodies involved differ in optimum reactivity temperatures, they are classified in the time-honored categories of “cold” and “warm” (Table 129-7). Autoantibody-mediated HAs may be seen in isolation (when they are called *idiopathic*) or as part of a systemic autoimmune disorder such as systemic lupus erythematosus. Here we discuss the most distinctive clinical pictures.

AUTOIMMUNE HEMOLYTIC ANEMIA (AIHA) Once a red cell is coated by an autoantibody (see [1] above), it will be destroyed by one or more mechanisms. In most cases, the Fc portion of the antibody will be recognized by the Fc receptor of macrophages, and this will trigger erythrophagocytosis. Thus, destruction of red cells will take place wherever macrophages are abundant, i.e., in the spleen, liver, or bone marrow; this is called *extravascular hemolysis* (Fig. 129-8). Because of the special anatomy of the spleen, this organ is particularly efficient in

trapping antibody-coated red cells, and often this is the predominant site of red cell destruction. In some cases, the nature of the antibody is such (usually an IgM antibody) that the antigen-antibody complex on the surface of red cells is able to activate complement (C); as a result, a large amount of membrane attack complex will form, and the red cells may be destroyed directly; this is known as *intravascular hemolysis*.

Clinical features AIHA is a serious condition; without appropriate treatment, it may have a mortality of approximately 10%. The onset is often abrupt and can be dramatic. The hemoglobin level can drop, within days, to as low as 4 g/dL; the massive red cell removal will produce jaundice; and sometimes the spleen is enlarged. When this triad is present, the suspicion of AIHA must be high. When hemolysis is (in part) intravascular, the telltale sign will be hemoglobinuria, which the patient may report or about which we must enquire or test for. The diagnostic test for AIHA is the direct antiglobulin test developed in 1945 by R. R. A. Coombs and known since by this name. The beauty of this test is that it detects directly the pathogenetic mediator of the disease, i.e., the presence of antibody on the red cells themselves. When the test is positive, it clinches the diagnosis; when it is negative, the diagnosis is unlikely. However, the sensitivity of the Coombs test varies depending on the technique that is used, and in doubtful cases, a repeat in a specialized lab is advisable; the term *Coombs-negative AIHA* is a last resort. In some cases, the autoantibody has a defined identity; it may be specific for an antigen belonging to the Rhesus system (it is often anti-e). In many cases, it is regarded as “nonspecific” because it reacts with virtually all types of red cells.

When AIHA develops in a person who is already known to have, for instance, systemic lupus or chronic lymphocytic leukemia (Table 129-7), we call it a complication; conversely, when AIHA presents on its own, it may be a pointer to an underlying condition that we ought to seek out. In both cases, what brings about AIHA remains, as in other autoimmune disorders, obscure. In some cases, AIHA can be associated, on first presentation or subsequently, with autoimmune thrombocytopenia (Evans’ syndrome).

TREATMENT AUTOIMMUNE HEMOLYTIC ANEMIA

Severe acute AIHA can be a medical emergency. The immediate treatment almost invariably includes transfusion of red cells. This may pose a special problem because, if the antibody involved is nonspecific, all of the blood units cross-matched will be incompatible. In these cases, it is often correct, paradoxically, to transfuse incompatible blood, with the rationale being that the transfused red cells will be destroyed no less but no more than the patient’s own red cells, but in the meantime, the patient stays alive. A situation like this requires close liaison and understanding between the clinical unit treating the patient and the blood transfusion/serology lab. Whenever the anemia is not immediately life-threatening, blood transfusion should be withheld (because compatibility problems may increase with each unit of blood transfused), and medical treatment started immediately with prednisone (1 mg/kg per day), which will produce a remission promptly in at least one-half of patients. Rituximab (anti-CD20) was regarded as second-line treatment, but it is increasingly likely that a relatively low dose (100 mg/wk × 4) of rituximab together with prednisone will become a first-line standard. It is especially encouraging that this approach seems to reduce the rate of relapse, a common occurrence in AIHA. For patients who do relapse or are refractory to medical treatment, one may have to consider splenectomy, which, although it does not cure the disease, can produce significant benefit by removing a major site of hemolysis, thus improving the anemia and/or reducing the need for other therapies (e.g., the dose of prednisone). Since the introduction of rituximab, azathioprine, cyclophosphamide, cyclosporine, and intravenous immunoglobulin have become second- or third-line agents. In very rare severe refractory cases, either autologous or allogeneic hematopoietic stem cell transplantation may have to be considered.

TABLE 129-7 CLASSIFICATION OF ACQUIRED IMMUNE HEMOLYTIC ANEMIAS

Clinical Setting	Type of Antibody	
	Cold, Mostly IgM, Optimal Temperature 4–30°C	Warm, Mostly IgG, Optimal Temperature 37°C; or Mixed
Primary	CAD	AIHA (idiopathic)
Secondary to viral infection	EBV	HIV
	CMV	Viral vaccines
	Other	
Secondary to other infection	Mycoplasma infection: paroxysmal cold hemoglobinuria	
Secondary to/associated with other disease	CAD in:	AIHA in:
	Waldenström’s disease	SLE
	Lymphoma	CLL
		Other malignancy
	Chronic inflammatory disorders (e.g., IBD)	
	After allogeneic HSCT	
Secondary to drugs: drug-induced immune hemolytic anemia	Small minority (e.g., with lenalidomide)	Majority: currently most common culprit drugs are cefotetan, ceftriaxone, piperacillin
	Drug-dependent: antibody destroys red cells only when drug present (e.g., rarely penicillin)	
	Drug-independent: antibody can destroy red cells even when drug no longer present (e.g., methyl dopa)	

Abbreviations: AIHA, autoimmune hemolytic anemia; CAD, cold agglutinin disease; CLL, chronic lymphocytic leukemia; CMV, cytomegalovirus; EBV, Epstein-Barr virus; HIV, human immunodeficiency virus; HSCT, hematopoietic stem cell transplantation; IBD, inflammatory bowel disease; SLE, systemic lupus erythematosus.

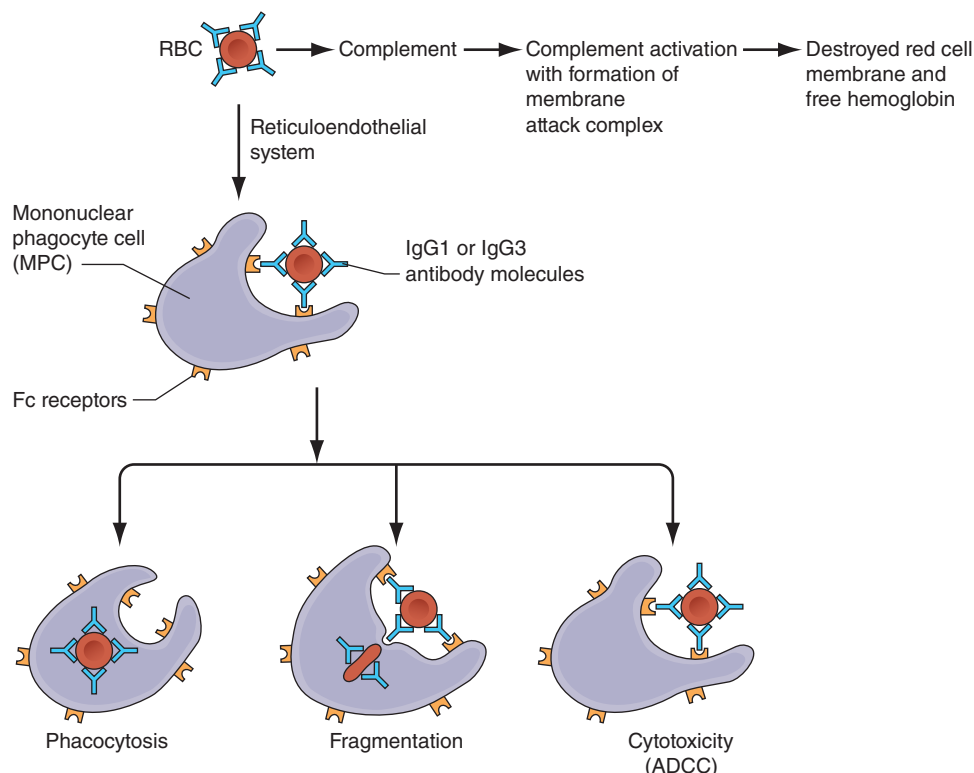


FIGURE 129-8 Mechanism of antibody-mediated immune destruction of red blood cells (RBCs). ADCC, antibody-dependent cell-mediated cytotoxicity. (From N Young et al: *Clinical Hematology*. Philadelphia, Elsevier, 2006; with permission.)

PAROXYSMAL COLD HEMOGLOBINURIA (PCH) PCH is a rather rare form of AIHA occurring mostly in children, usually triggered by a viral infection, usually self-limited, and characterized by the involvement of the so-called Donath-Landsteiner antibody. In vitro, this antibody has unique serologic features; it has anti-P specificity and binds to red cells only at a low temperature (optimally at 4°C), but when the temperature is shifted to 37°C, lysis of red cells takes place in the presence of complement. Consequently, in vivo there is intravascular hemolysis, resulting in hemoglobinuria. Clinically the differential diagnosis must include other causes of hemoglobinuria (Table 129-6), but the presence of the Donath-Landsteiner antibody will prove PCH. Active supportive treatment, including blood transfusion, is needed to control the anemia; subsequently, recovery is the rule.

COLD AGGLUTININ DISEASE (CAD) This designation is used for a form of chronic AIHA that usually affects the elderly and has special clinical and pathologic features. First, the term *cold* refers to the fact that the autoantibody involved reacts with red cells poorly or not at all at 37°C, whereas it reacts strongly at lower temperatures. As a result, hemolysis is more prominent the more the body is exposed to the cold. The antibody is usually IgM; usually it has an anti-I specificity (the I antigen is present on the red cells of almost everybody), and it may have a very high titer (1:100,000 or more has been observed). Second, the antibody is produced by an expanded clone of B lymphocytes, and sometimes its concentration in the plasma is high enough to show up as a spike in plasma protein electrophoresis, i.e., as a monoclonal gammopathy. Third, because the antibody is IgM, CAD is related to Waldenström's macroglobulinemia (WM) (Chap. 136), although in most cases, the other clinical features of this disease are not present. Thus, CAD must be regarded as a form of WM (i.e., as a low-grade mature B cell lymphoma) that manifests at an earlier stage precisely because the unique biologic properties of the IgM that it produces give the clinical picture of chronic HA.

In mild forms of CAD, avoidance of exposure to cold may be all that is needed to enable the patient to have a reasonably comfortable quality of life; but in more severe forms, the management of CAD is

not easy. Blood transfusion is not very effective because donor red cells are I positive and will be rapidly removed. Immunosuppressive/cytotoxic treatment with azathioprine or cyclophosphamide can reduce the antibody titer, but clinical efficacy is limited, and in view of the chronic nature of the disease, the side effects may prove unacceptable. Unlike in AIHA, prednisone and splenectomy are ineffective. Plasma exchange will remove antibody and is, therefore, in theory, a rational approach, but it is laborious and must be carried out at frequent intervals if it is to be beneficial. The management of CAD has changed significantly with the advent of rituximab; although its impact on CAD is not as great as on AIHA, up to 60% of patients respond, and remissions may be more durable with a rituximab-fludarabine combination. Given the long clinical course of CAD, it remains to be seen with what schedule or periodicity these agents will need to be administered.

Toxic Agents and Drugs A number of chemicals with oxidative potential, whether medicinal or not, can cause hemolysis even in people who are not G6PD deficient (see above). Examples are hyperbaric oxygen (or 100% oxygen), nitrates, chlorates, methylene blue, dapsone, cisplatin, and numerous aromatic (cyclic) compounds. Other chemicals may be hemolytic through nonoxidative, largely unknown mechanisms; examples include arsine, stibine, copper, and lead. The HA caused by lead poisoning is characterized by basophilic stippling; it is in fact a phenocopy of that seen in P5N deficiency (see above), suggesting it is mediated at least in part by lead inhibiting this enzyme.

In these cases, hemolysis appears to be mediated by a direct chemical action on red cells. But drugs can cause hemolysis through at least two other mechanisms. (1) A drug can behave as a hapten and induce antibody production; in rare subjects, this happens, for instance, with penicillin. Upon a subsequent exposure, red cells are caught, as innocent bystanders, in the reaction between penicillin and antipenicillin antibodies. Hemolysis will subside as soon as penicillin administration is stopped. (2) A drug can trigger, perhaps through mimicry, the production of an antibody against a red cell antigen. The best known example is methyldopa, an antihypertensive agent no longer in use, which in a small fraction of patients stimulated the production of the

660 Rhesus antibody anti-e. In patients who have this antigen, the anti-e is a true autoantibody, which then causes an autoimmune HA (see below). Usually this will gradually subside once methyl dopa is discontinued.

Severe intravascular hemolysis can be caused by the venom of certain snakes (cobras and vipers), and HA can also follow spider bites.

Paroxysmal Nocturnal Hemoglobinuria (PNH) PNH is an acquired chronic HA characterized by persistent intravascular hemolysis subject to recurrent exacerbations. In addition to hemolysis, there is often pancytopenia and a distinct tendency to venous thrombosis. This triad makes PNH a truly unique clinical condition; however, when not all of these three features are manifest on presentation, the diagnosis is often delayed, although it can always be made by appropriate laboratory investigations (see below).



PNH has about the same frequency in men and women and is encountered in all populations throughout the world, but it is a rare disease; its prevalence is estimated to be approximately 5 per million (it may be somewhat less rare in Southeast Asia and in the Far East). There is no evidence of inherited susceptibility. PNH has never been reported as a congenital disease, but it can present in small children or as late as in the seventies, although most patients are young adults.

CLINICAL FEATURES The patient may seek medical attention because, one morning, she or he passed blood instead of urine (Fig. 129-9). This distressing or frightening event may be regarded as the classical presentation; however, more frequently, this symptom is not noticed or is suppressed. Indeed, the patient often presents simply as a problem in the differential diagnosis of *anemia*, whether symptomatic or discovered incidentally. Sometimes, the anemia is associated from the outset with neutropenia, thrombocytopenia, or both, thus signaling an element of bone marrow failure (see below). Some patients may present with recurrent attacks of severe abdominal pain defying a specific diagnosis and eventually found to be related to thrombosis. When thrombosis affects the hepatic veins, it may produce acute hepatomegaly and ascites, i.e., a full-fledged Budd-Chiari syndrome, which, in the absence of liver disease, ought to raise the suspicion of PNH.

The *natural history* of PNH can extend over decades. Without treatment, the median survival is estimated to be about 8–10 years; in the past, the most common cause of death has been venous thrombosis, followed by infection secondary to severe neutropenia and hemorrhage secondary to severe thrombocytopenia. Rarely (estimated 1–2% of all cases), PNH may terminate in acute myeloid leukemia. On the other hand, full spontaneous recovery from PNH has been documented, albeit rarely.

LABORATORY INVESTIGATIONS AND DIAGNOSIS The most consistent blood finding is anemia, which may range from mild to moderate to very severe. The anemia is usually normomacrocyclic, with unremarkable red cell morphology. If the MCV is high, it is usually largely accounted

for by reticulocytosis, which may be quite marked (up to 20%, or up to 400,000/ μ L). The anemia may become microcytic if the patient is allowed to become iron deficient as a result of chronic urinary blood loss through hemoglobinuria. Unconjugated bilirubin is mildly or moderately elevated; LDH is typically markedly elevated (values in the thousands are common); and haptoglobin is usually undetectable. All of these findings make the diagnosis of hemolytic anemia compelling. Hemoglobinuria may be overt in a random urine sample; if it is not, it may be helpful to obtain serial urine samples, because hemoglobinuria can vary dramatically from day to day and even from hour to hour. The bone marrow is usually cellular, with marked to massive erythroid hyperplasia, often with mild to moderate dyserythropoietic features (not to be confused with myelodysplastic syndrome). At some stage of the disease, the marrow may become hypocellular or even frankly aplastic (see below).

The definitive diagnosis of PNH must be based on the demonstration that a substantial proportion of the patient's red cells have an increased susceptibility to complement (C), due to the deficiency on their surface of proteins (particularly CD59 and CD55) that normally protect the red cells from activated C. The sucrose hemolysis test is unreliable; in contrast, the acidified serum (Ham) test is highly reliable but is carried out only in a few labs. The gold standard today is flow cytometry, which can be carried out on granulocytes as well as on red cells. A bimodal distribution of cells, with a discrete population that is CD59 and CD55 negative, is diagnostic of PNH. In PNH patients, this population is at least 5% of the total red cells and at least 20% of the total granulocytes.

PATHOPHYSIOLOGY Hemolysis in PNH is mainly intravascular and is due to an intrinsic abnormality of the red cell, which makes it exquisitely sensitive to activated C, whether it is activated through the alternative pathway or through an antigen-antibody reaction. The former mechanism is mainly responsible for chronic hemolysis in PNH; the latter explains why the hemolysis can be dramatically exacerbated in the course of a viral or bacterial infection. Hypersusceptibility to C is due to deficiency of several protective membrane proteins (Fig. 129-10), of which CD59 is the most important, because it hinders the insertion into the membrane of C9 polymers. The molecular basis for the deficiency of these proteins has been pinpointed not to a defect in any of the respective genes, but rather to the shortage of a unique glycolipid molecule, GPI (Fig. 129-2), which, through a peptide bond, anchors these proteins to the surface membrane of cells. The shortage of GPI is due in turn to a mutation in an X-linked gene, called *PIG-A*, required for an early step in GPI biosynthesis. In virtually each patient, the *PIG-A* mutation is different. This is not surprising, because these mutations are not inherited; rather, each one takes place *de novo* in a hemopoietic stem cell (i.e., they are somatic mutations). As a result, the patient's marrow is a mosaic of mutant and nonmutant cells, and the peripheral blood always contains both PNH cells and normal (non-PNH) cells. Thrombosis is one of the most immediately life-threatening complications of PNH and yet one of the least understood in its pathogenesis. It could be that deficiency of CD59 on the PNH platelet causes inappropriate platelet activation; however, other mechanisms are possible.

BONE MARROW FAILURE (BMF) AND RELATIONSHIP BETWEEN PNH AND APLASTIC ANEMIA (AA) It is not unusual that patients with firmly established PNH have a previous history of well-documented AA; indeed, BMF preceding overt PNH is probably the rule rather than the exception. On the other hand, sometimes a patient with PNH becomes less hemolytic and more pancytopenic and ultimately has the clinical picture of AA. Because AA is probably an organ-specific autoimmune disease, in which T cells cause damage to hematopoietic stem cells, the same may be true of PNH, with the specific proviso that the damage spares PNH stem cells. *PIG-A* mutations can be demonstrated in normal people, and there is evidence from mouse models that PNH stem cells do not expand when the rest of the bone marrow is normal. Thus, we can visualize PNH as always having two components: failure of normal hematopoiesis and massive expansion of a PNH clone. Findings supporting this notion include skewing of the T cell repertoire and the demonstration of GPI-reactive T cells in patients with PNH.



FIGURE 129-9 Consecutive urine samples from a patient with paroxysmal nocturnal hemoglobinuria (PNH). The variation in the severity of hemoglobinuria within hours is probably unique to this condition.

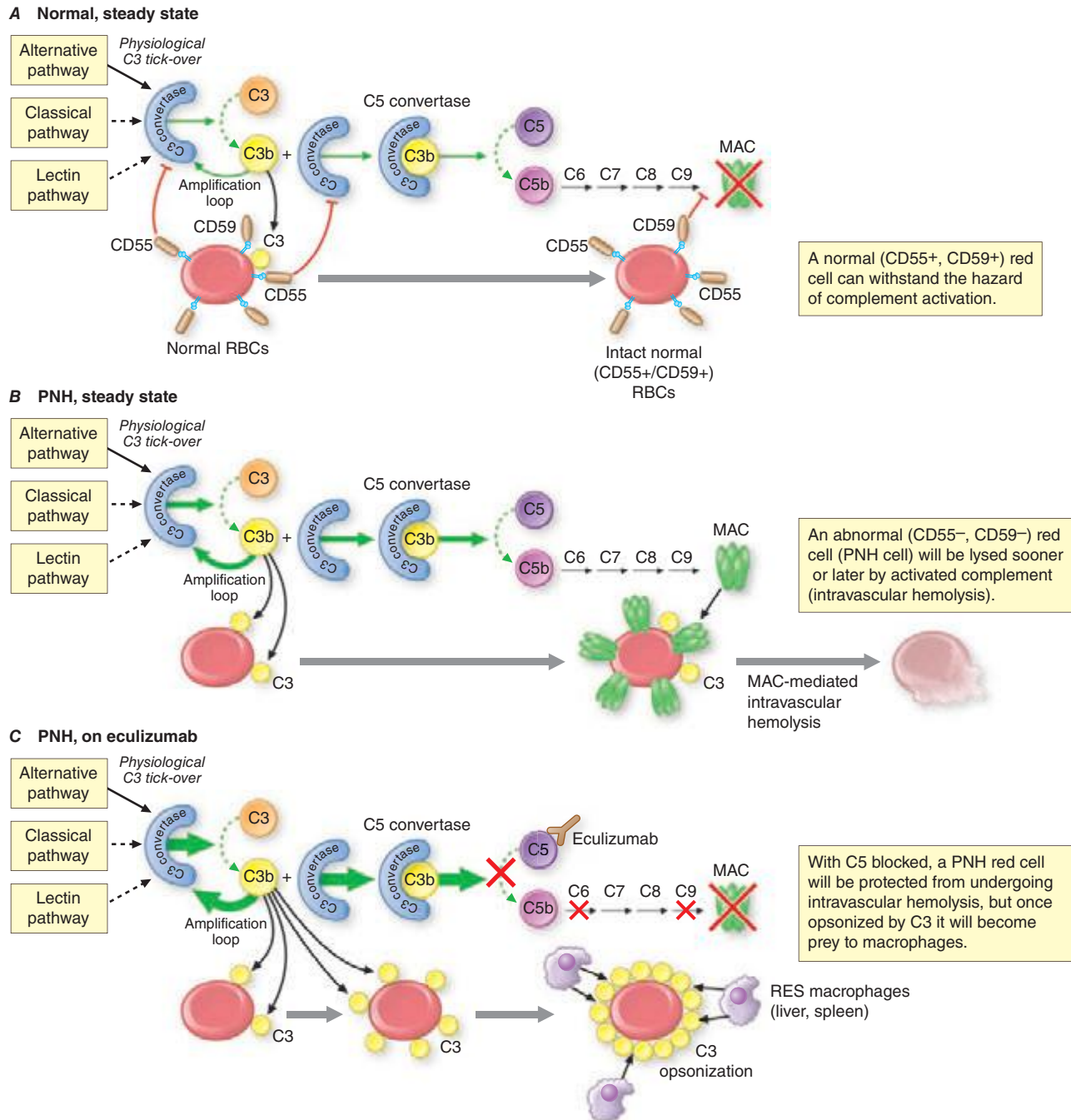


FIGURE 129-10 The complement cascade and the fate of red cells. **A.** Normal red cells are protected from complement activation and subsequent hemolysis by CD55 and CD59. These two proteins, being GPI-linked, are missing from the surface of PNH red cells as a result of a somatic mutation of the X-linked *PIG-A* gene that encodes a protein required for an early step of the GPI molecule biosynthesis. **B.** In the steady state, PNH erythrocytes suffer from spontaneous (tick-over) complement activation, with consequent intravascular hemolysis through formation of the membrane attack complex (MAC); when extra complement is activated through the classical pathway, an exacerbation of hemolysis will result. **C.** On eculizumab, PNH erythrocytes are protected from hemolysis from the inhibition of C5 cleavage; however, upstream complement activation may lead to C3 opsonization and possible extravascular hemolysis. GPI, glycosylphosphatidylinositol; PNH, paroxysmal nocturnal hemoglobinuria. (From L Luzzatto et al: *Haematologica* 95:523, 2010.)

TREATMENT PAROXYSMAL NOCTURNAL HEMOGLOBINURIA

Unlike other acquired hemolytic anemias, PNH may be a lifelong condition, and most patients receive supportive treatment only, including transfusion of filtered red cells¹ whenever necessary,

¹Now that filters with excellent retention of white cells are routinely used, the traditional washing of red cells, aiming to avoid white cell reactions triggering hemolysis, is no longer necessary and is wasteful.

which, for some patients, means quite frequently. Folic acid supplements (at least 3 mg/d) are mandatory; the serum iron should be checked periodically, and iron supplements should be administered as appropriate. Long-term glucocorticoids are not indicated because there is no evidence that they have any effect on chronic hemolysis; in fact, they are contraindicated because their side effects are considerable and potentially dangerous. A major advance in the management of PNH has been the development of a humanized monoclonal antibody, eculizumab, which binds to the complement component C5 near the site that, when cleaved, will trigger the distal part of the complement cascade leading to

662 the formation of membrane attack complex (MAC). In an international, placebo-controlled, randomized trial of 87 patients (so far the only controlled therapeutic trial in PNH) who had been selected on grounds of having severe hemolysis making them transfusion-dependent, eculizumab proved effective and was licensed in 2007. Eculizumab, by abrogating complement-dependent intravascular hemolysis, significantly improves the quality of life of PNH patients. One would expect that the need for blood transfusion would also be abrogated; indeed, this is the case in about one-half of patients, in many of whom there is also a rise in hemoglobin levels. In the remaining patients, however, the anemia remains sufficiently severe to require blood transfusion. One reason for this is that, once the distal complement pathway is blocked, red cells no longer destroyed by the MAC become opsonized by complement (C3) fragments and undergo extravascular hemolysis (Fig. 129-10). The extent to which this happens depends in part on a genetic polymorphism of the complement receptor CR1. Based on its half-life, eculizumab must be administered intravenously every 14 days. The only form of treatment that currently can provide a definitive cure for PNH is allogeneic BMT. When an HLA-identical sibling is available, BMT should be offered to any young patient with severe PNH; the availability of eculizumab has decreased significantly the proportion of patients receiving BMT.

For patients with the PNH-AA syndrome, immunosuppressive treatment with antithymocyte globulin and cyclosporine A may be indicated, especially in order to relieve severe thrombocytopenia and/or neutropenia in patients in whom these were the main problem(s); of course, this treatment will have little or no effect on hemolysis. Any patient who has had venous thrombosis or who has a genetically determined thrombophilic state in addition to PNH should be on regular anticoagulant prophylaxis. With thrombotic complications that do not resolve otherwise, thrombolytic treatment with tissue plasminogen activator may be indicated.

ANEMIA DUE TO ACUTE BLOOD LOSS

Blood loss causes anemia by two main mechanisms: (1) by the direct loss of red cells; and (2) if the loss of blood is protracted, it will gradually deplete iron stores, eventually resulting in iron deficiency. The latter type of anemia is covered in [Chap. 126](#); here we are concerned with the former type, i.e., *posthemorrhagic anemia*, which follows *acute* blood loss. This can be *external* (e.g., after trauma or obstetric hemorrhage) or *internal* (e.g., from bleeding in the gastrointestinal tract, rupture of the spleen, rupture of an ectopic pregnancy, subarachnoid hemorrhage). In any of these cases, after the sudden loss of a large amount of blood, there are three clinical/pathophysiologic stages. (1) At first, the dominant feature is hypovolemia, which poses a threat particularly to organs that normally have a high blood supply, like the brain and the kidneys; therefore, loss of consciousness and acute renal failure are major threats. It is important to note that at this stage an ordinary blood count will not show anemia, because the hemoglobin concentration is not affected. (2) Next, as an emergency response, baroreceptors and stretch receptors will cause release of vasopressin and other peptides, and the body will shift fluid from the extravascular to the intravascular compartment, producing hemodilution; thus, the hypovolemia gradually converts to anemia. The degree of anemia will reflect the amount of blood lost. If after 3 days the hemoglobin is, for example, 7 g/dL, it means that about half of the entire blood has been lost. (3) Provided bleeding does not continue, the bone marrow response will gradually ameliorate the anemia.

The diagnosis of acute posthemorrhagic anemia (APHA) is usually straightforward, although sometimes internal bleeding episodes (e.g., after a traumatic injury), even when large, may not be immediately obvious. Whenever an abrupt fall in hemoglobin has taken place, whatever history is given by the patient, APHA should be suspected. Supplementary history may have to be obtained by asking the appropriate questions, and appropriate investigations (e.g., a sonogram or an endoscopy) may have to be carried out.

TREATMENT ANEMIA DUE TO ACUTE BLOOD LOSS

With respect to treatment, a two-pronged approach is imperative. (1) In many cases, the blood lost needs to be replaced promptly. Unlike with many chronic anemias, when finding and correcting the cause of the anemia is the first priority and blood transfusion may not be even necessary because the body is adapted to the anemia, with acute blood loss the reverse is true; because the body is not adapted to the anemia, blood transfusion takes priority. (1) While the emergency is being confronted, it is imperative to stop the hemorrhage and to eliminate its source.

A special type of APHA is blood loss during and immediately after surgery, which can be substantial (e.g., up to 2 L in the case of a radical prostatectomy). Of course with elective surgical procedures, the patient's own stored blood may be available (through preoperative autologous blood donation), and in any case, blood loss ought to have been carefully monitored/measured. The fact that this blood loss is iatrogenic dictates that ever more effort should be invested in optimizing its management.

A Holy Grail of emergency medicine for a long time has been the idea of a blood substitute that would be universally available, suitable for all recipients, easy to store and to transport, safe, and as effective as blood itself. Two main paths have been pursued: (1) fluorocarbon synthetic chemicals that bind oxygen reversibly, and (2) artificially modified hemoglobins, known as hemoglobin-based oxygen carriers (HBOCs). Although there are numerous anecdotal reports of the use of both approaches in humans, and although HBOCs have reached the stage of phase 2–3 clinical trials, no “blood substitute” has yet become standard treatment.

130 Bone Marrow Failure Syndromes Including Aplastic Anemia and Myelodysplasia

Neal S. Young

The hypoproliferative anemias are normochromic, normocytic, or macrocytic and are characterized by a low reticulocyte count. Hypoproliferative anemia is also a prominent feature of hematologic diseases that are described as bone marrow failure states; these include aplastic anemia, myelodysplastic syndrome (MDS), pure red cell aplasia (PRCA), and myelophthisis. Anemia in these disorders is often not a solitary or even the major hematologic finding. More frequent in bone marrow failure is *pancytopenia*: anemia, leukopenia, and thrombocytopenia. Low blood counts in the marrow failure diseases result from deficient hematopoiesis, as distinguished from blood count depression due to peripheral destruction of red cells (hemolytic anemias), platelets (idiopathic thrombocytopenic purpura [ITP] or due to splenomegaly), and granulocytes (as in the immune leukopenias). Marrow damage and dysfunction also may be secondary to infection, inflammation, or cancer.

Hematopoietic failure syndromes are classified by dominant morphologic features of the bone marrow ([Table 130-1](#)). Although practical distinction among these syndromes usually is clear, some processes are so closely related that the diagnosis may be complex. Patients may seem to suffer from two or three related diseases simultaneously, or one diagnosis may appear to evolve into another. Many of these syndromes share an immune-mediated mechanism of marrow destruction and some element of genomic instability resulting in a higher rate of malignant transformation.

It is important that the internist and general practitioner recognize the marrow failure syndromes, as their prognosis may be poor if the

TABLE 130-1 DIFFERENTIAL DIAGNOSIS OF PANCYTOPENIA

Pancytopenia with Hypocellular Bone Marrow	
Acquired aplastic anemia	
Constitutional aplastic anemia (Fanconi anemia, dyskeratosis congenita)	
Some myelodysplasia	
Rare aleukemic leukemia	
Some acute lymphoid leukemia	
Some lymphomas of bone marrow	
Pancytopenia with Cellular Bone Marrow	
Primary bone marrow diseases	Secondary to systemic diseases
Myelodysplasia	Systemic lupus erythematosus
Paroxysmal nocturnal hemoglobinuria	Hypersplenism
Myelofibrosis	B ₁₂ folate deficiency
Some aleukemic leukemia	Overwhelming infection
Myelophthisis	Alcohol
Bone marrow lymphoma	Brucellosis
Hairy cell leukemia	Sarcoidosis
	Tuberculosis
	Leishmaniasis
Hypocellular Bone Marrow ± Cytopenia	
Q fever	
Legionnaires' disease	
Anorexia nervosa, starvation	
<i>Mycobacterium</i>	

patient is untreated; effective therapies are often available but sufficiently complicated in their choice and delivery so as to warrant the care of a hematologist or oncologist.

APLASTIC ANEMIA

DEFINITION

Aplastic anemia is pancytopenia with bone marrow hypocellularity. Acquired aplastic anemia is distinguished from iatrogenic aplasia, marrow hypocellularity after intensive cytotoxic chemotherapy for cancer. Aplastic anemia can also be constitutional: the genetic diseases Fanconi anemia and dyskeratosis congenita, although frequently associated with typical physical anomalies and the development of pancytopenia early in life, can also present as marrow failure in normal-appearing adults. Acquired aplastic anemia is often stereotypical in its manifestations, with the abrupt onset of low blood counts in a previously well young adult; seronegative hepatitis or a course of an incriminated medical drug may precede the onset. The diagnosis in these instances is uncomplicated. Sometimes blood count depression is moderate or incomplete, resulting in anemia, leukopenia, and thrombocytopenia in some combination. Aplastic anemia is related to both paroxysmal nocturnal hemoglobinuria (PNH; [Chap. 129](#)) and to MDS, and in some cases, a clear distinction among these disorders may not be possible.

EPIDEMIOLOGY



The incidence of acquired aplastic anemia in Europe and Israel is two cases per million persons annually. In Thailand and China, rates of five to seven per million have been established. In general, men and women are affected with equal frequency, but the age distribution is biphasic, with the major peak in the teens and twenties and a second rise in older adults.

ETIOLOGY

The origins of aplastic anemia have been inferred from several recurring clinical associations ([Table 130-2](#)); unfortunately, these relationships are not reliable in an individual patient and may not be etiologic. In addition, although most cases of aplastic anemia are idiopathic, little other than history separates these cases from those with a presumed etiology such as a drug exposure.

TABLE 130-2 CLASSIFICATION OF APLASTIC ANEMIA AND SINGLE CYTOPENIAS

Acquired	Inherited
Aplastic Anemia	
Secondary	Fanconi anemia
Radiation	Dyskeratosis congenita
Drugs and chemicals	Shwachman-Diamond syndrome
Regular effects	Reticular dysgenesis
Idiosyncratic reactions	Amegakaryocytic thrombocytopenia
Viruses	Familial aplastic anemias
Epstein-Barr virus (infectious mononucleosis)	Preleukemia (monosomy 7, etc.)
Hepatitis (non-A, non-B, non-C hepatitis)	Nonhematologic syndrome (Down, Dubowitz, Seckel)
Parvovirus B19 (transient aplastic crisis, PRCA)	
HIV-1 (AIDS)	
Immune diseases	
Eosinophilic fasciitis	
Hyperimmunoglobulinemia	
Large granular lymphocytosis (LGL)	
Thymoma/thymic carcinoma	
Graft-versus-host disease in immunodeficiency	
Paroxysmal nocturnal hemoglobinuria (PNH)	
Pregnancy	
Idiopathic	
Cytopenias	
PRCA (see Table 130-4)	Congenital PRCA (Diamond-Blackfan anemia)
Neutropenia/agranulocytosis	
Idiopathic	Kostmann syndrome
Drugs, toxins	Shwachman-Diamond syndrome
LGL	Reticular dysgenesis
Pure white cell aplasia (+/- thymoma)	
Thrombocytopenia	
Drugs, toxins	Amegakaryocytic thrombocytopenia
Idiopathic amegakaryocytic	Thrombocytopenia with absent radii

Abbreviation: PRCA, pure red cell aplasia.

Radiation Marrow aplasia is a major acute sequela of radiation. Radiation damages DNA; tissues dependent on active mitosis are particularly susceptible. Nuclear accidents involve not only power plant workers but also employees of hospitals, laboratories, and industry (food sterilization, metal radiography, etc.), as well as innocents exposed to stolen, misplaced, or misused sources. Whereas the radiation dose can be approximated from the rate and degree of decline in blood counts, dosimetry by reconstruction of the exposure can help to estimate the patient's prognosis and also to protect medical personnel from contact with radioactive tissue and excreta. MDS and leukemia, but probably not aplastic anemia, are late effects of radiation.

Chemicals Benzene is a notorious cause of bone marrow failure: epidemiologic, clinical, and laboratory data link benzene to aplastic anemia, acute leukemia, and blood and marrow abnormalities. For leukemia, incidence is correlated with cumulative exposure, but susceptibility must also be important, because only a minority of even heavily exposed workers develop myelotoxicity. The employment history is important, especially in industries where benzene is used for a secondary purpose, usually as a solvent. Benzene-related blood diseases have declined with regulation of industrial exposure. Although benzene is no longer generally available as a household solvent, exposure to its metabolites occurs in the normal diet and in the environment. The association between marrow failure and other chemicals is much less well substantiated.

TABLE 130-3 SOME DRUGS AND CHEMICALS ASSOCIATED WITH APLASTIC ANEMIA

Agents that regularly produce marrow depression as major toxicity in commonly used doses or normal exposures:

Cytotoxic drugs used in cancer chemotherapy: *alkylating agents, antimetabolites, antimicrobials*, some antibiotics

Agents that frequently but not inevitably produce marrow aplasia:

Benzene

Agents associated with aplastic anemia but with a relatively low probability:

Chloramphenicol

Insecticides

Antiprotozoals: *quinacrine* and chloroquine, mepacrine

Nonsteroidal anti-inflammatory drugs (including *phenylbutazone*, indomethacin, ibuprofen, sulindac, aspirin)

Anticonvulsants (*hydantoins, carbamazepine, phenacemide, felbamate*)

Heavy metals (*gold, arsenic, bismuth, mercury*)

Sulfonamides: some antibiotics, antithyroid drugs (methimazole, methylthiouracil, propylthiouracil), antidiabetes drugs (tolbutamide, chlorpropamide), carbonic anhydrase inhibitors (acetazolamide and methazolamide)

Antihistamines (*cimetidine, chlorpheniramine*)

D-Penicillamine

Estrogens (in pregnancy and in high doses in animals)

Agents whose association with aplastic anemia is more tenuous:

Other antibiotics (streptomycin, tetracycline, methicillin, mebendazole, trimethoprim/sulfamethoxazole, flucytosine)

Sedatives and tranquilizers (chlorpromazine, prochlorperazine, piperacetazine, chlordiazepoxide, meprobamate, methyprylon)

Allopurinol

Methyl dopa

Quinidine

Lithium

Guanidine

Potassium perchlorate

Thiocyanate

Carbimazole

Note: Terms set in italics show the most consistent association with aplastic anemia.

Drugs (Table 130-3) Many chemotherapeutic drugs have marrow suppression as a major toxicity; effects are dose dependent and will occur in all recipients. In contrast, idiosyncratic reactions to a large and diverse group of drugs may lead to aplastic anemia without a clear dose-response relationship. These associations rest largely on accumulated case reports until a large international study in Europe in the 1980s quantitated drug relationships, especially for nonsteroidal analgesics, sulfonamides, thyrostatic drugs, some psychotropics, penicillamine, allopurinol, and gold. Association does not equal causation: a drug may have been used to treat the first symptoms of bone marrow failure (antibiotics for fever or the preceding viral illness) or provoked the first symptom of a preexisting disease (petechiae by nonsteroidal anti-inflammatory agents administered to the thrombocytopenic patient). In the context of total drug use, idiosyncratic reactions, although individually devastating, are rare events. Risk estimates are usually lower when determined in population-based studies. Furthermore, the low absolute risk is also made more obvious: even a 10- or 20-fold increase in risk translates, in a rare disease, to just a handful of drug-induced aplastic anemia cases among hundreds of thousands of exposed persons.

Infections Hepatitis is the most common preceding infection, and posthepatitis marrow failure accounts for approximately 5% of etiologies in most series. Patients are usually young men who have recovered from a bout of liver inflammation 1 to 2 months earlier; the subsequent pancytopenia is very severe. The hepatitis is seronegative (non-A, non-B, non-C) and possibly due to an as yet undiscovered infectious agent. Fulminant liver failure in childhood also follows seronegative hepatitis, and marrow failure occurs at a high rate in these patients. Aplastic

anemia can rarely follow infectious mononucleosis. Parvovirus B19, the cause of transient aplastic crisis in hemolytic anemias and of some PRCAs (see below), does not usually cause generalized bone marrow failure. Mild blood count depression is frequent in the course of many viral and bacterial infections but resolves with the infection.

Immunologic Diseases Aplasia is a major consequence and the inevitable cause of death in *transfusion-associated graft-versus-host disease* (GVHD) that can occur after infusion of nonirradiated blood products to an immunodeficient recipient. Aplastic anemia is strongly associated with the rare collagen vascular syndrome *eosinophilic fasciitis* that is characterized by painful induration of subcutaneous tissues (**Chap. 382**). Thymoma and hypogammaglobulinemia are occasional associations with aplastic anemia. Pancytopenia with marrow hypoplasia can also occur in systemic lupus erythematosus (SLE).

Pregnancy Aplastic anemia very rarely may occur and recur during pregnancy and resolve with delivery or with spontaneous or induced abortion.

Paroxysmal Nocturnal Hemoglobinuria An acquired mutation in the *PIG-A* gene in a hematopoietic stem cell is required for the development of PNH, but *PIG-A* mutations probably occur commonly in normal individuals. If the *PIG-A* mutant stem cell proliferates, the result is a clone of progeny deficient in glycosylphosphatidylinositol-linked cell surface membrane proteins (**Chap. 129**). Small clones of deficient cells can be detected by sensitive flow cytometry tests in one-half or more of patients with aplastic anemia at the time of presentation. Functional studies of bone marrow from PNH patients, even those with mainly hemolytic manifestations, show evidence of defective hematopoiesis. Patients with an initial clinical diagnosis of PNH, especially younger individuals, may later develop frank marrow aplasia and pancytopenia; patients with an initial diagnosis of aplastic anemia may suffer from hemolytic PNH years after recovery of blood counts.

Constitutional Disorders Fanconi anemia, an autosomal recessive disorder, manifests as congenital developmental anomalies, progressive pancytopenia, and an increased risk of malignancy. Chromosomes in Fanconi anemia are peculiarly susceptible to DNA cross-linking agents, the basis for a diagnostic assay. Patients with Fanconi anemia typically have short stature, café au lait spots, and anomalies involving the thumb, radius, and genitourinary tract. At least 16 different genetic defects (all but one with an identified gene) have been defined; the most common, type A Fanconi anemia, is due to a mutation in *FANCA*. Most of the Fanconi anemia gene products form a protein complex that activates FANCD2 by monoubiquitination to play a role in the cellular response to DNA damage and especially interstrand cross-linking.

Dyskeratosis congenita is characterized by the triad of mucous membrane leukoplakia, dystrophic nails, reticular hyperpigmentation, and with the development of aplastic anemia in childhood. Dyskeratosis is due to mutations in genes of the telomere repair complex, which acts to maintain telomere length in replicating cells: the X-linked variety is due to mutations in the *DKC1* (*dyskerin*) gene; the more unusual autosomal dominant type is due to mutation in *TERC*, which encodes an RNA template, and *TERT*, which encodes the catalytic reverse transcriptase, telomerase. Mutations in *TNF2*, a component of the shelterin complex, proteins that bind the telomere DNA, also occur.

In Shwachman-Diamond syndrome, presentation is early in life with neutropenia with pancreatic insufficiency and malabsorption; most patients have compound heterozygous mutations in *SBDS* that may affect both ribosomal biogenesis (as in Diamond-Blackfan anemia; see below) and marrow stroma function.

While these constitutional syndromes can on occasion present in adults, genetic mutations are also risk factors for bone marrow failure. In the recently recognized telomeropathies, mutations in *TERT* and *TERC* have subtle effects on hematopoietic function. Typical presentations include not only severe but also moderate aplastic anemia, which can be chronic and not progressive, and isolated macrocytic anemia or thrombocytopenia. Physical anomalies are usually not found in

the patient, although early hair graying is a clue to the diagnosis. A careful family history may disclose pulmonary fibrosis and hepatic cirrhosis. Specific involvement of the bone marrow, liver, and lung is highly variable, as is penetrance of clinical phenotype, both within families and among kindreds. Variable penetrance means that *TERT* and *TERC* mutations represent risk factors for marrow failure, as family members with the same mutations may have normal or only slight hematologic abnormalities but more subtle evidence of (compensated) hematopoietic insufficiency.

PATHOPHYSIOLOGY

Bone marrow failure results from severe damage to the hematopoietic cell compartment. In aplastic anemia, replacement of the bone marrow by fat is apparent in the morphology of the biopsy specimen (Fig. 130-1) and magnetic resonance imaging (MRI) of the spine. Cells bearing the CD34 antigen, a marker of early hematopoietic cells, are greatly diminished, and in functional studies, committed and primitive progenitor cells are virtually absent; in vitro assays have suggested that the stem cell pool is reduced to $\leq 1\%$ of normal in severe disease at the time of presentation.

An intrinsic stem cell defect exists for the constitutional aplastic anemias: cells from patients with Fanconi anemia exhibit chromosome damage and death on exposure to certain chemical agents. Telomeres are short in some patients with aplastic anemia, due to heterozygous mutations in genes of the telomere repair complex. Telomeres may also shorten physiologically in acquired marrow failure due to replicative demands on a limited stem cell pool.

Drug Injury Extrinsic damage to the marrow follows massive physical or chemical insults such as high doses of radiation and toxic chemicals. For the more common idiosyncratic reaction to modest doses of medical drugs, altered drug metabolism has been invoked as a likely mechanism. The metabolic pathways of many drugs and chemicals, especially if they are polar and have limited water solubility, involve enzymatic degradation to highly reactive electrophilic compounds; these intermediates are toxic because of their propensity to bind to cellular macromolecules. For example, derivative hydroquinones and quinolones are responsible for benzene-induced tissue injury. Excessive generation of toxic intermediates or failure to detoxify the intermediates may be genetically determined and apparent only on specific drug challenge; the complexity and specificity of the pathways imply multiple susceptibility loci and would provide an explanation for the rarity of idiosyncratic drug reactions.

Immune-Mediated Injury The recovery of marrow function in some patients prepared for bone marrow transplantation with antilymphocyte globulin first suggested that aplastic anemia might be immune mediated. Consistent with this hypothesis was the frequent failure of simple bone marrow transplantation from a syngeneic twin, without conditioning cytotoxic chemotherapy, which also argued both *against* simple stem cell absence as the cause and *for* the presence of a host factor producing marrow failure. Laboratory data support an important role for the immune system in aplastic anemia. Blood and bone marrow cells of patients can suppress normal hematopoietic progenitor cell growth, and removal of T cells from aplastic anemia bone marrow improves colony formation in vitro. Increased numbers of activated

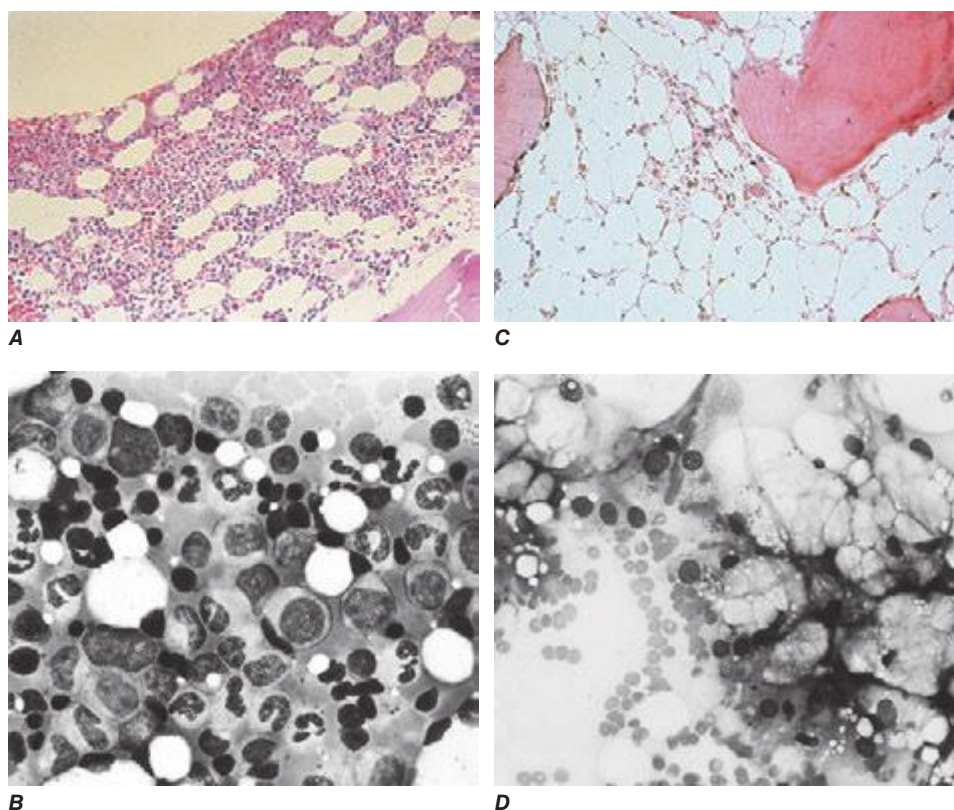


FIGURE 130-1 Normal and aplastic bone marrow. **A.** Normal bone marrow biopsy. **B.** Normal bone marrow aspirate smear. The marrow is normally 30–70% cellular, and there is a heterogeneous mix of myeloid, erythroid, and lymphoid cells. **C.** Aplastic anemia biopsy. **D.** Marrow smear in aplastic anemia. The marrow shows replacement of hematopoietic tissue by fat and only residual stromal and lymphoid cells.

cytotoxic T cell clones are observed in aplastic anemia patients and usually decline with successful immunosuppressive therapy; type 1 cytokines are implicated; and interferon γ (IFN- γ) induces Fas expression on CD34 cells, leading to apoptotic cell death. The early immune system events in aplastic anemia are not well understood, but an oligoclonal, T cell response implies antigenic stimulus. The rarity of aplastic anemia despite common exposures (medicines, seronegative hepatitis) suggests that genetically determined features of the immune response can convert a normal physiologic response into a sustained abnormal autoimmune process, including polymorphisms in histocompatibility antigens, cytokine genes, and genes that regulate T cell polarization and effector function.

CLINICAL FEATURES

History Aplastic anemia can appear abruptly or insidiously. Bleeding is the most common early symptom; a complaint of days to weeks of easy bruising, oozing from the gums, nose bleeds, heavy menstrual flow, and sometimes petechiae will have been noticed. With thrombocytopenia, massive hemorrhage is unusual, but small amounts of bleeding in the central nervous system can result in catastrophic intracranial or retinal hemorrhage. Symptoms of anemia are also frequent, including lassitude, weakness, shortness of breath, and a pounding sensation in the ears. Infection is an unusual first symptom in aplastic anemia (unlike in agranulocytosis, where pharyngitis, anorectal infection, or frank sepsis occurs early). A striking feature of aplastic anemia is the restriction of symptoms to the hematologic system, and patients often feel and look remarkably well despite drastically reduced blood counts. Systemic complaints and weight loss should point to other etiologies of pancytopenia. Prior drug use, chemical exposure, and preceding viral illnesses must often be elicited with repeated questioning. A family history of hematologic diseases or blood abnormalities, of pulmonary or liver fibrosis, or of early hair graying points to a telomeropathy.

666 Physical Examination Petechiae and ecchymoses are typical, and retinal hemorrhages may be present. Pelvic and rectal examinations can often be deferred but, when performed, should be undertaken with great gentleness to avoid trauma; these will often show bleeding from the cervical os and blood in the stool. Pallor of the skin and mucous membranes is common except in the most acute cases or those already transfused. Infection on presentation is unusual but may occur if the patient has been symptomatic for a few weeks. Lymphadenopathy and splenomegaly are highly atypical of aplastic anemia. Café au lait spots and short stature suggest Fanconi anemia; peculiar nails and leukoplakia suggest dyskeratosis congenita; early graying (and use of hair dyes to mask it!) suggests a telomerase defect.

LABORATORY STUDIES

Blood The smear shows large erythrocytes and a paucity of platelets and granulocytes. Mean corpuscular volume (MCV) is commonly increased. Reticulocytes are absent or few, and lymphocyte numbers may be normal or reduced. The presence of immature myeloid forms suggests leukemia or MDS; nucleated red blood cells (RBCs) suggest marrow fibrosis or tumor invasion; abnormal platelets suggest either peripheral destruction or MDS.

Bone Marrow The bone marrow is usually readily aspirated but dilute on smear, and the fatty biopsy specimen may be grossly pale on withdrawal; a “dry tap” instead suggests fibrosis or myelophthisis. In severe aplasia, the smear of the aspirated specimen shows only red cells, residual lymphocytes, and stromal cells; the biopsy (which should be >1 cm in length) is superior for determination of cellularity and shows mainly fat under the microscope, with hematopoietic cells occupying <25% of the marrow space; in the most serious cases, the biopsy is virtually all fat. The correlation between marrow cellularity and disease severity is imperfect, in part because marrow cellularity declines physiologically with aging. Additionally, some patients with moderate disease by blood counts will have empty iliac crest biopsies, whereas “hot spots” of hematopoiesis may be seen in severe cases. If an iliac crest specimen is inadequate, cells may also be obtained by aspiration from the sternum. Residual hematopoietic cells should have normal morphology, except for mildly megaloblastic erythropoiesis; megakaryocytes are invariably greatly reduced and usually absent. Granulomas may indicate an infectious etiology of the marrow failure.

Ancillary Studies Chromosome breakage studies of peripheral blood using diepoxybutane or mitomycin C should be performed on children and younger adults to exclude Fanconi anemia. Very short telomere length (available commercially) strongly suggests the presence of a telomerase or shelterin mutation, which can be pursued by family studies and nucleotide sequencing. Chromosome studies of bone marrow cells are often revealing in MDS and should be negative in typical aplastic anemia. Flow cytometry offers a sensitive diagnostic test for PNH. Serologic studies may show evidence of viral infection, such as Epstein-Barr virus and HIV. Posthepatitis aplastic anemia is seronegative. The spleen size should be determined by computed tomography (CT) scanning or ultrasound if the physical examination of the abdomen is unsatisfactory. Occasionally MRI may be helpful to assess the fat content of vertebrae in order to distinguish aplasia from MDS.

DIAGNOSIS

The diagnosis of aplastic anemia is usually straightforward, based on the combination of pancytopenia with a fatty bone marrow. Aplastic anemia is a disease of the young and should be a leading diagnosis in the pancytopenic adolescent or young adult. When pancytopenia is secondary, the primary diagnosis is usually obvious from either history or physical examination: the massive spleen of alcoholic cirrhosis, the history of metastatic cancer or SLE, or miliary tuberculosis on chest radiograph (Table 130-1).

Diagnostic problems can occur with atypical presentations and among related hematologic diseases. Although pancytopenia is most common, some patients with bone marrow hypocellularity have depression of only one or two of three blood lines, with later progression to pancytopenia. The bone marrow in constitutional aplastic

anemia is morphologically indistinguishable from the aspirate in acquired disease. The diagnosis can be suggested by family history, abnormal blood counts since childhood, or the presence of associated physical anomalies. Aplastic anemia may be difficult to distinguish from the hypocellular variety of MDS: MDS is favored by finding morphologic abnormalities, particularly of megakaryocytes and myeloid precursor cells, and typical cytogenetic abnormalities (see below).

PROGNOSIS

The natural history of severe aplastic anemia is rapid deterioration and death. Historically, provision first of RBC and later of platelet transfusions and effective antibiotics were of some benefit, but few patients show spontaneous recovery. The major prognostic determinant is the blood count. Severe disease has been defined by the presence of two of three parameters: absolute neutrophil count <500/ μ L, platelet count <20,000/ μ L, and corrected reticulocyte count <1% (or absolute reticulocyte count <60,000/ μ L). In the era of effective immunosuppressive therapies, absolute numbers of reticulocytes (>25,000/ μ L) and lymphocytes (>1000/ μ L) may be better predictors of response to treatment and long-term outcome.

TREATMENT APLASTIC ANEMIA

Severe acquired aplastic anemia can be cured by replacement of the absent hematopoietic cells (and the immune system) by stem cell transplant, or it can be ameliorated by suppression of the immune system to allow recovery of the patient's residual bone marrow function. Glucocorticoids are not of value as primary therapy. Suspect exposures to drugs or chemicals should be discontinued; however, spontaneous recovery of severe blood count depression is rare, and a waiting period before beginning treatment may not be advisable unless the blood counts are only modestly depressed.

HEMATOPOIETIC STEM CELL TRANSPLANTATION

This is the best therapy for the younger patient with a fully histocompatible sibling donor (Chap. 139e). Human leukocyte antigen (HLA) typing should be ordered as soon as the diagnosis of aplastic anemia is established in a child or younger adult. In transplant candidates, transfusion of blood from family members should be avoided so as to prevent sensitization to histocompatibility antigens, but limited numbers of blood products probably do not greatly affect outcome. For allogeneic transplant from fully matched siblings, long-term survival rates for children are approximately 90%. Transplant morbidity and mortality are increased among adults, due to the higher risk of chronic GVHD and serious infections.

Most patients do not have a suitable sibling donor. Occasionally, a full phenotypic match can be found within the family and serve as well. Far more available are other alternative donors, either unrelated but histocompatible volunteers or closely but not perfectly matched family members. High-resolution matching at HLA and more effective conditioning regimens and GVHD prophylaxis have led to improved survival rates in patients who proceed to alternative donor transplant, in some series approximating results with conventional sibling donors. Patients will be at risk for late complications, especially a higher rate of cancer, if radiation is used as a component of conditioning.

IMMUNOSUPPRESSION

The standard regimen of antithymocyte globulin (ATG) in combination with cyclosporine induces hematologic recovery (independence from transfusion and a leukocyte count adequate to prevent infection) in 60–70% of patients. Children do especially well, whereas older adult patients often suffer complications due to the presence of comorbidities. An early robust hematologic response correlates with long-term survival. Improvement in granulocyte number is generally apparent within 2 months of treatment. Most recovered patients continue to have some degree of blood count depression, the MCV remains elevated, and bone marrow cellularity returns toward normal very slowly if at all. Relapse (recurrent

pancytopenia) is frequent, often occurring as cyclosporine is discontinued; most, but not all, patients respond to reinstitution of immunosuppression, but some responders become dependent on continued cyclosporine administration. Development of MDS, with typical marrow morphologic or cytogenetic abnormalities, occurs in approximately 15% of treated patients, usually but not invariably associated with a return of pancytopenia, and some patients develop leukemia. A laboratory diagnosis of PNH can generally be made at the time of presentation of aplastic anemia by flow cytometry; recovered patients may have frank hemolysis if the PNH clone expands. Bone marrow examinations should be performed if there is an unfavorable change in blood counts.

Horse ATG is administered as intravenous infusions over 4 days. ATG binds to peripheral blood cells; therefore, platelet and granulocyte numbers may decrease further during active treatment. Serum sickness, a flulike illness with a characteristic cutaneous eruption and arthralgia, often develops approximately 10 days after initiating treatment. Methylprednisolone is administered with ATG to ameliorate the immune consequences of heterologous protein infusion. Excessive or extended glucocorticoid therapy is associated with avascular joint necrosis. Cyclosporine is administered orally at an initial high dose, with subsequent adjustment according to blood levels obtained every 2 weeks; rough levels should be between 150 and 200 ng/mL. The most important side effects are nephrotoxicity, hypertension, seizures, and opportunistic infections, especially *Pneumocystis jiroveci* (prophylactic treatment with monthly inhaled pentamidine is recommended).

Most patients with aplastic anemia lack a suitable marrow donor, and immunosuppression is the treatment of choice. Overall survival is equivalent with transplantation and immunosuppression. However, successful transplant cures marrow failure, whereas patients who recover adequate blood counts after immunosuppression remain at risk of relapse and malignant evolution. Because of excellent results in children and younger adults, allogeneic transplant should be performed if a suitable sibling donor is available. Increasing age and the severity of neutropenia are the most important factors weighing in the decision between transplant and immunosuppression in adults who have a matched family donor: older patients do better with ATG and cyclosporine, whereas transplant is preferred if granulocytopenia is profound.

Outcomes following both transplant and immunosuppression have improved with time. High doses of cyclophosphamide, without stem cell rescue, have been reported to produce durable hematologic recovery, without relapse or evolution to MDS, but this treatment can produce sustained severe fatal neutropenia, and response is often delayed.

OTHER THERAPIES

The effectiveness of androgens has not been verified in controlled trials, but occasional patients will respond or even demonstrate blood count dependence on continued therapy. Sex hormones upregulate telomerase gene activity *in vitro*, which is possibly also their mechanism of action in improving marrow function. For patients with moderate disease, especially if a telomere defect is present, or those with severe pancytopenia in whom immunosuppression has failed, a 3- to 4-month trial is appropriate.

Hematopoietic growth factors (HGFs) such as erythropoietin and granulocyte colony-stimulating factor (G-CSF) are not definitive therapy for severe aplastic anemia, and even their roles as adjuncts to immunosuppression are not clear. In research protocols, thrombopoietin mimetics have shown surprising activity in patients with refractory aplastic anemia, with patterns of blood count recovery suggesting that they act as stem cell stimulants.

SUPPORTIVE CARE

Meticulous medical attention is required so that the patient may survive to benefit from definitive therapy or, having failed treatment, to maintain a reasonable existence in the face of pancytopenia. First and most important, infection in the presence of severe

neutropenia must be aggressively treated by prompt institution of parenteral, broad-spectrum antibiotics, usually ceftazidime or a combination of an aminoglycoside, cephalosporin, and semisynthetic penicillin. Therapy is empirical and must not await results of culture, although specific foci of infection such as oropharyngeal or anorectal abscesses, pneumonia, sinusitis, and typhlitis (necrotizing colitis) should be sought on physical examination and with radiographic studies. When indwelling plastic catheters become contaminated, vancomycin should be added. Persistent or recrudescing fever implies fungal disease: *Candida* and *Aspergillus* are common, especially after several courses of antibacterial antibiotics. A major reason for the improved prognosis in aplastic anemia has been the development of better antifungal drugs and the timely institution of such therapy when infection is suspected. Granulocyte transfusions using G-CSF–mobilized peripheral blood may be effective in the treatment of overwhelming or refractory infections. Hand washing, the single best method of preventing the spread of infection, remains a neglected practice. Nonabsorbed antibiotics for gut decontamination are poorly tolerated and not of proven value. Total reverse isolation does not reduce mortality from infections.

Both platelet and erythrocyte numbers can be maintained by transfusion. Alloimmunization historically limited the usefulness of platelet transfusions and is now minimized by several strategies, including use of single donors to reduce exposure and physical or chemical methods to diminish leukocytes in the product; HLA-matched platelets are often effective in patients refractory to random donor products. Inhibitors of fibrinolysis such as aminocaproic acid have not been shown to relieve mucosal oozing; the use of low-dose glucocorticoids to induce “vascular stability” is unproven and not recommended. Whether platelet transfusions are better used prophylactically or only as needed remains unclear. Any rational regimen of prophylaxis requires transfusions once or twice weekly to maintain the platelet count $>10,000/\mu\text{L}$ (oozing from the gut increases precipitously at counts $<5000/\mu\text{L}$). Menstruation should be suppressed either by oral estrogens or nasal follicle-stimulating hormone/luteinizing hormone (FSH/LH) antagonists. Aspirin and other nonsteroidal anti-inflammatory agents inhibit platelet function and must be avoided.

RBCs should be transfused to maintain a normal level of activity, usually at a hemoglobin value of 70 g/L (90 g/L if there is underlying cardiac or pulmonary disease); a regimen of 2 units every 2 weeks will replace normal losses in a patient without a functioning bone marrow. In chronic anemia, the iron chelators, deferoxamine and deferasirox, should be added at approximately the fiftieth transfusion to avoid secondary hemochromatosis.

PURE RED CELL APLASIA

Other, more restricted forms of marrow failure occur, in which only a single circulating cell type is affected and the marrow shows corresponding absence or decreased numbers of specific precursor cells: aregenerative anemia as in PRCA (see below), thrombocytopenia with amegakaryocytosis (Chap. 140), and neutropenia without marrow myeloid cells in agranulocytosis (Chap. 80). In general, and in contrast to aplastic anemia and MDS, the unaffected lineages appear quantitatively and qualitatively normal. Agranulocytosis, the most frequent of these syndromes, is usually a complication of medical drug use (with agents similar to those related to aplastic anemia), either by a mechanism of direct chemical toxicity or by immune destruction. Agranulocytosis has an incidence similar to aplastic anemia but is especially frequent among older adults and in women. The syndrome should resolve with discontinuation of exposure, but significant mortality is attached to neutropenia in the older and often previously unwell patient. Both pure white cell aplasia (agranulocytosis without incriminating drug exposure) and amegakaryocytic thrombocytopenia are exceedingly rare and, like PRCA, appear to be due to destructive antibodies or lymphocytes and can respond to immunosuppressive therapies. In all of the single-lineage failure syndromes, progression to pancytopenia or leukemia is unusual.

Self-limited
Transient erythroblastopenia of childhood
Transient aplastic crisis of hemolysis (acute B19 parvovirus infection)
Fetal red blood cell aplasia
Nonimmune hydrops fetalis (in utero B19 parvovirus infection)
Hereditary pure red cell aplasia
Congenital pure red cell aplasia (Diamond-Blackfan anemia)
Acquired pure red cell aplasia
Cancer
Thymoma
Lymphoid malignancies (and more rarely other hematologic diseases)
Paraneoplastic to solid tumors
Connective tissue disorders with immunologic abnormalities
Systemic lupus erythematosus, juvenile rheumatoid arthritis, rheumatoid arthritis
Multiple endocrine gland insufficiency
Viruses
Persistent B19 parvovirus, hepatitis, adult T cell leukemia virus, Epstein-Barr virus
Pregnancy
Drugs
Especially phenytoin, azathioprine, chloramphenicol, procainamide, isoniazid
Antibodies to erythropoietin
Idiopathic

DEFINITION AND DIFFERENTIAL DIAGNOSIS

PRCA is characterized by anemia, reticulocytopenia, and absent or rare erythroid precursor cells in the bone marrow. The classification of PRCA is shown in [Table 130-4](#). In adults, PRCA is acquired. An identical syndrome can occur constitutionally: Diamond-Blackfan anemia, or congenital PRCA, is diagnosed at birth or in early childhood and often responds to glucocorticoid treatment; mutations in ribosome protein genes are etiologic. Temporary red cell failure occurs in transient aplastic crisis of hemolytic anemias due to acute parvovirus infection ([Chap. 221](#)) and in transient erythroblastopenia of childhood, which occurs in normal children.

CLINICAL ASSOCIATIONS AND ETIOLOGY

PRCA has important associations with immune system diseases. A small minority of cases occur with a thymoma. More frequently, red cell aplasia can be the major manifestation of large granular lymphocytosis or complicate chronic lymphocytic leukemia. Some patients may be hypogammaglobulinemic. Infrequently (compared to agranulocytosis), PRCA can be due to an idiosyncratic drug reaction. Subcutaneous administration of erythropoietin (EPO) has provoked PRCA mediated by neutralizing antibodies.

Like aplastic anemia, PRCA results from diverse mechanisms. Antibodies to RBC precursors are frequently present in the blood, but T cell inhibition

is probably the more common immune mechanism. Cytotoxic lymphocyte activity restricted by histocompatibility locus or specific for human T cell leukemia/lymphoma virus 1–infected cells and natural killer cell activity inhibitory of erythropoiesis have been demonstrated in particularly well-studied individual cases.

PERSISTENT PARVOVIRUS B19 INFECTION

Chronic parvovirus infection is an important, treatable cause of PRCA. This common virus causes a benign exanthem of childhood (fifth disease) and a polyarthralgia/arthritis syndrome in adults. In patients with underlying hemolysis (or any condition that increases demand for RBC production), parvovirus infection can cause a transient aplastic crisis and an abrupt but temporary worsening of the anemia due to failed erythropoiesis. In normal individuals, acute infection is resolved by production of neutralizing antibodies to the virus, but in the setting of congenital, acquired, or iatrogenic immunodeficiency, persistent viral infection may occur. The bone marrow shows red cell aplasia and the presence of giant pronormoblasts ([Fig. 130-2](#)), which is the cytopathic sign of B19 parvovirus infection. Viral tropism for human erythroid progenitor cells is due to its use of erythrocyte P antigen as a cellular receptor for entry. Direct cytotoxicity of virus causes anemia if demands on erythrocyte production are high; in normal individuals, the temporary cessation of red cell production is not clinically apparent, and skin and joint symptoms are mediated by immune complex deposition.

TREATMENT PURE RED CELL APLASIA

History, physical examination, and routine laboratory studies may disclose an underlying disease or a drug exposure. Thymoma should be sought by radiographic procedures. Tumor excision

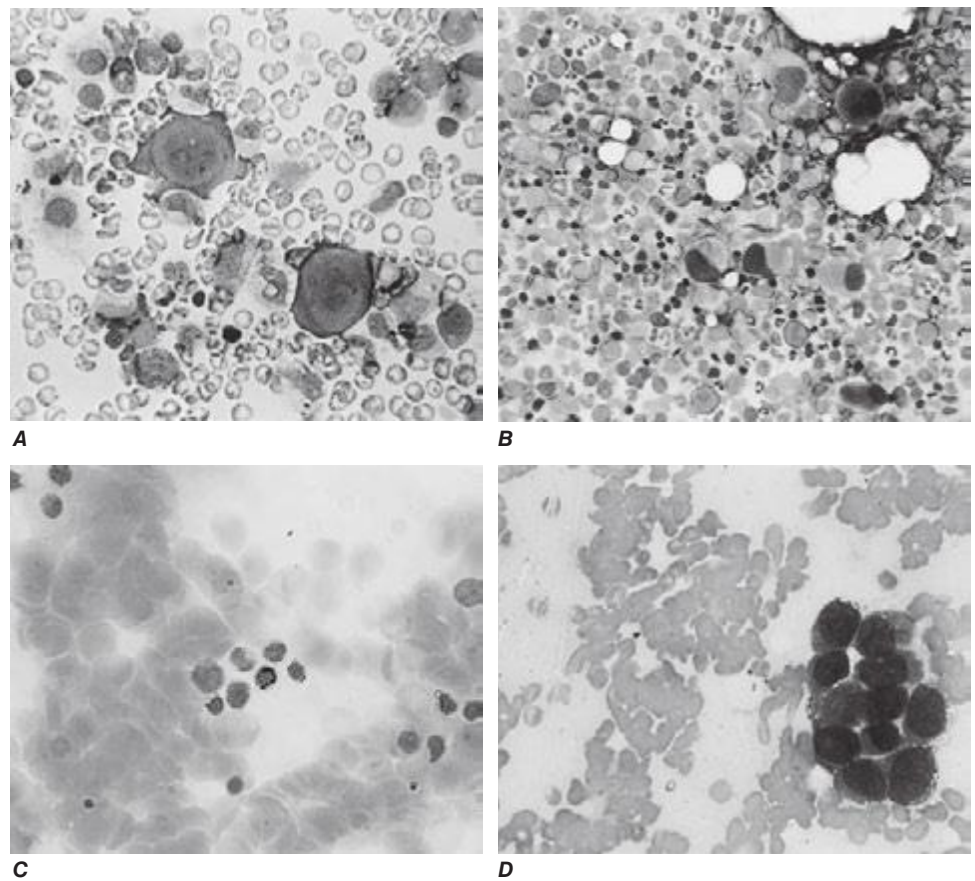


FIGURE 130-2 Pathognomonic cells in marrow failure syndromes. **A.** Giant pronormoblast, the cytopathic effect of B19 parvovirus infection of the erythroid progenitor cell. **B.** Uninuclear megakaryocyte and microblastic erythroid precursors typical of the 5q–myelodysplasia syndrome. **C.** Ringed sideroblast showing perinuclear iron granules. **D.** Tumor cells present on a touch preparation made from the marrow biopsy of a patient with metastatic carcinoma.

is indicated, but anemia does not necessarily improve with surgery. The diagnosis of parvovirus infection requires detection of viral DNA sequences in the blood (IgG and IgM antibodies are commonly absent). The presence of erythroid colonies has been considered predictive of response to immunosuppressive therapy in idiopathic PRCA.

Red cell aplasia is compatible with long-term survival with supportive care alone: a combination of erythrocyte transfusions and iron chelation. For persistent B19 parvovirus infection, almost all patients respond to intravenous immunoglobulin therapy (e.g., 0.4 g/kg daily for 5 days), although relapse and retreatment may be expected, especially in patients with AIDS. The majority of patients with idiopathic PRCA respond favorably to immunosuppression. Most first receive a course of glucocorticoids. Also effective are cyclosporine, ATG, azathioprine, and cyclophosphamide.

MYELOYDYSPLASIA

DEFINITION

The myelodysplastic syndromes (MDS) are a heterogeneous group of hematologic disorders broadly characterized by both (1) cytopenias due to bone marrow failure and (2) a high risk of development of acute

myeloid leukemia (AML). Anemia, often with thrombocytopenia and neutropenia, occurs with dysmorphic (abnormal appearing) and usually cellular bone marrow, which is evidence of ineffective blood cell production. In patients with “low-risk” MDS, marrow failure dominates the clinical course. In other patients, myeloblasts are present at diagnosis, chromosomes are abnormal, and the “high risk” is due to leukemic progression. MDS may be fatal due to the complications of pancytopenia or the incurability of leukemia, but a large proportion of patients will die of concurrent disease, the comorbidities typical in an elderly population. A clinically useful nosology of these often confusing entities was first developed by the French-American-British Cooperative Group in 1983. Five entities were defined: refractory anemia (RA), refractory anemia with ringed sideroblasts (RARS), refractory anemia with excess blasts (RAEB), refractory anemia with excess blasts in transformation (RAEB-t), and chronic myelomonocytic leukemia (CMML). The World Health Organization (WHO) classification (2002) recognized that the distinction between RAEB-t and AML is arbitrary and grouped them together as acute leukemia and that CMML behaves as a myeloproliferative disease; the WHO classification also separated refractory anemias with dysmorphic change restricted to erythroid lineage from those with multilineage changes. In a 2008 revision, specific categories for unilineage dysplasias were added (Table 130-5).

TABLE 130-5 WORLD HEALTH ORGANIZATION (WHO) CLASSIFICATION OF MYELOYDYSPLASTIC SYNDROMES/NEOPLASMS

Name	WHO Estimated Proportion of Patients with MDS	Peripheral Blood: Key Features	Bone Marrow: Key Features
<i>Refractory cytopenias with unilineage dysplasia (RCUD):</i>			
Refractory anemia (RA)	10–20%	Anemia <1% of blasts	Unilineage erythroid dysplasia (in ≥10% of cells) <5% blasts
Refractory neutropenia (RN)	<1%	Neutropenia <1% blasts	Unilineage granulocytic dysplasia <5% blasts
Refractory thrombocytopenia (RT)	<1%	Thrombocytopenia <1% blasts	Unilineage megakaryocytic dysplasia <5% blasts
Refractory anemia with ringed sideroblasts (RARS)	3–11%	Anemia No blasts	Unilineage erythroid dysplasia ≥15% of erythroid precursors are ringed sideroblasts <5% blasts
Refractory cytopenias with multilineage dysplasia (RCMD)	30%	Cytopenia(s) <1% blasts No Auer rods	Multilineage dysplasia ± ringed sideroblasts <5% blasts No Auer rods
Refractory anemia with excess blasts, type 1 (RAEB-1)	40%	Cytopenia(s) <5% blasts No Auer rods	Unilineage or multilineage dysplasia
Refractory anemia with excess blasts, type 2 (RAEB-2)		Cytopenia(s) 5–19% blasts ± Auer rods	Unilineage or multilineage dysplasia 10–19% blasts ± Auer rods
MDS associated with isolated del(5q) [del(5q)]	Uncommon	Anemia Normal or high platelet count <1% blasts	Isolated 5q31 chromosome deletion Anemia; hypolobated megakaryocytes <5% blasts
Childhood MDS, including refractory cytopenia of childhood (<i>provisional</i>) (RCC)	<1%	Pancytopenia	<5% marrow blasts for RCC Marrow usually hypocellular
MDS, unclassifiable (MDS-U)	?	Cytopenia ≤1% blasts	Does not fit other categories Dysplasia <5% blasts If no dysplasia, MDS-associated karyotype

Note: If peripheral blood blasts are 2–4%, the diagnosis is RAEB-1 even if marrow blasts are <5%. If Auer rods are present, the WHO considers the diagnosis RAEB-2 if the blast proportion is <20% (even if <10%), or acute myeloid leukemia (AML) if at least 20% blasts. For all subtypes, peripheral blood monocytes are $<1 \times 10^9/L$. Bicytopenia may be observed in RCUD subtypes, but pancytopenia with unilineage marrow dysplasia should be classified as MDS-U. Therapy-related MDS (t-MDS), whether due to alkylating agents or topoisomerase II inhibitors (t-MDS/t-AML) is now included in the WHO classification of myeloid neoplasms. The listing in this table excludes MDS/myeloproliferative neoplasm overlap categories, such as chronic myelomonocytic leukemia, juvenile myelomonocytic leukemia, and the provisional entity RARS with thrombocytosis.

Abbreviation: MDS, myelodysplastic syndrome.

The diagnosis of MDS may be a challenge, because sometimes subtle clinical and pathologic features must be distinguished and precise diagnostic categorization requires a hematopathologist knowledgeable in the latest classification scheme. Nonetheless, it is important that the internist and primary care physician be sufficiently familiar with MDS to expedite referral to a hematologist, both because many new therapies are now available to improve hematopoietic function and the judicious use of supportive care can improve the patient's quality of life.

EPIDEMIOLOGY

Idiopathic MDS is a disease of the elderly; the mean age at onset is older than 70 years. There is a slight male preponderance. MDS is a relatively common form of bone marrow failure, with reported incidence rates of 35 to >100 per million persons in the general population and 120 to >500 per million in the older adult. MDS is rare in children, but monocytic leukemia can be seen. Secondary or therapy-related MDS is not age related. Rates of MDS have increased over time, due to better recognition of the syndrome by physicians and an aging population.

ETIOLOGY AND PATHOPHYSIOLOGY

MDS is associated with environmental exposures such as radiation and benzene; other risk factors have been reported inconsistently. Secondary MDS occurs as a late toxicity of cancer treatment, usually a combination of radiation and the radiomimetic alkylating agents such as busulfan, nitrosourea, or procarbazine (with a latent period of 5–7 years) or the DNA topoisomerase inhibitors (2-year latency). Acquired aplastic anemia, Fanconi anemia, and other constitutional marrow failure diseases can evolve into MDS. However, the typical MDS patient does not have a suggestive environmental exposure history or a preceding hematologic disease. MDS is a disease of aging, suggesting random cumulative intrinsic and environmental damage to marrow cells.

MDS is a clonal hematopoietic stem cell disorder characterized by disordered cell proliferation and impaired differentiation, resulting in cytopenias and risk of progression to leukemia. Both chromosomal and genetic instability have been implicated, and both are likely aging-related. Cytogenetic abnormalities are found in approximately one-half of patients, and some of the same specific lesions are also seen in frank leukemia; aneuploidy (chromosome loss or gain) is more frequent than translocations. More sensitive assays, such as comparative genomic hybridization and single nucleotide polymorphism arrays, reveal chromosomal abnormalities in a large proportion of patients with normal conventional cytogenetics. Accelerated telomere attrition may destabilize the genome in marrow failure and predispose to acquisition of chromosomal lesions. Cytogenetic abnormalities are not random (loss of all or part of 5, 7, and 20, trisomy of 8) and may be related to etiology (11q23 following topoisomerase II inhibitors). The type and number of cytogenetic abnormalities strongly correlate with the probability of leukemic transformation and survival.

Genomics has illuminated the role of point mutations in the pathophysiology of MDS. Recurrent somatic mutations, acquired in the abnormal marrow cells and absent in the germline, have been identified in almost 100 genes. Many of the same genes are also mutated in AML without MDS, whereas others are distinctive in subtypes of MDS. A prominent example of the latter is the discovery of mutations in genes of the RNA splicing machinery, especially *SF3B1*, which strongly associate with sideroblastic anemia. Some mutations correlate with prognosis: spliceosome defects with favorable outcome, and mutations in *EZH2*, *TP53*, *RUNX1*, and *ASXL1* with poor outcome. Mutations and cytogenetic abnormalities are not independent: *TP53* mutations associate with complex cytogenetic abnormalities and *TET2* mutations with normal cytogenetics. Correlation and exclusion in the pattern of mutations indicate a functional genomic architecture. Analysis of deep sequencing results in patients whose MDS evolved to AML has shown evidence of clonal succession, with founder clones acquiring further mutations that allow clonal dominance. Furthermore, the prevalence of abnormal cells by morphology underestimates bone marrow involvement by MDS clones, as cells normal in appearance are apparently derived from the abnormal clones. Both presenting and

evolving hematologic manifestations result from the accumulation of multiple genetic lesions: loss of tumor-suppressor genes, activating oncogene mutations, epigenetic pathways that affect mRNA processing and methylation status, or other harmful alterations. Pathophysiology has been linked to mutations and chromosome abnormalities in some specific MDS syndromes. The 5q- deletion leads to heterozygous loss of a ribosomal protein gene that is also mutant in Diamond-Blackfan anemia, and both are characterized by deficient erythropoiesis. An immune pathophysiology may underlie trisomy 8 MDS, in which patients often experience improved blood counts after immunosuppressive therapy; there is T cell activity directed to hematopoietic progenitors, which the cytogenetically aberrant clone resists. However, in general for MDS, the role of the immune system and its cells and cytokines; the role of the hematopoietic stem cell niche, the microenvironment, and cell-cell interactions; the fate of normal cells in the Darwinian competitive environment of the dysplastic marrow; and how mutant cells produce marrow failure in MDS are not well understood.

CLINICAL FEATURES

Anemia dominates the early course. Most symptomatic patients complain of the gradual onset of fatigue and weakness, dyspnea, and pallor, but at least one-half the patients are asymptomatic, and their MDS is discovered only incidentally on routine blood counts. Previous chemotherapy or radiation exposure is an important historic fact. Fever and weight loss should point to a myeloproliferative rather than myelodysplastic process. MDS in childhood is rare and, when diagnosed, increases the likelihood of an underlying genetic disease. Children with Down syndrome are susceptible to MDS, and a family history may indicate a hereditary form of sideroblastic anemia, Fanconi anemia, or a telomeropathy. Inherited *GATA2* mutations, as in the MonoMAC syndrome (with increased susceptibility to viral, mycobacteria, and fungal infections, as well as deficient numbers of monocytes, natural killer cells, and B lymphocytes), also cause MDS in young patients.

The physical examination is remarkable for signs of anemia; approximately 20% of patients have splenomegaly. Some unusual skin lesions, including Sweet syndrome (febrile neutrophilic dermatosis), occur with MDS. Accompanying autoimmune syndromes are not infrequent. In the younger patient, stereotypical anomalies point to a constitutional syndrome (short stature, abnormal thumbs in Fanconi anemia; early graying in the telomeropathies; cutaneous warts in *GATA2* deficiency).

LABORATORY STUDIES

Blood Anemia is present in most cases, either alone or as part of bi- or pancytopenia; isolated neutropenia or thrombocytopenia is more unusual. Macrocytosis is common, and the smear may be dimorphic with a distinctive population of large red blood cells. Platelets are also large and lack granules. In functional studies, they may show marked abnormalities, and patients may have bleeding symptoms despite seemingly adequate numbers. Neutrophils are hypogranulated; have hyposegmented, ringed, or abnormally segmented nuclei; contain Döhle bodies; and may be functionally deficient. Circulating myeloblasts usually correlate with marrow blast numbers, and their quantity is important for classification and prognosis. The total white blood cell count (WBC) is usually normal or low, except in chronic myelomonocytic leukemia. As in aplastic anemia, MDS can be associated with a clonal population of PNH cells. Genetic testing is commercially available for constitutional syndromes.

Bone Marrow The bone marrow is usually normal or hypercellular, but in about 20% of cases, it is sufficiently hypocellular to be confused with aplasia. No single characteristic feature of marrow morphology distinguishes MDS, but the following are commonly observed: dyserythropoietic changes (especially nuclear abnormalities) and ringed sideroblasts in the erythroid lineage; hypogranulation and hyposegmentation in granulocytic precursors, with an increase in myeloblasts; and megakaryocytes showing reduced numbers of or disorganized

TABLE 130-6 INTERNATIONAL PROGNOSTIC SCORING SYSTEM (IPSS)

Prognostic Variable	Score Value			
	0	0.5	1	2
Bone marrow blasts (%)	<5%	5–10%	11–20%	21–30%
Karyotype ^a	Good	Intermediate	Poor	
Cytopenia ^b (lineages affected)	0 or 1	2 or 3		
Risk Group Scores	Score			
Low	0			
Intermediate-1	0.5–1			
Intermediate-2	1.5–2			
High	≥2.5			

^aGood, normal, -Y, del(5q), del(20q); poor, complex (≥3 abnormalities) or chromosome 7 abnormalities; intermediate, all other abnormalities. ^bCytopenias defined as hemoglobin <100 g/L, platelet count <100,000/μL, and absolute neutrophil count <1500/μL.

nuclei. Megaloblastic nuclei associated with defective hemoglobinization in the erythroid lineage are common. Prognosis strongly correlates with the proportion of marrow blasts. Cytogenetic analysis and fluorescent in situ hybridization can identify chromosomal abnormalities.

DIFFERENTIAL DIAGNOSIS

Deficiencies of vitamin B₁₂ or folate should be excluded by appropriate blood tests; vitamin B₆ deficiency can be assessed by a therapeutic trial of pyridoxine if the bone marrow shows ringed sideroblasts. Marrow dysplasia can be observed in acute viral infections, drug reactions, or chemical toxicity but should be transient. More difficult are the distinctions between hypocellular MDS and aplasia or between refractory anemia with excess blasts and early acute leukemia. The WHO considers the presence of 20% blasts in the marrow as the criterion that separates AML from MDS. In young patients, underlying, predisposing genetic diseases should be considered (see above).

PROGNOSIS

The median survival varies greatly from years for patients with 5q- or sideroblastic anemia to a few months in refractory anemia with excess blasts or severe pancytopenia associated with monosomy 7; an International Prognostic Scoring System (IPSS; [Table 130-6](#)) assists in making predictions. Even “low-risk” MDS has significant morbidity and mortality. Most patients die as a result of complications of pancytopenia and not due to leukemic transformation; perhaps one-third will succumb to other diseases unrelated to their MDS. Precipitous worsening of pancytopenia, acquisition of new chromosomal abnormalities on serial cytogenetic determination, increase in the number of blasts, and marrow fibrosis are all poor prognostic indicators. The outlook in therapy-related MDS, regardless of type, is extremely poor, and most patients will progress within a few months to refractory AML.

TREATMENT MYELOYDYSPLASIA

Historically, the therapy of MDS has been unsatisfactory, but new drugs recently have been approved for this disease. Several regimens appear to not only improve blood counts but to delay onset of leukemia and to improve survival. The choice of therapy for an individual patient, administration of treatment, and management of toxicities are complicated and require hematologic expertise.

Only hematopoietic stem cell transplantation offers cure of MDS. The current survival rate in selected patient cohorts is ~50% at 3 years and is improving. Results using unrelated matched donors are now similar to those obtained using siblings, and patients in their 50s and 60s have been successfully transplanted. Nevertheless, treatment-related mortality and morbidity increase with recipient age. Complicating the decision to undertake transplant is that the high-risk patient, for whom the procedure is most obviously indicated, has a high probability of a poor outcome from transplant-related mortality or disease relapse, whereas the low-risk patient,

who is more likely to tolerate transplant, also may do well for years with less aggressive therapies.

MDS has been regarded as particularly refractory to cytotoxic chemotherapy regimens, and as in AML in the older adult, drug toxicity is frequent and often fatal, and remissions if achieved are brief. Low doses of cytotoxic drugs have been administered for their “differentiation” potential, and from this experience, drug therapies have emerged based on pyrimidine analogues. These new drugs are classified as epigenetic modulators, believed to act through a demethylating mechanism to alter gene regulation and allow differentiation to mature blood cells from the abnormal MDS stem cell (although global methylation status has not correlated with clinical efficacy). Azacitidine and decitabine are two epigenetic modifiers frequently used in bone marrow failure clinics. Azacitidine improves blood counts and survival in MDS, compared to best supportive care. Azacitidine is usually administered subcutaneously, daily for 7 days, at 4-week intervals, for at least four cycles before assessing for response. Overall, generally improved blood counts with a decrease in transfusion requirements occurred in ~50% of patients in published trials. Response is dependent on continued drug administration, and most patients eventually will no longer respond and experience recurrent cytopenias or progression to AML. Decitabine is closely related to azacitidine and more potent; 30–50% of patients show responses in blood counts, with a duration of response of almost a year. Decitabine is usually administered by continuous intravenous infusion in regimens of varying doses and durations of 3 to 10 days in repeating cycles. The major toxicity of azacitidine and decitabine is myelosuppression, leading to worsened blood counts. Other symptoms associated with cancer chemotherapy frequently occur. Demethylating agents are frequently used in the high-risk patient who is not a candidate for stem cell transplant. In the lower risk patient, they are also effective, but alternative therapies should be considered.

Lenalidomide, a thalidomide derivative with a more favorable toxicity profile, is particularly effective in reversing anemia in MDS patients with 5q- syndrome; not only do a high proportion of these patients become transfusion independent with normal or near-normal hemoglobin levels, but their cytogenetics also become normal. The drug has many biologic activities, and it is unclear which is critical for clinical efficacy. Lenalidomide is administered orally. Most patients will improve within 3 months of initiating therapy. Toxicities include myelosuppression (worsening thrombocytopenia and neutropenia, necessitating blood count monitoring) and an increased risk of deep vein thrombosis and pulmonary embolism.

Immunosuppression, as used in aplastic anemia, also may produce sustained independence from transfusion and improve survival. ATG, cyclosporine, and the anti-CD52 monoclonal antibody alemtuzumab are especially effective in younger MDS patients (<60 years old) with more favorable IPSS scores and who bear the histocompatibility antigen HLA-DR15.

HGFs can improve blood counts but, as in most other marrow failure states, have been most beneficial to patients with the least severe pancytopenia. EPO alone or in combination with G-CSF can improve hemoglobin levels, but mainly in those with low serum EPO levels who have no or only a modest need for transfusions. Survival does not appear to be improved by G-CSF treatment alone but may be enhanced by erythropoietin and amelioration of anemia. G-CSF treatment alone failed to improve survival in a controlled trial.

The same principles of supportive care described for aplastic anemia apply to MDS. Despite improvements in drug therapy, many patients will be anemic for years. RBC transfusion support should be accompanied by iron chelation to prevent secondary hemochromatosis.

MYELOPHTHISIC ANEMIAS

Fibrosis of the bone marrow (see [Fig. 129-2](#)), usually accompanied by a characteristic blood smear picture called *leukoerythroblastosis*, can occur as a primary hematologic disease, called *myelofibrosis* or *myeloid metaplasia* ([Chap. 131](#)), and as a secondary process, called

672 *myelophthisis*. Myelophthisis, or secondary myelofibrosis, is reactive. Fibrosis can be a response to invading tumor cells, usually an epithelial cancer of breast, lung, or prostate origin or neuroblastoma. Marrow fibrosis may occur with infection of mycobacteria (both *Mycobacterium tuberculosis* and *Mycobacterium avium*), fungi, or HIV and in sarcoidosis. Intracellular lipid deposition in Gaucher's disease and obliteration of the marrow space related to absence of osteoclast remodeling in congenital osteopetrosis also can produce fibrosis. Secondary myelofibrosis is a late consequence of radiation therapy or treatment with radiomimetic drugs. Usually the infectious or malignant underlying processes are obvious. Marrow fibrosis can also be a feature of a variety of hematologic syndromes, especially chronic myeloid leukemia, multiple myeloma, lymphomas, myeloma, and hairy cell leukemia.

The pathophysiology has three distinct features: proliferation of fibroblasts in the marrow space (myelofibrosis); the extension of hematopoiesis into the long bones and into extramedullary sites, usually the spleen, liver, and lymph nodes (myeloid metaplasia); and ineffective erythropoiesis. The etiology of the fibrosis is unknown but most likely involves dysregulated production of growth factors: platelet-derived growth factor and transforming growth factor β have been implicated. Abnormal regulation of other hematopoietins would lead to localization of blood-producing cells in nonhematopoietic tissues and uncoupling of the usually balanced processes of stem cell proliferation and differentiation. Myelofibrosis is remarkable for pancytopenia despite very large numbers of circulating hematopoietic progenitor cells.

Anemia is dominant in secondary myelofibrosis, usually normocytic and normochromic. The diagnosis is suggested by the characteristic leukoerythroblastic smear (see Fig. 129-1). Erythrocyte morphology is highly abnormal, with circulating nucleated RBCs, teardrops, and shape distortions. WBC numbers are often elevated, sometimes mimicking a leukemoid reaction, with circulating myelocytes, promyelocytes, and myeloblasts. Platelets may be abundant and are often of giant size. Inability to aspirate the bone marrow, the characteristic "dry tap," can allow a presumptive diagnosis in the appropriate setting before the biopsy is decalcified.

The course of secondary myelofibrosis is determined by its etiology, usually a metastatic tumor or an advanced hematologic malignancy. Treatable causes must be excluded, especially tuberculosis and fungus. Transfusion support can relieve symptoms.

131 Polycythemia Vera and Other Myeloproliferative Neoplasms

Jerry L. Spivak

The World Health Organization (WHO) classification of the chronic myeloproliferative neoplasms (MPNs) includes eight disorders, some of which are rare or poorly characterized (Table 131-1) but all of which share an origin in a multipotent hematopoietic progenitor cell; overproduction of one or more of the formed elements of the blood without significant dysplasia; and a predilection to extramedullary hematopoiesis, myelofibrosis, and transformation at varying rates to acute leukemia. Within this broad classification, however, significant phenotypic heterogeneity exists. Some diseases such as chronic myelogenous leukemia (CML), chronic neutrophilic leukemia (CNL), and chronic eosinophilic leukemia (CEL) express primarily a myeloid phenotype, whereas in other diseases, such as polycythemia vera (PV), primary myelofibrosis (PMF), and essential thrombocytosis (ET), erythroid or megakaryocytic hyperplasia predominates. The latter three disorders, in contrast to the former three, also appear capable of transforming into each other.

TABLE 131-1 WORLD HEALTH ORGANIZATION CLASSIFICATION OF CHRONIC MYELOPROLIFERATIVE NEOPLASMS

Chronic myeloid leukemia, bcr-abl–positive
Chronic neutrophilic leukemia
Chronic eosinophilic leukemia, not otherwise specified
Polycythemia vera
Primary myelofibrosis
Essential thrombocytosis
Mastocytosis
Myeloproliferative neoplasms, unclassifiable

Such phenotypic heterogeneity has a genetic basis; CML is the consequence of the balanced translocation between chromosomes 9 and 22 [t(9;22)(q34;11)]; CNL has been associated with a t(15;19) translocation; and CEL occurs with a deletion or balanced translocations involving the *PDGFR α* gene. By contrast, to a greater or lesser extent, PV, PMF, and ET are characterized by a mutation, V617F, that causes constitutive activation of JAK2, a tyrosine kinase essential for the function of the erythropoietin and thrombopoietin receptors but not the granulocyte colony-stimulating factor receptor. This important distinction is also reflected in the natural histories of CML, CNL, and CEL, which are usually measured in years, and their high rate of leukemic transformation. By contrast, the natural history of PV, PMF, and ET is usually measured in decades, and transformation to acute leukemia is uncommon in PV and ET in the absence of exposure to mutagenic drugs. This chapter, therefore, will focus only on PV, PMF, and ET, because their clinical and genetic overlap is substantial even though their clinical courses are distinctly different.

The other chronic myeloproliferative neoplasms will be discussed in Chaps. 133 and 135e.

POLYCYTHEMIA VERA

PV is a clonal disorder involving a multipotent hematopoietic progenitor cell in which phenotypically normal red cells, granulocytes, and platelets accumulate in the absence of a recognizable physiologic stimulus. The most common of the chronic MPNs, PV occurs in 2.5 per 100,000 persons, sparing no adult age group and increasing with age to rates over 10/100,000. Familial transmission is infrequent, and women predominate among sporadic cases.

ETIOLOGY

The etiology of PV is unknown. Although nonrandom chromosome abnormalities such as deletion 20q and trisomy 8 and 9 have been documented in up to 30% of untreated PV patients, unlike CML, no consistent cytogenetic abnormality has been associated with the disorder. However, a mutation in the autoinhibitory pseudokinase domain of the tyrosine kinase JAK2—that replaces valine with phenylalanine (V617F), causing constitutive kinase activation—appears to have a central role in the pathogenesis of PV.

JAK2 is a member of an evolutionarily well-conserved, nonreceptor tyrosine kinase family and serves as the cognate tyrosine kinase for the erythropoietin and thrombopoietin receptors. It also functions as an obligate chaperone for these receptors in the Golgi apparatus and is responsible for their cell-surface expression. The conformational change induced in the erythropoietin and thrombopoietin receptors following binding to their respective cognate ligands, erythropoietin or thrombopoietin, leads to JAK2 autophosphorylation, receptor phosphorylation, and phosphorylation of proteins involved in cell proliferation, differentiation, and resistance to apoptosis. Transgenic animals lacking *JAK2* die as embryos from severe anemia. Constitutive activation of JAK2, on the other hand, explains the erythropoietin hypersensitivity, erythropoietin-independent erythroid colony formation, rapid terminal differentiation, increase in Bcl-X_L expression, and apoptosis resistance in the absence of erythropoietin that characterize the in vitro behavior of PV erythroid progenitor cells.

Importantly, the *JAK2* gene is located on the short arm of chromosome 9, and loss of heterozygosity on chromosome 9p due to mitotic recombination is the most common cytogenetic abnormality in PV. The segment of 9p involved contains the *JAK2* locus, and loss of heterozygosity in this region leads to homozygosity for *JAK2* V617F. More than 95% of PV patients express this mutation, as do approximately 50% of PMF and ET patients. Homozygosity for the mutation occurs in approximately 30% of PV patients and 60% of PMF patients but is rare in ET. Over time, a portion of PV *JAK2* V617F heterozygotes become homozygotes due to mitotic recombination, but usually not after 10 years of the disease. Most PV patients who do not express *JAK2* V617F express a mutation in exon 12 of the kinase and are not clinically different from those who do, nor do *JAK2* V617F heterozygotes differ clinically from homozygotes. Interestingly, the predisposition to acquire mutations in *JAK2* appears to be associated with a specific *JAK2* gene haplotype, GGCC. *JAK2* V617F is the basis for many of the phenotypic and biochemical characteristics of PV such as elevation of the leukocyte alkaline phosphatase (LAP) score; however, it cannot solely account for the entire PV phenotype and is probably not the initiating lesion in the three MPNs. First, PV patients with the same phenotype and documented clonal disease lack any mutation of *JAK2*. Second, ET and PMF patients have the same mutation but different clinical phenotypes. Third, familial PV can occur without the mutation, even when other members of the same family express it. Fourth, not all the cells of the malignant clone express *JAK2* V617F. Fifth, *JAK2* V617F has been observed in patients with long-standing idiopathic erythrocytosis. Sixth, in some patients, *JAK2* V617F appears to be acquired after another mutation. Finally, in some *JAK2* V617F-positive PV or ET patients, acute leukemia can occur in a *JAK2* V617F-negative progenitor cell. However, although *JAK2* V617F alone may not be sufficient to cause PV, it appears essential for the transformation of ET to PV, although not for its transformation to PMF.

CLINICAL FEATURES

Although isolated thrombocytosis, leukocytosis, or splenomegaly may be the initial presenting manifestation of PV, most often the disorder is first recognized by the incidental discovery of a high hemoglobin or hematocrit. With the exception of aquagenic pruritus, no symptoms distinguish PV from other causes of erythrocytosis.

Uncontrolled erythrocytosis causes hyperviscosity, leading to neurologic symptoms such as vertigo, tinnitus, headache, visual disturbances, and transient ischemic attacks (TIAs). Systolic hypertension is also a feature of the red cell mass elevation. In some patients, venous or arterial thrombosis may be the presenting manifestation of PV. Any vessel can be affected; but cerebral, cardiac, or mesenteric vessels are most commonly involved. Intraabdominal venous thrombosis is particularly common in young women and may be catastrophic if a sudden and complete obstruction of the hepatic vein occurs. Indeed, PV should be suspected in any patient who develops hepatic vein thrombosis. Digital ischemia, easy bruising, epistaxis, acid-peptic disease, or gastrointestinal hemorrhage may occur due to vascular stasis or thrombocytosis. Erythema, burning, and pain in the extremities, a symptom complex known as erythromelalgia, are other complications of the thrombocytosis of PV due to increased platelet stickiness. Given the large turnover of hematopoietic cells, hyperuricemia with secondary gout, uric acid stones, and symptoms due to hypermetabolism can also complicate the disorder.

DIAGNOSIS

When PV presents with erythrocytosis in combination with leukocytosis, thrombocytosis, or splenomegaly or a combination of these, the diagnosis is apparent. However, when patients present with an elevated hemoglobin or hematocrit alone, the diagnostic evaluation is more complex because of the many diagnostic possibilities (Table 131-2). Furthermore, unless the hemoglobin level is ≥ 20 g/dL (hematocrit $\geq 60\%$), it is not possible to distinguish true erythrocytosis from disorders causing plasma volume contraction. This is because uniquely in PV, in contrast to other causes of true erythrocytosis, there is expansion of the plasma volume, which can mask the elevated red

TABLE 131-2 CAUSES OF ERYTHROCYTOSIS

Relative Erythrocytosis	
Hemoconcentration secondary to dehydration, diuretics, ethanol abuse, androgens, or tobacco abuse	
Absolute Erythrocytosis	
Hypoxia	Tumors
Carbon monoxide intoxication	Hypernephroma
High-oxygen-affinity hemoglobin	Hepatoma
High altitude	Cerebellar hemangioblastoma
Pulmonary disease	Uterine myoma
Right to left cardiac or vascular shunts	Adrenal tumors
Sleep apnea syndrome	Meningioma
Hepatopulmonary syndrome	Pheochromocytoma
Renal Disease	Drugs
Renal artery stenosis	Androgens
Focal sclerosing or membranous glomerulonephritis	Recombinant erythropoietin
Postrenal transplantation	Familial (with normal hemoglobin function)
Renal cysts	Erythropoietin receptor mutation
Bartter's syndrome	<i>VHL</i> mutations (Chuvash polycythemia)
	2,3-BPG mutation
	Polycythemia vera

Abbreviations: 2,3-BPG, 2,3-bisphosphoglycerate; *VHL*, von Hippel-Lindau.

cell mass; thus, red cell mass and plasma volume determinations are necessary to establish the presence of an absolute erythrocytosis and to distinguish this from relative erythrocytosis due to a reduction in plasma volume alone (also known as *stress* or *spurious erythrocytosis* or *Gaisböck's syndrome*). Figure 77-18 illustrates a diagnostic algorithm for the evaluation of suspected erythrocytosis. Assay for *JAK2* mutations in the presence of a normal arterial oxygen saturation provides an alternative diagnostic approach to erythrocytosis when red cell mass and plasma volume determinations are not available; a normal serum erythropoietin level does not exclude the presence of PV, but an elevated erythropoietin level is more consistent with a secondary cause for the erythrocytosis.

Other laboratory studies that may aid in diagnosis include the red cell count, mean corpuscular volume, and red cell distribution width (RDW), particularly when the hematocrit or hemoglobin levels are less than 60% or 20 g/dL, respectively. Only three situations cause microcytic erythrocytosis: β thalassemia trait, hypoxic erythrocytosis, and PV. With β thalassemia trait, the RDW is normal, whereas with hypoxic erythrocytosis and PV, the RDW may be elevated due to associated iron deficiency. Today, however, the assay for *JAK2* V617F has superseded other tests for establishing the diagnosis of PV. Of course, in patients with associated acid-peptic disease, occult gastrointestinal bleeding may lead to a presentation with hypochromic, microcytic anemia, masking the presence of PV.

A bone marrow aspirate and biopsy provide no specific diagnostic information because these may be normal or indistinguishable from ET or PMF. Similarly, no specific cytogenetic abnormality is associated with the disease, and the absence of a cytogenetic marker does not exclude the diagnosis.

COMPLICATIONS

Many of the clinical complications of PV relate directly to the increase in blood viscosity associated with red cell mass elevation and indirectly to the increased turnover of red cells, leukocytes, and platelets with the attendant increase in uric acid and cytokine production. The latter appears to be responsible for constitutional symptoms. Peptic ulcer disease can also be due to *Helicobacter pylori* infection, the incidence of which is increased in PV, while the pruritus associated with this disorder may be a consequence of mast cell activation by *JAK2* V617F. A sudden increase in spleen size can be associated with painful splenic infarction. Myelofibrosis appears to be part of the natural history of the disease but is a reactive, reversible process that does not itself

674 impede hematopoiesis and by itself has no prognostic significance. In approximately 15% of patients, however, myelofibrosis is accompanied by significant extramedullary hematopoiesis, hepatosplenomegaly, and transfusion-dependent anemia, which are manifestations of stem cell failure. The organomegaly can cause significant mechanical discomfort, portal hypertension, and progressive cachexia. Although the incidence of acute nonlymphocytic leukemia is increased in PV, the incidence of acute leukemia in patients not exposed to chemotherapy or radiation therapy is low. Interestingly, chemotherapy, including hydroxyurea, has been associated with acute leukemia in *JAK2* V617F-negative stem cells in some PV patients. *Erythromelalgia* is a curious syndrome of unknown etiology associated with thrombocytosis, primarily involving the lower extremities and usually manifested by erythema, warmth, and pain of the affected appendage and occasionally digital infarction. It occurs with a variable frequency and is usually responsive to salicylates. Some of the central nervous system symptoms observed in patients with PV, such as ocular migraine, appear to represent a variant of erythromelalgia.

Left uncontrolled, erythrocytosis can lead to thrombosis involving vital organs such as the liver, heart, brain, or lungs. Patients with massive splenomegaly are particularly prone to thrombotic events because the associated increase in plasma volume masks the true extent of the red cell mass elevation measured by the hematocrit or hemoglobin level. A “normal” hematocrit or hemoglobin level in a PV patient with massive splenomegaly should be considered indicative of an elevated red cell mass until proven otherwise.

TREATMENT POLYCYTHEMIA VERA

PV is generally an indolent disorder, the clinical course of which is measured in decades, and its management should reflect its tempo. Thrombosis due to erythrocytosis is the most significant complication and often the presenting manifestation, and maintenance of the hemoglobin level at ≤ 140 g/L (14 g/dL; hematocrit $<45\%$) in men and ≤ 120 g/L (12 g/dL; hematocrit $<42\%$) in women is mandatory to avoid thrombotic complications. Phlebotomy serves initially to reduce hyperviscosity by bringing the red cell mass into the normal range while further expanding the plasma volume. Periodic phlebotomies thereafter serve to maintain the red cell mass within the normal range and to induce a state of iron deficiency that prevents an accelerated reexpansion of the red cell mass. In most PV patients, once an iron-deficient state is achieved, phlebotomy is usually only required at 3-month intervals. Neither phlebotomy nor iron deficiency increases the platelet count relative to the effect of the disease itself, and thrombocytosis is not correlated with thrombosis in PV, in contrast to the strong correlation between erythrocytosis and thrombosis in this disease. The use of salicylates as a tonic against thrombosis in PV patients is not only potentially harmful if the red cell mass is not controlled by phlebotomy, but is also an unproven remedy. Anticoagulants are only indicated when a thrombosis has occurred and can be difficult to monitor if the red cell mass is substantially elevated owing to the artifactual imbalance between the test tube anticoagulant and plasma that occurs when blood from these patients is assayed for prothrombin or partial thromboplastin activity. Asymptomatic hyperuricemia (<10 mg/dL) requires no therapy, but allopurinol should be administered to avoid further elevation of the uric acid when chemotherapy is used to reduce splenomegaly or leukocytosis or to treat pruritus. Generalized pruritus intractable to antihistamines or antidepressants such as doxepin can be a major problem in PV; interferon α (IFN- α), psoralens with ultraviolet light in the A range (PUVA) therapy, and hydroxyurea are other methods of palliation. Asymptomatic thrombocytosis requires no therapy unless the platelet count is sufficiently high to cause bleeding due an acquired form of von Willebrand’s disease in which there is adsorption and proteolysis of high-molecular-weight von Willebrand factor (VWF) multimers by the expanded platelet mass. Symptomatic splenomegaly can be treated with pegylated IFN- α . Pegylated IFN- α can also produce complete hematologic and molecular remissions in PV, and its role in this disorder is

currently under investigation. Anagrelide, a phosphodiesterase inhibitor, can reduce the platelet count and, if tolerated, is preferable to hydroxyurea because it lacks marrow toxicity and is protective against venous thrombosis. A reduction in platelet number may be necessary for the treatment of erythromelalgia or ocular migraine if salicylates are not effective or if the platelet count is sufficiently high to increase the risk of hemorrhage but only to the degree that symptoms are alleviated. Alkylating agents and radioactive sodium phosphate (^{32}P) are leukemogenic in PV, and their use should be avoided. If a cytotoxic agent must be used, hydroxyurea is preferred, but this drug does not prevent either thrombosis or myelofibrosis in PV, is itself leukemogenic, and should be used for as short a time as possible. Previously, PV patients with massive splenomegaly unresponsive to reduction by chemotherapy or interferon required splenectomy. However, with the introduction of the nonspecific *JAK2* inhibitor ruxolitinib, it has been possible in the majority of patients with PV complicated by myelofibrosis and myeloid metaplasia to reduce spleen size while at the same time alleviating constitutional symptoms due to cytokine release. This drug is currently undergoing clinical trials in PV patients intolerant of hydroxyurea. In some patients with end-stage disease, pulmonary hypertension may develop due to fibrosis or extramedullary hematopoiesis. A role for allogeneic bone marrow transplantation in PV has not been defined.

Most patients with PV can live long lives without functional impairment when their red cell mass is effectively managed with phlebotomy alone. Chemotherapy is never indicated to control the red cell mass unless venous access is inadequate.

PRIMARY MYELOFIBROSIS

Chronic PMF (other designations include *idiopathic myelofibrosis*, *agnogenic myeloid metaplasia*, or *myelofibrosis with myeloid metaplasia*) is a clonal disorder of a multipotent hematopoietic progenitor cell of unknown etiology characterized by marrow fibrosis, extramedullary hematopoiesis, and splenomegaly. PMF is the least common chronic MPN, and establishing this diagnosis in the absence of a specific clonal marker is difficult because myelofibrosis and splenomegaly are also features of both PV and CML. Furthermore, myelofibrosis and splenomegaly also occur in a variety of benign and malignant disorders (Table 131-3), many of which are amenable to specific therapies not effective in PMF. In contrast to the other chronic MPNs and so-called acute or malignant myelofibrosis, which can occur at any age, PMF primarily afflicts men in their sixth decade or later.

ETIOLOGY

The etiology of PMF is unknown. Nonrandom chromosome abnormalities such as 9p, 20q-, 13q-, trisomy 8 or 9, or partial trisomy 1q are common, but no cytogenetic abnormality specific to the disease has been identified. *JAK2* V617F is present in approximately 50% of PMF patients, and mutations in the thrombopoietin receptor *Mpl* occur in about 5%. Most of the rest have mutations in the

TABLE 131-3 DISORDERS CAUSING MYELOFIBROSIS

Malignant	Nonmalignant
Acute leukemia (lymphocytic, myelogenous, megakaryocytic)	HIV infection
Chronic myeloid leukemia	Hyperparathyroidism
Hairy cell leukemia	Renal osteodystrophy
Hodgkin’s disease	Systemic lupus erythematosus
Primary myelofibrosis	Tuberculosis
Lymphoma	Vitamin D deficiency
Multiple myeloma	Thorium dioxide exposure
Myelodysplasia	Gray platelet syndrome
Metastatic carcinoma	
Polycythemia vera	
Systemic mastocytosis	

calreticulin gene (*CALR*) that alter the carboxy-terminal portion of the gene product. The degree of myelofibrosis and the extent of extramedullary hematopoiesis are also not related. Fibrosis in this disorder is associated with overproduction of transforming growth factor β and tissue inhibitors of metalloproteinases, whereas osteosclerosis is associated with overproduction of osteoprotegerin, an osteoclast inhibitor. Marrow angiogenesis occurs due to increased production of vascular endothelial growth factor. Importantly, fibroblasts in PMF are polyclonal and not part of the neoplastic clone.

CLINICAL FEATURES

No signs or symptoms are specific for PMF. Many patients are asymptomatic at presentation, and the disease is usually detected by the discovery of splenic enlargement and/or abnormal blood counts during a routine examination. However, in contrast to its companion MPN, night sweats, fatigue, and weight loss are common presenting complaints. A blood smear will show the characteristic features of extramedullary hematopoiesis: teardrop-shaped red cells, nucleated red cells, myelocytes, and promyelocytes; myeloblasts may also be present (Fig. 131-1). Anemia, usually mild initially, is the rule, whereas the leukocyte and platelet counts are either normal or increased, but either can be depressed. Mild hepatomegaly may accompany the splenomegaly but is unusual in the absence of splenic enlargement; isolated lymphadenopathy should suggest another diagnosis. Both serum lactate dehydrogenase and alkaline phosphatase levels can be elevated. The LAP score can be low, normal, or high. Marrow is usually inaspirable due to the myelofibrosis (Fig. 131-2), and bone x-rays may reveal osteosclerosis. Exuberant extramedullary hematopoiesis can cause ascites; portal, pulmonary, or intracranial hypertension; intestinal or ureteral obstruction; pericardial tamponade; spinal cord compression; or skin nodules. Splenic enlargement can be sufficiently rapid to cause splenic infarction with fever and pleuritic chest pain. Hyperuricemia and secondary gout may ensue.

DIAGNOSIS

While the clinical picture described above is characteristic of PMF, all of the clinical features described can also be observed in PV or CML. Massive splenomegaly commonly masks erythrocytosis in PV, and reports of intraabdominal thrombosis in PMF most likely represent instances of unrecognized PV. In some patients with PMF, erythrocytosis has developed during the course of the disease. Furthermore, because many other disorders have features that overlap with PMF but respond to distinctly different therapies, the diagnosis of PMF is one of exclusion, which requires that the disorders listed in Table 131-3 be ruled out.

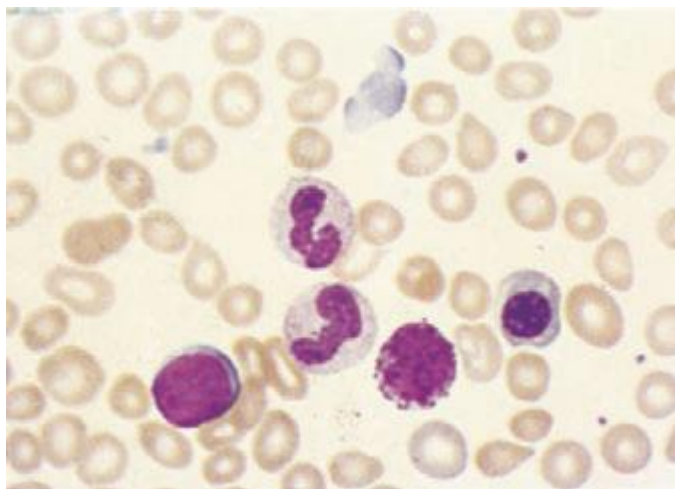


FIGURE 131-1 Teardrop-shaped red blood cells indicative of membrane damage from passage through the spleen, a nucleated red blood cell, and immature myeloid cells indicative of extramedullary hematopoiesis are noted. This peripheral blood smear is related to any cause of extramedullary hematopoiesis.

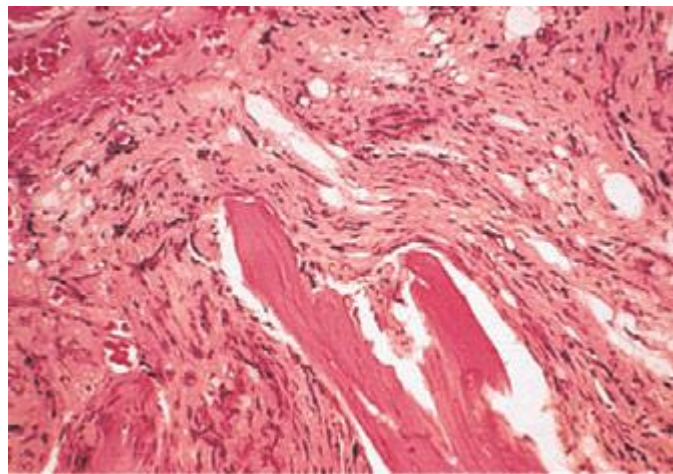


FIGURE 131-2 This marrow section shows the marrow cavity replaced by fibrous tissue composed of reticulin fibers and collagen. When this fibrosis is due to a primary hematologic process, it is called *myelofibrosis*. When the fibrosis is secondary to a tumor or a granulomatous process, it is called *myelophthisis*.

The presence of teardrop-shaped red cells, nucleated red cells, myelocytes, and promyelocytes establishes the presence of extramedullary hematopoiesis, while the presence of leukocytosis, thrombocytosis with large and bizarre platelets, and circulating myelocytes suggests the presence of an MPN as opposed to a secondary form of myelofibrosis (Table 131-3). Marrow is usually inaspirable due to increased marrow reticulin, but marrow biopsy will reveal a hypercellular marrow with trilineage hyperplasia and, in particular, increased numbers of megakaryocytes in clusters and with large, dysplastic nuclei. However, there are no characteristic bone marrow morphologic abnormalities that distinguish PMF from the other chronic MPNs. Splenomegaly due to extramedullary hematopoiesis may be sufficiently massive to cause portal hypertension and variceal formation. In some patients, exuberant extramedullary hematopoiesis can dominate the clinical picture. An intriguing feature of PMF is the occurrence of autoimmune abnormalities such as immune complexes, antinuclear antibodies, rheumatoid factor, or a positive Coombs' test. Whether these represent a host reaction to the disorder or are involved in its pathogenesis is unknown. Cytogenetic analysis of blood is useful both to exclude CML and for prognostic purposes, because complex karyotype abnormalities portend a poor prognosis in PMF. For unknown reasons, the number of circulating CD34+ cells is markedly increased in PMF (>15,000/ μ L) compared to the other chronic MPNs, unless they too develop myeloid metaplasia.

Importantly, approximately 50% of PMF patients, like patients with its companion myeloproliferative disorders PV and ET, express the *JAK2* V617F mutation, often as homozygotes. Such patients are usually older and have higher hematocrits than the patients who are *JAK2* V617F-negative, whereas PMF patients expressing an *MPL* mutation tend to be more anemic and have lower leukocyte counts. Somatic mutations in exon 9 of the calreticulin gene (*CALR*) have been found in a majority of patients with PMF and ET who lack mutations in either *JAK2* or *MPL*, and their clinical course appears to be more indolent than patients expressing either a *JAK2* or an *MPL* mutation.

COMPLICATIONS

Survival in PMF varies according to specific risk factors at diagnosis (Tables 131-4 and 131-5) but is shorter in most patients than in PV or ET patients. The natural history of PMF is one of increasing marrow failure with transfusion-dependent anemia and increasing organomegaly due to extramedullary hematopoiesis. As with CML, PMF can evolve from a chronic phase to an accelerated phase with constitutional symptoms and increasing marrow failure. About 10% of patients spontaneously transform to an aggressive form of acute leukemia for which therapy is usually ineffective. Additional important prognostic

TABLE 131-4 THREE CURRENT SCORING SYSTEMS FOR ESTIMATING PROGNOSIS IN PMF PATIENTS

Risk Factor	IPSS (2009) ^a	DIPSS (2010) ^b	DIPSS Plus (2011) ^c
Anemia (<10 g/dL)	X	X	X
Leukocytosis (>25,000/ μ L)	X	X	X
Peripheral blood blasts (\geq 1%)	X	X	X
Constitutional symptoms	X	X	X
Age (>65 years)	X	X	X
Unfavorable karyotype			X
Platelet count (<100,000/ μ L)			X
Transfusion dependence			X

^aBlood 113:2895, 2009. ^bBlood 115:1703, 2010. ^cJ Clin Oncol 29:392, 2011.

Note: The Dynamic International Prognostic Scoring System (DIPSS) was developed to determine if the International Prognostic Scoring System (IPSS) risk factors identified as important for survival at the time of primary myelofibrosis (PMF) diagnosis could also be used for risk stratification following their acquisition during the course of the disease. One point is assigned to each risk factor for IPSS scoring. For DIPSS, the same is true, but age >65 years, anemia, blood blasts, and constitutional symptoms are assigned 2 points each. The DIPSS Plus scoring system represents recognition that the addition of unfavorable karyotype, thrombocytopenia, and transfusion dependence improved the DIPSS risk stratification system for which additional points are assigned (Table 131-5). More recent studies suggest that mutational analysis of the *ASXL1*, *EZH2*, *SRSF2*, and *IDH1/2* genes further improves risk stratification for survival and leukemic transformation (Leukemia 27:1861, 2013).

factors for disease acceleration during the course of PMF include the presence of complex cytogenetic abnormalities, thrombocytopenia, and transfusion-dependent anemia. Most recently, mutations in the *ASXL1*, *EZH2*, *SRSF2*, and *IDH1/2* genes have been identified as risk factors for early death or transformation to acute leukemia and may prove to be more useful for PMF risk assessment than any clinical scoring system.

TREATMENT PRIMARY MYELOFIBROSIS

No specific therapy exists for PMF. The causes for anemia are multifarious and include ineffective erythropoiesis uncompensated by splenic extramedullary hematopoiesis, hemodilution due to splenomegaly, splenic sequestration, blood loss secondary to thrombocytopenia or portal hypertension, folic acid deficiency, systemic inflammation, and autoimmune hemolysis. Neither recombinant erythropoietin nor androgens such as danazol have proven to be consistently effective as therapy for anemia. Erythropoietin may worsen splenomegaly and will be ineffective if the serum erythropoietin level is >125 mU/L. Given the inflammatory milieu that characterizes PMF, corticosteroids can ameliorate anemia as well as constitutional symptoms such as fever, chills, night sweats, anorexia, and weight loss, and low-dose thalidomide together with prednisone has proved effective as well. Thrombocytopenia can be due to impaired marrow function, splenic sequestration, or autoimmune destruction

TABLE 131-5 IPSS AND DIPSS RISK STRATIFICATION SYSTEMS

Risk Categories ^a	Number of Risk Factors		
	IPSS	DIPSS	DIPSS PLUS
Low	0	0	0
Intermediate-1	1	1–2	1
Intermediate-2	2	3–4	2–3
High	\geq 3	>4	4–6

^aThe corresponding survival curves for each risk category can be found in the references cited in the footnotes of Table 131-4.

Abbreviations: DIPSS, Dynamic International Prognostic Scoring System; IPSS, International Prognostic Scoring System.

and may also respond to low-dose thalidomide together with prednisone. Splenomegaly is by far the most distressing and intractable problem for PMF patients, causing abdominal pain, portal hypertension, easy satiety, and cachexia, whereas surgical removal of a massive spleen is associated with significant postoperative complications including mesenteric venous thrombosis, hemorrhage, rebound leukocytosis and thrombocytosis, and hepatic extramedullary hematopoiesis with no amelioration of either anemia or thrombocytopenia when present. For unexplained reasons, splenectomy also increases the risk of blastic transformation. Splenic irradiation is, at best, temporarily palliative and associated with a significant risk of neutropenia, infection, and subsequent operative hemorrhage if splenectomy is attempted. Allopurinol can control significant hyperuricemia, and bone pain can be alleviated by local irradiation. The role of IFN- α is still undefined; its side effects are more pronounced in the older individuals, and it may exacerbate the bone marrow failure. The JAK2 inhibitor, ruxolitinib, has proved effective in reducing splenomegaly and alleviating constitutional symptoms in a majority of advanced PMF patients while also prolonging survival, although it does not significantly influence the *JAK2* V617F allele burden. Although anemia and thrombocytopenia are its major side effects, these are dose-dependent, and with time, anemia stabilizes and thrombocytopenia may improve. Allogeneic bone marrow transplantation is the only curative treatment for PMF and should be considered in younger patients; nonmyeloablative conditioning regimens may permit hematopoietic cell transplantation to be extended to older individuals, but this approach is currently under investigation.

ESSENTIAL THROMBOCYTOSIS

Essential thrombocytosis (other designations include *essential thrombocythemia*, *idiopathic thrombocytosis*, *primary thrombocytosis*, and *hemorrhagic thrombocythemia*) is a clonal disorder of unknown etiology involving a multipotent hematopoietic progenitor cell manifested clinically by overproduction of platelets without a definable cause. ET is an uncommon disorder, with an incidence of 1–2/100,000 and a distinct female predominance. No clonal marker is available to consistently distinguish ET from the more common nonclonal, reactive forms of thrombocytosis (Table 131-6), making its diagnosis difficult. Once considered a disease of the elderly and responsible for significant morbidity due to hemorrhage or thrombosis, with the widespread use of electronic cell counters, it is now clear that ET can occur at any age in adults and often without symptoms or disturbances of hemostasis. There is an unexplained female predominance in contrast to PMF or the reactive forms of thrombocytosis where no sex difference exists. Because no specific clonal marker is available, clinical criteria have been proposed to distinguish ET from the other chronic MPNs, which may also present with thrombocytosis but have differing prognoses and therapies (Table 131-6). These criteria do not establish clonality; therefore, they are truly useful only in identifying disorders such as CML, PV, or myelodysplasia, which can masquerade as ET, as opposed to actually establishing the presence of ET. Furthermore, as with “idiopathic” erythrocytosis, nonclonal benign forms of thrombocytosis

TABLE 131-6 CAUSES OF THROMBOCYTOSIS

Tissue inflammation: collagen vascular disease, inflammatory bowel disease	Hemorrhage
Malignancy	Iron-deficiency anemia
Infection	Surgery
Myeloproliferative disorders: polycythemia vera, primary myelofibrosis, essential thrombocytosis, chronic myelogenous leukemia	Rebound: Correction of vitamin B ₁₂ or folate deficiency, post-ethanol abuse
Myelodysplastic disorders: 5q–syndrome, idiopathic refractory sideroblastic anemia	Hemolysis
Postsplenectomy or hyposplenism	Familial: Thrombopoietin overproduction, <i>MPL</i> mutations

exist (such as hereditary overproduction of thrombopoietin) that are not widely recognized because we currently lack adequate diagnostic tools. Approximately 50% of ET patients carry the *JAK2* V617F mutation, but its absence does not exclude the disorder.

ETIOLOGY

Megakaryocytopoiesis and platelet production depend on thrombopoietin and its receptor *Mpl*. As in the case of early erythroid and myeloid progenitor cells, early megakaryocytic progenitors require the presence of interleukin 3 (IL-3) and stem cell factor for optimal proliferation in addition to thrombopoietin. Their subsequent development is also enhanced by the chemokine stromal cell-derived factor 1 (SDF-1). However, megakaryocyte maturation requires thrombopoietin.

Megakaryocytes are unique among hematopoietic progenitor cells because reduplication of their genome is endomitotic rather than mitotic. In the absence of thrombopoietin, endomitotic megakaryocytic reduplication and, by extension, the cytoplasmic development necessary for platelet production are impaired. Like erythropoietin, thrombopoietin is produced in both the liver and the kidneys, and an inverse correlation exists between the platelet count and plasma thrombopoietic activity. Unlike erythropoietin, thrombopoietin is only constitutively produced, and the plasma thrombopoietin level is controlled by the size of its progenitor cell pool. Also, in contrast to erythropoietin, but like its myeloid counterparts, granulocyte and granulocyte-macrophage colony-stimulating factors, thrombopoietin not only enhances the proliferation of its target cells but also enhances the reactivity of their end-stage product, the platelet. In addition to its role in thrombopoiesis, thrombopoietin also enhances the survival of multipotent hematopoietic stem cells and their bone marrow residence.

The clonal nature of ET was established by analysis of glucose-6-phosphate dehydrogenase isoenzyme expression in patients hemizygous for this gene, by analysis of X-linked DNA polymorphisms in informative female patients, and by the expression in patients of nonrandom, though variable, cytogenetic abnormalities. Although thrombocytosis is its principal manifestation, like the other chronic MPNs, a multipotent hematopoietic progenitor cell is involved in ET. Furthermore, a number of families have been described in which ET was inherited, in one instance as an autosomal dominant trait. In addition to ET, PMF and PV have also been observed in some kindreds. Like PMF, most patients who do not have *JAK2* mutations have *CALR* mutations.

CLINICAL FEATURES

Clinically, ET is most often identified incidentally when a platelet count is obtained during the course of a routine medical evaluation. Occasionally, review of previous blood counts will reveal that an elevated platelet count was present but overlooked for many years. No symptoms or signs are specific for ET, but these patients can have hemorrhagic and thrombotic tendencies expressed as easy bruising for the former and microvascular occlusive events for the latter such as erythromelalgia, ocular migraine, or a TIA. Physical examination is generally unremarkable except occasionally for mild splenomegaly. Splenomegaly is indicative of another MPN, in particular PV, PMF, or CML.

Anemia is unusual, but a mild neutrophilic leukocytosis is not. The blood smear is most remarkable for the number of platelets present, some of which may be very large. The large mass of circulating platelets may prevent the accurate measurement of serum potassium due to release of platelet potassium upon blood clotting. This type of hyperkalemia is a laboratory artifact and not associated with electrocardiographic abnormalities. Similarly, arterial oxygen measurements can be inaccurate unless thrombocythemic blood is collected on ice. The prothrombin and partial thromboplastin times are normal, whereas abnormalities of platelet function such as a prolonged bleeding time and impaired platelet aggregation can be present. However, despite much study, no platelet function abnormality is characteristic of ET, and no platelet function test predicts the risk of clinically significant bleeding or thrombosis.

The elevated platelet count may hinder marrow aspiration, but marrow biopsy usually reveals megakaryocyte hypertrophy and hyperplasia, as well as an overall increase in marrow cellularity. If marrow reticulin is increased, another diagnosis should be considered. The absence of stainable iron demands an explanation because iron deficiency alone can cause thrombocytosis, and absent marrow iron in the presence of marrow hypercellularity is a feature of PV.

Nonrandom cytogenetic abnormalities occur in ET but are uncommon, and no specific or consistent abnormality is notable, even those involving chromosomes 3 and 1, where the genes for thrombopoietin and its receptor *Mpl*, respectively, are located.

DIAGNOSIS

Thrombocytosis is encountered in a broad variety of clinical disorders (Table 131-6), in many of which production of cytokines is increased. The absolute level of the platelet count is not a useful diagnostic aid for distinguishing between benign and clonal causes of thrombocytosis. About 50% of ET patients express the *JAK2* V617F mutation. When *JAK2* V617F is absent, cytogenetic evaluation is mandatory to determine if the thrombocytosis is due to CML or a myelodysplastic disorder such as the 5q- syndrome. Because the bcr-abl translocation can be present in the absence of the Ph chromosome, and because bcr-abl reverse transcriptase polymerase chain reaction is associated with false-positive results, fluorescence in situ hybridization (FISH) analysis for bcr-abl is the preferred assay in patients with thrombocytosis in whom a cytogenetic study for the Ph chromosome is negative. *CALR* mutations are present in most patients who do not have *JAK2* mutations, but diagnostic tools to detect these mutations are not yet widespread. Anemia and ringed sideroblasts are not features of ET, but they are features of idiopathic refractory sideroblastic anemia, and in some of these patients, the thrombocytosis occurs in association with *JAK2* V617F expression. Splenomegaly should suggest the presence of another MPN, and in this setting, a red cell mass determination should be performed because splenomegaly can mask the presence of erythrocytosis. Importantly, what appears to be ET can evolve into PV or PMF after a period of many years, revealing the true nature of the underlying MPN. There is sufficient overlap of the *JAK2* V617F neutrophil allele burden between ET and PV that this cannot be used as a distinguishing diagnostic feature; only a red cell mass and plasma volume determination can distinguish PV from ET, and importantly in this regard, 64% of *JAK2* V617F-positive ET patients actually were found to have PV when red cell mass and plasma volume determinations were performed.

COMPLICATIONS

Perhaps no other condition in clinical medicine has caused otherwise astute physicians to intervene inappropriately more often than thrombocytosis, particularly if the platelet count is $>1 \times 10^6/\mu\text{L}$. It is commonly believed that a high platelet count causes intravascular stasis and thrombosis; however, no controlled clinical study has ever established this association, and in patients younger than age 60 years, the incidence of thrombosis was not greater in patients with thrombocytosis than in age-matched controls, and tobacco use appears to be the most important risk factor for thrombosis in ET patients.

To the contrary, very high platelet counts are associated primarily with hemorrhage due to acquired von Willebrand's disease. This is not meant to imply that an elevated platelet count cannot cause symptoms in an ET patient, but rather that the focus should be on the patient, not the platelet count. For example, some of the most dramatic neurologic problems in ET are migraine-related and respond only to lowering of the platelet count, whereas other symptoms such as erythromelalgia respond simply to platelet cyclooxygenase-1 inhibitors such as aspirin or ibuprofen, without a reduction in platelet number. Still others may represent an interaction between an atherosclerotic vascular system and a high platelet count, and others may have no relationship to the platelet count whatsoever. Recognition that PV can present with thrombocytosis alone as well as the discovery of previously unrecognized causes of hypercoagulability (Chap. 142) make the older literature on the complications of thrombocytosis unreliable.

678 ET can also evolve into PMF, but whether this is a feature of ET or represents PMF presenting initially with isolated thrombocytosis is unknown.

TREATMENT ESSENTIAL THROMBOCYTOSIS

Survival of patients with ET is not different than for the general population. An elevated platelet count in an asymptomatic patient without cardiovascular risk factors requires no therapy. Indeed, before any therapy is initiated in a patient with thrombocytosis, the cause of symptoms must be clearly identified as due to the elevated platelet count. When the platelet count rises above $1 \times 10^6/\mu\text{L}$, a substantial quantity of high-molecular-weight von Willebrand multimers are removed from the circulation and destroyed by the enlarged platelet mass, resulting in an acquired form of von Willebrand's disease. This can be identified by a reduction in ristocetin cofactor activity. In this situation, aspirin could promote hemorrhage. Bleeding in this situation usually responds to ϵ -aminocaproic acid, which can be given prophylactically before and after elective surgery. Plateletpheresis is at best a temporary and inefficient remedy that is rarely required. Importantly, ET patients treated with ^{32}P or alkylating agents are at risk of developing acute leukemia without any proof of benefit; combining either therapy with hydroxyurea increases this risk. If platelet reduction is deemed necessary on the basis of symptoms refractory to salicylates alone, pegylated IFN- α , the quinazoline derivative, anagrelide, or hydroxyurea can be used to reduce the platelet count, but none of these is uniformly effective or without significant side effects. Hydroxyurea and aspirin are more effective than anagrelide and aspirin for prevention of TIAs but not more effective for the prevention of other types of arterial thrombosis and are actually less effective for venous thrombosis. The effectiveness of hydroxyurea in preventing TIAs is because it is an NO donor. Normalizing the platelet count also does not prevent either arterial or venous thrombosis. The risk of gastrointestinal bleeding is also higher when aspirin is combined with anagrelide.

As more clinical experience is acquired, ET appears more benign than previously thought. Evolution to acute leukemia is more likely to be a consequence of therapy than of the disease itself. In managing patients with thrombocytosis, the physician's first obligation is to do no harm.

Heredity Certain syndromes with somatic cell chromosome aneuploidy, such as trisomy 21 noted in Down syndrome, are associated with an increased incidence of AML. Inherited diseases with defective DNA repair, e.g., Fanconi anemia, Bloom syndrome, and ataxia-telangiectasia, are also associated with AML. Congenital neutropenia (Kostmann syndrome) is a disease with mutations in the genes encoding the granulocyte colony-stimulating factor (G-CSF) receptor and, often, neutrophil elastase that may evolve into AML. Germline mutations of CCAAT/enhancer-binding protein α (*CEBPA*), runt-related transcription factor 1 (*RUNX1*), and tumor protein p53 (*TP53*) have also been associated with a higher predisposition to AML in some series.

Radiation High-dose radiation, like that experienced by survivors of the atomic bombs in Japan or nuclear reactor accidents, increases the risk of myeloid leukemias that peaks 5–7 years after exposure. Therapeutic radiation alone seems to add little risk of AML but can increase the risk in people also exposed to alkylating agents.

Chemical and Other Exposures Exposure to benzene, a solvent used in the chemical, plastic, rubber, and pharmaceutical industries, is associated with an increased incidence of AML. Smoking and exposure to petroleum products, paint, embalming fluids, ethylene oxide, herbicides, and pesticides have also been associated with an increased risk of AML.

Drugs Anticancer drugs are the leading cause of therapy-associated AML. Alkylating agent-associated leukemias occur on average 4–6 years after exposure, and affected individuals have aberrations in chromosomes 5 and 7. Topoisomerase II inhibitor-associated leukemias occur 1–3 years after exposure, and affected individuals often have aberrations involving chromosome 11q23. Newer agents for treatment of other hematopoietic malignancies and solid tumors are also under scrutiny for increased risk of AML. Chloramphenicol, phenylbutazone, and, less commonly, chloroquine and methoxyypsoralen can result in bone marrow failure that may evolve into AML.

CLASSIFICATION

The current categorization of AML uses the World Health Organization (WHO) classification (Table 132-1), which includes different biologically distinct groups based on clinical features and cytogenetic and molecular abnormalities in addition to morphology. In contrast to the previously used French-American-British (FAB) schema, the WHO classification places limited reliance on cytochemistry. A major difference between the WHO and the FAB systems is the blast cutoff for a diagnosis of AML as opposed to myelodysplastic syndrome (MDS); it is 20% in the WHO classification and 30% in the FAB. However, within the WHO classification, specific chromosomal rearrangements, i.e., t(8;21)(q22;q22), inv(16)(p13.1q22), t(16;16)(p13.1;q22), and t(15;17)(q22;q12), define AML even with <20% blasts.

Immunophenotype and Relevance to the WHO Classification The immunophenotype of human leukemia cells can be studied by multiparameter flow cytometry after the cells are labeled with monoclonal antibodies to cell-surface antigens. This can be important for separating AML from acute lymphoblastic leukemia (ALL) and identifying some subtypes of AML. For example, AML with minimal differentiation that is characterized by immature morphology and no lineage-specific cytochemical reactions may be diagnosed by flow-cytometric demonstration of the myeloid-specific antigens cluster designation (CD) 13 and/or 117. Similarly, acute megakaryoblastic leukemia can often be diagnosed only by expression of the platelet-specific antigens CD41 and/or CD61. Although flow cytometry is useful, widely used, and in some cases essential for the diagnosis of AML, it is supportive only in establishing the different subtypes of AML through the WHO classification.

Clinical Features and Relevance to the WHO Classification The WHO classification also considers clinical features in subdividing AML. For example, it identifies therapy-related AML as a separate entity that develops following prior therapy (e.g., alkylating agents, topoisomerase II inhibitors, ionizing radiation). It also identifies AML with

132 Acute Myeloid Leukemia

Guido Marcucci, Clara D. Bloomfield

INCIDENCE

Acute myeloid leukemia (AML) is a neoplastic disease characterized by infiltration of the blood, bone marrow, and other tissues by proliferative, clonal undifferentiated cells of the hematopoietic system. These leukemias comprise a spectrum of malignancies that, untreated, range from rapidly fatal to slowly growing. In 2013, the estimated number of new AML cases in the United States was 14,590. The incidence of AML is ~3.5 per 100,000 people per year, and the age-adjusted incidence is higher in men than in women (4.5 vs 3.1). AML incidence increases with age; it is 1.7 in individuals age <65 years and 15.9 in those age >65 years. The median age at diagnosis is 67 years.

ETIOLOGY

Heredity, radiation, chemical and other occupational exposures, and drugs have been implicated in the development of AML. No direct evidence suggests a viral etiology.

TABLE 132-1 WORLD HEALTH ORGANIZATION CLASSIFICATION OF ACUTE MYELOID LEUKEMIA (AML) AND RELATED NEOPLASMS^a**AML with recurrent genetic abnormalities**

- AML with t(8;21)(q22;q22); *RUNX1-RUNX1T1*^b
- AML with inv(16)(p13.1;q22) or t(16;16)(p13.1;q22); *CBFB-MYH11*^b
- Acute promyelocytic leukemia with t(15;17)(q22;q12); *PML-RARA*^b
- AML with t(9;11)(p22;q23); *MLLT3-MLL*
- AML with t(6;9)(p23;q34); *DEK-NUP214*
- AML with inv(3)(q21q26.2) or t(3;3)(q21;q26.2); *RPN1-EV1*
- AML (megakaryoblastic) with t(1;22)(p13;q13); *RBM15-MKL1*
- Provisional entity: AML with mutated *NPM1*
- Provisional entity: AML with mutated *CEBPA*

AML with myelodysplasia-related changes**Therapy-related myeloid neoplasms****AML, not otherwise specified**

- AML with minimal differentiation
- AML without maturation
- AML with maturation
- Acute myelomonocytic leukemia
- Acute monoblastic and monocytic leukemia
- Acute erythroid leukemia
- Acute megakaryoblastic leukemia
- Acute basophilic leukemia
- Acute panmyelosis with myelofibrosis

Myeloid sarcoma**Myeloid proliferations related to Down syndrome**

- Transient abnormal myelopoiesis
- Myeloid leukemia associated with Down syndrome

Blastic plasmacytoid dendritic cell neoplasm

^aFrom SH Swerdlow et al (eds): *World Health Organization Classification of Tumours of Haematopoietic and Lymphoid Tissues*. Lyon, IARC Press, 2008. ^bDiagnosis is AML regardless of blast count.

myelodysplasia-related changes based in part on medical history of an antecedent MDS or myelodysplastic/myeloproliferative neoplasm. The clinical features likely contribute to the prognosis of AML and have therefore been included in the classification.

**Genetic Findings and Relevance to the WHO Classification**

The WHO classification uses clinical, morphologic, and cytogenetic and/or molecular criteria to identify subtypes of AML and forces the clinician to take the appropriate steps to correctly identify the entity and thus tailor treatment(s) accordingly. The WHO classification is indeed the first AML classification that incorporates genetic (chromosomal and molecular) information. In this classification, subtypes of AML are recognized based on the presence or absence of specific recurrent genetic abnormalities. For example, the diagnosis of *acute promyelocytic leukemia* (APL) is based on the presence of either the t(15;17)(q22;q12) cytogenetic rearrangement or the *PML-RARA* fusion product of the translocation. A similar approach is taken with regard to core binding factor (CBF) AML that is now designated based on the presence of t(8;21)(q22;q22), inv(16)(p13.1;q22), or t(16;16)(p13.1;q22) or the respective fusion products *RUNX1-RUNX1T1* and *CBFB-MYH11*.

The WHO classification incorporates cytogenetics in the AML classification by recognizing a category of AML with recurrent genetic abnormalities and a category of AML with myelodysplasia-related changes (Table 132-1). The latter category is diagnosed not only by morphologic changes, but also in part by selected myelodysplasia-related cytogenetic abnormalities (e.g., complex karyotypes and unbalanced and balanced changes involving, among others, chromosomes 5, 7, and 11). Only one cytogenetic abnormality has been invariably associated with specific morphologic features: t(15;17)(q22;q12) with APL. Other chromosomal abnormalities have been associated primarily with one morphologic/immunophenotypic group, including inv(16)

(p13.1;q22) with AML with abnormal bone marrow eosinophils; t(8;21)(q22;q22) with slender Auer rods, expression of CD19, and increased normal eosinophils; and t(9;11)(p22;q23), and other translocations involving 11q23, with monocytic features. Recurring chromosomal abnormalities in AML may also be associated with specific clinical characteristics. More commonly associated with younger age are t(8;21) and t(15;17), and with older age, del(5q) and del(7q). Myeloid sarcomas (see below) are associated with t(8;21), and disseminated intravascular coagulation (DIC) is associated with t(15;17).

The WHO classification also incorporates molecular abnormalities by recognizing fusion genes that are products of recurrent cytogenetic aberrations or have been found mutated and may be involved in leukemogenesis. For instance, t(15;17) results in the fusion gene *PML-RARA* that encodes a chimeric protein, promyelocytic leukemia (Pml)–retinoic acid receptor α (Rara), which is formed by the fusion of the retinoic acid receptor α (*RARA*) gene from chromosome 17 and the promyelocytic leukemia (*PML*) gene from chromosome 15. The *RARA* gene encodes a member of the nuclear hormone receptor family of transcription factors. After binding retinoic acid, *RARA* can promote expression of a variety of genes. The 15;17 translocation juxtaposes *PML* with *RARA* in a head-to-tail configuration that is under the transcriptional control of *PML*. Three different breakpoints in the *PML* gene lead to various fusion protein isoforms. The Pml-Rara fusion protein tends to suppress gene transcription and blocks differentiation of the cells. Pharmacologic doses of the Rara ligand, all-*trans*-retinoic acid (tretinoin), relieve the block and promote hematopoietic cell differentiation (see below). Similar examples of molecular subtypes of the disease included in the category of AML with recurrent genetic abnormalities are those characterized by the leukemogenic fusion genes *RUNX1-RUNX1T1*, *CBFB-MYH11*, *MLLT3-MLL*, and *DEK-NUP214*, resulting, respectively, from t(8;21), inv(16) or t(16;16), t(9;11), and t(6;9)(p23;q34).

Two new provisional entities defined by the presence of gene mutations, rather than microscopic chromosomal abnormalities, have been added to the category of AML with recurrent genetic abnormalities: *AML with mutated nucleophosmin (nucleolar phosphoprotein B23, numatrin) (NPM1)* and *AML with mutated CEBPA*. AML with fms-related tyrosine kinase 3 (*FLT3*) mutations is not considered a distinct entity, although determining the presence of such mutations is recommended by WHO in patients with cytogenetically normal AML (CN-AML) because the relatively frequent *FLT3*-internal tandem duplication (ITD) carries a negative prognostic significance and therefore is clinically relevant. *FLT3* encodes a tyrosine kinase receptor important in the development of myeloid and lymphoid lineages. Activating mutations of *FLT3* are present in ~30% of adult AML patients due to ITDs in the juxtamembrane domain or point mutations of the activating loop of the kinase (called tyrosine kinase domain mutations). Aberrant activation of the *FLT3*-encoded protein provides increased proliferation and antiapoptotic signals to the myeloid progenitor cell. *FLT3*-ITD, the more common of the *FLT3* mutations, occurs preferentially in patients with CN-AML. The importance of identifying *FLT3*-ITD at diagnosis relates to the fact that not only is it a useful prognosticator but it also may predict response to specific treatment such as the tyrosine kinase inhibitors that are in clinical investigation.

PROGNOSTIC FACTORS

Several factors have been demonstrated to predict outcome of AML patients treated with chemotherapy, and they can be used for risk stratification and treatment guidance.

Chromosome findings at diagnosis are currently the most important independent prognostic factors. Several studies have categorized patients as having favorable, intermediate, or poor cytogenetic risk based on the presence of structural and/or numerical aberrations. Patients with t(15;17) have a very good prognosis (~85% cured), and those with t(8;21) and inv(16) have a good prognosis (~55% cured), whereas those with no cytogenetic abnormality have an intermediate outcome risk (~40% cured). Patients with a complex karyotype, t(6;9), inv(3), or -7 have a very poor prognosis. Another cytogenetic subgroup, the monosomal karyotype, has been suggested to adversely

680 impact the outcome of AML patients other than those with t(15;17), t(8;21), or inv(16) or t(16;16). The monosomal karyotype subgroup is defined by the presence of at least two autosomal monosomies (loss of chromosomes other than Y or X) or a single autosomal monosomy with additional structural abnormalities.

For patients lacking prognostic cytogenetic abnormalities, such as those with CN-AML, outcome prediction uses mutated or aberrantly expressed genes. *NPM1* mutations without concurrent presence of *FLT3*-ITD, and *CEBPA* mutations, especially if concurrently present in two different alleles, have been shown to predict favorable outcome, whereas *FLT3*-ITD predicts poor outcome. Given the proven prognostic importance of *NPM1* and *CEBPA* mutations and *FLT3*-ITD, molecular assessment of these genes at diagnosis has been incorporated in AML management guidelines by the National Comprehensive Cancer Network (NCCN) and the European LeukemiaNet (ELN). The same markers have also been incorporated in the definitions of the genetic groups of the ELN standardized reporting system, which are based on both cytogenetic and molecular abnormalities and used for comparing clinical features and treatment response among subsets of patients reported in different studies (Table 132-2). More recently, the prognostic impact of the genetic groups recognized by the ELN reporting system has been demonstrated. Thus, these genetic groups may also be used for risk stratification and treatment guidance.

In addition to *NPM1* and *CEBPA* mutations and *FLT3*-ITD, other molecular aberrations (Table 132-3) may in the future be routinely used for prognostication in AML and incorporated in the WHO classification and the ELN reporting system. Among these prognostic mutated genes are those encoding receptor tyrosine kinases (e.g., v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene homolog [*KIT*]), transcription factors (i.e., *RUNX1* and Wilms tumor 1 [*WT1*]), and epigenetic modifiers (i.e., additional sex combs like transcriptional regulator 1 [*ASXL1*], DNA (cytosine-5-)-methyltransferase 3 alpha [*DNMT3A*], isocitrate dehydrogenase 1 (NADP+), soluble [*IDH1*] and isocitrate dehydrogenase 2 (NADP+), mitochondrial [*IDH2*], lysine (K)-specific methyltransferase 2A [*KMT2A*, also known as *MLL*], and tet methylcytosine dioxygenase 2 [*TET2*]). Although *KIT* mutations are almost exclusively present in CBF AML and impact adversely the outcome, the remaining markers have been reported primarily in CN-AML. These gene mutations have been shown to be associated with outcome in multivariable analyses independently from other prognostic factors. However, for some of them, the prognostic impact (e.g., *TET2* mutations) or the type (adverse vs favorable) of prognostic impact (e.g., *IDH1*, *IDH2*) has been found in the majority, but not in all, of the reported studies.

An independent prognostic impact remains to be determined for mutated genes that are either associated primarily with unfavorable

TABLE 132-2 EUROPEAN LEUKEMIANET RECOMMENDED STANDARDIZED REPORTING FOR CORRELATION OF CYTOGENETIC AND MOLECULAR GENETIC DATA IN AML WITH CLINICAL DATA^a

Genetic Group	Subsets
Favorable	t(8;21)(q22;q22); <i>RUNX1</i> - <i>RUNX1T1</i> inv(16)(p13.1q22) or t(16;16)(p13.1;q22); <i>CBFB</i> - <i>MYH11</i> Mutated <i>NPM1</i> without <i>FLT3</i> -ITD (normal karyotype) Mutated <i>CEBPA</i> (normal karyotype)
Intermediate-I	Mutated <i>NPM1</i> and <i>FLT3</i> -ITD (normal karyotype) Wild-type <i>NPM1</i> and <i>FLT3</i> -ITD (normal karyotype) Wild-type <i>NPM1</i> without <i>FLT3</i> -ITD (normal karyotype)
Intermediate-II	t(9;11)(p22;q23); <i>MLL3</i> - <i>MLL</i> Cytogenetic abnormalities not classified as favorable or adverse
Adverse	inv(3)(q21q26.2) or t(3;3)(q21;q26.2); <i>RPN1</i> - <i>EV11</i> t(6;9)(p23;q34); <i>DEK</i> - <i>NUP214</i> t(v;11)(v;q23); <i>MLL</i> rearranged -5 or del(5q); -7; abn(17p); complex karyotype (≥3 abnormalities)

^aH Döhner et al: Blood 115:453, 2010.

Abbreviation: ITD, internal tandem duplication.

TABLE 132-3 MOLECULAR PROGNOSTIC MARKERS IN AML

Gene Symbol	Gene Location	Prognostic Impact
Genes Included in the WHO Classification and ELN Reporting System		
<i>NPM1</i> mutations	5q35.1	Favorable
<i>CEBPA</i> mutations	19q13.1	Favorable
<i>FLT3</i> -ITD	13q12	Adverse
Genes Encoding Receptor Tyrosine Kinases		
<i>KIT</i> mutation	4q12	Adverse
<i>FLT3</i> -TKD	13q12	Adverse
Genes Encoding Transcription Factors		
<i>RUNX1</i> mutations	21q22.12	Adverse
<i>WT1</i> mutations	11p13	Adverse
Genes Encoding Epigenetic Modifiers		
<i>ASXL1</i> mutations	20q11.21	Adverse
<i>DNMT3A</i> mutations	2p23.3	Adverse
<i>IDH</i> mutations (<i>IDH1</i> and <i>IDH2</i>)	2q34 & 15q26.1	Adverse
<i>MLL</i> -PTD	11q23	Adverse
<i>TET2</i> mutations	4q24	Adverse
Deregulated Genes		
<i>BAALC</i> overexpression	8q22.3	Adverse
<i>ERG</i> overexpression	21q22.3	Adverse
<i>MN1</i> overexpression	22q12.1	Adverse
<i>EV11</i> overexpression	3q26.2	Adverse
Deregulated MicroRNAs		
<i>miR-155</i> overexpression	21q21.3	Adverse
<i>miR-3151</i> overexpression	8q22.3	Adverse
<i>miR-181a</i> overexpression	1q32.1 and 9q33.3	Favorable

Abbreviations: AML, acute myeloid leukemia; ELN, European LeukemiaNet; ITD, internal tandem duplication; PTD, partial tandem duplication; TKD, tyrosine kinase domain; WHO, World Health Organization.

cytogenetic aberrations (e.g., *TP53*) or are found with a relatively lower frequency in AML patients like those encoding epigenetic modifiers (e.g., enhancer of zeste 2 polycomb repressive complex 2 subunit [*EZH2*]), phosphatases (e.g., protein tyrosine phosphatase, non-receptor type 11 [*PTPN11*]), putative transcription factors (e.g., PHD finger protein 6 [*PHF6*]), splicing factors (e.g., U2 small nuclear RNA auxiliary factor 1 [*U2AF1*]), and proteins involved in chromosome segregation and genome stability (e.g., structural maintenance of chromosomes 1A [*SMC1A*] or structural maintenance of chromosomes 3 [*SMC3*]). Finally, other mutated genes are recognized as predictors of treatment response to distinct therapies rather than prognosticators; for example, neuroblastoma RAS viral (v-ras) oncogene homolog (*NRAS*) and Kirsten rat sarcoma viral oncogene homolog (*KRAS*) predict a better response to high-dose cytarabine in CBF AML.

In addition to gene mutations, deregulation of the expression levels of coding genes and of short noncoding RNAs (microRNAs) have been reported to provide prognostic information (Table 132-3). Overexpression of genes such as brain and acute leukemia, cytoplasmic (BAALC), v-ets avian erythroblastosis virus E26 oncogene homologue (avian) (*ERG*), meningioma (disrupted in balanced translocation) 1 (*MN1*), and MDS1 and EV11 complex locus (*MECOM*, also known as *EV11*) have been found to be predictive for poor outcome, especially in CN-AML. Similarly, deregulated expression levels of microRNAs, naturally occurring noncoding RNAs that have been shown to regulate the expression of proteins involved in hematopoietic differentiation and survival pathways by degradation or translation inhibition of target coding RNAs, have been associated with prognosis in AML. Overexpression of *miR-155* and *miR-3151* has been found to affect outcome adversely in CN-AML, whereas overexpression of *miR-181a* predicts a favorable outcome both in CN-AML and cytogenetically abnormal AML.

Because prognostic molecular markers in AML are not mutually exclusive and often occur concurrently (>80% patients have at least

two or more prognostic gene mutations), the likelihood that distinct marker combinations may be more informative than single markers is being recognized.

Epigenetic changes (e.g., DNA methylation) and microRNAs are often involved in deregulation of genes involved in hematopoiesis, contribute to leukemogenesis, and are often associated with the previously discussed prognostic gene mutations. These changes not only have been shown to provide biologic insights into leukemogenic mechanisms, but also independent prognostic information. Indeed, it is anticipated that with the enormous progress made in DNA and RNA sequencing technology, additional genetic and epigenetic aberrations will soon be discovered and will contribute to classification and reporting systems and outcome risk determination in AML patients.

In addition to cytogenetics and/or molecular aberrations, several other factors are associated with outcome in AML. Age at diagnosis is one of the most important risk factors. Advancing age is associated with a poorer prognosis not only because of its influence on the ability to survive induction therapy due to coexisting comorbidities, but also because with each successive decade of age, a greater proportion of patients have an intrinsically more resistant disease. A prolonged symptomatic interval with cytopenias preceding diagnosis or a history of antecedent hematologic disorders including myeloproliferative neoplasms is often found in older patients and is a clinical feature associated with a lower complete remission (CR) rate and shorter survival time. The CR rate is lower in patients who have had anemia, leukopenia, and/or thrombocytopenia for >3 months before the diagnosis of AML when compared to those without such a history. Responsiveness to chemotherapy declines as the duration of the antecedent disorder(s) increases. AML developing after treatment with cytotoxic agents for other malignancies is usually difficult to treat successfully. Finally, it is likely that AML in older patients is also associated with poor outcome because of the presence of distinct biologic features that may increase the aggressiveness of the disease and reduce the likelihood of treatment response. The leukemic cells in older patients more commonly express the multidrug resistance 1 (MDR1) efflux pump that conveys resistance to natural product–derived agents such as the anthracyclines that are frequently incorporated into the initial treatment. In addition, older patients less frequently harbor favorable cytogenetic abnormalities [i.e., t(8;21), inv(16), and t(16;16)] and more frequently harbor adverse cytogenetic (e.g., complex and monosomal karyotypes) and/or molecular (e.g., *ASXL1*, *IDH2*, *RUNX1*, *TET2*) abnormalities.

Other factors independently associated with worse outcome are a low performance status that influences ability to survive induction therapy and thus respond to treatment and a high presenting leukocyte count that in some series is an adverse prognostic factor for attaining a CR. Among patients with hyperleukocytosis (>100,000/ μ L), early central nervous system bleeding and pulmonary leukostasis contribute to poor outcome with initial therapy.

Achievement of CR is associated with better outcome and longer survival. CR is defined after examination of both blood and bone marrow. The blood neutrophil count must be \geq 1000/ μ L and the platelet count \geq 100,000/ μ L. Hemoglobin concentration is not considered in determining CR. Circulating blasts should be absent. Although rare blasts may be detected in the blood during marrow regeneration, they should disappear on successive studies. The bone marrow should contain <5% blasts, and Auer rods should be absent. Extramedullary leukemia should not be present. Patients who achieve CR after one induction cycle have longer CR durations than those requiring multiple cycles.

CLINICAL PRESENTATION

Symptoms Patients with AML most often present with nonspecific symptoms that begin gradually or abruptly and are the consequence of anemia, leukocytosis, leukopenia or leukocyte dysfunction, or thrombocytopenia. Nearly half have had symptoms for \leq 3 months before the leukemia was diagnosed.

Half of patients mention fatigue as the first symptom, but most complain of fatigue or weakness at the time of diagnosis. Anorexia and weight loss are common. Fever with or without an identifiable

infection is the initial symptom in approximately 10% of patients. Signs of abnormal hemostasis (bleeding, easy bruising) are noted first in 5% of patients. On occasion, bone pain, lymphadenopathy, nonspecific cough, headache, or diaphoresis is the presenting symptom.

Rarely patients may present with symptoms from a myeloid sarcoma that is a tumor mass consisting of myeloid blasts occurring at anatomic sites other than bone marrow. Sites involved are most commonly the skin, lymph node, gastrointestinal tract, soft tissue, and testis. This rare presentation, often characterized by chromosome aberrations [e.g., monosomy 7, trisomy 8, *MLL* rearrangement, inv(16), trisomy 4, t(8;21)], may precede or coincide with AML.

Physical Findings Fever, splenomegaly, hepatomegaly, lymphadenopathy, sternal tenderness, and evidence of infection and hemorrhage are often found at diagnosis. Significant gastrointestinal bleeding, intrapulmonary hemorrhage, or intracranial hemorrhage occurs most often in APL. Bleeding associated with coagulopathy may also occur in monocytic AML and with extreme degrees of leukocytosis or thrombocytopenia in other morphologic subtypes. Retinal hemorrhages are detected in 15% of patients. Infiltration of the gingivae, skin, soft tissues, or meninges with leukemic blasts at diagnosis is characteristic of the monocytic subtypes and those with 11q23 chromosomal abnormalities.

Hematologic Findings Anemia is usually present at diagnosis and can be severe. The degree varies considerably, irrespective of other hematologic findings, splenomegaly, or duration of symptoms. The anemia is usually normocytic normochromic. Decreased erythropoiesis often results in a reduced reticulocyte count, and red blood cell (RBC) survival is decreased by accelerated destruction. Active blood loss also contributes to the anemia.

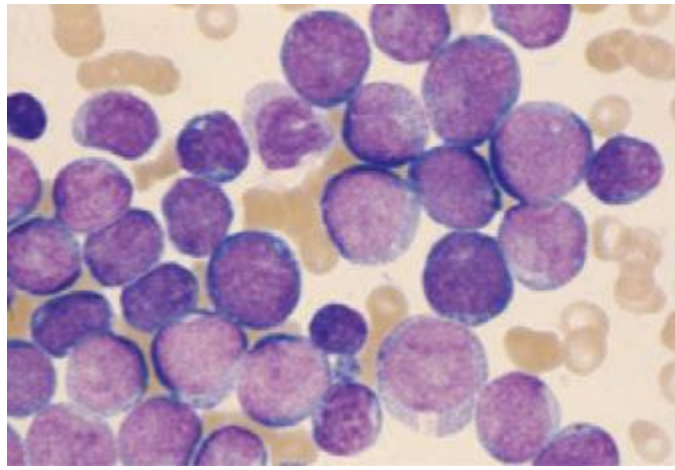
The median presenting leukocyte count is about 15,000/ μ L. Between 25 and 40% of patients have counts <5000/ μ L, and 20% have counts >100,000/ μ L. Fewer than 5% have no detectable leukemic cells in the blood. The morphology of the malignant cell varies in different subsets. In AML, the cytoplasm often contains primary (nonspecific) granules, and the nucleus shows fine, lacy chromatin with one or more nucleoli characteristic of immature cells. Abnormal rod-shaped granules called Auer rods are not uniformly present, but when they are, myeloid lineage is virtually certain (Fig. 132-1). Poor neutrophil function may be noted functionally by impaired phagocytosis and migration and morphologically by abnormal lobulation and deficient granulation.

Platelet counts <100,000/ μ L are found at diagnosis in ~75% of patients, and about 25% have counts <25,000/ μ L. Both morphologic and functional platelet abnormalities can be observed, including large and bizarre shapes with abnormal granulation and inability of platelets to aggregate or adhere normally to one another.

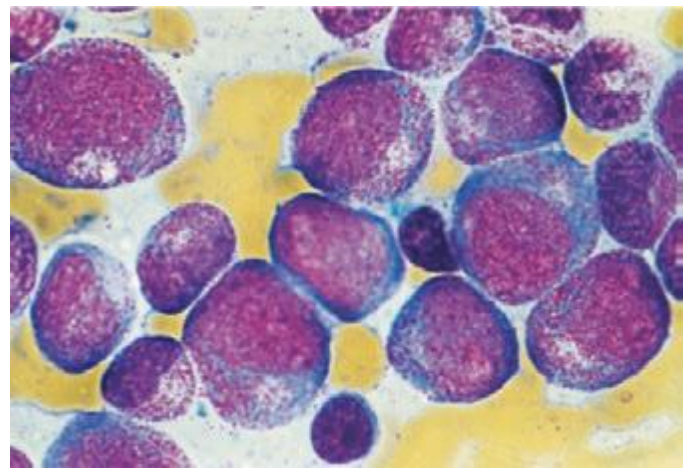
Pretreatment Evaluation Once the diagnosis of AML is suspected, a rapid evaluation and initiation of appropriate therapy should follow. In addition to clarifying the subtype of leukemia, initial studies should evaluate the overall functional integrity of the major organ systems, including the cardiovascular, pulmonary, hepatic, and renal systems (Table 132-4). Factors that have prognostic significance, either for achieving CR or for predicting the duration of CR, should also be assessed before initiating treatment, including cytogenetics and molecular markers (see above). Leukemic cells should be obtained from all patients and cryopreserved for future use as new tests and therapeutics become available. All patients should be evaluated for infection.

Most patients are anemic and thrombocytopenic at presentation. Replacement of the appropriate blood components, if necessary, should begin promptly. Because qualitative platelet dysfunction or the presence of an infection may increase the likelihood of bleeding, evidence of hemorrhage justifies the immediate use of platelet transfusion, even if the platelet count is only moderately decreased.

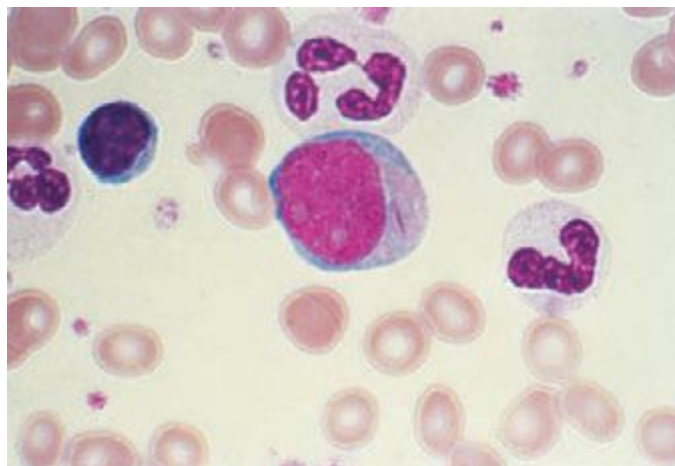
About 50% of patients have a mild to moderate elevation of serum uric acid at presentation. Only 10% have marked elevations, but renal precipitation of uric acid and the nephropathy that may result is a serious but uncommon complication. The initiation of chemotherapy may aggravate hyperuricemia, and patients are usually started immediately on allopurinol and hydration at diagnosis. Rasburicase (recombinant



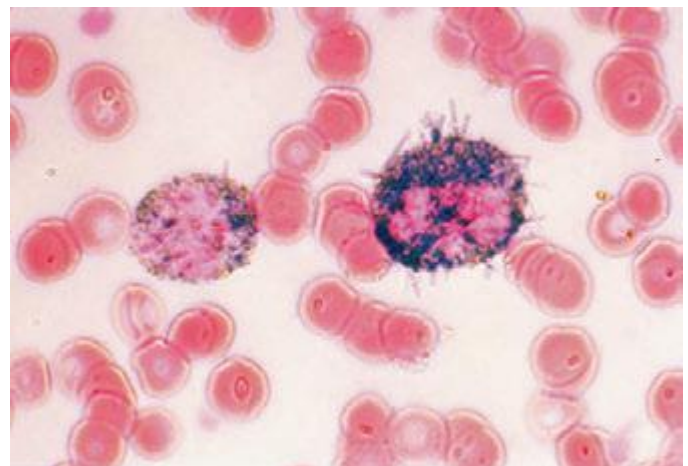
A



C



B



D

FIGURE 132-1 Morphology of acute myeloid leukemia (AML) cells. **A.** Uniform population of primitive myeloblasts with immature chromatin, nucleoli in some cells, and primary cytoplasmic granules. **B.** Leukemic myeloblast containing an Auer rod. **C.** Promyelocytic leukemia cells with prominent cytoplasmic primary granules. **D.** Peroxidase stain shows dark blue color characteristic of peroxidase in granules in AML.

uric oxidase) is also useful for treating uric acid nephropathy and often can normalize the serum uric acid level within hours with a single dose of treatment. The presence of high concentrations of lysozyme, a marker for monocytic differentiation, may be etiologic in renal tubular dysfunction, which could worsen other renal problems that arise during the initial phases of therapy.

TREATMENT ACUTE MYELOID LEUKEMIA

Treatment of the newly diagnosed patient with AML is usually divided into two phases, induction and postremission management (Fig. 132-2). The initial goal is to induce CR. Once CR is obtained, further therapy must be used to prolong survival and achieve cure. The initial induction treatment and subsequent postremission therapy are often chosen based on the patient's age. Intensifying therapy with traditional chemotherapy agents such as cytarabine and anthracyclines in younger patients (<60 years) appears to increase the cure rate of AML. In older patients, the benefit of intensive therapy is controversial; novel approaches for selecting patients predicted to be responsive to treatment and new therapies are being pursued.

INDUCTION CHEMOTHERAPY

The most commonly used CR induction regimens (for patients other than those with APL) consist of combination chemotherapy with cytarabine and an anthracycline (e.g., daunorubicin, idarubicin,

mitoxantrone). Cytarabine is a cell cycle S-phase-specific antimetabolite that becomes phosphorylated intracellularly to an active triphosphate form that interferes with DNA synthesis. Anthracyclines are DNA intercalators. Their primary mode of action is thought to be inhibition of topoisomerase II, leading to DNA breaks.

In younger adults (age <60 years), cytarabine is used either at standard dose (100–200 mg/m²) administered as a continuous intravenous infusion for 7 days or higher dose (2 g/m²) administered intravenously every 12 h for 6 days. With standard-dose cytarabine, anthracycline therapy generally consists of daunorubicin (60–90 mg/m²) or idarubicin (12 mg/m²) intravenously on days 1, 2, and 3 (the 7 and 3 regimen). Other agents can be added (i.e., cladribine) when 60 mg/m² of daunorubicin is used.

High-dose cytarabine-based regimens have also been shown to induce high CR rates. When given in high doses, higher intracellular levels of cytarabine may be achieved, thereby saturating the cytarabine-inactivating enzymes and increasing the intracellular levels of 1-β-D-arabinofuranylcytosine-triphosphate, the active metabolite incorporated into DNA. Thus, higher doses of cytarabine may increase the inhibition of DNA synthesis and thereby overcome resistance to standard-dose cytarabine. With high-dose cytarabine, daunorubicin 60 mg/m² or idarubicin 12 mg/m² is generally used.

The hematologic toxicity of high-dose cytarabine-based induction regimens has typically been greater than that associated with 7 and 3 regimens. Toxicity with high-dose cytarabine also includes pulmonary toxicity and significant and occasionally irreversible

TABLE 132-4 INITIAL DIAGNOSTIC EVALUATION AND MANAGEMENT OF ADULT PATIENTS WITH AML

History
Increasing fatigue or decreased exercise tolerance (anemia)
Excess bleeding or bleeding from unusual sites (DIC, thrombocytopenia)
Fevers or recurrent infections (neutropenia)
Headache, vision changes, nonfocal neurologic abnormalities (CNS leukemia or bleed)
Early satiety (splenomegaly)
Family history of AML (Fanconi, Bloom, or Kostmann syndromes or ataxia-telangiectasia)
History of cancer (exposure to alkylating agents, radiation, topoisomerase II inhibitors)
Occupational exposures (radiation, benzene, petroleum products, paint, smoking, pesticides)
Physical Examination
Performance status (prognostic factor)
Ecchymosis and oozing from IV sites (DIC, possible acute promyelocytic leukemia)
Fever and tachycardia (signs of infection)
Papilledema, retinal infiltrates, cranial nerve abnormalities (CNS leukemia)
Poor dentition, dental abscesses
Gum hypertrophy (leukemic infiltration, most common in monocytic leukemia)
Skin infiltration or nodules (leukemia infiltration, most common in monocytic leukemia)
Lymphadenopathy, splenomegaly, hepatomegaly
Back pain, lower extremity weakness [spinal granulocytic sarcoma, most likely in t(8;21) patients]
Laboratory and Radiologic Studies
CBC with manual differential cell count
Chemistry tests (electrolytes, creatinine, BUN, calcium, phosphorus, uric acid, hepatic enzymes, bilirubin, LDH, amylase, lipase)
Clotting studies (prothrombin time, partial thromboplastin time, fibrinogen, D-dimer)
Viral serologies (CMV, HSV-1, varicella-zoster)
RBC type and screen
HLA typing for potential allogeneic HSCT
Bone marrow aspirate and biopsy (morphology, cytogenetics, flow cytometry, molecular studies for <i>NPM1</i> and <i>CEBPA</i> mutations and <i>FLT3</i> -ITD)
Cryopreservation of viable leukemia cells
Myocardial function (echocardiogram or MUGA scan)
PA and lateral chest radiograph
Placement of central venous access device
Interventions for Specific Patients
Dental evaluation (for those with poor dentition)
Lumbar puncture (for those with symptoms of CNS involvement)
Screening spine MRI (for patients with back pain, lower extremity weakness, paresthesias)
Social work referral for patient and family psychosocial support
Counseling for All Patients
Provide patients with information regarding their disease, financial counseling, and support group contacts

Abbreviations: AML, acute myeloid leukemia; BUN, blood urea nitrogen; CBC, complete blood count; CMV, cytomegalovirus; CNS, central nervous system; DIC, disseminated intravascular coagulation; HLA, human leukocyte antigen; HSCT, hematopoietic stem cell transplantation; HSV, herpes simplex virus; IV, intravenous; LDH, lactate dehydrogenase; MRI, magnetic resonance imaging; MUGA, multigated acquisition; PA, posteroanterior; RBC, red blood (cell) count.

cerebellar toxicity. All patients treated with high-dose cytarabine must be closely monitored for cerebellar toxicity. Full cerebellar testing should be performed before each dose, and further high-dose cytarabine should be withheld if evidence of cerebellar toxicity develops. This toxicity occurs more commonly in patients with renal impairment and in those older than age 60 years. The increased toxicity observed with high-dose cytarabine has limited the use of this therapy in older AML patients.

Incorporation of novel and molecular targeting agents into these regimens is currently under investigation. For patients with *FLT3*-ITD AML, trials with tyrosine kinase inhibitors are ongoing. Patients with CBF AML may benefit from the combination of gemtuzumab ozogamicin, a monoclonal CD33 antibody linked to the cytotoxic agent calicheamicin, with induction and consolidation chemotherapies. This agent, initially approved for older patients with relapsed disease, has been withdrawn from the U.S. market at the request of the U.S. Food and Drug Administration due to concerns about the product's toxicity, including myelosuppression, infusion toxicity, and venoocclusive disease and the clinical benefit of the initially recommended higher doses. However, the aforementioned recent

results are encouraging and support the reintroduction of this agent into the therapeutic armamentarium for AML.

In older patients (age ≥ 60 years), the outcome is generally poor likely due to a higher induction treatment-related mortality rate and frequency of resistant disease, especially in patients with prior hematologic disorders (MDS or myeloproliferative syndromes) or who have received chemotherapy treatment for another malignancy or harbor cytogenetic and genetic abnormalities that adversely impact on clinical outcome. These patients should be considered for clinical trials. Alternatively, older patients can be also treated with the 7 and 3 regimen with standard-dose cytarabine and idarubicin (12 mg/m²), daunorubicin (45–90 mg/m²), or mitoxantrone (12 mg/m²). For patients older than 65 years, higher dose daunorubicin (90 mg/m²) has not shown benefit due to the increased toxicity and is not recommended. The combination of gemtuzumab ozogamicin with chemotherapy reduces the risk of relapse for patients age 50–70 years with previously untreated AML. Finally, older patients may be considered for single-agent therapies with clofarabine or hypomethylating agents (i.e., 5-azacitidine or decitabine). The latter are often used for patients unfit for more intensive therapies.

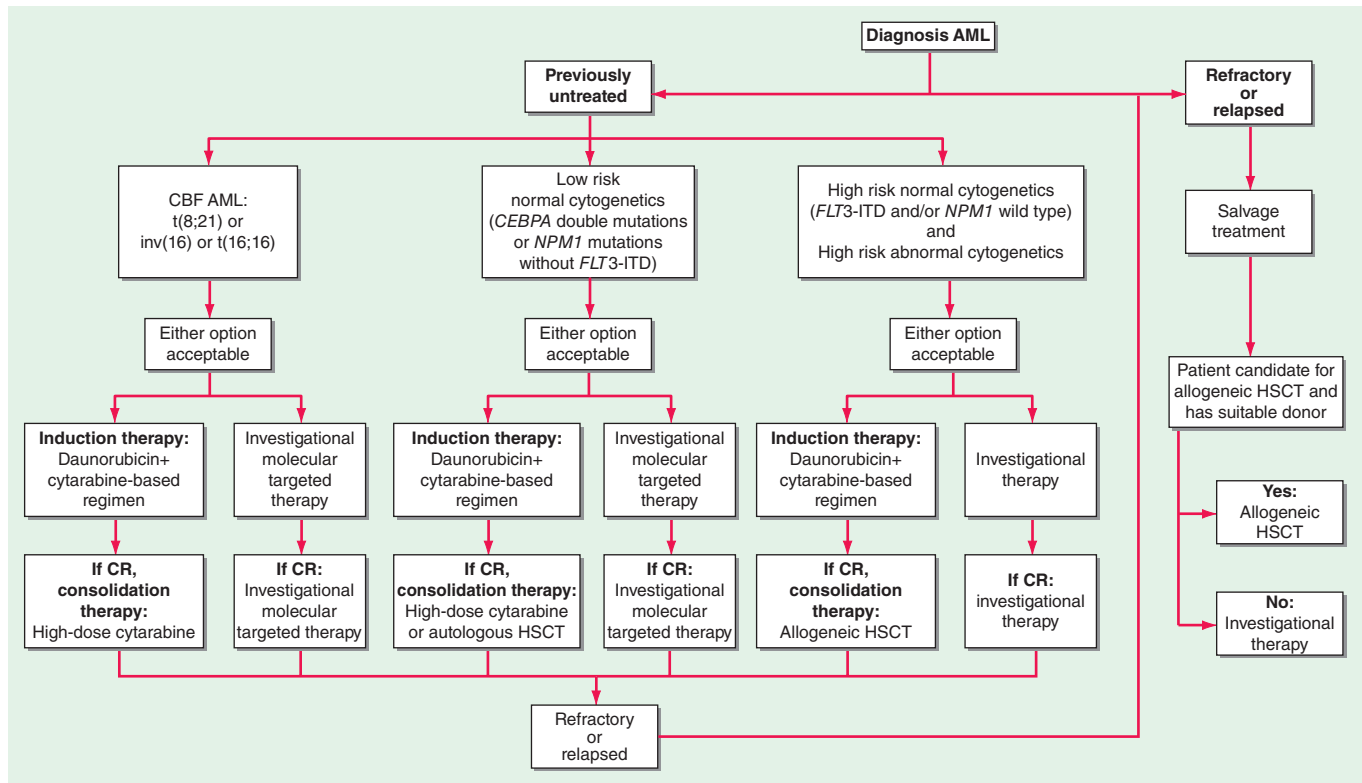


FIGURE 132-2 Flow chart for the therapy of newly diagnosed acute myeloid leukemia (AML). For all forms of AML except acute promyelocytic leukemia (APL), standard therapy includes a regimen based on a 7-day continuous infusion of cytarabine (100–200 mg/m² per day) and a 3-day course of daunorubicin (60–90 mg/m² per day) with or without additional drugs. Idarubicin (12–13 mg/m² per day) could be used in place of daunorubicin (not shown). Patients who achieve complete remission (CR) undergo postremission consolidation therapy, including sequential courses of high-dose cytarabine, autologous hematopoietic stem cell transplantation (HSCT), allogeneic HSCT, or novel therapies, based on their predicted risk of relapse (i.e., risk-stratified therapy). Patients with APL (see text for treatment) usually receive tretinoin and arsenic trioxide–based regimens with or without anthracycline-based chemotherapy and possibly maintenance with tretinoin. CBF, core binding factor; ITD, internal tandem duplication.

After one cycle of the 7 and 3 chemotherapy induction regimen, if persistence of leukemia is documented, the patient is usually re-treated with the same agents (cytarabine and the anthracycline) for 5 and 2 days, respectively. Our recommendation, however, is to consider changing therapy in this setting.

POSTREMISSION THERAPY

Induction of a durable first CR is critical to long-term disease-free survival in AML. However, without further therapy, virtually all patients experience relapse. Thus, postremission therapy is designed to eradicate residual leukemic cells to prevent relapse and prolong survival. The type of postremission therapy in AML is often based on age and cytogenetic and molecular risk.

For younger patients, most studies include intensive chemotherapy and allogeneic or autologous hematopoietic stem cell transplantation (HSCT). In the postremission setting, high-dose cytarabine for three to four cycles is more effective than standard-dose cytarabine. The Cancer and Leukemia Group B (CALGB), for example, compared the duration of CR in patients randomly assigned after remission to four cycles of high (3 g/m², every 12 h on days 1, 3, and 5), intermediate (400 mg/m² for 5 days by continuous infusion), or standard (100 mg/m² per day for 5 days by continuous infusion) doses of cytarabine. A dose-response effect for cytarabine in patients with AML who were age ≤60 years was demonstrated. High-dose cytarabine significantly prolonged CR and increased the fraction cured in patients with favorable [t(8;21) and inv(16)] and normal cytogenetics, but it had no significant effect on patients with other abnormal karyotypes. As discussed, high-dose cytarabine has increased toxicity in older patients. Therefore, in this age group, for patients without CBF AML, exploration of attenuated chemotherapy regimens has been pursued. However, because the outcome of older patients is poor, allogeneic

HSCT, when feasible, should be strongly considered. Postremission therapy is also a setting for introduction of new agents (Table 132-5).

Autologous HSCT preceded by one to two cycles of high-dose cytarabine is also an option for intensive consolidation therapy. Autologous HSCT has been generally applied to AML patients in the context of a clinical trial or when the risk of repetitive intensive chemotherapy represents a higher risk than the autologous HSCT (e.g., in patients with severe platelet alloimmunization) or when other factors including patient age, comorbid conditions, and fertility are considered.

Allogeneic HSCT is used in patients age <70–75 years with a human leukocyte antigen (HLA)-compatible donor who have high-risk cytogenetics. Selected high-risk patients are also considered for alternative donor transplants (e.g., mismatched unrelated, haploidentical related, and unrelated umbilical cord donors). In patients with CN-AML and high-risk molecular features such as FLT3-ITD, allogeneic HSCT is best applied in the context of clinical trials because the impact of aggressive therapy on outcome is unknown. For older patients, exploration of reduced-intensity allogeneic HSCT has been pursued.

Trials comparing intensive chemotherapy and autologous and allogeneic HSCT have shown improved duration of remission with allogeneic HSCT compared to autologous HSCT or chemotherapy alone. However, overall survival is generally not different; the improved disease control with allogeneic HSCT is erased by the increase in fatal toxicity. In fact, relapse following allogeneic HSCT occurs in only a small fraction of patients, but treatment-related toxicity is relatively high; complications include venoocclusive disease, graft-versus-host disease (GVHD), and infections. Autologous HSCT can be administered in young and older patients and uses the same preparative regimens. Patients subsequently receive their own stem cells collected while in remission. The toxicity is relatively low with

TABLE 132-5 SELECTED AGENTS UNDER STUDY FOR THE TREATMENT OF ACUTE MYELOID LEUKEMIA

Class of Drugs	Examples of Agents in Class
Inhibitors of Mutant proteins	
Tyrosine kinase inhibitors	Dasatinib, midostaurin, quizartinib, sorafenib
<i>IDH2</i> mutation inhibitor	AG-221
Epigenetic Targeting Compounds	
Demethylating agents	S110 (decitabine dinucleotide), oral azacitidine
Histone deacetylase inhibitors	Suberoylanilide hydroxamic acid (SAHA), MS275, LBH589
Inhibitors of Cell Proliferation	
Cell cycle inhibitors	Flavopiridol, CYC202 (R-roscovitine), SNS-032
Farnesyl transferase inhibitors	R115777, SCH66336
Aurora inhibitors	AZD1152, MLN-8237, AT9283
Inhibitors of Protein Synthesis and Degradation	
Aminopeptide inhibitors	Tosedostat
HSP-90 antagonists	17-Allylaminogeldanamycin (17-AAG), DMAG, or derivatives
Nedd8 activating enzyme (NAE) inhibitors	MLN4924
Cytotoxic Compounds	
Nucleoside analogues	Clofarabine, troxacitabine, elacytarabine, sapacitabine
Compounds with Immuno-Mediated Mechanisms	
Antibodies	CSL362 (anti-CD123), anti-CD33 (SGN33), anti-KIR
Immunomodulatory	Lenalidomide, interleukin 2, histamine dihydrochloride

autologous HSCT (5% mortality rate), but the relapse rate is higher than with allogeneic HSCT, due to the absence of the graft-versus-leukemia (GVL) effect seen with allogeneic HSCT and possible contamination of the autologous stem cells with residual tumor cells.

Prognostic factors may ultimately help to select the appropriate postremission therapy in patients in first CR. Our approach includes allogeneic HSCT in first CR for patients without favorable cytogenetics or genotype (e.g., patients who do not have *CEBPA* biallelic mutations or *NPM1* mutations without *FLT3*-ITD) and/or with other poor risk factors (e.g., an antecedent hematologic disorder or failure to attain remission with a single induction course). If a suitable HLA donor does not exist, investigational therapeutic approaches are considered. Indeed, postremission therapy is also a setting for introduction of new agents (Table 132-5). Because *FLT3*-ITD can be targeted with emerging novel inhibitors, patients with this molecular abnormality should be considered for clinical trials with these agents whenever possible.

Patients with the favorable CBF AML [i.e., t(8;21), inv(16), or t(16;16)] are treated with repetitive doses of high-dose cytarabine, which offers a high frequency of cure without the morbidity of transplant. Among AML patients with t(8;21) and inv(16), those with *KIT* mutations, who have a worse prognosis, may be considered for novel investigational studies, including tyrosine kinase inhibitors. The inclusion of gemtuzumab ozogamicin in induction and consolidation chemotherapy-based treatment has been reported to be beneficial in this subset of patients.

For patients in morphologic CR, immunophenotyping to detect minute populations of blasts or sensitive molecular assays (e.g., reverse transcriptase polymerase chain reaction [RT-PCR]) to detect AML-associated molecular abnormalities (e.g., *NPM1* mutation, the CBF AML *RUNX1/RUNX1T1* and *CBFB/MYH11* transcripts, the APL *PML/RARA* transcript), and the less sensitive metaphase cytogenetics or interphase cytogenetics by fluorescence in situ hybridization (FISH) to detect AML-associated cytogenetic aberrations, can be performed to assess whether clinically meaningful minimal

residual disease (MRD) is present at sequential time points during or after treatment. Detection of MRD may be a reliable discriminator between patients who will continue in CR and those who are destined to experience disease recurrence and therefore require early therapeutic intervention before clinical relapse occurs. Although assessment of MRD in bone marrow and/or blood during CR is routinely used in the clinic to anticipate clinical relapse and initiate timely salvage treatment for APL patients, for other cytogenetic and molecular subtypes of AML, this is an area of current investigation.

SUPPORTIVE CARE

Measures geared to supporting patients through several weeks of neutropenia and thrombocytopenia are critical to the success of AML therapy. Patients with AML should be treated in centers expert in providing supportive measures. Multilumen right atrial catheters should be inserted as soon as patients with newly diagnosed AML have been stabilized. They should be used thereafter for administration of intravenous medications and transfusions, as well as for blood drawing.

Adequate and prompt blood bank support is critical to therapy of AML. Platelet transfusions should be given as needed to maintain a platelet count $\geq 10,000/\mu\text{L}$. The platelet count should be kept at higher levels in febrile patients and during episodes of active bleeding or DIC. Patients with poor posttransfusion platelet count increments may benefit from administration of platelets from HLA-matched donors. RBC transfusions should be administered to keep the hemoglobin level $>80 \text{ g/L}$ (8 g/dL) in the absence of active bleeding, DIC, or congestive heart failure, which require higher hemoglobin levels. Blood products leukodepleted by filtration should be used to avert or delay alloimmunization as well as febrile reactions. Blood products should also be irradiated to prevent transfusion-associated GVHD. Cytomegalovirus (CMV)-negative blood products should be used for CMV-seronegative patients who are potential candidates for allogeneic HSCT. Leukodepleted products are also effective for these patients if CMV-negative products are not available.

Neutropenia (neutrophils $<500/\mu\text{L}$ or $<1000/\mu\text{L}$ and predicted to decline to $<500/\mu\text{L}$ over the next 48 h) can be part of the initial presentation and/or a side effect of the chemotherapy treatment in AML patients. Thus, infectious complications remain the major cause of morbidity and death during induction and postremission chemotherapy for AML. Antibacterial (i.e., quinolones) and antifungal (i.e., posaconazole) prophylaxis in the absence of fever is likely to be beneficial. For patients who are herpes simplex virus or varicella-zoster seropositive, antiviral prophylaxis should be initiated (e.g., acyclovir, valacyclovir).

Fever develops in most patients with AML, but infections are documented in only half of febrile patients. Early initiation of empirical broad-spectrum antibacterial and antifungal antibiotics has significantly reduced the number of patients dying of infectious complications (Chap. 104). An antibiotic regimen adequate to treat gram-negative organisms should be instituted at the onset of fever in a neutropenic patient after clinical evaluation, including a detailed physical examination with inspection of the indwelling catheter exit site and a perirectal examination, as well as procurement of cultures and radiographs aimed at documenting the source of fever. Specific antibiotic regimens should be based on antibiotic sensitivity data obtained from the institution at which the patient is being treated. Acceptable regimens for empiric antibiotic therapy include monotherapy with imipenem-cilastatin, meropenem, piperacillin/tazobactam, or an extended-spectrum antipseudomonal cephalosporin (cefepime or ceftazidime). The combination of an aminoglycoside with an antipseudomonal penicillin (e.g., piperacillin) or an aminoglycoside in combination with an extended-spectrum antipseudomonal cephalosporin should be considered in complicated or resistant cases. Aminoglycosides should be avoided if possible in patients with renal insufficiency. Empirical vancomycin should be added in neutropenic patients with catheter-related infections, blood cultures positive for gram-positive bacteria before final identification and susceptibility testing, hypotension or shock,

686 or known colonization with penicillin/cephalosporin-resistant pneumococci or methicillin-resistant *Staphylococcus aureus*. In special situations where decreased susceptibility to vancomycin, vancomycin-resistant organisms, or vancomycin toxicity is documented, other options including linezolid, daptomycin, and quinupristin/dalfopristin need to be considered.

Caspofungin (or a similar echinocandin), voriconazole, or liposomal amphotericin B should be considered for antifungal treatment if fever persists for 4–7 days following initiation of empiric antibiotic therapy. Amphotericin B has long been used for antifungal therapy. Although liposomal formulations have improved the toxicity profile of this agent, its use has been limited to situations with high risk of or documented mold infections. Caspofungin has been approved for empiric antifungal treatment. Voriconazole has also been shown to be equivalent in efficacy and less toxic than amphotericin B. Antibacterial and antifungal antibiotics should be continued until patients are no longer neutropenic, regardless of whether a specific source has been found for the fever.

Recombinant hematopoietic growth factors have been incorporated into clinical trials in AML. These trials have been designed to lower the infection rate after chemotherapy. Both G-CSF and granulocyte-macrophage colony-stimulating factor (GM-CSF) have reduced the median time to neutrophil recovery. This accelerated rate of neutrophil recovery, however, has not generally translated into significant reductions in infection rates or shortened hospitalizations. In most randomized studies, both G-CSF and GM-CSF have failed to improve the CR rate, disease-free survival, or overall survival. Although receptors for both G-CSF and GM-CSF are present on AML blasts, therapeutic efficacy is neither enhanced nor inhibited by these agents. The use of growth factors as supportive care for AML patients is controversial. We favor their use in elderly patients with complicated courses, those receiving intensive postremission regimens, patients with uncontrolled infections, or those participating in clinical trials.

TREATMENT FOR REFRACTORY OR RELAPSED AML

With the 7 and 3 regimen, 65–75% of younger and 50–60% of older patients with primary AML achieve CR. Two-thirds achieve CR after a single course of therapy, and one-third require two courses. Of patients who do not achieve CR, approximately 50% have a drug-resistant leukemia, and 50% do not achieve CR because of fatal complications of bone marrow aplasia or impaired recovery of normal stem cells. Patients with refractory disease after induction should be considered for salvage treatments, preferentially on clinical trials, before receiving allogeneic HSCT usually administered in patients who achieve a disease-free status. Because these patients are usually not cured even if they achieve second CR with salvage chemotherapy, allogeneic HSCT is a necessary therapeutic step.

In patients who relapse after achieving CR, the length of first CR is predictive of response to salvage chemotherapy treatment; patients with longer first CR (>12 months) generally relapse with drug-sensitive disease and have a higher chance of attaining a CR, even with the same chemotherapeutic agents used for first remission induction. Whether initial CR was achieved with one or two courses of chemotherapy and the type of postremission therapy may also predict achievement of second CR. Similar to patients with refractory disease, patients with relapsed disease are rarely cured by the salvage chemotherapy treatments. Therefore, patients who eventually achieve a second CR and are eligible for allogeneic HSCT should be transplanted.

Because achievement of a second CR with routine salvage therapies is relatively uncommon, especially in patients who relapse rapidly after achievement of first CR (<12 months), these patients and those lacking HLA-compatible donors or who are not candidates for allogeneic HSCT should be considered for innovative approaches on clinical trials (Table 132-5). The discovery of novel gene mutations and mechanisms of leukemogenesis that might represent actionable therapeutic targets has prompted the development of new targeting agents. In addition to kinase inhibitors for *FLT3*- and

KIT-mutated AML, other compounds targeting the aberrant activity of mutant proteins (e.g., IDH2 inhibitors) or biologic mechanisms deregulating epigenetics (e.g., histone deacetylase and DNA methyltransferase inhibitors), cell proliferation (e.g., farnesyl transferase inhibitors), protein synthesis (e.g., aminopeptide inhibitors) and folding (e.g., heat shock protein inhibitors), and ubiquitination, or with novel cytotoxic mechanisms (e.g., clofarabine, sapacitabine), are being tested in clinical trials. Furthermore, approaches with antibodies targeting commonly expressed leukemia blasts (e.g., CD33) or leukemia initiating cells (e.g., CD123) and immunomodulatory agents (e.g., lenalidomide) are also under investigation. Once these compounds have demonstrated safety and activity as single agents, investigation of combinations with other molecular targeting compounds and/or chemotherapy should be pursued.

TREATMENT OF ACUTE PROMYELOCYTIC LEUKEMIA

APL is a highly curable subtype of AML, and approximately 85% of these patients achieve long-term survival with current approaches. APL has long been shown to be responsive to cytarabine and daunorubicin, but previously patients treated with these drugs alone frequently died from DIC induced by the release of granule components by the chemotherapy-treated leukemic cells. However, the prognosis of APL patients has changed dramatically from adverse to favorable with the introduction of tretinoin, an oral drug that induces the differentiation of leukemic cells bearing the t(15;17), where disruption of the *RARA* gene encoding a retinoid acid receptor occurs. Tretinoin decreases the frequency of DIC but produces another complication called the APL differentiation syndrome. Occurring within the first 3 weeks of treatment, it is characterized by fever, fluid retention, dyspnea, chest pain, pulmonary infiltrates, pleural and pericardial effusions, and hypoxemia. The syndrome is related to adhesion of differentiated neoplastic cells to the pulmonary vasculature endothelium. Glucocorticoids, chemotherapy, and/or supportive measures can be effective for management of the APL differentiation syndrome. Temporary discontinuation of tretinoin is necessary in cases of severe APL differentiation syndrome (i.e., patients developing renal failure or requiring admission to the intensive care unit due to respiratory distress). The mortality rate of this syndrome is about 10%.

Tretinoin (45 mg/m² per day orally until remission is documented) plus concurrent anthracycline-based (i.e., idarubicin or daunorubicin) chemotherapy appears to be among the most effective treatment for APL, leading to CR rates of 90–95%. The role of cytarabine in APL induction and consolidation is controversial. The addition of cytarabine, although not demonstrated to increase the CR rate, seemingly decreases the risk for relapse. Following achievement of CR, patients should receive at least two cycles of anthracycline-based chemotherapy.

Arsenic trioxide has significant antileukemic activity and is being explored as part of initial treatment in clinical trials of APL. In a randomized trial, arsenic trioxide improved outcome if used after achievement of CR and before consolidation therapy with anthracycline-based chemotherapy. Patients receiving arsenic trioxide are at risk of APL differentiation syndrome, especially when it is administered during induction or salvage treatment after disease relapse. In addition, arsenic trioxide may prolong the QT interval, increasing the risk of cardiac arrhythmias.

Given the progress made in APL resulting in high cure rates, in recent years the goal has been to identify patients with low risk of relapse (i.e., those presenting with a leukocyte count $\leq 10,000/\mu\text{L}$) where attempts are being made to decrease the amount of therapy administered and to identify patients at greatest risk of relapse (i.e., those presenting with a leukocyte count $\geq 10,000/\mu\text{L}$) where new approaches can be developed to increase cure. A study compared the gold standard (tretinoin plus chemotherapy) in newly diagnosed non-high-risk APL with a chemotherapy-free combination of tretinoin and arsenic trioxide. An equivalent outcome was demonstrated between the two arms, and the chemotherapy-free regimen will likely become a new standard for non-high-risk APL patients.

Combinations of tretinoin, arsenic trioxide, and/or chemotherapy and/or gemtuzumab ozogamicin have shown favorable responses in high-risk APL patients at diagnosis.

Assessment of residual disease by RT-PCR amplification of the t(15;17) chimeric gene product *PML-RARA* following the final cycle of chemotherapy is an important step in the management of APL patients. Disappearance of the signal is associated with long-term disease-free survival; its persistence documented by two consecutive tests performed 2 weeks apart invariably predicts relapse. Sequential monitoring of RT-PCR for *PML-RARA* is now considered standard for postremission monitoring of APL, especially in high-risk patients.

The benefit from maintenance therapy with tretinoin has been documented in some studies and not in others. Thus, the use of tretinoin depends on which regimen has been used for induction and consolidation treatment and the risk category of the patients, with those with high-risk disease seemingly benefiting the most from maintenance therapy.

Patients in molecular, cytogenetic, or clinical relapse should be salvaged with arsenic trioxide with or without tretinoin; it produces meaningful responses in up to 85% of patients and can be followed by autologous or, less frequently, especially if RT-PCR positive for *PML-RARA*, allogeneic HSCT.

133 Chronic Myeloid Leukemia

Hagop Kantarjian, Jorge Cortes

Chronic myeloid leukemia (CML) is a clonal hematopoietic stem cell disorder. The disease is driven by the *BCR-ABL1* chimeric gene product, a constitutively active tyrosine kinase, resulting from a reciprocal balanced translocation between the long arms of chromosomes 9 and 22, t(9;22) (q34;q11.2), cytogenetically detected as the Philadelphia chromosome (Ph) (Fig. 133-1). Untreated, the course of CML may be biphasic or triphasic, with an early indolent or chronic phase, followed often by an accelerated phase and a terminal blastic phase. Before the era of selective *BCR-ABL1* tyrosine kinase inhibitors (TKIs), the median survival in CML was 3–7 years, and the 10-year survival rate was 30% or less. Introduced into CML therapy in 2000, TKIs have revolutionized the treatment, natural history, and prognosis of CML. Today, the estimated 10-year survival rate with imatinib mesylate, the first *BCR-ABL1* TKI approved, is 85%. Allogeneic stem cell transplantation (SCT), a curative but risky treatment approach, is now offered as second- or third-line therapy after failure of TKIs.

INCIDENCE AND EPIDEMIOLOGY

CML accounts for 15% of all cases of leukemia. There is a slight male preponderance (male:female ratio 1.6:1). The median age at diagnosis is 55–65 years. It is uncommon in children; only 3% of patients with CML are younger than 20 years. CML incidence increases slowly with age, with a steeper increase after the age of 40–50 years. The annual incidence of CML is 1.5 cases per 100,000 individuals. In the United States, this translates into 4500–5000 new cases per year. The incidence of CML has not changed over several decades. By extrapolation, the worldwide annual incidence of CML is about 100,000 cases. With a median survival of 6 years before 2000, the disease prevalence in the United States was 20,000–30,000 cases. With TKI therapy, the annual mortality has been reduced from 10–20% to about 2%. Therefore, the prevalence of CML in the United States is expected to continue to increase (about 80,000 in 2013) and reach a plateau of approximately 180,000 cases around 2030. The worldwide prevalence will depend on the treatment penetration of TKIs and their effect on reduction of worldwide annual mortality. Ideally, with full TKI treatment penetration, the worldwide prevalence should plateau at 35 times the incidence, or around 3 million patients.

ETIOLOGY

There are no familial associations in CML. The risk of developing CML is not increased in monozygotic twins or in relatives of patients. No etiologic agents are incriminated, and no associations exist with exposures to benzene or other toxins, fertilizers, insecticides, or viruses. CML is not a frequent secondary leukemia following therapy of other cancers with alkylating agents and/or radiation. Exposure to ionizing radiation (e.g., nuclear accidents, radiation treatment for ankylosing spondylitis or cervical cancer) has increased the risk of CML, which peaks at 5–10 years after exposure and is dose-related. The median time to development of CML among atomic bomb survivors was 6.3 years. Following the Chernobyl accident, the incidence of CML did not increase, suggesting that only large doses of radiation can cause CML. Because of adequate protection, the risk of CML is not increased in individuals working in the nuclear industry or among radiologists in recent times.

PATHOPHYSIOLOGY

The t(9;22) (q34;q11.2) is present in more than 90% of classical CML cases. It results from a balanced reciprocal translocation between the long arms of chromosomes 9 and 22. It is present in hematopoietic cells (myeloid, erythroid, megakaryocytes, and monocytes; less often mature B lymphocytes; rarely mature T lymphocytes, but not stromal cells), but not in other cells in the human body. As a result of the translocation, DNA sequences from the cellular oncogene *ABL1* are translocated next to the major breakpoint cluster region (*BCR*) gene on chromosome 22, generating a hybrid oncogene, *BCR-ABL1*. This fusion gene codes for a novel oncoprotein of molecular weight 210 kDa, referred to as p210^{BCR-ABL1} (Fig. 133-1B). This *BCR-ABL1* oncoprotein exhibits constitutive kinase activity that leads to excessive proliferation and reduced apoptosis of CML cells, endowing them with a growth advantage over their normal counterparts. Over time, normal hematopoiesis is suppressed, but normal stem cells can persist and may reemerge following effective therapy, for example with TKIs. In Ph-positive acute lymphocytic leukemia (ALL) and in rare cases of CML, the breakpoint in *BCR* is more centromeric, in a region called the minor *BCR* region (*mBCR*). As a result, a shorter sequence of *BCR* is fused to *ABL1*, with a consequent smaller *BCR-ABL1* oncoprotein, p190^{BCR-ABL1}. When occurring in Ph-positive CML, this translocation may predict for a worse outcome. A third rare breakpoint in *BCR* occurs telomeric to the major *BCR* region and is called *micro-BCR* (μ -*BCR*). It juxtaposes a larger fragment of the *BCR* gene to *ABL1* and produces a larger p230^{BCR-ABL1} oncoprotein, which is associated with a more indolent CML course.

The constitutive activation of *BCR-ABL1* results in autophosphorylation and activation of multiple downstream pathways that modify gene transcription, apoptosis, skeletal organization, and degradation of inhibitory proteins. These transduction pathways may involve RAS, mitogen-activated protein (MAP) kinases, signal transducers and activators of transcription (STAT), phosphatidylinositol-3-kinase (PI3k), MYC, and others. These interactions are mostly mediated through tyrosine phosphorylation and require binding of *BCR-ABL1* to adapter proteins such as GRB-2, CRK, CRK-like (CRK-L) protein, and Src homology containing proteins (SHC). *BCR-ABL1* TKIs bind to the *BCR-ABL1* kinase domain (KD), preventing the activation of transformation pathways and inhibiting downstream signaling. As a result, proliferation of CML cells is inhibited and apoptosis induced, leading to the reemergence of normal hematopoiesis. A plethora of signaling pathways have been implicated in *BCR-ABL1*-mediated cellular transformation. The emerging picture is a complex and redundant transformation network. An additional layer of complexity is related to differences in signal transduction between CML differentiated cells and early progenitors. Beta-catenin, Wnt1, Foxo3a, transforming growth factor β , interleukin-6, PP2A, SIRT1, and others have been implicated in CML stem cell survival.

Experimental models have established the causal relationship between the Ph-related *BCR-ABL1* molecular events and the development of CML. In animal models, expression of *BCR-ABL1* in normal hematopoietic cells produced CML-like disorders or lymphoid leukemia, demonstrating the leukemogenic potential of *BCR-ABL1* as a single oncogenic abnormality.

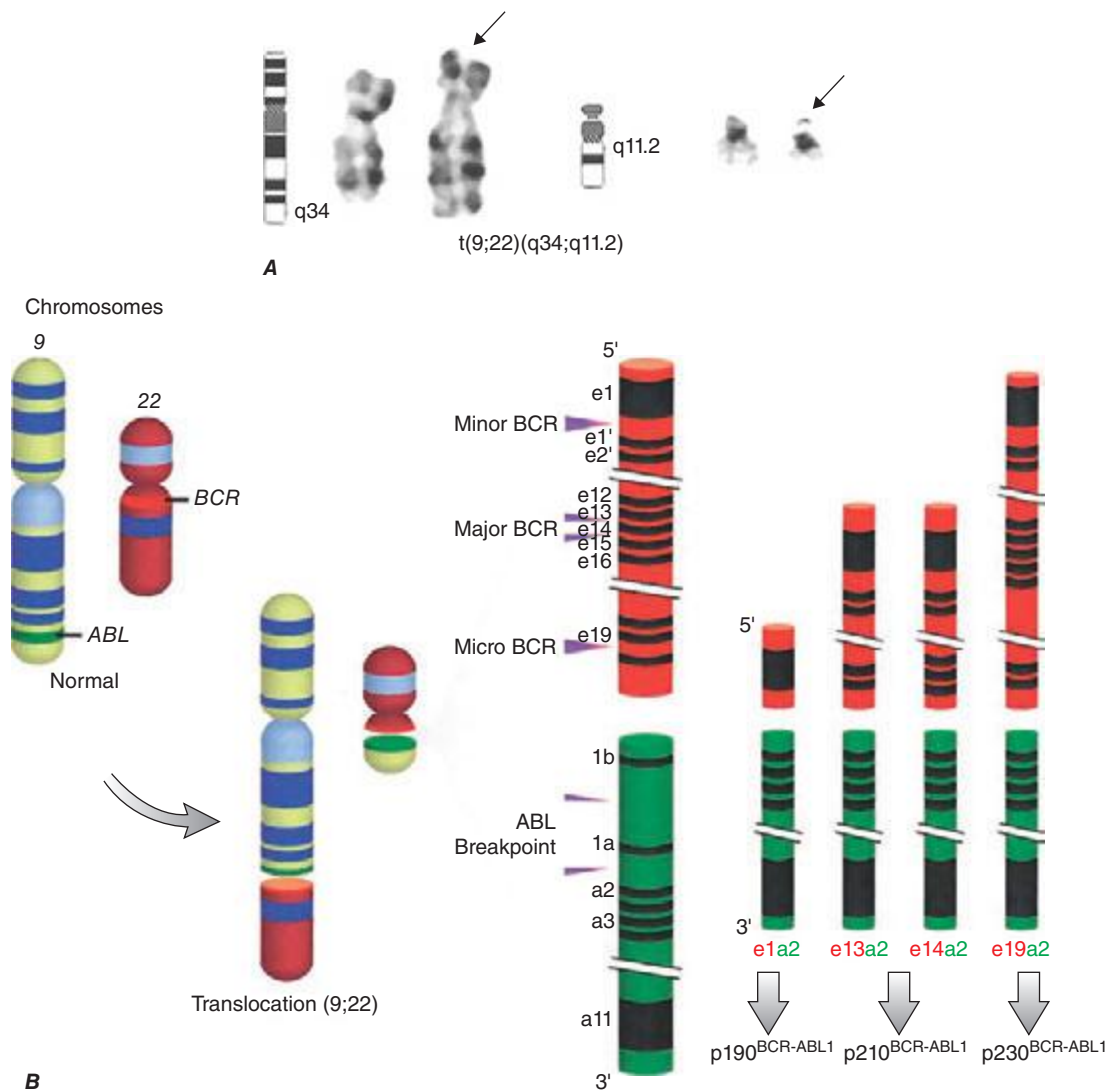


FIGURE 133-1 **A.** The Philadelphia (Ph) chromosome cytogenetic abnormality. **B.** Breakpoints in the long arms of chromosome 9 (*ABL* locus) and chromosome 22 (*BCR* regions) result in three different *BCR-ABL1* oncoprotein messages, p210^{BCR-ABL1} (most common message in chronic myeloid leukemia [CML]), p190^{BCR-ABL1} (present in two-thirds of patients with Ph-positive acute lymphocytic leukemia; rare in CML), and p230^{BCR-ABL1} (rare in CML and associated with an indolent course). (© 2013 The University of Texas MD Anderson Cancer Center.)

The cause of the *BCR-ABL1* molecular rearrangement is unknown. Molecular techniques that detect *BCR-ABL1* at a level of 1 in 10⁸ identify this molecular abnormality in the blood of up to 25% of normal adults and 5% of infants, but 0% of cord blood samples. This suggests that *BCR-ABL1* is not sufficient to cause overt CML in the overwhelming majority of individuals in whom it occurs. Because CML develops in only 1.5 of 100,000 individuals annually, it is evident that additional molecular events or poor immune recognition of the rearranged cells are needed to cause overt CML.

CML is defined by the presence of *BCR-ABL1* abnormality in a patient with a myeloproliferative neoplasm. In some patients with a typical morphologic picture of CML, the Ph abnormality is not detectable by standard cytogenetic analysis, but fluorescence in situ hybridization (FISH) and molecular studies (polymerase chain reaction [PCR]) detect *BCR-ABL1*. These patients have a course similar to Ph-positive CML and respond to TKI therapy. Many of the remaining patients have atypical morphologic or clinical features and belong to other diagnostic groups, such as atypical CML or chronic myelomonocytic leukemia. These individuals do not respond to TKI therapy and have a poor prognosis with a median survival of about 2–3 years. Detection of mutations in the granulocyte colony-stimulating factor receptor (*CSF3R*) in chronic neutrophilic leukemia and in some cases of atypical CML and of mutations in *SETBP1* in atypical CML confirmed that they are distinct entities.

The mechanisms associated with the transition of CML from a chronic to accelerated-blastic phase are poorly understood. They are often associated with characteristic chromosomal abnormalities such as a double Ph, trisomy 8, isochromosome 17 or deletion of 17p (loss of *TP53*), 20q-, and others. Molecular events associated with transformation include mutations in *TP53*, retinoblastoma 1 (*RB1*), myeloid transcription factors like Runx1, and cell cycle regulators like p16. A plethora of other mutations or functional abnormalities have been implicated in blastic transformation, but no unifying theme has emerged other than that *BCR-ABL1* itself induces genetic instability that leads to the acquisition of additional mutations and eventually to blastic transformation. In this frame of thinking, one critical effect of TKIs is their ability to stabilize the CML genome, leading to a much reduced transformation rate. In particular, the previously observed sudden blastic transformations (i.e., abrupt transformation to blastic phase in a patient who had been in cytogenetic response) have become uncommon, occurring rarely in younger patients in the first 1–2 years of TKI therapy (usually sudden lymphoid blastic transformations). Sudden transformations beyond the third year of TKI therapy are rare in patients who continue on TKI therapy. Moreover, initial experience suggests that the course of CML has become significantly more indolent, even without cytogenetic responses, in patients on TKI-based therapy compared to previous experience with hydroxyurea/busulfan.

Among patients developing resistance to TKIs, several resistance mechanisms have been observed. The most clinically relevant one is the development of different *ABL1* kinase domain mutations that prevent the binding of TKIs to the catalytic site (ATP binding site) of the kinase. More than 100 *BCR-ABL1* mutations have now been described, many of which confer relative or absolute resistance to imatinib. This has resulted in the development of second-generation TKIs (i.e., dasatinib, nilotinib, bosutinib) and of a third-generation TKI (ponatinib) with selective efficacy against T315I, a mutation of the gatekeeper residue of the kinase that causes resistance to all other TKIs.

CLINICAL PRESENTATION

The presenting signs and symptoms in CML depend on the availability of and access to health care procedures, including physical exams and screening tests. In the United States, because of the easy access to health care screening and physical exams, 50–60% of patients are diagnosed on routine blood tests and have minimal symptoms at presentation, such as fatigue. In geographic locations where access to health care is more limited, patients often present with high CML burden including splenomegaly, anemia, and related symptoms (abdominal pain, weight loss, fatigue), as well as a higher frequency of high-risk CML. Presenting findings in patients diagnosed in the United States are shown in [Table 133-1](#).

Symptoms Most patients with CML (90%) present in the indolent or chronic phase. Depending on the timing of diagnosis, patients are often asymptomatic (if the diagnosis is discovered during health care screening tests). Common symptoms, when present, are manifestations of anemia and splenomegaly. These may include fatigue, malaise, weight loss (if high leukemia burden), or early satiety and left upper quadrant pain or masses (from splenomegaly). Less common presenting findings include thrombotic or vasooclusive events (from severe leukocytosis or thrombocytosis). These include priapism, cardiovascular complications, myocardial infarction, venous thrombosis, visual disturbances, dyspnea and pulmonary insufficiency, drowsiness, loss of coordination, confusion, or cerebrovascular accidents. Bleeding diatheses findings include retinal hemorrhages, gastrointestinal bleeding,

and others. Patients who present with, or progress to, the accelerated or blastic phases have additional symptoms including unexplained fever, significant weight loss, severe fatigue, bone and joint aches, bleeding and thrombotic events, and infections.

Physical Findings Splenomegaly is the most common physical finding, occurring in 20–70% of patients depending on health care screening frequency. Other less common findings include hepatomegaly (10–20%), lymphadenopathy (5–10%), and extramedullary disease (skin or subcutaneous lesions). The latter indicates CML transformation if a biopsy confirms the presence of sheets of blasts. Other physical findings are manifestations of complications of high tumor burden described earlier (e.g., cardiovascular, cerebrovascular, bleeding). High basophil counts may be associated with histamine overproduction causing pruritus, diarrhea, flushing, and even gastrointestinal ulcers.

Hematologic and Marrow Findings In untreated CML, leukocytosis ranging from 10–500 × 10⁹/L is common. The peripheral blood differential shows left-shifted hematopoiesis with predominance of neutrophils and the presence of bands, myelocytes, metamyelocytes, promyelocytes, and blasts (usually ≤5%). Basophils and/or eosinophils are frequently increased. Thrombocytosis is common, but thrombocytopenia is rare and, when present, suggests a worse prognosis, disease acceleration, or an unrelated etiology. Anemia is present in one-third of patients. Cyclic oscillations of counts are noted in 25% of patients without treatment. Biochemical abnormalities include a low leukocyte alkaline phosphatase score and high levels of vitamin B₁₂, uric acid, lactic dehydrogenase, and lysozyme. The presence of unexplained and sustained leukocytosis, with or without splenomegaly, should lead to a marrow examination and cytogenetic analysis.

The bone marrow is hypercellular with marked myeloid hyperplasia and a high myeloid-to-erythroid ratio of 15–20:1. Marrow blasts are 5% or less; when higher, they carry a worse prognosis or represent acceleration (if they are ≥15%). Increased reticulin fibrosis (by Snook's silver stain) is common, with 30–40% of patients demonstrating grade 3–4 reticulin fibrosis. This was considered adverse in the pre-TKI era. With TKI therapy, reticulin fibrosis resolves in most patients and is not an indicator of poor prognosis. Collagen fibrosis (Wright-Giemsa stain) is rare at diagnosis. Disease progression with a “spent phase” of myelofibrosis (myelophthisis, or burnt-out marrow) was common with busulfan therapy (20–30%) but is rare with TKI therapy.

Cytogenetic and Molecular Findings The diagnosis of CML is straightforward and depends on documenting t(9;22)(q34;q11.2), which is found in 90% of cases. This is known as the Philadelphia-chromosome abnormality (discovered in Philadelphia) and was initially identified as a shortened chromosome, later identified to be chromosome 22 (22q-) (Fig. 133-1). Some patients may have complex translocations (variant Ph) involving three or more translocations that include chromosomes 9 and 22 and one or more other chromosomes. Others may have a “masked Ph,” involving translocations between chromosome 9 and a chromosome other than 22. The prognosis of these patients and their response to TKI therapy are similar to those in patients with Ph. About 5–10% of patients may have additional chromosomal abnormalities in the Ph-positive cells. These usually involve trisomy 8, a double Ph, isochromosome 17 or 17p deletion, 20q-, or others. This is referred to as clonal evolution and was historically a sign of adverse prognosis, particularly when trisomy 8, double Ph, or chromosome 17 abnormalities were noted.

Techniques such as FISH and PCR are now used to aid in the diagnosis of CML. They are more sensitive approaches to estimate the CML burden in patients on TKI therapy. They can be done on peripheral samples, and thus are less painful and more convenient. Patients with CML at diagnosis should have a FISH analysis to quantify the percentage of Ph-positive cells, if FISH is used to replace marrow cytogenetic analysis in monitoring response to therapy. FISH may not detect additional chromosomal abnormalities (clonal evolution); thus, a cytogenetic analysis is usually recommended at the time of diagnosis. The *BCR-ABL1* RNA message is usually one of two variants: e13a2 (formerly b2a2) and e14a2 (formerly b3a2). About 2–5% of patients

TABLE 133-1 PRESENTING SIGNS AND SYMPTOMS OF NEWLY DIAGNOSED PHILADELPHIA CHROMOSOME-POSITIVE CHRONIC MYELOID LEUKEMIA IN CHRONIC PHASE

Parameter	Percentage
Age ≥60 years (median)	18 (46)
Female gender	35–45
Splenomegaly	30
Hepatomegaly	5
Lymphadenopathy	5
Other extramedullary disease	2
Hemoglobin <10 g/dL	10–15
Platelets	
>450 × 10 ⁹ cells/L	30–35
<100 × 10 ⁹ cells/L	3–5
White blood cells ≥50 × 10 ⁹ cells/L	35–40
Marrow	
≥5% blasts	5
≥5% basophils	10–15
Peripheral blood	
≥3% blasts	8–10
≥7% basophils	10
Cytogenetic clonal evolution other than the Philadelphia chromosome	4–5
Sokal risk	
Low	60–65
Intermediate	25–30
High	10

690 may have other RNA fusion types (e.g., e1a2, e13a3, or e14a3). In these patients, the routine PCR primers may not amplify the *BCR-ABL1* transcripts, thus leading to false-negative results. Therefore, molecular studies at diagnosis are important to document the type and presence of *BCR-ABL1* transcripts to avoid erroneously “undetectable” *BCR-ABL1* transcripts on follow-up studies, with the misconception of a complete molecular response.

Both FISH and PCR studies can be falsely positive at low levels or falsely negative because of technical issues. Therefore, a diagnosis of CML must always rely on a marrow analysis with routine cytogenetics. The diagnostic bone marrow confirms the presence of the Ph chromosome, detects clonal evolution, i.e., chromosomal abnormalities in the Ph-positive cells (which may be prognostic), and also quantifies the percentage of marrow blasts and basophils. In 10% of patients, the percentage of marrow blasts and basophils can be significantly higher than in the peripheral blood, suggesting poorer prognosis or even disease transformation.

Monitoring patients on TKI therapy by cytogenetics, FISH, and molecular studies has become an important standard practice to assess response to therapy, emphasize compliance, evaluate possible treatment resistance, change TKI therapy, and order mutational analysis studies. It is thus important to recognize the comparability of these measures in monitoring response. A partial cytogenetic response is defined as the presence of 35% less Ph-positive metaphases by routine cytogenetic analysis. This is roughly equivalent to *BCR-ABL1* transcripts by the International Scale (IS) of 10% or less. A complete cytogenetic response refers to the absence of Ph-positive metaphases (0% Ph positivity). This is approximately equivalent to *BCR-ABL1* transcripts (IS) of 1% or less. A major molecular response refers to *BCR-ABL1* transcripts (IS) $\leq 0.1\%$, or roughly a 3-log or greater reduction of CML burden from baseline. A complete molecular response usually refers to *BCR-ABL1* transcripts (IS) $< 0.0032\%$ (undetectable by current techniques), roughly equivalent to a more than 4.5-log reduction of CML burden from baseline.

Findings in CML Transformation Progression of CML is usually associated with leukocytosis resistant to therapy, increasing anemia, fever and constitutional symptoms, and increased blasts and basophils in the peripheral blood or marrow. Criteria of accelerated-phase CML, historically associated with median survival of less than 1.5 years, include the presence of 15% or more peripheral blasts, 30% or more peripheral blasts plus promyelocytes, 20% or more peripheral basophils, cytogenetic clonal evolution (presence of chromosomal abnormalities in addition to Ph), and thrombocytopenia $< 100 \times 10^9/L$ (unrelated to

therapy). About 5–10% of patients present with de novo accelerated phase or blastic phase. The prognosis of de novo accelerated phase with TKI therapy has improved significantly, with an estimated 8-year survival rate of 75%. The median survival of accelerated phase evolving from chronic phase has also improved from a historical median survival of 18 months to an estimated 4-year survival rate of 70% on TKI therapy. Therefore, the criteria for accelerated-phase CML should be revisited because most have lost much of their prognostic significance. Blastic-phase CML is defined by the presence of 30% or more peripheral or marrow blasts or the presence of sheets of blasts in extramedullary disease (usually skin, soft tissues, or lytic bone lesions). Blastic-phase CML is commonly myeloid (60%) but can present uncommonly as erythroid, promyelocytic, monocytic, or megakaryocytic. Lymphoid blastic phase occurs in about 25% of patients. Lymphoblasts are terminal deoxynucleotide transferase positive and peroxidase negative (although occasionally with low positivity up to 3–5%) and express lymphoid markers (CD10, CD19, CD20, CD22). However, they also often express myeloid markers (50–80%), resulting in diagnostic confusion. This is important because, unlike other morphologic blastic phases, lymphoid blastic-phase CML is quite responsive to anti-ALL-type chemotherapy (e.g., hyper-CVAD [cyclophosphamide, vincristine, doxorubicin, and dexamethasone]) in combination with TKIs.

PROGNOSIS AND CML COURSE

Before the imatinib era, the annual mortality in CML was 10% in the first 2 years and 15–20% thereafter. The median survival time in CML was 3–7 years (with hydroxyurea-busulfan and interferon α). Without a curative option of allogeneic SCT, the course of CML was inexorable toward transformation to, and death from, accelerated or blastic phases. The disease stability was unpredictable, with some patients demonstrating sudden transformation to a blastic phase. With imatinib therapy, the annual mortality in CML has decreased to 2% in the first 12 years of observation. Half of the deaths are from factors other than CML, such as old age, accidents, suicides, other cancers, and other medical conditions (e.g., infections, surgical procedures). The estimated 8- to 10-year survival rate is now 85%, or 93% if only CML-related deaths are considered (Fig. 133-2). The course of CML has also become quite predictable. In the first 2 years of TKI therapy, rare sudden transformations are still noted (1–2%), usually lymphoid blastic transformations that respond to combinations of chemotherapy and TKIs followed by allogeneic SCT. These may be explained by the intrinsic mechanisms of sudden transformation already existing in the CML clones before the start of therapy that were not amenable to TKI inhibition, in particular imatinib. Second-generation TKIs (nilotinib,

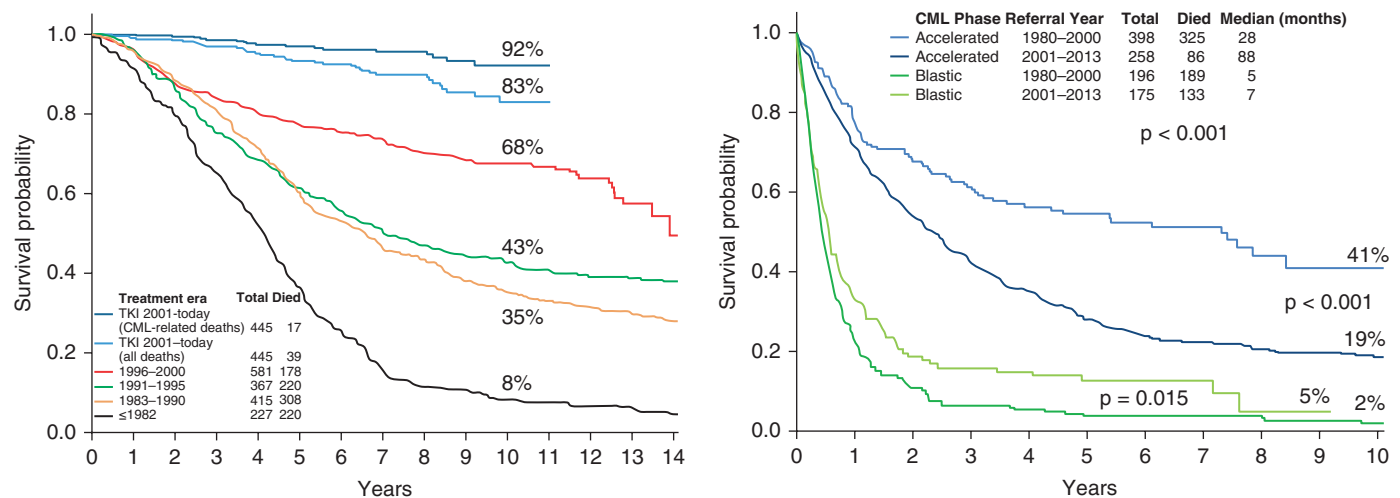


FIGURE 133-2 A. Survival in newly diagnosed chronic-phase chronic myeloid leukemia (CML) by era of therapy (M.D. Anderson Cancer Center experience from 1965 to present). Causes of non-CML deaths in 22 patients were other cancers (n = 7), postsurgical complications (n = 3), car accident (n = 2), suicide (n = 1), neurologic events (n = 3), cardiac (n = 3), pneumonia (n = 1), and unknown (n = 2). B. Survival in patients with accelerated- and blastic-phase CML referred to M.D. Anderson Cancer Center by era of therapy, demonstrating the significant survival benefit in the tyrosine kinase inhibitor (TKI) era in accelerated-phase CML but the modest benefit in blastic-phase CML. Referred cases included de novo and post-chronic-phase transformations.

dasatinib) used as frontline therapy have reduced the incidence of transformation in the first 2–3 years from 6–8% with imatinib to 2–4% with nilotinib or dasatinib. Disease transformation to accelerated or blastic phase is rare on continued TKI therapy, estimated at <1% annually in years 4–8 of follow-up on the original imatinib trials. Patients usually develop resistance in the form of cytogenetic relapse, followed by hematologic relapse and subsequent transformation, rather than the previously feared sudden transformations without the warning signals of cytogenetic-hematologic relapse.

Before the imatinib era, several pretreatment prognostic factors predicted for worse outcome in CML and have been incorporated into prognostic models and staging systems. These have included older age, significant splenomegaly, anemia, thrombocytopenia or thrombocytosis, high percentages of blasts and basophils (and/or eosinophils), marrow fibrosis, deletions in the long arm of chromosome 9, clonal evolution, and others. Different risk models and staging systems, derived from multivariate analyses, were proposed to define different risk groups. As with the introduction of cisplatin into testicular cancer therapy, the introduction of TKIs into CML therapy has nullified or lessened the prognostic impact of most of these prognostic factors and the significance of the CML models (e.g., Sokal, Hasford, European Treatment and Outcome Study [EUTOS]). Treatment-related prognostic factors have emerged as the most important prognostic factors in the era of imatinib therapy. Achievement of complete cytogenetic response has become the major therapeutic endpoint and is the only endpoint associated with improvement in survival. Achievement of a major molecular response is associated with decreased risk of events (relapse) and CML progression, may predict for differences in event-free survival (depending on the definition of an event) and for small differences in transformation rates, but has not been associated with survival prolongation. Among patients in complete cytogenetic response, survival is similar independent of whether they achieve a major molecular response or not. This may be due to the efficacy of salvage TKI therapies, which are and should be implemented at the first evidence of cytogenetic relapse. Achievement of complete molecular response (undetectable *BCR-ABL1* transcripts), particularly when durable (>2 years), may offer the possibility of durable molecular response (molecular cure rather than functional cure) in the context of investigational trials and may allow temporary therapy interruption in women eager to have babies. The lack of achievement of major or complete molecular responses should not be considered as “failure” of a particular TKI therapy and/or an indication to change the TKI or to consider allogeneic SCT.

Pretreatment prognostic factors and prognostic models have lost much of their clinical relevance to define prognosis and to select different therapies. However TKI-associated therapeutic responses have gained major clinical relevance and dictate appropriate and careful monitoring of patients to optimize their treatment.

TREATMENT CHRONIC MYELOID LEUKEMIA

The introduction of TKI therapy, first in the form of imatinib mesylate in 2001, has revolutionized the treatment and prognosis in CML. Before 2000, allogeneic SCT was frontline therapy, when available,

because of its potentially curative capacity. Otherwise, patients were offered interferon α therapy (approved for the treatment of CML in 1986), which had modest benefits (improving survival from a median of 3–4 years with hydroxyurea-busulfan to a median of 6–7 years), but also significant side effects. Other alternatives included hydroxyurea, busulfan, and other nonspecific chemotherapies. With TKI therapy, the estimated 10-year survival in CML is 85%. Since 2001, six agents have been approved by the U.S. Food and Drug Administration (FDA) for the treatment of CML. These include five oral *BCR-ABL1*-selective TKIs: imatinib (Gleevec), nilotinib (Tasigna), dasatinib (Sprycel), bosutinib (Bosulif), and ponatinib (Iclusig). Imatinib 400 mg orally daily, nilotinib 300 mg orally twice a day (on an empty stomach), and dasatinib 100 mg orally daily are approved for frontline therapy of CML. All three are also approved for salvage therapy (nilotinib 400 mg twice daily), in addition to bosutinib (500 mg daily) and ponatinib (45 mg daily). Imatinib, dasatinib (140 mg daily), bosutinib, and ponatinib are also approved for the treatment of CML transformation (accelerated and blastic phase), whereas nilotinib is only approved for chronic and accelerated phase. Dasatinib, nilotinib, and bosutinib are referred to as second-generation TKIs; ponatinib is referred to as a third-generation TKI. The sixth approved agent is omacetaxine (Synribo), a protein synthesis inhibitor with presumed more selective inhibition of the synthesis of the *BCR-ABL1* oncoprotein. It is approved for the treatment of chronic- and accelerated-phase CML after failure of two or more TKIs, at 1.25 mg/m² subcutaneously twice a day for 14 days for induction and for 7 days for consolidation-maintenance. Nilotinib is similar in structure to imatinib but 30 times more potent. Dasatinib and bosutinib are dual *SRC-ABL1* TKIs (dasatinib is reported to be 300 times more potent and bosutinib 30–50 times more potent than imatinib). Ponatinib is effective against wild-type and mutant *BCR-ABL1* clones. It is unique in being the only currently available *BCR-ABL1* TKI that is active against T315I, a gatekeeper mutant resistant to the other four TKIs (Table 133-2).

Imatinib, nilotinib, and dasatinib are all acceptable frontline therapies in CML. The long-term results of imatinib are very favorable. The 8-year follow-up results show a cumulative complete cytogenetic response rate (occurring at least once) of 83%, with 60–65% of patients being in complete cytogenetic response at 5-year follow-up. The estimated 8-year event-free survival rate is 81%, and the overall survival rate is 85%. Among patients continuing on imatinib, the annual rate of transformation to accelerated-blastic phase in years 4–8 is <1%. In two randomized studies, one comparing nilotinib 300 mg twice daily or 400 mg twice daily with imatinib (ENEST-nd) and the other comparing dasatinib 100 mg daily with imatinib (DASISION), the second-generation TKIs were associated with better outcomes in early surrogate endpoints, including higher rates of complete cytogenetic responses (85–87% vs 77–82%), major molecular responses (65–76% vs 46–63%), and undetectable *BCR-ABL1* transcripts (IS) (32–37% vs 15–30%), and lower rates of transformation to accelerated-blastic phase (2–4% vs 6%). However, neither study showed a survival benefit with second-generation TKIs (median follow-up times of 4–5 years). This may be because salvage

TABLE 133-2 MEDICAL THERAPEUTIC OPTIONS IN CHRONIC MYELOID LEUKEMIA

Agent (Brand Name)	Approved Indications	Dose Schedule	Notable Toxicities
Imatinib mesylate (Gleevec)	All phases	400 mg daily	See text
Dasatinib (Sprycel)	All phases	First-line: 100 mg daily Salvage: 140 mg daily	Myelosuppression; pleural and pericardial effusions; pulmonary hypertension
Nilotinib (Tasigna)	All phases except blastic phase	First-line: 300 mg twice daily Salvage: 400 mg twice daily	Diabetes; vasoocclusive disease; pancreatitis
Bosutinib (Bosulif)	All phases except frontline	500 mg daily	Diarrhea
Ponatinib (Iclusig)	All phases except frontline	45 mg daily (may consider lower starting doses in the future, e.g., 30 mg daily)	Skin rashes, pancreatitis; vasoocclusive disease (10–20%)
Omacetaxine mepesuccinate (Synribo)	Failure ≥ 2 tyrosine kinase inhibitors	1.25 mg/m ² subcutaneously twice daily for 14 days for induction; 7 days of maintenance every month	Myelosuppression

692 therapy with other TKIs (following close observation and treatment change at progression) provides highly effective salvage therapy that rebalances the negative effect of the relapse.

Salvage therapy in chronic phase with dasatinib, nilotinib, bosutinib, or ponatinib is associated with complete cytogenetic response rates of 30–60% of patients, depending on the salvage status (cytogenetic vs hematologic relapse), prior response to other TKIs, and the mutations at the time of relapse. Complete cytogenetic responses are generally durable, particularly in the absence of clonal evolution and mutations. Ponatinib is the only TKI active in the setting of T315I mutation, with complete cytogenetic response rates of 50–70%. The estimated 3- to 5-year survival rates with new TKIs as salvage are 70–80% (compared with <50% before their availability). For example, with dasatinib salvage after imatinib failure in chronic-phase CML, the major molecular response rates were 40–43%, the estimated 6-year survival rates were 74–83%, and progression-free survival rates were 40–51%. Thus, TKIs in the salvage setting have already reduced the annual mortality from the historical rate of 10–15% to ≤5%.

The goal of CML therapy is viewed differently in the context of research versus standard practice. In current practice, functional cure, defined as survival with CML similar to survival among normal individuals, is the current goal of therapy. CML is now considered an indolent disease, which, with appropriate TKI therapy, treatment compliance, careful monitoring, and early change to other TKIs as indicated, can be associated with close to normal survival. Therefore, in standard practice, achievement and maintenance of a complete cytogenetic response are the aims of therapy, because complete cytogenetic response is the only treatment-related factor associated with survival prolongation. Lack of achievement of a major molecular response (protects against events; associated with longer event-free survival) or of negative *BCR-ABL1* transcripts (offers the potential of TKI interruption on investigational studies) should not be considered indications to change TKI therapy or to consider allogeneic SCT. A general practice rule is to continue the particular TKI chosen at the most tolerable dose schedule not associated with grade 3–4 side effects or with bothersome chronic side effects, for as long as possible, until either cytogenetic relapse or the persistence of unacceptable side effects. These two factors (i.e., cytogenetic relapse and intolerable side effects as judged by the patient and treating physician) are the indicators of “failure” of a particular TKI therapy. Because of the increasing prevalence of CML (cost of TKI therapy) and the emerging long-term low rates of significant organ toxicities, the ultimate goal of CML therapy in the research setting is to achieve eradication of the disease (molecular cure) that is prolonged and durable, with recovery of nonneoplastic, nonclonal hematopoiesis off TKI therapy. The first step toward this aim is to obtain the highest rates of undetectable *BCR-ABL1* transcripts lasting for at least 2 or more years.

Recommendations provided by the National Comprehensive Cancer Network (NCCN) and by the European LeukemiaNet (ELN) discuss optimal/expected, suboptimal/warning, and failure response scenarios at different time points of TKI treatment duration. Unfortunately, they may have been misinterpreted in current practice, because oncologists often report that their aim of treatment is the achievement of major molecular response and disease eradication. Significantly, a substantial proportion of oncologists consider a change of TKI therapy in a patient in complete cytogenetic response if they note “loss of major molecular response” (increase of *BCR-ABL1* transcripts [IS] from <0.1% to >0.1%). This perception may be the result of confusion regarding the NCCN and ELN guidelines, which have been updated often as a result of maturing data and have multiple treatment endpoint considerations. Although such endpoints have been suggested by these recommendations as possible criteria for failure, it is important to emphasize that no randomized study has yet shown that a change of TKI treatment in patients with complete cytogenetic response because of a loss of major molecular response, versus changing at the time of cytogenetic relapse, has been shown to improve survival. This is likely because of the high efficacy of salvage TKI therapy at the time of cytogenetic relapse.

Side effects of TKIs are generally mild to moderate, although with long-term TKI therapy, they could affect the patient’s quality of life. Serious side effects occur in less than 5–10% of patients. With imatinib therapy, common mild to moderate side effects include fluid retention, weight gain, nausea, diarrhea, skin rashes, periorbital edema, bone or muscle aches, fatigue, and others (rates of 10–20%). In general, second-generation TKIs are associated with lower rates of these bothersome adverse events. However, dasatinib is associated with higher rates of myelosuppression (20–30%), particularly thrombocytopenia, and with pleural (10–25%) or pericardial effusions (≤5%). Nilotinib is associated with higher rates of hyperglycemia (10–20%), pruritus and skin rashes, and headaches. Nilotinib is also associated with rare events of pancreatitis (<5%). Bosutinib is associated with higher rates of early and self-limited gastrointestinal complications like diarrhea (50–70%). Ponatinib is associated with higher rates of skin rashes (10–15%), pancreatitis (5%), elevations of amylase/lipase (10%), and vasospastic/vasoocclusive events (10–20%). Nilotinib and dasatinib may cause prolongation of the QTc interval; therefore, they should be evaluated cautiously in patients with prolonged QTc interval on electrocardiogram (>470–480 ms), and drugs given for other medical conditions should have relatively smaller or no effects on QTc. These side effects can often be dose-dependent and are generally reversible with treatment interruptions and dose reductions. Dose reductions can be individualized. However, the lowest estimated effective doses of TKIs (from different studies and treatment practices) are imatinib 300 mg daily; nilotinib 200 mg twice daily; dasatinib 20 mg daily; bosutinib 300 mg daily; and ponatinib 15 mg daily.

With long-term follow-up, rare but clinically relevant serious toxicities are emerging. Renal dysfunction and renal failure (creatinine elevations >2–3 mg/dL) are observed in 2–3% of patients and reverse with TKI discontinuation and empirical use of other TKIs. Pulmonary hypertension has been reported with dasatinib (<1–2%) and should be considered in a patient with shortness of breath and a normal chest x-ray (echocardiogram with emphasis on measurement of pulmonary artery pressure). This may be reversible with dasatinib discontinuation and occasionally the use of sildenafil citrate. Systemic hypertension has been observed more often with ponatinib therapy, as well as other TKIs. Hyperglycemia and diabetes have been noted more frequently with nilotinib. Finally, mid- and small-vessel vasoocclusive and vasospastic events have been reported at low but significant rates with nilotinib and ponatinib and should be considered possibly TKI-related and represent indications to interrupt or reduce the dose of the TKI. These events include angina, coronary artery disease, myocardial infarction, peripheral arterial occlusive disease, transient ischemic attacks, cerebral vascular accidents, Raynaud’s phenomenon, and accelerated atherosclerosis. Although these events are uncommon (<5%), they are clinically significant for the patient’s long-term prognosis and occur at significantly higher rates than in the general population (5–20 times more often).

ALLOGENEIC STEM CELL TRANSPLANT

Allogeneic SCT, a curative modality in CML, is associated with long-term survival rates of 40–60% when implemented in the chronic phase. It is associated with early (1-year) mortality rates of 5–30%. Although the 5- to 10-year survival rates were reported to be around 50–60% (and considered as cure rates), about 10–15% of patients die in the subsequent 1–2 decades from subtle long-term complications of the transplant (rather than from CML relapse). These are related to chronic graft-versus-host disease (GVHD), organ dysfunction, development of second cancers, and hazard ratios for mortality higher than in the normal population. Other significant morbidities include infertility, chronic immune-mediated complications, cataracts, hip necrosis, and other morbidities affecting quality of life. The cure and early mortality rates in chronic-phase CML are also associated with several factors: patient age, duration of chronic phase, whether the donor is related or unrelated, degree of matching, preparative regimen, and others. In accelerated-phase

CML, the cure rates with allogeneic SCT are 20–40%, depending on the definition of acceleration. Patients with clonal evolution as the only criterion have cure rates of up to 40–50%. Patients undergoing allogeneic SCT in second chronic phase have cure rates of 40–50%. The cure rates with allogeneic SCT in blastic phase CML are \leq 15%. Post-allogeneic SCT strategies are now implemented in the setting of molecular or cytogenetic relapse or in hematologic relapse/transformation. These include the use of TKIs for prevention or treatment of relapse, donor lymphocyte infusions, and second allogeneic SCTs, among others. TKIs appear to be highly successful at reinducing cytogenetic/molecular remissions in the setting of cytogenetic or molecular relapse after allogeneic SCT.

Choice and Timing of Allogeneic SCT Allogeneic SCT was considered first-line CML therapy before 2000. The maturing positive experience with TKIs has now relegated its use to after first-line TKI failures. An important question is the optimal timing and sequence of TKIs and allogeneic SCT (whether allogeneic SCT should be used as second- or third-line therapy). Among patients who present with or evolve to blastic phase, combinations of chemotherapy and TKIs should be used to induce remission, followed by allogeneic SCT as soon as possible. The same applies to patients who evolve from chronic to accelerated phase. Patients with de novo accelerated-phase CML may do well with long-term TKI therapy (estimated 8-year survival rate 75%); the timing of allogeneic SCT depends on their optimal response to TKI (achievement of complete cytogenetic response). Among patients who relapse in chronic phase, the treatment sequence depends on several factors: (1) patient age and availability of appropriate donors; (2) risk of allogeneic SCT; (3) presence or absence of clonal evolution and mutations; (4) patient's prior history and comorbidities; and (5) patient and physician preferences (Table 133-3). Patients with T315I mutations at relapse should be offered ponatinib and considered for allogeneic SCT (because of the short follow-up with ponatinib). Patients with mutations involving Y253H, E255K/V, and F359V/C/I respond better to dasatinib or bosutinib. Patients with mutations involving V299L, T315A, and F317L/F//C respond better to nilotinib. Comorbidities such as diabetes, hypertension, pulmonary hypertension, chronic lung disease, cardiac conditions, and pancreatitis may influence the choice for or against a particular TKI. Patients with clonal evolution, unfavorable mutations, or lack of major/complete cytogenetic

response within 1 year of salvage TKI therapy have short remission durations and should consider allogeneic SCT as more urgent in the setting of salvage. Patients without clonal evolution or mutations at relapse and who achieve a complete cytogenetic response with TKI salvage, have long-lasting complete remissions and may delay the option of allogeneic SCT to third-line therapy. Finally, older patients (age 65–70 years or older) and those with high risk of mortality with allogeneic SCT may forgo this curative option for several years of disease control in chronic phase with or without cytogenetic response (Table 133-3). Historically, before the availability of TKIs, patients without cytogenetic response on interferon α or hydroxyurea had expected short median survival times (2–3 years) with expected rapid disease transformation. The maturing experience with TKIs suggests a different course, whereby patients may remain in chronic phase on TKI-based therapies (combinations including hydroxyurea, cytarabine, decitabine, and others), with or without cytogenetic response, for many years. Table 133-3 summarizes a general guidance to the choice of TKIs versus allogeneic SCT.

MONITORING THERAPY IN CML

Achievement of complete cytogenetic response by 12 months of imatinib therapy and its persistence later, the only consistent prognostic factor associated with survival, is now the main therapeutic endpoint in CML. Failure to achieve a complete cytogenetic response by 12 months or occurrence of later cytogenetic or hematologic relapse is considered as treatment failure and an indication to change therapy. Because salvage therapy with other TKIs reestablishes good outcome, it is important to ensure patient compliance to continued TKI therapy and change therapy at the first sign of cytogenetic relapse. Patients on frontline imatinib therapy should be closely monitored until documentation of complete cytogenetic response, at which time they can be monitored every 6 months with peripheral blood FISH and PCR studies (to check for concordance of results), or more frequently if there are concerns about changes in *BCR-ABL1* transcripts (e.g., every 3 months). Monitoring by molecular studies only is reasonable in patients who are in major molecular response. Cytogenetic relapse on imatinib is an indication of treatment failure and need to change TKI therapy. Mutational analysis in this instance helps in the selection of the next TKI and identifies mutations in 30–50% of patients. Mutational studies in patients in complete cytogenetic response (in whom there may be concerns of increasing *BCR-ABL1* transcripts) identify mutations in \leq 5% and are therefore not indicated. Earlier response has been identified as a prognostic factor for long-term outcome, including achievement of partial cytogenetic response (*BCR-ABL1* transcripts \leq 10%) by 3–6 months of therapy. Failure to achieve such a response on imatinib therapy has been associated with significantly worse survival in some studies (particularly when second-generation TKIs were not readily available as salvage therapy), but not in others (when they were).

The use of second-generation TKIs (nilotinib, dasatinib) as front-line therapy changed the monitoring approach slightly. Patients are expected to achieve complete cytogenetic response by 3–6 months of therapy. Failure to do so is associated with worse event-free survival, transformation rates, and survival. However, the 3- to 5-year estimated survival among such patients is still high, around 80–90%, which is better than what would be anticipated if such patients were offered allogeneic SCT at that time. Thus, this adverse response to therapy is considered a warning signal, but it is not known whether changing therapy to other TKIs at that time would improve longer term outcome.

TREATMENT OF ACCELERATED AND BLASTIC PHASES

Patients in accelerated or blastic phase may receive therapy with TKIs, preferably second- or third-generation TKIs (dasatinib, nilotinib, bosutinib, ponatinib), alone or in combination with chemotherapy, to reduce the CML burden, before undergoing allogeneic SCT. Response rates with single-agent TKIs range from 30 to 50% in accelerated phase and from 20 to 30% in blastic phase. Cytogenetic

TABLE 133-3 GENERAL SUGGESTIONS REGARDING THE USE OF TYROSINE KINASE INHIBITORS (TKIS) AND ALLOGENEIC STEM CELL TRANSPLANTATION (SCT) IN CHRONIC MYELOID LEUKEMIA (CML)

CML Phase	Use of TKI	Consideration of Allogeneic SCT
Accelerated or blastic	Interim therapy to achieve minimal CML burden	As soon as possible (exception: de novo accelerated phase)
Imatinib failure in chronic phase; T315I mutation	Ponatinib to achieve minimal CML burden	Depends on longer term follow-up results of ponatinib efficacy
Imatinib failure in chronic phase; no clonal evolution, no mutations, good initial response	Second-line kinase inhibitors long-term	Third-line after second-line TKI failures
Imatinib failure in chronic phase; clonal evolution or mutations, or no cytogenetic response to second-line TKI	Interim therapy to achieve minimal CML burden	Second-line
Older patients (\geq 65–70 years) after imatinib failure in chronic phase	Salvage TKIs as longer-term therapy	May forgo allogeneic SCT in favor of good quality of life and survival in chronic phase

Note: Mutations involving Y253H, E255K/V, or F359V/C/I: prefer dasatinib or bosutinib. Mutations involving V299L, T315A, or F317L/F//C: prefer nilotinib.

694 responses, particularly complete cytogenetic responses, are uncommon (10–30%) and transient in blastic phase. Studies of TKIs in combination with chemotherapy are ongoing; the general experience suggests that combined TKI-chemotherapy strategies increase the response rates and their durability and improve survival. In CML lymphoid blastic phase, the combination of anti-ALL chemotherapy with TKIs results in complete response rates of 60–70% and median survival times of 2–3 years (compared with historical response rates of 40–50% and median survival times of 12–18 months). This allows many patients to undergo allogeneic SCT in a state of minimal CML burden or secondary chronic phase, which are associated with higher cure rates. In CML nonlymphoid blastic phase, anti-AML chemotherapy combined with TKIs results in CR rates of 30–50% and median survival times of 9–12 months (compared with historical response rates of 20–30% and median survival times of 3–5 months). In accelerated phase, response to single TKIs is significant in conditions where “softer” accelerated phase criteria are considered (e.g., clonal evolution alone, thrombocytosis alone, significant splenomegaly or resistance to hydroxyurea, but without evidence of high blast and basophil percentages). In accelerated phase, combinations usually include TKIs with low-intensity chemotherapy such as low-dose cytarabine, low-dose idarubicin, decitabine, interferon α , hydroxyurea, or others.

OTHER TREATMENTS AND SPECIAL THERAPEUTIC CONSIDERATIONS

Interferon α Interferon α was a standard of care before 2000. Today, it is considered in combination with TKIs (an investigational approach), sometimes after CML failure on TKIs, occasionally in patients during pregnancy, or as part of investigational strategies with TKIs to eradicate residual molecular disease.

Chemotherapeutic Agents Hydroxyurea and busulfan were commonly used chemotherapeutic agents in the past. Hydroxyurea remains a safe and effective agent (at daily doses of 0.5–10 g) to reduce initial CML burden, as a temporary measure in between definitive therapies, or in combination with TKIs to sustain complete hematologic or cytogenetic responses. Busulfan is often used in allogeneic SCT preparative regimens. Because of its side effects (delayed myelosuppression, Addison-like disease, pulmonary and cardiac fibrosis, myelofibrosis), it is now only rarely used in the chronic management of CML. Low-dose cytarabine, decitabine, anthracyclines, 6-mercaptopurine, 6-thioguanine, thiopeta, anagrelide, and other agents are useful in different CML settings to control the disease burden.

Others Splenectomy is occasionally considered to alleviate symptoms of massive splenomegaly and/or hypersplenism. Splenic irradiation is rarely used, if at all, because of the postirradiation adhesions and complications. Leukapheresis is rarely used in patients presenting with extreme leukocytosis and leukostatic complications. Single doses of high-dose cytarabine or high doses of hydroxyurea, with tumor lysis management, may be as effective and less cumbersome.

Special Considerations Women with CML who become pregnant should discontinue TKI therapy immediately. Among 125 babies delivered to women with CML who discontinued TKI therapy as soon as the pregnancy was known, three babies were born with ocular, skeletal, and renal malformations, suggesting the uncommon teratogenicity of imatinib. There are no or little data with other TKIs. Control of CML during pregnancy can be managed with leukapheresis for severe symptomatic leukocytosis in the first trimester and with hydroxyurea subsequently until delivery. There are case reports of successful pregnancies and deliveries of normal babies with interferon α therapy and registry studies in essential thrombocytosis of its safety, but interferon α can be antiangiogenic and may increase the risk of spontaneous abortions.

Patients on TKI therapy may develop chromosomal abnormalities in the Ph-negative cells. These may involve loss of chromosome Y, trisomy 8, 20q-, chromosome 5 or 7 abnormalities, and others. Most chromosomal abnormalities disappear spontaneously on follow-up

and may be indicative of the genetic instability of the hematopoietic stem cells that predispose the patient to develop CML in the first place. Rarely, abnormalities involving chromosomes 5 or 7 may be truly clonal and evolve into myelodysplastic syndrome or acute myeloid leukemia. This is thought to be part of the natural course of patients in whom CML was suppressed and who live long enough to develop other hematologic malignancies.

GLOBAL ASPECTS OF CHRONIC MYELOID LEUKEMIA

Routine physical exams and blood tests in the United States and advanced countries result in early detection of CML in most patients. About 50–70% of patients with CML are diagnosed accidentally, and high-risk CML as defined by prognostic models (e.g., Sokal risk groups) is found in only 10–20% of patients. This is not the same situation in emerging nations (e.g., India, China, African countries, the Middle East), where most patients are diagnosed following evaluation for symptoms and many present with high tumor burdens, such as massive splenomegaly, and advanced phases of CML (high-risk CML documented in 30–50%). Therefore, the prognosis of such patients on TKI therapy may be worse than the published experience.

The high cost of TKI therapies (annual costs of \$90,000–\$140,000 in the United States; lower but variable in the rest of the world) makes the general affordability of such treatments difficult. Although TKI treatment penetration is high in nations where cost of therapy is not an issue (e.g., Sweden, European Union), it may be less so in other nations, even in advanced ones like the United States, where out-of-pocket expenses may be prohibitive to a subset of patients (perhaps 10–20%). Based on the sales of imatinib worldwide and charity free drug supplies, it is estimated that less than 30% of patients are treated with imatinib (or other TKIs) consistently. Although the estimated 10-year survival in CML is 85% in single-institution studies (e.g., M.D. Anderson Cancer Center), in national studies in countries with TKI affordability (Sweden) (Figs. 133-2 and 133-3) or in company-sponsored studies (where all patients have access to TKIs throughout their care), the estimated 10-year survival worldwide, even 12 years after the introduction of TKI therapies, is likely to be less than 50%. The Surveillance, Epidemiology, and End Results (SEER) data from

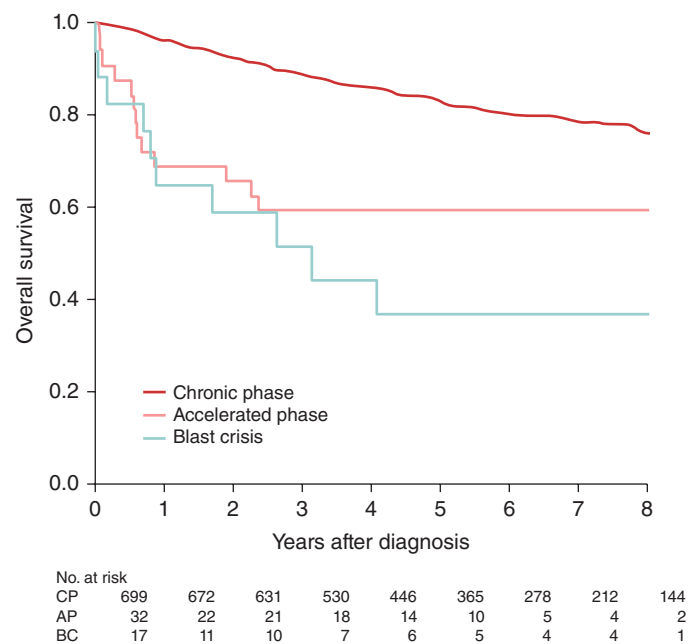


FIGURE 133-3 Survival in chronic (CP), accelerated (AP), and blastic crisis (BC) phases of chronic myeloid leukemia (CML) in the population-based Swedish national registry study. The accelerated- and blastic-phase cases are de novo presentations. The favorable outcome with de novo blastic phase may be due to use of 20% blasts or more to define blastic phase. (With permission from Dr. Martin Hoglund, Swedish CML Registry, 2013.)

the United States report an estimated 5-year survival rate of 60% in the era of TKIs.

The current high cost of TKI therapies poses two additional considerations. The first are the treatment pathways and guidelines in nations where TKIs may not be affordable by patients or the health care system. In these conditions, there are trends of pathways advocating frontline allogeneic SCT (a one-time cost of \$30,000–\$50,000) despite the associated mortality and morbidities. The second is the choice of frontline TKI therapy once imatinib becomes available in generic forms (hopefully at much lower annual prices, e.g., \$2,000–\$10,000). This will depend on the maturing data in randomized studies of second-generation TKIs versus imatinib in relation to important long-term outcome endpoints, particularly survival, but also event-free survival and transformation-free survival.

134 Malignancies of Lymphoid Cells

Dan L. Longo

Malignancies of lymphoid cells range from the most indolent to the most aggressive human malignancies. These cancers arise from cells of the immune system at different stages of differentiation, resulting in a wide range of morphologic, immunologic, and clinical findings. Insights on the normal immune system have allowed a better understanding of these sometimes confusing disorders.

Some malignancies of lymphoid cells almost always present as leukemia (i.e., primary involvement of bone marrow and blood), while others almost always present as lymphomas (i.e., solid tumors of the immune system). However, other malignancies of lymphoid cells can present as either leukemia or lymphoma. In addition, the clinical pattern can change over the course of the illness. This change is more often seen in a patient who seems to have a lymphoma and then develops the manifestations of leukemia over the course of the illness.

BIOLOGY OF LYMPHOID MALIGNANCIES: CONCEPTS OF THE WORLD HEALTH ORGANIZATION CLASSIFICATION OF LYMPHOID MALIGNANCIES

The classification of lymphoid cancers evolved steadily throughout the twentieth century. The distinction between leukemia and lymphoma was made early, and separate classification systems were developed for each. Leukemias were first divided into acute and chronic subtypes based on average survival. Chronic leukemias were easily subdivided into those of lymphoid or myeloid origin based on morphologic characteristics. However, a spectrum of diseases that were formerly all called *chronic lymphoid leukemia* has become apparent (Table 134-1). The acute leukemias were usually malignancies of blast cells with few identifying characteristics. When cytochemical stains became available, it was possible to divide these objectively into myeloid malignancies and acute leukemias of lymphoid cells. Acute leukemias of lymphoid cells have been subdivided based on morphologic characteristics by the French-American-British (FAB) group (Table 134-2). Using this system, lymphoid malignancies of small

TABLE 134-1 LYMPHOID DISORDERS THAT CAN PRESENT AS “CHRONIC LEUKEMIA” AND BE CONFUSED WITH TYPICAL B-CELL CHRONIC LYMPHOID LEUKEMIA

Follicular lymphoma	Prolymphocytic leukemia (B cell or T cell)
Splenic marginal zone lymphoma	Lymphoplasmacytic lymphoma
Nodal marginal zone lymphoma	Sézary's syndrome
Mantle cell lymphoma	Smoldering adult T-cell leukemia/lymphoma
Hairy cell leukemia	

TABLE 134-2 CLASSIFICATION OF ACUTE LYMPHOID LEUKEMIA (ALL)

Immunologic Subtype	% of Cases	FAB Subtype	Cytogenetic Abnormalities
Pre-B ALL	75	L1, L2	t(9;22), t(4;11), t(1;19)
T-cell ALL	20	L1, L2	14q11 or 7q34
B-cell ALL	5	L3	t(8;14), t(8;22), t(2;8)

Abbreviation: FAB, French-American-British classification.

uniform blasts (e.g., typical childhood acute lymphoblastic leukemia) were called L1, lymphoid malignancies with larger and more variable size cells were called L2, and lymphoid malignancies of uniform cells with basophilic and sometimes vacuolated cytoplasm were called L3 (e.g., typical Burkitt's lymphoma cells). Acute leukemias of lymphoid cells have also been subdivided based on immunologic (i.e., T cell vs B cell) and cytogenetic abnormalities (Table 134-2). Major cytogenetic subgroups include the t(9;22) (e.g., Philadelphia chromosome-positive acute lymphoblastic leukemia) and the t(8;14) found in the L3 or Burkitt's leukemia.

Non-Hodgkin's lymphomas were separated from Hodgkin's lymphoma by recognition of the Sternberg-Reed cells early in the twentieth century. The histologic classification for non-Hodgkin's lymphomas has been one of the most contentious issues in oncology. Imperfect morphologic systems were supplanted by imperfect immunologic systems, and poor reproducibility of diagnosis has hampered progress. In 1999, the World Health Organization (WHO) classification of lymphoid malignancies was devised through a process of consensus development among international leaders in hematopathology and clinical oncology. The WHO classification takes into account morphologic, clinical, immunologic, and genetic information and attempts to divide non-Hodgkin's lymphomas and other lymphoid malignancies into clinical/pathologic entities that have clinical and therapeutic relevance. This system is presented in Table 134-3. This system is clinically relevant and has a higher degree of diagnostic accuracy than those used previously. The possibilities for subdividing lymphoid malignancies are extensive. However, Table 134-3 presents in bold those malignancies that occur in at least 1% of patients. Specific lymphoma subtypes will be dealt with in more detail below.

Lymphomas occurring in fewer than 1% of patients with lymphoproliferative diseases are discussed in Chap. 135e, and lymphomas associated with HIV infection are discussed in Chap. 226.

GENERAL ASPECTS OF LYMPHOID MALIGNANCIES

ETIOLOGY AND EPIDEMIOLOGY

The relative frequency of the various lymphoid malignancies is shown in Fig. 134-1. Chronic lymphoid leukemia (CLL) is the most prevalent form of leukemia in Western countries. It occurs most frequently in older adults and is exceedingly rare in children. In 2014, 15,720 new cases were diagnosed in the United States, but because of the prolonged survival associated with this disorder, the total prevalence is many times higher. CLL is more common in men than in women and more common in whites than in blacks. This is an uncommon malignancy in Asia. The etiologic factors for typical CLL are unknown.

In contrast to CLL, acute lymphoid leukemias (ALLs) are predominantly cancers of children and young adults. The L3 or Burkitt's leukemia occurring in children in developing countries seems to be associated with infection by the Epstein-Barr virus (EBV) in infancy. However, the explanation for the etiology of more common subtypes of ALL is much less certain. Childhood ALL occurs more often in higher socioeconomic subgroups. Children with trisomy 21 (Down's syndrome) have an increased risk for childhood ALL as well as acute myeloid leukemia (AML). Exposure to high-energy radiation in early childhood increases the risk of developing T-cell ALL.

The etiology of ALL in adults is also uncertain. ALL is unusual in middle-aged adults but increases in incidence in the elderly. However, AML is still much more common in older patients. Environmental exposures, including certain industrial exposures, exposure to agricultural chemicals, and smoking, might increase the risk of developing

TABLE 134-3 WHO CLASSIFICATION OF LYMPHOID MALIGNANCIES

B Cell	T Cell	Hodgkin's Lymphoma
Precursor B-cell neoplasm Precursor B lymphoblastic leukemia/lymphoma (precursor B-cell acute lymphoblastic leukemia) includes subtypes with recurrent genetic abnormalities	Precursor T-cell neoplasm Precursor T lymphoblastic lymphoma/leukemia (precursor T cell acute lymphoblastic leukemia)	Nodular lymphocyte-predominant Hodgkin's lymphoma
Mature (peripheral) B-cell neoplasms B-cell chronic lymphocytic leukemia/small lymphocytic lymphoma B-cell prolymphocytic leukemia Lymphoplasmacytic lymphoma (Waldenström's macroglobulinemia) Splenic marginal zone B-cell lymphoma (± villous lymphocytes) Hairy cell leukemia Plasma cell myeloma/plasmacytoma Extranodal marginal zone B-cell lymphoma of MALT type Mantle cell lymphoma Follicular lymphoma Nodal marginal zone B-cell lymphoma (± monocytoid B cells) Diffuse large B-cell lymphoma (including subtypes) Burkitt's lymphoma/Burkitt's cell leukemia Primary mediastinal large B-cell lymphoma Plasmablastic lymphoma Primary effusion lymphoma Large B-cell lymphoma arising in HHV-8+ multicentric Castleman's disease Intravascular large B-cell lymphoma ALK+ large B-cell lymphoma	Mature (peripheral) T-cell neoplasms T-cell prolymphocytic leukemia T-cell granular lymphocytic leukemia Aggressive NK cell leukemia Adult T-cell lymphoma/leukemia (HTLV-1+) Extranodal NK/T-cell lymphoma, nasal type Enteropathy-type T-cell lymphoma Hepatosplenic γδ T-cell lymphoma Subcutaneous panniculitis-like T-cell lymphoma Mycosis fungoides/Sézary's syndrome Anaplastic large cell lymphoma, primary cutaneous type Peripheral T-cell lymphoma, not otherwise specified (NOS) Angioimmunoblastic T-cell lymphoma Anaplastic large cell lymphoma, ALK+ Primary cutaneous γδ T-cell lymphoma	Classical Hodgkin's lymphoma Nodular sclerosis classical Hodgkin's lymphoma Lymphocyte-rich classical Hodgkin's lymphoma Mixed-cellularity classical Hodgkin's lymphoma Lymphocyte-depletion classical Hodgkin's lymphoma

Note: Malignancies in bold occur in at least 1% of patients.

Abbreviations: HHV, human herpesvirus; HTLV, human T-cell lymphotropic virus; MALT, mucosa-associated lymphoid tissue; NK, natural killer; WHO, World Health Organization.

Source: Adapted from SH Swerdlow et al: *WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues*, 4th ed. World Health Organization, 2008.

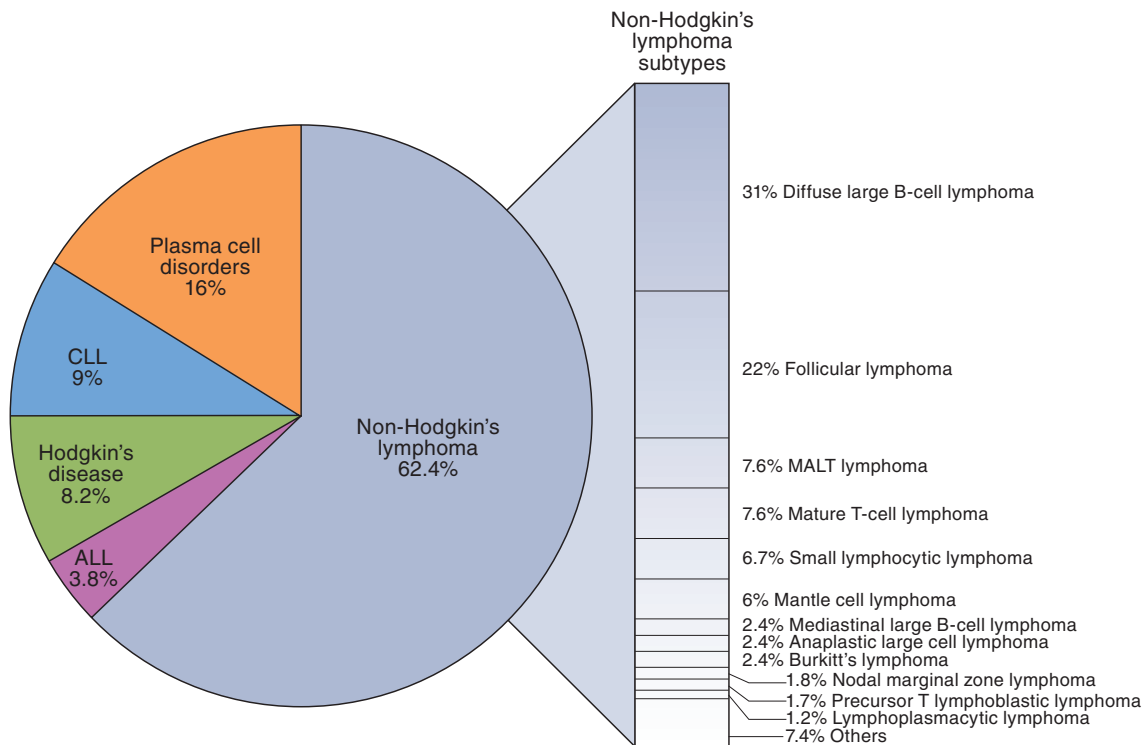


FIGURE 134-1 Relative frequency of lymphoid malignancies. ALL, acute lymphoid leukemia; CLL, chronic lymphoid leukemia; MALT, mucosa-associated lymphoid tissue.

ALL as an adult. ALL was diagnosed in 6020 persons and AML in 18,860 persons in the United States in 2014.

The preponderance of evidence suggests that Hodgkin's lymphoma is of B-cell origin. The incidence of Hodgkin's lymphoma appears fairly stable, with 9190 new cases diagnosed in 2014 in the United States. Hodgkin's lymphoma is more common in whites than in blacks and more common in males than in females. A bimodal distribution of age at diagnosis has been observed, with one peak incidence occurring in patients in their twenties and the other in those in their eighties. Some of the late age peak may be attributed to confusion among entities with similar appearance such as anaplastic large cell lymphoma and T-cell-rich B-cell lymphoma. Patients in the younger age groups diagnosed in the United States largely have the nodular sclerosing subtype of Hodgkin's lymphoma. Elderly patients, patients infected with HIV, and patients in Third World countries more commonly have mixed-cellularity Hodgkin's lymphoma or lymphocyte-depleted Hodgkin's lymphoma. Infection by HIV is a risk factor for developing Hodgkin's lymphoma. In addition, an association between infection by EBV and Hodgkin's lymphoma has been suggested. A monoclonal or oligoclonal proliferation of EBV-infected cells in 20–40% of the patients with Hodgkin's lymphoma has led to proposals for this virus having an etiologic role in Hodgkin's lymphoma. However, the matter is not settled definitively.

For unknown reasons, non-Hodgkin's lymphomas increased in frequency in the United States at the rate of 4% per year and increased 2–8% per year globally between 1950 and the late 1990s. The rate of increase in the past few years seems to be decreasing. About 70,800 new cases of non-Hodgkin's lymphoma were diagnosed in the United States in 2014 and nearly 360,000 cases worldwide. Non-Hodgkin's lymphomas are more frequent in the elderly and more frequent in men. Patients with both primary and secondary immunodeficiency states are predisposed to developing non-Hodgkin's lymphomas. These include patients with HIV infection; patients who have undergone organ transplantation; and patients with inherited immune deficiencies, the sicca syndrome, and rheumatoid arthritis.

The incidence of non-Hodgkin's lymphomas and the patterns of expression of the various subtypes differ geographically. T-cell lymphomas are more common in Asia than in Western countries, while certain subtypes of B-cell lymphomas such as follicular lymphoma are more common in Western countries. A specific subtype of non-Hodgkin's lymphoma known as the angiocentric nasal T/natural killer (NK) cell lymphoma has a striking geographic occurrence, being most frequent in Southern Asia and parts of Latin America. Another subtype of non-Hodgkin's lymphoma associated with infection by human T-cell lymphotropic virus (HTLV) 1 is seen particularly in southern Japan and the Caribbean (Chap. 225e).

A number of environmental factors have been implicated in the occurrence of non-Hodgkin's lymphoma, including infectious agents, chemical exposures, and medical treatments. Several studies have demonstrated an association between exposure to agricultural chemicals and an increased incidence of non-Hodgkin's lymphoma. Patients treated for Hodgkin's lymphoma can develop non-Hodgkin's lymphoma; it is unclear whether this is a consequence of the Hodgkin's lymphoma or its treatment. However, a number of non-Hodgkin's lymphomas are associated with infectious agents (Table 134-4). HTLV-1 infects T cells and leads directly to the development of adult T-cell lymphoma in a small percentage of infected patients. The cumulative lifetime risk of developing lymphoma in an infected patient is 2.5%. The virus is transmitted by infected lymphocytes ingested by nursing babies of infected mothers, bloodborne transmission, or sexually. The median age of patients with adult T-cell lymphoma is ~56 years, emphasizing the long latency. HTLV-1 is also the cause of tropical spastic paraparesis—a neurologic disorder that occurs somewhat more frequently than lymphoma and with shorter latency and usually from transfusion-transmitted virus (Chap. 225e).

EBV is associated with the development of Burkitt's lymphoma in Central Africa and the occurrence of aggressive non-Hodgkin's lymphomas in immunosuppressed patients in Western countries. The majority of primary central nervous system (CNS) lymphomas

TABLE 134-4 INFECTIOUS AGENTS ASSOCIATED WITH THE DEVELOPMENT OF LYMPHOID MALIGNANCIES

Infectious Agent	Lymphoid Malignancy
Epstein-Barr virus	Burkitt's lymphoma Post-organ transplant lymphoma Primary CNS diffuse large B-cell lymphoma Hodgkin's lymphoma Extranodal NK/T-cell lymphoma, nasal type
HTLV-1	Adult T-cell leukemia/lymphoma
HIV	Diffuse large B-cell lymphoma Burkitt's lymphoma
Hepatitis C virus	Lymphoplasmacytic lymphoma
<i>Helicobacter pylori</i>	Gastric MALT lymphoma
Human herpesvirus 8	Primary effusion lymphoma Multicentric Castlemans disease

Abbreviations: CNS, central nervous system; HIV, human immunodeficiency virus; HTLV, human T-cell lymphotropic virus; MALT, mucosa-associated lymphoid tissue; NK, natural killer.

are associated with EBV. EBV infection is strongly associated with the occurrence of extranodal nasal T/NK cell lymphomas in Asia and South America. Infection with HIV predisposes to the development of aggressive, B-cell non-Hodgkin's lymphoma. This may be through overexpression of interleukin 6 by infected macrophages. Infection of the stomach by the bacterium *Helicobacter pylori* induces the development of gastric MALT (mucosa-associated lymphoid tissue) lymphomas. This association is supported by evidence that patients treated with antibiotics to eradicate *H. pylori* have regression of their MALT lymphoma. The bacterium does not transform lymphocytes to produce the lymphoma; instead, a vigorous immune response is made to the bacterium, and the chronic antigenic stimulation leads to the neoplasia. MALT lymphomas of the skin may be related to *Borrelia* sp. infections, those of the eyes to *Chlamydomphila psittaci*, and those of the small intestine to *Campylobacter jejuni*.

Chronic hepatitis C virus infection has been associated with the development of lymphoplasmacytic lymphoma. Human herpesvirus 8 is associated with primary effusion lymphoma in HIV-infected persons and multicentric Castlemans disease, a diffuse lymphadenopathy associated with systemic symptoms of fever, malaise, and weight loss.

In addition to infectious agents, a number of other diseases or exposures may predispose to developing lymphoma (Table 134-5).

IMMUNOLOGY

All lymphoid cells are derived from a common hematopoietic progenitor that gives rise to lymphoid, myeloid, erythroid, monocyte, and megakaryocyte lineages. Through the ordered and sequential activation of a series of transcription factors, the cell first becomes committed to the lymphoid lineage and then gives rise to B and T cells. About 75% of all lymphoid leukemias and 90% of all lymphomas are of B-cell origin. A cell becomes committed to B-cell development when it begins to rearrange its immunoglobulin genes. The sequence of cellular changes,

TABLE 134-5 DISEASES OR EXPOSURES ASSOCIATED WITH INCREASED RISK OF DEVELOPMENT OF MALIGNANT LYMPHOMA

Inherited immunodeficiency disease	Autoimmune disease
Klinefelter's syndrome	Sjögren's syndrome
Chédiak-Higashi syndrome	Celiac sprue
Ataxia-telangiectasia syndrome	Rheumatoid arthritis and systemic lupus erythematosus
Wiskott-Aldrich syndrome	Chemical or drug exposures
Common variable immunodeficiency disease	Phenytoin
Acquired immunodeficiency diseases	Dioxin, phenoxy herbicides
iatrogenic immunosuppression	Radiation
HIV-1 infection	Prior chemotherapy and radiation therapy
Acquired hypogammaglobulinemia	

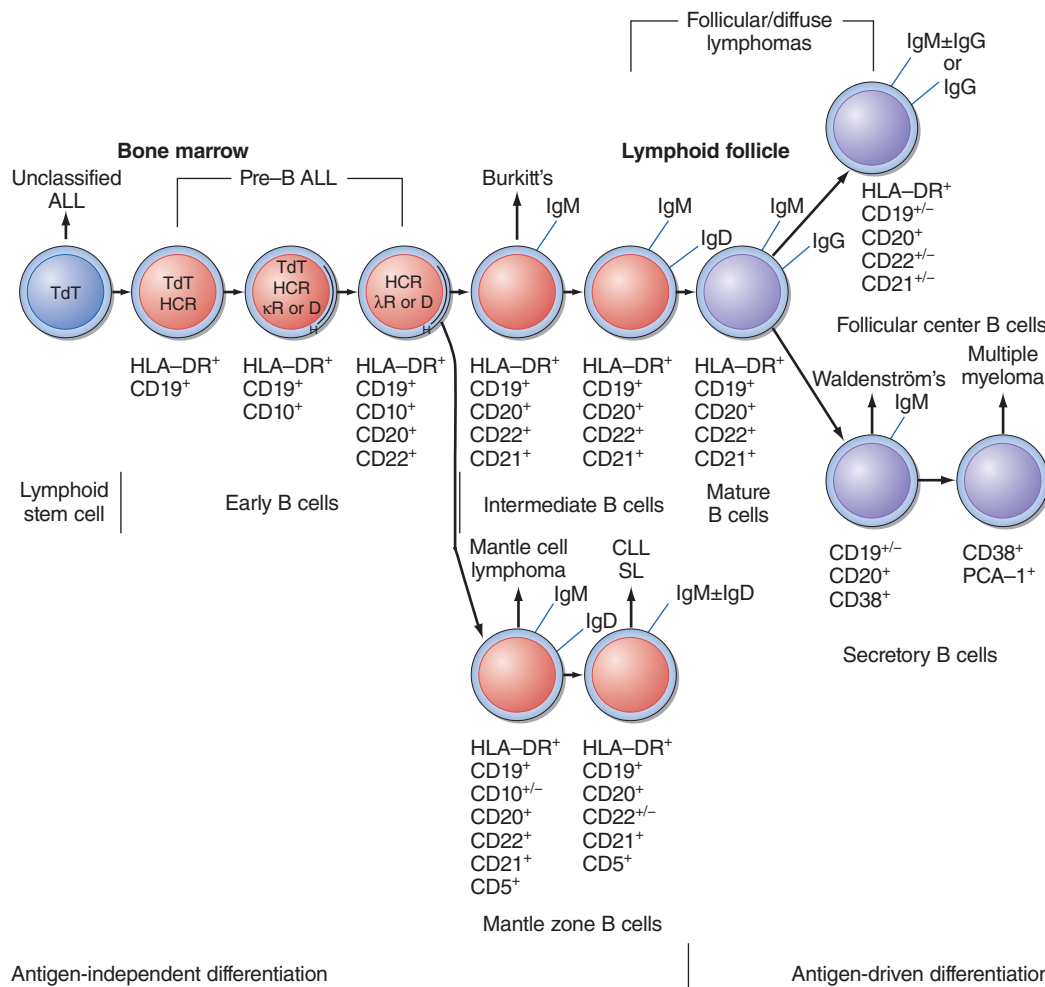


FIGURE 134-2 Pathway of normal B-cell differentiation and relationship to B-cell lymphomas. HLA-DR, CD10, CD19, CD20, CD21, CD22, CD5, and CD38 are cell markers used to distinguish stages of development. Terminal transferase (TdT) is a cellular enzyme. Immunoglobulin heavy chain gene rearrangement (HCR) and light chain gene rearrangement or deletion (κ R or D, λ R or D) occur early in B-cell development. The approximate normal stage of differentiation associated with particular lymphomas is shown. ALL, acute lymphoid leukemia; CLL, chronic lymphoid leukemia; SL, small lymphocytic lymphoma.

including changes in cell-surface phenotype, that characterizes normal B-cell development is shown in Fig. 134-2. A cell becomes committed to T-cell differentiation upon migration to the thymus and rearrangement of T-cell antigen receptor genes. The sequence of the events that characterize T-cell development is depicted in Fig. 134-3.

Although lymphoid malignancies often retain the cell-surface phenotype of lymphoid cells at particular stages of differentiation, this information is of little consequence. The so-called stage of differentiation of a malignant lymphoma does not predict its natural history. For example, the clinically most aggressive lymphoid leukemia is Burkitt's leukemia, which has the phenotype of a mature follicle center IgM-bearing B cell. Leukemias bearing the immunologic cell-surface phenotype of more primitive cells (e.g., pre-B ALL, CD10+) are less aggressive and more amenable to curative therapy than the "more mature" appearing Burkitt's leukemia cells. Furthermore, the apparent stage of differentiation of the malignant cell does not reflect the stage at which the genetic lesions that gave rise to the malignancy developed. For example, follicular lymphoma has the cell-surface phenotype of a follicle center cell, but its characteristic chromosomal translocation, the t(14;18), which involves juxtaposition of the antiapoptotic *bcl-2* gene next to the immunoglobulin heavy chain gene (see below), had to develop early in ontogeny as an error in the process of immunoglobulin gene rearrangement. Why the subsequent steps that led to transformation became manifest in a cell of follicle center differentiation is not clear.

The major value of cell-surface phenotyping is to aid in the differential diagnosis of lymphoid tumors that appear similar by light

microscopy. For example, benign follicular hyperplasia may resemble follicular lymphoma; however, the demonstration that all the cells bear the same immunoglobulin light chain isotype strongly suggests the mass is a clonal proliferation rather than a polyclonal response to an exogenous stimulus.

Malignancies of lymphoid cells are associated with recurring genetic abnormalities. While specific genetic abnormalities have not been identified for all subtypes of lymphoid malignancies, it is presumed that they exist. Genetic abnormalities can be identified at a variety of levels including gross chromosomal changes (i.e., translocations, additions, or deletions); rearrangement of specific genes that may or may not be apparent from cytogenetic studies; and overexpression, underexpression, or mutation of specific oncogenes. Altered expression or mutation of specific proteins is particularly important. Many lymphomas contain balanced chromosomal translocations involving the antigen receptor genes; immunoglobulin genes on chromosomes 2, 14, and 22 in B cells; and T-cell antigen receptor genes on chromosomes 7 and 14 in T cells. The rearrangement of chromosome segments to generate mature antigen receptors must create a site of vulnerability to aberrant recombination. B cells are even more susceptible to acquiring mutations during their maturation in germinal centers; the generation of antibody of higher affinity requires the introduction of mutations into the variable region genes in the germinal centers. Other nonimmunoglobulin genes, e.g., *bcl-6*, may acquire mutations as well.

In the case of diffuse large B-cell lymphoma, the translocation t(14;18) occurs in ~30% of patients and leads to overexpression of the

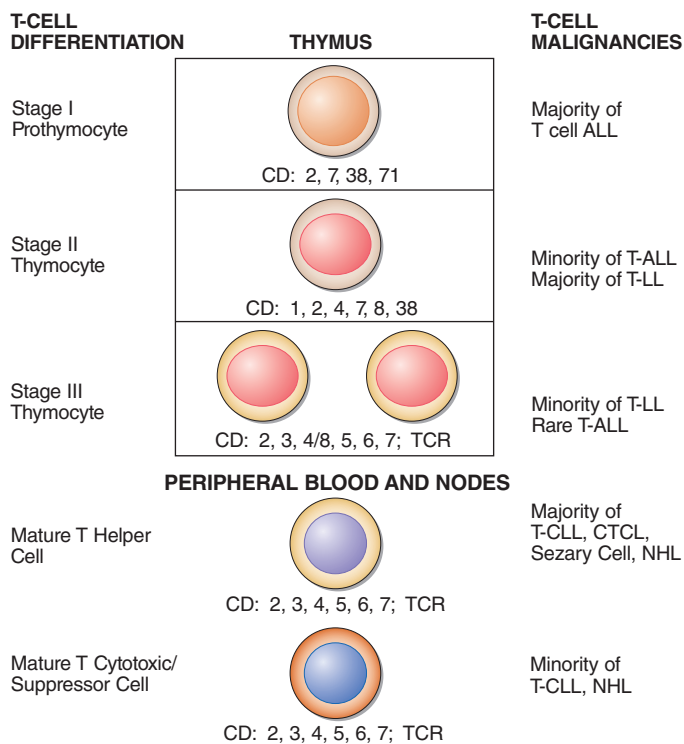


FIGURE 134-3 Pathway of normal T-cell differentiation and relationship to T-cell lymphomas. CD1, CD2, CD3, CD4, CD5, CD6, CD7, CD8, CD38, and CD71 are cell markers used to distinguish stages of development. T-cell antigen receptors (TCR) rearrange in the thymus, and mature T cells emigrate to nodes and peripheral blood. ALL, acute lymphoid leukemia; T-ALL, T-cell ALL; T-LL, T-cell lymphoblastic lymphoma; T-CLL, T-cell chronic lymphoid leukemia; CTCL, cutaneous T-cell lymphoma; NHL, non-Hodgkin's lymphoma.

bcl-2 gene found on chromosome 18. Some other patients without the translocation also overexpress the BCL-2 protein. This protein is involved in suppressing apoptosis—i.e., the mechanism of cell death most often induced by cytotoxic chemotherapeutic agents. A higher relapse rate has been observed in patients whose tumors overexpress the BCL-2 protein, but not in those patients whose lymphoma cells show only the translocation. Thus, particular genetic mechanisms have clinical ramifications.

Table 134-6 presents the most common translocations and associated oncogenes for various subtypes of lymphoid malignancies. In some cases, such as the association of the t(14;18) in follicular lymphoma, the t(2;5) in anaplastic large T/null cell lymphoma, the t(8;14) in Burkitt's lymphoma, and the t(11;14) in mantle cell lymphoma, the great majority of tumors in patients with these diagnoses display these abnormalities. In other types of lymphoma where a minority of the patients have tumors expressing specific genetic abnormalities, the defects may have prognostic significance. No specific genetic abnormalities have been identified in Hodgkin's lymphoma other than aneuploidy.

In typical B-cell CLL, trisomy 12 conveys a poorer prognosis. In ALL in both adults and children, genetic abnormalities have important prognostic significance. Patients whose tumor cells display the t(9;22) and translocations involving the *MLL* gene on chromosome 11q23 have a much poorer outlook than patients who do not have these translocations. Other genetic abnormalities that occur frequently in adults with ALL include the t(4;11) and the t(8;14). The t(4;11) is associated with younger age, female predominance, high white cell counts, and L1 morphology. The t(8;14) is associated with older age, male predominance, frequent CNS involvement, and L3 morphology. Both are associated with a poor prognosis. In childhood ALL, hyperdiploidy has been shown to have a favorable prognosis.

Gene profiling using array technology allows the simultaneous assessment of the expression of thousands of genes. This technology

TABLE 134-6 CYTOGENETIC TRANSLOCATION AND ASSOCIATED ONCOGENES OFTEN SEEN IN LYMPHOID MALIGNANCIES

Disease	Cytogenetic Abnormality	Oncogene
CLL/small lymphocytic lymphoma	t(14;15)(q32;q13)	—
MALT lymphoma	t(11;18)(q21;q21)	<i>API2/MALT, BCL-10</i>
Precursor B-cell acute lymphoid leukemia	t(9;22)(q34;q11) or variant	<i>BCR/ABL</i>
	t(4;11)(q21;q23)	<i>AF4, MLL1</i>
	t(12;21)	<i>TEL, AML1</i>
Precursor acute lymphoid leukemia	t(9;22)	<i>BCR, ABL</i>
	t(1;19)	<i>E2A, PBX</i>
	t(17;19)	<i>HLF, E2A</i>
	t(5;14)	<i>HOX11L2, CTIP2</i>
Mantle cell lymphoma	t(11;14)(q13;q32)	<i>BCL-1, IgH</i>
Follicular lymphoma	t(14;18)(q32;q21)	<i>BCL-2, IgH</i>
Diffuse large cell lymphoma	t(3;-(q27;-) ^a	<i>BCL-6</i>
	t(17;-(p13;-)	<i>p53</i>
Burkitt's lymphoma, Burkitt's leukemia	t(8;-(q24;-) ^a	<i>C-MYC</i>
CD30+ anaplastic large cell lymphoma	t(2;5)(p23;q35)	<i>ALK, NPM</i>
Lymphoplasmacytoid lymphoma	t(9;14)(p13;q32)	<i>PAX5, IgH</i>

^aNumerous sites of translocation may be involved with these genes.

Abbreviations: CLL, chronic lymphoid leukemia; IgH, immunoglobulin heavy chain; MALT, mucosa-associated lymphoid tissue.

provides the possibility to identify new genes with pathologic importance in lymphomas, the identification of patterns of gene expression with diagnostic and/or prognostic significance, and the identification of new therapeutic targets. Recognition of patterns of gene expression is complicated and requires sophisticated mathematical techniques. Early successes using this technology in lymphoma include the identification of previously unrecognized subtypes of diffuse large B-cell lymphoma whose gene expression patterns resemble either those of follicular center B cells or activated peripheral blood B cells. Patients whose lymphomas have a germinal center B-cell pattern of gene expression have a considerably better prognosis than those whose lymphomas have a pattern resembling activated peripheral blood B cells. This improved prognosis is independent of other known prognostic factors. Similar information is being generated in follicular lymphoma and mantle cell lymphoma. The challenge remains to provide information from such techniques in a clinically useful time frame.

APPROACH TO THE PATIENT: Lymphoid Cell Malignancies

Regardless of the type of lymphoid malignancy, the initial evaluation of the patient should include performance of a careful history and physical examination. These will help confirm the diagnosis, identify those manifestations of the disease that might require prompt attention, and aid in the selection of further studies to optimally characterize the patient's status to allow the best choice of therapy. It is difficult to overemphasize the importance of a carefully done history and physical examination. They might provide observations that lead to reconsidering the diagnosis, provide hints at etiology, clarify the stage, and allow the physician to establish rapport with the patient that will make it possible to develop and carry out a therapeutic plan.

For patients with ALL, evaluation is usually completed after a complete blood count, chemistry studies reflecting major organ function, a bone marrow biopsy with genetic and immunologic studies, and a lumbar puncture. The latter is necessary to rule out occult CNS involvement. At this point, most patients would

be ready to begin therapy. In ALL, prognosis is dependent on the genetic characteristics of the tumor, the patient's age, the white cell count, and the patient's overall clinical status and major organ function.

In CLL, the patient evaluation should include a complete blood count, chemistry tests to measure major organ function, serum protein electrophoresis, and a bone marrow biopsy. However, some physicians believe that the diagnosis would not always require a bone marrow biopsy. Patients often have imaging studies of the chest and abdomen looking for pathologic lymphadenopathy. Patients with typical B-cell CLL can be subdivided into three major prognostic groups. Those patients with only blood and bone marrow involvement by leukemia but no lymphadenopathy, organomegaly, or signs of bone marrow failure have the best prognosis. Those with lymphadenopathy and organomegaly have an intermediate prognosis, and patients with bone marrow failure, defined as hemoglobin <100 g/L (10 g/dL) or platelet count <100,000/ μ L, have the worst prognosis. The pathogenesis of the anemia or thrombocytopenia is important to discern. The prognosis is adversely affected when either or both of these abnormalities are due to progressive marrow infiltration and loss of productive marrow. However, either or both may be due to autoimmune phenomena or to hypersplenism that can develop during the course of the disease. These destructive mechanisms are usually completely reversible (glucocorticoids for autoimmune disease; splenectomy for hypersplenism) and do not influence disease prognosis.

Two popular staging systems have been developed to reflect these prognostic groupings (Table 134-7). Patients with typical B-cell CLL can have their course complicated by immunologic abnormalities, including autoimmune hemolytic anemia, autoimmune thrombocytopenia, and hypogammaglobulinemia. Patients with hypogammaglobulinemia benefit from regular (monthly) γ globulin administration. Because of expense, γ globulin is often withheld until the patient experiences a significant infection. These abnormalities do not have a clear prognostic significance and should not be used to assign a higher stage.

Two other features may be used to assess prognosis in B-cell CLL, but neither has yet been incorporated into a staging classification. At least two subsets of CLL have been identified based on the cytoplasmic expression of ZAP-70; expression of this protein, which is usually expressed in T cells, identifies a subgroup with poorer prognosis. A less powerful subsetting tool is CD38 expression. CD38+ tumors tend to have a poorer prognosis than CD38- tumors. A less easily measured feature, the presence of immunoglobulin variable region gene mutations, is also able to separate prognostic groups; patients with mutated immunoglobulin

TABLE 134-8 THE ANN ARBOR STAGING SYSTEM FOR HODGKIN'S LYMPHOMA

Stage	Definition
I	Involvement of a single lymph node region or lymphoid structure (e.g., spleen, thymus, Waldeyer's ring)
II	Involvement of two or more lymph node regions on the same side of the diaphragm (the mediastinum is a single site; hilar lymph nodes should be considered "lateralized" and, when involved on both sides, constitute stage II disease)
III	Involvement of lymph node regions or lymphoid structures on both sides of the diaphragm
III ₁	Subdiaphragmatic involvement limited to spleen, splenic hilar nodes, celiac nodes, or portal nodes
III ₂	Subdiaphragmatic involvement includes paraaortic, iliac, or mesenteric nodes plus structures in III ₁
IV	Involvement of extranodal site(s) beyond that designated as "E" More than one extranodal deposit at any location Any involvement of liver or bone marrow
A	No symptoms
B	Unexplained weight loss of >10% of the body weight during the 6 months before staging investigation Unexplained, persistent, or recurrent fever with temperatures >38°C during the previous month Recurrent drenching night sweats during the previous month
E	Localized, solitary involvement of extralymphatic tissue, excluding liver and bone marrow

variable region genes respond better to treatment and have better survival than those with unmutated immunoglobulin variable region genes.

The initial evaluation of a patient with Hodgkin's lymphoma or non-Hodgkin's lymphoma is similar. In both situations, the determination of an accurate anatomic stage is an important part of the evaluation. Staging is done using the Ann Arbor staging system originally developed for Hodgkin's lymphoma (Table 134-8).

Evaluation of patients with Hodgkin's lymphoma will typically include a complete blood count; erythrocyte sedimentation rate; chemistry studies reflecting major organ function; computed tomography (CT) scans of the chest, abdomen, and pelvis; and a bone marrow biopsy. Neither a positron emission tomography (PET) scan nor a gallium scan is absolutely necessary for primary staging, but one performed at the completion of therapy allows evaluation of persisting radiographic abnormalities, particularly the mediastinum. Knowing that the PET scan or gallium scan is abnormal before treatment can help in this assessment. In most cases, these studies will allow assignment of anatomic stage and the development of a therapeutic plan.

In patients with non-Hodgkin's lymphoma, the same evaluation described for patients with Hodgkin's lymphoma is usually carried out. In addition, serum levels of lactate dehydrogenase (LDH) and β_2 -microglobulin and serum protein electrophoresis are often included in the evaluation. Anatomic stage is assigned in the same manner as used for Hodgkin's lymphoma. However, the prognosis of patients with non-Hodgkin's lymphoma is best assigned using the International Prognostic Index (IPI) (Table 134-9). This is a powerful predictor of outcome in all subtypes of non-Hodgkin's lymphoma. Patients are assigned an IPI score based on the presence or absence of five adverse prognostic factors and may have none or all five of these adverse prognostic factors. Figure 134-4 shows the prognostic significance of this score in 1300 patients with all types of non-Hodgkin's lymphoma. With the addition of rituximab to CHOP (cyclophosphamide, doxorubicin, vincristine, and prednisone), treatment outcomes have improved and the original IPI has lost some of its discrimination power. A revised IPI has been proposed that better predicts outcome of rituximab plus chemotherapy-based programs (Table 134-9). CT scans are routinely used in the evaluation of patients with all subtypes of non-Hodgkin's lymphoma, but PET and

TABLE 134-7 STAGING OF TYPICAL B-CELL LYMPHOID LEUKEMIA

Stage	Clinical Features	Median Survival, Years
Rai System		
0: Low risk	Lymphocytosis only in blood and marrow	>10
I: Intermediate risk	Lymphocytosis + lymphadenopathy	7
II: Intermediate risk	Lymphocytosis + lymphadenopathy + splenomegaly \pm hepatomegaly	
III: High risk	Lymphocytosis + anemia	1.5
IV: High risk	Lymphocytosis + thrombocytopenia	
Binet System		
A	Fewer than three areas of clinical lymphadenopathy; no anemia or thrombocytopenia	>10
B	Three or more involved node areas; no anemia or thrombocytopenia	7
C	Hemoglobin \leq 10 g/dL and/or platelets <100,000/ μ L	2

TABLE 134-9 INTERNATIONAL PROGNOSTIC INDEX FOR NON-HODGKIN'S LYMPHOMA**Five clinical risk factors:**

- Age ≥ 60 years
- Serum lactate dehydrogenase levels elevated
- Performance status ≥ 2 (ECOG) or ≤ 70 (Karnofsky)
- Ann Arbor stage III or IV
- >1 site of extranodal involvement

Patients are assigned a number for each risk factor they have

Patients are grouped differently based on the type of lymphoma

For diffuse large B-cell lymphoma:

- | | |
|-------------------------------------|------------------------------------|
| 0, 1 factor = low risk: | 35% of cases; 5-year survival, 73% |
| 2 factors = low-intermediate risk: | 27% of cases; 5-year survival, 51% |
| 3 factors = high-intermediate risk: | 22% of cases; 5-year survival, 43% |
| 4, 5 factors = high risk: | 16% of cases; 5-year survival, 26% |

For diffuse large B-cell lymphoma treated with R-CHOP:

- | | |
|-------------------------|------------------------------------|
| 0 factor = very good: | 10% of cases; 5-year survival, 94% |
| 1, 2 factors = good: | 45% of cases; 5-year survival, 79% |
| 3, 4, 5 factors = poor: | 45% of cases; 5-year survival, 55% |

Abbreviations: ECOG, Eastern Cooperative Oncology Group; R-CHOP, rituximab, cyclophosphamide, doxorubicin, vincristine, prednisone.

gallium scans are much more useful in aggressive subtypes such as diffuse large B-cell lymphoma than in more indolent subtypes such as follicular lymphoma or small lymphocytic lymphoma. Although the IPI does divide patients with follicular lymphoma into subsets with distinct prognoses, the distribution of such patients is skewed toward lower-risk categories. A follicular lymphoma-specific IPI (FLIPI) has been proposed that replaces performance status with hemoglobin level (<120 g/L [<12 g/dL]) and number of extranodal sites with number of nodal sites (more than four). Low risk (zero or one factor) was assigned to 36% of patients, intermediate risk (two factors) to 37%, and poor risk (more than two factors) to 27% of patients.

CLINICAL FEATURES, TREATMENT, AND PROGNOSIS OF SPECIFIC LYMPHOID MALIGNANCIES

PRECURSOR CELL B-CELL NEOPLASMS

Precursor B-Cell Lymphoblastic Leukemia/Lymphoma The most common cancer in childhood is B-cell ALL. Although this disorder can also present as a lymphoma in either adults or children, presentation as lymphoma is rare.

The malignant cells in patients with precursor B-cell lymphoblastic leukemia are most commonly of pre-B cell origin. Patients typically

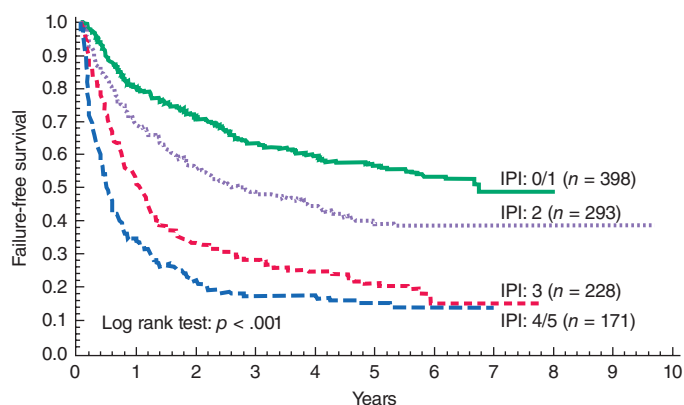


FIGURE 134-4 Relationship of International Prognostic Index (IPI) to survival. Kaplan-Meier survival curves for 1300 patients with various kinds of lymphoma stratified according to the IPI.

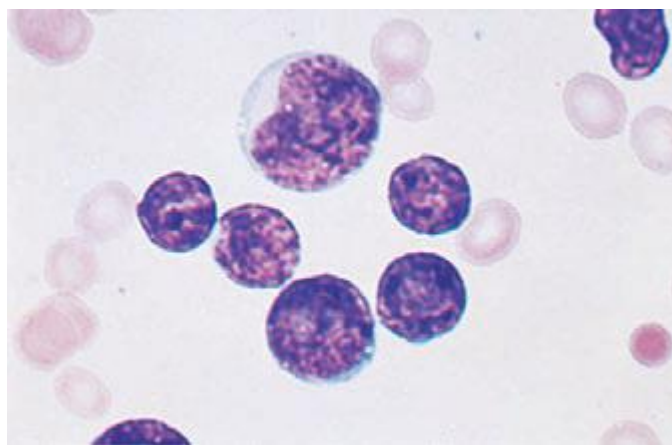


FIGURE 134-5 Acute lymphoblastic leukemia. The cells are heterogeneous in size and have round or convoluted nuclei, high nuclear/cytoplasmic ratio, and absence of cytoplasmic granules.

present with signs of bone marrow failure such as pallor, fatigue, bleeding, fever, and infection related to peripheral blood cytopenias. Peripheral blood counts regularly show anemia and thrombocytopenia but might show leukopenia, a normal leukocyte count, or leukocytosis based largely on the number of circulating malignant cells (Fig. 134-5). Extradural sites of disease are frequently involved in patients who present with leukemia, including lymphadenopathy, hepato- or splenomegaly, CNS disease, testicular enlargement, and/or cutaneous infiltration.

The diagnosis is usually made by bone marrow biopsy, which shows infiltration by malignant lymphoblasts. Demonstration of a pre-B cell immunophenotype (Fig. 134-2) and, often, characteristic cytogenetic abnormalities (Table 134-6) confirm the diagnosis. An adverse prognosis in patients with precursor B-cell ALL is predicted by a very high white cell count, the presence of symptomatic CNS disease, and unfavorable cytogenetic abnormalities. For example, t(9;22), frequently found in adults with B-cell ALL, has been associated with a very poor outlook. The bcr/abl kinase inhibitors have improved the prognosis.

TREATMENT PRECURSOR B-CELL LYMPHOBLASTIC LEUKEMIA

The treatment of patients with precursor B-cell ALL involves remission induction with combination chemotherapy, a consolidation phase that includes administration of high-dose systemic therapy and treatment to eliminate disease in the CNS, and a period of continuing therapy to prevent relapse and effect cure. The overall cure rate in children is 90%, whereas ~50% of adults are long-term disease-free survivors. This reflects the high proportion of adverse cytogenetic abnormalities seen in adults with precursor B-cell ALL.

Precursor B-cell lymphoblastic lymphoma is a rare presentation of precursor B-cell lymphoblastic malignancy. These patients often have a rapid transformation to leukemia and should be treated as though they had presented with leukemia. The few patients who present with the disease confined to lymph nodes have a high cure rate.

MATURE (PERIPHERAL) B-CELL NEOPLASMS

B-Cell Chronic Lymphoid Leukemia/Small Lymphocytic Lymphoma B-cell CLL/small lymphocytic lymphoma represents the most common lymphoid leukemia, and when presenting as a lymphoma, it accounts for ~7% of non-Hodgkin's lymphomas. Presentation can be as either leukemia or lymphoma. The major clinical characteristics of B-cell CLL/small lymphocytic lymphoma are presented in Table 134-10.

The diagnosis of typical B-cell CLL is made when an increased number of circulating lymphocytes (i.e., $>4 \times 10^9/L$ and usually

TABLE 134-10 CLINICAL CHARACTERISTICS OF PATIENTS WITH COMMON TYPES OF NON-HODGKIN'S LYMPHOMA (NHL)

Disease	Median Age, Years	Frequency in Children	% Male	Stage I/II vs III/IV, %	B Symptoms, %	Bone Marrow Involvement, %	Gastrointestinal Tract Involvement, %	% Surviving 5 Years
B-cell chronic lymphocytic leukemia/small lymphocytic lymphoma	65	Rare	53	9 vs 91	33	72	3	51
Mantle cell lymphoma	63	Rare	74	20 vs 80	28	64	9	27
Extranodal marginal zone B-cell lymphoma of MALT type	60	Rare	48	67 vs 33	19	14	50	74
Follicular lymphoma	59	Rare	42	33 vs 67	28	42	4	72
Diffuse large B-cell lymphoma	64	~25% of childhood NHL	55	54 vs 46	33	16	18	46
Burkitt's lymphoma	31	~30% of childhood NHL	89	62 vs 38	22	33	11	45
Precursor T-cell lymphoblastic lymphoma	28	~40% of childhood NHL	64	11 vs 89	21	50	4	26
Anaplastic large T/null-cell lymphoma	34	Common	69	51 vs 49	53	13	9	77
Peripheral T-cell NHL	61	~5% of childhood NHL	55	20 vs 80	50	36	15	25

Abbreviation: MALT, mucosa-associated lymphoid tissue.

>10 × 10⁹/L) is found (Fig. 134-6) that are monoclonal B cells expressing the CD5 antigen. Finding bone marrow infiltration by the same cells confirms the diagnosis. The peripheral blood smear in such patients typically shows many “smudge” or “basket” cells, nuclear remnants of cells damaged by the physical shear stress of making the blood smear. If cytogenetic studies are performed, trisomy 12 is found in 25–30% of patients. Abnormalities in chromosome 13 are also seen.

If the primary presentation is lymphadenopathy and a lymph node biopsy is performed, pathologists usually have little difficulty in making the diagnosis of small lymphocytic lymphoma based on morphologic findings and immunophenotype. However, even in these patients, 70–75% will be found to have bone marrow involvement and circulating monoclonal B lymphocytes are often present.

The differential diagnosis of typical B-cell CLL is extensive (Table 134-1). Immunophenotyping will eliminate the T-cell disorders and can often help sort out other B-cell malignancies. For example, only mantle cell lymphoma and typical B-cell CLL are usually CD5 positive. Typical B-cell small lymphocytic lymphoma can be confused with other B-cell disorders, including lymphoplasmacytic lymphoma (i.e., the tissue manifestation of Waldenström's macroglobulinemia), nodal marginal zone B-cell lymphoma, and mantle cell lymphoma. In addition, some small lymphocytic lymphomas have areas of large

cells that can lead to confusion with diffuse large B-cell lymphoma. An expert hematopathologist is vital for making this distinction.

Typical B-cell CLL is often found incidentally when a complete blood count is done for another reason. However, complaints that might lead to the diagnosis include fatigue, frequent infections, and new lymphadenopathy. The diagnosis of typical B-cell CLL should be considered in a patient presenting with an autoimmune hemolytic anemia or autoimmune thrombocytopenia. B-cell CLL has also been associated with red cell aplasia. When this disorder presents as lymphoma, the most common abnormality is asymptomatic lymphadenopathy, with or without splenomegaly. The staging systems predict prognosis in patients with typical B-cell CLL (Table 134-7). The evaluation of a new patient with typical B-cell CLL/small lymphocytic lymphoma will include many of the studies (Table 134-11) that are used in patients with other non-Hodgkin's lymphomas. In addition, particular attention needs to be given to detecting immune abnormalities such as autoimmune hemolytic anemia, autoimmune thrombocytopenia, hypogammaglobulinemia, and red cell aplasia. Molecular analysis of immunoglobulin gene sequences in CLL has demonstrated that about half the patients have tumors expressing mutated immunoglobulin genes and half have tumors expressing unmutated or germline immunoglobulin sequences. Patients with unmutated immunoglobulins tend to have a more aggressive clinical course and are less responsive to therapy. Unfortunately, immunoglobulin gene sequencing is not routinely available. CD38

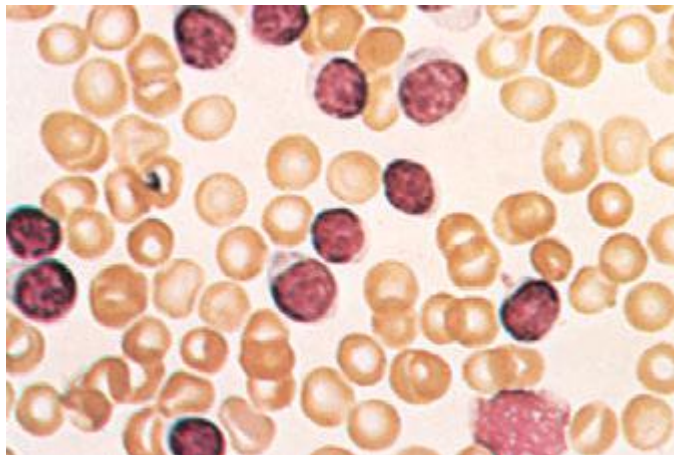


FIGURE 134-6 Chronic lymphocytic leukemia. The peripheral white blood cell count is high due to increased numbers of small, well-differentiated, normal-appearing lymphocytes. The leukemia lymphocytes are fragile, and substantial numbers of broken, smudged cells are usually also present on the blood smear.

TABLE 134-11 STAGING EVALUATION FOR NON-HODGKIN'S LYMPHOMA

Physical examination
Documentation of B symptoms
Laboratory evaluation
Complete blood counts
Liver function tests
Uric acid
Calcium
Serum protein electrophoresis
Serum β_2 -microglobulin
Chest radiograph
CT scan of abdomen, pelvis, and usually chest
Bone marrow biopsy
Lumbar puncture in lymphoblastic, Burkitt's, and diffuse large B-cell lymphoma with positive marrow biopsy
Gallium scan (SPECT) or PET scan in large cell lymphoma

Abbreviations: CT, computed tomography; PET, positron emission tomography; SPECT, single-photon emission computed tomography.

expression is said to be low in the better-prognosis patients expressing mutated immunoglobulin and high in poorer-prognosis patients expressing unmutated immunoglobulin, but this test has not been confirmed as a reliable means of distinguishing the two groups. ZAP-70 expression correlates with the presence of unmutated immunoglobulin genes, but the assay is not yet standardized and widely available.

TREATMENT B-CELL CHRONIC LYMPHOID LEUKEMIA/ SMALL LYMPHOCYTIC LYMPHOMA

Patients whose presentation is typical B-cell CLL with no manifestations of the disease other than bone marrow involvement and lymphocytosis (i.e., Rai stage 0 and Binet stage A; Table 134-7) can be followed without specific therapy for their malignancy. These patients have a median survival >10 years, and some will never require therapy for this disorder. If the patient has an adequate number of circulating normal blood cells and is asymptomatic, many physicians would not initiate therapy for patients in the intermediate stage of the disease manifested by lymphadenopathy and/or hepatosplenomegaly. However, the median survival for these patients is ~7 years, and most will require treatment in the first few years of follow-up. Patients who present with bone marrow failure (i.e., Rai stage III or IV or Binet stage C) will require initial therapy in almost all cases. These patients have a serious disorder with a median survival of only 1.5 years. It must be remembered that immune manifestations of typical B-cell CLL should be managed independently of specific antileukemia therapy. For example, glucocorticoid therapy for autoimmune cytopenias and γ globulin replacement for patients with hypogammaglobulinemia should be used whether or not antileukemia therapy is given.

Patients who present primarily with lymphoma and have a low IPI score have a 5-year survival of ~75%, but those with a high IPI score have a 5-year survival of <40% and are more likely to require early therapy.

The most common treatments for patients with typical B-cell CLL/small lymphocytic lymphoma have been chlorambucil or fludarabine, alone or in combination. Chlorambucil can be administered orally with few immediate side effects, while fludarabine is administered IV and is associated with significant immune suppression. However, fludarabine is by far the more active agent and is the only drug associated with a significant incidence of complete remission. The combination of rituximab (375–500 mg/m² day 1), fludarabine (25 mg/m² days 2–4 on cycle 1 and days 1–3 in subsequent cycles), and cyclophosphamide (250 mg/m² with fludarabine) achieves complete responses in 69% of patients, and those responses are associated with molecular remissions in half of the cases. Half the patients experience grade III or IV neutropenia. For young patients presenting with leukemia requiring therapy, regimens containing fludarabine are the treatment of choice. Because fludarabine is an effective second-line agent in patients with tumors unresponsive to chlorambucil, the latter agent is often chosen in elderly patients who require therapy. Bendamustine, an alkylating agent structurally related to nitrogen mustard, is highly effective and is vying with fludarabine as the primary treatment of choice. Patients who present with lymphoma (rather than leukemia) are also highly responsive to bendamustine, and some patients will receive a combination chemotherapy regimen used in other lymphomas such as CVP (cyclophosphamide, vincristine, and prednisone) or CHOP plus rituximab. Alemtuzumab (anti-CD52) is an antibody with activity in the disease, but it kills both B and T cells and is associated with more immune compromise than rituximab. Young patients with this disease can be candidates for bone marrow transplantation. Allogeneic bone marrow transplantation can be curative but is associated with a significant treatment-related mortality rate. Mini-transplants using immunosuppressive rather than myeloablative doses of preparative drugs are being studied (Chap. 139e). The use of autologous transplantation in patients with this disorder has been discouraging.

At least two newer anti-CD20 monoclonal antibodies have become available, ofatumumab and obinutuzumab. Both have activity in previously treated patients. Agents targeting signaling pathways, such as ibrutinib, an irreversible inhibitor of Bruton's tyrosine kinase, and idelalisib, an inhibitor of phosphoinositide-3-kinase delta, also have antitumor effects. The ideal combination and sequence of these therapies have not been defined.

Extranodal Marginal Zone B-Cell Lymphoma of MALT Type Extranodal marginal zone B-cell lymphoma of MALT type (MALT lymphoma) makes up ~8% of non-Hodgkin's lymphomas. This small cell lymphoma presents in extranodal sites. It was previously considered a small lymphocytic lymphoma or sometimes a pseudolymphoma. The recognition that the gastric presentation of this lymphoma was associated with *H. pylori* infection was an important step in recognizing it as a separate entity. The clinical characteristics of MALT lymphoma are presented in Table 134-10.

The diagnosis of MALT lymphoma can be made accurately by an expert hematopathologist based on a characteristic pattern of infiltration of small lymphocytes that are monoclonal B cells and CD5 negative. In some cases, transformation to diffuse large B-cell lymphoma occurs, and both diagnoses may be made in the same biopsy. The differential diagnosis includes benign lymphocytic infiltration of extranodal organs and other small cell B-cell lymphomas.

MALT lymphoma may occur in the stomach, orbit, intestine, lung, thyroid, salivary gland, skin, soft tissues, bladder, kidney, and CNS. It may present as a new mass, be found on routine imaging studies, or be associated with local symptoms such as upper abdominal discomfort in gastric lymphoma. Most MALT lymphomas are gastric in origin. At least two genetic forms of gastric MALT exist: one (accounting for ~50% of cases) characterized by t(11;18)(q21;q21) that juxtaposes the amino terminal of the *API2* gene with the carboxy terminal of the *MALT1* gene creating an *API2/MALT1* fusion product, and the other characterized by multiple sites of genetic instability including trisomies of chromosomes 3, 7, 12, and 18. About 95% of gastric MALT lymphomas are associated with *H. pylori* infection, and those that are do not usually express t(11;18). The t(11;18) usually results in activation of nuclear factor- κ B (NF- κ B), which acts as a survival factor for the cells. Lymphomas with t(11;18) translocations are genetically stable and do not evolve to diffuse large B-cell lymphoma. By contrast, t(11;18)-negative MALT lymphomas often acquire *BCL6* mutations and progress to aggressive histology lymphoma. MALT lymphomas are localized to the organ of origin in ~40% of cases and to the organ and regional lymph nodes in ~30% of patients. However, distant metastasis can occur—particularly with transformation to diffuse large B-cell lymphoma. Many patients who develop this lymphoma will have an autoimmune or inflammatory process such as Sjögren's syndrome (salivary gland MALT), Hashimoto's thyroiditis (thyroid MALT), *Helicobacter* gastritis (gastric MALT), *C. psittaci* conjunctivitis (ocular MALT), or *Borrelia* skin infections (cutaneous MALT).

Evaluation of patients with MALT lymphoma follows the pattern (Table 134-11) for staging a patient with non-Hodgkin's lymphoma. In particular, patients with gastric lymphoma need to have studies performed to document the presence or absence of *H. pylori* infection. Endoscopic studies including ultrasound can help define the extent of gastric involvement. Most patients with MALT lymphoma have a good prognosis, with a 5-year survival of ~75%. In patients with a low IPI score, the 5-year survival is ~90%, whereas it drops to ~40% in patients with a high IPI score.

TREATMENT MUCOSA-ASSOCIATED LYMPHOID TISSUE LYMPHOMA

MALT lymphoma is often localized. Patients with gastric MALT lymphomas who are infected with *H. pylori* can achieve remission in the 80% of cases with eradication of the infection. These remissions can be durable, but molecular evidence of persisting neoplasia is not

704 infrequent. After *H. pylori* eradication, symptoms generally improve quickly, but molecular evidence of persistent disease may be present for 12–18 months. Additional therapy is not indicated unless progressive disease is documented. Patients with more extensive disease or progressive disease are most often treated with single-agent chemotherapy such as chlorambucil. Combination regimens that include rituximab are also highly effective. Coexistent diffuse large B-cell lymphoma must be treated with combination chemotherapy (see below). The additional acquired mutations that mediate the histologic progression also convey *Helicobacter* independence to the growth.

Mantle Cell Lymphoma Mantle cell lymphoma makes up ~6% of all non-Hodgkin's lymphomas. This lymphoma was previously placed in a number of other subtypes. Its existence was confirmed by the recognition that these lymphomas have a characteristic chromosomal translocation, t(11;14), between the immunoglobulin heavy chain gene on chromosome 14 and the *bcl-1* gene on chromosome 11, and regularly overexpress the BCL-1 protein, also known as cyclin D1. Table 134-10 shows the clinical characteristics of mantle cell lymphoma.

The diagnosis of mantle cell lymphoma can be made accurately by an expert hematopathologist. As with all subtypes of lymphoma, an adequate biopsy is important. The differential diagnosis of mantle cell lymphoma includes other small cell B-cell lymphomas. In particular, mantle cell lymphoma and small lymphocytic lymphoma share a characteristic expression of CD5. Mantle cell lymphoma usually has a slightly indented nucleus.

The most common presentation of mantle cell lymphoma is with palpable lymphadenopathy, frequently accompanied by systemic symptoms. The median age is 63 years, and men are affected four times as commonly as women. Approximately 70% of patients will be stage IV at the time of diagnosis, with frequent bone marrow and peripheral blood involvement. Of the extranodal organs that can be involved, gastrointestinal involvement is particularly important to recognize. Patients who present with lymphomatous polyposis in the large intestine usually have mantle cell lymphoma. Table 134-11 outlines the evaluation of patients with mantle cell lymphoma. Patients who present with gastrointestinal tract involvement often have Waldeyer's ring involvement, and vice versa. The 5-year survival for all patients with mantle cell lymphoma is ~25%, with only occasional patients who present with a high IPI score surviving 5 years and ~50% of patients with a low IPI score surviving 5 years.

TREATMENT MANTLE CELL LYMPHOMA

Current therapies for mantle cell lymphoma are evolving. Patients with localized disease might be treated with combination chemotherapy followed by radiotherapy; however, these patients are exceedingly rare. For the usual presentation with disseminated disease, standard lymphoma treatments have been unsatisfactory, with the minority of patients achieving complete remission. Aggressive combination chemotherapy regimens followed by autologous or allogeneic bone marrow transplantation are frequently offered to younger patients. For the occasional elderly, asymptomatic patient, observation followed by single-agent chemotherapy might be the most practical approach. An intensive combination chemotherapy regimen originally used in the treatment of acute leukemia, HyperC-VAD (cyclophosphamide, vincristine, doxorubicin, dexamethasone, cytarabine, and methotrexate), in combination with rituximab, seems to be associated with better response rates, particularly in younger patients. Alternating two regimens, HyperC-VAD with rituximab added (R-HyperC-VAD) and rituximab plus high-dose methotrexate and cytarabine, can achieve complete responses in >80% of patients and an 8-year survival of 56%, comparable to regimens using high-dose therapy and autologous hematopoietic stem cell transplantation. Bendamustine plus rituximab has been found to induce complete responses in about 31% of patients, but the responses are generally not long lasting. Bortezomib and temsirolimus are

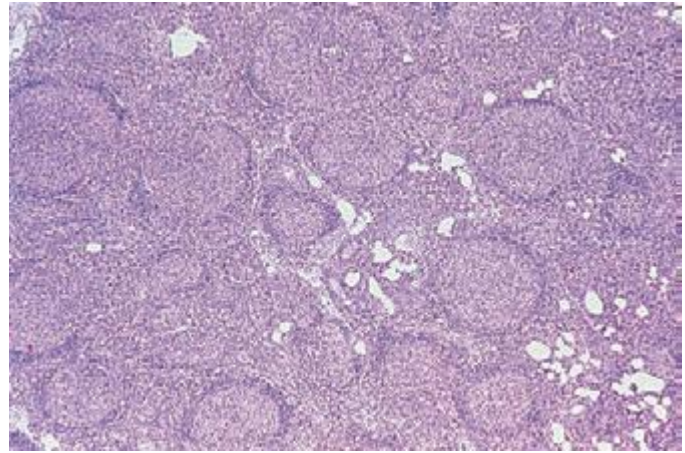


FIGURE 134-7 Follicular lymphoma. The normal nodal architecture is effaced by nodular expansions of tumor cells. Nodules vary in size and contain predominantly small lymphocytes with cleaved nuclei along with variable numbers of larger cells with vesicular chromatin and prominent nucleoli.

single agents that induce transient partial responses in a minority of patients and are being added to primary combinations.

Follicular Lymphoma Follicular lymphomas make up 22% of non-Hodgkin's lymphomas worldwide and at least 30% of non-Hodgkin's lymphomas diagnosed in the United States. This type of lymphoma can be diagnosed accurately on morphologic findings alone and has been the diagnosis in the majority of patients in therapeutic trials for “low-grade” lymphoma in the past. The clinical characteristics of follicular lymphoma are presented in Table 134-10.

Evaluation of an adequate biopsy by an expert hematopathologist is sufficient to make a diagnosis of follicular lymphoma. The tumor is composed of small cleaved and large cells in varying proportions organized in a follicular pattern of growth (Fig. 134-7). Confirmation of B-cell immunophenotype and the existence of the t(14;18) and abnormal expression of BCL-2 protein are confirmatory. The major differential diagnosis is between lymphoma and reactive follicular hyperplasia. The coexistence of diffuse large B-cell lymphoma must be considered. Patients with follicular lymphoma are often subclassified into those with predominantly small cells, those with a mixture of small and large cells, and those with predominantly large cells. Although this distinction cannot be made simply or very accurately, these subdivisions do have prognostic significance. Patients with follicular lymphoma with predominantly large cells have a higher proliferative fraction, progress more rapidly, and have a shorter overall survival with simple chemotherapy regimens.

The most common presentation for follicular lymphoma is with new, painless lymphadenopathy. Multiple sites of lymphoid involvement are typical, and unusual sites such as epitrochlear nodes are sometimes seen. However, essentially any organ can be involved, and extranodal presentations do occur. Most patients do not have fevers, sweats, or weight loss, and an IPI score of 0 or 1 is found in ~50% of patients. Fewer than 10% of patients have a high (i.e., 4 or 5) IPI score. The staging evaluation for patients with follicular lymphoma should include the studies shown in Table 134-11.

TREATMENT FOLLICULAR LYMPHOMA

Follicular lymphoma is one of the malignancies most responsive to chemotherapy and radiotherapy. In addition, tumors in as many as 25% of the patients undergo spontaneous regression—usually transient—without therapy. In an asymptomatic patient, no initial treatment and watchful waiting can be an appropriate management strategy and is particularly likely to be adopted for older patients

with advanced-stage disease. For patients who do require treatment, single-agent chlorambucil or cyclophosphamide or combination chemotherapy with CVP or CHOP is most frequently used. With adequate treatment, 50–75% of patients will achieve a complete remission. Although most patients relapse (median response duration is ~2 years), at least 20% of complete responders will remain in remission for >10 years. For the rare patients (15%) with localized follicular lymphoma, involved-field radiotherapy produces long-term disease-free survival in the majority.

A number of therapies have been shown to be active in the treatment of patients with follicular lymphoma. These include cytotoxic agents such as fludarabine, biologic agents such as interferon α , monoclonal antibodies with or without radionuclides, and lymphoma vaccines. In patients treated with a doxorubicin-containing combination chemotherapy regimen, interferon α given to patients in complete remission seems to prolong survival, but interferon toxicities can affect quality of life. The monoclonal antibody rituximab can cause objective responses in 35–50% of patients with relapsed follicular lymphoma, and radiolabeled antibodies appear to have response rates well in excess of 50%. The addition of rituximab to CHOP and other effective combination chemotherapy programs achieves prolonged overall survival and a decreased risk of histologic progression. Complete remissions can be noted in 85% or more of patients treated with R-CHOP, and median remission durations can exceed 6 or 7 years. Maintenance intermittent rituximab therapy can prolong remissions even further, although it is not completely clear that overall survival is prolonged. Some trials with tumor vaccines have been encouraging. Both autologous and allogeneic hematopoietic stem cell transplantations yield high complete response rates in patients with relapsed follicular lymphoma, and long-term remissions can occur in 40% or more of patients.

Patients with follicular lymphoma with a predominance of large cells have a shorter survival when treated with single-agent chemotherapy but seem to benefit from receiving an anthracycline-containing combination chemotherapy regimen plus rituximab. When their disease is treated aggressively, the overall survival for such patients is no lower than for patients with other follicular lymphomas, and the failure-free survival is superior.

Patients with follicular lymphoma have a high rate of histologic transformation to diffuse large B-cell lymphoma (5–7% per year). This is recognized ~40% of the time during the course of the illness by repeat biopsy and is present in almost all patients at autopsy. This transformation is usually heralded by rapid growth of lymph nodes—often localized—and the development of systemic symptoms such as fevers, sweats, and weight loss. Although these patients have a poor prognosis, aggressive combination chemotherapy regimens can sometimes cause a complete remission in the diffuse large B-cell lymphoma, at times leaving the patient with persisting follicular lymphoma. With more frequent use of R-CHOP to treat follicular lymphoma at diagnosis, it appears that the rate of histologic progression is decreasing. R-CHOP or bendamustine plus rituximab with intermittent rituximab maintenance for 2 years are the most commonly used treatment approaches.

Diffuse Large B-Cell Lymphoma Diffuse large B-cell lymphoma is the most common type of non-Hodgkin's lymphoma, representing approximately one-third of all cases. This lymphoma makes up the majority of cases in previous clinical trials of “aggressive” or “intermediate-grade” lymphoma. Table 134-10 shows the clinical characteristics of diffuse large B-cell lymphoma.

The diagnosis of diffuse large B-cell lymphoma can be made accurately by an expert hematopathologist (Fig. 134-8). Cytogenetic and molecular genetic studies are not necessary for diagnosis, but some evidence has accumulated that patients whose tumors overexpress the BCL-2 protein might be more likely to relapse than others. A subset of patients have tumors with mutations in *BCL6* and translocations involving *MYC*; these are called “double-hit” lymphomas and typically have more aggressive growth and are more poorly responsive to

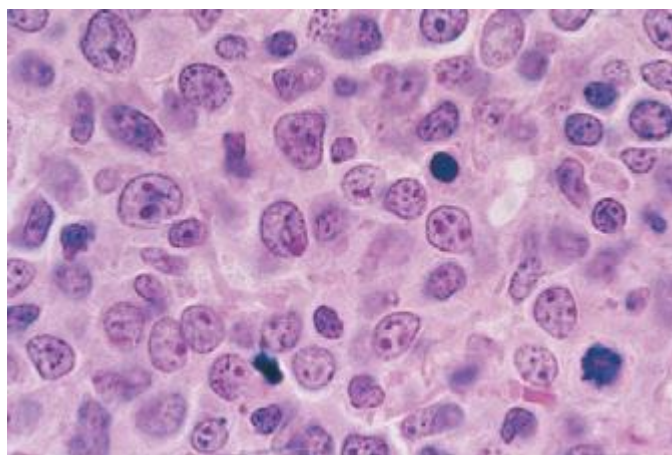


FIGURE 134-8 Diffuse large B-cell lymphoma. The neoplastic cells are heterogeneous but predominantly large cells with vesicular chromatin and prominent nucleoli.

treatment than other diffuse large B-cell lymphomas. Patients with prominent mediastinal involvement are sometimes diagnosed as a separate subgroup having primary mediastinal diffuse large B-cell lymphoma. This latter group of patients has a younger median age (i.e., 37 years) and a female predominance (66%). Subtypes of diffuse large B-cell lymphoma, including those with an immunoblastic subtype and tumors with extensive fibrosis, are recognized by pathologists but do not appear to have important independent prognostic significance.

Diffuse large B-cell lymphoma can present as either primary lymph node disease or at extranodal sites. More than 50% of patients will have some site of extranodal involvement at diagnosis, with the most common sites being the gastrointestinal tract and bone marrow, each being involved in 15–20% of patients. Essentially any organ can be involved, making a diagnostic biopsy imperative. For example, diffuse large B-cell lymphoma of the pancreas has a much better prognosis than pancreatic carcinoma but would be missed without biopsy. Primary diffuse large B-cell lymphoma of the brain is being diagnosed with increasing frequency. Other unusual subtypes of diffuse large B-cell lymphoma such as pleural effusion lymphoma and intravascular lymphoma have been difficult to diagnose and associated with a very poor prognosis.

Table 134-11 shows the initial evaluation of patients with diffuse large B-cell lymphoma. After a careful staging evaluation, ~50% of patients will be found to have stage I or II disease, and ~50% will have widely disseminated lymphoma. Bone marrow biopsy shows involvement by lymphoma in ~15% of cases, with marrow involvement by small cells more frequent than by large cells.

TREATMENT DIFFUSE LARGE B-CELL LYMPHOMA

The initial treatment of all patients with diffuse large B-cell lymphoma should be with a combination chemotherapy regimen. The most popular regimen in the United States is CHOP plus rituximab, although a variety of other anthracycline-containing combination chemotherapy regimens appear to be equally efficacious. Patients with stage I or nonbulky stage II disease can be effectively treated with three to four cycles of combination chemotherapy with or without subsequent involved-field radiotherapy. The need for radiation therapy is unclear. Cure rates of 70–80% in stage II disease and 85–90% in stage I disease can be expected.

For patients with bulky stage II, stage III, or stage IV disease, six to eight cycles of CHOP plus rituximab are usually administered. A large randomized trial showed the superiority of CHOP combined with rituximab over CHOP alone in elderly patients. A frequent approach would be to administer four cycles of therapy and then reevaluate. If the patient has achieved a complete remission after four cycles, two more cycles of treatment might be given and then therapy discontinued. Using this approach, 70–80% of patients can

706 be expected to achieve a complete remission, and 50–70% of complete responders will be cured. The chances for a favorable response to treatment are predicted by the IPI. In fact, the IPI was developed based on the outcome of patients with diffuse large B-cell lymphoma treated with CHOP-like regimens. For the 35% of patients with a low IPI score of 0–1, the 5-year survival is >70%, whereas for the 20% of patients with a high IPI score of 4–5, the 5-year survival is ~20%. The addition of rituximab to CHOP has improved each of those numbers by ~15%. A number of other factors, including molecular features of the tumor, levels of circulating cytokines and soluble receptors, and other surrogate markers, have been shown to influence prognosis. However, they have not been validated as rigorously as the IPI and have not been uniformly applied clinically.

Because a number of patients with diffuse large B-cell lymphoma are either initially refractory to therapy or relapse after apparently effective chemotherapy, 30–40% of patients will be candidates for salvage treatment at some point. Alternative combination chemotherapy regimens can induce complete remission in as many as 50% of these patients, but long-term disease-free survival is seen in ≤10%. Autologous bone marrow transplantation is superior to salvage chemotherapy at usual doses and leads to long-term disease-free survival in ~40% of patients whose lymphomas remain chemotherapy-sensitive after relapse.

Burkitt's Lymphoma/Leukemia Burkitt's lymphoma/leukemia is a rare disease in adults in the United States, making up <1% of non-Hodgkin's lymphomas, but it makes up ~30% of childhood non-Hodgkin's lymphoma. Burkitt's leukemia, or L3 ALL, makes up a small proportion of childhood and adult acute leukemias. Table 134-10 shows the clinical features of Burkitt's lymphoma.

Burkitt's lymphoma can be diagnosed morphologically by an expert hematopathologist with a high degree of accuracy. The cells are homogeneous in size and shape (Fig. 134-9). Demonstration of a very high proliferative fraction and the presence of the t(8;14) or one of its variants, t(2;8) (*c-myc* and the λ light chain gene) or t(8;22) (*c-myc* and the κ light chain gene), can be confirmatory. Burkitt's cell leukemia is recognized by the typical monotonous mass of medium-sized cells with round nuclei, multiple nucleoli, and basophilic cytoplasm with cytoplasmic vacuoles. Demonstration of surface expression of immunoglobulin and one of the above-noted cytogenetic abnormalities is confirmatory.

Three distinct clinical forms of Burkitt's lymphoma are recognized: endemic, sporadic, and immunodeficiency-associated. Endemic and sporadic Burkitt's lymphomas occur frequently in children in Africa, and the sporadic form occurs in Western countries.

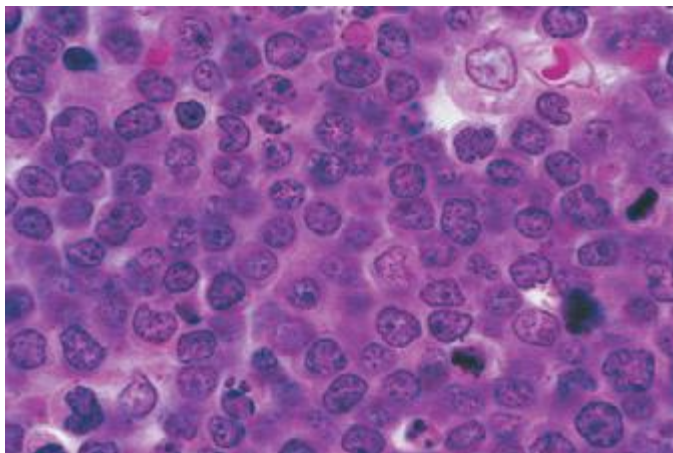


FIGURE 134-9 Burkitt's lymphoma. The neoplastic cells are homogeneous, medium-sized B cells with frequent mitotic figures, a morphologic correlate of high growth fraction. Reactive macrophages are scattered through the tumor, and their pale cytoplasm in a background of blue-staining tumor cells gives the tumor a so-called starry appearance.

Immunodeficiency-associated Burkitt's lymphoma is seen in patients with HIV infection.

Pathologists sometimes have difficulty distinguishing between Burkitt's lymphoma and diffuse large B-cell lymphoma. In the past, a separate subgroup of non-Hodgkin's lymphoma intermediate between the two was recognized. When tested, this subgroup could not be diagnosed accurately. Distinction between the two major types of B-cell aggressive non-Hodgkin's lymphoma can sometimes be made based on the extremely high proliferative fraction seen in patients with Burkitt's lymphoma (i.e., essentially 100% of tumor cells are in cycle) caused by *c-myc* deregulation.

Most patients in the United States with Burkitt's lymphoma present with peripheral lymphadenopathy or an intraabdominal mass. The disease is rapidly progressive and has a propensity to metastasize to the CNS. Initial evaluation should always include an examination of cerebrospinal fluid to rule out metastasis in addition to the other staging evaluations noted in Table 134-11. Once the diagnosis of Burkitt's lymphoma is suspected, a diagnosis must be made promptly, and staging evaluation must be accomplished expeditiously. This is the most rapidly progressive human tumor, and any delay in initiating therapy can adversely affect the patient's prognosis.

TREATMENT BURKITT'S LYMPHOMA

Treatment of Burkitt's lymphoma in both children and adults should begin within 48 h of diagnosis and involves the use of intensive combination chemotherapy regimens incorporating high doses of cyclophosphamide. Prophylactic therapy to the CNS is mandatory. Burkitt's lymphoma was one of the first cancers shown to be curable by chemotherapy. Today, cure can be expected in 70–80% of both children and young adults when effective therapy is administered precisely. Salvage therapy has been generally ineffective in patients in whom the initial treatment fails, emphasizing the importance of the initial treatment approach.

Other B-Cell Lymphoid Malignancies *B-cell prolymphocytic leukemia* involves blood and marrow infiltration by large lymphocytes with prominent nucleoli. Patients typically have a high white cell count, splenomegaly, and minimal lymphadenopathy. The chances for a complete response to therapy are poor.

Hairy cell leukemia is a rare disease that presents predominantly in older males. Typical presentation involves pancytopenia, although occasional patients will have a leukemic presentation. Splenomegaly is usual. The malignant cells appear to have “hairy” projections on light and electron microscopy and show a characteristic staining pattern with tartrate-resistant acid phosphatase. Bone marrow is typically not able to be aspirated, and biopsy shows a pattern of fibrosis with diffuse infiltration by the malignant cells. Patients with this disorder have monocytopenia and are prone to unusual infections, including infection by *Mycobacterium avium intracellulare*, and to vasculitic syndromes. Hairy cell leukemia is responsive to chemotherapy with interferon α , pentostatin, or cladribine, with the latter being the usually preferred treatment. Clinical complete remissions with cladribine occur in the majority of patients, and long-term disease-free survival is frequent. Many of these tumors have the V600E *BRAF* mutation and accordingly are responsive to *BRAF* inhibitors like vemurafenib.

Splenic marginal zone lymphoma involves infiltration of the splenic white pulp by small, monoclonal B cells. This is a rare disorder that can present as leukemia as well as lymphoma. Definitive diagnosis is often made at splenectomy, which is also an effective therapy. This is an extremely indolent disorder, but when chemotherapy is required, the most usual treatment has been chlorambucil.

Lymphoplasmacytic lymphoma is the tissue manifestation of Waldenström's macroglobulinemia (Chap. 136). Many of these tumors harbor a specific mutation, L265P, in *MYD88*, a change that leads to NF- κ B activation. This type of lymphoma has been associated with chronic hepatitis C virus infection, and an etiologic association has been proposed. Patients typically present with lymphadenopathy,

splenomegaly, bone marrow involvement, and occasionally peripheral blood involvement. The tumor cells do not express CD5. Patients often have a monoclonal IgM protein, high levels of which can dominate the clinical picture with the symptoms of hyperviscosity. Treatment of lymphoplasmacytic lymphoma can be aimed primarily at reducing the abnormal protein, if present, but will usually also involve chemotherapy. Chlorambucil, fludarabine, and cladribine have been used. The median 5-year survival for patients with this disorder is ~60%.

Nodal marginal zone lymphoma, also known as *monocytoid cell lymphoma*, represents ~1% of non-Hodgkin's lymphomas. This lymphoma has a slight female predominance and presents with disseminated disease (i.e., stage III or IV) in 75% of patients. Approximately one-third of patients have bone marrow involvement, and a leukemic presentation occasionally occurs. The staging evaluation and therapy should use the same approach as used for patients with follicular lymphoma. Approximately 60% of the patients with nodal marginal zone lymphoma will survive 5 years after diagnosis.

Other more uncommon B-cell malignancies are discussed in Chap. 135e.

PRECURSOR T-CELL MALIGNANCIES

Precursor T-Cell Lymphoblastic Leukemia/Lymphoma Precursor T-cell malignancies can present either as ALL or as an aggressive lymphoma. These malignancies are more common in children and young adults, with males more frequently affected than females.

Precursor T-cell ALL can present with bone marrow failure, although the severity of anemia, neutropenia, and thrombocytopenia is often less than in precursor B-cell ALL. These patients sometimes have very high white cell counts, a mediastinal mass, lymphadenopathy, and hepatosplenomegaly. Precursor T-cell lymphoblastic lymphoma is most often found in young men presenting with a large mediastinal mass and pleural effusions. Both presentations have a propensity to metastasize to the CNS, and CNS involvement is often present at diagnosis.

TREATMENT

PRECURSOR T-CELL LYMPHOBLASTIC LEUKEMIA/LYMPHOMA

Children with precursor T-cell ALL seem to benefit from very intensive remission induction and consolidation regimens. The majority of patients treated in this manner can be cured. Older children and young adults with precursor T-cell lymphoblastic lymphoma are also often treated with "leukemia-like" regimens. Patients who present with localized disease have an excellent prognosis. However, advanced age is an adverse prognostic factor. Adults with precursor T-cell lymphoblastic lymphoma who present with high LDH levels or bone marrow or CNS involvement are often offered bone marrow transplantation as part of their primary therapy.

MATURE (PERIPHERAL) T-CELL DISORDERS

Mycosis Fungoides Mycosis fungoides is also known as *cutaneous T-cell lymphoma*. This lymphoma is more often seen by dermatologists than internists. The median age of onset is in the mid-fifties, and the disease is more common in males and in blacks.

Mycosis fungoides is an indolent lymphoma with patients often having several years of eczematous or dermatitic skin lesions before the diagnosis is finally established. The skin lesions progress from patch stage to plaque stage to cutaneous tumors. Early in the disease, biopsies are often difficult to interpret, and the diagnosis may only become apparent by observing the patient over time. In advanced stages, the lymphoma can spread to lymph nodes and visceral organs. Patients with this lymphoma may develop generalized erythroderma and circulating tumor cells, called *Sézary's syndrome*.

Rare patients with localized early-stage mycosis fungoides can be cured with radiotherapy, often total-skin electron beam irradiation. More advanced disease has been treated with topical glucocorticoids, topical nitrogen mustard, phototherapy, psoralen with ultraviolet A

(PUVA), extracorporeal photopheresis, retinoids (bexarotene), electron beam radiation, interferon, antibodies, fusion toxins, histone deacetylase inhibitors, and systemic cytotoxic therapy. Unfortunately, these treatments are palliative.

Adult T-Cell Lymphoma/Leukemia Adult T-cell lymphoma/leukemia is one manifestation of infection by the HTLV-1 retrovirus. Patients can be infected through transplacental transmission, mother's milk, blood transfusion, and by sexual transmission of the virus. Patients who acquire the virus from their mother through breast milk are most likely to develop lymphoma, but the risk is still only 2.5% and the latency averages 55 years. Nationwide testing for HTLV-1 antibodies and the aggressive implementation of public health measures could theoretically lead to the disappearance of adult T-cell lymphoma/leukemia. Tropical spastic paraparesis, another manifestation of HTLV-1 infection (Chap. 225e), occurs after a shorter latency (1–3 years) and is most common in individuals who acquire the virus during adulthood from transfusion or sex.

The diagnosis of adult T-cell lymphoma/leukemia is made when an expert hematopathologist recognizes the typical morphologic picture, a T-cell immunophenotype (i.e., CD4 positive), and the presence in serum of antibodies to HTLV-1. Examination of the peripheral blood will usually reveal characteristic, pleomorphic abnormal CD4-positive cells with indented nuclei, which have been called "flower" cells (Fig. 134-10).

A subset of patients have a smoldering clinical course and long survival, but most patients present with an aggressive disease manifested by lymphadenopathy, hepatosplenomegaly, skin infiltration, pulmonary infiltrates, hypercalcemia, lytic bone lesions, and elevated LDH levels. The skin lesions can be papules, plaques, tumors, and ulcerations. Lung lesions can be either tumor or opportunistic infection in light of the underlying immunodeficiency in the disease. Bone marrow involvement is not usually extensive, and anemia and thrombocytopenia are not usually prominent. Although treatment with combination chemotherapy regimens can result in objective responses, true complete remissions are unusual, and the median survival of patients is ~7 months. A small phase II study reported a high response rate with interferon plus zidovudine and arsenic trioxide.

Anaplastic Large T/Null-Cell Lymphoma Anaplastic large T/null-cell lymphoma was previously usually diagnosed as undifferentiated carcinoma or malignant histiocytosis. Discovery of the CD30 (Ki-1) antigen and the recognition that some patients with previously unclassified malignancies displayed this antigen led to the identification of a new type of lymphoma. Subsequently, discovery of the t(2;5) and the resultant frequent overexpression of the anaplastic lymphoma kinase (ALK) protein confirmed the existence of this entity. This lymphoma accounts for ~2% of all non-Hodgkin's lymphomas. Table 134-10 shows the clinical characteristics of patients with anaplastic large T/null cell lymphoma.

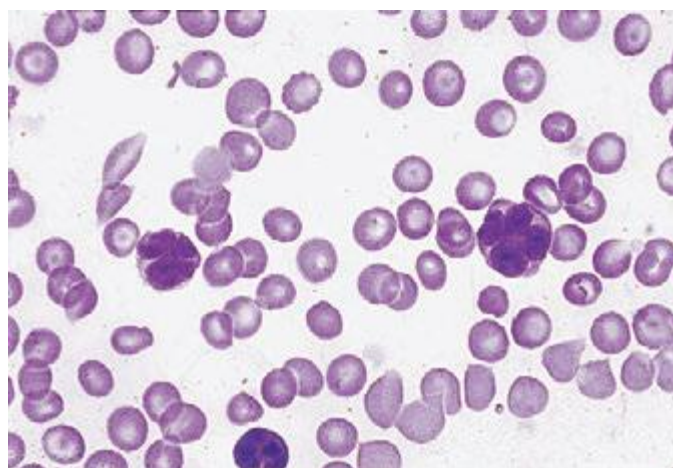


FIGURE 134-10 Adult T-cell leukemia/lymphoma. Peripheral blood smear showing leukemia cells with typical "flower-shaped" nucleus.

The diagnosis of anaplastic large T/null-cell lymphoma is made when an expert hematopathologist recognizes the typical morphologic picture and a T-cell or null-cell immunophenotype with CD30 positivity. Documentation of the t(2;5) and/or overexpression of ALK protein confirm the diagnosis. Some diffuse large B-cell lymphomas can also have an anaplastic appearance but have the same clinical course or response to therapy as other diffuse large B-cell lymphomas. A small percentage of anaplastic lymphomas are ALK negative.

Patients with anaplastic large T/null-cell lymphoma are typically young (median age, 33 years) and male (~70%). Some 50% of patients present in stage I/II, and the remainder present with more extensive disease. Systemic symptoms and elevated LDH levels are seen in about one-half of patients. Bone marrow and the gastrointestinal tract are rarely involved, but skin involvement is frequent. Some patients with disease confined to the skin have a different and more indolent disorder that has been termed *cutaneous anaplastic large T/null-cell lymphoma* and might be related to lymphomatoid papulosis.

TREATMENT ANAPLASTIC LARGE T/NULL-CELL LYMPHOMA

Treatment regimens appropriate for other aggressive lymphomas, such as diffuse large B-cell lymphoma, should be used in patients with anaplastic large T/null-cell lymphoma, with the exception that the B-cell-specific antibody, rituximab, is omitted. Surprisingly, given the anaplastic appearance, this disorder has the best survival rate of any aggressive lymphoma. The 5-year survival is >75%. While traditional prognostic factors such as the IPI predict treatment outcome, overexpression of the ALK protein is an important prognostic factor, with patients overexpressing this protein having a superior treatment outcome. The ALK inhibitor crizotinib appears highly active as well. In addition, the CD30 immunotoxin, brentuximab vedotin, is active in the disease.

Peripheral T-Cell Lymphoma The peripheral T-cell lymphomas make up a heterogeneous morphologic group of aggressive neoplasms that share a mature T-cell immunophenotype. They represent ~7% of all cases of non-Hodgkin's lymphoma. A number of distinct clinical syndromes are included in this group of disorders. Table 134-10 shows the clinical characteristics of patients with peripheral T-cell lymphoma.

The diagnosis of peripheral T-cell lymphoma, or any of its specific subtypes, requires an expert hematopathologist, an adequate biopsy, and immunophenotyping. Most peripheral T-cell lymphomas are CD4+, but a few will be CD8+, both CD4+ and CD8+, or have an NK cell immunophenotype. No characteristic genetic abnormalities have yet been identified, but translocations involving the T-cell antigen receptor genes on chromosomes 7 or 14 may be detected. The differential diagnosis of patients suspected of having peripheral T-cell lymphoma includes reactive T-cell infiltrative processes. In some cases, demonstration of a monoclonal T-cell population using T-cell receptor gene rearrangement studies will be required to make a diagnosis.

The initial evaluation of a patient with a peripheral T-cell lymphoma should include the studies in Table 134-11 for staging patients with non-Hodgkin's lymphoma. Unfortunately, patients with peripheral T-cell lymphoma usually present with adverse prognostic factors, with >80% of patients having an IPI score ≥ 2 and >30% having an IPI score ≥ 4 . As this would predict, peripheral T-cell lymphomas are associated with a poor outcome, and only 25% of the patients survive 5 years after diagnosis. Treatment regimens are the same as those used for diffuse large B-cell lymphoma (omitting rituximab), but patients with peripheral T-cell lymphoma have a poorer response to treatment. Because of this poor treatment outcome, hematopoietic stem cell transplantation is often considered early in the care of young patients.

A number of specific clinical syndromes are seen in the peripheral T-cell lymphomas. *Angioimmunoblastic T-cell lymphoma* is one of the more common subtypes, making up ~20% of T-cell lymphomas. These

patients typically present with generalized lymphadenopathy, fever, weight loss, skin rash, and polyclonal hypergammaglobulinemia. In some cases, it is difficult to separate patients with a reactive disorder from those with true lymphoma.

Extranodal T/NK-cell lymphoma of nasal type has also been called *angiocentric lymphoma* and was previously termed *lethal midline granuloma*. This disorder is more frequent in Asia and South America than in the United States and Europe. EBV is thought to play an etiologic role. Although most frequent in the upper airway, it can involve other organs. The course is aggressive, and patients frequently have the hemophagocytic syndrome. When marrow and blood involvement occur, distinction between this disease and leukemia might be difficult. Some patients will respond to aggressive combination chemotherapy regimens, but the overall outlook is poor.

Enteropathy-type intestinal T-cell lymphoma is a rare disorder that occurs in patients with untreated gluten-sensitive enteropathy. Patients are frequently wasted and sometimes present with intestinal perforation. The prognosis is poor. *Hepatosplenic $\gamma\delta$ T-cell lymphoma* is a systemic illness that presents with sinusoidal infiltration of the liver, spleen, and bone marrow by malignant T cells. Tumor masses generally do not occur. The disease is associated with systemic symptoms and is often difficult to diagnose. Treatment outcome is poor. *Subcutaneous panniculitis-like T-cell lymphoma* is a rare disorder that is often confused with panniculitis. Patients present with multiple subcutaneous nodules, which progress and can ulcerate. Hemophagocytic syndrome is common. Response to therapy is poor. The development of the hemophagocytic syndrome (profound anemia, ingestion of erythrocytes by monocytes and macrophages, elevated ferritin levels) in the course of any peripheral T-cell lymphoma is generally associated with a fatal outcome.

HODGKIN'S LYMPHOMA

Classical Hodgkin's Lymphoma Hodgkin's lymphoma occurs in 9000 patients in the United States each year, and the disease does not appear to be increasing in frequency. Most patients present with palpable lymphadenopathy that is nontender; in most patients, these lymph nodes are in the neck, supraclavicular area, and axilla. More than half the patients will have mediastinal adenopathy at diagnosis, and this is sometimes the initial manifestation. Subdiaphragmatic presentation of Hodgkin's lymphoma is unusual and more common in older males. One-third of patients present with fevers, night sweats, and/or weight loss—B symptoms in the Ann Arbor staging classification (Table 134-8). Occasionally, Hodgkin's lymphoma can present as a fever of unknown origin. This is more common in older patients who are found to have mixed-cellularity Hodgkin's lymphoma in an abdominal site. Rarely, the fevers persist for days to weeks, followed by afebrile intervals and then recurrence of the fever. This pattern is known as *Pel-Ebstein fever*. Hodgkin's lymphoma can occasionally present with unusual manifestations. These include severe and unexplained itching, cutaneous disorders such as erythema nodosum and ichthyosiform atrophy, paraneoplastic cerebellar degeneration and other distant effects on the CNS, nephrotic syndrome, immune hemolytic anemia and thrombocytopenia, hypercalcemia, and pain in lymph nodes on alcohol ingestion.

The diagnosis of Hodgkin's lymphoma is established by review of an adequate biopsy specimen by an expert hematopathologist. In the United States, most patients have nodular sclerosing Hodgkin's lymphoma, with a minority of patients having mixed-cellularity Hodgkin's lymphoma. Lymphocyte-predominant and lymphocyte-depleted Hodgkin's lymphoma are rare. Mixed-cellularity Hodgkin's lymphoma or lymphocyte-depletion Hodgkin's lymphoma are seen more frequently in patients infected by HIV (Fig. 134-11). Hodgkin's lymphoma is a tumor characterized by rare neoplastic cells of B-cell origin (immunoglobulin genes are rearranged but not expressed) in a tumor mass that is largely polyclonal inflammatory infiltrate, probably a reaction to cytokines produced by the tumor cells. The differential diagnosis of a lymph node biopsy suspicious for Hodgkin's lymphoma includes inflammatory processes, mononucleosis, non-Hodgkin's lymphoma, phenytoin-induced adenopathy, and nonlymphomatous malignancies.

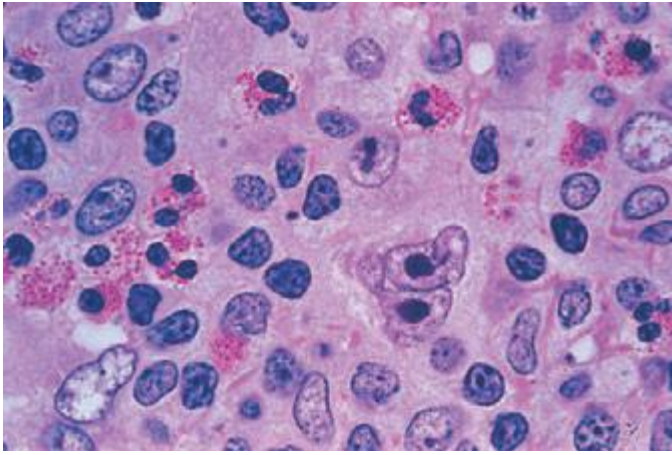


FIGURE 134-11 Mixed-cellularity Hodgkin's lymphoma. A Reed-Sternberg cell is present near the center of the field; a large cell with a bilobed nucleus and prominent nucleoli giving an "owl's eyes" appearance. The majority of the cells are normal lymphocytes, neutrophils, and eosinophils that form a pleomorphic cellular infiltrate.

The staging evaluation for a patient with Hodgkin's lymphoma would typically include a careful history and physical examination; complete blood count; erythrocyte sedimentation rate; serum chemistry studies including LDH; chest radiograph; CT scan of the chest, abdomen, and pelvis; and bone marrow biopsy. Many patients would also have a PET scan or a gallium scan. Although rarely used, a bipedal lymphangiogram can be helpful. PET and gallium scans are most useful to document remission. Staging laparotomies were once popular for most patients with Hodgkin's lymphoma but are now done rarely because of an increased reliance on systemic rather than local therapy.

TREATMENT CLASSICAL HODGKIN'S LYMPHOMA

Patients with localized Hodgkin's lymphoma are cured >90% of the time. In patients with good prognostic factors, extended-field radiotherapy has a high cure rate. Increasingly, patients with all stages of Hodgkin's lymphoma are treated initially with chemotherapy. Patients with localized or good-prognosis disease receive a brief course of chemotherapy followed by radiotherapy to sites of node involvement. Patients with more extensive disease or those with B symptoms receive a complete course of chemotherapy. The most popular chemotherapy regimen used in Hodgkin's lymphoma is a combination of doxorubicin, bleomycin, vinblastine, and dacarbazine (ABVD). Today, most patients in the United States receive ABVD, but a weekly chemotherapy regimen administered for 12 weeks called *Stanford V* is becoming increasingly popular, but it includes radiation therapy, which has been associated with life-threatening late toxicities such as premature coronary artery disease and second solid tumors. In Europe, a high-dose regimen called *BEACOPP* incorporating alkylating agents has become popular and might have a better response rate in very-high-risk patients. Long-term disease-free survival in patients with advanced disease can be achieved in >75% of patients who lack systemic symptoms and in 60–70% of patients with systemic symptoms.

Patients who relapse after primary therapy of Hodgkin's lymphoma can frequently still be cured. Patients who relapse after initial treatment with only radiotherapy have an excellent outcome when treated with chemotherapy. Patients who relapse after an effective chemotherapy regimen are usually not curable with subsequent chemotherapy administered at standard doses. However, patients with a long initial remission can be an exception to this rule. Autologous bone marrow transplantation can cure half of patients in whom effective chemotherapy regimens fail to induce durable remissions. The immunotoxin, brentuximab vedotin, a CD30-directed chemotherapy that selectively targets cells expressing CD30, is active in

the salvage setting and is being integrated into ABVD for initial treatment.

Because of the very high cure rate in patients with Hodgkin's lymphoma, long-term complications have become a major focus for clinical research. In fact, in some series of patients with early-stage disease, more patients died from late complications of therapy than from Hodgkin's lymphoma itself. This is particularly true in patients with localized disease. The most serious late side effects include second malignancies and cardiac injury. Patients are at risk for the development of acute leukemia in the first 10 years after treatment with combination chemotherapy regimens that contain alkylating agents plus radiation therapy. The risk for development of acute leukemia appears to be greater after MOPP-like (mechlorethamine, vincristine, procarbazine, prednisone) regimens than with ABVD. The risk of development of acute leukemia after treatment for Hodgkin's lymphoma is also related to the number of exposures to potentially leukemogenic agents (i.e., multiple treatments after relapse) and the age of the patient being treated, with those age >60 years at particularly high risk. The development of carcinomas as a complication of treatment for Hodgkin's lymphoma has become a major problem. These tumors usually occur ≥ 10 years after treatment and are associated with use of radiotherapy. For this reason, young women treated with thoracic radiotherapy for Hodgkin's lymphoma should institute screening mammograms 5–10 years after treatment, and all patients who receive thoracic radiotherapy for Hodgkin's lymphoma should be discouraged from smoking. Thoracic radiation also accelerates coronary artery disease, and patients should be encouraged to minimize risk factors for coronary artery disease such as smoking and elevated cholesterol levels. Cervical radiation therapy increases the risk of carotid atherosclerosis and stroke.

A number of other late side effects from the treatment of Hodgkin's lymphoma are well known. Patients who receive thoracic radiotherapy are at very high risk for the eventual development of hypothyroidism and should be observed for this complication; intermittent measurement of thyrotropin should be made to identify the condition before it becomes symptomatic. Lhermitte's syndrome occurs in ~15% of patients who receive thoracic radiotherapy. This syndrome is manifested by an "electric shock" sensation into the lower extremities on flexion of the neck. Infertility is a concern for all patients undergoing treatment for Hodgkin's lymphoma. In both women and men, the risk of permanent infertility is age-related, with younger patients more likely to recover fertility. In addition, treatment with ABVD increases the chances to retain fertility.

Nodular Lymphocyte-Predominant Hodgkin's Lymphoma Nodular lymphocyte-predominant Hodgkin's lymphoma is now recognized as an entity distinct from classical Hodgkin's lymphoma. Previous classification systems recognized that biopsies from a subset of patients diagnosed as having Hodgkin's lymphoma contained a predominance of small lymphocytes and rare Reed-Sternberg cells (Fig. 134-11). A subset of these patients have tumors with nodular growth pattern and a clinical course that varied from that of patients with classical Hodgkin's lymphoma. This is an unusual clinical entity and represents <5% of cases of Hodgkin's lymphoma.

Nodular lymphocyte-predominant Hodgkin's lymphoma has a number of characteristics that suggest its relationship to non-Hodgkin's lymphoma. These include a clonal proliferation of B cells and a distinctive immunophenotype; tumor cells express J chain and display CD45 and epithelial membrane antigen (EMA) and do not express two markers normally found on Reed-Sternberg cells, CD30 and CD15. This lymphoma tends to have a chronic, relapsing course and sometimes transforms to diffuse large B-cell lymphoma.

The treatment of patients with nodular lymphocyte-predominant Hodgkin's lymphoma is controversial. Some clinicians favor no treatment and merely close follow-up. In the United States, most physicians will treat localized disease with radiotherapy and disseminated disease with regimens used for patients with classical Hodgkin's lymphoma. Regardless of the therapy used, most series report a long-term survival of >80%.

The most common condition that pathologists and clinicians might confuse with lymphoma is reactive, atypical lymphoid hyperplasia. Patients might have localized or disseminated lymphadenopathy and might have the systemic symptoms characteristic of lymphoma. Underlying causes include a drug reaction to phenytoin or carbamazepine. Immune disorders such as rheumatoid arthritis and lupus erythematosus, viral infections such as cytomegalovirus and EBV, and bacterial infections such as cat-scratch disease may cause adenopathy (Chap. 79). In the absence of a definitive diagnosis after initial biopsy, continued follow-up, further testing, and repeated biopsies, if necessary, constitute the appropriate approach, rather than instituting therapy.

Specific conditions that can be confused with lymphoma include *Castleman's disease*, which can present with localized or disseminated lymphadenopathy; some patients have systemic symptoms. The disseminated form is often accompanied by anemia and polyclonal hypergammaglobulinemia, and the condition has been associated with overproduction of interleukin 6 (IL-6), in some cases produced by human herpesvirus 8 infection. Patients with localized disease can be treated effectively with local therapy, whereas the initial treatment for patients with disseminated disease is usually with systemic glucocorticoids. IL-6-directed therapy (tocilizumab) has produced short-term responses. Rituximab appears to produce longer remissions than tocilizumab.

Sinus histiocytosis with massive lymphadenopathy (Rosai-Dorfman disease) usually presents with bulky lymphadenopathy in children or young adults. The disease is usually nonprogressive and self-limited, but patients can manifest autoimmune hemolytic anemia.

Lymphomatoid papulosis is a cutaneous lymphoproliferative disorder that is often confused with anaplastic large cell lymphoma involving the skin. The cells of lymphomatoid papulosis are similar to those seen in lymphoma and stain for CD30, and T-cell receptor gene rearrangements are sometimes seen. However, the condition is characterized by waxing and waning skin lesions that usually heal, leaving small scars. In the absence of effective communication between the clinician and the pathologist regarding the clinical course in the patient, this disease will be misdiagnosed. Since the clinical picture is usually benign, misdiagnosis is a serious mistake.

ACKNOWLEDGMENT

James Armitage was a coauthor of this chapter in prior editions, and substantial material from those editions has been included here.

135e Less Common Hematologic Malignancies

Ayalew Tefferi, Dan L. Longo

The most common lymphoid malignancies are discussed in [Chap. 134](#), myeloid leukemias in [Chaps. 132 and 133](#), myelodysplastic syndromes in [Chap. 130](#), and myeloproliferative syndromes in [Chap. 131](#). This chapter will focus on the more unusual forms of hematologic malignancy. The diseases discussed here are listed in [Table 135e-1](#). Each of these entities accounts for less than 1% of hematologic neoplasms.

LYMPHOID MALIGNANCIES

Precursor B-cell and precursor T-cell neoplasms are discussed in [Chap. 134](#). All the lymphoid tumors discussed here are mature B cell or T cell, natural killer (NK) cell neoplasms.

MATURE B-CELL NEOPLASMS

B-Cell Prolymphocytic Leukemia (B-PLL) This is a malignancy of medium-sized (about twice the size of a normal small lymphocyte), round lymphocytes with a prominent nucleolus and light blue cytoplasm on Wright's stain. It dominantly affects the blood, bone marrow, and spleen and usually does not cause adenopathy. The median age of

affected patients is 70 years, and men are more often affected than women (male-to-female ratio is 1.6). This entity is distinct from chronic lymphoid leukemia (CLL) and does not develop as a consequence of that disease.

Clinical presentation is generally from symptoms of splenomegaly or incidental detection of an elevated white blood cell (WBC) count. The clinical course can be rapid. The cells express surface IgM (with or without IgD) and typical B-cell markers (CD19, CD20, CD22). CD23 is absent, and about one-third of cases express CD5. The CD5 expression along with the presence of the t(11;14) translocation in 20% of cases leads to confusion in distinguishing B-PLL from the leukemic form of mantle cell lymphoma. No reliable criteria for the distinction have emerged. About half of patients have mutation or loss of p53, and deletions have been noted in 11q23 and 13q14. Nucleoside analogues like fludarabine and cladribine and combination chemotherapy (cyclophosphamide, doxorubicin, vincristine, and prednisone [CHOP]) have produced responses. CHOP plus rituximab may be more effective than CHOP alone, but the disease is sufficiently rare that large series have not been reported. Splenectomy can produce palliation of symptoms but appears to have little or no impact on the course of the disease.

Splenic Marginal Zone Lymphoma (SMZL) This tumor of mainly small lymphocytes originates in the marginal zone of the spleen white pulp, grows to efface the germinal centers and mantle, and invades the red pulp. Splenic hilar nodes, bone marrow, and peripheral blood may be involved. The circulating tumor cells have short surface villi and are called villous lymphocytes. [Table 135e-2](#) shows differences in tumor cells of a number of neoplasms of small lymphocytes that aid in the differential diagnosis. SMZL cells express surface immunoglobulin and CD20, but are negative for CD5, CD10, CD43, and CD103. Lack of CD5 distinguishes SMZL from CLL, and lack of CD103 separates SMZL from hairy cell leukemia.

The median age of patients with SMZL is mid-fifties, and men and women are equally represented. Patients present with incidental or symptomatic splenomegaly or incidental detection of lymphocytosis in the peripheral blood with villous lymphocytes. Autoimmune anemia or thrombocytopenia may be present. The immunoglobulin produced by these cells contains somatic mutations that reflect transit through a germinal center, and ongoing mutations suggest that the mutation machinery has remained active. About 40% of patients have either deletions or translocations involving 7q21, the site of the *FLNC* gene (filamin C γ , involved in cross-linking actin filaments in the cytoplasm). *NOTCH2* mutations are seen in 25% of patients. Chromosome 8p deletions may also be noted. The genetic lesions typically found in extranodal marginal zone lymphomas [e.g., trisomy 3 and t(11;18)] are uncommon in SMZL.

The clinical course of disease is generally indolent with median survivals exceeding 10 years. Patients with elevated lactate dehydrogenase (LDH) levels, anemia, and hypoalbuminemia generally have a poorer prognosis. Long remissions can be seen after splenectomy. Rituximab is also active. A small fraction of patients undergo histologic progression to diffuse large B-cell lymphoma with a concomitant change to a more aggressive natural history. Experience with combination chemotherapy in SMZL is limited.

Hairy Cell Leukemia Hairy cell leukemia is a tumor of small lymphocytes with oval nuclei, abundant cytoplasm, and distinctive membrane projections (hairy cells). Patients have splenomegaly and diffuse bone marrow involvement. While some circulating cells are noted, the clinical picture is dominated by symptoms from the enlarged spleen and pancytopenia. The mechanism of the pancytopenia is not completely clear and may be mediated by both inhibitory cytokines and marrow replacement. The marrow has an increased level of reticulin fibers; indeed, hairy cell leukemia is a common cause of inability to aspirate bone marrow or so-called "dry tap" ([Table 135e-3](#)). Monocytopenia is profound and may explain a predisposition to atypical mycobacterial infection that is observed clinically. The tumor cells have strong expression of CD22, CD25, and CD103; soluble CD25 level in serum is an excellent tumor marker for disease activity. The cells also express tartrate-resistant acid phosphatase. The immunoglobulin genes are

TABLE 135e-1 UNUSUAL LYMPHOID AND MYELOID MALIGNANCIES

Lymphoid
Mature B-cell neoplasms
B-cell prolymphocytic leukemia
Splenic marginal zone lymphoma
Hairy cell leukemia
Nodal marginal zone B-cell lymphoma
Mediastinal large B-cell lymphoma
Intravascular large B-cell lymphoma
Primary effusion lymphoma
Lymphomatoid granulomatosis
Mature T-cell and natural killer (NK) cell neoplasms
T-cell prolymphocytic leukemia
T-cell large granular lymphocytic leukemia
Aggressive NK cell leukemia
Extranodal NK/T-cell lymphoma, nasal type
Enteropathy-type T-cell lymphoma
Hepatosplenic T-cell lymphoma
Subcutaneous panniculitis-like T-cell lymphoma
Blastic NK cell lymphoma
Primary cutaneous CD30+ T-cell lymphoma
Angioimmunoblastic T-cell lymphoma
Myeloid
Chronic neutrophilic leukemia
Chronic eosinophilic leukemia/hypereosinophilic syndrome
Histiocytic and Dendritic Cell Neoplasms
Histiocytic sarcoma
Langerhans cell histiocytosis
Langerhans cell sarcoma
Interdigitating dendritic cell sarcoma
Follicular dendritic cell sarcoma
Mast Cells
Mastocytosis
Cutaneous mastocytosis
Systemic mastocytosis
Mast cell sarcoma
Extracutaneous mastocytoma

TABLE 135e-2 IMMUNOPHENOTYPE OF TUMORS OF SMALL LYMPHOCYTES

	CD5	CD20	CD43	CD10	CD103	slg	CyclinD1
Follicular lymphoma	neg	pos	pos	pos	neg	pos	neg
Chronic lymphoid leukemia	pos	pos	pos	neg	neg	pos	neg
B-cell prolymphocytic leukemia	pos	pos	pos	neg	neg	pos	pos
Mantle cell lymphoma	pos	pos	pos	neg	neg	pos	pos
Splenic marginal zone lymphoma	neg	pos	neg	neg	neg	pos	neg
Hairy cell leukemia	neg	pos	?	neg	pos	pos	neg

Abbreviations: neg, negative; pos, positive.

rearranged and mutated, indicating the influence of a germinal center. No specific cytogenetic abnormality has been found, but most cases contain the activating *BRAF* mutation V600E.

The median age of affected patients is mid-fifties, and the male-to-female ratio is 5:1. Treatment options are numerous. Splenectomy is often associated with prolonged remission. Nucleosides including cladribine and deoxycoformycin are highly active but are also associated with further immunosuppression and can increase the risk of certain opportunistic infections. However, after brief courses of these agents, patients usually obtain very durable remissions during which immune function spontaneously recovers. Interferon α is also an effective therapy but is not as effective as nucleosides. Chemotherapy-refractory patients have responded to vemurafenib, a *BRAF* inhibitor.

Nodal Marginal Zone B-Cell Lymphoma This rare node-based disease bears an uncertain relationship with extranodal marginal zone lymphomas, which are often mucosa-associated and are called mucosa-associated lymphoid tissue (MALT) lymphomas, and SMZLs. Patients may have localized or generalized adenopathy. The neoplastic cell is a marginal zone B cell with monocytoid features and has been called monocytoid B-cell lymphoma in the past. Up to one-third of the patients may have extranodal involvement, and involvement of the lymph nodes can be secondary to the spread of a mucosal primary lesion. In authentic nodal primaries, the cytogenetic abnormalities associated with MALT lymphomas [trisomy 3 and t(11;18)] are very rare. The clinical course is indolent. Patients often respond to combination chemotherapy, although remissions have not been durable. Few patients have received CHOP plus rituximab, which is likely to be an effective approach to management.

Mediastinal (Thymic) Large B-Cell Lymphoma This entity was originally considered a subset of diffuse large B-cell lymphoma; however, additional study has identified it as a distinct entity with its own characteristic clinical, genetic, and immunophenotypic features. This is a disease that can be bulky in size but usually remains confined to the mediastinum. It can be locally aggressive, including progressing to produce a superior vena cava obstruction syndrome or pericardial effusion. About one-third of patients develop pleural effusions, and 5–10% can disseminate widely to kidney, adrenal, liver, skin, and even brain. The disease affects women more often than men (male-to-female ratio is 1:2–3), and the median age is 35–40 years.

The tumor is composed of sheets of large cells with abundant cytoplasm accompanied by variable, but often abundant, fibrosis. It is distinguished from nodular sclerosing Hodgkin's disease by the paucity of normal lymphoid cells and the absence of lacunar variants

of Reed-Sternberg cells. However, more than one-third of the genes that are expressed to a greater extent in primary mediastinal large B-cell lymphoma than in usual diffuse large B-cell lymphoma are also overexpressed in Hodgkin's disease, suggesting a possible pathogenetic relationship between the two entities that affect the same anatomic site. Tumor cells may overexpress *MAL*. The genome of tumor cells is characterized by frequent chromosomal gains and losses. The tumor cells in mediastinal large B-cell lymphoma express CD20, but surface immunoglobulin and HLA class I and class II molecules may be absent or incompletely expressed. Expression of lower levels of class II HLA identifies a subset with poorer prognosis. The cells are CD5 and CD10 negative but may show light staining with anti-CD30. The cells are CD45 positive, unlike cells of classical Hodgkin's disease.

Methotrexate, leucovorin, doxorubicin, cyclophosphamide, vincristine, prednisone, and bleomycin (MACOP-B) and rituximab plus CHOP are effective treatments, achieving 5-year survival of 75–87%. Dose-adjusted therapy with prednisone, etoposide, vincristine, cyclophosphamide, and doxorubicin (EPOCH) plus rituximab has produced 5-year survival of 97%. A role for mediastinal radiation therapy has not been definitively demonstrated, but it is frequently used, especially in patients whose mediastinal area remains positron emission tomography-avid after four to six cycles of chemotherapy.

Intravascular Large B-Cell Lymphoma This is an extremely rare form of diffuse large B-cell lymphoma characterized by the presence of lymphoma in the lumen of small vessels, particularly capillaries. It is also known as malignant angioendotheliomatosis or angiotropic large cell lymphoma. It is sufficiently rare that no consistent picture has emerged to define a clinical syndrome or its epidemiologic and genetic features. It is thought to remain inside vessels because of a defect in adhesion molecules and homing mechanisms, an idea supported by scant data suggesting absence of expression of β -1 integrin and ICAM-1. Patients commonly present with symptoms of small-vessel occlusion, skin lesions, or neurologic symptoms. The tumor cell clusters can promote thrombus formation. In general, the clinical course is aggressive and the disease is poorly responsive to therapy. Often a diagnosis is not made until very late in the course of the disease.

Primary Effusion Lymphoma This entity is another variant of diffuse large B-cell lymphoma that presents with pleural effusions, usually without apparent tumor mass lesions. It is most common in the setting of immune deficiency disease, especially AIDS, and is caused by human herpes virus 8 (HHV-8)/Kaposi's sarcoma herpes virus (KSHV). It is also known as body cavity-based lymphoma. Some patients have been previously diagnosed with Kaposi's sarcoma. It can also occur in the absence of immunodeficiency in elderly men of Mediterranean heritage, similar to Kaposi's sarcoma but even less common.

The malignant effusions contain cells positive for HHV-8/KSHV, and many are also co-infected with Epstein-Barr virus. The cells are large with large nuclei and prominent nucleoli that can be confused with Reed-Sternberg cells. The cells express CD20 and CD79a (immunoglobulin-signaling molecule), although they often do not express immunoglobulin. Some cases aberrantly express T-cell markers such as CD3 or rearranged T-cell receptor genes. No characteristic genetic lesions have been reported, but gains in chromosome 12 and X material has been seen, similar to other HIV-associated lymphomas. The clinical course is generally characterized by rapid progression and death within 6 months.

TABLE 135e-3 DIFFERENTIAL DIAGNOSIS OF "DRY TAP"—INABILITY TO ASPIRATE BONE MARROW

Dry taps occur in about 4% of attempts and are associated with:

Metastatic carcinoma infiltration	17%
Chronic myeloid leukemia	15%
Myelofibrosis	14%
Hairy cell leukemia	10%
Acute leukemia	10%
Lymphomas, Hodgkin's disease	9%
Normal marrow	Rare

Lymphomatoid Granulomatosis This is an angiocentric, angiodestructive lymphoproliferative disease comprised by neoplastic Epstein-Barr virus–infected monoclonal B cells accompanied and outnumbered by a polyclonal reactive T-cell infiltrate. The disease is graded based on histologic features such as cell number and atypia in the B cells. It is most often confused with extranodal NK–T cell lymphoma, nasal type, which can also be angiodestructive and is Epstein-Barr virus–related. The disease usually presents in adults (males > females) as a pulmonary infiltrate. Involvement is often entirely extranodal and can include kidney (32%), liver (29%), skin (25%), and brain (25%). The disease often but not always occurs in the setting of immune deficiency.

The disease can be remitting and relapsing in nature or can be rapidly progressive. The course is usually predicted by the histologic grade. The disease is highly responsive to combination chemotherapy and is curable in most cases. Some investigators have claimed that low-grade disease (grade I and II) can be treated with interferon α .

MATURE T-CELL AND NK CELL NEOPLASMS

T-Cell Prolymphocytic Leukemia This is an aggressive leukemia of medium-sized prolymphocytes involving the blood, marrow, nodes, liver, spleen, and skin. It accounts for 1–2% of all small lymphocytic leukemias. Most patients present with elevated WBC count (often >100,000/ μ L), hepatosplenomegaly, and adenopathy. Skin involvement occurs in 20%. The diagnosis is made from peripheral blood smear, which shows cells about 25% larger than those in small lymphocytes, with cytoplasmic blebs and nuclei that may be indented. The cells express T-cell markers like CD2, CD3, and CD7; two-thirds of patients have cells that are CD4+ and CD8–, and 25% have cells that are CD4+ and CD8+. T-cell receptor β chains are clonally rearranged. In 80% of patients, inversion of chromosome 14 occurs between q11 and q32. Ten percent have t(14;14) translocations that bring the T-cell receptor alpha/beta gene locus into juxtaposition with oncogenes *TCL1* and *TCL1b* at 14q32.1. Chromosome 8 abnormalities are also common. Deletions in the *ATM* gene are also noted. Activating *JAK3* mutations have also been reported.

The course of the disease is generally rapid, with median survival of about 12 months. Responses have been seen with the anti-CD52 antibody, nucleoside analogs, and CHOP chemotherapy. Small numbers of patients with T-cell prolymphocytic leukemia have also been treated with high-dose therapy and allogeneic bone marrow transplantation after remission has been achieved with conventional-dose therapy.

T-Cell Large Granular Lymphocytic Leukemia T-cell large granular lymphocytic leukemia (LGL leukemia) is characterized by increases in the number of LGLs in the peripheral blood (2000–20,000/ μ L) often accompanied by severe neutropenia, with or without concomitant anemia. Patients may have splenomegaly and frequently have evidence of systemic autoimmune disease, including rheumatoid arthritis, hypergammaglobulinemia, autoantibodies, and circulating immune complexes. Bone marrow involvement is mainly interstitial in pattern, with fewer than 50% lymphocytes on differential count. Usually the cells express CD3, T-cell receptors, and CD8; NK-like variants may be CD3–. The leukemic cells often express Fas and Fas ligand.

The course of the disease is generally indolent and dominated by the neutropenia. Paradoxically, immunosuppressive therapy with cyclosporine, methotrexate, or cyclophosphamide plus glucocorticoids can produce an increase in granulocyte counts. Nucleosides have been used anecdotally. Occasionally the disease can accelerate to a more aggressive clinical course.

Aggressive NK Cell Leukemia NK neoplasms are very rare, and they may follow a range of clinical courses from very indolent to highly aggressive. They are more common in Asians than whites, and the cells frequently harbor a clonal Epstein-Barr virus episome. The peripheral blood white count is usually not greatly elevated, but abnormal large lymphoid cells with granular cytoplasm are noted. The aggressive form is characterized by symptoms of fever and laboratory abnormalities of pancytopenia. Hepatosplenomegaly is common; node involvement is less common. Patients may have hemophagocytosis, coagulopathy, or multiorgan failure. Serum levels of Fas ligand are elevated.

The cells express CD2 and CD56 and do not have rearranged T-cell receptor genes. Deletions involving chromosome 6 are common. The disease can be rapidly progressive. Some forms of NK neoplasms are more indolent. They tend to be discovered incidentally with LGL lymphocytosis and do not manifest the fever and hepatosplenomegaly characteristic of the aggressive leukemia. The cells are also CD2 and CD56 positive, but they do not contain clonal forms of Epstein-Barr virus and are not accompanied by pancytopenia or autoimmune disease.

Extranodal NK/T-Cell Lymphoma, Nasal Type Like lymphomatoid granulomatosis, extranodal NK/T-cell lymphoma tends to be an angiocentric and angiodestructive lesion, but the malignant cells are not B cells. In most cases, they are CD56+ Epstein-Barr virus–infected cells; occasionally they are CD56– Epstein-Barr virus–infected cytotoxic T cells. They are most commonly found in the nasal cavity. Historically, this illness was called lethal midline granuloma, polymorphic reticulosis, and angiocentric immunoproliferative lesion. This form of lymphoma is prevalent in Asia, Mexico, and Central and South America; it affects males more commonly than females. When it spreads beyond the nasal cavity, it may affect soft tissue, the gastrointestinal tract, or the testis. In some cases, hemophagocytic syndrome may influence the clinical picture. Patients may have B symptoms. Many of the systemic manifestations of disease are related to the production of cytokines by the tumor cells and the cells responding to their signals. Deletions and inversions of chromosome 6 are common.

Many patients with extranodal NK/T-cell lymphoma, nasal type have excellent antitumor responses with combination chemotherapy regimens, particularly those with localized disease. Radiation therapy is often used after completion of chemotherapy. Four risk factors have been defined, including B symptoms, advanced stage, elevated LDH, and regional lymph node involvement. Patient survival is linked to the number of risk factors: 5-year survival is 81% for zero risk factors, 64% for one risk factor, 32% for two risk factors, and 7% for three or four risk factors. Combination regimens without anthracyclines have been touted as superior to CHOP, but data are sparse. High-dose therapy with stem cell transplantation has been used, but its role is unclear.

Enteropathy-Type T-Cell Lymphoma Enteropathy-type T-cell lymphoma is a rare complication of longstanding celiac disease. It most commonly occurs in the jejunum or the ileum. In adults, the lymphoma may be diagnosed at the same time as celiac disease, but the suspicion is that the celiac disease was a longstanding precursor to the development of lymphoma. The tumor usually presents as multiple ulcerating mucosal masses, but may also produce a dominant exophytic mass or multiple ulcerations. The tumor expresses CD3 and CD7 nearly always and may or may not express CD8. The normal-appearing lymphocytes in the adjacent mucosa often have a similar phenotype to the tumor. Most patients have the HLA genotype associated with celiac disease, HLA DQA1*0501 or DQB1*0201.

The prognosis of this form of lymphoma is typically (median survival is 7 months) poor, but some patients have a good response to CHOP chemotherapy. Patients who respond can develop bowel perforation from responding tumor. If the tumor responds to treatment, recurrence may develop elsewhere in the celiac disease–affected small bowel.

Hepatosplenic T-Cell Lymphoma Hepatosplenic T-cell lymphoma is a malignancy derived from T cells expressing the gamma/delta T-cell antigen receptor that affects mainly the liver and fills the sinusoids with medium-size lymphoid cells. When the spleen is involved, dominantly the red pulp is infiltrated. It is a disease of young people, especially young people with an underlying immunodeficiency or with an autoimmune disease that demands immunosuppressive therapy. The use of thiopurine and infliximab is particularly common in the history of patients with this disease. The cells are CD3+ and usually CD4– and CD8–. The cells may contain isochromosome 7q, often together with trisomy 8. The lymphoma has an aggressive natural history. Combination chemotherapy may induce remissions, but most patients relapse. Median survival is about 2 years. The tumor does not appear to respond to reversal of immunosuppressive therapy.

Subcutaneous Panniculitis-Like T-Cell Lymphoma Subcutaneous panniculitis-like T-cell lymphoma involves multiple subcutaneous collections of neoplastic T cells that are usually cytotoxic cells in phenotype (i.e., contain perforin and granzyme B and express CD3 and CD8). The rearranged T-cell receptor is usually alpha/beta-derived, but occasionally the gamma/delta receptors are involved, particularly in the setting of immunosuppression. The cells are negative for Epstein-Barr virus. Patients may have a hemophagocytic syndrome in addition to the skin infiltration; fever and hepatosplenomegaly may also be present. Nodes are generally not involved. Patients frequently respond to combination chemotherapy, including CHOP. When the disease is progressive, the hemophagocytic syndrome can be a component of a fulminant downhill course. Effective therapy can reverse the hemophagocytic syndrome.

Blastic NK Cell Lymphoma The neoplastic cells express NK cell markers, especially CD56, and are CD3 negative. They are large blastic-appearing cells and may produce a leukemia picture, but the dominant site of involvement is the skin. Morphologically, the cells are similar to the neoplastic cells in acute lymphoid and myeloid leukemia. No characteristic chromosomal abnormalities have been described. The clinical course is rapid, and the disease is largely unresponsive to typical lymphoma treatments.

Primary Cutaneous CD30+ T-Cell Lymphoma This tumor involves the skin and is composed of cells that appear similar to the cells of anaplastic T-cell lymphoma. Among cutaneous T-cell tumors, about 25% are CD30+ anaplastic lymphomas. If dissemination to lymph nodes occurs, it is difficult to distinguish between the cutaneous and systemic forms of the disease. The tumor cells are often CD4+, and the cells contain granules that are positive for granzyme B and perforin in 70% of cases. The typical t(2;5) of anaplastic T-cell lymphoma is absent; indeed, its presence should prompt a closer look for systemic involvement and a switch to a diagnosis of anaplastic T-cell lymphoma. This form of lymphoma has sporadically been noted as a rare complication of silicone on saline breast implants. Cutaneous CD30+ T-cell lymphoma often responds to therapy. Radiation therapy can be effective, and surgery can also produce long-term disease control. Five-year survival exceeds 90%.

Angioimmunoblastic T-Cell Lymphoma Angioimmunoblastic T-cell lymphoma is a systemic disease that accounts for about 15% of all T-cell lymphomas. Patients frequently have fever, advanced stage, diffuse adenopathy, hepatosplenomegaly, skin rash, polyclonal hypergammaglobulinemia, and a wide range of autoantibodies including cold agglutinins, rheumatoid factor, and circulating immune complexes. Patients may have edema, arthritis, pleural effusions, and ascites. The nodes contain a polymorphous infiltrate of neoplastic T cells and nonneoplastic inflammatory cells together with proliferation of high endothelial venules and follicular dendritic cells. The most common chromosomal abnormalities are trisomy 3, trisomy 5, and an extra X chromosome. Aggressive combination chemotherapy can induce regressions. The underlying immune defects make conventional lymphoma treatments more likely to produce infectious complications.

MYELOID MALIGNANCIES

The World Health Organization (WHO) system uses peripheral blood counts and smear analysis, bone marrow morphology, and cytogenetic and molecular genetic tests in order to classify myeloid malignancies into five major categories (Table 135e-4). In this chapter, we focus on chronic neutrophilic leukemia; atypical chronic myeloid leukemia, *BCR-ABL1* negative; chronic myelomonocytic leukemia; juvenile myelomonocytic leukemia; chronic eosinophilic leukemia, not otherwise specified; mastocytosis; myeloproliferative neoplasm (MPN), unclassifiable (MPN-U); myelodysplastic syndrome (MDS)/MPN, unclassifiable (MDS/MPN-U); refractory anemia with ring sideroblasts associated with marked thrombocytosis (RARS-T); and myeloid and lymphoid neoplasms with eosinophilia and abnormalities of *PDGFRA*, *PDGFRB*, or *FGFR1*. This chapter also includes histiocytic and dendritic cell neoplasms, transient myeloproliferative disorders,

TABLE 135e-4 WORLD HEALTH ORGANIZATION CLASSIFICATION OF MYELOID MALIGNANCIES

1. Acute myeloid leukemia (AML) and related precursor neoplasms^a
2. Myeloproliferative neoplasms (MPN)
 - 2.1. Chronic myelogenous leukemia, *BCR-ABL1* positive (CML)
 - 2.2. *BCR-ABL1*-negative MPN
 - 2.2.1. Polycythemia vera
 - 2.2.2. Primary myelofibrosis
 - 2.2.3. Essential thrombocythemia
 - 2.3. Chronic neutrophilic leukemia
 - 2.4. Chronic eosinophilic leukemia, not otherwise specified (CEL-NOS)
 - 2.5. Mastocytosis
 - 2.6. Myeloproliferative neoplasm, unclassifiable (MPN-U)
3. Myelodysplastic syndromes (MDS)
 - 3.1. Refractory cytopenia^b with unilineage dysplasia (RCUD)
 - 3.1.1. Refractory anemia (ring sideroblasts <15% of erythroid precursors)
 - 3.1.2. Refractory neutropenia
 - 3.1.3. Refractory thrombocytopenia
 - 3.2. Refractory anemia with ring sideroblasts (RARS; dysplasia limited to erythroid lineage and ring sideroblasts ≥15% of bone marrow erythroid precursors)
 - 3.3. Refractory cytopenia with multilineage dysplasia (RCMD; ring sideroblast count does not matter)
 - 3.4. Refractory anemia with excess blasts (RAEB)
 - 3.4.1. RAEB-1 (2–4% circulating or 5–9% marrow blasts)
 - 3.4.2. RAEB-2 (5–19% circulating or 10–19% marrow blasts or Auer rods present)
 - 3.5. MDS associated with isolated del(5q)
 - 3.6. MDS, unclassifiable (MDS-U)
4. MDS/MPN overlap
 - 4.1. Chronic myelomonocytic leukemia (CMML)
 - 4.2. Atypical chronic myeloid leukemia, *BCR-ABL1* negative (aCML)
 - 4.3. Juvenile myelomonocytic leukemia (JMML)
 - 4.4. MDS/MPN, unclassifiable (MDS/MPN-U)
 - 4.4.1. Provisional entity: Refractory anemia with ring sideroblasts associated with marked thrombocytosis (RARS-T)
5. Myeloid and lymphoid neoplasms with eosinophilia and abnormalities of *PDGFRA*, *PDGFRB*, or *FGFR1*^c
 - 5.1. Myeloid and lymphoid neoplasms with *PDGFRA* rearrangement
 - 5.2. Myeloid neoplasms with *PDGFRB* rearrangement
 - 5.3. Myeloid and lymphoid neoplasms with *FGFR1* abnormalities

^aAML-related precursor neoplasms include therapy-related MDS and myeloid sarcoma. ^bEither monocytopenia or bicytopenia: hemoglobin level <10 g/dL, absolute neutrophil count <1.8 × 10⁹/L, or platelet count <100 × 10⁹/L. However, higher blood counts do not exclude the diagnosis in the presence of unequivocal histologic/cytogenetic evidence for MDS. ^cGenetic rearrangements involving platelet-derived growth factor receptor α/β (*PDGFRA*/*PDGFRB*) or fibroblast growth factor receptor 1 (*FGFR1*).

and a broader discussion on primary eosinophilic disorders including hypereosinophilic syndrome (HES).

CHRONIC NEUTROPHILIC LEUKEMIA

Chronic neutrophilic leukemia (CNL) is characterized by mature neutrophilic leukocytosis with few or no circulating immature granulocytes. CNL is associated with activating mutations of the gene (*CSF3R*) encoding for the receptor for granulocyte colony-stimulating factor (G-CSF), also known as colony-stimulating factor 3 (CSF3). Patients with CNL might be asymptomatic at presentation but also display constitutional symptoms, splenomegaly, anemia, and thrombocytopenia. Median survival is approximately 2 years, and causes of death include leukemic transformation, progressive disease associated with severe cytopenias, and marked treatment-refractory leukocytosis. CNL is rare, with less than 200 reported cases. Median age at diagnosis is approximately 67 years, and the disease is equally prevalent in both genders.

Pathogenesis CSF3 is the main growth factor for granulocyte proliferation and differentiation. Accordingly, recombinant CSF3 is used for the treatment of severe neutropenia, including severe congenital neutropenia (SCN). Some patients with SCN acquire *CSF3R* mutations, and the frequency of such mutations is significantly higher (~80%) in patients who experience leukemic transformation. SCN-associated *CSF3R* mutations occur in the region of the gene coding for the cytoplasmic domain of CSF3R and result in truncation of the C-terminal-negative regulatory domain. A different class of *CSF3R* mutations is noted in ~90% of patients with CNL; these are mostly membrane proximal, with the most frequent being a C-to-T substitution at nucleotide 1853 (T618I). About 40% of the T618I-mutated cases also harbored *SETBP1* mutations. *CSF3R* T618I induces a lethal myeloproliferative disorder in a mouse model and is associated with in vitro sensitivity to JAK inhibition.

Diagnosis Diagnosis of CNL requires exclusion of the more common causes of neutrophilia including infections and inflammatory processes. In addition, one should be mindful of the association between some forms of metastatic cancer or plasma cell neoplasms with secondary neutrophilia. Neoplastic neutrophilia also occurs in other myeloid malignancies including atypical chronic myeloid leukemia and chronic myelomonocytic leukemia. Accordingly, the WHO diagnostic criteria for CNL are designed to exclude the possibilities of both secondary/reactive neutrophilia and leukocytosis associated with myeloid malignancies other than CNL (Table 135e-5): leukocytosis ($\geq 25 \times 10^9/L$), >80% segmented/band neutrophils, <10% immature myeloid cells, <1% circulating blasts, and absence of dysgranulopoiesis or monocytosis. Bone marrow in CNL is hypercellular and displays increased number and percentage of neutrophils with a very high myeloid-to-erythroid ratio and minimal left shift, myeloid dysplasia, or reticulin fibrosis.

Treatment Current treatment in CNL is largely palliative and sub-optimal in its efficacy. Several drugs alone or in combination have been tried, and none have shown remarkable efficacy. As such, allogeneic stem cell transplantation (ASCT) is reasonable to consider in the presence of symptomatic disease, especially in younger patients. Otherwise, cytoreductive therapy with hydroxyurea is probably as

good as any treatment, and a more intensive combination chemotherapy may not have additional value. However, response to hydroxyurea therapy is often transient, and some have successfully used interferon α as an alternative drug. Response to treatment with ruxolitinib (a JAK1 and JAK2 inhibitor) has been reported but has not been confirmed.

ATYPICAL CHRONIC MYELOID LEUKEMIA

Atypical chronic myeloid leukemia, *BCR-ABL1* negative (aCML) is formally classified under the MDS/MPN category of myeloid malignancies and is characterized by left shifted granulocytosis and dysgranulopoiesis. The differential diagnosis of aCML includes chronic myeloid leukemia (CML), which is distinguished by the presence of *BCR-ABL1*; CNL, which is distinguished by the absence of dysgranulopoiesis and presence of *CSF3R* mutations; and chronic myelomonocytic leukemia, which is distinguished by the presence of monocytosis (absolute monocyte count $>1 \times 10^9/L$). The WHO diagnostic criteria for aCML are listed in Table 135e-5 and include granulocytosis (WBC $\geq 13 \times 10^9/L$), neutrophilia with dysgranulopoiesis, $\geq 10\%$ immature granulocytes, <20% peripheral blood myeloblasts, <10% peripheral blood monocytes, <2% basophils, and absence of otherwise specific mutations such as *BCR-ABL1*. The bone marrow is hypercellular with granulocyte proliferation and dysplasia with or without erythroid or megakaryocytic dysplasia.

The molecular pathogenesis of aCML is incompletely understood; about one-fourth of the patients express *SETBP1* mutations, which are, however, also found in several other myeloid malignancies, including CNL and chronic myelomonocytic leukemia. *SETBP1* mutations in aCML were prognostically detrimental and mostly located between codons 858 and 871; similar mutations are seen with Schinzel-Giedion syndrome (a congenital disease with severe developmental delay and various physical stigmata including midface retraction, large forehead, and macroglossia).

In a series of 55 patients with WHO-defined aCML, median age at diagnosis was 62 years with female preponderance (57%); splenomegaly was reported in 54% of the patients, red cell transfusion requirement in 65%, abnormal karyotype in 20% (20q- and trisomy 8 being the most frequent), and leukemic transformation in 40%. Median survival was 25 months. Outcome was worse in patients with marked

TABLE 135e-5 WORLD HEALTH ORGANIZATION DIAGNOSTIC CRITERIA FOR CHRONIC NEUTROPHILIC LEUKEMIA (CNL); ATYPICAL CHRONIC MYELOID LEUKEMIA, *BCR-ABL1* NEGATIVE (aCML); AND CHRONIC MYELOMONOCYTIC LEUKEMIA (CMML)

Variables	CNL	aCML	CMML
PB leukocyte count	$\geq 25 \times 10^9/L$	$\geq 13 \times 10^9/L$	
PB segmented neutrophils/bands	>80%		
PB immature granulocytes ^a	<10%	$\geq 10\%$	
PB blast count	<1%		
PB monocyte count	$<1 \times 10^9/L$	$<1 \times 10^9/L$	$>1 \times 10^9/L$
PB increased neutrophils or precursors with dysgranulopoiesis	No	Yes	
PB basophil percentage		<2%	
PB monocyte percentage		<10%	
Bone marrow	\uparrow Neutrophils, number and % <5% blasts Normal neutrophilic maturation Megakaryocytes normal or left shifted	\uparrow Granulocyte proliferation Granulocytic dysplasia \pm erythroid/ megakaryocyte dysplasia	Dysplasia in ≥ 1 myeloid lineages or Clonal cytogenetic/molecular abnormality
<i>BCR-ABL1</i>	No	No	No
<i>PDGFRA</i> , <i>PDGFRB</i> , or <i>FGFR1</i> mutation	No	No	No
PB and BM blasts/promonocytes	<20%	<20%	<20%
Hepatosplenomegaly	\pm	\pm	\pm
Evidence for other MDS/MPN	No	No	No
Evidence for other MPN	No	No	No
Evidence for reactive leukocytosis ^b or monocytosis	No	No	No

^aImmature granulocytes include myeloblasts, promyelocytes, myelocytes, and metamyelocytes. ^bCauses of reactive neutrophilia include plasma cell neoplasms, solid tumor, infections, and inflammatory processes.

Abbreviations: BM, bone marrow; MDS, myelodysplastic syndromes; MPN, myeloproliferative neoplasms; PB, peripheral blood.

leukocytosis, transfusion requirement, and increased immature cells in the peripheral blood. Conventional chemotherapy is largely ineffective in the treatment of aCML. However, a favorable experience with ASCT was reported in nine patients; after a median follow-up of 55 months, the majority of the patients remained in complete remission.

CHRONIC MYELOMONOCYTIC LEUKEMIA

Chronic myelomonocytic leukemia (CMML) is classified under the WHO category of MDS/MPN and is defined by an absolute monocyte count (AMC) of $>1 \times 10^9/L$ in the peripheral blood. Median age at diagnosis ranges between 65 and 75 years, and there is a 2:1 male predominance. Clinical presentation is variable and depends on whether the disease presents with MDS-like or MPN-like phenotype; the former is associated with cytopenias and the latter with splenomegaly and features of myeloproliferation such as fatigue, night sweats, weight loss, and cachexia. About 20% of patients with CMML experience serositis involving the joints (arthritis), pericardium (pericarditis and pericardial effusion), pleura (pleural effusion), or peritoneum (ascites).

Pathogenesis Clonal cytogenetic abnormalities are seen in about one-third of patients with CMML and include trisomy 8 and abnormalities of chromosome 7. Almost all patients with CMML harbor somatic mutations involving epigenetic regulator genes (e.g., *ASXL1*, *TET2*), spliceosome pathway genes (e.g., *SRSF2*), DNA damage response genes (e.g., *TP53*), and tyrosine kinases/transcription factors (e.g., *KRAS*, *NRAS*, *CBL*, and *RUNX1*). However, none of these mutations are specific to CMML, and their precise pathogenetic contribution is unclear.

Diagnosis Reactive monocytosis is uncommon but has been reported in association with certain infections and inflammatory conditions. Clonal (i.e., neoplastic) monocytosis defines CMML but is also seen with juvenile myelomonocytic leukemia and acute myeloid leukemia with monocytic differentiation. The WHO diagnostic criteria for CMML are listed in Table 135e-5 and include persistent AMC $>1 \times 10^9/L$, absence of *BCR-ABL1*, absence of the *PDGFRA* or *PDGFRB* mutations, $<20\%$ blasts and promonocytes in the peripheral blood and bone marrow, and dysplasia involving one or more myeloid lineages.

The bone marrow in CMML is hypercellular with granulocytic and monocytic proliferation. Dysplasia is often present and may involve one, two, or all myeloid lineages. On immunophenotyping, the abnormal cells often express myelomonocytic antigens such as CD13 and CD33, with variable expression of CD14, CD68, CD64, and CD163. Monocytic-derived cells are almost always positive for the cytochemical nonspecific esterases (e.g., butyrate esterase), whereas normal granulocytic precursors are positive for lysozyme and chloroacetate esterase. In CMML, it is common to have a hybrid cytochemical staining pattern with cells expressing both chloroacetate and butyrate esterases simultaneously (dual esterase staining).

Prognosis A meta-analysis showed median survival of 1.5 years in CMML. Numerous prognostic systems have attempted to better define and stratify the natural history of CMML. One of these, the Mayo prognostic model, assigns one point each to the following four independent prognostic variables: AMC $>10 \times 10^9/L$, presence of circulating immature cells, hemoglobin <10 g/dL, and platelet count $<100,000/mL$. This model stratified patients into three risk groups: low (0 points), intermediate (1 point), and high (≥ 2 points), translating to median survival times of 32, 18, and 10 months, respectively.

A French study incorporated *ASXL1* mutational status in 312 CMML patients. In a multivariable model, independent predictors of poor survival were WBC $>15 \times 10^9/L$ (3 points), *ASXL1* mutations (2 points), age >65 years (2 points), platelet count $<100,000/mL$ (2 points), and hemoglobin <10 g/dL in females and <11 g/dL in males (2 points). This model stratified patients into three groups: low (0–4 points), intermediate (5–7 points), and high risk (8–12 points), with median survival times of not reached, 38.5 months, and 14.4 months, respectively.

Treatment Current treatment consists of hydroxyurea and supportive care, including red cell transfusions and use of erythropoiesis-stimulating agents (ESAs). The value of hydroxyurea was reinforced by

a randomized trial against oral etoposide. No other single or combination chemotherapy has been shown to be superior to hydroxyurea. ASCT is a viable treatment option for transplant-eligible patients with poor prognostic features. Given the MDS/MPN overlap phenotype and the presence of MDS-like genetic/methylation abnormalities in CMML, hypomethylating agents such as 5-azacitidine and decitabine have been used with limited efficacy.

JUVENILE MYELOMONOCYTIC LEUKEMIA

Juvenile myelomonocytic leukemia (JMML) is primarily a disease of early childhood and is included, along with CMML, in the MDS/MPN WHO category. Both CMML and JMML feature leukocytosis, monocytosis, and hepatosplenomegaly. Additional characteristic features in JMML include thrombocytopenia and elevated fetal hemoglobin. Myeloid progenitors in JMML display granulocyte-macrophage colony-stimulating factor (GM-CSF) hypersensitivity that has been attributed to dysregulated RAS/MAPK signaling. The latter is believed to result from mutually exclusive mutations involving *RAS*, *PTPN11*, and *NF1*. A third of patients with JMML that is not associated with Noonan's syndrome carry *PTPN11* mutations, whereas the incidence of *NF1* in patients without neurofibromatosis type 1 and *RAS* mutations is approximately 15% each. Drug therapy is relatively ineffective in JMML, and the treatment of choice is ASCT, which results in a 5-year survival of approximately 50%.

MDS/MPN-U

The WHO classifies patients with morphologic and laboratory features that resemble both MDS and MPN as *MDS/MPN overlap*. This category includes CMML, aCML, and JMML, which have been described above. In addition, MDS/MPN includes a fourth category referred to as *MDS/MPN, unclassifiable* (MDS/MPN-U). Diagnosis of MDS/MPN-U requires the presence of both MDS and MPN features that are not adequate to classify patients as CMML, aCML, or JMML. MDS/MPN includes the provisional category of RARS-T.

RARS-T is classified in the MDS/MPN category because it shares dysplastic features with RARS and myeloproliferative features with essential thrombocythemia (ET). In one study, 111 patients with RARS-T were compared with 33 patients with RARS. The frequency of *SF3B1* mutations in RARS-T (87%) was similar to that in RARS (85%). *JAK2 V617F* mutation was detected in 49% of RARS-T patients (including 48% of those mutated for *SF3B1*) but none of those with RARS. In RARS-T, *SF3B1* mutations were more frequent in females (95%) than in males (77%), and mean ring sideroblast counts were higher in *SF3B1*-mutated patients. Median overall survival was 6.9 years in *SF3B1*-mutated patients versus 3.3 years in unmutated patients. Six-year survival was 67% in *JAK2*-mutated patients versus 32% in unmutated patients. Multivariable analysis identified younger age and *JAK2* and *SF3B1* mutations as favorable factors.

In one series, 85 patients with non-RARS-T MDS/MPN, median age was 70 years, and 72% were males. Splenomegaly at presentation was present in 33%, thrombocytosis in 13%, leukocytosis in 18%, *JAK2* mutations in 30%, and abnormal karyotype in 51%; the most frequent cytogenetic abnormality was trisomy 8. Median survival was 12.4 months and favorably affected by thrombocytosis. Treatment with hypomethylating agents, immunomodulators, or ASCT did not appear to favorably affect survival.

MYELOPROLIFERATIVE NEOPLASM, UNCLASSIFIABLE (MPN-U)

The category of MPN-U includes MPN-like neoplasms that cannot be clearly classified as one of the other seven subcategories of MPN (Table 135e-4). Examples include patients presenting with unusual thrombocytosis or unexplained organomegaly with normal blood counts but found to carry MPN-characteristic mutations such as *JAK2* and *CALR* or display bone marrow morphology that is consistent with MPN. It is possible that some cases of MPN-U represent earlier disease stages in polycythemia vera (PV) or ET that fail to meet the threshold hemoglobin levels (18.5 g/dL in men or 16.5 g/dL in women) or platelet counts ($450 \times 10^9/L$) that are required by the WHO diagnostic criteria. Specific treatment interventions might not be necessary in asymptomatic patients

TABLE 135e-6 DIAGNOSIS OF CHRONIC EOSINOPHILIC LEUKEMIA AND HYPEREOSINOPHILIC SYNDROME

Required: Persistent eosinophilia $\geq 1500/\mu\text{L}$ in blood, increased marrow eosinophils, and myeloblasts $<20\%$ in blood or marrow.

1. Exclude all causes of reactive eosinophilia: allergy, parasites, infection, pulmonary disease (e.g., hypersensitivity pneumonitis, Loeffler's), and collagen vascular diseases
2. Exclude primary neoplasms associated with secondary eosinophilia: T-cell lymphomas, Hodgkin's disease, acute lymphoid leukemia, mastocytosis
3. Exclude other primary myeloid neoplasms that may involve eosinophils: chronic myeloid leukemia, acute myeloid leukemia with inv(16) or t(16;16) (p13;q22), other myeloproliferative syndromes, and myelodysplasia
4. Exclude T-cell reaction with increased interleukin 5 or other cytokine production

If these entities have been excluded and no evidence documents a clonal myeloid disorder, the diagnosis is hypereosinophilic syndrome.

If these entities have been excluded and the myeloid cells show a clonal chromosome abnormality or some other evidence of clonality and blast cells are present in the peripheral blood ($>2\%$) or are increased in the marrow (but $<20\%$), the diagnosis is chronic eosinophilic leukemia.

with MPN-U, whereas patients with arterial thrombotic complications might require cyto-reductive and aspirin therapy and those with venous thrombosis might require systemic anticoagulation.

TRANSIENT MYELOPROLIFERATIVE DISORDER (TMD)

TMD constitutes an often but not always transient phenomenon of abnormal megakaryoblast proliferation, which occurs in approximately 10% of infants with Down's syndrome. TMD is usually recognized at birth and either undergoes spontaneous regression (75% of cases) or progresses into acute megakaryoblastic leukemia (AMKL) (25% of cases). Almost all patients with TMD and TMD-derived AMKL display somatic *GATA1* mutations. TMD-associated *GATA1* mutations constitute exon 2 insertions, deletions, or missense mutations, affecting the N-terminal transactivation domain of GATA-1, and result in loss of full-length (50-kDa) GATA-1 and its replacement with a shorter isoform (40-kDa) that retains friend of GATA-1 (FOG-1) binding. In contrast, inherited forms of exon 2 *GATA1* mutations produce a phenotype with anemia, whereas exon 4 mutations that affect the N-terminal, FOG-1-interactive domain produce familial dyserythropoietic anemia with thrombocytopenia or X-linked macrothrombocytopenia.

EOSINOPHILIC DISORDERS

Eosinophilia refers to a peripheral blood absolute eosinophil count (AEC) that is above the upper normal limit of the reference range. The term *hypereosinophilia* is used when the AEC is $>1500 \times 10^9/\text{L}$. Eosinophilia is operationally classified as secondary (nonneoplastic proliferation of eosinophils) and primary (proliferation of eosinophils that is either neoplastic or otherwise unexplained) (Table 135e-6). Secondary eosinophilia is by far the most frequent cause of eosinophilia and is often associated with infections, especially those related to tissue-invasive helminths; allergic/vasculitic diseases; drugs; and metastatic cancer. Primary eosinophilia is the focus of this chapter and

is considered when a cause for secondary eosinophilia is not readily apparent.

Primary Eosinophilia Primary eosinophilia is classified as clonal or idiopathic. Diagnosis of clonal eosinophilia requires morphologic, cytogenetic, or molecular evidence of a myeloid neoplasm. Idiopathic eosinophilia is considered when both secondary and clonal eosinophilias have been ruled out as a possibility. HES is a subcategory of idiopathic eosinophilia with persistent AEC of $\geq 1.5 \times 10^9/\text{L}$ and associated with eosinophil-mediated organ damage (Table 135e-7). An HES-like disorder that is associated with clonal or phenotypically abnormal T cells is referred to as *lymphocytic variant hypereosinophilia* (Table 135e-7).

Clonal Eosinophilia Examples of clonal eosinophilia include eosinophilia associated with acute myeloid leukemia (AML), MDS, CML, mastocytosis, and MDS/MPN overlap. Myeloid neoplasm-associated eosinophilia also includes the WHO MPN subcategory of chronic eosinophilic leukemia, not otherwise specified (CEL-NOS) and the WHO myeloid malignancy subcategory referred to as myeloid/lymphoid neoplasms with eosinophilia and mutations involving platelet-derived growth factor receptor (*PDGFR*) α/β or fibroblast growth factor receptor 1 (*FGFR1*).

The diagnostic workup for clonal eosinophilia that is not associated with morphologically overt myeloid malignancy should start with peripheral blood mutation screening for *FIP1L1-PDGFR α* and *PDGFR β* mutations using fluorescence in situ hybridization (FISH) or reverse transcription polymerase chain reaction. This is crucial because such eosinophilia is easily treated with imatinib. If mutation screening is negative, a bone marrow examination with cytogenetic studies is indicated. In this regard, one must first pay attention to the presence or absence of 5q33, 4q12, or 8p11.2 translocations, which, if present, would suggest *PDGFR β* -, *PDGFR α* -, or *FGFR1*-rearranged clonal eosinophilia, respectively. The presence of 5q33 or 4q12 translocations predicts favorable response to treatment with imatinib mesylate, whereas 8p11.2 translocations are associated with aggressive myeloid malignancies that are refractory to current drug therapy.

CEL-NOS is considered in the presence of cytogenetic/morphologic evidence of a myeloid malignancy that is otherwise not classifiable. Specifically, CEL-NOS is distinguished from HES by the presence of either a cytogenetic abnormality or greater than 2% peripheral blood blasts or greater than 5% bone marrow blasts (Table 135e-7). HES or idiopathic eosinophilia is considered in the absence of both morphologic and molecular evidence of clonal eosinophilia. However, before making a working diagnosis of HES, one has to exclude lymphocytic variant hypereosinophilia by excluding the presence of phenotypically abnormal T lymphocytes (by flow cytometry) and clonal T-cell gene rearrangements.

Chronic Eosinophilic Leukemia, Not Otherwise Specified (CEL-NOS) CEL-NOS is a subset of clonal eosinophilia that is neither molecularly defined nor classified as an alternative clinicopathologically assigned myeloid malignancy. We prefer to use the term strictly in patients with an HES phenotype who also display either a clonal cytogenetic/molecular abnormality or excess blasts in the bone marrow or peripheral blood. The WHO defines CEL-NOS in the presence of an AEC $\geq 1.5 \times 10^9/\text{L}$

TABLE 135e-7 PRIMARY EOSINOPHILIA CLASSIFICATION

Variables	<i>PDGFRα</i> -, <i>PDGFRβ</i> -, or <i>FGFR1</i> -Mutated Eosinophilia	Chronic Eosinophilia, Not Otherwise Specified	Lymphocytic Variant Hypereosinophilia	Hypereosinophilic Syndrome
Absolute eosinophil count	$>600 \times 10^9/\text{L}$	$>1500 \times 10^9/\text{L}$	$>1500 \times 10^9/\text{L}$	$>1500 \times 10^9/\text{L}$
Peripheral blood blast $>2\%$	Yes or no	Yes or no	No	No
Bone marrow blast $>5\%$	Yes or no	Yes or No	No	No
Abnormal karyotype	Yes or no	Yes or no	No	No
<i>PDGFRα</i> , <i>PDGFRβ</i> , or <i>FGFR1</i> mutation	Yes	No	No	No
<i>BCR-ABL1</i>	No	No	No	No
Abnormal T lymphocyte phenotype or clonal T-cell clones	No	No	Yes	No
Eosinophil-mediated tissue damage	Yes or no	Yes or no	Yes or no	Yes

that is accompanied by either the presence of myeloblast excess (either >2% in the peripheral blood or 5–19% in the bone marrow) or evidence of myeloid clonality. Cytogenetic abnormalities in CEL, other than those that are associated with molecularly defined eosinophilic disorders, include trisomy 8 (the most frequent), t(10;11)(p14;q21), and t(7;12)(q11;p11). CEL-NOS does not respond to imatinib, and treatment strategies are often not different from those used in other similar MPNs and include ASCT for transplant-eligible patients with poor risk factors and participation in experimental treatment protocols otherwise.

PDGFR-Mutated Eosinophilia Both platelet-derived growth factor receptors α (*PDGFRA* located on chromosome 4q12) and β (*PDGFRB* located on chromosome 5q31-q32) are involved in MPN-relevant activating mutations. Clinical phenotype in both instances includes prominent blood eosinophilia and excellent response to imatinib therapy. In regard to *PDGFRA* mutations, the most popular is *FIP1L1-PDGFR*, a karyotypically occult del(4)(q12) that was described in 2003 as an imatinib-sensitive activating mutation. Functional studies have demonstrated transforming properties in cell lines and the induction of MPN in mice. Cloning of the *FIP1L1-PDGFR* fusion gene identified a novel molecular mechanism for generating this constitutively active fusion tyrosine kinase, wherein a ~800-kb interstitial deletion within 4q12 fuses the 5' portion of *FIP1L1* to the 3' portion of *PDGFRA*. *FIP1L1-PDGFR* occurs in a very small subset of patients who present with the phenotypic features of either systemic mastocytosis or HES, but the presence of the mutation reliably predicts complete hematologic and molecular response to imatinib therapy.

The association between eosinophilic myeloid malignancies and *PDGFRB* rearrangement was first characterized and published in 1994 when fusion of the tyrosine kinase–encoding region of *PDGFRB* to the *ets*-like gene, *ETV6* [*ETV6-PDGFRB*, t(5;12)(q33;p13)] was demonstrated. The fusion protein was transforming to cell lines and resulted in constitutive activation of *PDGFRB* signaling. Since then, several other *PDGFRB* fusion transcripts with similar disease phenotypes have been described, cell line transformation and myeloproliferative disease (MPD) induction in mice has been demonstrated, and imatinib therapy was proven effective when used.

FGFR1-Mutated Eosinophilia The 8p11 myeloproliferative syndrome (EMS) (also known as human stem cell leukemic/lymphoma syndrome) constitutes a clinical phenotype with features of both lymphoma and eosinophilic MPN and characterized by a fusion mutation that involves the gene for fibroblast growth factor receptor 1 (*FGFR1*), which is located on chromosome 8p11. In EMS, both myeloid and lymphoid lineage cells exhibit the 8p11 translocation, thus demonstrating the stem cell origin of the disease. The disease features several 8p11-linked chromosome translocations, and some of the corresponding fusion *FGFR1* mutants have been shown to transform cell lines and induce EMS- or CML-like disease in mice depending on the specific *FGFR1* partner gene (*ZNF198* or *BCR*, respectively). Consistent with this laboratory observation, some patients with *BCR-FGFR1* mutation manifest a more indolent CML-like disease. The mechanism of *FGFR1* activation in EMS is similar to that seen with *PDGFRB*-associated MPD; the tyrosine kinase domain of *FGFR1* is juxtaposed to a dimerization domain from the partner gene. EMS is aggressive and requires combination chemotherapy followed by ASCT.

Hypereosinophilic Syndrome (HES) Blood eosinophilia that is neither secondary nor clonal is operationally labeled as being idiopathic. HES is a subcategory of idiopathic eosinophilia with persistent increase of the AEC to $\geq 1.5 \times 10^9/L$ and presence of eosinophil-mediated organ damage, including cardiomyopathy, gastroenteritis, cutaneous lesions, sinusitis, pneumonitis, neuritis, and vasculitis. In addition, some patients manifest thromboembolic complications, hepatosplenomegaly, and either cytopenia or cytosis.

Bone marrow histologic and cytogenetic/molecular studies should be examined before a working diagnosis of HES is made. Additional blood studies that are currently recommended during the evaluation of HES include serum tryptase (an increased level suggests systemic

mastocytosis and warrants molecular studies to detect *FIP1L1-PDGFR*), T-cell immunophenotyping, and T-cell receptor antigen gene rearrangement analysis (a positive test suggests an underlying clonal or phenotypically abnormal T-cell disorder). In addition, initial evaluation in HES should include echocardiogram and measurement of serum troponin levels to screen for myocardial involvement by the disease.

Initial evaluation of the patient with eosinophilia should include tests that facilitate assessment of target organ damage, including complete blood count, chest x-ray, echocardiogram, and serum troponin level. An increased level of serum cardiac troponin has been shown to correlate with the presence of cardiomyopathy in HES. Typical echocardiographic findings in HES include ventricular apical thrombus, posterior mitral leaflet or tricuspid valve abnormality, endocardial thickening, dilated left ventricle, and pericardial effusion.

Glucocorticoids are the cornerstone of therapy in HES. Treatment with oral prednisone is usually started at 1 mg/kg per day and continued for 1–2 weeks before the dose is tapered slowly over the ensuing 2–3 months. If symptoms recur at a prednisone dose level of >10 mg/d, either hydroxyurea or interferon α is used as steroid-sparing agent. In patients who do not respond to usual therapy as outlined above, mepolizumab or alemtuzumab might be considered. Mepolizumab targets interleukin 5 (IL-5), a well-recognized survival factor for eosinophils. Alemtuzumab targets the CD52 antigen, which has been shown to be expressed by eosinophils but not by neutrophils.

MASTOCYTOSIS

Mast cell disease (MCD) is defined as tissue infiltration by morphologically and immunophenotypically abnormal mast cells. MCD is classified into two broad categories: cutaneous mastocytosis and systemic mastocytosis (SM). MCD in adults is usually systemic, and the clinical course can be either indolent or aggressive, depending on the respective absence or presence of impaired organ function. Symptoms and signs of MCD include urticaria pigmentosa, mast cell mediator release symptoms (e.g., headache, flushing, lightheadedness, syncope, anaphylaxis, pruritus, urticaria, angioedema, nausea, diarrhea, abdominal cramps), and organ damage (lytic bone lesions, osteoporosis, hepatosplenomegaly, cytopenia). Aggressive SM can be associated with another myeloid malignancy, including MPN, MDS, or MDS/MPN overlap (e.g., CMML), or present as overt mast cell leukemia. In general, life expectancy is near normal in indolent SM but significantly shortened in aggressive SM.

Diagnosis of SM is based on bone marrow examination that shows clusters of morphologically abnormal, spindle-shaped mast cells that are best evaluated by the use of immunohistochemical stains that are specific to mast cells (tryptase, CD117). In addition, mast cell immunophenotyping reveals aberrant CD25 expression by neoplastic mast cells. Other laboratory findings in SM include increased levels of serum tryptase, histamine and urine histamine metabolites, and prostaglandins. SM is associated with *KIT* mutations, usually *KIT* D816V, in the majority of patients. Accordingly, mutation screening for *KIT* D816V is diagnostically useful. However, the ability to detect *KIT* D816V depends on assay sensitivity and mast cell content of the test sample.

Both indolent and aggressive SM patients might experience mast cell mediator release symptoms, which are usually managed by both H_1 and H_2 histamine receptor blockers as well as cromolyn sodium. In addition, patients with propensity to vasodilatory shock should wear a medical alert bracelet and carry an Epi-Pen self-injector for self-administration of subcutaneous epinephrine. Urticaria pigmentosa shows variable response to both topical and systemic glucocorticoid therapy. Cytoreductive therapy is not recommended for indolent SM. In aggressive SM, either interferon α or cladribine is considered first-line therapy and benefits the majority of patients. In contrast, imatinib is ineffective in the treatment of *PDGFR*-unmutated SM.

DENDRITIC AND HISTIOCYTIC NEOPLASMS

Dendritic cell (DC) and histiocyte/macrophage neoplasms are extremely rare. DCs are antigen-presenting cells, whereas histiocyte/

macrophages are antigen-processing cells. Bone marrow myeloid stem cells (CD34+) give rise to monocyte (CD14+, CD68+, CD11c+, CD1a-) and DC (CD14-, CD11c+/-, CD1a+/c) precursors. Monocyte precursors, in turn, give rise to macrophages (CD14+, CD68+, CD11c+, CD163+, lysozyme+) and interstitial DCs (CD68+, CD1a-). DC precursors give rise to Langerhans cell DCs (Birbeck granules, CD1a+, S100+, langerin+) and plasmacytoid DCs (CD68+, CD123+). Follicular DCs (CD21+, CD23+, CD35+) originate from mesenchymal stem cells. Dendritic and histiocytic neoplasms are operationally classified into macrophage/histiocyte-related and DC-related neoplasms. The former includes histiocytic sarcoma/malignant histiocytosis and the latter Langerhans cell histiocytosis, Langerhans cell sarcoma, interdigitating DC sarcoma, and follicular DC sarcoma.

Histiocytic Sarcoma/Malignant Histiocytosis Histiocytic sarcoma represents malignant proliferation of mature tissue histiocytes and is often localized. Median age at diagnosis is estimated at 46 years with slight male predilection. Some patients might have history of lymphoma, MDS, or germ cell tumors at time of disease presentation. The three typical disease sites are lymph nodes, skin, and the gastrointestinal system. Patients may or may not have systemic symptoms including fever and weight loss, and other symptoms include hepatosplenomegaly, lytic bone lesions, and pancytopenia. Immunophenotype includes presence of histiocytic markers (CD68, lysozyme, CD11c, CD14) and absence of myeloid or lymphoid markers. Prognosis is poor, and treatment is often ineffective. The term malignant histiocytosis refers to a disseminated disease and systemic symptoms. Lymphoma-like treatment induces complete remissions in some patients, and median survival is estimated at 2 years.

Langerhans Cell Histiocytosis Langerhans cells (LCs) are specialized DCs that reside in mucocutaneous tissue and upon activation become specialized for antigen presentation to T cells. LC histiocytosis (LCH; also known as histiocytosis X) represents neoplastic proliferation of LCs (S-100+, CD1a+, and Birbeck granules on electron microscopy). LCH incidence is estimated at 5 per million, and the disease typically affects children with a male predilection. Presentation can be either unifocal (eosinophilic granuloma) or multifocal. The former usually affects bones and less frequently lymph nodes, skin, and lung, whereas the latter is more disseminated. Unifocal disease often affects older children and adults, whereas multisystem disease affects infants. LCH of the lung in adults is characterized by bilateral nodules. Prognosis

depends on organs involved. Only 10% of patients progress from unifocal to multiorgan disease. LCH of the lung might improve upon cessation of smoking.

Langerhans Cell Sarcoma Langerhans cell sarcoma (LCS) also represents neoplastic proliferation of LCs with overtly malignant morphology. The disease can present de novo or progress from antecedent LCH. There is a female predilection, and median age at diagnosis is estimated at 41 years. Immunophenotype is similar to that seen in LCH, and liver, spleen, lung, and bone are the usual sites of disease. Prognosis is poor, and treatment is generally ineffective.

Interdigitating Dendritic Cell Sarcoma Interdigitating DC sarcoma (IDCS), also known as reticulum cell sarcoma, represents neoplastic proliferation of interdigitating DCs. The disease is extremely rare and affects elderly adults with no sex predilection. Typical presentation is asymptomatic solitary lymphadenopathy. Immunophenotype includes S-100+ and negative for vimentin and CD1a. Prognosis ranges from benign local disease to widespread lethal disease.

Follicular Dendritic Cell Neoplasm Follicular DCs (FDCs) reside in B-cell follicles and present antigen to B cells. FDC neoplasms (FDCNs) are usually localized and often affect adults. FDCN might be associated with Castleman's disease in 10–20% of cases, and increased incidence in schizophrenia has been reported. Cervical lymph nodes are the most frequent site of involvement in FDCN, and other sites include maxillary, mediastinal, and retroperitoneal lymph nodes; oral cavity; gastrointestinal system; skin; and breast. Sites of metastasis include lung and liver. Immunophenotype includes CD21, CD35, and CD23. Clinical course is typically indolent, and treatment includes surgical excision followed by regional radiotherapy and sometimes systemic chemotherapy.

Hemophagocytic Syndromes Hemophagocytic syndrome (HPS) represents nonneoplastic proliferation and activation of macrophages that induce cytokine-mediated bone marrow suppression and features of intense phagocytosis in bone marrow and liver. HPS may result from genetic or acquired disorders of macrophages. The former entail genetically determined inability to regulate macrophage proliferation and activation. Acquired HPS is often precipitated by viral infections, most notably Epstein-Barr virus. HPS might also accompany certain malignancies such as T-cell lymphoma. Clinical course is often fulminant and fatal.

136 Plasma Cell Disorders

Nikhil C. Munshi, Dan L. Longo, Kenneth C. Anderson

The *plasma cell disorders* are monoclonal neoplasms related to each other by virtue of their development from common progenitors in the B-lymphocyte lineage. Multiple myeloma, Waldenström's macroglobulinemia, primary amyloidosis (**Chap. 137**), and the heavy chain diseases comprise this group and may be designated by a variety of synonyms such as *monoclonal gammopathies*, *paraproteinemias*, *plasma cell dyscrasias*, and *dysproteinemias*. Mature B lymphocytes destined to produce IgG bear surface immunoglobulin molecules of both M and G heavy chain isotypes with both isotypes having identical idiotypes (variable regions). Under normal circumstances, maturation to antibody-secreting plasma cells and their proliferation is stimulated by exposure to the antigen for which the surface immunoglobulin is specific; however, in the plasma cell disorders, the control over this process is lost. The clinical manifestations of all the plasma cell disorders relate to the expansion of the neoplastic cells, to the secretion of cell products (immunoglobulin molecules or subunits, lymphokines), and to some extent to the host's response to the tumor. **Normal development of B lymphocytes is discussed in Chap. 372e and depicted in Fig. 134-2.**

There are three categories of structural variation among immunoglobulin molecules that form antigenic determinants, and these are used to classify immunoglobulins. *Isotypes* are those determinants that distinguish among the main classes of antibodies of a given species and are the same in all normal individuals of that species. Therefore, isotypic determinants are, by definition, recognized by antibodies from a distinct species (heterologous sera) but not by antibodies from the same species (homologous sera). There are five heavy chain isotypes (M, G, A, D, E) and two light chain isotypes (κ , λ). *Allotypes* are distinct determinants that reflect regular small differences between individuals of the same species in the amino acid sequences of otherwise similar immunoglobulins. These differences are determined by allelic genes; by definition, they are detected by antibodies made in the same species. *Idiotypes* are the third category of antigenic determinants. They are unique to the molecules produced by a given clone of antibody-producing cells. Idiotypes are formed by the unique structure of the antigen-binding portion of the molecule.

Antibody molecules (**Fig. 136-1**) are composed of two heavy chains (~50,000 mol wt) and two light chains (~25,000 mol wt). Each chain has a constant portion (limited amino acid sequence variability) and a variable region (extensive sequence variability). The light and heavy chains are linked by disulfide bonds and are aligned so that their variable regions are adjacent to one another. This variable region forms the antigen recognition site of the antibody molecule; its unique structural features form idiotypes that are reliable markers for a particular clone of cells because each antibody is formed and secreted by a single clone. Because of the mechanics of the gene rearrangements necessary to specify the immunoglobulin variable regions (VDJ joining for the heavy chain, VJ joining for the light chain), a particular clone rearranges only one of the two chromosomes to produce an immunoglobulin molecule of only one light chain isotype and only one allotype (allelic exclusion) (**Fig. 136-1**). After exposure to antigen, the variable region may become associated with a new heavy chain isotype (class switch). Each clone of cells performs these sequential gene arrangements in a unique way. This results in each clone producing a unique immunoglobulin molecule. In most plasma cells, light chains are synthesized in slight excess, secreted as free light chains, and cleared by the kidney, but <10 mg of such light chains is excreted per day.

Electrophoretic analysis permits separation of components of the serum proteins (**Fig. 136-2**). The immunoglobulins move heterogeneously in an electric field and form a broad peak in the gamma region, which is usually increased in the sera of patients with plasma cell tumors. There is a sharp spike in this region called an *M component* (M for monoclonal). Less commonly, the M component may appear in the

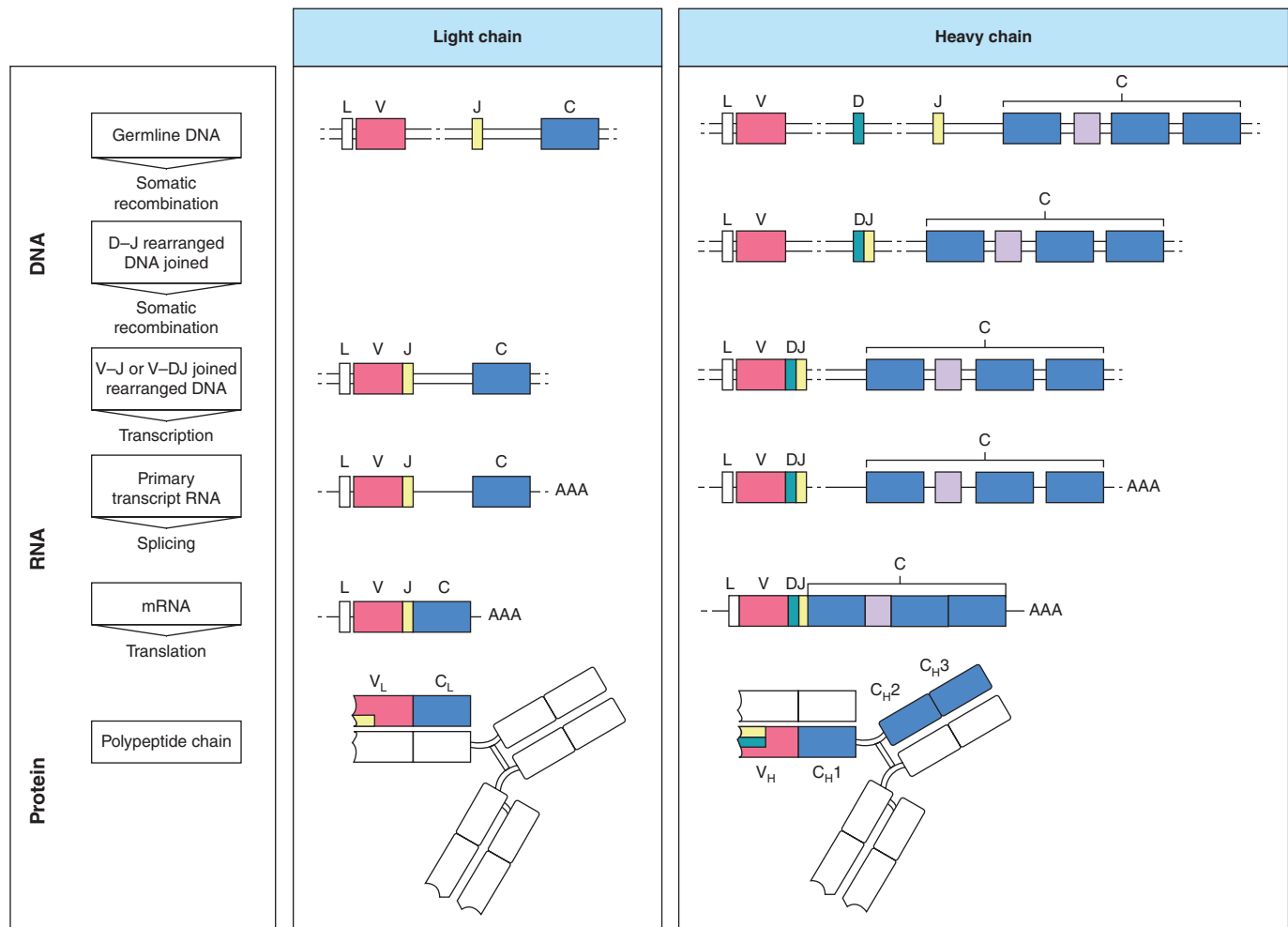
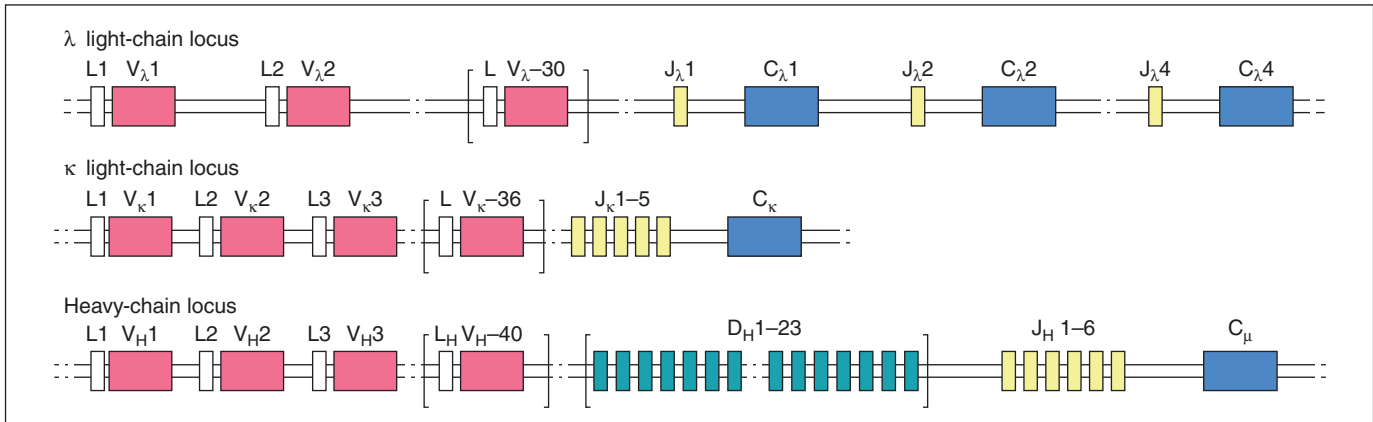


FIGURE 136-1 Immunoglobulin genetics and the relationship of gene segments to the antibody protein. The *top* portion of the figure is a schematic of the organization of the immunoglobulin genes, λ on chromosome 22, κ on chromosome 2, and the heavy chain locus on chromosome 14. The heavy chain locus is longer than 2 megabases, and some of the D region gene segments are only a few bases long, so the figure depicts the schematic relationship among the segments, not their actual size. The *bottom* portion of the figure outlines the steps in going from the noncontiguous germline gene segments to an intact antibody molecule. Two recombination events juxtapose the V-D-J (or V-J for light chains) segments. The rearranged gene is transcribed, and RNA splicing cuts out intervening sequences to produce an mRNA, which is then translated into an antibody light or heavy chain. The sites on the antibody that bind to antigen (the so called CDR3 regions) are encoded by D and J segments for heavy chains and the J segments for light chains. (From K Murphy: *Janeway's Immunobiology*, 8th ed. Garland Science, 2011.)

β_2 or α_2 globulin region. The monoclonal antibody must be present at a concentration of at least 5 g/L (0.5 g/dL) to be accurately quantitated by this method. This corresponds to $\sim 10^9$ cells producing the antibody. Confirmation of the type of immunoglobulin and that it is truly monoclonal is determined by immunoelectrophoresis that reveals a single heavy and/or light chain type. Hence immunoelectrophoresis

and electrophoresis provide qualitative and quantitative assessment of the M component, respectively. Once the presence of an M component has been confirmed, the amount of M component in the serum is a reliable measure of the tumor burden, making M component an excellent tumor marker to manage therapy, yet it is not specific enough to be used to screen asymptomatic patients. In addition to the plasma

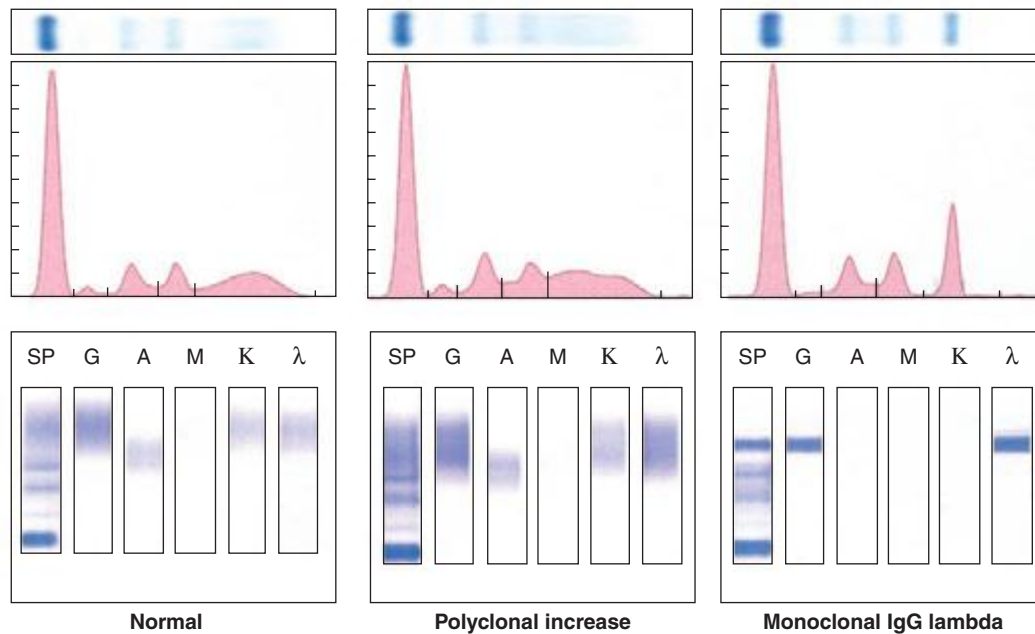


FIGURE 136-2 Representative patterns of serum electrophoresis and immunofixation. The upper panels represent agarose gel, middle panels are the densitometric tracing of the gel, and lower panels are immunofixation patterns. Panel on the left illustrates the normal pattern of serum protein on electrophoresis. Because there are many different immunoglobulins in the serum, their differing mobilities in an electric field produce a broad peak. In conditions associated with increases in polyclonal immunoglobulin, the broad peak is more prominent (middle panel). In monoclonal gammopathies, the predominance of a product of a single cell produces a “church spire” sharp peak, usually in the γ globulin region (right panel). The immunofixation (lower panel) identifies the type of immunoglobulin. For example, normal and polyclonal increase in immunoglobulins produce no distinct bands; however, the right panel shows distinct bands in IgG and lambda protein lanes, confirming the presence of IgG lambda monoclonal protein. (Courtesy of Dr. Neal I. Lindeman; with permission.)

cell disorders, M components may be detected in other lymphoid neoplasms such as chronic lymphocytic leukemia and lymphomas of B- or T-cell origin; nonlymphoid neoplasms such as chronic myeloid leukemia, breast cancer, and colon cancer; a variety of nonneoplastic conditions such as cirrhosis, sarcoidosis, parasitic diseases, Gaucher’s disease, and pyoderma gangrenosum; and a number of autoimmune conditions, including rheumatoid arthritis, myasthenia gravis, and cold agglutinin disease. Monoclonal proteins are also observed in immunosuppressed patients after organ transplant and, rarely, allogeneic transplant. At least two very rare skin diseases—lichen myxedematosus (also known as papular mucinosis) and necrobiotic xanthogranuloma—are associated with a monoclonal gammopathy. In papular mucinosis, highly cationic IgG is deposited in the dermis of patients. This organ specificity may reflect the specificity of the antibody for some antigenic component of the dermis. Necrobiotic xanthogranuloma is a histiocytic infiltration of the skin, usually of the face, that produces red or yellow nodules that can enlarge to plaques. Approximately 10% progress to myeloma. Five percent of patients with sensory motor neuropathy also have a monoclonal paraprotein.

The nature of the M component is variable in plasma cell disorders. It may be an intact antibody molecule of any heavy chain subclass, or it may be an altered antibody or fragment. Isolated light or heavy chains may be produced. In some plasma cell tumors such as extramedullary or solitary bone plasmacytomas, less than one-third of patients will have an M component. In ~20% of myelomas, only light chains are produced and, in most cases, are secreted in the urine as Bence Jones proteins. The frequency of myelomas of a particular heavy chain class is roughly proportional to the serum concentration, and therefore, IgG myelomas are more common than IgA and IgD myelomas. In approximately 1% of patients with myeloma, biclonal or triclonal gammopathy is observed.

MULTIPLE MYELOMA

DEFINITION

Multiple myeloma represents a malignant proliferation of plasma cells derived from a single clone. The tumor, its products, and the host

response to it result in a number of organ dysfunctions and symptoms, including bone pain or fracture, renal failure, susceptibility to infection, anemia, hypercalcemia, and occasionally clotting abnormalities, neurologic symptoms, and manifestations of hyperviscosity.

ETIOLOGY

The cause of myeloma is not known. Myeloma occurred with increased frequency in those exposed to the radiation of nuclear warheads in World War II after a 20-year latency. Myeloma has been seen more commonly than expected among farmers, wood workers, leather workers, and those exposed to petroleum products. A variety of chromosomal alterations have been found in patients with myeloma: hyperdiploidy, 13q14 deletions, translocations t(11;14)(q13;q32), t(4;14)(p16;q32), and t(14;16), and 17p13 deletions. Evidence is strong that errors in switch recombination—the genetic mechanism to change antibody heavy chain isotype—participate in the transformation process. However, no common molecular pathogenetic pathway has yet emerged. Genome sequencing studies have failed to identify any recurrent mutation with frequency >20%; *N-ras*, *K-ras*, and *B-raf* mutations are most common and combined occur in over 40% of patients. There is also evidence of complex clusters of subclonal variants at diagnosis that acquire additional mutations over time, indicative of genomic evolution that may drive disease progression. The neoplastic event in myeloma may involve cells earlier in B-cell differentiation than the plasma cell. Interleukin (IL) 6 may play a role in driving myeloma cell proliferation. It remains difficult to distinguish benign from malignant plasma cells based on morphologic criteria in all but a few cases (Fig. 136-3).

INCIDENCE AND PREVALENCE

An estimated 24,050 new cases of myeloma were diagnosed in 2014, and 11,090 people died from the disease in the United States. Myeloma increases in incidence with age. The median age at diagnosis is 70 years; it is uncommon under age 40. Males are more commonly affected than females, and blacks have nearly twice the incidence of whites. Myeloma accounts for 1.3% of all malignancies in whites and 2% in blacks, and 13% of all hematologic cancers in whites and 33% in blacks.

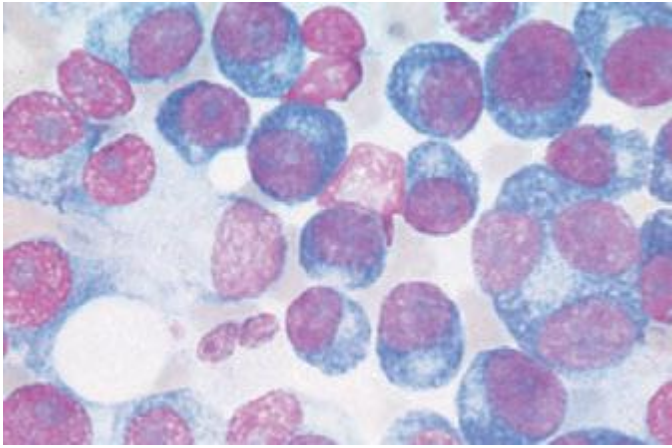


FIGURE 136-3 Multiple myeloma (marrow). The cells bear characteristic morphologic features of plasma cells, round or oval cells with an eccentric nucleus composed of coarsely clumped chromatin, a densely basophilic cytoplasm, and a perinuclear clear zone containing the Golgi apparatus. Binucleate and multinucleate malignant plasma cells can be seen.

GLOBAL CONSIDERATIONS



The incidence of myeloma is highest in African Americans and Pacific Islanders; intermediate in Europeans and North American whites; and lowest in people from developing countries including Asia. The higher incidence in more developed countries may result from the combination of a longer life expectancy and more frequent medical surveillance. Incidence of multiple myeloma in other ethnic groups including native Hawaiians, female Hispanics, American Indians from New Mexico, and Alaskan natives is higher relative to U.S. whites in the same geographic area. Chinese and Japanese populations have a lower incidence than whites. Immunoproliferative small-intestinal disease with alpha heavy chain disease is most prevalent in the Mediterranean area. Despite these differences in prevalence, the characteristics, response to therapy, and prognosis of myeloma are similar worldwide.

PATHOGENESIS AND CLINICAL MANIFESTATIONS

Multiple myeloma (MM) cells bind via cell-surface adhesion molecules to bone marrow stromal cells (BMSCs) and extracellular matrix (ECM), which triggers MM cell growth, survival, drug resistance, and migration in the bone marrow milieu (Fig. 136-4). These effects are due both to direct MM cell–BMSC binding and to induction of various cytokines, including IL-6, insulin-like growth factor type I (IGF-I), vascular endothelial growth factor (VEGF), and stromal cell–derived growth factor (SDF)-1 α . Growth, drug resistance, and migration are mediated via Ras/Raf/mitogen-activated protein kinase, PI3K/Akt, and protein kinase C signaling cascades, respectively.

Bone pain is the most common symptom in myeloma, affecting nearly 70% of patients. Unlike the pain of metastatic carcinoma, which often is worse at night, the pain of myeloma is precipitated by movement. Persistent localized pain in a patient with myeloma usually signifies a pathologic fracture. The bone lesions of myeloma are caused by the proliferation of tumor cells, activation of osteoclasts that destroy bone, and suppression of osteoblasts that form new bone. The increased osteoclast activity is mediated by osteoclast activating factors (OAFs) made by the myeloma cells (OAF activity can be mediated by several cytokines, including IL-1, lymphotoxin, VEGF, receptor activator of NF- κ B [RANK] ligand, macrophage inhibitory factor [MIP]-1 α , and tumor necrosis factor [TNF]). The bone lesions are lytic in nature and are rarely associated with osteoblastic new bone formation due to their suppression by dickhoff-1 (DKK-1) produced by myeloma cells. Therefore, radioisotopic bone scanning is less useful in diagnosis than is plain radiography. The bony lysis results in substantial mobilization of calcium from bone, and serious acute and chronic complications of hypercalcemia may dominate the clinical picture (see below). Localized bone lesions may expand to the point that mass lesions may be palpated, especially on the skull (Fig. 136-5), clavicles, and sternum; and the collapse of vertebrae may lead to spinal cord compression. The next most common clinical problem in patients with myeloma is susceptibility to bacterial infections. The most common infections are pneumonias and pyelonephritis, and the most frequent pathogens are *Streptococcus pneumoniae*, *Staphylococcus aureus*, and *Klebsiella pneumoniae* in the lungs and *Escherichia coli* and other gram-negative organisms in the urinary tract. In ~25% of patients, recurrent infections are the presenting features, and >75%

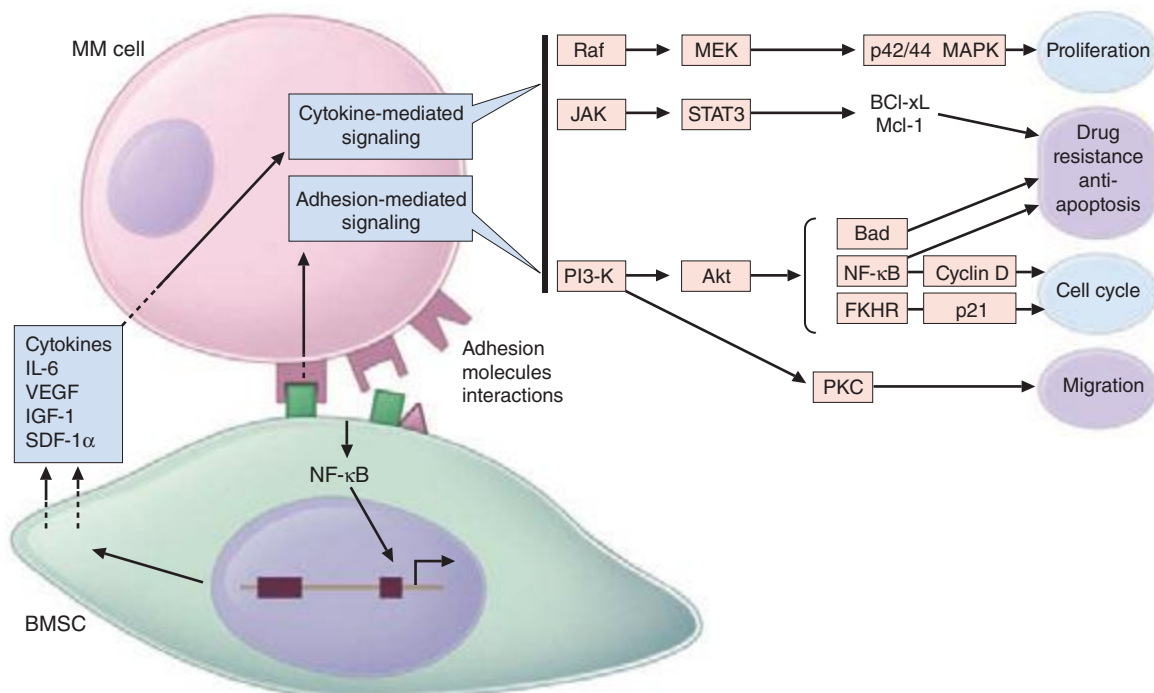


FIGURE 136-4 Pathogenesis of multiple myeloma. Multiple myeloma (MM) cells interact with bone marrow stromal cells (BMSCs) and extracellular matrix proteins via adhesion molecules, triggering adhesion-mediated signaling as well as cytokine production. This triggers cytokine-mediated signaling that provides growth, survival, and antiapoptotic effects as well as development of drug resistance.



FIGURE 136-5 Bony lesions in multiple myeloma. The skull demonstrates the typical “punched out” lesions characteristic of multiple myeloma. The lesion represents a purely osteolytic lesion with little or no osteoblastic activity. (Courtesy of Dr. Geraldine Schechter; with permission.)

of patients will have a serious infection at some time in their course. The susceptibility to infection has several contributing causes. First, patients with myeloma have diffuse hypogammaglobulinemia if the M component is excluded. The hypogammaglobulinemia is related to both decreased production and increased destruction of normal antibodies. Moreover, some patients generate a population of circulating regulatory cells in response to their myeloma that can suppress normal antibody synthesis. In the case of IgG myeloma, normal IgG antibodies are broken down more rapidly than normal because the catabolic rate for IgG antibodies varies directly with the serum concentration. The large M component results in fractional catabolic rates of 8–16% instead of the normal 2%. These patients have very poor antibody responses, especially to polysaccharide antigens such as those on bacterial cell walls. Most measures of T-cell function in myeloma are normal, but a subset of CD4+ cells may be decreased. Granulocyte lysozyme content is low, and granulocyte migration is not as rapid as normal in patients with myeloma, probably the result of a tumor product. There are also a variety of abnormalities in complement functions in myeloma patients. All these factors contribute to the immune deficiency of these patients. Some commonly used therapeutic agents, e.g., dexamethasone, suppress immune responses and increase susceptibility to bacterial and fungal infection, and bortezomib predisposes to herpesvirus reactivation.

Renal failure occurs in nearly 25% of myeloma patients, and some renal pathology is noted in more than 50%. Many factors contribute to this. Hypercalcemia is the most common cause of renal failure. Glomerular deposits of amyloid, hyperuricemia, recurrent infections, frequent use of nonsteroidal anti-inflammatory agents for pain control, use of iodinated contrast dye for imaging, bisphosphonate use, and occasional infiltration of the kidney by myeloma cells all may contribute to renal dysfunction. However, tubular damage associated with the excretion of light chains is almost always present. Normally, light chains are filtered, reabsorbed in the tubules, and catabolized. With the increase in the amount of light chains presented to the tubule, the tubular cells become overloaded with these proteins, and tubular damage results either directly from light chain toxic effects or indirectly from the release of intracellular lysosomal enzymes. The earliest manifestation of this tubular damage is the adult Fanconi’s syndrome (a type 2 proximal renal tubular acidosis), with loss of glucose and amino acids, as well as defects in the ability of the kidney to acidify and concentrate the urine. The proteinuria is not accompanied by hypertension, and the protein is nearly all light chains. Generally, very little albumin is in the urine because glomerular function is usually normal. When the glomeruli are involved, nonselective proteinuria

is also observed. Patients with myeloma also have a decreased anion gap [i.e., $\text{Na}^+ - (\text{Cl}^- + \text{HCO}_3^-)$] because the M component is cationic, resulting in retention of chloride. This is often accompanied by hyponatremia that is felt to be artificial (pseudohyponatremia) because each volume of serum has less water as a result of the increased protein. Renal dysfunction due to light chain deposition disease, light chain cast nephropathy, and amyloidosis is partially reversible with effective therapy. Myeloma patients are susceptible to developing acute renal failure if they become dehydrated.

Normocytic and normochromic anemia occurs in ~80% of myeloma patients. It is usually related to the replacement of normal marrow by expanding tumor cells, to the inhibition of hematopoiesis by factors made by the tumor, to reduced production of erythropoietin by the kidney, and to the effects of long-term therapy. In addition, mild hemolysis may contribute to the anemia. A larger than expected fraction of patients may have megaloblastic anemia due to either folate or vitamin B₁₂ deficiency. Granulocytopenia and thrombocytopenia are rare except when therapy-induced. Clotting abnormalities may be seen due to the failure of antibody-coated platelets to function properly; the interaction of the M component with clotting factors I, II, V, VII, or VIII; antibody to clotting factors; or amyloid damage of endothelium. Deep venous thrombosis is also observed with use of thalidomide, lenalidomide, or pomalidomide in combination with dexamethasone. Raynaud’s phenomenon and impaired circulation may result if the M component forms cryoglobulins, and hyperviscosity syndromes may develop depending on the physical properties of the M component (most common with IgM, IgG3, and IgA paraproteins). Hyperviscosity is defined based on the relative viscosity of serum as compared with water. Normal relative serum viscosity is 1.8 (i.e., serum is normally almost twice as viscous as water). Symptoms of hyperviscosity occur at a level greater than 4 centipoise (cP), which is usually reached at paraprotein concentrations of ~40 g/L (4 g/dL) for IgM, 50 g/L (5 g/dL) for IgG3, and 70 g/L (7 g/dL) for IgA; however, depending on chemical and physical properties of the paraprotein molecule, it can occasionally be observed at lower levels.

Although neurologic symptoms occur in a minority of patients, they may have many causes. Hypercalcemia may produce lethargy, weakness, depression, and confusion. Hyperviscosity may lead to headache, fatigue, shortness of breath, exacerbation or precipitation of heart failure, visual disturbances, ataxia, vertigo, retinopathy, somnolence, and coma. Bony damage and collapse may lead to cord compression, radicular pain, and loss of bowel and bladder control. Infiltration of peripheral nerves by amyloid can be a cause of carpal tunnel syndrome and other sensorimotor mono- and polyneuropathies. Neuropathy associated with monoclonal gammopathy of undetermined significance (MGUS) and myeloma is more frequently sensory than motor neuropathy and is associated with IgM more than other isotypes. In >50% of patients with neuropathy, the IgM monoclonal protein is directed against myelin-associated globulin (MAG). Sensory neuropathy is also a side effect of thalidomide and bortezomib therapy.

Many of the clinical features of myeloma, e.g., cord compression, pathologic fractures, hyperviscosity, sepsis, and hypercalcemia, can present as medical emergencies. Despite the widespread distribution of plasma cells in the body, tumor expansion is dominantly within bone and bone marrow and, for reasons unknown, rarely causes enlargement of spleen, lymph nodes, or gut-associated lymphatic tissue.

DIAGNOSIS AND STAGING

The diagnosis of myeloma requires marrow plasmacytosis (>10%), a serum and/or urine M component, and end organ damage detailed in [Table 136-1](#). Bone marrow plasma cells are CD138 and either monoclonal kappa or lambda light chain positive. The most important differential diagnosis in patients with myeloma involves their separation from individuals with MGUS or smoldering multiple myeloma (SMM). MGUS is vastly more common than myeloma, occurring in 1% of the population older than age 50 years and in up to 10% of individuals older than age 75 years. The diagnostic criteria for MGUS, SMM, and myeloma are described in [Table 136-1](#). Although ~1% of patients per year with MGUS go on to develop myeloma, all

TABLE 136-1 DIAGNOSTIC CRITERIA FOR MULTIPLE MYELOMA, MYELOMA VARIANTS, AND MONOCLONAL GAMMOPATHY OF UNDETERMINED SIGNIFICANCE

Monoclonal Gammopathy of Undetermined Significance (MGUS)
M protein in serum <30 g/L
Bone marrow clonal plasma cells <10%
No evidence of other B cell proliferative disorders
No myeloma-related organ or tissue impairment (no end organ damage, including bone lesions) ^a
Smoldering Multiple Myeloma (Asymptomatic Myeloma)
M protein in serum ≥30 g/L and/or
Bone marrow clonal plasma cells ≥10%
No myeloma-related organ or tissue impairment (no end organ damage, including bone lesions) ^a or symptoms
Symptomatic Multiple Myeloma
M protein in serum and/or urine
Bone marrow (clonal) plasma cells ^b or plasmacytoma
Myeloma-related organ or tissue impairment (end organ damage, including bone lesions)
Nonsecretory Myeloma
No M protein in serum and/or urine with immunofixation
Bone marrow clonal plasmacytosis ≥10% or plasmacytoma
Myeloma-related organ or tissue impairment (end organ damage, including bone lesions) ^a
Solitary Plasmacytoma of Bone
No M protein in serum and/or urine ^c
Single area of bone destruction due to clonal plasma cells
Bone marrow not consistent with multiple myeloma
Normal skeletal survey (and magnetic resonance imaging of spine and pelvis if done)
No related organ or tissue impairment (no end organ damage other than solitary bone lesion) ^a
POEMS Syndrome
All of the following four criteria must be met:
1. Polyneuropathy
2. Monoclonal plasma cell proliferative disorder
3. Any one of the following: (a) sclerotic bone lesions; (b) Castleman's disease; (c) elevated levels of vascular endothelial growth factor (VEGF)
4. Any one of the following: (a) organomegaly (splenomegaly, hepatomegaly, or lymphadenopathy); (b) extravascular volume overload (edema, pleural effusion, or ascites); (c) endocrinopathy (adrenal, thyroid, pituitary, gonadal, parathyroid, and pancreatic); (d) skin changes (hyperpigmentation, hypertrichosis, glomeruloid hemangiomas, plethora, acrocyanosis, flushing, and white nails); (e) papilledema; (f) thrombocytosis/polycythemia ^d

^aMyeloma-related organ or tissue impairment (end organ damage): calcium levels increased: serum calcium >0.25 mmol/L above the upper limit of normal or >2.75 mmol/L; renal insufficiency: creatinine >173 mmol/L; anemia: hemoglobin 2 g/dL below the lower limit of normal or hemoglobin <10 g/dL; bone lesions: lytic lesions or osteoporosis with compression fractures (magnetic resonance imaging or computed tomography may clarify); other: symptomatic hyperviscosity, amyloidosis, recurrent bacterial infections (>2 episodes in 12 months). ^bIf flow cytometry is performed, most plasma cells (>90%) will show a "neoplastic" phenotype. ^cA small M component may sometimes be present. ^dThese features should have no attributable other causes and have temporal relation with each other.

Abbreviation: POEMS, polyneuropathy, organomegaly, endocrinopathy, M-protein, and skin changes.

myeloma is preceded by MGUS. Non-IgG subtype, abnormal kappa/lambda free light chain ratio, and serum M protein >15 g/L (1.5 g/dL) are associated with higher incidence of progression of MGUS to myeloma. Absence of all three features predicts a 5% chance of progression, whereas higher risk MGUS with the presence of all three features predicts a 60% chance of progression over 20 years. The features responsible for higher risk of progression from SMM to MM are bone marrow plasmacytosis >10%, abnormal kappa/lambda free light chain ratio, and serum M protein >30 g/L (3 g/dL). Patients with only one of these three features have a 25% chance of progression to MM in 5 years, whereas patients with high-risk SMM with all three features

have a 76% chance of progression. There are two important variants of myeloma—solitary bone plasmacytoma and solitary extramedullary plasmacytoma. These lesions are associated with an M component in <30% of the cases, they may affect younger individuals, and both are associated with median survivals of ≥10 years. Solitary bone plasmacytoma is a single lytic bone lesion without marrow plasmacytosis. Extramedullary plasmacytomas usually involve the submucosal lymphoid tissue of the nasopharynx or paranasal sinuses without marrow plasmacytosis. Both tumors are highly responsive to local radiation therapy. If an M component is present, it should disappear after treatment. Solitary bone plasmacytomas may recur in other bony sites or evolve into myeloma. Extramedullary plasmacytomas rarely recur or progress.

The clinical evaluation of patients with myeloma includes a careful physical examination searching for tender bones and masses. Chest and bone radiographs may reveal lytic lesions or diffuse osteopenia. Magnetic resonance imaging (MRI) offers a sensitive means to document extent of bone marrow infiltration and cord or root compression in patients with pain syndromes. A complete blood count with differential may reveal anemia. Erythrocyte sedimentation rate is elevated. Rare patients (~1%) may have plasma cell leukemia with >2000 plasma cells/μL. This may be seen in disproportionate frequency in IgD (12%) and IgE (25%) myelomas. Serum calcium, urea nitrogen, creatinine, and uric acid levels may be elevated. Protein electrophoresis and measurement of serum immunoglobulins and free light chains are useful for detecting and characterizing M spikes, supplemented by immunoelectrophoresis, which is especially sensitive for identifying low concentrations of M components not detectable by protein electrophoresis. A 24-h urine specimen is necessary to quantitate Bence Jones protein excretion. Serum alkaline phosphatase is usually normal even with extensive bone involvement because of the absence of osteoblastic activity. It is also important to quantitate serum β₂-microglobulin and albumin (see below).

The serum M component will be IgG in 53% of patients, IgA in 25%, and IgD in 1%; 20% of patients will have only light chains in serum and urine. Dipsticks for detecting proteinuria are not reliable at identifying light chains, and the heat test for detecting Bence Jones protein is falsely negative in ~50% of patients with light chain myeloma. Fewer than 1% of patients have no identifiable M component; these patients usually have light chain myeloma in which renal catabolism has made the light chains undetectable in the urine. In most of these patients, light chains can now be detected by serum free light chain assay. IgD myeloma may also present with light chain disease. About two-thirds of patients with serum M components also have urinary light chains. The light chain isotype may have an impact on survival. Patients secreting lambda light chains have a significantly shorter overall survival than those secreting kappa light chains. Whether this is due to some genetically important determinant of cell proliferation or because lambda light chains are more likely to cause renal damage and form amyloid than are kappa light chains is unclear. The heavy chain isotype may have an impact on patient management as well. About half of patients with IgM paraproteins develop hyperviscosity compared with only 2–4% of patients with IgA and IgG M components. Among IgG myelomas, it is the IgG3 subclass that has the highest tendency to form both concentration- and temperature-dependent aggregates, leading to hyperviscosity and cold agglutination at lower serum concentrations.

PROGNOSIS

Serum β₂-microglobulin is the single most powerful predictor of survival and can substitute for staging. β₂-Microglobulin is a protein of 11,000 mol wt with homologies to the constant region of immunoglobulins that is the light chain of the class I major histocompatibility antigens (HLA-A, -B, -C) on the surface of every cell. Patients with β₂-microglobulin levels <0.004 g/L have a median survival of 43 months, and those with levels >0.004 g/L have a survival of only 12 months. Combination of serum β₂-microglobulin and albumin levels forms the basis for a three-stage International Staging System (ISS) (Table 136-2) that predicts survival. With the use of high-dose therapy and the newer agents, the

TABLE 136-2 RISK STRATIFICATION IN MYELOMA

Chromosomal Abnormalities		
Method	Standard Risk (80%) (expected survival 6–7+ years)	High Risk (20%) (expected survival 2–3 years)
Karyotype	No chromosomal aberration	Any abnormality on conventional karyotype
FISH	t(11;14)	Del(17p)
	t(6;14)	t(4;14)
	Del(13)	t(14;16) t(14;20)
International Staging System		
	Stage	Median Survival, Months
β_2 M <3.5, alb \geq 3.5	I (28%) ^a	62
β_2 M <3.5, alb <3.5 or β_2 M = 3.5–5.5	II (39%)	44
β_2 M >5.5	III (33%)	29
Other features suggesting high-risk disease:		
De novo plasma cell leukemia		
Extramedullary disease		
Elevated lactate dehydrogenase (LDH)		
High-risk gene expression profile		

^aPercentage of patients presenting at each stage.

Abbreviations: β_2 M, serum β_2 -microglobulin in mg/L; alb, serum albumin in g/dL; FISH, fluorescent in situ hybridization.

Durie-Salmon staging system is unable to predict outcome and thus is no longer used. High labeling index, circulating plasma cells, performance status, and high levels of lactate dehydrogenase are also associated with poor prognosis.

Other factors that may influence prognosis are the presence of cytogenetic abnormalities and hypodiploidy by karyotype, fluorescent in situ hybridization (FISH)-identified chromosome 17p deletion, and translocations t(4;14), (14;16), and t(14;20). Chromosome 13q deletion, previously thought to predict poor outcome, is not a predictor following the use of newer agents. Microarray profiling and comparative genomic hybridization have formed the basis for RNA- and DNA-based prognostic staging systems, respectively. The ISS system, along with cytogenetic changes, is the most widely used method for assessing prognosis (Table 136-2).

TREATMENT MULTIPLE MYELOMA

No specific intervention is indicated for patients with MGUS. Follow-up once a year or less frequently is adequate except in higher risk MGUS, where serum protein electrophoresis, complete blood count, creatinine, and calcium should be repeated every 6 months. A patient with MGUS and severe polyneuropathy is considered for therapeutic intervention if a causal relationship can be assumed, especially in absence of any other potential causes for neuropathy. Therapy can include plasmapheresis and occasionally rituximab in patients with IgM MGUS or myeloma-like therapy in those with IgG or IgA disease. About 10% of patients with myeloma are asymptomatic (SMM) and will have an indolent course demonstrating only very slow progression of disease over many years. For these patients, no specific therapeutic intervention is indicated, although early intervention with lenalidomide and dexamethasone may prevent progression from high-risk SMM to active MM. At present, patients with SMM only require antitumor therapy when the disease becomes symptomatic with development of anemia, hypercalcemia, progressive lytic bone lesions, renal dysfunction, or recurrent infections. Patients with solitary bone plasmacytomas and extramedullary plasmacytomas may be expected to enjoy

prolonged disease-free survival after local radiation therapy at a dose of around 40 Gy. There is a low incidence of occult marrow involvement in patients with solitary bone plasmacytoma. Such patients are usually identified because their serum M component falls slowly or disappears initially, only to return after a few months. These patients respond well to systemic therapy.

Patients with symptomatic and/or progressive myeloma require therapeutic intervention. In general, such therapy is of two sorts: (1) systemic therapy to control the progression of myeloma and (2) symptomatic supportive care to prevent serious morbidity from the complications of the disease. Therapy can significantly prolong survival and improve the quality of life for myeloma patients.

The therapy of myeloma includes an initial induction regimen followed by consolidation and/or maintenance therapy and, on subsequent progression, management of relapsed disease. The therapy is partly dictated by the patient's age and comorbidities, which may affect a patient's ability to undergo high-dose therapy and transplantation.

Thalidomide (200 mg daily), when combined with dexamethasone, achieved responses in two-thirds of newly diagnosed MM patients. Subsequently, lenalidomide (25 mg/d on days 1–21 every 4 weeks), an immunomodulatory derivative of thalidomide, and bortezomib (1.3 mg/m² on days 1, 4, 8, and 11 every 3 weeks), a proteasome inhibitor, have each been combined with dexamethasone (40 mg once every week) and obtained high response rates (>80%) in newly diagnosed patients with MM. Importantly, their superior toxicity profile with improved efficacy has made them the preferred agents for induction therapy. Efforts to improve the fraction of patients responding and the degree of response have involved adding agents to the treatment regimen. The combination of lenalidomide, bortezomib, and dexamethasone achieves close to a 100% response rate and 30% complete response rate, making it one of the preferred induction regimens in transplant-eligible patients. Other similar three-drug combinations (bortezomib, thalidomide, and dexamethasone or bortezomib, cyclophosphamide, and dexamethasone) also achieve >90% response rate. Herpes zoster prophylaxis is indicated if bortezomib is used, and neuropathy attendant to bortezomib can be decreased both by its subcutaneous administration and administration on a weekly schedule. Lenalidomide use requires prophylaxis for deep vein thrombosis (DVT) with either aspirin or warfarin or low-molecular-weight heparin if patients are at a greater risk of DVT. In patients receiving lenalidomide, stem cells should be collected within 6 months, because the continued use of lenalidomide may compromise the ability to collect adequate numbers of stem cells. Initial therapy is continued until maximal cytoreduction. In patients who are transplant candidates, alkylating agents such as melphalan should be avoided because they damage stem cells, leading to decreased ability to collect stem cells for autologous transplant.

In patients who are not transplant candidates due to physiologic age >70 years, significant cardiopulmonary problems, or other comorbid illnesses, the same two- or three-drug combinations described above are considered standard of care as induction therapy. Previously, therapy consisting of intermittent pulses of melphalan, an alkylating agent, with prednisone (MP; melphalan, 0.25 mg/kg per day, and prednisone, 1 mg/kg per day for 4 days) every 4–6 weeks was used. However, a number of studies have combined novel agents with MP and reported superior response and survival outcomes. In patients >65 years old, combining thalidomide with MP (MPT) obtains higher response rates and overall survival compared with MP alone. Similarly, significantly improved response (71 vs 35%) and overall survival (3-year survival 72 vs 59%) were observed with the combination of bortezomib and MP compared with MP alone. Lenalidomide added to MP followed by lenalidomide maintenance also prolonged progression-free survival compared with MP alone. These combinations of novel agents with MP also achieve high complete response rates (MPT, ~15%; MP plus bortezomib, ~30%; MP plus lenalidomide, ~20%; and MP, ~2–4%). Although combinations of MP with newer agents are an alternative

in these patients, most studies favor continuous therapy with non-MP-containing regimens (e.g., lenalidomide plus dexamethasone) due to longer term safety profile and efficacy.

Improvement in the serum M component may lag behind the symptomatic improvement. The fall in M component depends on the rate of tumor kill and the fractional catabolic rate of immunoglobulin, which in turn depends on the serum concentration (for IgG). Light chain excretion, with a functional half-life of ~6 h, may fall within the first week of treatment. Because urine light chain levels may relate to renal tubular function, they are not a reliable measure of tumor cell kill, especially in patients with renal dysfunction; however, improvements in serum free light chain measurement are often seen sooner. Although patients may not achieve complete remission, clinical responses may last for long periods of time.

High-dose therapy and consolidation/maintenance are standard practice in the majority of eligible patients. Randomized studies comparing standard-dose therapy to high-dose melphalan therapy (HDT) with hematopoietic stem cell support have shown that HDT can achieve high overall response rates, with up to 25–40% additional complete responses and prolonged progression-free and overall survival; however, few, if any, patients are cured. Although two successive HDTs (tandem transplantations) are more effective than single HDT, the benefit is only observed in the subset of patients who do not achieve a complete or very good partial response to the first transplantation, which is rare. Moreover, a randomized study failed to show any significant difference in overall survival between early transplantation after induction therapy versus delayed transplantation at relapse. These data allow an option to delay transplantation, especially with the availability of more agents and combinations. Allogeneic transplantations may also produce high response rates, but treatment-related mortality may be as high as 40%. Nonmyeloablative allogeneic transplantation can reduce toxicity but is recommended only under the auspices of a clinical trial to exploit an immune graft-versus-myeloma effect while avoiding attendant toxicity.

Maintenance therapy prolongs remissions following standard-dose regimens as well as HDT. Two phase 3 studies have demonstrated improved progression-free survival, and one study showed prolonged overall survival in patients receiving lenalidomide compared to placebo as maintenance therapy after HDT. In non-transplant candidates, another phase 3 study showed prolonged progression-free survival with lenalidomide maintenance after MP plus lenalidomide induction therapy. Although there is concern regarding an increased incidence of second primary malignancies in patients receiving lenalidomide maintenance, its benefits far outweigh the risk of progressive disease and death from myeloma. In patients with high-risk cytogenetics, lenalidomide and bortezomib have been combined and show promise as maintenance therapy after transplantation.

Relapsed myeloma can be treated with a number of agents including lenalidomide and/or bortezomib. These agents in combination with dexamethasone can achieve a partial response rate of up to 60% and a 10–15% complete response rate in patients with relapsed disease. The combination of bortezomib and liposomal doxorubicin is active in relapsed myeloma. Thalidomide, if not used as initial therapy, can achieve responses in refractory cases. The second-generation proteasome inhibitor carfilzomib and immunomodulatory agent pomalidomide have shown efficacy in relapsed and refractory MM, even MM refractory to lenalidomide and bortezomib. High-dose melphalan and stem cell transplantation, if not used earlier, also have activity as salvage therapy in patients with refractory disease.

The median overall survival of patients with myeloma is 7–8+ years, with subsets of younger patients surviving more than 10 years. The major causes of death are progressive myeloma, renal failure, sepsis, or therapy-related myelodysplasia. Nearly a quarter of patients die of myocardial infarction, chronic lung disease, diabetes, or stroke—all intercurrent illnesses related more to the age of the patient group than to the tumor.

Supportive care directed at the anticipated complications of the disease may be as important as primary antitumor therapy. Hypercalcemia generally responds well to bisphosphonates, glucocorticoid therapy, hydration, and natriuresis, and rarely requires calcitonin as well. Bisphosphonates (e.g., pamidronate 90 mg or zoledronate 4 mg once a month) reduce osteoclastic bone resorption and preserve performance status and quality of life, decrease bone-related complications, and may also have antitumor effects. Osteonecrosis of the jaw and renal dysfunction can occur in a minority of patients receiving aminobisphosphonate therapy. Treatments aimed at strengthening the skeleton such as fluorides, calcium, and vitamin D, with or without androgens, have been suggested, but are not of proven efficacy. Kyphoplasty or vertebroplasty should be considered in patients with painful collapsed vertebra. Iatrogenic worsening of renal function may be prevented by maintaining a high fluid intake to prevent dehydration and enhance excretion of light chains and calcium. In the event of acute renal failure, plasmapheresis is ~10 times more effective at clearing light chains than peritoneal dialysis; however, its role in reversing renal failure remains controversial. Importantly, reducing the protein load by effective antitumor therapy with agents such as bortezomib may result in improvement in renal function in over half of the patients. Use of lenalidomide in renal failure is possible but requires dose modification, because it is renally excreted. Urinary tract infections should be watched for and treated early. Plasmapheresis may be the treatment of choice for hyperviscosity syndromes. Although the pneumococcus is a dreaded pathogen in myeloma patients, pneumococcal polysaccharide vaccines may not elicit an antibody response. Prophylactic administration of intravenous γ globulin preparations is used in the setting of recurrent serious infections. Chronic oral antibiotic prophylaxis is not warranted. Patients developing neurologic symptoms in the lower extremities, severe localized back pain, or problems with bowel and bladder control may need emergency MRI and local radiation therapy and glucocorticoids if cord compression is identified. In patients in whom neurologic deficit is increasing or substantial, emergent surgical decompression may be necessary. Most bone lesions respond to analgesics and systemic therapy, but certain painful lesions may respond most promptly to localized radiation. The anemia associated with myeloma may respond to erythropoietin along with hematinics (iron, folate, cobalamin). The pathogenesis of the anemia should be established and specific therapy instituted, whenever possible.

WALDENSTRÖM'S MACROGLOBULINEMIA

In 1948, Waldenström described a malignancy of lymphoplasmacytoid cells that secreted IgM. In contrast to myeloma, the disease was associated with lymphadenopathy and hepatosplenomegaly, but the major clinical manifestation was hyperviscosity syndrome. The disease resembles the related diseases chronic lymphocytic leukemia, myeloma, and lymphocytic lymphoma. It originates from a post-germinal center B cell that has undergone somatic mutations and antigenic selection in the lymphoid follicle and has the characteristics of an IgM-bearing memory B cell. Waldenström's macroglobulinemia (WM) and IgM myeloma follow a similar clinical course, but therapeutic options are different. The diagnosis of IgM myeloma is usually reserved for patients with lytic bone lesions and predominant infiltration with CD138+ plasma cells in the bone marrow. Such patients are at greater risk of pathologic fractures than patients with WM.

A familial occurrence is common in WM, but its molecular bases are yet unclear. A distinct MYD88 L265P somatic mutation has been reported in over 90% of patients with WM and the majority of IgM MGUS. Presence of this mutation is now used as a diagnostic test to discriminate WM from marginal zone lymphomas (MZLs), IgM-secreting myeloma, and chronic lymphocytic leukemia (CLL) with plasmacytic differentiation. This mutation also explains the molecular pathogenesis of the disease, with involvement of Toll-like receptor (TLR) and interleukin 1 receptor (IL-1R) signaling leading to

718 activation of IL-1R-associated kinase (IRAK) 4 and IRAK1 followed by nuclear factor- κ B (NF- κ B) activation. The disease is similar to myeloma in being slightly more common in men and occurring with increased incidence with increasing age (median 64 years). There have been reports that the IgM in some patients with macroglobulinemia may have specificity for myelin-associated glycoprotein (MAG), a protein that has been associated with demyelinating disease of the peripheral nervous system and may be lost earlier and to a greater extent than the better known myelin basic protein in patients with multiple sclerosis. Sometimes patients with macroglobulinemia develop a peripheral neuropathy, and half of these patients are positive for anti-MAG antibody. The neuropathy may precede the appearance of the neoplasm. There is speculation that the whole process begins with a viral infection that may elicit an antibody response that cross-reacts with a normal tissue component.

Like myeloma, the disease involves the bone marrow, but unlike myeloma, it does not cause bone lesions or hypercalcemia. Bone marrow shows >10% infiltration with lymphoplasmacytic cells (surface IgM+, CD19+, CD20+, and CD22+, rarely CD5+, but CD10- and CD23-) with an increase in number of mast cells. Like myeloma, an M component is present in the serum in excess of 30 g/L (3 g/dL), but unlike myeloma, the size of the IgM paraprotein results in little renal excretion, and only ~20% of patients excrete light chains. Therefore, renal disease is not common. The light chain isotype is kappa in 80% of the cases. Patients present with weakness, fatigue, and recurrent infections similar to myeloma patients, but epistaxis, visual disturbances, and neurologic symptoms such as peripheral neuropathy, dizziness, headache, and transient paresis are much more common in macroglobulinemia. Physical examination reveals adenopathy and hepatosplenomegaly, and ophthalmoscopic examination may reveal vascular segmentation and dilation of the retinal veins characteristic of hyperviscosity states. Patients may have a normocytic, normochromic anemia, but rouleaux formation and a positive Coombs' test are much more common than in myeloma. Malignant lymphocytes are usually present in the peripheral blood. About 10% of macroglobulins are cryoglobulins. These are pure M components and are not the mixed cryoglobulins seen in rheumatoid arthritis and other autoimmune diseases. Mixed cryoglobulins are composed of IgM or IgA complexed with IgG, for which they are specific. In both cases, Raynaud's phenomenon and serious vascular symptoms precipitated by the cold may occur, but mixed cryoglobulins are not commonly associated with malignancy. Patients suspected of having a cryoglobulin based on history and physical examination should have their blood drawn into a warm syringe and delivered to the laboratory in a container of warm water to avoid errors in quantitating the cryoglobulin.

TREATMENT WALDENSTRÖM'S MACROGLOBULINEMIA

Control of serious hyperviscosity symptoms such as an altered state of consciousness or paresis can be achieved acutely by plasmapheresis because 80% of the IgM paraprotein is intravascular. The median survival of affected individuals is ~50 months, similar to that of MM. However, many patients with WM have indolent disease that does not require therapy. Pretreatment parameters including older age, male sex, general symptoms, and cytopenias define a high-risk population. Treatment is usually not initiated unless the disease is symptomatic or increasing anemia, hyperviscosity, lymphadenopathy, or hepatosplenomegaly is present. Bortezomib and bendamustine are two agents with significant efficacy in WM. Rituximab (anti-CD20) can produce responses, alone or combined with either of these two agents. Rituximab can produce IgM flare, so its use is initially withheld in patients with high IgM levels. Fludarabine (25 mg/m² per day for 5 days every 4 weeks) and cladribine (0.1 mg/kg per day for 7 days every 4 weeks) are also highly effective single agents. With identification of the MYD88 mutation, BTK and IRAK1/4 inhibitors are being evaluated and show significant responses. Although high-dose therapy plus autologous transplantation is an option, its use has declined due to the availability of other effective agents.

POEMS SYNDROME

The features of this syndrome are polyneuropathy, organomegaly, endocrinopathy, M-protein, and skin changes (POEMS). Diagnostic criteria are described in Table 136-1. Patients usually have a severe, progressive sensorimotor polyneuropathy associated with sclerotic bone lesions from myeloma. Polyneuropathy occurs in ~1.4% of myelomas, but the POEMS syndrome is only a rare subset of that group. Unlike typical myeloma, hepatomegaly and lymphadenopathy occur in about two-thirds of patients, and splenomegaly is seen in one-third. The lymphadenopathy frequently resembles Castleman's disease histologically, a condition that has been linked to IL-6 overproduction. The endocrine manifestations include amenorrhea in women and impotence and gynecomastia in men. Hyperprolactinemia due to loss of normal inhibitory control by the hypothalamus may be associated with other central nervous system manifestations such as papilledema and elevated cerebrospinal fluid pressure and protein. Type 2 diabetes mellitus occurs in about one-third of patients. Hypothyroidism and adrenal insufficiency are occasionally noted. Skin changes are diverse: hyperpigmentation, hypertrichosis, skin thickening, and digital clubbing. Other manifestations include peripheral edema, ascites, pleural effusions, fever, and thrombocytosis. Not all the components of POEMS syndrome may be present initially.

The pathogenesis of the disease is unclear, but high circulating levels of the proinflammatory cytokines IL-1, IL-6, VEGF, and TNF have been documented, and levels of the inhibitory cytokine transforming growth factor β are lower than expected. Treatment of the myeloma may result in an improvement in the other disease manifestations.

Patients are often treated similarly to those with myeloma. Plasmapheresis does not appear to be of benefit in POEMS syndrome. Patients presenting with isolated sclerotic lesions may have resolution of neuropathic symptoms after local therapy for plasmacytoma with radiotherapy. Similar to multiple myeloma, novel agents and high-dose therapy with autologous stem cell transplantation have been pursued in selected patients and have been associated with prolonged progression-free survival.

HEAVY CHAIN DISEASES

The heavy chain diseases are rare lymphoplasmacytic malignancies. Their clinical manifestations vary with the heavy chain isotype. Patients have absence of light chain and secrete a defective heavy chain that usually has an intact Fc fragment and a deletion in the Fd region. Gamma, alpha, and mu heavy chain diseases have been described, but no reports of delta or epsilon heavy chain diseases have appeared. Molecular biologic analysis of these tumors has revealed structural genetic defects that may account for the aberrant chain secreted.

GAMMA HEAVY CHAIN DISEASE (FRANKLIN'S DISEASE)

This disease affects individuals of widely different age groups and countries of origin. It is characterized by lymphadenopathy, fever, anemia, malaise, hepatosplenomegaly, and weakness. It is frequently associated with autoimmune diseases, especially rheumatoid arthritis. Its most distinctive symptom is palatal edema, resulting from involvement of nodes in Waldeyer's ring, and this may progress to produce respiratory compromise. The diagnosis depends on the demonstration of an anomalous serum M component (often <20 g/L [<2 g/dL]) that reacts with anti-IgG but not anti-light chain reagents. The M component is typically present in both serum and urine. Most of the paraproteins have been of the γ_1 subclass, but other subclasses have been seen. The patients may have thrombocytopenia, eosinophilia, and nondiagnostic bone marrow that may show increased numbers of lymphocytes or plasma cells that do not stain for light chain. Patients usually have a rapid downhill course and die of infection; however, some patients have survived 5 years with chemotherapy. Therapy is indicated when symptomatic and involves chemotherapeutic combinations used in low-grade lymphoma. Rituximab has also been reported to show efficacy.

ALPHA HEAVY CHAIN DISEASE (SELIGMANN'S DISEASE)

This is the most common of the heavy chain diseases. It is closely related to a malignancy known as Mediterranean lymphoma, a disease that

affects young persons in parts of the world where intestinal parasites are common, such as the Mediterranean, Asia, and South America. The disease is characterized by an infiltration of the lamina propria of the small intestine with lymphoplasmacytoid cells that secrete truncated alpha chains. Demonstrating alpha heavy chains is difficult because the alpha chains tend to polymerize and appear as a smear instead of a sharp peak on electrophoretic profiles. Despite the polymerization, hyperviscosity is not a common problem in alpha heavy chain disease. Without J chain–facilitated dimerization, viscosity does not increase dramatically. Light chains are absent from serum and urine. The patients present with chronic diarrhea, weight loss, and malabsorption and have extensive mesenteric and paraaortic adenopathy. Respiratory tract involvement occurs rarely. Patients may vary widely in their clinical course. Some may develop diffuse aggressive histologies of malignant lymphoma. Chemotherapy may produce long-term remissions. Rare patients appear to have responded to antibiotic therapy, raising the question of the etiologic role of antigenic stimulation, perhaps by some chronic intestinal infection. Chemotherapy plus antibiotics may be more effective than chemotherapy alone. Immunoproliferative small-intestinal disease (IPSID) is recognized as an infectious pathogen–associated human lymphoma that has association with *Campylobacter jejuni*. It involves mainly the proximal small intestine resulting in malabsorption, diarrhea, and abdominal pain. IPSID is associated with excessive plasma cell differentiation and produces truncated alpha heavy chain proteins lacking the light chains as well as the first constant domain. Early-stage IPSID responds to antibiotics (30–70% complete remission). Most untreated IPSID patients progress to lymphoplasmacytic and immunoblastic lymphoma. Patients not responding to antibiotic therapy are considered for treatment with combination chemotherapy used to treat low-grade lymphoma.

MU HEAVY CHAIN DISEASE

The secretion of isolated mu heavy chains into the serum appears to occur in a very rare subset of patients with CLL. The only features that may distinguish patients with mu heavy chain disease are the presence of vacuoles in the malignant lymphocytes and the excretion of kappa light chains in the urine. The diagnosis requires ultracentrifugation or gel filtration to confirm the nonreactivity of the paraprotein with the light chain reagents, because some intact macroglobulins fail to interact with these serums. The tumor cells seem to have a defect in the assembly of light and heavy chains, because they appear to contain both in their cytoplasm. There is no evidence that such patients should be treated differently from other patients with CLL ([Chap. 134](#)).

137 Amyloidosis

David C. Seldin, John L. Berk

GENERAL PRINCIPLES

Amyloidosis is the term for a group of protein folding disorders characterized by the extracellular deposition of insoluble polymeric protein fibrils in tissues and organs. A robust cellular machinery exists to chaperone proteins during the process of synthesis and secretion, to ensure that they achieve correct tertiary conformation and function, and to eliminate proteins that misfold. However, genetic mutation, incorrect processing, and other factors may favor misfolding, with consequent loss of normal protein function and intracellular or extracellular aggregation. Many diseases, ranging from cystic fibrosis to Alzheimer's disease, are now known to involve protein misfolding. In the amyloidoses, the aggregates are typically extracellular, and the misfolded protein subunits assume a common antiparallel, β -pleated sheet–rich structural conformation that leads to the formation of higher-order oligomers and then of fibrils with unique staining properties. The term *amyloid* was coined around 1854 by the pathologist Rudolf Virchow,

who thought that these deposits resembled starch (Latin *amylum*) under the microscope.

Amyloid diseases, defined by the biochemical nature of the protein composing the fibril deposits, are classified according to whether they are systemic or localized, whether they are acquired or inherited, and their clinical patterns ([Table 137-1](#)). The standard nomenclature is AX, where A indicates amyloidosis and X represents the protein present in the fibril. This chapter focuses primarily on the systemic forms. AL refers to amyloid composed of immunoglobulin light chains (LCs); this disorder, formerly termed *primary systemic amyloidosis*, arises from a clonal B cell or plasma cell disorder and can be associated with myeloma or lymphoma. AF refers to the *familial amyloidoses*, which are most commonly due to mutations in transthyretin (TTR), the transport protein for thyroid hormone and retinol-binding protein. AA amyloid is composed of the acute-phase reactant protein serum amyloid A (SAA) and occurs in the setting of chronic inflammatory or infectious diseases; for this reason, this type was formerly known as *secondary amyloidosis*. $A\beta_2M$ amyloid results from misfolded β_2 -microglobulin, occurring in individuals with long-standing renal disease who have undergone dialysis, typically for years. $A\beta$, the most common form of localized amyloidosis, is found in the brain of patients with Alzheimer's disease after abnormal proteolytic processing and aggregation of polypeptides derived from the amyloid precursor protein.

Diagnosis and treatment of the amyloidoses rest upon the histopathologic identification of amyloid deposits and immunohistochemical, biochemical, or genetic determination of amyloid type ([Fig. 137-1](#)). In the systemic amyloidoses, the clinically involved organs can be biopsied, but amyloid deposits may be found in any tissue of the body. Historically, blood vessels of the gingiva or rectal mucosa were often examined, but the most easily accessible tissue—positive in more than 80% of patients with systemic amyloidosis—is fat. After local anesthesia, fat is aspirated from the abdominal pannus with a 16-gauge needle. Fat globules expelled onto a glass slide can be stained, thus avoiding a surgical procedure. If this material is negative, more invasive biopsies of the kidney, heart, liver, or gastrointestinal tract can be considered in patients in whom amyloidosis is suspected. The regular β -sheet structure of amyloid deposits exhibits a unique “apple green” birefringence by polarized light microscopy when stained with Congo red dye; other regular protein structures (e.g., collagen) appear white under these conditions. The 10-nm-diameter fibrils can also be visualized by electron microscopy of paraformaldehyde-fixed tissue. Once amyloid is found, the protein type must be determined by immunohistochemistry, immunoelectron microscopy, or extraction and biochemical analysis employing mass spectrometry; gene sequencing is used to identify mutants causing AF amyloid. The patient's history, physical findings, and clinical presentation, including age and ethnic origin, organ system involvement, underlying diseases, and family history, may provide helpful clues as to the type of amyloid. However, there can be considerable overlap in clinical presentations, and accurate typing is essential to guide appropriate therapy.

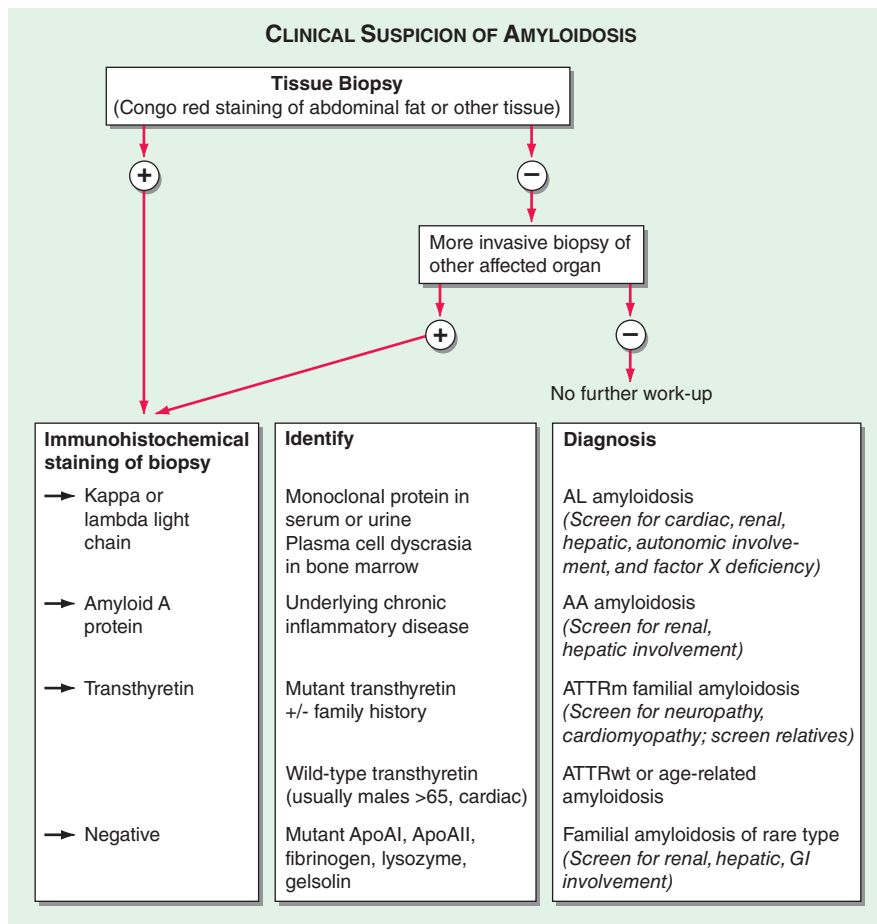
The mechanisms of fibril formation and tissue toxicity remain controversial. The “amyloid hypothesis,” as it is currently understood, proposes that precursor proteins undergo a process of reversible unfolding or misfolding; misfolded proteins form oligomeric aggregates, higher-order polymers, and then fibrils that deposit in tissues. Accumulating evidence suggests that the oligomeric intermediates may constitute the most toxic species. Oligomers are more capable than large fibrils of interacting with cells and inducing formation of reactive oxygen species and stress signaling. Ultimately, the fibrillar tissue deposits are likely to interfere with normal organ function. A more sophisticated understanding of the mechanisms leading to amyloid formation and cell and tissue dysfunction will continue to provide new targets for therapies.

The clinical syndromes of the amyloidoses are associated with relatively nonspecific alterations in routine laboratory tests. Blood counts are usually normal, although the erythrocyte sedimentation rate is frequently elevated. Patients with glomerular kidney involvement generally have proteinuria, often in the nephrotic range, leading to

TABLE 137-1 AMYLOID PRECURSOR PROTEINS AND THEIR CLINICAL SYNDROMES

Designation	Precursor	Clinical Syndrome	Clinical Involvement
Systemic Amyloidoses			
AL	Immunoglobulin light chain	Primary or myeloma-associated ^a	Any
AH	Immunoglobulin heavy chain	Rare variant of primary or myeloma-associated	Any
AA	Serum amyloid A protein	Secondary; reactive ^b	Renal, other
A β_2 M	β_2 -Microglobulin	Hemodialysis-associated	Synovial tissue, bone
ATTR	Transthyretin	Familial (mutant) Age-related (wild type)	Cardiac, peripheral and autonomic nerves
AApoAI	Apolipoprotein AI	Familial	Hepatic, renal
AApoAII	Apolipoprotein AII	Familial	Renal
AGel	Gelsolin	Familial	Cornea, cranial nerves, skin, renal
AFib	Fibrinogen A α	Familial	Renal
ALys	Lysozyme	Familial	Renal, hepatic
ALECT2	Leukocyte chemotactic factor 2	Undefined	Renal
Localized Amyloidoses			
A β	Amyloid β protein	Alzheimer's disease; Down syndrome	Central nervous system
ACys	Cystatin C	Cerebral amyloid angiopathy	Central nervous system, vascular
APrP	Prion protein	Spongiform encephalopathies	Central nervous system
AIAPP	Islet amyloid polypeptide (amylin)	Diabetes-associated	Pancreas
ACal	Calcitonin	Medullary carcinoma of the thyroid	Thyroid
AANF	Atrial natriuretic factor	Atrial fibrillation	Cardiac atria
APro	Prolactin	Endocrinopathy	Pituitary
ASgl	Semenogelin I	Age-related; incidental autopsy or biopsy finding	Seminal vesicles

^aLocalized AL deposits can occur in skin, conjunctiva, urinary bladder, and the tracheobronchial tree. ^bSecondary to chronic inflammation or infection or to a hereditary periodic fever syndrome such as familial Mediterranean fever.

**FIGURE 137-1** Algorithm for the diagnosis of amyloidosis and determination of type.

Clinical suspicion: unexplained nephropathy, cardiomyopathy, neuropathy, enteropathy, arthropathy, and macroglossia. ApoAI, apolipoprotein AI; ApoAII, apolipoprotein AII; GI, gastrointestinal.

hypoalbuminemia that may be severe; patients with serum albumin levels below 2 g/dL generally have pedal edema or anasarca. Amyloid cardiomyopathy is characterized by concentric ventricular hypertrophy and diastolic dysfunction associated with elevation of brain natriuretic peptide or N-terminal pro-brain natriuretic peptide as well as troponin. These cardiac biomarkers can be used for disease staging, prognostication, and disease activity monitoring in patients with AL amyloidosis. Notably, renal insufficiency can falsely elevate levels of these biomarkers. Recently, biomarkers of cardiac remodeling—i.e., matrix metalloproteinases and tissue inhibitors of metalloproteinases—have been found to be altered in the serum of patients with amyloid cardiomyopathy. Electrocardiographic and echocardiographic features of amyloid cardiomyopathy are described below. Patients with liver involvement, even when advanced, usually develop cholestasis with an elevated alkaline phosphatase concentration but minimal alteration of the aminotransferases and preservation of synthetic function. In AL amyloidosis, endocrine organs may be infiltrated with fibrils, and hypothyroidism, hypoadrenalism, or even hypopituitarism can occur. Although none of these findings is specific for amyloidosis, the presence of abnormalities in multiple organ systems should raise suspicion regarding this diagnosis.

AL AMYLOIDOSIS

Etiology and Incidence AL amyloidosis is most frequently caused by a clonal expansion of bone-marrow plasma cells that secrete a monoclonal immunoglobulin LC depositing as amyloid fibrils in tissues. Whether the clonal plasma

cells produce an LC that misfolds and leads to AL amyloidosis or an LC that folds properly, allowing the cells to inexorably expand over time and develop into multiple myeloma (Chap. 136), may depend upon primary sequence or other genetic or epigenetic factors. AL amyloidosis can occur with multiple myeloma or other B lymphoproliferative diseases, including non-Hodgkin's lymphoma (Chap. 134) and Waldenström's macroglobulinemia (Chap. 136). AL amyloidosis is the most common type of systemic amyloidosis diagnosed in North America. Its incidence has been estimated at 4.5 cases/100,000 population; however, ascertainment continues to be inadequate, and the true incidence may be much higher. AL amyloidosis, like other plasma cell diseases, usually occurs after age 40 and is often rapidly progressive and fatal if untreated.

Pathology and Clinical Features Amyloid deposits are usually widespread in AL amyloidosis and can be present in the interstitium of any organ outside the central nervous system. The amyloid fibril deposits are composed of full-length 23-kDa monoclonal immunoglobulin LCs as well as fragments. Accessory molecules co-deposited with LC fibrils (as well as with other amyloid fibrils) include serum amyloid P component, other proteins, glycosaminoglycans, and metal ions. Although all kappa and lambda LC subtypes have been identified in AL amyloid fibrils, lambda subtypes predominate. The lambda 6 subtype appears to have unique structural properties that predispose it to fibril formation, often in the kidney.

AL amyloidosis is often a rapidly progressive disease that presents as a pleiotropic set of clinical syndromes, recognition of which is key for initiation of the appropriate workup. Nonspecific symptoms of fatigue and weight loss are common; however, the diagnosis is rarely considered until symptoms referable to a specific organ develop. The kidneys are the most frequently involved organ and are affected in 70–80% of patients. Renal amyloidosis usually manifests as proteinuria, often in the nephrotic range and associated with hypoalbuminemia, secondary hypercholesterolemia and hypertriglyceridemia, and edema or anasarca. In some patients, interstitial rather than glomerular amyloid deposition can produce azotemia without proteinuria. The heart is the second most commonly affected organ (50–60% of patients), and cardiac involvement is the leading cause of death from AL amyloidosis. Early on, the electrocardiogram may show low voltage in the limb leads with a pseudo-infarct pattern. Echocardiographic features of disease include concentrically thickened ventricles and diastolic dysfunction with an abnormal strain pattern a “sparkly” appearance has been described but is often not seen with modern high-resolution echocardiographic techniques. Poor atrial contractility occurs even in sinus rhythm, and patients with cardiac amyloidosis are at risk for development of atrial thrombi and stroke. Cardiac MRI can show increased wall thickness, and characteristic enhancement of the sub-endocardium has been described following injection of gadolinium contrast. Nervous system symptoms include peripheral sensorimotor neuropathy and/or autonomic dysfunction manifesting as gastrointestinal motility disturbances (early satiety, diarrhea, constipation), impotence, orthostatic hypotension, and/or neurogenic bladder. Macroglossia (Fig. 137-2A), a pathognomonic sign of AL amyloidosis, is seen in only ~10% of patients. Liver involvement causes cholestasis

and hepatomegaly. The spleen is frequently involved, and there may be functional hyposplenism in the absence of significant splenomegaly. Many patients experience “easy bruising” due to amyloid deposits in capillaries or deficiency of clotting factor X, which can bind to amyloid fibrils; cutaneous ecchymoses appear, particularly around the eyes, producing the “raccoon-eye” sign (Fig. 137-2B), another uncommon but pathognomonic finding. Other findings include nail dystrophy (Fig. 137-2C), alopecia, and amyloid arthropathy with thickening of synovial membranes in the wrists and shoulders. The presence of a multisystemic illness or general fatigue along with any of these clinical syndromes should prompt a workup for amyloidosis.

Diagnosis Identification of an underlying clonal plasma cell or B lymphoproliferative process and a clonal LC are key to the diagnosis of AL amyloidosis. Serum protein electrophoresis and urine protein electrophoresis, although of value in multiple myeloma, are *not* useful screening tests if AL amyloidosis is suspected because the clonal LC or whole immunoglobulin often is not present in sufficient amounts to produce a monoclonal “M-spike” in the serum or LC (Bence Jones) protein in the urine. However, more than 90% of patients with AL amyloidosis have serum or urine monoclonal LC or whole immunoglobulin detectable by immunofixation electrophoresis of serum (SIFE) or urine (UIFE) (Fig. 137-3A) or by nephelometric measurement of “free” LCs (i.e., LCs circulating in monomeric form rather than in an immunoglobulin tetramer with heavy chain). Examining the ratio as well as the absolute amount of free LCs is essential, as renal insufficiency reduces LC clearance, elevating both isotypes. In addition, an increased percentage of plasma cells in the bone marrow—typically 5–30% of nucleated cells—is found in ~90% of patients. Kappa or lambda clonality should be demonstrated by flow cytometry, immunohistochemistry, or in situ hybridization for LC mRNA (Fig. 137-3B).

A monoclonal serum protein by itself is not diagnostic of amyloidosis, since monoclonal gammopathy of uncertain significance is common in older patients (Chap. 136). However, when monoclonal gammopathy of uncertain significance is found in patients with biopsy-proven amyloidosis, the AL type should be strongly suspected. Similarly, patients thought to have “smoldering myeloma” because of a modest elevation of bone-marrow plasma cells should be screened for AL amyloidosis if they have signs or symptoms of renal, cardiac, or neurologic disease. Accurate tissue amyloid typing is essential for appropriate treatment. Immunohistochemical staining of the amyloid deposits is useful if they bind one LC antibody in preference to the other; some AL deposits bind antibodies nonspecifically. Immunoelectron microscopy is more reliable, and mass spectrometry-based microsequencing of small amounts of protein extracted from fibril deposits can also be undertaken. In ambiguous cases, other forms of amyloidosis should be thoroughly excluded with appropriate genetic and other testing.

TREATMENT AL AMYLOIDOSIS

Extensive multisystemic involvement typifies AL amyloidosis, and the median survival period without treatment is usually only ~1–2 years from the time of diagnosis. Current therapies target the



FIGURE 137-2 Clinical signs of AL amyloidosis. **A.** Macroglossia. **B.** Periorbital ecchymoses. **C.** Fingernail dystrophy.

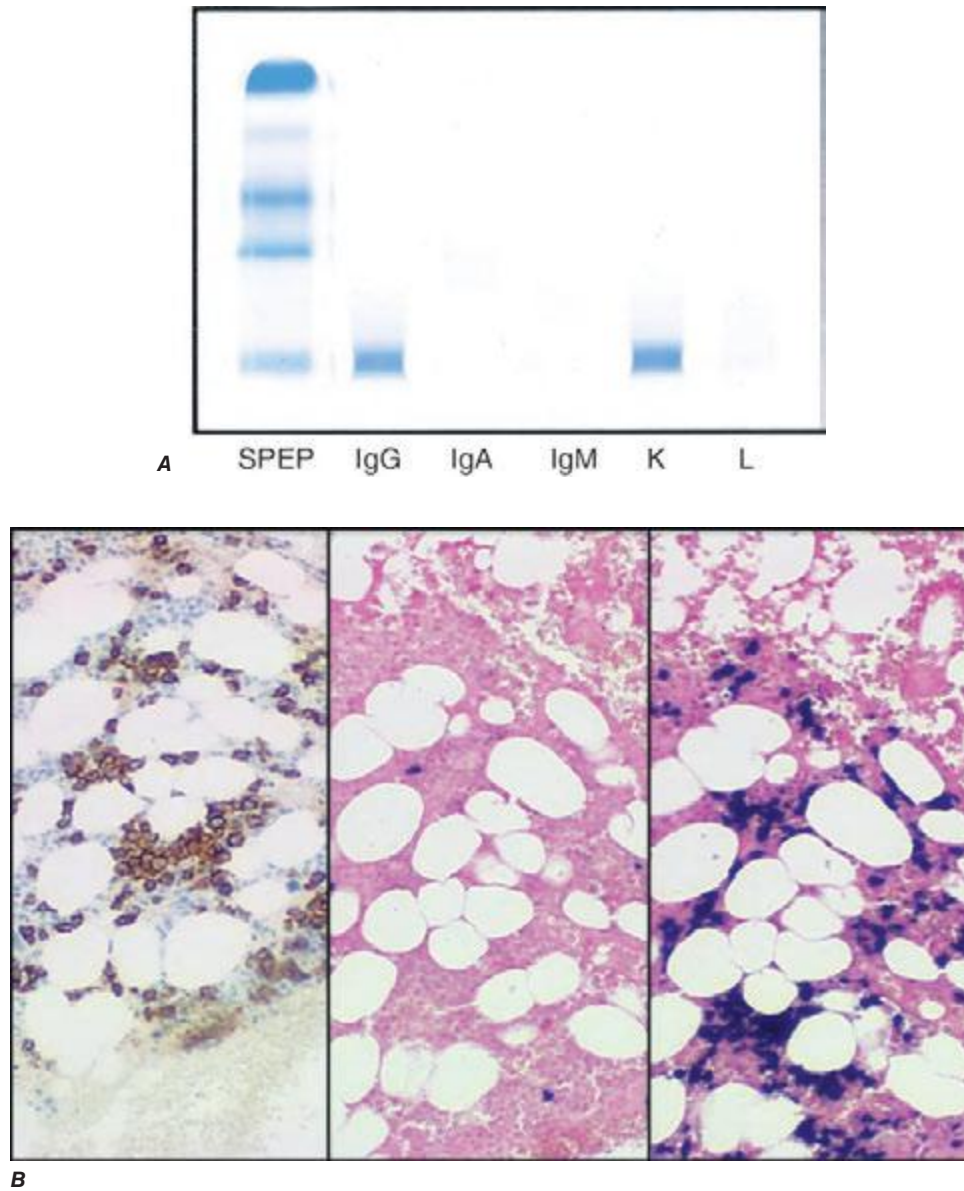


FIGURE 137-3 Laboratory features of AL amyloidosis. **A.** Serum immunofixation electrophoresis reveals an IgGκ monoclonal protein in this example; serum protein electrophoresis is often normal. **B.** Bone-marrow biopsy sections stained by immunohistochemistry with antibody to CD138 (syndecan, highly expressed on plasma cells) (left) or by in situ hybridization with fluorescein-tagged probes (Ventana Medical Systems) binding to κ mRNA (center) and λ mRNA (right) in plasma cells. (Photomicrograph courtesy of C. O'Hara; with permission.)

clonal bone-marrow plasma cells, using approaches employed for multiple myeloma. Treatment with oral melphalan and prednisone can decrease the plasma cell burden but rarely leads to complete hematologic remission, meaningful organ responses, or improved survival and is no longer widely used. The substitution of dexamethasone for prednisone produces a higher response rate and more durable remissions, although dexamethasone is not always well tolerated by patients with significant edema or cardiac disease. High-dose IV melphalan followed by autologous stem cell transplantation (HDM/SCT) produces complete hematologic responses in ~40% of treated patients, as determined by loss of clonal plasma cells in the bone marrow and disappearance of the monoclonal LC, as determined by SIFE/UIFE and free LC quantitation. Hematologic responses can be followed in the subsequent 6–12 months as improvements in organ function and quality of life. Hematologic responses appear to be more durable after HDM/SCT than in multiple myeloma, with remissions continuing in some patients beyond 15 years without additional treatment. Unfortunately, only about half of AL amyloidosis patients are suitable for aggressive treatment, and, even at specialized treatment centers, transplantation-related mortality rates are higher than those for other hematologic diseases

because of impaired organ function. Amyloid cardiomyopathy, poor nutritional and performance status, and multiorgan disease contribute to excess morbidity and mortality. A bleeding diathesis resulting from adsorption of clotting factor X to amyloid fibrils also increases mortality rates; however, this syndrome occurs in only 5–10% of patients. A randomized multicenter trial conducted in France compared oral melphalan and dexamethasone with HDM/SCT and failed to show a benefit of dose-intensive treatment, although the transplantation-related mortality rate in this study was very high. It has become clear that careful selection of patients and expert peritransplantation management are essential in reducing transplantation-related mortality.

For patients with impaired cardiac function or arrhythmias due to amyloid involvement of the myocardium, the median survival period is only ~6 months without treatment. In these patients, cardiac transplantation can be performed and followed by HDM/SCT to eliminate the noxious clone and prevent amyloid deposition in the transplanted heart or other organs.

Novel anti-plasma cell agents have been investigated for treatment of plasma cell diseases. The immunomodulators thalidomide, lenalidomide, and pomalidomide display activity; dosing may need

to be adjusted compared to their usage for myeloma. The proteasome inhibitor bortezomib has also been found to be effective in single-center and multicenter trials. Anti-fibril small molecules and humanized monoclonal antibodies are also being tested. Clinical trials are essential in improving therapy for this rare disease.

Supportive care is important for patients with any type of amyloidosis. For nephrotic syndrome, diuretics and supportive stockings can ameliorate edema; angiotensin-converting enzyme inhibitors should be used with caution and have not been shown to slow renal disease progression. Effective diuresis can be facilitated with albumin infusions to raise intravascular oncotic pressure. Congestive heart failure due to amyloid cardiomyopathy is best treated with diuretics; it is important to note that digitalis, calcium channel blockers, and beta blockers are relatively contraindicated as they can interact with amyloid fibrils and produce heart block and worsening heart failure. Amiodarone has been used for atrial and ventricular arrhythmias. Automatic implantable defibrillators have reduced effectiveness due to the thickened myocardium, but they may benefit some patients. Atrial ablation is an effective approach for atrial fibrillation. For conduction abnormalities, ventricular pacing may be indicated. Atrial contractile dysfunction is common in amyloid cardiomyopathy and is an indication for anticoagulation even in the absence of atrial fibrillation. Autonomic neuropathy can be treated with α agonists such as midodrine to support the blood pressure; gastrointestinal dysfunction may respond to motility or bulk agents. Nutritional supplementation, either oral or parenteral, is also important.

In localized AL disease, amyloid deposits can be produced by clonal plasma cells infiltrating local sites in the airways, bladder, skin, or lymph nodes (Table 137-1). These deposits may respond to surgical intervention or low-dose radiation therapy (typically only 20 Gy); systemic treatment generally is not appropriate. Patients should be referred to a center familiar with management of these rare manifestations of amyloidosis.

AA AMYLOIDOSIS

Etiology and Incidence AA amyloidosis can occur in association with almost any chronic inflammatory state (e.g., rheumatoid arthritis, inflammatory bowel disease, familial Mediterranean fever [Chap. 392], or other periodic fever syndromes) or chronic infections such as tuberculosis or subacute bacterial endocarditis. In the United States and Europe, AA amyloidosis has become less common, occurring in fewer than 2% of patients with these diseases, presumably because of advances in anti-inflammatory and antimicrobial therapies. It has also been described in association with Castleman's disease, and patients with AA amyloidosis should undergo CT scanning to look for such tumors as well as serologic and microbiologic studies. AA amyloidosis can also be seen without any identifiable underlying disease. AA is the only type of systemic amyloidosis that occurs in children.

Pathology and Clinical Features Organ involvement in AA amyloidosis usually begins in the kidneys. Hepatomegaly, splenomegaly, and autonomic neuropathy can also occur as the disease progresses; cardiomyopathy occurs, albeit rarely. The symptoms and signs of AA disease cannot be reliably distinguished from those of AL amyloidosis. AA amyloid fibrils are usually composed of an 8-kDa, 76-amino-acid N-terminal portion of the 12-kDa precursor protein SAA. This acute-phase apoprotein is synthesized in the liver and transported by high-density lipoprotein (HDL3) in the plasma. Several years of an underlying inflammatory disease causing chronic elevation of SAA levels usually precede fibril formation, although infections can lead to AA deposition more rapidly.

TREATMENT AA AMYLOIDOSIS

Primary therapy for AA amyloidosis consists of treatment of the underlying inflammatory or infectious disease. Treatment that suppresses or eliminates the inflammation or infection also decreases the SAA concentration. For familial Mediterranean fever, colchicine

at a dose of 1.2–1.8 mg/d is the standard treatment. However, colchicine has not been helpful for AA amyloidosis of other causes or for other amyloidoses. Tumor necrosis factor and interleukin 1 antagonists can be effective in syndromes related to cytokine elevation. For this disease, there is also a fibril-specific agent: eprodisate was designed to interfere with the interaction of AA amyloid protein with glycosaminoglycans and to prevent or disrupt fibril formation. The drug is well tolerated and delays progression of AA renal disease. Randomized phase III clinical trials with eprodisate are ongoing; the drug is not otherwise available.

ATTR AND AF AMYLOIDOSIS



The familial amyloidoses are autosomal dominant diseases in which, beginning in midlife, a variant (FINE) plasma protein forms amyloid deposits. These diseases are rare, with an estimated incidence of <1 case/100,000 population in the United States, although founder effects in isolated areas of Portugal, Sweden, and Japan have led to a much higher incidence. The most common form of AF amyloidosis is ATTRm in the updated nomenclature, caused by mutation of the abundant plasma protein transthyretin (TTR, also known as *prealbumin*). More than 100 TTR mutations are known, and most are associated with ATTR amyloidosis. One variant, V122I, has a carrier frequency that may be as high as 4% in the African-American population and is associated with late-onset cardiac amyloidosis. The actual incidence and penetrance of disease in the African-American population is the subject of ongoing research, but ATTR amyloidosis warrants consideration in the differential diagnosis of African-American patients who present with concentric cardiac hypertrophy and evidence of diastolic dysfunction, particularly in the absence of a history of hypertension. Other familial amyloidoses, caused by variant apolipoproteins AI or AII, gelsolin, fibrinogen A α , or lysozyme, are reported in only a few families worldwide. New amyloidogenic serum proteins continue to be identified periodically, including recently the leukocyte chemotactic factor LECT2, a cause of renal amyloidosis in Hispanic and Pakistani populations. To date, no mutation in the coding sequence for the LECT2 gene has been identified, so the heritability of ALECT2 is uncertain.

TTR deposits composed of unmutated fibrils occur with aging, and ATTRwt is being diagnosed with increasing frequency in Caucasian men >65 years of age with amyloid cardiomyopathy. Formerly termed senile systemic amyloidosis, ATTRwt has been found at autopsy in 25% of hearts from patients older than age 80 years. Why a wild type protein becomes amyloidogenic, and why patients bearing mutant TTR genes do not express disease until adulthood, remains a mystery.

Clinical Features and Diagnosis AF amyloidosis has a presentation that is variable but is usually consistent within kindreds affected by the same mutant protein. A family history makes AF disease more likely, but many patients present sporadically with new mutations. ATTR amyloidosis typically presents as a syndrome of familial amyloidotic polyneuropathy or familial amyloidotic cardiomyopathy. Peripheral neuropathy begins as a small-fiber lower-extremity sensory and motor neuropathy and progresses to the upper extremities. Autonomic neuropathy manifests as diarrhea with weight loss and orthostatic hypotension. Patients with TTR V30M, the most common mutation, have normal electrocardiograms but may develop conduction defects late in the disease. Patients with TTR T60A and several other mutations have myocardial thickening similar to that caused by AL amyloidosis, although heart failure is less common and long-term survival rates are usually better. Vitreous opacities caused by amyloid deposits are pathognomonic for ATTR amyloidosis.

Typical syndromes associated with other forms of AF disease include renal amyloidosis with mutant fibrinogen, lysozyme, or apolipoproteins; hepatic amyloidosis with apolipoprotein AI; and amyloidosis of cranial nerves and cornea with gelsolin. Patients with AF amyloidosis can present with clinical syndromes that mimic those of patients with AL disease. Rarely, AF carriers can develop AL disease or AF patients may have monoclonal gammopathy without AL.

724 Thus, it is important to screen both for plasma cell disorders and for mutations in patients with amyloidosis. Variant TTRs can usually be detected by isoelectric focusing, but DNA sequencing is now standard for diagnosis of ATTR and other AF mutations.

TREATMENT ATTR AMYLOIDOSIS

Without intervention, the survival period after onset of ATTR disease is 5–15 years. Orthotopic liver transplantation replaces the major source of variant TTR production with a source of normal TTR. While liver transplantation can slow disease progression and improve chances of survival, it does not reverse sensorimotor neuropathy. Liver transplants are most successful in young patients with early peripheral neuropathy; older patients with familial amyloidotic cardiomyopathy or advanced polyneuropathy often experience end-organ disease progression despite successful liver transplantation. Progressive disease has been attributed to accumulation of wild-type TTR in fibrillar deposits initiated by the mutant.

The rate-limiting step in ATTR amyloidosis is dissociation of the TTR tetramer into monomer followed by misfolding and aggregation. TTR tetramers can be stabilized by thyroxine binding or by small molecules such as the non-steroidal anti-inflammatory drug diflunisal or the rationally designed small-molecule therapeutic tafamidis. A placebo-controlled randomized trial of diflunisal demonstrated a reduction in the progression of polyneuropathy and maintenance of quality of life in patients with a wide variety of ATTR mutations who received the “repurposed” diflunisal. Tafamidis tested in a similar fashion in patients with the V30M ATTR mutation failed to meet its primary endpoints, but tafamidis was approved by the European Medicines Agency since most secondary endpoints favored the drug. These agents are now being investigated for effects on cardiomyopathy, and in ATTRwt. *In vitro* data and serendipitous observations in patients suggest that ATTRm disease can be ameliorated by “trans-suppression,” in which a T119M TTR variant stabilizes tetramers that also contain amyloidogenic subunits. Interestingly, in a large population study in Denmark, 0.5% of participants were heterozygous for the T119M allele, and this small group had higher levels of TTR in their blood, a reduced incidence of cerebrovascular disease, and a 5- to 10-year survival advantage compared with participants lacking this allele.

A β_2 M AMYLOIDOSIS

A β_2 M amyloid is composed of β_2 -microglobulin, the invariant chain of class I human leukocyte antigens, and produces rheumatologic manifestations in patients undergoing long-term hemodialysis. β_2 -Microglobulin is excreted by the kidney, and levels become elevated in end-stage renal disease. The molecular mass of β_2 M is 11.8 kDa—above the cutoff of some dialysis membranes. The incidence of this disease appears to be declining with the use of newer membranes in high-flow dialysis techniques. A β_2 M amyloidosis usually presents as carpal tunnel syndrome, persistent joint effusions, spondyloarthropathy, or cystic bone lesions. Carpal tunnel syndrome is often the first symptom. In the past, persistent joint effusions accompanied by mild discomfort were found in up to 50% of patients who had undergone dialysis for >12 years. Involvement is bilateral, and large joints (shoulders, knees, wrists, and hips) are most frequently affected. The synovial fluid is noninflammatory, and β_2 M amyloid can be found if the sediment is stained with Congo red. Although less common, visceral β_2 M amyloid deposits do occasionally occur in the gastrointestinal tract, heart, tendons, and subcutaneous tissues of the buttocks. There is no specific therapy for A β_2 M amyloidosis, but cessation of dialysis after renal allografting may lead to symptomatic improvement.

SUMMARY

A diagnosis of amyloidosis should be considered in patients with unexplained nephropathy, cardiomyopathy (particularly with diastolic

dysfunction), neuropathy (either peripheral or autonomic), enteropathy, or the pathognomonic soft tissue findings of macroglossia or periorbital ecchymoses. Pathologic identification of amyloid fibrils can be made with Congo red staining of aspirated abdominal fat or of an involved-organ biopsy specimen. Accurate typing by a combination of immunologic, biochemical, and genetic testing is essential in selecting appropriate therapy (Fig. 137-1). Systemic amyloidosis should not be considered an untreatable condition, as anti-plasma cell chemotherapy is highly effective in AL disease and targeted therapies are being developed for AA and ATTR disease. Tertiary referral centers can provide specialized diagnostic techniques and access to clinical trials for patients with these rare diseases.

ACKNOWLEDGMENT

This chapter represents a revised version of a chapter that was co-authored by Dr. Martha Skinner and Dr. David Seldin in previous editions of Harrison's Principles of Internal Medicine.

BLOOD GROUP ANTIGENS AND ANTIBODIES

The study of red blood cell (RBC) antigens and antibodies forms the foundation of transfusion medicine. Serologic studies initially characterized these antigens, but now the molecular composition and structure of many are known. Antigens, either carbohydrate or protein, are assigned to a blood group system based on the structure and similarity of the determinant epitopes. Other cellular blood elements and plasma proteins are also antigenic and can result in *alloimmunization*, the production of antibodies directed against the blood group antigens of another individual. These antibodies are called *alloantibodies*.

Antibodies directed against RBC antigens may result from “natural” exposure, particularly to carbohydrates that mimic some blood group antigens. Those antibodies that occur via natural stimuli are usually produced by a T cell-independent response (thus, generating no memory) and are IgM isotype. *Autoantibodies* (antibodies against autologous blood group antigens) arise spontaneously or as the result of infectious sequelae (e.g., from *Mycoplasma pneumoniae*) and are also often IgM. These antibodies are often clinically insignificant due to their low affinity for antigen at body temperature. However, IgM antibodies can activate the complement cascade and result in hemolysis. Antibodies that result from allogeneic exposure, such as transfusion or pregnancy, are usually IgG. IgG antibodies commonly bind to antigen at warmer temperatures and may hemolyze RBCs. Unlike IgM antibodies, IgG antibodies can cross the placenta and bind fetal erythrocytes bearing the corresponding antigen, resulting in hemolytic disease of the newborn, or *hydrops fetalis*.

Alloimmunization to leukocytes, platelets, and plasma proteins may also result in transfusion complications such as fevers and urticaria but generally does not cause hemolysis. Assay for these other alloantibodies is not routinely performed; however, they may be detected using special assays.

ABO ANTIGENS AND ANTIBODIES

The first blood group antigen system, recognized in 1900, was ABO, the most important in transfusion medicine. The major blood groups of this system are A, B, AB, and O. O type RBCs lack A or B antigens. These antigens are carbohydrates attached to a precursor backbone, may be found on the cellular membrane either as glycosphingolipids or glycoproteins, and are secreted into plasma and body fluids as glycoproteins. H substance is the immediate precursor on which the A and B antigens are added. This H substance is formed by the addition of fucose to the glycolipid or glycoprotein backbone. The subsequent addition of *N*-acetylgalactosamine creates the A antigen, whereas the addition of galactose produces the B antigen.

The genes that determine the A and B phenotypes are found on chromosome 9p and are expressed in a Mendelian codominant manner. The gene products are glycosyl transferases, which confer the enzymatic capability of attaching the specific antigenic carbohydrate. Individuals who lack the “A” and “B” transferases are phenotypically type “O,” whereas those who inherit both transferases are type “AB.” Rare individuals lack the H gene, which codes for fucose transferase, and cannot form H substance. These individuals are homozygous for the silent h allele (hh) and have Bombay phenotype (O_h).

The ABO blood group system is important because essentially all individuals produce antibodies to the ABH carbohydrate antigen that they lack. The naturally occurring anti-A and anti-B antibodies are termed *isoagglutinins*. Thus, type A individuals produce anti-B, whereas type B individuals make anti-A. Neither isoagglutinin is found in type AB individuals, whereas type O individuals produce both anti-A and anti-B. Thus, persons with type AB are “universal recipients” because they do not have antibodies against any ABO phenotype,

whereas persons with type O blood can donate to essentially all recipients because their cells are not recognized by any ABO isoagglutinins. The rare individuals with Bombay phenotype produce antibodies to H substance (which is present on all red cells except those of hh phenotype) as well as to both A and B antigens and are therefore compatible only with other hh donors.

In most people, A and B antigens are secreted by the cells and are present in the circulation. Nonsecretors are susceptible to a variety of infections (e.g., *Candida albicans*, *Neisseria meningitidis*, *Streptococcus pneumoniae*, *Haemophilus influenzae*) because many organisms may bind to polysaccharides on cells. Soluble blood group antigens may block this binding.

RH SYSTEM

The Rh system is the second most important blood group system in pretransfusion testing. The Rh antigens are found on a 30- to 32-kDa RBC membrane protein that has no defined function. Although >40 different antigens in the Rh system have been described, five determinants account for the vast majority of phenotypes. The presence of the D antigen confers Rh “positivity,” whereas persons who lack the D antigen are Rh negative. Two allelic antigen pairs, E/e and C/c, are also found on the Rh protein. The three Rh genes, E/e, D, and C/c, are arranged in tandem on chromosome 1 and inherited as a haplotype, i.e., cDE or Cde. Two haplotypes can result in the phenotypic expression of two to five Rh antigens.

The D antigen is a potent alloantigen. About 15% of individuals lack this antigen. Exposure of these Rh-negative people to even small amounts of Rh-positive cells, by either transfusion or pregnancy, can result in the production of anti-D alloantibody.

OTHER BLOOD GROUP SYSTEMS AND ALLOANTIBODIES

More than 100 blood group systems are recognized, composed of more than 500 antigens. The presence or absence of certain antigens has been associated with various diseases and anomalies; antigens also act as receptors for infectious agents. Alloantibodies of importance in routine clinical practice are listed in [Table 138e-1](#).

Antibodies to *Lewis system* carbohydrate antigens are the most common cause of incompatibility during pretransfusion screening. The Lewis gene product is a fucosyl transferase and maps to chromosome 19. The antigen is not an integral membrane structure but is adsorbed to the RBC membrane from the plasma. Antibodies to Lewis antigens are usually IgM and cannot cross the placenta. Lewis antigens may be adsorbed onto tumor cells and may be targets of therapy.

I system antigens are also oligosaccharides related to H, A, B, and Le. I and i are not allelic pairs but are carbohydrate antigens that differ only in the extent of branching. The i antigen is an unbranched chain that is converted by the I gene product, a glycosyltransferase, into a branched chain. The branching process affects all the ABH antigens, which become progressively more branched in the first 2 years of life. Some patients with cold agglutinin disease or lymphomas can produce anti-I autoantibodies that cause RBC destruction. Occasional patients with mononucleosis or *Mycoplasma pneumoniae* may develop cold agglutinins of either anti-I or anti-i specificity. Most adults lack i

TABLE 138e-1 RBC BLOOD GROUP SYSTEMS AND ALLOANTIGENS

Blood Group System	Antigen	Alloantibody	Clinical Significance
Rh (D, C/c, E/e)	RBC protein	IgG	HTR, HDN
Lewis (Le ^a , Le ^b)	Oligosaccharide	IgM/IgG	Rare HTR
Kell (K/k)	RBC protein	IgG	HTR, HDN
Duffy (Fy ^a /Fy ^b)	RBC protein	IgG	HTR, HDN
Kidd (Jk ^a /Jk ^b)	RBC protein	IgG	HTR (often delayed), HDN (mild)
I/i	Carbohydrate	IgM	None
MNSsU	RBC protein	IgM/IgG	Anti-M rare HDN, anti-S, -s, and -U HDN, HTR

Abbreviations: HDN, hemolytic disease of the newborn; HTR, hemolytic transfusion reaction; RBC, red blood cell.

expression; thus, finding a donor for patients with anti-i is not difficult. Even though most adults express I antigen, binding is generally low at body temperature. Thus, administration of warm blood prevents isoagglutination.

The *P* system is another group of carbohydrate antigens controlled by specific glycosyltransferases. Its clinical significance is in rare cases of syphilis and viral infection that lead to paroxysmal cold hemoglobinuria. In these cases, an unusual autoantibody to P is produced that binds to RBCs in the cold and fixes complement upon warming. Antibodies with these biphasic properties are called *Donath-Landsteiner antibodies*. The P antigen is the cellular receptor of parvovirus B19 and also may be a receptor for *Escherichia coli* binding to urothelial cells.

The *MNSsU* system is regulated by genes on chromosome 4. M and N are determinants on glycophorin A, an RBC membrane protein, and S and s are determinants on glycophorin B. Anti-S and anti-s IgG antibodies may develop after pregnancy or transfusion and lead to hemolysis. Anti-U antibodies are rare but problematic; virtually every donor is incompatible because nearly all persons express U.

The *Kell* protein is very large (720 amino acids), and its secondary structure contains many different antigenic epitopes. The immunogenicity of Kell is third behind the ABO and Rh systems. The absence of the Kell precursor protein (controlled by a gene on X) is associated with acanthocytosis, shortened RBC survival, and a progressive form of muscular dystrophy that includes cardiac defects. This rare condition is called the *McLeod phenotype*. The K_x gene is linked to the 91-kDa component of the NADPH-oxidase on the X chromosome, deletion or mutation of which accounts for about 60% of cases of chronic granulomatous disease.

The *Duffy* antigens are codominant alleles, Fy^a and Fy^b , that also serve as receptors for *Plasmodium vivax*. More than 70% of persons in malaria-endemic areas lack these antigens, probably from selective influences of the infection on the population. For unknown reasons, the lack of the Duffy antigen receptor for cytokines (DARC) is associated with mild neutropenia.

The *Kidd* antigens, Jk^a and Jk^b , may elicit antibodies transiently. A delayed hemolytic transfusion reaction that occurs with blood tested as compatible is often related to delayed appearance of anti- Jk^k .

PRETRANSFUSION TESTING

Pretransfusion testing of a potential recipient consists of the “type and screen.” The “forward type” determines the ABO and Rh phenotype of the recipient’s RBC by using antisera directed against the A, B, and D antigens. The “reverse type” detects isoagglutinins in the patient’s serum and should correlate with the ABO phenotype, or forward type.

The alloantibody screen identifies antibodies directed against other RBC antigens. The alloantibody screen is performed by mixing patient serum with type O RBCs that contain the major antigens of most blood group systems and whose extended phenotype is known. The specificity of the alloantibody is identified by correlating the presence or absence of antigen with the results of the agglutination.

Cross-matching is ordered when there is a high probability that the patient will require a packed RBC (PRBC) transfusion. Blood selected for cross-matching must be ABO compatible and lack antigens for which the patient has alloantibodies. Nonreactive cross-matching confirms the absence of any major incompatibility and reserves that unit for the patient.

In the case of Rh-negative patients, every attempt must be made to provide Rh-negative blood components to prevent alloimmunization to the D antigen. In an emergency, Rh-positive blood can be safely transfused to an Rh-negative patient who lacks anti-D; however, the recipient is likely to become alloimmunized and produce anti-D. Rh-negative women of childbearing age who are

transfused with products containing Rh-positive RBCs should receive passive immunization with anti-D (RhoGam or WinRho) to reduce or prevent sensitization.

BLOOD COMPONENTS

Blood products intended for transfusion are routinely collected as whole blood (450 mL) in various anticoagulants. Most donated blood is processed into components: PRBCs, platelets, and fresh-frozen plasma (FFP) or cryoprecipitate (Table 138e-2). Whole blood is first separated into PRBCs and platelet-rich plasma by slow centrifugation. The platelet-rich plasma is then centrifuged at high speed to yield one unit of random donor (RD) platelets and one unit of FFP. Cryoprecipitate is produced by thawing FFP to precipitate the plasma proteins and then separated by centrifugation.

Apheresis technology is used for the collection of multiple units of platelets from a single donor. These single-donor apheresis platelets (SDAP) contain the equivalent of at least six units of RD platelets and have fewer contaminating leukocytes than pooled RD platelets.

Plasma may also be collected by apheresis. Plasma derivatives such as albumin, intravenous immunoglobulin, antithrombin, and coagulation factor concentrates are prepared from pooled plasma from many donors and are treated to eliminate infectious agents.

WHOLE BLOOD

Whole blood provides both oxygen-carrying capacity and volume expansion. It is the ideal component for patients who have sustained acute hemorrhage of $\geq 25\%$ total blood volume loss. Whole blood is stored at 4°C to maintain erythrocyte viability, but platelet dysfunction and degradation of some coagulation factors occurs. In addition, 2,3-bisphosphoglycerate levels fall over time, leading to an increase in the oxygen affinity of the hemoglobin and a decreased capacity to deliver oxygen to the tissues, a problem with all red cell storage. Fresh whole blood avoids these problems, but it is typically used only in emergency settings (i.e., military). Whole blood is not readily available, because it is routinely processed into components.

PACKED RED BLOOD CELLS

This product increases oxygen-carrying capacity in the anemic patient. Adequate oxygenation can be maintained with a hemoglobin content of 70 g/L in the normovolemic patient without cardiac disease; however, comorbid factors may necessitate transfusion at a higher threshold. The decision to transfuse should be guided by the clinical situation and not by an arbitrary laboratory value. In the critical care setting, liberal use of transfusions to maintain near-normal levels of hemoglobin has not proven advantageous. In most patients requiring transfusion, levels of hemoglobin of 100 g/L are sufficient to keep oxygen supply from being critically low.

PRBCs may be modified to prevent certain adverse reactions. The majority of cellular blood products are now leukocyte reduced, and universal prestorage leukocyte reduction has been recommended. Prestorage filtration appears superior to bedside filtration as smaller amounts of cytokines are generated in the stored product. These PRBC units contain $< 5 \times 10^6$ donor white blood cells (WBCs), and their use

TABLE 138e-2 CHARACTERISTICS OF SELECTED BLOOD COMPONENTS

Component	Volume, mL	Content	Clinical Response
PRBC	180–200	RBCs with variable leukocyte content and small amount of plasma	Increase hemoglobin 10 g/L and hematocrit 3%
Platelets	50–70	5.5×10^{10} /RD unit	Increase platelet count 5000–10,000/ μ L
	200–400	$\geq 3 \times 10^{11}$ /SDAP product	CCI $\geq 10 \times 10^9$ /L within 1 h and $\geq 7.5 \times 10^9$ /L within 24 h posttransfusion
FFP	200–250	Plasma proteins—coagulation factors, proteins C and S, antithrombin	Increases coagulation factors about 2%
Cryoprecipitate	10–15	Cold-insoluble plasma proteins, fibrinogen, factor VIII, VWF	Topical fibrin glue, also 80 IU factor VIII

Abbreviations: CCI, corrected count increment; FFP, fresh-frozen plasma; PRBC, packed red blood cells; RBC, red blood cell; RD, random donor; SDAP, single-donor apheresis platelets; VWF, von Willebrand factor.

lowers the incidence of posttransfusion fever, cytomegalovirus (CMV) infections, and alloimmunization. Other theoretical benefits include less immunosuppression in the recipient and lower risk of infections. Plasma, which may cause allergic reactions, can be removed from cellular blood components by washing.

PLATELETS

Thrombocytopenia is a risk factor for hemorrhage, and platelet transfusion reduces the incidence of bleeding. The threshold for prophylactic platelet transfusion is 10,000/ μ L. In patients without fever or infections, a threshold of 5000/ μ L may be sufficient to prevent spontaneous hemorrhage. For invasive procedures, the usual target level is 50,000/ μ L platelets.

Platelets are given either as pools prepared from RDs or as SDAPs from a single donor. In an unsensitized patient without increased platelet consumption (splenomegaly, fever, disseminated intravascular coagulation [DIC]), two units of transfused RD per square-meter body surface area (BSA) is anticipated to increase the platelet count by approximately 10,000/ μ L. Patients who have received multiple transfusions may be alloimmunized to many HLA- and platelet-specific antigens and have little or no increase in their posttransfusion platelet counts. Patients who may require multiple transfusions are best served by receiving SDAP and leukocyte-reduced components to lower the risk of alloimmunization.

Refractoriness to platelet transfusion may be evaluated using the corrected count increment (CCI):

$$\text{CCI} = \frac{\text{posttransfusion count}(\mu\text{L}) - \text{pretransfusion count}(\mu\text{L})}{\text{number of platelets transfused} \times 10^{-11}} \times \text{BSA} (\text{m}^2)$$

where BSA is body surface area measured in square meters. The platelet count performed 1 h after the transfusion is acceptable if the CCI is $10 \times 10^9/\text{mL}$, and after 18–24 h an increment of $7.5 \times 10^9/\text{mL}$ is expected. Patients who have suboptimal responses are likely to have received multiple transfusions and have antibodies directed against class I HLA antigens. Refractoriness can be investigated by detecting anti-HLA antibodies in the recipient's serum. Patients who are sensitized will often react with 100% of the lymphocytes used for the HLA-antibody screen, and HLA-matched SDAPs should be considered for patients who require transfusion. Although ABO-identical HLA-matched SDAPs provide the best chance for increasing the platelet count, locating these products is difficult. Platelet cross-matching is available in some centers. Additional clinical causes for a low platelet CCI include fever, bleeding, splenomegaly, DIC, or medications in the recipient.

FRESH-FROZEN PLASMA

FFP contains stable coagulation factors and plasma proteins: fibrinogen, antithrombin, albumin, and proteins C and S. Indications for FFP include correction of coagulopathies, including the rapid reversal of warfarin; supplying deficient plasma proteins; and treatment of thrombotic thrombocytopenic purpura. FFP should not be routinely used to expand blood volume. FFP is an acellular component and does not transmit intracellular infections, e.g., CMV. Patients who are IgA-deficient and require plasma support should receive FFP from IgA-deficient donors to prevent anaphylaxis (see below).

CRYOPRECIPITATE

Cryoprecipitate is a source of fibrinogen, factor VIII, and von Willebrand factor (VWF). It is ideal for supplying fibrinogen to the volume-sensitive patient. When factor VIII concentrates are not available, cryoprecipitate may be used because each unit contains approximately 80 units of factor VIII. Cryoprecipitate may also supply VWF to patients with dysfunctional (type II) or absent (type III) von Willebrand's disease.

PLASMA DERIVATIVES

Plasma from thousands of donors may be pooled to derive specific protein concentrates, including albumin, intravenous immunoglobulin, antithrombin, and coagulation factors. In addition, donors who

have high-titer antibodies to specific agents or antigens provide hyperimmune globulins, such as anti-D (RhoGam, WinRho), and antisera to hepatitis B virus (HBV), varicella-zoster virus, CMV, and other infectious agents.

ADVERSE REACTIONS TO BLOOD TRANSFUSION

Adverse reactions to transfused blood components occur despite multiple tests, inspections, and checks. Fortunately, the most common reactions are not life threatening, although serious reactions can present with mild symptoms and signs. Some reactions can be reduced or prevented by modified (filtered, washed, or irradiated) blood components. When an adverse reaction is suspected, the transfusion should be stopped and reported to the blood bank for investigation.

Transfusion reactions may result from immune and nonimmune mechanisms. Immune-mediated reactions are often due to preformed donor or recipient antibody; however, cellular elements may also cause adverse effects. Nonimmune causes of reactions are due to the chemical and physical properties of the stored blood component and its additives.

Transfusion-transmitted viral infections are increasingly rare due to improved screening and testing. As the risk of viral infection is reduced, the relative risk of other reactions increases, such as hemolytic transfusion reactions and sepsis from bacterially contaminated components. Pretransfusion quality assurance improvements further increase the safety of transfusion therapy. Infections, like any adverse transfusion reaction, must be brought to the attention of the blood bank for appropriate studies (Table 138e-3).

IMMUNE-MEDIATED REACTIONS

Acute Hemolytic Transfusion Reactions Immune-mediated hemolysis occurs when the recipient has preformed antibodies that lyse donor erythrocytes. The ABO isoagglutinins are responsible for the majority of these reactions. However, alloantibodies directed against other RBC antigens, i.e., Rh, Kell, and Duffy, are responsible for more fatal hemolytic transfusion reactions.

Acute hemolytic reactions may present with hypotension, tachypnea, tachycardia, fever, chills, hemoglobinemia, hemoglobinuria, chest and/or flank pain, and discomfort at the infusion site. Monitoring

TABLE 138e-3 RISKS OF TRANSFUSION COMPLICATIONS

Reactions	Frequency, Episodes: Unit
Febrile (FNHTR)	• 1–4:100
Allergic	• 1–4:100
Delayed hemolytic	• 1:1000
TRALI	• 1:5000
Acute hemolytic	• 1:12,000
Fatal hemolytic	• 1:100,000
Anaphylactic	• 1:150,000
Infections^a	
Hepatitis B	• 1:220,000
Hepatitis C	• 1:1,800,000
HIV-1, -2	• 1:2,300,000
HTLV-1 and -2	• 1:2,993,000
Malaria	• 1:4,000,000
Other Complications	
RBC allosensitization	• 1:100
HLA allosensitization	• 1:10
Graft-versus-host disease	Rare

^aInfectious agents rarely associated with transfusion, theoretically possible, or of unknown risk include West Nile virus, hepatitis A virus, parvovirus B19, *Babesia microti* and *Babesia duncani* (babesiosis), *Borrelia burgdorferi* (Lyme disease), *Anaplasma phagocytophilum* (human granulocytic ehrlichiosis), *Trypanosoma cruzi* (Chagas' disease), *Treponema pallidum*, and human herpesvirus-8.

Abbreviations: FNHTR, febrile nonhemolytic transfusion reaction; HTLV, human T lymphotropic virus; RBC, red blood cell; TRALI, transfusion-related acute lung injury.

138e-4 the patient's vital signs before and during the transfusion is important to identify reactions promptly. When acute hemolysis is suspected, the transfusion must be stopped immediately, intravenous access maintained, and the reaction reported to the blood bank. A correctly labeled posttransfusion blood sample and any untransfused blood should be sent to the blood bank for analysis. The laboratory evaluation for hemolysis includes the measurement of serum haptoglobin, lactate dehydrogenase (LDH), and indirect bilirubin levels.

The immune complexes that result in RBC lysis can cause renal dysfunction and failure. Diuresis should be induced with intravenous fluids and furosemide or mannitol. Tissue factor released from the lysed erythrocytes may initiate DIC. Coagulation studies including prothrombin time (PT), activated partial thromboplastin time (aPTT), fibrinogen, and platelet count should be monitored in patients with hemolytic reactions.

Errors at the patient's bedside, such as mislabeling the sample or transfusing the wrong patient, are responsible for the majority of these reactions. The blood bank investigation of these reactions includes examination of the pre- and posttransfusion samples for hemolysis and repeat typing of the patient samples; direct antiglobulin test (DAT), sometimes called the *direct Coombs test*, of the posttransfusion sample; repeating the cross-matching of the blood component; and checking all clerical records for errors. DAT detects the presence of antibody or complement bound to RBCs in vivo (Fig. 138e-1).

Delayed Hemolytic and Serologic Transfusion Reactions Delayed hemolytic transfusion reactions (DHTRs) are not completely preventable.

These reactions occur in patients previously sensitized to RBC alloantigens who have a negative alloantibody screen due to low antibody levels. When the patient is transfused with antigen-positive blood, an anamnestic response results in the early production of alloantibody that binds donor RBCs. The alloantibody is detectable 1–2 weeks following the transfusion, and the posttransfusion DAT may become positive due to circulating donor RBCs coated with antibody or complement. The transfused, alloantibody-coated erythrocytes are cleared by the reticuloendothelial system. These reactions are detected most commonly in the blood bank when a subsequent patient sample reveals a positive alloantibody screen or a new alloantibody in a recently transfused recipient.

No specific therapy is usually required, although additional RBC transfusions may be necessary. Delayed serologic transfusion reactions are similar to DHTR, because the DAT is positive and alloantibody is detected; however, RBC clearance is not increased.

Febrile Nonhemolytic Transfusion Reaction The most frequent reaction associated with the transfusion of cellular blood components is a febrile nonhemolytic transfusion reaction (FNHTR). These reactions are characterized by chills and rigors and a $\geq 1^\circ\text{C}$ rise in temperature. FNHTR is diagnosed when other causes of fever in the transfused patient are ruled out. Antibodies directed against donor leukocyte and HLA antigens may mediate these reactions; thus, multiply transfused patients and multiparous women are felt to be at increased risk. Although anti-HLA antibodies may be demonstrated in the recipient's serum, investigation is not routinely done because of the mild nature

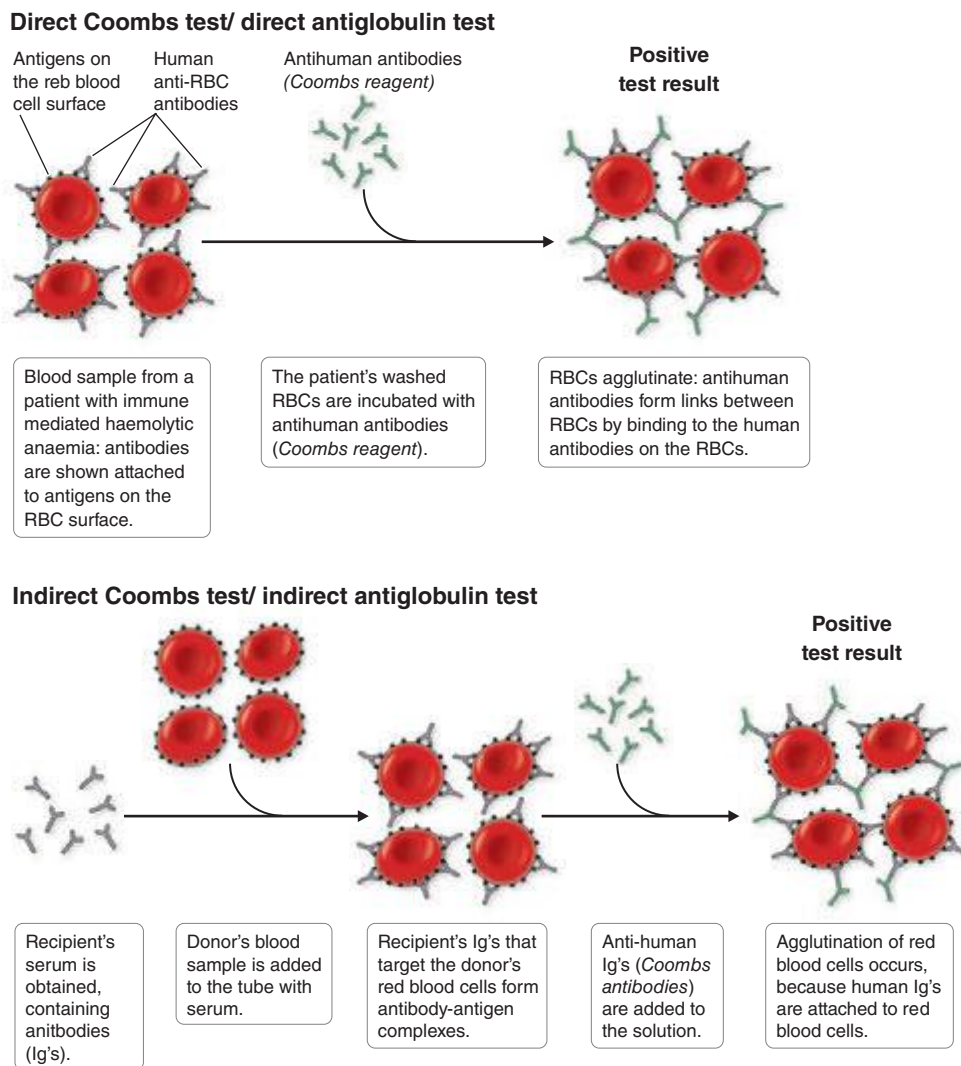


FIGURE 138E-1 Direct and indirect Coombs test. The direct Coombs (antiglobulin) test detects the presence of antibodies (or complement) on the surface of erythrocytes. The indirect Coombs (antiglobulin) test detects antibodies in the serum that may bind to donor erythrocytes. RBC, red blood cell. (Adapted from http://upload.wikimedia.org/wikipedia/commons/1/1c/coombs_test_schematic.png.)

of most FNHTR. The use of leukocyte-reduced blood products may prevent or delay sensitization to leukocyte antigens and thereby reduce the incidence of these febrile episodes. Cytokines released from cells within stored blood components may mediate FNHTR; thus, leukoreduction before storage may prevent these reactions.

Allergic Reactions Urticarial reactions are related to plasma proteins found in transfused components. Mild reactions may be treated symptomatically by temporarily stopping the transfusion and administering antihistamines (diphenhydramine, 50 mg orally or intramuscularly). The transfusion may be completed after the signs and/or symptoms resolve. Patients with a history of allergic transfusion reaction should be premedicated with an antihistamine. Cellular components can be washed to remove residual plasma for the extremely sensitized patient.

Anaphylactic Reaction This severe reaction presents after transfusion of only a few milliliters of the blood component. Symptoms and signs include difficulty breathing, coughing, nausea and vomiting, hypotension, bronchospasm, loss of consciousness, respiratory arrest, and shock. Treatment includes stopping the transfusion, maintaining vascular access, and administering epinephrine (0.5–1 mL of 1:1000 dilution subcutaneously). Glucocorticoids may be required in severe cases.

Patients who are IgA-deficient, <1% of the population, may be sensitized to this Ig class and are at risk for anaphylactic reactions associated with plasma transfusion. Individuals with severe IgA deficiency should therefore receive only IgA-deficient plasma and washed cellular blood components. Patients who have anaphylactic or repeated allergic reactions to blood components should be tested for IgA deficiency.

Graft-Versus-Host Disease Graft-versus-host disease (GVHD) is a frequent complication of allogeneic stem cell transplantation, in which lymphocytes from the donor attack and cannot be eliminated by an immunodeficient host. Transfusion-related GVHD is mediated by donor T lymphocytes that recognize host HLA antigens as foreign and mount an immune response, which is manifested clinically by the development of fever, a characteristic cutaneous eruption, diarrhea, and liver function abnormalities. GVHD can also occur when blood components that contain viable T lymphocytes are transfused to immunodeficient recipients or to immunocompetent recipients who share HLA antigens with the donor (e.g., a family donor). In addition to the aforementioned clinical features of GVHD, transfusion-associated GVHD (TA-GVHD) is characterized by marrow aplasia and pancytopenia. TA-GVHD is highly resistant to treatment with immunosuppressive therapies, including glucocorticoids, cyclosporine, antithymocyte globulin, and ablative therapy followed by allogeneic bone marrow transplantation. Clinical manifestations appear at 8–10 days, and death occurs at 3–4 weeks after transfusion.

TA-GVHD can be prevented by irradiation of cellular components (minimum of 2500 cGy) before transfusion to patients at risk. Patients at risk for TA-GVHD include fetuses receiving intrauterine transfusions, selected immunocompetent (e.g., lymphoma patients) or immunocompromised recipients, recipients of donor units known to be from a blood relative, and recipients who have undergone marrow transplantation. Directed donations by family members should be discouraged (they are not less likely to transmit infection); lacking other options, the blood products from family members should always be irradiated.

Transfusion-Related Acute Lung Injury Transfusion-related acute lung injury (TRALI) is the most common cause of transfusion related fatalities. The recipient develops symptoms of hypoxia ($P_{aO_2}/F_{IO_2} < 300$ mmHg) and signs of noncardiogenic pulmonary edema, including bilateral interstitial infiltrates on chest x-ray, either during or within 6 h of transfusion. Treatment is supportive, and patients usually recover without sequelae. TRALI usually results from the transfusion of donor plasma that contains high-titer anti-HLA class II antibodies that bind recipient leukocytes. The leukocytes aggregate in the pulmonary vasculature and release mediators that increase capillary permeability. Testing the donor's plasma for anti-HLA antibodies can support this diagnosis. The implicated donors are frequently multiparous women. The transfusion of plasma from male and nulliparous women donors

reduces the risk of TRALI. Recipient factors that are associated with increased risk of TRALI include smoking, chronic alcohol use, shock, liver surgery (transplantation), mechanical ventilation with >30 cmH₂O pressure support and positive fluid balance.

Posttransfusion Purpura This reaction presents as thrombocytopenia 7–10 days after platelet transfusion and occurs predominantly in women. Platelet-specific antibodies are found in the recipient's serum, and the most frequently recognized antigen is HPA-1a found on the platelet glycoprotein IIIa receptor. The delayed thrombocytopenia is due to the production of antibodies that react to both donor and recipient platelets. Additional platelet transfusions can worsen the thrombocytopenia and should be avoided. Treatment with intravenous immunoglobulin may neutralize the effector antibodies, or plasmapheresis can be used to remove the antibodies.

Alloimmunization A recipient may become alloimmunized to a number of antigens on cellular blood elements and plasma proteins. Alloantibodies to RBC antigens are detected during pretransfusion testing, and their presence may delay finding antigen-negative cross-match-compatible products for transfusion. Women of childbearing age who are sensitized to certain RBC antigens (i.e., D, c, E, Kell, or Duffy) are at risk for bearing a fetus with hemolytic disease of the newborn. Matching for D antigen is the only pretransfusion selection test to prevent RBC alloimmunization.

Alloimmunization to antigens on leukocytes and platelets can result in refractoriness to platelet transfusions. Once alloimmunization has developed, HLA-compatible platelets from donors who share similar antigens with the recipient may be difficult to find. Hence, prudent transfusion practice is directed at preventing sensitization through the use of leukocyte-reduced cellular components, as well as limiting antigenic exposure by the judicious use of transfusions and use of SDAPs.

NONIMMUNOLOGIC REACTIONS

Fluid Overload Blood components are excellent volume expanders, and transfusion may quickly lead to transfusion-associated circulatory overload (TACO). Dyspnea with $P_{aO_2} < 90\%$ on room air, bilateral infiltrates on chest x-ray, and systolic hypertension are found with TACO. Brain natriuretic peptide (BNP) is often elevated (>1.5) compared with pretransfusion levels. Monitoring the rate and volume of the transfusion and using a diuretic can minimize this problem.

Hypothermia Refrigerated (4°C) or frozen (–18°C or below) blood components can result in hypothermia when rapidly infused. Cardiac dysrhythmias can result from exposing the sinoatrial node to cold fluid. Use of an in-line warmer will prevent this complication.

Electrolyte Toxicity RBC leakage during storage increases the concentration of potassium in the unit. Neonates and patients in renal failure are at risk for hyperkalemia. Preventive measures, such as using fresh or washed RBCs, are warranted for neonatal transfusions because this complication can be fatal.

Citrate, commonly used to anticoagulate blood components, chelates calcium and thereby inhibits the coagulation cascade. Hypocalcemia, manifested by circumoral numbness and/or tingling sensation of the fingers and toes, may result from multiple rapid transfusions. Because citrate is quickly metabolized to bicarbonate, calcium infusion is seldom required in this setting. If calcium or any other intravenous infusion is necessary, it must be given through a separate line.

Iron Overload Each unit of RBCs contains 200–250 mg of iron. Symptoms and signs of iron overload affecting endocrine, hepatic, and cardiac function are common after 100 units of RBCs have been transfused (total-body iron load of 20 g). Preventing this complication by using alternative therapies (e.g., erythropoietin) and judicious transfusion is preferable and cost effective. Chelating agents, such as deferoxamine and deferasirox, are available, but the response is often suboptimal.

Hypotensive Reactions Transient hypotension may be noted among transfused patients who take angiotensin-converting enzyme (ACE) inhibitors. Because blood products contain bradykinin that is normally

degraded by ACE, patients on ACE inhibitors may have increased bradykinin levels that cause hypotension in the recipient. The blood pressure typically returns to normal without intervention.

Immunomodulation Transfusion of allogeneic blood is immunosuppressive. Multiply transfused renal transplant recipients are less likely to reject the graft, and transfusion may result in poorer outcomes in cancer patients and increase the risk of infections. Transfusion-related immunomodulation is thought to be mediated by transfused leukocytes. Leukocyte-depleted cellular products may cause less immunosuppression, although controlled data are unlikely to be obtained as the blood supply becomes universally leukocyte depleted.

INFECTIOUS COMPLICATIONS

The blood supply is initially screened by selecting healthy donors without high-risk lifestyles, medical conditions, or exposure to transmissible pathogens, such as intravenous drug use or visiting malaria endemic areas. Multiple tests performed on donated blood to detect the presence of infectious agents using nucleic acid amplification testing (NAAT) or evidence of prior infections by testing for antibodies to pathogens and sterility of platelet products further reduce the risk of transfusion-acquired infections.

Viral Infections • HEPATITIS C VIRUS (HCV) Blood donations are tested for antibodies to HCV and HCV RNA. The risk of acquiring HCV through transfusion is now calculated to be approximately 1 in 2,000,000 units. Infection with HCV may be asymptomatic or lead to chronic active hepatitis, cirrhosis, and liver failure.

HUMAN IMMUNODEFICIENCY VIRUS TYPE 1 (HIV-1) Donated blood is tested for antibodies to HIV-1, HIV-1 p24 antigen, and HIV RNA using NAAT. Approximately a dozen seronegative donors have been shown to harbor HIV RNA. The risk of HIV-1 infection per transfusion episode is 1 in 2,000,000. Antibodies to HIV-2 are also measured in donated blood. No cases of HIV-2 infection have been reported in the United States since 1992.

HEPATITIS B VIRUS (HBV) Donated blood is screened for HBV using assays for hepatitis B surface antigen (HbsAg). NAAT testing is not practical because of slow viral replication and lower levels of viremia. The risk of transfusion-associated HBV infection is several times greater than for HCV. Vaccination of individuals who require long-term transfusion therapy can prevent this complication.

OTHER HEPATITIS VIRUSES Hepatitis A virus is rarely transmitted by transfusion; infection is typically asymptomatic and does not lead to chronic disease. Other transfusion-transmitted viruses—TT virus, SEN virus, and GB virus C—do not cause chronic hepatitis or other disease states. Routine testing does not appear to be warranted.

WEST NILE VIRUS (WNV) Transfusion-transmitted WNV infections were documented in 2002. This RNA virus can be detected using NAAT; routine screening began in 2003. WNV infections range in severity from asymptomatic to fatal, with the older population at greater risk.

CYTOMEGALOVIRUS This ubiquitous virus infects $\geq 50\%$ of the general population and is transmitted by the infected “passenger” WBCs found in transfused PRBCs or platelet components. Cellular components that are leukocyte-reduced have a decreased risk of transmitting CMV, regardless of the serologic status of the donor. Groups at risk for CMV infections include immunosuppressed patients, CMV-seronegative transplant recipients, and neonates; these patients should receive leukocyte-depleted components or CMV seronegative products.

HUMAN T LYMPHOTROPIC VIRUS (HTLV) TYPE 1 Assays to detect HTLV-1 and -2 are used to screen all donated blood. HTLV-1 is associated with adult T cell leukemia/lymphoma and tropical spastic paraparesis in a small percentage of infected persons (Chap. 225e). The risk of HTLV-1 infection via transfusion is 1 in 641,000 transfusion episodes. HTLV-2 is not clearly associated with any disease.

PARVOVIRUS B19 Blood components and pooled plasma products can transmit this virus, the etiologic agent of erythema infectiosum, or fifth disease, in children. Parvovirus B19 shows tropism for erythroid

precursors and inhibits both erythrocyte production and maturation. Pure red cell aplasia, presenting either as acute aplastic crisis or chronic anemia with shortened RBC survival, may occur in individuals with an underlying hematologic disease, such as sickle cell disease or thalassemia (Chap. 130). The fetus of a seronegative woman is at risk for developing hydrops from this virus.

Bacterial Contamination The relative risk of transfusion-transmitted bacterial infection has increased as the absolute risk of viral infections has dramatically decreased.

Most bacteria do not grow well at cold temperatures; thus, PRBCs and FFP are not common sources of bacterial contamination. However, some gram-negative bacteria can grow at 1–6°C. *Yersinia*, *Pseudomonas*, *Serratia*, *Acinetobacter*, and *Escherichia* species have all been implicated in infections related to PRBC transfusion. Platelet concentrates, which are stored at room temperature, are more likely to contain skin contaminants such as gram-positive organisms, including coagulase-negative staphylococci. It is estimated that 1 in 1000–2000 platelet components is contaminated with bacteria. The risk of death due to transfusion-associated sepsis has been calculated at 1 in 17,000 for single-unit platelets derived from whole blood donation and 1 in 61,000 for apheresis product. Since 2004, blood banks have instituted methods to detect contaminated platelet components.

Recipients of transfusion contaminated with bacteria may develop fever and chills, which can progress to septic shock and DIC. These reactions may occur abruptly, within minutes of initiating the transfusion, or after several hours. The onset of symptoms and signs is often sudden and fulminant, which distinguishes bacterial contamination from an FNHTR. The reactions, particularly those related to gram-negative contaminants, are the result of infused endotoxins formed within the contaminated stored component.

When these reactions are suspected, the transfusion must be stopped immediately. Therapy is directed at reversing any signs of shock, and broad-spectrum antibiotics should be given. The blood bank should be notified to identify any clerical or serologic error. The blood component bag should be sent for culture and Gram stain.

Other Infectious Agents Various parasites, including those causing malaria, babesiosis, and Chagas’ disease, can be transmitted by blood transfusion. Geographic migration and travel of donors shift the incidence of these rare infections. Other agents implicated in transfusion transmission include dengue, chikungunya virus, variant Creutzfeldt-Jakob disease, *A. phagocytophilum*, and yellow fever vaccine virus, and the list will grow. Tests for some pathogens are available, such as *T. cruzi*, but not universally required, whereas others are being developed (*B. microti*). These infections should be considered in the transfused patient in the appropriate clinical setting.

ALTERNATIVES TO TRANSFUSION

Alternatives to allogeneic blood transfusions that avoid homologous donor exposures with attendant immunologic and infectious risks remain attractive. Autologous blood is the best option when transfusion is anticipated. However, the cost-benefit ratio of autologous transfusion remains high. No transfusion is a zero-risk event; clerical errors and bacterial contamination remain potential complications even with autologous transfusions. Additional methods of autologous transfusion in the surgical patient include preoperative hemodilution, recovery of shed blood from sterile surgical sites, and postoperative drainage collection. Directed or designated donation from friends and family of the potential recipient has not been safer than volunteer donor component transfusions. Such directed donations may in fact place the recipient at higher risk for complications such as GVHD and alloimmunization.

Granulocyte and granulocyte-macrophage colony-stimulating factors are clinically useful to hasten leukocyte recovery in patients with leukopenia related to high-dose chemotherapy. Erythropoietin stimulates erythrocyte production in patients with anemia of chronic renal failure and other conditions, thus avoiding or reducing the need for transfusion. This hormone can also stimulate erythropoiesis in the autologous donor to enable additional donation.

139e Hematopoietic Cell Transplantation

Frederick R. Appelbaum

Bone marrow transplantation was the original term used to describe the collection and transplantation of hematopoietic stem cells, but with the demonstration that peripheral blood and umbilical cord blood are also useful sources of stem cells, *hematopoietic cell transplantation* has become the preferred generic term for this process. The procedure is usually carried out for one of two purposes: (1) to replace an abnormal but nonmalignant lymphohematopoietic system with one from a normal donor or (2) to treat malignancy by allowing the administration of higher doses of myelosuppressive therapy than would otherwise be possible. The use of hematopoietic cell transplantation has been increasing, both because of its efficacy in selected diseases and because of increasing availability of donors. The Center for International Blood and Marrow Transplant Research (<http://www.cibmtr.org>) estimates that about 65,000 transplants are performed each year.

THE HEMATOPOIETIC STEM CELL

Several features of the hematopoietic stem cell make transplantation clinically feasible, including its remarkable regenerative capacity, its ability to home to the marrow space following intravenous injection, and the ability of the stem cell to be cryopreserved (Chap. 89e). Transplantation of a single stem cell can replace the entire lymphohematopoietic system of an adult mouse. In humans, transplantation of a small percentage of a donor's bone marrow volume regularly results in complete and sustained replacement of the recipient's entire lymphohematopoietic system, including all red cells, granulocytes, B and T lymphocytes, and platelets, as well as cells comprising the fixed macrophage population, including Kupffer cells of the liver, pulmonary alveolar macrophages, osteoclasts, Langerhans cells of the skin, and brain microglial cells. The ability of the hematopoietic stem cell to home to the marrow following intravenous injection is mediated, in part, by an interaction between CXCL12, also known as stromal cell-derived factor 1, produced by marrow stromal cells and the alpha-chemokine receptor CXCR4 found on stem cells. Homing is also influenced by the interaction of cell-surface molecules, termed *selectins*, including E- and L-selectin, on bone marrow endothelial cells with ligands, termed *integrins*, such as VLA-4, on early hematopoietic cells. Human hematopoietic stem cells can survive freezing and thawing with little, if any, damage, making it possible to remove and store a portion of the patient's own bone marrow for later reinfusion following treatment of the patient with high-dose myelotoxic therapy.

CATEGORIES OF HEMATOPOIETIC CELL TRANSPLANTATION

Hematopoietic cell transplantation can be described according to the relationship between the patient and the donor and by the anatomic source of stem cells. In ~1% of cases, patients have identical twins who can serve as donors. With the use of syngeneic donors, there is no risk of graft-versus-host disease (GVHD), which often complicates allogeneic transplantation, and unlike the use of autologous marrow, there is no risk that the stem cells are contaminated with tumor cells.

Allogeneic transplantation involves a donor and a recipient who are not genetically identical. Following allogeneic transplantation, immune cells transplanted with the stem cells or developing from them can react against the patient, causing GVHD. Alternatively, if the immunosuppressive preparative regimen used to treat the patient before transplant is inadequate, immunocompetent cells of the patient can cause graft rejection. The risks of these complications are greatly influenced by the degree of matching between donor and recipient for antigens encoded by genes of the major histocompatibility complex.

The human leukocyte antigen (HLA) molecules are responsible for binding antigenic proteins and presenting them to T cells. The antigens presented by HLA molecules may derive from exogenous

sources (e.g., during active infections) or may be endogenous proteins. If individuals are not HLA-matched, T cells from one individual will react strongly to the mismatched HLA, or "major antigens," of the second. Even if the individuals are HLA-matched, the T cells of the donor may react to differing endogenous or "minor antigens" presented by the HLA of the recipient. Reactions to minor antigens tend to be less vigorous. The genes of major relevance to transplantation include HLA-A, -B, -C, and -D; they are closely linked and therefore tend to be inherited as haplotypes, with only rare crossovers between them. Thus, the odds that any one full sibling will match a patient are one in four, and the probability that the patient has an HLA-identical sibling is $1 - (0.75)^n$, where n equals the number of siblings.

With current techniques, the risk of graft rejection is 1–3%, and the risk of severe, life-threatening acute GVHD is ~15% following transplantation between HLA-identical siblings. The incidence of graft rejection and GVHD increases progressively with the use of family member donors mismatched for one, two, or three antigens. Although survival following a one-antigen mismatched transplant is not markedly altered, survival following two- or three-antigen mismatched transplants is significantly reduced. Since the formation of the National Marrow Donor Program and other registries, it has become possible to identify HLA-matched unrelated donors for many patients. The genes encoding HLA antigens are highly polymorphic, and thus the odds of any two unrelated individuals being HLA identical are extremely low, somewhat less than 1 in 10,000. However, by identifying and typing >20 million volunteer donors, HLA-matched donors can now be found for ~60% of patients for whom a search is initiated, with higher rates among whites and lower rates among minorities and patients of mixed race. It takes, on average, 3–4 months to complete a search and schedule and initiate an unrelated donor transplant. With improvements in HLA typing and supportive care measures, survival following matched unrelated donor transplantation is essentially the same as that seen with HLA-matched siblings.

Autologous transplantation involves the removal and storage of the patient's own stem cells with subsequent reinfusion after the patient receives high-dose myeloablative therapy. Unlike allogeneic transplantation, there is no risk of GVHD or graft rejection with autologous transplantation. On the other hand, autologous transplantation lacks a graft-versus-tumor (GVT) effect, and the autologous stem cell product can be contaminated with tumor cells, which could lead to relapse. A variety of techniques have been developed to "purge" autologous products of tumor cells. Some use antibodies directed at tumor-associated antigens plus complement, antibodies linked to toxins, or antibodies conjugated to immunomagnetic beads. Another technique is positive selection of stem cells using antibodies to CD34, with subsequent column adherence or flow techniques to select normal stem cells while leaving tumor cells behind. All of these approaches can reduce the number of tumor cells from 1000- to 10,000-fold and are clinically feasible; however, no prospective randomized trials have yet shown that any of these approaches results in a decrease in relapse rates or improvements in disease-free or overall survival.

Bone marrow aspirated from the posterior and anterior iliac crests initially was the source of hematopoietic stem cells for transplantation. Typically, anywhere from 1.5 to 5×10^8 nucleated marrow cells per kilogram are collected for allogeneic transplantation. Several studies have found improved survival in the settings of both matched sibling and unrelated transplantation by transplanting higher numbers of bone marrow cells.

Hematopoietic stem cells circulate in the peripheral blood but in very low concentrations. Following the administration of certain hematopoietic growth factors, including granulocyte colony-stimulating factor (G-CSF) or granulocyte-macrophage colony-stimulating factor (GM-CSF), and during recovery from intensive chemotherapy, the concentration of hematopoietic progenitor cells in blood, as measured either by colony-forming units or expression of the CD34 antigen, increases markedly. This has made it possible to harvest adequate numbers of stem cells from the peripheral blood for transplantation. Donors are typically treated with 4 or 5 days of hematopoietic growth factor, following which stem cells are collected in one or two 4-h

pheresis sessions. In the autologous setting, transplantation of $>2.5 \times 10^6$ CD34 cells per kilogram, a number that can be collected in most circumstances, leads to rapid and sustained engraftment in virtually all cases. In the 10–20% of patients who fail to mobilize sufficient CD34+ cells with growth factor alone, the addition of plerixafor, an antagonist of CXCR4, may be useful. Compared to the use of autologous marrow, use of peripheral blood stem cells results in more rapid hematopoietic recovery, with granulocytes recovering to $500/\mu\text{L}$ by day 12 and platelets recovering to $20,000/\mu\text{L}$ by day 14. Although this more rapid recovery diminishes the morbidity rate of transplantation, no studies show improved survival.

Hesitation in studying the use of peripheral blood stem cells for allogeneic transplantation occurred because peripheral blood stem cell products contain as much as 1 log more T cells than are contained in the typical marrow harvest; in animal models, the incidence of GVHD is related to the number of T cells transplanted. Nonetheless, clinical trials have shown that the use of growth factor–mobilized peripheral blood stem cells from HLA-matched family members leads to faster engraftment without an increase in acute GVHD. Chronic GVHD may be increased with peripheral blood stem cells, but in trials conducted so far, this has been more than balanced by reductions in relapse rates and nonrelapse mortality rates, with the use of peripheral blood stem cells resulting in improved overall survival. However, in the setting of matched unrelated donor transplantation, use of peripheral blood results in more chronic GVHD without a compensatory survival advantage, favoring the use of bone marrow in this setting.

Umbilical cord blood contains a high concentration of hematopoietic progenitor cells, allowing for its use as a source of stem cells for transplantation. Cord blood transplantation from family members has been explored in the setting where the immediate need for transplantation precludes waiting the 9 or so months generally required for the baby to mature to the point of donating marrow. Use of cord blood results in slower engraftment and peripheral count recovery than seen with marrow but a lower incidence of GVHD, perhaps reflecting the low number of T cells in cord blood. Multiple cord blood banks have been developed to harvest and store cord blood for possible transplantation to unrelated patients from material that would otherwise be discarded. Currently more than 500,000 units are cryopreserved and available for use. The advantages of unrelated cord blood are rapid availability and decreased immune reactivity allowing for the use of partially matched units, which is of particular importance for those without matched unrelated donors. The risks of graft failure and transplant-related mortality are related to the dose of cord blood cells per kilogram, which previously limited the application of single cord blood transplantation to pediatric and smaller adult patients. Subsequent trials have found that the use of double cord transplants diminishes the risk of graft failure and early mortality even though only one of the donors ultimately engrafts. Survival rates are now similar with unrelated donor and cord blood transplantation.

THE TRANSPLANT PREPARATIVE REGIMEN

The treatment regimen administered to patients immediately preceding transplantation is designed to eradicate the patient's underlying disease and, in the setting of allogeneic transplantation, immunosuppress the patient adequately to prevent rejection of the transplanted marrow. The appropriate regimen therefore depends on the disease setting and source of marrow. For example, when transplantation is performed to treat severe combined immunodeficiency and the donor is a histocompatible sibling, no treatment is needed because no host cells require eradication and the patient is already too immunoincompetent to reject the transplanted marrow. For aplastic anemia, there is no large population of cells to eradicate, and high-dose cyclophosphamide plus antithymocyte globulin are sufficient to immunosuppress the patient adequately to accept the marrow graft. In the setting of thalassemia and sickle cell anemia, high-dose busulfan is frequently added to cyclophosphamide in order to eradicate hyperplastic host hematopoiesis. A variety of different regimens have been developed to treat malignant diseases. Most of these regimens include agents that

have high activity against the tumor in question at conventional doses and have myelosuppression as their predominant dose-limiting toxicity. Therefore, these regimens commonly include busulfan, cyclophosphamide, melphalan, thiotepea, carmustine, etoposide, and total-body irradiation in various combinations.

Although high-dose treatment regimens have typically been used in transplantation, the understanding that much of the antitumor effect of transplantation derives from an immunologically mediated GVT response has led investigators to ask if reduced-intensity conditioning regimens might be effective and more tolerable. Evidence for a GVT effect comes from studies showing that posttransplant relapse rates are lowest in patients who develop acute and chronic GVHD, higher in those without GVHD, and higher still in recipients of T cell–depleted allogeneic or syngeneic marrow. The demonstration that complete remissions can be obtained in many patients who have relapsed after transplant by simply administering viable lymphocytes from the original donor further strengthens the argument for a potent GVT effect. Accordingly, a variety of reduced-intensity regimens have been studied, ranging from the very minimum required to achieve engraftment (e.g., fludarabine plus 200 cGy total-body irradiation) to regimens of more immediate intensity (e.g., fludarabine plus melphalan). Studies to date document that engraftment can be readily achieved with less toxicity than seen with conventional transplantation. Furthermore, the severity of acute GVHD appears to be somewhat decreased. Complete sustained responses have been documented in many patients, particularly those with more indolent hematologic malignancies. In general, relapse rates are higher following reduced-intensity conditioning, but transplant-related mortality is lower, favoring the use of reduced-intensity conditioning in older patients and those with significant comorbidities. High-dose regimens are favored in younger, fitter patients.

THE TRANSPLANT PROCEDURE

Marrow is usually collected from the donor's posterior and sometimes anterior iliac crests, with the donor under general or spinal anesthesia. Typically, 10–15 mL/kg of marrow is aspirated, placed in heparinized media, and filtered through 0.3- and 0.2-mm screens to remove fat and bony spicules. The collected marrow may undergo further processing depending on the clinical situation, such as the removal of red cells to prevent hemolysis in ABO-incompatible transplants, the removal of donor T cells to prevent GVHD, or attempts to remove possible contaminating tumor cells in autologous transplantation. Marrow donation is safe, with only very rare complications reported.

Peripheral blood stem cells are collected by leukapheresis after the donor has been treated with hematopoietic growth factors or, in the setting of autologous transplantation, sometimes after treatment with a combination of chemotherapy and growth factors. Stem cells for transplantation are infused through a large-bore central venous catheter. Such infusions are usually well tolerated, although occasionally patients develop fever, cough, or shortness of breath. These symptoms typically resolve with slowing of the infusion. When the stem cell product has been cryopreserved using dimethyl sulfoxide, patients more often experience short-lived nausea or vomiting due to the odor and taste of the cryoprotectant.

ENGRAFTMENT

Peripheral blood counts usually reach their nadir several days to a week after transplant as a consequence of the preparative regimen; then cells produced by the transplanted stem cells begin to appear in the peripheral blood. The rate of recovery depends on the source of stem cells, the use of posttransplant growth factors, and the form of GVHD prophylaxis used. If marrow is the source of stem cells, recovery to 100 granulocytes/ μL occurs on average by day 16 and to $500/\mu\text{L}$ by day 22. Use of G-CSF–mobilized peripheral blood stem cells speeds the rate of recovery by ~1 week when compared to marrow, whereas engraftment following cord blood transplantation is typically delayed by ~1 week compared to marrow. Use of a myeloid growth factor (G-CSF or GM-CSF) after transplant can accelerate recovery by 3–5 days, whereas use of methotrexate to prevent GVHD delays engraftment by a similar

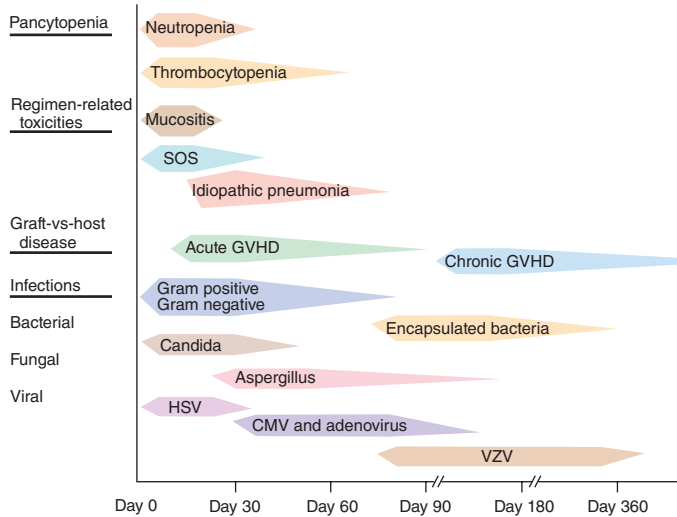


FIGURE 139e-1 Major syndromes complicating marrow transplantation. CMV, cytomegalovirus; GVHD, graft-versus-host disease; HSV, herpes simplex virus; SOS, sinusoidal obstructive syndrome (formerly venoocclusive disease); VZV, varicella-zoster virus. The size of the shaded area roughly reflects the period of risk of the complication.

period. Following allogeneic transplantation, engraftment can be documented using fluorescence in situ hybridization of sex chromosomes if donor and recipient are sex-mismatched or by analysis of a variable number of tandem repeats or short tandem repeat polymorphisms after DNA amplification.

COMPLICATIONS FOLLOWING HEMATOPOIETIC CELL TRANSPLANT

Early Direct Chemoradiotoxicities The transplant preparative regimen may cause a spectrum of acute toxicities that vary according to intensity of the regimen and the specific agents used, but frequently results in nausea, vomiting, and mild skin erythema (Fig. 139e-1). Regimens that include high-dose cyclophosphamide can result in hemorrhagic cystitis, which can usually be prevented by bladder irrigation or with the sulfhydryl compound mercaptoethanesulfonate (MESNA); rarely, acute hemorrhagic carditis is seen. Most high-dose preparative regimens will result in oral mucositis, which typically develops 5–7 days after transplant and often requires narcotic analgesia. Use of a patient-controlled analgesic pump provides the greatest patient satisfaction and results in a lower cumulative dose of narcotic. Keratinocyte growth factor (palifermin) can shorten the duration of mucositis by several days following autologous transplantation. Patients begin losing their hair 5–6 days after transplant and by 1 week are usually profoundly pancytopenic.

Depending on the intensity of the conditioning regimen, 3–10% of patients will develop sinusoidal obstruction syndrome (SOS) of the liver (formerly called venoocclusive disease), a syndrome that results from direct cytotoxic injury to hepatic-venular and sinusoidal endothelium, with subsequent deposition of fibrin and the development of a local hypercoagulable state. This chain of events leads to the clinical symptoms of tender hepatomegaly, ascites, jaundice, and fluid retention. These symptoms can develop any time during the first month after transplant, with the peak incidence at day 16. Predisposing factors include prior exposure to intensive chemotherapy, pretransplant hepatitis of any cause, and use of more intense conditioning regimens. The mortality rate of sinusoidal obstruction syndrome is ~30%, with progressive hepatic failure culminating in a terminal hepatorenal syndrome. Both thrombolytic and antithrombotic agents, such as tissue plasminogen activator, heparin, and prostaglandin E, have been studied as therapy, but none has proven of consistent major benefit in controlled trials, and all have significant toxicity. Studies with defibrotide, a polydeoxyribonucleotide, seem encouraging.

Although most pneumonias developing early after transplant are caused by infectious agents, in ~5% of patients a diffuse interstitial pneumonia will develop that is thought to be the result of direct

toxicity of high-dose preparative regimens. Bronchoalveolar lavage usually shows alveolar hemorrhage, and biopsies are typically characterized by diffuse alveolar damage, although some cases may have a more clearly interstitial pattern. High-dose glucocorticoids or antitumor necrosis factor therapies are sometimes used as treatment, although randomized trials testing their utility have not been reported.

Late Direct Chemoradiotoxicities Late complications of the preparative regimen include decreased growth velocity in children and delayed development of secondary sex characteristics. These complications can be partly ameliorated with the use of appropriate growth and sex hormone replacement. Most men become azoospermic, and most postpubertal women will develop ovarian failure, which should be treated. However, pregnancy is possible after transplantation, and patients should be counseled accordingly. Thyroid dysfunction, usually well compensated, is sometimes seen. Cataracts develop in 10–20% of patients and are most common in patients treated with total-body irradiation and those who receive glucocorticoid therapy after transplant for treatment of GVHD. Aseptic necrosis of the femoral head is seen in 10% of patients and is particularly frequent in those receiving chronic glucocorticoid therapy. Both acute and late chemoradiotoxicities (except those due to glucocorticoids and other agents used to treat GVHD) are considerably less frequent in recipients of reduced-intensity compared to high-dose preparative regimens.

Graft Failure Although complete and sustained engraftment is usually seen after transplant, occasionally marrow function either does not return or, after a brief period of engraftment, is lost. Graft failure after autologous transplantation can be the result of inadequate numbers of stem cells being transplanted, damage during ex vivo treatment or storage, or exposure of the patient to myelotoxic agents after transplant. Infections with cytomegalovirus (CMV) or human herpesvirus type 6 have also been associated with loss of marrow function. Graft failure after allogeneic transplantation can also be due to immunologic rejection of the graft by immunocompetent host cells. Immunologically based graft rejection is more common following use of less immunosuppressive preparative regimens, in recipients of T cell-depleted stem cell products, and in patients receiving grafts from HLA-mismatched donors or cord blood.

Treatment of graft failure usually involves removing all potentially myelotoxic agents from the patient's regimen and attempting a short trial of a myeloid growth factor. Persistence of lymphocytes of host origin in allogeneic transplant recipients with graft failure indicates immunologic rejection. Reinfusion of donor stem cells in such patients is usually unsuccessful unless preceded by a second immunosuppressive preparative regimen. Standard high-dose preparative regimens are generally tolerated poorly if administered within 100 days of a first transplant because of cumulative toxicities. However, use of regimens combining, for example, fludarabine plus low-dose total-body irradiation, or cyclophosphamide plus antithymocyte globulin, has been effective in some cases.

Graft-Versus-Host Disease GVHD is the result of allogeneic T cells that are transferred with the donor's stem cell inoculum reacting with antigenic targets on host cells. Acute GVHD usually occurs within the first 3 months after transplant with a peak onset around 4 weeks and is characterized by an erythematous maculopapular rash; by persistent anorexia or diarrhea, or both; and by liver disease with increased serum levels of bilirubin, alanine and aspartate aminotransferase, and alkaline phosphatase. Because many conditions can mimic acute GVHD, the diagnosis usually requires skin, liver, or endoscopic biopsy for confirmation. In all these organs, endothelial damage and lymphocytic infiltrates are seen. In skin, the epidermis and hair follicles are damaged; in liver, the small bile ducts show segmental disruption; and in intestines, destruction of the crypts and mucosal ulceration may be noted. A commonly used rating system for acute GVHD is shown in Table 139e-1. Grade I acute GVHD is of little clinical significance, does not affect the likelihood of survival, and does not require treatment. In contrast, grades II to IV GVHD are associated with significant symptoms and a poorer probability of survival, and they require aggressive therapy.

TABLE 139e-1 CLINICAL STAGING AND GRADING OF ACUTE GRAFT-VERSUS-HOST DISEASE

Clinical Stage	Skin	Liver—Bilirubin, μmol/L (mg/dL)	Gut
1	Rash <25% body surface	34–51 (2–3)	Diarrhea 500–1000 mL/d
2	Rash 25–50% body surface	51–103 (3–6)	Diarrhea 1000–1500 mL/d
3	Generalized erythroderma	103–257 (6–15)	Diarrhea >1500 mL/d
4	Desquamation and bullae	>257 (>15)	Ileus
Overall Clinical Grade	Skin Stage	Liver Stage	Gut Stage
I	1–2	0	0
II	1–3	1	1
III	1–3	2–3	2–3
IV	2–4	2–4	2–4

The incidence of acute GVHD is higher in recipients of stem cells from mismatched or unrelated donors, in older patients, and in patients unable to receive full doses of drugs used to prevent the disease.

One general approach to the prevention of GVHD is the administration of immunosuppressive drugs early after transplant. Combinations of methotrexate and either cyclosporine or tacrolimus are among the most effective and widely used regimens. Prednisone, anti-T cell antibodies, mycophenolate mofetil, sirolimus, and other immunosuppressive agents have also been or are being studied in various combinations. A second general approach to GVHD prevention is removal of T cells from the stem cell inoculum. While effective in preventing GVHD, T cell depletion has been associated with an increased incidence of graft failure, infectious complications, and tumor recurrence after transplant; as yet, it is unsettled whether T cell depletion improves cure rates in any specific setting.

Despite prophylaxis, significant acute GVHD will develop in ~30% of recipients of stem cells from matched siblings and in as many as 60% of those receiving stem cells from unrelated donors. The disease is usually treated with glucocorticoids, additional immunosuppressants, or monoclonal antibodies targeted against T cells or T cell subsets.

Chronic GVHD occurs most commonly between 3 months and 2 years after allogeneic transplant, developing in 20–50% of recipients. The disease is more common in older patients, in recipients of mismatched or unrelated stem cells, and in those with a preceding episode of acute GVHD. The disease resembles an autoimmune disorder with malar rash, sicca syndrome, arthritis, obliterative bronchiolitis, and bile duct degeneration and cholestasis. Single-agent prednisone or cyclosporine is standard treatment at present, although trials of other agents are under way. Mortality rates from chronic GVHD average around 15%, but range from 5–50% depending on severity. In most patients, chronic GVHD resolves, but it may require 1–3 years of immunosuppressive treatment before these agents can be withdrawn without the disease recurring. Because patients with chronic GVHD are susceptible to significant infection, they should receive prophylactic trimethoprim-sulfamethoxazole, and all suspected infections should be investigated and treated aggressively.

Although onset before or after 3 months after transplant is often used to discriminate between acute and chronic GVHD, occasional patients will develop signs and symptoms of acute GVHD after 3 months (late-onset acute GVHD), whereas others will exhibit signs and symptoms of both acute and chronic GVHD (overlap syndrome). There are as yet no data to suggest that these patients should be treated differently than those with classic acute or chronic GVHD.

From 3–5% of patients will develop an autoimmune disorder following allogeneic HCT, most commonly autoimmune hemolytic anemia or idiopathic thrombocytopenic purpura. Unrelated donor source and chronic GVHD are risk factors, but autoimmune disorders have been reported in patients with no obvious GVHD. Treatment is with prednisone, cyclosporine, or rituximab.

Infection Posttransplant patients, particularly recipients of allogeneic transplantation, require unique approaches to the problem of

infection. Early after transplantation, patients are profoundly neutropenic, and because the risk of bacterial infection is so great, most centers initiate antibiotic treatment once the granulocyte count falls to <500/μL. Fluconazole prophylaxis at a dose of 200–400 mg/d reduces the risk of candidal infections. Patients seropositive for herpes simplex should receive acyclovir prophylaxis. One approach to infection prophylaxis is shown in [Table 139e-2](#). Despite these prophylactic measures, most patients will develop fever and signs of infection after transplant. The management of patients who become febrile despite bacterial and fungal prophylaxis is a difficult challenge and is guided by individual aspects of the patient and by the institution's experience.

The general problem of infection in the immunocompromised host is discussed in Chap. 169.

Once patients engraft, the incidence of bacterial infection diminishes; however, patients, particularly allogeneic transplant recipients, remain at significant risk of infection. During the period from engraftment until about 3 months after transplant, the most common causes of infection are gram-positive bacteria, fungi (particularly *Aspergillus*), and viruses including CMV. CMV infection, which in the past was frequently seen and often fatal, can be prevented in seronegative patients transplanted from seronegative donors by the use of either seronegative blood products or products from which the white blood cells have been removed. In seropositive patients or patients transplanted from seropositive donors, the use of ganciclovir, either as prophylaxis beginning at the time of engraftment or initiated when CMV first reactivates as evidenced by development of antigenemia or viremia, can significantly reduce the risk of CMV disease. Foscarnet is effective for some patients who develop CMV antigenemia or infection despite the use of ganciclovir or who cannot tolerate the drug.

Pneumocystis jiroveci pneumonia, once seen in 5–10% of patients, can be prevented by treating patients with oral trimethoprim-sulfamethoxazole for 1 week before transplant and resuming the treatment once patients have engrafted.

The risk of infection diminishes considerably beyond 3 months after transplant unless chronic GVHD develops, requiring continuous immunosuppression. Most transplant centers recommend continuing trimethoprim-sulfamethoxazole prophylaxis while patients are receiving any immunosuppressive drugs and also recommend careful monitoring for late CMV reactivation. In addition, many centers recommend prophylaxis against varicella-zoster, using acyclovir for 1 year after transplant. Patients should be revaccinated against tetanus, diphtheria, *Haemophilus influenzae*, polio, and pneumococcal pneumonia starting at 12 months after transplant and against measles, mumps, and rubella (MMR), varicella-zoster virus, and possibly pertussis at 24 months.

TABLE 139e-2 APPROACH TO INFECTION PROPHYLAXIS IN ALLOGENEIC TRANSPLANT RECIPIENTS

Organism	Agent	Approach
Bacterial	Levofloxacin	750 mg PO or IV daily
Fungal	Fluconazole	400 mg PO qd to day 75 posttransplant
<i>Pneumocystis carinii</i>	Trimethoprim-sulfamethoxazole	1 double-strength tablet PO bid 2 days/week until day 180 or off immunosuppression
Viral		
Herpes simplex	Acyclovir	800 mg PO bid to day 30
Varicella-zoster	Acyclovir	800 mg PO bid to day 365
Cytomegalovirus	Ganciclovir	5 mg/kg IV bid for 7 days, then 5 (mg/kg)/d 5 days/week to day 100

TREATMENT OF SPECIFIC DISEASES USING HEMATOPOIETIC CELL TRANSPLANTATION

TREATMENT NONMALIGNANT DISEASES

IMMUNODEFICIENCY DISORDERS

By replacing abnormal stem cells with cells from a normal donor, hematopoietic cell transplantation can cure patients of a variety of immunodeficiency disorders including severe combined immunodeficiency, Wiskott-Aldrich syndrome, and Chédiak-Higashi syndrome. The widest experience has been with severe combined immunodeficiency disease, where cure rates of 90% can be expected with HLA-identical donors and success rates of 50–70% have been reported using haplotype-mismatched parents as donors (Table 139e-3).

APLASTIC ANEMIA

Transplantation from matched siblings after a preparative regimen of high-dose cyclophosphamide and antithymocyte globulin can cure up to 90% of patients age <40 years with severe aplastic anemia. Results in older patients and in recipients of mismatched family member or unrelated marrow are less favorable; therefore, a trial of immunosuppressive therapy is generally recommended for such patients before considering transplantation. Transplantation is effective in all forms of aplastic anemia including, for example, the syndromes associated with paroxysmal nocturnal hemoglobinuria and Fanconi's anemia. Patients with Fanconi's anemia are abnormally sensitive to the toxic effects of alkylating agents, and so less intensive preparative regimens must be used in their treatment (Chap. 130).

HEMOGLOBINOPATHIES

Marrow transplantation from an HLA-identical sibling following a preparative regimen of busulfan and cyclophosphamide can cure 80–90% of patients with thalassemia major. The best outcomes can

be expected if patients are transplanted before they develop hepatomegaly or portal fibrosis and if they have been given adequate iron chelation therapy. Among such patients, the probabilities of 5-year survival and disease-free survival are 95 and 90%, respectively. Although prolonged survival can be achieved with aggressive chelation therapy, transplantation is the only curative treatment for thalassemia. Transplantation is being studied as a curative approach to patients with sickle cell anemia. Two-year survival and disease-free survival rates of 90 and 80%, respectively, have been reported following matched sibling or cord blood transplantation. Decisions about patient selection and the timing of transplantation remain difficult, but transplantation represents a reasonable option for younger patients who suffer repeated crises or other significant complications and who have not responded to other interventions (Chap. 127).

OTHER NONMALIGNANT DISEASES

Theoretically, hematopoietic cell transplantation should be able to cure any disease that results from an inborn error of the lymphohematopoietic system. Transplantation has been used successfully to treat congenital disorders of white blood cells such as Kostmann's syndrome, chronic granulomatous disease, and leukocyte adhesion deficiency. Congenital anemias such as Blackfan-Diamond anemia can also be cured with transplantation. Infantile malignant osteopetrosis is due to an inability of the osteoclast to resorb bone, and because osteoclasts derive from the marrow, transplantation can cure this rare inherited disorder.

Hematopoietic cell transplantation has been used as treatment for a number of storage diseases caused by enzymatic deficiencies, such as Gaucher's disease, Hurler's syndrome, Hunter's syndrome, and infantile metachromatic leukodystrophy. Transplantation for these diseases has not been uniformly successful, but treatment early in the course of these diseases, before irreversible damage to extramedullary organs has occurred, increases the chance for success.

Transplantation is being explored as a treatment for severe acquired autoimmune disorders. These trials are based on studies demonstrating that transplantation can reverse autoimmune disorders in animal models and on the observation that occasional patients with coexisting autoimmune disorders and hematologic malignancies have been cured of both with transplantation.

TABLE 139e-3 ESTIMATED 5-YEAR SURVIVAL RATES FOLLOWING TRANSPLANTATION^a

Disease	Allogeneic, %	Autologous, %
Severe combined immunodeficiency	90	N/A
Aplastic anemia	90	N/A
Thalassemia	90	N/A
Acute myeloid leukemia		
First remission	55–60	50
Second remission	40	30
Acute lymphocytic leukemia		
First remission	50	40
Second remission	40	30
Chronic myeloid leukemia		
Chronic phase	70	ID
Accelerated phase	40	ID
Blast crisis	15	ID
Chronic lymphocytic leukemia	50	ID
Myelodysplasia	45	ID
Multiple myeloma	30	35
Non-Hodgkin's lymphoma		
First relapse/second remission	40	40
Hodgkin's disease		
First relapse/second remission	40	50

^aThese estimates are generally based on data reported by the International Bone Marrow Transplant Registry. The analysis has not been reviewed by their Advisory Committee.

Abbreviations: ID, insufficient data; N/A, not applicable.

TREATMENT MALIGNANT DISEASES

ACUTE LEUKEMIA

Allogeneic hematopoietic cell transplantation cures 15–20% of patients who do not achieve complete response from induction chemotherapy for acute myeloid leukemia (AML) and is the only form of therapy that can cure such patients. Cure rates of 30–35% are seen when patients are transplanted in second remission or in first relapse. The best results with allogeneic transplantation are achieved when applied during first remission, with disease-free survival rates averaging 55–60%. Meta-analyses of studies comparing matched related donor transplantation to chemotherapy for adult AML patients age <60 years show a survival advantage with transplantation. This advantage is greatest for those with unfavorable-risk AML and is lost in those with favorable-risk disease. The role of autologous transplantation in the treatment of AML is less well defined. The rates of disease recurrence with autologous transplantation are higher than those seen after allogeneic transplantation, and cure rates are somewhat less.

Similar to patients with AML, adults with acute lymphocytic leukemia who do not achieve a complete response to induction chemotherapy can be cured in 15–20% of cases with immediate transplantation. Cure rates improve to 30–50% in second remission, and therefore transplantation can be recommended for adults who

have persistent disease after induction chemotherapy or who have subsequently relapsed. Transplantation in first remission results in cure rates about 55%. Transplantation appears to offer a clear advantage over chemotherapy for patients with high-risk disease, such as those with Philadelphia chromosome–positive disease. Debate continues about whether adults with standard-risk disease should be transplanted in first remission or whether transplantation should be reserved until relapse. Autologous transplantation is associated with a higher relapse rate but a somewhat lower risk of nonrelapse mortality when compared to allogeneic transplantation. There is no obvious role of autologous transplantation for acute lymphocytic leukemia in first remission, and for second-remission patients, most experts recommend use of allogeneic stem cells if an appropriate donor is available.

CHRONIC LEUKEMIA

Allogeneic hematopoietic cell transplantation is the only therapy shown to cure a substantial portion of patients with chronic myeloid leukemia (CML). Five-year disease-free survival rates are 15–20% for patients transplanted for blast crisis, 25–50% for accelerated-phase patients, and 60–70% for chronic-phase patients, with cure rates as high as 80% at selected centers. However, with the availability of imatinib mesylate and other highly active tyrosine kinase inhibitors (TKIs), transplantation is generally reserved for those who fail to achieve a complete cytogenetic response with a TKI, relapse after an initial response, or are intolerant of the drugs (**Chap. 133**).

Allogeneic transplantation using a high-dose preparative regimen has rarely been used for chronic lymphocytic leukemia (CLL), in large part because of the chronic nature of the disease and because of the age profile of patients. In those cases where it was studied, complete remissions were achieved in the majority of patients, with disease-free survival rates of ~50% at 3 years, despite the advanced stage of the disease at the time of transplant. The marked antitumor effects have resulted in the increased use and study of allogeneic transplantation using reduced-intensity conditioning for the treatment of CLL.

MYELOYDYSPLASIA

Between 20 and 65% of patients with myelodysplasia appear to be cured with allogeneic transplantation. Results are better among younger patients and those with less advanced disease. However, patients with early-stage myelodysplasia can live for extended periods without intervention, and so transplantation is generally reserved for patients with an International Prognostic Scoring System (IPSS) score of Int-2 or for selected patients with an IPSS score of Int-1 who have other poor prognostic features (**Chap. 130**).

LYMPHOMA

Patients with disseminated intermediate- or high-grade non-Hodgkin's lymphoma who have not been cured by first-line chemotherapy and are transplanted in first relapse or second remission can still be cured in 40–50% of cases. This represents a clear advantage over results obtained with conventional-dose salvage chemotherapy. It is unsettled whether patients with high-risk disease benefit from transplantation in first remission. Most experts favor the use of autologous rather than allogeneic transplantation for patients with intermediate- or high-grade non-Hodgkin's lymphoma, because fewer complications occur with this approach and survival appears equivalent. For patients with recurrent disseminated indolent non-Hodgkin's lymphoma, autologous transplantation results in high response rates and improved progression-free survival compared to salvage chemotherapy. However, late relapses are seen after transplantation. The role of autologous transplantation in the initial

treatment of patients is debated. It may be indicated in the small subset of patients presenting with high-risk prognostic factors but is not clearly more effective in those in lower risk groups. Reduced-intensity conditioning regimens followed by allogeneic transplantation result in high response rates in patients with indolent lymphomas, but the exact role of this approach remains to be defined.

The role of transplantation in Hodgkin's disease is similar to that in intermediate- and high-grade non-Hodgkin's lymphoma. With transplantation, 5-year disease-free survival is 20–30% in patients who never achieve a first remission with standard chemotherapy and up to 70% for those transplanted in second remission. Transplantation has no defined role in first remission in Hodgkin's disease.

MYELOMA

Patients with myeloma who have progressed on first-line therapy can sometimes benefit from allogeneic or autologous transplantation. Prospective randomized studies demonstrate that the inclusion of autologous transplantation as part of the initial therapy of patients results in improved disease-free survival and overall survival. Further benefit is seen with the use of lenalidomide maintenance therapy following transplantation. The use of autologous transplantation followed by nonmyeloablative allogeneic transplantation has yielded mixed results.

SOLID TUMORS

Randomized trials evaluating autologous transplantation as treatment for primary or metastatic breast cancer have failed to show a consistent survival advantage with this approach, and therefore, there is no established role for transplantation in this disease.

Patients with testicular cancer in whom first-line platinum-containing chemotherapy has failed can still be cured in ~50% of cases if treated with high-dose chemotherapy with autologous stem cell support, an outcome better than that seen with low-dose salvage chemotherapy.

The use of high-dose chemotherapy with autologous stem cell support is being studied for several other solid tumors, including neuroblastoma and pediatric sarcomas. As in most other settings, the best results have been obtained in patients with limited amounts of disease and where the remaining tumor remains sensitive to conventional-dose chemotherapy. Few randomized trials of transplantation in these diseases have been completed.

Partial and complete responses have been reported following nonmyeloablative allogeneic transplantation for some solid tumors, most notably renal cell cancers. The GVT effect, well documented in the treatment of hematologic malignancies, may apply to selected solid tumors under certain circumstances.

POSTTRANSPLANT RELAPSE

Patients who relapse following autologous transplantation sometimes respond to further chemotherapy and may be candidates for possible allogeneic transplantation, particularly if the remission following the initial autologous transplant was long. Several options are available for patients who relapse following allogeneic transplantation. Of particular interest are the response rates seen with infusion of unirradiated donor lymphocytes. Complete responses in as many as 75% of patients with chronic myeloid leukemia, 40% in myelodysplasia, 25% in AML, and 15% in myeloma have been reported. Major complications of donor lymphocyte infusions include transient myelosuppression and the development of GVHD. These complications depend on the number of donor lymphocytes given and the schedule of infusions, with less GVHD seen with lower dose, fractionated schedules.

SECTION 3 DISORDERS OF HEMOSTASIS

140 Disorders of Platelets and Vessel Wall

Barbara A. Konkle

Hemostasis is a dynamic process in which the platelet and the blood vessel wall play key roles. Platelets become activated upon adhesion to von Willebrand factor (VWF) and collagen in the exposed subendothelium after injury. Platelet activation is also mediated through shear forces imposed by blood flow itself, particularly in areas where the vessel wall is diseased, and is also affected by the inflammatory state of the endothelium. The activated platelet surface provides the major physiologic site for coagulation factor activation, which results in further platelet activation and fibrin formation. Genetic and acquired influences on the platelet and vessel wall, as well as on the coagulation and fibrinolytic systems, determine whether normal hemostasis or bleeding or clotting symptoms will result.

THE PLATELET

Platelets are released from the megakaryocyte, likely under the influence of flow in the capillary sinuses. The normal blood platelet count is 150,000–450,000/ μL . The major regulator of platelet production is the hormone thrombopoietin (TPO), which is synthesized in the liver. Synthesis is increased with inflammation and specifically by interleukin 6. TPO binds to its receptor on platelets and megakaryocytes, by which it is removed from the circulation. Thus a reduction in platelet and megakaryocyte mass increases the level of TPO, which then stimulates platelet production. Platelets circulate with an average life span of 7–10 days. Approximately one-third of the platelets reside in the spleen, and this number increases in proportion to splenic size, although the platelet count rarely decreases to $<40,000/\mu\text{L}$ as the spleen enlarges. Platelets are physiologically very active, but are anucleate, and thus have limited capacity to synthesize new proteins.

Normal vascular endothelium contributes to preventing thrombosis by inhibiting platelet function (Chap. 78). When vascular endothelium is injured, these inhibitory effects are overcome, and platelets adhere to the exposed intimal surface primarily through VWF, a large multimeric protein present in both plasma and in the extracellular matrix of the subendothelial vessel wall. Platelet adhesion results in the generation of intracellular signals that lead to activation of the platelet glycoprotein (Gp) IIb/IIIa ($\alpha_{\text{IIb}}\beta_3$) receptor and resultant platelet aggregation.

Activated platelets undergo release of their granule contents, which include nucleotides, adhesive proteins, growth factors, and

procoagulants that serve to promote platelet aggregation and blood clot formation and influence the environment of the forming clot. During platelet aggregation, additional platelets are recruited to the site of injury, leading to the formation of an occlusive platelet thrombus. The platelet plug is stabilized by the fibrin mesh that develops simultaneously as the product of the coagulation cascade.

THE VESSEL WALL

Endothelial cells line the surface of the entire circulatory tree, totaling $1\text{--}6 \times 10^{13}$ cells, enough to cover a surface area equivalent to about six tennis courts. The endothelium is physiologically active, controlling vascular permeability, flow of biologically active molecules and nutrients, blood cell interactions with the vessel wall, the inflammatory response, and angiogenesis.

The endothelium normally presents an antithrombotic surface (Chap. 78) but rapidly becomes prothrombotic when stimulated, which promotes coagulation, inhibits fibrinolysis, and activates platelets. In many cases, endothelium-derived vasodilators are also platelet inhibitors (e.g., nitric oxide) and, conversely, endothelium-derived vasoconstrictors (e.g., endothelin) can also be platelet activators. The net effect of vasodilation and inhibition of platelet function is to promote blood fluidity, whereas the net effect of vasoconstriction and platelet activation is to promote thrombosis. Thus, blood fluidity and hemostasis are regulated by the balance of antithrombotic/prothrombotic and vasodilatory/vasoconstrictor properties of endothelial cells.

DISORDERS OF PLATELETS

THROMBOCYTOPENIA

Thrombocytopenia results from one or more of three processes: (1) decreased bone marrow production; (2) sequestration, usually in an enlarged spleen; and/or (3) increased platelet destruction. Disorders of production may be either inherited or acquired. In evaluating a patient with thrombocytopenia, a key step is to review the peripheral blood smear and to first rule out “pseudothrombocytopenia,” particularly in a patient without an apparent cause for the thrombocytopenia. Pseudothrombocytopenia (Fig. 140-1B) is an in vitro artifact resulting from platelet agglutination via antibodies (usually IgG, but also IgM and IgA) when the calcium content is decreased by blood collection in ethylenediamine tetraacetic (EDTA) (the anticoagulant present in tubes [purple top] used to collect blood for complete blood counts [CBCs]). If a low platelet count is obtained in EDTA-anticoagulated blood, a blood smear should be evaluated and a platelet count determined in blood collected into sodium citrate (blue top tube) or heparin (green top tube), or a smear of freshly obtained unanticoagulated blood, such as from a finger stick, can be examined.

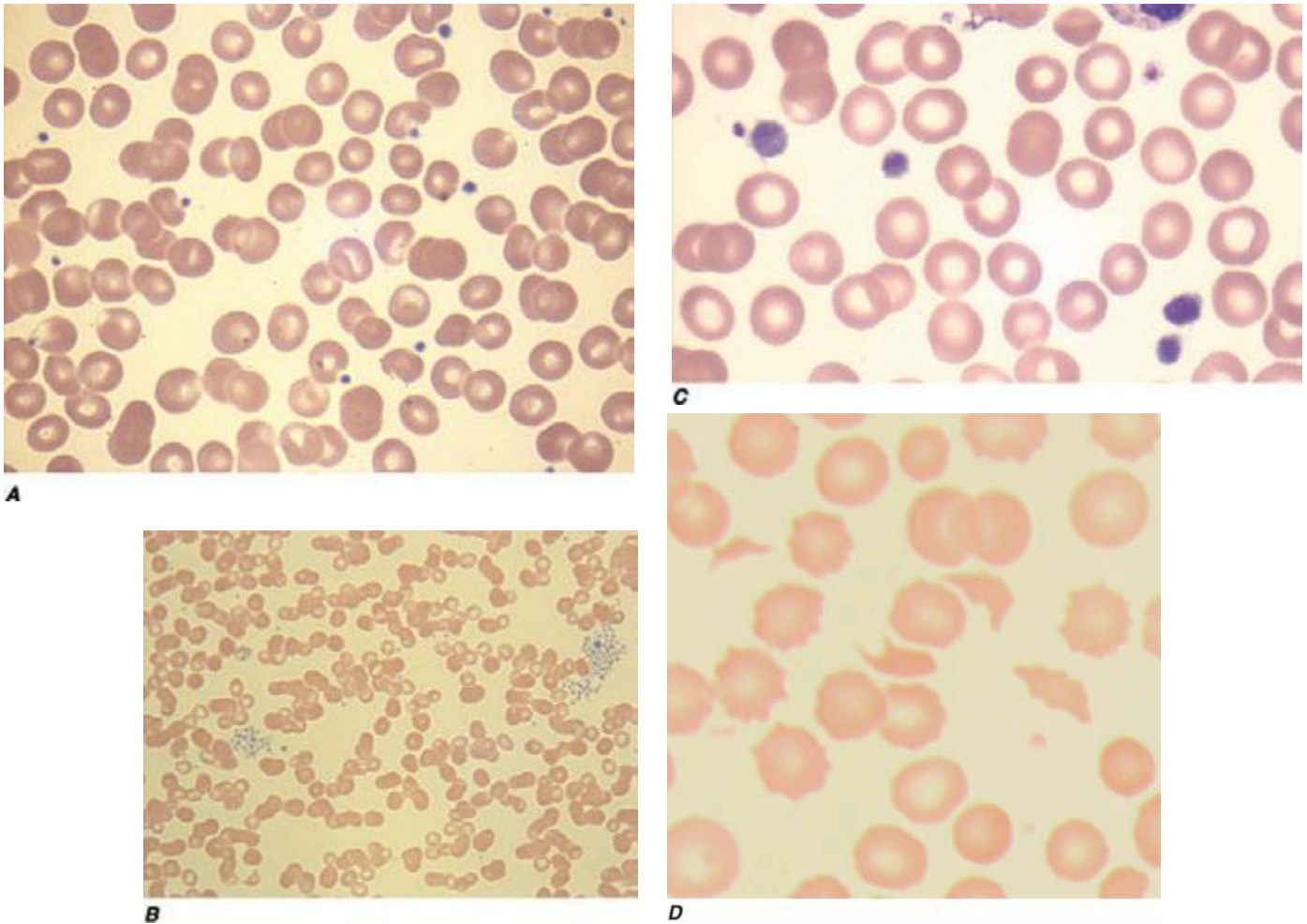


FIGURE 140-1 Photomicrographs of peripheral blood smears. **A.** Normal peripheral blood. **B.** Platelet clumping in pseudothrombocytopenia. **C.** Abnormal large platelet in autosomal dominant macrothrombocytopenia. **D.** Schistocytes and decreased platelets in microangiopathic hemolytic anemia.

APPROACH TO THE PATIENT: Thrombocytopenia

The history and physical examination, results of the CBC, and review of the peripheral blood smear are all critical components in the initial evaluation of thrombocytopenic patients (Fig. 140-2). The overall health of the patient and whether he or she is receiving drug treatment will influence the differential diagnosis. A healthy young adult with thrombocytopenia will have a much more limited differential diagnosis than an ill hospitalized patient who is receiving multiple medications. Except in unusual inherited disorders, decreased platelet production usually results from bone marrow disorders that also affect red blood cell (RBC) and/or white blood cell (WBC) production. Because myelodysplasia can present with isolated thrombocytopenia, the bone marrow should be examined in patients presenting with isolated thrombocytopenia who are older than 60 years of age. While inherited thrombocytopenia is rare, any prior platelet counts should be retrieved and a family history regarding thrombocytopenia obtained. A careful history of drug ingestion should be obtained, including nonprescription and herbal remedies, because drugs are the most common cause of thrombocytopenia.

The physical examination can document an enlarged spleen, evidence of chronic liver disease, and other underlying disorders. Mild to moderate splenomegaly may be difficult to appreciate in many individuals due to body habitus and/or obesity but can be easily assessed by abdominal ultrasound. A platelet count of approximately 5000–10,000 is required to maintain vascular integrity in the microcirculation. When the count is markedly decreased, petechiae

ALGORITHM FOR THROMBOCYTOPENIA EVALUATION

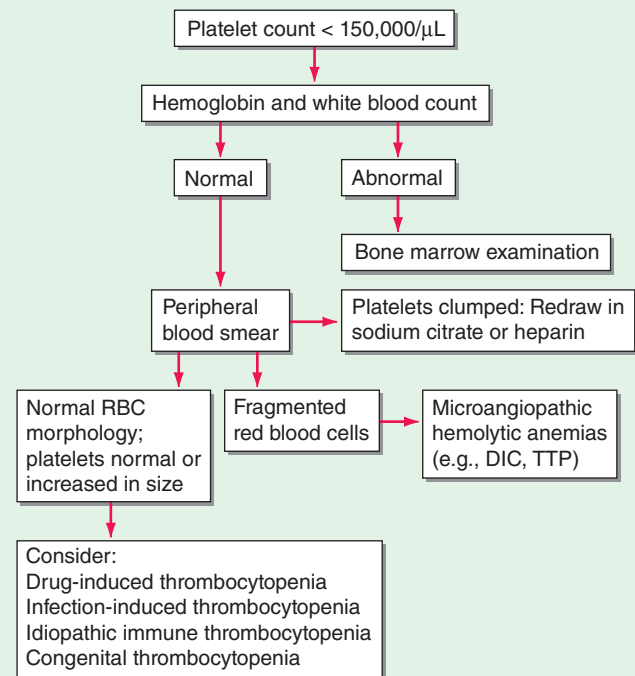


FIGURE 140-2 Algorithm for evaluating the thrombocytopenic patient. DIC, disseminated intravascular coagulation; RBC, red blood cell; TTP, thrombotic thrombocytopenic purpura.

first appear in areas of increased venous pressure, the ankles and feet in an ambulatory patient. Petechiae are pinpoint, nonblanching hemorrhages and are usually a sign of a decreased platelet number and not platelet dysfunction. Wet purpura, blood blisters that form on the oral mucosa, are thought to denote an increased risk of life-threatening hemorrhage in the thrombocytopenic patient. Excessive bruising is seen in disorders of both platelet number and function.

Infection-Induced Thrombocytopenia Many viral and bacterial infections result in thrombocytopenia and are the most common noniatrogenic cause of thrombocytopenia. This may or may not be associated with laboratory evidence of disseminated intravascular coagulation (DIC), which is most commonly seen in patients with systemic infections with gram-negative bacteria. Infections can affect both platelet production and platelet survival. In addition, immune mechanisms can be at work, as in infectious mononucleosis and early HIV infection. Late in HIV infection, pancytopenia and decreased and dysplastic platelet production are more common. Immune-mediated thrombocytopenia in children usually follows a viral infection and almost always resolves spontaneously. This association of infection with immune thrombocytopenic purpura is less clear in adults.

Bone marrow examination is often requested for evaluation of occult infections. A study evaluating the role of bone marrow examination in fever of unknown origin in HIV-infected patients found that for 86% of patients, the same diagnosis was established by less invasive techniques, notably blood culture. In some instances, however, the diagnosis can be made earlier; thus, a bone marrow examination and culture are recommended when the diagnosis is needed urgently or when other, less invasive methods have been unsuccessful.

Drug-Induced Thrombocytopenia Many drugs have been associated with thrombocytopenia. A predictable decrease in platelet count occurs after treatment with many chemotherapeutic drugs due to bone marrow suppression (Chap. 103e). Drugs that cause isolated thrombocytopenia and have been confirmed with positive laboratory testing are listed in Table 140-1, but all drugs should be suspect in a patient with thrombocytopenia without an apparent cause and should be stopped, or substituted, if possible. A helpful website, Platelets on the Internet (<http://www.ouhsc.edu/platelets/ditp.html>), lists drugs and supplements reported to have caused thrombocytopenia and the level of evidence supporting the association. Although not as well studied, herbal and over-the-counter preparations may also result in thrombocytopenia and should be discontinued in patients who are thrombocytopenic.

TABLE 140-1 DRUGS REPORTED AS DEFINITELY OR PROBABLY CAUSING ISOLATED THROMBOCYTOPENIA^a

Abciximab	Mirtazapine
Acetaminophen	Naproxen
Amiodarone	Oxaliplatin
Amlodipine	Penicillin
Ampicillin	Phenytoin
Carbamazepine	Piperacillin
Ceftriaxone	Quinine
Cephmandole	Quinidine
Ciprofloxacin	Ranitidine
Diazepam	Rosiglitazone
Eptifibatid	Roxifiban
Furosemide	Sulfisoxazole
Gold	Suramin
Haloperidol	Tirofiban
Heparin	Tranilast
Ibuprofen	Trimethoprim/sulfamethoxazole
Lorazepam	Vancomycin

^aBased on scoring requiring a compatible clinical picture and positive laboratory testing.

Source: Adapted from DM Arnold et al: J Thromb Hemost 11:169, 2013.

Classic drug-dependent antibodies are antibodies that react with specific platelet surface antigens and result in thrombocytopenia only when the drug is present. Many drugs are capable of inducing these antibodies, but for some reason, they are more common with quinine and sulfonamides. Drug-dependent antibody binding can be demonstrated by laboratory assays, showing antibody binding in the presence of, but not without, the drug present in the assay. The thrombocytopenia typically occurs after a period of initial exposure (median length 21 days), or upon reexposure, and usually resolves in 7–10 days after drug withdrawal. The thrombocytopenia caused by the platelet Gp IIb/IIIa inhibitory drugs, such as abciximab, differs in that it may occur within 24 h of initial exposure. This appears to be due to the presence of naturally occurring antibodies that cross-react with the drug bound to the platelet.

Heparin-Induced Thrombocytopenia Drug-induced thrombocytopenia due to heparin differs from that seen with other drugs in two major ways. (1) The thrombocytopenia is not usually severe, with nadir counts rarely <20,000/μL. (2) Heparin-induced thrombocytopenia (HIT) is not associated with bleeding and, in fact, markedly increases the risk of thrombosis. HIT results from antibody formation to a complex of the platelet-specific protein platelet factor 4 (PF4) and heparin. The anti-heparin/PF4 antibody can activate platelets through the FcγRIIIa receptor and also activate monocytes and endothelial cells. Many patients exposed to heparin develop antibodies to heparin/PF4, but do not appear to have adverse consequences. A fraction of those who develop antibodies will develop HIT, and a portion of those (up to 50%) will develop thrombosis (HITT).

HIT can occur after exposure to low-molecular-weight heparin (LMWH) as well as unfractionated heparin (UFH), although it is more common with the latter. Most patients develop HIT after exposure to heparin for 5–14 days (Fig. 140-3). It occurs before 5 days in those who were exposed to heparin in the prior few weeks or months (<~100 days) and have circulating anti-heparin/PF4 antibodies. Rarely, thrombocytopenia and thrombosis begin several days after all heparin has been stopped (termed *delayed-onset HIT*). The “4T’s” have been recommended to be used in a diagnostic algorithm for HIT: *thrombocytopenia*, *timing* of platelet count drop, *thrombosis* and other sequelae such as localized skin reactions, and *other* causes of thrombocytopenia not evident. Application of the 4T scoring system is very useful in excluding a diagnosis of HIT but will result in overdiagnosis of HIT in situations where thrombocytopenia and thrombosis due to other etiologies are common, such as in the intensive care unit. A scoring model based on broad expert opinion (the HIT Expert Probability [HEP] Score) has improved operating characteristics and may provide better utility as a scoring system.

LABORATORY TESTING FOR HIT HIT (anti-heparin/PF4) antibodies can be detected using two types of assays. The most widely available is an enzyme-linked immunoassay (ELISA) with PF4/polyanion complex as the antigen. Because many patients develop antibodies but do not develop clinical HIT, the test has a low specificity for the diagnosis

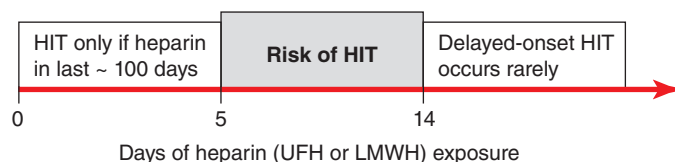


FIGURE 140-3 Time course of heparin-induced thrombocytopenia (HIT) development after heparin exposure. The timing of development after heparin exposure is a critical factor in determining the likelihood of HIT in a patient. HIT occurs early after heparin exposure in the presence of preexisting heparin/platelet factor 4 (PF4) antibodies, which disappear from circulation by ~100 days following a prior exposure. Rarely, HIT may occur later after heparin exposure (termed delayed-onset HIT). In this setting, heparin/PF4 antibody testing is usually markedly positive. HIT can occur after exposure to either unfractionated (UFH) or low-molecular-weight heparin (LMWH).

728 of HIT. This is especially true in patients who have undergone cardiopulmonary bypass surgery, where approximately 50% of patients develop these antibodies postoperatively. IgG-specific ELISAs increase specificity but may decrease sensitivity. The other assay is a platelet activation assay, most commonly the serotonin release assay, which measures the ability of the patient's serum to activate platelets in the presence of heparin in a concentration-dependent manner. This test has lower sensitivity but higher specificity than the ELISA. However, HIT remains a clinical diagnosis.

TREATMENT HEPARIN-INDUCED THROMBOCYTOPENIA

Early recognition is key in treatment of HIT, with prompt discontinuation of heparin and use of alternative anticoagulants if bleeding risk does not outweigh thrombotic risk. Thrombosis is a common complication of HIT, even after heparin discontinuation, and can occur in both the venous and arterial systems. Patients with higher anti-heparin/PF4 antibody titers may have a higher risk of thrombosis. In patients diagnosed with HIT, imaging studies to evaluate the patient for thrombosis (at least lower extremity duplex Doppler imaging) are recommended. Patients requiring anticoagulation should be switched from heparin to an alternative anticoagulant. The direct thrombin inhibitors (DTIs) argatroban and lepirudin are effective in HIT. The DTI bivalirudin and the antithrombin-binding pentasaccharide fondaparinux are also effective but not yet approved by the U.S. Food and Drug Administration (FDA) for this indication. Danaparoid, a mixture of glycosaminoglycans with anti-Xa activity, has been used extensively for the treatment of HIT; it is no longer available in the United States but is in other countries. HIT antibodies cross-react with LMWH, and these preparations should not be used in the treatment of HIT.

Because of the high rate of thrombosis in patients with HIT, anticoagulation should be considered, even in the absence of thrombosis. In patients with thrombosis, patients can be transitioned to warfarin, with treatment usually for 3–6 months. In patients without thrombosis, the duration of anticoagulation needed is undefined. An increased risk of thrombosis is present for at least 1 month after diagnosis; however, most thromboses occur early, and whether thrombosis occurs later if the patient is initially anticoagulated is unknown. Options include continuing anticoagulation until a few days after platelet recovery or for 1 month. Introduction of warfarin alone in the setting of HIT or HITT may precipitate thrombosis, particularly venous gangrene, presumably due to clotting activation and severely reduced levels of proteins C and S. Warfarin therapy, if started, should be overlapped with a DTI or fondaparinux and started after resolution of the thrombocytopenia and lessening of the prothrombotic state.

Immune Thrombocytopenic Purpura Immune thrombocytopenic purpura (ITP; also termed *idiopathic thrombocytopenic purpura*) is an acquired disorder in which there is immune-mediated destruction of platelets and possibly inhibition of platelet release from the megakaryocyte. In children, it is usually an acute disease, most commonly following an infection, and with a self-limited course. In adults, it is a more chronic disease, although in some adults, spontaneous remission occurs, usually within months of diagnosis. ITP is termed *secondary* if it is associated with an underlying disorder; autoimmune disorders, particularly systemic lupus erythematosus (SLE), and infections, such as HIV and hepatitis C, are common causes. The association of ITP with *Helicobacter pylori* infection is unclear.

ITP is characterized by mucocutaneous bleeding and a low, often very low, platelet count, with an otherwise normal peripheral blood cells and smear. Patients usually present either with ecchymoses and petechiae, or with thrombocytopenia incidentally found on a routine CBC. Mucocutaneous bleeding, such as oral mucosa, gastrointestinal, or heavy menstrual bleeding, may be present. Rarely, life-threatening, including central nervous system, bleeding can occur. Wet purpura (blood blisters in the mouth) and retinal hemorrhages may herald life-threatening bleeding.

LABORATORY TESTING IN ITP Laboratory testing for antibodies (serologic testing) is usually not helpful due to the low sensitivity and specificity of the current tests. Bone marrow examination can be reserved for those who have other signs or laboratory abnormalities not explained by ITP or in patients who do not respond to initial therapy. The peripheral blood smear may show large platelets, with otherwise normal morphology. Depending on the bleeding history, iron-deficiency anemia may be present.

Laboratory testing is performed to evaluate for secondary causes of ITP and should include testing for HIV infection and hepatitis C (and other infections if indicated). Serologic testing for SLE, serum protein electrophoresis, immunoglobulin levels to potentially detect hypogammaglobulinemia, selective testing for IgA deficiency or monoclonal gammopathies, and testing for *H. pylori* infection should be considered, depending on the clinical circumstance. If anemia is present, direct antiglobulin testing (Coombs' test) should be performed to rule out combined autoimmune hemolytic anemia with ITP (Evans' syndrome).

TREATMENT IMMUNE THROMBOCYTOPENIC PURPURA

The treatment of ITP uses drugs that decrease reticuloendothelial uptake of the antibody-bound platelet, decrease antibody production, and/or increase platelet production. The diagnosis of ITP does not necessarily mean that treatment must be instituted. Patients with platelet counts $>30,000/\mu\text{L}$ appear not to have increased mortality related to the thrombocytopenia.

Initial treatment in patients without significant bleeding symptoms, severe thrombocytopenia ($<5000/\mu\text{L}$), or signs of impending bleeding (such as retinal hemorrhage or large oral mucosal hemorrhages) can be instituted as an outpatient using single agents. Traditionally, this has been prednisone at 1 mg/kg, although Rh₀(D) immune globulin therapy (WinRho SDF), at 50–75 $\mu\text{g}/\text{kg}$, is also being used in this setting. Rh₀(D) immune globulin must be used only in Rh-positive patients because the mechanism of action is production of limited hemolysis, with antibody-coated cells "saturating" the Fc receptors, inhibiting Fc receptor function. Monitoring patients for 8 h after infusion is now advised by the FDA because of the rare complication of severe intravascular hemolysis. Intravenous gamma globulin (IVIgG), which is pooled, primarily IgG antibodies, also blocks the Fc receptor system, but appears to work primarily through different mechanism(s). IVIgG has more efficacy than anti-Rh₀(D) in postsplenectomized patients. IVIgG is dosed at 1–2 g/kg total, given over 1–5 days. Side effects are usually related to the volume of infusion and infrequently include aseptic meningitis and renal failure. All immunoglobulin preparations are derived from human plasma and undergo treatment for viral inactivation.

For patients with severe ITP and/or symptoms of bleeding, hospital admission and combined-modality therapy is given using high-dose glucocorticoids with IVIgG or anti-Rh₀(D) therapy and, as needed, additional immunosuppressive agents. Rituximab, an anti-CD20 (B cell) antibody, has shown efficacy in the treatment of refractory ITP, although long-lasting remission only occurs in approximately 30% of patients.

Splenectomy has been used for treatment of patients who relapse after glucocorticoids are tapered. Splenectomy remains an important treatment option; however, more patients than previously thought will go into a remission over time. Observation, if the platelet count is high enough, or intermittent treatment with anti-Rh₀(D) or IVIgG, or initiation of treatment with a TPO receptor agonist (see below) may be a reasonable approach to see if the ITP will resolve. Vaccination against encapsulated organisms (especially pneumococcus, but also meningococcus and *Haemophilus influenzae*, depending on patient age and potential exposure) is recommended before splenectomy. Accessory spleen(s) are a very rare cause of relapse.

TPO receptor agonists are now available for the treatment of ITP. This approach stems from the finding that many patients with ITP do not have increased TPO levels, as was previously hypothesized. TPO

levels reflect megakaryocyte mass, which is usually normal in ITP. TPO levels are not increased in the setting of platelet destruction. Two agents, one administered subcutaneously (romiplostim) and another orally (eltrombopag), are effective in raising platelet counts in patients with ITP and are recommended for adults at risk of bleeding who relapse after splenectomy or who have been unresponsive to at least one other therapy, particularly in those who have a contraindication to splenectomy. However, with the recognition that ITP will resolve spontaneously in some adult patients, short-term treatment with a TPO agonist can be considered before splenectomy in patients who need therapy.

Inherited Thrombocytopenia Thrombocytopenia is rarely inherited, either as an isolated finding or as part of a syndrome, and may be inherited in an autosomal dominant, autosomal recessive, or X-linked pattern. Many forms of autosomal dominant thrombocytopenia are now known to be associated with mutations in the nonmuscle myosin heavy chain *MYH9* gene. Interestingly, these include the May-Hegglin anomaly, and Sebastian, Epstein's, and Fechtner syndromes, all of which have distinct distinguishing features. A common feature of these disorders is large platelets (Fig. 140-1C). Autosomal recessive disorders include congenital amegakaryocytic thrombocytopenia, thrombocytopenia with absent radii, and Bernard-Soulier syndrome. The latter is primarily a functional platelet disorder due to absence of Gp Ib-IX-V, the VWF adhesion receptor. X-linked disorders include Wiskott-Aldrich syndrome and a dysmaturational syndrome resulting from a mutation in *GATA-1*, an important transcriptional regulator of hematopoiesis.

THROMBOTIC THROMBOCYTOPENIC PURPURA AND HEMOLYTIC-UREMIC SYNDROME

Thrombotic thrombocytopenic microangiopathies are a group of disorders characterized by thrombocytopenia, a microangiopathic hemolytic anemia evident by fragmented RBCs (Fig. 140-1D) and laboratory evidence of hemolysis, and microvascular thrombosis. They include thrombotic thrombocytopenic purpura (TTP) and hemolytic-uremic syndrome (HUS), as well as syndromes complicating bone marrow transplantation, certain medications and infections, pregnancy, and vasculitis. In DIC, although thrombocytopenia and microangiopathy are seen, a coagulopathy predominates, with consumption of clotting factors and fibrinogen resulting in an elevated prothrombin time (PT) and often activated partial thromboplastin time (aPTT). The PT and aPTT are characteristically normal in TTP or HUS.

Thrombotic Thrombocytopenic Purpura TTP and HUS were previously considered overlap syndromes. However, in the past few years, the pathophysiology of inherited and idiopathic TTP has become better understood and clearly differs from HUS. TTP was first described in 1924 by Eli Moschcowitz and characterized by a pentad of findings that include microangiopathic hemolytic anemia, thrombocytopenia, renal failure, neurologic findings, and fever. The full-blown syndrome is less commonly seen now, probably due to earlier diagnosis. The introduction of treatment with plasma exchange markedly improved the prognosis in patients, with a decrease in mortality from 85–100% to 10–30%.

The pathogenesis of inherited (Upshaw-Schulman syndrome) and idiopathic TTP is related to a deficiency of, or antibodies to, the metalloprotease ADAMTS13, which cleaves VWF. VWF is normally secreted as ultra-large multimers, which are then cleaved by ADAMTS13. The persistence of ultra-large VWF molecules is thought to contribute to pathogenic platelet adhesion and aggregation (Fig. 140-4). This defect alone, however, is not sufficient to result in TTP because individuals with a congenital absence of ADAMTS13 develop TTP only episodically. Additional provocative factors have not been defined. The level of ADAMTS13 activity, as well as antibodies, can now be detected by laboratory assays. Although assays with sufficient sensitivity and specificity to direct clinical management have yet to be clearly defined, ADAMTS13 activity levels of <10% are more clearly associated with idiopathic TTP.

VWF and Platelet Adhesion

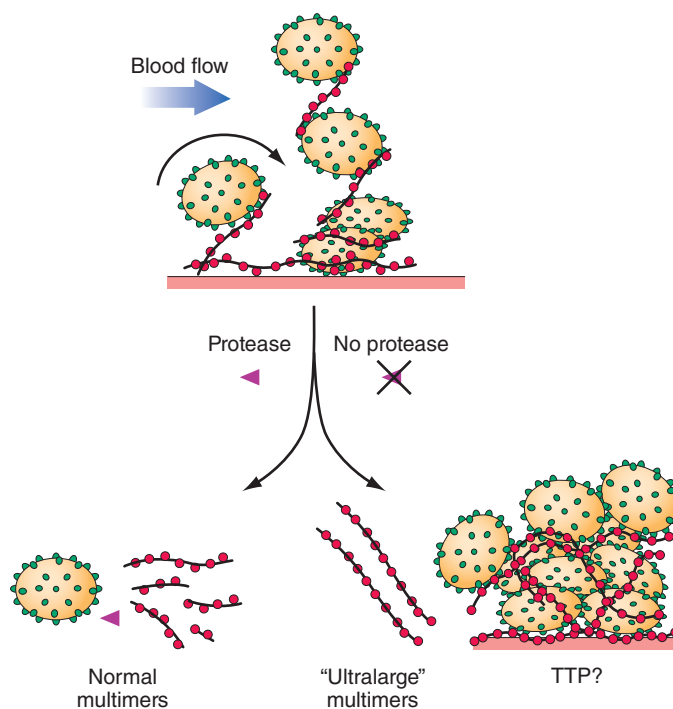


FIGURE 140-4 Pathogenesis of thrombotic thrombocytopenic purpura (TTP). Normally the ultra-high-molecular-weight multimers of von Willebrand factor (VWF) produced by the endothelial cells are processed into smaller multimers by a plasma metalloproteinase called ADAMTS13. In TTP, the activity of the protease is inhibited, and the ultra-high-molecular-weight multimers of VWF initiate platelet aggregation and thrombosis.

Idiopathic TTP appears to be more common in women than in men. No geographic or racial distribution has been defined. TTP is more common in patients with HIV infection and in pregnant women. TTP in pregnancy is not clearly related to ADAMTS13. Medication-related microangiopathic hemolytic anemia may be secondary to antibody formation (ticlopidine and possibly clopidogrel) or direct endothelial toxicity (cyclosporine, mitomycin C, tacrolimus, quinine), although this is not always so clear, and fear of withholding treatment, as well as lack of other treatment alternatives, results in broad application of plasma exchange. However, withdrawal, or reduction in dose, of endothelial toxic agents usually decreases the microangiopathy.

TREATMENT

THROMBOTIC THROMBOCYTOPENIC PURPURA

TTP is a devastating disease if not diagnosed and treated promptly. In patients presenting with new thrombocytopenia, with or without evidence of renal insufficiency and other elements of classic TTP, laboratory data should be obtained to rule out DIC and to evaluate for evidence of microangiopathic hemolytic anemia. Findings to support the TTP diagnosis include an increased lactate dehydrogenase and indirect bilirubin, decreased haptoglobin, and increased reticulocyte count, with a negative direct antiglobulin test. The peripheral smear should be examined for evidence of schistocytes (Fig. 140-1D). Polychromasia is usually also present due to the increased number of young red blood cells, and nucleated RBCs are often present, which is thought to be due to infarction in the microcirculatory system of the bone marrow.

Plasma exchange remains the mainstay of treatment of TTP. ADAMTS13 antibody-mediated TTP (idiopathic TTP) appears to respond best to plasma exchange. Plasma exchange is continued

730 until the platelet count is normal and signs of hemolysis are resolved for at least 2 days. Although never evaluated in clinical trials, the use of glucocorticoids seems a reasonable approach, but should only be used as an adjunct to plasma exchange. Additionally, other immunomodulatory therapies have been reported to be successful in refractory or relapsing TTP, including rituximab, vincristine, cyclophosphamide, and splenectomy. A significant relapse rate is noted; 25–45% of patients relapse within 30 days of initial “remission,” and 12–40% of patients have late relapses. Relapses are more frequent in patients with severe ADAMTS13 deficiency at presentation.

Hemolytic-Uremic Syndrome HUS is a syndrome characterized by acute renal failure, microangiopathic hemolytic anemia, and thrombocytopenia. It is seen predominantly in children and in most cases is preceded by an episode of diarrhea, often hemorrhagic in nature. *Escherichia coli* O157:H7 is the most frequent, although not only, etiologic serotype. HUS not associated with diarrhea is more heterogeneous in presentation and course. Atypical HUS (aHUS) due to genetic defects that result in chronic complement activation has been defined, and screening for mutations in complement regulatory genes is available.

TREATMENT HEMOLYTIC-UREMIC SYNDROME

Treatment of HUS is primarily supportive. In HUS associated with diarrhea, many (~40%) children require at least some period of support with dialysis; however, the overall mortality is <5%. In HUS not associated with diarrhea, the mortality is higher, approximately 26%. Plasma infusion or plasma exchange has not been shown to alter the overall course. ADAMTS13 levels are generally reported to be normal in HUS, although occasionally they have been reported to be decreased. In patients with atypical HUS, eculizumab therapy increases the platelet count and preserves renal function.

THROMBOCYTOSIS

Thrombocytosis is almost always due to (1) iron deficiency; (2) inflammation, cancer, or infection (reactive thrombocytosis); or (3) an underlying myeloproliferative process (essential thrombocythemia or polycythemia vera) (Chap. 131) or, rarely, the 5q- myelodysplastic process (Chap. 130). Patients presenting with an elevated platelet count should be evaluated for underlying inflammation or malignancy, and iron deficiency should be ruled out. Thrombocytosis in response to acute or chronic inflammation has not been clearly associated with an increased thrombotic risk. In fact, patients with markedly elevated platelet counts (>1.5 million), usually seen in the setting of a myeloproliferative disorder, have an increased risk of bleeding. This appears to be due, at least in part, to acquired von Willebrand disease (VWD) due to platelet-VWF binding and removal from the circulation.

QUALITATIVE DISORDERS OF PLATELET FUNCTION

Inherited Disorders of Platelet Function Inherited platelet function disorders are thought to be relatively rare, although the prevalence of mild disorders of platelet function is unclear, in part because our testing for such disorders is suboptimal. Rare qualitative disorders include the autosomal recessive disorders Glanzmann’s thrombasthenia (absence of the platelet Gp IIb/IIIa receptor) and Bernard-Soulier syndrome (absence of the platelet Gp Ib-IX-V receptor). Both are inherited in an autosomal recessive fashion and present with bleeding symptoms in childhood.

Platelet storage pool disorder (SPD) is the classic autosomal dominant qualitative platelet disorder. This results from abnormalities of platelet granule formation. It is also seen as a part of inherited disorders of granule formation, such as Hermansky-Pudlak syndrome. Bleeding symptoms in SPD are variable, but often are mild. The most common inherited disorders of platelet function prevent normal secretion of granule content and are termed *secretion defects*. Few of these abnormalities have been dissected at the molecular level but they likely result from various mutations.

TREATMENT INHERITED DISORDERS OF PLATELET DYSFUNCTION

Bleeding symptoms or prevention of bleeding in patients with severe platelet dysfunction frequently requires platelet transfusion. Care is taken to limit the risk of alloimmunization by limiting exposure, using HLA-matched leuko-depleted platelet concentrates for transfusion. Platelet disorders associated with milder bleeding symptoms frequently respond to desmopressin (1-deamino-8-D-arginine vasopressin [DDAVP]). DDAVP increases plasma VWF and factor VIII levels; it may also have a direct effect on platelet function. Particularly for mucosal bleeding symptoms, antifibrinolytic therapy (ε-aminocaproic acid or tranexamic acid) is used alone or in conjunction with DDAVP or platelet therapy.

Acquired Disorders of Platelet Function Acquired platelet dysfunction is common, usually due to medications, either intentionally as with antiplatelet therapy or unintentionally as with high-dose penicillins. Acquired platelet dysfunction occurs in uremia. This is likely multifactorial, but the resultant effect is defective adhesion and activation. The platelet defect is improved most by dialysis but may also be improved by increasing the hematocrit to 27–32%, giving DDAVP (0.3 µg/kg), or use of conjugated estrogens. Platelet dysfunction also occurs with cardiopulmonary bypass due to the effect of the artificial circuit on platelets, and bleeding symptoms respond to platelet transfusion. Platelet dysfunction seen with underlying hematologic disorders can result from nonspecific interference by circulating paraproteins or intrinsic platelet defects in myeloproliferative and myelodysplastic syndromes.

VON WILLEBRAND DISEASE

VWD is the most common inherited bleeding disorder. Estimates from laboratory data suggest a prevalence of approximately 1%, but data based on symptomatic individuals suggest that it is closer to 0.1% of the population. VWF serves two roles: (1) as the major adhesion molecule that tethers the platelet to the exposed subendothelium; and (2) as the binding protein for factor VIII (FVIII), resulting in significant prolongation of the FVIII half-life in circulation. The platelet-adhesive function of VWF is critically dependent on the presence of large VWF multimers, whereas FVIII binding is not. Most of the symptoms of VWD are “platelet-like” except in more severe VWD when the FVIII is low enough to produce symptoms similar to those found in FVIII deficiency (hemophilia A).

VWD has been classified into three major types, with four subtypes of type 2 (Table 140-2; Fig. 140-5). By far the most common type of VWD is type 1 disease, with a parallel decrease in VWF protein, VWF function, and FVIII levels, accounting for at least 80% of cases. Patients have predominantly mucosal bleeding symptoms, although postoperative bleeding can also be seen. Bleeding symptoms are very uncommon in infancy and usually manifest later in childhood with excessive bruising and epistaxis. Because these symptoms occur commonly in childhood, the clinician should particularly note bruising at sites unlikely to be traumatized and/or prolonged epistaxis requiring medical attention. Menorrhagia is a common manifestation of VWD. Menstrual bleeding resulting in anemia should warrant an evaluation for VWD and, if negative, functional platelet disorders. Frequently, mild type 1 VWD first manifests with dental extractions, particularly wisdom tooth extraction, or tonsillectomy.

Not all patients with low VWF levels have bleeding symptoms. Whether patients bleed or not will depend on the overall hemostatic balance they have inherited, along with environmental influences and the type of hemostatic challenges they experience. Although the inheritance of VWD is autosomal, many factors modulate both VWF levels and bleeding symptoms. These have not all been defined, but include blood type, thyroid hormone status, race, stress, exercise, and hormonal (both endogenous and exogenous) influences. Patients with type O blood have VWF protein levels of approximately one-half that of patients with AB blood type; and, in fact, the normal range for

TABLE 140-2 LABORATORY DIAGNOSIS OF VON WILLEBRAND DISEASE (VWD)

Type	aPTT	VWF Antigen	VWF Activity	FVIII Activity	Multimer
1	NI or ↑	↓	↓	↓	Normal distribution, decreased in quantity
2A	NI or ↑	↓	↓↓	↓	Loss of high- and intermediate-MW multimers
2B ^a	NI or ↑	↓	↓↓	↓	Loss of high-MW multimers
2M	NI or ↑	↓	↓↓	↓	Normal distribution, decreased in quantity
2N	↑↑	NI or ↓ ^b	NI or ↓ ^b	↓↓	Normal distribution
3	↑↑	↓↓	↓↓	↓↓	Absent

^aUsually also decreased platelet count. ^bFor type 2N, in the homozygous state, FVIII is very low; in the heterozygous state, it is only seen in conjunction with type 1 VWD.

Abbreviations: aPTT, activated partial thromboplastin time; F, factor; MW, molecular weight; NI, normal; VWF, von Willebrand factor.

patients with type O blood overlaps that which has been considered diagnostic for VWD. A mildly decreased VWF level should be viewed more as a risk factor for bleeding than as an actual disease.

Patients with type 2 VWD have functional defects; thus, the VWF antigen measurement is significantly higher than the test of function. For types 2A, 2B, and 2M VWD, platelet-binding and/or collagen binding VWF activity is decreased. In type 2A VWD, the impaired function is due either to increased susceptibility to cleavage by ADAMTS13,

resulting in loss of intermediate- and high-molecular-weight multimers, or to decreased secretion of these multimers by the cell. Type 2B VWD results from gain-of-function mutations that result in increased spontaneous binding of VWF to platelets in circulation, with subsequent clearance of this complex by the reticuloendothelial system. The resulting VWF in the patients' plasma lacks the highest molecular-weight multimers, and the platelet count is usually modestly reduced. Type 2M occurs as a consequence of a group of mutations that cause dysfunction but do not affect multimer structure.

Type 2N VWD is due to mutations in VWF that affect binding of FVIII.

As FVIII is stabilized by binding to VWF, the FVIII in patients with type 2N VWD has a very short half-life, and the FVIII level is markedly decreased. This is sometimes termed *autosomal hemophilia*. Type 3 VWD, or severe VWD, describes patients with virtually no VWF protein and FVIII levels <10%. Patients experience mucosal and joint bleeding, surgery-related bleeding, and other bleeding symptoms. Some patients with type 3 VWD, particularly those with large VWF gene deletions, are at risk of developing antibodies to infused VWF.

Acquired VWD is a rare disorder, most commonly seen in patients with underlying lymphoproliferative disorders, including monoclonal gammopathies of underdetermined significance (MGUS), multiple myeloma, and Waldenström's macroglobulinemia. It is seen most commonly in the setting of MGUS and should be suspected in patients, particularly elderly patients, with a new onset of severe mucosal bleeding symptoms. Laboratory evidence of acquired VWD is found in some patients with aortic valvular disease. Heyde's syndrome (aortic stenosis with gastrointestinal bleeding) is attributed to the presence of angiodysplasia of the gastrointestinal tract in patients with aortic stenosis. The shear stress on blood passing through the stenotic aortic valve appears to produce a change in VWF, making it susceptible to serum proteases. Consequently, large multimer forms are lost, leading to an acquired type 2 VWD, but return when the stenotic valve is replaced.

TREATMENT VON WILLEBRAND DISEASE

The mainstay of treatment for type 1 VWD is DDAVP (desmopressin), which results in release of VWF and FVIII from endothelial stores. DDAVP can be given intravenously or by a high-concentration intranasal spray (1.5 mg/mL). The peak activity when given intravenously is approximately 30 min, whereas it is 2 h when given intranasally. The usual dose is 0.3 µg/kg intravenously or two squirts (one in each nostril) for patients >50 kg (one squirt for those <50 kg). It is recommended that patients with VWD be tested with DDAVP to assess their response before using it. In patients who respond well (increase in laboratory values of two- to fourfold), it can be used for procedures with minor to moderate risk of bleeding. Depending on the procedure, additional doses may be needed; it is usually given every 12–24 h. Less frequent dosing may result in less tachyphylaxis, which occurs when synthesis cannot compensate for the released stores. The major side effect of DDAVP is hyponatremia due to decreased free water clearance. This occurs most commonly in the very young and the very old, but fluid restriction should be advised for all patients for the 24 h following each dose.

Some patients with types 2A and 2M VWD respond to DDAVP such that it can be used for minor procedures. For the other subtypes, for type 3 disease, and for major procedures requiring longer periods of normal hemostasis, VWF replacement can be given. Virally inactivated VWF-containing factor concentrates are safer than cryoprecipitate as the replacement product.

Antifibrinolytic therapy using either ε-aminocaproic acid or tranexamic acid is an important therapy, either alone or in an

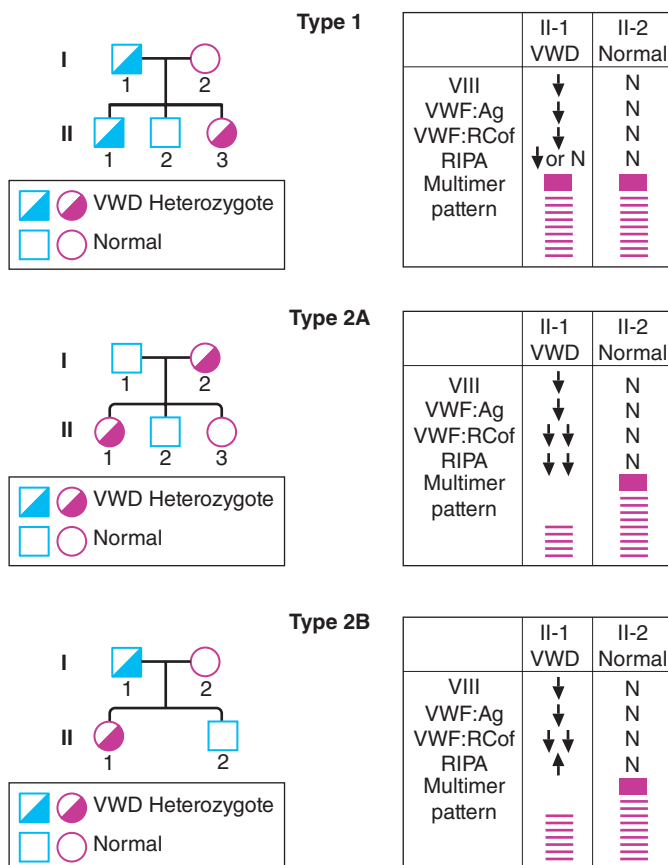


FIGURE 140-5 Pattern of inheritance and laboratory findings in von Willebrand disease (VWD). The assays of platelet function include a coagulation assay of factor VIII bound and carried by von Willebrand factor (VWF), abbreviated as VIII; immunoassay of total VWF protein (VWF:Ag); bioassay of the ability of patient plasma to support ristocetin-induced agglutination of normal platelets (VWF:RCof); and ristocetin-induced aggregation of patient platelets, abbreviated RIPA. The multimer pattern illustrates the protein bands present when plasma is electrophoresed in a polyacrylamide gel. The II-1 and II-2 columns refer to the phenotypes of the second-generation offspring.

732 adjunctive capacity, particularly for the prevention or treatment of mucosal bleeding. These agents are particularly useful in prophylaxis for dental procedures, with DDAVP for dental extractions and tonsillectomy, menorrhagia, and prostate procedures. It is contraindicated in the setting of upper urinary tract bleeding, due to the risk of ureteral obstruction.

141 Coagulation Disorders

Valder R. Arruda, Katherine A. High

DISORDERS OF THE VESSEL WALL

The vessel wall is an integral part of hemostasis, and separation of a fluid phase is artificial, particularly in disorders such as TTP or HIT that clearly involve the endothelium as well. Inflammation localized to the vessel wall, such as vasculitis, and inherited connective tissue disorders are abnormalities inherent to the vessel wall.

METABOLIC AND INFLAMMATORY DISORDERS Acute febrile illnesses may result in vascular damage. This can result from immune complexes containing viral antigens or the viruses themselves. Certain pathogens, such as the rickettsiae causing Rocky Mountain spotted fever, replicate in endothelial cells and damage them. Vascular purpura may occur in patients with polyclonal gammopathies but more commonly in those with monoclonal gammopathies, including Waldenström's macroglobulinemia, multiple myeloma, and cryoglobulinemia. Patients with mixed cryoglobulinemia develop a more extensive maculopapular rash due to immune complex-mediated damage to the vessel wall.

Patients with scurvy (vitamin C deficiency) develop painful episodes of perifollicular skin bleeding as well as more systemic bleeding symptoms. Vitamin C is needed to synthesize hydroxyproline, an essential constituent of collagen. Patients with Cushing's syndrome or on chronic glucocorticoid therapy develop skin bleeding and easy bruising due to atrophy of supporting connective tissue. A similar phenomenon is seen with aging, where following minor trauma, blood spreads superficially under the epidermis. This has been termed *senile purpura*. It is most common on skin that has been previously damaged by sun exposure.

Henoch-Schönlein, or anaphylactoid, purpura is a distinct, self-limited type of vasculitis that occurs in children and young adults. Patients have an acute inflammatory reaction with IgA and complement components in capillaries, mesangial tissues, and small arterioles leading to increased vascular permeability and localized hemorrhage. The syndrome is often preceded by an upper respiratory infection, commonly with streptococcal pharyngitis, or is triggered by drug or food allergies. Patients develop a purpuric rash on the extensor surfaces of the arms and legs, usually accompanied by polyarthralgias or arthritis, abdominal pain, and hematuria from focal glomerulonephritis. All coagulation tests are normal, but renal impairment may occur. Glucocorticoids can provide symptomatic relief but do not alter the course of the illness.

INHERITED DISORDERS OF THE VESSEL WALL Patients with inherited disorders of the connective tissue matrix, such as Marfan's syndrome, Ehlers-Danlos syndrome, and pseudoxanthoma elasticum, frequently report easy bruising. Inherited vascular abnormalities can result in increased bleeding. This is notably seen in hereditary hemorrhagic telangiectasia (HHT, or Osler-Weber-Rendu disease), a disorder where abnormal telangiectatic capillaries result in frequent bleeding episodes, primarily from the nose and gastrointestinal tract. Arteriovenous malformation (AVM) in the lung, brain, and liver may also occur in HHT. The telangiectasia can often be visualized on the oral and nasal mucosa. Signs and symptoms develop over time. Epistaxis begins, on average, at the age of 12 and occurs in >95% of affected individuals by middle age. Two genes involved in the pathogenesis are *eng* (endoglin) on chromosome 9q33-34 (so-called HHT type 1), associated with pulmonary AVM in 40% of cases, and *alk1* (activin-receptor-like kinase 1) on chromosome 12q13, associated with a much lower risk of pulmonary AVM.

ACKNOWLEDGMENT

Robert Handin, MD, contributed this chapter in the 16th edition and some materials from his chapter are included here.

Deficiencies of coagulation factors have been recognized for centuries. Patients with genetic deficiencies of plasma coagulation factors exhibit life-long recurrent bleeding episodes into joints, muscles, and closed spaces, either spontaneously or following an injury. The most common inherited factor deficiencies are the hemophilias, X-linked diseases caused by deficiency of factor (F) VIII (hemophilia A) or FIX (hemophilia B). Rare congenital bleeding disorders due to deficiencies of other factors, including FII (prothrombin), FV, FVII, FX, FXI, and FXIII, and fibrinogen are commonly inherited in an autosomal recessive manner (Table 141-1). Advances in characterization of the molecular bases of clotting factor deficiencies have contributed to better understanding of the disease phenotypes and may eventually allow more targeted therapeutic approaches through the development of small molecules, recombinant proteins, or cell and gene-based therapies.

Commonly used tests of hemostasis provide the initial screening for clotting factor activity (Fig. 141-1), and disease phenotype often correlates with the level of clotting activity. An isolated abnormal prothrombin time (PT) suggests FVII deficiency, whereas a prolonged activated partial thromboplastin time (aPTT) indicates most commonly hemophilia or FXI deficiency (Fig. 141-1). The prolongation of both PT and aPTT suggests deficiency of FV, FX, FII, or fibrinogen abnormalities. The addition of the missing factor at a range of doses to the subject's plasma will correct the abnormal clotting times; the result is expressed as a percentage of the activity observed in normal subjects.

Acquired deficiencies of plasma coagulation factors are more frequent than congenital disorders; the most common disorders include hemorrhagic diathesis of liver disease, disseminated intravascular coagulation (DIC), and vitamin K deficiency. In these disorders, blood coagulation is hampered by the deficiency of more than one clotting factor, and the bleeding episodes are the result of perturbation of both primary (coagulation) and secondary (e.g., platelet and vessel wall interactions) hemostasis.

The development of antibodies to coagulation plasma proteins, clinically termed *inhibitors*, is a relatively rare disease that often affects hemophilia A or B and FXI-deficient patients on repetitive exposure to the missing protein to control bleeding episodes. Inhibitors also occur among subjects without genetic deficiency of clotting factors (e.g., in the postpartum setting as a manifestation of underlying autoimmune or neoplastic disease or idiopathically). Rare cases of inhibitors to thrombin or FV have been reported in patients receiving topical bovine thrombin preparation as a local hemostatic agent in complex surgeries. The diagnosis of inhibitors is based on the same tests as those used to diagnose inherited plasma coagulation factor deficiencies. However, the addition of the missing protein to the plasma of a subject with an inhibitor does not correct the abnormal aPTT and/or PT tests (known as mixing tests). This is the major laboratory difference between deficiencies and inhibitors. Additional tests are required to measure the specificity of the inhibitor and its titer.

The treatment of these bleeding disorders often requires replacement of the deficient protein using recombinant or purified plasma-derived products or fresh-frozen plasma (FFP). Therefore, it is imperative to arrive at a proper diagnosis to optimize patient care without unnecessary exposure to suboptimal treatment and the risks of bloodborne disease.

HEMOPHILIA

PATHOGENESIS AND CLINICAL MANIFESTATIONS

Hemophilia is an X-linked recessive hemorrhagic disease due to mutations in the *F8* gene (hemophilia A or classic hemophilia) or *F9* gene (hemophilia B). The disease affects 1 in 10,000 males worldwide, in all ethnic groups; hemophilia A represents 80% of all cases. Male subjects

TABLE 141-1 GENETIC AND LABORATORY CHARACTERISTICS OF INHERITED COAGULATION DISORDERS

Clotting Factor Deficiency	Inheritance	Prevalence in General Population	Laboratory Abnormality ^a			Minimum Hemostatic Levels	Treatment	Plasma Half-Life
			aPTT	PT	TT			
Fibrinogen	AR	1 in 1,000,000	+	+	+	100 mg/dL	Cryoprecipitate	2–4 d
Prothrombin	AR	1 in 2,000,000	+	+	–	20–30%	FFP/PCC	3–4 d
Factor V	AR	1 in 1,000,000	+/-	+/-	–	15–20%	FFP	36 h
Factor VII	AR	1 in 500,000	–	+	–	15–20%	FFP/PCC	4–6 h
Factor VIII	X-linked	1 in 5,000	+	–	–	30%	FVIII concentrates	8–12 h
Factor IX	X-linked	1 in 30,000	+	–	–	30%	FIX concentrates	18–24 h
Factor X	AR	1 in 1,000,000	+/-	+/-	–	15–20%	FFP/PCC	40–60 h
Factor XI	AR	1 in 1,000,000	+	–	–	15–20%	FFP	40–70 h
Factor XII	AR	ND	+	–	–	^b	^b	60 h
HK	AR	ND	+	–	–	^b	^b	150 h
Prekallikrein	AR	ND	+	–	–	^b	^b	35 h
Factor XIII	AR	1 in 2,000,000	–	–	+/-	2–5%	Cryoprecipitate/FXIII concentrates	11–14 d

^aValues within normal range (–) or prolonged (+). ^bNo risk for bleeding; treatment is not indicated.

Abbreviations: aPTT, activated partial thromboplastin time; AR, autosomal recessive; FFP, fresh-frozen plasma; HK, high-molecular-weight kininogen; ND, not determined; PCC, prothrombin complex concentrates; PT, prothrombin time; TT, thrombin time.

are clinically affected; women, who carry a single mutated gene, are generally asymptomatic. Family history of the disease is absent in ~30% of cases, and in these cases, 80% of the mothers are carriers of the *de novo* mutated allele. More than 500 different mutations have been identified in the *F8* or *F9* genes of patients with hemophilia A or B, respectively. One of the most common hemophilia A mutations results from an inversion of the intron 22 sequence, and it is present in 40% of cases of severe hemophilia A. Advances in molecular diagnosis now permit precise identification of mutations, allowing accurate diagnosis of women carriers of the hemophilia gene in affected families.

Clinically, hemophilia A and hemophilia B are indistinguishable. The disease phenotype correlates with the residual activity of FVIII or FIX and can be classified as severe (<1%), moderate (1–5%), or mild (6–30%). In the severe and moderate forms, the disease is characterized by bleeding into the joints (hemarthrosis), soft tissues, and

muscles after minor trauma or even spontaneously. Patients with mild disease experience infrequent bleeding that is usually secondary to trauma. Among those with residual FVIII or FIX activity >25% of normal, the disease is discovered only by bleeding after major trauma or during routine presurgery laboratory tests. Typically, the global tests of coagulation show only an isolated prolongation of the aPTT assay. Patients with hemophilia have normal bleeding times and platelet counts. The diagnosis is made after specific determination of FVIII or FIX clotting activity.

Early in life, bleeding may present after circumcision or rarely as intracranial hemorrhages. The disease is more evident when children begin to walk or crawl. In the severe form, the most common bleeding manifestations are the recurrent hemarthroses, which can affect every joint but mainly affect knees, elbows, ankles, shoulders, and hips. Acute hemarthroses are painful, and clinical signs are local swelling and erythema. To avoid pain, the patient may adopt a fixed position, which leads eventually to muscle contractures. Very young children unable to communicate verbally show irritability and a lack of movement of the affected joint. Chronic hemarthroses are debilitating, with synovial thickening and synovitis in response to the intraarticular blood. After a joint has been damaged, recurrent bleeding episodes result in the clinically recognized “target joint,” which then establishes a vicious cycle of bleeding, resulting in progressive joint deformity that in critical cases requires surgery as the only therapeutic option. Hematomas into the muscle of distal parts of the limbs may lead to external compression of arteries, veins, or nerves that can evolve to a compartment syndrome.

Bleeding into the oropharyngeal spaces, central nervous system (CNS), or retroperitoneum is life threatening and requires immediate therapy. Retroperitoneal hemorrhages can accumulate large quantities of blood with formation of masses with calcification and inflammatory tissue reaction (pseudotumor syndrome) and also result in damage to the femoral nerve. Pseudotumors can also form in bones, especially long bones of the lower limbs. Hematuria is frequent among hemophilia patients, even in the absence of genitourinary pathology. It is often self-limited and may not require specific therapy.

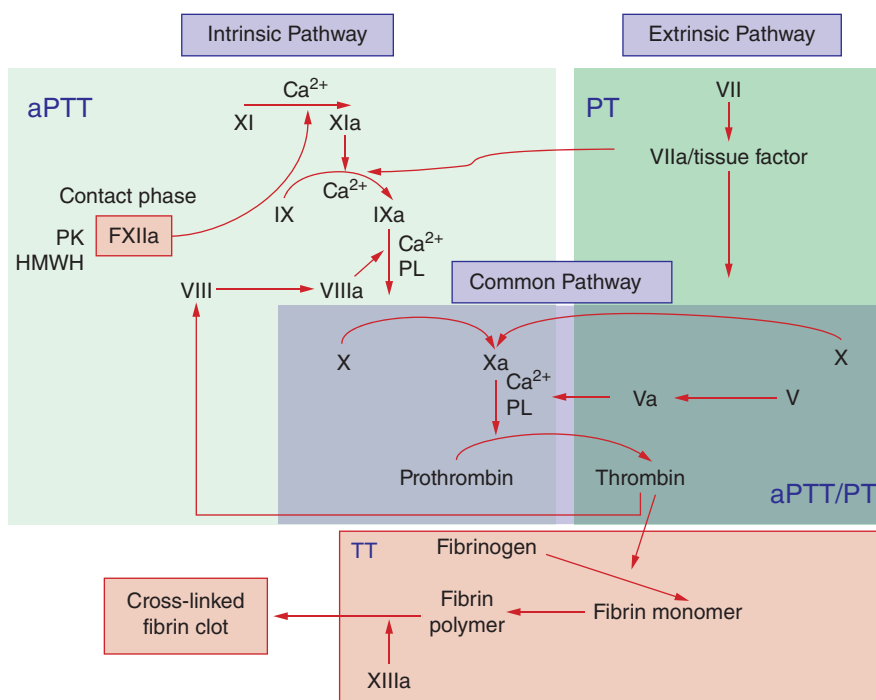


FIGURE 141-1 Coagulation cascade and laboratory assessment of clotting factor deficiency by activated partial thromboplastin time (aPTT), prothrombin time (PT), thrombin time (TT), and phospholipid (PL).

Without treatment, severe hemophilia has a limited life expectancy. Advances in the blood fractionation industry during World War II resulted in the realization that plasma could be used to treat hemophilia, but the volumes required to achieve even modest elevation of circulating factor levels limit the utility of plasma infusion as an approach to disease management. The discovery in the 1960s that the cryoprecipitate fraction of plasma was enriched for FVIII, and the eventual purification of FVIII and FIX from plasma, led to the introduction of home infusion therapy with factor concentrates in the 1970s. The availability of factor concentrates resulted in a dramatic improvement in life expectancy and in quality of life for people with severe hemophilia. However, the contamination of the blood supply with hepatitis viruses and, subsequently, HIV resulted in widespread transmission of these bloodborne infections within the hemophilia population; complications of HIV and of hepatitis C are now the leading causes of death among U.S. adults with severe hemophilia. The introduction of viral inactivation steps in the preparation of plasma-derived products in the mid-1980s greatly reduced the risk of HIV and hepatitis, and the risks were further reduced by the successful production of recombinant FVIII and FIX proteins, both licensed in the 1990s. It is uncommon for hemophilic patients born after 1985 to have contracted either hepatitis or HIV, and for these individuals, life expectancy is approximately 65 years. In fact, since 1998, no evidence of new infections with viral hepatitis or HIV has been reported in patients using blood products. Factor replacement therapy for hemophilia can be provided either in response to a bleeding episode or as a prophylactic treatment. Primary prophylaxis is defined as a strategy for maintaining the missing clotting factor at levels ~1% or higher on a regular basis in order to prevent bleeds, especially the onset of hemarthroses. Hemophilic boys receiving regular infusions of FVIII (3 days/week) or FIX (2 days/week) can reach puberty without detectable joint abnormalities. Prophylaxis has become gradually more common in young patients. The Centers for Disease Control and Prevention reported that 51% of children with severe hemophilia who are younger than age 6 years receive prophylaxis, increasing considerably from 33% in 1995. Although highly recommended, the high cost and difficulties in accessing peripheral veins in young patients and the potential infectious and thrombotic risks of long-term central vein catheters are important limiting factors for many young patients. Emerging data show that prophylaxis is also increasing among adults with severe hemophilia.

General considerations regarding the treatment of bleeds in hemophilia include the following: (1) Treatment should begin as soon as possible because symptoms often precede objective evidence of bleeding; because of the superior efficacy of early therapeutic intervention, classic symptoms of bleeding into the joint in a reliable patient, headaches, or automobile or other accidents require prompt replacement and further laboratory investigation. (2) Drugs that hamper platelet function, such as aspirin or aspirin-containing drugs, should be avoided; to control pain, drugs such as ibuprofen or propoxyphene are preferred. FVIII and FIX are dosed in units. One unit is defined as amount of FVIII (100 ng/mL) or FIX (5 µg/mL) in 1 mL of normal plasma. One unit of FVIII per kilogram of body weight increases the plasma FVIII level by 2%. One can calculate the dose needed to increase FVIII levels to 100% in a 70-kg severe hemophilia patient (<1%) using the simple formula below. Thus, 3500 units of FVIII will raise the circulating level to 100%.

$$\text{FVIII dose (IU)} = \frac{\text{Target FVIII levels} - \text{FVIII baseline levels}}{\text{body weight (kg)} \times 0.5 \text{ unit/kg}}$$

The doses for FIX replacement are different from those for FVIII, because FIX recovery after infusion is usually only 50% of the predicted value. Therefore, the formula for FIX replacement is as follows:

$$\text{FIX dose (IU)} = \frac{\text{Target FIX levels} - \text{FIX baseline levels}}{\text{body weight (kg)} \times 1 \text{ unit/kg}}$$

The FVIII half-life of 8–12 h requires injections twice a day to maintain therapeutic levels, whereas the FIX half-life is longer, ~24 h, so that once-a-day injection is sufficient. In specific situations such as after surgery, continuous infusion of factor may be desirable because of its safety in achieving sustained factor levels at a lower total cost.

Cryoprecipitate is enriched with FVIII protein (each bag contains ~80 IU of FVIII) and was commonly used for the treatment of hemophilia A decades ago; it is still in use in some developing countries, but because of the risk of bloodborne diseases, this product should be avoided in hemophilia patients when factor concentrates are available.

Mild bleeds such as uncomplicated hemarthroses or superficial hematomas require initial therapy with factor levels of 30–50%. Additional doses to maintain levels of 15–25% for 2 or 3 days are indicated for severe hemarthroses, especially when these episodes affect the “target joint.” Large hematomas, or bleeds into deep muscles, require factor levels of 50% or even higher if the clinical symptoms do not improve, and factor replacement may be required for a period of 1 week or longer. The control of serious bleeds including those that affect the oropharyngeal spaces, CNS, and the retroperitoneum require sustained protein levels of 50–100% for 7–10 days. Prophylactic replacement for surgery is aimed at achieving normal factor levels (100%) for a period of 7–10 days; replacement can then be tapered depending on the extent of the surgical wounds. Oral surgery is associated with extensive tissue damage that usually requires factor replacement for 1–3 days coupled with oral antifibrinolytic drugs.

NONTRANSFUSION THERAPY IN HEMOPHILIA

DDAVP (1-Amino-8-D-Arginine Vasopressin) DDAVP is a synthetic vasopressin analog that causes a transient rise in FVIII and von Willebrand factor (VWF), but not FIX, through a mechanism involving release from endothelial cells. Patients with moderate or mild hemophilia A should be tested to determine if they respond to DDAVP before a therapeutic application. DDAVP at doses of 0.3 µg/kg body weight, over a 20-min period, is expected to raise FVIII levels by two- to threefold over baseline, peaking between 30 and 60 min after infusion. DDAVP does not improve FVIII levels in severe hemophilia A patients, because there are no stores to release. Repeated dosing of DDAVP results in tachyphylaxis because the mechanism is an increase in release rather than de novo synthesis of FVIII and VWF. More than three consecutive doses become ineffective, and if further therapy is indicated, FVIII replacement is required to achieve hemostasis.

Antifibrinolytic Drugs Bleeding in the gums, gastrointestinal tract, and during oral surgery requires the use of oral antifibrinolytic drugs such as ε-amino caproic acid (EACA) or tranexamic acid to control local hemostasis. The duration of the treatment depending on the clinical indication is 1 week or longer. Tranexamic acid is given at doses of 25 mg/kg three to four times a day. EACA treatment requires a loading dose of 200 mg/kg (maximum of 10 g) followed by 100 mg/kg per dose (maximum 30 g/d) every 6 h. These drugs are not indicated to control hematuria because of the risk of formation of an occlusive clot in the lumen of genitourinary tract structures.

COMPLICATIONS

Inhibitor Formation The formation of alloantibodies to FVIII or FIX is currently the major complication of hemophilia treatment. The prevalence of inhibitors to FVIII is estimated to be between 5 and 10% of all cases and ~20% of severe hemophilia A patients. Inhibitors to FIX are detected in only 3–5% of all hemophilia B patients. The high-risk group for inhibitor formation includes severe deficiency (>80% of all cases of inhibitors), familial history of inhibitor, African descent, mutations in the FVIII or FIX gene resulting in deletion of large coding regions, or gross gene rearrangements. Inhibitors usually appear early in life, at a median of 2 years of age, and after 10 cumulative days of exposure. However, intensive replacement therapy such as for major surgery, intracranial bleeding, or trauma increases the risk of inhibitor formation for patients of all ages and degree of

clinical severity, which requires close laboratory monitoring in the following weeks.

The clinical diagnosis of an inhibitor is suspected when patients do not respond to factor replacement at therapeutic doses. Inhibitors increase both morbidity and mortality in hemophilia. Because early detection of an inhibitor is critical to a successful correction of the bleeding or to eradication of the antibody, most hemophilia centers perform annual screening for inhibitors. The laboratory test required to confirm the presence of an inhibitor is an aPTT with a mix (with normal plasma). In most hemophilia patients, a 1:1 mix with normal plasma completely corrects the aPTT. In inhibitor patients, the aPTT on a 1:1 mix is abnormally prolonged, because the inhibitor neutralizes the FVIII clotting activity of the normal plasma. The Bethesda assay uses a similar principle and defines the specificity of the inhibitor and its titer. The results are expressed in Bethesda units (BU), in which 1 BU is the amount of antibody that neutralizes 50% of the FVIII or FIX present in normal plasma after 2 h of incubation at 37°C. Clinically, inhibitor patients are classified as low responders or high responders, which provides guidelines for optimal therapy. Therapy for inhibitor patients has two goals: the control of acute bleeding episodes and the eradication of the inhibitor. For the control of bleeding episodes, low responders, those with titer <5 BU, respond well to high doses of human or porcine FVIII (50–100 U/kg), with minimal or no increase in the inhibitor titers. However, high-responder patients, those with initial inhibitor titer >10 BU or an anamnestic response in the antibody titer to >10 BU even if low titer initially, do not respond to FVIII or FIX concentrates. The control of bleeding episodes in high-responder patients can be achieved by using concentrates enriched for prothrombin, FVII, FIX, FX (prothrombin complex concentrates [PCCs] or activated PCCs [aPCCs]), and more recently recombinant activated factor VII (FVIIa) known as “bypass agents” (Fig. 141-1). The rates of therapeutic success have been higher for FVIIa than for PCC or aPCC. For eradication of the inhibitory antibody, immunosuppression alone is not effective. The most effective strategy is the immune tolerance induction (ITI) based on daily infusion of missing protein until the inhibitor disappears, typically requiring periods longer than 1 year, with success rates of approximately 60%. The management of patients with severe hemophilia and inhibitors resistant to ITI is challenging. The use of anti-CD20 monoclonal antibody (rituximab) combined with ITI was thought to be effective. Although this therapy may reduce the inhibitor titers in some cases, sustained eradication is uncommon and may require two to three infusions weekly of clotting factor concentrates.

Novel Therapeutic Approaches in Development for Hemophilia Clinical studies using long-acting clotting factors with prolonged half-lives are in the late phase of clinical testing, and these new-generation products (for FVIII and FIX) may facilitate prophylaxis by requiring fewer injections to maintain circulating levels above 1%.

The use of recombinant interleukin 11 in patients with moderate or mild hemophilia A unresponsive to DDAVP has been tested in early-phase clinical trials and may be an alternate therapeutic strategy for clinical situations that require transient increases in FVIII levels.

Gene therapy trials for hemophilia B using adeno-associated viral vectors are ongoing, and initial data are promising ([Chap. 91e](#)).

INFECTIOUS DISEASES

Hepatitis C virus (HCV) infection is the major cause of morbidity and the second leading cause of death in hemophilia patients exposed to older clotting factor concentrates. The vast majority of young patients treated with plasma-derived products from 1970 to 1985 became infected with HCV. It has been estimated that >80% of patients older than 20 years of age are HCV antibody positive as of 2006. The comorbidity of the underlying liver disease in hemophilia patients is clear when these individuals require invasive procedures; correction of both genetic and acquired (secondary to liver disease) deficiencies may be needed. Infection with HIV also swept the population of patients using plasma-derived concentrates two decades

ago. Co-infection of HCV and HIV, present in almost 50% of hemophilia patients, is an aggravating factor for the evolution of liver disease. The response to HCV antiviral therapy in hemophilia is restricted to <30% of patients and even poorer among those with both HCV and HIV infection. End-stage liver disease requiring organ transplantation may be curative for both the liver disease and for hemophilia.

EMERGING CLINICAL PROBLEMS IN AGING HEMOPHILIA PATIENTS

There has been continuous improvement of the management of hemophilia since the increase in the population of adults living beyond middle age in the developing world. The life expectancy of a patient with severe hemophilia is only ~10 years shorter than the general male population. In patients with mild or moderate hemophilia, life expectancy is approaching that of the male population without coagulopathy. Elderly hemophilia patients have different problems compared to the younger generation; they have more severe arthropathy and chronic pain, due to suboptimal treatment, and high rates of HCV and/or HIV infections.

Early data indicate that mortality from coronary artery disease is lower in hemophilia patients than the general male population. The underlying hypocoagulability probably provides a protective effect against thrombus formation, but it does not prevent atherogenesis. Similar to the general population, these patients are exposed to cardiovascular risk factors such as age, obesity, and smoking. Moreover, physical inactivity, hypertension, and chronic renal disease are commonly observed in hemophilia patients. In HIV patients on combined antiretroviral therapy, there may be a further increase in the risk of cardiovascular disease. Therefore, these patients should be carefully considered for preventive and therapeutic approaches to minimize the risk of cardiovascular disease.

Excessive replacement therapy should be avoided, and it is prudent to slowly infuse factor concentrates. Continuous infusion of clotting factor is preferable to bolus dosing in patients with cardiovascular risk factors undergoing invasive procedures. The management of an acute ischemic event and coronary revascularization should include the collaboration of hematologists and interventionalists. The early assumption that hemophilia would protect against occlusive vascular disease may change in this aging population. Cancer is a common cause of mortality in aging hemophilia patients because they are at risk for HIV- and HCV-related malignancies. Hepatocellular carcinoma (HCC) is the most prevalent primary liver cancer and a common cause of death in HIV-negative patients. The recommendations for cancer screening for the general population should be the same for age-matched hemophilia patients. Among those with high-risk HCV, a semiannual or annual ultrasound and a fetoprotein are recommended for HCC. Screening for urogenital neoplasm in the presence of hematuria or hematochezia may be delayed due to the underlying bleeding disease, thus preventing early intervention. Multidisciplinary interaction should facilitate the attempts to ensure optimal cancer prevention and treatment recommendations for those with hemophilia.

MANAGEMENT OF CARRIERS OF HEMOPHILIA

Usually hemophilia carriers, with factor levels of ~50% of normal, have not been considered to be at risk for bleeding. However, a wide range of values (22–116%) have been reported due to random inactivation of the X chromosomes (*Lyonization*). Therefore, it is important to measure the factor level of carriers to recognize those at risk of bleeding and to optimize preoperative and postoperative management. During pregnancy, both FVIII and FIX levels increase gradually until delivery. FVIII levels increase approximately two- to threefold compared to nonpregnant women, whereas an FIX increase is less pronounced. After delivery, there is a rapid fall in the pregnancy-induced rise of maternal clotting factor levels. This represents an imminent risk of bleeding that can be prevented by infusion of factor concentrate to levels of 50–70% for 3 days in the setting of vaginal delivery and up to 5 days for cesarean section. In mild cases, the use of DDAVP and/or antifibrinolytic drugs is recommended.

Factor XI is a zymogen of an active serine protease (FIXa) in the intrinsic pathway of blood coagulation that activates FIX (Fig. 141-1). There are two pathways for the formation of FIXa. In an aPTT-based assay, the protease is the result of activation by FXIIa in conjunction with high-molecular-weight kininogen and kallikrein. In vivo data suggest that thrombin is the physiologic activator of FIX. The generation of thrombin by the tissue factor/factor VIIa pathway activates FIX on the platelet surface that contributes to additional thrombin generation after the clot has formed and thus augments resistance to fibrinolysis through a thrombin-activated fibrinolytic inhibitor (TAFI).

Factor XI deficiency is a rare bleeding disorder that occurs in the general population at a frequency of one in a million. However, the disease is highly prevalent among Ashkenazi and Iraqi Jewish populations, reaching a frequency of 6% as heterozygotes and 0.1–0.3% as homozygotes. More than 65 mutations in the FXI gene have been reported, whereas fewer mutations (two to three) are found among affected Jewish populations.

Normal FXI clotting activity levels range from 70 to 150 U/dL. In heterozygous patients with moderate deficiency, FXI ranges from 20 to 70 U/dL, whereas in homozygous or double heterozygote patients, FXI levels are <1–20 U/dL. Patients with FXI levels <10% of normal have a high risk of bleeding, but the disease phenotype does not always correlate with residual FXI clotting activity. A family history is indicative of the risk of bleeding in the proband. Clinically, the presence of mucocutaneous hemorrhages such as bruises, gum bleeding, epistaxis, hematuria, and menorrhagia are common, especially following trauma. This hemorrhagic phenotype suggests that tissues rich in fibrinolytic activity are more susceptible to FXI deficiency. Postoperative bleeding is common but not always present, even among patients with very low FXI levels.

FXI replacement is indicated in patients with severe disease required to undergo a surgical procedure. A negative history of bleeding complications following invasive procedures does not exclude the possibility of an increased risk for hemorrhage.

TREATMENT FACTOR XI DEFICIENCY

The treatment of FXI deficiency is based on the infusion of FFP at doses of 15–20 mL/kg to maintain trough levels ranging from 10 to 20%. Because FXI has a half-life of 40–70 h, the replacement therapy can be given on alternate days. The use of antifibrinolytic drugs is beneficial to control bleeds, with the exception of hematuria or bleeds in the bladder. The development of an FXI inhibitor was observed in 10% of severely FXI-deficient patients who received replacement therapy. Patients with severe FXI deficiency who develop inhibitors usually do not bleed spontaneously. However, bleeding following a surgical procedure or trauma can be severe. In these patients, FFP and FXI concentrates should be avoided. The use of PCC/aPCC or recombinant activated FVII has been effective.

RARE BLEEDING DISORDERS

Collectively, the inherited disorders resulting from deficiencies of clotting factors other than FVIII, FIX, and FXI (Table 141-1) represent a group of rare bleeding diseases. The bleeding symptoms in these patients vary from asymptomatic (dysfibrinogenemia or FVII deficiency) to life-threatening (FX or FXIII deficiency). There is no pathognomonic clinical manifestation that suggests one specific disease, but overall, in contrast to hemophilia, hemarthrosis is a rare event and bleeding in the mucosal tract or after umbilical cord clamping is common. Individuals heterozygous for plasma coagulation deficiencies are often asymptomatic. The laboratory assessment for the specific deficient factor following screening with general coagulation tests (Table 141-1) will define the diagnosis.

Replacement therapy using FFP or prothrombin complex concentrates (containing prothrombin, FVII, FIX, and FX) provides adequate hemostasis in response to bleeds or as prophylactic treatment. The use

of PCC should be carefully monitored and avoided in patients with underlying liver disease, or those at high risk for thrombosis because of the risk of DIC.

FAMILIAL MULTIPLE COAGULATION DEFICIENCIES

There are several bleeding disorders characterized by the inherited deficiency of more than one plasma coagulation factor. To date, the genetic defects in two of these diseases have been characterized, and they provide new insights into the regulation of hemostasis by gene-encoding proteins outside blood coagulation.

Combined Deficiency of FV and FVIII Patients with combined FV and FVIII deficiency exhibit ~5% of residual clotting activity of each factor. Interestingly, the disease phenotype is a mild bleeding tendency, often following trauma. An underlying mutation has been identified in the endoplasmic reticulum/Golgi intermediate compartment (*ERGIC-53*) gene, a mannose-binding protein localized in the Golgi apparatus that functions as a chaperone for both FV and FVIII. In other families, mutations in the multiple coagulation factor deficiency 2 (*MCFD2*) gene have been defined; this gene encodes a protein that forms a Ca^{2+} -dependent complex with *ERGIC-53* and provides cofactor activity in the intracellular mobilization of both FV and FVIII.

Multiple Deficiencies of Vitamin K-Dependent Coagulation Factors Two enzymes involved in vitamin K metabolism have been associated with combined deficiency of all vitamin K-dependent proteins, including the procoagulant proteins prothrombin, VII, IX, and X and the anticoagulant proteins C and S. Vitamin K is a fat-soluble vitamin that is a cofactor for carboxylation of the gamma carbon of the glutamic acid residues in the vitamin K-dependent factors, a critical step for calcium and phospholipid binding of these proteins (Fig. 141-2). The enzymes γ -glutamylcarboxylase and epoxide reductase are critical for the metabolism and regeneration of vitamin K. Mutations in the genes encoding the γ -carboxylase (GGCX) or vitamin K epoxide reductase complex 1 (VKORC1) result in defective enzymes and thus in vitamin K-dependent factors with reduced activity, varying from 1 to 30% of normal. The disease phenotype is characterized by mild to severe bleeding episodes present from birth. Some patients respond to high doses of vitamin K. For severe bleeding, replacement therapy with FFP or PCC may be necessary to achieve full hemostatic control.

DISSEMINATED INTRAVASCULAR COAGULATION

DIC is a clinicopathologic syndrome characterized by widespread intravascular fibrin formation in response to excessive blood protease

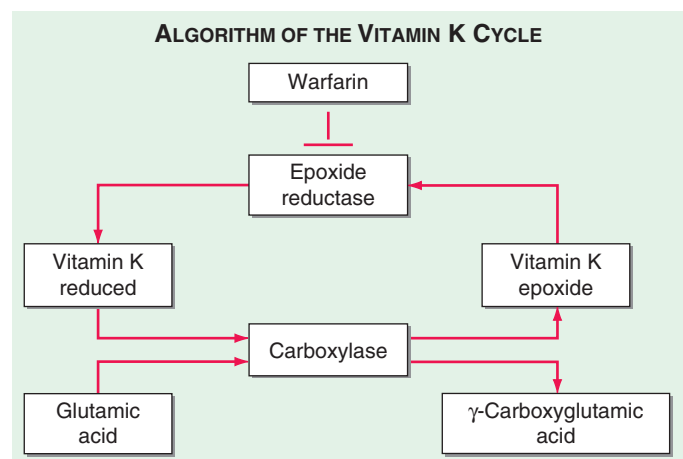


FIGURE 141-2 The vitamin K cycle. Vitamin K is a cofactor for the formation of γ -carboxyglutamic acid residues on coagulation proteins. Vitamin K-dependent γ -glutamylcarboxylase, the enzyme that catalyzes the vitamin K epoxide reductase, regenerates reduced vitamin K. Warfarin blocks the action of the reductase and competitively inhibits the effects of vitamin K.

TABLE 141-2 COMMON CLINICAL CAUSES OF DISSEMINATED INTRAVASCULAR COAGULATION

Sepsis	Immunologic Disorders
<ul style="list-style-type: none"> Bacterial: Staphylococci, streptococci, pneumococci, meningococci, gram-negative bacilli Viral Mycotic Parasitic Rickettsial 	<ul style="list-style-type: none"> Acute hemolytic transfusion reaction Organ or tissue transplant rejection Immunotherapy Graft-versus-host disease
Trauma and Tissue Injury	Drugs
<ul style="list-style-type: none"> Brain injury (gunshot) Extensive burns Fat embolism Rhabdomyolysis 	<ul style="list-style-type: none"> Fibrinolytic agents Aprotinin Warfarin (especially in neonates with protein C deficiency) Prothrombin complex concentrates Recreational drugs (amphetamines)
Vascular Disorders	Envenomation
<ul style="list-style-type: none"> Giant hemangiomas (Kasabach-Merritt syndrome) Large vessel aneurysms (e.g., aorta) 	<ul style="list-style-type: none"> Snake Insects
Obstetrical Complications	Liver Disease
<ul style="list-style-type: none"> Abruptio placentae Amniotic fluid embolism Dead fetus syndrome Septic abortion 	<ul style="list-style-type: none"> Fulminant hepatic failure Cirrhosis Fatty liver of pregnancy
Cancer	Miscellaneous
<ul style="list-style-type: none"> Adenocarcinoma (prostate, pancreas, etc.) Hematologic malignancies (acute promyelocytic leukemia) 	<ul style="list-style-type: none"> Shock Respiratory distress syndrome Massive transfusion

activity that overcomes the natural anticoagulant mechanisms. There are several underlying pathologies associated with DIC (Table 141-2).

The most common causes are bacterial sepsis, malignant disorders such as solid tumors or acute promyelocytic leukemia, and obstetric causes. DIC is diagnosed in almost one-half of pregnant women with abruptio placentae or with amniotic fluid embolism. Trauma, particularly to the brain, can also result in DIC. The exposure of blood to phospholipids from damaged tissue, hemolysis, and endothelial damage are all contributing factors to the development of DIC in this setting. Purpura fulminans is a severe form of DIC resulting from thrombosis of extensive areas of the skin; it affects predominantly young children following viral or bacterial infection, particularly those with inherited or acquired hypercoagulability due to deficiencies of the components of the protein C pathway. Neonates homozygous for protein C deficiency also present high risk for purpura fulminans with or without thrombosis of large vessels.

The central mechanism of DIC is the uncontrolled generation of thrombin by exposure of the blood to pathologic levels of tissue factor (Fig. 141-3). Simultaneous suppression of physiologic anticoagulant mechanisms and abnormal fibrinolysis further accelerate the process. Together, these abnormalities contribute to systemic fibrin deposition in small and midsize vessels. The duration and intensity of the fibrin deposition can compromise the blood supply of many organs, especially the lung, kidney, liver, and brain, with consequent organ failure. The sustained activation of coagulation results in consumption of clotting factors and platelets, which in turn leads to

systemic bleeding. This is further aggravated by secondary hyperfibrinolysis. Studies in animals demonstrate that the fibrinolytic system is indeed suppressed at the time of maximal activation of coagulation. Interestingly, in patients with acute promyelocytic leukemia, a severe hyperfibrinolytic state often occurs in addition to the coagulation activation. The release of several proinflammatory cytokines such as interleukin 6 and tumor necrosis factor α plays a central role in mediating the coagulation defects in DIC and symptoms associated with systemic inflammatory response syndrome (SIRS).

Clinical manifestations of DIC are related to the magnitude of the imbalance of hemostasis, to the underlying disease, or to both. The most common findings are bleeding ranging from oozing from venipuncture sites, petechiae, and ecchymoses to severe hemorrhage from the gastrointestinal tract, lung, or into the CNS. In chronic DIC, the bleeding symptoms are discrete and restricted to skin or mucosal surfaces. The hypercoagulability of DIC manifests as the occlusion of vessels in the microcirculation and resulting organ failure. Thrombosis of large vessels and cerebral embolism can also occur. Hemodynamic complications and shock are common among patients with acute DIC. The mortality ranges from 30 to >80% depending on the underlying disease, the severity of the DIC, and the age of the patient.

The diagnosis of clinically significant DIC is based on the presence of clinical and/or laboratory abnormalities of coagulation or thrombocytopenia. The laboratory diagnosis of DIC should prompt a search for the underlying disease if it is not already apparent. There is no single test that establishes the diagnosis of DIC. The laboratory investigation should include coagulation tests (aPTT, PT, thrombin time [TT]) and markers of fibrin degradation products (FDPs), in addition to platelet and red cell count and analysis of the blood smear. These tests should be repeated over a period of 6–8 h because an initially mild abnormality can change dramatically in patients with severe DIC.

Common findings include the prolongation of PT and/or aPTT; platelet counts $\mu 100,000/\mu\text{L}$, or a rapid decline in platelet numbers; the presence of schistocytes (fragmented red cells) in the blood smear; and elevated levels of FDP. The most sensitive test for DIC is the FDP level. DIC is an unlikely diagnosis in the presence of normal levels of FDP. The D-dimer test is more specific for detection of fibrin—but not fibrinogen—degradation products and indicates that the cross-linked fibrin has been digested by plasmin. Because fibrinogen has a prolonged half-life, plasma levels diminish acutely only in severe cases of DIC. High-grade DIC is also associated with levels of antithrombin III or plasminogen activity <60% of normal.

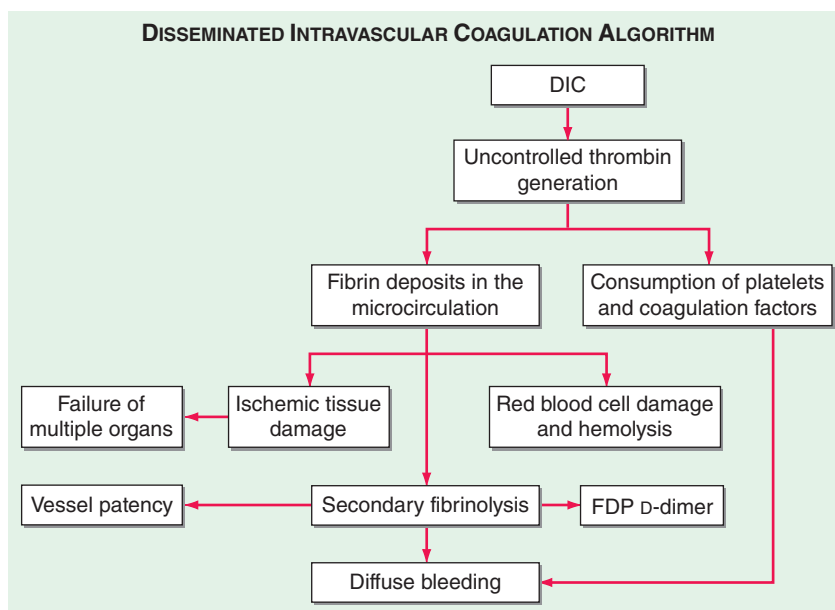


FIGURE 141-3 The pathophysiology of disseminated intravascular coagulation (DIC). Interactions between coagulation and fibrinolytic pathways result in bleeding and thrombosis in the microcirculation in patients with DIC. FDP, fibrin degradation product.

738 Chronic DIC Low-grade, compensated DIC can occur in clinical situations including giant hemangioma, metastatic carcinoma, or the dead fetus syndrome. Plasma levels of FDP or D-dimers are elevated. aPTT, PT, and fibrinogen values are within the normal range or high. Mild thrombocytopenia or normal platelet counts are also common findings. Red cell fragmentation is often detected but at a lower degree than in acute DIC.

Differential Diagnosis The differential diagnosis between DIC and severe liver disease is challenging and requires serial measurements of the laboratory parameters of DIC. Patients with severe liver disease are at risk for bleeding and manifest laboratory features including thrombocytopenia (due to platelet sequestration, portal hypertension, or hypersplenism), decreased synthesis of coagulation factors and natural anticoagulants, and elevated levels of FDP due to reduced hepatic clearance. However, in contrast to DIC, these laboratory parameters in liver disease do not change rapidly. Other important differential findings include the presence of portal hypertension or other clinical or laboratory evidence of an underlying liver disease.

Microangiopathic disorders such as thrombotic thrombocytopenic purpura present an acute clinical onset of illness accompanied by thrombocytopenia, red cell fragmentation, and multiorgan failure. However, there is no consumption of clotting factors or hyperfibrinolysis.

Over the last few years, several clinical trials on immune therapies for neoplasias using monoclonal antibodies or gene-modified T cells targeting tumor-specific antigens showed unwanted inflammatory responses with increased cytokine release. These complications are sometimes associated with increased D-dimers and decreased fibrinogen levels, cytopenias, and liver dysfunction; thus, careful screening tests for DIC are indicated.

TREATMENT DISSEMINATED INTRAVASCULAR COAGULATION

The morbidity and mortality associated with DIC are primarily related to the underlying disease rather than the complications of the DIC. The control or elimination of the underlying cause should therefore be the primary concern. Patients with severe DIC require control of hemodynamic parameters, respiratory support, and sometimes invasive surgical procedures. Attempts to treat DIC without accompanying treatment of the causative disease are likely to fail.

MANAGEMENT OF HEMORRHAGIC SYMPTOMS

Administration of FFP and/or platelet concentrates is indicated for patients with active bleeding or at high risk of bleeding, such as in preparation for invasive procedures or after chemotherapy. The control of bleeding in DIC patients with marked thrombocytopenia (platelet counts <10,000–20,000/ μ L) and low levels of coagulation factors will require replacement therapy. The PT (>1.5 times the normal) provides a good indicator of the severity of the clotting factor consumption. Replacement with FFP is indicated (1 unit of FFP increases most coagulation factors by 30% in an adult without DIC). Low levels of fibrinogen (<100 mg/dL) or brisk hyperfibrinolysis will require infusion of cryoprecipitate (plasma fraction enriched for fibrinogen, FVIII, and VWF). The replacement of 10 U of cryoprecipitate for every 2–3 U of FFP is sufficient to correct the hemostasis. The transfusion scheme must be adjusted according to the patient's clinical and laboratory evolution. Platelet concentrates at a dose of 1–2 U/10 kg body weight are sufficient for most DIC patients with severe thrombocytopenia. Clotting factor concentrates are not recommended for control of bleeding in DIC because of the limited efficacy afforded by replacement of single factors (FVIII or FIX concentrates) and the high risk of products containing traces of aPCCs that further aggravate the disease.

REPLACEMENT OF COAGULATION OR FIBRINOLYSIS INHIBITORS

Drugs to control coagulation such as heparin, antithrombin III (ATIII) concentrates, or antifibrinolytic drugs have all been tried in the

treatment of DIC. Low doses of continuous-infusion heparin (5–10 U/kg per h) may be effective in patients with low-grade DIC associated with solid tumor, acute promyelocytic leukemia, or in a setting with recognized thrombosis. Heparin is also indicated for the treatment of purpura fulminans during the surgical resection of giant hemangiomas and during removal of a dead fetus. In acute DIC, the use of heparin is likely to aggravate bleeding. To date, the use of heparin in patients with severe DIC has no proven survival benefit. The use of antifibrinolytic drugs, EACA, or tranexamic acid to prevent fibrin degradation by plasmin may reduce bleeding episodes in patients with DIC and confirmed hyperfibrinolysis. However, these drugs can increase the risk of thrombosis, and concomitant use of heparin is indicated. Patients with acute promyelocytic leukemia or those with chronic DIC associated with giant hemangiomas are among the few patients who may benefit from this therapy. The use of protein C concentrates to treat purpura fulminans associated with acquired protein C deficiency or meningococemia has been proven efficacious. The results from the replacement of ATIII in early-phase studies are promising but require further study.

Guidance for diagnosis and treatment of DIC had been proposed by the International Society of Thrombosis and Haemostasis. This initiative will permit more detailed clinical data on diagnosis and treatment of DIC. The clinical utility of these scoring systems and therapeutic recommendations contained in these guidelines is not yet known.

VITAMIN K DEFICIENCY Vitamin K–dependent proteins are a heterogeneous group, including clotting factor proteins and also proteins found in bone, lung, kidney, and placenta. Vitamin K mediates posttranslational modification of glutamate residues to γ -carboxylglutamate, a critical step for the activity of vitamin K–dependent proteins for calcium binding and proper assembly to phospholipid membranes (Fig. 141-2). Inherited deficiency of the functional activity of the enzymes involved in vitamin K metabolism, notably the GGCX or VKORC1 (see above), results in bleeding disorders. The amount of vitamin K in the diet is often limiting for the carboxylation reaction; thus recycling of the vitamin K is essential to maintain normal levels of vitamin K–dependent proteins. In adults, low dietary intake alone is seldom reason for severe vitamin K deficiency but may become common in association with the use of broad-spectrum antibiotics. Disease or surgical interventions that affect the ability of the intestinal tract to absorb vitamin K, either through anatomic alterations or by changing the fat content of bile salts and pancreatic juices in the proximal small bowel, can result in significant reduction of vitamin K levels. Chronic liver diseases such as primary biliary cirrhosis also deplete vitamin K stores. Neonatal vitamin K deficiency and the resulting hemorrhagic disease of the newborn have been almost entirely eliminated by routine administration of vitamin K to all neonates. Prolongation of PT values is the most common and earliest finding in vitamin K–deficient patients due to reduction in prothrombin, FVII, FIX, and FX levels. FVII has the shortest half-life among these factors that can prolong the PT before changes in the aPTT. Parenteral administration of vitamin K at a total dose of 10 mg is sufficient to restore normal levels of clotting factor within 8–10 h. In the presence of ongoing bleeding or a need for immediate correction before an invasive procedure, replacement with FFP or PCC is required. The latter should be avoided in patients with severe underlying liver disorders due to high risk of thrombosis. The reversal of excessive anticoagulant therapy with warfarin or warfarin-like drugs can be achieved by minimal doses of vitamin K (1 mg orally or by intravenous injection) for asymptomatic patients. This strategy can diminish the risk of bleeding while maintaining therapeutic anticoagulation for an underlying prothrombotic state.

In patients with life-threatening bleeds, the use of recombinant factor VIIa in nonhemophilia patients on anticoagulant therapy has been shown to be effective at restoring hemostasis rapidly, allowing emergency surgical intervention. However, patients with underlying vascular disease, vascular trauma and other comorbidities are at risk for thromboembolic complications that affect both arterial and venous

TABLE 141-3 COAGULATION DISORDERS AND HEMOSTASIS IN LIVER DISEASE

Bleeding	
Portal hypertension	
Esophageal varices	
Thrombocytopenia	
Splenomegaly	
Chronic or acute DIC	
Decreased synthesis of clotting factors	
Hepatocyte failure	
Vitamin K deficiency	
Systemic fibrinolysis	
DIC	
Dysfibrinogenemia	
Thrombosis	
Decreased synthesis of coagulation inhibitors: protein C, protein S, anti-thrombin	
Hepatocyte failure	
Vitamin K deficiency (protein C, protein S)	
Failure to clear activated coagulation proteins (DIC)	
Dysfibrinogenemia	
iatrogenic: Transfusion of prothrombin complex concentrates	
Antifibrinolytic agents: EACA, tranexamic acid	

Abbreviations: DIC, disseminated intravascular coagulation; EACA, ϵ -aminocaproic acid.

systems. Thus, the use of factor VIIa in this setting is limited to administration of low doses given for only a limited number of injections. Close monitoring for vascular complications is highly indicated.

COAGULATION DISORDERS ASSOCIATED WITH LIVER FAILURE The liver is central to hemostasis because it is the site of synthesis and clearance of most procoagulant and natural anticoagulant proteins and of essential components of the fibrinolytic system. Liver failure is associated with a high risk of bleeding due to deficient synthesis of procoagulant factors and enhanced fibrinolysis. Thrombocytopenia is common in patients with liver disease, and may be due to congestive splenomegaly (hypersplenism) or immune-mediated shortened platelet lifespan (primary biliary cirrhosis). In addition, several anatomic abnormalities secondary to underlying liver disease further promote the occurrence of hemorrhage (Table 141-3). Dysfibrinogenemia is a relatively common finding in patients with liver disease due to impaired fibrin polymerization. The development of DIC concomitant to chronic liver disease is not uncommon and may enhance the risk for bleeding. Laboratory evaluation is mandatory for an optimal therapeutic strategy, either to control ongoing bleeding or to prepare patients with liver disease for invasive procedures. Typically, these patients present with prolonged PT, aPTT, and TT depending on the degree of liver damage, thrombocytopenia, and normal or slight increase of FDP. Fibrinogen levels are diminished only in fulminant hepatitis, decompensated cirrhosis, or advanced liver disease, or in the presence of DIC. The presence of prolonged TT and normal fibrinogen and FDP levels suggest dysfibrinogenemia. FVIII levels are often normal or elevated in patients with liver failure, and decreased levels suggest superimposing DIC. Because FV is only synthesized in the hepatocyte and

is not a vitamin K–dependent protein, reduced levels of FV may be an indicator of hepatocyte failure. Normal levels of FV and low levels of FVII suggest vitamin K deficiency. Vitamin K levels may be reduced in patients with liver failure due to compromised storage in hepatocellular disease, changes in bile acids, or cholestasis that can diminish the absorption of vitamin K. Replacement of vitamin K may be desirable (10 mg given by slow intravenous injection) to improve hemostasis.

Treatment with FFP is the most effective to correct hemostasis in patients with liver failure. Infusion of FFP (5–10 mL/kg; each bag contains ~200 mL) is sufficient to ensure 10–20% of normal levels of clotting factors but not correction of PT or aPTT. Even high doses of FFP (20 mL/kg) do not correct the clotting times in all patients. Monitoring for clinical symptoms and clotting times will determine if repeated doses are required 8–12 h after the first infusion. Platelet concentrates are indicated when platelet counts are <10,000–20,000/ μ L to control an ongoing bleed or immediately before an invasive procedure if counts are <50,000/ μ L. Cryoprecipitate is indicated only when fibrinogen levels are less than 100 mg/mL; dosing is six bags for a 70-kg patient daily. Prothrombin complex concentrate infusion in patients with liver failure should be avoided due to the high risk of thrombotic complications. The safety of the use of antifibrinolytic drugs to control bleeding in patients with liver failure is not yet well defined and should be avoided.

LIVER DISEASE AND THROMBOEMBOLISM The clinical bleeding phenotype of hemostasis in patients with stable liver disease is often mild or even asymptomatic. However, as the disease progresses, the hemostatic balance is less stable and more easily disturbed than in healthy individuals. Furthermore, the hemostatic balance is compromised by comorbid complications such as infections and renal failure (Fig. 141-4). Based on the clinical bleeding complications in patients with cirrhosis and laboratory evidence of hypocoagulation such as a prolonged PT/aPTT, it has long been assumed that these patients are protected against thrombotic disease. Cumulative clinical experience, however, has demonstrated that these patients are at risk for thrombosis, especially those with advanced liver disease. Although hypercoagulability could explain the occurrence of venous thrombosis, according to Virchow's

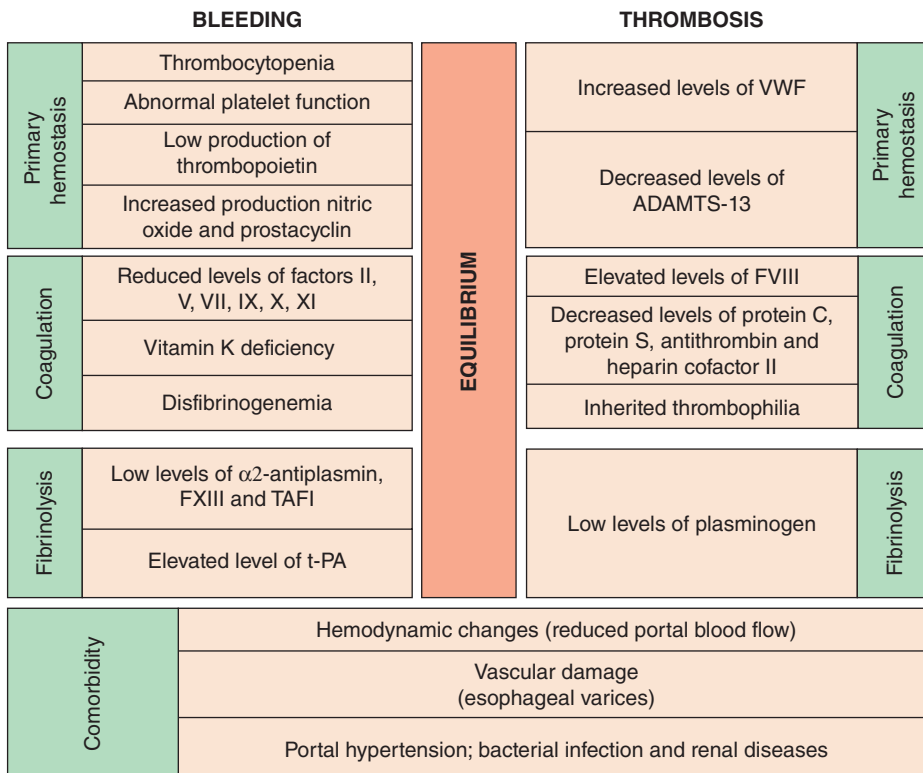


FIGURE 141-4 Balance of hemostasis in liver disease. TAFI, thrombin-activated fibrinolytic inhibitor; t-PA, tissue plasminogen activator; VWF, von Willebrand factor.

740 triad, hemodynamic changes and damaged vasculature may also be a contributing factor, and both processes may potentially also occur in patients with liver disease. Liver-related thrombosis, in particular, thrombosis of the portal and mesenteric veins, is common in patients with advanced cirrhosis. Hemodynamic changes, such as decreased portal flow, and evidence that inherited thrombophilia may enhance the risk for portal vein thrombosis in patients with cirrhosis suggest that hypercoagulability may play a role as well. Patients with liver disease develop deep vein thrombosis and pulmonary embolism at appreciable rates (ranging from 0.5 to 1.9%). The implication of these findings is relevant to the erroneous exclusion of thrombosis in patients with advanced liver disease, even in the presence of prolongation of routine clotting times, and caution should be advised on overcorrection of these laboratory abnormalities.

ACQUIRED INHIBITORS OF COAGULATION FACTORS An acquired inhibitor is an immune-mediated disease characterized by the presence of an autoantibody against a specific clotting factor. FVIII is the most common target of antibody formation, and is sometimes referred to as acquired hemophilia A, but inhibitors to prothrombin, FV, FIX, FX, and FXI are also reported. Acquired inhibitor to FVIII occurs predominantly in older adults (median age of 60 years), but occasionally in pregnant or postpartum women with no previous history of bleeding. In 50% of patients with inhibitors, no underlying disease is identified at the time of diagnosis. In the remaining patients, the causes are autoimmune diseases, malignancies (lymphomas, prostate cancer), dermatologic diseases, and pregnancy. Bleeding episodes occur commonly in soft tissues, the gastrointestinal or urinary tracts, and skin. In contrast to hemophilia, hemarthrosis is rare in these patients. Retroperitoneal hemorrhages and other life-threatening bleeding may appear suddenly. The overall mortality in untreated patients ranges from 8 to 22%, and most deaths occur within the first few weeks after presentation. The diagnosis is based on the prolonged aPTT with normal PT and TT. The aPTT remains prolonged after mixture of the test plasma with equal amounts of pooled normal plasma for 2 h at 37°C. The Bethesda assay using FVIII-deficient plasma as performed for inhibitor detection in hemophilia will confirm the diagnosis. Major bleeding is treated with bypass products such as PCC/aPCC or recombinant FVIIa. In contrast to hemophilia, inhibitors in nonhemophilic patients are typically responsive to immune suppression, and therapy should be initiated early for most cases. The first choice includes steroid or a combination of steroid with cytotoxic therapy (e.g., cyclophosphamide), with complete eradication of the inhibitors in more than 70% of patients. High-dose intravenous γ -globulin and anti-CD20 monoclonal antibody have been reported to be effective in patients with autoantibodies to FVIII; however, there is no firm evidence that these alternatives are superior to the first line of immunosuppressive drugs. Notably, relapse of the inhibitor to FVIII is relatively common (up to 20%) within the first 6 months following withdrawal of immunosuppression. Thus, after eradication, patients should be followed up regularly for early therapeutic intervention when indicated or prior to invasive procedure.

Topical plasma-derived bovine and human thrombin are commonly used in the United States and worldwide. These effective hemostatic sealants are used during major surgery such as for cardiovascular, thoracic, neurologic, pelvic, and trauma indications, as well as in the setting of extensive burns. The development of antibody formation to the xenoantigen or its contaminant (bovine clotting protein) has the potential to show cross-reactivity with human clotting factors that may hamper their function and induce bleeding.

Clinical features of these antibodies include bleeding from a primary hemostatic defect or coagulopathy that sometimes can be life threatening. The clinical diagnosis of these acquired coagulopathies is often complicated by the fact that the bleeding episodes may be detectable during or immediately following major surgery and could be assumed to be due to the procedure itself.

Notably, the risk of this complication is further increased by repeated exposure to topical thrombin preparations. Thus, a careful medical history of previous surgical interventions that may have occurred even decades earlier is critical to assessing risk.

The laboratory abnormalities are reflected by combined prolongation of the aPTT and PT that often fails to improve by transfusion of FFP and vitamin K. The abnormal laboratory tests cannot be corrected by mixing a test with equal parts of normal plasma that denotes the presence of inhibitory antibodies. The diagnosis of a specific antibody is obtained by the determination of the residual activity of human FV or other suspected human clotting factor. There are no commercially available assays specific for bovine thrombin coagulopathy.

There are no established treatment guidelines. Platelet transfusions have been used as a source of FV replacement for patients with FV inhibitors. Frequent injections of FFP and vitamin K supplementation may function as co-adjuvant rather than an effective treatment of the coagulopathy itself. Experience with recombinant FVIIa as a bypass agent is limited, and outcomes have been generally poor. Specific treatments to eradicate the antibodies based on immunosuppression with steroids, intravenous immunoglobulin, or serial plasmapheresis have been sporadically reported. Patients should be advised to avoid any topical thrombin sealant in the future.

Novel plasma-derived and recombinant human thrombin preparations for topical hemostasis have been approved by the U.S. Food and Drug Administration. These preparations have demonstrated hemostatic efficacy with reduced immunogenicity compared to the first generation of bovine thrombin products.

The presence of lupus anticoagulant can be associated with venous or arterial thrombotic disease. However, bleeding has also been reported in lupus anticoagulant; it is due to the presence of antibodies to prothrombin, which results in hypoprothrombinemia. Both disorders show a prolonged PTT that does not correct on mixing. To distinguish acquired inhibitors from lupus anticoagulant, note that the dilute Russell's viper venom test and the hexagonal-phase phospholipids test will be negative in patients with an acquired inhibitor and positive in patients with lupus anticoagulants. Moreover, lupus anticoagulant interferes with the clotting activity of many factors (FVIII, FIX, FXII, FXI), whereas acquired inhibitors are specific to a single factor.

142

Arterial and Venous Thrombosis

Jane E. Freedman, Joseph Loscalzo

OVERVIEW OF THROMBOSIS

GENERAL OVERVIEW

Thrombosis, the obstruction of blood flow due to the formation of clot, may result in tissue anoxia and damage, and it is a major cause of morbidity and mortality in a wide range of arterial and venous diseases and patient populations. In 2009 in the United States, an estimated 785,000 people had a new coronary thrombotic event, and about 470,000 had a recurrent ischemic episode. Each year, approximately 795,000 people have a new or recurrent stroke. It is estimated that 300,000–600,000 people each year have a pulmonary embolism or deep venous thrombotic event. In the nondiseased state, physiologic hemostasis reflects a delicate interplay between factors that promote and inhibit blood clotting, favoring the former. This response is crucial as it prevents uncontrolled hemorrhage and exsanguination following injury. In specific settings, the same processes that regulate normal hemostasis can cause pathologic thrombosis, leading to arterial or venous occlusion. Importantly, many commonly used therapeutic interventions may also alter the thrombotic–hemostatic balance adversely.

Hemostasis and thrombosis primarily involve the interplay among three factors: the vessel wall, coagulation proteins, and platelets. Many prevalent acute vascular diseases are due to thrombus formation within a vessel, including myocardial infarction, thrombotic cerebrovascular events, and venous thrombosis. Although the end result is vessel occlusion and tissue ischemia, the pathophysiologic processes

governing these pathologies have similarities as well as distinct differences. While many of the pathways regulating thrombus formation are similar to those that regulate hemostasis, the processes triggering thrombosis and, often, perpetuating the thrombus may be distinct and can vary in different clinical and genetic settings. In venous thrombosis, primary hypercoagulable states reflecting defects in the proteins governing coagulation and/or fibrinolysis or secondary hypercoagulable states involving abnormalities of blood vessels and blood flow or stasis lead to thrombosis. By contrast, arterial thrombosis is highly dependent on the state of the vessel wall, the platelet, and factors related to blood flow.

ARTERIAL THROMBOSIS

OVERVIEW OF ARTERIAL THROMBOSIS

In arterial thrombosis, the platelets and abnormalities of the vessel wall typically play a key role in vessel occlusion. Arterial thrombus forms via a series of sequential steps in which platelets adhere to the vessel wall, additional platelets are recruited, and thrombin is activated (Fig. 142-1). The regulation of platelet adhesion, activation, aggregation, and recruitment will be described in detail below. In addition, while the primary function of platelets is regulation of hemostasis, our understanding of their role in other processes, such as immunity, wound healing, and inflammation, continues to grow.

ARTERIAL THROMBOSIS AND VASCULAR DISEASE

Arterial thrombosis is a major cause of morbidity and mortality both in the United States and, increasingly, worldwide. Although the rates have declined in the United States, the overall burden remains high and accounts for approximately 33% of deaths. Overall, coronary heart disease is estimated to cause about 1 of every 5 deaths in the United States. In addition to the 785,000 Americans who will have a new coronary event, an additional 195,000 silent first myocardial infarctions are

projected to occur annually. Although the rate of strokes has fallen by a third, each year, about 795,000 people experience a new or recurrent stroke, although not all are caused by thrombotic occlusion of the vessel. Approximately 610,000 strokes are first events and 185,000 are recurrent events; it is estimated that 1 of every 18 deaths in the United States is due to stroke.

THE PLATELET

Many processes in platelets have parallels with other cell types, such as the presence of specific receptors and signaling pathways; however, unlike most cells, platelets lack a nucleus and are unable to adapt to changing biologic settings by altered gene transcription. Platelets sustain limited protein synthetic capacity from megakaryocyte-derived and intracellularly transported microRNA (miRNA) and messenger RNA (mRNA). Most of the molecules needed to respond to various stimuli, however, are maintained in storage granules and membrane compartments.

Platelets are disc-shaped, very small, anucleate cells (1–5 μm in diameter) that circulate in the blood at concentrations of 200–400,000/ μL , with an average lifespan of 7–10 days. Platelets are derived from megakaryocytes, polyploid hematopoietic cells found in the bone marrow. The primary regulator of platelet formation is thrombopoietin (TPO). The precise mechanism by which megakaryocytes produce and release fully formed platelets is unclear, but the process likely involves formation of proplatelets, pseudopod-like structures generated by the evagination of the cytoplasm from which platelets bud. Platelet granules are synthesized in megakaryocytes before thrombopoiesis and contain an array of prothrombotic, proinflammatory, and antimicrobial mediators. The two major types of platelet granules, alpha and dense, are distinguished by their size, abundance, and content. Alpha-granules contain soluble coagulation proteins, adhesion molecules, growth factors, integrins, cytokines, and inflammatory modulators. Platelet dense-granules are smaller than alpha-granules and less abundant. Whereas alpha-granules contain proteins that may be more important in the inflammatory response, dense-granules contain high concentrations of small molecules, including adenosine diphosphate (ADP) and serotonin, that influence platelet aggregation.

Platelet Adhesion (See Fig. 142-1) The formation of a thrombus is initiated by the adherence of platelets to the damaged vessel wall. Damage exposes subendothelial components responsible for triggering platelet reactivity, including collagen, von Willebrand factor, fibronectin, and other adhesive proteins, such as vitronectin and thrombospondin. The hemostatic response may vary, depending on the extent of damage, the specific proteins exposed, and flow conditions. Certain proteins are expressed on the platelet surface that subsequently regulate collagen-induced platelet adhesion, particularly under flow conditions, and include glycoprotein (GP) IV, GPVI, and the integrin $\alpha_2\beta_1$. The platelet GPIb-IX-V complex adhesive receptor is central both to platelet adhesion and to the initiation of platelet activation. Damage to the blood vessel wall exposes subendothelial von Willebrand factor and collagen to the circulating blood. The GPIb-IX-V complex binds to the exposed von Willebrand factor, causing platelets to adhere (Fig. 142-1). In addition, the engagement of the GPIb-IX-V complex with ligand induces

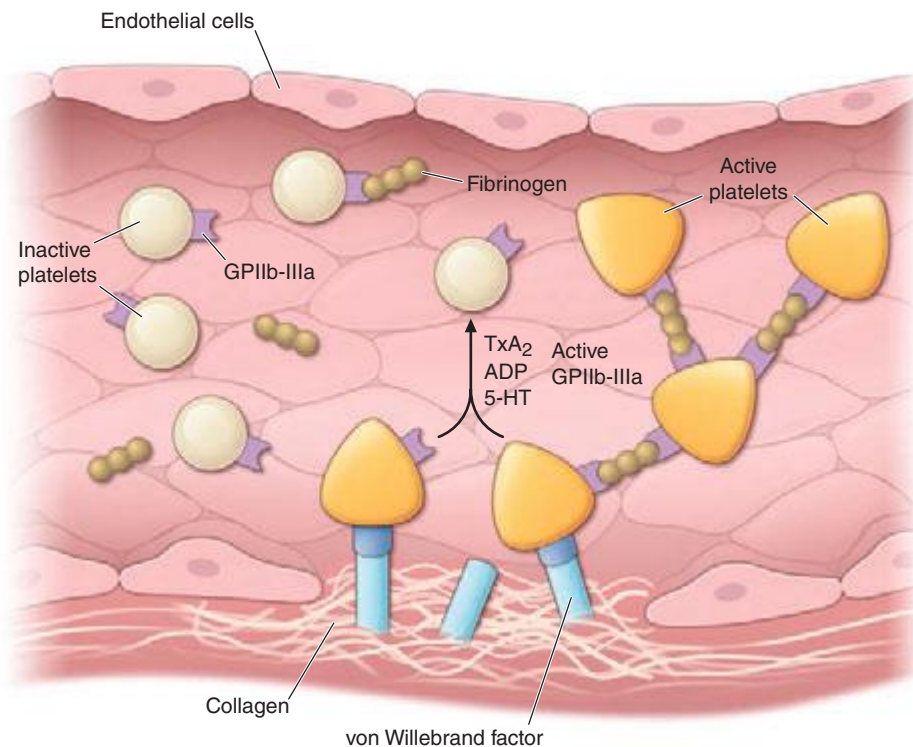


FIGURE 142-1 Platelet activation and thrombosis. Platelets circulate in an inactive form in the vasculature. Damage to the endothelium and/or external stimuli activates platelets that adhere to the exposed subendothelial von Willebrand factor and collagen. This adhesion leads to activation of the platelet, shape change, and the synthesis and release of thromboxane (TxA_2), serotonin (5-HT), and adenosine diphosphate (ADP). Platelet stimuli cause conformational change in the platelet integrin glycoprotein (GP) IIb/IIIa receptor, leading to the high-affinity binding of fibrinogen and the formation of a stable platelet thrombus.

742 signaling pathways that lead to platelet activation. von Willebrand factor-bound GPIb-IX-V promotes a calcium-dependent conformational change in the GPIIb/IIIa receptor, transforming it from an inactive low-affinity state to an active high-affinity receptor for fibrinogen.

Platelet Activation The activation of platelets is controlled by a variety of surface receptors that regulate various functions in the activation process. Platelet receptors control many distinct processes and are stimulated by a wide variety of agonists and adhesive proteins that result in variable degrees of activation. In general terms, the stimulation of platelet receptors triggers two specific processes: (1) activation of internal signaling pathways that lead to further platelet activation and granule release and (2) the capacity of the platelet to bind to other adhesive proteins/platelets. Both of these processes contribute to the formation of a thrombus. Stimulation of nonthrombotic receptors results in platelet adhesion or interaction with other vascular cells including endothelial cells, neutrophils, and mononuclear cells.

Many families and subfamilies of receptors are found on platelets that regulate a variety of platelet functions. These include the seven transmembrane receptor family, which is the main agonist-stimulated receptor family. Several seven transmembrane receptors are found on platelets, including the ADP receptors, prostaglandin receptors, lipid receptors, and chemokine receptors. Receptors for thrombin comprise the major seven transmembrane receptors found on platelets. Among this last group, the first identified was the protease activation receptor 1 (PAR1). The PAR class of receptors has a distinct mechanism of activation that involves specific cleavage of the N-terminus of thrombin, which, in turn, acts as a ligand for the receptor. Other PAR receptors are present on platelets, including PAR2 (not activated by thrombin) and PAR4. Adenosine receptors are responsible for transduction of ADP-induced signaling events, which are initiated by the binding of ADP to purinergic receptors on the platelet surface. There are several distinct ADP receptors, classified as P2X₁, P2Y₁, and P2Y₁₂. The activation of both the P2Y₁₂ and P2Y₁ receptors is essential for ADP-induced platelet aggregation. The thienopyridine derivatives, clopidogrel and prasugrel, are clinically used inhibitors of ADP-induced platelet aggregation.

Platelet Aggregation Activation of platelets results in a rapid series of signal transduction events, including tyrosine kinase, serine/threonine kinase, and lipid kinase activation. In unstimulated platelets, the major platelet integrin GPIIb/IIIa is maintained in an inactive conformation and functions as a low-affinity adhesion receptor for fibrinogen. This integrin is unique as it is only expressed on platelets. After stimulation, the interaction between fibrinogen and GPIIb/IIIa forms intercellular connections between platelets, leading to the formation of a platelet aggregate (Fig. 142-1). A calcium-sensitive conformational change in the extracellular domain of GPIIb/IIIa enables the high-affinity binding of soluble plasma fibrinogen as a result of a complex network of inside-out signaling events. The GPIIb/IIIa receptor serves as a bidirectional conduit with GPIIb/IIIa-mediated signaling (outside-in) occurring immediately after the binding of fibrinogen. This leads to additional intracellular signaling that further stabilizes the platelet aggregate and transforms platelet aggregation from a reversible to an irreversible process (inside-out).

THE ROLE OF PLATELETS AND THROMBOSIS IN INFLAMMATION

Inflammation plays an important role during the acute thrombotic phase of acute coronary syndromes. In the setting of acute upper respiratory infections, people are at higher risk of myocardial infarction and thrombotic stroke. Patients with acute coronary syndromes have not only increased interactions between platelets (homotypic aggregates), but also increased interactions between platelets and leukocytes (heterotypic aggregates) detectable in circulating blood. These latter aggregates form when platelets are activated and adhere to circulating leukocytes. Platelets bind via P-selectin (CD62P) expressed on the surface of activated platelets to the leukocyte receptor, P-selectin glycoprotein ligand 1 (PSGL-1). This association leads to increased expression of CD11b/CD18 (Mac-1) on leukocytes, which itself supports interactions with platelets partially via bivalent fibrinogen

linking this integrin with its platelet surface counterpart, GPIIb/IIIa. Platelet surface P-selectin also induces the expression of tissue factor on monocytes, which promotes fibrin formation.

In addition to platelet-monocyte aggregates, the immunomodulator, soluble CD40 ligand (CD40L or CD154), also reflects a link between thrombosis and inflammation. The CD40 ligand is a trimeric transmembrane protein of the tumor necrosis factor family and, with its receptor CD40, is an important contributor to the inflammatory process leading both to thrombosis and atherosclerosis. While many immunologic and vascular cells have been found to express CD40 and/or CD40 ligand, in platelets, CD40 ligand is rapidly translocated to the surface after stimulation and is upregulated in the newly formed thrombus. The surface-expressed CD40 ligand is cleaved from the platelet to generate a soluble fragment (soluble CD40 ligand).

Links have also been established among platelets, infection, immunity, and inflammation. Bacterial and viral infections are associated with a transient increase in the risk of acute thrombotic events, such as acute myocardial infarction and stroke. In addition, platelets contribute significantly to the pathophysiology and high mortality rates of sepsis. The expression, functionality, and signaling pathways of toll-like receptors (TLRs) have been established in platelets. Stimulation of platelet TLR2, TLR3, and TLR4 directly and indirectly activates the platelet's thrombotic and inflammatory responses, and live bacteria induce a proinflammatory response in platelets in a TLR2-dependent manner, suggesting a mechanism by which specific bacteria and bacterial components can directly activate platelet-dependent thrombosis.

GENETICS OF ARTERIAL THROMBOSIS



Some studies have associated arterial thrombosis with genetic variants (Table 142-1 A); however, the associations have been weak and not confirmed in larger series. Platelet count and mean platelet volume have been studied by genome-wide association studies (GWAS), and this approach identified signals located to noncoding regions. Of 15 quantitative trait loci associated with mean platelet volume and platelet count, one located at 12q24 is also a risk locus for coronary artery disease.

In the area of genetic variability and platelet function, studies have primarily dealt with pharmacogenetics, the field of pharmacology dealing with the interindividual variability in drug response based on genetic determinants (Table 142-2). This focus has been driven by the wide variability among individuals in terms of response to antithrombotic drugs and the lack of a common explanation for this variance. The best described is the issue of “aspirin resistance,” although heterogeneity for other antithrombotics (e.g., clopidogrel) has also been extensively examined. Primarily, platelet-dependent genetic determinants have been defined at the level of (1) drug effect, (2) drug compliance, and (3) drug metabolism. Many candidate platelet genes have been studied for their interaction with antiplatelet and antithrombotic agents.

Many patients have an inadequate response to the inhibitory effects of aspirin. Heritable factors contribute to the variability; however, ex vivo tests of residual platelet responsiveness after aspirin administration have not provided firm evidence for a pharmacogenetic interaction between aspirin and COX1 or other relevant platelet receptors. As such, currently, there is no clinical indication for genotyping to optimize aspirin's antiplatelet efficiency. For the platelet P2Y₁₂ receptor inhibitor clopidogrel, additional data suggest that genetics may affect the drug's responsiveness and utility. The responsible genetic variant appears not to be the expected P2Y₁₂ receptor but an enzyme responsible for drug metabolism. Clopidogrel is a prodrug, and liver metabolism by specific cytochrome P450 enzymes is required for activation. The genes encoding the CYP-dependent oxidative steps are polymorphic, and carriers of specific alleles of the CYP2C19 and CYP3A4 loci have increased platelet aggregability. Increased platelet activity has also been specifically associated with the CYP2C19*2 allele, which causes loss of platelet function in select patients. Because these are common genetic variants, this observation has been shown to be clinically relevant in large studies. In summary, although the loss-of-function polymorphisms in CYP2C19 is the strongest individual variable affecting pharmacokinetics and antiplatelet response to clopidogrel, it only

TABLE 142-1 HERITABLE CAUSES OF ARTERIAL AND VENOUS THROMBOSIS

A. Arterial Thrombosis	
<i>Platelet Receptors</i>	
β3 and α2 integrins	
P ₁ A2 polymorphism	
Fc(γ)RIIA	
GPIV T13254C polymorphism	
GPIb	
Thrombin receptor PAR1-5061 → D	
<i>Redox Enzymes</i>	
Plasma glutathione peroxidase	
H2 promoter haplotype	
Endothelial nitric oxide synthase	
–786T/C, –922A/G, –1468T/A	
Paraoxonase	
–107T allele, 192R allele	
<i>Homocysteine</i>	
Cystathionine β-synthase 833T → C	
5,10-Methylene tetrahydrofolate reductase (MTHFR) 677C → T	
B. Venous Thrombosis	
<i>Procoagulant Proteins</i>	
Fibrinogen	
–455G/A, –854G/A	
Prothrombin (20210G → A)	
<i>Protein C Anticoagulant Pathway</i>	
Factor V Leiden: 1691G → A (Arg506Gln)	
Thrombomodulin 1481C → T (Ala455Val)	
<i>Fibrinolytic Proteins with Known Polymorphisms</i>	
Tissue plasminogen activator (tPA)	
7351C/T, 20 099T/C in exon 6, 27 445T/A in intron 10	
Plasminogen activator inhibitor (PAI-1)	
4G/5G insertion/deletion polymorphism at position –675	
<i>Homocysteine</i>	
Cystathionine β-synthase 833T → C	
5,10-MTHFR 677C → T	

accounts for 5–12% of the variability in ADP-induced platelet aggregation on clopidogrel. In addition, genetic variables do not appear to significantly contribute to the clinical outcomes of patients treated with the P2Y₁₂ receptor antagonists prasugrel or ticagrelor.

VENOUS THROMBOSIS

OVERVIEW OF VENOUS THROMBOSIS

Coagulation is the process by which thrombin is activated and soluble plasma fibrinogen is converted into insoluble fibrin. These steps account for both normal hemostasis and the pathophysiologic processes influencing the development of venous thrombosis. The primary forms of venous thrombosis are deep vein thrombosis (DVT)

TABLE 142-2 GENETIC VARIATION AND PHARMACOGENETIC RESPONSES TO PLATELET INHIBITORS

Potential Gene Altered	Target Therapeutic Class	Specific Drug
P2Y ₁ and P2Y ₁₂ CYP2C19, CYP3A4, CYP3A5	ADP receptor inhibitors	Clopidogrel, prasugrel
COX1, COX2	Cyclooxygenase inhibitors	Aspirin
PIA1/A2	Receptor inhibitors	Abciximab, eptifibatid, tirofiban
INTB3, GPIbA	Glycoprotein IIb-IIIa receptor inhibitors	

TABLE 142-3 ACQUIRED CAUSES OF VENOUS THROMBOSIS

Surgery
Neurosurgery
Major abdominal surgery
Malignancy
Antiphospholipid syndrome
Other
Trauma
Pregnancy
Long-haul travel
Obesity
Oral contraceptives/hormone replacement
Myeloproliferative disorders
Polycythemia vera

in the extremities and the subsequent embolization to the lungs (pulmonary embolism), referred to together as venous thromboembolic disease. Venous thrombosis occurs due to heritable causes (Table 142-1 B) and acquired causes (Table 142-3).

DEEP VENOUS THROMBOSIS AND PULMONARY EMBOLISM

More than 200,000 new cases of venous thromboembolism occur each year. Of these cases, up to 30% of patients die within 30 days and one-fifth suffer sudden death due to pulmonary embolism; 30% go on to develop recurrent venous thromboembolism within 10 years. Data from the Atherosclerosis Risk in Communities (ARIC) study reported a 9% 28-day fatality rate from DVT and a 15% fatality rate from pulmonary embolism. Pulmonary embolism in the setting of cancer has a 25% fatality rate. The mean incidence of first DVT in the general population is 5 per 10,000 person-years; the incidence is similar in males and females when adjusting for factors related to reproduction and birth control and increases dramatically with age from 2 to 3 per 10,000 person-years at 30–49 years of age to 20 at 70–79 years of age.

OVERVIEW OF THE COAGULATION CASCADE AND ITS ROLE IN VENOUS THROMBOSIS

Coagulation is defined as the formation of fibrin by a series of linked enzymatic reactions in which each reaction product converts the subsequent inactive zymogen into an active serine protease (Fig. 142-2). This coordinated sequence is called the coagulation cascade and is a key mechanism for regulating hemostasis. Central to the function of the coagulation cascade is the principle of amplification: due to a series of linked enzymatic reactions, a small stimulus can lead to much greater quantities of fibrin, the end product that prevents hemorrhage at the site of vascular injury. In addition to the known risk factors relevant to hypercoagulopathy, stasis, and vascular dysfunction, newer areas of research have identified contributions from procoagulant microparticles, inflammatory cells, microvesicles, and fibrin structure.

The coagulation cascade is primarily initiated by vascular injury exposing tissue factor to blood components (Fig. 142-2). Tissue factor may also be found in bloodborne cell-derived microparticles and, under pathophysiologic conditions, in leukocytes or platelets. Plasma factor VII (FVII) is the ligand for and is activated (FVIIa) by binding to tissue factor exposed at the site of vessel damage. The binding of FVII/VIIa to tissue factor activates the downstream conversion of factor X (FX) to active FX (FXa). In an alternative reaction, the FVII/FVIIa–tissue factor complex initially converts FIX to FIXa, which then activates FX in conjunction with its cofactor factor VIII (FVIIIa). Factor Xa with its cofactor FVa converts prothrombin to thrombin, which then converts soluble plasma fibrinogen to insoluble fibrin, leading to clot or thrombus formation. Thrombin also activates FXIII to FXIIIa, a transglutaminase that covalently cross-links and stabilizes the fibrin clot. Formation of thrombi is affected by mechanisms governing fibrin structure and stability including specific fibrinogen variants and how they alter fibrin formation, strength and structure.

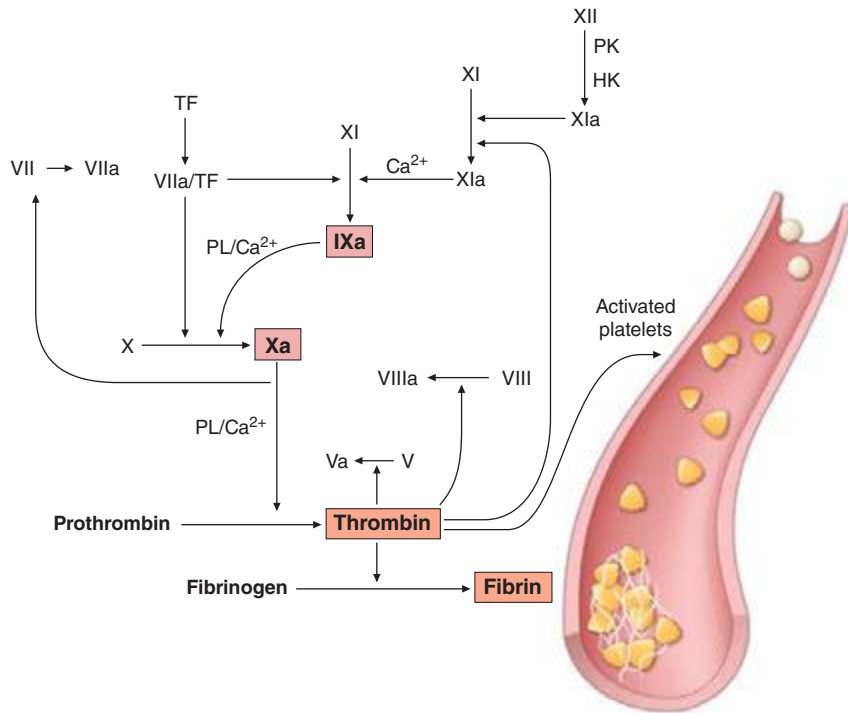


FIGURE 142-2 Summary of the coagulation pathways. Specific coagulation factors (“a” indicates activated form) are responsible for the conversion of soluble plasma fibrinogen into insoluble fibrin. This process occurs via a series of linked reactions in which the enzymatically active product subsequently converts the downstream inactive protein into an active serine protease. In addition, the activation of thrombin leads to stimulation of platelets. HK, high-molecular-weight kininogen; PK, prekallikrein; TF, tissue factor.

Several antithrombotic factors also regulate coagulation; these include antithrombin, tissue factor pathway inhibitor (TFPI), heparin cofactor II, and protein C/protein S. Under normal conditions, these factors limit the production of thrombin to prevent the perpetuation of coagulation and thrombus formation. Typically, after the clot has caused occlusion at the damaged site and begins to expand toward adjacent uninjured vessel segments, the anticoagulant reactions governed by the normal endothelium become pivotal in limiting the extent of this hemostatically protective clot.

RISK FACTORS FOR VENOUS THROMBOSIS

The risk factors for venous thrombosis are primarily related to hypercoagulability, which can be genetic (Table 142-1) or acquired, or due to immobilization and venous stasis. Independent predictors for recurrence include increasing age, obesity, malignant neoplasm, and acute extremity paresis. It is estimated that 5–8% of the U.S. population has a genetic risk factor known to predispose to venous thrombosis. Often, multiple risk factors are present in a single individual. Significant risk is incurred by major orthopedic, abdominal, or neurologic surgeries. Moderate risk is promoted by prolonged bedrest; certain types of cancer, pregnancy, hormone replacement therapy, or oral contraceptive use; and other sedentary conditions such as long-distance plane travel. It has been reported that the risk of developing a venous thromboembolic event doubles after air travel lasting 4 h, although the absolute risk remains low (1 in 6000). The relative risk of venous thromboembolism among pregnant or postpartum women is 4.3, and the overall incidence (absolute risk) is 199.7 per 100,000 woman-years.

GENETICS OF VENOUS THROMBOSIS



(See Table 142-2) Less common causes of venous thrombosis are those due to genetic variants. These abnormalities include loss-of-function mutations of endogenous anticoagulants as well as gain-of-function mutations of procoagulant proteins. Heterozygous antithrombin deficiency and homozygosity of the factor V Leiden mutation

significantly increase the risk of venous thrombosis. While homozygous protein C or protein S deficiencies are rare and may lead to fatal purpura fulminans, heterozygous deficiencies are associated with a moderate risk of thrombosis. Activated protein C impairs coagulation by proteolytic degradation of FVa. Patients resistant to the activity of activated protein C may have a point mutation in the FV gene located on chromosome 1, a mutant denoted factor V Leiden. Mildly increased risk has been attributed to elevated levels of procoagulant factors, as well as low levels of tissue factor pathway inhibitor. Polymorphisms of methylene tetrahydrofolate reductase as well as hyperhomocysteinemia have been shown to be independent risk factors for venous thrombosis, as well as arterial vascular disease; however, many of the initial descriptions of genetic variants and their associations with thromboembolism are being questioned in larger, more current studies.

FIBRINOLYSIS AND THROMBOSIS

Specific abnormalities in the fibrinolytic system have been associated with enhanced thrombosis. Factors such as elevated levels of tissue plasminogen activator (tPA) and plasminogen activator inhibitor type 1 (PAI-1) have been associated with decreased fibrinolytic activity and an increased risk of arterial thrombotic disease. Specific genetic variants have been associated with decreased fibrinolytic activity, including the 4G/5G insertion/deletion polymorphism in the (plasminogen activator type 1) *PAI-1* gene. Additionally, the 311-bp Alu insertion/deletion in tPA’s intron 8 has been associated with enhanced thrombosis; however, genetic abnormalities have not been associated consistently with altered function or tPA levels, raising questions about the relevant pathophysiologic mechanism. Thrombin-activatable fibrinolysis inhibitor (TAFI) is a carboxypeptidase that regulates fibrinolysis; elevated plasma TAFI levels have been associated with an increased risk of both DVT and cardiovascular disease.

The metabolic syndrome also is accompanied by altered fibrinolytic activity. This syndrome, which comprises abdominal fat (central obesity), altered glucose and insulin metabolism, dyslipidemia, and hypertension, has been associated with atherothrombosis. The mechanism for enhanced thrombosis appears to be due both to altered platelet function and to a procoagulant and hypofibrinolytic state. One of the most frequently documented prothrombotic abnormalities reported in this syndrome is an increase in plasma levels of PAI-1.

In addition to contributing to platelet function, inflammation plays a role in both coagulation-dependent thrombus formation and thrombus resolution. Both polymorphonuclear neutrophils and monocytes/macrophages contribute to multiple overlapping thrombotic functions, including fibrinolysis, chemokine and cytokine production, and phagocytosis.

THE DISTINCTION BETWEEN ARTERIAL AND VENOUS THROMBOSIS

Although there is overlap, venous thrombosis and arterial thrombosis are initiated differently, and clot formation progresses by somewhat distinct pathways. In the setting of stasis or states of hypercoagulability, venous thrombosis is activated with the initiation of the coagulation cascade primarily due to exposure of tissue factor; this leads to the formation of thrombin and the subsequent conversion of fibrinogen to fibrin. In the artery, thrombin formation also occurs, but thrombosis is primarily promoted by the adhesion of platelets to an injured vessel and stimulated by exposed extracellular matrix (Figs. 142-1 and 142-2). There is wide variation in individual responses to vascular injury, an

important determinant of which is the predisposition an individual has to arterial or venous thrombosis. This concept has been supported indirectly in prothrombotic animal models in which there is poor correlation between the propensity to develop venous versus arterial thrombosis.

Despite considerable progress in understanding the role of hypercoagulable states in venous thromboembolic disease, the contribution of hypercoagulability to arterial vascular disease is much less well understood. Although specific thrombophilic conditions, such as factor V Leiden and the prothrombin G20210A mutation, are risk factors for DVT, pulmonary embolism, and other venous thromboembolic events, their contribution to arterial thrombosis is less well defined. In fact, to the contrary, many of these thrombophilic factors have not been found to be clinically important risk factors for arterial thrombotic events, such as acute coronary syndromes.

Clinically, although the pathophysiology is distinct, arterial and venous thrombosis do share common risk factors, including age, obesity, cigarette smoking, diabetes mellitus, arterial hypertension, hyperlipidemia, and metabolic syndrome. Select genetic variants, including those of the glutathione peroxidase gene, have also been associated with arterial and venous thrombo-occlusive disease. Importantly, arterial and venous thrombosis may both be triggered by pathophysiologic stimuli responsible for activating inflammatory and oxidative pathways.

The diagnosis and treatment of ischemic heart disease are discussed in Chap. 293. Stroke diagnosis and management are discussed in Chap. 330. The diagnosis and management of DVT and pulmonary embolus are discussed in Chap. 300.

143 Antiplatelet, Anticoagulant, and Fibrinolytic Drugs

Jeffrey I. Weitz

Thromboembolic disorders are major causes of morbidity and mortality. Thrombosis can occur in arteries or veins. Arterial thrombosis is the most common cause of acute myocardial infarction (MI), ischemic stroke, and limb gangrene. Venous thromboembolism encompasses deep vein thrombosis (DVT), which can lead to postthrombotic syndrome, and pulmonary embolism (PE), which can be fatal or can result in chronic thromboembolic pulmonary hypertension.

Most arterial thrombi are superimposed on disrupted atherosclerotic plaque because plaque rupture exposes thrombogenic material in the plaque core to the blood. This material then triggers platelet aggregation and fibrin formation, which results in the generation of a platelet-rich thrombus that can temporarily or permanently occlude blood flow. In contrast, venous thrombi rarely form at sites of obvious vascular disruption. Although they can develop after surgical trauma to veins or secondary to indwelling venous catheters, venous thrombi usually originate in the valve cusps of the deep veins of the calf or in the muscular sinuses. Sluggish blood flow reduces the oxygen supply to the avascular valve cusps. Endothelial cells lining these valve cusps become activated and express adhesion molecules on their surface. Tissue factor-bearing leukocytes and microparticles adhere to these activated cells and induce coagulation. DNA extruded from neutrophils forms neutrophil extracellular traps (NETs) that provide a scaffold that traps red blood cells, promotes platelet adhesion and activation, and augments coagulation. Local thrombus formation is exacerbated by reduced clearance of activated clotting factors as a result of impaired blood flow. If the thrombi extend from the calf veins into the popliteal and more proximal veins of the leg, thrombus fragments can dislodge, travel to the lungs, and produce a PE.

Arterial and venous thrombi are composed of platelets, fibrin, and trapped red blood cells, but the proportions differ. Arterial thrombi

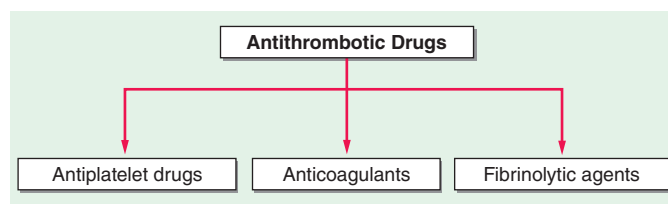


FIGURE 143-1 Classification of antithrombotic drugs.

are rich in platelets because of the high shear in the injured arteries. In contrast, venous thrombi, which form under low shear conditions, contain relatively few platelets and are predominantly composed of fibrin and trapped red cells. Because of the predominance of platelets, arterial thrombi appear white, whereas venous thrombi are red in color, reflecting the trapped red cells.

Antithrombotic drugs are used for prevention and treatment of thrombosis. Targeting the components of thrombi, these agents include (1) antiplatelet drugs, (2) anticoagulants, and (3) fibrinolytic agents (Fig. 143-1). With the predominance of platelets in arterial thrombi, strategies to attenuate arterial thrombosis focus mainly on antiplatelet agents, although, in the acute setting, often include anticoagulants and fibrinolytic agents. Anticoagulants are the mainstay of prevention and treatment of venous thromboembolism because fibrin is the predominant component of venous thrombi. Antiplatelet drugs are less effective than anticoagulants in this setting because of the limited platelet content of venous thrombi. Fibrinolytic therapy is used in selected patients with venous thromboembolism. For example, patients with massive or submassive PE can benefit from systemic or catheter-directed fibrinolytic therapy. Pharmaco-mechanical therapy also is used to restore blood flow in patients with extensive DVT involving the iliac and/or femoral veins.

ANTIPLATELET DRUGS

ROLE OF PLATELETS IN ARTERIAL THROMBOSIS

In healthy vasculature, circulating platelets are maintained in an inactive state by nitric oxide (NO) and prostacyclin released by endothelial cells lining the blood vessels. In addition, endothelial cells also express CD39 on their surface, a membrane-associated ecto-adenosine diphosphatase (ADPase) that degrades ADP released from activated platelets. When the vessel wall is damaged, release of these substances is impaired and subendothelial matrix is exposed. Platelets adhere to exposed collagen via $\alpha_2\beta_1$ and glycoprotein (Gp) VI and to von Willebrand factor (VWF) via Gp Iba and Gp IIb/IIIa ($\alpha_{IIb}\beta_3$)—receptors that are constitutively expressed on the platelet surface. Adherent platelets undergo a change in shape, secrete ADP from their dense granules, and synthesize and release thromboxane A_2 . Released ADP and thromboxane A_2 , which are platelet agonists, activate ambient platelets and recruit them to the site of vascular injury (Fig. 143-2).

Disruption of the vessel wall also exposes tissue factor-expressing cells to the blood. Tissue factor binds factor VIIa and initiates coagulation. Activated platelets potentiate coagulation by providing a surface that binds clotting factors and supports the assembly of activation complexes that enhance thrombin generation. In addition to converting fibrinogen to fibrin, thrombin serves as a potent platelet agonist and recruits more platelets to the site of vascular injury. Thrombin also amplifies its own generation by feedback activation of factors V, VIII, and XI and solidifies the fibrin network by activating factor XIII, which then cross-links the fibrin strands.

When platelets are activated, Gp IIb/IIIa, the most abundant receptor on the platelet surface, undergoes a conformational change that enables it to bind fibrinogen and, under high shear conditions, VWF. Divalent fibrinogen or multivalent VWF molecules bridge adjacent platelets together to form platelet aggregates. Fibrin strands, generated through the action of thrombin, then weave these aggregates together to form a platelet/fibrin mesh.

Antiplatelet drugs target various steps in this process. The commonly used drugs include aspirin, ADP receptor inhibitors, which

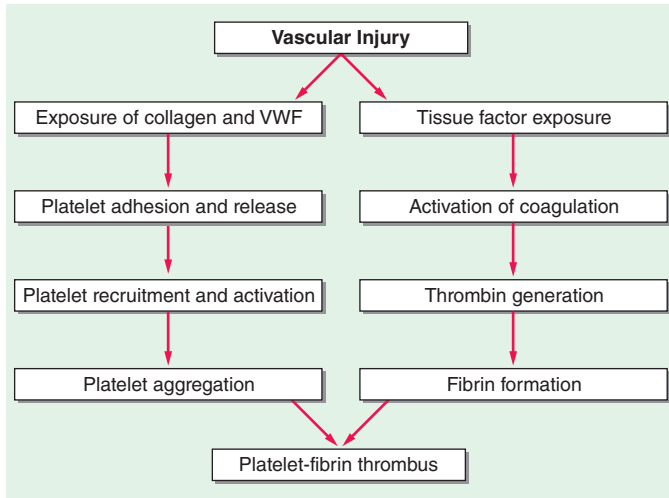


FIGURE 143-2 Coordinated role of platelets and the coagulation system in thrombogenesis.

Vascular injury simultaneously triggers platelet activation and aggregation and activation of the coagulation system. Platelet activation is initiated by exposure of subendothelial collagen and von Willebrand factor (VWF), onto which platelets adhere. Adherent platelets become activated and release ADP and thromboxane A_2 , platelet agonists that activate ambient platelets and recruit them to the site of injury. When platelets are activated, glycoprotein IIb/IIIa on their surface undergoes a conformational change that enables it to ligate fibrinogen and/or VWF and mediate platelet aggregation. Coagulation is triggered by tissue factor exposed at the site of injury. Tissue factor triggers thrombin generation. As a potent platelet agonist, thrombin amplifies platelet recruitment to the site of injury. Thrombin also converts fibrinogen to fibrin, and the fibrin strands then weave the platelet aggregates together to form a platelet/fibrin thrombus.

include the thienopyridines (clopidogrel and prasugrel) and ticagrelor, dipyridamole, and Gp IIb/IIIa antagonists.

ASPIRIN

The most widely used antiplatelet agent worldwide is aspirin. As a cheap and effective antiplatelet drug, aspirin serves as the foundation of most antiplatelet strategies.

Mechanism of Action Aspirin produces its antithrombotic effect by irreversibly acetylating and inhibiting platelet cyclooxygenase (COX)-1 (Fig. 143-3), a critical enzyme in the biosynthesis of thromboxane A_2 . At high doses (~1 g/d), aspirin also inhibits COX-2, an inducible COX isoform found in endothelial cells and inflammatory cells. In endothelial cells, COX-2 initiates the synthesis of prostacyclin, a potent vasodilator and inhibitor of platelet aggregation.

Indications Aspirin is widely used for secondary prevention of cardiovascular events in patients with coronary artery, cerebrovascular, or peripheral vascular disease. Compared with placebo, aspirin produces a 25% reduction in the risk of cardiovascular death, MI, or stroke. Aspirin is also used for primary prevention in patients whose estimated annual risk of MI is >1%, a point where its benefits are likely to outweigh harms. This includes patients older than age 40 years with two or more major risk factors for cardiovascular disease or men older than age 45 years and women over the age of 55 years with one or more such risk factors. Aspirin is equally effective in men and women. In men, aspirin mainly reduces the risk of MI, whereas in women, aspirin lowers the risk of stroke.

Dosages Aspirin is usually administered at doses of 75–325 mg once daily. Higher doses of aspirin are not more effective than lower aspirin doses, and some analyses suggest reduced efficacy with higher doses. Because the side effects of aspirin are dose-related, daily aspirin doses of 75–100 mg are recommended for most indications. When rapid

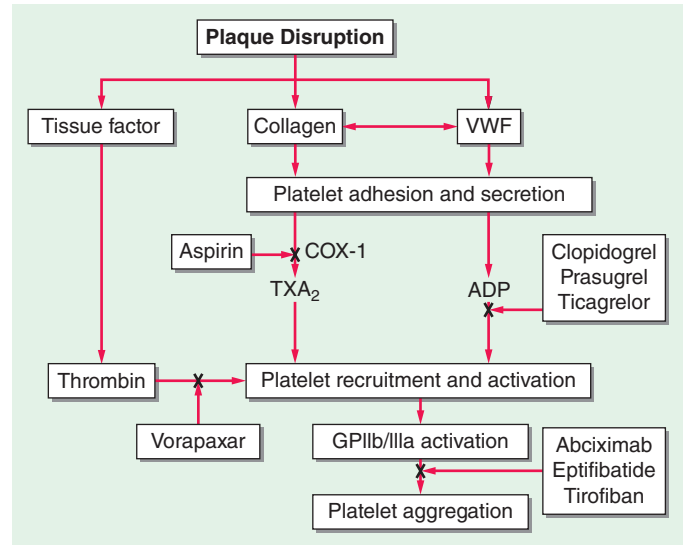


FIGURE 143-3 Site of action of antiplatelet drugs. Aspirin inhibits thromboxane A_2 (TXA₂) synthesis by irreversibly acetylating cyclooxygenase-1 (COX-1). Reduced TXA₂ release attenuates platelet activation and recruitment to the site of vascular injury. Clopidogrel and prasugrel irreversibly block P2Y₁₂, a key ADP receptor on the platelet surface; ticagrelor and prasugrel are reversible inhibitors of P2Y₁₂. Abciximab, eptifibatide, and tirofiban inhibit the final common pathway of platelet aggregation by blocking fibrinogen and von Willebrand factor binding to activated glycoprotein (Gp) IIb/IIIa. Vorapaxar inhibits thrombin-mediated platelet activation by targeting protease-activated receptor-1 (PAR-1), the major thrombin receptor on human platelets.

platelet inhibition is required, an initial aspirin dose of at least 160 mg should be given.

Side Effects The most common side effects are gastrointestinal and range from dyspepsia to erosive gastritis or peptic ulcers with bleeding and perforation. These side effects are dose-related. Use of enteric-coated or buffered aspirin in place of plain aspirin does not eliminate gastrointestinal side effects. The overall risk of major bleeding with aspirin is 1–3% per year. The risk of bleeding is increased two- to threefold when aspirin is given in conjunction with other antiplatelet drugs, such as clopidogrel, or with anticoagulants, such as warfarin. When dual or triple therapy is prescribed, low-dose aspirin should be given (75–100 mg daily). Eradication of *Helicobacter pylori* infection and administration of proton pump inhibitors may reduce the risk of aspirin-induced upper gastrointestinal bleeding in patients with peptic ulcer disease.

Aspirin should not be administered to patients with a history of aspirin allergy characterized by bronchospasm. This problem occurs in ~0.3% of the general population but is more common in those with chronic urticaria or asthma, particularly in individuals with nasal polyps or chronic rhinitis. Hepatic and renal toxicity are observed with aspirin overdose.

Aspirin Resistance Clinical aspirin resistance is defined as the failure of aspirin to protect patients from ischemic vascular events. This is not a helpful definition because it is made after the event occurs. Furthermore, it is not realistic to expect aspirin, which only blocks thromboxane A_2 -induced platelet activation, to prevent all vascular events.

Aspirin resistance has also been described biochemically as failure of the drug to produce its expected inhibitory effects on tests of platelet function, such as thromboxane A_2 synthesis or arachidonic acid-induced platelet aggregation. Potential causes of aspirin resistance include poor compliance, reduced absorption, drug-drug interaction with ibuprofen, and overexpression of COX-2. Unfortunately, the tests for aspirin resistance have not been well standardized, and there is little evidence that they identify patients at increased risk of recurrent vascular events, or that resistance can be reversed by giving

higher doses of aspirin or by adding other antiplatelet drugs. Until such information is available, testing for aspirin resistance remains a research tool.

ADP RECEPTOR ANTAGONISTS

The ADP receptor antagonists include the thienopyridines (clopidogrel and prasugrel) and ticagrelor. All of these drugs target P2Y₁₂, the key ADP receptor on platelets.

Thienopyridines • MECHANISM OF ACTION The thienopyridines are structurally related drugs that selectively inhibit ADP-induced platelet aggregation by irreversibly blocking P2Y₁₂ (Fig. 143-3). Clopidogrel and prasugrel are prodrugs that require metabolic activation by the hepatic cytochrome P450 (CYP) enzyme system. Prasugrel is about 10-fold more potent than clopidogrel and has a more rapid onset of action because of better absorption and more streamlined metabolic activation.

INDICATIONS When compared with aspirin in patients with recent ischemic stroke, recent MI, or a history of peripheral arterial disease, clopidogrel reduced the risk of cardiovascular death, MI, and stroke by 8.7%. Therefore, clopidogrel is more effective than aspirin but is also more expensive. In some patients, clopidogrel and aspirin are combined to capitalize on their capacity to block complementary pathways of platelet activation. For example, the combination of aspirin plus clopidogrel is recommended for at least 4 weeks after implantation of a bare metal stent in a coronary artery and for at least a year in those with a drug-eluting stent. Concerns about late in-stent thrombosis with drug-eluting stents have led some experts to recommend long-term use of clopidogrel plus aspirin for the latter indication. However, these recommendations are likely to change because the risk of late stent thrombosis is decreasing with the newer generation of drug-eluting coronary stents.

The combination of clopidogrel and aspirin is also effective in patients with unstable angina. Thus, in 12,562 such patients, the risk of cardiovascular death, MI, or stroke was 9.3% in those randomized to the combination of clopidogrel and aspirin and 11.4% in those given aspirin alone. This 20% relative risk reduction with combination therapy was highly statistically significant. However, combining clopidogrel with aspirin increases the risk of major bleeding to about 2% per year. This bleeding risk persists even if the daily dose of aspirin is ≤100 mg. Therefore, the combination of clopidogrel and aspirin should only be used when there is a clear benefit. For example, this combination has not proven to be superior to clopidogrel alone in patients with acute ischemic stroke or to aspirin alone for primary prevention in those at risk for cardiovascular events.

Prasugrel was compared with clopidogrel in 13,608 patients with acute coronary syndromes who were scheduled to undergo percutaneous coronary intervention. The incidence of the primary efficacy endpoint, a composite of cardiovascular death, MI, or stroke, was significantly lower with prasugrel than with clopidogrel (9.9% and 12.1%, respectively), mainly reflecting a reduction in the incidence of nonfatal MI. The incidence of stent thrombosis also was significantly lower with prasugrel (1.1% and 2.4%, respectively). However, these advantages were at the expense of significantly higher rates of fatal bleeding (0.4% and 0.1%, respectively) and life-threatening bleeding (1.4% and 0.9%, respectively) with prasugrel. Because patients older than age 75 years and those with a history of prior stroke or transient ischemic attack have a particularly high risk of bleeding, prasugrel should generally be avoided in older patients, and the drug is contraindicated in those with a history of cerebrovascular disease. Caution is required if prasugrel is used in patients weighing less than 60 kg or in those with renal impairment.

When prasugrel was compared with clopidogrel in 7243 patients with unstable angina or MI without ST-segment elevation, prasugrel failed to reduce the rate of the primary efficacy endpoint, which was a composite of cardiovascular death, MI, and stroke. Because of the negative results of this study, prasugrel is reserved for patients undergoing percutaneous coronary intervention. In this setting, prasugrel is usually given in conjunction with aspirin. To reduce the risk of bleeding, the daily aspirin dose should be ≤100 mg.

DOSING Clopidogrel is given once daily at a dose of 75 mg. Loading doses of clopidogrel are given when rapid ADP receptor blockade is desired. For example, patients undergoing coronary stenting are often given a loading dose of 300 mg, which produces inhibition of ADP-induced platelet aggregation in about 6 h; loading doses of 600 or 900 mg produce an even more rapid effect. After a loading dose of 60 mg, prasugrel is given once daily at a dose of 10 mg. Patients older than age 75 years or weighing less than 60 kg should receive a lower daily prasugrel dose of 5 mg.

SIDE EFFECTS The most common side effect of clopidogrel and prasugrel is bleeding. Because of its greater potency, bleeding is more common with prasugrel than clopidogrel. To reduce the risk of bleeding, clopidogrel and prasugrel should be stopped 5–7 days before major surgery. In patients taking clopidogrel or prasugrel who present with serious bleeding, platelet transfusion may be helpful.

Hematologic side effects, including neutropenia, thrombocytopenia, and thrombotic thrombocytopenic purpura, are rare.

THIENOPYRIDINE RESISTANCE The capacity of clopidogrel to inhibit ADP-induced platelet aggregation varies among subjects. This variability reflects, at least in part, genetic polymorphisms in the CYP isoenzymes involved in the metabolic activation of clopidogrel. Most important of these is CYP2C19. Clopidogrel-treated patients with the loss-of-function CYP2C19*2 allele exhibit reduced platelet inhibition compared with those with the wild-type CYP2C19*1 allele and experience a higher rate of cardiovascular events. This is important because estimates suggest that up to 25% of whites, 30% of African Americans, and 50% of Asians carry the loss-of-function allele, which would render them resistant to clopidogrel. Even patients with the reduced function CYP2C19*3, *4, or *5 alleles may derive less benefit from clopidogrel than those with the full-function CYP2C19*1 allele. Concomitant administration of clopidogrel with proton pump inhibitors, which are inhibitors of CYP2C19, produces a small reduction in the inhibitory effects of clopidogrel on ADP-induced platelet aggregation. The extent to which this interaction increases the risk of cardiovascular events remains controversial.

In contrast to their effect on the metabolic activation of clopidogrel, CYP2C19 polymorphisms appear to be less important determinants of the activation of prasugrel. Thus, no association was detected between the loss-of-function allele and decreased platelet inhibition or increased rate of cardiovascular events with prasugrel. The observation that genetic polymorphisms affecting clopidogrel absorption or metabolism influence clinical outcomes raises the possibilities that pharmacogenetic profiling may be useful to identify clopidogrel-resistant patients and that point-of-care assessment of the extent of clopidogrel-induced platelet inhibition may help detect patients at higher risk for subsequent cardiovascular events. Clinical trials designed to evaluate these possibilities have thus far been negative. Although administration of higher doses of clopidogrel can overcome a reduced response to clopidogrel, the clinical benefit of this approach is uncertain. Instead, prasugrel or ticagrelor may be better choices for these patients.

Ticagrelor As an orally active inhibitor of P2Y₁₂, ticagrelor differs from the thienopyridines in that ticagrelor does not require metabolic activation and it produces reversible inhibition of the ADP receptor.

MECHANISM OF ACTION Like the thienopyridines, ticagrelor inhibits P2Y₁₂. Because it does not require metabolic activation, ticagrelor has a more rapid onset and offset of action than clopidogrel, and it produces greater and more predictable inhibition of ADP-induced platelet aggregation than clopidogrel.

INDICATIONS When compared with clopidogrel in patients with acute coronary syndromes, ticagrelor produced a greater reduction in the primary efficacy endpoint—a composite of cardiovascular death, MI, and stroke at 1 year—than clopidogrel (9.8% and 11.7%, respectively; $p = .001$). This difference reflected a significant reduction in both cardiovascular death (4.0% and 5.1%, respectively; $p = .001$) and MI (5.8% and 6.9%, respectively; $p = .005$) with ticagrelor compared with

748 clopidogrel. Rates of stroke were similar with ticagrelor and clopidogrel (1.5% and 1.3%, respectively), and no difference in rates of major bleeding was noted. When minor bleeding was added to the major bleeding results, however, ticagrelor showed an increase relative to clopidogrel (16.1% and 14.6%, respectively; $p = .008$). Ticagrelor also was superior to clopidogrel in patients with acute coronary syndrome who underwent percutaneous coronary intervention or cardiac surgery. Based on these observations, some guidelines give ticagrelor preference over clopidogrel, particularly in higher risk patients.

DOSING Ticagrelor is initiated with an oral loading dose of 180 mg followed by 90 mg twice daily. The dose does not require adjustment in patients with renal impairment, but the drug should be used with caution in patients with hepatic disease and in those receiving potent inhibitors or inducers of CYP3A4 because ticagrelor is metabolized in the liver via CYP3A4. Ticagrelor is usually administered in conjunction with aspirin; the daily aspirin dose should not exceed 100 mg.

SIDE EFFECTS In addition to bleeding, the most common side effects of ticagrelor are dyspnea, which can occur in up to 15% of patients, and asymptomatic ventricular pauses. The dyspnea, which tends to occur soon after initiating ticagrelor, is usually self-limiting and mild in intensity. The mechanism responsible for this side effect is unknown.

To reduce the risk of bleeding, ticagrelor should be stopped 5–7 days prior to major surgery. Platelet transfusions are unlikely to be of benefit in patients with ticagrelor-related bleeding because the drug will bind to P2Y₁₂ on the transfused platelets.

DIPYRIDAMOLE

Dipyridamole is a relatively weak antiplatelet agent on its own, but an extended-release formulation of dipyridamole combined with low-dose aspirin, a preparation known as *Aggrenox*, is used for prevention of stroke in patients with transient ischemic attacks.

Mechanism of Action By inhibiting phosphodiesterase, dipyridamole blocks the breakdown of cyclic adenosine monophosphate (AMP). Increased levels of cyclic AMP reduce intracellular calcium and inhibit platelet activation. Dipyridamole also blocks the uptake of adenosine by platelets and other cells. This produces a further increase in local cyclic AMP levels because the platelet adenosine A₂ receptor is coupled to adenylate cyclase (Fig. 143-4).

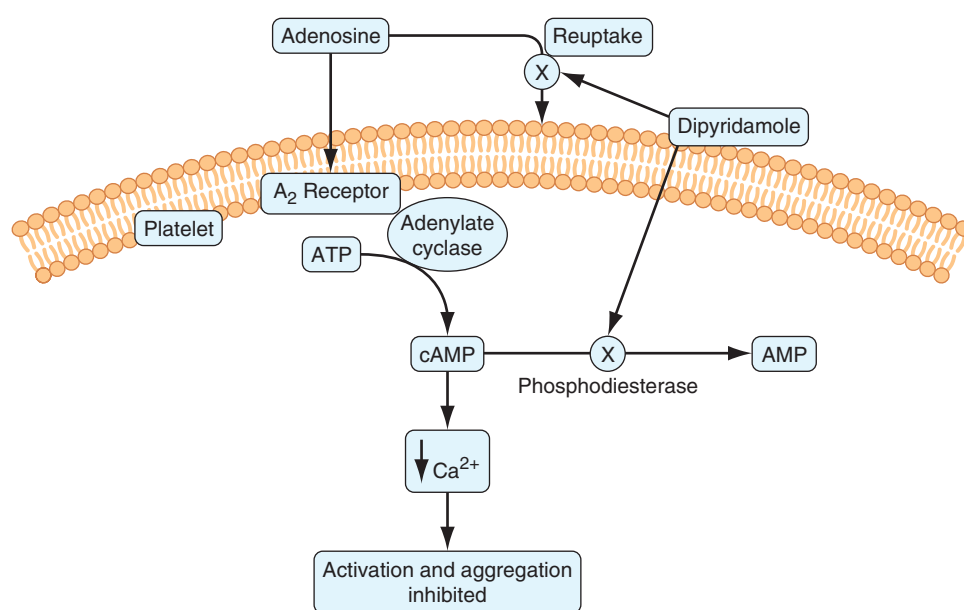


FIGURE 143-4 Mechanism of action of dipyridamole. Dipyridamole increases levels of cyclic AMP (cAMP) in platelets by (1) blocking the reuptake of adenosine and (2) inhibiting phosphodiesterase-mediated cyclic AMP degradation. By promoting calcium uptake, cyclic AMP reduces intracellular levels of calcium. This, in turn, inhibits platelet activation and aggregation.

Indications Dipyridamole plus aspirin was compared with aspirin or dipyridamole alone, or with placebo, in patients with an ischemic stroke or transient ischemic attack. The combination reduced the risk of stroke by 22.1% compared with aspirin and by 24.4% compared with dipyridamole. A second trial compared dipyridamole plus aspirin with aspirin alone for secondary prevention in patients with ischemic stroke. Vascular death, stroke, or MI occurred in 13% of patients given combination therapy and in 16% of those treated with aspirin alone. Another trial randomized 20,332 patients with noncardioembolic ischemic stroke to either Aggrenox or clopidogrel. The primary efficacy endpoint of recurrent stroke occurred in 9.0% of those given Aggrenox and in 8.8% of patients treated with clopidogrel. Although this difference was not statistically significant, the study failed to meet the prespecified margin to claim noninferiority of Aggrenox relative to clopidogrel. These results have dampened enthusiasm for the use of Aggrenox.

Because of its vasodilatory effects and the paucity of data supporting the use of dipyridamole in patients with symptomatic coronary artery disease, Aggrenox should not be used for stroke prevention in such patients. Clopidogrel is a better choice in this setting.

Dosing Aggrenox is given twice daily. Each capsule contains 200 mg of extended-release dipyridamole and 25 mg of aspirin.

Side Effects Because dipyridamole has vasodilatory effects, it must be used with caution in patients with coronary artery disease. Gastrointestinal complaints, headache, facial flushing, dizziness, and hypotension can also occur. These symptoms often subside with continued use of the drug.

GP IIB/IIIA RECEPTOR ANTAGONISTS

As a class, parenteral Gp IIB/IIIA receptor antagonists have an established niche in patients with acute coronary syndromes. The three agents in this class are abciximab, eptifibatid, and tirofiban.

Mechanism of Action A member of the integrin family of adhesion receptors, Gp IIB/IIIA is found on the surface of platelets and megakaryocytes. With about 80,000 copies per platelet, Gp IIB/IIIA is the most abundant receptor. Consisting of a noncovalently linked heterodimer, Gp IIB/IIIA is inactive on resting platelets. When platelets are activated, inside-outside signal transduction pathways trigger a conformational activation of the receptor. Once activated, Gp IIB/IIIA binds adhesive molecules, such as fibrinogen and, under high shear conditions, VWF. Binding is mediated by the Arg-Gly-Asp (RGD) sequence found on the α chains of fibrinogen and on VWF, and by the Lys-Gly-Asp (KGD) sequence located within a unique dodecapeptide domain on the γ chains of fibrinogen. Once bound, fibrinogen and/or VWF bridge adjacent platelets together to induce platelet aggregation.

Although abciximab, eptifibatid, and tirofiban all target the Gp IIB/IIIA receptor, they are structurally and pharmacologically distinct (Table 143-1). Abciximab is a Fab fragment of a humanized murine monoclonal antibody directed against the activated form of Gp IIB/IIIA. Abciximab binds to the activated receptor with high affinity and blocks the binding of adhesive molecules. In contrast, eptifibatid and tirofiban are synthetic small molecules. Eptifibatid is a cyclic heptapeptide that binds Gp IIB/IIIA because it incorporates the KGD motif, whereas tirofiban is a nonpeptidic tyrosine derivative that acts as an RGD mimetic. Abciximab has

TABLE 143-1 FEATURES OF GPIIB/IIIa ANTAGONISTS

Feature	Abciximab	Eptifibatide	Tirofiban
Description	Fab fragment of humanized mouse monoclonal antibody	Cyclical KGD-containing heptapeptide	Nonpeptidic RGD mimetic
Specific for Gp IIb/IIIa	No	Yes	Yes
Plasma half-life	Short (min)	Long (2.5 h)	Long (2.0 h)
Platelet-bound half-life	Long (days)	Short (s)	Short (s)
Renal clearance	No	Yes	Yes

a long half-life and can be detected on the surface of platelets for up to 2 weeks; eptifibatide and tirofiban have short half-lives.

Whereas eptifibatide and tirofiban are specific for Gp IIb/IIIa, abciximab also inhibits the closely related $\alpha_v\beta_3$ receptor, which binds vitronectin, and $\alpha_M\beta_2$, a leukocyte integrin. Inhibition of $\alpha_v\beta_3$ and $\alpha_M\beta_2$ may endow abciximab with anti-inflammatory and/or antiproliferative properties that extend beyond platelet inhibition.

Indications Abciximab and eptifibatide are used in patients undergoing percutaneous coronary interventions, particularly those who have not been pretreated with an ADP receptor antagonist. Tirofiban is used in high-risk patients with unstable angina. Eptifibatide also can be used for this indication.

Dosing All of the Gp IIb/IIIa antagonists are given as an IV bolus followed by an infusion. The recommended dose of abciximab is a bolus of 0.25 mg/kg followed by an infusion of 0.125 $\mu\text{g}/\text{kg}$ per minute to a maximum of 10 $\mu\text{g}/\text{kg}$ for 12 h. Eptifibatide is given as two 180 $\mu\text{g}/\text{kg}$ boluses given 10 min apart, followed by an infusion of 2.0 $\mu\text{g}/\text{kg}$ per minute for 18–24 h. Tirofiban is started at a rate of 0.4 $\mu\text{g}/\text{kg}$ per minute for 30 min; the drug is then continued at a rate of 0.1 $\mu\text{g}/\text{kg}$ per minute for up to 18 h. Because these agents are cleared by the kidneys, the doses of eptifibatide and tirofiban must be reduced in patients with renal insufficiency. Thus, the eptifibatide infusion is reduced to 1 $\mu\text{g}/\text{kg}$ per minute in patients with a creatinine clearance below 50 mL/min, whereas the dose of tirofiban is cut in half for patients with a creatinine clearance below 30 mL/min.

Side Effects In addition to bleeding, thrombocytopenia is the most serious complication. Thrombocytopenia is immune-mediated and is caused by antibodies directed against neoantigens on Gp IIb/IIIa that are exposed upon antagonist binding. With abciximab, thrombocytopenia occurs in up to 5% of patients. Thrombocytopenia is severe in ~1% of these individuals. Thrombocytopenia is less common with the other two agents, occurring in ~1% of patients.

NEW ANTIPLATELET AGENTS

New agents in advanced stages of development include cangrelor, a parenteral, rapidly acting, reversible inhibitor of P2Y₁₂, and vorapaxar, an orally active inhibitor of protease-activated receptor 1 (PAR-1), the major thrombin receptor on platelets (Fig. 143-3).

Cangrelor An adenosine analogue, cangrelor binds reversibly to P2Y₁₂ and inhibits its activity. The drug has a half-life of 3–6 min and is given IV as a bolus followed by an infusion. When stopped, platelet function recovers within 60 min. A trial comparing cangrelor with placebo during percutaneous coronary interventions and a study comparing cangrelor with clopidogrel after such procedures revealed little or no advantage of cangrelor. A third trial compared cangrelor (given as an IV bolus of 30 $\mu\text{g}/\text{kg}$ followed by an infusion of 4 $\mu\text{g}/\text{kg}$ per minute for at least 2 h, or for the duration of the procedure, whichever was longer) with a loading dose of clopidogrel (300 or 600 mg) in 11,145 patients undergoing urgent or elective percutaneous coronary intervention. The rate of the primary efficacy endpoint, a composite of death, MI, ischemia-driven revascularization, and stent thrombosis, was 4.7% in the cangrelor group and 5.9% in the clopidogrel group ($p = .005$). The rates of severe bleeding, the primary safety endpoint, were 0.16% and 0.11% in the cangrelor and clopidogrel groups, respectively. Using the same efficacy endpoint, a prespecified meta-analysis of the three trials

revealed a relative risk reduction of 19% with cangrelor compared with clopidogrel (3.8% and 4.7%, respectively) and a 40% reduction in stent thrombosis (0.5% and 0.8%, respectively) with no significant increase in serious bleeding. Based on these data, cangrelor is currently under regulatory review.

Vorapaxar An orally active PAR-1 antagonist, vorapaxar is slowly eliminated with a half-life of about 200 h. When compared with placebo in 12,944 patients with acute coronary syndrome without ST-segment elevation, vorapaxar failed to significantly reduce the primary efficacy endpoint, a composite of cardiovascular death, MI, stroke, recurrent ischemia requiring rehospitalization, and urgent coronary revascularization. Moreover, vorapaxar was associated with increased rates of bleeding, including intracranial bleeding.

In a second trial, vorapaxar was compared with placebo for secondary prevention in 26,449 patients with prior MI, ischemic stroke, or peripheral arterial disease. Overall, vorapaxar reduced the risk for cardiovascular death, MI, or stroke by 13%, but doubled the risk of intracranial bleeding. In the prespecified subgroup of 17,779 patients with prior MI, however, vorapaxar reduced the risk for cardiovascular death, MI, or stroke by 20% compared with placebo (from 9.7% to 8.1%, respectively). The rate of intracranial hemorrhage was higher with vorapaxar than with placebo (0.6% and 0.4%, respectively; $p = .076$) as was the rate of moderate or severe bleeding (3.4% and 2.1%, respectively; $P < 0.0001$). Based on these data, the drug is under consideration for regulatory approval in MI patients under the age of 75 years who have no history of stroke or transient ischemic attack and have a weight over 60 kg.

ANTICOAGULANTS

There are both parenteral and oral anticoagulants. The parenteral anticoagulants include heparin, low-molecular-weight heparin (LMWH), fondaparinux (a synthetic pentasaccharide), lepirudin, desirudin, bivalirudin, and argatroban. Currently available oral anticoagulants include warfarin; dabigatran etexilate, an oral thrombin inhibitor; and rivaroxaban and apixaban, oral factor Xa inhibitors. Edoxaban, a third oral factor Xa inhibitor, is undergoing regulatory review.

PARENTERAL ANTICOAGULANTS

Heparin A sulfated polysaccharide, heparin is isolated from mammalian tissues rich in mast cells. Most commercial heparin is derived from porcine intestinal mucosa and is a polymer of alternating D-glucuronic acid and N-acetyl-D-glucosamine residues.

MECHANISM OF ACTION Heparin acts as an anticoagulant by activating antithrombin (previously known as antithrombin III) and accelerating the rate at which antithrombin inhibits clotting enzymes, particularly thrombin and factor Xa. Antithrombin, the obligatory plasma cofactor for heparin, is a member of the serine protease inhibitor (serpin) superfamily. Synthesized in the liver and circulating in plasma at a concentration of $2.6 \pm 0.4 \mu\text{M}$, antithrombin acts as a suicide substrate for its target enzymes.

To activate antithrombin, heparin binds to the serpin via a unique pentasaccharide sequence that is found on one-third of the chains of commercial heparin (Fig. 143-5). Heparin chains without this pentasaccharide sequence have little or no anticoagulant activity. Once bound to antithrombin, heparin induces a conformational change in the reactive center loop of antithrombin that renders it more readily accessible to its target proteases. This conformational change enhances the rate at which antithrombin inhibits factor Xa by at least two orders of magnitude but has little effect on the rate of thrombin inhibition. To catalyze thrombin inhibition, heparin serves as a template that binds antithrombin and thrombin simultaneously. Formation of this ternary complex brings the enzyme in close apposition to the inhibitor, thereby

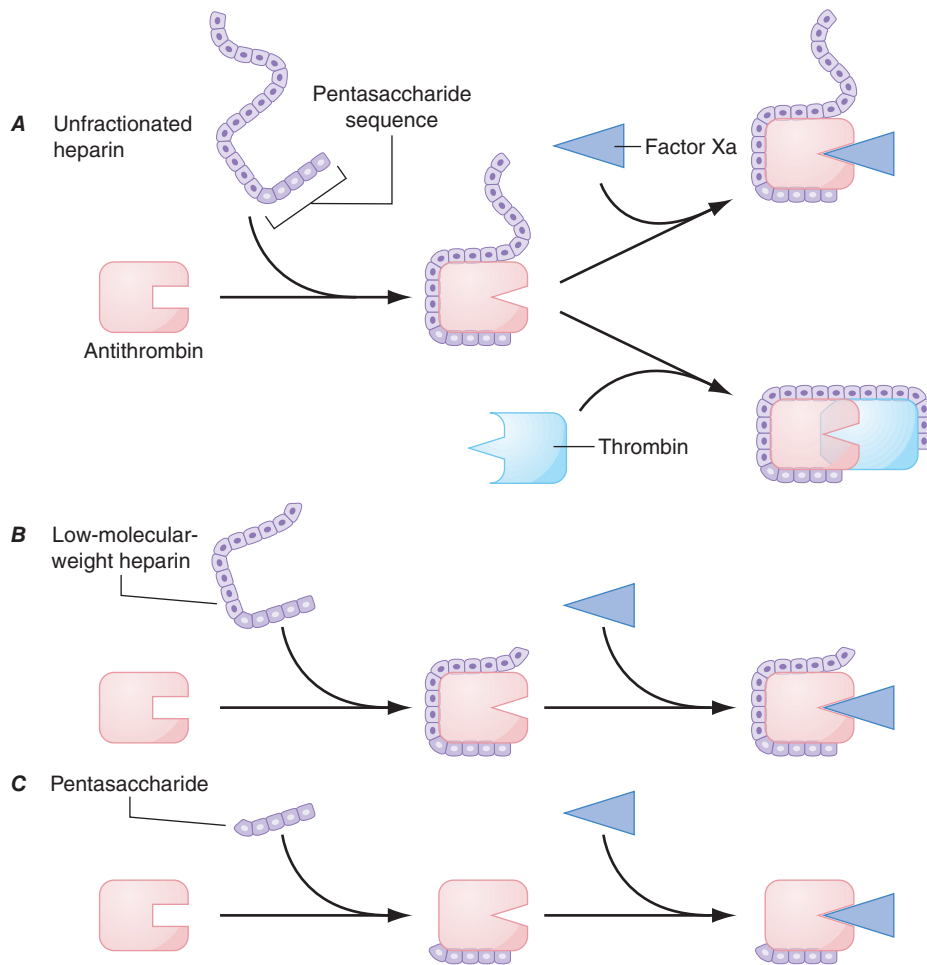


FIGURE 143-5 Mechanism of action of heparin, low-molecular-weight heparin (LMWH), and fondaparinux, a synthetic pentasaccharide. **A.** Heparin binds to antithrombin via its pentasaccharide sequence. This induces a conformational change in the reactive center loop of antithrombin that accelerates its interaction with factor Xa. To potentiate thrombin inhibition, heparin must simultaneously bind to antithrombin and thrombin. Only heparin chains composed of at least 18 saccharide units, which corresponds to a molecular weight of 5400, are of sufficient length to perform this bridging function. With a mean molecular weight of 15,000, all of the heparin chains are long enough to do this. **B.** LMWH has greater capacity to potentiate factor Xa inhibition by antithrombin than thrombin because, with a mean molecular weight of 4500–5000, at least half of the LMWH chains are too short to bridge antithrombin to thrombin. **C.** The pentasaccharide only accelerates factor Xa inhibition by antithrombin because the pentasaccharide is too short to bridge antithrombin to thrombin.

promoting the formation of a stable covalent thrombin-antithrombin complex.

Only pentasaccharide-containing heparin chains composed of at least 18 saccharide units (which correspond to a molecular weight of 5400) are of sufficient length to bridge thrombin and antithrombin together. With a mean molecular weight of 15,000, and a range of 5000–30,000, almost all of the chains of unfractionated heparin are long enough to do so. Consequently, by definition, heparin has equal capacity to promote the inhibition of thrombin and factor Xa by antithrombin and is assigned an anti-factor Xa to anti-factor IIa (thrombin) ratio of 1:1.

Heparin causes the release of tissue factor pathway inhibitor (TFPI) from the endothelium. A factor Xa-dependent inhibitor of tissue factor-bound factor VIIa, TFPI may contribute to the antithrombotic activity of heparin. Longer heparin chains induce the release of more TFPI than shorter ones.

PHARMACOLOGY Heparin must be given parenterally. It is usually administered SC or by continuous IV infusion. When used for therapeutic purposes, the IV route is most often employed. If heparin is given SC for treatment of thrombosis, the dose of heparin must be

high enough to overcome the limited bioavailability associated with this method of delivery.

In the circulation, heparin binds to the endothelium and to plasma proteins other than antithrombin. Heparin binding to endothelial cells explains its dose-dependent clearance. At low doses, the half-life of heparin is short because it binds rapidly to the endothelium. With higher doses of heparin, the half-life is longer because heparin is cleared more slowly once the endothelium is saturated. Clearance is mainly extrarenal; heparin binds to macrophages, which internalize and depolymerize the long heparin chains and secrete shorter chains back into the circulation. Because of its dose-dependent clearance mechanism, the plasma half-life of heparin ranges from 30 to 60 min with bolus IV doses of 25 and 100 units/kg, respectively.

Once heparin enters the circulation, it binds to plasma proteins other than antithrombin, a phenomenon that reduces its anticoagulant activity. Some of the heparin-binding proteins found in plasma are acute-phase reactants whose levels are elevated in ill patients. Others, such as high-molecular-weight multimers of VWF, are released from activated platelets or endothelial cells. Activated platelets also release platelet factor 4 (PF4), a highly cationic protein that binds heparin with high affinity. The large amounts of PF4 found in the vicinity of platelet-rich arterial thrombi can neutralize the anticoagulant activity of heparin. This phenomenon may attenuate heparin's capacity to suppress thrombus growth.

Because the levels of heparin-binding proteins in plasma vary from person to person, the anticoagulant response to fixed or weight-adjusted doses of heparin is unpredictable. Consequently, coagulation monitoring is essential to ensure that a therapeutic response is obtained. This is particularly important when heparin is administered for treatment of established thrombosis because a subtherapeutic anticoagulant response

may render patients at risk for recurrent thrombosis, whereas excessive anticoagulation increases the risk of bleeding.

MONITORING THE ANTICOAGULANT EFFECT Heparin therapy can be monitored using the activated partial thromboplastin time (aPTT) or anti-factor Xa level. Although the aPTT is the test most often used for this purpose, there are problems with this assay. aPTT reagents vary in their sensitivity to heparin, and the type of coagulometer used for testing can influence the results. Consequently, laboratories must establish a therapeutic aPTT range with each reagent-coagulometer combination by measuring the aPTT and anti-factor Xa level in plasma samples collected from heparin-treated patients. For most of the aPTT reagents and coagulometers in current use, therapeutic heparin levels are achieved with a two- to threefold prolongation of the aPTT.

Anti-factor Xa levels also can be used to monitor heparin therapy. With this test, therapeutic heparin levels range from 0.3 to 0.7 units/mL. Although this test is gaining in popularity, anti-factor Xa assays have yet to be standardized, and results can vary widely between laboratories.

Up to 25% of heparin-treated patients with venous thromboembolism require >35,000 units/d to achieve a therapeutic aPTT. These

patients are considered heparin resistant. It is useful to measure anti-factor Xa levels in heparin-resistant patients because many will have a therapeutic anti-factor Xa level despite a subtherapeutic aPTT. This dissociation in test results occurs because elevated plasma levels of fibrinogen and factor VIII, both of which are acute-phase proteins, shorten the aPTT but have no effect on anti-factor Xa levels. Heparin therapy in patients who exhibit this phenomenon is best monitored using anti-factor Xa levels instead of the aPTT. Patients with congenital or acquired antithrombin deficiency and those with elevated levels of heparin-binding proteins may also need high doses of heparin to achieve a therapeutic aPTT or anti-factor Xa level. If there is good correlation between the aPTT and the anti-factor Xa levels, either test can be used to monitor heparin therapy.

DOSING For prophylaxis, heparin is usually given in fixed doses of 5000 units SC two or three times daily. With these low doses, coagulation monitoring is unnecessary. In contrast, monitoring is essential when the drug is given in therapeutic doses. Fixed-dose or weight-based heparin nomograms are used to standardize heparin dosing and to shorten the time required to achieve a therapeutic anticoagulant response. At least two heparin nomograms have been validated in patients with venous thromboembolism and reduce the time required to achieve a therapeutic aPTT. Weight-adjusted heparin nomograms have also been evaluated in patients with acute coronary syndromes. After an IV heparin bolus of 5000 units or 70 units/kg, a heparin infusion rate of 12–15 units/kg per hour is usually administered. In contrast, weight-adjusted heparin nomograms for patients with venous thromboembolism use an initial bolus of 5000 units or 80 units/kg, followed by an infusion of 18 units/kg per h. Thus, patients with venous thromboembolism appear to require higher doses of heparin to achieve a therapeutic aPTT than do patients with acute coronary syndromes. This may reflect differences in the thrombus burden. Heparin binds to fibrin, and the amount of fibrin in patients with extensive DVT is greater than that in those with coronary thrombosis.

Heparin manufacturers in North America have traditionally measured heparin potency in USP units, with a unit defined as the concentration of heparin that prevents 1 mL of citrated sheep plasma from clotting for 1 h after calcium addition. In contrast, manufacturers in Europe measure heparin potency with anti-Xa assays using an international heparin standard for comparison. Because of problems with heparin contamination with oversulfated chondroitin sulfate, which the USP assay system does not detect, North American heparin manufacturers now use the anti-Xa assay to assess heparin potency. The use of international units in place of USP units results in a 10% reduction in heparin doses, which is a difference unlikely to affect patient care because monitoring will help to ensure that a therapeutic anticoagulant response has been achieved.

LIMITATIONS Heparin has pharmacokinetic and biophysical limitations (Table 143-2). The pharmacokinetic limitations reflect heparin's propensity to bind in a pentasaccharide-independent fashion to cells and plasma proteins. Heparin binding to endothelial cells explains its dose-dependent clearance, whereas binding to plasma proteins results in a variable anticoagulant response and can lead to heparin resistance.

TABLE 143-2 PHARMACOKINETIC AND BIOPHYSICAL LIMITATIONS OF HEPARIN

Limitations	Mechanism
Poor bioavailability at low doses	Binds to endothelial cells and macrophages
Dose-dependent clearance	Binds to macrophages
Variable anticoagulant response	Binds to plasma proteins whose levels vary from patient to patient
Reduced activity in the vicinity of platelet-rich thrombi	Neutralized by platelet factor 4 released from activated platelets
Limited activity against factor Xa incorporated in the prothrombinase complex and thrombin bound to fibrin	Reduced capacity of heparin-antithrombin complex to inhibit factor Xa bound to activated platelets and thrombin bound to fibrin

The biophysical limitations of heparin reflect the inability of the heparin-antithrombin complex to inhibit factor Xa when it is incorporated into the prothrombinase complex, the complex that converts prothrombin to thrombin, and to inhibit thrombin bound to fibrin. Consequently, factor Xa bound to activated platelets within platelet-rich thrombi has the potential to generate thrombin, even in the face of heparin. Once this thrombin binds to fibrin, it too is protected from inhibition by the heparin-antithrombin complex. Clot-associated thrombin can then trigger thrombus growth by locally activating platelets and amplifying its own generation through feedback activation of factors V, VIII, and XI. Further compounding the problem is the potential for heparin neutralization by the high concentrations of PF4 released from activated platelets within the platelet-rich thrombus.

SIDE EFFECTS The most common side effect of heparin is bleeding. Other complications include thrombocytopenia, osteoporosis, and elevated levels of transaminases.

Bleeding The risk of bleeding rises as the dose of heparin is increased. Concomitant administration of drugs that affect hemostasis, such as antiplatelet or fibrinolytic agents, increases the risk of bleeding, as does recent surgery or trauma. Heparin-treated patients with serious bleeding can be given protamine sulfate to neutralize the heparin. Protamine sulfate, a mixture of basic polypeptides isolated from salmon sperm, binds heparin with high affinity, and the resultant protamine-heparin complexes are then cleared. Typically, 1 mg of protamine sulfate neutralizes 100 units of heparin. Protamine sulfate is given IV. Anaphylactoid reactions to protamine sulfate can occur, and drug administration by slow IV infusion is recommended to reduce the risk.

THROMBOCYTOPENIA Heparin can cause thrombocytopenia. Heparin-induced thrombocytopenia (HIT) is an antibody-mediated process that is triggered by antibodies directed against neoantigens on PF4 that are exposed when heparin binds to this protein. These antibodies, which are usually of the IgG isotype, bind simultaneously to the heparin-PF4 complex and to platelet Fc receptors. Such binding activates the platelets and generates platelet microparticles. Circulating microparticles are prothrombotic because they express anionic phospholipids on their surface and can bind clotting factors and promote thrombin generation.

The clinical features of HIT are illustrated in Table 143-3. Typically, HIT occurs 5–14 days after initiation of heparin therapy, but it can manifest earlier if the patient has received heparin within the past 3 months. A platelet count below 100,000/ μ L or a 50% decrease in the platelet count from the pretreatment value should raise the suspicion of HIT in those receiving heparin. HIT is more common in surgical patients than in medical patients and, like many autoimmune disorders, occurs more frequently in females than in males.

HIT can be associated with thrombosis, either arterial or venous. Venous thrombosis, which manifests as DVT and/or PE, is more common than arterial thrombosis. Arterial thrombosis can manifest as ischemic stroke or acute MI. Rarely, platelet-rich thrombi in the distal aorta or iliac arteries can cause critical limb ischemia.

TABLE 143-3 FEATURES OF HEPARIN-INDUCED THROMBOCYTOPENIA

Features	Details
Thrombocytopenia	Platelet count of \leq 100,000/ μ L or a decrease in platelet count of \geq 50%
Timing	Platelet count falls 5–10 days after starting heparin
Type of heparin	More common with unfractionated heparin than low-molecular-weight heparin
Type of patient	More common in surgical patients and patients with cancer than general medical patients; more common in women than in men
Thrombosis	Venous thrombosis more common than arterial thrombosis

TABLE 143-4 MANAGEMENT OF HEPARIN-INDUCED THROMBOCYTOPENIA

Stop all heparin.
 Give an alternative anticoagulant, such as lepirudin, argatroban, bivalirudin, or fondaparinux.
 Do not give platelet transfusions.
 Do not give warfarin until the platelet count returns to its baseline level. If warfarin was administered, give vitamin K to restore the INR to normal.
 Evaluate for thrombosis, particularly deep vein thrombosis.

Abbreviation: INR, international normalized ratio.

The diagnosis of HIT is established using enzyme-linked assays to detect antibodies against heparin-PF4 complexes or with platelet activation assays. Enzyme-linked assays are sensitive but can be positive in the absence of any clinical evidence of HIT. The most specific diagnostic test is the serotonin release assay. This test is performed by quantifying serotonin release when washed platelets loaded with labeled serotonin are exposed to patient serum in the absence or presence of varying concentrations of heparin. If the patient serum contains the HIT antibody, heparin addition induces platelet activation and serotonin release.

Management of HIT is outlined in **Table 143-4**. Heparin should be stopped in patients with suspected or documented HIT, and an alternative anticoagulant should be administered to prevent or treat thrombosis. The agents most often used for this indication are parenteral direct thrombin inhibitors, such as lepirudin, argatroban, or bivalirudin, or factor Xa inhibitors, such as fondaparinux.

Patients with HIT, particularly those with associated thrombosis, often have evidence of increased thrombin generation that can lead to consumption of protein C. If these patients are given warfarin without a concomitant parenteral anticoagulant to inhibit thrombin or thrombin generation, the further decrease in protein C levels induced by the vitamin K antagonist can trigger skin necrosis. To avoid this problem, patients with HIT should be treated with a direct thrombin inhibitor or fondaparinux until the platelet count returns to normal levels. At this point, low-dose warfarin therapy can be introduced, and the thrombin inhibitor can be discontinued when the anticoagulant response to warfarin has been therapeutic for at least 2 days.

Osteoporosis Treatment with therapeutic doses of heparin for >1 month can cause a reduction in bone density. This complication has been reported in up to 30% of patients given long-term heparin therapy, and symptomatic vertebral fractures occur in 2–3% of these individuals.

Heparin causes bone loss both by decreasing bone formation and by enhancing bone resorption. Thus, heparin affects the activity of both osteoblasts and osteoclasts.

Elevated levels of transaminases Therapeutic doses of heparin are frequently associated with modest elevations in the serum levels of hepatic transaminases without a concomitant increase in the level of bilirubin. The levels of transaminases rapidly return to normal when the drug is stopped. The mechanism responsible for this phenomenon is unknown.

Low-Molecular-Weight Heparin Consisting of smaller fragments of heparin, LMWH is prepared from unfractionated heparin by controlled enzymatic or chemical depolymerization. The mean molecular weight of LMWH is about 5000, one-third the mean molecular weight of unfractionated heparin. LMWH has advantages over heparin (**Table 143-5**) and has replaced heparin for most indications.

MECHANISM OF ACTION Like heparin, LMWH exerts its anticoagulant activity by activating antithrombin. With a mean molecular weight of 5000, which corresponds to about 17 saccharide units, at least half of the pentasaccharide-containing chains of LMWH are too short to bridge thrombin to antithrombin (Fig. 143-5). However, these chains retain the capacity to accelerate factor Xa inhibition by antithrombin because this activity is largely the result of the conformational changes in antithrombin evoked by pentasaccharide binding. Consequently, LMWH catalyzes factor Xa inhibition by antithrombin more than

TABLE 143-5 ADVANTAGES OF LMWH OVER HEPARIN

Advantage	Consequence
Better bioavailability and longer half-life after subcutaneous injection	Can be given subcutaneously once or twice daily for both prophylaxis and treatment
Dose-independent clearance	Simplified dosing
Predictable anticoagulant response	Coagulation monitoring is unnecessary in most patients
Lower risk of heparin-induced thrombocytopenia	Safer than heparin for short- or long-term administration
Lower risk of osteoporosis	Safer than heparin for extended administration

Abbreviation: LMWH, low-molecular-weight heparin.

thrombin inhibition. Depending on their unique molecular weight distributions, LMWH preparations have anti-factor Xa to anti-factor IIa ratios ranging from 2:1 to 4:1.

PHARMACOLOGY Although usually given SC, LMWH also can be administered IV if a rapid anticoagulant response is needed. LMWH has pharmacokinetic advantages over heparin. These advantages reflect the fact that shorter heparin chains bind less avidly to endothelial cells, macrophages, and heparin-binding plasma proteins. Reduced binding to endothelial cells and macrophages eliminates the rapid, dose-dependent, and saturable mechanism of clearance that is a characteristic of unfractionated heparin. Instead, the clearance of LMWH is dose-independent and its plasma half-life is longer. Based on measurement of anti-factor Xa levels, LMWH has a plasma half-life of ~4 h. LMWH is cleared almost exclusively by the kidneys, and the drug can accumulate in patients with renal insufficiency.

LMWH exhibits about 90% bioavailability after SC injection. Because LMWH binds less avidly to heparin-binding proteins in plasma than heparin, LMWH produces a more predictable dose response, and resistance to LMWH is rare. With a longer half-life and more predictable anticoagulant response, LMWH can be given SC once or twice daily without coagulation monitoring, even when the drug is given in treatment doses. These properties render LMWH more convenient than unfractionated heparin. Capitalizing on this feature, studies in patients with venous thromboembolism have shown that home treatment with LMWH is as effective and safe as in-hospital treatment with continuous IV infusions of heparin. Outpatient treatment with LMWH streamlines care, reduces health care costs, and increases patient satisfaction.

MONITORING In the majority of patients, LMWH does not require coagulation monitoring. If monitoring is necessary, anti-factor Xa levels must be measured because most LMWH preparations have little effect on the aPTT. Therapeutic anti-factor Xa levels with LMWH range from 0.5 to 1.2 units/mL when measured 3–4 h after drug administration. When LMWH is given in prophylactic doses, peak anti-factor Xa levels of 0.2–0.5 units/mL are desirable.

Indications for LMWH monitoring include renal insufficiency and obesity. LMWH monitoring in patients with a creatinine clearance of ≤50 mL/min is advisable to ensure that there is no drug accumulation. Although weight-adjusted LMWH dosing appears to produce therapeutic anti-factor Xa levels in patients who are overweight, this approach has not been extensively evaluated in those with morbid obesity. It may also be advisable to monitor the anticoagulant activity of LMWH during pregnancy because dose requirements can change, particularly in the third trimester. Monitoring should also be considered in high-risk settings, such as in patients with mechanical heart valves who are given LMWH for prevention of valve thrombosis, and when LMWH is used in treatment doses in infants or children.

DOSING The doses of LMWH recommended for prophylaxis or treatment vary depending on the LMWH preparation. For prophylaxis, once-daily SC doses of 4000–5000 units are often used, whereas doses of 2500–3000 units are given when the drug is administered twice daily. For treatment of venous thromboembolism, a dose of

150–200 units/kg is given if the drug is administered once daily. If a twice-daily regimen is used, a dose of 100 units/kg is given. In patients with unstable angina, LMWH is given SC on a twice-daily basis at a dose of 100–120 units/kg.

SIDE EFFECTS The major complication of LMWH is bleeding. Meta-analyses suggest that the risk of major bleeding is lower with LMWH than with unfractionated heparin. HIT and osteoporosis are less common with LMWH than with unfractionated heparin.

Bleeding Like the situation with heparin, bleeding with LMWH is more common in patients receiving concomitant therapy with antiplatelet or fibrinolytic drugs. Recent surgery, trauma, or underlying hemostatic defects also increase the risk of bleeding with LMWH.

Although protamine sulfate can be used as an antidote for LMWH, protamine sulfate incompletely neutralizes the anticoagulant activity of LMWH because it only binds the longer chains of LMWH. Because longer chains are responsible for catalysis of thrombin inhibition by antithrombin, protamine sulfate completely reverses the anti-factor IIa activity of LMWH. In contrast, protamine sulfate only partially reverses the anti-factor Xa activity of LMWH because the shorter pentasaccharide-containing chains of LMWH do not bind to protamine sulfate. Consequently, patients at high risk for bleeding may be more safely treated with continuous IV unfractionated heparin than with SC LMWH.

Thrombocytopenia The risk of HIT is about fivefold lower with LMWH than with heparin. LMWH binds less avidly to platelets and causes less PF4 release. Furthermore, with lower affinity for PF4 than heparin, LMWH is less likely to induce the conformational changes in PF4 that trigger the formation of HIT antibodies.

LMWH should not be used to treat HIT patients because most HIT antibodies exhibit cross-reactivity with LMWH. This *in vitro* cross-reactivity is not simply a laboratory phenomenon because there are case reports of thrombosis when HIT patients were switched from heparin to LMWH.

Osteoporosis Because the risk of osteoporosis is lower with LMWH than with heparin, LMWH is the better choice for extended treatment.

Fondaparinux A synthetic analogue of the antithrombin-binding pentasaccharide sequence, fondaparinux differs from LMWH in several ways (Table 143-6). Fondaparinux is licensed for thromboprophylaxis in general medical or surgical patients and in high-risk orthopedic patients and as an alternative to heparin or LMWH for initial treatment of patients with established venous thromboembolism. Although widely used in Europe, as an alternative to heparin or LMWH in patients with acute coronary syndromes, fondaparinux is not licensed for this indication in the United States.

MECHANISM OF ACTION As a synthetic analogue of the antithrombin-binding pentasaccharide sequence found in heparin and LMWH, fondaparinux has a molecular weight of 1728. Fondaparinux binds only to antithrombin (Fig. 143-5) and is too short to bridge thrombin to antithrombin. Consequently, fondaparinux catalyzes factor Xa inhibition by antithrombin and does not enhance the rate of thrombin inhibition.

TABLE 143-6 COMPARISON OF LMWH AND FONDAPARINUX

Features	LMWH	Fondaparinux
Number of saccharide units	15–17	5
Catalysis of factor Xa inhibition	Yes	Yes
Catalysis of thrombin inhibition	Yes	No
Bioavailability after subcutaneous administration (%)	90	100
Plasma half-life (h)	4	17
Renal excretion	Yes	Yes
Induces release of tissue factor pathway inhibitor	Yes	No
Neutralized by protamine sulfate	Partially	No

PHARMACOLOGY Fondaparinux exhibits complete bioavailability after SC injection. With no binding to endothelial cells or plasma proteins, the clearance of fondaparinux is dose independent and its plasma half-life is 17 h. The drug is given SC once daily. Because fondaparinux is cleared unchanged via the kidneys, it is contraindicated in patients with a creatinine clearance <30 mL/min and should be used with caution in those with a creatinine clearance <50 mL/min.

Fondaparinux produces a predictable anticoagulant response after administration in fixed doses because it does not bind to plasma proteins. The drug is given at a dose of 2.5 mg once daily for prevention of venous thromboembolism. For initial treatment of established venous thromboembolism, fondaparinux is given at a dose of 7.5 mg once daily. The dose can be reduced to 5 mg once daily for those weighing <50 kg and increased to 10 mg for those >100 kg. When given in these doses, fondaparinux is as effective as heparin or LMWH for initial treatment of patients with DVT or PE and produces similar rates of bleeding.

Fondaparinux is used at a dose of 2.5 mg once daily in patients with acute coronary syndromes. When this prophylactic dose of fondaparinux was compared with treatment doses of enoxaparin in patients with non-ST-segment elevation acute coronary syndrome, there was no difference in the rate of cardiovascular death, MI, or stroke at 9 days. However, the rate of major bleeding was 50% lower with fondaparinux than with enoxaparin, a difference that likely reflects the fact that the dose of fondaparinux was lower than that of enoxaparin. In acute coronary syndrome patients who require percutaneous coronary intervention, there is a risk of catheter thrombosis with fondaparinux unless adjunctive heparin is given.

SIDE EFFECTS Fondaparinux does not cause HIT because it does not bind to PF4. In contrast to LMWH, there is no cross-reactivity of fondaparinux with HIT antibodies. Consequently, fondaparinux appears to be effective for treatment of HIT patients, although large clinical trials supporting its use are lacking.

The major side effect of fondaparinux is bleeding. There is no antidote for fondaparinux. Protamine sulfate has no effect on the anticoagulant activity of fondaparinux because it fails to bind to the drug. Recombinant activated factor VII reverses the anticoagulant effects of fondaparinux in volunteers, but it is unknown whether this agent controls fondaparinux-induced bleeding.

Parenteral Direct Thrombin Inhibitors Direct thrombin inhibitors bind directly to thrombin and block its interaction with its substrates. Approved parenteral direct thrombin inhibitors include recombinant hirudins (lepirudin and desirudin), argatroban, and bivalirudin (Table 143-7). Lepirudin and argatroban are licensed for treatment of patients with HIT, desirudin is licensed for thromboprophylaxis after elective hip arthroplasty, and bivalirudin is approved as an alternative to heparin in patients undergoing percutaneous coronary intervention, including those with HIT.

LEPIRUDIN AND DESIRUDIN Recombinant forms of hirudin, lepirudin, and desirudin are bivalent direct thrombin inhibitors that interact with the active site and exosite 1, the substrate-binding site on thrombin. For rapid anticoagulation, lepirudin is given by continuous IV infusion, but the drug can be given SC. Lepirudin has a plasma half-life of 60 min after IV infusion and is cleared by the kidneys. Consequently,

TABLE 143-7 COMPARISON OF THE PROPERTIES OF LEPIRUDIN, BIVALIRUDIN, AND ARGATROBAN

	Lepirudin/ Desirudin	Bivalirudin	Argatroban
Molecular mass	7000	1980	527
Site(s) of interaction with thrombin	Active site and exosite 1	Active site and exosite 1	Active site
Renal clearance	Yes	No	No
Hepatic metabolism	No	No	Yes
Plasma half-life (min)	60 (IV) 120–180 (SC)	25	45

754 lepirudin accumulates in patients with renal insufficiency. For thromboprophylaxis, desirudin is given SC twice daily in fixed doses; the half-life of desirudin is 2–3 h after SC injection.

A high proportion of lepirudin-treated patients develop antibodies against the drug; antibody formation is rare with SC desirudin. Although lepirudin-directed antibodies rarely cause problems, in a small subset of patients, they can delay lepirudin clearance and enhance its anticoagulant activity. Serious bleeding has been reported in some of these patients.

Lepirudin is usually monitored using the aPTT, and the dose is adjusted to maintain an aPTT that is 1.5–2.5 times the control. The aPTT is not an ideal test for monitoring lepirudin therapy because the clotting time plateaus with higher drug concentrations. Although the clotting time with ecarin, a snake venom that converts prothrombin to meizothrombin, provides a better index of lepirudin dose than the aPTT, the ecarin clotting time has yet to be standardized. When used for thromboprophylaxis, desirudin does not require monitoring.

ARGATROBAN A univalent inhibitor that targets the active site of thrombin, argatroban is metabolized in the liver. Consequently, this drug must be used with caution in patients with hepatic insufficiency. Argatroban is not cleared via the kidneys, so this drug is safer than lepirudin for HIT patients with renal insufficiency.

Argatroban is administered by continuous IV infusion and has a plasma half-life of ~45 min. The aPTT is used to monitor its anticoagulant effect, and the dose is adjusted to achieve an aPTT 1.5–3 times the baseline value, but not to exceed 100 s. Argatroban also prolongs the international normalized ratio (INR), a feature that can complicate the transitioning of patients to warfarin. This problem can be circumvented by using the levels of factor X to monitor warfarin in place of the INR. Alternatively, argatroban can be stopped for 2–3 h before INR determination.

BIVALIRUDIN A synthetic 20-amino-acid analogue of hirudin, bivalirudin is a divalent thrombin inhibitor. Thus, the N-terminus of bivalirudin interacts with the active site of thrombin, whereas its C-terminus binds to exosite 1. Bivalirudin has a plasma half-life of 25 min, the shortest half-life of all the parenteral direct thrombin inhibitors. Bivalirudin is degraded by peptidases and is partially excreted via the kidneys. When given in high doses in the cardiac catheterization laboratory, the anticoagulant activity of bivalirudin is monitored using the activated clotting time. With lower doses, its activity can be assessed using the aPTT.

Bivalirudin is licensed as an alternative to heparin in patients undergoing percutaneous coronary intervention. Bivalirudin also has been used successfully in HIT patients who require percutaneous coronary intervention or cardiac bypass surgery.

ORAL ANTICOAGULANTS

Current oral anticoagulant practice dates back almost 60 years to when the vitamin K antagonists were discovered as a result of investigations into the cause of hemorrhagic disease in cattle. Characterized by a decrease in prothrombin levels, this disorder is caused by ingestion of hay containing spoiled sweet clover. Hydroxycoumarin, which was isolated from bacterial contaminants in the hay, interferes with vitamin K metabolism, thereby causing a syndrome similar to vitamin K deficiency. Discovery of this compound provided the impetus for development of other vitamin K antagonists, including warfarin.

For many years, the vitamin K antagonists were the only available oral anticoagulants. This situation changed with the introduction of new oral anticoagulants, including dabigatran, which targets thrombin, and rivaroxaban, apixaban, and edoxaban, which target factor Xa.

Warfarin A water-soluble vitamin K antagonist initially developed as a rodenticide, warfarin is the coumarin derivative most often prescribed in North America. Like other vitamin K antagonists, warfarin interferes with the synthesis of the vitamin K-dependent clotting proteins, which include prothrombin (factor II) and factors VII, IX, and X. The synthesis of the vitamin K-dependent anticoagulant proteins, proteins C and S, is also reduced by vitamin K antagonists.

MECHANISM OF ACTION All of the vitamin K-dependent clotting factors possess glutamic acid residues at their N termini. A posttranslational modification adds a carboxyl group to the γ -carbon of these residues to generate γ -carboxyglutamic acid. This modification is essential for expression of the activity of these clotting factors because it permits their calcium-binding to negatively charged phospholipid surfaces. The γ -carboxylation process is catalyzed by a vitamin K-dependent carboxylase. Thus, vitamin K from the diet is reduced to vitamin K hydroquinone by vitamin K reductase (Fig. 143-6). Vitamin K hydroquinone serves as a cofactor for the carboxylase enzyme, which in the presence of carbon dioxide replaces the hydrogen on the γ -carbon of glutamic acid residues with a carboxyl group. During this process, vitamin K hydroquinone is oxidized to vitamin K epoxide, which is then reduced to vitamin K by vitamin K epoxide reductase.

Warfarin inhibits vitamin K epoxide reductase (VKOR), thereby blocking the γ -carboxylation process. This results in the synthesis of vitamin K-dependent clotting proteins that are only partially γ -carboxylated. Warfarin acts as an anticoagulant because these partially γ -carboxylated proteins have reduced or absent biologic activity. The onset of action of warfarin is delayed until the newly synthesized clotting factors with reduced activity gradually replace their fully active counterparts.

The antithrombotic effect of warfarin depends on a reduction in the functional levels of factor X and prothrombin, clotting factors that have half-lives of 24 and 72 h, respectively. Because the antithrombotic effect of warfarin is delayed, patients with established thrombosis or at high risk for thrombosis require concomitant treatment with a rapidly acting parenteral anticoagulant, such as heparin, LMWH, or fondaparinux, for at least 5 days.

PHARMACOLOGY Warfarin is a racemic mixture of R and S isomers. Warfarin is rapidly and almost completely absorbed from the

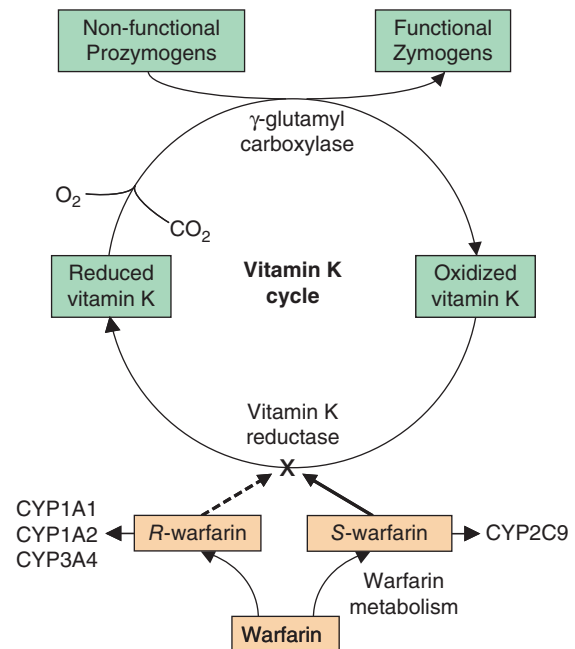


FIGURE 143-6 Mechanism of action of warfarin. A racemic mixture of S- and R-enantiomers, S-warfarin is most active. By blocking vitamin K epoxide reductase, warfarin inhibits the conversion of oxidized vitamin K into its reduced form. This inhibits vitamin K-dependent γ -carboxylation of factors II, VII, IX, and X because reduced vitamin K serves as a cofactor for a γ -glutamyl carboxylase that catalyzes the γ -carboxylation process, thereby converting prozymogens to zymogens capable of binding calcium and interacting with anionic phospholipid surfaces. S-warfarin is metabolized by CYP2C9. Common genetic polymorphisms in this enzyme can influence warfarin metabolism. Polymorphisms in the C1 subunit of vitamin K reductase (VKORC1) also can affect the susceptibility of the enzyme to warfarin-induced inhibition, thereby influencing warfarin dosage requirements.

TABLE 143-8 FREQUENCIES OF *CYP2C9* GENOTYPES AND *VKORC1* HAPLOTYPES IN DIFFERENT POPULATIONS AND THEIR EFFECT ON WARFARIN DOSE REQUIREMENTS

Genotype/Haplotype	Frequency, %			Dose Reduction Compared with Wild-Type
	Caucasians	African Americans (A/A)	Asians (A)	
<i>CYP2C9</i>				
*1/*1	70	90	95	–
*1/*2	17	2	0	22
*1/*3	9	3	4	34
*2/*2	2	0	0	43
*2/*3	1	0	0	53
*3/*3	0	0	1	76
<i>VKORC1</i>				
Non-A/non-A	37	82	7	–
Non-A/A	45	12	30	26
A/A	18	6	63	50

gastrointestinal tract. Levels of warfarin in the blood peak about 90 min after drug administration. Racemic warfarin has a plasma half-life of 36–42 h, and more than 97% of circulating warfarin is bound to albumin. Only the small fraction of unbound warfarin is biologically active.

Warfarin accumulates in the liver where the two isomers are metabolized via distinct pathways. *CYP2C9* mediates oxidative metabolism of the more active S isomer (Fig. 143-6). Two relatively common variants, *CYP2C9*2* and *CYP2C9*3*, encode an enzyme with reduced activity. Patients with these variants require lower maintenance doses of warfarin. Approximately 25% of Caucasians have at least one variant allele of *CYP2C9*2* or *CYP2C9*3*, whereas those variant alleles are less common in African Americans and Asians (Table 143-8). Heterozygosity for *CYP2C9*2* or *CYP2C9*3* decreases the warfarin dose requirement by 20–30% relative to that required in subjects with the wild-type *CYP2C9*1/1* alleles, whereas homozygosity for the *CYP2C9*2* or *CYP2C9*3* alleles reduces the warfarin dose requirement by 50–70%.

Consistent with their decreased warfarin dose requirement, subjects with at least one *CYP2C9* variant allele are at increased risk for bleeding. Compared with individuals with no variant alleles, the relative risks for warfarin-associated bleeding in *CYP2C9*2* or *CYP2C9*3* carriers are 1.9 and 1.8, respectively.

Polymorphisms in *VKORC1* also can influence the anticoagulant response to warfarin. Several genetic variations of *VKORC1* are in strong linkage disequilibrium and have been designated as non-A haplotypes. *VKORC1* variants are more prevalent than variants of *CYP2C9*. Asians have the highest prevalence of *VKORC1* variants, followed by Caucasians and African Americans (Table 143-8). Polymorphisms in *VKORC1* likely explain 30% of the variability in warfarin dose requirements. Compared with *VKORC1* non-A/non-A homozygotes, the warfarin dose requirement decreases by 25 and 50% in A haplotype heterozygotes and homozygotes, respectively. These findings prompted the Food and Drug Administration to amend the prescribing information for warfarin to indicate that lower initiation doses should be considered for patients with *CYP2C9* and *VKORC1* genetic variants. In addition to genotype data, other pertinent patient information has been incorporated into warfarin dosing algorithms. Although such algorithms help predict suitable warfarin doses, it remains unclear whether better dose identification improves patient outcome in terms of reducing hemorrhagic complications or recurrent thrombotic events.

In addition to genetic factors, the anticoagulant effect of warfarin is influenced by diet, drugs, and various disease states. Fluctuations in dietary vitamin K intake affect the activity of warfarin. A wide variety of drugs can alter absorption, clearance, or metabolism of warfarin. Because of the variability in the anticoagulant response to warfarin, coagulation monitoring is essential to ensure that a therapeutic response is obtained.

MONITORING Warfarin therapy is most often monitored using the prothrombin time, a test that is sensitive to reductions in the levels of prothrombin, factor VII, and factor X. The test is performed by adding thromboplastin, a reagent that contains tissue factor, phospholipid, and calcium, to citrated plasma and determining the time to clot formation. Thromboplastins vary in their sensitivity to reductions in the levels of the vitamin K–dependent clotting factors. Thus, less sensitive thromboplastins will trigger the administration of higher doses of warfarin to achieve a target prothrombin time. This is problematic because higher doses of warfarin increase the risk of bleeding.

The INR was developed to circumvent many of the problems associated with the prothrombin time. To calculate the INR, the patient's prothrombin time is divided by the mean normal prothrombin time, and this ratio is then multiplied by the international sensitivity index (ISI), which is an index of the sensitivity of the thromboplastin used for prothrombin time

determination to reductions in the levels of the vitamin K–dependent clotting factors. Highly sensitive thromboplastins have an ISI of 1.0. Most current thromboplastins have ISI values that range from 1.0 to 1.4.

Although the INR has helped to standardize anticoagulant practice, problems persist. The precision of INR determination varies depending on reagent-coagulometer combinations. This leads to variability in the INR results. Also complicating INR determination is unreliable reporting of the ISI by thromboplastin manufacturers. Furthermore, every laboratory must establish the mean normal prothrombin time with each new batch of thromboplastin reagent. To accomplish this, the prothrombin time must be measured in fresh plasma samples from at least 20 healthy volunteers using the same coagulometer that is used for patient samples.

For most indications, warfarin is administered in doses that produce a target INR of 2.0–3.0. An exception is patients with mechanical heart valves, particularly those in the mitral position or older ball and cage valves in the aortic position, where a target INR of 2.5–3.5 is recommended. Studies in atrial fibrillation demonstrate an increased risk of cardioembolic stroke when the INR falls to <1.7 and an increase in bleeding with INR values >4.5. These findings highlight the fact that vitamin K antagonists have a narrow therapeutic window. In support of this concept, a study in patients receiving long-term warfarin therapy for unprovoked venous thromboembolism demonstrated a higher rate of recurrent venous thromboembolism with a target INR of 1.5–1.9 compared with a target INR of 2.0–3.0.

DOSING Warfarin is usually started at a dose of 5–10 mg. Lower doses are used for patients with *CYP2C9* or *VKORC1* polymorphisms, which affect the pharmacodynamics or pharmacokinetics of warfarin and render patients more sensitive to the drug. The dose is then titrated to achieve the desired target INR. Because of its delayed onset of action, patients with established thrombosis or those at high risk for thrombosis are given concomitant initial treatment with a rapidly acting parenteral anticoagulant, such as heparin, LMWH, or fondaparinux. Early prolongation of the INR reflects reduction in the functional levels of factor VII. Consequently, concomitant treatment with the parenteral anticoagulant should be continued until the INR has been therapeutic for at least 2 consecutive days. A minimum 5-day course of parenteral anticoagulation is recommended to ensure that the levels of factor Xa and prothrombin have been reduced into the therapeutic range with warfarin.

Because warfarin has a narrow therapeutic window, frequent coagulation monitoring is essential to ensure that a therapeutic anticoagulant response is maintained. Even patients with stable warfarin dose requirements should have their INR determined every 3–4 weeks. More frequent monitoring is necessary when new medications are introduced because so many drugs enhance or reduce the anticoagulant effects of warfarin.

756 SIDE EFFECTS Like all anticoagulants, the major side effect of warfarin is bleeding. A rare complication is skin necrosis. Warfarin crosses the placenta and can cause fetal abnormalities. Consequently, warfarin should not be used during pregnancy.

Bleeding At least half of the bleeding complications with warfarin occur when the INR exceeds the therapeutic range. Bleeding complications may be mild, such as epistaxis or hematuria, or more severe, such as retroperitoneal or gastrointestinal bleeding. Life-threatening intracranial bleeding can also occur.

To minimize the risk of bleeding, the INR should be maintained in the therapeutic range. In asymptomatic patients whose INR is between 3.5 and 10, warfarin should be withheld until the INR returns to the therapeutic range. If the INR is over 10, oral vitamin K should be administered, at a dose of 2.5–5 mg, although there is no evidence that doing so reduces the bleeding risk. Higher doses of oral vitamin K (5–10 mg) produce more rapid reversal of the INR but may render patients temporarily resistant to warfarin when the drug is restarted. Patients with serious bleeding need more aggressive treatment. These patients should be given 5–10 mg of vitamin K by slow IV infusion. Additional vitamin K should be given until the INR is in the normal range. Treatment with vitamin K should be supplemented with fresh-frozen plasma as a source of the vitamin K–dependent clotting proteins. Four factor prothrombin complex concentrates, which contain all four vitamin K–dependent clotting proteins, are the treatment of choice for (1) life-threatening bleeds, (2) rapid restoration of the INR into the normal range in patients requiring urgent surgery or intervention, and (3) patients who cannot tolerate the volume load of fresh-frozen plasma.

Warfarin-treated patients who experience bleeding when their INR is in the therapeutic range require investigation into the cause of the bleeding. Those with gastrointestinal or genitourinary bleeding often have an underlying lesion.

Skin necrosis A rare complication of warfarin, skin necrosis usually is seen 2–5 days after initiation of therapy. Well-demarcated erythematous lesions form on the thighs, buttocks, breasts, or toes. Typically, the center of the lesion becomes progressively necrotic. Examination of skin biopsies taken from the border of these lesions reveals thrombi in the microvasculature.

Warfarin-induced skin necrosis is seen in patients with congenital or acquired deficiencies of protein C or protein S. Initiation of warfarin therapy in these patients produces a precipitous fall in plasma levels of proteins C or S, thereby eliminating this important anticoagulant pathway before warfarin exerts an antithrombotic effect through lowering of the functional levels of factor X and prothrombin. The resultant procoagulant state triggers thrombosis. Why the thrombosis is localized to the microvasculature of fatty tissues is unclear.

Treatment involves discontinuation of warfarin and reversal with vitamin K, if needed. An alternative anticoagulant, such as heparin or LMWH, should be given in patients with thrombosis. Protein C concentrate can be given to protein C–deficient patients to accelerate healing of the skin lesions; fresh-frozen plasma may be of value if protein C concentrate is unavailable and for those with protein S deficiency. Occasionally, skin grafting is necessary when there is extensive skin loss.

Because of the potential for skin necrosis, patients with known protein C or protein S deficiency require overlapping treatment with a parenteral anticoagulant when initiating warfarin therapy. Warfarin should be started in low doses in these patients, and the parenteral anticoagulant should be continued until the INR is therapeutic for at least 2–3 consecutive days.

Pregnancy Warfarin crosses the placenta and can cause fetal abnormalities or bleeding. The fetal abnormalities include a characteristic embryopathy, which consists of nasal hypoplasia and stippled epiphyses. The risk of embryopathy is highest if warfarin is given in the first

trimester of pregnancy. Central nervous system abnormalities can also occur with exposure to warfarin at any time during pregnancy. Finally, maternal administration of warfarin produces an anticoagulant effect in the fetus that can cause bleeding. This is of particular concern at delivery when trauma to the head during passage through the birth canal can lead to intracranial bleeding. Because of these potential problems, warfarin is contraindicated in pregnancy, particularly in the first and third trimesters. Instead, heparin, LMWH, or fondaparinux can be given during pregnancy for prevention or treatment of thrombosis.

Warfarin does not pass into the breast milk. Consequently, warfarin can safely be given to nursing mothers.

Special problems Patients with a lupus anticoagulant and those who need urgent or elective surgery present special challenges. Although observational studies suggested that patients with thrombosis complicating the antiphospholipid antibody syndrome required higher intensity warfarin regimens to prevent recurrent thromboembolic events, two randomized trials showed that targeting an INR of 2.0–3.0 is as effective as higher intensity treatment and produces less bleeding. Monitoring warfarin therapy can be problematic in patients with antiphospholipid antibody syndrome if the lupus anticoagulant prolongs the baseline INR; factor X levels can be used instead of the INR in such patients.

There is no need to stop warfarin before procedures associated with a low risk of bleeding; these include dental cleaning, simple dental extraction, cataract surgery, or skin biopsy. For procedures associated with a moderate or high risk of bleeding, warfarin should be stopped 5 days before the procedure to allow the INR to return to normal levels. Patients at high risk for thrombosis, such as those with mechanical heart valves, can be bridged with once- or twice-daily SC injections of LMWH when the INR falls to <2.0. The last dose of LMWH should be given 12–24 h before the procedure, depending on whether LMWH is administered twice or once daily. After the procedure, treatment with warfarin can be restarted.

New Oral Anticoagulants New oral anticoagulants are now available as alternatives to warfarin. These include dabigatran, which targets thrombin, and rivaroxaban, apixaban, and edoxaban, which target factor Xa. All of these drugs have a rapid onset and offset of action and have half-lives that permit once- or twice-daily administration. Designed to produce a predictable level of anticoagulation, the new oral agents are more convenient to administer than warfarin because they are given in fixed doses without routine coagulation monitoring.

MECHANISM OF ACTION The new oral anticoagulants are small molecules that bind reversibly to the active site of their target enzyme. [Table 143-9](#) summarizes the distinct pharmacologic properties of these agents.

INDICATIONS The new oral anticoagulants have been compared with warfarin for stroke prevention in patients with nonvalvular atrial fibrillation in four randomized trials that enrolled 71,683 patients. A meta-analysis of these data demonstrates that compared with warfarin, the new agents significantly reduce stroke or systemic embolism by 19% ($p = .001$), primarily driven by a 51% reduction in hemorrhagic stroke ($p < .0001$), and are associated with a 10% reduction in mortality ($p < .0001$). New oral anticoagulants reduce intracranial hemorrhage by

TABLE 143-9 COMPARISON OF THE PHARMACOLOGIC PROPERTIES OF THE NEW ORAL ANTICOAGULANTS

Characteristic	Rivaroxaban	Apixaban	Edoxaban	Dabigatran
Target	Factor Xa	Factor Xa	Factor Xa	Thrombin
Prodrug	No	No	No	Yes
Bioavailability	80%	60%	50%	6%
Dosing	qd (bid)	bid	qd	bid (qd)
Half-life	7–11 h	12 h	9–11 h	12–17 h
Renal	33% (66%)	25%	35%	80%
Monitoring	No	No	No	No
Interactions	3A4/P-gp	3A4/P-gp	P-gp	P-gp

Abbreviations: bid, twice a day; P-gp, P-glycoprotein; qd., once a day.

52% compared with warfarin ($p < .0001$), but increase gastrointestinal bleeding by about 24% ($p = .04$). Overall, the new agents demonstrate a favorable benefit-to-risk profile compared with warfarin, and their relative efficacy and safety are maintained across a wide spectrum of atrial fibrillation patients, including those over the age of 75 years and those with a prior history of stroke. Based on these findings, dabigatran, rivaroxaban, and apixaban are licensed as alternatives to warfarin for stroke prevention in nonvalvular atrial fibrillation, and edoxaban is under regulatory consideration for this indication. Nonvalvular atrial fibrillation is defined as that occurring in patients without mechanical heart valves or severe rheumatic valvular disease, particularly mitral stenosis and/or regurgitation.

Dabigatran, rivaroxaban, and apixaban have been compared with enoxaparin for thromboprophylaxis after elective hip or knee arthroplasty. Currently, only rivaroxaban and apixaban are licensed for this indication in the United States. Rivaroxaban and dabigatran are also licensed for treatment of DVT or PE. Apixaban and edoxaban have also been investigated for treatment of patients with venous thromboembolism, but have not yet been approved for this indication. Rivaroxaban is licensed in Europe for prevention of recurrent ischemic events in patients who have been stabilized after an acute coronary syndrome. In this setting, rivaroxaban is usually administered in conjunction with dual antiplatelet therapy with aspirin and clopidogrel.

DOSING For stroke prevention in patients with nonvalvular atrial fibrillation, rivaroxaban is given at a dose of 20 mg once daily with a dose reduction to 15 mg once daily in patients with a creatinine clearance of 15–49 mL/min; dabigatran is given at a dose of 150 mg twice daily with a dose reduction to 75 mg twice daily in those with a creatinine clearance of 15–30 mL/min; and apixaban is given at a dose of 5 mg twice daily with a dose reduction to 2.5 mg twice daily for patients with a creatinine >1.5 g/dL, for those 80 years of age or older, or for patients who weigh <60 kg.

For thromboprophylaxis after elective hip or knee replacement surgery, rivaroxaban is given at a dose of 10 mg once daily, whereas apixaban is given at a dose of 2.5 mg twice daily. For treatment of patients with DVT or PE, rivaroxaban is started at a dose of 15 mg twice daily for 3 weeks; the dose is then reduced to 20 mg once daily thereafter. After a minimum of a 5 day course of treatment with heparin or LMWH, dabigatran is given at a dose of 150 mg twice daily.

MONITORING Although designed to be administered without routine monitoring, there are situations where determination of the anticoagulant activity of the new oral anticoagulants can be helpful. These include assessment of adherence, detection of accumulation or overdose, identification of bleeding mechanisms, and determination of activity prior to surgery or intervention. For qualitative assessment of anticoagulant activity, the prothrombin time can be used for factor Xa inhibitors and the aPTT for dabigatran. Rivaroxaban and edoxaban prolong the prothrombin time more than apixaban. In fact, because apixaban has such a limited effect on the prothrombin time, anti-factor Xa assays are needed to assess its activity. The effect of the drugs on tests of coagulation varies depending on the time that the blood is drawn relative to the timing of the last dose of the drug and the reagents used to perform the tests. Chromogenic anti-factor Xa assays and a dilute thrombin clotting time with appropriate calibrators provide quantitative assays to measure the plasma levels of the factor Xa inhibitors and dabigatran, respectively.

SIDE EFFECTS Like all anticoagulants, bleeding is the most common side effect of the new oral anticoagulants. The new agents are associated with less intracranial bleeding than warfarin. The increased risk of intracranial bleeding with warfarin likely reflects the reduction in functional levels of factor VII, which precludes efficient thrombin generation at sites of microvascular bleeding in the brain. Because the new oral anticoagulants target downstream coagulation enzymes, they produce less impairment of hemostatic plug formation at sites of vascular injury.

A downside of the new oral anticoagulants is the increased risk of gastrointestinal bleeding. This likely occurs because unabsorbed active

drug in the gut exacerbates bleeding from lesions. Although dabigatran etexilate is a prodrug, only 7% is absorbed. Although the remainder passes through the gut, at least two-thirds is metabolically activated to dabigatran by gut esterases.

Dyspepsia occurs in up to 10% of patients treated with dabigatran; this problem improves with time and can be minimized by administering the drug with food. Dyspepsia is rare with rivaroxaban, apixaban, and edoxaban.

PERIPROCEDURAL MANAGEMENT Like warfarin, the new oral anticoagulants must be stopped before procedures associated with a moderate or high risk of bleeding. The drugs should be held for 1–2 days, or longer if renal function is impaired. Assessment of residual anticoagulant activity before procedures associated with a high bleeding risk is prudent.

MANAGEMENT OF BLEEDING There are no specific antidotes for the new oral anticoagulants. With minor bleeding, holding one or two doses of drug is usually sufficient. The approach to serious bleeding is similar to that with warfarin except that vitamin K administration is of no benefit. Thus, the anticoagulant and antiplatelet drugs should be held, the patient should be resuscitated with fluids and blood products as necessary, and, if possible, the bleeding site should be identified and managed. Coagulation testing will determine the extent of anticoagulation, and renal function should be assessed so that the half-life of the drug can be calculated. Timing of the last dose of anticoagulant is important; administration of oral activated charcoal may help to prevent absorption of drug administered in the past 2–4 h. If bleeding continues or is life-threatening, procoagulants, such as prothrombin complex concentrate (either unactivated or activated) or factor VIIa, can be administered, although the evidence of their effectiveness is limited. Dialysis removes dabigatran from the circulation in patients with renal impairment; dialysis does not remove rivaroxaban, apixaban, or edoxaban because unlike dabigatran, these drugs are highly protein-bound.

PREGNANCY As small molecules, the new oral anticoagulants can all pass through the placenta. Consequently, these agents are contraindicated in pregnancy, and when used by women of childbearing potential, appropriate contraception is important.

ONGOING INVESTIGATIONS Although the lack of antidotes has created concern about the risk of bleeding events in patients taking the new oral anticoagulants, emerging postmarketing data suggest that the rates of bleeding in the real-world setting are similar to those reported in the trials. Nonetheless, specific antidotes are under development. These include a humanized mouse monoclonal antibody fragment against dabigatran and a recombinant variant of factor Xa that serves as a decoy for the oral factor Xa inhibitors. Neither agent is currently available for clinical use.

FIBRINOLYTIC DRUGS

ROLE OF FIBRINOLYTIC THERAPY

Fibrinolytic drugs can be used to degrade thrombi and are administered systemically or can be delivered via catheters directly into the substance of the thrombus. Systemic delivery is used for treatment of acute MI, acute ischemic stroke, and most cases of massive PE. The goal of therapy is to produce rapid thrombus dissolution, thereby restoring antegrade blood flow. In the coronary circulation, restoration of blood flow reduces morbidity and mortality rates by limiting myocardial damage, whereas in the cerebral circulation, rapid thrombus dissolution decreases the neuronal death and brain infarction that produce irreversible brain injury. For patients with massive PE, the goal of thrombolytic therapy is to restore pulmonary artery perfusion.

Peripheral arterial thrombi and thrombi in the proximal deep veins of the leg are most often treated using catheter-directed thrombolytic therapy. Catheters with multiple side holes can be used to enhance drug delivery. In some cases, intravascular devices that fragment and extract the thrombus are used to hasten treatment. These devices can be used alone or in conjunction with fibrinolytic drugs.

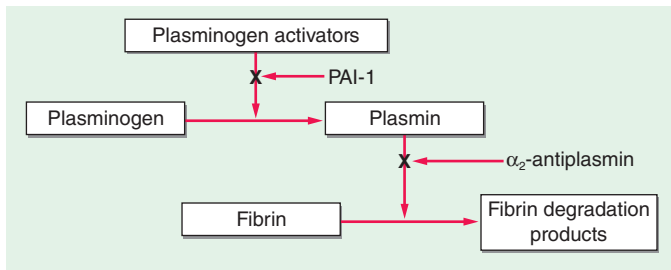


FIGURE 143-7 The fibrinolytic system and its regulation. Plasminogen activators convert plasminogen to plasmin. Plasmin then degrades fibrin into soluble fibrin degradation products. The system is regulated at two levels. Type 1 plasminogen activator inhibitor (PAI-1) regulates the plasminogen activators, whereas α_2 -antiplasmin serves as the major inhibitor of plasmin.

MECHANISM OF ACTION

Currently approved fibrinolytic agents include streptokinase; acylated plasminogen streptokinase activator complex (anistreplase); urokinase; recombinant tissue-type plasminogen activator (rtPA), which is also known as alteplase or activase; and two recombinant derivatives of rtPA, tenecteplase and reteplase. All of these agents act by converting plasminogen, the zymogen, to plasmin, the active enzyme (Fig. 143-7). Plasmin then degrades the fibrin matrix of thrombi and produces soluble fibrin degradation products.

Endogenous fibrinolysis is regulated at two levels. Plasminogen activator inhibitors, particularly the type 1 form (PAI-1), prevent excessive plasminogen activation by regulating the activity of tPA and urokinase-type plasminogen activator (uPA). Once plasmin is generated, it is regulated by plasmin inhibitors, the most important of which is α_2 -antiplasmin. The plasma concentration of plasminogen is twofold higher than that of α_2 -antiplasmin. Consequently, with pharmacologic doses of plasminogen activators, the concentration of plasmin that is generated can exceed that of α_2 -antiplasmin. In addition to degrading fibrin, unregulated plasmin can also degrade fibrinogen and other clotting factors. This process, which is known as the *systemic lytic state*, reduces the hemostatic potential of the blood and increases the risk of bleeding.

The endogenous fibrinolytic system is geared to localize plasmin generation to the fibrin surface. Both plasminogen and tPA bind to fibrin to form a ternary complex that promotes efficient plasminogen activation. In contrast to free plasmin, plasmin generated on the fibrin surface is relatively protected from inactivation by α_2 -antiplasmin, a feature that promotes fibrin dissolution. Furthermore, C-terminus lysine residues, exposed as plasmin degrades fibrin, serve as binding sites for additional plasminogen and tPA molecules. This creates a positive feedback that enhances plasmin generation. When used pharmacologically, the various plasminogen activators capitalize on these mechanisms to a lesser or greater extent.

Plasminogen activators that preferentially activate fibrin-bound plasminogen are considered fibrin-specific. In contrast, nonspecific plasminogen activators do not discriminate between fibrin-bound and circulating plasminogen. Activation of circulating plasminogen results in the generation of unopposed plasmin that can trigger the systemic lytic state. Alteplase and its derivatives are fibrin-specific plasminogen activators, whereas streptokinase, anistreplase, and urokinase are nonspecific agents.

STREPTOKINASE

Unlike other plasminogen activators, streptokinase is not an enzyme and does not directly convert plasminogen to plasmin. Instead, streptokinase forms a 1:1 stoichiometric complex with plasminogen. Formation of this complex induces a conformational change in plasminogen that exposes its active site (Fig. 143-8). The streptokinase-plasminogen complex then converts additional plasminogen to plasmin.

Streptokinase has no affinity for fibrin, and the streptokinase-plasminogen complex activates both free and fibrin-bound plasminogen. Activation of circulating plasminogen generates sufficient

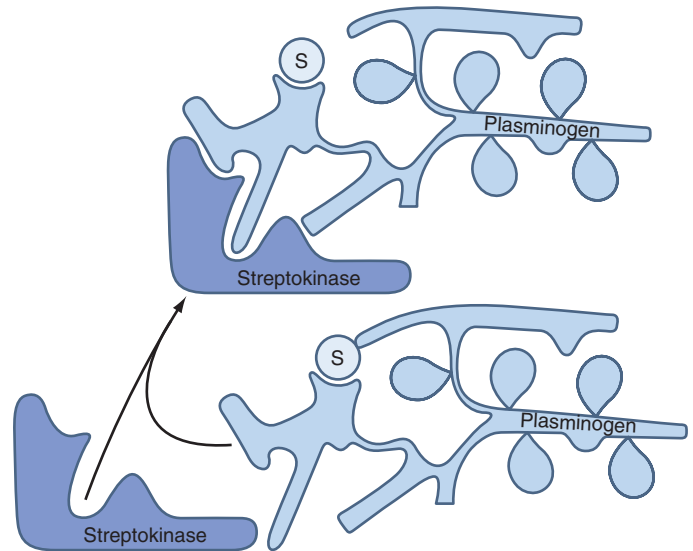


FIGURE 143-8 Mechanism of action of streptokinase. Streptokinase binds to plasminogen and induces a conformational change in plasminogen that exposes its active site. The streptokinase/plasminogen complex then serves as the activator of additional plasminogen.

amounts of plasmin to overwhelm α_2 -antiplasmin. Unopposed plasmin not only degrades fibrin in the occlusive thrombus but also induces a systemic lytic state.

When given systemically to patients with acute MI, streptokinase reduces mortality. For this indication, the drug is usually given as an IV infusion of 1.5 million units over 30–60 min. Patients who receive streptokinase can develop antibodies against the drug, as can patients with prior streptococcal infection. These antibodies can reduce the effectiveness of streptokinase.

Allergic reactions occur in ~5% of patients treated with streptokinase. These may manifest as a rash, fever, chills, and rigors. Although anaphylactic reactions can occur, these are rare. Transient hypotension is common with streptokinase and has been attributed to plasmin-mediated release of bradykinin from kininogen. The hypotension usually responds to leg elevation and administration of IV fluids and low doses of vasopressors, such as dopamine or norepinephrine.

ANISTREPLASE

To generate this drug, streptokinase is combined with equimolar amounts of Lys-plasminogen, a plasmin-cleaved form of plasminogen with a Lys residue at its N terminus. The active site of Lys-plasminogen that is exposed upon combination with streptokinase is then masked with an anisoyl group. After IV infusion, the anisoyl group is slowly removed by deacylation, giving the complex a half-life of ~100 min. This allows drug administration via a single bolus infusion.

Although it is more convenient to administer, anistreplase offers few mechanistic advantages over streptokinase. Like streptokinase, anistreplase does not distinguish between fibrin-bound and circulating plasminogen. Consequently, it too produces a systemic lytic state. Likewise, allergic reactions and hypotension are just as frequent with anistreplase as they are with streptokinase.

When anistreplase was compared with alteplase in patients with acute MI, reperfusion was obtained more rapidly with alteplase than with anistreplase. Improved reperfusion was associated with a trend toward better clinical outcomes and reduced mortality rate with alteplase. These results and the high cost of anistreplase have dampened the enthusiasm for its use.

UROKINASE

Urokinase is a two-chain serine protease derived from cultured fetal kidney cells with a molecular weight of 34,000. Urokinase converts plasminogen to plasmin directly by cleaving the Arg560-Val561 bond. Unlike streptokinase, urokinase is not immunogenic and allergic

reactions are rare. Urokinase produces a systemic lytic state because it does not discriminate between fibrin-bound and circulating plasminogen.

Despite many years of use, urokinase has never been systemically evaluated for coronary thrombolysis. Instead, urokinase is often employed for catheter-directed lysis of thrombi in the deep veins or the peripheral arteries. Because of production problems, the availability of urokinase is limited.

ALTEPLASE

A recombinant form of single-chain tPA, alteplase has a molecular weight of 68,000. Alteplase is rapidly converted into its two-chain form by plasmin. Although single- and two-chain forms of tPA have equivalent activity in the presence of fibrin, in its absence, single-chain tPA has tenfold lower activity.

Alteplase consists of five discrete domains (Fig. 143-9); the N-terminus A chain of two-chain alteplase contains four of these domains. Residues 4 through 50 make up the finger domain, a region that resembles the finger domain of fibronectin; residues 50 through 87 are homologous with epidermal growth factor, whereas residues 92 through 173 and 180 through 261, which have homology to the kringle domains of plasminogen, are designated as the first and second kringle, respectively. The fifth alteplase domain is the protease domain; it is located on the C-terminus B chain of two-chain alteplase.

The interaction of alteplase with fibrin is mediated by the finger domain and, to a lesser extent, by the second kringle domain. The affinity of alteplase for fibrin is considerably higher than that for fibrinogen. Consequently, the catalytic efficiency of plasminogen activation by alteplase is two to three orders of magnitude higher in the presence of fibrin than in the presence of fibrinogen. This phenomenon helps to localize plasmin generation to the fibrin surface.

Although alteplase preferentially activates plasminogen in the presence of fibrin, alteplase is not as fibrin-selective as was first predicted. Its fibrin specificity is limited because like fibrin, (DD)E, the major

soluble degradation product of cross-linked fibrin, binds alteplase and plasminogen with high affinity. Consequently, (DD)E is as potent as fibrin as a stimulator of plasminogen activation by alteplase. Whereas plasmin generated on the fibrin surface results in thrombolysis, plasmin generated on the surface of circulating (DD)E degrades fibrinogen. Fibrinogen degradation results in the accumulation of fragment X, a high-molecular-weight clottable fibrinogen degradation product. Incorporation of fragment X into hemostatic plugs formed at sites of vascular injury renders them susceptible to lysis. This phenomenon may contribute to alteplase-induced bleeding.

A trial comparing alteplase with streptokinase for treatment of patients with acute MI demonstrated significantly lower mortality with alteplase than with streptokinase, although the absolute difference was small. The greatest benefit was seen in patients age <75 years with anterior MI who presented <6 h after symptom onset.

For treatment of acute MI or acute ischemic stroke, alteplase is given as an IV infusion over 60–90 min. The total dose of alteplase usually ranges from 90 to 100 mg. Allergic reactions and hypotension are rare, and alteplase is not immunogenic.

TENECTEPLASE

Tenecteplase is a genetically engineered variant of tPA and was designed to have a longer half-life than tPA and to be resistant to inactivation by PAI-1. To prolong its half-life, a new glycosylation site was added to the first kringle domain (Fig. 143-9). Because addition of this extra carbohydrate side chain reduced fibrin affinity, the existing glycosylation site on the first kringle domain was removed. To render the molecule resistant to inhibition by PAI-1, a tetra-alanine substitution was introduced at residues 296–299 in the protease domain, the region responsible for the interaction of tPA with PAI-1.

Tenecteplase is more fibrin-specific than tPA. Although both agents bind to fibrin with similar affinity, the affinity of tenecteplase for (DD)E is significantly lower than that of tPA. Consequently, (DD)E does not stimulate systemic plasminogen activation by tenecteplase to the same extent as tPA. As a result, tenecteplase produces less fibrinogen degradation than tPA.

For coronary thrombolysis, tenecteplase is given as a single IV bolus. In a large phase III trial that enrolled >16,000 patients, the 30-day mortality rate with single-bolus tenecteplase was similar to that with accelerated-dose tPA. Although rates of intracranial hemorrhage were also similar with both treatments, patients given tenecteplase had fewer noncerebral bleeds and a reduced need for blood transfusions than those treated with tPA. The improved safety profile of tenecteplase likely reflects its enhanced fibrin specificity.

RETEPLASE

Retepulse is a single-chain, recombinant tPA derivative that lacks the finger, epidermal growth factor, and first kringle domains (Fig. 143-9). This truncated derivative has a molecular weight of 39,000. Retepulse binds fibrin more weakly than tPA because it lacks the finger domain. Because it is produced in *Escherichia coli*, reteplase is not glycosylated. This endows it with a plasma half-life longer than that of tPA. Consequently, reteplase is given as two IV boluses, which are separated by 30 min. Clinical trials have demonstrated that reteplase is at least as effective as streptokinase for treatment of acute MI, but the agent is not superior to tPA.

NEWER FIBRINOLYTIC AGENTS

Two new drugs are under investigation. These include desmoteplase (Fig. 143-9), a recombinant form of the full-length plasminogen activator isolated from the saliva of the vampire bat, and altimeprase, a truncated form of fibrolase, an enzyme isolated from the venom of the southern copperhead snake. Clinical studies with these agents have been disappointing. Desmoteplase, which is more fibrin-specific than tPA, was investigated for treatment of acute ischemic stroke. Patients presenting 3–9 h after symptom onset were randomized to one of two doses of desmoteplase or to placebo. Overall response rates were low and no different with desmoteplase than with placebo. The mortality rate was higher in the desmoteplase arms.

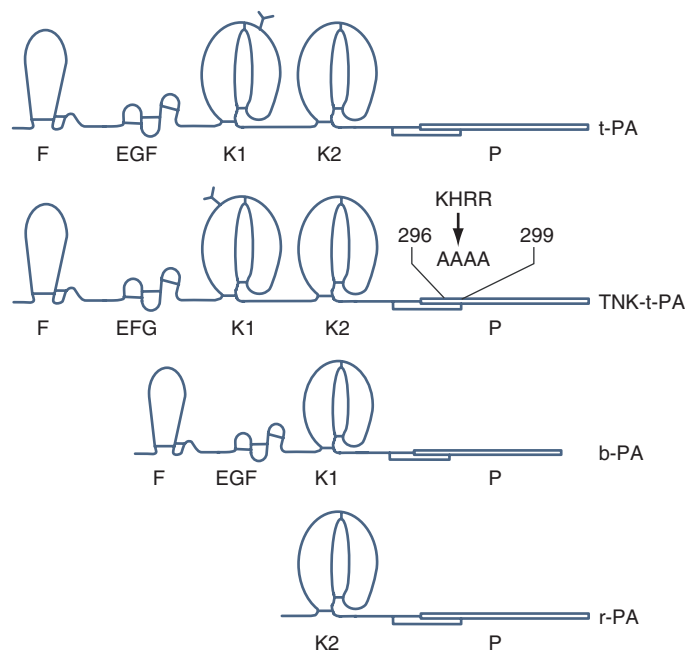


FIGURE 143-9 Domain structures of alteplase (tPA), tenecteplase (TNK-tPA), desmoteplase (b-PA), and reteplase (r-PA). The finger (F), epidermal growth factor (EGF), first and second kringles (K1 and K2, respectively), and protease (P) domains are illustrated. The glycosylation site (Y) on K1 has been repositioned in tenecteplase to endow it with a longer half-life. In addition, a tetra-alanine substitution in the protease domain renders tenecteplase resistant to type 1 plasminogen activator inhibitor (PAI-1) inhibition. Desmoteplase differs from alteplase and tenecteplase in that it lacks a K2 domain. Reteplase is a truncated variant that lacks the F, EGF, and K1 domains.

760 Alfimeprase is a metalloproteinase that degrades fibrin and fibrinogen in a plasmin-independent fashion. In the circulation, alfimeprase is rapidly inhibited by α_2 -macroglobulin. Consequently, the drug must be delivered via a catheter directly into the thrombus. Studies of alfimeprase for treatment of peripheral arterial occlusion or for restoration of flow in blocked central venous catheters were stopped due to lack of efficacy. The disappointing results with desmoteplase and alfimeprase highlight the challenges of introducing new fibrinolytic drugs.

CONCLUSIONS AND FUTURE DIRECTIONS

Thrombosis involves a complex interplay among the vessel wall, platelets, the coagulation system, and the fibrinolytic pathways. Activation of coagulation also triggers inflammatory pathways that may exacerbate

thrombosis. A better understanding of the biochemistry of blood coagulation and advances in structure-based drug design have identified new targets and resulted in the development of novel antithrombotic drugs. Well-designed clinical trials have provided detailed information on which drugs to use and when to use them. Despite these advances, however, thromboembolic disorders remain a major cause of morbidity and mortality. Therefore, the search for better targets and more potent antiplatelet, anticoagulant, and fibrinolytic drugs continues.

SECTION 1 BASIC CONSIDERATIONS IN INFECTIOUS DISEASES

144 Approach to the Patient with an Infectious Disease

Neeraj K. Surana, Dennis L. Kasper

HISTORICAL PERSPECTIVE

The origins of the field of infectious diseases are humble. The notion that communicable diseases were due to a *miasma* (“bad air”) can be traced back to at least the mid-sixteenth century. Not until the work of Louis Pasteur and Robert Koch in the late nineteenth century was there credible evidence supporting the germ theory of disease—i.e., that microorganisms are the direct cause of infections. In contrast to this relatively slow start, the twentieth century saw remarkable advances in the field of infectious diseases, and the etiologic agents of numerous infectious diseases were soon identified. Furthermore, the discovery of antibiotics and the advent of vaccines against some of the most deadly and debilitating infections greatly altered the landscape of human health. Indeed, the twentieth century saw the elimination of smallpox, one of the great scourges in the history of humanity. These remarkable successes prompted noted scholar Aidan Cockburn to write in a 1963 publication entitled *The Evolution and Eradication of Infectious Diseases*: “It seems reasonable to anticipate that within some measurable time . . . all the major infections will have disappeared.” Professor Cockburn was not alone in this view. Robert Petersdorf, a renowned infectious disease expert and former editor of this textbook, wrote in 1978 that “even with my great personal loyalties to infectious diseases, I cannot conceive a need for 309 more [graduating trainees in infectious diseases] unless they spend their time culturing each other.” Given the enormous growth of interest in the microbiome in the past 5 years, Dr. Petersdorf’s statement might have been ironically clairvoyant, although he could have had no idea what was in store for humanity, with an onslaught of new, emerging, and re-emerging infectious diseases.

Clearly, even with all the advances of the twentieth century, infectious diseases continue to represent a formidable challenge for patients and physicians alike. Furthermore, during the latter half of the century, several chronic diseases were demonstrated to be directly or indirectly caused by infectious microbes; perhaps the most notable examples are the associations of *Helicobacter pylori* with peptic ulcer disease and gastric carcinoma, human papillomavirus with cervical cancer, and hepatitis B and C viruses with liver cancer. In fact, ~16% of all malignancies are now known to be associated with an infectious cause. In addition, numerous emerging and re-emerging infectious diseases continue to have a dire impact on global health: HIV/AIDS, pandemic influenza, and severe acute respiratory syndrome (SARS) are but a few examples. The fear of weaponizing pathogens for bioterrorism is ever present and poses a potentially enormous threat to public health. Moreover, escalating antimicrobial resistance in clinically relevant microbes (e.g., *Mycobacterium tuberculosis*, *Staphylococcus aureus*, *Streptococcus pneumoniae*, *Plasmodium* species, and HIV) signifies that the administration of antimicrobial agents—once thought to be a panacea—requires appropriate stewardship. For all these reasons, infectious diseases continue to exert grim effects on individual patients as well as on international public health. Even with all the successes of the past century, physicians must be as thoughtful about infectious diseases now as they were at the beginning of the twentieth century.

GLOBAL CONSIDERATIONS



Infectious diseases remain the second leading cause of death worldwide. Although the rate of infectious disease–related deaths has decreased dramatically over the past 20 years, the absolute numbers of such deaths have remained relatively constant, totaling just over 12 million in 2010 (Fig. 144-1A). As shown in Fig. 144-1B, these deaths disproportionately affect low- and middle-income countries (Chap. 13e); in 2010, 23% of all deaths worldwide were related to infectious diseases, with rates >60% in most sub-Saharan African countries.

Given that infectious diseases are still a major cause of global mortality, understanding the local epidemiology of disease is critically important in evaluating patients. Diseases such as HIV/AIDS have decimated sub-Saharan Africa, with HIV-infected adults representing 15–26% of the total population in countries like Zimbabwe, Botswana, and Swaziland. Moreover, drug-resistant tuberculosis is rampant throughout the former Soviet-bloc countries, India, China, and South Africa. The ready availability of this type of information allows physicians to develop appropriate differential diagnoses and treatment plans for individual patients. Programs such as the Global Burden of Disease seek to quantify human losses (e.g., deaths, disability-adjusted life years) due to diseases by age, sex, and country over time; these data not only help inform local, national, and international health policy but can also help guide local medical decision-making. Even though some diseases (e.g., pandemic influenza, SARS) are seemingly geographically restricted, the increasing ease of rapid worldwide travel has raised concern about their swift spread around the globe. The world’s increasing interconnectedness has profound implications not only for the global economy but also for medicine and the spread of infectious diseases.

UNDERSTANDING THE MICROBIOTA

Normal, healthy humans are colonized with over 100 trillion bacteria as well as countless viruses, fungi, and archaea; taken together, these microorganisms outnumber human cells by 10–100 times (Chap. 86e). The major reservoir of these microbes is the gastrointestinal tract, but very substantial numbers of microbes live in the female genital tract, the oral cavity, and the nasopharynx. There is increasing interest in the skin and even the lungs as sites where microbial colonization might be highly relevant to the biology and disease susceptibility of the host. These commensal organisms provide the host with myriad benefits, from aiding in metabolism to shaping the immune system. With regard to infectious diseases, the vast majority of infections are caused by organisms that are part of the normal flora (e.g., *S. aureus*, *S. pneumoniae*, *Pseudomonas aeruginosa*), with relatively few infections due to organisms that are strictly pathogens (e.g., *Neisseria gonorrhoeae*, rabies virus). Perhaps it is not surprising that a general understanding of the microbiota is essential in the evaluation of infectious diseases. Individuals’ microbiotas likely have a major impact on their susceptibility to infectious diseases and even their responses to vaccines. Site-specific knowledge of the indigenous flora may facilitate appropriate interpretation of culture results, aid in selection of empirical antimicrobial therapy based on the likely causative agents, and provide additional impetus for rational antibiotic use to minimize the untoward effects of these drugs on the “beneficial” microbes that inhabit the body.

WHEN TO CONSIDER AN INFECTIOUS ETIOLOGY

The title of this chapter may appear to presuppose that the physician knows when a patient has an infectious disease. In reality, this chapter can serve only as a guide to the evaluation of a patient in whom an

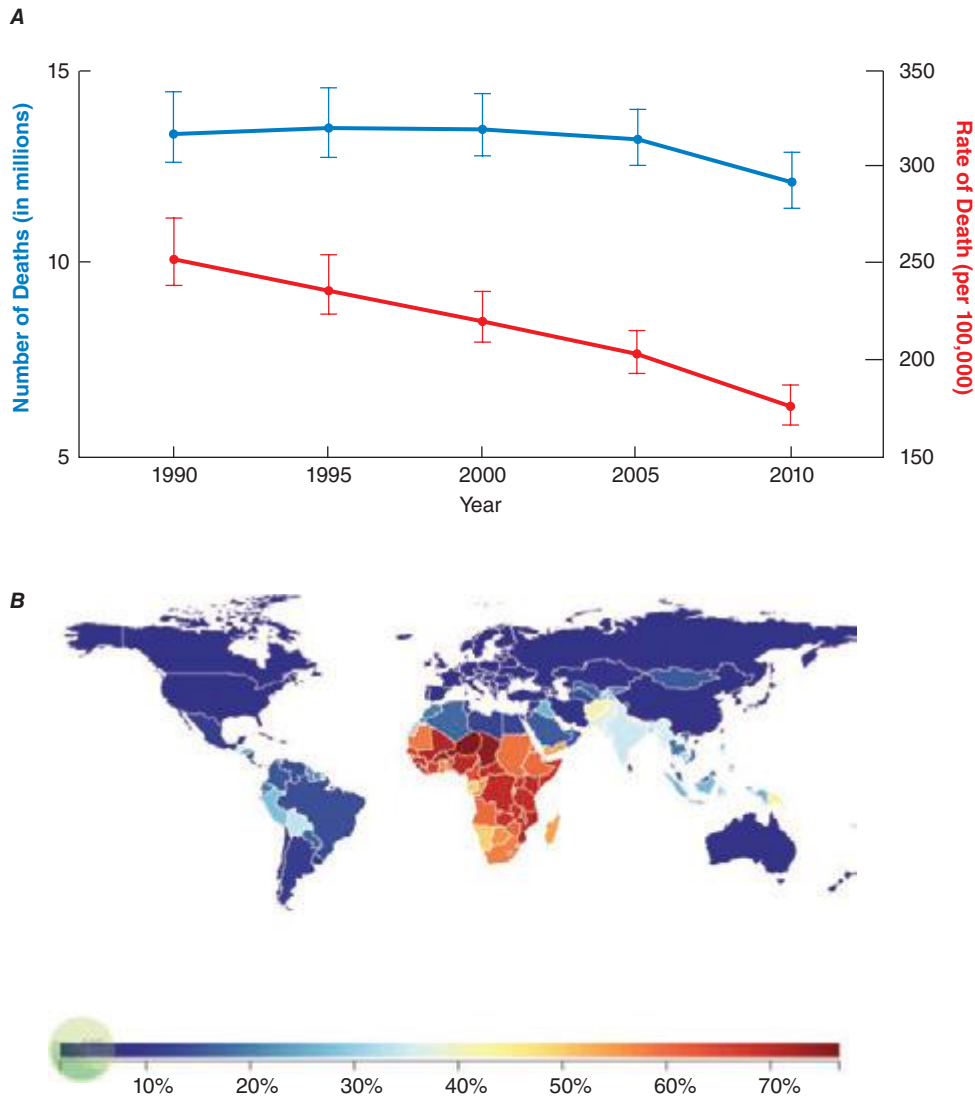


FIGURE 144-1 Magnitude of infectious disease–related deaths globally. **A.** The absolute number (blue line; left axis) and rate (red line; right axis) of infectious disease–related deaths throughout the world since 1990. **B.** A map depicting country-specific data for the percentages of total deaths that were attributable to communicable, maternal, neonatal, and nutritional disorders in 2010. (Source: Global Burden of Disease Study, Institute for Health Metrics and Evaluation.)

infectious disease is a possibility. Once a specific diagnosis is made, the reader should consult the subsequent chapters that deal with specific microorganisms in detail. The challenge for the physician is to recognize which patients may have an infectious disease as opposed to some other underlying disorder. This task is greatly complicated by the fact that infections have an infinite range of presentations, from acute life-threatening conditions (e.g., meningococemia) to chronic diseases of varying severity (e.g., *H. pylori*-associated peptic ulcer disease) to no symptoms at all (e.g., latent *M. tuberculosis* infection). While it is impossible to generalize about a presentation that encompasses all infections, common findings in the history, physical examination, and basic laboratory testing often suggest that the patient either has an infectious disease or should be more closely evaluated for one. This chapter focuses on these common findings and how they may direct the ongoing evaluation of the patient.

APPROACH TO THE PATIENT: Infectious Disease

See also Chap. 147.

HISTORY

As in all of medicine, obtaining a complete and thorough history is paramount in the evaluation of a patient with a possible infectious

disease. The history is critical for developing a focused differential diagnosis and for guiding the physical exam and initial diagnostic testing. Although detailing all the elements of a history is beyond the scope of this chapter, specific components relevant to infectious diseases require particular attention. In general, these aspects focus on two areas: (1) an exposure history that may identify microorganisms with which the patient may have come into contact and (2) host-specific factors that may predispose to the development of an infection.

Exposure History • HISTORY OF INFECTIONS OR EXPOSURE TO DRUG-RESISTANT MICROBES Knowledge about a patient's previous infections, with the associated microbial susceptibility profiles, is very helpful in determining possible etiologic agents. Specifically, knowing whether a patient has a history of infection with drug-resistant organisms (e.g., methicillin-resistant *S. aureus*, vancomycin-resistant *Enterococcus* species, enteric organisms that produce an extended-spectrum β -lactamase or carbapenemase) or may have been exposed to drug-resistant microbes (e.g., during a recent stay in a hospital, nursing home, or long-term acute-care facility) may alter the choice of empirical antibiotics. For example, a patient presenting with sepsis who is known to have a history of invasive infection with a multidrug-resistant isolate of *P. aeruginosa* should be treated empirically with an antimicrobial regimen that will cover this strain.

SOCIAL HISTORY Although the social history taken by physicians is often limited to inquiries about a patient's alcohol and tobacco use, a complete social history can offer a number of clues to the underlying diagnosis. Knowing whether the patient has any high-risk behaviors (e.g., unsafe sexual behaviors, IV drug use), potential hobby-associated exposures (e.g., avid gardening, with possible *Sporothrix schenckii* exposure), or occupational exposures (e.g., increased risk for *M. tuberculosis* exposure in funeral service workers) can facilitate diagnosis. The importance of the social history is exemplified by a case in 2009 in which a laboratory researcher died of a *Yersinia pestis* infection acquired during his work; although this patient had visited both an outpatient clinic and an emergency department, his records at both sites failed to include his occupation—information that potentially could have led quickly to appropriate treatment and infection control measures.

DIETARY HABITS As certain pathogens are associated with specific dietary habits, inquiring about a patient's diet can provide insight into possible exposures. For example, Shiga toxin-producing strains of *Escherichia coli* and *Toxoplasma gondii* are associated with the consumption of raw or undercooked meat; *Salmonella typhimurium*, *Listeria monocytogenes*, and *Mycobacterium bovis* with unpasteurized milk; *Leptospira* species, parasites, and enteric bacteria with unpurified water; and *Vibrio* species, norovirus, helminths, and protozoa with raw seafood.

ANIMAL EXPOSURES Because animals are often important vectors of infectious diseases, patients should be asked about exposures to any animals, including contact with their own pets, visits to petting zoos, or random encounters (e.g., home rodent infestation). For example, dogs can carry ticks that serve as agents for the transmission of several infectious diseases, including Lyme disease, Rocky Mountain spotted fever, and ehrlichiosis. Cats are associated with *Bartonella henselae* infection, reptiles with *Salmonella* infection, rodents with leptospirosis, and rabbits with tularemia (Chap. 167e).

TRAVEL HISTORY Attention should be paid to both international and domestic travel. Fever in a patient who has recently returned from abroad significantly broadens the differential diagnosis (Chap. 149); even a remote history of international travel may reflect patients' exposure to infections with pathogens such as *M. tuberculosis* or *Strongyloides stercoralis*. Similarly, domestic travel may have exposed patients to pathogens that are not normally found in their local environment and therefore may not routinely be considered in the differential diagnosis. For example, a patient who has recently visited California or Martha's Vineyard may have been exposed to *Coccidioides immitis* or *Francisella tularensis*, respectively. Beyond simply identifying locations that a patient may have visited, the physician needs to delve deeper to learn what kinds of activities and behaviors the patient engaged in during travel (e.g., the types of food and sources of water consumed, freshwater swimming, animal exposures) and whether the patient had the necessary immunizations and/or took the necessary prophylactic medications prior to travel; these additional exposures, which the patient may not think to report without specific prompting, are as important as exposures during a patient's routine daily living.

Host-Specific Factors Because many opportunistic infections (e.g., with *Pneumocystis jirovecii*, *Aspergillus* species, or JC virus) affect only immunocompromised patients, it is of vital importance to determine the immune status of the patient. Defects in the immune system may be due to an underlying disease (e.g., malignancy, HIV infection, malnutrition), a medication (e.g., chemotherapy, glucocorticoids, monoclonal antibodies to components of the immune system), a treatment modality (e.g., total body irradiation, splenectomy), or a primary immunodeficiency. The type of infection for which the patient is at increased risk varies with the specific type of immune defect (Chap. 375e). In concert with determining whether a patient is immunocompromised for any reason, the physician should

review the immunization record to ensure that the patient is adequately protected against vaccine-preventable diseases (Chap. 148).

PHYSICAL EXAMINATION

Similar to the history, a thorough physical examination is crucial in evaluating patients with an infectious disease. Some elements of the physical exam (e.g., skin, lymphatics) that are often performed in a cursory manner as a result of the ever-increasing pace of medical practice may help identify the underlying diagnosis. Moreover, serial exams are critical since new findings may appear as the illness progresses. A description of all the elements of a physical exam is beyond the scope of this chapter, but the following components have particular relevance to infectious diseases.

Vital Signs Given that elevations in temperature are often a hallmark of infection, paying close attention to the temperature may be of value in diagnosing an infectious disease. The idea that 37°C (98.6°F) is the normal human body temperature dates back to the nineteenth century and was initially based on axillary measurements. Rectal temperatures more accurately reflect the core body temperature and are 0.4°C (0.7°F) and 0.8°C (1.4°F) higher than oral and axillary temperatures, respectively. Although the definition of fever varies greatly throughout the medical literature, the most common definition, which is based on studies defining fever of unknown origin (Chap. 26), uses a temperature $\geq 38.3^\circ\text{C}$ (101°F). Although fever is very commonly associated with infection, it is also documented in many other diseases (Chap. 23). For every 1°C (1.8°F) increase in core temperature, the heart rate typically rises by 15–20 beats/min. Table 144-1 lists infections that are associated with relative bradycardia (*Faget's sign*), where patients have a lower heart rate than might be expected for a given body temperature. Although this pulse-temperature dissociation is not highly sensitive or specific for establishing a diagnosis, it is potentially useful in low-resource settings given its ready availability and simplicity.

Lymphatics There are ~600 lymph nodes throughout the body, and infections are an important cause of lymphadenopathy. A physical examination should include evaluation of lymph nodes in multiple

TABLE 144-1 CAUSES OF RELATIVE BRADYCARDIA

Infectious Causes	
Intracellular organisms	
Gram-negative bacteria	<i>Salmonella typhi</i> <i>Francisella tularensis</i> <i>Brucella</i> spp. <i>Coxiella burnetii</i> (Q fever) <i>Leptospira interrogans</i> <i>Legionella pneumophila</i> <i>Mycoplasma pneumoniae</i>
Tick-borne organisms	<i>Rickettsia</i> spp. <i>Orientia tsutsugamushi</i> (scrub typhus) <i>Babesia</i> spp.
Other	<i>Corynebacterium diphtheriae</i> <i>Plasmodium</i> spp. (malaria)
Viruses/viral infections	Yellow fever virus Dengue virus Viral hemorrhagic fevers ^a Viral myocarditis
Noninfectious Causes	
	Drug fever Beta blocker use Central nervous system lesions Malignant lymphoma Factitious fever

^aPrimarily early in the course of infection with Marburg or Ebola virus.

regions (e.g., popliteal, inguinal, epitrochlear, axillary, multiple cervical regions), with notation of the location, size (normal, <1 cm), presence or absence of tenderness, and consistency (soft, firm, or shotty) and of whether the nodes are matted (i.e., connected and moving together). Of note, palpable epitrochlear nodes are always pathologic. Of patients presenting with lymphadenopathy, 75% have localized findings, and the remaining 25% have generalized lymphadenopathy (i.e., that involving more than one anatomic region). Localized lymphadenopathy in the head and neck region is found in 55% of patients, inguinal lymphadenopathy in 14%, and axillary lymphadenopathy in 5%. Determining whether the patient has generalized versus localized lymphadenopathy can help narrow the differential diagnosis, as various infections present differently.

Skin The fact that many infections have cutaneous manifestations gives the skin examination particular importance in the evaluation of patients (Chaps. 24, 25e, 72, and 156). It is important to perform a complete skin exam, with attention to both front and back. Specific rashes are often extremely helpful in narrowing the differential diagnosis of an infection (Chaps. 24 and 25e). In numerous anecdotal instances, patients in the intensive care unit have had “fever of unknown origin” that was actually due to unrecognized pressure ulcers. Moreover, close examination of the distal extremities for splinter hemorrhages, Janeway lesions, or Osler’s nodes may yield evidence of endocarditis or other causes of septic emboli.

Foreign Bodies As previously mentioned, many infections are caused by members of the indigenous microbiota. These infections typically occur when these microbes escape their normal habitat and enter a new one. Thus, maintenance of epithelial barriers is one of the most important mechanisms in protection against infection. However, hospitalization of patients is often associated with breaches of these barriers—e.g., due to placement of IV lines, surgical drains, or tubes (such as endotracheal tubes and Foley catheters) that allow microorganisms to localize in sites to which they normally would not have access (Chap. 168). Accordingly, knowing what lines, tubes, and drains are in place is helpful in ascertaining what body sites might be infected.

DIAGNOSTIC TESTING

Laboratory and radiologic testing has advanced greatly over the past few decades and has become an important component in the evaluation of patients. The dramatic increase in the number of serologic diagnostics, antigen tests, and molecular diagnostics available to the physician has, in fact, revolutionized medical care. However, all of these tests should be viewed as adjuncts to the history and physical examination—not a replacement for them. The selection of initial tests should be based directly on the patient’s history and physical exam findings. Moreover, diagnostic testing should generally be limited to those conditions that are reasonably likely and treatable, important in terms of public health considerations, and/or capable of providing a definitive diagnosis that will consequently limit other testing.

White Blood Cell (WBC) Count Elevations in the WBC count are often associated with infection, though many viral infections are associated with leukopenia. It is important to assess the WBC differential, given that different classes of microbes are associated with various leukocyte types. For example, bacteria are associated with an increase in polymorphonuclear neutrophils, often with elevated levels of earlier developmental forms such as bands; viruses are associated with an increase in lymphocytes; and certain parasites are associated with an increase in eosinophils. Table 144-2 lists the major infectious causes of eosinophilia.

Inflammatory Markers The erythrocyte sedimentation rate (ESR) and the C-reactive protein (CRP) level are indirect and direct measures of the acute-phase response, respectively, that can be used to assess a patient’s general level of inflammation. Moreover, these markers can be followed serially over time to monitor disease

progress/resolution. It is noteworthy that the ESR changes relatively slowly, and its measurement more often than weekly usually is not useful; in contrast, CRP concentrations change rapidly, and daily measurements can be useful in the appropriate context. Although these markers are sensitive indicators of inflammation, neither is very specific. An extremely elevated ESR (>100 mm/h) has a 90% predictive value for a serious underlying disease (Table 144-3). Work is ongoing to identify other potentially useful inflammatory markers (e.g., procalcitonin, serum amyloid A protein); however, their clinical utility requires further validation.

Analysis of Cerebrospinal Fluid (CSF) Assessment of CSF is critical for patients with suspected meningitis or encephalitis. An opening pressure should always be recorded, and fluid should routinely be sent for cell counts, Gram’s stain and culture, and determination of glucose and protein levels. A CSF Gram’s stain typically requires >10⁵ bacteria/mL for reliable positivity; its specificity approaches 100%. Table 144-4 lists the typical CSF profiles for various infections. In general, CSF with a lymphocytic pleocytosis and a low glucose concentration suggests either infection (e.g., with *Listeria*, *M. tuberculosis*, or a fungus) or a noninfectious disorder (e.g., neoplastic meningitis, sarcoidosis). Bacterial antigen testing of CSF (e.g., latex agglutination tests for *Haemophilus influenzae* type b, group B *Streptococcus*, *S. pneumoniae*, and *Neisseria meningitidis*) is not recommended as a screening assay, given that these tests are no more sensitive than Gram’s stain; however, these assays can be helpful in presumptively identifying organisms seen on Gram’s stain. In contrast, other antigen tests (e.g., for *Cryptococcus*) and some CSF serologic testing (e.g., for *Treponema pallidum*, *Coccidioides*) are highly sensitive and are useful for select patients. In addition, polymerase chain reaction (PCR) analysis of CSF is increasingly being used for the diagnosis of bacterial (e.g., *N. meningitidis*, *S. pneumoniae*, mycobacteria) and viral (e.g., herpes simplex virus, enterovirus) infections; while these molecular tests permit rapid diagnosis with a high degree of sensitivity and specificity, they often do not allow determination of antimicrobial resistance profiles.

Cultures The mainstays of infectious disease diagnosis include the culture of infected tissue (e.g., surgical specimens) or fluid (e.g., blood, urine, sputum, purulence from a wound). Samples can be sent for culture of bacteria (aerobic or anaerobic), fungi, or viruses. Ideally, specimens are collected before the administration of antimicrobial therapy; in instances where this order of events is not clinically feasible, microscopic examination of the specimen (e.g., Gram-stained or potassium hydroxide [KOH]-treated preparations) is particularly important. Culture of the organism(s) allows identification of the etiologic agent, determination of the antimicrobial susceptibility profile, and—when there is concern about an outbreak—isolate typing. While cultures are extremely useful in the evaluation of patients, determining whether culture results are clinically meaningful or represent contamination (e.g., a non-*aureus*, non-*lugdunensis* staphylococcal species growing in a blood culture) can sometimes be challenging and requires an understanding of the patient’s immune status, exposure history, and microbiota. In some cases, serial cultures to demonstrate clearance of the organism may be helpful.

Pathogen-Specific Testing Numerous pathogen-specific tests (e.g., serology, antigen testing, PCR testing) are commercially available, and many hospitals now offer some of these tests in-house to facilitate rapid turnaround that ultimately enhances patient care. The reader is directed to relevant chapters on the pathogens of interest for specific details. Some of these tests (e.g., universal PCRs) identify organisms that currently are not cultivable and have unclear relationships to disease, thereby complicating diagnosis. As these tests become more commonplace and the work of the Human Microbiome Project progresses, the relevance of some of these previously unrecognized bacteria to human health will likely become more apparent.

TABLE 144-2 MAJOR INFECTIOUS CAUSES OF EOSINOPHILIA^a

Organ Involved	Organism	Exposure	Geographic Distribution	Degree of Eosinophilia ^b
Central nervous system	<i>Angiostrongylus</i>	Raw seafood	Asia	Mild
	<i>Gnathostoma</i>	Raw poultry and seafood	Asia	Moderate to extreme
Eye	<i>Loa loa</i>	Insect bite	Africa	Moderate (expatriates), mild (patients living in endemic areas)
	<i>Onchocerca</i>	Insect bite	Africa	Mild (expatriates), moderate (patients living in endemic areas)
Lung	<i>Chlamydia trachomatis</i>	Sexual transmission	Worldwide	Mild
	<i>Strongyloides</i>	Soil	Tropical	Moderate (acute), mild (chronic)
	<i>Toxocara canis/Toxocara cati</i> ^c	Dogs, soil	Worldwide	Moderate to extreme
	<i>Paragonimus</i>	Crabs and crayfish	Asia	Moderate (acute), mild (chronic)
	<i>Coccidioides immitis</i>	Soil	Southwestern United States	Mild (acute), extreme (disseminated)
	<i>Brugia malayi</i>	Insect bite	Asia	Mild to moderate
	<i>Pneumocystis jirovecii</i>	Air	Worldwide	Mild
Liver	<i>Schistosoma japonicum</i>	Freshwater swimming	Asia	Moderate (acute), mild (chronic)
	<i>Schistosoma mansoni</i>	Freshwater swimming	Africa, Middle East, Latin America	Moderate (acute), mild (chronic)
	<i>Fasciola</i>	Watercress	Worldwide	Moderate
	<i>Clonorchis</i>	Raw seafood	Asia	Mild to moderate
	<i>Opisthorchis</i>	Raw seafood	Asia	Mild to moderate
Intestines	<i>Ascaris</i> ^d	Raw fruits and vegetables, contaminated water	Worldwide	Mild to extreme
	Hookworm	Soil	Worldwide	Mild to moderate
	<i>Trichuris</i>	Raw fruits and vegetables, contaminated water	Tropical	Mild
	<i>Cystoisospora belli</i>	Contaminated water and food	Worldwide	Mild
	<i>Dientamoeba fragilis</i>	Unclear; spread via fecal-oral route	Worldwide	Mild
	<i>Capillaria</i>	Raw seafood	Asia	Extreme
	<i>Heterophyes</i>	Raw seafood	Asia, Middle East	Mild
	<i>Anisakis</i>	Raw seafood	Worldwide	Mild
	<i>Baylisascaris procyonis</i> ^e	Soil	North America	Moderate to extreme
	<i>Hymenolepis nana</i>	Contaminated water, soil	Worldwide	Mild
Bladder	<i>Schistosoma haematobium</i>	Freshwater swimming	Africa, Middle East	Moderate (acute), mild (chronic)
Muscle	<i>Trichinella</i>	Pork	Worldwide	Moderate to extreme
Lymphatics	<i>Wuchereria bancrofti</i> ^d	Insect bite	Tropical	Moderate to extreme ^f
	<i>Bartonella henselae</i>	Cats	Worldwide	Mild
Other	Recovery from bacterial or viral infections	—	—	Mild
	HIV	Contaminated bodily fluid	Worldwide	Mild
	<i>Cryptococcus neoformans</i>	Soil	Worldwide	Moderate to extreme (disseminated)

^aThere are numerous noninfectious causes of eosinophilia, such as atopic disease, DRESS (drug reaction with eosinophilia and systemic symptoms) syndrome, and pernicious anemia, which can cause mild eosinophilia; drug hypersensitivity and serum sickness, which can cause mild to moderate eosinophilia; collagen vascular disease, which can cause moderate eosinophilia; and malignancy, Churg-Strauss syndrome, and hyper-IgE syndromes, which can cause moderate to extreme eosinophilia. ^bMild: 500–1500 cells/μL; moderate: 1500–5000 cells/μL; extreme: >5000 cells/μL. ^cCan also affect the liver and the eyes. ^dCan also affect the lungs. ^eCan also affect the eyes and the central nervous system. ^fLevels are typically higher with pulmonary infections.

Radiology Imaging provides an important adjunct to the physical examination, allowing evaluation for lymphadenopathy in regions that are not externally accessible (e.g., mediastinum, intraabdominal sites), assessment of internal organs for evidence of infection, and facilitation of image-guided percutaneous sampling of deep spaces. The choice of imaging modality (e.g., CT, MRI, ultrasound, nuclear medicine, use of contrast) is best made in consultation with a radiologist to ensure that the results will address the physician's specific concerns.

TREATMENT

Physicians often must balance the need for empirical antibiotic treatment with the patient's clinical condition. When clinically feasible, it is best to obtain relevant samples (e.g., blood, CSF, tissue,

purulent exudate) for culture prior to the administration of antibiotics, as antibiotic treatment often makes subsequent diagnosis more difficult. Although a general maxim for antibiotic treatment is to use a regimen with as narrow a spectrum as possible (**Chap. 170**), empirical regimens are necessarily somewhat broad, given that a specific diagnosis has not yet been made. **Table 144-5** lists empirical antibiotic treatment regimens for commonly encountered infectious presentations. These regimens should be narrowed as appropriate once a specific diagnosis is made. In addition to antibiotics, there is sometimes a role for adjunctive therapies, such as intravenous immunoglobulin G (IVIG) pooled from healthy adults or hyperimmune globulin prepared from the blood of individuals with high titers of specific antibodies to select pathogens (e.g., cytomegalovirus, hepatitis B virus, rabies virus, vaccinia virus,

TABLE 144-3 CAUSES OF AN EXTREMELY ELEVATED ERYTHROCYTE SEDIMENTATION RATE (>100 mm/h)

Etiologic Category (% of Cases)	Specific Causes
Infectious diseases (35–40)	Subacute bacterial endocarditis Abscesses Osteomyelitis Tuberculosis Urinary tract infection
Inflammatory diseases (15–20)	Giant cell arteritis Rheumatoid arthritis Systemic lupus erythematosus
Malignancies (15–20)	Multiple myeloma Leukemias Lymphomas Carcinomas
Other (20–35)	Drug hypersensitivity reactions (drug fever) Ischemic tissue injury/trauma Renal diseases

Clostridium tetani, varicella-zoster virus, *Clostridium botulinum* toxin). Although the data suggesting efficacy are limited, IVIG is often used for patients with suspected staphylococcal or streptococcal toxic shock syndrome.

INFECTION CONTROL

When evaluating a patient with a suspected infectious disease, the physician must consider what infection control methods are necessary to prevent transmission of any possible infection to other people. In 2007, the U.S. Centers for Disease Control and Prevention published guidelines for isolation precautions that are available for download at www.cdc.gov/hicpac/2007IP/2007isolationPrecautions.html. Persons exposed to certain pathogens (e.g., *N. meningitidis*, HIV, *Bacillus anthracis*) should receive postexposure prophylaxis to prevent disease acquisition. (See relevant chapters for details on specific pathogens.)

WHEN TO OBTAIN AN INFECTIOUS DISEASE CONSULT

At times, primary physicians need assistance with patient management, from a diagnostic and/or therapeutic perspective. Multiple studies have demonstrated that an infectious disease consult is associated with positive outcomes for patients with various diseases. For example, in a prospective cohort study of patients with *S. aureus* bacteremia, infectious disease consultation was independently associated with a 56% reduction in 28-day mortality. In addition, infectious disease specialists provide other services (e.g., infection control, antimicrobial stewardship, management of outpatient antibiotic therapy, occupational exposure programs) that have been shown to benefit patients. Whenever such assistance would be advantageous to a patient with a possible infection, the primary physician should opt for an infectious disease consult. Specific situations that might prompt a consult include (1) difficult-to-diagnose patients with presumed infections, (2) patients who are not responding to treatment as expected, (3) patients with a complicated medical history (e.g., organ transplant recipients, patients immunosuppressed due to autoimmune or inflammatory conditions), and (4) patients with “exotic” diseases (i.e., diseases that are not typically seen within the region).

PERSPECTIVE

The study of infectious diseases is really a study of host-bacterial interactions and represents evolution by both the host and the bacteria—an endless struggle in which microbes have generally been more creative and adaptive. Given that nearly one-quarter of deaths worldwide are still related to infectious diseases, it is clear that the war against infectious diseases has not been won. For example, a cure for HIV infection is still lacking, there have been only marginal improvements in the methods for detection and treatment of tuberculosis after more than a half century of research, new infectious diseases (e.g., pandemic influenza, viral hemorrhagic fevers) continue to emerge, and the threat of microbial bioterrorism remains high. The subsequent chapters in Part 8 detail—on both a syndrome and a microbe-by-microbe basis—the current state of medical knowledge about infectious diseases. At their core, all of these chapters carry a similar message: Despite numerous advances in the diagnosis, treatment, and prevention of infectious diseases, much work and research are required before anyone can confidently claim that “all the major infections have disappeared.” In reality, this goal will never be attained, given the rapid adaptability of microbes.

TABLE 144-4 TYPICAL CSF PROFILES FOR MENINGITIS AND ENCEPHALITIS^a

	Normal	Bacterial Meningitis	Viral Meningitis	Fungal Meningitis ^b	Parasitic Meningitis	Tuberculous Meningitis	Encephalitis
WBC count (per μ L)	<5	>1000	25–500	40–600	150–2000	25–100	50–500
Differential of WBC	60–70% lymphocytes, \leq 30% monocytes/macrophages	$\uparrow\uparrow$ PMNs (\geq 80%)	Predominantly lymphocytes ^c	Lymphocytes or PMNs, depending on specific organism	$\uparrow\uparrow$ Eosinophils (\geq 50%) ^d	Predominantly lymphocytes ^c	Predominantly lymphocytes ^c
Gram's stain	Negative	Positive (in >60% of cases)	Negative	Rarely positive	Negative	Occasionally positive ^e	Negative
Glucose (mg/dL)	40–85	<40	Normal	\downarrow to normal	Normal	<50 in 75% of cases	Normal
Protein (mg/dL)	15–45	>100	20–80	150–300	50–200	100–200	50–100
Opening pressure (mmH ₂ O)	50–180	>300	100–350	160–340	Normal	150–280	Normal to \uparrow
Common causes	—	<i>Streptococcus pneumoniae</i> , <i>Neisseria meningitidis</i>	Enteroviruses	<i>Candida</i> , <i>Cryptococcus</i> , and <i>Aspergillus</i> spp.	<i>Angiostrongylus cantonensis</i> , <i>Gnathostoma spinigerum</i> , <i>Baylisascaris procyonis</i>	<i>Mycobacterium tuberculosis</i>	Herpesviruses, enteroviruses, influenza virus, rabies virus

^aNumbers indicate typical results, but actual results may vary. ^bCerebrospinal fluid characteristics depend greatly on the specific organism. ^cNeutrophils may predominate early in the disease course. ^dPatients typically have striking eosinophilia as well. ^eSensitivity can be increased by examination of a smear of protein coagulum (pellicle) and the use of acid-fast stains.

Abbreviations: PMNs, polymorphonuclear neutrophils; WBC, white blood cell.

TABLE 144-5 INITIAL EMPIRICAL ANTIBIOTIC THERAPY FOR COMMON INFECTIOUS DISEASE PRESENTATIONS^a

Clinical Syndrome	Common Etiologies	Antibiotic(s)	Comments	See Chapter(s)
Septic shock	<i>Staphylococcus aureus</i> , <i>Streptococcus pneumoniae</i> , enteric gram-negative bacilli	Vancomycin, 15 mg/kg q12h ^b ; plus A broad-spectrum antipseudomonal β-lactam (piperacillin-tazobactam, 4.5 g q6h; imipenem, 1 g q8h; meropenem, 1 g q8h; or cefepime, 1–2 g q8–12h)	—	325
Meningitis	<i>S. pneumoniae</i> , <i>Neisseria meningitidis</i>	Vancomycin, 15 mg/kg q12h ^b ; plus Ceftriaxone, 2 g q12h	Dexamethasone (0.15 mg/kg IV q6h for 2–4 d) should be added for patients with suspected or proven pneumococcal meningitis, with the first dose administered 10–20 min before the first dose of antibiotics.	164 and pathogen-specific chapters
CNS abscess	<i>Streptococcus</i> spp., <i>Staphylococcus</i> spp., anaerobes, gram-negative bacilli	Vancomycin, 15 mg/kg q12h ^b ; plus Ceftriaxone, 2 g q12h; plus Metronidazole, 500 mg q8h	—	164
Endocarditis	<i>S. aureus</i> , <i>Streptococcus</i> spp., coagulase-negative staphylococci	Vancomycin, 15 mg/kg q12h ^b ; plus Ceftriaxone, 2 g q12h	—	155
Pneumonia Community-acquired, outpatient	<i>S. pneumoniae</i> , <i>Mycoplasma pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Chlamydia pneumoniae</i>	Azithromycin, 500 mg PO × 1, then 250 mg PO qd × 4 days	If MRSA is a consideration, add vancomycin (15 mg/kg q12h ^b) or linezolid (600 mg q12h); daptomycin should not be used in patients with pneumonia.	153 and pathogen-specific chapters
Inpatient, non-ICU	Above plus <i>Legionella</i> spp.	A respiratory fluoroquinolone (moxifloxacin, 400 mg IV/PO qd; gemifloxacin, 320 mg PO qd; or levofloxacin, 750 mg IV/PO qd); or A β-lactam (cefotaxime, ceftriaxone, or ampicillin-sulbactam) <i>plus</i> azithromycin		
Inpatient, ICU	Above plus <i>S. aureus</i>	A β-lactam; plus Azithromycin <i>or</i> a respiratory fluoroquinolone		
Hospital-acquired pneumonia ^d	<i>S. pneumoniae</i> , <i>H. influenzae</i> , <i>S. aureus</i> , gram-negative bacilli (e.g., <i>Pseudomonas aeruginosa</i> , <i>Klebsiella pneumoniae</i> , <i>Acinetobacter</i> spp.)	An antipseudomonal β-lactam (cefepime, 1–2 g q8–12 h; ceftazidime, 2 g q8h; imipenem, 1 g q8h; meropenem, 1 g q8h; or piperacillin-tazobactam, 4.5 g q6h); plus An antipseudomonal fluoroquinolone (levofloxacin or ciprofloxacin, 400 mg q8h) <i>or</i> an aminoglycoside (amikacin, 20 mg/kg q24h ^e ; gentamicin, 7 mg/kg q24h ^e ; or tobramycin, 7 mg/kg q24h ^e)		
Complicated intra-abdominal infection			If MRSA is a consideration, add vancomycin (15 mg/kg q12h ^b).	159, 201 , and pathogen-specific chapters
Mild to moderate severity	Anaerobes (<i>Bacteroides</i> spp., <i>Clostridium</i> spp.), gram-negative bacilli (<i>Escherichia coli</i>), <i>Streptococcus</i> spp.	Cefoxitin, 2 g q6h; or A combination of metronidazole (500 mg q8–12h) <i>plus</i> cefazolin (1–2 g q8h) <i>or</i> cefuroxime (1.5 g q8h) <i>or</i> ceftriaxone (1–2 g q12–24h) <i>or</i> cefotaxime (1–2 g q6–8h)		
High-risk patient or high degree of severity	Same as above	A carbapenem (imipenem, 1 g q8h; meropenem, 1 g q8h; doripenem, 500 mg q8h); or Piperacillin-tazobactam, 3.375 g q6h ^f ; or A combination of metronidazole (500 mg q8–12h) <i>plus</i> an antipseudomonal cephalosporin (cefepime, 2 g q8–12h; ceftazidime, 2 g q8h) <i>or</i> an antipseudomonal fluoroquinolone (ciprofloxacin, 400 mg q12h; levofloxacin, 750 mg q24h)		

(Continued)

TABLE 144-5 INITIAL EMPIRICAL ANTIBIOTIC THERAPY FOR COMMON INFECTIOUS DISEASE PRESENTATIONS^a (CONTINUED)

Clinical Syndrome	Common Etiologies	Antibiotic(s)	Comments	See Chapter(s)
Skin and soft tissue infection	<i>S. aureus</i> , <i>Streptococcus pyogenes</i>	Dicloxacillin, 250–500 mg PO qid; or Cephalexin, 250–500 mg PO qid; or Clindamycin, 300–450 mg PO tid; or Nafcillin/oxacillin, 1–2 g q4h	If MRSA is a consideration, clindamycin, vancomycin (15 mg/kg q12h ^b), linezolid (600 mg IV/PO q12h), or TMP-SMX (1–2 double-strength tablets PO bid ^c) can be used.	156 and pathogen-specific chapters

^aThis table refers to immunocompetent adults with normal renal and hepatic function. All doses listed are for parenteral administration unless indicated otherwise. Local antimicrobial susceptibility profiles may influence the choice of antibiotic. Therapy should be tailored once a specific etiologic agent and its susceptibilities are identified. ^bTrough levels for vancomycin should be 15–20 \geq g/mL. ^cTrough levels for amikacin should be <4 \geq g/mL. ^dIn patients with late onset (i.e., after \leq 5 days of hospitalization) or risk factors for multidrug-resistant organisms. ^eTrough levels for gentamicin and tobramycin should be <1 \geq g/mL. ^fIf *P. aeruginosa* is a concern, the dosage may be increased to 3.375 g IV q4h or 4.5 g IV q6h. ^gData on the efficacy of TMP-SMX in skin and soft tissue infections are limited.

Abbreviations: CNS, central nervous system; ICU, intensive care unit; MRSA, methicillin-resistant *S. aureus*; TMP-SMX, trimethoprim-sulfamethoxazole.

145e Molecular Mechanisms of Microbial Pathogenesis

Gerald B. Pier

Over the past four decades, molecular studies of the pathogenesis of microorganisms have yielded an explosion of information about the various microbial and host molecules that contribute to the processes of infection and disease. These processes can be classified into several stages: microbial encounter with and entry into the host; microbial growth after entry; avoidance of innate host defenses; tissue invasion and tropism; tissue damage; and transmission to new hosts. *Virulence* is the measure of an organism's capacity to cause disease and is a function of the pathogenic factors elaborated by microbes. These factors promote *colonization* (the simple presence of potentially pathogenic microbes in or on a host), *infection* (attachment and growth of pathogens and avoidance of host defenses), and *disease* (often, but not always, the result of activities of secreted toxins or toxic metabolites). In addition, the host's inflammatory response to infection greatly contributes to disease and its attendant clinical signs and symptoms. The recent surge of interest in the role of the microbiota and its associated *microbiome*—the collection of microbial genomes residing in or on mammalian organisms—in the physiology of, susceptibility to, and response to infection and in immune system development has had an enormous impact on our understanding of host-pathogen interaction.

THE MICROBIOME

(See also Chap. 86e) We now understand that the indigenous microbial organisms living in close association with almost all animals are organized into complex communities that strongly modulate the ability of pathogenic microbes to become established in or on host surfaces. The sheer numbers of these microbes and their genomic variability vastly exceed the numbers of host cells and genes in a typical animal. Changes and differences in microbiomes within and between individuals, currently characterized by high-throughput DNA sequencing techniques and bioinformatic analysis, affect the development and control of the immune system as well as such diverse conditions as obesity, type 1 diabetes, cognition, neurologic states, autoimmune diseases, and infectious diseases of the skin, gastrointestinal tract, respiratory tract, and vagina. It has been more difficult to directly associate specific types of microbiomes with pathophysiological states and to assess how conserved or variable microbial species within human and animal microbiomes are evolving. Defining clusters of organisms associated with diseases may become more feasible as more data are obtained. Complicating this task are the results from the Human Microbiome Project suggesting a high level of variability among individuals in the components of the microbiome, although many individuals appear to maintain a fairly conserved microbiome throughout their lives. In the context of infectious diseases, clear changes and disruptions of the indigenous microbiome have a strong and often fundamental impact on the progression of infection. Such alterations can be associated with the effects of antibiotic and immunosuppressive drug use on the normal flora, with environmental changes, and with the impact of microbial virulence factors that displace the indigenous microbial flora to facilitate pathogen colonization. As the available technology for defining the microbiome expands, there is no doubt that the resulting data will markedly affect our concepts of and approaches to microbial pathogenesis and infectious disease treatment.

MICROBIAL ENTRY AND ADHERENCE

Entry Sites A microbial pathogen can potentially enter any part of a host organism. In general, the type of disease produced by a particular microbe is often a direct consequence of its route of entry into the body. The most common sites of entry are mucosal surfaces (the respiratory, alimentary, and urogenital tracts) and the skin. Ingestion, inhalation, and sexual contact are typical routes of microbial entry. Other portals of entry include sites of skin injury (cuts, bites, burns, trauma) along with injection via natural (i.e., vector-borne) or artificial

(i.e., needle-stick injury) routes. A few pathogens, such as *Schistosoma* species, can penetrate unbroken skin. The conjunctiva can serve as an entry point for pathogens of the eye, which occasionally spread systemically from that site.

Microbial entry usually relies on the presence of specific factors needed for persistence and growth in a tissue. Fecal-oral spread via the alimentary tract requires a biologic profile consistent with survival in the varied environments of the gastrointestinal tract (including the low pH of the stomach and the high bile content of the intestine) as well as in contaminated food or water outside the host. Organisms that gain entry via the respiratory tract survive well in small moist droplets produced during sneezing and coughing. Pathogens that enter by venereal routes often survive best in the warm moist environment of the urogenital mucosa and have restricted host ranges (e.g., *Neisseria gonorrhoeae*, *Treponema pallidum*, and HIV).

The biology of microbes entering through the skin is highly varied. Some of these organisms can survive in a broad range of environments, such as the salivary glands or alimentary tracts of arthropod vectors, the mouths of larger animals, soil, and water. A complex biology allows protozoan parasites such as *Plasmodium*, *Leishmania*, and *Trypanosoma* species to undergo morphogenic changes that permit transmission to mammalian hosts during insect feeding for blood meals. Plasmodia are injected as infective sporozoites from the salivary glands during mosquito feeding. *Leishmania* parasites are regurgitated as promastigotes from the alimentary tract of sandflies and injected by bite into a susceptible host. Trypanosomes are first ingested from infected hosts by reduviid bugs; the pathogens then multiply in the gastrointestinal tract of the insects and are released in feces onto the host's skin during subsequent feedings. Most microbes that land directly on intact skin are destined to die, as survival on the skin or in hair follicles requires resistance to fatty acids, low pH, and other antimicrobial factors on the skin. Once it is damaged (and particularly if it becomes necrotic), the skin can be a major portal of entry and growth for pathogens and elaboration of their toxic products. Burn wound infections and tetanus are clear examples. After animal bites, pathogens resident in the animal's saliva gain access to the victim's tissues through the damaged skin. Rabies is the paradigm for this pathogenic process; rabies virus grows in striated muscle cells at the site of inoculation.

Microbial Adherence Once in or on a host, most microbes must anchor themselves to a tissue or tissue factor; the possible exceptions are organisms that directly enter the bloodstream and multiply there. Specific ligands or adhesins for host receptors constitute a major area of study in the field of microbial pathogenesis. *Adhesins* comprise a wide range of surface structures, not only anchoring the microbe to a tissue and promoting cellular entry where appropriate but also eliciting host responses critical to the pathogenic process (Table 145e-1). Most microbes produce multiple adhesins specific for multiple host receptors. These adhesins are often redundant, are serologically variable, and act additively or synergistically with other microbial factors to promote microbial sticking to host tissues. In addition, some microbes adsorb host proteins onto their surface and utilize the natural host protein receptor for microbial binding and entry into target cells.

VIRAL ADHESINS All viral pathogens must bind to host cells, enter them, and replicate within them. Viral coat proteins serve as the ligands for cellular entry, and more than one ligand-receptor interaction may be needed; for example, HIV utilizes its envelope glycoprotein (gp) 120 to enter host cells by binding both to CD4 and to one of two receptors for chemokines (designated CCR5 and CXCR4). Similarly, the measles virus H glycoprotein binds to both CD46 and the membrane-organizing protein moesin on host cells. The gB and gC proteins on herpes simplex virus bind to heparan sulfate, although this adherence is not essential for entry but rather serves to concentrate virions close to the cell surface; this step is followed by attachment to mammalian cells mediated by the viral gD protein, with subsequent formation of a homotrimer of viral gB protein or a heterodimer of viral gH and gL proteins that permits fusion of the viral envelope with the host cell membrane. Herpes simplex virus can use a number of eukaryotic cell

TABLE 145e-1 EXAMPLES OF MICROBIAL LIGAND-RECEPTOR INTERACTIONS

Microorganism	Type of Microbial Ligand	Host Receptor
Viral Pathogens		
Influenza virus	Hemagglutinin	Sialic acid
Measles virus		
Vaccine strain	Hemagglutinin	CD46/moesin
Wild-type strains	Hemagglutinin	Signaling lymphocytic activation molecule (SLAM)
Human herpesvirus type 6	?	CD46
Herpes simplex virus	Glycoprotein C	Heparan sulfate
HIV	Surface glycoprotein	CD4 and chemokine receptors (CCR5 and CXCR4)
Epstein-Barr virus	Envelope protein	CD21 (CR2)
Adenovirus	Fiber protein	Coxsackie-adenovirus receptor (CAR)
Coxsackievirus	Viral coat proteins	CAR and major histocompatibility class I antigens
Bacterial Pathogens		
<i>Neisseria</i> spp.	Pili	Membrane cofactor protein (CD46)
<i>Pseudomonas aeruginosa</i>		
	Pili and flagella	Asialo-GM1
	Lipopolysaccharide	Cystic fibrosis transmembrane conductance regulator (CFTR)
<i>Escherichia coli</i>	Pili	Ceramides/mannose and digalactosyl residues
<i>Streptococcus pyogenes</i>	Hyaluronic acid capsule	CD44
<i>Yersinia</i> spp.	Invasin/accessory invasin locus	β_1 Integrins
<i>Bordetella pertussis</i>	Filamentous hemagglutinin	CR3
<i>Legionella pneumophila</i>	Adsorbed C3bi	CR3
<i>Mycobacterium tuberculosis</i>	Adsorbed C3bi	CR3; DC-SIGN ^a
Fungal Pathogens		
<i>Blastomyces dermatitidis</i>	WI-1	Possibly matrix proteins and integrins
<i>Candida albicans</i>	Int1p	Extracellular matrix proteins
Protozoal Pathogens		
<i>Plasmodium vivax</i>	Merozoite form	Duffy Fy antigen
<i>Plasmodium falciparum</i>	Erythrocyte-binding protein 175 (EBA-175)	Glycophorin A
<i>Entamoeba histolytica</i>	Surface lectin	N-Acetylglucosamine

^aA novel dendritic cell-specific C-type lectin.

surface receptors for entry, including the herpesvirus entry mediator (related to the tumor necrosis factor receptor), members of the immunoglobulin superfamily, the proteins nectin-1 and nectin-2, and modified heparan sulfate.

BACTERIAL ADHESINS Among the microbial adhesins studied in greatest detail are bacterial pili and flagella (Fig. 145e-1). *Pili* or *fimbriae* are commonly used by gram-negative bacteria for attachment to host cells and tissues; studies have identified similar factors produced by gram-positive organisms such as group B streptococci. In electron micrographs, these hairlike projections (up to several hundred per cell) may be confined to one end of the organism (polar pili) or distributed more evenly over the surface. An individual cell may have pili with a variety of functions. Most pili are made up of a major pilin protein subunit

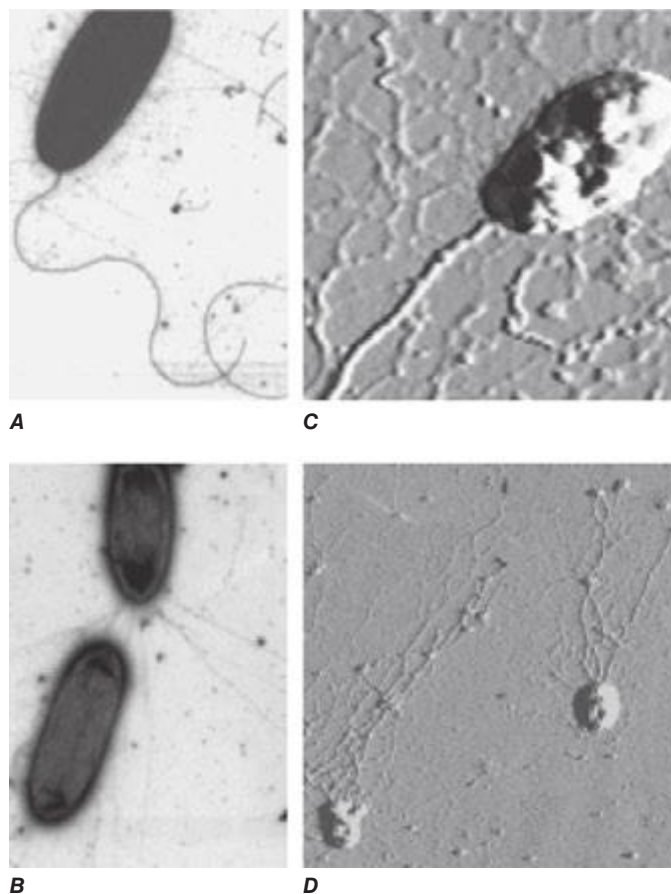


FIGURE 145e-1 Bacterial surface structures. **A** and **B**. Traditional electron micrographic images of fixed cells of *Pseudomonas aeruginosa*. Flagella (**A**) and pili (**B**) project out from the bacterial poles. **C** and **D**. Atomic force microscopic image of live *P. aeruginosa* freshly planted onto a smooth mica surface. This technology reveals the fine, three-dimensional detail of the bacterial surface structures. (Images courtesy of Drs. Martin Lee and Milan Bajmoczy, Harvard Medical School.)

(molecular weight, 17,000–30,000) that polymerizes to form the pilus. Many strains of *Escherichia coli* isolated from urinary tract infections express mannose-binding type 1 pili, whose binding to integral membrane glycoproteins called *uropilins* that coat the cells in the bladder epithelium is inhibited by D-mannose. Other strains produce the Pap (pyelonephritis-associated) or P pilus adhesin that mediates binding to digalactose (gal-gal) residues on globosides of the human P blood groups. Both of these types of pili have proteins located at the tips of the main pilus unit that are critical to the binding specificity of the whole pilus unit. Although immunization with the mannose-binding tip protein (FimH) of type 1 pili prevents experimental *E. coli* bladder infections in mice and monkeys, a human trial of this vaccine was not successful. *E. coli* cells causing diarrheal disease express pilus-like receptors for enterocytes on the small bowel, along with other receptors termed *colonization factors*.

The type IV pilus, a common type of pilus found in *Neisseria* species, *Moraxella* species, *Vibrio cholerae*, *Legionella pneumophila*, *Salmonella enterica* serovar Typhi, enteropathogenic *E. coli*, and *Pseudomonas aeruginosa*, often mediates adherence of organisms to target surfaces. Type IV pili tend to have a relatively conserved aminoterminal region and a more variable carboxyl-terminal region. For some species (e.g., *N. gonorrhoeae*, *Neisseria meningitidis*, and enteropathogenic *E. coli*), the pili are critical for attachment to mucosal epithelial cells. For others, such as *P. aeruginosa*, the pili only partially mediate the cells' adherence to host tissues and may in some circumstances inhibit colonization. For example, a recent study of *P. aeruginosa* colonization of the gastrointestinal tract of mice evaluated a bank of mutants in which all nonessential genes were interrupted; those mutants that were

unable to produce the type IVa pili were actually better able to colonize the gastrointestinal mucosa, although the basis for this observation was not identified. *V. cholerae* cells appear to use two different types of pili for intestinal colonization. Whereas interference with this stage of colonization would appear to be an effective antibacterial strategy, attempts to develop pilus-based vaccines for human diseases have not been highly successful to date.

Flagella are long appendages attached at either one or both ends of the bacterial cell (polar flagella) or distributed over the entire cell surface (peritrichous flagella). Flagella, like pili, are composed of a polymerized or aggregated basic protein. In flagella, the protein subunits form a tight helical structure and vary serologically with the species. Spirochetes such as *T. pallidum* and *Borrelia burgdorferi* have axial filaments similar to flagella running down the long axis of the center of the cell, and they “swim” by rotation around these filaments. Some bacteria can glide over a surface in the absence of obvious motility structures.

Other bacterial structures involved in adherence to host tissues include specific staphylococcal and streptococcal proteins that bind to human extracellular matrix proteins such as fibrin, fibronectin, fibrinogen, laminin, and collagen. Fibronectin appears to be a commonly used receptor for various pathogens; a particular amino acid sequence in fibronectin, Arg-Gly-Asp or RGD, is a critical target used by bacteria to bind to host tissues. Binding of a highly conserved *Staphylococcus aureus* surface protein, clumping factor A (ClfA), to fibrinogen has been implicated in many aspects of pathogenesis. Attempts to interrupt this interaction and prevent *S. aureus* sepsis in low-birth-weight infants by administering an intravenous IgG preparation derived from the plasma of individuals with high titers of antibody to ClfA failed to show efficacy in a clinical trial; however, this approach is being pursued in some vaccine formulations targeting this organism. The conserved outer-core portion of the lipopolysaccharide (LPS) of *P. aeruginosa* mediates binding to the cystic fibrosis transmembrane conductance regulator (CFTR) on airway epithelial cells—an event that appears to play a critical role in normal host resistance to infection by initiating recruitment of polymorphonuclear neutrophils (PMNs) to the lung mucosa to kill the cells via opsonophagocytosis. A large number of microbial pathogens encompassing major gram-positive bacteria (staphylococci and streptococci), gram-negative bacteria (major enteric species and coccobacilli), fungi (*Candida*, *Fusobacterium*, *Aspergillus*), and even eukaryotes (*Trichomonas vaginalis* and *Plasmodium falciparum*) express a surface polysaccharide composed of β -1-6-linked-poly-*N*-acetyl-D-glucosamine (PNAG). One of the functions of PNAG for some of these organisms is to promote binding to materials used in catheters and other types of implanted devices. This polysaccharide may be a critical factor in the establishment of device-related infections by pathogens such as staphylococci and *E. coli*. High-powered imaging techniques (e.g., atomic force microscopy) have revealed that bacterial cells have a nonhomogeneous surface that is probably attributable to different concentrations of cell surface molecules, including microbial adhesins, at specific places on the cell surface (Figs. 120-1C and 120-1D).

FUNGAL ADHESINS Several fungal adhesins have been described that mediate colonization of epithelial surfaces, particularly adherence to structures like fibronectin, laminin, and collagen. The product of the *Candida albicans* *INT1* gene, Int1p, bears similarity to mammalian integrins that bind to extracellular matrix proteins. The agglutinin-like sequence (ALS) adhesins are large cell-surface glycoproteins mediating adherence of pathogenic *Candida* to host tissues. These adhesins possess a conserved three-domain structure composed of an N-terminal domain that mediates adherence to host tissue receptors, a central motif consisting of a number of repeats of a conserved sequence of 36 amino acids, and a C-terminal domain that varies in length and sequence and contains a glycosylphosphatidylinositol (GPI) anchor addition site that allows binding of the adhesin to the fungal cell wall. Variability in the number of central domains in different ALS proteins characterizes different adhesins with specificity for different host receptors. The ALS adhesins are expressed under certain environmental conditions and are crucial for pathogenesis of fungal infections.

For several fungal pathogens that initiate infections after inhalation of infectious material, the inoculum is ingested by alveolar macrophages, in which the fungal cells transform to pathogenic phenotypes. Like *C. albicans*, *Blastomyces dermatitidis* binds to CD11b/CD18 integrins as well as to CD14 on macrophages. *B. dermatitidis* produces a 120-kDa surface protein, designated WI-1, that mediates this adherence. An unidentified factor on *Histoplasma capsulatum* also mediates binding of this fungal pathogen to the integrin surface proteins.

EUKARYOTIC PATHOGEN ADHESINS Eukaryotic parasites use complicated surface glycoproteins as adhesins, some of which are lectins (proteins that bind to specific carbohydrates on host cells). For example, *Plasmodium vivax*, one of six *Plasmodium* species causing malaria, binds (via Duffy-binding protein) to the Duffy blood group carbohydrate antigen Fy on erythrocytes. *Entamoeba histolytica*, the third leading cause of death from parasitic diseases, expresses two proteins that bind to the disaccharide galactose/*N*-acetyl galactosamine. Reports indicate that children with mucosal IgA antibody to one of these lectins are resistant to reinfection with virulent *E. histolytica*. A major surface glycoprotein (gp63) of *Leishmania* promastigotes is needed for these parasites to enter human macrophages—the principal target cell of infection. This glycoprotein promotes complement binding but inhibits complement lytic activity, allowing the parasite to use complement receptors for entry into macrophages; gp63 also binds to fibronectin receptors on macrophages. In addition, the pathogen can express a carbohydrate that mediates binding to host cells. Evidence suggests that, as part of hepatic granuloma formation, *Schistosoma mansoni* expresses a carbohydrate epitope related to the Lewis X blood group antigen that promotes adherence of helminthic eggs to vascular endothelial cells under inflammatory conditions.

Host Receptors Host receptors are found both on target cells (such as epithelial cells lining mucosal surfaces) and within the mucus layer covering these cells. Microbial pathogens bind to a wide range of host receptors to establish infection (Table 145e-1). Selective loss of host receptors for a pathogen may confer natural resistance to an otherwise susceptible population. For example, 70% of individuals in West Africa lack Fy antigens and are resistant to *P. vivax* infection. *S. enterica* serovar Typhi, the etiologic agent of typhoid fever, produces a pilus protein that binds to CFTR to enter the gastrointestinal submucosa after being ingested by enterocytes. As homozygous mutations in *CFTR* are the cause of the life-shortening disease cystic fibrosis, heterozygote carriers (e.g., 4–5% of individuals of European ancestry) may have had a selective advantage due to decreased susceptibility to typhoid fever.

Numerous virus–target cell interactions have been described, and it is now clear that different viruses can use similar host cell receptors for entry. The list of certain and likely host receptors for viral pathogens is long. Among the host membrane components that can serve as receptors for viruses are sialic acids, gangliosides, glycosaminoglycans, integrins and other members of the immunoglobulin superfamily, histocompatibility antigens, and regulators and receptors for complement components. A notable example of the effect of host receptors on the pathogenesis of infection has emerged from studies comparing the binding of avian influenza A subtype H5N1 with that of influenza A strains expressing the H1 subtype of hemagglutinin. The H1 subtypes tend to be highly pathogenic and transmissible from human to human, and they bind to a receptor composed of two sugar molecules: sialic acid linked α -2-6 to galactose. This receptor is expressed at high levels in the airway epithelium; when virus is shed from this surface, its transmission via coughing and aerosol droplets is facilitated. In contrast, the H5N1 avian influenza virus binds to sialic acid linked α -2-3 to galactose, and this receptor is expressed at high levels in pneumocytes in the alveoli. Infection in the alveoli is thought to underlie the high mortality rate associated with avian influenza but also the low interhuman transmissibility of this strain, which is not readily transported to the airways from which it can be expelled by coughing. Nonetheless, it was recently shown that H5 hemagglutinins can acquire mutations that vastly increase their transmissibility while not affecting their high level of lethality.

Once established on a mucosal or skin site, pathogenic microbes must replicate before causing full-blown infection and disease. Within cells, viral particles release their nucleic acids, which may be directly translated into viral proteins (positive-strand RNA viruses), transcribed from a negative strand of RNA into a complementary mRNA (negative-strand RNA viruses), or transcribed into a complementary strand of DNA (retroviruses); for DNA viruses, mRNA may be transcribed directly from viral DNA, either in the cell nucleus or in the cytoplasm. To grow, bacteria must acquire specific nutrients or synthesize them from precursors in host tissues. Many infectious processes are usually confined to specific epithelial surfaces—e.g., H1 subtype influenza to the respiratory mucosa, gonorrhea to the urogenital epithelium, shigellosis to the gastrointestinal epithelium. While there are multiple reasons for this specificity, one important consideration is the ability of these pathogens to obtain from these specific environments the nutrients needed for growth and survival.

Temperature restrictions also play a role in limiting certain pathogens to specific tissues. Rhinoviruses, a cause of the common cold, grow best at 33°C and replicate in cooler nasal tissues but not in the lung. Leprosy lesions due to *Mycobacterium leprae* are found in and on relatively cool body sites. Fungal pathogens that infect the skin, hair follicles, and nails (dermatophyte infections) remain confined to the cooler, exterior, keratinous layer of the epithelium.

A topic of major interest is the ability of many bacterial, fungal, and protozoal species to grow in multicellular masses referred to as *biofilms*. These masses are biochemically and morphologically quite distinct from the free-living individual cells referred to as *planktonic cells*. Growth in biofilms leads to altered microbial metabolism, production of extracellular virulence factors, and decreased susceptibility to biocides, antimicrobial agents, and host defense molecules and cells. *P. aeruginosa* growing on the bronchial mucosa during chronic infection, staphylococci and other pathogens growing on implanted medical devices, and dental pathogens growing on tooth surfaces to form plaque are several examples of microbial biofilm growth associated with human disease. Many other pathogens can form biofilms during *in vitro* growth. It is increasingly accepted that this mode of growth contributes to microbial virulence and induction of disease and that biofilm formation can also be an important factor in microbial survival outside the host, promoting transmission to additional susceptible individuals.

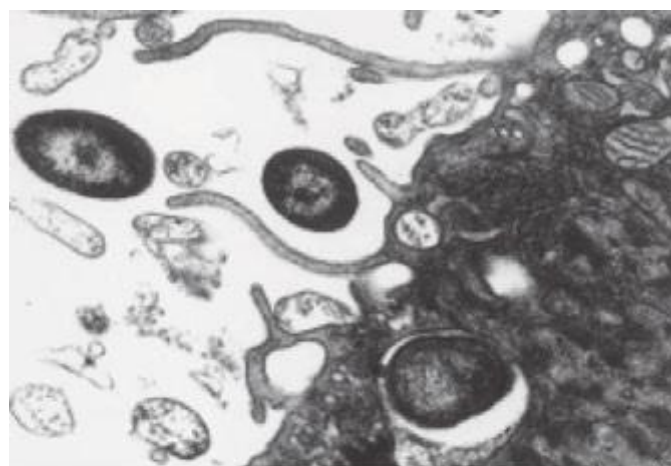
AVOIDANCE OF INNATE HOST DEFENSES

As microbes have interacted with mucosal/epithelial surfaces since the emergence of multicellular organisms, it is not surprising that multicellular hosts have a variety of innate surface defense mechanisms that can sense when pathogens are present and contribute to their elimination. The skin is acidic and is bathed with fatty acids toxic to many microbes. Skin pathogens such as staphylococci must tolerate these adverse conditions. Mucosal surfaces are covered by a barrier composed of a thick mucus layer that entraps microbes and facilitates their transport out of the body by such processes as mucociliary clearance, coughing, and urination. Mucous secretions, saliva, and tears contain antibacterial factors such as lysozyme and antimicrobial peptides as well as antiviral factors such as interferons (IFNs). Gastric acidity and bile salts are inimical to the survival of many ingested pathogens, and most mucosal surfaces—particularly the nasopharynx, the vaginal tract, and the gastrointestinal tract—contain a resident flora of commensal microbes that interfere with the ability of pathogens to colonize and infect a host. Major advances in the use of nucleic acid sequencing now allow extensive identification and characterization of the vast array of commensal organisms that have come to be referred to as the *microbiota*. In addition to its role in providing competition for mucosal colonization, acquisition of a normal microbiota is critical for proper development of the immune system, influencing maturation and differentiation of components of both the innate and acquired arms.

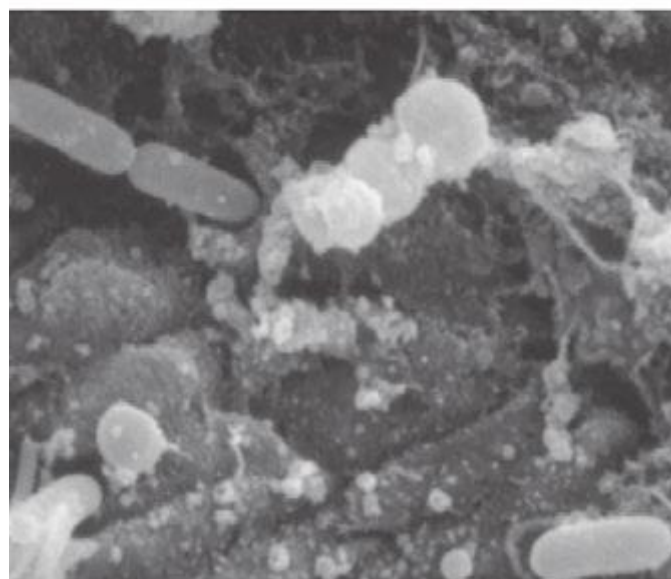
Pathogens that survive local antimicrobial factors must still contend with host endocytic, phagocytic, and inflammatory responses as well as with host genetic factors that determine the degree to which a

pathogen can survive and grow. The list of genes whose variants, usually by single-nucleotide polymorphisms, can affect host susceptibility and resistance to infection is rapidly expanding. A classic example is a 32-bp deletion in the gene for the HIV-1 co-receptor known as *chemokine receptor 5* (CCR5), which, when present in the homozygous state, confers high-level resistance to HIV-1 infection. The growth of viral pathogens entering skin or mucosal epithelial cells can be limited by a variety of host genetic factors, including production of IFNs, modulation of receptors for viral entry, and age- and hormone-related susceptibility factors; by nutritional status; and even by personal habits such as smoking and exercise.

Encounters with Epithelial Cells Over the past two decades, many pathogens have been shown to enter epithelial cells (Fig. 145e-2); they often use specialized surface structures that bind to receptors, with consequent internalization. However, the exact role and the importance of this process in infection and disease are not well defined for most of these pathogens. Microbial entry into host epithelial cells is seen as a means for dissemination to adjacent or deeper tissues or as a route to sanctuary to avoid ingestion and killing by professional



A



B

FIGURE 145e-2 Entry of bacteria into epithelial cells.

A. Internalization of *Pseudomonas aeruginosa* by cultured airway epithelial cells expressing wild-type cystic fibrosis transmembrane conductance regulator, the cell receptor for bacterial ingestion.

B. Entry of *P. aeruginosa* into murine tracheal epithelial cells after murine infection by the intranasal route.

phagocytes. Epithelial cell entry appears, for instance, to be a critical aspect of dysentery induction by *Shigella*.

Curiously, the less virulent strains of many bacterial pathogens are more adept at entering epithelial cells than are more virulent strains; examples include pathogens that lack the surface polysaccharide capsule needed to cause serious disease. Thus, for *Haemophilus influenzae*, *Streptococcus pneumoniae*, *Streptococcus agalactiae* (group B *Streptococcus*), and *Streptococcus pyogenes*, isogenic mutants or variants lacking capsules enter epithelial cells better than the wild-type, encapsulated parental forms that cause disseminated disease. These observations have led to the proposal that epithelial cell entry may be primarily a manifestation of host defense, resulting in bacterial clearance by both shedding of epithelial cells containing internalized bacteria and initiation of a protective and nonpathogenic inflammatory response. However, a possible consequence of this process could be the opening of a hole in the epithelium, potentially allowing uningested organisms to enter the submucosa. This scenario has been documented in murine *S. enterica* serovar Typhimurium infections and in experimental bladder infections with uropathogenic *E. coli*. In the latter system, bacterial pilus-mediated attachment to *uroplakins* induces exfoliation of the cells with attached bacteria. Subsequently, infection is produced by residual bacterial cells that invade the superficial bladder epithelium, where they can grow intracellularly into biofilm-like masses encased in an extracellular polysaccharide-rich matrix and surrounded by uroplakin. This mode of growth produces structures that have been referred to as *bacterial pods*. It is likely that at low bacterial inocula epithelial cell ingestion and subclinical inflammation are efficient means to eliminate pathogens, while at higher inocula a proportion of surviving bacterial cells enter the host tissue through the damaged mucosal surface and multiply, producing disease. Alternatively, failure of the appropriate epithelial cell response to a pathogen may allow the organism to survive on a mucosal surface where, if it avoids other host defenses, it can grow and cause a local infection. Along these lines, as noted above, *P. aeruginosa* is taken into epithelial cells by CFTR, a protein missing or nonfunctional in most severe cases of cystic fibrosis. The major clinical consequence of this disease is chronic airway-surface infection with *P. aeruginosa* in 80–90% of patients. The failure of airway epithelial cells to ingest and promote the removal of *P. aeruginosa* via a properly regulated inflammatory response has been proposed as a key component of the hypersusceptibility of cystic fibrosis patients to chronic airway infection with this organism.

Encounters with Phagocytes • PHAGOCYTOSIS AND INFLAMMATION

Phagocytosis of microbes is a major innate host defense that limits the growth and spread of pathogens. Phagocytes appear rapidly at sites of infection in conjunction with the initiation of inflammation. Ingestion of microbes by both tissue-fixed macrophages and migrating phagocytes probably accounts for the limited ability of most microbial agents to cause disease. A family of related molecules called *collectins*, *soluble defense collagens*, or *pattern-recognition molecules* are found in blood (mannose-binding lectins), in lung (surfactant proteins A and D), and most likely in other tissues as well and bind to carbohydrates on microbial surfaces to promote phagocyte clearance. Bacterial pathogens seem to be ingested principally by PMNs, while eosinophils are frequently found at sites of infection by protozoan or multicellular parasites. Successful pathogens, by definition, must avoid being cleared by professional phagocytes. One of several antiphagocytic strategies employed by bacteria and by the fungal pathogen *Cryptococcus neoformans* is to elaborate large-molecular-weight surface polysaccharide antigens, often in the form of a capsule that coats the cell surface. Most pathogenic bacteria produce such antiphagocytic capsules. On occasion, proteins or polypeptides form capsule-like coatings for organisms such as group A streptococci and *Bacillus anthracis*.

As activation of local phagocytes in tissues is a key step in initiating inflammation and migration of additional phagocytes into infected sites, much attention has been paid to microbial factors that initiate inflammation. These are usually conserved factors critical to the microbes' survival and are referred to as *pathogen-associated molecular patterns* (PAMPs). Cellular responses to microbial encounters with

phagocytes are governed largely by the structure of the microbial PAMPs that elicit inflammation, and detailed knowledge of these structures of bacterial pathogens has contributed greatly to our understanding of molecular mechanisms of microbial pathogenesis mediated by activation of host cell molecules such as TLRs (Fig. 145e-3). One of the best-studied systems involves the interaction of LPS from gram-negative bacteria and the GPI-anchored membrane protein CD14 found on the surface of professional phagocytes, including migrating and tissue-fixed macrophages and PMNs. A soluble form of CD14 is also found in plasma and on mucosal surfaces. A plasma protein, LPS-binding protein, transfers LPS to membrane-bound CD14 on myeloid cells and promotes binding of LPS to soluble CD14. Soluble CD14/LPS/LPS-binding protein complexes bind to many cell types and may be internalized to initiate cellular responses to microbial pathogens. It has been shown that peptidoglycan and lipoteichoic acid from gram-positive bacteria as well as cell-surface products of mycobacteria and spirochetes can interact with CD14 (Fig. 145e-3). Additional molecules, such as MD-2, also participate in the recognition of bacterial activators of inflammation.

GPI-anchored receptors do not have intracellular signaling domains; therefore, it is the TLRs that transduce signals for cellular activation due to LPS binding. Binding of microbial factors to TLRs to activate signal transduction occurs in the phagosome—and not on the surface—of dendritic cells that have internalized the microbe. This binding is probably due to the release of the microbial surface factor from the cell in the environment of the phagosome, where the liberated factor can bind to its cognate TLRs. TLRs initiate cellular activation through a series of signal-transducing molecules (Fig. 145e-3) that lead to nuclear translocation of the transcription factor NF- κ B (nuclear factor κ B), a master-switch for production of important inflammatory cytokines such as tumor necrosis factor α (TNF- α) and interleukin (IL) 1.

The initiation of inflammation can occur not only with LPS and peptidoglycan but also with viral particles and other microbial products such as polysaccharides, enzymes, and toxins. Bacterial flagella activate inflammation by binding of a conserved sequence to TLR5. Some pathogens (e.g., *Campylobacter jejuni*, *Helicobacter pylori*, and *Bartonella bacilliformis*) make flagella that lack this sequence and do not bind to TLR5; thus efficient host responses to infection are prevented. Bacteria also produce a high proportion of DNA molecules with unmethylated CpG residues that activate inflammation through TLR9. TLR3 recognizes double-stranded RNA, a pattern-recognition molecule produced by many viruses during their replicative cycle. TLR1 and TLR6 associate with TLR2 to promote recognition of acylated microbial proteins and peptides.

The myeloid differentiation factor 88 (MyD88) molecule and the Toll/IL-1R (TIR) domain-containing adapter protein (TIRAP) bind to the cytoplasmic domains of TLRs and also to receptors that are part of the IL-1 receptor families. Numerous studies have shown that MyD88/TIRAP-mediated transduction of signals from TLRs and other receptors is critical for innate resistance to infection, activating MAP-kinases and NF- κ B and thereby leading to production of cytokines/chemokines. Mice lacking MyD88 are more susceptible than normal mice to infections with a broad range of pathogens. In one study, nine children homozygous for defective MyD88 genes had recurrent infections with *S. pneumoniae*, *S. aureus*, and *P. aeruginosa*—three bacterial species showing increased virulence in MyD88-deficient mice; however, unlike these mice, the MyD88-deficient children seemed to have no greater susceptibility to other bacteria, viruses, fungi, or parasites. Another component of the MyD88-dependent signaling pathway is a molecule known as IL-1 receptor-associated kinase 4 (IRAK-4). Individuals with a homozygous deficiency in genes encoding this protein are at increased risk for *S. pneumoniae* and *S. aureus* infections and, to some degree, for *P. aeruginosa* infections as well.

In addition to their role in MyD88-mediated signaling, some TLRs (e.g., TLR3 and TLR4) can activate signal transduction via a MyD88-independent pathway involving TIR domain-containing, adapter-inducing IFN- β (TRIF) and the TRIF-related adapter molecule (TRAM). Signaling through TRIF and TRAM activates the production of both NF- κ B-dependent cytokines/chemokines and type 1 IFNs.

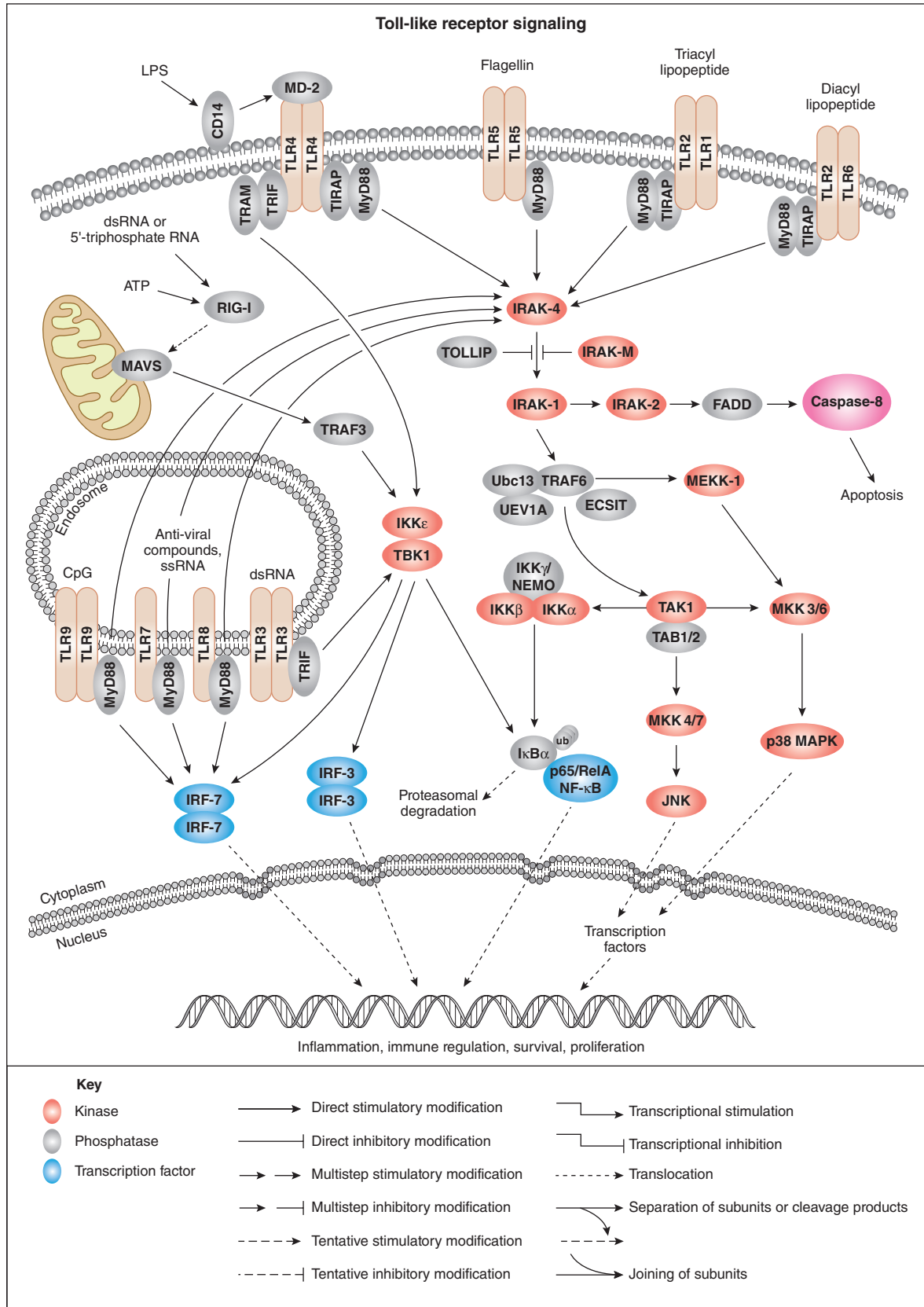


FIGURE 145e-3 Cellular signaling pathways for production of inflammatory cytokines in response to microbial products. Microbial cell-surface constituents interact with Toll-like receptors (TLRs), in some cases requiring additional factors such as MD-2, which facilitates the response to lipopolysaccharide (LPS) via TLR4. Although these constituents are depicted as interacting with the TLRs on the cell surface, TLRs contain extracellular leucine-rich domains that become localized to the lumen of the phagosome upon uptake of bacterial cells. The internalized TLRs can bind to microbial products. The TLRs are oligomerized, usually forming homodimers, and then bind to the general adapter protein MyD88 via the C-terminal Toll/IL-1R (TIR) domains, which also bind to TIRAP (TIR domain-containing adapter protein), a molecule that participates in the transduction of signals from TLRs 1, 2, 4, and 6. The MyD88/TIRAP complex activates signal-transducing molecules such as IRAK-4 (IL-1Rc-associated kinase 4), which in turn activates IRAK-1. This activation can be blocked by IRAK-M and Toll-interacting protein (TOLLIP). IRAK-1 activates TRAF6 (tumor necrosis factor receptor-associated factor 6), TAK1 (transforming growth factor β -activating kinase 1), and TAB1/2

The type 1 IFNs bind to the IFN- α receptor composed of two protein chains, IFNAR1 and IFNAR2. Humans produce three type 1 IFNs: IFN- α , IFN- β , and IFN- γ . These molecules activate another class of proteins known as the signal transducer and activator of transcription (STAT) complexes. The STAT factors are important in regulating immune system genes and thus play a critical role in responding to microbial infections.

Another intracellular complex of proteins found to be a major factor in the host cell response to infection is the *inflammasome* (Fig. 145e-4), in which inflammatory cytokines IL-1 and IL-18 are changed from their precursor to their active forms prior to secretion by the cysteine protease caspase-1. Within the inflammasome are additional proteins that are members of the nucleotide binding and oligomerization domain (NOD)-like receptor (NLR) family. Like the TLRs, NOD proteins sense the presence of the conserved microbial factors released inside a cell. Recognition of these PAMPs by NLRs leads to caspase-1 activation and to secretion of active IL-1 and IL-18 by an unknown mechanism. Studies of mice indicate that as many as four inflammasomes with different components are formed: the IPAF inflammasome, the NALP1 inflammasome, the cryopyrin/NALP3 inflammasome, and an inflammasome triggered by *Francisella tularensis* infection (Fig. 145e-4). The components depend on the type of stimulus driving inflammasome formation and activation.

A recent addition to the identified intracellular components responding to microbial infection is *autophagy*, initially described as an intracellular process for degradation and recycling of cellular components for reuse. Now it is clear that autophagy constitutes an early defense mechanism in which, after ingestion, microbial pathogens either within vacuoles or in the cytoplasm are delivered to lysosomal compartments for degradation. Avoidance of this process is critical if pathogens are to cause disease and can be achieved by multiple mechanisms, such as inhibition of proteins within the autophagic vacuole by shigellae, recruitment of host proteins to mask *Listeria monocytogenes*, and inhibition of formation of the vacuole by *L. pneumophila*.

ADDITIONAL INTERACTIONS OF MICROBIAL PATHOGENS AND PHAGOCYTES Other ways that microbial pathogens avoid destruction by phagocytes include production of factors that are toxic to these cells or that interfere with their chemotactic and ingestion function. Hemolysins, leukocidins, and the like are microbial proteins that can kill phagocytes that are attempting to ingest organisms elaborating these substances. For example, *S. aureus* elaborates a family of bicomponent leukocidins that bind to host receptors such as the HIV co-receptor CCR5 (which is also used by the Luke/D toxin) or—in the case of the Pantan-Valentine leukocidin—the receptor of the C5a component of activated complement (which is used by LukF/S). Streptolysin O made by *S. pyogenes* binds to cholesterol in phagocyte membranes and initiates a process of internal degranulation, with the release of normally granule-sequestered toxic components into the phagocyte's cytoplasm. *E. histolytica*, an intestinal protozoan that causes amebic dysentery, can disrupt phagocyte membranes after direct contact via the release of protozoal phospholipase A and pore-forming peptides.

MICROBIAL SURVIVAL INSIDE PHAGOCYTES Many important microbial pathogens use a variety of strategies to survive inside phagocytes (particularly macrophages) after ingestion. Inhibition of fusion of the phagocytic vacuole (the phagosome) containing the ingested microbe with the lysosomal granules containing antimicrobial substances (the lysosome) allows *Mycobacterium tuberculosis*, *S. enterica* serovar Typhi, and *Toxoplasma gondii* to survive inside macrophages. Some organisms, such as *L. monocytogenes*, escape into the phagocyte's cytoplasm to grow and eventually spread to other cells. Resistance to killing within the macrophage and subsequent growth are critical to successful infection by herpes-type viruses, measles virus, poxviruses, *Salmonella*, *Yersinia*, *Legionella*, *Mycobacterium*, *Trypanosoma*, *Nocardia*, *Histoplasma*, *Toxoplasma*, and *Rickettsia*. *Salmonella* species use a master regulatory system—in which the *PhoP/PhoQ* genes control other genes—to enter and survive within cells, with intracellular survival entailing structural changes in the cell envelope LPS.

TISSUE INVASION AND TISSUE TROPISM

Tissue Invasion Most viral pathogens cause disease by growth at skin or mucosal entry sites, but some pathogens spread from the initial site to deeper tissues. Virus can spread via the nerves (rabies virus) or plasma (picornaviruses) or within migratory blood cells (poliovirus, Epstein-Barr virus, and many others). Specific viral genes determine where and how individual viral strains can spread.

Bacteria may invade deeper layers of mucosal tissue via intracellular uptake by epithelial cells, traversal of epithelial cell junctions, or penetration through denuded epithelial surfaces. Among virulent *Shigella* strains and invasive strains of *E. coli*, outer-membrane proteins are critical to epithelial cell invasion and bacterial multiplication. *Neisseria* and *Haemophilus* species penetrate mucosal cells by poorly understood mechanisms before dissemination into the bloodstream. Staphylococci and streptococci elaborate a variety of extracellular enzymes, such as hyaluronidase, lipases, nucleases, and hemolysins, that are probably important in breaking down cellular and matrix structures and allowing the bacteria access to deeper tissues and blood. For example, staphylococcal α -hemolysin binds to a receptor, A-disintegrin and metalloprotease 10 (ADAM-10), to cause endothelial cell damage and disruption of vascular barrier function—events that are likely critical for systemic spread of *S. aureus* from an initial infectious site. Organisms that colonize the gastrointestinal tract can often translocate through the mucosa into the blood and, under circumstances in which host defenses are inadequate, cause bacteremia. *Yersinia enterocolitica* can invade the mucosa through the activity of the invasin protein. The complex milieu of the basement membrane-containing structures, such as laminin and collagen, that anchor epithelial cells to mucosal surfaces must often be breached. Numerous organisms express factors known as MSCRAMMs (microbial surface components recognizing adhesive matrix molecules). These MSCRAMMs promote bacterial attachment to factors in the host extracellular matrix, such as laminin, collagen, and fibronectin. Additional microbial proteases, along with the host's own surface-bound

(TAK1-binding protein 1/2). This signaling complex associates with the ubiquitin-conjugating enzyme Ubc13 and the Ubc-like protein UEV1A to catalyze the formation of a polyubiquitin chain on TRAF6. Polyubiquitination of TRAF6 activates TAK1, which, along with TAB1/2 (a protein that binds to lysine residue 63 in polyubiquitin chains via a conserved zinc-finger domain), phosphorylates the inducible kinase complex: IKK α , IKK β , and IKK γ . IKK γ is also called NEMO (nuclear factor κ B [NF- κ B] essential modulator). This large complex phosphorylates the inhibitory component of NF- κ B, I κ B α , resulting in release of I κ B α from NF- κ B. Phosphorylated (PP) I κ B is then ubiquitinated (ub) and degraded, and the two components of NF- κ B, p50 or Rel and p65, translocate to the nucleus, where they bind to regulatory transcriptional sites on target genes, many of which encode inflammatory proteins. In addition to inducing NF- κ B nuclear translocation, the TAK1/TAB1/2 complex activates MAP kinase transducers such as MKK 4/7 and MKK 3/6, which can lead to nuclear translocation of transcription factors such as AP1. TLR4 can also activate NF- κ B nuclear translocation via the MyD88-independent TRIF (TIR domain-containing adapter-inducing IFN- β) and TRAM (TRIF-related adapter molecule) cofactors. Intracellular TLRs 3, 7, 8, and 9 also use MyD88 and TRIF to activate IFN response factors 3 and 7 (IRF-3 and IRF-7), which also function as transcriptional factors in the nucleus. ATP, adenosine 5'-triphosphate; ECSIT, evolutionarily conserved signaling intermediate in Toll pathways; FADD, Fas-associated protein with death domain; JNK, c-Jun N-terminal kinase; MAVS, mitochondrial antiviral signaling protein; MEKK-1, MAP/ERK kinase kinase 1; p38 MAPK, p38 mitogen-activated protein kinase; RIG-1, retinoic acid-inducible gene 1; TBK1, TANK-binding kinase 1. (Pathway diagram reproduced courtesy of Cell Signaling Technology, Inc. [www.cellsignal.com].)

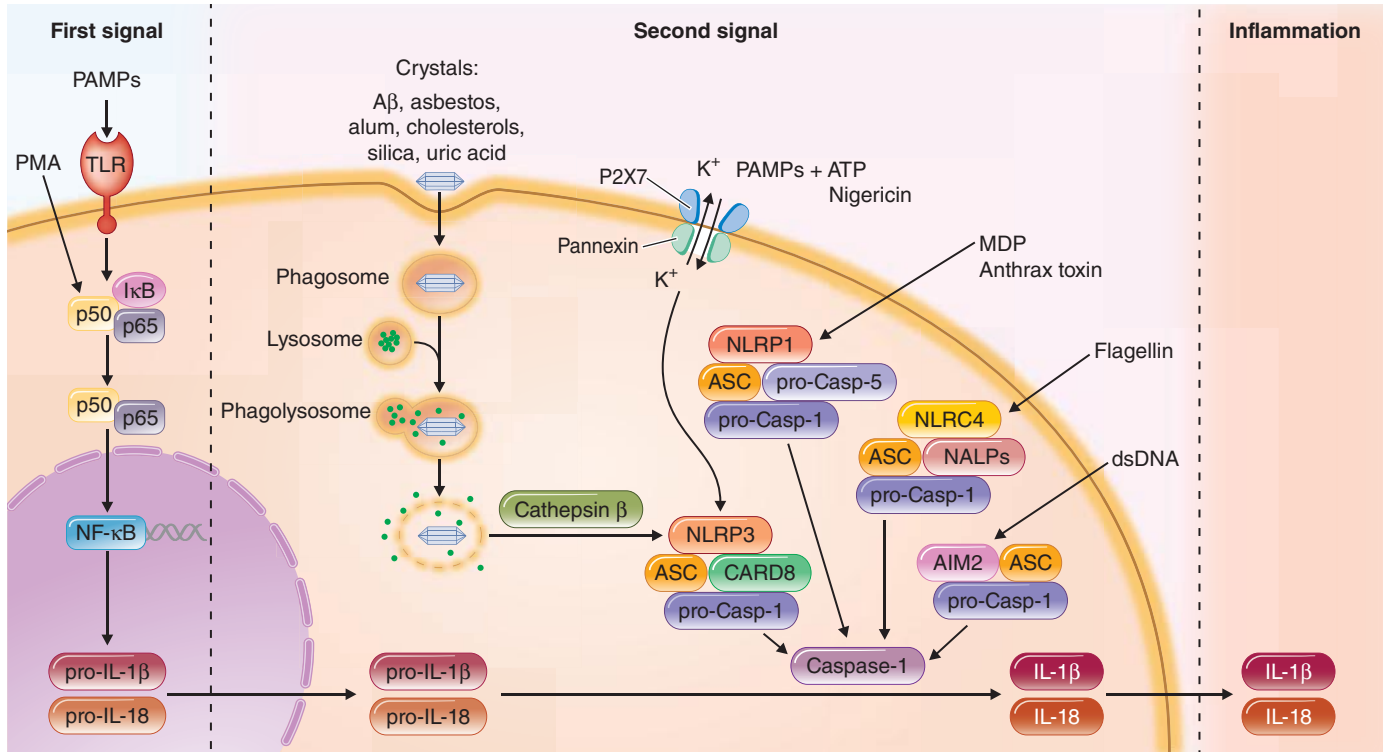


FIGURE 145e-4 Inflammasomes. The nucleotide-binding oligomerization domain-like receptor (NLR) family of proteins is involved in the regulation of innate immune responses. These proteins sense pathogen-associated molecular patterns (PAMPs) in the cytosol as well as the host-derived signals known as damage-associated molecular patterns (DAMPs). Certain NLRs induce the assembly of large caspase-1-activating complexes called *inflammasomes*. Activation of caspase-1 through autoproteolytic maturation leads to the processing and secretion of the proinflammatory cytokines interleukin 1 β (IL-1 β) and IL-18. So far, four inflammasomes have been identified and defined by the NLR protein that they contain: the NLRP1/NALP1b inflammasome; the NLRC4/IPAF inflammasome; the NLRP3/NALP3 inflammasome; and the AIM2 (absent in melanoma 2)-containing inflammasome. A β , amyloid β ; ASC, apoptosis-associated speck-like protein containing CARD; ATP, adenosine 5'-triphosphate; CARD8, caspase recruitment domain-containing protein 8; I κ B, inhibitor of κ B; IPAF, interleukin-converting enzyme protease-activating factor; MDP, muramyl dipeptide; NF- κ B, nuclear factor κ B; P2X7, purinergic P2X7 (receptor); PMA, phorbol myristate acetate; TLR, Toll-like receptor. (Pathway diagram reproduced with permission from Invivogen [www.invivogen.com/review-inflammasome].)

plasminogen and host matrix metalloproteases, then combine to degrade the extracellular matrix and promote microbial spread. Some bacteria (e.g., brucellae) can be carried from a mucosal site to a distant site by phagocytic cells that ingest but fail to kill the bacteria.

Fungal pathogens almost always take advantage of host immunocompromise to spread hematogenously to deeper tissues. The AIDS epidemic has resoundingly illustrated this principle: the immunodeficiency of many HIV-infected patients permits the development of life-threatening fungal infections of the lung, blood, and brain. Other than the capsule of *C. neoformans*, specific fungal antigens involved in tissue invasion are not well characterized. Both fungal pathogens and protozoal pathogens (e.g., *Plasmodium* species and *E. histolytica*) undergo morphologic changes to spread within a host. *C. albicans* undertakes a yeast-hyphal transformation wherein the hyphal forms are found where the fungus is infiltrating the mucosal barrier of tissues, while the yeast form grows on epithelial cell surfaces as well as on the tips of hyphae that have infiltrated tissues. Malarial parasites grow in liver cells as merozoites and are released into the blood to invade erythrocytes and become trophozoites. *E. histolytica* is found as both a cyst and a trophozoite in the intestinal lumen, through which this pathogen enters the host, but only the trophozoite form can spread systemically to cause amebic liver abscesses. Other protozoal pathogens, such as *T. gondii*, *Giardia lamblia*, and *Cryptosporidium*, also undergo extensive morphologic changes after initial infection to spread to other tissues.

Tissue Tropism The propensity of certain microbes to cause disease by infecting specific tissues has been known since the early days of bacteriology, yet the molecular basis for this propensity is understood somewhat better for viral pathogens than for other agents of infectious

disease. Specific receptor-ligand interactions clearly underlie the ability of certain viruses to enter cells within tissues and disrupt normal tissue function, but the mere presence of a receptor for a virus on a target tissue is not sufficient for tissue tropism. Factors in the cell, route of viral entry, viral capacity to penetrate into cells, viral genetic elements that regulate gene expression, and pathways of viral spread in a tissue all affect tissue tropism. Some viral genes are best transcribed in specific target cells, such as hepatitis B genes in liver cells and Epstein-Barr virus genes in B lymphocytes. The route of inoculation of poliovirus determines its neurotropism, although the molecular basis for this circumstance is not understood.

Compared with viral tissue tropism, the tissue tropism of bacterial and parasitic infections has not been as clearly elucidated, but studies of *Neisseria* species have provided insights. Both *N. gonorrhoeae*, which colonizes and infects the human genital tract, and *N. meningitidis*, which principally colonizes the human oropharynx but can spread to the brain, produce type IV pili (Tfp) that mediate adherence to host tissues. In the case of *N. gonorrhoeae*, the Tfp bind to a glucosamine-galactose-containing adhesin on the surface of cervical and urethral cells; in the case of *N. meningitidis*, the Tfp bind to cells in the human meninges and thus cross the blood-brain barrier. *N. meningitidis* expresses a capsular polysaccharide, while *N. gonorrhoeae* does not; however, there is no indication that this property plays a role in the different tissue tropisms displayed by these two bacterial species. *N. gonorrhoeae* can use cytidine monophosphate *N*-acetylneuraminic acid from host tissues to add *N*-acetylneuraminic acid (sialic acid) to its lipooligosaccharide O side chain, and this alteration appears to make the organism resistant to host defenses. Lactate, present at high levels on genital mucosal surfaces, stimulates sialylation of gonococcal lipooligosaccharide. Bacteria with sialic acid sugars in their capsules,

such as *N. meningitidis*, *E. coli* K1, and group B streptococci, have a propensity to cause meningitis, but this generalization has many exceptions. For example, all recognized serotypes of group B streptococci contain sialic acid in their capsules, but only one serotype (III) is responsible for most cases of group B streptococcal meningitis. Moreover, both *H. influenzae* and *S. pneumoniae* can readily cause meningitis, but these organisms do not have sialic acid in their capsules.

TISSUE DAMAGE AND DISEASE

Disease is a complex phenomenon resulting from tissue invasion and destruction, toxin elaboration, and host response. Viruses cause much of their damage by exerting a cytopathic effect on host cells and inhibiting host defenses. The growth of bacterial, fungal, and protozoal parasites in tissue, which may or may not be accompanied by toxin elaboration, can compromise tissue function and lead to disease. For some bacterial and possibly some fungal pathogens, toxin production is one of the best-characterized molecular mechanisms of pathogenesis, while host factors such as IL-1, TNF- α , kinins, inflammatory proteins, products of complement activation, and mediators derived from arachidonic acid metabolites (leukotrienes) and cellular degranulation (histamines) readily contribute to the severity of disease.

Viral Disease Viral pathogens are well known to inhibit host immune responses by a variety of mechanisms. Immune responses can be affected by decreasing production of most major histocompatibility complex molecules (adenovirus E3 protein), by diminishing cytotoxic T cell recognition of virus-infected cells (Epstein-Barr virus EBNA1 antigen and cytomegalovirus IE protein), by producing virus-encoded complement receptor proteins that protect infected cells from complement-mediated lysis (herpesvirus and vaccinia virus), by making proteins that interfere with the action of IFN (influenza virus and poxvirus), and by elaborating superantigen-like proteins (mouse mammary tumor virus and related retroviruses and the rabies nucleocapsid). Superantigens activate large populations of T cells that express particular subsets of the T cell receptor β protein, causing massive cytokine release and subsequent host reactions. Another molecular mechanism of viral virulence involves the production of peptide growth factors for host cells, which disrupt normal cellular growth, proliferation, and differentiation. In addition, viral factors can bind to and interfere with the function of host receptors for signaling molecules. Modulation of cytokine production during viral infection can stimulate viral growth inside cells with receptors for the cytokine, and virus-encoded cytokine homologues (e.g., the Epstein-Barr virus BCRF1 protein, which is highly homologous to the immunoinhibitory IL-10 molecule) can potentially prevent immune-mediated clearance of viral particles. Viruses can cause disease in neural cells by interfering with levels of neurotransmitters without necessarily destroying the cells, or they may induce either programmed cell death (apoptosis) to destroy tissues or inhibitors of apoptosis to allow prolonged viral infection of cells. For infection to spread, many viruses must be released from cells. In a newly identified function, viral protein U (Vpu) of HIV facilitates the release of virus, a process that is specific to certain cells. Mammalian cells produce a restriction factor involved in inhibiting the release of virus; for HIV, this factor is designated BST-2 (bone marrow stromal antigen 2)/HM1.24/CD317, or tetherin. Vpu of HIV interacts with tetherin, promoting release of infectious virus. Overall, disruption of normal cellular and tissue function due to viral infection, replication, and release promotes clinical disease.

Bacterial Toxins Among the first infectious diseases to be understood were those due to toxin-elaborating bacteria. Diphtheria, botulism, and tetanus toxins are responsible for the diseases associated with local infections due to *Corynebacterium diphtheriae*, *Clostridium botulinum*, and *Clostridium tetani*, respectively. *Clostridium difficile* is an anaerobic gram-positive organism that elaborates two toxins, A and B, responsible for disruption of the intestinal mucosa when organism numbers expand in the intestine, leading to antibiotic-associated diarrhea and potentially to pseudomembranous colitis. Enterotoxins produced by *E. coli*, *Salmonella*, *Shigella*, *Staphylococcus*, and

V. cholerae contribute to diarrheal disease caused by these organisms. Staphylococci, streptococci, *P. aeruginosa*, and *Bordetella* elaborate various toxins that cause or contribute to disease, including toxic shock syndrome toxin 1; erythrogenic toxin; exotoxins A, S, T, and U; and pertussis toxin. A number of bacterial toxins (e.g., cholera toxin, diphtheria toxin, pertussis toxin, *E. coli* heat-labile toxin, and *P. aeruginosa* exotoxin) have adenosine diphosphate ribosyl transferase activity; i.e., the toxins enzymatically catalyze the transfer of the adenosine diphosphate ribosyl portion of nicotinamide adenine diphosphate to target proteins and inactivate them. The staphylococcal enterotoxins, toxic shock syndrome toxin 1, and the streptococcal pyrogenic exotoxins behave as superantigens, stimulating certain T cells to proliferate without processing of the protein toxin by antigen-presenting cells. Part of this process involves stimulation of the antigen-presenting cells to produce IL-1 and TNF- α , which have been implicated in many clinical features of diseases like toxic shock syndrome and scarlet fever. A number of gram-negative pathogens (*Salmonella*, *Yersinia*, and *P. aeruginosa*) can inject toxins directly into host target cells by means of a complex set of proteins referred to as the *type III secretion system*. Loss or inactivation of this virulence system usually greatly reduces the capacity of a bacterial pathogen to cause disease.

Endotoxin The lipid A portion of gram-negative LPS has potent biologic activities that cause many of the clinical manifestations of gram-negative bacterial sepsis, including fever, muscle proteolysis, uncontrolled intravascular coagulation, and shock. The effects of lipid A appear to be mediated by the production of potent cytokines due to LPS binding to CD14 and signal transduction via TLRs, particularly TLR4. Cytokines exhibit potent hypothermic activity through effects on the hypothalamus; they also increase vascular permeability, alter the activity of endothelial cells, and induce endothelial-cell procoagulant activity. Numerous therapeutic strategies aimed at neutralizing the effects of endotoxin are under investigation, but so far the results have been disappointing. It has been suggested that this lack of success may be due to substantial differences between mouse and human inflammatory responses to factors such as endotoxin; thus drugs developed in mouse models of infection may not be applicable to the human response.

Invasion Many diseases are caused primarily by pathogens growing in tissue sites that are normally sterile. Pneumococcal pneumonia is mostly attributable to the growth of *S. pneumoniae* in the lung and the attendant host inflammatory response, although specific factors that enhance this process (e.g., pneumolysin) may be responsible for some of the pathogenic potential of the pneumococcus. Disease that follows bloodstream infection and invasion of the meninges by meningitis-producing bacteria such as *N. meningitidis*, *H. influenzae*, *E. coli* K1, and group B streptococci appears to be due solely to the ability of these organisms to gain access to these tissues, multiply in them, and provoke cytokine production leading to tissue-damaging host inflammation.

Specific molecular mechanisms accounting for tissue invasion by fungal and protozoal pathogens are less well described. Except for studies pointing to factors like capsule and melanin production by *C. neoformans* and possibly levels of cell wall glucans in some pathogenic fungi, the molecular basis for fungal invasiveness is not well defined. Melanin has been shown to protect the fungal cell against death caused by phagocyte factors such as nitric oxide, superoxide, and hypochlorite. Morphogenic variation and production of proteases (e.g., the *Candida* aspartyl proteinase) have been implicated in fungal invasion of host tissues.

If pathogens are to effectively invade host tissues (particularly the blood), they must avoid the major host defenses represented by complement and phagocytic cells. Bacteria most often elude these defenses through their surface polysaccharides—either capsular polysaccharides or long O-side-chain antigens characteristic of the smooth LPS of gram-negative bacteria. These molecules can prevent the activation and/or deposition of complement opsonins or can limit the access of phagocytic cells with receptors for complement opsonins to these molecules when they are deposited on the bacterial surface

145e-10 below the capsular layer. Another potential mechanism of microbial virulence is the ability of some organisms to present the capsule as an apparent self antigen through molecular mimicry. For example, the polysialic acid capsule of group B *N. meningitidis* is chemically identical to an oligosaccharide found on human brain cells.

Immunochemical studies of capsular polysaccharides have led to an appreciation of the tremendous chemical diversity that can result from the linking of a few monosaccharides. For example, three hexoses can link up in more than 300 different, potentially serologically distinct ways, while three amino acids have only six possible peptide combinations. Capsular polysaccharides have been used as effective vaccines against meningococcal meningitis as well as against pneumococcal and *H. influenzae* infections and may prove to be of value as vaccines against any organisms that express a nontoxic, immunogenic capsular polysaccharide. In addition, most encapsulated pathogens become virtually avirulent when capsule production is interrupted by genetic manipulation; this observation emphasizes the importance of this structure in pathogenesis. It is noteworthy that the capsule-like surface polysaccharide PNAG has been found as a conserved structure shared by many microbes but generally is a poor target for antibody-mediated immunity because of the propensity of most humans and animals—all colonized by PNAG-producing microbes—to produce a nonprotective type of antibody. Altering the structure of PNAG by removing the acetate substituents on the N-acetylglucosamine monomers yields an immunogenic form, deacetylated PNAG, that reportedly induces antibodies that protect animals against diverse microbial pathogens.

Host Response The inflammatory response of the host is critical for interruption and resolution of the infectious process but is often responsible for the signs and symptoms of disease. Infection promotes a complex series of host responses involving the complement, kinin, and coagulation pathways. The production of cytokines such as IL-1, IL-18, TNF- α , IFN- γ , and other factors regulated in part by the NF- κ B transcription factor leads to fever, muscle proteolysis, and other effects. An inability to kill or contain the microbe usually results in further damage due to the progression of inflammation and infection. For example, in many chronic infections, degranulation of host inflammatory cells can lead to release of host proteases, elastases, histamines, and other toxic substances that can degrade host tissues. Chronic inflammation in any tissue can lead to the destruction of that tissue and to clinical disease associated with loss of organ function, such as sterility from pelvic inflammatory disease caused by chronic infection with *N. gonorrhoeae*.

The nature of the host response elicited by the pathogen often determines the pathology of a particular infection. Local inflammation produces local tissue damage, while systemic inflammation, such as that seen during sepsis, can result in the signs and symptoms of septic shock. The severity of septic shock is associated with the degree of production of host effectors. Disease due to intracellular parasitism results from the formation of granulomas, wherein the host attempts to wall off the parasite inside a fibrotic lesion surrounded by fused

epithelial cells that make up so-called multinucleated giant cells. A number of pathogens, particularly anaerobic bacteria, staphylococci, and streptococci, provoke the formation of an abscess, probably because of the presence of zwitterionic surface polysaccharides such as the capsular polysaccharide of *Bacteroides fragilis*. The outcome of an infection depends on the balance between an effective host response that eliminates a pathogen and an excessive inflammatory response that is associated with an inability to eliminate a pathogen and with the resultant tissue damage that leads to disease.

TRANSMISSION TO NEW HOSTS

As part of the pathogenic process, most microbes are shed from the host, often in a form infectious for susceptible individuals. However, the rate of transmissibility may not necessarily be high, even if the disease is severe in the infected individual, as these traits are not linked. Most pathogens exit via the same route by which they entered: respiratory pathogens by aerosols from sneezing or coughing or through salivary spread, gastrointestinal pathogens by fecal-oral spread, sexually transmitted diseases by venereal spread, and vector-borne organisms by either direct contact with the vector through a blood meal or indirect contact with organisms shed into environmental sources such as water. Microbial factors that specifically promote transmission are not well characterized. Respiratory shedding is facilitated by overproduction of mucous secretions, with consequently enhanced sneezing and coughing. Diarrheal toxins such as cholera toxin, *E. coli* heat-labile toxins, and *Shigella* toxins probably facilitate fecal-oral spread of microbial cells in the high volumes of diarrheal fluid produced during infection. The ability to produce phenotypic variants that resist hostile environmental factors (e.g., the highly resistant cysts of *E. histolytica* shed in feces) represents another mechanism of pathogenesis relevant to transmission. Blood parasites such as *Plasmodium* species change phenotype after ingestion by a mosquito—a prerequisite for the continued transmission of this pathogen. Venereally transmitted pathogens may undergo phenotypic variation due to the production of specific factors to facilitate transmission, but shedding of these pathogens into the environment does not result in the formation of infectious foci.

SUMMARY

In summary, the molecular mechanisms used by pathogens to colonize, invade, infect, and disrupt the host are numerous and diverse. Each phase of the infectious process involves a variety of microbial and host factors interacting in a manner that can result in disease. Recognition of the coordinated genetic regulation of virulence factor elaboration when organisms move from their natural environment into the mammalian host emphasizes the complex nature of the host-parasite interaction. Fortunately, the need for diverse factors in successful infection and disease implies that a variety of therapeutic strategies may be developed to interrupt this process and thereby to prevent and treat microbial infections.

146 Genomics and Infectious Disease

Roby P. Bhattacharyya, Yonatan H. Grad,
Deborah T. Hung

Just as microscopy opened up the worlds of microbiology by providing a tool with which to visualize microorganisms, technological advances in genomics are now providing microbiologists with powerful new methods with which to characterize the genetic map underlying all

microbes with unprecedented resolution, thereby illuminating their complex and dynamic interactions with one another, the environment, and human health. The field of infectious disease genomics encompasses a vast frontier of active research that has the potential to transform clinical practice in relation to infectious diseases. While genetics has long played a key role in elucidating the process of infection and managing clinical infectious diseases, the ability to extend our thinking and our approaches beyond the study of single genes to an examination of the sequence, structure, and function of entire genomes is identifying new possibilities for research and opportunities to change clinical practice. From the development of diagnostics with unprecedented sensitivity, specificity, and speed to the design of novel public health interventions, technical and statistical genomic innovations are reshaping our understanding of the influence of the microbial world on human health and providing us with new tools to combat infection. This chapter explores the application of genomics methods to microbial pathogens and the infections they cause (Table 146-1). It discusses innovations that are driving the development of diagnostic approaches and the discovery of new pathogens; providing insight into novel therapeutic approaches and paradigms; and advancing methods in infectious disease epidemiology and the study of pathogen evolution that can inform infection control measures, public health responses to outbreaks, and vaccine development. We draw on examples in current practice and from the recent scientific literature as signposts that point toward the ways in which the insights from pathogen genomics may influence infectious diseases in the short and long terms. Table 146-2 provides definitions for a selection of important terms used in genomics.

MICROBIAL DIAGNOSTICS

The basic goals of a clinical microbiology laboratory are to establish the presence of a pathogen in a clinical sample, to identify the pathogen, and, when possible, to provide other information that can help guide clinical management and even prognosis, such as antibiotic susceptibility profiles or the presence of virulence factors. To date, clinical microbiology laboratories have largely approached these goals phenotypically by growth-based assays and biochemical testing. Bacteria, for instance, are algorithmically grouped into species by their characteristic microscopic appearance, nutrient requirements for growth, and ability to catalyze certain reactions. Antibiotic susceptibility is determined in most cases by assessing growth in the presence of antibiotic.

With the sequencing revolution paving the way to easy access of complete pathogen genomes (Fig. 146-1), we are now able to more systematically clarify the genetic basis of these observable phenotypes. Compared with traditional growth-based methods for bacterial diagnostics that dominate the clinical microbiology laboratory, nucleic

TABLE 146-1 CURRENT CLINICAL APPLICATIONS OF INFECTIOUS DISEASE GENOMICS

Application	Technology	Notes/Examples
Organism Identification		
Viral detection	PCR	Identification of HIV, HBV, HCV, respiratory viruses including influenza, and others for diagnosis and response to therapy
TB detection	PCR	Amplification of the <i>rhoB</i> gene for species-specific identification of <i>Mycobacterium tuberculosis</i>
Bacterial detection	PCR, NAAT	Identification of <i>Chlamydia</i> , <i>Neisseria gonorrhoeae</i> , <i>Clostridium difficile</i> , <i>Ehrlichia</i> , <i>Anaplasma</i> , and others
Bacterial detection	16S ribosomal gene PCR	Targeting of highly conserved regions of the 16S rRNA gene for identification of suspected bacterial infections undiagnosed by conventional methods
Pathogen Discovery		
Bacterial pathogens	Sequencing, metagenomic assembly	Unbiased "shotgun" sequencing of isolated nucleic acid from patient samples to identify associated pathogens; proofs-of-concept: new <i>Bradyrhizobium</i> species associated with cord colitis, <i>Escherichia coli</i> O104:H4 from 2011 diarrheal outbreak in Germany; research use only at this time
Viral pathogens	Microarray, sequencing	Hybridization of clinical samples to microarrays from phylogenetically diverse known viruses identified the SARS coronavirus and others. Direct sequencing has identified West Nile virus and the MERS coronavirus, among others. Use is primarily in research.
Antibiotic Resistance		
MRSA detection	PCR	Detection of the <i>mecA</i> gene, the genotypic cause of methicillin resistance in <i>Staphylococcus aureus</i>
VRE detection	PCR	Detection of the <i>vanA</i> or <i>vanB</i> genes, the main genotypic causes of vancomycin resistance in <i>Enterococcus</i>
MDR-TB detection	PCR, NAAT	Detection of polymorphisms in the <i>rhoB</i> gene from <i>M. tuberculosis</i> , which account for 95% of rifampin resistance. Other probes available for <i>inhA</i> and <i>katG</i> genes can detect up to 85% of isoniazid resistance.
Carbapenemase detection	PCR	Detection of genes encoding one of two enzymes, NDM-1 or KPC, that hydrolyze carbapenems; use in United States currently restricted to CDC
HIV resistance detection	Targeted sequencing	Targeted sequencing of specific genes with known resistance-conferring mutations; now standard of care prior to initial therapy in United States and Europe
Epidemiology		
Outbreak and epidemic tracking	Sequencing	Application to tracking outbreaks and epidemics on local and international scales, including spread of carbapenemase-producing <i>Klebsiella</i> , <i>S. aureus</i> , <i>M. tuberculosis</i> , <i>E. coli</i> , <i>Vibrio cholerae</i> , and influenza virus
Evolution and spread of pathogens	Sequencing	Sequencing collections of pathogens to shed light on pathogen dissemination, virulence factors, and antibiotic resistance determinants

Abbreviations: CDC, Centers for Disease Control and Prevention; HBV, hepatitis B virus; HCV, hepatitis C virus; MDR, multidrug-resistant; MERS, Middle East respiratory syndrome; MRSA, methicillin-resistant *Staphylococcus aureus*; NAAT, nucleic acid amplification test; PCR, polymerase chain reaction; SARS, severe acute respiratory syndrome; TB, tuberculosis; VRE, vancomycin-resistant enterococci.

TABLE 146-2 GLOSSARY OF SELECTED TERMS IN GENOMICS

Term	Definition
Contig	A DNA sequence representing a continuous fragment of a genome, assembled from overlapping sequences; relevant for de novo assembly of sequence data that do not align to previously sequenced genomes
Genome	The entire set of heritable genetic material within an organism
Horizontal gene transfer	The transfer of genes between organisms through mechanisms other than by clonal descent, such as through transformation, conjugation, or transduction
Metagenomics	Analysis of genetic material from multiple species directly from primary samples without requiring prior culture steps
Microarray	A collection of DNA oligonucleotides ("oligos") spatially arranged on a solid surface and used to detect or quantify sequences in a sample of interest that are complementary (and therefore bind) to one or more of the arrayed oligos
Mobile genetic element	DNA elements that can move within a genome and can be transferred between genomes through horizontal gene transfer (e.g., plasmids, bacteriophages, and transposons)
Multilocus sequence typing	A methodology for typing organisms based on DNA sequence fragments from a prespecified set of genes
Next-generation sequencing	High-throughput sequencing using a parallelized sequencing process that produces millions of sequences concurrently, far beyond the capacity of prior dye-terminator methods
Nucleic acid amplification test (NAAT)	Biochemical assay that evaluates for the presence of a particular string of nucleic acids through amplification by one of several methods, including polymerase and ligase chain reactions
Polymerase chain reaction (PCR)	A subset of NAAT used to amplify a specific region of DNA with specific oligonucleotide primers and a DNA polymerase
Transcriptome	The catalog of the full set of messenger RNA (mRNA) transcripts from a cell or organism, which are typically measured by microarray or by next-generation sequencing of complementary DNA (cDNA) via a process called RNA-Seq
Whole-genome sequencing	A process that determines the full DNA sequence of an organism's genome; has been greatly facilitated by next-generation sequencing technology

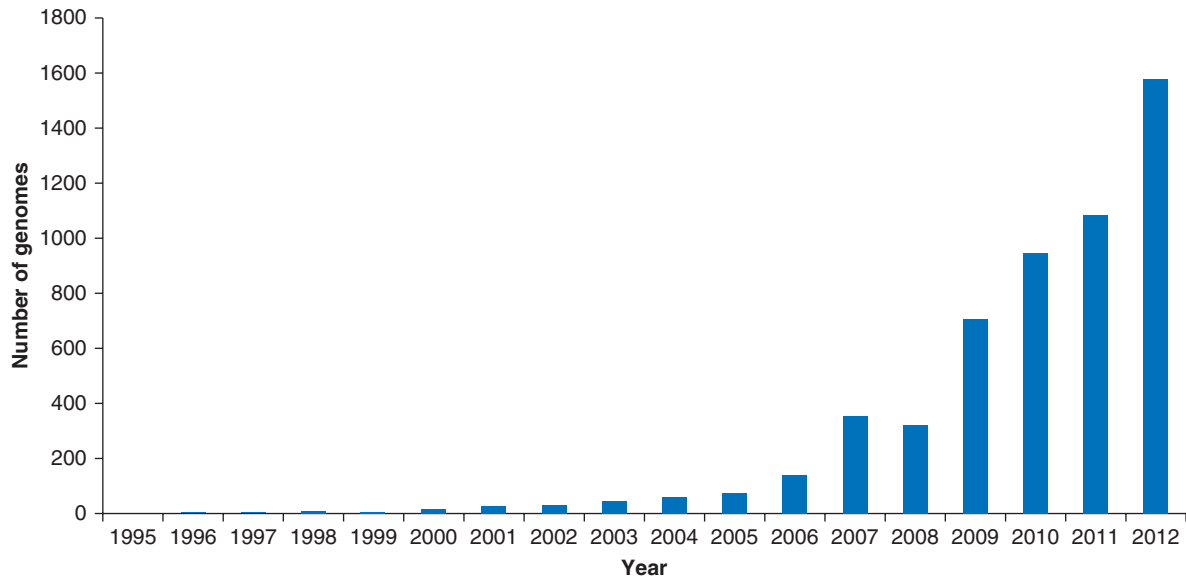


FIGURE 146-1 Completed bacterial genome sequences by year, through 2012. (Data compiled from www.genomesonline.org.)

acid-based diagnostics promise improved speed, sensitivity, specificity, and breadth of information. Bridging clinical and research laboratories, adaptations of genomic technologies have begun to deliver on this promise.

HISTORICAL LIMITATIONS AND PROGRESS THROUGH GENETIC APPROACHES

The molecular diagnostics revolution in the clinical microbiology laboratory is well under way, borne of necessity in the effort to identify microbes that are refractory to traditional culture methods. Historically, diagnosis of many so-called unculturable pathogens has relied largely on serology and antigen detection. However, these methods provide only limited clinical information because of their suboptimal sensitivity and specificity as well as the long delays that diminish their utility for real-time patient management. Newer tests to detect pathogens based on nucleic acid content have already offered improvements in the select cases to which they have been applied thus far.

Unlike direct pathogen detection, serologic diagnosis—measurement of the host's response to pathogen exposure—can typically be made only in retrospect, requiring both acute- and convalescent-phase sera. For chronic infections, distinguishing active from latent infection or identifying repeat exposure by serology alone can be difficult or impossible, depending on the syndrome. In addition, the sensitivity of serologic diagnosis varies with the organism and the patient's immune status. For instance, tuberculosis is notoriously difficult to identify by serologic methods; tuberculin skin testing using purified protein derivative (PPD) is especially insensitive in active disease and may be cross-reactive with vaccines or other mycobacteria. Even the newer interferon γ release assays (IGRAs), which measure cytokine release from T lymphocytes in response to *Mycobacterium tuberculosis*-specific antigens in vitro, have limited sensitivity in immunodeficient hosts. Neither PPD testing nor IGRAs can distinguish latent from active infection. Serologic Lyme disease diagnostics suffer similar limitations: in patients from endemic regions, the presence of IgG antibodies to *Borrelia burgdorferi* may reflect prior exposure rather than active disease, while IgM antibodies are imperfectly sensitive and specific (50% and 80%, respectively, in early disease). The complex nature of these tests, particularly in view of the nonspecific symptoms that may accompany Lyme disease, has had substantial implications on public perceptions of Lyme disease and antibiotic misuse in endemic areas. Similarly, syphilis, a chronic infection caused by *Treponema pallidum*, is notoriously difficult to stage by serology alone, requiring the use of multiple different nontreponemal (e.g., rapid protein reagin) and treponemal (e.g., fluorescent treponemal antibody) tests in conjunction with clinical suspicion. Complementing serology, antigen detection can

improve sensitivity and specificity in select cases but has been validated only for a limited set of infections. Typically, structural elements of pathogens are detected, including components of viral envelopes (e.g., hepatitis B surface antigen, HIV p24 antigen), cell surface markers in certain bacteria (e.g., *Streptococcus pneumoniae*, *Legionella pneumophila* serotype 1) or fungi (e.g., *Cryptococcus*, *Histoplasma*), and less specific fungal cell-wall components such as galactomannan and β -glucan (e.g., *Aspergillus* and other dimorphic fungi).

Given the impracticality of culture and the lack of sensitivity or sufficient clinical information afforded by serologic and antigenic methods, the push toward nucleic acid-based diagnostics originated in pursuit of viruses and fastidious bacteria, becoming part of the standard of care for select organisms in U.S. hospitals. Such tests, including polymerase chain reaction (PCR) and other nucleic acid amplification tests (NAATs), are now widely used for many viral infections, both chronic (e.g., HIV infection) and acute (e.g., influenza). This technique provides essential information about both the initial diagnosis and the response to therapy and in some cases genotypically predicts drug resistance. Indeed, progression from antigen detection to PCR transformed our understanding of the natural course of HIV infection, with profound implications for treatment (Fig. 146-2). In the early years of the AIDS pandemic, p24 antigenemia was detected in acute HIV infection but then disappeared for years before emerging again with progression to AIDS (Fig. 146-2B). Without a marker demonstrating viremia, the role of treatment during HIV infection prior to the development of clinical AIDS was uncertain, and monitoring treatment efficacy was challenging. With the emergence of PCR as a progressively more sensitive test (now able to detect as few as 20 copies of virus per milliliter of blood), viremia was recognized as a near-universal feature of HIV infection. This recognition has been transformative in guiding the initiation of therapy as well as adjustments in therapy and, together with the development of less toxic therapies, has helped to shape guidelines that now favor earlier introduction of antiretroviral therapy for HIV infection.

As they are for viruses, nucleic acid-based tests have become the diagnostic tests of choice for fastidious bacteria, including the common sexually transmitted intracellular bacterial pathogens *Neisseria gonorrhoeae* and *Chlamydia trachomatis* as well as the tick-borne *Ehrlichia chaffeensis* and *Anaplasma phagocytophilum*. More recently, nucleic acid amplification-based detection has offered improved sensitivity for diagnosis of the important nosocomial pathogen *Clostridium difficile*; NAATs can provide clinically relevant information on the presence of cytotoxins A and B as well as molecular markers of hypervirulence such as those characterizing the recently recognized North American pulsotype 1 (NAP1), which is found more frequently in

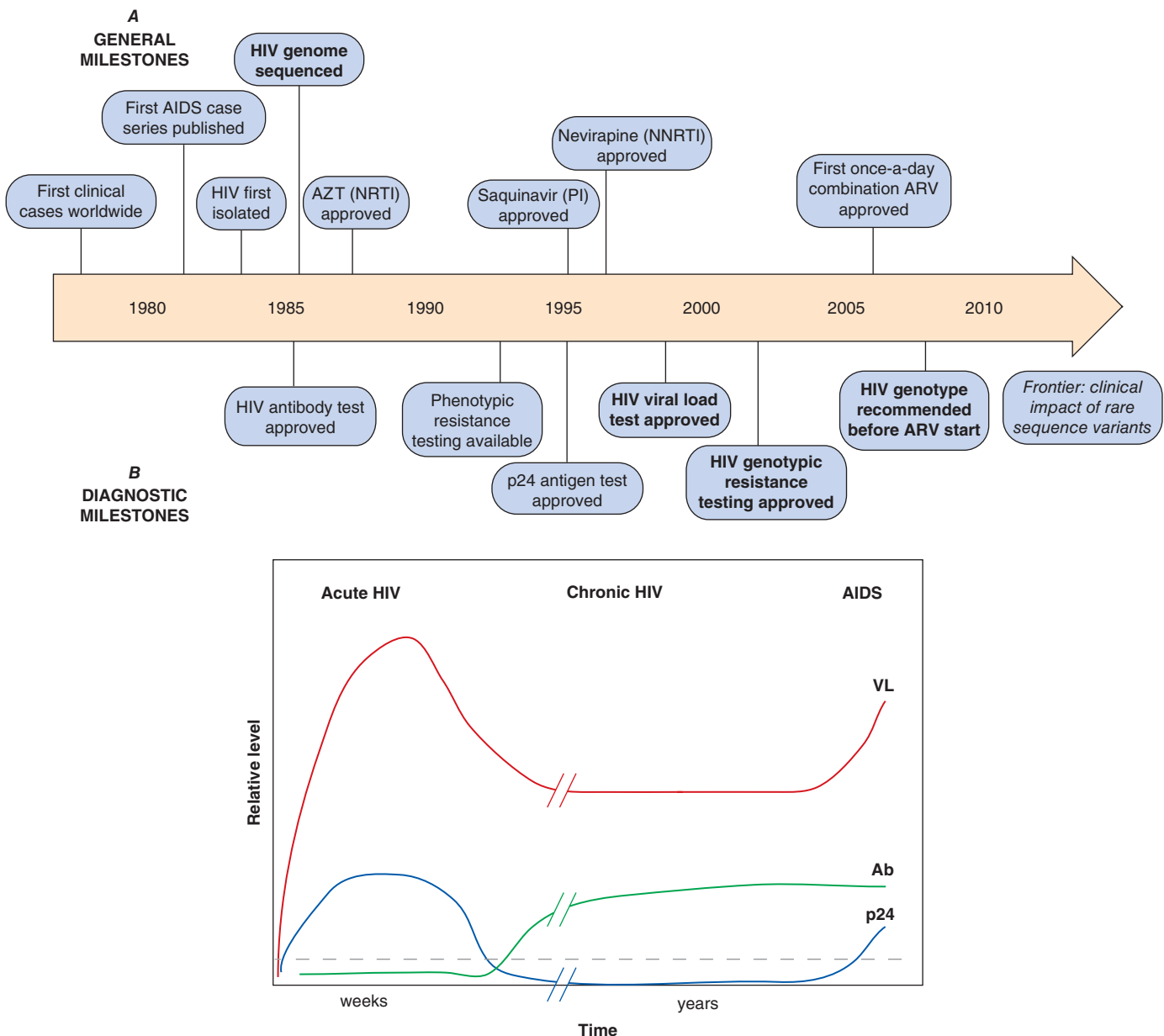


FIGURE 146-2 **A.** Timeline of select milestones in HIV management. Genomic advances are shown in bold type. The approvals and recommendations indicated apply to the United States. ARV, antiretroviral; AZT, zidovudine; NRTI, nucleoside reverse transcriptase (RT) inhibitor; NNRTI, non-nucleoside RT inhibitor; PI, protease inhibitor. **B.** Viral dynamics in the natural history of HIV infection. Three diagnostic markers are shown: HIV antibody (Ab), p24 antigen (p24), and viral load (VL). Dashed gray line represents limit of detection. (Adapted from data in HH Fiebig et al: *Dynamics of HIV viremia and antibody seroconversion in plasma donors: Implications for diagnosis and staging of primary HIV infection. AIDS 17:1871, 2003.*)

cases of severe illness. The importance of genomics in selecting loci for diagnostic assays and in monitoring test sensitivity was recently highlighted by the emergence in Sweden of a new variant of *C. trachomatis* containing a deletion that includes the gene targeted by a set of commercial NAATs. By evading detection through this deletion (which would have prompted the initiation of treatment), this strain came to be highly prevalent in some areas of Sweden. While nucleic acid–based tests remain the diagnostic approach of choice for fastidious bacteria, this example serves as a reminder of the need for careful development and ongoing monitoring of molecular diagnostics.

In contrast, for typical bacterial pathogens for which culture methods are well established, growth-based assays followed by biochemical tests still dominate in the clinical laboratory. Informed by decades of clinical microbiology, these tests have served clinicians well, yet the limitations of growth-based tests—in particular, the delays associated with waiting for growth—have left open opportunities for improvements. Molecular diagnostics, greatly informed by the vast quantity

of microbial genome sequences generated in recent years, offers a way forward. First, sequencing studies may identify key genes (or noncoding nucleic acids) that can be developed into targets for clinical assays using PCR or hybridization platforms. Second, sequencing itself may eventually become inexpensive and rapid enough to be performed routinely on clinical specimens, with consequent unbiased detection of pathogens.

ORGANISM IDENTIFICATION

In order to adapt nucleic acid detection to diagnostic tests and thus to identify pathogens on a wide scale, sequences must be identified that are conserved enough within a species to identify the diversity of strains that may be encountered in various clinical settings, yet divergent enough to distinguish one species from another. Until recently, this problem has been solved for bacteria by targeting the element of a bacterial genome that is most highly conserved within a species: the 16S ribosomal RNA (rRNA) subunit. At present, 16S PCR amplification from

772 tissue specimens can be performed by specialty laboratories, though its sensitivity and clinical utility to date have remained somewhat limited because, for instance, of inhibitory molecules often found in clinical tissue samples that prevent reliable, sensitive PCR amplification. As such barriers are reduced through technological advances and as the causes of culture-negative infection are clarified (perhaps in part through sequencing efforts), these tests may become both more accessible and more helpful.

With the wealth of sequencing data now available, other regions beyond 16S rRNA can be targeted for bacterial species identification. These other genomic loci can provide additional information about a clinical isolate that is relevant to patient management. For instance, detection of the presence—or potentially even the expression—of toxin genes such as those for *C. difficile* toxins A and B or Shiga toxin may provide clinicians with additional information that will help distinguish commensals or colonizing bacteria from pathogens and thus aid in prognostication as well as diagnosis.

While amplification tests such as PCR exemplify one approach to nucleic acid detection, other approaches exist, including detection by hybridization. Although not currently used in the clinical realm, techniques for detection and identification of pathogens by hybridization to microarrays are being developed for other purposes. Of note, these different detection techniques require different degrees of conservation. Highly sensitive amplification methods require a high degree of sequence identity between PCR primer pairs and their short, specific target sequences; even a single base-pair mismatch (particularly near the 3' end of the primer) may interfere with detection. In contrast, hybridization-based tests are more tolerant of mismatch and thus can be used to detect important regions that may be less precisely conserved within a species, thus potentially allowing detection of clinical isolates from a given species with greater diversity between isolates. Such assays take advantage of the predictable binding interactions of nucleic acids. The applicability of hybridization-based methods toward either DNA or RNA opens up the possibility of expression profiling, which can uncover phenotypic information from nucleic acid content.

Both PCR and hybridization methods target specific, known organisms. At the other extreme, as sequencing costs and turnaround times decrease, direct metagenomic sequencing from patient samples is becoming increasingly feasible. This shotgun sequencing approach is unbiased—i.e., is able to detect any microbial sequence, however divergent or unexpected. This new approach brings its own set of challenges, however, including the need to recognize pathogenic sequences against a background of expected host and commensal sequences and to distinguish true pathogens from either colonizers or laboratory contaminants. In a powerful example of this new frontier of sequencing-based clinical diagnosis, investigators diagnosed neuroleptospirosis in a child with an unexplained encephalitis syndrome by finding sequences corresponding to the *Leptospira* genus in cerebrospinal fluid from the patient. Rapid (<48-h) sequencing and analysis informed the patient's care in real time, leading to life-saving targeted antibiotic therapy for an unexpected diagnosis that was impossible to make through standard laboratory testing. The diagnosis was retrospectively confirmed through both convalescent serologies and PCR using primers designed on the basis of sequencing data.

PATHOGEN DISCOVERY

In addition to clinical diagnostic applications, novel genomic technologies, including whole-genome sequencing, are being applied to clinical research specimens with a goal of identifying new pathogens in a variety of circumstances. The tremendous sensitivity and unbiased nature of sequencing is also ideal in searching clinical samples for unknown or unsuspected pathogens.

Causal inference in infectious diseases has progressed since the time of Koch, whose historical postulates provided a rigorous framework for attributing a disease to a microorganism. According to an updated version of Koch's postulates, an organism, whether it can be cultured or not, should induce disease upon introduction into a healthy host if it is to be implicated as a causative pathogen. Current sequencing

technologies are ideal for advancing this modern version of Koch's postulates because they can identify candidate causal pathogens with unprecedented sensitivity and in an unbiased way, unencumbered by limitations such as culturability. Yet, as direct sequencing on primary patient samples greatly expands our ability to recognize associations between microbes and disease states, critical thinking and experimentation will continue to be vital to establishing causality.

Virus discovery in particular has been greatly facilitated by new nucleic acid technology. These frontiers were first notably explored with high-density microarrays containing spatially arrayed sequences from a phylogenetically diverse collection of viruses. Although biased toward those with homology to known viruses, novel viruses in clinical samples were successfully identified on the basis of their ability to hybridize to these prespecified sequences. This methodology famously contributed to identification of the coronavirus causing severe acute respiratory syndrome (SARS). Once discovered, this SARS coronavirus was rapidly sequenced: the full genome was assembled in April 2003, less than 6 months after recognition of the first case. This accomplishment illustrated the advancing power and speed of new diagnostic technologies.

With the advent of next-generation sequencing, unbiased pathogen discovery is now possible through a process known as *metagenomic assembly* (Fig. 146-3). Sequences of random nucleotide fragments can be generated from clinical specimens with no a priori knowledge of pathogen identity through a process called *shotgun sequencing*. This collection of sequences can then be computationally aligned to host (i.e., human) sequences, with aligned sequences removed and remaining sequences compared with other known genomes to detect the presence of known microorganisms. Sequence fragments that remain unaligned suggest the presence of an additional organism that cannot be matched to a known, characterized genome; these reads can be assembled into contiguous nucleic acid stretches that can be compared to known sequences to construct the genome of a potentially novel organism. Assembled genomes (or parts of genomes) can then be compared to known genomes to infer the phylogeny of new organisms and identify related classes or traits. Thus, not only can this process identify unanticipated pathogens; it can even identify undiscovered organisms. Some early applications of sequencing on clinical samples have centered around the discovery of novel viruses, including such emerging pathogens as West Nile virus, SARS coronavirus, and the Middle East respiratory syndrome coronavirus (MERS-CoV) that has caused severe respiratory illnesses in healthy adults, as well as viral causes of myriad other conditions, from tropical hemorrhagic fevers to diarrhea in newborns.

More recently, metagenomic assembly has been successfully extended to bacterial pathogen discovery. Investigators identified a new bacterial species associated with “cord colitis”—a rare antibiotic-responsive, culture-negative colitis in recipients of umbilical cord-blood stem cells—by sequencing colon biopsy samples from affected patients and matched controls. A single dominant species emerged from metagenomic assembly in samples from patients that was absent from control samples. The presence of this species was confirmed by PCR and fluorescence in situ hybridization on primary tissue samples. On the basis of its similarity to other known species, the organism was named *Bradyrhizobium enterica*, a novel species from a genus that has proved difficult to culture and thus would have been hard to identify by other means. Correlation versus causation remains an open question; therefore, further efforts will be required to make such links.

As metagenomic sequencing and assembly techniques become more robust, this technology holds great promise for identifying microorganisms that are associated with clinical conditions of unknown etiology. Conventional methods already have unexpectedly linked numerous conditions with specific agents of infection—e.g., cervical and oropharyngeal cancers with human papillomavirus, Kaposi's sarcoma with human herpesvirus 8, and certain lymphomas with Epstein-Barr virus. Sequencing techniques offer unprecedented sensitivity and specificity for identifying foreign nucleic acid sequences that may suggest other conditions—from malignancies to inflammatory conditions to unexplained fevers or other clinical syndromes—associated with organisms

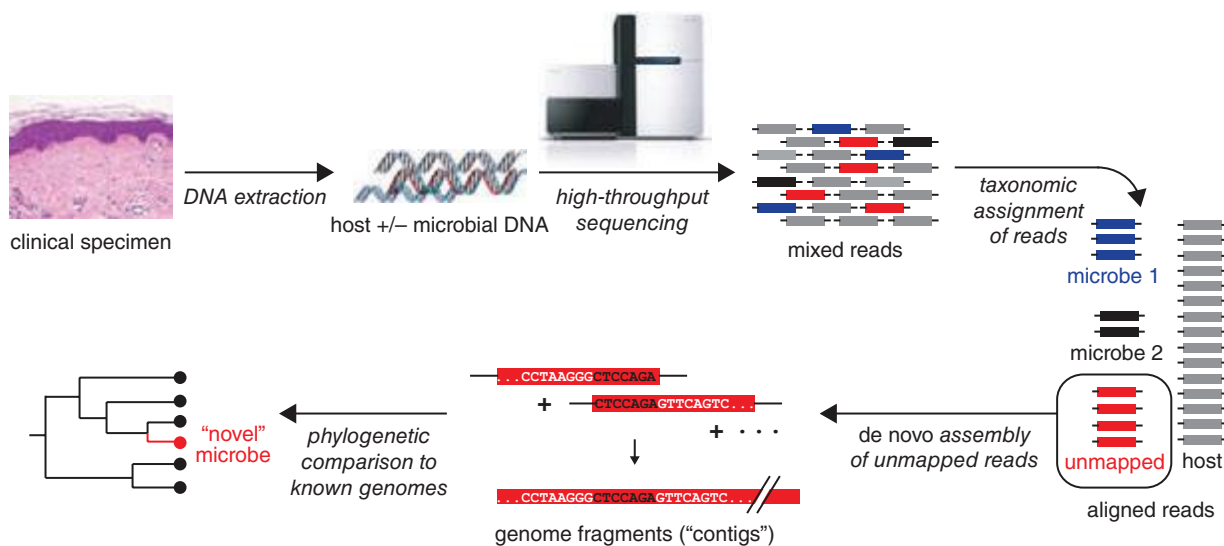


FIGURE 146-3 Workflow of metagenomic assembly for pathogen discovery. DNA is isolated from a specimen of interest (e.g., tissue, body fluid) containing a mixture of host DNA and nucleic acids from coexisting microbes, either commensal or pathogenic. All DNA (and RNA if a reverse transcription step is added) is then sequenced, yielding a mixture of DNA sequence fragments (“reads”) from organisms present. These reads are then aligned to existing reference genomes for the host or any known microbes, leaving reads that do not align (“map”) to any known sequence. These unmapped reads are then computationally assembled de novo into the largest contiguous stretches of DNA possible (“contigs”), representing fragments of previously unsequenced genomes. These genome fragments (contigs) are then mapped onto a phylogenetic tree based on their sequence. Some may represent known but as-yet-unsequenced organisms, while others will represent novel species. (Figure prepared with valuable input from Dr. Ami S. Bhatt, personal communication.)

from viruses to bacteria to parasites. As sequencing-based discovery expands, microbes may be found to be associated with conditions not classically thought of as infectious. Studies of bowel flora in laboratory animals and even humans are already beginning to suggest correlations between microbe composition and various aspects of metabolic and cardiovascular health. Improved methods for pathogen detection will continue to uncover unexpected correlations between microbes and disease states, but the mere presence of a microbe does not establish causality. Fortunately, once the relatively laborious and computationally intensive metagenomic sequencing and assembly efforts have identified a pathogen, further detection can easily be undertaken with targeted methods such as PCR or hybridization, which are much more straightforward and scalable. This capacity should facilitate the additional careful investigation that will be required to progress beyond correlation and to draw causal inference.

ANTIBIOTIC RESISTANCE

At present, antibiotic resistance in bacteria and fungi is determined by isolating a single colony from a cultured clinical specimen and testing its growth in the presence of a drug. The requirement for multiple growth steps in these conventional assays has several consequences. First, only culturable pathogens can be readily processed. Second, this process requires considerable infrastructure to support the sterile environment required for culture-based testing of diverse organisms. Finally, and perhaps most significantly, even the fastest-growing organisms require 1–2 days of processing for identification and 2–3 days for determination of susceptibilities. Slower-growing organisms take even longer: for instance, weeks must pass before drug-resistant *M. tuberculosis* can be identified by growth phenotype. Given the clinical imperative in serious illness to begin effective therapy early, this inherent delay in susceptibility determination has obvious implications for empirical antibiotic use: broad-spectrum antibiotics often must be chosen up front in situations where it is later shown that preferred narrower-spectrum drugs would have been effective or even that no antibiotics were appropriate (i.e., in viral infections). With this strategy, the empirical choice can be incorrect, often with devastating consequences. Real-time identification of the infecting organism and information on its susceptibility profile would guide initial therapy and support judicious antibiotic use, ideally improving patient outcomes while aiding in the ever-escalating struggle with antibiotic

resistance by reserving the use of broad-spectrum agents for cases in which they are truly needed.

Molecular diagnostics and sequencing offer a way to accelerate detection of a pathogen’s antibiotic susceptibility profile. If a genotype that confers resistance can be identified, this genotype can be targeted for molecular detection. In infectious disease, this approach has most convincingly come to fruition for HIV (Fig. 146-2A). (In a conceptually parallel application of genomic analysis, molecular detection of certain resistance determinants in cancers is beginning to inform chemotherapeutic selection.) Extensive sequencing of HIV strains and correlations drawn between viral genotypes and phenotypic resistance have delineated the majority of mutations in key HIV genes, such as reverse transcriptase, protease, and integrase, that confer resistance to the antiretroviral agents that target these proteins. For instance, the single-amino-acid substitution K103N in the HIV reverse transcriptase gene predicts resistance to the first-line nonnucleoside reverse transcriptase inhibitor efavirenz, and its detection thus informs a clinician to choose a different agent. The effects of these common mutations on HIV susceptibility to various drugs—as well as on viral fitness—are curated in publicly available databases. Thus, genotypes are now routinely used to predict drug resistance in HIV, as phenotypic resistance assays are far more cumbersome than targeted sequencing. Indeed, current recommendations in the United States are to sequence virus from a patient’s blood before initiating antiretroviral therapy, which is then tailored to the predicted resistance phenotype. As new targeted therapies are introduced, this targeted sequencing-based approach to drug resistance will likely prove important in other viral infections (e.g., hepatitis C).

For several reasons, the challenge of predicting antibiotic susceptibility from genotype has not yet been met in bacteria to the same degree as in HIV. In general, bacteria have evolved diverse resistance mechanisms to most antibiotics; thus, the task cannot be reduced to probing for a single genetic lesion, target, or mechanism. For instance, at least five distinct modes of resistance to fluoroquinolones are known: reduced import, increased efflux, mutated target sites, drug modification, and shielding of the target sites by expression of another protein. Further, we lack a comprehensive compendium of genetic elements conferring resistance, and new mechanisms and genes emerge regularly in the face of antibiotic deployment. As bacteria have far more complex genomes than viruses, with thousands of genes on their chromosomes and the capacity for acquiring many more through

774 horizontal gene transfer of plasmids and mobile genetic elements within and even between species, the task of not only defining all current but also predicting all future mechanisms at a genetic level is daunting or perhaps impossible.

Despite these challenges, in a few select cases where the genotypic basis for resistance has been well defined, genotypic assays for antibiotic resistance are already being introduced into clinical practice. One important example is the detection of methicillin-resistant *Staphylococcus aureus* (MRSA). *S. aureus* is one of the most common and serious bacterial pathogens of humans, particularly in health care settings. Resistance to methicillin, the most effective antistaphylococcal antibiotic, has become very common even in community-acquired strains. The alternative to methicillin, vancomycin, is effective against MRSA but measurably inferior to methicillin against methicillin-susceptible *S. aureus* (MSSA). Analysis of clinical MRSA isolates has demonstrated that the molecular basis for resistance to methicillin in essentially all cases stems from the expression of an alternative penicillin-binding protein (PBP2A) encoded by the gene *mecA*, which is found within a transferable genetic element called *mec*. This mobile cassette has spread rapidly through the *S. aureus* population via horizontal gene transfer and selection from widespread antibiotic use. Because resistance is essentially always due to the presence of the *mec* cassette, MRSA is amenable to molecular detection. In recent years, a PCR test for the presence of the *mec* cassette, which saves hours to days compared with standard culture-based methods, has been approved by the U.S. Food and Drug Administration.

Additional molecular diagnostics are being implemented in the evaluation of bacterial antibiotic resistance. Vancomycin-resistant enterococci (VRE) harbor one of a limited number of *van* genes responsible for resistance to this important antibiotic by altering the mechanism for cell wall cross-linking that vancomycin inhibits. Detection of one of these genes by PCR indicates resistance. Identification of two carbapenemase-encoding plasmids—NDM-1 and KPC—responsible for a significant fraction of carbapenem resistance (though not for all such resistance) has led to the development of a multiplexed PCR assay to detect these important resistance elements. Because carbapenems are broad-spectrum antibiotics frequently reserved for multidrug-resistant bacteria (particularly enteric gram-negative bacilli) and are often used as antibiotics of last resort, the initial appearance of resistance and the subsequent increase in its prevalence have caused considerable concern. Therefore, even though other mechanisms for carbapenem resistance exist, a rapid PCR test for the two plasmids encoding these two carbapenemases has been developed to aid in both diagnostic and infection control efforts. Efforts are under way to extend this multiplexed PCR assay to other plasmid-borne carbapenemases and thus to make it more comprehensive.

The power and application of molecular genetic tests are not limited to high-income settings. With the increasing burden of drug-resistant tuberculosis in the developing world, a molecular diagnostic test has now been developed to detect rifampin-resistant tuberculosis. The genetic basis for rifampin resistance has been well defined by targeted sequencing: characteristic mutations in the molecular target of rifampin, RNA polymerase, account for the vast majority of rifampin-resistant strains of *M. tuberculosis*. A PCR assay that can detect both the *M. tuberculosis* organism and a rifampin-resistant allele of RNA polymerase from clinical samples has recently been approved. Since rifampin resistance frequently accompanies resistance to other antibiotics, this test can suggest the possible presence of multidrug-resistant *M. tuberculosis* within hours instead of weeks.

Despite differences in relative genome complexity, HIV genotypic screening for antiretroviral resistance offers one framework for broadening efforts at genotypic assays for nonviral antibiotic resistance. As whole-genome pathogen sequencing has become increasingly feasible and inexpensive (Fig. 146-1), significant efforts have been launched to sequence hundreds to thousands of antibiotic-sensitive and -resistant isolates of a given pathogen in order to more comprehensively define resistance-conferring genetic elements. In parallel with advancing sequencing technologies, progress in computational techniques, bioinformatics and statistics, and data storage as well as experimental

confirmatory testing of hypotheses will be needed to move toward the ambitious goal of a comprehensive compendium of antibiotic resistance determinants. Open sharing and careful curation of new sequence information will be of paramount importance.

Yet no matter how thorough and carefully curated such a genotype-phenotype database is, history suggests that comprehensively cataloguing resistance in nonviral pathogens, with new mechanisms continuously emerging, will be challenging at best. Even identifying and itemizing current resistance mutations is a daunting prospect: nonviral genomes are much larger than viral ones, and their abundance and diversity are such that hundreds to thousands of genetic differences often exist between clinical isolates, of which perhaps only one may cause resistance. For example, increasing resistance to artemisinin, one of the most effective new agents for malaria, has prompted recent large-scale efforts to identify the basis for resistance. While such studies have identified promising leads, no clear mechanism has emerged; in fact, a single genetic lesion alone may not fully account for resistance. Especially with multiple possible resistance mechanisms for a given antibiotic as well as ongoing evolutionary pressure resulting in the development and acquisition of new modes of resistance, a genotypic approach to diagnosing antibiotic resistance is likely to be imperfect.

We have already observed the accumulation of new or unanticipated modes of resistance from ongoing evolutionary pressure caused by the widespread clinical use of antibiotics. Even with MRSA, perhaps the best-studied case of antibiotic resistance and a model of relative simplicity with a single known monogenic resistance determinant (*mecA*), a genotype-based approach to resistance detection proved flawed. One limitation was a recall of the initial commercial genotypic resistance assay that was deployed for the identification of MRSA. A clinical isolate of *S. aureus* emerged in Belgium that expressed a variant of the *mec* cassette not detected by the assay's PCR primers. New primers were added to detect this new variant, and the assay was re-approved for use. More recently, an even more divergent but functionally analogous gene called *mecC*, which confers methicillin resistance but evades PCR detection by this assay, was found. This series of events exemplifies the need for ongoing monitoring of any genotypic resistance assay. A second limitation is that a contradiction can occur between genotypic and phenotypic evidence for resistance. Up to 5% of MSSA strains carry a copy of the *mecA* gene that is either nonfunctional or not expressed. Thus, the erroneous identification of these strains as MRSA by genotypic detection would lead to administration of the inferior antibiotic vancomycin rather than the preferred β -lactam therapy.

These examples illustrate one of the prime challenges of moving beyond growth-based assays: genotype is merely a proxy for the resistance phenotype that directly informs patient care. One alternative approach currently under development attempts to circumvent the limitations of genotypic resistance testing by returning to a phenotypic approach, albeit one informed by genomic methods: transcriptional profiles serve as a rapid phenotypic signature for antibiotic response. Conceptually, since dying cells are transcriptionally distinct from cells fated to survive, susceptible bacteria enact different transcriptional profiles after antibiotic exposure that are different from the profiles of resistant strains, independent of the mechanism of resistance. These differences can be measured and, since transcription is one of the most rapid responses to cell stress (minutes to hours), can be used to determine whether cells are resistant or susceptible much more rapidly than is possible if one waits for growth in the presence of antibiotics (days). Like DNA, RNA can be readily detected through predictable rules governing base pairing via either amplification or hybridization-based methods. Changes in a carefully selected set of transcripts form an expression signature that can represent the total cellular response to antibiotic without requiring full characterization of the entire transcriptome. Preliminary proof-of-concept studies suggest that this approach may identify antibiotic susceptibility on the basis of transcriptional phenotype much more quickly than is possible with growth-based assays.

Because of its sensitivity in detecting even very rare nucleic acid fragments, sequencing is now permitting studies of unprecedented depth into complex populations of cells and tissues. The strength of

this depth and sensitivity applies not only to the detection of rare, novel pathogens in a sea of host signal but also to the identification of heterogeneous pathogen subpopulations in a single host that may differ, for example, in drug resistance profiles or pathogenesis determinants. Future studies will be needed to elucidate the clinical significance of these variable subpopulations, even as deep sequencing is now providing unprecedented levels of detail about majority and minority members of this population.

HOST-BASED DIAGNOSTICS

While pathogen-based diagnostics continue to be the mainstay for diagnosing infection, serologic testing has long been the basis of a strategy to diagnose infection by measuring host responses. Here, too, the application of genomics is now being explored to improve upon this approach, given the previously described limitations of serologic testing. Rather than using antibody responses as a retrospective biomarker for infection, recent efforts have focused on transcriptomic analysis of the host response as a new direction with diagnostic implications for human disease. For instance, while pathogen-based diagnostic tests to distinguish active from latent tuberculosis infection have proved elusive, recent work shows that the transcriptional profile of circulating white blood cells exhibits a differential pattern of expression of nearly 400 transcripts that distinguish active from latent tuberculosis; this expression pattern is driven in part by changes in interferon-inducible genes in the myeloid lineage. In a validation cohort, this transcriptional signature was able to distinguish patients with active versus latent disease, to distinguish tuberculosis infection from other pulmonary inflammatory states or infections, and to track responses to treatment in as little as 2 weeks, with normalization of expression toward that of patients without active disease over 6 months of effective therapy. Such a test could play an important role not only in the management of patients but also as a marker of efficacy in clinical trials of new therapeutic agents. Similarly, other investigators have been trying to identify host transcriptional signatures in circulating blood cells that are distinct in influenza A infection from those in upper respiratory infections caused by certain other viruses or bacteria. These signatures also varied with phase of infection and showed promise in distinguishing exposed subjects who will become symptomatic from those who will not. These results suggest that profiling of host transcriptional dynamics could augment the information obtained from studies of pathogens, both enhancing diagnosis and monitoring the progression of illness and the response to therapy.

In this era of genome-wide association studies and attempts to move toward personalized medicine, genomic approaches are also being applied to the identification of host genetic loci and factors that contribute to infection susceptibility. Such loci will have undergone strong selection among populations in which the disease is endemic. By identifying the beneficial genetic alleles among individuals who survive in such settings, markers for susceptibility or resistance are being discovered; these markers can be translated into diagnostic tests to identify susceptible individuals in order to implement preventive or prophylactic interventions. Further, such studies may offer mechanistic insight into the pathogenesis of infection and inform new methods of therapeutic intervention. Such beneficial genetic associations were recognized long before the advent of genomics, as in the protective effects of the negative Duffy blood group or heterozygous hemoglobin abnormalities against *Plasmodium* infection. Genomic methods enable more systematic and widespread investigations of the host to identify not only people with altered susceptibility to numerous diseases (e.g., HIV infection, tuberculosis, and cholera) but also host factors that contribute to and thus might predict the severity of disease.

THERAPEUTICS

Genomics has the potential to impact infectious disease therapeutics in two ways. By transforming the speed of diagnostic information acquisition or the type of diagnostic information that can be attained, it can influence therapeutic decision-making. Alternatively, by opening up new avenues to understanding pathogenesis, providing new

ways to disrupt infection, and delineating new approaches to antibiotic discovery, it can facilitate the development of new therapeutic agents.

GENOMIC DIAGNOSTICS INFORMING THERAPEUTICS

Efforts at antibiotic discovery are declining, with few new agents in the pipeline and even fewer entering the market. This phenomenon is due in part to the lack of economic incentives for the private sector; however, it is also attributable in part to the enormous challenges involved in the discovery and development of antibiotics. For obvious market-related reasons, nearly all efforts have focused on broad-spectrum antibiotics; the development of a chemical entity that works across an extremely diverse set of organisms (i.e., more divergent from each other than a human is from an amoeba) is far more challenging than the development of an agent that is designed to target a single bacterial species. Nevertheless, the concept of narrow-spectrum antibiotics has heretofore been rejected because of the lack of early diagnostic information that would guide the selection of such agents. Thus, rapid diagnostics providing antibiotic susceptibility information that can guide antibiotic selection in real time have the potential to alter and simplify antibiotic strategies by allowing a paradigm shift away from broad-spectrum drugs and toward narrow-spectrum agents. Such a paradigm shift clearly would have additional implications for the current escalation of antibiotic resistance.

In yet another diagnostic paradigm with the potential to impact therapeutic interventions, genomics is opening new avenues to a better understanding not only of different host susceptibilities to infection but also of different host responses to therapy. In a sense, the promise of “personalized medicine” has been a tantalizing holy grail. Some signs now point to the realization of this goal. For example, the role of glucocorticoids in tuberculous meningitis has been long debated. Recently, polymorphisms in the human genetic locus *LTA4H*, which encodes a leukotriene-modifying enzyme, were found to modulate the inflammatory response to tuberculosis. Patients with tuberculous meningitis who were homozygous for the proinflammatory *LTA4H* allele were most helped by adjunctive glucocorticoid treatment, while those who were homozygous for the anti-inflammatory allele were negatively affected by steroid treatment. This anti-inflammatory adjunct has become the standard of care in tuberculous meningitis, but this study suggests that perhaps only a subset of patients benefit and further suggests a genetic means of prospectively identifying this subset. Thus, genomic diagnostic tests may eventually inform diagnosis, prognosis, and treatment decisions by revealing the pathogenic potential of the microbe and detecting host responses to both infection and therapy.

GENOMICS IN DRUG AND VACCINE DEVELOPMENT

Genomic technologies are already dramatically changing research on host–pathogen interactions, with a goal of increasingly influencing the process of therapeutic discovery and development. Sequencing offers several possible avenues into antimicrobial therapeutic discovery. First, genomic-scale molecular methods have paved the way for comprehensive identification of all essential genes encoded within a pathogen’s genome, with consequent systematic identification of all possible vulnerabilities within a pathogen that could be targeted therapeutically. Second, transcriptional profiling can offer insights into mechanisms of action of new candidate drugs that emerge from screens. For instance, the transcriptional signature of cell wall disruptors (e.g., β -lactams) is distinct from that of DNA-damaging agents (e.g., fluoroquinolones) or protein synthesis inhibitors (e.g., aminoglycosides). Thus, transcriptional analysis of a pathogen’s response to a new antibiotic can either suggest a mechanism of action or flag compounds for prioritization because of a potentially novel activity. In an alternative genomic strategy for determining mechanisms of action, an RNA interference approach followed by targeted sequencing has been used to identify genes required for antitrypanosomal drug efficacy. This approach provided new insights into the mechanism of action of drugs that have been in use for decades for human African trypanosomiasis. Third, sequencing can readily identify the most conserved regions of a pathogen’s genomes and corresponding gene products; this information is invaluable in narrowing antigen candidates for vaccine development. These surface

776 proteins can be expressed recombinantly and tested for the ability to elicit a serologic response and protective immunity. This process, termed *reverse vaccinology*, has proved particularly useful for pathogens that are difficult to culture or poorly immunogenic, as was the case with the development of a vaccine for *Neisseria meningitidis* serogroup B.

Large-scale gene content analysis from sequencing or expression profiling enables new research directions that provide novel insights into the interplay of pathogen and host during infection or colonization. One important goal of such research is to suggest new therapeutic approaches to disrupt this interaction in favor of the host. Indeed, one of the most immediate applications of next-generation sequencing technology has come from simply characterizing human pathogens and related commensal or environmental strains and then finding genomic correlates for pathogenicity. For instance, as *Escherichia coli* varies from a simple nonpathogenic, lab-adapted strain (K-12) to a Shiga toxin-producing enterohemorrhagic gastrointestinal pathogen (O157:H7), it displays up to a 25% difference in gene content, even though its phylogenetic classification stays the same. Although this is an extreme example, it is not an isolated case. Some isolates of *Enterococcus*—notorious for its increasing incidence of resistance to common antibiotics such as ampicillin, vancomycin, and aminoglycosides—also contain recently acquired genetic material comprising up to 25% of the genome on mobile genetic elements. This fact suggests that horizontal gene transfer may play an important role in the organism's adaptation as a nosocomial pathogen. On closer study, this genome expansion has been demonstrated to be associated with loss of regulatory elements called CRISPRs (clustered, regularly interspaced short palindromic repeats). Loss of CRISPR elements, which protect the bacterial genome from invasion by certain foreign genetic materials, may thus facilitate the acquisition of antibiotic resistance-conferring genetic elements. While loss of this regulation appears to impose a competitive disadvantage in antibiotic-free environments, these drug-resistant strains thrive in the presence of even some of the most useful antienterococcal therapies. In addition to insights gained from genome sequencing, extension of unbiased whole-transcriptome sequencing (RNA-Seq) efforts to bacteria is beginning to identify unexpected regulatory, noncoding RNAs in many diverse species. While the functional implications of these new transcripts are as yet largely unknown, the presence of such features—conserved across many bacterial species—implies evolutionary importance and suggests areas for future study and possible new therapeutic avenues.

Thus, genomic studies are already beginning to transform our understanding of infection, offering evidence of virulence factors or toxins and providing insight into ongoing evolution of pathogenicity and drug resistance. One goal of such studies is to identify therapeutic agents that can disrupt the pathogenic process; there is currently much interest in the theoretical concept of antivirulence drugs that inhibit virulence factors rather than killing the pathogen outright as a means to intervene in infection. Further, as sequencing becomes increasingly accessible and efficient, large-scale studies with unprecedented statistical power to associate clinical outcomes with pathogen and host genotypes and thus to further reveal vulnerabilities in the infection process that can be targeted for disruption are being initiated. Although this is just the beginning, such studies point to a tantalizing future in which the clinician is armed with genomic predictors of infection outcome and therapeutic response to guide clinical decision-making.

EPIDEMIOLOGY OF INFECTIOUS DISEASES

Epidemiologic studies of infectious diseases have several main goals: to identify and characterize outbreaks, to describe the pattern and dynamics of an infectious disease as it spreads through populations, and to identify interventions that can limit or reduce the burden of disease. One classic, paradigmatic example is John Snow's elucidation of the origin of the 1854 London cholera outbreak. Snow used careful geographic mapping of cases to determine that the likely source of the outbreak was contaminated water from the Broad Street pump, and, by removing the pump handle, he aborted the outbreak. Whereas that intervention was undertaken without knowledge of the causative agent

of cholera, advances in microbiology and genomics have expanded the purview of epidemiology, which now considers not just the disease but also the pathogen, its virulence factors, and the complex relationships between microbial and host populations.

Through the use of novel genomic tools such as high-throughput sequencing, the diversity of a microbial population can now be rapidly described with unprecedented resolution, with discrimination between isolates that have single-nucleotide differences across the entire genome and advancement beyond prior approaches that relied on phenotypes (such as antibiotic resistance testing) or genetic markers (such as multilocus sequence typing). The development of statistical methods grounded in molecular genetics and evolutionary theory has established analytical approaches that translate descriptions of microbial population diversity and structure into insights into the origin and history of pathogen spread. By linking phylogenetic reconstruction with epidemiologic and demographic data, genomic epidemiology provides the opportunity to track transmission from person to person, to infer transmission patterns of both pathogens and sequence elements that confer phenotypes of interest, and to estimate the transmission dynamics of outbreaks.

DECIPHERING PERSON-TO-PERSON TRANSMISSION

The use of comparisons of whole-genome sequencing to infer person-to-person transmission and identify point-source outbreaks of pathogens has proved useful in hospital infection control settings. As reported in a seminal paper in 2010, a study of MRSA in a Thai hospital demonstrated that whole-genome sequencing can be used to infer transmission of a pathogen from patient to patient within a hospital setting through integration of the analysis of accumulation of mutations over time with dates and hospital locations of the infections. Since that time, multiple instances of the use of whole-genome sequencing to define and motivate interventions aimed at interrupting transmission chains have been reported. In another MRSA outbreak in a special-care baby unit in Cambridge, United Kingdom, whole-genome sequencing extended the traditional infection control analysis, which relies on typing organisms by their antibiotic susceptibilities, to sequencing of isolates from clinical samples. This approach identified an otherwise unrecognized outbreak of a specific MRSA strain that was occurring against a background of the usual pattern of infections caused by a diverse circulating population of MRSA strains. The analysis showed evidence of transmission among mothers within the special-care baby unit and in the community and demonstrated the key role of MRSA carriage in a single health care provider in the persistence of the outbreak. MRSA decolonization of the health care provider terminated the outbreak. In yet another example, in response to the observation of 18 cases of infection by carbapenemase-producing *Klebsiella pneumoniae* over 6 months at the National Institutes of Health Clinical Research Center, genome sequencing of the isolates was used to discriminate between the possibilities that these cases represented multiple, independent introductions into the health care system or a single introduction with subsequent transmission. On the basis of network and phylogenetic analysis of genomic and epidemiologic data, the authors reconstructed the likely relationships among the isolates from patient to patient, demonstrating that the spread of resistant *Klebsiella* infection was in fact due to nosocomial transmission of a single strain.

Uncovering of unexpected transmission events by genomic epidemiology studies is motivating renewed questioning of pathogen ecology and modes of transmission. For example, the rise in prevalence of infections with nontuberculous mycobacteria, including *Mycobacterium abscessus*, among patients with cystic fibrosis (CF) has led to speculation about the possible role of patient-to-patient transmission in the CF community; however, conventional typing approaches have lacked the resolution to define population structure accurately, a critical component of inferring transmission. Past infection control guidelines discounted the possibility of acquisition of nontuberculous mycobacteria in health care settings, as no strong evidence for such transmission had been described. In a whole-genome sequencing study of *M. abscessus* isolates from patients with CF, an analytical approach using genome sequencing, epidemiology, and

Bayesian modeling examined the likelihood of transmission between patients within a CF center; the authors found nearly identical isolates in a number of patients and observed that these isolates were less diverse than isolates from a single individual. Because no clear epidemiologic link places the infected patients in the same place at the same time, this finding highlights a need to explore preexisting notions of circumstances required for transmission and a reconsideration of *M. abscessus* infection control guidelines. Similar studies of other pathogens—particularly those that share human, other animal host, and environmental reservoirs—will continue to advance our insight into the relative roles and prominence of sources of infection as well as the modes of spread through populations, thereby establishing evidence-based strategies for prevention and intervention.

As increasing numbers of studies aim to carefully define the origins and spread of infectious agents using the high-resolution lens of whole-genome sequencing, fundamental questions are arising with regard to our understanding of infection in a single individual and the process of a single transmission event. For example, a better understanding of a pathogen population's diversity within a single infected individual is a critical component in interpreting the relationship among isolates from different patients. While we have traditionally thought of individuals as infected with a single bacterial strain, a recent sequencing study of multiple colonies of *S. aureus* from a single individual showed a “cloud” of diversity; this finding raises a number of questions that will be important to address as this field develops: What is the clinical significance of this diversity? What are the processes that generate and limit diversity? What amount of diversity is transmitted under different conditions and routes of transmission? How do the answers to these questions vary by infectious organism, type of infection, and host and in response to treatment? More comprehensive descriptions of diversity, population dynamics, transmission bottlenecks, and the forces that shape and influence the growth and spread of microbial populations will be a critically important focus of future investigations.

RECONSTRUCTING THE ORIGINS AND DYNAMICS OF PATHOGEN SPREAD

In addition to reconstructing the transmission chains of local outbreaks, genomics-based epidemiologic methods are providing insight into broad-scale geographic and temporal spread of pathogens. A classic example has been the study of cholera, the dehydrating diarrheal illness caused by infection with *Vibrio cholerae*. Cholera first spread worldwide from the Indian subcontinent in the 1800s and has since caused seven pandemics; the seventh pandemic has been ongoing since the 1960s. An investigation into the geographic patterns of

cholera spread in the seventh pandemic used genome sequences from a global collection of 154 *V. cholerae* strains representing isolates from 1957–2010. This investigation revealed that the seventh pandemic has comprised at least three overlapping waves spreading out from the Indian subcontinent (Fig. 146-4). Further, analysis of the genome of an isolate of *V. cholerae* from the 2010 outbreak of cholera in Haiti showed it to be more closely related to isolates from South Asia than to isolates from neighboring Latin America, a result supporting the hypothesis that the outbreak was derived from *V. cholerae* introduced into Haiti by human travel (likely from Nepal) rather than by environmental or more geographically proximal sources. A subsequent study that dated the time to the most recent common ancestor of a population of *V. cholerae* isolates from Haiti provided further support for a single point-source introduction from Nepal.

Increasing numbers of investigations into the spread of many pathogens—thus far including strains of *S. aureus*, *S. pneumoniae*, *Chlamydia*, *Salmonella*, *Shigella*, *E. coli*, *C. difficile*, West Nile virus, rabies virus, and dengue virus—are contributing to a growing atlas of maps describing routes, patterns, and tempos of microbial diversification and dissemination. Large-scale efforts like the 100K Foodborne Pathogen Genome Project, which aims to sequence the genomes of 100,000 strains of food-borne pathogens collected from sources including food, the environment, and farm animals, are possible because of advances in sequencing technologies. Such studies will yield a vast amount of data that can be used to investigate diversity and microbiologic links within distinct niches and the patterns of spread from one niche to another. The increasingly broad adoption of genome sequencing by health care and public health institutions will ensure that the available catalog of genome sequences and associated epidemiologic data will grow very rapidly. With higher-resolution description of microbial diversity and of the dynamics of that diversity over time and across epidemiologic and demographic boundaries and evolutionary niches, we will gain even greater insights into the relationships of transmission routes and patterns of historic spread.

PREDICTING EPIDEMIC POTENTIAL

Defining pathogen transmissibility is a critical step in the development of public health surveillance and intervention strategies as this information can help predict the epidemic potential of an outbreak. Transmissibility can be estimated by a variety of methods, including inference from the growth rate of an epidemic together with the generation time of an infection (the mean interval between infection of an index case and of the people infected by that index case). Genome

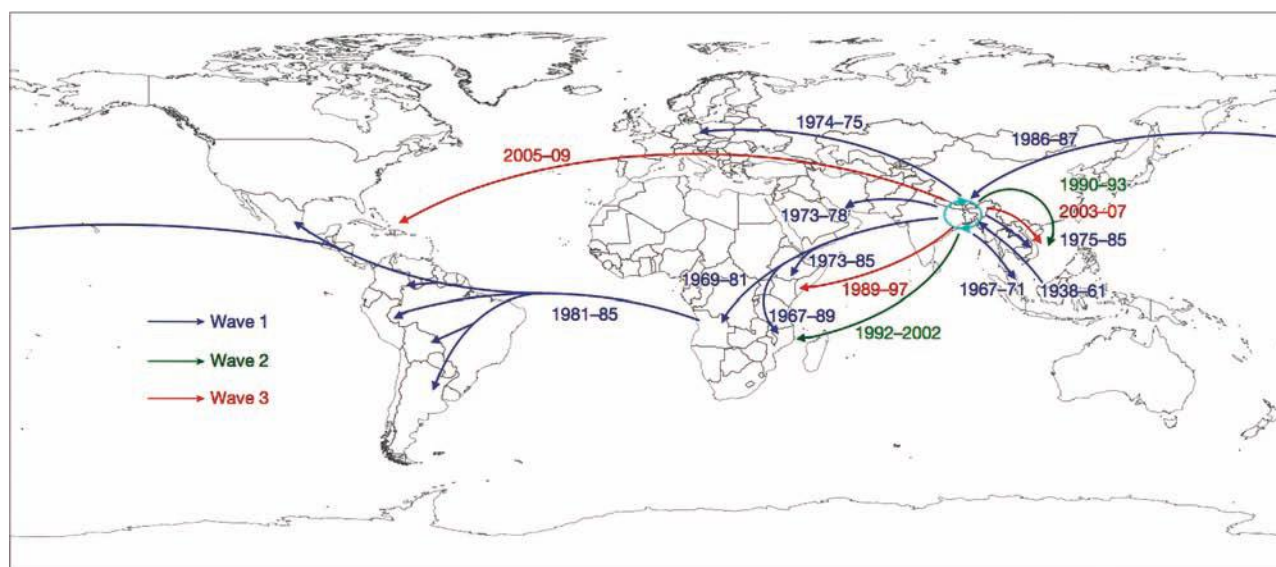


FIGURE 146-4 Transmission events inferred from phylogenetic reconstruction of 154 *Vibrio cholerae* isolates from the seventh cholera pandemic. Date ranges represent estimated time to the most recent common ancestor for strains transmitted from source to destination locations, based on a Bayesian model of the phylogeny. (Reprinted with permission from A Mutreja et al: Evidence for several waves of global transmission in the seventh cholera pandemic. *Nature* 477:462, 2011.)

778 sequencing and analysis of a well-sampled population provide another method by which to derive similar fundamental epidemiologic parameters. One key measure of transmissibility is the basic reproduction number (R_0), defined as the number of secondary infections generated from a single primary infectious case. When the basic reproduction number is greater than 1 ($R_0 > 1$), an outbreak has epidemic potential; when it is less than 1 ($R_0 < 1$), the outbreak will become extinct. On the basis of sequences from influenza samples obtained from infected patients very early in the 2009 H1N1 influenza pandemic, the basic reproduction number was estimated through a population genomic analysis at 1.2; this compared to estimates of 1.4–1.6 based on several epidemiologic analyses. In addition, with the assumption of a molecular clock model, sequences of H1N1 samples together with information about when and where the samples were obtained have been used to estimate the date and location of the pandemic's origin, providing insight into disease origins and dynamics. Because the magnitude and intensity of the public health response are guided by the predicted size of an outbreak, the ability of genomic methods to elucidate a pathogen's origin and epidemic potential adds an important dimension to the contributions of these methods to infectious disease epidemiology.

PROVIDING INSIGHT INTO PATHOGEN EVOLUTION

Beyond describing transmission and dynamics, pathogen genomics can provide insight into the evolution of pathogens and the interaction of selective pressures, the host, and pathogen populations, which can have implications for vaccine or therapeutic development. From a clinical perspective, this process is central to the acquisition of antibiotic resistance, the generation of increasing pathogenicity or new virulence traits, the evasion of host immunity and clearance (leading to chronic infection), and vaccine efficacy.

Microbial genomes evolve through a variety of mechanisms, including mutation, duplication, insertion, deletion, recombination, and horizontal gene transfer. Segmented viruses (e.g., influenza virus) can reassort gene segments within multiply infected cells. The pandemic 2009 H1N1 influenza A virus, for example, appears to have been generated through reassortment of several avian, swine, and human influenza strains. Such potential for the evolution of novel pandemic strains has precipitated concern about the possible evolution to transmissibility of virulent strains that have been associated with high mortality rates but have not yet exhibited efficient human infectivity. Controversial experiments with H5N1 avian influenza virus, for example, have defined five mutations that render the virus transmissible, at least in ferrets—the animal model system for human influenza.

The continual antigenic evolution of seasonal influenza offers an example of how studies of pathogen evolution can impact surveillance and vaccine development. Frequent updates to the annual influenza vaccine are needed to ensure protection against the dominant strains. These updates are based on an ability to anticipate which viral populations from a pool of substantial locally and globally diverse circulating viruses will predominate in the upcoming season. Toward that end, sequencing-based studies of influenza virus dynamics have shed light on the global spread of influenza, providing concrete data on patterns of spread and helping to elucidate the origins, emergence, and circulation of novel strains. Through analysis of more than 1000 influenza A H3N2 virus isolates over the 2002–2007 influenza seasons, Southeast Asia was identified as the usual site from which diversity originates and spreads worldwide. Further studies of global isolate collections have shed further light on the diversity of circulating virus, showing that some strains persist and circulate outside of Asia for multiple seasons.

Not only do genomic epidemiology studies have the potential to help guide vaccine selection and development; they are also helping to track what happens to pathogens circulating in the population in response to vaccination. By describing pathogen evolution under the selective pressure of a vaccinated population, such studies can play a key role in surveillance and identification of virulence determinants and perhaps may even help to predict the future evolution of escape from vaccine protection. The 7-valent pneumococcal conjugate vaccine (PCV-7) targeted the seven serotypes of *S. pneumoniae* responsible for

the majority of cases of invasive disease at the time of its introduction in 2000; since then, PCV-7 has dramatically reduced the incidence of pneumococcal disease and mortality. Population genomic analysis of the sequences of more than 600 Massachusetts pneumococcal isolates from 2001–2007 has shown that preexisting rare nonvaccine serotypes are replacing vaccine serotypes and that some strains have persisted despite vaccination by recombining the vaccine-targeted capsule locus with a cassette of capsule genes from non-vaccine-targeted serotypes.

GLOBAL CONSIDERATIONS



While cutting-edge genomic technologies are largely implemented in the developed world, their application to infectious diseases offers perhaps the biggest potential impact in less developed regions where the burden of these infections is greatest. This globalization of genomic technology and its extensions has already begun in each of the areas of focus highlighted in this chapter; it has occurred both through the application of advanced technologies to samples collected in the developing world and through the adaptation and importation of technologies directly to the developing world for on-site implementation as they become more globally accessible. Genomic characterization of the pathogens responsible for important global illnesses such as tuberculosis, malaria, trypanosomiasis, and cholera has led to insights in diagnosis, treatment, and infection control. For instance, the nucleic acid–based test developed for rapid diagnosis of *M. tuberculosis* infection and detection of rifampin resistance is being priced for implementation in field settings in Africa and Asia where tuberculosis is most prevalent. The potential to diagnose multi-drug-resistant tuberculosis in hours instead of weeks or months may truly revolutionize treatment and control of this common and devastating illness. High-resolution genomic tracking of the spread of cholera has yielded insights into which public health measures may prove most effective in controlling local epidemics. Overall, sequencing efforts have become exponentially cheaper with each passing year. As these technologies synergize with efforts to globalize information-technology resources, global implementation of genomic methods promises to spread state-of-the-art methods for diagnosis, treatment, and epidemic tracking of infections to areas that need these capabilities the most.

SUMMARY

By illuminating the genetic information that encodes the most fundamental processes of life, genomic technologies are transforming many aspects of medicine. In infectious diseases, methods such as next-generation sequencing and genome-scale expression analysis offer information of unprecedented depth about individual microbes as well as microbial communities. This information is expanding our understanding of the interactions of these microorganisms and communities with one another, with their human hosts, and with the environment. Despite significant progress and the abundant genomic data now available, technological and financial barriers continue to impede the widespread adoption of large-scale pathogen sequencing in clinical, public health, and research settings. As even vaster amounts of data are generated, innovations in storage, development of bioinformatics tools to manipulate the data, standardization of methods, and training of end-users in both the research and clinical realms will be required. The cost-effectiveness and applicability of whole-genome sequencing, particularly in the clinic, remain to be studied, and studies of the impact of genome sequencing on patient outcomes will be needed to clarify the contexts in which these new methodologies can make the greatest contributions to patient well-being. The ongoing efforts to overcome limitations through collaboration, teaching, and reduction of financial obstacles should be applauded and expanded. With advances in genomic technologies and computational analysis, our ability to detect, characterize, treat, monitor, prevent, and control infections has progressed rapidly in recent years and will continue to do so, with the hope of heralding a new era where the clinician is better armed to combat infection and promote human health.

147 Approach to the Acutely Ill Infected Febrile Patient

Tamar F. Barlam, Dennis L. Kasper

The physician treating the acutely ill febrile patient must be able to recognize infections that require emergent attention. If such infections are not adequately evaluated and treated at initial presentation, the opportunity to alter an adverse outcome may be lost. In this chapter, the clinical presentations of and approach to patients with relatively common infectious disease emergencies are discussed. These infectious processes and their treatments are discussed in detail in other chapters.

APPROACH TO THE PATIENT: Acute Febrile Illness

Before the history is elicited and a physical examination is performed, an immediate assessment of the patient's general appearance can yield valuable information. The perceptive physician's subjective sense that a patient is septic or toxic often proves accurate. Visible agitation or anxiety in a febrile patient can be a harbinger of critical illness.

HISTORY Presenting symptoms are frequently nonspecific. Detailed questions should be asked about the onset and duration of symptoms and about changes in severity or rate of progression over time. Host factors and comorbid conditions may increase the risk of infection with certain organisms or of a more fulminant course than is usually seen. Lack of splenic function, alcoholism with significant liver disease, IV drug use, HIV infection, diabetes, malignancy, organ transplantation, and chemotherapy all predispose to specific infections and frequently to increased severity. The patient should be questioned about factors that might help identify a nidus for invasive infection, such as recent upper respiratory tract infections, influenza, or varicella; prior trauma; disruption of cutaneous barriers due to lacerations, burns, surgery, body piercing, or decubiti; and the presence of foreign bodies, such as nasal packing after rhinoplasty, tampons, or prosthetic joints. Travel, contact with pets or other animals, or activities that might result in tick or mosquito exposure can lead to diagnoses that would not otherwise be considered. Recent dietary intake, medication use, social or occupational contact with ill individuals, vaccination history, recent sexual contacts, and menstrual history may be relevant. A review of systems should focus on any neurologic signs or sensorium alterations, rashes or skin lesions, and focal pain or tenderness and should also include a general review of respiratory, gastrointestinal, or genitourinary symptoms.

PHYSICAL EXAMINATION A complete physical examination should be performed, with special attention to several areas that are sometimes given short shrift in routine examinations. Assessment of the patient's general appearance and vital signs, skin and soft tissue examination, and the neurologic evaluation are of particular importance.

The patient may appear either anxious and agitated or lethargic and apathetic. Fever is usually present, although elderly patients and compromised hosts (e.g., patients who are uremic or cirrhotic and those who are taking glucocorticoids or nonsteroidal anti-inflammatory drugs) may be afebrile despite serious underlying infection. Measurement of blood pressure, heart rate, and respiratory rate helps determine the degree of hemodynamic and metabolic compromise. The patient's airway must be evaluated to rule out the risk of obstruction from an invasive oropharyngeal infection.

The etiologic diagnosis may become evident in the context of a thorough skin examination ([Chap. 24](#)). Petechial rashes are

typically seen with meningococemia or Rocky Mountain spotted fever (RMSF; [see Fig. 25e-16](#)); erythroderma is associated with toxic shock syndrome (TSS) and drug fever. The soft tissue and muscle examination is critical. Areas of erythema or dusky skin, edema, and tenderness may indicate underlying necrotizing fasciitis, myositis, or myonecrosis. The neurologic examination must include a careful assessment of mental status for signs of early encephalopathy. Evidence of nuchal rigidity or focal neurologic findings should be sought.

DIAGNOSTIC WORKUP After a quick clinical assessment, diagnostic material should be obtained rapidly and antibiotic and supportive treatment begun. Blood (for cultures; baseline complete blood count with differential; measurement of serum electrolytes, blood urea nitrogen, serum creatinine, and serum glucose; and liver function tests) can be obtained at the time an IV line is placed and before antibiotics are administered. The blood lactate concentration also should be measured. Three sets of blood cultures should be performed for patients with possible acute endocarditis. Asplenic patients should have a buffy coat examined for bacteria; these patients can have $>10^6$ organisms per milliliter of blood (compared with 10^4 /mL in patients with an intact spleen). Blood smears from patients at risk for severe parasitic disease, such as malaria or babesiosis ([Chap. 250e](#)), must be examined for the diagnosis and quantitation of parasitemia. Blood smears may also be diagnostic in ehrlichiosis and anaplasmosis.

Patients with possible meningitis should have cerebrospinal fluid (CSF) drawn before the initiation of antibiotic therapy. Focal findings, depressed mental status, or papilledema should be evaluated by brain imaging prior to lumbar puncture, which, in this setting, could initiate herniation. *Antibiotics should be administered before imaging but after blood for cultures has been drawn.* If CSF cultures are negative, blood cultures will provide the diagnosis in 50–70% of cases. Molecular diagnostic techniques (e.g., broad-range 16S rRNA gene polymerase chain reaction testing for bacterial meningitis pathogens) are of increasing importance in the rapid diagnosis of life-threatening infections.

Focal abscesses necessitate immediate CT or MRI as part of an evaluation for surgical intervention. Other diagnostic procedures, such as wound cultures, should not delay the initiation of treatment for more than minutes. Once emergent evaluation, diagnostic procedures, and (if appropriate) surgical consultation (see below) have been completed, other laboratory tests can be conducted. Appropriate radiography, computed axial tomography, MRI, urinalysis, erythrocyte sedimentation rate and C-reactive protein determination, and transthoracic or transesophageal echocardiography all may prove important.

TREATMENT THE ACUTELY ILL PATIENT

In the acutely ill patient, empirical antibiotic therapy is critical and should be administered without undue delay. Increased prevalence of antibiotic resistance in community-acquired bacteria must be considered when antibiotics are selected. [Table 147-1](#) lists first-line empirical regimens for infections considered in this chapter. In addition to the rapid initiation of antibiotic therapy, several of these infections require urgent surgical attention. Neurosurgical evaluation for subdural empyema, otolaryngologic surgery for possible mucormycosis, and cardiothoracic surgery for critically ill patients with acute endocarditis are as important as antibiotic therapy. For infections such as necrotizing fasciitis and clostridial myonecrosis, rapid surgical intervention supersedes other diagnostic or therapeutic maneuvers.

Adjunctive treatments may reduce morbidity and mortality rates and include dexamethasone for bacterial meningitis or IV immunoglobulin for TSS and necrotizing fasciitis caused by group A *Streptococcus*. Adjunctive therapies should usually be initiated within the first hours of treatment; however, dexamethasone for

TABLE 147-1 EMPIRICAL TREATMENT FOR COMMON INFECTIOUS DISEASE EMERGENCIES^a

Clinical Syndrome	Possible Etiologies	Treatment	Comments	See Chap(s).
Sepsis without a Clear Focus				
Septic shock	<i>Pseudomonas</i> spp., gram-negative enteric bacilli, <i>Staphylococcus</i> spp., <i>Streptococcus</i> spp.	Vancomycin (15 mg/kg q12h) ^b plus gentamicin (5 mg/kg per day) plus either Piperacillin/tazobactam (3.375 g q4h) or cefepime (2 g q8h) ^c	Adjust treatment when culture data become available.	172, 173, 186, 189, 325
Overwhelming post-splenectomy sepsis	<i>Streptococcus pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Neisseria meningitidis</i>	Ceftriaxone (2 g q12h) plus vancomycin (15 mg/kg q12h) ^b	If a β -lactam-sensitive strain is identified, vancomycin can be discontinued.	325
Babesiosis	<i>Babesia microti</i> (U.S.), <i>B. divergens</i> (Europe)	Clindamycin (600 mg q8h) plus quinine (650 mg q8h)	Atovaquone and azithromycin can be used in less severe disease and are associated with fewer side effects. Treatment with doxycycline (100 mg bid) for potential co-infection with <i>Borrelia burgdorferi</i> or <i>Anaplasma</i> spp. may be prudent.	246e, 249
Sepsis with Skin Findings				
Meningococcemia	<i>N. meningitidis</i>	Penicillin (4 mU q4h) or ceftriaxone (2 g q12h)	Consider protein C replacement, if available, in fulminant meningococcemia. Drotrecogin alfa (activated) is no longer produced.	180
Rocky Mountain spotted fever (RMSF)	<i>Rickettsia rickettsii</i>	Doxycycline (100 mg bid)	If both meningococcemia and RMSF are being considered, use ceftriaxone (2 g q12h) plus doxycycline (100 mg bid). If RMSF is diagnosed, doxycycline is the proven superior agent.	211
Purpura fulminans	<i>S. pneumoniae</i> , <i>H. influenzae</i> , <i>N. meningitidis</i>	Ceftriaxone (2 g q12h) plus vancomycin (15 mg/kg q12h) ^b	If a β -lactam-sensitive strain is identified, vancomycin can be discontinued.	171, 180, 182, 325
Erythroderma: toxic shock syndrome	Group A <i>Streptococcus</i> , <i>Staphylococcus aureus</i>	Vancomycin (15 mg/kg q12h) ^b plus clindamycin (600 mg q8h)	If a penicillin- or oxacillin-sensitive strain is isolated, these agents are superior to vancomycin (penicillin, 2 mU q4h; or oxacillin, 2 g IV q4h). The site of toxigenic bacteria should be debrided; IV immunoglobulin can be used in severe cases. ^d	172, 173
Sepsis with Soft Tissue Findings				
Necrotizing fasciitis	Group A <i>Streptococcus</i> , mixed aerobic/anaerobic flora, CA-MRSA ^e	Vancomycin (15 mg/kg q12h) ^b plus clindamycin (600 mg q8h) plus gentamicin (5 mg/kg q8h)	Urgent surgical evaluation is critical. Adjust treatment when culture data become available.	156, 172, 173
Clostridial myonecrosis	<i>Clostridium perfringens</i>	Penicillin (2 mU q4h) plus clindamycin (600 mg q8h)	Urgent surgical evaluation is critical.	179
Neurologic Infections				
Bacterial meningitis	<i>S. pneumoniae</i> , <i>N. meningitidis</i>	Ceftriaxone (2 g q12h) plus vancomycin (15 mg/kg q12h) ^b	If a β -lactam-sensitive strain is identified, vancomycin can be discontinued. If the patient is >50 years old or has comorbid disease, add ampicillin (2 g q4h) for <i>Listeria</i> coverage. Dexamethasone (10 mg q6h \times 4 days) improves outcome in adults with meningitis (especially pneumococcal) and cloudy CSF, positive CSF Gram's stain, or a CSF leukocyte count >1000/mL.	164
Brain abscess, suppurative intracranial infections	<i>Streptococcus</i> spp., <i>Staphylococcus</i> spp., anaerobes, gram-negative bacilli	Vancomycin (15 mg/kg q12h) ^b plus metronidazole (500 mg q8h) plus ceftriaxone (2 g q12h)	Urgent surgical evaluation is critical. If a penicillin- or oxacillin-sensitive strain is isolated, these agents are superior to vancomycin (penicillin, 4 mU q4h; or oxacillin, 2 g q4h).	164
Cerebral malaria	<i>Plasmodium falciparum</i>	Artesunate (2.4 mg/kg IV at 0, 12, and 24 h; then once daily) ^f or quinine (IV loading dose of 20 mg salt/kg; then 10 mg/kg q8h)	Do not use glucocorticoids. Use IV quinidine if IV quinine is not available. During IV quinidine treatment, blood pressure and cardiac function should be monitored continuously and blood glucose periodically.	246e, 248
Spinal epidural abscess	<i>Staphylococcus</i> spp., gram-negative bacilli	Vancomycin (15 mg/kg q12h) ^b plus ceftriaxone (2 g q24h)	Surgical evaluation is essential. If a penicillin- or oxacillin-sensitive strain is isolated, these agents are superior to vancomycin (penicillin, 4 mU q4h; or oxacillin, 2 g q4h).	456
Focal Infections				
Acute bacterial endocarditis	<i>S. aureus</i> , β -hemolytic streptococci, HACEK group, ^g <i>Neisseria</i> spp., <i>S. pneumoniae</i>	Ceftriaxone (2 g q12h) plus vancomycin (15 mg/kg q12h) ^b	Adjust treatment when culture data become available. Surgical evaluation is essential.	155

^aThese empirical regimens include coverage for gram-positive pathogens that are resistant to β -lactam antibiotics. Local resistance patterns should be considered and may alter the need for empirical vancomycin. ^bA vancomycin loading dose of 20–25 mg/kg can be considered in critically ill patients. ^c β -Lactam antibiotics may exhibit unpredictable pharmacodynamics in sepsis. Prolonged or continuous infusions can be considered. ^dThe optimal dose of IV immunoglobulin has not been determined, but the median dose in observational studies is 2 g/kg (total dose administered for 1–5 days). ^eCommunity-acquired methicillin-resistant *S. aureus*. ^fIn the United States, artesunate must be obtained through the Centers for Disease Control and Prevention. For patients diagnosed with severe malaria, full doses of parenteral antimalarial treatment should be started with whichever recommended antimalarial agent is first available. ^g*Haemophilus* species, *Aggregatibacter* species, *Cardiobacterium hominis*, *Eikenella corrodens*, and *Kingella kingae*.

bacterial meningitis must be given before or at the time of the first dose of antibiotic. Glucocorticoids can also be harmful, sometimes resulting in worse outcomes—e.g., when given in the setting of cerebral malaria or viral hepatitis.

SPECIFIC PRESENTATIONS

The infections considered below according to common clinical presentation can have rapidly catastrophic outcomes, and their immediate recognition and treatment can be life-saving. Recommended empirical therapeutic regimens are presented in [Table 147-1](#).

SEPSIS WITHOUT AN OBVIOUS FOCUS OF PRIMARY INFECTION

Patients initially have a brief prodrome of nonspecific symptoms and signs that progresses quickly to hemodynamic instability with hypotension, tachycardia, tachypnea, respiratory distress, and altered mental status. Disseminated intravascular coagulation (DIC) with clinical evidence of a hemorrhagic diathesis is a poor prognostic sign.

Septic Shock (See also [Chap. 325](#)) Patients with bacteremia leading to septic shock may have a primary site of infection (e.g., pneumonia, pyelonephritis, or cholangitis) that is not evident initially. Elderly patients with comorbid conditions, hosts compromised by malignancy and neutropenia, and patients who have recently undergone a surgical procedure or hospitalization are at increased risk for an adverse outcome. Gram-negative bacteremia with organisms such as *Pseudomonas aeruginosa* or *Escherichia coli* and gram-positive infection with organisms such as *Staphylococcus aureus* (including methicillin-resistant *S. aureus* [MRSA]) or group A streptococci can present as intractable hypotension and multiorgan failure. Treatment can usually be initiated empirically on the basis of the presentation, host factors ([Chap. 325](#)), and local patterns of bacterial resistance. Outcomes are worse when antimicrobial treatment is delayed or when the responsible pathogen ultimately proves not to be susceptible to the initial regimen. Broad-spectrum antimicrobial agents are therefore recommended and should be instituted rapidly, preferably within the first hour after presentation. Risk factors for fungal infection should be assessed, as the incidence of fungal septic shock is increasing. Biomarkers such as C-reactive protein and procalcitonin have not proved reliable diagnostically but, when measured over time, can facilitate appropriate de-escalation of therapy. Glucocorticoids should be considered only for patients with severe sepsis who do not respond to fluid resuscitation and vasopressor therapy.

Overwhelming Infection in Asplenic Patients (See also [Chap. 325](#))

Patients without splenic function are at risk for overwhelming bacterial sepsis. Asplenic adult patients succumb to sepsis at 58 times the rate of the general population. Most infections are thought to occur within the first 2 years after splenectomy, with a mortality rate of ~50%, but the increased risk persists throughout life. In asplenia, encapsulated bacteria cause the majority of infections. Adults, who are more likely to have antibody to these organisms, are at lower risk than children. *Streptococcus pneumoniae* is the most common isolate, causing 50–70% of cases, but the risk of infection with *Haemophilus influenzae* or *Neisseria meningitidis* is also high. Severe clinical manifestations of infections due to *E. coli*, *S. aureus*, group B streptococci, *P. aeruginosa*, *Bordetella holmesii*, and *Capnocytophaga*, *Babesia*, and *Plasmodium* species have been described.

Babesiosis (See also [Chap. 249](#)) A history of recent travel to endemic areas raises the possibility of infection with *Babesia*. Between 1 and 4 weeks after a tick bite, the patient experiences chills, fatigue, anorexia, myalgia, arthralgia, shortness of breath, nausea, and headache; ecchymosis and/or petechiae are occasionally seen. The tick that most commonly transmits *Babesia*, *Ixodes scapularis*, also transmits *Borrelia burgdorferi* (the agent of Lyme disease) and *Anaplasma*; coinfection can occur, resulting in more severe disease. Infection with the European species *Babesia divergens* is more frequently fulminant than that due to the U.S. species *Babesia microti*. *B. divergens* causes a febrile syndrome with hemolysis, jaundice, hemoglobinemia, and renal failure and is associated with a mortality rate of >40%. Severe babesiosis

is especially common in asplenic hosts but does occur in hosts with normal splenic function, particularly those >60 years of age and those with underlying immunosuppressive conditions such as HIV infection or malignancy. Complications include renal failure, acute respiratory failure, and DIC.

Other Sepsis Syndromes Tularemia ([Chap. 195](#)) is seen throughout the United States but occurs primarily in Arkansas, Missouri, South Dakota, and Oklahoma. This disease is associated with wild rabbit, tick, and tabanid fly contact. It can be transmitted by arthropod bite, handling of infected animal carcasses, consumption of contaminated food and water, or inhalation. The typhoidal form can be associated with gram-negative septic shock and a mortality rate of >30%, especially in patients with underlying comorbid or immunosuppressive conditions. Plague occurs infrequently in the United States ([Chap. 196](#)), primarily after contact with ground squirrels, prairie dogs, or chipmunks, but is endemic in other parts of the world, with >90% of all cases occurring in Africa. The septic form is particularly rare and is associated with shock, multiorgan failure, and a 30% mortality rate. These infections should be considered in the appropriate epidemiologic setting. The Centers for Disease Control and Prevention lists *Francisella tularensis* and *Yersinia pestis* (the agents of tularemia and plague, respectively) along with *Bacillus anthracis* (the agent of anthrax) as important organisms that might be used for bioterrorism ([Chap. 261e](#)).

SEPSIS WITH SKIN MANIFESTATIONS

(See also [Chap. 24](#)) Maculopapular rashes may reflect early meningococcal or rickettsial disease but are usually associated with nonemergent infections. Exanthems are usually viral. Primary HIV infection commonly presents with a rash that is typically maculopapular and involves the upper part of the body but can spread to the palms and soles. The patient is usually febrile and can have lymphadenopathy, severe headache, dysphagia, diarrhea, myalgias, and arthralgias. Recognition of this syndrome provides an opportunity to prevent transmission and to institute treatment and monitoring early on.

Petechial rashes caused by viruses are seldom associated with hypotension or a toxic appearance, although there can be exceptions (e.g., severe measles or arboviral infection). Petechial rashes limited to the distribution of the superior vena cava are rarely associated with severe disease. In other settings, petechial rashes require more urgent attention.

Meningococcemia (See also [Chap. 180](#))

Almost three-quarters of patients with *N. meningitidis* bacteremia have a rash. Meningococcemia most often affects young children (i.e., those 6 months to 5 years old). In sub-Saharan Africa, the high prevalence of serogroup A meningococcal disease has been a threat to public health for more than a century. Thousands of deaths occur annually in this area, which is known as the “meningitis belt,” and large epidemic waves occur approximately every 8–12 years. Serogroups W135 and X are also important emerging pathogens in Africa. In the United States, sporadic cases and outbreaks occur in day-care centers, schools (grade school through college, particularly among college freshmen living in residential halls), and army barracks. Household contacts of index cases are at 400–800 times greater risk of disease than the general population. Patients may exhibit fever, headache, nausea, vomiting, myalgias, changes in mental status, and meningismus. However, the rapidly progressive form of disease is not usually associated with meningitis. The rash is initially pink, blanching, and maculopapular, appearing on the trunk and extremities, but then becomes hemorrhagic, forming petechiae. Petechiae are first seen at the ankles, wrists, axillae, mucosal surfaces, and palpebral and bulbar conjunctiva, with subsequent spread on the lower extremities and to the trunk. A cluster of petechiae may be seen at pressure points—e.g., where a blood pressure cuff has been inflated. In rapidly progressive meningococcemia (10–20% of cases), the petechial rash quickly becomes purpuric (see [Fig. 70-5](#)), and patients develop DIC, multiorgan failure, and shock; 50–60% of these patients die, and survivors often require extensive debridement or amputation of gangrenous extremities.

782 Hypotension with petechiae for <12 h is associated with significant mortality. Cyanosis, coma, oliguria, metabolic acidosis, and elevated partial thromboplastin time also are associated with a fatal outcome. Correction of protein C deficiency may improve outcome. Antibiotics given in the office by the primary care provider before hospital evaluation and admission may improve prognosis; this observation suggests that early initiation of treatment may be life-saving. Meningococcal conjugate vaccines are protective against serogroups A, C, Y and W135 and are recommended for children 11–18 years of age and for other high-risk patients.

Rocky Mountain Spotted Fever (See also Chap. 211) RMSF is a tick-borne disease caused by *Rickettsia rickettsii* that occurs throughout North and South America. Up to 40% of patients do not report a history of a tick bite, but a history of travel or outdoor activity (e.g., camping in tick-infested areas) can often be ascertained. For the first 3 days, headache, fever, malaise, myalgias, nausea, vomiting, and anorexia are documented. By day 3, half of patients have skin findings. Blanching macules develop initially on the wrists and ankles and then spread over the legs and trunk. The lesions become hemorrhagic and are frequently petechial. The rash spreads to palms and soles later in the course. The centripetal spread is a classic feature of RMSF but occurs in a minority of patients. Moreover, 10–15% of patients with RMSF never develop a rash. The patient can be hypotensive and develop noncardiogenic pulmonary edema, confusion, lethargy, and encephalitis progressing to coma. The CSF contains 10–100 cells/ μ L, usually with a predominance of mononuclear cells. The CSF glucose level is often normal; the protein concentration may be slightly elevated. Renal and hepatic injury as well as bleeding secondary to vascular damage are noted. For untreated infections, mortality rates are 20–30%. Delayed recognition and treatment are associated with a greater risk of death; Native Americans, children 5–9 years of age, adults >70 years old, and persons with underlying immunosuppression also are at increased risk of death.



Other rickettsial diseases cause significant morbidity and mortality worldwide. *Mediterranean spotted fever* caused by *Rickettsia conorii* is found in Africa, southwestern and south-central Asia, and southern Europe. Patients have fever, flu-like symptoms, and an inoculation eschar at the site of the tick bite. A maculopapular rash develops within 1–7 days, involving the palms and soles but sparing the face. Elderly patients or those with diabetes, alcoholism, uremia, or congestive heart failure are at risk for severe disease characterized by neurologic involvement, respiratory distress, and gangrene of the digits. Mortality rates associated with this severe form of disease approach 50%. *Epidemic typhus*, caused by *Rickettsia prowazekii*, is transmitted in louse-infested environments and emerges in conditions of extreme poverty, war, and natural disaster. Patients experience a sudden onset of high fevers, severe headache, cough, myalgias, and abdominal pain. A maculopapular rash develops (primarily on the trunk) in more than half of patients and can progress to petechiae and purpura. Serious signs include delirium, coma, seizures, noncardiogenic pulmonary edema, skin necrosis, and peripheral gangrene. Mortality rates approached 60% in the preantibiotic era and continue to exceed 10–15% in contemporary outbreaks. *Scrub typhus*, caused by *Orientia tsutsugamushi* (a separate genus in the family Rickettsiaceae), is transmitted by larval mites or chiggers and is one of the most common infections in southeastern Asia and the western Pacific. The organism is found in areas of heavy scrub vegetation (e.g., along riverbanks). Patients may have an inoculation eschar and may develop a maculopapular rash. Severe cases progress to pneumonia, meningoencephalitis, DIC, and renal failure. Mortality rates range from 1% to 35%.

If recognized in a timely fashion, rickettsial disease is very responsive to treatment. Doxycycline (100 mg twice daily for 3–14 days) is the treatment of choice for both adults and children. The newer macrolides and chloramphenicol may be suitable alternatives, but mortality rates are higher when a tetracycline-based treatment is not given.

Purpura Fulminans (See also Chaps. 180 and 325) Purpura fulminans is the cutaneous manifestation of DIC and presents as large ecchymotic areas and hemorrhagic bullae. Progression of petechiae to

purpura, ecchymoses, and gangrene is associated with congestive heart failure, septic shock, acute renal failure, acidosis, hypoxia, hypotension, and death. Purpura fulminans has been associated primarily with *N. meningitidis* but, in splenectomized patients, may be associated with *S. pneumoniae*, *H. influenzae*, and *S. aureus*.

Ecthyma Gangrenosum Septic shock caused by *P. aeruginosa* or *Aeromonas hydrophila* can be associated with ecthyma gangrenosum (see Figs. 189-1 and 25e-35): hemorrhagic vesicles surrounded by a rim of erythema with central necrosis and ulceration. These gram-negative bacteremias are most common among patients with neutropenia, extensive burns, and hypogammaglobulinemia.

Other Emergent Infections Associated with Rash *Vibrio vulnificus* and other noncholera *Vibrio* bacteremic infections (Chap. 193) can cause focal skin lesions and overwhelming sepsis in hosts with chronic liver disease, iron storage disorders, diabetes, renal insufficiency, or other immunocompromising conditions. After ingestion of contaminated raw shellfish, typically oysters from the Gulf Coast, there is a sudden onset of malaise, chills, fever, and hypotension. The patient develops bullous or hemorrhagic skin lesions, usually on the lower extremities, and 75% of patients have leg pain. The mortality rate can be as high as 50–60%, particularly when the patient presents with hypotension. Outcomes are improved when patients are treated with tetracycline-containing regimens. Other infections, caused by agents such as *Aeromonas*, *Klebsiella*, and *E. coli*, can cause hemorrhagic bullae and death due to overwhelming sepsis in cirrhotic patients. *Capnocytophaga canimorsus* can cause septic shock in asplenic patients. Infection typically follows a dog bite. Patients present with fever, chills, myalgia, vomiting, diarrhea, dyspnea, confusion, and headache. Findings can include an exanthem or erythema multiforme (see Figs. 70-9 and 25e-25), cyanotic mottling or peripheral cyanosis, petechiae, and ecchymosis. About 30% of patients with this fulminant form die of overwhelming sepsis and DIC, and survivors may require amputation because of gangrene.

Erythroderma TSS (Chaps. 172 and 173) is usually associated with erythroderma. The patient presents with fever, malaise, myalgias, nausea, vomiting, diarrhea, and confusion. There is a sunburn-type rash that may be subtle and patchy but is usually diffuse and is found on the face, trunk, and extremities. Erythroderma, which desquamates after 1–2 weeks, is more common in *Staphylococcus*-associated than in *Streptococcus*-associated TSS. Hypotension develops rapidly—often within hours—after the onset of symptoms. Multiorgan failure occurs. Early renal failure may precede hypotension and distinguishes this syndrome from other septic shock syndromes. There may be no indication of a primary focal infection, although possible cutaneous or mucosal portals of entry for the organism can be ascertained when a careful history is taken. Colonization rather than overt infection of the vagina or a postoperative wound, for example, is typical with staphylococcal TSS, and the mucosal areas appear hyperemic but not infected. Streptococcal TSS is more often associated with skin or soft tissue infection (including necrotizing fasciitis), and patients are more likely to be bacteremic. TSS caused by *Clostridium sordellii* is associated with childbirth or with skin injection of black-tar heroin. The diagnosis of TSS is defined by the clinical criteria of fever, rash, hypotension, and multiorgan involvement. The mortality rate is 5% for menstruation-associated TSS, 10–15% for nonmenstrual TSS, 30–70% for streptococcal TSS, and up to 90% for obstetric *C. sordellii* TSS.

Viral Hemorrhagic Fevers Viral hemorrhagic fevers (Chaps. 233 and 234)



are zoonotic illnesses caused by viruses that reside in either animal reservoirs or arthropod vectors. These diseases occur worldwide and are restricted to areas where the host species live. They are caused by four major groups of viruses: Arenaviridae (e.g., Lassa fever in Africa), Bunyaviridae (e.g., Rift Valley fever in Africa; hantavirus hemorrhagic fever with renal syndrome in Asia; or Crimean-Congo hemorrhagic fever, which has an extensive geographic distribution), Filoviridae (e.g., Ebola and Marburg virus infections in Africa), and Flaviviridae (e.g., yellow fever in Africa and South America and dengue in Asia, Africa, and the Americas). Lassa fever

and Ebola and Marburg virus infections are also transmitted from person to person. The vectors for most viral fevers are found in rural areas; dengue and yellow fever are important exceptions. After a prodrome of fever, myalgias, and malaise, patients develop evidence of vascular damage, petechiae, and local hemorrhage. Shock, multifocal hemorrhaging, and neurologic signs (e.g., seizures or coma) predict a poor prognosis. Dengue (Chap. 233) is the most common arboviral disease worldwide. More than half a million cases of dengue hemorrhagic fever occur each year, with at least 12,000 deaths. Patients have a triad of symptoms: hemorrhagic manifestations, evidence of plasma leakage, and platelet counts of $<100,000/\mu\text{L}$. Mortality rates are 10–20%. If dengue shock syndrome develops, mortality rates can reach 40%. Supportive care to maintain blood pressure and intravascular volume with careful volume-replacement therapy is key to survival. Ribavirin also may be useful against Arenaviridae and Bunyaviridae.

SEPSIS WITH A SOFT TISSUE/MUSCLE PRIMARY FOCUS

See also Chap. 156.

Necrotizing Fasciitis This infection is characterized by extensive necrosis of the subcutaneous tissue and fascia. It may arise at a site of minimal trauma or postoperative incision and may also be associated with recent varicella, childbirth, or muscle strain. The most common causes of necrotizing fasciitis are group A streptococci alone (Chap. 173), the incidence of which has been increasing for the past two decades, and a mixed facultative and anaerobic flora (Chap. 156). Diabetes mellitus, IV drug use, chronic liver or renal disease, and malignancy are associated risk factors. Physical findings are initially minimal compared with the severity of pain and the degree of fever. The examination is often unremarkable except for soft tissue edema and erythema. The infected area is red, hot, shiny, swollen, and exquisitely tender. In untreated infection, the overlying skin develops blue-gray patches after 36 h, and cutaneous bullae and necrosis develop after 3–5 days. Necrotizing fasciitis due to a mixed flora, but not that due to group A streptococci, can be associated with gas production. Without treatment, pain decreases because of thrombosis of the small blood vessels and destruction of the peripheral nerves—an ominous sign. The mortality rate is 15–34% overall, $>70\%$ in association with TSS, and nearly 100% without surgical intervention. Necrotizing fasciitis may also be due to *Clostridium perfringens* (Chap. 179); in this condition, the patient is extremely toxic and the mortality rate is high. Within 48 h, rapid tissue invasion and systemic toxicity associated with hemolysis and death ensue. The distinction between this entity and clostridial myonecrosis is made by muscle biopsy. Necrotizing fasciitis caused by community-acquired MRSA also has been reported.

Clostridial Myonecrosis (See also Chap. 179) Myonecrosis is often associated with trauma or surgery but can develop spontaneously. The incubation period is usually 12–24 h long, and massive necrotizing gangrene develops within hours of onset. Systemic toxicity, shock, and death can occur within 12 h. The patient's pain and toxic appearance are out of proportion to physical findings. On examination, the patient is febrile, apathetic, tachycardic, and tachypneic and may express a feeling of impending doom. Hypotension and renal failure develop later, and hyperalertness is evident preterminally. The skin over the affected area is bronze-brown, mottled, and edematous. Bullous lesions with serosanguineous drainage and a mousy or sweet odor can develop. Crepitus can occur secondary to gas production in muscle tissue. The mortality rate is $>65\%$ for spontaneous myonecrosis, which is often associated with *Clostridium septicum* or *C. tertium* and underlying malignancy. The mortality rates associated with trunk and limb infection are 63% and 12%, respectively, and any delay in surgical treatment increases the risk of death.

NEUROLOGIC INFECTIONS WITH OR WITHOUT SEPTIC SHOCK

Bacterial Meningitis (See also Chap. 164) Bacterial meningitis is one of the most common infectious disease emergencies involving the central nervous system. Although hosts with cell-mediated immune deficiency (including transplant recipients, diabetic patients, elderly patients, and cancer patients receiving certain chemotherapeutic

agents) are at particular risk for *Listeria monocytogenes* meningitis, most cases in adults are due to *S. pneumoniae* (30–60%) and *N. meningitidis* (10–35%). The classic presentation of fever, meningismus, and altered mental status is seen in only one-half to two-thirds of patients. The elderly can present without fever or meningeal signs. Cerebral dysfunction is evidenced by confusion, delirium, and lethargy that can progress to coma. In some cases, the presentation is fulminant, with sepsis and brain edema; papilledema at presentation is unusual and suggests another diagnosis (e.g., an intracranial lesion). Focal signs, including cranial nerve palsies (IV, VI, VII), can be seen in 10–20% of cases; 50–70% of patients have bacteremia. A poor outcome is associated with coma, hypotension, a pneumococcal etiology, respiratory distress, a CSF glucose level of $<0.6\text{ mmol/L}$ ($<<0\text{ mg/dL}$), a CSF protein level of $>2.5\text{ g/L}$, a peripheral white blood cell count of $<5000/\mu\text{L}$, and a serum sodium level of $<135\text{ mmol/L}$. Rapid initiation of treatment is essential; the odds of an unfavorable outcome may increase by 30% for each hour that treatment is delayed. Mortality also increases linearly with age of the patient.

Suppurative Intracranial Infections (See also Chap. 164) In suppurative intracranial infections, rare intracranial lesions present along with sepsis and hemodynamic instability. Rapid recognition of the toxic patient with central neurologic signs is crucial to improvement of the dismal prognosis of these entities. *Subdural empyema* arises from the paranasal sinus in 60–70% of cases. Microaerophilic streptococci and staphylococci are the predominant etiologic organisms. The patient is toxic, with fever, headache, and nuchal rigidity. Of all patients, 75% have focal signs and 6–20% die. Despite improved survival rates, 15–44% of patients are left with permanent neurologic deficits. *Septic cavernous sinus thrombosis* follows a facial or sphenoid sinus infection; 70% of cases are due to staphylococci (including MRSA), and the remainder are due primarily to aerobic or anaerobic streptococci. A unilateral or retroorbital headache progresses to a toxic appearance and fever within days. Three-quarters of patients have unilateral periorbital edema that becomes bilateral and then progresses to ptosis, proptosis, ophthalmoplegia, and papilledema. The mortality rate is as high as 30%. *Septic thrombosis of the superior sagittal sinus* spreads from the ethmoid or maxillary sinuses and is caused by *S. pneumoniae*, other streptococci, and staphylococci. The fulminant course is characterized by headache, nausea, vomiting, rapid progression to confusion and coma, nuchal rigidity, and brainstem signs. If the sinus is totally thrombosed, the mortality rate exceeds 80%.

Brain Abscess (See also Chap. 164) Brain abscess often occurs without systemic signs. Almost half of patients are afebrile, and presentations are more consistent with a space-occupying lesion in the brain; 70% of patients have headache and/or altered mental status, 50% have focal neurologic signs, and 25% have papilledema. Abscesses can present as single or multiple lesions resulting from contiguous foci or hematogenous infection, such as endocarditis. The infection progresses over several days from cerebritis to an abscess with a mature capsule. More than half of infections are polymicrobial, with an etiology consisting of aerobic bacteria (primarily streptococcal species) and anaerobes. Abscesses arising hematogenously are especially apt to rupture into the ventricular space, causing a sudden and severe deterioration in clinical status and a high mortality rate. Otherwise, mortality is low but morbidity is high (30–55%). Patients presenting with stroke and a parameningeal infectious focus, such as sinusitis or otitis, may have a brain abscess, and physicians must maintain a high level of suspicion. Prognosis worsens in patients with a fulminant course, delayed diagnosis, abscess rupture into the ventricles, multiple abscesses, or abnormal neurologic status at presentation.

Cerebral Malaria (See also Chap. 248) This entity should be urgently considered if patients who have recently traveled to areas endemic for malaria present with a febrile illness and lethargy or other neurologic signs. Fulminant malaria is caused by *Plasmodium falciparum* and is associated with temperatures of $>40^\circ\text{C}$ ($>104^\circ\text{F}$), hypotension, jaundice, adult respiratory distress syndrome, and bleeding. By definition, any patient with a change in mental status or repeated

784 seizure in the setting of fulminant malaria has cerebral malaria. In adults, this nonspecific febrile illness progresses to coma over several days; occasionally, coma occurs within hours and death within 24 h. Nuchal rigidity and photophobia are rare. On physical examination, symmetric encephalopathy is typical, and upper motor neuron dysfunction with decorticate and decerebrate posturing can be seen in advanced disease. Unrecognized infection results in a 20–30% mortality rate.

Intracranial and Spinal Epidural Abscesses (See also Chap. 456) Spinal and intracranial epidural abscesses (SEAs and ICEAs) can result in permanent neurologic deficits, sepsis, and death. At-risk patients include those with diabetes mellitus; IV drug use; chronic alcohol abuse; recent spinal trauma, surgery, or epidural anesthesia; and other comorbid conditions, such as HIV infection. Fungal epidural abscess and meningitis can follow epidural or paraspinal glucocorticoid infections. In the United States and Canada, where early treatment of otitis and sinusitis is typical, ICEA is rare but the number of cases of SEA is on the rise. In Africa and areas with limited access to health care, SEAs and ICEAs cause significant morbidity and mortality. ICEAs typically present as fever, mental status changes, and neck pain, while SEAs often present as fever, localized spinal tenderness, and back pain. ICEAs are typically polymicrobial, whereas SEAs are most often due to hematogenous seeding, with staphylococci the most common etiologic agent. Early diagnosis and treatment, which may include surgical drainage, minimize rates of mortality and permanent neurologic sequelae. Outcomes are worse for SEA due to MRSA, infection at a higher vertebral-body level, impaired neurologic status on presentation, and dorsal rather than ventral location of the abscess. Elderly patients and persons with renal failure, malignancy, and other comorbidities also have less favorable outcomes.

Other Focal Syndromes with a Fulminant Course Infection at virtually any primary focus (e.g., osteomyelitis, pneumonia, pyelonephritis, or cholangitis) can result in bacteremia and sepsis. Lemierre's disease—jugular septic thrombophlebitis caused by *Fusobacterium necrophorum*—is associated with metastatic infectious emboli (primarily to the lung) and sepsis, with mortality rates of >15%. TSS has been associated with focal infections such as septic arthritis, peritonitis, sinusitis, and wound infection. Rapid clinical deterioration and death can be associated with destruction of the primary site of infection, as is seen in endocarditis and in infections of the oropharynx (e.g., Ludwig's angina or epiglottitis, in which edema suddenly compromises the airway).

Rhinocerebral Mucormycosis (See also Chap. 242) Individuals with diabetes or immunocompromising conditions are at risk for invasive rhinocerebral mucormycosis. Patients present with low-grade fever, dull sinus pain, diplopia, decreased mental status, decreased ocular motion, chemosis, proptosis, dusky or necrotic nasal turbinates, and necrotic hard-palate lesions that respect the midline. Without rapid recognition and intervention, the process continues on an inexorable invasive course, with high mortality rates.

Acute Bacterial Endocarditis (See also Chap. 155) This entity presents with a much more aggressive course than subacute endocarditis. Bacteria such as *S. aureus*, *S. pneumoniae*, *L. monocytogenes*, *Haemophilus* species, and streptococci of groups A, B, and G attack native valves. Native-valve endocarditis caused by *S. aureus* (including MRSA strains) is increasing, particularly in health care settings. Mortality rates range from 10% to 40%. The host may have comorbid conditions such as underlying malignancy, diabetes mellitus, IV drug use, or alcoholism. The patient presents with fever, fatigue, and malaise <2 weeks after onset of infection. On physical examination, a changing murmur and congestive heart failure may be noted. Hemorrhagic macules on palms or soles (*Janeway lesions*) sometimes develop. Petechiae, Roth's spots, splinter hemorrhages, and

splenomegaly are unusual. Rapid valvular destruction, particularly of the aortic valve, results in pulmonary edema and hypotension. Myocardial abscesses can form, eroding through the septum or into the conduction system and causing life-threatening arrhythmias or high-degree conduction block. Large friable vegetations can result in major arterial emboli, metastatic infection, or tissue infarction. Older patients with *S. aureus* endocarditis are especially likely to present with nonspecific symptoms—a circumstance that delays diagnosis and worsens prognosis. Rapid intervention is crucial for a successful outcome.

Inhalational Anthrax (See also Chap. 261e) Inhalational anthrax, the most severe form of disease caused by *B. anthracis*, had not been reported in the United States for more than 25 years until the use of this organism as an agent of bioterrorism in 2001. Patients presented with malaise, fever, cough, nausea, drenching sweats, shortness of breath, and headache. Rhinorrhea was unusual. All patients had abnormal chest roentgenograms at presentation. Pulmonary infiltrates, mediastinal widening, and pleural effusions were the most common findings. Hemorrhagic meningitis was seen in 38% of these patients. Survival was more likely when antibiotics were given during the prodromal period and when multidrug regimens were used. In the absence of urgent intervention with antimicrobial agents and supportive care, inhalational anthrax progresses rapidly to hypotension, cyanosis, and death.

Avian and Swine Influenza (See also Chap. 224) Human cases of avian influenza have occurred primarily in Southeast Asia, particularly Vietnam (H5N1) and China (H7N9). Avian influenza should be considered in patients with severe respiratory tract illness, particularly if they have been exposed to poultry. Patients present with high fever, an influenza-like illness, and lower respiratory tract symptoms; this illness can progress rapidly to bilateral pneumonia, acute respiratory distress syndrome, multiorgan failure, and death. Early antiviral treatment with neuraminidase inhibitors should be initiated along with aggressive supportive measures. Unlike avian influenza, for which human-to-human transmission has been rare so far and has not been sustained, a novel swine-associated influenza A/H1N1 virus has spread rapidly throughout the world. Patients most at risk of severe disease are children <5 years of age, elderly persons, patients with underlying chronic conditions, and pregnant women. Obesity also has been identified as a risk factor for severe illness.

Hantavirus Pulmonary Syndrome (See also Chap. 233) Hantavirus pulmonary syndrome has been documented in the United States (primarily the southwestern states), Canada, and South America. Most cases occur in rural areas and are associated with exposure to rodents. Patients present with a nonspecific viral prodrome of fever, malaise, myalgias, nausea, vomiting, and dizziness that may progress to pulmonary edema and respiratory failure. Hantavirus pulmonary syndrome causes myocardial depression and increased pulmonary vascular permeability; therefore, careful fluid resuscitation and use of pressor agents are crucial. Aggressive cardiopulmonary support during the first few hours of illness can be life-saving. The early onset of thrombocytopenia may help distinguish this syndrome from other febrile illnesses in an appropriate epidemiologic setting.

CONCLUSION

Acutely ill febrile patients with the syndromes discussed in this chapter require close observation, aggressive supportive measures, and—in most cases—admission to intensive care units. The most important task of the physician is to distinguish these patients from other infected febrile patients whose illness will not progress to fulminant disease. The alert physician must recognize the acute infectious disease emergency and then proceed with appropriate urgency.

148 Immunization Principles and Vaccine Use

Anne Schuchat, Lisa A. Jackson

Few medical interventions of the past century can rival the effect that immunization has had on longevity, economic savings, and quality of life. Seventeen diseases are now preventable through vaccines routinely administered to children and adults in the United States (Table 148-1), and most vaccine-preventable diseases of childhood are at historically low levels (Table 148-2). Health care providers deliver the vast majority of vaccines in the United States in the course of providing routine health services and therefore play an integral role in the nation's public health system.

VACCINE IMPACT

Direct and Indirect Effects Immunizations against specific infectious diseases protect individuals against infection and thereby prevent symptomatic illnesses. Specific vaccines may blunt the severity of clinical illness (e.g., rotavirus vaccines and severe gastroenteritis) or reduce complications (e.g., zoster vaccines and postherpetic neuralgia). Some immunizations also reduce transmission of infectious disease agents from immunized people to others, thereby reducing the impact of infection spread. This indirect impact is known as *herd immunity*. The level of immunization in a population that is required to achieve indirect protection of unimmunized people varies substantially with the specific vaccine.

Since childhood vaccines have become widely available in the United States, major declines in rates of vaccine-preventable diseases among both children and adults have become evident (Table 148-2). For example, vaccination of children <5 years of age against seven types of *Streptococcus pneumoniae* led to a >90% overall reduction in invasive disease caused by those types. A series of childhood vaccines targeting 13 vaccine-preventable diseases in a single birth cohort leads to prevention of 42,000 premature deaths and 20 million illnesses and saves nearly \$70 billion (U.S.).

Control, Elimination, and Eradication of Vaccine-Preventable Diseases

Immunization programs are associated with the goals of controlling,

TABLE 148-1 DISEASES PREVENTABLE WITH VACCINES ROUTINELY ADMINISTERED IN THE UNITED STATES TO CHILDREN AND/OR ADULTS

Condition	Target Population(s) for Routine Use
Pertussis	Children, adolescents, adults
Diphtheria	Children, adolescents, adults
Tetanus	Children, adolescents, adults
Poliomyelitis	Children
Measles	Children
Mumps	Children
Rubella, congenital rubella syndrome	Children
Hepatitis B	Children
<i>Haemophilus influenzae</i> type b infection	Children
Hepatitis A	Children
Influenza	Children, adolescents, adults
Varicella	Children
Invasive pneumococcal disease	Children, older adults
Meningococcal disease	Adolescents
Rotavirus infection	Infants
Human papillomavirus infection, cervical and anogenital cancers	Adolescents and young adults
Zoster	Older adults

TABLE 148-2 DECLINE IN VACCINE-PREVENTABLE DISEASES IN THE UNITED STATES FOLLOWING WIDESPREAD IMPLEMENTATION OF NATIONAL VACCINE RECOMMENDATIONS

Condition	Annual No. of Prevacine Cases (Average)	No. of Cases Reported in 2012 ^a	Reduction (%) in Cases After Widespread Vaccination
Smallpox	29,005	0	100
Diphtheria	21,053	1	≥99
Measles	530,217	55	≥99
Mumps	162,344	229	≥99
Pertussis	200,752	48,277	76
Polio (paralytic)	16,316	0	100
Rubella	47,745	9	>99
Congenital rubella syndrome	152	2	99
Tetanus	580	37	94
<i>Haemophilus influenzae</i> type b infection	20,000	30 ^b	99
Hepatitis A	117,333	2,890 ^c	98
Hepatitis B (acute)	66,232	18,800 ^c	72
Invasive pneumococcal infection: all ages	63,067	31,600 ^d	50
Invasive pneumococcal infection: <5 years of age	16,069	1,800 ^d	89
Varicella	4,085,120	216,511	95

^aExcept for cases of hepatitis A and hepatitis B, for which 2011 figures are shown. ^bAn additional 13 type b infections are estimated to have occurred among 210 reports of *H. influenzae* infection caused by unknown types among children <5 years of age. ^cData are from the CDC's Viral Hepatitis Surveillance, 2011. ^dData are from the CDC's Active Bacterial Core Surveillance 2012 Provisional Report.

Source: Adapted from SW Roush et al: JAMA 298:2155, 2007; and MMWR 62(33): 669, 2013.


eliminating, or eradicating a disease. *Control* of a vaccine-preventable disease reduces poor illness outcomes and often limits the disruptive impacts associated with outbreaks of disease in communities, schools, and institutions. Control programs can also reduce absences from work for ill persons and for parents caring for sick children, decrease absences from school, and limit health care utilization associated with treatment visits.

Elimination of a disease is a more demanding goal than control, usually requiring the reduction to zero of cases in a defined geographic area but sometimes defined as reduction in the indigenous sustained transmission of an infection in a geographic area. As of 2013, the United States had eliminated indigenous transmission of measles, rubella, poliomyelitis, and diphtheria. Importation of pathogens from other parts of the world continues to be important, and public health efforts are intended to react promptly to such cases and to limit forward spread of the infectious agent.

Eradication of a disease is achieved when its elimination can be sustained without ongoing interventions. The only vaccine-preventable disease of humans that has been globally eradicated thus far is smallpox. Although smallpox vaccine is no longer given routinely, the disease has not reemerged naturally because all chains of human transmission were interrupted through earlier vaccination efforts and humans were the only natural reservoir of the virus. Currently, a major health initiative is targeting the global eradication of polio. Sustained transmission of polio has been eliminated from most nations but has never been interrupted in three countries—Afghanistan, Nigeria, and Pakistan—while recent outbreaks in Syria and the Horn of Africa underscore that other countries remain at risk for importation until these reservoirs have been addressed. Detection of a case of disease that has been targeted for eradication or elimination is considered a sentinel event that could permit the infectious agent to become reestablished in the community or region. Therefore, such episodes must be promptly reported to public health authorities.

786 Outbreak Detection and Control Clusters of cases of a vaccine-preventable disease detected in an institution, a medical practice, or a community may signal important changes in the pathogen, vaccine, or environment. Several factors can give rise to increases in vaccine-preventable disease, including (1) low rates of immunization that result in an accumulation of susceptible people (e.g., measles resurgence among vaccination abstainers); (2) changes in the infectious agent that permit it to escape vaccine-induced protection (e.g., non-vaccine-type pneumococci); (3) waning of vaccine-induced immunity (e.g., pertussis among adolescents and adults vaccinated in early childhood); and (4) point-source introductions of large inocula (e.g., food-borne exposure to hepatitis A virus). Reporting episodes of outbreak-prone diseases to public health authorities can facilitate recognition of clusters that require further interventions.

PUBLIC HEALTH REPORTING Recognition of suspected cases of diseases targeted for elimination or eradication—along with other diseases that require urgent public health interventions, such as contact tracing, administration of chemo- or immunoprophylaxis, or epidemiologic investigation for common-source exposure—is typically associated with special reporting requirements. Many diseases against which vaccines are routinely used, including measles, pertussis, *Haemophilus influenzae* type b invasive disease, and varicella, are nationally notifiable. Clinicians and laboratory staff have a responsibility to report some vaccine-preventable disease occurrences to local or state public health authorities according to specific case-definition criteria. All providers should be aware of state or city disease-reporting requirements and the best ways to contact public health authorities. A prompt response to vaccine-preventable disease outbreaks can greatly enhance the effectiveness of control measures.

 **GLOBAL CONSIDERATIONS** Several international health initiatives currently focus on reducing vaccine-preventable diseases in regions throughout the world. These efforts include improving access to new and underutilized vaccines, such as pneumococcal conjugate, rotavirus, human papillomavirus (HPV), and meningococcal A conjugate vaccines. The American Red Cross, the World Health Organization (WHO), the United Nations Foundation, the United Nations Children's Fund (UNICEF), and the Centers for Disease Control and Prevention (CDC) are partners in the Measles & Rubella Initiative, which targets reduction of worldwide measles deaths by 95% from 2000 to 2015. During 2000–2011, global measles mortality rates declined by 71%—i.e., from an estimated 548,000 deaths in 2000 to 158,000 deaths in 2011. Rotary International, UNICEF, the CDC, and the WHO are leading partners in the global eradication of polio, an endeavor that reduced the annual number of paralytic polio cases from 350,000 in 1988 to <250 in 2012. The GAVI Alliance and the Bill and Melinda Gates Foundation have brought substantial momentum to global efforts to reduce vaccine-preventable diseases, expanding on earlier efforts by the WHO, UNICEF, and governments in developed and developing countries.

Enhancing Immunization in Adults Although immunization has become a centerpiece of routine pediatric medical visits, it has not been as well integrated into routine health care visits for adults. This chapter focuses on immunization principles and vaccine use in adults. Accumulating evidence suggests that immunization coverage can be increased through efforts directed at consumer-, provider-, institution-, and system-level factors. The literature suggests that the application of multiple strategies is more effective at raising coverage rates than is the use of any single strategy.

RECOMMENDATIONS FOR ADULT IMMUNIZATIONS The CDC's Advisory Committee on Immunization Practices (ACIP) is the main source of recommendations for administration of vaccines approved by the U.S. Food and Drug Administration (FDA) for use in children and adults in the U.S. civilian population. The ACIP is a federal advisory committee that consists of 15 voting members (experts in fields associated with immunization) appointed by the Secretary of the U.S. Department of Health and Human Services; 8 ex officio members representing federal agencies; and 26 nonvoting representatives of various liaison

organizations, including major medical societies and managed-care organizations. The ACIP recommendations are available at www.cdc.gov/vaccines/hcp/acip-recs/. These recommendations are harmonized to the greatest extent possible with vaccine recommendations made by other organizations, including the American College of Obstetricians and Gynecologists, the American Academy of Family Physicians, and the American College of Physicians.

ADULT IMMUNIZATION SCHEDULES Immunization schedules for adults in the United States are updated annually and can be found online (www.cdc.gov/vaccines/schedules/hcp/adult.html). In January, the schedules are published in *American Family Physician*, *Annals of Internal Medicine*, and *Morbidity and Mortality Weekly Report* (www.cdc.gov/mmwr). The adult immunization schedules for 2013 are summarized in **Fig. 148-1**. Additional information and specifications are contained in the footnotes to these schedules. In the time between annual publications, additions and changes to schedules are published as Notices to Readers in *Morbidity and Mortality Weekly Report*.

IMMUNIZATION PRACTICE STANDARDS

Administering immunizations to adults involves a number of processes, such as deciding whom to vaccinate, assessing vaccine contraindications and precautions, providing vaccine information statements (VISs), ensuring appropriate storage and handling of vaccines, administering vaccines, and maintaining vaccine records. In addition, provider reporting of adverse events that follow vaccination is an essential component of the vaccine safety monitoring system.

Deciding Whom to Vaccinate Every effort should be made to ensure that adults receive all indicated vaccines as expeditiously as possible. When adults present for care, their immunization history should be assessed and recorded, and this information should be used to identify needed vaccinations according to the most current version of the adult immunization schedule. Decision-support tools incorporated into electronic health records can provide prompts for needed vaccinations. Standing orders, which are often used for routinely indicated vaccines (e.g., influenza and pneumococcal vaccines), permit a nurse or another approved licensed practitioner to administer vaccines without a specific physician order, thus lowering barriers to adult immunization.

Assessing Contraindications and Precautions Before vaccination, all patients should be screened for contraindications and precautions. A *contraindication* is a condition that increases the risk of a serious adverse reaction to vaccination. A vaccine should not be administered when a contraindication is documented. For example, a history of an anaphylactic reaction to a dose of vaccine or to a vaccine component is a contraindication for further doses. A *precaution* is a condition that may increase the risk of an adverse event or that may compromise the ability of the vaccine to evoke immunity (e.g., administering measles vaccine to a person who has recently received a blood transfusion and may consequently have transient passive immunity to measles virus). Normally, a vaccine is not administered when a precaution is noted. However, situations may arise when the benefits of vaccination outweigh the estimated risk of an adverse event, and the provider may decide to vaccinate the patient despite the precaution.

In some cases, contraindications and precautions are temporary and may lead to mere deferral of vaccination until a later time. For example, moderate or severe acute illness with or without fever is generally considered a transient precaution to vaccination and results in postponement of vaccine administration until the acute phase has resolved; thus the superimposition of adverse effects of vaccination on the underlying illness and the mistaken attribution of a manifestation of the underlying illness to the vaccine are avoided. Contraindications and precautions to vaccines licensed in the United States for use in civilian adults are summarized in **Table 148-3**. It is important to recognize conditions that are *not* contraindications in order not to miss opportunities for vaccination. For example, in most cases, mild acute illness (with or without fever), a history of a mild to moderate local reaction to a previous dose of the vaccine, and breast-feeding are not contraindications to vaccination.

Recommended Immunizations for Adults by Age

Talk to your healthcare professional about these vaccines:	If you are this age,					
	19–21 years	22–26 years	27–49 years	50–59 years	60–64 years	65+ years
Influenza (Flu) ¹	Get a flu vaccine every year					
Tetanus, diphtheria, pertussis (Td/Tdap) ²	Get a Tdap vaccine once, then a Td booster vaccine every 10 years					
Varicella (Chickenpox)	2 doses					
HPV Vaccine for Women ³	3 doses					
HPV Vaccine for Men ³	3 doses	3 doses				
Zoster (Shingles)					1 dose	
Measles, mumps, rubella (MMR) ⁴	1 or 2 doses					
Pneumococcal (Pneumonia) ⁵		1-3 doses				1 dose
Meningococcal		1 or more doses				
Hepatitis A		2 doses				
Hepatitis B		3 doses				

Boxes this color show that the vaccine is recommended for all adults unless your healthcare professional tells you that you cannot safely receive the vaccine or that you do not need it.

Boxes this color show when the vaccine is recommended for adults with certain risks related to their health, job or lifestyle that put them at higher risk for serious diseases. Talk to your healthcare professional to see if you are at higher risk.

No recommendation

FOOTNOTES:

(Influenza vaccine)¹ There are several flu vaccines available—talk to your healthcare professional about which flu vaccine is right for you.

(Tdap vaccine)² Pregnant women are recommended to get Tdap vaccine with each pregnancy to increase protection for infants who are too young for vaccination but at highest risk for severe illness and death from pertussis (whooping cough).

(HPV vaccine)³ There are two HPV vaccines but only one HPV vaccine (Gardasil®) should be given to men. Gay men or men who have sex with men who are 22 through 26 years old should get HPV vaccine if they haven't already started or completed the series.

(MMR vaccine)⁴ If you were born in 1957 or after, and don't have a record of being vaccinated or having had these infections, talk to your healthcare professional about how many doses you may need.

(Pneumococcal vaccine)⁵ There are two different types of pneumococcal vaccine: PCV13 and PPSV23. Talk with your healthcare professional to find out if one or both pneumococcal vaccines are recommended for you.

If you are traveling outside of the United States, you may need additional vaccines. Ask your healthcare professional which vaccines you may need.

For more information, call toll free 1-800-CDC-INFO (1-800-232-4636) or visit <http://www.cdc.gov/vaccines>

FIGURE 148-1 Recommended adult immunization schedules, United States, 2013. For complete statements by the Advisory Committee on Immunization Practices (ACIP), visit www.cdc.gov/vaccines/hcp/acip-recs/.

(Continues)

Recommended Immunizations for Adults by Medical Condition

Talk to your healthcare professional about these vaccines:	If you have this health condition,								
	Pregnancy	Weakened immune system (not human immuno-deficiency virus [HIV])	HIV Infection		Kidney disease or poor kidney function	Asplenia (if you do not have a spleen or it does not work well)	Heart disease, chronic lung disease, chronic alcoholism	Diabetes (Type 1 and Type 2)	Chronic Liver Disease
			CD4 count is less than 200	CD4 count is 200 or greater					
Influenza (Flu) ¹	Get a flu vaccine every year								
Tetanus, diphtheria, pertussis (Td/Tdap)	1 dose Tdap each pregnancy	Get a Tdap vaccine once, then a Td booster every 10 years							
Varicella (Chickenpox)	SHOULD NOT GET VACCINE			2 doses					
HPV Vaccine for Women ²	3 doses through age 26 yrs								
HPV Vaccine for Men ²	3 doses through age 26 yrs			3 doses through age 21 yrs					
Zoster (Shingles) ³	SHOULD NOT GET VACCINE			1 dose for those 60 yrs and older					
Measles, mumps, rubella (MMR) ⁴	SHOULD NOT GET VACCINE			1 or 2 doses					
Pneumococcal (PPSV23)	1 or 2 doses	1 or 2 doses							
Pneumococcal (PCV13)		1 dose				1 dose			
Meningococcal	1 or more doses				1 or more doses		1 or more doses		
Hepatitis A	2 doses							2 doses	
Hepatitis B	3 doses		3 doses		3 doses		3 doses		

Boxes this color show that the vaccine is recommended for all adults who have not been vaccinated, unless your healthcare professional tells you that you cannot safely receive the vaccine or that you do not need it.

Boxes this color show when the vaccine is recommended for adults with certain risks related to their health, job or lifestyle that put them at higher risk for serious diseases. Talk to your healthcare professional to see if you are at higher risk.

Boxes this color indicate the adult should NOT get this vaccine.

No recommendation

FOOTNOTES:

Last updated August 2013 • CS241388-A

(Influenza vaccine)¹ There are several flu vaccines available—talk to your healthcare professional about which flu vaccine is right for you.

(HPV vaccine)² There are two HPV vaccines but only one HPV vaccine (Gardasil®) should be given to men. Gay men or men who have sex with men who are 22 through 26 years old should get HPV vaccine if they haven't already started or completed the series.

(Zoster)³ You should get zoster vaccine even if you've had shingles before.

(MMR vaccine)⁴ If you were born in 1957 or after, and don't have a record of being vaccinated or having had these infections, talk to your healthcare professional about how many doses you may need.

FIGURE 148-1 (Continued)

TABLE 148-3 CONTRAINDICATIONS AND PRECAUTIONS FOR COMMONLY USED VACCINES IN ADULTS

Vaccine Formulation	Contraindications and Precautions
All vaccines	<p>Contraindication</p> <p>Severe allergic reaction (e.g., anaphylaxis) after a previous vaccine dose or to a vaccine component</p> <p>Precaution</p> <p>Moderate or severe acute illness with or without fever. Defer vaccination until illness resolves.</p>
Td	<p>Precautions</p> <p>GBS within 6 weeks after a previous dose of TT-containing vaccine</p> <p>History of arthus-type hypersensitivity reactions after a previous dose of TD- or DT-containing vaccines (including MCV4). Defer vaccination until at least 10 years have elapsed since the last dose.</p>
Tdap	<p>Contraindication</p> <p>History of encephalopathy (e.g., coma or prolonged seizures) not attributable to another identifiable cause within 7 days of administration of a vaccine with pertussis components, such as DTaP or Tdap</p> <p>Precautions</p> <p>GBS within 6 weeks after a previous dose of TT-containing vaccine</p> <p>Progressive or unstable neurologic disorder, uncontrolled seizures, or progressive encephalopathy. Defer vaccination until a treatment regimen has been established and the condition has stabilized.</p> <p>History of arthus-type hypersensitivity reactions after a previous dose of TT- or DT-containing vaccines (including MCV4). Defer vaccination until at least 10 years have elapsed since the last dose.</p>
HPV	<p>Contraindication</p> <p>History of immediate hypersensitivity to yeast (for Gardasil)</p> <p>Precaution</p> <p>Pregnancy. If a woman is found to be pregnant after initiation of the vaccination series, the remainder of the 3-dose regimen should be delayed until after completion of the pregnancy. If a vaccine dose has been administered during pregnancy, no intervention is needed. Exposure to Gardasil during pregnancy should be reported to Merck (800-986-8999); exposure to Cervarix during pregnancy should be reported to GlaxoSmithKline (888-452-9622).</p>
MMR	<p>Contraindications</p> <p>History of immediate hypersensitivity reaction to gelatin^a or neomycin</p> <p>Pregnancy</p> <p>Known severe immunodeficiency (e.g., hematologic and solid tumors; chemotherapy; congenital immunodeficiency; long-term immunosuppressive therapy; severe immunocompromise due to HIV infection)</p> <p>Precautions</p> <p>Recent receipt (within 11 months) of antibody-containing blood product</p> <p>History of thrombocytopenia or thrombocytopenic purpura</p>
Varicella	<p>Contraindications</p> <p>Pregnancy</p> <p>Known severe immunodeficiency</p> <p>History of immediate hypersensitivity reaction to gelatin^a or neomycin</p> <p>Precaution</p> <p>Recent receipt (within 11 months) of antibody-containing blood product</p>
Influenza, inactivated, injectable	<p>Precautions</p> <p>History of severe allergic reaction (e.g., anaphylaxis) to egg protein^b (note: not a precaution for Flublok recombinant influenza vaccine, which is approved for persons 18–49 years of age and is manufactured without the use of eggs)</p> <p>History of GBS within 6 weeks after a previous influenza vaccine dose</p>
Influenza, live attenuated nasal spray	<p>Contraindications</p> <p>History of severe allergic reaction (e.g., anaphylaxis) to egg protein^b</p> <p>Age ≥50 years</p> <p>Pregnancy</p> <p>Immunosuppression, including that caused by medications or by HIV infection; known severe immunodeficiency (e.g., hematologic and solid tumors; chemotherapy; congenital immunodeficiency; long-term immunosuppressive therapy; severe immunocompromise due to HIV infection)</p> <p>Certain chronic medical conditions, such as diabetes mellitus; chronic pulmonary disease (including asthma); chronic cardiovascular disease (except hypertension); renal, hepatic, neurologic/neuromuscular, hematologic, or metabolic disorders</p> <p>Close contact with severely immunosuppressed persons who require a protected environment, such as isolation in a bone marrow transplantation unit</p> <p>Close contact with persons with lesser degrees of immunosuppression (e.g., persons receiving chemotherapy or radiation therapy who are not being cared for in a protective environment; persons with HIV infection) is <i>not</i> a contraindication or a precaution. Health care personnel in neonatal intensive care units or oncology clinics may receive live attenuated influenza vaccine.</p> <p>Precautions</p> <p>History of GBS within 6 weeks of a previous influenza vaccine dose</p> <p>Receipt of specific antiviral agents (i.e., amantadine, rimantadine, zanamivir, or oseltamivir) with 48 h before vaccination</p>

(Continued)

TABLE 148-3 CONTRAINDICATIONS AND PRECAUTIONS FOR COMMONLY USED VACCINES IN ADULTS (CONTINUED)

Vaccine Formulation	Contraindications and Precautions
Pneumococcal polysaccharide	None, other than those listed for all vaccines
Pneumococcal conjugate	None, other than those listed for all vaccines
Hepatitis A	Precaution Pregnancy
Hepatitis B	Contraindication History of immediate hypersensitivity to yeast
Meningococcal conjugate	Contraindication History of severe allergic reaction to dry natural rubber (latex) (certain vaccine formulations; see text)
Meningococcal polysaccharide	Contraindication History of severe allergic reaction to dry natural rubber (latex)
Zoster	Contraindications Age <50 years Pregnancy Known severe immunodeficiency History of immediate hypersensitivity reaction to gelatin ^a or neomycin Precaution Receipt of specific antiviral agents (i.e., acyclovir, famciclovir, or valacyclovir) within 24 h before vaccination

^aExtreme caution must be exercised in administering MMR, varicella, or zoster vaccine to persons with a history of anaphylactic reaction to gelatin or gelatin-containing products. Before administration, skin testing for sensitivity to gelatin can be considered. However, no specific protocols for this purpose have been published. ^bRecommendations for safely administering influenza vaccine to persons with egg allergies are reported in the annual ACIP recommendations for influenza vaccination (www.cdc.gov/vaccines/hcp/acip-recs/vacc-specific/flu.html).

Abbreviations: DT, diphtheria toxoid; DTaP, diphtheria, tetanus, and pertussis; GBS, Guillain-Barré syndrome; HPV, human papillomavirus; MCV4, quadrivalent meningococcal conjugate vaccine; MMR, measles, mumps, and rubella; Td, tetanus and diphtheria toxoids; Tdap, tetanus and diphtheria toxoids and acellular pertussis; TT, tetanus toxoid.

HISTORY OF IMMEDIATE HYPERSENSITIVITY TO A VACCINE COMPONENT A severe allergic reaction (e.g., anaphylaxis) to a previous dose of a vaccine or to one of its components is a contraindication to vaccination. While most vaccines have many components, substances to which individuals are most likely to have had a severe allergic reaction include egg protein, gelatin, and yeast. In addition, although natural rubber (latex) is not a vaccine component, some vaccines are supplied in vials or syringes that contain natural rubber latex. These vaccines can be identified by the product insert and should not be administered to persons who report a severe (anaphylactic) allergy to latex unless the benefit of vaccination clearly outweighs the risk for a potential allergic reaction. The much more common local or contact hypersensitivity to latex, such as to medical gloves (which contain synthetic latex that is not linked to allergic reactions), is *not* a contraindication to administration of a vaccine supplied in a vial or syringe that contains natural rubber latex. Vaccines routinely indicated for adults that, as of December 2012, were sometimes supplied in a vial or syringe containing natural rubber include Havrix hepatitis A vaccine (syringe); Vaqta hepatitis A vaccine (vial and syringe); Engerix-B hepatitis B vaccine (syringe); Recombivax HB hepatitis B vaccine (vial); Cervarix HPV vaccine (syringe); Fluarix, Fluvirin, Agriflu, and Flucelvax influenza vaccines (syringe); Adacel and Boostrix Tdap (tetanus and diphtheria toxoids and acellular pertussis) vaccines (syringe); Td (tetanus and diphtheria toxoids) vaccines (syringe); Twinrix hepatitis A and B vaccine (syringe); and Menomune meningococcal polysaccharide vaccine (vial).

PREGNANCY Live-virus vaccines are contraindicated during pregnancy because of the theoretical risk that vaccine virus replication will cause congenital infection or have other adverse effects on the fetus. Most live-virus vaccines, including varicella vaccine, are not secreted in breast milk; therefore, breast-feeding is not a contraindication for live-virus or other vaccines. Pregnancy is not a contraindication to administration of inactivated vaccines, but most are avoided during pregnancy because relevant safety data are limited. Two inactivated vaccines, Tdap vaccine and inactivated influenza vaccine, are routinely recommended for pregnant women in the United States. Tdap vaccine is recommended during each pregnancy, regardless of prior vaccination status, in order to prevent pertussis in neonates. Annual influenza vaccination is recommended for all persons 6 months of age and older, regardless of pregnancy status. Some other vaccines, such as

meningococcal vaccines, may be given to pregnant women in certain circumstances.

IMMUNOSUPPRESSION Live-virus vaccines elicit an immune response due to replication of the attenuated (weakened) vaccine virus that is contained by the recipient's immune system. In persons with compromised immune function, enhanced replication of vaccine viruses is possible and could lead to disseminated infection with the vaccine virus. For this reason, live-virus vaccines are contraindicated for persons with severe immunosuppression, the definition of which may vary with the vaccine. Severe immunosuppression may be caused by many disease conditions, including HIV infection and hematologic or generalized malignancy. In some of these conditions, all affected persons are severely immunocompromised. In others (e.g., HIV infection), the degree to which the immune system is compromised depends on the severity of the condition, which in turn depends on the stage of disease or treatment. For example, measles-mumps-rubella (MMR) vaccine may be given to HIV-infected persons who are not severely immunocompromised. Severe immunosuppression may also be due to therapy with immunosuppressive agents, including high-dose glucocorticoids. In this situation, the dose, duration, and route of administration may influence the degree of immunosuppression.

VACCINE INFORMATION STATEMENTS

A VIS is a one-page (two-sided) information sheet produced by the CDC that informs vaccine recipients (or their parents or legal representatives) about the benefits and risks of a vaccine. VISs are mandated by the National Childhood Vaccine Injury Act (NCVIA) of 1986 and—whether the vaccine recipient is a child or an adult—must be provided for any vaccine covered by the Vaccine Injury Compensation Program. As of July 2011, vaccines that are covered by the NCVIA and that are licensed for use in adults include Td, Tdap, hepatitis A, hepatitis B, HPV, trivalent inactivated influenza, trivalent live intranasal influenza, MMR, 13-valent pneumococcal conjugate, meningococcal, polio, and varicella vaccines. When combination vaccines for which no separate VIS exists are given (e.g., hepatitis A and B combination vaccine), all relevant VISs should be provided. VISs also exist for some vaccines not covered by the NCVIA, such as pneumococcal polysaccharide, Japanese encephalitis, rabies, zoster, typhoid, and yellow fever vaccines. The use of these VISs is encouraged but is not mandated.

All current VISs are available on the Internet at two websites: the CDC's Vaccines & Immunizations site (www.cdc.gov/vaccines/hcp/vis/) and the Immunization Action Coalition's site (www.immunize.org/vis/). (The latter site also includes translations of the VISs.) VISs from these sites can be downloaded and printed.

STORAGE AND HANDLING

Injectable vaccines are packaged in multidose vials, single-dose vials, or manufacturer-filled single-dose syringes. The live attenuated nasal-spray influenza vaccine is packaged in single-dose sprayers. Oral typhoid vaccine is packaged in capsules. Some vaccines, such as MMR, varicella, zoster, and meningococcal polysaccharide vaccines, come as lyophilized (freeze-dried) powders that must be reconstituted (i.e., mixed with a liquid diluent) before use. The lyophilized powder and the diluent come in separate vials. Diluents are not interchangeable but rather are specifically formulated for each type of vaccine; only the specific diluent provided by the manufacturer for each type of vaccine should be used. Once lyophilized vaccines have been reconstituted, their shelf-life is limited and they must be stored under appropriate temperature and light conditions. For example, varicella and zoster vaccines must be protected from light and administered within 30 min of reconstitution; MMR vaccine likewise must be protected from light but can be used up to 8 h after reconstitution. Single-dose vials of meningococcal polysaccharide vaccine must be used within 30 min of reconstitution, while multidose vials must be used within 35 days.

Vaccines are stored either at refrigerator temperature (2–8°C) or at freezer temperature (–15°C or colder). In general, inactivated vaccines (e.g., inactivated influenza, pneumococcal polysaccharide, and meningococcal conjugate vaccines) are stored at refrigerator temperature, while vials of lyophilized-powder live-virus vaccines (e.g., varicella, zoster, and MMR vaccines) are stored at freezer temperature. Diluents for lyophilized vaccines may be stored at refrigerator or room temperature. Live attenuated influenza vaccine—a live-virus liquid formulation administered by nasal spray—is stored at refrigerator temperature.

Vaccine storage and handling errors can result in the loss of vaccines worth millions of dollars, and administration of improperly stored vaccines may elicit inadequate immune responses in patients. To improve the standard of vaccine storage and handling practices, the CDC has published detailed guidance (available at www.cdc.gov/vaccines/recs/storage/toolkit/storage-handling-toolkit.pdf). For vaccine storage, the CDC recommends stand-alone units—i.e., self-contained units that either refrigerate or freeze but do not do both—as these units maintain the required temperatures better than combination refrigerator/freezer units. Dormitory-style combined refrigerator/freezer units should never be used for vaccine storage.

The temperature of refrigerators and freezers used for vaccine storage must be monitored and the temperature recorded at least twice each workday. Ideally, continuous thermometers that measure and record temperature all day and all night are used, and minimum and maximum temperatures are read and documented each workday. The CDC recommends the use of calibrated digital thermometers with a probe in a glycol-filled bottle; more detailed information on specifications of storage units and temperature-monitoring devices is provided at the link given above.

ADMINISTRATION OF VACCINES

Most parenteral vaccines recommended for routine administration to adults in the United States are given by either the IM or the SC route; one influenza vaccine formulation approved for use in adults 18–64 years of age is given intradermally. Live-virus vaccines such as varicella, zoster, and MMR are given SC. Most inactivated vaccines are given IM, except for meningococcal polysaccharide vaccine, which is given SC. The 23-valent pneumococcal polysaccharide vaccine may be given either IM or SC, but IM administration is preferred because it is associated with a lower risk of injection-site reactions.

Vaccines given to adults by the SC route are administered with a 5/8-inch needle into the upper outer-triceps area. Vaccines administered to adults by the IM route are injected into the deltoid muscle



FIGURE 148-2 Technique for IM administration of vaccine. (Photo credit: James Gathany, Centers for Disease Control and Prevention; accessible at Public Health Image Library, www.cdc.gov. PHIL ID#9420.)

(Fig. 148-2) with a needle whose length should be selected on the basis of the recipient's sex and weight to ensure adequate penetration into the muscle. Current guidelines indicate that, for men and women weighing <152 lbs (<70 kg), a 1-inch needle is sufficient; for women weighing 152–200 lbs (70–90 kg) and men weighing 152–260 lbs (70–118 kg), a 1- to 1.5-inch needle is needed; and for women weighing >200 lbs (>90 kg) and men weighing >260 lbs (>118 kg), a 1.5-inch needle is required. Additional illustrations of vaccine injection locations and techniques may be found at www.immunize.org/catg.d/p2020a.pdf.

Aspiration, the process of pulling back on the plunger of the syringe after skin penetration but prior to injection, is not necessary because no large blood vessels are present at the recommended vaccine injection sites.

Multiple vaccines can be administered at the same visit; indeed, administration of all needed vaccines at one visit is encouraged. Studies have shown that vaccines are as effective when administered simultaneously as they are individually, and simultaneous administration of multiple vaccines is not associated with an increased risk of adverse effects. If more than one vaccine must be administered in the same limb, the injection sites should be separated by 1–2 inches so that any local reactions can be differentiated. If a vaccine and an immune globulin preparation are administered simultaneously (e.g., Td vaccine and tetanus immune globulin), a separate anatomic site should be used for each injection.

For certain vaccines (e.g., HPV vaccine and hepatitis B vaccine), multiple doses are required for an adequate and persistent antibody response. The recommended vaccination schedule specifies the interval between doses. Many adults who receive the first dose in a multiple-dose vaccine series do not complete the series or do not receive subsequent doses within the recommended interval. For example, at least one-third of adults who receive the first dose of hepatitis B vaccine in the three-dose series do not complete the series. In these circumstances, vaccine efficacy and/or the duration of protection may be compromised. Providers should implement recall systems that will prompt patients to return for subsequent doses in a vaccination series at the appropriate intervals. With the exception of oral typhoid vaccination, an interruption in the schedule does not require restarting of the entire series or the addition of extra doses.

Syncope may follow vaccination, especially in adolescents and young adults. Serious injuries, including skull fracture and cerebral hemorrhage, have occurred. Adolescents and adults should be seated or lying down during vaccination. The majority of reported syncope episodes after vaccination occur within 15 min. The ACIP recommends that vaccine providers strongly consider observing patients, particularly adolescents, with patients seated or lying down for 15 min after vaccination. If syncope develops, patients should be observed until the symptoms resolve.

792 *Anaphylaxis* is a rare complication of vaccination. All facilities providing immunizations should have an emergency kit containing aqueous epinephrine for administration in the event of a systemic anaphylactic reaction.

MAINTENANCE OF VACCINE RECORDS

All vaccines administered should be fully documented in the patient's permanent medical record. Documentation should include the date of administration, the name or common abbreviation of the vaccine, the vaccine lot number and manufacturer, the administration site, the VIS edition, the date the VIS was provided, and the name, address, and title of the person who administered the vaccine. Increasing use of two-dimensional bar codes on vaccine vials and syringes that can be scanned for data entry into compatible electronic medical records and immunization information systems may facilitate more complete and accurate recording of required information.

VACCINE SAFETY MONITORING AND ADVERSE EVENT REPORTING

Prelicensure Evaluations of Vaccine Safety Before vaccines are licensed by the FDA, they are evaluated in clinical trials with volunteers. These trials are conducted in three progressive phases. Phase 1 trials are small, usually involving fewer than 100 volunteers. Their purposes are to provide a basic evaluation of safety and to identify common adverse events. Phase 2 trials, which are larger and may involve several hundred participants, collect additional information on safety and are usually designed to evaluate immunogenicity as well. Data gained from phase 2 trials can be used to determine the composition of the vaccine, the number of doses required, and a profile of common adverse events. Vaccines that appear promising are evaluated in phase 3 trials, which typically involve several hundred to several thousand volunteers and are generally designed to demonstrate vaccine efficacy and provide additional information on vaccine safety.

Postlicensure Monitoring of Vaccine Safety After licensure, a vaccine's safety is assessed by several mechanisms. The NCVIA of 1986 requires health care providers to report certain adverse events that follow vaccination. As a mechanism for that reporting, the Vaccine Adverse Event Reporting System (VAERS) was established in 1990 and is jointly managed by the CDC and the FDA. This safety surveillance system collects reports of adverse events associated with vaccines currently licensed in the United States. *Adverse events* are defined as untoward events that occur after immunization and that might be caused by the vaccine product or vaccination process. While the VAERS was established in response to the NCVIA, any adverse event following vaccination—whether in a child or an adult, and whether or not it is believed to have actually been caused by vaccination—may be reported through the VAERS. The adverse events that health care providers are required to report are listed in the reportable-events table on the VAERS website at vaers.hhs.gov/reportable.htm. Approximately 30,000 VAERS reports are filed annually, with ~13% reporting serious events resulting in hospitalization, life-threatening illness, disability, or death.

Anyone can file a VAERS report, including health care providers, manufacturers, and vaccine recipients or their parents or guardians. VAERS reports may be submitted online (vaers.hhs.gov/esub/step1) or by completing a paper form requested by email (info@vaers.org), phone (800-822-7967), or fax (877-721-0366). The VAERS form asks for the following information: the type of vaccine received; the timing of vaccination; the time of onset of the adverse event; and the recipient's current illnesses or medications, history of adverse events following vaccination, and demographic characteristics (e.g., age and sex). This information is entered into a database. The individual who reported the adverse event then receives a confirmation letter by mail with a VAERS identification number that can be used if additional information is submitted later. In selected cases of serious adverse reaction, the patient's recovery status may be followed up at 60 days and 1 year after vaccination. The FDA and the CDC have access to VAERS data and use this information to monitor vaccine safety and conduct research studies. VAERS data (minus personal information) are also available to the public.

While the VAERS provides useful information on vaccine safety, this passive reporting system has important limitations. One is that

events following vaccination are merely reported; the system cannot assess whether a given type of event occurs more often than expected after vaccination. A second is that event reporting is incomplete and is biased toward events that are believed to be more likely to be due to vaccination and that occur relatively soon after vaccination. To obtain more systematic information on adverse events occurring in both vaccinated and unvaccinated persons, the Vaccine Safety Datalink project was initiated in 1991. Directed by the CDC, this project includes nine managed-care organizations in the United States; member databases include information on immunizations, medical conditions, demographics, laboratory results, and medication prescriptions. The Department of Defense oversees a similar system monitoring the safety of immunizations among active-duty military personnel. In addition, postlicensure evaluations of vaccine safety may be conducted by the vaccine manufacturer. In fact, such evaluations are often required by the FDA as a condition of vaccine licensure.

CONSUMER ACCESS TO AND DEMAND FOR IMMUNIZATION

By removing barriers to the consumer or patient, providers and health care institutions can improve vaccine use. Financial barriers have traditionally been important constraints, particularly among uninsured adults. Even for insured adults, out-of-pocket costs associated with newer, more expensive adult vaccines (e.g., zoster vaccine) are an obstacle to be overcome. After influenza vaccine was included by Medicare for all beneficiaries in 1993, coverage among persons ≥ 65 years of age doubled (from ~30% in 1989 to >60% in 1997). Other strategies that enhance patients' access to vaccination include extended office hours (e.g., evening and weekend hours) and scheduled vaccination-only clinics where waiting times are reduced. Provision of vaccines outside the "medical home" (e.g., through occupational clinics, universities, pharmacies, and retail settings) can expand access for adults who do not make medical visits frequently. Increasing proportions of adults are being vaccinated in these settings.

Health promotion efforts aimed at increasing the demand for immunization are common. Direct-to-consumer advertising by pharmaceutical companies has been used for some newer adolescent and adult vaccines. Efforts to raise consumer demand for vaccines have not increased immunization rates unless implemented in conjunction with other strategies that target strengthening of provider practices or reduction of consumer barriers. Attitudes and beliefs related to vaccination can be considerable impediments to consumer demand. Many adults view vaccines as important for children but are less familiar with vaccinations targeting disease prevention in adults. Several vaccines are recommended for adults with certain medical risk factors, but self-identification as a high-risk individual is relatively rare. Communication research suggests that many adults with chronic diseases may be more motivated to receive a vaccine by a desire to protect their family members rather than to reduce their own risk. Some vaccines are explicitly recommended for persons at relatively low risk of serious complications, with the goal of reducing the risk of transmission to higher-risk contacts. For example, for protection of newborns, vaccinations against influenza and pertussis are recommended for pregnant women and for others who will be around the infant.

STRATEGIES FOR PROVIDERS AND HEALTH CARE FACILITIES

Recommendation from the Provider Health care providers can have great influence on patients with regard to immunization. A recommendation from a doctor or nurse carries more weight than do recommendations from professional societies or endorsements by celebrities. Providers should be well informed about vaccine risks and benefits so that they can address patients' common concerns. The CDC, the American College of Physicians, and the American Academy of Family Physicians review and update the schedule for adult immunization on an annual basis and also have developed educational materials to facilitate provider-patient discussions about vaccination (www.cdc.gov/vaccines/hcp.htm).

System Supports Medical offices can incorporate a variety of methods to ensure that providers consistently offer specific immunizations to patients with indications for specific vaccines. Decision-support tools

have been incorporated into some electronic health records to alert the provider when specific vaccines are indicated. Manual or automated reminders and standing orders have been discussed (see “Deciding Whom to Vaccinate,” above) and have consistently improved vaccination coverage in both office and hospital settings. Most clinicians’ estimates of their own performance diverge from objective measurements of their patients’ immunization coverage; quantitative assessment and feedback have been shown in pediatric and adolescent practices to increase immunization performance significantly. Some health plans have instituted incentives for providers with high rates of immunization coverage. Specialty providers, including obstetrician–gynecologists, may be the only providers serving some high-risk patients with indications for selected vaccines (e.g., Tdap, influenza, or pneumococcal polysaccharide vaccine).

Immunization Requirements Vaccination against selected communicable diseases is required for attendance at many universities and colleges as well as for service in the U.S. military or in some occupational settings (e.g., child care, laboratory, veterinary, and health care). Immunizations are recommended and sometimes required for travel to certain countries (Chap. 149).

Vaccination of Health Care Staff A particular area of focus for medical settings is vaccination of health care workers, including those with and without direct patient-care responsibilities. The Joint Commission (which accredits health care organizations), the CDC’s Healthcare Infection Control Practices Advisory Committee, and the ACIP all recommend influenza vaccination of all health care personnel; recommendations also focus on requiring documentation of declination for providers who do not accept annual influenza vaccination. As part of their participation in the Centers for Medicare and Medicaid Services’ Hospital Inpatient Quality Reporting program, acute-care hospitals are required to report the proportion of their health care personnel who have received seasonal influenza vaccine. Some institutions and jurisdictions have added mandates on influenza vaccination of health care workers and have expanded on earlier requirements related to vaccination or proof of immunity for hepatitis B, measles, mumps, rubella, and varicella.

VACCINATION IN NONMEDICAL SETTINGS

Receipt of vaccination in medical offices is most frequent among young children and adults ≥ 65 years of age. People in these age groups make more office visits and are more likely to receive care in a consistent “medical home” than are older children, adolescents, and nonelderly adults. Vaccination outside the medical home can expand access to those whose health care visits are limited and reduce the burden on busy clinical practices. In some locations, financial constraints related to inventory and storage requirements have led providers to stock few or no vaccines. Outside private office and hospital settings, vaccination may also occur at health department venues, workplaces, retail sites (including pharmacies and supermarkets), and schools or colleges.

When vaccines are given in nonmedical settings, it remains important for standards of immunization practice to be followed. Consumers should be provided with information on how to report adverse events (e.g., via provision of a VIS), and procedures should ensure that documentation of vaccine administration is forwarded to the primary care provider and the state or city public health immunization registry. Detailed documentation may be required for employment, school attendance, and travel. Personalized health records can help consumers keep track of their immunizations, and some occupational health clinics have incorporated automated immunization reports that help employees stay up-to-date with recommended vaccinations. Some pharmacy chain establishments are using automated systems to report immunization information to the state or local immunization information system.

PERFORMANCE MONITORING

Tracking of immunization coverage at national, state, institution, and practice levels can yield feedback to practitioners and programs

and facilitate quality improvement. Healthcare Effectiveness Data and Information Set (HEDIS) measures related to adult immunization facilitate comparison of health plans. The CDC’s National Immunization Survey and National Health Interview Survey provide selected information on immunization coverage among adults and track progress toward achievement of Healthy People 2020 targets for immunization coverage. Influenza and pneumococcal vaccine coverage rates have been higher among persons ≥ 65 years of age (60–70%) than among high-risk 18- to 64-year-olds. Figures on state-specific immunization coverage with pneumococcal polysaccharide and influenza vaccines (as measured through the CDC’s Behavioral Risk Factor Surveillance System) reveal substantial geographic variation in coverage. There are persistent disparities in adult immunization coverage rates between whites and racial and ethnic minorities. In contrast, racial and economic disparities in immunization of young children have been dramatically reduced during the past 20 years. Much of this progress is attributed to the Vaccines for Children Program, which since 1994 has entitled uninsured children to receive free vaccines.

FUTURE TRENDS

Although most vaccines developed in the twentieth century targeted common acute infectious diseases of childhood, more recently developed vaccines prevent chronic conditions prevalent among adults. Hepatitis B vaccine prevents hepatitis B–related cirrhosis and hepatocellular carcinoma, zoster vaccine prevents shingles and postherpetic neuralgia, and HPV vaccine prevents some types of cervical cancer, genital warts, and anogenital cancers and may also prevent some oropharyngeal cancers (although this outcome was not studied in prelicensure randomized controlled trials). New targets of vaccine development and research may further broaden the definition of vaccine-preventable disease. Research is ongoing on vaccines to prevent insulin-dependent diabetes mellitus, nicotine addiction, and Alzheimer’s disease. Expanding strategies for vaccine development are incorporating molecular approaches such as DNA, vector, and peptide vaccines. New technologies, such as the use of transdermal and other needle-less routes of administration, are being applied to vaccine delivery.

149 Health Recommendations for International Travel

Jay S. Keystone, Phyllis E. Kozarsky

According to the World Tourism Organization, international tourist arrivals grew dramatically from 25 million in 1950 to >1 billion in 2012. Not only are more people traveling; travelers are seeking more exotic and remote destinations. Travel from industrialized to developing regions has been increasing, with Asia and the Pacific, Africa, and the Middle East now emerging destinations. Figure 149-1 summarizes the monthly incidence of health problems during travel in developing countries. Studies continue to show that 50–75% of short-term travelers to the tropics or subtropics report some health impairment. Most of these health problems are minor: only 5% require medical attention, and $<1\%$ require hospitalization. Although infectious agents contribute substantially to morbidity among travelers, these pathogens account for only $\sim 1\%$ of deaths in this population. Cardiovascular disease and injuries are the most frequent causes of death among travelers from the United States, accounting for 49% and 22% of deaths, respectively. Age-specific rates of death due to cardiovascular disease are similar among travelers and nontravelers. In contrast, rates of death due to injury (the majority from

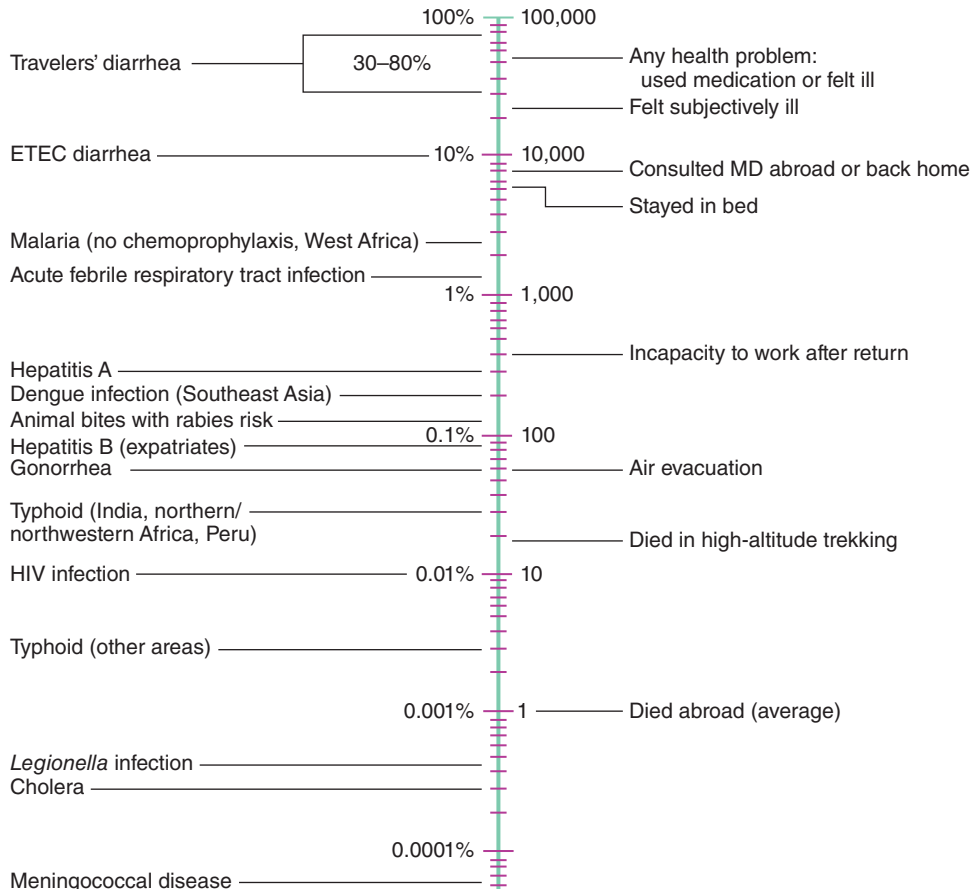


FIGURE 149-1 Monthly incidence rates of health problems during stays in developing countries. ETEC, enterotoxigenic *Escherichia coli*. (From R Steffen et al: *Int J Antimicrob Agents* 21:89, 2003.)

motor vehicle, drowning, or aircraft accidents) are several times higher among travelers. Motor vehicle accidents account for >40% of travelers' deaths that are not due to cardiovascular disease or preexisting illness.

GENERAL ADVICE

Health maintenance recommendations are based not only on the traveler's destination but also on assessment of risk, which is determined by such variables as health status, specific itinerary, purpose of travel, season, and lifestyle during travel. Detailed information regarding country-specific risks and recommendations may be obtained from the Centers for Disease Control and Prevention (CDC) publication *Health Information for International Travel* (available at www.cdc.gov/travel).

Fitness for travel is an issue of growing concern in view of the increased numbers of elderly and chronically ill individuals journeying to exotic destinations (see "Travel and Special Hosts," below). Since most commercial aircraft are pressurized to 2500 m (8000 ft) above sea level (corresponding to a $P_{a_{O_2}}$ of ~55 mmHg), individuals with serious cardiopulmonary problems or anemia should be evaluated before travel. In addition, those who have recently had surgery, a myocardial infarction, a cerebrovascular accident, or a deep-vein thrombosis may be at high risk for adverse events during flight. A summary of current recommendations regarding fitness to fly has been published by the Aerospace Medical Association Air Transport Medicine Committee (www.asma.org/publications/medical-publications-for-airline-travel). A pretravel health assessment may be advisable for individuals considering particularly adventurous recreational activities, such as mountain climbing and scuba diving.

IMMUNIZATIONS FOR TRAVEL

Immunizations for travel fall into three broad categories: *routine* (childhood/adult boosters that are necessary regardless of travel), *required* (immunizations that are mandated by international regulations for entry into certain areas or for border crossings), and

recommended (immunizations that are desirable because of travel-related risks). Required and recommended vaccines commonly given to travelers are listed in [Table 149-1](#).

Routine Immunizations • DIPHTHERIA, TETANUS, AND POLIO Diphtheria ([Chap. 175](#)) continues to be a problem worldwide. Large outbreaks have occurred in countries that do not have rigorous vaccination programs or that have reduced their public vaccination programs. Serologic surveys show that tetanus ([Chap. 177](#)) antibodies are lacking in many North Americans, especially in women over the age of 50. The risk of polio ([Chap. 228](#)) to the international traveler is extremely low, and wild-type poliovirus has been eradicated from the Western Hemisphere and Europe. However, studies in the United States suggest that 12% of adult travelers are unprotected against at least one poliovirus serogroup. In addition, challenges continue to be faced by polio eradication programs. Foreign travel offers an ideal opportunity to have these immunizations updated. With the recent increase in pertussis among adults, the diphtheria–tetanus–acellular pertussis (Tdap) combination is now recommended for adults as a once-only replacement for the 10-year tetanus–diphtheria (Td) booster.

MEASLES Measles (rubeola) continues to be a major cause of morbidity and death in the developing world ([Chap. 229](#)). Several outbreaks of measles in the United States and Canada have been linked to imported cases, especially from Europe, where large outbreaks have occurred recently. The group at highest risk consists of persons born after 1956 and vaccinated before 1980, in many of whom primary vaccination failed. The measles–mumps–rubella (MMR) vaccine is typically used; its coverage of rubella also addresses a growing concern in some areas of Eastern Europe and Asia.

INFLUENZA Influenza ([Chap. 224](#))—possibly the most common vaccine-preventable infection in travelers—occurs year-round in the tropics and during the summer months in the Southern Hemisphere (coinciding with the winter months in the Northern Hemisphere). One prospective study showed that influenza developed in 1% of travelers to Southeast Asia per month of stay. Annual vaccination should be considered for all travelers who do not have a contraindication. Travel-related influenza continues to occur during summer months in Alaska and the Northwest Territories of Canada among cruise-ship passengers and staff. The speed of global spread of the pandemic H1N1 virus once again illustrates why influenza immunization is so important for travelers.

PNEUMOCOCCAL INFECTION Regardless of travel, pneumococcal vaccine ([Chap. 171](#)) should be administered routinely to persons over the age of 65 and to persons at high risk of serious infection, including those with chronic heart, lung, or kidney disease; those who have been splenectomized; and those who have sickle cell disease.

Required Immunizations • YELLOW FEVER Documentation of vaccination against yellow fever ([Chap. 233](#)) may be required or recommended as a condition for entry into or passage through countries of sub-Saharan Africa and equatorial South America, where the disease is endemic or epidemic, or (by the International Health Regulations) for entry into countries at risk of having the infection introduced. This

TABLE 149-1 VACCINES COMMONLY USED FOR TRAVEL IN ADULTS

Vaccine	Primary Series	Booster Interval
Cholera (Dukoral [®] inactivated whole-cell recombinant subunit; available in Canada and Europe)	1 dose	2 years
Hepatitis A (Havrix), 1440 enzyme immunoassay U/mL	2 doses, 6–12 months apart, IM	None required
Hepatitis A (VAQTA, AVAXIM, EPAXAL)	2 doses, 6–18 months apart, IM	None required
Hepatitis A/B combined (Twinrix)	3 doses at 0, 1, and 6 months 0, 7, and 21–30 days plus booster at 1 year, IM	None required <i>except</i> 12 months (once only, for accelerated schedule)
Hepatitis B (Engerix B): accelerated schedule	3 doses at 0, 1, and 2 months 0, 7, and 21 days plus booster at 1 year, IM	12 months, once only
Hepatitis B (Engerix B or Recombivax): standard schedule	3 doses at 0, 1, and 6 months, IM	None required
Immune globulin (hepatitis A prevention)	1 dose IM	Intervals of 3–5 months, depending on initial dose
Japanese encephalitis (Ixiaro)	2 doses at 0 and 28 days, IM	>1 year after primary series (optimal booster schedule not yet determined)
Meningococcus, quadrivalent (Menomune [polysaccharide], Menactra, Menveo [conjugate])	1 dose (Menactra/Menveo IM; Menomune SC)	>3 years (optimal booster schedule not yet determined)
Rabies human diploid cell vaccine (Imovax), rabies vaccine absorbed (RVA), or purified chick embryo cell vaccine (RabAvert)	3 doses at 0, 7, and 21 or 28 days, IM	None required except with exposure
Typhoid Ty21a, oral live attenuated (Vivotif)	1 capsule every other day × 4 doses	5 years
Typhoid Vi capsular polysaccharide, injectable (Typhim Vi)	1 dose IM	2 years
Yellow fever	1 dose SC	10 years

[®]Cross-protects against enterotoxigenic *Escherichia coli* and provides 30–50% protection against travelers' diarrhea.

vaccine is given only by state-authorized yellow fever centers, and its administration must be documented on an official International Certificate of Vaccination. A registry of U.S. clinics that provide the vaccine is available from the CDC (www.cdc.gov/travel). Recent data suggest that fewer than 50% of travelers entering areas endemic for yellow fever are immunized; this lack of coverage is a serious problem, as 13 countries in Central and South America and 30 countries in Africa harbor the illness. Severe adverse events associated with this vaccine have recently increased in incidence. First-time vaccine recipients may present with a syndrome characterized as either neurotropic (1 case per 125,000 doses) or viscerotropic (overall, 1 case per 250,000 doses; among persons 60–69 years of age, 1 case per 100,000 doses; and among persons ≥ 70 years of age, 1 case per 40,000 doses). Immunosuppression and thymic disease increase the risk of these adverse events (www.cdc.gov/vaccines/hcp/vis/vis-statements/yf.pdf).

MENINGOCOCCAL MENINGITIS Protection against meningitis with one of the quadrivalent (preferably conjugate) vaccines is required for entry into Saudi Arabia during the Hajj (**Chap. 180**).

INFLUENZA Both seasonal and pandemic H1N1 vaccines (the latter, where available) were required for entry into Saudi Arabia during the Hajj in 2013.

Recommended Immunizations • HEPATITIS A AND B Hepatitis A (**Chap. 360**) is one of the most common vaccine-preventable infections of travelers. The risk is six times greater for travelers who stray from the usual tourist routes. The mortality rate for hepatitis A increases with age, reaching almost 2% among individuals over age 50. Of the four hepatitis A vaccines currently available in North America (two in the United States), all are interchangeable and have an efficacy of >95%. Hepatitis A vaccine is currently given to all children in the United States. Since the most frequently identified risk factor for hepatitis A in the United States is international travel, and since morbidity and mortality increase with age, it seems appropriate that all adults be immune prior to travel.

Long-stay overseas workers appear to be at considerable risk for hepatitis B infection (**Chap. 360**). The recommendation that all travelers be immunized against hepatitis B before departure is supported by two studies showing that 17% of the assessed travelers who received health care abroad had some type of injection; according to the World Health Organization, nonsterile equipment is used for up to 75% of all injections given in parts of the developing world. A 3-week accelerated schedule of the combined hepatitis A and B vaccine has been approved in the United States. Although no data are available on the specific risk of infection with hepatitis B virus among U.S. travelers, ~240 million people in the world have chronic infection. All children and adolescents in the United States are immunized against this illness. Hepatitis B vaccination should be considered for all travelers.

TYPHOID FEVER Most cases of typhoid fever in North America are due to travel, with ~300 cases seen per year in the United States. The attack rate for typhoid fever (**Chap. 190**) is 1 case per 30,000 travelers per month of travel to the developing world. However, attack rates in India, Senegal, and North Africa are tenfold higher; rates are especially high among travelers to relatively remote destinations and among immigrants and their families who have returned to their homelands to visit friends or relatives (VFRs). Between 1999 and 2006 in the United States, 66% of imported cases involved the latter group. Unfortunately, data show that the causative organism has become increasingly resistant to fluoroquinolone antibiotics (especially in those cases acquired on the Indian subcontinent). Both of the available vaccines—one oral (live) and the other injectable (polysaccharide)—have efficacy rates of ~70%. In some countries, a combined hepatitis A/typhoid vaccine is available.

MENINGOCOCCAL MENINGITIS Although the risk of meningococcal disease among travelers has not been quantified, it is likely to be higher among travelers who live with poor indigenous populations in overcrowded conditions (**Chap. 180**). Because of its enhanced ability to prevent nasal carriage (compared with the older polysaccharide vaccine), a quadrivalent conjugate vaccine is the product of choice (regardless of age) for immunization of persons traveling to sub-Saharan Africa during the dry season or to areas of the world where there are epidemics. The vaccine, which protects against serogroups A, C, Y, and W-135, has an efficacy rate of >90%.

JAPANESE ENCEPHALITIS The risk of Japanese encephalitis (**Chap. 233**), an infection transmitted by mosquitoes in rural Asia and Southeast Asia, can be as high as ~1 case per 5000 travelers per month of stay in an endemic area. Most infections are asymptomatic, with a very small proportion of infected persons becoming ill. However, among those who do become ill, severe neurologic sequelae are common. Most symptomatic infections among U.S. residents have involved military personnel or their families. The vaccine efficacy rate is >90%. The vaccine is recommended for persons staying >1 month in rural endemic areas or for shorter periods if their activities (e.g., camping, bicycling, hiking) in these areas will increase exposure risk.

CHOLERA The risk of cholera (**Chap. 193**) is extremely low, with ~1 case per 500,000 journeys to endemic areas. Cholera vaccine, not currently available in the United States, was rarely recommended but was considered for aid and health care workers in refugee camps or in disaster-stricken/war-torn areas. A more effective oral cholera vaccine is available in other countries.

796 RABIES Domestic animals, primarily dogs, are the major transmitters of rabies in developing countries (**Chap. 232**). Several studies have shown that the risk of rabies posed by a dog bite in an endemic area translates into 1–3.6 cases per 1000 travelers per month of stay. Countries where canine rabies is highly endemic include Mexico, the Philippines, Sri Lanka, India, Thailand, China, and Vietnam. The two vaccines available in the United States provide >90% protection. Rabies vaccine is recommended for long-stay travelers, particularly children (who tend to play with animals and may not report bites), and for persons who may be occupationally exposed to rabies in endemic areas; however, in a large-scale study, almost 50% of potential exposures occurred within the first month of travel. Even after receipt of a preexposure rabies vaccine series, two postexposure doses are required. Travelers who have had the preexposure series do not require rabies immune globulin (which is often unavailable in developing countries) if they are exposed to the disease.

PREVENTION OF MALARIA AND OTHER INSECT-BORNE DISEASES

It is estimated that more than 30,000 American and European travelers develop malaria each year (**Chap. 248**). The risk to travelers is highest in Oceania and sub-Saharan Africa (estimated at 1:5 and 1:50 per month of stay, respectively, among persons not using chemoprophylaxis); intermediate in malarious areas on the Indian subcontinent and in Southeast Asia (1:250–1:1000 per month); and low in South and Central America (1:2500–1:10,000 per month). Of the 1925 cases of malaria reported in 2011 in the United States (the highest figure in 40 years), 90% of those due to *Plasmodium falciparum* occurred in travelers returning or emigrating from Africa and Oceania. VFRs are at the highest risk of acquiring malaria and may die of the disease if their immunity has waned after living outside an endemic area for a number of years. According to data from the CDC, VFRs accounted for 59% of severe malaria cases in the United States in 2011. With the worldwide increase in chloroquine- and multidrug-resistant falciparum malaria, decisions about chemoprophylaxis have become more difficult. The case-fatality rate for falciparum malaria in the United States is 4%; however, *in only one-third of patients who die is the diagnosis of malaria considered before death.*

Several studies indicate that fewer than 50% of travelers adhere to basic recommendations for malaria prevention. Keys to the prevention of malaria include both personal protection measures against mosquito bites (especially between dusk and dawn) and malaria chemoprophylaxis. The former measures entail the use of DEET-containing insect repellents, permethrin-impregnated bed nets and clothing, screened sleeping accommodations, and protective clothing. Thus, in regions where infections such as malaria are transmitted, DEET products (25–50%) are recommended, even for children and infants at birth. Studies suggest that concentrations of DEET above ~50% do not offer a marked increase in protection time against mosquitoes. The CDC also recommends picaridin, oil of lemon eucalyptus (PMD, parmenthane-3,8-diol), and IR3535 (3-[N-butyl-N-acetyl]-aminopropionic acid, ethyl ester). In general, higher concentrations of any active ingredient provide a longer duration of protection. Personal protection measures also help prevent other insect-transmitted illnesses, such as dengue fever (**Chap. 233**). Over the past decade, the incidence of dengue has markedly increased, particularly in the Caribbean region, Latin America, Southeast Asia, and (more recently) Africa. Chikungunya, another mosquito-borne infection that clinically resembles dengue fever but with arthralgia and arthritis instead of myalgia, has recently crossed to the Western Hemisphere; many thousands of cases are now occurring in the Caribbean. Both dengue and chikungunya viruses are transmitted by an urban-dwelling mosquito that bites primarily at dawn and dusk.

Table 149-2 lists the currently recommended drugs of choice for prophylaxis of malaria, by destination.

PREVENTION OF GASTROINTESTINAL ILLNESS

Diarrhea, the leading cause of illness in travelers (**Chap. 160**), is usually a short-lived, self-limited condition. However, 40% of affected individuals need to alter their scheduled activities, and another 20% are confined to bed. The most important determinant of risk is the destination. Incidence rates per 2-week stay have been reported to be

TABLE 149-2 MALARIA CHEMOSUPPRESSIVE REGIMENS, ACCORDING TO GEOGRAPHIC AREA^a

Geographic Area	Drug of Choice ^b	Alternatives
Central America (north of Panama), Iraq, Turkey, northern Argentina, and Paraguay	Chloroquine	Atovaquone-proguanil ^c Doxycycline Mefloquine Primaquine
South America (but not northern Argentina or Paraguay, where chloroquine may be used); Central America (only Panama east of the Canal); Asia (including Southeast Asia); Africa; and Oceania	Doxycycline Atovaquone-proguanil ^c Mefloquine	
Thai-Myanmar and Thai-Cambodian borders, central Vietnam	Atovaquone-proguanil ^c Doxycycline	

^aSee CDC's *Health Information for International Travel 2014* (www.cdc.gov/travel). ^bIn all areas where chloroquine can still be used, the other drugs listed may be used as alternatives.

^cMalarone.

Note: See also **Chap. 248**.

as low as 8% in industrialized countries and as high as 55% in parts of Africa, Central and South America, and Southeast Asia. Infants and young adults are at particularly high risk for gastrointestinal illness and for complications such as dehydration. Recent reviews suggest that there is little correlation between dietary indiscretions and the occurrence of travelers' diarrhea. Earlier studies of U.S. students in Mexico showed that eating meals in restaurants and cafeterias or consuming food from street vendors was associated with increased risk. For further discussion, see "Precautions," below.

Etiology (See also **Table 160-3**) The most frequently identified pathogens causing travelers' diarrhea are enterotoxigenic and enteroaggregative *Escherichia coli* (**Chap. 186**), although in some parts of the world (notably northern Africa and Southeast Asia) *Campylobacter* infections (**Chap. 192**) appear to predominate. Other common causative organisms include *Salmonella* (**Chap. 190**), *Shigella* (**Chap. 191**), rotavirus (**Chap. 227**), and norovirus (**Chap. 227**). The latter virus has caused numerous outbreaks on cruise ships. Except for giardiasis (**Chap. 254**), parasitic infections are uncommon causes of travelers' diarrhea in short-term travelers. A growing problem for travelers is the development of antibiotic resistance among many bacterial pathogens. Examples include strains of *Campylobacter* resistant to quinolones and strains of *E. coli*, *Shigella*, and *Salmonella* resistant to trimethoprim-sulfamethoxazole. *E. coli* O157 is very rarely a cause of travelers' diarrhea.

Precautions Some experts think that it is not only *what* travelers eat but also where they eat that puts them at risk of illness. Food sold by street vendors can carry a high risk, and restaurant hygiene can be a major problem over which the traveler has no control. In addition to discretion in choosing the source of food and water, general precautions include eating foods piping hot; avoiding foods that are raw or poorly cooked; and drinking only boiled or commercially bottled beverages, particularly those that are carbonated. Heating kills diarrhea-causing organisms, whereas freezing does not; therefore, ice cubes made from unpurified water should be avoided. In spite of these recommendations, the literature has repeatedly documented dietary indiscretions by 98% of travelers within the first 72 h after arrival at their destination. The maxim "Boil it, cook it, peel it, or forget it!" is easy to remember but apparently difficult to follow.

Self-Treatment (See also **Table 160-5**) As travelers' diarrhea often occurs despite rigorous food and water precautions, travelers should carry medications for self-treatment. An antibiotic is useful in reducing the frequency of bowel movements and the duration of illness in moderate to severe diarrhea. The standard regimen is a 3-day course of a quinolone taken twice daily (or, in the case of some newer

formulations, once daily). However, studies have shown that one double dose of a quinolone may be equally effective. For diarrhea acquired in areas such as Thailand, where >90% of *Campylobacter* infections are quinolone resistant, azithromycin may be a better alternative. Rifaximin, a poorly absorbed rifampin derivative, is highly effective against noninvasive bacterial pathogens such as enterotoxigenic and enteroaggregative *E. coli*. The current approach to self-treatment of travelers' diarrhea for the typical short-term traveler is to carry three once-daily doses of an antibiotic and to use as many doses as necessary to resolve the illness. If neither high fever nor blood in the stool accompanies the diarrhea, loperamide should be taken in combination with the antibiotic; studies have shown that this combination is more effective than an antibiotic alone and does not prolong illness.

Prophylaxis Prophylaxis of travelers' diarrhea with bismuth subsalicylate is widely used but only ~60% effective. For certain individuals (e.g., athletes, persons with a repeated history of travelers' diarrhea, and persons with chronic diseases), a single daily dose of a quinolone, azithromycin, or rifaximin during travel of <1 month's duration is 75–90% efficacious in preventing travelers' diarrhea. Probiotics have been only ~20% effective as prophylaxis. In Europe and Canada, an oral subunit cholera vaccine that cross-protects against enterotoxigenic *E. coli* (Dukoral) has been shown to provide 30–50% protection against travelers' diarrhea.

Illness After Return Although extremely common, acute travelers' diarrhea is usually self-limited or amenable to antibiotic therapy. Persistent bowel problems after the traveler returns home have a less well-defined etiology and may require medical attention from a specialist. Infectious agents (e.g., *Giardia lamblia*, *Cyclospora cayotensis*, *Entamoeba histolytica*) appear to be responsible for only a small proportion of cases with persistent bowel symptoms. By far the most common causes of persistent diarrhea after travel are postinfectious sequelae such as lactose intolerance and irritable bowel syndrome. A meta-analysis showed that postinfectious irritable bowel syndrome lasting months to years may occur in as many as 4–13% of cases. When no infectious etiology can be identified, a trial of metronidazole therapy for presumed giardiasis, a strict lactose-free diet for 1 week, or a several-week trial of high-dose hydrophilic mucilloid (plus an osmotic laxative such as lactulose or PEG 3350 for persons with alternating diarrhea and constipation) relieves the symptoms of many patients.

PREVENTION OF OTHER TRAVEL-RELATED PROBLEMS

Travelers are at high risk for *sexually transmitted diseases* (Chap. 163). Surveys have shown that large numbers of travelers engage in casual sex, and there is a reluctance to use condoms consistently. An increasing number of travelers are being diagnosed with illnesses such as *schistosomiasis* (Chap. 259), *dengue* (Chap. 233), *chikungunya* (Chap. 233), and *tick-borne rickettsial disease* (Chap. 211). Travelers should be cautioned to avoid bathing, swimming, or wading in freshwater lakes, streams, or rivers in parts of northeastern South America, the Caribbean, Africa, and Southeast Asia. Insect repellents are important for prevention not only of malaria but also of other vector-borne diseases. Prevention of *travel-associated injury* depends mostly on common-sense precautions. Riding on motorcycles (especially without helmets) and in overcrowded public vehicles is not recommended; in developing countries, individuals should *never* travel by road in rural areas after dark. Of persons who die during travel, fewer than 1% die of infection, whereas 40% die in motor vehicle accidents. Excessive alcohol use has been a significant factor in motor vehicle accidents, drownings, assaults, and injuries. Travelers are cautioned to avoid walking barefoot because of the risk of hookworm and *Strongyloides* infections (Chap. 257) and snakebites (Chap. 474).

THE TRAVELER'S MEDICAL KIT

A traveler's medical kit is strongly advisable. The contents may vary widely, depending on the itinerary, duration of stay, style of travel, and local medical facilities. While many medications are available abroad (often over the counter), directions for their use may be nonexistent or in a foreign language, or a product may be outdated

or counterfeit. For example, a multicountry study in Southeast Asia showed that a mean of 53% (range, 21–92%) of antimalarial products were counterfeit or contained inadequate amounts of active drug. The sale and marketing of such medications constitute a growing industry. In the medical kit, the short-term traveler should consider carrying an analgesic; an antiarrheal agent and an antibiotic for self-treatment of travelers' diarrhea; antihistamines; a laxative; oral rehydration salts; a sunscreen with broad-spectrum protection (UVA and UVB, with the latter at a level of at least 30 SPF); a DEET-containing or equivalent insect repellent for the skin; an insecticide for clothing (permethrin); and, if necessary, an antimalarial drug. To these medications, the long-stay traveler might add a broad-spectrum general-purpose antibiotic (levofloxacin or azithromycin), an antibacterial eye and skin ointment, and a topical antifungal cream. Regardless of the duration of travel, a first-aid kit containing such items as scissors, tweezers, and bandages should be considered. A practical approach to self-treatment of infections in the long-stay traveler who carries a once-daily dose of antibiotics (e.g., levofloxacin) is to use 3 tablets "below the waist" (bowel and bladder infections) and 6 tablets "above the waist" (skin and respiratory infections).

TRAVEL AND SPECIAL HOSTS

PREGNANCY AND TRAVEL

(See also Chap. 8) A woman's medical history and itinerary, the quality of medical care at her destinations, and her degree of flexibility determine whether travel is wise during pregnancy. According to the American College of Obstetrics and Gynecology, the safest part of pregnancy in which to travel is between 18 and 24 weeks, when there is the least danger of spontaneous abortion or premature labor. Some obstetricians prefer that women stay within a few hundred miles of home after the 28th week of pregnancy in case problems arise. In general, however, healthy women may be advised that it is acceptable to travel.

Relative contraindications to international travel during pregnancy include a history of miscarriage, premature labor, incompetent cervix, or toxemia. General medical problems such as diabetes, heart failure, severe anemia, or a history of thromboembolic disease also should prompt the pregnant woman to postpone her travels. Finally, regions in which the pregnant woman and her fetus may be at excessive risk (e.g., those at high altitudes, those where live-virus vaccines are required, and those where multidrug-resistant malaria is endemic) are not ideal destinations during any trimester.

Malaria Malaria during pregnancy carries a significant risk of morbidity and death. Levels of parasitemia are highest and failure to clear the parasites after treatment is most frequent among primigravidae. Severe disease, with complications such as cerebral malaria, massive hemolysis, and renal failure, is especially likely in pregnancy. Fetal sequelae include spontaneous abortion, stillbirth, preterm delivery, and congenital infection. Chloroquine and mefloquine are considered to be safe in all trimesters.

Enteric Infections Pregnant travelers must be extremely cautious regarding their food and beverage intake. Dehydration due to travelers' diarrhea can lead to inadequate placental blood flow. Infections such as toxoplasmosis, hepatitis E, and listeriosis also can cause serious sequelae in pregnancy.

The mainstay of therapy for travelers' diarrhea is rehydration. Loperamide may be used if necessary. For self-treatment, azithromycin may be the best option. Although quinolones are increasingly being used safely during pregnancy and rifaximin is poorly absorbed from the gastrointestinal tract, these drugs are not approved for this indication.

Because of the serious problems encountered when infants are given local foods and beverages, women are strongly encouraged to breast-feed when traveling with a neonate. A nursing mother with travelers' diarrhea should not stop breast-feeding but should increase her fluid intake.

Air Travel and High-Altitude Destinations Commercial air travel is not a risk to the healthy pregnant woman or to the fetus. The higher radiation levels reported at altitudes of >10,500 m (>35,000 ft) should pose

798 no problem for the healthy pregnant traveler. Since each airline has a policy regarding pregnancy and flying, it is best to check with the specific carrier when booking reservations. Domestic air travel is usually permitted until the 36th week, whereas international air travel is generally curtailed after the 32nd week.

There are no known risks for pregnant women who travel to high-altitude destinations and stay for short periods. However, there are likewise no data on the safety of pregnant women at altitudes of >4500 m (15,000 ft).

THE HIV-INFECTED TRAVELER

(See also Chap. 226) The HIV-infected traveler is at special risk of serious infections due to a number of pathogens that may be more prevalent at travel destinations than at home. However, the degree of risk depends primarily on the state of the immune system at the time of travel. For persons whose CD4+ T cell counts are normal or >500/ μ L, data suggest no greater risk during travel than for persons without HIV infection. Individuals with AIDS (CD4+ T cell counts of <200/ μ L) and others who are symptomatic need special counseling and should visit a travel medicine practitioner before departure, especially when traveling to the developing world.

Several countries routinely deny entry to HIV-positive individuals for prolonged stay, even though these restrictions do not appear to decrease rates of transmission of the virus. In general, HIV testing is required for individuals who wish to stay abroad >3 months or who intend to work or study abroad. Some countries will accept an HIV serologic test done within 6 months of departure, whereas others will not accept a blood test done at any time in the traveler's home country. Border officials often have the authority to make inquiries of individuals entering a country and to check the medications they are carrying. If antiretroviral drugs are identified, the person may be barred from entering the country. Information on testing requirements for specific countries is available from consular offices but is subject to frequent change.

Immunizations All of the HIV-infected traveler's routine immunizations should be up to date (Chap. 148). The response to immunization may be impaired at CD4+ T cell counts of <200/ μ L and in some cases at even higher counts. Thus HIV-infected persons should be vaccinated as early as possible to ensure adequate immune responses. For patients receiving antiretroviral therapy, at least 3 months must elapse before regenerated CD4+ T cells can be considered fully functional; therefore, vaccination of these patients should be delayed. However, when the risk of illness is high or the sequelae of illness are serious, immunization is recommended. In certain circumstances, it may be prudent to check the adequacy of the serum antibody response before departure.

Because of the increased risk of infections due to *Streptococcus pneumoniae* and other bacterial pathogens that cause pneumonia after influenza, the conjugate pneumococcal vaccine (Pneumovax 13) followed by the 23-valent polysaccharide vaccine (Pneumovax) as well as influenza vaccine should be administered. The estimated rates of response to influenza vaccine are >80% among persons with asymptomatic HIV infection and <50% among those with AIDS.

In general, live attenuated vaccines are contraindicated for persons with immune dysfunction. Because measles (rubeola) can be a severe or lethal infection in HIV-positive patients, these patients should receive the measles vaccine (or the combination MMR vaccine) unless the CD4+ T cell count is <200/ μ L. Between 18% and 58% of symptomatic HIV-infected vaccinees develop adequate measles antibody titers, and 50–100% of asymptomatic HIV-infected persons seroconvert.

It is recommended that the live yellow fever vaccine not be given to HIV-infected travelers. Although the potential adverse effects of a live vaccine in an HIV-infected individual are always a consideration, there appear to have been no reported cases of illness in those who have inadvertently received this vaccine. Nonetheless, if the CD4+ T cell count is <200/ μ L, an alternative itinerary that poses no risk of exposure to yellow fever is recommended. If the traveler is passing through or traveling to an area where the vaccine is required but the disease risk is low, a physician's waiver should be issued.

A transient increase in HIV viremia (lasting days to weeks) has been demonstrated in HIV-infected individuals after immunization against influenza, pneumococcal infection, and tetanus (Chap. 226). At this point, however, no evidence indicates that this transient increase is detrimental.

Gastrointestinal Illness Decreased levels of gastric acid, abnormal gastrointestinal mucosal immunity, other complications of HIV infection, and medications taken by HIV-infected patients make travelers' diarrhea especially problematic in these individuals. Travelers' diarrhea is likely to occur more frequently, to be more severe, to be accompanied by bacteremia, and to be more difficult to treat. *Cryptosporidium*, *Isospora belli*, and *Microsporidium* infections, although uncommon, are associated with increased morbidity and mortality rates in AIDS patients.

The HIV-infected traveler must be careful to consume only appropriately prepared foods and beverages and may benefit from antibiotic prophylaxis for travelers' diarrhea. Sulfonamides (as used to prevent pneumocystosis) are ineffective because of widespread resistance.

Other Travel-Related Infections Data are lacking on the severity of many vector-borne diseases in HIV-infected individuals. Malaria is especially severe in asplenic persons and in those with AIDS. The HIV load doubles during malaria, with subsidence in ~8–9 weeks; the significance of this increase in viral load is unknown.

Visceral leishmaniasis (Chap. 251) has been reported in numerous HIV-infected travelers. Diagnosis may be difficult, given that splenomegaly and hyperglobulinemia are often lacking and serologic results are frequently negative. Sandfly bites may be prevented by evening use of insect repellents.

Certain respiratory illnesses, such as histoplasmosis and coccidioidomycosis, cause greater morbidity and mortality among patients with AIDS. Although tuberculosis is common among HIV-infected persons (especially in developing countries), its acquisition by the short-term HIV-infected traveler has not been reported as a major problem. From a prospective study, it is estimated that for travelers not engaged in health care the risk of tuberculosis infection is ~3% per year of travel.

Medications Adverse events due to medications and drug interactions are common and raise complex issues for HIV-infected persons. Rates of cutaneous reaction (e.g., increased cutaneous sensitivity to sulfonamides) are unusually high among patients with AIDS. Since zidovudine is metabolized by hepatic glucuronidation, inhibitors of this process may elevate serum levels of the drug. Concomitant administration of the antimalarial drug mefloquine and the antiretroviral agent ritonavir may result in decreased plasma levels of ritonavir; mefloquine may also interact with many of the other protease inhibitors. In contrast, no significant influence of concomitant mefloquine administration on plasma levels of indinavir or nelfinavir was detected in two HIV-infected travelers. Serum levels of mefloquine may be lowered with the use of efavirenz or nevirapine. There are also potential interactions between atovaquone-proguanil (Malarone) and many of the protease inhibitors as well as between Malarone and the nonnucleoside reverse transcriptase inhibitors (NNRTIs). Because of the increase in antiretroviral agents and the lack of accumulated data on their interactions with antimalarial agents, decisions about malaria chemoprophylaxis continue to be difficult; with a short duration of travel, an interaction may be inconsequential. However, doxycycline appears to have no clinically significant interactions with either the protease inhibitors or the NNRTIs. With regard to malaria treatment, a great hypothetical concern is that the antimalarial drugs lumefantrine (combined with artemisinin in Coartem) and halofantrine may interact with HIV protease inhibitors and NNRTIs since drugs in the latter two categories are known to be potent inhibitors of cytochrome P450. In keeping current with antiretroviral drug interactions, a website from the University of Liverpool (www.hiv-druginteractions.org) is helpful.

CHRONIC ILLNESS, DISABILITY, AND TRAVEL

Chronic health problems need not prevent travel, but special measures can make the journey safer and more comfortable.

Heart Disease Cardiovascular events are the main cause of deaths among travelers and of in-flight emergencies on commercial aircraft. Extra supplies of all medications should be kept in carry-on luggage, along with a copy of a recent electrocardiogram and the name and telephone number of the traveler's physician at home. Pacemakers are not affected by airport security devices, although electronic telephone checks of pacemaker function cannot be transmitted by international satellites. Travelers with electronic defibrillators should carry a note to that effect and ask for hand screening. A traveler may benefit from supplemental oxygen; since oxygen delivery systems are not standard, supplementary oxygen should be ordered by the traveler's physician well before flight time. Travelers may benefit from aisle seating and should walk, perform stretching and flexing exercises, consider wearing support hose, and remain hydrated during the flight to prevent venous thrombosis and pulmonary embolism.

Chronic Lung Disease Chronic obstructive pulmonary disease is one of the most common diagnoses in patients who require emergency-department evaluation for symptoms occurring during airline flights. The best predictor of the development of in-flight problems is the sea-level PaO_2 . A PaO_2 of at least 72 mmHg corresponds to an in-flight arterial PaO_2 of ~55 mmHg when the cabin is pressurized to 2500 m (8000 ft). If the traveler's baseline PaO_2 is <72 mmHg, the provision of supplemental oxygen should be considered. Contraindications to flight include active bronchospasm, lower respiratory infection, lower-limb deep-vein phlebitis, pulmonary hypertension, and recent thoracic surgery (within the preceding 3 weeks) or pneumothorax. Decreased outdoor activity at the destination should be considered if air pollution is excessive.

Diabetes Mellitus Alterations in glucose control and changes in insulin requirements are common problems among patients with diabetes who travel. Changes in time zone, in the amount and timing of food intake, and in physical activity demand vigilant assessment of metabolic control. Because of the risk of foot ulcers, travelers should wear closed footwear that has been proven to be comfortable. The traveler with diabetes should pack medication (including a bottle of regular insulin for emergencies), insulin syringes and needles, equipment and supplies for glucose monitoring, and snacks in carry-on luggage. Insulin is stable for ~3 months at room temperature but should be kept as cool as possible. The name and telephone number of the home physician and a card and bracelet listing the patient's medical problems and the type and dose of insulin used should accompany the traveler. In order to facilitate international border crossings, travelers should carry a physician's letter authorizing the carriage of needles and syringes. In traveling eastward (e.g., from the United States to Europe), the morning insulin dose on arrival may need to be decreased. The blood glucose can then be checked during the day to determine whether additional insulin is required. For flights westward, with lengthening of the day, an additional dose of regular insulin may be required.

Other Special Groups Other groups for whom special travel measures are encouraged include patients undergoing dialysis, those with transplants, and those with other disabilities. Up to 13% of travelers have some disability, but few advocacy groups and tour companies dedicate themselves to this growing population. Medication interactions are a source of serious concern for these travelers, and appropriate medical information should be carried, along with the home physician's name and telephone number. Some travelers taking glucocorticoids carry stress doses in case they become ill. Immunization of these immunocompromised travelers may result in less than adequate protection. Thus the traveler and the physician must carefully consider which destinations are appropriate.

TRAVEL HEALTH INSURANCE

Today, more elderly or chronically ill individuals travel and more of these individuals journey to remote locations and enjoy adventurous activities. Illness or injury abroad is not uncommon and is best considered before the journey. Persons who develop health problems abroad may incur enormous out-of-pocket expenses. Thus prospective travelers

should consider purchasing additional travel health insurance and should check with their health insurance company regarding whether they have coverage for illness or injury overseas. Unfortunately, many insurance companies will not cover pre-existing illness if it is the reason for trip cancellation or illness abroad. Most countries do not accept routine health insurance from other countries unless there is a special traveler supplement. In most circumstances, travelers are asked to pay in cash for services rendered on an emergency basis, whether in a physician's office, in an emergency or urgent care center, or even in a hospital. There are several types of travel insurance. It is wise to purchase *trip cancellation insurance*, especially, for example, if the traveler has an underlying chronic illness and may need to cancel a trip due to an exacerbation of disease. *Travel health insurance* will cover expenses in the event that medical care abroad is needed. *Evacuation insurance* will cover medical evacuation, usually to a medical center in another location where it is deemed that the care is similar to that available in the traveler's home country. The cost of medical evacuation can easily exceed \$100,000 US. There are a number of travel insurance providers, and it is very important to read the fine print carefully and to determine exactly what each company provides, thereby ensuring an appropriate fit for the individual's particular circumstances. The U.S. Department of State website lists travel health insurance companies (http://travel.state.gov/travel/tips/emergencies/emergencies_5981.html).

MEDICAL TOURISM

Travel for the purpose of obtaining health care abroad has received a great deal of attention in the medical literature and the media. According to the annual U.S. Department of Commerce In-Flight Survey, there were ~500,000 overseas trips during 2006 in which health treatment was at least one purpose of travel. Lower cost is usually cited as the motivation for this type of tourism, and an entire industry has flourished as a result of this phenomenon. However, the quality of facilities, assistance services, and care is neither uniform nor regulated; thus, in most instances, responsibility for assessing the suitability of an individual program or facility lies solely with the traveler. Persons considering this option must recognize that they are almost always at a disadvantage when being treated in a foreign country, particularly if there are complications. Concerns to be addressed include the quality of the health care facility and its staff; language and cultural differences that may impede accurate interpretation of both verbal and nonverbal communication; religious and ethical differences that may be encountered over issues such as efforts to preserve life and limb or the provision of care for the terminally ill; lack of familiarity with the local medical system; limited access of the care provider to the patient's medical history; the use of unfamiliar drugs and medicines; the relative difficulty of arranging follow-up care back in the United States; and the possibility that such follow-up care may be fraught with problems should there be complications. If serious issues arise, legal recourse may be difficult or impossible. Patients planning to travel abroad to obtain health care, particularly when surgery is involved, should be immunized for hepatitis B and should consider having baseline hepatitis B and C and HIV tests preoperatively. Prevalence rates of hepatitis B and C and HIV infection vary considerably around the world and are generally higher in developing regions than in the United States and Western Europe. The latest information available on the safety of the blood supply outside the United States is the World Health Organization's Global Database on Blood Safety based on data from 2011 (www.who.int/bloodsafety/global_database/en). Persons researching the accreditation status of overseas facilities should note that, although these facilities may be part of a chain, they are surveyed and accredited individually. Accreditation resources include (1) the Joint Commission International (www.jointcommissioninternational.org), (2) the Australian Council for Healthcare Standards International (www.achs.org.au/achs-international/), and (3) the Canadian Council on Health Services (www.cchsa.ca). The American Medical Association also offers guidelines for medical tourism (www.ama-assn.org/ama/pub/upload/mm/31/medicaltourism.pdf).

The most common medical problems encountered by travelers after their return home are diarrhea, fever, respiratory illnesses, and skin diseases (Fig. 149-2). Frequently ignored problems are fatigue and emotional stress, especially in long-stay travelers. The approach to diagnosis requires some knowledge of geographic medicine, in

particular the epidemiology and clinical presentation of infectious disorders. A geographic history should focus on the traveler's exact itinerary, including dates of arrival and departure; exposure history (food indiscretions, drinking-water sources, freshwater contact, sexual activity, animal contact, insect bites); location and style of travel (urban vs. rural, first-class hotel accommodation vs. camping); immunization history; and use of antimalarial chemosuppression. Recently,

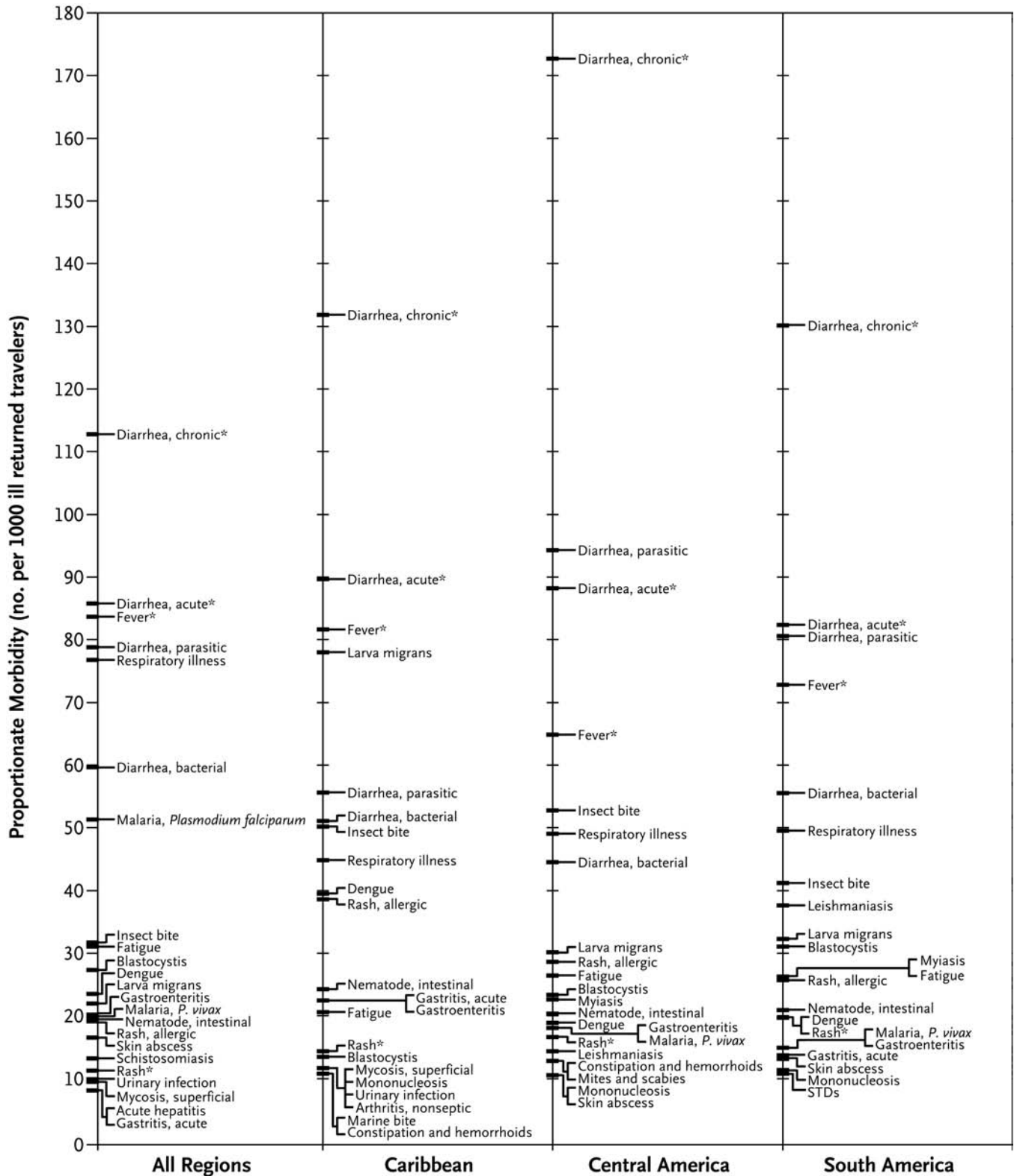


FIGURE 149-2 Proportionate morbidity among ill travelers returning from the developing world, according to region of travel. The proportions (not incidence rates) are shown for each of the top 22 specific diagnoses among all ill returned travelers within each region. STDs, sexually transmitted diseases. Asterisks indicate syndromic diagnoses for which specific etiologies could not be assigned. (Reprinted from DO Freedman et al: *N Engl J Med* 354:119, 2006.)

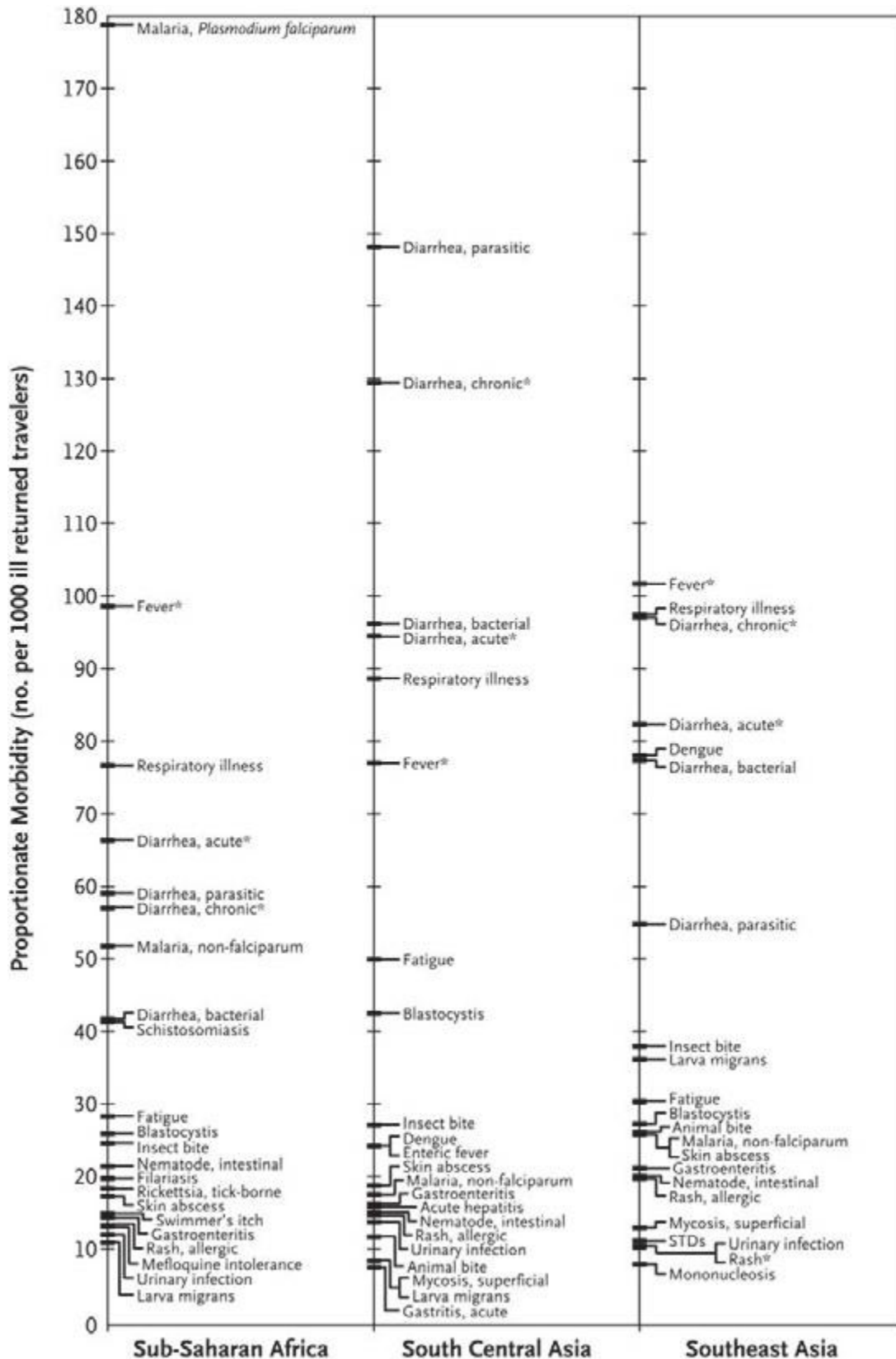


FIGURE 149-2 (Continued)

some travelers who have been hospitalized abroad have been shown on return to be colonized with multidrug-resistant bacteria such as Enterobacteriaceae producing extended-spectrum β -lactamases and bacteria producing NDM-1 (New Delhi metallo- β -lactamase 1).

DIARRHEA

See "Prevention of Gastrointestinal Illness," above.

FEVER

Fever in a traveler who has returned from a malarious area should be considered a medical emergency because death from *P. falciparum* malaria can follow an illness of only several days' duration. Although "fever from the tropics" does not always have a tropical cause, malaria should be the first diagnosis considered. The risk of *P. falciparum* malaria is highest among travelers returning from Africa or Oceania

TABLE 149-3 ETIOLOGY AND GEOGRAPHIC DISTRIBUTION OF SYSTEMIC FEBRILE ILLNESS IN RETURNED TRAVELERS N = 3907

Etiology	Percentage of Cases ^a					
	Caribbean	Central America	South America	Sub-Saharan Africa	South-Central Asia	Southeast Asia
Malaria	<1	13	13	62	14	13
Dengue	23	12	14	<1	14	32
Mononucleosis	7	7	8	1	2	3
<i>Rickettsia</i>	0	0	0	6	1	2
<i>Salmonella</i>	2	3	2	<1	14	3

^aBold type is for emphasis only.

Source: Revised from Table 2 in DO Freedman et al: *N Engl J Med* 354:119, 2006.

and among those who become symptomatic within the first 2 months after return. Other important causes of fever after travel include viral hepatitis (A and E), typhoid and paratyphoid fever, bacterial enteritis, arboviral infections (e.g., dengue fever), rickettsial infections (including tick typhus, scrub typhus, and Q fever), and—in rare instances—leptospirosis, acute HIV infection, and amebic liver abscess. A cooperative study by GeoSentinel (an emerging infectious disease surveillance group established by the CDC and the International Society of Travel Medicine) showed that, among 3907 febrile returned travelers, malaria was acquired most often in Africa, dengue in Southeast Asia and the Caribbean, typhoid fever in southern Asia, and rickettsial infections (tick typhus) in South Africa (Table 149-3). Outbreaks of dengue, previously considered to be very rare in Africa, have been documented recently in Angola, Kenya, and Tanzania. However, in at least 25% of cases, no etiology of the fever can be found and it resolves spontaneously. Clinicians should keep in mind that no present-day antimalarial agent guarantees protection from malaria and that some immunizations (notably, that against typhoid fever) are only partially protective.

When no specific diagnosis is forthcoming, the following investigations, where applicable, are suggested: complete blood count, liver function tests, thick/thin blood films or rapid diagnostic testing for malaria (repeated several times if necessary), urinalysis, urine and blood cultures (repeated once), chest x-ray, and collection of an acute-phase serum sample to be held for subsequent examination along with a paired convalescent-phase serum sample.

SKIN DISEASES

Pyodermas, sunburn, insect bites, skin ulcers, and cutaneous larva migrans are the most common skin conditions affecting travelers after their return home. In those with persistent skin ulcers, a diagnosis of cutaneous leishmaniasis, mycobacterial infection, or fungal infection should be considered. Careful, complete inspection of the skin is important in detecting the rickettsial eschar in a febrile patient or the central breathing hole in a “boil” due to myiasis.

EMERGING INFECTIOUS DISEASES

In recent years, travel and commerce have fostered the worldwide spread of HIV infection, led to the reemergence of cholera as a global health threat, and created considerable fear about the possible spread of novel respiratory diseases, including those caused by influenza viruses (H5N1, H1N1, and H7N9). For travelers, there are more common, everyday concerns. One of the largest outbreaks of dengue fever ever documented is now raging in Latin America and Southeast Asia; chikungunya virus has spread rapidly from Africa to southern Asia, southern Europe, and, for the first time in the Western Hemisphere, the Caribbean; schistosomiasis is being described in previously unaffected lakes in Africa; and antibiotic-resistant strains of sexually transmitted and enteric pathogens are emerging at an alarming rate in the developing world. In addition, concerns have been raised about the potential for bioterrorism involving not only standard strains of unusual agents but mutant strains as well.

CONCLUSIONS

The growth of global travel and migration now demand that the clinician become as familiar as possible with travel medicine. Practitioners may choose either to refer their patients to a travel clinic before departure or to acquire knowledge that enables them to provide pre-travel counseling and to prescribe appropriate vaccinations and chemoprophylaxis. It is equally important for physicians seeing ill returned travelers to be familiar with common post-travel syndromes and diseases, particularly those that may have been acquired in the developing world, and to identify other physicians who can assist with com-

plex post-travel illnesses. The CDC publishes a biennial text, *Health Information for International Travel* (accessed through their website at www.cdc.gov/travel) that provides pretravel health recommendations. The International Society of Travel Medicine (www.istm.org) publishes a list of travel clinics, and the American Society of Tropical Medicine and Hygiene (www.astmh.org) publishes a list of clinical tropical medicine specialists.

As Nobel Laureate Dr. Joshua Lederberg pointed out, “The microbe that felled one child in a distant continent yesterday can reach yours today and seed a global pandemic tomorrow.” The vigilant clinician understands that the importance of a thorough travel history cannot be overemphasized.

150e Laboratory Diagnosis of Infectious Diseases

Alexander J. McAdam, Andrew B. Onderdonk

The laboratory diagnosis of infection requires the demonstration—either direct or indirect—of viral, bacterial, fungal, or parasitic agents in tissues, fluids, or excreta of the host. Clinical microbiology laboratories are responsible for processing these specimens and also for determining the antibiotic susceptibility of bacterial and fungal pathogens. Traditionally, detection of pathogenic agents has relied largely on either the microscopic visualization of pathogens in clinical material or the growth of microorganisms in the laboratory. Identification generally is based on phenotypic characteristics such as fermentation profiles for bacteria, cytopathic effects in tissue culture for viral agents, and microscopic morphology for fungi and parasites. These techniques are reliable but are often time-consuming. Increasingly, the use of nucleic acid probes is becoming a standard method for detection, quantitation, and/or identification in the clinical microbiology laboratory, gradually replacing phenotypic characterization and microscopic visualization methods. This chapter discusses general concepts of diagnostic testing, with an emphasis on detection of bacteria. Detection of viral, fungal, and parasitic pathogens is discussed in greater detail in separate chapters (see Chaps. 214e, 235, and 245e, respectively).

DETECTION METHODS

Reappraisal of the methods employed in the clinical microbiology laboratory has led to the development of strategies for detection of pathogenic agents through nonvisual biologic signal detection systems. A *biologic signal* is generated by detection of a material that can be reproducibly differentiated from other substances present in the sample. Key issues in the use of a biologic signal are distinguishing it from background noise and translating it into meaningful information. Examples of useful materials for detection of biologic signals applicable to clinical microbiology include structural components of bacteria, fungi, and viruses; specific antigens; metabolic end products; unique DNA or RNA base sequences; enzymes; toxins or other proteins; and surface polysaccharides.

A detector is used to sense a signal and discriminate between that signal and background noise. Detection systems range from the trained eyes of a technologist assessing morphologic variations to electronic instruments such as gas-liquid chromatographs or mass spectrometers. The sensitivity with which signals can be detected varies widely. It is essential to use a detection system that discerns small amounts of signal even when biologic background noise is present—i.e., that is both sensitive and specific. Common detection systems include immunofluorescence; chemiluminescence for DNA/RNA probes; flame ionization detection of short- or long-chain fatty acids; and detection of substrate utilization or end-product formation as color changes, of enzyme activity as a change in light absorbance, of turbidity changes as a measure of growth, of cytopathic effects in cell lines, and of particle agglutination as a measure of antigen presence.

Amplification enhances the sensitivity with which weak signals can be detected. The most common microbiologic amplification technique is growth of a single bacterium into a discrete, visible colony on an agar plate or into a suspension containing many identical organisms. The advantage of growth as an amplification method is that it requires only an appropriate growth medium; the disadvantage is the amount of time required. More rapid amplification of biologic signals can be achieved with techniques such as polymerase chain reaction (PCR), ligase chain reaction (LCR), and transcription-mediated amplification (TMA), all of which target the pathogen's DNA/RNA; enzyme immunoassays (EIAs) for antigens and antibodies; electronic amplification (for gas-liquid chromatography assays); antibody capture methods (for concentration and/or separation); and selective filtration or centrifugation.

DIRECT DETECTION

Direct detection refers to detection of pathogens without the use of culture. Molecular methods of direct detection are discussed below.

MICROSCOPY AND STAINING

The field of microbiology was defined largely by the development and use of the microscope. The examination of specimens by microscopic methods rapidly provides useful diagnostic information. Staining techniques permit organisms to be seen more clearly.

The simplest method for microscopic evaluation is the *wet mount*, which is used, for example, to examine cerebrospinal fluid (CSF) for the presence of *Cryptococcus neoformans*, with India ink as a background against which to visualize large-capsuled yeast cells. Wet mounts with dark-field illumination also are used to detect spirochetes in genital lesions and *Borrelia* or *Leptospira* in blood. Skin scrapings and hair samples can be examined with the use of either 10% KOH wet-mount preparations or the *calcofluor white method* and ultraviolet illumination to detect fungal elements as fluorescing structures. Staining of wet mounts—e.g., with lactophenol cotton blue stain for fungal elements—often is used for morphologic identification.

Bacteria are difficult to see by light microscopy unless they are *stained*. Although simple one-step stains can be used, differential stains are more common.

Gram's Stain Gram's stain differentiates between organisms with thick peptidoglycan cell walls (gram-positive) and those with thin peptidoglycan cell walls and outer membranes that can be dissolved with alcohol or acetone (gram-negative). Morphology and Gram's stain characteristics often can be used to categorize stained organisms into groups such as streptococci, staphylococci, and clostridia (Table 150e-1). Gram's stain is particularly useful for examining sputum for polymorphonuclear leukocytes (PMNs) and bacteria. Sputum specimens from immunocompetent patients with ≥ 25 PMNs and < 10 epithelial cells per low-power field often provide clinically useful information. However, the presence in "sputum" samples of > 10 epithelial cells per low-power field and of multiple bacterial types suggests contamination with oral microflora. Despite the difficulty of discriminating between normal microflora and pathogens, Gram's stain may prove useful for specimens from areas with a large resident microflora if a useful biologic marker (signal) is available. Gram's staining of vaginal swab specimens can be used to detect epithelial cells covered with gram-positive bacteria in the absence of lactobacilli and the presence of gram-negative rods—a scenario regarded as a sign of bacterial vaginosis. Similarly, examination of stained stool specimens for leukocytes is useful as a screening procedure before testing for *Clostridium difficile* toxin or other enteric pathogens.

The examination of samples from normally sterile body sites (e.g., CSF or joint, pleural, or peritoneal fluid) with Gram's stain is useful for determining whether bacteria and/or PMNs are present. The sensitivity is such that $> 10^4$ bacteria/mL should be detected. Centrifugation often is performed before staining to concentrate specimens thought to contain low numbers of organisms. This simple method is particularly useful for examination of CSF for bacteria and white blood cells or examination of sputum for mycobacteria.

Acid-Fast Stain The acid-fast stain identifies acid-fast bacteria (AFB; mycobacteria) by their retention of carbol fuchsin dye after acid/organic solvent disruption. Modifications of this procedure allow the differentiation of *Actinomyces* from *Nocardia* or other weakly (or partially) acid-fast organisms. The acid-fast stain is applied to sputum, other fluids, and tissue samples when *Mycobacterium* species are suspected. Because few AFB may be present in an entire smear, even when the specimen has been concentrated by centrifugation, identification of the pink/red AFB against the blue background of the counterstain requires a trained eye. An alternative method for detection of AFB is the auramine-rhodamine fluorescent dye stain.

TABLE 150e-1 INTERPRETATION OF GRAM'S STAIN^a

Morphology	Additional Features	Species
Gram-Negative Bacteria		
Bacilli	Regular bacilli	Enterobacteriaceae (<i>Escherichia coli</i> , <i>Salmonella</i> spp., <i>Shigella</i> spp., others) <i>Pseudomonas aeruginosa</i> <i>Acinetobacter baumannii</i> <i>Stenotrophomonas maltophilia</i>
	Small pleomorphic coccobacilli	<i>Haemophilus influenzae</i> , other <i>Haemophilus</i> spp.
Cocci	Diplococci	<i>Neisseria meningitidis</i> , <i>Neisseria gonorrhoeae</i> , <i>Moraxella catarrhalis</i>
Curved	Comma-shaped	<i>Vibrio</i> spp.
	Comma- or S-shaped	<i>Campylobacter</i> spp.
Gram-Positive Bacteria		
Bacilli	Small regular bacilli	<i>Listeria monocytogenes</i>
	Small pleomorphic bacilli in irregular palisading (picket-fence) arrangement	<i>Corynebacterium</i> spp.
	Large regular bacilli, sometimes with spores	<i>Bacillus</i> spp., <i>Clostridium</i> spp.
	Bacilli arranged in branching chains	<i>Nocardia</i> spp., <i>Actinomyces</i> spp.
	Pale purple or unstained "ghost cells" that are acid-fast	<i>Mycobacterium</i> spp.
Cocci	Cocci arranged in pairs and short chains	<i>Streptococcus pneumoniae</i> , <i>Enterococcus</i> spp.
	Cocci arranged in long chains	<i>Streptococcus pyogenes</i> , <i>Streptococcus agalactiae</i> , other <i>Streptococcus</i> spp.
	Cocci arranged in clusters	<i>Staphylococcus</i> spp., <i>Micrococcus</i> spp.

^aSome important bacteria cannot be seen with Gram's stain because they are too small or too slender or do not retain the stain. These bacteria include *Treponema pallidum*, *Borrelia burgdorferi*, *Chlamydia* spp., *Mycoplasma* spp., and *Ureaplasma* spp.

Fluorochrome Stains Fluorochrome stains such as acridine orange are used to identify white blood cells, yeasts, and bacteria in body fluids. Capsular, flagellar, and spore stains are used for identification or demonstration of characteristic structures.

Immunofluorescent Stains The *direct* immunofluorescent antibody technique uses antibody coupled to a fluorescent compound (e.g., fluorescein) and directed at a specific antigenic target to visualize organisms. When samples are examined under appropriate conditions, the fluorescing compound absorbs ultraviolet light and re-emits light at a higher wavelength that is visible to the human eye. In the *indirect* immunofluorescent antibody technique, an unlabeled (target) antibody binds a specific antigen. The specimen is then stained with a fluorochrome-labeled antibody directed at the target antibody. Because each unlabeled target antibody attached to the appropriate antigen has multiple sites for attachment of the second antibody, the visual signal is amplified. Immunofluorescence is used to detect viral antigens (e.g., cytomegalovirus, herpes simplex virus, and respiratory viruses) within cultured cells or clinical specimens as well as many difficult-to-grow bacterial agents (e.g., *Legionella pneumophila*) in clinical specimens.

MACROSCOPIC ANTIGEN DETECTION

Latex agglutination assays and EIAs are rapid and inexpensive methods for identifying organisms, extracellular toxins, and viral agents by means of protein and polysaccharide antigens. Such assays may be performed directly on clinical samples or after growth of organisms on agar plates or in viral cell cultures. Antibodies coupled to a reporter (such as latex particles or an enzyme) are used for detection of antibody-antigen binding reactions.

Direct agglutination of bacterial cells with specific antibody is simple but relatively insensitive; latex agglutination and EIAs are more sensitive. Some cell-associated antigens, such as capsular polysaccharides and lipopolysaccharides, can be detected by agglutination of a suspension of bacterial cells when antibody is added; this method is useful for typing of the somatic antigens of *Shigella* and *Salmonella*. EIAs employ antibodies coupled to an enzyme, and an antigen-antibody reaction results in the conversion of a colorless substrate to a colored product. Most of these assays provide information about whether antigen is present but do not quantify the antigen. EIAs are also useful

for detecting bacterial toxins—e.g., toxins produced by Shiga toxin-producing *Escherichia coli*.

Rapid and simple immunoassays for antigens of group A *Streptococcus*, influenza virus, and respiratory syncytial virus can be used in the clinical setting without a specialized diagnostic laboratory. Such tests usually are reasonably specific but may have only modest sensitivity.

DETECTION OF PATHOGENIC AGENTS BY SEROLOGIC METHODS

Measurement of serum antibody provides an indirect marker for past or current infection with a specific viral agent or other pathogens, including *Brucella*, *Legionella*, *Rickettsia*, and *Helicobacter pylori*. Serologic methods can be used to determine whether an individual has protective antibody levels or is infected by a specific pathogen. Determination of an antibody level as a measure of current immunity is important in the case of viral agents for which there are vaccines, such as rubella virus and varicella-zoster virus; assays for this purpose normally use one or two dilutions of serum for a qualitative determination of protective antibody levels. Quantitative serologic assays to detect increases in antibody titers most often employ paired serum samples obtained at the onset of illness and 10–14 days later (i.e., acute- and convalescent-phase samples). Since the incubation period before symptoms are noted may be long enough for an antibody response to occur, the demonstration of acute-phase antibody alone is often insufficient to establish the diagnosis of active infection as opposed to past exposure. A fourfold increase in total antibody titer between the acute- and convalescent-phase samples is regarded as evidence for active infection. In addition, IgM may be useful as a measure of an early, acute-phase antibody response. For certain viral agents, such as Epstein-Barr virus, the antibodies produced may be directed at different antigens during different phases of the infection. For this reason, most laboratories test for antibody directed at both viral capsid antigens and antigens associated with recently infected host cells to determine the stage of infection.

DETECTION OF PATHOGENIC AGENTS BY CULTURE

SPECIMEN COLLECTION AND TRANSPORT

To culture bacterial, fungal, or viral pathogens, an appropriate sample must be placed into the proper medium for growth. The success of efforts to identify a specific pathogen often depends on the

collection and transport process coupled to a laboratory-processing algorithm suitable for the specific sample/agent. In some instances, it is better for specimens to be plated at the time of collection rather than first being transported to the laboratory (e.g., urethral swabs being cultured for *Neisseria gonorrhoeae*). In general, the more rapidly a specimen is plated onto appropriate media, the better the chance is for isolating bacterial pathogens. Deep tissue or fluid (pus) samples are more likely to give useful culture results than are superficial swab specimens. **Table 150e-2** lists procedures for collection and transport of common specimens. Because there are many pathogen-specific paradigms for these procedures, it is important to seek advice from the microbiology laboratory when in doubt about a particular situation.

ISOLATION OF BACTERIAL PATHOGENS

Isolation of pathogens from clinical material relies on the use of artificial media that support bacterial growth. Such media are composed of agar, nutrients, and sometimes substances that inhibit the growth of other bacteria. Broth is employed for growth of organisms from specimens with few bacteria, such as peritoneal dialysis fluid or CSF, or from samples in which anaerobes or other fastidious organisms may be present. Broths that allow the growth of small numbers of organisms may be subcultured onto solid medium once growth is detected. The use of liquid medium for all specimens is not worthwhile.

Two basic strategies are used to isolate pathogenic bacteria. The first is to employ enriched media that support the growth of any bacteria that may be present at a site that is normally sterile, such as blood or CSF. The second strategy is to use selective media to isolate specific bacterial species from samples that contain many bacteria under normal conditions (e.g., stool or genital tract secretions). Antimicrobial agents or other substances are incorporated into the agar medium to inhibit growth of all but the bacteria of interest. After incubation, organisms that grow on such media are characterized further to determine whether they are pathogens (**Fig. 150e-1**).

BLOOD CULTURE

The detection of microbial pathogens in blood is difficult because the number of organisms present in the sample is often low and the organisms' integrity and ability to replicate may be damaged by humoral defense mechanisms or antimicrobial agents. Automated blood culture systems detect production of gas (mainly CO₂) by bacteria or yeasts growing in broth in a blood culture bottle. Because the bottles are monitored frequently, a positive culture often is detected more rapidly than by manual techniques, and important information, including the results of Gram's stain and preliminary susceptibility assays, can be obtained sooner. Several factors affect the yield of blood culture from bacteremic patients. Increasing the volume of blood tested increases the chance of a positive culture. For example, a sample-size increase from 10 to 20 mL of blood increases the proportion of positive cultures by ~30%, although this effect is less pronounced in patients with bacterial endocarditis. Obtaining multiple samples for culture (up to three per 24-h period) also increases the chance of detecting a bacterial pathogen. Prolonged culture and blind subculture for detection of most fastidious bacteria (e.g., HACEK organisms) are not needed with automated blood culture systems.

Automated systems also have been applied to the detection of microbial growth from specimens other than blood, such as peritoneal and other normally sterile fluids. *Mycobacterium* species can be detected in certain automated systems if appropriate liquid media are used for culture. Although automated blood culture systems are more sensitive than lysis-centrifugation methods (e.g., Isolator) for yeasts and most bacteria, lysis-centrifugation culture is recommended for filamentous fungi, *Histoplasma capsulatum*, and some fastidious bacteria (*Legionella* and *Bartonella*).

IDENTIFICATION METHODS

Once bacteria are isolated, characteristics that are readily detectable after growth on agar media (colony size, color, hemolytic reactions,

odor, microscopic appearance) may suggest a species, but definitive identification requires additional tests. Identification methods include classic biochemical phenotyping, which is still the most common approach, as well as more sophisticated methods such as mass spectrometry, gas chromatography, and nucleic acid tests (see below).

Biochemical Phenotyping Classic biochemical identification of bacteria entails tests for protein or carbohydrate antigens, the production of specific enzymes, the ability to metabolize specific substrates and carbon sources (such as carbohydrates), or the production of certain metabolites. Rapid versions of some of these tests are available, and many common organisms can be identified on the first day of growth. Other organisms, particularly gram-negative bacteria, require more extensive testing, either manual or automated.

Automated systems allow rapid phenotypic identification of bacterial pathogens. Most of these systems are based on biotyping techniques in which isolates are grown on multiple substrates and the reaction pattern is compared with known patterns for various bacterial species. This procedure is relatively fast. Commercially available systems include miniaturized fermentation, coding to simplify recording of results, and probability calculations for the most likely pathogens. If the biotyping approach is automated and the reading process is coupled to computer-based data analysis, rapidly growing organisms (such as Enterobacteriaceae) can be identified within hours of detection on agar plates.

Several systems use substrates for preformed bacterial enzymes for identification within 2–3 h. These systems do not rely on bacterial growth per se to determine whether a substrate has been used. They employ a heavy inoculum in which specific bacterial enzymes are present in amounts sufficient to convert substrate to product rapidly. In addition, some systems use fluorogenic substrate/end-product detection methods to increase sensitivity through signal amplification.

Gas-Liquid Chromatography Gas-liquid chromatography often is used to detect metabolic end products of bacterial fermentation. One common application is identification of short-chain fatty acids produced by obligate anaerobes during glucose fermentation. Because the types and relative concentrations of volatile acids differ among the various genera and species that make up this group of organisms, such information serves as a metabolic fingerprint for a particular isolate.

Gas-liquid chromatography can be coupled to a sophisticated signal-analysis software system for identification and quantitation of long-chain fatty acids (LCFAs) in the outer membranes and cell walls of bacteria and fungi. For any particular species, the types and relative concentrations of LCFAs are distinctive enough to allow differentiation even from closely related species. An organism may be identified definitively within a few hours after detection of growth on appropriate media.

Matrix-Assisted Laser Desorption/Ionization/Time-of-Flight Mass Spectrometry (MALDI-TOF MS)

MALDI-TOF MS is a rapid and accurate method for identifying microorganisms by protein analysis. The organism is mixed with a chemical matrix, dried on a target plate, and then pulsed with a laser. The laser ionizes and vaporizes the microbial proteins, which then travel through a charged vacuum chamber to a detector. The time of flight to the detector is measured for the individual proteins, and the resulting pattern (or fingerprint) is compared with a library of known patterns for various microorganisms to identify the test organism.

The primary clinical advantage of MALDI-TOF is that it takes only minutes to identify an organism, whereas several hours are needed for conventional phenotyping. MALDI-TOF is highly accurate for identification of most bacteria and yeasts grown on solid agar or in blood culture broth. Other potential uses of MALDI-TOF include identification of bacteria and yeasts directly from clinical specimens (e.g., urine), detection of β -lactamase activity, and strain typing of bacteria; however, these applications are still under development.

TABLE 150e-2 INSTRUCTIONS FOR COLLECTION AND TRANSPORT OF SPECIMENS FOR CULTURE

Note: It is absolutely essential that the microbiology laboratory be informed of the site of origin of the sample to be cultured and the infections that are suspected. This information determines the selection of culture media and the duration of culture.

Type of Culture (Synonyms)	Specimen	Minimal Volume	Container	Other Considerations
Blood				
Blood, routine (blood culture for aerobes, anaerobes, and yeasts)	Whole blood	10 mL in each of 2 bottles for adults and children; 5 mL, if possible, in aerobic bottles for infants; less for neonates	See below. ^a	See below. ^b
Blood for fungi (not yeasts)/ <i>Mycobacterium</i> spp.	Whole blood	10 mL in each of 2 bottles, as for routine blood cultures, or in Isolator tube requested from laboratory	Same as for routine blood culture	Specify "hold for extended incubation," since fungal agents may require 4 weeks to grow.
Blood, Isolator (lysis centrifugation)	Whole blood	10 mL	Isolator tubes	Use mainly for isolation of fungi, <i>Mycobacterium</i> , and other fastidious aerobes and for elimination of antibiotics from cultured blood in which organisms are concentrated by centrifugation.
Respiratory Tract				
Nose	Swab from nares	1 swab	Sterile Culturette or similar transport system containing holding medium	Swabs made of calcium alginate may be used.
Throat	Swab of posterior pharynx, ulcerations, or areas of suspected purulence	1 swab	Sterile Culturette or similar swab specimen collection system containing holding medium	See below. ^c
Sputum	Fresh sputum (not saliva)	2 mL	Commercially available sputum collection system or similar sterile container with screw cap	<i>Cause for rejection:</i> Care must be taken to ensure that the specimen is sputum and not saliva. Examination of Gram's stain, with numbers of epithelial cells and polymorphonuclear leukocytes noted, can be an important part of the evaluation process. Induced sputum specimens should not be rejected.
Bronchial aspirates	Transtracheal aspirate, bronchoscopy specimen, or bronchial aspirate	1 mL of aspirate or brush in transport medium	Sterile aspirate or bronchoscopy tube, bronchoscopy brush in a separate sterile container	Special precautions may be required, depending on diagnostic considerations (e.g., <i>Pneumocystis</i>).
Stool				
Stool for routine culture; stool for <i>Salmonella</i> , <i>Shigella</i> , and <i>Campylobacter</i>	Fresh, randomly collected stool (preferably) or rectal swab	1 g of stool or 2 rectal swabs	Plastic-coated cardboard cup or plastic cup with tight-fitting lid. Other leakproof containers also are acceptable.	If <i>Vibrio</i> spp. are suspected, the laboratory must be notified, and appropriate collection/transport methods should be used.
Stool for <i>Yersinia</i> , <i>Escherichia coli</i> O157	Fresh, randomly collected stool	1 g	Plastic-coated cardboard cup or plastic cup with tight-fitting lid	<i>Limitations:</i> Procedure requires enrichment techniques.
Stool for <i>Aeromonas</i> and <i>Plesiomonas</i>	Fresh, randomly collected stool	1 g	Plastic-coated cardboard cup or plastic cup with tight-fitting lid	<i>Limitations:</i> Stool should not be cultured for these organisms unless also cultured for other enteric pathogens.
Urogenital Tract				
Urine	Clean-voided urine specimen or urine collected by catheter	0.5 mL	Sterile, leakproof container with screw cap or special urine transfer tube	See below. ^d
Urogenital secretions	Vaginal or urethral secretions, cervical swabs, uterine fluid, prostatic fluid, etc.	1 swab or 0.5 mL of fluid	Vaginal and rectal swabs transported in Amies transport medium or similar holding medium for group B <i>Streptococcus</i> ; direct inoculation preferred for <i>Neisseria gonorrhoeae</i>	Vaginal swab samples for "routine culture" should be discouraged whenever possible unless a particular pathogen is suspected. For detection of multiple organisms (e.g., group B <i>Streptococcus</i> , <i>Trichomonas</i> , <i>Chlamydia</i> , or <i>Candida</i> spp.), 1 swab per test should be obtained.
Body Fluids, Aspirates, and Tissues				
Cerebrospinal fluid (lumbar puncture)	Spinal fluid	1 mL for routine cultures; ≥5 mL for <i>Mycobacterium</i>	Sterile tube with tight-fitting cap	Do not refrigerate; transfer to laboratory as soon as possible.
Body fluids	Aseptically aspirated body fluids	1 mL for routine cultures	Sterile tube with tight-fitting cap. Specimen may be left in syringe used for collection if the syringe is capped before transport.	For some body fluids (e.g., peritoneal lavage samples), increased volumes are helpful for isolation of small numbers of bacteria.

Biopsy and aspirated materials	Tissue removed at surgery, bone, anticoagulated bone marrow, biopsy samples, or other specimens from normally sterile areas	1 mL of fluid or a 1-g piece of tissue	Sterile Culturette-type swab or similar transport system containing holding medium. Sterile bottle or jar should be used for tissue specimens.	Accurate identification of specimen and source is critical. Enough tissue should be collected for both microbiologic and histopathologic evaluations.
Wounds	Purulent material or abscess contents obtained from wound or abscess without contamination by normal microflora	2 swabs or 0.5 mL of aspirated pus	Culturette swab or similar transport system or sterile tube with tight-fitting screw cap. For simultaneous anaerobic cultures, send specimen in anaerobic transport device or closed syringe.	<i>Collection:</i> When possible, abscess contents or other fluids should be collected in a syringe (rather than with a swab) to provide an adequate sample volume and an anaerobic environment.
Special Recommendations				
Fungi	Specimen types listed above may be used. When urine or sputum is cultured for fungi, a first morning specimen usually is preferred.	1 mL or as specified above for individual listing of specimens. Large volumes may be useful for urinary fungi.	Sterile, leakproof container with tight-fitting cap	<i>Collection:</i> Specimen should be transported to microbiology laboratory within 1 h of collection. Contamination with normal flora from skin, rectum, vaginal tract, or other body surfaces should be avoided.
<i>Mycobacterium</i> (acid-fast bacilli)	Sputum, tissue, urine, body fluids	10 mL of fluid or small piece of tissue. Swabs should not be used.	Sterile container with tight-fitting cap	Detection of <i>Mycobacterium</i> spp. is improved by use of concentration techniques. Smears and cultures of pleural, peritoneal, and pericardial fluids often have low yields. Multiple cultures from the same patient are encouraged. Culturing in liquid media shortens time to detection.
<i>Legionella</i>	Pleural fluid, lung biopsy, bronchoalveolar lavage fluid, bronchial/transbronchial biopsy	1 mL of fluid; any size tissue sample, although a 0.5-g sample should be obtained when possible	—	Rapid transport to laboratory is critical.
Anaerobic organisms	Aspirated specimens from abscesses or body fluids	1 mL of aspirated fluid, 1 g of tissue, or 2 swabs	An appropriate anaerobic transport device is required. ^e	Specimens cultured for obligate anaerobes should be cultured for facultative bacteria as well. Fluid or tissue is preferred to swabs.
Viruses ^f	Respiratory secretions, wash aspirates from respiratory tract, nasal swabs, blood samples (including buffy coats), vaginal and rectal swabs, swab specimens from suspicious skin lesions, stool samples (in some cases)	1 mL of fluid, 1 swab, or 1 g of stool in each appropriate transport medium	Fluid or stool samples in sterile containers or swab samples in viral Culturette devices (kept on ice but not frozen) are generally suitable. Plasma samples and buffy coats in sterile collection tubes should be kept at 4–8°C. If specimens are to be shipped or kept for a long time, freezing at –80°C is usually adequate.	Most samples for culture are transported in holding medium containing antibiotics to prevent bacterial overgrowth and viral inactivation. Many specimens should be kept cool but not frozen, provided they are transported promptly to the laboratory. Procedures and transport media vary with the agent to be cultured and the duration of transport.

^aFor samples from adults, two bottles (smaller for pediatric samples) should be used: one with dextrose phosphate, tryptic soy, or another appropriate broth and the other with thioglycollate or another broth containing reducing agents appropriate for isolation of obligate anaerobes. For children, from whom only limited volumes of blood can be obtained, only an aerobic culture should be done unless there is specific concern about anaerobic sepsis (e.g., with abdominal infections). For special situations (e.g., suspected fungal infection, culture-negative endocarditis, or mycobacteremia), different blood collection systems may be used (Isolator systems; see table).

^bCollection: An appropriate disinfecting technique should be used on both the bottle septum and the patient. Do not allow air bubbles to get into anaerobic broth bottles. Special considerations: There is no more important clinical microbiology test than the detection of bloodborne pathogens. The rapid identification of bacterial and fungal agents is a major determinant of patients' survival. Bacteria may be present in blood either continuously (as in endocarditis, overwhelming sepsis, and the early stages of salmonellosis and brucellosis) or intermittently (as in most other bacterial infections, in which bacteria are shed into the blood on a sporadic basis). Most blood culture systems employ two separate bottles containing broth medium: one that is vented in the laboratory for the growth of facultative and aerobic organisms and one that is maintained under anaerobic conditions. In cases of suspected continuous bacteremia/fungemia, two or three samples should be drawn before the start of therapy, with additional sets obtained if fastidious organisms are thought to be involved. For intermittent bacteremia, two or three samples should be obtained at least 1 h apart during the first 24 h.

^cNormal microflora in the throat includes α -hemolytic streptococci, saprophytic *Neisseria* spp., diphtheroids, and *Staphylococcus* spp. Aerobic culture of the throat ("routine") includes screening for and identification of β -hemolytic *Streptococcus* spp. and other potentially pathogenic organisms. Although considered components of the normal microflora, organisms such as *Staphylococcus aureus*, *Haemophilus influenzae*, and *Streptococcus pneumoniae* will be identified by most laboratories, if requested. When *Neisseria gonorrhoeae* or *Corynebacterium diphtheriae* is suspected, a special culture request is recommended.

^d(1) When clean-voided specimens, midvoid specimens, and Foley or indwelling catheter specimens yield 50,000 organisms/mL and no more than three species are isolated, the organisms should be identified. Neither indwelling catheter tips nor urine from the bag of a catheterized patient should be cultured. (2) Straight-catheterized, bladder-tap, and similar urine specimens should undergo a complete workup (identification and susceptibility testing) for all potentially pathogenic organisms regardless of colony count. (3) Certain clinical problems (e.g., acute dysuria in women) may warrant identification and susceptibility testing of isolates present at concentrations of >50,000 organisms/mL.

^eAspirated specimens in capped syringes or other transport devices designed to limit oxygen exposure are suitable for the cultivation of obligate anaerobes. A variety of commercially available transport devices may be used. Contamination of specimens with normal microflora from the skin, rectum, vaginal vault, or another body site should be avoided. Collection containers for aerobic culture (such as dry swabs) and inappropriate specimens (such as refrigerated samples; expectorated sputum; stool; gastric aspirates; and vaginal, throat, nose, and rectal swabs) should be rejected as unsuitable.

^fLaboratories generally use diverse methods to detect viral agents, and the specific requirements for each specimen should be checked before a sample is sent.

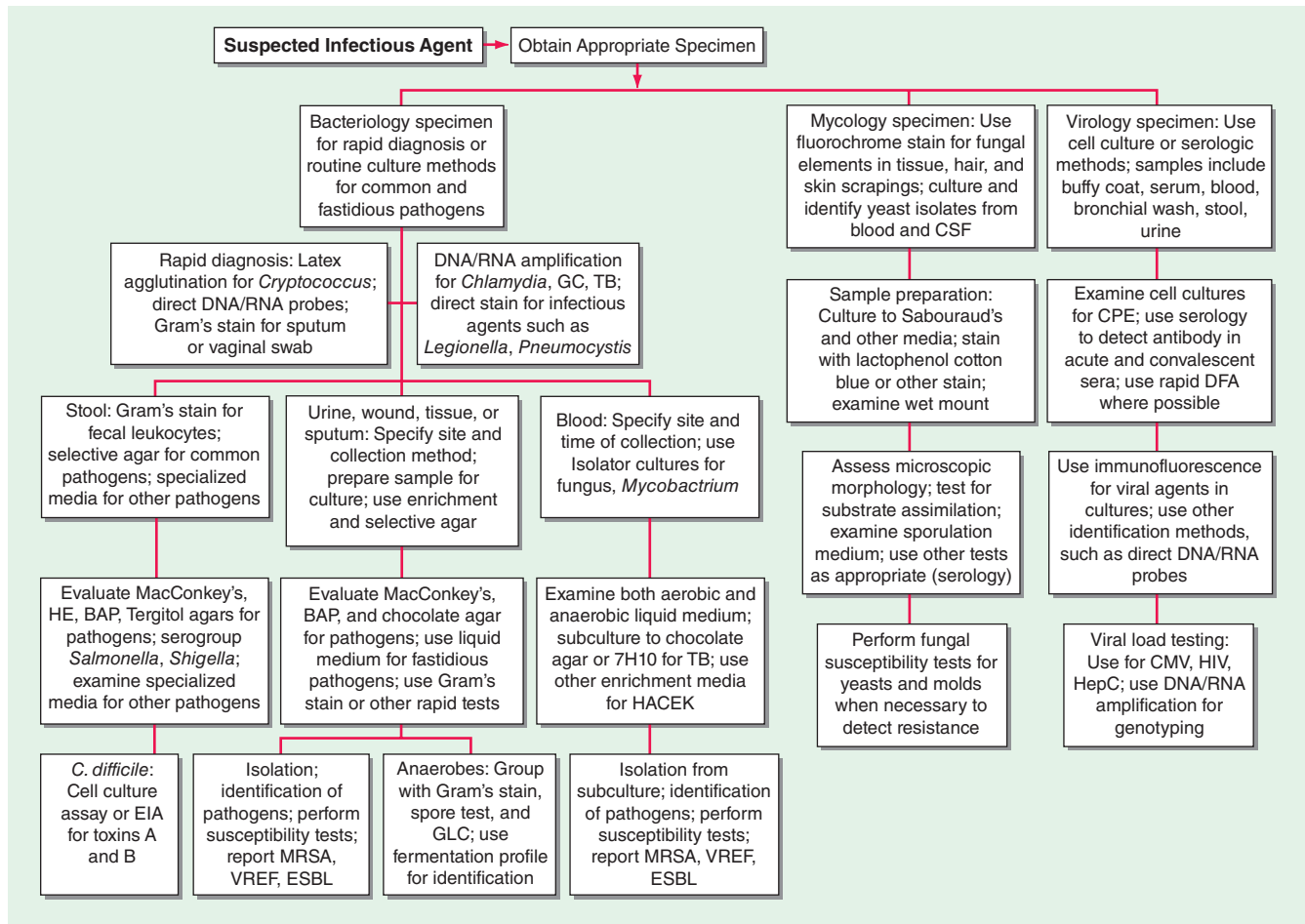


FIGURE 150e-1 Common specimen-processing algorithms used in clinical microbiology laboratories. BAP, blood agar plate; CMV, cytomegalovirus; CPE, cytopathic effects; CSF, cerebrospinal fluid; DFA, direct fluorescent antibody; EIA, enzyme immunoassay; ESBL, extended-spectrum β -lactamase; GBS, group B *Streptococcus*; GC, *Neisseria gonorrhoeae*; GLC, gas-liquid chromatography; HACEK, *Haemophilus aphrophilus/parainfluenzae/paraphrophilus*, *Aggregatibacter actinomycetemcomitans*, *Cardiobacterium hominis*, *Eikenella corrodens*, and *Kingella kingae*; HE, Hektoen enteric medium; HepC, hepatitis C virus; HIV, human immunodeficiency virus; MRSA, methicillin-resistant *Staphylococcus aureus*; TB, *Mycobacterium tuberculosis*; VREF, vancomycin-resistant *Enterococcus faecium*.

NUCLEIC ACID TESTS

Techniques for the detection and quantitation of specific DNA and RNA base sequences in clinical specimens have become powerful tools for the diagnosis of bacterial, viral, parasitic, and fungal infections. Nucleic acid tests are used for four purposes. First, they are used to detect, and sometimes to quantify, specific pathogens in clinical specimens. Second, such tests are used for identification of organisms (usually bacteria) that are difficult to identify by conventional methods. Third, nucleic acid tests are used to determine whether two or more isolates of the same pathogen are closely related (i.e., whether they belong to the same “clone” or “strain”). Fourth, these tests are used to predict the sensitivity of organisms to chemotherapeutic agents. Current technology encompasses a wide array of methods for amplification and signal detection, some of which have been approved by the U.S. Food and Drug Administration (FDA) for clinical diagnosis.

Use of nucleic acid tests generally involves lysis of intact cells or viruses and denaturation of the DNA or RNA to render it single-stranded. Probe(s) or primer(s) complementary to the pathogen-specific target sequence are hybridized to the target sequence in a solution or on a solid support, depending on the system employed. In situ hybridization of a probe to a target also is possible and allows the use of probes with agents present in tissue specimens. Once the probe(s) or primer(s) have been hybridized to the target (biologic signal), a variety of strategies may be employed to detect, amplify, and/or quantify the target-probe complex (Fig. 150e-2).

PROBES FOR DIRECT DETECTION OF PATHOGENS IN CLINICAL SPECIMENS

Nucleic acid probes are used for direct detection of pathogens in clinical specimens without amplification of the target strand of DNA or RNA. Such tests detect a relatively short sequence of bases specific for a particular pathogen on single-stranded DNA or RNA by hybridization of a complementary sequence of bases (probe) coupled to a reporter system that serves as the signal for detection. Nucleic acid probes are available commercially for direct detection of various bacterial and parasitic pathogens, including *Chlamydia trachomatis*, *N. gonorrhoeae*, and group A *Streptococcus*. A combined assay to detect and differentiate agents of vaginitis/vaginosis (*Gardnerella vaginalis*, *Trichomonas vaginalis*, and *Candida* species) also has been approved. An assortment of probes is available for confirming the identity of cultured pathogens, including some dimorphic molds, *Mycobacterium* species, and other bacteria (e.g., *Campylobacter* species, *Streptococcus* species, and *Staphylococcus aureus*). Probes for the direct detection of bacterial pathogens often are aimed at highly conserved 16S ribosomal RNA sequences, of which there are many more copies than there are of any single genomic DNA sequence in a bacterial cell. The sensitivity and specificity of probe assays for direct detection are comparable to those of more traditional assays, including EIA and culture.

In an alternative probe assay called *hybrid capture*, an RNA probe anneals to a DNA target, and the resulting DNA/RNA hybrid is captured on a solid support by antibody specific for DNA/RNA hybrids (concentration/amplification) and detected by chemiluminescent-labeled antibody specific for DNA/RNA hybrids. Hybrid capture

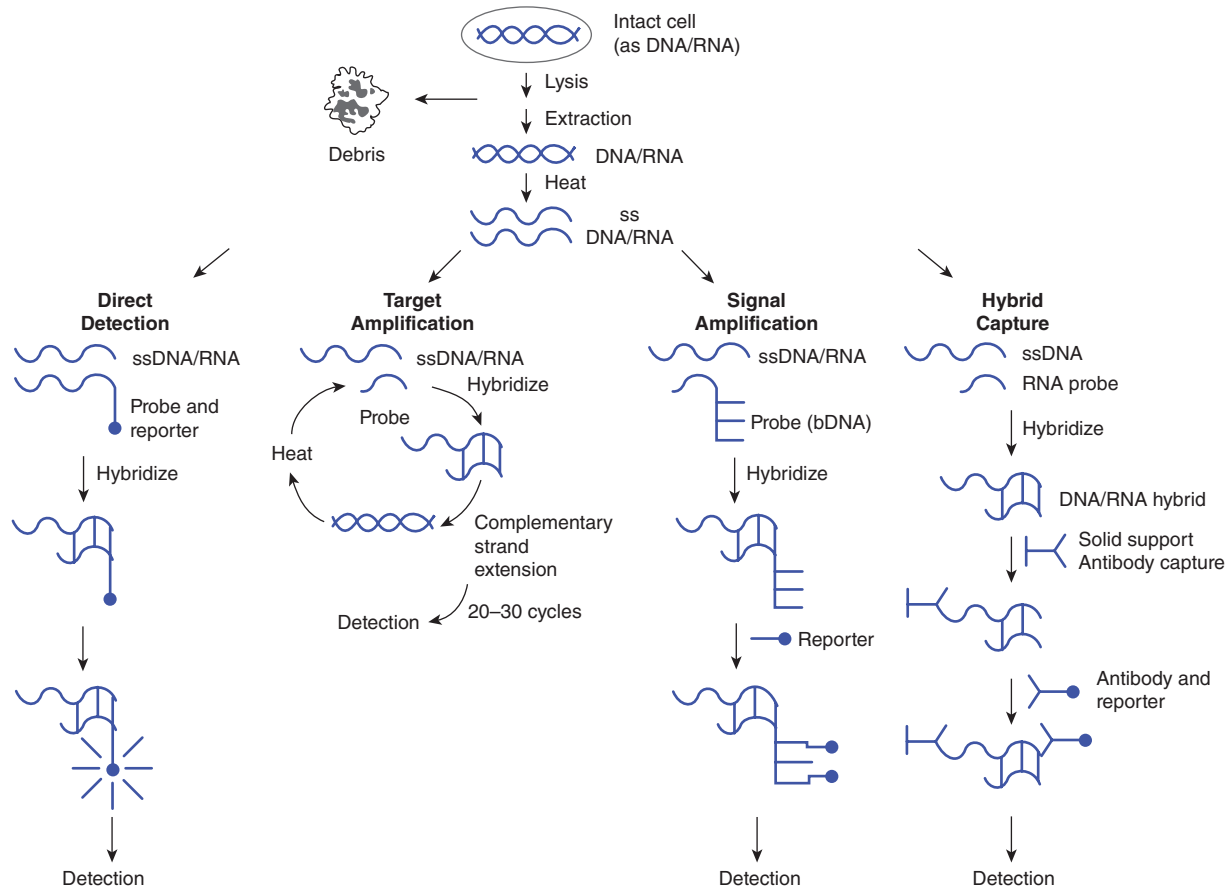


FIGURE 150e-2 Strategies for amplification and/or detection of a target–probe complex. DNA or RNA extracted from microorganisms is heated to create single-stranded (ss) DNA/RNA containing appropriate target sequences. These target sequences may be hybridized directly (direct detection) with probes attached to reporter molecules; they may be amplified by repetitive cycles of complementary strand extension (polymerase chain reaction) before attachment of a reporter probe; or the original target–probe signal may be amplified via hybridization with an additional probe containing multiple copies of a secondary reporter target sequence (branched-chain DNA, or bDNA). DNA/RNA hybrids also can be “captured” on a solid support (hybrid capture), with antibody to the DNA/RNA hybrids used to concentrate them and a second antibody coupled to a reporter molecule attached to the captured hybrid.

assays are available for *C. trachomatis*, *N. gonorrhoeae*, and human papillomavirus.

Many laboratories have developed their own probes for pathogens; however, unless a method-validation protocol for diagnostic testing has been performed, federal law in the United States restricts the use of such probes to research.

NUCLEIC ACID AMPLIFICATION TEST (NAAT) STRATEGIES

There are several methods for amplification (copying) of small numbers of molecules of nucleic acid to readily detectable levels. These NAATs include PCR, LCR, strand displacement amplification, and self-sustaining sequence replication. In each case, exponential amplification of a pathogen-specific DNA or RNA sequence depends on primers that anneal to the target sequence. The amplified nucleic acid can be detected after the reaction is complete or (in real-time detection) as amplification proceeds. The sensitivity of NAATs is far greater than that of traditional assay methods such as culture. PCR, the first and still the most common NAAT, requires repeated heating of the DNA to separate the two complementary strands of the double helix, hybridization of a primer sequence to the appropriate target sequence, target amplification using PCR for complementary strand extension, and signal detection via a labeled probe. Methods for the monitoring of PCR after each amplification cycle—via either incorporation of fluorescent dyes into the DNA during primer extension or use of fluorescent probes capable of fluorescence resonance energy transfer—have decreased the time required to detect a specific target. An alternative NAAT employs transcription-mediated amplification, in which an RNA target

sequence is converted to DNA, which then is exponentially transcribed into an RNA target. The advantage of this method is that only a single heating/annealing step is required for amplification. Identification of otherwise difficult-to-identify bacteria involves an initial amplification of a highly conserved region of 16S rDNA by PCR. Automated sequencing of several hundred bases is then performed, and the sequence information is compared with large databases containing sequence information for thousands of different organisms. Although 16S sequencing is not as rapid as other methods and is still relatively expensive for routine use in a clinical microbiology laboratory, it is becoming the definitive method for identification of unusual or difficult-to-cultivate organisms.

QUANTITATIVE NUCLEIC ACID TEST STRATEGIES

With the advent of newer therapeutic regimens for HIV-associated disease, cytomegalovirus infection, and hepatitis B and C virus infections, the response to therapy has been monitored by determining both genotype and viral load at various times after treatment initiation. Quantitative NAATs are available for HIV (PCR), cytomegalovirus (PCR), hepatitis B virus (PCR), and hepatitis C virus (PCR and TMA). Many laboratories have validated and perform quantitative assays for these and other pathogens (e.g., Epstein-Barr virus), using analyte-specific reagents for NAATs.

Branched-chain DNA (bDNA) testing is an alternative to NAATs for quantitative nucleic acid testing. In such testing, bDNA attaches to a site different from the target-binding sequence of the original probe. Chemiluminescence-labeled oligonucleotides can then bind to multiple repeating sequences on the bDNA. The amplified bDNA signal

150e-8 is detected by chemiluminescence. bDNA assays for viral load of HIV, hepatitis B virus, and hepatitis C virus have been approved by the FDA. The advantage of bDNA assays over PCR is that only a single heating/annealing step is required to hybridize the target-binding probe to the target sequence for amplification.

APPLICATIONS OF NUCLEIC ACID TESTS

The number of FDA-approved NAATS has increased dramatically and includes tests for *Mycobacterium tuberculosis*, *N. gonorrhoeae*, *C. trachomatis*, group B *Streptococcus*, and methicillin-resistant *S. aureus*. FDA-approved multiplex NAAT panels for detection of several respiratory or gastrointestinal pathogens also are available. Again, many laboratories have used commercially available reagents and analyte-specific reagents to create laboratory-developed tests for diagnostic use. Nucleic acid tests are useful to detect and identify difficult-to-grow or noncultivable bacterial pathogens such as *Legionella*, *Ehrlichia*, *Rickettsia*, *Babesia*, *Borrelia*, and *Tropheryma whipplei*. In addition, methods for rapid detection of agents of public health concern, such as *Francisella tularensis*, *Bacillus anthracis*, smallpox virus, and *Yersinia pestis*, have been developed and are available in regional (state) laboratories and at the Centers for Disease Control and Prevention.

Nucleic acid tests are also used to determine how close the relationship is among different isolates of the same species of pathogen. The demonstration that bacteria of a single clone have infected multiple patients in the context of a possible source of transmission (e.g., a health care provider) offers confirmatory evidence for an outbreak. Pulsed-field gel electrophoresis remains the gold standard for bacterial strain analysis. This method involves the use of restriction enzymes that recognize rare sequences of nucleotides to digest bacterial DNA, resulting in large DNA fragments. These fragments are separated by gel electrophoresis with variable polarity of the electrophoretic current and then are visualized. Similar band patterns (i.e., differences in ≤ 3 bands) suggest that different bacterial isolates are closely related, or clonal. Simpler methods of strain typing include sequencing of single or multiple genes and PCR-based amplification of repetitive DNA sequences in the bacterial chromosome. Whole bacterial genome sequencing has been used in investigation of some outbreaks, but this method is still under development.

Future applications of nucleic acid testing probably will include the replacement of culture for identification of many pathogens with additional multiplex NAAT and solid-state DNA/RNA chip technology, in which thousands of unique nucleic acid sequences can be detected on a single silicon chip.

SUSCEPTIBILITY TESTING OF BACTERIA

A principal responsibility of the clinical microbiology laboratory is to determine which antimicrobial agents inhibit a specific bacterial isolate. Such testing is used for patient care and for monitoring of infection control problems, such as methicillin-resistant *S. aureus* or vancomycin-resistant *Enterococcus faecium*. Two approaches are useful. The first is a qualitative assessment of susceptibility, with responses

categorized as susceptible, resistant, or intermediate. This approach can involve either the placement of paper disks containing antibiotics on an agar surface inoculated with the bacterial strain to be tested (Kirby-Bauer or disk-diffusion method), with measurement of the zones of growth inhibition after incubation, or the use of broth cultures containing a set concentration of antibiotic (breakpoint method). These methods have been calibrated carefully against quantitative methods and clinical experience with each antibiotic, and zones of inhibition and breakpoints have been determined on a species-by-species basis.

The second approach is to inoculate the test strain of bacteria into a series of cultures with increasing concentrations of antibiotic. The lowest concentration of antibiotic that inhibits visible microbial growth of the bacteria is known as the *minimal inhibitory concentration* (MIC). If tubes in which no growth is seen are subcultured, the minimal concentration of antibiotic required to kill 99.9% of the starting inoculum also can be determined (*minimal bactericidal concentration*, or MBC). The MIC value can be given a categorical interpretation of susceptible, resistant, or intermediate and so is more widely used than the MBC. Quantitative susceptibility testing using microbroth dilution in microwell plates or other miniaturized testing platforms has been automated and is used commonly in clinical laboratories. The epsilon-meter test (E-test), a novel method for determination of the MIC, uses a plastic strip with a known gradient of antibiotic concentrations along its length. When the strip is placed on the surface of an agar plate seeded with the bacterial strain to be tested, antibiotic diffuses into the medium, and bacterial growth is inhibited. For some organisms, such as obligate anaerobes and some β -hemolytic streptococci, routine susceptibility testing generally is not performed because of the difficulty of growing the organisms or the predictable sensitivity of most isolates to specific antibiotics.

SUSCEPTIBILITY TESTING OF FUNGI

With the advent of many new agents for treating yeasts and systemic fungal agents, the need for testing of individual isolates for susceptibility to specific antifungal agents has increased. In the past, few laboratories participated in such testing because of a lack of standard methods like those available for testing bacterial agents. However, several systems have been approved for antifungal susceptibility testing. These methods, which determine the minimal fungicidal concentration (MFC), are similar to the broth microdilution methods used to determine the MIC for bacteria. The E-test method is approved for testing the susceptibility of yeasts to fluconazole, itraconazole, and flucytosine, and disk diffusion can be used to test the susceptibility of *Candida* species to fluconazole and voriconazole. Methods for determining the MFC of a drug against fungal agents such as *Aspergillus* species are technically difficult, and most clinical laboratories refer requests for such testing to reference laboratories.

ANTIVIRAL TESTING

See Chaps. 214e and 215e.

151e Climate Change and Infectious Disease

Aaron S. Bernstein

The release of greenhouse gases—principally carbon dioxide—into Earth’s atmosphere since the late nineteenth century has contributed to a climate unfamiliar to our species, *Homo sapiens*. This new climate has already altered the epidemiology of some infectious diseases. Continued accumulation of greenhouse gases in the atmosphere will further alter the planet’s climate. In some cases climate change may establish conditions favoring the emergence of infectious diseases, while in others it may render areas that are presently suitable for certain diseases unsuitable. This chapter presents the current state of knowledge regarding the known and prospective infectious-disease consequences of climate change.

OVERVIEW

The term *climate change* refers to long-term alterations in temperature, precipitation, wind, humidity, and other components of weather. Over the past 2.5 million years, the earth has warmed and cooled, cycling between glacial and interglacial periods during which average global temperatures moved up and down by 4–7°C. During the last glacial period, which ended roughly 12,000 years ago, global temperatures were, on average, 5°C cooler than in the mid-twentieth century (Fig. 151e-1).

The present climate period, known as the Holocene, is remarkable for its stability: temperatures have largely remained within a range of 2–3°C. This stability has enabled the successful population and cultivation of much of the earth’s landmass by humanity. Current climate change differs from that in the past not only because its primary cause is human activities but also because its pace is faster. The 5°C of warming that occurred at the end of the last ice age took roughly 5000 years, whereas such a temperature increment may occur within the next 150 years unless the release of greenhouse gases is substantially reduced in coming decades. The current rate of warming on Earth is

unprecedented in the last 50 million years. Climate science, although still a relatively new discipline, has provided an ever-clearer picture of how the changing chemistry of the atmosphere has influenced, and will continue to influence, the global climate.

GREENHOUSE GASES

Greenhouse gases (Table 151e-1 and Fig. 151e-2) are a group of gases in Earth’s atmosphere that absorb infrared radiation and thus retain heat inside the atmosphere. In the absence of these gases, the earth’s average temperature would be about 33°C colder. Carbon dioxide, released into the atmosphere primarily from fossil fuel combustion and deforestation, has had the greatest effect on climate since the Industrial Revolution. Of note, the Swedish scientist Svante Arrhenius first suggested in the late nineteenth century that the addition of carbon dioxide to the earth’s atmosphere would increase the planet’s surface temperature. Water vapor is the most abundant and a highly potent greenhouse gas but, given its short atmospheric life span and sensitivity to temperature, is not a major factor in recently observed climate change.

The atmosphere, some of the aerosols suspended in it, and clouds reflect a portion of incoming solar radiation back toward space. The remainder reaches Earth’s surface, where it is absorbed and some is then emitted back at the atmosphere. The earth emits energy absorbed from the sun at longer wavelengths, primarily infrared, that greenhouse gases are able to absorb. The change in wavelength that occurs as solar radiation is absorbed and re-emitted from the earth’s surface is fundamental to the greenhouse effect (Fig. 151e-3).

TEMPERATURE

Climate change has become nearly synonymous with global warming, as a clear signal from rising greenhouse gas concentrations has been an increase in the mean global surface temperature of ~0.85°C since 1880. However, this mean warming belies warming that is occurring much faster in certain regions. The Arctic has warmed twice as fast overall, and winters are warming faster than summers. Nighttime minimum temperatures are also rising faster than daytime high temperatures. Each of these nuances bears upon the incidence of infectious diseases in general and vector-borne disease specifically.

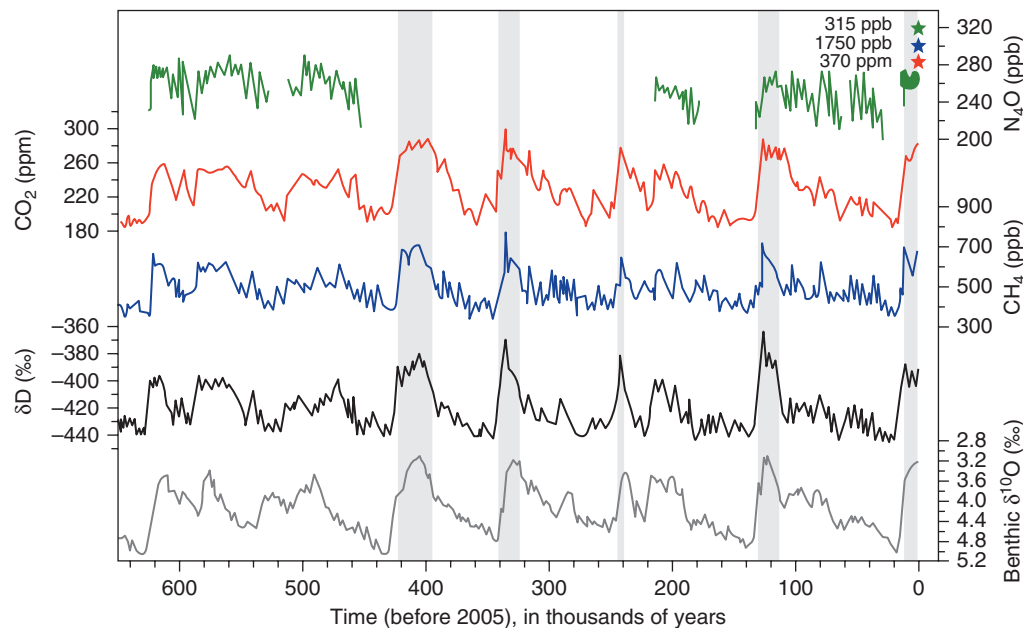


FIGURE 151e-1 Overview of the earth’s temperature and primary greenhouse gases over the last 600,000 years. Variations of deuterium (δD ; black) serve as a proxy for temperature. Atmospheric concentrations of greenhouse gases— CO_2 (red), CH_4 (blue), and nitrous oxide (N_2O ; green)—were derived from air trapped within Antarctic ice cores and from recent atmospheric measurements. Shaded areas indicate interglacial periods. Benthic $\delta^{18}O$ marine records (dark gray) are a proxy for global ice-volume fluctuations and can be compared to the ice core data. Downward trends in the benthic $\delta^{18}O$ curve reflect increasing ice volumes on land. The stars and labels indicate atmospheric concentrations as of the year 2000. CO_2 levels surpassed 400 ppm as of 2013 and are rising at a rate of 2–2.5 ppm per year. (From Intergovernmental Panel on Climate Change Fourth Assessment Report. Working Group I, Chapter 6, Figure 6.3. Cambridge University Press, 2007.)

TABLE 151e-1 GREENHOUSE GASES: SOURCES, SINKS, AND FORCINGS

Gas	Human Sources	Sink ^a	Radiative Forcing ^b (95% Confidence Interval)
Carbon dioxide (CO ₂)	Fossil fuel combustion, deforestation	Uptake by oceans (~30%), plants	1.68 (1.33–2.03)
Methane (CH ₄)	Fossil fuel production, ruminant animals, decomposition in landfills	Hydroxyl radicals in the troposphere	0.97 (0.74–1.20)
Nitrous oxide (N ₂ O)	Fertilizer, fossil fuel combustion, biomass burning, livestock manure	Photolysis in the stratosphere	0.17 (0.14–0.23)
Halocarbons	Refrigerants, electrical insulation, aluminum production	Hydroxyl radicals in the troposphere, sunlight in the stratosphere	0.18 (0.01–0.35)

^aIn this table, a *sink* refers to the place where greenhouse gases are naturally stored or the mechanism through which they are destroyed. ^bRadiative forcing, measured in watts per meter squared, refers to how much an entity can alter the balance of incoming and outgoing radiation to and from Earth's atmosphere. It is measured relative to a preindustrial (i.e., 1750) baseline. Greenhouse gases have a positive "forcing"; that is, on balance, they increase the amount of radiation (and specifically infrared radiation) that is retained in Earth's atmosphere. (Sources: Intergovernmental Panel on Climate Change Fifth Assessment Report, Working Group 1, Chapter 8; American Chemical Society "Greenhouse gas sources and sinks," available at www.acs.org/content/acs/en/climate-science/greenhouse-gases/sources-and-sinks.html)

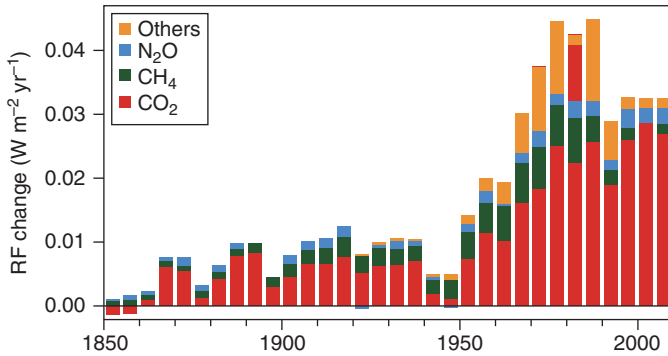


FIGURE 151e-2 Acceleration of radiative forcing (RF) from release of major greenhouse gases, 1850–2011. For definition of radiative forcing, see footnote b to Table 151e-1. (From Intergovernmental Panel on Climate Change Fifth Assessment Report, Working Group 1, Chapter 8, Figure 8.6, p. 677.)

A moderate projection based on the best available scientific evidence suggests that average global temperatures will warm an additional 1.4–3.1°C by 2100 as compared to the period 1986–2005. Because of climate change, extreme heat waves have already become more common and are expected to be even more frequent later in this century. Besides contributing directly to morbidity and mortality in human populations, heat waves wilt crops and are expected to contribute substantially to predicted agricultural losses. For example, the 2010 heat wave in Russia, which was unprecedented in its severity, contributed to hundreds of forest fires that generated enough air pollution to kill an estimated 56,000 people and that burned 300,000 acres of crops, including roughly 25% of the nation's wheat fields. Nutritional deficiencies underlie a substantial portion of the global burden of many infectious diseases.

PRECIPITATION

In addition to changing temperature, the emission of greenhouse gases and the consequent increase in energy in Earth's atmosphere have influenced the planet's water cycle. Since 1950, substantial increases in the heaviest precipitation events (i.e., those above the 95th percentile) have been observed in Europe and North America. While trends over

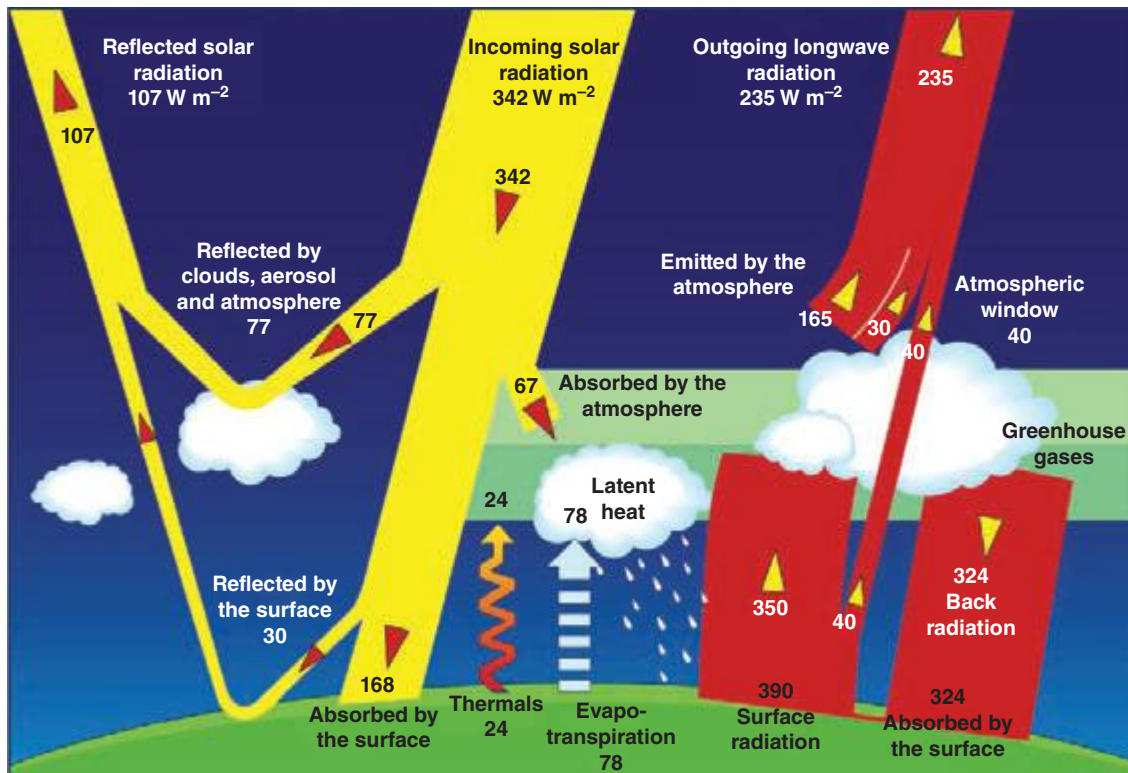


FIGURE 151e-3 Earth's energy balance. (Courtesy of NASA CERES project. Data from Trenberth KE et al: Earth's global energy budget. Bull Am Meteor Soc 90:311, 2009.)

that same interval are less clear in other regions because of limited data, regions of Southeast Asia and southern South America have likely experienced increases in heavy precipitation as well. Other areas have seen greater drought, notably southern Australia and the southwestern United States.

A warmer atmosphere holds more water vapor; specifically, there is 6–7.5% more water vapor per degree (Celsius) of warming in the lower atmosphere. For areas that have traditionally had more precipitation on average, warming tends to promote heavier precipitation events. In contrast, in regions prone to drought, warming tends to result in greater periods between rainfalls and in the risk of drought.

HURRICANES

The world's oceans have absorbed 90% of the excess heat that greenhouse gases have kept in Earth's atmosphere since the 1960s. Ocean heat provides energy for hurricanes, and warmer years tend to have greater hurricane activity. Atlantic hurricanes are the best studied and have the most data available. An analysis of satellite observations from 1983 to 2005 has shown a trend toward increasing severity—albeit decreasing frequency—of Atlantic hurricanes. Modeling of future tropical cyclones suggests that their intensity may increase 2–11% by 2100 and that the average storm will bring 20% more rainfall.

SEA LEVEL RISE

Between 1901 and 2010, the global sea level rose ~200 mm, or ~1.7 mm per year on average. From 1993 to 2010, the rate of rise nearly doubled—i.e., to 3.2 mm annually. Most of this sea level rise has resulted from the thermal expansion of water. Glacial ice melt is the second greatest factor, and its contribution is accelerating. By 2100, global sea level may rise by ≥ 500 mm from 1986–2005 levels, with an annual rate of rise of 8–16 mm at the century's end. A large section of the West Antarctic ice sheet has begun to fall apart, and its melting alone may cause sea level to rise by ≥ 3 m in coming centuries.

Sea level rise is not uniform. The rate of rise on the eastern seaboard of North America has been roughly double the global rate. Compounding sea level rise is the subsidence of coastal areas due to human settlement. In the absence of levee upgrades, an estimated 170 million people living near coasts worldwide will be at risk of flooding in 2100 because of the combined effects of subsidence, erosion, and sea level rise.

Along with extreme storms and overuse of coastal aquifers, rising seas also contribute to salinization of coastal groundwater. About 1 billion people rely on coastal aquifers for potable water.

EL NIÑO SOUTHERN OSCILLATION

The *El Niño Southern Oscillation* (ENSO) refers to periodic changes in water temperature in the eastern Pacific Ocean that occur roughly every 5 years. ENSO cycles have dramatic effects on weather around the globe. Warmer-than-average water temperatures in the eastern Pacific define *El Niño* events (see below), whereas cooler-than-average water temperatures define La Niña periods. Evidence is accruing that climate change may be increasing the frequency and severity of *El Niño* events.

El Niño events drive alterations in weather worldwide (Fig. 151e-4) and are associated with extreme events and consequently higher rates of morbidity and mortality. Hurricane Mitch, one of the most powerful hurricanes ever observed, with winds reaching 290 km/h, dropped 1–1.8 m (3–6 feet) of rain over 72 h on parts of Honduras and Nicaragua. As a result of this storm, 11,000 people died and 2.7 million were displaced. Outbreaks of cholera, leptospirosis, and dengue occurred in the storm's aftermath.

POPULATION MIGRATION AND CONFLICT

The final common outcome of all climate change effects is population migration. Sea level rise, extreme heat and precipitation, droughts, and salinization of water supplies all conspire to make regions (including some inhabited by humans for millennia) uninhabitable. Among climate change migrants in the near future may be the inhabitants of low-lying South Pacific islands that are vulnerable to sea level rise and residents of the Alaskan archipelago, where melting of permafrost has rendered traditional means of cold food storage difficult.

Climate change may also be contributing to humanitarian crises and conflicts. A severe 2011 drought in East Africa may have incited the Somali famine that resulted in 1 million refugees; mortality rates reached 7.4/10,000 in some refugee camps. Crop losses associated with the 2010



FIGURE 151e-4 Characteristic weather anomalies, by season, during *El Niño* events. (Source: www.cpc.ncep.noaa.gov/products/precip/CWlink/ENSO/ENSO-Global-Impacts/High-Resolution/.)

EFFECTS OF CLIMATE CHANGE ON INFECTIOUS DISEASE

The incidence of most, if not all, infectious diseases depends on climate. For any given infection, however, climate change is but one of many factors that determine disease epidemiology, and often it is not the most influential factor. Even in instances in which climate change creates conditions favorable to the spread of infections, diseases may be kept in check through interventions such as vector control or antibiotic treatment.

Detecting climate-change influence on an emerging human disease can be challenging. Research with animal pathogens, which in most instances is less well monitored and intervened upon than that with their human counterparts, has suggested how climate change may influence disease spread. For example, the life cycle of nematode parasites of caribou and musk oxen shortens as temperatures rise. As the Arctic has warmed, higher nematode burdens and consequently higher rates of morbidity and mortality have been observed. Other examples from animals, such as the spread of the protozoan parasite *Perkinsus marinus* in oysters, demonstrate how warming can enable range expansion of pathogens previously held in check by colder temperatures.

As these and other examples from studies of animals make clear, the influence of climate change on infectious diseases can be pronounced. The following sections deal with the infectious diseases for which research has explored the influence of climate change.

VECTOR-BORNE DISEASE

Because insects are cold-blooded, ambient temperature dictates their geographic distribution. With increases in temperatures (in particular, nighttime minimum temperatures), insects are freed to move poleward and up mountainsides. At the same time, as new areas become climatically suitable, current mosquito habitats may become unsuitable as a result of heat extremes.

In addition, insects tend to be sensitive to water availability. Mosquitoes that transmit malaria, dengue, and other infections may breed in pools of water created by heavy downpours. As has been observed in the Amazon, breeding pools can also appear during periods of drought when rivers recede and leave behind stagnant pools of water for *Anopheles* mosquitoes. These circumstances have raised interest in the potentially favorable impact of water-cycle intensification on the spread of mosquito-borne disease.

Malaria • TEMPERATURE Higher temperatures promote higher mosquito-biting rates, shorter parasite reproductive cycles, and the potential for the survival of mosquito vectors of *Plasmodium* infection in locations previously too cold to sustain them. Recent modeling experiments identify highland areas of East Africa and South America as perhaps most vulnerable to increased malarial incidence as a result of rising temperatures. In addition, a recent analysis of interannual malaria in Ecuador and Colombia has documented a greater incidence of malaria at higher altitudes in warmer years. Highland populations may be more vulnerable to malaria epidemics because they lack immunity.

Although rising temperature has the potential to expand the viable range of disease, malaria incidence is not associated with temperature in a strictly linear fashion. While mosquitoes and parasites may adapt to a warming climate, the present optimal temperature for malaria transmission is ~25°C, with a range of transmission temperatures between 16°C and 34°C. Rising temperatures also can have differential effects on parasite development during external incubation and on the mosquitoes' gonotrophic cycle. Asynchrony between these two temperature-sensitive processes has been shown to decrease the vectorial capacity of mosquitoes.¹

¹ rVc is the vectorial capacity relative to the vector-to-human population ratio and is defined by the equation $rVc = a^2 b_h b_m e^{-\mu_m n} / \mu_m$ where a is the vector biting rate; b_h is the probability of vector-to-human transmission per bite; b_m is the probability of human-to-vector infection per bite; n is the duration of the extrinsic incubation period; and μ_m is the vector mortality rate.

PRECIPITATION The abundance of *Anopheles* mosquitoes is strongly correlated with the availability of surface water pools for mosquito breeding, and biting rates have been linked to soil moisture (a surrogate for breeding pools). Research in the East African highlands has documented that increased variance in rainfall over time has strengthened the association between precipitation and disease incidence. These disease-promoting effects of precipitation may be countered by the potential for extreme rainfall to flush mosquito larvae from breeding sites.

PROJECTIONS Climate models have begun to deliver output on regional scales, permitting projections of climate-suitable regions to assist national and local health authorities. Climate models speak to the temperature and precipitation ranges necessary for malaria transmission but do not—and cannot—account for the capacity of malaria control programs to halt the spread of disease. The global reduction in malaria distribution over the past century makes it clear that, even with climate change, malaria occurs in far fewer places today because of public health interventions.

Despite intensive efforts, malaria remains the single greatest vector-borne disease cause of morbidity and death in the world. Particularly in regions that are most affected by malaria and where the public health infrastructure is inadequate to contain it, climate modeling may provide a useful tool in determining where the disease may spread. Recent modeling studies in sub-Saharan Africa have suggested that, although East African nations may encompass regions that will become more climatically suitable for malaria over this century, West African nations may not. By 2100, temperatures in West Africa may largely exceed those optimal for malaria transmission, and the climate may become drier; in contrast, higher temperatures and changes in precipitation may allow malaria to move up the mountainsides of East African countries. Climate change may create conditions favorable to malaria in subtropical and temperate regions of the Americas, Europe, and Asia as well.

Dengue Like malaria epidemics, dengue fever epidemics depend on temperature (Fig. 151e-5). Higher temperatures increase the rate of larval development and accelerate the emergence of adult *Aedes* mosquitoes. The daily temperature range may also influence dengue virus transmission, with a smaller range corresponding to a higher transmission potential. Temperatures <15°C or >36°C also substantially reduce mosquito feeding. In a *Rhesus* model of dengue, viral replication can occur in as little as 7 days with temperatures of >32–35°C; at 30°C, replication takes ≥12 days; and replication does not reliably occur at 26°C. Research on dengue in New Caledonia has shown peak transmission at ~32°C, reflecting combined effects of a shorter extrinsic incubation period, a higher feeding frequency, and more rapid development of mosquitoes. Along with temperature, peak relative humidity is a strong predictor of dengue outbreaks.

The association between dengue epidemics and precipitation is less consistent in the peer-reviewed literature, possibly because of the mosquito vector's greater reliance on domestic breeding sites than on natural pools of water. For instance, in some studies, increased access to a piped water supply has been linked to dengue epidemics, presumably because of associated increased domestic water storage. Nonetheless, several studies have established rainfall as a predictor of the seasonal timing of dengue epidemics.

The current global distribution of dengue largely overlaps the geographic spread of *Aedes* mosquitoes (Fig. 151e-6). The presence of *Aedes* without dengue endemicity in large regions of North and South America and Africa illustrates the relevance of variables other than climate to disease incidence. Nevertheless, coupled climatic-epidemiologic modeling suggests dramatic shifts in the relative vectorial capacity for dengue by the end of this century should little or no mitigation of greenhouse gas emissions occur (Fig. 151e-7).

Other Arbovirus Infections Climate change may favor increased geographic spread of other arboviral diseases, including Chikungunya virus disease, West Nile virus disease, and eastern equine encephalitis. Chikungunya virus disease emerged in Italy in 2007, having previously

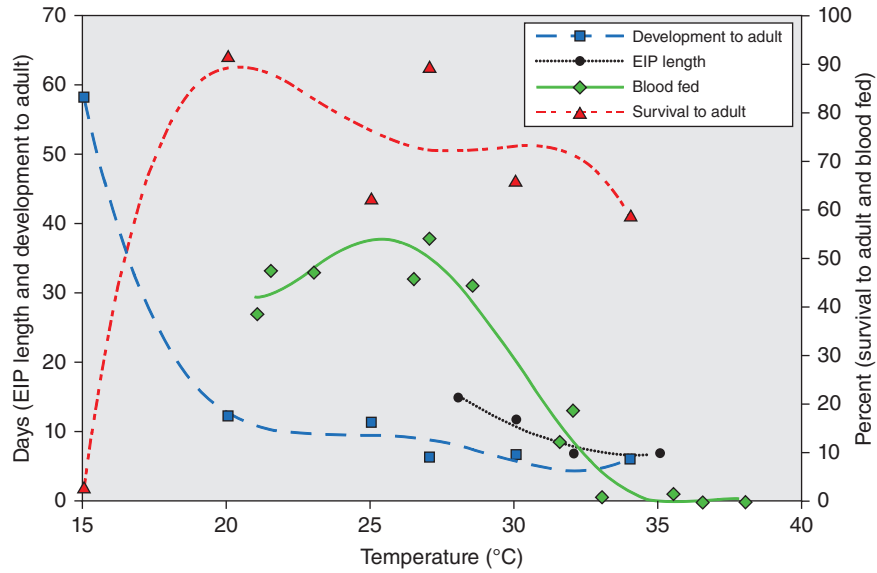


FIGURE 151e-5 Effects of temperature on variables associated with dengue transmission. Shown are the number of days required for development of immature *Aedes aegypti* mosquitoes to adults, the length of the dengue virus type 2 extrinsic incubation period (EIP), the percentage of *A. aegypti* mosquitoes that complete a blood meal within 30 min after a blood source is made available, and the percentage of hatched *A. aegypti* larvae surviving to adulthood. (Reproduced from CW Morin et al: *Climate and dengue transmission: evidence and implications. Environ Health Perspect* 121:1264, 2013.)

been mostly a disease of African nations. Climate models predict that, should competent vectors be present, conditions will be suitable for Chikungunya virus to gain a foothold in Western Europe, especially France, in the first half of the twenty-first century. In North America, areas favorable to West Nile virus outbreaks are expected to shift northward in this century. Current hotspots in North America are the California Central Valley, southwestern Arizona, southern Texas, and Louisiana, which have both compatible climates and avian reservoirs for the disease. By mid-century, the upper Midwest and New England will be more climatically suited to West Nile virus; by the end of the century, the area of risk may shift even further north to southern Canada.

Whether the disease will ultimately move northward will depend on reservoir availability and mosquito control programs, among other factors.

Lyme Disease In the past few decades, *Ixodes scapularis*, the primary tick vector for Lyme disease as well as for anaplasmosis and babesiosis in New England, has become established in Canada because of warming temperatures. With climate change, the range of the tick is expected to expand further (Fig. 151e-8).

Lyme disease, caused by the spirochete *Borrelia burgdorferi*, is the most commonly reported vector-borne disease in North America, with ~30,000 cases per year. The model used in Fig. 151e-8 showed 95%

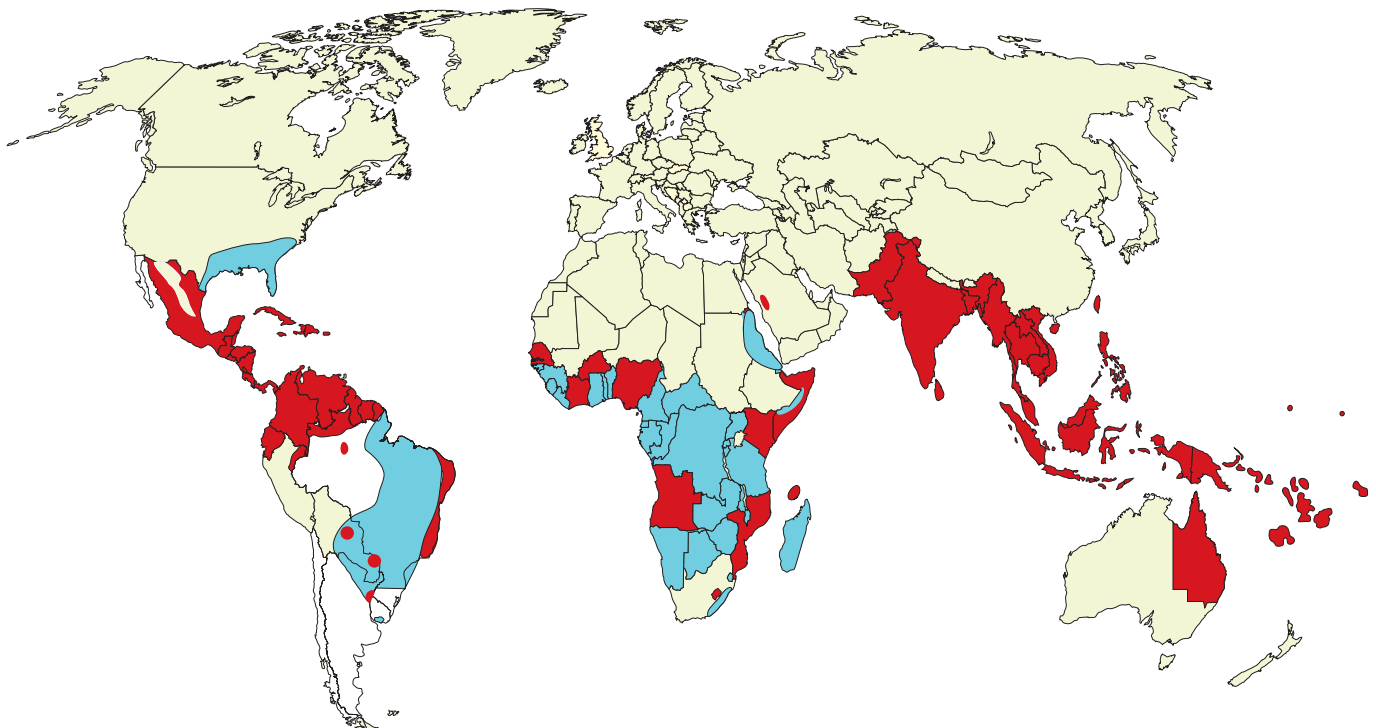


FIGURE 151e-6 Distribution of *Aedes aegypti* mosquitoes (turquoise) and dengue fever epidemics (red). (Map produced by the Agricultural Research Service of the U.S. Department of Agriculture.)

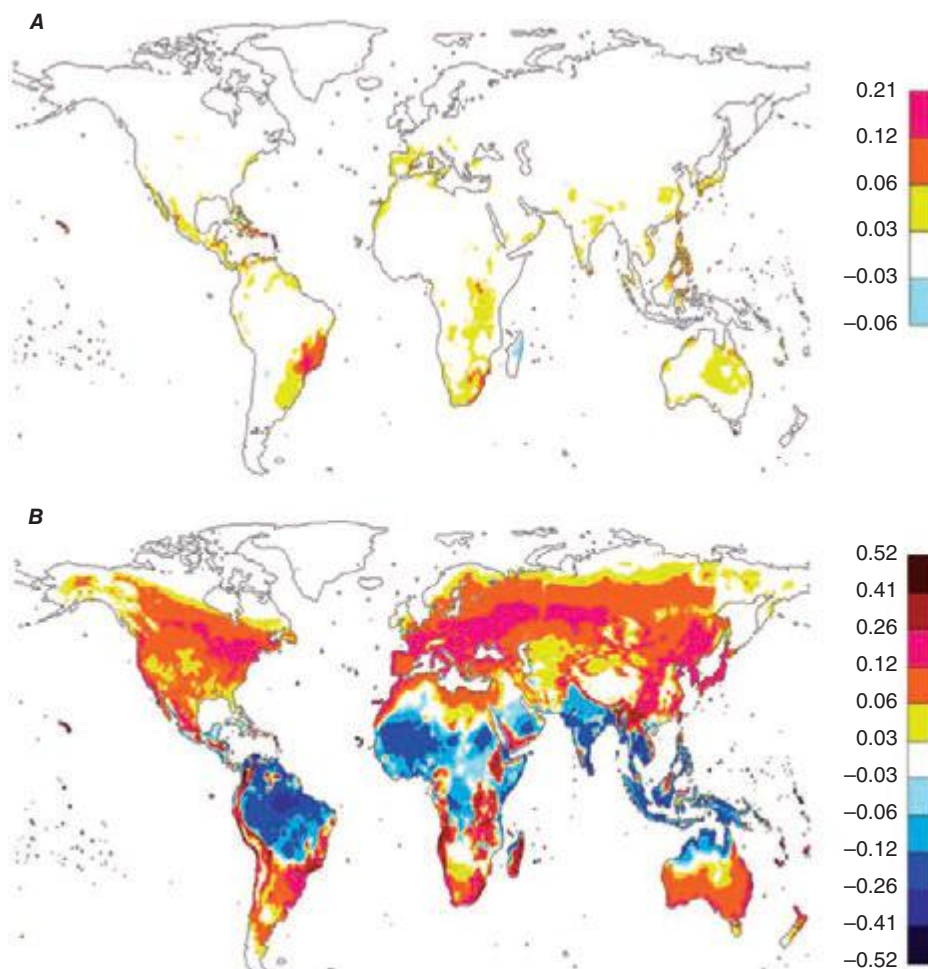


FIGURE 151e-7 Trend of annually averaged global dengue epidemic potential (rVc). Differences in rVc are based on 30-year averages of temperature and daily temperature range. **A.** Differences between 1980–2009 and 1901–1930. **B.** Differences between 2070–2099 and 1980–2009. The mean value of rVc was averaged from five global climate models under RCP8.5, a scenario of high greenhouse-gas emission. The color bar describes the values of the rVc . (From J Liu-Helmerson et al: Vectorial capacity of *Aedes aegypti*: effects of temperature and implications for global dengue epidemic potential. *PLoS ONE* 9:e89783, 2014 [doi:10.1371/journal.pone.0089783].)

accuracy in predicting current *I. scapularis* distribution and suggests substantial expansion of tick habitat and consequently of populations at risk for the diseases this tick transmits, particularly in Quebec, Iowa, and Arkansas, by 2080. Of note, some areas on the Gulf Coast may become less suitable for ticks by the end of the century.

WATERBORNE DISEASE

Outbreaks of waterborne disease are associated with heavy rainfall events. A review of 548 waterborne disease outbreaks in the United States found that 51% were preceded by precipitation levels above

the 90th percentile. Since 1900, most regions of the United States except the Southwest and Hawaii have experienced an increase in heavy downpours (Fig. 151e-9), with the greatest intensification of the water cycle in New England and Alaska. Climate models suggest that by 2100 daily heavy-precipitation events, which are defined as a cumulative daily amount that now occurs once every 20 years, will increase nationwide (Fig. 151e-10). This scenario may be from two to as much as five times more likely, depending on the extent of greenhouse gas emission reductions achieved early in the twenty-first century.

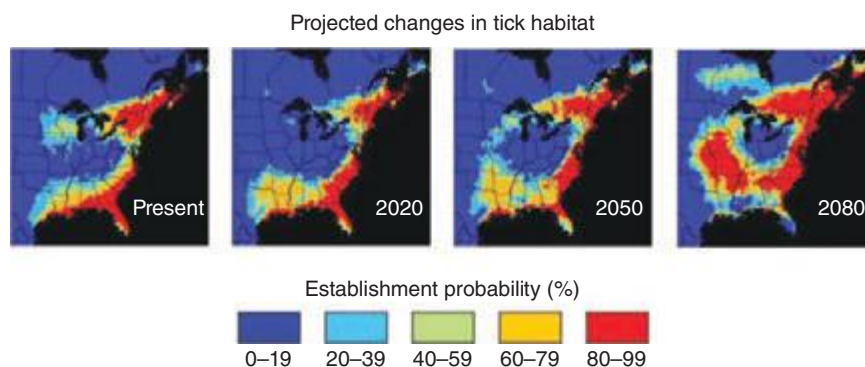


FIGURE 151e-8 Present and projected probability of establishment of *Ixodes scapularis*. (From U.S. National Climate Assessment 2014, adapted from JS Brownstein et al: Effect of climate change on Lyme disease risk in North America. *Ecohealth* 2:38, 2005.)

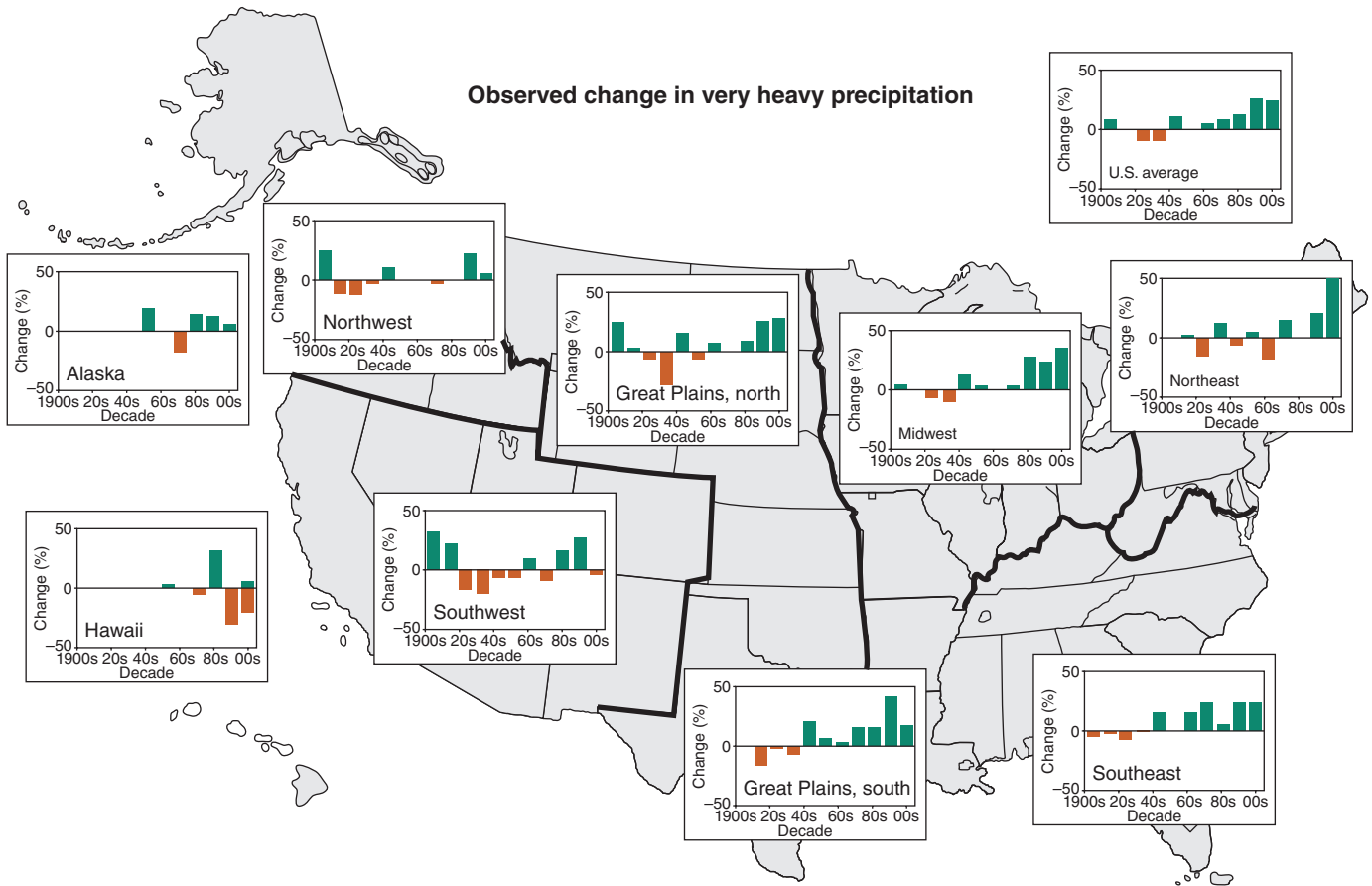


FIGURE 151e-9 Percentage changes in the annual amount of precipitation falling in very heavy events, defined as the heaviest 1% of all daily events from 1901 to 2012 for each region. Changes are relative to a 1901–1960 average for all regions except values for Alaska and Hawaii, which are relative to the 1951–1980 average. (From U.S. National Climate Assessment 2014, NOAA National Climate Data Center/Cooperative Institute for Climate and Satellites, North Carolina.)

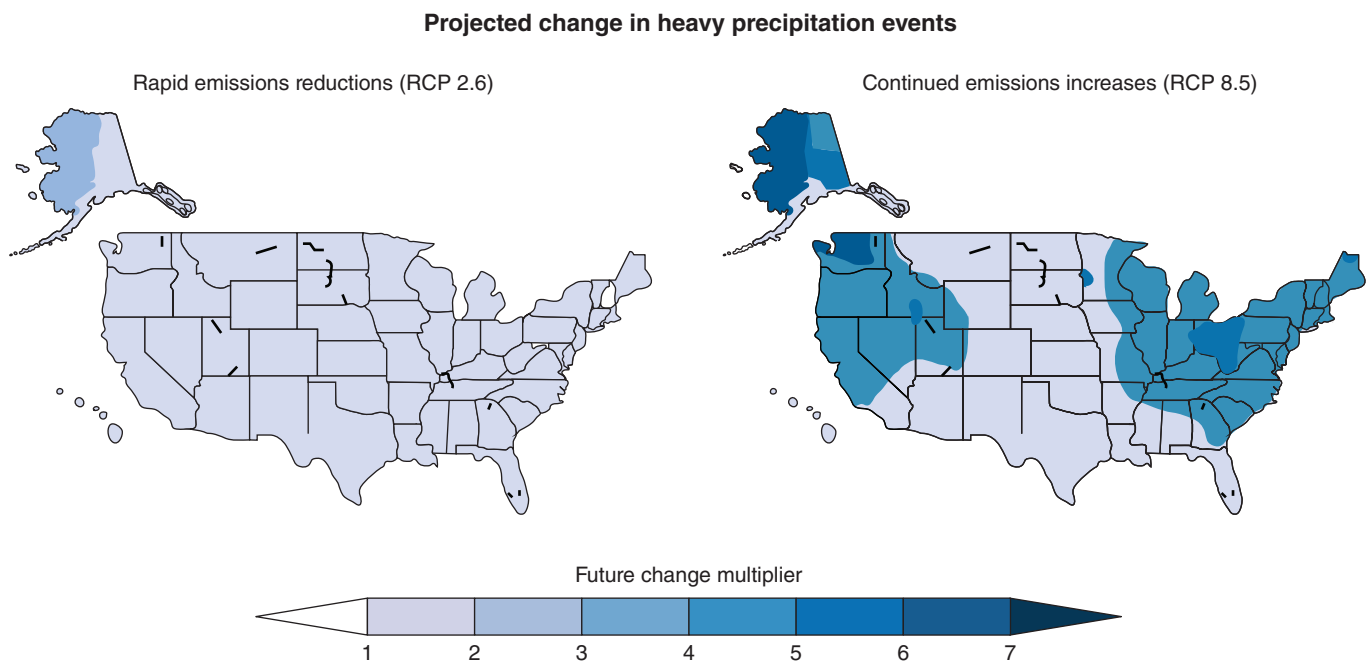


FIGURE 151e-10 Increased frequency of extreme daily precipitation events (defined as a daily amount that now occurs once in 20 years) by the latter part of the twenty-first century (2081–2100) compared to the frequency in the latter part of the twentieth century (1981–2000). A representative concentration pathway (RCP) describes a plausible climate future based on a net radiative forcing (e.g., 2.6 or 8.5) in 2100. (From U.S. National Climate Assessment 2014, NOAA National Climate Data Center/Cooperative Institute for Climate and Satellites, North Carolina.)

Most disease outbreaks after heavy precipitation occur through contamination of drinking-water supplies. While outbreaks related to surface-water contamination generally occur within a month of the precipitation event, disease outbreaks from groundwater contamination tend to occur ≥ 2 months later. According to a review of published reports of waterborne disease outbreaks, *Vibrio* and *Leptospira* species are the pathogens most commonly involved in the wake of heavy precipitation.

Combined Sewer Systems Roughly 40 million people in the United States and millions more around the world rely on combined sewer systems in which storm water and sanitary wastewater are conveyed in the same pipe to treatment facilities. These systems were designed on the basis of the nineteenth-century climate, in which heavy downpours were less frequent than they are today. The frequency of combined sewer overflows resulting in untreated sewage discharge, usually into freshwater bodies, has been increasing in cities worldwide. Overflows are associated with discharges of heavy metals and other chemical pollutants as well as a variety of pathogens. Outbreaks of hepatitis A, *Escherichia coli* O157:H7 infection, and cryptosporidial disease have been associated with sewer overflows in the United States.

Rising Temperatures and *Vibrio* Species Warmer temperatures favor proliferation of *Vibrio* species and disease outbreaks, as has been demonstrated in countries surrounding the Baltic Sea, Chile, Israel, northwestern Spain, and the U.S. Pacific Northwest. Around the Baltic Sea, outbreaks of *Vibrio* infection may be particularly likely because of faster warming near the poles and the sea's relatively low salt content. In 2004, a *Vibrio parahaemolyticus* outbreak occurred from consumption of Alaskan oysters. This pathogen was unknown in Alaskan oysters prior to this event and extended the known geographic range of the disease 1000 km northward.

ENSO-RELATED OUTBREAKS

In the past, El Niño events were used as a model to investigate the potential for extreme weather-related infectious disease epidemics occurring in association with climate change. Recent evidence has indicated that climate change itself may be strengthening El Niño events. These events tend to promote epidemic infections in certain regions (Fig. 151e-11).

Associations of El Niño with outbreaks of Rift Valley fever in eastern and southern Africa have been known since the 1950s. El Niño favors wet conditions suitable for the insect vectors of the disease in

these regions. Given the strong association between El Niño conditions and disease incidence, models have successfully predicted Rift Valley fever epidemics in humans and animals. In the 2006–2007 El Niño season, for example, outbreaks of Rift Valley fever were accurately predicted 2–6 weeks prior to epidemics in Somalia, Kenya, and Tanzania.

El Niño has had inconsistent associations with malaria incidence in African countries. Some of the strongest associations between El Niño and malaria have been identified in South Africa and Swaziland, where available data on incidence are relatively robust; however, even in these instances, the observed increased risk did not reach statistical significance. A stronger link to El Niño has been found in several studies done in South America. Research on malaria incidence in Colombia between 1960 and 2006 found that a 1°C temperature rise contributed to a 20% increase in incidence.

El Niño years are often associated with an increased incidence of dengue. Research on dengue outbreaks in Thailand from 1996 to 2005 revealed that 15–22% of the variance in monthly dengue disease incidence was attributable to El Niño. In South America, data on dengue outbreaks between 1995 and 2010 showed an increased incidence during the El Niño events of 1997–1998 and 2006–2007.

CLIMATE CHANGE, POPULATION DISPLACEMENT, AND INFECTIOUS DISEASE EPIDEMICS

For many reasons, including freshwater shortages, flooding, food shortages, and climate change-driven conflicts, climate change has and will continue to put pressure on human populations to move. Human migrations have long been associated with epidemic disease in the migrating populations themselves and in the communities in which they settle. The specific pathogens and patterns of disease that may appear after population migration relate to endemic diseases present in the migrant populations.

Large-scale migrations are common after extreme precipitation events. Hurricane Katrina, for instance, displaced about 1 million people from the U.S. Gulf Coast. Among Katrina refugees, outbreaks of respiratory, diarrheal, and skin diseases were most common. While attribution of a single weather event to increased greenhouse gas emissions is difficult, research can provide information on the likelihood of such events. It is expected, for example, that warming by 1°C increases the odds of a storm as strong as or stronger than Katrina by two- to sevenfold.

In the developing world, infectious disease outbreaks associated with population displacement due to extreme weather events may be

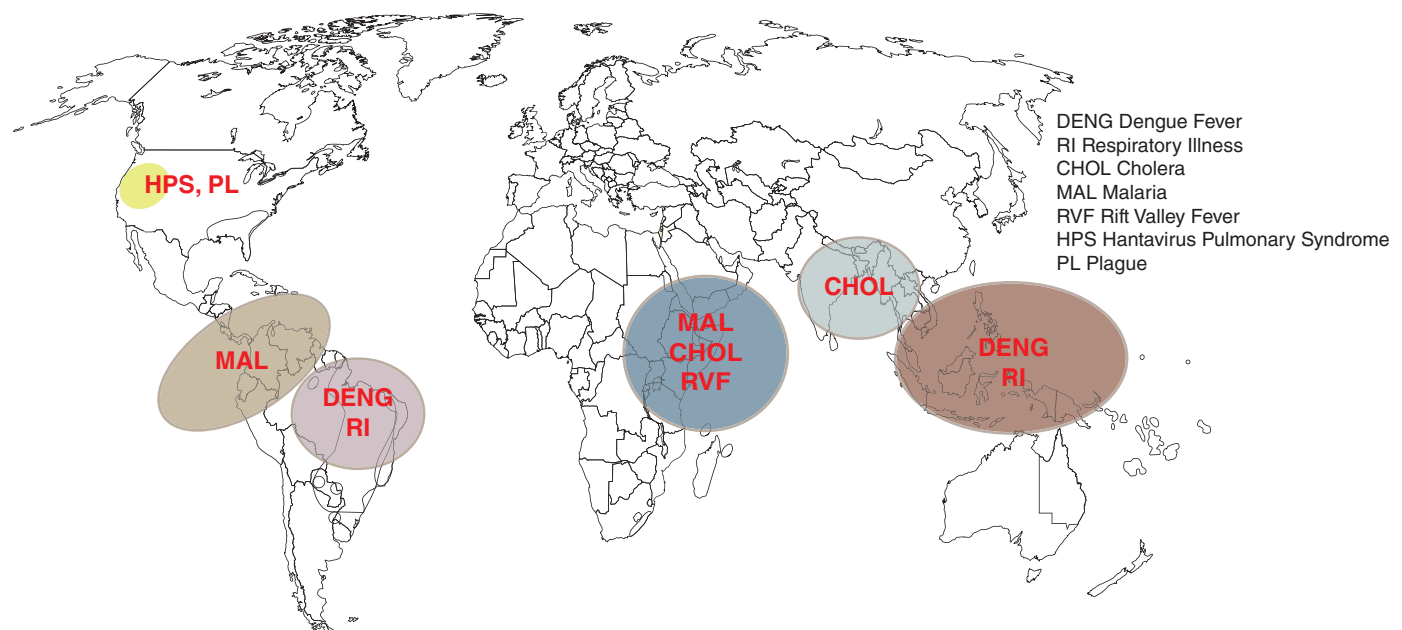


FIGURE 151e-11 Characteristic patterns of disease outbreaks associated with El Niño events, determined on the basis of 2006–2007 conditions. (From A Anyamba et al: *Developing global climate anomalies suggest potential disease risks for 2006–2007*. *Int J Health Geogr* 5:60, 2006.)

especially hard to detect and respond to. Mitigation of disease risk requires overlaying of climate-related migration risk with foci of disease epidemics.

A BROADER VIEW OF CLIMATE CHANGE AND HEALTH

Climate change has far-reaching implications for the distribution and spread of infectious diseases worldwide. However, the greatest disease burdens related to climate change may not be due to infections. Because climate change disrupts the foundations of health, such as access to safe drinking water and food, it has the potential to undermine progress against major existing health problems such as malnutrition. In addition, resource scarcity and climate instability are increasingly associated with conflicts. Scholars have argued that events related to climate change were a factor in the revolutions of the Arab Spring and the Syrian civil war.

The public health response to climate change entails both mitigation and adaptation measures. *Mitigation* represents primary prevention and involves the reduction of greenhouse gas emissions into the atmosphere. Although no clear safety threshold of greenhouse gas emissions has been agreed upon, national governments from the major industrialized countries have agreed to set a warming target of $<2^{\circ}\text{C}$ above preindustrial levels by 2050; the attainment of this goal will require reducing greenhouse gas emissions by 40–70% below 2010 levels. To date, no international agreement exists to facilitate this reduction. *Adaptation* represents secondary prevention and is aimed at reducing the harms associated with sea level rise, heat waves, floods, droughts, wildfires, and other greenhouse gas-driven events. The efficacy of adaptation is constrained by the challenges inherent in predicting the precise location, duration, and severity of extreme weather events and flooding related to sea level rise, among other considerations.

Andrew W. Artenstein

Wars are an unfortunate but apparently inescapable consequence of human socialization. In the past quarter-century alone, there have been dozens of major armed conflicts worldwide. Several of these wars have been multinational in scope and have involved the deployment of large numbers of ground forces from their home countries to distinct areas of conflict in developing portions of the world, such as southwest and central Asia (e.g., Iraq, Afghanistan) and Africa.

Troops who are deployed as combatants or in other military capacities on foreign soil are at risk of acquiring infectious diseases endemic to that region on the basis of intimate human and environmental exposures and their immunologic naiveté with regard to local endemic or enzootic pathogens. This risk is magnified by the crowded social conditions engendered by mass troop deployments, infrastructure destruction, and population displacement; it is further amplified by vulnerabilities in public health, such as the lapses in hygiene and sanitation that invariably accompany armed conflicts. The clinical spectrum of infectious illness acquired in this setting includes acute infections in the combat theater, acute infections with delayed symptoms, and chronic or relapsing infections. The latter two scenarios have the potential to cause illness in veterans returning from foreign wars.

The impact of acute infectious diseases of war, which were once a major cause of noncombat mortality, has significantly lessened in modern conflicts, largely because of the use of preventive vaccines and the early institution of antimicrobial therapy. Nonetheless, these acute diseases remain an important cause of morbidity in deployed military personnel (Table 152e-1). Many acute infectious diseases, such as influenza, meningococcal meningitis, hepatitis A, and adenoviral respiratory disease, can largely be prevented by routine vaccination of troops. Others, such as bacterial gastroenteritis and viral respiratory tract infections, continue to represent common causes of minor morbidity among deployed forces. The incidence of several other acute infections, such as malaria and dengue, can be favorably impacted—although not completely abrogated—by the use of chemoprophylaxis, personal protective measures, and vector control. Uncommonly, infections that have short incubation periods and are acquired just days before leaving a combat theater may become clinically manifest only upon the return of troops to their countries of origin. Such was the case in a cluster of African tick typhus that occurred during short-term U.S. troop deployments to Somalia and Botswana in the early 1990s. However, because most acute infections with brief clinical incubation periods are self-limited or responsive to treatment, they are not typically seen in returning war veterans and will not be addressed further in this chapter.

A number of infectious diseases acquired in a theater of military operations may become apparent in veterans only after they have returned to their home countries. Although their incidences have not been precisely defined, these complications of military service—manifesting clinically with either acute or chronic signs and symptoms—may cause significant morbidity in affected veterans and in some settings may endanger public health through secondary transmission or contamination of the blood supply. Of a sample of nearly 53,000 U.S. veterans of the Persian Gulf War (1990–1991), 7% were diagnosed with an infectious disease in the aftermath of the war; no excess risk of mortality was observed over a 6-year follow-up comparison with nondeployed veterans.

This chapter focuses on infectious diseases that have occurred or have been a source of concern in returning veterans of foreign wars over the past quarter-century. During this period, several pathogens have been associated with disease in this population, as discussed below. Some pathogens have been associated with only rare case reports in war veterans, and some, given their epidemiology, may pose

TABLE 152e-1 ACUTE INFECTIOUS DISEASES OF WAR THAT HAVE BRIEF CLINICAL INCUBATION PERIODS AND THEREFORE ARE LIKELY TO CAUSE SYMPTOMATIC ILLNESS IN MILITARY PERSONNEL DURING DEPLOYMENT

Bacterial	
Meningococcal meningitis	
Gastroenteritis (enterotoxigenic or enteroinvasive <i>Escherichia coli</i> , <i>Shigella</i> , <i>Salmonella</i> , <i>Campylobacter</i>)	
Typhoid fever	
Cholera	
Typhus (epidemic, murine, scrub)	
African tick-bite fever	
Leptospirosis	
Sexually transmitted diseases (gonorrhea, chlamydial infection, genital herpes, genital warts, chancroid, syphilis)	
Combat wound infections	
Q fever	
Viral	
Norovirus gastroenteritis	
West Nile virus infection	
Crimean-Congo hemorrhagic fever	
Influenza	
Adenoviral respiratory disease	
Viral upper respiratory tract infections	
Dengue	
Alphavirus infections (e.g., Chikungunya, O'nyong-nyong, or Sindbis virus)	
Tickborne encephalitis	
Sandfly fever	
Hantavirus syndromes	
Parasitic	
Malaria	
Gastroenteritis (cryptosporidiosis, giardiasis, amebiasis)	
Ophthalmomyiasis	

a risk in future conflicts. In general, it is practical to classify infections with delayed signs and symptoms related to prolonged incubation periods or significant clinical latency in terms of their potential to manifest clinically as acute illnesses or chronic/relapsing diseases. Table 152e-2 provides details regarding the epidemiology, clinical characteristics, diagnosis, therapy, and prevention of infectious diseases of concern in returning war veterans. Figure 152e-1 illustrates a differential diagnostic approach—based on prominent signs or symptoms—to suspected infections in this population.

ACUTE INFECTIOUS DISEASES WITH DELAYED CLINICAL PRESENTATIONS

Malaria (See Chap. 248) Malaria, which is due to infection with *Plasmodium* species of protozoa, has historically caused significant battlefield morbidity and lost duty time among armed forces; these phenomena have been affirmed by recent U.S. military experiences in Africa and central Asia. Because of its worldwide prevalence and its pathophysiology, malaria remains an important cause of infection both during military operations and in returning war veterans.

The risk of malaria is exacerbated by several factors inherent to war: inadequate shelter promoting increased troop exposure to vectors; abeyance of governmental programs for vector control; and ecologic changes leading to an increased vector presence in the contested environment. Because of the complex life cycle of the parasite and the predilection of *P. vivax* and *P. ovale* to remain latent in their liver stages of development for prolonged periods, malaria in foreign-stationed troops may become clinically apparent only after their return home. In the aftermath of the Vietnam War, more than 13,000 cases of malaria—the vast majority due to *P. vivax*—were imported into the United States. Of the 7683 Soviet troops diagnosed with *P. vivax*

TABLE 152e-2 INFECTIOUS DISEASES ASSOCIATED WITH DELAYED CLINICAL MANIFESTATIONS THAT MAY PRESENT WITH ACUTE, CHRONIC, OR RELAPSING COURSES IN VETERANS RETURNING FROM RECENT FOREIGN WARS

Disease	Pathogen	Global Epidemiology	Mode of Transmission	Clinical Incubation Period	Symptoms/Signs	Diagnosis	Treatment	Prevention	Chapter
Acute Clinical Course									
Malaria	<i>Plasmodium falciparum</i> , <i>P. malariae</i> , <i>P. vivax</i> <i>P. ovale</i>	Asia, Africa, South and Central America As above; limited in Africa; foci in Asia	Vector (mosquito)	7–339 days; rare reports of <i>P. falciparum</i> 2 years and <i>P. vivax</i> 5 years after exposure	Fever (possibly cyclical, every 48 or 72 h), rigors, headache, night sweats, weakness, altered mental status	Serial Giemsa-stained blood smears demonstrating intraerythrocytic parasites; rapid antigen capture assay	Chloroquine (if acquired in chloroquine-sensitive area); mefloquine or atovaquone + proguanil or quinine + doxycycline; primaquine for <i>P. vivax</i>	Bed nets, insect repellents, vector control, chemoprophylaxis	248
Hepatitis A	Hepatitis A virus (HAV)	Worldwide: highest prevalence in developing world	Fecal-oral	15–50 days	Fever, malaise, anorexia, nausea, abdominal pain, jaundice, elevated aminotransferases	Serology: anti-HAV IgM	Supportive; no specific treatment	Food and water hygiene; passive immunization with pooled immunoglobulin; active immunization with killed vaccine	360
Hepatitis B	Hepatitis B virus (HBV)	Worldwide; highest prevalence in developing world	Percutaneous or sexual exposure to infected body fluids	45–180 days	Malaise ± fever, anorexia, nausea, vomiting, abdominal pain, jaundice, elevated aminotransferases	Serology: HBV surface antigen positive, IgM anti-HBV core antibody	Supportive; no specific therapy for acute disease	Personal protective measures; screening of blood supply; use of sterile needles; immunization with recombinant vaccine	360
Hepatitis E	Hepatitis E virus (HEV)	Asia, north and west Africa, Mexico	Fecal-oral	15–64 days	Fever, malaise, anorexia, nausea, abdominal pain, jaundice, elevated aminotransferases	Serology: anti-HEV IgM	Supportive; no specific therapy for acute disease	Food and water hygiene	360
Rabies	Rhabdovirus (member of genus <i>Lyssavirus</i>)	Worldwide: in developing world, domestic and feral animals; in developed world, wild animals	Exposure to saliva of infected animals; rarely, airborne transmission in bat roosting caves	9 days to >1 year (rare)	Heralded by headache, fever, apprehension, paresthesias near site of exposure; progression to paresis, muscle spasm, dysphagia, hydrophobia, delirium, seizures	Direct fluorescent antibody staining of brain tissue (animal or human) or skin on back of neck; RT-PCR on CSF, saliva, tissue	Supportive; no specific therapy for acute disease	Before exposure: active vaccination. After exposure: treatment to prevent active disease (wound care; passive immunization with human rabies immune globulin infiltrated into site of exposure; active immunization with vaccine)	232
Leptospirosis	<i>Leptospira interrogans</i> serovars	Worldwide	Contact of skin or mucous membranes with fresh-water or soil contaminated with urine of infected animals	10–21 days	Fever, headache, myalgia, conjunctival suffusion, possibly in association with jaundice, meningitis, mental status changes, hemolysis, hepatitis, myocarditis, and pneumonia. Illness may be biphasic and become chronic.	Serology: IgM antibodies; organism isolation in culture from blood, urine, or CSF during acute phase of illness	Oral doxycycline for mild disease; parenteral ceftriaxone or penicillin for severe disease	Personal protective measures when exposure cannot be avoided; doxycycline prophylaxis in high-risk exposures	208

Scrub typhus	<i>Orientia tsutsugamushi</i>	Central, southeastern, and eastern Asia	Vector (chiggers [mite larvae])	6–21 days	Fever, headache, myalgia, arthralgia, conjunctival injection, malaise, cough, transient maculopapular rash, eschar at site of mite attachment	Serology: Weil-Felix test detects cross-reacting antibodies to <i>Proteus mirabilis</i> OX-K.	Doxycycline or chloramphenicol	Personal protective measures; doxycycline prophylaxis	211
Chronic or Relapsing Clinical Course									
Cutaneous leishmaniasis	Old World: <i>Leishmania major</i> , <i>L. tropica</i> , <i>L. infantum-donovani</i> New World: <i>L. mexicana</i> complex, <i>L. braziliensis</i> complex	Southwest and central Asia, China, Africa, Central and South America	Vector (<i>Phlebotomus</i> spp. sandfly)	2–8 weeks (up to 18 months)	Single or multiple chronic, painless, nodular or ulcerative skin lesions ± eschar near inoculation sites on head and limbs, associated with spontaneous healing and scarring after 5–12 months; rare chronic or relapsing forms	Parasites in Wright-Giemsa-stained skin scraping, slit skin smear, or biopsy specimen; culture or PCR for speciation	Vigilant monitoring for spontaneous resolution; pentavalent antimony; oral azoles	Personal protective measures (sandflies can penetrate mosquito nets; permethrin coating enhances efficacy); vector and animal reservoir control	251
Visceral leishmaniasis (kala-azar)	<i>L. infantum-donovani</i>	Southwest and central Asia, Brazil, India, China, Africa	Vector (<i>Phlebotomus</i> spp. sandfly)	2–14 months (rarely up to 2 years)	Chronic fever, cachexia, anorexia, weight loss, pancytopenia, hepatosplenomegaly	Parasites in tissue (bone marrow, liver, lymph node, spleen) on histopathologic staining, culture, or PCR; rK39 serologic assay	Liposomal amphotericin B, pentavalent antimony, miltefosine	Personal protective measures, vector and animal reservoir control	251
Tuberculosis	<i>Mycobacterium tuberculosis</i>	Worldwide	Inhalation of aerosols from individuals with active pulmonary tuberculosis	1 month to many years	Fever, weight loss, night sweats, cachexia, cough, hemoptysis, pulmonary infiltrates	Sputum acid-fast smears, mycobacterial culture, nucleic acid amplification	Multidrug therapy (number depends on risk of drug resistance) including combination of isoniazid, rifampin, pyrazinamide, ethambutol, and others	Personal protective measures; chemoprophylaxis for skin test conversion	202
Q fever	<i>Coxiella burnetii</i>	Worldwide	Inhalation of aerosols from infected secretions of livestock herd or parturient animal; ingestion of raw milk	2–5 weeks for acute disease; chronic manifestations may present indolently over many years	Acute: undifferentiated febrile illness, atypical pneumonia Chronic: granulomatous hepatitis, culture-negative endocarditis, osteomyelitis (including vertebral), fever of unknown origin	Serology: complement-fixation antibody titer to phase I antigen	Doxycycline (with hydroxychloroquine for endocarditis)	Avoidance of unpasteurized products	211
Brucellosis	<i>Brucella abortus</i> , <i>B. melitensis</i> , <i>B. suis</i>	Worldwide; especially prevalent in southwest and central Asia, India	Inhalation of infected animal materials, ingestion of raw milk, direct animal contact	5 days to months	Skeletal, genitourinary, or pulmonary disease; chronic meningitis	Organism isolation from blood, bone marrow; <i>Brucella</i> -specific antibody	Doxycycline + rifampin or streptomycin	Personal protective measures, avoidance of unpasteurized dairy products	194e

(Continued)

TABLE 152e-2 INFECTIOUS DISEASES ASSOCIATED WITH DELAYED CLINICAL MANIFESTATIONS THAT MAY PRESENT WITH ACUTE, CHRONIC, OR RELAPSING COURSES IN VETERANS RETURNING FROM RECENT FOREIGN WARS (CONTINUED)

Disease	Pathogen	Global Epidemiology	Mode of Transmission	Clinical Incubation Period	Symptoms/Signs	Diagnosis	Treatment	Prevention	Chapter
Melioidosis	<i>Burkholderia pseudomallei</i>	Southeast Asia	Percutaneous inoculation from environmental sources; respiratory route, including possible inhalation in dusts raised by helicopter blades during military operations	1–21 days; may become clinically apparent or relapse years after exposure	Chronic, suppurative skin and soft tissue infections; osteomyelitis; chronic cavity pneumonia; septicemia	Isolation of organism in culture	Ceftazidime or carbapenem; possible use of trimethoprim-sulfamethoxazole as eradication therapy	Personal protective measures and wound hygiene	189
Echinococcal (hydatid) disease	<i>Echinococcus granulosus</i> ; <i>E. multilocularis</i>	Worldwide (<i>E. granulosus</i>); north Asia (<i>E. multilocularis</i>)	Ingestion of eggs from feces-contaminated soil	Months to years	Symptoms related to progressive enlargement of cystic lesions in liver, lungs, brain, bone	Immunologic assay (ELISA) in setting of consistent imaging picture	Albendazole therapy followed by surgical excision (with or without instillation of cysticidal agent)	Animal hygiene and control measures	260
Schistosomiasis	<i>Schistosoma haematobium</i> (urinary tract); <i>S. mansoni</i> , <i>S. japonicum</i> (biliary tract); <i>S. mekongi</i>	<i>S. mansoni</i> : Africa, southwest Asia, South America; <i>S. japonicum</i> : Far East; <i>S. mekongi</i> : southeast Asia; <i>S. haematobium</i> : Africa, southwest Asia	Freshwater via exposure to snail intermediate hosts	14–84 days for acute manifestations (Katayama fever); chronic manifestations over years	Katayama fever: fever, headache, malaise, dry cough, dyspnea, transient urticarial rash Chronic: Terminal hematuria, dysuria, frequency (<i>S. haematobium</i>); abdominal pain, eosinophilia, hepatosplenomegaly, sequelae of portal hypertension (<i>S. mansoni</i> , <i>S. japonicum</i>)	Urine or fecal examination for eggs; bladder biopsy; serology for antibodies to schistosomes	Praziquantel	Personal protective measures in freshwater; snail control via molluscicides	259
Histoplasmosis	<i>Histoplasma capsulatum</i>	Africa, Americas, Far East	Inhalation of airborne environmental conidia	3–17 days for acute manifestations; chronic manifestations over months to years	Chronic, systemic disease with fever, weight loss, fatigue, cytopenias, hepatosplenomegaly; chronic pulmonary disease similar to tuberculosis	Fungal visualization in stained tissue specimens; serum or urine antigen detection	Itraconazole or ketoconazole	Personal protective measures in selected high-risk environments, such as caves, pigeon roosts, chicken houses	236
Extraintestinal amebiasis	<i>Entamoeba histolytica</i>	Worldwide	Fecal-oral	11–21 days	Fever, abdominal pain, watery diarrhea; right-upper-quadrant tenderness, hepatomegaly	Stool antigen assay; serum amebic antibodies in liver abscess	Metronidazole + paromomycin or diloxanide furoate; dehydroemetine + paromomycin or diloxanide furoate	Food and water hygiene	247
Giardiasis	<i>Giardia lamblia</i>	Worldwide	Fecal-oral	7–10 days	Chronic diarrhea, abdominal pain, bloating, nausea, flatulence due to postinfectious irritable bowel syndrome or protein-losing enteropathy; postinfection fatigue	Fecal, string test, or duodenal aspirate microscopy; stool antigen assay	Metronidazole for ongoing infection; no specific antiparasitic therapy for postinfectious syndromes	Food and water hygiene	254

Cryptosporidiosis	<i>Cryptosporidium</i> spp.	Worldwide	Fecal-oral	3–6 days	Symptoms as above; chronic watery diarrhea, with or without fever, abdominal pain, nausea in immunocompromised hosts	Fecal microscopy or intestinal biopsy; stool antigen assay	No specific antiparasitic therapy for postinfectious syndromes	Food and water hygiene	254
Strongyloidiasis	<i>Strongyloides stercoralis</i>	Tropical and subtropical climates	Fecal-oral as initial route; persistent infection can result from autoinfection in human host	11–28 days	Abdominal pain, persistent diarrhea, urticaria; disseminated disease can cause wasting, pulmonary symptoms, eosinophilia (predilection for immunocompromised hosts)	Stool antigen detection assay; serology	Ivermectin; thiabendazole or albendazole as second-line alternatives	Personal protective measures, including wearing shoes in endemic areas (infective larvae penetrate intact skin)	257
Sandfly fever (convalescence)	Phleboviruses	Africa, Asia, South and Central America	Vector (<i>Phlebotomus</i> spp. sandfly)	Weeks to months for convalescent symptoms	Depression, fatigue, generalized weakness	Serology	No specific therapy	Personal protective measures; vector control	233
Relapsing fever	<i>Borrelia recurrentis</i> (louse-borne, epidemic), <i>Borrelia</i> spp. (tick-borne, endemic)	Worldwide	Vector (body louse; soft tick)	4–18 days initially, with relapses after 7- to 10-day intervals	Recurrent episodes of fever, rigors, diaphoresis, headache, myalgias, arthralgias, asthenia lasting 3–6 days and alternating with symptom-free periods	Spirochetes on stained peripheral-blood smear during febrile episodes	Tetracycline or erythromycin; antibiotic treatment may lead to Jarisch-Herxheimer reaction with fever, rigor, hypotension within 2 h of initiation	Personal protective measures; vector control	209
Brill-Zinsser disease	<i>Rickettsia prowazekii</i>	Worldwide	Vector (body louse)	Recrudescence years after primary episode of epidemic typhus	Mild febrile illness with systemic symptoms and macular eruption	Serology	Doxycycline or chloramphenicol	Vector control; personal protective measures; appropriate treatment of initial typhus episode	211
Chronic wound infection	<i>Acinetobacter</i> spp., other gram-negative organisms; <i>Staphylococcus aureus</i> , including MRSA; invasive molds (<i>Aspergillus</i> , <i>Fusarium</i> , <i>Mucor</i> , <i>Absidia</i> spp.); atypical mycobacteria (<i>M. chelonae</i> , <i>M. abscessus</i>) associated with draining sinuses	Worldwide	Inoculation via combat wounds and penetrating injury	Weeks to months; progression of acute infection	Chronic pain, swelling, ± serosanguineous or purulent drainage of infected site with or without constitutional symptoms/signs	Culture of tissue	Guided by results of cultures and in vitro antibiotic susceptibility testing; carbapenem ± amikacin as empirical therapy for multidrug-resistant <i>Acinetobacter</i> ; colistin if resistant	Adequate initial wound debridement and treatment of acute soft-tissue infection; removal of foreign bodies; strict adherence to infection control precautions to prevent nosocomial infection	Miscellaneous

(Continued)

TABLE 152e-2 INFECTIOUS DISEASES ASSOCIATED WITH DELAYED CLINICAL MANIFESTATIONS THAT MAY PRESENT WITH ACUTE, CHRONIC, OR RELAPSING COURSES IN VETERANS RETURNING FROM RECENT FOREIGN WARS (CONTINUED)

Disease	Pathogen	Global Epidemiology	Mode of Transmission	Clinical Incubation Period	Symptoms/Signs	Diagnosis	Treatment	Prevention	Chapter
Chronic osteomyelitis	<i>Acinetobacter baumannii</i> and other gram-negative pathogens; <i>S. aureus</i> , including MRSA	Worldwide	Foreign bodies (projectiles); contiguous spread from soft tissue infection	Weeks to months	Chronic pain, swelling of infected site with or without constitutional symptoms/signs	Culture of bone biopsy	Treatment as above	Adequate initial wound debridement and treatment of soft tissue infection; removal of foreign bodies; strict adherence to infection control precautions to prevent nosocomial infection	158
Leprosy	<i>Mycobacterium leprae</i>	Southeast and south Asia, tropical Africa, Brazil	Skin, mucous membrane, or fomite contact	4–10 years	Skin lesions, cutaneous hypoesthesia, peripheral-nerve enlargement	Acid-fast bacilli in skin biopsy	Dapsone + rifampin ± clofazimine	Multidrug therapy to prevent chronic sequelae	203
Filariasis	<i>Wuchereria bancrofti</i> , <i>Brugia malayi</i> , <i>Brugia timori</i>	<i>Wuchereria</i> : tropical and subtropical areas of Africa, Asia, Americas; <i>Brugia</i> : southeast and south Asia	Vector (mosquito)	5–18 months	Pain and swelling of arms, legs, genitalia; headache, nausea, fatigue, eosinophilia	Identification of microfilariae on nocturnal, Giemsa-stained peripheral-blood smear	Diethylcarbamazine or ivermectin to possibly reduce microfilarial counts in blood; no proven effective drug therapy for lymphatic disease	Vector control, personal protective measures	258
Visceral larva migrans	<i>Toxocara canis</i> , <i>T. cati</i>	Worldwide	Ingestion of eggs from feces-contaminated soil	Weeks to months	Eosinophilia, rash, fever, pneumonitis, chronic abdominal pain	Serology	No proven effective therapy; usually self-limited	Food and personal hygiene	256
Hepatitis C	Hepatitis C virus (HCV)	Worldwide	Bloodborne, less efficient sexual transmission	Years	Chronic hepatitis leading to cirrhosis in minority	Serology: HCV antibody	Combination antiviral therapy	Bloodborne precautions; condom use	362
HIV-1 infection	Human immunodeficiency virus type 1	Worldwide	Bloodborne, sexual	Years	Constitutional symptoms and signs, opportunistic infections	Serology, viral PCR	Combination highly active antiretroviral therapy	Bloodborne precautions; condom use	226
HTLV-I infection	Human T cell lymphotropic virus type I	Foci of infection in Central and South America, sub-Saharan Africa	Bloodborne, sexual	Years	T cell leukemia/lymphoma, chronic progressive myelopathy/tropical spastic paraparesis	Serology, viral PCR	No specific antiviral therapy	Bloodborne precautions; condom use	225e

Abbreviations: CSF, cerebrospinal fluid; MRSA, methicillin-resistant *S. aureus*; PCR, polymerase chain reaction; RT-PCR, reverse transcription PCR.

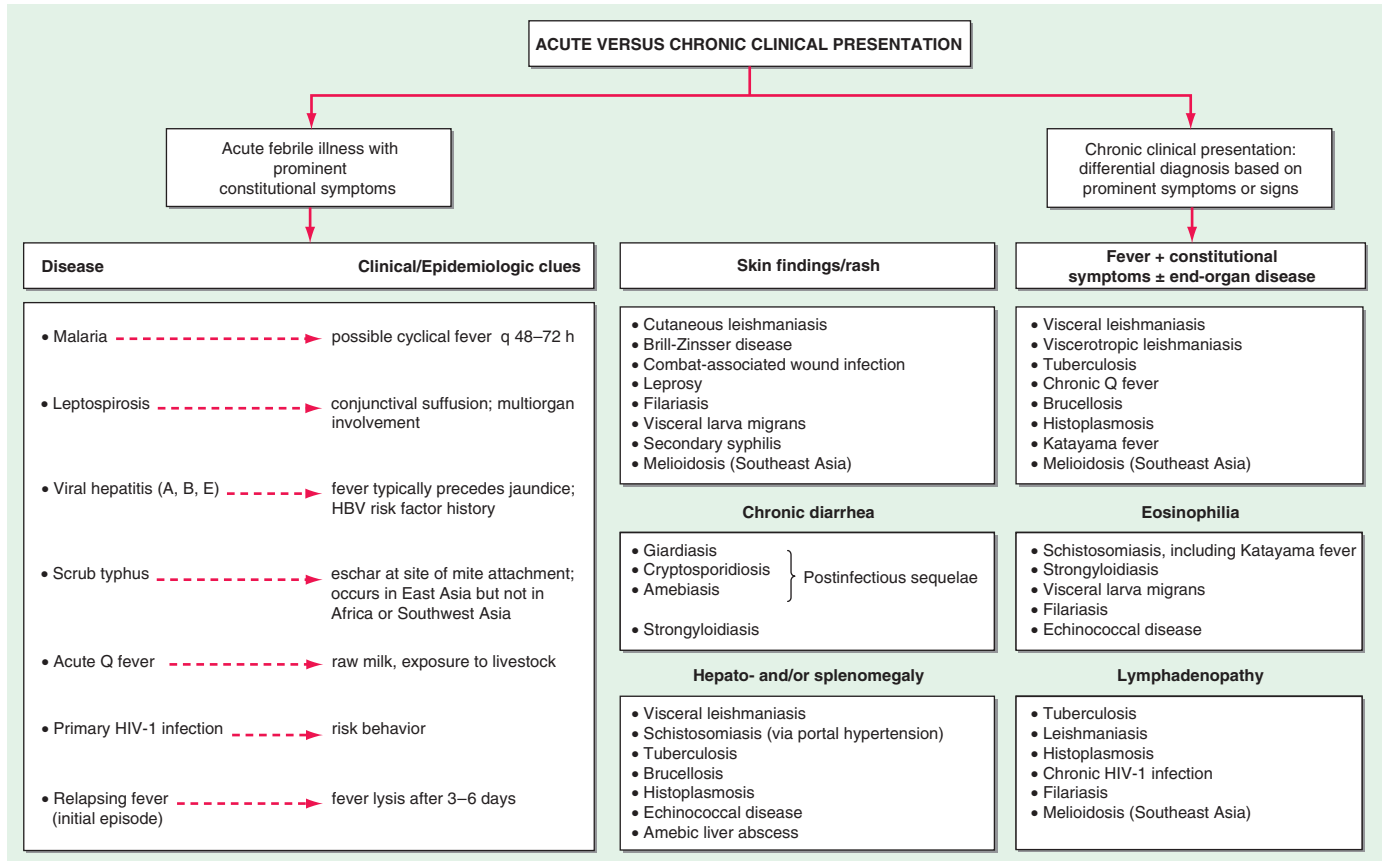


FIGURE 152e-1 Syndromic approach to the differential diagnosis of suspected infection in a veteran who has returned from a foreign war in southwest Asia, central Asia, or Africa at least 2 weeks prior to clinical presentation. HBV, hepatitis B virus.

malaria acquired during the U.S.S.R.'s war in Afghanistan in the 1980s, 76% developed clinical manifestations >1 month after their return to the U.S.S.R., with some cases developing as long as 3 years later.

Imported malaria with prolonged clinical latency periods remains a problem among veterans returning from wars in endemic areas. In a cluster of 112 cases of imported disease in U.S. Marines returning to the United States from deployment to Somalia in 1993, falciparum malaria was diagnosed as late as 12 weeks after return; some cases due to *P. vivax* were diagnosed after an additional 2 months. In an outbreak of imported *P. vivax* malaria among U.S. Army Rangers deployed to Afghanistan in 2002, the median time to diagnosis was nearly 8 months after return.

Although malaria is largely preventable through the combined use of vector control, personal protective measures (e.g., bed nets, insect repellent, long sleeves, permethrin-treated clothing), and chemoprophylaxis, nonadherence to these measures and/or to chemotherapy (including terminal primaquine prophylaxis to eradicate the liver stage of *P. vivax*) appears to be responsible for the majority of cases in recent U.S. military experiences. However, there is also evidence to support chemoprophylaxis failures in a small subgroup of cases of *P. falciparum* and *P. vivax* malaria acquired during U.S. deployments to Somalia. Thus, imported malaria in returning war veterans remains possible despite appropriate adherence to chemoprophylaxis.

Viral Hepatitis (See Chap. 360) The incidence of viral hepatitis, once a major scourge of military campaigns and their aftermath, has declined considerably over the past half-century of military engagements. Although more than 115,000 cases of viral hepatitis, most due to hepatitis A virus, were reported among Soviet troops serving in Afghanistan in the 1980s, only rare reports of hepatitis A and B were noted during the massive, short-term deployment of U.S. troops to the Persian Gulf in the early 1990s. Hepatitis A and E, endemic in many parts of the developing world, present clinically as acute infections transmitted by the fecal-oral route and can be largely controlled with interventions

practiced widely among military forces from industrialized countries: appropriate food and water hygiene and pre-deployment vaccination against hepatitis A. Hepatitis B contamination of serum-stabilized yellow fever vaccine caused a large outbreak of the disease among U.S. forces during World War II; such events are currently unlikely, given the use of modern virus-inactivation techniques in vaccine manufacturing. Despite their potential—as a consequence of their long clinical incubation and latency periods—to cause disease in returning veterans, hepatitis B and hepatitis C have been acquired relatively rarely in theater because of risk factor mitigation: routine drug testing of modern armies and screening of the blood supply to exclude viral contamination.

Rabies (See Chap. 232) Deployed soldiers are often in close contact with feral dogs and other potentially rabid animals in both urban and remote combat environments. In the period 2001–2010, 643 animal bites were documented during medical encounters among U.S. forces in the combat theaters of southwest and central Asia; the majority of these potential exposures were dog bites. Of bitten personnel, 18% received rabies postexposure prophylaxis. Recently, a U.S. soldier developed rapidly progressive signs and symptoms of rabies 8 months after a dog bite in Afghanistan and died within 17 days of illness onset. This case, which represents the first rabies death in nearly 40 years in a member of the U.S. military who acquired the infection overseas, reinforces both the lethality of the disease and the need to practice vigilant exposure precautions among deployed military personnel and to administer postexposure prophylaxis to troops who are bitten by animals in high-risk areas.

CHRONIC OR RELAPSING INFECTIOUS DISEASES ACQUIRED IN THE THEATER OF WAR

Leishmaniasis (See Chap. 251) Various forms of leishmaniasis may be acquired during military deployment and may present with myriad chronic clinical manifestations in war veterans. Because the protozoal

pathogens (*Leishmania* species) are endemic throughout much of southwest and central Asia, their associated diseases have occurred in veterans returning from several recent conflicts there. The widespread distribution of various species of *Leishmania* elsewhere throughout the developing world suggests that leishmaniasis may complicate future conflicts as well.

Leishmaniasis may present clinically as cutaneous, mucocutaneous, or visceral disease; all forms are transmitted to humans by the bite of infected phlebotomine sandflies via zoonotic (small mammal) reservoirs. In rare circumstances, infection may be transmitted through blood transfusion. Transmission to humans is enhanced by factors that bring them into close proximity to animal reservoirs, such as life in dense, mobile populations; disruption of ecologic niches; and infrastructural breakdown. All these factors are common sequelae of war.

At least 1300 cases of cutaneous leishmaniasis caused by *L. major* or *L. tropica* have been diagnosed in American soldiers deployed to Iraq and Afghanistan over the past decade; however, the actual burden of infection may be higher due to underreporting, as lesions spontaneously resolve in many cases. The infection manifests clinically as one or more chronic, painless skin ulcers or nodules that may persist for 6–12 months. Rarely, lesions of cutaneous leishmaniasis may disseminate locally or diffusely.

Visceral disease (*kala-azar*) is typically caused by *L. donovani* and may be life-threatening. There have been at least five confirmed reports of U.S. veterans returning from recent deployments with classic visceral leishmaniasis associated with chronic fever, weight loss, pancytopenia, hypergammaglobulinemia, and organomegaly. As systemic leishmanial infection is known to manifest clinically years after exposure and may recrudescence if host immunity wanes due to unrelated causes, it remains possible that additional cases may yet surface.

Chronic Diarrhea Although acute bacterial gastroenteritis remains a major noncombat cause of morbidity and duty days lost during troop deployments, chronic illness is unusual. However, selected bacterial and parasitic enteric pathogens may cause chronic infections or illnesses in returning veterans. Although such infections have been uncommonly noted in recent deployments, they pose potential threats in future wars because of their worldwide prevalence.

Giardiasis (Chap. 254), amebiasis (Chap. 247), and cryptosporidiosis (Chap. 254), which usually cause self-limited protozoal gastroenteritis in immunocompetent hosts, may result in persistent symptoms in immunocompromised populations or when complicated by secondary illness. *Giardia* infection has been associated with chronic diarrhea due to postinfectious irritable bowel syndrome and with chronic signs and symptoms of systemic illness in association with postinfection fatigue or protein-losing enteropathy. Cryptosporidiosis also may cause chronic diarrhea or malabsorptive syndromes in immunocompromised individuals. Amebic infection of the colon may be associated with several serious complications, including perforation, fistulae, and obstruction; extraintestinal spread of amebiasis may result in hepatic invasion leading to abscess formation.

Systemic Illness Due to Enteric Pathogens Certain helminthic infections are endemic in many parts of the developing world and may pose continuing risks to infected military personnel after their return. Larvae of the intestinal nematode *Strongyloides stercoralis* (Chap. 257) either may be passed in the feces and develop into the infective stage in the external environment or may persist in the human small intestine and initiate new infective cycles in a process known as *autoinfection*. Autoinfection with *Strongyloides* may result in chronic clinical manifestations such as pruritus, rash, abdominal pain, weight loss, diarrhea, and eosinophilia. In immunocompromised hosts, chronic strongyloidiasis may cause a life-threatening hyperinfection syndrome, likely triggered by high parasite burdens and resulting in a multiorgan, systemic illness consistent with severe inflammatory response syndrome. In some cases, *Strongyloides* hyperinfection syndrome may be complicated by gram-negative sepsis or meningitis related to bacterial seeding from parasitic involvement of the lungs or gastrointestinal tract. Although not described in association with recent wars, chronic strongyloidiasis was an uncommon phenomenon affecting a small number

of World War II and Vietnam War veterans; one study estimated that there were up to 400 affected individuals still living in Great Britain. The pathogen may pose a risk to troops deployed in the future to tropical and subtropical regions of the world where the parasite is endemic.

Chronic schistosomiasis (Chap. 259) results from intravascular infection by trematode parasites whose larval forms penetrate the skin of humans through contact with freshwater inhabited by the snail intermediate host. The pathogens are widely distributed throughout large portions of the developing world. A chronic inflammatory response in the portal circulation to the eggs of *S. mansoni* and *S. japonicum* leads to fibrosing disease in the liver and eventually to cirrhosis. Similar pathophysiologic events occur in the vascular plexus of the human genitourinary tract in response to chronic *S. haematobium* infection, leading to fibrosing changes in the human bladder and ureters—a precursor to bladder cancer. Rarely, individuals with chronic schistosomiasis develop a syndrome of persistent or relapsing bacteremia with *Salmonella typhi*, which is the etiologic agent of typhoid fever and is not otherwise a typical infectious cause of chronic disease in veterans.

Other Chronic Infections/Syndromes Awareness of the potential threat of troop exposure to agents of biological warfare (Chap. 261e) has been heightened over the past two decades by revelations regarding Iraq's state-sponsored chemical weapons program in the 1990s, the known broad availability of such technology, and escalations of global and geopolitical conflicts as well as of international acts of terrorism. Most of the high-risk pathogens posing a threat of bioterrorism cause acute clinical manifestations; however, selected agents, such as those causing Q fever and brucellosis, may result in chronic diseases, whether exposure is natural or intentional. Isolated cases of naturally acquired Q fever and brucellosis have been reported in recent U.S. veterans of the wars in Iraq and Afghanistan. To date, there has been no confirmed evidence of infections related to exposure to biological weapons in returning war veterans.

HIV-1 infection (Chap. 226), ubiquitous throughout the world, continues to pose a potential bloodborne and sexually transmitted risk to soldiers engaged in armed conflict in high-prevalence areas. Several reports describe war veterans returning to their countries of origin harboring HIV-1; in some of these cases, novel viral genotypes have been imported into the population. Tuberculosis (Chap. 202) also is endemic throughout much of the developing world and is prevalent in several areas of recent multinational conflicts. Although there is no evidence that active, chronic tuberculosis has affected veterans of recent wars, the rate of tuberculin skin test conversion, which indicates new infections, was noted to be 2.5% among U.S. military personnel deployed to southwest Asia in the early 2000s.

Several chronic infections that pose a risk to war veterans tend to recur or become clinically active in immunocompromised individuals and may be particularly aggressive in this population. Latent infections such as leishmaniasis, tuberculosis, histoplasmosis, brucellosis, Q fever, and strongyloidiasis in otherwise healthy veterans returning from war may become clinically expressed only much later in the setting of chronic glucocorticoid use, monoclonal antibody therapy, organ transplantation, cytotoxic chemotherapy, advanced HIV-1 disease, hematologic malignancy, or other immunosuppressive conditions. Thus, physicians should remain vigilant regarding the potential development of symptomatic disease due to such latent microbes in immunologically compromised veterans who may have initially acquired an infection years previously while serving in the military.

A number of syndromes of possible infectious etiology, some of which may present with chronic clinical manifestations, have been noted in veterans returning from recent wars. After the Gulf War in 1990–1991, numerous veterans from several countries experienced constellations of various common, nonspecific symptoms, including fatigue, musculoskeletal pain, sleep disturbances, and difficulty concentrating. Despite exhaustive investigations and several hypotheses regarding potential etiologies of this chronic multisystemic illness, including infectious agents, no unifying or single cause has been identified. In a randomized, placebo-controlled trial, a prolonged course of

doxycycline failed to alleviate such symptoms at 1 year. In 2003, a small outbreak of acute idiopathic eosinophilic pneumonia was reported among U.S. troops serving in southwest Asia. Although a thorough investigation failed to elucidate an infectious etiology, it is noteworthy that symptoms developed up to 11 months after arrival in the theater of combat; this time frame suggests that such cases may become clinically manifest after return to the home front.

Chronic Wound Infections and Osteomyelitis War wounds, an important cause of morbidity in all armed conflicts, are at high risk of infection due to contamination with environmental bacteria and the presence of retained foreign bodies. In recent conflicts, improved survival rates due to enhanced and expedited care of combat casualties have had the unintended consequence of increasing the potential for infectious complications, a situation exacerbated by repeated and at times prolonged exposure to health care environments and their associated pathogens. Many wounds sustained in recent wars have resulted from penetrating soft-tissue trauma and open fractures of the extremities—injuries attributable to improvised explosive devices used as antipersonnel weapons and to body armor that leaves the limbs unprotected. Cultures of samples taken at the time of injury at a combat support hospital in Iraq revealed that most contaminated wounds harbored gram-positive commensal skin bacteria; other investigators, however, have noted an early predominance of gram-negative bacteria, including multidrug-resistant (MDR) pathogens.

Approximately 3% of nearly 17,000 combat injuries sustained between 2003 and 2009 in U.S. military operations in Iraq and Afghanistan involved soft tissue infections. Although it is not clear how many of these infections became chronic or progressed to involve deeper tissue structures, a significant number were managed in tertiary care facilities, many on the home front. The bacteriology of infected combat wounds comprises predominantly gram-negative bacilli and MDR organisms. Broad-spectrum antimicrobial prophylaxis administered at the time of injury appears to be a risk factor for subsequent infection; nosocomial acquisition of health care-associated MDR pathogens likely contributes as well. Invasive fungal infections have recently emerged as a significant cause of morbidity and death in the context of combat wounds.

During the past decade of wars in Iraq and Afghanistan, MDR strains of *Acinetobacter baumannii* (Chap. 187) have emerged as

important pathogens in both wound and bloodstream infections among returning veterans treated at U.S. health care facilities. The majority of isolates display in vitro susceptibility to amikacin and variable susceptibility to carbapenems but are largely resistant to other commonly used antimicrobial agents. Antimicrobial treatment should be guided by in vitro susceptibility data; patients who are critically ill, are immunocompromised, or have significant medical co-morbidities may benefit from combination therapy. Colistin (polymyxin E) has been shown to be clinically effective against *Acinetobacter* infections caused by isolates resistant to both aminoglycosides and carbapenems. Mortality rates have been low among immunocompetent hosts receiving appropriate antimicrobial treatment and undergoing debridement; however, *Acinetobacter* infections in immunocompromised individuals are associated with higher mortality risk. Strict adherence to hand-washing and other infection control procedures is important to limit the nosocomial spread of MDR organisms.

Chronic osteomyelitis related to either extension of a contiguous soft tissue infection or an infected prosthesis also represents a burgeoning problem for wounded veterans of recent wars. Limited microbiologic data have shown a predominance of gram-negative etiologic agents—most often *Acinetobacter* and *Pseudomonas aeruginosa*—in the initial episodes of osteomyelitis but a shift to staphylococcal isolates in the majority of recurrent cases—a change that may perhaps be related to nosocomial acquisition. Relapses have been noted to occur 1 month to 1 year after treatment of the initial infection.

Veterans with traumatic brain injury, who have accounted for 22% of American casualties in recent wars in Iraq and Afghanistan, are at risk for infectious complications due to several factors: the presence of foreign bodies or prosthetic material related to their traumatic wounds; acquisition of a wide range of nosocomial infections during repeated interactions with the health care system; and injury-induced cognitive changes that may increase impulsivity and risk-taking behaviors. In line with the last-mentioned factor, this subgroup of veterans may be at heightened risk for substance abuse and other practices that expose them to various bloodborne and sexually transmitted infections. Moreover, they may be at risk for post-neurosurgical complications, such as pyogenic meningitis due to multidrug-resistant *Acinetobacter* species.

SECTION 2

CLINICAL SYNDROMES: COMMUNITY-ACQUIRED INFECTIONS

153 Pneumonia

Lionel A. Mandell, Richard G. Wunderink

DEFINITION

Pneumonia is an infection of the pulmonary parenchyma. Despite being the cause of significant morbidity and mortality, pneumonia is often misdiagnosed, mistreated, and underestimated. In the past, pneumonia was typically classified as community-acquired (CAP), hospital-acquired (HAP), or ventilator-associated (VAP). Over the past two decades, however, some persons presenting with onset of pneumonia as outpatients have been found to be infected with the multidrug-resistant (MDR) pathogens previously associated with HAP. Factors responsible for this phenomenon include the development and widespread use of potent oral antibiotics, earlier transfer of patients out of acute-care hospitals to their homes or various lower-acuity facilities, increased use of outpatient IV antibiotic therapy, general aging of the

population, and more extensive immunomodulatory therapies. The potential involvement of these MDR pathogens has led to a designation for a new category of pneumonia—*health care-associated pneumonia* (HCAP)—that is distinct from CAP. Conditions associated with HCAP and the likely pathogens are listed in [Table 153-1](#).

Although the new classification system has been helpful in designing empirical antibiotic strategies, it is not without its disadvantages. Not all MDR pathogens are associated with all risk factors (Table 153-1). Moreover, HCAP is a distillation of multiple risk factors, and each patient must be considered individually. For example, the risk of infection with MDR pathogens for a nursing home resident who has dementia but can independently dress, ambulate, and eat is quite different from the risk for a patient who is in a chronic vegetative state with a tracheostomy and a percutaneous feeding tube in place. In addition, risk factors for MDR infection do not preclude the development of pneumonia caused by the usual CAP pathogens.

This chapter deals with pneumonia in patients who are not considered to be immunocompromised. **Pneumonia in severely immunocompromised patients, some of whom overlap with the groups**

TABLE 153-1 CLINICAL CONDITIONS ASSOCIATED WITH AND LIKELY PATHOGENS IN HEALTH CARE–ASSOCIATED PNEUMONIA

Condition	Pathogen			
	MRSA	<i>Pseudomonas aeruginosa</i>	<i>Acinetobacter</i> spp.	MDR Enterobacteriaceae
Hospitalization for ≥ 48 h	√	√	√	√
Hospitalization for ≥ 2 days in prior 3 months	√	√	√	√
Nursing home or extended-care-facility residence	√	√	√	√
Antibiotic therapy in preceding 3 months		√		√
Chronic dialysis	√			
Home infusion therapy	√			
Home wound care	√			
Family member with MDR infection	√			√

Abbreviations: MDR, multidrug-resistant; MRSA, methicillin-resistant *Staphylococcus aureus*.

PATHOPHYSIOLOGY

Pneumonia results from the proliferation of microbial pathogens at the alveolar level and the host’s response to those pathogens. Microorganisms gain access to the lower respiratory tract in several ways. The most common is by aspiration from the oropharynx. Small-volume aspiration occurs frequently during sleep (especially in the elderly) and in patients with decreased levels of consciousness. Many pathogens are inhaled as contaminated droplets. Rarely, pneumonia occurs via hematogenous spread (e.g., from tricuspid endocarditis) or by contiguous extension from an infected pleural or mediastinal space.

Mechanical factors are critically important in host defense. The hairs and turbinates of the nares capture larger inhaled particles before they reach the lower respiratory tract. The branching architecture of the tracheobronchial tree traps microbes on the airway lining, where mucociliary clearance and local antibacterial factors either clear or kill the potential pathogen. The gag reflex and the cough mechanism offer critical protection from aspiration. In addition, the normal flora adhering to mucosal cells of the oropharynx, whose components are remarkably constant, prevents pathogenic bacteria from binding and thereby decreases the risk of pneumonia caused by these more virulent bacteria.

When these barriers are overcome or when microorganisms are small enough to be inhaled to the alveolar level, resident alveolar macrophages are extremely efficient at clearing and killing pathogens. Macrophages are assisted by proteins that are produced by the alveolar epithelial cells (e.g., surfactant proteins A and D) and that have intrinsic opsonizing properties or antibacterial or antiviral activity. Once engulfed by the macrophage, the pathogens—even if they are not killed—are eliminated via either the mucociliary elevator or the lymphatics and no longer represent an infectious challenge. Only when the capacity of the alveolar macrophages to ingest or kill the microorganisms is exceeded does clinical pneumonia become manifest. In that situation, the alveolar macrophages initiate the inflammatory response to bolster lower respiratory tract defenses. The host inflammatory response, rather than proliferation of microorganisms, triggers the clinical syndrome of pneumonia. The release of inflammatory mediators, such as interleukin 1 and tumor necrosis factor, results in fever. Chemokines, such as interleukin 8 and granulocyte colony-stimulating factor, stimulate the release of neutrophils and their attraction to the lung, producing both peripheral leukocytosis and increased purulent secretions. Inflammatory mediators released by macrophages and the newly recruited neutrophils create an alveolar capillary leak equivalent to that seen in the acute respiratory distress syndrome, although in pneumonia this leak is localized (at least initially). Even erythrocytes can cross the alveolar-capillary membrane, with consequent hemoptysis. The capillary leak results in a radiographic infiltrate and rales detectable on auscultation, and hypoxemia results from alveolar filling. Moreover, some bacterial pathogens appear to interfere with the hypoxemic vasoconstriction that would normally occur with fluid-filled alveoli, and this interference can result in severe hypoxemia. Increased respiratory drive in the systemic inflammatory response syndrome (Chap. 325) leads to respiratory alkalosis. Decreased compliance due to capillary leak, hypoxemia, increased respiratory drive, increased secretions, and occasionally infection-related bronchospasm all lead to dyspnea. If severe enough, the changes in lung mechanics secondary to reductions in lung volume and compliance and the intrapulmonary shunting of blood may cause respiratory failure and the patient’s death.

PATHOLOGY

Classic pneumonia evolves through a series of pathologic changes. The initial phase is one of edema, with the presence of a proteinaceous exudate—and often of bacteria—in the alveoli. This phase is rarely evident in clinical or autopsy specimens because of the rapid transition to the

red hepatization phase. The presence of erythrocytes in the cellular intraalveolar exudate gives this second stage its name, but neutrophil influx is more important with regard to host defense. Bacteria are occasionally seen in pathologic specimens collected during this phase. In the third phase, gray hepatization, no new erythrocytes are extravasating, and those already present have been lysed and degraded. The neutrophil is the predominant cell, fibrin deposition is abundant, and bacteria have disappeared. This phase corresponds with successful containment of the infection and improvement in gas exchange. In the final phase, resolution, the macrophage reappears as the dominant cell type in the alveolar space, and the debris of neutrophils, bacteria, and fibrin has been cleared, as has the inflammatory response.

This pattern has been described best for lobar pneumococcal pneumonia and may not apply to pneumonia of all etiologies, especially viral or *Pneumocystis* pneumonia. In VAP, respiratory bronchiolitis may precede the development of a radiologically apparent infiltrate. Because of the microaspiration mechanism, a bronchopneumonia pattern is most common in nosocomial pneumonias, whereas a lobar pattern is more common in bacterial CAP. Despite the radiographic appearance, viral and *Pneumocystis* pneumonias represent alveolar rather than interstitial processes.

COMMUNITY-ACQUIRED PNEUMONIA

ETIOLOGY

The extensive list of potential etiologic agents in CAP includes bacteria, fungi, viruses, and protozoa. Newly identified pathogens include metapneumoviruses, the coronaviruses responsible for severe acute respiratory syndrome (SARS) and Middle East respiratory syndrome, and community-acquired strains of methicillin-resistant *Staphylococcus aureus* (MRSA). Most cases of CAP, however, are caused by relatively few pathogens (Table 153-2). Although *Streptococcus pneumoniae* is most common, other organisms also must be considered in light of the patient’s risk factors and severity of illness. Separation of potential agents into either “typical” bacterial pathogens or “atypical” organisms may be helpful. The former category includes *S. pneumoniae*, *Haemophilus influenzae*, and (in selected patients) *S. aureus* and gram-negative bacilli such as *Klebsiella pneumoniae* and *Pseudomonas aeruginosa*. The “atypical” organisms include *Mycoplasma pneumoniae*, *Chlamydia pneumoniae*, and *Legionella* species (in inpatients) as well as respiratory viruses such as influenza viruses, adenoviruses, human metapneumovirus, and respiratory syncytial viruses. Viruses may be responsible for a large proportion of CAP cases that require hospital admission, even in adults. Atypical organisms cannot be cultured on standard media or seen on Gram’s stain. The frequency and importance of atypical pathogens have significant implications for therapy. These organisms are intrinsically resistant to all β-lactam agents and must be treated with a macrolide, a fluoroquinolone, or a tetracycline. In the ~10–15% of CAP cases that are polymicrobial, the etiology usually includes a combination of typical and atypical pathogens.

Anaerobes play a significant role only when an episode of aspiration has occurred days to weeks before presentation of pneumonia. The

TABLE 153-2 MICROBIAL CAUSES OF COMMUNITY-ACQUIRED PNEUMONIA, BY SITE OF CARE

Outpatients	Hospitalized Patients	
	Non-ICU	ICU
<i>Streptococcus pneumoniae</i>	<i>S. pneumoniae</i>	<i>S. pneumoniae</i>
<i>Mycoplasma pneumoniae</i>	<i>M. pneumoniae</i>	<i>Staphylococcus aureus</i>
<i>Haemophilus influenzae</i>	<i>Chlamydia pneumoniae</i>	<i>Legionella</i> spp.
<i>C. pneumoniae</i>	<i>H. influenzae</i>	Gram-negative bacilli
Respiratory viruses ^a	<i>Legionella</i> spp.	<i>H. influenzae</i>
	Respiratory viruses ^a	

^aInfluenza A and B viruses, human metapneumovirus, adenoviruses, respiratory syncytial viruses, parainfluenza viruses.

Note: Pathogens are listed in descending order of frequency. ICU, intensive care unit.

combination of an unprotected airway (e.g., in patients with alcohol or drug overdose or a seizure disorder) and significant gingivitis constitutes the major risk factor. Anaerobic pneumonias are often complicated by abscess formation and by significant empyemas or parapneumonic effusions.

S. aureus pneumonia is well known to complicate influenza infection. However, MRSA has been reported as the primary etiologic agent of CAP. While this entity is still relatively uncommon, clinicians must be aware of its potentially serious consequences, such as necrotizing pneumonia. Two important developments have led to this problem: the spread of MRSA from the hospital setting to the community and the emergence of genetically distinct strains of MRSA in the community. The former circumstance is more likely to result in HCAP, whereas the novel community-acquired MRSA (CA-MRSA) strains may infect healthy individuals with no association with health care.

Unfortunately, despite a careful history and physical examination as well as routine radiographic studies, the causative pathogen in a case of CAP is difficult to predict with any degree of certainty; in more than one-half of cases, a specific etiology is never determined. Nevertheless, epidemiologic and risk factors may suggest the involvement of certain pathogens (Table 153-3).

EPIDEMIOLOGY

More than 5 million CAP cases occur annually in the United States; usually, 80% of the affected patients are treated as outpatients and 20% as inpatients. The mortality rate among outpatients is usually $\leq 1\%$, whereas among hospitalized patients the rate can range from $\sim 12\%$ to 40%, depending on whether treatment is provided in or outside of the intensive care unit (ICU). CAP results in more than 1.2 million hospitalizations and more than 55,000 deaths annually. The overall yearly cost associated with CAP is estimated at \$12 billion. The incidence rates are highest at the extremes of age. The overall annual rate in the United States is 12 cases/1000 persons, but the figure increases to 12–18/1000 among children < 4 years of age and to 20/1000 among persons > 60 years of age.

TABLE 153-3 EPIDEMIOLOGIC FACTORS SUGGESTING POSSIBLE CAUSES OF COMMUNITY-ACQUIRED PNEUMONIA

Factor	Possible Pathogen(s)
Alcoholism	<i>Streptococcus pneumoniae</i> , oral anaerobes, <i>Klebsiella pneumoniae</i> , <i>Acinetobacter</i> spp., <i>Mycobacterium tuberculosis</i>
COPD and/or smoking	<i>Haemophilus influenzae</i> , <i>Pseudomonas aeruginosa</i> , <i>Legionella</i> spp., <i>S. pneumoniae</i> , <i>Moraxella catarrhalis</i> , <i>Chlamydia pneumoniae</i>
Structural lung disease (e.g., bronchiectasis)	<i>P. aeruginosa</i> , <i>Burkholderia cepacia</i> , <i>Staphylococcus aureus</i>
Dementia, stroke, decreased level of consciousness	Oral anaerobes, gram-negative enteric bacteria
Lung abscess	CA-MRSA, oral anaerobes, endemic fungi, <i>M. tuberculosis</i> , atypical mycobacteria
Travel to Ohio or St. Lawrence river valleys	<i>Histoplasma capsulatum</i>
Travel to southwestern United States	Hantavirus, <i>Coccidioides</i> spp.
Travel to Southeast Asia	<i>Burkholderia pseudomallei</i> , avian influenza virus
Stay in hotel or on cruise ship in previous 2 weeks	<i>Legionella</i> spp.
Local influenza activity	Influenza virus, <i>S. pneumoniae</i> , <i>S. aureus</i>
Exposure to bats or birds	<i>H. capsulatum</i>
Exposure to birds	<i>Chlamydia psittaci</i>
Exposure to rabbits	<i>Francisella tularensis</i>
Exposure to sheep, goats, parturient cats	<i>Coxiella burnetii</i>

Abbreviations: CA-MRSA, community-acquired methicillin-resistant *Staphylococcus aureus*; COPD, chronic obstructive pulmonary disease.

The risk factors for CAP in general and for pneumococcal pneumonia in particular have implications for treatment regimens. Risk factors for CAP include alcoholism, asthma, immunosuppression, institutionalization, and an age of ≥ 70 years. In the elderly, factors such as decreased cough and gag reflexes as well as reduced antibody and Toll-like receptor responses increase the likelihood of pneumonia. Risk factors for pneumococcal pneumonia include dementia, seizure disorders, heart failure, cerebrovascular disease, alcoholism, tobacco smoking, chronic obstructive pulmonary disease, and HIV infection. CA-MRSA pneumonia is more likely in patients with skin colonization or infection with CA-MRSA. Enterobacteriaceae tend to infect patients who have recently been hospitalized and/or received antibiotic therapy or who have comorbidities such as alcoholism, heart failure, or renal failure. *P. aeruginosa* is a particular problem in patients with severe structural lung disease, such as bronchiectasis, cystic fibrosis, or severe chronic obstructive pulmonary disease. Risk factors for *Legionella* infection include diabetes, hematologic malignancy, cancer, severe renal disease, HIV infection, smoking, male gender, and a recent hotel stay or ship cruise. (Many of these risk factors would now reclassify as HCAP some cases that were previously designated CAP.)

CLINICAL MANIFESTATIONS

CAP can vary from indolent to fulminant in presentation and from mild to fatal in severity. Manifestations of progression and severity include both constitutional findings and those limited to the lung and associated structures. In light of the pathobiology of the disease, many of the findings are to be expected.

The patient is frequently febrile with tachycardia or may have a history of chills and/or sweats. Cough may be either nonproductive or productive of mucoid, purulent, or blood-tinged sputum. Gross hemoptysis is suggestive of CA-MRSA pneumonia. Depending on severity, the patient may be able to speak in full sentences or may be very short of breath. If the pleura is involved, the patient may experience pleuritic chest pain. Up to 20% of patients may have gastrointestinal symptoms such as nausea, vomiting, and/or diarrhea. Other symptoms may include fatigue, headache, myalgias, and arthralgias.

Findings on physical examination vary with the degree of pulmonary consolidation and the presence or absence of a significant pleural effusion. An increased respiratory rate and use of accessory muscles of respiration are common. Palpation may reveal increased or decreased tactile fremitus, and the percussion note can vary from dull to flat, reflecting underlying consolidated lung and pleural fluid, respectively. Crackles, bronchial breath sounds, and possibly a pleural friction rub may be heard on auscultation. The clinical presentation may not be so obvious in the elderly, who may initially display new-onset or worsening confusion and few other manifestations. Severely ill patients may have septic shock and evidence of organ failure.

DIAGNOSIS

When confronted with possible CAP, the physician must ask two questions: Is this pneumonia, and, if so, what is the likely etiology? The former question is typically answered by clinical and radiographic methods, whereas the latter requires the aid of laboratory techniques.

Clinical Diagnosis The differential diagnosis includes both infectious and noninfectious entities such as acute bronchitis, acute exacerbations of chronic bronchitis, heart failure, pulmonary embolism, hypersensitivity pneumonitis, and radiation pneumonitis. The importance of a careful history cannot be overemphasized. For example, known cardiac disease may suggest worsening pulmonary edema, while underlying carcinoma may suggest lung injury secondary to irradiation.

Unfortunately, the sensitivity and specificity of the findings on physical examination are less than ideal, averaging 58% and 67%, respectively. Therefore, chest radiography is often necessary to differentiate CAP from other conditions. Radiographic findings may include risk factors for increased severity (e.g., cavitation or multilobar involvement). Occasionally, radiographic results suggest an etiologic diagnosis. For example, pneumatoceles suggest infection with *S. aureus*, and an upper-lobe cavitating lesion suggests tuberculosis. CT may be of value in a patient with suspected postobstructive

806 pneumonia caused by a tumor or foreign body or suspected cavitory disease. For outpatients, the clinical and radiologic assessments are usually all that is done before treatment for CAP is started since most laboratory results are not available soon enough to influence initial management significantly. In certain cases, the availability of rapid point-of-care outpatient diagnostic tests can be very important; for example, rapid diagnosis of influenza virus infection can prompt specific anti-influenza drug treatment and secondary prevention.

Etiologic Diagnosis The etiology of pneumonia usually cannot be determined solely on the basis of clinical presentation. Except for CAP patients admitted to the ICU, no data exist to show that treatment directed at a specific pathogen is statistically superior to empirical therapy. The benefit of establishing a microbial etiology can therefore be questioned, particularly in light of the cost of diagnostic testing. However, a number of reasons can be advanced for attempting an etiologic diagnosis. Identification of an unexpected pathogen allows narrowing of the initial empirical regimen, thereby decreasing antibiotic selection pressure and lessening the risk of resistance. Pathogens with important public safety implications, such as *Mycobacterium tuberculosis* and influenza virus, may be found in some cases. Finally, without culture and susceptibility data, trends in resistance cannot be followed accurately, and appropriate empirical therapeutic regimens are harder to devise.

GRAM'S STAIN AND CULTURE OF SPUTUM The main purpose of the sputum Gram's stain is to ensure that a sample is suitable for culture. However, Gram's staining may also identify certain pathogens (e.g., *S. pneumoniae*, *S. aureus*, and gram-negative bacteria) by their characteristic appearance. To be adequate for culture, a sputum sample must have >25 neutrophils and <10 squamous epithelial cells per low-power field. The sensitivity and specificity of the sputum Gram's stain and culture are highly variable. Even in cases of proven bacteremic pneumococcal pneumonia, the yield of positive cultures from sputum samples is ≤50%.

Some patients, particularly elderly individuals, may not be able to produce an appropriate expectorated sputum sample. Others may already have started a course of antibiotics that can interfere with culture results at the time a sample is obtained. Inability to produce sputum can be a consequence of dehydration, and the correction of this condition may result in increased sputum production and a more obvious infiltrate on chest radiography. For patients admitted to the ICU and intubated, a deep-suction aspirate or bronchoalveolar lavage sample (obtained either via bronchoscopy or non-bronchoscopically) has a high yield on culture when sent to the microbiology laboratory as soon as possible. Since the etiologies in severe CAP are somewhat different from those in milder disease (Table 153-2), the greatest benefit of staining and culturing respiratory secretions is to alert the physician of unsuspected and/or resistant pathogens and to permit appropriate modification of therapy. Other stains and cultures (e.g., specific stains for *M. tuberculosis* or fungi) may be useful as well.

BLOOD CULTURES The yield from blood cultures, even when samples are collected before antibiotic therapy, is disappointingly low. Only 5–14% of cultures of blood from patients hospitalized with CAP are positive, and the most frequently isolated pathogen is *S. pneumoniae*. Since recommended empirical regimens all provide pneumococcal coverage, a blood culture positive for this pathogen has little, if any, effect on clinical outcome. However, susceptibility data may allow narrowing of antibiotic therapy in appropriate cases. Because of the low yield and the lack of significant impact on outcome, blood cultures are no longer considered *de rigueur* for all hospitalized CAP patients. Certain high-risk patients—including those with neutropenia secondary to pneumonia, asplenia, complement deficiencies, chronic liver disease, or severe CAP—should have blood cultured.

URINARY ANTIGEN TESTS Two commercially available tests detect pneumococcal and *Legionella* antigen in urine. The test for *Legionella pneumophila* detects only serogroup 1, but this serogroup accounts for most community-acquired cases of Legionnaires' disease in the United States. The sensitivity and specificity of the *Legionella* urine antigen

test are as high as 90% and 99%, respectively. The pneumococcal urine antigen test also is quite sensitive and specific (80% and >90%, respectively). Although false-positive results can be obtained with samples from pneumococcus-colonized children, the test is generally reliable. Both tests can detect antigen even after the initiation of appropriate antibiotic therapy.

POLYMERASE CHAIN REACTION Polymerase chain reaction (PCR) tests, which amplify a microorganism's DNA or RNA, are available for a number of pathogens. PCR of nasopharyngeal swabs has become the standard for diagnosis of respiratory viral infection. In addition, PCR can detect the nucleic acid of *Legionella* species, *M. pneumoniae*, *C. pneumoniae*, and mycobacteria. In patients with pneumococcal pneumonia, an increased bacterial load in whole blood documented by PCR is associated with an increased risk of septic shock, the need for mechanical ventilation, and death. Clinical availability of such a test could conceivably help identify patients suitable for ICU admission.

SEROLOGY A fourfold rise in specific IgM antibody titer between acute- and convalescent-phase serum samples is generally considered diagnostic of infection with the pathogen in question. In the past, serologic tests were used to help identify atypical pathogens as well as selected unusual organisms such as *Coxiella burnetii*. Recently, however, they have fallen out of favor because of the time required to obtain a final result for the convalescent-phase sample.

BIOMARKERS A number of substances can serve as markers of severe inflammation. The two currently in use are C-reactive protein (CRP) and procalcitonin (PCT). Levels of these acute-phase reactants increase in the presence of an inflammatory response, particularly to bacterial pathogens. CRP may be of use in the identification of worsening disease or treatment failure, and PCT may play a role in determining the need for antibacterial therapy. These tests should not be used on their own but, when interpreted in conjunction with other findings from the history, physical examination, radiology, and laboratory tests, may help with antibiotic stewardship and appropriate management of seriously ill patients with CAP.

TREATMENT COMMUNITY-ACQUIRED PNEUMONIA

SITE OF CARE

The cost of inpatient management exceeds that of outpatient treatment by a factor of 20, and hospitalization accounts for most CAP-related expenditures. Thus the decision to admit a patient with CAP to the hospital has considerable implications. Certain patients clearly can be managed at home, and others clearly require treatment in the hospital, but the choice is sometimes difficult. Tools that objectively assess the risk of adverse outcomes, including severe illness and death, can minimize unnecessary hospital admissions. There are currently two sets of criteria: the Pneumonia Severity Index (PSI), a prognostic model used to identify patients at low risk of dying; and the CURB-65 criteria, a severity-of-illness score.

To determine the PSI, points are given for 20 variables, including age, coexisting illness, and abnormal physical and laboratory findings. On the basis of the resulting score, patients are assigned to one of five classes with the following mortality rates: class 1, 0.1%; class 2, 0.6%; class 3, 2.8%; class 4, 8.2%; and class 5, 29.2%. Determination of the PSI is often impractical in a busy emergency-department setting because of the number of variables that must be assessed. However, clinical trials demonstrate that routine use of the PSI results in lower admission rates for class 1 and class 2 patients. Patients in class 3 could ideally be admitted to an observation unit until a further decision can be made.

The CURB-65 criteria include five variables: confusion (C); urea >7 mmol/L (U); respiratory rate ≥30/min (R); blood pressure, systolic ≤90 mmHg or diastolic ≤60 mmHg (B); and age ≥65 years. Patients with a score of 0, among whom the 30-day mortality rate is 1.5%, can be treated outside the hospital. With a score of 2, the 30-day mortality rate is 9.2%, and patients should be admitted to the hospital.

TABLE 153-4 RISK FACTORS FOR EARLY DETERIORATION IN CAP

Multilobar infiltrates	Hypoalbuminemia
Severe hypoxemia (arterial saturation <90%)	Neutropenia
Severe acidosis (pH <7.30)	Thrombocytopenia
Mental confusion	Hyponatremia
Severe tachypnea (>30 breaths/min)	Hypoglycemia

Among patients with scores of ≥ 3 , mortality rates are 22% overall; these patients may require ICU admission.

It is not clear which assessment tool is superior. Whichever system is used, these objective criteria must always be tempered by careful consideration of factors relevant to individual patients, including the ability to comply reliably with an oral antibiotic regimen and the resources available to the patient outside the hospital.


Neither PSI nor CURB-65 is accurate in determining the need for ICU admission. Septic shock or respiratory failure in the emergency department is an obvious indication for ICU care. However, mortality rates are higher among less ill patients who are admitted to the floor and then deteriorate than among equally ill patients monitored in the ICU. A variety of scores have been proposed to identify patients most likely to have early deterioration (Table 153-4). Most factors in these scores are similar to the minor severity criteria proposed by the Infectious Diseases Society of America (IDSA) and the American Thoracic Society (ATS) in their guidelines for the management of CAP.

ANTIBIOTIC RESISTANCE

Antimicrobial resistance is a significant problem that threatens to diminish our therapeutic armamentarium. Misuse of antibiotics results in increased antibiotic selection pressure that can affect resistance locally or even globally by clonal dissemination. For CAP, the main resistance issues currently involve *S. pneumoniae* and CA-MRSA.

S. pneumoniae In general, pneumococcal resistance is acquired (1) by direct DNA incorporation and remodeling resulting from contact with closely related oral commensal bacteria, (2) by the process of natural transformation, or (3) by mutation of certain genes.

The minimal inhibitory concentration (MIC) cutoffs for penicillin in pneumonia are ≤ 2 $\mu\text{g/mL}$ for susceptibility, >2 – 4 $\mu\text{g/mL}$ for intermediate, and ≥ 8 $\mu\text{g/mL}$ for resistant. A change in susceptibility thresholds resulted in a dramatic decrease in the proportion of pneumococcal isolates considered nonsusceptible. For meningitis, MIC thresholds remain at the former higher levels. Fortunately, resistance to penicillin appeared to plateau even before the change in MIC thresholds. Pneumococcal resistance to β -lactam drugs is due solely to low-affinity penicillin-binding proteins. Risk factors for penicillin-resistant pneumococcal infection include recent antimicrobial therapy, an age of <2 years or >65 years, attendance at day-care centers, recent hospitalization, and HIV infection.


 In contrast to penicillin resistance, resistance to macrolides is increasing through several mechanisms. *Target-site modification* caused by ribosomal methylation in 23S rRNA encoded by the *ermB* gene results in high-level resistance (MICs, ≥ 64 $\mu\text{g/mL}$) to macrolides, lincosamides, and streptogramin B–type antibiotics. The *efflux mechanism* encoded by the *mef* gene (*M phenotype*) is usually associated with low-level resistance (MICs, 1–32 $\mu\text{g/mL}$). These two mechanisms account for ~45% and ~65%, respectively, of resistant pneumococcal isolates in the United States. High-level resistance to macrolides is more common in Europe, whereas lower-level resistance predominates in North America.

Pneumococcal resistance to fluoroquinolones (e.g., ciprofloxacin and levofloxacin) has been reported. Changes can occur in one or both target sites (topoisomerases II and IV) from mutations in the *gyrA* and *parC* genes, respectively. In addition, an efflux pump may play a role in pneumococcal resistance to fluoroquinolones.

Isolates resistant to drugs from three or more antimicrobial classes with different mechanisms of action are considered MDR strains. The propensity for an association of pneumococcal resistance to penicillin with reduced susceptibility to other drugs, such as macrolides, tetracyclines, and trimethoprim-sulfamethoxazole, also is of concern. In the United States, 58.9% of penicillin-resistant pneumococcal isolates from blood are also resistant to macrolides.

The most important risk factor for antibiotic-resistant pneumococcal infection is use of a specific antibiotic within the previous 3 months. Therefore, a patient's history of prior antibiotic treatment is a critical factor in avoiding the use of an inappropriate antibiotic.

CA-MRSA CAP due to MRSA may be caused by the classic hospital-acquired strains or by the more recently identified, genotypically and phenotypically distinct community-acquired strains. Most infections with the former strains have been acquired either directly or indirectly by contact with the health care environment and would now be classified as HCAP. However, in some hospitals, the CA-MRSA strains are displacing the classic hospital-acquired strains—a trend suggesting that the newer strains may be more robust and blurring this distinction.

 Methicillin resistance in *S. aureus* is determined by the *mecA* gene, which encodes for resistance to all β -lactam drugs. At least five *staphylococcal chromosomal cassette mec* (SCCmec) types have been described. The typical hospital-acquired strain usually has type II or III, whereas CA-MRSA has a type IV SCCmec element. CA-MRSA isolates tend to be less resistant than the older hospital-acquired strains and are often susceptible to trimethoprim-sulfamethoxazole, clindamycin, and tetracycline in addition to vancomycin and linezolid. However, the most important distinction is that CA-MRSA strains also carry genes for superantigens, such as enterotoxins B and C and Panton-Valentine leukocidin, a membrane-tropic toxin that can create cytolytic pores in polymorphonuclear neutrophils, monocytes, and macrophages.

Gram-Negative Bacilli A detailed discussion of resistance among gram-negative bacilli is beyond the scope of this chapter (Chap. 186). Fluoroquinolone resistance among isolates of *Escherichia coli* from the community appears to be increasing. *Enterobacter* species are typically resistant to cephalosporins; the drugs of choice for use against these bacteria are usually fluoroquinolones or carbapenems. Similarly, when infections due to bacteria producing extended-spectrum β -lactamases are documented or suspected, a fluoroquinolone or a carbapenem should be used; these MDR strains are more likely to be involved in HCAP.

INITIAL ANTIBIOTIC MANAGEMENT

Since the etiology of CAP is rarely known at the outset of treatment, initial therapy is usually empirical, designed to cover the most likely pathogens (Table 153-5). In all cases, antibiotic treatment should be initiated as expeditiously as possible. The CAP treatment guidelines in the United States (summarized in Table 153-5) represent joint statements from the IDSA and the ATS; the Canadian guidelines come from the Canadian Infectious Disease Society and the Canadian Thoracic Society. In all these guidelines, coverage is always provided for the pneumococcus and the atypical pathogens. In contrast, guidelines from some European countries do not always include atypical coverage based on local epidemiologic data. The U.S./Canadian approach is supported by retrospective data derived from administrative databases including thousands of patients. Atypical pathogen coverage provided by the addition of a macrolide to a cephalosporin or by the use of a fluoroquinolone alone has been consistently associated with a significant reduction in mortality rates compared with those for β -lactam coverage alone.

Therapy with a macrolide or a fluoroquinolone within the previous 3 months is associated with an increased likelihood of infection with a resistant strain of *S. pneumoniae*. For this reason, a fluoroquinolone-based regimen should be used for patients recently given a macrolide, and vice versa (Table 153-5).

TABLE 153-5 EMPIRICAL ANTIBIOTIC TREATMENT OF COMMUNITY-ACQUIRED PNEUMONIA**Outpatients**

1. Previously healthy and no antibiotics in past 3 months
 - A macrolide (clarithromycin [500 mg PO bid] or azithromycin [500 mg PO once, then 250 mg qd]) *or*
 - Doxycycline (100 mg PO bid)
2. Comorbidities or antibiotics in past 3 months: select an alternative from a different class
 - A respiratory fluoroquinolone (moxifloxacin [400 mg PO qd], gemifloxacin [320 mg PO qd], levofloxacin [750 mg PO qd]) *or*
 - A β -lactam (preferred: high-dose amoxicillin [1 g tid] or amoxicillin/clavulanate [2 g bid]; alternatives: ceftriaxone [1–2 g IV qd], cefpodoxime [200 mg PO bid], cefuroxime [500 mg PO bid]) *plus* a macrolide^a
3. In regions with a high rate of “high-level” pneumococcal macrolide resistance,^b consider alternatives listed above for patients with comorbidities.

Inpatients, Non-ICU

- A respiratory fluoroquinolone (e.g., moxifloxacin [400 mg PO or IV qd] or levofloxacin [750 mg PO or IV qd])
- A β -lactam^c (e.g., ceftriaxone [1–2 g IV qd], ampicillin [1–2 g IV q4–6h], cefotaxime [1–2 g IV q8h], ertapenem [1 g IV qd]) *plus* a macrolide^d (e.g., oral clarithromycin or azithromycin [as listed above] or IV azithromycin [1 g once, then 500 mg qd])

Inpatients, ICU

- A β -lactam^c (e.g., ceftriaxone [2 g IV qd], ampicillin-sulbactam [2 g IV q8h], or cefotaxime [1–2 g IV q8h]) *plus* either azithromycin or a fluoroquinolone (as listed above for inpatients, non-ICU)

Special Concerns

If Pseudomonas is a consideration:

- An antipseudomonal β -lactam (e.g., piperacillin/tazobactam [4.5 g IV q6h], cefepime [1–2 g IV q12h], imipenem [500 mg IV q6h], meropenem [1 g IV q8h]) *plus* either ciprofloxacin (400 mg IV q12h) or levofloxacin (750 mg IV qd)
- The above β -lactams *plus* an aminoglycoside (amikacin [15 mg/kg qd] or tobramycin [1.7 mg/kg qd]) *plus* azithromycin
- The above β -lactams^c *plus* an aminoglycoside *plus* an antipneumococcal fluoroquinolone

If CA-MRSA is a consideration:

- Add linezolid (600 mg IV q12h) or vancomycin (15 mg/kg q12h initially, with adjusted doses)

^aDoxycycline (100 mg PO bid) is an alternative to the macrolide. ^bMICs of >16 μ g/mL in 25% of isolates. ^cA respiratory fluoroquinolone should be used for penicillin-allergic patients. ^dDoxycycline (100 mg IV q12h) is an alternative to the macrolide. ^eFor penicillin-allergic patients, use a respiratory fluoroquinolone and aztreonam (2 g IV q8h). ^fFor penicillin-allergic patients, substitute aztreonam.

Abbreviations: CA-MRSA, community-acquired methicillin-resistant *Staphylococcus aureus*; ICU, intensive care unit.

Once the etiologic agent(s) and susceptibilities are known, therapy may be altered to target the specific pathogen(s). However, this decision is not always straightforward. If blood cultures yield *S. pneumoniae* sensitive to penicillin after 2 days of treatment with a macrolide plus a β -lactam or with a fluoroquinolone alone, should therapy be switched to penicillin alone? The concern here is that a β -lactam alone would not be effective in the potential 15% of cases with atypical co-infection. No standard approach exists. In all cases, the individual patient and the various risk factors must be considered.

Management of bacteremic pneumococcal pneumonia also is controversial. Data from nonrandomized studies suggest that combination therapy (especially macrolide/ β -lactam) is associated with a lower mortality rate than monotherapy, particularly in severely ill patients. The exact reason is unknown, but possible explanations include an additive or synergistic antibacterial effect, antimicrobial tolerance, atypical co-infection, or the immunomodulatory effects of the macrolides.

For CAP patients admitted to the ICU, the risk of infection with *P. aeruginosa* or CA-MRSA is increased. Empirical coverage should

be considered when a patient has risk factors or a Gram's stain suggestive of these pathogens (Table 153–5). If CA-MRSA is suspected, either linezolid or vancomycin can be added to the initial empirical regimen; however, there is increasing concern about vancomycin's loss of potency against MRSA, poor penetration into epithelial lining fluid, and lack of effect on toxin production relative to linezolid.

Although hospitalized patients have traditionally received initial therapy by the IV route, some drugs—particularly the fluoroquinolones—are very well absorbed and can be given orally from the outset to select patients. For patients initially treated IV, a switch to oral treatment is appropriate as long as the patient can ingest and absorb the drugs, is hemodynamically stable, and is showing clinical improvement.

The duration of treatment for CAP has generated considerable interest. Patients were previously treated for 10–14 days, but studies with fluoroquinolones and telithromycin suggest that a 5-day course is sufficient for otherwise uncomplicated CAP. Even a single dose of ceftriaxone has been associated with a significant cure rate. A longer course may be required for patients with bacteremia, metastatic infection, or infection with a virulent pathogen such as *P. aeruginosa* or CA-MRSA.

GENERAL CONSIDERATIONS

In addition to appropriate antimicrobial therapy, certain general considerations apply in dealing with CAP, HCAP, or HAP/VAP. Adequate hydration, oxygen therapy for hypoxemia, and assisted ventilation when necessary are critical to successful treatment. Patients with severe CAP who remain hypotensive despite fluid resuscitation may have adrenal insufficiency and may respond to glucocorticoid treatment. The value of adjunctive therapy, such as glucocorticoids, statins, and angiotensin-converting enzyme inhibitors, remains unproven in the management of CAP.

Failure to Improve Patients slow to respond to therapy should be reevaluated at about day 3 (sooner if their condition is worsening rather than simply not improving), and several possible scenarios should be considered. A number of noninfectious conditions mimic pneumonia, including pulmonary edema, pulmonary embolism, lung carcinoma, radiation and hypersensitivity pneumonitis, and connective tissue disease involving the lungs. If the patient truly has CAP and empirical treatment is aimed at the correct pathogen, lack of response may be explained in a number of ways. The pathogen may be resistant to the drug selected, or a sequestered focus (e.g., lung abscess or empyema) may be blocking access of the antibiotic(s) to the pathogen. The patient may be getting either the wrong drug or the correct drug at the wrong dose or frequency of administration. Another possibility is that CAP is the correct diagnosis but an unsuspected pathogen (e.g., CA-MRSA, *M. tuberculosis*, or a fungus) is the cause. Nosocomial superinfections—both pulmonary and extrapulmonary—are other possible explanations for a patient's failure to improve or deterioration. In all cases of delayed response or deteriorating condition, the patient must be carefully reassessed and appropriate studies initiated, possibly including such diverse procedures as CT or bronchoscopy.

Complications As in other severe infections, common complications of severe CAP include respiratory failure, shock and multiorgan failure, coagulopathy, and exacerbation of comorbid illnesses. Three particularly noteworthy conditions are metastatic infection, lung abscess, and complicated pleural effusion. Metastatic infection (e.g., brain abscess or endocarditis) is very unusual and will require a high degree of suspicion and a detailed workup for proper treatment. Lung abscess may occur in association with aspiration or with infection caused by a single CAP pathogen, such as CA-MRSA, *P. aeruginosa*, or (rarely) *S. pneumoniae*. Aspiration pneumonia is typically a polymicrobial infection involving both aerobes and anaerobes. A significant pleural effusion should be tapped for

both diagnostic and therapeutic purposes. If the fluid has a pH of <7, a glucose level of <2.2 mmol/L, and a lactate dehydrogenase concentration of >1000 U/L or if bacteria are seen or cultured, then it should be completely drained; a chest tube is often required and video-assisted thoracoscopy may be needed for late treatment or difficult cases.

Follow-Up Fever and leukocytosis usually resolve within 2–4 days in otherwise healthy patients with CAP, but physical findings may persist longer. Chest radiographic abnormalities are slowest to resolve (4–12 weeks), with the speed of clearance depending on the patient's age and underlying lung disease. Patients may be discharged from the hospital once their clinical conditions, including comorbidities, are stable. The site of residence after discharge (nursing home, home with family, home alone) is an important discharge timing consideration, particularly for elderly patients. For a hospitalized patient, a follow-up radiograph ~4–6 weeks later is recommended. If relapse or recurrence is documented, particularly in the same lung segment, the possibility of an underlying neoplasm must be considered.

PROGNOSIS

The prognosis of CAP depends on the patient's age, comorbidities, and site of treatment (inpatient or outpatient). Young patients without comorbidity do well and usually recover fully after ~2 weeks. Older patients and those with comorbid conditions can take several weeks longer to recover fully. The overall mortality rate for the outpatient group is <1%. For patients requiring hospitalization, the overall mortality rate is estimated at 10%, with ~50% of deaths directly attributable to pneumonia.

PREVENTION

The main preventive measure is vaccination (**Chap. 148**). Recommendations of the Advisory Committee on Immunization Practices should be followed for influenza and pneumococcal vaccines.

A pneumococcal polysaccharide vaccine (PPV23) and a protein conjugate pneumococcal vaccine (PCV13) are available in the United States. The former product contains capsular material from 23 pneumococcal serotypes; in the latter, capsular polysaccharide from 13 of the most frequent pneumococcal pathogens affecting children is linked to an immunogenic protein. PCV13 produces T cell-dependent antigens that result in long-term immunologic memory. Administration of this vaccine to children has led to an overall decrease in the prevalence of antimicrobial-resistant pneumococci and in the incidence of invasive pneumococcal disease among both children and adults. However, vaccination can be followed by the replacement of vaccine serotypes with nonvaccine serotypes, as was seen with serotypes 19A and 35B after introduction of the original 7-valent conjugate vaccine. PCV13 now is also recommended for the elderly and for younger immunocompromised patients. Because of an increased risk of pneumococcal infection, even among patients without obstructive lung disease, smokers should be strongly encouraged to stop smoking.

Two forms of influenza vaccine are available: intramuscular inactivated vaccine and intranasal live-attenuated cold-adapted vaccine. The latter is contraindicated in immunocompromised patients. In the event of an influenza outbreak, unprotected patients at risk from complications should be vaccinated immediately and given chemoprophylaxis with either oseltamivir or zanamivir for 2 weeks—i.e., until vaccine-induced antibody levels are sufficiently high.

HEALTH CARE–ASSOCIATED PNEUMONIA

HCAP represents a transition between classic CAP and typical HAP. The definition of HCAP is still in some flux because of a lack of consistent large-scale studies. Several early studies were limited to patients with culture-positive pneumonia. In these studies, the incidence of MDR pathogens in HCAP was as high as or higher than in HAP/VAP.

MRSA in particular was more common in HCAP than in traditional HAP/VAP. Conversely, prospective studies in nontertiary-care centers have found a low incidence of MDR pathogens in HCAP.

The patients at greatest risk for HCAP are not well defined. Patients from nursing homes are not always at elevated risk for infection with MDR pathogens. Careful evaluation of nursing home residents with pneumonia suggests that their risk of MDR infection is low if they have not recently received antibiotics and are independent in most activities of daily living. Recent hospitalization (i.e., in the preceding 90 days) is also a major risk factor for infection with MDR pathogens. Conversely, nursing home patients are at increased risk of infection with influenza virus and other atypical pneumonia pathogens. Undue concern about MDR pathogens occasionally results in failure to cover atypical pathogens when treating nursing home patients. In addition, patients receiving home infusion therapy or undergoing chronic dialysis are probably at particular risk for MRSA pneumonia but may not be at greater risk for infection with *Pseudomonas* or *Acinetobacter* than are other patients who develop CAP.

In general, the management of HCAP due to MDR pathogens is similar to that of MDR HAP/VAP. This topic will therefore be covered in subsequent sections on HAP and VAP. The prognosis of HCAP is intermediate between that of CAP and VAP and is closer to that of HAP.

VENTILATOR-ASSOCIATED PNEUMONIA

Most hospital-acquired pneumonia research has focused on VAP. However, the information and principles based on this research can be applied to non-ICU HAP and HCAP as well. The greatest difference between VAP and HCAP/HAP studies is the dependence on expectorated sputum for a microbiologic diagnosis of VAP (as for that of CAP), which is further complicated by frequent colonization by pathogens in patients with HAP or HCAP. Therefore, most of the literature has focused on HCAP or HAP resulting in intubation, where, once again, access to the lower respiratory tract facilitates an etiologic diagnosis.

Etiology Potential etiologic agents of VAP include both MDR and non-MDR bacterial pathogens (**Table 153-6**). The non-MDR group is nearly identical to the pathogens found in severe CAP (**Table 153-2**); it is not surprising that such pathogens predominate if VAP develops in the first 5–7 days of the hospital stay. However, if patients have other risk factors for HCAP, MDR pathogens are a consideration, even early in the hospital course. The relative frequency of individual MDR pathogens can vary significantly from hospital to hospital and even between different critical care units within the same institution. Most hospitals have problems with *P. aeruginosa* and MRSA, but other MDR pathogens are often institution-specific. Less commonly, fungal and viral pathogens cause VAP, usually affecting severely immunocompromised patients. Rarely, community-associated viruses cause mini-epidemics, usually when introduced by ill health care workers.

TABLE 153-6 MICROBIOLOGIC CAUSES OF VENTILATOR-ASSOCIATED PNEUMONIA

Non-MDR Pathogens	MDR Pathogens
<i>Streptococcus pneumoniae</i>	<i>Pseudomonas aeruginosa</i>
Other <i>Streptococcus</i> spp.	MRSA
<i>Haemophilus influenzae</i>	<i>Acinetobacter</i> spp.
MSSA	Antibiotic-resistant
Antibiotic-sensitive	Enterobacteriaceae
Enterobacteriaceae	<i>Enterobacter</i> spp.
<i>Escherichia coli</i>	ESBL-positive strains
<i>Klebsiella pneumoniae</i>	<i>Klebsiella</i> spp.
<i>Proteus</i> spp.	<i>Legionella pneumophila</i>
<i>Enterobacter</i> spp.	<i>Burkholderia cepacia</i>
<i>Serratia marcescens</i>	<i>Aspergillus</i> spp.

Abbreviations: ESBL, extended-spectrum β -lactamase; MDR, multidrug-resistant; MRSA, methicillin-resistant *Staphylococcus aureus*; MSSA, methicillin-sensitive *S. aureus*.

810 Epidemiology Pneumonia is a common complication among patients requiring mechanical ventilation. Prevalence estimates vary between 6 and 52 cases per 100 patients, depending on the population studied. On any given day in the ICU, an average of 10% of patients will have pneumonia—VAP in the overwhelming majority of cases. The frequency of diagnosis is not static but changes with the duration of mechanical ventilation, with the highest hazard ratio in the first 5 days and a plateau in additional cases (1% per day) after ~2 weeks. However, the cumulative rate among patients who remain ventilated for as long as 30 days is as high as 70%. These rates often do not reflect the recurrence of VAP in the same patient. Once a ventilated patient is transferred to a chronic-care facility or to home, the incidence of pneumonia drops significantly, especially in the absence of other risk factors for pneumonia. However, in chronic ventilator units, purulent tracheobronchitis becomes a significant issue, often interfering with efforts to wean patients off mechanical ventilation (**Chap. 323**).

Three factors are critical in the pathogenesis of VAP: colonization of the oropharynx with pathogenic microorganisms, aspiration of these organisms from the oropharynx into the lower respiratory tract, and compromise of the normal host defense mechanisms. Most risk factors and their corresponding prevention strategies pertain to one of these three factors (**Table 153-7**).

The most obvious risk factor is the endotracheal tube, which bypasses the normal mechanical factors preventing aspiration. While the presence of an endotracheal tube may prevent large-volume aspiration, microaspiration is actually exacerbated by secretions pooling above the cuff. The endotracheal tube and the concomitant need

for suctioning can damage the tracheal mucosa, thereby facilitating tracheal colonization. In addition, pathogenic bacteria can form a glycocalyx biofilm on the tube's surface that protects them from both antibiotics and host defenses. The bacteria can also be dislodged during suctioning and can reinoculate the trachea, or tiny fragments of glycocalyx can embolize to distal airways, carrying bacteria with them.

In a high percentage of critically ill patients, the normal oropharyngeal flora is replaced by pathogenic microorganisms. The most important risk factors are antibiotic selection pressure, cross-infection from other infected/colonized patients or contaminated equipment, and malnutrition. Of these factors, antibiotic exposure poses the greatest risk by far. Pathogens such as *P. aeruginosa* almost never cause infection in patients without prior exposure to antibiotics. The recent emphasis on hand hygiene has lowered the cross-infection rate.

How the lower respiratory tract defenses become overwhelmed remains poorly understood. Almost all intubated patients experience microaspiration and are at least transiently colonized with pathogenic bacteria. However, only around one-third of colonized patients develop VAP. Colony counts increase to high levels, sometimes days before the development of clinical pneumonia; these increases suggest that the final step in VAP development, independent of aspiration and oropharyngeal colonization, is the overwhelming of host defenses. Severely ill patients with sepsis and trauma appear to enter a state of immunoparalysis several days after admission to the ICU—a time that corresponds to the greatest risk of developing VAP. The mechanism of this immunosuppression is not clear, although several factors have been suggested. Hyperglycemia affects neutrophil function, and trials suggest that keeping the blood sugar level close to normal with exogenous insulin may have beneficial effects, including a decreased risk of infection. More frequent transfusions also adversely affect the immune response.

Clinical Manifestations The clinical manifestations are generally the same in VAP as in all other forms of pneumonia: fever, leukocytosis, increase in respiratory secretions, and pulmonary consolidation on physical examination, along with a new or changing radiographic infiltrate. The frequency of abnormal chest radiographs before the onset of pneumonia in intubated patients and the limitations of portable radiographic technique make interpretation of radiographs more difficult than in patients who are not intubated. Other clinical features may include tachypnea, tachycardia, worsening oxygenation, and increased minute ventilation.

Diagnosis No single set of criteria is reliably diagnostic of pneumonia in a ventilated patient. The inability to identify such patients compromises efforts to prevent and treat VAP and even calls into question estimates of the impact of VAP on mortality rates.

Application of clinical criteria consistently results in overdiagnosis of VAP, largely because of three common findings in at-risk patients: (1) tracheal colonization with pathogenic bacteria in patients with endotracheal tubes, (2) multiple alternative causes of radiographic infiltrates in mechanically ventilated patients, and (3) the high frequency of other sources of fever in critically ill patients. The differential diagnosis of VAP includes a number of entities such as atypical pulmonary edema, pulmonary contusion, alveolar hemorrhage, hypersensitivity pneumonitis, acute respiratory distress syndrome, and pulmonary embolism. Clinical findings in ventilated patients with fever and/or leukocytosis may have alternative causes, including antibiotic-associated diarrhea, sinusitis, urinary tract infection, pancreatitis, and drug fever. Conditions mimicking pneumonia are often documented in patients in whom VAP has been ruled out by accurate diagnostic techniques. Most of these alternative diagnoses do not require antibiotic treatment; require antibiotics different from those used to treat VAP; or require some additional intervention, such as surgical drainage or catheter removal, for optimal management.

This diagnostic dilemma has led to debate and controversy. The major question is whether a quantitative-culture approach as a means of eliminating false-positive clinical diagnoses is superior to the clinical approach

TABLE 153-7 PATHOGENIC MECHANISMS AND CORRESPONDING PREVENTION STRATEGIES FOR VENTILATOR-ASSOCIATED PNEUMONIA

Pathogenic Mechanism	Prevention Strategy
Oropharyngeal colonization with pathogenic bacteria	
Elimination of normal flora	Avoidance of prolonged antibiotic courses
Large-volume oropharyngeal aspiration around time of intubation	Short course of prophylactic antibiotics for comatose patients ^a
Gastroesophageal reflux	Postpyloric enteral feeding ^b ; avoidance of high gastric residuals, prokinetic agents
Bacterial overgrowth of stomach	Avoidance of prophylactic agents that raise gastric pH ^b ; selective decontamination of digestive tract with nonabsorbable antibiotics ^b
Cross-infection from other colonized patients	Hand washing, especially with alcohol-based hand rub; intensive infection control education ^c ; isolation; proper cleaning of reusable equipment
Large-volume aspiration	Endotracheal intubation; rapid-sequence intubation technique; avoidance of sedation; decompression of small-bowel obstruction
Microaspiration around endotracheal tube	
Endotracheal intubation	Noninvasive ventilation ^a
Prolonged duration of ventilation	Daily awakening from sedation, ^a weaning protocols ^a
Abnormal swallowing function	Early percutaneous tracheostomy ^a
Secretions pooled above endotracheal tube	Head of bed elevated ^c ; continuous aspiration of subglottic secretions with specialized endotracheal tube ^e ; avoidance of reintubation; minimization of sedation and patient transport
Altered lower respiratory host defenses	Tight glycemic control ^b ; lowering of hemoglobin transfusion threshold

^aStrategies demonstrated to be effective in at least one randomized controlled trial.

^bStrategies with negative randomized trials or conflicting results.

enhanced by principles learned from quantitative-culture studies. The most recent IDSA/ATS guidelines for HAP/VAP suggest that either approach is clinically valid.

QUANTITATIVE-CULTURE APPROACH The essence of the quantitative-culture approach is to discriminate between colonization and true infection by determining the bacterial burden. The more distal in the respiratory tree the diagnostic sampling, the more specific the results and therefore the lower the threshold of growth necessary to diagnose pneumonia and exclude colonization. For example, a quantitative endotracheal aspirate yields proximate samples, and the diagnostic threshold is 10^6 cfu/mL. The protected specimen brush method, in contrast, obtains distal samples and has a threshold of 10^3 cfu/mL. Conversely, sensitivity declines as more distal secretions are obtained, especially when they are collected blindly (i.e., by a technique other than bronchoscopy). Additional tests that may increase the diagnostic yield include Gram's staining, differential cell counts, staining for intracellular organisms, and detection of local protein levels elevated in response to infection.

The Achilles heel of the quantitative approach is the effect of antibiotic therapy. With sensitive microorganisms, a single antibiotic dose can reduce colony counts below the diagnostic threshold. Recent changes in antibiotic therapy are the most significant. After 3 days, the operating characteristics of the tests are almost the same as if no antibiotic therapy has been given. Conversely, colony counts above the diagnostic threshold during antibiotic therapy suggest that the current antibiotics are ineffective. Even the normal host response may be sufficient to reduce quantitative-culture counts below the diagnostic threshold if sampling is delayed. In short, expertise in quantitative-culture techniques is critical, with a specimen obtained as soon as pneumonia is suspected and before antibiotic therapy is initiated or changed.

In a study comparing the quantitative with the clinical approach, use of bronchoscopic quantitative cultures resulted in significantly less antibiotic use at 14 days after study entry and in lower rates of mortality and severity-adjusted mortality at 28 days. In addition, more alternative sites of infection were found in patients randomized to the quantitative-culture strategy. A critical aspect of this study was that antibiotic treatment was initiated only in patients whose gram-stained respiratory sample was positive or who displayed signs of hemodynamic instability. Fewer than one-half as many patients were treated for pneumonia in the bronchoscopy group, and only one-third as many microorganisms were cultured.

CLINICAL APPROACH The lack of specificity of a clinical diagnosis of VAP has led to efforts to improve the diagnostic criteria. The Clinical Pulmonary Infection Score (CPIS) was developed by weighting of the various clinical criteria usually used for the diagnosis of VAP (Table 153-8). Use of the CPIS allows the selection of low-risk patients who may need only short-course antibiotic therapy or no treatment at all. Moreover, studies have demonstrated that the absence of bacteria in gram-stained endotracheal aspirates makes pneumonia an unlikely cause of fever or pulmonary infiltrates. These findings, coupled with a heightened awareness of the alternative diagnoses possible in patients with suspected VAP, can prevent inappropriate overtreatment for pneumonia. Furthermore, data show that the absence of an MDR pathogen in tracheal aspirate cultures eliminates the need for MDR coverage when empirical antibiotic therapy is narrowed. Since the most likely explanations for the mortality benefit of bronchoscopic quantitative cultures are decreased antibiotic selection pressure (which reduces the risk of subsequent infection with MDR pathogens) and identification of alternative sources of infection, a clinical diagnostic approach that incorporates such principles may result in similar outcomes.

Other large randomized studies that did not demonstrate a similar beneficial impact of quantitative culture on outcomes did not tightly link antibiotic treatment to the results of quantitative culture and other tests. Given the conflicting results only partially explained by methodologic issues, the IDSA/ATS guidelines therefore suggest that the choice depends on availability and local expertise.

TABLE 153-8 CLINICAL PULMONARY INFECTION SCORE (CPIS)

Criterion	Score
Fever (°C)	
≥38.5 but ≤38.9	1
>39 or <36	2
Leukocytosis	
<4000 or >11,000/ μ L	1
Bands >50%	1 (additional)
Oxygenation (mmHg)	
Pa _o ₂ /FI _o ₂ <250 and no ARDS	2
Chest radiograph	
Localized infiltrate	2
Patchy or diffuse infiltrate	1
Progression of infiltrate (no ARDS or CHF)	2
Tracheal aspirate	
Moderate or heavy growth	1
Same morphology on Gram's stain	1 (additional)
Maximal score ^a	12

^aAt the time of the original diagnosis, the progression of the infiltrate is not known and results of tracheal aspirate culture are often unavailable; thus, the maximal score is initially 8–10.

Abbreviations: ARDS, acute respiratory distress syndrome; CHF, congestive heart failure.

TREATMENT VENTILATOR-ASSOCIATED PNEUMONIA

Many studies have demonstrated higher mortality rates with initially inappropriate empirical antibiotic therapy. The key to appropriate antibiotic management of VAP is an appreciation of the resistance patterns of the most likely pathogens in a given patient.

ANTIBIOTIC RESISTANCE

If not for the higher risk of infection with MDR pathogens (Table 153-1), VAP could be treated with the same antibiotics used for severe CAP. However, antibiotic selection pressure leads to the frequent involvement of MDR pathogens by selecting either for drug-resistant isolates of common pathogens (MRSA and Enterobacteriaceae producing extended-spectrum β -lactamases or carbapenemases) or for intrinsically resistant pathogens (*P. aeruginosa* and *Acinetobacter* species). Frequent use of β -lactam drugs, especially cephalosporins, appears to be the major risk factor for infection with MRSA and extended-spectrum β -lactamase-positive strains.

P. aeruginosa has demonstrated the ability to develop resistance to all routinely used antibiotics. Unfortunately, even if initially sensitive, *P. aeruginosa* isolates also have a propensity to develop resistance during treatment. Either de-repression of resistance genes or selection of resistant clones within the large bacterial inoculum associated with most pneumonias may be the cause. *Acinetobacter* species, *Stenotrophomonas maltophilia*, and *Burkholderia cepacia* are intrinsically resistant to many of the empirical antibiotic regimens employed (see later in this chapter). VAP caused by these pathogens emerges during treatment of other infections, and resistance is always evident at initial diagnosis.

EMPIRICAL THERAPY

Recommended options for empirical therapy are listed in Table 153-9. Treatment should be started once diagnostic specimens have been obtained. The major factor in the selection of agents is the presence of risk factors for MDR pathogens. Choices among the various options listed depend on local patterns of resistance and—a very important factor—the patient's prior antibiotic exposure.

The majority of patients *without* risk factors for MDR infection can be treated with a single agent. The major difference from CAP is the markedly lower incidence of atypical pathogens in VAP; the exception is *Legionella*, which can be a nosocomial pathogen, especially with breakdowns in the treatment of potable water in the hospital.

TABLE 153-9 EMPIRICAL ANTIBIOTIC TREATMENT OF HEALTH CARE–ASSOCIATED PNEUMONIA

Patients without Risk Factors for MDR Pathogens	
Ceftriaxone (2 g IV q24h) or cefotaxime (2 g IV q6–8 h) or	
Moxifloxacin (400 mg IV q24h), ciprofloxacin (400 mg IV q8h), or levofloxacin (750 mg IV q24h) or	
Ampicillin/sulbactam (3 g IV q6h) or	
Ertapenem (1 g IV q24h)	
Patients with Risk Factors for MDR Pathogens	
1. A β -lactam:	
	Ceftazidime (2 g IV q8h) or ceftipime (2 g IV q8–12h) or
	Piperacillin/tazobactam (4.5 g IV q6h) or
	Imipenem (500 mg IV q6h or 1 g IV q8h), or meropenem (1 g IV q8h)
	plus
2. A second agent active against gram-negative bacterial pathogens:	
	Gentamicin or tobramycin (7 mg/kg IV q24h) or amikacin (20 mg/kg IV q24h) or
	Ciprofloxacin (400 mg IV q8h) or levofloxacin (750 mg IV q24h)
	plus
3. An agent active against gram-positive bacterial pathogens:	
	Linezolid (600 mg IV q12h) or
	Vancomycin (15 mg/kg q12h initially with adjusted doses)

Abbreviation: MDR, multidrug-resistant.

The standard recommendation for patients *with* risk factors for MDR infection is for three antibiotics: two directed at *P. aeruginosa* and one at MRSA. The choice of a β -lactam agent provides the greatest variability in coverage, yet the use of the broadest-spectrum agent—a carbapenem, even in an antibiotic combination—still represents inappropriate initial therapy in 10–15% of cases.

SPECIFIC TREATMENT

Once an etiologic diagnosis is made, broad-spectrum empirical therapy can be modified to specifically address the known pathogen. For patients with MDR risk factors, antibiotic regimens can be reduced to a single agent in more than one-half of cases and to a two-drug combination in more than one-quarter of cases. Only a minority of cases require a complete course with three drugs. A negative tracheal-aspirate culture or growth below the threshold for quantitative cultures, especially if the sample was obtained before any antibiotic change, strongly suggests that antibiotics should be discontinued. Identification of other confirmed or suspected sites of infection may require ongoing antibiotic therapy, but the spectrum of pathogens (and the corresponding antibiotic choices) may be different from those for VAP. If the CPIS decreases over the first 3 days, antibiotics should be stopped after 8 days. An 8-day course of therapy is just as effective as a 2-week course and is associated with less frequent emergence of antibiotic-resistant strains.

The major controversy regarding specific therapy for VAP concerns the need for ongoing combination treatment of *Pseudomonas* infection. No randomized controlled trials have demonstrated a benefit of combination therapy with a β -lactam and an aminoglycoside, nor have subgroup analyses in other trials found a survival benefit with such a regimen. The unacceptably high rates of clinical failure and death for VAP caused by *P. aeruginosa* despite combination therapy (see “Failure to Improve,” later) indicate that better regimens are needed—including, perhaps, aerosolized antibiotics. VAP caused by MRSA is associated with a 40% clinical failure rate when treated with standard-dose vancomycin. One proposed solution is the use of high-dose individualized treatment, although the risk of renal toxicity increases with this strategy. In addition, the MIC of vancomycin has been increasing, and a high percentage of clinical failures occur when the MIC is in the upper range of sensitivity (i.e., 1.5–2 $\mu\text{g}/\text{mL}$). Linezolid appears to be 15% more efficacious than even adjusted-dose vancomycin and is clearly preferred in

patients with renal insufficiency and those infected with high-MIC isolates of MRSA.

FAILURE TO IMPROVE

Treatment failure is not uncommon in VAP, especially that caused by MDR pathogens. In addition to the 40% failure rate for MRSA infection treated with vancomycin, VAP due to *Pseudomonas* has a 50% failure rate, no matter what the regimen. Causes of clinical failure vary with the pathogen(s) and the antibiotic(s). Inappropriate therapy can usually be minimized by use of the recommended triple-drug regimen (Table 153-9). However, the emergence of β -lactam resistance during therapy is an important problem, especially in infection with *Pseudomonas* and *Enterobacter* species. Recurrent VAP caused by the same pathogen is possible because the biofilm on endotracheal tubes allows reintroduction of the microorganism. However, studies of VAP caused by *Pseudomonas* show that approximately half of recurrent cases are caused by a new strain. Inadequate local levels of vancomycin are the likely cause of treatment failure in VAP due to MRSA.

Treatment failure is very difficult to diagnose. Pneumonia due to a new superinfection, the presence of extrapulmonary infection, and drug toxicity must be considered in the differential diagnosis of treatment failure. Serial CPIS calculations appear to track the clinical response accurately, while repeat quantitative cultures may clarify the microbiologic response. A persistently elevated or rising CPIS by day 3 of therapy is likely to indicate treatment failure. The most sensitive component of the CPIS is improvement in oxygenation.

COMPLICATIONS

Apart from death, the major complication of VAP is prolongation of mechanical ventilation, with corresponding increases in length of stay in the ICU and in the hospital. In most studies, an additional week of mechanical ventilation resulting from VAP is common. The additional expense of this complication often warrants costly and aggressive efforts at prevention.

In rare cases, some types of necrotizing pneumonia (e.g., that due to *P. aeruginosa*) result in significant pulmonary hemorrhage. More commonly, necrotizing infections result in the long-term complications of bronchiectasis and parenchymal scarring leading to recurrent pneumonias. The long-term complications of pneumonia are underappreciated. Pneumonia results in a catabolic state in a patient already nutritionally at risk. The muscle loss and general debilitation from an episode of VAP often require prolonged rehabilitation and, in the elderly, commonly result in an inability to return to independent function and the need for nursing home placement.

FOLLOW-UP

Clinical improvement, if it occurs, is usually evident within 48–72 h of the initiation of antimicrobial treatment. Because findings on chest radiography often worsen initially during treatment, they are less helpful than clinical criteria as an indicator of clinical response in severe pneumonia. Seriously ill patients with pneumonia often undergo follow-up chest radiography daily, at least until they are being weaned off mechanical ventilation.

Prognosis VAP is associated with significant mortality. Crude mortality rates of 50–70% have been reported, but the real issue is attributable mortality. Many patients with VAP have underlying diseases that would result in death even if VAP did not occur. Attributable mortality exceeded 25% in one matched-cohort study, while more recent studies have suggested much lower rates. Patients who develop VAP are at least twice as likely to die as those who do not. Some of the variability in VAP mortality rates is clearly related to the type of patient and ICU studied. VAP in trauma patients is not associated with attributable mortality, possibly because many of the patients were otherwise healthy before being injured. However, the causative pathogen also plays a major role. Generally, MDR pathogens are associated with

significantly greater attributable mortality than non-MDR pathogens. Pneumonia caused by some pathogens (e.g., *S. maltophilia*) is simply a marker for a patient whose immune system is so compromised that death is almost inevitable.

Prevention (Table 153-7) Because of the significance of the endotracheal tube as a risk factor for VAP, the most important preventive intervention is to avoid endotracheal intubation or minimize its duration. Successful use of noninvasive ventilation via a nasal or full-face mask avoids many of the problems associated with endotracheal tubes. Strategies that minimize the duration of ventilation through daily holding of sedation and formal weaning protocols also have been highly effective in preventing VAP.

Unfortunately, a tradeoff in risks is sometimes required. Aggressive attempts to extubate early may result in reintubation(s) and increase aspiration, posing a risk of VAP. Heavy continuous sedation increases the risk, but self-extubation because of insufficient sedation also is a risk. The tradeoffs also apply to antibiotic therapy. Short-course antibiotic prophylaxis can decrease the risk of VAP in comatose patients requiring intubation, and data suggest that antibiotics decrease VAP rates in general. However, the major benefit appears to be a decrease in the incidence of early-onset VAP, which is usually caused by the less pathogenic non-MDR microorganisms. Conversely, prolonged courses of antibiotics consistently increase the risk of VAP caused by the more lethal MDR pathogens. Despite its virulence and associated mortality, VAP caused by *Pseudomonas* is rare among patients who have not recently received antibiotics.

Minimizing the amount of microaspiration around the endotracheal tube cuff also is a strategy for avoidance of VAP. Simply elevating the head of the bed (at least 30° above horizontal but preferably 45°) decreases VAP rates. Specially modified endotracheal tubes that allow removal of the secretions pooled above the cuff also may prevent VAP. The risk-to-benefit ratio of transporting the patient outside the ICU for diagnostic tests or procedures should be carefully considered, since VAP rates are increased among transported patients.

Emphasis on the avoidance of agents that raise gastric pH and on oropharyngeal decontamination has been diminished by the equivocal and conflicting results of recent clinical trials. The role in the pathogenesis of VAP that is played by the overgrowth of bacterial components of the bowel flora in the stomach also has been downplayed. MRSA and the nonfermenters *P. aeruginosa* and *Acinetobacter* species are not normally part of the bowel flora but reside primarily in the nose and on the skin, respectively. Therefore, emphasis on controlling overgrowth of the bowel flora may be relevant only in certain populations, such as liver transplant recipients and patients who have undergone other major intraabdominal procedures or who have bowel obstruction.

In outbreaks of VAP due to specific pathogens, the possibility of a breakdown in infection control measures (particularly contamination of reusable equipment) should be investigated. Even high rates of pathogens that are already common in a particular ICU may be a result of cross-infection. Education and reminders of the need for consistent hand washing and other infection-control practices can minimize this risk.

HOSPITAL-ACQUIRED PNEUMONIA

While significantly less well studied than VAP, HAP in nonintubated patients—both inside and outside the ICU—is similar to VAP. The main differences are the higher frequency of non-MDR pathogens and the better underlying host immunity in nonintubated patients. The lower frequency of MDR pathogens allows monotherapy in a larger proportion of cases of HAP than of VAP.

The only pathogens that may be more common in the non-VAP population are anaerobes. The greater risk of macroaspiration by nonintubated patients and the lower oxygen tensions in the lower respiratory tract of these patients increase the likelihood of a role for anaerobes. While more common in patients with HAP, anaerobes are usually only contributors to polymicrobial pneumonias except in patients with large-volume aspiration or in the setting of bowel obstruction/ileus. As in the management of CAP, specific therapy

targeting anaerobes probably is not indicated (unless gross aspiration is a concern) since many of the recommended antibiotics are active against anaerobes.

Diagnosis is even more difficult for HAP in the nonintubated patient than for VAP. Lower respiratory tract samples appropriate for culture are considerably more difficult to obtain from nonintubated patients. Many of the underlying diseases that predispose a patient to HAP are also associated with an inability to cough adequately. Since blood cultures are infrequently positive (<15% of cases), the majority of patients with HAP do not have culture data on which antibiotic modifications can be based. Therefore, de-escalation of therapy is less likely in patients with risk factors for MDR pathogens. Despite these difficulties, the better host defenses in non-ICU patients result in lower mortality rates than are documented for VAP. In addition, the risk of antibiotic failure is lower in HAP.

154 Lung Abscess

Rebecca M. Baron, Miriam Baron Barshak

Lung abscess represents necrosis and cavitation of the lung following microbial infection. Lung abscesses can be single or multiple but usually are marked by a single dominant cavity >2 cm in diameter.

ETIOLOGY

The low prevalence of lung abscesses makes them difficult to study in randomized controlled trials. Although the incidence of lung abscesses has decreased in the postantibiotic era, they are still a source of significant morbidity and mortality.

Lung abscesses are usually characterized as either primary (~80% of cases) or secondary. *Primary* lung abscesses usually arise from aspiration, are often caused principally by anaerobic bacteria, and occur in the absence of an underlying pulmonary or systemic condition. *Secondary* lung abscesses arise in the setting of an underlying condition, such as a postobstructive process (e.g., a bronchial foreign body or tumor) or a systemic process (e.g., HIV infection or another immunocompromising condition). Lung abscesses can also be characterized as acute (<4–6 weeks in duration) or chronic (~40% of cases).

EPIDEMIOLOGY

The majority of the existing epidemiologic information involves primary lung abscesses. In general, middle-aged men are more commonly affected than middle-aged women. The major risk factor for primary lung abscesses is aspiration. Patients at particular risk for aspiration, such as those with altered mental status, alcoholism, drug overdose, seizures, bulbar dysfunction, prior cerebrovascular or cardiovascular events, or neuromuscular disease, are most commonly affected. In addition, patients with esophageal dysmotility or esophageal lesions (strictures or tumors) and those with gastric distention and/or gastroesophageal reflux, especially those who spend substantial time in the recumbent position, are at risk for aspiration.

It is widely thought that colonization of the gingival crevices by anaerobic bacteria or microaerophilic streptococci (especially in patients with gingivitis and periodontal disease), combined with a risk of aspiration, is important in the development of lung abscesses. In fact, many physicians consider it extremely rare for lung abscesses to develop in the absence of teeth as a nidus for bacterial colonization.

The importance of these risk factors in the development of lung abscesses is highlighted by a significant reduction in abscess incidence in the late 1940s that coincided with a change in oral surgical technique: beginning at that time, these operations were no longer performed with the patient in the seated position without a cuffed endotracheal tube, and the frequency of perioperative aspiration events was thus decreased. In addition, the introduction of penicillin around

TABLE 154-1 EXAMPLES OF MICROBIAL PATHOGENS THAT CAN CAUSE LUNG ABSCESSSES

Clinical Condition	Pathogens
Primary lung abscess (usually with risk factors for aspiration)	Anaerobes (e.g., <i>Peptostreptococcus</i> spp., <i>Prevotella</i> spp., <i>Bacteroides</i> spp., <i>Streptococcus milleri</i>), microaerophilic streptococci
Secondary lung abscess (often with underlying immunocompromise)	<i>Staphylococcus aureus</i> , gram-negative rods (e.g., <i>Pseudomonas aeruginosa</i> , Enterobacteriaceae), <i>Nocardia</i> spp., <i>Aspergillus</i> spp., Mucorales, <i>Cryptococcus</i> spp., <i>Legionella</i> spp., <i>Rhodococcus equi</i> , <i>Pneumocystis jirovecii</i>
Embolitic lesions	<i>Staphylococcus aureus</i> (often from endocarditis), <i>Fusobacterium necrophorum</i> (Lemierre's disease; see text for details)
Endemic infections (with or without underlying immunocompromise)	<i>Mycobacterium tuberculosis</i> (as well as <i>Mycobacterium avium</i> and <i>Mycobacterium kansasii</i>), <i>Coccidioides</i> spp., <i>Histoplasma capsulatum</i> , <i>Blastomyces</i> spp., parasites (e.g., <i>Entamoeba histolytica</i> , <i>Paragonimus westermani</i> , <i>Strongyloides stercoralis</i>)
Miscellaneous conditions	Bacterial pathogen (often <i>S. aureus</i>) after influenza or other viral infection, <i>Actinomyces</i> spp.

the same time significantly reduced the incidence of and mortality rate from lung abscess.

PATHOGENESIS

Primary Lung Abscesses The development of primary lung abscesses is thought to originate when chiefly anaerobic bacteria (as well as microaerophilic streptococci) in the gingival crevices are aspirated into the lung parenchyma in a susceptible host (Table 154-1). Thus, patients who develop primary lung abscesses usually carry an overwhelming burden of aspirated material or are unable to clear the bacterial load. Pneumonitis develops initially (exacerbated in part by tissue damage caused by gastric acid); then, over a period of 7–14 days, the anaerobic bacteria produce parenchymal necrosis and cavitation whose extent depends on the host–pathogen interaction (Fig. 154-1). Anaerobes are thought to produce more extensive tissue necrosis in polymicrobial infections in which virulence factors of the various bacteria can act synergistically to cause more significant tissue destruction.

Secondary Lung Abscesses The pathogenesis of secondary abscesses depends on the predisposing factor. For example, in cases of bronchial

obstruction from malignancy or a foreign body, the obstructing lesion prevents clearance of oropharyngeal secretions, leading to abscess development. With underlying systemic conditions (e.g., immunosuppression after bone marrow or solid organ transplantation), impaired host defense mechanisms lead to increased susceptibility to development of lung abscesses caused by a broad range of pathogens, including opportunistic organisms (Table 154-1).

Lung abscesses also arise from septic emboli, either in tricuspid valve endocarditis (often involving *Staphylococcus aureus*) or in Lemierre's syndrome, in which an infection begins in the pharynx (classically involving *Fusobacterium necrophorum*) and then spreads to the neck and the carotid sheath (which contains the jugular vein) to cause septic thrombophlebitis.

PATHOLOGY AND MICROBIOLOGY

Primary Lung Abscesses In primary lung abscesses, the dependent segments (posterior upper lobes and superior lower lobes) are the most common locations, given the predisposition of aspirated materials to be deposited in these areas. Generally, the right lung is affected more commonly than the left because the right mainstem bronchus is less angulated. In secondary abscesses, the location of the abscess may vary with the underlying cause.

The microbiology of primary lung abscesses is often polymicrobial, primarily including anaerobic organisms as well as microaerophilic streptococci (Table 154-1). The retrieval and culture of anaerobes can be complicated by the contamination of samples with microbes from the oral cavity, the need for expeditious transport of the cultures to the laboratory, the need for early plating with special culture techniques, the prolonged time required for culture growth, and the need for collection of specimens prior to administration of antibiotics. When attention is paid to these factors, rates of recovery of specific isolates have been reported to be as high as 78%.

Because it is not clear that knowing the identity of the causative anaerobic isolate alters the response to treatment of a primary lung abscess, practice has shifted away from the use of specialized techniques to obtain material for culture, such as transtracheal aspiration and bronchoalveolar lavage with protected brush specimens that allow recovery of culture material while avoiding contamination from the oral cavity. When no pathogen is isolated from a primary lung abscess (which is the case as often as 40% of the time), the abscess is termed a *nonspecific lung abscess*, and the presence of anaerobes is often presumed. A *putrid lung abscess* refers to foul-smelling breath, sputum, or empyema and is essentially diagnostic of an anaerobic lung abscess.

Secondary Lung Abscesses In contrast, the microbiology of secondary lung abscesses can encompass quite a broad bacterial spectrum, with infection by *Pseudomonas aeruginosa* and other gram-negative rods most common. In addition, a broad array of pathogens can be identified in patients from certain endemic areas and in specific clinical scenarios (e.g., a significant incidence of fungal infections among immunosuppressed patients following bone marrow or solid organ transplantation). Because immunocompromised hosts and patients without the classic presentation of a primary lung abscess can be infected with a wide array of unusual organisms (Table 154-1), it is of special importance to obtain culture material in order to target therapy.

CLINICAL MANIFESTATIONS

Clinical manifestations may initially be similar to those of pneumonia, with fevers, cough, sputum production, and chest pain; a more chronic and indolent presentation that includes night sweats, fatigue, and anemia is often observed with anaerobic lung abscesses. A subset of patients with putrid lung abscesses may report discolored phlegm and foul-tasting or foul-smelling sputum. Patients with lung abscesses due to non-anaerobic organisms, such as *S. aureus*, may present with a more fulminant course characterized by high fevers and rapid progression.

Findings on physical examination may include fevers, poor dentition, and/or gingival disease as well as amphoric and/or cavernous breath sounds on lung auscultation. Additional findings may include digital clubbing and the absence of a gag reflex.

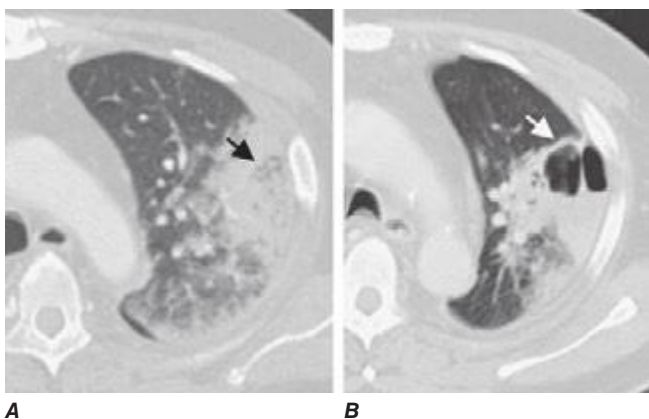


FIGURE 154-1 Representative chest CT scans demonstrating development of lung abscesses. This patient was immunocompromised due to underlying lymphoma and developed severe *Pseudomonas aeruginosa* pneumonia, as represented by a left lung infiltrate with concern for central regions of necrosis (panel A, black arrow). Two weeks later, areas of cavitation with air fluid levels were visible in this region and were consistent with the development of lung abscesses (panel B, white arrow). (Images provided by Dr. Ritu Gill, Division of Chest Radiology, Brigham and Women's Hospital, Boston.)

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of lung abscesses includes other noninfectious processes that result in cavitary lung lesions, including lung infarction, malignancy, sequestration, vasculitides (e.g., granulomatosis with polyangiitis), lung cysts or bullae containing fluid, and septic emboli (e.g., from tricuspid valve endocarditis).

DIAGNOSIS

The presence of a lung abscess is determined by chest imaging. Although a chest radiograph usually detects a thick-walled cavity with an air-fluid level, computed tomography (CT) permits better definition and may provide earlier evidence of cavitation. CT may also yield additional information regarding a possible underlying cause of lung abscess, such as malignancy, and may help distinguish a peripheral lung abscess from a pleural infection. This distinction has important implications for treatment, because a pleural space infection, such as an empyema, may require urgent drainage.

As described earlier (see “Pathology and Microbiology,” above), more invasive diagnostics (such as transtracheal aspiration) were traditionally undertaken for primary lung abscesses, whereas empirical therapy that includes drugs targeting anaerobic organisms currently is used more often. While sputum can be collected noninvasively for Gram’s stain and culture, which may yield a pathogen, it is likely that the infection will be polymicrobial, and culture results may not reflect the presence of anaerobic organisms. Many physicians consider putrid-smelling sputum to be virtually diagnostic of an anaerobic infection.

When a secondary lung abscess is present or empirical therapy fails to elicit a response, sputum and blood cultures are advised in addition to serologic studies for opportunistic pathogens (e.g., viruses and fungi causing infections in immunocompromised hosts). Additional diagnostics, such as bronchoscopy with bronchoalveolar lavage or protected brush specimen collection and CT-guided percutaneous needle aspiration, can be undertaken. Risks posed by these more invasive diagnostics include spillage of abscess contents into the other lung (with bronchoscopy) and pneumothorax and bronchopleural fistula development (with CT-guided needle aspiration). However, early diagnostics in secondary abscesses, especially in immunocompromised hosts, are particularly important, because the patients involved may be especially fragile and at risk for infection with a broad array of pathogens and, therefore, less likely than other patients to respond to empirical therapy.

TREATMENT LUNG ABSCESS

The availability of antibiotics in the 1940s and 1950s established therapy with this drug class as the primary approach to the treatment of lung abscess. Previously, surgery had been relied upon much more frequently. For many decades, penicillin was the antibiotic of choice for primary lung abscesses in light of its anaerobic coverage; however, because oral anaerobes can produce β -lactamases, clindamycin has proved superior to penicillin in clinical trials. For primary lung abscesses, the recommended regimens are (1) clindamycin (600 mg IV three times daily); then, with the disappearance of fever and clinical improvement, 300 mg PO four times daily) or (2) an IV-administered β -lactam/ β -lactamase combination, followed—once the patient’s condition is stable—by orally administered amoxicillin-clavulanate. This therapy should be continued until imaging demonstrates that the lung abscess has cleared or regressed to a small scar. Treatment duration may range from 3–4 weeks to as long as 14 weeks. One small study suggested that moxifloxacin (400 mg/d PO) is as effective and well tolerated as ampicillin-sulbactam. Notably, metronidazole is not effective as a single agent: it covers anaerobic organisms but not the microaerophilic streptococci that are often components of the mixed flora of primary lung abscesses.

In secondary lung abscesses, antibiotic coverage should be directed at the identified pathogen, and a prolonged course (until resolution of the abscess is documented) is often required. Treatment regimens and courses vary widely, depending on the immune state of the host and the identified pathogen. Other interventions may be necessary as well, such as relief of an obstructing lesion or treatment directed at the underlying condition predisposing the patient to lung abscess. Similarly, if the condition of patients with presumed primary lung abscess fails to improve, additional studies to rule out an underlying predisposing cause for a secondary lung abscess are indicated.

Although it can take as long as 7 days for patients receiving appropriate therapy to defervesce, as many as 10–20% of patients may not respond at all, with continued fevers and progression of the abscess cavity on imaging. An abscess >6–8 cm in diameter is less likely to respond to antibiotic therapy without additional interventions. Options for patients who do not respond to antibiotics and whose additional diagnostic studies fail to identify an additional pathogen that can be treated include surgical resection and percutaneous drainage of the abscess, especially in poor surgical candidates. Possible complications of percutaneous drainage include bacterial contamination of the pleural space as well as pneumothorax and hemothorax.

COMPLICATIONS

Larger cavity size on presentation may correlate with the development of persistent cystic changes (pneumatocoeles) or bronchiectasis. Additional possible complications include recurrence of abscesses despite appropriate therapy, extension to the pleural space with development of empyema, life-threatening hemoptysis, and massive aspiration of lung abscess contents.

PROGNOSIS AND PREVENTION

Reported mortality rates for primary abscesses have been as low as 2%, while rates for secondary abscesses are generally higher—as high as 75% in some case series. Other poor prognostic factors include an age >60, the presence of aerobic bacteria, sepsis at presentation, symptom duration of >8 weeks, and abscess size >6 cm.

Mitigation of underlying risk factors may be the best approach to prevention of lung abscesses, with attention directed toward airway protection, oral hygiene, and minimized sedation with elevation of the head of the bed for patients at risk for aspiration. Prophylaxis against certain pathogens in at-risk patients (e.g., recipients of bone marrow or solid organ transplants or patients whose immune systems are significantly compromised by HIV infection) may be undertaken.

APPROACH TO THE PATIENT: Lung Abscess

For patients with a lung abscess and a low likelihood of malignancy (e.g., smokers <45 years old) and with risk factors for aspiration, it is reasonable to administer empirical treatment and then to pursue further evaluation if therapy does not elicit a response. However, some clinicians may opt for up-front cultures, even in primary lung abscesses. In patients with risk factors for malignancy or other underlying conditions (especially immunocompromised hosts) or with an atypical presentation, earlier diagnostics should be considered, such as bronchoscopy with biopsy or CT-guided needle aspiration. Bronchoscopy should be performed early in patients whose history, symptoms, or imaging findings are consistent with possible bronchial obstruction. In patients from areas endemic for tuberculosis or patients with other risk factors for tuberculosis (e.g., underlying HIV infection), induced sputum samples should be examined early in the workup to rule out this disease.

155 Infective Endocarditis

Adolf W. Karchmer

The prototypic lesion of infective endocarditis, the *vegetation* (Fig. 155-1), is a mass of platelets, fibrin, microcolonies of microorganisms, and scant inflammatory cells. Infection most commonly involves heart valves but may also occur on the low-pressure side of a ventricular septal defect, on mural endocardium damaged by aberrant jets of blood or foreign bodies, or on intracardiac devices themselves. The analogous process involving arteriovenous shunts, arterio-arterial shunts (patent ductus arteriosus), or a coarctation of the aorta is called *infective endarteritis*.

Endocarditis can be classified according to the temporal evolution of disease, the site of infection, the cause of infection, or the predisposing risk factor (e.g., injection drug use). While each classification criterion provides therapeutic and prognostic insight, none is sufficient alone. *Acute endocarditis* is a hectically febrile illness that rapidly damages cardiac structures, seeds extracardiac sites, and, if untreated, progresses to death within weeks. *Subacute endocarditis* follows an indolent course; causes structural cardiac damage only slowly, if at all; rarely metastasizes; and is gradually progressive unless complicated by a major embolic event or a ruptured mycotic aneurysm.

In developed countries, the incidence of endocarditis ranges from 4 to 7 cases per 100,000 population per year and has remained relatively stable during recent decades. While congenital heart diseases remain a constant predisposition, predisposing conditions in developed countries have shifted from chronic rheumatic heart disease (still a common predisposition in developing countries) to illicit IV drug use, degenerative valve disease, and intracardiac devices. The incidence of endocarditis is notably increased among the elderly. In developed countries, 25–35% of cases of native valve endocarditis (NVE) are associated with health care, and 16–30% of all cases of endocarditis involve prosthetic valves. The risk of prosthesis infection is greatest during the first 6–12 months after valve replacement; gradually declines to a low, stable rate thereafter; and is similar for mechanical and bioprosthetic devices. The incidence of endocarditis involving cardiovascular implantable electronic devices (CIED), primarily permanent pacemakers and implantable cardioverter-defibrillators, ranges from 0.5 to 1.14 cases per 1000 device recipients and is higher among patients with an implantable cardioverter-defibrillator than among those with a permanent pacemaker.

ETIOLOGY

Although many species of bacteria and fungi cause sporadic episodes of endocarditis, a few bacterial species cause the majority of cases (Table 155-1). The oral cavity, skin, and upper respiratory tract are

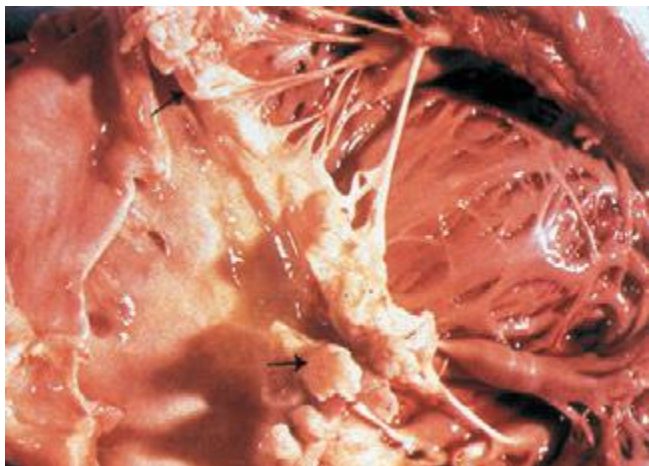


FIGURE 155-1 Vegetations (arrows) due to viridans streptococcal endocarditis involving the mitral valve.

the respective primary portals for viridans streptococci, staphylococci, and HACEK organisms (*Haemophilus* species, *Aggregatibacter aphrophilus*, *A. actinomycetemcomitans*, *Cardiobacterium* species, *Eikenella* species, and *Kingella* species). *Streptococcus gallolyticus* subspecies *gallolyticus* (formerly *S. bovis* biotype 1) originates from the gastrointestinal tract, where it is associated with polyps and colonic tumors, and enterococci enter the bloodstream from the genitourinary tract. Health care–associated NVE, most commonly caused by *Staphylococcus aureus*, coagulase-negative staphylococci (CoNS), and enterococci, may have either a nosocomial onset (55%) or a community onset (45%); community-onset cases develop in patients who have had extensive contact with the health care system over the preceding 90 days. Endocarditis complicates 6–25% of episodes of catheter-associated *S. aureus* bacteremia; the higher rates are detected in high-risk patients studied by transesophageal echocardiography (TEE) (see “Echocardiography,” later).

Prosthetic valve endocarditis (PVE) arising within 2 months of valve surgery is generally nosocomial, the result of intraoperative contamination of the prosthesis or a bacteremic postoperative complication. This nosocomial origin is reflected in the primary microbial causes: *S. aureus*, CoNS, facultative gram-negative bacilli, diphtheroids, and fungi. The portals of entry and organisms causing cases beginning >12 months after surgery are similar to those in community-acquired NVE. PVE due to CoNS that presents 2–12 months after surgery often represents delayed-onset nosocomial infection. Regardless of the time of onset after surgery, at least 68–85% of CoNS strains that cause PVE are resistant to methicillin.

Endocarditis related to a permanent pacemaker or an implantable cardioverter-defibrillator involves the device or the endothelium at points of device contact. Occasionally, there is concurrent aortic or mitral valve infection. One-third of cases of CIED endocarditis present within 3 months after device implantation or manipulation, one-third present at 4–12 months, and one-third present at >1 year. *S. aureus* and CoNS, both of which are commonly resistant to methicillin, cause the majority of cases.

Injection drug use–associated endocarditis, especially that involving the tricuspid valve, is commonly caused by *S. aureus*, which in many cases is resistant to methicillin. Left-sided valve infections in addicts have a more varied etiology. In addition to the usual causes of endocarditis, these cases can be due to *Pseudomonas aeruginosa* and *Candida* species, and sporadic cases can be caused by unusual organisms such as *Bacillus*, *Lactobacillus*, and *Corynebacterium* species. Polymicrobial endocarditis occurs among injection drug users. HIV infection in drug users does not significantly influence the causes of endocarditis.

From 5% to 15% of patients with endocarditis have negative blood cultures; in one-third to one-half of these cases, cultures are negative because of prior antibiotic exposure. The remainder of these patients are infected by fastidious organisms, such as nutritionally variant bacteria (now designated *Granulicatella* and *Abiotrophia* species), HACEK organisms, *Coxiella burnetii*, and *Bartonella* species. Some fastidious organisms occur in characteristic geographic settings (e.g., *C. burnetii* and *Bartonella* species in Europe, *Brucella* species in the Middle East). *Tropheryma whippelii* causes an indolent, culture-negative, afebrile form of endocarditis.

PATHOGENESIS

The undamaged endothelium is resistant to infection by most bacteria and to thrombus formation. Endothelial injury (e.g., at the site of impact of high-velocity blood jets or on the low-pressure side of a cardiac structural lesion) allows either direct infection by virulent organisms or the development of a platelet-fibrin thrombus—a condition called *nonbacterial thrombotic endocarditis* (NBTE). This thrombus serves as a site of bacterial attachment during transient bacteremia. The cardiac conditions most commonly resulting in NBTE are mitral regurgitation, aortic stenosis, aortic regurgitation, ventricular septal defects, and complex congenital heart disease. NBTE also arises as a result of a hypercoagulable state; this gives rise to *marantic endocarditis* (uninfected vegetations seen in patients with malignancy and chronic diseases) and to bland vegetations

TABLE 155-1 ORGANISMS CAUSING MAJOR CLINICAL FORMS OF ENDOCARDITIS

Organism	Percentage of Cases							
	Native Valve Endocarditis		Prosthetic Valve Endocarditis at Indicated Time of Onset (Months) after Valve Surgery			Endocarditis in Injection Drug Users		
	Community-Acquired (n = 1718)	Health Care-Associated (n = 1110)	<2 (n = 144)	2–12 (n = 31)	>12 (n = 194)	Right-Sided (n = 346)	Left-Sided (n = 204)	Total (n = 675) ^a
Streptococci ^b	40	13	1	9	31	5	15	12
Pneumococci	2	—	—	—	—	—	—	—
Enterococci ^c	9	16	8	12	11	2	24	9
<i>Staphylococcus aureus</i>	28	52 ^d	22	12	18	77	23	57
Coagulase-negative staphylococci	5	11	33	32	11	—	—	—
Fastidious gram-negative coccobacilli (HACEK group) ^e	3	—	—	—	6	—	—	—
Gram-negative bacilli	1	1	13	3	6	5	13	7
<i>Candida</i> spp.	<1	1	8	12	1	—	12	4
Polymicrobial/miscellaneous	3	3	3	6	5	8	10	7
Diphtheroids	—	<1	6	—	3	—	—	0.1
Culture-negative	9	3	5	6	8	3	3	3

^aThe total number of cases is larger than the sum of right- and left-sided cases because the location of infection was not specified in some cases. ^bIncludes viridans streptococci; *Streptococcus galloyticus*; other non-group A, groupable streptococci; and *Abiotrophia* and *Granulicatella* spp. (nutritionally variant, pyridoxal-requiring streptococci). ^cPrimarily *E. faecalis* or nonspiciated isolates; occasionally *E. faecium* or other, less likely species. ^dMethicillin resistance is common among these *S. aureus* strains. ^eIncludes *Haemophilus* spp., *Aggregatibacter aphrophilus*, *Aggregatibacter actinomycetemcomitans*, *Cardiobacterium hominis*, *Eikenella* spp., and *Kingella* spp.

Note: Data are compiled from multiple studies.

complicating systemic lupus erythematosus and the antiphospholipid antibody syndrome.

Organisms that cause endocarditis enter the bloodstream from mucosal surfaces, the skin, or sites of focal infection. Except for more virulent bacteria (e.g., *S. aureus*) that can adhere directly to intact endothelium or exposed subendothelial tissue, microorganisms in the blood adhere at sites of NBTE. The organisms that commonly cause endocarditis have surface adhesin molecules, collectively called microbial surface components recognizing adhesin matrix molecules (MSCRAMMs), that mediate adherence to NBTE sites or injured endothelium. Adherence is facilitated by fibronectin-binding proteins present on many gram-positive bacteria; by clumping factor (a fibrinogen- and fibrin-binding surface protein) on *S. aureus*; by fibrinogen-binding surface proteins (Fss2), collagen-binding surface protein (Ace), and Ebp pili (the latter mediating platelet adherence) in *Enterococcus faecalis*; and by glucans or FimA (a member of the family of oral mucosal adhesins) on streptococci. Fibronectin-binding proteins are required for *S. aureus* invasion of intact endothelium; thus these surface proteins may facilitate infection of previously normal valves. If resistant to the bactericidal activity of serum and the microbicidal peptides released locally by platelets, adherent organisms proliferate to form dense microcolonies. Microorganisms also induce platelet deposition and a localized procoagulant state by eliciting tissue factor from the endothelium or, in the case of *S. aureus*, from monocytes as well. Fibrin deposition combines with platelet aggregation and microorganism proliferation to generate an infected vegetation. Organisms deep in vegetations are metabolically inactive (nongrowing) and relatively resistant to killing by antimicrobial agents. Proliferating surface organisms are shed into the bloodstream continuously.

The clinical manifestations of endocarditis—other than constitutional symptoms, which probably result from cytokine production—arise from damage to intracardiac structures; embolization of vegetation fragments, leading to infection or infarction of remote tissues; hematogenous infection of sites during bacteremia; and tissue injury due to the deposition of circulating immune complexes or immune responses to deposited bacterial antigens.

CLINICAL MANIFESTATIONS

The clinical endocarditis syndrome is highly variable and spans a continuum between acute and subacute presentations. NVE, PVE, and endocarditis due to injection drug use share clinical and laboratory

manifestations (Table 155-2). The causative microorganism is primarily responsible for the temporal course of endocarditis. β -Hemolytic streptococci, *S. aureus*, and pneumococci typically result in an acute course, although *S. aureus* occasionally causes subacute disease. Endocarditis caused by *Staphylococcus lugdunensis* (a coagulase-negative species) or by enterococci may present acutely. Subacute endocarditis is typically caused by viridans streptococci, enterococci, CoNS, and the HACEK group. Endocarditis caused by *Bartonella* species, *T. whipplei*, or *C. burnetii* is exceptionally indolent.

TABLE 155-2 CLINICAL AND LABORATORY FEATURES OF INFECTIVE ENDOCARDITIS

Feature	Frequency, %
Fever	80–90
Chills and sweats	40–75
Anorexia, weight loss, malaise	25–50
Myalgias, arthralgias	15–30
Back pain	7–15
Heart murmur	80–85
New/worsened regurgitant murmur	20–50
Arterial emboli	20–50
Splenomegaly	15–50
Clubbing	10–20
Neurologic manifestations	20–40
Peripheral manifestations (Osler's nodes, subungual hemorrhages, Janeway lesions, Roth's spots)	2–15
Petechiae	10–40
Laboratory manifestations	
Anemia	70–90
Leukocytosis	20–30
Microscopic hematuria	30–50
Elevated erythrocyte sedimentation rate	60–90
Elevated C-reactive protein level	>90
Rheumatoid factor	50
Circulating immune complexes	65–100
Decreased serum complement	5–40

818 In patients with subacute presentations, fever is typically low-grade and rarely exceeds 39.4°C (103°F); in contrast, temperatures of 39.4°–40°C (103°–104°F) are often noted in acute endocarditis. Fever may be blunted in patients who are elderly, are severely debilitated, or have renal failure.

Cardiac Manifestations Although heart murmurs are usually indicative of the predisposing cardiac pathology rather than of endocarditis, valvular damage and ruptured chordae may result in new regurgitant murmurs. In acute endocarditis involving a normal valve, murmurs may be absent initially but ultimately are detected in 85% of cases. Congestive heart failure (CHF) develops in 30–40% of patients as a consequence of valvular dysfunction. Occasionally, CHF is due to endocarditis-associated myocarditis or an intracardiac fistula. Heart failure due to aortic valve dysfunction progresses more rapidly than does that due to mitral valve dysfunction. Extension of infection beyond valve leaflets into adjacent annular or myocardial tissue results in perivalvular abscesses, which in turn may cause intracardiac fistulae with new murmurs. Abscesses may burrow from the aortic valve annulus through the epicardium, causing pericarditis, or into the upper ventricular septum, where they may interrupt the conduction system, leading to varying degrees of heart block. Mitral perivalvular abscesses, which are usually more distant from the conduction system, only rarely cause conduction abnormalities; if such abnormalities occur in this setting, the conduction pathway is most likely disrupted near the atrioventricular node or in the proximal bundle of His. Emboli to a coronary artery occur in 2% of patients and may result in myocardial infarction.

Noncardiac Manifestations The classic nonsuppurative peripheral manifestations of subacute endocarditis (e.g., Janeway lesions; Fig. 155-2A) are related to prolonged infection; with early diagnosis and treatment, these have become infrequent. In contrast, septic embolization mimicking some of these lesions (subungual hemorrhage, Osler's nodes) is common in patients with acute *S. aureus* endocarditis (Fig. 155-2B). Musculoskeletal pain usually remits promptly with treatment but must be distinguished from focal metastatic infections (e.g., spondylodiscitis), which may complicate 10–15% of cases. Hematogenously seeded focal infection occurs most often in the skin, spleen, kidneys, skeletal system, and meninges. Arterial emboli, one-half of which precede the diagnosis, are clinically apparent in up to 50% of patients. Endocarditis caused by *S. aureus*, vegetations >10 mm in diameter (as measured by echocardiography), and infection involving the mitral valve, especially the anterior leaflet, are independently associated with an increased risk of embolization. Symptoms, pain, or ischemia-induced dysfunction relate to the organ or area suffering embolic arterial occlusion (e.g., kidney, spleen, bowel, extremity). Cerebrovascular emboli presenting as strokes or occasionally as encephalopathy complicate 15–35% of cases of endocarditis. Again, one-half of these events precede the diagnosis of endocarditis. The frequency of stroke is 8 per 1000 patient-days during the week prior to diagnosis; the figure falls to 4.8 and 1.7 per 1000 patient-days during the first and second weeks of effective antimicrobial therapy, respectively. This decline exceeds that which can be attributed to change in vegetation size. Only 3% of strokes occur after 1 week of effective therapy. Emboli occurring late during or after effective therapy do not in themselves constitute evidence of failed antimicrobial treatment.



A



B

FIGURE 155-2 A. Janeway lesions on toe (left) and plantar surface (right) of the foot in subacute *Neisseria mucosa* endocarditis. (Image courtesy of Rachel Baden, MD.) B. Septic emboli with hemorrhage and infarction due to acute *Staphylococcus aureus* endocarditis.

TABLE 155-3 THE MODIFIED DUKE CRITERIA FOR THE CLINICAL DIAGNOSIS OF INFECTIVE ENDOCARDITIS^a

Major Criteria	
1. Positive blood culture	Typical microorganism for infective endocarditis from two separate blood cultures Viridans streptococci, <i>Streptococcus gallolyticus</i> , HACEK group organisms, <i>Staphylococcus aureus</i> , or Community-acquired enterococci in the absence of a primary focus, or Persistently positive blood culture, defined as recovery of a microorganism consistent with infective endocarditis from: Blood cultures drawn >12 h apart; or All of 3 or a majority of ≥4 separate blood cultures, with first and last drawn at least 1 h apart or Single positive blood culture for <i>Coxiella burnetii</i> or phase I IgG antibody titer of >1:800
2. Evidence of endocardial involvement	Positive echocardiogram ^b Oscillating intracardiac mass on valve or supporting structures or in the path of regurgitant jets or in implanted material, in the absence of an alternative anatomic explanation, or Abscess, or New partial dehiscence of prosthetic valve, or New valvular regurgitation (increase or change in preexisting murmur not sufficient)
Minor Criteria	
1. Predisposition: predisposing heart conditions ^c or injection drug use	
2. Fever ≥38.0°C (≥100.4°F)	
3. Vascular phenomena: major arterial emboli, septic pulmonary infarcts, mycotic aneurysm, intracranial hemorrhage, conjunctival hemorrhages, Janeway lesions	
4. Immunologic phenomena: glomerulonephritis, Osler's nodes, Roth's spots, rheumatoid factor	
5. Microbiologic evidence: positive blood culture but not meeting major criterion, as noted previously, ^d or serologic evidence of active infection with an organism consistent with infective endocarditis	

^aDefinite endocarditis is defined by documentation of two major criteria, of one major criterion and three minor criteria, or of five minor criteria. See text for further details. ^bTransesophageal echocardiography is required for optimal assessment of possible prosthetic valve endocarditis or complicated endocarditis. ^cValvular disease with stenosis or regurgitation, presence of a prosthetic valve, congenital heart disease including corrected or partially corrected conditions (except isolated atrial septal defect, repaired ventricular septal defect, or closed patent ductus arteriosus), prior endocarditis, or hypertrophic cardiomyopathy. ^dExcluding single positive cultures for coagulase-negative staphylococci and diphtheroids, which are common culture contaminants, or for organisms that do not cause endocarditis frequently, such as gram-negative bacilli.

Source: Adapted from JS Li et al: Clin Infect Dis 30:633, 2000. With permission from Oxford University Press.

Other neurologic complications include aseptic or purulent meningitis, intracranial hemorrhage due to hemorrhagic infarcts or ruptured mycotic aneurysms, and seizures. (*Mycotic aneurysms* are focal dilations of arteries occurring at points in the artery wall that have been weakened by infection in the vasa vasorum or where septic emboli have lodged.) Microabscesses in brain and meninges occur commonly in *S. aureus* endocarditis; surgically drainable intracerebral abscesses are infrequent.

Immune complex deposition on the glomerular basement membrane causes diffuse hypocomplementemic glomerulonephritis and renal dysfunction, which typically improve with effective antimicrobial therapy. Embolic renal infarcts cause flank pain and hematuria but rarely cause renal dysfunction.

Manifestations of Specific Predisposing Conditions Almost 50% of endocarditis associated with injection drug use is limited to the tricuspid valve and presents with fever but with faint or no murmur and no peripheral manifestations. Septic pulmonary emboli, which are common with tricuspid endocarditis, cause cough, pleuritic chest pain, nodular pulmonary infiltrates, or occasionally pyopneumothorax. Infection of the aortic or mitral valves presents with the typical clinical features of endocarditis, including peripheral manifestations.

If not associated with a retained intracardiac device or masked by the symptoms of concurrent comorbid illness, health care-associated endocarditis has typical manifestations. CIED endocarditis may be associated with obvious or cryptic generator pocket infection and results in fever, minimal murmur, and pulmonary symptoms due to septic emboli. Late-onset PVE presents with typical clinical features. In cases arising within 60 days of valve surgery (early onset), typical symptoms may be obscured by comorbidity associated with recent surgery. In both early-onset and more delayed presentations, paravalvular infection is common and often results in partial valve dehiscence, regurgitant murmurs, CHF, or disruption of the conduction system.

DIAGNOSIS

In order to avoid delayed or missed diagnosis, careful clinical, microbiologic, and echocardiographic evaluation should be pursued when febrile patients have endocarditis predispositions, cardiac or noncardiac features of endocarditis, or microbiologic findings consistent with endocarditis (e.g., a stroke or splenic infarct, multiple positive blood cultures for an endocarditis-associated organism).

The Duke Criteria The diagnosis of infective endocarditis is established with certainty only when vegetations are examined histologically and microbiologically. Nevertheless, a highly sensitive and specific diagnostic schema—known as the *modified Duke criteria*—is based on clinical, laboratory, and echocardiographic findings commonly encountered in patients with endocarditis (Table 155-3). While developed as a research tool rather than for patient management, the criteria can be a helpful diagnostic tool. If the criteria are to be maximally helpful in evaluating patients, appropriate data must be collected. Furthermore, clinical judgment must be exercised in order to use the criteria effectively. Documentation of two major criteria, of one major criterion and three minor criteria, or of five minor criteria allows a clinical diagnosis of definite endocarditis. The diagnosis of endocarditis is rejected if an alternative diagnosis is established, if symptoms resolve and do not recur with ≤4 days of antibiotic therapy, or if surgery or autopsy after ≤4 days of antimicrobial therapy yields no histologic evidence of endocarditis. Illnesses not classified as definite endocarditis or rejected as such are considered cases of possible infective endocarditis when either one major and one minor criterion or three minor criteria are fulfilled. Requiring some clinical features of endocarditis for classification as possible infective endocarditis increases the specificity of the schema without significantly reducing its sensitivity. Unless there are extenuating circumstances, patients with definite or possible endocarditis are treated as such.

The criteria emphasize bacteremia and echocardiographic findings typical of endocarditis. The requirement for multiple positive blood cultures over time is consistent with the continuous low-density

bacteremia characteristic of endocarditis. Among patients with untreated endocarditis who ultimately have a positive blood culture, 95% of all blood cultures are positive. The diagnostic criteria attach significance to the species of organism isolated from blood cultures. To fulfill a major criterion, the isolation of an organism that causes both endocarditis and bacteremia in the absence of endocarditis (e.g., *S. aureus*, enterococci) must take place repeatedly (i.e., persistent bacteremia) and in the absence of a primary focus of infection. Organisms that rarely cause endocarditis but commonly contaminate blood cultures (e.g., diphtheroids, CoNS) must be isolated repeatedly if their isolation is to serve as a major criterion.

Blood Cultures Isolation of the causative microorganism from blood cultures is critical for diagnosis and for planning treatment. In patients with suspected NVE, PVE, or CIED endocarditis who have not received antibiotics during the prior 2 weeks, three 2-bottle blood

820 culture sets, separated from one another by at least 2 h, should be obtained from different venipuncture sites over 24 h. If the cultures remain negative after 48–72 h, two or three additional blood culture sets should be obtained, and the laboratory should be consulted for advice regarding optimal culture techniques. Pending culture results, empirical antimicrobial therapy should be withheld initially from hemodynamically stable patients with suspected subacute endocarditis, especially those who have received antibiotics within the preceding 2 weeks. Thus, if necessary, additional blood culture sets can be obtained without the confounding effect of empirical treatment. Patients with acute endocarditis or with deteriorating hemodynamics who may require urgent surgery should receive empirical treatment immediately after three sets of blood cultures are obtained over several hours.

Non-Blood-Culture Tests Serologic tests can be used to implicate organisms that are difficult to recover by blood culture: *Brucella*, *Bartonella*, *Legionella*, *Chlamydia psittaci*, and *C. burnetii*. Pathogens can also be identified in vegetations by culture, microscopic examination with special stains (i.e., the periodic acid–Schiff stain for *T. whipplei*), or direct fluorescence antibody techniques and by the use of polymerase chain reaction to recover unique microbial DNA or DNA encoding the 16S or 28S ribosomal unit (16S rRNA or 28S rRNA); sequencing of these DNAs allows identification of bacteria and fungi, respectively.

Echocardiography Echocardiography anatomically confirms and measures vegetations, detects intracardiac complications, and assesses cardiac function (Fig. 155-3). Transthoracic echocardiography (TTE) is noninvasive and exceptionally specific; however, it cannot image vegetations <2 mm in diameter, and in 20% of patients it is technically inadequate because of emphysema or body habitus. TTE detects vegetations in 65–80% of patients with definite clinical endocarditis but is not optimal for evaluating prosthetic valves or detecting intracardiac complications. TEE is safe and detects vegetations in >90% of patients with definite endocarditis; nevertheless, initial studies may yield false-negative results in 6–18% of endocarditis patients. When endocarditis is likely, a negative TEE result does not exclude the diagnosis but rather warrants repetition of the study once or twice in 7–10 days. TEE is the optimal method for the diagnosis of PVE, the detection of myocardial abscess, valve perforation, or intracardiac fistulae and for the detection of vegetations in patients with CIED. In patients with CIED and negative blood cultures, a mass adherent to the lead is likely to be a bland thrombosis rather than an infected vegetation.

Because *S. aureus* bacteremia is associated with a high prevalence of endocarditis, routine echocardiographic evaluation (TTE or preferably TEE) is recommended in these patients. Patients with nosocomial *S. aureus* bacteremia are at increased risk of endocarditis if one or more of the following are present: positive blood cultures for 2–4 days, hemodialysis dependency, a permanent intracardiac device, spine infection, nonvertebral osteomyelitis, or an endocarditis-predisposing valve abnormality. Ideally, these patients should be evaluated with TEE. In patients with none of these findings, the risk of endocarditis is low and evaluation with TTE may suffice.

Experts favor echocardiographic evaluation of all patients with a clinical diagnosis of endocarditis; however, the test should not be used to screen patients with a low probability of endocarditis (e.g., patients with unexplained fever). An American Heart Association approach to the use of echocardiography for evaluation of patients with suspected endocarditis is illustrated in (Fig. 155-4).

Other Studies Many studies that are not diagnostic—i.e., complete blood count, creatinine determination, liver function tests, chest radiography, and electrocardiography—are important in the management of patients with endocarditis. The erythrocyte sedimentation rate, C-reactive protein level, and circulating immune complex titer are commonly increased in endocarditis (Table 155-2). Cardiac catheterization is useful primarily to assess coronary artery patency in older individuals who are to undergo surgery for endocarditis.

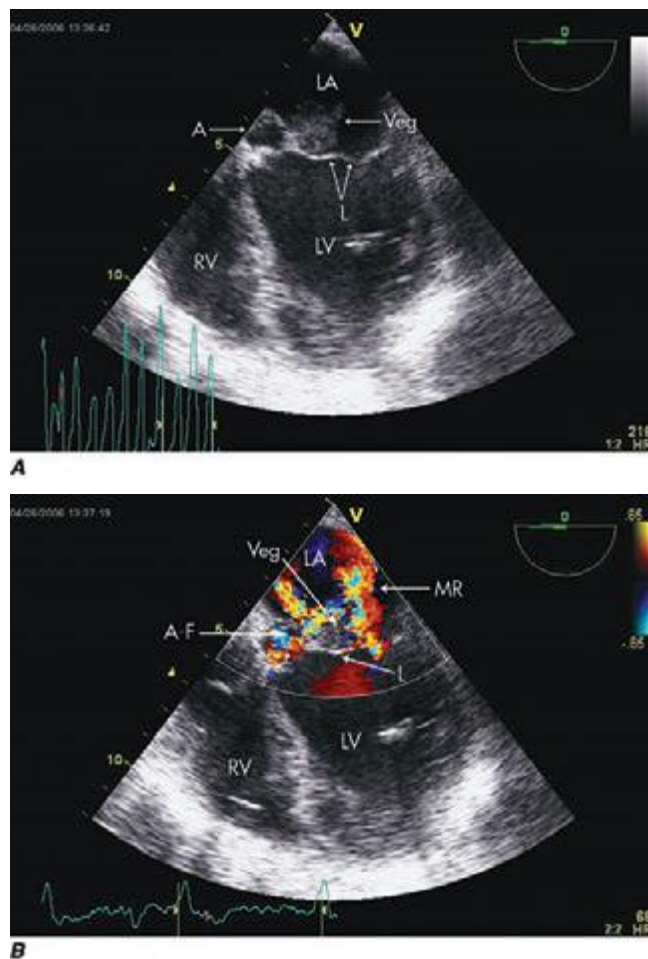


FIGURE 155-3 Imaging of a mitral valve infected with *Staphylococcus aureus* by low-esophageal, four-chamber-view, transesophageal echocardiography (TEE). **A**, Two-dimensional echocardiogram showing a large vegetation with an adjacent echolucent abscess cavity. **B**, Color-flow Doppler image showing severe mitral regurgitation through both the abscess-fistula and the central valve orifice. A, abscess; A-F, abscess-fistula; L, valve leaflets; LA, left atrium; LV, left ventricle; MR, mitral central valve regurgitation; RV, right ventricle; veg, vegetation. (With permission of Andrew Burger, MD.)

TREATMENT INFECTIVE ENDOCARDITIS

ANTIMICROBIAL THERAPY

To cure endocarditis, all bacteria in the vegetation must be killed. However, it is difficult to eradicate these bacteria because local host defenses are deficient and because the bacteria are largely nongrowing and metabolically inactive and thus are less easily killed by antibiotics. Accordingly, therapy must be bactericidal and prolonged. Antibiotics are generally given parenterally to achieve serum concentrations that, through passive diffusion, result in effective concentrations in the depths of the vegetation. To select effective therapy requires knowledge of the susceptibility of the causative microorganisms. The decision to initiate treatment empirically must balance the need to establish a microbiologic diagnosis against the potential progression of disease or the need for urgent surgery (see “Blood Cultures,” earlier). Simultaneous infection at other sites (such as the meninges), allergies, end-organ dysfunction, interactions with concomitantly administered medications, and risks of adverse events must be considered in the selection of therapy.

Although given for several weeks longer, the regimens recommended for the treatment of PVE (except that caused by staphylococci) are similar to those used to treat NVE (Table 155-4). Recommended doses and durations of therapy should be followed

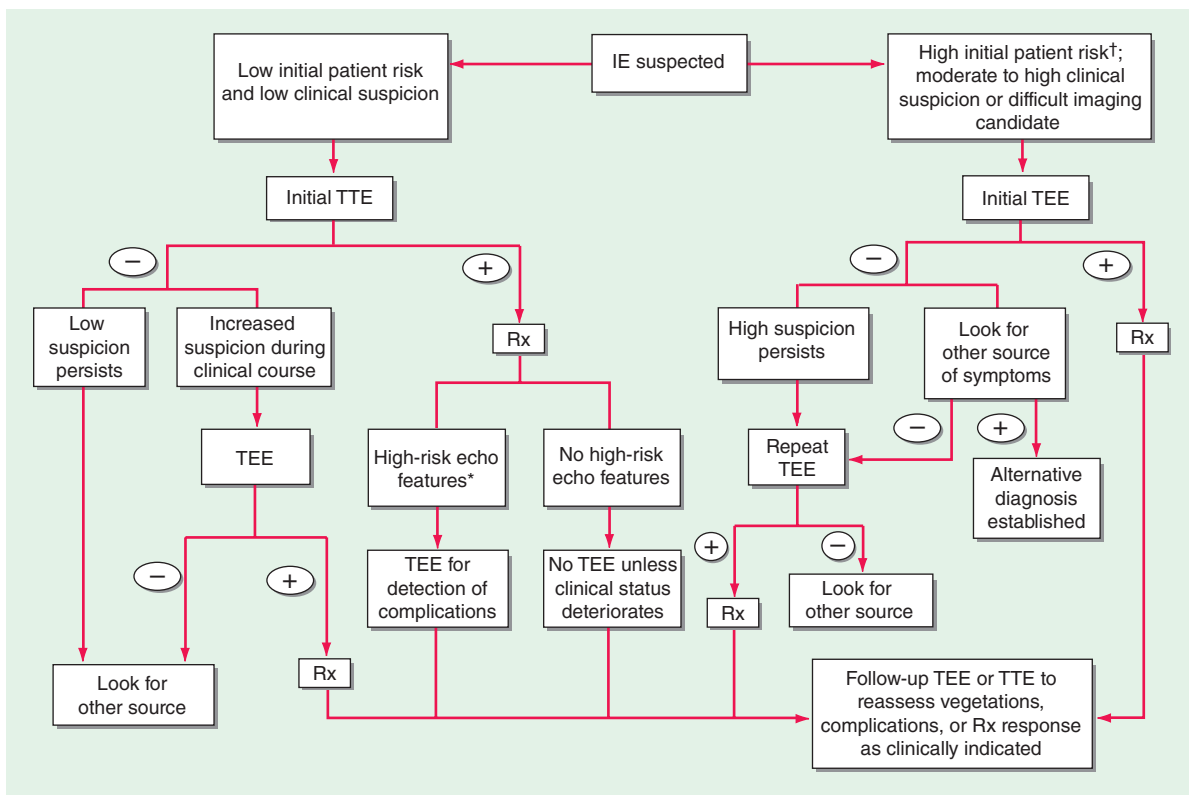


FIGURE 155-4 The diagnostic use of transesophageal and transtracheal echocardiography (TEE and TTE, respectively). †High initial patient risk for infective endocarditis (IE), as listed in Table 155-8, or evidence of intracardiac complications (new regurgitant murmur, new electrocardiographic conduction changes, or congestive heart failure). *High-risk echocardiographic features include large vegetations, valve insufficiency, paravalvular infection, or ventricular dysfunction. Rx indicates initiation of antibiotic therapy. (Reproduced with permission from *Diagnosis and Management of Infective Endocarditis and Its Complications*. *Circulation* 98:2936, 1998. © 1998 American Heart Association.)

unless alterations are required by end-organ dysfunction or adverse events.

Organism-Specific Therapies • STREPTOCOCCI Optimal therapy for streptococcal endocarditis is based on the minimal inhibitory concentration (MIC) of penicillin for the causative isolate (Table 155-4). The 2-week penicillin/gentamicin or ceftriaxone/gentamicin regimens should not be used to treat PVE or complicated NVE. Caution should be exercised in considering aminoglycoside-containing regimens for the treatment of patients at increased risk for aminoglycoside toxicity. The regimens recommended for relatively penicillin-resistant streptococci are advocated for treatment of group B, C, or G streptococcal endocarditis. Nutritionally variant organisms (*Granulicatella* or *Abiotrophia* species) and *Gemella* species are treated with the regimens for moderately penicillin-resistant streptococci, as is PVE caused by these organisms or by streptococci with a penicillin MIC of >0.1 $\mu\text{g}/\text{mL}$ (Table 155-4).

ENTEROCOCCI Enterococci are resistant to oxacillin, nafcillin, and the cephalosporins and are only inhibited—not killed—by penicillin, ampicillin, teicoplanin (not available in the United States), and vancomycin. To kill enterococci requires the synergistic interaction of a cell wall–active antibiotic that is effective at achievable serum concentrations (penicillin, ampicillin, vancomycin, or teicoplanin) and an aminoglycoside (gentamicin or streptomycin) to which the isolate does not exhibit high-level resistance. An isolate's resistance to cell wall–active agents or its ability to replicate in the presence of gentamicin at ≥ 500 $\mu\text{g}/\text{mL}$ or streptomycin at 1000–2000 $\mu\text{g}/\text{mL}$ —a phenomenon called *high-level aminoglycoside resistance*—indicates that the ineffective antimicrobial agent cannot participate in the interaction to produce killing. High-level resistance to gentamicin predicts that tobramycin, netilmicin, amikacin, and kanamycin also will be ineffective. In fact, even when enterococci are not highly resistant to gentamicin, it is difficult to predict the ability of these

other aminoglycosides to participate in synergistic killing; consequently, they should not, in general, be used to treat enterococcal endocarditis. High concentrations of ampicillin plus ceftriaxone or cefotaxime, by expanded binding of penicillin-binding proteins, also kill *E. faecalis* in vitro and in animal models of endocarditis.

Enterococci must be tested for high-level resistance to streptomycin and gentamicin, β -lactamase production, and susceptibility to penicillin and ampicillin (MIC, <8 $\mu\text{g}/\text{mL}$) and to vancomycin (MIC, ≤ 4 $\mu\text{g}/\text{mL}$) and teicoplanin (MIC ≤ 2 $\mu\text{g}/\text{mL}$). If the isolate produces β -lactamase, ampicillin/sulbactam or vancomycin can be used as the cell wall–active component; if the penicillin/ampicillin MIC is ≥ 8 $\mu\text{g}/\text{mL}$, vancomycin can be considered; and if the vancomycin MIC is ≥ 8 $\mu\text{g}/\text{mL}$, penicillin or ampicillin can be considered. In the absence of high-level resistance, gentamicin or streptomycin should be used as the aminoglycoside (Table 155-4). Although the dose of gentamicin used to achieve bactericidal synergy in treating enterococcal endocarditis is smaller than that used in standard therapy, nephrotoxicity (or vestibular toxicity with streptomycin) is not uncommon during treatment lasting 4–6 weeks. Regimens in which the aminoglycoside component is given for only 2–3 weeks have been curative and associated with less nephrotoxicity than those using longer courses of gentamicin. Thus regimens wherein gentamicin is administered for only 2–3 weeks are preferred by some.

If there is high-level resistance to both gentamicin and streptomycin, a synergistic bactericidal effect cannot be achieved by the addition of an aminoglycoside; thus no aminoglycoside should be given. Instead, an 8- to 12-week course of a single cell wall–active agent can be considered; for *E. faecalis* endocarditis, high doses of ampicillin combined with ceftriaxone or cefotaxime are suggested (Table 155-4). Nonrandomized comparative studies suggest that ampicillin-ceftriaxone may be as effective as (and less nephrotoxic than) penicillin or ampicillin plus an aminoglycoside in the treatment of *E. faecalis* endocarditis. Given the reduced risk

TABLE 155-4 ANTIBIOTIC TREATMENT FOR INFECTIVE ENDOCARDITIS CAUSED BY COMMON ORGANISMS^a

Organism	Drug (Dose, Duration)	Comments
Streptococci		
Penicillin-susceptible ^b streptococci, <i>S. gallolyticus</i>	<ul style="list-style-type: none"> Penicillin G (2–3 mU IV q4h for 4 weeks) Ceftriaxone (2 g/d IV as a single dose for 4 weeks) Vancomycin^c (15 mg/kg IV q12h for 4 weeks) 	<p>—</p> <p>Can use ceftriaxone in patients with nonimmediate penicillin allergy.</p> <p>Use vancomycin in patients with severe or immediate β-lactam allergy.</p>
Relatively penicillin-resistant ^f	<ul style="list-style-type: none"> Penicillin G (2–3 mU IV q4h) or ceftriaxone (2 g IV qd) for 2 weeks plus Gentamicin^d (3 mg/kg qd IV or IM, as a single dose^e or divided into equal doses q8h for 2 weeks) 	Avoid 2-week regimen when risk of aminoglycoside toxicity is increased and in prosthetic valve or complicated endocarditis.
Moderately penicillin-resistant ^g streptococci, nutritionally variant organisms, or <i>Gemella</i> species	<ul style="list-style-type: none"> Penicillin G (4 mU IV q4h) or ceftriaxone (2 g IV qd) for 4 weeks plus Gentamicin^d (3 mg/kg qd IV or IM, as a single dose^e or divided into equal doses q8h for 2 weeks) Vancomycin^c as noted above for 4 weeks 	Penicillin alone at this dose for 6 weeks or with gentamicin during the initial 2 weeks is preferred for prosthetic valve endocarditis caused by streptococci with penicillin MICs of ≤ 0.1 $\mu\text{g/mL}$.
	<ul style="list-style-type: none"> Penicillin G (4–5 mU IV q4h) or ceftriaxone (2 g IV qd) for 6 weeks plus Gentamicin^d (3 mg/kg qd IV or IM as a single dose^e or divided into equal doses q8h for 6 weeks) Vancomycin^c as noted above for 4 weeks 	Preferred for prosthetic valve endocarditis caused by streptococci with penicillin MICs of >0.1 $\mu\text{g/mL}$.
		Regimen is preferred by some.
Enterococci^h		
	<ul style="list-style-type: none"> Penicillin G (4–5 mU IV q4h) plus gentamicin^d (1 mg/kg IV q8h), both for 4–6 weeks Ampicillin (2 g IV q4h) plus gentamicin^d (1 mg/kg IV q8h), both for 4–6 weeks Vancomycin^c (15 mg/kg IV q12h) plus gentamicin^d (1 mg/kg IV q8h), both for 4–6 weeks 	Can use streptomycin (7.5 mg/kg q12h) in lieu of gentamicin if there is not high-level resistance to streptomycin.
	<ul style="list-style-type: none"> Ampicillin (2 g IV q4h) plus ceftriaxone (2 g IV q12h), both for 6 weeks 	—
		Use vancomycin plus gentamicin for penicillin-allergic patients (or desensitize to penicillin) and for isolates resistant to penicillin/ampicillin.
		Use for <i>E. faecalis</i> isolates with high-level resistance to gentamicin and streptomycin or for patients at high risk for aminoglycoside nephrotoxicity (see text).
Staphylococci		
MSSA infecting native valves (no foreign devices)	<ul style="list-style-type: none"> Nafcillin, oxacillin, or flucloxacillin (2 g IV q4h for 4–6 weeks) Cefazolin (2 g IV q8h for 4–6 weeks) Vancomycin^c (15 mg/kg IV q12h for 4–6 weeks) 	<p>Can use penicillin (4 mU q4h) if isolate is penicillin-susceptible (does not produce β-lactamase).</p> <p>Can use cefazolin regimen for patients with nonimmediate penicillin allergy.</p> <p>Use vancomycin for patients with immediate (urticarial) or severe penicillin allergy; see text regarding addition of gentamicin, fusidic acid, or rifampin.</p>
MRSA infecting native valves (no foreign devices)	<ul style="list-style-type: none"> Vancomycin^c (15 mg/kg IV q8–12h for 4–6 weeks) 	No role for routine use of rifampin (see text). Consider alternative treatment (see text) for MRSA with vancomycin MIC >1.0 or persistent bacteremia during vancomycin therapy.
MSSA infecting prosthetic valves	<ul style="list-style-type: none"> Nafcillin, oxacillin, or flucloxacillin (2 g IV q4h for 6–8 weeks) plus Gentamicin^d (1 mg/kg IM or IV q8h for 2 weeks) plus Rifampinⁱ (300 mg PO q8h for 6–8 weeks) 	Use gentamicin during initial 2 weeks; determine susceptibility to gentamicin before initiating rifampin (see text); if patient is highly allergic to penicillin, use regimen for MRSA; if β -lactam allergy is of the minor nonimmediate type, cefazolin can be substituted for oxacillin/nafcillin.
MRSA infecting prosthetic valves	<ul style="list-style-type: none"> Vancomycin^c (15 mg/kg IV q12h for 6–8 weeks) plus Gentamicin^d (1 mg/kg IM or IV q8h for 2 weeks) plus Rifampinⁱ (300 mg PO q8h for 6–8 weeks) 	Use gentamicin during initial 2 weeks; determine gentamicin susceptibility before initiating rifampin (see text).
HACEK Organisms		
	<ul style="list-style-type: none"> Ceftriaxone (2 g/d IV as a single dose for 4 weeks) Ampicillin/sulbactam (3 g IV q6h for 4 weeks) 	<p>Can use another third-generation cephalosporin at comparable dosage.</p> <p>—</p>

(continued)

TABLE 155-4 ANTIBIOTIC TREATMENT FOR INFECTIVE ENDOCARDITIS CAUSED BY COMMON ORGANISMS^a (CONTINUED)

Organism	Drug (Dose, Duration)	Comments
<i>Coxiella burnetii</i>	<ul style="list-style-type: none"> Doxycycline (100 mg PO q12h) <i>plus</i> hydroxychloroquine (200 mg PO q8h), both for 18 (native valve) or 24 (prosthetic valve) months 	Follow serology to monitor response during treatment (antiphase I IgG and IgA decreased 4-fold and IgM antiphase II negative) and thereafter for relapse.
<i>Bartonella</i> spp.	<ul style="list-style-type: none"> Ceftriaxone (2 g IV q24h) or ampicillin (2 g IV q4h) or doxycycline (100 mg q12h PO) for 6 weeks <i>plus</i> Gentamicin (1 mg/kg IV q8h for 3 weeks) 	If patient is highly allergic to β -lactams, use doxycycline.

^aDoses are for adults with normal renal function. Doses of gentamicin, streptomycin, and vancomycin must be adjusted for reduced renal function. Ideal body weight is used to calculate doses of gentamicin and streptomycin per kilogram (men = 50 kg + 2.3 kg per inch over 5 feet; women = 45.5 kg + 2.3 kg per inch over 5 feet). ^bMIC, ≤ 0.1 $\mu\text{g}/\text{mL}$. ^cVancomycin dose is based on actual body weight. Adjust for trough level of 10–15 $\mu\text{g}/\text{mL}$ for streptococcal and enterococcal infections and 15–20 $\mu\text{g}/\text{mL}$ for staphylococcal infections. ^dAminoglycosides should not be administered as single daily doses for enterococcal endocarditis and should be introduced as part of the initial treatment. Target peak and trough serum concentrations of divided-dose gentamicin 1 h after a 20- to 30-min infusion or IM injection are ~ 3.5 $\mu\text{g}/\text{mL}$ and ≤ 1 $\mu\text{g}/\text{mL}$, respectively; target peak and trough serum concentrations of streptomycin (timing as with gentamicin) are 20–35 $\mu\text{g}/\text{mL}$ and < 10 $\mu\text{g}/\text{mL}$, respectively. ^eNetilmicin (4 mg/kg qd, as a single dose) can be used in lieu of gentamicin. ^fMIC, > 0.1 $\mu\text{g}/\text{mL}$ and < 0.5 $\mu\text{g}/\text{mL}$. ^gMIC, ≥ 0.5 $\mu\text{g}/\text{mL}$ and < 8 $\mu\text{g}/\text{mL}$. ^hAntimicrobial susceptibility must be evaluated; see text. ⁱRifampin increases warfarin and dicumarol requirements for anticoagulation.

Abbreviations: MIC, minimal inhibitory concentration; MRSA, methicillin-resistant *S. aureus*; MSSA, methicillin-sensitive *S. aureus*.

of nephrotoxicity with ampicillin-ceftriaxone therapy, this regimen may also be preferred in patients who are at increased risk for aminoglycoside nephrotoxicity.

If the enterococcal isolate is resistant to all of the commonly used agents, suppression of bacteremia followed by surgical treatment should be considered. The role of newer agents potentially active against multidrug-resistant enterococci (quinupristin/dalfopristin [*E. faecium* only], linezolid, and daptomycin) in the treatment of endocarditis has not been established.

STAPHYLOCOCCI The regimens used to treat staphylococcal endocarditis (Table 155-4) are based not on coagulase production but rather on the presence or absence of a prosthetic valve or foreign device, the native valve(s) involved, and the susceptibility of the isolate to penicillin, methicillin, and vancomycin. All staphylococci are considered penicillin-resistant until shown not to produce penicillinase. Similarly, methicillin resistance has become so prevalent among staphylococci that empirical therapy should be initiated with a regimen that covers methicillin-resistant organisms and should later be revised if the isolate proves to be susceptible to methicillin. The addition of 3–5 days of gentamicin to a β -lactam antibiotic or vancomycin to enhance therapy for native mitral or aortic valve endocarditis has not improved survival rates and may be associated with nephrotoxicity. Neither this addition nor the addition of fusidic acid or rifampin is recommended.

For treatment of endocarditis caused by *methicillin-resistant S. aureus* (MRSA), vancomycin, dosed to achieve trough concentrations of 15–20 $\mu\text{g}/\text{mL}$, is recommended, with the caveat that this regimen may be associated with nephrotoxicity. Although resistance to vancomycin among staphylococci is rare, reduced vancomycin susceptibility among MRSA strains is increasingly encountered. Isolates with a vancomycin MIC of 4–16 $\mu\text{g}/\text{mL}$ have intermediate susceptibility and are referred to as *vancomycin-intermediate S. aureus* (VISA). Isolates with an MIC of 2 $\mu\text{g}/\text{mL}$ may harbor subpopulations with higher MICs. These *heteroresistant VISA* (hVISA) isolates are not detectable by routine susceptibility testing. Because of the pharmacokinetics/pharmacodynamics of vancomycin, killing of MRSA with a vancomycin MIC of > 1.0 $\mu\text{g}/\text{mL}$ is unpredictable, even with aggressive vancomycin dosing. Although not approved by the U.S. Food and Drug Administration for this indication, daptomycin (6 mg/kg [or, as some experts prefer, 8–10 mg/kg] IV once daily) has been recommended as an alternative to vancomycin, particularly for left-sided endocarditis caused by VISA, hVISA, or isolates with a vancomycin MIC of > 1.0 $\mu\text{g}/\text{mL}$. These isolates should be tested to document daptomycin susceptibility. Daptomycin activity against MRSA—even against some isolates with reduced daptomycin susceptibility—is enhanced by the addition of nafcillin or ceftaroline. Case reports suggest that either the latter combinations or ceftaroline

alone (600 mg IV q8h) may be effective in recalcitrant MRSA endocarditis. Nevertheless, a discussion of treatment of endocarditis in which MRSA bacteremia persists despite therapy is beyond the scope of this chapter and requires consultation with an infectious disease specialist. The efficacy of linezolid for left-sided MRSA endocarditis has not been established. Although not widely adopted by other groups, the recommendation of the British Society for Antimicrobial Chemotherapy is that a second drug be added to vancomycin (rifampin) or to daptomycin (rifampin, gentamicin, or linezolid) for the treatment of NVE due to MRSA.

Methicillin-susceptible *S. aureus* endocarditis that is uncomplicated and limited to the tricuspid or pulmonic valve can often be treated with a 2-week course that combines oxacillin or nafcillin (but not vancomycin) with gentamicin. However, patients with prolonged fever (≥ 5 days) during therapy or multiple septic pulmonary emboli should receive standard-duration therapy. Vancomycin plus gentamicin for 2 weeks as treatment for right-sided endocarditis caused by MRSA yields suboptimal results; thus this entity is treated for 4 weeks with vancomycin or daptomycin (6 mg/kg as a single daily dose).

Staphylococcal PVE is treated for 6–8 weeks with a multidrug regimen. Rifampin is an essential component because it kills staphylococci that are adherent to foreign material in a biofilm. Two other agents (selected on the basis of susceptibility testing) are combined with rifampin to prevent in vivo emergence of resistance. Because many staphylococci (particularly MRSA and *Staphylococcus epidermidis*) are resistant to gentamicin, the isolate's susceptibility to gentamicin or an alternative agent should be established before rifampin treatment is begun. If the isolate is resistant to gentamicin, then another aminoglycoside, a fluoroquinolone (chosen on the basis of susceptibility), or another active agent should be substituted for gentamicin.

OTHER ORGANISMS In the absence of meningitis, endocarditis caused by *Streptococcus pneumoniae* isolates with a penicillin MIC of ≤ 1 $\mu\text{g}/\text{mL}$ can be treated with IV penicillin (4 million units every 4 h), ceftriaxone (2 g/d as a single dose), or cefotaxime (at a comparable dosage). Infection caused by pneumococcal strains with a penicillin MIC of ≥ 2 $\mu\text{g}/\text{mL}$ should be treated with vancomycin. If meningitis is suspected or present, treatment with vancomycin plus ceftriaxone—at the doses advised for meningitis—should be initiated until susceptibility results are known. Definitive therapy should then be selected on the basis of meningitis breakpoints (penicillin MIC, 0.06 $\mu\text{g}/\text{mL}$; or ceftriaxone MIC, 0.5 $\mu\text{g}/\text{mL}$). *P. aeruginosa* endocarditis is treated with an antipseudomonal penicillin (ticarcillin or piperacillin) and high doses of tobramycin (8 mg/kg per day in three divided doses). Endocarditis caused by Enterobacteriaceae is treated with a potent β -lactam antibiotic plus an aminoglycoside. Corynebacterial

824 endocarditis is treated with a penicillin plus an aminoglycoside (if the organism is susceptible to the aminoglycoside) or with vancomycin, which is highly bactericidal for most strains. Therapy for *Candida* endocarditis consists of amphotericin B plus flucytosine and early surgery; long-term (if not indefinite) suppression with an oral azole is advised. Echinocandin treatment of *Candida* endocarditis has been effective in sporadic cases; nevertheless, the role of echinocandins in this setting has not been established.

Empirical Therapy In designing therapy (largely with antimicrobials and doses from Table 155-4 to target putative microorganisms) to be administered before culture results are known or when cultures are negative, clinical clues (e.g., acute vs. subacute presentation, site of infection, patient's predispositions) as well as epidemiologic clues to etiology must be considered. Thus empirical therapy for acute endocarditis in an injection drug user should cover MRSA and gram-negative bacilli. Treatment with vancomycin plus gentamicin, initiated immediately after blood samples are obtained for culture, covers these organisms as well as many other potential causes. Similarly, treatment of health care-associated endocarditis must cover MRSA. In the treatment of culture-negative episodes, marantic endocarditis must be excluded and fastidious organisms sought by serologic testing. In the absence of prior antibiotic therapy, it is unlikely that *S. aureus*, CoNS, or enterococcal infection will present with negative blood cultures; thus, in this situation, recommended empirical therapy targets not these organisms but rather nutritionally variant organisms, the HACEK group, and *Bartonella* species. Pending the availability of diagnostic data, blood culture-negative subacute NVE is treated with gentamicin plus ampicillin-sulbactam (12 g every 24 h) or ceftriaxone; doxycycline (100 mg twice daily) is added for enhanced *Bartonella* coverage. For culture-negative PVE, vancomycin, gentamicin, cefepime, and rifampin should be used if the prosthetic valve has been in place for ≤ 1 year. Empirical therapy for infected prosthetic valves in place for > 1 year is similar to that for culture-negative NVE. If cultures may be negative because of confounding by prior antibiotic administration, broader empirical therapy may be indicated, with particular attention to pathogens that are likely to be inhibited by the specific prior therapy.

CIED Endocarditis Antimicrobial therapy for CIED endocarditis is adjunctive to complete device removal. The antimicrobial selected is based on the causative organism and should be used as recommended for NVE (Table 155-4). Bacteremic CIED infection may be complicated by coincident NVE or remote-site infection (e.g., osteomyelitis). A 4- to 6-week course of endocarditis-targeted therapy is recommended for patients with CIED endocarditis and for those with bacteremia that continues during ongoing antimicrobial therapy after device removal. Although *S. aureus* bacteremia (and persistent CoNS bacteremia) in patients who have a CIED in place is likely—in the absence of another source—to reflect endocarditis and should be managed accordingly, not all bloodstream infections in these patients indicate endocarditis. If evidence suggesting endocarditis is lacking, bloodstream infection due to gram-negative bacilli, streptococci, enterococci, and *Candida* species may not indicate device infection. However, in the absence of another source, relapse after antimicrobial therapy increases the likelihood of CIED endocarditis and warrants treatment as such.

Outpatient Antimicrobial Therapy Fully compliant, clinically stable patients who are no longer bacteremic, are not febrile, and have no clinical or echocardiographic findings that suggest an impending complication may complete therapy as outpatients. Careful follow-up and a stable home setting are necessary, as are predictable IV access and use of antimicrobial agents that are stable in solution. Recommended regimens should not be compromised to accommodate outpatient therapy.

Monitoring Antimicrobial Therapy Measurement of the serum bactericidal titer—the highest dilution of the patient's serum during therapy that kills 99.9% of the standard inoculum of the infecting organism—is not recommended for assessment of standard

regimens but may be useful for assessment of the treatment of endocarditis caused by unusual organisms. Serum concentrations of aminoglycosides and vancomycin should be monitored and doses adjusted to avoid or address toxicity.

Antibiotic toxicities, including allergic reactions, occur in 25–40% of patients and commonly arise after several weeks of therapy. Blood tests to detect renal, hepatic, and hematologic toxicity should be performed periodically.

Blood cultures should be repeated daily until sterile in patients with endocarditis due to *S. aureus* or difficult-to-treat organisms, rechecked if there is recrudescence fever, and performed again 4–6 weeks after therapy to document cure. Blood cultures become sterile within 2 days after the start of appropriate therapy when infection is caused by viridans streptococci, enterococci, or HACEK organisms. In *S. aureus* endocarditis, β -lactam therapy results in sterile cultures in 3–5 days, whereas in MRSA endocarditis, positive cultures may persist for 7–9 days with vancomycin or daptomycin treatment. MRSA bacteremia persisting despite an adequate dosage of vancomycin may indicate infection due to a strain with reduced vancomycin susceptibility and therefore may point to a need for alternative therapy. When fever persists for 7 days despite appropriate antibiotic therapy, patients should be evaluated for paravalvular abscess, extracardiac abscesses (spleen, kidney), or complications (embolic events). Recrudescence fever raises the possibility of these complications but also of drug reactions or complications of hospitalization. Vegetations become smaller with effective therapy; however, 3 months after cure, 50% are unchanged and 25% are slightly larger.

SURGICAL TREATMENT

Intracardiac and central nervous system complications are important causes of morbidity and death due to infective endocarditis. In some cases, effective treatment for these complications requires surgery. The indications for cardiac surgical treatment of endocarditis (Table 155-5) have been derived from observational studies and expert opinion. The strength of individual indications varies; thus the risks and benefits as well as the timing of surgery must be individualized (Table 155-6). From 25% to 40% of patients with left-sided endocarditis undergo cardiac surgery during active infection, with slightly higher surgery rates for PVE than NVE. Intracardiac complications (which are most reliably detected by TEE) and CHF are the most commonly cited indications for surgery. The benefit of surgery has been assessed primarily in studies comparing populations of medically and surgically treated patients matched for the necessity of

TABLE 155-5 INDICATIONS FOR CARDIAC SURGICAL INTERVENTION IN PATIENTS WITH ENDOCARDITIS

Surgery Required for Optimal Outcome
Moderate to severe congestive heart failure due to valve dysfunction
Partially dehiscent unstable prosthetic valve
Persistent bacteremia despite optimal antimicrobial therapy
Lack of effective microbicidal therapy (e.g., fungal or <i>Brucella</i> endocarditis)
<i>S. aureus</i> prosthetic valve endocarditis with an intracardiac complication
Relapse of prosthetic valve endocarditis after optimal antimicrobial therapy
Surgery to Be Strongly Considered for Improved Outcome ^a
Perivalvular extension of infection
Poorly responsive <i>S. aureus</i> endocarditis involving the aortic or mitral valve
Large (> 10 mm in diameter) hypermobile vegetations with increased risk of embolism, particularly with prior embolic event or with significant valve dysfunction
Persistent unexplained fever (≥ 10 days) in culture-negative native valve endocarditis
Poorly responsive or relapsed endocarditis due to highly antibiotic-resistant enterococci or gram-negative bacilli

^aSurgery must be carefully considered; findings are often combined with other indications to prompt surgery.

TABLE 155-6 TIMING OF CARDIAC SURGICAL INTERVENTION IN PATIENTS WITH ENDOCARDITIS

Timing	Indication for Surgical Intervention	
	Strong Supporting Evidence	Conflicting Evidence, but Majority of Opinions Favor Surgery
Emergent (same day)	Acute aortic regurgitation plus preclosure of mitral valve Sinus of Valsalva abscess ruptured into right heart Rupture into pericardial sac	
Urgent (within 1–2 days)	Valve obstruction by vegetation Unstable (dehiscenced) prosthesis Acute aortic or mitral regurgitation with heart failure (New York Heart Association class III or IV) Septal perforation Perivalvular extension of infection with or without new electrocardiographic conduction system changes Lack of effective antibiotic therapy	Major embolus plus persisting large vegetation (>10 mm in diameter)
Elective (earlier usually preferred)	Vegetation diameter >10 mm plus severe aortic or mitral valve dysfunction ^a Progressive paravalvular prosthetic regurgitation Valve dysfunction plus persisting infection after ≥7–10 days of antimicrobial therapy Fungal (mold) endocarditis	Staphylococcal prosthetic valve endocarditis Early prosthetic valve endocarditis (≤2 months after valve surgery) Fungal endocarditis (<i>Candida</i> spp.) Antibiotic-resistant organisms

^aSupported by a single-institution randomized trial showing benefit from early surgery. Implementation requires clinical judgment.

Source: Adapted from L Olaison, G Petterson: *Infect Dis Clin North Am* 16:453, 2002.

surgery (indications assessed in studies as *propensity*), with adjustments for predictors of death (comorbidities) and timing of the surgical intervention. Although study results vary, surgery for currently advised indications appears to convey a significant survival benefit (27–55%) that becomes apparent only with follow-up for ≥6 months. During the initial weeks after surgery, mortality risk may appear increased (disease + surgery-related mortality).

Indications • CONGESTIVE HEART FAILURE Moderate to severe refractory CHF caused by new or worsening valve dysfunction is the major indication for cardiac surgery. At 6 months of follow-up, patients with left-sided endocarditis and moderate to severe heart failure due to valve dysfunction who are treated medically have a 50% mortality rate, while among matched patients who undergo surgery the mortality rate is 15%. The survival benefit with surgery, which is most predictable among patients with the most weighty indications (*propensity*), is seen in both NVE and PVE. Surgery can relieve functional stenosis due to large vegetations or restore competence to damaged regurgitant valves by repair or replacement.

PERIVALVULAR INFECTION This complication, which is most common with aortic valve infection, occurs in 10–15% of native valve and 45–60% of prosthetic valve infections. It is suggested by persistent unexplained fever during appropriate therapy, new electrocardiographic conduction disturbances, or pericarditis. TEE with color Doppler is the test of choice to detect perivalvular abscesses (sensitivity, ≥85%). For optimal outcome, surgery is required, especially when fever persists, fistulae develop, prostheses are dehiscenced and unstable, or infection relapses after appropriate treatment. Cardiac rhythm must be monitored since high-grade heart block may require insertion of a pacemaker.

UNCONTROLLED INFECTION Continued positive blood cultures or otherwise-unexplained persistent fevers (in patients with either blood culture-positive or -negative endocarditis) despite optimal antibiotic therapy may reflect uncontrolled infection and may warrant surgery. Surgical treatment is also advised for endocarditis caused by organisms against which effective antimicrobial therapy is lacking (e.g., yeasts, fungi, *P. aeruginosa*, other highly resistant gram-negative bacilli, *Brucella* species).

S. AUREUS ENDOCARDITIS The mortality rate for *S. aureus* PVE exceeds 50% with medical treatment but is reduced to 25% with surgical treatment. In patients with intracardiac complications associated

with *S. aureus* PVE, surgical treatment reduces the mortality rate twentyfold. Surgical treatment should be considered for patients with *S. aureus* native aortic or mitral valve infection who have TTE-demonstrable vegetations and remain septic during the initial week of therapy. Isolated tricuspid valve endocarditis, even with persistent fever, rarely requires surgery.

PREVENTION OF SYSTEMIC EMBOLI Death and persisting morbidity may result from cerebral or coronary artery emboli. Predicting a high risk of systemic embolization by echocardiographic determination of vegetation size and anatomy does not by itself identify those patients in whom surgery to prevent emboli will result in increased chances of survival. Net benefits from surgery to prevent emboli are most likely when other surgical benefits can be achieved simultaneously—e.g., repair of a moderately dysfunctional valve or debridement of a paravalvular abscess. Only 3.5% of patients undergo surgery solely to prevent systemic emboli. Valve repair, with the consequent avoidance of prosthesis insertion, improves the benefit-to-risk ratio of surgery performed to address vegetations.

CIED ENDOCARDITIS Removal of all hardware is recommended for patients with established CIED infection (pocket or intracardiac lead) or erosion of the device through the skin. Percutaneous lead extraction is preferred. With lead vegetations of >3 cm and the resulting risk of a pulmonary embolus or with retained hardware after attempted percutaneous extraction, surgical removal should be considered. Removal of the infected CIED during the initial hospitalization is associated with increased 30-day and 1-year survival rates over those attained with antibiotic therapy and device retention. If necessary, the CIED can be reimplemented percutaneously or surgically (epicardial leads) at a new site after at least 10–14 days of effective antimicrobial therapy. CIEDs should be removed and replaced subsequently when patients undergo valve surgery for endocarditis.

Timing of Cardiac Surgery With the more life-threatening indications for surgery (valve dysfunction and severe CHF, paravalvular abscess, major prosthesis dehiscence), early surgery—i.e., during the initial week of therapy—is associated with a greater chance of survival than later surgery. With less compelling indications, surgery may reasonably be delayed to allow further treatment as well as improvement in overall health (Table 155-6). After 14 days of recommended antibiotic therapy, excised valves are culture-negative in 99% and 50% of patients with streptococcal and *S. aureus* endocarditis,

826 respectively. Recrudescence of endocarditis on a new implanted prosthetic valve follows surgery for active NVE and PVE in 2% and 6–15% of patients, respectively. These frequencies do not justify the risk of an adverse outcome due to a delay in surgery, particularly in patients with severe heart failure, valve dysfunction, and uncontrolled staphylococcal infections. Delay is justified when infection is controlled and CHF is resolved with medical therapy.

Neurologic complications of endocarditis may be exacerbated as a consequence of cardiac surgery. The risk of neurologic deterioration is related to the type of neurologic complication and the interval between the complication and surgery. Whenever feasible, cardiac surgery should be delayed for 2–3 weeks after a nonhemorrhagic embolic infarction and for 4 weeks after a cerebral hemorrhage. A ruptured mycotic aneurysm should be treated before cardiac surgery.

Antibiotic Therapy after Cardiac Surgery Organisms have been detected on Gram's stain—or their DNA has been detected by polymerase chain reaction—in excised valves from 45% of patients who have successfully completed the recommended therapy for endocarditis. In only 7% of these patients are the organisms, most of which are unusual and antibiotic resistant, cultured from the valve. Detection of organisms or their DNA does not necessarily indicate antibiotic failure; in fact, relapse of endocarditis after surgery is uncommon. Thus, when valve cultures are negative in uncomplicated NVE caused by susceptible organisms, the duration of preoperative plus postoperative treatment should equal the total duration of recommended therapy, with ~2 weeks of treatment administered after surgery. For endocarditis complicated by paravalvular abscess, partially treated PVE, or cases with culture-positive valves, a full course of therapy should be given postoperatively.

Extracardiac Complications Splenic abscess develops in 3–5% of patients with endocarditis. Effective therapy requires either image-guided percutaneous drainage or splenectomy. Mycotic aneurysms occur in 2–15% of endocarditis patients; one-half of these cases involve the cerebral arteries and present as headaches, focal neurologic symptoms, or hemorrhage. Cerebral aneurysms should be monitored by angiography. Some will resolve with effective antimicrobial therapy, but those that persist, enlarge, or leak should be treated surgically if possible. Extracerebral aneurysms present as local pain, a mass, local ischemia, or bleeding; these aneurysms are treated surgically.

OUTCOME

Factors that can adversely affect outcome include older age, severe comorbid conditions and diabetes, delayed diagnosis, involvement of prosthetic valves or the aortic valve, an invasive (*S. aureus*) or antibiotic-resistant (*P. aeruginosa*, yeast) pathogen, intracardiac and major neurologic complications, and an association with health care. Death and poor outcome often are related not to failure of antibiotic therapy but rather to the interactions of comorbidities and endocarditis-related end-organ complications. In developed countries, overall survival rates are 80–85%; however, rates vary considerably among subpopulations of endocarditis patients. Survival rates for patients with NVE caused by viridans streptococci, HACEK organisms, or enterococci (susceptible to synergistic therapy) are 85–90%. For *S. aureus* NVE in patients who do not inject drugs, survival rates are 55–70%, whereas 85–90% of injection drug users survive this infection. PVE beginning within 2 months of valve replacement results in mortality rates of 40–50%, whereas rates are only 10–20% in later-onset cases.

PREVENTION

To prevent endocarditis (long a goal in clinical practice), past expert committees have supported systemic antibiotic administration prior to many bacteremia-inducing procedures. A reappraisal of the evidence for antibiotic prophylaxis for endocarditis by the American Heart Association and the European Society of Cardiology culminated in guidelines advising its more restrictive use. At best, the

TABLE 155-7 ANTIBIOTIC REGIMENS FOR PROPHYLAXIS OF ENDOCARDITIS IN ADULTS WITH HIGH-RISK CARDIAC LESIONS^{a,b}

A. Standard oral regimen	Amoxicillin: 2 g PO 1 h before procedure
B. Inability to take oral medication	Ampicillin: 2 g IV or IM within 1 h before procedure
C. Penicillin allergy	1. Clarithromycin or azithromycin: 500 mg PO 1 h before procedure
	2. Cephalexin ^c : 2 g PO 1 h before procedure
	3. Clindamycin: 600 mg PO 1 h before procedure
D. Penicillin allergy, inability to take oral medication	1. Cefazolin ^c or ceftriaxone ^c : 1 g IV or IM 30 min before procedure
	2. Clindamycin: 600 mg IV or IM 1 h before procedure

^aDosing for children: for amoxicillin, ampicillin, cephalexin, or cefadroxil, use 50 mg/kg PO; cefazolin, 25 mg/kg IV; clindamycin, 20 mg/kg PO or 25 mg/kg IV; clarithromycin, 15 mg/kg PO; and vancomycin, 20 mg/kg IV. ^bFor high-risk lesions, see Table 155-8. Prophylaxis is not advised for other lesions. ^cDo not use cephalosporins in patients with immediate hypersensitivity (urticaria, angioedema, anaphylaxis) to penicillin.

Source: Table created using the guidelines published by the American Heart Association and the European Society of Cardiology (W Wilson et al: *Circulation* 116:1736, 2007; and G Habib et al: *Eur Heart J* 30:2369, 2009).

benefit of antibiotic prophylaxis is minimal. Most endocarditis cases do not follow a procedure. Although dental treatments have been widely considered to predispose to endocarditis, such infection occurs no more frequently in patients who are undergoing dental treatment than in matched controls who are not. Furthermore, the frequency and magnitude of bacteremia associated with dental procedures and a routine day's activities (e.g., tooth brushing and flossing) are similar; because dental procedures are infrequent events, exposure of cardiac structures to bacteremic oral-cavity organisms is notably greater from routine daily activities than from dental care. The relation of gastrointestinal and genitourinary procedures to subsequent endocarditis is even more tenuous than that of dental procedures. In addition, cost-effectiveness and cost-benefit estimates suggest that antibiotic prophylaxis represents a poor use of resources.

Nevertheless, studies in animal models suggest that antibiotic prophylaxis may be effective. Thus it is possible that rare cases of endocarditis are prevented. Weighing the potential benefits, potential adverse events, and costs associated with antibiotic prophylaxis, the American Heart Association and the European Society of Cardiology now recommend prophylactic antibiotics (Table 155-7) only for those patients at highest risk for severe morbidity or death from endocarditis (Table 155-8). Maintaining good dental hygiene is essential. Prophylaxis is recommended only when there is manipulation of gingival tissue or the periapical region of the teeth or perforation of the oral mucosa (including surgery on the respiratory tract). Prophylaxis is not advised for patients undergoing gastrointestinal or genitourinary tract procedures. High-risk patients should be treated before or

TABLE 155-8 HIGH-RISK CARDIAC LESIONS FOR WHICH ENDOCARDITIS PROPHYLAXIS IS ADVISED BEFORE DENTAL PROCEDURES

Prosthetic heart valves
Prior endocarditis
Unrepaired cyanotic congenital heart disease, including palliative shunts or conduits
Completely repaired congenital heart defects during the 6 months after repair
Incompletely repaired congenital heart disease with residual defects adjacent to prosthetic material
Valvulopathy developing after cardiac transplantation ^a

^aNot a target population for prophylaxis according to recommendations of the European Society of Cardiology.

Source: Table created using the guidelines published by the American Heart Association and the European Society of Cardiology (W Wilson et al: *Circulation* 116:1736, 2007; and G Habib et al: *Eur Heart J* 30:2369, 2009).

when they undergo procedures on an infected genitourinary tract or on infected skin and soft tissue.

In patients with aortic or mitral valve regurgitation or a prosthetic valve, treatment of acute Q fever with doxycycline plus hydroxychloroquine (for doses, see Table 155-4) for 12 months is highly effective in preventing *C. burnetii* endocarditis.

The National Institute for Health and Clinical Excellence in the United Kingdom has advised discontinuation of all antibiotic prophylaxis for endocarditis. Limited surveillance studies have not detected increased viridans streptococcal endocarditis subsequent to the promulgation of guidelines that are more restrictive or advise no prophylaxis.

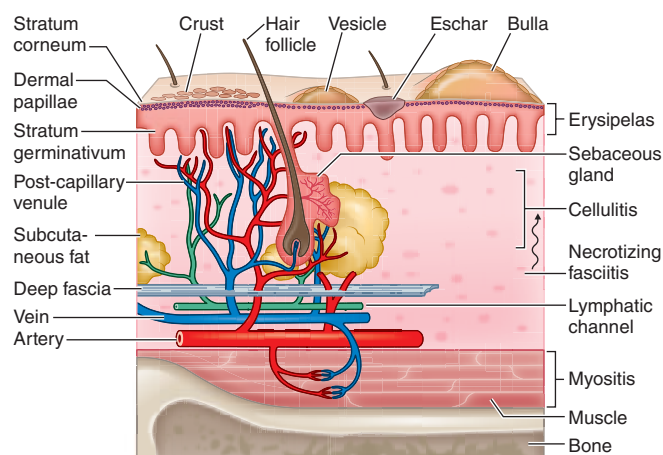


FIGURE 156-1 Structural components of the skin and soft tissue, superficial infections, and infections of the deeper structures. The rich capillary network beneath the dermal papillae plays a key role in the localization of infection and in the development of the acute inflammatory reaction.

156 Infections of the Skin, Muscles, and Soft Tissues

Dennis L. Stevens

Skin and soft tissue infections occur in all races, all ethnic groups, and all geographic locations, although some have unique geographic niches. In modern times, the frequency and severity of some skin and soft tissue infections have increased for several reasons. First, microbes are rapidly disseminated throughout the world via efficient air travel, acquiring genes for virulence factors and antibiotic resistance. Second, natural disasters, such as earthquakes, tsunamis, tornadoes, and hurricanes, appear to be increasing in frequency, and the injuries sustained during these events commonly cause major skin and soft-tissue damage that predisposes to infection. Third, trauma and casualties resulting from combat and terrorist activities can markedly damage or destroy tissues and provide both endogenous and exogenous pathogens with ready access to deeper structures. Unfortunately, because the marvels of modern medicine may not be available during human-instigated and natural disasters, primary treatment may be delayed and the likelihood of severe infection and death increased.

ANATOMIC RELATIONSHIPS: CLUES TO THE DIAGNOSIS OF SOFT TISSUE INFECTIONS

Skin and soft tissue infections have been common human afflictions for centuries. However, between 2000 and 2004, hospital admissions for skin and soft tissue infections rose by 27%, a remarkable increase that was attributable largely to the emergence of the USA300 clone of methicillin-resistant *Staphylococcus aureus* (MRSA). This chapter provides an anatomic approach to understanding the types of soft tissue infections and the diverse microbes responsible.

Protection against infection of the epidermis depends on the mechanical barrier afforded by the stratum corneum, since the epidermis itself is devoid of blood vessels (Fig. 156-1). Disruption of this layer by burns or bites, abrasions, foreign bodies, primary dermatologic disorders (e.g., herpes simplex, varicella, ecthyma gangrenosum), surgery, or vascular or pressure ulcer allows penetration of bacteria to the deeper structures. Similarly, the hair follicle can serve as a portal either for components of the normal flora (e.g., *Staphylococcus*) or for extrinsic bacteria (e.g., *Pseudomonas* in hot-tub folliculitis). Intracellular infection of the squamous epithelium with vesicle formation may arise from cutaneous inoculation, as in infection with herpes simplex virus (HSV) type 1; from the dermal capillary plexus, as in varicella and infections due to other viruses associated with viremia; or from cutaneous nerve roots, as in herpes zoster. Bacteria infecting the epidermis, such as *Streptococcus pyogenes*, may be translocated laterally to deeper structures via lymphatics, an event that results in the rapid superficial spread of erysipelas. Later, engorgement or obstruction of

lymphatics causes flaccid edema of the epidermis, another characteristic of erysipelas.

The rich plexus of capillaries beneath the dermal papillae provides nutrition to the stratum germinativum, and physiologic responses of this plexus produce important clinical signs and symptoms. For example, infective vasculitis of the plexus results in petechiae, Osler's nodes, Janeway lesions, and palpable purpura, which, if present, are important clues to the existence of endocarditis (Chap. 155). In addition, metastatic infection within this plexus can result in cutaneous manifestations of disseminated fungal infection (Chap. 240), gonococcal infection (Chap. 181), *Salmonella* infection (Chap. 190), *Pseudomonas* infection (i.e., ecthyma gangrenosum; Chap. 189), meningococemia (Chap. 180), and staphylococcal infection (Chap. 172). The plexus also provides bacteria with access to the circulation, thereby facilitating local spread or bacteremia. The postcapillary venules of this plexus are a prominent site of polymorphonuclear leukocyte sequestration, diapedesis, and chemotaxis to the site of cutaneous infection.

Amplification of these physiologic mechanisms by excessive levels of cytokines or bacterial toxins causes leukostasis, venous occlusion, and pitting edema. Edema with purple bullae, ecchymosis, and cutaneous anesthesia suggests loss of vascular integrity and necessitates exploration of the deeper structures for evidence of necrotizing fasciitis or myonecrosis. An early diagnosis requires a high level of suspicion in instances of unexplained fever and of pain and tenderness in the soft tissue, even in the absence of acute cutaneous inflammation.

Table 156-1 indicates the chapters in which the infections described below are discussed in greater detail. Many of these infections are illustrated in the chapters cited or in Chap. 25e (Atlas of Rashes Associated with Fever).

INFECTIONS ASSOCIATED WITH VESICLES

(Table 156-1) Vesicle formation due to infection is caused by viral proliferation within the epidermis. In varicella and variola, viremia precedes the onset of a diffuse centripetal rash that progresses from macules to vesicles, then to pustules, and finally to scabs over the course of 1–2 weeks. Vesicles of varicella have a “dewdrop” appearance and develop in crops randomly about the trunk, extremities, and face over 3–4 days. Herpes zoster occurs in a single dermatome; the appearance of vesicles is preceded by pain for several days. Zoster may occur in persons of any age but is most common among immunosuppressed individuals and elderly patients, whereas most cases of varicella occur in young children. Vesicles due to HSV are found on or around the

Lesion, Clinical Syndrome	Infectious Agent	See Also Chap(s).
Vesicles		
Smallpox	Variola virus	261e
Chickenpox	Varicella-zoster virus	217
Shingles (herpes zoster)	Varicella-zoster virus	217
Cold sores, herpetic whitlow, herpes gladiatorum	Herpes simplex virus	216
Hand-foot-and-mouth disease	Coxsackievirus A16	228
Orf	Parapoxvirus	220e
Molluscum contagiosum	Molluscum contagiosum poxvirus	220e
Rickettsialpox	<i>Rickettsia akari</i>	211
Blistering distal dactylitis	<i>Staphylococcus aureus</i> or <i>Streptococcus pyogenes</i>	172, 173
Bullae		
Staphylococcal scalded-skin syndrome	<i>S. aureus</i>	172
Necrotizing fasciitis	<i>S. pyogenes</i> , <i>Clostridium</i> spp., mixed aerobes and anaerobes	173, 179, 201
Gas gangrene	<i>Clostridium</i> spp.	179
Halophilic vibrio	<i>Vibrio vulnificus</i>	193
Crusted lesions		
Bullous impetigo/ecthyma	<i>S. aureus</i>	172
Impetigo contagiosa	<i>S. pyogenes</i>	173
Ringworm	Superficial dermatophyte fungi	243
Sporotrichosis	<i>Sporothrix schenckii</i>	243
Histoplasmosis	<i>Histoplasma capsulatum</i>	236
Coccidioidomycosis	<i>Coccidioides immitis</i>	237
Blastomycosis	<i>Blastomyces dermatitidis</i>	238
Cutaneous leishmaniasis	<i>Leishmania</i> spp.	251
Cutaneous tuberculosis	<i>Mycobacterium tuberculosis</i>	202
Nocardiosis	<i>Nocardia asteroides</i>	199
Folliculitis		
Furunculosis	<i>S. aureus</i>	172
Hot-tub folliculitis	<i>Pseudomonas aeruginosa</i>	189
Swimmer's itch	<i>Schistosoma</i> spp.	259
Acne vulgaris	<i>Propionibacterium acnes</i>	71
Papular and nodular lesions		
Fish-tank or swimming-pool granuloma	<i>Mycobacterium marinum</i>	204
Creeping eruption (cutaneous larva migrans)	<i>Ancylostoma braziliense</i>	256
Dracunculiasis	<i>Dracunculus medinensis</i>	258
Cercarial dermatitis	<i>Schistosoma mansoni</i>	259
Verruca vulgaris	Human papillomaviruses 1, 2, 4	222
Condylomata acuminata (anogenital warts)	Human papillomaviruses 6, 11, 16, 18	222
Onchocerciasis nodule	<i>Onchocerca volvulus</i>	258
Cutaneous myiasis	<i>Dermatobia hominis</i>	475
Verruca peruana	<i>Bartonella bacilliformis</i>	197
Cat-scratch disease	<i>Bartonella henselae</i>	197
Lepromatous leprosy	<i>Mycobacterium leprae</i>	203
Secondary syphilis (papulosquamous and nodular lesions, condylomata lata)	<i>Treponema pallidum</i>	206
Tertiary syphilis (nodular gummatous lesions)	<i>T. pallidum</i>	206
Ulcers with or without eschars		
Anthrax	<i>Bacillus anthracis</i>	261e
Ulceroglandular tularemia	<i>Francisella tularensis</i>	195, 261e
Bubonic plague	<i>Yersinia pestis</i>	196, 261e
Buruli ulcer	<i>Mycobacterium ulcerans</i>	204
Leprosy	<i>M. leprae</i>	203
Cutaneous tuberculosis	<i>M. tuberculosis</i>	202
Chancroid	<i>Haemophilus ducreyi</i>	182
Primary syphilis	<i>T. pallidum</i>	206
Erysipelas	<i>S. pyogenes</i>	173
Cellulitis	<i>Staphylococcus</i> spp., <i>Streptococcus</i> spp., various other bacteria	Various
Necrotizing fasciitis		
Streptococcal gangrene	<i>S. pyogenes</i>	173
Fournier's gangrene	Mixed aerobic and anaerobic bacteria	201
Staphylococcal necrotizing fasciitis	Methicillin-resistant <i>S. aureus</i>	172
Myositis and myonecrosis		
Pyomyositis	<i>S. aureus</i>	172
Streptococcal necrotizing myositis	<i>S. pyogenes</i>	173
Gas gangrene	<i>Clostridium</i> spp.	179
Nonclostridial (crepitant) myositis	Mixed aerobic and anaerobic bacteria	201
Synergistic nonclostridial anaerobic myonecrosis	Mixed aerobic and anaerobic bacteria	201

lips (HSV-1) or genitals (HSV-2) but also may appear on the head and neck of young wrestlers (herpes gladiatorum) or on the digits of health care workers (herpetic whitlow). Recurrent herpes labialis (HSV-1) and herpes genitalis commonly follow primary infection. Coxsackievirus A16 characteristically causes vesicles on the hands, feet, and mouth of children. Orf is caused by a DNA virus related to smallpox virus and infects the fingers of individuals who work around goats and sheep. Molluscum contagiosum virus induces flaccid vesicles on the skin of healthy and immunocompromised individuals. Although variola (smallpox) in nature was eradicated as of 1977, post-millennial terrorist events have renewed interest in this devastating infection (Chap. 261e). Viremia beginning after an incubation period of 12 days is followed by a diffuse maculopapular rash, with rapid evolution to vesicles, pustules, and then scabs. Secondary cases can occur among close contacts.

Rickettsialpox begins after mite-bite inoculation of *Rickettsia akari* into the skin. A papule with a central vesicle evolves to form a 1- to 2.5-cm painless crusted black eschar with an erythematous halo and proximal adenopathy. While more common in the northeastern United States and the Ukraine in 1940–1950, rickettsialpox has recently been described in Ohio, Arizona, and Utah. Blistering dactylitis is a painful, vesicular, localized *S. aureus* or group A streptococcal infection of the pulps of the distal digits of the hands.

INFECTIONS ASSOCIATED WITH BULLAE

(Table 156-1) Staphylococcal scalded-skin syndrome (SSSS) in neonates is caused by a toxin (exfoliatin) from phage group II *S. aureus*. SSSS must be distinguished from toxic epidermal necrolysis (TEN), which occurs primarily in adults, is drug-induced, and is associated with a higher mortality rate. Punch biopsy with frozen section is useful in making this distinction since the cleavage plane is the stratum corneum in SSSS and the stratum germinativum in TEN (Fig. 156-1). Intravenous γ -globulin is a promising treatment for TEN. Necrotizing fasciitis and gas gangrene also induce bulla formation (see “Necrotizing Fasciitis,” below). Halophilic vibrio infection can be as aggressive and fulminant as necrotizing fasciitis; a helpful clue in its diagnosis is a history of exposure to waters of the Gulf of Mexico or the Atlantic seaboard or (in a patient with cirrhosis) the ingestion of raw seafood. The etiologic organism (*Vibrio vulnificus*) is highly susceptible to tetracycline.

INFECTIONS ASSOCIATED WITH CRUSTED LESIONS

(Table 156-1) Impetigo contagiosa is caused by *S. pyogenes*, and bullous impetigo is due to *S. aureus*. Both skin lesions may have an early bullous stage but then appear as thick crusts with a golden-brown color. Epidemics of impetigo caused by MRSA have been reported. Streptococcal lesions are most common among children 2–5 years of age, and epidemics may occur in settings of poor hygiene, particularly among children in lower socioeconomic settings in tropical climates. It is important to recognize impetigo contagiosa because of its relationship to poststreptococcal glomerulonephritis. Rheumatic fever is not a complication of skin infection caused by *S. pyogenes*. Superficial dermatophyte infection (ringworm) can occur on any skin surface, and skin scrapings with KOH staining are diagnostic. Primary infections with dimorphic fungi such as *Blastomyces dermatitidis* and *Sporothrix schenckii* can initially present as crusted skin lesions resembling ringworm. Disseminated infection with *Coccidioides immitis* can also involve the skin, and biopsy and culture should be performed on crusted lesions in patients from endemic areas. Crusted nodular lesions caused by *Mycobacterium chelonae* have been described in HIV-seropositive patients. Treatment with clarithromycin looks promising.

FOLLICULITIS

(Table 156-1) Hair follicles serve as portals for a number of bacteria, although *S. aureus* is the most common cause of localized folliculitis. Sebaceous glands empty into hair follicles and ducts and, if these portals are blocked, form sebaceous cysts that may resemble staphylococcal abscesses or may become secondarily infected. Infection of sweat

glands (hidradenitis suppurativa) also can mimic infection of hair follicles, particularly in the axillae. Chronic folliculitis is uncommon except in acne vulgaris, where constituents of the normal flora (e.g., *Propionibacterium acnes*) may play a role.

Diffuse folliculitis occurs in two settings. *Hot-tub folliculitis* is caused by *Pseudomonas aeruginosa* in waters that are insufficiently chlorinated and maintained at temperatures of 37–40°C. Infection is usually self-limited, although bacteremia and shock have been reported. *Swimmer's itch* occurs when a skin surface is exposed to water infested with freshwater avian schistosomes. Warm water temperatures and alkaline pH are suitable for mollusks that serve as intermediate hosts between birds and humans. Free-swimming schistosomal cercariae readily penetrate human hair follicles or pores but quickly die and elicit a brisk allergic reaction, causing intense itching and erythema.

PAPULAR AND NODULAR LESIONS

(Table 156-1) Raised lesions of the skin occur in many different forms. *Mycobacterium marinum* infections of the skin may present as cellulitis or as raised erythematous nodules. Similar lesions caused by *Mycobacterium abscessus* and *M. chelonae* have been described among patients undergoing cosmetic laser surgery and tattooing, respectively. Erythematous papules are early manifestations of cat-scratch disease (with lesions developing at the primary site of inoculation of *Bartonella henselae*) and bacillary angiomatosis (also caused by *B. henselae*). Raised serpiginous or linear eruptions are characteristic of cutaneous larva migrans, which is caused by burrowing larvae of dog or cat hookworms (*Ancylostoma braziliense*) and which humans acquire through contact with soil that has been contaminated with dog or cat feces. Similar burrowing raised lesions are present in dracunculiasis caused by migration of the adult female nematode *Dracunculus medinensis*. Nodules caused by *Onchocerca volvulus* measure 1–10 cm in diameter and occur mostly in persons bitten by *Simulium* flies in Africa. The nodules contain the adult worm encased in fibrous tissue. Migration of microfilariae into the eyes may result in blindness. Verruga peruana is caused by *Bartonella bacilliformis*, which is transmitted to humans by the sandfly *Phlebotomus*. This condition can take the form of single gigantic lesions (several centimeters in diameter) or multiple small lesions (several millimeters in diameter). Numerous subcutaneous nodules may also be present in cysticercosis caused by larvae of *Taenia solium*. Multiple erythematous papules develop in schistosomiasis; each represents a cercarial invasion site. Skin nodules as well as thickened subcutaneous tissue are prominent features of lepromatous leprosy. Large nodules or gummas are features of tertiary syphilis, whereas flat papulosquamous lesions are characteristic of secondary syphilis. Human papillomavirus may cause singular warts (verruca vulgaris) or multiple warts in the anogenital area (condylomata acuminata). The latter are major problems in HIV-infected individuals.

ULCERS WITH OR WITHOUT ESCHARS

(Table 156-1) Cutaneous anthrax begins as a pruritic papule, which develops within days into an ulcer with surrounding vesicles and edema and then into an enlarging ulcer with a black eschar. Cutaneous anthrax may cause chronic nonhealing ulcers with an overlying dirty-gray membrane, although lesions may also mimic psoriasis, eczema, or impetigo. Ulceroglandular tularemia may have associated ulcerated skin lesions with painful regional adenopathy. Although buboes are the major cutaneous manifestation of plague, ulcers with eschars, papules, or pustules are also present in 25% of cases.

Mycobacterium ulcerans typically causes chronic skin ulcers on the extremities of individuals living in the tropics. *Mycobacterium leprae* may be associated with cutaneous ulcerations in patients with lepromatous leprosy related to Lucio's phenomenon, in which immune-mediated destruction of tissue bearing high concentrations of *M. leprae* bacilli occurs, usually several months after initiation of effective therapy. *Mycobacterium tuberculosis* also may cause ulcerations, papules, or erythematous macular lesions of the skin in both immunocompetent and immunocompromised patients.

Decubitus ulcers are due to tissue hypoxemia secondary to pressure-induced vascular insufficiency and may become secondarily infected with components of the skin and gastrointestinal flora, including anaerobes. Ulcerative lesions on the anterior shins may be due to pyoderma gangrenosum, which must be distinguished from similar lesions of infectious etiology by histologic evaluation of biopsy sites. Ulcerated lesions on the genitals may be either painful (chancroid) or painless (primary syphilis).

ERYSIPELAS

(Table 156-1) Erysipelas is due to *S. pyogenes* and is characterized by an abrupt onset of fiery-red swelling of the face or extremities. The distinctive features of erysipelas are well-defined indurated margins, particularly along the nasolabial fold; rapid progression; and intense pain. Flaccid bullae may develop during the second or third day of illness, but extension to deeper soft tissues is rare. Treatment with penicillin is effective; swelling may progress despite appropriate treatment, although fever, pain, and the intense red color diminish. Desquamation of the involved skin occurs 5–10 days into the illness. Infants and elderly adults are most commonly afflicted, and the severity of systemic toxicity varies.

CELLULITIS

(Table 156-1) Cellulitis is an acute inflammatory condition of the skin that is characterized by localized pain, erythema, swelling, and heat. It may be caused by indigenous flora colonizing the skin and appendages (e.g., *S. aureus* and *S. pyogenes*) or by a wide variety of exogenous bacteria. Because the exogenous bacteria involved in cellulitis occupy unique niches in nature, a thorough history (including epidemiologic data) provides important clues to etiology. When there is drainage, an open wound, or an obvious portal of entry, Gram's stain and culture provide a definitive diagnosis. In the absence of these findings, the bacterial etiology of cellulitis is difficult to establish, and in some cases staphylococcal and streptococcal cellulitis may have similar features. Even with needle aspiration of the leading edge or a punch biopsy of the cellulitis tissue itself, cultures are positive in only 20% of cases. This observation suggests that relatively low numbers of bacteria may cause cellulitis and that the expanding area of erythema within the skin may be a direct effect of extracellular toxins or of the soluble mediators of inflammation elicited by the host.

Bacteria may gain access to the epidermis through cracks in the skin, abrasions, cuts, burns, insect bites, surgical incisions, and IV catheters. Cellulitis caused by *S. aureus* spreads from a central localized infection, such as an abscess, folliculitis, or an infected foreign body (e.g., a splinter, a prosthetic device, or an IV catheter). MRSA is rapidly replacing methicillin-sensitive *S. aureus* (MSSA) as a cause of cellulitis in both inpatient and outpatient settings. Cellulitis caused by MSSA or MRSA is usually associated with a focal infection, such as a furuncle, a carbuncle, a surgical wound, or an abscess; the U.S. Food and Drug Administration preferentially refers to these types of infection as *purulent cellulitis*. In contrast, cellulitis due to *S. pyogenes* is a more rapidly spreading, diffuse process that is frequently associated with lymphangitis and fever and should be referred to as *nonpurulent cellulitis*. Recurrent streptococcal cellulitis of the lower extremities may be caused by organisms of group A, C, or G in association with chronic venous stasis or with saphenous venectomy for coronary artery bypass surgery. Streptococci also cause recurrent cellulitis among patients with chronic lymphedema resulting from elephantiasis, lymph node dissection, or Milroy's disease. Recurrent staphylococcal cutaneous infections are more common among individuals who have eosinophilia and elevated serum levels of IgE (Job's syndrome) and among nasal carriers of staphylococci. Cellulitis caused by *Streptococcus agalactiae* (group B *Streptococcus*) occurs primarily in elderly patients and those with diabetes mellitus or peripheral vascular disease. *Haemophilus influenzae* typically causes periorbital cellulitis in children in association with sinusitis, otitis media, or epiglottitis. It is unclear whether this form of cellulitis will (like meningitis) become less common as a result of the impressive efficacy of the *H. influenzae* type b vaccine.

Many other bacteria also cause cellulitis. It is fortunate that these organisms occur in such characteristic settings that a good history provides useful clues to the diagnosis. Cellulitis associated with cat bites and, to a lesser degree, with dog bites is commonly caused by *Pasteurella multocida*, although in the latter case *Staphylococcus intermedius* and *Capnocytophaga canimorsus* also must be considered. Sites of cellulitis and abscesses associated with dog bites and human bites also contain a variety of anaerobic organisms, including *Fusobacterium*, *Bacteroides*, aerobic and anaerobic streptococci, and *Eikenella corrodens*. *Pasteurella* is notoriously resistant to dicloxacillin and nafcillin but is sensitive to all other β -lactam antimicrobial agents as well as to quinolones, tetracycline, and erythromycin. Ampicillin/clavulanate, ampicillin/sulbactam, and cefoxitin are good choices for the treatment of animal or human bite infections. *Aeromonas hydrophila* causes aggressive cellulitis in tissues surrounding lacerations sustained in freshwater (lakes, rivers, and streams). This organism remains sensitive to aminoglycosides, fluoroquinolones, chloramphenicol, trimethoprim-sulfamethoxazole, and third-generation cephalosporins; it is resistant to ampicillin, however.

P. aeruginosa causes three types of soft tissue infection: ecthyma gangrenosum in neutropenic patients, hot-tub folliculitis, and cellulitis following penetrating injury. Most commonly, *P. aeruginosa* is introduced into the deep tissues when a person steps on a nail. Treatment includes surgical inspection and drainage, particularly if the injury also involves bone or joint capsule. Choices for empirical treatment while antimicrobial susceptibility data are awaited include an aminoglycoside, a third-generation cephalosporin (ceftazidime, cefoperazone, or cefotaxime), a semisynthetic penicillin (ticarcillin, mezlocillin, or piperacillin), or a fluoroquinolone (although drugs of the last class are not indicated for the treatment of children <13 years old).

Gram-negative bacillary cellulitis, including that due to *P. aeruginosa*, is most common among hospitalized, immunocompromised hosts. Cultures and sensitivity tests are critically important in this setting because of multidrug resistance (**Chap. 189**).

The gram-positive aerobic rod *Erysipelothrix rhusiopathiae* is most often associated with fish and domestic swine and causes cellulitis primarily in bone renderers and fishmongers. *E. rhusiopathiae* remains susceptible to most β -lactam antibiotics (including penicillin), erythromycin, clindamycin, tetracycline, and cephalosporins but is resistant to sulfonamides, chloramphenicol, and vancomycin. Its resistance to vancomycin, which is unusual among gram-positive bacteria, is of potential clinical significance since this agent is sometimes used in empirical therapy for skin infection. Fish food containing the water flea *Daphnia* is sometimes contaminated with *M. marinum*, which can cause cellulitis or granulomas on skin surfaces exposed to the water in aquariums or injured in swimming pools. Rifampin plus ethambutol has been an effective therapeutic combination in some cases, although no comprehensive studies have been undertaken. In addition, some strains of *M. marinum* are susceptible to tetracycline or to trimethoprim-sulfamethoxazole.

NECROTIZING FASCIITIS

(Table 156-1) Necrotizing fasciitis, formerly called streptococcal gangrene, may be associated with group A *Streptococcus* or mixed aerobic-anaerobic bacteria or may occur as a component of gas gangrene caused by *Clostridium perfringens*. Strains of MRSA that produce the Panton-Valentine leukocidin (PVL) toxin have been reported to cause necrotizing fasciitis. Early diagnosis may be difficult when pain or unexplained fever is the only presenting manifestation. Swelling then develops and is followed by brawny edema and tenderness. With progression, dark-red induration of the epidermis appears, along with bullae filled with blue or purple fluid. Later the skin becomes friable and takes on a bluish, maroon, or black color. By this stage, thrombosis of blood vessels in the dermal papillae (Fig. 156-1) is extensive. Extension of infection to the level of the deep fascia causes this tissue to take on a brownish-gray appearance. Rapid spread occurs along fascial planes, through venous channels and lymphatics. Patients in the later stages are toxic and frequently manifest shock and multiorgan failure.

Necrotizing fasciitis caused by mixed aerobic-anaerobic bacteria begins with a breach in the integrity of a mucous membrane barrier, such as the mucosa of the gastrointestinal or genitourinary tract. The portal can be a malignancy, a diverticulum, a hemorrhoid, an anal fissure, or a urethral tear. Other predisposing factors include peripheral vascular disease, diabetes mellitus, surgery, and penetrating injury to the abdomen. Leakage into the perineal area results in a syndrome called *Fournier's gangrene*, characterized by massive swelling of the scrotum and penis with extension into the perineum or the abdominal wall and the legs.

Necrotizing fasciitis caused by *S. pyogenes* has increased in frequency and severity since 1985. There are two distinct clinical presentations: those with no portal of entry and those with a defined portal of entry. Infections in the first category often begin deep at the site of a non-penetrating minor trauma, such as a bruise or a muscle strain. Seeding of the site via transient bacteremia is likely, although most patients deny antecedent streptococcal infection. The affected patients present with only severe pain and fever. Late in the course, the classic signs of necrotizing fasciitis, such as purple (violaceous) bullae, skin sloughing, and progressive toxicity, develop. In infections of the second type, *S. pyogenes* may reach the deep fascia from a site of cutaneous infection or penetrating trauma. These patients have early signs of superficial skin infection with progression to necrotizing fasciitis. In either case, toxicity is severe, and renal impairment may precede the development of shock. In 20–40% of cases, myositis occurs concomitantly, and, as in gas gangrene (see below), serum creatine phosphokinase levels may be markedly elevated. Necrotizing fasciitis due to mixed aerobic-anaerobic bacteria may be associated with gas in deep tissue, but gas usually is not present when the cause is *S. pyogenes* or MRSA. Prompt surgical exploration down to the deep fascia and muscle is essential. Necrotic tissue must be surgically removed, and Gram's staining and culture of excised tissue are useful in establishing whether group A streptococci, mixed aerobic-anaerobic bacteria, MRSA, or *Clostridium* species are present (see "Treatment," below).

MYOSITIS AND MYONECROSIS

(Table 156-1) Muscle involvement can occur with viral infection (e.g., influenza, dengue, or coxsackievirus B infection) or parasitic invasion (e.g., trichinellosis, cysticercosis, or toxoplasmosis). Although myalgia develops in most of these infections, severe muscle pain is the hallmark of pleurodynia (coxsackievirus B), trichinellosis, and bacterial infection. Acute rhabdomyolysis predictably occurs with clostridial and streptococcal myositis but may also be associated with influenza virus, echovirus, coxsackievirus, Epstein-Barr virus, and *Legionella* infections.

Pyomyositis is usually due to *S. aureus*, is common in tropical areas, and generally has no known portal of entry. Cases of pyomyositis caused by MRSA producing the PVL toxin have been described among children in the United States. Muscle infection begins at the exact site of blunt trauma or muscle strain. Infection remains localized, and shock does not develop unless organisms produce toxic shock syndrome toxin 1 or certain enterotoxins and the patient lacks antibodies to the toxin produced by the infecting organisms. In contrast, *S. pyogenes* may induce primary myositis (referred to as *streptococcal necrotizing myositis*) in association with severe systemic toxicity. Myonecrosis occurs concomitantly with necrotizing fasciitis in ~50% of cases. Both are part of the streptococcal toxic shock syndrome.

Gas gangrene usually follows severe penetrating injuries that result in interruption of the blood supply and introduction of soil into wounds. Such cases of traumatic gangrene are usually caused by the clostridial species *C. perfringens*, *C. septicum*, and *C. histolyticum*. Rarely, latent or recurrent gangrene can occur years after penetrating trauma; dormant spores that reside at the site of previous injury are most likely responsible. Spontaneous nontraumatic gangrene among patients with neutropenia, gastrointestinal malignancy, diverticulosis, or recent radiation therapy to the abdomen is caused by several clostridial species, of which *C. septicum* is the most commonly involved. The tolerance of this anaerobe to oxygen probably explains why it can initiate infection spontaneously in normal tissue anywhere in the body.

Gas gangrene of the uterus, especially that due to *Clostridium sordellii*, historically occurred as a consequence of illegal or self-induced abortion and nowadays also follows spontaneous abortion, vaginal delivery, and cesarean section. *C. sordellii* has also been implicated in medically induced abortion. Postpartum *C. sordellii* infections in young, previously healthy women present as a unique clinical picture: little or no fever, lack of a purulent discharge, refractory hypotension, extensive peripheral edema and effusions, hemoconcentration, and a markedly elevated white blood cell count. The infection is almost uniformly fatal, with death ensuing rapidly. *C. sordellii* and *C. novyi* have also been associated with cutaneous injection of black tar heroin; mortality rates are lower among the affected individuals, probably because their aggressive injection-site infections are readily apparent and diagnosis is therefore prompt.

Synergistic nonclostridial anaerobic myonecrosis, also known as necrotizing cutaneous myositis and synergistic necrotizing cellulitis, is a variant of necrotizing fasciitis caused by mixed aerobic and anaerobic bacteria with the exclusion of clostridial organisms (see "Necrotizing Fasciitis," above).

DIAGNOSIS

This chapter emphasizes the physical appearance and location of lesions within the soft tissues as important diagnostic clues. Other crucial considerations in narrowing the differential diagnosis are the temporal progression of the lesions as well as the patient's travel history, animal exposure or bite history, age, underlying disease status, and lifestyle. However, even the astute clinician may find it challenging to diagnose all infections of the soft tissues by history and inspection alone. Soft tissue radiography, CT (Fig. 156-2), and MRI may be useful in determining the depth of infection and should be performed when the patient has rapidly progressing lesions or evidence of a systemic inflammatory response syndrome. These tests are particularly valuable for defining a localized abscess or detecting gas in tissue. Unfortunately, they may reveal only soft tissue swelling and thus are not specific for fulminant infections such as necrotizing fasciitis or myonecrosis caused by group A *Streptococcus* (Fig. 156-2), where gas is not found in lesions.

Aspiration of the leading edge or punch biopsy with frozen section may be helpful if the results of imaging tests are positive, but false-negative results occur in ~80% of cases. There is some evidence that aspiration alone may be superior to injection and aspiration with normal saline. Frozen sections are especially useful in distinguishing SSSS from TEN and are quite valuable in cases of necrotizing fasciitis. Open surgical inspection, with debridement as indicated, is clearly the best way to determine the extent and severity of infection and to obtain material for Gram's staining and culture. Such an aggressive approach is important and may be lifesaving if undertaken early in the course of fulminant infections where there is evidence of systemic toxicity.



FIGURE 156-2 CT showing edema and inflammation of the left chest wall in a patient with necrotizing fasciitis and myonecrosis caused by group A *Streptococcus*.

A full description of the treatment of all the clinical entities described herein is beyond the scope of this chapter. As a guide to the clinician in selecting appropriate treatment, the antimicrobial agents useful in the most common and the most fulminant cutaneous infections are listed in **Table 156-2**.

Furuncles, carbuncles, and abscesses caused by MRSA and MSSA are common, and their treatment depends upon the size of the lesion. Furuncles <2.5 cm in diameter are usually treated with moist heat. Those that are larger (4.5 cm of erythema and induration) require surgical drainage, and the occurrence of these larger lesions in association with fever, chills, or leukocytosis requires both drainage and antibiotic treatment. A study in children demonstrated that surgical drainage of abscesses (mean diameter, 3.8 cm) was as effective when used alone as when combined with trimethoprim-sulfamethoxazole treatment. However, the rate of recurrence of

new lesions was lower in the group undergoing both drainage and antibiotic treatment.

Early and aggressive surgical exploration is essential in cases of suspected necrotizing fasciitis, myositis, or gangrene in order to (1) visualize the deep structures, (2) remove necrotic tissue, (3) reduce compartment pressure, and (4) obtain suitable material for Gram's staining and for aerobic and anaerobic cultures. Appropriate empirical antibiotic treatment for mixed aerobic-anaerobic infections could consist of ampicillin/sulbactam, ceftiofur, or the following combination: (1) clindamycin (600–900 mg IV every 8 h) or metronidazole (500 mg every 6 h) plus (2) ampicillin or ampicillin/sulbactam (1.5–3 g IV every 6 h) plus (3) gentamicin (1–1.5 mg/kg every 8 h). Group A streptococcal and clostridial infection of the fascia and/or muscle carries a mortality rate of 20–50% with penicillin treatment. In experimental models of streptococcal and clostridial necrotizing fasciitis/myositis, clindamycin has exhibited markedly superior efficacy, but no comparative clinical trials have been performed. A retrospective study of children with invasive group A streptococcal

TABLE 156-2 TREATMENT OF COMMON INFECTIONS OF THE SKIN

Diagnosis/Condition	Primary Treatment	Alternative Treatment	See Also Chap(s).
Animal bite (prophylaxis or early infection) ^a	Amoxicillin/clavulanate, 875/125 mg PO bid	Doxycycline, 100 mg PO bid	167e
Animal bite ^a (established infection)	Ampicillin/sulbactam, 1.5–3 g IV q6h	Clindamycin, 600–900 mg IV q8h, <i>plus</i> Ciprofloxacin, 400 mg IV q12h, <i>or</i> ceftiofur, 2 g IV q6h	167e
Bacillary angiomatosis	Erythromycin, 500 mg PO qid	Doxycycline, 100 mg PO bid	197
Herpes simplex (primary genital)	Acyclovir, 400 mg PO tid for 10 days	Famciclovir, 250 mg PO tid for 5–10 days, <i>or</i> Valacyclovir, 1000 mg PO bid for 10 days	216
Herpes zoster (immunocompetent host >50 years of age)	Acyclovir, 800 mg PO 5 times daily for 7–10 days	Famciclovir, 500 mg PO tid for 7–10 days, <i>or</i> Valacyclovir, 1000 mg PO tid for 7 days	217
Cellulitis (staphylococcal or streptococcal ^{b,c})	Nafcillin or oxacillin, 2 g IV q4–6h	Cefazolin, 1–2 g q8h, <i>or</i> Ampicillin/sulbactam, 1.5–3 g IV q6h, <i>or</i> Erythromycin, 0.5–1 g IV q6h, <i>or</i> Clindamycin, 600–900 mg IV q8h	172, 173
MRSA skin infection ^d	Vancomycin, 1 g IV q12h	Linezolid, 600 mg IV q12h	172
Necrotizing fasciitis (group A streptococcal ^b)	Clindamycin, 600–900 mg IV q6–8h, <i>plus</i> Penicillin G, 4 million units IV q4h	Clindamycin, 600–900 mg IV q6–8h, <i>plus</i> Cephalosporin (first- or second-generation)	173
Necrotizing fasciitis (mixed aerobes and anaerobes)	Ampicillin, 2 g IV q4h, <i>plus</i> Clindamycin, 600–900 mg IV q6–8h, <i>plus</i> Ciprofloxacin, 400 mg IV q6–8h	Vancomycin, 1 g IV q6h, <i>plus</i> Metronidazole, 500 mg IV q6h, <i>plus</i> Ciprofloxacin, 400 mg IV q6–8h	201
Gas gangrene	Clindamycin, 600–900 mg IV q6–8h, <i>plus</i> Penicillin G, 4 million units IV q4–6h	Clindamycin, 600–900 mg IV q6–8h, <i>plus</i> Ceftiofur, 2 g IV q6h	179

^a*Pasteurella multocida*, a species commonly associated with both dog and cat bites, is resistant to cephalexin, dicloxacillin, clindamycin, and erythromycin. *Eikenella corrodens*, a bacterium commonly associated with human bites, is resistant to clindamycin, penicillinase-resistant penicillins, and metronidazole but is sensitive to trimethoprim-sulfamethoxazole and fluoroquinolones. ^bThe frequency of erythromycin resistance in group A *Streptococcus* is currently ~5% in the United States but has reached 70–100% in some other countries. Most, but not all, erythromycin-resistant group A streptococci are susceptible to clindamycin. Approximately 90% of *Staphylococcus aureus* strains are sensitive to clindamycin, but resistance—both intrinsic and inducible—is increasing. ^cSevere hospital-acquired *S. aureus* infections or community-acquired *S. aureus* infections that are not responding to the β-lactam antibiotics recommended in this table may be caused by methicillin-resistant strains, requiring a switch to vancomycin, daptomycin, or linezolid. ^dSome strains of methicillin-resistant *S. aureus* (MRSA) remain sensitive to tetracycline and trimethoprim-sulfamethoxazole. Daptomycin (4 mg/kg IV q24h) or tigecycline (100-mg loading dose followed by 50 mg IV q12h) is an alternative treatment for MRSA.

infection demonstrated higher survival rates with clindamycin treatment than with β -lactam antibiotic therapy. Hyperbaric oxygen treatment also may be useful in gas gangrene due to clostridial species. Antibiotic treatment should be continued until all signs of systemic toxicity have resolved, all devitalized tissue has been removed, and granulation tissue has developed (Chaps. 173, 179, and 201).

In summary, infections of the skin and soft tissues are diverse in presentation and severity and offer a great challenge to the clinician. This chapter provides an approach to diagnosis and understanding of the pathophysiologic mechanisms involved in these infections. More in-depth information is found in chapters on specific infections.

157 Infectious Arthritis

Lawrence C. Madoff

Although *Staphylococcus aureus*, *Neisseria gonorrhoeae*, and other bacteria are the most common causes of infectious arthritis, various mycobacteria, spirochetes, fungi, and viruses also infect joints (Table 157-1). Since acute bacterial infection can destroy articular cartilage rapidly, all inflamed joints must be evaluated without delay to exclude noninfectious processes and determine appropriate antimicrobial therapy and drainage procedures. For more detailed information on infectious arthritis caused by specific organisms, the reader is referred to the chapters on those organisms.

Acute bacterial infection typically involves a single joint or a few joints. Subacute or chronic monoarthritis or oligoarthritis suggests mycobacterial or fungal infection; episodic inflammation is seen in syphilis, Lyme disease, and the reactive arthritis that follows enteric infections and chlamydial urethritis. Acute polyarticular inflammation occurs as an immunologic reaction during the course of endocarditis, rheumatic fever, disseminated neisserial infection, and acute hepatitis B. Bacteria and viruses occasionally infect multiple joints, the former most commonly in persons with rheumatoid arthritis.

APPROACH TO THE PATIENT: Infectious Arthritis

Aspiration of synovial fluid—an essential element in the evaluation of potentially infected joints—can be performed without difficulty in most cases by the insertion of a large-bore needle into the site of maximal fluctuance or tenderness or by the route of easiest access. Ultrasonography or fluoroscopy may be used to guide aspiration of difficult-to-localize effusions of the hip and, occasionally, the shoulder and other joints. Normal synovial fluid contains <180 cells (predominantly mononuclear cells) per microliter. Synovial cell counts averaging 100,000/ μ L (range, 25,000–250,000/ μ L), with >90% neutrophils, are characteristic of acute bacterial infections. Crystal-induced, rheumatoid, and other noninfectious inflammatory arthritides usually are associated with <30,000–50,000 cells/ μ L; cell counts of 10,000–30,000/ μ L, with 50–70% neutrophils and the remainder lymphocytes, are common in mycobacterial and fungal infections. Definitive diagnosis of an infectious process relies on identification of the pathogen in stained smears of synovial fluid, isolation of the pathogen from cultures of synovial fluid and blood, or detection of microbial nucleic acids and proteins by nucleic acid amplification (NAA)-based assays and immunologic techniques.

ACUTE BACTERIAL ARTHRITIS

PATHOGENESIS

Bacteria enter the joint from the bloodstream; from a contiguous site of infection in bone or soft tissue; or by direct inoculation during surgery, injection, animal or human bite, or trauma. In hematogenous infection, bacteria escape from synovial capillaries, which have no limiting basement membrane, and within hours provoke neutrophilic infiltration of the synovium. Neutrophils and bacteria enter the joint space; later, bacteria adhere to articular cartilage. Degradation of cartilage begins within 48 h as a result of increased intraarticular pressure, release of proteases and cytokines from chondrocytes and synovial macrophages, and invasion of the cartilage by bacteria and inflammatory cells. Histologic studies reveal bacteria lining the synovium and

TABLE 157-1 DIFFERENTIAL DIAGNOSIS OF ARTHRITIS SYNDROMES

Acute Monoarticular Arthritis	Chronic Monoarticular Arthritis	Polyarticular Arthritis
<i>Staphylococcus aureus</i>	<i>Mycobacterium tuberculosis</i>	<i>Neisseria meningitidis</i>
<i>Streptococcus pneumoniae</i>	Nontuberculous mycobacteria	<i>N. gonorrhoeae</i>
β -Hemolytic streptococci	<i>Borrelia burgdorferi</i>	Nongonococcal bacterial arthritis
Gram-negative bacilli	<i>Treponema pallidum</i>	Bacterial endocarditis
<i>Neisseria gonorrhoeae</i>	<i>Candida</i> spp.	<i>Candida</i> spp.
<i>Candida</i> spp.	<i>Sporothrix schenckii</i>	Poncet's disease (tuberculous rheumatism)
Crystal-induced arthritis	<i>Coccidioides immitis</i>	Hepatitis B virus
Fracture	<i>Blastomyces dermatitidis</i>	Parvovirus B19
Hemarthrosis	<i>Aspergillus</i> spp.	HIV
Foreign body	<i>Cryptococcus neoformans</i>	Human T-lymphotropic virus type I
Osteoarthritis	<i>Nocardia</i> spp.	Rubella virus
Ischemic necrosis	<i>Brucella</i> spp.	Arthropod-borne viruses
Monoarticular rheumatoid arthritis	Legg-Calvé-Perthes disease	Sickle cell disease flare
	Osteoarthritis	Reactive arthritis
		Serum sickness
		Acute rheumatic fever
		Inflammatory bowel disease
		Systemic lupus erythematosus
		Rheumatoid arthritis/Still's disease
		Other vasculitides
		Sarcoidosis

834 cartilage as well as abscesses extending into the synovium, cartilage, and—in severe cases—subchondral bone. Synovial proliferation results in the formation of a pannus over the cartilage, and thrombosis of inflamed synovial vessels develops. Bacterial factors that appear important in the pathogenesis of infective arthritis include various surface-associated adhesins in *S. aureus* that permit adherence to cartilage and endotoxins that promote chondrocyte-mediated breakdown of cartilage.

MICROBIOLOGY

The hematogenous route of infection is the most common route in all age groups, and nearly every bacterial pathogen is capable of causing septic arthritis. In infants, group B streptococci, gram-negative enteric bacilli, and *S. aureus* are the most common pathogens. Since the advent of the *Haemophilus influenzae* vaccine, the predominant causes among children <5 years of age have been *S. aureus*, *Streptococcus pyogenes* (group A *Streptococcus*), and (in some centers) *Kingella kingae*. Among young adults and adolescents, *N. gonorrhoeae* is the most commonly implicated organism. *S. aureus* accounts for most nongonococcal isolates in adults of all ages; gram-negative bacilli, pneumococci, and β -hemolytic streptococci—particularly groups A and B but also groups C, G, and F—are involved in up to one-third of cases in older adults, especially those with underlying comorbid illnesses.

Infections after surgical procedures or penetrating injuries are due most often to *S. aureus* and occasionally to other gram-positive bacteria or gram-negative bacilli. Infections with coagulase-negative staphylococci are unusual except after the implantation of prosthetic joints or arthroscopy. Anaerobic organisms, often in association with aerobic or facultative bacteria, are found after human bites and when decubitus ulcers or intraabdominal abscesses spread into adjacent joints. Polymicrobial infections complicate traumatic injuries with extensive contamination. Bites and scratches from cats and other animals may introduce *Pasteurella multocida* or *Bartonella henselae* into joints either directly or hematogenously, and bites from humans may introduce *Eikenella corrodens* or other components of the oral flora. Penetration of a sharp object through a shoe is associated with *Pseudomonas aeruginosa* arthritis in the foot.

NONGONOCOCCAL BACTERIAL ARTHRITIS

Epidemiology Although hematogenous infections with virulent organisms such as *S. aureus*, *H. influenzae*, and pyogenic streptococci occur in healthy persons, there is an underlying host predisposition in many cases of septic arthritis. Patients with rheumatoid arthritis have the highest incidence of infective arthritis (most often secondary to *S. aureus*) because of chronically inflamed joints; glucocorticoid therapy; and frequent breakdown of rheumatoid nodules, vasculitic ulcers, and skin overlying deformed joints. Diabetes mellitus, glucocorticoid therapy, hemodialysis, and malignancy all carry an increased risk of infection with *S. aureus* and gram-negative bacilli. Tumor necrosis factor inhibitors (e.g., etanercept, infliximab), which increasingly are used for the treatment of rheumatoid arthritis, predispose to mycobacterial infections and possibly to other pyogenic bacterial infections and could be associated with septic arthritis in this population. Pneumococcal infections complicate alcoholism, deficiencies of humoral immunity, and hemoglobinopathies. Pneumococci, *Salmonella* species, and *H. influenzae* cause septic arthritis in persons infected with HIV. Persons with primary immunoglobulin deficiency are at risk for mycoplasmal arthritis, which results in permanent joint damage if tetracycline and replacement therapy with IV immunoglobulin are not administered promptly. IV drug users acquire staphylococcal and streptococcal infections from their own flora and acquire pseudomonal and other gram-negative infections from drugs and injection paraphernalia.

Clinical Manifestations Some 90% of patients present with involvement of a single joint—most commonly the knee; less frequently the hip; and still less often the shoulder, wrist, or elbow. Small joints of the hands and feet are more likely to be affected after direct inoculation or a bite. Among IV drug users, infections of the spine, sacroiliac joints, and sternoclavicular joints (Fig. 157-1) are more common than infections of the appendicular skeleton. Polyarticular infection is most



FIGURE 157-1 Acute septic arthritis of the sternoclavicular joint.

A man in his forties with a history of cirrhosis presented with a new onset of fever and lower neck pain. He had no history of IV drug use or previous catheter placement. Jaundice and a painful swollen area over his left sternoclavicular joint were evident on physical examination. Cultures of blood drawn at admission grew group B *Streptococcus*. The patient recovered after treatment with IV penicillin. (Courtesy of Francisco M. Marty, MD, Brigham and Women's Hospital, Boston; with permission.)

common among patients with rheumatoid arthritis and may resemble a flare of the underlying disease.

The usual presentation consists of moderate to severe pain that is uniform around the joint, effusion, muscle spasm, and decreased range of motion. Fever in the range of 38.3–38.9°C (101–102°F) and sometimes higher is common but may not be present, especially in persons with rheumatoid arthritis, renal or hepatic insufficiency, or conditions requiring immunosuppressive therapy. The inflamed, swollen joint is usually evident on examination except in the case of a deeply situated joint such as the hip, shoulder, or sacroiliac joint. Cellulitis, bursitis, and acute osteomyelitis, which may produce a similar clinical picture, should be distinguished from septic arthritis by their greater range of motion and less-than-circumferential swelling. A focus of extraarticular infection, such as a boil or pneumonia, should be sought. Peripheral-blood leukocytosis with a left shift and elevation of the erythrocyte sedimentation rate or C-reactive protein level are common.

Plain radiographs show evidence of soft-tissue swelling, joint-space widening, and displacement of tissue planes by the distended capsule. Narrowing of the joint space and bony erosions indicate advanced infection and a poor prognosis. Ultrasound is useful for detecting effusions in the hip, and CT or MRI can demonstrate infections of the sacroiliac joint, the sternoclavicular joint, and the spine very well.

Laboratory Findings Specimens of peripheral blood and synovial fluid should be obtained before antibiotics are administered. Blood cultures are positive in up to 50–70% of *S. aureus* infections but are less frequently positive in infections due to other organisms. The synovial fluid is turbid, serosanguineous, or frankly purulent. Gram-stained smears confirm the presence of large numbers of neutrophils. Levels of total protein and lactate dehydrogenase in synovial fluid are elevated, and the glucose level is depressed; however, these findings are not specific for infection, and measurement of these levels is not necessary for diagnosis. The synovial fluid should be examined for crystals, because gout and pseudogout can resemble septic arthritis clinically, and infection and crystal-induced disease occasionally occur together. Organisms are seen on synovial fluid smears in nearly three-quarters of infections with *S. aureus* and streptococci and in 30–50% of infections due to gram-negative and other bacteria. Cultures of synovial fluid are positive in >90% of cases. Inoculation of synovial fluid into bottles containing liquid media for blood cultures increases the yield

of a culture, especially if the pathogen is a fastidious organism or the patient is taking an antibiotic. NAA-based assays for bacterial DNA, when available, can be useful for the diagnosis of partially treated or culture-negative bacterial arthritis.

TREATMENT NONGONOCOCCAL BACTERIAL ARTHRITIS

Prompt administration of systemic antibiotics and drainage of the involved joint can prevent destruction of cartilage, postinfectious degenerative arthritis, joint instability, or deformity. Once samples of blood and synovial fluid have been obtained for culture, empirical antibiotics should be given that are directed against the bacteria visualized on smears or the pathogens that are likely in light of the patient's age and risk factors. Initial therapy should consist of IV administration of bactericidal agents; direct instillation of antibiotics into the joint is not necessary to achieve adequate levels in synovial fluid and tissue. An IV third-generation cephalosporin such as cefotaxime (1 g every 8 h) or ceftriaxone (1–2 g every 24 h) provides adequate empirical coverage for most community-acquired infections in adults when smears show no organisms. IV vancomycin (1 g every 12 h) is used if there are gram-positive cocci on the smear. If methicillin-resistant *S. aureus* is an unlikely pathogen (e.g., when it is not widespread in the community), either oxacillin or nafcillin (2 g every 4 h) should be given. In addition, an aminoglycoside or third-generation cephalosporin should be given to IV drug users and to other patients in whom *P. aeruginosa* may be the responsible agent.

Definitive therapy is based on the identity and antibiotic susceptibility of the bacteria isolated in culture. Infections due to staphylococci are treated with oxacillin, nafcillin, or vancomycin for 4 weeks. Pneumococcal and streptococcal infections due to penicillin-susceptible organisms respond to 2 weeks of therapy with penicillin G (2 million units IV every 4 h); infections caused by *H. influenzae* and by strains of *Streptococcus pneumoniae* that are resistant to penicillin are treated with cefotaxime or ceftriaxone for 2 weeks. Most enteric gram-negative infections can be cured in 3–4 weeks by a second- or third-generation cephalosporin given IV or by a fluoroquinolone such as levofloxacin (500 mg IV or PO every 24 h). *P. aeruginosa* infection should be treated for at least 2 weeks with a combination regimen composed of an aminoglycoside plus either an extended-spectrum penicillin such as mezlocillin (3 g IV every 4 h) or an antipseudomonal cephalosporin such as ceftazidime (1 g IV every 8 h). If tolerated, this regimen is continued for an additional 2 weeks; alternatively, a fluoroquinolone such as ciprofloxacin (750 mg PO twice daily) is given by itself or with the penicillin or cephalosporin in place of the aminoglycoside.

Timely drainage of pus and necrotic debris from the infected joint is required for a favorable outcome. Needle aspiration of readily accessible joints such as the knee may be adequate if loculations or particulate matter in the joint does not prevent its thorough decompression. Arthroscopic drainage and lavage may be employed initially or within several days if repeated needle aspiration fails to relieve symptoms, decrease the volume of the effusion and the synovial white cell count, and clear bacteria from smears and cultures. In some cases, arthrotomy is necessary to remove loculations and debride infected synovium, cartilage, or bone. Septic arthritis of the hip is best managed with arthrotomy, particularly in young children, in whom infection threatens the viability of the femoral head. Septic joints do not require immobilization except for pain control before symptoms are alleviated by treatment. Weight bearing should be avoided until signs of inflammation have subsided, but frequent passive motion of the joint is indicated to maintain full mobility. Although addition of glucocorticoids to antibiotic treatment improves the outcome of *S. aureus* arthritis in experimental animals, no clinical trials have evaluated this approach in humans.

GONOCOCCAL ARTHRITIS

Epidemiology Although its incidence has declined in recent years, gonococcal arthritis (Chap. 181) has accounted for up to 70% of episodes of infectious arthritis in persons <40 years of age in the

United States. Arthritis due to *N. gonorrhoeae* is a consequence of bacteremia arising from gonococcal infection or, more frequently, from asymptomatic gonococcal mucosal colonization of the urethra, cervix, or pharynx. Women are at greatest risk during menses and during pregnancy and overall are two to three times more likely than men to develop disseminated gonococcal infection (DGI) and arthritis. Persons with complement deficiencies, especially of the terminal components, are prone to recurrent episodes of gonococcemia. Strains of gonococci that are most likely to cause DGI include those which produce transparent colonies in culture, have the type IA outer-membrane protein, or are of the AUH-auxotroph type.

Clinical Manifestations and Laboratory Findings The most common manifestation of DGI is a syndrome of fever, chills, rash, and articular symptoms. Small numbers of papules that progress to hemorrhagic pustules develop on the trunk and the extensor surfaces of the distal extremities. Migratory arthritis and tenosynovitis of the knees, hands, wrists, feet, and ankles are prominent. The cutaneous lesions and articular findings are believed to be the consequence of an immune reaction to circulating gonococci and immune-complex deposition in tissues. Thus, cultures of synovial fluid are consistently negative, and blood cultures are positive in fewer than 45% of patients. Synovial fluid may be difficult to obtain from inflamed joints and usually contains only 10,000–20,000 leukocytes/ μ L.

True gonococcal septic arthritis is less common than the DGI syndrome and always follows DGI, which is unrecognized in one-third of patients. A single joint such as the hip, knee, ankle, or wrist is usually involved. Synovial fluid, which contains >50,000 leukocytes/ μ L, can be obtained with ease; the gonococcus is evident only occasionally in Gram-stained smears, and cultures of synovial fluid are positive in fewer than 40% of cases. Blood cultures are almost always negative.

Because it is difficult to isolate gonococci from synovial fluid and blood, specimens for culture should be obtained from potentially infected mucosal sites. NAA-based urine tests also may be positive. Cultures and Gram-stained smears of skin lesions are occasionally positive. All specimens for culture should be plated onto Thayer-Martin agar directly or in special transport media at the bedside and transferred promptly to the microbiology laboratory in an atmosphere of 5% CO₂, as generated in a candle jar. NAA-based assays are extremely sensitive in detecting gonococcal DNA in synovial fluid. A dramatic alleviation of symptoms within 12–24 h after the initiation of appropriate antibiotic therapy supports a clinical diagnosis of the DGI syndrome if cultures are negative.

TREATMENT GONOCOCCAL ARTHRITIS

Initial treatment consists of ceftriaxone (1 g IV or IM every 24 h) to cover possible penicillin-resistant organisms. Once local and systemic signs are clearly resolving, the 7-day course of therapy can be completed with an oral fluoroquinolone such as ciprofloxacin (500 mg twice daily) if the organism is known to be susceptible. If penicillin-susceptible organisms are isolated, amoxicillin (500 mg three times daily) may be used. Suppurative arthritis usually responds to needle aspiration of involved joints and 7–14 days of antibiotic treatment. Arthroscopic lavage or arthrotomy is rarely required. Patients with DGI should be treated for *Chlamydia trachomatis* infection unless this infection is ruled out by appropriate testing.

It is noteworthy that arthritis symptoms similar to those seen in DGI occur in meningococcemia. A dermatitis-arthritis syndrome, purulent monoarthritis, and reactive polyarthritis have been described. All respond to treatment with IV penicillin.

SPIROCHETAL ARTHRITIS

LYME DISEASE

Lyme disease (Chap. 210) due to infection with the spirochete *Borrelia burgdorferi* causes arthritis in up to 60% of persons who are not treated. Intermittent arthralgias and myalgias—but not arthritis—occur within days or weeks of inoculation of the spirochete by the *Ixodes* tick. Later,

836 there are three patterns of joint disease: (1) Fifty percent of untreated persons experience intermittent episodes of monoarthritis or oligoarthritis involving the knee and/or other large joints. The symptoms wax and wane without treatment over months, and each year 10–20% of patients report loss of joint symptoms. (2) Twenty percent of untreated persons develop a pattern of waxing and waning arthralgias. (3) Ten percent of untreated patients develop chronic inflammatory synovitis that results in erosive lesions and destruction of the joint. Serologic tests for IgG antibodies to *B. burgdorferi* are positive in more than 90% of persons with Lyme arthritis, and an NAA-based assay detects *Borrelia* DNA in 85%.

TREATMENT LYME ARTHRITIS

Lyme arthritis generally responds well to therapy. A regimen of oral doxycycline (100 mg twice daily for 30 days), oral amoxicillin (500 mg four times daily for 30 days), or parenteral ceftriaxone (2 g/d for 2–4 weeks) is recommended. Patients who do not respond to a total of 2 months of oral therapy or 1 month of parenteral therapy are unlikely to benefit from additional antibiotic therapy and are treated with anti-inflammatory agents or synovectomy. Failure of therapy is associated with host features such as the human leukocyte antigen DR4 (HLA-DR4) genotype, persistent reactivity to OspA (outer-surface protein A), and the presence of hLFA-1 (human leukocyte function-associated antigen 1), which cross-reacts with OspA.

SYPHILITIC ARTHRITIS

Articular manifestations occur in different stages of syphilis (Chap. 206). In early congenital syphilis, periarticular swelling and immobilization of the involved limbs (*Parrot's pseudoparalysis*) complicate osteochondritis of long bones. *Clutton's joint*, a late manifestation of congenital syphilis that typically develops between ages 8 and 15 years, is caused by chronic painless synovitis with effusions of large joints, particularly the knees and elbows. Secondary syphilis may be associated with arthralgias, with symmetric arthritis of the knees and ankles and occasionally of the shoulders and wrists, and with sacroiliitis. The arthritis follows a subacute to chronic course with a mixed mononuclear and neutrophilic synovial-fluid pleocytosis (typical cell counts, 5000–15,000/ μ L). Immunologic mechanisms may contribute to the arthritis, and symptoms usually improve rapidly with penicillin therapy. In tertiary syphilis, Charcot's joint results from sensory loss due to tabes dorsalis. Penicillin is not helpful in this setting.

MYCOBACTERIAL ARTHRITIS

Tuberculous arthritis (Chap. 202) accounts for ~1% of all cases of tuberculosis and 10% of extrapulmonary cases. The most common presentation is chronic granulomatous monoarthritis. An unusual syndrome, *Poncet's disease*, is a reactive symmetric form of polyarthritis that affects persons with visceral or disseminated tuberculosis. No mycobacteria are found in the joints, and symptoms resolve with antituberculous therapy.

Unlike tuberculous osteomyelitis (Chap. 158), which typically involves the thoracic and lumbar spine (50% of cases), tuberculous arthritis primarily involves the large weight-bearing joints, in particular the hips, knees, and ankles, and only occasionally involves smaller non-weight-bearing joints. Progressive monoarticular swelling and pain develop over months or years, and systemic symptoms are seen in only half of all cases. Tuberculous arthritis occurs as part of a disseminated primary infection or through late reactivation, often in persons with HIV infection or other immunocompromised hosts. Coexistent active pulmonary tuberculosis is unusual.

Aspiration of the involved joint yields fluid with an average cell count of 20,000/ μ L, with ~50% neutrophils. Acid-fast staining of the fluid yields positive results in fewer than one-third of cases, and cultures are positive in 80%. Culture of synovial tissue taken at biopsy is positive in ~90% of cases and shows granulomatous inflammation in most. NAA methods can shorten the time to diagnosis to 1 or 2 days. Radiographs reveal peripheral erosions at the points of synovial attachment,

periarticular osteopenia, and eventually joint-space narrowing. Therapy for tuberculous arthritis is the same as that for tuberculous pulmonary disease, requiring the administration of multiple agents for 6–9 months. Therapy is more prolonged in immunosuppressed individuals such as those infected with HIV.

Various atypical mycobacteria (Chap. 204) found in water and soil may cause chronic indolent arthritis. Such disease results from trauma and direct inoculation associated with farming, gardening, or aquatic activities. Smaller joints, such as the digits, wrists, and knees, are usually involved. Involvement of tendon sheaths and bursae is typical. The mycobacterial species involved include *Mycobacterium marinum*, *M. avium-intracellulare*, *M. terrae*, *M. kansasii*, *M. fortuitum*, and *M. chelonae*. In persons who have HIV infection or are receiving immunosuppressive therapy, hematogenous spread to the joints has been reported for *M. kansasii*, *M. avium* complex, and *M. haemophilum*. Diagnosis usually requires biopsy and culture, and therapy is based on antimicrobial susceptibility patterns.

FUNGAL ARTHRITIS

Fungi are an unusual cause of chronic monoarticular arthritis. Granulomatous articular infection with the endemic dimorphic fungi *Coccidioides immitis*, *Blastomyces dermatitidis*, and (less commonly) *Histoplasma capsulatum* (Fig. 157-2) results from hematogenous seeding or direct extension from bony lesions in persons with disseminated disease. Joint involvement is an unusual complication of sporotrichosis (infection with *Sporothrix schenckii*) among gardeners and other persons who work with soil or sphagnum moss. Articular sporotrichosis is six times more common among men than among women, and alcoholics and other debilitated hosts are at risk for polyarticular infection.

Candida infection involving a single joint—usually the knee, hip, or shoulder—results from surgical procedures, intraarticular injections, or (among critically ill patients with debilitating illnesses such as diabetes mellitus or hepatic or renal insufficiency and patients receiving immunosuppressive therapy) hematogenous spread. *Candida* infections in IV drug users typically involve the spine, sacroiliac joints, or other fibrocartilaginous joints. Unusual cases of arthritis due to *Aspergillus* species, *Cryptococcus neoformans*, *Pseudallescheria boydii*, and the dematiaceous fungi also have resulted from direct inoculation or disseminated hematogenous infection in immunocompromised persons. In the United States, a 2012 national outbreak of fungal arthritis (and meningitis) caused by *Exserohilum rostratum* was linked to intraspinal and intraarticular injection of a contaminated preparation of methylprednisolone acetate.

The synovial fluid in fungal arthritis usually contains 10,000–40,000 cells/ μ L, with ~70% neutrophils. Stained specimens and cultures of synovial tissue often confirm the diagnosis of fungal arthritis when studies of synovial fluid give negative results. Treatment consists of drainage and lavage of the joint and systemic administration of an antifungal agent directed at a specific pathogen. The doses and duration of therapy are the same as for disseminated disease (see Part 8, Section 16). Intraarticular instillation of amphotericin B has been used in addition to IV therapy.

VIRAL ARTHRITIS


Viruses produce arthritis by infecting synovial tissue during systemic infection or by provoking an immunologic reaction that involves joints. As many as 50% of women report persistent arthralgias, and 10% report frank arthritis within 3 days of the rash that follows natural infection with rubella virus and within 2–6 weeks after receipt of live-virus vaccine. Episodes of symmetric inflammation of fingers, wrists, and knees uncommonly recur for >1 year, but a syndrome of chronic fatigue, low-grade fever, headaches, and myalgias can persist for months or years. IV immunoglobulin has been helpful in selected cases. Self-limited monoarticular or migratory polyarthritis may develop within 2 weeks of the parotitis of mumps; this sequela is more common among men than among women. Approximately



FIGURE 157-2 Chronic arthritis caused by *Histoplasma capsulatum* in the left knee. **A.** A man in his sixties from El Salvador presented with a history of progressive knee pain and difficulty walking for several years. He had undergone arthroscopy for a meniscal tear 7 years before presentation (without relief) and had received several intraarticular glucocorticoid injections. The patient developed significant deformity of the knee over time, including a large effusion in the lateral aspect. **B.** An x-ray of the knee showed multiple abnormalities, including severe medial femorotibial joint-space narrowing, several large subchondral cysts within the tibia and the patellofemoral compartment, a large suprapatellar joint effusion, and a large soft tissue mass projecting laterally over the knee. **C.** MRI further defined these abnormalities and demonstrated the cystic nature of the lateral knee abnormality. Synovial biopsies demonstrated chronic inflammation with giant cells, and cultures grew *H. capsulatum* after 3 weeks of incubation. All clinical cystic lesions and the effusion resolved after 1 year of treatment with itraconazole. The patient underwent a left total knee replacement for definitive treatment. (Courtesy of Francisco M. Marty, MD, Brigham and Women's Hospital, Boston; with permission.)

10% of children and 60% of women develop arthritis after infection with parvovirus B19. In adults, arthropathy sometimes occurs without fever or rash. Pain and stiffness, with less prominent swelling (primarily of the hands but also of the knees, wrists, and ankles), usually resolve within weeks, although a small proportion of patients develop chronic arthropathy.

About 2 weeks before the onset of jaundice, up to 10% of persons with acute hepatitis B develop an immune complex-mediated, serum sickness-like reaction with maculopapular rash, urticaria, fever, and arthralgias. Less common developments include symmetric arthritis involving the hands, wrists, elbows, or ankles and morning stiffness that resembles a flare of rheumatoid arthritis. Symptoms resolve at the time jaundice develops. Many persons with chronic hepatitis C infection report persistent arthralgia or arthritis, both in the presence and in the absence of cryoglobulinemia.

 Painful arthritis involving larger joints often accompanies the fever and rash of several arthropod-borne viral infections, including those caused by chikungunya, O'nyong-nyong, Ross River, Mayaro, and Barmah Forest viruses (Chap. 233). Symmetric arthritis involving the hands and wrists may occur during the convalescent phase of infection with lymphocytic choriomeningitis virus. Patients infected with an enterovirus frequently report arthralgias, and echovirus has been isolated from patients with acute polyarthrititis.

Several arthritis syndromes are associated with HIV infection. Reactive arthritis with painful lower-extremity oligoarthritis often follows an episode of urethritis in HIV-infected persons. HIV-associated reactive arthritis appears to be extremely common among persons with the HLA-B27 haplotype, but sacroiliac joint disease is unusual and is seen mostly in the absence of HLA-B27. Up to one-third of HIV-infected persons with psoriasis develop psoriatic arthritis. Painless monoarthropathy and persistent symmetric polyarthropathy occasionally complicate HIV infection. Chronic persistent oligoarthritis of the shoulders, wrists, hands, and knees occurs in women infected with human T-lymphotropic virus type I. Synovial thickening, destruction of articular cartilage, and leukemic-appearing atypical lymphocytes in synovial fluid are characteristic, but progression to T cell leukemia is unusual.

PARASITIC ARTHRITIS

Arthritis due to parasitic infection is rare. The guinea worm *Dracunculus medinensis* may cause destructive joint lesions in the lower extremities as migrating gravid female worms invade joints or cause ulcers in adjacent

soft tissues that become secondarily infected. Hydatid cysts infect bones in 1–2% of cases of infection with *Echinococcus granulosus*. The expanding destructive cystic lesions may spread to and destroy adjacent joints, particularly the hip and pelvis. In rare cases, chronic synovitis has been associated with the presence of schistosomal eggs in synovial biopsies. Monoarticular arthritis in children with lymphatic filariasis appears to respond to therapy with diethylcarbamazine even in the absence of microfilariae in synovial fluid. Reactive arthritis has been attributed to hookworm, *Strongyloides*, *Cryptosporidium*, and *Giardia* infection in case reports, but confirmation is required.

POSTINFECTIOUS OR REACTIVE ARTHRITIS

Reactive polyarthrititis develops several weeks after ~1% of cases of nongonococcal urethritis and 2% of enteric infections, particularly those due to *Yersinia enterocolitica*, *Shigella flexneri*, *Campylobacter jejuni*, and *Salmonella* species. Only a minority of these patients have the other findings of classic reactive arthritis, including urethritis, conjunctivitis, uveitis, oral ulcers, and rash. Studies have identified microbial DNA or antigen in synovial fluid or blood, but the pathogenesis of this condition is poorly understood.

Reactive arthritis is most common among young men (except after *Yersinia* infection) and has been linked to the HLA-B27 locus as a potential genetic predisposing factor. Patients report painful, asymmetric oligoarthritis that affects mainly the knees, ankles, and feet. Low-back pain is common, and radiographic evidence of sacroiliitis is found in patients with long-standing disease. Most patients recover within 6 months, but prolonged recurrent disease is more common in cases that follow chlamydial urethritis. Anti-inflammatory agents help relieve symptoms, but the role of prolonged antibiotic therapy in eliminating microbial antigen from the synovium is controversial.

Migratory polyarthrititis and fever constitute the usual presentation of acute rheumatic fever in adults (Chap. 381). This presentation is distinct from that of poststreptococcal reactive arthritis, which also follows infections with group A *Streptococcus* but is not migratory, lasts beyond the typical 3-week maximum of acute rheumatic fever, and responds poorly to aspirin.

INFECTIONS IN PROSTHETIC JOINTS

Infection complicates 1–4% of total joint replacements. The majority of infections are acquired intraoperatively or immediately postoperatively as a result of wound breakdown or infection; less commonly,

838 these joint infections develop later after joint replacement and are the result of hematogenous spread or direct inoculation. The presentation may be acute, with fever, pain, and local signs of inflammation, especially in infections due to *S. aureus*, pyogenic streptococci, and enteric bacilli. Alternatively, infection may persist for months or years without causing constitutional symptoms when less virulent organisms, such as coagulase-negative staphylococci or diphtheroids, are involved. Such indolent infections usually are acquired during joint implantation and are discovered during evaluation of chronic unexplained pain or after a radiograph shows loosening of the prosthesis; the erythrocyte sedimentation rate and C-reactive protein level are usually elevated in such cases.

The diagnosis is best made by needle aspiration of the joint; accidental introduction of organisms during aspiration must be avoided meticulously. Synovial fluid pleocytosis with a predominance of polymorphonuclear leukocytes is highly suggestive of infection, since other inflammatory processes uncommonly affect prosthetic joints. Culture and Gram's stain usually yield the responsible pathogen. Sonication of explanted prosthetic material can improve the yield of culture, presumably by breaking up bacterial biofilms on the surfaces of prostheses. Use of special media for unusual pathogens such as fungi, atypical mycobacteria, and *Mycoplasma* may be necessary if routine and anaerobic cultures are negative.

TREATMENT PROSTHETIC JOINT INFECTIONS

Treatment includes surgery and high doses of parenteral antibiotics, which are given for 4–6 weeks because bone is usually involved. In most cases, the prosthesis must be replaced to cure the infection. Implantation of a new prosthesis is best delayed for several weeks or months because relapses of infection occur most commonly within this time frame. In some cases, reimplantation is not possible, and the patient must manage without a joint, with a fused joint, or even with amputation. Cure of infection without removal of the prosthesis is occasionally possible in cases that are due to streptococci or pneumococci and that lack radiologic evidence of loosening of the prosthesis. In these cases, antibiotic therapy must be initiated within several days of the onset of infection, and the joint should be drained vigorously by open arthrotomy or arthroscopically. In selected patients who prefer to avoid the high morbidity rate associated with joint removal and reimplantation, suppression of the infection with antibiotics may be a reasonable goal. A high cure rate with retention of the prosthesis has been reported when the combination of oral rifampin and ciprofloxacin is given for 3–6 months to persons with staphylococcal prosthetic joint infection of short duration. This approach, which is based on the ability of rifampin to kill organisms adherent to foreign material and in the stationary growth phase, requires confirmation in prospective trials.

PREVENTION

To avoid the disastrous consequences of infection, candidates for joint replacement should be selected with care. Rates of infection are particularly high among patients with rheumatoid arthritis, persons who have undergone previous surgery on the joint, and persons with medical conditions requiring immunosuppressive therapy. Perioperative antibiotic prophylaxis, usually with cefazolin, and measures to decrease intraoperative contamination, such as laminar flow, have lowered the rates of perioperative infection to <1% in many centers. After implantation, measures should be taken to prevent or rapidly treat extra-articular infections that might give rise to hematogenous spread to the prosthesis. The effectiveness of prophylactic antibiotics for the prevention of hematogenous infection after dental procedures has not been demonstrated; in fact, viridans streptococci and other components of the oral flora are extremely unusual causes of prosthetic joint infection. Accordingly, the American Dental Association and the American Academy of Orthopaedic Surgeons do not recommend antibiotic

prophylaxis for most dental patients with total joint replacements and have stated that there is no convincing evidence to support its use. Similarly, guidelines issued by the American Urological Association and the American Academy of Orthopaedic Surgeons do not recommend the use of prophylactic antibiotics for most patients with prosthetic joints who are undergoing urologic procedures but state that prophylaxis should be considered in certain situations—e.g., for patients (especially immunocompromised patients) who are undergoing a procedure posing a relatively high risk of bacteremia (such as lithotripsy or surgery involving bowel segments).

ACKNOWLEDGMENTS

The contributions of James H. Maguire and the late Scott J. Thaler to this chapter in earlier editions are gratefully acknowledged.

158 Osteomyelitis

Werner Zimmerli

Osteomyelitis, an infection of bone, can be caused by various microorganisms that arrive at bone through different routes. Spontaneous hematogenous osteomyelitis may occur in otherwise healthy individuals, whereas local microbial spread mainly affects either individuals who have underlying disease (e.g., vascular insufficiency) or patients who have compromised skin or other tissue barriers, with consequent exposure of bone. The latter situation typically follows surgery involving bone, such as sternotomy or orthopedic repair.

The manifestations of osteomyelitis are different in children and adults. In children circulating microorganisms seed mainly long bones, whereas in adults the vertebral column is the most commonly affected site.

Management of osteomyelitis differs greatly depending on whether an implant is involved. The most important aim of the management of either type of osteomyelitis is to prevent progression to chronic osteomyelitis by rapid diagnosis and prompt treatment. Device-related bone and joint infection necessitates a multidisciplinary approach requiring antibiotic therapy and, in many cases, surgical removal of the device. The optimal duration of antibiotic treatment has not been established for any type of osteomyelitis in clinical trials. Therefore, the recommendations for therapy in this chapter reflect only expert opinions.

CLASSIFICATION

There is no generally accepted, comprehensive system for classification of osteomyelitis, primarily because of the multifaceted presentation of this infection. Different specialists are confronted with different facets of bone disease. Most often, however, general practitioners or internists are the first to encounter patients with the initial signs and symptoms of osteomyelitis. These primary care physicians should be able to recognize this disease in any of its forms. Osteomyelitis cases can be classified by various criteria, including pathogenesis, duration of infection, location of infection, and presence or absence of foreign material. The widely used Cierny-Mader staging system classifies osteomyelitis according to anatomic site, comorbidity, and radiographic findings, with stratification of long-bone osteomyelitis to optimize surgical management; this system encompasses both systemic and local factors affecting immune status, metabolism, and local vascularity.

Any of three mechanisms can underlie osteomyelitis: (1) hematogenous spread; (2) spread from a contiguous site following surgery; and (3) secondary infection in the setting of vascular insufficiency or concomitant neuropathy. Hematogenous osteomyelitis in adults typically involves the vertebral column. In only about half of patients can a primary focus be detected. The most common primary foci of infection are the urinary tract, skin/soft tissue, intravascular catheterization

sites, and the endocardium. Spread from a contiguous source follows either bone trauma or surgical intervention. Wound infection leading to osteomyelitis typically occurs after cardiovascular intervention involving the sternum, orthopedic repair, or prosthetic joint insertion. Osteomyelitis secondary to vascular insufficiency or peripheral neuropathy most often follows chronic, progressively deep skin and soft tissue infection of the foot. The most common underlying condition is diabetes. In diabetes that is poorly controlled, the *diabetic foot syndrome* is caused by skin, soft tissue, and bone ischemia combined with motor, sensory, and autonomic neuropathy.

Classification of osteomyelitis according to the duration of infection, although ill defined (because there is no clear time limit for the transition from acute to chronic osteomyelitis), is useful because the management of acute and chronic osteomyelitis differs. Whereas acute osteomyelitis can generally be treated with antibiotics alone, antibiotic treatment for chronic osteomyelitis should be combined with debridement surgery. Acute hematogenous or contiguous osteomyelitis evolves over a short period—i.e., a few days or weeks. In contrast, subacute or chronic osteomyelitis lasts for weeks or months before treatment is started. Typical examples of a subacute course are vertebral osteomyelitis due to tuberculosis or brucellosis and delayed implant-associated infections caused mainly by low-virulence microorganisms (coagulase-negative staphylococci, *Propionibacterium acnes*). Chronic osteomyelitis develops when insufficient therapy leads to persistence or recurrence, most often after sternal, mandibular, or foot infection.

Classification by location distinguishes among cases in the long bones, the vertebral column, and the periarticular bones. Long bones are generally involved after hematogenous seeding in children or contiguous spread following trauma or surgery. The risk of vertebral osteomyelitis in adults increases with age. Periarticular osteomyelitis, which complicates septic arthritis that has not been adequately treated, is especially common in periprosthetic joint infection.

Osteomyelitis involving a foreign device requires surgical management for cure. Even acute implant-associated infection calls for prolonged antimicrobial therapy. Therefore, identification of this type of disease is of practical importance.

VERTEBRAL OSTEOMYELITIS

PATHOGENESIS

Vertebral osteomyelitis, also referred to as disk-space infection, septic diskitis, spondylodiskitis, or spinal osteomyelitis, is the most common manifestation of hematogenous bone infection in adults. This designation reflects a pathogenic process leading to involvement of the adjacent vertebrae and the corresponding intervertebral disk. In adults, the disk is avascular. Microorganisms invade via the segmental arterial circulation in adjacent endplates and then spread into the disk. Alternative routes of infection are retrograde seeding through the prevertebral venous plexus and direct inoculation during spinal surgery, epidural infiltration, or trauma. In the setting of implant surgery, microorganisms are inoculated either during the procedure or, if wound healing is impaired, in the early postoperative period.

EPIDEMIOLOGY

Vertebral osteomyelitis occurs more often in male than in female patients (ratio, 1.5:1). The overall incidence is 2.4 cases/100,000 population. There is a clear age-dependent increase from 0.3 cases/100,000 at ages <20 years to 6.5 cases/100,000 at ages >70 years. The observed increase in reported cases during the past two decades may reflect improvements in diagnosis resulting from the broad availability of MRI technology. In addition, the fraction of cases of vertebral osteomyelitis acquired in association with health care is certainly increasing as a consequence of the rising number of spinal interventions and local infiltrations.

MICROBIOLOGY

Vertebral osteomyelitis is typically classified as pyogenic or nonpyogenic. However, this distinction is arbitrary because, in “nonpyogenic” cases (tuberculous, brucellar), macroscopic pus formation (caseous necrosis, abscess) is quite common. A more accurate scheme is to

classify cases as acute or subacute/chronic. Whereas the microbiologic spectrum of acute cases is similar in different parts of the world, the spectrum of subacute/chronic cases varies according to the geographic region. The great majority of cases are monomicrobial in etiology. Of episodes of acute vertebral osteomyelitis, 40–50% are caused by *Staphylococcus aureus*, 12% by streptococci, and 20% by gram-negative bacilli—mainly *Escherichia coli* (9%) and *Pseudomonas aeruginosa* (6%). Subacute vertebral osteomyelitis is typically caused by *Mycobacterium tuberculosis* or *Brucella* species in regions where these microorganisms are endemic. Osteomyelitis due to viridans streptococci also has a subacute presentation; these infections most often occur as secondary foci in patients with endocarditis. In vertebral osteomyelitis due to *Candida* species, the diagnosis is often delayed by several weeks; this etiology should be suspected in IV drug users who do not use sterile paraphernalia. In implant-associated spinal osteomyelitis, coagulase-negative staphylococci and *P. acnes*—which, in the absence of an implant, are generally considered contaminants—typically cause low-grade (chronic) infections. As an exception, coagulase-negative staphylococci can cause native spinal osteomyelitis in cases of prolonged bacteremia (e.g., in patients with infected pacemaker electrodes or implanted vascular catheters that are not promptly removed).

CLINICAL MANIFESTATIONS

The signs and symptoms of vertebral osteomyelitis are nonspecific. Only about half of patients develop fever >38°C (100.4°F), perhaps because analgesic drugs are frequently used by these patients. Back pain is the leading initial symptom (>85% of cases). The location of the pain corresponds to the site of infection: the cervical spine in ~10% of cases, the thoracic spine in 30%, and the lumbar spine in 60%. One exception is involvement at the thoracic level in two-thirds of cases of tuberculous osteomyelitis and at the lumbar level in only one-third. This difference is due to direct mycobacterial spread via pleural or mediastinal lymph nodes in pulmonary tuberculosis.

Neurologic deficits, such as radiculopathy, weakness, or sensory loss, are observed in about one-third of cases of vertebral osteomyelitis. In brucellar vertebral osteomyelitis, neurologic impairment is less frequent; in tuberculous osteomyelitis, it is about twice as frequent as in cases of other etiologies. Neurologic signs and symptoms are caused mostly by spinal epidural abscess. This complication starts with severe localized back pain and progresses to radicular pain, reflex changes, sensory abnormalities, motor weakness, bowel and bladder dysfunction, and paralysis.

A primary focus should always be sought but is found in only half of cases. Overall, endocarditis is identified in ~10% of patients. In osteomyelitis caused by viridans streptococci, endocarditis is the source in about half of patients.

Implant-associated spinal osteomyelitis can present as either early- or late-onset infection. Early-onset infection is diagnosed within 30 days after implant placement. *S. aureus* is the most common pathogen. Wound healing impairment and fever are the leading findings. Late-onset infection is diagnosed beyond 30 days after surgery, with low-virulence organisms such as coagulase-negative staphylococci or *P. acnes* as typical infecting agents. Fever is rare. One-quarter of patients have a sinus tract. Because of the delayed course and the lack of classic signs of infection, rapid diagnosis requires a high degree of suspicion.

DIAGNOSIS

Leukocytosis and neutrophilia have low levels of diagnostic sensitivity (only 65% and 40%, respectively). In contrast, an increased erythrocyte sedimentation rate or C-reactive protein (CRP) level has been reported in 98% and 100% of cases, respectively; thus, these tests are helpful in excluding vertebral osteomyelitis. The fraction of blood cultures that yield positive results depends heavily on whether the patient has been pretreated with antibiotics; across studies, the range is from 30% to 78%. In view of this low rate of positive blood culture after antibiotic treatment, such therapy should be withheld until microbial growth is proven unless the patient has sepsis syndrome. In patients with negative blood cultures, CT-guided or open biopsy is needed. Whether a CT-guided biopsy with a negative result is repeated or followed by

840 open biopsy depends on the experience of personnel at the specific center. Bone samples should be cultured for aerobic, anaerobic, and fungal agents, with a portion of the sample sent for histopathologic study. In cases with a subacute/chronic presentation, a suggestive history, or a granuloma detected during histopathologic analysis, mycobacteria and brucellae also should be sought. When blood and tissue cultures are negative despite suggestive histopathology, broad-range polymerase chain reaction analysis of biopsy specimens or aspirated pus should be considered. This technique allows detection of unusual pathogens such as *Tropheryma whipplei*.

Given that signs and symptoms of osteomyelitis are nonspecific, the clinical differential diagnosis of febrile back pain is broad, including pyelonephritis, pancreatitis, and viral syndromes. In addition, multiple noninfectious pathologies of the vertebral column, such as osteoporotic fracture, seronegative spondylitis (ankylosing spondylitis, psoriasis, reactive arthritis, enteropathic arthritis), and spinal stenosis must be considered.

Imaging procedures are the most important tools not only for the diagnosis of vertebral osteomyelitis but also for the detection of pyogenic complications and alternative conditions (e.g., bone metastases or osteoporotic fractures). Plain radiography is a reasonable first step in evaluating patients without neurologic symptoms and may reveal an alternative diagnosis. Because of its low sensitivity, plain radiography generally is not helpful in acute osteomyelitis, but it can be useful in subacute or chronic cases. The gold standard is MRI, which should be performed expeditiously in patients with neurologic impairment in order to rule out a herniated disk or to detect pyogenic complications in a timely manner. Even if the pathologic findings on MRI suggest vertebral osteomyelitis, alternative diagnoses should be considered, especially when blood cultures are negative. The most common alternative diagnosis is erosive osteochondrosis. Septic bone necrosis, gouty spondylodiskitis, and erosive diskovertebral lesions (Andersson lesions) in ankylosing spondylitis may likewise mimic vertebral osteomyelitis. CT is less sensitive than MRI but may be helpful in guiding a percutaneous biopsy. In the future, positron-emission tomography (PET) with ¹⁸F-fluorodeoxyglucose, which has a high degree of diagnostic accuracy, may be an alternative imaging procedure when MRI is contraindicated. ¹⁸F-fluorodeoxyglucose PET should be considered for patients with implants and patients in whom several foci are suspected.

TREATMENT VERTEBRAL OSTEOMYELITIS

The aims of therapy for vertebral osteomyelitis are (1) elimination of the pathogen(s), (2) protection from further bone loss, (3) relief of back pain, (4) prevention of complications, and (5) stabilization, if needed.

ANTIMICROBIAL THERAPY

Table 158-1 summarizes suggested antimicrobial regimens for infections attributable to the most common etiologic agents. For optimal antimicrobial therapy, identification of the infecting agent is required. Therefore, in patients without sepsis syndrome, antibiotics should not be administered until the pathogen is identified in a blood culture, a bone biopsy, or an aspirated pus collection. Traditionally, bone infections are at least initially treated by the IV route. Unfortunately, relevant controlled trials are lacking, and the preference for the IV route is not evidence based. There are no good arguments for the assumption that IV therapy is superior to oral administration if the following requirements are met: (1) optimal antibiotic spectrum, (2) excellent bioavailability of the oral drug, (3) clinical studies confirming efficacy of the oral drug, (4) normal intestinal function, and (5) no vomiting. However, a short initial course of parenteral therapy with a β -lactam antibiotic may lower the risk of emergence of fluoroquinolone resistance, especially if *P. aeruginosa* infection is treated with ciprofloxacin or staphylococcal infection with the combination of a fluoroquinolone plus rifampin. These suggestions are based on observational studies and expert opinion. There are no data from controlled trials on the optimal duration of therapy. Most experts suggest 6 weeks for patients

TABLE 158-1 ANTIBIOTIC THERAPY FOR OSTEOMYELITIS IN ADULTS WITHOUT IMPLANTS^a

Microorganism	Antimicrobial Agent (Dose, ^b Route)
<i>Staphylococcus</i> spp.	
Methicillin-susceptible	Nafcillin or oxacillin ^c (2 g IV q6h) followed by Rifampin (300–450 mg PO q12h) <i>plus</i> levofloxacin (750 mg PO q24h or 500 mg PO q12h)
Methicillin-resistant	Vancomycin ^d (15 mg/kg IV q12h) or daptomycin (>6–8 mg/kg IV q24h) followed by Rifampin (300–450 mg PO q12h) plus Levofloxacin (750 mg PO q24h or 500 mg PO q12h) or TMP-SMX ^e (1 double-strength tablet PO q8h) or fusidic acid (500 mg PO q8h)
<i>Streptococcus</i> spp.	Penicillin G ^f (5 million units IV q6h) or ceftriaxone (2 g IV q24h)
Enterobacteriaceae	
Quinolone-susceptible	Ciprofloxacin (750 mg PO q24h)
Quinolone-resistant ^g	Imipenem (500 mg IV q6h)
<i>Pseudomonas aeruginosa</i>	Cefepime or ceftazidime (2 g IV q8h) <i>plus</i> an aminoglycoside ^h or Piperacillin-tazobactam (4.5 g IV q8h) <i>plus</i> an aminoglycoside ^h for 2–4 weeks followed by Ciprofloxacin ^h (750 mg PO q12h)
Anaerobes	Clindamycin (600 mg IV q6–8h) for 2–4 weeks followed by Clindamycin ⁱ (300 mg PO q6h)

^aUnless otherwise indicated, the total duration of antimicrobial treatment is generally 6 weeks. ^bAll dosages are for adults with normal renal function. ^cWhen the patient has delayed-type penicillin hypersensitivity, cefuroxime (1.5 g IV q6–8h) can be administered. When the patient has immediate-type penicillin hypersensitivity, the penicillin should be replaced by vancomycin (1 g IV q12h). ^dTarget vancomycin trough level: 15–20 μ g/mL. ^eTrimethoprim-sulfamethoxazole. A double-strength tablet contains 160 mg of trimethoprim and 800 mg of sulfamethoxazole. ^fIncluding isolates producing extended-spectrum β -lactamase. ^gThe need for addition of an aminoglycoside has not yet been proven. However, this addition may decrease the risk of emergence of resistance to the β -lactam. ^hThe rationale for starting ciprofloxacin treatment only after pretreatment with a β -lactam is the increased risk of emergence of quinolone resistance in the presence of a heavy bacterial load. ⁱAlternatively, penicillin G (5 million units IV q6h) or ceftriaxone (2 g IV q24h) can be used against gram-positive anaerobes (e.g., *Propionibacterium acnes*), and metronidazole (500 mg IV/PO q8h) can be used against gram-negative anaerobes (e.g., *Bacteroides* spp.).

Source: From W Zimmerli: N Engl J Med 362:1022, 2010. © Massachusetts Medical Society. Reprinted with permission.

who have acute osteomyelitis without an implant. According to an observational study, prolonging antibiotic therapy beyond 6 weeks does not improve the rate of recovery or lower the risk of recurrence. However, prolonged antibiotic therapy is recommended for patients with abscesses that have not been drained and patients with spinal implants. Treatment efficacy should be regularly monitored through inquiries about signs and symptoms (fever, pain) and assessment for signs of inflammation (elevated CRP concentrations). Follow-up MRI is appropriate only for patients with pyogenic complications, since the correlation between clinical healing and improvement on MRI is very poor.

Surgical treatment generally is not needed in acute hematogenous vertebral osteomyelitis. However, it is always necessary in implant-associated spinal infection. Early infections (those occurring up to 30 days after internal stabilization) can be cured with debridement, implant retention, and a 3-month course of antibiotics (**Table 158-2**). In contrast, in late infection with a duration of >30 days, implant removal and a 6-week-course of antibiotics (**Table 158-1**)

TABLE 158-2 ANTIBIOTIC THERAPY FOR OSTEOMYELITIS ASSOCIATED WITH ORTHOPEDIC DEVICES

Microorganism	Antimicrobial Agent ^a (Dose, Route)
<i>Staphylococcus</i> spp.	Recommendation for initial treatment phase (2 weeks with implant)
Methicillin-susceptible	Rifampin (450 mg PO/IV q12h ^b) plus Nafcillin or oxacillin ^c (2 g IV q6h)
Methicillin-resistant	Rifampin (450 mg PO/IV q12h ^b) plus Vancomycin (15 mg/kg IV q12h) or daptomycin (6–8 mg/kg IV q24h)
<i>Staphylococcus</i> spp.	Recommendation after completion of initial treatment phase
	Rifampin (450 mg PO q12h ^b) plus Levofloxacin (750 mg PO q24h or 500 mg PO q12h or ciprofloxacin (750 mg PO q12h) or fusidic acid (500 mg PO q8h) or TMP-SMX ^d (1 double-strength tablet PO q8h) or minocycline (100 mg PO q12h) or linezolid (600 mg PO q12h) or clindamycin (1200–1350 mg/d PO in 3 or 4 divided doses)
<i>Streptococcus</i> spp. ^e	Penicillin G ^c (18–24 million units/d IV in 6 divided doses) or ceftriaxone (2 g IV q24h) for 4 weeks followed by Amoxicillin (750–1000 mg PO q6–8h) or clindamycin (1200–1350 mg/d PO in 3 or 4 divided doses)
<i>Enterococcus</i> spp. ^f	
Penicillin-susceptible	Penicillin G ^c (24 million units/d IV in 6 divided doses) or ampicillin or amoxicillin ^g (2 g IV q4–6h)
Penicillin-resistant	Vancomycin (15 mg/kg IV q12h) or daptomycin (6–8 mg/kg IV q24h) or linezolid (600 mg IV/PO q12h)
Enterobacteriaceae	A β -lactam selected in light of in vitro susceptibility profile for 2 weeks ^h followed by Ciprofloxacin (750 mg PO q12h)
<i>Enterobacter</i> spp. ⁱ and nonfermenters ^j (e.g., <i>Pseudomonas aeruginosa</i>)	Cefepime or ceftazidime (2 g IV q8h) or meropenem (1 g IV q8h ^k) for 2–4 weeks followed by Ciprofloxacin (750 mg PO q12h)
<i>Propionibacterium</i> spp.	Penicillin G ^c (18–24 million units/d IV in 6 divided doses) or clindamycin (600–900 mg IV q8h) for 2–4 weeks followed by Amoxicillin (750–1000 mg PO q6–8h) or clindamycin (1200–1350 mg/d PO in 3 or 4 divided doses)
Gram-negative anaerobes (e.g., <i>Bacteroides</i> spp.)	Metronidazole (500 mg IV/PO q8h)
Mixed bacteria (without methicillin-resistant staphylococci)	Ampicillin-sulbactam (3 g IV q6h) or amoxicillin-clavulanate ^l (2.2 g IV q6h) or piperacillin-tazobactam (4.5 g IV q8h) or imipenem (500 mg IV q6h) or meropenem (1 g IV q8h ^k) for 2–4 weeks followed by Individualized oral regimens chosen in light of antimicrobial susceptibility

^aAntimicrobial agents should be chosen in light of the isolate's in vitro susceptibility, the patient's drug allergies and intolerances, potential drug interactions, and contraindications to specific drugs. All dosages recommended are for adults with normal renal and hepatic function. See text for total durations of antibiotic treatment. ^bOther dosages and intervals of administration with equivalent success rates have been reported. ^cWhen the patient has delayed-type penicillin hypersensitivity, cefazolin (2 g IV q8h) can be administered. When the patient has immediate-type penicillin hypersensitivity, the penicillin should be replaced by vancomycin (1 g IV q12h). ^dTrimethoprim-sulfamethoxazole. A double-strength tablet contains 160 mg of trimethoprim and 800 mg of sulfamethoxazole. ^eDetermination of the minimal inhibitory concentration (MIC) of penicillin is advisable. ^fCombination therapy with an aminoglycoside is optional since its superiority to monotherapy for prosthetic joint infection is unproved. When using combination therapy, monitor signs of aminoglycoside ototoxicity and nephrotoxicity; the latter is potentiated by other nephrotoxic agents (e.g., vancomycin). ^gFor patients with hypersensitivity to penicillin, see treatment options for penicillin-resistant enterococci. ^hCiprofloxacin (PO or IV) can be administered to patients with hypersensitivity to β -lactams. ⁱCeftriaxone and ceftazidime should not be administered for treatment targeting *Enterobacter* species, even strains that test susceptible in the laboratory, but can be used against nonfermenters. Strains producing extended-spectrum β -lactamases should not be treated with any cephalosporin, including cefepime. *Enterobacter* infections can also be treated with ertapenem (1 g IV q24h); however, ertapenem is not effective against *Pseudomonas* spp. and other nonfermenters. ^jAddition of an aminoglycoside is optional. Use of two active drugs can be considered in light of the patient's clinical condition. ^kThe recommended dosage is in line with the guidelines of the Infectious Diseases Society of America. In Europe, 2 g IV q8h is suggested for *P. aeruginosa* infections. ^lNot available as an IV formulation in the United States.

Source: Modified from W Zimmerli et al: *N Engl J Med* 351:1645, 2004. © Massachusetts Medical Society. Reprinted with permission.

are required for complete elimination of the infection. If implants cannot be removed, oral suppressive long-term treatment should follow the initial course of IV antibiotics. The optimal duration of suppressive therapy is unknown. However, if antibiotic therapy is discontinued after, for example, 1 year, close clinical and laboratory (CRP) follow-up is needed.

COMPLICATIONS

Complications include persistent pain, persistently increased CRP levels, and new-onset or persistent neurologic impairment. In cases of persistent pain with or without signs of inflammation, paravertebral, epidural, or psoas abscesses (Fig. 158-1) must be sought. Epidural abscesses occur in 15–20% of cases. This complication is more common in the cervical column (30%) than in the lumbar spine

(12%). Persistent pain despite normalization of CRP values indicates mechanical complications such as severe osteonecrosis or spinal instability. These patients require a consult with an experienced orthopedic surgeon.

OSTEOMYELITIS IN LONG BONES

PATHOGENESIS

Osteomyelitis in long bones is a consequence of hematogenous seeding, exogenous contamination during trauma (open fracture), or perioperative contamination during orthopedic repairs. Its presentation is either acute (with a duration of days to a few weeks) or chronic. Hematogenous infection in long bones typically occurs in children. Ineffectively treated hematogenous osteomyelitis during childhood can progress to chronic disease. In adults, the leading pathogenic source is exogenous infection, mainly associated with internal fixation



FIGURE 158-1 CT scan of acute vertebral osteomyelitis (L1/L2) due to *Staphylococcus aureus* in a 64-year-old man. Low-grade fever persisted despite appropriate IV antibiotic therapy. The scan revealed a psoas abscess on the right side.



FIGURE 158-2 A 42-year-old man who had had a malleolar fracture 6 weeks previously had persistent pain and slight inflammation after orthopedic repair. His infection was treated with oral antibiotics without debridement surgery. This insufficient management of an implant-associated *Staphylococcus aureus* infection was complicated by a sinus tract.

Patients may present with persisting pain, subtle local signs of inflammation, intermittent discharge of pus, or fluctuating erythema over the scar (Fig. 158-2).

DIAGNOSIS

The diagnostic workup for acute hematogenous long-bone osteomyelitis is similar to that for vertebral osteomyelitis. Bone remodeling and thus marker uptake are increased for at least 1 year after surgery. Therefore, the three-phase bone scan is not useful during this interval. However, in late recurrences it allows rapid diagnosis at low cost. If the results are positive, CT is required in order to estimate the extent of inflamed tissue and to detect bone necrosis (sequestrs). Implant-associated infection should be suspected if CRP values do not return to the normal range or rise after an initial decrease. Clinical and laboratory suspicion should prompt surgical exploration and sampling.

In chronic osteomyelitis of >1 year's duration, single-photon emission CT plus conventional CT (SPECT/CT) is a good option, either with ^{99m}Tc methylene diphosphonate (^{99m}Tc -MDP)-labeled leukocytes or with labeled monoclonal antibodies to granulocytes. Surgical debridement is needed for diagnostic (biopsy culture, histology) and therapeutic reasons.

TREATMENT OSTEOMYELITIS IN LONG BONES

Treatment for acute hematogenous infection in long bones is identical to that for acute vertebral osteomyelitis (Table 158-1). The suggested duration of antibiotic therapy is 4–6 weeks. In contrast to chronic or implant-associated osteomyelitis, acute hematogenous infection does not require surgical intervention. Initial IV administration of antimicrobial agents is followed by long-term oral treatment. The duration of the initial IV phase of therapy has not been defined. The IV course can be as short as a couple of days if a drug with excellent bioavailability is available. In case of recurrence of chronic osteomyelitis as well as in each type of exogenous osteomyelitis (acute, chronic, with or without an implant), a combination of surgical debridement, obliteration of dead space, and long-term antibiotic therapy is needed.

The therapeutic aims in patients whose infections are associated with internal fixation devices are consolidation of the fracture and prevention of chronic osteomyelitis. Stable implants can be maintained except in patients with uncontrolled sepsis. Appropriate antimicrobial therapies are listed in Table 158-2. The cure rate for

devices. Chronic osteomyelitis can recur after a symptom-free interval of >50 years. Such recurrences are most common among elderly patients who developed osteomyelitis in the preantibiotic era.

EPIDEMIOLOGY

In adults, most cases of long-bone osteomyelitis are posttraumatic or postsurgical; less frequently, late recurrence arises from hematogenous infections during childhood. The risk of infection depends on the type of fracture. After closed fracture, implant-associated infection occurs in fewer than 1% of patients. In contrast, after open fracture, the risk of osteomyelitis ranges from ~2% up to 16%, with the precise figure depending on the degree of tissue damage during trauma.

MICROBIOLOGY

The spectrum of microorganisms causing hematogenous long-bone osteomyelitis does not differ from that in vertebral osteomyelitis. *S. aureus* is most commonly isolated from adult patients. In rare cases, mycobacteria or fungal agents such as *Cryptococcus* species, *Sporothrix schenckii*, *Blastomyces dermatitidis*, or *Coccidioides* species are found in patients who live or have traveled in endemic regions. Impaired cellular immunity (e.g., in HIV infection or after transplantation) predisposes to these etiologies. Coagulase-negative staphylococci are the second most common etiologic agents (after *S. aureus*) in implant-associated osteomyelitis. After open fracture, contiguous long-bone osteomyelitis is typically caused by gram-negative bacilli or a polymicrobial mixture of organisms.

CLINICAL MANIFESTATIONS

The leading symptoms in adults with primary or recurrent hematogenous long-bone osteomyelitis are pain and low-grade fever. Infection occasionally manifests as clinical sepsis and local signs of inflammation (erythema and swelling). After internal fixation, osteomyelitis can be classified as acute (≤ 3 weeks) or chronic. Acute long-bone osteomyelitis manifests as signs of surgical site infection, such as erythema and impaired wound healing. Acute implant-associated infection may also follow hematogenous seeding at any time after implantation of a device. Typical symptoms are new-onset pain and signs of sepsis. Chronic infections are usually caused by low-virulence microorganisms or occur after ineffective treatment of early-onset infection.

early staphylococcal implant-associated infections treated with a fluoroquinolone plus rifampin is >90%. Rifampin is efficacious against staphylococcal biofilms of ≤ 3 weeks' duration. Similarly, fluoroquinolones are active against biofilms formed by gram-negative bacilli. In these cases, an initial 2-week course of IV therapy with a β -lactam is suggested in order to minimize the risk of emergence of resistance to the oral drugs. The total duration of treatment is 3 months, and the device can be retained even after antibiotics have been discontinued. In contrast, in cases caused by rifampin-resistant staphylococci or fluoroquinolone-resistant gram-negative bacilli, the hardware should be removed after consolidation of the fracture and before discontinuation of antibiotics. These patients are treated with an oral antibiotic (suppressive therapy) as long as retention of the hardware is necessary.

COMPLICATIONS

The main complication of long-bone osteomyelitis is the persistence of infection with progression to chronic osteomyelitis. This risk is especially high after internal fixation of an open fracture and among patients with implant-associated osteomyelitis that is treated without surgical debridement. In chronic osteomyelitis, recurrent sinus tracts result in severe damage to skin and soft tissue (Fig. 158-2). Patients who have chronic open wounds need a therapeutic approach combining orthopedic repair and plastic reconstructive surgery.

PERIPROSTHETIC JOINT INFECTION

PATHOGENESIS

Implanted foreign material is highly susceptible to local infection due to local immunodeficiency around the device. Infection occurs by either the exogenous or the hematogenous route. More rarely, contiguous spread from adjacent sites of osteomyelitis or deep soft-tissue infection may cause periprosthetic joint infection (PJI). The fact that foreign devices are covered with host proteins such as fibronectin favors the adherence of staphylococci and the formation of a biofilm that resists phagocytosis.

EPIDEMIOLOGY

The risk of infection manifesting during the first 2 postoperative years varies according to the joint. It is lowest after hip and knee arthroplasty (0.3–1.5%) and highest after ankle and elbow replacement (4–10%). The risk of hematogenous PJI is highest in the early postoperative period. However, hematogenous seeding occurs throughout life, and most cases therefore develop >2 years after implantation.

MICROBIOLOGY

About 70% of cases of PJI are caused by staphylococci (*S. aureus* and coagulase-negative staphylococci), 10% by streptococci, 10% by gram-negative bacilli, and the rest by various other microorganisms. All microorganisms can cause PJI, including fungi and mycobacteria. *P. acnes* causes up to one-third of episodes of periprosthetic shoulder infection.

CLASSIFICATION AND CLINICAL MANIFESTATIONS

PJI is traditionally classified as early (<3 months after implantation), delayed (3–24 months after surgery), or late (>2 years after implantation). For therapeutic decision-making (see below), it is more useful to classify PJI as (1) acute hematogenous PJI with <3 weeks of symptoms, (2) early postinterventional PJI manifesting within 1 month after surgery, and (3) chronic PJI with symptom duration of >3 weeks.

Acute exogenous PJI typically presents with local signs of infection (Fig. 158-3). In contrast, acute hematogenous PJI, most often caused by *S. aureus*, is characterized by new-onset pain that initially is not accompanied by prominent local inflammatory signs. In most cases, an ongoing sepsis syndrome dominates the clinical picture. Key findings in chronic PJI are joint effusion, local pain, implant loosening, and occasionally a sinus tract. Chronic PJI is most commonly caused



FIGURE 158-3 Early periprosthetic joint infection of the left hip caused by group B streptococci in a 68-year-old woman.

by low-virulence microorganisms such as coagulase-negative staphylococci or *P. acnes*. These infections are characterized by nonspecific symptoms, such as chronic pain caused by low-grade inflammation or early loosening.

DIAGNOSIS

Blood tests such as the measurement of CRP (elevated levels, ≥ 10 mg/L) and erythrocyte sedimentation rate (elevated rates, ≥ 30 mm/h) are sensitive (91–97%) but not specific (70–78%). Synovial fluid cell counts are ~90% sensitive and specific, with threshold values of 1700 leukocytes/ μ L in periprosthetic knee infection and 4200 leukocytes/ μ L in periprosthetic hip infection. During debridement surgery, at least three but optimally six tissue samples should be obtained for culture and histopathology. If implant material (modular parts, screws, or the prosthesis) is removed, sonication of this material followed by culture and/or use of molecular methods to examine the sonicate fluid allows the detection of microorganisms in biofilms.

The three-phase bone scan is very sensitive for detecting PJI but is not specific. As mentioned above, this test does not differentiate bone remodeling from infection and therefore is not useful during at least the first year after implantation. CT and MRI detect soft tissue infection, prosthetic loosening, and bone erosion, but imaging artifacts caused by metal implants limit their use. 18 F-fluorodeoxyglucose PET is an alternative method with fair sensitivity and specificity for the detection of PJI. However, this technique is not yet an established procedure for this purpose.

TREATMENT PERIPROSTHETIC JOINT INFECTION

Treatment of PJI requires a multidisciplinary approach involving an experienced orthopedic surgeon, an infectious disease specialist, a plastic reconstructive surgeon, and a microbiologist. Therefore, most patients are referred to a specialized center. In general, the goal of treatment is cure—i.e., a pain-free functional joint with complete eradication of the infecting pathogen(s). However, for patients with severe comorbidity, lifelong suppressive antimicrobial therapy may be preferred. As a rule, antimicrobial therapy without surgical intervention is not curative but merely suppressive. There are four curative surgical options: debridement and implant retention, one-stage implant exchange, two-stage implant exchange, and implant removal without replacement. Implant retention offers a good chance of infection-free survival (>80%) only if the following conditions are fulfilled: (1) acute infection, (2) stable implant, (3) pathogen susceptible to a biofilm-active antimicrobial agent (see below), and (4) skin and soft tissue in good condition.

Table 158-2 summarizes pathogen-specific antimicrobial therapy for PJI. Initial IV therapy is followed by long-term oral antibiotics. Efficacious treatment is best defined in staphylococcal implant-associated infections. Rifampin exhibits excellent activity against

844 biofilms composed of susceptible staphylococci. Because of the risk of rapid emergence of resistance, rifampin must always be combined with another effective antibiotic. If gram-negative infections are treated with implant retention, fluoroquinolones should be used because of their activity against gram-negative biofilms.

PREVENTION OF HEMATOGENOUS INFECTION

As mentioned above, hematogenous seeding may occur throughout life. This risk is highest during *S. aureus* bacteremia from a distant focus. Therefore, documented bacterial infections should be promptly treated in patients with prosthetic joints. However, according to a large, prospective, case-control study, the risk of prosthetic hip or knee infection is not increased following dental procedures. Therefore, antibiotic prophylaxis is not needed during dental work.

STERNAL OSTEOMYELITIS

PATHOGENESIS

Sternal osteomyelitis occurs primarily after sternal surgery (with the entry of exogenous organisms) and more rarely by hematogenous seeding or contiguous extension from adjacent sites of sternocostal arthritis. Exogenous sternal osteomyelitis after open sternal surgery is also called *deep sternal wound infection*. Exogenous infection may also follow minor sternal trauma, sternal fracture, and manubriosternal septic arthritis. Tuberculous sternal osteomyelitis typically manifests during hematogenous seeding in children or as reactivated infection in adults. Reactivation is sometimes preceded by blunt trauma. In rare cases, tuberculous sternal osteomyelitis is caused by continuous infection from an infected internal mammary lymph node.

EPIDEMIOLOGY

The incidence of poststernotomy wound infection varies from 0.5% to 5%, but figures are even higher among patients with risk factors such as diabetes, obesity, chronic renal failure, emergency surgery, use of bilateral internal mammary arteries, and reexploration for bleeding. Rapid diagnosis and correct management of superficial sternal wound infection prevent its progression to sternal osteomyelitis. Primary (hematogenous) sternal osteomyelitis accounts for only 0.3% of all cases of osteomyelitis. Risk factors are IV drug use, HIV infection, radiotherapy, blunt trauma, cardiopulmonary resuscitation, alcohol abuse, liver cirrhosis, and hemoglobinopathy.

MICROBIOLOGY

Poststernotomy osteomyelitis is generally caused by *S. aureus* (40–50% of cases), coagulase-negative staphylococci (15–30%), enterococci (5–12%), or gram-negative bacilli (15–25%). Fungal infections caused by *Candida* species also play a role. The fact that ~20% of cases are polymicrobial is indicative of exogenous superinfection during therapy. Hematogenous sternal osteomyelitis is caused most commonly by *S. aureus*. Other microorganisms play a role in special populations—e.g., *P. aeruginosa* in IV drug users, *Salmonella* species in individuals with sickle cell anemia, and *M. tuberculosis* in patients from endemic areas who have previously had tuberculosis.

CLINICAL MANIFESTATIONS

Exogenous sternal osteomyelitis manifests as fever, increased local pain, erythema, wound discharge, and sternal instability (Fig. 158-4). Contiguous mediastinitis is a feared complication, occurring in ~10–30% of patients with sternal osteomyelitis. Hematogenous sternal osteomyelitis is characterized by sternal pain, swelling, and erythema. In addition, most patients have systemic signs and symptoms of sepsis.

The differential diagnosis of hematogenous sternal osteomyelitis includes immunologic processes typically presenting as systemic or multifocal inflammation of the sternum or the sternoclavicular or sternocostal joints (e.g., SAPHO [synovitis, acne, pustulosis, hyperostosis, osteitis], vasculitis, and chronic multifocal relapsing osteomyelitis).



FIGURE 158-4 Sternal osteomyelitis caused by *Staphylococcus epidermidis* 5 weeks after sternotomy for aortocoronary bypass in a 72-year-old man.

DIAGNOSIS

In primary sternal osteomyelitis, the diagnostic workup does not differ from that in other types of hematogenous osteomyelitis (see above). When a patient has grown up in regions where tuberculosis is endemic, a specific workup for mycobacterial infection should be performed, especially if osteomyelitis had its onset after a blunt sternal trauma. In secondary sternal osteomyelitis, leukocyte counts may be normal, but the CRP level is >100 mg/L in most cases. Tissue sampling for microbiologic studies is crucial. In osteomyelitis associated with sternal wires, low-virulence microorganisms, such as coagulase-negative staphylococci, play an important role. In order to differentiate between colonization and infection, samples from at least three deep biopsies should be subjected to microbiologic examination. Superficial swab cultures are not diagnostic and may be misleading. No studies have compared the value of the various imaging modalities in suspected primary sternal osteomyelitis. However, MRI is the current gold standard for detection of each type of osteomyelitis.

TREATMENT STERNAL OSTEOMYELITIS

In cases of deep sternal wound infection, antibiotic therapy should be started immediately after samples have been obtained for microbiologic analyses in order to control clinical sepsis. To protect a newly inserted heart valve, initial treatment should be directed against staphylococci, with consideration of the local susceptibility pattern. In centers with a high prevalence of methicillin-resistant *S. aureus*, vancomycin or daptomycin should be added to a broad-spectrum β -lactam drug. As soon as cultures of blood and/or deep wound biopsies have confirmed the pathogen's identity and susceptibility pattern, treatment should be optimized and narrowed accordingly. Tables 158-1 and 158-2 show appropriate therapeutic choices for the most frequently identified microorganisms causing sternal osteomyelitis in the absence and presence, respectively, of an implanted device. In a recent observational study of patients with staphylococcal deep sternal wound infection, the use of a rifampin-containing regimen was predictive of success. The optimal duration of antibiotic therapy has not been established. In acute sternal osteomyelitis without hardware, a 6-week course is the rule. In patients with remaining sternal wires, treatment duration is generally prolonged to 3 months (Table 158-2). Like other types of tuberculous bone infection, tuberculous sternal osteomyelitis is treated for 6–12 months.

Primary sternal osteomyelitis can generally be treated without surgery. In contrast, in secondary sternal osteomyelitis, debridement is always required. This procedure should be performed by a team of experienced surgeons, since mediastinitis, bone infection,

and skin and soft tissue damage may need to be treated during the same intervention.

PROGNOSIS

Primary sternal osteomyelitis poses a minimal mortality risk. In contrast, the in-hospital mortality rates from secondary sternal osteomyelitis are 15–30% after sternal surgery.

FOOT OSTEOMYELITIS

PATHOGENESIS

Osteomyelitis of the foot usually occurs in patients with diabetes, peripheral arterial insufficiency, or peripheral neuropathy and after foot surgery. These entities are often linked to each other, especially in diabetic patients with late complications. However, foot osteomyelitis is also seen in patients with isolated peripheral neuropathy and can manifest as implant-associated osteomyelitis in patients without comorbidity due to a deep wound infection after foot surgery (hallux valgus surgery, arthrodesis, total ankle arthroplasty). Foot osteomyelitis is acquired almost exclusively by the exogenous route. It is a complication of deep pressure ulcers and of impaired wound healing after surgery.

EPIDEMIOLOGY

The incidence of diabetic foot infection is 30–40 cases/1000 persons with diabetes per year. The condition starts with skin and soft tissue lesions and progresses to osteomyelitis, especially in patients with risk factors. About 60–80% of patients with diabetic foot infection have confirmed osteomyelitis. Diabetic foot osteomyelitis increases the risk of amputation. With adequate management of the early stage of diabetic foot infections, the rate of amputation can be lowered.

RISK FACTORS

Risk factors for diabetic foot infection are (1) peripheral motor, sensory, and autonomic neuropathy; (2) neuro-osteoarthropathic deformities (Charcot foot; Fig. 158-5); (3) arterial insufficiency; (4) uncontrolled hyperglycemia; (5) disabilities such as reduced vision; and (6) maladaptive behavior.

MICROBIOLOGY

The correlation between cultures from bone biopsy and those from wound swabs or even deep soft tissue punctures is poor. Consistent

results have been found in only 13–43% of cases in various studies. The correlation is better when *S. aureus* is isolated (40–50%) than when anaerobes (20–35%), gram-negative bacilli (20–30%), or coagulase-negative staphylococci (0–20%) are identified. When only bone biopsy samples are considered, the leading pathogens are *S. aureus* (30–40%), anaerobes (10–20%), and various gram-negative bacilli (30–40%). The precise distribution depends on whether the patient already has been treated with antibiotics. Anaerobes are especially prevalent in chronic wounds. Pretreatment typically selects for *P. aeruginosa* or enterococci.

DIAGNOSIS

In many cases, foot osteomyelitis can be diagnosed clinically, without imaging procedures. Most clinicians rely on the “probe-to-bone” test, which has a positive predictive value of ~90% in populations with a high pretest probability. Thus, in a patient with diabetes who is hospitalized for a chronic deep foot ulcer, the diagnosis of foot osteomyelitis is highly probable if bone can be directly touched with a metal instrument. In a patient with a lower pretest probability, MRI should be performed because of its high degree of sensitivity (80–100%) and specificity (80–90%). Plain radiography has a sensitivity of only 30–90% and a specificity of only 50–90%; it may be considered for follow-up of patients with confirmed diabetic foot osteomyelitis.

TREATMENT FOOT OSTEOMYELITIS

As mentioned above, correlation between cultures of bone and those of wound swabs or wound punctures is poor. Antibiotic treatment should be based on bone culture. If no bone biopsy is performed, empirical therapy chosen in light of the most common infecting agents and the type of clinical syndrome should be given. Wound debridement combined with a 4- to 6-week course of antibiotics has been shown to render amputation unnecessary in about two-thirds of patients. According to the 2012 Infectious Diseases Society of America Clinical Practice Guideline for the Diagnosis and Treatment of Diabetic Foot Infections, the following management strategies should be considered. If a foot ulcer is clinically infected, prompt empirical antimicrobial therapy may prevent progression to osteomyelitis. When the risk of methicillin-resistant *S. aureus* is considered high, an agent active against these strains (e.g., vancomycin) should be chosen. If the patient has not recently received antibiotics, the spectrum of the selected antibiotic must include gram-positive cocci (e.g., clindamycin, ampicillin-sulbactam). If the patient has received antibiotics within the past month, the spectrum of empirical antibiotics should include gram-negative bacilli (e.g., clindamycin plus a fluoroquinolone). If the patient has risk factors for *Pseudomonas* infection (previous colonization, residence in a warm climate, frequent exposure of the foot to water), an empirical anti-pseudomonal agent (e.g., piperacillin-tazobactam, cefepime) is indicated. If osteomyelitis is suspected either on clinical grounds (probe to bone) or on the basis of imaging procedures (MRI), bone biopsy should be performed. If not all infected bone is surgically removed, the patient should be treated for 4–6 weeks in line with the identified pathogen(s) and their susceptibility. Treatment should initially be given by the IV route. Whether therapy can later be administered by the oral route depends on the bioavailability of oral drugs that cover the infecting agents. If dead bone cannot be removed, long-term therapy (at least 3 months) should be considered. In such cases, cure of osteomyelitis is usually the exception, and repetitive suppressive treatment may be needed.



FIGURE 158-5 Neuropathic joint disease (Charcot foot) complicated by chronic foot osteomyelitis in a 78-year old woman with diabetes mellitus complicated by severe neuropathy.

159 Intraabdominal Infections and Abscesses

Miriam Baron Barshak, Dennis L. Kasper

Intraperitoneal infections generally arise because a normal anatomic barrier is disrupted. This disruption may occur when the appendix, a diverticulum, or an ulcer ruptures; when the bowel wall is weakened by ischemia, tumor, or inflammation (e.g., in inflammatory bowel disease); or with adjacent inflammatory processes, such as pancreatitis or pelvic inflammatory disease, in which enzymes (in the former case) or organisms (in the latter) may leak into the peritoneal cavity. Whatever the inciting event, once inflammation develops and organisms usually contained within the bowel or another organ enter the normally sterile peritoneal space, a predictable series of events takes place. Intraabdominal infections occur in two stages: peritonitis and—if the patient survives this stage and goes untreated—abscess formation. The types of microorganisms predominating in each stage of infection are responsible for the pathogenesis of disease.

PERITONITIS

Peritonitis is a life-threatening event that is often accompanied by bacteremia and sepsis syndrome (Chap. 325). The peritoneal cavity is large but is divided into compartments. The upper and lower peritoneal cavities are divided by the transverse mesocolon; the greater omentum extends from the transverse mesocolon and from the lower pole of the stomach to line the lower peritoneal cavity. The pancreas, duodenum, and ascending and descending colon are located in the anterior retroperitoneal space; the kidneys, ureters, and adrenals are found in the posterior retroperitoneal space. The other organs, including liver, stomach, gallbladder, spleen, jejunum, ileum, transverse and sigmoid colon, cecum, and appendix, are within the peritoneal cavity. The cavity is lined with a serous membrane that can serve as a conduit for fluids—a property exploited in peritoneal dialysis (Fig. 159-1).

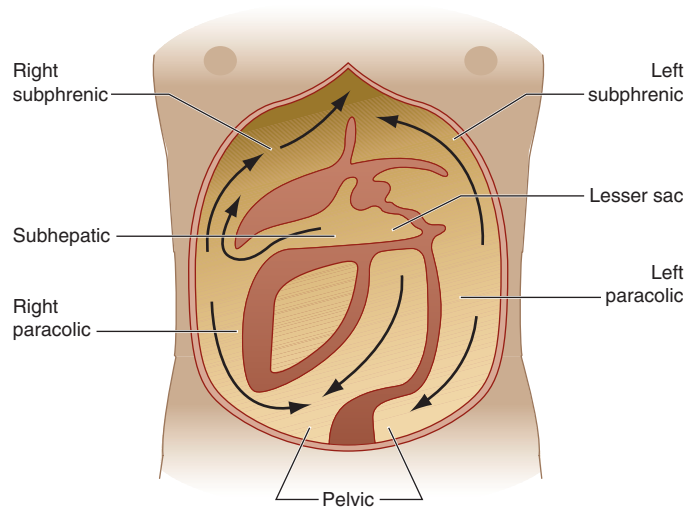


FIGURE 159-1 Diagram of the intraperitoneal spaces, showing the circulation of fluid and potential areas for abscess formation. Some compartments collect fluid or pus more often than others. These compartments include the pelvis (the lowest portion), the subphrenic spaces on the right and left sides, and Morrison's pouch, which is a posterosuperior extension of the subhepatic spaces and is the lowest part of the paravertebral groove when a patient is recumbent. The falciform ligament separating the right and left subphrenic spaces appears to act as a barrier to the spread of infection; consequently, it is unusual to find bilateral subphrenic collections. (Reprinted with permission from B Lorber [ed]: *Atlas of Infectious Diseases, vol VII: Intra-abdominal Infections, Hepatitis, and Gastroenteritis*. Philadelphia, Current Medicine, 1996, p 1.13.)

A small amount of serous fluid is normally present in the peritoneal space, with a protein content (consisting mainly of albumin) of <30 g/L and <300 white blood cells (WBCs, generally mononuclear cells) per microliter. In bacterial infections, leukocyte recruitment into the infected peritoneal cavity consists of an early influx of polymorphonuclear leukocytes (PMNs) and a prolonged subsequent phase of mononuclear cell migration. The phenotype of the infiltrating leukocytes during the course of inflammation is regulated primarily by resident-cell chemokine synthesis.

PRIMARY (SPONTANEOUS) BACTERIAL PERITONITIS

Peritonitis is either primary (without an apparent source of contamination) or secondary. The types of organisms found and the clinical presentations of these two processes are different. In adults, primary bacterial peritonitis (PBP) occurs most commonly in conjunction with cirrhosis of the liver (frequently the result of alcoholism). However, the disease has been reported in adults with metastatic malignant disease, postnecrotic cirrhosis, chronic active hepatitis, acute viral hepatitis, congestive heart failure, systemic lupus erythematosus, and lymphedema as well as in patients with no underlying disease. Although PBP virtually always develops in patients with preexisting ascites, it is, in general, an uncommon event, occurring in ≤10% of cirrhotic patients. The cause of PBP has not been established definitively but is believed to involve hematogenous spread of organisms in a patient in whom a diseased liver and altered portal circulation result in a defect in the usual filtration function. Organisms multiply in ascites, a good medium for growth. The proteins of the complement cascade have been found in peritoneal fluid, with lower levels in cirrhotic patients than in patients with ascites of other etiologies. The opsonic and phagocytic properties of PMNs are diminished in patients with advanced liver disease. Cirrhosis is associated with alterations in the gut microbiota, including an increased prevalence of potentially pathogenic bacteria such as Enterobacteriaceae. Small-intestinal bacterial overgrowth is frequently present in advanced stages of liver cirrhosis and has been linked with pathologic bacterial translocation and PBP. Factors promoting these changes in cirrhosis may include deficiencies in Paneth cell defensins, reduced intestinal motility, decreased pancreaticobiliary secretions, and portal-hypertensive enteropathy.

The presentation of PBP differs from that of secondary peritonitis. The most common manifestation is fever, which is reported in up to 80% of patients. Ascites is found but virtually always predates infection. Abdominal pain, an acute onset of symptoms, and peritoneal irritation during physical examination can be helpful diagnostically, but the absence of any of these findings does not exclude this often-subtle diagnosis. Nonlocalizing symptoms (such as malaise, fatigue, or encephalopathy) without another clear etiology should also prompt consideration of PBP in a susceptible patient. It is vital to sample the peritoneal fluid of any cirrhotic patient with ascites and fever. The finding of >250 PMNs/μL is diagnostic for PBP, according to Conn (<http://jac.oxfordjournals.org/cgi/content/full/47/3/369>). This criterion does not apply to secondary peritonitis (see below). The microbiology of PBP is also distinctive. While enteric gram-negative bacilli such as *Escherichia coli* are most commonly encountered, gram-positive organisms such as streptococci, enterococci, or even pneumococci are sometimes found. In an important development, widespread use of quinolones to prevent PBP in high-risk subgroups of patients, frequent hospitalizations, and exposure to broad-spectrum antibiotics have led to a change in flora of infections in patients with cirrhosis, with more gram-positive bacteria in recent years. Risk factors for multiresistant infections include nosocomial origin of infection, long-term norfloxacin prophylaxis, recent infection with multiresistant bacteria, and recent use of β-lactam antibiotics. In PBP, a single organism is typically isolated; anaerobes are found less frequently in PBP than in secondary peritonitis, in which a mixed flora including anaerobes is the rule. In fact, if PBP is suspected and multiple organisms including anaerobes are recovered from the peritoneal fluid, the diagnosis must be reconsidered and a source of secondary peritonitis sought.



FIGURE 159-2 Pneumoperitoneum. Free air under the diaphragm on an upright chest film suggests the presence of a bowel perforation and associated peritonitis. (Image courtesy of Dr. John Braver; with permission.)

The diagnosis of PBP is not easy. It depends on the exclusion of a primary intraabdominal source of infection. Contrast-enhanced CT is useful in identifying an intraabdominal source for infection. It may be difficult to recover organisms from cultures of peritoneal fluid, presumably because the burden of organisms is low. However, the yield can be improved if 10 mL of peritoneal fluid is placed directly into a blood culture bottle. Because bacteremia frequently accompanies PBP, blood should be cultured simultaneously. To maximize the yield, culture samples should be collected prior to administration of antibiotics. No specific radiographic studies are helpful in the diagnosis of PBP. A plain film of the abdomen would be expected to show ascites. Chest and abdominal radiography should be performed in patients with abdominal pain to exclude free air, which signals a perforation (Fig. 159-2).

TREATMENT PRIMARY BACTERIAL PERITONITIS

Treatment for PBP is directed at the isolate from blood or peritoneal fluid. Gram's staining of peritoneal fluid often gives negative results in PBP. Therefore, until culture results become available, therapy should cover gram-negative aerobic bacilli and gram-positive cocci. Third-generation cephalosporins such as cefotaxime (2 g q8h, administered IV) provide reasonable initial coverage in moderately ill patients. Broad-spectrum antibiotics, such as penicillin/ β -lactamase inhibitor combinations (e.g., piperacillin/tazobactam, 3.375 g q6h IV for adults with normal renal function) or ceftriaxone (2 g q24h IV), are also options. Broader empirical coverage aimed at resistant hospital-acquired gram-negative bacteria (e.g., treatment with carbapenem) may be appropriate for nosocomially acquired PBP until culture results become available. Empirical coverage for anaerobes is not necessary. A mortality benefit from albumin (1.5 g/kg of body weight within 6 h of detection and 1.0 g/kg on day 3) has been demonstrated for patients who present with serum creatinine levels ≥ 1 mg/dL, blood urea nitrogen levels ≥ 30 mg/dL, or total bilirubin levels ≥ 4 mg/dL but not for patients who do not meet these criteria. After the infecting organism is identified, therapy should be narrowed to target the specific pathogen. Patients with PBP usually respond within 72 h to appropriate antibiotic therapy. Antimicrobial

treatment can be administered for as little as 5 days if rapid improvement occurs and blood cultures are negative, but a course of up to 2 weeks may be required for patients with bacteremia and for those whose improvement is slow. Persistence of WBCs in the ascitic fluid after therapy should prompt a search for additional diagnoses.

Prevention • PRIMARY PREVENTION Several observational studies and a meta-analysis raise the concern that proton pump inhibitor therapy may increase the risk of PBP. No prospective studies have yet addressed whether avoidance of such therapy may prevent PBP. Nonselective beta blockers may prevent secondary bacterial peritonitis. A 2012 guideline from the American Association for the Study of Liver Diseases recommends chronic antibiotic prophylaxis with a regimen described in the next section for patients who are at highest risk for PBP—that is, those with an ascitic-fluid total protein level <1.5 g/dL along with impaired renal function (creatinine, ≥ 1.2 mg/dL; blood urea nitrogen, ≥ 25 mg/dL; or serum sodium, ≤ 130 mg/dL) and/or liver failure (Child-Pugh score, ≥ 9 ; and bilirubin, ≥ 3 mg/dL). A 7-day course of antibiotic prophylaxis is recommended for patients with cirrhosis and gastrointestinal bleeding.

SECONDARY PREVENTION PBP has a high rate of recurrence. Up to 70% of patients experience a recurrence within 1 year. Antibiotic prophylaxis is recommended for patients with a history of PBP to reduce this rate to $<20\%$ and improve short-term survival rates. Prophylactic regimens for adults with normal renal function include fluoroquinolones (ciprofloxacin, 750 mg weekly; norfloxacin, 400 mg/d) or trimethoprim-sulfamethoxazole (one double-strength tablet daily). However, long-term administration of broad-spectrum antibiotics in this setting has been shown to increase the risk of severe staphylococcal infections.

SECONDARY PERITONITIS

Secondary peritonitis develops when bacteria contaminate the peritoneum as a result of spillage from an intraabdominal viscus. The organisms found almost always constitute a mixed flora in which facultative gram-negative bacilli and anaerobes predominate, especially when the contaminating source is colonic. Early in the course of infection, when the host response is directed toward containment, exudate containing fibrin and PMNs is found. Early death in this setting is attributable to gram-negative bacillary sepsis and to potent endotoxins circulating in the bloodstream (Chap. 325). Gram-negative bacilli, particularly *E. coli*, are common bloodstream isolates, but *Bacteroides fragilis* bacteremia also occurs. The severity of abdominal pain and the clinical course depend on the inciting process. The organisms isolated from the peritoneum also vary with the source of the initial process and the normal flora at that site. Secondary peritonitis can result primarily from chemical irritation and/or bacterial contamination. For example, as long as the patient is not achlorhydric, a ruptured gastric ulcer will release low-pH gastric contents that will serve as a chemical irritant. The normal flora of the stomach comprises the same organisms found in the oropharynx but in lower numbers. Thus, the bacterial burden in a ruptured ulcer is negligible compared with that in a ruptured appendix. The normal flora of the colon below the ligament of Treitz contains $\sim 10^{11}$ anaerobic organisms/g of feces but only 10^8 aerobes/g; therefore, anaerobic species account for 99.9% of the bacteria. Leakage of colonic contents (pH 7–8) does not cause significant chemical peritonitis, but infection is intense because of the heavy bacterial load.

Depending on the inciting event, local symptoms may occur in secondary peritonitis—for example, epigastric pain from a ruptured gastric ulcer. In appendicitis (Chap. 356), the initial presenting symptoms are often vague, with periumbilical discomfort and nausea followed in a number of hours by pain more localized to the right lower quadrant. Unusual locations of the appendix (including a retrocecal position) can complicate this presentation further. Once infection has spread to the peritoneal cavity, pain increases, particularly with infection involving the parietal peritoneum, which is innervated extensively. Patients usually lie motionless, often with knees drawn up to avoid stretching the nerve fibers of the peritoneal cavity. Coughing and sneezing, which increase pressure within the peritoneal cavity, are associated with sharp

848 pain. There may or may not be pain localized to the infected or diseased organ from which secondary peritonitis has arisen. Patients with secondary peritonitis generally have abnormal findings on abdominal examination, with marked voluntary and involuntary guarding of the anterior abdominal musculature. Later findings include tenderness, especially rebound tenderness. In addition, there may be localized findings in the area of the inciting event. In general, patients are febrile, with marked leukocytosis and a left shift of the WBCs to band forms.

While recovery of organisms from peritoneal fluid is easier in secondary than in primary peritonitis, a tap of the abdomen is rarely the procedure of choice in secondary peritonitis. An exception is in cases involving trauma, where the possibility of a hemoperitoneum may need to be excluded early. Emergent studies (such as abdominal CT) to find the source of peritoneal contamination should be undertaken if the patient is hemodynamically stable; unstable patients may require surgical intervention without prior imaging.

TREATMENT SECONDARY PERITONITIS

Treatment for secondary peritonitis includes early administration of antibiotics aimed particularly at aerobic gram-negative bacilli and anaerobes (see below). Mild to moderate disease can be treated with many drugs covering these organisms, including broad-spectrum penicillin/β-lactamase inhibitor combinations (e.g., ticarcillin/clavulanate, 3.1 g q4–6h IV), ceftazidime (2 g q4–6h IV), or a combination of either a fluoroquinolone (e.g., levofloxacin, 750 mg q24h IV) or a third-generation cephalosporin (e.g., ceftriaxone, 2 g q24h IV) plus metronidazole (500 mg q8h IV). Patients in intensive care units should receive imipenem (500 mg q6h IV), meropenem (1 g q8h IV), or combinations of drugs, such as ampicillin plus metronidazole plus ciprofloxacin. The role of enterococci and *Candida* species in mixed infections is controversial. Secondary peritonitis usually requires both surgical intervention to address the inciting process and antibiotics to treat early bacteremia, to decrease the incidence of abscess formation and wound infection, and to prevent distant spread of infection. Although surgery is rarely indicated in PBP in adults, it may be life-saving in secondary peritonitis. Recombinant human activated protein C (APC) was considered at one time for treatment of severe sepsis from causes including secondary peritonitis but was withdrawn from the market in 2011 after it was determined that the drug was associated with an increased risk of bleeding and that evidence for its beneficial effects was inadequate. Thus APC should not be used for sepsis or septic shock outside randomized clinical trials.

Peritonitis may develop as a complication of abdominal surgeries. These infections may be accompanied by localizing pain and/or nonlocalizing signs or symptoms such as fever, malaise, anorexia, and toxicity. As a nosocomial infection, postoperative peritonitis may be associated with organisms such as staphylococci, components of the gram-negative hospital microflora, and the microbes that cause PBP and secondary peritonitis, as described above.

PERITONITIS IN PATIENTS UNDERGOING CONTINUOUS AMBULATORY PERITONEAL DIALYSIS

A third type of peritonitis arises in patients who are undergoing continuous ambulatory peritoneal dialysis (CAPD). Unlike PBP and secondary peritonitis, which are caused by endogenous bacteria, CAPD-associated peritonitis usually involves skin organisms. The pathogenesis of infection is similar to that of intravascular device-related infection, in which skin organisms migrate along the catheter, which both serves as an entry point and exerts the effects of a foreign body. Exit-site or tunnel infection may or may not accompany CAPD-associated peritonitis. Like PBP, CAPD-associated peritonitis is usually caused by a single organism. Peritonitis is, in fact, the most common reason for discontinuation of CAPD. Improvements in equipment design, especially the Y-set connector, have resulted in a decrease from one case of peritonitis per 9 months of CAPD to one case per 24 months.

The clinical presentation of CAPD peritonitis resembles that of secondary peritonitis in that diffuse pain and peritoneal signs are common. The dialysate is usually cloudy and contains >100 WBCs/μL, >50% of which are neutrophils. However, the number of cells depends in part on dwell time. According to a guideline from the International Society for Peritoneal Dialysis (2010), for patients undergoing automated peritoneal dialysis who present during their nighttime treatment and whose dwell time is much shorter than with CAPD, the clinician should use the percentage of PMNs rather than the absolute number of WBCs to diagnose peritonitis. As the normal peritoneum has very few PMNs, a proportion above 50% is strong evidence of peritonitis even if the absolute WBC count does not reach 100/μL. Meanwhile, patients undergoing automated peritoneal dialysis without a daytime exchange who present with abdominal pain may have no fluid to withdraw, in which case 1 L of dialysate should be infused and permitted to dwell a minimum of 1–2 h, then drained, examined for turbidity, and sent for cell count with differential and culture. The differential (with a shortened dwell time) may be more useful than the absolute WBC count. In equivocal cases or in patients with systemic or abdominal symptoms in whom the effluent appears clear, a second exchange is performed, with a dwell time of at least 2 h. Clinical judgment should guide initiation of therapy.

The most common organisms are *Staphylococcus* species, which accounted for ~45% of cases in one series. Historically, coagulase-negative staphylococcal species were identified most commonly in these infections, but these isolates have more recently been decreasing in frequency. *Staphylococcus aureus* is more often involved among patients who are nasal carriers of the organism than among those who are not, and this organism is the most common pathogen in overt exit-site infections. Gram-negative bacilli and fungi such as *Candida* species are also found. Vancomycin-resistant enterococci and vancomycin-intermediate *S. aureus* have been reported to produce peritonitis in CAPD patients. The finding of more than one organism in dialysate culture should prompt evaluation for secondary peritonitis. As with PBP, culture of dialysate fluid in blood culture bottles improves the yield. To facilitate diagnosis, several hundred milliliters of removed dialysis fluid should be concentrated by centrifugation before culture.

TREATMENT CAPD PERITONITIS

Empirical therapy for CAPD peritonitis should be directed at *S. aureus*, coagulase-negative *Staphylococcus*, and gram-negative bacilli until the results of cultures become available. Guidelines suggest that agents should be chosen on the basis of local experience with resistant organisms. In some centers, a first-generation cephalosporin such as cefazolin (for gram-positive bacteria) and a fluoroquinolone or a third-generation cephalosporin such as ceftazidime (for gram-negative bacteria) may be reasonable; in areas with high rates of infection with methicillin-resistant *S. aureus*, vancomycin should be used instead of cefazolin, and gram-negative coverage may need to be broadened—e.g., with an aminoglycoside, ceftazidime, cefepime, or carbapenem. Broad coverage including vancomycin should be particularly considered for toxic patients and for those with exit-site infections. Vancomycin should also be included in the regimen if the patient has a history of colonization or infection with methicillin-resistant *S. aureus* or has a history of severe allergy to penicillins and cephalosporins. Loading doses are administered intraperitoneally; doses depend on the dialysis method and the patient's renal function. Antibiotics are given either continuously (i.e., with each exchange) or intermittently (i.e., once daily, with the dose allowed to remain in the peritoneal cavity for at least 6 h). If the patient is severely ill, IV antibiotics should be added at doses appropriate for the patient's degree of renal failure. The clinical response to an empirical treatment regimen should be rapid; if the patient has not responded after 48–96 h of treatment, new samples should be collected for cell counts and cultures, and catheter removal should be considered. For patients who lack exit-site or tunnel infection, the typical duration of antibiotic treatment is 14 days. For patients with exit-site or tunnel infection, catheter removal should be considered,

and a longer duration of antibiotic therapy (up to 21 days) may be appropriate. In fungal infections, the catheter should be removed immediately.

TUBERCULOUS PERITONITIS

See Chap. 202.

INTRAABDOMINAL ABSCESSES

INTRAPERITONEAL ABSCESSES

Abscess formation is common in untreated peritonitis if overt gram-negative sepsis either does not develop or develops but is not fatal. In experimental models of abscess formation, mixed aerobic and anaerobic organisms have been implanted intraperitoneally. Without therapy directed at anaerobes, animals develop intraabdominal abscesses. As in humans, these experimental abscesses may stud the peritoneal cavity, lie within the omentum or mesentery, or even develop on the surface of or within viscera such as the liver.

Pathogenesis and Immunity There is often disagreement about whether an abscess represents a disease state or a host response. In a sense, it represents both: while an abscess is an infection in which viable infecting organisms and PMNs are contained in a fibrous capsule, it is also a process by which the host confines microbes to a limited space, thereby preventing further spread of infection. In any event, abscesses do cause significant symptoms, and patients with abscesses can be quite ill. Experimental work has helped to define both the host cells and the bacterial virulence factors responsible—most notably in the case of *B. fragilis*. This organism, although accounting for only 0.5% of the normal colonic flora, is the anaerobe most frequently isolated from intraabdominal infections, is especially prominent in abscesses, and is the most common anaerobic bloodstream isolate. On clinical grounds, therefore, *B. fragilis* appears to be uniquely virulent. Moreover, *B. fragilis* acts alone to cause abscesses in animal models of intraabdominal infection, whereas most other *Bacteroides* species must act synergistically with a facultative organism to induce abscess formation.

Of the several virulence factors identified in *B. fragilis*, one is critical: the capsular polysaccharide complex found on the bacterial surface. This complex comprises at least eight distinct surface polysaccharides. Structural analysis of these polysaccharides has shown an unusual motif of oppositely charged sugars. Polysaccharides having these *zwitterionic* characteristics, such as polysaccharide A, evoke a host response in the peritoneal cavity that localizes bacteria into abscesses. *B. fragilis* and polysaccharide A have been found to adhere to primary mesothelial cells *in vitro*; this adherence, in turn, stimulates the production of tumor necrosis factor α and intercellular adhesion molecule 1 by peritoneal macrophages. Although abscesses characteristically contain PMNs, the process of abscess induction depends on the stimulation of T lymphocytes by these unique *zwitterionic* polysaccharides. The stimulated CD4⁺ T lymphocytes secrete leukoattractant cytokines and chemokines. The alternative pathway of complement and fibrinogen also participate in abscess formation.

While antibodies to the capsular polysaccharide complex enhance bloodstream clearance of *B. fragilis*, CD4⁺ T cells are critical in immunity to abscesses. When administered subcutaneously, *B. fragilis* polysaccharide A has immunomodulatory characteristics and stimulates CD4⁺ T regulatory cells via an interleukin 2–dependent mechanism to produce interleukin 10. Interleukin 10 downregulates the inflammatory response, thereby preventing abscess formation.

Clinical Presentation Of all intraabdominal abscesses, 74% are intraperitoneal or retroperitoneal and are not visceral. Most intraperitoneal abscesses result from fecal spillage from a colonic source, such as an inflamed appendix. Abscesses can also arise from other processes. They usually form within weeks of the development of peritonitis and may be found in a variety of locations from omentum to mesentery, pelvis to psoas muscles, and subphrenic space to a visceral organ such as the liver, where they may develop either on the surface of the organ or within it. Periappendiceal and diverticular abscesses occur commonly. Diverticular abscesses are least likely to rupture. Infections

of the female genital tract and pancreatitis are also among the more common causative events. When abscesses occur in the female genital tract—either as a primary infection (e.g., tuboovarian abscess) or as an infection extending into the pelvic cavity or peritoneum—*B. fragilis* figures prominently among the organisms isolated. *B. fragilis* is not found in large numbers in the normal vaginal flora. For example, it is encountered less commonly in pelvic inflammatory disease and endometritis without an associated abscess. In pancreatitis with leakage of damaging pancreatic enzymes, inflammation is prominent. Therefore, clinical findings such as fever, leukocytosis, and even abdominal pain do not distinguish pancreatitis itself from complications such as pancreatic pseudocyst, pancreatic abscess (Chap. 371), or intraabdominal collections of pus. Especially in cases of necrotizing pancreatitis, in which the incidence of local pancreatic infection may be as high as 30%, needle aspiration under CT guidance is performed to sample fluid for culture. Many centers prescribe preemptive antibiotics for patients with necrotizing pancreatitis. Imipenem is frequently used for this purpose because it reaches high tissue levels in the pancreas (although it is not unique in this regard). Recent randomized controlled studies have not demonstrated a benefit from this practice, and some guidelines no longer recommend preemptive antibiotics for patients with acute pancreatitis. If needle aspiration yields infected fluid in the setting of acute necrotizing pancreatitis, antibiotic treatment is appropriate in conjunction with surgical and/or percutaneous drainage of infected material. Infected pseudocysts that occur remotely from acute pancreatitis are unlikely to be associated with significant amounts of necrotic tissue and may be treated with either surgical or percutaneous catheter drainage in conjunction with appropriate antibiotic therapy.

Diagnosis Scanning procedures have considerably facilitated the diagnosis of intraabdominal abscesses. Abdominal CT probably has the highest yield, although ultrasonography is particularly useful for the right upper quadrant, kidneys, and pelvis. Both indium-labeled WBCs and gallium tend to localize in abscesses and may be useful in finding a collection. Because gallium is taken up in the bowel, indium-labeled WBCs may have a slightly greater yield for abscesses near the bowel. Neither indium-labeled WBC nor gallium scans serve as a basis for a definitive diagnosis, however; both need to be followed by other, more specific studies, such as CT, if an area of possible abnormality is identified. Abscesses contiguous with or contained within diverticula are particularly difficult to diagnose with scanning procedures. Although barium should not be injected if a perforation is suspected, a barium enema occasionally may detect a diverticular abscess not diagnosed by other procedures. If one study is negative, a second study sometimes reveals a collection. Although exploratory laparotomy has been less commonly used since the advent of CT, this procedure still must be undertaken on occasion if an abscess is strongly suspected on clinical grounds.

TREATMENT INTRAPERITONEAL ABSCESSES

An algorithm for the management of patients with intraabdominal (including intraperitoneal) abscesses by percutaneous drainage is presented in Fig. 159-3. The treatment of intraabdominal infections involves the determination of the initial focus of infection, the administration of broad-spectrum antibiotics targeting the organisms involved, and the performance of a drainage procedure if one or more definitive abscesses have formed. Antimicrobial therapy, in general, is adjunctive to drainage and/or surgical correction of an underlying lesion or process in intraabdominal abscesses. Unlike the intraabdominal abscesses resulting from most causes, for which drainage of some kind is generally required, abscesses associated with diverticulitis usually wall off locally after rupture of a diverticulum, so that surgical intervention is not routinely required.

A number of agents exhibit excellent activity against aerobic gram-negative bacilli. Because death in intraabdominal sepsis is linked to gram-negative bacteremia, empirical therapy for intraabdominal infection always needs to include adequate coverage of gram-negative aerobic, facultative, and anaerobic organisms. Even

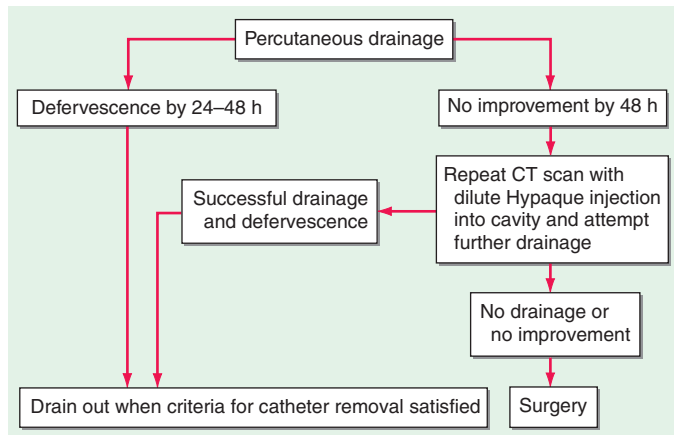


FIGURE 159-3 Algorithm for the management of patients with intraabdominal abscesses using percutaneous drainage.

Antimicrobial therapy should be administered concomitantly. (Reprinted with permission from B Lorber [ed]: *Atlas of Infectious Diseases, vol VII: Intra-abdominal Infections, Hepatitis, and Gastroenteritis*. Philadelphia, Current Medicine, 1996, p 1.30, as adapted from OD Rotstein, RL Simmons, in SL Gorbach et al [eds]: *Infectious Diseases*. Philadelphia, Saunders, 1992, p 668.)

if anaerobes are not cultured from clinical specimens, they still must be covered by the therapeutic regimen. Empirical antibiotic therapy should be the same as that discussed above for secondary peritonitis.

VISCERAL ABSCESSSES

Liver Abscesses The liver is the organ most subject to the development of abscesses. In one study of 540 intraabdominal abscesses, 26% were visceral. Liver abscesses made up 13% of the total number, or 48% of all visceral abscesses. Liver abscesses may be solitary or multiple; they may arise from hematogenous spread of bacteria or from local spread from contiguous sites of infection within the peritoneal cavity. In the past, appendicitis with rupture and subsequent spread of infection was the most common source for a liver abscess. Currently, associated disease of the biliary tract is most common. Pylephlebitis (suppurative thrombosis of the portal vein), usually arising from infection in the pelvis but sometimes from infection elsewhere in the peritoneal cavity, is another common source for bacterial seeding of the liver.

Fever is the most common presenting sign of liver abscess. Some patients, particularly those with associated disease of the biliary tract, have symptoms and signs localized to the right upper quadrant, including pain, guarding, punch tenderness, and even rebound tenderness. Nonspecific symptoms, such as chills, anorexia, weight loss, nausea, and vomiting, may also develop. Only 50% of patients with liver abscesses, however, have hepatomegaly, right-upper-quadrant tenderness, or jaundice; thus, one-half of patients have no symptoms or signs to direct attention to the liver. Fever of unknown origin may be the only manifestation of liver abscess, especially in the elderly. Diagnostic studies of the abdomen, especially the right upper quadrant, should be a part of any workup for fever of unknown origin. The single most reliable laboratory finding is an elevated serum concentration of alkaline phosphatase, which is documented in 70% of patients with liver abscesses. Other tests of liver function may yield normal results, but 50% of patients have elevated serum levels of bilirubin, and 48% have elevated concentrations of aspartate aminotransferase. Other laboratory findings include leukocytosis in 77% of patients, anemia (usually normochromic, normocytic) in 50%, and hypoalbuminemia in 33%. Concomitant bacteremia is found in one-third to one-half of patients. A liver abscess is sometimes suggested by chest radiography, especially if a new elevation of the right hemidiaphragm is seen; other suggestive findings include a right basilar infiltrate and a right pleural effusion.

Imaging studies are the most reliable methods for diagnosing liver abscesses. These studies include ultrasonography, CT (Fig. 159-4),



FIGURE 159-4 Multilocular liver abscess on CT scan. Multiple or multilocular abscesses are more common than solitary abscesses. (Reprinted with permission from B Lorber [ed]: *Atlas of Infectious Diseases, vol VII: Intra-abdominal Infections, Hepatitis, and Gastroenteritis*. Philadelphia, Current Medicine, 1996, Fig. 1.22.)

indium-labeled WBC or gallium scan, and MRI. More than one such study may be required.

Organisms recovered from liver abscesses vary with the source. In liver infection arising from the biliary tree, enteric gram-negative aerobic bacilli and enterococci are common isolates. *Klebsiella pneumoniae* liver abscess has been well described in Southeast Asia for more than 20 years and has become an emerging syndrome in North America and elsewhere. These community-acquired infections have been linked to a virulent hypermucoviscous *K. pneumoniae* phenotype and to a specific genotype. The typical syndrome includes liver abscess, bacteremia, and metastatic infection. Ampicillin/amoxicillin therapy started within the previous 30 days has been associated with increased risk for this syndrome, presumably because of selection for the causative strain. Unless previous surgery has been performed, anaerobes are not generally involved in liver abscesses arising from biliary infections. In contrast, in liver abscesses arising from pelvic and other intraperitoneal sources, a mixed flora including both aerobic and anaerobic species is common; *B. fragilis* is the species most frequently isolated. With hematogenous spread of infection, usually only a single organism is encountered; this species may be *S. aureus* or a streptococcal species such as one in the *Streptococcus milleri* group. Results of cultures obtained from drain sites are not reliable for defining the etiology of infections. Liver abscesses may also be caused by *Candida* species; such abscesses usually follow fungemia in patients receiving chemotherapy for cancer and often present when PMNs return after a period of neutropenia. Amebic liver abscesses are not an uncommon problem (Chap. 247). Amebic serologic testing gives positive results in >95% of cases. In addition, polymerase chain reaction (PCR) testing has been used in recent years. Negative results from these studies help to exclude this diagnosis.

TREATMENT LIVER ABSCESSSES

(Fig. 159-3) Drainage is the mainstay of therapy for intraabdominal abscesses, including liver abscesses; the approach can be either percutaneous (with a pigtail catheter kept in place or possibly with a device that can perform pulse lavage to fragment and evacuate the semisolid contents of a liver abscess) or surgical. However, there is growing interest in medical management alone for pyogenic liver abscesses. The drugs used for empirical therapy include the same ones used in intraabdominal sepsis and secondary bacterial peritonitis. Usually, blood cultures and a diagnostic aspirate of abscess contents should be obtained before the initiation of empirical therapy, with antibiotic choices adjusted when the results of Gram's staining and culture become available. Cases treated without

definitive drainage generally require longer courses of antibiotic therapy. When percutaneous drainage was compared with open surgical drainage, the average length of hospital stay for the former was almost twice that for the latter, although both the time required for fever to resolve and the mortality rate were the same for the two procedures. The mortality rate was appreciable despite treatment, averaging 15%. Several factors predict the failure of percutaneous drainage and therefore may favor primary surgical intervention. These factors include the presence of multiple, sizable abscesses; viscous abscess contents that tend to plug the catheter; associated disease (e.g., disease of the biliary tract) requiring surgery; the presence of yeast; communication with an untreated obstructed biliary tree; or the lack of a clinical response to percutaneous drainage in 4–7 days.

Treatment of candidal liver abscesses often entails initial administration of amphotericin B or liposomal amphotericin, with subsequent fluconazole therapy (Chap. 240). In some cases, therapy with fluconazole alone (6 mg/kg daily) may be used—e.g., in clinically stable patients whose infecting isolate is susceptible to this drug.

Splenic Abscesses Splenic abscesses are much less common than liver abscesses. The incidence of splenic abscesses has ranged from 0.14% to 0.7% in various autopsy series. The clinical setting and the organisms isolated usually differ from those for liver abscesses. The degree of clinical suspicion for splenic abscess needs to be high because this condition is frequently fatal if left untreated. Even in the most recently published series, diagnosis was made only at autopsy in 37% of cases. Although splenic abscesses may arise occasionally from contiguous spread of infection or from direct trauma to the spleen, hematogenous spread of infection is more common. Bacterial endocarditis is the most common associated infection (Chap. 155). Splenic abscesses can develop in patients who have received extensive immunosuppressive therapy (particularly those with malignancy involving the spleen) and in patients with hemoglobinopathies or other hematologic disorders (especially sickle cell anemia).

Although ~50% of patients with splenic abscesses have abdominal pain, the pain is localized to the left upper quadrant in only one-half of these cases. Splenomegaly is found in ~50% of cases. Fever and leukocytosis are generally present; the development of fever preceded diagnosis by an average of 20 days in one series. Left-sided chest findings may include abnormalities to auscultation, and chest radiographic findings may include an infiltrate or a left-sided pleural effusion. CT scan of the abdomen has been the most sensitive diagnostic tool. Ultrasonography can yield the diagnosis but is less sensitive. Liver-spleen scan or gallium scan may also be useful. Streptococcal species are the most common bacterial isolates from splenic abscesses, followed by *S. aureus*—presumably reflecting the associated endocarditis. An increase in the prevalence of gram-negative aerobic isolates from splenic abscesses has been reported; these organisms often derive from a urinary tract focus, with associated bacteremia, or from another intraabdominal source. *Salmonella* species are seen fairly commonly, especially in patients with sickle cell hemoglobinopathy. Anaerobic species accounted for only 5% of isolates in the largest collected series, but the reporting of a number of “sterile abscesses” may indicate that optimal techniques for the isolation of anaerobes were not used.

TREATMENT SPLENIC ABSCESSSES

Because of the high mortality figures reported for splenic abscesses, splenectomy with adjunctive antibiotics has traditionally been considered standard treatment and remains the best approach for complex, multilocular abscesses or multiple abscesses. However, percutaneous drainage has worked well for single, small (<3-cm) abscesses in some studies and may also be useful for patients with high surgical risk. Patients undergoing splenectomy should be vaccinated against encapsulated organisms (*Streptococcus pneumoniae*, *Haemophilus influenzae*, *Neisseria meningitidis*). The most important factor in successful treatment of splenic abscesses is early diagnosis.

Perinephric and Renal Abscesses Perinephric and renal abscesses are not common. The former accounted for only ~0.02% of hospital admissions and the latter for ~0.2% in Altemeier’s series of 540 intraabdominal abscesses. Before antibiotics became available, most renal and perinephric abscesses were hematogenous in origin, usually complicating prolonged bacteremia, with *S. aureus* most commonly recovered. Now, in contrast, >75% of perinephric and renal abscesses arise from a urinary tract infection. Infection ascends from the bladder to the kidney, with pyelonephritis preceding abscess development. Bacteria may directly invade the renal parenchyma from medulla to cortex. Local vascular channels within the kidney may also facilitate the transport of organisms. Areas of abscess developing within the parenchyma may rupture into the perinephric space. The kidneys and adrenal glands are surrounded by a layer of perirenal fat that, in turn, is surrounded by Gerota’s fascia, which extends superiorly to the diaphragm and inferiorly to the pelvic fat. Abscesses extending into the perinephric space may track through Gerota’s fascia into the psoas or transversalis muscles, into the anterior peritoneal cavity, superiorly to the subdiaphragmatic space, or inferiorly to the pelvis. Of the risk factors that have been associated with the development of perinephric abscesses, the most important is concomitant nephrolithiasis obstructing urinary flow. Of patients with perinephric abscess, 20–60% have renal stones. Other structural abnormalities of the urinary tract, prior urologic surgery, trauma, and diabetes mellitus have also been identified as risk factors.

The organisms most frequently encountered in perinephric and renal abscesses are *E. coli*, *Proteus* species, and *Klebsiella* species. *E. coli*, the aerobic species most commonly found in the colonic flora, seems to have unique virulence properties in the urinary tract, including factors promoting adherence to uroepithelial cells. The urease of *Proteus* species splits urea, thereby creating a more alkaline and more hospitable environment for bacterial proliferation. *Proteus* species are frequently found in association with large struvite stones caused by the precipitation of magnesium ammonium sulfate in an alkaline environment. These stones serve as a nidus for recurrent urinary tract infection. Although a single bacterial species is usually recovered from a perinephric or renal abscess, multiple species may also be found. If a urine culture is not contaminated with periurethral flora and is found to contain more than one organism, a perinephric abscess or renal abscess should be considered in the differential diagnosis. Urine cultures may also be polymicrobial in cases of bladder diverticulum.

Candida species can cause renal abscesses. This fungus may spread to the kidney hematogenously or by ascension from the bladder. The hallmark of the latter route of infection is ureteral obstruction with large fungal balls.

The presentation of perinephric and renal abscesses is quite nonspecific. Flank pain and abdominal pain are common. At least 50% of patients are febrile. Pain may be referred to the groin or leg, particularly with extension of infection. The diagnosis of perinephric abscess, like that of splenic abscess, is frequently delayed, and the mortality rate in some series is appreciable, although lower than in the past. Perinephric or renal abscess should be most seriously considered when a patient presents with symptoms and signs of pyelonephritis and remains febrile after 4 or 5 days of treatment. Moreover, when a urine culture yields a polymicrobial flora, when a patient is known to have renal stones, or when fever and pyuria coexist with a sterile urine culture, these diagnoses should be entertained.

Renal ultrasonography and abdominal CT are the most useful diagnostic modalities. If a renal or perinephric abscess is diagnosed, nephrolithiasis should be excluded, especially when a high urinary pH suggests the presence of a urea-splitting organism.

TREATMENT PERINEPHRIC AND RENAL ABSCESSSES

Treatment for perinephric and renal abscesses, like that for other intraabdominal abscesses, includes drainage of pus and antibiotic therapy directed at the organism(s) recovered. For perinephric abscesses, percutaneous drainage is usually successful.

852 Psoas Abscesses The psoas muscle is another location in which abscesses are encountered. Psoas abscesses may arise from a hematogenous source, by contiguous spread from an intraabdominal or pelvic process, or by contiguous spread from nearby bony structures (e.g., vertebral bodies). Associated osteomyelitis due to spread from bone to muscle or from muscle to bone is common in psoas abscesses. When Pott's disease was common, *Mycobacterium tuberculosis* was a frequent cause of psoas abscess. Currently, either *S. aureus* or a mixture of enteric organisms including aerobic and anaerobic gram-negative bacilli is usually isolated from psoas abscesses in the United States. *S. aureus* is most likely to be isolated when a psoas abscess arises from hematogenous spread or a contiguous focus of osteomyelitis; a mixed enteric flora is the most likely etiology when the abscess has an intra-abdominal or pelvic source. Patients with psoas abscesses frequently present with fever, lower abdominal or back pain, or pain referred to the hip or knee. CT is the most useful diagnostic technique.

TREATMENT PSOAS ABSCESSSES

Treatment includes surgical drainage and the administration of an antibiotic regimen directed at the inciting organism(s).


Pancreatic Abscesses See Chap. 371.

ACKNOWLEDGMENT

The substantial contributions of Dori F. Zaleznik, MD, to this chapter in previous editions are gratefully acknowledged.

160 Acute Infectious Diarrheal Diseases and Bacterial Food Poisoning

Regina C. LaRocque, Edward T. Ryan, Stephen B. Calderwood

 Acute diarrheal disease is a leading cause of illness globally and is associated with an estimated 1.4 million deaths per year. Among children <5 years of age, diarrheal disease is second only to lower respiratory infection as the most common infectious cause of death. The incidence rate of diarrheal disease among children in low- and middle-income countries is estimated to be 2.9 episodes per child per year, for a total of 1.7 billion episodes annually. The morbidity from diarrhea is also significant. Recurrent intestinal infections are associated with physical and mental stunting, wasting, micronutrient

deficiencies, and malnutrition. In short, diarrheal disease is a driving factor in global morbidity and mortality.

The wide range of clinical manifestations of acute gastrointestinal illnesses is matched by the wide variety of infectious agents involved, including viruses, bacteria, and parasites (Table 160-1). This chapter discusses factors that enable gastrointestinal pathogens to cause disease, reviews host defense mechanisms, and delineates an approach to the evaluation and treatment of patients presenting with acute diarrhea. Individual organisms causing acute gastrointestinal illnesses are discussed in detail in subsequent chapters.

PATHOGENIC MECHANISMS

Enteric pathogens have developed a variety of tactics to overcome host defenses. Understanding the virulence factors employed by these organisms is important in the diagnosis and treatment of clinical disease.

INOCULUM SIZE

The number of microorganisms that must be ingested to cause disease varies considerably from species to species. For *Shigella*, enterohemorrhagic *Escherichia coli*, *Giardia lamblia*, or *Entamoeba*, as few as 10–100 bacteria or cysts can produce infection, while 10⁵–10⁸ *Vibrio cholerae* organisms must be ingested to cause disease. The infective dose of *Salmonella* varies widely, depending on the species, host, and food vehicle. The ability of organisms to overcome host defenses has important implications for transmission; *Shigella*, enterohemorrhagic *E. coli*, *Entamoeba*, and *Giardia* can spread by person-to-person contact, whereas under some circumstances *Salmonella* may have to grow in food for several hours before reaching an effective infectious dose.

ADHERENCE

Many organisms must adhere to the gastrointestinal mucosa as an initial step in the pathogenic process; thus, organisms that can compete with the normal bowel flora and colonize the mucosa have an important advantage in causing disease. Specific cell-surface proteins involved in attachment of bacteria to intestinal cells are important virulence determinants. *V. cholerae*, for example, adheres to the brush border of small-intestinal enterocytes via specific surface adhesins, including the toxin-coregulated pilus and other accessory colonization factors. Enterotoxigenic *E. coli*, which causes watery diarrhea, produces an adherence protein called *colonization factor antigen* that is necessary for colonization of the upper small intestine by the organism prior to the production of enterotoxin. Enteropathogenic *E. coli*, an agent of diarrhea in young children, and enterohemorrhagic *E. coli*, which causes hemorrhagic colitis and the hemolytic-uremic syndrome, produce virulence determinants that allow these organisms to attach to and efface the brush border of the intestinal epithelium.

TOXIN PRODUCTION

The production of one or more exotoxins is important in the pathogenesis of numerous enteric organisms. Such toxins include *enterotoxins*, which cause watery diarrhea by acting directly on secretory mechanisms

TABLE 160-1 GASTROINTESTINAL PATHOGENS CAUSING ACUTE DIARRHEA

Mechanism	Location	Illness	Stool Findings	Examples of Pathogens Involved
Noninflammatory (enterotoxin)	Proximal small bowel	Watery diarrhea	No fecal leukocytes; mild or no increase in fecal lactoferrin	<i>Vibrio cholerae</i> , enterotoxigenic <i>Escherichia coli</i> (LT and/or ST), enteroaggregative <i>E. coli</i> , <i>Clostridium perfringens</i> , <i>Bacillus cereus</i> , <i>Staphylococcus aureus</i> , <i>Aeromonas hydrophila</i> , <i>Plesiomonas shigelloides</i> , rotavirus, norovirus, enteric adenoviruses, <i>Giardia lamblia</i> , <i>Cryptosporidium</i> spp., <i>Cyclospora</i> spp., microsporidia
Inflammatory (invasion or cytotoxin)	Colon or distal small bowel	Dysentery or inflammatory diarrhea	Fecal polymorphonuclear leukocytes; substantial increase in fecal lactoferrin	<i>Shigella</i> spp., <i>Salmonella</i> spp., <i>Campylobacter jejuni</i> , enterohemorrhagic <i>E. coli</i> , enteroinvasive <i>E. coli</i> , <i>Yersinia enterocolitica</i> , <i>Listeria monocytogenes</i> , <i>Vibrio parahaemolyticus</i> , <i>Clostridium difficile</i> , <i>A. hydrophila</i> , <i>P. shigelloides</i> , <i>Entamoeba histolytica</i> , <i>Klebsiella oxytoca</i>
Penetrating	Distal small bowel	Enteric fever	Fecal mononuclear leukocytes	<i>Salmonella typhi</i> , <i>Y. enterocolitica</i>

Abbreviations: LT, heat-labile enterotoxin; ST, heat-stable enterotoxin.

in the intestinal mucosa; *cytotoxins*, which cause destruction of mucosal cells and associated inflammatory diarrhea; and *neurotoxins*, which act directly on the central or peripheral nervous system.

The prototypical enterotoxin is cholera toxin, a heterodimeric protein composed of one A and five B subunits. The A subunit contains the enzymatic activity of the toxin, while the B subunit pentamer binds holotoxin to the enterocyte surface receptor, the ganglioside G_{M1} . After the binding of holotoxin, a fragment of the A subunit is translocated across the eukaryotic cell membrane into the cytoplasm, where it catalyzes the adenosine diphosphate ribosylation of a guanosine triphosphate-binding protein and causes persistent activation of adenylate cyclase. The end result is an increase of cyclic adenosine monophosphate in the intestinal mucosa, which increases Cl^- secretion and decreases Na^+ absorption, leading to a loss of fluid and the production of diarrhea.

Enterotoxigenic strains of *E. coli* may produce a protein called *heat-labile enterotoxin* (LT) that is similar to cholera toxin and causes secretory diarrhea by the same mechanism. Alternatively, enterotoxigenic strains of *E. coli* may produce *heat-stable enterotoxin* (ST), one form of which causes diarrhea by activation of guanylate cyclase and elevation of intracellular cyclic guanosine monophosphate. Some enterotoxigenic strains of *E. coli* produce both LT and ST.

Bacterial cytotoxins, in contrast, destroy intestinal mucosal cells and produce the syndrome of dysentery, with bloody stools containing inflammatory cells. Enteric pathogens that produce such cytotoxins include *Shigella dysenteriae* type 1, *Vibrio parahaemolyticus*, and *Clostridium difficile*. *S. dysenteriae* type 1 and Shiga toxin-producing strains of *E. coli* produce potent cytotoxins and have been associated with outbreaks of hemorrhagic colitis and hemolytic-uremic syndrome.

Neurotoxins are usually produced by bacteria outside the host and therefore cause symptoms soon after ingestion. Included are the staphylococcal and *Bacillus cereus* toxins, which act on the central nervous system to produce vomiting.

INVASION

Dysentery may result not only from the production of cytotoxins but also from bacterial invasion and destruction of intestinal mucosal cells. Infections due to *Shigella* and enteroinvasive *E. coli* are characterized by the organisms' invasion of mucosal epithelial cells, intraepithelial multiplication, and subsequent spread to adjacent cells. *Salmonella* causes inflammatory diarrhea by invasion of the bowel mucosa but generally is not associated with the destruction of enterocytes or the full clinical syndrome of dysentery. *Salmonella typhi* and *Yersinia enterocolitica* can penetrate intact intestinal mucosa, multiply intracellularly in Peyer's patches and intestinal lymph nodes, and then disseminate through the bloodstream to cause enteric fever, a syndrome characterized by fever, headache, relative bradycardia, abdominal pain, splenomegaly, and leukopenia.

HOST DEFENSES

Given the enormous number of microorganisms ingested with every meal, the normal host must combat a constant influx of potential enteric pathogens. Studies of infections in patients with alterations in defense mechanisms have led to a greater understanding of the variety of ways in which the normal host can protect itself against disease.

INTESTINAL MICROBIOTA

The large numbers of bacteria that normally inhabit the intestine (*the intestinal microbiota*) act as an important host defense mechanism, preventing colonization by potential enteric pathogens. Persons with fewer intestinal bacteria, such as infants who have not yet developed normal enteric colonization or patients receiving antibiotics, are at significantly greater risk of developing infections with enteric pathogens. The composition of the intestinal microbiota is as important as the number of organisms present. More than 99% of the normal colonic microbiota is made up of anaerobic bacteria, and the acidic pH and volatile fatty acids produced by these organisms appear to be critical elements in resistance to colonization.

GASTRIC ACID

The acidic pH of the stomach is an important barrier to enteric pathogens, and an increased frequency of infections due to *Salmonella*, *G. lamblia*, and a variety of helminths has been reported among patients who have undergone gastric surgery or are achlorhydric for some other reason. Neutralization of gastric acid with antacids, proton pump inhibitors, or H_2 blockers—a common practice in the management of hospitalized patients—similarly increases the risk of enteric colonization. In addition, some microorganisms can survive the extreme acidity of the gastric environment; rotavirus, for example, is highly stable to acidity.


INTESTINAL MOTILITY

Normal peristalsis is the major mechanism for clearance of bacteria from the proximal small intestine. When intestinal motility is impaired (e.g., by treatment with opiates or other antimotility drugs, anatomic abnormalities, or hypomotility states), the frequency of bacterial overgrowth and infection of the small bowel with enteric pathogens is increased. Some patients whose treatment for *Shigella* infection consists of diphenoxylate hydrochloride with atropine (Lomotil) experience prolonged fever and shedding of organisms, while patients treated with opiates for mild *Salmonella* gastroenteritis have a higher frequency of bacteremia than those not treated with opiates.

IMMUNITY

Both cellular immune responses and antibody production play important roles in protection from enteric infections. Humoral immunity to enteric pathogens consists of systemic IgG and IgM as well as secretory IgA. The mucosal immune system may be the first line of defense against many gastrointestinal pathogens. The binding of bacterial antigens to the luminal surface of M cells in the distal small bowel and the subsequent presentation of antigens to subepithelial lymphoid tissue lead to the proliferation of sensitized lymphocytes. These lymphocytes circulate and populate all of the mucosal tissues of the body as IgA-secreting plasma cells.

GENETIC DETERMINANTS

 Host genetic variation influences susceptibility to diarrheal diseases. People with blood group O show increased susceptibility to disease due to *V. cholerae*, *Shigella*, *E. coli* O157, and norovirus. Polymorphisms in genes encoding inflammatory mediators have been associated with the outcome of infection with enteroaggregative *E. coli*, enterotoxin-producing *E. coli*, *Salmonella*, *C. difficile*, and *V. cholerae*.

APPROACH TO THE PATIENT:

Infectious Diarrhea or Bacterial Food Poisoning

The approach to the patient with possible infectious diarrhea or bacterial food poisoning is shown in Fig. 160-1.

HISTORY

The answers to questions with high discriminating value can quickly narrow the range of potential causes of diarrhea and help determine whether treatment is needed. Important elements of the narrative history are detailed in Fig. 160-1.

PHYSICAL EXAMINATION

The examination of patients for signs of dehydration provides essential information about the severity of the diarrheal illness and the need for rapid therapy. Mild dehydration is indicated by thirst, dry mouth, decreased axillary sweat, decreased urine output, and slight weight loss. Signs of moderate dehydration include an orthostatic fall in blood pressure, skin tenting, and sunken eyes (or, in infants, a sunken fontanelle). Signs of severe dehydration include lethargy, obtundation, feeble pulse, hypotension, and frank shock.

DIAGNOSTIC APPROACH

After the severity of illness is assessed, the clinician must distinguish between *inflammatory* and *noninflammatory* disease. Using the history and epidemiologic features of the case as guides, the clinician

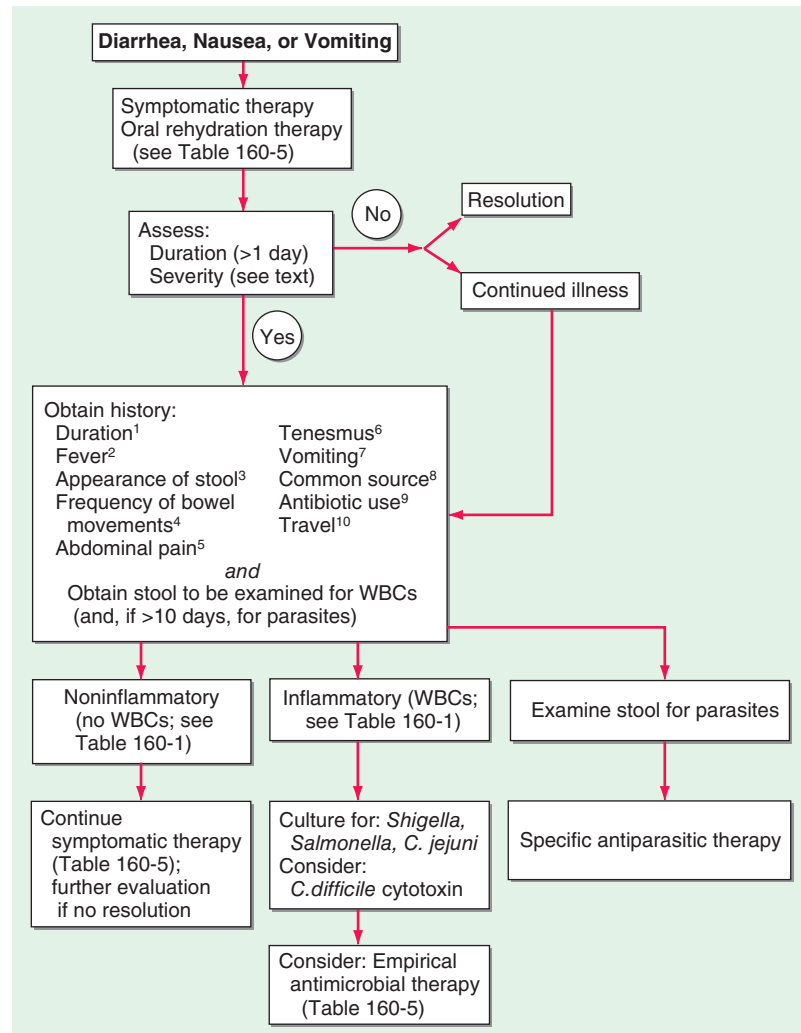


FIGURE 160-1 Clinical algorithm for the approach to patients with community-acquired infectious diarrhea or bacterial food poisoning.

Key to superscripts: 1. Diarrhea lasting >2 weeks is generally defined as chronic; in such cases, many of the causes of acute diarrhea are much less likely, and a new spectrum of causes needs to be considered. 2. Fever often implies invasive disease, although fever and diarrhea may also result from infection outside the gastrointestinal tract, as in malaria. 3. Stools that contain blood or mucus indicate ulceration of the large bowel. Bloody stools without fecal leukocytes should alert the laboratory to the possibility of infection with Shiga toxin-producing enterohemorrhagic *Escherichia coli*. Bulky white stools suggest a small-intestinal process that is causing malabsorption. Profuse “rice-water” stools suggest cholera or a similar toxigenic process. 4. Frequent stools over a given period can provide the first warning of impending dehydration. 5. Abdominal pain may be most severe in inflammatory processes like those due to *Shigella*, *Campylobacter*, and necrotizing toxins. Painful abdominal muscle cramps, caused by electrolyte loss, can develop in severe cases of cholera. Bloating is common in giardiasis. An appendicitis-like syndrome should prompt a culture for *Yersinia enterocolitica* with cold enrichment. 6. Tenesmus (painful rectal spasms with a strong urge to defecate but little passage of stool) may be a feature of cases with proctitis, as in shigellosis or amebiasis. 7. Vomiting implies an acute infection (e.g., a toxin-mediated illness or food poisoning) but can also be prominent in a variety of systemic illnesses (e.g., malaria) and in intestinal obstruction. 8. Asking patients whether anyone else they know is sick is a more efficient means of identifying a common source than is constructing a list of recently eaten foods. If a common source seems likely, specific foods can be investigated. See text for a discussion of bacterial food poisoning. 9. Current antibiotic therapy or a recent history of treatment suggests *Clostridium difficile* diarrhea (Chap. 161). Stop antibiotic treatment if possible and consider tests for *C. difficile* toxins. Antibiotic use may increase the risk of chronic intestinal carriage following salmonellosis. 10. See text (and Chap. 149) for a discussion of traveler’s diarrhea. (After TS Steiner, RL Guerrant: Principles and syndromes of enteric infection, in Mandell, Douglas, and Bennett’s Principles and Practice of Infectious Diseases, 7th ed, GL Mandell et al [eds]. Philadelphia, Churchill Livingstone, 2010, pp 1335–1351; RL Guerrant, DA Bobak: N Engl J Med 325:327, 1991; with permission.)

can then rapidly evaluate the need for further efforts to define a specific etiology and for therapeutic intervention. Examination of a stool sample may supplement the narrative history. Grossly bloody or mucoid stool suggests an inflammatory process. A test for fecal leukocytes (preparation of a thin smear of stool on a glass slide, addition of a drop of methylene blue, and examination of the wet mount) can suggest inflammatory disease in patients with diarrhea, although the predictive value of this test is still debated. A test for fecal lactoferrin, which is a marker of fecal leukocytes, is more

sensitive and is available in latex agglutination and enzyme-linked immunosorbent assay formats. Causes of acute infectious diarrhea, categorized as inflammatory and noninflammatory, are listed in Table 160-1.

POST-DIARRHEA COMPLICATIONS


Chronic complications may follow the resolution of an acute diarrheal episode. The clinician should inquire about prior diarrheal illness if the conditions listed in Table 160-2 are observed.

TABLE 160-2 POST-DIARRHEA COMPLICATIONS OF ACUTE INFECTIOUS DIARRHEAL ILLNESS

Complication	Comments
Chronic diarrhea	Occurs in ~1% of travelers with acute diarrhea
• Lactase deficiency	
• Small-bowel bacterial overgrowth	• Protozoa account for ~1/3 of cases
• Malabsorption syndromes (tropical and celiac sprue)	
Initial presentation or exacerbation of inflammatory bowel disease	May be precipitated by traveler's diarrhea
Irritable bowel syndrome	Occurs in ~10% of travelers with traveler's diarrhea
Reactive arthritis	Particularly likely after infection with invasive organisms (<i>Shigella</i> , <i>Salmonella</i> , <i>Campylobacter</i> , <i>Yersinia</i>)
Hemolytic-uremic syndrome (hemolytic anemia, thrombocytopenia, and renal failure)	Follows infection with Shiga toxin-producing bacteria (<i>Shigella dysenteriae</i> type 1 and enterohemorrhagic <i>Escherichia coli</i>)
Guillain-Barré syndrome	Particularly likely after <i>Campylobacter</i> infection

EPIDEMIOLOGY

TRAVEL HISTORY

 Of the several million people who travel from temperate industrialized countries to tropical regions of Asia, Africa, and Central and South America each year, 20–50% experience a sudden onset of abdominal cramps, anorexia, and watery diarrhea; thus *traveler's diarrhea* is the most common travel-related infectious illness (Chap. 149). The time of onset is usually 3 days to 2 weeks after the traveler's arrival in a resource-poor area; most cases begin within the first 3–5 days. The illness is generally self-limited, lasting 1–5 days. The high rate of diarrhea among travelers to underdeveloped areas is related to the ingestion of contaminated food or water.

The organisms that cause traveler's diarrhea vary considerably with location (Table 160-3), as does the pattern of antimicrobial resistance. In all areas, enterotoxigenic and enteroaggregative strains of *E. coli* are the most common isolates from persons with the classic secretory traveler's diarrhea syndrome. Infection with *Campylobacter jejuni* is especially common in areas of Asia.

LOCATION

Closed and semi-closed communities, including day-care centers, schools, residential facilities, and cruise ships, are important settings for outbreaks of enteric infections. Norovirus, which is highly contagious and robust in surviving on surfaces, is the most common etiologic agent associated with outbreaks of acute gastroenteritis. Other common organisms, often spread by fecal-oral contact in such communities, are *Shigella*, *C. jejuni*, and *Cryptosporidium*. Rotavirus is rarely a cause of pediatric diarrheal outbreaks in the United States since rotavirus vaccination was broadly recommended in 2006. Similarly, hospitals are sites in which enteric infections are concentrated. Diarrhea is one of the most common manifestations of nosocomial infections. *C. difficile* is the predominant cause of nosocomial diarrhea among adults in the United States, and outbreaks of norovirus infection are common in health care settings. *Klebsiella oxytoca* has been identified as a cause of antibiotic-associated hemorrhagic colitis. Enteropathogenic *E. coli* has been associated with outbreaks of diarrhea in nurseries for newborns. One-third of elderly patients in chronic-care institutions develop a significant diarrheal illness each year; more than one-half of these cases are caused by cytotoxin-producing *C. difficile*. Antimicrobial therapy can predispose to pseudomembranous colitis by altering the normal colonic flora and allowing the multiplication of *C. difficile* (Chap. 161).

TABLE 160-3 CAUSES OF TRAVELER'S DIARRHEA

Etiologic Agent	Approximate Percentage of Cases	Comments
Bacteria	50–75	
Enterotoxigenic <i>Escherichia coli</i>	10–45	Single most important agent
Enteroaggregative <i>E. coli</i>	5–35	Emerging enteric pathogen with worldwide distribution
<i>Campylobacter jejuni</i>	5–25	More common in Asia
<i>Shigella</i>	0–15	Major cause of dysentery
<i>Salmonella</i>	0–15	
Others	0–5	Including <i>Aeromonas</i> , <i>Plesiomonas</i> , and <i>Vibrio cholerae</i>
Viruses	0–20	
Norovirus	0–10	Associated with cruise ships
Rotavirus	0–5	Particularly common among children
Parasites	0–10	
<i>Giardia lamblia</i>	0–5	Affects hikers and campers who drink from freshwater streams
<i>Cryptosporidium</i>	0–5	Resistant to chlorine treatment
<i>Entamoeba histolytica</i>	<1	
<i>Cyclospora</i>	<1	
Other	0–10	
Acute food poisoning ^a	0–5	
No pathogen identified	10–50	

^aFor etiologic agents, see Table 160-4.

Source: After DR Hill et al: The practice of travel medicine: Guidelines by the Infectious Diseases Society of America. Clin Infect Dis 43:1499, 2006.

AGE

Globally, most morbidity and mortality from enteric pathogens involves children <5 years of age. Breast-fed infants are protected from contaminated food and water and derive some protection from maternal antibodies, but their risk of infection rises dramatically when they begin to eat solid foods. Exposure to rotavirus is universal, with most children experiencing their first infection in the first or second year of life if not vaccinated. Older children and adults are more commonly infected with norovirus. Other organisms with higher attack rates among children than among adults include enterotoxigenic, enteropathogenic, and enterohemorrhagic *E. coli*; *Shigella*; *C. jejuni*; and *G. lamblia*.

HOST IMMUNE STATUS

Immunocompromised hosts are at elevated risk of acute and chronic infectious diarrhea. Individuals with defects in cell-mediated immunity (including those with AIDS) are at particularly high risk of invasive enteropathies, including salmonellosis, listeriosis, and cryptosporidiosis. Individuals with hypogammaglobulinemia are at particular risk of *C. difficile* colitis and giardiasis. Patients with cancer are more likely to develop *C. difficile* infection as a result of chemotherapy and frequent hospitalizations. Infectious diarrhea can be life-threatening in immunocompromised hosts, with complications including bacteremia and metastatic seeding of infection. Furthermore, dehydration may compromise renal function and increase the toxicity of immunosuppressive drugs.

BACTERIAL FOOD POISONING

If the history and the stool examination indicate a noninflammatory etiology of diarrhea and there is evidence of a common-source outbreak, questions concerning the ingestion of specific foods and the time of onset of the diarrhea after a meal can provide clues to the bacterial cause of the illness. Potential causes of bacterial food poisoning are shown in Table 160-4.

Incubation Period, Organism	Symptoms	Common Food Sources
1–6 h		
<i>Staphylococcus aureus</i>	Nausea, vomiting, diarrhea	Ham, poultry, potato or egg salad, mayonnaise, cream pastries
<i>Bacillus cereus</i>	Nausea, vomiting, diarrhea	Fried rice
8–16 h		
<i>Clostridium perfringens</i>	Abdominal cramps, diarrhea (vomiting rare)	Beef, poultry, legumes, gravies
<i>B. cereus</i>	Abdominal cramps, diarrhea (vomiting rare)	Meats, vegetables, dried beans, cereals
>16 h		
<i>Vibrio cholerae</i>	Watery diarrhea	Shellfish, water
Enterotoxigenic <i>Escherichia coli</i>	Watery diarrhea	Salads, cheese, meats, water
Enterohemorrhagic <i>E. coli</i>	Bloody diarrhea	Ground beef, roast beef, salami, raw milk, raw vegetables, apple juice
<i>Salmonella</i> spp.	Inflammatory diarrhea	Beef, poultry, eggs, dairy products
<i>Campylobacter jejuni</i>	Inflammatory diarrhea	Poultry, raw milk
<i>Shigella</i> spp.	Dysentery	Potato or egg salad, lettuce, raw vegetables
<i>Vibrio parahaemolyticus</i>	Dysentery	Mollusks, crustaceans

Bacterial disease caused by an enterotoxin elaborated outside the host, such as that due to *Staphylococcus aureus* or *B. cereus*, has the shortest incubation period (1–6 h) and generally lasts <12 h. Most cases of staphylococcal food poisoning are caused by contamination from infected human carriers. Staphylococci can multiply at a wide range of temperatures; thus, if food is left to cool slowly and remains at room temperature after cooking, the organisms will have the opportunity to form enterotoxin. Outbreaks following picnics where potato salad, mayonnaise, and cream pastries have been served offer classic examples of staphylococcal food poisoning. Diarrhea, nausea, vomiting, and abdominal cramping are common, while fever is less so.

B. cereus can produce either a syndrome with a short incubation period—the *emetic* form, mediated by a staphylococcal type of enterotoxin—or one with a longer incubation period (8–16 h)—the *diarrheal* form, caused by an enterotoxin resembling *E. coli* LT, in which diarrhea and abdominal cramps are characteristic but vomiting is uncommon. The emetic form of *B. cereus* food poisoning is associated with contaminated fried rice; the organism is common in uncooked rice, and its heat-resistant spores survive boiling. If cooked rice is not refrigerated, the spores can germinate and produce toxin. Frying before serving may not destroy the preformed, heat-stable toxin.

Food poisoning due to *Clostridium perfringens* also has a slightly longer incubation period (8–14 h) and results from the survival of heat-resistant spores in inadequately cooked meat, poultry, or legumes. After ingestion, toxin is produced in the intestinal tract, causing moderately severe abdominal cramps and diarrhea; vomiting is rare, as is fever. The illness is self-limited, rarely lasting >24 h.

Not all food poisoning has a bacterial cause. Nonbacterial agents of short-incubation food poisoning include capsaicin, which is found in hot peppers, and a variety of toxins found in fish and shellfish (Chap. 474).

LABORATORY EVALUATION

Many cases of noninflammatory diarrhea are self-limited or can be treated empirically, and in these instances the clinician may not need to determine a specific etiology. Potentially pathogenic *E. coli* cannot be distinguished from normal fecal flora by routine culture, and tests to detect enterotoxins are not available in most clinical laboratories. In situations in which cholera is a concern, stool should be cultured on

selective media such as thiosulfate–citrate–bile salts–sucrose (TCBS) or tellurite–taurocholate–gelatin (TTG) agar. A latex agglutination test has made the rapid detection of rotavirus in stool practical for many laboratories, while reverse-transcriptase polymerase chain reaction (PCR) and specific antigen enzyme immunoassays have been developed for the identification of norovirus. Stool specimens should be examined by immunofluorescence-based rapid assays or (less sensitive) standard microscopy for *Giardia* cysts or *Cryptosporidium* if the level of clinical suspicion regarding the involvement of these organisms is high.

All patients with fever and evidence of inflammatory disease acquired outside the hospital should have stool cultured for *Salmonella*, *Shigella*, and *Campylobacter*. *Salmonella* and *Shigella* can be selected on MacConkey agar as non-lactose-fermenting (colorless) colonies or can be grown on *Salmonella-Shigella* agar or in selenite enrichment broth, both of which inhibit most organisms except these pathogens. Evaluation of nosocomial diarrhea should initially focus on *C. difficile*; stool culture for other pathogens in this setting has an extremely low yield and is not cost-effective. Toxins A and B produced by pathogenic strains of *C. difficile* can be detected by rapid enzyme immunoassays, latex agglutination tests, or PCR (Chap. 161). Isolation of *C. jejuni* requires inoculation of fresh stool onto selective growth medium and incubation at 42°C in a microaerophilic atmosphere. In many laboratories in the United States, *E. coli* O157:H7 is among the most common pathogens isolated from visibly bloody stools. Strains of this enterohemorrhagic serotype can be identified in specialized laboratories by serotyping but also can be identified presumptively in hospital laboratories as lactose-fermenting, indole-positive colonies of sorbitol nonfermenters (white colonies) on sorbitol MacConkey plates. If the clinical presentation suggests the possibility of intestinal amebiasis, stool should be examined by a rapid antigen detection assay or by (less sensitive and less specific) microscopy.

TREATMENT INFECTIOUS DIARRHEA OR BACTERIAL FOOD POISONING

In many cases, a specific diagnosis is not necessary or not available to guide treatment. The clinician can proceed with the information obtained from the history, stool examination, and evaluation of dehydration severity. Empirical regimens for the treatment of traveler's diarrhea are listed in Table 160-5.

The mainstay of treatment is adequate rehydration. The treatment of cholera and other dehydrating diarrheal diseases was revolutionized by the promotion of oral rehydration solution (ORS), the efficacy of which depends on the fact that glucose-facilitated absorption of sodium and water in the small intestine remains intact in the presence of cholera toxin. The use of ORS has reduced mortality rates for cholera from >50% (in untreated cases) to <1%. A number of ORS formulas have been used. Initial preparations were based on the treatment of patients with cholera and included a solution containing 3.5 g of sodium chloride, 2.5 g of sodium bicarbonate (or 2.9 g of sodium citrate), 1.5 g of potassium chloride, and 20 g of glucose (or 40 g of sucrose) per liter of water. Such a preparation can still be used for the treatment of severe cholera. Many causes of secretory diarrhea, however, are associated with less electrolyte loss than occurs in cholera. Beginning in 2002, the World Health Organization recommended a “reduced-osmolarity/reduced-salt” ORS that is better tolerated and more effective than classic ORS. This preparation contains 2.6 g of sodium chloride, 2.9 g of trisodium citrate, 1.5 g of potassium chloride, and 13.5 g of glucose (or 27 g of sucrose) per liter of water. ORS formulations containing rice or cereal as the carbohydrate source may be even more effective than glucose-based solutions. Patients who are severely dehydrated or in whom vomiting precludes the use of oral therapy should receive IV solutions such as Ringer's lactate.

Although most secretory forms of traveler's diarrhea (usually due to enterotoxigenic or enteroaggregative *E. coli* or to *Campylobacter*) can be treated effectively with rehydration, bismuth subsalicylate,

TABLE 160-5 TREATMENT OF TRAVELER'S DIARRHEA ON THE BASIS OF CLINICAL FEATURES*

Clinical Syndrome	Suggested Therapy
Watery diarrhea (no blood in stool, no fever), 1 or 2 unformed stools per day without distressing enteric symptoms	Oral fluids (oral rehydration solution, Pedialyte, Lytren, or flavored mineral water) and saltine crackers
Watery diarrhea (no blood in stool, no fever), 1 or 2 unformed stools per day with distressing enteric symptoms	Bismuth subsalicylate (for adults): 30 mL or 2 tablets (262 mg/tablet) every 30 min for 8 doses; or loperamide ^b : 4 mg initially followed by 2 mg after passage of each unformed stool, not to exceed 8 tablets (16 mg) per day (prescription dose) or 4 caplets (8 mg) per day (over-the-counter dose); drugs can be taken for 2 days
Watery diarrhea (no blood in stool, no distressing abdominal pain, no fever), >2 unformed stools per day	Antibacterial drug ^c plus (for adults) loperamide ^b (see dose above)
Dysentery (passage of bloody stools) or fever (>37.8°C)	Antibacterial drug ^c
Vomiting, minimal diarrhea	Bismuth subsalicylate (for adults; see dose above)
Diarrhea in infants (<2 years old)	Fluids and electrolytes (oral rehydration solution, Pedialyte, Lytren); continue feeding, especially with breast milk; seek medical attention for moderate dehydration, fever lasting >24 h, bloody stools, or diarrhea lasting more than several days

*All patients should take oral fluids (Pedialyte, Lytren, or flavored mineral water) plus saltine crackers. If diarrhea becomes moderate or severe, if fever persists, or if bloody stools or dehydration develops, the patient should seek medical attention. ^bLoperamide should not be used by patients with fever or dysentery; its use may prolong diarrhea in patients with infection due to *Shigella* or other invasive organisms. ^cThe recommended antibacterial drugs are as follows:

If the level of suspicion is low for fluoroquinolone-resistant *Campylobacter*:

Adults: (1) A fluoroquinolone such as ciprofloxacin, 750 mg as a single dose or 500 mg bid for 3 days; levofloxacin, 500 mg as a single dose or 500 mg qd for 3 days; or norfloxacin, 800 mg as a single dose or 400 mg bid for 3 days. (2) Azithromycin, 1000 mg as a single dose or 500 mg qd for 3 days. (3) Rifaximin, 200 mg tid or 400 mg bid for 3 days (not recommended for use in dysentery).

Children: Azithromycin, 10 mg/kg on day 1, 5 mg/kg on days 2 and 3 if diarrhea persists.

If fluoroquinolone-resistant *Campylobacter* is suspected (for example, following travel to Southeast Asia):

Adults: Azithromycin (at above dose for adults). *Children*: Same as for children traveling to other areas (see above).

Source: After DR Hill et al: The practice of travel medicine: Guidelines by the Infectious Diseases Society of America. *Clin Infect Dis* 43:1499, 2006.

or antiperistaltic agents, antimicrobial agents can shorten the duration of illness from 3–4 days to 24–36 h. Changes in diet have not been shown to have an impact on the duration of illness, while the efficacy of probiotics continues to be debated. Most individuals who present with dysentery (bloody diarrhea and fever) should be treated empirically with an antimicrobial agent (e.g., a fluoroquinolone or a macrolide) pending microbiologic analysis of stool. Individuals with shigellosis should receive a 3- to 7-day course. Individuals with *Campylobacter* infection often benefit from antimicrobial treatment as well. Because of widespread resistance of *Campylobacter* to fluoroquinolones, especially in parts of Asia, a macrolide antibiotic such as erythromycin or azithromycin may be preferred for this infection.

Treatment of salmonellosis must be tailored to the individual patient. Since administration of antimicrobial agents often prolongs intestinal colonization with *Salmonella*, these drugs are usually reserved for individuals at high risk of complications from disseminated salmonellosis, such as young children, patients with prosthetic devices, elderly patients, and immunocompromised persons. Antimicrobial agents should not be administered to individuals

(especially children) in whom enterohemorrhagic *E. coli* infection is suspected. Laboratory studies of enterohemorrhagic *E. coli* strains have demonstrated that a number of antibiotics induce replication of Shiga toxin-producing lambdoid bacteriophages, thereby significantly increasing toxin production by these strains. Clinical studies have supported these laboratory results, and antibiotics may increase by twentyfold the risk of hemolytic-uremic syndrome and renal failure during enterohemorrhagic *E. coli* infection. A clinical clue in the diagnosis of the latter infection is bloody diarrhea with low fever or none at all.

PROPHYLAXIS

Improvements in hygiene to limit fecal-oral spread of enteric pathogens will be necessary if the prevalence of diarrheal diseases is to be significantly reduced in developing countries. Travelers can reduce their risk of diarrhea by eating only hot, freshly cooked food; by avoiding raw vegetables, salads, and unpeeled fruit; and by drinking only boiled or treated water and avoiding ice. Historically, few travelers to tourist destinations adhere to these dietary restrictions. Bismuth subsalicylate is an inexpensive agent for the prophylaxis of traveler's diarrhea; it is taken at a dosage of 2 tablets (525 mg) four times a day. Treatment appears to be effective and safe for up to 3 weeks, but adverse events such as temporary darkening of the tongue and tinnitus can occur. A meta-analysis suggests that probiotics may lessen the likelihood of traveler's diarrhea by ~15%. Prophylactic antimicrobial agents, although effective, are not generally recommended for the prevention of traveler's diarrhea except when travelers are immunosuppressed or have other underlying illnesses that place them at high risk for morbidity from gastrointestinal infection. The risk of side effects and the possibility of developing an infection with a drug-resistant organism or with more harmful, invasive bacteria make it more reasonable to institute an empirical short course of treatment if symptoms develop. If prophylaxis is indicated, the nonabsorbed antibiotic rifaximin can be considered.

The possibility of exerting a major impact on the worldwide morbidity and mortality associated with diarrheal diseases has led to intense efforts to develop effective vaccines against the common bacterial and viral enteric pathogens. An effective rotavirus vaccine is currently available. Vaccines against *S. typhi* and *V. cholerae* also are available, although the protection they offer is incomplete and/or short lived. At present, there is no effective commercially available vaccine against *Shigella*, enterotoxigenic *E. coli*, *Campylobacter*, nontyphoidal *Salmonella*, norovirus, or intestinal parasites.

161 Clostridium difficile Infection, Including Pseudomembranous Colitis

Dale N. Gerding, Stuart Johnson

DEFINITION

Clostridium difficile infection (CDI) is a unique colonic disease that is acquired most often in association with antimicrobial use and the consequent disruption of the normal colonic microbiota. The most commonly diagnosed diarrheal illness acquired in the hospital, CDI results from the ingestion of spores of *C. difficile* that vegetate, multiply, and secrete toxins, causing diarrhea and pseudomembranous colitis (PMC) in the most severe cases.

ETIOLOGY AND EPIDEMIOLOGY

C. difficile is an obligately anaerobic, gram-positive, spore-forming bacillus whose spores are found widely in nature, particularly in the environment of hospitals and chronic-care facilities. CDI occurs

858 frequently in hospitals and nursing homes (or shortly after discharge from these facilities) where the level of antimicrobial use is high and the environment is contaminated by *C. difficile* spores.

Clindamycin, ampicillin, and cephalosporins were the first antibiotics associated with CDI. The second- and third-generation cephalosporins, particularly cefotaxime, ceftriaxone, cefuroxime, and ceftazidime, are frequently responsible for this condition, and the fluoroquinolones (ciprofloxacin, levofloxacin, and moxifloxacin) are the most recent drug class to be implicated in hospital outbreaks. Penicillin/ β -lactamase-inhibitor combinations such as ticarcillin/clavulanate and piperacillin/tazobactam pose significantly less risk. However, all antibiotics, including vancomycin and metronidazole (the agents most commonly used to treat CDI), carry a risk of subsequent CDI. Rare cases are reported in patients without prior antibiotic exposure.

C. difficile is acquired exogenously—most frequently in the hospital or nursing home, but also possibly in the outpatient setting—and is carried in the stool of both symptomatic and asymptomatic patients. The rate of fecal colonization is often $\geq 20\%$ among adult patients hospitalized for >1 week; in contrast, the rate is 1–3% among community residents. Community-onset CDI without recent hospitalization, nursing home residence, or outpatient health-care contact probably accounts for $\leq 10\%$ of all cases. The risk of *C. difficile* acquisition increases in proportion to the length of hospital stay. Asymptomatic fecal carriage of *C. difficile* in healthy neonates is very common, with repeated colonization by multiple strains in infants (<1 year old), but associated disease in these infants is extremely rare if it occurs at all. Spores of *C. difficile* are found on environmental surfaces (where the organism can persist for months) and on the hands of hospital personnel who fail to practice good hand hygiene. Hospital epidemics of CDI have been attributed to a single *C. difficile* strain and to multiple strains present simultaneously. Other identified risk factors for CDI include older age, greater severity of underlying illness, gastrointestinal surgery, use of electronic rectal thermometers, enteral tube feeding, and antacid treatment. Use of proton pump inhibitors may be a risk factor, but this risk is probably modest, and no firm data have implicated these agents in patients who are not already receiving antibiotics.

PATHOLOGY AND PATHOGENESIS

Spores of toxigenic *C. difficile* are ingested, survive gastric acidity, germinate in the small bowel, and colonize the lower intestinal tract, where they elaborate two large toxins: toxin A (an enterotoxin) and toxin B (a cytotoxin). These toxins initiate processes resulting in the disruption of epithelial-cell barrier function, diarrhea, and pseudomembrane formation. Toxin A is a potent neutrophil chemoattractant, and both toxins glucosylate the guanosine triphosphate (GTP)-binding proteins of the Rho subfamily that regulate the actin cell cytoskeleton. Data from studies using molecular disruption of toxin genes in isogenic mutants suggest that toxin B is the more important virulence factor. This possibility, if confirmed, might account for the occurrence of clinical disease caused by toxin A–negative strains. Disruption of the cytoskeleton results in loss of cell shape, adherence, and tight junctions, with consequent fluid leakage. A third toxin, binary toxin CDT, was previously found in only $\sim 6\%$ of strains but is present in all isolates of the widely recognized epidemic NAP1/BI/027 strain (see “Global Considerations,” below); this toxin is related to *C. perfringens* iota toxin. Its role in the pathogenesis of CDI has not yet been defined.

The pseudomembranes of PMC are confined to the colonic mucosa and initially appear as 1- to 2-mm whitish-yellow plaques. The intervening mucosa appears unremarkable, but, as the disease progresses, the pseudomembranes coalesce to form larger plaques and become confluent over the entire colon wall (Fig. 161-1). The whole colon is usually involved, but 10% of patients have rectal sparing. Viewed microscopically, the pseudomembranes have a mucosal attachment point and contain necrotic leukocytes, fibrin, mucus, and cellular debris. The epithelium is eroded and necrotic in focal areas, with neutrophil infiltration of the mucosa.

Patients colonized with *C. difficile* were initially thought to be at high risk for CDI. However, four prospective studies have shown that colonized patients who have not previously had CDI actually have a

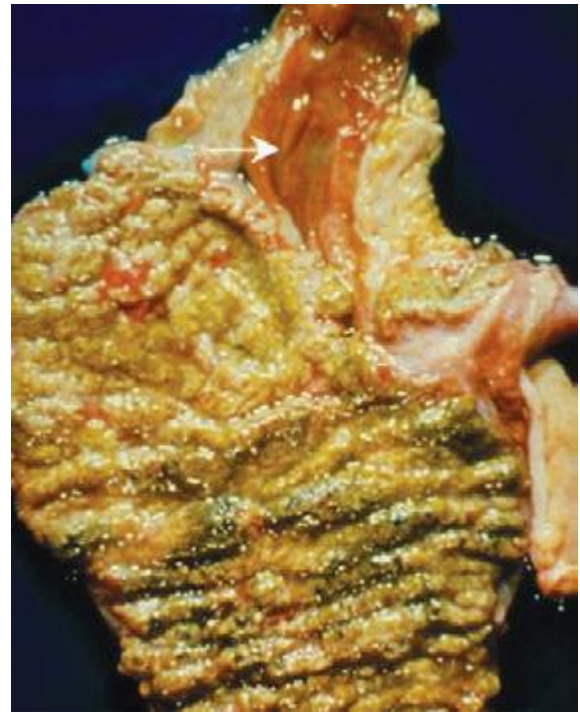


FIGURE 161-1 Autopsy specimen showing confluent pseudomembranes covering the cecum of a patient with pseudomembranous colitis. Note the sparing of the terminal ileum (arrow).

decreased risk of CDI. At least three events are proposed as essential for the development of CDI (Fig. 161-2). Exposure to antimicrobial agents is the first event and establishes susceptibility to *C. difficile* infection, most likely through disruption of the normal gastrointestinal microbiota. The second event is exposure to toxigenic *C. difficile*. Given that the majority of patients do not develop CDI after the first two events, a third event is clearly essential for its occurrence. Candidate third events include exposure to a *C. difficile* strain of particular virulence, exposure to antimicrobial agents especially likely

Pathogenesis model for *C. difficile* enteric disease

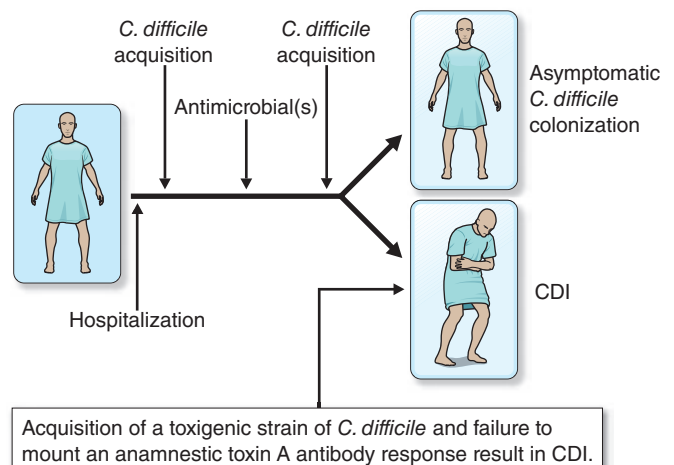


FIGURE 161-2 Pathogenesis model for hospital-acquired *Clostridium difficile* infection (CDI). At least three events are integral to *C. difficile* pathogenesis: (1) Exposure to antibiotics establishes susceptibility to infection. (2) Once susceptible, the patient may acquire nontoxigenic (nonpathogenic) or toxigenic strains of *C. difficile* as a second event. (3) Acquisition of toxigenic *C. difficile* may be followed by asymptomatic colonization or CDI, depending on one or more additional events (e.g., an inadequate host anamnestic IgG response to *C. difficile* toxin A).

to cause CDI, and an inadequate host immune response. The host anamnestic serum IgG antibody response to toxin A of *C. difficile* is the most likely third event that determines which patients develop diarrhea and which patients remain asymptomatic. In all probability, the majority of people first develop antibody to *C. difficile* toxins when colonized asymptotically during the first year of life or after CDI in childhood. Infants are thought not to develop symptomatic CDI because they lack suitable mucosal toxin receptors that develop later in life. In adulthood, serum levels of IgG antibody to toxin A increase more in response to infection in individuals who become asymptomatic carriers than in those who develop CDI. For persons who develop CDI, development of increasing levels of antitoxin A during treatment correlates with a lower risk of recurrence of CDI. A clinical trial using monoclonal antibodies to both toxin A and toxin B in addition to standard therapy showed rates of recurrence significantly lower than those obtained with placebo plus standard therapy.

GLOBAL CONSIDERATIONS



Rates and severity of CDI in the United States, Canada, and Europe increased markedly after the year 2000. Rates in U.S. hospitals tripled between 2000 and 2005. In 2005, hospitals in Montreal, Quebec, reported rates four times higher than the 1997 baseline, with directly attributable mortality of 6.9% (increased from 1.5%). An epidemic strain, variously known as toxinotype III, REA type BI, polymerase chain reaction (PCR) ribotype 027, and pulsed-field type NAP1 and thus collectively designated NAP1/BI/027, is thought to account for much of the increase in incidence and has been found in North America, Europe, and Asia. It is now recognized that two clones of NAP1/BI/027 originated in the United States and Canada and spread to the United Kingdom, Europe, and Asia. The epidemic organism is characterized by (1) an ability to produce 16–23 times as much toxin A and toxin B as control strains *in vitro*; (2) the presence of a third toxin (binary toxin CDT); and (3) high-level resistance to all fluoroquinolones. New strains have been and probably will continue to be implicated in outbreaks, including a strain (toxinotype V, ribotype 078) commonly found in food animals that also carries binary toxin and has been associated with high mortality risk in human infections. In the past 5 years, rates of CDI in the United Kingdom have markedly decreased, and the frequency of the NAP1/BI/027 strain in the countries of the European Union has likewise decreased. However, there has been no evidence of decreased rates of CDI or a decreased incidence of NAP1/BI/027 in North America; the latter strain still causes 25–35% of all CDIs in most regions of the United States.

CLINICAL MANIFESTATIONS

Diarrhea is the most common manifestation caused by *C. difficile*. Stools are almost never grossly bloody and range from soft and unformed to watery or mucoid in consistency, with a characteristic odor. Patients may have as many as 20 bowel movements per day.

Clinical and laboratory findings include fever in 28% of cases, abdominal pain in 22%, and leukocytosis in 50%. When adynamic ileus (which is seen on x-ray in ~20% of cases) results in cessation of stool passage, the diagnosis of CDI is frequently overlooked. A clue to the presence of unsuspected CDI in these patients is unexplained leukocytosis, with $\geq 15,000$ white blood cells (WBCs)/ μL . Such patients are at high risk for complications of *C. difficile* infection, particularly toxic megacolon and sepsis.

C. difficile diarrhea recurs after treatment in ~15–30% of cases, and this figure may be increasing. Recurrences may represent either relapses due to the same strain or reinfections with a new strain. Susceptibility to recurrence of clinical CDI is likely a result of continued fecal-microbiota disruption caused by the antibiotic used to treat CDI.

DIAGNOSIS

The diagnosis of CDI is based on a combination of clinical criteria: (1) diarrhea (≥ 3 unformed stools per 24 h for ≥ 2 days) with no other recognized cause plus (2) toxin A or B detected in the stool, toxin-producing *C. difficile* detected in the stool by PCR or culture, or pseudomembranes seen in the colon. PMC is a more advanced form of CDI and is visualized at endoscopy in only ~50% of patients with diarrhea who have a positive stool culture and toxin assay for *C. difficile*. Endoscopy is a rapid diagnostic tool in seriously ill patients with suspected PMC and an acute abdomen, but a negative result in this examination does not rule out CDI.

Despite the array of tests available for *C. difficile* and its toxins (Table 161-1), no single traditional test has high sensitivity, high specificity, and rapid turnaround. Most laboratory tests for toxins, including enzyme immunoassays, lack sensitivity. However, testing of multiple additional stool specimens is not recommended. Nucleic acid amplification tests, including PCR assays, have now been approved for diagnostic purposes and appear to be both rapid and sensitive while retaining high specificity. Testing of asymptomatic patients is not recommended except for epidemiologic study purposes. In particular, so-called tests of cure following treatment are not recommended because >50% of patients continue to harbor the organism and toxin after diarrhea has ceased and test results do not always predict recurrence of CDI. Thus these results should not be used to restrict placement of patients in long-term-care or nursing home facilities.

TREATMENT CLOSTRIDIUM DIFFICILE INFECTION

PRIMARY CDI

When possible, discontinuation of any ongoing antimicrobial administration is recommended as the first step in treatment of CDI. Earlier studies indicated that 15–23% of patients respond to this simple measure. However, with the advent of the current epidemic strain and the associated rapid clinical deterioration of some patients,

TABLE 161-1 RELATIVE SENSITIVITY AND SPECIFICITY OF DIAGNOSTIC TESTS FOR *CLOSTRIDIUM DIFFICILE* INFECTION (CDI)

Type of Test	Relative Sensitivity ^a	Relative Specificity ^a	Comment
Stool culture for <i>C. difficile</i>	++++	+++	Most sensitive test; specificity of ++++ if the <i>C. difficile</i> isolate tests positive for toxin; with clinical data, is diagnostic of CDI; turnaround time too slow for practical use
Cell culture cytotoxin test on stool	+++	++++	With clinical data, is diagnostic of CDI; highly specific but not as sensitive as stool culture; slow turnaround time
Enzyme immunoassay for toxin A or toxins A and B in stool	++ to +++	+++	With clinical data, is diagnostic of CDI; rapid results, but not as sensitive as stool culture or cell culture cytotoxin test
Enzyme immunoassay for <i>C. difficile</i> common antigen in stool	+++ to ++++	+++	Detects glutamate dehydrogenase found in toxigenic and non-toxigenic strains of <i>C. difficile</i> and other stool organisms; more sensitive and less specific than enzyme immunoassay for toxins; rapid results
Nucleic acid amplification tests for <i>C. difficile</i> toxin A or B gene in stool	++++	++++	Detect toxigenic <i>C. difficile</i> in stool; newly approved for clinical testing, but appears to be more sensitive than enzyme immunoassay toxin testing and at least as specific
Colonoscopy or sigmoidoscopy	+	++++	Highly specific if pseudomembranes are seen; insensitive compared with other tests

^aAccording to both clinical and test-based criteria. +++++, >90%; +++, 71–90%; ++, 51–70%; +, ~50%.

860 prompt initiation of specific CDI treatment has become the standard. Empirical treatment is appropriate if CDI is strongly suspected on clinical grounds. General treatment guidelines include hydration and the avoidance of antiperistaltic agents and opiates, which may mask symptoms and possibly worsen disease. Nevertheless, antiperistaltic agents have been used safely with vancomycin or metronidazole for mild to moderate CDI.

Oral administration of vancomycin, fidaxomicin, or metronidazole is recommended for CDI treatment. IV vancomycin is ineffective for CDI, and fidaxomicin is available only for oral administration; when IV metronidazole is administered, fecal bactericidal drug concentrations are achieved during acute diarrhea; however, in the presence of adynamic ileus, IV metronidazole treatment of CDI has failed. Two large clinical trials comparing vancomycin and fidaxomicin indicated comparable resolution of diarrhea (~90% of patients) as well as significantly reduced rates of recurrent CDI with fidaxomicin from rates with vancomycin. In previous randomized trials, diarrhea response rates to oral therapy with vancomycin or metronidazole were ≥94%, but four observational studies found that response rates for metronidazole had declined to 62–78%. Although the mean time to resolution of diarrhea is 2–4 days, the response to metronidazole may be much slower. Treatment should not be deemed a failure until a drug has been given for at least 6 days. On the basis of data for shorter courses of vancomycin and the results of two large-scale clinical trials, it is recommended that vancomycin, fidaxomicin, and metronidazole be given for at least 10 days. Metronidazole is not approved for CDI by the U.S. Food and Drug Administration (FDA), but most patients with mild to moderate illness respond to 500 mg given by mouth three times a day for 10 days; extension of the treatment period may be needed for slow responders. In addition to the reports of increases in metronidazole failures, a prospective, randomized, double-blind, placebo-controlled study has demonstrated the superiority of vancomycin over metronidazole for treatment of severe CDI. The severity assessment score in that study included age as well as laboratory parameters (elevated temperature, low albumin level, or elevated WBC count), documentation of PMC by endoscopy, and treatment of CDI in the intensive care unit. Although a validated severity score is not available, it is important to initiate treatment with oral vancomycin for patients

who appear seriously ill, particularly if they have a high WBC count (>15,000/μL) or a creatinine level that is ≥1.5 times higher than the premorbid value (Table 161-2). In addition, a randomized blinded trial compared a toxin-binding polymer, tolevamer, with two antibiotic regimens for treatment of CDI and showed that vancomycin was superior to metronidazole for all patients regardless of severity. Small randomized trials of nitazoxanide, bacitracin, rifaximin, and fusidic acid for treatment of CDI have been conducted. These drugs have not been extensively studied, shown to be superior, or approved by the FDA for CDI, but they provide potential alternatives to vancomycin, fidaxomicin, and metronidazole.

RECURRENT CDI

Overall, ~15–30% of successfully treated patients experience recurrences of CDI, either as relapses caused by the original organism or as reinfections following treatment. Rates of CDI recurrence are significantly lower among patients treated with fidaxomicin rather than vancomycin. Rates of recurrence are comparable with vancomycin and metronidazole. Recurrence rates are higher among patients ≥65 years old, those who continue to take antibiotics while being treated for CDI, and those who remain in the hospital after the initial episode of CDI. Patients who have a first recurrence of CDI have a high rate of second recurrence (33–65%). In the first recurrence, re-treatment with metronidazole is comparable to treatment with vancomycin (Table 161-2), and fidaxomicin is superior to vancomycin in reducing the risk of further recurrences in patients who have had one recurrence. Recurrent CDI, once thought to be relatively mild, has now been documented to pose a significant (11%) risk of serious complications (shock, megacolon, perforation, colectomy, or death within 30 days). There is no standard treatment for multiple recurrences, but long or repeated metronidazole courses should be avoided because of potential neurotoxicity. The use of vancomycin in tapering doses or with pulse dosing every other day for 2–8 weeks may be the most practical approach to treatment of patients with multiple recurrences. Other approaches include the administration of vancomycin followed by the yeast *Saccharomyces boulardii*; the administration of vancomycin followed by a fecal microbiota transplant given via nasoduodenal tube, colonoscopy, or enema; and the intentional colonization of the patient with a nontoxigenic

TABLE 161-2 RECOMMENDATIONS FOR THE TREATMENT OF *CLOSTRIDIUM DIFFICILE* INFECTION (CDI)^a

Clinical Setting	Treatment(s)	Comments
Initial episode, mild to moderate	Metronidazole (500 mg tid × 10–14 d)	Vancomycin (125 mg qid × 10–14 d) may be more effective than metronidazole. Fidaxomicin (200 mg bid × 10 d) is another alternative.
Initial episode, severe	Vancomycin (125 mg qid × 10–14 d)	Indicators of severe disease may include leukocytosis (≥15,000 white blood cells/μL) and a creatinine level ≥1.5 times the premorbid value. Fidaxomicin is an alternative.
Initial episode, severe complicated or fulminant	Vancomycin (500 mg PO or via nasogastric tube) plus metronidazole (500 mg IV q8h) plus consider Rectal instillation of vancomycin (500 mg in 100 mL of normal saline as a retention enema q6–8h)	Severe complicated or fulminant CDI is defined as severe CDI with the addition of hypotension, shock, ileus, or toxic megacolon. The duration of treatment may need to be >2 weeks and is dictated by response. Consider using tigecycline (50 mg IV q12h after a 100-mg loading dose) in place of metronidazole.
First recurrence	Same as for initial episode	Adjust treatment if severity of CDI has changed with recurrence. Consider fidaxomicin, which significantly decreases the likelihood of additional recurrences.
Second recurrence	Vancomycin in taper/pulse regimen	Typical taper/pulse regimen: 125 mg qid × 10–14 d, then bid × 1 week, then daily × 1 week, then q2–3d for 2–8 weeks.
Multiple recurrences	Consider the following options: <ul style="list-style-type: none"> • Repeat vancomycin taper/pulse • Vancomycin (500 mg qid × 10 d) plus <i>Saccharomyces boulardii</i> (500 mg bid × 28 d) • Vancomycin (125 mg qid × 10–14 d); then stop vancomycin and start rifaximin (400 mg bid × 2 weeks) • Nitazoxanide (500 mg bid × 10 d) • Fecal microbiota transplantation • IV immunoglobulin (400 mg/kg) 	The only controlled studies that included patients with one or more recurrent CDI episodes were with vancomycin and <i>S. boulardii</i> , which showed borderline significance compared with vancomycin plus placebo, and fecal microbiota transplantation, which was highly significant compared with a high-dose course of vancomycin. (The vancomycin taper was not compared.)

^aAll agents are given orally unless otherwise specified.

162 Urinary Tract Infections, Pyelonephritis, and Prostatitis

Kalpana Gupta, Barbara W. Trautner

strain of *C. difficile*. None of these biotherapeutic approaches has been approved by the FDA for use in the United States. Other non-FDA-approved antibiotic strategies include (1) sequential treatment with vancomycin (125 mg four times daily for 10–14 days) followed by rifaximin (400 mg twice daily for 14 days) and (2) treatment with nitazoxanide (500 mg twice daily for 7 days). IV immunoglobulin, which has also been used with variable success, presumably provides antibodies to *C. difficile* toxins.

SEVERE COMPLICATED OR FULMINANT CDI

Fulminant (rapidly progressive and severe) CDI presents the most difficult treatment challenge. Patients with fulminant disease often do not have diarrhea, and their illness mimics an acute surgical abdomen. Sepsis (hypotension, fever, tachycardia, leukocytosis) may result from severe CDI. An acute abdomen (with or without toxic megacolon) may include signs of obstruction, ileus, colon-wall thickening and ascites on abdominal CT, and peripheral-blood leukocytosis ($\geq 20,000$ WBCs/ μ L). With or without diarrhea, the differential diagnosis of an acute abdomen, sepsis, or toxic megacolon should include CDI if the patient has received antibiotics in the past 2 months. Cautious sigmoidoscopy or colonoscopy to visualize PMC and abdominal CT are the best diagnostic tests in patients without diarrhea.

Medical management of fulminant CDI is suboptimal because of the difficulty of delivering oral fidaxomicin, metronidazole, or vancomycin to the colon in the presence of ileus (Table 161-2). The combination of vancomycin (given via nasogastric tube and by retention enema) plus IV metronidazole has been used with some success in uncontrolled studies, as has IV tigecycline in small-scale uncontrolled studies. Surgical colectomy may be life-saving if there is no response to medical management. If possible, colectomy should be performed before the serum lactate level reaches 5 mmol/L. The incidence of fulminant CDI requiring colectomy appears to be increasing in the evolving epidemic; however, morbidity and death associated with colectomy may be reduced by performing instead a laparoscopic ileostomy followed by colon lavage with polyethylene glycol and vancomycin infusion into the colon via the ileostomy.

PROGNOSIS

The mortality rate attributed to CDI, previously found to be 0.6–3.5%, has reached 6.9% in recent outbreaks and rises progressively with increasing age. Most patients recover, but recurrences are common.

PREVENTION AND CONTROL

Strategies for the prevention of CDI are of two types: those aimed at preventing transmission of the organism to the patient and those aimed at reducing the risk of CDI if the organism is transmitted. Transmission of *C. difficile* in clinical practice has been prevented by gloving of personnel, elimination of the use of contaminated electronic thermometers, and use of hypochlorite (bleach) solution for environmental decontamination of patients' rooms. Hand hygiene is critical; hand washing is recommended in CDI outbreaks because alcohol hand gels are not sporicidal. CDI outbreaks have been best controlled by restricting the use of specific antibiotics, such as clindamycin and second- and third-generation cephalosporins. Outbreaks of CDI due to clindamycin-resistant strains have resolved promptly when clindamycin use is restricted. Future preventive strategies are likely to include use of monoclonal antibodies, vaccines, and biotherapeutics containing live organisms that will restore colonization protection in the microbiota.

Urinary tract infection (UTI) is a common and painful human illness that, fortunately, is rapidly responsive to modern antibiotic therapy. In the preantibiotic era, UTI caused significant morbidity. Hippocrates, writing about a disease that appears to have been acute cystitis, said that the illness could last for a year before either resolving or worsening to involve the kidneys. When chemotherapeutic agents used to treat UTI were introduced in the early twentieth century, they were relatively ineffective, and persistence of infection after 3 weeks of therapy was common. Nitrofurantoin, which became available in the 1950s, was the first tolerable and effective agent for the treatment of UTI.

Since the most common manifestation of UTI is acute cystitis and since acute cystitis is far more prevalent among women than among men, most clinical research on UTI has involved women. Many studies have enrolled women from college campuses or large health maintenance organizations in the United States. Therefore, when reviewing the literature and recommendations concerning UTI, clinicians must consider whether the findings are applicable to their patient populations.

DEFINITIONS

UTI may be asymptomatic (subclinical infection) or symptomatic (disease). Thus, the term *urinary tract infection* encompasses a variety of clinical entities, including asymptomatic bacteriuria (ASB), cystitis, prostatitis, and pyelonephritis. The distinction between symptomatic UTI and ASB has major clinical implications. Both UTI and ASB connote the presence of bacteria in the urinary tract, usually accompanied by white blood cells and inflammatory cytokines in the urine. However, ASB occurs in the absence of symptoms attributable to the bacteria in the urinary tract and does not usually require treatment, while UTI has more typically been assumed to imply symptomatic disease that warrants antimicrobial therapy. Much of the literature concerning UTI, particularly catheter-associated infection, does not differentiate between UTI and ASB. In this chapter, the term *UTI* denotes symptomatic disease; *cystitis*, symptomatic infection of the bladder; and *pyelonephritis*, symptomatic infection of the kidneys. *Uncomplicated UTI* refers to acute cystitis or pyelonephritis in nonpregnant outpatient women without anatomic abnormalities or instrumentation of the urinary tract; the term *complicated UTI* encompasses all other types of UTI. *Recurrent UTI* is not necessarily complicated; individual episodes can be uncomplicated and treated as such. *Catheter-associated bacteriuria* can be either symptomatic (CAUTI) or asymptomatic.

EPIDEMIOLOGY AND RISK FACTORS

Except among infants and the elderly, UTI occurs far more commonly in females than in males. During the neonatal period, the incidence of UTI is slightly higher among males than among females because male infants more commonly have congenital urinary tract anomalies. After 50 years of age, obstruction from prostatic hypertrophy becomes common in men, and the incidence of UTI is almost as high among men as among women. Between 1 year and ~50 years of age, UTI and recurrent UTI are predominantly diseases of females. The prevalence of ASB is ~5% among women between ages 20 and 40 and may be as high as 40–50% among elderly women and men.

As many as 50–80% of women in the general population acquire at least one UTI during their lifetime—uncomplicated cystitis in most cases. Recent use of a diaphragm with spermicide, frequent sexual intercourse, and a history of UTI are independent risk factors for acute cystitis. Cystitis is temporally related to recent sexual intercourse in a dose-response manner, with an increased relative risk ranging from 1.4 with one episode of intercourse to 4.8 with five episodes of intercourse in the preceding week. In healthy postmenopausal women, sexual activity, diabetes mellitus, and incontinence are risk factors for UTI.

Many factors predisposing women to cystitis also increase the risk of pyelonephritis. Factors independently associated with pyelonephritis in young healthy women include frequent sexual intercourse, a new sexual partner, a UTI in the previous 12 months, a maternal history of UTI, diabetes, and incontinence. The common risk factors for cystitis and pyelonephritis are not surprising given that pyelonephritis typically arises through the ascent of bacteria from the bladder to the upper urinary tract. However, pyelonephritis can occur without clear antecedent cystitis.

About 20–30% of women who have had one episode of UTI will have recurrent episodes. Early recurrence (within 2 weeks) is usually regarded as relapse rather than reinfection and may indicate the need to evaluate the patient for a sequestered focus. Intracellular pods of infecting organisms within the bladder epithelium have been demonstrated in animal models of UTI, but the importance of this phenomenon in humans is not yet clear. The rate of recurrence ranges from 0.3 to 7.6 infections per patient per year, with an average of 2.6 infections per year. It is not uncommon for multiple recurrences to follow an initial infection, resulting in clustering of episodes. Clustering may be related temporally to the presence of a new risk factor or to the sloughing of the protective outer bladder epithelial layer in response to bacterial attachment during acute cystitis. The likelihood of a recurrence decreases with increasing time since the last infection. A case-control study of predominantly white premenopausal women with recurrent UTI identified frequent sexual intercourse, use of spermicide, a new sexual partner, a first UTI before 15 years of age, and a maternal history of UTI as independent risk factors for recurrent UTI. The only consistently documented behavioral risk factors for recurrent UTI include frequent sexual intercourse and spermicide use. In postmenopausal women, major risk factors for recurrent UTI include a history of premenopausal UTI and anatomic factors affecting bladder emptying, such as cystoceles, urinary incontinence, and residual urine.

In pregnant women, ASB has clinical consequences, and both screening for and treatment of this condition are indicated. Specifically, ASB during pregnancy is associated with preterm birth and perinatal death of the fetus and with pyelonephritis in the mother. A Cochrane meta-analysis found that treatment of ASB in pregnant women decreased the risk of pyelonephritis by 75%.

The majority of men with UTI have a functional or anatomic abnormality of the urinary tract, most commonly urinary obstruction secondary to prostatic hypertrophy. That said, not all men with UTI have detectable urinary abnormalities; this point is particularly relevant for men ≤ 45 years of age. Lack of circumcision is also associated with an increased risk of UTI because *Escherichia coli* is more likely to colonize the glans and prepuce and subsequently migrate into the urinary tract.

Women with diabetes have been found to have a two- to threefold higher rate of ASB and UTI than women without diabetes; there is insufficient evidence to make a corresponding statement about men. Increased duration of diabetes and the use of insulin rather than oral medication are also associated with a higher risk of UTI among women with diabetes. Poor bladder function, obstruction in urinary flow, and incomplete voiding are additional factors commonly found in patients with diabetes that increase the risk of UTI. Impaired cytokine secretion may contribute to ASB in diabetic women.

ETIOLOGY



The uropathogens causing UTI vary by clinical syndrome but are usually enteric gram-negative rods that have migrated to the urinary tract. The susceptibility patterns of these organisms vary by clinical syndrome and by geography. In acute uncomplicated cystitis in the United States, the etiologic agents are highly predictable: *E. coli* accounts for 75–90% of isolates; *Staphylococcus saprophyticus* for 5–15% (with particularly frequent isolation from younger women); and *Klebsiella*, *Proteus*, *Enterococcus*, and *Citrobacter* species, along with other organisms, for 5–10%. Similar etiologic agents are found in Europe and Brazil. The spectrum of agents causing uncomplicated pyelonephritis is similar, with *E. coli* predominating. In complicated UTI (e.g., CAUTI), *E. coli* remains the predominant organism, but other aerobic gram-negative rods, such as *Pseudomonas*

aeruginosa and *Klebsiella*, *Proteus*, *Citrobacter*, *Acinetobacter*, and *Morganella* species, also are frequently isolated. Gram-positive bacteria (e.g., enterococci and *Staphylococcus aureus*) and yeasts are also important pathogens in complicated UTI. Data on etiology and resistance are generally obtained from laboratory surveys and should be understood in the context that organism identification is performed only in cases in which urine is sent for culture—i.e., typically, when complicated UTI or pyelonephritis is suspected. The available data demonstrate a worldwide increase in the resistance of *E. coli* to antibiotics commonly used to treat UTI. North American and European surveys from women with acute cystitis have documented resistance rates of >20% to trimethoprim-sulfamethoxazole (TMP-SMX) and to ciprofloxacin in some regions. In community-acquired infections, the increased prevalence of uropathogens producing extended-spectrum β -lactamases has left few oral options for therapy. Since resistance rates vary by local geographic region, with individual patient characteristics, and over time, it is important to use current and local data when choosing a treatment regimen.

PATHOGENESIS

The urinary tract can be viewed as an anatomic unit united by a continuous column of urine extending from the urethra to the kidneys. In the majority of UTIs, bacteria establish infection by ascending from the urethra to the bladder. Continuing ascent up the ureter to the kidney is the pathway for most renal parenchymal infections. However, introduction of bacteria into the bladder does not inevitably lead to sustained and symptomatic infection. The interplay of host, pathogen, and environmental factors determines whether tissue invasion and symptomatic infection will ensue (Fig. 162-1). For example, bacteria often enter the bladder after sexual intercourse, but normal voiding and innate host defense mechanisms in the bladder eliminate these organisms. Any foreign body in the urinary tract, such as a urinary catheter or stone, provides an inert surface for bacterial colonization. Abnormal micturition and/or significant residual urine volume promotes true infection. In the simplest of terms, anything that increases the likelihood of bacteria entering the bladder and staying there increases the risk of UTI.

Bacteria can also gain access to the urinary tract through the bloodstream. However, hematogenous spread accounts for <2% of documented UTIs and usually results from bacteremia caused by relatively virulent organisms, such as *Salmonella* and *S. aureus*. Indeed, the isolation of either of these pathogens from a patient without a catheter or other instrumentation warrants a search for a bloodstream source. Hematogenous infections may produce focal abscesses or areas of pyelonephritis within a kidney and result in positive urine cultures. The pathogenesis of candiduria is distinct in that the hematogenous route is common. The presence of *Candida* in the urine of a noninstrumented

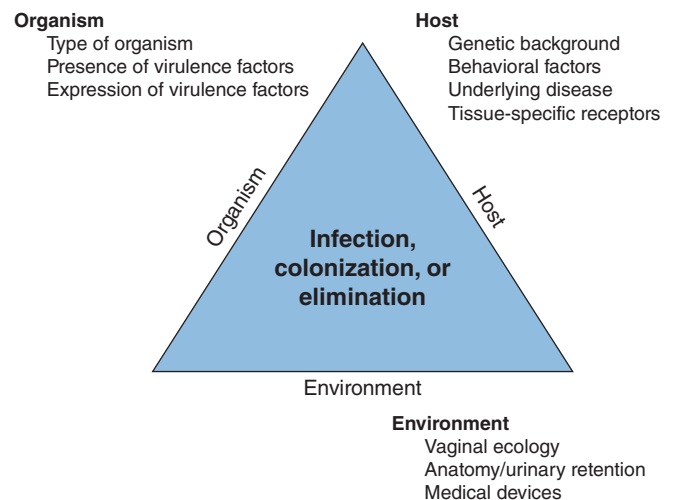


FIGURE 162-1 Pathogenesis of urinary tract infection. The relationship among specific host, pathogen, and environmental factors determines the clinical outcome.

immunocompetent patient implies either genital contamination or potentially widespread visceral dissemination.

Environmental Factors • VAGINAL ECOLOGY In women, vaginal ecology is an important environmental factor affecting the risk of UTI. Colonization of the vaginal introitus and periurethral area with organisms from the intestinal flora (usually *E. coli*) is the critical initial step in the pathogenesis of UTI. Sexual intercourse is associated with an increased risk of vaginal colonization with *E. coli* and thereby increases the risk of UTI. Nonoxynol-9 in spermicide is toxic to the normal vaginal microflora and thus is likewise associated with an increased risk of *E. coli* vaginal colonization and bacteriuria. In postmenopausal women, the previously predominant vaginal lactobacilli are replaced with colonizing gram-negative bacteria. The use of topical estrogens to prevent UTI in postmenopausal women is controversial; given the side effects of systemic hormone replacement, oral estrogens should not be used to prevent UTI.

ANATOMIC AND FUNCTIONAL ABNORMALITIES Any condition that permits urinary stasis or obstruction predisposes the individual to UTI. Foreign bodies such as stones or urinary catheters provide an inert surface for bacterial colonization and formation of a persistent biofilm. Thus, vesicoureteral reflux, ureteral obstruction secondary to prostatic hypertrophy, neurogenic bladder, and urinary diversion surgery create an environment favorable to UTI. In persons with such conditions, *E. coli* strains lacking typical urinary virulence factors are often the cause of infection. Inhibition of ureteral peristalsis and decreased ureteral tone leading to vesicoureteral reflux are important in the pathogenesis of pyelonephritis in pregnant women. Anatomic factors—specifically, the distance of the urethra from the anus—are considered to be the primary reason why UTI is predominantly an illness of young women rather than of young men.



Host Factors The genetic background of the host influences the individual's susceptibility to recurrent UTI, at least among women. A familial disposition to UTI and to pyelonephritis is well documented. Women with recurrent UTI are more likely to have had their first UTI before the age of 15 years and to have a maternal history of UTI. A component of the underlying pathogenesis of this familial predisposition to recurrent UTI may be persistent vaginal colonization with *E. coli*, even during asymptomatic periods. Vaginal and periurethral mucosal cells from women with recurrent UTI bind threefold more uropathogenic bacteria than do mucosal cells from women without recurrent infection. Epithelial cells from women who are non-secretors of certain blood group antigens may possess specific types of receptors to which *E. coli* can bind, thereby facilitating colonization and invasion. Mutations in host response genes (e.g., those coding for Toll-like receptors and the interleukin 8 receptor) also have been linked to recurrent UTI and pyelonephritis. Polymorphisms in the interleukin 8–specific receptor gene *CXCR1* are associated with increased susceptibility to pyelonephritis. Lower-level expression of *CXCR1* on the surface of neutrophils impairs neutrophil-dependent host defense against bacterial invasion of the renal parenchyma.



Microbial Factors An anatomically normal urinary tract presents a stronger barrier to infection than a compromised urinary tract. Thus, strains of *E. coli* that cause invasive symptomatic infection of the urinary tract in otherwise normal hosts often possess and express genetic virulence factors, including surface adhesins that mediate binding to specific receptors on the surface of uroepithelial cells. The best-studied adhesins are the P fimbriae, hairlike protein structures that interact with a specific receptor on renal epithelial cells. (The letter P denotes the ability of these fimbriae to bind to blood group antigen P, which contains a D-galactose-D-galactose residue.) P fimbriae are important in the pathogenesis of pyelonephritis and subsequent bloodstream invasion from the kidney.

Another adhesin is the type 1 pilus (fimbria), which all *E. coli* strains possess but not all *E. coli* strains express. Type 1 pili are thought to play a key role in initiating *E. coli* bladder infection; they mediate binding to uroplakins on the luminal surface of bladder uroepithelial cells. The binding of type 1 fimbriae of *E. coli* to receptors on uroepithelial cells

initiates a complex series of signaling events that leads to apoptosis and exfoliation of uroepithelial cells, with the attached *E. coli* organisms carried away in the urine.

APPROACH TO THE PATIENT: Clinical Syndromes

The most important issue to be addressed when a UTI is suspected is the characterization of the clinical syndrome as ASB, uncomplicated cystitis, pyelonephritis, prostatitis, or complicated UTI. This information will shape the diagnostic and therapeutic approach.

ASYMPTOMATIC BACTERIURIA

A diagnosis of ASB can be considered only when the patient does not have local or systemic symptoms referable to the urinary tract. The clinical presentation is usually that of a patient who undergoes a screening urine culture for a reason unrelated to the genitourinary tract and is incidentally found to have bacteriuria. The presence of systemic signs or symptoms such as fever, altered mental status, and leukocytosis in the setting of a positive urine culture does not merit a diagnosis of symptomatic UTI unless other potential etiologies have been considered.

CYSTITIS

The typical symptoms of cystitis are dysuria, urinary frequency, and urgency. Nocturia, hesitancy, suprapubic discomfort, and gross hematuria are often noted as well. Unilateral back or flank pain is generally an indication that the upper urinary tract is involved. Fever also is an indication of invasive infection of either the kidney or the prostate.

PYELONEPHRITIS

Mild pyelonephritis can present as low-grade fever with or without lower-back or costovertebral-angle pain, whereas severe pyelonephritis can manifest as high fever, rigors, nausea, vomiting, and flank and/or loin pain. Symptoms are generally acute in onset, and symptoms of cystitis may not be present. Fever is the main feature distinguishing cystitis and pyelonephritis. The fever of pyelonephritis typically exhibits a high spiking “picket-fence” pattern and resolves over 72 h of therapy. Bacteremia develops in 20–30% of cases of pyelonephritis. Patients with diabetes may present with obstructive uropathy associated with acute papillary necrosis when the sloughed papillae obstruct the ureter. Papillary necrosis may also be evident in some cases of pyelonephritis complicated by obstruction, sickle cell disease, analgesic nephropathy, or combinations of these conditions. In the rare cases of bilateral papillary necrosis, a rapid rise in the serum creatinine level may be the first indication of the condition. *Emphysematous* pyelonephritis is a particularly severe form of the disease that is associated with the production of gas in renal and perinephric tissues and occurs almost exclusively in diabetic patients (Fig. 162-2). *Xanthogranulomatous* pyelonephritis occurs when chronic urinary obstruction (often by staghorn calculi), together with chronic infection, leads to suppurative destruction of renal tissue (Fig. 162-3). On pathologic examination, the residual renal tissue frequently has a yellow coloration, with infiltration by lipid-laden macrophages. Pyelonephritis can also be complicated by intraparenchymal abscess formation; this situation should be suspected when a patient has continued fever and/or bacteremia despite antibacterial therapy.

PROSTATITIS

Prostatitis includes both infectious and noninfectious abnormalities of the prostate gland. Infections can be acute or chronic, are almost always bacterial in nature, and are far less common than the noninfectious entity *chronic pelvic pain syndrome* (formerly known as chronic prostatitis). Acute bacterial prostatitis presents as dysuria, frequency, and pain in the prostatic pelvic or perineal area. Fever and chills are usually present, and symptoms of bladder outlet obstruction are common. Chronic bacterial prostatitis presents

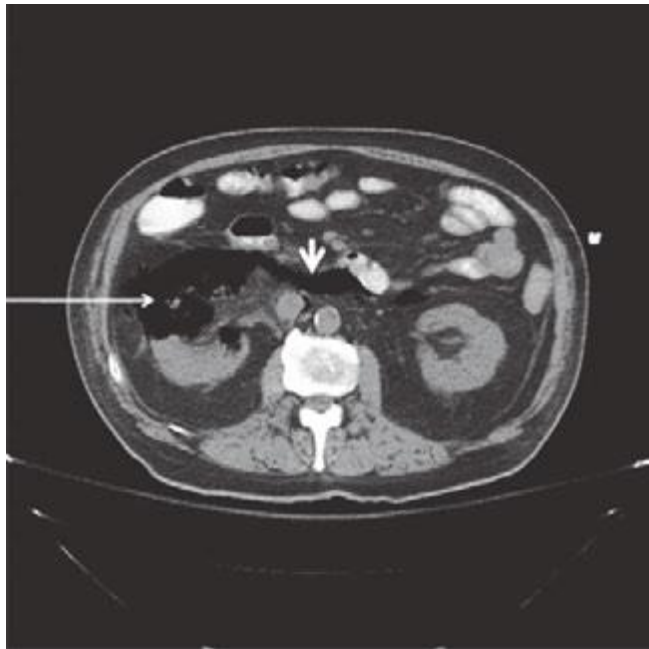


FIGURE 162-2 Emphysematous pyelonephritis. Infection of the right kidney of a diabetic man by *Escherichia coli*, a gas-forming, facultative anaerobic uropathogen, has led to destruction of the renal parenchyma (arrow) and tracking of gas through the retroperitoneal space (arrowhead).

more insidiously as recurrent episodes of cystitis, sometimes with associated pelvic and perineal pain. Men who present with recurrent cystitis should be evaluated for a prostatic focus.

COMPLICATED UTI

Complicated UTI presents as a symptomatic episode of cystitis or pyelonephritis in a man or woman with an anatomic predisposition to infection, with a foreign body in the urinary tract, or with factors predisposing to a delayed response to therapy.

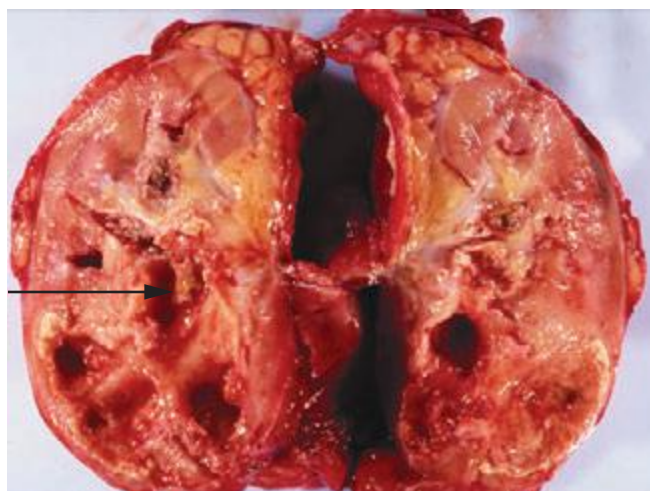
DIAGNOSTIC TOOLS

History The diagnosis of any of the UTI syndromes or ASB begins with a detailed history (Fig. 162-4). The history given by the patient

has a high predictive value in uncomplicated cystitis. A meta-analysis evaluating the probability of acute UTI on the basis of history and physical findings concluded that, in women presenting with at least one symptom of UTI (dysuria, frequency, hematuria, or back pain) and without complicating factors, the probability of acute cystitis or pyelonephritis is 50%. The even higher rates of accuracy of self-diagnosis among women with recurrent UTI probably account for the success of patient-initiated treatment of recurrent cystitis. If vaginal discharge and complicating factors are absent and risk factors for UTI are present, then the probability of UTI is close to 90%, and no laboratory evaluation is needed. Similarly, a combination of dysuria and urinary frequency in the absence of vaginal discharge increases the probability of UTI to 96%. Further laboratory evaluation with dipstick testing or urine culture is not necessary in such patients before the initiation of definitive therapy.

When the patient's history is applied as a diagnostic tool, it is important to recall that the studies included in the meta-analysis cited above did not enroll children, adolescents, pregnant women, men, or patients with complicated UTI. One significant concern is that sexually transmitted disease—that caused by *Chlamydia trachomatis* in particular—may be inappropriately treated as UTI. This concern is particularly relevant for female patients under the age of 25. The differential diagnosis to be considered when women present with dysuria includes cervicitis (*C. trachomatis*, *Neisseria gonorrhoeae*), vaginitis (*Candida albicans*, *Trichomonas vaginalis*), herpetic urethritis, interstitial cystitis, and noninfectious vaginal or vulvar irritation. Women with more than one sexual partner and inconsistent use of condoms are at high risk for both UTI and sexually transmitted disease, and symptoms alone do not always distinguish between these conditions.

The Urine Dipstick Test, Urinalysis, and Urine Culture Useful diagnostic tools include the urine dipstick test and urinalysis, both of which provide point-of-care information, and the urine culture, which can retrospectively confirm a prior diagnosis. Understanding the parameters of the dipstick test is important in interpreting its results. Only members of the family Enterobacteriaceae convert nitrate to nitrite, and enough nitrite must accumulate in the urine to reach the threshold of detection. If a woman with acute cystitis is forcing fluids and voiding frequently, the dipstick test for nitrite is less likely to be positive, even when *E. coli* is present. The leukocyte esterase test detects this enzyme in the host's polymorphonuclear leukocytes in the urine, whether the cells are intact or lysed. Many reviews have attempted to describe the diagnostic accuracy of dipstick testing. The bottom line for clinicians is that a urine dipstick test can confirm the diagnosis of uncomplicated cystitis in a patient with a reasonably high pretest probability of this



A



B

FIGURE 162-3 Xanthogranulomatous pyelonephritis. **A.** This photograph shows extensive destruction of renal parenchyma due to long-standing suppurative inflammation. The precipitating factor was obstruction by a staghorn calculus, which has been removed, leaving a depression (arrow). The mass effect of xanthogranulomatous pyelonephritis can mimic renal malignancy. **B.** A large staghorn calculus (arrow) is seen obstructing the renal pelvis and calyceal system. The lower pole of the kidney shows areas of hemorrhage and necrosis with collapse of cortical areas. (Images courtesy of Dharam M. Ramnani, MD, Virginia Urology Pathology Laboratory, Richmond, VA.)

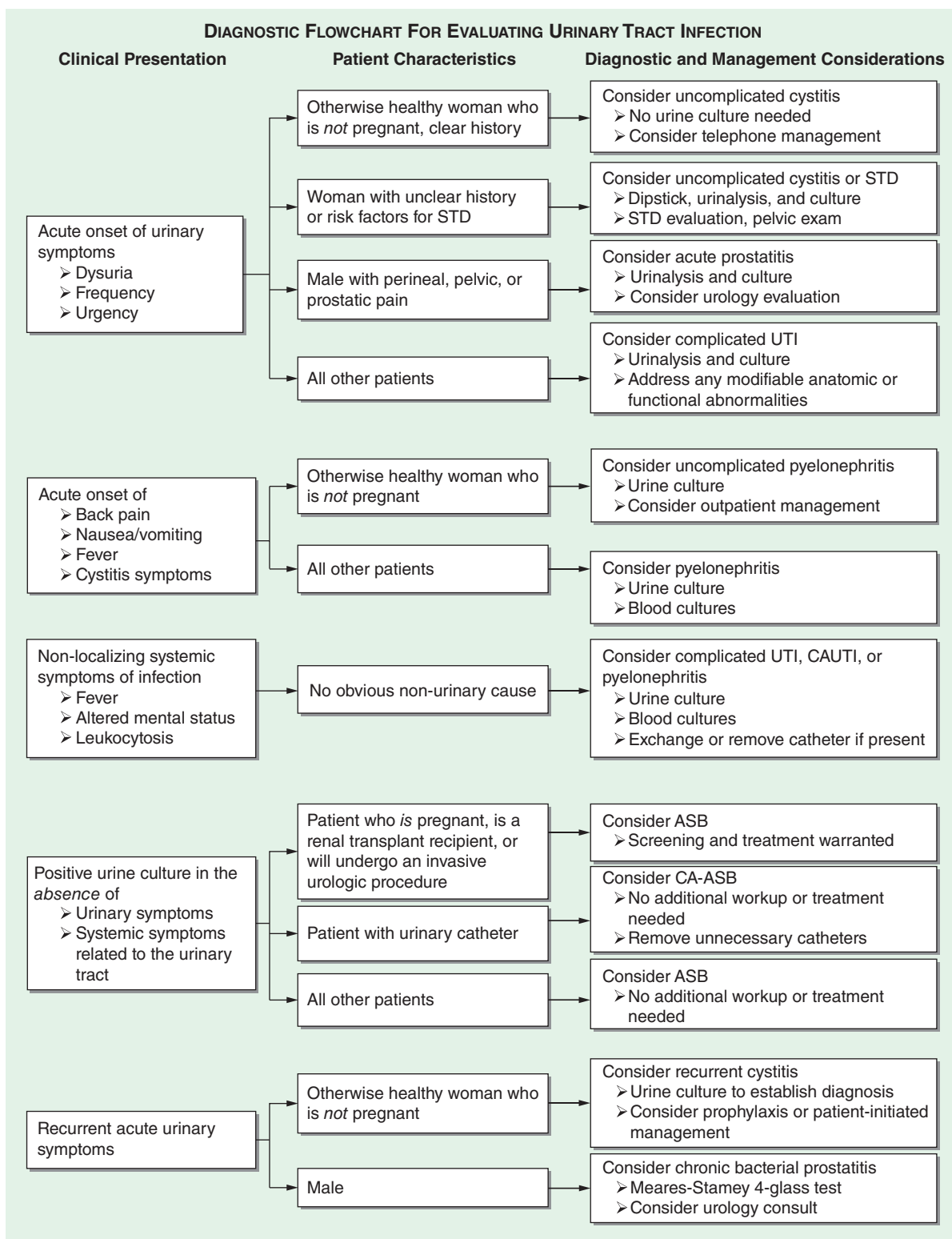


FIGURE 162-4 Diagnostic approach to urinary tract infection (UTI). STD, sexually transmitted disease; CAUTI, catheter-associated UTI; ASB, asymptomatic bacteriuria; CA-ASB, catheter-associated ASB.

disease. Either nitrite or leukocyte esterase positivity can be interpreted as a positive result. Blood in the urine also may suggest a diagnosis of UTI. A dipstick test negative for both nitrite and leukocyte esterase in the same type of patient should prompt consideration of other explanations for the patient's symptoms and collection of urine for culture. A negative dipstick test is not sufficiently sensitive to rule out bacteriuria in pregnant women, in whom it is important to detect all episodes of bacteriuria. Performance characteristics of the dipstick test

differ in men (highly specific) and in noncatheterized nursing home residents (highly sensitive).

Urine microscopy reveals pyuria in nearly all cases of cystitis and hematuria in ~30% of cases. In current practice, most hospital laboratories use an automated system rather than manual examination for urine microscopy. A machine aspirates a sample of the urine and then classifies the particles in the urine by size, shape, contrast, light scatter, volume, and other properties. These automated systems can be overwhelmed by high

866 numbers of dysmorphic red blood cells, white blood cells, or crystals; in general, counts of bacteria are less accurate than are counts of red and white blood cells. The authors' clinical recommendation is that the patient's symptoms and presentation should outweigh an incongruent result on automated urinalysis.

The detection of bacteria in a urine culture is the diagnostic "gold standard" for UTI; unfortunately, however, culture results do not become available until 24 h after the patient's presentation. Identifying specific organism(s) can require an additional 24 h. Studies of women with symptoms of cystitis have found that a colony count threshold of $>10^2$ bacteria/mL is more sensitive (95%) and specific (85%) than a threshold of 10^5 /mL for the diagnosis of acute cystitis in women. In men, the minimal level indicating infection appears to be 10^3 /mL. Urine specimens frequently become contaminated with the normal microbial flora of the distal urethra, vagina, or skin. These contaminants can grow to high numbers if the collected urine is allowed to stand at room temperature. In most instances, a culture that yields mixed bacterial species is contaminated except in settings of long-term catheterization, chronic urinary retention, or the presence of a fistula between the urinary tract and the gastrointestinal or genital tract.

DIAGNOSIS

The approach to diagnosis is influenced by which of the clinical UTI syndromes is suspected (Fig. 162-4).

Uncomplicated Cystitis in Women Uncomplicated cystitis in women can be treated on the basis of history alone. However, if the symptoms are not specific or if a reliable history cannot be obtained, then a urine dipstick test should be performed. A positive nitrite or leukocyte esterase result in a woman with one symptom of UTI increases the probability of UTI from 50% to ~80%, and empirical treatment can be considered without further testing. In this setting, a negative dipstick result does not rule out UTI, and a urine culture, close clinical follow-up, and possibly a pelvic examination are recommended. These recommendations are made with the caveat that no factors associated with complicated UTI, such as pregnancy, are known to be present.

Cystitis in Men The signs and symptoms of cystitis in men are similar to those in women, but this disease differs in several important ways in the male population. Collection of urine for culture is strongly recommended when a man has symptoms of UTI, as the documentation of bacteriuria can differentiate the less common syndromes of acute and chronic bacterial prostatitis from the very common entity of chronic pelvic pain syndrome, which is not associated with bacteriuria and thus is not usually responsive to antibacterial therapy. If the diagnosis is unclear, localization cultures using the two- or four-glass Meares-Stamey test (urine collection after prostate massage) should be undertaken to differentiate between bacterial and nonbacterial prostatic syndromes, and the patient should be referred to a urologist. Men with febrile UTI often have an elevated serum level of prostate-specific antigen as well as an enlarged prostate and enlarged seminal vesicles on ultrasound—findings indicative of prostate involvement. In 85 men with febrile UTI, symptoms of urinary retention, early recurrence of UTI, hematuria at follow-up, and voiding difficulties were predictive of surgically correctable disorders. Men with none of these symptoms had normal upper and lower urinary tracts on urologic workup.

Asymptomatic Bacteriuria The diagnosis of ASB involves both microbiologic and clinical criteria. The microbiologic criterion is usually $\geq 10^5$ bacterial CFU/mL except in catheter-associated disease, in which $\geq 10^2$ CFU/mL is the cutoff. The clinical criterion is that the person has no signs or symptoms referable to UTI.

TREATMENT URINARY TRACT INFECTIONS

Antimicrobial therapy is warranted for any symptomatic UTI. The choice of antimicrobial agent and the dose and duration of therapy depend on the site of infection and the presence or absence of complicating conditions. Each category of UTI warrants a different approach based on the particular clinical syndrome.



Antimicrobial resistance among uropathogens varies from region to region and impacts the approach to empirical treatment of UTI. *E. coli* ST131 is the predominant multilocus sequence type found worldwide as the cause of multidrug-resistant UTI. Recommendations for treatment must be considered in the context of local resistance patterns and national differences in some agents' availability. For example, fosfomycin and pivmecillinam are not available in all countries but are considered first-line options where they are available because they retain activity against a majority of uropathogens that produce extended-spectrum β -lactamases. Thus, therapeutic choices should depend on local resistance, drug availability, and individual patient factors such as recent travel and antimicrobial use.

UNCOMPLICATED CYSTITIS IN WOMEN

Since the species and antimicrobial susceptibilities of the bacteria that cause acute uncomplicated cystitis are highly predictable, many episodes of uncomplicated cystitis can be managed over the telephone (Fig. 162-4). Most patients with other UTI syndromes require further diagnostic evaluation. Although the risk of serious complications with telephone management appears to be low, studies of telephone management algorithms generally have involved otherwise healthy white women who are at low risk for complications of UTI.

In 1999, TMP-SMX was recommended as the first-line agent for treatment of uncomplicated UTI in the published guidelines of the Infectious Diseases Society of America. Antibiotic resistance among uropathogens causing uncomplicated cystitis has since increased, appreciation of the importance of collateral damage (as defined below) has increased, and newer agents have been studied. Unfortunately, there is no longer a single best agent for acute uncomplicated cystitis.

Collateral damage refers to the adverse ecologic effects of antimicrobial therapy, including killing of the normal flora and selection of drug-resistant organisms. Outbreaks of *Clostridium difficile* infection offer an example of collateral damage in the hospital environment. The implication of collateral damage in this context is that a drug that is highly efficacious for the treatment of UTI is not necessarily the optimal first-line agent if it also has pronounced secondary effects on the normal flora or is likely to change resistance patterns. Drugs used for UTI that have a minimal effect on fecal flora include pivmecillinam, fosfomycin, and nitrofurantoin. In contrast, trimethoprim, TMP-SMX, quinolones, and ampicillin affect the fecal flora more significantly; these drugs are notably the agents for which rising resistance levels have been documented.



Several effective therapeutic regimens are available for acute uncomplicated cystitis in women (Table 162-1). Well-studied first-line agents include TMP-SMX and nitrofurantoin. Second-line agents include fluoroquinolone and β -lactam compounds. Single-dose fosfomycin treatment for acute cystitis is widely used in Europe but has produced mixed results in randomized trials. There is increasing experience with the use of fosfomycin against UTIs (including complicated infections) caused by multidrug-resistant *E. coli*. Pivmecillinam is not currently available in the United States or Canada but is a popular agent in some European countries. The pros and cons of other therapies are discussed briefly below.

Traditionally, TMP-SMX has been recommended as first-line treatment for acute cystitis, and it remains appropriate to consider the use of this drug in regions with resistance rates not exceeding 20%. TMP-SMX resistance has clinical significance: in TMP-SMX-treated patients with resistant isolates, the time to symptom resolution is longer and rates of both clinical and microbiologic failure are higher. Individual host factors associated with an elevated risk of UTI caused by a strain of *E. coli* resistant to TMP-SMX include recent use of TMP-SMX or another antimicrobial agent and recent travel to an area with high rates of TMP-SMX resistance. The optimal setting for empirical use of TMP-SMX is uncomplicated UTI in a female patient who has an established relationship with the practitioner and who can thus seek further care if her symptoms do not respond promptly.

TABLE 162-1 TREATMENT STRATEGIES FOR ACUTE UNCOMPLICATED CYSTITIS

Drug and Dose	Estimated Clinical Efficacy, %	Estimated Bacterial Efficacy, ^a %	Common Side Effects
Nitrofurantoin, 100 mg bid × 5–7 d	84–95	86–92	Nausea, headache
TMP-SMX, 1 DS tablet bid × 3 d	90–100	91–100	Rash, urticaria, nausea, vomiting, hematologic abnormalities
Fosfomycin, 3-g single-dose sachet	70–91	78–83	Diarrhea, nausea, headache
Pivmecillinam, 400 mg bid × 3–7 d	55–82	74–84	Nausea, vomiting, diarrhea
Fluoroquinolones, dose varies by agent; 3-d regimen	85–95	81–98	Nausea, vomiting, diarrhea, headache, drowsiness, insomnia
β-Lactams, dose varies by agent; 5- to 7-d regimen	79–98	74–98	Diarrhea, nausea, vomiting, rash, urticaria

^aMicrobial response as measured by reduction of bacterial counts in the urine.

Note: Efficacy rates are averages or ranges calculated from the data and studies included in the 2010 Infectious Diseases Society of America/European Society of Clinical Microbiology and Infectious Diseases Guideline for Treatment of Uncomplicated UTI. TMP-SMX, trimethoprim-sulfamethoxazole; DS, double-strength.

Resistance to nitrofurantoin remains low despite >60 years of use. Since this drug affects bacterial metabolism in multiple pathways, several mutational steps are required for the development of resistance. Nitrofurantoin remains highly active against *E. coli* and most non-*E. coli* isolates. *Proteus*, *Pseudomonas*, *Serratia*, *Enterobacter*, and yeasts are all intrinsically resistant to this drug. Although nitrofurantoin has traditionally been prescribed as a 7-day regimen, similar microbiologic and clinical efficacies are noted with a 5-day course of nitrofurantoin or a 3-day course of TMP-SMX for treatment of women with acute cystitis; 3-day courses of nitrofurantoin are not recommended for acute cystitis. Nitrofurantoin does not reach significant levels in tissue and cannot be used to treat pyelonephritis.

Most fluoroquinolones are highly effective as short-course therapy for cystitis; the exception is moxifloxacin, which may not reach adequate urinary levels. The fluoroquinolones commonly used for UTI include ofloxacin, ciprofloxacin, and levofloxacin. The main concern about fluoroquinolone use for acute cystitis is the propagation of fluoroquinolone resistance, not only among uropathogens but also among other organisms causing more serious and difficult-to-treat infections at other sites. Fluoroquinolone use is also a factor driving the emergence of *C. difficile* outbreaks in hospital settings. Most experts now call for restricting fluoroquinolones to specific instances of uncomplicated cystitis in which other antimicrobial agents are not suitable. Quinolone use in certain populations, including adults >60 years of age, has been associated with an increased risk of Achilles tendon rupture.

Except for pivmecillinam, β-lactam agents generally have not performed as well as TMP-SMX or fluoroquinolones in acute cystitis. Rates of pathogen eradication are lower and relapse rates are higher with β-lactam drugs. The generally accepted explanation is that β-lactams fail to eradicate uropathogens from the vaginal reservoir. A proposed role for intracellular biofilm communities is intriguing. Many strains of *E. coli* that are resistant to TMP-SMX are also resistant to amoxicillin and cephalixin; thus, these drugs should be used only for patients infected with susceptible strains.

Urinary analgesics are appropriate in certain situations to speed resolution of bladder discomfort. The urinary tract analgesic phenazopyridine is widely used but can cause significant nausea. Combination analgesics containing urinary antiseptics (methenamine, methylene blue), a urine-acidifying agent (sodium phosphate), and an antispasmodic agent (hyoscyamine) also are available.

PYELONEPHRITIS

Since patients with pyelonephritis have tissue-invasive disease, the treatment regimen chosen should have a very high likelihood of eradicating the causative organism and should reach therapeutic blood levels quickly. High rates of TMP-SMX-resistant *E. coli* in patients with pyelonephritis have made fluoroquinolones the first-line therapy for acute uncomplicated pyelonephritis. Whether the fluoroquinolones are given orally or parenterally depends on the patient's tolerance for oral intake. A randomized clinical trial demonstrated that a 7-day course of therapy with oral ciprofloxacin (500 mg twice daily, with or without an initial IV 400-mg dose) was highly effective for the initial management of pyelonephritis in the outpatient setting. Oral TMP-SMX (one double-strength tablet twice daily for 14 days) also is effective for treatment of acute uncomplicated pyelonephritis if the uropathogen is known to be susceptible. If the pathogen's susceptibility is not known and TMP-SMX is used, an initial IV 1-g dose of ceftriaxone is recommended. Oral β-lactam agents are less effective than the fluoroquinolones and should be used with caution and close follow-up. Options for parenteral therapy for uncomplicated pyelonephritis include fluoroquinolones, an extended-spectrum cephalosporin with or without an aminoglycoside, or a carbapenem. Combinations of a β-lactam and a β-lactamase inhibitor (e.g., ampicillin-sulbactam, ticarcillin-clavulanate, piperacillin-tazobactam) or imipenem-cilastatin can be used in patients with more complicated histories, previous episodes of pyelonephritis, or recent urinary tract manipulations; in general, the treatment of such patients should be guided by urine culture results. Once the patient has responded clinically, oral therapy should be substituted for parenteral therapy.

UTI IN PREGNANT WOMEN

Nitrofurantoin, ampicillin, and the cephalosporins are considered relatively safe in early pregnancy. One retrospective case-control study suggesting an association between nitrofurantoin and birth defects has not been confirmed. Sulfonamides should clearly be avoided both in the first trimester (because of possible teratogenic effects) and near term (because of a possible role in the development of kernicterus). Fluoroquinolones are avoided because of possible adverse effects on fetal cartilage development. Ampicillin and the cephalosporins have been used extensively in pregnancy and are the drugs of choice for the treatment of asymptomatic or symptomatic UTI in this group of patients. For pregnant women with overt pyelonephritis, parenteral β-lactam therapy with or without aminoglycosides is the standard of care.

UTI IN MEN

Since the prostate is involved in the majority of cases of febrile UTI in men, the goal in these patients is to eradicate the prostatic infection as well as the bladder infection. A 7- to 14-day course of a fluoroquinolone or TMP-SMX is recommended if the uropathogen is susceptible. If acute bacterial prostatitis is suspected, antimicrobial therapy should be initiated after urine and blood are obtained for cultures. Therapy can be tailored to urine culture results and should be continued for 2–4 weeks. For documented chronic bacterial prostatitis, a 4- to 6-week course of antibiotics is often necessary. Recurrences, which are not uncommon in chronic prostatitis, often warrant a 12-week course of treatment.

COMPLICATED UTI

Complicated UTI (other than that discussed above) occurs in a heterogeneous group of patients with a wide variety of structural and functional abnormalities of the urinary tract and kidneys. The range of species and their susceptibility to antimicrobial agents are likewise heterogeneous. As a consequence, therapy for complicated UTI must be individualized and guided by urine culture results. Frequently, a patient with complicated UTI will have prior urine culture data that can be used to guide empirical therapy while current culture results are awaited. Xanthogranulomatous pyelonephritis is treated with nephrectomy. Percutaneous drainage can be used

868 as the initial therapy in emphysematous pyelonephritis and can be followed by elective nephrectomy as needed. Papillary necrosis with obstruction requires intervention to relieve the obstruction and to preserve renal function.

ASYMPTOMATIC BACTERIURIA

Treatment of ASB does not decrease the frequency of symptomatic infections or complications except in pregnant women, persons undergoing urologic surgery, and perhaps neutropenic patients and renal transplant recipients. Treatment of ASB in pregnant women and patients undergoing urologic procedures should be directed by urine culture results. In all other populations, screening for and treatment of ASB are discouraged. The majority of cases of catheter-associated bacteriuria are asymptomatic and do not warrant antimicrobial therapy.

CATHETER-ASSOCIATED UTI

Multiple institutions have released guidelines for the treatment of CAUTI, which is defined by bacteriuria and symptoms in a catheterized patient. The signs and symptoms either are localized to the urinary tract or can include otherwise unexplained systemic manifestations, such as fever. The accepted threshold for bacteriuria to meet the definition of CAUTI is $\geq 10^3$ CFU/mL, while the threshold for bacteriuria to meet the definition of ASB is $\geq 10^5$ CFU/mL.

The formation of biofilm—a living layer of uropathogens—on the urinary catheter is central to the pathogenesis of CAUTI and affects both therapeutic and preventive strategies. Organisms in a biofilm are relatively resistant to killing by antibiotics, and eradication of a catheter-associated biofilm is difficult without removal of the device itself. Furthermore, because catheters provide a conduit for bacteria to enter the bladder, bacteriuria is inevitable with long-term catheter use.

The typical signs and symptoms of UTI, including pain, urgency, dysuria, fever, peripheral leukocytosis, and pyuria, have less predictive value for the diagnosis of infection in catheterized patients. Furthermore, the presence of bacteria in the urine of a patient who is febrile and catheterized does not necessarily predict CAUTI, and other explanations for the fever should be considered.

The etiology of CAUTI is diverse, and urine culture results are essential to guide treatment. Fairly good evidence supports the practice of catheter change during treatment for CAUTI. The goal is to remove biofilm-associated organisms that could serve as a nidus for reinfection. Pathology studies reveal that many patients with long-term catheters have occult pyelonephritis. A randomized trial in persons with spinal cord injury who were undergoing intermittent catheterization found that relapse was more common after 3 days of therapy than after 14 days. In general, a 7- to 14-day course of antibiotics is recommended, but further studies on the optimal duration of therapy are needed.

In the setting of long-term catheter use, systemic antibiotics, bladder-acidifying agents, antimicrobial bladder washes, topical disinfectants, and antimicrobial drainage-bag solutions have all been ineffective at preventing the onset of bacteriuria and have been associated with the emergence of resistant organisms. The best strategy for prevention of CAUTI is to avoid insertion of unnecessary catheters and to remove catheters once they are no longer necessary. Evidence is insufficient to recommend suprapubic catheters and condom catheters as alternatives to indwelling urinary catheters as a means to prevent CAUTI. However, intermittent catheterization may be preferable to long-term indwelling urethral catheterization in certain populations (e.g., spinal cord-injured persons) to prevent both infectious and anatomic complications. Antimicrobial catheters impregnated with silver or nitrofurazone have not been shown to provide significant clinical benefit in terms of reducing rates of symptomatic UTI.

CANDIDURIA

The appearance of *Candida* in the urine is an increasingly common complication of indwelling catheterization, particularly for

patients in the intensive care unit, those taking broad-spectrum antimicrobial drugs, and those with underlying diabetes mellitus. In many studies, >50% of urinary *Candida* isolates have been found to be non-*albicans* species. The clinical presentation varies from an asymptomatic laboratory finding to pyelonephritis and even sepsis. Removal of the urethral catheter results in resolution of candiduria in more than one-third of asymptomatic cases. Treatment of asymptomatic patients does not appear to decrease the frequency of recurrence of candiduria. Treatment is recommended for patients who have symptomatic cystitis or pyelonephritis and for those who are at high risk for disseminated disease. High-risk patients include those with neutropenia, those who are undergoing urologic manipulation, those who are clinically unstable, and low-birth-weight infants. Fluconazole (200–400 mg/d for 14 days) reaches high levels in urine and is the first-line regimen for *Candida* infections of the urinary tract. Although instances of successful eradication of candiduria by some of the newer azoles and echinocandins have been reported, these agents are characterized by only low-level urinary excretion and thus are not recommended. For *Candida* isolates with high levels of resistance to fluconazole, oral flucytosine and/or parenteral amphotericin B are options. Bladder irrigation with amphotericin B generally is not recommended.

PREVENTION OF RECURRENT UTI IN WOMEN

Recurrence of uncomplicated cystitis in reproductive-age women is common, and a preventive strategy is indicated if recurrent UTIs are interfering with a patient's lifestyle. The threshold of two or more symptomatic episodes per year is not absolute; decisions about interventions should take the patient's preferences into account.

Three prophylactic strategies are available: continuous, postcoital, and patient-initiated therapy. Continuous prophylaxis and postcoital prophylaxis usually entail low doses of TMP-SMX, a fluoroquinolone, or nitrofurantoin. These regimens are all highly effective during the period of active antibiotic intake. Typically, a prophylactic regimen is prescribed for 6 months and then discontinued, at which point the rate of recurrent UTI often returns to baseline. If bothersome infections recur, the prophylactic program can be reinstated for a longer period. Selection of resistant strains in the fecal flora has been documented in studies of women taking prophylactic antibiotics for 12 months.

Patient-initiated therapy involves supplying the patient with materials for urine culture and with a course of antibiotics for self-medication at the first symptoms of infection. The urine culture is refrigerated and delivered to the physician's office for confirmation of the diagnosis. When an established and reliable patient-provider relationship exists, the urine culture can be omitted as long as the symptomatic episodes respond completely to short-course therapy and are not followed by relapse.

PROGNOSIS

Cystitis is a risk factor for recurrent cystitis and pyelonephritis. ASB is common among elderly and catheterized patients but does not in itself increase the risk of death. The relationships among recurrent UTI, chronic pyelonephritis, and renal insufficiency have been widely studied. In the absence of anatomic abnormalities, recurrent infection in children and adults does not lead to chronic pyelonephritis or to renal failure. Moreover, infection does not play a primary role in chronic interstitial nephritis; the primary etiologic factors in this condition are analgesic abuse, obstruction, reflux, and toxin exposure. In the presence of underlying renal abnormalities (particularly obstructing stones), infection as a secondary factor can accelerate renal parenchymal damage. In spinal cord-injured patients, use of a long-term indwelling bladder catheter is a well-documented risk factor for bladder cancer. Chronic bacteriuria resulting in chronic inflammation is one possible explanation for this observation.

163 Sexually Transmitted Infections: Overview and Clinical Approach

Jeanne M. Marrazzo, King K. Holmes

CLASSIFICATION AND EPIDEMIOLOGY

Worldwide, most adults acquire at least one sexually transmitted infection (STI), and many remain at risk for complications. Each year, for example, an estimated 14 million persons in the United States acquire a new genital human papillomavirus (HPV) infection, and many of these individuals are at risk for genital neoplasias. Certain STIs, such as syphilis, gonorrhea, HIV infection, hepatitis B, and chancroid, are most concentrated within “core populations” characterized by high rates of partner change, multiple concurrent partners, or “dense,” highly connected sexual networks—e.g., involving sex workers and their clients, some men who have sex with men (MSM), and persons involved in the use of illicit drugs, particularly crack cocaine and methamphetamine. Other STIs are distributed more evenly throughout societies. For example, chlamydial infections, genital infections with HPV, and genital herpes can spread widely, even in relatively low-risk populations.

In general, the product of three factors determines the initial rate of spread of any STI within a population: rate of sexual exposure of susceptible to infectious people, efficiency of transmission per exposure, and duration of infectivity of those infected. Accordingly, efforts to prevent and control STIs aim to decrease the rate of sexual exposure of susceptibles to infected persons (e.g., through individual counseling and efforts to change the norms of sexual behavior and through a variety of STI control efforts aimed at reducing the proportion of the population infected), to decrease the duration of infectivity (through early diagnosis and curative or suppressive treatment), and to decrease the efficiency of transmission (e.g., through promotion of condom use and safer sexual practices, through use of effective vaccines, and recently through male circumcision).



In all societies, STIs rank among the most common of all infectious diseases, with >30 infections now classified as predominantly sexually transmitted or as frequently sexually transmissible (Table 163-1). In developing countries, with three-quarters of the world's population and 90% of the world's STIs, factors such as population growth (especially in adolescent and young-adult age groups), rural-to-urban migration, wars, limited or no provision of reproductive health services for women, and poverty create exceptional vulnerability to disease resulting from unprotected sex. During the 1990s in China, Russia, the other states of the former Soviet Union, and South Africa, internal social structures changed rapidly as borders opened to the West, unleashing enormous new epidemics of HIV infection and other STIs. Despite advances in the provision of highly effective antiretroviral therapy worldwide, HIV remains the leading cause of death in some developing countries, and HPV and hepatitis B virus (HBV) remain important causes of cervical and hepatocellular carcinoma, respectively—two of the most common malignancies in the developing world. Sexually transmitted herpes simplex virus (HSV) infections now cause most genital ulcer disease throughout the world and an increasing proportion of cases of genital herpes in developing countries with generalized HIV epidemics, where the positive-feedback loop between HSV and HIV transmission is a growing, intractable problem. Despite this consistent link, randomized trials evaluating the efficacy of antiviral therapy in suppressing HSV in both HIV-uninfected and HIV-infected persons have not demonstrated a protective effect against acquisition or transmission of HIV. The World Health Organization estimated that 448 million new cases of four curable STIs—gonorrhea, chlamydial infection, syphilis, and trichomoniasis—occurred in 2005. Up to 50% of women of reproductive age in developing countries have bacterial vaginosis (arguably acquired sexually). All of these curable infections have been associated with increased risk of HIV transmission or acquisition.

TABLE 163-1 SEXUALLY TRANSMITTED AND SEXUALLY TRANSMISSIBLE MICROORGANISMS

Bacteria	Viruses	Other ^a
Transmitted in Adults Predominantly by Sexual Intercourse		
<i>Neisseria gonorrhoeae</i>	HIV (types 1 and 2)	<i>Trichomonas vaginalis</i>
<i>Chlamydia trachomatis</i>	Human T cell lymphotropic virus type 1	<i>Pthirus pubis</i>
<i>Treponema pallidum</i>	Herpes simplex virus type 2	
<i>Haemophilus ducreyi</i>	Human papillomavirus (multiple genital genotypes)	
<i>Klebsiella (Calymmatobacterium) granulomatis</i>	Hepatitis B virus ^b	
<i>Ureaplasma urealyticum</i>	Molluscum contagiosum virus	
<i>Mycoplasma genitalium</i>		
Sexual Transmission Repeatedly Described but Not Well Defined or Not the Predominant Mode		
<i>Mycoplasma hominis</i>	Cytomegalovirus	(?) Epstein-Barr virus
<i>Gardnerella vaginalis</i> and other vaginal bacteria	Human T cell lymphotropic virus type 2	Human herpesvirus type 8
Group B <i>Streptococcus</i>	Hepatitis C virus	<i>Candida albicans</i>
<i>Mobiluncus</i> spp.	(?) Hepatitis D virus	<i>Sarcoptes scabiei</i>
<i>Helicobacter cinaedi</i>	Herpes simplex virus type 1	
<i>Helicobacter fennelliae</i>		
Transmitted by Sexual Contact Involving Oral-Fecal Exposure; of Declining Importance in Men Who Have Sex with Men		
<i>Shigella</i> spp.	Hepatitis A virus	<i>Giardia lamblia</i>
<i>Campylobacter</i> spp.		<i>Entamoeba histolytica</i>

^aIncludes protozoa, ectoparasites, and fungi. ^bAmong U.S. patients for whom a risk factor can be ascertained, most hepatitis B virus infections are transmitted sexually.

In the United States, the prevalence of antibody to HSV-2 began to fall in the late 1990s, especially among adolescents and young adults; the decline is presumably due to delayed sexual debut, increased condom use, and lower rates of multiple (four or more) sex partners, as is well documented by the U.S. Youth Risk Behavior Surveillance System. The estimated annual incidence of HBV infection has also declined dramatically since the mid-1980s; this decrease is probably attributable more to adoption of safer sexual practices and reduced needle sharing among injection drug users than to use of hepatitis B vaccine, for which coverage among young adults (including those at high risk for this infection) initially was very limited. Genital HPV remains the most common sexually transmitted pathogen in this country, infecting 60% of a cohort of initially HPV-negative, sexually active Washington state college women within 5 years in a study conducted from 1990 to 2000. The scale-up of HPV vaccine coverage among young women has already shown promise in reducing the incidence of infection with the HPV types included in the vaccines and of conditions associated with these viruses.

In industrialized countries, fear of HIV infection since the mid-1980s, coupled with widespread behavioral interventions and better-organized systems of care for the curable STIs, initially helped curb the transmission of the latter diseases. However, foci of hyperendemic transmission persist in the southeastern United States and in most large U.S. cities. Rates of gonorrhea and syphilis remain higher in the United States than in any other Western industrialized country.

In the United States, the Centers for Disease Control and Prevention (CDC) has compiled reported rates of STIs since 1941. The incidence of reported gonorrhea peaked at 468 cases per 100,000 population in the mid-1970s and fell to a low of 98 cases per 100,000 in 2012. With increased testing and more sensitive tests, the incidence of reported *Chlamydia trachomatis* infection has been increasing steadily since reporting began in 1984, reaching an all-time peak of 457.6 cases per 100,000 in 2011. The incidence of primary and secondary syphilis per 100,000 peaked at 71 cases in 1946, fell rapidly to 3.9 cases in 1956, ranged from ~10 to 15 cases through 1987 (with markedly increased

870 rates among MSM and African Americans), and then fell to a nadir of 2.1 cases in 2000–2001 (with rates falling most rapidly among heterosexual African Americans). Unfortunately, since 1996, with the introduction of highly active antiretroviral therapy, the increased use of “serosorting” (i.e., the avoidance of unprotected sex with HIV-serodiscordant partners but not with HIV-seroconcordant partners, a strategy that provides no protection against STIs other than HIV infection), and an ongoing epidemic of methamphetamine use, gonorrhea, syphilis, and chlamydial infection have had a remarkable resurgence among MSM in North America and Europe, where outbreaks of a rare type of chlamydial infection (lymphogranuloma venereum [LGV]) that had virtually disappeared during the AIDS era have occurred. In 2012, ~75% of primary and secondary syphilis cases reported to the CDC were in MSM. These developments have resulted in a high degree of co-infection with HIV and other sexually transmitted pathogens (particularly syphilis and LGV), primarily among MSM.

MANAGEMENT OF COMMON SEXUALLY TRANSMITTED DISEASE (STD) SYNDROMES

Although other chapters discuss management of specific STIs, delineating treatment based on diagnosis of a specific infection, most patients are actually managed (at least initially) on the basis of presenting symptoms and signs and associated risk factors, even in industrialized countries. Table 163-2 lists some of the most common clinical STD syndromes and their microbial etiologies. Strategies for their management are outlined below. Chapters 225e and 226 address the management of infections with human retroviruses.

STD care and management begin with risk assessment and proceed to clinical assessment, diagnostic testing or screening, treatment, and prevention. Indeed, the routine care of any patient begins with risk assessment (e.g., for risk of heart disease, cancer). STD/HIV risk assessment is important in primary care, urgent care, and emergency care settings as well as in specialty clinics providing adolescent, HIV/AIDS, prenatal, and family planning services. STD/HIV risk assessment guides detection and interpretation of symptoms that could reflect an STD; decisions on screening or prophylactic/preventive treatment; risk reduction counseling and intervention (e.g., hepatitis B vaccination); and treatment of partners of patients with known infections. Consideration of routine demographic data (e.g., gender, age, area of residence) is a simple first step in STD/HIV risk assessment. For example, national guidelines strongly recommend routine screening of sexually active females ≤25 years of age for *C. trachomatis* infection. Table 163-3 provides a set of 11 STD/HIV risk-assessment questions that clinicians can pose verbally or that health care systems can adapt (with yes/no responses) into a routine self-administered questionnaire for use in clinics. The initial framing statement gives permission to discuss topics that may be perceived as sensitive or socially unacceptable by providers and patients alike.

Risk assessment is followed by clinical assessment (elicitation of information on specific current symptoms and signs of STDs). Confirmatory diagnostic tests (for persons with symptoms or signs) or screening tests (for those without symptoms or signs) may involve microscopic examination, culture, antigen detection tests, nucleic acid amplification tests (NAATs), or serology. Initial syndrome-based treatment should cover the most likely causes. For certain syndromes, results of rapid tests can narrow the spectrum of this initial therapy (e.g., saline microscopy of vaginal fluid for women with vaginal discharge, Gram’s stain of urethral discharge for men with urethral discharge, rapid plasma reagin test for genital ulcer). After the institution of treatment, STD management proceeds to the “4 Cs” of prevention and control: contact tracing (see “Prevention and Control of STIs,” below), ensuring compliance with therapy, and counseling on risk reduction, including condom promotion and provision.

Consistent with current guidelines, all adults should be screened for infection with HIV-1 at least once and more frequently if they are at elevated risk for acquisition of this infection.

URETHRITIS IN MEN

Urethritis in men produces urethral discharge, dysuria, or both, usually without frequency of urination. Causes include *Neisseria gonorrhoeae*,

TABLE 163-2 MAJOR SEXUALLY TRANSMITTED DISEASE SYNDROMES AND SEXUALLY TRANSMITTED MICROBIAL ETIOLOGIES

Syndrome	Sexually Transmitted Microbial Etiologies
AIDS	HIV types 1 and 2
Urethritis: males	<i>Neisseria gonorrhoeae</i> , <i>Chlamydia trachomatis</i> , <i>Mycoplasma genitalium</i> , <i>Ureaplasma urealyticum</i> (subspecies <i>urealyticum</i>), <i>Trichomonas vaginalis</i> , HSV
Epididymitis	<i>C. trachomatis</i> , <i>N. gonorrhoeae</i>
Lower genital tract infections: females	
Cystitis/urethritis	<i>C. trachomatis</i> , <i>N. gonorrhoeae</i> , HSV
Mucopurulent cervicitis	<i>C. trachomatis</i> , <i>N. gonorrhoeae</i> , <i>M. genitalium</i>
Vulvitis	<i>Candida albicans</i> , HSV
Vulvovaginitis	<i>C. albicans</i> , <i>T. vaginalis</i>
BV	BV-associated bacteria (see text)
Acute pelvic inflammatory disease	<i>N. gonorrhoeae</i> , <i>C. trachomatis</i> , BV-associated bacteria, <i>M. genitalium</i> , group B streptococci
Infertility	<i>N. gonorrhoeae</i> , <i>C. trachomatis</i> , BV-associated bacteria
Ulcerative lesions of the genitalia	HSV-1, HSV-2, <i>Treponema pallidum</i> , <i>Haemophilus ducreyi</i> , <i>C. trachomatis</i> (LGV strains), <i>Klebsiella</i> (<i>Calymmatobacterium</i>) <i>granulomatis</i>
Complications of pregnancy/ puerperium	Several agents implicated
Intestinal infections	
Proctitis	<i>C. trachomatis</i> , <i>N. gonorrhoeae</i> , HSV, <i>T. pallidum</i>
Proctocolitis or enterocolitis	<i>Campylobacter</i> spp., <i>Shigella</i> spp., <i>Entamoeba histolytica</i> , <i>Helicobacter</i> spp., other enteric pathogens
Enteritis	<i>Giardia lamblia</i>
Acute arthritis with urogenital infection or viremia	<i>N. gonorrhoeae</i> (e.g., DGI), <i>C. trachomatis</i> (e.g., reactive arthritis), HBV
Genital and anal warts	HPV (30 genital types)
Mononucleosis syndrome	CMV, HIV, EBV
Hepatitis	Hepatitis viruses, <i>T. pallidum</i> , CMV, EBV
Neoplasias	
Squamous cell dysplasias and cancers of the cervix, anus, vulva, vagina, or penis	HPV (especially types 16, 18, 31, 45)
Kaposi’s sarcoma, body-cavity lymphomas	HHV-8
T cell leukemia	HTLV-1
Hepatocellular carcinoma	HBV
Tropical spastic paraparesis	HTLV-1
Scabies	<i>Sarcoptes scabiei</i>
Pubic lice	<i>Phthirus pubis</i>

Abbreviations: BV, bacterial vaginosis; CMV, cytomegalovirus; DGI, disseminated gonococcal infection; EBV, Epstein-Barr virus; HBV, hepatitis B virus; HHV-8, human herpesvirus type 8; HPV, human papillomavirus; HSV, herpes simplex virus; HTLV, human T cell lymphotropic virus; LGV, lymphogranuloma venereum.

C. trachomatis, *Mycoplasma genitalium*, *Ureaplasma urealyticum*, *Trichomonas vaginalis*, HSV, and adenovirus.

Until recently, *C. trachomatis* caused ~30–40% of cases of nongonococcal urethritis (NGU), particularly in heterosexual men; however, the proportion of cases due to this organism has probably declined in some populations served by effective chlamydial-control programs, and older men with urethritis appear less likely to have chlamydial infection. HSV and *T. vaginalis* each cause a small proportion of NGU cases in the United States. Recently, multiple studies have consistently implicated *M. genitalium* as a probable cause of many

TABLE 163-3 ELEVEN-QUESTION STD/HIV RISK ASSESSMENT

Framing Statement:	
In order to provide the best care for you today and to understand your risk for certain infections, it is necessary for us to talk about your sexual behavior.	
Screening Questions:	
(1) Do you have any reason to think you might have a sexually transmitted infection? If so, what reason?	
(2) For all adolescents <18 years old: Have you begun having any kind of sex yet?	
STD History:	
(3) Have you ever had any sexually transmitted infections or any genital infections? If so, which ones?	
Sexual Preference:	
(4) Have you had sex with men, women, or both?	
Injection Drug Use:	
(5) Have you ever injected yourself ("shot up") with drugs? (If yes, have you ever shared needles or injection equipment?)	
(6) Have you ever had sex with a gay or bisexual man or with anyone who had ever injected drugs?	
Characteristics of Partner(s):	
(7) Has your sex partner had any sexually transmitted infections? If so, which ones?	
(8) Has your sex partner had other sex partners during the time you've been together?	
STD Symptoms Checklist:	
(9) Have you recently developed any of these symptoms?	
For Men	For Women
(a) Discharge of pus (drip) from the penis	(a) Abnormal vaginal discharge (increased amount, abnormal odor, abnormal yellow color)
(b) Genital sores (ulcers) or rash	(b) Genital sores (ulcers), rash, or itching
Sexual Practices, Past 2 Months (for patients answering yes to any of the above questions, to guide examination and testing):	
(10) Now I'd like to ask what parts of your body may have been sexually exposed to an STD (e.g., your penis, mouth, vagina, anus).	
Query about Interest in STD Screening Tests (for patients answering no to all of the above questions):	
(11) Would you like to be tested for HIV or any other STDs today? (If yes, clinician can explore which STD and why.)	

Source: Adapted from JR Curtis, KK Holmes, in KK Holmes et al (eds): *Sexually Transmitted Diseases*, 4th ed. New York, McGraw-Hill, 2008.

Chlamydia-negative cases. Fewer studies than in the past have implicated *Ureaplasma*; the ureaplasmas have been differentiated into *U. urealyticum* and *Ureaplasma parvum*, and a few studies suggest that *U. urealyticum*—but not *U. parvum*—is associated with NGU. Coliform bacteria can cause urethritis in men who practice insertive anal intercourse. The initial diagnosis of urethritis in men currently includes specific tests only for *N. gonorrhoeae* and *C. trachomatis*; it does not yet include testing for *Mycoplasma* or *Ureaplasma* species. The following summarizes the approach to the patient with suspected urethritis:

1. **Establish the presence of urethritis.** If proximal-to-distal "milking" of the urethra does not express a purulent or mucopurulent discharge, even after the patient has not voided for several hours (or preferably overnight), a Gram's-stained smear of an anterior urethral specimen obtained by passage of a small urethrogenital swab 2–3 cm into the urethra usually reveals ≥ 5 neutrophils per 1000 \times field in areas containing cells; in gonococcal infection, such a smear usually reveals gram-negative intracellular diplococci as well. Alternatively, the centrifuged sediment of the first 20–30 mL of voided urine—ideally collected as the first morning specimen—can be examined for inflammatory cells, either by microscopy showing ≥ 10 leukocytes per high-power field or by the leukocyte esterase test. Patients with symptoms who lack objective evidence of urethritis may have functional rather than organic problems and generally do not benefit from repeated courses of antibiotics.

2. **Evaluate for complications or alternative diagnoses.** A brief history and examination will exclude epididymitis and systemic complications, such as disseminated gonococcal infection (DGI) and reactive arthritis. Although digital examination of the prostate gland seldom contributes to the evaluation of sexually active young men with urethritis, men with dysuria who lack evidence of urethritis as well as sexually inactive men with urethritis should undergo prostate palpation, urinalysis, and urine culture to exclude bacterial prostatitis and cystitis.
3. **Evaluate for gonococcal and chlamydial infection.** An absence of typical gram-negative diplococci on Gram's-stained smear of urethral exudate containing inflammatory cells warrants a preliminary diagnosis of NGU, as this test is 98% sensitive for the diagnosis of gonococcal urethral infection. However, an increasing proportion of men with symptoms and/or signs of urethritis are simultaneously assessed for infection with *N. gonorrhoeae* and *C. trachomatis* by "multiplex" NAATs of first-voided urine. The urine specimen tested should consist of the first 10–15 mL of the stream, and, if possible, patients should not have voided for the prior 2 h. Culture or NAAT for *N. gonorrhoeae* may yield positive results when Gram's staining is negative; certain strains of *N. gonorrhoeae* can result in negative urethral Gram's stains in up to 30% of cases of urethral infection. Results of tests for gonococcal and chlamydial infection predict the patient's prognosis (with greater risk for recurrent NGU if neither chlamydiae nor gonococci are found than if either is detected) and can guide both the counseling given to the patient and the management of the patient's sexual partner(s).
4. **Treat urethritis promptly while test results are pending.**

TREATMENT URETHRITIS IN MEN

Table 163-4 summarizes the steps in management of sexually active men with urethral discharge and/or dysuria.

In practice, if Gram's stain does not reveal gonococci, urethritis is treated with a regimen effective for NGU, such as azithromycin or doxycycline. Both are effective, although azithromycin may give better results in *M. genitalium* infection. If gonococci are demonstrated by Gram's stain or if no diagnostic tests are performed to exclude gonorrhea definitively, treatment should include parenteral cephalosporin therapy for gonorrhea (**Chap. 181**) plus oral azithromycin, primarily for additive activity against *N. gonorrhoeae* given concerns about evolving antibiotic resistance. Azithromycin also treats *C. trachomatis*, which often causes urethral co-infection in men with gonococcal urethritis. Ideally, sexual partners should be tested for gonorrhea and chlamydial infection; regardless of whether they are tested for these infections, however, they should receive the same regimen given to the male index case. Patients with confirmed persistence or recurrence of urethritis after treatment should be re-treated with the initial regimen if they did not comply with the original treatment or were reexposed to an untreated partner. Otherwise, an intraurethral swab specimen and a first-voided urine sample should be tested for *T. vaginalis* (currently done by culture, although NAATs are more sensitive and are approved for the diagnosis of trichomoniasis in women). If compliance with initial treatment is confirmed and reexposure to an untreated sex partner is deemed unlikely, the recommended treatment is with metronidazole or tinidazole (2 g by mouth in a single dose) plus azithromycin (1 g by mouth in a single dose); the azithromycin component is especially important if this drug has not been given during initial therapy.

EPIDIDYMITIS

Acute epididymitis, almost always unilateral, produces pain, swelling, and tenderness of the epididymis, with or without symptoms or signs of urethritis. This condition must be differentiated from testicular torsion, tumor, and trauma. Torsion, a surgical emergency, usually occurs in

Usual Causes	Usual Initial Evaluation
<i>Chlamydia trachomatis</i>	Demonstration of urethral discharge or pyuria
<i>Neisseria gonorrhoeae</i>	Exclusion of local or systemic complications
<i>Mycoplasma genitalium</i>	
<i>Ureaplasma urealyticum</i>	Urethral Gram's stain to confirm urethritis, detect gram-negative diplococci
<i>Trichomonas vaginalis</i>	
Herpes simplex virus	Test for <i>N. gonorrhoeae</i> , <i>C. trachomatis</i>

Initial Treatment for Patient and Partners		
Treat gonorrhea (unless excluded):		
Ceftriaxone, 250 mg IM ^a	plus	Azithromycin, 1 g PO

Management of Recurrence

Confirm objective evidence of urethritis. If patient was reexposed to untreated or new partner, repeat treatment of patient and partner.

If patient was not reexposed, consider infection with *T. vaginalis*^b or doxycycline-resistant *M. genitalium*^c or *Ureaplasma*, and consider treatment with metronidazole, azithromycin, or both.

^aNeither oral cephalosporins nor fluoroquinolones are recommended for treatment of gonorrhea in the United States because of the emergence of increasing fluoroquinolone resistance in *N. gonorrhoeae*, especially (but not only) among men who have sex with men, and decreasing susceptibility of a still-small proportion of gonococci to ceftriaxone (Fig. 163-1). Updates on the emergence of antimicrobial resistance in *N. gonorrhoeae* can be obtained from the Centers for Disease Control and Prevention at <http://www.cdc.gov/std>. ^bIn men, the diagnosis of *T. vaginalis* infection requires culture, DNA testing, or nucleic acid amplification testing (where available) of early-morning first-voided urine sediment or of a urethral swab specimen obtained before voiding. ^c*M. genitalium* is often resistant to doxycycline and azithromycin but is usually susceptible to the fluoroquinolone moxifloxacin. Until nucleic acid amplification testing for *M. genitalium* becomes commercially available, moxifloxacin can be considered for treatment of refractory nongonococcal, nonchlamydial urethritis.

the second or third decade of life and produces a sudden onset of pain, elevation of the testicle within the scrotal sac, rotation of the epididymis from a posterior to an anterior position, and absence of blood flow on Doppler examination or ^{99m}Tc scan. Persistence of symptoms after a course of therapy for epididymitis suggests the possibility of testicular tumor or of a chronic granulomatous disease, such as tuberculosis. In sexually active men under age 35, acute epididymitis is caused most frequently by *C. trachomatis* and less commonly by *N. gonorrhoeae* and is usually associated with overt or subclinical urethritis. Acute epididymitis occurring in older men or following urinary tract instrumentation is usually caused by urinary pathogens. Similarly, epididymitis in men who have practiced insertive rectal intercourse is often caused by Enterobacteriaceae. These older men usually have no urethritis but do have bacteriuria.

TREATMENT EPIDIDYMITIS

Ceftriaxone (250 mg as a single dose IM) followed by doxycycline (100 mg by mouth twice daily for 10 days) constitutes effective treatment for epididymitis caused by *N. gonorrhoeae* or *C. trachomatis*. Neither oral cephalosporins nor fluoroquinolones are recommended for treatment of gonorrhea in the United States because of resistance in *N. gonorrhoeae*, especially (but not only) among MSM (Fig. 163-1). Oral levofloxacin (500 mg once daily for 10 days) is also effective for syndrome-based initial treatment of epididymitis when infection with Enterobacteriaceae is suspected; however, this regimen should be combined with effective therapy for possible gonococcal or chlamydial infection unless bacteriuria with Enterobacteriaceae is confirmed.

URETHRITIS AND THE URETHRAL SYNDROME IN WOMEN

C. trachomatis, *N. gonorrhoeae*, and occasionally HSV cause symptomatic urethritis—known as the urethral syndrome in women—that is characterized by “internal” dysuria (usually without urinary urgency or frequency), pyuria, and an absence of *Escherichia coli* and other

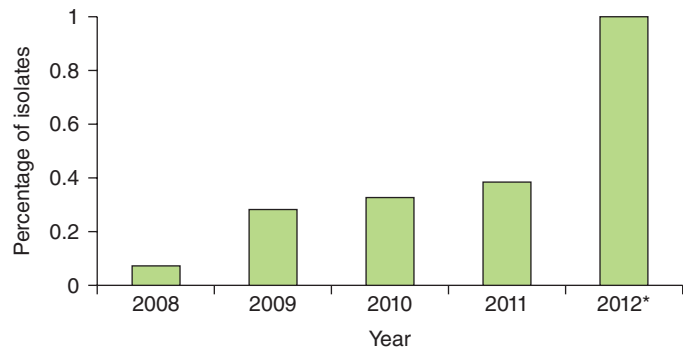


FIGURE 163-1 Proportion of *Neisseria gonorrhoeae* isolates with elevated ceftriaxone minimum inhibitory concentrations (MICs), United States 2008–2012. Elevated resistance is defined by ceftriaxone MICs of ≥ 0.125 $\mu\text{g/mL}$. *Preliminary (January–June). (From the Centers for Disease Control and Prevention: Gonococcal Isolate Surveillance Project [GISP], 2013.)

uropathogens at counts of $\geq 10^2/\text{mL}$ in urine. In contrast, the dysuria associated with vulvar herpes or vulvovaginal candidiasis (and perhaps with trichomoniasis) is often described as “external,” being caused by painful contact of urine with the inflamed or ulcerated labia or introitus. Acute onset, association with urinary urgency or frequency, hematuria, or suprapubic bladder tenderness suggests bacterial cystitis. Among women with symptoms of acute bacterial cystitis, costovertebral pain and tenderness or fever suggests acute pyelonephritis. **The management of bacterial urinary tract infection (UTI) is discussed in Chap. 162.**

Signs of vulvovaginitis, coupled with symptoms of external dysuria, suggest vulvar infection (e.g., with HSV or *Candida albicans*). Among dysuric women without signs of vulvovaginitis, bacterial UTI must be differentiated from the urethral syndrome by assessment of risk, evaluation of the pattern of symptoms and signs, and specific microbiologic testing. An STI etiology of the urethral syndrome is suggested by young age, more than one current sexual partner, a new partner within the past month, a partner with urethritis, or coexisting mucopurulent cervicitis (see below). The finding of a single urinary pathogen, such as *E. coli* or *Staphylococcus saprophyticus*, at a concentration of $\geq 10^2/\text{mL}$ in a properly collected specimen of midstream urine from a dysuric woman with pyuria indicates probable bacterial UTI, whereas pyuria with $<10^2$ conventional uropathogens per milliliter of urine (“sterile” pyuria) suggests acute urethral syndrome due to *C. trachomatis* or *N. gonorrhoeae*. Gonorrhea and chlamydial infection should be sought by specific tests (e.g., NAATs of vaginal secretions collected with a swab). Among dysuric women with sterile pyuria caused by infection with *N. gonorrhoeae* or *C. trachomatis*, appropriate treatment alleviates dysuria. The role of *M. genitalium* in the urethral syndrome in women remains undefined.

VULVOVAGINAL INFECTIONS

Abnormal Vaginal Discharge If directly questioned about vaginal discharge during routine health checkups, many women acknowledge having nonspecific symptoms of vaginal discharge that do not correlate with objective signs of inflammation or with actual infection. However, unsolicited reporting of abnormal vaginal discharge does suggest bacterial vaginosis or trichomoniasis. Specifically, an abnormally increased amount or an abnormal odor of the discharge is associated with one or both of these conditions. Cervical infection with *N. gonorrhoeae* or *C. trachomatis* does not often cause an increased amount or abnormal odor of discharge; however, when these pathogens cause cervicitis, they—like *T. vaginalis*—often result in an increased number of neutrophils in vaginal fluid, which thus takes on a yellow color. Vulvar conditions such as genital herpes or vulvovaginal candidiasis can cause vulvar pruritus, burning, irritation, or lesions as well as external dysuria (as urine passes over the inflamed vulva or areas of epithelial disruption) or vulvar dyspareunia.

Certain vulvovaginal infections may have serious sequelae. Trichomoniasis, bacterial vaginosis, and vulvovaginal candidiasis have all been associated with increased risk of acquisition of HIV infection; bacterial vaginosis may promote HIV transmission from HIV-infected women to their male sex partners. Vaginal trichomoniasis and bacterial vaginosis early in pregnancy independently predict premature onset of labor. Bacterial vaginosis can also lead to anaerobic bacterial infection of the endometrium and salpinges. Vaginitis may be an early and prominent feature of toxic shock syndrome, and recurrent or chronic vulvovaginal candidiasis develops with increased frequency among women who have systemic illnesses, such as diabetes mellitus or HIV-related immunosuppression (although only a very small proportion of women with recurrent vulvovaginal candidiasis in industrialized countries actually have a serious predisposing illness).

Thus vulvovaginal symptoms or signs warrant careful evaluation, including speculum and pelvic examination, simple rapid diagnostic tests, and appropriate therapy specific for the anatomic site and type of infection. Unfortunately, a survey in the United States indicated that clinicians seldom perform the tests required to establish the cause of such symptoms. Further, comparison of telephone and office management of vulvovaginal symptoms has documented the inaccuracy of the former, and comparison of evaluations by nurse-midwives with those by physician-practitioners showed that the practitioners'

clinical evaluations correlated poorly both with the nurses' evaluations and with diagnostic tests. The diagnosis and treatment of the three most common types of vaginal infection are summarized in **Table 163-5**.

Inspection of the vulva and perineum may reveal tender genital ulcerations or fissures (typically due to HSV infection or vulvovaginal candidiasis) or discharge visible at the introitus before insertion of a speculum (suggestive of bacterial vaginosis or trichomoniasis). Speculum examination permits the clinician to discern whether the discharge in fact looks abnormal and whether any abnormal discharge in the vagina emanates from the cervical os (mucoid and, if abnormal, yellow) or from the vagina (not mucoid, since the vaginal epithelium does not produce mucus). Symptoms or signs of abnormal vaginal discharge should prompt testing of vaginal fluid for pH, for a fishy odor when mixed with 10% KOH, and for certain microscopic features when mixed with saline (motile trichomonads and/or "clue cells") and with 10% KOH (pseudohyphae or hyphae indicative of vulvovaginal candidiasis). Additional objective laboratory tests useful for establishing the cause of abnormal vaginal discharge include rapid point-of-care tests for bacterial vaginosis and *T. vaginalis*, as described below; a DNA probe test (the Affirm test) to detect *T. vaginalis* and *C. albicans* as well as the increased concentrations of *Gardnerella vaginalis* associated with bacterial vaginosis; and a NAAT for *T. vaginalis*. Gram's

TABLE 163-5 DIAGNOSTIC FEATURES AND MANAGEMENT OF VAGINAL INFECTION

Feature	Normal Vaginal Examination	Vulvovaginal Candidiasis	Trichomonal Vaginitis	Bacterial Vaginosis
Etiology	Uninfected; lactobacilli predominant	<i>Candida albicans</i>	<i>Trichomonas vaginalis</i>	Associated with <i>Gardnerella vaginalis</i> , various anaerobic and/or noncultured bacteria, and mycoplasmas
Typical symptoms	None	Vulvar itching and/or irritation	Profuse discharge; vulvar itching	Malodorous, slightly increased discharge
Discharge				
Amount	Variable; usually scant	Scant	Often profuse	Moderate
Color ^a	Clear or translucent	White	White or yellow	White or gray
Consistency	Nonhomogeneous, flocculent	Clumped; adherent plaques	Homogeneous	Homogeneous, low viscosity; uniformly coats vaginal walls
Inflammation of vulvar or vaginal epithelium	None	Erythema of vaginal epithelium, introitus; vulvar dermatitis, fissures common	Erythema of vaginal and vulvar epithelium; colpitis macularis	None
pH of vaginal fluid ^b	Usually ≤4.5	Usually ≤4.5	Usually ≥5	Usually >4.5
Amine ("fishy") odor with 10% KOH	None	None	May be present	Present
Microscopy ^c	Normal epithelial cells; lactobacilli predominant	Leukocytes, epithelial cells; mycelia or pseudomycelia in up to 80% of <i>C. albicans</i> culture-positive persons with typical symptoms	Leukocytes; motile trichomonads seen in 80–90% of symptomatic patients, less often in the absence of symptoms	Clue cells; few leukocytes; no lactobacilli or only a few out-numbered by profuse mixed microbiota, nearly always including <i>G. vaginalis</i> plus anaerobic species on Gram's stain (Nugent's score ≥7)
Other laboratory findings		Isolation of <i>Candida</i> spp.	Isolation of <i>T. vaginalis</i> or positive NAAT ^d	
Usual treatment	None	Azole cream, tablet, or suppository—e.g., miconazole (100-mg vaginal suppository) or clotrimazole (100-mg vaginal tablet) once daily for 7 days Fluconazole, 150 mg orally (single dose)	Metronidazole or tinidazole, 2 g orally (single dose) Metronidazole, 500 mg PO bid for 7 days	Metronidazole, 500 mg PO bid for 7 days Metronidazole gel, 0.75%, one applicator (5 g) intravaginally once daily for 5 days Clindamycin, 2% cream, one full applicator vaginally each night for 7 days
Usual management of sexual partner	None	None; topical treatment if candidal dermatitis of penis is detected	Examination for sexually transmitted infection; treatment with metronidazole, 2 g PO (single dose)	None

^aColor of discharge is best determined by examination against the white background of a swab. ^bA pH determination is not useful if blood is present or if the test is performed on endocervical secretions. ^cTo detect fungal elements, vaginal fluid is digested with 10% KOH prior to microscopic examination; to examine for other features, fluid is mixed (1:1) with physiologic saline. Gram's stain is also excellent for detecting yeasts (less predictive of vulvovaginitis) and pseudomycelia or mycelia (strongly predictive of vulvovaginitis) and for distinguishing normal flora from the mixed flora seen in bacterial vaginosis, but it is less sensitive than the saline preparation for detection of *T. vaginalis*. ^dNAAT, nucleic acid amplification test (where available).

874 staining of vaginal fluid can be used to score alterations in the vaginal microbiota but is used primarily for research purposes and requires familiarity with the morphotypes and scale involved.

TREATMENT VAGINAL DISCHARGE

Patterns of treatment for vaginal discharge vary widely. In developing countries, where clinics or pharmacies often dispense treatment based on symptoms alone without examination or testing, oral treatment with metronidazole—particularly with a 7-day regimen—provides reasonable coverage against both trichomoniasis and bacterial vaginosis, the usual causes of symptoms of vaginal discharge. Metronidazole treatment of sex partners prevents reinfection of women with trichomoniasis, even though it does not help prevent the recurrence of bacterial vaginosis. Guidelines for syndromic management promulgated by the World Health Organization suggest consideration of treatment for cervical infection and for trichomoniasis, bacterial vaginosis, and vulvovaginal candidiasis in women with symptoms of abnormal vaginal discharge. However, it is important to note that the majority of chlamydial and gonococcal cervical infections produce no symptoms.

In industrialized countries, clinicians treating symptoms and signs of abnormal vaginal discharge should, at a minimum, differentiate between bacterial vaginosis and trichomoniasis, because optimal management of patients and partners differs for these two conditions (as discussed briefly below).

Vaginal Trichomoniasis (See also Chap. 254) Symptomatic trichomoniasis characteristically produces a profuse, yellow, purulent, homogeneous vaginal discharge and vulvar irritation, sometimes with visible inflammation of the vaginal and vulvar epithelium and petechial lesions on the cervix (the so-called strawberry cervix, usually evident only by colposcopy). The pH of vaginal fluid—normally <4.7—usually rises to ≥ 5 . In women with typical symptoms and signs of trichomoniasis, microscopic examination of vaginal discharge mixed with saline reveals motile trichomonads in most culture-positive cases. However, saline microscopy probably detects only one-half of all cases, and, especially in the absence of symptoms or signs, culture or NAAT is usually required for detection of the organism. NAAT for *T. vaginalis* is as sensitive as or more sensitive than culture, and NAAT of urine has disclosed surprisingly high prevalences of this pathogen among men at several STD clinics in the United States. Treatment of asymptomatic as well as symptomatic cases reduces rates of transmission and prevents later development of symptoms.

TREATMENT VAGINAL TRICHOMONIASIS

Only nitroimidazoles (e.g., metronidazole and tinidazole) consistently cure trichomoniasis. A single 2-g oral dose of metronidazole is effective and much less expensive than the alternatives. Tinidazole has a longer half-life than metronidazole, causes fewer gastrointestinal symptoms, and may be useful in treating trichomoniasis that fails to respond to metronidazole. Treatment of sexual partners—facilitated by dispensing metronidazole to the female patient to give to her partner(s), with a warning about avoiding the concurrent use of alcohol—significantly reduces both the risk of reinfection and the reservoir of infection; treating the partner is the standard of care. Intravaginal treatment with 0.75% metronidazole gel is not reliable for vaginal trichomoniasis. Systemic use of metronidazole is recommended throughout pregnancy. In a large randomized trial, metronidazole treatment of trichomoniasis during pregnancy did not reduce—and in fact actually increased—the frequency of perinatal morbidity; thus routine screening of asymptomatic pregnant women for trichomoniasis is not recommended.

Bacterial Vaginosis Bacterial vaginosis (formerly termed *nonspecific vaginitis*, *Haemophilus vaginitis*, *anaerobic vaginitis*, or *Gardnerella-associated*

vaginal discharge) is a syndrome of complex etiology that is characterized by symptoms of vaginal malodor and a slightly to moderately increased white discharge, which appears homogeneous, is low in viscosity, and evenly coats the vaginal mucosa. Bacterial vaginosis has been associated with increased risk of acquiring several other genital infections, including those caused by HIV, *C. trachomatis*, and *N. gonorrhoeae*. Other risk factors include recent unprotected vaginal intercourse, having a female sex partner, and vaginal douching. Although bacteria associated with bacterial vaginosis have been detected under the foreskin of uncircumcised men, metronidazole treatment of male partners has not reduced the rate of recurrence among affected women.

Among women with bacterial vaginosis, culture of vaginal fluid has shown markedly increased prevalences and concentrations of *G. vaginalis*, *Mycoplasma hominis*, and several anaerobic bacteria (e.g., *Mobiluncus*, *Prevotella* [formerly *Bacteroides*], and some *Peptostreptococcus* species) as well as an absence of hydrogen peroxide-producing *Lactobacillus* species that constitute most of the normal vaginal microbiota and help protect against certain cervical and vaginal infections. Broad-range polymerase chain reaction (PCR) amplification of 16S rDNA in vaginal fluid, with subsequent identification of specific bacterial species by various methods, has documented an even greater and unexpected bacterial diversity, including several unique species not previously cultivated (e.g., three species in the order Clostridiales that appear to be specific for bacterial vaginosis and are associated with metronidazole treatment failure [Fig. 163-2]). Also detected are DNA sequences related to *Atopobium vaginae*, an organism that is strongly associated with bacterial vaginosis, is resistant to metronidazole, and is also associated with recurrent bacterial vaginosis after metronidazole treatment. Other genera newly implicated in bacterial vaginosis include *Megasphaera*, *Leptotrichia*, *Eggerthella*, and *Dialister*.

Bacterial vaginosis is conventionally diagnosed clinically with the Amsel criteria, which include any three of the following four clinical abnormalities: (1) objective signs of increased white homogeneous vaginal discharge; (2) a vaginal discharge pH of >4.5 ; (3) liberation of a distinct fishy odor (attributable to volatile amines such as trimethylamine) immediately after vaginal secretions are mixed with a 10% solution of KOH; and (4) microscopic demonstration of “clue cells” (vaginal epithelial cells coated with coccobacillary organisms, which have a granular appearance and indistinct borders; Fig. 163-3) on a wet mount prepared by mixing vaginal secretions with normal saline in a ratio of $\sim 1:1$.

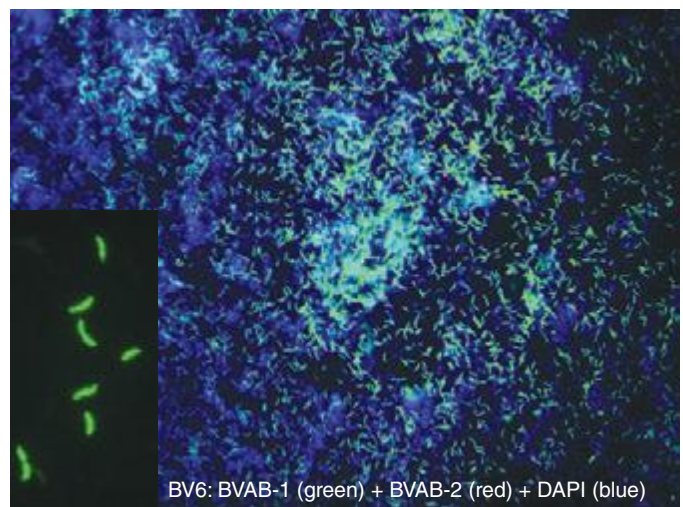


FIGURE 163-2 Broad-range polymerase chain reaction amplification of 16S rDNA in vaginal fluid from a woman with bacterial vaginosis shows a field of bacteria hybridizing with probes for bacterial vaginosis-associated bacterium 1 (BVAB-1, visible as a thin, curved green rod) and for BVAB-2 (red). The inset shows that BVAB-1 has a morphology similar to that of *Mobiluncus* (curved rod). (Reprinted with permission from DN Fredricks et al: *N Engl J Med* 353:1899, 2005.)

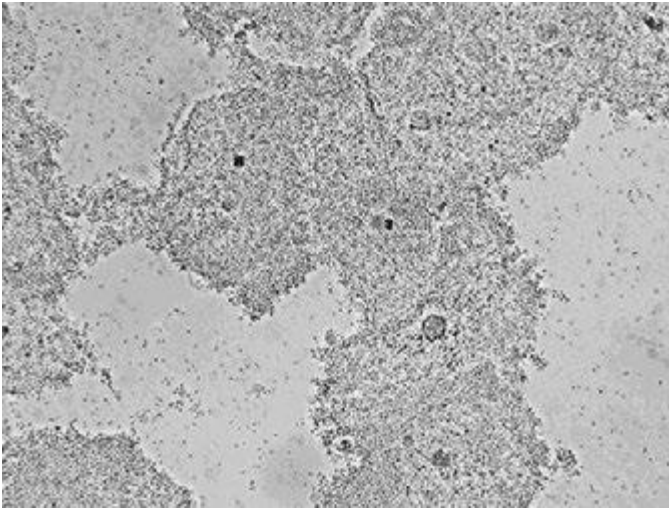


FIGURE 163-3 Wet mount of vaginal fluid showing typical clue cells from a woman with bacterial vaginosis. Note the obscured epithelial cell margins and the granular appearance attributable to many adherent bacteria ($\times 400$). (Photograph provided by Lorna K. Rabe, reprinted with permission from S Hillier et al, in KK Holmes et al [eds]: *Sexually Transmitted Diseases*, 4th ed. New York, McGraw-Hill, 2008.)

TREATMENT BACTERIAL VAGINOSIS

The standard dosage of oral metronidazole for the treatment of bacterial vaginosis is 500 mg twice daily for 7 days. The single 2-g oral dose of metronidazole recommended for trichomoniasis produces significantly lower short-term cure rates and should not be used. Intravaginal treatment with 2% clindamycin cream (one full applicator [5 g containing 100 mg of clindamycin phosphate] each night for 7 nights) or with 0.75% metronidazole gel (one full applicator [5 g containing 37.5 mg of metronidazole] twice daily for 5 days) is also approved for use in the United States and does not elicit systemic adverse reactions; the response to both of these treatments is similar to the response to oral metronidazole. Other alternatives include oral clindamycin (300 mg twice daily for 7 days), clindamycin ovules (100 g intravaginally once at bedtime for 3 days), and oral tinidazole (1 g daily for 5 days or 2 g daily for 3 days). Unfortunately, recurrence over the long term (i.e., several months later) is distressingly common after either oral or intravaginal treatment. A randomized trial comparing intravaginal gel containing 37.5 mg of metronidazole with a suppository containing 500 mg of metronidazole plus nystatin (the latter not marketed in the United States) showed significantly higher rates of recurrence with the 37.5-mg regimen; this result suggests that higher metronidazole dosages may be important in topical intravaginal therapy. Recurrences can be significantly lessened with the twice-weekly use of suppressive intravaginal metronidazole gel. As stated above, treatment of male partners with metronidazole does not prevent recurrence of bacterial vaginosis.

Efforts to replenish numbers of vaginal lactobacilli that produce hydrogen peroxide and probably sustain vaginal health have been unsuccessful. While one randomized trial of orally ingested lactobacilli found reduced rates of recurrent bacterial vaginosis, this result has not yet been either confirmed or refuted, and a randomized multicenter trial in the United States found no benefit of repeated intravaginal inoculation of a vaginal peroxide-producing *Lactobacillus* species following treatment of bacterial vaginosis with metronidazole. A meta-analysis of 18 studies concluded that bacterial vaginosis during pregnancy substantially increased the risk of preterm delivery and of spontaneous abortion. However, most studies of topical intravaginal treatment of bacterial vaginosis with clindamycin during pregnancy have not reduced adverse pregnancy outcomes. Numerous trials of oral metronidazole treatment during pregnancy have given inconsistent results, and a 2013 Cochrane review concluded that antenatal treatment of women with

bacterial vaginosis—even those with previous preterm delivery—did not reduce the risk of preterm delivery. The U.S. Preventive Services Task Force thus recommends against routine screening of pregnant women for bacterial vaginosis.

Vulvovaginal Pruritus, Burning, or Irritation Vulvovaginal candidiasis produces vulvar pruritus, burning, or irritation, generally without symptoms of increased vaginal discharge or odor. Genital herpes can produce similar symptoms, with lesions sometimes difficult to distinguish from the fissures and inflammation caused by candidiasis. Signs of vulvovaginal candidiasis include vulvar erythema, edema, fissures, and tenderness. With candidiasis, a white scanty vaginal discharge sometimes takes the form of white thrush-like plaques or cottage cheese-like curds adhering loosely to the vaginal epithelium. *C. albicans* accounts for nearly all cases of symptomatic vulvovaginal candidiasis, which probably arise from endogenous strains of *C. albicans* that have colonized the vagina or the intestinal tract. Complicated vulvovaginal candidiasis includes cases that recur four or more times per year; are unusually severe; are caused by non-*albicans* *Candida* species; or occur in women with uncontrolled diabetes, debilitation, immunosuppression, or pregnancy.

In addition to compatible clinical symptoms, the diagnosis of vulvovaginal candidiasis usually involves the demonstration of pseudohyphae or hyphae by microscopic examination of vaginal fluid mixed with saline or 10% KOH or subjected to Gram's staining. Microscopic examination is less sensitive than culture but correlates better with symptoms. Culture is typically reserved for cases that do not respond to standard first-line antimycotic agents and is undertaken to rule out imidazole or azole resistance (often associated with *Candida glabrata*) or before the initiation of suppressive antifungal therapy for recurrent disease.

TREATMENT VULVOVAGINAL PRURITUS, BURNING, OR IRRITATION

Symptoms and signs of vulvovaginal candidiasis warrant treatment, usually intravaginal administration of any of several imidazole antibiotics (e.g., miconazole or clotrimazole) for 3–7 days or of a single dose of oral fluconazole (Table 163-5). Over-the-counter marketing of such preparations has reduced the cost of care and made treatment more convenient for many women with recurrent yeast vulvovaginitis. However, most women who purchase these preparations do not have vulvovaginal candidiasis, whereas many do have other vaginal infections that require different treatment. Therefore, only women with classic symptoms of vulvar pruritus and a history of previous episodes of yeast vulvovaginitis documented by an experienced clinician should self-treat. Short-course topical intravaginal azole drugs are effective for the treatment of uncomplicated vulvovaginal candidiasis (e.g., clotrimazole, two 100-mg vaginal tablets daily for 3 days; or miconazole, a 1200-mg vaginal suppository as a single dose). Single-dose oral treatment with fluconazole (150 mg) is also effective and is preferred by many patients. Management of complicated cases (see above) and those that do not respond to the usual intravaginal or single-dose oral therapy often involves prolonged or periodic oral therapy; this situation is discussed extensively in the 2010 CDC STD treatment guidelines (<http://www.cdc.gov/std/treatment>). Treatment of sexual partners is not routinely indicated.

Other Causes of Vaginal Discharge or Vaginitis In the ulcerative vaginitis associated with staphylococcal toxic shock syndrome, *Staphylococcus aureus* should be promptly identified in vaginal fluid by Gram's stain and by culture. In desquamative inflammatory vaginitis, smears of vaginal fluid reveal neutrophils, massive vaginal epithelial-cell exfoliation with increased numbers of parabasal cells, and gram-positive cocci; this syndrome may respond to treatment with 2% clindamycin cream, often given in combination with topical steroid preparations for several weeks. Additional causes

876 of vaginitis and vulvovaginal symptoms include retained foreign bodies (e.g., tampons), cervical caps, vaginal spermicides, vaginal antiseptic preparations or douches, vaginal epithelial atrophy (in postmenopausal women or during prolonged breast-feeding in the postpartum period), allergic reactions to latex condoms, vaginal aphthae associated with HIV infection or Behçet's syndrome, and vestibulitis (a poorly understood syndrome).

MUCOPURULENT CERVICITIS

Mucopurulent cervicitis (MPC) refers to inflammation of the columnar epithelium and subepithelium of the endocervix and of any contiguous columnar epithelium that lies exposed in an ectopic position on the ectocervix. MPC in women represents the “silent partner” of urethritis in men, being equally common and often caused by the same agents (*N. gonorrhoeae*, *C. trachomatis*, or—as shown by case-control studies—*M. genitalium*); however, MPC is more difficult than urethritis to recognize. As the most common manifestation of these serious bacterial infections in women, MPC can be a harbinger or sign of upper genital tract infection, also known as *pelvic inflammatory disease* (PID; see below). In pregnant women, MPC can lead to obstetric complications. In a prospective study in Seattle of 167 consecutive patients with MPC (defined on the basis of yellow endocervical mucus or ≥ 30 polymorphonuclear leukocytes [PMNs]/1000 \times microscopic field) who were seen at STD clinics during the 1980s, slightly more than one-third of cervicovaginal specimens tested for *C. trachomatis*, *N. gonorrhoeae*, *M. genitalium*, HSV, and *T. vaginalis* revealed no identifiable etiology (Fig. 163-4). More recently, a study in Baltimore using NAATs for these pathogens still failed to identify a microbiologic etiology in nearly one-half of the 133 women with MPC.

The diagnosis of MPC rests on the detection of cardinal signs at the cervix, including yellow mucopurulent discharge from the cervical os, endocervical bleeding upon gentle swabbing, and edematous cervical ectopy (see below); the latter two findings are somewhat more common with MPC due to chlamydial infection, but signs alone do not allow a distinction between the causative pathogens. Unlike the endocervicitis produced by gonococcal or chlamydial infection, cervicitis caused by HSV produces ulcerative lesions on the stratified squamous epithelium of the ectocervix as well as on the columnar epithelium. Yellow cervical mucus on a white swab removed from the endocervix indicates the presence of PMNs. Gram's staining may confirm their

presence, although it adds relatively little to the diagnostic value of assessment for cervical signs. The presence of ≥ 20 PMNs/1000 \times microscopic field within strands of cervical mucus not contaminated by vaginal squamous epithelial cells or vaginal bacteria indicates endocervicitis. Detection of intracellular gram-negative diplococci in carefully collected endocervical mucus is quite specific but $\leq 50\%$ sensitive for gonorrhea. Therefore, specific and sensitive tests for *N. gonorrhoeae* as well as for *C. trachomatis* (e.g., NAATs) are always indicated in the evaluation of MPC.

TREATMENT MUCOPURULENT CERVICITIS

Although the above criteria for MPC are neither highly specific nor highly predictive of gonococcal or chlamydial infection in some settings, the 2010 CDC STD guidelines call for consideration of empirical treatment for MPC, pending test results, in most patients. Presumptive treatment with antibiotics active against *C. trachomatis* should be provided for women at increased risk for this common STI (risk factors: age <25 years, new or multiple sex partners, and unprotected sex), especially if follow-up cannot be ensured. Concurrent therapy for gonorrhea is indicated if the prevalence of this infection is substantial in the relevant patient population (e.g., young adults, a clinic with documented high prevalence). In this situation, therapy should include a single-dose regimen effective for gonorrhea plus treatment for chlamydial infection, as outlined in Table 163-4 for the treatment of urethritis. In settings where gonorrhea is much less common than chlamydial infection, initial therapy for chlamydial infection alone suffices, pending test results for gonorrhea. The etiology and potential benefit of treatment for endocervicitis not associated with gonorrhea or chlamydial infection have not been established. Although the antimicrobial susceptibility of *M. genitalium* is not yet well defined, the organism frequently persists after doxycycline therapy, and it currently seems reasonable to use azithromycin to treat possible *M. genitalium* infection in such cases. With resistance of *M. genitalium* to azithromycin now recognized, moxifloxacin may be a reasonable alternative. The sexual partner(s) of a woman with MPC should be examined and given a regimen similar to that chosen for the woman unless results of tests for gonorrhea or chlamydial infection in either partner warrant different therapy or no therapy.

CERVICAL ECTOPY

Cervical ectopy, often mislabeled “cervical erosion,” is easily confused with infectious endocervicitis. Ectopy represents the presence of the one-cell-thick columnar epithelium extending from the endocervix out onto the visible ectocervix. In ectopy, the cervical os may contain clear or slightly cloudy mucus but usually not yellow mucus. Colposcopy shows intact epithelium. Normally found during adolescence and early adulthood, ectopy gradually recedes through the second and third decades of life, as squamous metaplasia replaces the ectopic columnar epithelium. Oral contraceptive use favors the persistence or reappearance of ectopy, while smoking apparently accelerates squamous metaplasia. Cauterization of ectopy is not warranted. Ectopy may render the cervix more susceptible to infection with *N. gonorrhoeae*, *C. trachomatis*, or HIV.

PELVIC INFLAMMATORY DISEASE

The term *pelvic inflammatory disease* usually refers to infection that ascends from the cervix or vagina to involve the endometrium and/or fallopian tubes. Infection can extend beyond the reproductive tract to cause pelvic peritonitis, generalized peritonitis, perihepatitis, perisplenitis, or pelvic abscess. Rarely, infection not related to specific sexually transmitted pathogens extends secondarily to the pelvic organs (1) from adjacent foci of inflammation (e.g., appendicitis, regional ileitis, or diverticulitis) or bacterial vaginosis, (2) as a result of hematogenous dissemination (e.g., of tuberculosis or staphylococcal bacteremia), or (3) as a complication of certain tropical diseases (e.g., schistosomiasis). Intrauterine infection can be primary (spontaneously occurring and usually sexually transmitted) or secondary to invasive

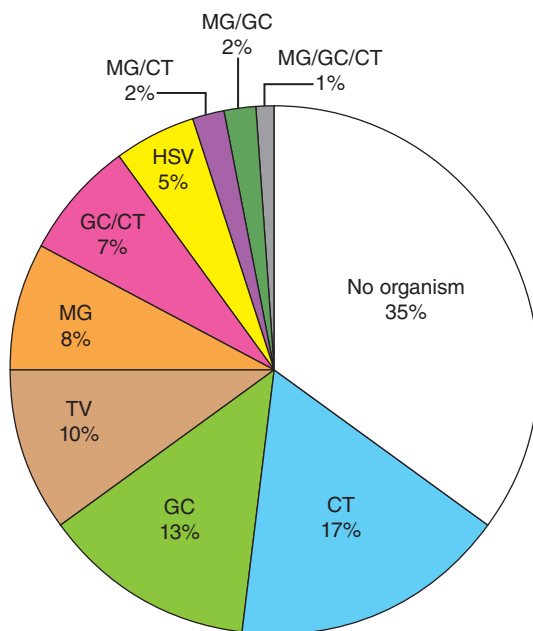


FIGURE 163-4 Organisms detected among female sexually transmitted disease clinic patients with mucopurulent cervicitis (n = 167). CT, *Chlamydia trachomatis*; GC, gonococcus; MG, *Mycoplasma genitalium*; TV, *Trichomonas vaginalis*; HSV, herpes simplex virus. (Courtesy of Dr. Lisa Manhart; with permission.)

intrauterine surgical procedures (e.g., dilation and curettage, termination of pregnancy, insertion of an intrauterine device [IUD], or hysterosalpingography) or to parturition.

Etiology The agents most often implicated in acute PID include the primary causes of endocervicitis (e.g., *N. gonorrhoeae* and *C. trachomatis*) and organisms that can be regarded as components of an altered vaginal microbiota. In general, PID is most often caused by *N. gonorrhoeae* where there is a high incidence of gonorrhea—e.g., in inner-city populations in the United States. In case-control studies, *M. genitalium* has also been significantly associated with histopathologic diagnoses of endometritis and with salpingitis.

Anaerobic and facultative organisms (especially *Prevotella* species, peptostreptococci, *E. coli*, *Haemophilus influenzae*, and group B streptococci) as well as genital mycoplasmas have been isolated from the peritoneal fluid or fallopian tubes in a varying proportion (typically one-fourth to one-third) of women with PID studied in the United States. The difficulty of determining the exact microbial etiology of an individual case of PID—short of using invasive procedures for specimen collection—has implications for the approach to empirical antimicrobial treatment of this infection.

Epidemiology In the United States, the estimated annual number of initial visits to physicians' offices for PID by women 15–44 years of age fell from an average of 400,000 during the 1980s to 250,000 in 1999 and then to 90,000 in 2011. Hospitalizations for acute PID in the United States also declined steadily throughout the 1980s and early 1990s but have remained fairly constant at 70,000–100,000 per year since 1995. Important risk factors for acute PID include the presence of endocervical infection or bacterial vaginosis, a history of salpingitis or of recent vaginal douching, and recent insertion of an IUD. Certain other iatrogenic factors, such as dilation and curettage or cesarean section, can increase the risk of PID, especially among women with endocervical gonococcal or chlamydial infection or bacterial vaginosis. Symptoms of *N. gonorrhoeae*-associated and *C. trachomatis*-associated PID often begin during or soon after the menstrual period; this timing suggests that menstruation is a risk factor for ascending infection from the cervix and vagina. Experimental inoculation of the fallopian tubes of nonhuman primates has shown that repeated exposure to *C. trachomatis* leads to the greatest degree of tissue inflammation and damage; thus, immunopathology probably contributes to the pathogenesis of chlamydial salpingitis. Women using oral contraceptives appear to be at decreased risk of symptomatic PID, and tubal sterilization reduces the risk of salpingitis by preventing intraluminal spread of infection into the tubes.

Clinical Manifestations • **ENDOMETRITIS: A CLINICAL PATHOLOGIC SYNDROME**

A study of women with clinically suspected PID who were undergoing both endometrial biopsy and laparoscopy showed that those with endometritis alone differed from those who also had salpingitis in significantly less often having lower-quadrant, adnexal, or cervical motion or abdominal rebound tenderness; fever; or elevated C-reactive protein levels. In addition, women with endometritis alone differed from those with neither endometritis nor salpingitis in more often having gonorrhea, chlamydial infection, and risk factors such as douching or IUD use. Thus, women with endometritis alone were intermediate between those with neither endometritis nor salpingitis and those with salpingitis with respect to risk factors, clinical manifestations, cervical infection prevalence, and elevated C-reactive protein level. Women with endometritis alone are at lower risk of subsequent tubal occlusion and resulting infertility than are those with salpingitis.

SALPINGITIS Symptoms of nontuberculous salpingitis classically evolve from a yellow or malodorous vaginal discharge caused by MPC and/or bacterial vaginosis to midline abdominal pain and abnormal vaginal bleeding caused by endometritis and then to bilateral lower abdominal and pelvic pain caused by salpingitis, with nausea, vomiting, and increased abdominal tenderness if peritonitis develops.

The abdominal pain in nontuberculous salpingitis is usually described as dull or aching. In some cases, pain is lacking or atypical, but active inflammatory changes are found in the course of an

unrelated evaluation or procedure, such as a laparoscopic evaluation for infertility. Abnormal uterine bleeding precedes or coincides with the onset of pain in ~40% of women with PID, symptoms of urethritis (dysuria) occur in 20%, and symptoms of proctitis (anorectal pain, tenesmus, and rectal discharge or bleeding) are occasionally seen in women with gonococcal or chlamydial infection.

Speculum examination shows evidence of MPC (yellow endocervical discharge, easily induced endocervical bleeding) in the majority of women with gonococcal or chlamydial PID. Cervical motion tenderness is produced by stretching of the adnexal attachments on the side toward which the cervix is pushed. Bimanual examination reveals uterine fundal tenderness due to endometritis and abnormal adnexal tenderness due to salpingitis that is usually, but not necessarily, bilateral. Adnexal swelling is palpable in about one-half of women with acute salpingitis, but evaluation of the adnexae in a patient with marked tenderness is not reliable. The initial temperature is >38°C in only about one-third of patients with acute salpingitis. Laboratory findings include elevation of the erythrocyte sedimentation rate (ESR) in 75% of patients with acute salpingitis and elevation of the peripheral white blood cell count in up to 60%.

Unlike nontuberculous salpingitis, genital tuberculosis often occurs in older women, many of whom are postmenopausal. Presenting symptoms include abnormal vaginal bleeding, pain (including dysmenorrhea), and infertility. About one-quarter of these women have had adnexal masses. Endometrial biopsy shows tuberculous granulomas and provides optimal specimens for culture.

PERIHEPATITIS AND PERIAPPENDICITIS Pleuritic upper-abdominal pain and tenderness, usually localized to the right upper quadrant (RUQ), develop in 3–10% of women with acute PID. Symptoms of perihepatitis arise during or after the onset of symptoms of PID and may overshadow lower abdominal symptoms, thereby leading to a mistaken diagnosis of cholecystitis. In perhaps 5% of cases of acute salpingitis, early laparoscopy reveals perihepatic inflammation ranging from edema and erythema of the liver capsule to exudate with fibrinous adhesions between the visceral and parietal peritoneum. When treatment is delayed and laparoscopy is performed late, dense “violinsting” adhesions can be seen over the liver; chronic exertional or positional RUQ pain ensues when traction is placed on the adhesions. Although perihepatitis, also known as the *Fitz-Hugh-Curtis syndrome*, was for many years specifically attributed to gonococcal salpingitis, most cases are now attributed to chlamydial salpingitis. In patients with chlamydial salpingitis, serum titers of microimmunofluorescent antibody to *C. trachomatis* are typically much higher when perihepatitis is present than when it is absent.

Physical findings include RUQ tenderness and usually include adnexal tenderness and cervicitis, even in patients whose symptoms do not suggest salpingitis. Results of liver function tests and RUQ ultrasonography are nearly always normal. The presence of MPC and pelvic tenderness in a young woman with subacute pleuritic RUQ pain and normal ultrasonography of the gallbladder points to a diagnosis of perihepatitis.

Periappendicitis (appendiceal serositis without involvement of the intestinal mucosa) has been found in ~5% of patients undergoing appendectomy for suspected appendicitis and can occur as a complication of gonococcal or chlamydial salpingitis.

Among women with salpingitis, HIV infection is associated with increased severity of salpingitis and with tuboovarian abscess requiring hospitalization and surgical drainage. Nonetheless, among women with HIV infection and salpingitis, the clinical response to conventional antimicrobial therapy (coupled with drainage of tuboovarian abscess, when found) has usually been satisfactory.

Diagnosis Treatment appropriate for PID must not be withheld from patients who have an equivocal diagnosis; it is better to err on the side of overdiagnosis and overtreatment. On the other hand, it is essential to differentiate between salpingitis and other pelvic pathology, particularly surgical emergencies such as appendicitis and ectopic pregnancy.

Nothing short of laparoscopy definitively identifies salpingitis, but routine laparoscopy to confirm suspected salpingitis is generally

878 impractical. Most patients with acute PID have lower abdominal pain of <3 weeks' duration, pelvic tenderness on bimanual pelvic examination, and evidence of lower genital tract infection (e.g., MPC). Approximately 60% of such patients have salpingitis at laparoscopy, and perhaps 10–20% have endometritis alone. Among the patients with these findings, a rectal temperature >38°C, a palpable adnexal mass, and elevation of the ESR to >15 mm/h also raise the probability of salpingitis, which has been found at laparoscopy in 68% of patients with one of these additional findings, 90% of patients with two, and 96% of patients with three. However, only 17% of all patients with laparoscopy-confirmed salpingitis have had all three additional findings.

In a woman with pelvic pain and tenderness, increased numbers of PMNs (30 per 1000× microscopic field in strands of cervical mucus) or leukocytes outnumbering epithelial cells in vaginal fluid (in the absence of trichomonal vaginitis, which also produces PMNs in vaginal discharge) increase the predictive value of a clinical diagnosis of acute PID, as do onset with menses, history of recent abnormal menstrual bleeding, presence of an IUD, history of salpingitis, and sexual exposure to a male with urethritis. Appendicitis or another disorder of the gut is favored by the early onset of anorexia, nausea, or vomiting; the onset of pain later than day 14 of the menstrual cycle; or unilateral pain limited to the right or left lower quadrant. Whenever the diagnosis of PID is being considered, serum assays for human β -chorionic gonadotropin should be performed; these tests are usually positive with ectopic pregnancy. Ultrasonography and magnetic resonance imaging (MRI) can be useful for the identification of tuboovarian or pelvic abscess. MRI of the tubes can also show increased tubal diameter, intratubal fluid, or tubal wall thickening in cases of salpingitis.

The primary and uncontested value of laparoscopy in women with lower abdominal pain is for the exclusion of other surgical problems. Some of the most common or serious problems that may be confused with salpingitis (e.g., acute appendicitis, ectopic pregnancy, corpus luteum bleeding, ovarian tumor) are unilateral. Unilateral pain or pelvic mass, although not incompatible with PID, is a strong indication for laparoscopy unless the clinical picture warrants laparotomy instead. Atypical clinical findings such as the absence of lower genital tract infection, a missed menstrual period, a positive pregnancy test, or failure to respond to appropriate therapy are other common indications for laparoscopy. Endometrial biopsy is relatively sensitive and specific for the diagnosis of endometritis, which correlates well with the presence of salpingitis.

Vaginal or endocervical swab specimens should be examined by NAATs for *N. gonorrhoeae* and *C. trachomatis*. At a minimum, vaginal fluid should be evaluated for the presence of PMNs, and endocervical secretions ideally should be assessed by Gram's staining for PMNs and gram-negative diplococci, which indicate gonococcal infection. The clinical diagnosis of PID made by expert gynecologists is confirmed by laparoscopy or endometrial biopsy in ~90% of women who also have cultures positive for *N. gonorrhoeae* or *C. trachomatis*. Even among women with no symptoms suggestive of acute PID who were attending an STD clinic or a gynecology clinic in Pittsburgh, endometritis was significantly associated with endocervical gonorrhea or chlamydial infection or with bacterial vaginosis, being detected in 26%, 27%, and 15% of women with these conditions, respectively.

TREATMENT PELVIC INFLAMMATORY DISEASE

Recommended combination regimens for ambulatory or parenteral management of PID are presented in **Table 163-6**. Women managed as outpatients should receive a combined regimen with broad activity, such as ceftriaxone (to cover possible gonococcal infection) followed by doxycycline (to cover possible chlamydial infection). Metronidazole can be added, if tolerated, to enhance activity against anaerobes; this addition should be strongly considered if bacterial vaginosis is documented. Although few methodologically sound clinical trials (especially with prolonged follow-up) have been

TABLE 163-6 COMBINATION ANTIMICROBIAL REGIMENS RECOMMENDED FOR OUTPATIENT TREATMENT OR FOR PARENTERAL TREATMENT OF PELVIC INFLAMMATORY DISEASE

Outpatient Regimens ^a	Parenteral Regimens
Ceftriaxone (250 mg IM once)	Initiate parenteral therapy with either of the following regimens; continue parenteral therapy until 48 h after clinical improvement; then change to outpatient therapy, as described in the text
plus Doxycycline (100 mg PO bid for 14 days)	
plus^b Metronidazole (500 mg PO bid for 14 days)	Regimen A Cefotetan (2 g IV q12h) or cefoxitin (2 g IV q6h)
	plus Doxycycline (100 mg IV or PO q12h)
	Regimen B Clindamycin (900 mg IV q8h)
	plus Gentamicin (loading dose of 2 mg/kg IV or IM, then maintenance dose of 1.5 mg/kg q8h)

^aSee text for discussion of options in the patient who is intolerant of cephalosporins. ^bThe addition of metronidazole is recommended by some experts, particularly if bacterial vaginosis is present.

Source: Adapted from Centers for Disease Control and Prevention: MMWR Recomm Rep 59(RR-12):1, 2010.

conducted, one meta-analysis suggested a benefit of providing good coverage against anaerobes.

The CDC STD treatment guidelines recommend initiation of empirical treatment for PID in sexually active young women and other women at risk for PID if they are experiencing pelvic or lower abdominal pain, if no other cause for the pain can be identified, and if pelvic examination reveals one or more of the following criteria for PID: cervical motion tenderness, uterine tenderness, or adnexal tenderness. Women with suspected PID can be treated as either outpatients or inpatients. In the multicenter Pelvic Inflammatory Disease Evaluation and Clinical Health (PEACH) trial, 831 women with mild to moderately severe symptoms and signs of PID were randomized to receive either inpatient treatment with IV cefoxitin and doxycycline or outpatient treatment with a single IM dose of cefoxitin plus oral doxycycline. Short-term clinical and microbiologic outcomes and long-term outcomes were equivalent in the two groups. Nonetheless, hospitalization should be considered when (1) the diagnosis is uncertain and surgical emergencies such as appendicitis and ectopic pregnancy cannot be excluded, (2) the patient is pregnant, (3) pelvic abscess is suspected, (4) severe illness or nausea and vomiting preclude outpatient management, (5) the patient has HIV infection, (6) the patient is assessed as unable to follow or tolerate an outpatient regimen, or (7) the patient has failed to respond to outpatient therapy. Some experts also prefer to hospitalize adolescents with PID for initial therapy, although younger women do as well as older women on outpatient therapy.

Currently, oral cephalosporins, doxycycline, and the fluoroquinolones do not provide reliable coverage for gonococcal infection. Thus, adequate oral treatment of women with serious intolerance to cephalosporins is a challenge. If penicillins are an option, amoxicillin/clavulanic acid combined with doxycycline has elicited a short-term clinical response in one trial. If fluoroquinolones are the only option and if the community prevalence and individual risk of gonorrhea are known to be low, oral levofloxacin (500 mg once daily) or ofloxacin (400 mg twice daily) for 14 days, with or without metronidazole, may be considered; moreover, clinical trials performed outside the United States support the effectiveness of oral moxifloxacin. In this case, it is imperative to perform a sensitive diagnostic test for gonorrhea (ideally, culture to test for antimicrobial susceptibility) before initiation of therapy. For women whose PID involves quinolone-resistant *N. gonorrhoeae*, treatment is uncertain but could include parenteral

gentamicin or oral azithromycin, although the latter agent has not been studied for this purpose.

For hospitalized patients, the following two parenteral regimens (Table 163-6) have given nearly identical results in a multicenter randomized trial:

1. Doxycycline plus either cefotetan or ceftiofloxacin: Administration of these drugs should be continued by the IV route for at least 48 h after the patient's condition improves and then followed with oral doxycycline (100 mg twice daily) to complete 14 days of therapy.
2. Clindamycin plus gentamicin in patients with normal renal function: Once-daily administration of gentamicin (with combination of the total daily dose into a single daily dose) has not been evaluated in PID but has been efficacious in other serious infections and could be substituted. Treatment with these drugs should be continued for at least 48 h after the patient's condition improves and then followed with oral doxycycline (100 mg twice daily) or clindamycin (450 mg four times daily) to complete 14 days of therapy. In cases with tuboovarian abscess, clindamycin rather than doxycycline for continued therapy provides better coverage for anaerobic infection.

FOLLOW-UP

Hospitalized patients should show substantial clinical improvement within 3–5 days. Women treated as outpatients should be clinically reevaluated within 72 h. A follow-up telephone survey of women seen in an emergency department and given a prescription for 10 days of oral doxycycline for PID found that 28% never filled the prescription and 41% stopped taking the medication early (after an average of 4.1 days), often because of persistent symptoms, lack of symptoms, or side effects. Women not responding favorably to ambulatory therapy should be hospitalized for parenteral therapy and further diagnostic evaluations, including a consideration of laparoscopy. Male sex partners should be evaluated and treated empirically for gonorrhea and chlamydial infection. After completion of treatment, tests for persistent or recurrent infection with *N. gonorrhoeae* or *C. trachomatis* should be performed if symptoms persist or recur or if the patient has not complied with therapy or has been reexposed to an untreated sex partner.

SURGERY

Surgery is necessary for the treatment of salpingitis only in the face of life-threatening infection (such as rupture or threatened rupture of a tuboovarian abscess) or for drainage of an abscess. Conservative surgical procedures are usually sufficient. Pelvic abscesses can often be drained by posterior colpotomy, and peritoneal lavage can be used for generalized peritonitis.

Prognosis Late sequelae include infertility due to bilateral tubal occlusion, ectopic pregnancy due to tubal scarring without occlusion, chronic pelvic pain, and recurrent salpingitis. The overall post-salpingitis risk of infertility due to tubal occlusion in a large study in Sweden was 11% after one episode of salpingitis, 23% after two episodes, and 54% after three or more episodes. A University of Washington study found a sevenfold increase in the risk of ectopic pregnancy and an eightfold increase in the rate of hysterectomy after PID.

Prevention A randomized controlled trial designed to determine whether selective screening for chlamydial infection reduces the risk of subsequent PID showed that women randomized to undergo screening had a 56% lower rate of PID over the following year than did women receiving the usual care without screening. This report helped prompt U.S. national guidelines for risk-based chlamydial screening of young women to reduce the incidence of PID and the prevalence of post-PID sequelae, while also reducing sexual transmission of *C. trachomatis*. The CDC and the U.S. Preventive Services Task Force recommend that sexually active women ≤ 25 years of age be screened

for genital chlamydial infection annually. Despite this recommendation, screening coverage in many primary care settings remains low.

ULCERATIVE GENITAL OR PERIANAL LESIONS

Genital ulceration reflects a set of important STIs, most of which sharply increase the risk of sexual acquisition and shedding of HIV. In a 1996 study of genital ulcers in 10 of the U.S. cities with the highest rates of primary syphilis, PCR testing of ulcer specimens demonstrated HSV in 62% of patients, *Treponema pallidum* (the cause of syphilis) in 13%, and *Haemophilus ducreyi* (the cause of chancroid) in 12–20%. Today, genital herpes represents an even higher proportion of genital ulcers in the United States and other industrialized countries.

In Asia and Africa, chancroid (Fig. 163-5) was once considered the most common type of genital ulcer, followed in frequency by primary syphilis and then genital herpes (Fig. 163-6). With increased efforts to control chancroid and syphilis and widespread use of broad-spectrum antibiotics to treat STI-related syndromes, together with more frequent recurrences or persistence of genital herpes attributable to HIV infection, PCR testing of genital ulcers now clearly implicates genital herpes as by far the most common cause of genital ulceration in most developing countries. LGV due to *C. trachomatis* (Fig. 163-7) and donovanosis (granuloma inguinale, due to *Klebsiella granulomatis*; see Fig. 198e-1) continue to cause genital ulceration in some developing countries. LGV virtually disappeared in industrialized countries during the first 20 years of the HIV pandemic, but outbreaks are again occurring in Europe (including the United Kingdom), in North America, and in Australia. In these outbreaks, LGV typically presents as proctitis, with or without anal lesions, in men who report unprotected receptive anal intercourse, very often in association with HIV and/or hepatitis C virus infection; the latter may be an acute infection acquired through the same exposure. Other causes of genital ulcers include (1) candidiasis and traumatized genital warts—both readily recognized; (2) lesions due to genital involvement by more widespread dermatoses; (3) cutaneous manifestations of systemic diseases such as genital mucosal ulceration in Stevens-Johnson syndrome or Behçet's disease; (4) superinfections of lesions that may originally have been sexually acquired (for example, methicillin-resistant *S. aureus* complicating a genital ulcer due to HSV-2); and (5) localized drug reactions, such as the ulcers occasionally seen with topical paromomycin cream or boric acid preparations.

Diagnosis Although most genital ulcerations cannot be diagnosed confidently on clinical grounds alone, clinical findings (Table 163-7)



FIGURE 163-5 Chancroid: multiple, painful, punched-out ulcers with undermined borders on the labia occurring after autoinoculation.



FIGURE 163-6 Genital herpes. A relatively mild, superficial ulcer is typically seen in episodic outbreaks. (Courtesy of Michael Remington, University of Washington Virology Research Clinic.)



FIGURE 163-7 Lymphogranuloma venereum (LGV): striking tender lymphadenopathy occurring at the femoral and inguinal lymph nodes, separated by a groove made by Poupart's ligament. This "sign-of-the-groove" is not considered specific for LGV; for example, lymphomas may present with this sign.

TABLE 163-8 INITIAL MANAGEMENT OF GENITAL OR PERIANAL ULCER

Causative Pathogens

Herpes simplex virus (HSV)
Treponema pallidum (primary syphilis)
Haemophilus ducreyi (chancroid)

Usual Initial Laboratory Evaluation

Dark-field exam (if available), direct FA, or PCR for *T. pallidum*
 RPR, VDRL, or EIA serologic test for syphilis^a
 Culture, direct FA, ELISA, or PCR for HSV
 HSV-2-specific serology (consider)
 In chancroid-endemic area: PCR or culture for *H. ducreyi*

Initial Treatment

Herpes confirmed or suspected (history or sign of vesicles):

Treat for genital herpes with acyclovir, valacyclovir, or famciclovir.

Syphilis confirmed (dark-field, FA, or PCR showing *T. pallidum*, or RPR reactive):

Benzathine penicillin (2.4 million units IM once to patient, to recent [e.g., within 3 months] seronegative partner[s], and to all seropositive partners)^b

Chancroid confirmed or suspected (diagnostic test positive, or HSV and syphilis excluded, and persistent lesion):

Ciprofloxacin (500 mg PO as single dose) or
 Ceftriaxone (250 mg IM as single dose) or
 Azithromycin (1 g PO as single dose)

^aIf results are negative but primary syphilis is suspected, treat presumptively when indicated by epidemiologic and sexual risk assessment; repeat in 1 week. ^bThe same treatment regimen is also effective in HIV-infected persons with early syphilis.

Abbreviations: EIA, enzyme immunoassay; ELISA, enzyme-linked immunosorbent assay; FA, fluorescent antibody; HSV, herpes simplex virus; PCR, polymerase chain reaction; RPR, rapid plasma reagin; VDRL, Venereal Disease Research Laboratory.

and epidemiologic considerations can usually guide initial management (Table 163-8) pending results of specific tests. Clinicians should order a rapid serologic test for syphilis in all cases of genital ulcer. To evaluate lesions except those highly characteristic of infection with HSV (i.e., those with herpetic vesicles), dark-field microscopy, direct immunofluorescence, and PCR for *T. pallidum* can be useful but are rarely available today in most countries. It is important to note that 30% of syphilitic chancres—the primary ulcer of syphilis—are associated with an initially nonreactive syphilis serology. All patients presenting with genital ulceration should be counseled and tested for HIV infection.

Typical vesicles or pustules or a cluster of painful ulcers preceded by vesiculopustular lesions suggests genital herpes. These typical clinical manifestations make detection of the virus optional; however, many patients want confirmation of the diagnosis, and differentiation of

TABLE 163-7 CLINICAL FEATURES OF GENITAL ULCERS

Feature	Syphilis	Herpes	Chancroid	Lymphogranuloma Venereum	Donovanosis
Incubation period	9–90 days	2–7 days	1–14 days	3 days–6 weeks	1–4 weeks (up to 6 months)
Early primary lesions	Papule	Vesicle	Pustule	Papule, pustule, or vesicle	Papule
No. of lesions	Usually one	Multiple	Usually multiple, may coalesce	Usually one; often not detected, despite lymphadenopathy	Variable
Diameter	5–15 mm	1–2 mm	Variable	2–10 mm	Variable
Edges	Sharply demarcated, elevated, round, or oval	Erythematous	Undermined, ragged, irregular	Elevated, round, or oval	Elevated, irregular
Depth	Superficial or deep	Superficial	Excavated	Superficial or deep	Elevated
Base	Smooth, nonpurulent, relatively nonvascular	Serous, erythematous, nonvascular	Purulent, bleeds easily	Variable, nonvascular	Red and velvety, bleeds readily
Induration	Firm	None	Soft	Occasionally firm	Firm
Pain	Uncommon	Frequently tender	Usually very tender	Variable	Uncommon
Lymphadenopathy	Firm, nontender, bilateral	Firm, tender, often bilateral with initial episode	Tender, may suppurate, loculated, usually unilateral	Tender, may suppurate, loculated, usually unilateral	None; pseudobuboes

Source: From RM Ballard, in KK Holmes et al (eds): *Sexually Transmitted Diseases*, 4th ed. New York, McGraw-Hill, 2008.

HSV-1 from HSV-2 has prognostic implications, because the latter causes more frequent genital recurrences.

Painless, nontender, indurated ulcers with firm, nontender inguinal adenopathy suggest primary syphilis. If results of dark-field examination and a rapid serologic test for syphilis are initially negative, presumptive therapy should be provided on the basis of the individual's risk. For example, with increasing rates of syphilis among MSM in the United States, most experts would not withhold therapy for this infection pending watchful waiting and/or subsequent detection of seroconversion. Repeated serologic testing for syphilis 1 or 2 weeks after treatment of seronegative primary syphilis usually demonstrates seroconversion.

"Atypical" or clinically trivial ulcers may be more common manifestations of genital herpes than classic vesiculopustular lesions. Specific tests for HSV in such lesions are therefore indicated (**Chap. 216**). Commercially available type-specific serologic tests for serum antibody to HSV-2 may give negative results, especially when patients present early with the initial episode of genital herpes or when HSV-1 is the cause of genital herpes (as is often the case today). Furthermore, a positive test for antibody to HSV-2 does not prove that the current lesions are herpetic, because nearly one-fifth of the general population of the United States (and no doubt a higher proportion of those at risk for other STIs) becomes seropositive for HSV-2 during early adulthood. Although even "type-specific" tests for HSV-2 that are commercially available in the United States are not 100% specific, a positive HSV-2 serology does enable the clinician to tell the patient that he or she has probably had genital herpes, should learn to recognize symptoms, and should avoid sex during recurrences. In addition, because genital shedding and sexual transmission of HSV-2 often occur in the absence of symptoms and signs of recurrent herpetic lesions, persons who have a history of genital herpes or who are seropositive for HSV-2 should consider the use of condoms or suppressive antiviral therapy, both of which can reduce the risk of HSV-2 transmission to a sexual partner.

Demonstration of *H. ducreyi* by culture (or by PCR, where available) is most useful when ulcers are painful and purulent, especially if inguinal lymphadenopathy with fluctuance or overlying erythema is noted; if chancroid is prevalent in the community; or if the patient has recently had a sexual exposure elsewhere in a chancroid-endemic area (e.g., a developing country). Enlarged, fluctuant lymph nodes should be aspirated for culture or PCR to detect *H. ducreyi* as well as for Gram's staining and culture to rule out the presence of other pyogenic bacteria.

When genital ulcers persist beyond the natural history of initial episodes of herpes (2–3 weeks) or of chancroid or syphilis (up to 6 weeks) and do not resolve with syndrome-based antimicrobial therapy, then—in addition to the usual tests for herpes, syphilis, and chancroid—biopsy is indicated to exclude donovanosis, carcinoma, and other nonvenereal dermatoses. If not performed previously, HIV serology should be standard because chronic, persistent genital herpes is common in AIDS.

TREATMENT ULCERATIVE GENITAL OR PERIANAL LESIONS

Immediate syndrome-based treatment for acute genital ulcerations (after collection of all necessary diagnostic specimens at the first visit) is often appropriate before all test results become available because patients with typical initial or recurrent episodes of genital or anorectal herpes can benefit from prompt oral antiviral therapy (**Chap. 216**); because early treatment of sexually transmitted causes of genital ulcers decreases further transmission; and because many patients do not return for test results and treatment. A thorough assessment of the patient's sexual-risk profile and medical history is critical in determining the course of initial management. The patient who has risk factors consistent with exposure to syphilis (e.g., a male patient who reports sex with other men or who has HIV infection) should generally receive initial treatment for syphilis. Empirical therapy for chancroid should be considered if there has been an exposure in an area of the world where chancroid occurs or if regional lymph node suppuration is evident. In resource-poor

settings lacking ready access to diagnostic tests, this approach to syndromic treatment for syphilis and chancroid has helped bring these two diseases under control. Finally, empirical antimicrobial therapy may be indicated if ulcers persist and the diagnosis remains unclear after a week of observation despite attempts to diagnose herpes, syphilis, and chancroid.

PROCTITIS, PROCTOCOLITIS, ENTEROCOLITIS, AND ENTERITIS

Sexually acquired *proctitis*, with inflammation limited to the rectal mucosa (the distal 10–12 cm), results from direct rectal inoculation of typical STD pathogens. In contrast, inflammation extending from the rectum to the colon (*proctocolitis*), involving both the small and the large bowel (*enterocolitis*), or involving the small bowel alone (*enteritis*) can result from ingestion of typical intestinal pathogens through oral–anal exposure during sexual contact. Anorectal pain and mucopurulent, bloody rectal discharge suggest proctitis or proctocolitis. Proctitis commonly produces tenesmus (causing frequent attempts to defecate, but not true diarrhea) and constipation, whereas proctocolitis and enterocolitis more often cause true diarrhea. In all three conditions, anoscopy usually shows mucosal exudate and easily induced mucosal bleeding (i.e., a positive "wipe test"), sometimes with petechiae or mucosal ulcers. Exudate should be sampled for Gram's staining and other microbiologic studies. Sigmoidoscopy or colonoscopy shows inflammation limited to the rectum in proctitis or disease extending at least up into the sigmoid colon in proctocolitis.

The AIDS era brought an extraordinary shift in the clinical and etiologic spectrum of intestinal infections among MSM. The number of cases of the acute intestinal STIs described above fell as high-risk sexual behaviors became less common in this group. At the same time, the number of AIDS-related opportunistic intestinal infections increased rapidly, many associated with chronic or recurrent symptoms. The incidence of these opportunistic infections has since fallen with increasingly widespread coverage of HIV-infected persons with effective antiretroviral therapy. Two species initially isolated in association with intestinal symptoms in MSM—now known as *Helicobacter cinaedi* and *H. fennelliae*—have both been isolated from the blood of HIV-infected men and other immunosuppressed persons, often in association with a syndrome of multifocal dermatitis and arthritis.

Acquisition of HSV, *N. gonorrhoeae*, or *C. trachomatis* (including LGV strains of *C. trachomatis*) during receptive anorectal intercourse causes most cases of infectious proctitis in women and MSM. Primary and secondary syphilis can also produce anal or anorectal lesions, with or without symptoms. Gonococcal or chlamydial proctitis typically involves the most distal rectal mucosa and the anal crypts and is clinically mild, without systemic manifestations. In contrast, primary proctitis due to HSV and proctocolitis due to the strains of *C. trachomatis* that cause LGV usually produce severe anorectal pain and often cause fever. Perianal ulcers and inguinal lymphadenopathy, most commonly due to HSV, can also occur with LGV or syphilis. Sacral nerve root radiculopathies, usually presenting as urinary retention, laxity of the anal sphincter, or constipation, may complicate primary herpetic proctitis. In LGV, rectal biopsy typically shows crypt abscesses, granulomas, and giant cells—findings resembling those in Crohn's disease; such findings should always prompt rectal culture and serology for LGV, which is a curable infection. Syphilis can also produce rectal granulomas, usually in association with infiltration by plasma cells or other mononuclear cells. Syphilis, LGV, and HSV infection involving the rectum can produce perirectal adenopathy that is sometimes mistaken for malignancy; syphilis, LGV, HSV infection, and chancroid involving the anus can produce inguinal adenopathy because anal lymphatics drain to inguinal lymph nodes.

Diarrhea and abdominal bloating or cramping pain without anorectal symptoms and with normal findings on anoscopy and sigmoidoscopy occur with inflammation of the small intestine (enteritis) or with proximal colitis. In MSM without HIV infection, enteritis is often attributable to *Giardia lamblia*. Sexually acquired proctocolitis is most often due to *Campylobacter* or *Shigella* species.

TREATMENT PROCTITIS, PROCTOCOLITIS, ENTEROCOLITIS, AND ENTERITIS

Acute proctitis in persons who have practiced receptive anal intercourse is usually sexually acquired. Such patients should undergo anoscopy to detect rectal ulcers or vesicles and petechiae after swabbing of the rectal mucosa; to examine rectal exudates for PMNs and gram-negative diplococci; and to obtain rectal swab specimens for testing for rectal gonorrhea, chlamydial infection, herpes, and syphilis. Pending test results, patients with proctitis should receive empirical syndromic treatment—e.g., with ceftriaxone (a single IM dose of 250 mg for gonorrhea) plus doxycycline (100 mg by mouth twice daily for 7 days) for possible chlamydial infection plus treatment for herpes or syphilis if indicated. If LGV proctitis is proven or suspected, the recommended treatment is doxycycline (100 mg by mouth twice daily for 21 days); alternatively, 1 g of azithromycin once a week for 3 weeks is likely to be effective but is little studied.

PREVENTION AND CONTROL OF STIs

Prevention and control of STIs require the following:

1. Reduction of the average rate of sexual exposure to STIs through alteration of sexual risk behaviors and behavioral norms among both susceptible and infected persons in all population groups. The necessary changes include reduction in the total number of sexual partners and the number of concurrent sexual partners.
2. Reduction of the efficiency of transmission through the promotion of safer sexual practices, the use of condoms during casual or commercial sex, vaccination against HBV and HPV infection, male circumcision (which reduces risk of acquisition of HIV infection, chancroid, and perhaps other STIs), and a growing number of other approaches (e.g., early detection and treatment of other STIs to reduce the efficiency of sexual transmission of HIV). Longitudinal studies have shown that consistent condom use is associated with significant protection of both males and females against all STIs that have been examined, including HIV, HPV, and HSV infections as well as gonorrhea and chlamydial infection. The only exceptions are probably sexually transmitted *Pthirus pubis* and *Sarcoptes scabiei* infestations.
3. Shortening of the duration of infectivity of STIs through early detection and curative or suppressive treatment of patients and their sexual partners.

Financial and time constraints imposed by many clinical practices, along with the reluctance of some clinicians to ask questions about stigmatized sexual behaviors, often curtail screening and prevention services. As outlined in Fig. 163-8, the success of clinicians' efforts to detect and treat STIs depends in part on societal efforts to teach young people how to recognize symptoms of STIs; to motivate individuals with symptoms to seek care promptly; to educate persons who are at risk but have no symptoms about what tests they should undergo routinely; and to make high-quality, appropriate care accessible, affordable, and acceptable, especially to the young indigent patients most likely to acquire an STI.

Because many infected individuals develop no symptoms or fail to recognize and report symptoms, clinicians should routinely perform an STI risk assessment for teenagers and young adults as a guide to selective screening. As stated earlier, U.S. Preventive Services Task Force Guidelines recommend screening sexually active female patients ≤ 25 years of age for *C. trachomatis* whenever they present for health care (at least once a year); older women should be tested if they have more than one sexual partner, have begun a new sexual relationship since the previous test, or have another STI diagnosed. In women 25–29 years of age, chlamydial infection is uncommon but still may reach a prevalence of 3–5% in some settings; information provided by women in this age group on a sex partner's concurrency (whether

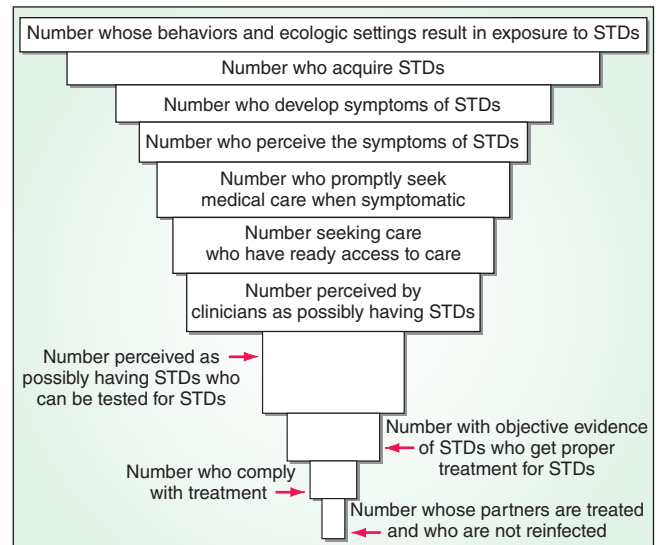


FIGURE 163-8 Critical control points for preventive and clinical interventions against sexually transmitted diseases (STDs).

(Adapted from HT Waller and MA Piot: *Bull World Health Organ* 41:75, 1969 and 43:1, 1970; and from "Resource allocation model for public health planning—a case study of tuberculosis control," *Bull World Health Organ* 48 [Suppl], 1973.)

a male partner has had another sex partner during the time they have been together) is helpful in identifying women at increased risk. In some regions of the United States, widespread selective screening and treatment of young women for cervical *C. trachomatis* infection have been associated with a 50–60% drop in prevalence. Such screening and treatment also protect the individual woman from PID. Sensitive urine-based genetic amplification tests permit expansion of screening to men, teenage boys, and girls in settings where examination is not planned or is impractical (e.g., during preparticipation sports examinations or during initial medical evaluation of adolescent girls). Vaginal swabs—collected either by the health care provider at a pelvic examination or by the woman herself—are highly sensitive and specific for the diagnosis of chlamydial and gonococcal infection; they are now the preferred type of specimen for screening and diagnosis of these infections.

Although gonorrhea is now substantially less common than chlamydial infection in industrialized countries, screening tests for *N. gonorrhoeae* are still appropriate for women and teenage girls attending STD clinics and for sexually active teens and young women from areas of high gonorrhea prevalence. Multiplex NAATs that combine screening for *N. gonorrhoeae* and *C. trachomatis*—and, more recently, for *T. vaginalis*—in a single low-cost assay now facilitate the prevention and control of these infections for populations at high risk.

All patients who have newly detected STIs or are at high risk for STIs according to routine risk assessment as well as all pregnant women should be encouraged to undergo serologic testing for syphilis and HIV infection, with appropriate HIV counseling before and after testing. Randomized trials have shown that risk-reduction counseling of patients with STIs significantly lowers subsequent risk of acquiring an STI; such counseling should now be considered a standard component of STI management. Preimmunization serologic testing for antibody to HBV is indicated for unvaccinated persons who are known to be at high risk, such as MSM and people who use injection drugs. In most young persons, however, it is more cost-effective to vaccinate against HBV without serologic screening. It is important to recognize that, while immunization against HBV has contributed to marked reductions in the incidence of infection with this virus, the majority of new cases that do occur are acquired through sex. In 2006, the Advisory Committee on Immunization Practices (ACIP) of the CDC recommended the following: (1) Universal hepatitis B vaccination should be implemented for all unvaccinated adults in

settings in which a high proportion of adults have risk factors for HBV infection (e.g., STD clinics, HIV testing and treatment facilities, drug-abuse treatment and prevention settings, health care settings targeting services to injection drug users or MSM, and correctional facilities). (2) In other primary care and specialty medical settings in which adults at risk for HBV infection receive care, health care providers should inform all patients about the health benefits of vaccination, the risk factors for HBV infection, and the persons for whom vaccination is recommended and should vaccinate adults who report risk factors for HBV infection as well as any adult who requests protection from HBV infection. To promote vaccination in all settings, health care providers should implement standing orders to identify adults recommended for hepatitis B vaccination, should administer hepatitis B vaccine as part of routine clinical services, should not require acknowledgment of an HBV infection risk factor for adult vaccination, and should use available reimbursement mechanisms to remove financial barriers to hepatitis B vaccination.

In 2007, the ACIP recommended routine immunization of 9- to 26-year-old girls and women with the quadrivalent HPV vaccine (against HPV types 6, 11, 16, and 18) approved by the U.S. Food and Drug Administration; the optimal age for recommended vaccination is 11–12 years because of the very high risk of HPV infection after sexual debut. In 2009, the ACIP added bivalent HPV vaccine (against types 6 and 11) as an option and expanded the groups in which immunization (with either quadrivalent or bivalent vaccine) is safe and effective to include boys and men 9–26 years old. HPV vaccines offering broader protection against additional oncogenic HPV types are anticipated. Since 2011, the ACIP has recommended routine administration of quadrivalent HPV vaccine to boys at 11 or 12 years of age and to males 13–21 years of age who have not yet been vaccinated or who have not completed the three-dose vaccine series; men 22–26 years of age may also be vaccinated.

Partner notification is the process of identifying and informing partners of infected patients about possible exposure to an STI and of examining, testing, and treating partners as appropriate. In a series of 22 reports concerning partner notification during the 1990s, index patients with gonorrhea or chlamydial infection named a mean of 0.75–1.6 partners, of whom one-fourth to one-third were infected; those with syphilis named 1.8–6.3 partners, with one-third to one-half infected; and those with HIV infection named 0.76–5.31 partners, with up to one-fourth infected. Persons who transmit infection or who have recently been infected and are still in the incubation period usually have no symptoms or only mild symptoms and seek medical attention only when notified of their exposure. Therefore, the clinician must encourage patients to participate in partner notification, must ensure that exposed persons are notified and treated, and must guarantee confidentiality to all involved. In the United States, local health departments often offer assistance in partner notification, treatment, and/or counseling. It seems both feasible and most useful to notify those partners exposed within the patient's likely period of infectiousness, which is often considered the preceding 1 month for gonorrhea, 1–2 months for chlamydial infection, and up to 3 months for early syphilis.

Persons with a new-onset STI always have a *source* contact who gave them the infection; in addition, they may have a *secondary* (*spread* or *exposed*) contact with whom they had sex after becoming infected. The identification and treatment of these two types of contacts have different objectives. Treatment of the source contact (often a casual contact) benefits the community by preventing further transmission and benefits the source contact; treatment of the recently exposed secondary contact (typically a spouse or another steady sexual partner) prevents the development of serious complications (such as PID) in the partner, reinfection of the index patient, and further spread of infection. A survey of a random sample of U.S. physicians found that most instructed patients to abstain from sex during treatment, to use condoms, and to inform their sex partners after being diagnosed with gonorrhea, chlamydial infection, or syphilis; physicians sometimes gave the patients drugs for their partners. However, follow-up of the partners by physicians was infrequent. A randomized trial compared patients' delivery of therapy to partners exposed to gonorrhea or chlamydial infection with

conventional notification and advice to partners to seek evaluation for STD; patients' delivery of partners' therapy, also known as *expedited partner therapy* (EPT), significantly reduced combined rates of reinfection of the index patient with *N. gonorrhoeae* or *C. trachomatis*. State-by-state variations in regulations governing this approach have not been well defined, but the 2010 CDC STD treatment guidelines and the EPT final report of 2006 (<http://www.cdc.gov/std/treatment/EPTFinalReport2006.pdf>) describe its potential use. Currently, EPT is commonly used by many practicing physicians. Its legal status varies by state, but EPT is now permissible in 38 states and potentially allowable in another 9. (Updated information on the legal status of EPT is available at <http://www.cdc.gov/std/ept>.)

In summary, clinicians and public health agencies share responsibility for the prevention and control of STIs. In the current health care environment, the role of primary care clinicians has become increasingly important in STI prevention as well as in diagnosis and treatment, and the resurgence of bacterial STIs like syphilis and LGV among MSM—particularly those co-infected with HIV—emphasizes the need for risk assessment and routine screening.

164 Meningitis, Encephalitis, Brain Abscess, and Empyema

Karen L. Roos, Kenneth L. Tyler

Acute infections of the nervous system are among the most important problems in medicine because early recognition, efficient decision making, and rapid institution of therapy can be lifesaving. These distinct clinical syndromes include acute bacterial meningitis, viral meningitis, encephalitis, focal infections such as brain abscess and subdural empyema, and infectious thrombophlebitis. Each may present with a nonspecific prodrome of fever and headache, which in a previously healthy individual may initially be thought to be benign, until (with the exception of viral meningitis) altered consciousness, focal neurologic signs, or seizures appear. Key goals of early management are to emergently distinguish between these conditions, identify the responsible pathogen, and initiate appropriate antimicrobial therapy.

APPROACH TO THE PATIENT: Meningitis, Encephalitis, Brain Abscess, and Empyema

(**Figure 164-1**) The first task is to identify whether an infection predominantly involves the subarachnoid space (*meningitis*) or whether there is evidence of either generalized or focal involvement of brain tissue in the cerebral hemispheres, cerebellum, or brainstem. When brain tissue is directly injured by a bacterial or viral infection, the disease is referred to as *encephalitis*, whereas focal infections involving brain tissue are classified as either *cerebritis* or *abscess*, depending on the presence or absence of a capsule.

Nuchal rigidity (“stiff neck”) is the pathognomonic sign of meningeal irritation and is present when the neck resists passive flexion. Kernig’s and Brudzinski’s signs are also classic signs of meningeal irritation. *Kernig’s sign* is elicited with the patient in the supine position. The thigh is flexed on the abdomen, with the knee flexed; attempts to passively extend the knee elicit pain when meningeal irritation is present. *Brudzinski’s sign* is elicited with the patient in the supine position and is positive when passive flexion of the neck results in spontaneous flexion of the hips and knees. Although commonly tested on physical examinations, the sensitivity and specificity of Kernig’s and Brudzinski’s signs are uncertain. Both may be absent or reduced in very young or elderly patients, immunocompromised individuals, or patients with a severely depressed

mental status. The high prevalence of cervical spine disease in older individuals may result in false-positive tests for nuchal rigidity.

Initial management can be guided by several considerations: (1) Empirical therapy should be initiated promptly whenever bacterial meningitis is a significant diagnostic consideration. (2) All patients who have had recent head trauma, are immunocompromised, have known malignant lesions or central nervous system (CNS) neoplasms, or have focal neurologic findings, papilledema, or a depressed level of consciousness should undergo computed tomography (CT) or magnetic resonance imaging (MRI) of the brain prior to lumbar puncture (LP). In these cases empirical antibiotic therapy should not be delayed pending test results but should be administered prior to neuroimaging and LP. (3) A significantly depressed

level of consciousness (e.g., somnolence, coma), seizures, or focal neurologic deficits do not occur in viral meningitis; patients with these symptoms should be hospitalized for further evaluation and treated empirically for bacterial and viral meningoencephalitis. (4) Immunocompetent patients with a normal level of consciousness, no prior antimicrobial treatment, and a cerebrospinal fluid (CSF) profile consistent with viral meningitis (lymphocytic pleocytosis and a normal glucose concentration) can often be treated as outpatients if appropriate contact and monitoring can be ensured. Failure of a patient with suspected viral meningitis to improve within 48 h should prompt a reevaluation including follow-up neurologic and general medical examination and repeat imaging and laboratory studies, often including a second LP.

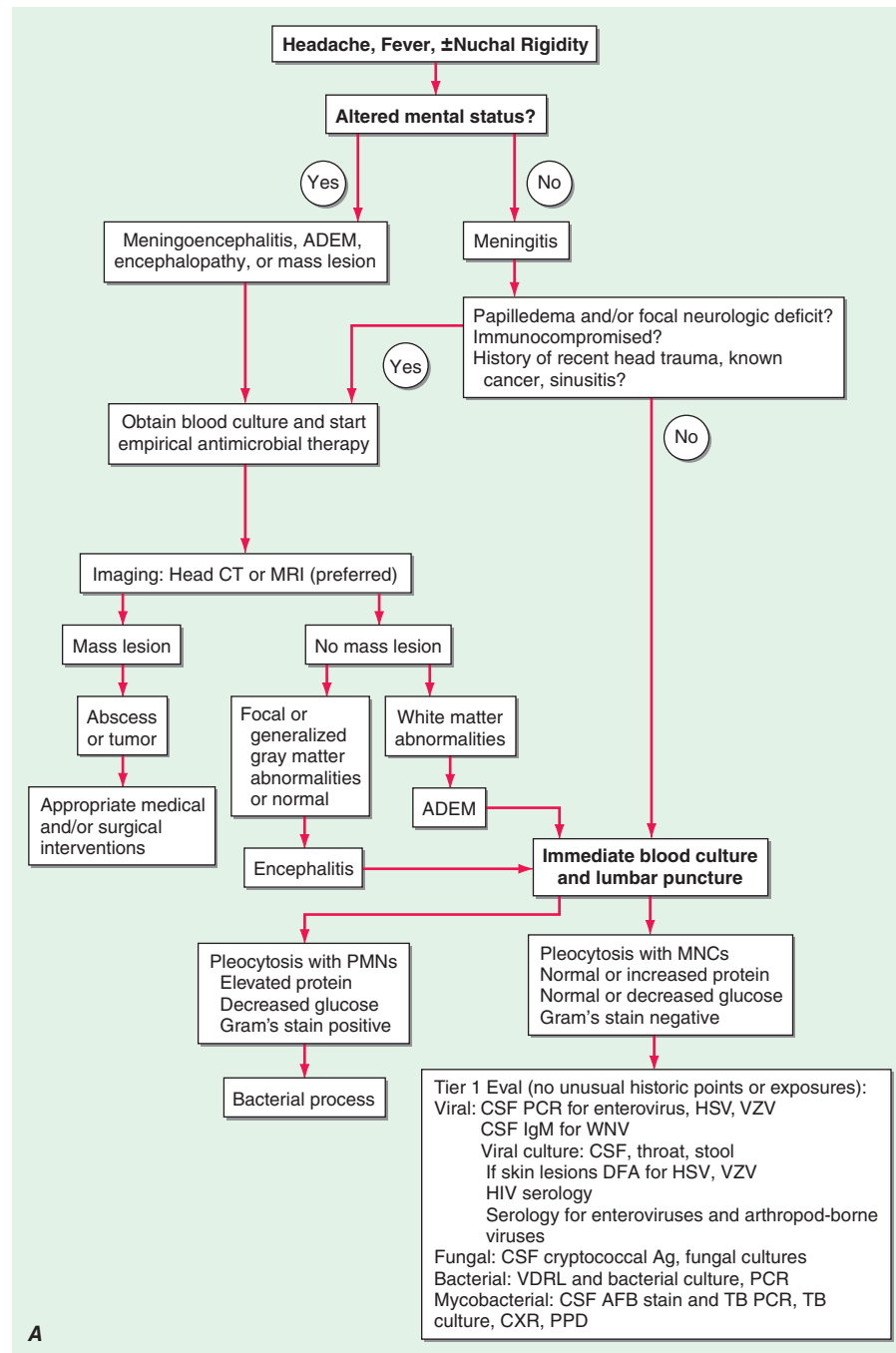


FIGURE 164-1 The management of patients with suspected central nervous system (CNS) infection. ADEM, acute disseminated encephalomyelitis; AFB, acid-fast bacillus; Ag, antigen; CSF, cerebrospinal fluid; CT, computed tomography; CTFV, Colorado tick fever virus; CXR, chest x-ray; DFA, direct fluorescent antibody; EBV, Epstein-Barr virus; HHV, human herpesvirus; HSV, herpes simplex virus; LCMV, lymphocytic choriomeningitis virus; MNCs, mononuclear cells; MRI, magnetic resonance imaging; PCR, polymerase chain reaction; PMNs, polymorphonuclear leukocytes; PPD, purified protein derivative; TB, tuberculosis; VDRL, Venereal Disease Research Laboratory; VZV, varicella-zoster virus; WNV, West Nile virus.

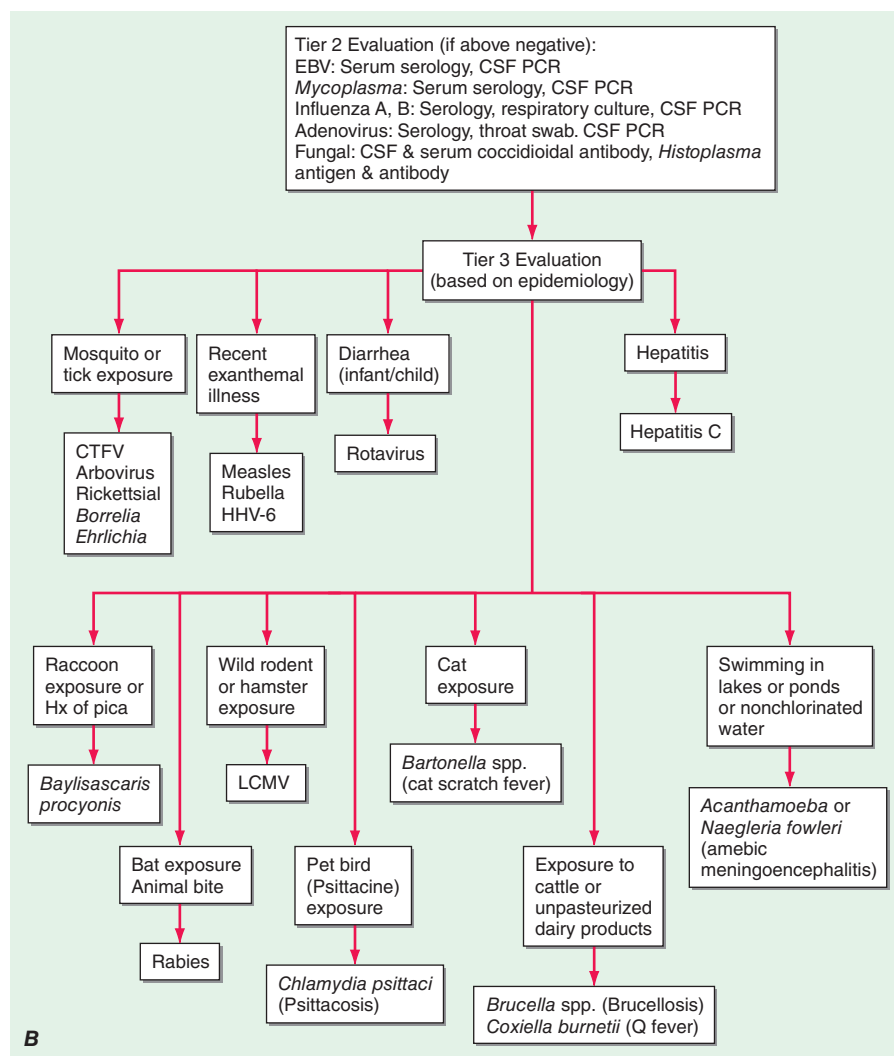


FIGURE 164-1 (Continued)

ACUTE BACTERIAL MENINGITIS

DEFINITION

Bacterial meningitis is an acute purulent infection within the subarachnoid space. It is associated with a CNS inflammatory reaction that may result in decreased consciousness, seizures, raised intracranial pressure (ICP), and stroke. The meninges, subarachnoid space, and brain parenchyma are all frequently involved in the inflammatory reaction (*meningoencephalitis*).

EPIDEMIOLOGY

Bacterial meningitis is the most common form of suppurative CNS infection, with an annual incidence in the United States of >2.5 cases/100,000 population. The organisms most often responsible for community-acquired bacterial meningitis are *Streptococcus pneumoniae* (~50%), *Neisseria meningitidis* (~25%), group B streptococci (~15%), and *Listeria monocytogenes* (~10%). *Haemophilus influenzae* type b accounts for <10% of cases of bacterial meningitis in most series. *N. meningitidis* is the causative organism of recurring epidemics of meningitis every 8 to 12 years.

ETIOLOGY

S. pneumoniae (Chap. 173) is the most common cause of meningitis in adults >20 years of age, accounting for nearly half the reported cases (1.1 per 100,000 persons per year). There are a number of predisposing conditions that increase the risk of pneumococcal meningitis, the most important of which is pneumococcal pneumonia. Additional risk factors include coexisting acute or chronic pneumococcal sinusitis or

otitis media, alcoholism, diabetes, splenectomy, hypogammaglobulinemia, complement deficiency, and head trauma with basilar skull fracture and CSF rhinorrhea. The mortality rate remains ~20% despite antibiotic therapy.

The incidence of meningitis due to *N. meningitidis* (Chap. 180) has decreased with the routine immunization of 11- to 18-year-olds with the quadrivalent (serogroups A, C, W-135, and Y) meningococcal glycoconjugate vaccine. The vaccine does not contain serogroup B, which is responsible for one-third of cases of meningococcal disease. The presence of petechial or purpuric skin lesions can provide an important clue to the diagnosis of meningococcal infection. In some patients the disease is fulminant, progressing to death within hours of symptom onset. Infection may be initiated by nasopharyngeal colonization, which can result in either an asymptomatic carrier state or invasive meningococcal disease. The risk of invasive disease following nasopharyngeal colonization depends on both bacterial virulence factors and host immune defense mechanisms, including the host's capacity to produce antimeningococcal antibodies and to lyse meningococci by both classic and alternative complement pathways. Individuals with deficiencies of any of the complement components, including properdin, are highly susceptible to meningococcal infections.

Gram-negative bacilli cause meningitis in individuals with chronic and debilitating diseases such as diabetes, cirrhosis, or alcoholism and in those with chronic urinary tract infections. Gram-negative meningitis can also complicate neurosurgical procedures, particularly craniotomy, and head trauma associated with CSF rhinorrhea or otorrhea.

Otitis, mastoiditis, and sinusitis are predisposing and associated conditions for meningitis due to *Streptococci* sp., gram-negative anaerobes,

886 *Staphylococcus aureus*, *Haemophilus* sp., and Enterobacteriaceae. Meningitis complicating endocarditis may be due to viridans streptococci, *S. aureus*, *Streptococcus bovis*, the HACEK group (*Haemophilus* sp., *Actinobacillus actinomycetemcomitans*, *Cardiobacterium hominis*, *Eikenella corrodens*, *Kingella kingae*), or enterococci.

Group B *Streptococcus*, or *Streptococcus agalactiae*, was previously responsible for meningitis predominantly in neonates, but it has been reported with increasing frequency in individuals >50 years of age, particularly those with underlying diseases.

L. monocytogenes (Chap. 176) is an increasingly important cause of meningitis in neonates (<1 month of age), pregnant women, individuals >60 years, and immunocompromised individuals of all ages. Infection is acquired by ingesting foods contaminated by *Listeria*. Foodborne human listerial infection has been reported from contaminated coleslaw, milk, soft cheeses, and several types of “ready-to-eat” foods, including delicatessen meat and uncooked hotdogs.

The frequency of *H. influenzae* type b (Hib) meningitis in children has declined dramatically since the introduction of the Hib conjugate vaccine, although rare cases of Hib meningitis in vaccinated children have been reported. More frequently, *H. influenzae* causes meningitis in unvaccinated children and older adults, and non-b *H. influenzae* is an emerging pathogen.

S. aureus and coagulase-negative staphylococci (Chap. 172) are important causes of meningitis that occurs following invasive neurosurgical procedures, particularly shunting procedures for hydrocephalus, or as a complication of the use of subcutaneous Ommaya reservoirs for administration of intrathecal chemotherapy.

and complications of bacterial meningitis result from the immune response to the invading pathogen rather than from direct bacteria-induced tissue injury. As a result, neurologic injury can progress even after the CSF has been sterilized by antibiotic therapy.

The lysis of bacteria with the subsequent release of cell-wall components into the subarachnoid space is the initial step in the induction of the inflammatory response and the formation of a purulent exudate in the subarachnoid space (Fig. 164-2). Bacterial cell-wall components, such as the lipopolysaccharide (LPS) molecules of gram-negative bacteria and teichoic acid and peptidoglycans of *S. pneumoniae*, induce meningeal inflammation by stimulating the production of inflammatory cytokines and chemokines by microglia, astrocytes, monocytes, microvascular endothelial cells, and CSF leukocytes. In experimental models of meningitis, cytokines including tumor necrosis factor alpha (TNF- α) and interleukin 1 β (IL-1 β) are present in CSF within 1–2 h of intracisternal inoculation of LPS. This cytokine response is quickly followed by an increase in CSF protein concentration and leukocytosis. Chemokines (cytokines that induce chemotactic migration in leukocytes) and a variety of other proinflammatory cytokines are also produced and secreted by leukocytes and tissue cells that are stimulated by IL-1 β and TNF- α . In addition, bacteremia and the inflammatory cytokines induce the production of excitatory amino acids, reactive oxygen and nitrogen species (free oxygen radicals, nitric oxide, and peroxynitrite), and other mediators that can induce death of brain cells, especially in the dentate gyrus of the hippocampus.

Much of the pathophysiology of bacterial meningitis is a direct consequence of elevated levels of CSF cytokines and chemokines. TNF- α and

PATHOPHYSIOLOGY

The most common bacteria that cause meningitis, *S. pneumoniae* and *N. meningitidis*, initially colonize the nasopharynx by attaching to nasopharyngeal epithelial cells. Bacteria are transported across epithelial cells in membrane-bound vacuoles to the intravascular space or invade the intravascular space by creating separations in the apical tight junctions of columnar epithelial cells. Once in the bloodstream, bacteria are able to avoid phagocytosis by neutrophils and classic complement-mediated bactericidal activity because of the presence of a polysaccharide capsule. Bloodborne bacteria can reach the intraventricular choroid plexus, directly infect choroid plexus epithelial cells, and gain access to the CSF. Some bacteria, such as *S. pneumoniae*, can adhere to cerebral capillary endothelial cells and subsequently migrate through or between these cells to reach the CSF. Bacteria are able to multiply rapidly within CSF because of the absence of effective host immune defenses. Normal CSF contains few white blood cells (WBCs) and relatively small amounts of complement proteins and immunoglobulins. The paucity of the latter two prevents effective opsonization of bacteria, an essential prerequisite for bacterial phagocytosis by neutrophils. Phagocytosis of bacteria is further impaired by the fluid nature of CSF, which is less conducive to phagocytosis than a solid tissue substrate.

A critical event in the pathogenesis of bacterial meningitis is the inflammatory reaction induced by the invading bacteria. Many of the neurologic manifestations

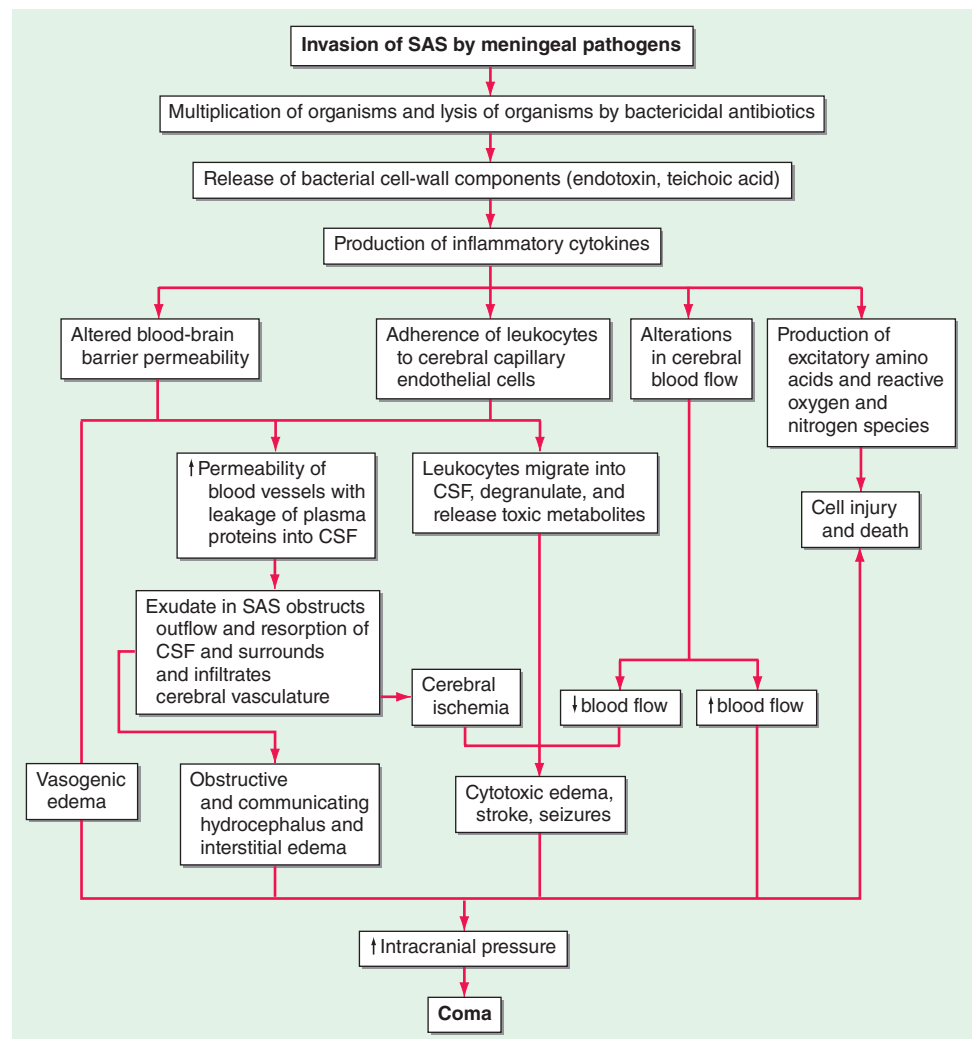


FIGURE 164-2 The pathophysiology of the neurologic complications of bacterial meningitis. CSF, cerebrospinal fluid; SAS, subarachnoid space.

IL-1 β act synergistically to increase the permeability of the blood-brain barrier, resulting in induction of vasogenic edema and the leakage of serum proteins into the subarachnoid space (Fig. 164-2). The subarachnoid exudate of proteinaceous material and leukocytes obstructs the flow of CSF through the ventricular system and diminishes the resorptive capacity of the arachnoid granulations in the dural sinuses, leading to obstructive and communicating hydrocephalus and concomitant interstitial edema.

Inflammatory cytokines upregulate the expression of selectins on cerebral capillary endothelial cells and leukocytes, promoting leukocyte adherence to vascular endothelial cells and subsequent migration into the CSF. The adherence of leukocytes to capillary endothelial cells increases the permeability of blood vessels, allowing for the leakage of plasma proteins into the CSF, which adds to the inflammatory exudate. Neutrophil degranulation results in the release of toxic metabolites that contribute to cytotoxic edema, cell injury, and death. Contrary to previous beliefs, CSF leukocytes probably do little to contribute to the clearance of CSF bacterial infection.

During the very early stages of meningitis, there is an increase in cerebral blood flow, soon followed by a decrease in cerebral blood flow and a loss of cerebrovascular autoregulation (Chap. 330). Narrowing of the large arteries at the base of the brain due to encroachment by the purulent exudate in the subarachnoid space and infiltration of the arterial wall by inflammatory cells with intimal thickening (*vasculitis*) also occur and may result in ischemia and infarction, obstruction of branches of the middle cerebral artery by thrombosis, thrombosis of the major cerebral venous sinuses, and thrombophlebitis of the cerebral cortical veins. The combination of interstitial, vasogenic, and cytotoxic edema leads to raised ICP and coma. Cerebral herniation usually results from the effects of cerebral edema, either focal or generalized; hydrocephalus and dural sinus or cortical vein thrombosis may also play a role.

CLINICAL PRESENTATION

Meningitis can present as either an acute fulminant illness that progresses rapidly in a few hours or as a subacute infection that progressively worsens over several days. The classic triad of meningitis is fever, headache, and nuchal rigidity, but the classic triad may not be present. A decreased level of consciousness occurs in >75% of patients and can vary from lethargy to coma. Fever and either headache, stiff neck, or an altered level of consciousness will be present in nearly every patient with bacterial meningitis. Nausea, vomiting, and photophobia are also common complaints.

Seizures occur as part of the initial presentation of bacterial meningitis or during the course of the illness in 20–40% of patients. Focal seizures are usually due to focal arterial ischemia or infarction, cortical venous thrombosis with hemorrhage, or focal edema. Generalized seizure activity and status epilepticus may be due to hyponatremia, cerebral anoxia, or, less commonly, the toxic effects of antimicrobial agents.

Raised ICP is an expected complication of bacterial meningitis and the major cause of obtundation and coma in this disease. More than 90% of patients will have a CSF opening pressure >180 mmH₂O, and 20% have opening pressures >400 mmH₂O. Signs of increased ICP include a deteriorating or reduced level of consciousness, papilledema, dilated poorly reactive pupils, sixth nerve palsies, decerebrate posturing, and the Cushing reflex (bradycardia, hypertension, and irregular respirations). The most disastrous complication of increased ICP is cerebral herniation. The incidence of herniation in patients with bacterial meningitis has been reported to occur in as few as 1% to as many as 8% of cases.

Specific clinical features may provide clues to the diagnosis of individual organisms and are discussed in more detail in specific chapters devoted to individual pathogens. The most important of these clues is the rash of meningococemia, which begins as a diffuse erythematous maculopapular rash resembling a viral exanthem; however, the skin lesions of meningococemia rapidly become petechial. Petechiae are found on the trunk and lower extremities, in the mucous membranes and conjunctiva, and occasionally on the palms and soles.

DIAGNOSIS

When bacterial meningitis is suspected, blood cultures should be immediately obtained and empirical antimicrobial and adjunctive dexamethasone therapy initiated without delay (Table 164-1). The diagnosis of bacterial meningitis is made by examination of the CSF (Table 164-2). The need to obtain neuroimaging studies (CT or MRI) prior to LP requires clinical judgment. In an immunocompetent patient with no known history of recent head trauma, a normal level of consciousness, and no evidence of papilledema or focal neurologic deficits, it is considered safe to perform LP without prior neuroimaging studies. If LP is delayed in order to obtain neuroimaging studies, empirical antibiotic therapy should be initiated after blood cultures are obtained. Antibiotic therapy initiated a few hours prior to LP will not significantly alter the CSF WBC count or glucose concentration, nor is it likely to prevent visualization of organisms by Gram's stain or detection of bacterial nucleic acid by polymerase chain reaction (PCR) assay.

The classic CSF abnormalities in bacterial meningitis (Table 164-2) are (1) polymorphonuclear (PMN) leukocytosis (>100 cells/ μ L in 90%), (2) decreased glucose concentration (<2.2 mmol/L [<40 mg/dL] and/or CSF/serum glucose ratio of <0.4 in ~60%), (3) increased protein concentration (>0.45 g/L [>45 mg/dL] in 90%), and (4) increased opening pressure (>180 mmH₂O in 90%). CSF bacterial cultures are positive in >80% of patients, and CSF Gram's stain demonstrates organisms in >60%.

CSF glucose concentrations <2.2 mmol/L (<40 mg/dL) are abnormal, and a CSF glucose concentration of zero can be seen in bacterial meningitis. Use of the CSF/serum glucose ratio corrects for hyperglycemia

TABLE 164-1 ANTIBIOTICS USED IN EMPIRICAL THERAPY OF BACTERIAL MENINGITIS AND FOCAL CENTRAL NERVOUS SYSTEM INFECTIONS^a

Indication	Antibiotic
Preterm infants to infants <1 month	Ampicillin + cefotaxime
Infants 1–3 months	Ampicillin + cefotaxime or ceftriaxone
Immunocompetent children >3 months and adults <55	Cefotaxime, ceftriaxone, or cefepime + vancomycin
Adults >55 and adults of any age with alcoholism or other debilitating illnesses	Ampicillin + cefotaxime, ceftriaxone or cefepime + vancomycin
Hospital-acquired meningitis, posttraumatic or postneurosurgery meningitis, neutropenic patients, or patients with impaired cell-mediated immunity	Ampicillin + ceftazidime or meropenem + vancomycin

Antimicrobial Agent	Total Daily Dose and Dosing Interval	
	Child (>1 month)	Adult
Ampicillin	300 (mg/kg)/d, q6h	12 g/d, q4h
Cefepime	150 (mg/kg)/d, q8h	6 g/d, q8h
Cefotaxime	225–300 (mg/kg)/d, q6h	12 g/d, q4h
Ceftriaxone	100 (mg/kg)/d, q12h	4 g/d, q12h
Ceftazidime	150 (mg/kg)/d, q8h	6 g/d, q8h
Gentamicin	7.5 (mg/kg)/d, q8h ^b	7.5 (mg/kg)/d, q8h
Meropenem	120 (mg/kg)/d, q8h	6 g/d, q8h
Metronidazole	30 (mg/kg)/d, q6h	1500–2000 mg/d, q6h
Nafcillin	100–200 (mg/kg)/d, q6h	9–12 g/d, q4h
Penicillin G	400,000 (U/kg)/d, q4h	20–24 million U/d, q4h
Vancomycin	45–60 (mg/kg)/d, q6h	45–60 (mg/kg)/d, q6–12h ^b

^aAll antibiotics are administered intravenously; doses indicated assume normal renal and hepatic function. ^bDoses should be adjusted based on serum peak and trough levels: gentamicin therapeutic level: peak: 5–8 μ g/mL; trough: <2 μ g/mL; vancomycin therapeutic level: peak: 25–40 μ g/mL; trough: 5–15 μ g/mL.

TABLE 164-2 CEREBROSPINAL FLUID (CSF) ABNORMALITIES IN BACTERIAL MENINGITIS

Opening pressure	>180 mmH ₂ O
White blood cells	10/μL to 10,000/μL; neutrophils predominate
Red blood cells	Absent in nontraumatic tap
Glucose	<2.2 mmol/L (<40 mg/dL)
CSF/serum glucose	<0.4
Protein	>0.45 g/L (>45 mg/dL)
Gram's stain	Positive in >60%
Culture	Positive in >80%
Latex agglutination	May be positive in patients with meningitis due to <i>Streptococcus pneumoniae</i> , <i>Neisseria meningitidis</i> , <i>Haemophilus influenzae</i> type b, <i>Escherichia coli</i> , group B streptococci
Limulus lysate	Positive in cases of gram-negative meningitis
PCR	Detects bacterial DNA

Abbreviation: PCR, polymerase chain reaction

that may mask a relative decrease in the CSF glucose concentration. The CSF glucose concentration is low when the CSF/serum glucose ratio is <0.6. A CSF/serum glucose ratio <0.4 is highly suggestive of bacterial meningitis but may also be seen in other conditions, including fungal, tuberculous, and carcinomatous meningitis. It takes from 30 min to several hours for the concentration of CSF glucose to reach equilibrium with blood glucose levels; therefore, administration of 50 mL of 50% glucose (D50) prior to LP, as commonly occurs in emergency room settings, is unlikely to alter CSF glucose concentration significantly unless more than a few hours have elapsed between glucose administration and LP.

A 16S rRNA conserved sequence broad-based bacterial PCR can detect small numbers of viable and nonviable organisms in CSF and is expected to be useful for making a diagnosis of bacterial meningitis in patients who have been pretreated with oral or parenteral antibiotics and in whom Gram's stain and CSF culture are negative. When the broad-range PCR is positive, a PCR that uses specific bacterial primers to detect the nucleic acid of *S. pneumoniae*, *N. meningitidis*, *Escherichia coli*, *L. monocytogenes*, *H. influenzae*, and *S. agalactiae* can be obtained based on the clinical suspicion of the meningeal pathogen. The latex agglutination (LA) test for the detection of bacterial antigens of *S. pneumoniae*, *N. meningitidis*, *H. influenzae* type b, group B *Streptococcus*, and *E. coli* K1 strains in the CSF has been useful for making a diagnosis of bacterial meningitis but is being replaced by the CSF bacterial PCR assay. The CSF LA test has a specificity of 95–100% for *S. pneumoniae* and *N. meningitidis*, so a positive test is virtually diagnostic of bacterial meningitis caused by these organisms. However, the sensitivity of the CSF LA test is only 70–100% for detection of *S. pneumoniae* and 33–70% for detection of *N. meningitidis* antigens, so a negative test does not exclude infection by these organisms. The Limulus amebocyte lysate assay is a rapid diagnostic test for the detection of gram-negative endotoxin in CSF and thus for making a diagnosis of gram-negative bacterial meningitis. The test has a specificity of 85–100% and a sensitivity approaching 100%. Thus, a positive Limulus amebocyte lysate assay occurs in virtually all patients with gram-negative bacterial meningitis, but false positives may occur.

Almost all patients with bacterial meningitis will have neuroimaging studies performed during the course of their illness. MRI is preferred over CT because of its superiority in demonstrating areas of cerebral edema and ischemia. In patients with bacterial meningitis, diffuse meningeal enhancement is often seen after the administration of gadolinium. Meningeal enhancement is not diagnostic of meningitis but occurs in any CNS disease associated with increased blood-brain barrier permeability.

Petechial skin lesions, if present, should be biopsied. The rash of meningococcemia results from the dermal seeding of organisms with vascular endothelial damage, and biopsy may reveal the organism on Gram's stain.

DIFFERENTIAL DIAGNOSIS

Viral meningoencephalitis, and particularly herpes simplex virus (HSV) encephalitis, can mimic the clinical presentation of bacterial meningitis (see “Viral Encephalitis,” below). HSV encephalitis typically presents with headache, fever, altered consciousness, focal neurologic deficits (e.g., dysphasia, hemiparesis), and focal or generalized seizures. The findings on CSF studies, neuroimaging, and electroencephalogram (EEG) distinguish HSV encephalitis from bacterial meningitis. The typical CSF profile with viral CNS infections is a lymphocytic pleocytosis with a normal glucose concentration, in contrast to the PMN pleocytosis and hypoglycorrhachia characteristic of bacterial meningitis. MRI abnormalities (other than meningeal enhancement) are not seen in uncomplicated bacterial meningitis. By contrast, in HSV encephalitis, on T2-weighted, fluid-attenuated inversion recovery (FLAIR) and diffusion-weighted MRI images, high signal intensity lesions are seen in the orbitofrontal, anterior, and medial temporal lobes in the majority of patients within 48 h of symptom onset. Some patients with HSV encephalitis have a distinctive periodic pattern on EEG (see below).

Rickettsial disease can resemble bacterial meningitis (**Chap. 211**). Rocky Mountain spotted fever (RMSF) is transmitted by a tick bite and caused by the bacteria *Rickettsia rickettsii*. The disease may present acutely with high fever, prostration, myalgia, headache, nausea, and vomiting. Most patients develop a characteristic rash within 96 h of the onset of symptoms. The rash is initially a diffuse erythematous maculopapular rash that may be difficult to distinguish from that of meningococcemia. It progresses to a petechial rash, then to a purpuric rash, and if untreated, to skin necrosis or gangrene. The color of the lesions changes from bright red to very dark red, then yellowish-green to black. The rash typically begins in the wrist and ankles and then spreads distally and proximally within a matter of a few hours, involving the palms and soles. Diagnosis is made by immunofluorescent staining of skin biopsy specimens. Ehrlichiosis are also transmitted by a tick bite. These are small gram-negative coccobacilli of which two species cause human disease. *Anaplasma phagocytophilum* causes human granulocytic ehrlichiosis (anaplasmosis), and *Ehrlichia chaffeensis* causes human monocytic ehrlichiosis. The clinical and laboratory manifestations of the infections are similar. Patients present with fever, headache, confusion, nausea, and vomiting. Twenty percent of patients have a maculopapular or petechial rash. There is laboratory evidence of leukopenia, thrombocytopenia, and anemia, and mild to moderate elevations in alanine aminotransferases, alkaline phosphatase, and lactate dehydrogenase. Patients with RMSF and those with ehrlichial infections may have an altered level of consciousness ranging from mild lethargy to coma, confusion, focal neurologic signs, cranial nerve palsies, hyperreflexia, and seizures.

Focal suppurative CNS infections (see below), including subdural and epidural empyema and brain abscess, should also be considered, especially when focal neurologic findings are present. MRI should be performed promptly in all patients with suspected meningitis who have focal features, both to detect the intracranial infection and to search for associated areas of infection in the sinuses or mastoid bones.

A number of noninfectious CNS disorders can mimic bacterial meningitis. Subarachnoid hemorrhage (SAH; **Chap. 330**) is generally the major consideration. Other possibilities include chemical meningitis due to rupture of tumor contents into the CSF (e.g., from a cystic glioma or craniopharyngioma epidermoid or dermoid cyst); drug-induced hypersensitivity meningitis; carcinomatous or lymphomatous meningitis; meningitis associated with inflammatory disorders such as sarcoid, systemic lupus erythematosus (SLE), and Behçet's syndrome; pituitary apoplexy; and uveomeningitic syndromes (Vogt-Koyanagi-Harada syndrome).

On occasion, subacutely evolving meningitis (**Chap. 165**) may be considered in the differential diagnosis of acute meningitis. The principal causes include *Mycobacterium tuberculosis* (**Chap. 202**), *Cryptococcus neoformans* (**Chap. 239**), *Histoplasma capsulatum* (**Chap. 236**), *Coccidioides immitis* (**Chap. 237**), and *Treponema pallidum* (**Chap. 206**).

EMPIRICAL ANTIMICROBIAL THERAPY

(Table 164-1) Bacterial meningitis is a medical emergency. The goal is to begin antibiotic therapy within 60 min of a patient's arrival in the emergency room. Empirical antimicrobial therapy is initiated in patients with suspected bacterial meningitis before the results of CSF Gram's stain and culture are known. *S. pneumoniae* (Chap. 171) and *N. meningitidis* (Chap. 180) are the most common etiologic organisms of community-acquired bacterial meningitis. Due to the emergence of penicillin- and cephalosporin-resistant *S. pneumoniae*, empirical therapy of community-acquired suspected bacterial meningitis in children and adults should include a combination of dexamethasone, a third- or fourth-generation cephalosporin (e.g., ceftriaxone, cefotaxime, or cefepime), and vancomycin, plus acyclovir, as HSV encephalitis is the leading disease in the differential diagnosis, and doxycycline during tick season to treat tick-borne bacterial infections. Ceftriaxone or cefotaxime provides good coverage for susceptible *S. pneumoniae*, group B streptococci, and *H. influenzae* and adequate coverage for *N. meningitidis*. Cefepime is a broad-spectrum fourth-generation cephalosporin with in vitro activity similar to that of cefotaxime or ceftriaxone against *S. pneumoniae* and *N. meningitidis* and greater activity against *Enterobacter* species and *Pseudomonas aeruginosa*. In clinical trials, cefepime has been demonstrated to be equivalent to cefotaxime in the treatment of penicillin-sensitive pneumococcal and meningococcal meningitis, and it has been used successfully in some patients with meningitis due to *Enterobacter* species and *P. aeruginosa*. Ampicillin should be added to the empirical regimen for coverage of *L. monocytogenes* in individuals <3 months of age, those >55, or those with suspected impaired cell-mediated immunity because of chronic illness, organ transplantation, pregnancy, malignancy, or immunosuppressive therapy. Metronidazole is added to the empirical regimen to cover gram-negative anaerobes in patients with otitis, sinusitis, or mastoiditis. In hospital-acquired meningitis, and particularly meningitis following neurosurgical procedures, staphylococci and gram-negative organisms including *P. aeruginosa* are the most common etiologic organisms. In these patients, empirical therapy should include a combination of vancomycin and ceftazidime, cefepime, or meropenem. Ceftazidime, cefepime, or meropenem should be substituted for ceftriaxone or cefotaxime in neurosurgical patients and in neutropenic patients, because ceftriaxone and cefotaxime do not provide adequate activity against CNS infection with *P. aeruginosa*. Meropenem is a carbapenem antibiotic that is highly active in vitro against *L. monocytogenes*, has been demonstrated to be effective in cases of meningitis caused by *P. aeruginosa*, and shows good activity against penicillin-resistant pneumococci. In experimental pneumococcal meningitis, meropenem was comparable to ceftriaxone and inferior to vancomycin in sterilizing CSF cultures. The number of patients with bacterial meningitis enrolled in clinical trials of meropenem has not been sufficient to definitively assess the efficacy of this antibiotic.

SPECIFIC ANTIMICROBIAL THERAPY

Meningococcal Meningitis (Table 164-3) Although ceftriaxone and cefotaxime provide adequate empirical coverage for *N. meningitidis*, penicillin G remains the antibiotic of choice for meningococcal meningitis caused by susceptible strains. Isolates of *N. meningitidis* with moderate resistance to penicillin have been identified and are increasing in incidence worldwide. CSF isolates of *N. meningitidis* should be tested for penicillin and ampicillin susceptibility, and if resistance is found, cefotaxime or ceftriaxone should be substituted for penicillin. A 7-day course of intravenous antibiotic therapy is adequate for uncomplicated meningococcal meningitis. The index case and all close contacts should receive chemoprophylaxis with a 2-day regimen of rifampin (600 mg every 12 h for 2 days in adults and 10 mg/kg every 12 h for 2 days in children >1 year). Rifampin is not recommended in pregnant women. Alternatively, adults can be

TABLE 164-3 ANTIMICROBIAL THERAPY OF CENTRAL NERVOUS SYSTEM BACTERIAL INFECTIONS BASED ON PATHOGEN^a

Organism	Antibiotic
<i>Neisseria meningitidis</i>	
Penicillin-sensitive	Penicillin G or ampicillin
Penicillin-resistant	Ceftriaxone or cefotaxime
<i>Streptococcus pneumoniae</i>	
Penicillin-sensitive	Penicillin G
Penicillin-intermediate	Ceftriaxone or cefotaxime or cefepime
Penicillin-resistant	Ceftriaxone (or cefotaxime or cefepime) + vancomycin
Gram-negative bacilli (except <i>Pseudomonas</i> spp.)	Ceftriaxone or cefotaxime
<i>Pseudomonas aeruginosa</i>	Ceftazidime or cefepime or meropenem
<i>Staphylococci</i> spp.	
Methicillin-sensitive	Nafcillin
Methicillin-resistant	Vancomycin
<i>Listeria monocytogenes</i>	Ampicillin + gentamicin
<i>Haemophilus influenzae</i>	Ceftriaxone or cefotaxime or cefepime
<i>Streptococcus agalactiae</i>	Penicillin G or ampicillin
<i>Bacteroides fragilis</i>	Metronidazole
<i>Fusobacterium</i> spp.	Metronidazole

^aDoses are as indicated in Table 164-1.

treated with one dose of azithromycin (500 mg) or one intramuscular dose of ceftriaxone (250 mg). Close contacts are defined as those individuals who have had contact with oropharyngeal secretions, either through kissing or by sharing toys, beverages, or cigarettes.

Pneumococcal Meningitis Antimicrobial therapy of pneumococcal meningitis is initiated with a cephalosporin (ceftriaxone, cefotaxime, or cefepime) and vancomycin. All CSF isolates of *S. pneumoniae* should be tested for sensitivity to penicillin and the cephalosporins. Once the results of antimicrobial susceptibility tests are known, therapy can be modified accordingly (Table 164-3). For *S. pneumoniae* meningitis, an isolate of *S. pneumoniae* is considered to be susceptible to penicillin with a minimal inhibitory concentration (MIC) <0.06 µg/mL and to be resistant when the MIC is >0.12 µg/mL. Isolates of *S. pneumoniae* that have cephalosporin MICs ≤0.5 µg/mL are considered sensitive to the cephalosporins (cefotaxime, ceftriaxone, cefepime). Those with MICs of 1 µg/mL are considered to have intermediate resistance, and those with MICs ≥2 µg/mL are considered resistant. For meningitis due to pneumococci, with cefotaxime or ceftriaxone MICs ≤0.5 µg/mL, treatment with cefotaxime or ceftriaxone is usually adequate. For MIC >1 µg/mL, vancomycin is the antibiotic of choice. Rifampin can be added to vancomycin for its synergistic effect but is inadequate as monotherapy because resistance develops rapidly when it is used alone.

A 2-week course of intravenous antimicrobial therapy is recommended for pneumococcal meningitis.

Patients with *S. pneumoniae* meningitis should have a repeat LP performed 24–36 h after the initiation of antimicrobial therapy to document sterilization of the CSF. Failure to sterilize the CSF after 24–36 h of antibiotic therapy should be considered presumptive evidence of antibiotic resistance. Patients with penicillin- and cephalosporin-resistant strains of *S. pneumoniae* who do not respond to intravenous vancomycin alone may benefit from the addition of intraventricular vancomycin. The intraventricular route of administration is preferred over the intrathecal route because adequate concentrations of vancomycin in the cerebral ventricles are not always achieved with intrathecal administration.

Listeria Meningitis Meningitis due to *L. monocytogenes* is treated with ampicillin for at least 3 weeks (Table 164-3). Gentamicin is added in critically ill patients (2 mg/kg loading dose, then 7.5 mg/kg per day

890 given every 8 h and adjusted for serum levels and renal function). The combination of trimethoprim (10–20 mg/kg per day) and sulfamethoxazole (50–100 mg/kg per day) given every 6 h may provide an alternative in penicillin-allergic patients.

Staphylococcal Meningitis Meningitis due to susceptible strains of *S. aureus* or coagulase-negative staphylococci is treated with nafcillin (Table 164-3). Vancomycin is the drug of choice for methicillin-resistant staphylococci and for patients allergic to penicillin. In these patients, the CSF should be monitored during therapy. If the CSF is not sterilized after 48 h of intravenous vancomycin therapy, then either intraventricular or intrathecal vancomycin, 20 mg once daily, can be added.

Gram-Negative Bacillary Meningitis The third-generation cephalosporins—cefotaxime, ceftriaxone, and ceftazidime—are equally efficacious for the treatment of gram-negative bacillary meningitis, with the exception of meningitis due to *P. aeruginosa*, which should be treated with ceftazidime, cefepime, or meropenem (Table 164-3). A 3-week course of intravenous antibiotic therapy is recommended for meningitis due to gram-negative bacilli.

ADJUNCTIVE THERAPY

The release of bacterial cell-wall components by bactericidal antibiotics leads to the production of the inflammatory cytokines IL-1 β and TNF- α in the subarachnoid space. Dexamethasone exerts its beneficial effect by inhibiting the synthesis of IL-1 β and TNF- α at the level of mRNA, decreasing CSF outflow resistance, and stabilizing the blood-brain barrier. The rationale for giving dexamethasone 20 min before antibiotic therapy is that dexamethasone inhibits the production of TNF- α by macrophages and microglia only if it is administered before these cells are activated by endotoxin. Dexamethasone does not alter TNF- α production once it has been induced. The results of clinical trials of dexamethasone therapy in meningitis due to *H. influenzae*, *S. pneumoniae*, and *N. meningitidis* have demonstrated its efficacy in decreasing meningeal inflammation and neurologic sequelae such as the incidence of sensorineural hearing loss.

A prospective European trial of adjunctive therapy for acute bacterial meningitis in 301 adults found that dexamethasone reduced the number of unfavorable outcomes (15 vs. 25%, $p = .03$) including death (7 vs. 15%, $p = .04$). The benefits were most striking in patients with pneumococcal meningitis. Dexamethasone (10 mg intravenously) was administered 15–20 min before the first dose of an antimicrobial agent, and the same dose was repeated every 6 h for 4 days. These results were confirmed in a second trial of dexamethasone in adults with pneumococcal meningitis. Therapy with dexamethasone should ideally be started 20 min before, or not later than concurrent with, the first dose of antibiotics. It is unlikely to be of significant benefit if started >6 h after antimicrobial therapy has been initiated. Dexamethasone may decrease the penetration of vancomycin into CSF, and it delays the sterilization of CSF in experimental models of *S. pneumoniae* meningitis. As a result, to assure reliable penetration of vancomycin into the CSF, children and adults are treated with vancomycin in a dose of 45–60 mg/kg per day. Alternatively, vancomycin can be administered by the intraventricular route.

One of the concerns for using dexamethasone in adults with bacterial meningitis is that in experimental models of meningitis, dexamethasone therapy increased hippocampal cell injury and reduced learning capacity. This has not been the case in clinical series. The efficacy of dexamethasone therapy in preventing neurologic sequelae is different between high- and low-income countries. Three large randomized trials in low-income countries (sub-Saharan Africa, Southeast Asia) failed to show benefit in subgroups of patients. The lack of efficacy of dexamethasone in these trials has been attributed to late presentation to the hospital with more advanced disease, antibiotic pretreatment, malnutrition, infection with HIV, and treatment of patients with probable, but not microbiologically proven, bacterial meningitis. The results of these clinical

trials suggest that patients in sub-Saharan Africa and those in low-income countries with negative CSF Gram's stain and culture should not be treated with dexamethasone.

INCREASED INTRACRANIAL PRESSURE

Emergency treatment of increased ICP includes elevation of the patient's head to 30–45°, intubation and hyperventilation (Paco₂ 25–30 mmHg), and mannitol. Patients with increased ICP should be managed in an intensive care unit; accurate ICP measurements are best obtained with an ICP monitoring device.

Treatment of increased intracranial pressure is discussed in detail in Chap. 330.

PROGNOSIS

Mortality rate is 3–7% for meningitis caused by *H. influenzae*, *N. meningitidis*, or group B streptococci; 15% for that due to *L. monocytogenes*; and 20% for *S. pneumoniae*. In general, the risk of death from bacterial meningitis increases with (1) decreased level of consciousness on admission, (2) onset of seizures within 24 h of admission, (3) signs of increased ICP, (4) young age (infancy) and age >50, (5) the presence of comorbid conditions including shock and/or the need for mechanical ventilation, and (6) delay in the initiation of treatment. Decreased CSF glucose concentration (<2.2 mmol/L [<40 mg/dL]) and markedly increased CSF protein concentration (>3 g/L [>300 mg/dL]) have been predictive of increased mortality and poorer outcomes in some series. Moderate or severe sequelae occur in ~25% of survivors, although the exact incidence varies with the infecting organism. Common sequelae include decreased intellectual function, memory impairment, seizures, hearing loss and dizziness, and gait disturbances.

ACUTE VIRAL MENINGITIS

CLINICAL MANIFESTATIONS

Immunocompetent adult patients with viral meningitis usually present with headache, fever, and signs of meningeal irritation coupled with an inflammatory CSF profile (see below). Headache is almost invariably present and often characterized as frontal or retroorbital and frequently associated with photophobia and pain on moving the eyes. Nuchal rigidity is present in most cases but may be mild and present only near the limit of neck anteflexion. Constitutional signs can include malaise, myalgia, anorexia, nausea and vomiting, abdominal pain, and/or diarrhea. Patients often have mild lethargy or drowsiness; however, profound alterations in consciousness, such as stupor, coma, or marked confusion, do not occur in viral meningitis and suggest the presence of encephalitis or other alternative diagnoses. Similarly, seizures or focal neurologic signs or symptoms or neuroimaging abnormalities indicative of brain parenchymal involvement are not typical of viral meningitis and suggest the presence of encephalitis or another CNS infectious or inflammatory process.

ETIOLOGY

Using a variety of diagnostic techniques, including CSF PCR, culture, and serology, a specific viral cause can be found in 60–90% of cases of viral meningitis. The most important agents are enteroviruses (including echoviruses and coxsackieviruses in addition to numbered enteroviruses), varicella-zoster virus (VZV), HSV (HSV-2 > HSV-1), HIV, and arboviruses (Table 164-4). CSF cultures are positive in 30–70% of patients, with the frequency of isolation depending on the specific viral agent. Approximately two-thirds of culture-negative cases of “aseptic” meningitis have a specific viral etiology identified by CSF PCR testing (see below).

EPIDEMIOLOGY

Viral meningitis is not a nationally reportable disease; however, it has been estimated that the incidence is ~60,000–75,000 cases per year. In temperate climates, there is a substantial increase in cases during the nonwinter months, reflecting the seasonal predominance of enterovirus and arthropod-borne virus (arbovirus) infections in the summer and fall, with a peak monthly incidence of about 1 reported case per 100,000 population.

TABLE 164-4 VIRUSES CAUSING ACUTE MENINGITIS AND ENCEPHALITIS IN NORTH AMERICA

Acute Meningitis	
Common	Less Common
Enteroviruses (coxsackieviruses, echoviruses, and human enteroviruses 68–71)	Herpes simplex virus 1
Varicella-zoster virus	Human herpesvirus 6
Herpes simplex virus 2	Cytomegalovirus
Epstein-Barr virus	Lymphocytic choriomeningitis virus
Arthropod-borne viruses	Mumps
HIV	
Acute Encephalitis	
Common	Less Common
Herpesviruses	Rabies
Cytomegalovirus ^a	Eastern equine encephalitis virus
Herpes simplex virus 1^b	
Herpes simplex virus 2	
Human herpesvirus 6	
Varicella-zoster virus	Powassan virus
Epstein-Barr virus	Cytomegalovirus ^a
Arthropod-borne viruses	Colorado tick fever virus
La Crosse virus	Mumps
West Nile virus^c	
St. Louis encephalitis virus	
Enteroviruses	

^aImmunocompromised host. ^bThe most common cause of sporadic encephalitis.

^cThe most common cause of epidemic encephalitis.

LABORATORY DIAGNOSIS

CSF Examination The most important laboratory test in the diagnosis of viral meningitis is examination of the CSF. The typical profile is a pleocytosis, a normal or slightly elevated protein concentration (0.2–0.8 g/L [20–80 mg/dL]), a normal glucose concentration, and a normal or mildly elevated opening pressure (100–350 mmH₂O). Organisms are *not* seen on Gram's stain of CSF. The total CSF cell count in viral meningitis is typically 25–500/μL, although cell counts of several thousand/μL are occasionally seen, especially with infections due to lymphocytic choriomeningitis virus (LCMV) and mumps virus. Lymphocytes are typically the predominant cell. Rarely, PMNs may predominate in the first 48 h of illness, especially with infections due to echovirus 9, West Nile virus, eastern equine encephalitis (EEE) virus, or mumps. A PMN pleocytosis occurs in 45% of patients with West Nile virus (WNV) meningitis and can persist for a week or longer before shifting to a lymphocytic pleocytosis. PMN pleocytosis with low glucose may also be a feature of cytomegalovirus (CMV) infections in immunocompromised hosts. Despite these exceptions, the presence of a CSF PMN pleocytosis in a patient with suspected viral meningitis in whom a specific diagnosis has not been established should prompt consideration of alternative diagnoses, including bacterial meningitis or parameningeal infections. The CSF glucose concentration is typically normal in viral infections, although it may be decreased in 10–30% of cases due to mumps or LCMV. Rare instances of decreased CSF glucose concentration occur in cases of meningitis due to echoviruses and other enteroviruses, HSV-2, and VZV. As a rule, a lymphocytic pleocytosis with a low glucose concentration should suggest fungal or tuberculous meningitis, *Listeria* meningoencephalitis, or noninfectious disorders (e.g., sarcoid, neoplastic meningitis).

A number of tests measuring levels of various CSF proteins, enzymes, and mediators—including C-reactive protein, lactic acid, lactate dehydrogenase, neopterin, quinolinic acid, IL-1β, IL-6, soluble IL-2 receptor, β₂-microglobulin, and TNF—have been proposed as potential discriminators between viral and bacterial meningitis or as markers of specific types of viral infection (e.g., infection with HIV),

but they remain of uncertain sensitivity and specificity and are not widely used for diagnostic purposes.

Polymerase Chain Reaction Amplification of Viral Nucleic Acid Amplification of viral-specific DNA or RNA from CSF using PCR amplification has become the single most important method for diagnosing CNS viral infections. In both enteroviral and HSV infections of the CNS, CSF PCR has become the diagnostic procedure of choice and is substantially more sensitive than viral cultures. HSV CSF PCR is also an important diagnostic test in patients with recurrent episodes of “aseptic” meningitis, many of whom have amplifiable HSV DNA in CSF despite negative viral cultures. CSF PCR is also used routinely to diagnose CNS viral infections caused by CMV, Epstein-Barr virus (EBV), VZV, and human herpesvirus 6 (HHV-6). CSF PCR tests are available for WNV but are not as sensitive as detection of WNV-specific CSF IgM. PCR is also useful in the diagnosis of CNS infection caused by *Mycoplasma pneumoniae*, which can mimic viral meningitis and encephalitis. PCR of throat washings may assist in diagnosis of enteroviral and mycoplasmal CNS infections. PCR of stool specimens may also assist in diagnosis of enteroviral infections (see below).

Viral Culture The sensitivity of CSF cultures for the diagnosis of viral meningitis and encephalitis, in contrast to its utility in bacterial infections, is generally poor. In addition to CSF, specific viruses may also be isolated from throat swabs, stool, blood, and urine. Enteroviruses and adenoviruses may be found in feces; arboviruses, some enteroviruses, and LCMV in blood; mumps and CMV in urine; and enteroviruses, mumps, and adenoviruses in throat washings. During enteroviral infections, viral shedding in stool may persist for several weeks. The presence of enterovirus in stool is not diagnostic and may result from residual shedding from a previous enteroviral infection; it also occurs in some asymptomatic individuals during enteroviral epidemics.

Serologic Studies For many arboviruses including WNV, serologic studies remain important diagnostic tools. Serum antibody determination is less useful for viruses with high seroprevalence rates in the general population such as HSV, VZV, CMV, and EBV. For viruses with low seroprevalence rates, diagnosis of acute viral infection can be made by documenting seroconversion between acute-phase and convalescent sera (typically obtained after 2–4 weeks) or by demonstrating the presence of virus-specific IgM antibodies. For viruses with high seroprevalence such as VZV and HSV, demonstration of synthesis of virus-specific antibodies in CSF, as shown by an increased IgG index or the presence of CSF IgM antibodies, may be useful and can provide presumptive evidence of CNS infection. Although serum and CSF IgM antibodies generally persist for only a few months after acute infection, there are exceptions to this rule. For example, WNV serum IgM has been shown to persist in some patients for >1 year following acute infection. Unfortunately, the delay between onset of infection and the host's generation of a virus-specific antibody response often means that serologic data are useful mainly for the retrospective establishment of a specific diagnosis, rather than in aiding acute diagnosis or management. In the case of EBV, demonstration of antibody responses consistent with recent/acute infection (e.g., IgM viral capsid antibody, antibody against early antigen, absence of antibody against EBV-associated nuclear antigen) may assist in diagnosis.

CSF oligoclonal gamma globulin bands occur in association with a number of viral infections. The associated antibodies are often directed against viral proteins. Oligoclonal bands also occur commonly in certain noninfectious neurologic diseases (e.g., multiple sclerosis) and may be found in nonviral infections (e.g., neurosyphilis, Lyme neuroborreliosis).

Other Laboratory Studies All patients with suspected viral meningitis should have a complete blood count and differential, liver and renal function tests, erythrocyte sedimentation rate (ESR), and C-reactive protein, electrolytes, glucose, creatine kinase, aldolase, amylase, and lipase. Neuroimaging studies (MRI preferable to CT) are not absolutely necessary in patients with uncomplicated viral meningitis but should be performed in patients with altered consciousness, seizures, focal neurologic signs or symptoms, atypical CSF profiles, or underlying immunocompromising treatments or conditions.

The most important issue in the differential diagnosis of viral meningitis is to consider diseases that can mimic viral meningitis, including (1) untreated or partially treated bacterial meningitis; (2) early stages of meningitis caused by fungi, mycobacteria, or *Treponema pallidum* (neurosyphilis), in which a lymphocytic pleocytosis is common, cultures may be slow growing or negative, and hypoglycorrhachia may not be present early; (3) meningitis caused by agents such as *Mycoplasma*, *Listeria* spp., *Brucella* spp., *Coxiella* spp., *Leptospira* spp., and *Rickettsia* spp.; (4) parameningeal infections; (5) neoplastic meningitis; and (6) meningitis secondary to noninfectious inflammatory diseases, including autoimmune and hypersensitivity meningitis, SLE and other rheumatologic diseases, sarcoidosis, Behçet's syndrome, and the uveomeningitic syndromes. Studies in children >28 days of age suggest that the presence of CSF protein >0.5 g/L (sensitivity 89%, specificity 78%) and elevated serum procalcitonin levels >0.5 ng/mL (sensitivity 89%, specificity 89%) were clues to the presence of bacterial as opposed to "aseptic" meningitis. A variety of clinical algorithms for differentiating bacterial from aseptic meningitis have been developed. One such prospectively validated system, the *bacterial meningitis score*, suggests that the probability of bacterial meningitis is 0.3% or less (negative predictive value 99.7%, 95% confidence interval 99.6–100%) in children with CSF pleocytosis who have (1) a negative CSF Gram's stain, (2) CSF neutrophil count <1000 cells/ μ L, (3) CSF protein <80 mg/dL, (4) peripheral absolute neutrophil count of <10,000 cells/ μ L, and (5) no prior history or current presence of seizures.

SPECIFIC VIRAL ETIOLOGIES

Enteroviruses (EV) (Chap. 228) are the most common cause of viral meningitis, accounting for >85% of cases in which a specific etiology can be identified. Cases may either be sporadic or occur in clusters. EV71 has produced large epidemics of neurologic disease outside the United States, especially in Southeast Asia, but most recently reported cases in the United States have been sporadic. Enteroviruses are the most likely cause of viral meningitis in the summer and fall months, especially in children (<15 years), although cases occur at reduced frequency year round. Although the incidence of enteroviral meningitis declines with increasing age, some outbreaks have preferentially affected older children and adults. Meningitis outside the neonatal period is usually benign. Patients present with sudden onset of fever; headache; nuchal rigidity; and often constitutional signs, including vomiting, anorexia, diarrhea, cough, pharyngitis, and myalgias. The physical examination should include a careful search for stigmata of enterovirus infection, including exanthems, hand-foot-mouth disease, herpangina, pleurodynia, myopericarditis, and hemorrhagic conjunctivitis. The CSF profile is typically a lymphocytic pleocytosis (100–1000 cells/ μ L) with normal glucose and normal or mildly elevated protein concentration. However, up to 15% of patients, most commonly young infants rather than older children or adults, have a normal CSF leukocyte count. In rare cases, PMNs may predominate during the first 48 h of illness. CSF reverse transcriptase PCR (RT-PCR) is the diagnostic procedure of choice and is both sensitive (>95%) and specific (>100%). CSF PCR has the highest sensitivity if performed within 48 h of symptom onset, with sensitivity declining rapidly after day 5 of symptoms. PCR of throat washings or stool specimens may be positive for several weeks, and positive results can help support the diagnosis of an acute enteroviral infection. The sensitivity of routine enteroviral PCRs for detecting EV71 is low, and specific testing may be required. Treatment is supportive, and patients usually recover without sequelae. Chronic and severe infections can occur in neonates and in individuals with hypo- or agammaglobulinemia.

Arbovirus infections (Chap. 233) occur predominantly in the summer and early fall. Arboviral meningitis should be considered when clusters of meningitis and encephalitis cases occur in a restricted geographic region during the summer or early fall. In the United States, the most important causes of arboviral meningitis and encephalitis are WNV, St. Louis encephalitis virus, and the California encephalitis group of viruses. In WNV epidemics, avian deaths may serve as sentinel infections for subsequent human disease. A history of tick

exposure or travel or residence in the appropriate geographic area should suggest the possibility of Colorado tick fever virus or Powassan virus infection, although nonviral tick-borne diseases, including RMSF and Lyme neuroborreliosis, may present similarly. Arbovirus meningoencephalitis is typically associated with a CSF lymphocytic pleocytosis, normal glucose concentration, and normal or mildly elevated protein concentration. However, ~45% of patients with WNV meningoencephalitis have CSF neutrophilia, which can persist for a week or more. The rarity of hypoglycorrhachia in WNV infection, the absence of positive Gram's stains, and the negative cultures help distinguish these patients from those with bacterial meningitis. Definitive diagnosis of arboviral meningoencephalitis is based on demonstration of viral-specific IgM in CSF or seroconversion. The prevalence of CSF IgM increases progressively during the first week after infection, peaking at >80% in patients with neuroinvasive disease; as a result, repeat studies may be needed when disease suspicion is high and an early study is negative. CSF PCR tests are available for some viruses in selected diagnostic laboratories and at the Centers for Disease Control and Prevention (CDC), but in the case of WNV, sensitivity (~70%) of CSF PCR is less than that of CSF serology. WNV CSF PCR may be useful in immunocompromised patients who may have absent or reduced antibody responses.

HSV meningitis (Chap. 216) has been increasingly recognized as a major cause of viral meningitis in adults, and overall, it is probably second in importance to enteroviruses as a cause of viral meningitis, accounting for 5% of total cases overall and undoubtedly a higher frequency of those cases occurring in adults and/or outside of the summer-fall period when enterovirus infections are increasingly common. In adults, the majority of cases of uncomplicated meningitis are due to HSV-2, whereas HSV-1 is responsible for 90% of cases of HSV encephalitis. HSV meningitis occurs in ~25–35% of women and ~10–15% of men at the time of an initial (primary) episode of genital herpes. Of these patients, 20% go on to have recurrent attacks of meningitis. Diagnosis of HSV meningitis is usually by HSV CSF PCR because cultures may be negative, especially in patients with recurrent meningitis. Demonstration of intrathecal synthesis of HSV-specific antibody may also be useful in diagnosis, although antibody tests are less sensitive and less specific than PCR and may not become positive until after the first week of infection. Although a history of or the presence of HSV genital lesions is an important diagnostic clue, many patients with HSV meningitis give no history and have no evidence of active genital herpes at the time of presentation. Most cases of recurrent viral or "aseptic" meningitis, including cases previously diagnosed as Mollaret's meningitis, are due to HSV.

VZV meningitis should be suspected in the presence of concurrent chickenpox or shingles. However, it is important to recognize that VZV is being increasingly identified as an important cause of both meningitis and encephalitis in patients without rash. The frequency of VZV as a cause of meningitis is extremely variable, ranging from as low as 3% to as high as 20% in different series. Diagnosis is usually based on CSF PCR, although the sensitivity of this test may not be as high as for the other herpesviruses. VZV serologic studies complement PCR testing, and the diagnosis of VZV CNS infection can be made by the demonstration of VZV-specific intrathecal antibody synthesis and/or the presence of VZV CSF IgM antibodies, or by positive CSF cultures.

EBV infections may also produce aseptic meningitis, with or without associated infectious mononucleosis. The presence of atypical lymphocytes in the CSF or peripheral blood is suggestive of EBV infection but may occasionally be seen with other viral infections. EBV is almost never cultured from CSF. Serum and CSF serology can help establish the presence of acute infection, which is characterized by IgM viral capsid antibodies (VCAs), antibodies to early antigens (EAs), and the absence of antibodies to EBV-associated nuclear antigen (EBNA). CSF PCR is another important diagnostic test, although false-positive results may reflect viral reactivation associated with other infectious or inflammatory processes or the presence of latent viral DNA in lymphocytes recruited due to other inflammatory conditions.

HIV meningitis should be suspected in any patient presenting with a viral meningitis with known or suspected risk factors for HIV infection.

Meningitis may occur following primary infection with HIV in 5–10% of cases and less commonly at later stages of illness. Cranial nerve palsies, most commonly involving cranial nerves V, VII, or VIII, are more common in HIV meningitis than in other viral infections. Diagnosis can be confirmed by detection of HIV genome in blood or CSF. Seroconversion may be delayed, and patients with negative HIV serologies who are suspected of having HIV meningitis should be monitored for delayed seroconversion. **For further discussion of HIV infection, see Chap. 226.**

Mumps (Chap. 231e) should be considered when meningitis occurs in the late winter or early spring, especially in males (male-to-female ratio 3:1). With the widespread use of the live attenuated mumps vaccine in the United States since 1967, the incidence of mumps meningitis has fallen by >95%; however, mumps remains a potential source of infection in nonimmunized individuals and populations. Rare cases (10–100:100,000 vaccinated individuals) of vaccine-associated mumps meningitis have been described, with onset typically 2–4 weeks after vaccination. The presence of parotitis, orchitis, oophoritis, pancreatitis, or elevations in serum lipase and amylase is suggestive of mumps meningitis; however, their absence does not exclude the diagnosis. Clinical meningitis was previously estimated to occur in 10–30% of patients with mumps parotitis; however, in a recent U.S. outbreak of nearly 2600 cases of mumps, only 11 cases of meningitis were identified, suggesting the incidence may be lower than previously suspected. Mumps infection confers lifelong immunity, so a documented history of previous infection excludes this diagnosis. Patients with meningitis have a CSF pleocytosis that can exceed 1000 cells/ μ L in 25%. Lymphocytes predominate in 75%, although CSF neutrophilia occurs in 25%. Hypoglycorrhachia, occurs in 10–30% of patients and may be a clue to the diagnosis when present. Diagnosis is typically made by culture of virus from CSF or by detecting IgM antibodies or seroconversion. CSF PCR is available in some diagnostic and research laboratories.

LCMV infection (Chap. 233) should be considered when aseptic meningitis occurs in the late fall or winter and in individuals with a history of exposure to house mice (*Mus musculus*), pet or laboratory rodents (e.g., hamsters, rats, mice), or their excreta. Some patients have an associated rash, pulmonary infiltrates, alopecia, parotitis, orchitis, or myopericarditis. Laboratory clues to the diagnosis of LCMV, in addition to the clinical findings noted above, may include the presence of leukopenia, thrombocytopenia, or abnormal liver function tests. Some cases present with a marked CSF pleocytosis (>1000 cells/ μ L) and hypoglycorrhachia (<30%). Diagnosis is based on serology and/or culture of virus from CSF.

TREATMENT ACUTE VIRAL MENINGITIS

Treatment of almost all cases of viral meningitis is primarily symptomatic and includes use of analgesics, antipyretics, and antiemetics. Fluid and electrolyte status should be monitored. Patients with suspected bacterial meningitis should receive appropriate empirical therapy pending culture results (see above). Hospitalization may not be required in immunocompetent patients with presumed viral meningitis and no focal signs or symptoms, no significant alteration in consciousness, and a classic CSF profile (lymphocytic pleocytosis, normal glucose, negative Gram's stain) if adequate provision for monitoring at home and medical follow-up can be ensured. Immunocompromised patients; patients with significant alteration in consciousness, seizures, or the presence of focal signs and symptoms suggesting the possibility of encephalitis or parenchymal brain involvement; and patients who have an atypical CSF profile should be hospitalized. Oral or intravenous acyclovir may be of benefit in patients with meningitis caused by HSV-1 or -2 and in cases of severe EBV or VZV infection. Data concerning treatment of HSV, EBV, and VZV meningitis are extremely limited. Seriously ill patients should probably receive intravenous acyclovir (15–30 mg/kg per day in three divided doses), which can be followed by an oral drug such as acyclovir (800 mg five times daily), famciclovir (500 mg tid), or valacyclovir (1000 mg tid) for a total course of 7–14 days. Patients

who are less ill can be treated with oral drugs alone. Patients with HIV meningitis should receive highly active antiretroviral therapy (**Chap. 226**). There is no specific therapy of proven benefit for patients with arboviral encephalitis, including that caused by WNV.

Patients with viral meningitis who are known to have deficient humoral immunity (e.g., X-linked agammaglobulinemia) and who are not already receiving either intramuscular gamma globulin or intravenous immunoglobulin (IVIg) should be treated with these agents. Intraventricular administration of immunoglobulin through an Ommaya reservoir has been tried in some patients with chronic enteroviral meningitis who have not responded to intramuscular or intravenous immunoglobulin.

Vaccination is an effective method of preventing the development of meningitis and other neurologic complications associated with poliovirus, mumps, measles, rubella, and varicella infection. A live attenuated VZV vaccine (Varivax) is available in the United States. Clinical studies indicate an effectiveness rate of 70–90% for this vaccine, but a booster may be required after ~10 years to maintain immunity. A related vaccine (Zostavax) is recommended for prevention of herpes zoster (shingles) in adults over the age of 60. An inactivated varicella vaccine is available for transplant recipients and others for whom live viral vaccines are contraindicated

PROGNOSIS

In adults, the prognosis for full recovery from viral meningitis is excellent. Rare patients complain of persisting headache, mild mental impairment, incoordination, or generalized asthenia for weeks to months. The outcome in infants and neonates (<1 year) is less certain; intellectual impairment, learning disabilities, hearing loss, and other lasting sequelae have been reported in some studies.

VIRAL ENCEPHALITIS

DEFINITION

In contrast to viral meningitis, where the infectious process and associated inflammatory response are limited largely to the meninges, in encephalitis the brain parenchyma is also involved. Many patients with encephalitis also have evidence of associated meningitis (meningoencephalitis) and, in some cases, involvement of the spinal cord or nerve roots (encephalomyelitis, encephalomyelorradiculitis).

CLINICAL MANIFESTATIONS

In addition to the acute febrile illness with evidence of meningeal involvement characteristic of meningitis, the patient with encephalitis commonly has an altered level of consciousness (confusion, behavioral abnormalities), or a depressed level of consciousness ranging from mild lethargy to coma, and evidence of either focal or diffuse neurologic signs and symptoms. Patients with encephalitis may have hallucinations, agitation, personality change, behavioral disorders, and, at times, a frankly psychotic state. Focal or generalized seizures occur in many patients with encephalitis. Virtually every possible type of focal neurologic disturbance has been reported in viral encephalitis; the signs and symptoms reflect the sites of infection and inflammation. The most commonly encountered focal findings are aphasia, ataxia, upper or lower motor neuron patterns of weakness, involuntary movements (e.g., myoclonic jerks, tremor), and cranial nerve deficits (e.g., ocular palsies, facial weakness). Involvement of the hypothalamic-pituitary axis may result in temperature dysregulation, diabetes insipidus, or the development of the syndrome of inappropriate secretion of antidiuretic hormone (SIADH). Even though neurotropic viruses typically cause pathologic injury in distinct regions of the CNS, variations in clinical presentations make it impossible to reliably establish the etiology of a specific case of encephalitis on clinical grounds alone (see “Differential Diagnosis,” below).

ETIOLOGY

In the United States, there are an estimated ~20,000 cases of encephalitis per year, although the actual number of cases is likely to be significantly larger. Despite comprehensive diagnostic efforts, the

894 majority of cases of acute encephalitis of suspected viral etiology remain of unknown cause. Hundreds of viruses are capable of causing encephalitis, although only a limited subset is responsible for most cases in which a specific cause is identified (Table 164-4). The most commonly identified viruses causing sporadic cases of acute encephalitis in immunocompetent adults are herpesviruses (HSV, VZV, EBV). Epidemics of encephalitis are caused by arboviruses, which belong to several different viral taxonomic groups including *Alphaviruses* (e.g., EEE virus, western equine encephalitis virus), *Flaviviruses* (e.g., WNV, St. Louis encephalitis virus, Japanese encephalitis virus, Powassan virus), and *Bunyaviruses* (e.g., California encephalitis virus serogroup, La Crosse virus). Historically, the largest number of cases of arbovirus encephalitis in the United States has been due to St. Louis encephalitis virus and the California encephalitis virus serogroup. However, since 2002, WNV has been responsible for the majority of arbovirus meningitis and encephalitis cases in the United States. WNV caused 2873 confirmed cases of neuroinvasive disease (encephalitis, meningitis, or myelitis) in 2012 with 286 deaths. States reporting >200 cases included Texas (1868 cases), California (479), Louisiana (335), Illinois (290), Mississippi (249), South Dakota (203), and Michigan (202). In 2013, there were 1140 neuroinvasive cases with 100 deaths. States reporting >100 cases included California (357 cases), Colorado (315), Nebraska (213), Texas (157), South Dakota (148), North Dakota (123), and Illinois (106). It is important to recognize that WNV epidemics are unpredictable and that cases have occurred in every state in the continental United States. New causes of viral CNS infections are constantly appearing, as evidenced by the outbreak of cases of encephalitis in Southeast Asia caused by Nipah virus, a newly identified member of the Paramyxoviridae family; of meningitis in Europe caused by Toscana virus, an arbovirus belonging to the Bunyavirus family; and of neurologic disorders associated with major epidemics of Chikungunya virus, a togavirus, in Africa, India, and Southeast Asia. Parechoviruses including human parechovirus 3 (HPeV3), members of the Picornavirus family, have recently been reported as causes of fever, sepsis, and meningitis in infants (age <3 months) in the United States and abroad.

LABORATORY DIAGNOSIS

CSF Examination CSF examination should be performed in all patients with suspected viral encephalitis unless contraindicated by the presence of severely increased ICP. Ideally at least 20 mL should be collected with 5–10 mL stored frozen for later studies as needed. The characteristic CSF profile is indistinguishable from that of viral meningitis and typically consists of a lymphocytic pleocytosis, a mildly elevated protein concentration, and a normal glucose concentration. A CSF pleocytosis (>5 cells/ μ L) occurs in >95% of immunocompetent patients with documented viral encephalitis. In rare cases, a pleocytosis may be absent on the initial LP but present on subsequent LPs. Patients who are severely immunocompromised by HIV infection, glucocorticoid or other immunosuppressant drugs, chemotherapy, or lymphoreticular malignancies may fail to mount a CSF inflammatory response. CSF cell counts exceed 500/ μ L in only about 10% of patients with encephalitis. Infections with certain arboviruses (e.g., EEE virus or California encephalitis virus), mumps, and LCMV may occasionally result in cell counts >1000/ μ L, but this degree of pleocytosis should suggest the possibility of nonviral infections or other inflammatory processes. Atypical lymphocytes in the CSF may be seen in EBV infection and less commonly with other viruses, including CMV, HSV, and enteroviruses. Increased numbers of plasmacytoid or Mollaret-like large mononuclear cells have been reported in WNV encephalitis. Polymorphonuclear pleocytosis occurs in ~45% of patients with WNV encephalitis and is also a common feature in CMV myelofolliculitis in immunocompromised patients. Large numbers of CSF PMNs may be present in patients with encephalitis due to EEE virus, echovirus 9, and, more rarely, other enteroviruses. However, persisting CSF neutrophilia should prompt consideration of bacterial infection, leptospirosis, amebic infection, and noninfectious processes such as acute hemorrhagic leukoencephalitis. About 20% of patients with encephalitis will have a significant number of red blood cells (>500/ μ L)

in the CSF in a nontraumatic tap. The pathologic correlate of this finding may be a hemorrhagic encephalitis of the type seen with HSV; however, CSF red blood cells occur with similar frequency and in similar numbers in patients with nonherpetic focal encephalitis. A decreased CSF glucose concentration is distinctly unusual in viral encephalitis and should suggest the possibility of bacterial, fungal, tuberculous, parasitic, leptospiral, syphilitic, sarcoid, or neoplastic meningitis. Rare patients with mumps, LCMV, or advanced HSV encephalitis and many patients with CMV myelofolliculitis have low CSF glucose concentrations.

CSF PCR

CSF PCR has become the primary diagnostic test for CNS infections caused by CMV, EBV, HHV-6, and enteroviruses (see “Viral Meningitis,” above). In the case of VZV CNS infection, CSF PCR and detection of virus-specific IgM or intrathecal antibody synthesis both provide important aids to diagnosis. The sensitivity and specificity of CSF PCRs vary with the virus being tested. The sensitivity (~96%) and specificity (~99%) of HSV CSF PCR are equivalent to or exceed those of brain biopsy. It is important to recognize that HSV CSF PCR results need to be interpreted after considering the likelihood of disease in the patient being tested, the timing of the test in relationship to onset of symptoms, and the prior use of antiviral therapy. A negative HSV CSF PCR test performed by a qualified laboratory at the appropriate time during illness in a patient with a high likelihood of HSV encephalitis based on clinical and laboratory abnormalities significantly reduces the likelihood of HSV encephalitis but does not exclude it. For example, in a patient with a pretest probability of 35% of having HSV encephalitis, a negative HSV CSF PCR reduces the posttest probability to ~2%, and for a patient with a pretest probability of 60%, a negative test reduces the posttest probability to ~6%. In both situations, a positive test makes the diagnosis almost certain (98–99%). There have been several recent reports of initially negative HSV CSF PCR tests that were obtained early (\leq 72 h) following symptom onset and that became positive when repeated 1–3 days later. The frequency of positive HSV CSF PCRs in patients with herpes encephalitis also decreases as a function of the duration of illness, with only ~20% of cases remaining positive after \geq 14 days. PCR results are generally not affected by \leq 1 week of antiviral therapy. In one study, 98% of CSF specimens remained PCR-positive during the first week of initiation of antiviral therapy, but the numbers fell to ~50% by 8–14 days and to ~21% by >15 days after initiation of antiviral therapy.

The sensitivity and specificity of CSF PCR tests for viruses other than HSV have not been definitively characterized. Enteroviral (EV) CSF PCR appears to have a sensitivity and specificity of >95%. EV PCR sensitivity for EV71 may be considerably lower (~30% in some reports). Parechoviruses are also not detected by standard EV RT-PCRs. The specificity of EBV CSF PCR has not been established. Positive EBV CSF PCRs associated with positive tests for other pathogens have been reported and may reflect reactivation of EBV latent in lymphocytes that enter the CNS as a result of an unrelated infectious or inflammatory process. In patients with CNS infection due to VZV, CSF antibody and PCR studies should be considered complementary, because patients may have evidence of intrathecal synthesis of VZV-specific antibodies and negative CSF PCRs. In the case of WNV infection, CSF PCR appears to be less sensitive (~70% sensitivity) than detection of WNV-specific CSF IgM, although PCR testing remains useful in immunocompromised patients who may not mount an effective anti-WNV antibody response.

CSF Culture CSF culture is generally of limited utility in the diagnosis of acute viral encephalitis. Culture may be insensitive (e.g., >95% of patients with HSV encephalitis have negative CSF cultures as do virtually all patients with EBV-associated CNS disease) and often takes too long to significantly affect immediate therapy.

Serologic Studies and Antigen Detection The basic approach to the serodiagnosis of viral encephalitis is identical to that discussed earlier for viral meningitis. Demonstration of WNV IgM antibodies is diagnostic of WNV encephalitis because IgM antibodies do not cross the

blood-brain barrier, and their presence in CSF is therefore indicative of intrathecal synthesis. Timing of antibody collection may be important because the rate of CSF WNV IgM seropositivity increases by ~10% per day during the first week after illness onset, reaching 80% or higher on day 7 after symptom onset. In patients with HSV encephalitis, both antibodies to HSV-1 glycoproteins and glycoprotein antigens have been detected in the CSF. Optimal detection of both HSV antibodies and antigen typically occurs after the first week of illness, limiting the utility of these tests in acute diagnosis. Nonetheless, HSV CSF antibody testing is of value in selected patients whose illness is >1 week in duration and who are CSF PCR-negative for HSV. In the case of VZV infection, CSF antibody tests may be positive when PCR fails to detect viral DNA, and both tests should be considered complementary rather than mutually exclusive.

MRI, CT, and EEG Patients with suspected encephalitis almost invariably undergo neuroimaging studies and often EEG. These tests help identify or exclude alternative diagnoses and assist in the differentiation between a focal, as opposed to a diffuse, encephalitic process. Focal findings in a patient with encephalitis should always raise the possibility of HSV encephalitis. Examples of focal findings include: (1) areas of increased signal intensity in the frontotemporal, cingulate, or insular regions of the brain on T2-weighted, FLAIR, or diffusion-weighted MRI (Fig. 164-3); (2) focal areas of low absorption, mass effect, and contrast enhancement on CT; or (3) periodic focal temporal lobe spikes on a background of slow or low-amplitude (“flattened”) activity on EEG. Approximately 10% of patients with PCR-documented HSV encephalitis will have a normal MRI, although nearly 80% will have abnormalities in the temporal lobe, and an additional 10% in extratemporal regions. The lesions are typically hyperintense on T2-weighted images. The addition of FLAIR and diffusion-weighted images to the standard MRI sequences enhances sensitivity. Children with HSV encephalitis may have atypical patterns of MRI lesions and often show involvement of brain regions outside the frontotemporal areas. CT is less sensitive than MRI and is normal in up to 20–35% of patients. EEG abnormalities occur in >75% of PCR-documented cases of HSV encephalitis; they typically involve the temporal lobes but are often nonspecific. Some

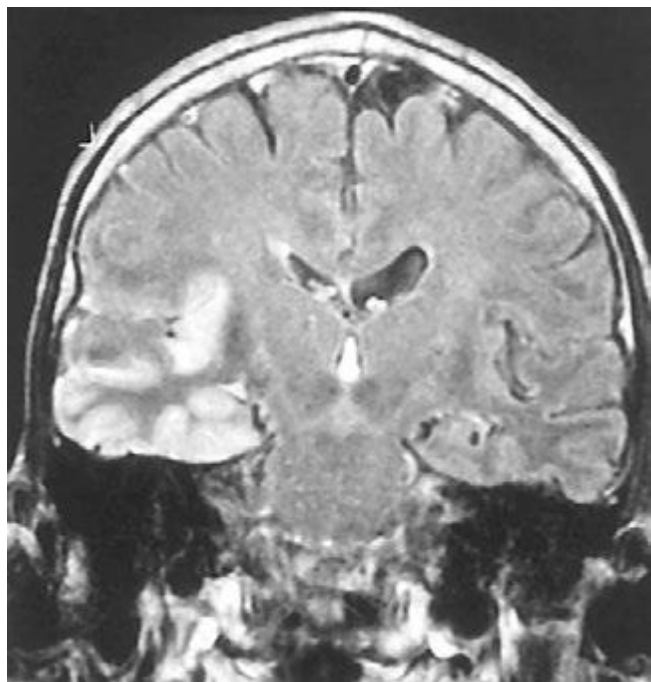


FIGURE 164-3 Coronal fluid-attenuated inversion recovery (FLAIR) magnetic resonance image from a patient with herpes simplex encephalitis. Note the area of increased signal in the right temporal lobe (left side of image) confined predominantly to the gray matter. This patient had predominantly unilateral disease; bilateral lesions are more common but may be quite asymmetric in their intensity.

patients with HSV encephalitis have a distinctive EEG pattern consisting of periodic, stereotyped, sharp-and-slow complexes originating in one or both temporal lobes and repeating at regular intervals of 2–3 s. The periodic complexes are typically noted between days 2 and 15 of the illness and are present in two-thirds of pathologically proven cases of HSV encephalitis.

Significant MRI abnormalities are found in only approximately two-thirds of patients with WNV encephalitis, a frequency less than that with HSV encephalitis. When present, abnormalities often involve deep brain structures, including the thalamus, basal ganglia, and brainstem, rather than the cortex and may only be apparent on FLAIR images. EEGs in patients with WNV encephalitis typically show generalized slowing that may be more anteriorly prominent rather than the temporally predominant pattern of sharp or periodic discharges more characteristic of HSV encephalitis. Patients with VZV encephalitis may show multifocal areas of hemorrhagic and ischemic infarction, reflecting the tendency of this virus to produce a CNS vasculopathy rather than a true encephalitis. Immunocompromised adult patients with CMV often have enlarged ventricles with areas of increased T2 signal on MRI outlining the ventricles and subependymal enhancement on T1-weighted postcontrast images. Table 164-5 highlights specific diagnostic test results in encephalitis that can be useful in clinical decision making.

Brain Biopsy Brain biopsy is now generally reserved for patients in whom CSF PCR studies fail to lead to a specific diagnosis, who have focal abnormalities on MRI, and who continue to show progressive clinical deterioration despite treatment with acyclovir and supportive therapy.

DIFFERENTIAL DIAGNOSIS

Infection by a variety of other organisms can mimic viral encephalitis. In studies of biopsy-proven HSV encephalitis, common infectious

TABLE 164-5 USE OF DIAGNOSTIC TESTS IN ENCEPHALITIS

The best test for WNV encephalitis is the CSF IgM antibody test. The prevalence of positive CSF IgM tests increases by about 10% per day after illness onset and reaches 70–80% by the end of the first week. Serum WNV IgM can provide evidence for recent WNV infection, but in the absence of other findings does not establish the diagnosis of neuroinvasive disease (meningitis, encephalitis, acute flaccid paralysis).

Approximately 80% of patients with proven HSV encephalitis have MRI abnormalities involving the temporal lobes. This percentage likely increases to >90% when FLAIR and diffusion-weighted MRI sequences are also used. The absence of temporal lobe lesions on MRI reduces the likelihood of HSV encephalitis and should prompt consideration of other diagnostic possibilities.

The CSF HSV PCR test may be negative in the first 72 h of symptoms of HSV encephalitis. A repeat study should be considered in patients with an initial early negative PCR in whom diagnostic suspicion of HSV encephalitis remains high and no alternative diagnosis has yet been established.

Detection of intrathecal synthesis (increased CSF/serum HSV antibody ratio corrected for breakdown of the blood-brain barrier) of HSV-specific antibody may be useful in diagnosis of HSV encephalitis in patients in whom only late (>1 week after onset) CSF specimens are available and PCR studies are negative. Serum serology alone is of no value in diagnosis of HSV encephalitis due to the high seroprevalence rate in the general population.

Negative CSF viral cultures are of no value in excluding the diagnosis of HSV or EBV encephalitis.

VZV CSF IgM antibodies may be present in patients with a negative VZV CSF PCR. Both tests should be performed in patients with suspected VZV CNS disease.

The specificity of EBV CSF PCR for diagnosis of CNS infection is unknown. Positive tests may occur in patients with a CSF pleocytosis due to other causes. Detection of EBV CSF IgM or intrathecal synthesis of antibody to VCA supports the diagnosis of EBV encephalitis. Serologic studies consistent with acute EBV infection (e.g., IgM VCA, presence of antibodies against EA but not against EBNA) can help support the diagnosis.

Abbreviations: CNS, central nervous system; CSF, cerebrospinal fluid; DWI, diffusion-weighted imaging; EA, early antigen; EBNA, EBV-associated nuclear antigen; EBV, Epstein-Barr virus; FLAIR, fluid-attenuated inversion recovery; HSV, herpes simplex virus; IgM, immunoglobulin M; MRI, magnetic resonance imaging; PCR, polymerase chain reaction; VCA, viral capsid antibody; VZV, varicella-zoster virus; WNV, West Nile virus.

896 mimics of focal viral encephalitis included mycobacteria, fungi, rickettsiae, *Listeria*, *Mycoplasma*, and other bacteria (including *Bartonella* sp.). Autoimmune causes of encephalitis, including those associated with antibodies against N-methyl-D-aspartate (NMDA) receptor, voltage-gated potassium channels (VGKC), α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA), and γ -aminobutyric acid (GABA) receptors, and GAD-65, have been increasingly recognized as causes of encephalitis that can mimic that caused by viral infection. In most cases, diagnosis is made by detection of the specific autoantibodies in serum and/or CSF. NMDA receptor antibodies have recently been reported in some patients with HSE encephalitis, and their presence should not exclude appropriate testing and treatment for HSV encephalitis. Autoimmune encephalitis may also be associated with specific cancers (paraneoplastic) and onconeural antibodies (e.g., anti-Hu, Yo, Ma2, amphiphysin, CRMP5, CV2) (Chap. 122). Subacute or chronic forms of encephalitis may occur in association with autoantibodies against thyroglobulin and thyroperoxidase (Hashimoto's encephalopathy) and with prion diseases.

Infection caused by the amoeba *Naegleria fowleri* can also cause acute meningoencephalitis (primary amoebic meningoencephalitis), whereas that caused by *Acanthamoeba* and *Balamuthia* more typically produces subacute or chronic granulomatous amoebic meningoencephalitis. *Naegleria* thrive in warm, iron-rich pools of water, including those found in drains, canals, and both natural and human-made outdoor pools. Infection has typically occurred in immunocompetent children with a history of swimming in potentially infected water. The CSF, in contrast to the typical profile seen in viral encephalitis, often resembles that of bacterial meningitis with a neutrophilic pleocytosis and hypoglycorrhachia. Motile trophozoites can be seen in a wet mount of warm, fresh CSF. There have been an increasing number of cases of *Balamuthia mandrillaris* amoebic encephalitis mimicking acute viral encephalitis in children and immunocompetent adults. This organism has also been associated with encephalitis in recipients of transplanted organs from a donor with unrecognized infection. No effective treatment has been identified, and mortality approaches 100%.

Encephalitis can be caused by the raccoon pinworm *Baylisascaris procyonis*. Clues to the diagnosis include a history of raccoon exposure, especially of playing in or eating dirt potentially contaminated with raccoon feces. Most patients are children, and many have an associated eosinophilia.

Once nonviral causes of encephalitis have been excluded, the major diagnostic challenge is to distinguish HSV from other viruses that cause encephalitis. This distinction is particularly important because in virtually every other instance the therapy is supportive, whereas specific and effective antiviral therapy is available for HSV, and its efficacy is enhanced when it is instituted early in the course of infection. HSV encephalitis should be considered when clinical features suggesting involvement of the inferomedial frontotemporal regions of the brain are present, including prominent olfactory or gustatory hallucinations, anosmia, unusual or bizarre behavior or personality alterations, or memory disturbance. HSV encephalitis should always be suspected in patients with signs and symptoms consistent with acute encephalitis with focal findings on clinical examination, neuroimaging studies, or EEG. The diagnostic procedure of choice in these patients is CSF PCR analysis for HSV. A positive CSF PCR establishes the diagnosis, and a negative test dramatically reduces the likelihood of HSV encephalitis (see above).



The anatomic distribution of lesions may provide an additional clue to diagnosis. Patients with rapidly progressive encephalitis and prominent brainstem signs, symptoms, or neuroimaging abnormalities may be infected by flaviviruses (WNV, St. Louis encephalitis virus, Japanese encephalitis virus), HSV, rabies, or *L. monocytogenes*. Significant involvement of deep gray matter structures, including the basal ganglia and thalamus, should also suggest possible flavivirus infection. These patients may present clinically with prominent movement disorders (tremor, myoclonus) or parkinsonian features. Patients with WNV infection can also present with a poliomyelitis-like acute flaccid paralysis, as can patients infected with EV71 and, less commonly, other enteroviruses. Acute flaccid paralysis is

characterized by the acute onset of a lower motor neuron type of weakness with flaccid tone, reduced or absent reflexes, and relatively preserved sensation. The complete eradication of polio remains an ongoing challenge despite a continuing World Health Organization poliovirus elimination campaign. Three hundred forty-one cases of polio (almost all due to serotype 1) have been reported in 2013 from eight countries (Somalia 183 cases, Pakistan 63, Nigeria 51, Kenya 14, Syria 13, Afghanistan 9, Ethiopia 6, and Cameroon 2). There have been small outbreaks of poliomyelitis associated with vaccine strains of virus that have reverted to virulence through mutation or recombination with circulating wild-type enteroviruses in Hispaniola, China, the Philippines, Indonesia, Nigeria, and Madagascar.

Epidemiologic factors may provide important clues to the diagnosis of viral meningitis or encephalitis. Particular attention should be paid to the season of the year; the geographic location and travel history; and possible exposure to animal bites or scratches, rodents, and ticks. Although transmission from the bite of an infected dog remains the most common cause of rabies worldwide, in the United States very few cases of dog rabies occur, and the most common risk factor is exposure to bats—although a clear history of a bite or scratch is often lacking. The classic clinical presentation of encephalitic (furious) rabies is fever, fluctuating consciousness, and autonomic hyperactivity. Phobic spasms of the larynx, pharynx, neck muscles, and diaphragm can be triggered by attempts to swallow water (*hydrophobia*) or by inspiration (*aerophobia*). Patients may also present with paralytic (dumb) rabies characterized by acute ascending paralysis. Rabies due to the bite of a bat has a different clinical presentation than classic rabies due to a dog or wolf bite. Patients present with focal neurologic deficits, myoclonus, seizures, and hallucinations; phobic spasms are not a typical feature. Patients with rabies have a CSF lymphocytic pleocytosis and may show areas of increased T2 signal abnormality in the brainstem, hippocampus, and hypothalamus. Diagnosis can be made by finding rabies virus antigen in brain tissue or in the neural innervation of hair follicles at the nape of the neck. PCR amplification of viral nucleic acid from CSF and saliva or tears may also enable diagnosis. Serology is frequently negative in both serum and CSF in the first week after onset of infection, which limits its acute diagnostic utility. No specific therapy is available, and cases are almost invariably fatal, with isolated survivors having devastating neurologic sequelae.

State public health authorities provide a valuable resource concerning isolation of particular agents in individual regions. Regular updates concerning the number, type, and distribution of cases of arboviral encephalitis can be found on the CDC and U.S. Geological Survey (USGS) websites (<http://www.cdc.gov> and <http://diseasemaps.usgs.gov>).

TREATMENT VIRAL ENCEPHALITIS

Specific antiviral therapy should be initiated when appropriate. Vital functions, including respiration and blood pressure, should be monitored continuously and supported as required. In the initial stages of encephalitis, many patients will require care in an intensive care unit. Basic management and supportive therapy should include careful monitoring of ICP, fluid restriction, avoidance of hypotonic intravenous solutions, and suppression of fever. Seizures should be treated with standard anticonvulsant regimens, and prophylactic therapy should be considered in view of the high frequency of seizures in severe cases of encephalitis. As with all seriously ill, immobilized patients with altered levels of consciousness, encephalitic patients are at risk for aspiration pneumonia, stasis ulcers and decubiti, contractures, deep venous thrombosis and its complications, and infections of indwelling lines and catheters.

Acyclovir is of benefit in the treatment of HSV and should be started empirically in patients with suspected viral encephalitis, especially if focal features are present, while awaiting viral diagnostic studies. Treatment should be discontinued in patients found not to have HSV encephalitis, with the possible exception of patients with severe encephalitis due to VZV or EBV. HSV, VZV, and EBV all encode an enzyme, deoxythymidine (thymidine) kinase, that phosphorylates acyclovir to produce acyclovir-5'-monophosphate.

Host cell enzymes then phosphorylate this compound to form a triphosphate derivative. It is the triphosphate that acts as an antiviral agent by inhibiting viral DNA polymerase and by causing premature termination of nascent viral DNA chains. The specificity of action depends on the fact that uninfected cells do not phosphorylate significant amounts of acyclovir to acyclovir-5'-monophosphate. A second level of specificity is provided by the fact that the acyclovir triphosphate is a more potent inhibitor of viral DNA polymerase than of the analogous host cell enzymes.

Adults should receive a dose of 10 mg/kg of acyclovir intravenously every 8 h (30 mg/kg per day total dose) for 14–21 days. CSF PCR can be repeated at the completion of this course, with PCR-positive patients receiving additional treatment, followed by a repeat CSF PCR test. Neonatal HSV CNS infection is less responsive to acyclovir therapy than HSV encephalitis in adults; it is recommended that neonates with HSV encephalitis receive 20 mg/kg of acyclovir every 8 h (60 mg/kg per day total dose) for a minimum of 21 days.

Prior to intravenous administration, acyclovir should be diluted to a concentration ≤ 7 mg/mL. (A 70-kg person would receive a dose of 700 mg, which would be diluted in a volume of 100 mL.) Each dose should be infused slowly over 1 h, rather than by rapid or bolus infusion, to minimize the risk of renal dysfunction. Care should be taken to avoid extravasation or intramuscular or subcutaneous administration. The alkaline pH of acyclovir can cause local inflammation and phlebitis (9%). Dose adjustment is required in patients with impaired renal glomerular filtration. Penetration into CSF is excellent, with average drug levels $\sim 50\%$ of serum levels. Complications of therapy include elevations in blood urea nitrogen and creatinine levels (5%), thrombocytopenia (6%), gastrointestinal toxicity (nausea, vomiting, diarrhea) (7%), and neurotoxicity (lethargy or obtundation, disorientation, confusion, agitation, hallucinations, tremors, seizures) (1%). Acyclovir resistance may be mediated by changes in either the viral deoxythymidine kinase or DNA polymerase. To date, acyclovir-resistant isolates have not been a significant clinical problem in immunocompetent individuals. However, there have been reports of clinically virulent acyclovir-resistant HSV isolates from sites outside the CNS in immunocompromised individuals, including those with AIDS.

Oral antiviral drugs with efficacy against HSV, VZV, and EBV, including acyclovir, famciclovir, and valacyclovir, have not been evaluated in the treatment of encephalitis either as primary therapy or as supplemental therapy following completion of a course of parenteral acyclovir. A recently completed National Institute of Allergy and Infectious Disease (NIAID)/National Institute of Neurological Disorders and Stroke-sponsored phase III trial of supplemental oral valacyclovir therapy (2 g tid for 3 months) following the initial 14- to 21-day course of therapy with parenteral acyclovir (www.clinicaltrials.gov, identifier NCT00031486) was terminated early due to low enrollment. Although analysis was compromised due to low numbers, no differences were seen in the 12-month endpoints including dementia rating scale, mini-mental state exam, and Glasgow coma score in patients receiving valacyclovir versus placebo. The role of adjunctive intravenous glucocorticoids in treatment of HSV and VZV infection remains unclear, with most guidelines considering the existing supportive evidence weak and recommendation for possible use based on expert opinion only.

Ganciclovir and foscarnet, either alone or in combination, are often used in the treatment of CMV-related CNS infections, although their efficacy remains unproven. Cidofovir (see below) may provide an alternative in patients who fail to respond to ganciclovir and foscarnet, although data concerning its use in CMV CNS infections are extremely limited.

Ganciclovir is a synthetic nucleoside analogue of 2'-deoxyguanosine. The drug is preferentially phosphorylated by virus-induced cellular kinases. Ganciclovir triphosphate acts as a competitive inhibitor of the CMV DNA polymerase, and its incorporation into nascent viral DNA results in premature chain termination. Following intravenous administration, CSF concentrations of ganciclovir are 25–70% of

coincident plasma levels. The usual dose for treatment of severe neurologic illnesses is 5 mg/kg every 12 h given intravenously at a constant rate over 1 h. Induction therapy is followed by maintenance therapy of 5 mg/kg every day for an indefinite period. Induction therapy should be continued until patients show a decline in CSF pleocytosis and a reduction in CSF CMV DNA copy number on quantitative PCR testing (where available). Doses should be adjusted in patients with renal insufficiency. Treatment is often limited by the development of granulocytopenia and thrombocytopenia (20–25%), which may require reduction in or discontinuation of therapy. Gastrointestinal side effects, including nausea, vomiting, diarrhea, and abdominal pain, occur in $\sim 20\%$ of patients. Some patients treated with ganciclovir for CMV retinitis have developed retinal detachment, but the causal relationship to ganciclovir treatment is unclear. Valganciclovir is an orally bioavailable prodrug that can generate high serum levels of ganciclovir, although studies of its efficacy in treating CMV CNS infections are limited.

Foscarnet is a pyrophosphate analogue that inhibits viral DNA polymerases by binding to the pyrophosphate-binding site. Following intravenous infusion, CSF concentrations range from 15 to 100% of coincident plasma levels. The usual dose for serious CMV-related neurologic illness is 60 mg/kg every 8 h administered by constant infusion over 1 h. Induction therapy for 14–21 days is followed by maintenance therapy (60–120 mg/kg per day). Induction therapy may need to be extended in patients who fail to show a decline in CSF pleocytosis and a reduction in CSF CMV DNA copy number on quantitative PCR tests (where available). Approximately one-third of patients develop renal impairment during treatment, which is reversible following discontinuation of therapy in most, but not all, cases. This is often associated with elevations in serum creatinine and proteinuria and is less frequent in patients who are adequately hydrated. Many patients experience fatigue and nausea. Reductions in serum calcium, magnesium, and potassium occur in $\sim 15\%$ of patients and may be associated with tetany, cardiac rhythm disturbances, or seizures.

Cidofovir is a nucleotide analogue that is effective in treating CMV retinitis and equivalent to or better than ganciclovir in some experimental models of murine CMV encephalitis, although data concerning its efficacy in human CMV CNS disease are limited. The usual dose is 5 mg/kg intravenously once weekly for 2 weeks, then biweekly for two or more additional doses, depending on clinical response. Patients must be prehydrated with normal saline (e.g., 1 L over 1–2 h) prior to each dose and treated with probenecid (e.g., 1 g 3 h before cidofovir and 1 g 2 and 8 h after cidofovir). Nephrotoxicity is common; the dose should be reduced if renal function deteriorates.

Intravenous ribavirin (15–25 mg/kg per day in divided doses given every 8 h) has been reported to be of benefit in isolated cases of severe encephalitis due to California encephalitis (La Crosse) virus. Ribavirin might be of benefit for the rare patients, typically infants or young children, with severe adenovirus or rotavirus encephalitis and in patients with encephalitis due to LCMV or other arenaviruses. However, clinical trials are lacking. Hemolysis, with resulting anemia, has been the major side effect limiting therapy.

No specific antiviral therapy of proven efficacy is currently available for treatment of WNV encephalitis. Patients have been treated with α -interferon, ribavirin, an Israeli IVIg preparation that contains high-titer anti-WNV antibody (Omr-IgG-am) (www.clinicaltrials.gov, identifier NCT00069316 and 0068055), and humanized monoclonal antibodies directed against the viral envelope glycoprotein (www.clinicaltrials.gov, identifier NCT00927953 and 00515385). WNV chimeric vaccines, in which WNV envelope and premembrane proteins are inserted into the background of another flavivirus, are already undergoing human clinical testing and have been found to be both safe and immunogenic in healthy adults but have not yet been tested for disease prevention in humans (www.clinicaltrials.gov, identifier NCT00746798, 00442169, 00094718, and 00537147). Both chimeric and killed inactivated WNV vaccines have been found to be safe and effective in preventing equine WNV infection, and

898 several effective flavivirus vaccines are already in human use, creating optimism that a safe and effective human WNV vaccine can also be developed.

SEQUELAE

There is considerable variation in the incidence and severity of sequelae in patients surviving viral encephalitis. In the case of EEE virus infection, nearly 80% of survivors have severe neurologic sequelae. At the other extreme are infections due to EBV, California encephalitis virus, and Venezuelan equine encephalitis virus, where severe sequelae are unusual. For example, approximately 5–15% of children infected with La Crosse virus have a residual seizure disorder, and 1% have persistent hemiparesis. Detailed information about sequelae in patients with HSV encephalitis treated with acyclovir is available from the NIAID-Collaborative Antiviral Study Group (CASG) trials. Of 32 acyclovir-treated patients, 26 survived (81%). Of the 26 survivors, 12 (46%) had no or only minor sequelae, 3 (12%) were moderately impaired (gainfully employed but not functioning at their previous level), and 11 (42%) were severely impaired (requiring continuous supportive care). The incidence and severity of sequelae were directly related to the age of the patient and the level of consciousness at the time of initiation of therapy. Patients with severe neurologic impairment (Glasgow coma score 6) at initiation of therapy either died or survived with severe sequelae. Young patients (<30 years) with good neurologic function at initiation of therapy did substantially better (100% survival, 62% with no or mild sequelae) compared with their older counterparts (>30 years; 64% survival, 57% no or mild sequelae). Some recent studies using quantitative HSV CSF PCR tests indicate that clinical outcome following treatment also correlates with the amount of HSV DNA present in CSF at the time of presentation. Many patients with WNV infection have sequelae, including cognitive impairment; weakness; and hyper- or hypokinetic movement disorders, including tremor, myoclonus, and parkinsonism. In a large longitudinal study of prognosis in 156 patients with WNV infection, the mean time to achieve recovery (defined as 95% of maximal predicted score on specific validated tests) was 112–148 days for fatigue, 121–175 days for physical function, 131–139 days for mood, and 302–455 days for mental function (the longer interval in each case representing patients with neuroinvasive disease).

SUBACUTE MENINGITIS

CLINICAL MANIFESTATIONS

Patients with subacute meningitis typically have an unrelenting headache, stiff neck, low-grade fever, and lethargy for days to several weeks before they present for evaluation. Cranial nerve abnormalities and night sweats may be present. **This syndrome overlaps that of chronic meningitis, discussed in detail in Chap. 165.**

ETIOLOGY

Common causative organisms include *M. tuberculosis*, *C. neoformans*, *H. capsulatum*, *C. immitis*, and *T. pallidum*. Initial infection with *M. tuberculosis* is acquired by inhalation of aerosolized droplet nuclei. Tuberculous meningitis in adults does not develop acutely from hematogenous spread of tubercle bacilli to the meninges. Rather, millet seed-sized (miliary) tubercles form in the parenchyma of the brain during hematogenous dissemination of tubercle bacilli in the course of primary infection. These tubercles enlarge and are usually caseating. The propensity for a caseous lesion to produce meningitis is determined by its proximity to the subarachnoid space (SAS) and the rate at which fibrous encapsulation develops. Subependymal caseous foci cause meningitis via discharge of bacilli and tuberculous antigens into the SAS. Mycobacterial antigens produce an intense inflammatory reaction that leads to the production of a thick exudate that fills the basilar cisterns and surrounds the cranial nerves and major blood vessels at the base of the brain.



Fungal infections are typically acquired by the inhalation of airborne fungal spores. The initial pulmonary infection may be asymptomatic or present with fever, cough, sputum production,

and chest pain. The pulmonary infection is often self-limited. A localized pulmonary fungal infection can then remain dormant in the lungs until there is an abnormality in cell-mediated immunity that allows the fungus to reactivate and disseminate to the CNS. The most common pathogen causing fungal meningitis is *C. neoformans*. This fungus is found worldwide in soil and bird excreta. *H. capsulatum* is endemic to the Ohio and Mississippi River valleys of the central United States and to parts of Central and South America. *C. immitis* is endemic to the desert areas of the southwest United States, northern Mexico, and Argentina.

Syphilis is a sexually transmitted disease that is manifest by the appearance of a painless chancre at the site of inoculation. *T. pallidum* invades the CNS early in the course of syphilis. Cranial nerves VII and VIII are most frequently involved.

LABORATORY DIAGNOSIS

The classic CSF abnormalities in tuberculous meningitis are as follows: (1) elevated opening pressure, (2) lymphocytic pleocytosis (10–500 cells/ μ L), (3) elevated protein concentration in the range of 1–5 g/L, and (4) decreased glucose concentration in the range of 1.1–2.2 mmol/L (20–40 mg/dL). *The combination of unrelenting headache, stiff neck, fatigue, night sweats, and fever with a CSF lymphocytic pleocytosis and a mildly decreased glucose concentration is highly suspicious for tuberculous meningitis.* The last tube of fluid collected at LP is the best tube to send for a smear for acid-fast bacilli (AFB). If there is a pellicle in the CSF or a cobweb-like clot on the surface of the fluid, AFB can best be demonstrated in a smear of the clot or pellicle. Positive smears are typically reported in only 10–40% of cases of tuberculous meningitis in adults. Cultures of CSF take 4–8 weeks to identify the organism and are positive in ~50% of adults. Culture remains the gold standard to make the diagnosis of tuberculous meningitis. PCR for the detection of *M. tuberculosis* DNA should be sent on CSF if available, but the sensitivity and specificity on CSF have not been defined. The CDC recommends the use of nucleic acid amplification tests for the diagnosis of pulmonary tuberculosis.

The characteristic CSF abnormalities in fungal meningitis are a mononuclear or lymphocytic pleocytosis, an increased protein concentration, and a decreased glucose concentration. There may be eosinophils in the CSF in *C. immitis* meningitis. Large volumes of CSF are often required to demonstrate the organism on India ink smear or grow the organism in culture. If spinal fluid examined by LP on two separate occasions fails to yield an organism, CSF should be obtained by high-cervical or cisternal puncture.

The cryptococcal polysaccharide antigen test is a highly sensitive and specific test for cryptococcal meningitis. A reactive CSF cryptococcal antigen test establishes the diagnosis. The detection of the *Histoplasma* polysaccharide antigen in CSF establishes the diagnosis of a fungal meningitis but is not specific for meningitis due to *H. capsulatum*. It may be falsely positive in coccidioidal meningitis. The CSF complement fixation antibody test is reported to have a specificity of 100% and a sensitivity of 75% for coccidioidal meningitis.

The diagnosis of syphilitic meningitis is made when a reactive serum treponemal test (fluorescent treponemal antibody absorption test [FTA-ABS] or microhemagglutination assay–*T. pallidum* [MHA-TP]) is associated with a CSF lymphocytic or mononuclear pleocytosis and an elevated protein concentration, or when the CSF Venereal Disease Research Laboratory (VDRL) test is positive. A reactive CSF FTA-ABS is not definitive evidence of neurosyphilis. The CSF FTA-ABS can be falsely positive from blood contamination. A negative CSF VDRL does not rule out neurosyphilis. A negative CSF FTA-ABS or MHA-TP rules out neurosyphilis.

TREATMENT SUBACUTE MENINGITIS

Empirical therapy of tuberculous meningitis is often initiated on the basis of a high index of suspicion without adequate laboratory support. Initial therapy is a combination of isoniazid (300 mg/d), rifampin (10 mg/kg per day), pyrazinamide (30 mg/kg per day in divided doses), ethambutol (15–25 mg/kg per day in divided doses),

and pyridoxine (50 mg/d). When the antimicrobial sensitivity of the *M. tuberculosis* isolate is known, ethambutol can be discontinued. If the clinical response is good, pyrazinamide can be discontinued after 8 weeks and isoniazid and rifampin continued alone for the next 6–12 months. A 6-month course of therapy is acceptable, but therapy should be prolonged for 9–12 months in patients who have an inadequate resolution of symptoms of meningitis or who have positive mycobacterial cultures of CSF during the course of therapy. Dexamethasone therapy is recommended for HIV-negative patients with tuberculous meningitis. The dose is 12–16 mg/d for 3 weeks, and then tapered over 3 weeks.

Meningitis due to *C. neoformans* in non-HIV, nontransplant patients is treated with induction therapy with amphotericin B (AmB) (0.7 mg/kg IV per day) plus flucytosine (100 mg/kg per day in four divided doses) for at least 4 weeks if CSF culture results are negative after 2 weeks of treatment. Therapy should be extended for a total of 6 weeks in the patient with neurologic complications. Induction therapy is followed by consolidation therapy with fluconazole 400 mg/d for 8 weeks. Organ transplant recipients are treated with liposomal AmB (3–4 mg/kg per day) or AmB lipid complex (ABLC) 5 mg/kg per day plus flucytosine (100 mg/kg per day in four divided doses) for at least 2 weeks or until CSF culture is sterile. Follow CSF yeast cultures for sterilization rather than the cryptococcal antigen titer. This treatment is followed by an 8- to 10-week course of fluconazole (400–800 mg/d [6–12 mg/kg] PO). If the CSF culture is sterile after 10 weeks of acute therapy, the dose of fluconazole is decreased to 200 mg/d for 6 months to a year. Patients with HIV infection are treated with AmB or a lipid formulation plus flucytosine for at least 2 weeks, followed by fluconazole for a minimum of 8 weeks. HIV-infected patients may require indefinite maintenance therapy with fluconazole 200 mg/d. Meningitis due to *H. capsulatum* is treated with AmB (0.7–1.0 mg/kg per day) for 4–12 weeks. A total dose of 30 mg/kg is recommended. Therapy with AmB is not discontinued until fungal cultures are sterile. After completing a course of AmB, maintenance therapy with itraconazole 200 mg two or three times daily is initiated and continued for at least 9 months to a year. *C. immitis* meningitis is treated with either high-dose fluconazole (1000 mg daily) as monotherapy or intravenous AmB (0.5–0.7 mg/kg per day) for >4 weeks. Intrathecal AmB (0.25–0.75 mg/d three times weekly) may be required to eradicate the infection. Lifelong therapy with fluconazole (200–400 mg daily) is recommended to prevent relapse. AmBisome (5 mg/kg per day) or AmB lipid complex (5 mg/kg per day) can be substituted for AmB in patients who have or who develop significant renal dysfunction. The most common complication of fungal meningitis is hydrocephalus. Patients who develop hydrocephalus should receive a CSF diversion device. A ventriculostomy can be used until CSF fungal cultures are sterile, at which time the ventriculostomy is replaced by a ventriculoperitoneal shunt.

Syphilitic meningitis is treated with aqueous penicillin G in a dose of 3–4 million units intravenously every 4 h for 10–14 days. An alternative regimen is 2.4 million units of procaine penicillin G intramuscularly daily with 500 mg of oral probenecid four times daily for 10–14 days. Either regimen is followed with 2.4 million units of benzathine penicillin G intramuscularly once a week for 3 weeks. The standard criterion for treatment success is reexamination of the CSF. The CSF should be reexamined at 6-month intervals for 2 years. The cell count is expected to normalize within 12 months, and the VDRL titer to decrease by two dilutions or revert to nonreactive within 2 years of completion of therapy. Failure of the CSF pleocytosis to resolve or an increase in the CSF VDRL titer by two or more dilutions requires retreatment.

CHRONIC ENCEPHALITIS

PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY

Clinical Features and Pathology Progressive multifocal leukoencephalopathy (PML) is characterized pathologically by multifocal areas of demyelination of varying size distributed throughout the brain but

sparing the spinal cord and optic nerves. In addition to demyelination, there are characteristic cytologic alterations in both astrocytes and oligodendrocytes. Astrocytes are enlarged and contain hyperchromatic, deformed, and bizarre nuclei and frequent mitotic figures. Oligodendrocytes have enlarged, densely staining nuclei that contain viral inclusions formed by crystalline arrays of JC virus (JCV) particles. Patients often present with visual deficits (45%), typically a homonymous hemianopia; mental impairment (38%) (dementia, confusion, personality change); weakness, including hemi- or monoparesis; and ataxia. Seizures occur in ~20% of patients, predominantly in those with lesions abutting the cortex.

Almost all patients have an underlying immunosuppressive disorder or are receiving immunomodulatory therapy. In recent series, the most common associated conditions were AIDS (80%), hematologic malignancies (13%), transplant recipients (5%), and chronic inflammatory diseases (2%). It has been estimated that up to 5% of AIDS patients will develop PML. There have been over 400 reported cases of PML occurring in patients being treated for multiple sclerosis and inflammatory bowel disease with natalizumab, a humanized monoclonal antibody that inhibits lymphocyte trafficking into CNS and bowel mucosa by binding to α_4 integrins. Overall risk in these patients has been estimated at ~3.4 PML cases per 1000 treated patients, but the risk depends on a variety of factors including anti-JCV antibody serostatus, prior immunosuppressive therapy use, and duration of natalizumab therapy. Patients who lack detectable JCV antibody have a risk of developing PML of <0.1 case/1000 patients, whereas those who are JCV seropositive and have been exposed to prior immunosuppressive therapy and have received >24 months of natalizumab therapy have a risk of >1 case/100 treated patients. PML cases have also been reported in patients receiving other humanized monoclonal antibodies with immunomodulatory activity including efalizumab and rituximab, although the relative risks have not been clearly established. The basic clinical and diagnostic features appear to be similar in HIV-associated PML and PML associated with immunomodulatory drugs with the exception of an increased likelihood of peripheral enhancement in MRIs of PML lesions in immunomodulatory cases. In natalizumab-associated PML, patients will also almost invariably develop clinical and radiographic worsening of lesions with discontinuation of therapy, attributed to development of immune reconstitution inflammatory syndrome (IRIS).

Diagnostic Studies The diagnosis of PML is frequently suggested by MRI. MRI reveals multifocal asymmetric, coalescing white matter lesions located periventricularly, in the centrum semiovale, in the parietal-occipital region, and in the cerebellum. These lesions have increased signal on T2 and FLAIR images and decreased signal on T1-weighted images. HIV-PML lesions are classically nonenhancing (90%), but patients with immunomodulatory drug associated PML may have peripheral ring enhancement. PML lesions are not typically associated with edema or mass effect. CT scans, which are less sensitive than MRI for the diagnosis of PML, often show hypodense nonenhancing white matter lesions.

The CSF is typically normal, although mild elevation in protein and/or IgG may be found. Pleocytosis occurs in <25% of cases, is predominantly mononuclear, and rarely exceeds 25 cells/ μ L. PCR amplification of JCV DNA from CSF has become an important diagnostic tool. The presence of a positive CSF PCR for JCV DNA in association with typical MRI lesions in the appropriate clinical setting is diagnostic of PML, reflecting the assay's relatively high specificity (92–100%); however, sensitivity is variable, and a negative CSF PCR does not exclude the diagnosis. In HIV-negative patients and HIV-positive patients not receiving highly active antiviral therapy (HAART), sensitivity is likely 70–90%. In HAART-treated patients, sensitivity may be closer to 60%, reflecting the lower JCV CSF viral load in this relatively more immunocompetent group. Studies with quantitative JCV CSF PCR indicate that patients with low JCV loads (<100 copies/ μ L) have a generally better prognosis than those with higher viral loads. Patients with negative CSF PCR studies may require brain biopsy for definitive diagnosis. In biopsy or necropsy specimens of brain, JCV antigen and nucleic acid can be detected by immunocytochemistry, in situ hybridization, or PCR amplification.

900 Serologic studies are of no utility in diagnosis due to high basal seroprevalence level, but may contribute to risk stratification in patients contemplating therapy with immunomodulatory drugs such as natalizumab.

TREATMENT PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY

No effective therapy for PML is available. There are case reports of potential beneficial effects of the 5-HT_{2a} receptor antagonist mirtazapine, which may inhibit binding of JCV to its receptor on oligodendrocytes. Retrospective noncontrolled studies have also suggested a possible beneficial effect of treatment with interferon- α . Neither of these agents has been tested in randomized controlled clinical trials. A prospective multicenter clinical trial to evaluate the efficacy of the antimalarial drug mefloquine failed to show benefit. Intravenous and/or intrathecal cytarabine were not shown to be of benefit in a randomized controlled trial in HIV-associated PML, although some experts suggest that cytarabine may have therapeutic efficacy in situations where breakdown of the blood-brain barrier allows sufficient CSF penetration. A randomized controlled trial of cidofovir in HIV-associated PML also failed to show significant benefit. Because PML almost invariably occurs in immunocompromised individuals, any therapeutic interventions designed to enhance or restore immunocompetence should be considered. Perhaps the most dramatic demonstration of this is disease stabilization and, in rare cases, improvement associated with the improvement in the immune status of HIV-positive patients with AIDS following institution of HAART. In HIV-positive PML patients treated with HAART, 1-year survival is ~50%, although up to 80% of survivors may have significant neurologic sequelae. HIV-positive PML patients with higher CD4 counts (>300/ μ L) and low or nondetectable HIV viral loads have a better prognosis than those with lower CD4 counts and higher viral loads. Although institution of HAART enhances survival in HIV-positive PML patients, the associated immune reconstitution in patients with an underlying opportunistic infection such as PML may also result in a severe CNS inflammatory syndrome (IRIS) associated with clinical worsening, CSF pleocytosis, and the appearance of new enhancing MRI lesions. Patients receiving natalizumab or other immunomodulatory antibodies, who are suspected of having PML, should have therapy immediately halted and circulating antibodies removed by plasma exchange. Patients should be closely monitored for development of IRIS, which is generally treated with intravenous glucocorticoids, although controlled clinical trials of efficacy remain lacking.

SUBACUTE SCLEROSING PANENCEPHALITIS (SSPE)

SSPE is a rare chronic, progressive demyelinating disease of the CNS associated with a chronic nonpermissive infection of brain tissue with measles virus. The frequency has been estimated at 1 in 100,000–500,000 measles cases. An average of five cases per year are reported in the United States. The incidence has declined dramatically since the introduction of a measles vaccine. Most patients give a history of primary measles infection at an early age (2 years), which is followed after a latent interval of 6–8 years by the development of a progressive neurologic disorder. Some 85% of patients are between 5 and 15 years old at diagnosis. Initial manifestations include poor school performance and mood and personality changes. Typical signs of a CNS viral infection, including fever and headache, do not occur. As the disease progresses, patients develop progressive intellectual deterioration, focal and/or generalized seizures, myoclonus, ataxia, and visual disturbances. In the late stage of the illness, patients are unresponsive, quadriparetic, and spastic, with hyperactive tendon reflexes and extensor plantar responses.

Diagnostic Studies MRI is often normal early, although areas of increased T2 signal develop in the white matter of the brain and brainstem as disease progresses. The EEG may initially show only nonspecific

slowing, but with disease progression, patients develop a characteristic periodic pattern with bursts of high-voltage, sharp, slow waves every 3–8 s, followed by periods of attenuated (“flat”) background. The CSF is acellular with a normal or mildly elevated protein concentration and a markedly elevated gamma globulin level (>20% of total CSF protein). CSF antimeasles antibody levels are invariably elevated, and oligoclonal antimeasles antibodies are often present. Measles virus can be cultured from brain tissue using special cocultivation techniques. Viral antigen can be identified immunocytochemically, and viral genome can be detected by in situ hybridization or PCR amplification.

TREATMENT SUBACUTE SCLEROSING PANENCEPHALITIS

No definitive therapy for SSPE is available. Treatment with isoprinosine (Inosiplex, 100 mg/kg per day), alone or in combination with intrathecal or intraventricular interferon- α , has been reported to prolong survival and produce clinical improvement in some patients but has never been subjected to a controlled clinical trial.

PROGRESSIVE RUBELLA PANENCEPHALITIS


This is an extremely rare disorder that primarily affects males with congenital rubella syndrome, although isolated cases have been reported following childhood rubella. After a latent period of 8–19 years, patients develop progressive neurologic deterioration. The manifestations are similar to those seen in SSPE. CSF shows a mild lymphocytic pleocytosis, slightly elevated protein concentration, markedly increased gamma globulin, and rubella virus-specific oligoclonal bands. No therapy is available. Universal prevention of both congenital and childhood rubella through the use of the available live attenuated rubella vaccine would be expected to eliminate the disease.

BRAIN ABSCESS

DEFINITION

A brain abscess is a focal, suppurative infection within the brain parenchyma, typically surrounded by a vascularized capsule. The term *cerebritis* is often employed to describe a nonencapsulated brain abscess.

EPIDEMIOLOGY

 A bacterial brain abscess is a relatively uncommon intracranial infection, with an incidence of ~0.3–1.3:100,000 persons per year. Predisposing conditions include otitis media and mastoiditis, paranasal sinusitis, pyogenic infections in the chest or other body sites, penetrating head trauma or neurosurgical procedures, and dental infections. In immunocompetent individuals the most important pathogens are *Streptococcus* spp. (anaerobic, aerobic, and viridans [40%]), Enterobacteriaceae (*Proteus* spp., *E. coli* sp., *Klebsiella* spp. [25%]), anaerobes (e.g., *Bacteroides* spp., *Fusobacterium* spp. [30%]), and staphylococci (10%). In immunocompromised hosts with underlying HIV infection, organ transplantation, cancer, or immunosuppressive therapy, most brain abscesses are caused by *Nocardia* spp., *Toxoplasma gondii*, *Aspergillus* spp., *Candida* spp., and *C. neoformans*. In Latin America and in immigrants from Latin America, the most common cause of brain abscess is *Taenia solium* (neurocysticercosis). In India and East Asia, mycobacterial infection (tuberculoma) remains a major cause of focal CNS mass lesions.

ETIOLOGY

A brain abscess may develop (1) by direct spread from a contiguous cranial site of infection, such as paranasal sinusitis, otitis media, mastoiditis, or dental infection; (2) following head trauma or a neurosurgical procedure; or (3) as a result of hematogenous spread from a remote site of infection. In up to 25% of cases, no obvious primary source of infection is apparent (cryptogenic brain abscess).

Approximately one-third of brain abscesses are associated with otitis media and mastoiditis, often with an associated cholesteatoma.

Otogenic abscesses occur predominantly in the temporal lobe (55–75%) and cerebellum (20–30%). In some series, up to 90% of cerebellar abscesses are otogenic. Common organisms include streptococci, *Bacteroides* spp., *Pseudomonas* spp., *Haemophilus* spp., and Enterobacteriaceae. Abscesses that develop as a result of direct spread of infection from the frontal, ethmoidal, or sphenoidal sinuses and those that occur due to dental infections are usually located in the frontal lobes. Approximately 10% of brain abscesses are associated with paranasal sinusitis, and this association is particularly strong in young males in their second and third decades of life. The most common pathogens in brain abscesses associated with paranasal sinusitis are streptococci (especially *Streptococcus milleri*), *Haemophilus* spp., *Bacteroides* spp., *Pseudomonas* spp., and *S. aureus*. Dental infections are associated with ~2% of brain abscesses, although it is often suggested that many “cryptogenic” abscesses are in fact due to dental infections. The most common pathogens in this setting are streptococci, staphylococci, *Bacteroides* spp., and *Fusobacterium* spp.

Hematogenous abscesses account for ~25% of brain abscesses. Hematogenous abscesses are often multiple, and multiple abscesses often (50%) have a hematogenous origin. These abscesses show a predilection for the territory of the middle cerebral artery (i.e., posterior frontal or parietal lobes). Hematogenous abscesses are often located at the junction of the gray and white matter and are often poorly encapsulated. The microbiology of hematogenous abscesses is dependent on the primary source of infection. For example, brain abscesses that develop as a complication of infective endocarditis are often due to viridans streptococci or *S. aureus*. Abscesses associated with pyogenic lung infections such as lung abscess or bronchiectasis are often due to streptococci, staphylococci, *Bacteroides* spp., *Fusobacterium* spp., or Enterobacteriaceae. Abscesses that follow penetrating head trauma or neurosurgical procedures are frequently due to methicillin-resistant *S. aureus* (MRSA), *S. epidermidis*, Enterobacteriaceae, *Pseudomonas* spp., and *Clostridium* spp. Enterobacteriaceae and *P. aeruginosa* are important causes of abscesses associated with urinary sepsis. Congenital cardiac malformations that produce a right-to-left shunt, such as tetralogy of Fallot, patent ductus arteriosus, and atrial and ventricular septal defects, allow bloodborne bacteria to bypass the pulmonary capillary bed and reach the brain. Similar phenomena can occur with pulmonary arteriovenous malformations. The decreased arterial oxygenation and saturation from the right-to-left shunt and polycythemia may cause focal areas of cerebral ischemia, thus providing a nidus for microorganisms that bypassed the pulmonary circulation to multiply and form an abscess. Streptococci are the most common pathogens in this setting.

PATHOGENESIS AND HISTOPATHOLOGY

Results of experimental models of brain abscess formation suggest that for bacterial invasion of brain parenchyma to occur, there must be preexisting or concomitant areas of ischemia, necrosis, or hypoxemia in brain tissue. The intact brain parenchyma is relatively resistant to infection. Once bacteria have established infection, brain abscess frequently evolves through a series of stages, influenced by the nature of the infecting organism and by the immunocompetence of the host. The early cerebritis stage (days 1–3) is characterized by a perivascular infiltration of inflammatory cells, which surround a central core of coagulative necrosis. Marked edema surrounds the lesion at this stage. In the late cerebritis stage (days 4–9), pus formation leads to enlargement of the necrotic center, which is surrounded at its border by an inflammatory infiltrate of macrophages and fibroblasts. A thin capsule of fibroblasts and reticular fibers gradually develops, and the surrounding area of cerebral edema becomes more distinct than in the previous stage. The third stage, early capsule formation (days 10–13), is characterized by the formation of a capsule that is better developed on the cortical than on the ventricular side of the lesion. This stage correlates with the appearance of a ring-enhancing capsule on neuroimaging studies. The final stage, late capsule formation (day 14 and beyond), is defined by a well-formed necrotic center surrounded by a dense collagenous capsule. The surrounding area of cerebral edema has regressed, but marked gliosis with large numbers of reactive

astrocytes has developed outside the capsule. This gliotic process may contribute to the development of seizures as a sequela of brain abscess.

CLINICAL PRESENTATION

A brain abscess typically presents as an expanding intracranial mass lesion rather than as an infectious process. Although the evolution of signs and symptoms is extremely variable, ranging from hours to weeks or even months, most patients present to the hospital 11–12 days following onset of symptoms. The classic clinical triad of headache, fever, and a focal neurologic deficit is present in <50% of cases. The most common symptom in patients with a brain abscess is headache, occurring in >75% of patients. The headache is often characterized as a constant, dull, aching sensation, either hemicranial or generalized, and it becomes progressively more severe and refractory to therapy. Fever is present in only 50% of patients at the time of diagnosis, and its absence should not exclude the diagnosis. The new onset of focal or generalized seizure activity is a presenting sign in 15–35% of patients. Focal neurologic deficits including hemiparesis, aphasia, or visual field defects are part of the initial presentation in >60% of patients.

The clinical presentation of a brain abscess depends on its location, the nature of the primary infection if present, and the level of the ICP. Hemiparesis is the most common localizing sign of a frontal lobe abscess. A temporal lobe abscess may present with a disturbance of language (dysphasia) or an upper homonymous quadrantanopia. Nystagmus and ataxia are signs of a cerebellar abscess. Signs of raised ICP—papilledema, nausea and vomiting, and drowsiness or confusion—can be the dominant presentation of some abscesses, particularly those in the cerebellum. Meningismus is not present unless the abscess has ruptured into the ventricle or the infection has spread to the subarachnoid space.

DIAGNOSIS

Diagnosis is made by neuroimaging studies. MRI (Fig. 164-4) is better than CT for demonstrating abscesses in the early (cerebritis) stages and is superior to CT for identifying abscesses in the posterior fossa. Cerebritis appears on MRI as an area of low-signal intensity on T1-weighted images with irregular postgadolinium enhancement and as an area of increased signal intensity on T2-weighted images. Cerebritis is often not visualized by CT scan, but when present, appears as an area of hypodensity. On a contrast-enhanced CT scan, a mature brain abscess appears as a focal area of hypodensity surrounded by ring enhancement with surrounding edema (hypodensity). On contrast-enhanced T1-weighted MRI, a mature brain abscess has a capsule that enhances surrounding a hypodense center and surrounded by a hypodense area of edema. On T2-weighted MRI, there is a hyperintense central area of pus surrounded by a well-defined hypointense capsule and a hyperintense surrounding area of edema. It is important to recognize that the CT and MRI appearance, particularly of the capsule, may be altered by treatment with glucocorticoids. The distinction between a brain abscess and other focal CNS lesions such as primary or metastatic tumors may be facilitated by the use of diffusion-weighted imaging sequences on which a brain abscess typically shows increased signal due to restricted diffusion of the abscess cavity with corresponding low signal on apparent diffusion coefficient images.

Microbiologic diagnosis of the etiologic agent is most accurately determined by Gram's stain and culture of abscess material obtained by CT-guided stereotactic needle aspiration. Aerobic and anaerobic bacterial cultures and mycobacterial and fungal cultures should be obtained. Up to 10% of patients will also have positive blood cultures. LP should not be performed in patients with known or suspected focal intracranial infections such as abscess or empyema; CSF analysis contributes nothing to diagnosis or therapy, and LP increases the risk of herniation.

Additional laboratory studies may provide clues to the diagnosis of brain abscess in patients with a CNS mass lesion. About 50% of patients have a peripheral leukocytosis, 60% an elevated ESR, and 80% an elevated C-reactive protein. Blood cultures are positive in ~10% of cases overall but may be positive in >85% of patients with abscesses due to *Listeria*.

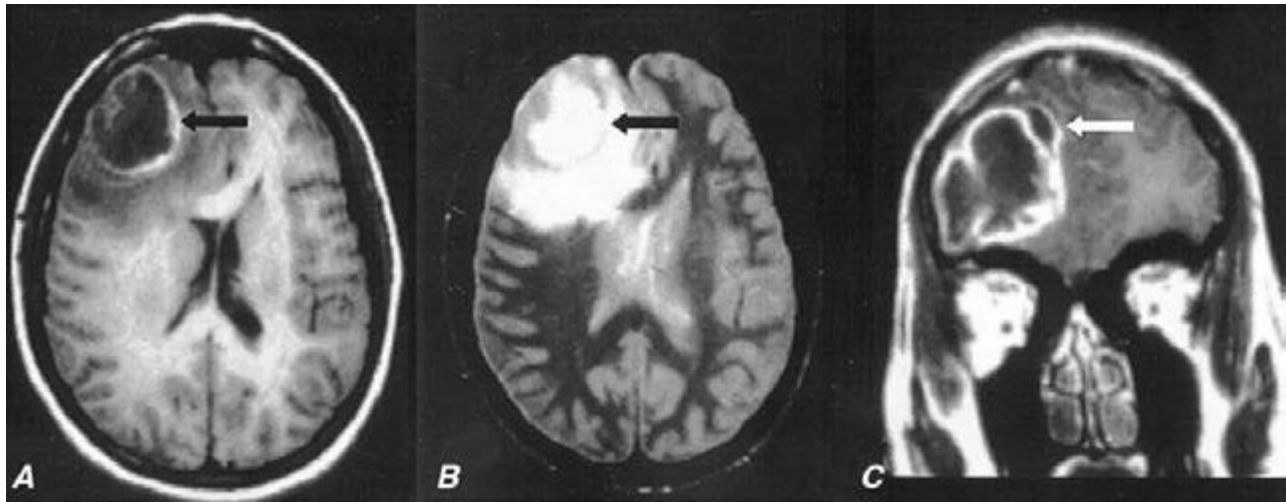


FIGURE 164-4 Pneumococcal brain abscess. Note that the abscess wall has hyperintense signal on the axial T1-weighted magnetic resonance imaging (MRI) (**A**, black arrow), hypointense signal on the axial proton density images (**B**, black arrow), and enhances prominently after gadolinium administration on the coronal T1-weighted image (**C**). The abscess is surrounded by a large amount of vasogenic edema and has a small “daughter” abscess (**C**, white arrow). (Courtesy of Joseph Lurito, MD; with permission.)

DIFFERENTIAL DIAGNOSIS

Conditions that can cause headache, fever, focal neurologic signs, and seizure activity include brain abscess, subdural empyema, bacterial meningitis, viral meningoencephalitis, superior sagittal sinus thrombosis, and acute disseminated encephalomyelitis. When fever is absent, primary and metastatic brain tumors become the major differential diagnosis. Less commonly, cerebral infarction or hematoma can have an MRI or CT appearance resembling brain abscess.

TREATMENT BRAIN ABSCESS

Optimal therapy of brain abscesses involves a combination of high-dose parenteral antibiotics and neurosurgical drainage. Empirical therapy of community-acquired brain abscess in an immunocompetent patient typically includes a third- or fourth-generation cephalosporin (e.g., cefotaxime, ceftriaxone, or cefepime) and metronidazole (see Table 164-1 for antibiotic dosages). In patients with penetrating head trauma or recent neurosurgical procedures, treatment should include ceftazidime as the third-generation cephalosporin to enhance coverage of *Pseudomonas* spp. and vancomycin for coverage of staphylococci. Meropenem plus vancomycin also provides good coverage in this setting.

Aspiration and drainage of the abscess under stereotactic guidance are beneficial for both diagnosis and therapy. Empirical antibiotic coverage should be modified based on the results of Gram's stain and culture of the abscess contents. Complete excision of a bacterial abscess via craniotomy or craniectomy is generally reserved for multiloculated abscesses or those in which stereotactic aspiration is unsuccessful.

Medical therapy alone is not optimal for treatment of brain abscess and should be reserved for patients whose abscesses are neurosurgically inaccessible, for patients with small (<2–3 cm) or nonencapsulated abscesses (cerebritis), and for patients whose condition is too tenuous to allow performance of a neurosurgical procedure. All patients should receive a minimum of 6–8 weeks of parenteral antibiotic therapy. The role, if any, of supplemental oral antibiotic therapy following completion of a standard course of parenteral therapy has never been adequately studied.

In addition to surgical drainage and antibiotic therapy, patients should receive prophylactic anticonvulsant therapy because of the high risk (~35%) of focal or generalized seizures. Anticonvulsant therapy is continued for at least 3 months after resolution of the abscess, and decisions regarding withdrawal are then based on the EEG. If the EEG is abnormal, anticonvulsant therapy should be continued. If the EEG is normal, anticonvulsant therapy can be slowly

withdrawn, with close follow-up and repeat EEG after the medication has been discontinued.

Glucocorticoids should not be given routinely to patients with brain abscesses. Intravenous dexamethasone therapy (10 mg every 6 h) is usually reserved for patients with substantial periabscess edema and associated mass effect and increased ICP. Dexamethasone should be tapered as rapidly as possible to avoid delaying the natural process of encapsulation of the abscess.

Serial MRI or CT scans should be obtained on a monthly or twice-monthly basis to document resolution of the abscess. More frequent studies (e.g., weekly) are probably warranted in the subset of patients who are receiving antibiotic therapy alone. A small amount of enhancement may remain for months after the abscess has been successfully treated.

PROGNOSIS

The mortality rate of brain abscess has declined in parallel with the development of enhanced neuroimaging techniques, improved neurosurgical procedures for stereotactic aspiration, and improved antibiotics. In modern series, the mortality rate is typically <15%. Significant sequelae, including seizures, persisting weakness, aphasia, or mental impairment, occur in ≥20% of survivors.

NONBACTERIAL CAUSES OF INFECTIOUS FOCAL CNS LESIONS

ETIOLOGY

Neurocysticercosis is the most common parasitic disease of the CNS worldwide. Humans acquire cysticercosis by the ingestion of food contaminated with the eggs of the parasite *T. solium*. Toxoplasmosis is a parasitic disease caused by *T. gondii* and acquired from the ingestion of undercooked meat and from handling cat feces.

CLINICAL PRESENTATION

The most common manifestation of neurocysticercosis is new-onset partial seizures with or without secondary generalization. Cysticerci may develop in the brain parenchyma and cause seizures or focal neurologic deficits. When present in the subarachnoid or ventricular spaces, cysticerci can produce increased ICP by interference with CSF flow. Spinal cysticerci can mimic the presentation of intraspinal tumors. When the cysticerci first lodge in the brain, they frequently cause little in the way of an inflammatory response. As the cysticercal cyst degenerates, it elicits an inflammatory response that may present clinically as a seizure. Eventually the cyst dies, a process that may take several years and is typically associated with resolution of the inflammatory response and, often, abatement of seizures.

Primary *Toxoplasma* infection is often asymptomatic. However, during this phase parasites may spread to the CNS, where they become latent. Reactivation of CNS infection is almost exclusively associated with immunocompromised hosts, particularly those with HIV infection. During this phase patients present with headache, fever, seizures, and focal neurologic deficits.

DIAGNOSIS

The lesions of neurocysticercosis are readily visualized by MRI or CT scans. Lesions with viable parasites appear as cystic lesions. The scolex can often be visualized on MRI. Lesions may appear as contrast-enhancing lesions surrounded by edema. A very early sign of cyst death is hypointensity of the vesicular fluid on T2-weighted images when compared with CSF. Parenchymal brain calcifications are the most common finding and evidence that the parasite is no longer viable. MRI findings of toxoplasmosis consist of multiple lesions in the deep white matter, the thalamus, and basal ganglia and at the gray-white junction in the cerebral hemispheres. With contrast administration, the majority of the lesions enhance in a ringed, nodular, or homogeneous pattern and are surrounded by edema. In the presence of the characteristic neuroimaging abnormalities of *T. gondii* infection, serum IgG antibody to *T. gondii* should be obtained and, when positive, the patient should be treated.

TREATMENT INFECTIOUS FOCAL CNS LESIONS

Anticonvulsant therapy is initiated when the patient with neurocysticercosis presents with a seizure. There is controversy about whether or not anthelmintic therapy should be given to all patients, and recommendations are based on the stage of the lesion. Cysticerci appearing as cystic lesions in the brain parenchyma with or without pericystic edema or in the subarachnoid space at the convexity of the cerebral hemispheres should be treated with anticysticidal therapy. Cysticidal drugs accelerate the destruction of the parasites, resulting in a faster resolution of the infection. Albendazole and praziquantel are used in the treatment of neurocysticercosis. Approximately 85% of parenchymal cysts are destroyed by a single course of albendazole, and ~75% are destroyed by a single course of praziquantel. The dose of albendazole is 15 mg/kg per day in two doses for 8 days. The dose of praziquantel is 50 mg/kg per day for 15 days, although a number of other dosage regimens are also frequently cited. Prednisone or dexamethasone is given with anticysticidal therapy to reduce the host inflammatory response to degenerating parasites. Many, but not all, experts recommend anticysticidal therapy for lesions that are surrounded by a contrast-enhancing ring. There is universal agreement that calcified lesions do not need to be treated with anticysticidal therapy. Antiepileptic therapy can be stopped once the follow-up CT scan shows resolution of the lesion. Long-term antiepileptic therapy is recommended when seizures occur after resolution of edema and resorption or calcification of the degenerating cyst.

CNS toxoplasmosis is treated with a combination of sulfadiazine, 1.5–2.0 g orally qid, plus pyrimethamine, 100 mg orally to load, then 75–100 mg orally qd, plus folinic acid, 10–15 mg orally qd. Folinic acid is added to the regimen to prevent megaloblastic anemia. Therapy is continued until there is no evidence of active disease on neuroimaging studies, which typically takes at least 6 weeks, and then the dose of sulfadiazine is reduced to 2–4 g/d and pyrimethamine to 50 mg/d. Clindamycin plus pyrimethamine is an alternative therapy for patients who cannot tolerate sulfadiazine, but the combination of pyrimethamine and sulfadiazine is more effective.

SUBDURAL EMPYEMA

A subdural empyema (SDE) is a collection of pus between the dura and arachnoid membranes (Fig. 164-5).

EPIDEMIOLOGY

SDE is a rare disorder that accounts for 15–25% of focal suppurative CNS infections. Sinusitis is the most common predisposing condition

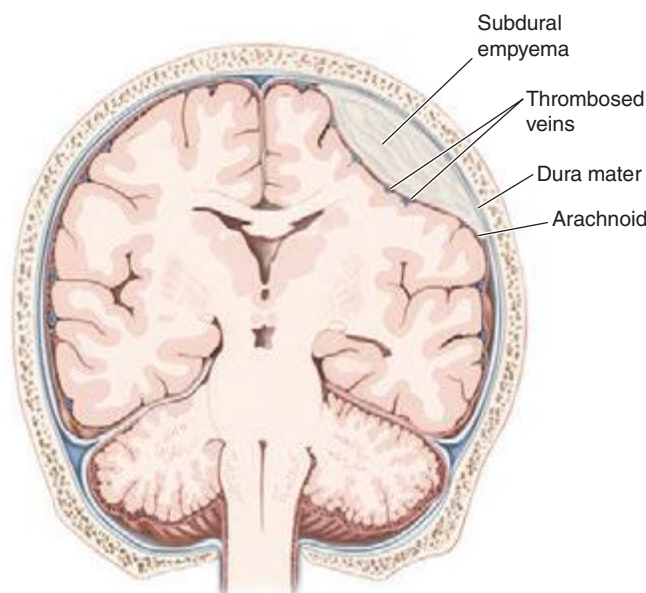


FIGURE 164-5 Subdural empyema.

and typically involves the frontal sinuses, either alone or in combination with the ethmoid and maxillary sinuses. Sinusitis-associated empyema has a striking predilection for young males, possibly reflecting sex-related differences in sinus anatomy and development. It has been suggested that SDE may complicate 1–2% of cases of frontal sinusitis severe enough to require hospitalization. As a consequence of this epidemiology, SDE shows an ~3:1 male/female predominance, with 70% of cases occurring in the second and third decades of life. SDE may also develop as a complication of head trauma or neurosurgery. Secondary infection of a subdural effusion may also result in empyema, although secondary infection of hematomas, in the absence of a prior neurosurgical procedure, is rare.

ETIOLOGY

Aerobic and anaerobic streptococci, staphylococci, Enterobacteriaceae, and anaerobic bacteria are the most common causative organisms of sinusitis-associated SDE. Staphylococci and gram-negative bacilli are often the etiologic organisms when SDE follows neurosurgical procedures or head trauma. Up to one-third of cases are culture-negative, possibly reflecting difficulty in obtaining adequate anaerobic cultures.

PATHOPHYSIOLOGY

Sinusitis-associated SDE develops as a result of either retrograde spread of infection from septic thrombophlebitis of the mucosal veins draining the sinuses or contiguous spread of infection to the brain from osteomyelitis in the posterior wall of the frontal or other sinuses. SDE may also develop from direct introduction of bacteria into the subdural space as a complication of a neurosurgical procedure. The evolution of SDE can be extremely rapid because the subdural space is a large compartment that offers few mechanical barriers to the spread of infection. In patients with sinusitis-associated SDE, suppuration typically begins in the upper and anterior portions of one cerebral hemisphere and then extends posteriorly. SDE is often associated with other intracranial infections, including epidural empyema (40%), cortical thrombophlebitis (35%), and intracranial abscess or cerebritis (>25%). Cortical venous infarction produces necrosis of underlying cerebral cortex and subcortical white matter, with focal neurologic deficits and seizures (see below).

CLINICAL PRESENTATION

A patient with SDE typically presents with fever and a progressively worsening headache. The diagnosis of SDE should always be suspected in a patient with known sinusitis who presents with new CNS signs or symptoms. Patients with underlying sinusitis frequently have symptoms related to this infection. As the infection progresses, focal

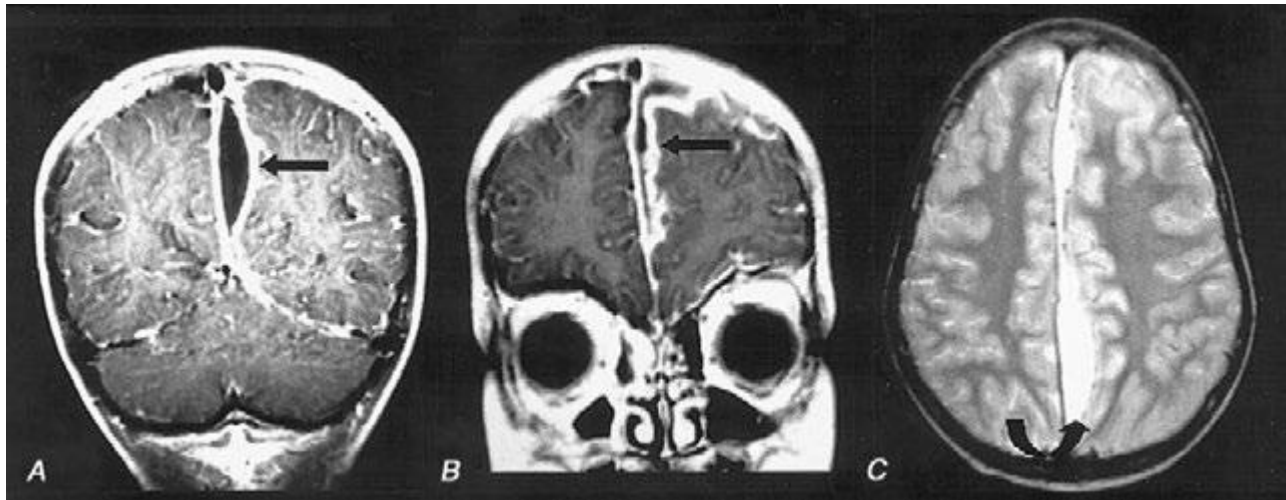


FIGURE 164-6 Subdural empyema. There is marked enhancement of the dura and leptomeninges (**A, B**, straight arrows) along the left medial hemisphere. The pus is hypointense on T1-weighted images (**A, B**) but markedly hyperintense on the proton density-weighted (**C**, curved arrow) image. (Courtesy of Joseph Lurito, MD; with permission.)

neurologic deficits, seizures, nuchal rigidity, and signs of increased ICP commonly occur. Headache is the most common complaint at the time of presentation; initially it is localized to the side of the subdural infection, but then it becomes more severe and generalized. Contralateral hemiparesis or hemiplegia is the most common focal neurologic deficit and can occur from the direct effects of the SDE on the cortex or as a consequence of venous infarction. Seizures begin as partial motor seizures that then become secondarily generalized. Seizures may be due to the direct irritative effect of the SDE on the underlying cortex or result from cortical venous infarction (see above). In untreated SDE, the increasing mass effect and increase in ICP cause progressive deterioration in consciousness, leading ultimately to coma.

DIAGNOSIS

MRI (Fig. 164-6) is superior to CT in identifying SDE and any associated intracranial infections. The administration of gadolinium greatly improves diagnosis by enhancing the rim of the empyema and allowing the empyema to be clearly delineated from the underlying brain parenchyma. Cranial MRI is also extremely valuable in identifying sinusitis, other focal CNS infections, cortical venous infarction, cerebral edema, and cerebritis. CT may show a crescent-shaped hypodense lesion over one or both hemispheres or in the interhemispheric fissure. Frequently the degree of mass effect, exemplified by midline shift, ventricular compression, and sulcal effacement, is far out of proportion to the mass of the SDE.

CSF examination should be avoided in patients with known or suspected SDE because it adds no useful information and is associated with the risk of cerebral herniation.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of the combination of headache, fever, focal neurologic signs, and seizure activity that progresses rapidly to an altered level of consciousness includes subdural hematoma, bacterial meningitis, viral encephalitis, brain abscess, superior sagittal sinus thrombosis, and acute disseminated encephalomyelitis. The presence of nuchal rigidity is unusual with brain abscess or epidural empyema and should suggest the possibility of SDE when associated with significant focal neurologic signs and fever. Patients with bacterial meningitis also have nuchal rigidity but do not typically have focal deficits of the severity seen with SDE.

TREATMENT SUBDURAL EMPYEMA

SDE is a medical emergency. Emergent neurosurgical evacuation of the empyema, either through craniotomy, craniectomy, or burr-hole drainage, is the definitive step in the management of this infection. Empirical antimicrobial therapy for community-acquired SDE

should include a combination of a third-generation cephalosporin (e.g., cefotaxime or ceftriaxone), vancomycin, and metronidazole (see Table 164-1 for dosages). Patients with hospital-acquired SDE may have infections due to *Pseudomonas* spp. or MRSA and should receive coverage with a carbapenem (e.g., meropenem) and vancomycin. Metronidazole is not necessary for antianaerobic therapy when meropenem is being used. Parenteral antibiotic therapy should be continued for a minimum of 3–4 weeks after SDE drainage. Patients with associated cranial osteomyelitis may require longer therapy. Specific diagnosis of the etiologic organisms is made based on Gram's stain and culture of fluid obtained via either burr holes or craniotomy; the initial empirical antibiotic coverage can be modified accordingly.

PROGNOSIS

Prognosis is influenced by the level of consciousness of the patient at the time of hospital presentation, the size of the empyema, and the speed with which therapy is instituted. Long-term neurologic sequelae, which include seizures and hemiparesis, occur in up to 50% of cases.

CRANIAL EPIDURAL ABSCESS

Cranial epidural abscess is a suppurative infection occurring in the potential space between the inner skull table and dura (Fig. 164-7).

ETIOLOGY AND PATHOPHYSIOLOGY

Cranial epidural abscess is less common than either brain abscess or SDE and accounts for <2% of focal suppurative CNS infections. A

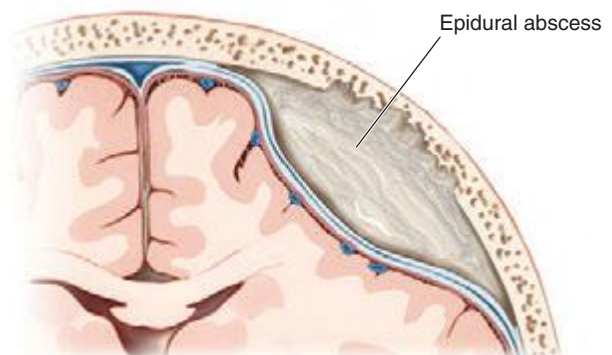


FIGURE 164-7 Cranial epidural abscess is a collection of pus between the dura and the inner table of the skull.

cranial epidural abscess develops as a complication of a craniotomy or compound skull fracture or as a result of spread of infection from the frontal sinuses, middle ear, mastoid, or orbit. An epidural abscess may develop contiguous to an area of osteomyelitis, when craniotomy is complicated by infection of the wound or bone flap, or as a result of direct infection of the epidural space. Infection in the frontal sinus, middle ear, mastoid, or orbit can reach the epidural space through retrograde spread of infection from septic thrombophlebitis in the emissary veins that drain these areas or by way of direct spread of infection through areas of osteomyelitis. Unlike the subdural space, the epidural space is really a potential rather than an actual compartment. The dura is normally tightly adherent to the inner skull table, and infection must dissect the dura away from the skull table as it spreads. As a result, epidural abscesses are often smaller than SDEs. Cranial epidural abscesses, unlike brain abscesses, only rarely result from hematogenous spread of infection from extracranial primary sites. The bacteriology of a cranial epidural abscess is similar to that of SDE (see above). The etiologic organisms of an epidural abscess that arises from frontal sinusitis, middle-ear infections, or mastoiditis are usually streptococci or anaerobic organisms. Staphylococci or gram-negative organisms are the usual cause of an epidural abscess that develops as a complication of craniotomy or compound skull fracture.

CLINICAL PRESENTATION

Patients present with fever (60%), headache (40%), nuchal rigidity (35%), seizures (10%), and focal deficits (5%). Development of symptoms may be insidious, as the empyema usually enlarges slowly in the confined anatomic space between the dura and the inner table of the skull. Periorbital edema and Pott's puffy tumor, reflecting underlying associated frontal bone osteomyelitis, are present in ~40%. In patients with a recent neurosurgical procedure, wound infection is invariably present, but other symptoms may be subtle and can include altered mental status (45%), fever (35%), and headache (20%). The diagnosis should be considered when fever and headache follow recent head trauma or occur in the setting of frontal sinusitis, mastoiditis, or otitis media.

DIAGNOSIS

Cranial MRI with gadolinium enhancement is the procedure of choice to demonstrate a cranial epidural abscess. The sensitivity of CT is limited by the presence of signal artifacts arising from the bone of the inner skull table. The CT appearance of an epidural empyema is that of a lens or crescent-shaped hypodense extraaxial lesion. On MRI, an epidural empyema appears as a lentiform or crescent-shaped fluid collection that is hyperintense compared to CSF on T2-weighted images. On T1-weighted images, the fluid collection may be either isointense or hypointense compared to brain. Following the administration of gadolinium, there is linear enhancement of the dura on T1-weighted images. In distinction to subdural empyema, signs of mass effect or other parenchymal abnormalities are uncommon.

TREATMENT EPIDURAL ABSCESS

Immediate neurosurgical drainage is indicated. Empirical antimicrobial therapy, pending the results of Gram's stain and culture of the purulent material obtained at surgery, should include a combination of a third-generation cephalosporin, vancomycin, and metronidazole (Table 164-1). Ceftazidime or meropenem should be substituted for ceftriaxone or cefotaxime in neurosurgical patients. Metronidazole is not necessary for antianaerobic coverage in patients receiving meropenem. When the organism has been identified, antimicrobial therapy can be modified accordingly. Antibiotics should be continued for 3–6 weeks after surgical drainage. Patients with associated osteomyelitis may require additional therapy.

PROGNOSIS

The mortality rate is <5% in modern series, and full recovery is the rule in most survivors.

SUPPURATIVE THROMBOPHLEBITIS

DEFINITION

Suppurative intracranial thrombophlebitis is septic venous thrombosis of cortical veins and sinuses. This may occur as a complication of bacterial meningitis; SDE; epidural abscess; or infection in the skin of the face, paranasal sinuses, middle ear, or mastoid.

ANATOMY AND PATHOPHYSIOLOGY

The cerebral veins and venous sinuses have no valves; therefore, blood within them can flow in either direction. The superior sagittal sinus is the largest of the venous sinuses (Fig. 164-8). It receives blood from the frontal, parietal, and occipital superior cerebral veins and the diploic veins, which communicate with the meningeal veins. Bacterial meningitis is a common predisposing condition for septic thrombosis of the superior sagittal sinus. The diploic veins, which drain into the superior sagittal sinus, provide a route for the spread of infection from the meninges, especially in cases where there is purulent exudate near areas of the superior sagittal sinus. Infection can also spread to the superior sagittal sinus from nearby SDE or epidural abscess. Dehydration from vomiting, hypercoagulable states, and immunologic abnormalities, including the presence of circulating antiphospholipid antibodies, also contribute to cerebral venous sinus thrombosis. Thrombosis may extend from one sinus to another, and at autopsy, thrombi of different histologic ages can often be detected in several sinuses. Thrombosis of the superior sagittal sinus is often associated with thrombosis of superior cortical veins and small parenchymal hemorrhages.

The superior sagittal sinus drains into the transverse sinuses (Fig. 164-8). The transverse sinuses also receive venous drainage from small veins from both the middle ear and mastoid cells. The transverse sinus becomes the sigmoid sinus before draining into the internal jugular vein. Septic transverse/sigmoid sinus thrombosis can be a complication of acute and chronic otitis media or mastoiditis. Infection spreads from the mastoid air cells to the transverse sinus via the emissary veins or by direct invasion. The cavernous sinuses are inferior to the superior sagittal sinus at the base of the skull. The cavernous sinuses receive blood from the facial veins via the superior and inferior ophthalmic veins. Bacteria in the facial veins enter the cavernous sinus via these veins. Bacteria in the sphenoid and ethmoid sinuses can spread to the cavernous sinuses via the small emissary veins. The sphenoid and ethmoid sinuses are the most common sites of primary infection resulting in septic cavernous sinus thrombosis.

CLINICAL MANIFESTATIONS

Septic thrombosis of the superior sagittal sinus presents with headache, fever, nausea and vomiting, confusion, and focal or generalized seizures. There may be a rapid development of stupor and coma.

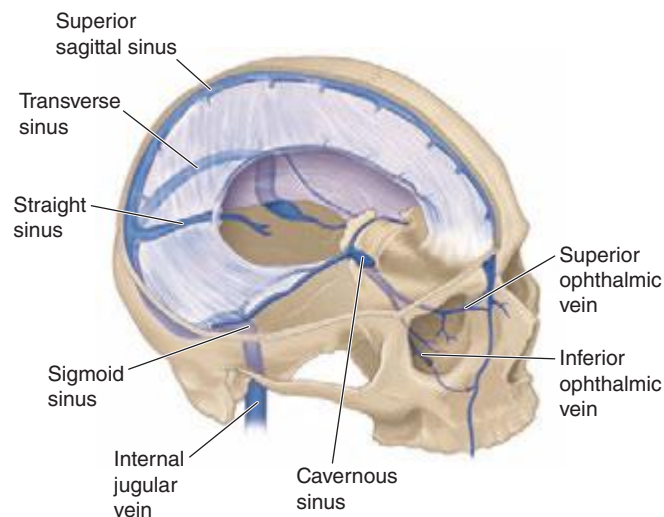


FIGURE 164-8 Anatomy of the cerebral venous sinuses.

906 Weakness of the lower extremities with bilateral Babinski's signs or hemiparesis is often present. When superior sagittal sinus thrombosis occurs as a complication of bacterial meningitis, nuchal rigidity and Kernig's and Brudzinski's signs may be present.

The oculomotor nerve, the trochlear nerve, the abducens nerve, the ophthalmic and maxillary branches of the trigeminal nerve, and the internal carotid artery all pass through the cavernous sinus (see Fig. 455-4). The symptoms of *septic cavernous sinus thrombosis* are fever, headache, frontal and retroorbital pain, and diplopia. The classic signs are ptosis, proptosis, chemosis, and extraocular dysmotility due to deficits of cranial nerves III, IV, and VI; hyperesthesia of the ophthalmic and maxillary divisions of the fifth cranial nerve and a decreased corneal reflex may be detected. There may be evidence of dilated, tortuous retinal veins and papilledema.

Headache and earache are the most frequent symptoms of *transverse sinus thrombosis*. A transverse sinus thrombosis may also present with otitis media, sixth nerve palsy, and retroorbital or facial pain (*Gradenigo's syndrome*). Sigmoid sinus and internal jugular vein thrombosis may present with neck pain.

DIAGNOSIS

The diagnosis of septic venous sinus thrombosis is suggested by an absent flow void within the affected venous sinus on MRI and confirmed by magnetic resonance venography, CT angiogram, or the venous phase of cerebral angiography. The diagnosis of thrombophlebitis of intracerebral and meningeal veins is suggested by the presence of intracerebral hemorrhage but requires cerebral angiography for definitive diagnosis.

TREATMENT SUPPURATIVE THROMBOPHLEBITIS

Septic venous sinus thrombosis is treated with antibiotics, hydration, and removal of infected tissue and thrombus in septic lateral or cavernous sinus thrombosis. The choice of antimicrobial therapy is based on the bacteria responsible for the predisposing or associated condition. Optimal duration of therapy is unknown, but antibiotics are usually continued for 6 weeks or until there is radiographic evidence of resolution of thrombosis. Anticoagulation with dose-adjusted intravenous heparin is recommended for aseptic venous sinus thrombosis and in the treatment of septic venous sinus thrombosis complicating bacterial meningitis in patients who have progressive neurologic deterioration despite antimicrobial therapy and intravenous fluids. The presence of a small intracerebral hemorrhage from septic thrombophlebitis is not an absolute contraindication to heparin therapy. Successful management of aseptic venous sinus thrombosis has been reported with surgical thrombectomy, catheter-directed urokinase therapy, and a combination of intrathrombus recombinant tissue plasminogen activator (rtPA) and intravenous heparin, but there are not enough data to recommend these therapies in septic venous sinus thrombosis.

165 Chronic and Recurrent Meningitis

Walter J. Koroshetz, Avindra Nath

Chronic inflammation of the meninges (pia, arachnoid, and dura) can produce profound neurologic disability and may be fatal if not successfully treated. Chronic meningitis is diagnosed when a characteristic neurologic syndrome exists for >4 weeks and is associated with a persistent inflammatory response in the cerebrospinal fluid (CSF) (white blood cell count >5/ μ L). The causes are varied, and appropriate treatment depends on identification of the etiology. Five categories of disease account for most cases of chronic meningitis: (1) meningeal infections, (2) malignancy, (3) autoimmune inflammatory disorders, (4) chemical meningitis, and (5) parameningeal infections.

TABLE 165-1 SYMPTOMS AND SIGNS OF CHRONIC MENINGITIS

Symptom	Sign
Chronic headache	\pm Papilledema
Neck or back pain/stiffness	Brudzinski's or Kernig's sign of meningeal irritation
Change in personality	Altered mental status—drowsiness, inattention, disorientation, memory loss, frontal release signs (grasp, suck, snout), perseveration
Facial weakness	Peripheral seventh CN paresis
Double vision	Paresis of CNs III, IV, VI
Diminished Vision	Papilledema, optic atrophy
Hearing loss	Eighth CN paresis
Arm or leg weakness	Myelopathy or radiculopathy
Numbness in arms or legs	Myelopathy or radiculopathy
Urinary retention/incontinence	Myelopathy or radiculopathy
Clumsiness	Frontal lobe dysfunction (hydrocephalus)
	Ataxia

Abbreviation: CN, cranial nerve.

CLINICAL PATHOPHYSIOLOGY

Neurologic manifestations of chronic meningitis (**Table 165-1**) are determined by the anatomic location of the inflammation and its consequences. Persistent headache with or without a stiff neck, hydrocephalus, cranial neuropathies, radiculopathies, and cognitive or personality changes are the cardinal features. These can occur alone or in combination. When they appear in combination, widespread dissemination of the inflammatory process along CSF pathways has occurred. In some cases, the presence of an underlying systemic illness points to a specific agent or class of agents as the probable cause. The diagnosis of chronic meningitis is usually made when the clinical presentation prompts the physician to examine the CSF for signs of inflammation. CSF is produced by the choroid plexus of the cerebral ventricles, exits through narrow foramina into the subarachnoid space surrounding the brain and spinal cord, circulates around the base of the brain and over the cerebral hemispheres, and is resorbed by arachnoid villi projecting into the superior sagittal sinus. CSF flow provides a pathway for rapid spread of infectious and other infiltrative processes over the brain, spinal cord, and cranial and spinal nerve roots. Spread from the subarachnoid space into brain parenchyma may occur via the arachnoid cuffs that surround blood vessels that penetrate brain tissue (Virchow-Robin spaces).

Intracranial Meningitis Nociceptive nerve fibers of the meninges are stimulated by the inflammatory process, resulting in headache, neck pain, or back pain. Obstruction of CSF pathways at the foramina or arachnoid villi may produce *hydrocephalus* and symptoms of raised intracranial pressure (ICP), including headache, vomiting, apathy or drowsiness, gait instability, papilledema, visual loss, impaired upgaze, or palsy of the sixth cranial nerve (CN) (**Chap. 455**). Cognitive and behavioral changes during the course of chronic meningitis may also result from vascular damage, which may similarly produce seizures, stroke, or myelopathy. Inflammatory deposits seeded via the CSF circulation are often prominent around the brainstem and cranial nerves and along the undersurface of the frontal and temporal lobes. Such cases, termed *basal meningitis*, often present as multiple cranial neuropathies, with decreased vision (CN II), facial weakness (CN VII), decreased hearing (CN VIII), diplopia (CNs III, IV, and VI), sensory or motor abnormalities of the oropharynx (CNs IX, X, and XII), decreased olfaction (CN I), or decreased facial sensation and masseter weakness (CN V).

Spinal Meningitis Injury may occur to motor and sensory roots as they traverse the subarachnoid space and penetrate the meninges. These cases present as multiple radiculopathies with combinations of radicular pain, sensory loss, motor weakness, and urinary or fecal incontinence. Meningeal inflammation can encircle the cord, resulting in a myelopathy. Patients with slowly progressive involvement of multiple cranial nerves and/or spinal nerve roots are likely to have chronic

meningitis. Electrophysiologic testing (electromyography, nerve conduction studies, and evoked response testing) may be helpful in determining whether there is involvement of cranial and spinal nerve roots.

Systemic Manifestations In some patients, evidence of systemic disease provides clues to the underlying cause of chronic meningitis. A complete history of travel, sexual practice, and exposure to infectious agents should be sought. Infectious causes are often associated with fever, malaise, anorexia, and signs of localized or disseminated infection outside the nervous system. Infectious causes are of major concern in the immunosuppressed patient, especially in patients with AIDS, in whom chronic meningitis may present without headache or fever. Noninfectious inflammatory disorders often produce systemic manifestations, but meningitis may be the initial manifestation. Carcinomatous meningitis may or may not be accompanied by clinical evidence of the primary neoplasm.

APPROACH TO THE PATIENT: Chronic Meningitis

The occurrence of chronic headache, hydrocephalus, cranial neuropathy, radiculopathy, and/or cognitive decline in a patient should prompt consideration of a lumbar puncture for evidence of meningeal inflammation. On occasion, the diagnosis is made when an imaging study (CT or MRI) shows contrast enhancement of the meninges, which is always concerning with the exception of dural enhancement after lumbar puncture, neurosurgical procedures, or spontaneous CSF leakage. Once chronic meningitis is confirmed by CSF examination, effort is focused on identifying the cause (Tables 165-2 and 165-3) by (1) further analysis of the CSF, (2) diagnosis of an underlying systemic infection or noninfectious inflammatory condition, or (3) pathologic examination of meningeal biopsy specimens.

Two clinical forms of chronic meningitis exist. In the first, the symptoms are chronic and persistent, whereas in the second there are recurrent, discrete episodes of illness. In the latter group, all symptoms, signs, and CSF parameters of meningeal inflammation resolve completely between episodes without specific therapy. In such patients, the likely etiologies include herpes simplex virus (HSV) type 2; chemical meningitis due to episodic leakage from an epidermoid tumor, craniopharyngioma, or cholesteatoma into CSF; primary autoimmune inflammatory conditions, including Vogt-Koyanagi-Harada syndrome, Behçet's syndrome, systemic lupus erythematosus (SLE), and Mollaret's meningitis; and drug hypersensitivity with repeated administration of the offending agent.

The epidemiologic history is of considerable importance and may provide direction for selection of laboratory studies. Pertinent features include a history of tuberculosis or exposure to a likely case; past travel to areas endemic for fungal infections (the San Joaquin Valley in California and southwestern states for coccidioidomycosis, midwestern states for histoplasmosis, southeastern states for blastomycosis); travel to the Mediterranean region or ingestion of imported unpasteurized dairy products (*Brucella*); time spent in wooded areas endemic for Lyme disease; exposure to sexually transmitted disease (syphilis); exposure of an immunocompromised host to pigeons and their droppings (*Cryptococcus*); gardening (*Sporothrix schenckii*); ingestion of poorly cooked meat or contact with a household cat (*Toxoplasma gondii*); residence in Thailand or Japan (*Gnathostoma spinigerum*), Latin America (*Paracoccidioides brasiliensis*), or the South Pacific (*Angiostrongylus cantonensis*); rural residence and raccoon exposure (*Baylisascaris procyonis*); and residence in Latin America, the Philippines, or Southeast Asia (*Taenia solium/cysticercosis*).

The presence of focal cerebral signs in a patient with chronic meningitis suggests the possibility of a brain abscess or other

TABLE 165-2 INFECTIOUS CAUSES OF CHRONIC MENINGITIS

Causative Agent	CSF Formula	Helpful Diagnostic Tests	Risk Factors and Systemic Manifestations
Common Bacterial Causes			
Partially treated suppurative meningitis	Mononuclear or mixed mononuclear-polymorphonuclear cells	CSF culture and Gram's stain	History consistent with acute bacterial meningitis and incomplete treatment
Parameningeal infection	Mononuclear or mixed polymorphonuclear-mononuclear cells	Contrast-enhanced CT or MRI to detect parenchymal, subdural, epidural, or sinus infection	Otitis media, pleuropulmonary infection, right-to-left cardiopulmonary shunt for brain abscess; focal neurologic signs; neck, back, ear, or sinus tenderness
<i>Mycobacterium tuberculosis</i>	Mononuclear cells except polymorphonuclear cells in early infection (commonly <500 WBC/ μ L); low CSF glucose, high protein	Tuberculin skin test may be negative; AFB culture of CSF (sputum, urine, gastric contents if indicated); tuberculostearic acid detection in CSF; identify tubercle bacillus on acid-fast stain of CSF or protein pellicle; PCR	Exposure history; previous tuberculous illness; immunosuppressed, anti-TNF therapy or AIDS; young children; fever, meningismus, night sweats, miliary TB on x-ray or liver biopsy; stroke due to arteritis
Lyme disease (Bannwarth's syndrome) <i>Borrelia burgdorferi</i>	Mononuclear cells; elevated protein	Serum Lyme antibody titer; Western blot confirmation; (patients with syphilis may have false-positive Lyme titer)	History of tick bite or appropriate exposure history; erythema chronicum migrans skin rash; arthritis, radiculopathy, Bell's palsy, meningoen- cephalitis—multiple sclerosis-like syndrome
Syphilis (secondary, tertiary) <i>Treponema pallidum</i>	Mononuclear cells; elevated protein	CSF VDRL; serum VDRL (or RPR); fluorescent treponemal antibody-absorbed (FTA) or MHA-TP; serum VDRL may be negative in tertiary syphilis	Appropriate exposure history; HIV-seropositive individuals at increased risk of aggressive infection; "dementia"; cerebral infarction due to endarteritis
Uncommon Bacterial Causes			
<i>Actinomyces</i>	Polymorphonuclear cells	Anaerobic culture	Parameningeal abscess or sinus tract (oral or dental focus); pneumonitis
<i>Nocardia</i>	Polymorphonuclear; occasionally mononuclear cells; often low glucose	Isolation may require weeks; weakly acid fast	Associated brain abscess may be present
<i>Brucella</i>	Mononuclear cells (rarely polymorphonuclear); elevated protein; often low glucose	CSF antibody detection; serum antibody detection	Intake of unpasteurized dairy products; exposure to goats, sheep, cows; fever, arthralgia, myalgia, vertebral osteomyelitis
Whipple's disease <i>Tropheryma whipplei</i>	Mononuclear cells	Biopsy of small bowel or lymph node; CSF PCR for <i>T. whipplei</i> ; brain and meningeal biopsy (with PAS stain and EM examination)	Diarrhea, weight loss, arthralgias, fever; dementia, ataxia, paresis, ophthalmoplegia, oculomastocytary myoclonus

(Continued)

TABLE 165-2 INFECTIOUS CAUSES OF CHRONIC MENINGITIS (CONTINUED)

Causative Agent	CSF Formula	Helpful Diagnostic Tests	Risk Factors and Systemic Manifestations
Rare Bacterial Causes			
Leptospirosis (occasionally if left untreated may last 3–4 weeks)			
Fungal Causes			
<i>Cryptococcus neoformans</i>	Mononuclear cells; count not elevated in some patients with AIDS	India ink or fungal wet mount of CSF (budding yeast); blood and urine cultures; antigen detection in CSF	AIDS and immune suppression; pigeon exposure; skin and other organ involvement due to disseminated infection
<i>Coccidioides immitis</i>	Mononuclear cells (sometimes 10–20% eosinophils); often low glucose	Antibody detection in CSF and serum	Exposure history—southwestern U.S.; increased virulence in dark-skinned races
<i>Candida</i> sp.	Polymorphonuclear or mononuclear	Fungal stain and culture of CSF	IV drug abuse; post surgery; prolonged IV therapy; disseminated candidiasis
<i>Histoplasma capsulatum</i>	Mononuclear cells; low glucose	Fungal stain and culture of large volumes of CSF; antigen detection in CSF, serum, and urine; antibody detection in serum, CSF	Exposure history—Ohio and central Mississippi River Valley; AIDS; mucosal lesions
<i>Blastomyces dermatitidis</i>	Mononuclear cells	Fungal stain and culture of CSF; biopsy and culture of skin, lung lesions; antibody detection in serum	Midwestern and southeastern U.S.; usually systemic infection; abscesses, draining sinus, ulcers
<i>Aspergillus</i> sp.	Mononuclear or polymorphonuclear	CSF culture	Sinusitis; granulocytopenia or immunosuppression
<i>Sporothrix schenckii</i>	Mononuclear cells	Antibody detection in CSF and serum; CSF culture	Traumatic inoculation; IV drug use; ulcerated skin lesion
Rare Fungal Causes			
<i>Xylohypha</i> (formerly <i>Cladosporium</i>) <i>trichoides</i> and other dark-walled (dematiaceous) fungi such as <i>Curvularia</i> , <i>Drechslera</i> ; <i>Mucor</i> , and, after water aspiration, <i>Pseudallescheria boydii</i> , iatrogenic <i>Exserohilum rostratum</i> infection following spinal blocks			
Protozoal Causes			
<i>Toxoplasma gondii</i>	Mononuclear cells	Biopsy or response to empirical therapy in clinically appropriate context (including presence of antibody in serum)	Usually with intracerebral abscesses; common in HIV-seropositive patients
Trypanosomiasis <i>Trypanosoma gambiense</i> , <i>T. rhodesiense</i>	Mononuclear cells, elevated protein	Elevated CSF IgM; identification of trypanosomes in CSF and blood smear	Endemic in Africa; chancre, lymphadenopathy; prominent sleep disorder
Rare Protozoal Causes			
<i>Acanthamoeba</i> sp. causing granulomatous amebic encephalitis and meningoencephalitis in immunocompromised and debilitated individuals. <i>Balamuthia mandrillaris</i> causing chronic meningoencephalitis in immunocompetent hosts.			
Helminthic Causes			
Cysticercosis (infection with cysts of <i>Taenia solium</i>)	Mononuclear cells; may have eosinophils; glucose level may be low	Indirect hemagglutination assay in CSF; ELISA immunoblotting in serum	Usually with multiple cysts in basal meninges and hydrocephalus; cerebral cysts, muscle calcification
<i>Gnathostoma spinigerum</i>	Eosinophils, mononuclear cells	Peripheral eosinophilia	History of eating raw fish; common in Thailand and Japan; subarachnoid hemorrhage; painful radiculopathy
<i>Angiostrongylus cantonensis</i>	Eosinophils, mononuclear cells	Recovery of worms from CSF	History of eating raw shellfish; common in tropical Pacific regions; often benign
<i>Baylisascaris procyonis</i> (raccoon ascarid)	Eosinophils, mononuclear cells		Infection follows accidental ingestion of <i>B. procyonis</i> eggs from raccoon feces; fatal meningoencephalitis
Rare Helminthic Causes			
<i>Trichinella spiralis</i> (trichinosis); <i>Fasciola hepatica</i> (liver fluke), <i>Echinococcus</i> cysts; <i>Schistosoma</i> sp. The former may produce a lymphocytic pleocytosis whereas the latter two may produce an eosinophilic response in CSF associated with cerebral cysts (<i>Echinococcus</i>) or granulomatous lesions of brain or spinal cord			
Viral Causes			
Mumps	Mononuclear cells	Antibody in serum	No prior mumps or immunization; may produce meningoencephalitis; may persist for 3–4 weeks
Lymphocytic choriomeningitis	Mononuclear cells	Antibody in serum	Contact with rodents or their excreta; may persist for 3–4 weeks
Echovirus	Mononuclear cells; may have low glucose	Virus isolation from CSF	Congenital hypogammaglobulinemia; history of recurrent meningitis
HIV (acute retroviral syndrome)	Mononuclear cells	p24 antigen in serum and CSF; high level of HIV viremia.	HIV risk factors; rash, fever, lymphadenopathy; lymphopenia in peripheral blood; syndrome may persist long enough to be considered as “chronic meningitis”; or chronic meningitis may develop in later stages (AIDS) due to HIV
Herpes simplex (HSV)	Mononuclear cells	PCR for HSV, CMV DNA; CSF antibody for HSV, EBV	Recurrent meningitis due to HSV-2 (rarely HSV-1) often associated with genital recurrences; EBV associated with myeloradiculopathy, CMV with polyradiculopathy

Abbreviations: AFB, acid-fast bacillus; CMV, cytomegalovirus; CSF, cerebrospinal fluid; CT, computed tomography; EBV, Epstein-Barr virus; ELISA, enzyme-linked immunosorbent assay; EM, electron microscopy; FTA, fluorescent treponemal antibody absorption test; HSV, herpes simplex virus; MHA-TP, microhemagglutination assay—*T. pallidum*; MRI, magnetic resonance imaging; PAS, periodic acid–Schiff; PCR, polymerase chain reaction; RPR, rapid plasma reagin test; TB, tuberculosis; VDRL, Venereal Disease Research Laboratories test.

TABLE 165-3 NONINFECTIOUS CAUSES OF CHRONIC MENINGITIS

Causative Agents	CSF Formula	Helpful Diagnostic Tests	Risk Factors and Systemic Manifestations
Malignancy	Mononuclear cells, elevated protein, low glucose	Repeated cytologic examination of large volumes of CSF; CSF exam by polarizing microscopy; clonal lymphocyte markers; deposits on nerve roots or meninges seen on myelogram or contrast-enhanced MRI; meningeal biopsy	Metastatic cancer of breast, lung, stomach, or pancreas; melanoma, lymphoma, leukemia; meningeal gliomatosis; meningeal sarcoma; cerebral dysgerminoma; meningeal melanoma or B cell lymphoma
Chemical compounds (may cause recurrent meningitis)	Mononuclear or PMNs, low glucose, elevated protein; xanthochromia from subarachnoid hemorrhage in week prior to presentation with "meningitis"	Contrast-enhanced CT scan or MRI; cerebral angiogram to detect aneurysm	History of recent injection into the subarachnoid space; history of sudden onset of headache; recent resection of acoustic neuroma or craniopharyngioma; epidermoid tumor of brain or spine, sometimes with dermoid sinus tract; pituitary apoplexy
Primary inflammation			
CNS sarcoidosis	Mononuclear cells; elevated protein; often low glucose	Serum and CSF angiotensin-converting enzyme levels; biopsy of extraneural affected tissues or brain lesion/meningeal biopsy	CN palsy, especially of CN VII; hypothalamic dysfunction, especially diabetes insipidus; abnormal chest radiograph; peripheral neuropathy or myopathy
Vogt-Koyanagi-Harada syndrome (recurrent meningitis)	Mononuclear cells		Recurrent meningoencephalitis with uveitis, retinal detachment, alopecia, lightening of eyebrows and lashes, dysacusia, cataracts, glaucoma
Isolated granulomatous angiitis of the nervous system	Mononuclear cells, elevated protein	Angiography or meningeal biopsy	Subacute dementia; multiple cerebral infarctions; recent zoster ophthalmicus
Systemic lupus erythematosus	Mononuclear or PMNs	Anti-DNA antibody, antinuclear antibodies	Encephalopathy; seizures; stroke; transverse myelopathy; rash; arthritis
Behçet's syndrome (recurrent meningitis)	Mononuclear or PMNs, elevated protein		Oral and genital aphthous ulcers; iridocyclitis; retinal hemorrhages; pathergic lesions at site of skin puncture
Chronic benign lymphocytic meningitis	Mononuclear cells		Recovery in 2–6 months, diagnosis by exclusion
Mollaret's meningitis (recurrent meningitis)	Large endothelial cells and PMNs in first hours, followed by mononuclear cells	PCR for herpes; MRI/CT to rule out epidermoid tumor or dural cyst	Recurrent meningitis; exclude HSV-2; rare cases due to HSV-1; occasional case associated with dural cyst
Drug hypersensitivity	PMNs; occasionally mononuclear cells or eosinophils	Complete blood count (eosinophilia)	Exposure to nonsteroidal anti-inflammatory agents, sulfonamides, isoniazid, tolmetin, ciprofloxacin, penicillin, carbamazepine, lamotrigine, IV immunoglobulin, OKT3 antibodies, phenazopyridine; improvement after discontinuation of drug; recurrence with repeat exposure
Granulomatosis with polyangiitis (Wegener's)	Mononuclear cells	Chest and sinus radiographs; urinalysis; ANCA antibodies in serum	Associated sinus, pulmonary, or renal lesions; CN palsies; skin lesions; peripheral neuropathy

Other: multiple sclerosis, Sjögren's syndrome, monogenic autoinflammatory disorders, and rarer forms of vasculitis (e.g., Cogan's syndrome)

Abbreviations: ANCA, antineutrophil cytoplasmic antibodies; CN, cranial nerve; CSF, cerebrospinal fluid; CT, computed tomography; HSV, herpes simplex virus; MRI, magnetic resonance imaging; PCR, polymerase chain reaction; PMNs, polymorphonuclear cells.

parameningeal infection; identification of a potential source of infection (chronic draining ear, sinusitis, right-to-left cardiac or pulmonary shunt, chronic pleuropulmonary infection) supports this diagnosis. In some cases, diagnosis may be established by recognition and biopsy of unusual skin lesions (Behçet's syndrome, SLE, cryptococcosis, blastomycosis, Lyme disease, sporotrichosis, trypanosomiasis, IV drug use) or enlarged lymph nodes (lymphoma, sarcoid, tuberculosis, HIV, secondary syphilis, or Whipple's disease). A careful ophthalmologic examination may reveal uveitis (Vogt-Koyanagi-Harada syndrome, sarcoid, or central nervous system [CNS] lymphoma), keratoconjunctivitis sicca (Sjögren's syndrome), or iridocyclitis (Behçet's syndrome) and is essential to assess visual loss from papilledema. Aphthous oral lesions, genital ulcers, and hypopyon suggest Behçet's syndrome. Hepatosplenomegaly suggests lymphoma, sarcoid, tuberculosis, or brucellosis. Herpetic lesions in the genital area or on the thighs suggest HSV-2 infection. A breast nodule, a suspicious pigmented skin lesion, focal bone pain, or an abdominal mass directs attention to possible carcinomatous meningitis.

IMAGING

Once the clinical syndrome is recognized as a potential manifestation of chronic meningitis, proper analysis of the CSF is essential. However, if the possibility of raised ICP exists, a brain imaging study should be performed before lumbar puncture. If ICP is elevated because of a mass lesion, brain swelling, or a block in ventricular CSF

outflow (obstructive hydrocephalus), then lumbar puncture carries the potential risk of brain herniation. Obstructive hydrocephalus usually requires direct ventricular drainage. In patients with open CSF flow pathways, elevated ICP can still occur due to impaired resorption of CSF by arachnoid villi. In such patients, lumbar puncture is usually safe, but repetitive or continuous lumbar drainage may be necessary to prevent abrupt deterioration and death from raised ICP. In some patients, especially those with cryptococcal meningitis, fatal levels of raised ICP can occur without enlarged ventricles.

Contrast-enhanced MRI or CT studies of the brain and spinal cord can identify meningeal enhancement, parameningeal infections (including brain abscess), encasement of the spinal cord (malignancy, inflammation or infection), or nodular deposits on the meninges or nerve roots (malignancy or sarcoidosis) (Fig. 165-1). Imaging studies are also useful to localize areas of meningeal disease prior to meningeal biopsy.

Angiographic studies can identify evidence of cerebral arteritis in patients with chronic meningitis and stroke.

CEREBROSPINAL FLUID ANALYSIS

The CSF pressure should be measured and samples sent for bacterial, fungal, and tuberculous culture; Venereal Disease Research Laboratories (VDRL) test; cell count and differential; Gram's stain; and measurement of glucose and protein. Wet mount for fungus and parasites, india ink preparation and culture, culture

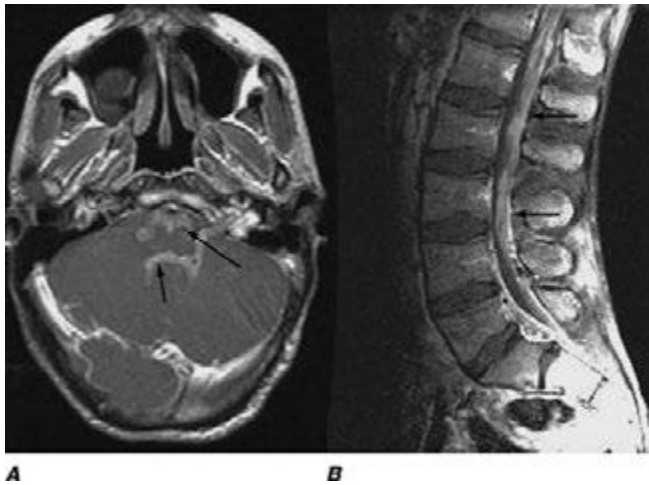


FIGURE 165-1 Primary central nervous system lymphoma. A 24-year-old man, immunosuppressed due to intestinal lymphangiectasia, developed multiple cranial neuropathies. CSF findings consisted of 100 lymphocytes/ μL and a protein of 2.5 g/L (250 mg/dL); cytology and cultures were negative. Gadolinium-enhanced T1 MRI revealed diffuse, multifocal meningeal enhancement surrounding the brainstem (A), spinal cord, and cauda equina (B).

for fastidious bacteria and fungi, assays for cryptococcal antigen and oligoclonal immunoglobulin bands, and cytology should be performed. Other specific CSF tests (Tables 165-2 and 165-3) or blood tests and cultures should be ordered as indicated on the basis of the history, physical examination, or preliminary CSF results (i.e., eosinophilic, mononuclear, or polymorphonuclear meningitis). Rapid diagnosis may be facilitated by serologic tests and polymerase chain reaction (PCR) testing to identify DNA sequences in the CSF that are specific for the suspected pathogen. In patients with suspected fungal infections, when other tests are negative, assays for beta-glucan may be a useful adjunct in establishing the diagnosis.

In most categories of chronic (not recurrent) meningitis, mononuclear cells predominate in the CSF. When neutrophils predominate after 3 weeks of illness, the principal etiologic considerations are *Nocardia asteroides*, *Actinomyces israelii*, *Brucella*, *Mycobacterium tuberculosis* (5–10% of early cases only), various fungi (*Blastomyces dermatitidis*, *Candida albicans*, *Histoplasma capsulatum*, *Aspergillus* spp., *Pseudallescheria boydii*, *Cladophialophora bantiana*), and non-infectious causes (SLE, exogenous chemical meningitis). When eosinophils predominate or are present in limited numbers in a primarily mononuclear cell response in the CSF, the differential diagnosis includes parasitic diseases (*A. cantonensis*, *G. spinigerum*, *B. procyonis*, or *Toxocara canis* infection), cysticercosis, schistosomiasis, echinococcal disease, *T. gondii* infection), fungal infections (6–20% eosinophils along with a predominantly lymphocyte pleocytosis, particularly with coccidioid meningitis), neoplastic disease (lymphoma, leukemia, metastatic carcinoma), or other inflammatory processes (sarcoidosis, hypereosinophilic syndrome).

It is often necessary to broaden the number of diagnostic tests if the initial workup does not reveal the cause. In addition, repeated samples of large volumes of CSF may be required to diagnose certain infectious and malignant causes of chronic meningitis. Flow cytometry for malignant cells may be useful in patients with suspected carcinomatous meningitis. Lymphomatous or carcinomatous meningitis may be diagnosed by examination of sections cut from a cell block formed by spinning down the sediment from a large volume of CSF. The diagnosis of fungal meningitis may require large volumes of CSF for culture of sediment. If standard lumbar puncture is unrewarding, a cervical cisternal tap to sample CSF near to the basal meninges may be fruitful.

LABORATORY INVESTIGATION

In addition to the CSF examination, an attempt should be made to uncover pertinent underlying illnesses. Tuberculin skin test, chest

radiograph, urine analysis and culture, blood count and differential, renal and liver function tests, alkaline phosphatase, sedimentation rate, antinuclear antibody, anti-Ro antibody, anti-La antibody, and serum angiotensin-converting enzyme level are often indicated. In some cases, a thorough search for a systemic site of infection is indicated. Pulmonary foci of infection may be present, particularly with fungal or tuberculous disease. Hence a CT or MRI of the chest and a sputum examination may be helpful. Abnormalities can be pursued by bronchoscopy or transthoracic needle biopsy. A tuberculin skin test is often placed, although the test has limited specificity and sensitivity for diagnosis of active disease. Liver or bone marrow biopsy may be diagnostic in some cases of miliary tuberculosis, disseminated fungal infection, sarcoidosis, or metastatic malignancy. Positron emission tomography with fluorodeoxyglucose may be useful in identifying a systemic site for biopsy in patients with suspected carcinomatous meningitis or sarcoidosis when other tests are unrevealing. Genetic testing can identify mutations that cause rare monogenic autoinflammatory disorders.

MENINGEAL BIOPSY

If CSF is not diagnostic then a meningeal biopsy should be strongly considered in patients who are severely disabled, who need chronic ventricular decompression, or whose illness is progressing rapidly. The activities of the surgeon, pathologist, microbiologist, and cytologist should be coordinated so that a large enough sample is obtained and the appropriate cultures and histologic and molecular studies, including electron-microscopic and PCR studies, are performed. The diagnostic yield of meningeal biopsy can be increased by targeting regions that enhance with contrast on MRI or CT. With current microsurgical techniques, most areas of the basal meninges can be accessed for biopsy via a limited craniotomy. In a series from the Mayo Clinic reported by TM Cheng et al. (*Neurosurgery* 34:590, 1994), MRI demonstrated meningeal enhancement in 47% of patients undergoing meningeal biopsy. Biopsy of an enhancing region was diagnostic in 80% of cases; biopsy of nonenhancing regions was diagnostic in only 9%; sarcoid (31%) and metastatic adenocarcinoma (25%) were the most common conditions identified. Tuberculosis is the most common condition identified in many reports from outside the United States.

APPROACH TO THE ENIGMATIC CASE

In approximately one-third of cases, the diagnosis is not known despite careful evaluation of CSF and potential extraneural sites of disease. A number of the organisms that cause chronic meningitis may take weeks to be identified by cultures. In enigmatic cases, several options are available, determined by the extent of the clinical deficits and rate of progression. It is prudent to wait until cultures are finalized if the patient is asymptomatic or symptoms are mild and not progressive. Unfortunately, in many cases progressive neurologic deterioration occurs, and rapid treatment is required. Ventricular-peritoneal shunts may be placed to relieve hydrocephalus, but the risk of disseminating the undiagnosed inflammatory process into the abdomen must be considered.

Empirical Treatment Diagnosis of the causative agent is essential because effective therapies exist for many etiologies of chronic meningitis, but if the condition is left untreated, progressive damage to the CNS and cranial nerves and roots is likely to occur. Occasionally, empirical therapy must be initiated when all attempts at diagnosis fail. In general, empirical therapy in the United States consists of antimycobacterial agents, amphotericin for fungal infection, or glucocorticoids for noninfectious inflammatory causes. It is important to direct empirical therapy of lymphocytic meningitis at tuberculosis, particularly if the condition is associated with low CSF glucose and sixth and other CN palsies, since untreated disease can be devastating within weeks. Patients on prolonged anti-tumor necrosis factor therapy who develop chronic meningitis also should be treated empirically with antituberculous therapy if the etiology is uncertain. In the Mayo Clinic series, the most useful empirical

therapy was administration of glucocorticoids rather than antituberculous therapy. Carcinomatous or lymphomatous meningitis may be difficult to diagnose initially, but the diagnosis becomes evident with time.

THE IMMUNOSUPPRESSED PATIENT

Chronic meningitis is not uncommon in the course of HIV infection. Pleocytosis and mild meningeal signs often occur at the onset of HIV infection, and occasionally low-grade meningitis persists. Toxoplasmosis commonly presents as intracranial abscesses and also may be associated with meningitis. Other important causes of chronic meningitis in AIDS include infection with *Cryptococcus*, *Nocardia*, *Candida*, or other fungi; syphilis; and lymphoma (Fig. 165-1). Toxoplasmosis, cryptococcosis, nocardiosis, and other fungal infections are important etiologic considerations in individuals with immunodeficiency states other than AIDS, including those due to immunosuppressive medications. Because of the increased risk of chronic meningitis and the attenuation of clinical signs of meningeal irritation in immunosuppressed individuals, CSF examination should be performed for any persistent headache or unexplained change in mental state.

ACKNOWLEDGMENT

Morton N. Swartz contributed to earlier editions of this chapter.

The skin is an essential component of immunity, protecting the host from potential pathogens in the environment. Breaches in this protective barrier thus represent a form of immunocompromise that predisposes the patient to infection. Thermal burns may cause massive destruction of the integument as well as derangements in humoral and cellular immunity, permitting the development of infection caused by environmental opportunists and components of the host's skin flora.

EPIDEMIOLOGY

Over the past decade, the estimated incidence of burn injuries in the United States has steadily declined; still, however, >1 million burn injuries are brought to medical attention each year. While many burn injuries are minor and require little or no intervention, 183,000 cases were reported between 2002 and 2011 to the National Burn Repository from specialized burn care facilities; of the 45,000 persons hospitalized for these injuries, 60% required intensive care and 20,000 had major burns involving at least 25% of the total body surface area. The majority of burn patients are men. Children under the age of 5 account for ~20% of all reported cases. Scalds, structural fires, and flammable liquids and gases are the major causes of burns, but electrical, chemical, and smoking-related sources also are important. Burns predispose to infection by damaging the protective barrier function of the skin, thus facilitating the entry of pathogenic microorganisms, and by inducing systemic immunosuppression. It is therefore not surprising that multiorgan failure and infectious complications are the major causes of morbidity and death in serious burn injury. More than 3000 patients in the United States die of burn-related infections each year, and 6 of the top 10 complications recently identified by the American Burn Association's 10-year review are infectious: pneumonia (4.6%), septicemia (2.7%), cellulitis/traumatic injury (2.6%), respiratory failure (2.5%), wound infection (2.2%), another infection (2.0%), renal failure (1.5%), line infection (1.4%), acute respiratory distress syndrome (1.2%), and arrhythmia (1.0%).

PATHOPHYSIOLOGY

Loss of the cutaneous barrier facilitates entry of the patient's own flora and of organisms from the hospital environment into the burn wound. Initially, the wound is colonized with gram-positive bacteria from the surrounding tissue, but the number of bacteria grows rapidly beneath the burn eschar, reaching $\sim 8.4 \times 10^3$ cfu/g on day 4 after the burn. The avascularity of the eschar, along with the impairment of local immune responses, favors further bacterial colonization and proliferation. By day 7, the wound is colonized with other microbes, including gram-positive bacteria, gram-negative bacteria, and yeasts derived from the gastrointestinal and upper respiratory flora. Invasive infection—localized and/or systemic—occurs when these bacteria penetrate viable tissue. In addition, a role for biofilms has been recognized in experimental animal models of burn-wound infection. (*Biofilms* are surface-associated communities of bacteria, often embedded in a matrix, that allow the microbes to persist and to resist the effects of host immunity and antimicrobial agents.)

Streptococci and staphylococci were the predominant causes of burn-wound infection in the preantibiotic era and remain important pathogens at present. With the advent of antimicrobial agents, *Pseudomonas aeruginosa* became a major problem in burn-wound management. In animal models of cutaneous thermal injury and wound infection with *Pseudomonas*, there is an early, steady increase of neutrophils in the skin and bacterial dissemination to lungs and spleen within 72 h. Less common anaerobic bacteria are typically found in infections of electrical burns or when open wound dressings are used. The widespread use of topical and more effective antimicrobial agents has resulted in a decline in bacterial wound infections

and the emergence of fungi (particularly *Candida albicans*, *Aspergillus* species, and the agents of mucormycosis) as increasingly important pathogens in burn-wound patients. Herpes simplex virus also has been found in burn wounds, especially those on the neck and face and those associated with inhalation injury. Cytomegalovirus viremia has been described in up to 71% of seropositive burn patients in prospective studies, and high levels (>1000 copies/mL) have been associated with increased duration of mechanical ventilation and longer stay in the intensive care unit (ICU).

Autopsy reports from patients with severe thermal burns over the last decade have identified *P. aeruginosa*, *Escherichia coli*, *Klebsiella pneumoniae*, and *Staphylococcus aureus* in association with mortality, independent of the percentage of total body surface area burned, the percentage of full-thickness burn, inhalation injury, and day of death after the burn. Indeed, burn trauma patients who acquire secondary *P. aeruginosa* infection have a fourfold greater mortality rate than those without *P. aeruginosa*. Historically, mortality rates among burn patients infected with *P. aeruginosa* have been as high as 77% over a 25-year period. In addition, *Acinetobacter calcoaceticus-baumannii* is among the top pathogens in some burn centers.

The cascade of events that follow a severe burn injury and that lead to multiorgan system failure and death is thought to represent a two-step process. The burn injury itself, with ensuing hypovolemia and tissue hypoxia, is followed by invasive infection arising from large amounts of devitalized tissue. The frequency of infection parallels the extent and severity of the burn injury. Severe burn injuries cause a state of immunosuppression that affects innate and adaptive immune responses. The substantial impact of immunocompromise on infection is due to effects on both the cellular and the humoral arms of the immune system. For example, decreases in the number and activity of circulating helper T cells, increases in suppressor T cells, decreases in production and release of monocytes and macrophages, and diminution in levels of immunoglobulin follow major burns. Neutrophil and complement functions also are impaired after burns. The increased levels of multiple cytokines detected in burn patients are compatible with the widely held belief that the inflammatory response becomes dysregulated in these individuals; bacterial cell products play a potent role in inducing proinflammatory mediators that contribute to this uncontrolled systemic inflammatory response. Increased permeability of the gut wall to bacteria and their components (e.g., endotoxin) also contributes to immune dysregulation and sepsis. Thus, the burn patient is predisposed to infection at remote sites (see below) as well as at the sites of burn injury. Another contributor to secondary immunosuppression after burn injuries is the endocrine system; increasing levels of vasopressin, aldosterone, cortisol, glucagon, growth hormone, catecholamines, and other hormones that directly affect lymphocyte proliferation, secretion of proinflammatory cytokines, natural killer cell activity, and suppressive T cells are seen.

CLINICAL MANIFESTATIONS AND DIAGNOSIS

Since clinical indications of wound infection are difficult to interpret, wounds must be monitored carefully for changes that may reflect infection. A margin of erythema frequently surrounds the sites of burns and by itself is not usually indicative of infection. Signs of infection include the conversion of a partial-thickness to a full-thickness burn, color changes (e.g., the appearance of a dark brown or black discoloration of the wound), the new appearance of erythema or violaceous edema in normal tissue at the wound margins, the sudden separation of the eschar from subcutaneous tissues, and the degeneration of the wound with the appearance of a new eschar.

Early surgical excision of devitalized tissue is now widely used, and burn-wound infections can be classified in relation to the excision site as (1) burn-wound impetigo (infection characterized by loss of epithelium from a previously re-epithelialized surface, as seen in a partial-thickness burn that is allowed to close by secondary intention, a grafted burn, or a healed skin donor site); (2) burn-related surgical-wound infection (purulent infection of excised burn and donor sites that have not yet epithelialized, accompanied by positive



FIGURE 166e-1 Cellulitis complicating a burn wound of the arm, with extension of the infection to adjacent healthy tissue. (Courtesy of Dr. Robert L. Sheridan, Massachusetts General Hospital, Boston; with permission.)

cultures); (3) burn-wound cellulitis (extension of infection to surrounding healthy tissue; Fig. 166e-1); and (4) invasive infection in unexcised burn wounds (infection that is secondary to a partial- or full-thickness burn wound and is manifested by separation of the eschar or by violaceous, dark brown, or black discoloration of the eschar; Fig. 166e-2). The appearance of a green discoloration of the wound or subcutaneous fat (Fig. 166e-3) or the development of ecthyma gangrenosum (see Fig. 25e-35) at a remote site points to a diagnosis of invasive *P. aeruginosa* infection.

Changes in body temperature, hypotension, tachycardia, altered mentation, neutropenia or neutrophilia, thrombocytopenia, and renal failure may result from invasive burn wounds and sepsis. However, because profound alterations in homeostasis occur as a consequence of burns per se and because inflammation without infection is a normal component of these injuries, the assessment of these changes is complicated. Alterations in body temperature, for example, are attributable to thermoregulatory dysfunction; tachycardia and hyperventilation accompany the metabolic changes induced by extensive burn injury and are not necessarily indicative of bacterial sepsis.

Given the difficulty of evaluating burn wounds solely on the basis of clinical observation and laboratory data, wound biopsies are necessary



FIGURE 166e-2 A severe upper-extremity burn infected with *Pseudomonas aeruginosa*. The wound requires additional debridement. Note the dark brown to black discoloration of the eschar. (Courtesy of Dr. Robert L. Sheridan, Massachusetts General Hospital, Boston; with permission.)

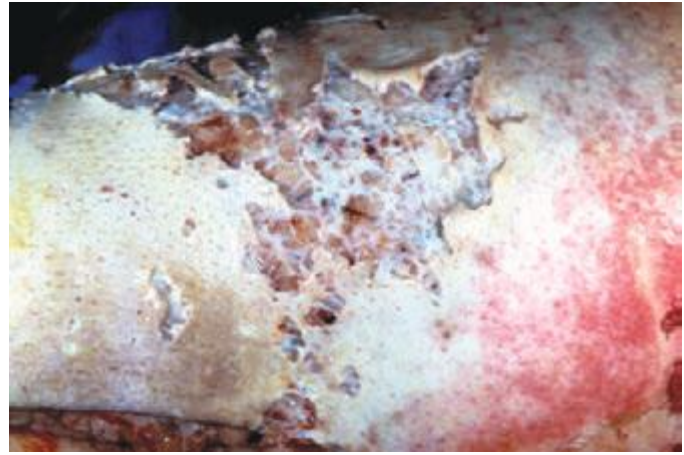


FIGURE 166e-3 A burn wound infected with *Pseudomonas aeruginosa*, with liquefaction of tissue. Note the green discoloration at the wound margins, which is suggestive of *Pseudomonas* infection. (Courtesy of Dr. Robert L. Sheridan, Massachusetts General Hospital, Boston; with permission.)

for definitive diagnosis of infection. The timing of these biopsies can be guided by clinical changes, but in some centers burn wounds are routinely biopsied at regular intervals. The biopsy specimen is examined for histologic evidence of bacterial invasion, and quantitative microbiologic cultures are performed. The presence of $>10^5$ viable bacteria per gram of tissue is highly suggestive of invasive infection and of a dramatically increased risk of sepsis. Histopathologic evidence of the invasion of viable tissue and the presence of microorganisms in unburned blood vessels and lymphatics is a more definitive indicator of infection. A blood culture positive for the same organism seen in large quantities in biopsied tissue is a reliable indicator of burn sepsis. Surface cultures may provide some indication of the microorganisms present in the hospital environment but are not indicative of the etiology of infection. This noninvasive technique may be of use in determining the flora present in excised burn areas or in areas where the skin is too thin for biopsy (e.g., over the ears, eyes, or digits). Rapid identification of organisms and institution of appropriate therapy are critical to the survival of patients with severe burn injury; polymerase chain reaction (PCR) is now being used for rapid identification of specific pathogens, sometimes in <6 h, to allow earlier treatment interventions.

In addition to infection of the burn wound itself, a number of other infections due to the immunosuppression caused by extensive burns and the manipulations necessary for clinical care put burn patients at risk. Pneumonia, now the most common infectious complication among hospitalized burn patients, is most often acquired nosocomially via the respiratory route. The incidence of ventilator-associated pneumonia among burn patients is 22–30 cases per 1000 ventilator-days—more than double that among surgical or medical ICU cohorts; this infection usually results from colonization of the lower respiratory tract and parenchyma because of sustained microaspiration. Among the risk factors associated with secondary pneumonia are inhalation injury, intubation, full-thickness chest wall burns, cutaneous thermal injuries, immobility, blood transfusions, and uncontrolled wound sepsis with hematogenous spread. Septic pulmonary emboli also may occur. Suppurative thrombophlebitis may complicate the vascular catheterization necessary for fluid and nutritional support in burns. Endocarditis, urinary tract infection, bacterial chondritis (particularly in patients with burned ears), and intraabdominal infection also complicate serious burn injury. Staphylococcal scalded skin syndrome due to burn-wound infection with *S. aureus* has been described as a rare complication. Finally, burn surgical-wound infections contribute to morbidity and have been found in up to 39% of patients; these infections often result in repeat skin grafting and prolonged hospitalization.

The ultimate goal of burn-wound management is closure and healing of the wound. Early surgical excision of burned tissue, with extensive debridement of necrotic tissue and grafting of skin or skin substitutes, greatly decreases mortality rates associated with severe burns. In addition, the four widely used topical antimicrobial agents—silver sulfadiazine cream, mafenide acetate cream, silver nitrate cream, and nanocrystalline silver dressings—dramatically decrease the bacterial burden of burn wounds and reduce the incidence of burn-wound infection; these agents are routinely applied to partial- and full-thickness burns. The bactericidal properties of silver are related to its effect on respiratory enzymes on bacterial cell walls; its interaction with structural proteins causes keratinocyte and fibroblast toxicity that can delay wound healing if silver-based compounds are used indiscriminately. All four agents are broadly active against many bacteria and some fungi and are useful before bacterial colonization is established. Silver sulfadiazine is often used initially, but its value can be limited by bacterial resistance, poor wound penetration, or toxicity (leukopenia). Mafenide acetate has broader activity against gram-negative bacteria. The cream penetrates eschars and thus can prevent or treat infection beneath them; its use without dressings allows regular examination of the wound area. The foremost disadvantages of mafenide acetate are that it can inhibit carbonic anhydrase, resulting in metabolic acidosis, and that it elicits hypersensitivity reactions in up to 7% of patients. This agent is most often used when gram-negative bacteria invade the burn wound and when treatment with silver sulfadiazine fails. The activity of mafenide acetate against gram-positive bacteria is limited. Nanocrystalline silver dressings provide broader antimicrobial coverage than any other available topical preparation, exhibiting activity against methicillin-resistant *S. aureus* (MRSA) and vancomycin-resistant enterococci (VRE), moderate ability to penetrate eschars, and limited toxicity. In addition, this approach provides controlled and prolonged release of nanocrystalline silver into the wound, limiting the number of dressing changes and therefore reducing the risk of nosocomial infections as well as the cost of treatment. Mupirocin, a topical antimicrobial agent used to eradicate nasal colonization with MRSA, is increasingly being used in burn units where MRSA is prevalent. The efficacy of mupirocin in reducing burn-wound bacterial counts and preventing systemic infections is comparable to that of silver sulfadiazine.

In recent years, rates of fungal infection have increased in burn patients. When superficial fungal infection occurs, nystatin may be mixed with silver sulfadiazine or mafenide acetate as topical therapy. A small study found that nystatin powder (6 million units/g) was effective for treatment of superficial and deep burn-wound infections caused by *Aspergillus* or *Fusarium* species. In addition to these products, moisture-retention ointments with antimicrobial properties can promote rapid autolysis, debridement, and moist healing of partial-thickness wounds.

When invasive wound infection is diagnosed, topical therapy should be changed to mafenide acetate. *Subeschar clysis* (the direct instillation of an antibiotic, often piperacillin, into wound tissues under the eschar) is a useful adjunct to surgical and systemic antimicrobial therapy. Systemic treatment with antibiotics active against the pathogens present in the wound should be instituted. In the absence of culture data, treatment should be broad in spectrum, covering organisms commonly encountered in that particular burn unit. Such coverage is usually achieved with an antibiotic active

against gram-positive pathogens (e.g., oxacillin, 2 g IV every 4 h) and with a drug active against *P. aeruginosa* and other gram-negative rods (e.g., mezlocillin, 3 g IV every 4 h; gentamicin, 5 mg/kg IV per day). In penicillin-allergic patients, vancomycin (1 g IV every 12 h) may be substituted for oxacillin (and is efficacious against MRSA), and ciprofloxacin (400 mg IV every 12 h) may be substituted for mezlocillin. Oxazolidinone antibiotics like linezolid have demonstrated efficacy in reducing bacterial growth and toxic shock syndrome toxin 1 levels in animal models of MRSA burn-wound infections.

Patients with burn wounds frequently have alterations in metabolism and renal clearance mechanisms that mandate the monitoring of serum antibiotic levels. The levels achieved with standard doses are often subtherapeutic.

Treatment of infections caused by emerging resistant pathogens remains a challenge in the care of burn patients. MRSA, resistant enterococci, multidrug-resistant gram-negative rods, and Enterobacteriaceae producing extended-spectrum β -lactamases have been associated with burn-wound infections and identified in burn-unit outbreaks. Strict infection-control practices (including microbiologic surveillance in burn units) and appropriate antimicrobial therapy remain important measures in reducing rates of infection due to resistant organisms.

In general, prophylactic systemic antibiotics have no role in the management of burn wounds and can, in fact, lead to colonization with resistant microorganisms. In some studies, antibiotic prophylaxis has been associated with increases in secondary infections of the upper and lower respiratory tract and the urinary tract as well as with prolonged hospitalization. An exception involves cases requiring burn-wound manipulation. Since procedures such as debridement, excision, or grafting frequently result in bacteremia, prophylactic systemic antibiotics are administered at the time of wound manipulation; the specific agents used should be chosen on the basis of data obtained by wound culture or data on the hospital's resident flora.

The use of oral antibiotics for selective digestive decontamination (SDD) to decrease bacterial colonization and the risk of burn-wound infection is controversial and has not been widely adopted. In a randomized, double-blind, placebo-controlled trial in patients with burns involving >20% of the total body surface area, SDD was associated with reduced mortality rates in the burn ICU and in the hospital and also with a reduced incidence of pneumonia. The effects of SDD on the normal anaerobic bowel flora must be taken into consideration before this approach is used.

Strategies to reduce or limit systemic spread of wound infections, particularly to the lung, may be useful adjuncts to therapy. Some of these strategies are aimed at reducing neutrophilic inflammation at the site of injury, which can accelerate biofilm formation, particularly by *P. aeruginosa*. For example, in animal models of cutaneous burns with *P. aeruginosa* wound inoculation, a single dose of azithromycin administered early reduces rates of *Pseudomonas* infection and systemic spread to lung and spleen and appears to have effects similar to those of classic anti-*Pseudomonas* agents, such as tobramycin. The extent to which azithromycin can be administered early in humans to prevent dissemination remains to be studied.

All burn-injury patients should undergo tetanus booster immunization if they have completed primary immunization but have not received a booster dose in the past 5 years. Patients without prior immunization should receive tetanus immune globulin and undergo primary immunization.

167e Infectious Complications of Bites

Lawrence C. Madoff, Florencia Pereyra

The skin is an essential component of nonspecific immunity, protecting the host from potential pathogens in the environment. Breaches in this protective barrier thus represent a form of immunocompromise that predisposes the patient to infection. Bites and scratches from animals and humans allow the inoculation of microorganisms past the skin's protective barrier into deeper, susceptible host tissues.

Each year in the United States, millions of animal-bite wounds are sustained. The vast majority are inflicted by pet dogs and cats, which number >100 million; the annual incidence of dog and cat bites has been reported as 300 bites per 100,000 population. Other bite wounds are a consequence of encounters with animals in the wild or in occupational settings. While many of these wounds require minimal or no therapy, a significant number result in infection, which may be life-threatening. The microbiology of bite-wound infections in general reflects the oropharyngeal flora of the biting animal, although organisms from the soil, the skin of the animal and the victim, and the animal's feces may also be involved.

DOG BITES

In the United States, dogs bite >4.7 million people each year and are responsible for 80% of all animal-bite wounds, an estimated 15–20% of which become infected. Each year, 800,000 Americans seek medical attention for dog bites; of those injured, 386,000 require treatment in an emergency department, with >1000 emergency department visits each day and about a dozen deaths per year. Most dog bites are provoked and are inflicted by the victim's pet or by a dog known to the victim. These bites are frequently sustained during efforts to break up a dogfight. Children are more likely than adults to sustain canine bites, with the highest incidence of 6 bites/1000 population among boys 5–9 years old. Victims are more often male than female, and bites most often involve an upper extremity. Among children <4 years old, two-thirds of all these injuries involve the head or neck. Infection typically manifests 8–24 h after the bite as pain at the site of injury with cellulitis accompanied by purulent, sometimes foul-smelling discharge. Septic arthritis and osteomyelitis may develop if a canine tooth penetrates synovium or bone. Systemic manifestations (e.g., fever, lymphadenopathy, and lymphangitis) also may occur. The microbiology of dog-bite wound infections is usually mixed and includes β -hemolytic streptococci, *Pasteurella* species, *Staphylococcus* species (including methicillin-resistant *Staphylococcus aureus* [MRSA]), *Eikenella corrodens*, and *Capnocytophaga canimorsus*. Many wounds also include anaerobic bacteria such as *Actinomyces*, *Fusobacterium*, *Prevotella*, and *Porphyromonas* species.

While most infections resulting from dog-bite injuries are localized to the area of injury, many of the microorganisms involved are capable of causing systemic infection, including bacteremia, meningitis, brain abscess, endocarditis, and chorioamnionitis. These infections are particularly likely in hosts with edema or compromised lymphatic drainage in the involved extremity (e.g., after a bite on the arm in a woman who has undergone mastectomy) and in patients who are immunocompromised by medication or disease (e.g., glucocorticoid use, systemic lupus erythematosus, acute leukemia, or hepatic cirrhosis). In addition, dog bites and scratches may result in systemic illnesses such as rabies (Chap. 232) and tetanus (Chap. 177).

Infection with *C. canimorsus* following dog-bite wounds may result in fulminant sepsis, disseminated intravascular coagulation, and renal failure, particularly in hosts who have impaired hepatic function, who have undergone splenectomy, or who are immunosuppressed. This thin gram-negative rod is difficult to culture on most solid media but grows in a variety of liquid media. The bacteria are occasionally seen within polymorphonuclear leukocytes on Wright-stained smears of

peripheral blood from septic patients. Tularemia (Chap. 195) also has been reported to follow dog bites.


CAT BITES

Although less common than dog bites, cat bites and scratches result in infection in more than half of all cases. Because the cat's narrow, sharp canine teeth penetrate deeply into tissue, cat bites are more likely than dog bites to cause septic arthritis and osteomyelitis; the development of these conditions is particularly likely when punctures are located over or near a joint, especially in the hand. Women sustain cat bites more frequently than do men. These bites most often involve the hands and arms. Both bites and scratches from cats are prone to infection from organisms in the cat's oropharynx. *Pasteurella multocida*, a normal component of the feline oral flora, is a small gram-negative coccobacillus implicated in the majority of cat-bite wound infections. Like that of dog-bite wound infections, however, the microflora of cat-bite wound infections is usually mixed. Other microorganisms causing infection after cat bites are similar to those causing dog-bite wound infections.

The same risk factors for systemic infection following dog-bite wounds apply to cat-bite wounds. *Pasteurella* infections tend to advance rapidly, often within hours, causing severe inflammation accompanied by purulent drainage; *Pasteurella* may also be spread by respiratory droplets from animals, resulting in pneumonia or bacteremia. Like dog-bite wounds, cat-bite wounds may result in the transmission of rabies or in the development of tetanus. Infection with *Bartonella henselae* causes cat-scratch disease (Chap. 197) and is an important late consequence of cat bites and scratches. Tularemia (Chap. 195) also has been reported to follow cat bites.

OTHER ANIMAL BITES

Infections have been attributed to bites from many animal species. Often these bites are sustained as a consequence of occupational exposure (farmers, laboratory workers, veterinarians) or recreational exposure (hunters and trappers, wilderness campers, owners of exotic pets). Generally, the microflora of bite wounds reflects the oral flora of the biting animal. Most members of the cat family, including feral cats, harbor *P. multocida*. Bite wounds from aquatic animals such as alligators or piranhas may contain *Aeromonas hydrophila*. Venomous snakebites (Chap. 474) result in severe inflammatory responses and tissue necrosis—conditions that render these injuries prone to infection. The snake's oral flora includes many species of aerobes and anaerobes, such as *Pseudomonas aeruginosa*, *Serratia marcescens*, *Proteus* species, *Staphylococcus epidermidis*, *Bacteroides fragilis*, and *Clostridium* species. Bites from nonhuman primates are highly susceptible to infection with pathogens similar to those isolated from human bites (see below). Bites from Old World monkeys (*Macaca*) may also result in the transmission of B virus (*Herpesvirus simiae*, cercopithecine herpesvirus), a cause of serious infection of the human central nervous system. Bites of seals, walruses, and polar bears may cause a chronic suppurative infection known as *seal finger*, which is probably due to one or more species of *Mycoplasma* colonizing these animals.

 Small rodents, including rats, mice, and gerbils, as well as animals that prey on rodents may transmit *Streptobacillus moniliformis* (a microaerophilic, pleomorphic gram-negative rod) or *Spirillum minor* (a spirochete); these organisms cause a clinical illness known as *rat-bite fever*. The vast majority of cases in the United States are streptobacillary, whereas *Spirillum* infection occurs mainly in Asia.

In the United States, the risk of rodent bites is usually greatest among laboratory workers or inhabitants of rodent-infested dwellings (particularly children). Rat-bite fever is distinguished from acute bite-wound infection by its typical manifestation after the initial wound has healed. Streptobacillary disease follows an incubation period of 3–10 days. Fever, chills, myalgias, headache, and severe migratory arthralgias are usually followed by a maculopapular rash, which characteristically involves the palms and soles and may become confluent or purpuric. Complications include endocarditis, myocarditis, meningitis, pneumonia, and abscesses in many organs. *Haverhill fever* is an

167e-2 *S. moniliformis* infection acquired from contaminated milk or drinking water and has similar manifestations. Streptobacillary rat-bite fever was frequently fatal in the preantibiotic era. The differential diagnosis includes Rocky Mountain spotted fever, Lyme disease, leptospirosis, and secondary syphilis. The diagnosis is made by direct observation of the causative organisms in tissue or blood, by culture of the organisms on enriched media, or by serologic testing with specific agglutinins.

Spirillum infection (referred to in Japan as *sodoku*) causes pain and purple swelling at the site of the initial bite, with associated lymphangitis and regional lymphadenopathy, after an incubation period of 1–4 weeks. The systemic illness includes fever, chills, and headache. The original lesion may eventually progress to an eschar. The infection is diagnosed by direct visualization of the spirochetes in blood or tissue or by animal inoculation.

Finally, NO-1 (CDC nonoxidizer group 1) is a bacterium associated with dog- and cat-bite wounds. Infections in which NO-1 has been isolated have tended to manifest locally (i.e., as abscess and cellulitis). These infections have occurred in healthy persons with no underlying illness and in some instances have progressed from localized to systemic illnesses. The phenotypic characteristics of NO-1 are similar to those of asaccharolytic *Acinetobacter* species; i.e., NO-1 is oxidase-, indole-, and urease-negative. To date, all strains identified have been shown to be susceptible to aminoglycosides, β -lactam antibiotics, tetracyclines, quinolones, and sulfonamides.

HUMAN BITES

Human bites may be self-inflicted; may be sustained by medical personnel caring for patients; or may take place during fights, domestic abuse, or sexual activity. Human-bite wounds become infected more frequently (~10–15% of the time) than do bites inflicted by other animals. These infections reflect the diverse oral microflora of humans, which includes multiple species of aerobic and anaerobic bacteria. Common aerobic isolates include viridans streptococci, *S. aureus*, *E. corrodens* (which is particularly common in clenched-fist injury; see below), and *Haemophilus influenzae*. Anaerobic species, including *Fusobacterium nucleatum* and *Prevotella*, *Porphyromonas*, and *Peptostreptococcus* species, are isolated from 50% of wound infections due to human bites; many of these isolates produce β -lactamases. The oral flora of hospitalized and debilitated patients often includes Enterobacteriaceae in addition to the usual organisms. Hepatitis B, hepatitis C, herpes simplex virus infection, syphilis, tuberculosis, actinomycosis, and tetanus have been reported to be transmitted by human bites; it is biologically possible to transmit HIV through human bites, although this event is quite unlikely.

Human bites are categorized as either *occlusional* injuries, which are inflicted by actual biting, or *clenched-fist* injuries, which are sustained when the fist of one individual strikes the teeth of another, causing traumatic laceration of the hand. For several reasons, clenched-fist injuries, which are sometimes referred to as “fight bite” and which are more common than occlusional injuries, result in particularly serious infections. The deep spaces of the hand, including the bones, joints, and tendons, are frequently inoculated with organisms in the course of such injuries. The clenched position of the fist during injury, followed by extension of the hand, may further promote the introduction of bacteria as contaminated tendons retract beneath the skin’s surface. Moreover, medical attention is often sought only after frank infection develops.

APPROACH TO THE PATIENT: Animal or Human Bites

A careful history should be elicited, including the type of biting animal, the type of attack (provoked or unprovoked), and the amount of time elapsed since injury. Local and regional public-health authorities should be contacted to determine whether an individual species could be rabid and/or to locate and observe the biting animal when rabies prophylaxis may be indicated ([Chap. 232](#)).

Suspicious human-bite wounds should provoke careful questioning regarding domestic or child abuse. Details on antibiotic allergies, immunosuppression, splenectomy, liver disease, mastectomy, and immunization history should be obtained. The wound should be inspected carefully for evidence of infection, including redness, exudate, and foul odor. The type of wound (puncture, laceration, or scratch); the depth of penetration; and the possible involvement of joints, tendons, nerves, and bones should be assessed. It is often useful to include a diagram or photograph of the wound in the medical record. In addition, a general physical examination should be conducted and should include an assessment of vital signs as well as an evaluation for evidence of lymphangitis, lymphadenopathy, dermatologic lesions, and functional limitations. Injuries to the hand warrant consultation with a hand surgeon for the assessment of tendon, nerve, and muscular damage. Radiographs should be obtained when bone may have been penetrated or a tooth fragment may be present. Culture and Gram’s staining of all infected wounds are essential; anaerobic cultures should be undertaken if abscesses, devitalized tissue, or foul-smelling exudate is present. A small-tipped swab may be used to culture deep punctures or small lacerations. It is also reasonable to culture samples from apparently uninfected wounds due to bites inflicted by animals other than dogs and cats, since the microorganisms causing disease are less predictable in these cases. The white blood cell count should be determined and the blood cultured if systemic infection is suspected.

TREATMENT BITE-WOUND INFECTIONS

WOUND MANAGEMENT

Wound closure is controversial in bite injuries. Many authorities prefer not to attempt primary closure of wounds that are or may become infected, choosing instead to irrigate these wounds copiously, debride devitalized tissue, remove foreign bodies, and approximate the wound edges. Delayed primary closure may be undertaken after the risk of infection is over. Small uninfected wounds may be allowed to close by secondary intention. Puncture wounds due to cat bites should be left unsutured because of the high rate at which they become infected. Facial wounds are usually sutured after thorough cleaning and irrigation because of the importance of a good cosmetic result in this area and because anatomic factors such as an excellent blood supply and the absence of dependent edema lessen the risk of infection.

ANTIBIOTIC THERAPY

Established Infection Antibiotics should be administered for all established bite-wound infections and should be chosen in light of the most likely potential pathogens, as indicated by the biting species and by Gram’s stain and culture results ([Table 167e-1](#)). For dog and cat bites, antibiotics should be effective against *S. aureus*, *Pasteurella* species, *C. canimorsus*, streptococci, and oral anaerobes. For human bites, agents with activity against *S. aureus*, *H. influenzae*, and β -lactamase-positive oral anaerobes should be used. The combination of an extended-spectrum penicillin with a β -lactamase inhibitor (amoxicillin/clavulanic acid, ticarcillin/clavulanic acid, ampicillin/sulbactam) appears to offer the most reliable coverage for these pathogens. Second-generation cephalosporins (cefuroxime, cefoxitin) also offer substantial coverage. The choice of antibiotics for penicillin-allergic patients (particularly those in whom immediate-type hypersensitivity makes the use of cephalosporins hazardous) is more difficult and is based primarily on in vitro sensitivity since data on clinical efficacy are inadequate. The combination of an antibiotic active against gram-positive cocci and anaerobes (such as clindamycin) with trimethoprim-sulfamethoxazole or a fluoroquinolone, which is active against many of the other potential pathogens, would appear reasonable. In vitro data suggest that azithromycin alone provides coverage against most commonly isolated bite-wound pathogens. As MRSA becomes

TABLE 167e-1 MANAGEMENT OF WOUND INFECTIONS FOLLOWING ANIMAL AND HUMAN BITES

Biting Species	Commonly Isolated Pathogens	Preferred Antibiotic(s) ^a	Alternative in Penicillin-Allergic Patient	Prophylaxis Advised for Early Uninfected Wounds	Other Considerations
Dog	<i>Staphylococcus aureus</i> , <i>Pasteurella multocida</i> , anaerobes, <i>Capnocytophaga canimorsus</i>	Amoxicillin/clavulanate (250–500 mg PO tid) or ampicillin/sulbactam (1.5–3.0 g IV q6h)	Clindamycin (150–300 mg PO qid) plus either TMP-SMX (1 DS tablet PO bid) or ciprofloxacin (500 mg PO bid)	Sometimes ^b	Consider rabies prophylaxis.
Cat	<i>P. multocida</i> , <i>S. aureus</i> , anaerobes	Amoxicillin/clavulanate or ampicillin/sulbactam as above	Clindamycin plus TMP-SMX as above or a fluoroquinolone	Usually	Consider rabies prophylaxis. Carefully evaluate for joint/bone penetration.
Human, occlusional	Viridans streptococci, <i>S. aureus</i> , <i>Haemophilus influenzae</i> , anaerobes	Amoxicillin/clavulanate or ampicillin/sulbactam as above	Erythromycin (500 mg PO qid) or a fluoroquinolone	Always	
Human, clenched-fist	As for occlusional, plus <i>Eikenella corrodens</i>	Ampicillin/sulbactam as above or imipenem (500 mg q6h)	Cefoxitin ^c	Always	Examine for tendon, nerve, or joint involvement.
Monkey	As for human bite	As for human bite	As for human bite	Always	For macaque monkeys, consider B virus prophylaxis with acyclovir.
Snake	<i>Pseudomonas aeruginosa</i> , <i>Proteus</i> spp., <i>Bacteroides fragilis</i> , <i>Clostridium</i> spp.	Ampicillin/sulbactam as above	Clindamycin plus TMP-SMX as above or a fluoroquinolone	Sometimes, especially with venomous snakes	Administer antivenin for venomous snakebite.
Rodent	<i>Streptobacillus moniliformis</i> , <i>Leptospira</i> spp., <i>P. multocida</i>	Penicillin VK (500 mg PO qid)	Doxycycline (100 mg PO bid)	Sometimes	—

^aAntibiotic choices should be based on culture data when available. These suggestions for empirical therapy need to be tailored to individual circumstances and local conditions. IV regimens should be used for hospitalized patients. A single IV dose of antibiotics may be given to patients who will be discharged after initial management. ^bProphylactic antibiotics are suggested for severe or extensive wounds, facial wounds, and crush injuries; when bone or joint may be involved; and when comorbidity is present (see text). ^cMay be hazardous in patients with immediate-type hypersensitivity to penicillin.

Abbreviations: DS, double-strength; TMP-SMX, trimethoprim-sulfamethoxazole.

more common in the community and evidence of its transmission between humans and their animal contacts increases, empirical use of agents active against MRSA should be considered in high-risk situations while culture results are awaited.

Antibiotics are generally given for 10–14 days, but the response to therapy must be carefully monitored. Failure to respond should prompt a consideration of diagnostic alternatives and surgical evaluation for possible drainage or debridement. Complications such as osteomyelitis or septic arthritis mandate a longer duration of therapy.

Management of *C. canimorsus* sepsis requires a 2-week course of IV penicillin G (2 million units IV every 4 h) and supportive measures. Alternative agents for the treatment of *C. canimorsus* infection include cephalosporins and fluoroquinolones. Serious infection with *P. multocida* (e.g., pneumonia, sepsis, or meningitis) also should be treated with IV penicillin G. Alternative agents include second- or third-generation cephalosporins or ciprofloxacin.

Bites by venomous snakes (**Chap. 474**) may not require antibiotic treatment. Because it is often difficult to distinguish signs of infection from tissue damage caused by the envenomation, many authorities continue to recommend treatment directed against the snake's oral flora—i.e., the administration of broadly active agents such as ceftriaxone (1–2 g IV every 12–24 h) or ampicillin/sulbactam (1.5–3.0 g IV every 6 h).

Seal finger appears to respond to doxycycline (100 mg twice daily for a duration guided by the response to therapy).

Presumptive or Prophylactic Therapy The use of antibiotics for patients presenting early (within 8 h) after bite injury is controversial. Although symptomatic infection frequently will not yet have manifested at this point, many early wounds will harbor pathogens, and many will become infected. Studies of antibiotic prophylaxis for wound infections are limited and have often included only small numbers of cases in which various types of wounds have been managed according to various protocols. A meta-analysis of eight

randomized trials of prophylactic antibiotics in patients with dog-bite wounds demonstrated a reduction in the rate of infection by 50% with prophylaxis. However, in the absence of sound clinical trials, many clinicians base the decision to treat bite wounds with empirical antibiotics on the species of the biting animal; the location, severity, and extent of the bite wound; and the existence of comorbid conditions in the host. All human- and monkey-bite wounds should be treated presumptively because of the high rate of infection. Most cat-bite wounds, particularly those involving the hand, should be treated. Other factors favoring treatment for bite wounds include severe injury, as in crush wounds; potential bone or joint involvement; involvement of the hands or genital region; host immunocompromise, including that due to liver disease or splenectomy; and prior mastectomy on the side of an involved upper extremity. When prophylactic antibiotics are administered, they are usually given for 3–5 days.

Rabies and Tetanus Prophylaxis Rabies prophylaxis, consisting of both passive administration of rabies immune globulin (with as much of the dose as possible infiltrated into and around the wound) and active immunization with rabies vaccine, should be given in consultation with local and regional public health authorities for some animal bites and scratches as well as for certain nonbite exposures (**Chap. 232**). Rabies is endemic in a variety of animals, including dogs and cats in many areas of the world. Many local health authorities require the reporting of all animal bites. A tetanus booster immunization should be given if the patient has undergone primary immunization but has not received a booster dose in the past 5 years. Patients who have not previously completed primary immunization should be immunized and should also receive tetanus immune globulin. Elevation of the site of injury is an important adjunct to antimicrobial therapy. Immobilization of the infected area, especially the hand, also is beneficial.

SECTION 3

CLINICAL SYNDROMES: HEALTH CARE–ASSOCIATED INFECTIONS

168 Infections Acquired in Health Care Facilities

Robert A. Weinstein

The costs of hospital-acquired (nosocomial) and other health care-associated infections are great. These infections have affected as many as 1.7 million patients at a cost of ~\$28–33 billion and 99,000 lives in U.S. hospitals annually. Although efforts to lower infection risks have been challenged by the numbers of immunocompromised patients, antibiotic-resistant bacteria, fungal and viral superinfections, and invasive devices and procedures, a prevailing viewpoint—often termed “zero tolerance”—is that almost all health care-associated infections should be avoidable with strict application of evidence-based prevention guidelines ([Table 168-1](#)). In fact, rates of device-related infections—historically, the largest drivers of risk—have fallen steadily

over the past few years. Unfortunately, at the same time, antimicrobial-resistant pathogens have risen in number and are estimated to contribute to ~23,000 deaths in and outside of hospitals annually. This chapter reviews health care-associated and device-related infections as well as basic surveillance, prevention, control, and treatment activities.

ORGANIZATION, RESPONSIBILITIES, AND INCREASING SCRUTINY OF HEALTH CARE–ASSOCIATED INFECTION PROGRAMS

The standards of the Joint Commission require all accredited hospitals to have active programs for surveillance, prevention, and control of nosocomial infections. Education of physicians in infection control and health care epidemiology is required in infectious disease fellowship programs and is available in online courses. Concerns over “patient safety” have led to federal legislation that prevents U.S. hospitals from upgrading Medicare charges to pay for hospital costs resulting from at least 14 specific nosocomial events ([Table 168-2](#)) and have prompted national efforts to publicly

TABLE 168-1 SOURCES OF INFECTION CONTROL GUIDELINES AND OVERSIGHT

Organization	Role	Major Constituents	Website
Joint Commission	Regulatory	Hospitals, long-term-care facilities, laboratories	www.jointcommission.org
CAP	Regulatory	Laboratories	www.cap.org
OSHA	Regulatory	Workers	www.osha.gov
CMS	Regulatory	Medicare/Medicaid providers	www.cms.hhs.gov
PQRI	Regulatory and advisory	Eligible professionals	www.cms.hhs.gov/pqri/
HHS Action Plan	Regulatory and advisory	Health care and infection prevention personnel	www.hhs.gov/ash/initiatives/hai/actionplan/
CDC			
DHQP	Advisory	Health care facilities and personnel	www.cdc.gov/nceid/dhqp/
HICPAC	Advisory	Health care facilities and personnel	www.cdc.gov/hicpac/
NIOSH	Advisory	Workers	www.cdc.gov/niosh/
AHRQ	Advisory	Broad (e.g., health care personnel)	www.ahrq.gov
NQF	Advisory	Broad (e.g., health care personnel)	www.qualityforum.org
IOM	Advisory	Broad (e.g., health care personnel)	www.iom.edu
Federal Influenza Planning	Advisory	Health care and public health personnel	www.flu.gov/planning-preparedness/hospital#
Trust for America's Health	Advisory	Broad (e.g., the public)	healthyamericans.org
CSTE	Advisory and professional society	Public health personnel	www.cste.org
IDSA	Professional society	Infectious disease physicians/researchers	www.idsociety.org
SHEA	Professional society	Health care epidemiologists	www.shea-online.org
HIS	Professional society	Health care epidemiologists	www.his.org.uk
APIC	Professional society	Infection preventionists	www.apic.org
MedQIC	Quality improvement	Broad (e.g., health care personnel)	www.qualitynet.org
IHI	Quality improvement	Broad (e.g., health care personnel)	www.ihl.org
Leapfrog Group	Quality improvement	Broad (payers, consumers, employers, and health care personnel)	www.leapfroggroup.org
NSQIP	Quality improvement	Surgery services	www.acsnsqip.org

Abbreviations: AHRQ, Agency for Healthcare Research and Quality; APIC, Association for Professionals in Infection Control and Epidemiology; CAP, College of American Pathologists; CDC, Centers for Disease Control and Prevention; CMS, Centers for Medicare & Medicaid Services; CSTE, Council of State and Territorial Epidemiologists; DHQP, Division of Healthcare Quality Promotion; HHS, Health and Human Services; HICPAC, Healthcare Infection Control Practices Advisory Committee; HIS, Hospital Infection Society; IDSA, Infectious Diseases Society of America; IHI, Institute for Healthcare Improvement; IOM, Institute of Medicine; MedQIC, Medicare Quality Improvement Community; NIOSH, National Institute for Occupational Safety and Health; NQF, National Quality Forum; NSQIP, National Surgical Quality Improvement Program; OSHA, Occupational Safety & Health Administration; PQRI, Physician Quality Reporting Initiative; SHEA, Society for Healthcare Epidemiology of America.

report on processes of patient care (e.g., timely administration and appropriateness of perioperative antibiotic prophylaxis) and patient outcomes (e.g., surgical wound infection rates). Neither the carrot (pay-for-performance) nor the stick (nonpayment for preventable infections) appears to have impacted infection rates. The effect of public attention may be more positive; in 2009, the U.S. Department

of Health and Human Services released a major interagency Action Plan to Prevent Healthcare-Associated Infections, including a list of 5-year national prevention targets that are mostly on track (Table 168-3).

SURVEILLANCE

Traditionally, infection preventionists have surveyed inpatients for infections acquired in hospitals (defined as those neither present nor incubating at the time of admission). Surveillance most often requires review of microbiology laboratory results, “shoe-leather” epidemiology on nursing wards, and application of standardized definitions of infection. Progressively more infection-control programs use computerized hospital databases for algorithm-driven electronic surveillance (e.g., of vascular catheter and surgical wound infections) that removes observer bias and, by so doing, provides data that are more reliable for interfacility comparisons. Although infection surveillance in nursing homes and some long-term acute-care hospitals (LTACHs) is still in its formative stage, the role of these facilities in the transmission of antimicrobial-resistant pathogens will require their increased attention to infection surveillance and control.

Most hospitals aim surveillance at infections associated with high-level morbidity or expense. Quality-improvement activities in infection control have led to increased surveillance of personnel compliance with infection control policies (e.g., adherence to influenza vaccination recommendations). In the spirit of “what is measured improves,” the majority of states now require public reporting of processes for prevention of health care-associated infection and/or patient outcomes. As a result, in some locales, the surveillance pendulum is swinging

TABLE 168-2 HEALTH CARE–ACQUIRED CONDITIONS NOT ELIGIBLE FOR ADDITIONAL FEDERAL PAYMENT*

Foreign objects retained after surgery
Air embolism
Blood incompatibilities
Decubitus ulcers (stages III and IV)
Fractures/other injuries from falls or trauma
Catheter-associated urinary tract infections
Vascular catheter-associated infections
Manifestations of poor glycemic control
Surgical-site infection or mediastinitis following coronary artery bypass graft
Surgical-site infection following certain orthopedic procedures
Surgical-site infection following bariatric surgery for obesity
Surgical-site infection following cardiac electronic device implantation
Venous thromboembolism (after hip or knee replacement)
Iatrogenic pneumothorax with venous catheterization

*Based on the U.S. Federal Deficit Reduction Act of 2005. As of October 2012, Medicare stopped paying additional money to hospitals for these 14 health care–acquired conditions. See www.cms.gov/HospitalAcqCond/ (last accessed November 13, 2014).

TABLE 168-3 SUMMARY OF PROGRESS TOWARD THE NINE NATIONAL TARGETS FOR ELIMINATION OF HEALTH CARE–ASSOCIATED INFECTIONS, U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES: MIDPOINT EVALUATION

Metric	Source	National 5-Year Prevention Target	On Track to Meet 2013 Targets?
Bloodstream infections	NHSN	50% reduction	Yes ^a
Adherence to central-line insertion practices	NHSN	100% adherence	Retired
<i>Clostridium difficile</i> diagnosed in hospital	HCUP	30% reduction	No
<i>C. difficile</i> infections	NHSN	30% reduction	No ^a
Urinary tract infections	NHSN	25% reduction	Yes ^a
MRSA invasive infections (population)	EIP	50% reduction	Yes ^a
MRSA bacteremia (hospital)	NHSN	25% reduction	Yes
Surgical-site infections	NHSN	25% reduction	Yes
Surgical Care Improvement Project Measures	SCIP	95% adherence	Yes

^aExamples of changes: Catheter-related bloodstream infections decreased to ~1.7/1000 catheter-days; *C. difficile* infections increased to ~11.2 cases/10,000 discharges; catheter-related urinary tract infections decreased to ~3.1/1000 catheter-days; and hospital-onset MRSA invasive infections decreased to ~4.5/100,000 persons.

Abbreviations: EIP, CDC's Emerging Infections Program; HCUP, Agency for Healthcare Research and Quality's Healthcare Cost and Utilization Project; MRSA, methicillin-resistant *Staphylococcus aureus*; NHSN, CDC's National Healthcare Safety Network; SCIP, Surgical Care Improvement Project.

Source: Adapted from www.hhs.gov/ash/initiatives/hai/nationaltargets/ (last accessed November 13, 2014).

back to use of “house-wide” surveillance, and many states now require that hospitals use the Centers for Disease Control and Prevention's (CDC's) National Healthcare Safety Network (NHSN) reporting system to provide uniform definitions and to facilitate transmission of data. Increasing reliance on the NHSN by states to facilitate public reporting has led to participation by more than 12,000 facilities (~4700 of the ~5700 acute-care hospitals in the United States, ~540 LTACHs, ~270 inpatient rehabilitation facilities, ~6000 outpatient dialysis facilities, ~300 ambulatory surgery centers, ~150 long-term-care facilities). This level of participation provides a nationwide view of health care–associated infections and represents a watershed in potential access to national rates of antimicrobial use and resistance.

Results of surveillance are expressed as rates. In general, for example, 5–10% of patients develop nosocomial infections. However, such broad statistics have little value unless qualified by duration of risk, by site of infection, by patient population, and by exposure to risk factors. To account for some of these variables, the CDC now uses a Standardized Infection Ratio (SIR; www.cdc.gov/hai/national-annual-sir/) as part of NHSN rate reporting. Meaningful denominators for infection rates include the number of patients exposed to a specific risk (e.g., patients using mechanical ventilators) or the number of intervention days (e.g., 1000 patient-days on a ventilator). As use of invasive devices such as indwelling bladder catheters has purposely been decreased, the denominators have become smaller, but the fact that patients who still require such devices may be those at intrinsically higher risk (potential numerators) may paradoxically increase rates when device-days account for the denominator. Temporal trends in rates should be reviewed, and rates should be compared with regional and national benchmarks that incorporate the SIR. Interhospital comparisons still may be misleading because of the wide range in risk factors and severity of underlying illnesses. Process measures (e.g., adherence to hand hygiene) do not usually require risk adjustment, and outcome measures (e.g., cardiac surgery wound-infection rates) can identify hospitals with outlier infection rates (e.g., in the top deciles) for further evaluation. Moreover, temporal analysis of a hospital's infection rates can help to determine whether control measures are succeeding and where increased efforts should be focused.

EPIDEMIOLOGIC BASIS AND GENERAL MEASURES FOR PREVENTION AND CONTROL

Nosocomial infections follow basic epidemiologic patterns that can help to direct prevention and control measures. Nosocomial pathogens have reservoirs, are transmitted by largely predictable routes, and require susceptible hosts. Reservoirs and sources exist in the inanimate environment (e.g., residual *Clostridium difficile* spores on frequently touched surfaces in patients' rooms) and in the animate environment (e.g., infected or colonized health care workers, patients, and hospital visitors). The mode of transmission usually is either cross-infection (e.g., indirect spread of pathogens from one patient to another on the inadequately cleaned hands of hospital personnel) or autoinoculation (e.g., aspiration of oropharyngeal flora into the lungs along an endotracheal tube). Occasionally, pathogens (e.g., group A streptococci and many respiratory viruses) are spread from person to person via large infectious droplets released by coughing or sneezing. Much less common—but often devastating in terms of epidemic risk—is true airborne spread of small or droplet nuclei (as in nosocomial chickenpox) or common-source spread (e.g., by contaminated IV fluids). Factors that increase host susceptibility include underlying conditions, abnormalities of innate defense (e.g., due to genetic polymorphisms; see [Chap. 82](#)), and medical-surgical interventions and procedures that compromise host defenses.

Hospitals' infection-control programs must determine general and specific control measures. Given the prominence of cross-infection, hand hygiene is cited traditionally as the most important preventive measure. Health care workers' rates of adherence to hand-hygiene recommendations are abysmally low (often <50%). Reasons cited include inconvenience, time pressures, and skin damage from frequent washing. Sinkless alcohol rubs are quick and highly effective and actually improve hand condition since they contain emollients and allow the retention of natural protective oils that would be removed with repeated rinsing. Use of alcohol hand rubs between patient contacts is recommended for all health care workers except when hands are visibly soiled or after care of a patient who is part of a health care facility outbreak of infection with *C. difficile*, whose spores resist killing by alcohol and require mechanical removal. In these cases, washing with soap and running water is recommended. A number of innovative systems have been developed to track hand-hygiene adherence in real time and to provide feedback; although this approach is exciting, sustained improvements in rates remain to be seen.

NOSOCOMIAL AND DEVICE-RELATED INFECTIONS

The fact that >25–50% of nosocomial infections are due to the combined effect of the patient's own flora and invasive devices highlights the importance of improvements in the use and design of such devices. Intensive education, “bundling” of evidence-based interventions ([Table 168-4](#)), and use of checklists to facilitate adherence have reduced infection rates ([Table 168-3](#)) through improved asepsis in handling and earlier removal of invasive devices. It is especially noteworthy that turnover or shortages of trained personnel jeopardize safe and effective patient care and have been associated with increased infection rates.

URINARY TRACT INFECTIONS

Urinary tract infections (UTIs) account for ~30–40% of nosocomial infections; up to 3% of bacteriuric patients develop bacteremia. Although UTIs contribute at most 15% to prolongation of hospital stay and may have an attributable cost in the range of only \$1300, these infections are reservoirs and sources for spread of antibiotic-resistant bacteria. Most nosocomial UTIs are associated with preceding instrumentation or indwelling bladder catheters, which create a 3–7% risk of infection each day. UTIs generally are caused by pathogens that spread up the periurethral space from the patient's perineum or gastrointestinal tract—the most common pathogenesis in women—or via intraluminal contamination of urinary catheters, usually due to cross-infection by caregivers who are irrigating catheters or emptying drainage bags. Pathogens come occasionally from

TABLE 168-4 EXAMPLES OF EVIDENCE-BASED “BUNDLED INTERVENTIONS” TO PREVENT COMMON HEALTH CARE–ASSOCIATED INFECTIONS AND OTHER ADVERSE EVENTS**Prevention of Central Venous Catheter Infections***Catheter insertion bundle:*

- Educate personnel about catheter insertion and care.
- Use chlorhexidine to prepare the insertion site.
- Use maximal barrier precautions and asepsis during catheter insertion.
- Consolidate insertion supplies (e.g., in an insertion kit or cart).
- Use a checklist to enhance adherence to the “insertion bundle.”
- Empower nurses to halt insertion if asepsis is breached.

Catheter maintenance bundle:

- Cleanse patients daily with chlorhexidine.
- Maintain clean, dry dressings.
- Enforce hand hygiene among health care workers.
- Ask daily: Is the catheter needed? Remove catheter if not needed or used.

Prevention of Ventilator-Associated Events

- Elevate head of bed to 30–45 degrees.
- Decontaminate oropharynx regularly with chlorhexidine (controversial).
- Give “sedation vacation” and assess readiness to extubate daily.
- Use peptic ulcer disease prophylaxis.
- Use deep-vein thrombosis prophylaxis (unless contraindicated).

Prevention of Surgical-Site Infections

- Choose a surgeon wisely.
- Administer prophylactic antibiotics within 1 h before surgery; discontinue within 24 h.
- Limit any hair removal to the time of surgery; use clippers or do not remove hair at all.
- Prepare surgical site with chlorhexidine-alcohol.
- Maintain normal perioperative glucose levels (cardiac surgery patients).^a
- Maintain perioperative normothermia (colorectal surgery patients).^a

Prevention of Urinary Tract Infections

- Place bladder catheters only when absolutely needed (e.g., to relieve obstruction), not solely for the provider’s convenience.
- Use aseptic technique for catheter insertion and urinary tract instrumentation.
- Minimize manipulation or opening of drainage systems.
- Ask daily: Is the bladder catheter needed? Remove catheter if not needed.

Prevention of Pathogen Cross-Transmission

- Cleanse hands with alcohol hand rub before and after all contacts with patients or their environments.

^aThese components of care are supported by clinical trials and experimental evidence in the specified populations; they may prove valuable for other surgical patients as well.

Source: Adapted from information presented at the following websites: www.cdc.gov/hicpac/pubs.html; www.cdc.gov/HAI/prevent/prevention.html; www.ihl.org.

inadequately disinfected urologic equipment and rarely from contaminated supplies.

Hospitals should monitor essential performance measures for preventing nosocomial UTIs (Table 168-4). Prompts to clinicians to assess a patient’s need for continued use of an indwelling bladder catheter can improve removal rates and lessen the risk of UTI. Guidelines for managing postoperative urinary retention (e.g., with bladder scanners) also may limit the use or duration of catheterization. Other approaches to the prevention of UTIs have included the use of topical meatal antimicrobial agents, drainage bag disinfectants, and anti-infective catheters. None of the latter three measures is considered routine.

Administration of systemic antimicrobial agents for other purposes decreases the risk of UTI during the first 4 days of catheterization, after which resistant bacteria or yeasts emerge as pathogens. Prophylactic antibiotic administration at the time of catheter removal has been reported to decrease the risk of UTI. Selective decontamination of the gut also is associated with a reduced risk. Again, however, none of these approaches is routine.

Irrigation of catheters, with or without antimicrobial agents, may actually increase the risk of infection. A condom catheter for men without bladder obstruction may be more acceptable than an indwelling catheter and may lessen the risk of UTI if maintained carefully. The role of suprapubic catheters in preventing infection is not well defined.

Treatment of UTIs is based on the results of quantitative urine cultures (Chap. 162). The most common pathogens are *Escherichia coli*, nosocomial gram-negative bacilli, enterococci, and *Candida*. Several caveats apply in the treatment of institutionally acquired infection. First, in patients with chronic indwelling bladder catheters, especially those in long-term-care facilities, “catheter flora”—microorganisms living on encrustations within the catheter lumen—may differ from actual urinary tract pathogens. Therefore, for suspected UTI in the setting of chronic catheterization (especially in women), it is useful to replace the bladder catheter and to obtain a freshly voided urine specimen. Second, as in all nosocomial infections, at the time treatment is initiated on the basis of a positive culture, it is useful to repeat the culture to verify the persistence of infection. Third, the frequency with which UTIs occur may lead to the erroneous assumption that the urinary tract alone is the source of infection in a febrile hospitalized patient. Fourth, recovery of *Staphylococcus aureus* from urine cultures may result from hematogenous seeding and may indicate an occult systemic infection. Finally, although *Candida* is now the most common pathogen in nosocomial UTIs among patients on intensive care units (ICUs), treatment of candiduria is often unsuccessful and is recommended only when there is upper-pole or bladder-wall invasion, obstruction, neutropenia, or immunosuppression.

PNEUMONIA

Historically, pneumonia has accounted for ~10–15% of nosocomial infections; ventilator-associated pneumonia (VAP) occurred in 1 to >4 patients per 1000 ventilator-days, and these infections were reported as responsible for a mean of 10 extra hospital days and \$23,000 in extra costs per episode. Most cases of bacterial nosocomial pneumonia are caused by aspiration of endogenous or hospital-acquired oropharyngeal (and occasionally gastric) flora. Nosocomial pneumonias are associated with more deaths than are infections at any other body site. However, attributable mortality rates suggest that the risk of dying from nosocomial pneumonia is affected greatly by other factors, including comorbidities, inadequate antibiotic treatment, and the involvement of specific pathogens (particularly *Pseudomonas aeruginosa* or *Acinetobacter*). Surveillance and accurate diagnosis of pneumonia have been problematic in hospitals because many patients, especially those in the ICU, have abnormal chest roentgenographs, fever, and leukocytosis potentially attributable to multiple causes. This diagnostic uncertainty has led to a refocus from VAP to “ventilator-associated events” (VAEs), conditions, and complications, for which worsening physiologic parameters, such as oxygenation, are key metrics. Early data suggest that ~5–10% of patients using mechanical ventilators develop VAEs. Viral pneumonias, which are particularly important in pediatric and immunocompromised patients, are discussed in the virology section and in Chap. 153.

Risk factors for nosocomial pneumonia include those events that increase colonization by potential pathogens (e.g., prior antimicrobial therapy, contaminated ventilator circuits or equipment, or decreased gastric acidity); those that facilitate aspiration of oropharyngeal contents into the lower respiratory tract (e.g., intubation, decreased levels of consciousness, or presence of a nasogastric tube); and those that reduce host defense mechanisms in the lung and permit overgrowth of aspirated pathogens (e.g., chronic obstructive pulmonary disease, extremes of age, or upper abdominal surgery).

Control measures for pneumonia (Table 168-4) are aimed at frequent testing of readiness for extubation, remediation of risk factors in patient care (e.g., minimizing aspiration-prone supine positioning), and aseptic care of respirator equipment (e.g., disinfecting or sterilizing all inline reusable components such as nebulizers, replacing tubing/breathing circuits only if required because of malfunction or visible soiling—rather than on the basis of duration of use—to lessen

the number of breaks in the system, and teaching aseptic technique for suctioning). Although the benefits of selective decontamination of the oropharynx and gut with nonabsorbable antimicrobial agents and/or use of short-course postintubation systemic antibiotics have been controversial, a randomized multicenter trial demonstrated lowered ICU mortality rates among patients on mechanical ventilation who underwent oropharyngeal decontamination.

Among the logical preventive measures that require further investigation are placement of endotracheal tubes that provide channels for subglottic drainage of secretions, which has been associated with reduced infection risks during short-term postoperative use, and non-invasive mechanical ventilation whenever feasible. Use of silver-coated endotracheal tubes may lessen risk of VAP but is not considered routine. It is noteworthy that reducing the rate of VAP often has not reduced overall ICU mortality; this fact suggests that this infection is a marker for patients with an otherwise-heightened risk of death.

The most likely pathogens for nosocomial pneumonia and treatment options are discussed in **Chap. 153**. Several considerations regarding diagnosis and treatment are worth emphasizing. First, clinical criteria for diagnosis (e.g., fever, leukocytosis, development of purulent secretions, new or changing radiographic infiltrates, changes in oxygen requirement or ventilator settings) have high sensitivity but relatively low specificity. These criteria are most useful for selecting patients for bronchoscopic or nonbronchoscopic procedures that yield lower respiratory tract samples protected from upper-tract contamination; quantitative cultures of such specimens have diagnostic sensitivities in the range of 80%. Second, early-onset nosocomial pneumonia, which manifests within the first 4 days of hospitalization, is most often caused by community-acquired pathogens such as *Streptococcus pneumoniae* and *Haemophilus* species, although some studies have challenged this view. Late-onset pneumonias most commonly are due to *S. aureus*, *P. aeruginosa*, *Enterobacter* species, *Klebsiella pneumoniae*, or *Acinetobacter*. When invasive techniques are used to diagnose VAP, the proportion of isolates accounted for by gram-negative bacilli decreases from 50–70% to 35–45%. Infection is polymicrobial in as many as 20–40% of cases. The role of anaerobic bacteria in VAP is not well defined. Third, one multicenter study suggested that 8 days is an appropriate duration of therapy for nosocomial pneumonia, with a longer duration (15 days in that study) when the pathogen is *Acinetobacter* or *P. aeruginosa*. Finally, in febrile patients (particularly those who have endotracheal or gastric tubes inserted through the nares), occult respiratory tract infections, especially bacterial sinusitis and otitis media, should be considered.

SURGICAL WOUND INFECTIONS

Wound infections occur in ~500,000 patients each year, account for ~15–20% of nosocomial infections, contribute up to 7–10 extra postoperative hospital days, and result in \$3000 to \$29,000 in extra costs, depending on the operative procedure and pathogen(s). The average wound infection has an incubation period of 5–7 days—longer than many postoperative stays. For this reason and because many procedures are now performed on an outpatient basis, the incidence of wound infections has become more difficult to assess. These infections usually are caused by the patient's endogenous or hospital-acquired skin and mucosal flora and occasionally are due to airborne spread of skin squames that may be shed into the wound from members of the operating-room team. True airborne spread of infection through droplet nuclei is rare in operating rooms unless there is a “disseminator” (e.g., of group A streptococci or staphylococci) among the staff. In general, the common risks for postoperative wound infection are related to the surgeon's technical skill, the patient's underlying conditions (e.g., diabetes mellitus, obesity) or advanced age, and inappropriate timing of antibiotic prophylaxis. Additional risks include the presence of drains, prolonged preoperative hospital stays, shaving of operative sites by razor the day before surgery, long duration of surgery, and infection at remote sites (e.g., untreated UTI).

The substantial literature related to risk factors for surgical-site infections and the recognized morbidity and cost of these infections have led to national prevention efforts and to recommendations for

“bundling” preventive measures (Table 168-4). Additional measures include attention to technical surgical issues (e.g., avoiding open or prophylactic drains), operating-room asepsis, and preoperative therapy for active infection. Reporting surveillance results to surgeons has been associated with reductions in infection rates. Preoperative administration of intranasal mupirocin to patients colonized with *S. aureus*, preoperative antiseptic bathing, and intra- and postoperative oxygen supplementation have been controversial because of conflicting study results, but evidence seems mostly to favor these interventions.

The process of diagnosing and treating wound infections begins with a careful assessment of the surgical site in the febrile postoperative patient. Diagnosis of deeper organ-space infections or subphrenic abscesses requires a high index of suspicion and the use of CT or MRI. Diagnosis of infections of prosthetic devices, such as orthopedic implants, may be particularly difficult and often requires the use of interventional radiographic techniques to obtain periprosthetic specimens for culture. Cultures of periprosthetic joint tissue obtained at surgery may miss pathogens that are cloistered in prosthesis-adherent biofilms; cultures of sonicates from explanted prosthetic joints have been more sensitive, particularly for patients who have received antimicrobial agents within 2 weeks of surgery.

The most common pathogens in postoperative wound infections are *S. aureus*, coagulase-negative staphylococci, and enteric and anaerobic bacteria. In rapidly progressing postoperative infections manifesting within 24–48 h of a surgical procedure, the level of suspicion regarding group A streptococcal or clostridial infection (**Chaps. 173 and 179**) should be high. Treatment of postoperative wound infections requires drainage or surgical excision of infected or necrotic material and antibiotic therapy aimed at the most likely or laboratory-confirmed pathogens.

INFECTIONS RELATED TO VASCULAR ACCESS AND MONITORING

Intravascular device-related bacteremias cause ~10–15% of nosocomial infections; central vascular catheters (CVCs) account for most of these bloodstream infections. Past national estimates indicated that as many as 200,000 bloodstream infections associated with CVCs occurred each year in the United States, with attributable mortality rates of 12–25%, an excess mean length of hospital stay of 12 days, and an estimated cost of \$3700 to \$29,000 per episode; one-third to one-half of these episodes occurred in ICUs. However, infection rates have dropped steadily (Table 168-3) since the publication of guidelines by the Healthcare Infection Control Practices Advisory Committee (HICPAC) in 2002. With increasing care of seriously ill patients in the community, vascular catheter-associated bloodstream infections acquired in outpatient settings are becoming more frequent. Broader surveillance for infections—outside ICUs and even outside hospitals—will be needed.

Catheter-related bloodstream infections derive largely from the cutaneous microflora of the insertion site, with pathogens migrating extraluminally to the catheter tip, usually during the first week after insertion. In addition, contamination of the hubs of CVCs or of the ports of “needle-less” systems may lead to intraluminal infection over longer periods, particularly with surgically implanted or cuffed catheters. Intrinsic (during the manufacturing process) or extrinsic (on-site in a health care facility) contamination of infusate, although rare, is the most common cause of epidemic device-related bloodstream infection; extrinsic contamination may cause up to half of endemic bacteremias related to arterial infusions used for hemodynamic monitoring. The most common pathogens isolated from vascular device-associated bacteremias include coagulase-negative staphylococci, *S. aureus* (with ≥50% of isolates in the United States resistant to methicillin), enterococci, nosocomial gram-negative bacilli, and *Candida*. Many pathogens, especially staphylococci, produce extracellular polysaccharide biofilms that facilitate attachment to catheters and provide sanctuary from antimicrobial agents. “Quorum-sensing” proteins, a target for future interventions, help bacterial cells communicate during biofilm development.

Evidence-based bundles of control measures (Table 168-4) have been strikingly effective, eliminating almost all CVC-associated infections in one ICU study. Additional control measures for infections associated with vascular access include use of a chlorhexidine-impregnated

916 patch at the skin-catheter junction; daily bathing of ICU patients with chlorhexidine; application of semitransparent access-site dressings (for ease of bathing and site inspection and protection of the site from secretions); avoidance of the femoral site for catheterization because of a higher risk of infection (most likely related to the density of the skin flora); rotation of peripheral catheters to a new site at specified intervals (e.g., every 72–96 h), which may be facilitated by use of an IV therapy team; and application of aseptic technique when accessing pressure transducers or other vascular ports.

Unresolved issues include the role of gut translocation rather than vascular access sites as a cause of primary bacteremia in immunocompromised patients and the implications for surveillance definitions; the best frequency for rotation of CVC sites (given that guidewire-assisted catheter changes at the same site do not lessen and can even increase infection risk); the appropriate role of mupirocin ointment, a topical antibiotic with excellent antistaphylococcal activity, in site care; the relative degrees of risk posed by peripherally inserted central catheters (PICC lines); and the risk-benefit of prophylactic use of heparin (to avoid catheter thrombi, which may be associated with increased risk of infection) or of vancomycin or alcohol (as catheter flushes or “locks”—i.e., concentrated anti-infective solutions instilled into the catheter lumen) for high-risk patients.

Vascular device–related infection is suspected on the basis of the appearance of the catheter site or the presence of fever or bacteremia without another source in patients with vascular catheters. The diagnosis is confirmed by the recovery of the same species of microorganism from peripheral-blood cultures (preferably two samples drawn from peripheral veins by separate venipunctures) and from semiquantitative or quantitative cultures of the vascular catheter tip. Less commonly used diagnostic measures include (1) differential (faster) time to positivity (>2 h) for blood drawn through the vascular access device than for a sample from a peripheral vein and (2) differences in quantitative cultures (a threefold or greater “step-up”) for blood samples drawn simultaneously from a peripheral vein and from a CVC, which should show the step-up if infected. When infusion-related sepsis is considered (e.g., because of the abrupt onset of fever or shock temporally related to infusion therapy), a sample of the infusate or blood product should be retained for culture.

Therapy for vascular access–related infection is directed at the pathogen recovered from the blood and/or infected site. Important considerations in treatment are the need for an echocardiogram (to evaluate the patient for endocarditis), the duration of therapy, and the need to remove potentially infected catheters. In one report, approximately one-fourth of patients with intravascular catheter–associated *S. aureus* bacteremia who were studied by transesophageal echocardiography had evidence of endocarditis; this test may be useful in determining the appropriate duration of treatment.

Detailed consensus guidelines for the management of intravascular catheter–related infections have been published and recommend catheter removal in most cases of bacteremia or fungemia due to nontunneled CVCs. When attempting to salvage a potentially infected catheter, some clinicians use the “antibiotic lock” technique, which may facilitate penetration of infected biofilms, in addition to systemic antimicrobial therapy (see www.idsociety.org/Other_Guidelines/).

The authors of the consensus treatment guidelines advise that the decision to remove a tunneled catheter or implanted device suspected of being the source of bacteremia or fungemia should be based on the severity of the patient’s illness, the strength of evidence that the device is infected, the presence of local or systemic complications, an assessment of the specific pathogens, and the patient’s response to antimicrobial therapy if the catheter or device is initially retained. For patients with track-site infection, successful therapy without catheter removal is unusual. For patients with suppurative venous thrombophlebitis, excision of affected veins is usually required.

ISOLATION TECHNIQUES

Written policies for the isolation of infectious patients are a standard component of infection control programs. To replace its prior pathogen-specific guidelines, the CDC published recommendations in 2006 for

the control of multidrug-resistant organisms in health care settings; in 2007, the CDC published a revised edition of its basic isolation guidelines to provide updated recommendations for all components of health care, including acute-care hospitals and long-term, ambulatory, and home-care settings (see www.cdc.gov/hicpac/pdf/isolation/Isolation2007.pdf).

Standard precautions are designed for the care of all patients in hospitals and aim to reduce the risk of transmission of microorganisms from both recognized and unrecognized sources. These precautions include gloving as well as hand cleansing for potential contact with (1) blood; (2) all other body fluids, secretions, and excretions, whether or not they contain visible blood; (3) nonintact skin; and (4) mucous membranes. Depending on exposure risks, standard precautions also include use of masks, eye protection, and gowns.


Precautions for the care of patients with potentially contagious clinical syndromes (e.g., acute diarrhea) or with suspected or diagnosed colonization or infection by transmissible pathogens are based on probable routes of transmission: *airborne*, *droplet*, or *contact*, for which personnel don, at a minimum, N95 respirators, surgical face masks, or glove and gown, respectively. Sets of precautions may be combined for diseases that have more than one route of transmission (e.g., contact and airborne isolation for varicella).

Some prevalent antibiotic-resistant pathogens, particularly those that colonize the gastrointestinal tract (e.g., vancomycin-resistant enterococci [VRE] and even multidrug-resistant gram-negative bacilli such as carbapenemase-producing strains of *K. pneumoniae* [KPCs]), may be present on *intact* skin of patients in hospitals (the “fecal patina”). This issue has led some experts to recommend gloving for all contact with patients who are acutely ill and/or in high-risk units, such as ICUs or LTACHs. Wearing gloves does not replace the need for hand hygiene because hands sometimes (in up to 20% of interactions) become contaminated during wearing or removal of gloves.

EPIDEMIC AND EMERGING PROBLEMS

Outbreaks are always big news but probably account for <5% of nosocomial infections. The investigation and control of nosocomial epidemics require that infection control personnel (1) develop a case definition, (2) confirm that an outbreak really exists (since apparent epidemics may actually be pseudo-outbreaks due to surveillance or laboratory artifacts), (3) review aseptic practices and disinfectant use, (4) determine the extent of the outbreak, (5) perform an epidemiologic investigation to determine modes of transmission, (6) work closely with microbiology personnel to culture for common sources or personnel carriers as appropriate and to type epidemiologically important isolates, and (7) heighten surveillance to judge the effect of control measures. Control measures generally include reinforcing routine aseptic practices and hand hygiene, ensuring appropriate isolation of cases (and instituting cohort isolation and nursing if needed), and implementing further controls on the basis of the investigation’s findings. Examples of some emerging and potential epidemic problems follow.

VIRAL RESPIRATORY INFECTIONS: PANDEMIC INFLUENZA

 Infections caused by the severe acute respiratory syndrome (SARS)–associated coronavirus challenged health care systems globally in 2003 ([Chap. 223](#)), and in 2012 Middle East respiratory syndrome coronavirus (MERS-CoV) emerged as a more geographically localized problem ([Chap. 223](#)). For SARS, basic infection-control measures helped to keep the worldwide case and death counts at ~8000 and ~800, respectively, although the virus was unforgiving of lapses in protocol adherence or laboratory biosafety. The epidemiology of SARS—spread largely in households once patients were ill or in hospitals—contrasts markedly with that of influenza ([Chap. 224](#)), which is often contagious a day before symptom onset; can spread rapidly in the community among nonimmune persons; and, even in its seasonal variety, kills as many as 35,000 persons each year in the United States.

Control of seasonal influenza has depended on (1) the use of effective vaccines, with increasingly broad evidence-based recommendations for vaccination of children, the general public, and health care

workers; (2) the use of antiviral medications for early treatment and for prophylaxis as part of outbreak control, especially for high-risk patients and in high-risk settings like nursing homes or hospitals; and (3) infection control (surveillance and droplet precautions) for symptomatic patients. Controversial infection-control issues have been the questionable role of airborne spread of influenza and the need to mandate influenza vaccination of health care workers because of the embarrassingly low rates of vaccination in this high-risk group.

With the occurrence of localized outbreaks of avian (H5N1) influenza in Asia over the past few years, concerns about potential pandemic influenza led to (1) recommendations for universal “respiratory hygiene and cough etiquette” (basically, “cover your cough”), as described and promoted in the CDC’s 2007 *Guideline for Isolation Precautions*, and for “source containment” (e.g., use of face masks and spatial separation) for outpatients with potentially infectious respiratory illnesses; (2) re-examinations of the value in the 1918–1919 influenza pandemic of nonpharmacologic interventions, such as “social distancing” (e.g., closing of schools and community venues); and (3) debate about the level of respiratory protection required for health care workers (i.e., whether to use the higher-efficiency N95 respirators recommended for airborne isolation rather than the surgical masks used for droplet precautions).

In the spring of 2009, a novel strain of influenza virus—H1N1 or “swine flu” virus—caused the first influenza pandemic in four decades. Recombinant events that create new strains (e.g., H7N9) continue to challenge global efforts at infection control and vaccine development (Chap. 224).

NOSOCOMIAL DIARRHEA

A new, more virulent strain of *C. difficile*—NAP1/BI/027—emerged in North America, and overall rates of *C. difficile*-associated diarrhea (Chap. 161) have increased, especially among older patients, in U.S. hospitals during the past few years. *C. difficile* control measures include judicious use of all antibiotics, especially fluoroquinolone antibiotics that have been implicated in driving these changes; heightened suspicion for “atypical” presentations (e.g., toxic megacolon or leukemoid reaction without diarrhea); and early diagnosis, treatment, and contact precautions. To improve diagnosis, use of more sensitive polymerase chain reaction–based rather than enzyme immunoassay–based testing of diarrheal stool is now recommended, with resultant artificial doubling of infection rates in some hospitals. Preliminary data suggest a role for probiotics in the prevention of *C. difficile*-associated diarrhea in patients in whom systemic antibiotic therapy is being initiated. Fecal transplantation has had dramatic results in the treatment of relapsing cases of *C. difficile*-associated diarrhea (Chap. 161). Successes with fecal transplants and probiotics have called attention to the potential role of manipulation of the intestinal microbiome as a broader infection-control strategy.

Outbreaks of norovirus infection (Chap. 227) in U.S. and European health care facilities appear to continue to increase in frequency or at least in reporting, with the virus often introduced by ill visitors or staff. This pathogen should be suspected when nausea and vomiting are prominent aspects of bacterial culture–negative diarrheal syndromes. Contact precautions may need to be augmented by aggressive environmental cleaning (given the persistence of norovirus on inanimate objects), prevention of secondary cases in cleaning staff through an emphasis on the use of personal protective equipment and hand hygiene, and active exclusion of ill staff and visitors.

CHICKENPOX

Infection control practitioners institute a varicella exposure investigation and control plan whenever health care workers have been exposed to chickenpox (Chap. 217) or have worked while having or during the 24 h before developing chickenpox. The names of exposed workers and patients are obtained; medical histories are reviewed, and (if necessary) serologic tests for immunity are conducted; physicians are notified of susceptible exposed patients; postexposure prophylaxis with a preparation of varicella-zoster immune globulin (VZIG) is considered for immunocompromised or pregnant contacts, with

administration as soon as possible (but as long as 10 days after exposure) (Table 217-1); varicella vaccine is recommended or preemptive use of acyclovir is considered as an alternative strategy in other susceptible persons; and susceptible exposed employees are furloughed during the at-risk period for disease (8–21 days or, if VZIG has been administered, 28 days). Routine varicella vaccination of children and susceptible employees has made nosocomial spread less common and less problematic.

TUBERCULOSIS

Important measures for the control of tuberculosis (Chap. 202) include prompt recognition, isolation, and treatment of cases; recognition of atypical presentations (e.g., lower-lobe infiltrates without cavitation); use of negative-pressure, 100% exhaust, private isolation rooms with closed doors and at least 6–12 air changes per hour; use of N95 respirators by caregivers entering isolation rooms; possible use of high-efficiency particulate air filter units and/or ultraviolet lights for disinfecting air when other engineering controls are not feasible or reliable; and follow-up testing of susceptible personnel who have been exposed to infectious patients before isolation. The use of serologic tests, rather than skin tests, in the diagnosis of latent tuberculosis for infection control purposes has become common, mostly for logistic reasons. As tuberculosis once again is on the decline in the United States, we need to remember that the price of freedom—in this instance, from a communicable disease—is eternal vigilance.

GROUP A STREPTOCOCCAL INFECTIONS

The potential for an outbreak of group A streptococcal infection (Chap. 173) should be considered when even one or two nosocomial cases occur. Most outbreaks involve surgical wounds and are due to the presence of an asymptomatic carrier in the operating room. Investigation can be confounded by carriage at extrapharyngeal sites such as the rectum and vagina. Health care workers in whom carriage has been linked to nosocomial transmission of group A streptococci are removed from the patient-care setting and are not permitted to return until carriage has been eliminated by antimicrobial therapy.

FUNGAL INFECTIONS

Fungal spores are common in the environment, particularly on dusty surfaces. When dusty areas are disturbed during hospital repairs or renovation, the spores become airborne. Inhalation of spores by immunosuppressed (especially neutropenic) patients creates a risk of pulmonary and/or paranasal sinus infection and disseminated aspergillosis (Chap. 241). Routine surveillance among neutropenic patients for infections with filamentous fungi, such as *Aspergillus* and *Fusarium*, helps hospitals to determine whether they are facing environmental risks. As a matter of routine, hospitals should inspect and clean air-handling equipment, review all planned renovations with infection control personnel and subsequently construct appropriate barriers, remove immunosuppressed patients from renovation sites, and consider the use of high-efficiency particulate air intake filters for rooms housing immunosuppressed patients.

A major multistate iatrogenic outbreak of meningitis, localized spinal or paraspinal infection, and arthritis due to *Exserohilum rostratum* was recognized in 2012 and traced to contamination of an injectable preservative-free steroid product produced by a single compounding pharmacy (Chap. 241).

LEGIONELLOSIS

Nosocomial *Legionella* pneumonia (Chap. 184) is most often due to contamination of potable water and predominantly affects immunosuppressed patients, particularly those receiving glucocorticoid medications. The risk varies greatly within and among geographic regions, depending on the extent of hospital water contamination and on specific hospital practices (e.g., inappropriate use of nonsterile water in respiratory therapy equipment). Laboratory-based surveillance for nosocomial *Legionella* should be performed, and a diagnosis of legionellosis should probably be considered more often than it is. If nosocomial cases are detected, environmental samples (e.g., tap water)

918 should be cultured. If cultures yield *Legionella* and if typing of clinical and environmental isolates reveals a correlation, eradication measures should be pursued. An alternative approach is to periodically culture tap water in wards housing high-risk patients. If *Legionella* is found, a concerted effort should be made to culture samples from all patients with nosocomial pneumonia for *Legionella*.

ANTIBIOTIC-RESISTANT BACTERIA

Emerging multidrug-resistant bacteria like KPCs are harbingers of a potential “postantibiotic” era. Control of antibiotic resistance depends on close laboratory surveillance, with early detection of problems; on aggressive reinforcement of routine asepsis; on implementation of barrier precautions for all colonized and/or infected patients; on use of patient-surveillance cultures to more fully ascertain the extent of patient colonization; on antimicrobial stewardship to lessen ecologic pressures; and on timely initiation of an epidemiologic investigation when rates increase. Molecular typing (e.g., pulsed-field gel electrophoresis and, most recently, whole-genome sequencing) can help differentiate an outbreak due to a single strain (which necessitates an emphasis on hand hygiene and an evaluation of potential common-source exposures) from a polyclonal outbreak (which requires an emphasis on antibiotic prudence and device bundles; Table 168-4). Continuing emergence of multidrug-resistant organisms suggests that control efforts have been insufficient and that regional or broader (national and global) strategies and interventions are urgently needed (see www.cdc.gov/drugresistance/threat-report-2013/ and www.gov.uk/government/publications/uk-5-year-antimicrobial-resistance-strategy-2013-to-2018/).

Currently, several antibiotic resistance problems are of particular concern. First, over the past decade or so, the emergence of community-associated methicillin-resistant *S. aureus* (CA-MRSA) has been dramatic in many countries, with as many as 50% of community-acquired “staph infections” in some U.S. cities now caused by strains resistant to β -lactam antibiotics (Chap. 172). The incursion of CA-MRSA into hospitals is well documented and has impacted surveillance and control of nosocomial MRSA infections.

Second, in the ongoing global reemergence of nosocomial multidrug-resistant gram-negative bacilli, new problems include plasmid-mediated resistance to fluoroquinolones, metallo- β -lactamase-mediated resistance to carbapenems, KPCs, and panresistant strains of *Acinetobacter*. The problematic New Delhi metallo- β -lactamase (NDM) is plasmid-mediated, has been highly successful in inter-genus transmission, and has quickly become a global threat (see wwwnc.cdc.gov/eid/article/17/10/11-0655_article.htm). For several years, KPCs were a very focal problem in the United States (predominantly in Brooklyn, NY), but more recently these strains have become a national threat. Many multidrug-resistant gram-negative bacilli are susceptible only to colistin, a drug that is consequently being “rediscovered,” or to no available agents.

Third, there has been renewed recognition of the role of nursing homes, and now LTACHs, in the spread of resistant gram-negative bacilli such as KPCs. In some LTACHs, as many as 30–50% of patients may be colonized with KPCs.

Fourth, there has been increasing community-based spread of *E. coli* strains harboring an enzyme, CTX-M, that renders them broadly resistant to β -lactam antibiotics. Given the community focus of spread, these strains may be seen as a gram-negative version of CA-MRSA.

Finally, clinical infections with MRSA strains exhibiting high-level vancomycin resistance due to VRE-derived plasmids have been reported in a few patients—almost all in the United States and most in Michigan—in the setting of prolonged or repeated treatment with vancomycin and/or VRE colonization. Much more common is vancomycin “MIC creep”: an increasing prevalence of MRSA strains that exhibit upper-limit susceptibility to vancomycin.

Colonized personnel who are implicated in nosocomial transmission of multidrug-resistant pathogens and patients who pose a threat can be decontaminated, depending on the pathogen. In a few ICUs,

nonabsorbed antimicrobial agents for gastrointestinal decontamination of patients have been used successfully as a temporary emergency control measure for outbreaks of infection due to gram-negative bacilli. Potentially, manipulation of patients’ intestinal microbiome could be a more durable strategy to control outbreaks of multidrug-resistant pathogens that have a gastrointestinal reservoir.



In several trials over the past 10 years, source control—i.e., removal of patients’ fecal patina—by daily bathing with chlorhexidine has reduced the risk of bacteremia in ICU patients. “Search-and-destroy” methods—i.e., active surveillance cultures to detect and isolate the “resistance iceberg” of patients colonized with MRSA—in nonoutbreak settings are credited with elimination of nosocomial MRSA in the Netherlands and Denmark. In a recent multicenter trial in the United States, universal source control with chlorhexidine and nasal mupirocin was significantly more effective for controlling MRSA than was a search-and-destroy approach and led to control of other pathogens as well, providing a broad (“horizontal”) rather than a narrower (“vertical”) intervention (see www.ahrq.gov/professionals/systems/hospital/universal_icu_decolonization/). For some pathogens, such as VRE, enforcement of environmental cleaning also reduces cross-transmission risk.

Because the excessive use of broad-spectrum antibiotics underlies many resistance problems, “antibiotic stewardship” has been promulgated actively. The main tenets are to restrict the use of particular agents to narrowly defined indications in order to limit selective pressure on the nosocomial flora and, when broad-spectrum therapy is begun empirically in critically ill patients, to “de-escalate” treatment as soon as possible on the basis of the results of culture and susceptibility tests.

BIOTERRORISM AND OTHER “SURGE-EVENT” PREPAREDNESS



The horrific attack on the World Trade Center in New York City on September 11, 2001; the subsequent mailings of anthrax spores in the United States; the Boston Marathon bombing in 2013; and ongoing revelations of terrorist plans and activities in many other countries as well as the United States have made bioterrorism a prominent source of concern to hospital infection-control programs. The essentials for hospital preparedness entail education, internal and external communication, and risk assessment. Up-to-date information is available from the CDC (see www.bt.cdc.gov).

EMPLOYEE HEALTH SERVICE ISSUES

An institution’s employee health service is a critical component of its infection control efforts. New employees should be processed through the service, where a contagious-disease history can be taken; evidence of immunity to a variety of diseases, such as hepatitis B, chickenpox, measles, mumps, and rubella, can be sought; immunizations for hepatitis B, measles, mumps, rubella, varicella, and pertussis (the only vaccine-preventable childhood disease that is on the rise again in the United States) can be given as needed; baseline tuberculosis testing can be performed; and education about personal responsibility for infection control can be initiated. Evaluations of employees should be codified to meet the requirements of accrediting and regulatory agencies.

The employee health service must have protocols for dealing with workers exposed to contagious diseases (e.g., influenza) and those percutaneously or mucosally exposed to the blood of patients infected with HIV or hepatitis B or C virus. For example, postexposure HIV prophylaxis (PEP) with combination antiretroviral agents is recommended, as indicated; free consultation is available from the CDC-funded PEPLine (888-HIV-4911). Protocols are also needed for dealing with caregivers who have common contagious diseases (such as chickenpox, group A streptococcal infection, influenza or another respiratory infection, or infectious diarrhea) and for those who have less common but high-visibility public health problems (such as chronic hepatitis B or C or HIV infection) for which exposure-control guidelines have been published by the CDC and by the Society for Healthcare Epidemiology of America.

This chapter considers aspects of infection unique to patients receiving transplanted tissue. The evaluation of infections in transplant recipients involves consideration of both the donor and the recipient of the transplanted cells or organ. Two central issues are of paramount importance: (1) infectious agents (particularly viruses, but also bacteria, fungi, and parasites) can be introduced into the recipient by the donor; and (2) treatment of the recipient with medicine to prevent rejection can suppress normal immune responses, greatly increasing susceptibility to infection. Thus, what might have been a latent or asymptomatic infection in an immunocompetent donor or in the recipient prior to therapy can become a life-threatening problem when the recipient becomes immunosuppressed. The pretransplantation evaluation of each patient should be guided by an analysis of both (1) what infections the recipient is currently harboring, since organisms that exist in a state of latency or dormancy before the procedure may cause fatal disease when the patient receives immunosuppressive treatment; and (2) what organisms are likely to be transmitted by the donor, particularly those to which the recipient may be naïve.

PRETRANSPLANTATION EVALUATION

The Donor A variety of organisms have been transmitted by organ transplantation. Transmission of infections that may have been latent or not clinically apparent in the donor has resulted in the development of specific donor-screening protocols. Results from routine blood bank studies, including those for antibodies to *Treponema pallidum* (syphilis), *Trypanosoma cruzi*, hepatitis B and C viruses, HIV-1 and -2, human T-lymphotropic virus types 1 and 2 (HTLV-1 and -2), and West Nile virus (WNV), should be documented. Serologic studies should be ordered to identify latent infection with viruses such as herpes simplex virus types 1 and 2 (HSV-1, HSV-2), varicella-zoster virus (VZV), cytomegalovirus (CMV), Epstein-Barr virus (EBV), Kaposi's sarcoma-associated herpesvirus (KSHV); acute infection with hepatitis A virus; and infection with the common parasite *Toxoplasma gondii*. Donors should be screened, when relevant, for viruses such as rabies virus and lymphocytic choriomeningitis virus as well as for parasites such as *Strongyloides stercoralis* and *Schistosoma* species. Clinicians caring for prospective organ donors should examine chest radiographs for evidence of granulomatous disease (e.g., caused by mycobacteria or fungi) and should perform skin testing or obtain blood for immune cell–based assays that detect active or latent *Mycobacterium tuberculosis* infection. An investigation of the donor's dietary habits (e.g., consumption of raw meat or fish or of unpasteurized dairy products), occupations or avocations (e.g., gardening or spelunking), and travel history (e.g., travel to areas with endemic fungi) also is indicated and may mandate additional testing. Creutzfeldt-Jakob disease has been transmitted through corneal transplants. Whether it can be transmitted by transfused blood is not known. Variant Creutzfeldt-Jakob disease can be transmitted with transfused non-leukodepleted blood, posing a theoretical risk to transplant recipients.

The Recipient It is expected that the recipient will have been even more comprehensively assessed than the donor. Additional studies recommended for the recipient include evaluation for acute respiratory viruses and gastrointestinal pathogens in the immediate pretransplantation period. An important caveat is that, because of immune dysfunction resulting from chemotherapy or underlying chronic disease, serologic testing of the recipient may prove less reliable than usual.

The Donor Cells/Organ Careful attention to the sterility of the medium used to process the donor organ, combined with meticulous microbiologic evaluation, reduces rates of transmission of bacteria (or, rarely, yeasts) that may be present or grow in the organ culture medium. From 2% to >20% of donor kidneys are estimated to be contaminated with bacteria—in most cases, with the organisms that colonize the skin

or grow in the tissue culture medium used to bathe the donor organ while it awaits implantation. The reported rate of bacterial contamination of transplanted stem cells (bone marrow, peripheral blood, cord blood) is as high as 17% but most commonly is ~1%. The use of enrichment columns and monoclonal antibody depletion procedures results in a higher incidence of contamination. In one series of patients receiving contaminated stem cells, 14% had fever or bacteremia, but none died. Results of cultures performed at the time of cryopreservation and at the time of thawing were helpful in guiding therapy for the recipient.

INFECTIONS IN HEMATOPOIETIC STEM CELL TRANSPLANT RECIPIENTS

Transplantation of hematopoietic stem cells (HSCs) from bone marrow or from peripheral or cord blood for cancer, immunodeficiency, or autoimmune disease most often results in a transient state of complete immunologic incompetence. Immediately after myeloablative chemotherapy and transplantation, both innate immune cells (phagocytes, dendritic cells, natural killer cells) and adaptive immune cells (T and B cells) are absent, and the host is extremely susceptible to infection. The reconstitution that follows transplantation has been likened to maturation of the immune system in neonates. The analogy does not entirely predict infections seen in HSC transplant recipients, however, because the stem cells mature in an old host who has several latent infections already. The choice among the current variety of methods for obtaining stem cells is determined by availability and by the need to optimize the chances of a cure for an individual recipient. One strategy is autologous HSC transplantation, in which the donor and the recipient are the same. After chemotherapy, stem cells are collected and are purged (ex vivo) of residual neoplastic populations. Allogeneic HSC transplantation has the advantage of providing a graft-versus-tumor effect. In this case, the recipient is matched to varying degrees for human leukocyte antigens (HLAs) with a donor who may be related or unrelated. In some individuals, nonmyeloablative therapy (mini-allo transplantation) is used and permits recipient cells to persist for some time after transplantation while preserving the graft-versus-tumor effect and sparing the recipient myeloablative therapy. Cord-blood transplantation is increasingly utilized in adults; two independent cord-blood units are typically required for suitable neutrophil engraftment early after transplantation, even though only one of the units is likely to provide long-term engraftment. In each circumstance, a different balance is struck among the toxicity of conditioning therapy, the need for a maximal graft-versus-target effect, short-term and long-term infectious complications, and the risk of graft-versus-host disease (GVHD; acute versus chronic). The various approaches differ in terms of reconstitution speed, cell lineages introduced, and likelihood of GVHD—all factors that can produce distinct effects on the risk of infection after transplantation (Table 169-1). Despite these caveats, most infections occur in a predictable time frame after transplantation (Table 169-2).

BACTERIAL INFECTIONS

In the first month after HSC transplantation, infectious complications are similar to those in granulocytopenic patients receiving chemotherapy for acute leukemia (Chap. 104). Because of the anticipated 1- to 4-week duration of neutropenia and the high rate of bacterial infection in this population, many centers give prophylactic antibiotics to patients upon initiation of myeloablative therapy. Quinolones decrease the incidence of gram-negative bacteremia among these patients. Bacterial infections are common in the first few days after HSC transplantation. The organisms involved are predominantly those found on skin, mucosa, or IV catheters (*Staphylococcus aureus*, coagulase-negative staphylococci, streptococci) or aerobic bacteria that colonize the bowel (*Escherichia coli*, *Klebsiella*, *Pseudomonas*). *Bacillus cereus*, although rare, has emerged as a pathogen early after transplantation and can cause meningitis, which is unusual in these patients. Chemotherapy, use of broad-spectrum antibiotics, and delayed reconstitution of humoral immunity place HSC transplant patients at risk for diarrhea and colitis caused by *Clostridium difficile* overgrowth and toxin production.

TABLE 169-1 RISK OF INFECTION, BY TYPE OF HEMATOPOIETIC STEM CELL TRANSPLANT

Type of Hematopoietic Stem Cell Transplant	Source of Stem Cells	Risk of Early Infection: Neutrophil Depletion	Risk of Late Infection: Impaired T and B Cell Function	Risk of Ongoing Infection: GVHD and Iatrogenic Immunosuppression	Graft vs. Tumor Effect
Autologous	Recipient (self)	High risk; neutrophil recovery sometimes prolonged	~1 year	Minimal to no risk of GVHD and late-onset severe infection	None (–)
Syngeneic (genetic twin)	Identical twin	Low risk; 1–2 weeks for neutrophil recovery	~1 year	Minimal risk of GVHD and late-onset severe infection	+/-
Allogeneic related	Sibling	Low risk; 1–2 weeks for neutrophil recovery	~1 year	Minimal to moderate risk of GVHD and late-onset severe infection	++
Allogeneic related	Child/parent (haploidentical)	Intermediate risk; 2–3 weeks for neutrophil recovery	1–2 years	Moderate risk of GVHD and late-onset severe infection	++++
Allogeneic unrelated adult	Unrelated donor	Intermediate risk; 2–3 weeks for neutrophil recovery	1–2 years	High risk of GVHD and late-onset severe infection	++++
Allogeneic unrelated cord blood	Unrelated cord-blood units (x2)	Intermediate to high risk; neutrophil recovery sometimes prolonged	Prolonged	Minimal to moderate risk of GVHD and late-onset severe infection	++++
Allogeneic mini (nonmyeloablative)	Donor (transiently coexisting with recipient cells)	Low risk; neutrophil counts close to normal	1–2+ years	Variable risk of GVHD and late-onset severe infection ^a	++++ (but develops slowly)

^aDepending on the disparity of the match (major and minor histocompatibility antigens), GVHD may be severe or mild, the requirement for immunosuppression intense or minimal, and the risk of severe late infections coordinate with the degree of immunosuppression.

Abbreviation: GVHD, graft-versus-host disease.

Beyond the first few days of neutropenia, infections with nosocomial pathogens (e.g., vancomycin-resistant enterococci, *Stenotrophomonas maltophilia*, *Acinetobacter* species, and extended-spectrum β -lactamase-producing gram-negative bacteria) as well as with filamentous bacteria (e.g., *Nocardia* species) become more common. Vigilance is indicated, particularly for patients with a history of active or known latent tuberculosis, even when they have been appropriately pretreated. A form of bacterial colitis among cord-blood recipients has occurred 90–300 days after transplantation, responds to antimicrobial agents such as metronidazole, and—as determined by polymerase chain reaction (PCR) of biopsy specimens—may be attributed to the bacterium *Bradyrhizobium enterica* (related to *B. japonicum*). Episodes of bacteremia due to encapsulated organisms mark the late posttransplantation period (>6 months after HSC reconstitution); patients who have undergone splenectomy and those with persistent hypogammaglobulinemia are at particular risk.

FUNGAL INFECTIONS

Beyond the first week after transplantation, fungal infections become increasingly common, particularly among patients who have received broad-spectrum antibiotics. As in most granulocytopenic patients,

Candida infections are most commonly seen in this setting. However, with increased use of prophylactic fluconazole, infections with resistant fungi—in particular, *Aspergillus* and other non-*Aspergillus* molds (*Rhizopus*, *Fusarium*, *Scedosporium*, *Penicillium*)—have become more common, prompting some centers to replace fluconazole with agents such as micafungin, voriconazole, or posaconazole. The role of antifungal prophylaxis with these different agents, in contrast to empirical treatment for suspected infection that is based on a positive β -D-glucan assay or galactomannan antigen test, remains controversial (Chap. 104). Documented infection should be aggressively treated, ideally with agents of proven activity. In patients with GVHD who require prolonged or indefinite courses of glucocorticoids and other immunosuppressive agents (e.g., cyclosporine, tacrolimus [FK 506, Prograf], mycophenolate mofetil [Cellcept], rapamycin [sirolimus, Rapamune], antithymocyte globulin, or anti-CD52 antibody [alemtuzumab, Campath—an antilymphocyte and antimonocyte monoclonal antibody]), there is a high risk of fungal infection (usually with *Candida* or *Aspergillus*) even after engraftment and resolution of neutropenia. These patients are also at high risk for reactivation of latent fungal infection (histoplasmosis, coccidioidomycosis, or blastomycosis) in areas where endemic fungi reside and after involvement

TABLE 169-2 COMMON SOURCES OF INFECTIONS AFTER HEMATOPOIETIC STEM CELL TRANSPLANTATION

Infection Site	Period after Transplantation		
	Early (<1 Month)	Middle (1–4 Months)	Late (>6 Months)
Disseminated	Aerobic bacteria (gram-negative, gram-positive)	<i>Candida</i> , <i>Aspergillus</i> , EBV	Encapsulated bacteria (<i>Streptococcus pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Neisseria meningitidis</i>)
Skin and mucous membranes	HSV	HHV-6	VZV, HPV (warts)
Lungs	Aerobic bacteria (gram-negative, gram-positive), <i>Candida</i> , <i>Aspergillus</i> , other molds, HSV	CMV, seasonal respiratory viruses, <i>Pneumocystis</i> , <i>Toxoplasma</i>	<i>Pneumocystis</i> , <i>Nocardia</i> , <i>S. pneumoniae</i>
Gastrointestinal tract	<i>Clostridium difficile</i>	CMV, adenovirus, <i>Bradyrhizobium enterica</i> (cord blood cells)	EBV, CMV, <i>B. enterica</i> (cord blood cells)
Kidney		BK virus, adenovirus	
Brain		HHV-6, <i>Toxoplasma</i>	<i>Toxoplasma</i> , JC virus (rare)
Bone marrow		CMV, HHV-6	CMV, HHV-6

Abbreviations: CMV, cytomegalovirus; EBV, Epstein-Barr virus; HHV-6, human herpesvirus type 6; HPV, human papillomavirus; HSV, herpes simplex virus; VZV, varicella-zoster virus.

in activities such as gardening or caving. Prolonged use of central venous catheters for parenteral nutrition (lipids) increases the risk of fungemia with *Malassezia*. Some centers administer prophylactic antifungal agents to these patients. Because of the high and prolonged risk of *Pneumocystis jirovecii* pneumonia (especially among patients being treated for hematologic malignancies), most patients receive maintenance prophylaxis with trimethoprim-sulfamethoxazole (TMP-SMX) starting 1 month after engraftment and continuing for at least 1 year.

PARASITIC INFECTIONS

The regimen just described for the fungal pathogen *Pneumocystis* may also protect patients seropositive for the parasite *T. gondii*, which can cause pneumonia, visceral disease (occasionally), and central nervous system (CNS) lesions (more commonly). The advantages of maintaining HSC transplant recipients on daily TMP-SMX for 1 year after transplantation include some protection against *Listeria monocytogenes* and nocardial disease as well as late infections with *Streptococcus pneumoniae* and *Haemophilus influenzae*, which stem from the inability of the immature immune system to respond to polysaccharide antigens.

With increasing international travel, parasitic diseases typically restricted to particular environmental niches may pose a risk of reactivation in certain patients after HSC transplantation. Thus, in recipients with an appropriate history who were not screened and/or treated before transplantation or in patients with recent exposures, evaluation for infection with *Strongyloides*, *Leishmania*, schistosomes, trypanosomes, or various parasitic causes of diarrheal illness (*Giardia*, *Entamoeba*, *Cryptosporidium*, microsporidia) may be warranted.

VIRAL INFECTIONS

HSC transplant recipients are susceptible to infection with a variety of viruses, including primary and reactivation syndromes caused by most human herpesviruses (Table 169-3) and acute infections caused by viruses that circulate in the community.

TABLE 169-3 HERPESVIRUS SYNDROMES OF TRANSPLANT RECIPIENTS

Virus	Reactivation Disease
Herpes simplex virus type 1	Oral lesions
	Esophageal lesions
	Pneumonia (primarily HSC transplant recipients)
	Hepatitis (rare)
Herpes simplex virus type 2	Anogenital lesions
	Hepatitis (rare)
Varicella-zoster virus	Zoster (can disseminate)
Cytomegalovirus	Associated with graft rejection
	Fever and malaise
	Bone marrow failure
	Pneumonitis
Epstein-Barr virus	Gastrointestinal disease
	B cell lymphoproliferative disease/lymphoma
	Oral hairy leukoplakia (rare)
Human herpesvirus type 6	Fever
	Delayed monocyte/platelet engraftment
	Encephalitis (rare)
Human herpesvirus type 7	Undefined
Kaposi's sarcoma-associated virus	Kaposi's sarcoma
	Primary effusion lymphoma (rare)
	Multicentric Castleman's disease (rare)
	Marrow aplasia (rare)

Abbreviation: HSC, hematopoietic stem cell.

Herpes Simplex Virus Within the first 2 weeks after transplantation, most patients who are seropositive for HSV-1 excrete the virus from the oropharynx. The ability to isolate HSV declines with time. Administration of prophylactic acyclovir (or valacyclovir) to seropositive HSC transplant recipients has been shown to reduce mucositis and prevent HSV pneumonia (a rare condition reported almost exclusively in allogeneic HSC transplant recipients). Both esophagitis (usually due to HSV-1) and anogenital disease (commonly caused by HSV-2) may be prevented with acyclovir prophylaxis. **For further discussion, see Chap. 216.**

Varicella-Zoster Virus Reactivation of VZV manifests as herpes zoster and may occur within the first month but more commonly occurs several months after transplantation. Reactivation rates are ~40% for allogeneic HSC transplant recipients and 25% for autologous recipients. Localized zoster can spread rapidly in an immunosuppressed patient. Fortunately, disseminated disease can usually be controlled with high doses of acyclovir. Because of frequent dissemination among patients with skin lesions, acyclovir is given prophylactically in some centers to prevent severe disease. Low doses of acyclovir appear to be effective in preventing reactivation of VZV. However, acyclovir can also suppress the development of VZV-specific immunity. Thus, its administration for only 6 months after transplantation does not prevent zoster from occurring when treatment is stopped. Administration of low doses of acyclovir for an entire year after transplantation is effective and may eliminate most cases of posttransplantation zoster, even among cord-blood recipients. **For further discussion, see Chap. 217.**

Cytomegalovirus The onset of CMV disease (interstitial pneumonia, bone marrow suppression, graft failure, hepatitis/colitis) usually begins 30–90 days after HSC transplantation, when the granulocyte count is adequate but immunologic reconstitution has not occurred. CMV disease rarely develops earlier than 14 days after transplantation and may become evident as late as 4 months after the procedure. It is of greatest concern in the second month after transplantation, particularly in allogeneic HSC transplant recipients. In cases in which the donor marrow is depleted of T cells (to prevent GVHD or eliminate a T cell tumor) and in cord-blood recipients, the disease may manifest earlier. The use of alemtuzumab to prevent GVHD in nonmyeloablative transplantation has been associated with an increase in CMV disease. Patients who receive ganciclovir for prophylaxis, preemptive treatment, or treatment (see below) may develop recurrent CMV infection even later than 4 months after transplantation, as treatment appears to delay the development of the normal immune response to CMV infection. Although CMV disease may present as isolated fever, granulocytopenia, thrombocytopenia, or gastrointestinal disease, the foremost cause of death from CMV infection in the setting of HSC transplantation is pneumonia.

With the standard use of CMV-negative or filtered blood products, CMV infection should be a major risk in allogeneic transplantation only when the recipient is CMV-seropositive and the donor is CMV-seronegative. This situation is the reverse of that in solid organ transplant recipients. CMV reactivates from latent reservoirs present in the recipient at a time when donor T cells (especially cord-blood T cells) are too immature to control CMV replication. If the T cells from the donor have never encountered CMV and the recipient carries the virus, the patient is at maximal risk of severe disease. Reactivation disease or superinfection with another strain from the donor also can occur in CMV-positive recipients, but clinical manifestations are typically less severe, presumably because of CMV-specific memory in transplanted donor T cells. Most patients infected with CMV who undergo HSC transplantation excrete virus, with or without clinical findings. Serious CMV disease is much more common among allogeneic than autologous recipients and is often associated with GVHD. In addition to pneumonia and marrow suppression (and, less often, graft failure), manifestations of CMV disease in HSC transplant recipients include fever with or without arthralgias, myalgias, hepatitis, and esophagitis. CMV ulcerations occur in both the lower and the upper gastrointestinal tract, and it may be difficult to distinguish diarrhea due to GVHD from that due to CMV infection. The finding of CMV in

922 the liver of a patient with GVHD does not necessarily mean that CMV is responsible for hepatic enzyme abnormalities. It is interesting that the ocular and neurologic manifestations of CMV infections, which are common in patients with AIDS, are uncommon in patients who develop disease after transplantation.

Management of CMV disease in HSC transplant recipients includes strategies directed at prophylaxis, preemptive therapy (suppression of silent replication), and treatment of disease. Prophylaxis results in a lower incidence of disease at the cost of treating many patients who otherwise would not require therapy. Because of the high fatality rate associated with CMV pneumonia in these patients and the difficulty of early diagnosis of CMV infection, prophylactic IV ganciclovir (or oral valganciclovir) has been used in some centers and has been shown to prevent CMV disease during the period of maximal vulnerability (from engraftment to day 120 after transplantation). Ganciclovir also prevents HSV reactivation and reduces the risk of VZV reactivation; thus acyclovir prophylaxis should be discontinued when ganciclovir is administered. The foremost problem with the administration of ganciclovir relates to adverse effects, which include dose-related bone marrow suppression (thrombocytopenia, leukopenia, anemia, and pancytopenia). Because the frequency of CMV pneumonia is lower among autologous HSC transplant recipients (2–7%) than among allogeneic HSC transplant recipients (10–40%), prophylaxis in the former group will not become the rule until a less toxic oral antiviral agent becomes available. Several are under study.

Preemptive treatment of CMV—that is, initiation of therapy with drugs only after CMV is detected in blood by a nucleic acid amplification test (NAAT)—is used at most centers. To limit variability between tests, the World Health Organization (WHO) has developed an international reference standard for measurement of CMV load by NAAT-based assays. Because of toxic drug side effects (e.g., neutropenia and bone marrow suppression), the preemptive approach has supplanted prophylactic therapy; it has also replaced treatment of all seropositive (recipient and/or donor) HSC transplants with an antiviral agent (typically ganciclovir). A positive test (or increasing viral load) prompts the initiation of preemptive therapy with ganciclovir. Preemptive approaches that target patients who have quantitative NAAT evidence of CMV infection can still lead to unnecessary treatment of many individuals with drugs that have adverse effects on the basis of a laboratory test that is not highly predictive of disease; however, invasive disease, particularly in the form of pulmonary infection, is difficult to treat and is associated with high mortality rates. When prophylaxis or preemptive therapy is stopped, late manifestations of CMV replication may occur, although by then the HSC transplant patient is often equipped with improved graft function and is better able to combat disease. Cord-blood transplant recipients are especially vulnerable to disease caused by members of the human herpesvirus family, including CMV. Implementation of the WHO standard for CMV load measurement will facilitate large-scale comparative studies and thus the establishment of optimal guidelines for distinct patient subsets.

CMV pneumonia in HSC transplant recipients (unlike that in other clinical settings) is often treated with both IV immunoglobulin (IVIg) and ganciclovir. In patients who cannot tolerate ganciclovir, foscarnet is a useful alternative, although it may produce nephrotoxicity and electrolyte imbalance. When neither ganciclovir nor foscarnet is clinically tolerated, cidofovir can be used; however, its efficacy is less well established, and its side effects include nephrotoxicity. A lipid-conjugate form of cidofovir and an oral antiviral agent, maribavir, are in clinical trials. Case reports have suggested that the immunosuppressive agent leflunomide may be active in this setting, but controlled studies are lacking. Transfusion of CMV-specific T cells from the donor has decreased viral load in a small series of patients; this result suggests that immunotherapy (e.g., banked T cells) may play a role in the management of this disease in the future. **For further discussion, see Chap. 219.**

Human Herpesviruses 6 and 7 Human herpesvirus type 6 (HHV-6), the cause of roseola in children, is a ubiquitous herpesvirus that is reactivated (as determined by quantitative plasma PCR) in ~50% of HSC

transplant recipients 2–4 weeks after transplantation. Reactivation is more common among patients requiring glucocorticoids for GVHD and among those receiving second transplants. Reactivation of HHV-6, primarily type B, may be associated with delayed monocyte and platelet engraftment. Limbic encephalitis developing after transplantation has been associated with HHV-6 in cerebrospinal fluid (CSF). The causality of the association is not well defined; in several cases, plasma viremia was detected long before the onset of encephalitis. Nevertheless, most patients with encephalitis had very high viral loads in plasma at the time of CNS illness, and viral antigen has been detected in hippocampal astrocytes. HHV-6 DNA is sometimes found in lung samples after transplantation. However, its role in pneumonitis is unclear, as co-pathogens are frequently present. While HHV-6 is susceptible to foscarnet or cidofovir (and possibly to ganciclovir) *in vitro*, the efficacy of antiviral treatment has not been well studied. Little is known about the related herpesvirus HHV-7 or its role in posttransplantation infection. **For further discussion, see Chap. 219.**

Epstein-Barr Virus Primary EBV infection can be fatal to HSC transplant recipients; EBV reactivation can cause EBV-B cell lymphoproliferative disease (EBV-LPD), which may also be fatal to patients taking immunosuppressive drugs. Latent EBV infection of B cells leads to several interesting phenomena in HSC transplant recipients. The marrow ablation that occurs as part of the HSC transplantation procedure may sometimes eliminate latent EBV from the host. Infection can then be reacquired immediately after transplantation by transfer of infected donor B cells. Rarely, transplantation from a seronegative donor may result in a cure. The recipient is then at risk for a second primary infection.

EBV-LPD can develop in the recipient's B cells (if any survive marrow ablation) but is more likely to be a consequence of outgrowth of infected donor cells. Both lytic replication and latent replication of EBV are more likely during immunosuppression (e.g., they are associated with GVHD and the use of antibodies to T cells). Although less likely in autologous transplantation, reactivation can occur in T cell-depleted autologous recipients (e.g., patients being given antibodies to T cells for the treatment of a T cell lymphoma with marrow depletion). EBV-LPD, which can become apparent as early as 1–3 months after engraftment, can cause high fevers and cervical adenopathy resembling the symptoms of infectious mononucleosis but more commonly presents as an extranodal mass. The incidence of EBV-LPD among allogeneic HSC transplant recipients is 0.6–1%, which contrasts with figures of ~5% for renal transplant recipients and up to 20% for cardiac transplant patients. In all cases, EBV-LPD is more likely to occur with high-dose, prolonged immunosuppression, especially that caused by the use of antibodies to T cells, glucocorticoids, and calcineurin inhibitors (e.g., cyclosporine, tacrolimus). Cord-blood recipients constitute another high-risk group because of delayed T cell function. Ganciclovir, administered to preempt CMV disease, may reduce EBV lytic replication and thereby diminish the pool of B cells that can become newly infected and give rise to LPD. Increasing evidence indicates that replacement of calcineurin inhibitors with mTOR inhibitors (e.g., rapamycin) exerts an antiproliferative effect on EBV-infected B cells that decreases the likelihood of development of LPD or unrelated proliferative disorders associated with transplant-related immunosuppression.

PCR can be used to monitor EBV production after HSC transplantation. High or increasing viral loads predict an enhanced likelihood of EBV-LPD development and should prompt rapid reduction of immunosuppression and a search for nodal or extranodal disease. If reduction of immunosuppression does not have the desired effect, administration of a monoclonal antibody to CD20 (e.g., rituximab) for the treatment of B cell lymphomas that express this surface protein has elicited dramatic responses and currently constitutes first-line therapy for CD20-positive EBV-LPD. However, long-term suppression of new antibody responses accompanies therapy, and recurrences are not infrequent. Additional B cell-directed antibodies, including anti-CD22, are under study. The role of antiviral drugs is uncertain because no available agents have been documented to have activity

against the different forms of latent EBV infection. Diminishing lytic replication and virion production in these patients would theoretically produce a statistical decrease in the frequency of latent disease by decreasing the number of virions available to cause additional infection. In case reports and animal studies, ganciclovir and/or high-dose zidovudine, together with other agents, has been used to eradicate EBV-LPD and CNS lymphomas, another EBV-associated complication of transplantation. Both interferon and retinoic acid have been employed in the treatment of EBV-LPD, as has IVIg, but no large-scale prospective studies have assessed the efficacy of any of these agents. Several additional drugs are undergoing preclinical evaluation. Standard chemotherapeutic regimens are used if disease persists after reduction of immunosuppressive agents and administration of antibodies. EBV-specific T cells generated from the donor have been used experimentally to prevent and treat EBV-LPD in allogeneic recipients, and efforts are under way to increase the activity and specificity of ex vivo-generated T cells. **For further discussion, see Chap. 218.**

Human Herpesvirus 8 (KSHV) The EBV-related gammaherpesvirus KSHV, which is causally associated with Kaposi's sarcoma, primary effusion lymphoma, and multicentric Castleman's disease, has rarely resulted in disease in HSC transplant recipients, although some cases of virus-associated marrow aplasia have been reported in the peri-transplantation period. The relatively low seroprevalence of KSHV in the population and the limited duration of profound T cell suppression after HSC transplantation provide a plausible explanation for the currently low incidence of KSHV disease compared with that in recipients of solid organ transplants and patients with HIV infection. **For further discussion, see Chap. 219.**

Other (Non-Herpes) Viruses The diagnosis of pneumonia in HSC transplant recipients poses special problems. Because patients have undergone treatment with multiple chemotherapeutic agents and sometimes irradiation, their differential diagnosis should include—in addition to bacterial and fungal pneumonia—CMV pneumonitis, pneumonia of other viral etiologies, parasitic pneumonia, diffuse alveolar hemorrhage, and chemical- or radiation-associated pneumonitis. Since fungi and viruses (e.g., influenza A and B viruses, respiratory syncytial virus [RSV], parainfluenza virus [types 1–4], adenovirus, enterovirus, bocavirus, human metapneumovirus, coronavirus, and rhinovirus [increasingly detected by multiplex PCR]) also can cause pneumonia in this setting, it is important to obtain a specific diagnosis. Diagnostic modalities include Gram's stain, microbiologic culture, antigen testing, and—increasingly—multipathogen PCR and mass spectrometry assays. *M. tuberculosis* has been an uncommon cause of pneumonia among HSC transplant recipients in Western countries (accounting for <0.1–0.2% of cases) but is common in Hong Kong (5.5%) and in countries where the prevalence of tuberculosis is high. The recipient's exposure history is clearly critical in an assessment of posttransplantation infections.

Both RSV and parainfluenza viruses, particularly type 3, can cause severe or even fatal pneumonia in HSC transplant recipients. Infections with both of these agents sometimes occur as disastrous nosocomial epidemics. Therapy with palivizumab or ribavirin for RSV infection remains controversial. New agents, some host-directed, are under study. Influenza also occurs in HSC transplant recipients and generally mirrors the presence of infection in the community. Progression to pneumonia is more common when infection occurs early after transplantation and when the recipient is lymphopenic. The neuraminidase inhibitors oseltamivir (oral) and zanamivir (aerosolized) are active against both influenza A virus and influenza B virus and are a reasonable treatment option. Parenteral forms of neuraminidase inhibitors such as peramivir (intravenous) and several new oral agents remain in trial status. An important preventive measure is immunization of household members, hospital staff members, and other frequent contacts. Adenoviruses can be isolated from HSC transplant recipients at rates varying from 5% to ≥18%. Like CMV infection, adenovirus infection usually occurs in the first to third month after transplantation and is often asymptomatic, although pneumonia, hemorrhagic cystitis/nephritis, severe gastroenteritis with hemorrhage, and fatal

disseminated infection have been reported and may be strain-specific. A role for cidofovir therapy has been suggested, but the efficacy of this agent in adenovirus infection remains to be determined. Banked virus-specific T cell therapy is under study for adenovirus infection (as well as for CMV and EBV infections).

Although diverse respiratory viruses can sometimes cause severe pneumonia and respiratory failure in HSC transplant recipients, mild or even asymptomatic infection may be more common. For example, rhinoviruses and coronaviruses are frequent co-pathogens in HSC transplant recipients; however, whether they independently contribute to significant pulmonary infection is not known. At present, the overall contribution of these viral respiratory pathogens to the burden of lower respiratory tract disease in HSC transplant recipients requires further study. Infections with parvovirus B19 (presenting as anemia or occasionally as pancytopenia) and disseminated enteroviruses (sometimes fatal) can occur. Parvovirus B19 infection can be treated with IVIg (**Chap. 221**).

Rotaviruses are a cause of gastroenteritis in HSC transplant recipients, more frequently in children. Norovirus is a common cause of vomiting and diarrhea, and symptoms can be prolonged in HSC recipients. The polyomavirus BK virus is found at high titers in the urine of patients who are profoundly immunosuppressed. BK viraemia may be associated with hemorrhagic cystitis in these patients. In contrast to its incidence among patients with impaired T cell function due to AIDS (4–5%), progressive multifocal leukoencephalopathy caused by the related JC virus is relatively rare among HSC transplant recipients (**Chap. 164**). When transmitted by mosquitoes or by blood transfusion, WNV can cause encephalitis and death after HSC transplantation.

INFECTIONS IN SOLID ORGAN TRANSPLANT RECIPIENTS

Rates of morbidity and mortality among recipients of solid organ transplants (SOTs) are reduced by the use of effective antibiotics. The organisms that cause acute infections in recipients of SOTs are different from those that infect HSC transplant recipients because SOT recipients do not go through a period of neutropenia. As the transplantation procedure involves major surgery, however, SOT recipients are subject to infections at anastomotic sites and to wound infections. Compared with HSC transplant recipients, SOT patients are immunosuppressed for longer periods (often permanently). Thus they are susceptible to many of the same organisms as patients with chronically impaired T cell immunity (**Chap. 104, especially Table 104-1**). Moreover, the persistent HLA mismatch between recipient immune cells (e.g., effector T cells) and the donor organ (allograft) places the organ at permanently increased risk of infection.

During the early period (<1 month after transplantation; **Table 169-4**), infections are most commonly caused by extracellular bacteria (staphylococci, streptococci, enterococci, and *E. coli* and other gram-negative organisms, including nosocomial organisms with broad antibiotic resistance), which often originate in surgical wound or anastomotic sites. The type of transplant largely determines the spectrum of infection. In subsequent weeks, the consequences of the administration of agents that suppress cell-mediated immunity become apparent, and acquisition—or, more commonly, reactivation—of viruses, mycobacteria, endemic fungi, and parasites (from the recipient or from the transplanted organ) can occur. CMV infection is often a problem, particularly in the first 6 months after transplantation, and may present as severe systemic disease or as infection of the transplanted organ. HHV-6 reactivation (assessed by plasma PCR) occurs within the first 2–4 weeks after transplantation and may be associated with fever, leukopenia, and very rare cases of encephalitis. Data suggest that replication of HHV-6 and HHV-7 may exacerbate CMV-induced disease. CMV is associated not only with generalized immunosuppression but also with organ-specific, rejection-related syndromes: glomerulopathy in kidney transplant recipients, bronchiolitis obliterans in lung transplant recipients, vasculopathy in heart transplant recipients, and the vanishing bile duct syndrome in liver transplant recipients. A complex interplay between increased CMV

TABLE 169-4 COMMON INFECTIONS AFTER SOLID ORGAN TRANSPLANTATION, BY SITE OF INFECTION

Infected Site	Period after Transplantation		
	Early (<1 Month)	Middle (1–4 Months)	Late (>6 Months)
Donor organ	Bacterial and fungal infections of the graft, anastomotic site, and surgical wound	CMV infection	EBV infection (may present in allograft organ)
Systemic	Bacteremia and candidemia (often resulting from central venous catheter colonization)	CMV infection (fever, bone marrow suppression)	CMV infection, especially in patients given early posttransplantation prophylaxis; EBV proliferative syndromes (may occur in donor organs)
Lung	Bacterial aspiration pneumonia with prevalent nosocomial organisms associated with intubation and sedation (highest risk in lung transplantation)	<i>Pneumocystis</i> infection; CMV pneumonia (highest risk in lung transplantation); <i>Aspergillus</i> infection (highest risk in lung transplantation)	<i>Pneumocystis</i> infection; granulomatous lung diseases (nocardial and reactivated fungal and mycobacterial diseases)
Kidney	Bacterial and fungal (<i>Candida</i>) infections (cystitis, pyelonephritis) associated with urinary tract catheters (highest risk in kidney transplantation)	Kidney transplantation: BK virus infection (associated with nephropathy); JC virus infection	Kidney transplantation: bacterial infections (late urinary tract infections, usually not associated with bacteremia); BK virus infection (nephropathy, graft failure, generalized vasculopathy)
Liver and biliary tract	Cholangitis	CMV hepatitis	CMV hepatitis
Heart		<i>Toxoplasma gondii</i> infection (highest risk in heart transplantation); endocarditis (<i>Aspergillus</i> and gram-negative organisms more common than in general population)	<i>T. gondii</i> (highest risk in heart transplantation)
Gastrointestinal tract	Peritonitis, especially after liver transplantation	Colitis secondary to <i>Clostridium difficile</i> infection (risk can persist)	Colitis secondary to <i>C. difficile</i> infection (risk can persist)
Central nervous system		<i>Listeria</i> infection (meningitis); <i>T. gondii</i> infection; CMV infection	Listerial meningitis; cryptococcal meningitis; nocardial abscess; JC virus–associated PML

Abbreviations: CMV, cytomegalovirus; EBV, Epstein-Barr virus; PML, progressive multifocal leukoencephalopathy.

replication and enhanced graft rejection is well established: elevated immunosuppression leads to increased CMV replication, which is associated with graft rejection. For this reason, considerable attention has been focused on the diagnosis, prophylaxis, and treatment of CMV infection in SOT recipients. Early transmission of WNV to transplant recipients from a donated organ or transfused blood has been reported; however, the risk of WNV acquisition has been reduced by implementation of screening procedures. In rare instances, rabies virus and lymphocytic choriomeningitis virus also have been acutely transmitted in this setting; although accompanied by distinct clinical syndromes, both viral infections have resulted in fatal encephalitis. As screening for unusual viruses is not routine, only vigilant assessment of the prospective donor is likely to prevent the use of an infected organ.

Beyond 6 months after transplantation, infections characteristic of patients with defects in cell-mediated immunity—e.g., infections with *Listeria*, *Nocardia*, *Rhodococcus*, mycobacteria, various fungi, and other intracellular pathogens—may be a problem. International patients and global travelers may experience reactivation of dormant infections with trypanosomes, *Leishmania*, *Plasmodium*, *Strongyloides*, and other parasites. Reactivation of latent *M. tuberculosis* infection, while rare in Western nations, is far more common among persons from developing countries. The recipient is typically the source, although reactivation and spread from the donor organ can occur. While pulmonary disease remains most common, atypical sites can be involved and mortality rates can be high (up to 30%). Vigilance, prophylaxis/preemptive therapy (when indicated), and rapid diagnosis and treatment of infections can be lifesaving in SOT recipients, who, unlike most HSC transplant recipients, continue to be immunosuppressed.

SOT recipients are susceptible to EBV-LPD from as early as 2 months to many years after transplantation. The prevalence of this complication is increased by potent and prolonged use of T cell-suppressive drugs. Decreasing the degree of immunosuppression may in some cases reverse the condition. Among SOT patients, those with heart and lung transplants—who receive the most intensive immunosuppressive regimens—are most likely to develop EBV-LPD, particularly in the lungs. Although the disease usually originates in recipient B cells, several cases of donor origin, particularly in the transplanted

organ, have been noted. High organ-specific content of B lymphoid tissues (e.g., bronchus-associated lymphoid tissue in the lung), anatomic factors (e.g., lack of access of host T cells to the transplanted organ because of disturbed lymphatics), and differences in major histocompatibility loci between the host T cells and the organ (e.g., lack of cell migration or lack of effective T cell/macrophage/dendritic cell cooperation) may result in defective elimination of EBV-infected B cells. SOT recipients are also highly susceptible to the development of Kaposi's sarcoma and, less frequently, to the B cell-proliferative disorders associated with KSHV, such as primary effusion lymphoma and multicentric Castleman's disease. Kaposi's sarcoma is 550–1000 times more common among SOT recipients than in the general population, can develop very rapidly after transplantation, and can also occur in the allograft. However, because the seroprevalence of KSHV is very low in Western countries, Kaposi's sarcoma is not common. Recipients (or donors) from Iceland, the Middle East, Mediterranean countries, and Africa are at highest risk of disease. Data suggest that a switch of immunosuppressive agents—from calcineurin inhibitors (cyclosporine, tacrolimus) to mTor pathway-active agents (sirolimus, everolimus)—after adequate wound healing may significantly reduce the likelihood of development of Kaposi's sarcoma and perhaps of EBV-LPD and certain other posttransplantation malignancies.

KIDNEY TRANSPLANTATION

See Table 169-4.

Early Infections Bacteria often cause infections that develop in the period immediately after kidney transplantation. There is a role for perioperative antibiotic prophylaxis, and many centers give cephalosporins to decrease the risk of postoperative complications. Urinary tract infections developing soon after transplantation are usually related to anatomic alterations resulting from surgery. Such early infections may require prolonged treatment (e.g., 6 weeks of antibiotic administration for pyelonephritis). Urinary tract infections that occur >6 months after transplantation may be treated for shorter periods because they do not seem to be associated with the high rate of pyelonephritis or relapse seen with infections that occur during the first 3 months.

TABLE 169-5 PROPHYLACTIC REGIMENS COMMONLY USED TO DECREASE RISK OF INFECTION IN TRANSPLANT RECIPIENTS^a

Risk Factor	Organism	Prophylactic Drug	Examination(s) ^b
Travel to or residence in area with known risk of endemic fungal infection	<i>Histoplasma</i> , <i>Blastomyces</i> , <i>Coccidioides</i>	Triazoles considered in context of clinical and laboratory assessment	Chest radiography, antigen testing, serology
Latent herpesviruses	HSV, VZV, CMV, EBV	Acyclovir after HSC transplantation to prevent HSV and VZV infection or reactivation; ganciclovir to prevent CMV infection, with possible effect on EBV/KSHV/HHV-6 infections in some settings	Serologic tests for HSV, VZV, CMV, HHV-6, EBV, KSHV; PCR
Latent fungi and parasites	<i>Pneumocystis jirovecii</i> , <i>Toxoplasma gondii</i>	Trimethoprim-sulfamethoxazole (or alternatives)	Serologic test for <i>Toxoplasma</i>
History of exposure to active or latent tuberculosis	<i>Mycobacterium tuberculosis</i>	Isoniazid in patients with recent seroconversion or positive chest imaging and/or no previous treatment	Chest imaging; TST and/or cell-based assay

^aFor information on latent infection with hepatitis B or C virus, see [Chap. 362](#). ^bSerologic examination, tuberculin skin test, and interferon assays may be less reliable after transplantation.

Abbreviations: CMV, cytomegalovirus; EBV, Epstein-Barr virus; HHV-6, human herpesvirus type 6; HSC, hematopoietic stem cell; HSV, herpes simplex virus; KSHV, Kaposi's sarcoma-associated herpesvirus; PCR, polymerase chain reaction; TST, tuberculin skin test; VZV, varicella-zoster virus.

Prophylaxis with TMP-SMX for the first 4–6 months after transplantation decreases the incidence of early and middle-period infections (see below, [Table 169-4](#), and [Table 169-5](#)).

Middle-Period Infections Because of continuing immunosuppression, kidney transplant recipients are predisposed to lung infections characteristic of those in patients with T cell deficiency (i.e., infections with intracellular bacteria, mycobacteria, nocardiae, fungi, viruses, and parasites). A high mortality rate associated with *Legionella pneumophila* infection ([Chap. 184](#)) led to the closing of renal transplant units in hospitals with endemic legionellosis.

About 50% of all renal transplant recipients presenting with fever 1–4 months after transplantation have evidence of CMV disease; CMV itself accounts for the fever in more than two-thirds of cases and thus is the predominant pathogen during this period. CMV infection ([Chap. 219](#)) may also present as arthralgias, myalgias, or organ-specific symptoms. During this period, this infection may represent primary disease (in the case of a seronegative recipient of a kidney from a seropositive donor) or may represent reactivation disease or superinfection. Patients may have atypical lymphocytosis. Unlike immunocompetent patients, however, they rarely have lymphadenopathy or splenomegaly. Therefore, clinical suspicion and laboratory confirmation are necessary for diagnosis. The clinical syndrome may be accompanied by bone marrow suppression (particularly leukopenia). CMV also causes glomerulopathy and is associated with an increased incidence of other opportunistic infections. Because of the frequency and severity of disease, a considerable effort has been made to prevent and treat CMV infection in renal transplant recipients. An immune globulin preparation enriched with antibodies to CMV was used by many centers in the past in an effort to protect the group at highest risk for severe infection (seronegative recipients of seropositive kidneys). However, with the development of effective oral antiviral agents, CMV immune globulin is no longer used. Ganciclovir (or valganciclovir) is beneficial for prophylaxis (when indicated) and for the treatment of serious CMV disease. The availability of valganciclovir has allowed most centers to move to oral prophylaxis for transplant recipients. Infection with the other herpesviruses may become evident within 6 months after transplantation or later. Early after transplantation, HSV may cause either oral or anogenital lesions that are usually responsive to acyclovir. Large ulcerating lesions in the anogenital area may lead to bladder and rectal dysfunction and may predispose the patient to bacterial infection. VZV may cause fatal disseminated infection in nonimmune kidney transplant recipients, but in immune patients reactivation zoster usually does not disseminate outside the dermatome; thus disseminated VZV infection is a less fearsome complication in kidney transplantation than in HSC transplantation. HHV-6 reactivation may take place and (although usually asymptomatic) may be associated with fever, rash, marrow suppression, or rare instances of renal impairment, hepatitis, colitis, or encephalitis.

EBV disease is more serious; it may present as an extranodal proliferation of B cells that invade the CNS, nasopharynx, liver, small bowel, heart, and other organs, including the transplanted kidney. The disease is diagnosed by the finding of a mass of proliferating EBV-positive B cells. The incidence of EBV-LPD is elevated among patients who acquire EBV infection from the donor and among patients given high doses of cyclosporine, tacrolimus, glucocorticoids, and anti-T cell antibodies. Disease may regress once immunocompetence is restored. KSHV infection can be transmitted with the donor kidney and result in development of Kaposi's sarcoma, although it more often represents reactivation of latent infection of the recipient. Kaposi's sarcoma often appears within 1 year after transplantation, although the time of onset ranges widely (1 month to ~20 years). Avoidance of immunosuppressive agents that inhibit calcineurin has been associated with less Kaposi's sarcoma, less EBV disease, and even less CMV replication. The use of rapamycin (sirolimus) has independently led to regression of Kaposi's sarcoma.

The papovaviruses BK virus and JC virus (polyomavirus hominis types 1 and 2) have been cultured from the urine of kidney transplant recipients (as they have from that of HSC transplant recipients) in the setting of profound immunosuppression. High levels of BK virus replication detected by PCR in urine and blood are predictive of pathology, especially in the setting of renal transplantation. JC virus may rarely cause similar disease in kidney transplantation. Urinary excretion of BK virus and BK viremia are associated with the development of ureteral strictures, polyomavirus-associated nephropathy (1–10% of renal transplant recipients), and (less commonly) generalized vasculopathy. Timely detection and early reduction of immunosuppression are critical and can reduce rates of graft loss related to polyomavirus-associated nephropathy from 90% to 10–30%. Therapeutic responses to IVIg, quinolones, leflunomide, and cidofovir have been reported, but the efficacy of these agents has not been substantiated through adequate clinical study. Most centers approach the problem by reducing immunosuppression in an effort to enhance host immunity and decrease viral titers. JC virus is associated with rare cases of progressive multifocal leukoencephalopathy. Adenoviruses may persist and cause hemorrhagic nephritis/cystitis with continued immunosuppression in these patients, but disseminated disease like that seen in HSC transplant recipients is much less common.

Kidney transplant recipients are also subject to infections with other intracellular organisms. These patients may develop pulmonary infections with *Mycobacterium*, *Aspergillus*, and *Mucor* species as well as infections with other pathogens in which the T cell/macrophage axis plays an important role. *L. monocytogenes* is a common cause of bacteremia ≥ 1 month after renal transplantation and should be seriously considered in renal transplant recipients presenting with fever and headache. Kidney transplant recipients may develop *Salmonella* bacteremia, which can lead to endovascular infections and require prolonged therapy. Pulmonary infections with *Pneumocystis* are common unless the patient is maintained on TMP-SMX prophylaxis.

926 Acute interstitial nephritis caused by TMP-SMX is rare. However, because transient increases in creatinine (artificial) and hyperkalemia (manageable) can occur, early discontinuation of prophylaxis, especially after kidney transplantation, is recommended by some groups. Although additional monitoring is indicated, the benefits of TMP-SMX in kidney transplant recipients may outweigh the risks; otherwise, second-line prophylactic agents should be used. *Nocardia* infection (Chap. 199) may present in the skin, bones, and lungs or in the CNS, where it usually takes the form of single or multiple brain abscesses. Nocardiosis generally occurs ≥ 1 month after transplantation and may follow immunosuppressive treatment for an episode of rejection. Pulmonary manifestations most commonly consist of localized disease with or without cavities, but the disease may be disseminated. The diagnosis is made by culture of the organism from sputum or from the involved nodule. As it is for *P. jirovecii* infection, prophylaxis with TMP-SMX is often efficacious in the prevention of nocardiosis.

Toxoplasmosis can occur in seropositive patients but is less common than in other transplantation settings, usually developing in the first few months after kidney transplantation. Again, TMP-SMX is helpful in prevention. In endemic areas, histoplasmosis, coccidioidomycosis, and blastomycosis may cause pulmonary infiltrates or disseminated disease.

Late Infections Late infections (>6 months after kidney transplantation) may involve the CNS and include CMV retinitis as well as other CNS manifestations of CMV disease. Patients (particularly those whose immunosuppression has been increased) are at risk for subacute meningitis due to *Cryptococcus neoformans*. Cryptococcal disease may present in an insidious manner (sometimes as a skin infection before the development of clear CNS findings). *Listeria* meningitis may have an acute presentation and requires prompt therapy to avoid a fatal outcome. TMP-SMX prophylaxis may reduce the frequency of *Listeria* infections.

Patients who continue to take glucocorticoids are predisposed to ongoing infection. “Transplant elbow,” a recurrent bacterial infection in and around the elbow that is thought to result from a combination of poor tensile strength of the skin of steroid-treated patients and steroid-induced proximal myopathy, requires patients to push themselves up with their elbows to get out of chairs. Bouts of cellulitis (usually caused by *S. aureus*) recur until patients are provided with elbow protection.

Kidney transplant recipients are susceptible to invasive fungal infections, including those due to *Aspergillus* and *Rhizopus*, which may present as superficial lesions before dissemination. Mycobacterial infection (particularly that with *Mycobacterium marinum*) can be diagnosed by skin examination. Infection with *Prototheca wickerhamii* (an achlorophyllic alga) has been diagnosed by skin biopsy. Warts caused by human papillomaviruses (HPVs) are a late consequence of persistent immunosuppression; imiquimod or other forms of local therapy are usually satisfactory. Merkel cell carcinoma, a rare and aggressive neuroendocrine skin tumor whose frequency is increased fivefold in elderly SOT (especially kidney) recipients, is causally linked to a novel polyomavirus, Merkel cell polyomavirus.

Notably, although BK virus replication and virus-associated disease can be detected far earlier, polyomavirus-associated nephropathy is clinically diagnosed in a median of ~ 300 days and thus qualifies as a late-onset disease. With the establishment of better screening procedures (e.g., urine cytology, urine nucleic acid load, plasma PCR), disease onset is being detected earlier (see “Middle-Period Infections,” above) and preemptive strategies (decrease or modification of immunosuppression) are being instituted more promptly, as the efficacy of antiviral therapy is not well established.

HEART TRANSPLANTATION

Early Infections Sternal wound infection and mediastinitis are early complications of heart transplantation. An indolent course is common, with fever or a mildly elevated white blood cell count preceding the development of site tenderness or drainage. Clinical suspicion based on evidence of sternal instability and failure to heal may lead to

the diagnosis. Common microbial residents of the skin (e.g., *S. aureus*, including methicillin-resistant strains, and *Staphylococcus epidermidis*) as well as gram-negative organisms (e.g., *Pseudomonas aeruginosa*) and fungi (e.g., *Candida*) are often involved. In rare cases, mediastinitis in heart transplant recipients can also be due to *Mycoplasma hominis* (Chap. 212); since this organism requires an anaerobic environment for growth and may be difficult to see on conventional medium, the laboratory should be alerted that its involvement is suspected. *M. hominis* mediastinitis has been cured with a combination of surgical debridement (sometimes requiring muscle-flap placement) and the administration of clindamycin and tetracycline. Organisms associated with mediastinitis may sometimes be cultured from pericardial fluid.

Middle-Period Infections *T. gondii* (Chap. 253) residing in the heart of a seropositive donor may be transmitted to a seronegative recipient. Thus serologic screening for *T. gondii* infection is important before and in the months after cardiac transplantation. Rarely, active disease can be introduced at the time of transplantation. The overall incidence of toxoplasmosis is so high in the setting of heart transplantation that some prophylaxis is always warranted. Although alternatives are available, the most frequently used agent is TMP-SMX, which prevents infection with *Pneumocystis* as well as with *Nocardia* and several other bacterial pathogens. CMV also has been transmitted by heart transplantation. *Toxoplasma*, *Nocardia*, and *Aspergillus* can cause CNS infections. *L. monocytogenes* meningitis should be considered in heart transplant recipients with fever and headache.

CMV infection is associated with poor outcomes after heart transplantation. The virus is usually detected 1–2 months after transplantation, causes early signs and laboratory abnormalities (usually fever and atypical lymphocytosis or leukopenia and thrombocytopenia) at 2–3 months, and can produce severe disease (e.g., pneumonia) at 3–4 months. An interesting observation is that seropositive recipients usually develop viremia faster than patients whose primary CMV infection is a consequence of transplantation. Between 40% and 70% of patients develop symptomatic CMV disease in the form of (1) CMV pneumonia, the form most likely to be fatal; (2) CMV esophagitis and gastritis, sometimes accompanied by abdominal pain with or without ulcerations and bleeding; and (3) the CMV syndrome, consisting of CMV in the bloodstream along with fever, leukopenia, thrombocytopenia, and hepatic enzyme abnormalities. Ganciclovir is efficacious in the treatment of CMV infection; prophylaxis with ganciclovir or possibly with other antiviral agents, as described for renal transplantation, may reduce the overall incidence of CMV-related disease.

Late Infections EBV infection usually presents as a lymphoma-like proliferation of B cells late after heart transplantation, particularly in patients maintained on intense immunosuppressive therapy. A subset of heart and heart-lung transplant recipients may develop early fulminant EBV-LPD (within 2 months). Treatment includes the reduction of immunosuppression (if possible), the use of glucocorticoid and calcineurin inhibitor-sparing regimens, and the consideration of therapy with anti-B cell antibodies (rituximab and possibly others). Immunomodulatory and antiviral agents continue to be studied. Ganciclovir prophylaxis for CMV disease may indirectly reduce the risk of EBV-LPD through reduced spread of replicating EBV to naïve B cells. Aggressive chemotherapy is a last resort, as discussed earlier for HSC transplant recipients. KSHV-associated disease, including Kaposi’s sarcoma and primary effusion lymphoma, has been reported in heart transplant recipients. GVHD prophylaxis with sirolimus may decrease the risk of both rejection and outgrowth of KSHV-infected cells. Antitumor therapy is discussed in Chap. 103e. Prophylaxis for *Pneumocystis* infection is required for these patients (see “Lung Transplantation, Late Infections,” below).

LUNG TRANSPLANTATION

Early Infections It is not surprising that lung transplant recipients are predisposed to the development of pneumonia. The combination of ischemia and the resulting mucosal damage, together with accompanying denervation and lack of lymphatic drainage, probably contributes to the high rate of pneumonia (66% in one series). The

prophylactic use of high doses of broad-spectrum antibiotics for the first 3–4 days after surgery may decrease the incidence of pneumonia. Gram-negative pathogens (Enterobacteriaceae and *Pseudomonas* species) are troublesome in the first 2 weeks after surgery (the period of maximal vulnerability). Pneumonia can also be caused by *Candida* (possibly as a result of colonization of the donor lung), *Aspergillus*, and *Cryptococcus*. Many centers use antifungal prophylaxis (typically fluconazole or liposomal amphotericin B) for the first 1–2 weeks.

Mediastinitis may occur at an even higher rate among lung transplant recipients than among heart transplant recipients and most commonly develops within 2 weeks of surgery. In the absence of prophylaxis, pneumonitis due to CMV (which may be transmitted as a consequence of transplantation) usually presents between 2 weeks and 3 months after surgery, with primary disease occurring later than reactivation disease.

Middle-Period Infections The incidence of CMV infection, either reactivated or primary, is 75–100% if either the donor or the recipient is seropositive for CMV. CMV-induced disease after solid organ transplantation appears to be most severe in recipients of lung and heart-lung transplants. Whether this severity relates to the mismatch in lung antigen presentation and host immune cells or is attributable to nonimmunologic factors is not known. More than half of lung transplant recipients with symptomatic CMV disease have pneumonia. Difficulty in distinguishing the radiographic picture of CMV infection from that of other infections or from organ rejection further complicates therapy. CMV can also cause bronchiolitis obliterans in lung transplants. The development of pneumonitis related to HSV has led to the prophylactic use of acyclovir. Such prophylaxis may also decrease rates of CMV disease, but ganciclovir is more active against CMV and is also active against HSV. The prophylaxis of CMV infection with IV ganciclovir—or increasingly with valganciclovir, the oral alternative—is recommended for lung transplant recipients. Antiviral alternatives are discussed in the earlier section on HSC transplantation. Although the overall incidence of serious disease is decreased during prophylaxis, late disease may occur when prophylaxis is stopped—a pattern observed increasingly in recent years. With recovery from peritransplantation complications and, in many cases, a decrease in immunosuppression, the recipient is often better equipped to combat late infection.

Late Infections The incidence of *Pneumocystis* infection (which may present with a paucity of findings) is high among lung and heart-lung transplant recipients. Some form of prophylaxis for *Pneumocystis* pneumonia is indicated in all organ transplant situations (Table 169-5). Prophylaxis with TMP-SMX for 12 months after transplantation may be sufficient to prevent *Pneumocystis* disease in patients whose immunosuppression is not increased.

As in other transplant recipients, EBV infection in lung and heart-lung recipients may cause either a mononucleosis-like syndrome or EBV-LPD. The tendency of the B cell blasts to present in the lung appears to be greater after lung transplantation than after the transplantation of other organs, possibly because of a rich source of B cells in bronchus-associated lymphoid tissue. Reduction of immunosuppression and switching of regimens, as discussed in earlier sections, cause remission in some cases, but mTOR inhibitors such as rapamycin may contribute to lung toxicity. Airway compression can be fatal, and rapid intervention may therefore become necessary. The approach to EBV-LPD is similar to that described in other sections.

LIVER TRANSPLANTATION

Early Infections As in other transplantation settings, early bacterial infections are a major problem after liver transplantation. Many centers administer systemic broad-spectrum antibiotics for the first 24 h or sometimes longer after surgery, even in the absence of documented infection. However, despite prophylaxis, infectious complications are common and correlate with the duration of the surgical procedure and the type of biliary drainage. An operation lasting >12 h is associated with an increased likelihood of infection. Patients who have a choledochojejunostomy with drainage of the biliary duct to a Roux-en-Y jejunal

bowel loop have more fungal infections than those whose bile is drained via anastomosis of the donor common bile duct to the recipient common bile duct. Overall, liver transplant patients have a high incidence of fungal infections, and the occurrence of fungal (often candidal) infection in the setting of choledochojejunostomy correlates with retransplantation, elevated creatinine levels, long procedures, transfusion of >40 units of blood, reoperation, preoperative use of glucocorticoids, prolonged treatment with antibacterial agents, and fungal colonization 2 days before and 3 days after surgery. Many centers give antifungal agents prophylactically in this setting.

Peritonitis and intraabdominal abscesses are common complications of liver transplantation. Bacterial peritonitis or localized abscesses may result from biliary leaks. Early leaks are especially common with live-donor liver transplants. Peritonitis in liver transplant recipients is often polymicrobial, frequently involving enterococci, aerobic gram-negative bacteria, staphylococci, anaerobes, or *Candida* and sometimes involving other invasive fungi. Only one-third of patients with intraabdominal abscesses have bacteremia. Abscesses within the first month after surgery may occur not only in and around the liver but also in the spleen, pericolic area, and pelvis. Treatment includes antibiotic administration and drainage as necessary.

Middle-Period Infections The development of postsurgical biliary stricture predisposes patients to cholangitis. The incidence of strictures is increased in live-donor liver transplantation. Transplant recipients who develop cholangitis may have high spiking fevers and rigors but often lack the characteristic signs and symptoms of classic cholangitis, including abdominal pain and jaundice. Although these findings may suggest graft rejection, rejection is typically accompanied by marked elevation of liver function enzymes. In contrast, in cholangitis in transplant recipients, results of liver function tests (with the possible exception of alkaline phosphatase levels) are often within the normal range. Definitive diagnosis of cholangitis in liver transplant recipients requires demonstration of aggregated neutrophils in bile duct biopsy specimens. Unfortunately, invasive studies of the biliary tract (either T-tube cholangiography or endoscopic retrograde cholangiopancreatography) may themselves lead to cholangitis. For this reason, many clinicians recommend an empirical trial of therapy with antibiotics covering gram-negative organisms and anaerobes before these procedures are undertaken as well as antibiotic coverage if procedures are eventually performed.

Reactivation of viral hepatitis is a common complication of liver transplantation (Chap. 360). Recurrent hepatitis B and C infections, for which transplantation may be performed, are problematic. To prevent hepatitis B virus reinfection, prophylaxis with an optimal antiviral agent or combination of agents (lamivudine, adefovir, entecavir) and hepatitis B immune globulin is currently recommended, although the optimal dose, route, and duration of therapy remain controversial. Success in preventing reinfection with hepatitis B virus has increased in recent years. Complications related to hepatitis C infection are the most common reason for liver transplantation in the United States. Reinfection of the graft with hepatitis C virus occurs in all patients, with a variable time frame. Studies of aggressive pretransplantation treatment of selected recipients with antiviral agents and prophylactic/preemptive regimens are ongoing. However, early posttransplantation initiation of treatment for histologically documented disease with the classic combination of ribavirin and pegylated interferon has produced sustained responses at rates in the range of 25–40%. Several protease and polymerase inhibitors that block production of hepatitis C virus as well as regimens that spare interferon and a monoclonal antibody to the virus are undergoing preclinical and clinical trials for prevention and or control of infection after transplantation (Chap. 360).

As in other transplantation settings, reactivation disease with herpesviruses is common (Table 169-3). Herpesviruses can be transmitted in donor organs. Although CMV hepatitis occurs in ~4% of liver transplant recipients, it is usually not so severe as to require retransplantation. Without prophylaxis, CMV disease develops in the majority of seronegative recipients of organs from CMV-positive donors, but fatality rates are lower among liver transplant recipients than among lung or heart-lung transplant recipients. Disease due to CMV can also

928 be associated with the vanishing bile duct syndrome after liver transplantation. Patients respond to treatment with ganciclovir; prophylaxis with oral forms of ganciclovir or high-dose acyclovir may decrease the frequency of disease. A role for HHV-6 reactivation in early posttransplantation fever and leukopenia has been proposed, although the more severe sequelae described in HSC transplantation are unusual. HHV-6 and HHV-7 appear to exacerbate CMV disease in this setting. EBV-LPD after liver transplantation shows a propensity for involvement of the liver, and such disease may be of donor origin. See previous sections for discussion of EBV infections in solid organ transplantation.

PANCREAS TRANSPLANTATION

Pancreas transplantation is most frequently performed together with or after kidney transplantation, although it may be performed alone. Transplantation of the pancreas can be complicated by early bacterial and yeast infections. Most pancreatic transplants are drained into the bowel, and the rest are drained into the bladder. A cuff of duodenum is used in the anastomosis between the pancreatic graft and either the gut or the bladder. Bowel drainage poses a risk of early intraabdominal and allograft infections with enteric bacteria and yeasts. These infections can result in loss of the graft. Bladder drainage causes a high rate of urinary tract infection and sterile cystitis; however, such infection can usually be cured with appropriate antimicrobial agents. In both procedures, prophylactic antimicrobial agents are commonly used at the time of surgery. Aggressive immunosuppression, especially when the patient receives a kidney and a pancreas from different donors, is associated with late-onset systemic fungal and viral infections; thus many centers administer an antifungal drug and an antiviral agent (ganciclovir or a congener) for extended prophylaxis.

Issues related to the development of CMV infection, EBV-LPD, and infections with opportunistic pathogens in patients receiving a pancreatic transplant are similar to those in other SOT recipients.

COMPOSITE TISSUE TRANSPLANTATION

Composite tissue allotransplantation (CTA) is a new field in which, rather than a single organ, multiple tissue types composing a major body part are transplanted. The sites involved have included an upper extremity, the face, the trachea, the knee, and the abdominal wall. The numbers of recipients are limited. The different procedures and the associated infectious complications vary. Nevertheless, some early trends related to infectious complications have become apparent, as very intense and prolonged immunosuppression is typically required to prevent rejection. For example, in the early postoperative period, bacterial infections are especially frequent in facial transplant recipients. Perioperative prophylaxis is tailored to the organisms likely to complicate the different procedures. As in SOT recipients, complicated CMV infections have been observed in several CTA settings, particularly when the recipient is seronegative and the donor is seropositive. In some patients, anti-CMV immune globulin in addition to ganciclovir (as used in HSC transplant recipients with CMV pneumonia) was needed to control disease, and ganciclovir resistance requiring alternative therapies developed in several patients. Infectious complications from reactivation of other members of the human herpesvirus family and other latent viruses also caused significant morbidity, as discussed for SOT recipients. Prophylaxis for CMV infection, *P. jirovecii* infection, toxoplasmosis, and fungal infection is administered for several months on the basis of the limited studies available.

MISCELLANEOUS INFECTIONS IN SOLID ORGAN TRANSPLANTATION

Indwelling IV Catheter Infections The prolonged use of indwelling IV catheters for administration of medications, blood products, and nutrition is common in diverse transplantation settings and poses a risk of local and bloodstream infections. Exit-site infection is most commonly caused by staphylococcal species. Bloodstream infection most frequently develops within 1 week of catheter placement or in patients who become neutropenic. Coagulase-negative staphylococci are the most common isolates from blood. Although infective endocarditis in HSC transplant recipients is uncommon, the incidence of endocarditis in SOT recipients has been estimated to be as high as 1%, and this infection is associated with excessive high mortality in this

population. Although staphylococci predominate, the involvement of fungal and gram-negative organisms may be more common than in the general population.

For further discussion of differential diagnosis and therapeutic options, see Chap. 104.

Tuberculosis The incidence of tuberculosis within the first 12 months after solid organ transplantation is greater than that observed after HSC transplantation (0.23–0.79%) and ranges broadly worldwide (1.2–15%), reflecting the prevalence of tuberculosis in local populations. Lesions suggesting prior tuberculosis on chest radiography, older age, diabetes, chronic liver disease, GVHD, and intense immunosuppression are predictive of tuberculosis reactivation and development of disseminated disease in a host with latent disease. Tuberculosis has rarely been transmitted from the donor organ. In contrast to the low mortality rate among HSC transplant recipients, mortality rates among SOT recipients are reported to be as high as 30%. Vigilance is indicated, as the presentation of disease is often extrapulmonary (gastrointestinal, genitourinary, central nervous, endocrine, musculoskeletal, laryngeal) and atypical; tuberculosis in this setting sometimes manifests as fever of unknown origin. Careful elicitation of a history and direct evaluation of both the recipient and the donor prior to transplantation are optimal. Skin testing of the recipient with purified protein derivative may be unreliable because of chronic disease and/or immunosuppression. Cell-based assays that measure interferon γ and/or cytokine production may prove more sensitive in the future. Isoniazid toxicity has not been a significant problem except in the setting of liver transplantation. Therefore, appropriate prophylaxis should be used (see recommendations from the Centers for Disease Control and Prevention [CDC] at www.cdc.gov/tb/topic/treatment/lubi.htm). An assessment of the need to treat latent disease should include careful consideration of the possibility of a false-negative test result. Pending final confirmation of suspected tuberculosis, aggressive multidrug treatment in accordance with the guidelines of the CDC, the Infectious Diseases Society of America, and the American Thoracic Society is indicated because of the high mortality rates among these patients. Altered drug metabolism (e.g., upon coadministration of antituberculous medications and certain immunosuppressive agents) can be managed with careful monitoring of drug levels and appropriate dose adjustment. Close follow-up of hepatic enzymes is warranted. Drug-resistant tuberculosis is especially problematic in these individuals (**Chap. 202**).

Virus-Associated Malignancies In addition to malignancy associated with gammaherpesvirus infection (EBV, KSHV) and simple warts (HPV), other tumors that are virus-associated or suspected of being virus-associated are more likely to develop in transplant recipients, particularly those who require long-term immunosuppression, than in the general population. The interval to tumor development is usually >1 year. Transplant recipients develop nonmelanoma skin or lip cancers that, in contrast to de novo skin cancers, have a high ratio of squamous cells to basal cells. HPV may play a major role in these lesions. Cervical and vulvar carcinomas, which are quite clearly associated with HPV, develop with increased frequency in female transplant recipients. The frequency of Merkel cell carcinoma associated with Merkel cell polyomavirus is also increased in transplant recipients; however, it is unclear whether recipients infected with HTLV-1 are at increased risk of leukemia. Among renal transplant recipients, rates of melanoma are modestly increased and rates of cancers of the kidney and bladder are increased.

VACCINATION OF TRANSPLANT RECIPIENTS

(**See also Chap. 148**) In addition to receiving antibiotic prophylaxis, transplant recipients should be vaccinated against likely pathogens (**Table 169-6**). In the case of HSC transplant recipients, optimal responses cannot be achieved until after immune reconstitution, despite previous immunization of both donor and recipient. Recipients of an allogeneic HSC transplant must be reimmunized if they are to be protected against pathogens. The situation is less clear-cut in the case of autologous transplantation. T and B cells in the peripheral blood may reconstitute the immune response if they are transferred

TABLE 169-6 VACCINATION OF HEMATOPOIETIC STEM CELL TRANSPLANT (HSCT) AND SOLID ORGAN TRANSPLANT (SOT) RECIPIENTS

Vaccine	Type of Transplantation	
	HSCT	SOT ^a
<i>Streptococcus pneumoniae</i> , <i>Haemophilus influenzae</i> , <i>Neisseria meningitidis</i>	Immunize after transplantation. See CDC-ACIP recommendations. (For <i>S. pneumoniae</i> , a new primary series may be indicated.)	Immunize before transplantation. See CDC-ACIP recommendations. (For <i>S. pneumoniae</i> , a booster dose of polysaccharide vaccine after 5 years is recommended.)
Influenza	Vaccinate in the fall. Vaccinate close contacts.	Vaccinate in the fall. Vaccinate close contacts.
Polio	Administer inactivated vaccine.	Administer inactivated vaccine.
Measles/mumps/rubella	Immunize 24 months after transplantation if GVHD is absent.	Immunize before transplantation.
Diphtheria, pertussis, tetanus	Reimmunize after transplantation with primary series, DTaP. See IDSA 2013 recommendations (www.idsociety.org/Other_Guidelines/#immunizationFortheCompromisedHost).	Immunize or boost before transplantation with Tdap; give boosters at 10-year intervals or as required.
Hepatitis B and A	Reimmunize after transplantation. See recommendations.	Immunize before transplantation.
Human papillomavirus	Recommendations are pending (www.cdc.gov/std/hpv/stdfact-hpv-vaccine-hcp.htm).	Recommendations are pending.

^aImmunizations should be given before solid organ transplantation whenever possible.

Abbreviations: CDC, Centers for Disease Control and Prevention; ACIP, Advisory Committee on Immunization Practices; DTaP, full-level diphtheria and tetanus toxoids and acellular pertussis, adsorbed; GVHD, graft-versus-host disease; Tdap, tetanus toxoid, reduced diphtheria toxoid, and acellular pertussis; IDSA, Infectious Diseases Society of America.

Note: Recommendations from the CDC should be checked regularly as they frequently change upon receipt of new clinical information and new formulations of specific vaccines.

in adequate numbers. However, cancer patients (particularly those with Hodgkin's disease, in whom vaccination has been extensively studied) who are undergoing chemotherapy do not respond normally to immunization, and titers of antibodies to infectious agents fall more rapidly than in healthy individuals. Therefore, even immunosuppressed patients who have not undergone HSC transplantation may need booster vaccine injections. If memory cells are specifically eliminated as part of a stem cell "cleanup" procedure, it will be necessary to reimmunize the recipient with a new primary series. Optimal times for immunizations of different transplant populations are being evaluated. Yearly immunization of household and other contacts (including health care personnel) against influenza benefits the patient by preventing local spread.

In the absence of compelling data as to optimal timing, it is reasonable to administer the pneumococcal and *H. influenzae* type b conjugate vaccines to both autologous and allogeneic HSC transplant recipients beginning 12 months after transplantation. A series that includes both the 13-valent pneumococcal conjugate vaccine (Prevnar) and the 23-valent pneumococcal polysaccharide vaccine (Pneumovax) is now recommended (according to CDC guidelines). The pneumococcal and *H. influenzae* type b vaccines are particularly important for patients who have undergone splenectomy. The *Neisseria meningitidis* polysaccharide conjugate vaccine (Menactra or Menveo) also is recommended. In addition, diphtheria, tetanus, acellular pertussis, and inactivated polio vaccines can all be given at these same intervals (12 months and, as required, 24 months after transplantation). Some authorities recommend a new primary series for tetanus/diphtheria/pertussis and inactivated poliovirus vaccines beginning 12 months after transplantation. Vaccination to prevent hepatitis B and hepatitis A (both killed

vaccines) also seems advisable. A formal recommendation regarding immunization with the tetravalent HPV virus-like particle vaccine (Gardasil) after HSC transplantation has not been issued. However, HPV vaccination, which can prevent genital warts as well as specific cancers, is recommended through age 26 for healthy young adults who previously have not been vaccinated or have not received the full series. Live-virus measles/mumps/rubella (MMR) vaccine can be given to autologous HSC transplant recipients 24 months after transplantation and to most allogeneic HSC transplant recipients at the same point if they are not receiving maintenance therapy with immunosuppressive drugs and do not have ongoing GVHD. The risk of spread from a household contact is low for MMR vaccine. In parts of the world where live poliovirus vaccine is used, patients as well as contacts should be advised to receive only the killed vaccine. In the rare setting where both donor and recipient are VZV naïve and the recipient is no longer receiving acyclovir or ganciclovir prophylaxis, the patient should be counseled to receive varicella-zoster immune globulin (VariZIG) up to 10 days after an exposure to a person with chickenpox or uncovered zoster; such patients should avoid close contact with persons recently vaccinated with Varivax. A formal recommendation regarding Varivax immunization of such patients is not currently available. Neither patients nor their household contacts should receive vaccinia vaccine unless they have been exposed to smallpox virus. Among patients who have active GVHD and/or are taking high maintenance doses of glucocorticoids, it may be prudent to avoid all live-virus vaccines.

In the case of SOT recipients, administration of all the usual vaccines and of the indicated booster doses should be completed before immunosuppression, if possible, to maximize responses. For patients taking immunosuppressive agents, the administration of pneumococcal vaccine should be repeated every 5 years. No data are available for the meningococcal vaccine, but it is probably reasonable to administer it along with the pneumococcal vaccine. *H. influenzae* conjugate vaccine is safe and should be efficacious in this population; therefore, its administration before transplantation is recommended. Booster doses of this vaccine are not recommended for adults. SOT recipients who continue to receive immunosuppressive drugs should not receive live-virus vaccines. A person in this group who is exposed to measles should be given measles immune globulin. Similarly, an immunocompromised patient who is seronegative for varicella and who comes into contact with a person who has chickenpox should be given varicella-zoster immune globulin as soon as possible (optimally within 96 h; up to 10 days after contact); if this is not possible, a 10- to 14-day course of acyclovir therapy should be started immediately. Upon the discontinuation of treatment, clinical disease may still occur in a small number of patients; thus vigilance is indicated. Rapid re-treatment with acyclovir should limit the symptoms of disease. Household contacts of transplant recipients can receive live attenuated VZV vaccine, but vaccinees should avoid direct contact with the patient if a rash develops. Virus-like particle vaccines have been licensed for the prevention of infection with several HPV serotypes most commonly implicated in cervical and anal carcinomas and in anogenital and laryngeal warts. These vaccines are not live; however, no information is yet available about their immunogenicity or efficacy in transplant recipients.

Immunocompromised patients who travel may benefit from some but not all vaccines (Chaps. 148 and 149). In general, these patients should receive any killed or inactivated vaccine preparation appropriate to the area they are visiting; this recommendation includes the vaccines for Japanese encephalitis, hepatitis A and B, poliomyelitis, meningococcal infection, and typhoid. The live typhoid vaccines are not recommended for use in most immunocompromised patients, but an inactivated or purified polysaccharide typhoid vaccine can be used. Live yellow fever vaccine should not be administered. On the other hand, primary immunization or boosting with the purified-protein hepatitis B vaccine is indicated. Inactivated hepatitis A vaccine should also be used in the appropriate setting (Chap. 148). A vaccine is now available that provides dual protection against hepatitis A and hepatitis B. If hepatitis A vaccine is not administered, travelers should consider receiving passive protection with immune globulin (the dose depending on the duration of travel in the high-risk area).

170 Treatment and Prophylaxis of Bacterial Infections

David C. Hooper, Erica S. Shenoy, Christy A. Varughese

Antimicrobial agents have had a major impact on human health. Together with vaccines, they have contributed to reduced mortality, extended lifespan, and enhanced quality of life. Among drugs used in human medicine, however, they are distinctive in that their use promotes the occurrence of drug resistance in the pathogens they are designed to treat as well as in other “bystander” organisms. Indeed, the history of antimicrobial development has been driven in large part by the medical need engendered by the emergence of resistance to each generation of agents. Thus, the careful and appropriate use of antimicrobial drugs is particularly important not only for optimizing efficacy and minimizing adverse effects but also for minimizing the risk of resistance and preserving the value of existing agents. Although this chapter focuses on antibacterial agents, the optimal use of all antimicrobials depends on an understanding of each drug’s mechanism of action, spectrum of activity, mechanisms of resistance, pharmacology, and adverse effect profile. This information is then applied in the context of the patient’s clinical presentation, underlying conditions, and epidemiology to define the site and likely nature of the infection or other condition and thus to choose the best therapy. Gathering of microbiologic information is important for refining therapeutic choices on the basis of documented pathogen and susceptibility data whenever possible; this information also makes it possible to choose more targeted therapy, thereby reducing the risk of selection of resistant bacteria. Durations of therapy are chosen according to the nature of the infection and the patient’s response to treatment and are informed by clinical studies when they are available, with the understanding that shorter courses are less likely than longer ones to promote the emergence of resistance. This chapter provides specific information that is necessary for making informed choices among antibacterial agents.

MECHANISMS OF ACTION AND RESISTANCE

The mechanisms of action of and resistance to antibacterial agents are discussed in detail in the text and are summarized for the most commonly used groups of agents in [Table 170-1](#). A schematic of antibacterial targets is provided in [Fig. 170-1](#).

MECHANISMS OF ACTION

Multiple essential components of bacterial cell structures and metabolism have been the targets of antibacterial agents used in clinical medicine, and the interaction of an agent with its target results in either inhibition of bacterial growth and replication (*bacteriostatic effect*) or bacterial killing (*bactericidal effect*). In general, targets have been chosen because they either do not exist in mammalian cells and physiology or are sufficiently different from their bacterial counterparts to allow selective bacterial targeting. Treatment with bacteriostatic agents is effective when the patient’s host defenses are sufficient to contribute to eradication of the infecting pathogen. In patients with impaired host defenses (e.g., neutropenia) or infections at body sites with impaired or limited host defenses (e.g., meningitis and endocarditis), bactericidal agents are generally preferred.

Inhibition of Cell Wall Synthesis The bacterial cell wall, which is external to the cytoplasmic membrane and has no counterpart in mammalian cells, protects bacterial cells from lysis under low osmotic conditions. The cell wall is a cross-linked peptidoglycan composed of a polymer of alternating units of *N*-acetylglucosamine (NAG) and *N*-acetylmuramic acid (NAM), four-amino-acid stem peptides linked to each NAM, and

a peptide cross-bridge that links adjacent stem peptides to form a net-like structure. Several steps in peptidoglycan synthesis are targets of antibacterial agents. Inhibition of cell wall synthesis generally results in a bactericidal effect that is linked to cell lysis. This effect results not only from the blocking of new cell-wall formation but from the uninhibited action of cell wall–remodeling enzymes called *autolysins*, which cleave peptidoglycan as part of normal cell-wall growth.

In gram-positive bacteria the peptidoglycan is the most external cell structure, but in gram-negative bacteria an asymmetric lipid outer membrane is external to the peptidoglycan and contains diffusion channels called *porins*. The space between the cytoplasmic membrane peptidoglycan and the outer membrane is referred to as the *periplasmic space*. Most antibacterial drugs enter the gram-negative bacterial cell through a porin channel, since the outer membrane is a major diffusion barrier. Although the peptidoglycan layer is thicker in gram-positive (20–80 nm) than in gram-negative (1 nm) bacteria, peptidoglycan itself constitutes only a limited diffusion barrier for antibacterial agents.

β-LACTAMS The β-lactam drugs, including penicillins, cephalosporins, monobactams, and carbapenems, target transpeptidase enzymes (also called *penicillin-binding proteins*, or PBPs) involved in the stem-peptide cross-linking step.

GLYCOPEPTIDES The glycopeptides, including vancomycin, teicoplanin, telavancin, dalbavancin, and oritavancin, bind the two terminal *D*-alanine residues of the stem peptide, hindering the glycosyltransferase involved in polymerizing NAG–NAM units. Telavancin also binds to the lipid II intermediate that delivers cell-wall precursor subunits. Likewise, dalbavancin and oritavancin interact with the cell membrane, and oritavancin may also inhibit transpeptidases. Both β-lactams and glycopeptides interact with their targets external to the cytoplasmic membrane.

BACITRACIN (TOPICAL) AND FOSFOMYCIN These agents interrupt enzymatic steps in the production of peptidoglycan precursors in the cytoplasm.

Inhibition of Protein Synthesis Most inhibitors of bacterial protein synthesis target bacterial ribosomes, whose difference from eukaryotic ribosomes allows selective antibacterial action. Some inhibitors bind to the 30S ribosomal subunit and others to the 50S subunit. Most protein synthesis–inhibiting agents are bacteriostatic; aminoglycosides are an exception and are bactericidal.

AMINOGLYCOSIDES Aminoglycosides (amikacin, gentamicin, kanamycin, netilmicin, streptomycin, tobramycin) bind irreversibly to 16S ribosomal RNA (rRNA) of the 30S ribosomal subunit, blocking the translocation of peptidyl transfer RNA (tRNA) from the A (aminoacyl) to the P (peptidyl) site and, at low concentrations, causing misreading of messenger RNA (mRNA) codons and thus causing the introduction of incorrect amino acids into the peptide chain; at higher concentrations, translocation of the peptide chain is blocked. Cellular uptake of aminoglycosides is dependent on the electrochemical gradient across the bacterial membrane. Under anaerobic conditions, this gradient is reduced, with a consequent reduction in the uptake and activity of the aminoglycosides. Spectinomycin is a related aminocyclitol antibiotic that also binds to 16S rRNA of the 30S ribosomal subunit but at a different site. This drug inhibits translocation of the growing peptide chain but does not trigger codon misreading and produces only a bacteriostatic effect.

TETRACYCLINES AND GLYCYLCYCLES Tetracyclines (doxycycline, minocycline, tetracycline) bind reversibly to the 16S rRNA of the 30S ribosomal subunit and block the binding of aminoacyl tRNA to the ribosomal A site, thereby inhibiting peptide elongation. Active transport of tetracyclines into bacterial but not mammalian cells contributes to the selectivity of these agents. Tigecycline, a derivative of minocycline and

TABLE 170-1 MECHANISMS OF ACTION OF AND RESISTANCE TO ANTIBACTERIAL AGENTS

Antibacterial Agent(s)	Major Target	Mechanism(s) of Action	Mechanism(s) of Resistance
β -Lactams (penicillins, cephalosporins, monobactams, carbapenems)	Cell wall synthesis	Bind cell wall cross-linking enzymes (PBPs, transpeptidases)	1. Drug inactivation by β -lactamases 2. Altered PBP targets 3. Reduced diffusion through porin channels
Glycopeptides (vancomycin, teicoplanin, telavancin, dalbavancin, oritavancin)	Cell wall synthesis	Block cell wall glycosyltransferases by binding D-Ala-D-Ala stem-peptide terminus Telavancin, dalbavancin, and oritavancin: affect membrane function Oritavancin: inhibits transpeptidases	1. Altered D-Ala-D-Ala target (D-Ala-D-Lac) 2. Increased D-Ala-D-Ala target binding at sites distant from cell wall synthesis enzymes
Bacitracin	Cell wall synthesis	Blocks lipid carrier of cell wall precursors	Active drug efflux
Fosfomycin	Cell wall synthesis	Blocks linkage of stem peptide to NAG by enoyltransferase	1. Target enzyme overexpression 2. Drug-modifying enzymes
Aminoglycosides (gentamicin, tobramycin, amikacin)	Protein synthesis	Bind 30S ribosomal subunit Block translocation of peptide chain Cause misreading of mRNA	1. Drug-modifying enzymes 2. Methylation at ribosome binding site 3. Decreased permeation to target due to active efflux
Tetracyclines (tetracycline, doxycycline, minocycline)	Protein synthesis	Bind 30S ribosomal subunit Inhibit peptide elongation	1. Active drug efflux 2. Ribosomal protection proteins
Tigecycline	Protein synthesis	Same as tetracyclines	Active drug efflux (pumps different from those affecting tetracyclines)
Macrolides (erythromycin, clarithromycin, azithromycin) and ketolide (telithromycin)	Protein synthesis	Bind 50S ribosomal subunit Block peptide chain exit	1. Methylation at ribosome binding site 2. Active drug efflux
Lincosamides (clindamycin)	Protein synthesis	Bind 50S ribosomal subunit Block peptide bond formation	Methylation at ribosome binding site
Streptogramins (quinupristin, dalfopristin)	Protein synthesis	Same as macrolides	1. Same as macrolides 2. Drug-modifying enzymes
Chloramphenicol	Protein synthesis	Binds 50S ribosomal subunit Blocks aminoacyl tRNA positioning	Drug-modifying enzymes
Oxazolidinones (linezolid, tedizolid)	Protein synthesis	Bind 50S ribosomal subunit Inhibit initiation of peptide synthesis	1. Altered rRNA binding site 2. Methylation of ribosome binding site
Mupirocin	Protein synthesis	Blocks isoleucyl tRNA synthetase	1. Acquired resistant tRNA synthetase (drug bypass) 2. Altered native tRNA synthetase target
Sulfonamides (sulfadiazine, sulfisoxazole, and sulfamethoxazole)	Folate synthesis	Inhibit dihydropteroate synthetase	Acquired resistant dihydropteroate synthetase (drug bypass)
Trimethoprim	Folate synthesis	Inhibits dihydrofolate reductase	Acquired resistant dihydrofolate reductase (drug bypass)
Quinolones (norfloxacin, ciprofloxacin, ofloxacin, levofloxacin, moxifloxacin, gemifloxacin)	DNA synthesis	Inhibit DNA gyrase and DNA topoisomerase IV Enzyme-DNA-drug complex: blocks DNA replication apparatus	1. Altered target(s) 2. Active efflux 3. Protection of target from drug 4. Drug-modifying enzyme (ciprofloxacin)
Rifamycins (rifampin, rifabutin, rifapentine)	RNA synthesis	Inhibit RNA polymerase	Altered target
Nitrofurantoin	Nucleic acid synthesis	Reduce reactive drug derivatives that damage DNA	Altered drug-activating enzymes
Metronidazole	Nucleic acid synthesis	Reduce reactive drug derivatives that damage DNA	1. Altered drug-activating enzyme 2. Acquired detoxifying enzymes 3. Active efflux
Polymyxins (polymyxin B and polymyxin E [colistin])	Cell membrane	Bind LPS and disrupt both outer and cytoplasmic membranes	Altered cell membrane charge with reduced drug binding
Daptomycin	Cell membrane	Produces membrane channel and membrane leakage	Altered cell membrane with reduced drug binding

Abbreviations: LPS, lipopolysaccharide; NAG, *N*-acetylglucosamine; PBP, penicillin-binding protein.

the only available glycylycine, acts similarly to the tetracyclines but is distinctive for its ability to circumvent the most common mechanisms of resistance to the tetracyclines.

MACROLIDES AND KETOLIDES In contrast to the aminoglycosides and tetracyclines, the macrolides (azithromycin, clarithromycin, erythromycin) and ketolides (telithromycin) bind to the 23S rRNA of the 50S ribosomal subunit. These agents block translocation of the growing peptide chain by binding to the tunnel from which the chain exits the ribosome.

LINCOSAMIDES Clindamycin is the only lincosamide in clinical use. It binds to the 23S rRNA of the 50S ribosomal subunit, interacting with both the ribosomal A and P sites and blocking peptide bond formation.

STREPTOGRAMINS The only streptogramin in clinical use is a combination of quinupristin, a group B streptogramin, and dalfopristin, a group A streptogramin. Both components bind to 23S rRNA of the 50S ribosome: dalfopristin binds to both the A and P sites of the peptidyl transferase center, and quinupristin binds to a site that overlaps the macrolide-binding site, blocking the emergence of nascent peptide

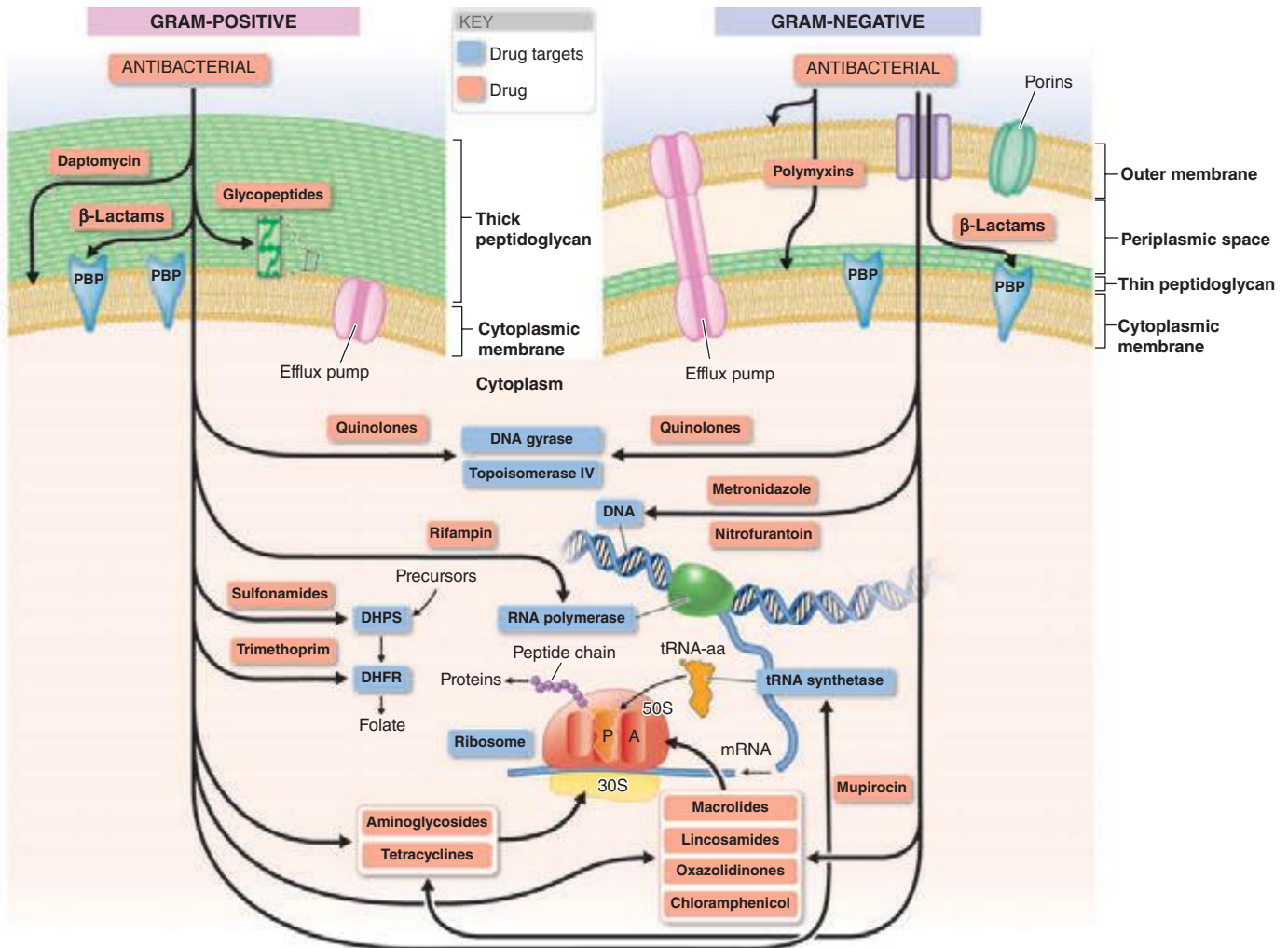


FIGURE 170-1 Antibacterial targets. A, aminoacyl site; DHFR, dihydrofolate reductase; DHPS, dihydropteroate synthetase; P, peptidyl site; PBP, penicillin-binding protein; tRNA-aa, aminoacyl tRNA.

from the ribosome. The combination is bactericidal, but macrolide-resistant bacteria exhibit cross-resistance to quinupristin, and the remaining activity of dalbapristin alone is bacteriostatic.

CHLORAMPHENICOL Chloramphenicol binds reversibly to the 23S rRNA of the 50S subunit in a manner that interferes with the proper positioning of the aminoacyl component of tRNA in the A site. This site of binding is near those of the macrolides and lincosamides.

OXAZOLIDINONES Linezolid and tedizolid are the only oxazolidinones in clinical use. They bind directly to the A site in the 23S rRNA of the 50S ribosomal subunit and block binding of aminoacyl tRNA, inhibiting the initiation of protein synthesis.

MUPIROCIN Mupirocin (pseudomonic acid) is used topically. It competes with isoleucine for binding to isoleucyl tRNA synthetase, depleting stores of isoleucyl tRNA and thereby inhibiting protein synthesis.

Inhibition of Bacterial Metabolism Available inhibitors (antimetabolites) target the pathway for synthesis of folate, which is a cofactor in a number of one-carbon transfer reactions involved in the synthesis of some nucleic acids, including pyrimidine, thymidine, and all purines (adenine and guanine), as well as some amino acids (methionine and serine) and acetyl coenzyme A (CoA). Two sequential steps in folate synthesis are targeted. The selective antibacterial effect stems from the inability of mammalian cells to synthesize folate; they depend instead on exogenous sources. Antibacterial activity, however, may be reduced in the presence of high exogenous concentrations of the end products of the folate pathway (e.g., thymidine and purines) that may occur in

some infections, resulting from local breakdown of leukocytes and host tissues.

SULFONAMIDES Sulfonamides, including sulfadiazine, sulfisoxazole, and sulfamethoxazole, inhibit dihydropteroate synthetase, which adds *p*-aminobenzoic acid (PABA) to pteridine, producing dihydropteroate. Sulfonamides are structural analogues of PABA and act as competing enzyme substrates.

TRIMETHOPRIM Subsequent steps in folate synthesis are catalyzed by dihydrofolate synthase, which adds glutamate to dihydropteroate, and dihydrofolate reductase, which then generates the final product, tetrahydrofolate. Trimethoprim is a structural analogue of pteridine and inhibits dihydrofolate reductase. Trimethoprim is available alone but is most often used in combination products that also contain sulfamethoxazole and thus block two sequential steps in folate synthesis.

Inhibition of DNA and RNA Synthesis or Activity A variety of antibacterial agents act on these processes.

QUINOLONES The quinolones include nalidixic acid, the first agent in the class, and newer, more widely used fluorinated derivatives (fluoroquinolones), including norfloxacin, ciprofloxacin, levofloxacin, moxifloxacin, and gemifloxacin. The quinolones are synthetic compounds that inhibit bacterial DNA synthesis by interacting with the DNA complexes of two essential enzymes, DNA gyrase and DNA topoisomerase IV, which alter DNA topology. Quinolones trap enzyme–DNA complexes in such a way that they block movement of the DNA replication apparatus and can generate lethal double-strand breaks

in DNA, resulting in bactericidal activity. Although mammalian cells also have type II DNA topoisomerases like gyrase and topoisomerase IV, the structures of the mammalian enzymes are sufficiently different from those of the bacterial enzymes that quinolones have substantially selective antibacterial activity.

RIFAMYCINS Rifampin, rifabutin, and rifapentine are semisynthetic derivatives of rifamycin B and bind the β subunit of bacterial RNA polymerase, thereby blocking elongation of mRNA. Their action is highly selective for the bacterial enzyme over mammalian RNA polymerases.

NITROFURANTOIN The reduction of nitrofurantoin, a nitrofurantoin compound, by bacterial enzymes produces highly reactive derivatives that are thought to cause DNA strand breakage. Nitrofurantoin is used only for the treatment of lower urinary tract infections.

METRONIDAZOLE Metronidazole is a synthetic nitroimidazole with activity limited to anaerobic bacteria and certain anaerobic protozoa. Reduction of its nitro group by the electron-transport system in anaerobic bacteria produces reactive intermediates that damage DNA and result in bactericidal activity. Both nitrofurantoin and metronidazole have selective antibacterial activity because the reducing activity needed to generate active derivatives is generated only by bacterial and not mammalian enzymes.

Disruption of Membrane Integrity The integrity of the bacterial cytoplasmic membrane—and, in gram-negative bacteria, the outer membrane—is important for bacterial viability. Two bactericidal drugs have membrane targets.

POLYMYXINS The polymyxins, including polymyxin B and polymyxin E (colistin), are cationic cyclic polypeptides that disrupt the cytoplasmic membrane and the outer membrane (the latter by binding lipopolysaccharide).

DAPTOMYCIN Daptomycin is a lipopeptide that binds the cytoplasmic membrane of gram-positive bacteria in the presence of calcium, generating a channel that leads to leakage of cytoplasmic potassium ions and membrane depolarization.

MECHANISMS OF RESISTANCE

Bacteria use a wide variety of mechanisms to block or circumvent the activity of antibacterial agents. Although myriad, these mechanisms can generally be grouped into three categories: (1) altered or bypass targets that exhibit reduced binding of the drug, (2) altered access of the drug to its target by reductions in uptake or increases in active efflux, and (3) a modification of the drug that reduces its activity. These mechanisms result from either mutations in bacterial chromosomal genes occurring spontaneously during bacterial DNA replication or the acquisition of new genes by DNA transfer from other bacteria or uptake of exogenous DNA. New genes are most often acquired on self-replicating plasmids or other DNA elements transferred from other bacteria. However, some bacteria, such as *Streptococcus pneumoniae* and *Neisseria gonorrhoeae*, can also take up fragments of environmental DNA from related species and recombine that DNA directly into their own chromosomes, a process called *transformation*. Not uncommonly, resistant bacteria have combinations of resistance mechanisms either within one category or among categories, and many plasmids contain more than one resistance gene. Thus, plasmid acquisition itself can in many cases confer resistance to multiple antibacterial agents.

Many antibacterial drugs are derived from natural products of microbial species. Some genes encoding resistance to these drugs may have evolved and been mobilized onto plasmids from a protection mechanism in the producer organism or in other surviving bacteria in the exposed environment. Exposure to antibacterial agents either in nature or from human or animal use results in the selection of resistant strains within an otherwise susceptible bacterial population. Because the patterns and extent of resistance may differ among settings, initial choices of antibacterial drugs should be based, whenever possible, on local susceptibility data and should be modified as needed as soon as specific microbiology susceptibility data become available.

β -Lactams The most common mechanism of resistance to β -lactams is their degradation by β -lactamases, enzymes that break down the core β -lactam ring and destroy drug activity. Different β -lactamases degrade different β -lactams. Some β -lactamases are encoded on the bacterial chromosome, and their activity contributes to the susceptibility profile of a particular species. Because other β -lactamases are encoded by acquired plasmids, their resistance profiles may be present in some strains of a species but not others. In gram-positive bacteria β -lactamases are secreted into the extracellular environment, whereas in gram-negative bacteria these enzymes are secreted into the periplasmic space between the cytoplasmic and outer membranes. Thus, in gram-negative bacteria, access of β -lactams both to their target PBPs and to β -lactamases requires diffusion across the outer membrane, generally through the porin channels.

Most strains of *Staphylococcus aureus* produce a plasmid-encoded β -lactamase that degrades penicillin but not semisynthetic penicillins, such as oxacillin and nafcillin. The most common plasmid-encoded β -lactamases of gram-negative bacteria are able to inactivate all penicillins and most earlier-generation cephalosporins. Extended-spectrum β -lactamase (ESBL) variants of these early enzymes that can degrade later-generation cephalosporins (ceftriaxone, cefotaxime, ceftazidime) as well as the monobactam aztreonam have now emerged and are widely disseminated. Some ESBLs also degrade the fourth-generation cephalosporin cefepime. Carbapenems (imipenem, meropenem, ertapenem, doripenem) are not degraded by ESBLs, but additional β -lactamases, called *carbapenemases*, that degrade carbapenems and most if not all other β -lactams have begun to emerge.

The chromosomal β -lactamase of *Klebsiella pneumoniae* preferentially degrades penicillins but not cephalosporins. In contrast, the chromosomal β -lactamase of *Enterobacter* and related genera, AmpC, can degrade almost all cephalosporins but is normally expressed in small amounts. Mutations that cause increased amounts of AmpC to be produced confer full resistance to penicillins and cephalosporins; the exceptions are ceftiofex and cefepime, which are relatively stable to AmpC. Resistance to cefepime can develop, however, through the combined effects of increased AmpC production and decreased porin diffusion channels. Genes encoding AmpC have also been found on plasmids but are less common than plasmid-encoded ESBLs.

Inhibitors of β -lactamases such as clavulanate, sulbactam, and tazobactam have been developed and paired with amoxicillin and ticarcillin, ampicillin, and piperacillin, respectively. These inhibitors have little or no antibacterial activity of their own but inhibit plasmid-mediated β -lactamases, including ESBLs but not AmpC enzymes.

Resistance to β -lactams also occurs through alterations in their target PBPs. In *S. pneumoniae*, *N. gonorrhoeae*, and *Neisseria meningitidis*, resistance to penicillin occurs by recombination of transformed DNA from related species. In staphylococci, resistance to methicillin and other β -lactams occurs by the acquisition of the *mec* gene, which encodes PBP2a with reduced drug affinity. Ceftaroline is the only β -lactam that has affinity for PBP2a and is thus active against methicillin-resistant staphylococcal strains.

Glycopeptides Resistance to vancomycin in enterococci is due to the acquisition of a set of *van* genes that result in (1) the production of D-alanine-D-lactate—instead of the normal D-alanine-D-alanine—at the end of the peptidoglycan stem peptide and (2) the reduction of existing D-alanine-D-alanine terminated peptides. Vancomycin binds D-alanine-D-lactate with a thousandfold lower affinity than D-alanine-D-alanine. In a small number of cases, the *van* gene cassettes have been transferred from enterococci to *S. aureus*, with the consequent generation of full vancomycin resistance. Particularly in patients receiving prolonged courses of vancomycin, intermediate resistance to this drug has developed in *S. aureus* by a different mechanism: multiple chromosomal mutations that result in a thickened and poorly cross-linked cell wall in which multiple distant D-alanine-D-alanine stem peptide termini exist and bind vancomycin, impeding its access to the binding sites proximal to the cell membrane where new cell-wall synthesis occurs and where binding would block transpeptidase and transglycosylase enzymes. Susceptibility to telavancin, dalbavancin, and oritavancin is

934 also reduced in strains that exhibit resistance or intermediate susceptibility to vancomycin, although in some cases strains may still be classified as susceptible on the basis of clinical interpretive criteria.

Aminoglycosides The most common mechanism of resistance is due to acquisition of plasmid genes encoding transferase enzymes that modify aminoglycosides by the addition of acetyl, adenylyl, or phosphate groups; these added groups decrease the drugs' binding affinity to their ribosomal target site. Transferases differ in which aminoglycosides they modify, and amikacin resistance occurs less often than resistance to gentamicin or tobramycin. More recently, plasmids encoding methyltransferases that modify the ribosomal site of aminoglycoside binding and confer resistance to all aminoglycosides have been found in enteric gram-negative bacteria. For streptomycin, a ribosomal protein mutation may cause resistance. In *Pseudomonas aeruginosa*, resistance may also occur through mutations causing increased expression of a chromosomally encoded efflux pump, MexXY.

Tetracyclines and Glycylcyclines For tetracyclines, resistance is most often plasmid mediated and attributable either to active efflux pumps, which are generally specific for tetracyclines, or to proteins that protect the ribosome from tetracycline action. Resistance to the glycylcycline tigecycline, which is not affected by the usual tetracycline resistance mechanisms, can occur through mutations that cause overexpression of certain broad-spectrum efflux pumps in *Proteus* species.

Macrolides, Ketolides, Lincosamides, and Streptogramins Resistance to macrolides, clindamycin, and quinupristin is most often due to plasmid-acquired methylases that modify the drug binding site on the ribosome. Resistance to quinupristin by this mechanism renders the quinupristin-dalfopristin combination bacteriostatic rather than bactericidal. Telithromycin, a ketolide, has an additional binding site on the ribosome and remains active in the presence of these methylases. Acquired genes encoding active efflux pumps can also contribute to resistance to macrolides in streptococci and resistance to macrolides, clindamycin, and dalfopristin in staphylococci. Plasmid-acquired drug-modifying enzymes in staphylococci can also cause resistance to quinupristin and dalfopristin. Macrolide resistance due to 23S rRNA mutations is uncommon in staphylococci and streptococci because of the multiple copies of the rRNA genes on the chromosomes of these species; such resistance may occur more frequently, however, in mycobacteria and *Helicobacter pylori*, which have only single chromosomal copies of these rRNA genes.

Chloramphenicol Resistance to chloramphenicol is most often due to a plasmid-encoded drug-modifying acetyltransferase.

Oxazolidinones Linezolid resistance has been seen in enterococci more often than in staphylococci and, in both organisms, is due to mutations in multiple copies of the 23S rRNA genes that reduce drug binding to the ribosome. A plasmid-acquired ribosomal methylase gene that confers resistance to both chloramphenicol and linezolid has also been found in some strains of staphylococci but is not yet widespread. Tedizolid may still be active against some but not all linezolid-resistant strains.

Mupirocin Resistance to mupirocin occurs by either mutation in the target leucyl-tRNA synthetase (low-level resistance) or the acquisition of a plasmid-encoded resistant tRNA synthetase (high-level resistance).

Sulfonamides and Trimethoprim Resistance to both of these antimetabolites is due to plasmid-acquired genes encoding resistant enzymes that bypass the inhibition of the native sensitive enzymes—a resistant dihydropteroate synthetase in the case of sulfonamides and a resistant dihydrofolate reductase in the case of trimethoprim.

Quinolones Resistance to quinolones is most often due either to chromosomal mutations altering the target enzymes DNA gyrase and DNA topoisomerase IV, with consequent reduction in drug binding, or to mutations that increase the expression of native broad-spectrum efflux pumps for which quinolones (among other compounds) are substrates. In addition, three types of genes can confer reduced susceptibility or low-level resistance by protecting target enzymes, modifying some

quinolones, or pumping quinolones out of the cell (efflux). These genes are located on multidrug resistance plasmids that have spread worldwide. Their presence can promote the selection of higher levels of quinolone resistance linked to resistance to other antibacterial drugs that is encoded on the same plasmid.

Rifampin and Rifabutin Single mutations in the β subunit of RNA polymerase can cause high-level resistance to rifampin. Thus rifampin and other rifamycins are used for treatment of infections only in combination with other antibacterial drugs in order to prevent resistance.

Metronidazole Acquired resistance to metronidazole in *Bacteroides* species is rare. Such resistance has been reported in strains that lack endogenous nitroreductase activity or that have acquired *nim* genes responsible for further reduction of DNA-damaging nitroso intermediates to an inactive derivative. Active efflux and enhanced DNA repair mechanisms also have been associated with resistance.

Nitrofurantoin Resistance to nitrofurantoin in *Escherichia coli* can emerge through a series of mutations that progressively decrease the nitroreductase activity necessary for generating active nitrofurantoin metabolites.

Polymyxins Because of emerging multidrug resistance in gram-negative bacteria, colistin and polymyxin B are being used increasingly for infections due to resistant Enterobacteriaceae, *P. aeruginosa*, and *Acinetobacter* species. Rates of resistance vary. Resistance can emerge during therapy through mutations that cause reductions in the negative charge of the gram-negative bacterial cell surface, thereby reducing binding of the positively charged colistin.

Daptomycin The mechanisms of resistance to daptomycin are complex and involve mutations in several genes that can alter cell membrane charge and reduce daptomycin binding. Resistance to daptomycin is relatively infrequent but has emerged in some *S. aureus* strains with intermediate vancomycin susceptibility from patients treated with vancomycin but not with daptomycin. In some methicillin-resistant *S. aureus* (MRSA) strains, daptomycin resistance has been linked to acquired susceptibility to β -lactams; combinations of daptomycin and nafcillin have been successful for treatment of patients infected with resistant strains when daptomycin alone or in combination with other agents has failed. The mechanism of this effect is not yet clear.

PHARMACOKINETICS AND PHARMACODYNAMICS

The term *pharmacokinetics* describes the disposition of a drug in the body, whereas *pharmacodynamics* describes the determinants of drug action on the pathogen in relation to pharmacokinetic factors. An understanding of the principles governing these two areas is required for effective drug selection, dosing, and prevention of toxicities.

PHARMACOKINETICS

The process of drug disposition has four principal phases: absorption, distribution, metabolism, and excretion. These components determine the time course of drug concentrations in serum and subsequently the concentrations in other tissues and body fluids.

Absorption When a drug is given by a particular route, *absorption* is defined as the percentage of the dose that reaches the systemic circulation. For example, since IV administration provides direct access to the systemic circulation, 100% of a drug dose given IV is usually absorbed. The level of absorption becomes more relevant when non-IV routes are used—e.g., the oral, IM, SC, and topical routes. The percentage of a drug that is absorbed is termed its *bioavailability*. Examples of antibacterial agents with a high oral bioavailability include metronidazole, levofloxacin, and linezolid. IV administration and oral dosing for highly bioavailable agents usually give equivalent results. Many factors can affect a drug's oral bioavailability, including the timing of food consumption relative to drug administration, drug-metabolizing enzymes, efflux transporters, concentration-dependent solubility, and acid degradation. Underlying conditions such as diarrhea or ileus can also affect the site of drug absorption and thereby alter bioavailability. Certain orally administered drugs have lower bioavailability because

of the *first-pass effect*—the process by which drugs are absorbed in the small intestine through the portal circulation and then directly transported to the liver for metabolism.

Distribution *Distribution* describes the process by which a drug transfers reversibly between the general circulation and the tissues. After absorption into the general circulation and the central compartment (the extensively perfused organs), the drug will also distribute into the peripheral compartment (less well-perfused tissues). The volume of distribution (Vd) is a pharmacokinetic parameter that describes the amount of drug in the body at a given time relative to the measured serum concentration. Properties such as the drug's lipophilicity, partition coefficient within different body tissues, and protein binding; blood flow; and pH can affect the volume of distribution. Drugs with a small volume of distribution are limited to certain areas within the body (typically extracellular fluid), whereas those with a higher volume of distribution penetrate extensively into tissues throughout the body. Antibacterial drugs can bind to serum proteins, and a given drug is usually described as either poorly or highly protein bound. Only the unbound (free) drug is active and available to exert antibacterial effects. For example, because tigecycline is highly protein bound and also has a large volume of distribution, concentrations of free drug in the serum are low.

Metabolism *Metabolism* is the chemical transformation of a drug by the body. This modification can occur within several areas; the liver is the organ most commonly involved. Drugs are metabolized by enzymes, but enzyme systems have a finite capacity to metabolize a substrate drug. If a drug is given in a dose at which the concentration does not exceed the rate of metabolism, then the metabolic process is generally linear. If the dose exceeds the amount that can be metabolized, drug accumulation and potential toxicity may occur. Drugs are metabolized through phase I or phase II reactions. In phase I reactions, the drug is made more polar through dealkylation, hydroxylation, oxidation, and deamination. Polarity facilitates drug removal from the body. Phase II reactions, which include glucuronidation, sulfation, and acetylation, result in compounds larger and more polar than the parent drug. Both phases usually inactivate the parent drug, but some drugs are rendered more active. The cytochrome P450 (CYP) enzyme system is responsible for phase I reactions and is generally found in the liver. CYP3A4 is a common subfamily within this system that is responsible for the majority of drug metabolism. Antibacterial drugs can be substrates, inhibitors, or inducers of a particular CYP enzyme. Inducers, such as rifampin, can increase the production of CYP enzymes and consequently increase the metabolism of other drugs. Inhibitors, such as quinupristin-dalfopristin, cause a decrease in enzyme activity (or competition for CYP substrate) and therefore an increase in the concentration of the interacting drug.

Excretion *Excretion* describes the body's mechanisms of drug elimination. Drugs can be eliminated through more than one mechanism. Renal clearance is the most common route and includes elimination through glomerular filtration, tubular secretion, and/or passive diffusion. Some agents have nonrenal clearance and rely on the biliary tree or the intestine for excretion. Excretion affects the half-life of a drug—i.e., the time it takes for the blood concentration of a drug to decrease by one-half. This value can range from minutes to days. Approximately five to seven half-lives are required for a drug to reach steady state when multiple doses are given in a time frame shorter than the half-life itself. Drug half-life and overall clearance can be extended if the organ responsible for clearance is impaired. Patients with renal or hepatic impairment may require dose adjustments that take delayed clearance into account and prevent toxicities from drug accumulation. For example, imipenem is cleared predominantly through glomerular filtration, and in the presence of renal impairment the dosing interval is typically increased to account for the increased half-life.

PHARMACODYNAMICS

The term *pharmacodynamics* describes the relationship between the serum concentrations that determine the efficacy of the drug and the serum concentrations that produce the toxic effects of the drug. For an antibacterial agent, the pharmacodynamic focus is the type of drug

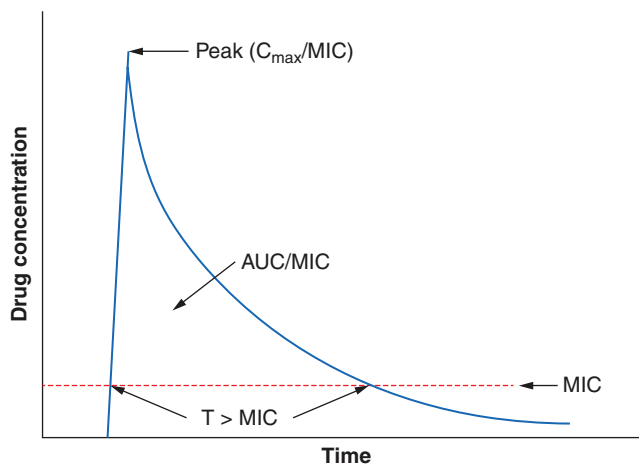


FIGURE 170-2 Pharmacokinetic and pharmacodynamic model predicting efficacy of antibacterial drugs. AUC, area under the time–concentration curve; C_{max} , peak serum concentration of drug; MIC, minimal inhibitory concentration; T>MIC, duration of drug concentrations above the MIC.

exposure needed for optimal antibacterial effect in relation to the minimal inhibitory concentration (MIC)—the lowest drug concentration that inhibits the visible growth of a microorganism under standardized laboratory conditions. Antibacterial effect usually correlates with one of the following parameters: (1) ratio of peak serum concentration to the MIC (C_{max}/MIC), (2) ratio of the area under the concentration–time curve to the MIC (AUC/MIC), or (3) duration of concentrations above the MIC (T>MIC) (Fig. 170-2).

For *concentration-dependent* killing agents, as the designation implies, the higher the drug concentration, the higher the rate and extent of bacterial killing. Aminoglycosides fit into the C_{max}/MIC model of pharmacodynamics activity, and a particular peak serum concentration is often targeted to achieve optimal killing. Fluoroquinolones exemplify antibacterial agents for which the AUC/MIC is a predictor of efficacy. For example, studies have found that an AUC/MIC ratio of >30 will maximize killing of *S. pneumoniae* by fluoroquinolones. In contrast, *time-dependent* killing agents reach a ceiling at which higher concentrations do not result in increased effect. Instead, these agents are active against bacteria only when the drug concentration is above the MIC. The T>MIC predicts clinical efficacy for all β -lactams. The longer the concentration of the β -lactam remains above the MIC for an infecting pathogen during the dosing interval, the greater the killing effect. For some drug classes, such as aminoglycosides, a *postantibiotic effect*—the delayed regrowth of surviving bacteria after exposure to an antibiotic—supports less frequent dosing.

APPROACH TO THERAPY

The approach to antibiotic therapy is driven by host factors, site of infection, and local resistance profiles of suspected or known pathogens. Further, national and local drug shortages and formulary restrictions can affect available therapies. Regular monitoring of the patient and collection of laboratory data should be undertaken to streamline antibacterial therapy as appropriate and to investigate the possibility of treatment failure if the patient fails to respond appropriately.

EMPIRICAL AND DIRECTED THERAPY

Therapy is considered *empirical* when the causative agent has yet to be determined and therapeutic decisions are based on the severity of illness and the clinician's assessment of likely pathogens in light of the clinical syndrome, the patient's medical conditions and prior therapy, and relevant epidemiologic factors. For patients with severe illness, empirical therapy often takes the form of an antibacterial combination that provides broad coverage of diverse agents and thus ensures adequate treatment of possible pathogens while additional data are being collected. *Directed* therapy is predicated on identification of the

936 pathogen, determination of its susceptibility profile, and establishment of the extent of the infection. Directed therapy generally allows the use of more targeted and narrower-spectrum antibacterial agents than does empirical therapy.

Information on epidemiology, exposures, and local antibacterial susceptibility patterns can help guide empirical therapy. When empirical treatment is clinically appropriate, care should be taken to obtain clinical specimens for microbiologic analysis before the initiation of therapy and to de-escalate therapy as new information is obtained about the patient's clinical condition and the causal pathogens. De-escalation to the point of directed therapy can limit unnecessary risks to the patient as well as the risk of emergence of antibacterial resistance.

SITE OF INFECTION

The site of infection is a consideration in antibacterial therapy, largely because of the differing abilities of drugs to penetrate and achieve adequate concentrations at particular body sites. For example, to be effective in the treatment of meningitis, an agent must (1) be able to cross the blood–brain barrier and reach adequate concentrations in the cerebrospinal fluid (CSF) and (2) be active against the relevant pathogen(s). Dexamethasone, administered with or 15–20 min before the first dose of an antibacterial drug, has been shown to improve outcomes in patients with acute bacterial meningitis, but its use may reduce penetration of some antibacterial agents, such as vancomycin, into the CSF. In this case, rifampin is added because its penetration is not reduced by dexamethasone. Infections at other sites where either pathogens are protected from normal host defenses or penetration of an antibacterial drug is suboptimal include osteomyelitis, prostatitis, intraocular infections, and abscesses. In such cases, consideration must be given to the mechanism of drug delivery (e.g., intravitreal injections) as well as to the role of interventions to drain, debride, or otherwise reduce the barriers to effective antibacterial therapy.

HOST FACTORS

Host factors, including immune function, pregnancy, allergies, age, renal and hepatic function, drug–drug interactions, comorbid conditions, and occupational or social exposures, should be considered.

Immune Dysfunction Patients with deficits in immune function that blunt the response to bacterial infection, including neutropenia, deficient humoral immunity, and asplenia (either surgical or functional), are all at increased risk of severe bacterial infection. Such patients should be treated aggressively and often broadly in the early stages of suspected infection pending results of microbiologic tests. For asplenic patients, treatment should include coverage of encapsulated organisms, particularly *S. pneumoniae*, that may cause rapidly life-threatening infection.

Pregnancy Pregnancy affects decisions regarding antibacterial therapy in two respects. First, pregnancy is associated with an increased risk of particular infections (e.g., those caused by *Listeria*). Second, the potential risks to the fetus that are posed by specific drugs must be considered. As for other drugs, the safety of the vast majority of antibacterial agents in pregnancy has not been established, and such agents are grouped in categories B and C by the U.S. Food and Drug Administration. Drugs in categories D and X are contraindicated in pregnancy or lactation due to established risks. The risks associated with antibacterial use in pregnancy and during lactation are summarized in [Table 170-2](#).

Allergies Allergies to antibiotics are among the most common allergies reported, and an allergy history should be obtained whenever possible before therapy is chosen. A detailed allergy history can shed light on the type of reaction experienced previously and on whether rechallenge with the same or a related medication is advisable (and, if so, under what circumstances). Allergies to the penicillins are most common. Although as many as 10% of patients may report an allergy to penicillin, studies suggest that up to 90% of these patients could tolerate a penicillin or cephalosporin. Adverse effects ([Table 170-3](#)) should be distinguished from true allergies to ensure appropriate selection of antibacterial therapy.

Drug–Drug Interactions Patients commonly receive other drugs that may interact with antibacterial agents. A summary of the most common drug–drug interactions, by antibacterial class, is provided in [Table 170-4](#).

Exposures Exposures, both occupational and social, may provide clues to likely pathogens. When relevant, inquiries about exposure to ill contacts, animals, insects, and water should be included in the history, along with sites of residence and travel.

Other Host Factors Age, renal and hepatic function, and comorbid conditions are all considerations in the choice of and schedule for therapy. Dose adjustments should be made accordingly. In patients with decreased or unreliable oral absorption, IV therapy may be preferred to ensure adequate blood levels of drug and delivery of the antibacterial agent to the site of infection.

DURATION OF THERAPY

Whether empirical or directed, the duration of therapy should be planned in most clinical situations. Guidelines that synthesize available literature and expert opinion provide recommendations on therapy duration that are based on infecting organism, organ system, and patient factors. For example, the American Heart Association has published guidelines endorsed by the Infectious Diseases Society of America (IDSA) on diagnosis, antibacterial therapy, and management of complications of infective endocarditis. Similar guidelines from the IDSA exist for bacterial meningitis, catheter-associated urinary tract infections, intraabdominal infections, community- and hospital-acquired pneumonia, and other infections.

FAILURE OF THERAPY

If a patient does not respond to therapy, investigations often should include the collection of additional specimens for microbiologic testing and imaging as indicated. Failure to respond can be the result of an antibacterial regimen that does not address the underlying causative organism, the development of resistance during therapy, or the existence of a focus of infection at a site poorly penetrated by systemic therapy. Some infections may also require surgical interventions for cure (e.g., large abscesses, myonecrosis). Fever due to allergic drug reactions can sometimes complicate assessment of the patient's response to antibacterial treatment.

EXPERT GUIDANCE

Selected websites with the most up-to-date information and guidance for the clinician include the following:

- Johns Hopkins ABX Guide (www.hopkins-abxguide.org)
- IDSA Practice Guidelines (www.idsociety.org/IDSA_Practice_Guidelines/)
- Center for Disease Dynamics, Economics and Policy Resistance Map (www.cddep.org/map)
- CDC Antibiotic/Antimicrobial Resistance (www.cdc.gov/drug-resistance/)

CLINICAL USE OF ANTIBACTERIAL AGENTS

The clinical application of antibacterial therapy is guided by the spectrum of the agent and the suspected or known target pathogen. Infections for which specific antibacterial agents are among the drugs of choice are listed, along with associated pathogens and susceptibility data, in [Table 170-5](#). Resistance rates of specific organisms are dynamic and should be taken into account in the approach to antibacterial therapy. While national resistance rates can serve as a reference, the most useful reference for the clinician is the most recent local laboratory antibiogram, which provides details on local resistance patterns, often on an annual or semiannual basis.

β-LACTAMS

The β-lactam class of antibiotics consists of penicillins, cephalosporins, carbapenems, and monobactams. The term β-lactam reflects the drugs' four-membered lactam ring, which is their core structure. The differing

TABLE 170-2 RISKS ASSOCIATED WITH USE OF ANTIBACTERIAL DRUGS IN PREGNANCY AND LACTATION

Pregnancy Category ^a	Antibacterial Drug	Fetal Risk Recommendation ^b	Breast-Feeding Risk Recommendation ^b
B	Azithromycin	Limited human data. Animal data suggest low risk.	Limited human data; probably compatible
	Cephalosporins (including cephalexin, cefuroxime, cefixime, cefpodoxime, cefotaxime, ceftriaxone)	Compatible	Compatible
	Clindamycin	Compatible	Compatible
	Ertapenem	No human data; probably compatible	Limited human data; probably compatible
	Erythromycin	Compatible (excluding estolate salt)	Compatible
	Meropenem	No human data. Animal data suggest low risk.	No human data; probably compatible
	Metronidazole	Human data suggest low risk.	Interrupt breast-feeding for 12–24 h after single 2-g dose. Limited human data; potential toxicity in divided doses
	Nitrofurantoin	Human data suggest risk in third trimester.	Limited human data; probably compatible. Higher risk associated with younger infants and those with G6PD deficiency
	Penicillins (including amoxicillin, ampicillin, cloxacillin)	Compatible	Compatible
	Quinupristin-dalfopristin	Compatible. Maternal benefit must far outweigh risk to embryo/fetus.	No human data; potential toxicity
C	Vancomycin	Compatible	Limited human data; probably compatible
	Chloramphenicol	Compatible	Limited human data; potential toxicity
	Fluoroquinolones	Human data suggest low risk.	Limited human data; probably compatible
	Clarithromycin	Limited human data. Animal data suggest high risk.	No human data; probably compatible
	Imipenem/cilastatin	Limited human data. Animal data suggest low risk.	Limited human data; probably compatible
	Linezolid	Compatible. Maternal benefit must far outweigh risk to embryo/fetus.	No human data; potential toxicity
	Telavancin	No human data. Animal studies have revealed evidence of teratogenicity. ^c	No human data. Animal studies have revealed evidence of teratogenicity. ^c
	Tedizolid	Limited data. Embryo-fetal studies in mice, rats, and rabbits have demonstrated fetal developmental toxicities. Use only if benefit outweighs risk.	Excreted in the breastmilk of rats; unknown in humans; caution use
	Dalbavancin	Limited human data. At high doses in animal studies, delayed fetal maturation, increased embryo and offspring death. Use only if benefit outweighs risk.	Excreted in the breastmilk of animals; unknown in humans; caution use
	Oritavancin	Limited human data. Studies in rats and rabbits demonstrated no harm at 25% of recommended human dose. Use only if benefit outweighs risk.	Excreted in the breastmilk of rats; unknown in humans; caution use
C/D	Amikacin	Human data suggest low risk.	Compatible
	Gentamicin	Human data suggest low risk.	Compatible
D	Kanamycin	Human data suggest risk.	Limited human data; probably compatible
	Streptomycin	Human data suggest risk.	Compatible
	Sulfonamides	Human data suggest risk in third trimester.	Limited human data; potential toxicity. Avoid in ill, stressed, premature infants and in infants with hyperbilirubinemia or G6PD deficiency.
	Tetracyclines	Contraindicated in second and third trimesters	Compatible
	Tigecycline	Human data suggest risk in second and third trimesters.	No human data; potential toxicity

^a**Category B:** Either animal reproduction studies have failed to demonstrate a risk to the fetus, and there are no adequate and well-controlled studies in pregnant women; or animal studies have shown an adverse effect, but adequate and well-controlled studies in pregnant women have failed to demonstrate a risk to the fetus in any trimester. **Category C:** Animal reproduction studies have shown an adverse effect on the fetus, and there are no adequate and well-controlled studies in humans, but potential benefits may warrant use of the drug in pregnant women despite potential risks. **Category D:** There is positive evidence of human fetal risk based on adverse reaction data from investigational or marketing experience or studies in humans, but potential benefits may warrant use of the drug in pregnant women despite potential risks. ^bFetal risk recommendation and breast-feeding risk recommendation adapted from GG Briggs et al, eds: *Drugs in Pregnancy and Lactation*, 9th ed. Philadelphia, Lippincott Williams and Wilkins, 2011; and the U.S. Food and Drug Administration (Drugs@FDA).

^cA registry has been established to monitor pregnancy outcomes of pregnant women exposed to telavancin. Physicians are encouraged to register pregnant patients, or pregnant women may enroll themselves by calling 1-855-633-8479.

Abbreviation: G6PD, glucose-6-phosphate dehydrogenase.

side chains among the agents of this family determine the spectrum of activity. All β -lactams exert a bactericidal effect by inhibiting bacterial cell-wall synthesis. The β -lactams are classified as time-dependent killing agents; therefore, their clinical efficacy is best correlated with the proportion of the dosing interval during which the drug levels remain above the MIC for the pathogenic organism.

Penicillins and β -Lactamase Inhibitors Penicillin, the first β -lactam, was discovered in 1928 by Alexander Fleming. Natural penicillins, such

as penicillin G, are active against non- β -lactamase-producing gram-positive and gram-negative bacteria, anaerobes, and some gram-negative cocci. Penicillin G is used for penicillin-susceptible streptococcal infections, pneumococcal and meningococcal meningitis, enterococcal endocarditis, and syphilis. The antistaphylococcal penicillins, which have potent activity against methicillin-susceptible *S. aureus* (MSSA), include nafcillin, oxacillin, dicloxacillin, and flucloxacillin. Aminopenicillins, such as ampicillin and amoxicillin, provide added coverage beyond penicillin against gram-negative cocci, such as *Haemophilus influenzae*,

TABLE 170-3 COMMON ADVERSE REACTIONS TO ANTIBACTERIAL AGENTS

Antibacterial(s)	Potential Adverse Effects	Comments
β-Lactams	Hypersensitivity reactions	Range from rash to anaphylaxis. Cross-reactivity among β-lactams is related to chemical structure and side chain similarity.
	Neurotoxicity	More commonly described with cefepime and imipenem, but likely a class effect. Risk is increased in patients with history of seizures, renal impairment, and advanced age.
	Neutropenia/hematologic reactions	May be related to high doses and prolonged duration
Vancomycin	Nephrotoxicity	Risk increases with vancomycin trough levels >20 µg/mL or concomitant administration with other potentially nephrotoxic agents. The effect is usually reversible.
	“Red man syndrome”	Can be managed with a slower vancomycin infusion and pretreatment with antihistamine
Telavancin	QT prolongation	May falsely affect INR, PT, aPTT. Perform these tests before the next dose of telavancin (when serum drug levels are at their nadir).
	Interference with coagulation tests	
	Taste disturbances	
	Nephrotoxicity	
Oritavancin	Interference with coagulation tests	May falsely affect INR, PT, aPTT. Perform these tests at least 24 h after the dose was administered
Dalbavancin, oritavancin	Similar to vancomycin without “red man syndrome”	
Daptomycin	Myopathy	Monitor CPK levels during therapy. Rhabdomyolysis has been reported but appears to be rare.
	Eosinophilic pneumonia	
Aminoglycosides	Nephrotoxicity	Associated with prolonged use; usually reversible
	Ototoxicity	Can cause both vestibular and cochlear toxicity. Ototoxicity may be irreversible.
Fluoroquinolones	QTc prolongation	Moxifloxacin appears more likely than other quinolones to exert this effect. Risk of arrhythmia increases when these drugs are given concomitantly with other QTc-prolonging agents.
	Tendinitis	Risk is greater among elderly and patients receiving steroids.
	Dysglycemia	
	Exacerbation of myasthenia gravis	
Rifampin	Hepatotoxicity	Risk is greater when drug is given with other antituberculosis agents. When rifampin is given alone, LFT values may be transiently elevated without symptoms.
	Orange discoloration of body fluids	
Tetracyclines and glycyclines	Photosensitivity	High incidence of diarrhea, nausea, vomiting
	Gastrointestinal distress	
Macrolides	Gastrointestinal distress	Erythromycin is occasionally used as a therapeutic agent for some gastric motility disorders.
	QTc prolongation	Azithromycin use is associated with an increased risk of death from cardiovascular causes among patients at high baseline risk.
Metronidazole	Peripheral neuropathy	Associated with prolonged use
Clindamycin	Diarrhea and pseudomembranous colitis	
Linezolid, tedizolid	Myelosuppression	Associated with prolonged use
	Optic and peripheral neuropathy	Associated with prolonged use
TMP-SMX	Hypersensitivity reactions	Allergy usually associated with sulfonamide moiety
	Nephrotoxicity	Associated with high doses
	Hematologic effects	Associated with prolonged use
Nitrofurantoin	Pneumonitis and other pulmonary reactions	Associated with prolonged use
	Peripheral neuropathy	Associated with accumulation of nitrofurantoin in renal failure. Avoid use in renal impairment.
Fosfomycin	Gastrointestinal effects	
Polymyxins	Nephrotoxicity	Associated with high dose
	Neurotoxicity	Neuromuscular blockade and muscle weakness are well described and usually reversible.
Quinupristin-dalfopristin	Arthralgias and myalgias	
Chloramphenicol	Bone marrow suppression	Aplastic anemia or hematopoietic toxicity

Note: All systemic antibiotics have the potential to alter abdominal flora and induce *Clostridium difficile* infection.

Abbreviations: aPTT, activated partial thromboplastin time; CPK, creatine phosphokinase; INR, international normalized ratio; LFT, liver function test; PT, prothrombin time; TMP-SMX, trimethoprim-sulfamethoxazole.

and some Enterobacteriaceae, including *E. coli*, *Proteus mirabilis*, *Salmonella*, and *Shigella*. The aminopenicillins are hydrolyzed by many common β-lactamases. These drugs are commonly used for otitis media, respiratory tract infections, intraabdominal infections, endocarditis, meningitis, and urinary tract infections. The antipseudomonal penicillins include ticarcillin and piperacillin. These penicillin groups generally offer adequate anaerobic coverage; the exceptions are *Bacteroides*

species (such as *Bacteroides fragilis*), which produce β-lactamases and are generally resistant. The rising prevalence of β-lactamase-producing bacteria has led to the increased use of β-lactam/β-lactamase inhibitor combinations, such as ampicillin-sulbactam, amoxicillin-clavulanate, ticarcillin-clavulanate, and piperacillin-tazobactam. The β-lactamase inhibitors themselves do not have antibacterial activity (with the exception of sulbactam, which has activity against *Acinetobacter baumannii*)

TABLE 170-4 SIGNIFICANT ANTIBACTERIAL DRUG INTERACTIONS

Antibacterial(s)	Interacting Agent(s)	Potential Effect and Management
Nafcillin	Warfarin, cyclosporine, tacrolimus	Decreased levels of warfarin, cyclosporine via CYP3A4 induction. Monitor levels of affected drug closely if drugs are given concomitantly.
Ceftriaxone	Calcium-containing IV solutions	Concomitant use is contraindicated in neonates (<28 days); the combination can lead to precipitation of ceftriaxone-calcium particulate. Ceftriaxone and calcium-containing solutions can be given to infants >28 days of age provided they are given sequentially and the lines are thoroughly flushed between infusions.
Carbapenems	Valproic acid	Decreased levels of valproic acid. Monitor valproic acid levels closely if drugs are given concomitantly.
Linezolid, tedizolid	Serotonergic and adrenergic agents (e.g., SSRIs, vasopressors)	Increased levels of serotonergic and adrenergic agents. Monitor for serotonin syndrome. Tedizolid may have less potential than linezolid to cause this drug interaction.
Quinupristin dalfopristin	Substrates of cytochrome CYP3A4 (e.g., warfarin, ritonavir, cyclosporine, diazepam, verapamil)	Can result in increased levels of interacting drug. Erythromycin and clarithromycin are more potent CYP3A4 inhibitors than azithromycin. Avoid concomitant administration if possible.
Fluoroquinolones	Theophylline ^a	Can result in theophylline toxicity
	Sucralfate; antacids containing aluminum, calcium, or magnesium; ferrous sulfate and zinc-containing multivitamins	Can result in subtherapeutic fluoroquinolone levels. Administer fluoroquinolone 2 h before or 6 h after interacting drug.
	Tizanidine ^a	Can result in increased levels of tizanidine and hypotensive, sedative effects. Monitor for side effects if drugs are given concomitantly.
Rifampin	Substrates of cytochrome CYP3A4 (e.g., warfarin, ritonavir, cyclosporine, diazepam, verapamil, protease inhibitors, voriconazole)	Can result in decreased levels of interacting drug. Avoid concomitant use if possible. If giving drugs concomitantly, monitor drug levels if possible.
	Substrates of cytochrome CYP2C19 (e.g., omeprazole, lansoprazole)	
	Substrates of cytochrome CYP2C9 (e.g., warfarin, tolbutamide)	
	Substrates of cytochrome CYP2C8 (e.g., repaglinide, rosiglitazone)	
	Substrates of cytochrome CYP2B6 (e.g., efavirenz)	
	Hormone therapy (e.g., norethindrone)	
Tetracyclines	Antacids or drugs containing calcium, magnesium, iron, or aluminum	Can result in decreased absorption of tetracyclines. Administer tetracycline 2 h before or 6 h after interacting drug.
Macrolides ^b	Substrates of cytochrome CYP3A4 (e.g., warfarin, ritonavir, cyclosporine, diazepam, verapamil)	Avoid concomitant administration if possible.
	QTc-prolonging agents (e.g., fluoroquinolones, sotalol)	Increased risk of cardiotoxicity and arrhythmias. Monitor QTc.
	Protease inhibitors (e.g., ritonavir)	Can result in increased levels of both macrolides and protease inhibitors. Avoid concomitant use if possible.
Metronidazole	Cimetidine	Cimetidine can increase levels of macrolides.
	Ethanol	Can result in disulfiram-like reaction. Ethanol may be present in some formulations of oral drug suspensions (e.g., ritonavir).
TMP-SMX	Warfarin	Can increase warfarin levels. Monitor INR closely if drugs are given concomitantly.
	Phenytoin	Increased effect of warfarin. Monitor levels closely if drugs are given concomitantly.
	Methotrexate	Increased levels of phenytoin. Monitor levels closely if drugs are given concomitantly.
Oritavancin	Substrates of cytochrome CYP3A4 (e.g., cyclosporine, warfarin) and CYP2D6 (e.g., aripiprazole)	Increased levels of methotrexate. Monitor levels closely if drugs are given concomitantly.
	Substrates of cytochrome CYP2C19 (e.g., omeprazole) and CYP2C9 (e.g., warfarin)	Can result in decreased levels of interacting drug. Avoid concomitant use if possible. If giving drugs concomitantly, monitor drug levels if possible.
		Can result in increased levels of interacting drug. Avoid concomitant use if possible. If giving drugs concomitantly, monitor drug levels if possible.

^aDrug reaction described with ciprofloxacin only. ^bClarithromycin and erythromycin are potent CYP3A4 inhibitors; the probability of a drug interaction with azithromycin is lower.

Abbreviations: INR, international normalized ratio; SSRI, selective serotonin-reuptake inhibitor; TMP-SMX, trimethoprim-sulfamethoxazole.

but typically inhibit the *S. aureus* class A β -lactamase, β -lactamases of *H. influenzae* and *Bacteroides* species, and a number of plasmid-encoded β -lactamases. These combination agents are typically used when broader-spectrum coverage is needed—e.g., in pneumonia and intraabdominal infections. Piperacillin-tazobactam is a useful agent for broad coverage in febrile neutropenic patients. The combination

agents, however, are not effective against organisms that produce AmpC β -lactamases or carbapenemases.

Cephalosporins The cephalosporin drug class encompasses five generations determined by spectrum of antibacterial activity. The first generation (cefazolin, cefadroxil, cephalexin) largely has activity

TABLE 170-5 DRUG INDICATIONS FOR SPECIFIC INFECTIONS, ASSOCIATED PATHOGENS, AND SAMPLE SUSCEPTIBILITY RATES

Antimicrobial(s)	Infections	Common Pathogens (% Susceptible); Resistance as Noted ^a
Penicillin G	Syphilis; yaws; leptospirosis; streptococcal infections; pneumococcal infections; actinomycosis; oral and periodontal infections; meningococcal meningitis and meningococcemia; viridans streptococcal endocarditis; clostridial myonecrosis; tetanus; rat-bite fever; <i>Pasteurella multocida</i> infections; erysipeloid (<i>Erysipelothrix rhusiopathiae</i>)	<i>Neisseria meningitidis</i> ; viridans streptococci (73%); <i>Streptococcus pneumoniae</i> (92% nonmeningitis; 65% meningitis)
Ampicillin, amoxicillin	Salmonellosis; acute otitis media; <i>Haemophilus influenzae</i> meningitis and epiglottitis; <i>Listeria monocytogenes</i> meningitis; <i>Enterococcus faecalis</i> UTI	<i>Escherichia coli</i> (52%); <i>H. influenzae</i> (70%); <i>Salmonella</i> spp. (91%)
Nafcillin, oxacillin	MSSA bacteremia and endocarditis	<i>Staphylococcus aureus</i> (68%); coagulase-negative staphylococci (47%)
Piperacillin-tazobactam	Intraabdominal infections (facultative enteric gram-negative bacilli and obligate anaerobes); infections caused by mixed flora (aspiration pneumonia, diabetic foot ulcers); infections caused by <i>Pseudomonas aeruginosa</i>	<i>P. aeruginosa</i> (88%) ^b
Cefazolin	<i>E. coli</i> UTI; surgical prophylaxis; MSSA bacteremia and endocarditis	<i>E. coli</i> (85%)
Cefoxitin, cefotetan	Intraabdominal infections and pelvic inflammatory disease	<i>Bacteroides fragilis</i> (60%) ^c
Ceftriaxone	Gonococcal infections; pneumococcal meningitis; viridans streptococcal endocarditis; salmonellosis and typhoid fever; hospital-acquired infections caused by nonpseudomonal facultative gram-negative enteric bacilli	<i>S. pneumoniae</i> (93%); ^d <i>E. coli</i> (93%); <i>Klebsiella pneumoniae</i> (89%)
Ceftazidime, cefepime	Hospital-acquired infections caused by facultative gram-negative bacilli and <i>Pseudomonas</i> spp.	<i>P. aeruginosa</i> (89%)
Ceftaroline	CAP caused by <i>S. pneumoniae</i> , MSSA, <i>H. influenzae</i> , <i>K. pneumoniae</i> , <i>Klebsiella oxytoca</i> , and <i>E. coli</i> ; acute bacterial skin and skin-structure infections caused by MSSA, MRSA, <i>Streptococcus pyogenes</i> , <i>Streptococcus agalactiae</i> , <i>E. coli</i> , <i>K. pneumoniae</i> , and <i>K. oxytoca</i>	Mostly susceptible; four strains of MRSA with ceftaroline MICs >4 µg/mL reported in isolates from a single Greek hospital ^e
Imipenem, meropenem	Intraabdominal infections, infections caused by <i>Enterobacter</i> spp. and ESBL-producing gram-negative bacilli	<i>P. aeruginosa</i> (76% and 83%); <i>Acinetobacter calcoaceticus-baumannii</i> complex (81% and 82%)
Ertapenem	CAP; complicated UTIs, including pyelonephritis; acute pelvic infections; complicated intraabdominal infections; complicated skin and skin-structure infections, excluding diabetic foot infections accompanied by osteomyelitis or caused by <i>P. aeruginosa</i>	<i>Enterobacter cloacae</i> (87%); <i>K. pneumoniae</i> (97%)
Aztreonam	HAIs caused by facultative gram-negative bacilli and <i>Pseudomonas</i> in penicillin-allergic patients	<i>P. aeruginosa</i> (76%)
Vancomycin	Bacteremia, endocarditis, and other invasive disease caused by MRSA; pneumococcal meningitis; oral formulation for CDAD	<i>S. aureus</i> (100%); <i>E. faecalis</i> (89%); <i>E. faecium</i> (24%)
Telavancin	Hospital- and ventilator-associated pneumonia or skin and soft tissue infections caused by MRSA	<i>S. aureus</i> (100%)
Dalbavancin, oritavancin	Complicated skin and soft tissue infections	<i>S. aureus</i> (100%)
Daptomycin	VRE infections; MRSA bacteremia	<i>E. faecalis</i> (99.9%); ^f <i>E. faecium</i> (99.7%); ^f <i>S. aureus</i> (99.9%) ^f
Gentamicin, tobramycin, amikacin	Combined with penicillin for staphylococcal, enterococcal, or streptococcal endocarditis; combined with β-lactam for gram-negative bacteremia; pyelonephritis	<i>E. coli</i> (gentamicin, 91%); <i>P. aeruginosa</i> (amikacin, 87%; gentamicin, 81%); <i>A. calcoaceticus-baumannii</i> complex (amikacin, 68%; gentamicin, 83%)
Azithromycin, clarithromycin, erythromycin	<i>Legionella</i> , <i>Campylobacter</i> , and <i>Mycoplasma</i> infections; CAP; GAS pharyngitis in penicillin-allergic patients; bacillary angiomatosis; gastric infections due to <i>Helicobacter pylori</i> ; MAI infections	<i>S. pneumoniae</i> (59%); group A streptococci (78%); <i>H. pylori</i> (75%) ^g
Clindamycin	Severe, invasive GAS infections (with β-lactam); infections caused by obligate anaerobes; infections caused by susceptible staphylococci	<i>S. aureus</i> (67%)
Doxycycline, minocycline	Acute bacterial exacerbations of chronic bronchitis; granuloma inguinale; brucellosis (with streptomycin); tularemia; glanders; melioidosis; spirochetal infections caused by <i>Borrelia</i> (Lyme disease and relapsing fever; doxycycline); infections caused by <i>Vibrio vulnificus</i> ; some <i>Aeromonas</i> infections; infections due to <i>Stenotrophomonas</i> (minocycline); plague; ehrlichiosis; chlamydial infections (doxycycline); granulomatous infections due to <i>Mycobacterium marinum</i> (minocycline); rickettsial infections; mild CAP; skin and soft tissue infections caused by gram-positive cocci (e.g., CA-MRSA infections); leptospirosis; syphilis; and actinomycosis in the penicillin-allergic patient	<i>S. pneumoniae</i> (75%); <i>S. aureus</i> (94%)
Tigecycline	CAP caused by <i>S. pneumoniae</i> , <i>H. influenzae</i> , or <i>Legionella pneumophila</i> ; complicated skin infections caused by <i>E. coli</i> , MRSA, MSSA, <i>S. pyogenes</i> , <i>Streptococcus anginosus</i> , <i>S. agalactiae</i> , <i>B. fragilis</i> ; complicated intraabdominal infections caused by <i>E. coli</i> , vancomycin-susceptible <i>E. faecalis</i> , <i>Citrobacter freundii</i> , <i>Enterobacter cloacae</i> , <i>K. pneumoniae</i> , <i>K. oxytoca</i> , <i>Bacteroides</i> spp., <i>Clostridium perfringens</i> , and <i>Peptostreptococcus</i> spp.	Mostly susceptible, though case reports of resistance in <i>A. baumannii</i> and <i>K. pneumoniae</i>

(Continued)

TABLE 170-5 DRUG INDICATIONS FOR SPECIFIC INFECTIONS, ASSOCIATED PATHOGENS, AND SAMPLE SUSCEPTIBILITY RATES (CONTINUED)

Antimicrobial(s)	Infections	Common Pathogens (% Susceptible); Resistance as Noted ^a
TMP-SMX	Community-acquired UTI; CA-MRSA skin and soft tissue infections	<i>E. coli</i> (73%); <i>S. aureus</i> (96%)
Sulfonamides	Nocardial infections; leprosy (dapson); toxoplasmosis (sulfadiazine)	Unknown
Ciprofloxacin, levofloxacin, moxifloxacin	CAP (levofloxacin and moxifloxacin); UTI; bacterial gastroenteritis; hospital-acquired gram-negative enteric infections; <i>Pseudomonas</i> infections (ciprofloxacin and levofloxacin)	<i>S. pneumoniae</i> (99%); <i>E. coli</i> (80%); <i>P. aeruginosa</i> (ciprofloxacin, 72%; levofloxacin, 69%); <i>Salmonella</i> spp. (ciprofloxacin, 98%; levofloxacin, 100%)
Rifampin	Staphylococcal foreign body infections in combination with other antistaphylococcal agents; <i>Legionella</i> pneumonia; <i>Mycobacterium tuberculosis</i> ; atypical nontuberculous mycobacterial infection; pneumococcal meningitis when organisms are susceptible or response is delayed	<i>S. aureus</i> (99%), although staphylococci rapidly develop resistance with monotherapy
Metronidazole	Obligate anaerobic gram-negative bacteria (e.g., <i>Bacteroides</i> spp); abscess in lung, brain, or abdomen; bacterial vaginosis; CDAD	Mostly susceptible; resistance very rare
Linezolid, tedizolid	VRE; uncomplicated and complicated skin and soft tissue infections caused by MSSA and MRSA; CAP with concurrent bacteremia; hospital-acquired pneumonia	Mostly susceptible; resistance occasionally seen in VRE
Chloramphenicol	HAI due to gram-positive and gram-negative organisms resistant to standard alternatives (e.g., <i>Burkholderia</i>)	Unknown
Colistin	HAI due to gram-negative bacilli resistant to all other chemotherapy (e.g., <i>P. aeruginosa</i> , <i>Acinetobacter</i> spp., and <i>Stenotrophomonas maltophilia</i>)	<i>P. aeruginosa</i> (case reports, outbreaks)
Quinupristin-dalfopristin	VRE; complicated skin and skin-structure infections due to MSSA and <i>S. pyogenes</i>	<i>E. faecalis</i> (<20%); ^h <i>E. faecium</i> (>90%) ^h
Mupirocin	Topical application to nares for <i>S. aureus</i> decolonization	<i>S. aureus</i> (74–100%) ⁱ
Nitrofurantoin	UTI caused by most gram-negative bacilli and some gram-positive organisms; prophylaxis in recurrent cystitis	<i>E. coli</i> (92%); <i>E. faecalis</i> (99%)
Fosfomycin	UTI caused by most gram-negative bacilli and some gram-positive organisms; prophylaxis in recurrent cystitis	Unknown

^aUnless otherwise noted, susceptibility rates are based on isolates from the Massachusetts General Hospital Clinical Microbiology Laboratory collected between January and December 2012. Local rates will vary. ^bThe Center for Disease Dynamics, Economics and Policy Resistance Map, Washington, DC. ^cS Sepehri et al: Prevalence of antimicrobial resistance among clinical isolates of *Bacteroides fragilis* group in Canada in 2010–2011: CANWARD Surveillance Study. Abstract C2-1814, presented at the 51st Interscience Conference on Antimicrobial Agents and Chemotherapy, 2011. Available at www.can-c.com/posters/ICAAC2011/Sepehri%20Prevalence%20Bfragilis%20ICAAC2011.pdf. ^dGV Doern et al: Clin Infect Dis 41:139, 2005. ^eR Mendes et al: J Antimicrob Chemother 67:1321, 2012. ^fHS Sader et al: J Chemother 23:200, 2011. ^gJ Torres et al: J Clin Microbiol 39:2677, 2001. ^hWS Oh et al: Antimicrob Agents Chemother 49:5176, 2005. ⁱAE Simor et al: Antimicrob Agents Chemother 51:3880, 2007.

Abbreviations: CA-MRSA, community-acquired MRSA; CAP, community-acquired pneumonia; CA-UTI, community-acquired UTI; CDAD, *Clostridium difficile*-associated diarrhea; ESBL, extended-spectrum β -lactamase; GAS, group A streptococcal; HAI, hospital-acquired infection; MAI, *Mycobacterium avium-intracellulare*; MIC, minimal inhibitory concentration; MRSA, methicillin-resistant *Staphylococcus aureus*; MSSA, methicillin-susceptible *S. aureus*; TMP-SMX, trimethoprim-sulfamethoxazole; UTI, urinary tract infection; VRE, vancomycin-resistant *Enterococcus*.

against gram-positive bacteria, with some additional activity against *E. coli*, *P. mirabilis*, and *K. pneumoniae*. First-generation cephalosporins are commonly used for infections caused by MSSA and streptococci (e.g., skin and soft tissue infections). Cefazolin is a popular choice for surgical prophylaxis against skin organisms. The second generation (cefamandole, cefuroxime, cefaclor, cefprozil, cefuroxime axetil, cefoxitin, cefotetan) has additional activity against *H. influenzae* and *Moraxella catarrhalis*. Cefoxitin and cefotetan have potent activity against anaerobes as well. Second-generation cephalosporins are used to treat community-acquired pneumonia because of their activity against *S. pneumoniae*, *H. influenzae*, and *M. catarrhalis*. They are also used for other mild or moderate infections, such as acute otitis media and sinusitis. The third-generation cephalosporins are characterized by greater potency against gram-negative bacilli and reduced potency against gram-positive cocci. These cephalosporins, which include cefoperazone, cefotaxime, ceftazidime, ceftriaxone, cefdinir, cefixime, and cefpodoxime, are used for infections caused by Enterobacteriaceae, although resistance is an increasing concern. It is noteworthy that ceftazidime is the only third-generation cephalosporin with activity against *P. aeruginosa* but lacks activity against gram-positive bacteria. This drug is frequently used for pulmonary infections in cystic fibrosis and febrile neutropenia. Ceftriaxone penetrates the CSF and can be used to treat meningitis caused by *H. influenzae*, *N. meningitidis*, and susceptible strains of *S. pneumoniae*. It is also used for the treatment of later-stage Lyme disease. The fourth generation includes cefepime and ceftipime, broad-coverage agents that provide potent activity against both gram-negative bacilli, including *P. aeruginosa*, and gram-positive cocci. The fourth generation has clinical applications similar to those of the third generation and can be used in bacteremia, pneumonia, skin and soft tissue infections, and urinary tract infections caused by

susceptible bacteria. Cefepime is also commonly used in febrile neutropenia. Ceftaroline, a fifth-generation cephalosporin, differs from the other cephalosporins in its added activity against MRSA, which is resistant to all other β -lactams. Ceftaroline's gram-negative activity is similar to that of the third-generation cephalosporins but does not include *P. aeruginosa*. Ceftaroline is efficacious in community-acquired pneumonia and skin infections, but few data are available on its use for more serious infections, such as bacteremia.

Carbapenems With a few exceptions for cefepime, all penicillins and cephalosporins are ineffective in the presence of ESBLs. Carbapenems, including doripenem, imipenem, meropenem, and ertapenem, offer the most reliable coverage for strains containing ESBLs. All carbapenems have broad activity against gram-positive cocci, gram-negative bacilli, and anaerobes. None is active against MRSA, but all are active against MSSA, *Streptococcus* species, and Enterobacteriaceae. Ertapenem is the only carbapenem that has poor activity against *P. aeruginosa* and *Acinetobacter*. Imipenem is active against penicillin-susceptible *Enterococcus faecalis* but not *Enterococcus faecium*. Carbapenems are not active against Enterobacteriaceae containing carbapenemases. *Stenotrophomonas maltophilia* and some *Bacillus* species are intrinsically resistant to carbapenems because of a zinc-dependent carbapenemase.

Monobactams Aztreonam is the sole monobactam. Its activity is limited to gram-negative bacteria and includes *P. aeruginosa* and most other Enterobacteriaceae. This drug is inactivated by ESBLs and carbapenemases. The principal use for aztreonam is as an alternative to penicillins, cephalosporins, or carbapenems in patients with serious β -lactam allergy. Aztreonam is structurally related to ceftazidime and should be used cautiously in individuals with a serious ceftazidime allergy. It is commonly used in febrile neutropenia and intraabdominal

942 infections. Aztreonam does not penetrate the CSF and should not be used for treatment of meningitis.

Adverse Reactions to β -Lactam Drugs Agents within the β -lactam class are known for several adverse effects. Gastrointestinal side effects, mainly diarrhea, are common, but hypersensitivity reactions constitute the most common adverse effect of β -lactams. The reactions' severity can range from rash to anaphylaxis, but the rate of true anaphylactic reactions is only 0.05%. An individual with an accelerated IgE-mediated reaction to one β -lactam agent may still receive another agent within the class, but caution should be taken to choose a β -lactam that has a dissimilar side chain and a low level of cross-reactivity. For example, the second-, third-, and fourth-generation cephalosporins and the carbapenems display very low cross-reactivity in patients with penicillin allergy. Aztreonam is the only β -lactam that has no cross-reactivity with the penicillin group. In cases of severe allergy, desensitization (a graded challenge) to the indicated β -lactam, with close monitoring, may be warranted if other antibacterial options are not suitable.

β -Lactams can rarely cause serum sickness, Stevens-Johnson syndrome, nephropathy, hematologic reactions, and neurotoxicity. Neutropenia appears to be related to high doses or prolonged use. Neutropenia and interstitial nephritis caused by β -lactams generally resolve upon discontinuation of the agent. Imipenem and cefepime are associated with an increased risk of seizure, but this risk is likely a class effect and related to high doses or doses that are not adjusted in renal impairment.

GLYCOPEPTIDES

The glycopeptide antibiotics include vancomycin and telavancin. Vancomycin has activity against staphylococci (including MRSA and coagulase-negative staphylococci), streptococci (including *S. pneumoniae*), and enterococci. It is not active against gram-negative organisms. Vancomycin also displays activity against *Bacillus* species, *Corynebacterium jeikeium*, *Listeria monocytogenes*, and gram-positive anaerobes such as *Peptostreptococcus*, *Actinomyces*, *Clostridium*, and *Propionibacterium* species. Vancomycin has several important clinical uses. It is used for serious infections caused by MRSA, including health care-associated pneumonia, bacteremia, osteomyelitis, and endocarditis. It is also commonly used for skin and soft tissue infections. Oral vancomycin is not absorbed systemically and is reserved for the treatment of *Clostridium difficile* infection. Vancomycin is also an alternative for the treatment of infections caused by MSSA in patients who cannot tolerate β -lactams. Resistance to vancomycin is a rising concern. Strains of vancomycin-intermediate *S. aureus* (VISA) and vancomycin-resistant enterococci (VRE) are not uncommon. Vancomycin appears to be a concentration-dependent killer, with AUC/MIC ratio being the best predictor of efficacy (Fig. 170-2). Guidelines recommend targeting a vancomycin trough level of 15–20 $\mu\text{g/mL}$ in MRSA infections in order to maintain an AUC/MIC ratio >400 . When using vancomycin, clinicians should monitor for nephrotoxicity. The risk increases when trough levels are $>20 \mu\text{g/mL}$. Concomitant therapy with other nephrotoxic agents, such as aminoglycosides, also increases the risk of nephrotoxicity. Ototoxicity was reported with early formulations of vancomycin but is currently uncommon because purer formulations are available. Both of these adverse effects are reversible upon discontinuation of vancomycin. Clinicians should be aware of the “red man syndrome,” a common reaction that presents as a rapid onset of erythematous rash or pruritus on the head, face, neck, and upper trunk. This reaction is caused by histamine release from basophils and mast cells and can be treated with diphenhydramine and slowing of the vancomycin infusion.

Telavancin, dalbavancin, and oritavancin are structurally similar to vancomycin and are referred to as *lipoglycopeptides*. They have antibacterial activity against *S. aureus* (including MRSA and some strains of VISA and vancomycin-resistant *S. aureus* [VRSA]), streptococci, and enterococci. They also have good activity against anaerobic gram-positive organisms except for *Lactobacillus* and some *Clostridium* species. The clinical efficacy of telavancin has been demonstrated in both skin and soft tissue infections and nosocomial pneumonia, and the efficacy of dalbavancin and oritavancin has been shown in skin and soft

tissue infections. The vancomycin resistance phenotype may reduce the potency of all three lipoglycopeptides, but the rate of resistance to these drugs among *S. aureus* and enterococci has been low. Adverse effects of telavancin include insomnia, a metallic taste, nephrotoxicity, and gastrointestinal side effects. Clinicians should be aware of the potential for electrocardiographic QTc prolongation that can increase the risk of cardiac arrhythmias when telavancin is used concomitantly with other QTc-prolonging agents. Telavancin may interfere with certain coagulation tests (e.g., causing false elevations in prothrombin time). Dalbavancin and oritavancin have safety profiles similar to that of vancomycin.

LIPOPEPTIDES

Daptomycin is a lipopeptide antibiotic with activity against a broad range of gram-positive organisms. This drug is active against staphylococci (including MRSA and coagulase-negative staphylococci), streptococci, and enterococci. Daptomycin remains active against enterococci that are resistant to vancomycin. In addition, it exhibits activity against *Bacillus*, *Corynebacterium*, *Peptostreptococcus*, and *Clostridium* species. Daptomycin's pharmacodynamic parameter for efficacy is concentration-dependent killing. Resistance to daptomycin is rare, but MICs may be higher for VISA strains. Daptomycin is efficacious in skin and soft tissue infections, bacteremia, endocarditis, and osteomyelitis. It is an important alternative for MRSA and other gram-positive infections when bactericidal therapy is needed and vancomycin cannot be used. Daptomycin is generally well tolerated, and its main toxicity consists of elevation of creatinine phosphokinase (CPK) levels and myopathy. CPK should be monitored during daptomycin treatment, and the drug should be discontinued if muscular toxicities occur. There have also been case reports of reversible eosinophilic pneumonia associated with daptomycin use.

AMINOGLYCOSIDES

The aminoglycosides are a class of antibacterial agents with concentration-dependent activity against most gram-negative organisms. The most commonly used aminoglycosides are gentamicin, tobramycin, and amikacin, although others, such as streptomycin, kanamycin, neomycin, and paromomycin, may be used in special circumstances. Aminoglycosides have a significant dose-dependent post-antibiotic effect, meaning that they have an antibacterial effect even after serum drug levels are undetectable. The postantibiotic effect and concentration-dependent killing form the rationale behind extended-interval aminoglycoside dosing, in which a larger dose is given once daily rather than smaller doses multiple times daily. Aminoglycosides are active against gram-negative bacilli, such as Enterobacteriaceae, *P. aeruginosa*, and *Acinetobacter*. They also enhance the activity of cell wall-active agents such as β -lactams or vancomycin in some gram-positive bacteria, including staphylococci and enterococci. This combination therapy is termed *synergistic* because the effect of both agents provides a killing effect greater than would be predicted from the effects of either agent alone. Amikacin and streptomycin have activity against *Mycobacterium tuberculosis*, and amikacin has activity against *Mycobacterium avium-intracellulare*. The aminoglycosides do not have activity against anaerobes, *S. maltophilia*, or *Burkholderia cepacia*. Aminoglycosides are used in clinical practice in a variety of infections caused by gram-negative organisms, including bacteremia and urinary tract infections. They are frequently used alone or in combination for the treatment of *P. aeruginosa* infection. When used in combination with a cell wall-active agent, gentamicin and streptomycin are also important for the treatment of gram-positive bacterial endocarditis. All aminoglycosides can cause nephrotoxicity and ototoxicity. The risk of nephrotoxicity is related to the dose and duration of therapy as well as the concomitant use of other nephrotoxic agents. Nephrotoxicity is usually reversible, but ototoxicity can be irreversible.

MACROLIDES AND KETOLIDES

The macrolides (azithromycin, clarithromycin, erythromycin) and ketolides (telithromycin) are classes of antibiotics that inhibit protein synthesis. Compared with erythromycin (the older antibiotic), azithromycin and clarithromycin have better oral absorption and tolerability.

Azithromycin, clarithromycin, and telithromycin all have broader spectra of activity than erythromycin, which is less frequently used. These agents are commonly used in the treatment of upper and lower respiratory tract infections caused by *S. pneumoniae*, *H. influenzae*, *M. catarrhalis*, and atypical organisms (e.g., *Chlamydia pneumoniae*, *Legionella pneumophila*, and *Mycoplasma pneumoniae*); group A streptococcal pharyngitis in penicillin-allergic patients; and nontuberculous mycobacterial infections (e.g., caused by *M. marinum* and *M. chelonae*) as well as in the prophylaxis and treatment of *M. avium-intracellulare* infection in patients with HIV/AIDS and in combination therapy for *H. pylori* infection and bartonellosis. Enterobacteriaceae, *Pseudomonas* species, and *Acinetobacter* species are intrinsically resistant to macrolides as a result of decreased membrane permeability, although azithromycin is active against gram-negative diarrheal pathogens. The major adverse effects of this drug class include nausea, vomiting, diarrhea and abdominal pain, prolongation of QT_c interval, exacerbation of myasthenia gravis, and tinnitus. Azithromycin specifically has been associated with an increased risk of death, especially among patients with underlying heart disease, because of the risk of QT_c interval prolongation and torsades de pointes. Erythromycin, clarithromycin, and telithromycin inhibit the CYP3A4 hepatic drug-metabolizing enzyme and can result in increased levels of coadministered drugs, including benzodiazepines, statins, warfarin, cyclosporine, and tacrolimus. Azithromycin does not inhibit CYP3A4 and lacks these drug–drug interactions.

CLINDAMYCIN

Clindamycin is a lincosamide antibiotic and is bacteriostatic against some organisms and bactericidal against others. It is used most often to treat bacterial infections caused by anaerobes (e.g., *B. fragilis*, *Clostridium perfringens*, *Fusobacterium* species, *Prevotella melaninogenicus*, and *Peptostreptococcus* species) and susceptible staphylococci and streptococci. Clindamycin is used for treatment of dental infections, anaerobic lung abscess, and skin and soft tissue infections. It is used together with bactericidal agents (penicillins or vancomycin) to inhibit new toxin synthesis in the treatment of streptococcal or staphylococcal toxic shock syndrome. Other uses include treatment of infections caused by *Capnocytophaga canimorsus*, a component of combination therapy for malaria and babesiosis, and therapy for toxoplasmosis. Clindamycin has excellent oral bioavailability. Adverse effects include nausea, vomiting, diarrhea, *C. difficile*-associated diarrhea and pseudomembranous colitis, maculopapular rash, and (rarely) Stevens-Johnson syndrome.

TETRACYCLINES AND GLYCYLCYCLINES

The tetracyclines (doxycycline, minocycline, and tetracycline) and the glycylicyclines (tigecycline) inhibit protein synthesis and are bacteriostatic. These drugs have wide clinical uses. They are used in the treatment of skin and soft tissue infections caused by gram-positive cocci (including MRSA), spirochetal infections (e.g., Lyme disease, syphilis, leptospirosis, and relapsing fever), rickettsial infections (e.g., Rocky Mountain spotted fever), atypical pneumonia, sexually transmitted infections (e.g., *Chlamydia trachomatis* infection, lymphogranuloma venereum, and granuloma inguinale), infections with *Nocardia* and *Actinomyces*, brucellosis, tularemia, Whipple's disease, and malaria. Tigecycline, the only approved agent in the glycylicycline class, is a derivative of minocycline and is indicated in the treatment of infections due to MRSA, vancomycin-sensitive enterococci, many Enterobacteriaceae, and *Bacteroides* species. Tigecycline has no activity against *P. aeruginosa*. It has been used in combination with colistin for the treatment of serious infections with multidrug-resistant gram-negative organisms. A pooled analysis of 13 clinical trials found an increased risk of death and treatment failure among patients treated with tigecycline alone. Tetracyclines have reduced absorption when coadministered with calcium- and iron-containing compounds, including milk, and doses should be spaced at least 2 h apart. The major adverse reactions to both of these classes are nausea, vomiting, diarrhea, and photosensitivity. Tetracyclines have been associated with fetal bone-growth abnormalities and should be avoided during pregnancy and in the treatment of children <8 years old.

TRIMETHOPRIM-SULFAMETHOXAZOLE

Trimethoprim-sulfamethoxazole (TMP-SMX) is an antibiotic whose two components both inhibit folate synthesis and produce antibacterial activity. TMP-SMX is active against gram-positive bacteria such as staphylococci and streptococci; however, its use against MRSA is usually limited to community-acquired infections, and its activity against *Streptococcus pyogenes* may not be reliable. TMP-SMX is also active against many gram-negative bacteria, including *H. influenzae*, *E. coli*, *P. mirabilis*, *N. gonorrhoeae*, and *S. maltophilia*. TMP-SMX does not have activity against anaerobes or *P. aeruginosa*. It has many uses because of its wide spectrum of activity and high oral bioavailability. Urinary tract infections, skin and soft tissue infections, and respiratory tract infections are among the common uses. Another important indication is for both prophylaxis and treatment of *Pneumocystis jirovecii* infections in immunocompromised patients. Resistance to TMP-SMX has limited its use against many Enterobacteriaceae. Resistance rates among urinary isolates of *E. coli* are almost 25% in the United States. The most common adverse reactions associated with TMP-SMX are gastrointestinal effects such as nausea, vomiting, and diarrhea. In addition, rash is a common allergic reaction and may preclude the subsequent use of other sulfonamides. With prolonged use, leukopenia, thrombocytopenia, and granulocytopenia can develop. TMP-SMX can also cause nephrotoxicity, hyperkalemia, and hyponatremia, which are more common at high doses. TMP-SMX has several important interactions with other drugs (Table 170-4), including warfarin, phenytoin, and methotrexate.

FLUOROQUINOLONES

The fluoroquinolones include norfloxacin, ciprofloxacin, ofloxacin, levofloxacin, moxifloxacin, and gemifloxacin. Ciprofloxacin and levofloxacin have the broadest spectrum of activity against gram-negative bacteria, including *P. aeruginosa* (similar to that of third-generation cephalosporins). Because of the risk of selection of resistance during fluoroquinolone treatment of serious pseudomonal infections, these agents are usually used in combination with an antipseudomonal β -lactam. Levofloxacin, moxifloxacin, and gemifloxacin have additional gram-positive activity, including that against *S. pneumoniae* and some strains of MSSA, and are used for treatment of community-acquired pneumonia. Strains of MRSA are commonly resistant to all fluoroquinolones. Moxifloxacin is used as one component of second-line regimens for multidrug-resistant tuberculosis. Fluoroquinolones exhibit concentration-dependent killing, are well absorbed orally, and have elimination half-lives that usually support once- or twice-daily dosing. Oral coadministration with compounds containing high concentrations of aluminum, magnesium, or calcium can reduce fluoroquinolone absorption. Their penetration into prostate tissue supports their use for bacterial prostatitis. Fluoroquinolones are generally well tolerated but can cause CNS stimulatory effects, including seizures; glucose dysregulation; and tendinopathy associated with Achilles tendon rupture, particularly in older patients, organ transplant recipients, and patients taking glucocorticoids. Worsening of myasthenia gravis also has been associated with quinolone use. Moxifloxacin causes modest prolongation of the QT_c interval and should be used with caution in patients receiving other QT_c-prolonging drugs.

RIFAMYCINS

The rifamycins include rifampin, rifabutin, and rifapentine. Rifampin is the most commonly used rifamycin. For almost all therapeutic indications, it is used in combination with other agents to reduce the likelihood of selection of high-level rifampin resistance. Rifampin is used foremost in the treatment of mycobacterial infections—specifically, as a mainstay of combination therapy for *M. tuberculosis* infection or as a single agent in the treatment of latent *M. tuberculosis* infection. In addition, it is often used in the treatment of nontuberculous mycobacterial infection. Rifampin is used in combination regimens for the treatment of staphylococcal infections, particularly prosthetic valve endocarditis and bone infections with retained hardware. It is a component of combination therapy for brucellosis (with doxycycline) and leprosy (with dapsone for tuberculoid leprosy and with dapsone

944 and clofazimine for lepromatous disease). Rifampin can be used alone for prophylaxis in close contacts of patients with *H. influenzae* or *N. meningitidis* meningitis. The drug has high oral bioavailability, which is further enhanced when it is taken on an empty stomach. Rifampin has several adverse effects, including elevated aminotransferase levels (14%), rash (1–5%), and gastrointestinal events such as nausea, vomiting, and diarrhea (1–2%). Its many clinically relevant interactions with other drugs mandate the clinician's careful review of the patient's medications before rifampin initiation to assess safety and the need for additional monitoring.

METRONIDAZOLE

Metronidazole is used in the treatment of anaerobic bacterial infections as well as infections caused by protozoa (e.g., amebiasis, giardiasis, trichomoniasis). It is the agent of choice as a component of combination therapy for polymicrobial abscesses in the lung, brain, or abdomen, the etiology of which often includes anaerobic bacteria, and for bacterial vaginosis, pelvic inflammatory disease, mild to moderate *C. difficile*-associated diarrhea, and anaerobic infections, such as those due to *Bacteroides*, *Fusobacterium*, and *Prevotella* species. Metronidazole is bactericidal against anaerobic bacteria and exhibits concentration-dependent killing. It has high oral bioavailability and tissue penetration, including penetration of the blood–brain barrier. The majority of *Actinomyces*, *Propionibacterium*, and *Lactobacillus* species are intrinsically resistant to metronidazole. The major adverse effects include nausea, diarrhea, and a metallic taste. Concomitant ingestion of alcohol may result in a disulfiram-like reaction, and patients are usually instructed to avoid alcohol during treatment. Long-term treatment carries the risk of leukopenia, neutropenia, peripheral neuropathy, and central nervous system toxicity manifesting as confusion, dysarthria, ataxia, nystagmus, and ophthalmoparesis. Through metronidazole's effect on the CYP2C9 drug-metabolizing enzyme, its coadministration with warfarin can result in decreased metabolism and enhanced anticoagulant effects that require close monitoring. Concomitant administration of metronidazole with lithium can result in increased serum levels of lithium and associated toxicity; coadministration with phenytoin can result in phenytoin toxicity and possibly decreased levels of metronidazole.

OXAZOLIDINONES

Linezolid is a bacteriostatic agent and is indicated for serious infections due to resistant gram-positive bacteria, such as MRSA and VRE. The intrinsic resistance of gram-negative bacteria is mediated primarily by endogenous efflux pumps. Linezolid has excellent oral bioavailability. Adverse effects include myelosuppression and ocular and peripheral neuropathy with prolonged therapy. Peripheral neuropathy may be irreversible. Linezolid is a weak, reversible monoamine oxidase inhibitor, and coadministration with sympathomimetics and foods rich in tyramine should be avoided. Linezolid has been associated with serotonin syndrome when coadministered with selective serotonin-reuptake inhibitors. Tedizolid has properties similar to those of linezolid, but with lower dosing it may be less likely to cause adverse hematologic and neuropathic effects.

NITROFURANTOIN

Nitrofurantoin's antibacterial activity results from the drug's conversion to highly reactive intermediates that can damage DNA and other macromolecules. Nitrofurantoin is bactericidal, and its action is concentration dependent. It displays activity against a range of gram-positive bacteria, including *S. aureus*, *Staphylococcus epidermidis*, *Staphylococcus saprophyticus*, *E. faecalis*, *Streptococcus agalactiae*, group D streptococci, viridans streptococci, and corynebacteria, as well as gram-negative organisms, including *E. coli* and *Enterobacter*, *Neisseria*, *Salmonella*, and *Shigella* species. Nitrofurantoin is used primarily in the treatment of urinary tract infections and is preferred in the treatment of such infections in pregnancy. It may be used for the prevention of recurrent cystitis. Recently, there has been interest in the use of nitrofurantoin for treatment of urinary tract infections caused by ESBL-producing Enterobacteriaceae such as *E. coli*,

although resistance has been growing in Latin America and parts of Europe. Coadministration with magnesium should be avoided because of decreased absorption, and patients should be encouraged to take the drug with food to increase its bioavailability and decrease the risk of adverse effects, which include nausea, vomiting, and diarrhea. Nitrofurantoin may also cause pulmonary fibrosis and drug-induced hepatitis. Because the risk of adverse reactions increases with age, the use of nitrofurantoin in elderly patients is not recommended. Patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency are at elevated risk for nitrofurantoin-associated hemolytic anemia.

POLYMYXINS

Colistin and polymyxin B act by disrupting cell membrane integrity and are active against the nonenteric pathogens *P. aeruginosa* and *A. baumannii* but not against *Burkholderia*. These drugs also exhibit activity against many Enterobacteriaceae, with the exceptions of *Proteus*, *Providencia*, and *Serratia* species. They lack activity against gram-positive bacteria. Polymyxins are bactericidal and are available in IV formulations. Colistimethate is converted to the active form (colistin) in plasma. Polymyxins are most often used for infections due to pathogens resistant to multiple other antibacterial agents, including urinary tract infections, hospital-acquired pneumonia, and bloodstream infections. Nebulized formulations have been used for adjunctive treatment of refractory ventilator-associated pneumonia. The most important adverse effect is dose-dependent reversible nephrotoxicity. Neurotoxicity, including paresthesias, muscle weakness, and confusion, is reversible and less common than nephrotoxicity.

QUINUPRISTIN-DALFOPRISTIN

Quinupristin-dalfopristin is a member of the streptogramin class of antibiotics and kills bacteria by inhibiting protein synthesis. The antibacterial spectrum of quinupristin-dalfopristin includes staphylococci (including MRSA), streptococci, and *E. faecium* (but not *E. faecalis*). This drug is also active against *Corynebacterium* species and *L. monocytogenes*. Quinupristin-dalfopristin is not reliably active against gram-negative organisms. It exhibits concentration-dependent killing, with an AUC/MIC ratio predicting efficacy. The clinical use of quinupristin-dalfopristin is largely for infections due to vancomycin-resistant *E. faecium* and other gram-positive bacterial infections. The drug has demonstrated efficacy in a variety of infections, including urinary tract infections, bone and joint infections, and bacteremia. Adverse effects associated with quinupristin-dalfopristin include infusion-related reactions, arthralgias, and myalgias. The arthralgias and myalgias may be severe enough to warrant drug discontinuation. Quinupristin-dalfopristin inhibits the CYP3A4 drug-metabolizing enzyme, with consequent drug interactions (Table 170-4).

FOSFOMYCIN

Fosfomycin is a phosphonic acid antibiotic that has greater activity in acidic environments and is excreted in its active form in the urine. Thus, its use is primarily for prophylaxis and treatment of uncomplicated cystitis. The drug is administered as a single 3-g dose that results in high urine concentrations for up to 48 h. Fosfomycin is active against *S. aureus*, vancomycin-susceptible and vancomycin-resistant enterococci, and a wide range of gram-negative organisms, including *E. coli*, *Enterobacter* species, *S. marcescens*, *P. aeruginosa*, and *K. pneumoniae*. Notably, the vast majority of ESBL-producing Enterobacteriaceae are susceptible to fosfomycin. *A. baumannii* and *Burkholderia* species are resistant. The emergence of resistance to fosfomycin has not been observed during treatment of cystitis but has been documented during treatment of respiratory tract infections and osteomyelitis. The few adverse effects that have been reported include nausea and diarrhea.

CHLORAMPHENICOL

The use of chloramphenicol is limited by its potentially serious toxicities. When other agents are contraindicated or ineffective, chloramphenicol represents an alternative treatment for infections, including meningitis caused by susceptible bacteria such as *N. meningitidis*,

H. influenzae, and *S. pneumoniae*. It has also been used for the treatment of anthrax, brucellosis, *Burkholderia* infections, chlamydial infections, clostridial infections, ehrlichiosis, rickettsial infections, and typhoid fever. Adverse reactions include aplastic anemia, myelosuppression, and gray baby syndrome. Chloramphenicol inhibits the CYP2C19 and CYP3A4 drug-metabolizing enzymes and consequently increases levels of many classes of drugs.

APPROACH TO PROPHYLAXIS OF INFECTION

Antibacterial prophylaxis is indicated only in selected circumstances (Table 170-6) and should be supported by well-designed studies or expert panel recommendations. In all cases the risk or severity of the infection to be prevented should be greater than the adverse consequences of antibacterial therapy, including the potential for selection of resistance. In addition, the timing and duration of antibacterial treatment should be targeted for maximal effect and minimal required exposure. Prophylaxis of surgical site infections targets bacteria that may contaminate the wound during the surgical procedure, including the skin flora of the patient or operating team and the air in the operating room. Delivery of the antibacterial drug within 1 h before the surgical incision is most effective. For prolonged procedures, redosing may be necessary to maintain effective blood and tissue levels until the

wound is closed. In patients with nasal carriage of *S. aureus*, preoperative decolonization with nasal mupirocin reduces the rate of *S. aureus* surgical site infections and is generally recommended for high-risk procedures such as cardiac surgery and orthopedic implantation of prosthetic devices. For dental procedures, preprocedure antibacterial drugs are given to prevent transient bacteremia and the seeding of certain high-risk cardiac lesions. Prophylaxis is also used in nonprocedural settings in certain patients who have recurrent infections or who are at risk of serious infection from a specific exposure (e.g., close contact with a patient with meningococcal meningitis). Extension of prophylaxis beyond the period of infection risk (24 h in the case of surgical procedures) does not add further benefit and may increase the risk of resistance selection or *C. difficile* disease.

ANTIMICROBIAL STEWARDSHIP

In an era of increasing prevalence of multidrug-resistant bacteria and with a substantial amount of inappropriate antimicrobial use, the need for rational antimicrobial prescribing has never been greater. *Antimicrobial stewardship* describes the practice of promoting the selection of the appropriate drug, dosage, route, and duration of antimicrobial therapy. Antimicrobial stewardship programs implement a variety of strategies to (1) improve patient care through appropriate

TABLE 170-6 PROPHYLAXIS OF BACTERIAL INFECTIONS IN ADULTS

Condition	Antibacterial Agents ^a	Timing or Duration of Prophylaxis
Surgical		
Clean (cardiac, thoracic, neurologic, orthopedic, vascular, plastic)	Cefazolin (vancomycin, ^b clindamycin)	1 h before incision; redose with long procedures
Clean (ophthalmic)	Topical neomycin-polymyxin B-gramicidin, topical moxifloxacin	Every 5–15 min for 5 doses immediately prior to procedure
Clean-contaminated (head and neck)	Cefazolin + metronidazole, ampicillin-sulbactam ^f (clindamycin)	1 h before incision; redose with long procedures
Clean-contaminated (hysterectomy, gastroduodenal, biliary, unobstructed small intestine, urologic)	Cefazolin, ampicillin-sulbactam ^f (clindamycin + aminoglycoside, aztreonam, or fluoroquinolone)	1 h before incision; redose with long procedures
Clean-contaminated (colorectal, appendectomy)	Cefazolin + metronidazole, ampicillin-sulbactam, ^c ertapenem (clindamycin + aminoglycoside, aztreonam, or fluoroquinolone)	1 h before incision; redose with long procedures
Dirty (ruptured viscus)	Therapeutic regimen directed at anaerobes and gram-negative bacteria (e.g., ceftriaxone + metronidazole)	1 h before incision; redose with long procedures; continue for 3–5 days after procedure
Dirty (traumatic wound)	Therapeutic regimen: cefazolin (clindamycin ± aminoglycoside, aztreonam, or fluoroquinolone)	1 h before incision; redose with long procedures; continue for 3–5 days after procedure
Nonsurgical		
Dental, oral, or upper respiratory procedures in patients with high-risk cardiac lesions (prosthetic valves, congenital heart defects, prior endocarditis)	Amoxicillin PO, ampicillin IM (clindamycin PO, IV)	Oral agents 1 h before procedure; injection 30 min before procedure
Recurrent <i>S. aureus</i> skin infections ^d	Mupirocin ^e	Intranasal application for 5 days
Recurrent cellulitis associated with lymphatic disruption ^d	Benzathine penicillin IM monthly, oral penicillin or erythromycin twice daily	Undefined
Recurrent cystitis in women ^d	Nitrofurantoin, TMP-SMX, fluoroquinolone	After sexual intercourse or 3 times weekly for up to 1 year
Bite wounds	Amoxicillin-clavulanate (doxycycline, moxifloxacin)	3–5 days
Recurrent spontaneous bacterial peritonitis in cirrhotic patients ^d	Fluoroquinolone ^f	Undefined
Recurrent pneumococcal meningitis in patient with CSF leak or humoral immune defect ^d	Penicillin	Undefined
Exposure to patient with meningococcal meningitis	Rifampin, ciprofloxacin	2 days (rifampin), single dose (ciprofloxacin)
High-risk neutropenia (ANC, ≤100/μL for >7 days) ^d	Levofloxacin or ciprofloxacin ^f	Until neutropenia resolves or fever dictates use of other antibacterials

^aRegimens in parentheses are alternatives for patients allergic to β-lactams. ^bVancomycin may be given together with cefazolin to patients known to be colonized with methicillin-resistant *Staphylococcus aureus*. ^cCefoxitin or cefotetan may also be considered. ^dNot considered routine for all patients, but an acceptable consideration among alternative approaches. ^eUsually coupled with bathing with chlorhexidine-containing skin antiseptic. ^fChoice of fluoroquinolone prophylaxis must be balanced against the risk of selection of resistance.

Abbreviations: ANC, absolute neutrophil count; CSF, cerebrospinal fluid; TMP-SMX, trimethoprim-sulfamethoxazole.

946 antimicrobial use; (2) decrease the development of resistance within patients and populations; (3) reduce the incidence of adverse effects; and (4) control costs.

Infections caused by resistant pathogens result in significant morbidity and mortality as well as increased health care costs. Antimicrobial stewardship programs are typically multidisciplinary and often include infectious disease physicians, clinical pharmacists (usually with special training in infectious disease), clinical microbiologists, information systems specialists, infection prevention and control practitioners, and epidemiologists. These teams employ a variety of approaches to achieving the program's goals.

Established strategies of antimicrobial stewardship programs include (1) prospective audit of antimicrobial use, with intervention and feedback; (2) formulary restriction; and (3) preauthorization. *Prospective audit and feedback* are usually undertaken by an infectious disease physician or a pharmacist. In this process, orders for broad-spectrum antimicrobials (e.g., carbapenems) or high-impact agents (e.g., linezolid, daptomycin) are reviewed on a regular basis for appropriateness. In circumstances in which an antimicrobial is used in the absence of an appropriate indication, the stewardship program team intervenes and recommends an alternative to the primary team caring for the patient. This process has been successful in several

quasi-experimental studies, resulting in declines in use of broad-spectrum drugs and decreases in adverse events, such as *C. difficile* infection. *Formulary restriction* is the inclusion of a limited set of antimicrobial agents in a hospital formulary for the purpose of limiting indiscriminate use of antimicrobials in the absence of demonstrated benefit. Such restriction coincidentally serves to reduce costs. *Preauthorization* is the practice of requiring clinicians to obtain approval before using selected antimicrobials. Approval may be provided electronically with sophisticated Computerized Provider Order Entry (CPOE) software, after specific criteria for use are met, or after communication with an infectious disease specialist as designated by the stewardship program. These strategies have led to a decrease in *C. difficile* infections and to improvements in drug susceptibility patterns.

Additional strategies used in specific health-care settings are guidelines and pathways, dose optimization, parenteral-to-oral conversion, and de-escalation of therapy. Antimicrobial stewardship is an evolving area and an increasingly active area of research aimed at identifying the best practices. The IDSA, in collaboration with several other professional organizations, has published guidelines for developing institutional antimicrobial stewardship programs (www.idsociety.org/Antimicrobial_Agents/).

SECTION 5 DISEASES CAUSED BY GRAM-POSITIVE BACTERIA

PART 8

Infectious Diseases

171 Pneumococcal Infections

David Goldblatt, Katherine L. O'Brien

In the late nineteenth century, pairs of micrococci were first recognized in the blood of rabbits injected with human saliva by both Louis Pasteur, working in France, and George Sternberg, an American army physician. The important role of these micrococci in human disease was not appreciated at that time. By 1886, when the organism was designated “pneumokokkus” and *Diplococcus pneumoniae*, the pneumococcus had been isolated by many independent investigators, and its role in the etiology of pneumonia was well known. In the 1930s, pneumonia was the third leading cause of death in the United States (after heart disease and cancer) and was responsible for ~7% of all deaths both in the United States and in Europe. While pneumonia was caused by a host of pathogens, lobar pneumonia—a pattern more likely to be caused by the pneumococcus—accounted for approximately one-half of all pneumonia deaths in the United States in 1929. In 1974, the organism was reclassified as *Streptococcus pneumoniae*.

MICROBIOLOGY

Etiologic Agent Pneumococci are spherical gram-positive bacteria of the genus *Streptococcus*. Within this genus, cell division occurs along a single axis, and bacteria grow in chains or pairs—hence the name *Streptococcus*, from the Greek *streptos*, meaning “twisted,” and *kokkos*, meaning “berry.” At least 22 streptococcal species are recognized and are divided further into groups based on their hemolytic properties. *S. pneumoniae* belongs to the α -hemolytic group that characteristically produces a greenish color on blood agar because of the reduction of iron in hemoglobin (Fig. 171-1). The bacteria are fastidious and grow best in 5% CO₂ but require a source of catalase (e.g., blood) for growth on agar plates, where they develop mucoid (smooth/shiny) colonies. Pneumococci without a capsule produce colonies with a rough surface. Unlike that of other α -hemolytic streptococci, their growth is inhibited in the presence of optochin (ethylhydrocupreine hydrochloride), and they are bile soluble.

In common with other gram-positive bacteria, pneumococci have a cell membrane beneath a cell wall, which in turn is covered by a

polysaccharide capsule. Pneumococci are divided into serogroups or serotypes based on capsular polysaccharide structure, as distinguished with rabbit polyclonal antisera; capsules swell in the presence of specific antiserum (the Quellung reaction). The most recently discovered serotypes, 6C, 6D, and 11E, have been identified with monoclonal antibodies and by serologic, genetic, and biochemical means, respectively. The currently recognized 93 serotypes fall into 21 serogroups, and each serogroup contains two to five serotypes with closely related capsules. The capsule protects the bacteria from phagocytosis by host cells in the absence of type-specific antibody and is arguably the most



FIGURE 171-1 Pneumococci growing on blood agar, illustrating α hemolysis and optochin sensitivity (zone around optochin disk). Inset: Gram's stain, illustrating gram-positive diplococci. (Photographs courtesy of Paul Turner, Shoklo Malaria Research Unit, Thailand.)

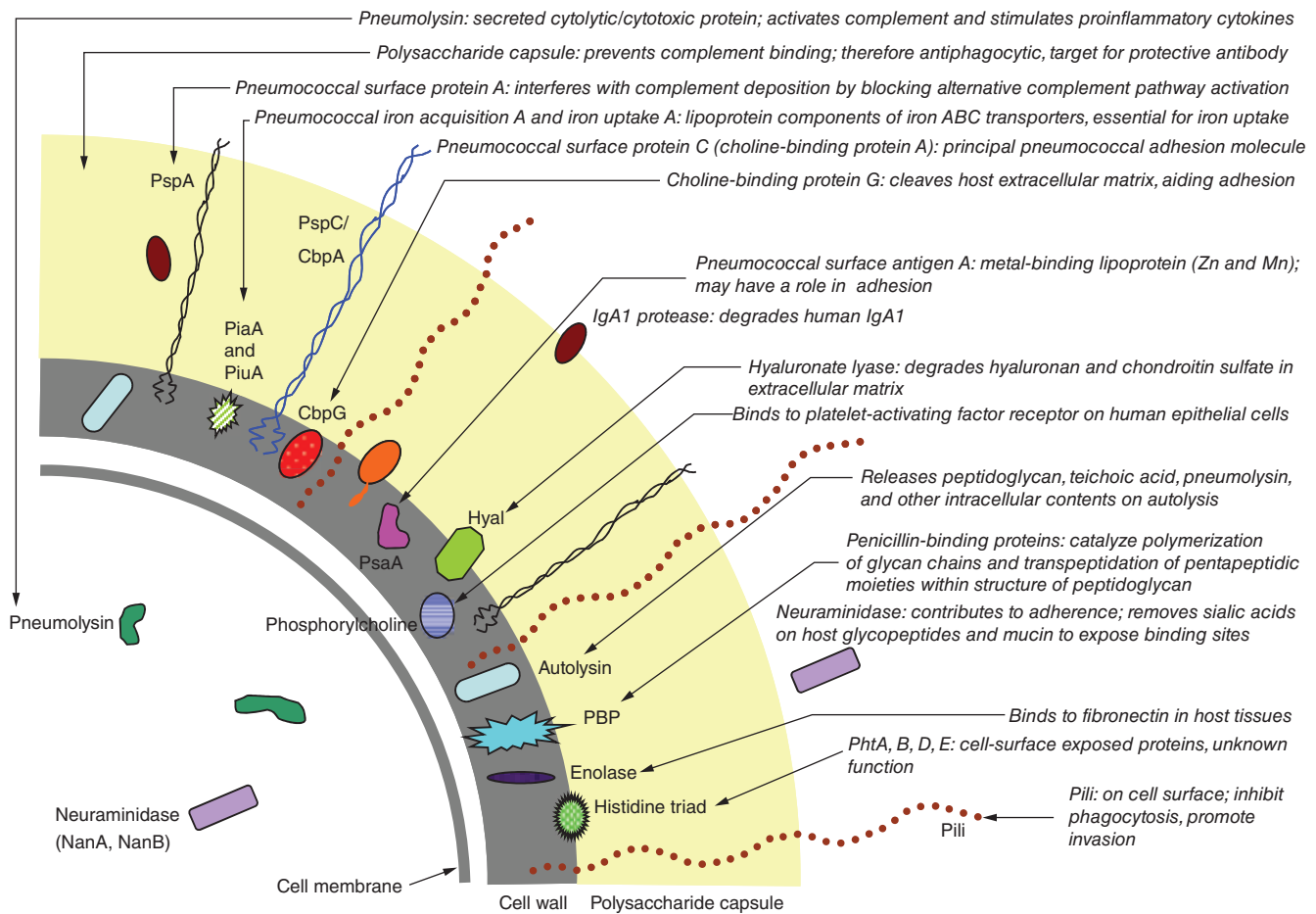


FIGURE 171-2 Schematic diagram of the pneumococcal cell surface, with key antigens and their roles highlighted.

important determinant of pneumococcal virulence. Unencapsulated variants tend not to cause invasive disease.

Virulence Factors Within the cytoplasm, cell membrane, and cell wall, many molecules that may play a role in pneumococcal pathogenesis and virulence have been identified (Fig. 171-2). These proteins are often involved in direct interactions with host tissues or in concealment of the bacterial surface from host defense mechanisms. Pneumolysin is a secreted cytotoxin thought to result in cytolysis of cells and tissues, and LytA enhances pathogenesis. A number of cell wall proteins interfere with the complement pathway, thus inhibiting complement deposition and preventing lysis and/or opsonophagocytosis. The pneumococcal H inhibitor (Hic) impedes the formation of C3 convertase, while pneumococcal surface protein C (PspC), also known as choline-binding protein A (CbpA), binds factor H and is thought to accelerate the breakdown of C3. PspA and CbpA inhibit the deposition of or degrade C3b. The numerous pneumococcal proteins thought to be involved in adhesion include the ubiquitous surface-anchored sialidase (neuraminidase) NanA, which cleaves sialic acid on host cells and proteins, and pneumococcal surface adhesin A (PsaA). Pili recently recognized by electron microscopy also may play an important role in binding to cells. Some of the antigens mentioned above are potential vaccine candidates (see “Prevention,” below).

Although the capsule surrounding the cell wall of *S. pneumoniae* is the basis for categorization by serotype, the behavior and pathogenic potential of a serotype may also be related to the genetic origin of the strain. Molecular typing is therefore of considerable interest. Initially, techniques such as pulsed-field gel electrophoresis were used to determine genetic relatedness; such techniques have been superseded by sequencing of housekeeping genes to define a clone (multilocus sequence typing, MLST). For *S. pneumoniae*, alleles at each of the loci *aroE*, *gdh*, *gki*, *recP*, *spi*, *xpt*, and *ddl* are sequenced and compared with all of the known alleles at that locus. Sequences identical to a known allele are assigned the same allele number,

whereas those differing from any known allele—even at a single nucleotide site—are assigned new numbers. Software for assignment of alleles at each locus is available on the pneumococcal MLST website (spneumoniae.mlst.net), and the allelic profile of each isolate and its consequent sequence type are generated. With the advent of high-throughput and relatively inexpensive sequencing techniques, whole-genome sequencing will soon supersede MLST.

EPIDEMIOLOGY

Pneumococcal infections remain a significant global cause of morbidity and death, particularly among children and the elderly. Rapid and dramatic changes in the epidemiology of this disease during the past decade in several developed countries followed the licensure and routine childhood administration of pneumococcal polysaccharide–protein conjugate vaccine (PCV). With PCV introduction in developing and middle-income countries, additional profound changes in pneumococcal ecology and disease epidemiology are likely. The disease burden and serotype distribution in the PCV era may be different than expected because of concomitant secular trends in pneumococcal disease, the impact of antibiotic use on pneumococcal strain ecology, and surveillance system attributes that can themselves affect analysis of epidemiologic features.

Serotype Distribution Not all pneumococcal serotypes are equally likely to cause disease; serotype distribution varies by age, disease syndrome, and geography. Geographic differences may be driven by variation in the burden of disease rather than by true serotype distribution differences. Most data on serotype distribution are related to pediatric invasive pneumococcal disease (IPD, defined as infection of a normally sterile site); much less information on global distribution is available for disease in adults. Among children <5 years of age, five to seven serotypes cause >60% of IPD cases in most parts of the world, seven serotypes (1, 5, 6A, 6B, 14, 19F, and 23F) account for ~60% of cases in all areas of the world, but in any given region these

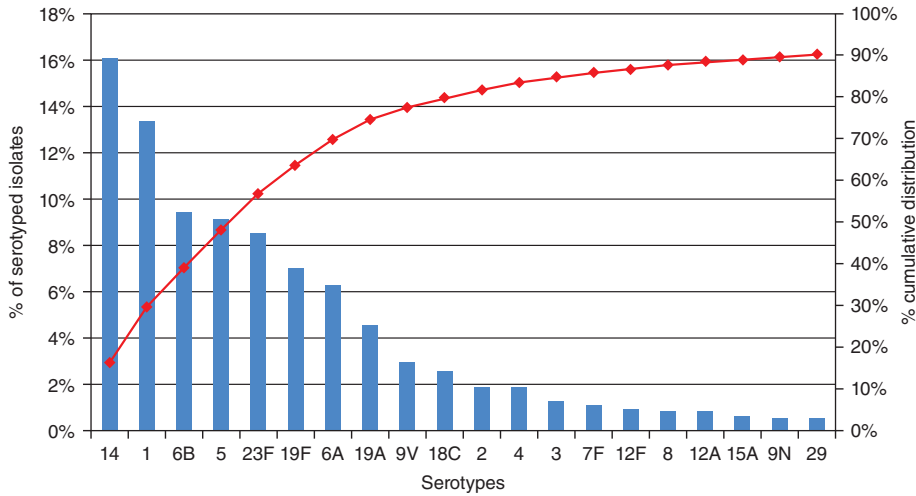


FIGURE 171-3 Meta-analysis of available global pneumococcal serotype data, adjusted for regional disease incidence. The red line shows cumulative incidence, as indicated on the right-hand Y axis. (Source: Global Serotype Project Report for the Pneumococcal Advance Market Commitment Target Product Profile; available at <http://www.gavi.org/library/gavi-documents/amc/tpp-codebook/>.)

seven serotypes may not all rank as the most common disease strains (Fig. 171-3). Some serotypes (e.g., types 1 and 5) not only tend to cause disease in areas with a high disease burden but also cause waves of disease in lower-burden areas (e.g., Europe) or outbreaks (e.g., in military barracks; meningitis in sub-Saharan Africa). The broader range of serotypes causing disease among adults than among children is apparent from a comparison of the coverage of existing multiserotype vaccines in different age groups. For example, data from the United States for 2006–2007 on the serotypes causing IPD indicated that a polysaccharide vaccine containing 23 serotypes (PPSV23) would cover 84% of cases among children <5 years of age and 76% of those among persons 18–64 years of age but only 65% of those among persons ≥65 years of age.

Nasopharyngeal Carriage Pneumococci are intermittent inhabitants of the healthy human nasopharynx and are transmitted by respiratory droplets. In children, pneumococcal nasopharyngeal ecology varies by geographic region, socioeconomic status, climate, degree of crowding, and particularly intensity of exposure to other children, with children in day-care settings having higher rates of colonization. In developed-world settings, children serve as the major vectors of pneumococcal transmission. By 1 year of age, ~50% of children have had at least one episode of pneumococcal colonization. Cross-sectional prevalence data show rates of pneumococcal carriage ranging from 20% to 50% among children <5 years of age and from 5% to 15% among young and middle-aged adults; Fig. 171-4 shows relevant data from the United Kingdom. Data on colonization rates among healthy elderly individuals are limited. In developing-world settings, pneumococcal acquisition occurs much earlier, sometimes within the first few days after birth, and nearly all infants have had at least one episode of colonization by 2 months of age. Cross-sectional studies show that up to the age of 5 years, 70–90% of children carry *S. pneumoniae* in the nasopharynx, and a significant proportion of adults (sometimes >40%) also are colonized. Their high rates of colonization make adults an important source of transmission and may affect community transmission dynamics.

Invasive Disease and Pneumonia IPD develops when *S. pneumoniae* invades the bloodstream and seeds other organs or directly reaches the cerebrospinal fluid (CSF) by local extension. Pneumonia may follow aspiration of pneumococci, although only 10–30% of such cases are associated with a positive blood culture (and thus contribute to the measured burden of IPD). The dramatic variation of IPD rates with age is illustrated by data from the United States for 1998–1999, a period prior to PCV introduction. Rates of IPD were highest among children

<2 years of age and among adults ≥65 years of age (188 and 60 cases/100,000, respectively; Fig. 171-5). Since the introduction of PCV, IPD rates among infants and children in the United States have fallen by >75%, a decrease driven by the near elimination of vaccine-serotype IPD. A similar impact of PCV on vaccine-serotype IPD rates has been consistently observed in countries where PCV has been introduced into the routine pediatric vaccination schedule. However, changes in the non-vaccine-serotype IPD rate in various countries have been heterogeneous; the interpretation of this heterogeneity is a complex issue. In the United States, Canada, and Australia, rates of non-vaccine-serotype IPD have increased but the magnitude of the increase is generally small relative to the substantial reductions in vaccine-serotype IPD. In contrast, in other settings (e.g., Alaska Native communities and the United Kingdom), the reduction in vaccine-serotype IPD has been offset by notable increases in rates of disease caused by non-vaccine serotypes. Explanations for the heterogeneity of findings include replacement

disease resulting from vaccine pressure, changes in clinical case investigation, secular trends unrelated to PCV use, antibiotic pressure selecting for resistant organisms, changes in surveillance or reporting systems, rapidity of introduction, and inclusion of a catch-up campaign. A recent systematic review concludes that serotype replacement in IPD follows the use of PCV7 but that the magnitude of this phenomenon is small relative to the reduction in disease from vaccine serotypes. The net effect of PCV is to reduce the rate of pneumococcal disease both in the age group targeted for vaccination and in unvaccinated age groups.

Pneumonia is the most common of the serious pneumococcal disease syndromes and poses special challenges from a clinical and public health perspective. Most cases of pneumococcal pneumonia are not associated with bacteremia, and in these cases a definitive etiologic diagnosis is difficult. As a result, estimates of disease burden focus primarily on IPD rates and fail to include the major portion of the burden of serious pneumococcal disease. Among children, PCV trials designed to collect efficacy data on syndrome-based outcomes (e.g., radiographically confirmed pneumonia, clinically diagnosed pneumonia) have revealed the burden of culture-negative pneumococcal pneumonia.

The case-fatality ratios (CFRs) for pneumococcal pneumonia and IPD vary by age, underlying medical condition, and access to care. In addition, the CFR for pneumococcal pneumonia varies with the

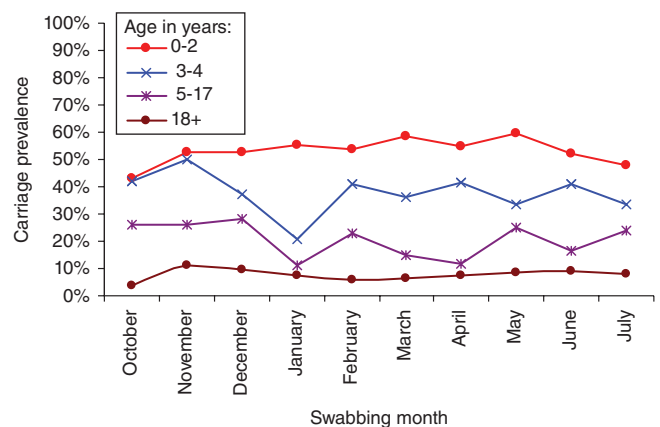


FIGURE 171-4 Prevalence of pneumococcal carriage in adults and children resident in the United Kingdom who had nasopharyngeal swabs collected monthly for 10 months (no seasonal trend; *t* test trend, >.05). (Data adapted from D Goldblatt et al: *J Infect Dis* 192:387, 2005.)

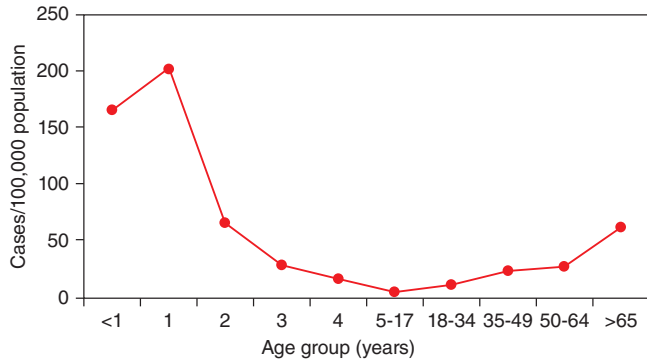


FIGURE 171-5 Rates of invasive pneumococcal disease before the introduction of pneumococcal conjugate vaccine, by age group: United States, 1998. (Source: CDC, *Active Bacterial Core Surveillance/Emerging Infectious Program Network*, 2000. Data adapted from *MMWR* 49[RR-9], 2000.)

severity of disease at presentation (rather than according to whether the pneumonia episode is associated with bacteremia) and with the patient's age (from <5% among hospitalized patients 18–44 years old to >12% among those >65 years old, even when appropriate and timely management is available). Notably, the likelihood of death in the first 24 h of hospitalization did not change substantially with the introduction of antibiotics; this surprising observation highlights the fact that the pathophysiology of severe pneumococcal pneumonia among adults reflects a rapidly progressive cascade of events that often unfolds irrespective of antibiotic administration. Management in an intensive care unit can provide critical support for the patient through the acute period, with lower CFRs.

Rates of pneumococcal disease vary by season, with higher rates in colder than in warmer months in temperate climates; by sex, with males more often affected than females; and by risk group, with risk factors including underlying medical conditions, behavioral issues, and ethnic group. In the United States, some Native American populations (including Alaska natives) and African Americans have higher rates of disease than the general population; the increased risk is probably attributable to socioeconomic conditions and the prevalence of underlying risk factors for pneumococcal disease. Medical conditions that increase the risk of pneumococcal infection are listed in [Table 171-1](#). Outbreaks of disease are well recognized in crowded settings with susceptible individuals, such as infant day-care facilities, military barracks, and nursing homes. Furthermore, there is a clear association between preceding viral respiratory disease (especially but not exclusively influenza) and risk of secondary pneumococcal infections. The significant role of pneumococcal pneumonia in the morbidity and mortality associated with seasonal and pandemic influenza is increasingly recognized.

Antibiotic Resistance Reduced pneumococcal susceptibility to penicillin was first noted in 1967, but not until the 1990s did reduced antibiotic susceptibility emerge as a significant clinical and public health issue, with an increasing prevalence of pneumococcal isolates resistant to single or multiple classes of antibiotics and a rising absolute magnitude of minimal inhibitory concentrations (MICs). Strains with reduced susceptibility to penicillin G, cefotaxime, ceftriaxone, macrolides, and other antibiotics are now found worldwide and account for a significant proportion of disease-causing strains in many locations, especially among children. Vancomycin resistance has not yet been observed in clinical pneumococcal strains. Lack of antimicrobial susceptibility is clearly related to a subset of serotypes, many of which disproportionately cause disease among children. The vicious cycle of antibiotic exposure, selection of resistant organisms in the nasopharynx, and transmission of these organisms within the community, leading to difficult-to-treat infections and increased antibiotic exposure, has been interrupted to some extent by the introduction and routine use of PCV. The clinical implications of pneumococcal antimicrobial nonsusceptibility are addressed below in the section on treatment.

TABLE 171-1 CLINICAL RISK GROUPS FOR PNEUMOCOCCAL INFECTION

Clinical Risk Group	Examples
Asplenia or splenic dysfunction	Sickle cell disease, celiac disease
Chronic respiratory disease	Chronic obstructive pulmonary disease, bronchiectasis, cystic fibrosis, interstitial lung fibrosis, pneumoconiosis, bronchopulmonary dysplasia, aspiration risk, neuromuscular disease (e.g., cerebral palsy), severe asthma
Chronic heart disease	Ischemic heart disease, congenital heart disease, hypertension with cardiac complications, chronic heart failure
Chronic kidney disease	Nephrotic syndrome, chronic renal failure, renal transplantation
Chronic liver disease	Cirrhosis, biliary atresia, chronic hepatitis
Diabetes mellitus	Diabetes mellitus requiring insulin or oral hypoglycemic drugs
Immunocompromise/immunosuppression	HIV infection, common variable immunodeficiency, leukemia, lymphoma, Hodgkin's disease, multiple myeloma, generalized malignancy, chemotherapy, organ or bone marrow transplantation, systemic glucocorticoid treatment for >1 month at a dose equivalent to ≥ 20 mg/d (children, ≥ 1 mg/kg per day)
Cochlear implants	...
Cerebrospinal fluid leaks	...
Miscellaneous	Infancy and old age; prior hospitalization; alcoholism; malnutrition; cigarette smoking; day-care center attendance; residence in military training camps, prisons, homeless shelters

Note: Groups for whom pneumococcal vaccines are recommended by the Advisory Committee on Immunization Practices can be found at www.cdc.gov/vaccines/schedules/.

PATHOGENESIS

Pneumococci colonize the human nasopharynx from an early age; colonization acquisition events are generally described as asymptomatic, but evidence exists to associate acquisition with mild respiratory symptoms, especially in the very young. From the nasopharynx, the bacteria spread either via the bloodstream to distant sites (e.g., brain, joint, bones, peritoneal cavity) or locally to mucosal surfaces where they can cause otitis media or pneumonia. Direct spread from the nasopharynx to the central nervous system (CNS) can occur in rare cases of skull base fracture, although most cases of pneumococcal meningitis are secondary to hematogenous spread. Pneumococci can cause disease in almost any organ or part of the body; however, otitis media, pneumonia, bacteremia, and meningitis are most common. Colonization is a relatively frequent event, yet disease is rare. In the nasopharynx, pneumococci survive in mucus secreted by epithelial cells, where they can avoid local immune factors such as leukocytes and complement. The mucus itself is a component of local defense mechanisms, and the flow of mucus (driven in part by cilia in what is known as the *mucociliary escalator*) effects mechanical clearance of pneumococci. While many colonization episodes are of short duration, longitudinal studies in adults and children have revealed persistent colonization with a specific serotype over many months. Colonization eventually results in the development of capsule-specific serum IgG, which is thought to play a role in mediating clearance of bacteria from the nasopharynx. IgG antibodies to surface-exposed cell wall or secreted proteins also appear in the circulation in an age-dependent fashion or after colonization; the biologic role of these antibodies is less clear. Recent acquisition of a new colonizing serotype is more likely to be associated with subsequent invasion, presumably as a result of the absence of type-specific immunity. Intercurrent viral infections make the host more susceptible to pneumococcal colonization, and pneumococcal disease in a colonized individual often follows perturbation of the nasopharyngeal mucosa by such infections. Local cytokine production after a viral infection is thought to upregulate adhesion factors in the respiratory epithelium, allowing pneumococci to adhere via a variety of surface adhesin molecules, including PsaA, PspA,

950 CbpA, PspC, Hyl, pneumolysin, and the neuraminidases (Fig. 171-2). Adhesion coupled with inflammation induced by pneumococcal factors such as peptidoglycans and teichoic acids results in invasion. It is the inflammation induced by various bacterium-derived factors that is responsible for the pathology associated with pneumococcal infection. Cell wall-derived teichoic acids and peptidoglycans induce a variety of cytokines, including the proinflammatory cytokines interleukin (IL) 1, IL-6, and tumor necrosis factor, and activate complement via the alternative pathway. Polymorphonuclear leukocytes are thus attracted, and an intense inflammatory response is initiated. Pneumolysin also is important in local pathology, inducing proinflammatory cytokine production by local monocytes.

The pneumococcal capsule, consisting of polysaccharides with antiphagocytic properties due to resistance to the deposition of complement, plays an important role in pathogenesis. While most capsular types can cause human disease, certain capsular types are more commonly isolated from sites of infection. The reason for the dominance of some serotypes over others in IPD, as depicted in Fig. 171-3, is unclear.

HOST DEFENSE MECHANISMS

Innate Immunity As described above, intact respiratory epithelium and a host of nonspecific or innate immune factors (e.g., mucus, splenic function, complement, neutrophils, and macrophages) constitute the first line of defense against pneumococci. Physical factors such as the cough reflex and the mucociliary escalator are important in clearing bacteria from the lungs. Immunologic factors are critical as well: C-reactive protein (CRP) binds phosphorylcholine in the pneumococcal cell wall, inducing complement activation and leading to bacterial clearance; Toll-like receptor 2 (TLR2) recognizes both pneumococcal lipoteichoic acid and cell wall peptidoglycan; and in animal models, the absence of host TLR2 leads to more severe infection and impaired clearance of nasopharyngeal colonization. TLR4 appears to be necessary for the proinflammatory effect of pneumolysin on macrophages. The importance of TLR recognition is underlined by descriptions of an inherited deficiency of human IL-1 receptor-associated kinase 4 (IRAK-4) that manifests as an unusual susceptibility to infection with bacteria, including *S. pneumoniae*. IRAK-4 is essential for the normal functioning of several TLRs. Other factors that interfere with these nonspecific mechanisms (e.g., viral infections, cystic fibrosis, bronchiectasis, complement deficiency, and chronic obstructive pulmonary disease) all predispose to the development of pneumococcal pneumonia. Patients who lack a spleen or have abnormal splenic function (e.g., persons with sickle cell disease) are at high risk of developing overwhelming pneumococcal disease.

Acquired Immunity Acquired immunity induced via contact following colonization or through cross-reactive antigens rests largely on the development of serum IgG antibody specific for the pneumococcal capsular polysaccharide. Nearly all polysaccharides are T cell-independent antigens; B cells can make antibodies to such antigens without T cell help. However, in children <1–2 years old, such B cell responses are poorly developed. This delayed ontogeny of capsule-specific IgG in young children is associated with susceptibility to pneumococcal infection (Fig. 171-5). The extremely high risk of pneumococcal infection in the absence of serum immunoglobulin (i.e., in conditions such as agammaglobulinemia) highlights the important role of capsular antibody in protection against disease. Each serotype's capsule is chemically distinct; thus immunity tends to be serotype specific, although some cross-immunity exists. For example, conjugate vaccine-induced antibodies to serotype 6B prevent infection due to serotype 6A. However, cross-protection against serotypes within serogroups is not universal; for instance, antibodies to serotype 19F do not appear to confer protection against disease caused by serotype 19A. Antibodies to surface-exposed or secreted pneumococcal proteins (such as pneumolysin, PsaA, and PspA) also appear in the circulation with increasing age of the host, but their functional significance remains unclear. Data from murine models suggest that CD4+ T cells may play a role in preventing pneumococcal colonization and disease, and recent experimental data

derived from humans suggest that IL-17-secreting CD4+ T cells may be relevant.

APPROACH TO THE PATIENT: Pneumococcal Infections

There is no pathognomonic presentation of pneumococcal disease; patients may present with a range of syndromes and with more than one clinical syndrome (e.g., pneumonia and meningitis). *S. pneumoniae* can infect nearly any body tissue, manifesting as disease ranging in severity from mild and self-limited to life-threatening. The differential diagnosis of common clinical syndromes such as pneumonia, otitis media, fever of unknown origin, and meningitis should always include pneumococcal infection. A microbiologically confirmed diagnosis is made in only a minority of pneumococcal cases since, in most circumstances (and especially in pneumonia and otitis media), fluid from the site of infection is not available for etiologic determination. Empirical therapy that includes appropriate treatment for *S. pneumoniae* is often indicated.

Algorithms for assessment and management of ill children have been developed for use in the developing world or in other settings where evaluation by a trained physician may not be feasible. Children who present with ominous signs such as an inability to drink, convulsions, lethargy, and severe malnutrition are categorized as having very severe disease without further evaluation by the community health care worker, are given antibiotics, and are immediately referred to a hospital for diagnosis and management. Children who present with cough and tachypnea (the latter defined according to specific age strata) are further stratified into severity categories based on the presence or absence of lower chest wall indrawing and are managed accordingly with either antibiotics alone or antibiotics and referral to a hospital facility. Children with cough but no tachypnea are categorized as having a nonpneumonia respiratory illness.

CLINICAL MANIFESTATIONS

The clinical manifestations of pneumococcal disease depend on the site of infection and the duration of illness. Clinical syndromes are classified as noninvasive (e.g., otitis media and nonbacteremic pneumonia) or invasive (e.g., bacteremic pneumonia). The pathogenesis of noninvasive illness involves contiguous spread from the nasopharynx or skin; invasive disease involves infection of a normally sterile body fluid or follows bacteremia.

Pneumonia Pneumonia is the most common serious pneumococcal syndrome and is considered invasive when associated with a positive blood culture. Pneumococcal pneumonia can present as a mild community-acquired infection at one extreme and as a life-threatening disease requiring intubation and intensive support at the other.

PRESENTING MANIFESTATIONS The presentation of pneumococcal pneumonia does not reliably distinguish it from pneumonia of other etiologies. In a subset of cases, pneumococcal pneumonia is recognized at the outset as associated with a viral upper respiratory infection and is characterized by the abrupt onset of cough and dyspnea accompanied by fever, shaking chills, and myalgias. The cough evolves from nonpurulent to productive of sputum that is purulent and sometimes tinged with blood. Patients may describe stabbing pleuritic chest pain and significant dyspnea indicating involvement of the parietal pleura. Among the elderly, the presenting clinical symptoms may be less specific, with confusion or malaise but without fever or cough. In such cases, a high index of suspicion is required because failure to treat pneumococcal pneumonia promptly in an elderly patient is likely to result in rapid evolution of the infection, with increased severity, morbidity, and risk of death.

FINDINGS ON PHYSICAL EXAMINATION The clinical signs associated with pneumococcal pneumonia among adults include tachypnea

(>30 breaths/min) and tachycardia, hypotension in severe cases, and fever in most cases (although not in all elderly patients). Respiratory signs are varied, including dullness to percussion in areas of the chest with significant consolidation, crackles on auscultation, reduced expansion of the chest in some cases as a result of splinting to reduce pain, bronchial breathing in a minority of cases, pleural rub in occasional cases, and cyanosis in cases with significant hypoxemia. Among infants with severe pneumonia, chest wall indrawing and nasal flaring are common. Nonrespiratory findings can include upper abdominal pain if the diaphragmatic pleura is involved as well as mental status changes, particularly confusion in elderly patients.

DIFFERENTIAL DIAGNOSIS The differential diagnosis of pneumococcal pneumonia includes cardiac conditions such as myocardial infarction and heart failure with atypical pulmonary edema; pulmonary conditions such as atelectasis; and pneumonia caused by viral pathogens, mycoplasmas, *Haemophilus influenzae*, *Klebsiella pneumoniae*, *Staphylococcus aureus*, *Legionella*, or (in HIV-infected and otherwise immunocompromised hosts) *Pneumocystis*. In cases with abdominal symptoms, the differential diagnosis includes cholecystitis, appendicitis, perforated peptic ulcer disease, and subphrenic abscesses. The challenge in cases with abdominal symptoms is to remember to include pneumococcal pneumonia—a nonabdominal process—in the differential diagnosis.

DIAGNOSIS Some authorities advocate treating uncomplicated, non-severe, community-acquired pneumonia without determining the microbiologic etiology, given that this information is unlikely to alter clinical management. However, efforts to identify the cause of pneumonia are important when the disease is more severe and when the diagnosis of pneumonia is not clearly established. The gold standard for etiologic diagnosis of pneumococcal pneumonia is pathologic examination of lung tissue. In lieu of that procedure, evidence of an infiltrate on chest radiography warrants a diagnosis of pneumonia. However, cases of pneumonia without radiographic evidence do occur. An infiltrate can be absent either early in the course of the illness or with dehydration; upon rehydration, an infiltrate usually appears. The radiographic appearance of pneumococcal pneumonia is varied; it classically consists of lobar or segmental consolidation (Fig. 171-6) but in some cases is patchy. More than one lobe is involved in ~30% of cases. Consolidation may be associated with a small pleural effusion



FIGURE 171-6 Chest radiograph depicting classic lobar pneumococcal pneumonia in the right lower lobe of an elderly patient's lung.

or empyema in complicated cases. In children, “round pneumonia,” a distinctly spherical consolidation on chest radiography, is associated with a pneumococcal etiology. Round pneumonia is uncommon in adults. *S. pneumoniae* is not the only cause of such lesions; other causes, especially cancer, should be considered.

Blood drawn from patients with suspected pneumococcal pneumonia can be used for supportive or definitive diagnostic tests. Blood cultures are positive for pneumococci in a minority (<30%) of cases of pneumococcal pneumonia. Nonspecific findings include an elevated polymorphonuclear leukocyte count (>15,000/ μ L in most cases and upward of 40,000/ μ L in some), leukopenia in <10% of cases (a poor prognostic sign associated with a fatal outcome), and elevated values in liver function tests (e.g., both conjugated and unconjugated hyperbilirubinemia). Anemia, low serum albumin levels, hyponatremia, and elevated serum creatinine levels are all found in ~20–30% of patients.

Urinary pneumococcal antigen assays have facilitated etiologic diagnosis. In adults, among whom the prevalence of pneumococcal nasopharyngeal colonization is relatively low, a positive pneumococcal urinary antigen test has a high predictive value. The same is not true for children, in whom a positive urinary antigen test can reflect the mere presence of *S. pneumoniae* in the nasopharynx.

Most cases of pneumococcal pneumonia are diagnosed by Gram's staining and culture of sputum. The utility of a sputum specimen is directly related to its quality and the patient's antibiotic treatment status.

COMPLICATIONS Empyema is the most common focal complication of pneumococcal pneumonia, occurring in <5% of cases. When fluid in the pleural space is accompanied by fever and leukocytosis (even low-grade) after 4–5 days of appropriate antibiotic treatment for pneumococcal pneumonia, empyema should be considered. Parapneumonic effusions are more common than empyema, representing a self-limited inflammatory response to pneumonia. Pleural fluid with frank pus, bacteria (detected by microscopic examination), or a pH of ≤ 7.1 indicates empyema and demands aggressive and complete drainage, usually through chest tube insertion.

Meningitis Pneumococcal meningitis typically presents as a pyogenic condition that is clinically indistinguishable from meningitis of other bacterial etiologies. Meningitis can be the primary presenting pneumococcal syndrome or a complication of other conditions such as skull fracture, otitis media, bacteremia, or mastoiditis. Now that *H. influenzae* type b vaccine is routinely used, *S. pneumoniae* and *Neisseria meningitidis* are the most common bacterial causes of meningitis in both adults and children. Pyogenic meningitis, including that due to *S. pneumoniae*, is associated clinically with findings that include severe, generalized, gradual-onset headache, fever, and nausea as well as specific CNS manifestations such as stiff neck, photophobia, seizures, and confusion. Clinical signs include a toxic appearance, altered consciousness, bradycardia, and hypertension indicative of increased intracranial pressure. A small proportion of adult patients have Kernig's or Brudzinski's sign or cranial nerve palsies (particularly of the third and sixth cranial nerves).

A definitive diagnosis of pneumococcal meningitis rests on the examination of CSF for (1) evidence of turbidity (visual inspection); (2) elevated protein level, elevated white blood cell count, and reduced glucose concentration (quantitative measurement); and (3) specific identification of the etiologic agent (culture, Gram's staining, antigen testing, or polymerase chain reaction [PCR]). A blood culture positive for *S. pneumoniae* in conjunction with clinical manifestations of meningitis also is considered confirmatory. Among adults, detection of pneumococcal antigen in urine is considered highly specific because of the low prevalence of nasopharyngeal colonization in this age group.

The mortality rate for pneumococcal meningitis is ~20%. In addition, up to 50% of survivors experience acute or chronic complications, including deafness, hydrocephalus, and mental retardation in children and diffuse brain swelling, subarachnoid bleeding, hydrocephalus, cerebrovascular complications, and hearing loss in adults.

Other Invasive Syndromes *S. pneumoniae* can cause other invasive syndromes involving virtually any body site. These syndromes include

952 primary bacteremia without other sites of infection (bacteremia without a source; occult bacteremia), osteomyelitis, septic arthritis, endocarditis, pericarditis, and peritonitis. The essential diagnostic approach is collection of fluid from the site of infection by sterile technique and examination by Gram's staining, culture, and—when relevant—capsular antigen assay or PCR. Hemolytic-uremic syndrome can complicate invasive pneumococcal disease.

Noninvasive Syndromes The two major noninvasive syndromes caused by *S. pneumoniae* are sinusitis and otitis media; the latter is the most common pneumococcal syndrome and most often affects young children. The manifestations of otitis media include the acute onset of severe pain, fever, deafness, and tinnitus, most frequently in the setting of a recent upper respiratory tract infection. Clinical signs include a red, swollen, often bulging tympanic membrane with reduced movement on insufflation or tympanography. Redness of the tympanic membrane is not sufficient for the diagnosis of otitis media.

Pneumococcal sinusitis is also a complication of upper respiratory tract infections and presents with facial pain, congestion, fever, and—in many cases—persistent nighttime cough. A definitive diagnosis is made by aspiration and culture of sinus material; however, presumptive treatment is most commonly initiated after application of a strict set of clinical diagnostic criteria.

TREATMENT PNEUMOCOCCAL INFECTIONS

Historically, the activity of penicillin against pneumococci made parenteral penicillin G the drug of choice for disease caused by susceptible organisms, including community-acquired pneumonia. For susceptible strains, penicillin G remains the most commonly used agent, with daily doses ranging from 50,000 U/kg for minor infections to 300,000 U/kg for meningitis. Other parenteral β -lactam drugs, such as ampicillin, cefotaxime, ceftriaxone, and cefuroxime, can be used against penicillin-susceptible strains but offer little advantage over penicillin. Macrolides and cephalosporins are alternatives for penicillin-allergic patients. While agents such as clindamycin, tetracycline, and trimethoprim-sulfamethoxazole exhibit some activity against pneumococci, resistance to these agents is frequently encountered in different parts of the world.

Penicillin-resistant pneumococci were first described in the mid-1960s, at which point tetracycline- and macrolide-resistant strains had already been reported. Multidrug-resistant strains were first described in the 1970s, but it was during the 1990s that pneumococcal drug resistance reached pandemic proportions. The use of antibiotics selects for resistant pneumococci, and strains resistant to β -lactam agents and to multiple drugs are now found all over the world. The emergence of high rates of macrolide and fluoroquinolone resistance also has been described.

The molecular basis of penicillin resistance in *S. pneumoniae* is the alteration of penicillin-binding protein (PBP) genes by transformation and horizontal transfer of DNA from related streptococcal species. Such alteration of PBPs results in lower affinity for penicillins. Depending on the specific PBP(s) and the number of PBPs altered, the level of resistance ranges from intermediate to high. For many years, penicillin susceptibility breakpoints have been defined by MICs as follows: susceptible, ≤ 0.06 $\mu\text{g/mL}$; intermediate, 0.12–1.0 $\mu\text{g/mL}$; and resistant, ≥ 2.0 $\mu\text{g/mL}$. However, in vitro results often were not predictive of the response of a patient to treatment for pneumococcal diseases other than meningitis. New recommendations have been based on the revised penicillin G breakpoints established in 2008 by the Clinical and Laboratory Standards Institute. For IV treatment of meningitis with at least 24 million units per day in 8 divided doses, the susceptibility breakpoint remains ≤ 0.06 $\mu\text{g/mL}$, and MICs of ≥ 0.12 $\mu\text{g/mL}$ indicate resistance. For IV treatment of nonmeningeal infections with 12 million units per day in 6 divided doses, the breakpoints are ≤ 2 $\mu\text{g/mL}$ for susceptible organisms, 4 $\mu\text{g/mL}$ for intermediate organisms, and ≥ 8 $\mu\text{g/mL}$ for resistant organisms; a dosage of 18–24 million units per day is recommended for strains with MICs in the intermediate category. The original

breakpoints remain the same for oral treatment of nonmeningeal infections with penicillin V.

Although guidelines for antibiotic therapy should be driven in part by local patterns of resistance, guidelines from national organizations in many countries (e.g., the Infectious Diseases Society of America/American Thoracic Society, the British Thoracic Society, and the European Respiratory Society) lay out evidence-based approaches. The following guidelines for the treatment of individual sepsis syndromes are based on those advocated by the American Academy of Pediatrics and published in the 2012 *Red Book*.

MENINGITIS LIKELY OR PROVEN TO BE DUE TO *S. PNEUMONIAE*

As a result of the increased prevalence of resistant pneumococci, first-line therapy for persons ≥ 1 month of age is a combination of vancomycin (adults, 30–60 mg/kg per day; infants and children, 60 mg/kg per day) and cefotaxime (adults, 8–12 g/d in 4–6 divided doses; children, 225–300 mg/kg per day in 1 dose or 2 divided doses) or ceftriaxone (adults, 4 g/d in 1 dose or 2 divided doses; children, 100 mg/kg per day in 1 dose or 2 divided doses). If children are hypersensitive to β -lactam agents (penicillins and cephalosporins), rifampin (adults, 600 mg/d; children, 20 mg/d in 1 dose or 2 divided doses) can be substituted for cefotaxime or ceftriaxone. A repeat lumbar puncture should be considered after 48 h if the organism is not susceptible to penicillin and information on cephalosporin sensitivity is not yet available, if the patient's clinical condition does not improve or deteriorates, or if dexamethasone has been administered and may be compromising clinical evaluation. When antibiotic sensitivity data become available, treatment should be modified accordingly. If the isolate is sensitive to penicillin, vancomycin can be discontinued and penicillin can replace the cephalosporin, or cefotaxime or ceftriaxone can be continued alone. If the isolate displays any resistance to penicillin but is susceptible to the cephalosporins, vancomycin can be discontinued and cefotaxime or ceftriaxone continued. If the isolate exhibits any resistance to penicillin and is not susceptible to cefotaxime and ceftriaxone, vancomycin and high-dose cefotaxime or ceftriaxone can be continued; rifampin may be added as well if the isolate is susceptible and the patient's clinical condition is worsening, if the CSF remains positive for bacteria, or if the MIC of the cephalosporin in question against the infecting strain is high. Some physicians advocate the use of glucocorticoids in children >6 months old, but this recommendation remains controversial and is not universally considered the standard of care. Glucocorticoids significantly reduce rates of mortality, severe hearing loss, and neurologic sequelae in adults and should be administered to those with community-acquired bacterial meningitis. If dexamethasone is given to either adults or children, it should be administered before or in conjunction with the first antibiotic dose.

INVASIVE INFECTIONS (EXCLUDING MENINGITIS)

In previously well children with noncritical illness, therapy with a recommended antibiotic should be instigated at the following dosages: penicillin G, 250,000–400,000 units/kg per day (in divided doses 4–6 h apart); cefotaxime, 75–100 mg/d (doses 8 h apart); or ceftriaxone, 50–75 mg/d (doses 12–24 h apart). For critically ill children, including those who have myocarditis or multilobular pneumonia with hypoxia or hypotension, vancomycin may be added if the isolate may possibly be resistant to β -lactam drugs, with its use reviewed once susceptibility data become available. If the organism is resistant to β -lactam agents, therapy should be modified on the basis of clinical response and susceptibility to other antibiotics. Clindamycin or vancomycin can be used as a first-line agent for children with severe β -lactam hypersensitivity, but vancomycin should not be continued if the organism is shown to be sensitive to other non- β -lactam antibiotics.

For outpatient management, amoxicillin (1 g every 8 h) provides effective treatment for virtually all cases of pneumococcal pneumonia. Neither cephalosporins nor quinolones, which are far more expensive, offer any advantage over amoxicillin. Levofloxacin

(500–750 mg/d as a single dose) and moxifloxacin (400 mg/d as a single dose) also are highly likely to be effective in the United States except in patients who come from closed populations where these drugs are used widely or who have themselves been treated recently with a quinolone. Clindamycin (600–1200 mg/d every 6 h) is effective in 90% of cases and azithromycin (500 mg on day 1 followed by 250–500 mg/d) or clarithromycin (500–750 mg/d as a single dose) in 80% of cases. Treatment failure resulting in bacteremic disease due to macrolide-resistant isolates has been amply documented in patients given azithromycin empirically. As noted above, rates of resistance to all these antibiotics are relatively low in some countries and much higher in others; high-dose amoxicillin remains the best option worldwide.

The optimal duration of treatment for pneumococcal pneumonia is uncertain, but its continuation for at least 5 days once the patient becomes afebrile appears to be a prudent approach. Cases with a second focus of infection (e.g., empyema or septic arthritis) require longer therapy.

ACUTE OTITIS MEDIA

Amoxicillin (80–90 mg/kg per day) is recommended for children with acute otitis media except in situations where observation and symptom-based treatment without antibiotics are advocated. These situations include nonsevere illness and an uncertain diagnosis in children 6 months to 2 years of age and nonsevere illness (even if the diagnosis seems certain) in children >2 years of age. Although the optimal duration of therapy has not been conclusively established, a 10-day course is recommended for younger children and for children with severe disease at any age. For children >6 years old who have mild or moderate disease, a course of 5–7 days is considered adequate. Patients whose illness fails to respond should be reassessed at 48–72 h. If acute otitis media is confirmed and antibiotic treatment has not been started, administration of amoxicillin should be commenced. If antibiotic therapy fails, a change is indicated. Failure to respond to second-line antibiotics as well indicates that myringotomy or tympanocentesis may need to be undertaken in order to obtain samples for culture.

The above recommendations can also be followed for the treatment of sinusitis. Detailed information on the further management of these conditions in children has been published by the American Academy of Pediatrics and the American Academy of Family Physicians.

PREVENTION

Measures to prevent pneumococcal disease include vaccination against *S. pneumoniae* and influenza viruses, reduction of comorbidities that increase the risk of pneumococcal disease, and prevention of antibiotic overuse, which fuels pneumococcal resistance.

Capsular Polysaccharide Vaccines The 23-valent pneumococcal polysaccharide vaccine (PPSV23), containing 25 µg of each capsular polysaccharide, has been licensed for use since 1983. Recommendations for its use vary by country. The U.S. Advisory Committee on Immunization Practices recommends PPSV23 for all persons ≥65 years of age and for those 2–64 years of age who have underlying medical conditions that put them at increased risk for pneumococcal disease or, if infected, disease of increased severity (Table 171-1; see also www.cdc.gov/vaccines/schedules/). The committee recently updated their recommendations to include the combined use of PPSV23 and a conjugate vaccine in at-risk individuals (see “Polysaccharide–Protein Conjugate Vaccines,” below). Revaccination 5 years after the first dose is recommended for persons >2 years of age who have underlying medical conditions but not routinely for those whose only indication is an age of ≥65 years. PPSV23 does not induce an anamnestic response, and antibody concentrations wane over time; thus revaccination is particularly important for individuals with conditions resulting in loss of antibody. Concerns about repeated revaccination have focused on safety (i.e., local reactions) and the induction of immune hyporesponsiveness.

Neither the clinical relevance nor the biologic basis of hyporesponsiveness is clear, but, given the possibility of its occurrence, more than one revaccination has not been recommended.

The effectiveness of PPSV23 against IPD, pneumococcal pneumonia, all-cause pneumonia, and death is controversial, with wide variation in observations. The many published meta-analyses of PPSV efficacy have often reached opposing conclusions with regard to a given clinical entity. Generally, observational studies cite greater effectiveness than do controlled clinical trials. The consensus is that PPSV is effective against IPD but is less effective or ineffective against nonbacteremic pneumococcal pneumonia. However, published trials, observational studies, and meta-analyses contradict this view. Efficacy is often lower in the elderly and in immunodeficient patients whose condition is associated with reduced antibody responses to vaccines than in younger, healthier populations. When PPSV is effective, the duration of protection following a single dose of vaccine is estimated to be ~5 years.

What is not disputed is that improved pneumococcal vaccines are needed for adults. Even in the setting of routine pneumococcal conjugate vaccination of infants (which indirectly protects adults from vaccine-serotype strains), disease caused by serotypes not represented in the conjugate vaccine continues to be a significant burden among adults.

Polysaccharide–Protein Conjugate Vaccines Infants and young children respond poorly to PPSV, which contains T cell–independent antigens. Consequently, another class of pneumococcal vaccines, the PCVs, were developed specifically for infants and young children. The first product, a 7-valent PCV, was licensed in 2000 in the United States. Three PCV products—containing 7, 10, and 13 serotypes, respectively—are currently (2014) commercially available. The serotypes included in these PCV formulations are important causes of IPD and antibiotic resistance among young children. Randomized controlled trials have demonstrated a high degree of efficacy of PCVs against vaccine-serotype IPD as well as efficacy against pneumonia, otitis media, nasopharyngeal colonization, and all-cause mortality. PCVs are recommended by the World Health Organization for inclusion in routine childhood immunization schedules worldwide, especially in countries with high infant mortality rates.

The United States was the first country to introduce PCV and therefore has the longest experience with its community-wide effects. The introduction of PCV in the United States has resulted in a >90% reduction in vaccine-serotype IPD among the whole population (Fig. 171-7). This decline has been noted not only in those age groups immunized but also in adults and is attributable to the near elimination of vaccine-serotype nasopharyngeal colonization in immunized infants, which reduces spread to adults. This protection of unimmunized community members through vaccination of a subset of the community is termed *the indirect effect*. Increases in colonization with—and concomitantly in disease due to—non-vaccine-serotype strains (i.e., replacement colonization and disease) have been seen; however, the absolute rate increases in IPD caused by non-vaccine serotypes are generally small, especially relative to decreases in vaccine-serotype IPD (see “Epidemiology,” above). Since vaccine-serotype strains are more commonly resistant to antibiotics than are non-vaccine serotypes, use of PCV has also resulted in dramatic declines in the proportion and absolute rates of drug-resistant pneumococcal disease. The recommendations of the Advisory Committee on Immunization Practices for the use of conjugate vaccines can be found at www.cdc.gov/MMWR/pdf/wk/mm5909.pdf. Recently, PCV has been shown to prevent pneumococcal infection in HIV-infected adults. In the United States, PCV13 followed by a dose of PPSV23 is now recommended for all immunocompromised children and adults.

Other Prevention Strategies Pneumococcal disease can also be averted through the prevention of illnesses that predispose individuals to pneumococcal infections. Relevant measures include influenza vaccination and improved management and control of diabetes, HIV infection, heart disease, and lung disease. Finally, the reduction of antibiotic misuse is a strategy for the prevention of pneumococcal disease in that

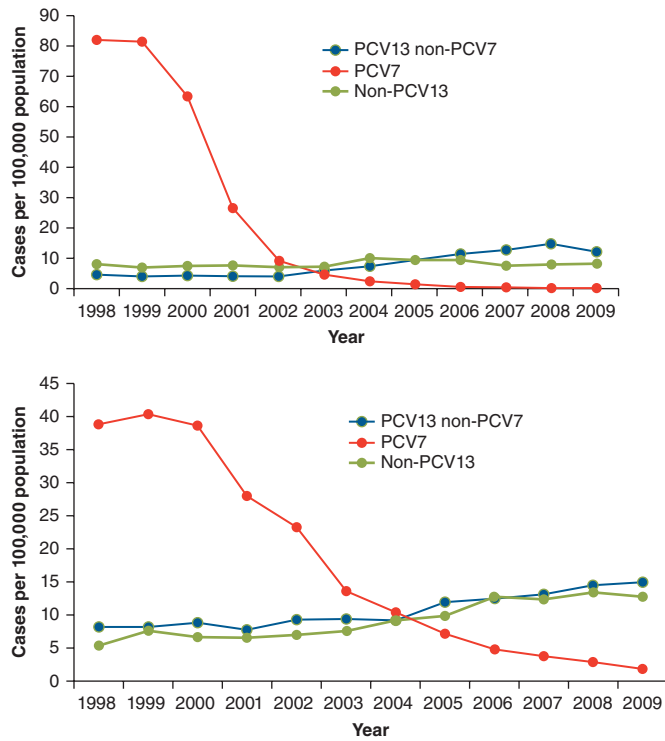


FIGURE 171-7 Changes in invasive pneumococcal disease (IPD) incidence, by serotype group, among children <5 years old (top) and adults >65 years old (bottom), 1998–2009. 7-Valent pneumococcal conjugate vaccine (PCV7) was introduced in the United States for routine administration to infants and young children during the second half of 2000, while PCV13 was introduced in 2010, the year following this surveillance period. PCV7 serotypes include serotypes 4, 6B, 9V, 14, 18C, 19F, and 23F as well as cross-reactive serotype 6A. PCV13 serotypes include the PCV7 serotypes as well as serotypes 1, 3, 5, 6A, 7F, and 19A. (Reprinted with permission from Dr. M. Moore, Centers for Disease Control and Prevention.)

antimicrobial resistance directly and indirectly perpetuates organism transmission and disease in the community.

WEBSITES

American Academy of Pediatrics RED BOOK. The report of the Committee on Infectious Diseases: aapredbook.aappublications.org; Pneumococcal Regional Serotype Distribution for Pneumococcal AMC TPP: www.gavialliance.org/library/documents/amc/tpp-codebook/

172 Staphylococcal Infections

Franklin D. Lowy

Staphylococcus aureus, the most virulent of the many staphylococcal species, has demonstrated its versatility by remaining a major cause of morbidity and mortality worldwide despite the availability of numerous effective antistaphylococcal antibiotics. *S. aureus* is a pluripotent pathogen, causing disease through both toxin- and non-toxin-mediated mechanisms. This organism is responsible for numerous nosocomial and community-based infections that range from relatively minor skin and soft tissue infections to life-threatening systemic infections.

The “other” staphylococci, collectively designated *coagulase-negative staphylococci* (CoNS), are considerably less virulent than *S. aureus* but remain important pathogens in infections that are primarily associated with prosthetic devices.

MICROBIOLOGY AND TAXONOMY

Staphylococci, gram-positive cocci in the family Micrococcaceae, form grapelike clusters on Gram’s stain (Fig. 172-1). These organisms (~1 μm in diameter) are catalase-positive (unlike streptococcal species), non-motile, aerobic, and facultatively anaerobic. They are capable of prolonged survival on environmental surfaces under varying conditions. Some species have a relatively broad host range, including mammals and birds, whereas for others the host range is quite narrow—i.e., limited to one or two closely related animals.

More than 30 staphylococcal species are pathogenic. Identification of the more clinically important species has generally relied on a series of biochemical tests. Automated diagnostic systems, kits for biochemical characterization, and DNA-based assays are available for species identification. With few exceptions, *S. aureus* is distinguished from other staphylococcal species by its production of coagulase, a surface enzyme that converts fibrinogen to fibrin. Latex kits that detect both protein A and clumping factor also distinguish *S. aureus* from most other staphylococcal species. *S. aureus* ferments mannitol, is positive for protein A, and produces DNase. On blood agar plates, *S. aureus* tends to form golden β -hemolytic colonies; in contrast, CoNS produce small white nonhemolytic colonies. Increasingly, sequence-based methods (e.g., 16S rRNA) are being used to identify different staphylococcal species.

Determining whether multiple staphylococcal isolates from different patients are the same or different is often relevant when there is concern that a nosocomial outbreak is due to a common point source (e.g., a contaminated medical instrument). Molecular typing methods, such as pulsed-field gel electrophoresis and sequence-based techniques (e.g., staphylococcal protein A [SpA] typing), have increasingly been used for this purpose. More recently, whole-genome sequencing has enhanced the ability to discriminate among clinical isolates.

S. AUREUS INFECTIONS

EPIDEMIOLOGY

S. aureus is both a commensal and an opportunistic pathogen. Approximately 30% of healthy persons are colonized with *S. aureus*, with a smaller percentage (~10%) persistently colonized. The rate of colonization is elevated among insulin-dependent diabetics, HIV-infected patients, patients undergoing hemodialysis, injection drug users, and individuals with skin damage. The anterior nares and oropharynx are frequent sites of human colonization, although the skin (especially when damaged), vagina, axilla, and perineum may also be colonized. These colonization sites serve as a reservoir for future infections.

Transmission of *S. aureus* most frequently results from direct personal contact. Colonization of different body sites allows transfer from one person to another during contact. Spread of staphylococci in aerosols of respiratory or nasal secretions from heavily colonized individuals has also been reported. Most individuals who develop

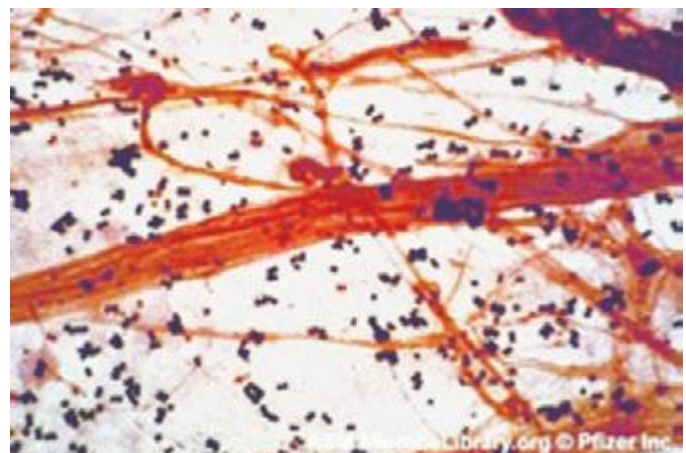


FIGURE 172-1 Gram’s stain of *S. aureus* in a sputum sample. (From ASM MicrobeLibrary.org. © Pfizer, Inc.)

S. aureus infections become infected with a strain that is already a part of their own commensal flora. Breaches of the skin or mucosal membrane allow *S. aureus* to initiate infection.

Some diseases increase the risk of *S. aureus* infection; diabetes, for example, combines an increased rate of *S. aureus* colonization and the use of injectable insulin with the possibility of impaired leukocyte function. Individuals with congenital or acquired qualitative or quantitative defects of polymorphonuclear leukocytes (PMNs) are at increased risk of *S. aureus* infections; this group includes neutropenic patients (e.g., those receiving chemotherapeutic agents), those with chronic granulomatous disease, and those with Job's or Chédiak-Higashi syndrome. Other groups at risk include individuals with end-stage renal disease, HIV infection, skin abnormalities, or prosthetic devices.

S. aureus is a leading cause of health care-associated infections (Chap. 168). It is the most common cause of surgical wound infections and is second only to CoNS as a cause of primary bacteremia. These isolates are generally resistant to multiple antibiotics; thus available therapeutic options are limited. In the community, *S. aureus* remains an important cause of skin and soft tissue infections, respiratory infections, and (among injection drug users) infective endocarditis. The increasing use of home infusion therapy is another cause of community-acquired staphylococcal infections.



In the past two decades, there has been a dramatic change in the epidemiology of infections due to methicillin-resistant *S. aureus* (MRSA). In addition to its major role as a nosocomial pathogen, MRSA has become an established community-based pathogen. Numerous outbreaks of community-associated MRSA (CA-MRSA) infections have been reported in both rural and urban settings in widely separated regions throughout the world. The outbreaks have occurred among such diverse groups as children, prisoners, athletes, Native Americans, and drug users. Risk factors common to these outbreaks include poor hygienic conditions, close contact, contaminated material, and damaged skin. These infections have been caused by a limited number of MRSA strains. In the United States, strain USA300 (defined by pulsed-field gel electrophoresis) has been the predominant clone. In other geographic regions of the world, different strains of CA-MRSA have been responsible for these community-based outbreaks. Although the majority of infections caused by these strains have involved the skin and soft tissue, 5–10% have been invasive and potentially life-threatening. CA-MRSA strains have also been responsible for an increasing number of nosocomial infections. Of concern has been the apparent capacity of CA-MRSA to cause disease in immunocompetent individuals.

PATHOGENESIS

General Concepts *S. aureus* is a pyogenic pathogen known for its capacity to induce abscess formation at sites of both local and metastatic infections. This classic pathologic response to *S. aureus* defines the framework within which the infection will progress. The bacteria elicit an inflammatory response characterized by an initial intense infiltration of PMNs and a subsequent infiltration of macrophages and fibroblasts. Either the host cellular response (including the deposition of fibrin and collagen) contains the infection, or infection spreads to the adjoining tissue or the bloodstream.

In toxin-mediated staphylococcal disease, infection is not invariably present. For example, once toxin has been elaborated into food, staphylococcal food poisoning can develop in the absence of viable bacteria. In staphylococcal toxic shock syndrome (TSS), conditions allowing toxin elaboration at colonization sites (e.g., the presence of a superabsorbent tampon) suffice for initiation of clinical illness.



The *S. aureus* Genome The complete genomes of numerous strains of *S. aureus* have now been fully sequenced. Among the interesting revelations are (1) the high degree of nucleotide sequence similarity of the core genomes of different strains; (2) acquisition of a relatively large amount of genetic information by horizontal transfer from other bacterial species; and (3) the presence of unique “pathogenicity” or “genomic” islands—mobile genetic elements that

contain clusters of enterotoxin and exotoxin genes and/or antimicrobial resistance determinants. Among the genes in these islands are those carrying *mecA*, the gene responsible for methicillin resistance. Methicillin resistance-containing islands have been designated staphylococcal cassette chromosome *mec* (SCC*mec*) types and range in size from ~20 to 60 kb. To date, 11 SCC*mec* types have been identified. Among the more common types, types 1–3 are traditionally associated with nosocomial MRSA isolates, whereas types 4–6 have been associated with the epidemic CA-MRSA strains.



A limited number of MRSA clones have been responsible for most community- and hospital-associated infections worldwide.

A comparison of these strains with those from earlier outbreaks (e.g., the phage 80/81 strains from the 1950s) has revealed preservation of the nucleotide sequence over time. This observation suggests that these strains possess determinants that facilitate survival and spread.



Regulation of Virulence Gene Expression In both toxin-mediated and non-toxin-mediated diseases due to *S. aureus*, the expression of virulence determinants associated with infection depends on a series of regulatory genes (e.g., accessory gene regulator [*agr*] and staphylococcal accessory regulator [*sar*]) that coordinately control the expression of many virulence genes. The regulatory gene *agr* is part of a quorum-sensing signal transduction pathway that senses and responds to bacterial density. Staphylococcal surface proteins are synthesized during the bacterial exponential growth phase *in vitro*. In contrast, many secreted proteins, such as a toxin, the enterotoxins, and assorted enzymes, are released during the postexponential growth phase in response to transcription of the effector molecule of *agr*, RNAIII.

It has been hypothesized that these regulatory genes serve a similar function *in vivo*. Successful invasion requires the sequential expression of these different bacterial elements. Bacterial adhesins are needed to initiate colonization of host tissue surfaces. The subsequent release of various enzymes enables the colony to obtain nutritional support and permits bacteria to spread to adjacent tissues. Studies with strains in which these regulatory genes are inactivated show reduced virulence in several animal models of *S. aureus* infection.

Pathogenesis of Invasive *S. aureus* Infection Staphylococci are opportunists. For these organisms to invade the host and cause infection, some or all of the following steps are necessary: contamination and colonization of host tissue surfaces, breach of cutaneous or mucosal barriers, establishment of a localized infection, invasion, evasion of the host response, and metastatic spread. Colonizing strains or strains transferred from other individuals are introduced into damaged skin, a wound, or the bloodstream. Recurrences of *S. aureus* infections are common, apparently because of the capacity of these pathogens to survive, to persist in a quiescent state in various tissues, and then to cause recrudescence infections when suitable conditions arise.

***S. AUREUS* COLONIZATION OF BODY SURFACES** The anterior nares is one of the primary sites of staphylococcal colonization in humans. Colonization appears to involve the attachment of *S. aureus* to keratinized epithelial cells of the anterior nares. Other factors that may contribute to colonization include the influence of other resident nasal flora and their bacterial density, host factors, and nasal mucosal damage (e.g., that resulting from inhalational drug use). Other colonized body sites, such as damaged skin, the groin, and the oropharynx, may be particularly important reservoirs for CA-MRSA strains.

INOCULATION AND COLONIZATION OF TISSUE SURFACES Staphylococci may be introduced into tissue as a result of minor abrasions, administration of medications such as insulin, or establishment of IV access with catheters. After their introduction into a tissue site, bacteria replicate and colonize the host tissue surface. A family of structurally related *S. aureus* surface proteins referred to as MSCRAMMs (microbial surface components recognizing adhesive matrix molecules) plays an important role in mediating adherence to these sites. By adhering to exposed matrix molecules (e.g., fibrinogen, fibronectin), MSCRAMMs such as clumping factor and collagen-binding protein enable the bacteria to colonize different tissue surfaces; these proteins contribute to

956 the pathogenesis of invasive infections such as endocarditis and septic arthritis by facilitating the adherence of *S. aureus* to surfaces with exposed fibrinogen or collagen.

Although CoNS are classically known for their ability to elaborate biofilms and to colonize prosthetic devices, *S. aureus* also possesses the genes responsible for biofilm formation, such as the intercellular adhesion (*ica*) locus. Binding to these devices occurs in a stepwise fashion, involving staphylococcal adherence to serum constituents that have coated the device surface and subsequent biofilm elaboration. *S. aureus* is thus a frequent cause of biomedical-device infections.

INVASION After colonization, staphylococci replicate at the initial site of infection, elaborating enzymes that include serine proteases, hyaluronidases, thermolysins, and lipases. These enzymes facilitate bacterial survival and local spread across tissue surfaces, although their precise role in infections is not well defined. The lipases may facilitate survival in lipid-rich areas such as the hair follicles, where *S. aureus* infections are often initiated. The *S. aureus* toxin Pantone-Valentine leukocidin is cytolytic to PMNs, macrophages, and monocytes. Strains elaborating this toxin have been epidemiologically linked with cutaneous and more serious infections caused by strains of CA-MRSA. MSCRAMMs also appear to play an important role in the ability of *S. aureus* to spread and cause disease at other tissue sites.

Constitutional findings may result from either localized or systemic infections. The staphylococcal cell wall—consisting of alternating N-acetyl muramic acid and N-acetyl glucosamine units in combination with an additional cell wall component, lipoteichoic acid—can initiate an inflammatory response that includes the sepsis syndrome. Staphylococcal α toxin, which causes pore formation in various eukaryotic cells, can also initiate an inflammatory response with findings suggestive of sepsis.

EVASION OF HOST DEFENSE MECHANISMS Staphylococci have a multitude of immune evasion strategies that are critical to their success as invasive pathogens. They possess an antiphagocytic polysaccharide microcapsule. Most human *S. aureus* infections are due to capsular types 5 and 8. The zwitterionic (both negatively and positively charged) *S. aureus* capsule plays a critical role in the induction of abscess formation. Protein A, an MSCRAMM unique to *S. aureus*, acts as an Fc receptor, binding the Fc portion of IgG subclasses 1, 2, and 4 and preventing opsonophagocytosis by PMNs. Both chemotaxis inhibitory protein of staphylococci (CHIPS, a secreted protein) and extracellular adherence protein (EAP, a surface protein) interfere with PMN migration to sites of infection.

An additional potential mechanism of *S. aureus* evasion is its capacity for intracellular survival. Both professional and nonprofessional phagocytes internalize staphylococci. Internalization by these cells may provide a sanctuary that protects bacteria against the host's defenses. The intracellular environment favors the phenotypic expression of *S. aureus* small-colony variants. Small-colony variants are found in patients receiving antimicrobial therapy (e.g., with aminoglycosides) and in those with cystic fibrosis or osteomyelitis. These variants, whether intra- or extracellular, may facilitate prolonged staphylococcal survival in different tissue sites and enhance the likelihood of recurrences. Finally, *S. aureus* can survive within PMNs and may use these cells to spread and to seed other tissue sites.

PATHOGENESIS OF COMMUNITY-ACQUIRED MRSA INFECTIONS A number of virulence determinants have been identified as contributing to the pathogenesis of CA-MRSA infections. There is a strong epidemiologic association linking the presence of the gene for the Pantone-Valentine leukocidin with skin and soft tissue infections as well as with necrotizing postinfluenza pneumonia. Other determinants that play a role in the pathogenesis of these infections include the arginine catabolic mobile element (ACME), a cluster of unique genes that may facilitate evasion of host defense mechanisms; phenol-soluble modulins, a family of cytolytic peptides; and a toxin.

Host Response to *S. aureus* Infection The primary host response to *S. aureus* infection is the recruitment of PMNs. These cells are attracted to infection sites by bacterial components such as formylated peptides or peptidoglycan as well as by the cytokines tumor necrosis factor

(TNF) and interleukins (ILs) 1 and 6, which are released by activated macrophages and endothelial cells.

Although most individuals have antibodies to staphylococci, it is not clear that antibody levels are qualitatively or quantitatively sufficient to protect against infection. Although anticapsular and anti-MSCRAMM antibodies facilitate opsonization in vitro and have been protective against infection in several animal models, they have not yet successfully prevented staphylococcal infections in clinical trials.

Pathogenesis of Toxin-Mediated Disease *S. aureus* produces three types of toxin: cytotoxins, pyrogenic toxin superantigens, and exfoliative toxins. Both epidemiologic data and studies in animals suggest that antitoxin antibodies are protective against illness in TSS, staphylococcal food poisoning, and staphylococcal scalded-skin syndrome (SSSS). Illness develops after toxin synthesis and absorption and the subsequent toxin-initiated host response.

ENTEROTOXIN AND TOXIC SHOCK SYNDROME TOXIN 1 (TSST-1) The pyrogenic toxin superantigens are a family of small-molecular-size, structurally similar proteins that are responsible for two diseases: TSS and food poisoning. TSS results from the ability of enterotoxins and TSST-1 to function as T cell mitogens. In the normal process of antigen presentation, the antigen is first processed within the cell, and peptides are then presented in the major histocompatibility complex (MHC) class II groove, initiating a measured T cell response. In contrast, enterotoxins bind directly to the invariant region of MHC—outside the MHC class II groove. The enterotoxins can then bind T cell receptors via the $v\beta$ chain; this binding results in a dramatic overexpansion of T cell clones (up to 20% of the total T cell population). The consequence of this T cell expansion is a “cytokine storm,” with the release of inflammatory mediators that include interferon γ , IL-1, IL-6, TNF- α , and TNF- β . The resulting multisystem disease produces a constellation of findings that mimic those in endotoxin shock; however, the pathogenic mechanisms differ. The release of endotoxin from the gastrointestinal tract may synergistically enhance the toxin's effects.

A different region of the enterotoxin molecule is responsible for the symptoms of food poisoning. The enterotoxins are heat stable and can survive conditions that kill the bacteria. Illness results from the ingestion of preformed toxin. As a result, the incubation period is short (1–6 h). The toxin stimulates the vagus nerve and the vomiting center of the brain. It also appears to stimulate intestinal peristaltic activity.

EXFOLIATIVE TOXINS AND SSSS The exfoliative toxins are responsible for SSSS. The toxins that produce disease in humans are of two serotypes: ETA and ETB. These toxins are serine proteases, which cleave desmosomal cadherins in the superficial layer of the skin, triggering exfoliation. The result is a split in the epidermis at the granular level, which is responsible for the superficial desquamation of the skin that typifies this illness.

DIAGNOSIS

Staphylococcal infections are readily diagnosed by Gram's stain (Fig. 172-1) and microscopic examination of abscess contents or of infected tissue. Routine culture of infected material usually yields positive results, and blood cultures are sometimes positive even when infections are localized to extravascular sites. *S. aureus* is rarely a blood culture contaminant. Polymerase chain reaction (PCR)-based assays have been applied to the rapid diagnosis of *S. aureus* infection and are increasingly used in clinical microbiology laboratories. A number of point-of-care tests are now available to screen patients for colonization with MRSA. Determining whether patients with documented *S. aureus* bacteremia also have infective endocarditis or a metastatic focus of infection remains a diagnostic challenge. Uniformly positive blood cultures suggest an endovascular infection such as endocarditis (see “Bacteremia, Sepsis, and Infective Endocarditis,” below).

CLINICAL SYNDROMES (Table 172-1)

Skin and Soft Tissue Infections *S. aureus* causes a variety of cutaneous infections, many of which can also be caused by group A streptococci

TABLE 172-1 COMMON ILLNESSES CAUSED BY STAPHYLOCOCCUS AUREUS

Skin and Soft Tissue Infections
Folliculitis
Abscess, furuncle, carbuncle
Cellulitis
Impetigo
Mastitis
Surgical wound infections
Musculoskeletal Infections
Septic arthritis
Osteomyelitis (hematogenous or contiguous spread)
Pyomyositis
Psoas abscess
Respiratory Tract Infections
Ventilator-associated or nosocomial pneumonia
Septic pulmonary emboli
Postviral pneumonia (e.g., influenza)
Empyema
Bacteremia and its Complications
Sepsis, septic shock
Metastatic foci of infection (kidney, joints, bone, lung)
Infective endocarditis
Infective Endocarditis
Injection drug use–associated
Native-valve
Prosthetic-valve
Nosocomial
Device-Related Infections (e.g., intravascular catheters, prosthetic joints)
Toxin-Mediated Illnesses
Toxic shock syndrome
Food poisoning
Staphylococcal scalded-skin syndrome
Invasive Infections Associated with Community-Acquired Methicillin-Resistant <i>S. aureus</i>
Necrotizing fasciitis
Waterhouse-Friderichsen syndrome
Necrotizing pneumonia
Purpura fulminans

or (less commonly) other streptococcal species. Common factors predisposing to *S. aureus* cutaneous infection include chronic skin conditions (e.g., eczema), skin damage (e.g., insect bites, minor trauma), injections (e.g., in diabetes, injection drug use), and poor personal hygiene. These infections are characterized by the formation of pus-containing blisters, which often begin in hair follicles and spread to adjoining tissues. *Folliculitis* is a superficial infection that involves the hair follicle, with a central area of purulence (pus) surrounded by induration and erythema. *Furuncles* (boils) are more extensive, painful lesions that tend to occur in hairy, moist regions of the body and extend from the hair follicle to become a true abscess with an area of central purulence. *Carbuncles* are most often located in the lower neck and are even more severe and painful, resulting from the coalescence of other lesions that extend to a deeper layer of the subcutaneous tissue. In general, furuncles and carbuncles are readily apparent, with pus often expressible or discharging from the abscess. Other cutaneous *S. aureus* infections include impetigo and cellulitis. *S. aureus* is one of the most common causes of surgical wound infection.

Mastitis develops in 1–3% of nursing mothers. This infection of the breast, which generally presents within 2–3 weeks after delivery, is characterized by findings that range from cellulitis to abscess formation. Systemic signs, such as fever and chills, are often present in more severe cases.

Musculoskeletal Infections *S. aureus* is among the most common causes of bone infections—both those resulting from hematogenous dissemination and those arising from contiguous spread from a soft tissue site. *Hematogenous osteomyelitis* in children most often involves the long bones. Infections present as fever and bone pain or with a child's reluctance to bear weight. The white blood cell count and erythrocyte sedimentation rate are often elevated. Blood cultures are positive in ~50% of cases. When necessary, bone biopsies for culture and histopathologic examination are usually diagnostic. Routine x-rays may be normal for up to 14 days after the onset of symptoms. ^{99m}Tc-phosphonate scanning often detects early evidence of infection. MRI is more sensitive than other techniques in establishing a radiologic diagnosis.

In adults, hematogenous osteomyelitis involving the long bones is less common. However, *vertebral osteomyelitis* is among the more common clinical presentations. Vertebral bone infections are most often seen in patients with endocarditis, those undergoing hemodialysis, diabetics, and injection drug users. These infections may present as intense back pain and fever but may also be clinically occult, presenting as chronic back pain and low-grade fever. *S. aureus* is the most common cause of epidural abscess, a complication that can result in neurologic compromise. Patients complain of difficulty voiding or walking and of radicular pain in addition to the symptoms associated with their osteomyelitis. Surgical intervention in this setting often constitutes a medical emergency. MRI most reliably establishes the diagnosis (Fig. 172-2).

Bone infections that result from contiguous spread tend to develop from soft tissue infections, such as those associated with diabetic or vascular ulcers, surgery, or trauma. Exposure of bone, a draining fistulous tract, failure to heal, or continued drainage suggests involvement of underlying bone. Bone involvement is established by bone culture and histopathologic examination (revealing evidence of PMN infiltration). Contamination of culture material from adjacent tissue can make the diagnosis of osteomyelitis difficult in the absence of pathologic confirmation. In addition, it is sometimes hard to distinguish radiologically between osteomyelitis and overlying soft tissue infection with underlying osteitis.

In both children and adults, *S. aureus* is the most common cause of *septic arthritis* in native joints. This infection is rapidly progressive and may be associated with extensive joint destruction if left untreated. It presents as intense pain on motion of the affected joint, swelling, and fever. Aspiration of the joint reveals turbid fluid, with >50,000 PMNs/ μ L and gram-positive cocci in clusters on Gram's stain (Fig. 172-1). In

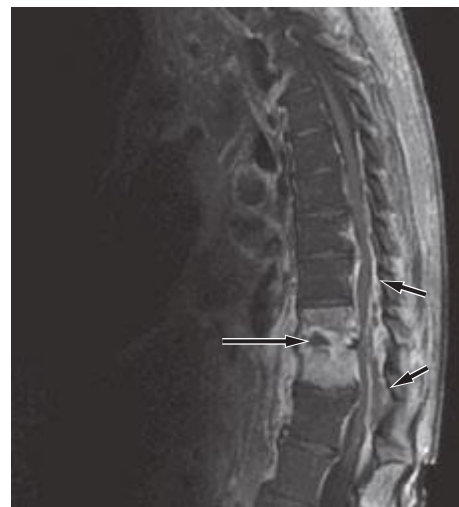


FIGURE 172-2 *S. aureus* vertebral osteomyelitis and epidural abscess involving the thoracic disk between T9 and T10. Sagittal postcontrast MRI of the spine illustrates destruction of the T9–T10 intervertebral space with enhancement (arrow). There is impingement on the thoracic cord and an epidural collection extending from T9 through T11 (short arrows).

958 adults, septic arthritis may result from trauma, surgery, or hematogenous dissemination. The most commonly involved joints include the knees, shoulders, hips, and phalanges. Infection frequently develops in joints previously damaged by osteoarthritis or rheumatoid arthritis. Iatrogenic infections resulting from aspiration or injection of agents into the joint also occur. In these settings, the patient experiences increased pain and swelling in the involved joint in association with fever.

Pyomyositis is an unusual infection of skeletal muscles that is seen primarily in tropical climates but also occurs in immunocompromised and HIV-infected patients. It is believed to arise from occult bacteremia. *Pyomyositis* presents as fever, swelling, and pain overlying the involved muscle. Aspiration of fluid from the involved tissue yields pus. Although a history of trauma may be associated with the infection, its pathogenesis is poorly understood.

Respiratory Tract Infections Respiratory tract infections caused by *S. aureus* occur in selected clinical settings. *S. aureus* is a cause of serious respiratory tract infections in newborns and infants; these infections present with shortness of breath, fever, and respiratory failure. Chest x-ray may reveal pneumatoceles (shaggy, thin-walled cavities). Pneumothorax and empyema are recognized complications.

In adults, nosocomial *S. aureus* pulmonary infections are common among intubated patients in intensive care units. Nasally colonized patients are at increased risk of these infections. The clinical presentation is no different from that encountered in pulmonary infections of other bacterial etiologies. Patients produce increased volumes of purulent sputum and develop respiratory distress, fever, and new pulmonary infiltrates. Distinguishing bacterial pneumonia from respiratory failure or other causes of new pulmonary infiltrates in critically ill patients is often difficult and relies on a constellation of clinical, radiologic, and laboratory findings.

Community-acquired respiratory tract infections due to *S. aureus* usually follow viral infections—most commonly influenza. Patients may present with fever, bloody sputum production, and midlung-field pneumatoceles or multiple, patchy pulmonary infiltrates. Diagnosis is made by sputum Gram's stain and culture. Blood cultures, although useful, are usually negative.

Bacteremia, Sepsis, and Infective Endocarditis *S. aureus* bacteremia may be complicated by sepsis, endocarditis, vasculitis, or metastatic seeding (establishment of suppurative collections at other tissue sites). The frequency of metastatic seeding during bacteremia has been estimated to be as high as 31%. Among the more commonly seeded tissue sites are bones, joints, kidneys, and lungs.

Recognition of these complications by clinical and laboratory diagnostic methods alone is often difficult. Comorbid conditions that are frequently seen in association with *S. aureus* bacteremia and that increase the risk of complications include diabetes, HIV infection, and renal insufficiency. Other host factors associated with an increased risk of complications include presentation with community-acquired *S. aureus* bacteremia (except in injection drug users), lack of an identifiable primary focus of infection, and the presence of prosthetic devices or material.

Clinically, *S. aureus* sepsis presents in a manner similar to that documented for sepsis due to other bacteria. The well-described progression of hemodynamic changes—beginning with respiratory alkalosis and clinical findings of hypotension and fever—is commonly seen. The microbiologic diagnosis is established by positive blood cultures.

The overall incidence of *S. aureus* endocarditis has increased over the past 20 years. *S. aureus* is now the leading cause of endocarditis worldwide, accounting for 25–35% of cases. This increase is due, at least in part, to the increased use of intravascular devices. Studies of patients with *S. aureus* bacteremia and intravascular catheters that used transesophageal echocardiography found an infective endocarditis incidence of ~25%. Other factors associated with an increased risk of endocarditis are injection drug use, hemodialysis, the presence of intravascular prosthetic devices at the time of bacteremia, and immunosuppression. Patients with implantable cardiac devices (e.g., permanent pacemakers) are at increased risk of endocarditis or device-related infections. Despite the availability of effective antibiotics, mortality rates

from these infections continue to range from 20% to 40%, depending on both the host and the nature of the infection. Complications of *S. aureus* endocarditis include cardiac valvular insufficiency, peripheral emboli, metastatic seeding, and central nervous system (CNS) involvement (e.g., mycotic aneurysms, embolic strokes).

S. aureus endocarditis is encountered in four clinical settings: (1) right-sided endocarditis in association with injection drug use, (2) left-sided native-valve endocarditis, (3) prosthetic-valve endocarditis, and (4) nosocomial endocarditis. In each of these settings, the diagnosis is suspected by recognition of clinical stigmata suggestive of endocarditis. These findings include cardiac manifestations, such as new or changing cardiac valvular murmurs; cutaneous evidence, such as vasculitic lesions, Osler's nodes, or Janeway lesions; evidence of right- or left-sided embolic disease; and a history suggesting a risk for *S. aureus* bacteremia. In the absence of antecedent antibiotic therapy, blood cultures are almost uniformly positive. Transthoracic echocardiography, while less sensitive than transesophageal echocardiography, is less invasive and may establish the presence of valvular vegetations. The Duke criteria (see Table 155-3) are now commonly used to help establish the likelihood of this diagnosis.

Acute right-sided tricuspid valvular *S. aureus* endocarditis is most often seen in injection drug users. The classic presentation includes a high fever, a toxic clinical appearance, pleuritic chest pain, and the production of purulent (sometimes bloody) sputum. Chest x-rays or CT scans reveal evidence of septic pulmonary emboli (small, peripheral, circular lesions that may cavitate with time) (Fig. 172-3). A high percentage of affected patients have no history of antecedent valvular damage. At the outset of their illness, patients may present with fever alone, without cardiac or other localizing findings. As a result, a high index of clinical suspicion is essential for diagnosis.

Individuals with antecedent cardiac valvular damage more commonly present with left-sided native-valve endocarditis involving the damaged valve. These patients tend to be older than those with right-sided endocarditis, their prognosis is worse, and their incidence of complications (including peripheral emboli, cardiac decompensation, and metastatic seeding) is higher.

S. aureus is one of the more common causes of prosthetic-valve endocarditis. This infection is especially fulminant in the early postoperative period and is associated with a high mortality rate. In most instances, medical therapy alone is not sufficient and urgent valve replacement is necessary. Patients are prone to develop valvular insufficiency or myocardial abscesses originating from the region of valve implantation.

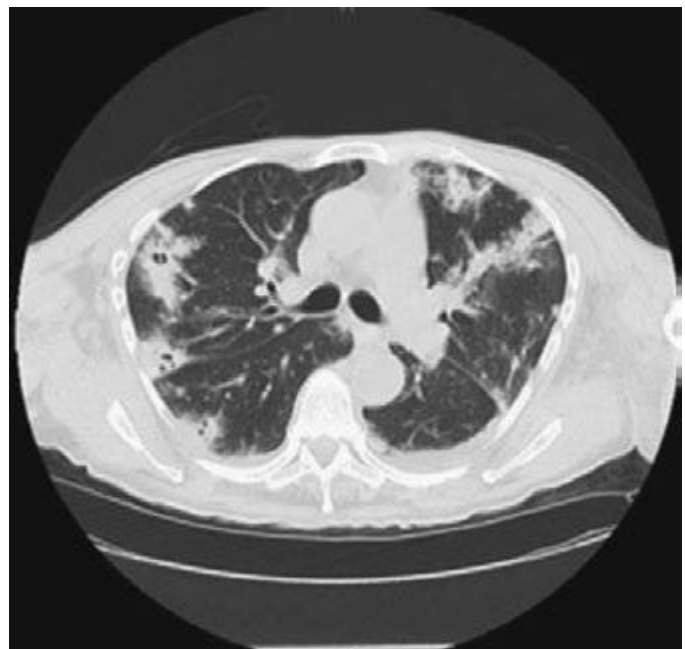


FIGURE 172-3 CT scan illustrating septic pulmonary emboli in a patient with methicillin-resistant *Staphylococcus aureus* bacteremia.

The increased frequency of nosocomial endocarditis (15–30% of cases, depending on the series) reflects in part the increased use of intravascular devices. This form of endocarditis is most commonly caused by *S. aureus*. Because patients often are critically ill, are receiving antibiotics for various other indications, and have comorbid conditions, the diagnosis is often missed.

Urinary Tract Infections Urinary tract infections (UTIs) are infrequently caused by *S. aureus*. The presence of *S. aureus* in the urine generally suggests hematogenous dissemination. Ascending *S. aureus* infections occasionally result from instrumentation of the genitourinary tract.

Prosthetic Device–Related Infections *S. aureus* accounts for a large proportion of prosthetic device–related infections. These infections often involve intravascular catheters, prosthetic valves, orthopedic devices, peritoneal catheters, pacemakers, left-ventricular-assist devices, and vascular grafts. In contrast with the more indolent presentation of CoNS infections, *S. aureus* device-related infections are often more acute, with both localized and systemic manifestations. The latter infections also tend to progress more rapidly. It is relatively common for a pyogenic collection to be present at the device site. Aspiration of these collections and performance of blood cultures are important components in establishing a diagnosis. *S. aureus* infections tend to occur more commonly soon after implantation unless the device is used for access (e.g., intravascular or hemodialysis catheters). In the latter instance, infections can occur at any time. As in most prosthetic-device infections, successful therapy usually involves removal of the device. Left in place, the device is a potential nidus for either persistent or recurrent infections.

Infections Associated with Community-Acquired MRSA Although the skin and soft tissues are the most common sites of infection associated with CA-MRSA, 5–10% of these infections are invasive and can even be life-threatening. The latter unique infections, including necrotizing fasciitis, necrotizing pneumonia, and sepsis with Waterhouse-Friderichsen syndrome or purpura fulminans, were rarely associated with *S. aureus* prior to the emergence of CA-MRSA. These life-threatening infections reflect the increased virulence of CA-MRSA strains.

Toxin-Mediated Diseases • FOOD POISONING *S. aureus* is among the most common causes of foodborne outbreaks of infection in the United States. Staphylococcal food poisoning results from the inoculation of toxin-producing *S. aureus* into food by colonized food handlers. Toxin is then elaborated in such growth-promoting food as custards, potato salad, or processed meats. Even if the bacteria are killed by warming, the heat-stable toxin is not destroyed. The onset of illness is rapid, occurring within 1–6 h of ingestion. The illness is characterized by nausea and vomiting, although diarrhea, hypotension, and dehydration may also occur. The differential diagnosis includes diarrhea of other etiologies, especially that caused by similar toxins (e.g., the toxins elaborated by *Bacillus cereus*). The rapidity of onset, the absence of fever, and the epidemic nature of the presentation (without second-degree spread) arouse suspicion of staphylococcal food poisoning. Symptoms generally resolve within 8–10 h. The diagnosis can be established by the demonstration of bacteria or the documentation of enterotoxin in the implicated food. Treatment is entirely supportive.

TOXIC SHOCK SYNDROME TSS gained attention in the early 1980s, when a nationwide outbreak occurred in the United States among young, otherwise healthy, menstruating women. Epidemiologic investigation demonstrated that these cases were associated with the use of a highly absorbent tampon that had recently been introduced to the market. Subsequent studies established the role of TSST-1 in these illnesses. Withdrawal of the tampon from the market resulted in a rapid decline in the incidence of this disease. However, menstrual and nonmenstrual cases continue to be reported. Nonmenstrual cases are frequently seen in patients with surgical or postpartum wound infections.

The clinical presentation is similar in menstrual and nonmenstrual TSS. Evidence of clinical *S. aureus* infection is not a prerequisite.

TABLE 172-2 CASE DEFINITION OF *S. AUREUS* TOXIC SHOCK SYNDROME

1. Fever: temperature of $\geq 38.9^{\circ}\text{C}$ ($\geq 102^{\circ}\text{F}$)
2. Hypotension: systolic blood pressure of ≤ 90 mmHg or orthostatic hypotension (orthostatic drop in diastolic blood pressure by ≥ 15 mmHg, orthostatic syncope, or orthostatic dizziness)
3. Diffuse macular rash, with desquamation 1–2 weeks after onset (including the palms and soles)
4. Multisystem involvement
 - a. Hepatic: bilirubin or aminotransferase levels ≥ 2 times normal
 - b. Hematologic: platelet count $\leq 100,000/\mu\text{L}$
 - c. Renal: blood urea nitrogen or serum creatinine level ≥ 2 times the normal upper limit
 - d. Mucous membranes: vaginal, oropharyngeal, or conjunctival hyperemia
 - e. Gastrointestinal: vomiting or diarrhea at onset of illness
 - f. Muscular: severe myalgias or serum creatine phosphokinase level ≥ 2 times the normal upper limit
 - g. Central nervous system: disorientation or alteration in consciousness without focal neurologic signs and in the absence of fever and hypotension
5. Negative serologic or other tests for measles, leptospirosis, and Rocky Mountain spotted fever as well as negative blood or cerebrospinal fluid cultures for organisms other than *S. aureus*

Source: M Wharton et al: Case definitions for public health surveillance. MMWR 39:1, 1990; with permission.

TSS results from the elaboration of an enterotoxin or the structurally related enterotoxin-like TSST-1. More than 90% of menstrual cases are caused by TSST-1, whereas a high percentage of nonmenstrual cases are caused by enterotoxins. TSS begins with relatively nonspecific flu-like symptoms. In menstrual cases, the onset usually comes 2 or 3 days after the start of menstruation. Patients present with fever, hypotension, and erythroderma of variable intensity. Mucosal involvement is common (e.g., conjunctival hyperemia). The illness can rapidly progress to symptoms that include vomiting, diarrhea, confusion, myalgias, and abdominal pain. These symptoms reflect the multisystemic nature of the disease, with involvement of the liver, kidneys, gastrointestinal tract, and/or CNS. Desquamation of the skin occurs during convalescence, usually 1–2 weeks after the onset of illness. Laboratory findings may include azotemia, leukocytosis, hypoalbuminemia, thrombocytopenia, and liver function abnormalities.

Diagnosis of TSS still depends on a constellation of findings rather than one specific finding and on a lack of evidence of other possible infections (Table 172-2). Other diagnoses to be considered are drug toxicities, viral exanthems, Rocky Mountain spotted fever, sepsis, and Kawasaki disease. Illness occurs only in persons who lack antibody to TSST-1. Recurrences are possible if antibody fails to develop after the illness.

STAPHYLOCOCCAL SCALDED-SKIN SYNDROME SSSS primarily affects newborns and children. The illness may vary from a localized blister to exfoliation of much of the skin surface. The skin is usually fragile and often tender, with thin-walled, fluid-filled bullae. Gentle pressure results in rupture of the lesions, leaving denuded underlying skin (*Nikolsky's sign*; Fig. 172-4). The mucous membranes are usually spared. In more generalized infection, there are often constitutional symptoms, including fever, lethargy, and irritability with poor feeding. Significant amounts of fluid can be lost in more extensive cases. Illness usually follows localized infection at one of a number of possible sites. SSSS is much less common among adults but can follow infections caused by exfoliative toxin–producing strains.

PREVENTION

Primary prevention of *S. aureus* infections in the hospital setting involves hand washing and careful attention to appropriate isolation procedures. Through careful screening for MRSA carriage and strict isolation practices, several Scandinavian countries have been remarkably



FIGURE 172-4 Evidence of staphylococcal scalded-skin syndrome in a 6-year-old boy. Nikolsky's sign, with separation of the superficial layer of the outer epidermal layer, is visible. (Reprinted with permission from LA Schenfeld et al: *N Engl J Med* 342:1178, 2000. © 2000 Massachusetts Medical Society. All rights reserved.)

successful at preventing the introduction and dissemination of MRSA in hospitals.

Decolonization strategies, using both universal and targeted approaches with topical agents (e.g., mupirocin) to eliminate nasal colonization and/or chlorhexidine to eliminate cutaneous colonization with *S. aureus*, have been successful in some clinical settings (e.g., intensive care units) where the risk of infection is high. An analysis of clinical trials suggests that there may also be a reduction in the incidence of postsurgical infections among persons who are nasally colonized with *S. aureus*.

“Bundling” (the application of selected medical interventions in a sequence of prescribed steps) has reduced rates of nosocomial infections related to such procedures as the insertion of intravenous catheters, in which staphylococci are among the most common pathogens (see Table 168-4). A number of immunization strategies to prevent *S. aureus* infections—both active (e.g., capsular polysaccharide–protein conjugate vaccine) and passive (e.g., clumping factor antibody)—have been investigated. However, none has been successful for either prophylaxis or therapy in clinical trials.

COAGULASE-NEGATIVE STAPHYLOCOCCAL INFECTIONS

Although considerably less virulent than *S. aureus*, CoNS are among the most common causes of prosthetic-device infections. Approximately half of the identified CoNS species have been associated with human infections. Of these species, *Staphylococcus epidermidis* is the most common human pathogen. This component of the normal human flora is found on the skin (where it is the most abundant bacterial species) as well as in the oropharynx and vagina. *Staphylococcus saprophyticus*, a novobiocin-resistant species, is a common pathogen in UTIs.

PATHOGENESIS

S. epidermidis is the CoNS species most often associated with prosthetic-device infections. Infection is a two-step process, with initial adhesion to the device followed by colonization. *S. epidermidis* is uniquely adapted to colonize these devices by its capacity to elaborate the extracellular polysaccharide (glycocalyx or slime) that facilitates formation of a protective biofilm on the device surface.

Implanted prosthetic material is rapidly coated with host serum or tissue constituents such as fibrinogen or fibronectin. These molecules serve as potential bridging ligands, facilitating initial bacterial attachment to the device surface. A number of staphylococcal surface-associated proteins, such as autolysin (AtlE), fibrinogen-binding protein, and accumulation-associated protein (AAP), may play a role in attachment to either modified or unmodified prosthetic surfaces. The polysaccharide intercellular adhesin facilitates subsequent

staphylococcal colonization and accumulation on the device surface. In *S. epidermidis*, intercellular adhesin (*ica*) genes are more commonly found in strains associated with device infections than in strains associated with colonization of mucosal surfaces. Biofilm appears to act as a barrier protecting bacteria from host defense mechanisms as well as from antibiotics, while providing a suitable environment for bacterial survival. Poly- γ -DL-glutamic acid is secreted by *S. epidermidis* and provides protection against neutrophil phagocytosis.

Two additional staphylococcal species, *Staphylococcus lugdunensis* and *Staphylococcus schleiferi*, produce more serious infections (native-valve endocarditis and osteomyelitis) than do other CoNS. The basis for this enhanced virulence is not known, although both species appear to share more virulence determinants with *S. aureus* (e.g., clumping factor and lipase) than do other CoNS.

The capacity of *S. saprophyticus* to cause UTIs in young women appears to be related to its enhanced capacity to adhere to uroepithelial cells. A 160-kDa hemagglutinin/adhesin may contribute to this affinity.

DIAGNOSIS

Although the detection of CoNS at sites of infection or in the bloodstream is not difficult by standard microbiologic culture methods, interpretation of these results is frequently problematic. Because these organisms are present in large numbers on the skin, they often contaminate cultures. It has been estimated that only 10–25% of blood cultures positive for CoNS reflect true bacteremia. Similar problems arise with cultures obtained from other sites. Among the clinical findings suggestive of true bacteremia are fever, evidence of local infection (e.g., erythema or purulent drainage at the IV catheter site), leukocytosis, and systemic signs of sepsis. Laboratory findings suggestive of true bacteremia include multiple isolations of the same strain (i.e., the same species with the same antibiogram or a closely related DNA fingerprint) from separate cultures, growth of the strain within 48 h, and bacterial growth in both aerobic and anaerobic bottles.

CLINICAL SYNDROMES

CoNS cause a diverse array of prosthetic device–related infections, including those that involve prosthetic cardiac valves and joints, vascular grafts, intravascular devices, and CNS shunts. In all of these settings, the clinical presentation is similar. The signs of localized infection are often subtle, the rate of disease progression is slow, and the systemic findings are often limited. Signs of infection, such as purulent drainage, pain at the site, or loosening of prosthetic implants, are sometimes evident. Fever is frequently but not always present, and there may be mild leukocytosis. Acute-phase reactant levels, the erythrocyte sedimentation rate, and the C-reactive protein concentration may be elevated.

Infections that are not associated with prosthetic devices are infrequent, although native-valve endocarditis due to CoNS has accounted for ~5% of cases in some reviews. *S. lugdunensis* appears to be a more aggressive pathogen in this setting, causing greater mortality and rapid valvular destruction with abscess formation.

TREATMENT STAPHYLOCOCCAL INFECTIONS

GENERAL PRINCIPLES OF THERAPY

Surgical incision and drainage of all suppurative collections constitute the most important therapeutic intervention for staphylococcal infections. The emergence of MRSA in the community has increased the importance of culturing all collections in order to identify pathogens and to determine antimicrobial susceptibility. Successful therapy for prosthetic-device infections generally requires device removal. In situations in which removal is not possible or the infection is due to CoNS, an initial attempt at medical therapy without device removal may be warranted. Because of the well-recognized risk of complications associated with *S. aureus* bacteremia (e.g., endocarditis, metastatic foci of infection), therapy is generally prolonged (4–6 weeks) unless the patient is identified as being among those individuals who are at low risk for complications.

DURATION OF ANTIMICROBIAL THERAPY

Debate continues regarding the duration of therapy for bacteremic *S. aureus* infections. Patients with “complicated” bacteremia are at increased risk of endocarditis and metastatic infections. Among the findings associated with an increased risk of complicated bacteremia are persistently positive blood cultures 96 h after institution of therapy, acquisition of the infection in the community, failure to remove a removable focus of infection (i.e., an intravascular catheter), and infection with cutaneous or embolic manifestations. For immunocompetent patients in whom short-course therapy is planned, transesophageal echocardiography to rule out endocarditis is warranted because neither clinical nor laboratory findings can reliably detect cardiac involvement. In addition, an aggressive radiologic investigation to identify potential metastatic collections is indicated. All symptomatic body sites must be carefully evaluated.

CHOICE OF ANTIMICROBIAL AGENTS

The choice of antimicrobial agents to treat both coagulase-positive and coagulase-negative staphylococcal infections has become increasingly problematic because of the prevalence of multidrug-resistant strains. Staphylococcal resistance to most antibiotic families, including β -lactams, aminoglycosides, fluoroquinolones, and (to a lesser extent) glycopeptides, has increased. This trend is more apparent with CoNS: >80% of nosocomial isolates are resistant to methicillin, and these methicillin-resistant strains are usually resistant to most other antibiotics as well. Because the selection of antimicrobial agents for *S. aureus* infections is similar to that for CoNS infections, treatment options for these pathogens are discussed together and are summarized in [Table 172-3](#).

As a result of the widespread dissemination of plasmids containing the enzyme penicillinase, few strains of staphylococci ($\leq 5\%$) remain susceptible to penicillin. However, penicillin remains the drug of choice against susceptible strains if the laboratory can reliably test for penicillin susceptibility. Penicillin-resistant isolates are treated with semisynthetic penicillinase-resistant penicillins (SPRPs), such as oxacillin or nafcillin. Methicillin, the first of the SPRPs, is now used infrequently. Cephalosporins are alternative therapeutic agents for these infections. Second- and third-generation cephalosporins do not offer a therapeutic advantage over first-generation cephalosporins for the treatment of staphylococcal infections. The carbapenems have excellent activity against methicillin-sensitive *S. aureus* but not against MRSA.

The isolation of MRSA was reported within 1 year of the introduction of methicillin. Since then, the prevalence of MRSA has steadily increased. In many hospitals, 40–50% of *S. aureus* isolates are now resistant to methicillin. Resistance to methicillin indicates resistance to all SPRPs as well as to all cephalosporins (except ceftaroline). Production of a novel penicillin-binding protein (PBP2a) is responsible for methicillin resistance. This protein is synthesized by the *mecA* gene, which (as stated above) is part of a large mobile genetic element—a pathogenicity or genomic island—called *SCCmec*. It is hypothesized that this genetic material was acquired via horizontal transfer from a related staphylococcal species, such as *Staphylococcus sciuri*. Phenotypic expression of methicillin resistance may be constitutive (i.e., expressed in all organisms in a population) or heterogeneous (i.e., displayed by only a proportion of the total organism population). Detection of methicillin resistance in the clinical microbiology laboratory can be difficult if the strain expresses heterogeneous resistance. Therefore, susceptibility studies are routinely performed at reduced temperatures ($\leq 35^\circ\text{C}$ for 24 h), with increased concentrations of salt in the medium to enhance the expression of resistance. In addition to PCR-based techniques, a number of rapid methods for the detection of methicillin resistance have been developed.

In light of decreasing susceptibility of MRSA isolates to vancomycin, both vancomycin and daptomycin are now recommended as the drugs of choice for the treatment of MRSA infections. Vancomycin is less effective than SPRPs for the treatment of infections due to methicillin-susceptible strains. Alternatives to SPRPs should be used only after careful consideration in patients with a history of serious β -lactam allergies.

Three types of staphylococcal resistance to vancomycin have emerged. (1) Minimal inhibitory concentration (MIC) “creep” refers to the incremental increase in vancomycin MICs that has been detected in various geographic areas. Studies suggest that infections due to *S. aureus* strains with vancomycin MICs of $>1 \mu\text{g}/\text{mL}$ may not respond as well to vancomycin therapy as those due to strains with MICs of $<1 \mu\text{g}/\text{mL}$. Some authorities (e.g., *The Medical Letter*) have recommended choosing an alternative agent in this setting. (2) In 1997, an *S. aureus* strain with reduced susceptibility to vancomycin (vancomycin-intermediate *S. aureus* [VISA]) was reported from Japan. Subsequently, additional VISA clinical isolates were reported. These strains were all resistant to methicillin and many other antimicrobial agents. The VISA strains appear to evolve (under vancomycin selective pressure) from strains that are susceptible to vancomycin but are heterogeneous, with a small proportion of the bacterial population expressing the resistance phenotype. The mechanism of VISA resistance is in part due to an abnormally thick cell wall. Vancomycin is trapped by the abnormal peptidoglycan cross-linking and is unable to gain access to its target site. (3) In 2002, the first clinical isolate of fully vancomycin-resistant *S. aureus* was reported. Resistance in this and several additional clinical isolates was due to the presence of *vanaA*, the gene responsible for expression of vancomycin resistance in enterococci. This observation suggested that resistance was acquired as a result of horizontal conjugal transfer from a vancomycin-resistant strain of *Enterococcus faecalis*. Several patients had both MRSA and vancomycin-resistant enterococci cultured from infection sites. The *vanaA* gene is responsible for the synthesis of the dipeptide D-Ala-D-Lac in place of D-Ala-D-Ala. Vancomycin cannot bind to the altered peptide.

Daptomycin, a parenteral bactericidal agent with antistaphylococcal activity, is approved for the treatment of bacteremia (including right-sided endocarditis) and complicated skin infections. It is not effective in respiratory infections. This drug has a novel mechanism of action: it disrupts the cytoplasmic membrane. Staphylococcal resistance to daptomycin, sometimes developing during therapy, has been reported.

Linezolid—the first oxazolidinone—is bacteriostatic against staphylococci and offers the advantage of comparable bioavailability after oral or parenteral administration. Cross-resistance with other inhibitors of protein synthesis has not been detected. However, resistance to linezolid has been reported. Serious adverse reactions to linezolid include thrombocytopenia, occasional cases of neutropenia, and rare instances of peripheral and optic neuropathy.

Tedizolid, a second oxazolidinone released in 2014, is available as both oral and parenteral preparations. It has enhanced in vitro activity against antibiotic-resistant gram-positive bacteria, including staphylococci. Tedizolid is administered once a day.

Ceftaroline is a fifth-generation cephalosporin with bactericidal activity against MRSA (including strains with reduced susceptibility to vancomycin and daptomycin). It is approved for use in nosocomial pneumonias and for skin and soft tissue infections.

The parenteral streptogramin antibiotic quinupristin/dalfopristin displays bactericidal activity against all staphylococci, including VISA strains. This drug has been used successfully to treat serious MRSA infections. In cases of resistance to erythromycin or clindamycin, quinupristin/dalfopristin is bacteriostatic against staphylococci. There are limited data on the efficacy of either quinupristin/dalfopristin or linezolid for the treatment of infective endocarditis.

Telavancin is a parenteral lipoglycopeptide derivative of vancomycin that is approved for the treatment of complicated skin and soft tissue infections and for nosocomial pneumonia. The drug has two targets: the cell wall and the cell membrane. It remains active against VISA strains. Because of its nephrotoxicity, it should be avoided in patients with renal disease.

Dalbavancin is a long-acting, parenterally administered lipoglycopeptide that has been used to treat skin and soft tissue infections. Because of its long half-life, it can be administered on a weekly basis. There are limited data on its use in the treatment of invasive staphylococcal infections.

TABLE 172-3 ANTIMICROBIAL THERAPY FOR STAPHYLOCOCCAL INFECTIONS^a

Sensitivity/Resistance of Isolate	Drug of Choice	Alternative(s)	Comments
Parenteral Therapy for Serious Infections			
Sensitive to penicillin	Penicillin G (4 mU q4h)	Nafcillin or oxacillin (2 g q4h), cefazolin (2 g q8h), vancomycin (1 g q12h ^b)	Fewer than 5% of isolates are sensitive to penicillin. The clinical microbiology laboratory must verify that the strain is not a β -lactamase producer.
Sensitive to methicillin	Nafcillin or oxacillin (2 g q4h)	Cefazolin (2 g q8h), vancomycin (15–20 mg/kg q8–12h ^b)	Patients with penicillin allergy can be treated with a cephalosporin if the allergy does not involve an anaphylactic or accelerated reaction; desensitization to β -lactams may be indicated in selected cases of serious infection when maximal bactericidal activity is needed (e.g., prosthetic-valve endocarditis ^c). Type A β -lactamase may rapidly hydrolyze cefazolin and reduce its efficacy in endocarditis. Vancomycin is a less effective option.
Resistant to methicillin	Vancomycin (15–20 mg/kg q8–12h ^b), daptomycin (6 mg/kg IV q24h ^{b,d}) for bacteremia, endocarditis, and complicated skin infections	Linezolid (600 mg q12h PO or IV), ceftaroline (600 mg IV q12h)	Sensitivity testing is necessary before an alternative drug is selected. For some serious infections, higher doses of daptomycin have been used. Quinupristin/dalfopristin is bactericidal against methicillin-resistant isolates unless the strain is resistant to erythromycin or clindamycin. The efficacy of adjunctive therapy is not well established in many settings. Both linezolid and quinupristin/dalfopristin have had in vitro activity against most VISA and VRSA strains. See footnote for treatment of prosthetic-valve endocarditis. ^c
Resistant to methicillin with intermediate or complete resistance to vancomycin ^e	Daptomycin (6 mg/kg q24h ^{b,d}) for bacteremia, endocarditis, and complicated skin infections	Same as for methicillin-resistant strains; check antibiotic susceptibilities or Ceftaroline (600 mg IV q12h) Newer agents include tedizolid (200 mg administered once daily either IV or orally) or dalbavancin (two IV doses: 1000 mg followed in 1 week by 500 mg). Both drugs are approved only for the treatment of skin and soft tissue infections.	Same as for methicillin-resistant strains; check antibiotic susceptibilities. Ceftaroline is used either alone or in combination with daptomycin.
Not yet known (i.e., empirical therapy)	Vancomycin (15–20 mg/kg q8–12h ^b), daptomycin (6 mg/kg q24h ^{b,d}) for bacteremia, endocarditis, and complicated skin infections	—	Empirical therapy is given when the susceptibility of the isolate is not known. Vancomycin is recommended for suspected community- or hospital-acquired <i>Staphylococcus aureus</i> infections because of the increased frequency of methicillin-resistant strains in the community.
Oral Therapy for Skin and Soft Tissue Infections			
Sensitive to methicillin	Dicloxacillin (500 mg qid), cephalexin (500 mg qid)	Minocycline or doxycycline (100 mg q12h ^b), TMP-SMX (1 or 2 ds tablets bid), clindamycin (300–450 mg/kg tid), linezolid (600 mg PO q12h), tedizolid (200 mg PO q24h)	It is important to know the antibiotic susceptibility of isolates in the specific geographic region. All drainage should be cultured.
Resistant to methicillin	Clindamycin (300–450 mg/kg tid), TMP-SMX (1 or 2 ds tablets bid), minocycline or doxycycline (100 mg q12h ^b), linezolid (600 mg bid) or tedizolid (200 mg once daily)	Same options as under “Drug of Choice”	It is important to know the antibiotic susceptibility of isolates in the specific geographic region. All drainage should be cultured.

^aRecommended dosages are for adults with normal renal and hepatic function. ^bThe dosage must be adjusted for patients with reduced creatinine clearance. ^cFor the treatment of prosthetic-valve endocarditis, the addition of gentamicin (1 mg/kg q8h) and rifampin (300 mg PO q8h) is recommended, with adjustment of the gentamicin dosage if the creatinine clearance rate is reduced. ^dDaptomycin cannot be used for the treatment of pneumonia. ^eVancomycin-resistant *S. aureus* isolates from clinical infections have been reported.

Abbreviations: ds, double-strength; TMP-SMX, trimethoprim-sulfamethoxazole; VISA, vancomycin-intermediate *S. aureus*; VRSA, vancomycin-resistant *S. aureus*.

Source: Modified with permission from FD Lowy: N Engl J Med 339:520, 1998 (© 1998 Massachusetts Medical Society. All rights reserved.); C Liu et al: Clin Infect Dis 52:285, 2011; DL Stevens et al: Clin Infect Dis 59:148, 2014; and Med Lett Drugs Ther 56:39, 2014.

Although the quinolones are active against staphylococci in vitro, the frequency of staphylococcal resistance to these agents has increased progressively, especially among methicillin-resistant isolates. Of particular concern in MRSA is the possible emergence of quinolone resistance during therapy. Therefore, quinolones are not recommended for the treatment of MRSA infections. Resistance to the quinolones is most commonly chromosomal and results from mutations of the topoisomerase IV or DNA gyrase genes, although multidrug efflux pumps may also contribute. Although the newer quinolones exhibit increased in vitro activity against staphylococci, it is uncertain whether this increase translates into enhanced in vivo activity.

Tigecycline, a broad-spectrum minocycline analogue, has bacteriostatic activity against MRSA and is approved for use in skin and soft tissue infections as well as intraabdominal infections caused by *S. aureus*. Other antibiotics, such as minocycline and trimethoprim-sulfamethoxazole, have been used successfully to treat MRSA infections in cases of vancomycin toxicity or intolerance.

Combinations of antistaphylococcal agents have been used to enhance bactericidal activity in the treatment of serious infections such as endocarditis or osteomyelitis. In selected instances (e.g., right-sided endocarditis), drug combinations are also used to shorten the duration of therapy. Among the antimicrobial agents

used in combinations are rifampin, aminoglycosides (e.g., gentamicin), and fusidic acid (not readily available in the United States). To date, clinical studies have not documented a therapeutic benefit; recent reports have raised concern about the potential nephrotoxicity of gentamicin and about adverse drug reactions from the addition of rifampin. As a result, the use of gentamicin in combination with β -lactams or other antimicrobial agents is no longer routinely recommended for the treatment of endocarditis. Rifampin continues to be used for the treatment of prosthetic device–related infections and for osteomyelitis.

The combination of daptomycin with a β -lactam antibiotic has been successfully used to treat patients with persistent MRSA bacteremia, even those infected with isolates that exhibit reduced susceptibility to daptomycin. The combination appears to enhance the bactericidal activity of daptomycin, perhaps by reducing the bacterial cell surface charge and thus allowing more daptomycin binding.

ANTIMICROBIAL THERAPY FOR SELECTED SETTINGS

When necessary, the use of oral antistaphylococcal agents for uncomplicated skin and soft tissue infections is usually successful. For other infections, parenteral therapy is indicated.

S. aureus endocarditis is usually an acute, life-threatening infection. Thus, prompt collection of blood for cultures must be followed immediately by empirical antimicrobial therapy. For life-threatening *S. aureus* native-valve endocarditis, therapy with a β -lactam is recommended. If a MRSA strain is isolated, vancomycin (15–20 mg/kg every 8–12 h, given in equal doses up to a total of 2 g) or daptomycin (6 mg/kg every 24 h) is recommended. The vancomycin dose should be adjusted on the basis of trough vancomycin levels. Patients are generally treated for 4–6 weeks, with duration depending on whether there are complications. For prosthetic-valve endocarditis, surgery in addition to antibiotic therapy is often necessary. The combination of a β -lactam agent—or, if the isolate is β -lactam-resistant, vancomycin (30 mg/kg every 24 h, given in doses up to a total of 2 g) or daptomycin (6 mg/kg every 24 h)—with an aminoglycoside (gentamicin, 1 mg/kg IV every 8 h) and rifampin (300 mg orally or IV every 8 h) for ≥ 6 weeks is recommended.

For hematogenous osteomyelitis or septic arthritis in children, a 4-week course of therapy is usually adequate. In adults, treatment is often more prolonged. For chronic forms of osteomyelitis, surgical debridement is necessary in combination with antimicrobial therapy. For joint infections, a critical component of therapy is the repeated aspiration or arthroscopy of the affected joint to prevent damage from leukocytes. The combination of rifampin with ciprofloxacin has been used successfully to treat prosthetic joint infections, especially when the device cannot be removed. The efficacy of this combination may reflect enhanced activity against staphylococci in biofilms as well as the attainment of effective intracellular concentrations.

The choice of empirical therapy for staphylococcal infections depends in part on susceptibility data for the local geographic area. Increasingly, vancomycin and daptomycin are the drugs of choice for both community- and hospital-acquired infections. The increase in CA-MRSA skin and soft tissue infections has drawn attention to the need for initiation of appropriate empirical therapy. Oral agents that have been effective against these isolates include clindamycin, trimethoprim-sulfamethoxazole, doxycycline, linezolid, and tedizolid.

THERAPY FOR TOXIC SHOCK SYNDROME

Supportive therapy with reversal of hypotension is the mainstay of therapy for TSS. Both fluids and pressors may be necessary. Tampons or other packing material should be promptly removed. The role of antibiotics is less clear. Some investigators recommend a combination of clindamycin and a semisynthetic penicillin or vancomycin (if the isolate is resistant to methicillin). Clindamycin is advocated because, as a protein synthesis inhibitor, it reduces toxin synthesis *in vitro*. Linezolid also appears to be effective. A semisynthetic penicillin or glycopeptide is suggested to eliminate any potential focus of infection as well as to eradicate persistent carriage that might increase the likelihood of recurrent illness.

Anecdotal reports document the successful use of IV immunoglobulin to treat TSS. The role of glucocorticoids in the treatment of this disease is uncertain.

THERAPY FOR OTHER TOXIN-MEDIATED DISEASES

Therapy for staphylococcal food poisoning is entirely supportive. For SSSS, antistaphylococcal therapy targets the primary site of infection.

173 Streptococcal Infections

Michael R. Wessels

Many varieties of streptococci are found as part of the normal flora colonizing the human respiratory, gastrointestinal, and genitourinary tracts. Several species are important causes of human disease. Group A *Streptococcus* (GAS, *Streptococcus pyogenes*) is responsible for streptococcal pharyngitis, one of the most common bacterial infections of school-age children, and for the postinfectious syndromes of acute rheumatic fever (ARF) and poststreptococcal glomerulonephritis (PSGN). Group B *Streptococcus* (GBS, *Streptococcus agalactiae*) is the leading cause of bacterial sepsis and meningitis in newborns and a major cause of endometritis and fever in parturient women. Viridans streptococci are the most common cause of bacterial endocarditis. Enterococci, which are morphologically similar to streptococci, are now considered a separate genus on the basis of DNA homology studies. Thus, the species previously designated as *Streptococcus faecalis* and *Streptococcus faecium* have been renamed *Enterococcus faecalis* and *Enterococcus faecium*, respectively. The enterococci are discussed in [Chap. 174](#).


Streptococci are gram-positive, spherical to ovoid bacteria that characteristically form chains when grown in liquid media. Most streptococci that cause human infections are facultative anaerobes, although some are strict anaerobes. Streptococci are relatively fastidious organisms, requiring enriched media for growth in the laboratory. Clinicians and clinical microbiologists identify streptococci by several classification systems, including hemolytic pattern, Lancefield group, species name, and common or trivial name. Many streptococci associated with human infection produce a zone of complete (β) hemolysis around the bacterial colony when cultured on blood agar. The β -hemolytic streptococci can be classified by the Lancefield system, a serologic grouping based on the reaction of specific antisera with bacterial cell-wall carbohydrate antigens. With rare exceptions, organisms belonging to Lancefield groups A, B, C, and G are all β -hemolytic, and each is associated with characteristic patterns of human infection. Other streptococci produce a zone of partial (α) hemolysis, often imparting a greenish appearance to the agar. These α -hemolytic streptococci are further identified by biochemical testing and include *Streptococcus pneumoniae* ([Chap. 171](#)), an important cause of pneumonia, meningitis, and other infections, and the several species referred to collectively as the *viridans streptococci*, which are part of the normal oral flora and are important agents of subacute bacterial endocarditis. Finally, some streptococci are nonhemolytic, a pattern sometimes called γ hemolysis. Among the organisms classified serologically as group D streptococci, the enterococci are classified as a distinct genus ([Chap. 174](#)). The classification of the major streptococcal groups causing human infections is outlined in [Table 173-1](#).

GROUP A STREPTOCOCCI

Lancefield's group A consists of a single species, *S. pyogenes*. As its species name implies, this organism is associated with a variety of suppurative infections. In addition, GAS can trigger the postinfectious syndromes of ARF (which is uniquely associated with *S. pyogenes* infection; [Chap. 381](#)) and PSGN ([Chap. 338](#)).

Lancefield Group	Representative Species	Hemolytic Pattern	Typical Infections
A	<i>S. pyogenes</i>	β	Pharyngitis, impetigo, cellulitis, scarlet fever
B	<i>S. agalactiae</i>	β	Neonatal sepsis and meningitis, puerperal infection, urinary tract infection, diabetic ulcer infection, endocarditis
C, G	<i>S. dysgalactiae</i> subsp. <i>equisimilis</i>	β	Cellulitis, bacteremia, endocarditis
D	Enterococci ^a : <i>E. faecalis</i> , <i>E. faecium</i> Nonenterococci: <i>S. gallolyticus</i> (formerly <i>S. bovis</i>)	Usually nonhemolytic Usually nonhemolytic	Urinary tract infection, nosocomial bacteremia, endocarditis Bacteremia, endocarditis
Variable or nongroupable	Viridans streptococci: <i>S. sanguis</i> , <i>S. mitis</i>	α	Endocarditis, dental abscess, brain abscess
	<i>Intermedius</i> or <i>milleri</i> group: <i>S. intermedius</i> , <i>S. anginosus</i> , <i>S. constellatus</i>	Variable	Brain abscess, visceral abscess
	Anaerobic streptococci ^b : <i>Peptostreptococcus magnus</i>	Usually nonhemolytic	Sinusitis, pneumonia, empyema, brain abscess, liver abscess

^aSee Chap. 174. ^bSee Chap. 201.

 Worldwide, GAS infections and their postinfectious sequelae (primarily ARF and rheumatic heart disease) account for an estimated 500,000 deaths per year. Although data are incomplete, the incidence of all forms of GAS infection and that of rheumatic heart disease are thought to be tenfold higher in resource-limited countries than in developed countries (Fig. 173-1).

PATHOGENESIS

GAS elaborates a number of cell-surface components and extracellular products important in both the pathogenesis of infection and the human immune response. The cell wall contains a carbohydrate antigen that may be released by acid treatment. The reaction of such acid extracts with group A-specific antiserum is the basis for definitive identification of a streptococcal strain as *S. pyogenes*. The major surface protein of GAS is M protein, which occurs in more than 100 antigenically distinct types and is the basis for the serotyping of strains with specific antisera. The M protein molecules are fibrillar structures anchored in the cell wall of the organism that extend as hairlike projections away from the cell surface. The amino acid sequence of the distal or amino-terminal portion of the M protein molecule is quite variable, accounting for the antigenic variation of the different M types, while more proximal regions of the protein are relatively conserved. A newer technique for assignment of M type to GAS isolates uses the polymerase chain reaction to amplify the variable region of

the *emm* gene, which encodes M protein. DNA sequence analysis of the amplified gene segment can be compared with an extensive database (developed at the Centers for Disease Control and Prevention [CDC]) for assignment of *emm* type. This method eliminates the need for typing sera, which are available in only a few reference laboratories. The presence of M protein on a GAS isolate correlates with its capacity to resist phagocytic killing in fresh human blood. This phenomenon appears to be due, at least in part, to the binding of plasma fibrinogen to M protein molecules on the streptococcal surface, which interferes with complement activation and deposition of opsonic complement fragments on the bacterial cell. This resistance to phagocytosis may be overcome by M protein-specific antibodies; thus individuals with antibodies to a given M type acquired as a result of prior infection are protected against subsequent infection with organisms of the same M type but not against that with different M types.

GAS also elaborates, to varying degrees, a polysaccharide capsule composed of hyaluronic acid. The production of large amounts of capsule by certain strains imparts a characteristic mucoid appearance to the colonies. The capsular polysaccharide plays an important role in protecting GAS from ingestion and killing by phagocytes. In contrast to M protein, the hyaluronic acid capsule is a weak immunogen, and antibodies to hyaluronate have not been shown to be important in protective immunity. The presumed explanation is the apparent structural identity between streptococcal hyaluronic acid and the hyaluronic

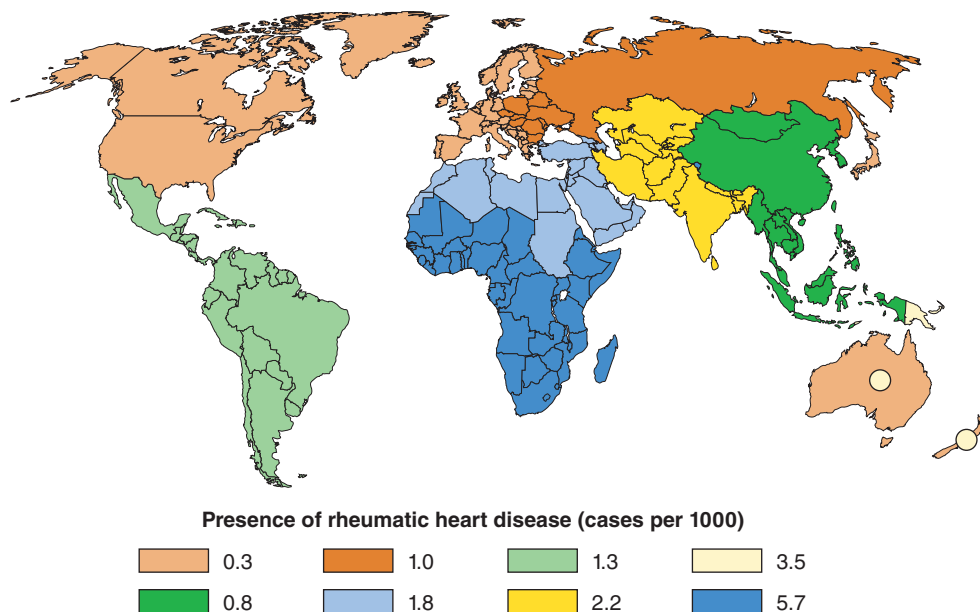


FIGURE 173-1 Prevalence of rheumatic heart disease in children 5–14 years old. The circles within Australia and New Zealand represent indigenous populations (and also Pacific Islanders in New Zealand). (From JR Carapetis et al: *Lancet Infect Dis* 5:685, 2005, with permission.)

acid of mammalian connective tissues. The capsular polysaccharide may also play a role in GAS colonization of the pharynx by binding to CD44, a hyaluronic acid-binding protein expressed on human pharyngeal epithelial cells.

GAS produces a large number of extracellular products that may be important in local and systemic toxicity and in the spread of infection through tissues. These products include streptolysins S and O, toxins that damage cell membranes and account for the hemolysis produced by the organisms; streptokinase; DNases; SpyCEP, a serine protease that cleaves and inactivates the chemoattractant cytokine interleukin 8, thereby inhibiting neutrophil recruitment to the site of infection; and several pyrogenic exotoxins. Previously known as erythrogenic toxins, the pyrogenic exotoxins cause the rash of scarlet fever. Since the mid-1980s, pyrogenic exotoxin-producing strains of GAS have been linked to unusually severe invasive infections, including necrotizing fasciitis and the streptococcal toxic shock syndrome (TSS). Several extracellular products stimulate specific antibody responses useful for serodiagnosis of recent streptococcal infection. Tests for these antibodies are used primarily for detection of preceding streptococcal infection in cases of suspected ARF or PSGN.

CLINICAL MANIFESTATIONS

Pharyngitis Although seen in patients of all ages, GAS pharyngitis is one of the most common bacterial infections of childhood, accounting for 20–40% of all cases of exudative pharyngitis in children; it is rare among those under the age of 3. Younger children may manifest streptococcal infection with a syndrome of fever, malaise, and lymphadenopathy without exudative pharyngitis. Infection is acquired through contact with another individual carrying the organism. Respiratory droplets are the usual mechanism of spread, although other routes, including food-borne outbreaks, have been well described. The incubation period is 1–4 days. Symptoms include sore throat, fever and chills, malaise, and sometimes abdominal complaints and vomiting, particularly in children. Both symptoms and signs are quite variable, ranging from mild throat discomfort with minimal physical findings to high fever and severe sore throat associated with intense erythema and swelling of the pharyngeal mucosa and the presence of purulent exudate over the posterior pharyngeal wall and tonsillar pillars. Enlarged, tender anterior cervical lymph nodes commonly accompany exudative pharyngitis.

The differential diagnosis of streptococcal pharyngitis includes the many other bacterial and viral etiologies (Table 173-2). Streptococcal infection is an unlikely cause when symptoms and signs suggestive of viral infection are prominent (conjunctivitis, coryza, cough, hoarseness, or discrete ulcerative lesions of the buccal or pharyngeal mucosa). Because of the range of clinical presentations of streptococcal pharyngitis and the large number of other agents that can produce the same clinical picture, diagnosis of streptococcal pharyngitis on clinical grounds alone is not reliable. The throat culture remains the diagnostic gold standard. Culture of a throat specimen that is properly collected (i.e., by vigorous rubbing of a sterile swab over both tonsillar pillars) and properly processed is the most sensitive and specific means of definitive diagnosis. A rapid diagnostic kit for latex agglutination or enzyme immunoassay of swab specimens is a useful adjunct to throat culture. While precise figures on sensitivity and specificity vary, rapid diagnostic kits generally are >95% specific. Thus a positive result can be relied upon for definitive diagnosis and eliminates the need for throat culture. However, because rapid diagnostic tests are less sensitive than throat culture (relative sensitivity in comparative studies, 55–90%), a negative result should be confirmed by throat culture.

TREATMENT GAS PHARYNGITIS

In the usual course of uncomplicated streptococcal pharyngitis, symptoms resolve after 3–5 days. The course is shortened little by treatment, which is given primarily to prevent suppurative complications and ARF. Prevention of ARF depends on eradication of the organism from the pharynx, not simply on resolution of symptoms,

TABLE 173-2 INFECTIOUS ETIOLOGIES OF ACUTE PHARYNGITIS

Organism	Associated Clinical Syndrome(s)
Viruses	
Rhinovirus	Common cold
Coronavirus	Common cold
Adenovirus	Pharyngoconjunctival fever
Influenza virus	Influenza
Parainfluenza virus	Cold, croup
Coxsackievirus	Herpangina, hand-foot-and-mouth disease
Herpes simplex virus	Gingivostomatitis (primary infection)
Epstein-Barr virus	Infectious mononucleosis
Cytomegalovirus	Mononucleosis-like syndrome
HIV	Acute (primary) infection syndrome
Bacteria	
Group A streptococci	Pharyngitis, scarlet fever
Group C or G streptococci	Pharyngitis
Mixed anaerobes	Vincent's angina
<i>Arcanobacterium haemolyticum</i>	Pharyngitis, scarlatiniform rash
<i>Neisseria gonorrhoeae</i>	Pharyngitis
<i>Treponema pallidum</i>	Secondary syphilis
<i>Francisella tularensis</i>	Pharyngeal tularemia
<i>Corynebacterium diphtheriae</i>	Diphtheria
<i>Yersinia enterocolitica</i>	Pharyngitis, enterocolitis
<i>Yersinia pestis</i>	Plague
Chlamydiae	
<i>Chlamydia pneumoniae</i>	Bronchitis, pneumonia
<i>Chlamydia psittaci</i>	Psittacosis
Mycoplasmas	
<i>Mycoplasma pneumoniae</i>	Bronchitis, pneumonia

and requires 10 days of penicillin treatment (Table 173-3). A first-generation cephalosporin, such as cephalexin or cefadroxil, may be substituted for penicillin in cases of penicillin allergy if the nature of the allergy is not an immediate hypersensitivity reaction (anaphylaxis or urticaria) or another potentially life-threatening manifestation (e.g., severe rash and fever).

TABLE 173-3 TREATMENT OF GROUP A STREPTOCOCCAL INFECTIONS

Infection	Treatment ^a
Pharyngitis	Benzathine penicillin G (1.2 mU IM) or penicillin V (250 mg PO tid or 500 mg PO bid) × 10 days (Children <27 kg: Benzathine penicillin G [600,000 units IM] or penicillin V [250 mg PO bid or tid] × 10 days)
Impetigo	Same as pharyngitis
Erysipelas/cellulitis	Severe: Penicillin G (1–2 mU IV q4h) Mild to moderate: Procaine penicillin (1.2 mU IM bid)
Necrotizing fasciitis/myositis	Surgical debridement plus penicillin G (2–4 mU IV q4h) plus clindamycin ^b (600–900 mg IV q8h)
Pneumonia/empyema	Penicillin G (2–4 mU IV q4h) plus drainage of empyema
Streptococcal toxic shock syndrome	Penicillin G (2–4 mU IV q4h) plus clindamycin ^b (600–900 mg IV q8h) plus IV immunoglobulin ^b (2 g/kg as a single dose)

^aPenicillin allergy: A first-generation cephalosporin, such as cephalexin or cefadroxil, may be substituted for penicillin in cases of penicillin allergy if the nature of the allergy is not an immediate hypersensitivity reaction (anaphylaxis or urticaria) or another potentially life-threatening manifestation (e.g., severe rash and fever). Alternative agents for oral therapy are erythromycin (10 mg/kg PO qid, up to a maximum of 250 mg per dose) and azithromycin (a 5-day course at a dose of 12 mg/kg once daily, up to a maximum of 500 mg/d). Vancomycin is an alternative for parenteral therapy. ^bEfficacy unproven, but recommended by several experts. See text for discussion.



Alternative agents are erythromycin and azithromycin. Azithromycin is more expensive but offers the advantages of better gastrointestinal tolerability, once-daily dosing, and a 5-day treatment course. Resistance to erythromycin and other macrolides is common among isolates from several countries, including Spain, Italy, Finland, Japan, and Korea. Macrolide resistance may be becoming more prevalent elsewhere with the increasing use of this class of antibiotics. In areas with resistance rates exceeding 5–10%, macrolides should be avoided unless results of susceptibility testing are known.

Follow-up culture after treatment is no longer routinely recommended but may be warranted in selected cases, such as those involving patients or families with frequent streptococcal infections or those occurring in situations in which the risk of ARF is thought to be high (e.g., when cases of ARF have recently been reported in the community).

COMPLICATIONS Suppurative complications of streptococcal pharyngitis have become uncommon with the widespread use of antibiotics for most symptomatic cases. These complications result from the spread of infection from the pharyngeal mucosa to deeper tissues by direct extension or by the hematogenous or lymphatic route and may include cervical lymphadenitis, peritonsillar or retropharyngeal abscess, sinusitis, otitis media, meningitis, bacteremia, endocarditis, and pneumonia. Local complications, such as peritonsillar or parapharyngeal abscess formation, should be considered in a patient with unusually severe or prolonged symptoms or localized pain associated with high fever and a toxic appearance. Nonsuppurative complications include ARF (**Chap. 381**) and PSGN (**Chap. 338**), both of which are thought to result from immune responses to streptococcal infection. Penicillin treatment of streptococcal pharyngitis has been shown to reduce the likelihood of ARF but not that of PSGN.

BACTERIOLOGIC TREATMENT FAILURE AND THE ASYMPTOMATIC CARRIER STATE Surveillance cultures have shown that up to 20% of individuals in certain populations may have asymptomatic pharyngeal colonization with GAS. There are no definitive guidelines for management of these asymptomatic carriers or of asymptomatic patients who still have a positive throat culture after a full course of treatment for symptomatic pharyngitis. A reasonable course of action is to give a single 10-day course of penicillin for symptomatic pharyngitis and, if positive cultures persist, not to re-treat unless symptoms recur. Studies of the natural history of streptococcal carriage and infection have shown that the risk both of developing ARF and of transmitting infection to others is substantially lower among asymptomatic carriers than among individuals with symptomatic pharyngitis. Therefore, overly aggressive attempts to eradicate carriage probably are not justified under most circumstances. An exception is the situation in which an asymptomatic carrier is a potential source of infection to others. Outbreaks of food-borne infection and nosocomial puerperal infection have been traced to asymptomatic carriers who may harbor the organisms in the throat, vagina, or anus or on the skin.

TREATMENT ASYMPTOMATIC PHARYNGEAL COLONIZATION WITH GAS

When a carrier is transmitting infection to others, attempts to eradicate carriage are warranted. Data are limited on the best regimen to clear GAS after penicillin alone has failed. Regimens reported to have efficacy superior to that of penicillin alone for eradication of carriage include (1) oral clindamycin (7 mg/kg; 300 mg maximum) three times daily for 10 days or (2) penicillin (as recommended for treatment of pharyngitis in Table 173-3) plus oral rifampin (10 mg/kg; 300 mg maximum) twice daily for the first 4 days of treatment. A 10-day course of oral vancomycin (250 mg four times daily) and rifampin (600 mg twice daily) has eradicated rectal colonization.

Scarlet Fever Scarlet fever consists of streptococcal infection, usually pharyngitis, accompanied by a characteristic rash (**Fig. 173-2**).



FIGURE 173-2 Scarlet fever exanthem. Finely punctate erythema has become confluent (scarlatiniform); petechiae can occur and have a linear configuration within the exanthem in body folds (Pastia's lines). (From Fitzpatrick, Johnson, Wolff: *Color Atlas and Synopsis of Clinical Dermatology*, 4th ed, New York, McGraw-Hill, 2001, with permission.)

The rash arises from the effects of one of several toxins, currently designated *streptococcal pyrogenic exotoxins* and previously known as *erythrogenic* or *scarlet fever toxins*. In the past, scarlet fever was thought to reflect infection of an individual lacking toxin-specific immunity with a toxin-producing strain of GAS. Susceptibility to scarlet fever was correlated with results of the Dick test, in which a small amount of erythrogenic toxin injected intradermally produced local erythema in susceptible individuals but elicited no reaction in those with specific immunity. Subsequent studies have suggested that development of the scarlet fever rash may reflect a hypersensitivity reaction requiring prior exposure to the toxin. For reasons that are not clear, scarlet fever has become less common in recent years, although strains of GAS that produce pyrogenic exotoxins continue to be prevalent in the population. The symptoms of scarlet fever are the same as those of pharyngitis alone. The rash typically begins on the first or second day of illness over the upper trunk, spreading to involve the extremities but sparing the palms and soles. The rash is made up of minute papules, giving a characteristic “sandpaper” feel to the skin. Associated findings include circumoral pallor, “strawberry tongue” (enlarged papillae on a coated tongue, which later may become denuded), and accentuation of the rash in skinfolds (*Pastia's lines*). Subsidence of the rash in 6–9 days is followed after several days by desquamation of the palms and soles. The differential diagnosis of scarlet fever includes other causes of fever and generalized rash, such as measles and other viral exanthems, Kawasaki disease, TSS, and systemic allergic reactions (e.g., drug eruptions).

Skin and Soft Tissue Infections GAS—and occasionally other streptococcal species—can cause a variety of infections involving the skin, subcutaneous tissues, muscles, and fascia. While several clinical syndromes offer a useful means for classification of these infections, not all cases fit exactly into one category. The classic syndromes are general guides to predicting the level of tissue involvement in a particular patient, the probable clinical course, and the likelihood that surgical intervention or aggressive life support will be required.

IMPETIGO (PYODERMA) Impetigo, a superficial infection of the skin, is caused primarily by GAS and occasionally by other streptococci or *Staphylococcus aureus*. Impetigo is seen most often in young children, tends to occur during warmer months, and is more common in semi-tropical or tropical climates than in cooler regions. Infection is more



FIGURE 173-3 Impetigo contagiosa is a superficial streptococcal or *Staphylococcus aureus* infection consisting of honey-colored crusts and erythematous weeping erosions. Occasionally, bullous lesions may be seen. (Courtesy of Mary Spraker, MD; with permission.)

common among children living under conditions of poor hygiene. Prospective studies have shown that colonization of unbroken skin with GAS precedes clinical infection. Minor trauma, such as a scratch or an insect bite, may then serve to inoculate organisms into the skin. Impetigo is best prevented, therefore, by attention to adequate hygiene. The usual sites of involvement are the face (particularly around the nose and mouth) and the legs, although lesions may occur at other locations. Individual lesions begin as red papules, which evolve quickly into vesicular and then pustular lesions that break down and coalesce to form characteristic honeycomb-like crusts (Fig. 173-3). Lesions generally are not painful, and patients do not appear ill. Fever is not a feature of impetigo and, if present, suggests either infection extending to deeper tissues or another diagnosis. The classic presentation of impetigo usually poses little diagnostic difficulty. Cultures of impetiginous lesions often yield *S. aureus* as well as GAS. In almost all cases, streptococci are isolated initially and staphylococci appear later, presumably as secondary colonizing flora. In the past, penicillin was nearly always effective against these infections. However, an increasing frequency of penicillin treatment failure suggests that *S. aureus* may have become more prominent as a cause of impetigo. Bullous impetigo due to *S. aureus* is distinguished from typical streptococcal infection by more extensive, bullous lesions that break down and leave thin paper-like crusts instead of the thick amber crusts of streptococcal impetigo. Other skin lesions that may be confused with impetigo include herpetic lesions—either those of orolabial herpes simplex or those of chickenpox or zoster. Herpetic lesions can generally be distinguished by their appearance as more discrete, grouped vesicles and by a positive Tzanck test. In difficult cases, cultures of vesicular fluid should yield GAS in impetigo and the responsible virus in herpesvirus infections.

TREATMENT STREPTOCOCCAL IMPETIGO

Treatment of streptococcal impetigo is the same as that for streptococcal pharyngitis. In view of evidence that *S. aureus* has become a relatively frequent cause of impetigo, empirical regimens should cover both streptococci and *S. aureus*. For example, either dicloxacillin or cephalexin can be given at a dose of 250 mg four times daily for 10 days. Topical mupirocin ointment is also effective. Culture may be indicated to rule out methicillin-resistant *S. aureus*, especially if the response to empirical treatment is unsatisfactory. ARF is not a sequela to streptococcal skin infections, although PSGN may follow either skin or throat infection. The reason for this difference is

not known. One hypothesis is that the immune response necessary for development of ARF occurs only after infection of the pharyngeal mucosa. In addition, the strains of GAS that cause pharyngitis are generally of different M protein types than those associated with skin infections; thus the strains that cause pharyngitis may have rheumatogenic potential, while the skin-infecting strains may not.

CELLULITIS Inoculation of organisms into the skin may lead to *cellulitis*: infection involving the skin and subcutaneous tissues. The portal of entry may be a traumatic or surgical wound, an insect bite, or any other break in skin integrity. Often, no entry site is apparent. One form of streptococcal cellulitis, *erysipelas*, is characterized by a bright red appearance of the involved skin, which forms a plateau sharply demarcated from surrounding normal skin (Fig. 173-4). The lesion is warm to the touch, may be tender, and appears shiny and swollen. The skin often has a *peau d'orange* texture, which is thought to reflect involvement of superficial lymphatics; superficial blebs or bullae may form, usually 2–3 days after onset. The lesion typically develops over a few hours and is associated with fever and chills. Erysipelas tends to occur on the malar area of the face (often with extension over the bridge of the nose to the contralateral malar region) and the lower extremities. After one episode, recurrence at the same site—sometimes years later—is not uncommon. Classic cases of erysipelas, with typical features, are almost always due to β -hemolytic streptococci, usually GAS and occasionally group C or G. Often, however, the appearance of streptococcal cellulitis is not sufficiently distinctive to permit a specific diagnosis on clinical grounds. The area involved may not be typical for erysipelas, the lesion may be less intensely red than usual and may fade into surrounding skin, and/or the patient may appear only mildly ill. In such cases, it is prudent to broaden the spectrum of empirical antimicrobial therapy to include other pathogens, particularly *S. aureus*, that can produce cellulitis with the same appearance. Staphylococcal infection should be suspected if cellulitis develops around a wound or an ulcer.

Streptococcal cellulitis tends to develop at anatomic sites in which normal lymphatic drainage has been disrupted, such as sites of prior cellulitis, the arm ipsilateral to a mastectomy and axillary lymph node dissection, a lower extremity previously involved in deep venous thrombosis or chronic lymphedema, or the leg from which a saphenous vein has been harvested for coronary artery bypass grafting. The organism may enter via a dermal breach some distance from the eventual site of clinical cellulitis. For example, some patients with recurrent leg cellulitis following saphenous vein removal stop having recurrent episodes only after treatment of tinea pedis on the affected



FIGURE 173-4 Erysipelas is a streptococcal infection of the superficial dermis and consists of well-demarcated, erythematous, edematous, warm plaques.

968 extremity. Fissures in the skin presumably serve as a portal of entry for streptococci, which then produce infection more proximally in the leg at the site of previous injury. Streptococcal cellulitis may also involve recent surgical wounds. GAS is among the few bacterial pathogens that typically produce signs of wound infection and surrounding cellulitis within the first 24 h after surgery. These wound infections are usually associated with a thin exudate and may spread rapidly, either as cellulitis in the skin and subcutaneous tissue or as a deeper tissue infection (see below). Streptococcal wound infection or localized cellulitis may also be associated with *lymphangitis*, manifested by red streaks extending proximally along superficial lymphatics from the infection site.

TREATMENT STREPTOCOCCAL CELLULITIS

See Table 173-3 and Chap. 156.

DEEP SOFT-TISSUE INFECTIONS *Necrotizing fasciitis* (*hemolytic streptococcal gangrene*) involves the superficial and/or deep fascia investing the muscles of an extremity or the trunk. The source of the infection is either the skin, with organisms introduced into tissue through trauma (sometimes trivial), or the bowel flora, with organisms released during abdominal surgery or from an occult enteric source, such as a diverticular or appendiceal abscess. The inoculation site may be inapparent and is often some distance from the site of clinical involvement; e.g., the introduction of organisms via minor trauma to the hand may be associated with clinical infection of the tissues overlying the shoulder or chest. Cases associated with the bowel flora are usually polymicrobial, involving a mixture of anaerobic bacteria (such as *Bacteroides fragilis* or anaerobic streptococci) and facultative organisms (usually gram-negative bacilli). Cases unrelated to contamination from bowel organisms are most commonly caused by GAS alone or in combination with other organisms (most often *S. aureus*). Overall, GAS is implicated in ~60% of cases of necrotizing fasciitis. The onset of symptoms is usually quite acute and is marked by severe pain at the site of involvement, malaise, fever, chills, and a toxic appearance. The physical findings, particularly early on, may not be striking, with only minimal erythema of the overlying skin. Pain and tenderness are usually severe. In contrast, in more superficial cellulitis, the skin appearance is more abnormal, but pain and tenderness are only mild or moderate. As the infection progresses (often over several hours), the severity and extent of symptoms worsen, and skin changes become more evident, with the appearance of dusky or mottled erythema and edema. The marked tenderness of the involved area may evolve into anesthesia as the spreading inflammatory process produces infarction of cutaneous nerves.

Although myositis is more commonly due to *S. aureus* infection, GAS occasionally produces abscesses in skeletal muscles (*streptococcal myositis*), with little or no involvement of the surrounding fascia or overlying skin. The presentation is usually subacute, but a fulminant form has been described in association with severe systemic toxicity, bacteremia, and a high mortality rate. The fulminant form may reflect the same basic disease process seen in necrotizing fasciitis, but with the necrotizing inflammatory process extending into the muscles themselves rather than remaining limited to the fascial layers.

TREATMENT DEEP SOFT-TISSUE INFECTIONS

Once necrotizing fasciitis is suspected, early surgical exploration is both diagnostically and therapeutically indicated. Surgery reveals necrosis and inflammatory fluid tracking along the fascial planes above and between muscle groups, without involvement of the muscles themselves. The process usually extends beyond the area of clinical involvement, and extensive debridement is required. Drainage and debridement are central to the management of necrotizing fasciitis; antibiotic treatment is a useful adjunct (Table 173-3), but surgery is life-saving. Treatment for streptococcal myositis consists of surgical drainage—usually by an open procedure that permits evaluation of the extent of infection and ensures

adequate debridement of involved tissues—and high-dose penicillin (Table 173-3).

Pneumonia and Empyema GAS is an occasional cause of pneumonia, generally in previously healthy individuals. The onset of symptoms may be abrupt or gradual. Pleuritic chest pain, fever, chills, and dyspnea are the characteristic manifestations. Cough is usually present but may not be prominent. Approximately one-half of patients with GAS pneumonia have an accompanying pleural effusion. In contrast to the sterile parapneumonic effusions typical of pneumococcal pneumonia, those complicating streptococcal pneumonia are almost always infected. The empyema fluid is usually visible by chest radiography on initial presentation, and its volume may increase rapidly. These pleural collections should be drained early, as they tend to become loculated rapidly, resulting in a chronic fibrotic reaction that may require thoracotomy for removal.

Bacteremia, Puerperal Sepsis, and Streptococcal Toxic Shock Syndrome GAS bacteremia is usually associated with an identifiable local infection. Bacteremia occurs rarely with otherwise uncomplicated pharyngitis, occasionally with cellulitis or pneumonia, and relatively frequently with necrotizing fasciitis. Bacteremia without an identified source raises the possibility of endocarditis, an occult abscess, or osteomyelitis. A variety of focal infections may arise secondarily from streptococcal bacteremia, including endocarditis, meningitis, septic arthritis, osteomyelitis, peritonitis, and visceral abscesses. GAS is occasionally implicated in infectious complications of childbirth, usually endometritis and associated bacteremia. In the preantibiotic era, puerperal sepsis was commonly caused by GAS; currently, it is more often caused by GBS. Several nosocomial outbreaks of puerperal GAS infection have been traced to an asymptomatic carrier, usually someone present at delivery. The site of carriage may be the skin, throat, anus, or vagina.

Beginning in the late 1980s, several reports described patients with GAS infections associated with shock and multisystem organ failure. This syndrome was called *streptococcal TSS* because it shares certain features with staphylococcal TSS. In 1993, a case definition for streptococcal TSS was formulated (Table 173-4). The general features of the illness include fever, hypotension, renal impairment, and respiratory distress syndrome. Various types of rash have been described, but rash usually does not develop. Laboratory abnormalities include a marked shift to the left in the white blood cell differential, with many immature granulocytes; hypocalcemia; hypoalbuminemia; and thrombocytopenia, which usually becomes more pronounced on the second or third day of illness. In contrast to patients with staphylococcal TSS, the majority with streptococcal TSS are bacteremic. The most common associated infection is a soft tissue infection—necrotizing

TABLE 173-4 PROPOSED CASE DEFINITION FOR THE STREPTOCOCCAL TOXIC SHOCK SYNDROME^a

- I. Isolation of group A streptococci (*Streptococcus pyogenes*)
 - A. From a normally sterile site
 - B. From a nonsterile site
- II. Clinical signs of severity
 - A. Hypotension *and*
 - B. ≥2 of the following signs
 1. Renal impairment
 2. Coagulopathy
 3. Liver function impairment
 4. Adult respiratory distress syndrome
 5. A generalized erythematous macular rash that may desquamate
 6. Soft tissue necrosis, including necrotizing fasciitis or myositis; or gangrene

^aAn illness fulfilling criteria IA, IIA, and IIB is defined as a *definite* case. An illness fulfilling criteria IB, IIA, and IIB is defined as a *probable* case if no other etiology for the illness is identified.

Source: Modified from Working Group on Severe Streptococcal Infections: JAMA 269:390, 1993.

fasciitis, myositis, or cellulitis—although a variety of other associated local infections have been described, including pneumonia, peritonitis, osteomyelitis, and myometritis. Streptococcal TSS is associated with a mortality rate of $\geq 30\%$, with most deaths secondary to shock and respiratory failure. Because of its rapidly progressive and lethal course, early recognition of the syndrome is essential. Patients should receive aggressive supportive care (fluid resuscitation, pressors, and mechanical ventilation) in addition to antimicrobial therapy and, in cases associated with necrotizing fasciitis, surgical debridement. Exactly why certain patients develop this fulminant syndrome is not known. Early studies of the streptococcal strains isolated from these patients demonstrated a strong association with the production of pyrogenic exotoxin A. This association has been inconsistent in subsequent case series. Pyrogenic exotoxin A and several other streptococcal exotoxins act as superantigens to trigger release of inflammatory cytokines from T lymphocytes. Fever, shock, and organ dysfunction in streptococcal TSS may reflect, in part, the systemic effects of superantigen-mediated cytokine release.

TREATMENT STREPTOCOCCAL TOXIC SHOCK SYNDROME

In light of the possible role of pyrogenic exotoxins or other streptococcal toxins in streptococcal TSS, treatment with clindamycin has been advocated by some authorities (Table 173-3), who argue that, through its direct action on protein synthesis, clindamycin is more effective in rapidly terminating toxin production than is penicillin—a cell-wall agent. Support for this view comes from studies of an experimental model of streptococcal myositis, in which mice given clindamycin had a higher rate of survival than those given penicillin. Comparable data on the treatment of human infections are not available, although retrospective analysis has suggested a better outcome when patients with invasive soft-tissue infection are treated with clindamycin rather than with cell wall-active antibiotics. Although clindamycin resistance in GAS is uncommon ($<2\%$ among U.S. isolates), it has been documented. Thus, if clindamycin is used for initial treatment of a critically ill patient, penicillin should be given as well until the antibiotic susceptibility of the streptococcal isolate is known. IV immunoglobulin has been used as adjunctive therapy for streptococcal TSS (Table 173-3). Pooled immunoglobulin preparations contain antibodies capable of neutralizing the effects of streptococcal toxins. Anecdotal reports and case series have suggested favorable clinical responses to IV immunoglobulin, but no adequately powered, prospective, controlled trials have been reported.

PREVENTION

No vaccine against GAS is commercially available. A formulation that consists of recombinant peptides containing epitopes of 26 M-protein types has undergone phase 1 and 2 testing in volunteers. Early results indicate that the vaccine is well tolerated and elicits type-specific antibody responses. Vaccines based on a conserved region of M protein or on a mixture of other conserved GAS protein antigens are in earlier stages of development.

Household contacts of individuals with invasive GAS infection (e.g., bacteremia, necrotizing fasciitis, or streptococcal TSS) are at greater risk of invasive infection than the general population. Asymptomatic pharyngeal colonization with GAS has been detected in up to 25% of persons with >4 h/d of same-room exposure to an index case. However, antibiotic prophylaxis is not routinely recommended for contacts of patients with invasive disease because such an approach (if effective) would require treatment of hundreds of contacts to prevent a single case.

STREPTOCOCCI OF GROUPS C AND G

Group C and group G streptococci are β -hemolytic bacteria that occasionally cause human infections similar to those caused by GAS. Strains that form small colonies on blood agar (<0.5 mm) are generally members of the *Streptococcus milleri* (*Streptococcus intermedius*,

Streptococcus anginosus) group (see “Viridans Streptococci,” below). Large-colony group C and G streptococci of human origin are now considered a single species, *Streptococcus dysgalactiae* subspecies *equisimilis*. These organisms have been associated with pharyngitis, cellulitis and soft tissue infections, pneumonia, bacteremia, endocarditis, and septic arthritis. Puerperal sepsis, meningitis, epidural abscess, intraabdominal abscess, urinary tract infection, and neonatal sepsis have also been reported. Group C or G streptococcal bacteremia most often affects elderly or chronically ill patients and, in the absence of obvious local infection, is likely to reflect endocarditis. Septic arthritis, sometimes involving multiple joints, may complicate endocarditis or develop in its absence. Distinct streptococcal species of Lancefield group C cause infections in domesticated animals, especially horses and cattle; some human infections are acquired through contact with animals or consumption of unpasteurized milk. These zoonotic organisms include *Streptococcus equi* subspecies *zooepidemicus* and *S. equi* subspecies *equi*.

TREATMENT GROUP C OR G STREPTOCOCCAL INFECTION

Penicillin is the drug of choice for treatment of group C or G streptococcal infections. Antibiotic treatment is the same as for similar syndromes due to GAS (Table 173-3). Patients with bacteremia or septic arthritis should receive IV penicillin (2–4 mU every 4 h). All group C and G streptococci are sensitive to penicillin; nearly all are inhibited in vitro by concentrations of ≤ 0.03 $\mu\text{g}/\text{mL}$. Occasional isolates exhibit tolerance: although inhibited by low concentrations of penicillin, they are killed only by significantly higher concentrations. The clinical significance of tolerance is unknown. Because of the poor clinical response of some patients to penicillin alone, the addition of gentamicin (1 mg/kg every 8 h for patients with normal renal function) is recommended by some authorities for treatment of endocarditis or septic arthritis due to group C or G streptococci; however, combination therapy has not been shown to be superior to penicillin treatment alone. Patients with joint infections often require repeated aspiration or open drainage and debridement for cure; the response to treatment may be slow, particularly in debilitated patients and those with involvement of multiple joints. Infection of prosthetic joints almost always requires prosthesis removal in addition to antibiotic therapy.

GROUP B STREPTOCOCCI

Identified first as a cause of mastitis in cows, streptococci belonging to Lancefield’s group B have since been recognized as a major cause of sepsis and meningitis in human neonates. GBS is also a frequent cause of peripartum fever in women and an occasional cause of serious infection in nonpregnant adults. Since the widespread institution of prenatal screening for GBS in the 1990s, the incidence of neonatal infection per 1000 live births has fallen from ~ 2 –3 cases to ~ 0.6 case. During the same period, GBS infection in adults with underlying chronic illnesses has become more common; adults now account for a larger proportion of invasive GBS infections than do newborns. Lancefield group B consists of a single species, *S. agalactiae*, which is definitively identified with specific antiserum to the group B cell wall-associated carbohydrate antigen. A streptococcal isolate can be classified presumptively as GBS on the basis of biochemical tests, including hydrolysis of sodium hippurate (in which 99% of isolates are positive), hydrolysis of bile esculin (in which 99–100% are negative), bacitracin susceptibility (in which 92% are resistant), and production of CAMP factor (in which 98–100% are positive). CAMP factor is a phospholipase produced by GBS that causes synergistic hemolysis with β lysin produced by certain strains of *S. aureus*. Its presence can be demonstrated by cross-streaking of the test isolate and an appropriate staphylococcal strain on a blood agar plate. GBS organisms causing human infections are encapsulated by one of ten antigenically distinct polysaccharides. The capsular polysaccharide is an important virulence factor. Antibodies to the capsular polysaccharide afford protection against GBS of the same (but not of a different) capsular type.

Two general types of GBS infection in infants are defined by the age of the patient at presentation. *Early-onset infections* occur within the first week of life, with a median age of 20 h at onset. Approximately half of these infants have signs of GBS disease at birth. The infection is acquired during or shortly before birth from the colonized maternal genital tract. Surveillance studies have shown that 5–40% of women are vaginal or rectal carriers of GBS. Approximately 50% of infants delivered vaginally by carrier mothers become colonized, although only 1–2% develop clinically evident infection. Prematurity, prolonged labor, obstetric complications, and maternal fever are risk factors for early-onset infection. The presentation of early-onset infection is the same as that of other forms of neonatal sepsis. Typical findings include respiratory distress, lethargy, and hypotension. Essentially all infants with early-onset disease are bacteremic, one-third to one-half have pneumonia and/or respiratory distress syndrome, and one-third have meningitis.

Late-onset infections occur in infants 1 week to 3 months old and, in rare instances, in older infants (mean age at onset, 3–4 weeks). The infecting organism may be acquired during delivery (as in early-onset cases) or during later contact with a colonized mother, nursery personnel, or another source. Meningitis is the most common manifestation of late-onset infection and in most cases is associated with a strain of capsular type III. Infants present with fever, lethargy or irritability, poor feeding, and seizures. The various other types of late-onset infection include bacteremia without an identified source, osteomyelitis, septic arthritis, and facial cellulitis associated with submandibular or preauricular adenitis.

TREATMENT GROUP B STREPTOCOCCAL INFECTION IN NEONATES

Penicillin is the agent of choice for all GBS infections. Empirical broad-spectrum therapy for suspected bacterial sepsis, consisting of ampicillin and gentamicin, is generally administered until culture results become available. If cultures yield GBS, many pediatricians continue to administer gentamicin, along with ampicillin or penicillin, for a few days until clinical improvement becomes evident. Infants with bacteremia or soft tissue infection should receive penicillin at a dosage of 200,000 units/kg per day in divided doses. For meningitis, infants ≤ 7 days of age should receive 250,000–450,000 units/kg per day in three divided doses; infants > 7 days of age should receive 450,000–500,000 units/kg per day in four divided doses. Meningitis should be treated for at least 14 days because of the risk of relapse with shorter courses.

PREVENTION

The incidence of GBS infection is unusually high among infants of women with risk factors: preterm delivery, early rupture of membranes (> 24 h before delivery), prolonged labor, fever, or chorioamnionitis. Because the usual source of the organisms infecting a neonate is the mother's birth canal, efforts have been made to prevent GBS infections by the identification of high-risk carrier mothers and their treatment with various forms of antibiotic prophylaxis or immunoprophylaxis. Prophylactic administration of ampicillin or penicillin to such patients during delivery reduces the risk of infection in the newborn. This approach has been hampered by logistical difficulties in identifying colonized women before delivery; the results of vaginal cultures early in pregnancy are poor predictors of carrier status at delivery. The CDC recommends screening for anogenital colonization at 35–37 weeks of pregnancy by a swab culture of the lower vagina and anorectum; intrapartum chemoprophylaxis is recommended for culture-positive women and for women who, regardless of culture status, have previously given birth to an infant with GBS infection or have a history of GBS bacteriuria during pregnancy. Women whose culture status is unknown and who develop premature labor (< 37 weeks), prolonged rupture of membranes (> 18 h), or intrapartum fever or who have a

positive intrapartum nucleic acid amplification test for GBS should also receive intrapartum chemoprophylaxis. The recommended regimen for chemoprophylaxis is a loading dose of 5 million units of penicillin G followed by 2.5 million units every 4 h until delivery. Cefazolin is an alternative for women with a history of penicillin allergy who are thought not to be at high risk for anaphylaxis. For women with a history of immediate hypersensitivity, clindamycin may be substituted, but only if the colonizing isolate has been demonstrated to be susceptible. If susceptibility testing results are not available or indicate resistance, vancomycin should be used in this situation.

Treatment of all pregnant women who are colonized or have risk factors for neonatal infection will result in exposure of up to one-third of pregnant women and newborns to antibiotics, with the attendant risks of allergic reactions and selection for resistant organisms. Although still in the developmental stages, a GBS vaccine may ultimately offer a better solution to prevention. Because transplacental passage of maternal antibodies produces protective antibody levels in newborns, efforts are under way to develop a vaccine against GBS that can be given to childbearing-age women before or during pregnancy. Results of phase I clinical trials of GBS capsular polysaccharide–protein conjugate vaccines suggest that a multivalent conjugate vaccine would be safe and highly immunogenic.

INFECTION IN ADULTS

The majority of GBS infections in otherwise healthy adults are related to pregnancy and parturition. Peripartum fever, the most common manifestation, is sometimes accompanied by symptoms and signs of endometritis or chorioamnionitis (abdominal distention and uterine or adnexal tenderness). Blood and vaginal swab cultures are often positive. Bacteremia is usually transitory but occasionally results in meningitis or endocarditis. Infections in adults that are not associated with the peripartum period generally involve individuals who are elderly or have an underlying chronic illness, such as diabetes mellitus or a malignancy. Among the infections that develop with some frequency in adults are cellulitis and soft tissue infection (including infected diabetic skin ulcers), urinary tract infection, pneumonia, endocarditis, and septic arthritis. Other reported infections include meningitis, osteomyelitis, and intraabdominal or pelvic abscesses. Relapse or recurrence of invasive infection weeks to months after a first episode is documented in ~4% of cases.

TREATMENT GROUP B STREPTOCOCCAL INFECTION IN ADULTS

GBS is less sensitive to penicillin than GAS, requiring somewhat higher doses. Adults with serious localized infections (pneumonia, pyelonephritis, abscess) should receive doses of ~12 million units of penicillin G daily; patients with endocarditis or meningitis should receive 18–24 million units per day in divided doses. Vancomycin is an acceptable alternative for penicillin-allergic patients.

NONENTEROCOCCAL GROUP D STREPTOCOCCI

The main nonenterococcal group D streptococci that cause human infections were previously considered a single species, *Streptococcus bovis*. The organisms encompassed by *S. bovis* have been reclassified into two species, each of which has two subspecies: *Streptococcus gallolyticus* subspecies *gallolyticus*, *S. gallolyticus* subspecies *pasteurianus*, *Streptococcus infantarius* subspecies *infantarius*, and *S. infantarius* subspecies *coli*. Endocarditis caused by these organisms is often associated with neoplasms of the gastrointestinal tract—most frequently, a colon carcinoma or polyp—but is also reported in association with other bowel lesions. When occult gastrointestinal lesions are carefully sought, abnormalities are found in $> 60\%$ of patients with endocarditis due to *S. gallolyticus* or *S. infantarius*. In contrast to the enterococci, nonenterococcal group D streptococci like these organisms are reliably killed by penicillin as a single agent, and penicillin is the agent of choice for the infections they cause.

VIRIDANS AND OTHER STREPTOCOCCI

VIRIDANS STREPTOCOCCI

Consisting of multiple species of α -hemolytic streptococci, the viridans streptococci are a heterogeneous group of organisms that are important agents of bacterial endocarditis (Chap. 155). Several species of viridans streptococci, including *Streptococcus salivarius*, *Streptococcus mitis*, *Streptococcus sanguis*, and *Streptococcus mutans*, are part of the normal flora of the mouth, where they live in close association with the teeth and gingiva. Some species contribute to the development of dental caries.

Previously known as *Streptococcus morbillorum*, *Gemella morbillorum* has been placed in a separate genus, along with *Gemella haemolysans*, on the basis of genetic-relatedness studies. These species resemble viridans streptococci with respect to habitat in the human host and associated infections.

The transient viridans streptococcal bacteremia induced by eating, toothbrushing, flossing, and other sources of minor trauma, together with adherence to biologic surfaces, is thought to account for the predilection of these organisms to cause endocarditis (see Fig. 155-1). Viridans streptococci are also isolated, often as part of a mixed flora, from sites of sinusitis, brain abscess, and liver abscess.

Viridans streptococcal bacteremia occurs relatively frequently in neutropenic patients, particularly after bone marrow transplantation or high-dose chemotherapy for cancer. Some of these patients develop a sepsis syndrome with high fever and shock. Risk factors for viridans streptococcal bacteremia include chemotherapy with high-dose cytosine arabinoside, prior treatment with trimethoprim-sulfamethoxazole or a fluoroquinolone, treatment with antacids or histamine antagonists, mucositis, and profound neutropenia.

The *S. milleri* group (also referred to as the *S. intermedius* or *S. anginosus* group) includes three species that cause human disease: *S. intermedius*, *S. anginosus*, and *Streptococcus constellatus*. These organisms are often considered viridans streptococci, although they differ somewhat from other viridans streptococci in both their hemolytic pattern (they may be α -, β -, or nonhemolytic) and the disease syndromes they cause. This group commonly produces suppurative infections, particularly abscesses of brain and abdominal viscera, and infections related to the oral cavity or respiratory tract, such as peritonsillar abscess, lung abscess, and empyema.

TREATMENT INFECTION WITH VIRIDANS STREPTOCOCCI

Isolates from neutropenic patients with bacteremia are often resistant to penicillin; thus these patients should be treated presumptively with vancomycin until the results of susceptibility testing become available. Viridans streptococci isolated in other clinical settings usually are sensitive to penicillin.

ABIOTROPHIA AND GRANULICATELLA SPECIES (NUTRITIONALLY VARIANT STREPTOCOCCI)

Occasional isolates cultured from the blood of patients with endocarditis fail to grow when subcultured on solid media. These *nutritionally variant streptococci* require supplemental thiol compounds or active forms of vitamin B₆ (pyridoxal or pyridoxamine) for growth in the laboratory. The nutritionally variant streptococci are generally grouped with the viridans streptococci because they cause similar types of infections. However, they have been reclassified on the basis of 16S ribosomal RNA sequence comparisons into two separate genera: *Abiotrophia*, with a single species (*Abiotrophia defectiva*), and *Granulicatella*, with three species associated with human infection (*Granulicatella adiacens*, *Granulicatella para-adiacens*, and *Granulicatella elegans*).

TREATMENT INFECTION WITH NUTRITIONALLY VARIANT STREPTOCOCCI

Treatment failure and relapse appear to be more common in cases of endocarditis due to nutritionally variant streptococci than in those due to the usual viridans streptococci. Thus the addition of

gentamicin (1 mg/kg every 8 h for patients with normal renal function) to the penicillin regimen is recommended for endocarditis due to the nutritionally variant organisms.

OTHER STREPTOCOCCI

Streptococcus suis is an important pathogen in swine and has been reported to cause meningitis in humans, usually in individuals with occupational exposure to pigs. Strains of *S. suis* associated with human infections have generally reacted with Lancefield group R typing serum and sometimes with group D typing serum as well. Isolates may be α - or β -hemolytic and are sensitive to penicillin. *Streptococcus iniae*, a pathogen of fish, has been associated with infections in humans who have handled live or freshly killed fish. Cellulitis of the hand is the most common form of human infection, although bacteremia and endocarditis have been reported. *Anaerobic streptococci*, or *peptostreptococci*, are part of the normal flora of the oral cavity, bowel, and vagina. **Infections caused by the anaerobic streptococci are discussed in Chap. 201.**

174 Enterococcal Infections

Cesar A. Arias, Barbara E. Murray

Enterococci have been recognized as potential human pathogens for more than a century, but only in recent years have these organisms acquired prominence as important causes of nosocomial infections. The ability of enterococci to survive and/or disseminate in the hospital environment and to acquire antibiotic resistance determinants makes the treatment of some enterococcal infections in critically ill patients a difficult challenge. Enterococci were first mentioned in the French literature in 1899; the “entérocoque” was found in the human gastrointestinal tract and was noted to have the potential to produce significant disease. Indeed, the first pathologic description of an enterococcal infection dates to the same year. A clinical isolate from a patient who died as a consequence of endocarditis was initially designated *Micrococcus zymogenes*, was later named *Streptococcus faecalis* subspecies *zymogenes*, and would now be classified as *Enterococcus faecalis*. The ability of this isolate to cause severe disease in both rabbits and mice illustrated its potential lethality in the appropriate settings.

ETIOLOGY

Enterococci are gram-positive organisms. In clinical specimens, they are usually observed as single cells, diplococci, or short chains (Fig. 174-1), although long chains are noted with some strains. Enterococci were originally classified as streptococci because organisms of the two genera share many morphologic and phenotypic characteristics, including a generally negative catalase reaction. Only DNA hybridization studies and later 16S rRNA sequencing clearly demonstrated that enterococci should be grouped as a genus distinct from the streptococci. Nonetheless, unlike the majority of streptococci, enterococci hydrolyze esculin in the presence of 40% bile salts and grow at high salt concentrations (e.g., 6.5%) and at high temperatures (46°C). Enterococci are usually reported by the clinical laboratory to be nonhemolytic on the basis of their inability to lyse the ovine or bovine red blood cells (RBCs) commonly used in agar plates; however, some strains of *E. faecalis* do lyse RBCs from humans, horses, and rabbits. The majority of clinically relevant enterococcal species hydrolyze pyrrolidonyl- β -naphthylamide (PYR); this characteristic is helpful in differentiating enterococci from organisms of the *Streptococcus gallolyticus* group (formerly known as *S. bovis*), which includes *S. gallolyticus*, *Streptococcus pasteurianus*, and *Streptococcus infantarius*, and from *Leuconostoc* species. Although at least 18 species of enterococci have been isolated from human infections, the overwhelming

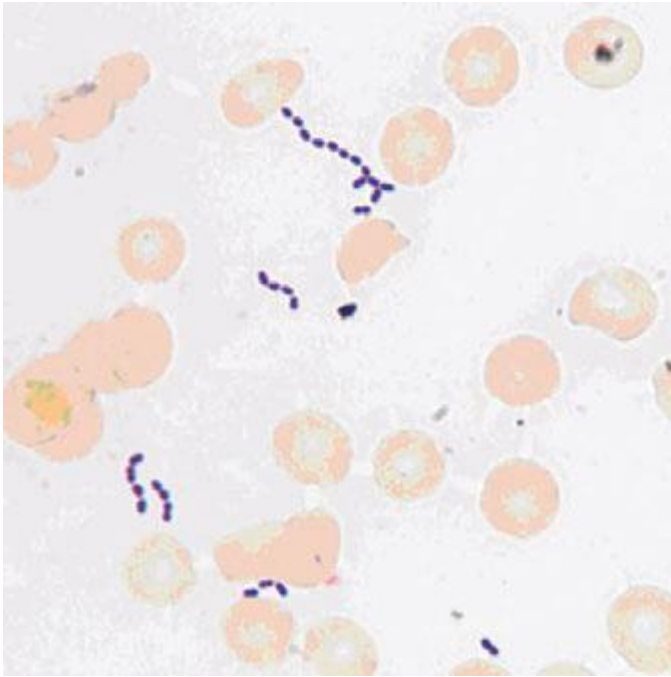


FIGURE 174-1 Gram's stain of cultured blood from a patient with enterococcal bacteremia. Oval gram-positive bacterial cells are arranged as diplococci and short chains. (Courtesy of Audrey Wanger, PhD.)

majority of cases are caused by two species, *E. faecalis* and *Enterococcus faecium*. Less frequently isolated species include *Enterococcus gallinarum*, *Enterococcus durans*, *Enterococcus hirae*, and *Enterococcus avium*.

PATHOGENESIS

Enterococci are normal inhabitants of the large bowel of human adults, although they usually make up <1% of the culturable intestinal microflora. In the healthy human gastrointestinal tract, enterococci are typical symbionts that coexist with other gastrointestinal bacteria; in fact, the utility of certain enterococcal strains as probiotics in the treatment of diarrhea suggests their possible role in maintaining the homeostatic equilibrium of the bowel. Enterococci are intrinsically resistant to a variety of commonly used antibacterial drugs. One of the most important factors that disrupts this equilibrium and promotes increased gastrointestinal colonization by enterococci is the administration of antimicrobial agents. In particular, antibiotics that are excreted in the bile and have broad-spectrum activity (e.g., certain cephalosporins that target anaerobes and gram-negative bacteria) are usually associated with the recovery of higher numbers of enterococci from feces. This increased colonization appears to be due not only to the simple enterococcal replacement in a “biologic niche” after the eradication of competing components of the flora, but also (at least in mice) to the suppression—upon reduction of the gram-negative microflora by antibiotics—of important immunologic signals (e.g., by the lectin RegIII γ) that help keep enterococcal counts low in the normal human bowel. Several studies have shown that higher levels of gastrointestinal colonization are a critical factor in the pathogenesis of enterococcal infections. However, the mechanisms by which enterococci successfully colonize the bowel and gain access to the lymphatics and/or bloodstream remain incompletely understood.

Several vertebrate, worm, and insect models have been developed to study the role of possible pathogenic determinants in both *E. faecalis* and *E. faecium*. Three main groups of virulence factors may increase the ability of enterococci to colonize the gastrointestinal tract and/or cause disease. The first group, *enterococcal secreted factors*, are molecules released outside the bacterial cell that contribute to the process of infection. The best-studied of these molecules include enterococcal hemolysin/cytolysin and two enterococcal proteases (gelatinase

and serine protease). Enterococcal cytolysin is a heterodimeric toxin produced by some strains of *E. faecalis* that is capable of lysing human RBCs as well as polymorphonuclear leukocytes and macrophages. *E. faecalis* gelatinase and serine protease are thought to mediate virulence by several mechanisms, including the degradation of host tissues and the modification of critical components of the immune system. Mutants lacking the genes corresponding to these proteins are highly attenuated in experimental peritonitis, endocarditis, and endophthalmitis.

A second group of virulence factors, *enterococcal surface components*, are thought to contribute to bacterial attachment to extracellular matrix molecules in the human host. Several molecules on the surface of enterococci have been characterized and shown to play a role in the pathogenesis of enterococcal infections. Among the characterized adhesins is aggregation substance of *E. faecalis*, which mediates the attachment of bacterial cells to each other, thereby facilitating conjugative plasmid exchange. Several lines of evidence indicate that aggregation substance and enterococcal cytolysin act synergistically to increase the virulence potential of *E. faecalis* strains in experimental endocarditis. The surface protein adhesin of collagen of *E. faecalis* (Ace) and its *E. faecium* homologue (Acm) recognize adhesive matrix molecules (MSCRAMMs) involved in bacterial attachment to host proteins such as collagen, fibronectin, and fibrinogen; both Ace and Acm are important in the pathogenesis of experimental endocarditis. Pili of gram-positive bacteria have been shown to be important mediators of attachment to and invasion of host tissues and are considered potential targets for immunotherapy. Both *E. faecalis* and *E. faecium* have surface pili. Mutants of *E. faecalis* lacking pili are attenuated in biofilm production, experimental endocarditis, and urinary tract infections (UTIs). Other surface proteins that share structural homology with MSCRAMMs and appear to play a role in enterococcal attachment to the host and in virulence include the *E. faecalis* surface protein Esp and its *E. faecium* homologue Esp_{fm}, the second collagen adhesin of *E. faecium* (Scm), the surface proteins of *E. faecium* (Fms), SgrA (which binds to components of the basal lamina), and EcbA (which binds to collagen type V). Additional surface components apparently associated with pathogenicity include the Elr protein (a protein from the WxL family) and polysaccharides, which are thought to interfere with phagocytosis of the organism by host immune cells. Some *E. faecalis* strains appear to harbor at least three distinct classes of capsular polysaccharide; some of these polysaccharides play a role in virulence and are potential targets for immunotherapy.

The third group of virulence factors has not been well characterized but consists of the *E. faecalis* stress protein Gls24, which has been associated with enterococcal resistance to bile salts and appears to be important in the pathogenesis of endocarditis, and the *hyl_{Efm}*-containing plasmids of *E. faecium*, which are transferable between strains and increase gastrointestinal colonization by *E. faecium*. In mouse peritonitis, acquisition of these plasmids increased the lethality of a commensal strain of *E. faecium*. Recently, a gene encoding a regulator of oxidative stress (AsrR) has been identified as an important virulence factor of *E. faecium*.



The ability to sequence bacterial genomes has increased our understanding of bacterial diversity, evolution, pathogenesis, and mechanisms of antibiotic resistance. The genome sequences of more than 560 enterococcal strains are currently available, and some have been entirely closed and annotated. Sequence analysis has shown that the genetic diversity of enterococci is related in large part to the acquisition of exogenous DNA and the mobilization of large chromosomal regions, resulting in recombination of the “core” genomes. In addition, analyses indicate that *E. faecium* harbors a malleable *accessory genome* incorporating a substantial content of exogenous elements, including DNA from phages. Indeed, a hospital-associated *E. faecium* clade that contains most clinical and outbreak-associated strains is the predominant genetic lineage circulating in hospitals around the world. This clade appears to be evolving rapidly, and genomic comparisons suggest that this lineage emerged 75 years ago—a time point that coincides with the introduction of antimicrobial drugs—and evolved from animal strains, not from human commensal isolates. An initial genomic

separation within *E. faecium* appears to have occurred ~3000 years ago, simultaneous with urbanization and domestication of animals. This genomic information provides new clues with regard to the evolution of enterococci from commensal organisms to important nosocomial pathogens.

EPIDEMIOLOGY

According to the National Healthcare Safety Network of the Centers for Disease Control and Prevention, enterococci are the second most common organisms (after staphylococci) isolated from hospital-associated infections in the United States. Although *E. faecalis* remains the predominant species recovered from nosocomial infections, the isolation of *E. faecium* has increased substantially in the past 20 years. In fact, *E. faecium* is now almost as common as *E. faecalis* as an etiologic agent of hospital-associated infections. This point is important, because *E. faecium* is by far the most resistant and challenging enterococcal species to treat; indeed, more than 80% of *E. faecium* isolates recovered in U.S. hospitals are resistant to vancomycin, and more than 90% are resistant to ampicillin (historically the most effective β -lactam agent against enterococci). Resistance to vancomycin and ampicillin in *E. faecalis* isolates is much less common (~7% and ~4%, respectively).

The dynamics of enterococcal transmission and dissemination in the hospital environment have been extensively studied, with a focus on vancomycin-resistant enterococci (VRE). These studies have revealed that VRE colonization of the gastrointestinal tract is a critical step in the natural history of enterococcal disease and that a substantial proportion of patients colonized with VRE remain colonized for prolonged periods (sometimes >1 year) and are more likely to develop an *Enterococcus*-related illness (e.g., bacteremia). The most important factors associated with VRE colonization and persistence in the gut include prolonged hospitalization; long courses of antibiotic therapy; hospitalization in long-term-care facilities, surgical units, and/or intensive care units; organ transplantation; renal failure (particularly in patients undergoing hemodialysis) and/or diabetes; high Acute Physiology and Chronic Health Evaluation (APACHE) scores; and physical proximity to patients infected or colonized with VRE or these patients' rooms. Once a patient becomes colonized with VRE, several key factors are involved in the organisms' dissemination in the hospital environment. VRE can survive exposure to heat and certain disinfectants and have been found on numerous inanimate objects in the hospital, including bed rails, medical equipment, doorknobs, gloves, telephones, and computer keyboards. Thus health care workers and the environment play pivotal roles in enterococcal transmission from patient to patient, and infection control measures are crucial in breaking the chain of transmission. Moreover, two meta-analyses have found that, independent of the patient's clinical status, VRE infection increases the risk of death over that among individuals infected with a glycopeptide-susceptible enterococcal strain.



The epidemiology of enterococcal disease and the emergence of VRE have followed slightly different trends in other parts of the world than in the United States. In Europe, the emergence of VRE in the mid-1980s was seen primarily in isolates recovered from animals and healthy humans rather than from hospitalized patients. The presence of VRE was associated with the use of the glycopeptide avoparcin as a growth promoter in animal feeds; this association prompted the European Union to ban the use of this compound in animal husbandry in 1996. However, after an initial decrease in the isolation of VRE from animals and humans, the prevalence of hospital-associated VRE infections has slowly increased in certain European countries, with important regional differences. For example, rates of vancomycin resistance among *E. faecium* clinical isolates in Europe are highest in Greece, the United Kingdom, and Portugal (10–30%), whereas rates in the Scandinavian countries and the Netherlands are <1%. These regional differences have been attributed in part to the implementation of aggressive “search-and-destroy” policies of infection control in countries such as the Netherlands; these policies have kept the frequency of nosocomial methicillin-resistant *Staphylococcus*

aureus (MRSA) and VRE very low. Despite regional differences, rates of VRE continue to be much lower in Europe than in the United States. The reasons are not totally understood, although it has been postulated that this difference is related to the higher levels of human antibiotic use in the United States. Rates of enterococcal resistance to vancomycin in some Latin American countries are also lower (~4%) than those in the United States. Conversely, in Asia, rates of vancomycin resistance among enterococci appear to be similar to those in U.S. hospitals.



As mentioned above, genomic analyses of vancomycin-resistant *E. faecium* in different parts of the world suggest that the emergence and dissemination of these organisms in the hospital environment worldwide are due to the success of a unique hospital-associated genetic clade that acquired the genes responsible for vancomycin resistance as well as other antibiotic resistance determinants.

CLINICAL SYNDROMES

URINARY TRACT INFECTION AND PROSTATITIS

Enterococci are well-known causes of nosocomial UTI—the most common infection caused by these organisms (Chap. 162). Enterococcal UTIs are usually associated with indwelling catheters, instrumentation, or anatomic abnormalities of the genitourinary tract, and it is often challenging to differentiate between true infection and colonization (particularly in patients with chronic indwelling catheters). The presence of leukocytes in the urine in conjunction with systemic manifestations (e.g., fever) or local signs and symptoms of infection with no other explanation and a positive urine culture ($\geq 10^5$ colony-forming units [CFU]/mL) suggests the diagnosis. Moreover, enterococcal UTIs often occur in critically ill patients whose comorbidities may obscure the diagnosis. In many cases, removal of the indwelling catheter may suffice to eradicate the organism without specific antimicrobial therapy. In rare circumstances, UTIs caused by enterococci may run a complicated course, with the development of pyelonephritis and perinephric abscesses that may be a portal of entry for bloodstream infections (see below). Enterococci are also known causes of chronic prostatitis, particularly in patients whose urinary tract has been manipulated surgically or endoscopically. These infections can be difficult to treat because the agents most potent against enterococci (i.e., aminopenicillins and glycopeptides) penetrate prostatic tissue poorly. Chronic prostatic infection can be a source of recurrent enterococcal bacteremia.

BACTEREMIA AND ENDOCARDITIS

Bacteremia without endocarditis is one of the most common presentations of enterococcal disease. Intravascular catheters and other devices are commonly associated with these bacteremic episodes (Chap. 168). Other well-known sources of enterococcal bacteremia include the gastrointestinal and hepatobiliary tracts; pelvic and intraabdominal foci; and, less frequently, wound infections, UTIs, and bone infections. In the United States, enterococci are ranked second (after coagulase-negative staphylococci) as etiologic agents of central line-associated bacteremia. Patients with enterococcal bacteremia usually have comorbidities and have been in the hospital for prolonged periods; they commonly have received several courses of antibiotics. Several studies indicate that the isolation of *E. faecium* from the blood may lead to worse outcomes and higher mortality rates than when other enterococcal species are isolated; this finding may be related to the higher prevalence of vancomycin and ampicillin resistance in *E. faecium* than in other enterococcal species, with the consequent reduction of therapeutic options. In many cases (usually when the gastrointestinal tract is the source), enterococcal bacteremia may be polymicrobial, with gram-negative organisms isolated at the same time. In addition, several cases have now been documented in which enterococcal bacteremia was associated with *Strongyloides stercoralis* hyperinfection syndrome in immunocompromised patients.

Enterococci are important causes of community- and health care-associated endocarditis, ranking second after staphylococci in the

974 latter infections. The presumed initial source of bacteremia leading to endocarditis is the gastrointestinal or genitourinary tract—e.g., in patients who have malignant and inflammatory conditions of the gut or have undergone procedures in which these tracts are manipulated. The affected patients tend to be male and elderly and to have other debilitating diseases and heart conditions. Both prosthetic and native valves can be involved; mitral and aortic valves are affected most often. Community-associated endocarditis (usually caused by *E. faecalis*) also occurs in patients with no apparent risk factors or cardiac abnormalities. Endocarditis in women of childbearing age has been well described. The typical presentation of enterococcal endocarditis is a subacute course of fever, weight loss, malaise, and cardiac murmur; typical stigmata of endocarditis (e.g., petechiae, Osler's nodes, Roth's spots) are found in only a minority of patients. Atypical manifestations include arthralgias and manifestations of metastatic disease (splenic abscesses, hiccups, pain in the left flank, pleural effusion, and spondylodiscitis). Embolic complications are variable and can affect the brain. Heart failure is a common complication of enterococcal endocarditis, and valve replacement may be critical in curing this infection, particularly when multidrug-resistant organisms or major complications are involved. The duration of therapy is usually 4–6 weeks, with more prolonged courses suggested for multidrug-resistant isolates in the absence of valvular replacement.

MENINGITIS

Enterococcal meningitis is an uncommon disease (accounting for only ~4% of meningitis cases) that is usually associated with neurosurgical interventions and conditions such as shunts, central nervous system (CNS) trauma, and cerebrospinal fluid (CSF) leakage. In some instances—usually in patients with a debilitating condition, such as cardiovascular or congenital heart disease, chronic renal failure, malignancy, receipt of immunosuppressive therapy, or HIV/AIDS—presumed hematogenous seeding of the meninges is seen in infections such as endocarditis or bacteremia. Fever and changes in mental status are common, whereas overt meningeal signs are less so. CSF findings are consistent with bacterial infection—i.e., pleocytosis with a predominance of polymorphonuclear leukocytes (average, ~500/μL), an elevated serum protein level (usually >100 mg/dL), and a decreased glucose concentration (average, 28 mg/dL). Gram's staining yields a positive result in about half of cases, with a high rate of organism recovery from CSF cultures; the most common species isolated are *E. faecalis* and *E. faecium*. Complications include hydrocephalus, brain abscesses, and stroke. As mentioned before for bacteremia, an association with *Strongyloides* hyperinfection has also been documented.

INTRAABDOMINAL, PELVIC, AND SOFT TISSUE INFECTIONS

As mentioned earlier, enterococci are part of the commensal flora of the gastrointestinal tract and can produce spontaneous peritonitis in cirrhotic individuals and in patients undergoing chronic ambulatory peritoneal dialysis (Chap. 159). These organisms are commonly found (usually along with other bacteria, including enteric gram-negative species and anaerobes) in clinical samples from intraabdominal and pelvic collections. The presence of enterococci in intraabdominal infections is sometimes considered to be of little clinical relevance. Several studies have shown that the role of enterococci in intraabdominal infections originating in the community and involving previously healthy patients is minor, because surgery and broad-spectrum antimicrobial drugs that do not target enterococci are often sufficient to treat these infections successfully. In the last few decades, however, these organisms have become prominent as a cause of intraabdominal infections in hospitalized patients because of the emergence and spread of vancomycin resistance among enterococci and the increase in rates of nosocomial infections due to multidrug-resistant *E. faecium* isolates. In fact, several studies have now documented treatment failures due to enterococci, with consequently increased rates of postoperative complications and death among patients with intraabdominal infections. Thus, anti-enterococcal therapy is recommended for nosocomial peritonitis in immunocompromised and severely ill patients who have had

a prolonged hospital stay, have undergone multiple procedures, have persistent abdominal sepsis and collections, or have risk factors for the development of endocarditis (e.g., prosthetic or damaged heart valves). Conversely, specific treatment for enterococci in the first episode of intraabdominal infections originating in the community and affecting previously healthy patients with no important cardiac risk factors for endocarditis does not appear to be beneficial.

Enterococci are commonly isolated from soft tissue infections (Chap. 156), particularly those involving surgical wounds (Chap. 168). In fact, these organisms rank third as agents of nosocomial surgical-site infections, with *E. faecalis* the most frequently isolated species. The clinical relevance of enterococci in some of these infections—as in intraabdominal infections—is a matter of debate; differentiating between colonization and true infection is sometimes challenging, although in some cases enterococci have been recovered from lung, liver, and skin abscesses. Diabetic foot and decubitus ulcers are often colonized with enterococci and may be the portal of entry for bone infections.

OTHER INFECTIONS

Enterococci are well-known causes of neonatal infections, including sepsis (mostly late-onset), bacteremia, meningitis, pneumonia, and UTI. Outbreaks of enterococcal sepsis in neonatal units have been well documented. Risk factors for enterococcal disease in newborns include prematurity, low birth weight, indwelling devices, and abdominal surgery. Enterococci have also been described as etiologic agents of bone and joint infections, including vertebral osteomyelitis, usually in patients with underlying conditions such as diabetes or endocarditis. Similarly, enterococci have been isolated from bone infections in patients who have undergone arthroplasty or reconstruction of fractures with the placement of hardware. Because enterococci can produce a biofilm that is likely to alter the efficacy of otherwise active anti-enterococcal agents, treatment of infections that involve foreign material is challenging, and removal of the hardware may be necessary to eradicate the infection. Rare cases of enterococcal pneumonia, lung abscess, and spontaneous empyema have been described.

TREATMENT ENTEROCOCCAL INFECTIONS

GENERAL PRINCIPLES

Enterococci are intrinsically resistant and/or tolerant to several antimicrobial agents (with *tolerance* defined as lack of killing by drug concentrations 32 times higher than the minimal inhibitory concentration [MIC]). Monotherapy for endocarditis with a β -lactam antibiotic (to which many enterococci are tolerant) has produced disappointing results, with low cure rates at the end of therapy. However, the addition of an aminoglycoside to a cell wall-active agent (a β -lactam or a glycopeptide) increases cure rates and eradicates the organisms; moreover, this combination is synergistic and bactericidal in vitro. Therefore, for many decades, combination therapy with a cell wall-active agent and an aminoglycoside has been the standard of care for endovascular infections caused by enterococci. This synergistic effect can be explained, at least in part, by the increased penetration of the aminoglycoside into the bacterial cell, presumably as a result of cell wall alterations produced by the β -lactam (or glycopeptide). Nonetheless, attaining synergistic bactericidal activity in the treatment of severe enterococcal infections has become increasingly difficult because of the development of resistance to virtually all antibiotics available for this purpose.

The treatment of *E. faecalis* differs substantially from that of *E. faecium* (Tables 174-1 and 174-2), mainly because of differences in resistance profiles (see below). For example, resistance to ampicillin and vancomycin is rare in *E. faecalis*, whereas these antibiotics are only infrequently useful against current isolates of *E. faecium*. Moreover, as a consequence of the challenges and therapeutic limitations posed by the emergence of drug resistance in enterococci,

TABLE 174-1 SUGGESTED REGIMENS FOR THE MANAGEMENT OF INFECTIONS CAUSED BY ENTEROCOCCUS FAECALIS

Clinical Syndrome	Suggested Therapeutic Options ^a
Endovascular infections (including endocarditis)	<ul style="list-style-type: none"> • Ampicillin^b (12 g/d IV in divided doses q4h or by continuous infusion) or penicillin (18–30 million units/d IV in divided doses q4h or by continuous infusion) plus an aminoglycoside^c • Ampicillin^b (12 g/d IV in divided doses q4h) plus ceftriaxone (2 g IV q12h) • Vancomycin^d (15 mg/kg IV per dose) plus an aminoglycoside^c • High-dose daptomycin^e ± another active agent^f • Ampicillin^b plus imipenem
Nonendovascular bacteremia ^g	<ul style="list-style-type: none"> • Ampicillin (12 g/d IV in divided doses q4h) or penicillin (18 million units/d IV in divided doses q4h) ± an aminoglycoside^c or ceftriaxone • Vancomycin^d (15 mg/kg IV per dose) • High-dose daptomycin^e ± another active agent^f • Linezolid (600 mg IV/PO q12h)
Meningitis	<ul style="list-style-type: none"> • Ampicillin (20–24 g/d IV in divided doses q4h) or penicillin (24 million units/d IV in divided doses q4h) plus an aminoglycoside^{ch} or ceftriaxone • Vancomycin (500–750 mg IV q6h)^d plus an aminoglycoside^c or rifampin • Linezolid • High-dose daptomycin^e (plus intrathecal daptomycin) ± another active agent^f
Urinary tract infections (uncomplicated)	<ul style="list-style-type: none"> • Fosfomycin (3 g PO, one dose)ⁱ • Ampicillin (500 mg IV or PO q6h) • Nitrofurantoin (100 mg PO q6h)

^aAuthors' preferences are underlined for each category; many of these regimens are off-label. ^bIn rare cases, β-lactamase-producing isolates may be found. Because these isolates are not detected by conventional minimal inhibitory concentration determination, additional tests (e.g., the nitrocefin disk) are recommended for isolates from endocarditis. The use of ampicillin/sulbactam (12–24 g/d) is suggested in these cases. ^cOnly if the organism does not exhibit high-level resistance (HLR) to aminoglycosides. HLR is assessed by the clinical microbiology laboratory only for gentamicin or streptomycin, because gentamicin (1–1.5 mg/kg IV q8h) and streptomycin (15 mg/kg per day IV/IM, in two divided doses) are the only two recommended aminoglycosides. The test used to detect HLR is the growth of enterococci on agar containing gentamicin (500 μg/mL) or streptomycin (2000 μg/mL). If HLR is documented, the aminoglycoside will not act synergistically with the other agent in the combination. HLR to gentamicin implies lack of synergism with tobramycin and with amikacin. ^dVancomycin is recommended only as an alternative to β-lactam agents in cases of allergy, toxicity, and inability to desensitize. Cerebrospinal fluid (CSF) concentrations should be determined in meningitis. Vancomycin-resistant strains of *E. faecalis* have been reported. ^eConsider doses of 8–10 mg/kg per day if used in combination and 10–12 mg/kg per day if used alone. Close monitoring of creatine phosphokinase levels is recommended throughout therapy because of possible rhabdomyolysis. ^fPotentially active agents may include an aminoglycoside (if HLR is not detected), ampicillin, ceftaroline, tigecycline, or a fluoroquinolone (which, if the isolate is susceptible, may be favored in meningitis). ^gIn selected cases of catheter-associated bacteremia, removal of the catheter and a short course of therapy (~5–7 days) may be sufficient. A single positive blood culture that is likely to be associated with a catheter in a patient who is otherwise doing well may not require therapy after removal of the catheter. Patients at high risk for endovascular infections or with severe disease may benefit from synergistic combination therapy. ^hThe addition of intrathecal or intraventricular therapy with gentamicin (2–10 mg/d if the organism does not exhibit HLR) or vancomycin (10–20 mg/d when the isolate is susceptible) has been suggested by some authorities. The addition of systemic rifampin (a good CSF-penetrating agent) may be considered. The combination of ampicillin and ceftriaxone may have clinical benefit (by analogy with endocarditis), but no cases treated with this combination have been reported. ⁱApproved by the Food and Drug Administration only for uncomplicated urinary tract infections caused by vancomycin-susceptible *E. faecalis*.

TABLE 174-2 SUGGESTED REGIMENS FOR THE MANAGEMENT OF INFECTIONS CAUSED BY VANCOMYCIN- AND AMPICILLIN-RESISTANT ENTEROCOCCUS FAECIUM

Clinical Syndrome	Suggested Therapeutic Options ^a
Endovascular infections (including endocarditis)	<ul style="list-style-type: none"> • High-dose daptomycin^b plus another agent^c ± an aminoglycoside^d • Q/D^e (22.5 mg/kg per day in divided doses q8h) ± another active agent^f • Linezolid^g (600 mg IV q12h) • High-dose ampicillin (if MIC is ≤64 μg/mL) ± an aminoglycoside^d
Nonendovascular bacteremia ^g	<ul style="list-style-type: none"> • High-dose daptomycin^b ± another agent^c ± an aminoglycoside^d • Q/D (22.5 mg/kg per day in divided doses q8h) ± another active agent^f • Linezolid (600 mg IV q12h)
Meningitis	<ul style="list-style-type: none"> • Linezolid (600 mg IV q12h) ± another CSF-penetrating active agent^h • Q/D (22.5 mg/kg per day in divided doses q8h plus intraventricular Q/D)ⁱ ± another active agent^h • High-dose daptomycin^b (plus intraventricular daptomycin) ± another CSF-penetrating active agent^{h,j}
Urinary tract infections	<ul style="list-style-type: none"> • Fosfomycin (3 g PO, one dose)^k • Nitrofurantoin (100 mg PO q6h) • Ampicillin or amoxicillin (2 g IV/PO q4–6h)^l

^aAuthors' preferences are underlined for each category; many of these regimens are off-label. ^bConsider doses of 8–10 mg/kg per day if used in combination and 10–12 mg/kg per day if used alone (off-label). Close monitoring of creatine phosphokinase levels is recommended throughout therapy because of possible rhabdomyolysis. ^cPotentially active agents may include ampicillin or ceftaroline (even if the infecting strain is resistant in vitro) or tigecycline. In vitro synergism of daptomycin with some β-lactam agents is observed against some isolates that subsequently become nonsusceptible to daptomycin during therapy. Consider combination therapy if the daptomycin minimal inhibitory concentration (MIC) is ≥3 μg/mL. ^dOnly if the organism does not exhibit high-level resistance to aminoglycosides (see Table 174-1, footnote c). ^eQuinupristin-dalfopristin (Q/D) and linezolid are listed in the American Heart Association's recommendations for the treatment of endocarditis caused by vancomycin- and ampicillin-resistant *E. faecium*. ^fAgents that may be useful in combination with Q/D (if the isolate is susceptible to each agent) include doxycycline with rifampin (one reported case) and fluoroquinolones (one reported case). ^gIn selected cases of catheter-associated bacteremia, removal of the catheter and a short course of therapy (~5–7 days) may be sufficient. A single positive blood culture that is likely to be associated with a catheter in a patient who is otherwise doing well may not require therapy after removal of the catheter. ^hFluoroquinolone antibiotics (e.g., moxifloxacin) and rifampin (if the isolate is susceptible to each agent) reach therapeutic levels in the cerebrospinal fluid (CSF). ⁱIntrathecal Q/D (1–5 mg/d) has been used in combination with Q/D systemic therapy in meningitis. If Q/D is chosen, simultaneous use of both systemic and intrathecal therapy is suggested. ^jIntrathecal gentamicin (2–10 mg/d) if high-level resistance is not detected. Intraventricular daptomycin has been used in two cases of meningitis. ^kApproved by the Food and Drug Administration only for uncomplicated urinary tract infections caused by vancomycin-susceptible *E. faecalis*. ^lConcentrations of amoxicillin and ampicillin in urine far exceed those in serum and may be potentially effective even against isolates with high MICs. Doses up to 12 g/d are suggested for isolates with MICs of ≥64 μg/mL.

976 valve replacement may need to be considered in the treatment of endocarditis caused by multidrug-resistant enterococci. Less severe infections are often related to indwelling intravascular catheters; removal of the catheter increases the likelihood of enterococcal eradication by a subsequent short course of appropriate antimicrobial therapy.

CHOICE OF ANTIMICROBIAL AGENTS

Among the β -lactams, the most active are the aminopenicillins (ampicillin, amoxicillin) and ureidopenicillins (i.e., piperacillin); next most active are penicillin G and imipenem. For *E. faecium*, a combination of high-dose ampicillin (up to 30 g/d) plus an aminoglycoside has been suggested—even for ampicillin-resistant strains if the MIC is ≤ 64 $\mu\text{g/mL}$ —because a plasma ampicillin concentration of >100 $\mu\text{g/mL}$ can be achieved at high doses. The only two aminoglycosides recommended for synergistic therapy in severe enterococcal infections are gentamicin and streptomycin. The use of amikacin is discouraged, tobramycin should never be used against *E. faecium*, and aminoglycoside monotherapy is not effective. Vancomycin is an alternative to β -lactam drugs for the treatment of *E. faecalis* infections but is less useful against *E. faecium* because resistance is common.

As mentioned above, use of the aminoglycoside–ampicillin combination for *E. faecalis* infections has become increasingly problematic because of toxicity in critically ill patients and increased rates of high-level resistance to aminoglycosides. A recent observational, nonrandomized, comparative study encompassing a multicenter cohort was conducted in 17 Spanish hospitals and 1 Italian hospital; this study found that the combination of ampicillin and ceftriaxone is as effective as ampicillin plus gentamicin in the treatment of *E. faecalis* endocarditis, with less risk of toxicity. Therefore, this regimen should be considered in patients at risk for aminoglycoside toxicity and could be considered for all patients.

Linezolid and quinupristin/dalfopristin (Q/D) are two agents approved by the U.S. Food and Drug Administration (FDA) for the treatment of some VRE infections (Table 174–2). Linezolid is not bactericidal, and its use in severe endovascular infections has produced mixed results; therefore, it is recommended only as an alternative to other agents. In addition, linezolid may cause significant toxicities (thrombocytopenia, peripheral neuropathy, and optic neuritis) when used in regimens given for >2 weeks. Nonetheless, linezolid may play a role in the treatment of enterococcal meningitis and other CNS infections, although clinical data are limited. Q/D is not active against most *E. faecalis* isolates, and its in vivo efficacy against *E. faecium* may often be compromised by resistance (see below). Adverse reactions to Q/D are common, including pain and inflammation at the infusion site and severe arthralgias and myalgias leading to discontinuation of treatment. Thus, Q/D should be used with caution and probably combined with other agents (Table 174–2).

The lipopeptide daptomycin is a bactericidal antibiotic with potent in vitro activity against all enterococci. Although daptomycin is not approved by the FDA for the treatment of VRE or *E. faecium* infections, it has been used alone (at high dosage) or in combination with other agents (ampicillin, ceftaroline, and tigecycline) with apparent success against multidrug-resistant enterococcal infections (Tables 174–1 and 174–2). The main adverse reactions to daptomycin are elevated creatine phosphokinase levels and eosinophilic pneumonitis (rare). Daptomycin is not useful against pulmonary infections because the pulmonary surfactant inhibits its antibacterial activity. Although the glycolcycline drug tigecycline is active in vitro against all enterococci (regardless of the isolates' vancomycin susceptibility), its use as monotherapy for endovascular or

severe enterococcal infections is not recommended because of low attainable blood levels. Telavancin, a lipoglycopeptide approved by the FDA for the treatment of skin and soft tissue infections as well as hospital-associated pneumonia, is active against vancomycin-susceptible enterococci but not VRE. Oritavancin, a compound of the same class that is active against VRE, has recently been approved by the FDA for the treatment of bacterial skin and soft tissue infections and may offer promise for the treatment of VRE in the future.

ANTIMICROBIAL RESISTANCE

As mentioned above, resistance to β -lactam agents continues to be observed only infrequently in *E. faecalis*, although rare outbreaks caused by β -lactamase-producing isolates have occurred in the United States and Argentina. However, ampicillin resistance is common in *E. faecium*. The mechanism of this resistance is related to a penicillin-binding protein (PBP) designated PBP5, which is the target of β -lactam antibiotics. PBP5 exhibits lower affinity for ampicillin and can synthesize cell wall in the presence of this antibiotic, even when other PBPs are inhibited. Two common mechanisms of high-level ampicillin resistance (MIC, >64 $\mu\text{g/mL}$) in clinical strains are (1) mutations in the PBP5-encoding gene that further decrease the protein's affinity for ampicillin and (2) hyperproduction of PBP5. These factors preclude the use of all β -lactam agents in the treatment of *E. faecium* infections.

Vancomycin is a glycopeptide antibiotic that inhibits cell wall peptidoglycan synthesis in susceptible enterococci and has been widely used against enterococcal infections in clinical practice when the utility of β -lactams is limited by resistance, allergy, or adverse reactions. This effect is mediated by binding of the antibiotic to peptidoglycan precursors (UDP-MurNAC-pentapeptides) upon their exit from the bacterial cell cytoplasm. The interaction of vancomycin with the peptidoglycan is specific and involves the last two D-alanine residues of the precursor. The first isolates of VRE were documented in 1986, and vancomycin resistance (particularly in *E. faecium*) has since increased considerably around the world. The mechanism involves the replacement of the last D-alanine residue of peptidoglycan precursors with D-lactate or D-serine, with consequent high- and low-level resistance, respectively. There is significant heterogeneity among isolates, but either substitution substantially decreases the affinity of vancomycin for the peptidoglycan; with the D-lactate substitution, the MIC is increased by up to 1000-fold. Vancomycin-resistant organisms also produce enzymes that destroy the D-alanine-D-alanine ending precursors, ensuring that additional binding sites for vancomycin are not available.

High-level resistance to aminoglycosides (of which gentamicin and streptomycin are the only two tested by clinical laboratories) abolishes the synergism observed between cell wall-active agents and the aminoglycoside. This important phenotype is routinely sought in isolates from serious infections (Tables 174–1 and 174–2). The laboratory reports high-level resistance as gentamicin and streptomycin MICs of >500 $\mu\text{g/mL}$ and >2000 $\mu\text{g/mL}$, respectively (agar dilution method) or as "SYN-R" (resistance to synergism). Genes encoding aminoglycoside-modifying enzymes are usually the cause of high-level resistance to these compounds and are widely disseminated among enterococci, decreasing the options for the treatment of severe enterococcal infections. The aforementioned enterococcal resistance to newer antibiotics such as linezolid (usually due to mutations in the 23S rRNA genes and the presence of an rRNA methylase), Q/D, daptomycin (involving major changes in cell membrane homeostasis), and tigecycline further reduces therapeutic alternatives.

DIPHTHERIA

Diphtheria is a nasopharyngeal and skin infection caused by *Corynebacterium diphtheriae*. Toxigenic strains of *C. diphtheriae* produce a protein toxin that causes systemic toxicity, myocarditis, and polyneuropathy. The toxin is associated with the formation of pseudomembranes in the pharynx during respiratory diphtheria. While toxigenic strains most frequently cause pharyngeal diphtheria, nontoxigenic strains commonly cause cutaneous disease.

ETIOLOGY

C. diphtheriae is a gram-positive bacillus that is unencapsulated, nonmotile, and nonsporulating. The organism was first identified microscopically in 1883 by Klebs and a year later was isolated in pure culture by Löffler in Robert Koch's laboratory. The bacteria have a characteristic club-shaped bacillary appearance and typically form clusters of parallel rays, or *palisades*, that are referred to as "Chinese characters." The specific laboratory media recommended for the cultivation of *C. diphtheriae* rely upon tellurite, colistin, or nalidixic acid for the organism's selective isolation from other autochthonous pharyngeal microbes. *C. diphtheriae* may be isolated from individuals with both nontoxigenic (*tox*⁻) and toxigenic (*tox*⁺) phenotypes. Uchida and Pappenheimer demonstrated that corynebacteriophage beta carries the structural gene *tox*, which encodes diphtheria toxin, and that a family of closely related corynebacteriophages are responsible for toxigenic conversion of *tox*⁻ *C. diphtheriae* to the *tox*⁺ phenotype. Moreover, lysogenic conversion from a nontoxigenic to a toxigenic phenotype has been shown to occur in situ. Growth of toxigenic strains of *C. diphtheriae* under iron-limiting conditions leads to the optimal expression of diphtheria toxin and is believed to be a pathogenic mechanism during human infection.

EPIDEMIOLOGY

While in many regions diphtheria has been controlled in recent years with effective vaccination, there have been sporadic outbreaks in the United States and Europe. Diphtheria is still common in the Caribbean, Latin America, and the Indian subcontinent, where mass immunization programs are not enforced. Large-scale epidemics of diphtheria have occurred in the post-Soviet independent states. Additional outbreaks have been reported in Algeria, China, and Ecuador.

C. diphtheriae is transmitted via the aerosol route, usually during close contact with an infected person. There are no significant reservoirs other than humans. The incubation period for respiratory diphtheria is 2–5 days, but disease onset has occurred as late as 10 days after exposure. Prior to the vaccination era, most individuals over the age of 10 were immune to *C. diphtheriae*; infants were protected by maternal IgG antibodies but became susceptible after ~6 months of age. Thus, the disease primarily affected children and nonimmune young adults. In temperate regions, respiratory diphtheria occurs year-round but is most common during winter months.

The development of diphtheria antitoxin in 1898 by von Behring and of the diphtheria toxoid vaccine in 1924 by Ramon led to the near-elimination of diphtheria in Western countries. The annual incidence rate in the United States peaked in 1921 at 191 cases per 100,000 population. In contrast, since 1980, the annual figure in the United States has been <5 cases per 100,000. Nevertheless, pockets of colonization persist in North America, particularly in South Dakota, Ontario, and recently the state of Washington. Immunity to diphtheria induced by childhood vaccination gradually decreases in adulthood. An estimated 30% of men 60–69 years old have antitoxin titers below the protective

level. In addition to older age and lack of vaccination, risk factors for diphtheria outbreaks include alcoholism, low socioeconomic status, crowded living conditions, and Native American ethnic background. An outbreak of diphtheria in Seattle, Washington, between 1972 and 1982 comprised 1100 cases, most of which were cutaneous. During the 1990s in the states of the former Soviet Union, a much larger diphtheria epidemic included more than 150,000 cases and more than 5000 deaths. Clonally related toxigenic *C. diphtheriae* strains of the ET8 complex were associated with this outbreak. Given that the ET8 complex expressed a toxin against which the prevalent diphtheria toxoid vaccine was effective, the epidemic was attributed to failure of the public health infrastructure to effectively vaccinate the population. Beginning in 1998, this epidemic was controlled by mass vaccination programs. During the epidemic, the incidence rate was high among individuals between 16 and 50 years of age. Socioeconomic instability, migration, deteriorating public health programs, frequent vaccine shortages, delayed implementation of vaccination and treatment in response to cases, and lack of public education and awareness were contributing factors.

Significant outbreaks of diphtheria and diphtheria-related mortality continue to be reported from many developing countries, particularly in Africa and Asia. Statistics collected by the World Health Organization indicated the occurrence of ~7000 reported diphtheria cases in 2008 and ~5000 diphtheria deaths in 2004. Although ~82% of the global population has been adequately vaccinated, only 26% of countries have successfully vaccinated >80% of individuals in all districts.

Cutaneous diphtheria is usually a secondary infection that follows a primary skin lesion due to trauma, allergy, or autoimmunity. Most often, these isolates lack the *tox* gene and thus do not express diphtheria toxin. In tropical latitudes, cutaneous diphtheria is more common than respiratory diphtheria. In contrast to respiratory disease, cutaneous diphtheria is not reportable in the United States. Nontoxigenic strains of *C. diphtheriae* have also been associated with pharyngitis in Europe, causing outbreaks among men who have sex with men and persons who use illicit IV drugs.

PATHOGENESIS AND IMMUNOLOGY

Diphtheria toxin produced by *tox*⁺ strains of *C. diphtheriae* is the primary virulence factor in clinical disease. The toxin is synthesized in precursor form; is released as a 535-amino-acid, single-chain protein; and, in sensitive species (e.g., guinea pigs and humans, but not mice or rats), has a 50% lethal dose of ~100 ng/kg of body weight. The toxin is produced in the pseudomembranous lesion and is taken up in the bloodstream, from which it is distributed to all organ systems in the body. Once bound to its cell surface receptor (a heparin-binding epidermal growth factor–like precursor), the toxin is internalized by receptor-mediated endocytosis and enters the cytosol from an acidified early endosomal compartment. In vitro, the toxin may be separated into two chains by digestion with serine proteases: the N-terminal A fragment and the C-terminal B fragment. Delivery of the A fragment into the eukaryotic cell cytosol results in irreversible inhibition of protein synthesis by NAD⁺-dependent ADP-ribosylation of elongation factor 2. The eventual result is the death of the cell.

In 1926, Ramon at the Institut Pasteur found that formalinization of diphtheria toxin resulted in the production of a nontoxic but highly immunogenic diphtheria toxoid. Subsequent studies showed that immunization with diphtheria toxoid elicited antibodies that neutralized the toxin and prevented most disease manifestations. In the 1930s, mass immunization of children and susceptible adults with diphtheria toxoid commenced in the United States and Europe.

Individuals with a diphtheria antitoxin titer of >0.01 U/mL are at low risk of disease. In populations where a majority of individuals have protective antitoxin titers, the carrier rate for toxigenic strains of *C. diphtheriae* decreases and the overall risk of diphtheria among susceptible individuals is reduced. Nevertheless, individuals with nonprotective titers may contract diphtheria through either travel or exposure to individuals who have recently returned from regions where the disease is endemic.

978 Characteristic pathologic findings of diphtheria include mucosal ulcers with a pseudomembranous coating composed of an inner band of fibrin and a luminal band of neutrophils. Initially white and firmly adherent, in advanced diphtheria the pseudomembranes turn gray or even green or black as necrosis progresses. Mucosal ulcers result from toxin-induced necrosis of the epithelium accompanied by edema, hyperemia, and vascular congestion of the submucosal base. A significant fibrinosuppurative exudate from the ulcer develops into the pseudomembrane. Ulcers and pseudomembranes in severe respiratory diphtheria may extend from the pharynx into medium-sized bronchial airways. Expanding and sloughing membranes may result in fatal airway obstruction.

APPROACH TO THE PATIENT: Diphtheria

Diphtheria, though rare in the United States and other developed countries, should be considered when a patient has severe pharyngitis, particularly when there is difficulty swallowing, respiratory compromise, or signs of systemic disease (e.g., myocarditis or generalized weakness). The leading causes of pharyngitis are respiratory viruses (rhinoviruses, influenza viruses, parainfluenza viruses, coronaviruses, adenoviruses; ~25% of cases), group A streptococci (15–30%), group C streptococci (~5%), atypical bacteria such as *Mycoplasma pneumoniae* and *Chlamydia pneumoniae* (15–20% in some series), and other viruses such as herpes simplex virus (~4%) and Epstein-Barr virus (<1% in infectious mononucleosis). Less common causes are acute HIV infection, gonorrhea, fusobacterial infection (e.g., Lemierre's syndrome), thrush due to *Candida albicans* or other *Candida* species, and diphtheria. The presence of a pharyngeal pseudomembrane or an extensive exudate should prompt consideration of diphtheria (Figure 175-1).



FIGURE 175-1 Respiratory diphtheria due to toxigenic *C. diphtheriae* producing exudative pharyngitis in a 47-year-old female patient displaying neck edema and a pseudomembrane extending from the uvula to the pharyngeal wall. The characteristic white pseudomembrane is caused by diphtheria toxin-mediated necrosis of the respiratory epithelial layer, producing a fibrinous coagulable exudate. Submucosal edema adds to airway narrowing. The pharyngitis is acute in onset, and respiratory obstruction from the pseudomembrane may occur in severe cases. Inoculation of pseudomembrane fragments or submembranous swabs onto Löffler's or tellurite selective medium reveals *C. diphtheriae*. (Photograph by P. Strebel, MD, used by permission. From R. Kadirova et al: *J Infect Dis* 181:S110, 2000. With permission of Oxford University Press.)

CLINICAL MANIFESTATIONS

Respiratory Diphtheria The clinical diagnosis of diphtheria is based on the constellation of sore throat; adherent tonsillar, pharyngeal, or nasal pseudomembranous lesions; and low-grade fever. In addition, diagnosis requires the isolation of *C. diphtheriae* or histopathologic isolation of compatible gram-positive organisms. The Centers for Disease Control and Prevention (CDC) recognizes *confirmed* respiratory diphtheria (laboratory proven or epidemiologically linked to a culture-confirmed case) and *probable* respiratory diphtheria (clinically compatible but not laboratory proven or epidemiologically linked). Carriers are defined as individuals who have positive cultures for *C. diphtheriae* and who either are asymptomatic or have symptoms but lack pseudomembranes. Most patients seek medical care for sore throat and fever several days into the illness. Occasionally, weakness, dysphagia, headache, and voice change are the initial manifestations. Neck edema and difficulty breathing are evident in more advanced cases and carry a poor prognosis.

The systemic manifestations of diphtheria stem from the effects of diphtheria toxin and include weakness as a result of neurotoxicity and cardiac arrhythmias or congestive heart failure due to myocarditis. Most commonly, the pseudomembranous lesion is located in the tonsillopharyngeal region. Less commonly, the lesions are located in the larynx, nares, and trachea or bronchial passages. Large pseudomembranes are associated with severe disease and a poor prognosis. A few patients develop massive swelling of the tonsils and present with “bull-neck” diphtheria, which results from massive edema of the submandibular and paratracheal region and is further characterized by foul breath, thick speech, and stridorous breathing. The diphtheritic pseudomembrane is gray or whitish and sharply demarcated. Unlike the exudative lesion associated with streptococcal pharyngitis, the pseudomembrane in diphtheria is tightly adherent to the underlying tissues. Attempts to dislodge the membrane may cause bleeding. Hoarseness suggests laryngeal diphtheria, in which laryngoscopy may be diagnostically helpful.

Cutaneous Diphtheria This dermatosis is characterized by punched-out ulcerative lesions with necrotic sloughing or pseudomembrane formation (Figure 175-2). The diagnosis requires cultivation of *C. diphtheriae* from lesions, which most commonly occur on the lower and upper extremities, head, and trunk.

Infections Due to Non-diphtheriae *Corynebacterium* Species and Nontoxigenic *C. diphtheriae* Non-diphtheriae species of *Corynebacterium* and related genera (discussed below) as well as nontoxigenic strains of *C. diphtheriae* itself have been found in bloodstream and respiratory infections, often in individuals with immunosuppression or chronic respiratory disease. These organisms can cause disease manifestations and should not necessarily be dismissed as colonizers.



FIGURE 175-2 Cutaneous diphtheria due to nontoxigenic *C. diphtheriae* on the lower extremity. (From the Centers for Disease Control and Prevention.)

Other Clinical Manifestations *C. diphtheriae* causes rare cases of endocarditis and septic arthritis, most often in patients with preexisting risk factors, such as abnormal cardiac valves, injection drug use, or cirrhosis.

COMPLICATIONS

Airway obstruction poses a significant early risk in patients presenting with advanced diphtheria. Pseudomembranes may slough and obstruct the airway or may advance to the larynx or into the tracheobronchial tree. Children are particularly prone to obstruction because of their small airways.



Polyneuropathy and myocarditis are late toxic manifestations of diphtheria. During a diphtheria outbreak in the Kyrgyz Republic in 1999, myocarditis was found in 22% and neuropathy in 5% of 656 hospitalized patients. The mortality rate was 7% among patients with myocarditis as opposed to 2% among those without myocardial manifestations. The median time to death in hospitalized patients was 4.5 days. Myocarditis is typically associated with dysrhythmia of the conduction tract and dilated cardiomyopathy.

Polyneuropathy is seen 3–5 weeks after the onset of diphtheria and has a slow indolent course. However, patients may develop severe and prolonged neurologic abnormalities. The disorders typically occur in the mouth and neck, with lingual or facial numbness as well as dysphonia, dysphagia, and cranial nerve paresthesias. More ominous signs include weakness of respiratory and abdominal muscles and paresis of the extremities. Sensory manifestations and sensory ataxia also are observed. Cranial nerve dysfunction typically precedes disturbances of the trunk and extremities because of proximity to the site of infection. Autonomic dysfunction also is associated with polyneuropathy and can lead to hypotension. Polyneuropathy is typically reversible in patients who survive the acute phase.

Other complications of diphtheria include pneumonia, renal failure, encephalitis, cerebral infarction, pulmonary embolism, and serum sickness from antitoxin therapy.

DIAGNOSIS

The diagnosis of diphtheria is based on clinical signs and symptoms plus laboratory confirmation. Respiratory diphtheria should be considered in patients with sore throat, pharyngeal exudates, and fever. Other symptoms may include hoarseness, stridor, or palatal paralysis. The presence of a pseudomembrane should prompt strong consideration of diphtheria. Once a clinical diagnosis of diphtheria is made, diphtheria antitoxin should be obtained and administered as rapidly as possible.

Laboratory diagnosis of diphtheria is based either on cultivation of *C. diphtheriae* or toxigenic *Corynebacterium ulcerans* from the site of infection or on the demonstration of local lesions with characteristic histopathology. *Corynebacterium pseudodiphtheriticum*, a nontoxigenic organism, is a common component of the normal throat flora and does not pose a significant risk. Throat samples should be submitted to the laboratory for culture with the notation that diphtheria is being considered. This information should prompt cultivation on special selective medium and subsequent biochemical testing to differentiate *C. diphtheriae* from other nasopharyngeal commensal corynebacteria. All laboratory isolates of *C. diphtheriae*, including nontoxigenic strains, should be submitted to the CDC.

A diagnosis of cutaneous diphtheria requires laboratory confirmation since the lesions are not characteristic and are indistinguishable from other dermatoses. Diphtheritic ulcers occasionally—but not consistently—have a punched-out appearance (Fig. 175-2). Patients in whom cutaneous diphtheria is identified should have the nasopharynx cultured for *C. diphtheriae*. The laboratory medium for cutaneous diphtheria specimens is the same as that used for respiratory diphtheria: Löffler's or Tinsdale's selective medium in addition to nonselective medium such as blood agar. As has been mentioned, respiratory diphtheria remains a notifiable disease in the United States, whereas cutaneous diphtheria is not.

TREATMENT DIPHTHERIA

DIPHTHERIA ANTITOXIN

Prompt administration of diphtheria antitoxin is critical in the management of respiratory diphtheria. Diphtheria antitoxin, a horse antiserum, is effective in reducing the extent of local disease as well as the risk of complications of myocarditis and neuropathy. Rapid institution of antitoxin therapy is also associated with a significant reduction in mortality risk. Because diphtheria antitoxin cannot neutralize cell-bound toxin, prompt initiation is important. This product, which is no longer commercially available in the United States, can be obtained from the CDC by calling the Bacterial Vaccine Preventable Disease Branch of the National Immunization Program at 404-639-8257 (8:00 A.M. to 4:30 P.M., U.S. Eastern time) or, at other hours, the Emergency Operations Center at 770-488-7100; the relevant website is www.cdc.gov/diphtheria/dat.html. The current protocol for the use of diphtheria antitoxin involves a test dose to rule out immediate hypersensitivity. Patients who demonstrate hypersensitivity require desensitization before a full therapeutic dose of antitoxin is administered.

ANTIMICROBIAL THERAPY

Antibiotics are used in the management of diphtheria primarily to prevent transmission to susceptible contacts. Antibiotics also prevent further toxin production and reduce the severity of local infection. Recommended treatment options for patients with respiratory diphtheria are as follows:

- Procaine penicillin G, 600,000 U IM q12h (for children: 12,500–25,000 U/kg IM q12h) until the patient can swallow comfortably; then oral penicillin V, 125–250 mg qid to complete a 14-day course
- Erythromycin, 500 mg IV q6h (for children: 40–50 mg/kg per day IV in two or four divided doses) until the patient can swallow comfortably; then 500 mg PO qid to complete a 14-day course



A clinical study in Vietnam found that penicillin was associated with a more rapid resolution of fever and a lower rate of bacterial resistance than erythromycin; however, relapses were more common in the penicillin group. Erythromycin therapy targets protein synthesis and thus offers the presumed benefit of stopping toxin synthesis more quickly than a cell wall-active β -lactam agent. Alternative therapeutic agents for patients who are allergic to penicillin or cannot take erythromycin include rifampin and clindamycin. Eradication of *C. diphtheriae* should be documented after antimicrobial therapy is complete. A repeat throat culture 2 weeks later is recommended. For patients in whom the organism is not eradicated after a 14-day course of erythromycin or penicillin, an additional 10-day course followed by repeat culture is recommended. Drug-resistant strains of *C. diphtheriae* exist, and several reports have described multidrug-resistant strains, predominantly in Southeast Asia. Drug resistance should be considered when efforts at pathogen eradication fail.

Cutaneous diphtheria should be treated as described above for respiratory disease. Individuals infected with toxigenic strains should receive antitoxin. It is important to treat the underlying cause of the dermatoses in addition to the superinfection with *C. diphtheriae*.

Patients who recover from respiratory or cutaneous diphtheria should have antitoxin levels measured. If diphtheria antitoxin has been administered, this test should be performed 6 months later. Patients who recover from respiratory or cutaneous diphtheria should receive the appropriate vaccine to ensure the development of protective antibody titers.

MANAGEMENT STRATEGIES

Patients in whom diphtheria is suspected should be hospitalized in respiratory isolation rooms, with close monitoring of cardiac and respiratory function. A cardiac workup is recommended to assess the possibility of myocarditis. In patients with extensive pseudomembranes, an anesthesiology or an ear, nose, and throat consultation is recommended because of the possible need for

980 tracheostomy or intubation. In some settings, pseudomembranes can be removed surgically. Treatment with glucocorticoids has not been shown to reduce the risk of myocarditis or polyneuropathy.

PROGNOSIS

Fatal pseudomembranous diphtheria typically occurs in patients with nonprotective antibody titers and in unimmunized patients. The pseudomembrane may actually increase in size from the time it is first noted. Risk factors for death include bullneck diphtheria; myocarditis with ventricular tachycardia; atrial fibrillation; complete heart block; an age of >60 years or <6 months; alcoholism; extensive pseudomembrane elongation; and laryngeal, tracheal, or bronchial involvement. Another important predictor of fatal outcome is the interval between the onset of local disease and the administration of antitoxin. Cutaneous diphtheria has a low mortality rate and is rarely associated with myocarditis or peripheral neuropathy.

PREVENTION

Vaccination Sustained campaigns for vaccination of children and adequate boosting vaccination of adults are responsible for the exceedingly low incidence of diphtheria in most developed nations. Currently, diphtheria toxoid vaccine is coadministered with tetanus vaccine (with or without acellular pertussis). DTaP (a full-level diphtheria and tetanus toxoids and acellular pertussis vaccine) is currently recommended for children up to the age of 7; DTaP replaced the earlier whole-cell pertussis vaccine DTP in 1997. Tdap is a tetanus toxoid, reduced diphtheria toxoid, and acellular pertussis vaccine formulated for adolescents and adults. Tdap was licensed for use in the United States in 2005 and is the recommended booster vaccine for children 11–12 years old and the recommended catch-up vaccine for children 7–10 and 13–18 years of age. It is recommended that all adults (i.e., persons >19 years old) receive a single dose of Tdap if they have not received it previously, regardless of the interval since the last dose of Td (tetanus and reduced-dose diphtheria toxoids, adsorbed). Tdap vaccination is a priority for health care workers, pregnant women, adults anticipating contact with infants, and adults not previously vaccinated for pertussis. Adults who have received acellular pertussis vaccine should continue to receive decennial Td booster vaccinations. **The vaccine schedule is detailed in Chap. 148.**

Prophylaxis Administration to Contacts Close contacts of diphtheria patients should undergo throat culture to determine whether they are carriers. After samples for throat culture are obtained, antimicrobial prophylaxis should be considered for all contacts, even those whose cultures are negative. The options are 7–10 days of oral erythromycin or one dose of IM benzathine penicillin G (1.2 million units for persons ≥6 years of age or 600,000 units for children <6 years of age).

Contacts of diphtheria patients whose immunization status is uncertain should receive the appropriate diphtheria toxoid-containing vaccine. The Tdap vaccine (rather than Td) is now the booster vaccine of choice for adults who have not recently received an acellular pertussis-containing vaccine. Carriers of *C. diphtheriae* in the community should be treated and vaccinated when identified.

OTHER CORYNEBACTERIAL AND RHODOCOCCLUS INFECTIONS

Nondiphtherial corynebacteria, referred to as *diphtheroids* or *coryneforms*, are frequently considered colonizers or contaminants; however, they have been associated with invasive disease, particularly in immunocompromised patients. These organisms have been isolated from the bloodstream, particularly in association with catheter infection, endocarditis, prosthetic valve infection, meningitis, neurosurgical shunt infection, brain abscess, and peritonitis and often in the setting of chronic ambulatory peritoneal dialysis, osteomyelitis, septic arthritis, urinary tract infection, empyema, and pneumonia, among other infections. Patients infected with these organisms usually have significant medical comorbidity or are immunosuppressed. The nondiphtherial coryneforms are a diverse collection of bacteria that

are taxonomically grouped together in the genus *Corynebacterium* on the basis of their 16S rDNA signature nucleotides. Despite the shared rDNA signatures, these isolates are quite diverse. For example, their guanine-cytosine content ranges from 45% to 70%. Several nondiphtheroid corynebacteria, including *Corynebacterium jeikeium* and *Corynebacterium urealyticum*, are associated with resistance to multiple antibiotics. *Rhodococcus equi* is associated with necrotizing pneumonia and granulomatous infection, particularly in immunocompromised individuals.

MICROBIOLOGY AND LABORATORY DIAGNOSIS

These organisms are non-acid-fast, catalase-positive, aerobic or facultatively anaerobic rods. Their colonial morphologies vary widely; some species are small and α-hemolytic (similar to lactobacilli), whereas others form large white colonies (similar to yeasts). Many nondiphtherial coryneforms require special media, such as Löffler's, Tinsdale's, or telluride medium. These cultivation idiosyncrasies have led to a complex taxonomic categorization of the organisms.

EPIDEMIOLOGY

Humans are the natural reservoirs for several nondiphtherial coryneforms, including *C. xerosis*, *C. pseudodiphtheriticum*, *C. striatum*, *C. minutissimum*, *C. jeikeium*, *C. urealyticum*, and *Arcanobacterium haemolyticum*. Animal reservoirs are responsible for carriage of *Arcanobacterium pyogenes*, *C. ulcerans*, and *C. pseudotuberculosis*. Soil is the natural reservoir for *R. equi*.

C. pseudodiphtheriticum is a component of the normal flora of the human pharynx and skin. *C. xerosis* is found on the skin, nasopharynx, and conjunctiva; *C. auris* in the external auditory canal; and *C. striatum* in the anterior nares and on the skin. *C. jeikeium* and *C. urealyticum* are found in the axilla, groin, and perineum, particularly in hospitalized patients. Infections with *C. ulcerans* and *C. pseudotuberculosis* have been associated with the consumption of raw milk from infected cattle.

C. ulcerans This organism causes a diphtheria-like illness and produces both diphtheria toxin and a dermonecrotic toxin. The organism is a commensal in horses and cattle and has been isolated from cow's milk. *C. ulcerans* causes exudative pharyngitis, primarily during summer months, in rural areas, and among individuals exposed to cattle. In contrast to diphtheria, this infection is considered a zoonosis whose person-to-person transmission has not been documented. Nevertheless, treatment with antitoxin and antibiotics should be initiated when respiratory *C. ulcerans* is identified, and a contact investigation (including throat cultures to determine the need for antimicrobial prophylaxis and, in unimmunized contacts, administration of the appropriate diphtheria toxoid-containing vaccine) should be conducted. The organism grows on Löffler's, Tinsdale's, and telluride agars as well as blood agar. In addition to exudative pharyngitis, cutaneous disease due to *C. ulcerans* has been reported. *C. ulcerans* is susceptible to a wide panel of antibiotics. Erythromycin and macrolides appear to be the first-line agents.

C. pseudotuberculosis (ovis) Infection caused by *C. pseudotuberculosis* is rare and is reported almost exclusively from Australia. *C. pseudotuberculosis* causes suppurative granulomatous lymphadenitis and an eosinophilic pneumonia syndrome among individuals who handle horses, cattle, goats, and deer or who drink raw milk. The organism is an important veterinary pathogen, causing suppurative lymphadenitis, abscesses, and pneumonia, but is rarely a human pathogen. Successful treatment with erythromycin or tetracycline has been reported, with surgery also performed when indicated.

C. jeikeium (Group JK) Originally described in American hospitals, *C. jeikeium* infection was subsequently reported in Europe. After a 1976 survey of diseases caused by nondiphtherial corynebacteria, CDC group JK emerged as an important opportunistic pathogen among neutropenic and HIV-infected patients. The organism has now been designated a separate species. *C. jeikeium* forms small,

gray to white, glistening, nonhemolytic colonies on blood agar. It lacks urease and nitrate reductase and does not ferment most carbohydrates. The predominant syndrome associated with *C. jeikeium* is sepsis with pneumonia, endocarditis, meningitis, osteomyelitis, and epidural abscess. Risk factors for *C. jeikeium* infection include hematologic malignancy, neutropenia from comorbid conditions, prolonged hospitalization, exposure to multiple antibiotics, and skin disruption. There is evidence that *C. jeikeium* is part of the inguinal, axillary, genital, and perirectal flora of hospitalized patients.

Broad-spectrum antimicrobial therapy appears to select for colonization. Gram's staining shows gram-positive coccobacillary forms slightly resembling streptococci. Moreover, *C. jeikeium* is resistant to all antibiotics tested except vancomycin. Effective therapy involves removal of the infectious source, whether a catheter, prosthetic joint, or prosthetic valve. Efforts have been made to prevent *C. jeikeium* infection by improving hygienic conditions for high-risk patients in intensive care settings with antibacterial soap.

***C. urealyticum* (Group D2)** Identified as a urease-positive nondiphtherial *Corynebacterium* in 1972, *C. urealyticum* is an opportunistic pathogen causing sepsis and urinary tract infection. *C. urealyticum* appears to be the etiologic agent of a severe urinary tract syndrome known as *alkaline-encrusted cystitis*, a chronic inflammatory bladder infection associated with deposition of ammonium magnesium phosphate on the surface and walls of ulcerating lesions in the bladder. In addition, *C. urealyticum* has been associated with pneumonia, peritonitis, endocarditis, osteomyelitis, and wound infection. It is similar to *C. jeikeium* in its resistance to most antibiotics except vancomycin. Vancomycin therapy has been used successfully in severe infections.

***C. minutissimum* (Erythrasma)** Erythrasma is a cutaneous infection producing reddish-brown, macular, scaly, pruritic intertriginous patches. The dermatologic presentation under the Wood's lamp is of coral red fluorescence. *C. minutissimum* appears to be a common cause of erythrasma, although there is evidence for a polymicrobial etiology in certain settings. This microbe has also been associated with bacteremia in patients with hematologic malignancy. Erythrasma responds to topical erythromycin, clarithromycin, clindamycin, or fusidic acid, although more severe infections may require oral macrolide therapy.

Other Nondiphtherial Corynebacterial Infections *C. xerosis* is a human commensal found in the conjunctiva, nasopharynx, and skin. This nontoxigenic organism is occasionally identified as a source of invasive infection in immunocompromised or postoperative patients and prosthetic joint recipients. *C. striatum* is found in the anterior nares, skin, face, and upper torso of healthy individuals. Also nontoxigenic, this organism has been associated with invasive opportunistic infections in severely ill or immunocompromised patients. *C. amycolatum* is isolated from human skin and is identified on the basis of a unique 16S ribosomal RNA sequence associated with opportunistic infection. *C. glucuronolyticum* is a nonlipophilic species that causes male genitourinary tract infections such as prostatitis and urethritis. These infections may be successfully treated with a wide variety of antibacterial agents, including β -lactams, rifampin, aminoglycosides, or vancomycin; however, the organism appears to be resistant to fluoroquinolones, macrolides, and tetracyclines. *C. imitans* has been identified in eastern Europe as a nontoxigenic cause of pharyngitis. *C. auris* has been identified in children with otitis media; it is susceptible to fluoroquinolones, rifampin, tetracycline, and vancomycin but resistant to penicillin G and variably susceptible to macrolides. *C. pseudodiphtheriticum* (*C. hoffmanii*) is a nontoxigenic species that is part of the normal human flora. Human infections—particularly endocarditis of either prosthetic or natural valves and invasive pneumonia—have been reported only rarely. Although *C. pseudodiphtheriticum* may be isolated from the nasopharynx of patients with suspected diphtheria,

it is part of the normal flora and does not produce diphtheria toxin. *C. propinquum*, a close relative of *C. pseudodiphtheriticum*, is part of CDC group ANF-3 and is isolated from the human respiratory tract and blood. *C. afermentans* subspecies *lipophilum* belongs to CDC group ANF-1 and has been isolated from human blood and abscesses. *C. accolens* has been isolated from wound drainage, throat swabs, and sputum and is typically identified as a satellite of staphylococcal organisms; this species has been associated with endocarditis. *C. bovis* is a veterinary commensal that has not been clearly associated with human disease. *C. aquaticum* is a water-dwelling organism that is occasionally isolated from patients using medical devices (e.g., for chronic ambulatory peritoneal dialysis or venous access).

Rhodococcus *Rhodococcus* species are phylogenetically related to the corynebacteria. These gram-positive coccobacilli have been associated with tuberculosis-like infections in humans with granulomatous pathology. While *R. equi* is best known, other species have been identified, including *R. (Gordonia) bronchialis*, *R. (Tsukamurella) aurantia-cus*, *R. luteus*, *R. erythropolis*, *R. rhodochrous*, and *R. rubropertinctus*.

R. equi has been recognized as a cause of pneumonia in horses since the 1920s and as a cause of related infections in cattle, sheep, and swine. It is found in soil as an environmental microbe. The organisms vary in length; appear as spherical to long, curved, clubbed rods; and produce large irregular mucoid colonies. *R. equi* cannot ferment carbohydrates or liquefy gelatin and is often acid fast. An intracellular pathogen of macrophages, *R. equi* can cause granulomatous necrosis and caseation. This organism has most commonly been identified in pulmonary infection, but infections of brain, bone, and skin also have been reported. Most commonly, *R. equi* disease manifests as nodular cavitary pneumonia of the upper lobe—a picture similar to that seen in tuberculosis or nocardiosis. Most patients are immunocompromised, often by HIV infection. Subcutaneous nodular lesions have also been identified. The involvement of *R. equi* should be considered when any patient presents with a tuberculosis-like syndrome.

Infection due to *R. equi* has been treated successfully with antibiotics that penetrate intracellularly, including macrolides, clindamycin, rifampin, and trimethoprim-sulfamethoxazole. β -Lactam antibiotics have not been useful. The organism is routinely susceptible to vancomycin, which is considered the drug of choice.

Other Related Species • **ACTINOMYCES PYOGENES** This organism, a well-known pathogen of cattle, sheep, goats, and pigs, causes seasonal leg ulcers in rural Thailand. A few human cases of sepsis, endocarditis, septic arthritis, pneumonia, meningitis, and empyema have been reported. This species is susceptible to β -lactams, tetracyclines, aminoglycosides, and fluoroquinolones.

ARCANOBACTERIUM HAEMOLYTICUM *A. haemolyticum* was identified as an agent of wound infections in U.S. soldiers in the South Pacific during World War II. It appears to be a human commensal of the nasopharynx and skin, but has also been implicated in pharyngitis and chronic skin ulcers. In contrast to the much more common pharyngitis caused by *Streptococcus pyogenes*, *A. haemolyticum* pharyngitis is associated with a scarlatiniform rash on the trunk and proximal extremities in about half of cases; this illness is occasionally confused with toxic shock syndrome. Because *A. haemolyticum* pharyngitis primarily affects teenagers, it has been postulated that the rash-pharyngitis syndrome may represent copathogenicity, synergy, or opportunistic secondary infection with Epstein-Barr virus. *A. haemolyticum* has also been reported as a cause of bacteremia, soft tissue infections, osteomyelitis, and cavitary pneumonia, predominantly in the setting of underlying diabetes mellitus. The organism is susceptible to β -lactams, macrolides, fluoroquinolones, clindamycin, vancomycin, and doxycycline. Penicillin resistance has been reported.

176 *Listeria monocytogenes* Infections

Elizabeth L. Hohmann, Daniel A. Portnoy

Listeria monocytogenes is a food-borne pathogen that can cause serious infections, particularly in pregnant women and immunocompromised individuals. A ubiquitous saprophytic environmental bacterium, *L. monocytogenes* is also a facultative intracellular pathogen with a broad host range. Humans are probably accidental hosts for this microorganism. *L. monocytogenes* is of interest not only to clinicians but also to basic scientists as a model intracellular pathogen that is used to study basic mechanisms of microbial pathogenesis and host immunity.

MICROBIOLOGY

L. monocytogenes is a facultatively anaerobic, nonsporulating, gram-positive rod that grows over a broad temperature range, including refrigeration temperatures. This organism is motile during growth at low temperatures but much less so at 37°C. The vast majority of cases of human listerial disease can be traced to serotypes 1/2a, 1/2b, and 4. *L. monocytogenes* is weakly β -hemolytic on blood agar, and (as detailed below) its β -hemolysin is an essential determinant of its pathogenicity.

PATHOGENESIS

Infections with *L. monocytogenes* follow ingestion of contaminated food that contains the bacteria at high concentrations. The conversion from environmental saprophyte to pathogen involves the coordinate regulation of bacterial determinants of pathogenesis that mediate entry into cells, intracellular growth, and cell-to-cell spread. Many of the organism's pathogenic strategies can be examined experimentally in tissue culture models of infection (Fig. 176-1). Like other enteric pathogens, *L. monocytogenes* induces its own internalization by cells that are not normally phagocytic. Its entry into cells is mediated by host surface proteins classified as internalins. Internalin-mediated entry is important in the crossing of intestinal, blood-brain, and fetoplacental barriers, although how *L. monocytogenes* traffics from the intestine to the brain or fetus is only beginning to be investigated. In a pregnant guinea pig model of infection, *L. monocytogenes* was shown

to traffic from maternal organs to the placenta; surprisingly, however, it also trafficked from the placenta back to maternal organs. These data are consistent with a model in which miscarriage can be viewed as a host defense strategy to eliminate a nidus of infection.

An essential determinant of the pathogenesis of *L. monocytogenes* is its β -hemolysin, listeriolysin O (LLO). LLO is a pore-forming, cholesterol-dependent cytolysin. (Related cytolysins include streptolysin O, pneumolysin, and perfringolysin O, all of which are produced by extracellular pathogens.) LLO is largely responsible for mediating the rupture of the phagosomal membrane that forms after phagocytosis of *L. monocytogenes*. LLO probably acts by insertion into an acidifying phagosome, which prevents the vesicle's maturation. In addition, LLO acts as a translocation pore for one or both of the *L. monocytogenes* phospholipases that also contribute to vacuolar lysis. LLO synthesis and activity are controlled at multiple levels to ensure that its lytic activity is limited to acidic vacuoles and does not affect the cytosol. Mutations in LLO that influence its synthesis, cytosolic half-life, or pH optimum cause premature toxicity to infected cells. There is an inverse relationship between toxicity and virulence—i.e., the more cytotoxic the strain, the less virulent it is in animals. This relationship may seem paradoxical, but, as an intracellular pathogen, *L. monocytogenes* benefits from leaving its host cell unharmed.

Shortly after exposure to the mammalian-cell cytosol, *L. monocytogenes* expresses a surface protein, ActA, that mediates the nucleation of host actin filaments to propel the bacteria intra- and intercellularly. ActA mimics host proteins of the Wiskott-Aldrich syndrome protein (WASP) family by promoting the actin nucleation properties of the Arp2/3 complex. Thus, *L. monocytogenes* can enter the cytosol of almost any eukaryotic cell or cell extract and can exploit a conserved and essential actin-based motility system. Other pathogens as diverse as certain *Shigella*, *Mycobacterium*, *Rickettsia*, and *Burkholderia* species use a related pathogenic strategy that allows cell-to-cell spread without exposure to the extracellular milieu.

IMMUNE RESPONSE

The innate and acquired immune responses to *L. monocytogenes* have been studied extensively in mice. Shortly after IV injection, most bacteria are found in Kupffer cells in the liver, with some organisms in splenic dendritic cells and macrophages. Listeriae that survive the bactericidal activity of initially infected macrophages grow in the cytosol and spread from cell to cell. *L. monocytogenes* triggers three innate immune pathways: a MyD88-dependent pathway leading to inflammatory cytokines, a STING/IRF3 pathway leading to a type I interferon response; and low-level inflammasome activation. Neutrophils are crucial to host defense during the first 24 h of infection, whereas an influx of activated macrophages from the bone marrow is critical subsequently. Mice that survive sublethal infection clear the organisms within a week, with consequent sterile immunity. Studies with knockout mice have been instrumental in dissecting the roles played by chemokines and cytokines during infection. For example, interferon γ , tumor necrosis factor, and CCR2 are essential in controlling infection. While innate immunity is sufficient to control infection, the acquired immune response is required for sterile immunity. Immunity is cell mediated; antibody plays no measurable role. The critical effector cells are cytotoxic (CD8+) T cells that recognize and lyse infected cells; the resulting extracellular bacteria are killed by circulating activated phagocytes.

A hallmark of the *L. monocytogenes* model is that killed vaccines do not provide protective immunity. The explanation for this fundamental observation is multifactorial, involving the generation of appropriate cytokines and the compartmentalization of bacterial proteins for antigen processing and presentation. Because the organism has the capacity to induce a robust cell-mediated immune response, attenuated strains have been engineered to express foreign antigens and are undergoing clinical studies as therapeutic vaccines for cancer.

EPIDEMIOLOGY

L. monocytogenes usually enters the body via the gastrointestinal tract in foods. Listeriosis is most often sporadic, although outbreaks

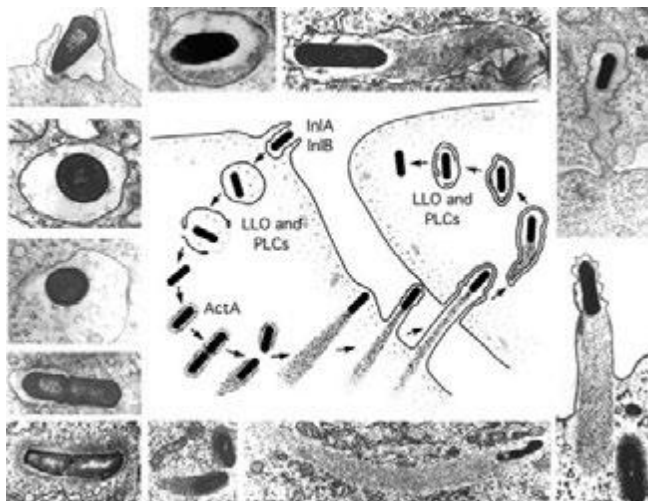


FIGURE 176-1 Stages in the intracellular life cycle of *Listeria monocytogenes*. The central diagram depicts cell entry, escape from a vacuole, actin nucleation, actin-based motility, and cell-to-cell spread. Surrounding the diagram are representative electron micrographs from which it was derived. ActA, surface protein mediating nucleation of host actin filaments to propel bacteria intra- and intercellularly; LLO, listeriolysin O; PLCs, phospholipases C; Inl, internalin. See text for further details. (Adapted with permission from LG Tilney, DA Portnoy: *J Cell Biol* 109:1597, 1989. © Rockefeller University Press.)

do occur. No epidemiologic or clinical evidence supports person-to-person transmission (other than vertical transmission from mother to fetus) or waterborne infection. In line with its survival and multiplication at refrigeration temperatures, *L. monocytogenes* is commonly found in processed and unprocessed foods of animal and plant origin, especially soft cheeses, delicatessen meats, hot dogs, milk, and cold salads; fresh fruits and vegetables can also transmit the organism. Because food supplies are increasingly centralized and normal hosts tolerate the organism well, outbreaks may not be immediately apparent. The U.S. Food and Drug Administration has a zero-tolerance policy for *L. monocytogenes* in ready-to-eat foods.

DIAGNOSIS

Symptoms of listerial infection overlap greatly with those of other infectious diseases. Timely diagnosis requires that the illness be considered in groups at risk: pregnant women; elderly persons; neonates; individuals immunocompromised by organ transplantation, cancer, or treatment with tumor necrosis factor antagonists or glucocorticoids; and patients with a variety of chronic medical conditions, including alcoholism, diabetes, renal disease, and rheumatologic and hepatic illnesses. Meningitis in older adults (especially with parenchymal brain involvement or subcortical brain abscess) should trigger consideration of *L. monocytogenes* infection and treatment. Listeriosis occasionally affects healthy, young, nonpregnant individuals. HIV-infected patients are at risk; however, listeriosis seems to be prevented by trimethoprim-sulfamethoxazole (TMP-SMX) prophylaxis targeting other AIDS-related infections. The diagnosis is typically made by culture of blood, cerebrospinal fluid (CSF), or amniotic fluid. *L. monocytogenes* may be confused with “diphtheroids” or pneumococci in Gram-stained CSF or may be gram-variable and confused with *Haemophilus* species. Polymerase chain reaction diagnostics have been described but are not widely available, and serology is not clinically useful.

CLINICAL MANIFESTATIONS

Listerial infections present as several clinical syndromes, of which meningitis and septicemia are most common. Monocytosis is seen in infected rabbits but is not a hallmark of human infection.

Gastroenteritis Appreciated only since the outbreaks of the late 1980s, listerial gastroenteritis typically develops within 48 h of ingestion of a large inoculum of bacteria in contaminated foods. Attack rates are high (50–100%). *L. monocytogenes* is neither sought nor found in routine fecal cultures, but its involvement should be considered in outbreaks when cultures for other likely pathogens are negative. Sporadic intestinal illness appears to be uncommon. Manifestations include fever, diarrhea, headache, and constitutional symptoms. The largest reported outbreak occurred in an Italian school system and included 1566 individuals; ~20% of patients were hospitalized, but only one person had a positive blood culture. Isolated gastrointestinal illness does not require antibiotic treatment. Surveillance studies show that 0.1–5% of healthy asymptomatic adults may have stool cultures positive for the organism.

Bacteremia *L. monocytogenes* septicemia presents with fever, chills, and myalgias/arthralgias and cannot be differentiated from septicemia involving other organisms. Meningeal symptoms, focal neurologic findings, or mental status changes may suggest the diagnosis. Bacteremia is documented in 70–90% of cancer patients with listeriosis. A nonspecific flulike illness with fever is a common presentation in pregnant women. Endocarditis of prosthetic and native valves is an uncommon complication, with reported fatality rates of 35–50% in case series. A lumbar puncture is often prudent, although not necessary, in pregnant women without central nervous system (CNS) symptoms.

Meningitis *L. monocytogenes* causes ~5–10% of all cases of community-acquired bacterial meningitis in adults in the United States. Case-fatality rates are reported to be 15–26% and do not appear to have changed over time. This diagnosis should be considered in all older

or chronically ill adults with “aseptic” meningitis. The presentation is more frequently subacute (with illness developing over several days) than in meningitis of other bacterial etiologies, and nuchal rigidity and meningeal signs are less common. Photophobia is infrequent. Focal findings and seizures are common in some but not all series. The CSF profile in listerial meningitis most often shows white blood cell counts in the range of 100–5000/μL (rarely higher); 75% of patients have counts below 1000/μL, usually with a neutrophil predominance more modest than that in other bacterial meningitides. Low glucose levels and positive results on Gram’s staining are found ~30–40% of the time. Hydrocephalus can occur.

Meningoencephalitis and Focal CNS Infection *L. monocytogenes* can directly invade the brain parenchyma, producing either cerebritis or focal abscess. Approximately 10% of cases of CNS infection are macroscopic abscesses resulting from bacteremic seeding; the affected patients often have positive blood cultures. Concurrent meningitis can exist, but the CSF may appear normal. Abscesses can be misdiagnosed as metastatic or primary tumors and, in rare instances, occur in the cerebellum and the spinal cord. Invasion of the brainstem results in a characteristic severe rhombencephalitis, usually in otherwise healthy older adults (although there are numerous other infectious and noninfectious causes of this syndrome). The presentation may be biphasic, with a prodrome of fever and headache followed by asymmetric cranial nerve deficits, cerebellar signs, and hemiparetic and hemisensory deficits. Respiratory failure can occur. The subacute course and the often minimally abnormal CSF findings may delay the diagnosis, which may be suggested by MRI showing ring-enhancing lesions after gadolinium contrast and hyperintense lesions on diffusion-weighted imaging. MRI is superior to CT for the diagnosis of these infections.

Infection in Pregnant Women and Neonates Listeriosis in pregnancy is a severe and important infection. The usual presentation is a nonspecific acute or subacute febrile illness with myalgias, arthralgias, backache, and headache. Pregnant women with listeriosis are usually bacteremic. This syndrome should prompt blood cultures, especially if there is no other reasonable explanation. Involvement of the CNS is rare in the absence of other risk factors. Preterm delivery is a common complication, and the diagnosis may be made only post-partum. As many as 70–90% of fetuses from infected women can become infected. Prepartum treatment of bacteremic women enhances the chances of delivery of a healthy infant. Women usually do well after delivery: maternal deaths are very rare, even when the diagnosis is made late in pregnancy or post-partum. Overall mortality rates for fetuses infected in utero approach 50% in some series; among live-born neonates treated with antibiotics, mortality rates are much lower (~20%). *Granulomatosis infantiseptica* is an overwhelming listerial fetal infection with miliary microabscesses and granulomas, most often in the skin, liver, and spleen. Less severe neonatal infection acquired in utero presents at birth. “Late-onset” neonatal illness typically develops ~10–30 days post-partum. Mothers of infants with late-onset disease are not ill.

TREATMENT

INFECTIONS CAUSED BY LISTERIA MONOCYTOGENES

ANTIBIOTICS

No clinical trials have compared antimicrobial agents for the treatment of *L. monocytogenes* infections. Data from studies conducted in vitro and in animals as well as observational clinical data indicate that ampicillin is the drug of choice, although penicillin also is highly active. Adults should receive IV ampicillin at high doses (2 g every 6 h). Many experts recommend the addition of gentamicin for synergy (1.0–1.7 mg/kg every 8 h); retrospective uncontrolled trials are not conclusive, but one study suggests that gentamicin may not help. TMP-SMX, given IV, is the best alternative for the penicillin-allergic patient (15–20 mg of TMP/kg per day in divided doses every 6–8 h). The dosages recommended cover CNS infection and

984 bacteremia (see below for duration); dosages must be reduced for patients with renal insufficiency. One small nonrandomized study supports a combination of ampicillin and TMP-SMX. Case reports document success with vancomycin, imipenem, meropenem, linezolid, tetracycline, and macrolides, although there are also reports of clinical failure or disease development with some of these agents. Acquired resistance to antimicrobial agents has been sought but not found in large strain collections. Cephalosporins are *not* effective and should not be used. Neonates should receive ampicillin and gentamicin at doses based on weight.

DURATION

The duration of therapy depends on the syndrome: 2 weeks for bacteremia, 3 weeks for meningitis, 6–8 weeks for brain abscess/encephalitis, and 4–6 weeks for endocarditis in both neonates and adults. Early-onset neonatal disease may be more severe and should be treated for >2 weeks.

COMPLICATIONS AND PROGNOSIS

Many individuals who are promptly diagnosed and treated recover fully, but permanent neurologic sequelae are common in patients with brain abscess or rhombencephalitis. Focal infections of visceral organs; the eye; the pleural, peritoneal, and pericardial spaces; the bones; and both native and prosthetic joints have all been reported. Of 100 live-born, treated neonates in one series, 60% recovered fully, 24% died, and 13% had long-term neurologic or other complications.

PREVENTION

Healthy persons should take standard precautions to prevent food-borne illness: fully cooking meats, washing fresh vegetables, carefully cleaning utensils, and avoiding unpasteurized dairy products. In addition, persons at risk for listeriosis, including pregnant women, should avoid soft cheeses (hard cheeses and yogurt are not problematic) and should avoid or thoroughly reheat ready-to-eat and delicatessen foods.

Maternal tetanus is defined by the WHO as tetanus occurring during pregnancy or within 6 weeks after the conclusion of pregnancy (whether with birth, miscarriage, or abortion).

ETIOLOGY

C. tetani is an anaerobic, gram-positive, spore-forming rod whose spores are highly resilient and can survive readily in the environment throughout the world. Spores resist boiling and many disinfectants. In addition, *C. tetani* spores and bacilli survive in the intestinal systems of many animals, and fecal carriage is common. The spores or bacteria enter the body through abrasions, wounds, or (in the case of neonates) the umbilical stump. Once in a suitable anaerobic environment, the organisms grow, multiply, and release tetanus toxin, an exotoxin that enters the nervous system and causes disease. Very low concentrations of this highly potent toxin can result in tetanus (minimum lethal human dose, 2.5 ng/kg).

In 20–30% of cases of tetanus, no puncture entry wound is found. Superficial abrasions to the limbs are the commonest infection sites in adults. Deeper infections (e.g., attributable to open fracture, abortion, or drug injection) are associated with more severe disease and worse outcomes. In neonates, infection of the umbilical stump can result from inadequate umbilical cord care; in some cultures, for example, the cord is cut with grass or animal dung is applied to the stump. Circumcision or ear-piercing also can result in neonatal tetanus.

EPIDEMIOLOGY

Tetanus is a rare disease in the developed world. Two cases of neonatal tetanus have occurred in the United States since 1989. Between 2001 and 2008, a total of 231 cases of tetanus were reported to the U.S. national surveillance system. Most cases occur in incompletely vaccinated or unvaccinated individuals. Vaccination status is known in 50% of cases reported in the United States between 1972 and 2009; among these cases, only 16% of patients had had three or more doses of tetanus toxoid.

Persons >60 years of age are at greater risk of tetanus because antibody levels decrease over time. One-third of recent cases in the United States were in persons >65 years old. Injection drug users—particularly those injecting heroin subcutaneously (“skin-popping”)—are increasingly recognized as a high-risk group (15% of all cases in 2001–2008). In 2004, an outbreak of tetanus occurred in the United Kingdom, which had previously reported low rates among drug users. The reasons for this outbreak remain unclear but are thought to involve a combination of heroin contamination, skin-popping, and incomplete vaccination. Since then, only seven sporadic cases have been reported in the United Kingdom.

PATHOGENESIS

Genome sequencing of *C. tetani* has allowed identification of several exotoxins and virulence factors. Only those bacteria producing tetanus toxin (*tetanospasm*) can cause tetanus. Although closely related to the botulinum toxins in structure and mode of action, tetanus toxin undergoes retrograde transport into the central nervous system and thus produces clinical effects different from those caused by the botulinum toxins, which remain at the neuromuscular junction.

Toxin is transported by intra-axonal transport to motor nuclei of the cranial nerves or ventral horns of the spinal cord. Tetanus toxin is produced as a single 150-kDa protein that is cleaved to produce heavy (100-kDa) and light (50-kDa) chains linked by a disulfide bond and noncovalent forces. The carboxy terminal of the heavy chain binds to specific membrane components in presynaptic α -motor nerve terminals; evidence suggests binding to both polysialogangliosides and membrane proteins. This binding results in toxin internalization and uptake into the nerves. Once inside the neuron, the toxin enters a retrograde transport pathway, whereby it is transported proximally to the motor neuron body in what appears to be a highly specific process. Unlike other components of the endosomal contents, which undergo acidification following internalization, tetanus toxin is transported in a carefully regulated pH-neutral environment that prevents an acid-induced conformational change that would result in light-chain expulsion into the surrounding cytosol.

177 Tetanus

C. Louise Thwaites, Lam Minh Yen



Tetanus is an acute disease manifested by skeletal muscle spasm and autonomic nervous system disturbance. It is caused by a powerful neurotoxin produced by the bacterium *Clostridium tetani* and is completely preventable by vaccination. *C. tetani* is found throughout the world, and tetanus commonly occurs where the vaccination coverage rate is low. In developed countries, the disease is seen occasionally in individuals who are incompletely vaccinated. In any setting, established tetanus is a severe disease with a high mortality rate.

DEFINITION

Tetanus is diagnosed on clinical grounds (sometimes with supportive laboratory confirmation of the presence of *C. tetani*; see “Diagnosis,” below), and case definitions are often used to facilitate clinical and epidemiologic assessments. The Centers for Disease Control and Prevention (CDC) defines tetanus as “the acute onset of hypertonia or ... painful muscular contractions (usually of the muscles of the jaw and neck) and generalized muscle spasms without other apparent medical cause.” *Neonatal* tetanus is defined by the World Health Organization (WHO) as “an illness occurring in a child who has the normal ability to suck and cry in the first 2 days of life but who loses this ability between days 3 and 28 of life and becomes rigid and has spasms.” Given the unique presentation of neonatal tetanus, the history generally permits accurate classification of the illness with a high degree of probability.

The next stage in toxin trafficking is less clearly understood but involves tetanus toxin's escaping normal lysosomal degradation processes and undergoing translocation across the synapse to the GABA-ergic presynaptic inhibitory interneuron terminals. Here the light chain, which is a zinc-dependent endopeptidase, cleaves vesicle-associated membrane protein 2 (VAMP2, also known as *synaptobrevin*). This molecule is necessary for presynaptic binding and release of neurotransmitter; thus tetanus toxin prevents transmitter release and effectively blocks inhibitory interneuron discharge. The result is unregulated activity in the motor nervous system. Similar activity in the autonomic system accounts for the characteristic features of skeletal muscle spasm and autonomic system disturbance. The increased circulating catecholamine levels in severe tetanus are associated with cardiovascular complications.

Relatively little is known about the processes of recovery from tetanus. Recovery can take several weeks. Peripheral nerve sprouting is involved in recovery from botulism, and similar central nervous system sprouting may occur in tetanus. Other evidence suggests toxin degradation as a mechanism of recovery.

APPROACH TO THE PATIENT:

Tetanus

The clinical manifestations of tetanus occur only after tetanus toxin has reached presynaptic inhibitory nerves. Once these effects become apparent, there may be little that can be done to affect disease progression. Treatment should not be delayed while the results of laboratory tests are awaited. Management strategies aim to neutralize remaining unbound toxin and support vital functions until the effects of the toxin have worn off. Recent interest has focused on intrathecal methods of antitoxin administration to neutralize toxin within the central nervous system and limit disease progression (see "Treatment," below).

CLINICAL MANIFESTATIONS

Tetanus produces a wide spectrum of clinical features that are broadly divided into generalized (including neonatal) and local. In the usually mild form of local tetanus, only isolated areas of the body are affected

and only small areas of local muscle spasm may be apparent. If the cranial nerves are involved in localized cephalic tetanus, the pharyngeal or laryngeal muscles may spasm, with consequent aspiration or airway obstruction, and the prognosis may be poor. In the typical progression of generalized tetanus (Fig. 177-1), muscles of the face and jaw often are affected first, presumably because of the shorter distances toxin must travel up motor nerves to reach presynaptic terminals. Neonates typically present with inability to suck.

In assessing prognosis, the speed at which tetanus develops is important. The incubation period (time from wound to first symptom) and the period of onset (time from first symptom to first generalized spasm) are of particular significance; shorter times are associated with worse outcome. In neonatal tetanus, the younger the infant is when symptoms occur, the worse the prognosis.

The commonest initial symptoms are trismus (lockjaw), muscle pain and stiffness, back pain, and difficulty swallowing. In neonates, difficulty in feeding is the usual presentation. As the disease progresses, muscle spasm develops. Generalized muscle spasm can be very painful. Commonly, the laryngeal muscles are involved early or even in isolation. This is a life-threatening event as complete airway obstruction may ensue. Spasm of the respiratory muscles results in respiratory failure. Without ventilatory support, respiratory failure is the commonest cause of death in tetanus. Spasms strong enough to produce tendon avulsions and crush fractures have been reported, but this outcome is rare.

Autonomic disturbance is maximal during the second week of severe tetanus, and death due to cardiovascular events becomes the major risk. Blood pressure is usually labile, with rapid fluctuations from high to low accompanied by tachycardia. Episodes of bradycardia and heart block can also occur. Autonomic involvement is evidenced by gastrointestinal stasis, sweating, increased tracheal secretions, and acute (often high-output) renal failure.

DIAGNOSIS

The diagnosis of tetanus is based on clinical findings. As stated above, treatment should not be delayed while laboratory tests are conducted. Culture of *C. tetani* from a wound provides supportive evidence. Serum anti-tetanus immunoglobulin G may also be measured in a sample taken before the administration of antitoxin or immunoglobulin. Serum levels >0.1 IU/mL are deemed protective and do not support the diagnosis of tetanus. If levels are below this threshold, a

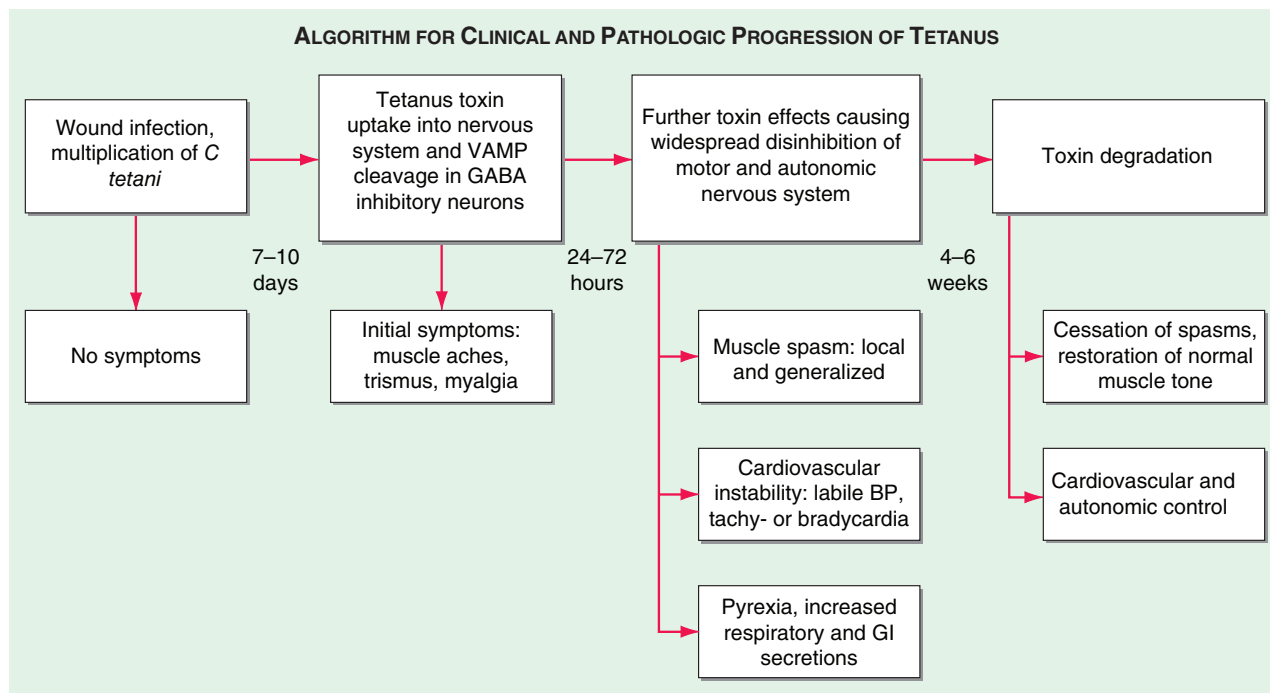


FIGURE 177-1 Clinical and pathologic progression of tetanus. BP, blood pressure; GABA, γ -aminobutyric acid; GI, gastrointestinal; VAMP, vesicle-associated membrane protein (synaptobrevin).

986 bioassay for serum tetanus toxin may be helpful, but a negative result does not exclude the diagnosis. Polymerase chain reaction also has been used for detection of tetanus toxin, but its sensitivity is unknown.

The few conditions that mimic generalized tetanus include strychnine poisoning and dystonic reactions to antidopaminergic drugs. Abdominal muscle rigidity is characteristically continuous in tetanus but is episodic in the latter two conditions. Cephalic tetanus can be confused with other causes of trismus, such as oropharyngeal infection. Hypocalcemia and meningoencephalitis are included in the differential diagnosis of neonatal tetanus.

TREATMENT TETANUS

If possible, the entry wound should be identified, cleaned, and debrided of necrotic material in order to remove anaerobic foci of infection and prevent further toxin production. Metronidazole (400 mg rectally or 500 mg IV every 6 h for 7 days) is the preferred antibiotic. An alternative is penicillin (100,000–200,000 IU/kg per day), although this drug theoretically may exacerbate spasms. Failure to remove pockets of ongoing infection may result in recurrent or prolonged tetanus.

Antitoxin should be given early in an attempt to deactivate any circulating tetanus toxin and prevent its uptake into the nervous system. Two preparations are available: human tetanus immune globulin (TIG) and equine antitoxin. TIG is the preparation of choice, as it is less likely to be associated with anaphylactoid reactions. Recommended therapy is 3000–5000 IU of TIG as a single IM dose, a portion of which should be injected around the wound. Equine-derived antitoxin is available widely and is used in low-income countries at a dosage of 10,000–20,000 U administered IM as a single dose or as divided doses after testing for hypersensitivity. Some evidence indicates that intrathecal administration of TIG inhibits disease progression and leads to a better outcome. The results of relevant studies have been supported by a meta-analysis of trials involving both adults and neonates, with TIG doses of 50–1500 IU administered intrathecally.

Spasms are controlled by heavy sedation with benzodiazepines. Chlorpromazine and phenobarbital are commonly used worldwide, and IV magnesium sulfate has been used as a muscle relaxant. A significant problem with all these treatments is that the doses necessary to control spasms also cause respiratory depression; thus, in resource-limited settings without mechanical ventilators, controlling spasms while maintaining adequate ventilation is problematic, and respiratory failure is a common cause of death. In locations with ventilation equipment, severe spasms are best controlled with a combination of sedatives or magnesium and relatively short-acting, cardiovascularly inert, nondepolarizing neuromuscular blocking agents that allow titration against spasm intensity. Infusions of propofol have also been used successfully to control spasms and provide sedation.

It is important to establish a secure airway early in severe tetanus. Ideally, patients should be nursed in calm, quiet environments because light and noise can trigger spasms. Tracheal secretions are increased in tetanus, and dysphagia due to pharyngeal involvement combined with hyperactivity of laryngeal muscles makes endotracheal intubation difficult. Patients may need ventilator support for several weeks. Thus tracheostomy is the usual method of securing the airway in severe tetanus.

Cardiovascular instability in severe tetanus is notoriously difficult to treat. Rapid fluctuations in blood pressure and heart rate can occur. Cardiovascular stability is improved by increasing sedation with IV magnesium sulfate (plasma concentration, 2–4 mmol/L or titrated against disappearance of the patella reflex), morphine, or other sedatives. In addition, drugs acting specifically on the cardiovascular system (e.g., esmolol, calcium antagonists, and inotropes) may be required. Short-acting drugs that allow rapid titration are preferred; particular care should be taken when longer-acting β antagonists are administered, as their use has been associated with hypotensive cardiac arrest.

Complications arising from treatment are common and include thrombophlebitis associated with diazepam injection, ventilator-associated pneumonia, central-line infections, and septicemia. In some centers, prophylaxis against deep-vein thrombosis and thromboembolism is routine.

Recovery from tetanus may take 4–6 weeks. Patients must be given a full primary course of immunization, as tetanus toxin is poorly immunogenic and the immune response following natural infection is inadequate.

PROGNOSIS

Rapid development of tetanus is associated with more severe disease and poorer outcome; it is important to note time of onset and length of incubation period. More sophisticated modeling has revealed other important predictors of prognosis (Table 177-1). Few studies have formally addressed long-term outcomes of tetanus. However, it is generally accepted that recovery is typically complete unless periods of hypoventilation have been prolonged or other complications have ensued. Studies of children and neonates have suggested a higher incidence of neurologic sequelae. Neonates may be at increased risk of learning disabilities, behavioral problems, cerebral palsy, and deafness.

PREVENTION

Tetanus is prevented by good wound care and immunization (Chap. 148). In neonates, use of safe, clean delivery and cord-care practices as well as maternal vaccination are essential. The WHO guidelines for tetanus vaccination consist of a primary course of three doses in infancy, boosters at 4–7 and 12–15 years of age, and one booster in adulthood. In the United States, the CDC suggests an additional dose at 14–16 months and boosters every 10 years. “Catch-up” schedules recommend a three-dose primary course for unimmunized adolescents followed by two further doses. For persons who have received a complete primary course in childhood but no further boosters, two doses at least 4 weeks apart are recommended.

Standard WHO recommendations for prevention of maternal and neonatal tetanus call for administration of two doses of tetanus toxoid at least 4 weeks apart to previously unimmunized pregnant women. However, in high-risk areas, a more intensive approach has been successful, with all women of childbearing age receiving a primary course along with education on safe delivery and postnatal practices.

Individuals sustaining tetanus-prone wounds should be immunized if their vaccination status is incomplete or unknown or if their last booster was given >10 years earlier. Patients sustaining wounds not classified as clean or minor should also undergo passive immunization with TIG. It is recommended that tetanus toxoid be given in conjunction with diphtheria toxoid in a preparation with or without acellular pertussis: DTaP for children <7 years old, Td for 7- to 9-year olds, and Tdap for children >9 years old and adults.

In the early 1980s, tetanus caused more than 1 million deaths a year, accounting for an estimated 5% of maternal deaths and 14% of all neonatal deaths. In 1989, the World Health



TABLE 177-1 FACTORS ASSOCIATED WITH A POOR PROGNOSIS IN TETANUS

Adult Tetanus	Neonatal Tetanus
Age >70 years	Younger age, premature birth
Incubation period <7 days	Incubation period <6 days
Short time from first symptom to admission	Delay in hospital admission
Puerperal, IV, postsurgery, burn entry site	Grass used to cut cord
Period of onset ^a <48 h	Low birth weight
Heart rate >140 beats/min ^b	Fever on admission
Systolic blood pressure >140 mmHg ^b	
Severe disease or spasms ^b	
Temperature >38.5°C ^b	

^aTime from first symptom to first generalized spasm. ^bAt hospital admission.

Assembly adopted a resolution to eliminate neonatal tetanus by the year 2000; elimination was defined as <1 case/1000 live births in every district in every country. By 1999, elimination was still to be achieved in 57 countries and the deadline was extended until 2005, with the additional target of eliminating maternal tetanus (tetanus occurring during pregnancy or within 6 weeks of its end). Ratification of the Millennium Development Goals, in particular goal 4 (achieving a two-thirds reduction in the mortality rate among children under 5 by 2015), has further focused attention on reducing deaths from vaccine-preventable disease, particularly in the first 4 weeks of life.

Because vaccination reduces the incidence of neonatal tetanus by an estimated 94%, immunization of pregnant women with two doses of tetanus toxoid at least 4 weeks apart has been the primary method of maternal and neonatal tetanus elimination. In some areas, all women of childbearing age have been targeted as a means of increasing vaccination coverage. In addition, educational programs have focused on improving hygiene during the birth process, an intervention that in itself is estimated to reduce neonatal tetanus deaths by up to 40%.

The latest available data show that 34 countries have yet to eliminate maternal and neonatal tetanus, although incidence has declined significantly. Worldwide, deaths from neonatal tetanus fell by 92% between 1990 and 2008; in the latter year, with 84% of newborns protected from the disease by maternal vaccination, there were an estimated 59,000 neonatal tetanus deaths. Despite this relative success, immunization programs need to be ongoing as there is no tetanus herd immunity effect and *C. tetani* contamination of soil and feces is widespread.

178 Botulism

Susan Maslanka, Agam K. Rao

Botulism, recognized at least since the eighteenth century, is a neuroparalytic disease caused by botulinum toxin, one of the most toxic substances known. While initially thought to be caused only by the ingestion of botulinum toxin in contaminated food (food-borne botulism), three additional forms caused by in situ toxin production after germination of spores in either a wound or the intestine are now recognized worldwide: wound botulism, infant botulism, and adult intestinal colonization botulism. In addition to occurring in these recognized natural forms of the disease, botulism symptoms have been reported in patients receiving injections of botulinum toxin for cosmetic or therapeutic purposes (iatrogenic botulism). Moreover, botulism was reported after inhalation of botulinum toxin in a laboratory setting. All forms of botulism manifest as a relatively distinct clinical syndrome of symmetric cranial-nerve palsies followed by descending bilateral flaccid paralysis of voluntary muscles, which may progress to respiratory compromise and death. The mainstays of therapy are meticulous intensive care and treatment with antitoxin as soon as botulism is suspected and before other illnesses have been ruled out.

ETIOLOGY AND PATHOGENESIS

Seven serologically distinct confirmed serotypes of botulinum toxin (A through G) have been confirmed. Botulinum toxin is produced by four recognized species of clostridia: *Clostridium botulinum*, *Clostridium argentinense*, *Clostridium baratii*, and *Clostridium butyricum*. Certain strains may produce more than one serotype. All are anaerobic gram-positive organisms that form subterminal spores; *C. botulinum* and *C. argentinense* spores have been recovered from the environment. The spores survive environmental conditions and ordinary cooking procedures. Toxin production, however, requires a rare confluence of product storage conditions: an anaerobic environment, a pH of >4.6, low salt and sugar concentrations, and temperatures of >4°C. Although commonly ingested, spores do not normally germinate and produce toxin in the adult human intestine.

Food-borne botulism is caused by consumption of foods contaminated with botulinum toxin; no confirmed host-specific factors are involved in the disease. *Wound botulism* is caused by contamination of wounds with *C. botulinum* spores, subsequent spore germination, and toxin production in the anaerobic milieu of an abscess or a wound. *Infant botulism* results from absorption of toxin produced in situ by toxigenic clostridia colonizing the intestine of children ≤1 year of age. Colonization is thought to occur because the normal bowel microbiota is not yet fully established; this theory is supported by studies in animals. *Adult intestinal colonization botulism*, a very rare form that is poorly understood, has a pathology similar to that of infant botulism but occurs in adults; typically, patients have some anatomic or functional bowel abnormality or have recently used antibiotics that may help toxigenic clostridia compete more successfully against the normal bowel microbiota. Despite antitoxin treatment, relapse due to intermittent intraluminal production of toxin may be observed in patients with adult intestinal colonization botulism.

Regardless of how exposure occurs, botulinum neurotoxin enters the vascular system and is transported to peripheral cholinergic nerve terminals, including neuromuscular junctions, postganglionic parasympathetic nerve endings, and peripheral ganglia. Botulinum toxin is a zinc-endopeptidase protein of ~150 kDa, consisting of a 100-kDa heavy chain and a 50-kDa light chain. Steps in neurotoxin activity include (1) heavy-chain binding to nerve terminals, (2) internalization in endocytic vesicles, (3) translocation of the light chain to cytosol, and (4) light-chain serotype-specific cleavage of one of several proteins involved in the release of the neurotransmitter acetylcholine. Inhibition of acetylcholine release by any of the seven toxin serotypes results in characteristic flaccid paralysis. Recovery follows sprouting of new nerve terminals.

All botulinum toxin serotypes have been demonstrated to cause botulism in nonhuman primates. Human cases associated with serotypes A, B, E, and F are reported each year. Serotype A produces the most severe syndrome, with the greatest proportion of patients requiring mechanical ventilation. Serotype B appears to cause milder disease than type A in both food-borne and infant botulism. Serotype E, most often associated with foods of aquatic origin, produces a syndrome of variable severity. The rare cases of illness caused by toxin serotype F, whether in infants or adults, are characterized by rapid progression to quadriplegia and respiratory failure but also by relatively rapid recovery. Recent studies have shown that at least some serotypes can be differentiated into subtypes through neurotoxin gene sequencing; however, the impact of these subtype differences on clinical illness is not yet known.

EPIDEMIOLOGY



Botulism occurs worldwide, but the number of cases reported varies among countries and regions. The variation may be due not only to actual differences in incidence but also to (1) availability of resources to identify botulism, a rare disease; (2) differences in reporting requirements; and (3) limited external access to data collections. There is no universal surveillance system to capture worldwide botulism incidence. However, 30 countries currently participate in voluntary reporting of botulism cases to the European Union through an established surveillance system that includes standardized case definitions similar to those used in the United States and Canada. Other countries (e.g., Argentina, China, Thailand, Japan) maintain independent botulism surveillance.

Food-Borne Botulism From 1899 to 2011, 1225 food-borne botulism events (single cases or outbreaks) were reported in the United States; from 1990 to 2000, a median of 23 cases were reported annually. Most such events (~80%) involve vegetables or fish/aquatic animals, usually home-preserved (canned, jarred). Native communities in both the United States (Alaska) and Canada have a high incidence of food-borne botulism due to traditional food-preparation practices; 85% of all cases in Canada occur in Native communities. Outbreaks typically involve two or three cases; however, one restaurant-associated outbreak in 1977 affected 59 persons. Worldwide, the highest incidence rate is reported from Georgia and Armenia in the southern Caucasus region, where illness is also associated with home-canning practices. Outbreaks in Asia are

988 attributable to consumption of home-preserved fish or vegetable products such as bean curd and bamboo shoots. In parts of Europe, including Poland, France, and Germany, illness is often associated with home-preserved meat such as ham or sausage. Since 1950, commercial products have rarely been implicated in botulism in the United States, and botulism from commercial products is most often attributed to consumer error in storage or cooking. However, manufacturer deficiencies do occur. In 2007, botulism developed in eight persons in the United States who consumed a commercially canned hot-dog chili sauce. Significant deficiencies discovered by regulatory authorities involved 91 different products and resulted in the recall of 111 million cans of food.

Wound Botulism This form of disease was first recognized in 1951 as a result of a review of the clinical records on an accidental injury in 1943. Between 1943 and 2011, 491 cases of wound botulism were reported in the United States; 97% of cases reported after 1990 were associated with injection drug use. The typical patient was a 30- to 50-year-old resident of the western United States with a long history of black-tar heroin injection. In the early 2000s, wound botulism associated with drug use emerged in Europe, and at least two case clusters have been reported.

Infant Botulism More than 3900 infant botulism cases have been reported worldwide (84% in the United States alone) since this form of the disease was first recognized in 1976; ~80–100 cases (commonly caused by serotypes A and B) are reported annually in the United States.

Adult Intestinal Colonization Botulism This form of botulism is difficult to confirm because it is poorly understood and because no clear guidelines are available to help differentiate it from other adult botulism cases. Often these cases are caused by *C. baratii* type F, but the involvement of both *C. botulinum* type A and *C. butyricum* type E has been reported.

Iatrogenic Botulism Paralysis of variable severity has followed injection of licensed botulinum toxin products for treatment of conditions involving hypertonicity of large muscle groups. The U.S. Food and Drug Administration received 658 reports of adverse events related to botulinum toxin use—some very serious—between 1997 and 2006. Although some patients had symptoms consistent with botulism, no cases were laboratory confirmed. Injection of approved doses of licensed products for cosmetic purposes has not been associated with botulism. However, four cases of laboratory-confirmed botulism resulted from illegal injection of research-grade toxin for cosmetic purposes in a U.S. medical facility in 2004.

Inhalational Botulism Inhalational botulism does not occur naturally. One report from Germany has described botulism resulting from possible inhalational exposure to botulinum toxin in a laboratory incident.

Intentional Botulism Botulinum toxin has been “weaponized” by governments and terrorist organizations. An attack might use aerosolization of toxin or contamination of foods or beverages ranging in scope from small-scale tampering to contamination of a widely distributed food item. An unnatural event may be suggested by unusual relationships between patients (e.g., a visit to the same building), atypical exposure vehicles, or atypical toxin serotypes.

CLINICAL MANIFESTATIONS

The distinctive clinical syndrome of botulism consists of symmetric cranial-nerve palsies followed by bilateral descending flaccid paralysis that may progress to respiratory failure and death. The incubation period from ingestion of contaminated food to onset of symptoms in food-borne botulism is usually 8–36 h but can be as long as 10 days and is dose dependent. Incubation periods of 4–17 days have been documented in wound botulism associated with accidental injury. However, estimation is difficult in cases involving injection drug users because most of these patients inject drugs several times daily. Similarly, the incubation period for infant botulism has not been established, but the fact that the illness has affected infants <3 days old suggests that this interval may be very short.

Cranial nerve deficits may include some of the following: diplopia, dysarthria, dysphonia, ptosis, ophthalmoplegia, facial paralysis, and impaired gag reflex. Pupillary reflexes may be depressed, and fixed or dilated pupils are sometimes noted. Autonomic symptoms such as dizziness, dry mouth, and very dry, occasionally sore throat are common. Constipation due to paralytic ileus is nearly universal, and urinary retention is also common. Patients are afebrile and remain alert and oriented. Respiratory failure may occur due to either paralysis of the diaphragm and accessory breathing muscles or pharyngeal collapse secondary to cranial nerve paralysis. Weakness descends, often rapidly, from the head to involve the neck, arms, thorax, and legs; occasionally, weakness is asymmetric. Deep tendon reflexes may be normal or may progressively disappear. Paresthesias, while rare, have been reported and may represent secondary nerve compression from immobility due to paralysis. Absence of cranial nerve palsies or their onset after the appearance of other true neurologic symptoms makes botulism highly unlikely. Nausea, vomiting, and abdominal pain may precede or follow the onset of paralysis in food-borne botulism. Infants with botulism typically present with reduced ability to suck and swallow, constipation, weakened voice, ptosis, sluggish pupils, hypotonia, and floppy neck; as in adults, illness can progress to generalized flaccidity and respiratory compromise.

Even when intubated, patients can respond to questions by moving their fingers or toes unless paralysis has affected the digits. In some instances, unfortunately, the severe ptosis, expressionless face, and weak phonation of patients with botulism have been interpreted as signs of mental status changes due to alcohol intoxication, drug overdose, encephalitis, or meningitis—a conclusion that delays an accurate diagnosis. Because of skeletal muscle paralysis, patients experiencing respiratory distress may appear placid and detached even as they near respiratory arrest. Death in untreated botulism is usually due to airway obstruction from pharyngeal muscle paralysis and inadequate tidal volume resulting from paralysis of diaphragmatic and accessory respiratory muscles. Death can also result from hospital-associated infections and other sequelae of long-term paralysis, hospitalization, and mechanical ventilation.

A history of preparing home-canned foods may assist with the diagnosis. Patients with wound botulism may or may not have a discernible wound or abscess. A history of injection drug use or the presence of track marks may raise suspicion of the diagnosis. Clinical improvement follows sprouting of new nerve terminals and may take weeks to months. Patients often require outpatient rehabilitation therapy and may experience residual deficits.

DIAGNOSIS

Botulism is diagnosed primarily on clinical grounds, with laboratory confirmation by specific tests that identify botulinum toxin in clinical and food samples. In the setting of an outbreak with multiple patients presenting to the same treatment facility, the diagnosis is apparent as long as physicians recognize that cases within a cluster may have varied signs and symptoms. The temporal occurrence of two or more cases with symptoms compatible with botulism is essentially pathognomonic because other illnesses that resemble botulism do not usually occur in clusters. In lone (sporadic) cases, the diagnosis is often missed. The rarity of this disease prevents many physicians from gaining experience with its clinical diagnosis, and some patients present with signs and symptoms that do not fit the classic pattern. Assessing clinical characteristics of other paralytic illnesses in single cases is sometimes critical to rule in or rule out the diagnosis of botulism.

In adults, a food history and the names of contacts who may have shared foods should be obtained before illness progresses to respiratory failure; specific questions should include information about the consumption of home-preserved and/or exotic foods and of products requiring refrigeration that have been left at room temperature in sealed plastic containers or bags. A history of recent consumption of home-canned food substantially enhances the probability of food-borne botulism.

Ascertainment of the patient’s behavioral history related to injection drug use is critical to the diagnosis of wound botulism unless an accidental wound is evident. Caretakers’ observations up to and including the onset of symptoms are vital to the diagnosis of infant

botulism. A history of recent abdominal surgery or antibiotic use may be important in the diagnosis of adult intestinal colonization botulism.

Differential Diagnosis The illnesses most commonly considered in the differential diagnosis of adult botulism cases include Guillain-Barré syndrome (GBS), myasthenia gravis, stroke syndromes, Eaton-Lambert syndrome, and tick paralysis. Less likely possibilities are poisoning by tetrodotoxin, shellfish, or a host of rarer agents and antimicrobial drug-associated paralysis. A thorough history and a meticulous physical examination can effectively eliminate most alternative diagnoses, but a workup for other diagnoses should not delay treatment with botulinum antitoxin.

GBS, a rare autoimmune demyelinating polyneuropathy that often follows an acute infection, presents most often as an ascending paralysis. Case clusters are rare. Occasional GBS cases present as the Miller Fisher variant, whose characteristic triad of ophthalmoplegia, ataxia, and areflexia is easily mistaken for the early descending paralysis of botulism. Protein levels in cerebrospinal fluid (CSF) are elevated in GBS; because this increase may be delayed until several days after symptom onset, an early lumbar puncture with a negative result may need to be repeated. In contrast, CSF findings are generally normal in botulism, although marginally elevated CSF protein concentrations have been reported. In experienced hands, electromyography may differentiate GBS from botulism.

The edrophonium (Tensilon) test is sometimes of value in distinguishing botulism (usually a negative result) from myasthenia gravis (usually a positive result).

In most cerebrovascular accidents, physical examination reveals asymmetry of paralysis and upper motor neuron signs. Brain imaging can reveal the rare basilar stroke that produces symmetric bulbar palsies. Eaton-Lambert syndrome usually manifests as proximal limb weakness in a patient already debilitated by cancer. Tick paralysis is a rare flaccid condition closely resembling botulism and caused by neurotoxins of certain ticks.

Botulism-Specific Laboratory Tests Botulism is confirmed in the laboratory by demonstration of toxin in clinical specimens (e.g., serum, stool, gastric aspirate, and sterile-water enema samples) or in samples of ingested foods. Isolation of toxigenic clostridia from stool also provides evidence of botulism. Wound cultures yielding the organism are highly suggestive in symptomatic cases. The universally accepted method for confirmation of botulism is the mouse bioassay, which is available only in specialized laboratories. Specific guidance about what specimens to collect should be obtained from the testing laboratory because the requirements vary with the form of botulism suspected. Clinical specimens collected early in the hospital admission process should be submitted for testing; toxin results may be negative if specimens are collected >7 days after symptom onset. Because of the extreme potency of botulinum toxin, a test may yield a negative result even when a patient has botulism; thus, a negative result does not exclude this diagnosis. In suspected wound botulism, material from abscesses should be collected in anaerobic culture tubes. New laboratory tests for botulism are being developed but remain experimental. Nerve conduction studies showing reduced amplitude of motor potentials—with or without potentiation by rapid repetitive stimulation in weak muscles—and needle electromyography showing small-motor-unit action potentials are consistent with botulism; these results and those that make alternative diagnoses more likely may be useful. Standard blood work and radiologic studies are not useful in diagnosing botulism.

TREATMENT BOTULISM

The cornerstones of treatment for botulism are meticulous intensive care and immediate administration of botulinum antitoxin. Because antitoxin is most beneficial early in the course of clinical illness, it should be administered as soon as botulism is suspected and before the time-consuming workup for other illnesses is complete. Persons of all ages (including infants) in whom botulism is suspected should be hospitalized immediately so that respiratory

failure—the usual cause of death—can be detected and managed promptly. Vital capacity should be monitored frequently and mechanical ventilation provided as needed. Botulinum antitoxin can limit the progression of illness because it neutralizes toxin molecules in the circulation that have not yet bound to nerve endings. However, antitoxin does not reverse existing paralysis, which may take weeks to improve. In the United States, there are two licensed antitoxin products: Botulism Antitoxin Heptavalent® (BAT; Emergent Biosolutions, Rockville, MD), an equine-derived heptavalent (A through G) product enzymatically de-specciated for treatment of all forms of adult botulism and infant cases not involving serotypes A and B; and Botulism Immune Globulin Intravenous (BabyBIG®; California Department of Public Health, Sacramento, CA), a human-derived product for treating infant botulism caused by serotype A and/or B only. Antitoxin is also available in some other countries. Aminoglycosides and other medications that block the neuromuscular junction may potentiate botulism and should be avoided.

In wound botulism, suspect wounds and abscesses should be cleaned, debrided, and drained promptly. The role of penicillin and metronidazole in treatment and decolonization is unclear. It has been hypothesized that antimicrobial agents may increase circulating botulinum toxin from lysis of bacterial cells.

Person-to-person transmission of botulism does not occur. Universal precautions are the only infection-control measures required during inpatient care.

NOTIFICATION, EXPERT CONSULTATION, AND ANTITOXIN PROVISION

Every botulism case is a public health emergency. Antitoxin is not universally available. Some countries maintain stockpiles of antitoxin for immediate response, whereas others must access supplies from other nations when an outbreak occurs.

In the United States, clinicians must report every suspected case, regardless of form, on an emergency basis to their state health department. The state health department may put the physician in contact with the 24/7 botulism consultation service at the Centers for Disease Control and Prevention (CDC) through the CDC Emergency Operations Center (770-488-7100) or a locally available service. The botulism consultant will review the case and determine whether botulism is likely. If indicated, the consultant will coordinate laboratory confirmation at appropriate testing facilities and facilitate emergency shipment of antitoxin for all adult cases and for infant cases not involving serotypes A and B. In this country, botulinum antitoxin for noninfant cases is available exclusively from the CDC. Physicians who see suspected infant botulism cases should contact the California Department of Public Health Infant Botulism Treatment and Prevention Program (510-231-7600), which provides 24-h consultation and distributes antitoxin (BabyBIG) for the treatment of infant botulism nationwide. Except in cases involving infants who reside in California, laboratory testing requests must still be authorized by the state health department where the infant is located or by the CDC.

PREVENTION

No prophylaxis or licensed vaccine is available against botulism. Home-canning instructions and equipment have changed over the years. Up-to-date canning instructions from reliable sources (e.g., the U.S. Department of Agriculture or the U.S. Food and Drug Administration) should be followed to ensure food safety. Processed food should be stored properly and heated thoroughly prior to consumption. Because of the possible presence of spores, honey should not be given to infants (≤12 months of age). Injection of illicit drugs should be avoided. All wounds should be meticulously cleaned to eliminate possible contamination with bacterial spores. Clinicians should educate individuals or family members of at-risk individuals, including infants, illegal drug users, and preparers of home-preserved foods.

ACKNOWLEDGMENT

The authors thank Dr. Jeremy Sobel for his valued contributions to the previous version of this chapter.

179 Gas Gangrene and Other Clostridial Infections

Amy E. Bryant, Dennis L. Stevens

The genus *Clostridium* encompasses more than 60 species that may be commensals of the gut microflora or may cause a variety of infections in humans and animals through the production of a plethora of proteinaceous exotoxins. *C. tetani* and *C. botulinum*, for example, cause specific clinical disease by elaborating single but highly potent toxins. In contrast, *C. perfringens* and *C. septicum* cause aggressive necrotizing infections that are attributable to multiple toxins, including bacterial proteases, phospholipases, and cytotoxins.

ETIOLOGIC AGENT

Vegetative cells of *Clostridium* species are pleomorphic, rod-shaped, and arranged singly or in short chains (Fig. 179-1); the cells have rounded or sometimes pointed ends. Although clostridia stain gram-positive in the early stages of growth, they may appear to be gram-negative or gram-variable later in the growth cycle or in infected tissue specimens. Most strains are motile by means of peritrichous flagella; *C. septicum* swarms on solid media. Nonmotile species include *C. perfringens*, *C. ramosum*, and *C. innocuum*. Most species are obligately anaerobic, although clostridial tolerance to oxygen varies widely; some species (e.g., *C. septicum*, *C. tertium*) will grow but will not sporulate in air.

Clostridia produce more protein toxins than any other bacterial genus, and more than 25 clostridial toxins lethal to mice have been identified. These proteins include neurotoxins, enterotoxins, cytotoxins, collagenases, permeases, necrotizing toxins, lipases, lecithinases, hemolysins, proteinases, hyaluronidases, DNases, ADP-ribosyltransferases, and neuraminidases. Botulinum and tetanus neurotoxins are the most potent toxins known, with lethal doses of 0.2–10 ng/kg for humans. Epsilon toxin, a 33-kDa protein produced by *C. perfringens* types B and D, causes edema and hemorrhage in the brain, heart, spinal cord, and kidneys of animals. It is among the most lethal of the clostridial toxins and is considered a potential agent of bioterrorism (Chap. 261e). The genomic sequences of some pathogenic clostridia are now available and are likely to facilitate a comprehensive approach to understanding the virulence factors involved in clostridial pathogenesis.

EPIDEMIOLOGY AND TRANSMISSION

Clostridium species are widespread in nature, forming endospores that are commonly found in soil, feces, sewage, and marine sediments. The ecology of *C. perfringens* in soil is greatly influenced by the degree and

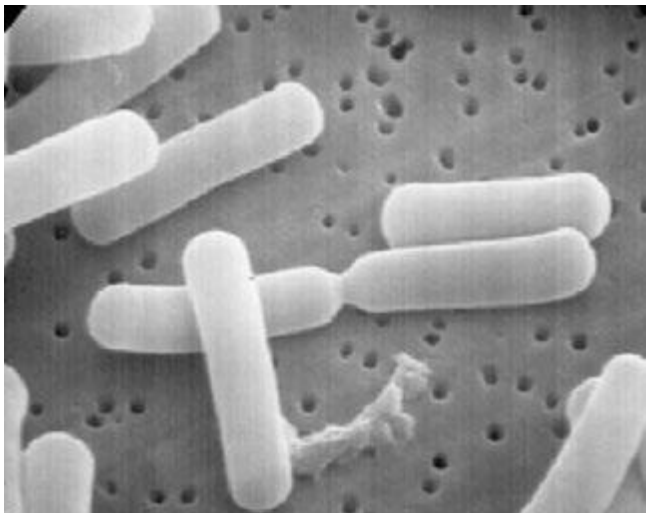


FIGURE 179-1 Scanning electron micrograph of *C. perfringens*.

duration of animal husbandry in a given location and is relevant to the incidence of gas gangrene caused by contamination of war wounds with soil. For example, the incidence of clostridial gas gangrene is higher in agricultural regions of Europe than in the Sahara Desert of Africa. Similarly, the incidences of tetanus and food-borne botulism are clearly related to the presence of clostridial spores in soil, water, and many foods. Clostridia are present in large numbers in the indigenous microbiota of the intestinal tract of humans and animals, in the female genital tract, and on the oral mucosa. It should be noted that not all commensal clostridia are toxigenic.



Clostridial infections remain a serious public health concern worldwide. In developing nations, food poisoning, necrotizing enterocolitis, and gas gangrene are common because large portions of the population are poor and have little or no immediate access to health care. These infections remain prevalent in developed countries as well. Gas gangrene commonly follows knife or gunshot wounds or vehicular accidents or develops as a complication of surgery or gastrointestinal carcinoma. Severe clostridial infections have emerged as a health threat to injection drug users and to women undergoing childbirth or abortion. Historically, clostridial gas gangrene has been the scourge of the battlefield. The global political situation portends another possible scenario involving mass casualties of war or terrorism, with extensive injuries conducive to gas gangrene. Thus there is an ongoing need to develop novel strategies to prevent or attenuate the course of clostridial infections in both civilians and military personnel. Vaccination against exotoxins important in pathogenesis would be of great benefit in developing nations and could also be used safely in at-risk populations such as the elderly, patients with diabetes who may require lower-limb surgery due to trauma or poor circulation, and those undergoing intestinal surgery. Moreover, a hyperimmune globulin would be a valuable tool for prophylaxis in victims of acute traumatic injury or for attenuation of the spread of infection in patients with established gas gangrene.

CLINICAL SYNDROMES

Life-threatening clostridial infections range from intoxications (e.g., food poisoning, tetanus) to necrotizing enteritis/colitis, bacteremia, myonecrosis, and toxic shock syndrome (TSS). **Tetanus and botulism are discussed in Chaps. 177 and 178, respectively. Colitis due to *C. difficile* is discussed in Chap. 161.**

CLOSTRIDIAL WOUND CONTAMINATION

Of open traumatic wounds, 30–80% reportedly are contaminated with clostridial species. In the absence of devitalized tissue, the presence of clostridia does not necessarily lead to infection. In traumatic injuries, clostridia are isolated with equal frequency from both suppurative and well-healing wounds. Thus, diagnosis and treatment of clostridial infection should be based on clinical signs and symptoms and not solely on bacteriologic findings.

POLYMICROBIAL INFECTIONS INVOLVING CLOSTRIDIA

Clostridial species may be found in polymicrobial infections also involving microbial components of the indigenous flora. In these infections, clostridia often appear in association with non-spore-forming anaerobes and facultative or aerobic organisms. Head and neck infections, conjunctivitis, brain abscess, sinusitis, otitis, aspiration pneumonia, lung abscess, pleural empyema, cholecystitis, septic arthritis, and bone infections all may involve clostridia. These conditions are often associated with severe local inflammation but may lack the characteristic systemic signs of toxicity and rapid progression seen in other clostridial infections. In addition, clostridia are isolated from ~66% of intraabdominal infections in which the mucosal integrity of the bowel or respiratory system has been compromised. In this setting, *C. ramosum*, *C. perfringens*, and *C. bifermentans* are the most commonly isolated species. Their presence does not invariably lead to a poor outcome. Clostridia have been isolated from suppurative infections of the female genital tract (e.g., ovarian or pelvic abscess) and from diseased gallbladders. Although the most frequently isolated species is *C. perfringens*, gangrene is not typically observed; however, gas formation in

the biliary system can lead to emphysematous cholecystitis, especially in diabetic patients. *C. perfringens* in association with mixed aerobic and anaerobic microbes can cause aggressive life-threatening type I necrotizing fasciitis or Fournier's gangrene.

The treatment of mixed aerobic/anaerobic infection of the abdomen, perineum, or gynecologic organs should be based on Gram's staining, culture, and antibiotic sensitivity information. Reasonable empirical treatment consists of ampicillin or ampicillin/sulbactam combined with either clindamycin or metronidazole (Table 179-1). Broader gram-negative coverage may be necessary if the patient has recently been hospitalized or treated with antibiotics. Such coverage can be obtained by substituting ticarcillin/clavulanic acid, piperacillin/sulbactam, or a penem antibiotic for ampicillin or by adding a fluoroquinolone or an aminoglycoside to the regimen. Empirical treatment should be given for 10–14 days or until the patient's clinical condition improves.

ENTERIC CLOSTRIDIAL INFECTIONS

C. perfringens type A is one of the most common bacterial causes of food-borne illness in the United States and Canada. The foods typically implicated include improperly cooked meat and meat products (e.g., gravy) in which residual spores germinate and proliferate during slow cooling or insufficient reheating. Illness results from the ingestion of food containing at least $\sim 10^8$ viable vegetative cells, which sporulate in the alkaline environment of the small intestine, producing *C. perfringens* enterotoxin in the process. The diarrhea that develops within 7–30 h of ingestion of contaminated food is generally mild and self-limiting; however, in the very young, the elderly, and the immunocompromised, symptoms are more severe and occasionally fatal. Enterotoxin-producing *C. perfringens* has been implicated as an etiologic agent of persistent diarrhea in elderly patients in nursing homes and tertiary-care institutions and has been considered to play a role in antibiotic-associated diarrhea without pseudomembranous colitis.



C. perfringens strains associated with food poisoning possess the gene (*cpe*) coding for enterotoxin, which acts by forming pores in host cell membranes. *C. perfringens* strains isolated from non-food-borne diseases, such as antibiotic-associated and sporadic diarrhea, carry *cpe* on a plasmid that may be transmitted to other strains. Several methods have been described for the detection of *C. perfringens* enterotoxin in feces, including cell culture assay (Vero cells), enzyme-linked immunosorbent assay, reversed-phase latex agglutination, and polymerase chain reaction (PCR) amplification of *cpe*. Each method has its advantages and limitations.



Enteritis necroticans (gas gangrene of the bowel) is a fulminating clinical illness characterized by extensive necrosis of the intestinal mucosa and wall. Cases can occur sporadically in

adults or as epidemics in people of all ages. Enteritis necroticans is caused by α toxin- and β toxin-producing strains of *C. perfringens* type C; β toxin is located on a plasmid and is mainly responsible for pathogenesis. This life-threatening infection causes ischemic necrosis of the jejunum. In Papua New Guinea during the 1960s, enteritis necroticans (known in that locale as *pigbel*) was found to be the most common cause of death in childhood; it was associated with pig feasts and occurred both sporadically and in outbreaks. Intramuscular immunization against the β toxin resulted in a decreased incidence of the disease in Papua New Guinea, although the condition remains common. Enteritis necroticans has also been recognized in the United States, the United Kingdom, Germany (where it is known as *darmbrand*), and other developed nations; especially affected are adults who are malnourished or who have diabetes, alcoholic liver disease, or neutropenia.

Necrotizing enterocolitis, a disease resembling enteritis necroticans but associated with *C. perfringens* type A, has been found in North America in previously healthy adults. It is also a serious gastrointestinal disease of low-birth-weight (premature) infants hospitalized in neonatal intensive care units. The etiology and pathogenesis of this disease have remained enigmatic for more than four decades. Pathologic similarities between necrotizing enterocolitis and enteritis necroticans include the pattern of small-bowel necrosis involving the submucosa, mucosa, and muscularis; the presence of gas dissecting the tissue planes; and the degree of inflammation. In contrast to enteritis necroticans, which most commonly involves the jejunum, necrotizing enterocolitis affects the ileum and frequently the ileocecal valve. Both diseases may manifest as intestinal gas cysts, although this feature is more common in necrotizing enterocolitis. The sources of the gas, which contains hydrogen, methane, and carbon dioxide, are probably the fermentative activities of intestinal bacteria, including clostridia. Epidemiologic data support an important role for *C. perfringens* or other gas-producing microorganisms (e.g., *C. neonatale*, certain other clostridia, or *Klebsiella* species) in the pathogenesis of necrotizing enterocolitis.

Patients with suspected clostridial enteric infection should undergo nasogastric suction and receive IV fluids. Pyrantel is given by mouth, and the bowel is rested by fasting. Benzylpenicillin (1 mU) is given IV every 4 h, and the patient is observed for complications requiring surgery. Patients with mild cases recover without surgical intervention. If surgical indications are present (gas in the peritoneal cavity, absent bowel sounds, rebound tenderness, abdominal rigidity), however, the mortality rate ranges from 35% to 100%; a fatal outcome is due in part to perforation of the intestine.

As *pigbel* continues to be a common disease in Papua New Guinea, consideration should be given to the use of a *C. perfringens*

TABLE 179-1 TREATMENT OF CLOSTRIDIAL INFECTIONS

Condition	Antibiotic Treatment	Penicillin Allergy	Adjunctive Treatment/Note
Wound contamination	None	—	Treatment should be based on clinical signs and symptoms as listed below and not solely on bacteriologic findings.
Polymicrobial anaerobic infections involving clostridia (e.g., abdominal wall, gynecologic)	Ampicillin (2 g IV q4h) plus Clindamycin (600–900 mg IV q6–8h) plus Ciprofloxacin (400 mg IV q6–8 h)	Vancomycin (1 g IV q12h) plus Metronidazole (500 mg IV q6h) plus Ciprofloxacin (400 mg IV q6–8h)	Empirical therapy should be initiated. Therapy should be based on Gram's stain and culture results and on sensitivity data when available. Add gram-negative coverage if indicated (see text).
Clostridial sepsis	Penicillin, 3–4 mU IV q4–6h plus Clindamycin (600–900 mg IV q6–8h)	Clindamycin alone or Metronidazole (as above) or Vancomycin (as above)	Transient bacteremia without signs of systemic toxicity may be clinically insignificant.
Gas gangrene	Penicillin G (4 mU IV q4–6 h) plus Clindamycin (600–900 mg IV q6–8h)	Cefoxitin (2 g IV q6h) plus Clindamycin (600–900 mg IV q6–8h)	Emergent surgical exploration and thorough debridement are extremely important. Hyperbaric oxygen therapy may be considered after surgery and antibiotic initiation.

992 type C β toxoid vaccine in local areas. Two doses given 3–4 months apart are preventive.

CLOSTRIDIAL BACTEREMIA

Clostridium species are important causes of bloodstream infections. Molecular epidemiologic studies of anaerobic bacteremia have identified *C. perfringens* and *C. tertium* as the two most frequently isolated species; these organisms cause up to 79% and 5%, respectively, of clostridial bacteremias. Occasionally, *C. perfringens* bacteremia occurs in the absence of an identifiable infection at another site. When associated with myonecrosis, bacteremia has a grave prognosis.

C. septicum is also commonly associated with bacteremia. This species is isolated only rarely from the feces of healthy individuals but may be found in the normal appendix. More than 50% of patients whose blood cultures are positive for this organism have some gastrointestinal anomaly (e.g., diverticular disease) or underlying malignancy (e.g., carcinoma of the colon). In addition, a clinically important association of *C. septicum* bacteremia with neutropenia of any origin—and, more specifically, with neutropenic enterocolitis involving the terminal ileum or cecum—has been observed. Patients with diabetes mellitus, severe atherosclerotic cardiovascular disease, or anaerobic myonecrosis (gas gangrene) also may develop *C. septicum* bacteremia. *C. septicum* has been recovered from the bloodstream of cirrhotic patients, as have *C. perfringens*, *C. bifermentans*, and other clostridia. Infections of the bloodstream by *C. sordellii* and *C. perfringens* have been associated with TSS.

Bloodstream infection by *C. tertium*, either alone or in combination with *C. septicum* or *C. perfringens*, can be found in patients with serious underlying disease such as malignancy or acute pancreatitis, with or without neutropenic enterocolitis; the frequency has not been systematically studied. *C. tertium* may present special problems in terms of both identification and treatment. This organism may stain gram-negative; is aerotolerant; and is resistant to metronidazole, clindamycin, and cephalosporins.

Other clostridia from the *C. clostridioforme* group (including *C. clostridioforme*, *C. hathewayi*, and *C. boltea*) can cause bacteremia.

The clinical importance of recognizing clostridial bacteremia—especially that due to *C. septicum*—and starting appropriate treatment immediately (Table 179-1) cannot be overemphasized. Patients with this condition usually are gravely ill, and infection may metastasize to distant anatomic sites, resulting in spontaneous myonecrosis (see next section). Alternative methods to identify bacteremia-causing clostridial species, such as PCR or other rapid diagnostic tests, are not currently available. Anaerobic blood cultures and Gram's stain interpretation remain the best diagnostic tests at this point.

CLOSTRIDIAL SKIN AND SOFT-TISSUE INFECTIONS

Histotoxic clostridial species such as *C. perfringens*, *C. histolyticum*, *C. septicum*, *C. novyi*, and *C. sordellii* cause aggressive necrotizing infections of the skin and soft tissues. These infections are attributable in part to the elaboration of bacterial proteases, phospholipases, and cytotoxins. Necrotizing clostridial soft-tissue infections are rapidly progressive and are characterized by marked tissue destruction, gas in the tissues, and shock; they frequently end in death. Severe pain, crepitus, brawny induration with rapid progression to skin sloughing, violaceous bullae, and marked tachycardia are characteristics found in the majority of patients.

Clostridial Myonecrosis (Gas Gangrene) • TRAUMATIC GAS GANGRENE *C. perfringens* myonecrosis (gas gangrene) is one of the most fulminant gram-positive bacterial infections of humans. Even with appropriate antibiotic therapy and management in an intensive care unit, tissue destruction can progress rapidly. Gas gangrene is accompanied by bacteremia, hypotension, and multiorgan failure and is invariably fatal if untreated. Gas gangrene is a true emergency and requires immediate surgical debridement.

The development of gas gangrene requires an anaerobic environment and contamination of a wound with spores or vegetative organisms. Devitalized tissue, foreign bodies, and ischemia reduce

locally available oxygen levels and favor outgrowth of vegetative cells and spores. Thus conditions predisposing to traumatic gas gangrene include crush-type injury, laceration of large or medium-sized arteries, and open fractures of long bones that are contaminated with soil or bits of clothing containing the bacterial spores. Gas gangrene of the abdominal wall and flanks follows penetrating injuries such as knife or gunshot wounds that are sufficient to compromise intestinal integrity, with resultant leakage of the bowel contents into the soft tissues. Proximity to fecal sources of bacteria is a risk factor for cases following hip surgery, adrenaline injections into the buttocks, or amputation of the leg for ischemic vascular disease. In the last decade, cutaneous gas gangrene caused by *C. perfringens*, *C. novyi*, and *C. sordellii* has been described in the United States and northern Europe among persons injecting black-tar heroin subcutaneously.

The incubation period for traumatic gas gangrene can be as short as 6 h and is usually <4 days. The infection is characterized by the sudden onset of excruciating pain at the affected site and the rapid development of a foul-smelling wound containing a thin serosanguineous discharge and gas bubbles. Brawny edema and induration develop and give way to cutaneous blisters containing bluish to maroon-colored fluid. Such tissue later may become liquefied and slough. The margin between healthy and necrotic tissue often advances several inches per hour despite appropriate antibiotic therapy, and radical amputation remains the single best life-saving intervention. Shock and organ failure frequently accompany gas gangrene; when patients become bacteremic, the mortality rate exceeds 50%.

Diagnosis of traumatic gas gangrene is not difficult because the infection always begins at the site of significant trauma, is associated with gas in the tissue, and is rapidly progressive. Gram's staining of drainage or tissue biopsy is usually definitive, demonstrating large gram-positive (or gram-variable) rods, an absence of inflammatory cells, and widespread soft-tissue necrosis.

SPONTANEOUS (NONTRAUMATIC) GAS GANGRENE Spontaneous gas gangrene generally occurs via hematogenous seeding of normal muscle with histotoxic clostridia—principally *C. perfringens*, *C. septicum*, and *C. novyi* and occasionally *C. tertium*—from a gastrointestinal tract portal of entry (as in colonic malignancy, inflammatory bowel disease, diverticulitis, necrotizing enterocolitis, cecitis, or distal ileitis or after gastrointestinal surgery). These gastrointestinal pathologies permit bacterial access to the bloodstream; consequently, aerotolerant *C. septicum* can proliferate in normal tissues. Patients surviving bacteremia or spontaneous gangrene due to *C. septicum* should undergo aggressive diagnostic studies to rule out gastrointestinal pathology.

Additional predisposing host factors include leukemia, lymphoproliferative disorders, cancer chemotherapy, radiation therapy, and AIDS. Cyclic, congenital, or acquired neutropenia also is strongly associated with an increased incidence of spontaneous gas gangrene due to *C. septicum*; in such cases, necrotizing enterocolitis, cecitis, or distal ileitis is common, particularly among children.

The first symptom of spontaneous gas gangrene may be confusion followed by the abrupt onset of excruciating pain in the absence of trauma. These findings, along with fever, should heighten suspicion of spontaneous gas gangrene. However, because of the lack of an obvious portal of entry, the correct diagnosis is frequently delayed or missed. The infection is characterized by rapid progression of tissue destruction with demonstrable gas in the tissue (Fig. 179-2). Swelling increases, and bullae filled with clear, cloudy, hemorrhagic, or purplish fluid appear. The surrounding skin has a purple hue, which may reflect vascular compromise resulting from the diffusion of bacterial toxins into surrounding tissues. Invasion of healthy tissue rapidly ensues, with quick progression to shock and multiple-organ failure. Mortality rates in this setting range from 67 to 100% among adults; among children, the mortality rate is 59%, with the majority of deaths occurring within 24 h of onset.

PATHOGENESIS OF GAS GANGRENE In traumatic gas gangrene, organisms are introduced into devitalized tissue. It is important to recognize that for *C. perfringens* and *C. novyi*, trauma must be sufficient to interrupt the blood supply and thereby to establish an optimal anaerobic



FIGURE 179-2 Radiograph of patient with spontaneous gas gangrene due to *C. septicum*, demonstrating gas in the affected arm and shoulder.

environment for growth of these species. These conditions are not strictly required for the more aerotolerant species such as *C. septicum* and *C. tertium*, which can seed normal tissues from gastrointestinal lesions. Once introduced into an appropriate niche, the organisms proliferate locally and elaborate exotoxins.

The major *C. perfringens* extracellular toxins implicated in gas gangrene are α toxin and θ toxin. A lethal hemolysin that has both phospholipase C and sphingomyelinase activities, α toxin has been implicated as the major virulence factor of *C. perfringens*: immunization of mice with the C-terminal domain of α toxin provides protection against lethal challenge with *C. perfringens*, and isogenic α toxin-deficient mutant strains of *C. perfringens* are not lethal in a murine model of gas gangrene. It has been shown in experimental models that the severe pain, rapid progression, marked tissue destruction, and absence of neutrophils in *C. perfringens* gas gangrene are attributable in large part to a toxin-induced occlusion of blood vessels by heterotypic aggregates of platelets and neutrophils. The formation of these aggregates, which occurs within minutes, is largely mediated by α toxin's ability to activate the platelet adhesion molecule gpIIb/IIIa (Fig. 179-3); the implication is that platelet glycoprotein inhibitors (e.g., eptifibatide, abciximab) may be therapeutic for maintaining tissue blood flow.

C. perfringens θ toxin (*perfringolysin*) is a member of the thiol-activated cytolysin family known as cholesterol-dependent cytolysins, which includes streptolysin O from group A *Streptococcus*, pneumolysin from *Streptococcus pneumoniae*, and several other toxins. Cholesterol-dependent cytolysins bind as oligomers to cholesterol in host cell membranes. At high concentrations, these toxins form ring-like pores resulting in cell lysis. At sublytic concentrations, θ toxin hyperactivates phagocytes and vascular endothelial cells.

Cardiovascular collapse and end-organ failure occur late in the course of *C. perfringens* gas gangrene and are largely attributable to both direct and indirect effects of α and θ toxins. In experimental models, θ toxin causes markedly reduced systemic vascular resistance but increased cardiac output (i.e., “warm shock”), probably via induction of endogenous mediators (e.g., prostacyclin, platelet-activating factor)

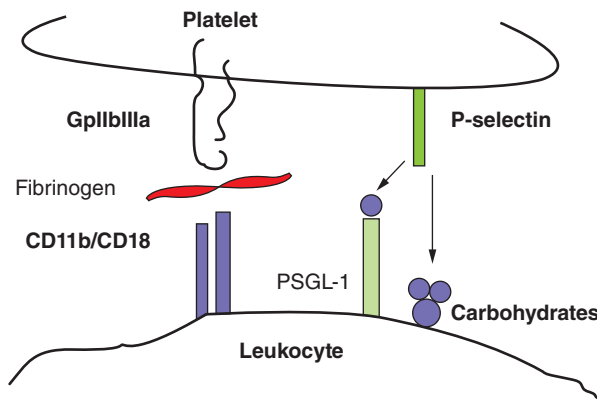


FIGURE 179-3 Schematic illustration of the molecular mechanisms of *C. perfringens* α toxin-induced platelet/neutrophil aggregates. Homotypic aggregates of platelets (not shown) and heterotypic aggregates of platelets and leukocytes are due to a toxin-induced activation of the platelet fibrinogen receptor gpIIb/IIIa and upregulation of leukocyte CD11b/CD18. Binding of fibrinogen (red) bridges the connection between these adhesion molecules on adjacent cells. An auxiliary role for α toxin-induced upregulation of platelet P-selectin and its binding to leukocyte P-selectin glycoprotein ligand 1 (PSGL-1) or other leukocyte surface carbohydrates also has been demonstrated.

that cause vasodilation. This effect is similar to that observed in gram-negative sepsis. In sharp contrast, α toxin directly suppresses myocardial contractility; the consequence is profound hypotension due to a sudden reduction in cardiac output. The roles of other endogenous mediators, such as cytokines (e.g., tumor necrosis factor, interleukin 1, interleukin 6) and vasodilators (e.g., bradykinin) have not been fully elucidated.

C. septicum produces four main toxins— α toxin (lethal, hemolytic, necrotizing activity), β toxin (DNase), γ toxin (hyaluronidase), and Δ toxin (septicolysin, an oxygen-labile hemolysin)—as well as a protease and a neuraminidase. Unlike the α toxin of *C. perfringens*, that of *C. septicum* does not possess phospholipase activity. The mechanisms remain to be fully elucidated, but it is likely that each of these toxins contributes uniquely to *C. septicum* gas gangrene.

TREATMENT GAS GANGRENE

Patients with suspected gas gangrene (either traumatic or spontaneous) should undergo prompt surgical inspection of the infected site. Direct examination of a Gram-stained smear of the involved tissues is of major importance. Characteristic histologic findings in clostridial gas gangrene include widespread tissue destruction, a paucity of leukocytes in infected tissues in conjunction with an accumulation of leukocytes in adjacent vessels (Fig. 179-4), and the presence of gram-positive rods (with or without spores). CT and MRI are invaluable for determining whether the infection is localized or is spreading along fascial planes, and needle aspiration or punch biopsy may provide an etiologic diagnosis in at least 20% of cases. However, these techniques should not replace surgical exploration, Gram's staining, and histopathologic examination. When spontaneous gas gangrene is suspected, blood should be cultured since bacteremia usually precedes cutaneous manifestations by several hours.

For patients with evidence of clostridial gas gangrene, thorough emergent surgical debridement is of extreme importance. All devitalized tissue should be widely resected back to healthy viable muscle and skin so as to remove conditions that allow anaerobic organisms to continue proliferating. Closure of traumatic wounds or compound fractures should be delayed for 5–6 days until it is certain that these sites are free of infection.

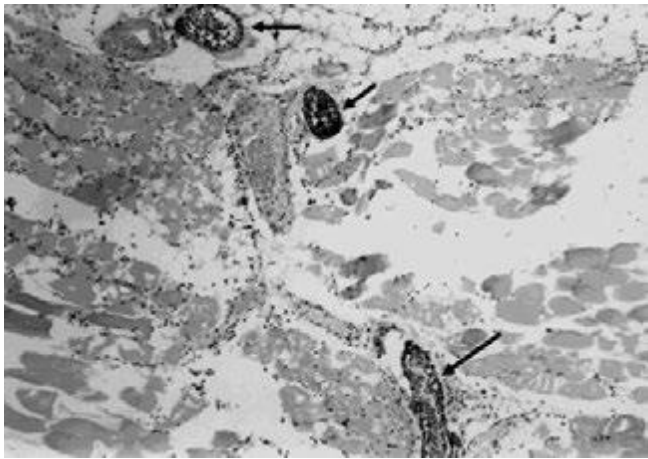


FIGURE 179-4 Histopathology of experimental gas gangrene due to *C. perfringens*, demonstrating widespread muscle necrosis, a paucity of leukocytes in infected tissues, and accumulation of leukocytes in adjacent vessels (arrows). These features are due to the effects of α and θ toxins on muscle cells, platelets, leukocytes, and endothelial cells.

Antibiotic treatment of traumatic or spontaneous gas gangrene (Table 179-1) consists of the administration of penicillin and clindamycin for 10–14 days. Penicillin is recommended on the basis of in vitro sensitivity data; clindamycin is recommended because of its superior efficacy over penicillin in animal models of *C. perfringens* gas gangrene and in some clinical reports. Controlled clinical trials comparing the efficacy of these agents in humans have not been performed. In the penicillin-allergic patient, clindamycin may be used alone. The superior efficacy of clindamycin is probably due to its ability to inhibit bacterial protein toxin production, its insensitivity to the size of the bacterial load or the stage of bacterial growth, and its ability to modulate the host's immune response.

C. tertium is resistant to penicillin, cephalosporins, and clindamycin. Appropriate antibiotic therapy for *C. tertium* infection is vancomycin (1 g every 12 h IV) or metronidazole (500 mg every 8 h IV).

The value of adjunctive treatment with hyperbaric oxygen (HBO) for gas gangrene remains controversial. Basic science studies suggest that HBO can inhibit the growth of *C. perfringens* but not that of the more aerotolerant *C. septicum*. In vitro, blood and macerated muscle inhibit the bactericidal potential of HBO. Numerous studies in animals demonstrate little efficacy of HBO alone, whereas antibiotics alone—especially those that inhibit bacterial protein synthesis—confer marked benefits. Addition of HBO to the therapeutic regimen provides some additional benefit, but only if surgery and antibiotic administration precede HBO treatment.

In conclusion, gas gangrene is a rapidly progressive infection whose outcome depends on prompt recognition, emergent surgery, and timely administration of antibiotics that inhibit toxin production. Gas gangrene associated with bacteremia probably represents a later stage of illness and is associated with the worst outcomes. Emergent surgical debridement is crucial to ensure survival, and ancillary procedures (e.g., CT or MRI) or transport to HBO units should not delay this intervention. Some trauma centers associated with HBO units may have special expertise in managing these aggressive infections, but proximity and speed of transfer must be carefully weighed against the need for haste.

PROGNOSIS OF GAS GANGRENE The prognosis for patients with gas gangrene is more favorable when the infection involves an extremity rather than the trunk or visceral organs, since debridement of the latter sites is more difficult. Gas gangrene is most likely to progress to shock and death in patients with associated bacteremia and intravascular hemolysis. Mortality rates are highest for patients in shock at the time

of diagnosis. Mortality rates are relatively high among patients with spontaneous gas gangrene, especially that due to *C. septicum*. Survivors of gas gangrene may undergo multiple debridements and face long periods of hospitalization and rehabilitation.

PREVENTION OF GAS GANGRENE Initial aggressive debridement of devitalized tissue can reduce the risk of gas gangrene in contaminated deep wounds. Interventions to be avoided include prolonged application of tourniquets and surgical closure of traumatic wounds; patients with compound fractures are at significant risk for gas gangrene if the wound is closed surgically. Vaccination against a toxin is protective in experimental animal models of *C. perfringens* gas gangrene but has not been investigated in humans. In addition, as mentioned above, a hyperimmune globulin would represent a significant advance for prophylaxis in victims of acute traumatic injury or for attenuation of the spread of infection in patients with established gas gangrene.

Toxic Shock Syndrome Clostridial infection of the endometrium, particularly that due to *C. sordellii*, can develop after gynecologic procedures, childbirth, or abortion (spontaneous or elective, surgical or medical) and, once established, proceeds rapidly to TSS and death. Systemic manifestations, including edema, effusions, profound leukocytosis, and hemoconcentration, are followed by the rapid onset of hypotension and multiple-organ failure. Elevation of the hematocrit to 75–80% and leukocytosis of 50,000–200,000 cells/ μ L, with a left shift, are characteristic of *C. sordellii* infection. Pain may not be a prominent feature, and fever is typically absent. In one series, 18% of 45 cases of *C. sordellii* infection were associated with normal childbirth, 11% with medically induced abortion, and 0.4% with spontaneous abortion; the case-fatality rate was 100% in these groups. Of the infections in this series that were not related to gynecologic procedures or childbirth, 22% occurred in injection drug users, and 50% of these patients died. Other infections followed trauma or surgery (42%), mostly in healthy persons, and 53% of these patients died. Overall, the mortality rate was 69% (31 of 45 cases). Of patients who succumbed, 85% died within 2–6 days after infection onset or following procedures.

Early diagnosis of *C. sordellii* infections often proves difficult for several reasons. First, the prevalence of these infections is low. Second, the initial symptoms are nonspecific and frankly misleading. Early in the course, the illness resembles any number of infectious diseases, including viral syndromes. Given these vague symptoms and an absence of fever, physicians usually do not aggressively pursue additional diagnostic tests. The absence of local evidence of infection and the lack of fever make early diagnosis of *C. sordellii* infection particularly problematic in patients who develop deep-seated infection following childbirth, therapeutic abortion, gastrointestinal surgery, or trauma. Such patients are frequently evaluated for pulmonary embolization, gastrointestinal bleeding, pyelonephritis, or cholecystitis. Unfortunately, such delays in diagnosis increase the risk of death, and, as in most necrotizing soft-tissue infections, patients are hypotensive with evidence of organ dysfunction by the time local signs and symptoms become apparent. In contrast, infection is more readily suspected in injection drug users presenting with local swelling, pain, and redness at injection sites; early recognition probably contributes to the lower mortality rates in this group.

Physicians should suspect *C. sordellii* infection in patients who present within 2–7 days after injury, surgery, drug injection, childbirth, or abortion and who report pain, nausea, vomiting, and diarrhea but are afebrile. There is little information regarding appropriate treatment for *C. sordellii* infections. In fact, the interval between onset of symptoms and death is often so short that there is little time to initiate empirical antimicrobial therapy. Indeed, anaerobic cultures of blood and wound aspirates are time-consuming, and many hospital laboratories do not routinely perform antimicrobial sensitivity testing on anaerobes. Antibiotic susceptibility data from older studies suggest that *C. sordellii*, like most clostridia, is susceptible to β -lactam antibiotics, clindamycin, tetracycline, and chloramphenicol but is resistant to aminoglycosides and sulfonamides. Antibiotics that suppress toxin synthesis (e.g., clindamycin) may possibly prove useful as therapeutic

adjuncts since they are effective in necrotizing infections due to other toxin-producing gram-positive organisms.

Other Clostridial Skin and Soft-Tissue Infections *Crepitant cellulitis* (also called *anaerobic cellulitis*) occurs principally in diabetic patients and characteristically involves subcutaneous tissues or retroperitoneal tissues, whereas the muscle and fascia are not involved. This infection can progress to fulminant systemic disease.

Cases of *C. histolyticum* infection with cellulitis, abscess formation, or endocarditis have also been documented in injection drug users. Endophthalmitis due to *C. sordellii* or *C. perfringens* has been described. *C. ramosum* is also isolated frequently from clinical specimens, including blood and both intraabdominal and soft tissues. This species may be resistant to clindamycin and multiple cephalosporins.

SECTION 6 DISEASES CAUSED BY GRAM-NEGATIVE BACTERIA

180 Meningococcal Infections

Andrew J. Pollard

DEFINITION

Infection with *Neisseria meningitidis* most commonly manifests as asymptomatic colonization in the nasopharynx of healthy adolescents and adults. Invasive disease occurs rarely, usually presenting as either bacterial meningitis or meningococcal septicemia. Patients may also present with occult bacteremia, pneumonia, septic arthritis, conjunctivitis, and chronic meningococcemia.

ETIOLOGY AND MICROBIOLOGY

N. meningitidis is a gram-negative aerobic diplococcus that colonizes humans only and that causes disease after transmission to a susceptible individual. Several related organisms have been recognized, including the pathogen *N. gonorrhoeae* and the commensals *N. lactamica*, *N. flavescens*, *N. mucosa*, *N. sicca*, and *N. subflava*. *N. meningitidis* is a catalase- and oxidase-positive organism that utilizes glucose and maltose to produce acid.

Meningococci associated with invasive disease are usually encapsulated with polysaccharide, and the antigenic nature of the capsule determines an organism's serogroup (Table 180-1). In total, 13 serogroups have been identified (A–D, X–Z, 29E, W, H–J, and L), but just 6 serogroups—A, B, C, X, Y, and W (formerly W135)—account for the majority of cases of invasive disease. Acapsular meningococci are commonly isolated from the nasopharynx in studies of carriage; the lack of capsule often is a result of phase variation of capsule expression, but as many as 16% of isolates lack the genes for capsule synthesis and assembly. These “capsule-null” meningococci and those that express

capsules other than A, B, C, X, Y, and W are only rarely associated with invasive disease and are most commonly identified in the nasopharynx of asymptomatic carriers.

Beneath the capsule, meningococci are surrounded by an outer phospholipid membrane containing lipopolysaccharide (LPS, endotoxin) and multiple outer-membrane proteins (Figs. 180-1 and 180-2). Antigenic variability in porins expressed in the outer membrane defines the serotype (PorB) and serosubtype (PorA) of the organism, and structural differences in LPS determine the immunotype. Serologic methods for typing of meningococci are restricted by the limited availability of serologic reagents that can distinguish among the organisms' highly variable surface proteins. Where available, high-throughput antigen gene sequencing has superseded serology for meningococcal typing. A large database of antigen gene sequences for the outer-membrane proteins PorA, PorB, FetA, Opa, and factor H-binding protein is available online (www.neisseria.org). The number of specialized iron-regulated proteins found in the meningococcal outer membrane (e.g., FetA and transferrin-binding proteins) highlights the organisms' dependence on iron from human sources. A thin peptidoglycan cell wall separates the outer membrane from the cytoplasmic membrane.



The structure of meningococcal populations involved in local and global spread has been studied with multilocus enzyme electrophoresis (MLEE), which characterizes isolates according to differences in the electrophoretic mobility of cytoplasmic enzymes. However, this technique has mostly been replaced by multilocus sequence typing (MLST), in which meningococci are characterized by sequence types assigned on the basis of sequences of internal fragments of seven housekeeping genes. The online MLST database currently includes more than 27,000 meningococcal isolates and 10,500 unique sequence types (pubmlst.org/neisseria/). A limited number of hyperinvasive lineages of *N. meningitidis* have been recognized

TABLE 180-1 STRUCTURE OF THE POLYSACCHARIDE CAPSULE OF COMMON DISEASE-CAUSING MENINGOCOCCI

Meningococcal Serogroup	Chemical Structure of Oligosaccharide	Current Disease Epidemiology
A	2-Acetamido-2-deoxy-D-mannopyranosyl phosphate	Epidemic disease mainly in sub-Saharan Africa; sporadic cases worldwide
B	α -2,8-N-acetylneuraminic acid	Sporadic cases worldwide; propensity to cause hyperendemic disease
C	α -2,9-O-acetylneuraminic acid	Small outbreaks and sporadic disease
Y	4-O- α -D-glucopyranosyl-N-acetylneuraminic acid	Sporadic disease and occasional small institutional outbreaks
W	4-O- α -D-galactopyranosyl-N-acetylneuraminic acid	Sporadic disease; outbreaks of disease associated with mass gatherings; epidemics in sub-Saharan Africa
X	(α 1 \rightarrow 4) N-acetyl-D-glucosamine-1-phosphate	Sporadic disease and large outbreaks in the meningitis belt of Africa

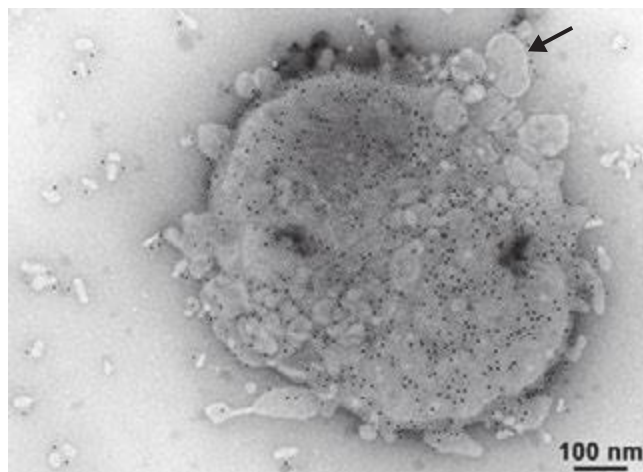


FIGURE 180-1 Electron micrograph of *Neisseria meningitidis*. Black dots are gold-labeled polyclonal antibodies binding surface opacity proteins. Blebs of outer membrane can be seen being released from the bacterial surface (arrow). (Photo courtesy of D. Ferguson, Oxford University.)

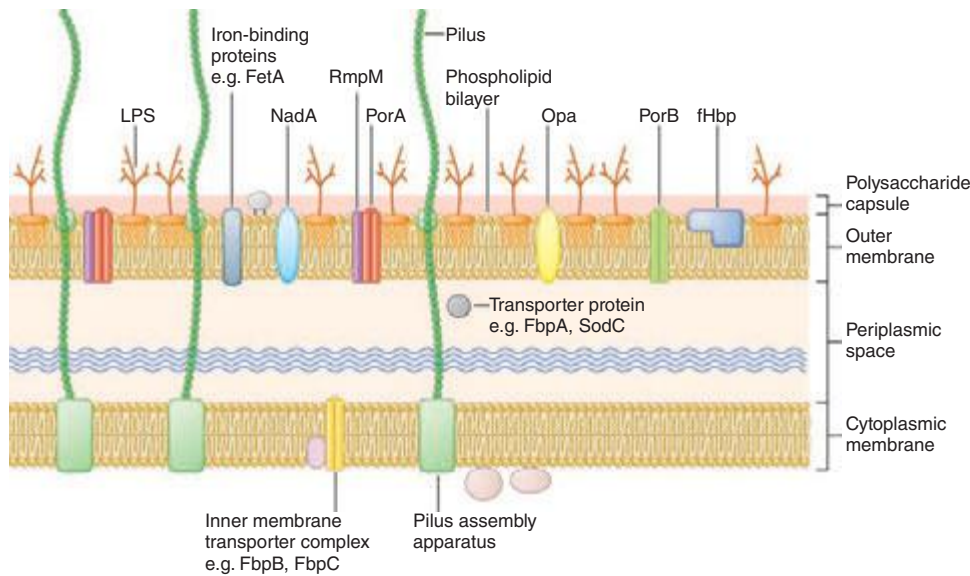


FIGURE 180-2 Cross-section through surface structures of *Neisseria meningitidis*. LPS, lipopolysaccharide. (Reprinted with permission from M Sadarangani, AJ Pollard: *Lancet Infect Dis* 10:112, 2010.)

and are responsible for the majority of cases of invasive meningococcal disease worldwide. The apparent genetic stability of these meningococcal clones over decades and during wide geographic spread indicates that they are well adapted to the nasopharyngeal environment of the host and to efficient transmission. While MLST has become established as the main method of genotyping meningococci in many reference laboratories over the past decade, whole-genome sequencing is set to replace this approach in the decade ahead, with almost 1000 genomes already available in the United Kingdom's national library (www.meningitis.org/genome-library).

The group B meningococcal genome is >2 megabases in length and contains 2158 coding regions. Many genes undergo phase variation that makes it possible to control their expression; this capacity is likely to be important in meningococcal adaptation to the host environment and evasion of the immune response. Meningococci can obtain DNA from their environment and can acquire new genes—including the capsular operon—such that *capsule switching* from one serogroup to another can occur.

EPIDEMIOLOGY

Patterns of Disease Up to 500,000 cases of meningococcal disease are thought to occur worldwide each year, and ~10% of the individuals affected die. There are several patterns of disease: epidemic, outbreak (small clusters of cases), hyperendemic, and sporadic or endemic.

Epidemics have continued since the original descriptions of meningococcal disease, especially affecting the sub-Saharan meningitis belt of Africa, where tens to hundreds of thousands of cases (caused mainly by serogroup A but also by serogroups W and X) may be reported over a season and rates may be as high as 1000 cases per 100,000 population. Serogroup A epidemics took place in Europe and North America after the First and Second World Wars, and serogroup A outbreaks have been documented over the past 30 years in New Zealand, China, Nepal, Mongolia, India, Pakistan, Poland, and Russia.

Clusters of cases occur where there is an opportunity for increased transmission—i.e., in (semi-)closed communities such as schools, colleges, universities, military training centers, and refugee camps. Recently, such clusters have been especially strongly linked with a particular clone (sequence type 11) that is mainly associated with the serogroup C or W capsule but was first described in association with serogroup B. Wider and more prolonged community outbreaks (hyperendemic disease) due to single clones of serogroup B meningococci account for ≥ 10 cases per 100,000. Regions affected in the past decade include the U.S. Pacific Northwest, New Zealand (both

islands), and the province of Normandy in France.

Most countries now experience predominantly sporadic cases (0.3–5 cases per 100,000 population), with many different disease-causing clones involved and usually no clear epidemiologic link between one case and another. The disease rate and the distribution of meningococcal strains vary in different regions of the world and also in any one location over time. For example, in the United States, the rate of meningococcal disease fell from 1.2 cases per 100,000 population in 1997 to <0.15 case per 100,000 in 2012 (Fig. 180-3). Meningococcal disease in the United States was previously dominated by serogroups B and C; however, serogroup Y emerged during the 1990s and became more common than serogroup C in 2007. In contrast, rates of disease in England and Wales rose to >5 cases per 100,000 during the 1990s because of an increase in cases caused by the

ST11 serogroup C clone. As a result of a mass immunization program against serogroup C in 1999, almost all cases in the United Kingdom are now attributed to serogroup B (Fig. 180-4). Over the last decade, most industrialized nations have seen a general decrease in meningococcal disease; this decrease is linked to immunization against serogroup C meningococci in Europe, Canada, and Australia and to adolescent immunization programs for A, C, Y and W in the United States. However, other factors, including changes in population immunity (probably the explanation for the cyclic nature of meningococcal disease rates) as well as a reduction in smoking and passive exposure to tobacco smoke (driven by bans on smoking in buildings and public spaces) across wealthy countries are likely to have contributed to the fall in cases.

Factors Associated with Disease Risk and Susceptibility The principal determinant of disease susceptibility is age, with the peak incidence in the first year of life (Fig. 180-5). The susceptibility of the very young presumably results from an absence of specific adaptive immunity in combination with very close contact with colonized individuals, including parents. Compared with other age groups, infants appear to be particularly susceptible to serogroup B disease: >30% of serogroup B cases in the United States occur during the first year of life. In the early

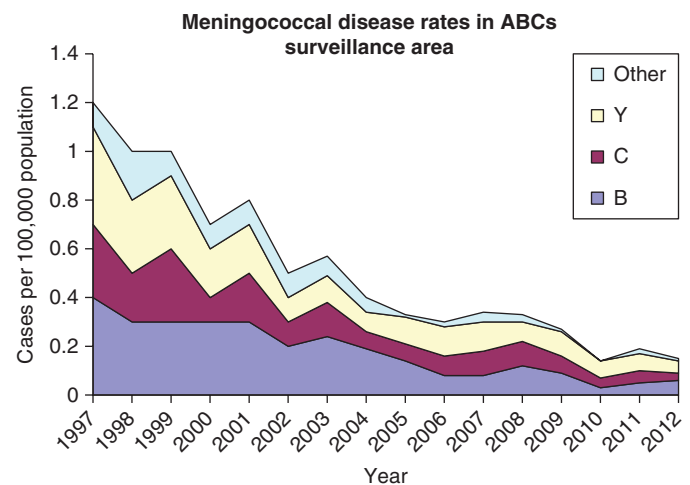


FIGURE 180-3 Meningococcal disease in the United States over time. ABCs, active bacterial cores. (Adapted from ABC Surveillance data, Centers for Disease Control and Prevention; www.cdc.gov.)

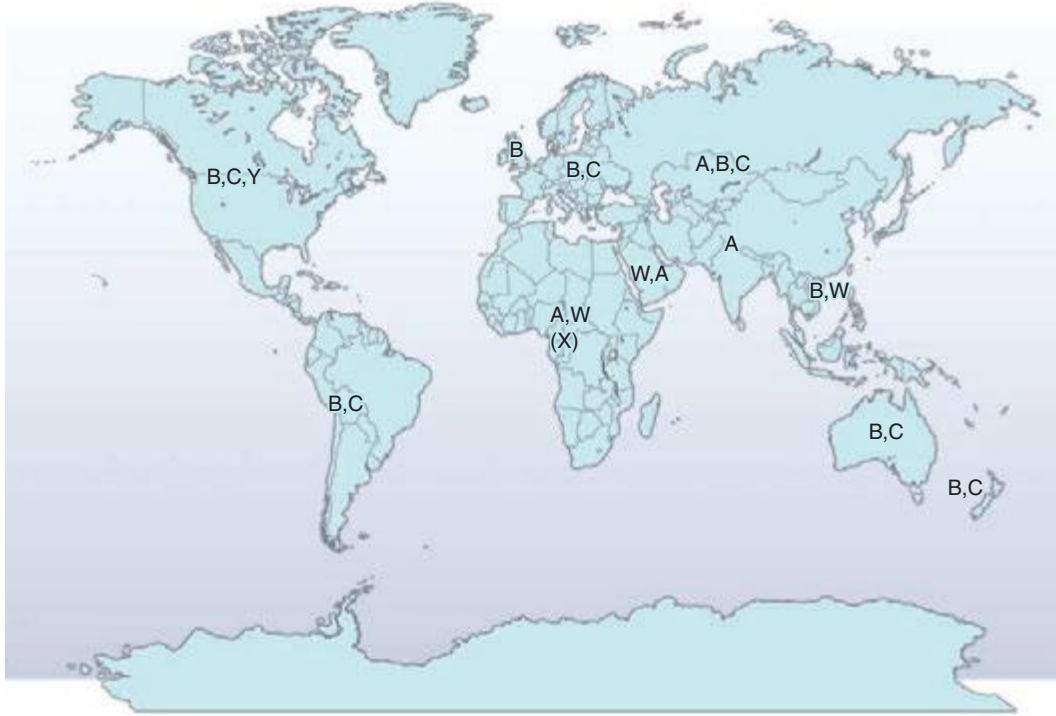


FIGURE 180-4 Global distribution of meningococcal serogroups, 1999–2009.

1990s in North America, the median ages for patients with disease due to serogroups B, C, Y, and W were 6, 17, 24, and 33 years, respectively.



After early childhood, a second peak of disease occurs among adolescents and young adults (15–25 years of age) in Europe and North America. It is thought that this peak relates to social behaviors and environmental exposures in this age group, as discussed below. Most cases of infection with *N. meningitidis* in developed countries today are sporadic, and the rarity of the disease suggests that individual susceptibility may be important. A number of factors probably contribute to individual susceptibility, including the host's genetic constitution, environment, and contact with a carrier or a case.

The best-documented genetic association with meningococcal disease is complement deficiency, chiefly of the terminal complement components (C5–9), properdin, or factor D; such a deficiency increases the risk of disease by up to 600-fold and may result in recurrent attacks. Complement components are believed to be important for the bactericidal activity of serum, which is considered the principal mechanism of immunity against invasive meningococcal disease. However, when investigated, complement deficiency is found in only a very small proportion of individuals with meningococcal

disease (0.3%). Conversely, 7–20% of persons whose disease is caused by the less common serogroups (W, X, Y, Z, 29E) have a complement deficiency. Complement deficiency appears to be associated with serogroup B disease only rarely. Individuals with recurrences of meningococcal disease, particularly those caused by non-B serogroups, should be assessed for complement deficiency by measurement of total hemolytic complement activity. There is also limited evidence that hyposplenism (through reduction in phagocytic capacity) and hypogammaglobulinemia (through absence of specific antibody) increase the risk of meningococcal disease. Genetic studies have revealed various associations with disease susceptibility, including complement and mannose-binding lectin deficiency, single-nucleotide polymorphisms in Toll-like receptor (TLR) 4 and complement factor H, and variants of Fc gamma receptors.

Factors that increase the chance of a susceptible individual's acquiring *N. meningitidis* via the respiratory route also increase the risk of meningococcal disease. Acquisition occurs through close contact with carriers as a result of overcrowding (e.g., in poor socioeconomic settings, in refugee camps, during the Hajj pilgrimage to Mecca, and during freshman-year residence in college dormitories) and certain social behaviors (e.g., attendance at bars and nightclubs, kissing). Secondary cases may occur in close contacts of an index case (e.g., household members and persons kissing the infected individual); the risk to these contacts may be as high as 1000 times the background rate in the population. Factors that damage the nasopharyngeal epithelium also increase the risk of both colonization with *N. meningitidis* and invasive disease. The most important of these factors are cigarette smoking (odds ratio, 4.1) and passive exposure to cigarette smoke. In addition, recent viral respiratory tract infection, infection with *Mycoplasma* species, and winter or the dry season have been associated with meningococcal disease; all of these factors presumably either increase the expression of adhesion molecules in the nasopharynx, thus enhancing meningococcal adhesion, or facilitate meningococcal invasion of the bloodstream.

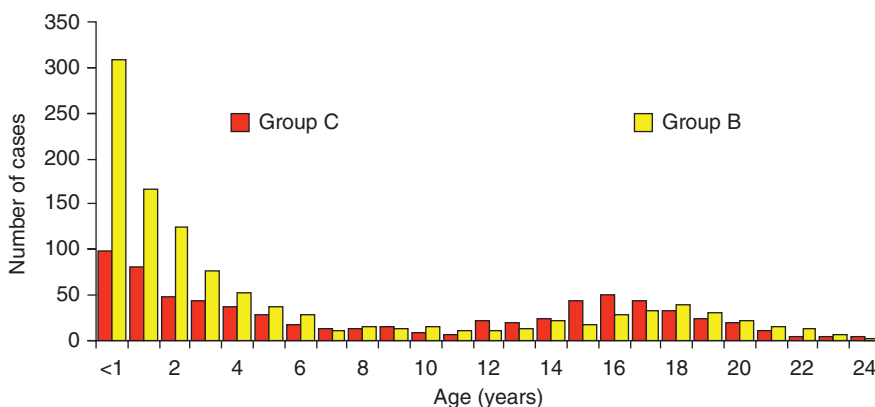


FIGURE 180-5 Age distribution of serogroups B and C meningococcal disease in England and Wales, 1998–1999. (Health Protection Agency, UK; www.hpa.org.uk.)

N. meningitidis has evolved as an effective colonizer of the human nasopharynx, with asymptomatic infection rates of >25% described in some series of adolescents and young adults and among residents of crowded communities. Point-prevalence studies reveal widely divergent rates of carriage for different types of meningococci. This variation suggests that some types may be adapted to a short duration of carriage with frequent transmission to maintain the population, while others may be less efficiently transmitted but may overcome this disadvantage by colonizing for a long period. Despite the high rates of carriage among adolescents and young adults, only ~10% of adults carry meningococci, and colonization is very rare in early childhood. Many of the same factors that increase the risk of meningococcal disease also increase the risk of carriage, including smoking, crowding, and respiratory viral infection. Colonization of the nasopharynx involves a series of interactions of meningococcal adhesins (e.g., Opa proteins and pili) with their ligands on the epithelial mucosa. *N. meningitidis* produces an IgA1 protease that is likely to reduce interruption of colonization by mucosal IgA.

Colonization should be considered the normal state of meningococcal infection, with an increased risk of invasion being the unfortunate consequence (for both host and organism) of adaptations of hyperinvasive meningococcal lineages. The meningococcal capsule is an important virulence factor: acapsular strains only very rarely cause invasive disease. The capsule provides resistance to phagocytosis and may be important in preventing desiccation during transmission between hosts. Antigenic diversity in surface structures and an ability to vary levels of their expression have probably evolved as important factors in maintaining meningococcal populations within and between individual hosts.

Invasion through the mucosa into the blood occurs rarely, usually within a few days of acquisition of an invasive strain by a susceptible individual. Only occasional cases of prolonged colonization prior to invasion have been documented. Once the organism is in the bloodstream, its growth may be limited if the individual is partially immune, although bacteremia may allow seeding of another site, such as the meninges or the joints. Alternatively, unchecked proliferation may continue, resulting in high bacterial counts in the circulation. During growth, meningococci release blebs of outer membrane (Fig. 180-1) containing outer-membrane proteins and LPS. Endotoxin binds cell-bound CD14 in association with TLR4 to initiate an inflammatory cascade with the release of high levels of various mediators, including tumor necrosis factor (TNF) α , soluble TNF receptor, interleukin (IL) 1, IL-1 receptor antagonist, IL-1 β , IL-6, IL-8, IL-10, plasminogen-activator inhibitor 1 (PAI-1), and leukemia inhibitory factor. Soluble CD14-bound endotoxin acts as a mediator of endothelial activation. The severity of meningococcal disease is related both to the levels of endotoxin in the blood and to the magnitude of the inflammatory response. The latter is determined to some extent by polymorphisms in the inflammatory response genes (and their inhibitors), and the release of the inflammatory cascade heralds the development of meningococcal septicemia (meningococcemia). Endothelial injury is central to many clinical features of meningococcemia, including increased vascular permeability, pathologic changes in vascular tone, loss of thromboresistance, intravascular coagulation, and myocardial dysfunction. Endothelial injury leads to increased vascular permeability (attributed to loss of glycosaminoglycans and endothelial proteins), with subsequent gross proteinuria. Leakage of fluid and electrolytes into the tissues from capillaries (“capillary leak syndrome”) leads to hypovolemia, tissue edema, and pulmonary edema. Initial compensation results in vasoconstriction and tachycardia, although cardiac output eventually falls. While resuscitation fluids may restore circulating volume, tissue edema will continue to increase, and, in the lung, the consequence may be respiratory failure.

Intravascular thrombosis (caused by activation of procoagulant pathways in association with upregulation of tissue factor on the endothelium) occurs in some patients with meningococcal disease and results in purpura fulminans and infarction of areas of skin or even of whole limbs. At the same time, multiple anticoagulant pathways

are downregulated through loss of endothelial thrombomodulin and protein C receptors and decreases in levels of antithrombin III, protein C, protein S, and tissue factor pathway inhibitor. Thrombolysis is also profoundly impaired in meningococcal sepsis through the release of high levels of PAI-1.

Shock in meningococcal septicemia appears to be attributable to a combination of factors, including hypovolemia, which results from the capillary leak syndrome secondary to endothelial injury, and myocardial depression, which is driven by hypovolemia, hypoxia, metabolic derangements (e.g., hypocalcemia), and cytokines (e.g., IL-6). Decreased perfusion of tissues as a result of intravascular thrombosis, vasoconstriction, tissue edema, and reduced cardiac output in meningococcal septicemia can cause widespread organ dysfunction, including renal impairment and—later in the disease—a decreased level of consciousness due to central nervous system involvement.

Bacteria that reach the meninges cause a local inflammatory response—with release of a spectrum of cytokines similar to that seen in septicemia—that presents clinically as meningitis and is thought to determine the severity of neuronal injury. Local endothelial injury may result in cerebral edema and rapid onset of raised intracranial pressure in some cases.

CLINICAL MANIFESTATIONS

As discussed above, the most common form of infection with *N. meningitidis* is asymptomatic carriage of the organism in the nasopharynx. Despite the location of infection in the upper airway, meningococcal pharyngitis is rarely reported; however, upper respiratory tract symptoms are common prior to presentation with invasive disease. It is not clear whether these symptoms relate to preceding viral infection (which may promote meningococcal acquisition) or to meningococcal acquisition itself. After acquiring the organism, susceptible individuals develop disease manifestations in 1–10 days (usually <4 days, although colonization for 11 weeks has been documented).

Along the spectrum of presentations of meningococcal disease, the most common clinical syndromes are meningitis and meningococcal septicemia. In fulminant cases, death may occur within hours of the first symptoms. Occult bacteremia is also recognized and, if untreated, progresses in two-thirds of cases to focal infection, including meningitis or septicemia. Meningococcal disease may also present as pneumonia, pyogenic arthritis or osteomyelitis, purulent pericarditis, endophthalmitis, conjunctivitis, primary peritonitis, or (rarely) urethritis. Perhaps because it is difficult to diagnose, pneumococcal pneumonia is not commonly reported but is associated with serogroups Y, W, and Z and appears most often to affect individuals >10 years of age.

Rash A nonblanching rash (petechial or purpuric) develops in >80% of cases of meningococcal disease; however, the rash is often absent early in the illness. Usually initially blanching in nature (macules, maculopapules, or urticaria) and indistinguishable from more common viral rashes, the rash of meningococcal infection becomes petechial or frankly purpuric over the hours after onset. In the most severe cases, large purpuric lesions develop (purpura fulminans). Some patients (including those with overwhelming sepsis) may have no rash. While petechial rash and fever are important signs of meningococcal disease, fewer than 10% of children (and, in some clinical settings, fewer than 1% of patients) with this presentation are found to have meningococcal disease. Most patients presenting with a petechial or purpuric rash have a viral infection (Table 180-2). The skin lesions exhibit widespread endothelial necrosis and occlusion of small vessels in the dermis and subcutaneous tissues, with a neutrophilic infiltrate.

Meningitis Meningococcal meningitis commonly presents as non-specific manifestations, including fever, vomiting, and (especially in infants and young children) irritability, and is indistinguishable from other forms of bacterial meningitis unless there is an associated petechial or purpuric rash, which occurs in two-thirds of cases. Headache is rarely reported in early childhood but is more common in later childhood and adulthood. When headache is present, the following features, in association with fever or a history of fever, are suggestive of bacterial meningitis: neck stiffness, photophobia, decreased

TABLE 180-2 COMMON CAUSES OF PETECHIAL OR PURPURIC RASHES

Enteroviruses
Influenza and other respiratory viruses
Measles virus
Epstein-Barr virus
Cytomegalovirus
Parvovirus
Deficiency of protein C or S (including postvaricella protein S deficiency)
Platelet disorders (e.g., idiopathic thrombocytopenic purpura, drug effects, bone marrow infiltration)
Henoch-Schönlein purpura, connective tissue disorders, trauma (including nonaccidental injuries in children)
Pneumococcal, streptococcal, staphylococcal, or gram-negative bacterial sepsis

level of consciousness, seizures or status epilepticus, and focal neurologic signs. Classic signs of meningitis, such as neck stiffness and photophobia, are often absent in infants and young children with bacterial meningitis, who more usually present with fever and irritability and may have a bulging fontanelle.

While 30–50% of patients present with a meningitis syndrome alone, up to 40% of meningitis patients also present with some features of septicemia. Most deaths from meningococcal meningitis alone (i.e., without septicemia) are associated with raised intracranial pressure presenting as a reduced level of consciousness, relative bradycardia and hypertension, focal neurologic signs, abnormal posturing, and signs of brainstem involvement—e.g., unequal, dilated, or poorly reactive pupils; abnormal eye movement; and impaired corneal responses (**Chap. 328**).

Septicemia Meningococcal septicemia alone accounts for up to 20% of cases of meningococcal disease. The condition may progress from early nonspecific symptoms to death within hours. Mortality rates among children with this syndrome have been high (25–40%), but early aggressive management (as discussed below) may reduce the figure to <10%. Early symptoms are nonspecific and suggest an influenza-like illness with fever, headache, and myalgia accompanied by vomiting and abdominal pain. As discussed above, the rash, if present, may appear to be viral early in the course until petechiae or purpuric lesions develop. Purpura fulminans occurs in severe cases, with multiple large purpuric lesions and signs of peripheral ischemia. Surveys of patients have indicated that limb pain, pallor (including a mottled appearance and cyanosis), and cold hands and feet may be prominent. Shock is manifested by tachycardia, poor peripheral perfusion, tachypnea, and oliguria. Decreased cerebral perfusion leads to confusion, agitation, or decreased level of consciousness. With progressive shock, multiorgan failure ensues; hypotension is a late sign in children, who more commonly present with compensated shock (tachycardia, poor peripheral perfusion, and normal blood pressure). Poor outcome is associated with an absence of meningism, hypotension, young age, coma, relatively low temperature (<38°C), leukopenia, and thrombocytopenia. Spontaneous hemorrhage (pulmonary, gastric, or cerebral) may result from consumption of coagulation factors and thrombocytopenia.

Chronic Meningococcemia Chronic meningococcemia, which is rarely recognized, presents as repeated episodes of petechial rash associated with fever, joint pain, features of arthritis, and splenomegaly that may progress to acute meningococcal septicemia if untreated. During the relapsing course, bacteremia characteristically clears without treatment and then recurs. The differential diagnosis includes bacterial endocarditis, acute rheumatic fever, Henoch-Schönlein purpura, infectious mononucleosis, disseminated gonococcal infection, and immune-mediated vasculitis. This condition has been associated with complement deficiencies in some cases and with inadequate sulfonamide therapy in others.



A study from the Netherlands found that half of isolates from patients with chronic meningococcemia had an underacylated lipid A (part of the surface LPS molecule) due to an *lpxL1* gene

mutation, which markedly reduces the inflammatory response to endotoxin.

Postmeningococcal Reactive Disease In a small proportion of patients, an immune complex disease develops ~4–10 days after the onset of meningococcal disease, with manifestations that include a maculopapular or vasculitic rash (2% of cases), arthritis (up to 8% of cases), iritis (1%), pericarditis, and/or polyserositis associated with fever. The immune complexes involve meningococcal polysaccharide antigen and result in immunoglobulin and complement deposition with an inflammatory infiltrate. These features resolve spontaneously without sequelae. It is important to recognize this condition since a new onset of fever and rash can lead to concerns about relapse of meningococcal disease and unnecessarily prolonged antibiotic treatment.

DIAGNOSIS

Like other invasive bacterial infections, meningococcal disease may produce elevations of the white blood cell (WBC) count and of values for inflammatory markers (e.g., C-reactive protein and procalcitonin levels or the erythrocyte sedimentation rate). Values may be normal or low in rapidly progressive disease, and a lack of rise in these laboratory test values does not exclude the diagnosis. However, in the presence of fever and a petechial rash, these elevations are suggestive of meningococcal disease. In patients with severe meningococcal septicemia, common laboratory findings include hypoglycemia, acidosis, hypokalemia, hypocalcemia, hypomagnesemia, hypophosphatemia, anemia, and coagulopathy.

Although meningococcal disease is often diagnosed on clinical grounds, in suspected meningococcal meningitis or meningococcemia, blood should routinely be sent for culture to confirm the diagnosis and to facilitate public health investigations; blood cultures are positive in up to 75% of cases. Culture media containing sodium polyanethol sulfonate, which may inhibit meningococcal growth, should be avoided. Meningococcal viability is reduced if there is a delay in transport of the specimen to the microbiology laboratory for culture or in plating of cerebrospinal fluid (CSF) samples. In countries where treatment with antibiotics before hospitalization is recommended for meningococcal disease, the majority of clinically suspected cases are culture negative. Real-time polymerase chain reaction (PCR) analysis of whole-blood samples increases the diagnostic yield by >40%, and results obtained with this method may remain positive for several days after administration of antibiotics. Indeed, in the United Kingdom, more than half of clinically suspected cases are currently identified by PCR.

Unless contraindications exist (raised intracranial pressure, uncorrected shock, disordered coagulation, thrombocytopenia, respiratory insufficiency, local infection, ongoing convulsions), lumbar puncture should be undertaken to identify and confirm the etiology of suspected meningococcal meningitis, whose presentation cannot be distinguished from that of meningitis of other bacterial causes. Some authorities have recommended a CT brain scan prior to lumbar puncture because of the risk of cerebral herniation in patients with raised intracranial pressure. However, a normal CT scan is not uncommon in the presence of raised intracranial pressure in meningococcal meningitis, and the decision to perform a lumbar puncture should be made on clinical grounds. CSF features of meningococcal meningitis (elevated protein level and WBC count, decreased glucose level) are indistinguishable from those of other types of bacterial meningitis unless a gram-negative diplococcus is identified. (Gram's staining is up to 80% sensitive for meningococcal meningitis.) CSF should be submitted for culture (sensitivity, 90%) and (where available) PCR analysis. CSF antigen testing with latex agglutination is insensitive and should be replaced by molecular diagnosis when possible.

Lumbar puncture should generally be avoided in meningococcal septicemia, as positioning for the procedure may critically compromise the patient's circulation in the context of hypovolemic shock. Delayed lumbar puncture may still be useful when the diagnosis is uncertain, particularly if molecular diagnostic technology is available.

In other types of focal infection, culture and PCR analysis of normally sterile body fluids (e.g., synovial fluid) may aid in the diagnosis. Although some authorities have recommended cultures of scrapings

1000 or aspirates from skin lesions, this procedure adds little to the diagnostic yield when compared with a combination of blood culture and PCR analysis. Urinary antigen testing also is insensitive, and serologic testing for meningococcal infection has not been adequately studied. Because *N. meningitidis* is a component of the normal human nasopharyngeal flora, identification of the organism on throat swabs has no diagnostic value.

TREATMENT MENINGOCOCCAL INFECTIONS

Death from meningococcal disease is associated most commonly with hypovolemic shock (meningococemia) and occasionally with raised intracranial pressure (meningococcal meningitis). Therefore, management should focus on the treatment of these urgent clinical issues in addition to the administration of specific antibiotic therapy. Delayed recognition of meningococcal disease or its associated physiologic derangements, together with inadequate emergency management, is associated with poor outcome. Since the disease is rare, protocols for emergency management have been developed (see www.meningitis.org).

Airway patency may be compromised if the level of consciousness is depressed as a result of shock (impaired cerebral perfusion) or raised intracranial pressure; this situation may require intervention. In meningococemia, pulmonary edema and pulmonary oligemia (presenting as hypoxia) require oxygen therapy or elective endotracheal intubation. In cases with shock, aggressive fluid resuscitation (with replacement of the circulating volume several times in severe cases) and inotropic support may be necessary to maintain cardiac output. If shock persists after volume resuscitation at 40 mL/kg, the risk of pulmonary edema is high, and elective intubation is recommended to improve oxygenation and decrease the work of breathing. Metabolic derangements, including hypoglycemia, acidosis, hypokalemia, hypocalcemia, hypomagnesemia, hypophosphatemia, anemia, and coagulopathy, should be anticipated and corrected. In the presence of raised intracranial pressure, management includes correction of coexistent shock and neurointensive care to maintain cerebral perfusion.

Empirical antibiotic therapy for suspected meningococcal disease consists of a third-generation cephalosporin such as ceftriaxone (75–100 mg/kg per day [maximum, 4 g/d] in one or two divided IV doses) or cefotaxime (200 mg/kg per day [maximum, 8 g/d] in four divided IV doses) to cover the various other (potentially penicillin-resistant) bacteria that may produce an indistinguishable clinical syndrome. Although unusual in most isolates, reduced meningococcal sensitivity to penicillin (a minimal inhibitory concentration of 0.12–1.0 µg/mL) has been reported widely.

Both meningococcal meningitis and meningococcal septicemia are conventionally treated for 7 days, although courses of 3–5 days may be equally effective. Furthermore, a single dose of ceftriaxone or an oily suspension of chloramphenicol has been used successfully in resource-poor settings. No data are available to guide the duration of treatment for meningococcal infection at other foci (e.g., pneumonia, arthritis); antimicrobial therapy is usually continued until clinical and laboratory evidence of infection has resolved. Cultures usually become sterile within 24 h of initiation of appropriate antibiotic chemotherapy.

The use of glucocorticoids for adjunctive treatment of meningococcal meningitis remains controversial since no relevant studies have had sufficient power to determine true efficacy. One large study in adults did indicate a trend toward benefit, and in clinical practice a decision to use glucocorticoids usually must precede a definite microbiologic diagnosis. Therapeutic doses of glucocorticoids are not recommended in meningococcal septicemia, but many intensivists recommend replacement glucocorticoid doses for patients who have refractory shock in association with impaired adrenal gland responsiveness.

Various other adjunctive therapies for meningococcal disease have been considered, but few have been subjected to clinical trials and none can currently be recommended. An antibody to LPS (HA1A) failed to confer a demonstrable benefit. Recombinant

bactericidal/permeability-increasing protein (which is not currently available) was tested in a study that had inadequate power to show an effect on mortality rates; however, there were trends toward lower mortality rates among patients who received a complete infusion, and this group also had fewer amputations, fewer blood-product transfusions, and a significantly improved functional outcome. Given that protein C concentrations are reduced in meningococcal disease, the use of activated protein C has been considered since a survival benefit was demonstrated in adult sepsis trials; however, trials in pediatric sepsis (of particular relevance for meningococcal disease) found no benefit and indicated a potential risk of bleeding complications with use of activated protein C.

The postmeningococcal immune-complex inflammatory syndrome has been treated with nonsteroidal anti-inflammatory agents until spontaneous resolution occurs.

COMPLICATIONS

About 10% of patients with meningococcal disease die despite the availability of antimicrobial therapy and other intensive medical interventions. The most common complication of meningococcal disease (10% of cases) is scarring after necrosis of purpuric skin lesions, for which skin grafting may be necessary. The lower limbs are most often affected; next in frequency are the upper limbs, the trunk, and the face. On average, 13% of the skin surface area is involved. Amputations are necessary in 1–2% of survivors of meningococcal disease because of a loss of tissue viability after peripheral ischemia or compartment syndromes. Unless there is local infection, amputation should usually be delayed to allow the demarcation between viable and nonviable tissue to become apparent. Approximately 5% of patients with meningococcal disease suffer hearing loss, and 7% have neurologic complications. In one study pain was reported by 21% of survivors, and in a recent analysis of serogroup B meningococcal disease (the MOSAIC study) as many as one-quarter of survivors had psychological disorders. In some investigations, the rate of complications is higher for serogroup C disease (mostly associated with the ST11 clone) than for serogroup B disease. In patients with severe hypovolemic shock, renal perfusion may be impaired and prerenal failure is common, but permanent renal replacement therapy is rarely needed.

Several studies suggest adverse psychosocial outcomes after meningococcal disease, with reduced quality of life, lowered self-esteem, and poorer neurologic development, including increased rates of attention deficit/hyperactivity disorder and special educational needs. Other studies have not found evidence of such outcomes.

PROGNOSIS

Several prognostic scoring systems have been developed to identify patients with meningococcal disease who are least likely to survive. Factors associated with a poorer prognosis are shock; young age (infancy), old age, and adolescence; coma; purpura fulminans; disseminated intravascular coagulation; thrombocytopenia; leukopenia; absence of meningitis; metabolic acidosis; low plasma concentrations of antithrombin and proteins S and C; high blood levels of PAI-1; and a low erythrocyte sedimentation rate or C-reactive protein level. The Glasgow Meningococcal Septicaemia Prognostic Score (GMSPS) is probably the best-performing scoring system studied so far and may be clinically useful for severity assessment in meningococcal disease. However, scoring systems do not direct the clinician to specific interventions, and the priority in management should be recognition of compromised airways, breathing, or circulation and direct, urgent intervention. Most patients improve rapidly with appropriate antibiotics and supportive therapy. Fulminant meningococemia is more likely to result in death or ischemic skin loss than is meningitis; optimal emergency management may reduce mortality rates among the most severely affected patients.

PREVENTION

Since mortality rates in meningococcal disease remain high despite improvements in intensive care management, immunization is the

only rational approach to prevention at a population level. Secondary cases are common among household and “kissing” contacts of cases, and secondary prophylaxis with antibiotic therapy is widely recommended for these contacts (see below).

Polysaccharide Vaccines Purified meningococcal capsular polysaccharide has been used for immunization since the 1960s. Meningococcal polysaccharide vaccines are currently formulated as either bivalent (serogroups A and C) or quadrivalent (serogroups A, C, Y, and W), with 50 µg of each polysaccharide per dose. Local reactions (erythema, induration, and tenderness) may occur in up to 40% of vaccinees, but serious adverse events (including febrile convulsions in young children) are very rarely reported. In adults, the vaccines are immunogenic, but immunity appears to be relatively short-lived (with antibody levels above baseline for only 2–10 years), and booster doses do not induce a further rise in antibody concentration. Indeed, a state of immunologic hyporesponsiveness has been widely reported to follow booster doses of plain polysaccharide vaccines. The repeating units of these vaccines cross-link B cell receptors to drive specific memory B cells to become plasma cells and produce antibody. Because meningococcal polysaccharides are T cell-independent antigens, no memory B cells are produced after immunization, and the memory B-cell pool is depleted such that fewer polysaccharide-specific cells are available to respond to a subsequent dose of vaccine (Fig. 180-6). The clinical relevance of hyporesponsiveness is unknown. Plain polysaccharide

vaccines generally are not immunogenic in early childhood, possibly because marginal-zone B cells are involved in polysaccharide responses and maturation of the splenic marginal zone is not complete until 18 months to 2 years of age. The efficacy of the meningococcal serogroup C component is >90% in young adults; no efficacy data are available for the serogroup Y and W polysaccharides in this age group.

Group A meningococcal polysaccharides are exceptional in that they have been found to be effective in preventing disease at all ages. Two doses administered 2–3 months apart to children 3–18 months of age or a single dose administered to older children or adults has a protective efficacy rate of >95%. The vaccine has been widely used in the control of meningococcal disease in the African meningitis belt. The duration of protection appears to be only 3–5 years.

There is no meningococcal serogroup B plain polysaccharide vaccine because α -2,8-*N*-acetylneuraminic acid is expressed on the surface of neural cells in the fetus such that the B polysaccharide is perceived as “self” and therefore is not immunogenic in humans.

Conjugate Vaccines The poor immunogenicity of plain polysaccharide vaccines in infancy has been overcome by chemical conjugation of the polysaccharides to a carrier protein (CRM₁₉₇, tetanus toxoid, or diphtheria toxoid). Conjugates that contain monovalent serogroup C polysaccharide and quadrivalent vaccines with A, C, Y, and W polysaccharides have been developed, as have vaccines including various other antigen combinations (e.g., tetanus conjugates with serogroup C

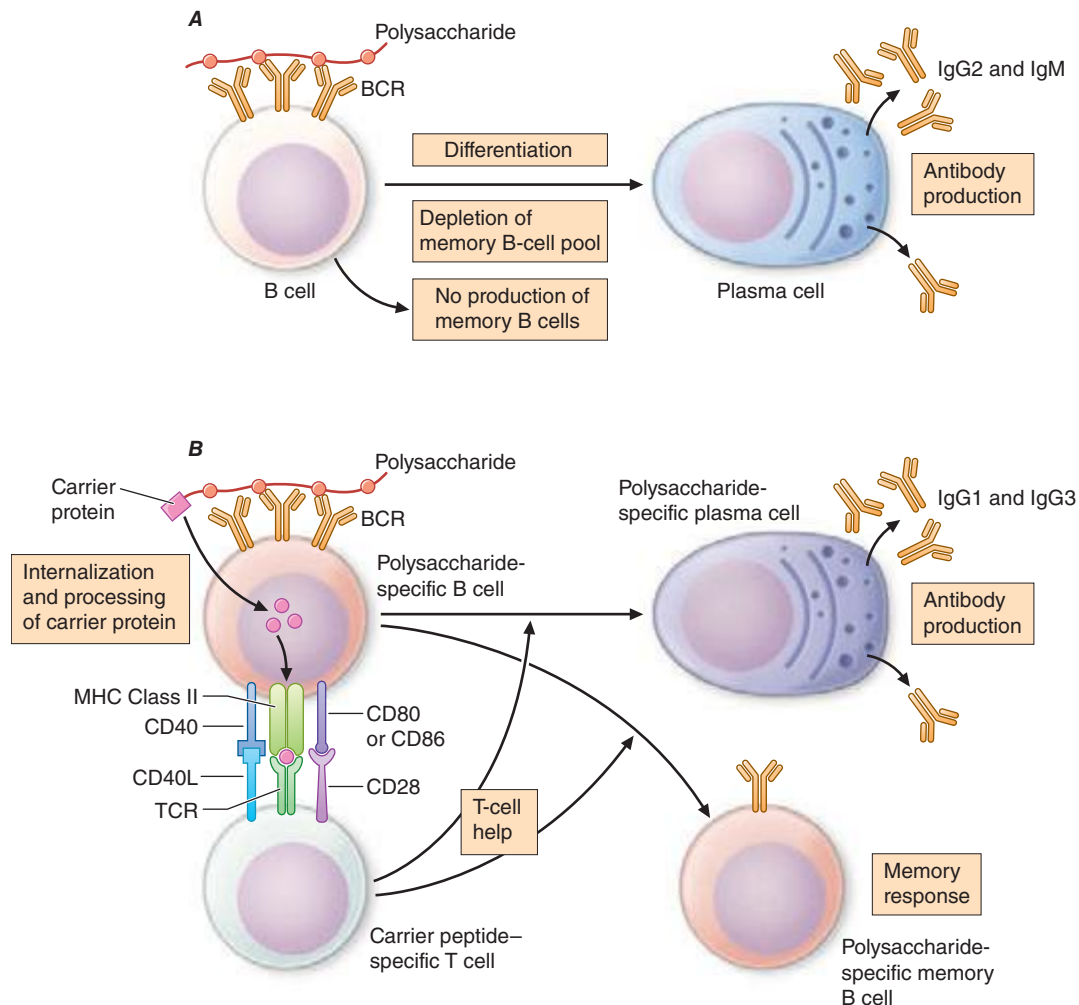


FIGURE 180-6 **A.** Polysaccharides from the encapsulated bacteria that cause disease in early childhood stimulate B cells by cross-linking the BCR and driving the production of immunoglobulins. There is no production of memory B cells, and the B-cell pool may be depleted by this process such that subsequent immune responses are decreased. **B.** The carrier protein from protein-polysaccharide conjugate vaccines is processed by the polysaccharide-specific B cell, and peptides are presented to carrier peptide-specific T cells, with the consequent production of both plasma cells and memory B cells. BCR, B-cell receptor; MHC, major histocompatibility complex; TCR, T-cell receptor. (Reprinted from AJ Pollard et al: *Nat Rev Immunol* 9:213, 2009.)

1002 and/or Y polysaccharide with *Haemophilus influenzae* type b polysaccharide). After immunization, peptides from the carrier protein are conventionally thought to be presented to peptide-specific T cells in association with major histocompatibility complex (MHC) class II molecules (some recent data suggesting that carrier protein peptide may actually be presented in association with an oligosaccharide and MHCII) by polysaccharide-specific B cells; the result is a T cell-dependent immune response that allows production of antibody and generation of an expanded B-cell memory pool. Unlike responses to booster doses of plain polysaccharides, responses to booster doses of conjugate vaccines have the characteristics of memory responses. Indeed, conjugate vaccines overcome the hyporesponsiveness induced by plain polysaccharides by replenishing the memory pool. The reactogenicity of conjugate vaccines is similar to that of plain polysaccharide vaccines.



The first widespread use of serogroup C meningococcal conjugate vaccine (MenC) came in 1999 in the United Kingdom after a rise in serogroup C disease. A mass vaccination campaign involving all individuals <19 years of age was undertaken, and the number of laboratory-confirmed serogroup C cases fell from 955 in 1998–1999 to just 29 in 2011–2012. The effectiveness of the immunization program was attributed both to direct protection of immunized persons and to reduced transmission of the organism in the population as a result of decreased rates of colonization among the immunized (herd immunity). Data on immunogenicity and effectiveness have shown that the duration of protection is short when the vaccine is administered in early childhood; thus booster doses are needed to maintain population immunity. In contrast, immunity after a dose of vaccine given in adolescence appears to be prolonged.

The first quadrivalent conjugate meningococcal vaccine containing A, C, Y, and W polysaccharides conjugated to diphtheria toxoid was initially recommended for all children >11 years of age in the United States in 2005. In 2007 the license was extended to high-risk children 2–10 years of age. In the same year, the vaccine was licensed in Canada for persons 2–55 years of age. Uptake was slow, but current U.S. data suggest an efficacy rate of 82% in the first year after vaccination, with waning to 59% at 3–6 years after vaccination. Limited data from the U.S. Vaccine Adverse Events Reporting System indicated that there might be a short-term increase in the risk of Guillain-Barré syndrome after immunization with the diphtheria conjugate vaccine; however, further investigation has not confirmed this finding. Quadrivalent conjugate vaccines with tetanus or CRM₁₉₇ as carrier protein are now available in many countries.

A monovalent serogroup A vaccine, manufactured in India, was licensed in 2010 and rolled out to countries in the sub-Saharan African meningitis belt. There is strong evidence that this vaccine has been highly effective in controlling epidemic meningococcal disease in the region, with some evidence of a >90% reduction in disease in vaccinated populations. However, disease caused by serogroup X and W persists.



Vaccines Based on Subcapsular Antigens The lack of immunogenicity of the serogroup B capsule has led to the development of vaccines based on subcapsular antigens. Various surface components have been studied in early-phase clinical trials. Outer-membrane vesicles (OMVs) containing outer-membrane proteins, phospholipid, and LPS can be extracted from cultures of *N. meningitidis* by detergent treatment (Fig. 180-7). OMVs prepared in this way were used in efficacy trials with a Norwegian outbreak strain and reduced the incidence of group B disease among 14- to 16-year-old schoolchildren by 53%. Similarly, OMV vaccines constructed from local outbreak strains in Cuba and New Zealand have had reported efficacy rates of >70%. These OMV vaccines appear to produce strain-specific immune responses, with only limited cross-protection, and are therefore best suited to clonal outbreaks (e.g., those in Cuba and New Zealand as well as others in Norway and the province of Normandy in France).

Several purified surface proteins have been evaluated in phase I clinical trials but have not yet been developed further because of antigenic variability or poor immunogenicity (e.g., transferrin-binding

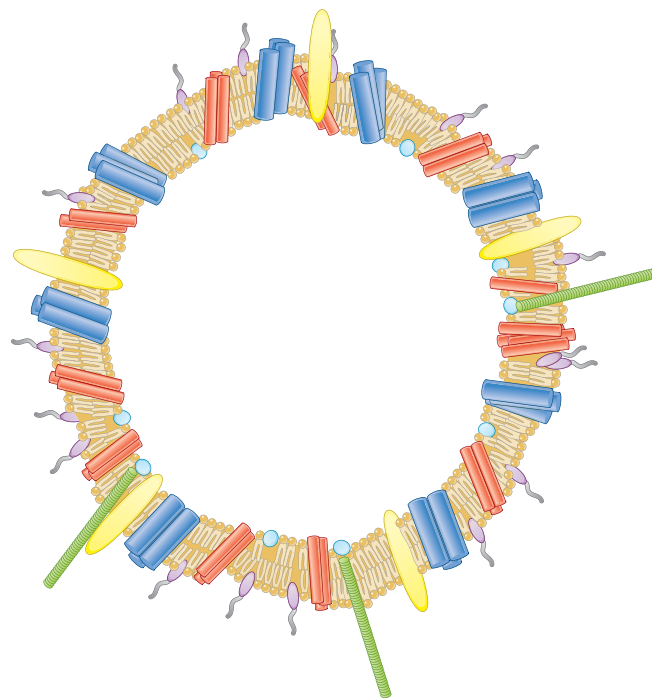


FIGURE 180-7 Illustration of meningococcal outer-membrane vesicle containing outer-membrane structures.

proteins, neisserial surface protein A). Other vaccine candidates have been identified since sequencing of the meningococcal genome. A combination vaccine that includes the New Zealand OMV vaccine and three recombinant proteins (neisserial adhesin A, factor H-binding protein, and neisserial heparin-binding antigen) is immunogenic in infancy and has been licensed for use in Europe and Australia. Recommendations for its use are pending. Finally, a highly immunogenic vaccine based on two variants of the lipoprotein factor H-binding protein is undergoing clinical evaluation

MANAGEMENT OF CONTACTS

Close (household and kissing) contacts of individuals with meningococcal disease are at increased risk (up to 1000 times the rate for the general population) of developing secondary disease; a secondary case follows as many as 3% of sporadic cases. About one-fifth of secondary cases are actually co-primary cases—i.e., cases that occur soon after the primary case and in which transmission is presumed to have originated from the same third party. The rate of secondary cases is highest during the week after presentation of the index case. The risk falls rapidly but remains above baseline for up to 1 year after the index case; 30% of secondary cases occur in the first week, 20% in the second week, and most of the remainder over the next 6 weeks. In outbreaks of meningococcal disease, mass prophylaxis has been used; however, limited data support population intervention, and significant concerns have arisen about adverse events and the development of resistance. For these reasons, prophylaxis is usually restricted to (1) persons at greatest risk who are intimate and/or household contacts of the index case and (2) health care workers who have been directly exposed to respiratory secretions. In most cases, members of wider communities (e.g., at schools or colleges) are not offered prophylaxis.



The aim of prophylaxis is to eradicate colonization of close contacts with the strain that has caused invasive disease in the index case. Prophylaxis should be given to all contacts at the same time to avoid recolonization by meningococci transmitted from untreated contacts and should also be used as soon as possible to treat early disease in secondary cases. If the index patient is treated with an antibiotic that does not reliably clear colonization (e.g., penicillin), he or she should be given a prophylactic agent at the end of treatment to prevent relapse or onward transmission. Although rifampin has been

most widely used and studied, it is not the optimal agent because it fails to eradicate carriage in 15–20% of cases, rates of adverse events have been high, compliance is affected by the need for four doses, and emerging resistance has been reported. Ceftriaxone as a single IM or IV injection is highly (97%) effective in carriage eradication and can be used at all ages and in pregnancy. Reduced susceptibility of isolates to ceftriaxone has occasionally been reported. Ciprofloxacin or ofloxacin is preferred in some countries; these agents are highly effective and can be administered by mouth but are not recommended in pregnancy. Resistance to fluoroquinolones has been reported in some meningococci in North America, Europe, and Asia.

In documented serogroup A, C, Y, or W disease, contacts may be offered immunization (preferably with a conjugate vaccine) in addition to chemoprophylaxis to provide protection beyond the duration of antibiotic therapy. Mass vaccination has been used successfully to control disease during outbreaks in closed communities (educational and military establishments) as well as during epidemics in open communities.

181 Gonococcal Infections

Sanjay Ram, Peter A. Rice

DEFINITION

Gonorrhea is a sexually transmitted infection (STI) of epithelium and commonly manifests as cervicitis, urethritis, proctitis, and conjunctivitis. If untreated, infections at these sites can lead to local complications such as endometritis, salpingitis, tuboovarian abscess, Bartholinitis, peritonitis, and perihepatitis in female patients; periurethritis and epididymitis in male patients; and ophthalmia neonatorum in newborns. Disseminated gonococemia is an uncommon event whose manifestations include skin lesions, tenosynovitis, arthritis, and (in rare cases) endocarditis or meningitis.

MICROBIOLOGY

Neisseria gonorrhoeae is a gram-negative, nonmotile, non-spore-forming organism that grows singly and in pairs (i.e., as monococci and diplococci, respectively). Exclusively a human pathogen, the gonococcus contains, on average, three genome copies per coccal unit; this polyploidy permits a high level of antigenic variation and the survival of the organism in its host. Gonococci, like all other *Neisseria* species, are oxidase positive. They are distinguished from other neisseriae by their ability to grow on selective media and to use glucose but not maltose, sucrose, or lactose.

EPIDEMIOLOGY



The incidence of gonorrhea has declined significantly in the United States, but there were still ~311,000 newly reported cases in 2012. Gonorrhea remains a major public health problem worldwide, is a significant cause of morbidity in developing countries, and may play a role in enhancing transmission of HIV.

Gonorrhea predominantly affects young, nonwhite, unmarried, less educated members of urban populations. The number of reported cases probably represents half of the true number of cases—a discrepancy resulting from underreporting, self-treatment, and nonspecific treatment without a laboratory-proven diagnosis. The number of reported new cases of gonorrhea in the United States rose from ~250,000 in the early 1960s to a high of 1.01 million in 1978. The recorded incidence of gonorrhea in modern times peaked in 1975, with 468 reported new cases per 100,000 population in the United States. This peak was attributable to the interaction of several variables, including improved accuracy of diagnosis, changes in patterns of contraceptive use, and changes in sexual behavior. The incidence of the disease has since declined gradually and is currently estimated at 120 cases per 100,000,

a figure that is still the highest among industrialized countries. A further decline in the overall incidence of gonorrhea in the United States over the past quarter-century may reflect increased condom use resulting from public health efforts to curtail HIV transmission. At present, the attack rate in the United States is highest among 15- to 19-year-old women and 20- to 24-year-old men; 60% of all reported cases occur in the preceding two groups together. From the standpoint of ethnicity, rates are highest among African Americans and lowest among persons of Asian or Pacific Island descent.

The incidence of gonorrhea is higher in developing countries than in industrialized nations. The exact incidence of any STI is difficult to ascertain in developing countries because of limited surveillance and variable diagnostic criteria. Studies in Africa have clearly demonstrated that nonulcerative STIs such as gonorrhea (in addition to ulcerative STIs) are an independent risk factor for the transmission of HIV (Chap. 226).

Gonorrhea is transmitted from males to females more efficiently than in the opposite direction. The rate of transmission to a woman during a single unprotected sexual encounter with an infected man is ~50–70%. Oropharyngeal gonorrhea occurs in ~20% of women who practice fellatio with infected partners. Transmission in either direction by cunnilingus is rare.

In any population, there exists a small minority of individuals who have high rates of new-partner acquisition. These “core-group members” or “high-frequency transmitters” are vital in sustaining STI transmission at the population level. Another instrumental factor in sustaining gonorrhea in the population is the large number of infected individuals who are asymptomatic or have minor symptoms that are ignored. These persons, unlike symptomatic individuals, may not cease sexual activity and therefore continue to transmit the infection. This situation underscores the importance of contact tracing and empirical treatment of the sex partners of index cases.

PATHOGENESIS, IMMUNOLOGY, AND ANTIMICROBIAL RESISTANCE

Outer-Membrane Proteins • Pili Fresh clinical isolates of *N. gonorrhoeae* initially form piliated (fimbriated) colonies distinguishable on translucent agar. Pilus expression is rapidly switched off with unselected subculture because of rearrangements in pilus genes. This change is a basis for antigenic variation of gonococci. Piliated strains adhere better to cells derived from human mucosal surfaces and are more virulent in organ culture models and human inoculation experiments than nonpiliated variants. In a fallopian tube explant model, pili mediate gonococcal attachment to nonciliated columnar epithelial cells. This event initiates gonococcal phagocytosis and transport through these cells to intercellular spaces near the basement membrane or directly into the subepithelial tissue. Pili are also essential for genetic competence and transformation of *N. gonorrhoeae*, which permit horizontal transfer of genetic material between different gonococcal lineages in vivo.

OPACITY-ASSOCIATED PROTEIN Another gonococcal surface protein that is important in adherence to epithelial cells is opacity-associated protein (Opa, formerly called protein II). Opa contributes to intergonococcal adhesion, which is responsible for the opaque nature of gonococcal colonies on translucent agar and the organism’s adherence to a variety of eukaryotic cells, including polymorphonuclear leukocytes (PMNs). Certain Opa variants promote invasion of epithelial cells, and this effect has been linked with the ability of Opa to bind vitronectin, glycosaminoglycans, and several members of the carcinoembryonic antigen-related cell adhesion molecule (CEACAM) receptor family. *N. gonorrhoeae* Opa proteins that bind CEACAM1, which is expressed by primary CD4+ T lymphocytes, suppress the activation and proliferation of these lymphocytes. This phenomenon may serve to explain the transient decrease in CD4+ T lymphocyte counts associated with gonococcal infection. Select Opa proteins can engage CEACAM3, which is expressed on neutrophils, with consequent nonopsonic phagocytosis (i.e., phagocytosis independent of antibody and complement) and killing of bacteria.

1004 PORIN Porin (previously designated protein I) is the most abundant gonococcal surface protein, accounting for >50% of the organism's total outer-membrane protein. Porin molecules exist as trimers that provide anion-transporting aqueous channels through the otherwise hydrophobic outer membrane. Porin exhibits stable interstrain antigenic variation and forms the basis for gonococcal serotyping. Two main serotypes have been identified: PorB.1A strains are often associated with disseminated gonococcal infection (DGI), whereas PorB.1B strains usually cause local genital infections only. DGI strains are generally resistant to the killing action of normal human serum and do not incite a significant local inflammatory response; therefore, they may not cause symptoms at genital sites. These characteristics may be related to the ability of PorB.1A strains to bind to complement-inhibitory molecules, resulting in a diminished inflammatory response. Porin can translocate to the cytoplasmic membrane of host cells—a process that could initiate gonococcal endocytosis and invasion.

OTHER OUTER-MEMBRANE PROTEINS Other notable outer-membrane proteins include H.8, a lipoprotein that is present in high concentration on the surface of all gonococcal strains and is an excellent target for antibody-based diagnostic testing. Transferrin-binding proteins (Tbp1 and Tbp2) and lactoferrin-binding protein are required for scavenging iron from transferrin and lactoferrin in vivo. Transferrin and iron have been shown to enhance the attachment of iron-deprived *N. gonorrhoeae* to human endometrial cells. IgA1 protease is produced by *N. gonorrhoeae* and may protect the organism from the action of mucosal IgA.

Lipooligosaccharide Gonococcal lipooligosaccharide (LOS) consists of a lipid A and a core oligosaccharide that lacks the repeating O-carbohydrate antigenic side chain seen in other gram-negative bacteria (Chap. 145e). Gonococcal LOS possesses marked endotoxic activity and contributes to the local cytotoxic effect in a fallopian tube model. LOS core sugars undergo a high degree of phase variation under different conditions of growth; this variation reflects genetic regulation and expression of glycotransferase genes that dictate the carbohydrate structure of LOS. These phenotypic changes may affect interactions of *N. gonorrhoeae* with elements of the humoral immune system (antibodies and complement) and may also influence direct binding of organisms to both professional phagocytes and nonprofessional phagocytes (epithelial cells). For example, gonococci that are sialylated at their LOS sites bind complement factor H and inhibit the alternative pathway of complement. LOS sialylation may also decrease nonopsonic Op-mediated association with neutrophils and inhibit the oxidative burst in PMNs. The binding of the unsialylated terminal lactosamine residue of LOS to an asialoglycoprotein receptor on male epithelial cells facilitates adherence and subsequent gonococcal invasion of these cells. Moreover, oligosaccharide structures in LOS can modulate host immune responses. For example, the terminal monosaccharide expressed by LOS determines the C-type lectin receptor on dendritic cells that is targeted by the bacteria. In turn, the specific C-type lectin receptor engaged influences whether a T_H1- or T_H2-type response is elicited; the latter response may be less favorable for clearance of gonococcal infection.

Host Factors In addition to gonococcal structures that interact with epithelial cells, host factors seem to be important in mediating entry of gonococci into nonphagocytic cells. Activation of phosphatidylcholine-specific phospholipase C and acidic sphingomyelinase by *N. gonorrhoeae*, which results in the release of diacylglycerol and ceramide, is a requirement for the entry of *N. gonorrhoeae* into epithelial cells. Ceramide accumulation within cells leads to apoptosis, which may disrupt epithelial integrity and facilitate entry of gonococci into subepithelial tissue. Release of chemotactic factors as a result of complement activation contributes to inflammation, as does the toxic effect of LOS in provoking the release of inflammatory cytokines.

The importance of humoral immunity in host defenses against neisserial infections is best illustrated by the predisposition of persons deficient in terminal complement components (C5 through C9) to recurrent bacteremic gonococcal infections and to recurrent meningococcal meningitis or meningococcemia. Gonococcal porin induces

T cell–proliferative responses in persons with urogenital gonococcal disease. A significant increase in porin-specific interleukin (IL) 4–producing CD4+ as well as CD8+ T lymphocytes is seen in individuals with mucosal gonococcal disease. A portion of these lymphocytes that show a porin-specific T_H2-type response could traffic to mucosal surfaces and play a role in immune protection against the disease. Few data clearly indicate that protective immunity is acquired from a previous gonococcal infection, although bactericidal and opsonophagocytic antibodies to porin and LOS may offer partial protection. On the other hand, women who are infected and acquire high levels of antibody to another outer-membrane protein, Rmp (reduction modifiable protein, formerly called protein III), may be especially likely to become reinfected with *N. gonorrhoeae* because Rmp antibodies block the effect of bactericidal antibodies to porin and LOS. Rmp shows little, if any, interstrain antigenic variation; therefore, Rmp antibodies potentially may block antibody-mediated killing of all gonococci. The mechanism of blocking has not been fully characterized, but Rmp antibodies may noncompetitively inhibit binding of porin and LOS antibodies because of the proximity of these structures in the gonococcal outer membrane. In male volunteers who have no history of gonorrhea, the net effect of these events may influence the outcome of experimental challenge with *N. gonorrhoeae*. Because Rmp bears extensive homology to enterobacterial OmpA and meningococcal class 4 proteins, it is possible that these blocking antibodies result from prior exposure to cross-reacting proteins from these species and also play a role in first-time infection with *N. gonorrhoeae*.

Gonococcal Resistance to Antimicrobial Agents It is no surprise that *N. gonorrhoeae*, with its remarkable capacity to alter its antigenic structure and adapt to changes in the microenvironment, has become resistant to numerous antibiotics. The first effective agents against gonorrhea were the sulfonamides, which were introduced in the 1930s and became ineffective within a decade. Penicillin was then used as the drug of choice for the treatment of gonorrhea. By 1965, 42% of gonococcal isolates had developed low-level resistance to penicillin G. Resistance due to the production of penicillinase arose later.



Gonococci become fully resistant to antibiotics either by chromosomal mutations or by acquisition of R factors (plasmids). Two types of chromosomal mutations have been described. The first type, which is drug specific, is a single-step mutation leading to high-level resistance. The second type involves mutations at several chromosomal loci that combine to determine the level as well as the pattern of resistance. Strains with mutations in chromosomal genes were first observed in the late 1950s. As recently as 2007, chromosomal mutations accounted for resistance to penicillin, tetracycline, or both in ~16% of strains surveyed in the United States.

β-Lactamase (penicillinase)–producing strains of *N. gonorrhoeae* (PPNG) carrying plasmids with the P_c determinant had rapidly spread worldwide by the early 1980s. *N. gonorrhoeae* strains with plasmid-borne tetracycline resistance (TRNG) can mobilize some β-lactamase plasmids, and PPNG and TRNG occur together, sometimes along with strains exhibiting chromosomally mediated resistance (CMRNG). Penicillin, ampicillin, and tetracycline are no longer reliable for the treatment of gonorrhea and should not be used.



Quinolone-containing regimens were also recommended for treatment of gonococcal infections; the fluoroquinolones offered the advantage of antichlamydial activity when administered for 7 days. However, quinolone-resistant *N. gonorrhoeae* (QRNG) appeared soon after these agents were first used to treat gonorrhea. QRNG is particularly common in the Pacific Islands (including Hawaii) and Asia, where, in certain areas, all gonococcal strains are now resistant to quinolones. At present, QRNG is also common in parts of Europe and the Middle East. In the United States, QRNG has been identified in midwestern and eastern areas as well as in states on the Pacific coast, where resistant strains were first seen. Alterations in DNA gyrase and topoisomerase IV have been implicated as mechanisms of fluoroquinolone resistance.

Resistance to spectinomycin, which has been used in the past as an alternative agent, has been reported. Because this agent usually is not

associated with resistance to other antibiotics, spectinomycin can be reserved for use against multidrug-resistant strains of *N. gonorrhoeae*. Nevertheless, outbreaks caused by strains resistant to spectinomycin have been documented in Korea and England when the drug has been used for primary treatment of gonorrhea.

Third-generation cephalosporins have remained highly effective as single-dose therapy for gonorrhea, but the recent isolation of strains highly resistant to ceftriaxone (minimal inhibitory concentrations [MICs], 2 µg/mL) in Japan and some European countries is cause for concern. Even though the MICs of ceftriaxone against certain strains may reach 0.015–0.125 µg/mL (higher than the MICs of 0.0001–0.008 µg/mL for fully susceptible strains), these levels are greatly exceeded in the blood, the urethra, and the cervix when the routinely recommended parenteral dose of ceftriaxone is administered. The rising MICs of oral cefixime (the previously recommended alternative oral third-generation cephalosporin) against *N. gonorrhoeae*, combined with this drug's limited capacity to reach levels sufficiently higher than MICs in the blood, the urethra, the cervix, and especially the pharynx, have resulted in the removal of cefixime from the list of first-line agents for treatment of uncomplicated gonorrhea. All *N. gonorrhoeae* strains with reduced susceptibility to ceftriaxone and cefixime (i.e., cephalosporin-intermediate/resistant strains) contain (1) a mosaic *penA* allele, which is the principal resistance determinant and encodes a penicillin-binding protein (PBP2) whose sequence differs in 60 amino acids from that of wild-type PBP2, and (2) additional genetic resistance determinants that are also required for high-level penicillin resistance.

CLINICAL MANIFESTATIONS

Gonococcal Infections in Men Acute urethritis is the most common clinical manifestation of gonorrhea in male patients. The usual incubation period after exposure is 2–7 days, although the interval can be longer and some men remain asymptomatic. Strains of the PorB.1A serotype tend to cause a greater proportion of cases of mild and asymptomatic urethritis than do PorB.1B strains. Urethral discharge and dysuria, usually without urinary frequency or urgency, are the major symptoms. The discharge initially is scant and mucoid but becomes profuse and purulent within a day or two. Gram's staining of the urethral discharge may reveal PMNs and gram-negative intracellular moncocci and diplococci (Fig. 181-1). The clinical manifestations of gonococcal urethritis are usually more severe and overt than those of nongonococcal urethritis, including urethritis caused by *Chlamydia trachomatis* (Chap. 213); however, exceptions are common, and it is often impossible to differentiate the causes of urethritis on clinical grounds alone. The majority of cases of urethritis seen in the United States today are not caused by *N. gonorrhoeae* and/or *C. trachomatis*. Although a number of other organisms may be responsible, many cases do not have a specific etiologic agent identified.

Most symptomatic men with gonorrhea seek treatment and cease to be infectious. The remaining men, who are largely asymptomatic, accumulate in number over time and constitute about two-thirds of all infected men at any point in time; together with men incubating

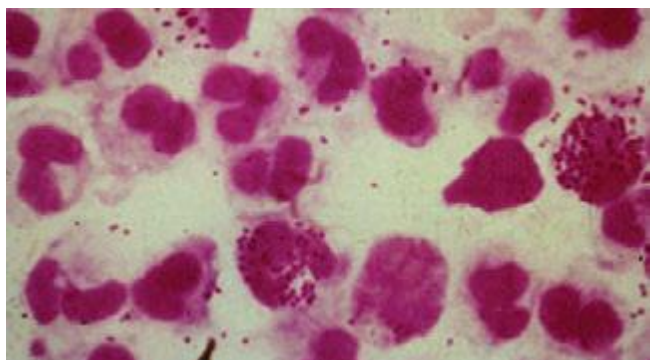


FIGURE 181-1 Gram's stain of urethral discharge from a male patient with gonorrhea shows gram-negative intracellular moncocci and diplococci. (From the Public Health Agency of Canada.)

the organism (who shed the organism but are asymptomatic), they serve as the source of spread of infection. Before the antibiotic era, symptoms of urethritis persisted for ~8 weeks. Epididymitis is now an uncommon complication, and gonococcal prostatitis occurs rarely, if at all. Other unusual local complications of gonococcal urethritis include edema of the penis due to dorsal lymphangitis or thrombophlebitis, submucous inflammatory "soft" infiltration of the urethral wall, periurethral abscess or fistula, inflammation or abscess of Cowper's gland, and seminal vesiculitis. Balanitis may develop in uncircumcised men.

Gonococcal Infections in Women • GONOCOCCAL CERVICITIS Mucopurulent cervicitis is a common STI diagnosis in American women and may be caused by *N. gonorrhoeae*, *C. trachomatis*, and other organisms, including *Mycoplasma genitalium* (Chap. 212). Cervicitis may coexist with candidal or trichomonal vaginitis. *N. gonorrhoeae* primarily infects the columnar epithelium of the cervical os. Bartholin's glands occasionally become infected.

Women infected with *N. gonorrhoeae* usually develop symptoms. However, the women who either remain asymptomatic or have only minor symptoms may delay in seeking medical attention. These minor symptoms may include scant vaginal discharge issuing from the inflamed cervix (without vaginitis or vaginosis per se) and dysuria (often without urgency or frequency) that may be associated with gonococcal urethritis. Although the incubation period of gonorrhea is less well defined in women than in men, symptoms usually develop within 10 days of infection and are more acute and intense than those of chlamydial cervicitis.

The physical examination may reveal a mucopurulent discharge (mucopus) issuing from the cervical os. Because Gram's stain is not sensitive for the diagnosis of gonorrhea in women, specimens should be submitted for culture or a nonculture assay (see "Laboratory Diagnosis," below). Edematous and friable cervical ectopy and endocervical bleeding induced by gentle swabbing are more often seen in chlamydial infection. Gonococcal infection may extend deep enough to produce dyspareunia and lower abdominal or back pain. In such cases, it is imperative to consider a diagnosis of pelvic inflammatory disease (PID) and to administer treatment for that disease (Chaps. 163 and 213).

N. gonorrhoeae may also be recovered from the urethra and rectum of women with cervicitis, but these are rarely the only infected sites. Urethritis in women may produce symptoms of internal dysuria, which is often attributed to "cystitis." Pyuria in the absence of bacteriuria seen on Gram's stain of unspun urine, accompanied by urine cultures that fail to yield >10² colonies of bacteria usually associated with urinary tract infection, signifies the possibility of urethritis due to *C. trachomatis*. Urethral infection with *N. gonorrhoeae* may also occur in this context, but in this instance urethral cultures are usually positive.

GONOCOCCAL VAGINITIS The vaginal mucosa of healthy women is lined by stratified squamous epithelium and is rarely infected by *N. gonorrhoeae*. However, gonococcal vaginitis can occur in an estrogenic women (e.g., prepubertal girls and postmenopausal women), in whom the vaginal stratified squamous epithelium is often thinned down to the basilar layer, which can be infected by *N. gonorrhoeae*. The intense inflammation of the vagina makes the physical (speculum and bimanual) examination extremely painful. The vaginal mucosa is red and edematous, and an abundant purulent discharge is often present. Infection in the urethra and in Skene's and Bartholin's glands often accompanies gonococcal vaginitis. Inflamed cervical erosion or abscesses in nabothian cysts may also occur. Coexisting cervicitis may result in pus in the cervical os.

Anorectal Gonorrhea Because the female anatomy permits the spread of cervical exudate to the rectum, *N. gonorrhoeae* is sometimes recovered from the rectum of women with uncomplicated gonococcal cervicitis. The rectum is the sole site of infection in only 5% of women with gonorrhea. Such women are usually asymptomatic but occasionally have acute proctitis manifested by anorectal pain or pruritus, tenesmus, purulent rectal discharge, and rectal bleeding. Among men

who have sex with men (MSM), the frequency of gonococcal infection, including rectal infection, fell by $\geq 90\%$ throughout the United States in the early 1980s, but a resurgence of gonorrhoea among MSM has been documented in several cities since the 1990s. Gonococcal isolates from the rectum of MSM tend to be more resistant to antimicrobial agents than are gonococcal isolates from other sites. Gonococcal isolates with a mutation in *mtrR* (multiple transferable resistance repressor) or in the promoter region of the gene that encodes for this transcriptional repressor develop increased resistance to antimicrobial hydrophobic agents such as bile acids and fatty acids in feces and thus are found with increased frequency in MSM. This situation may have been responsible for higher rates of failure of treatment for rectal gonorrhoea with older regimens consisting of penicillin or tetracyclines.

Pharyngeal Gonorrhoea Pharyngeal gonorrhoea is usually mild or asymptomatic, although symptomatic pharyngitis does occasionally occur with cervical lymphadenitis. The mode of acquisition is oral-genital sexual exposure, with fellatio being a more efficient means of transmission than cunnilingus. In certain female adolescent populations in the United States, pharyngeal gonorrhoea has become as common as genital gonorrhoea. Most cases resolve spontaneously, and transmission from the pharynx to sexual contacts is rare. Pharyngeal infection almost always coexists with genital infection. Swabs from the pharynx should be plated directly onto gonococcal selective media. Pharyngeal colonization with *Neisseria meningitidis* needs to be differentiated from that with other *Neisseria* species.

Ocular Gonorrhoea in Adults Ocular gonorrhoea in an adult usually results from autoinoculation of *N. gonorrhoeae* from an infected genital site. As in genital infection, the manifestations range from severe to occasionally mild or asymptomatic disease. The variability in clinical manifestations may be attributable to differences in the ability of the infecting strain to elicit an inflammatory response. Infection may result in a markedly swollen eyelid, severe hyperemia and chemosis, and a profuse purulent discharge. The massively inflamed conjunctiva may be draped over the cornea and limbus. Lytic enzymes from the infiltrating PMNs occasionally cause corneal ulceration and rarely cause perforation.

Prompt recognition and treatment of this condition are of paramount importance. Gram's stain and culture of the purulent discharge establish the diagnosis. Genital cultures should also be performed.

Gonorrhoea in Pregnant Women, Neonates, and Children Gonorrhoea in pregnancy can have serious consequences for both the mother and the infant. Recognition of gonorrhoea early in pregnancy also identifies a population at risk for other STIs, particularly chlamydial infection, syphilis, and trichomoniasis. The risks of salpingitis and PID—conditions associated with a high rate of fetal loss—are highest during the first trimester. Pharyngeal infection, most often asymptomatic, may be more common during pregnancy because of altered sexual practices. Prolonged rupture of the membranes, premature delivery, chorioamnionitis, funisitis (infection of the umbilical cord stump), and sepsis in the infant (with *N. gonorrhoeae* detected in the newborn's gastric aspirate during delivery) are common complications of maternal gonococcal infection at term. Other conditions and microorganisms, including *Mycoplasma hominis*, *Ureaplasma urealyticum*, *C. trachomatis*, and bacterial vaginosis (often accompanied by infection with *Trichomonas vaginalis*), have been associated with similar complications.

The most common form of gonorrhoea in neonates is ophthalmia neonatorum, which results from exposure to infected cervical secretions during parturition. Ocular neonatal instillation of a prophylactic agent (e.g., 1% silver nitrate eye drops or ophthalmic preparations containing erythromycin or tetracycline) prevents ophthalmia neonatorum but is not effective for its treatment, which requires systemic antibiotics. The clinical manifestations are acute and usually begin 2–5 days after birth. An initial nonspecific conjunctivitis with a serosanguineous discharge is followed by tense edema of the eyelids, chemosis, and a profuse, thick, purulent discharge. Corneal ulcerations that result in nebulae or perforation may lead to anterior synechiae, anterior staphyloma, panophthalmitis, and blindness. Infections described at other mucosal sites in infants, including vaginitis, rhinitis, and anorectal

infection, are likely to be asymptomatic. Pharyngeal colonization has been demonstrated in 35% of infants with gonococcal ophthalmia, and coughing is the most prominent symptom in these cases. Septic arthritis (see below) is the most common manifestation of systemic infection or DGI in the newborn. The onset usually comes at 3–21 days of age, and polyarticular involvement is common. Sepsis, meningitis, and pneumonia are seen in rare instances.

Any STI in children beyond the neonatal period raises the possibility of sexual abuse. Gonococcal vulvovaginitis is the most common manifestation of gonococcal infection in children beyond infancy. Anorectal and pharyngeal infections are common in these children and are frequently asymptomatic. The urethra, Bartholin's and Skene's glands, and the upper genital tract are rarely involved. All children with gonococcal infection should also be evaluated for chlamydial infection, syphilis, and possibly HIV infection.

Gonococcal Arthritis (DGI) DGI (gonococcal arthritis) results from gonococcal bacteremia. In the 1970s, DGI occurred in ~0.5–3% of persons with untreated gonococcal mucosal infection. The lower incidence of DGI at present is probably attributable to a decline in the prevalence of particular strains that are likely to disseminate. DGI strains resist the bactericidal action of human serum and generally do not incite inflammation at genital sites, probably because of limited generation of chemotactic factors. Strains recovered from DGI cases in the 1970s were often of the PorB.1A serotype, were highly susceptible to penicillin, and had special growth requirements—including arginine, hypoxanthine, and uracil—that made the organism more fastidious and more difficult to isolate.

Menstruation is a risk factor for dissemination, and approximately two-thirds of cases of DGI are in women. In about half of affected women, symptoms of DGI begin within 7 days of onset of menses. Complement deficiencies, especially of the components involved in the assembly of the membrane attack complex (C5 through C9), predispose to neisserial bacteremia, and persons with more than one episode of DGI should be screened with an assay for total hemolytic complement activity.

The clinical manifestations of DGI have sometimes been classified into two stages: a bacteremic stage, which is less common today, and a joint-localized stage with suppurative arthritis. A clear-cut progression usually is not evident. Patients in the bacteremic stage have higher temperatures, and chills more frequently accompany their fever. Painful joints are common and often occur together with tenosynovitis and skin lesions. Polyarthralgias usually include the knees, elbows, and more distal joints; the axial skeleton is generally spared. Skin lesions are seen in ~75% of patients and include papules and pustules, often with a hemorrhagic component (Fig. 181-2; see also Fig. 25e-44). Other manifestations of noninfectious dermatitis, such as nodular lesions, urticaria, and erythema multiforme, have been described. These lesions are usually on the extremities and number between 5 and 40. The differential diagnosis of the bacteremic stage of DGI includes reactive arthritis, acute rheumatoid arthritis, sarcoidosis, erythema nodosum, drug-induced arthritis, and viral infections (e.g., hepatitis B and acute HIV infection). The distribution of joint symptoms in reactive arthritis differs from that in DGI (Fig. 181-3), as do the skin and genital manifestations (Chap. 384).

Suppurative arthritis involves one or two joints, most often the knees, wrists, ankles, and elbows (in decreasing order of frequency); other joints occasionally are involved. Most patients who develop gonococcal septic arthritis do so without prior polyarthralgias or skin lesions; in the absence of symptomatic genital infection, this disease cannot be distinguished from septic arthritis caused by other pathogens. The differential diagnosis of acute arthritis in young adults is discussed in Chap. 157. Rarely, osteomyelitis complicates septic arthritis involving small joints of the hand.

Gonococcal endocarditis, although rare today, was a relatively common complication of DGI in the preantibiotic era, accounting for about one-quarter of reported cases of endocarditis. Another unusual complication of DGI is meningitis.

Gonococcal Infections in HIV-Infected Persons The association between gonorrhoea and the acquisition of HIV has been demonstrated in

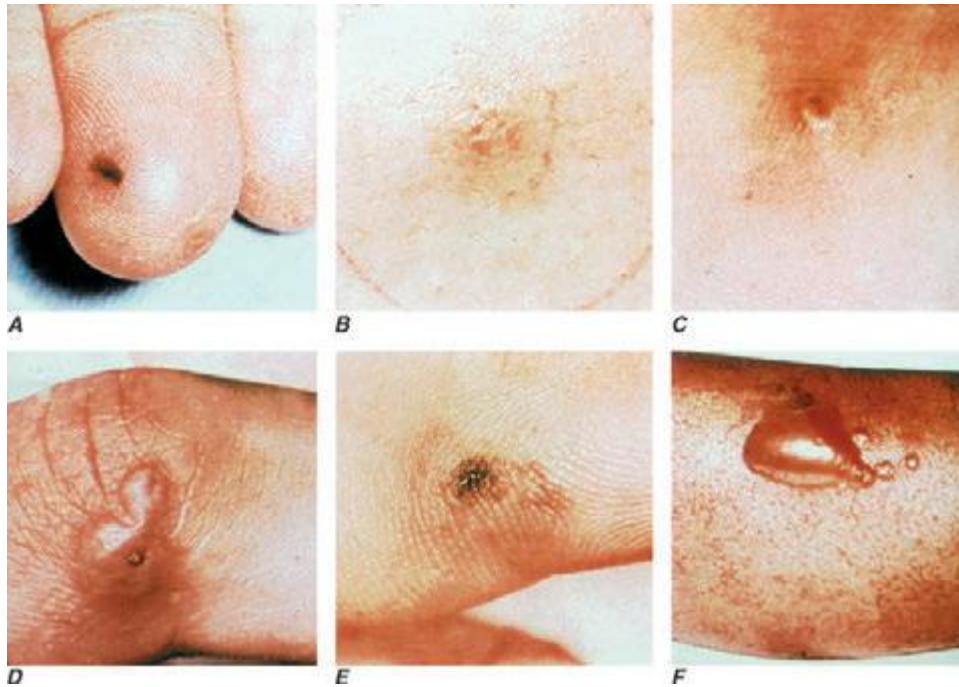


FIGURE 181-2 Characteristic skin lesions in patients with proven gonococcal bacteremia. The lesions are in various stages of evolution. **A.** Very early petechia on finger. **B.** Early papular lesion, 7 mm in diameter, on lower leg. **C.** Pustule with central eschar resulting from early petechial lesion. **D.** Pustular lesion on finger. **E.** Mature lesion with central necrosis (black) on hemorrhagic base. **F.** Bullae on anterior tibial surface. (Reprinted with permission from KK Holmes et al: *Disseminated gonococcal infection*. *Ann Intern Med* 74:979, 1971.)

several well-controlled studies, mainly in Kenya and Zaire. The non-ulcerative STIs enhance the transmission of HIV by three- to fivefold; transmission of HIV-infected immune cells and increased viral shedding by persons with urethritis or cervicitis may contribute (Chap. 226). HIV has been detected by polymerase chain reaction (PCR) more commonly in ejaculates from HIV-positive men with gonococcal urethritis than in those from HIV-positive men with nongonococcal urethritis. PCR positivity diminishes by twofold after appropriate therapy for urethritis. Not only does gonorrhea enhance the transmission of HIV, but it may also increase the individual's risk for acquisition of HIV. A proposed mechanism is the significantly greater number of CD4+ T lymphocytes and dendritic cells that can be infected by HIV in endocervical secretions

from women with nonulcerative STIs than in those from women with ulcerative STIs.

LABORATORY DIAGNOSIS

A rapid diagnosis of gonococcal infection in men may be obtained by Gram's staining of urethral exudates (Fig. 181-1). The detection of gram-negative intracellular monococci and diplococci is usually highly specific and sensitive in diagnosing gonococcal urethritis in symptomatic males but is only ~50% sensitive in diagnosing gonococcal cervicitis. Samples should be collected with Dacron or rayon swabs. Part of the sample should be inoculated onto a plate of modified Thayer-Martin or other gonococcal selective medium for culture. It is important to process all samples immediately because gonococci do not tolerate drying. If plates cannot be incubated immediately, they can be held safely for several hours at room temperature in candle extinction jars prior to incubation. If processing is to occur within 6 h, transport of specimens may be facilitated by the use of nonnutritive swab transport systems such as Stuart or Amies medium. For longer holding periods (e.g., when specimens for culture are to be mailed), culture media with self-contained CO₂-generating systems (such as the JEMBEC or Gono-Pak systems) may be used. Specimens should also be obtained for the diagnosis of chlamydial infection (Chap. 213).

PMNs are often seen in the endocervix on a Gram's stain, and an abnormally increased number (≥ 30 PMNs per field in five 1000 \times oil-immersion microscopic fields) establishes the presence of an inflammatory discharge. Unfortunately, the presence or absence of gram-negative intracellular monococci or diplococci in cervical smears does not accurately predict which patients have gonorrhea, and the diagnosis in this setting should be made by culture or another suitable nonculture diagnostic method. The sensitivity of a single endocervical culture is ~80–90%. If a history of rectal sex is elicited, a rectal wall swab (uncontaminated with feces) should be cultured. A presumptive diagnosis of gonorrhea cannot be made on the basis of gram-negative diplococci in smears from the pharynx, where other *Neisseria* species are components of the normal flora.

Increasingly, nucleic acid probe tests are being substituted for culture for the direct detection of *N. gonorrhoeae* in urogenital

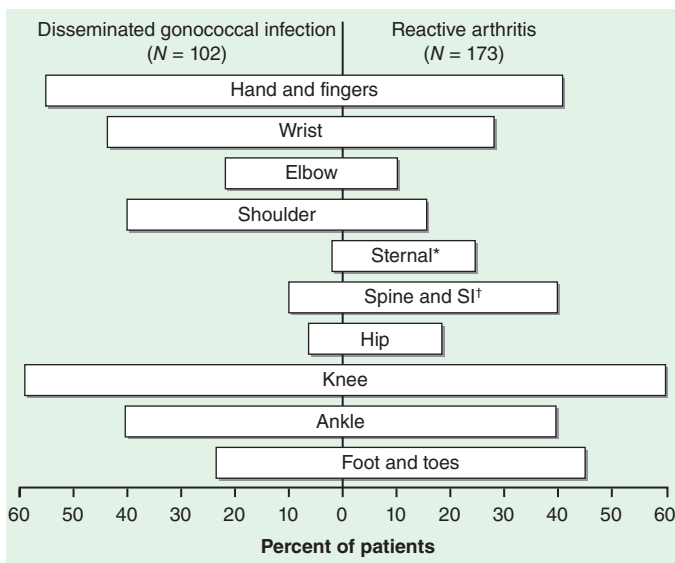


FIGURE 181-3 Distribution of joints with arthritis in 102 patients with disseminated gonococcal infection and 173 patients with reactive arthritis. *Includes the sternoclavicular joints. †SI, sacroiliac joint.

1008 specimens. A common assay uses a nonisotopic chemiluminescent DNA probe that hybridizes specifically with gonococcal 16S ribosomal RNA; this assay is as sensitive as conventional culture techniques. A disadvantage of non-culture-based assays is that *N. gonorrhoeae* cannot be grown from the transport systems. Thus a culture-confirmatory test and formal antimicrobial susceptibility testing, if needed, cannot be performed. Nucleic acid amplification tests (NAATs), including the Roche Cobas® Amplicor, Gen-Probe APTIMA COMBO 2®, and BD ProbeTec™ ET, also detect *C. trachomatis* and are more sensitive than culture identification of either *N. gonorrhoeae* or *C. trachomatis*. The Gen-Probe and BD tests offer the advantage that urine samples can be tested with a sensitivity similar to or greater than that obtained when urethral or cervical swab samples are assessed by other non-NAATs or culture, respectively. Several amplification tests are now available on semi-automated or fully automated platforms.

Because of the legal implications, the preferred method for the diagnosis of gonococcal infection in children is a standardized culture. Two positive NAATs, each targeting a different nucleic acid sequence, may be substituted for culture of the cervix or the urethra as legal evidence of infection in children. Although nonculture tests for gonococcal infection have not been approved by the U.S. Food and Drug Administration for use with specimens obtained from the pharynx and rectum of infected children, NAATs from these sites are preferred for diagnostic evaluation in adult victims of suspected sexual abuse, especially if the NAATs have been evaluated by the local laboratory and found to be superior. Cultures should be obtained from the pharynx and anus of both girls and boys, the urethra of boys, and the vagina of girls; cervical specimens are not recommended for prepubertal girls. For boys with a urethral discharge, a meatal specimen of the discharge is adequate for culture. Presumptive colonies of *N. gonorrhoeae* should be identified definitively by at least two independent methods.

Blood should be cultured in suspected cases of DGI. The use of Isolator blood culture tubes may enhance the yield. The probability of positive blood cultures decreases after 48 h of illness. Synovial fluid should be inoculated into blood culture broth medium and plated onto chocolate agar rather than selective medium because this fluid is not likely to be contaminated with commensal bacteria. Gonococci are infrequently recovered from early joint effusions containing <20,000 leukocytes/μL but may be recovered from effusions containing >80,000 leukocytes/μL. The organisms are seldom recovered from blood and synovial fluid of the same patient.

TREATMENT GONOCOCCAL INFECTIONS

Treatment failure can lead to continued transmission and the emergence of antibiotic resistance. The importance of adequate treatment with a regimen that the patient will adhere to cannot be overemphasized. Thus highly effective single-dose regimens have been developed for uncomplicated gonococcal infections. The modified 2010 treatment guidelines for gonococcal infections from the Centers for Disease Control and Prevention (CDC) are summarized in [Table 181-1](#). Rising MICs of cefixime worldwide have led the CDC to discontinue its recommendation of this agent as first-line treatment for uncomplicated gonorrhea. The recommendations for uncomplicated gonorrhea apply to HIV-infected as well as HIV-uninfected patients.

Currently, a single IM dose of the third-generation cephalosporin ceftriaxone is the mainstay of therapy for uncomplicated gonococcal infection of the urethra, cervix, rectum, or pharynx and almost always results in an effective cure. Quinolone-containing regimens are no longer recommended in the United States as first-line treatment because of widespread resistance. A recent multicenter trial of treatment for uncomplicated gonorrhea in the United States showed ≥99.5% efficacy of two combination regimens: (1) gemifloxacin (320 mg, single oral dose) plus azithromycin (2 g, single

oral dose) or (2) azithromycin (2 g, single oral dose) plus gentamicin (a single IM dose of 240 mg or, in individuals who weigh ≤45 kg, 5 mg/kg).

Because co-infection with *C. trachomatis* occurs frequently, initial treatment regimens must also incorporate an agent (e.g., azithromycin or doxycycline) that is effective against chlamydial infection. Pregnant women with gonorrhea, who should not take doxycycline, should receive concurrent treatment with a macrolide antibiotic for possible chlamydial infection. A single 1-g dose of azithromycin, which is effective therapy for uncomplicated chlamydial infections, results in an unacceptably low cure rate (93%) for gonococcal infections and should not be used alone. A single 2-g dose of azithromycin, particularly in the extended-release microsphere formulation, delivers azithromycin to the lower gastrointestinal tract, thereby improving tolerability. Azithromycin is effective against sensitive strains, but this drug is expensive, causes gastrointestinal distress, and is not recommended for routine or first-line treatment of gonorrhea. Spectinomycin has been used as an alternative agent for the treatment of uncomplicated gonococcal infections in penicillin-allergic persons outside the United States but is not currently available in this country. Of note, the limited effectiveness of spectinomycin for the treatment of pharyngeal infection reduces its utility in populations among whom such infection is common, such as MSM.

Persons with uncomplicated infections who receive ceftriaxone do not need a test of cure; however, cultures for *N. gonorrhoeae* should be performed if symptoms persist after therapy with an established regimen, and any gonococci isolated should be tested for antimicrobial susceptibility. Persons given an alternative regimen should return for a test of cure targeting the infected anatomic site. This test ideally should be a culture. If culture is not readily available and a NAAT is positive, every effort should be made to perform a confirmatory culture. All positive cultures for test of cure should undergo antimicrobial susceptibility testing. Because of high rates of reinfection with *N. gonorrhoeae* and *C. trachomatis* within 6 months, repeat testing is recommended 3 months after treatment.

Symptomatic gonococcal pharyngitis is more difficult to eradicate than genital infection. Persons who cannot tolerate ceftriaxone and those in whom quinolones are contraindicated may be treated with spectinomycin if it is available, but this agent results in a cure rate of ≤52%. Persons given spectinomycin should have a pharyngeal sample cultured 3–5 days after treatment as a test of cure. A single 2-g dose of azithromycin may be used in areas where rates of resistance to azithromycin are low.

Treatments for gonococcal epididymitis and PID are discussed in [Chap. 163](#). Ocular gonococcal infections in older children and adults should be managed with a single dose of ceftriaxone combined with saline irrigation of the conjunctivae (both undertaken expeditiously), and patients should undergo a careful ophthalmologic evaluation that includes a slit-lamp examination.

DGI may require higher dosages and longer durations of therapy (Table 181-1). Hospitalization is indicated if the diagnosis is uncertain, if the patient has localized joint disease that requires aspiration, or if the patient cannot be relied on to comply with treatment. Open drainage is necessary only occasionally—e.g., for management of hip infections that may be difficult to drain percutaneously. Nonsteroidal anti-inflammatory agents may be indicated to alleviate pain and hasten clinical improvement of affected joints.

Gonococcal meningitis and endocarditis should be treated in the hospital with high-dose IV ceftriaxone (1–2 g every 12 h); therapy should continue for 10–14 days for meningitis and for at least 4 weeks for endocarditis. All persons who experience more than one episode of DGI should be evaluated for complement deficiency.

TABLE 181-1 RECOMMENDED TREATMENT FOR GONOCOCCAL INFECTIONS: ADAPTED FROM THE 2010 GUIDELINES OF THE CENTERS FOR DISEASE CONTROL AND PREVENTION

Diagnosis	Treatment of Choice ^a
Uncomplicated gonococcal infection of the cervix, urethra, pharynx ^b , or rectum	
First-line regimen	Ceftriaxone (250 mg IM, single dose) <i>plus</i> Treatment for <i>Chlamydia</i> if chlamydial infection is not ruled out: Azithromycin (1 g PO, single dose) <i>or</i> Doxycycline (100 mg PO bid for 7 days)
Alternative regimens ^c	Cefixime (400 mg PO, single dose) <i>or</i> Ceftizoxime (500 mg IM, single dose) <i>or</i> Cefotaxime (500 mg IM, single dose) <i>or</i> Spectinomycin (2 g IM, single dose) ^{d,e} <i>or</i> Cefotetan (1 g IM, single dose) <i>plus</i> probenecid (1 g PO, single dose) ^d <i>or</i> Cefoxitin (2 g IM, single dose) <i>plus</i> probenecid (1 g PO, single dose) ^d
Epididymitis	See Chap. 163
Pelvic inflammatory disease	See Chap. 163
Gonococcal conjunctivitis in an adult	Ceftriaxone (1 g IM, single dose) ^f
Ophthalmia neonatorum ^g	Ceftriaxone (25–50 mg/kg IV, single dose, not to exceed 125 mg)
Disseminated gonococcal infection ^h	
Initial therapy ⁱ	
Patient tolerant of β-lactam drugs	Ceftriaxone (1 g IM or IV q24h; recommended) <i>or</i> Cefotaxime (1 g IV q8h) <i>or</i> Ceftizoxime (1 g IV q8h)
Patients allergic to β-lactam drugs	Spectinomycin (2 g IM q12h) ^d
Continuation therapy ^j	Cefixime (400 mg PO bid)
Meningitis or endocarditis	See text ^k

^aTrue failure of treatment with a recommended regimen is rare and should prompt an evaluation for reinfection, infection with a drug-resistant strain, or an alternative diagnosis. ^bCeftriaxone is the only agent recommended for treatment of pharyngeal infection. ^cSee text for follow-up of persons with infection who are treated with alternative regimens. ^dSpectinomycin, cefotetan, and cefoxitin, which are alternative agents, currently are unavailable or in short supply in the United States. ^eSpectinomycin may be ineffective for the treatment of pharyngeal gonorrhea. ^fPlus lavage of the infected eye with saline solution (once). ^gProphylactic regimens are discussed in the text. ^hHospitalization is indicated if the diagnosis is uncertain, if the patient has frank arthritis with an effusion, or if the patient cannot be relied on to adhere to treatment. ⁱAll initial regimens should be continued for 24–48 h after clinical improvement begins, at which time the switch may be made to an oral agent (e.g., cefixime or a quinolone) if antimicrobial susceptibility can be documented by culture of the causative organism. If no organism is isolated and the diagnosis is secure, then treatment with ceftriaxone should be continued for at least 1 week. Treatment for chlamydial infection (as above) should be given if this infection has not been ruled out. ^kHospitalization is indicated to exclude suspected meningitis or endocarditis.

PREVENTION AND CONTROL

Condoms, if properly used, provide effective protection against the transmission and acquisition of gonorrhea as well as other infections that are transmitted to and from genital mucosal surfaces. Spermicidal preparations used with a diaphragm or cervical sponges impregnated with nonoxynol 9 offer some protection against gonorrhea and chlamydial infection. However, the frequent use of preparations that contain nonoxynol 9 is associated with mucosal disruption that paradoxically may enhance the risk of HIV infection in the event of exposure. All patients should be instructed to refer sex partners for evaluation and treatment. All sex partners of persons with gonorrhea should be evaluated and treated for *N. gonorrhoeae* and *C. trachomatis* infections if their last contact with the patient took place within 60 days before the onset of symptoms or the diagnosis of infection in the patient. If the patient's last sexual encounter was >60 days before onset of symptoms or diagnosis, the patient's most recent sex partner should be treated. Partner-delivered medications or prescriptions for

medications to treat gonorrhea and chlamydial infection diminish the likelihood of reinfection (or relapse) in the infected patient. In states where it is legal, this approach is an option for partner management. Patients should be instructed to abstain from sexual intercourse until therapy is completed and until they and their sex partners no longer have symptoms. Greater emphasis must be placed on prevention by public health education, individual patient counseling, and behavior modification. Sexually active persons, especially adolescents, should be offered screening for STIs. For male patients, a NAAT on urine or a urethral swab may be used for screening. Preventing the spread of gonorrhea may help reduce the transmission of HIV. No effective vaccine for gonorrhea is yet available, but efforts to test several candidates are under way.

ACKNOWLEDGMENT

The authors acknowledge the contributions of Dr. King K. Holmes and Dr. Stephen A. Morse to the chapter on this subject in earlier editions.

182 Haemophilus and Moraxella Infections

Timothy F. Murphy

HAEMOPHILUS INFLUENZAE

MICROBIOLOGY

Haemophilus influenzae was first recognized in 1892 by Pfeiffer, who erroneously concluded that the bacterium was the cause of influenza. *H. influenzae* is a small ($1 \times 0.3\text{-}\mu\text{m}$) gram-negative organism of variable shape; thus, it is often described as a pleomorphic coccobacillus. In clinical specimens such as cerebrospinal fluid (CSF) and sputum, *H. influenzae* frequently stains only faintly with safranin and therefore can easily be overlooked.

H. influenzae grows both aerobically and anaerobically. Its aerobic growth requires two factors: hemin (X factor) and nicotinamide adenine dinucleotide (V factor). These requirements are used in the clinical laboratory to identify the bacterium. Caution must be used to distinguish *H. influenzae* from *H. haemolyticus*, a respiratory tract commensal that has identical growth requirements. *H. haemolyticus* has classically been distinguished from *H. influenzae* by the hemolysis of the former species on horse blood agar. However, a significant proportion of isolates of *H. haemolyticus* have now been recognized as nonhemolytic. Analysis of various genotypic and phenotypic markers, including 16S ribosomal sequences, superoxide dismutase, outer-membrane protein P6, protein D, and fuculose kinase, can be used to distinguish these two species.

Six major serotypes of *H. influenzae* have been identified; designated *a* through *f*, they are based on antigenically distinct polysaccharide capsules. In addition, some strains lack a polysaccharide capsule and are referred to as nontypable strains. Type b and nontypable strains are the most relevant strains clinically (Table 182-1), although encapsulated strains other than type b can cause disease. *H. influenzae* was the first free-living organism to have its entire genome sequenced.

The antigenically distinct type b capsule is a linear polymer composed of ribosyl-ribitol phosphate. Strains of *H. influenzae* type b (Hib) cause disease primarily in infants and children <6 years of age. Nontypable strains are primarily mucosal pathogens but occasionally cause invasive disease.

EPIDEMIOLOGY AND TRANSMISSION

H. influenzae, an exclusively human pathogen, is spread by airborne droplets or by direct contact with secretions or fomites. Colonization with nontypable *H. influenzae* is a dynamic process; new strains are acquired and other strains are replaced periodically.

The widespread use of Hib conjugate vaccines in many industrialized countries has resulted in striking decreases in the rate of nasopharyngeal colonization by Hib and in the incidence of Hib infection (Fig. 182-1). However, the majority of the world's children remain unimmunized. Worldwide, invasive Hib disease occurs predominantly in unimmunized children and in those who have not



completed the primary immunization series. Certain groups have a higher incidence of invasive Hib disease than the general population, including African-American children and Native American groups. Although this increased incidence has not yet been accounted for, several factors may be relevant, including age at exposure to the bacterium, socioeconomic conditions, and genetic differences.

TABLE 182-1 CHARACTERISTICS OF TYPE b AND NONTYPABLE STRAINS OF HAEMOPHILUS INFLUENZAE

Feature	Type b Strains	Nontypable Strains
Capsule	Ribosyl-ribitol phosphate	Unencapsulated
Pathogenesis	Invasive infections due to hematogenous spread	Mucosal infections due to contiguous spread
Clinical manifestations	Meningitis and invasive infections in incompletely immunized infants and children	Otitis media in infants and children; lower respiratory tract infections in adults with chronic bronchitis
Evolutionary history	Basically clonal	Genetically diverse
Vaccine	Highly effective conjugate vaccines	None available; under development

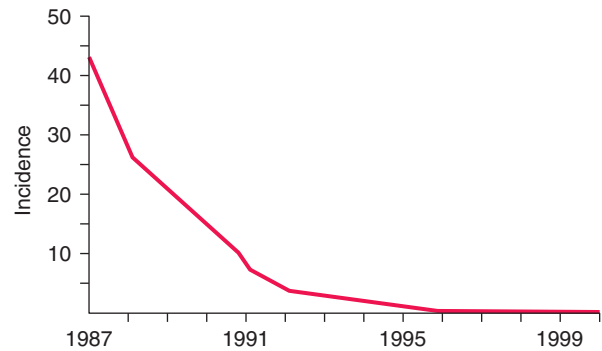


FIGURE 182-1 Estimated incidence (rate per 100,000) of invasive disease due to *Haemophilus influenzae* type b among children <5 years of age: 1987–2000. Fewer than 40 cases per year have been reported since 2000. (Data from the Centers for Disease Control and Prevention.)

completed the primary immunization series. Certain groups have a higher incidence of invasive Hib disease than the general population, including African-American children and Native American groups. Although this increased incidence has not yet been accounted for, several factors may be relevant, including age at exposure to the bacterium, socioeconomic conditions, and genetic differences.

PATHOGENESIS

Hib strains cause systemic disease by invasion and hematogenous spread from the respiratory tract to distant sites such as the meninges, bones, and joints. The type b polysaccharide capsule is an important virulence factor affecting the bacterium's ability to avoid opsonization and cause systemic disease.

Nontypable strains cause disease by local invasion of mucosal surfaces. Otitis media results when bacteria reach the middle ear by way of the eustachian tube. Adults with chronic bronchitis experience recurrent lower respiratory tract infection due to nontypable strains. In addition, persistent nontypable *H. influenzae* colonization of the lower airways of adults with chronic obstructive pulmonary disease (COPD) contributes to the airway inflammation that is a hallmark of the disease. Nontypable strains that cause infection in adults with COPD differ in pathogenic potential and genome content from strains that cause otitis media. In the middle ear, nontypable strains form biofilms. More resistant to host clearance mechanisms and to antibiotics than are planktonic bacteria, biofilms are associated with chronic and recurrent otitis media. The incidence of invasive disease caused by nontypable strains is low. Strains that cause invasive disease are genetically and phenotypically diverse.

IMMUNE RESPONSE

Antibody to the capsule is important in protection from infection by Hib strains. The level of (maternally acquired) serum antibody to the capsular polysaccharide, which is a polymer of polyribitol ribose phosphate (PRP), declines from birth to 6 months of age and, in the absence of vaccination, remains low until ~2 or 3 years of age. The age at the antibody nadir correlates with that of the peak incidence of type b disease. Antibody to PRP then appears partly as a result of exposure to Hib or cross-reacting antigens. Systemic Hib disease is unusual after the age of 6 years because of the presence of protective antibody. Vaccines in which PRP is conjugated to protein carrier molecules have been developed and are now used widely. These vaccines generate an antibody response to PRP in infants and effectively prevent invasive infections in infants and children.

Since nontypable strains lack a capsule, the immune response to infection is directed at noncapsular antigens. These antigens have generated considerable interest as immune targets and potential vaccine components. The human immune response to nontypable strains appears to be strain-specific, a characteristic that accounts in part for the propensity of these strains to cause recurrent otitis media and recurrent exacerbations of chronic bronchitis in immunocompetent hosts.

CLINICAL MANIFESTATIONS

Hib The most serious manifestation of infection with Hib is *meningitis* (Chap. 164), which primarily affects children <2 years of age. The clinical manifestations of Hib meningitis are similar to those of meningitis caused by other bacterial pathogens. Fever and altered central nervous system function are the most common features at presentation. Nuchal rigidity may or may not be evident. Subdural effusion, the most common complication, is suspected when, despite 2 or 3 days of appropriate antibiotic therapy, the infant has seizures, hemiparesis, or continued obtundation. The overall mortality rate from Hib meningitis is ~5%, and the morbidity rate is high. Of survivors, 6% have permanent sensorineural hearing loss, and about one-fourth have a significant handicap of some type. If more subtle handicaps are sought, up to half of survivors are found to have some neurologic sequelae, such as partial hearing loss and delayed language development.

Epiglottitis (Chap. 44) is a life-threatening Hib infection involving cellulitis of the epiglottis and supraglottic tissues. It can lead to acute upper airway obstruction. Its unique epidemiologic features are its occurrence in an older age group (2–7 years old) than other Hib infections and its absence among Navajo Indians and Alaskan Eskimos. Sore throat and fever rapidly progress to dysphagia, drooling, and airway obstruction. Epiglottitis also occurs in adults.

Cellulitis (Chap. 156) due to Hib occurs in young children. The most common location is on the head or neck, and the involved area sometimes takes on a characteristic bluish-red color. Most patients have bacteremia, and 10% have an additional focus of infection.

Hib causes *pneumonia* in infants. The infection is clinically indistinguishable from other types of bacterial pneumonia (e.g., pneumococcal pneumonia) except that Hib is more likely to involve the pleura. Several less common invasive conditions can be important clinical manifestations of Hib infection in children. These include osteomyelitis, septic arthritis, pericarditis, orbital cellulitis, endophthalmitis, urinary tract infection, abscesses, and bacteremia without an identifiable focus.

Non-type b encapsulated strains of *H. influenzae* (types a, c, d, e, and f) are unusual causes of invasive infection manifested predominantly by bacteremia and pneumonia. Most such infections occur in the setting of underlying conditions.

Nontypable *H. influenzae* Nontypable *H. influenzae* is the most common bacterial cause of exacerbations of COPD; these exacerbations are characterized by increased cough, sputum production, and shortness of breath. Fever is low-grade, and no infiltrates are evident on chest x-ray. Nontypable strains also cause community-acquired bacterial pneumonia in adults, especially among patients with COPD or AIDS. The clinical features of *H. influenzae* pneumonia are similar to those of other types of bacterial pneumonia, including pneumococcal pneumonia.

Nontypable *H. influenzae* is one of the three most common causes of childhood otitis media (the other two being *Streptococcus pneumoniae* and *Moraxella catarrhalis*) (Chap. 44). Infants are febrile and irritable, while older children report ear pain. Symptoms of viral upper respiratory infection often precede otitis media. The diagnosis is made by pneumatic otoscopy. An etiologic diagnosis, although not routinely sought, can be established by tympanocentesis and culture of middle-ear fluid. Clinical features associated with *H. influenzae* otitis media include a history of recurrent episodes, treatment failure, concomitant conjunctivitis, bilateral otitis media, and recent antimicrobial therapy. The increasing use of pneumococcal polysaccharide conjugate vaccines in infants is resulting in a relative increase in the proportion of otitis media cases that are caused by *H. influenzae*.

Nontypable *H. influenzae* also causes puerperal sepsis and is an important cause of neonatal bacteremia. These nontypable strains, which are closely related to *H. haemolyticus*, tend to be of biotype IV and cause invasive disease after colonizing the female genital tract.

Nontypable *H. influenzae* causes sinusitis (Chap. 44) in adults and children. In addition, the bacterium is a less common cause of various invasive infections. These infections include empyema, adult epiglottitis, pericarditis, cellulitis, septic arthritis, osteomyelitis, endocarditis, cholecystitis, intraabdominal infections, urinary tract infections,

mastoiditis, aortic graft infection, and bacteremia without a detectable focus. While most *H. influenzae* invasive infections in countries where Hib vaccines are used widely are caused by nontypable strains, there is no convincing evidence of an increased incidence of infection by nontypable *H. influenzae* as a result of use of Hib vaccines. Continued monitoring will be important. Many patients with *H. influenzae* bacteremia have an underlying condition, such as HIV infection, cardiopulmonary disease, alcoholism, or cancer.

DIAGNOSIS

The most reliable method for establishing a diagnosis of Hib infection is recovery of the organism in culture. The presence of gram-negative coccobacilli in Gram-stained CSF is strong evidence for Hib meningitis. Recovery of the organism from CSF confirms the diagnosis. Cultures of other normally sterile body fluids, such as blood, joint fluid, pleural fluid, pericardial fluid, and subdural effusion, are confirmatory in other infections.

Detection of PRP is an important adjunct to culture in rapid diagnosis of Hib meningitis. Immunoelectrophoresis, latex agglutination, coagglutination, and enzyme-linked immunosorbent assay are effective in detecting PRP. These assays are particularly helpful when patients have received prior antimicrobial therapy and thus are especially likely to have negative cultures.

Because nontypable *H. influenzae* is primarily a mucosal pathogen, it is a component of a mixed flora; thus etiologic diagnosis is challenging. Nontypable *H. influenzae* infection is strongly suggested by the predominance of gram-negative coccobacilli among abundant polymorphonuclear leukocytes in a Gram-stained sputum specimen from a patient in whom pneumonia is suspected. Although bacteremia is detectable in a small proportion of patients with pneumonia due to nontypable *H. influenzae*, most such patients have negative blood cultures.

A diagnosis of otitis media is based on the detection by pneumatic otoscopy of fluid in the middle ear. An etiologic diagnosis requires tympanocentesis but is not routinely sought. An invasive procedure is also required to determine the etiology of sinusitis; thus, treatment is often empirical once the diagnosis is suspected in light of clinical symptoms and sinus radiographs.

TREATMENT HAEMOPHILUS INFLUENZAE

Initial therapy for meningitis due to Hib should consist of a cephalosporin such as ceftriaxone or cefotaxime. For children, the dosage of ceftriaxone is 75–100 mg/kg daily given in two doses 12 h apart. The pediatric dosage of cefotaxime is 200 mg/kg daily given in four doses 6 h apart. Adult dosages are 2 g every 12 h for ceftriaxone and 2 g every 4–6 h for cefotaxime. An alternative regimen for initial therapy is ampicillin (200–300 mg/kg daily in four divided doses) plus chloramphenicol (75–100 mg/kg daily in four divided doses). Therapy should continue for a total of 1–2 weeks.

Administration of glucocorticoids to patients with Hib meningitis reduces the incidence of neurologic sequelae. The presumed mechanism is reduction of the inflammation induced by bacterial cell-wall mediators of inflammation when cells are killed by antimicrobial agents. Dexamethasone (0.6 mg/kg per day intravenously in four divided doses for 2 days) is recommended for the treatment of Hib meningitis in children >2 months of age.

Invasive infections other than meningitis are treated with the same antimicrobial agents. For epiglottitis, the dosage of ceftriaxone is 50 mg/kg daily, and the dosage of cefotaxime is 150 mg/kg daily, given in three divided doses 8 h apart. Epiglottitis constitutes a medical emergency, and maintenance of an airway is critical. The duration of therapy is determined by the clinical response. A course of 1–2 weeks is usually appropriate.

Many infections caused by nontypable strains of *H. influenzae*, such as otitis media, sinusitis, and exacerbations of COPD, can be treated with oral antimicrobial agents. Approximately 20–35% of nontypable strains produce β -lactamase (with the exact proportion

1012 depending on geographic location), and these strains are resistant to ampicillin. Several agents have excellent activity against nontypable *H. influenzae*, including amoxicillin/clavulanic acid, various extended-spectrum cephalosporins, and the macrolides azithromycin and clarithromycin. Fluoroquinolones are highly active against *H. influenzae* and are useful in adults with exacerbations of COPD. However, fluoroquinolones are not currently recommended for the treatment of children or pregnant women because of possible effects on articular cartilage.



In addition to β -lactamase production, alteration of penicillin-binding proteins—a second mechanism of ampicillin resistance—has been detected in isolates of *H. influenzae*. Although rare in the United States, these β -lactamase-negative ampicillin-resistant strains are common in Japan and are increasing in prevalence in Europe. Continued monitoring of the evolving antimicrobial susceptibility patterns of *H. influenzae* will be important.

PREVENTION



Vaccination (See also [Chap. 148](#)) Two conjugate vaccines that prevent invasive infections with Hib in infants and children are licensed in the United States. In addition to eliciting protective antibody, these vaccines prevent disease by reducing rates of pharyngeal colonization with Hib. The widespread use of conjugate vaccines has dramatically reduced the incidence of Hib disease in developed countries. Even though the manufacture of Hib vaccines is costly, vaccination is cost-effective. The Global Alliance for Vaccines and Immunizations has recognized the underuse of Hib conjugate vaccines.

The disease burden has been reduced in developing countries that have implemented routine vaccination (e.g., The Gambia, Chile). An important obstacle to more widespread vaccination is the lack of data on the epidemiology and burden of Hib disease in many developing countries.

All children should be immunized with an Hib conjugate vaccine, receiving the first dose at ~2 months of age, the rest of the primary series at 2–6 months of age, and a booster dose at 12–15 months of age. Specific recommendations vary for the different conjugate vaccines. The reader is referred to the recommendations of the American Academy of Pediatrics ([Chap. 148](#) and www.cispiimmunize.org).

Currently, no vaccines are available specifically for the prevention of disease caused by nontypable *H. influenzae*. However, a vaccine that contains protein D—a surface protein of *H. influenzae*—conjugated to pneumococcal polysaccharides is licensed in other countries and is used widely in Europe. The vaccine has shown partial efficacy in preventing *H. influenzae* otitis media in clinical trials. Additional progress in the development of vaccines against nontypable *H. influenzae* is anticipated.

Chemoprophylaxis The risk of secondary disease is greater than normal among household contacts of patients with Hib disease. Therefore, all children and adults (except pregnant women) in households with an index case and at least one incompletely immunized contact <4 years of age should receive prophylaxis with oral rifampin. When two or more cases of invasive Hib disease have occurred within 60 days at a child-care facility attended by incompletely vaccinated children, administration of rifampin to all attendees and personnel is indicated, as is recommended for household contacts. Chemoprophylaxis is not indicated in nursery and child-care contacts of a single index case. The reader is referred to the recommendations of the American Academy of Pediatrics.

HAEMOPHILUS DUCREYI

Haemophilus ducreyi is the etiologic agent of chancroid ([Chap. 163](#)), a sexually transmitted disease characterized by genital ulceration and inguinal adenitis. In addition to being a cause of morbidity in itself, chancroid is associated with HIV infection because of the role played by genital ulceration in HIV transmission. Chancroid increases the

efficiency of transmission of and the degree of susceptibility to HIV infection.

MICROBIOLOGY

H. ducreyi is a highly fastidious coccobacillary gram-negative bacterium whose growth requires X factor (hemin). Although, in light of this requirement, the bacterium has been classified in the genus *Haemophilus*, DNA homology and chemotaxonomic studies have established substantial differences between *H. ducreyi* and other *Haemophilus* species. Taxonomic reclassification of the organism is likely in the future but awaits further study. Ulcers contain predominantly T cells. The fact that patients who have had chancroid may have repeated infections indicates that infection does not confer protection.

EPIDEMIOLOGY AND PREVALENCE



The prevalence of chancroid has declined in the United States and worldwide. However, prevalence data must be interpreted with caution because of the difficulty of establishing a diagnosis. The infection appears to be more common in developing countries. Transmission is predominantly heterosexual, and cases in males have outnumbered those in females by ratios of 3:1 to 25:1 during outbreaks. Contact with commercial sex workers and illicit drug use are strongly associated with chancroid.

CLINICAL MANIFESTATIONS AND DIFFERENTIAL DIAGNOSIS

Infection is acquired as the result of a break in the epithelium during sexual contact with an infected individual. After an incubation period of 4–7 days, the initial lesion—a papule with surrounding erythema—appears. In 2 or 3 days, the papule evolves into a pustule, which spontaneously ruptures and forms a sharply circumscribed ulcer that generally is not indurated ([Fig. 182-2](#)). The ulcers are painful and bleed easily; little or no inflammation of the surrounding skin is evident. Approximately half of patients develop enlarged, tender inguinal lymph nodes, which frequently become fluctuant and spontaneously rupture. Patients usually seek medical care after 1–3 weeks of painful symptoms.

The presentation of chancroid does not usually include all of the typical clinical features and is sometimes atypical. Multiple ulcers can coalesce to form giant ulcers. Ulcers can appear and then resolve, with inguinal adenitis ([Fig. 182-2](#)) and suppuration following 1–3 weeks later; this clinical picture can be confused with that of lymphogranuloma venereum ([Chap. 213](#)). Multiple small ulcers can resemble



FIGURE 182-2 Chancroid with characteristic penile ulcers and associated left inguinal adenitis (bubo).

folliculitis. Other differential diagnostic considerations include the various infections causing genital ulceration, such as primary syphilis, secondary syphilis (condyloma latum), genital herpes, and donovanosis. In rare cases, chancroid lesions become secondarily infected with bacteria; the result is extensive inflammation.

DIAGNOSIS

Clinical diagnosis of chancroid is often inaccurate, and laboratory confirmation should be attempted in suspected cases. An accurate diagnosis of chancroid relies on culture of *H. ducreyi* from the lesion or from an aspirate of suppurative lymph nodes. Since the organism can be difficult to grow, the use of selective and supplemented media is necessary. No polymerase chain reaction assay for *H. ducreyi* is commercially available; such tests can be performed by Clinical Laboratory Improvement Amendment (CLIA)-certified clinical laboratories that have developed their own assays.

A probable diagnosis of chancroid can be made when the following criteria are met: (1) one or more painful genital ulcers; (2) no evidence of *Treponema pallidum* infection by dark-field examination of ulcer exudate or by a negative serologic test for syphilis performed at least 7 days after ulcer onset; (3) a typical clinical presentation for chancroid; and (4) a negative test for herpes simplex virus in the ulcer exudate.

TREATMENT HAEMOPHILUS DUCREYI

Treatment regimens recommended by the Centers for Disease Control and Prevention include (1) a single 1-g oral dose of azithromycin; (2) ceftriaxone (250 mg intramuscularly in a single dose); (3) ciprofloxacin (500 mg by mouth twice a day for 3 days); and (4) erythromycin base (500 mg by mouth three times a day for 7 days). Isolates from patients who do not respond promptly to treatment should be tested for antimicrobial resistance. In patients with HIV infection, healing may be slow and longer courses of treatment may be necessary. Clinical treatment failure in HIV-seropositive patients may reflect co-infection, especially with herpes simplex virus. Contacts of patients with chancroid should be identified and treated, whether or not symptoms are present, if they have had sexual contact with the patient during the 10 days preceding the patient's onset of symptoms.

MORAXELLA CATARRHALIS

MICROBIOLOGY

M. catarrhalis is an unencapsulated gram-negative diplococcus whose ecologic niche is the human respiratory tract. The organism was initially designated *Micrococcus catarrhalis*. Its name was changed to *Neisseria catarrhalis* in 1970 because of phenotypic similarities to commensal *Neisseria* species. On the basis of more rigorous analysis of genetic relatedness, *Moraxella catarrhalis* is now the widely accepted name for this species.

EPIDEMIOLOGY



Nasopharyngeal colonization by *M. catarrhalis* is common in infancy, with colonization rates ranging between 33% and 100% and depending on geographic location. Several factors probably account for this geographic variation, including living conditions, day-care attendance, hygiene, household smoking, and population genetics. The prevalence of colonization decreases steadily with age.

The widespread use of pneumococcal conjugate vaccines in some countries has resulted in alterations in patterns of nasopharyngeal colonization in resident populations. A relative increase in colonization by nonvaccine pneumococcal serotypes, nontypable *H. influenzae*, and *M. catarrhalis* has occurred. These changes in colonization patterns may be altering the distribution of pathogens of both otitis media and sinusitis in children.

PATHOGENESIS

M. catarrhalis causes mucosal infections of the respiratory tract by contiguous spread from its colonizing site in the upper airway. A

preceding viral upper respiratory tract infection is a common inciting event for otitis media. In exacerbations of COPD, the acquisition of new strains is critical for pathogenesis. Strains exhibit substantial genetic diversity and differences in virulence properties.

The expression of several adhesin molecules with differing specificities for various host cell receptors reflects the importance of adherence to the respiratory epithelial surface in the pathogenesis of infection. *M. catarrhalis* invades multiple cell types. Its intracellular residence in lymphoid tissue provides a potential reservoir for persistence in the human respiratory tract. Like many gram-negative bacteria, *M. catarrhalis* sheds vesicles into the surrounding environment. The vesicles are internalized by host cells and mediate several virulence mechanisms, including induction of inflammation and delivery of β -lactamase, that can promote the survival of co-pathogens.

CLINICAL MANIFESTATIONS

In children, *M. catarrhalis* causes predominantly mucosal infections when the bacterium migrates from the nasopharynx to the middle ear or the sinuses (Chap. 44). The inciting event for both otitis media and sinusitis is often a preceding viral infection. Overall, cultures of middle-ear fluid obtained by tympanocentesis indicate that *M. catarrhalis* causes 15–20% of cases of acute otitis media. Acute otitis media caused by *M. catarrhalis* or nontypable *H. influenzae* is clinically milder than otitis media caused by *S. pneumoniae*, with less fever and a lower prevalence of a red bulging tympanic membrane. However, substantial overlap makes it impossible to predict etiology in an individual child on the basis of clinical features.

A small proportion of viral upper respiratory tract infections are complicated by bacterial sinusitis. Cultures of sinus puncture aspirates show that *M. catarrhalis* accounts for ~20% of cases of acute bacterial sinusitis in children and for a smaller proportion in adults.

M. catarrhalis is a common cause of exacerbations in adults with COPD. The bacterium has been overlooked in this clinical setting because it has long been considered to be a commensal and because it is easily mistaken for commensal *Neisseria* species in cultures of respiratory secretions (see “Diagnosis,” below). Several independent lines of evidence have established *M. catarrhalis* as a pathogen in COPD. These include (1) the demonstration of *M. catarrhalis* in the lower airways during exacerbations, (2) the association of exacerbation with acquisition of new strains, (3) elevations of inflammatory markers in association with *M. catarrhalis*, and (4) the development of specific immune responses following infection. *M. catarrhalis* is the second most common bacterial cause of COPD exacerbations (after *H. influenzae*), as shown in a 10-year prospective study; the distribution of exacerbations associated with new-strain acquisitions is shown in Fig. 182-3. Not included are culture-negative cases or cases from which a pathogen had been previously isolated. With the application of rigorous clinical criteria for defining the etiology of exacerbations (both culture-positive and culture-negative), ~10% of all exacerbations in the same study were caused by *M. catarrhalis*. The clinical features of an exacerbation due to *M. catarrhalis* are similar to those of exacerbations due to other bacterial pathogens, including *H. influenzae* and *S. pneumoniae*. The cardinal symptoms are cough with increased sputum production, sputum purulence, and dyspnea in comparison with baseline symptoms.

Pneumonia due to *M. catarrhalis* occurs in the elderly, particularly in the setting of underlying cardiopulmonary disease, but is infrequent. Invasive infections, such as bacteremia, endocarditis, neonatal meningitis, and septic arthritis, are rare.

DIAGNOSIS

Tympanocentesis is required for etiologic diagnosis of otitis media, but this procedure is not performed routinely. Therefore, treatment of otitis media is generally empirical. Similarly, an etiologic diagnosis of sinusitis requires an invasive procedure and thus is usually not available to the clinician. Isolation of *M. catarrhalis* from an expectorated sputum sample from an adult experiencing clinical symptoms of an exacerbation is suggestive, but not diagnostic, of *M. catarrhalis* as the cause.

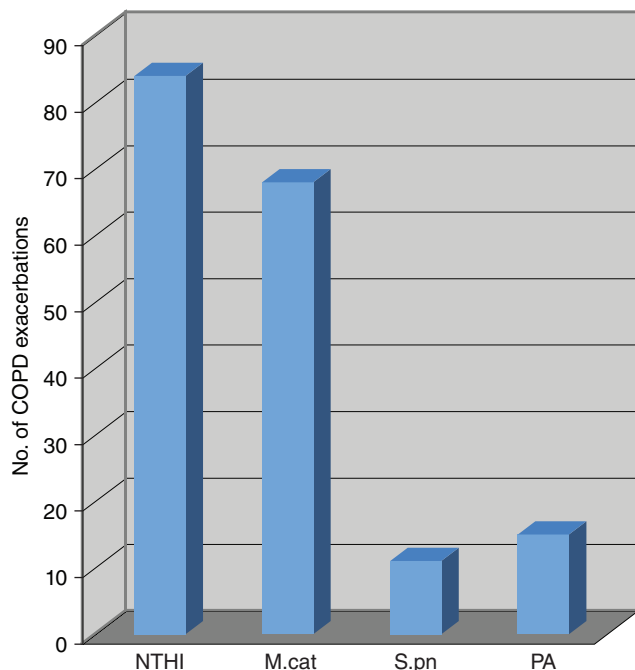


FIGURE 182-3 Cumulative results of a prospective study (1994–2004) of bacterial infection in chronic obstructive pulmonary disease (COPD) showing etiology of exacerbations. The numbers of exacerbations shown indicate the acquisition of a new strain simultaneous with clinical symptoms of an exacerbation. NTHI, nontypable *H. influenzae*; M.cat, *M. catarrhalis*; S.pn, *Streptococcus pneumoniae*; PA, *Pseudomonas aeruginosa*. (Adapted from TF Murphy, GI Parameswaran: *Clin Infect Dis* 49:124, 2009, with permission. © 2009 Infectious Diseases Society of America.)

Upon culture, colonies of *M. catarrhalis* resemble commensal neisseriae that are part of the normal upper airway flora. As mentioned above, the difficulty in distinguishing colonies of *M. catarrhalis* from neisserial colonies in cultures of respiratory secretions explains in part why *M. catarrhalis* has been overlooked as a pathogen. In contrast to these *Neisseria* species, *M. catarrhalis* colonies can be slid across the agar surface without disruption (the “hockey puck sign”). In addition, after 48 h of growth, *M. catarrhalis* colonies take on a pink color and tend to be larger than neisserial colonies. A variety of biochemical tests can distinguish *M. catarrhalis* from neisseriae. Kits that rely on these biochemical reactions are commercially available.

TREATMENT **MORAXELLA CATARRHALIS**

M. catarrhalis rapidly acquired Δ -lactamases during the 1970s and 1980s; antimicrobial susceptibility patterns have remained relatively stable since that time, with >90% of strains now producing Δ -lactamase and thus resistant to amoxicillin. Otitis media in children and exacerbations of COPD in adults are generally managed empirically with antimicrobial agents that are active against *S. pneumoniae*, *H. influenzae*, and *M. catarrhalis*. Most strains of *M. catarrhalis* are susceptible to amoxicillin/clavulanic acid, extended-spectrum cephalosporins, newer macrolides (azithromycin, clarithromycin), trimethoprim-sulfamethoxazole, and fluoroquinolones.

THE HACEK GROUP

HACEK organisms are a group of fastidious, slow-growing, gram-negative bacteria whose growth requires an atmosphere of carbon dioxide. Species belonging to this group include several *Haemophilus* species, *Aggregatibacter* (formerly *Actinobacillus*) species, *Cardiobacterium hominis*, *Eikenella corrodens*, and *Kingella kingae*. HACEK bacteria normally reside in the oral cavity and have been associated with local infections in the mouth. They are also known to cause severe systemic infections—most often bacterial endocarditis, which can develop on either native or prosthetic valves (Chap. 155).

In large series, 0.8–6% of cases of infective endocarditis are attributable to HACEK organisms, most often *Aggregatibacter* species, *Haemophilus* species, and *C. hominis*. Invasive infection typically occurs in patients with a history of cardiac valvular disease, often in the setting of a recent dental procedure or nasopharyngeal infection. The aortic and mitral valves are most commonly affected. Compared with non-HACEK endocarditis, HACEK endocarditis occurs in younger patients and is more frequently associated with embolic, vascular, and immunologic manifestations but less commonly associated with congestive heart failure. The clinical course of HACEK endocarditis tends to be subacute, particularly with *Aggregatibacter* or *Cardiobacterium*. However, *K. kingae* endocarditis may have a more aggressive presentation. Systemic embolization is common. The overall prevalence of major emboli associated with HACEK endocarditis ranges from 28% to 71% in different series. On echocardiography, valvular vegetations are seen in up to 85% of patients. *Aggregatibacter* and *Haemophilus* species cause mitral valve vegetations most often; *Cardiobacterium* is associated with aortic valve vegetations. The microbiology laboratory should be alerted when a HACEK organism is being considered; however, most cultures that ultimately yield a HACEK organism become positive within the first week, especially with improved culture systems such as BACTEC. In addition, polymerase chain reaction (PCR) techniques (e.g., of cardiac valves) and other tools, such as matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) mass spectrometry performed directly on agar colonies, are facilitating the diagnosis of HACEK infections. Because of the organisms' slow growth, antimicrobial testing may be difficult, and β -lactamase production may not be detected. Resistance is most commonly noted in *Haemophilus* and *Aggregatibacter* species. E-test methodology may increase the accuracy of susceptibility testing. The overall prognosis in HACEK endocarditis is excellent and significantly better than that in endocarditis caused by non-HACEK pathogens.

Haemophilus Species *Haemophilus parainfluenzae* is the most common species isolated from cases of HACEK endocarditis. Of patients with HACEK endocarditis due to *Haemophilus* species, 60% have been ill for <2 months before presentation, and 19–50% develop congestive heart failure. Mortality rates as high as 30–50% were reported in older series; however, more recent studies have documented mortality rates of <5%. *H. parainfluenzae* has been isolated from other infections, such as meningitis; brain, dental, pelvic, and liver abscess; pneumonia; urinary tract infection; and septicemia.

Aggregatibacter Species The species of *Aggregatibacter* that most frequently cause infective endocarditis are *A. actinomycetemcomitans*, *A. (formerly Haemophilus) aphrophilus*, and *A. paraphrophilus*. *Aggregatibacter* is associated with prosthetic valve endocarditis more often than are *Haemophilus* species. *A. actinomycetemcomitans* can be

isolated from soft tissue infections and abscesses in association with *Actinomyces israelii*. Typically, patients who develop endocarditis with *Aggregatibacter* have periodontal disease or have recently undergone dental procedures in the setting of underlying cardiac valvular damage. The disease is insidious; patients may be sick for several months before diagnosis. Frequent complications include embolic phenomena, congestive heart failure, and renal failure. *A. actinomycetemcomitans* has been isolated from patients with brain abscess, meningitis, endophthalmitis, parotitis, osteomyelitis, urinary tract infection, pneumonia, and empyema, among other infections, while *A. aphrophilus* is often associated with bone and joint infection.

Cardiobacterium hominis *C. hominis* primarily causes endocarditis in patients with underlying valvular heart disease or with prosthetic valves. This organism most frequently affects the aortic valve. Many patients have signs and symptoms of long-standing infection before diagnosis, with evidence of arterial embolization, vasculitis, cerebrovascular accidents, immune complex glomerulonephritis, or arthritis at presentation. Embolization, mycotic aneurysms, and congestive heart failure are common complications. A second species, *C. valvarum*, has now been described in association with endocarditis.

Eikenella corrodens *E. corrodens* is most frequently recovered from sites of infection in conjunction with other bacterial species. Clinical sources of *E. corrodens* include sites of human bite wounds (clenched-fist injuries), endocarditis, soft tissue infections, osteomyelitis, head and neck infections, respiratory infections, chorioamnionitis, gynecologic infections associated with intrauterine devices, meningitis, brain abscesses, and visceral abscesses. This organism is the least common cause of HACEK endocarditis.

Kingella kingae Because of improved microbiologic methodology and molecular methods such as real-time PCR, the isolation of *K. kingae* is increasingly common. Inoculation of clinical specimens (e.g., synovial fluid) into aerobic blood culture bottles enhances recovery of this organism. More than half of cases of *K. kingae* infection are bone and joint infections; the majority of the remaining infections are infective endocarditis and bacteremia. PCR studies of joint fluid can identify *K. kingae* in culture-negative cases. Some studies now demonstrate that *K. kingae* has surpassed *Staphylococcus aureus* as the leading cause of septic arthritis in children. Invasive *K. kingae* infections with bacteremia are associated with upper respiratory tract infections and stomatitis in 80% of cases. Rates of oropharyngeal colonization with *K. kingae* are highest in the first 3 years of life (detected in ~10% of children), coinciding with an increased incidence of skeletal infections due to this organism. *K. kingae* bacteremia can present with a petechial rash similar to that seen in *Neisseria meningitidis* sepsis.

Infective endocarditis, unlike other infections with *K. kingae*, occurs in older children and adults. The majority of patients have preexisting valvular disease. There is a high incidence of complications, including arterial emboli, cerebrovascular accidents, tricuspid insufficiency, and congestive heart failure with cardiovascular collapse.

TREATMENT HACEK ENDOCARDITIS

(Table 183e-1) Ceftriaxone (2 g/d) is first-line therapy for HACEK endocarditis. Data on the use of levofloxacin (500–750 mg/d) for HACEK endocarditis remain limited, but this drug can be considered an alternative for treatment of patients intolerant of β -lactam therapy. Of note, *Eikenella* is resistant to clindamycin, metronidazole, and aminoglycosides.

Native-valve endocarditis should be treated for 4 weeks with antibiotics, whereas prosthetic-valve endocarditis requires 6 weeks of therapy. The cure rates for HACEK prosthetic-valve endocarditis appear to be high. Unlike prosthetic-valve endocarditis caused by other gram-negative organisms, HACEK endocarditis is often cured with antibiotic treatment alone—i.e., without surgical intervention.

TABLE 183e-1 TREATMENT OF INFECTIONS CAUSED BY HACEK GROUP ORGANISMS

Organisms	Preferred Therapy	Alternative Agents	Comments
<i>Haemophilus</i> spp.	Ceftriaxone (2 g/d)	Ampicillin/sulbactam (3 g of ampicillin q6h)	Ampicillin/sulbactam resistance has been described in <i>Haemophilus</i> and <i>Aggregatibacter</i> spp.
<i>Aggregatibacter actinomycetemcomitans</i> , <i>A. aphrophilus</i> , <i>A. paraphrophilus</i> , other species		Levofloxacin (500–750 mg/d)	Data on use of levofloxacin for endocarditis therapy are limited. Fluoroquinolones are not recommended for treatment of patients <18 years of age.
<i>Cardiobacterium hominis</i>			
<i>Eikenella corrodens</i>		Penicillin (16–18 million units q4h) or ampicillin (2 g q4h)	Penicillin or ampicillin can be used if the organism is susceptible. However, because of the slow growth of HACEK bacteria, antimicrobial testing may be difficult, and β -lactamase production may not be detected.
<i>Kingella kingae</i>			

OTHER GRAM-NEGATIVE BACTERIA

Achromobacter xylosoxidans *Achromobacter* (previously *Alcaligenes xylosoxidans*) is probably part of the endogenous intestinal flora and has been isolated from a variety of water sources, including well water, IV fluids, and humidifiers. Immunocompromised hosts, including patients with cancer and postchemotherapy neutropenia, cirrhosis, chronic renal failure, and cystic fibrosis, are at increased risk. Nosocomial outbreaks and pseudo-outbreaks of *A. xylosoxidans* infection have been attributed to contaminated fluids, and clinical illness has been associated with isolates from many sites, including blood (often in the setting of intravascular devices). Community-acquired *A. xylosoxidans* bacteremia usually occurs in the setting of pneumonia. Metastatic skin lesions are present in one-fifth of cases. The reported mortality rate is as high as 67%—a figure similar to rates for other bacteremic gram-negative pneumonias.

TREATMENT ACHROMOBACTER XYLOSOXIDANS INFECTIONS

(Table 183e-2) Treatment is based on in vitro susceptibility testing of all clinically relevant isolates; multidrug resistance is common. Meropenem, tigecycline, and colistin are typically the most active agents.

Aeromonas Species More than 85% of *Aeromonas* infections are caused by *A. hydrophila*, *A. caviae*, and *A. veronii* biovar *sobria*. *Aeromonas* proliferates in potable water, freshwater, and soil. It remains controversial whether *Aeromonas* is a cause of bacterial gastroenteritis; asymptomatic colonization of the intestinal tract with *Aeromonas* occurs frequently. However, rare cases of hemolytic-uremic syndrome following bloody diarrhea have been shown to be secondary to the presence of *Aeromonas*.

Aeromonas causes sepsis and bacteremia in infants with multiple medical problems and in immunocompromised hosts, particularly those with cancer or hepatobiliary disease. *A. caviae* is associated with health care–related bacteremia. *Aeromonas* infection and sepsis can occur in patients with trauma (including severe trauma with myonecrosis) and in burn patients exposed to *Aeromonas* by environmental (freshwater or soil) contamination of their wounds. Reported mortality rates range from 25% among immunocompromised adults with sepsis to >90% among patients with myonecrosis. *Aeromonas* can produce ecthyma gangrenosum (hemorrhagic vesicles surrounded by a rim of erythema with central necrosis and ulceration; see Fig. 25e-35) resembling the lesions seen in *Pseudomonas aeruginosa* infection. This organism causes nosocomial infections related to catheters, surgical incisions, or use of leeches. Other manifestations include necrotizing fasciitis, meningitis, peritonitis, pneumonia, and ocular infections.

TREATMENT AEROMONAS INFECTIONS

(Table 183e-2) *Aeromonas* species are generally susceptible to fluoroquinolones (e.g., ciprofloxacin at a dosage of 500 mg every 12 h PO or 400 mg every 12 h IV), third- and fourth-generation cephalosporins,

carbapenems, and aminoglycosides, but resistance has been described to all those agents. Because *Aeromonas* can produce various β -lactamases, including carbapenemases, susceptibility testing must be used to guide therapy. Antibiotic prophylaxis (e.g., with ciprofloxacin) is indicated when medicinal leeches are used.

Capnocytophaga Species This genus of fastidious, fusiform, gram-negative coccobacilli is facultatively anaerobic and requires an atmosphere enriched in carbon dioxide for optimal growth. *C. ochracea*, *C. gingivalis*, *C. haemolytica*, and *C. sputigena* have been associated with sepsis in immunocompromised hosts, particularly neutropenic patients with oral ulcerations. These species have been isolated from many other sites as well, usually as part of a polymicrobial infection. Most *Capnocytophaga* infections are contiguous with the oropharynx (e.g., periodontal disease, respiratory tract infections, cervical abscesses, and endophthalmitis).

TABLE 183e-2 TREATMENT OPTIONS FOR OTHER SELECTED GRAM-NEGATIVE BACTERIA

Organism	Treatment Options	Comments
<i>Achromobacter xylosoxidans</i>	Meropenem, tigecycline, colistin	Treatment should be based on in vitro susceptibility testing; multidrug resistance is common among these organisms.
<i>Aeromonas</i> spp.	Fluoroquinolones (e.g., ciprofloxacin), third- and fourth-generation cephalosporins, carbapenems, aminoglycosides	
<i>Elizabethkingia/Chryseobacterium</i> spp.	Fluoroquinolones, trimethoprim-sulfamethoxazole, piperacillin/tazobactam	
<i>Rhizobium radiobacter</i>	Fluoroquinolones, third- and fourth-generation cephalosporins, carbapenems	
<i>Shewanella</i> spp.	Fluoroquinolones, third- and fourth-generation cephalosporins, β -lactam/ β -lactamase inhibitors, carbapenems, aminoglycosides	
<i>Capnocytophaga</i> spp.	Ampicillin/sulbactam	Penicillin should be used if the isolate is known to be susceptible.
<i>Pasteurella multocida</i>	Ampicillin/sulbactam or ceftriaxone	Penicillin should be used if the isolate is known to be susceptible. <i>P. multocida</i> is also susceptible to tetracyclines and fluoroquinolones.

C. canimorsus and *C. cynodegmi* are endogenous to the canine mouth (Chap. 167e). Patients infected with these species frequently have a history of dog bites or of canine exposure without scratches or bites. Asplenia, glucocorticoid therapy, and alcohol abuse are predisposing conditions that can be associated with severe sepsis with shock and disseminated intravascular coagulation. Patients typically have a petechial rash that can progress from purpuric lesions to gangrene. Meningitis, endocarditis, cellulitis, osteomyelitis, and septic arthritis also have been associated with these organisms.

TREATMENT CAPNOCYTOPHAGA INFECTIONS

(Table 183e-2) Because of increasing β -lactamase production, a penicillin derivative plus a β -lactamase inhibitor—such as ampicillin/sulbactam (1.5–3.0 g of ampicillin every 6 h)—is currently recommended for empirical treatment of infections caused by *Capnocytophaga* species. If the isolate is known to be susceptible, infections with *C. canimorsus* should be treated with penicillin (12–18 million units every 4 h). *Capnocytophaga* is also susceptible to clindamycin (600–900 mg every 6–8 h). This regimen or ampicillin/sulbactam should be given prophylactically to asplenic patients who have sustained dog-bite injuries.

Elizabethkingia/Chryseobacterium Species *Elizabethkingia meningoseptica* (formerly *Chryseobacterium meningosepticum*) is an important cause of nosocomial infections, including outbreaks due to contaminated fluids (e.g., disinfectants and aerosolized antibiotics) and sporadic infections due to indwelling devices, feeding tubes, and other fluid-associated apparatuses. Nosocomial *E. meningoseptica* infection usually involves neonates or patients with underlying immunosuppression (e.g., related to malignancy or diabetes). *E. meningoseptica* has been reported to cause meningitis (primarily in neonates), pneumonia, sepsis, endocarditis, bacteremia, and soft tissue infections. Most published reports have originated from Taiwan. *Chryseobacterium indologenes* has caused bacteremia, sepsis, and pneumonia, typically in immunocompromised patients with indwelling devices.

TREATMENT ELIZABETHKINGIA/CHRYSEOBACTERIUM INFECTIONS

(Table 183e-2) These organisms are often susceptible to fluoroquinolones and trimethoprim-sulfamethoxazole. They may be susceptible to β -lactam/ β -lactamase inhibitor combinations such as piperacillin/tazobactam but can possess extended-spectrum β -lactamases and metallo- β -lactamases. Susceptibility testing should be performed.

Pasteurella multocida *P. multocida* is a bipolar-staining, gram-negative coccobacillus that colonizes the respiratory and gastrointestinal tracts of domestic animals; oropharyngeal colonization rates are 70–90% in

cats and 50–65% in dogs. *P. multocida* can be transmitted to humans through bites or scratches, via the respiratory tract from contact with contaminated dust or infectious droplets, or via deposition of the organism on injured skin or mucosal surfaces during licking. Most human infections affect skin and soft tissue; almost two-thirds of these infections are caused by cats. Patients at the extremes of age or with serious underlying disorders (e.g., cirrhosis, diabetes) are at increased risk for systemic manifestations, including meningitis, peritonitis, osteomyelitis and septic arthritis, endocarditis, and septic shock, but cases have also occurred in healthy individuals of all ages. If inhaled, *P. multocida* can cause acute respiratory tract infection, particularly in patients with underlying sinus and pulmonary disease.

TREATMENT PASTEURELLA MULTOCIDA INFECTIONS

P. multocida is susceptible to penicillin, ampicillin, ampicillin/sulbactam, second- and third-generation cephalosporins, tetracyclines, and fluoroquinolones. β -lactamase-producing strains have been reported.

MISCELLANEOUS ORGANISMS

Rhizobium (formerly *Agrobacterium*) *radiobacter* has usually been associated with infection in the presence of medical devices, including intravascular catheter-related infections, prosthetic-joint and prosthetic-valve infections, and peritonitis caused by dialysis catheters. Cases of endophthalmitis after cataract surgery also have been described. Most *R. radiobacter* infections occur in immunocompromised hosts, especially individuals with malignancy or HIV infection. Strains are usually susceptible to fluoroquinolones, third- and fourth-generation cephalosporins, and carbapenems (Table 183e-2).

Shewanella putrefaciens and *S. algae* are ubiquitous organisms found primarily in seawater. Devitalized tissues can become colonized with *Shewanella* and serve as a nidus for systemic infection. *Shewanella* species cause skin and soft tissue infections, chronic ulcers of the lower extremities, ear infections, biliary tract infections, pneumonia, necrotizing fasciitis, bacteremia, and sepsis. A fulminant course is associated with cirrhosis, diabetes mellitus, malignancy, or other severe underlying conditions. Organisms are often susceptible to fluoroquinolones, third- and fourth-generation cephalosporins, β -lactam/ β -lactamase inhibitors, carbapenems, and aminoglycosides (Table 183e-2).

Chromobacterium violaceum has been responsible for life-threatening infections with severe sepsis and metastatic abscesses, particularly in children with defective neutrophil function (e.g., those with chronic granulomatous disease). *Ochrobactrum anthropi* causes infections related to central venous catheters in compromised hosts; other invasive infections have been described. Other organisms implicated in human infections include *Weeksella* species; various CDC groups, such as Ve-1 and Ve-2; *Flavimonas* species; *Sphingobacterium* species; and *Oligella urethralis*. The reader is advised to consult subspecialty texts and references for further guidance on these organisms.

184 Legionella Infections

Victor L. Yu, M. Luisa Pedro-Botet, Yusen E. Lin

Legionellosis refers to the two clinical syndromes caused by bacteria of the genus *Legionella*. *Pontiac fever* is an acute, febrile, self-limited illness that has been serologically linked to *Legionella* species, whereas *Legionnaires' disease* is the designation for pneumonia caused by these species. Legionnaires' disease was first recognized in 1976, when an outbreak of pneumonia took place at a Philadelphia hotel during an American Legion convention.

MICROBIOLOGY

The family Legionellaceae comprises more than 50 species with more than 70 serogroups. The species *L. pneumophila* causes 80–90% of human infections and includes at least 16 serogroups; serogroups 1, 4, and 6 are most commonly implicated in human infections. To date, 18 species other than *L. pneumophila* have been associated with human infections, among which *L. micdadei* (Pittsburgh pneumonia agent), *L. bozemanii*, *L. dumoffii*, and *L. longbeachae* are the most common. Members of the Legionellaceae are aerobic gram-negative bacilli that do not grow on routine microbiologic media. Buffered charcoal yeast extract (BCYE) agar is the medium used to grow *Legionella*.

ECOLOGY AND TRANSMISSION



The natural habitats for *L. pneumophila* are aquatic bodies, including lakes and streams. *L. longbeachae* has been isolated from natural soil. Commercial potting soil has been suggested as the reservoir for *L. longbeachae* infections in Australia and New Zealand. *Legionella* can survive under a wide range of environmental conditions; for example, the organisms can live for years in refrigerated water samples. Natural bodies of water contain only small numbers of legionellae. However, once the organisms enter human-constructed aquatic reservoirs (such as drinking-water systems), they can grow and proliferate. Factors known to enhance colonization by and amplification of legionellae include warm temperatures (25°–42°C) and the presence of scale and sediment. *L. pneumophila* can form microcolonies within biofilms; its eradication from drinking-water systems requires disinfectants that can penetrate the biofilm. The presence

of symbiotic microorganisms, including algae, amebas, ciliated protozoa, and other water-dwelling bacteria, promotes the growth of *Legionella*. The organisms can invade and multiply within free-living protozoa.

Heavy rainfall and flooding can result in the entry of high numbers of legionellae into water-distribution systems, leading to an upsurge of cases. Large buildings over three stories high are commonly colonized with *Legionella*. Sporadic community-acquired Legionnaires' disease has been linked to colonization of hotels, office buildings, factories, and even private homes. Drinking-water systems in hospitals and extended-care facilities have been the source for health care-associated Legionnaires' disease.

In contrast, cooling towers and evaporative condensers have been overestimated as sources of *Legionella* causing human illness. Early investigations that implicated cooling towers antedated the discovery that the organism could also exist in drinking water. In many outbreaks attributed to cooling towers, cases of Legionnaires' disease continued to occur despite disinfection of the towers; drinking water was found to be the actual source. Koch's postulates have never been fulfilled for *Legionella* links to cooling tower-associated outbreaks as they have been for hospital-acquired Legionnaires' disease. Nevertheless, cooling towers have, in rare instances, been implicated in community-acquired outbreaks, including an outbreak in Murcia, Spain. As mentioned above, *L. longbeachae* infections have been linked to potting soil, but the mode of transmission remains to be clarified.

Multiple modes of transmission of *Legionella* to humans exist, including aerosolization, aspiration, and direct instillation into the lungs during respiratory tract manipulations. Aspiration is now known to be the predominant mode of transmission, but it is unclear whether *Legionella* enters the lungs via oropharyngeal colonization or directly via the drinking of contaminated water. Oropharyngeal colonization with *Legionella* has been demonstrated in patients undergoing transplantation. Nasogastric tubes have been linked to hospital-acquired Legionnaires' disease; microaspiration of contaminated water was the hypothesized mode of transmission. Surgery with general anesthesia is a known risk factor that is consistent with aspiration. Especially compelling is the reported 30% incidence of postoperative Legionnaires' disease among patients undergoing head and neck surgery at a hospital with a contaminated water supply; aspiration is a recognized postoperative complication in such cases. One observational study showed that patients with hospital-acquired Legionnaires' disease underwent endotracheal intubation significantly more often and for a significantly longer duration than patients with hospital-acquired pneumonias of other etiologies.

Aerosolization of *Legionella* by devices filled with tap water, including whirlpools, nebulizers, and humidifiers, has been linked to cases in patients. An ultrasonic mist machine in the produce section of a grocery store has been the source in community outbreaks. Pontiac fever has been linked to *Legionella*-containing aerosols from water-using machinery, a cooling tower, air conditioners, and whirlpools.

EPIDEMIOLOGY



Community-Acquired Pneumonia The incidence of Legionnaires' disease depends on the degree of contamination of the aquatic reservoir, the immune status of the persons exposed to water from that reservoir, the intensity of exposure, and the availability of specialized laboratory tests on which the correct diagnosis can be based. Numerous prospective studies have ranked *Legionella* among the top four microbial causes of community-acquired pneumonia, finding that it accounts for 2–13% of cases. (*Streptococcus pneumoniae*, *Haemophilus influenzae*, and *Chlamydia pneumoniae* are usually ranked first, second, and third, respectively.) On the basis of a multihospital study of community-acquired pneumonia in Ohio, the Centers for Disease Control and Prevention (CDC) estimated that only 3% of community-acquired cases of Legionnaires' disease are diagnosed as such. Observational studies of community-acquired pneumonia showed that Legionnaires' disease was largely unrecognized unless *Legionella* diagnostic testing was routinely applied to all

patients with pneumonia; such studies in Spain and Germany resulted in detection of increased numbers of cases throughout Europe. It is likely that observational studies in Taiwan and Australia will have a similar result, with more cases identified throughout Asia as the index of suspicion rises.

Hospital-Acquired Pneumonia *Legionella* is responsible for 10–50% of cases of nosocomial pneumonia when a hospital's water system is colonized with the organism. The incidence of hospital-acquired Legionnaires' disease depends on the degree of contamination of drinking water, as defined by the rate of positivity of distal water sites; in contrast, the use of quantitative criteria of the number of colony-forming units per milliliter has proven useless.



Proactive culture of the hospital water supply has increased the detection of hospital-acquired Legionnaires' disease and simultaneously allowed expeditious diagnosis resulting in early administration of antibiotic therapy. In the early years after its recognition, Legionnaires' disease was documented primarily in the United States. As diagnostic modalities (especially the urinary antigen test) became more widely used, cases were documented in European hospitals. Likewise, following the enactment of public health guidelines in Taiwan, cases attributable to hospital tap water were found in Taiwanese hospitals. Risk factors for Legionnaires' disease include cigarette smoking, chronic lung disease, advanced age, prior hospitalization with discharge within 10 days before onset of pneumonia symptoms, and immunosuppression. Immunosuppressive conditions that predispose to Legionnaires' disease include transplantation, HIV infection, and treatment with glucocorticoids or tumor necrosis factor α antagonists. However, in a large prospective study of community-acquired pneumonia, 28% of patients with Legionnaires' disease did not have these classic risk factors. Hospital-acquired cases are now being recognized among neonates and immunosuppressed children.

Pneumonia in Transplant Recipients Transplant recipients appear to be at unusually high risk of *Legionella* pneumonia. This elevated risk may be due to diagnostic bias, given the extensive workup for opportunistic pathogens with pneumonic symptoms as well as the long-standing immunosuppression in this population of patients. Legionnaires' disease usually occurs in the 3 months after transplantation. Cavitation is seen on chest radiograph more frequently in transplant recipients, and mortality rates are higher.

Pontiac Fever Pontiac fever occurs in epidemics. The high attack rate (>90%) reflects airborne transmission.

PATHOGENESIS AND IMMUNITY

Legionella enters the lungs through aspiration or direct inhalation. Attachment to host cells is mediated by bacterial type IV pili, heat-shock proteins, a major outer-membrane protein, and complement. Because the organism possesses pili that mediate adherence to respiratory tract epithelial cells, conditions that impair mucociliary clearance, including cigarette smoking, lung disease, or alcoholism, predispose to Legionnaires' disease.

Both innate and adaptive immune responses play a role in host defense. Toll-like receptors mediate recognition of *L. pneumophila* in alveolar macrophages and enhance early neutrophil recruitment to the site of infection. Alveolar macrophages phagocytose legionellae by a conventional or a coiling mechanism. After phagocytosis, *L. pneumophila* evades intracellular killing by inhibiting phagosomal-lysosome fusion. Although many legionellae are killed, some proliferate intracellularly until the cells rupture; the bacteria are then phagocytosed again by newly recruited phagocytes, and the cycle begins anew. The role of neutrophils in immunity appears to be minimal: neutropenic patients are not predisposed to Legionnaires' disease. Although *L. pneumophila* is susceptible to oxygen-dependent microbiologic systems in vitro, it resists killing by neutrophils. The humoral immune system is active against *Legionella*. Type-specific IgM and IgG antibodies are measurable within weeks of infection. In vitro, antibodies promote killing of *Legionella* by phagocytes (neutrophils, monocytes, and alveolar macrophages). Immunized animals

1016 develop a specific antibody response, with subsequent resistance to *Legionella* challenge. However, antibodies neither enhance lysis by complement nor inhibit intracellular multiplication within phagocytes.



The genome of *L. pneumophila* has been sequenced. A broad range of membrane transporters within the genome are thought to optimize the use of nutrients in water and soil.

Some *L. pneumophila* strains are clearly more virulent than others, although the precise factors mediating virulence remain uncertain. For example, although multiple strains may colonize water-distribution systems, only a few cause disease in patients exposed to water from these systems. At least one surface epitope of *L. pneumophila* serogroup 1 is associated with virulence. Monoclonal antibody subtype mAb2 has been linked to virulence. *L. pneumophila* serogroup 6 is more commonly involved in hospital-acquired Legionnaires' disease and is especially likely to be associated with a poor outcome.

CLINICAL AND LABORATORY FEATURES

Pontiac Fever Pontiac fever is an acute, self-limiting, flu-like illness with an incubation period of 24–48 h. Pneumonia does not develop. Malaise, fatigue, and myalgias are the most common symptoms, occurring in 97% of cases. Fever (usually with chills) develops in 80–90% of cases and headache in 80%. Other symptoms (seen in fewer than 50% of cases) include arthralgias, nausea, cough, abdominal pain, and diarrhea. Modest leukocytosis with a neutrophilic predominance is sometimes detected. Complete recovery occurs within a few days; antibiotic therapy is unnecessary. A few patients may experience lassitude for some weeks after recovery. The diagnosis is established by antibody seroconversion. Pontiac fever due to *L. longbeachae* has been reported in individuals exposed to potting soil.

Legionnaires' Disease (Pneumonia) Legionnaires' disease is often included in the differential diagnosis of "atypical pneumonia," along with pneumonia due to *C. pneumoniae*, *Chlamydia psittaci*, *Mycoplasma pneumoniae*, *Coxiella burnetii*, and some viruses. The clinical similarities among "atypical" pneumonias include a nonproductive cough with a low frequency of grossly purulent sputum. The clinical manifestations of Legionnaires' disease are usually more severe than those of most "atypical" pneumonias. The course and prognosis of *Legionella* pneumonia more closely resemble those of bacteremic pneumococcal pneumonia than those of pneumonia due to other "atypical" pathogens. Patients with community-acquired Legionnaires' disease are significantly more likely than patients with pneumonia of other etiologies to be admitted to an intensive care unit (ICU) on presentation.

The incubation period for Legionnaires' disease is usually 2–10 days, although slightly longer incubation periods have been documented. Fever is almost universal. In one observational study, 20% of patients had temperatures in excess of 40°C (104°F). The symptoms and signs may range from a mild cough and a slight fever to stupor with widespread pulmonary infiltrates and multisystem failure. The mild cough of Legionnaires' disease is only slightly productive. Sometimes the sputum is streaked with blood. Chest pain—either pleuritic or nonpleuritic—can be a prominent feature and, when coupled with hemoptysis, can lead to an incorrect diagnosis of pulmonary embolism. Shortness of breath is reported by one-third to one-half of patients. Gastrointestinal difficulties are often pronounced; abdominal pain, nausea, and vomiting affect 10–20% of patients. Diarrhea (watery rather than bloody) is reported in 25–50% of cases. The most common neurologic abnormalities are confusion or changes in mental status; however, the multitudinous neurologic symptoms reported range from headache and lethargy to encephalopathy. Nonspecific symptoms—malaise, fatigue, anorexia, and headache—are reported early in the illness. Myalgias and arthralgias are uncommon but are prominent in a few patients. Upper respiratory symptoms, including coryza, are rare.

Relative bradycardia has been overemphasized as a useful diagnostic finding; it occurs primarily in older patients with severe pneumonia. Rales are detected by chest examination early in the course, and evidence of consolidation is found as the disease progresses. Abdominal examination may reveal generalized or local tenderness.

TABLE 184-1 CLINICAL CLUES SUGGESTIVE OF LEGIONNAIRES' DISEASE

Diarrhea
High fever (>40°C; >104°F)
Numerous neutrophils but no organisms revealed by Gram's staining of respiratory secretions
Hyponatremia (serum sodium level <131 mg/dL)
Failure to respond to β -lactam drugs (penicillins or cephalosporins) and aminoglycoside antibiotics
Occurrence of illness in an environment in which the potable water supply is known to be contaminated with <i>Legionella</i>
Onset of symptoms within 10 days after discharge from the hospital (hospital-acquired legionellosis manifesting after discharge or transfer)

Although the clinical manifestations often considered classic for Legionnaires' disease may suggest the diagnosis (Table 184-1), prospective comparative studies have shown that clinical manifestations are generally nonspecific and that Legionnaires' disease is not readily distinguishable from pneumonia of other etiologies. In a review of 13 studies of community-acquired pneumonia, clinical manifestations that occurred significantly more often in Legionnaires' disease included diarrhea, neurologic findings (including confusion), and a temperature of >39°C. Hyponatremia, elevated values in liver function tests, and hematuria also occurred more frequently in Legionnaires' disease. Other laboratory abnormalities include creatine phosphokinase elevation, hypophosphatemia, serum creatinine elevation, and proteinuria.

Sporadic cases of Legionnaires' disease appear to be more severe than outbreak-associated and hospital-acquired cases, presumably because their diagnosis is delayed. Results of the German CAPNETZ Study showed that, among cases of community-acquired *Legionella* pneumonia, ambulatory patients were as common as hospitalized patients.

Extrapulmonary Legionellosis Because the portal of entry for *Legionella* is the lung in virtually all cases, extrapulmonary manifestations usually result from bloodborne dissemination from the lung. *Legionella* has been identified in lymph nodes, spleen, liver, or kidneys in autopsied cases. Sinusitis, peritonitis, pyelonephritis, skin and soft tissue infection, septic arthritis, and pancreatitis have developed predominantly in immunosuppressed patients. The most severe sequela, neurologic dysfunction, is rare but can be debilitating. The most common neurologic deficits in the long term—ataxia and speech difficulties—result from cerebellar dysfunction.

We speculate that cardiac abnormalities in patients without pneumonia are caused by *Legionella*-contaminated water entering through an intravenous site, chest tube, or surgical wound, with subsequent seeding of a prosthetic valve, the myocardium, or the pericardium. This scenario is supported by cases occurring at Stanford University Hospital in which sternal wound infections and prosthetic valve endocarditis due to *L. pneumophila* were observed. The source was a sink in the postoperative surgical recovery ward.

Chest Radiography Virtually all patients with Legionnaires' disease have abnormal chest radiographs showing pulmonary infiltrates at the time of clinical presentation. In a few cases of hospital-acquired disease, fever and respiratory tract symptoms have preceded the radiographic appearance of the infiltrate. Radiologic findings are nonspecific. Pleural effusion is evident in 28–63% of patients on hospital admission. In immunosuppressed patients, especially those receiving glucocorticoids, distinctive rounded nodular opacities may be seen; these lesions may expand and cavitate (Fig. 184-1). Likewise, abscesses can occur in immunosuppressed hosts. The progression of infiltrates and pleural effusion on chest radiography despite appropriate antibiotic therapy within the first week is common, and radiographic improvement lags behind clinical improvement by several days. Complete clearing of infiltrates requires 1–4 months. Computed tomography (CT) is more sensitive than chest radiography, may show more extensive disease, and should be performed if fever persists during treatment with presumably effective antibiotics (Fig. 184-2).



FIGURE 184-1 Chest radiographic findings in a 52-year-old man who presented with pneumonia subsequently diagnosed as Legionnaires' disease. The patient was a cigarette smoker with chronic obstructive pulmonary disease and alcoholic cardiomyopathy; he had received glucocorticoids. *Legionella pneumophila* was identified by direct fluorescent antibody staining and culture of sputum. **Left:** Baseline chest radiograph showing long-standing cardiomegaly. **Center:** Admission chest radiograph showing new rounded opacities. **Right:** Chest radiograph taken 3 days after admission, during treatment with erythromycin.

DIAGNOSIS

Given the nonspecific clinical manifestations of Legionnaires' disease and the high mortality rates for untreated Legionnaires' disease, *Legionella* testing—especially the *Legionella* urinary antigen test—is recommended for all patients with pneumonia, including patients with ambulatory pneumonia and hospitalized children. *Legionella* cultures should be made more widely available because the urinary antigen test can diagnose only *L. pneumophila* serogroup 1. Hospitals in which the drinking water is known to be colonized with *Legionella* species should have *Legionella* cultures routinely available.

The diagnosis of Legionnaires' disease requires special microbiologic tests (Table 184-2). The sensitivity of bronchoscopy specimens is similar to that of sputum samples for culture on selective media; if sputum is not available, bronchoscopy specimens may yield the organism. Bronchoalveolar lavage fluid gives higher yields than bronchial wash specimens. Thoracentesis should be performed if pleural effusion is found, and the fluid should be evaluated by direct fluorescent antibody (DFA) staining, culture, and the antigen assay designed for use with urine.

Stains Gram's staining of material from normally sterile sites, such as pleural fluid or lung tissue, occasionally suggests the diagnosis; efforts to detect *Legionella* in sputum by Gram's staining typically reveal numerous leukocytes but no organisms. When they are visualized, the organisms appear as small, pleomorphic, faint, gram-negative bacilli. *L. micdadei* organisms can be detected as weakly or partially acid-fast bacilli in clinical specimens.

The DFA stain is rapid and highly specific but is less sensitive than culture because large numbers of organisms are required for microscopic visualization. This test is more likely to be positive in advanced than in early disease.

Culture The definitive method for diagnosis of *Legionella* infection is isolation of the organism from respiratory secretions, although culture for 3–5 days is required. Antibiotics added to the medium suppress the growth of competing flora from nonsterile sites, and dyes color the colonies and assist in identification. The use of multiple selective BCYE media is necessary for maximal sensitivity. When culture plates are overgrown with other microflora, pretreatment of the specimen with acid or heat can markedly improve the yield. *L. pneumophila* is often isolated from sputum that is not grossly or microscopically purulent; sputum containing more than 25 epithelial cells per high-power field (a finding that classically suggests contamination) may still yield *L. pneumophila*.

Antibody Detection Antibody testing of both acute- and convalescent-phase sera is necessary. A fourfold rise in titer is diagnostic; 12 weeks are often required for the detection of an antibody response. A single titer of 1:128 in a patient with pneumonia constitutes circumstantial evidence for Legionnaires' disease. The CDC uses a titer of 1:256 as presumptive evidence for Legionnaires' disease. Serology is of use

primarily in epidemiologic studies. The specificity of serology for *Legionella* species other than *L. pneumophila* is uncertain; there is cross-reactivity within *Legionella* species and with some gram-negative bacilli. Serology is used as the criterion for the diagnosis of Pontiac fever.

Urinary Antigen The assay for *Legionella* soluble antigen in urine is second only to culture in terms of sensitivity and is highly specific. A rapid immunochromatographic assay is commercially available (BinaxNOW; Alere, Waltham, MA). This assay is relatively inexpensive and easy to perform. Its drawback is that the urinary antigen test is reliable only for *L. pneumophila* serogroup 1, which causes ~80% of *Legionella* infections. Cross-reactivity with other *L. pneumophila* serogroups and other *Legionella* species has been detected in up to 22% of urine samples from patients with culture-proven cases. Antigen in urine is detectable 3 days after the onset of clinical disease and disappears over 2 months; positivity can be prolonged when patients receive glucocorticoids. The test is not affected by antibiotic administration.

Molecular Methods DFA stains can identify a number of *Legionella* species. Both polyclonal and monoclonal antibody stains are commercially available. Polymerase chain reaction (PCR) with DNA probes is being applied in-house in selected hospitals but is not yet commercially available. PCR has proven somewhat useful in the identification of *Legionella* from environmental water specimens. Epidemiologic links cannot easily be made with PCR because the infecting pathogen is not available for molecular subtyping.

Procalcitonin can be used as an indicator of severity of illness in patients in ICUs. Clinical response to antibiotics can be monitored by procalcitonin levels.

TREATMENT LEGIONELLA INFECTION

Because *Legionella* is an intracellular pathogen, antibiotics that can attain high intracellular concentrations are most likely to be effective. The dosages for various drugs used in the treatment of *Legionella* infection are listed in Table 184-3.

The macrolides (especially azithromycin) and the respiratory quinolones are now the antibiotics of choice and are effective as monotherapy. Compared with erythromycin, the newer macrolides have superior in vitro activity, display greater intracellular activity, reach higher concentrations in respiratory secretions and lung tissue, and have fewer adverse effects. The pharmacokinetics of the newer macrolides and quinolones also allow once- or twice-daily dosing. Quinolones are the preferred antibiotics for transplant recipients because both macrolides and rifampin interact pharmacologically with cyclosporine and tacrolimus. Retrospective uncontrolled studies have shown that complications of pneumonia are fewer and clinical response is more rapid in patients receiving quinolones than in those receiving macrolides. Initial therapy should be given by the

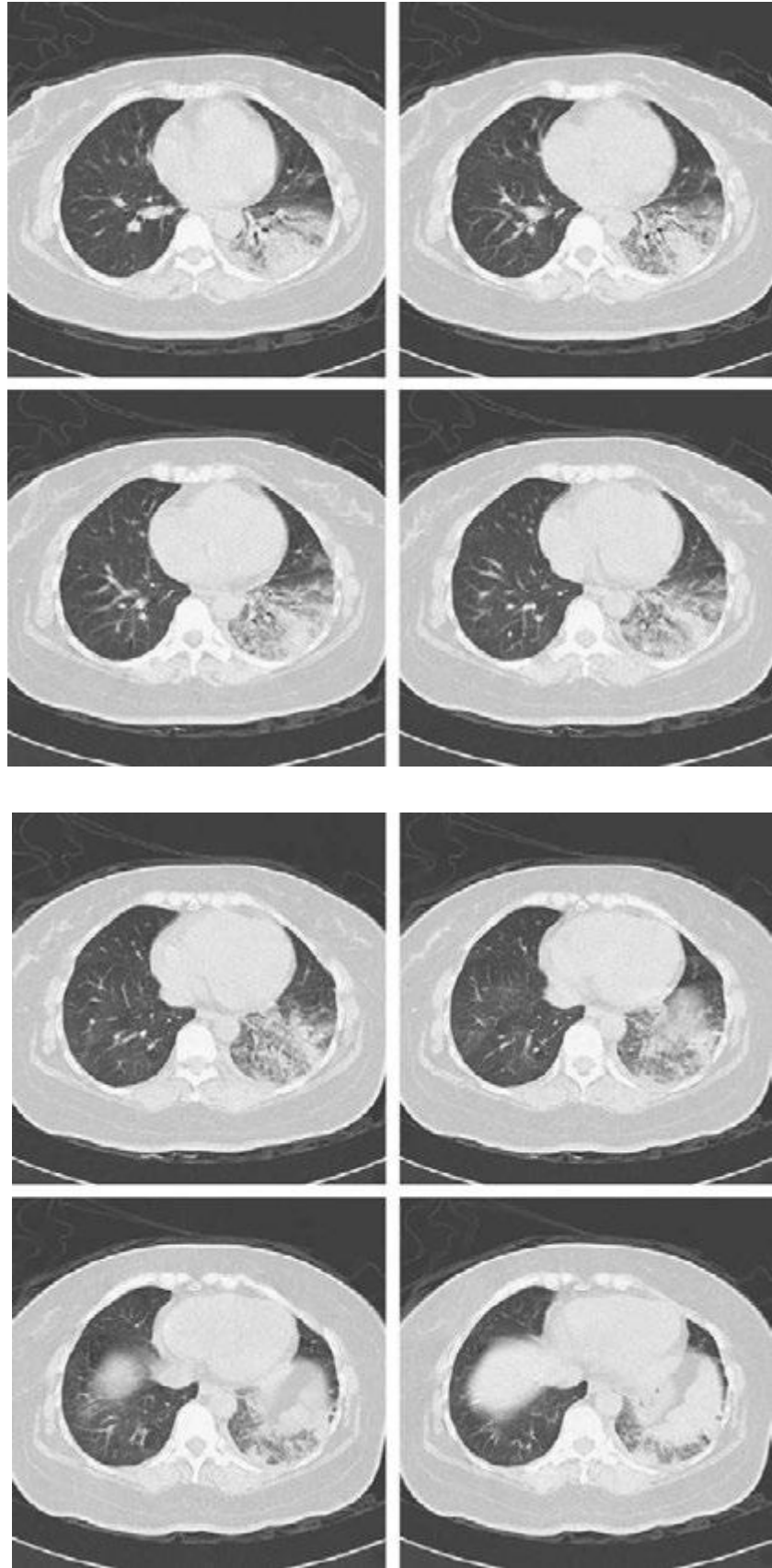


FIGURE 184-2 Computed tomography (CT) scans of a 49-year-old woman with no underlying conditions who presented with community-acquired pneumonia. CT revealed multilobar infiltrates, some of which were not as prominent on chest x-ray. Cultures of both the patient's sputum and her home water supply yielded *Legionella pneumophila* serogroup 1. (Images courtesy of Dr. Wen-Chien Ko, National Cheng Kung University Hospital, Tainan, Taiwan.)

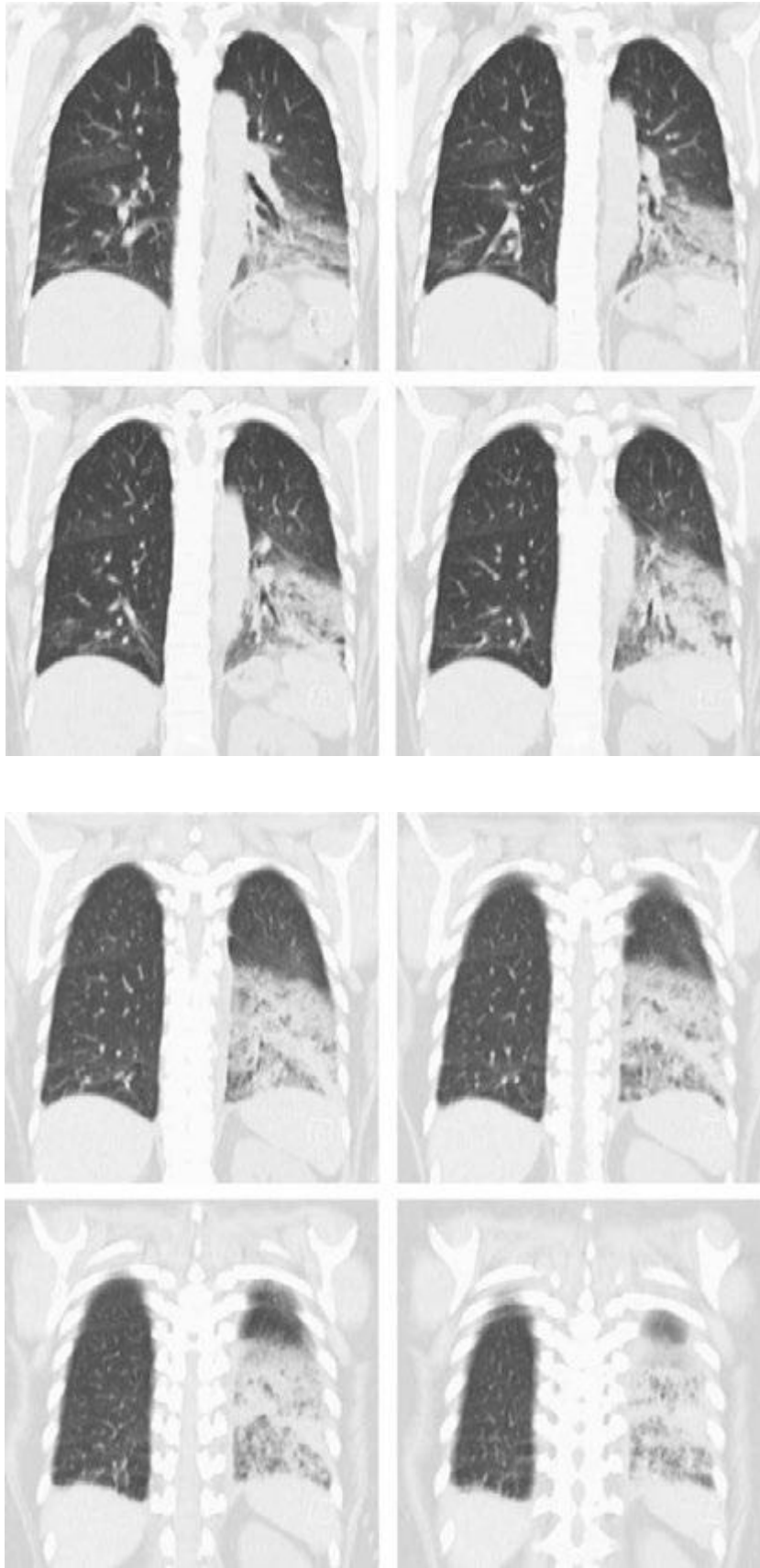


FIGURE 184-2 (Continued)

TABLE 184-2 UTILITY OF SPECIAL LABORATORY TESTS FOR THE DIAGNOSIS OF LEGIONNAIRES' DISEASE

Test	Sensitivity, %	Specificity, %
Culture		
Sputum ^a	80	100
Transtracheal aspirate	90	100
Direct fluorescent antibody staining of sputum	50–70	96–99
Urinary antigen testing ^b	70	100
Antibody serology ^c	40–60	96–99

^aUse of multiple selective media with dyes. ^bReliable only for *L. pneumophila* serogroup 1. ^cIgG and IgM testing of both acute- and convalescent-phase sera. A single titer of $\geq 1:256$ is considered presumptive, whereas a fourfold rise in titer between the acute and convalescent phases is considered definitive. Titers peak at 3 months.

IV route. A clinical response usually occurs within 3–5 days, after which oral therapy can be substituted. The total duration of therapy in the immunocompetent host is 10–14 days.

Alternative agents include tetracycline and its analogues doxycycline and minocycline. Tigecycline is active in vitro, but clinical experience with this drug is minimal. Anecdotal reports have described both successes and failures with trimethoprim-sulfamethoxazole, imipenem, and clindamycin.

For critically ill patients, the authors use combination regimens of azithromycin, a quinolone, and/or rifampin. This practice is empirical and is not supported by comparative studies. Rifampin is highly active in vitro and in cell models. Its interaction with other medications and its side effect of reversible hyperbilirubinemia can be minimized by limiting the duration of therapy to 3–5 days. A longer course of therapy (3 weeks) may be appropriate for immunosuppressed patients and those with advanced disease. For azithromycin, with its long half-life, a 5- to 10-day course is sufficient.

Pontiac fever requires only symptom-based treatment, not antimicrobial therapy.

PROGNOSIS

Mortality rates for Legionnaires' disease vary with the patient's underlying disease, the patient's immune status, the severity of pneumonia, and the timing of administration of appropriate antimicrobial therapy.

TABLE 184-3 ANTIBIOTIC THERAPY FOR LEGIONELLA INFECTION

Antimicrobial Agent	Dosage ^a
Macrolides	
Azithromycin	500 mg ^b PO or IV ^c q24h
Clarithromycin	500 mg PO or IV ^c q12h
Quinolones	
Levofloxacin	750 mg IV q24h
	500 mg ^b PO q24h
Ciprofloxacin	400 mg IV q8h
	750 mg PO q12h
Moxifloxacin	400 mg ^b PO q24h
Ketolide	
Telithromycin	800 mg PO q24h
Tetracyclines	
Doxycycline	100 mg ^b PO or IV q12h
Minocycline	100 mg ^b PO or IV q12h
Tetracycline	500 mg PO or IV q6h
Tigecycline	100-mg IV load, then 50 mg IV q12h
Others	
Trimethoprim-sulfamethoxazole	160/800 mg IV q8h
	160/800 mg PO q12h
Rifampin ^d	300–600 mg PO or IV q12h

^aDosages are derived from clinical experience. ^bThe authors recommend doubling the first dose. ^cThe IV formulation is not available in some countries. ^dRifampin should be used only in combination with a macrolide or a quinolone.

Mortality rates are highest (80%) among immunosuppressed patients who do not receive appropriate antimicrobial therapy early in the course of illness. With timely antibiotic treatment, mortality rates from community-acquired Legionnaires' disease among immunocompetent patients range from 0 to 11%; without treatment, the figure may be as high as 31%. In a study of survivors of an outbreak of community-acquired Legionnaires' disease, sequelae of fatigue, neurologic symptoms, and weakness were found in 63–75% of patients 17 months after receipt of antibiotics.

PREVENTION

Routine environmental culture of hospital water supplies for *Legionella* is recommended as an approach to the prevention of hospital-acquired Legionnaires' disease. Guidelines mandating this proactive approach have been adopted throughout Europe and in several U.S. states. The presence of *Legionella* in the water supply mandates the use of specialized laboratory tests (especially culture on selective media and the urinary antigen test) for patients with hospital-acquired pneumonia. A 30% cutoff for the presence of *Legionella* in water from multiple hospital sites prompts an increased index of suspicion. When the 30% cutoff point is exceeded, diagnostic tests for *Legionella* need to be applied in all cases of hospital-acquired pneumonia, and measures directed at eliminating the organism from the water supply should be considered. Quantitative criteria at a given water site (colony-forming units [CFU]/mL) have proven unreliable and inconsistent in the prediction of disease.

Studies have shown that neither a high degree of outward cleanliness of the water system nor routine application of maintenance measures decreases the frequency or intensity of *Legionella* contamination. Thus, engineering guidelines and building codes, although routinely advocated as preventive measures, have little impact on the presence of *Legionella*.

Environmental cultures for *Legionella* from cold-water taps, hot-water taps, the hot-water recirculating line, and water-storage tanks will reveal the source of hospital-acquired infections. Disinfection of the hospital drinking-water system is an effective preventive measure for hospital-acquired cases of Legionnaires' diseases because this system is the reservoir for *Legionella*. In geographic areas where the climate is semitropical, cold-water lines may be colonized by *Legionella*.

Copper-silver ionization is a reliable method for eradication of *Legionella*. Unlike the efficacy of chlorine dioxide decontamination and chlorination, that of ionization is not affected by high water temperature. Ionization systems are easy to install, and the ions are odorless, with minimal adverse effects. The efficacy of copper-silver ionization has been documented in hospitals worldwide. A comprehensive review of 10 published studies concluded that copper-silver ionization is effective for *Legionella* control as long as ion levels are monitored. If cold-water colonization by *Legionella* is the source of an outbreak, chlorine dioxide and monochloramine offer advantages. Chlorine dioxide, often the least expensive option, penetrates biofilms better and is less corrosive than chlorine. The major disadvantage of chlorine dioxide is the need to maintain an effective residual throughout the drinking-water system, especially in the hot-water system. Eradication of *Legionella* by chlorine dioxide may require several months—a drawback in outbreak situations. Monochloramine is a promising approach in disinfection. Hyperchlorination is no longer recommended because of its expense, carcinogenicity, corrosive effects on piping, and unreliable efficacy.

Point-of-use disposable water filters (0.2 μ m) may be an economical and effective option in high-risk areas (e.g., ICUs and transplantation units). These filters can be used in an outbreak situation for a limited period.

Ineffective yet expensive methods that are often promulgated include removal of stagnation ("dead legs") in the water-distribution system and replacement or disinfection/cleaning of distal outlets. Infection control personnel should oversee the selection of disinfection technology and should apply evidence-based criteria when making their choice. Managers of health care facilities should not be given the primary responsibility for selection and subsequent monitoring of measures to eliminate and control *Legionella*.

Karina A. Top, Scott A. Halperin

Pertussis is an acute infection of the respiratory tract caused by *Bordetella pertussis*. The name *pertussis* means “violent cough,” which aptly describes the most consistent and prominent feature of the illness. The inspiratory sound made at the end of an episode of paroxysmal coughing gives rise to the common name for the illness, “whooping cough.” However, this feature is variable: it is uncommon among infants ≤ 6 months of age and is frequently absent in older children and adults. The Chinese name for pertussis is “the 100-day cough,” which accurately describes the clinical course of the illness. The identification of *B. pertussis* was first reported by Bordet and Gengou in 1906, and vaccines were produced over the following two decades.

MICROBIOLOGY


Of the 10 identified species in the genus *Bordetella*, only four are of major medical significance. *B. pertussis* infects only humans and is the most important *Bordetella* species causing human disease. *B. parapertussis* causes an illness in humans that is similar to pertussis but is typically milder; co-infections with *B. parapertussis* and *B. pertussis* have been documented. With improved polymerase chain reaction (PCR) diagnostic methodology, up to 20% of patients with a pertussis-like syndrome have been found to be infected with *B. holmesii*, formerly thought to be an unusual cause of bacteremia. *B. bronchiseptica* is an important pathogen of domestic animals that causes kennel cough in dogs, atrophic rhinitis and pneumonia in pigs, and pneumonia in cats. Both respiratory infection and opportunistic infection due to *B. bronchiseptica* are occasionally reported in humans. *B. petrii*, *B. hinzii*, and *B. ansorpii* have been isolated from patients who are immunocompromised.



Bordetella species are gram-negative pleomorphic aerobic bacilli that share common genotypic characteristics. *B. pertussis* and *B. parapertussis* are the most similar of the species, but *B. parapertussis* does not express the gene coding for pertussis toxin. *B. pertussis* is a slow-growing fastidious organism that requires selective medium and forms small, glistening, bifurcated colonies. Suspicious colonies are presumptively identified as *B. pertussis* by direct fluorescent antibody testing or by agglutination with species-specific antiserum. *B. pertussis* is further differentiated from other *Bordetella* species by biochemical and motility characteristics.

B. pertussis produces a wide array of toxins and biologically active products that are important in its pathogenesis and in immunity. Most of these virulence factors are under the control of a single genetic locus that regulates their production, resulting in antigenic modulation and phase variation. Although these processes occur both in vitro and in vivo, their importance in the pathobiology of the organism is unknown; they may play a role in intracellular persistence and person-to-person spread. The organism's most important virulence factor is *pertussis toxin*, which is composed of a B oligomer-binding subunit and an enzymatically active A protomer that ADP-ribosylates a guanine nucleotide-binding regulatory protein (G protein) in target cells, producing a variety of biologic effects. Pertussis toxin has important mitogenic activity, affects the circulation of lymphocytes, and serves as an adhesin for bacterial binding to respiratory ciliated cells. Other important virulence factors and adhesins are *filamentous hemagglutinin*, a component of the cell wall, and *pertactin*, an outer-membrane protein. *Fimbriae*, bacterial appendages that play a role in bacterial attachment, are the major antigens against which agglutinating antibodies are directed. These agglutinating antibodies have historically been the primary means of serotyping *B. pertussis* strains. Other virulence factors include tracheal cytotoxin, which causes respiratory epithelial damage; adenylate cyclase toxin, which impairs host immune-cell function; dermonecrotic toxin, which may contribute to respiratory mucosal damage; and lipooligosaccharide, which has properties similar to those of other gram-negative bacterial endotoxins.

EPIDEMIOLGY

 Pertussis is a highly communicable disease, with attack rates of 80–100% among unimmunized household contacts and 20% within households in well-immunized populations. The infection has a worldwide distribution, with cyclical outbreaks every 3–5 years (a pattern that has persisted despite widespread immunization). Pertussis occurs in all months; however, in North America, its activity peaks in summer and autumn.

In developing countries, pertussis remains an important cause of infant morbidity and death. The reported incidence of pertussis worldwide has decreased as a result of improved vaccine coverage (Fig. 185-1). However, coverage rates are still $< 50\%$ in many developing nations; the World Health Organization (WHO) estimates that 90% of the burden of pertussis is in developing regions. In addition, overreporting of immunization coverage and underreporting of disease result in substantial underestimation of the global burden of pertussis. The WHO estimates that there were 195,000 deaths from pertussis among children in 2008.

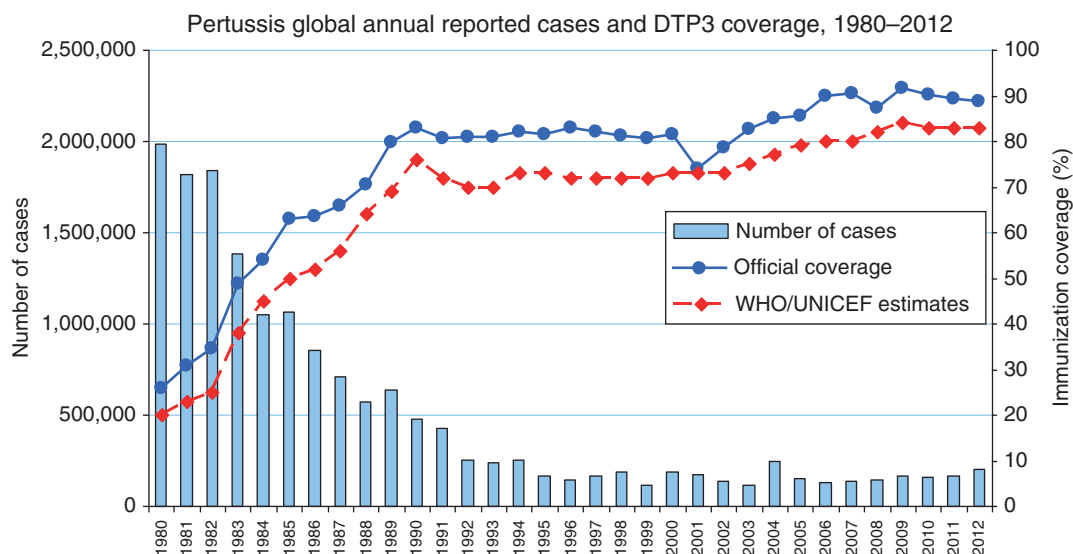


FIGURE 185-1 Global annual reported cases of pertussis and rate of coverage with DTP3 (diphtheria toxoid, tetanus toxoid, and pertussis vaccine; 3 doses), 1980–2012. (© World Health Organization, 2013. All rights reserved. From www.who.int/immunization/monitoring_surveillance/burden/vpd/surveillance_type/passive/Pertussis_coverage.JPG. Source: WHO/IVB database, 2013.)

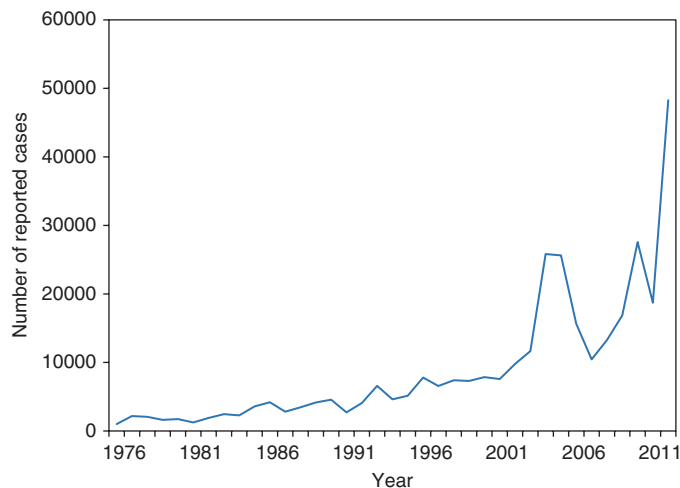


FIGURE 185-2 Reported cases of pertussis by year—United States, 1976–2012. (From the Centers for Disease Control and Prevention, www.cdc.gov/pertussis/surv-reporting/cases-by-year.html. Accessed December 17, 2013.)

Before the institution of widespread immunization programs in the developed world, pertussis was one of the most common infectious causes of morbidity and death. In the United States before the 1940s, between 115,000 and 270,000 cases of pertussis were reported annually, with an average yearly rate of 150 cases per 100,000 population. With universal childhood immunization, the number of reported cases fell by >95%, and mortality rates decreased even more dramatically. Only 1010 cases of pertussis were reported in 1976 (Fig. 185-2). After that historic low, rates of pertussis slowly increased. In recent years, pertussis epidemics have been reported with increasing frequency worldwide. The United States experienced widespread outbreaks of pertussis in 2005, 2010, and 2012 at levels not seen in 40–50 years (>40,000 reported cases in 2012).

Although thought of as a disease of childhood, pertussis can affect people of all ages and is increasingly being identified as a cause of prolonged coughing illness in adolescents and adults. In unimmunized populations, pertussis incidence peaks during the preschool years, and well over half of children have the disease before reaching adulthood. In highly immunized populations such as those in North America, the peak incidence is among infants <1 year of age who have not completed the three-dose primary immunization series. An increase in pertussis incidence among adolescents and adults began in the late 1990s and led to the introduction of an adolescent booster across North America by 2006. While the disease burden among adolescents has started to decrease, children 7–10 years of age have recently emerged as a high-risk group. In major outbreaks in 2010 and 2012, the incidence of pertussis among children 10 years of age, most of whom were fully immunized, was as high as that among infants <6 months of age. Although adults contribute a smaller proportion of reported cases of pertussis than do children and adolescents, this difference may be related to a greater degree of underrecognition and underreporting. A number of studies of prolonged coughing illness suggest that *B. pertussis* may be the etiologic agent in 12–30% of adults with cough that does not improve within 2 weeks. In one study of the efficacy of an acellular pertussis vaccine in adolescents and adults, the incidence of pertussis in the placebo group was 3.7–4.5 cases per 1000 person-years. Although this prospective cohort study yielded a lower estimate than the studies of cough illness, its results still translate to 600,000–800,000 cases of pertussis annually among adults in the United States.

Severe morbidity and high mortality rates, however, are restricted almost entirely to infants. In Canada, there were 16 deaths from pertussis between 1991 and 2001; all those who died were infants ≤6 months of age. Similarly, in the United States between 1993 and 2004, all pertussis deaths and 86% of hospitalizations for pertussis involved infants ≤3 months of age. Although school-age children are the source of infection for most households, adults are the likely source for cases

in high-risk infants and may serve as the reservoir of infection between epidemic years.

PATHOGENESIS

Infection with *B. pertussis* is initiated by attachment of the organism to the ciliated epithelial cells of the nasopharynx. Attachment is mediated by surface adhesins (e.g., pertactin and filamentous hemagglutinin), which bind to the integrin family of cell-surface proteins, probably in conjunction with pertussis toxin. The role of fimbriae in adhesion and in maintenance of infection has not been fully delineated. At the site of attachment, the organism multiplies, producing a variety of other toxins that cause local mucosal damage (tracheal cytotoxin, dermonecrotic toxin). Impairment of host defense by *B. pertussis* is mediated by pertussis toxin and adenylate cyclase toxin. There is local cellular invasion, with intracellular bacterial persistence; however, systemic dissemination does not occur. Systemic manifestations (lymphocytosis) result from the effects of the toxins.

The pathogenesis of the clinical manifestations of pertussis is poorly understood. It is not known what causes the hallmark paroxysmal cough. A pivotal role for pertussis toxin has been proposed. Proponents of this position point to the efficacy of preventing clinical symptoms with a vaccine containing only pertussis toxin. Detractors counter that pertussis toxin is not the critical factor because paroxysmal cough also occurs in patients infected with *B. parapertussis*, which does not produce pertussis toxin. It is thought that neurologic events in pertussis, such as seizures and encephalopathy, are due to hypoxia from coughing paroxysms or apnea rather than to the effects of specific bacterial products. *B. pertussis* pneumonia, which occurs in up to 10% of infants with pertussis, is usually a diffuse bilateral primary infection. In older children and adults with pertussis, pneumonia is often due to secondary bacterial infection with streptococci or staphylococci. Deaths from pertussis among young infants are frequently associated with very high levels of leukocytosis and pulmonary hypertension.

IMMUNITY

Both humoral and cell-mediated immunity are thought to be important in pertussis. Antibodies to pertussis toxin, filamentous hemagglutinin, pertactin, and fimbriae are all protective in animal models. Pertussis agglutinins were correlated with protection in early studies of whole-cell pertussis vaccines. Serologic correlates of protection conferred by acellular pertussis vaccines have not been established, although antibody to pertactin, fimbriae, and (to a lesser degree) pertussis toxin correlated best with protection in two efficacy trials. The duration of immunity after whole-cell pertussis vaccination is short-lived, with little protection remaining after 10–12 years. Recent studies have demonstrated early waning of immunity—i.e., within 2–4 years after the fifth dose of acellular pertussis vaccine in children who received acellular pertussis vaccine for their primary series in infancy. These data suggest that boosters may be needed more frequently than every 10 years, as previously thought. Although immunity after natural infection was thought to be lifelong, seroepidemiologic evidence demonstrates that it clearly is not and that subsequent episodes of clinical pertussis are prevented by intermittent subclinical infection.

CLINICAL MANIFESTATIONS

Pertussis is a prolonged coughing illness with clinical manifestations that vary by age (Table 185-1). Although not uncommon among adolescents and adults, classic pertussis is most often seen in preschool and school-age children. After an incubation period averaging 7–10 days, an illness develops that is indistinguishable from the common cold and is characterized by coryza, lacrimation, mild cough, low-grade fever, and malaise. After 1–2 weeks, this *catarrhal phase* evolves into the *paroxysmal phase*: the cough becomes more frequent and spasmodic with repetitive bursts of 5–10 coughs, often within a single expiration. Posttussive vomiting is frequent, with a mucous plug occasionally expelled at the end of an episode. The episode may be terminated by an audible whoop, which occurs upon rapid inspiration against a closed glottis at the end of a paroxysm. During a spasm, there may be impressive neck-vein distension, bulging eyes, tongue

TABLE 185-1 CLINICAL FEATURES OF PERTUSSIS, BY AGE GROUP AND DIAGNOSTIC STATUS

Feature	Percentage of Patients		
	Adolescents and Adults		Children
	Laboratory Confirmation	No Laboratory Confirmation	
Cough	95–100	95–100	95–100
Prolonged	60–80	60–80	60–95
Paroxysmal	60–90	50–90	80–95
Sleep-disturbing	50–80	50–80	90–100
Whoop	10–40	5–30	40–80
Posttussive vomiting	20–50	5–30	80–90

protrusion, and cyanosis. Paroxysms may be precipitated by noise, eating, or physical contact. Between attacks, the patient's appearance is normal but increasing fatigue is evident. The frequency of paroxysmal episodes varies widely, from several per hour to 5–10 per day. Episodes are often worse at night and interfere with sleep. Weight loss is not uncommon as a result of the illness's interference with eating. Most complications occur during the paroxysmal stage. Fever is uncommon and suggests bacterial superinfection.

After 2–4 weeks, the coughing episodes become less frequent and less severe—changes heralding the onset of the *convalescent phase*. This phase can last 1–3 months and is characterized by gradual resolution of coughing episodes. For 6–12 months, intercurrent viral infections may be associated with a recrudescence of paroxysmal cough.

Not all individuals who develop pertussis have classic disease. The clinical manifestations in adolescents and adults are more often atypical. In a German study of pertussis in adults, more than two-thirds had paroxysmal cough and more than one-third had whoop. Adult illness in North America differs from this experience: the cough may be severe and prolonged but is less frequently paroxysmal, and a whoop is uncommon. Vomiting with cough is the best predictor of pertussis as the cause of prolonged cough in adults. Other predictive features are a cough at night, sweating episodes between paroxysms of coughing, and exposure to other individuals with a prolonged coughing illness.

COMPLICATIONS

Complications are frequently associated with pertussis and are more common among infants than among older children or adults. Subconjunctival hemorrhages, abdominal and inguinal hernias, pneumothoraces, and facial and truncal petechiae can result from increased intrathoracic pressure generated by severe fits of coughing. Weight loss can follow decreased caloric intake. In a series of more than 1100 children <2 years of age who were hospitalized with pertussis, 27.1% had apnea, 9.4% had pneumonia, 2.6% had seizures, and 0.4% had encephalopathy; 10 children (0.9%) died. Pneumonia is reported in <5% of adolescents and adults and increases in frequency after 50 years of age. In contrast to the primary *B. pertussis* pneumonia that develops in infants, pneumonia in adolescents and adults with pertussis is usually caused by a secondary infection with encapsulated organisms such as *Streptococcus pneumoniae* or *Haemophilus influenzae*. Pneumothorax, severe weight loss, inguinal hernia, rib fracture, carotid artery aneurysm, and cough syncope have all been reported in adolescents and adults with pertussis.

DIAGNOSIS

If the classic symptoms of pertussis are present, clinical diagnosis is not difficult. However, particularly in older children and adults, it is difficult to differentiate infections caused by *B. pertussis* and *B. parapertussis* from other respiratory tract infections on clinical grounds. Therefore, laboratory confirmation should be attempted in all cases. Lymphocytosis (an absolute lymphocyte count of $>10^8$ – $10^9/L$) is common among young children, in whom it is unusual with other infections, but not among adolescents and adults. Culture of nasopharyngeal secretions remains the gold standard of diagnosis, although DNA

detection by PCR has replaced culture in many laboratories because of increased sensitivity and quicker results. Appropriate PCR methodology must include primers to differentiate among *B. pertussis*, *B. parapertussis*, and *B. holmesii*. The best specimen is collected by nasopharyngeal aspiration, in which a fine flexible plastic catheter attached to a 10-mL syringe is passed into the nasopharynx and withdrawn while gentle suction is applied. Since *B. pertussis* is highly sensitive to drying, secretions for culture should be inoculated without delay onto appropriate medium (Bordet-Gengou or Regan-Lowe), or the catheter should be flushed with a phosphate-buffered saline solution for culture and/or PCR. An alternative to the aspirate is a Dacron or rayon nasopharyngeal swab; again, inoculation of culture plates should be immediate or an appropriate transport medium (e.g., Regan-Lowe charcoal medium) should be used. Results of PCR can be available within hours; cultures become positive by day 5 of incubation. *B. pertussis* and *B. parapertussis* can be differentiated by agglutination with specific antisera or by direct immunofluorescence.

Nasopharyngeal cultures in untreated pertussis remain positive for a mean of 3 weeks after the onset of illness; these cultures become negative within 5 days of the institution of appropriate antimicrobial therapy. The duration of a positive PCR in untreated pertussis or after therapy is not known but exceeds that of positive cultures. Since much of the period during which the organism can be recovered from the nasopharynx falls into the catarrhal phase, when the etiology of the infection is not suspected, there is only a small window of opportunity for culture-proven diagnosis. Cultures from infants and young children are more frequently positive than those from older children and adults; this difference may reflect earlier presentation of the former age group for medical care. Direct fluorescent antibody tests of nasopharyngeal secretions for direct diagnosis may still be available in some laboratories but should not be used because of poor sensitivity and specificity. Pseudo-outbreaks of pertussis have been reported as a result of false-positive PCR results. Greater standardization of PCR methodology can alleviate this problem.

As a result of the difficulties with laboratory diagnosis of pertussis in adolescents, adults, and patients who have been symptomatic for >4 weeks, increasing attention is being given to serologic diagnosis. Enzyme immunoassays detecting IgA and IgG antibodies to pertussis toxin, filamentous hemagglutinin, pertactin, and fimbriae have been developed and assessed for reproducibility. Two- or fourfold increases in antibody titer are suggestive of pertussis, although cross-reactivity of some antigens (such as filamentous hemagglutinin and pertactin) among *Bordetella* species makes it difficult to depend diagnostically on seroconversion involving a single type of antibody. Late presentation for medical care and prior immunization also complicate serologic diagnosis because the first sample obtained may in fact be a convalescent-phase specimen. Criteria for serologic diagnosis based on comparison of results for a single serum specimen with established population values are gaining acceptance, and serologic measurement of antibody to pertussis toxin is becoming more widely standardized and available for diagnostic purposes, particularly in outbreak settings and for surveillance.

DIFFERENTIAL DIAGNOSIS

A child presenting with paroxysmal cough, posttussive vomiting, and whoop is likely to have an infection caused by *B. pertussis* or *B. parapertussis*; lymphocytosis increases the likelihood of a *B. pertussis* etiology. Viruses such as respiratory syncytial virus and adenovirus have been isolated from patients with clinical pertussis but probably represent co-infection.

In adolescents and adults, who often do not have paroxysmal cough or whoop, the differential diagnosis of a prolonged coughing illness is more extensive. Pertussis should be suspected when any patient has a cough that does not improve within 14 days, a paroxysmal cough of any duration, a cough followed by vomiting (adolescents and adults), or any respiratory symptoms after contact with a laboratory-confirmed case of pertussis. Other etiologies to consider include infections caused by *Mycoplasma pneumoniae*, *Chlamydia pneumoniae*, adenovirus, influenza virus, and other respiratory viruses. Use of angiotensin-converting

TABLE 185-2 ANTIMICROBIAL THERAPY FOR PERTUSSIS

Drug	Adult Daily Dose	Frequency	Duration, Days	Comments
Erythromycin estolate	1–2 g	3 divided doses	7–14	Frequent gastrointestinal side effects
Clarithromycin	500 mg	2 divided doses	7	—
Azithromycin	500 mg on day 1, 250 mg subsequently	1 daily dose	5	—
Trimethoprim-sulfamethoxazole	160 mg of trimethoprim, 800 mg of sulfamethoxazole	2 divided doses	14	For patients allergic to macrolides; data on effectiveness limited

enzyme (ACE) inhibitors, reactive airway disease, and gastroesophageal reflux disease are well-described noninfectious causes of prolonged cough in adults.

TREATMENT PERTUSSIS

ANTIBIOTICS

The purpose of antibiotic therapy for pertussis is to eradicate the infecting bacteria from the nasopharynx; therapy does not substantially alter the clinical course unless given early in the catarrhal phase. Macrolide antibiotics are the drugs of choice for treatment of pertussis (Table 185-2); macrolide-resistant *B. pertussis* strains have been reported but are rare. Trimethoprim-sulfamethoxazole is recommended as an alternative for individuals allergic to macrolides.

SUPPORTIVE CARE

Young infants have the highest rates of complication and death from pertussis; therefore, most infants (and older children with severe disease) should be hospitalized. A quiet environment may decrease the stimulation that can trigger paroxysmal episodes. Use of β -adrenergic agonists and/or glucocorticoids has been advocated by some authorities but has not been proven to be effective. Cough suppressants are not effective and play no role in the management of pertussis.

INFECTION CONTROL MEASURES

Hospitalized patients with pertussis should be placed in respiratory isolation, with the use of precautions appropriate for pathogens spread by large respiratory droplets. Isolation should continue for 5 days after initiation of macrolide therapy or, in untreated patients, for 3 weeks (i.e., until nasopharyngeal cultures are consistently negative).

PREVENTION

Chemoprophylaxis Because the risk of transmission of *B. pertussis* within households is high, chemoprophylaxis is widely recommended for household contacts of pertussis cases. The effectiveness of chemoprophylaxis, although unproven, is supported by several epidemiologic studies of institutional and community outbreaks. In the only randomized placebo-controlled study, erythromycin estolate (50 mg/kg per day in three divided doses; maximum dose, 1 g/d) was effective in reducing the incidence of bacteriologically confirmed pertussis by 67%; however, there was no decrease in the incidence of clinical disease. Despite these disappointing results, many authorities continue to recommend chemoprophylaxis, particularly in households with members at high risk of severe disease (children <1 year of age, pregnant women). Data are not available on use of the newer macrolides for chemoprophylaxis, but these drugs are commonly used because of their increased tolerability and their effectiveness.

Immunization (See also Chap. 148) The mainstay of pertussis prevention is active immunization. Pertussis vaccine became widely used in North America after 1940; the reported number of pertussis cases subsequently fell by >90%. Whole-cell pertussis vaccines are prepared through the heating, chemical inactivation, and purification of whole *B. pertussis* organisms. Despite their efficacy (average estimate, 85%; range for different products, 30–100%), whole-cell pertussis vaccines are associated with adverse events—both common (fever; injection-site pain, erythema, and swelling; irritability) and uncommon (febrile seizures, hypotonic hyporesponsive episodes). Alleged associations of whole-cell pertussis vaccine with encephalopathy, sudden infant death syndrome, and autism, although not substantiated, have spawned an active anti-immunization lobby. The development of acellular pertussis vaccines, which are effective and less reactogenic, has greatly alleviated concerns about the inclusion of pertussis vaccine in the combined infant immunization series.



Although whole-cell vaccines are still used extensively in developing regions of the world, acellular pertussis vaccines are used exclusively for childhood immunization in much of the developed world. In North America, acellular pertussis vaccines for children are given as a three-dose primary series at 2, 4, and 6 months of age, with a reinforcing dose at 15–18 months of age and a booster dose at 4–6 years of age.

Although a wide variety of acellular pertussis vaccines were developed, only a few are still widely marketed; all contain pertussis toxoid and filamentous hemagglutinin. One acellular pertussis vaccine also contains pertactin, and another contains pertactin and two types of fimbriae. In light of phase 3 efficacy studies, most experts have concluded that two-component acellular pertussis vaccines are more effective than monocomponent vaccines and that the addition of pertactin increases efficacy still more. The further addition of fimbriae appears to enhance protective efficacy against milder disease. In two studies, protection conferred by pertussis vaccines correlated best with the production of antibody to pertactin, fimbriae, and pertussis toxin.

Adult formulations of acellular pertussis vaccines have been shown to be safe, immunogenic, and efficacious in clinical trials in adolescents and adults and are now recommended for routine immunization of these groups in several countries, including the United States. In this country, adolescents should receive a dose of the adult-formulation diphtheria–tetanus–acellular pertussis vaccine at the preadolescence physician's visit, and all unvaccinated adults should receive a single dose of this combined vaccine. In addition, in the United States, pertussis immunization is specifically recommended for health care workers and for women during each pregnancy to increase passive transfer of maternal antibodies to the fetus. Pertussis vaccine coverage among U.S. adolescents was 78.2% in 2011, but coverage among adults is low (2.1% as of 2007). Further improvements in adult vaccine coverage may permit better control of pertussis across the age spectrum, with collateral protection of infants too young to be immunized. However, more effective vaccines with longer-lasting protection will ultimately be needed to control this disease.


186 Diseases Caused by Gram-Negative Enteric Bacilli

Thomas A. Russo, James R. Johnson

GENERAL FEATURES AND PRINCIPLES

Escherichia coli, *Klebsiella*, *Proteus*, *Enterobacter*, *Serratia*, *Citrobacter*, *Morganella*, *Providencia*, *Cronobacter*, and *Edwardsiella* are gram-negative enteric bacilli that are members of the family Enterobacteriaceae. *Salmonella*, *Shigella*, and *Yersinia*, also in the family Enterobacteriaceae, are discussed in Chaps. 190, 191, and 196, respectively. These pathogens cause a wide variety of infections involving diverse anatomic sites in both healthy and compromised hosts. Increasing antimicrobial resistance in this group has put them at the forefront of an evolving public health crisis. In addition, new infectious syndromes have emerged. Therefore, a thorough knowledge of clinical presentations and appropriate therapeutic choices is necessary for optimal outcomes.

EPIDEMIOLOGY

 *E. coli*, *Klebsiella*, *Proteus*, *Enterobacter*, *Serratia*, *Citrobacter*, *Morganella*, *Providencia*, *Cronobacter*, and *Edwardsiella* are components of the normal animal and human colonic microbiota and/or the microbiota of a variety of environmental habitats, including long-term-care facilities (LTCFs) and hospitals. As a result, except for certain pathotypes of intestinal pathogenic *E. coli*, these genera are global pathogens. The incidence of infection due to these agents is increasing because of the combination of an aging population and increasing antimicrobial resistance. In healthy humans, *E. coli* is the predominant species of gram-negative bacilli (GNB) in the colonic flora; *Klebsiella* and *Proteus* are less prevalent. GNB (primarily *E. coli*, *Klebsiella*, and *Proteus*) only transiently colonize the oropharynx and skin of healthy individuals. In contrast, in LTCFs and hospital settings, a variety of GNB emerge as the dominant microbiota of both mucosal and skin surfaces, particularly in association with antimicrobial use, severe illness, and extended length of stay. LTCFs are emerging as an important reservoir for resistant GNB. This colonization may lead to subsequent infection; for example, oropharyngeal colonization may lead to pneumonia. Interestingly, the use of ampicillin or amoxicillin was associated with an increased risk of subsequent infection due to the hypervirulent variant of *Klebsiella pneumoniae* in Taiwan; this association suggests that changes in the quantity or prevalence of colonizing bacteria may be important. *Serratia* and *Enterobacter* infection may be acquired through a variety of infusates (e.g., medications, blood products). *Edwardsiella* infections are acquired through freshwater and marine environment exposures and are most common in Southeast Asia.

STRUCTURE AND FUNCTION

Enteric GNB possess an extracytoplasmic outer membrane, which consists of a lipid bilayer with associated proteins, lipoproteins, and

polysaccharides (capsule, lipopolysaccharide). The outer membrane interfaces with the external environment, including the human host. A variety of components of the outer membrane are critical determinants in pathogenesis (e.g., capsule) and antimicrobial resistance (e.g., permeability barrier, efflux pumps).

PATHOGENESIS

Multiple bacterial virulence factors are required for the pathogenesis of infections caused by GNB. Possession of specialized virulence genes defines pathogens and enables them to infect the host efficiently. Hosts and their cognate pathogens have been co-adapting throughout evolutionary history. During the host-pathogen “chess match” over time, various and redundant strategies have emerged in both the pathogens and their hosts (Table 186-1).

Intestinal pathogenic mechanisms are discussed below. The members of the Enterobacteriaceae family that cause extraintestinal infections are primarily extracellular pathogens and therefore share certain pathogenic features. Innate immunity (including the activities of complement, antimicrobial peptides, and professional phagocytes) and humoral immunity are the principal host defense components. Both susceptibility to and severity of infection are increased with dysfunction or deficiencies of these components. By contrast, the virulence traits of intestinal pathogenic *E. coli*—i.e., the distinctive strains that can cause diarrheal disease—are for the most part different from those of extraintestinal pathogenic *E. coli* (ExPEC) and other GNB that cause extraintestinal infections. This distinction reflects site-specific differences in host environments and defense mechanisms.

A given strain usually possesses multiple adhesins for binding to a variety of host cells (e.g., in *E. coli*: type 1, S, and F1C fimbriae; P pili). Nutrient acquisition (e.g., of iron via siderophores) requires many genes that are necessary but not sufficient for pathogenesis. The ability to resist the bactericidal activity of complement and phagocytes in the absence of antibody (e.g., as conferred by capsule or O antigen of lipopolysaccharide) is one of the defining traits of an extracellular pathogen. Tissue damage (e.g., as mediated by hemolysin in the case of *E. coli*) may facilitate spread within the host. Without doubt, many important virulence genes await identification (Chap. 145e).

The ability to induce septic shock is another defining feature of these genera. GNB are the most common causes of this potentially lethal syndrome. Pathogen-associated molecular pattern molecules (PAMPs; e.g., the lipid A moiety of lipopolysaccharide) stimulate a proinflammatory host response via pattern recognition receptors (e.g., Toll-like or C-type lectin receptors) that activate host defense signaling pathways; if overly exuberant, this response results in shock (Chap. 325). Direct bacterial damage of host tissue (e.g., by toxins) or collateral damage from the host response can result in the release of damage-associated molecular pattern molecules (DAMPs; e.g., HMGB1) that can propagate a detrimental proinflammatory host response.

Many antigenic variants (serotypes) exist in most genera of GNB. For example, *E. coli* has more than 150 O-specific antigens and more than 80 capsular antigens. This antigenic variability, which permits immune evasion and allows recurrent infection by different strains of the same species, has impeded vaccine development (Chap. 148).

TABLE 186-1 INTERACTIONS OF EXTRAINTESTINAL PATHOGENIC *ESCHERICHIA COLI* WITH THE HUMAN HOST: A PARADIGM FOR EXTRACELLULAR, EXTRAINTESTINAL GRAM-NEGATIVE BACTERIAL PATHOGENS


Bacterial Goal	Host Obstacle	Bacterial Solution
Extraintestinal attachment	Flow of urine, mucociliary blanket	Multiple adhesins (e.g., type 1, S, and F1C fimbriae; P pili)
Nutrient acquisition for growth	Nutrient sequestration (e.g., iron via intracellular storage and extracellular scavenging via lactoferrin and transferrin)	Cellular lysis (e.g., hemolysin), multiple mechanisms for competing for iron (e.g., siderophores) and other nutrients
Initial avoidance of host bactericidal activity	Complement, phagocytic cells, antimicrobial peptides	Capsular polysaccharide, lipopolysaccharide
Dissemination (within host and between hosts)	?	Irritant tissue damage resulting in increased excretion (e.g., toxins such as hemolysin)
Late avoidance of host bactericidal activity	Acquired immunity (e.g., specific antibodies), treatment with antibiotics	?Cell entry, acquisition of antimicrobial resistance

E. coli can cause either intestinal or extraintestinal infection, depending on the particular pathotype, and *Edwardsiella tarda* can cause both intestinal and extraintestinal infection. *Klebsiella* primarily causes extraintestinal infection, but hemorrhagic colitis has been associated with a toxin-producing variant of *Klebsiella oxytoca*. Depending on both the host and the pathogen, nearly every organ or body cavity can be infected with GNB. *E. coli* and—to a lesser degree—*Klebsiella* account for most extraintestinal infections due to GNB and are the most virulent pathogens within this group; this virulence is demonstrated by the ability of *E. coli* and *Klebsiella pneumoniae* (primarily the hypervirulent variant) to cause severe infections in healthy, ambulatory hosts from the community. However, the other genera are also important, especially among LTCF residents and hospitalized patients, in large part because of the intrinsic or acquired antimicrobial resistance of these organisms and the increasing number of individuals with compromised host defenses. The mortality rate is substantial in many GNB infections and correlates with the severity of illness. Especially problematic are pneumonia and bacteremia (arising from any source), particularly when complicated by organ failure (severe sepsis) and/or shock, for which the associated mortality rates are 20–50%.

DIAGNOSIS

Isolation of GNB from sterile sites almost always implies infection, whereas their isolation from nonsterile sites, particularly from open soft-tissue wounds and the respiratory tract, requires clinical correlation to differentiate colonization from infection. Tentative laboratory identification based on lactose fermentation and indole production (described for each genus below), which usually is possible before final identification of the organism and determination of its antimicrobial susceptibilities, may help to guide empirical antimicrobial therapy.


TREATMENT INFECTIONS CAUSED BY GRAM-NEGATIVE ENTERIC BACILLI

 (See also Chap. 170) Evidence indicates that initiation of appropriate empirical antimicrobial therapy early in the course of GNB infections (particularly serious infections) leads to improved outcomes. The ever-increasing prevalence of multidrug-resistant (MDR) and extensively drug-resistant (XDR) GNB; the lag between published (historical) and current resistance rates; and variations by species, geographic location, regional antimicrobial use, and hospital site (e.g., intensive care units [ICUs] versus wards) necessitate familiarity with evolving patterns of antimicrobial resistance for the selection of appropriate empirical therapy. Factors predictive of isolate resistance include recent antimicrobial use, a health care association (e.g., recent or ongoing hospitalization, dialysis, residence in an LTCF), or international travel (e.g., to Asia, Latin America, Africa, southern Europe). For appropriately selected patients, it may be prudent initially, while susceptibility results are awaited, to use two potentially active agents with the rationale that at least one agent will be active. If broad-spectrum treatment has been initiated, it is critical to switch to the most appropriate narrower-spectrum agent when information on antimicrobial susceptibility becomes available. Such responsible antimicrobial stewardship will slow down the ever-escalating cycle of selection for increasingly resistant bacteria, decrease the likelihood of *Clostridium difficile* infection, decrease costs, and maximize the useful longevity of available antimicrobial agents. Likewise, it is important to avoid treatment of patients who are colonized but not infected (e.g., who have a positive sputum culture without evidence of pneumonia). At present, the most reliably active agents against enteric GNB are the carbapenems (e.g., imipenem), the aminoglycoside amikacin, the fourth-generation cephalosporin cefepime, the β -lactam/ β -lactamase inhibitor combination piperacillin-tazobactam, and the polymyxins (e.g., colistin or polymyxin B). The number of antimicrobials effective against certain Enterobacteriaceae is shrinking. Truly pan-resistant GNB exist, and it is unlikely that new agents will come to market in

the short term. Accordingly, the presently available antimicrobials must be used judiciously.

β -Lactamases, which inactivate β -lactam agents, are the most important mediators of resistance to these drugs in GNB. Decreased permeability and/or active efflux of β -lactam agents, although less common, may occur alone or in combination with β -lactamase-mediated resistance.

Broad-spectrum β -lactamases (e.g., TEM, SHV), which mediate resistance to many penicillins and first-generation cephalosporins, are frequently expressed in enteric GNB. These enzymes are inhibited by β -lactamase inhibitors (e.g., clavulanate, sulbactam, tazobactam). They usually do not hydrolyze third- and fourth-generation cephalosporins or cephamycins (e.g., cefoxitin).

 *Extended-spectrum β -lactamases* (ESBLs; e.g., CTX-M, SHV, TEM) are modified broad-spectrum enzymes that confer resistance to the same drugs as well as to third-generation cephalosporins, aztreonam, and (in some instances) fourth-generation cephalosporins. GNB that express ESBLs may also possess porin mutations that result in decreased uptake of cephalosporins and β -lactam/ β -lactamase inhibitor combinations. The prevalence of acquired ESBL production, particularly of CTX-M-type enzymes, is increasing in GNB worldwide, in large part due to the presence of the responsible genes on large transferable plasmids with linked or associated resistance to fluoroquinolones, trimethoprim-sulfamethoxazole (TMP-SMX), aminoglycosides, and tetracyclines. To date, ESBLs are most prevalent in *K. pneumoniae*, *K. oxytoca*, and *E. coli* but also occur (and are probably underrecognized) in *Enterobacter*, *Citrobacter*, *Proteus*, *Serratia*, and other enteric GNB. At present, the rough regional prevalence of ESBL-producing GNB is India > China > rest of Asia, Latin America, Africa, southern Europe > northern Europe > United States, Canada, and Australia. International travel to high-prevalence regions increases the likelihood of colonization with these strains. ESBL-producing GNB were initially described in hospitals (ICUs > wards) and LTCFs, where outbreaks occurred in association with extensive use of third-generation cephalosporins. However, over the last decade, the incidence of uncomplicated cystitis due to CTX-M ESBL-containing *E. coli* has increased worldwide (including in the United States) among healthy ambulatory women without health care or antimicrobial exposure. Antimicrobial use in food animals has also been implicated in the rise of ESBLs.

The carbapenems are the most reliably active β -lactam agents against ESBL-expressing strains. Clinical experience with alternatives is more limited, but, for organisms susceptible to piperacillin-tazobactam (minimal inhibitory concentration [MIC], ≤ 4 $\mu\text{g}/\text{mL}$), this agent—at a dosage of 4.5 g q6h—may offer a carbapenem-sparing alternative, at least for *E. coli*. The role of tigecycline is unclear despite its excellent in vitro activity; *Proteus*, *Morganella*, and *Providencia* are inherently resistant, and attainable serum and urine levels are low. Therefore, caution appears to be prudent, especially with serious infections, until more clinical data become available. Oral options for the treatment of strains expressing CTX-M ESBLs are limited, with fosfomycin being the most reliably active agent (see section below on the treatment of extraintestinal *E. coli* infections).

AmpC β -lactamases, when induced or stably derepressed to high levels of expression, confer resistance to the same substrates as ESBLs plus the cephamycins (e.g., cefoxitin and cefotetan). The genes encoding these enzymes are primarily chromosomally located and therefore may not exhibit the linked or associated resistance to fluoroquinolones, TMP-SMX, aminoglycosides, and tetracyclines that is common with ESBLs. These enzymes are problematic for the clinician: resistance may develop during therapy with third-generation cephalosporins, resulting in clinical failure, particularly in the setting of bacteremia. Although chromosomal AmpC β -lactamases are present in nearly all members of the Enterobacteriaceae family, the risk of clinically significant induction of high expression levels or selection of stably derepressed mutants with cephalosporin treatment is greatest with *Enterobacter cloacae* and *Enterobacter aerogenes*, lower with *Serratia marcescens* and *Citrobacter freundii*, and lowest with *Providencia* and *Morganella morganii*. In addition, rare

strains of *E. coli*, *K. pneumoniae*, and other Enterobacteriaceae have acquired plasmids containing inducible AmpC β -lactamase genes. Carbapenems are a viable treatment option. The fourth-generation cephalosporin cefepime may be an appropriate option if the concomitant presence of an ESBL can be excluded and source control is achieved. Other carbapenem-sparing alternatives to consider if isolates are susceptible *in vitro* are fluoroquinolones, piperacillin-tazobactam, TMP-SMX, tigecycline, and aminoglycosides, although clinical data are limited.



Carbapenemases (e.g., KPC [class A]; NDM-1, VIM, and IMP [class B]; and OXA-48 [class D]) confer resistance to the same drugs as ESBLs and also to cephamycins and carbapenems. Similar to ESBLs, carbapenemases are usually encoded on large transferable plasmids, which often encode linked resistance to fluoroquinolones, TMP-SMX, tetracyclines, and aminoglycosides. Unfortunately, carbapenemase-producing Enterobacteriaceae are becoming increasingly common, particularly in Asia, and infection with these strains is associated with elevated mortality rates. This reality has prompted the Centers for Disease Control and Prevention (CDC) to categorize carbapenem-resistant Enterobacteriaceae as an “urgent threat” to health care. Carbapenemase production by Enterobacteriaceae is most prevalent in *K. pneumoniae* and *E. coli* but has been described in nearly all members of the family. Automated susceptibility systems may be unreliable for detection of carbapenemases. An elevated MIC or a diminished zone diameter for meropenem or imipenem should prompt genotypic confirmation, if available. Alternatively, the phenotype can be confirmed with a modified Hodge test (which detects classes A, B, and D, although results can be false positive) and/or inhibition tests with boronic acid (class A), EDTA (class B), or dipicolinic acid (class B). Carbapenem resistance may also occur in the absence of carbapenemase production and can be mediated by AmpC β -lactamase and ESBL production coupled with modifications in permeability/efflux.

For treatment of carbapenem-resistant Enterobacteriaceae, tigecycline and colistin are the parenteral agents with the most reliable *in vitro* activity. However, because tigecycline reaches only low serum and urine concentrations, caution is warranted in using it to treat bacteremia and perhaps urinary tract infection (UTI), although a few case reports describe some success with tigecycline therapy for UTI. Colistin has nephrotoxic and neurotoxic potential. Furthermore, increasing resistance has been described to both of these agents. Thus the clinician is left with few or no therapeutic options. Aminoglycosides may have some utility if active. Fosfomycin is often active *in vitro*, but clinical data are limited, concerns exist about the development of resistance with monotherapy, and no parenteral formulation is available in the United States. Although control data are lacking, combination therapy is being used in this setting with the goals of increasing efficacy and decreasing the emergence of resistance.

Resistance to fluoroquinolones usually is due to alterations of the target site (DNA gyrase and/or topoisomerase IV), with or without decreased permeability, active efflux, or protection of the target site. Resistance to this drug class is increasingly prevalent among GNB and is associated with resistance to other antimicrobial classes; for example, 20–80% of ESBL-producing enteric GNB are also resistant to fluoroquinolones. At present, quinolones should be considered unreliable as empirical therapy for infections due to GNB in critically ill patients.

In this era of increasing antimicrobial resistance, it is critical to culture the local site of infection before the initiation of antimicrobial therapy and, for systemically ill patients, to obtain blood samples for culture. Antimicrobial resistance may not always be identified by *in vitro* testing; therefore, it is important to assess the clinical response to treatment. Moreover, as discussed above, resistance may emerge during therapy through the induction or stable derepression of AmpC β -lactamases. In addition, drainage of abscesses, resection of necrotic tissue, and removal of infected foreign bodies are often required for cure. GNB are commonly involved in polymicrobial infections, in which the role of each individual pathogen is uncertain

(Chap. 201). Although some GNB are more pathogenic than others, it is usually prudent, if possible, to design an antimicrobial regimen active against all of the GNB identified, because each is capable of pathogenicity in its own right. Lastly, for patients treated initially with a broad-spectrum empirical regimen, the regimen should be de-escalated as expeditiously as possible once susceptibility results are known and the patient has responded to therapy.

PREVENTION

(See also Chap. 168) Avoidance of inappropriate antimicrobial use is a key measure in preventing infections due to antimicrobial-resistant strains and the further development of antimicrobial resistance. Antimicrobial stewardship programs should be adopted to facilitate achievement of this goal. Diligent adherence to hand-hygiene protocols by health care personnel, cleaning/disinfection of objects that come into contact with patients (e.g., stethoscopes and blood pressure cuffs), and contact precautions should be implemented for patients colonized or infected with carbapenem-resistant (and perhaps other XDR) GNB. Avoidance of the use of indwelling devices (e.g., urinary and intravascular catheters, endotracheal tubes) and, when such devices are necessary, placement according to an appropriate protocol decrease infection risk. Likewise, protocols for daily use evaluation and removal as soon as possible should be implemented. Patient positioning (e.g., head of bed at $\geq 30^\circ$) and good oral hygiene decrease the incidence of pneumonia among ventilated patients. Increasing data support the implementation of universal decolonization to prevent infection in ICU patients.

ESCHERICHIA COLI INFECTIONS



Strains of *E. coli* are united by a core genome of ~2000 genes. A strain's ability to cause infections and the nature of such infections are defined by ancillary genes that encode various virulence factors. This experiment of nature is fluid and ongoing, as demonstrated by the recent evolution of Shiga toxin-producing enteroaggregative *E. coli*.

COMMENSAL STRAINS

For the most part, commensal *E. coli* variants, which constitute the bulk of the normal facultative intestinal flora in most humans, confer benefits to the host (e.g., resistance to colonization with pathogenic organisms). These strains generally lack the specialized virulence traits that enable extraintestinal and intestinal pathogenic *E. coli* strains to cause disease outside and within the gastrointestinal tract, respectively. However, even commensal *E. coli* strains can be involved in extraintestinal infections in the presence of an aggravating factor, such as a foreign body (e.g., a urinary catheter), host compromise (e.g., local anatomic or functional abnormalities, such as urinary or biliary tract obstruction or systemic immunocompromise), or an inoculum that is large or contains a mixture of bacterial species (e.g., fecal contamination of the peritoneal cavity).

EXTRAIESTINAL PATHOGENIC STRAINS



ExPEC strains are the most common enteric GNB to cause community-acquired and health care-associated bacterial infections. The emerging propensity of these strains to acquire new antimicrobial resistance mechanisms (e.g., ESBL and carbapenemase production) has posed challenges in managing ExPEC infection. One clonal group—ST131, the members of which are usually resistant to fluoroquinolones and increasingly express an ESBL (CTX-M)—has undergone global dissemination.

Like commensal *E. coli* (but in contrast to intestinal pathogenic *E. coli*), ExPEC strains are often found in the intestinal flora of healthy individuals and do not cause gastroenteritis in humans. Entry from their site of colonization (e.g., the colon, vagina, or oropharynx) into a normally sterile extraintestinal site (e.g., the urinary tract, peritoneal cavity, or lungs) is the rate-limiting step for infection. ExPEC strains have acquired genes encoding diverse extraintestinal virulence factors that enable the bacteria to cause infections outside the gastrointestinal

1028 tract in both normal and compromised hosts (Table 186-1). These virulence genes define ExPEC and, for the most part, are distinct from those that enable intestinal pathogenic strains to cause diarrheal disease (Table 186-2). All age groups, all types of hosts, and nearly all organs and anatomic sites are susceptible to infection by ExPEC. Even previously healthy hosts can become severely ill or die when infected with ExPEC; however, adverse outcomes are more common among hosts with comorbid illnesses and host defense abnormalities. The diversity and the medical and economic impact of ExPEC infections are evident from consideration of the following specific syndromes.

Extraintestinal Infectious Syndromes • URINARY TRACT INFECTION The urinary tract is the site most frequently infected by ExPEC. An exceedingly common infection among ambulatory patients, UTI accounts for 1% of ambulatory care visits in the United States and is second only to lower respiratory tract infection among infections responsible for hospitalization. UTIs are best considered by clinical syndrome (e.g., uncomplicated cystitis, pyelonephritis, and catheter-associated UTIs) and within the context of specific hosts (e.g., premenopausal women, compromised hosts; Chap. 162). *E. coli* is the single most common pathogen for all UTI syndrome/host group combinations. Each year in the United States, *E. coli* causes 80–90% of an estimated 6–8 million episodes of uncomplicated cystitis in premenopausal women. Furthermore, 20% of women with an initial cystitis episode develop frequent recurrences.

Uncomplicated cystitis, the most common acute UTI syndrome, is characterized by dysuria, urinary frequency, and suprapubic pain. Fever and/or back pain suggests progression to pyelonephritis. Even with appropriate treatment of pyelonephritis, fever may take 5–7 days to resolve completely. Persistently elevated or increasing fever and neutrophil counts should prompt evaluation for intrarenal or perinephric abscess and/or obstruction. Renal parenchymal damage and loss of renal function during pyelonephritis occur primarily with urinary obstruction, which can be preexisting or, rarely, occurs de novo in diabetic patients who develop renal papillary necrosis as a result of kidney infection. Pregnant women are at unusually high risk for developing pyelonephritis, which can adversely affect the outcome of pregnancy. As a result, prenatal screening for and treatment of asymptomatic bacteriuria are standard. Prostatic infection is a potential complication of UTI in men. The diagnosis and treatment of UTI, as detailed in Chap. 162, should be tailored to the individual host, the nature and site of infection, and local patterns of antimicrobial susceptibility.

ABDOMINAL AND PELVIC INFECTION The abdomen/pelvis is the second most common site of extraintestinal infection due to *E. coli*. A wide

variety of clinical syndromes occur in this location, including acute peritonitis secondary to fecal contamination, spontaneous bacterial peritonitis, dialysis-associated peritonitis, diverticulitis, appendicitis, intraperitoneal or visceral abscesses (hepatic, pancreatic, splenic), infected pancreatic pseudocysts, and septic cholangitis and/or cholecystitis. In intraabdominal infections, *E. coli* can be isolated either alone or (as often occurs) in combination with other facultative and/or anaerobic members of the intestinal flora (Chap. 159).

PNEUMONIA *E. coli* is not usually considered a cause of pneumonia (Chap. 153). Indeed, enteric GNB account for only 1–3% of cases of community-acquired pneumonia, in part because these organisms only transiently colonize the oropharynx in a minority of healthy individuals. However, rates of oral colonization with *E. coli* and other GNB increase with severity of illness and antibiotic use. Consequently, GNB are a more common cause of pneumonia among residents of LTCFs and are the most common cause (60–70% of cases) of hospital-acquired pneumonia (Chap. 168), particularly among postoperative and ICU patients (e.g., ventilator-associated pneumonia). Pulmonary infection is usually acquired by small-volume aspiration but occasionally occurs via hematogenous spread, in which case multifocal nodular infiltrates can be seen. Tissue necrosis, probably due to bacterial cytotoxins, is common. Despite significant institutional variation, *E. coli* is generally the third or fourth most commonly isolated GNB in hospital-acquired pneumonia, accounting for 5–8% of episodes in both U.S.-based and Europe-based studies. Regardless of the host, pneumonia due to ExPEC is a serious disease, with high crude and attributable mortality rates (20–60% and 10–20%, respectively).

MENINGITIS (See also Chap. 164) *E. coli* is one of the two leading causes of neonatal meningitis, the other being group B *Streptococcus*. Most *E. coli* strains that cause neonatal meningitis possess the K1 capsular antigen and derive from a limited number of meningitis-associated clonal groups. Ventriculomegaly commonly occurs. After the first month of life, *E. coli* meningitis is uncommon, occurring predominantly in the setting of surgical or traumatic disruption of the meninges or in the presence of cirrhosis. In patients with cirrhosis who develop meningitis, the meninges are presumably seeded as a result of poor hepatic clearance of portal vein bacteremia.

CELLULITIS/MUSCULOSKELETAL INFECTION *E. coli* contributes frequently to infections of decubitus ulcers and occasionally to infections of ulcers and wounds of the lower extremity in diabetic patients and other hosts with neurovascular compromise. Osteomyelitis secondary to contiguous spread can occur in these settings. *E. coli* also causes cellulitis or infections of burn sites and surgical wounds (accounting for ~10% of surgical site infections), particularly when the infection originates

TABLE 186-2 INTestinal Pathogenic *E. coli*

Pathotype	Epidemiology	Clinical Syndrome ^a	Defining Molecular Trait	Responsible Genetic Element ^b
STEC/EHEC/STEAEC	Food, water, person-to-person; all ages, industrialized countries	Hemorrhagic colitis, hemolytic-uremic syndrome	Shiga toxin	Lambda-like Stx1- or Stx2-encoding bacteriophage
ETEC	Food, water; young children in and travelers to developing countries	Traveler's diarrhea	Heat-stable and labile enterotoxins, colonization factors	Virulence plasmid(s)
EPEC	Person-to-person; young children and neonates in developing countries	Watery diarrhea, persistent diarrhea	Localized adherence, attaching and effacing lesion on intestinal epithelium	EPEC adherence factor plasmid pathogenicity island (locus for enterocyte effacement [LEE])
EIEC	Food, water; children in and travelers to developing countries	Dysentery	Invasion of colonic epithelial cells, intracellular multiplication, cell-to-cell spread	Multiple genes contained primarily in a large virulence plasmid
EAEC	?Food, water; children in and travelers to developing countries; all ages, industrialized countries	Traveler's diarrhea, acute diarrhea, persistent diarrhea	Aggregative/diffuse adherence, virulence factors regulated by AggR	Chromosomal or plasmid-associated adherence and toxin genes

Abbreviations: EAEC, enteroaggregative *E. coli*; EHEC, enterohemorrhagic *E. coli*; EIEC, enteroinvasive *E. coli*; EPEC, enteropathogenic *E. coli*; ETEC, enterotoxigenic *E. coli*; STEAEC, Shiga toxin-producing enteroaggregative *E. coli*; STEC, Shiga toxin-producing *E. coli*. ^a Classic syndromes; see text for details on disease spectrum. ^b Pathogenesis involves multiple genes, including genes in addition to those listed.

close to the perineum. Hematogenously acquired osteomyelitis, especially of vertebral bodies, is more commonly caused by *E. coli* than is generally appreciated; this organism accounts for up to 10% of cases in some series (Chap. 158). *E. coli* occasionally causes orthopedic device-associated infection or septic arthritis and rarely causes hematogenous myositis. Upper-leg myositis or fasciitis due to *E. coli* should prompt an evaluation for an abdominal source with contiguous spread.

ENDOVASCULAR INFECTION Despite being one of the most common causes of bacteremia, *E. coli* rarely seeds native heart valves. When the organism does seed native valves, it usually does so in the setting of prior valvular disease. *E. coli* infections of aneurysms, the portal vein (*pylphlebitis*), and vascular grafts are quite uncommon.

MISCELLANEOUS INFECTIONS *E. coli* can cause infection in nearly every organ and anatomic site. It occasionally causes postoperative mediastinitis or complicated sinusitis and uncommonly causes endophthalmitis, ecthyma gangrenosum, or brain abscess.

BACTEREMIA *E. coli* bacteremia can arise from primary infection at any extraintestinal site. In addition, primary *E. coli* bacteremia can arise from percutaneous intravascular devices or transrectal prostate biopsy or from the increased intestinal mucosal permeability seen in neonates and in the settings of neutropenia and chemotherapy-induced mucositis, trauma, and burns. Roughly equal proportions of *E. coli* bacteremia cases originate in the community and in health care settings. In most studies, *E. coli* and *Staphylococcus aureus* are the two most common blood isolates of clinical significance. Isolation of *E. coli* from the blood is almost always clinically significant and is typically accompanied by the sepsis syndrome, severe sepsis (sepsis-induced dysfunction of at least one organ or system), or septic shock (Chap. 325).

The urinary tract is the most common source of *E. coli* bacteremia, accounting for one-half to two-thirds of episodes. Bacteremia from a urinary tract source is particularly common among patients with pyelonephritis, urinary tract obstruction, or urinary instrumentation in the presence of infected urine. The abdomen is the second most common source, accounting for 25% of episodes. Although biliary obstruction (stones, tumor) and overt bowel disruption, which typically are readily apparent, are responsible for many of these cases, some abdominal sources (e.g., abscesses) are remarkably silent clinically and require identification via imaging studies (e.g., CT). Therefore, the physician should be cautious in designating the urinary tract as the source of *E. coli* bacteremia in the absence of characteristic signs and symptoms of UTI. Soft tissue, bone, pulmonary, and intravascular catheter infections are other sources of *E. coli* bacteremia.

Diagnosis Strains of *E. coli* that cause extraintestinal infections usually grow both aerobically and anaerobically within 24 h on standard diagnostic media and are easily identified by the clinical microbiology laboratory according to routine biochemical criteria. More than 90% of ExPEC strains are rapid lactose fermenters and are indole positive.

TREATMENT EXTRAINTESTINAL *E. COLI* INFECTIONS

In the past, most *E. coli* isolates were highly susceptible to a broad range of antimicrobial agents. Unfortunately, this situation has changed. In general, the high prevalence of resistance precludes empirical use of ampicillin and amoxicillin-clavulanate, even for community-acquired infections. The prevalence of resistance to first-generation cephalosporins and TMP-SMX is increasing among community-acquired strains in the United States (with current rates of 10–40%) and is even higher outside North America. Until recently, TMP-SMX was the drug of choice for the treatment of uncomplicated cystitis in many locales. Although continued empirical use of TMP-SMX will predictably result in ever-diminishing cure rates, a wholesale switch to alternative agents (e.g., fluoroquinolones) will just as predictably accelerate the widespread emergence of resistance to these antimicrobial classes, as has already occurred in some areas. More than 90% of isolates that cause uncomplicated cystitis remain susceptible to nitrofurantoin and fosfomicin.

The prevalence of resistance to fluoroquinolones among *E. coli* isolates from U.S. outpatients has increased steadily over the last decade (i.e., from 3% in 2000 to 17.1% in 2010, according to one survey). Resistance rates are generally higher in the ambulatory setting outside the United States and are even higher in populations for which fluoroquinolone prophylaxis is used extensively (e.g., patients with leukemia, transplant recipients, and patients with cirrhosis) and among isolates from LTCFs and hospitals. For example, the National Healthcare Safety Network (NHSN) reported fluoroquinolone resistance in 41.8% of central line-associated bloodstream infection (CLABSI) *E. coli* isolates in 2009–2010, and the International Nosocomial Infection Control Consortium (INICC) reported that 53.4% of ICU *E. coli* isolates were resistant to quinolones in 2004–2009. Furthermore, the NHSN reported 19% resistance to third- and fourth-generation cephalosporins in CLABSI *E. coli* isolates, and the INICC found that 66.6% of ICU *E. coli* isolates were resistant to third-generation cephalosporins.

ESBL-producing strains are increasingly prevalent among both health care-associated (5–10%) and ambulatory isolates (region-dependent figures). An increasing number of reports describe community-acquired UTIs caused by *E. coli* strains that produce CTX-M ESBLs. Data suggest that acquisition of CTX-M-producing, fluoroquinolone-resistant strains may result from consumption of meat products from food animals treated with third- and fourth-generation cephalosporins and fluoroquinolones. Oral treatment options for such strains are limited; however, in vitro and limited clinical data indicate that, for cystitis, fosfomicin and nitrofurantoin appear to be useful options. Carbapenems and amikacin are the most predictably active agents overall, but carbapenemase-producing strains are on the rise (1–5% among health care-associated isolates in the United States and higher rates in many other countries). Tigecycline and the polymyxins, with or without a second agent, have been used most frequently against these extremely resistant isolates.

This evolving antimicrobial resistance—a source of serious concern—necessitates not only the increasing use of broad-spectrum agents but also the use of the most appropriate narrower-spectrum agent whenever possible and the avoidance of treatment of colonized but uninfected patients.

INTESTINAL PATHOGENIC STRAINS

Pathotypes Certain strains of *E. coli* are capable of causing diarrheal disease. Other important intestinal pathogens are discussed in Chaps. 160, 161, and 190–193. At least in the industrialized world, intestinal pathogenic strains of *E. coli* are rarely encountered in the fecal flora of healthy persons and instead appear to be essentially obligate pathogens. These strains have evolved a special ability to cause enteritis, enterocolitis, and colitis when ingested in sufficient quantities by a naive host. At least five distinct pathotypes of intestinal pathogenic *E. coli* exist: (1) Shiga toxin-producing *E. coli* (STEC), which includes the subsets of enterohemorrhagic *E. coli* (EHEC) and the recently evolved Shiga toxin-producing enteroaggregative *E. coli* (STEAEC); (2) enterotoxigenic *E. coli* (ETEC); (3) enteropathogenic *E. coli* (EPEC); (4) enteroinvasive *E. coli* (EIEC); and (5) enteroaggregative *E. coli* (EAEC). Diffusely adherent *E. coli* (DAEC) and cytotodetaching *E. coli* are additional putative pathotypes. Lastly, a variant termed adherent invasive *E. coli* (AIEC) has been associated with Crohn's disease (although a causal role remains unproven) but does not cause acute diarrheal disease. Transmission occurs predominantly via contaminated food and water for ETEC, STEC/EHEC/STEAEC, EIEC, and EAEC and by person-to-person spread for EPEC (and occasionally STEC/EHEC/STEAEC). Gastric acidity confers some protection against infection; therefore, persons with decreased stomach acid levels are especially susceptible. Humans are the major reservoir (except for STEC/EHEC, with regard to which bovines are the main concern); host range appears to be dictated by species-specific attachment factors. Although there is some overlap, each pathotype possesses a largely unique combination of virulence traits that results in a distinctive intestinal pathogenic mechanism (Table 186-2). These strains are largely incapable of causing disease outside the


1030 intestinal tract. Except in the cases of STEC/EHEC/STEAEC and EAEC, disease due to this group of pathogens occurs primarily in developing countries.

ENTEROHEMORRHAGIC *E. COLI*/SHIGA TOXIN–PRODUCING ENTEROAGGREGATIVE *E. COLI* STEC/EHEC/STEAEC strains constitute an emerging group of pathogens that can cause hemorrhagic colitis and the hemolytic-uremic syndrome (HUS). Several large outbreaks resulting from the consumption of fresh produce (e.g., lettuce, spinach, sprouts) and of undercooked ground beef have received significant attention in the media. An outbreak in central Europe in 2011 due to STEAEC (O104:H4) that was probably transmitted by sprouts, with some subsequent human-to-human transmission, resulted in more than 800 cases of HUS and 54 deaths. Within this group of organisms, O157:H7 is the most prominent serotype, but many other serotypes have also been associated with these syndromes, including O6, O26, O45, O55, O91, O103, O111, O113, O121, O145, and OX3.


The ability of STEC/EHEC/STEAEC to produce Shiga toxin (Stx2 and/or Stx1) or related toxins is a critical factor in the occurrence of clinical disease. *Shigella dysenteriae* strains that produce the closely related Shiga toxin Stx can cause the same syndrome. Stx2 and its Stx2C variant (which may be variably present in combination with Stx2 and/or Stx1) appear to be more important than Stx1 in the development of HUS. All Shiga toxins studied to date are multimers comprising one enzymatically active A subunit and five identical B subunits that mediate binding to globosyl ceramides, which are membrane-associated glycolipids expressed on certain host cells. As in ricin, the A subunit cleaves an adenine from the host cell's 28S rRNA, thereby irreversibly inhibiting ribosomal function and potentially leading to apoptosis. Stx2-mediated activation of complement may also play a role in the development of HUS.

Additional properties, such as acid tolerance and epithelial cell adherence, are necessary for full pathogenicity among STEC strains. Most disease-causing isolates possess the chromosomal locus for enterocyte effacement (LEE). This pathogenicity island was first described in EPEC strains and contains genes that mediate adherence to intestinal epithelial cells and a system that subverts host cells by the translocation of bacterial proteins (type III secretion system). EHEC strains make up the subgroup of STEC strains that possess *stx*₁ and/or *stx*₂ as well as LEE. STEAEC (LEE-negative) evolved from EAEC via the acquisition of a number of genes, including those that encode Stx2, the Iha adhesin, tellurite resistance, a type VI secretion system, and the CTX-M-15 ESBL.


Domesticated ruminant animals, particularly cattle and young calves, serve as the major reservoir for STEC/EHEC. Ground beef—the most common food source of STEC/EHEC strains—is often contaminated during processing. Furthermore, manure from cattle or other animals (including that in the form of fertilizer) can contaminate produce (potatoes, lettuce, spinach, sprouts, fallen fruits, nuts, strawberries), and fecal runoff from this source can contaminate water systems. Dairy products and petting zoos are additional sources of infection. By contrast, humans appear to be the reservoir for STEAEC. It is estimated that <10² colony-forming units (CFU) of STEC/EHEC/STEAEC can cause disease. Therefore, not only can low levels of food or environmental contamination (e.g., in water swallowed while swimming) result in disease, but person-to-person transmission (e.g., at day-care centers and in institutions) is an important route for secondary spread. Laboratory-associated infections also occur. Illness due to this group of pathogens occurs both as outbreaks and as sporadic cases, with a peak incidence in the summer months.

 In contrast to other intestinal pathotypes, STEC/EHEC/STEAEC causes infections more frequently in industrialized countries than in developing regions. O157:H7 strains are the fourth most commonly reported cause of bacterial diarrhea in the United States (after *Campylobacter*, *Salmonella*, and *Shigella*). Colonization of the colon and perhaps the ileum results in symptoms after an incubation period of 3 or 4 days. Colonic edema and an initial nonbloody secretory diarrhea may develop into the STEC/EHEC/STEAEC hallmark syndrome of grossly bloody diarrhea (identified by

history or examination) in >90% of cases. Significant abdominal pain and fecal leukocytes are common (70% of cases), whereas fever is not; absence of fever can incorrectly lead to consideration of noninfectious conditions (e.g., intussusception and inflammatory or ischemic bowel disease). Occasionally, infections caused by *C. difficile*, *K. oxytoca* (see “*Klebsiella* Infections,” below), *Campylobacter*, and *Salmonella* present in a similar fashion. STEC/EHEC disease is usually self-limited, lasting 5–10 days. An uncommon but feared complication of this infection is HUS, which occurs 2–14 days after diarrhea in 2–8% of cases, most often affecting very young or elderly patients. Distinctive features of STEAEC infection, as compared with classical STEC/EHEC disease, include a higher incidence among adults, especially young women, and a higher rate of HUS (~20%). It is estimated that >50% of all cases of HUS in the United States and 90% of HUS cases in children are caused by STEC/EHEC. This complication is mediated by the systemic translocation of Shiga toxins. Erythrocytes may serve as carriers of Stx to endothelial cells located in the small vessels of the kidney and brain. The subsequent development of thrombotic microangiopathy (perhaps with direct toxin-mediated effects on various nonendothelial cells) commonly produces some combination of fever, thrombocytopenia, renal failure, and encephalopathy. Although the mortality rate with dialysis support is <10%, residual renal and neurologic dysfunction may persist.

 **ENTEROTOXIGENIC *E. COLI*** In tropical or developing countries, ETEC is a major cause of endemic diarrhea. After weaning, children in these locales commonly experience several episodes of ETEC infection during the first 3 years of life. The incidence of disease diminishes with age, a pattern that correlates with the development of mucosal immunity to colonization factors (i.e., adhesins). In industrialized countries, infection usually follows travel to endemic areas, although occasional food-borne outbreaks occur. ETEC is the most common agent of traveler's diarrhea, causing 25–75% of cases. The incidence of infection may be decreased by prudent avoidance of potentially contaminated fluids and foods, particularly items that are poorly cooked, unpeeled, or unrefrigerated (Chap. 149). ETEC infection is uncommon in the United States, but outbreaks secondary to consumption of food products imported from endemic areas have occurred. A large inoculum (10⁶–10¹⁰ CFU) is needed to produce disease, which usually develops after an incubation period of 12–72 h.

After adherence of ETEC via colonization factors (e.g., CFA/I, CS1-6), disease is mediated primarily by a heat-labile toxin (LT-1) and/or a heat-stable toxin (STa) that causes net fluid secretion via activation of adenylate cyclase (LT-1) and/or guanylate cyclase (STa) in the jejunum and ileum. The result is watery diarrhea accompanied by cramps. LT-1 consists of an A and a B subunit and is structurally and functionally similar to cholera toxin. Strong binding of the B subunit to the GM₁ ganglioside on intestinal epithelial cells leads to the intracellular translocation of the A subunit, which functions as an ADP-ribosyltransferase. Mature STa is an 18- or 19-amino-acid secreted peptide whose biologic activity is mediated by binding to the guanylate cyclase C found in the brush-border membrane of enterocytes and results in increased intracellular concentrations of cyclic GMP. Characteristically absent in ETEC-mediated disease are histopathologic changes within the small bowel; mucus, blood, and inflammatory cells in stool; and fever. The disease spectrum ranges from a mild illness to a life-threatening cholera-like syndrome. Although symptoms are usually self-limited (typically lasting for 3 days), infection may result in significant morbidity and mortality (mostly from profound volume depletion) when access to health care or suitable rehydration fluids is limited and when small and/or undernourished children are affected.

 **ENTEROPATHOGENIC *E. COLI*** EPEC causes disease primarily in young children, including neonates. The first *E. coli* pathotype recognized as an agent of diarrheal disease, EPEC was responsible for outbreaks of infantile diarrhea (including some outbreaks in hospital nurseries) in industrialized countries in the 1940s and 1950s. At present, EPEC infection is an uncommon cause of diarrhea in developed countries but is an important cause of diarrhea (both sporadic and epidemic) among infants in developing countries. Breast-feeding diminishes the incidence of EPEC infection. Rapid person-to-person spread

may occur. Upon colonization of the small bowel, symptoms develop after a brief incubation period (1 or 2 days). Initial localized adherence via bundle-forming pili leads to a characteristic effacement of microvilli, with the formation of cuplike, actin-rich pedestals mediated by factors in the LEE. Diarrhea production is a complex and regulated process in which host cell modulation by a type III secretion system plays an important role. Strains lacking bundle-forming pili have been categorized as atypical EPEC (aEPEC); increasing data support a role for these strains as intestinal pathogens. Diarrheal stool often contains mucus but not blood. Although EPEC diarrhea is usually self-limited (lasting 5–15 days), it may persist for weeks.



ENTEROINVASIVE *E. COLI* EIEC, a relatively uncommon cause of diarrhea, is rarely identified in the United States, although a few food-related outbreaks have been described. In developing countries, sporadic disease is infrequently recognized in children and travelers. EIEC shares many genetic and clinical features with *Shigella*, both of which evolved from a common ancestor. However, unlike *Shigella*, EIEC produces disease only with a large inoculum (10^8 – 10^{10} CFU), with onset generally following an incubation period of 1–3 days. Initially, enterotoxins are believed to induce secretory small-bowel diarrhea. Subsequently, colonization and invasion of the colonic mucosa, followed by replication therein and cell-to-cell spread, result in the development of inflammatory colitis characterized by fever, abdominal pain, tenesmus, and scant stool containing mucus, blood, and inflammatory cells. Symptoms are usually self-limited (7–10 days).



ENTEROAGGREGATIVE AND DIFFUSELY ADHERENT *E. COLI* EAEC has been described primarily in developing countries and in young children. However, recent studies indicate that it may be a relatively common cause of diarrhea in all age groups in industrialized countries. EAEC has also been recognized increasingly as an important cause of traveler's diarrhea. It is highly adapted to humans, the probable reservoir. A large inoculum is required for infection, which usually manifests as watery and sometimes persistent diarrhea in healthy, malnourished, and HIV-infected hosts. In vitro, the organisms exhibit a diffuse or "stacked-brick" pattern of adherence to small-intestine epithelial cells. Virulence factors that probably are necessary for disease are regulated in part by the transcriptional activator AggR and include the aggregative adherence fimbriae (AAF/I-III); the Hda adhesin; the mucinase Pic; the enterotoxins Pet, EAST-1, ShET1, and HlyE; and dispersin, an antiaggregation protein that promotes mucosal spread. Some strains of DAEC are capable of causing diarrheal disease, primarily in children 2–6 years of age in some developing countries, and may perhaps cause traveler's diarrhea. The Afa/Dr adhesins may contribute to the pathogenesis of such infections.

Diagnosis A practical approach to the evaluation of diarrhea is to distinguish noninflammatory from inflammatory cases; the latter is suggested by grossly bloody or mucoid stool or a positive test for fecal leukocytes (Chap. 160). ETEC, EPEC, and DAEC cause noninflammatory diarrhea and are uncommon in the United States; in this country, the incidence of EAEC infection, which also causes noninflammatory diarrhea, may be underrecognized. The diagnosis of these infections requires specialized assays (e.g., polymerase chain reaction–based tests for pathotype-specific genes) that are not routinely available and are rarely needed because the diseases are self-limited. ETEC causes the majority and EAEC a minority of cases of noninflammatory traveler's diarrhea. Definitive diagnosis generally is not necessary. Empirical antimicrobial (or symptom-based) treatment, along with rehydration therapy, is a reasonable approach. If diarrhea persists for >10 days despite treatment, *Giardia* or *Cryptosporidium* (or, in immunocompromised hosts, certain other microbial agents) should be sought. The diagnosis of infection with EIEC, a rare cause of inflammatory diarrhea in the United States, also requires specialized assays. The CDC now recommends that all patients with community-acquired diarrhea, whether inflammatory or not, be evaluated for STEC/EHEC/STEAEC infection by simultaneous culture (which is important for outbreak detection and control) and assay for the detection of Shiga toxin or its associated genes. The reasons for this recommendation are that bloody

stool is not always present and detection of fecal white blood cells is not optimally sensitive for the diagnosis of STEC/EHEC/STEAEC infection. The use of both tests increases the rate of identification of infection over rates obtained with either test alone. O157 STEC/EHEC may be identified via culture by screening for *E. coli* strains that do not ferment sorbitol, with subsequent serotyping and testing for Shiga toxin. Selective or screening media are not available for the culture of non-O157 strains. Detection of Shiga toxins or toxin genes via DNA-based, enzyme-linked immunosorbent, and cytotoxicity assays offers the advantages of rapidity plus detection of non-O157 STEC/EHEC/STEAEC strains. Specimens positive for toxin but culture-negative for O157 should be forwarded to the local or state public health laboratory.

TREATMENT INTestinal *E. COLI* INFECTIONS

(See also Chap. 128) The mainstay of treatment for all diarrheal syndromes is replacement of water and electrolytes. This measure is especially important for STEC/EHEC/STEAEC infection because appropriate volume expansion may decrease renal damage and improve outcome. The use of prophylactic antibiotics to prevent traveler's diarrhea generally should be discouraged, especially in light of high rates of antimicrobial resistance. However, in selected patients (e.g., those who cannot afford a brief illness or are predisposed to infection), the use of rifaximin, which is nonabsorbable and is well tolerated, is reasonable. When stools are free of mucus and blood, early patient-initiated treatment of traveler's diarrhea with a fluoroquinolone or azithromycin decreases the duration of illness, and the use of loperamide may halt symptoms within a few hours. Although dysentery caused by EIEC is self-limited, treatment hastens the resolution of symptoms, particularly in severe cases. In contrast, antimicrobial therapy for STEC/EHEC/STEAEC infection (the presence of which is suggested by grossly bloody diarrhea without fever) should be avoided because antibiotics may increase the incidence of HUS (possibly via increased production/release of Stx). The role of plasmapheresis and inhibition of C5 (eculizumab) in the treatment of HUS is unresolved.

KLEBSIELLA INFECTIONS



K. pneumoniae is the most important *Klebsiella* species from a medical standpoint, causing community-acquired, LTCF-acquired, and nosocomial infections. *K. oxytoca* is primarily a pathogen in LTCF and hospital settings. *Klebsiella* species are broadly prevalent in the environment and colonize mucosal surfaces of mammals. In healthy humans, the prevalence of *K. pneumoniae* colonization is 5–35% in the colon and 1–5% in the oropharynx; the skin is usually colonized only transiently. Person-to-person spread is the predominant mode of acquisition. Most *Klebsiella* infections in Western countries are caused by "classic" *K. pneumoniae* (cKP) and occur in hospitals and LTCFs. The most common clinical syndromes due to cKP are pneumonia, UTI, abdominal infection, intravascular device infection, surgical site infection, soft tissue infection, and subsequent bacteremia. cKP strains have gained notoriety because their propensity for acquiring antimicrobial resistance determinants makes treatment challenging. Clonal group ST258, many members of which produce the KPC carbapenemase, is undergoing international dissemination. The spread of NDM-1 carbapenemase-producing strains from India in association with medical tourism has captured the attention of physicians and the lay press.

cKP strains appear to be phenotypically and clinically distinct from hypervirulent *K. pneumoniae* (hvKP), an emerging variant that was first recognized in Taiwan in 1986. Although hvKP infections have occurred globally in all ethnic groups, the majority have been reported in the Asian Pacific Rim. This concentration of cases raises the question of whether a geo-specific distribution of the organism or increased susceptibility of Asian hosts is responsible. In contrast to the usual health care-associated venue for cKP infections in the West, hvKP causes serious life- and organ-threatening infections in younger, healthy individuals from the community and can spread metastatically

1032 from primary sites of infection. hvKP infection initially was characterized and distinguished from traditional infections due to cKP by (1) presentation as community-acquired pyogenic liver abscess (**Fig. 186-1, top**), (2) occurrence in patients lacking a history of hepatobiliary disease, and (3) a propensity for metastatic spread to distant sites (e.g., eyes, central nervous system, lungs), which occurred in 11–80% of cases. More recently, this variant has been recognized as the cause of a variety of serious community-acquired extrahepatic abscesses/infections in the absence of liver involvement, including pneumonia, meningitis, endophthalmitis (**Fig. 186-1, middle**), splenic abscess, and necrotizing fasciitis. The affected individuals often have diabetes mellitus and are of Asian ethnicity; however, nondiabetics and all ethnic groups can be affected. Survivors often suffer catastrophic morbidity, such as loss of vision and neurologic sequelae.

K. pneumoniae subspecies *rhinoscleromatis* is the causative agent of rhinoscleroma, a granulomatous mucosal upper respiratory infection that progresses slowly (over months or years) and causes necrosis and occasionally obstruction of the nasal passages. *K. pneumoniae* subspecies *ozaenae* has been implicated as a cause of chronic atrophic rhinitis and rarely of invasive disease in compromised hosts. These two *K. pneumoniae* subspecies are usually isolated from patients in tropical climates and are genomically distinct from both cKP and hvKP.

INFECTIOUS SYNDROMES



Pneumonia Although cKP accounts for only a small proportion of cases of community-acquired pneumonia in Western countries (**Chap. 153**), cKP and *K. oxytoca* are common causes of pneumonia among LTCF residents and hospitalized patients because of increased rates of oropharyngeal colonization. Mechanical ventilation is an important risk factor. In Asia and South Africa, community-acquired pneumonia due to hvKP is becoming increasingly common and often occurs in younger patients with no underlying disease. *Klebsiella* is also a common cause of pneumonia in severely malnourished children in developing countries.

As in all pneumonias due to enteric GNB, production of purulent sputum and evidence of airspace disease are typical. Presentation with earlier, less extensive infection is now more common than that with the classically described lobar infiltrate and bulging fissure. Pulmonary infection due to hvKP that has spread metastatically (e.g., from a hepatic abscess) usually includes nodular bilateral densities, more commonly in the lower lobes. Pulmonary necrosis, pleural effusion, and empyema can occur with disease progression.

UTI cKP accounts for only 1–2% of UTI episodes among otherwise healthy adults but for 5–17% of episodes of complicated UTI, including infections associated with indwelling urinary catheters. UTI due to hvKP presents more commonly as renal or prostatic abscess due to bacteremic spread than as ascending infection.



Abdominal Infection cKP causes a spectrum of abdominal infections similar to that caused by *E. coli* but is less frequently isolated from these infections. hvKP is a common cause of monomicrobial community-acquired pyogenic liver abscess and in the Asian Pacific Rim has been recovered with steadily increasing frequency over the past two decades, replacing *E. coli* as the most common pathogen causing this syndrome. hvKP is increasingly described as a cause of spontaneous bacterial peritonitis and splenic abscess.



Other Infections cKP- and *K. oxytoca*-mediated cellulitis or soft tissue infection most frequently affects devitalized tissue (e.g., decubitus and diabetic ulcers, burn sites) and immunocompromised hosts. cKP and *K. oxytoca* cause some cases of surgical site infection and nosocomial sinusitis in addition to occasional cases of osteomyelitis contiguous to soft tissue infection, nontropical myositis, and meningitis (both during the neonatal period and after neurosurgery). By contrast, hvKP has become an important cause of community-acquired

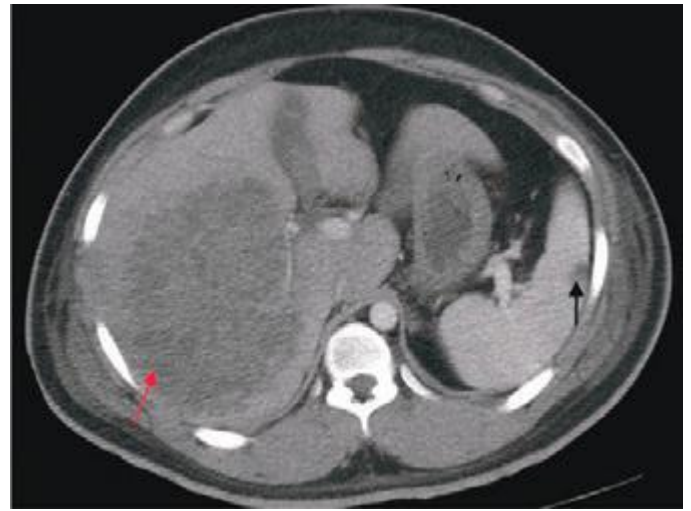


FIGURE 186-1 New hypervirulent variant of *K. pneumoniae* (hvKP). **Top:** Abdominal CT scan of a previously healthy 24-year-old Vietnamese man shows a primary liver abscess (red arrow) with metastatic spread to the spleen (black arrow). (Courtesy of Drs. Chiu-Bin Hsaio and Diana Pomakova.) **Middle:** A previously healthy 33-year-old Chinese man presented with endophthalmitis. (From *Virulence* 4:2, 1-12 Feb. 15, 2013.) **Bottom:** A hypermucoviscous phenotype (which does not necessarily equate with a mucoid phenotype) has been associated with hvKP strains. This phenotype has been semiquantitatively defined by a positive “string test” (formation of a viscous string >5 mm long when bacterial colonies on an agar plate are stretched by an inoculation loop). (Courtesy of Dr. Russo.)

monomicrobial necrotizing fasciitis; meningitis; brain, subdural, and epidural abscess; and endophthalmitis (Fig. 186-1, *middle*), particularly in the Asian Pacific Rim but also globally. Cytotoxin-producing strains of *K. oxytoca* have been implicated as a cause of hemorrhagic antibiotic-associated non-*C. difficile* colitis.

Bacteremia *Klebsiella* infection at any site can produce bacteremia. Infections of the urinary tract, respiratory tract, and abdomen (especially hepatic abscess) each account for 15–30% of episodes of *Klebsiella* bacteremia. Intravascular device-related infections account for another 5–15% of episodes, and surgical site and miscellaneous infections account for the rest. *Klebsiella* is a cause of sepsis in neonates and of bacteremia in neutropenic patients. Like enteric GNB in general, *Klebsiella* rarely causes endocarditis or endovascular infection.

DIAGNOSIS

Klebsiellae are readily isolated and identified in the laboratory. These organisms usually ferment lactose, although the subspecies *rhinoscleromatis* and *ozaenae* are nonfermenters and are indole negative. hvKP usually possesses a hypermucoviscous phenotype (Fig. 186-1, *bottom*), although the sensitivity and specificity of this test are undefined and probably less than optimal. A better diagnostic test for hvKP is desirable.

TREATMENT KLEBSIELLA INFECTIONS



cKP and *K. oxytoca* have similar antibiotic resistance profiles. These species are intrinsically resistant to ampicillin and ticarcillin, and nitrofurantoin is inconsistently active against them. NHSN data for 2009–2010 documented resistance to third- and fourth-generation cephalosporins in 28.9% of CLABSI isolates of cKP and *K. oxytoca*, and INICC data for 2004–2009 identified resistance to third-generation cephalosporins in 76.3% of ICU isolates of *K. pneumoniae*. This increasing resistance is mediated primarily by plasmid-encoded ESBLs. In addition, such plasmids usually encode resistance to aminoglycosides, tetracyclines, and TMP-SMX. Furthermore, isolates of cKP that produce CTX-M ESBLs have been obtained from ambulatory patients with no recent health care contact (see the section on the treatment of extraintestinal *E. coli* infections for treatment considerations). Resistance to β -lactam/ β -lactamase inhibitor combinations and cephamycins independent of ESBL-encoding plasmids has also been described with increasing frequency, particularly in Latin America. The prevalence of fluoroquinolone resistance is 15–20% overall and is 50% among ESBL-containing strains. Given both the undesirability of treating the latter strains with penicillins or cephalosporins and the fluoroquinolone resistance often associated with ESBLs, empirical treatment of serious or health care-associated cKP and *K. oxytoca* infections with amikacin or carbapenems is prudent, as dictated by local susceptibilities. Predictably, however, the ESBL-driven use of carbapenems has selected for strains of cKP and *K. oxytoca* that express carbapenemases. NHSN data for 2009–2010 documented resistance to carbapenems in 12.8% of CLABSI isolates of cKP and *K. oxytoca*. Treatment of infections due to strains that produce carbapenemases is highly challenging; increasingly, these strains are nearly pan-resistant. The optimal choice for therapy is unclear. Tigecycline and the polymyxins (e.g., colistin) are the most active agents in vitro and are used most frequently. However, resistance to these agents is already emerging, and strains of cKP resistant to all known antimicrobial agents have been described in the United States and globally. Combination therapy is often used in this setting.

PROTEUS INFECTIONS

Proteus mirabilis causes 90% of *Proteus* infections, which occur in the community, LTCFs, and hospitals. *Proteus vulgaris* and *Proteus penneri* are associated primarily with infections acquired in LTCFs or hospitals. *Proteus* species are part of the colonic flora of a wide variety of mammals, birds, fish, and reptiles. The ability of these GNB to generate histamine

from contaminated fish has implicated them in the pathogenesis of scombroid (fish) poisoning (Chap. 474). *P. mirabilis* colonizes healthy humans (prevalence, 50%), whereas *P. vulgaris* and *P. penneri* are isolated primarily from individuals with underlying disease. The urinary tract is by far the most common site of *Proteus* infection, with adhesins, flagella, IgA-IgG protease, iron acquisition systems, and urease representing the principal known urovirulence factors. *Proteus* less commonly causes infection at a variety of other extraintestinal sites.

INFECTIOUS SYNDROMES

UTI Most *Proteus* infections arise from the urinary tract. *P. mirabilis* causes only 1–2% of UTIs in healthy women, and *Proteus* species collectively cause only 5% of hospital-acquired UTIs. However, *Proteus* is responsible for 10–15% of cases of complicated UTI, primarily those associated with catheterization; indeed, among UTI isolates from chronically catheterized patients, the prevalence of *Proteus* is 20–45%. This high prevalence is due in part to bacterial production of urease, which hydrolyzes urea to ammonia and results in alkalization of the urine. Alkalization of urine, in turn, leads to precipitation of organic and inorganic compounds, which contributes to formation of struvite and carbonate-apatite crystals, formation of biofilms on catheters, and/or development of frank calculi. *Proteus* becomes associated with the stones and biofilms; thereafter, it usually can be eradicated only by removal of the stones or the catheter. Over time, staghorn calculi may form within the renal pelvis and lead to obstruction and renal failure. Thus, urine samples with unexplained alkalinity should be cultured for *Proteus*, and identification of a *Proteus* species in urine should prompt consideration of an evaluation for urolithiasis.

Other Infections *Proteus* occasionally causes pneumonia (primarily in LTCF residents or hospitalized patients), nosocomial sinusitis, intraabdominal abscesses, biliary tract infection, surgical site infection, soft tissue infection (especially decubitus and diabetic ulcers), and osteomyelitis (primarily contiguous); in rare cases, it causes nontropical myositis. In addition, *Proteus* uncommonly causes neonatal meningitis, with the umbilicus frequently implicated as the source; this disease is often complicated by development of a cerebral abscess. Otogenic brain abscess also occurs.

Bacteremia The majority of *Proteus* bacteremia episodes originate from the urinary tract; however, any of the less common sites of infection as well as intravascular devices are also potential sources. Endovascular infection is rare. *Proteus* species are occasional agents of sepsis in neonates and of bacteremia in neutropenic patients.

DIAGNOSIS

Proteus is readily isolated and identified in the laboratory. Most strains are lactose negative, produce H_2S , and demonstrate characteristic swarming motility on agar plates. *P. mirabilis* and *P. penneri* are indole negative, whereas *P. vulgaris* is indole positive. The inability to produce ornithine decarboxylase differentiates *P. penneri* from *P. mirabilis*.

TREATMENT PROTEUS INFECTIONS

P. mirabilis is usually susceptible to most antimicrobial agents except tetracycline, nitrofurantoin, the polymyxins, and tigecycline. Resistance to ampicillin and first-generation cephalosporins has been acquired by 10–50% of strains. Overall, 10–15% of *P. mirabilis* isolates are resistant to fluoroquinolones; 5% of isolates in the United States now produce ESBLs. Furthermore, isolates of *P. mirabilis* that produce CTX-M ESBLs have been recovered from ambulatory patients with no recent health care contact (see the section on the treatment of extraintestinal *E. coli* infections for treatment considerations). *P. vulgaris* and *P. penneri* exhibit more extensive drug resistance than does *P. mirabilis*. Resistance to ampicillin and first-generation cephalosporins is the rule, and 30–40% of isolates are resistant to fluoroquinolones. Induction or selection of variants with stable derepression of chromosomal AmpC β -lactamase may

1034 occur with *P. vulgaris* isolates. Carbapenems, fourth-generation cephalosporins (e.g., cefepime), amikacin, TMP-SMX, and fosfomicin display excellent activity against *Proteus* species (90–100% of isolates susceptible).

ENTEROBACTER AND CRONOBACTER INFECTIONS

E. cloacae and *E. aerogenes* are responsible for most *Enterobacter* infections (65–75% and 15–25%, respectively); *Cronobacter sakazakii* (formerly *Enterobacter sakazakii*) and *Enterobacter gergoviae* are less commonly isolated (1% for each). *Enterobacter* species cause primarily health care–related infections. The organisms are widely prevalent in foods, environmental sources (including equipment at health care facilities), and a variety of animals. Few healthy humans are colonized, but the percentage increases significantly with LTCF residence or hospitalization. Although colonization is an important prelude to infection, direct introduction via IV lines (e.g., contaminated IV fluids or pressure monitors) also occurs. Extensive antibiotic resistance has developed in *Enterobacter* species and probably has contributed to the emergence of the organisms as prominent nosocomial pathogens. Individuals who have previously received antibiotic treatment, have comorbid disease, and are ICU residents are at greatest risk for infection. *Enterobacter* causes a spectrum of extraintestinal infections similar to that described for other GNB.

INFECTIOUS SYNDROMES

Pneumonia, UTI (particularly catheter-related), intravascular device–related infection, surgical site infection, and abdominal infection (primarily postoperative or related to devices such as biliary stents) are the most common syndromes encountered. Nosocomial sinusitis, meningitis related to neurosurgical procedures (including use of intracranial pressure monitors), osteomyelitis, and endophthalmitis after eye surgery are less frequent. *C. sakazakii* is associated with neonatal bacteremia, necrotizing enterocolitis, and meningitis (which is often complicated by brain abscess or ventriculitis); contaminated formula has been implicated as a source for such infections. *Enterobacter* bacteremia can result from infection at any anatomic site. In bacteremia of unclear origin, the contamination of IV fluids or medications, blood components or plasma derivatives, catheter-flushing fluids, pressure monitors, and dialysis equipment should be considered, particularly in an outbreak setting. *Enterobacter* can also cause bacteremia in neutropenic patients. *Enterobacter* endocarditis is rare, occurring primarily in association with illicit IV drug use or prosthetic valves.

DIAGNOSIS

Enterobacter is readily isolated and identified in the laboratory. Most strains are lactose positive and indole negative.

TREATMENT ENTEROBACTER INFECTIONS



Significant antimicrobial resistance exists among *Enterobacter* strains. Ampicillin and first- and second-generation cephalosporins have little or no activity. Extensive use of third-generation cephalosporins can induce or select for variants with stable derepression of AmpC β -lactamase, which confers resistance to these agents as well as monobactams (e.g., aztreonam) and—in many cases— β -lactam/ β -lactamase inhibitor combinations. Resistance may emerge during therapy; in one study, this phenomenon was documented in 20% of clinical isolates. De novo resistance should be considered when clinical deterioration follows initial improvement, and third-generation cephalosporins should be avoided in the treatment of serious *Enterobacter* infections. Cefepime is stable in the presence of AmpC β -lactamases; thus, it is a suitable option for treatment of *Enterobacter* infections so long as no coexistent ESBL is present. Detection of ESBLs in *Enterobacter* is difficult because of the presence of AmpC β -lactamase; nonetheless, their prevalence (particularly in *E. cloacae*) is known to be variable worldwide but is generally increasing and is now 5–50% overall. This increase is evidenced by HSN data, which documented resistance

to third- and fourth-generation cephalosporins in 37.4% of CLABSI *Enterobacter* isolates in the United States; fortunately, carbapenems, amikacin, and tigecycline have generally retained excellent activity (90–99% susceptibility) and fluoroquinolones have good activity (85–95% susceptibility). Once susceptibility data become available, it is critical to de-escalate the antimicrobial regimen whenever possible.

SERRATIA INFECTIONS



S. marcescens causes the majority (>90%) of *Serratia* infections; *Serratia liquefaciens*, *Serratia rubidaea*, *Serratia fonticola*, *Serratia grimesii*, *Serratia plymuthica*, and *Serratia odorifera* are isolated occasionally. *Serratiae* are found primarily in the environment (including in health care institutions), particularly in moist settings. *Serratiae* have been isolated from a variety of animals, insects, and plants, but healthy humans are rarely colonized. In LTCFs or hospitals, reservoirs for the organisms include the hands and fingernails of health care personnel, food, milk (on neonatal units), sinks, respiratory and other medical equipment or devices, pressure monitors, IV solutions or parenteral medications (particularly those generated by compounding pharmacies), prefilled syringes and multiple-access medication vials (e.g., heparin, saline), blood products (e.g., platelets), hand soaps and lotions, irrigation solutions, and even disinfectants. Infection results from either direct inoculation (e.g., via IV fluid) or colonization (primarily of the respiratory tract). Sporadic infection is most common, but epidemics (often involving MDR strains in adult and neonatal ICUs) and common-source outbreaks also occur. The spectrum of extraintestinal infections caused by *Serratia* is similar to that for other GNB. *Serratia* species are usually considered causative agents of health care–associated infection and account for 1–3% of hospital-acquired infections. However, population-based laboratory surveillance studies in Canada and Australia demonstrated that community-acquired infections occur more commonly than was previously appreciated.

INFECTIOUS SYNDROMES

The respiratory tract, the genitourinary tract, intravascular devices, the eye (contact lens–associated keratitis and other ocular infections), surgical wounds, and the bloodstream (from contaminated infusions) are the most common sites of *Serratia* infection; the former five sites are the most common sources of *Serratia* bacteremia. Soft tissue infections (including myositis, fasciitis, mastitis), osteomyelitis, abdominal and biliary tract infection (postprocedural), and septic arthritis (primarily from intraarticular injections) occur less commonly. *Serratiae* are uncommon causes of neonatal or postsurgical meningitis and of bacteremia in neutropenic patients. Endocarditis is rare.

DIAGNOSIS

Serratiae are readily cultured and identified by the laboratory and are usually lactose and indole negative. Some *S. marcescens* strains and *S. rubidaea* are red pigmented.

TREATMENT SERRATIA INFECTIONS



Most *Serratia* strains (>80%) are resistant to ampicillin, amoxicillin-clavulanate, ampicillin-sulbactam, first-generation cephalosporins, cephamycins, nitrofurantoin, and colistin. In general, >90% of *Serratia* isolates are susceptible to other antibiotics appropriate for use against GNB. Induction or selection of variants with stable derepression of chromosomal AmpC β -lactamases may develop during therapy. Both in the United States and globally, the prevalence of ESBL-producing isolates is generally low (<5%), but rates of 20–30% have been reported in Asia and Latin America. Acquisition of carbapenemase-encoding genes is uncommon but increasing.

CITROBACTER INFECTIONS

C. freundii and *Citrobacter koseri* cause most human *Citrobacter* infections, which are epidemiologically and clinically similar to *Enterobacter* infections. *Citrobacter* species are commonly present in

water, food, soil, and certain animals. *Citrobacter* is part of the normal fecal flora in a minority of healthy humans, but colonization rates are higher in LTCFs and hospitals—the settings in which nearly all *Citrobacter* infections occur. *Citrobacter* species account for 1–2% of nosocomial infections. The affected hosts are usually immunocompromised or have comorbid disease. *Citrobacter* causes extraintestinal infections similar to those described for other GNB.

INFECTIOUS SYNDROMES

The urinary tract accounts for 40–50% of *Citrobacter* infections. Less commonly involved sites include the biliary tree (particularly with stones or obstruction), the respiratory tract, surgical sites, soft tissue (e.g., decubitus ulcers), the peritoneum, and intravascular devices. Osteomyelitis (usually from a contiguous focus), adult central nervous system infection (from neurosurgical or other types of meningeal disruption), and myositis occur rarely. *Citrobacter* (primarily *C. koseri*) also causes 1–2% of neonatal meningitis cases, of which 50–80% are complicated by brain abscess. Further, case reports in adults suggest that *C. koseri* infection has a predilection for abscess formation. Bacteremia is most often due to UTI, biliary/abdominal infection, or intravascular device infection. *Citrobacter* occasionally causes bacteremia in neutropenic patients. Endocarditis and endovascular infections are rare.

DIAGNOSIS

Citrobacter species are readily isolated and identified; 35–50% of isolates are lactose positive, and 100% are oxidase negative. *C. freundii* is indole negative, whereas *C. koseri* is indole positive.

TREATMENT CITROBACTER INFECTIONS

C. freundii is more extensively resistant to antibiotics than is *C. koseri*. More than 90% of isolates are resistant to ampicillin and first- and second-generation cephalosporins. *Citrobacter* species (except *C. koseri*) possess AmpC β -lactamases; induction or selection of variants with stable derepression may develop during therapy. Resistance to antipseudomonal penicillins, aztreonam, fluoroquinolones, gentamicin, and third-generation cephalosporins is variable but increasing. The prevalence of ESBL-producing isolates is <5%. Carbapenems, amikacin, cefepime, tigecycline (with which clinical experience is limited), fosfomycin (which is available in the United States only as an oral formulation), and colistin (which is an agent of last resort because of potential toxicities) are most active, with >90% of strains susceptible.

MORGANELLA AND PROVIDENCIA INFECTIONS

M. morgani, *Providencia stuartii*, and (less frequently) *Providencia rettgeri* are the members of their respective genera that cause human infections. The epidemiologic associations, pathogenic properties, and clinical manifestations of these organisms resemble those of *Proteus* species. However, *Morganella* and *Providencia* occur more commonly among LTCF residents; to a lesser degree, they affect hospitalized patients. In settings with extensive use of polymyxins and tigecycline, these organisms may become increasingly common because of their intrinsic resistance to these agents.

INFECTIOUS SYNDROMES

These species are primarily urinary tract pathogens, causing UTIs that are most often associated with long-term (>30-day) catheterization. Such infections commonly lead to biofilm formation and catheter encrustation (sometimes causing catheter obstruction) or to the development of struvite bladder or renal stones (sometimes causing renal obstruction and serving as foci for relapse). *Morganella* is also commonly isolated from snakebite infection. Other, less common infectious syndromes include surgical site infection, soft tissue infection (primarily involving decubitus and diabetic ulcers), burn site infection, pneumonia (particularly ventilator-associated), intravascular device infection, and intraabdominal infection. Rarely, the other extraintestinal infections described for GNB also occur. Bacteremia is uncommon; any infected site can serve as the source, but the urinary

tract accounts for most cases, with the next most common sources being surgical site, soft tissue, and hepatobiliary infections.

DIAGNOSIS

M. morgani and *Providencia* are readily isolated and identified. Nearly all isolates are lactose negative and indole positive.

TREATMENT MORGANELLA AND PROVIDENCIA INFECTIONS



Morganella and *Providencia* may be extensively resistant to antibiotics. Most isolates are resistant to ampicillin, first-generation cephalosporins, nitrofurantoin, fosfomycin, tigecycline, and the polymyxins; 40% are resistant to fluoroquinolones. *Morganella* and *Providencia* possess inducible AmpC β -lactamases; clinically significant induction or selection of stably derepressed mutants may develop during therapy. Resistance to antipseudomonal penicillins, aztreonam, gentamicin, TMP-SMX, and second- and third-generation cephalosporins is emerging but is still variably prevalent. The β -lactamase inhibitor tazobactam increases susceptibility to β -lactam agents, but sulbactam and clavulanic acid do not. Carbapenems, amikacin, and cefepime are the most active agents (>90% of isolates susceptible); however, resistance to the carbapenems, when present, is a concern because of the inherent resistance of *Morganella* and *Providencia* to the polymyxins and tigecycline. Removal of a colonized catheter or stone is critical for eradication of UTI.

EDWARDSIELLA INFECTIONS



E. tarda is the only member of the genus *Edwardsiella* that is associated with human disease. This organism is found predominantly in freshwater and marine environments and in the associated aquatic animal species. Human acquisition occurs primarily during interaction with these reservoirs and ingestion of inadequately cooked aquatic animals. *E. tarda* infection is rare in the United States; recently reported cases are mostly from Southeast Asia. This pathogen shares clinical features with *Salmonella* species (as an intestinal pathogen; Chap. 190), *Vibrio vulnificus* (as an extraintestinal pathogen; Chap. 193) and *Aeromonas hydrophila* (as both an intestinal and extraintestinal pathogen; Chap. 183e).

INFECTIOUS SYNDROMES

Gastroenteritis is the predominant infectious syndrome (50–80% of infections). Self-limiting watery diarrhea is most common, but severe colitis also occurs. The most common extraintestinal infection is wound infection due to direct inoculation, which is often associated with freshwater, marine, or snake-related injuries. Other infectious syndromes result from invasion of the gastrointestinal tract and subsequent bacteremia. Most afflicted hosts have comorbidities (e.g., hepatobiliary disease, iron overload, cancer, or diabetes mellitus). A primary bacteremic syndrome, sometimes complicated by meningitis, has a 40% case-fatality rate. Visceral (primarily hepatic) and intra-peritoneal abscesses also occur. Endocarditis and empyema have been described.

DIAGNOSIS

Although *E. tarda* can readily be isolated and identified, most laboratories do not routinely seek to identify it in stool samples. Production of hydrogen sulfide is a characteristic biochemical property.

TREATMENT EDWARDSIELLA INFECTIONS

E. tarda is susceptible to most antimicrobial agents appropriate for use against GNB. Gastroenteritis is generally self-limiting, but treatment with a fluoroquinolone may hasten resolution. In the setting

1036 of severe sepsis, fluoroquinolones, third- and fourth-generation cephalosporins, carbapenems, and amikacin—either alone or in combination—are the safest choices pending susceptibility data.

INFECTIONS CAUSED BY MISCELLANEOUS GENERA

Species of *Hafnia*, *Kluyvera*, *Cedecea*, *Pantoea*, *Ewingella*, *Leclercia*, and *Photorhabdus* are occasionally isolated from diverse clinical specimens, including blood, sputum, urine, cerebrospinal fluid, joint fluid, bile, and wounds. These organisms are rare and usually cause infection in a compromised host or in the setting of an invasive procedure or a foreign body. Cephalosporinases from *Kluyvera* have been implicated as the progenitors of CTX-M ESBLs.

187 *Acinetobacter* Infections

David L. Paterson, Anton Y. Peleg

Infections with bacteria of the genus *Acinetobacter* are established as a significant problem worldwide. *Acinetobacter baumannii* is particularly formidable because of its propensity to acquire antibiotic resistance determinants. Endemic infections caused by strains of *A. baumannii* resistant to multiple antibiotic classes, including carbapenems, are a serious concern in many specialized hospital units, especially intensive care units (ICUs). The foremost implication of infection with carbapenem-resistant *A. baumannii* is the need to use “last-line” antibiotics such as colistin, polymyxin B, or tigecycline; these options have the potential to render these bacteria resistant to all available antibiotics.

DEFINITION

Acinetobacter species are oxidase-negative, nonfermenting, short, gram-negative bacilli. They were traditionally thought of as nonmotile—a characteristic from which the genus name was derived (from the Greek *akineto*, meaning “nonmotile”). However, recent work has shown that *Acinetobacter* organisms demonstrate motility under certain growth conditions. The bacteria grow well at 37°C in aerobic conditions on a range of laboratory media (e.g., blood agar). Some species may not grow on MacConkey agar. Differentiation of *Acinetobacter* species is difficult with the means typically available to most clinical microbiology laboratories, including commercial semiautomated identification systems. The commonly used matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) systems are undergoing evaluation for species-level identification of *Acinetobacter*. DNA–DNA hybridization is a method used for speciation in reference laboratories. Naturally occurring oxacillinase genes (*bla_{OXA}*) have been identified in several *Acinetobacter* species, and their detection by polymerase chain reaction can aid in species identification.

ETIOLOGY

Widely distributed in nature, *Acinetobacter* species can be found in water, in soil, and on vegetables. *Acinetobacter* is a component of the human skin flora and is sometimes identified as a contaminant in blood samples collected for culture. Fecal carriage can be detected in both healthy and hospitalized individuals. However, despite the ubiquity of some *Acinetobacter* species, the natural habitat of the most clinically important species, *A. baumannii*, remains to be fully defined.

EPIDEMIOLOGY

A. baumannii infections have been diagnosed in patients on all inhabited continents. The vast majority of infections occur in hospitalized patients and other patients with significant

health-care contact. Outbreaks of carbapenem-resistant *A. baumannii* are particularly problematic. A significant issue is the introduction of carbapenem-resistant *A. baumannii* into hospitals as a result of medical transfers, especially from hospitals where the organism is highly endemic.

The Americas In 1991 and 1992, outbreaks of carbapenem-resistant *A. baumannii* infection occurred in a hospital in New York City. Subsequently, numerous other hospitals in the United States and South America have had outbreaks of carbapenem-resistant *A. baumannii*. Infections with *A. baumannii* among military personnel from the United States and Canada injured in Iraq or Afghanistan were widely observed beginning in 2002. *Acinetobacter* was one of the most common causes of bloodstream infections and bone and soft tissue infections after war-related injury. An epidemiologic investigation revealed that *A. baumannii* could be grown from environmental sites in field hospitals and that the environmental strains were closely related genotypically to clinical isolates.

Europe *A. baumannii* infections have posed a substantial clinical challenge in many parts of Europe since the early 1980s. Three clones (European clones I, II, and III) have been the predominant causes of *A. baumannii* infection in hospitals in Europe. Carbapenem resistance in *A. baumannii* is a significant issue in many European countries, most notably the United Kingdom, Greece, Italy, Spain, and Turkey.

Asia, Australia, the Middle East, and Africa Although surveillance data are sparse from many countries in these regions, problems with carbapenem-resistant *A. baumannii* abound. Community-acquired infections are well described in northern Australia and some parts of Asia. These infections may be more likely in men >45 years of age who have histories of cigarette smoking, alcoholism, diabetes mellitus, or chronic obstructive airway disease. Community-acquired strains are more susceptible to antimicrobial agents than are hospital-acquired strains, but the clinical presentation of community-acquired disease is quite distinct and is characterized by overwhelming infection with severe pneumonia, septic shock, and multiorgan failure.

PATHOGENESIS

A. baumannii colonizes patients exposed to heavily contaminated hospital environments or to the hands of health care workers in these locations. Emerging data suggest that the organism can be found in the air in rooms of patients infected with *Acinetobacter*. Colonization of the upper airways in mechanically ventilated patients may lead to nosocomial pneumonia. Colonization of the skin may lead to central line-associated bloodstream infection, catheter-associated urinary tract infection (UTI), wound infection, or postneurosurgical meningitis. Throat carriage and microaspiration may be involved in the pathogenesis of community-acquired pneumonia due to *A. baumannii*.

Much less is known about the virulence mechanisms of and host responses to *A. baumannii* than about these aspects of other pathogenic gram-negative bacteria. Because of the emergence of multidrug-resistant strains, including those resistant to all available antibiotics, the impetus to study *A. baumannii* pathogenesis has grown. Novel targets for antibacterial drug development are desperately required, and drugs that have antivirulence mechanisms may provide new therapeutic options. Specific virulence mechanisms in *A. baumannii* include iron acquisition and transport systems; outer-membrane protein A (OmpA), which mediates mammalian cell adhesion, invasion, and cytotoxicity through mitochondrial damage and initiation of caspase-dependent apoptosis; lipopolysaccharide (LPS); and proteins important in the formation of biofilm on abiotic and biotic surfaces. Biofilm formation on abiotic surfaces is dependent on a pilus assembly system, which in turn is controlled by a traditional two-component regulatory system mediated by *bfmR*. Also important in biofilm formation are a gene that encodes a biofilm-associated protein (Bap); OmpA; the quorum-sensing gene *abaI*, which controls the secretion of 3-hydroxy- C_{12} -homoserine lactone; and the *pga* locus, which is essential for the production of the polysaccharide poly- β -1,6-*N*-acetylglucosamine. Most recently, a global virulence regulator known



as GacSA was described as important in regulating *A. baumannii* biofilms, motility, growth in human serum, and virulence in a mammalian infection model.

New model systems for the study of *A. baumannii* infection, including both nonmammalian (invertebrate) and mammalian models, have been described. Furthermore, the use of *A. baumannii* transposon-generated mutant libraries to screen for mutants with attenuated growth in human biologic fluids (serum and ascites fluid) has allowed the identification of new virulence mechanisms. These include phospholipase D; capsule production mediated by *ptk* and *epsA*; penicillin-binding protein 7/8 encoded by the *pbpG* gene; and a glycosyltransferase important for LPS biosynthesis encoded by the *lpsB* gene.

The LPS of *A. baumannii* appears to play a significant role in eliciting host responses. In studies with knockout mice, Toll-like receptor 4 and CD14 were shown to be important in host recognition, signaling, and cytokine production in response to *A. baumannii*. Humoral responses targeting iron-regulated outer-membrane proteins and the O-polysaccharide component of LPS also have been described.

APPROACH TO THE PATIENT: *Acinetobacter* Infection

Acinetobacter must be considered in the differential diagnosis of hospital-acquired pneumonia, central line-associated bloodstream infection, posttraumatic wound infection in military personnel, and postneurosurgical meningitis.

CLINICAL MANIFESTATIONS

Pneumonia It may be difficult to distinguish between upper-airway colonization with *A. baumannii* and hospital-acquired pneumonia. An estimated 5–10% of cases of ventilator-associated pneumonia are due to *A. baumannii*, although much regional variation exists. Typically, patients with *A. baumannii* ventilator-associated pneumonia have had a prolonged stay in an ICU; in outbreak situations, however, patients may acquire the infection within days of arrival in an ICU.



Community-acquired pneumonia due to *A. baumannii* has been described in tropical regions of Australia and Asia. The disease typically occurs during the “wet” season among people with a history of alcohol abuse. Infection may result in fulminant pneumonia requiring admission to an ICU, with a mortality rate of ~50%.

Bloodstream Infection Although *A. baumannii* accounts for only ~1–2% of nosocomial bloodstream infections, crude mortality rates from these infections may be as high as 40%. Sources of bloodstream infection are typically a central line or underlying pneumonia, UTI, or wound infection.

Traumatic Battlefield and Other Wounds *A. baumannii* is a well-known pathogen in burn units. This organism is commonly isolated from wounds of combat casualties; it was the most commonly isolated organism in one assessment of combat victims with open tibial fractures but did not appear to contribute directly to persistent nonunion or the need for amputation.

Meningitis *A. baumannii* may cause meningitis following neurosurgical procedures. Patients typically have an external ventricular drain in situ.

Urinary Tract Infection *A. baumannii* is an occasional cause of catheter-associated UTI. It is highly unusual for this organism to cause uncomplicated UTI in healthy women.

Other Clinical Manifestations A small number of case reports describe *Acinetobacter* prosthetic-valve endocarditis and endophthalmitis/keratitis. The latter is sometimes related to contact lens use or eye surgery.

DIAGNOSIS

Acinetobacter infection should be suspected when plump coccobacilli are seen in Gram’s-stained respiratory tract secretions, blood cultures, or cerebrospinal fluid. Sometimes the organisms are difficult to

de-stain. Given their small size, they may be misidentified as either gram-negative or gram-positive cocci.

TREATMENT ACINETOBACTER INFECTION (TABLE 187-1)

Treatment is hampered by the remarkable ability of *A. baumannii* to upregulate or acquire antibiotic resistance determinants. The most prominent example is that of β -lactamases, including those capable of inactivating carbapenems, cephalosporins, and penicillins. These enzymes, which include the OXA-type β -lactamases (e.g., OXA-23), the metallo- β -lactamases (e.g., NDM), and rarely KPC-type carbapenemases, are typically resistant to currently available β -lactamase inhibitors such as clavulanate or tazobactam. Plasmids that harbor genes encoding these β -lactamases may also harbor genes encoding resistance to aminoglycosides and sulfur antibiotics. The end result is that carbapenem-resistant *A. baumannii* may become truly multidrug resistant.

Selection of empirical antibiotic therapy when *A. baumannii* is suspected is challenging and must rely on a knowledge of local epidemiology. Receipt of prompt, effective antibiotic therapy is the goal. Given the diversity of resistance mechanisms in *A. baumannii*, definitive therapy should be based on the results of antimicrobial susceptibility testing. Carbapenems (imipenem, meropenem, and doripenem but not ertapenem) have long been thought of as the agents of choice for serious *A. baumannii* infections. However, the clinical utility of carbapenems is now widely jeopardized by the production of carbapenemases, as described above. Sulbactam may be an alternative to carbapenems. Unlike other β -lactamase inhibitors (e.g., clavulanic acid and tazobactam), sulbactam has intrinsic activity against *Acinetobacter*; this activity is mediated by the drug’s binding to penicillin-binding protein 2 rather than by its ability to inhibit β -lactamases. Sulbactam is commercially available in a combined formulation with either ampicillin or cefoperazone and may also be available as a single agent in some countries. Despite the absence of randomized clinical trials, sulbactam seems to be equivalent to carbapenems in clinical effectiveness against susceptible strains.

Therapy for carbapenem-resistant *A. baumannii* is particularly problematic. The only currently available choices are polymyxins (colistin and polymyxin B) and tigecycline. Neither option is perfect. Polymyxins may be nephrotoxic and neurotoxic. Definition of the optimal dose and schedule for administration of polymyxins to patients in vulnerable groups (e.g., those requiring renal replacement therapy) remains challenging, and emergence of resistance in association with monotherapy is a concern. Conventional doses of tigecycline may not result in serum concentrations adequate to treat bloodstream infections. Resistance of *A. baumannii* to tigecycline may develop during treatment with this drug.

As a consequence of these issues with the polymyxins and tigecycline, combination therapy is now favored for carbapenem-resistant *Acinetobacter*. However, in a randomized controlled trial, 30-day mortality was not reduced by the addition of rifampin to colistin. Nevertheless, a significant increase in microbiologic eradication was observed in the colistin plus rifampin arm over that attained with

TABLE 187-1 TREATMENT OPTIONS FOR ACINETOBACTER INFECTIONS

Antibiotic	Comments
Sulbactam	Intrinsic activity against <i>Acinetobacter</i> , not linked to β -lactamase inhibition
Trimethoprim-sulfamethoxazole	May be an option for urinary tract infection or wound infection
Meropenem	Carbapenem resistance now widespread
Amikacin	May be an option for carbapenem-resistant strains
Tigecycline	May be an option for carbapenem-resistant strains but inappropriate for urinary tract infection, bloodstream infection, or meningitis
Colistin or polymyxin B	May be an option for carbapenem-resistant strains, but pharmacokinetics not yet well understood

1038 colistin alone. Combinations of polymyxins with a carbapenem look more promising and are being evaluated in prospective clinical trials. Fosfomycin has poor activity against *Acinetobacter* and should not be relied upon for treatment. Clearly, new treatment options are needed for serious *Acinetobacter* infections.

COMPLICATIONS AND PROGNOSIS

Given the propensity of *A. baumannii* to cause infections in seriously ill patients in ICUs, it is not surprising that *A. baumannii* infections are associated with high mortality rates. Thus a pertinent question is whether *A. baumannii* infections are associated with high attributable mortality rates after the severity of illness is controlled for. A number of studies have addressed this issue but have had disparate results. Whether the discrepant results can be explained purely by methodologic differences is unknown at present.

PREVENTION

Multidrug-resistant *A. baumannii* clearly causes outbreaks of infection and then establishes endemicity. In endemic situations, a small number of strain types predominate. In the 1991–1992 outbreaks in New York City, for example, two strain types accounted for more than 80% of carbapenem-resistant isolates. This “oligoclonality” plainly demonstrates the potential importance of infection control interventions in response to outbreaks of multidrug-resistant *A. baumannii* infection.

The hospital environment is an important reservoir of organisms capable of colonizing patients and causing infection. Environmental sources of *A. baumannii* include computer keyboards, glucometers, multidose medication vials, IV nutrition, inadequately sterilized reusable arterial pressure transducers, ventilator tubing, suction catheters, humidifiers, containers of distilled water, urine collection jugs, and moist bedding articles. Pulsatile-lavage wound treatment—a high-pressure irrigation system used to debride wounds—has been associated with an outbreak of *A. baumannii* infection.

Contaminated inanimate objects should be removed from the patient-care environment or subjected to enhanced environmental cleaning. Although contact-isolation procedures (use of gloves and gowns when dealing with colonized patients or their environment), accommodation of patients in single rooms, and improved hand hygiene are critical, attention to the patient-care environment may be the only measure that leads to control of outbreaks of *A. baumannii* infection. One study found that *Acinetobacter* can be cultured from the air in rooms of patients with *A. baumannii* infection; the infection-control implications are not yet clear.

diseases and whether it is protective against some recently emergent medical problems, such as childhood-onset asthma and obesity.

ETIOLOGIC AGENT

Helicobacter pylori *H. pylori* is a gram-negative bacillus that has naturally colonized humans for at least 100,000 years, and probably throughout human evolution. It lives in gastric mucus, with a small proportion of the bacteria adherent to the mucosa and possibly a very small number of the organisms entering cells or penetrating the mucosa; the organism’s distribution is never systemic. Its spiral shape and flagella render *H. pylori* motile in the mucus environment. The organism has several acid-resistance mechanisms, most notably a highly expressed urease that catalyzes urea hydrolysis to produce buffering ammonia. *H. pylori* is microaerophilic (i.e., requires low levels of oxygen), is slow-growing, and requires complex growth media in vitro.

Other *Helicobacter* Species A very small proportion of gastric *Helicobacter* infections are due to species other than *H. pylori*, possibly acquired as zoonoses. These non-*pylori* gastric helicobacters are associated with low-level inflammation and occasionally with disease. In immunocompromised hosts, several nongastric (intestinal) *Helicobacter* species can cause disease with clinical features resembling those of *Campylobacter* infections; these species are covered in [Chap. 192](#).

EPIDEMIOLOGY

Prevalence and Risk Factors The prevalence of *H. pylori* among adults is <30% in most parts of the United States and in other developed countries as opposed to >80% in most developing countries. In the United States, prevalence varies with age: up to 50% of 60-year-old persons, ~20% of 30-year-old persons, and fewer than 10% of children are colonized. *H. pylori* is usually acquired in childhood. The age association is due mostly to a birth-cohort effect whereby current 60-year-olds were more commonly colonized as children than are current children. Spontaneous acquisition or loss of *H. pylori* in adulthood is uncommon. Childhood acquisition explains why the main risk factors for infection are markers of crowding and social deprivation in childhood.

Transmission Humans are the only important reservoir of *H. pylori*. Children may acquire the organism from their parents (most often the primary caregiver) or from other children. The former is more common in developed countries and the latter in less developed countries. Whether transmission takes place more often by the fecal-oral or the oral-oral route is unknown, but *H. pylori* is easily cultured from vomitus and gastroesophageal refluxate and is less easily cultured from stool.

PATHOLOGY AND PATHOGENESIS

H. pylori colonization induces *chronic superficial gastritis*, a tissue response in the stomach that includes infiltration of the mucosa by both mononuclear and polymorphonuclear cells. (The term *gastritis* should be used specifically to describe histologic features; it has also been used to describe endoscopic appearances and even symptoms, but these features do not correlate with microscopic findings or even with the presence of *H. pylori*.) Although *H. pylori* is capable of numerous adaptations that prevent excessive stimulation of the immune system, colonization is accompanied by a considerable persistent local and systemic immune response, including the production of antibodies and cell-mediated responses. However, these responses are ineffective in clearing the bacterium. This inefficient clearing appears to be due in part to *H. pylori*’s downregulation of the immune system, which fosters its own persistence.


Most *H. pylori*-colonized persons do not develop clinical sequelae. That some persons develop overt disease whereas others do not is related to a combination of factors: bacterial strain differences, host susceptibility to disease, and environmental factors.

Bacterial Virulence Factors Several *H. pylori* virulence factors are more common among strains that are associated with disease than among those that are not. The *cag* island is a group of genes

188 *Helicobacter pylori* Infections

John C. Atherton, Martin J. Blaser

DEFINITION

 *Helicobacter pylori* colonizes the stomach in ~50% of the world’s human population, essentially for life unless eradicated by antibiotic treatment. Colonization with this organism is the main risk factor for peptic ulceration ([Chap. 348](#)) as well as for gastric adenocarcinoma and gastric mucosa-associated lymphoid tissue (MALT) lymphoma ([Chap. 109](#)). Treatment for *H. pylori* has revolutionized the management of peptic ulcer disease, providing a permanent cure in most cases. Such treatment also represents first-line therapy for patients with low-grade gastric MALT lymphoma. Treatment of *H. pylori* is of no benefit in the treatment of gastric adenocarcinoma, but prevention of *H. pylori* colonization could potentially prevent gastric malignancy and peptic ulceration. In contrast, increasing evidence indicates that lifelong *H. pylori* colonization may offer some protection against complications of gastroesophageal reflux disease (GERD), including esophageal adenocarcinoma. Recent research has focused on whether *H. pylori* colonization is also a risk factor for some extragastric



that encodes a bacterial type IV secretion system. Through this system, an effector protein, CagA, is translocated into epithelial cells, where it may be transformed by phosphorylation and induces host cell signal transduction; proliferative, cytoskeletal, and inflammatory changes in the cell result. The protein at the tip of the secretory apparatus, CagL, binds to integrins on the cell surface, transducing further signaling. Finally, soluble components of the peptidoglycan cell wall enter the cell, mediated by the same secretory system. These components are recognized by the emergency intracellular bacterial receptor Nod1, which stimulates a proinflammatory cytokine response resulting in enhanced gastric inflammation. Carriage of *cag*-positive strains increases the risk of peptic ulcer or gastric adenocarcinoma. A second major virulence factor is the vacuolating cytotoxin VacA, which forms pores in cell membranes. VacA is polymorphic, and carriage of more active forms also increases the risk of disease. Other bacterial factors that are associated with increased disease risk include adhesins, such as BabA (which binds to blood group antigens on epithelial cells), and incompletely characterized factors, such as another recently described bacterial type 4 secretion system.

Host Genetic and Environmental Factors The best-characterized host determinants of disease are genetic polymorphisms leading to enhanced activation of the innate immune response, including polymorphisms in cytokine genes or in genes encoding bacterial recognition proteins such as Toll-like receptors. For example, colonized people with polymorphisms in the interleukin 1 gene that increase the production of this cytokine in response to *H. pylori* infection are at increased risk of gastric adenocarcinoma. In addition, environmental cofactors are important in pathogenesis. Smoking increases the risks of duodenal ulcers and gastric cancer in *H. pylori*-positive individuals. Diets high in salt and preserved foods increase cancer risk, whereas diets high in antioxidants and vitamin C are modestly protective.

Distribution of Gastritis and Differential Disease Risk The pattern of gastric inflammation is associated with disease risk: antral-predominant gastritis is most closely linked with duodenal ulceration, whereas pan-gastritis is linked with gastric ulceration and adenocarcinoma. This difference probably explains why patients with duodenal ulceration are not at high risk of developing gastric adenocarcinoma later in life, despite being colonized by *H. pylori*.

PATHOGENESIS OF DUODENAL ULCERATION How gastric colonization causes duodenal ulceration is now becoming more clear. *H. pylori*-induced inflammation of the gastric antrum diminishes the number of somatostatin-producing D cells. Because somatostatin inhibits gastrin release, gastrin levels are higher than in *H. pylori*-negative persons, and these higher levels lead to increased meal-stimulated acid secretion from the relatively spared gastric corpus. How this situation increases duodenal ulcer risk remains controversial, but the increased acid secretion may contribute to the formation of the potentially protective gastric metaplasia found in the duodenum of duodenal ulcer patients. Gastric metaplasia in the duodenum may become colonized by *H. pylori* and subsequently inflamed and ulcerated.

PATHOGENESIS OF GASTRIC ULCERATION AND GASTRIC ADENOCARCINOMA The pathogenesis of these conditions is less well understood, although both arise in association with pan- or corpus-predominant gastritis. The hormonal changes described above still occur, but the inflammation in the gastric corpus means that it produces less acid (hypochlorhydria) despite hypergastrinemia. Gastric ulcers usually occur

at the junction of antral and corpus-type mucosa, an area that is often particularly inflamed. Gastric cancer probably stems from progressive DNA damage and the survival of abnormal epithelial cell clones. The DNA damage is thought to be due principally to reactive oxygen and nitrogen species arising from inflammatory cells, perhaps in relation to other bacteria that survive in a hypochlorhydric stomach. Longitudinal analyses of gastric biopsy specimens taken years apart from the same patient show that the common *intestinal* type of gastric adenocarcinoma follows stepwise changes from simple gastritis to gastric atrophy, intestinal metaplasia, and dysplasia. A second, *diffuse* type of gastric adenocarcinoma found more commonly in younger adults may arise directly from chronic gastritis without atrophic changes.

CLINICAL MANIFESTATIONS

Essentially all *H. pylori*-colonized persons have histologic gastritis, but only ~10–15% develop associated illnesses such as peptic ulceration, gastric adenocarcinoma, or gastric lymphoma (Fig. 188-1). Rates among women are less than half of those among men for both diseases.



Peptic Ulcer Disease Worldwide, >80% of duodenal ulcers and >60% of gastric ulcers are related to *H. pylori* colonization (Chap. 348). However, in particular, the proportion of gastric ulcers caused by aspirin and nonsteroidal anti-inflammatory drugs (NSAIDs) is increasing, and in many developed countries these drugs have overtaken *H. pylori* as a cause of gastric ulceration. The main lines of evidence supporting an ulcer-promoting role for *H. pylori* are that (1) the presence of the organism is a risk factor for the development of ulcers, (2) non-NSAID-induced ulcers rarely develop in the absence of *H. pylori*, (3) eradication of *H. pylori* virtually abolishes long-term ulcer relapse, and (4) experimental *H. pylori* infection of gerbils can cause gastric ulceration.

Gastric Adenocarcinoma and Lymphoma Prospective nested case-control studies have shown that *H. pylori* colonization is a risk factor for adenocarcinomas of the distal (noncardia) stomach (Chap. 109). Long-term experimental infection of gerbils also may result in gastric adenocarcinoma. Moreover, *H. pylori* may induce primary gastric

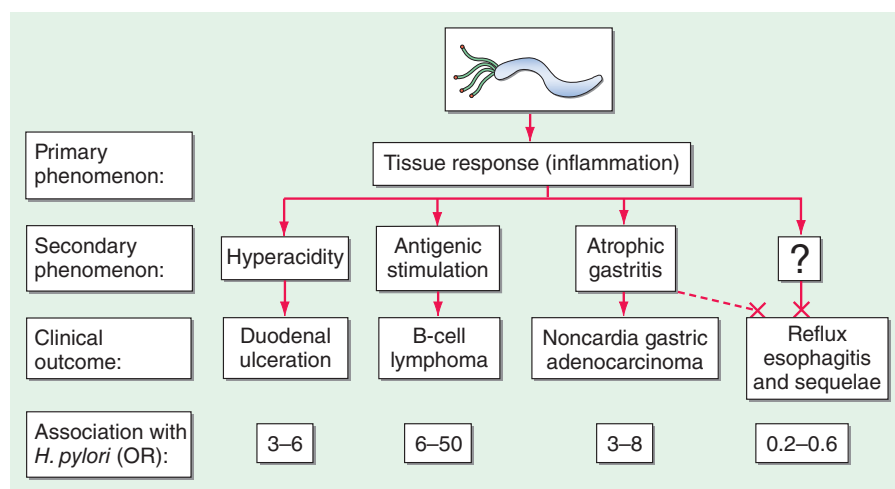


FIGURE 188-1 Schematic of the relationships between colonization with *Helicobacter pylori* and diseases of the upper gastrointestinal tract. Essentially all persons colonized with *H. pylori* develop a host response, which is generally termed *chronic gastritis*. The nature of the host's interaction with the particular bacterial population determines the clinical outcome. *H. pylori* colonization increases the lifetime risk of peptic ulcer disease, noncardia gastric cancer, and B-cell non-Hodgkin's gastric lymphoma (odds ratios [ORs] for all, >3). In contrast, a growing body of evidence indicates that *H. pylori* colonization (especially with *cagA*⁺ strains) protects against adenocarcinoma of the esophagus (and the sometimes related gastric cardia) and premalignant lesions such as Barrett's esophagus (OR, <1). Although the incidences of peptic ulcer disease (cases not due to nonsteroidal anti-inflammatory drugs) and noncardia gastric cancer are declining in developed countries, the incidence of adenocarcinoma of the esophagus is increasing. (Adapted from MJ Blaser: Hypothesis: The changing relationships of *Helicobacter pylori* and humans: Implications for health and disease. *J Infect Dis* 179:1523, 1999, with permission.)

1040 lymphoma, although this condition is much less common. Many low-grade gastric B-cell lymphomas are dependent on *H. pylori* for continuing growth and proliferation, and these tumors may regress either fully or partially after *H. pylori* eradication. However, they require careful short- and long-term monitoring, and some necessitate additional treatment with chemotherapeutic agents.

Functional Dyspepsia Many patients have upper gastrointestinal symptoms but have normal results on upper gastrointestinal endoscopy (so-called functional or nonulcer dyspepsia; **Chap. 348**). Because *H. pylori* is common, some of these patients will be colonized with the organism. *H. pylori* eradication leads to symptom resolution a little more commonly (from 0 to 7% in different studies) than does placebo treatment. Whether such patients have peptic ulcers in remission at the time of endoscopy or whether a small subgroup of patients with “true” functional dyspepsia respond to *H. pylori* treatment is unclear.

Protection Against Peptic Esophageal Disease, Including Esophageal Adenocarcinoma Much interest has focused on a protective role for *H. pylori* against GERD (**Chap. 347**), Barrett’s esophagus (**Chap. 347**), and adenocarcinoma of the esophagus and gastric cardia (**Chap. 109**). The main lines of evidence for this role are (1) that there is a temporal relationship between a falling prevalence of gastric *H. pylori* colonization and a rising incidence of these conditions; (2) that, in most studies, the prevalence of *H. pylori* colonization (especially with proinflammatory *cagA*⁺ strains) is significantly lower among patients with these esophageal diseases than among control participants; and (3) that, in prospective nested studies (see above), the presence of *H. pylori* is inversely related to these cancers. The mechanism underlying this protective effect is likely *H. pylori*-induced hypochlorhydria. Because, at the individual level, GERD symptoms may decrease, worsen, or remain unchanged after *H. pylori* treatment, concerns about GERD should not affect decisions about whether to treat *H. pylori* when an indication exists.

Other Pathologies *H. pylori* has an increasingly recognized role in other gastric pathologies. It may be one initial precipitant of autoimmune gastritis and pernicious anemia and also may predispose some patients to iron deficiency through occult blood loss and/or hypochlorhydria and reduced iron absorption. In addition, several extragastric pathologies have been linked with *H. pylori* colonization, although evidence of causality is less strong. Studies of *H. pylori* treatment in idiopathic thrombocytopenic purpura have consistently described improvement in or even normalization of platelet counts. Potentially important but even more controversial associations are with ischemic heart disease and cerebrovascular disease. However, the strength of the latter associations is reduced if confounding factors are taken into account, and most authorities consider the associations to be noncausal. Several studies have shown an inverse association of *cagA*⁺ *H. pylori* with childhood-onset asthma, hay fever, and atopic disorders. These associations have been shown to be causal in animal models, but causality in humans and the size of any effect have not been established.

DIAGNOSIS

Tests for *H. pylori* fall into two groups: tests that require upper gastrointestinal endoscopy and simpler tests that can be performed in the clinic (**Table 188-1**).

Endoscopy-Based Tests Endoscopy is usually unnecessary in the initial management of young patients with simple dyspepsia but is commonly used to exclude malignancy and make a positive diagnosis in older patients or those with “alarm” symptoms. If endoscopy is performed, the most convenient biopsy-based test is the biopsy urease test, in which one large or two small gastric biopsy specimens are placed into a gel containing urea and an indicator. The presence of *H. pylori* urease leads to a pH alteration and therefore to a color change, which often occurs within minutes but can require up to 24 h. Histologic examination of biopsy specimens for *H. pylori* also is accurate, provided that a special stain (e.g., a modified Giemsa or silver stain) permitting optimal visualization of the organism is used. If biopsy specimens are obtained from both antrum and corpus, histologic study yields

TABLE 188-1 TESTS COMMONLY USED TO DETECT *HELICOBACTER PYLORI*

Test	Advantages	Disadvantages
Tests Based on Endoscopic Biopsy		
Biopsy urease test	Quick, simple	Some commercial tests not fully sensitive before 24 h
Histology	May give additional histologic information	Sensitivity dependent on experience and use of special stains
Culture	Permits determination of antibiotic susceptibility	Sensitivity dependent on experience
Noninvasive Tests		
Serology	Inexpensive and convenient; not affected by recent antibiotics or proton pump inhibitors to the same extent as breath and stool tests	Cannot be used for early follow-up after treatment; some commercial kits inaccurate, and most less accurate than urea breath test
¹³ C urea breath test	Inexpensive and simpler than endoscopy; useful for follow-up after treatment	Requires fasting; not as convenient as blood or stool tests
Stool antigen test	Inexpensive and convenient; useful for follow-up after treatment; may be useful in children	Stool-based tests are disliked by people from some cultures

additional information, including the degree and pattern of inflammation and the presence of any atrophy, metaplasia, or dysplasia. Microbiologic culture is most specific but may be insensitive because of difficulty with *H. pylori* isolation. Once the organism is cultured, its identity as *H. pylori* can be confirmed by its typical appearance on Gram’s stain and its positive reactions in oxidase, catalase, and urease tests. Moreover, the organism’s susceptibility to antibiotics can be determined, and this information can be clinically useful in difficult cases. The occasional biopsy specimens containing the less common non-*pylori* gastric helicobacters give only weakly positive results in the biopsy urease test. Positive identification of these bacteria requires visualization of the characteristic long, tight spirals in histologic sections; they cannot easily be cultured.

Noninvasive Tests Noninvasive *H. pylori* testing is the norm if gastric cancer does not need to be excluded by endoscopy. The best-established test (and a very accurate one) is the *urea breath test*. In this simple test, the patient drinks a solution of urea labeled with the nonradioactive isotope ¹³C and then blows into a tube. If *H. pylori* urease is present, the urea is hydrolyzed, and labeled carbon dioxide is detected in breath samples. The *stool antigen test*, a simple and accurate test using monoclonal antibodies specific for *H. pylori* antigens, is more convenient and potentially less expensive than the urea breath test, but some patients dislike sampling stool. The simplest tests for ascertaining *H. pylori* status are *serologic assays* measuring specific IgG levels in serum by enzyme-linked immunosorbent assay or immunoblot. The best of these tests are as accurate as other diagnostic methods, but many commercial tests—especially rapid office tests—do not perform well.

Use of Tests to Assess Treatment Success The urea breath test, the stool antigen test, and biopsy-based tests can all be used to assess the success of treatment (**Fig. 188-2**). However, because these tests are dependent on *H. pylori* load, their use <4 weeks after treatment may yield false-negative results. Furthermore, these tests are unreliable if performed within 4 weeks of intercurrent treatment with antibiotics or bismuth compounds or within 2 weeks of the discontinuation of proton pump inhibitor (PPI) treatment. In the assessment of treatment success, noninvasive tests are normally preferred; however, after gastric ulceration, endoscopy should be repeated to ensure healing and to exclude gastric carcinoma by further histologic sampling. Serologic tests are not used to monitor treatment success, as the gradual drop in titer of *H. pylori*-specific antibodies is too slow to be of practical use.

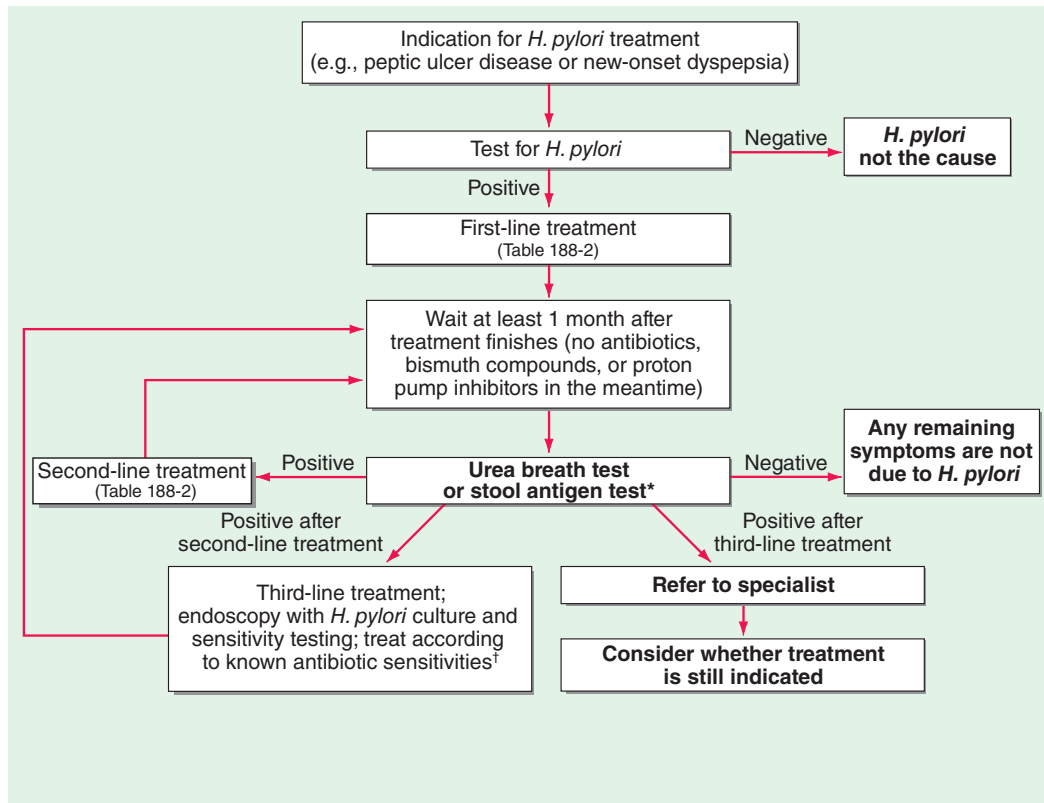


FIGURE 188-2 Algorithm for the management of *Helicobacter pylori* infection. *Note that either the urea breath test or the stool antigen test can be used in this algorithm. Occasionally, endoscopy and a biopsy-based test are used instead of either of these tests in follow-up after treatment. The main indication for these invasive tests is gastric ulceration; in this condition, as opposed to duodenal ulceration, it is important to check healing and to exclude underlying gastric adenocarcinoma. However, even in this situation, patients undergoing endoscopy may still be receiving proton pump inhibitor therapy, which precludes *H. pylori* testing. Thus a urea breath test or a stool antigen test is still required at a suitable interval after the end of therapy to determine whether treatment has been successful (see text). †Some authorities use empirical third-line regimens, of which several have been described.

TREATMENT HELICOBACTER PYLORI INFECTION

INDICATIONS

The most clear-cut indications for treatment are *H. pylori*-related duodenal or gastric ulceration or low-grade gastric B-cell lymphoma. Whether or not the ulcers are currently active, *H. pylori* should be eradicated in patients with documented ulcer disease to prevent relapse (Fig. 188-2). Testing for *H. pylori* and treatment if the results are positive also have been advocated in uninvestigated simple dyspepsia, but only when the prevalence of *H. pylori* in the community is >20% are these measures more cost-effective than simply treating the dyspepsia with PPIs. Guidelines have recommended *H. pylori* treatment in functional dyspepsia in case the patient is one of the perhaps 0–7% who will benefit from such treatment (beyond placebo effects). Some guidelines also recommend treatment of conditions not definitively known to respond to *H. pylori* eradication, including idiopathic thrombocytopenic purpura, vitamin B12 deficiency, and iron-deficiency anemia (in the last instance, only when other causes have been carefully excluded). Test-and-treat has emerged as a common clinical practice in recent years despite the lack of direct evidence that it is advantageous; whether this practice will survive the scrutiny of time and further study remains to be determined. For individuals with a strong family history of gastric cancer, treatment to eradicate *H. pylori* in the hope of reducing their cancer risk is reasonable but of unproven value. Currently, widespread community screening for and treatment of *H. pylori* as primary prophylaxis for gastric cancer and peptic ulcers are not recommended in most countries, mainly because the extent of the consequent reduction in cancer risk is not known. Several studies have found a modestly reduced cancer risk after treatment, but the period of follow-up is still fairly short and the size of the effect in different

populations remains unclear. Other reasons not to treat *H. pylori* in asymptomatic populations at present include (1) the adverse side effects (which are common and can be severe in rare cases) of the multiple-antibiotic regimens used; (2) antibiotic resistance, which may emerge in *H. pylori* or other incidentally carried bacteria; (3) the anxiety that may arise in otherwise healthy people, especially if treatment is unsuccessful; and (4) the existence of a subset of people who will develop GERD symptoms after treatment, although, on average, *H. pylori* treatment does not affect GERD symptoms or severity. Despite the absence of screening strategies, many doctors treat *H. pylori* if it is known to be present (particularly in children and younger adults), even when the patient is asymptomatic. The rationale is that it reduces patient concern and may reduce future gastric cancer risk and that any reduction in risk is likely to be greater in younger patients. However, such practices do not factor in any potential benefits of *H. pylori* colonization. Overall, despite widespread clinical activity in this area, most treatment of asymptomatic *H. pylori* carriage is given without a firm evidence base.

REGIMENS

Although *H. pylori* is susceptible to a wide range of antibiotics in vitro, monotherapy is not usually successful, probably because of inadequate antibiotic delivery to the colonization niche. Failure of monotherapy prompted the development of multidrug regimens, the most successful of which are triple and quadruple combinations. Current regimens consist of a PPI and two or three antimicrobial agents given for 7–14 days (Table 188-2). Research on optimizing drug combinations to increase efficacy continues, and guidelines are likely to change as the field develops and as countries increasingly tailor treatment to suit local antibiotic resistance patterns and economic needs.

TABLE 188-2 COMMONLY RECOMMENDED TREATMENT REGIMENS FOR *HELICOBACTER PYLORI*

Regimen ^a (Duration)	Drug 1	Drug 2	Drug 3	Drug 4
Regimen 1: OCM (7–14 days) ^b	Omeprazole (20 mg bid ^c)	Clarithromycin (500 mg bid)	Metronidazole (500 mg bid)	—
Regimen 2: OCA (7–14 days) ^b	Omeprazole (20 mg bid ^c)	Clarithromycin (500 mg bid)	Amoxicillin (1 g bid)	—
Regimen 3: OBTM (14 days)^d	Omeprazole (20 mg bid ^c)	Bismuth subsalicylate (2 tabs qid)	Tetracycline HCl (500 mg qid)	Metronidazole (500 mg tid)
Regimen 4^e: sequential (5 days + 5 days)	Omeprazole (20 mg bid ^c)	Amoxicillin (1 g bid)	—	—
	Omeprazole (20 mg bid ^c)	Clarithromycin (500 mg bid)	Tinidazole (500 mg bid ^g)	—
Regimen 5^f: concomitant (14 days)	Omeprazole (20 mg bid ^c)	Amoxicillin (1 g bid)	Clarithromycin (500 mg bid)	Tinidazole (500 mg bid ^g)
Regimen 6 ^h : OAL (10 days)	Omeprazole (20 mg bid ^c)	Amoxicillin (1 g bid)	Levofloxacin (500 mg bid)	—

^aThe recommended first-line regimens for most of the world are shown in **bold** type. ^bThese regimens should be used only for populations in which the prevalence of clarithromycin-resistant strains is known to be <20%. In practice, this restriction limits the regimens' appropriate range mainly to northern Europe. Meta-analyses show that a 14-day course of therapy is slightly superior to a 7-day course. ^cMany authorities and some guidelines recommend doubling this dose of omeprazole, as trials show resultant increased efficacy with some antibiotic combinations. Omeprazole may be replaced with any proton pump inhibitor at an equivalent dosage. ^dData supporting this regimen come mainly from Europe and are based on the use of bismuth subcitrate (1 tablet qid) and metronidazole (400 mg tid). This is a recommended first-line regimen in most countries and is the recommended second-line regimen in northern Europe. ^eData supporting this regimen come mainly from Europe. This regimen may be used as an alternative to regimen 3. ^fThis regimen may be used as an alternative to regimen 3 or 4. ^gMetronidazole (500 mg bid) may be used as an alternative. ^hData supporting this regimen come mainly from Europe. It is used as second-line treatment in many countries (particularly where quadruple therapy is used as the first-line regimen) and as third-line treatment in others. This regimen may be less effective where rates of quinolone use are high.



The two most important factors in successful *H. pylori* treatment are the patient's close compliance with the regimen and the use of drugs to which the patient's strain of *H. pylori* has not acquired resistance. Treatment failure following minor lapses in compliance is common and often leads to acquired resistance to metronidazole or clarithromycin. To stress the importance of compliance, written instructions should be given to the patient, and minor side effects of the regimen should be explained. Increasing levels of *H. pylori* resistance to clarithromycin, quinolones, and—to a lesser extent—metronidazole are of growing concern and are thought to be responsible for the reduced efficacy of previously popular clarithromycin-based triple-therapy regimens worldwide. Treatment with these regimens is now virtually confined to certain northern European countries where the use of clarithromycin (or azithromycin) for respiratory infections has not been widespread and resistance rates in *H. pylori* are still low. Strains of *H. pylori* with some degree of in vitro resistance to metronidazole are common but still may be eradicated with metronidazole-containing regimens, which have only slightly reduced efficacy in vivo. Assessment of antibiotic susceptibilities before treatment would be optimal but is not usually undertaken because endoscopy and mucosal biopsy are necessary to obtain *H. pylori* for culture and because most microbiology laboratories are inexperienced in *H. pylori* culture. In the absence of susceptibility information, the patient's history of (even distant) antibiotic use for other conditions should be ascertained; use of the previously administered agent(s) should then be avoided if possible, particularly in the case of clarithromycin (e.g., previous use for upper respiratory infection) and quinolones. If initial *H. pylori* treatment fails, the usual approach is empirical re-treatment with another drug regimen (Table 188-2). The third-line approach should ideally be endoscopy, biopsy, and culture plus treatment based on documented antibiotic sensitivities. However, empirical third-line therapies are often used.

Non-*pylori* gastric helicobacters are treated in the same way as *H. pylori*. However, in the absence of trials, it is unclear whether a positive outcome always represents successful treatment or whether it is sometimes due to natural clearance of the bacteria.

PREVENTION



Carriage of *H. pylori* has considerable public health significance in developed countries, where it is associated with peptic ulcer disease and gastric adenocarcinoma, and in developing countries, where gastric adenocarcinoma may be an even more common cause of cancer death late in life. If mass prevention were contemplated, vaccination would be the most obvious method, and experimental immunization of animals has given promising results. However, given that *H. pylori* has co-evolved with its human host over

millennia, preventing or eliminating colonization on a population basis may have biological and clinical costs. For example, lifelong absence of *H. pylori* is a risk factor for GERD complications, including esophageal adenocarcinoma. We have speculated that the disappearance of *H. pylori* may also be associated with an increased risk of other emergent diseases reflecting aspects of the current Western lifestyle, such as childhood-onset asthma and allergy.

189

Infections Due to *Pseudomonas* Species and Related Organisms

Reuben Ramphal

The pseudomonads are a heterogeneous group of gram-negative bacteria that have in common an inability to ferment lactose. Formerly classified in the genus *Pseudomonas*, the members of this group have been assigned to three medically important genera—*Pseudomonas*, *Burkholderia*, and *Stenotrophomonas*—whose biologic behaviors encompass both similarities and marked differences and whose genetic repertoires differ in many respects. The pathogenicity of most pseudomonads is based on opportunism; the exceptions are the organisms that cause melioidosis (*Burkholderia pseudomallei*) and glanders (*Burkholderia mallei*), which can be considered as primary pathogens.

Pseudomonas aeruginosa, the major pathogen of the group, is a significant cause of infections in hospitalized patients and in patients with cystic fibrosis (CF; **Chap. 313**). Cytotoxic chemotherapy, mechanical ventilation, and broad-spectrum antibiotic therapy probably paved the way for colonization and infection of increasing numbers of hospitalized patients by this organism. Thus most conditions predisposing to *P. aeruginosa* infections have involved host compromise and/or broad-spectrum antibiotic use. The other members of the genus *Pseudomonas*—*Pseudomonas putida*, *Pseudomonas fluorescens*, and *Pseudomonas stutzeri*—infect humans infrequently.

The genus *Burkholderia* comprises more than 40 species, of which *Burkholderia cepacia* is most frequently encountered in Western countries. Like *P. aeruginosa*, *B. cepacia* is both a nosocomial pathogen and a cause of infection in CF. The other medically important members of this genus are *B. pseudomallei* and *B. mallei*, the etiologic agents of melioidosis and glanders, respectively.

The genus *Stenotrophomonas* contains one species of medical significance, *Stenotrophomonas maltophilia* (previously classified in the genera *Pseudomonas* and *Xanthomonas*). This organism is strictly an opportunist that “overgrows” in the setting of potent broad-spectrum antibiotic use.

PSEUDOMONAS AERUGINOSA

EPIDEMIOLOGY

P. aeruginosa is found in most moist environments. Soil, plants, vegetables, tap water, and countertops are all potential reservoirs for this microbe, as it has simple nutritional needs. Given the ubiquity of *P. aeruginosa*, simple contact with the organism is not sufficient for colonization or infection. Clinical and experimental observations suggest that *P. aeruginosa* infection often occurs concomitantly with host defense compromise, mucosal trauma, physiologic derangement, and antibiotic-mediated suppression of normal flora. Thus, it comes as no surprise that the majority of *P. aeruginosa* infections occur in intensive care units (ICUs), where these factors frequently converge. The organism is initially acquired from environmental sources, but patient-to-patient spread also occurs in clinics and families.

In the past, burned patients appeared to be unusually susceptible to *P. aeruginosa*. For example, in 1959–1963, *Pseudomonas* burn-wound sepsis was the principal cause of death in 60% of burned patients dying at the U.S. Army Institute of Surgical Research. For reasons that are unclear, *P. aeruginosa* infection in burns is no longer the major problem that it was during the 1950s and 1960s. Similarly, in the 1960s, *P. aeruginosa* appeared as a common pathogen in patients receiving cytotoxic chemotherapy at many institutions in the United States, but it subsequently diminished in importance. Despite this subsidence, *P. aeruginosa* remains one of the most feared pathogens in this population because of its high attributable mortality rate.



In some parts of Asia and Latin America, *P. aeruginosa* continues to be the most common cause of gram-negative bacteremia in neutropenic patients.

In contrast to the trends for burned patients and neutropenic patients in the United States, the incidence of *P. aeruginosa* infections among patients with CF has not changed. *P. aeruginosa* remains the most common contributing factor to respiratory failure in CF and is responsible for the majority of deaths among CF patients.

LABORATORY FEATURES

P. aeruginosa is a nonfastidious, motile, gram-negative rod that grows on most common laboratory media, including blood and MacConkey agars. It is easily identified in the laboratory on primary-isolation agar plates by pigment production that confers a yellow to dark green or even bluish appearance. Colonies have a shiny “gun-metal” appearance and a characteristic fruity odor. Two of the identifying biochemical characteristics of *P. aeruginosa* are an inability to ferment lactose on MacConkey agar and a positive reaction in the oxidase test. Most strains are identified on the basis of these readily detectable laboratory features even before extensive biochemical testing is done. Some isolates from CF patients are easily identified by their mucoid appearance, which is due to the production of large amounts of the mucoid exopolysaccharide or alginate.

PATHOGENESIS

Unraveling the mechanisms that underlie disease caused by *P. aeruginosa* has proved challenging. Of the common gram-negative bacteria, no other species produces such a large number of putative virulence factors (Table 189-1). Yet *P. aeruginosa* rarely initiates an infectious process in the absence of host injury or compromise, and few of its putative virulence factors have been shown definitively to be involved in disease in humans. Despite its metabolic versatility and possession of multiple colonizing factors, *P. aeruginosa* exhibits no competitive advantage over enteric bacteria in the human gut; neither is it a normal inhabitant of the human gastrointestinal tract, despite the host's continuous environmental exposure to the organism.

Virulence Attributes Involved in Acute *P. aeruginosa* Infections • MOTILITY AND COLONIZATION A general tenet of bacterial pathogenesis is that most bacteria must adhere to surfaces or colonize a host niche in order to initiate disease. Most pathogens examined thus far possess adherence factors called *adhesins*. *P. aeruginosa* is no exception. Among its many adhesins are its pili, which demonstrate adhesive properties for a variety of cells and adhere best to injured cell surfaces. In the organism's flagellum, the

TABLE 189-1 MAIN PUTATIVE VIRULENCE FACTORS OF *PSEUDOMONAS AERUGINOSA*

Substance/Organelle	Function	Virulence in Animal Disease
Pili	Adhesion to cells	?
Flagella	Adhesion, motility, inflammation	Yes
Lipopolysaccharide	Antiphagocytic activity, inflammation	Yes
Type III secretion system	Toxic activity (ExoU, ExoS)	Yes
Type II secretion system	Toxic activity	Yes
Proteases	Proteolytic activity	?
Phospholipases	Cytotoxicity	?
Exotoxin A	Cytotoxicity	?

flagellin molecule binds to cells, and the flagellar cap attaches to mucins through the recognition of glycan chains. Other *P. aeruginosa* adhesins include the outer core of the lipopolysaccharide (LPS) molecule, which binds to the cystic fibrosis transmembrane conductance regulator (CFTR) and aids in internalization of the organism, and the alginate coat of mucoid strains, which enhances adhesion to cells and mucins. In addition, membrane proteins and lectins have been proposed as colonization factors. The deletion of any given adhesin is not sufficient to abrogate the ability of *P. aeruginosa* to colonize surfaces. Motility is important in host invasion via mucosal surfaces in some animal models; however, nonmotile strains are not uniformly avirulent.

EVASION OF HOST DEFENSES The transition from bacterial colonization to disease requires the evasion of host defenses followed by invasion of the microorganism. *P. aeruginosa* appears to be well equipped for evasion. Attached bacteria inject four known toxins (ExoS or ExoU, ExoT, and ExoY) via a type III secretion system that allows the bacteria to evade phagocytic cells either by direct cytotoxicity or by inhibition of phagocytosis. Mutants with defects in this system fail to disseminate in some animal models of infection. The type II secretion system as a whole secretes toxins that can kill animals, and some of its secreted toxins, such as exotoxin A, have the potential to kill phagocytic cells. Multiple proteases secreted by this system may degrade host effector molecules, such as cytokines and chemokines, that are released in response to infection. Thus this system may also contribute to host evasion.

TISSUE INJURY Among gram-negative bacteria, *P. aeruginosa* probably produces the largest number of substances that are toxic to cells and thus may injure tissues. The toxins secreted by its type III secretion system are capable of tissue injury. However, their delivery requires the adherence of the organism to cells. Thus, the effects of these toxins are likely to be local or to depend on the presence of vast numbers of bacteria. On the other hand, diffusible toxins, secreted by the organism's type II secretion system, can act freely wherever they come into contact with cells. Possible effectors include exotoxin A, four different proteases, and at least two phospholipases; in addition to these secreted toxins, rhamnolipids, pyocyanin, and hydrocyanic acid are produced by *P. aeruginosa* and are all capable of inducing host injury.

INFLAMMATORY COMPONENTS The inflammatory components of *P. aeruginosa*—e.g., the inflammatory responses to the lipid A component of LPSs and to flagellin, mediated through the Toll-like receptor (TLR) system (principally TLR4 and TLR5)—have been thought to represent important factors in disease causation. Although these inflammatory responses are required for successful defense against *P. aeruginosa* (i.e., in their absence, animals are defenseless against *P. aeruginosa* infection), florid responses are likely to result in disease. When the sepsis syndrome and septic shock develop in *P. aeruginosa* infection, they are probably the result of the host response to one or both of these substances, but injury to the lung by *Pseudomonas* toxins may also result in sepsis syndromes, possibly by causing cell death and the release of cellular components (e.g., heat-shock proteins) that may activate the TLR or another proinflammatory system.

1044 Chronic *P. aeruginosa* Infections Chronic infection due to *P. aeruginosa* occurs mainly in the lungs in the setting of structural pulmonary diseases. The classic example is CF; others include bronchiectasis and chronic relapsing panbronchiolitis, a disease seen in Japan and some Pacific Islands. Hallmarks of these illnesses are altered mucociliary clearance leading to mucus stasis and mucus accumulation in the lungs. There is probably a common factor that selects for *P. aeruginosa* colonization in these lung diseases—perhaps the adhesiveness of *P. aeruginosa* for mucus, a phenomenon that is not noted for most other common gram-negative bacteria, and/or the ability of *P. aeruginosa* to evade host defenses in mucus. Furthermore, *P. aeruginosa* seems to evolve in ways that allow its prolonged survival in the lung without an early fatal outcome for the host. The strains found in CF patients exhibit minimal production of virulence factors. Some strains even lose the ability to produce pili and flagella, and most become complement-sensitive because of the loss of the O side chain of their LPS molecules. An example of the impact of these changes is found in the organism's discontinuation of the production of flagellin (probably its most strongly proinflammatory molecule) when it encounters purulent mucus. This response probably dampens the host's response, allowing the organism to survive in mucus. *P. aeruginosa* is also believed to lose the ability to secrete many of its injectable toxins during growth in mucus. Although the alginate coat is thought to play a role in the organism's survival, alginate is not essential, as nonmucoid strains may also predominate for long periods. In short, virulence in chronic infections may be mediated by the chronic but attenuated host inflammatory response, which injures the lungs over decades.

CLINICAL MANIFESTATIONS

P. aeruginosa causes infections at almost all sites in the body but shows a rather strong predilection for the lungs. The infections encountered most commonly in hospitalized patients are described below.

Bacteremia Crude mortality rates exceeding 50% have been reported among patients with *P. aeruginosa* bacteremia. Consequently, this clinical entity has been much feared, and its management has been attempted with the use of multiple antibiotics. Recent publications report attributable mortality rates of 28–44%, with the precise figure depending on the adequacy of treatment and the seriousness of the underlying disease. In the past, the patient with *P. aeruginosa* bacteremia classically was neutropenic or had a burn injury. Today, however, a minority of such patients have bacteremic *P. aeruginosa* infections. Rather, *P. aeruginosa* bacteremia is seen most often in patients in ICUs.

The clinical presentation of *P. aeruginosa* bacteremia rarely differs from that of sepsis in general (Chap. 324). Patients are usually febrile, but those who are most severely ill may be in shock or even hypothermic. The only point differentiating this entity from gram-negative sepsis of other causes may be the distinctive skin lesions (ecthyma gangrenosum) of *Pseudomonas* infection, which occur almost exclusively in markedly neutropenic patients and patients with AIDS. These small or large, painful, reddish, maculopapular lesions have a geographic margin; they are initially pink, then darken to purple, and finally become black and necrotic (Fig. 189-1). Histopathologic studies indicate that the lesions are due to vascular invasion and are teeming with bacteria. Although similar lesions may occur in aspergillosis and



FIGURE 189-1 Ecthyma gangrenosum in a neutropenic patient 3 days after onset.

mucormycosis, their presence suggests *P. aeruginosa* bacteremia as the most likely diagnosis.

TREATMENT *P. AERUGINOSA* BACTEREMIA

(Table 189-2) Antimicrobial treatment of *P. aeruginosa* bacteremia has been controversial. Before 1971, the outcome of *Pseudomonas* bacteremia in febrile neutropenic patients treated with the available agents—gentamicin and the polymyxins—was dismal. However, treatment with carbenicillin, with or without an aminoglycoside, significantly improved outcomes. Concurrently, several retrospective analyses suggested that the use of two agents that were synergistic against gram-negative pathogens in vitro resulted in better outcomes in neutropenic patients. Thus, combination therapy became the standard of care—first for *P. aeruginosa* bacteremia in febrile neutropenic patients and then for all *P. aeruginosa* infections in neutropenic or nonneutropenic patients.

With the introduction of newer antipseudomonal drugs, a number of studies have revisited the choice between combination treatment and monotherapy for *Pseudomonas* bacteremia. Although the majority of experts still favor combination therapy, most of these observational studies indicate that a single modern antipseudomonal β -lactam agent to which the isolate is sensitive is as efficacious as a combination. Even in patients at greatest risk of early death from *P. aeruginosa* bacteremia (i.e., those with fever and neutropenia), empirical antipseudomonal monotherapy is deemed to be as efficacious as empirical combination therapy by the practice guidelines of the Infectious Diseases Society of America. One firm conclusion is that monotherapy with an aminoglycoside is not optimal.

There are, of course, institutions and countries where rates of susceptibility of *P. aeruginosa* to first-line antibiotics are <80%. Thus, when a septic patient with a high probability of *P. aeruginosa* infection is encountered in such settings, empirical combination therapy should be administered until the pathogen is identified and susceptibility data become available. Thereafter, whether one or two agents should be continued remains a matter of individual preference. Recent studies suggest that extended infusions of β -lactams such as cefepime or piperacillin-tazobactam may result in better outcomes of *Pseudomonas* bacteremia and possibly *Pseudomonas* pneumonia.

Acute Pneumonia Respiratory infections are the most common of all infections caused by *P. aeruginosa*. This organism appears first or second among the causes of ventilator-associated pneumonia (VAP). However, much debate centers on the actual role of *P. aeruginosa* in VAP. Many of the relevant data are based on cultures of sputum or endotracheal tube aspirates and may represent nonpathogenic colonization of the tracheobronchial tree, biofilms on the endotracheal tube, or simple tracheobronchitis.

Older reports of *P. aeruginosa* pneumonia described patients with an acute clinical syndrome of fever, chills, cough, and necrotizing pneumonia indistinguishable from other gram-negative bacterial pneumonias. The traditional accounts described a fulminant infection. Chest radiographs demonstrated bilateral pneumonia, often with nodular densities with or without cavities. This picture is now remarkably rare. Today, the typical patient is on a ventilator, has a slowly progressive infiltrate, and has been colonized with *P. aeruginosa* for days. While some cases may progress rapidly over 48–72 h, they are the exceptions. Nodular densities are not commonly seen. However, infiltrates may go on to necrosis. Necrotizing pneumonia has also been seen in the community (e.g., after inhalation of hot-tub water contaminated with *P. aeruginosa*). The typical patient has fever, leukocytosis, and purulent sputum, and the chest radiograph shows a new infiltrate or the expansion of a preexisting infiltrate. Chest examination generally detects rales or dullness. Of course, such findings are quite common among ventilated patients in the ICU. A sputum Gram's stain showing mainly polymorphonuclear leukocytes (PMNs) in conjunction with a culture positive for *P. aeruginosa* in this setting suggests a diagnosis of acute *P. aeruginosa* pneumonia. There is no consensus about whether

TABLE 189-2 ANTIBIOTIC TREATMENT OF INFECTIONS DUE TO *PSEUDOMONAS AERUGINOSA* AND RELATED SPECIES

Infection	Antibiotics and Dosages	Other Considerations
Bacteremia		
Nonneutropenic host	Monotherapy: Ceftazidime (2 g q8h IV) or cefepime (2 g q12h IV) Combination therapy: Piperacillin/tazobactam (3.375 g q4h IV) or imipenem (500 mg q6h IV) or meropenem (1 g q8h IV) or doripenem (500 mg q8h IV) plus Amikacin (7.5 mg/kg q12h or 15 mg/kg q24h IV)	Add an aminoglycoside for patients in shock and in regions or hospitals where rates of resistance to the primary β -lactam agents are high. Tobramycin may be used instead of amikacin (susceptibility permitting). The duration of therapy is 7 days for nonneutropenic patients. Neutropenic patients should be treated until no longer neutropenic.
Neutropenic host	Cefepime (2 g q8h IV) or all other agents (except doripenem) in above dosages	
Endocarditis	Antibiotic regimens as for bacteremia for 6–8 weeks	Resistance during therapy is common. Surgery is required for relapse.
Pneumonia	Drugs and dosages as for bacteremia, except that the available carbapenems should not be the sole primary drugs because of high rates of resistance during therapy	IDSA guidelines recommend the addition of an aminoglycoside or ciprofloxacin. The duration of therapy is 10–14 days.
Bone infection, malignant otitis externa	Cefepime or ceftazidime at the same dosages as for bacteremia; aminoglycosides not a necessary component of therapy; ciprofloxacin (500–750 mg q12h PO) may be used	Duration of therapy varies with the drug used (e.g., 6 weeks for a β -lactam agent; at least 3 months for oral therapy except in puncture-wound osteomyelitis, for which the duration should be 2–4 weeks).
Central nervous system infection	Ceftazidime or cefepime (2 g q8h IV) or meropenem (1 g q8h IV)	Abscesses or other closed-space infections may require drainage. The duration of therapy is ≥ 2 weeks.
Eye infection		
Keratitis/ulcer	Topical therapy with tobramycin/ciprofloxacin/levofloxacin eyedrops	Use maximal strengths available or compounded by pharmacy. Therapy should be administered for 2 weeks or until the resolution of eye lesions, whichever is shorter.
Endophthalmitis	Ceftazidime or cefepime as for central nervous system infection plus Topical therapy	
Urinary tract infection	Ciprofloxacin (500 mg q12h PO) or levofloxacin (750 mg q24h) or any aminoglycoside (total daily dose given once daily)	Relapse may occur if an obstruction or a foreign body is present. The duration of therapy for complicated UTI is 7–10 days (up to 2 weeks for pyelonephritis).
Multidrug-resistant <i>P. aeruginosa</i> infection	Colistin (100 mg q12h IV) for the shortest possible period to obtain a clinical response	Doses used have varied. Dosage adjustment is required in renal failure. Inhaled colistin may be added for pneumonia (100 mg q12h).
<i>Stenotrophomonas maltophilia</i> infection	TMP-SMX (1600/320 mg q12h IV) plus ticarcillin/clavulanate (3.1 g q4h IV) for 14 days	Resistance to all agents is increasing. Levofloxacin or tigecycline may be alternatives, but there is little published clinical experience with these agents.
<i>Burkholderia cepacia</i> infection	Meropenem (1 g q8h IV) or TMP-SMX (1600/320 mg q12h IV) for 14 days	Resistance to both agents is increasing. Do not use them in combination because of possible antagonism.
Melioidosis, glanders	Ceftazidime (2 g q6h) or meropenem (1 g q8h) or imipenem (500 mg q6h) for 2 weeks followed by TMP-SMX (1600/320 mg q12h PO) for 3 months	

Abbreviations: IDSA, Infectious Diseases Society of America; TMP-SMX, trimethoprim-sulfamethoxazole.

an invasive procedure (e.g., bronchoalveolar lavage or protected-brush sampling of the distal airways) is superior to tracheal aspiration to obtain samples for lung cultures in order to substantiate the occurrence of *P. aeruginosa* pneumonia and prevent antibiotic overuse.

TREATMENT ACUTE PNEUMONIA

(Table 189-2) Therapy for *P. aeruginosa* pneumonia has been unsatisfactory. Reports suggest mortality rates of 40–80%, but how many of these deaths are attributable to underlying disease remains unknown. The drugs of choice for *P. aeruginosa* pneumonia are similar to those given for bacteremia. A potent antipseudomonal β -lactam drug is the mainstay of therapy. Failure rates were high when aminoglycosides were used as single agents, possibly because of their poor penetration into the airways and their binding to airway secretions. Thus a strong case cannot be made for the inclusion of the aminoglycoside component in regimens used against fully susceptible organisms, especially given the evidence that aminoglycosides are not optimally active in the lungs at concentrations normally reached after IV administration. Nonetheless, aminoglycosides

are commonly used in clinical practice. Some experts suggest the combination of a β -lactam agent and an antipseudomonal fluoroquinolone instead when combination therapy is desired.



Chronic Respiratory Tract Infections *P. aeruginosa* is responsible for chronic infections of the airways associated with a number of underlying or predisposing conditions—most commonly CF (Chap. 313). A state of chronic colonization beginning early in childhood is seen in some Asian populations with chronic or diffuse panbronchiolitis, a disease of unknown etiology. *P. aeruginosa* is one of the organisms that colonizes damaged bronchi in bronchiectasis, a disease secondary to multiple causes in which profound structural abnormalities of the airways result in mucus stasis.

TREATMENT CHRONIC RESPIRATORY TRACT INFECTIONS

Optimal management of chronic *P. aeruginosa* lung infection has not been determined. Patients respond clinically to antipseudomonal therapy, but the organism is rarely eradicated. Because

1046 eradication is unlikely, the aim of treatment for chronic infection is to quell exacerbations of inflammation. The regimens used are similar to those used for pneumonia, but an aminoglycoside is almost always added because resistance is common in chronic disease. However, it may be appropriate to use an inhaled aminoglycoside preparation in order to maximize airway drug levels.

Endovascular Infections Infective endocarditis due to *P. aeruginosa* is a disease of IV drug users whose native valves are involved. This organism has also been reported to cause prosthetic valve endocarditis. Sites of prior native-valve injury due to the injection of foreign material such as talc or fibers probably serve as niduses for bacterial attachment to the heart valve. The manifestations of *P. aeruginosa* endocarditis resemble those of other forms of endocarditis in IV drug users except that the disease is more indolent than *Staphylococcus aureus* endocarditis. While most disease involves the right side of the heart, left-sided involvement is not rare and multivalvular disease is common. Fever is a common manifestation, as is pulmonary involvement (due to septic emboli to the lungs). Hence, patients may also experience chest pain and hemoptysis. Involvement of the left side of the heart may lead to signs of cardiac failure, systemic emboli, and local cardiac involvement with sinus of Valsalva abscesses and conduction defects. Skin manifestations are rare in this disease, and ecthyma gangrenosum is not seen. The diagnosis is based on positive blood cultures along with clinical signs of endocarditis.

TREATMENT ENDOVASCULAR INFECTIONS

(Table 189-2) It has been customary to use synergistic antibiotic combinations in treating *P. aeruginosa* endocarditis because of the development of resistance during therapy with a single antipseudomonal β -lactam agent. Which combination therapy is preferable is unclear, as all combinations have failed. Cases of *P. aeruginosa* endocarditis that relapse during or fail to respond to therapy are often caused by resistant organisms and may require surgical therapy. Other considerations for valve replacement are similar to those in other forms of endocarditis (**Chap. 155**).

Bone and Joint Infections *P. aeruginosa* is an infrequent cause of bone and joint infections. However, *Pseudomonas* bacteremia or infective endocarditis caused by the injection of contaminated illicit drugs has been documented to result in vertebral osteomyelitis and sternoclavicular joint arthritis. The clinical presentation of vertebral *P. aeruginosa* osteomyelitis is more indolent than that of staphylococcal osteomyelitis. The duration of symptoms in IV drug users with vertebral osteomyelitis due to *P. aeruginosa* varies from weeks to months. Fever is not uniformly present; when present, it tends to be low grade. There may be mild tenderness at the site of involvement. Blood cultures are usually negative unless there is concomitant endocarditis. The erythrocyte sedimentation rate (ESR) is generally elevated. Vertebral osteomyelitis due to *P. aeruginosa* has also been reported in the elderly, in whom it originates from urinary tract infections (UTIs). The infection generally involves the lumbosacral area because of a shared venous drainage (Batson's plexus) between the lumbosacral spine and the pelvis. Sternoclavicular septic arthritis due to *P. aeruginosa* is seen almost exclusively in IV drug users. This disease may occur with or without endocarditis, and a primary site of infection often is not found. Plain radiographs show joint or bone involvement. Treatment of these forms of disease is generally successful.

Pseudomonas osteomyelitis of the foot most often follows puncture wounds through sneakers and mostly affects children. The main manifestation is pain in the foot, sometimes with superficial cellulitis around the puncture wound and tenderness on deep palpation of the wound. Multiple joints or bones of the foot may be involved. Systemic symptoms are generally absent, and blood cultures are usually negative. Radiographs may or may not be abnormal, but the bone scan is usually positive, as are magnetic resonance imaging (MRI) studies. Needle

aspiration usually yields a diagnosis. Prompt surgery, with exploration of the nail puncture tract and debridement of the involved bones and cartilage, is generally recommended in addition to antibiotic therapy.

Central Nervous System (CNS) Infections CNS infections due to *P. aeruginosa* are relatively rare. Involvement of the CNS is almost always secondary to a surgical procedure or head trauma. The entity seen most often is postoperative or posttraumatic meningitis. Subdural or epidural infection occasionally results from contamination of these areas. Embolic disease arising from endocarditis in IV drug users and leading to brain abscesses has also been described. The cerebrospinal fluid (CSF) profile of *P. aeruginosa* meningitis is no different from that of pyogenic meningitis of any other etiology.

TREATMENT CENTRAL NERVOUS SYSTEM INFECTIONS

(Table 189-2) Treatment of *Pseudomonas* meningitis is difficult; little information has been published, and no controlled trials in humans have been undertaken. However, the general principles involved in the treatment of meningitis apply, including the need for high doses of bactericidal antibiotics to attain high drug levels in the CSF. The agent with which there is the most published experience in *P. aeruginosa* meningitis is ceftazidime, but other antipseudomonal β -lactam drugs that reach high CSF concentrations, such as cefepime and meropenem, have also been used successfully. Other forms of *P. aeruginosa* CNS infection, such as brain abscesses and epidural and subdural empyema, generally require surgical drainage in addition to antibiotic therapy.

Eye Infections Eye infections due to *P. aeruginosa* occur mainly as a result of direct inoculation into the tissue during trauma or surface injury by contact lenses. Keratitis and corneal ulcers are the most common types of eye disease and are often associated with contact lenses (especially the extended-wear variety). Keratitis can be slowly or rapidly progressive, but the classic description is disease progressing over 48 h to involve the entire cornea, with opacification and sometimes perforation. *P. aeruginosa* keratitis should be considered a medical emergency because of the rapidity with which it can progress to loss of sight. *P. aeruginosa* endophthalmitis secondary to bacteremia is the most devastating of *P. aeruginosa* eye infections. The disease is fulminant, with severe pain, chemosis, decreased visual acuity, anterior uveitis, vitreous involvement, and panophthalmitis.

TREATMENT EYE INFECTIONS

(Table 189-2) The usual therapy for keratitis is the administration of topical antibiotics. Therapy for endophthalmitis includes the use of high-dose local and systemic antibiotics (to achieve higher drug concentrations in the eye) and vitrectomy.

Ear Infections *P. aeruginosa* infections of the ears vary from mild swimmer's ear to serious life-threatening infections with neurologic sequelae. Swimmer's ear is common among children and results from infection of moist macerated skin of the external ear canal. Most cases resolve with treatment, but some patients develop chronic drainage. Swimmer's ear is managed with topical antibiotic agents (otic solutions). The most serious form of *Pseudomonas* infection involving the ear has been given various names: two of these designations, *malignant otitis externa* and *necrotizing otitis externa*, are now used for the same entity. This disease was originally described in elderly diabetic patients, in whom the majority of cases still occur. However, it has also been described in patients with AIDS and in elderly patients without underlying diabetes or immunocompromise. The usual presenting symptoms are decreased hearing and ear pain, which may be severe and lancinating. The pinna is usually painful, and the external canal may be tender. The ear canal almost always shows signs of inflammation, with granulation tissue and exudate. Tenderness anterior to the tragus may

extend as far as the temporomandibular joint and mastoid process. A small minority of patients have systemic symptoms. Patients in whom the diagnosis is made late may present with cranial nerve palsies or even with cavernous venous sinus thrombosis. The ESR is invariably elevated (≥ 100 mm/h). The diagnosis is made on clinical grounds in severe cases; however, the “gold standard” is a positive technetium-99 bone scan in a patient with otitis externa due to *P. aeruginosa*. In diabetic patients, a positive bone scan constitutes presumptive evidence for this diagnosis and should prompt biopsy or empirical therapy.

TREATMENT EAR INFECTIONS

(Table 189-2) Given the infection of the ear cartilage, sometimes with mastoid or petrous ridge involvement, patients with malignant (necrotizing) otitis externa are treated as for osteomyelitis.

Urinary Tract Infections UTIs due to *P. aeruginosa* generally occur as a complication of a foreign body in the urinary tract, an obstruction in the genitourinary system, or urinary tract instrumentation or surgery. However, UTIs caused by *P. aeruginosa* have been described in pediatric outpatients without stones or evident obstruction.

TREATMENT URINARY TRACT INFECTIONS

(Table 189-2) Most *P. aeruginosa* UTIs are considered complicated infections that must be treated longer than uncomplicated cystitis. In general, a 7- to 10-day course of treatment suffices, with up to 2 weeks of therapy in cases of pyelonephritis. Urinary catheters, stents, or stones should be removed to prevent relapse, which is common and may be due not to resistance but rather to factors such as a foreign body that has been left in place or an ongoing obstruction.

Skin and Soft Tissue Infections Besides pyoderma gangrenosum in neutropenic patients, folliculitis and other papular or vesicular lesions due to *P. aeruginosa* have been extensively described and are collectively referred to as *dermatitis*. Multiple outbreaks have been linked to whirlpools, spas, and swimming pools. To prevent such outbreaks, the growth of *P. aeruginosa* in the home and in recreational environments must be controlled by proper chlorination of water. Most cases of hot-tub folliculitis are self-limited, requiring only the avoidance of exposure to the contaminated source of water.



Toe-web infections occur especially often in the tropics, and the “green nail syndrome” is caused by *P. aeruginosa* paronychia, which results from frequent submersion of the hands in water. In the latter entity, the green discoloration results from diffusion of pyocyanin into the nail bed. *P. aeruginosa* remains a prominent cause of burn wound infections in some parts of the world. The management of these infections is best left to specialists in burn wound care.



Infections in Febrile Neutropenic Patients In febrile neutropenia, *P. aeruginosa* has historically been the organism against which empirical coverage is always essential. Although in Western countries these infections are now less common, their importance has not diminished because of persistently high mortality rates. In other parts of the world as well, *P. aeruginosa* continues to be a significant problem in febrile neutropenia, causing a larger proportion of infections in febrile neutropenic patients than any other single organism. For example, *P. aeruginosa* was responsible for 28% of documented infections in 499 febrile neutropenic patients in one study from the Indian subcontinent and for 31% of such infections in another. In a large study of infections in leukemia patients from Japan, *P. aeruginosa* was the most frequently documented cause of bacterial infection. In studies performed in North America, northern Europe, and Australia, the incidence of *P. aeruginosa* bacteremia in febrile neutropenia was quite variable. In a review of 97 reports published in 1987–1994, the incidence was reported to be 1–2.5% among febrile neutropenic patients given empirical therapy and 5–12% among

microbiologically documented infections. The most common clinical syndromes encountered were bacteremia, pneumonia, and soft tissue infections manifesting mainly as ecthyma gangrenosum.

TREATMENT INFECTIONS IN FEBRILE NEUTROPENIC PATIENTS

(Table 189-2) Compared with rates three decades ago, improved rates of response to antibiotic therapy have been reported in many studies. A study of 127 patients demonstrated a reduction in the mortality rate from 71% to 25% with the introduction of ceftazidime and imipenem. Because neutrophils—the normal host defenses against this organism—are absent in febrile neutropenic patients, maximal doses of antipseudomonal β -lactam antibiotics should be used for the management of *P. aeruginosa* bacteremia in this setting.

Infections in Patients with AIDS Both community- and hospital-acquired *P. aeruginosa* infections were documented in patients with AIDS before the advent of antiretroviral therapy. Since the introduction of protease inhibitors, *P. aeruginosa* infections in AIDS patients have been seen less frequently but still occur, particularly in the form of sinusitis. The clinical presentation of *Pseudomonas* infection (especially pneumonia and bacteremia) in AIDS patients is remarkable in that, although the illness may appear not to be severe, the infection may nonetheless be fatal. Patients with bacteremia may have only a low-grade fever and may present with ecthyma gangrenosum. Pneumonia, with or without bacteremia, is perhaps the most common type of *P. aeruginosa* infection in AIDS patients. Patients with AIDS and *P. aeruginosa* pneumonia exhibit the classic clinical signs and symptoms of pneumonia, such as fever, productive cough, and chest pain. The infection may be lobar or multilobar and shows no predisposition for any particular location. The most striking feature is the high frequency of cavitory disease.

TREATMENT INFECTIONS IN PATIENTS WITH AIDS

Therapy for any of these conditions in AIDS patients is no different from that in other patients. However, relapse is the rule unless the patient's CD4+ T cell count rises to $>50/\mu\text{L}$ or suppressive antibiotic therapy is given. In attempts to achieve cures and prevent relapses, therapy tends to be more prolonged than in the case of an immunocompetent patient.



Multidrug-Resistant Infections (Table 189-2) *P. aeruginosa* has a notorious propensity to develop antibiotic resistance. During three decades, the impact of resistance was minimized by the rapid development of potent antipseudomonal agents. However, the situation has recently changed, with the worldwide selection of strains carrying determinants that mediate resistance to β -lactams, fluoroquinolones, and aminoglycosides. This situation has been compounded by the lack of development of new classes of antipseudomonal drugs for nearly two decades. Physicians now resort to drugs such as colistin and polymyxin, which were discarded decades ago. These alternative approaches to the management of multiresistant *P. aeruginosa* infections were first used some time ago in CF patients, who receive colistin (polymyxin E) IV and by aerosol despite its renal toxicity. Colistin is rapidly becoming the last-resort agent of choice, even in non-CF patients infected with multiresistant *P. aeruginosa*.

The clinical outcome of multidrug-resistant *P. aeruginosa* infections treated with colistin is difficult to judge from case reports, especially given the many drugs used in the complicated management of these patients. Although earlier reports described marginal efficacy and serious nephrotoxicity and neurotoxicity, recent reports have been more encouraging. Because colistin shows synergy with other antimicrobial agents *in vitro*, it may be possible to reduce the dosage—and thus the toxicity—of this drug when it is combined with drugs such as rifampin

OTHER PSEUDOMONADS

STENOTROPHOMONAS MALTOPHILIA

S. maltophilia is the only potential human pathogen among a genus of ubiquitous organisms found in the rhizosphere (i.e., the soil that surrounds the roots of plants). The organism is an opportunist that is acquired from the environment but is even more limited than *P. aeruginosa* in its ability to colonize patients or cause infections. Immunocompromise is not sufficient to permit these events; rather, major perturbations of the human flora are usually necessary for the establishment of *S. maltophilia*. Accordingly, most cases of human infection occur in the setting of very broad-spectrum antibiotic therapy with agents such as advanced cephalosporins and carbapenems, which eradicate the normal flora and other pathogens. The remarkable ability of *S. maltophilia* to resist virtually all classes of antibiotics is attributable to the possession of antibiotic efflux pumps and of two β -lactamases (L1 and L2) that mediate β -lactam resistance, including that to carbapenems. It is fortunate that the virulence of *S. maltophilia* appears to be limited. Although a serine protease is present in some strains, virulence is probably a result of the host's inflammatory response to components of the organism such as LPS and flagellin. *S. maltophilia* is most commonly found in the respiratory tract of ventilated patients, where the distinction between its roles as a colonizer and as a pathogen is often difficult to make. However, *S. maltophilia* does cause pneumonia and bacteremia in such patients, and these infections have led to septic shock. Also common is central venous line-associated infection (with or without bacteremia), which has been reported most often in patients with cancer. *S. maltophilia* is a rare cause of ecthyma gangrenosum in neutropenic patients. It has been isolated from ~5% of CF patients but is not believed to be a significant pathogen in this setting.

TREATMENT *S. MALTOPHILIA* INFECTIONS

The intrinsic resistance of *S. maltophilia* to most antibiotics renders infection difficult to treat. The antibiotics to which it is most often (although not uniformly) susceptible are trimethoprim-sulfamethoxazole (TMP-SMX), ticarcillin/clavulanate, levofloxacin, and tigecycline (Table 189-2). Consequently, a combination of TMP-SMX and ticarcillin/clavulanate is recommended for initial therapy. Catheters must be removed in the treatment of bacteremia to hasten cure and prevent relapses. The treatment of VAP due to *S. maltophilia* is much more difficult than that of bacteremia, with the frequent development of resistance during therapy.

BURKHOLDERIA CEPACIA

B. cepacia gained notoriety as the cause of a rapidly fatal syndrome of respiratory distress and septicemia (the "cepacia syndrome") in CF patients. Previously, it had been recognized as an antibiotic-resistant nosocomial pathogen (then designated *P. cepacia*) in ICU patients. Patients with chronic granulomatous disease are also predisposed to *B. cepacia* lung disease. The organism has been reclassified into nine subgroups, only some of which are common in CF. *B. cepacia* is an environmental organism that inhabits moist environments and is found in the rhizosphere. This organism possesses multiple virulence factors that may play roles in disease as well as colonizing factors that are capable of binding to lung mucus—an ability that may explain the predilection of *B. cepacia* for the lungs in CF. *B. cepacia* secretes elastase and possesses components of an injectable toxin-secretion system like that of *P. aeruginosa*; its LPS is among the most potent of all LPSs in stimulating an inflammatory response in the lungs. Inflammation may be the major cause of the lung disease seen in the cepacia syndrome. The organism can penetrate epithelial surfaces by virtue of motility and inhibition of host innate immune defenses. Besides infecting the

lungs in CF, *B. cepacia* appears as an airway colonizer during broad-spectrum antibiotic therapy and is a cause of VAP, catheter-associated infections, and wound infections.

TREATMENT *B. CEPACIA* INFECTIONS

B. cepacia is intrinsically resistant to many antibiotics. Therefore, treatment must be tailored according to sensitivities. TMP-SMX, meropenem, and doxycycline are the most effective agents in vitro and may be started as first-line agents (Table 189-2). Some strains are susceptible to third-generation cephalosporins and fluoroquinolones, and these agents may be used against isolates known to be susceptible. Combination therapy for serious pulmonary infection (e.g., in CF) is suggested when multidrug-resistant strains are implicated; the combination of meropenem and TMP-SMX may be antagonistic, however. Resistance to all agents used has been reported during therapy.

BURKHOLDERIA PSEUDOMALLEI



B. pseudomallei is the causative agent of melioidosis, a disease of humans and animals that is geographically restricted to Southeast Asia and northern Australia, with occasional cases in countries such as India and China. This organism may be isolated from individuals returning directly from these endemic regions and from military personnel who have served in endemic regions and then returned home after stops in Europe. Symptoms of this illness may develop only at a later date because of the organism's ability to cause latent infections. *B. pseudomallei* is found in soil and water. Humans and animals are infected by inoculation, inhalation, or ingestion; only rarely is the organism transmitted from person to person. Humans are not colonized without being infected. Among the pseudomonads, *B. pseudomallei* is perhaps the most virulent. Host compromise is not an essential prerequisite for disease, although many patients have common underlying medical diseases (e.g., diabetes or renal failure). *B. pseudomallei* is a facultative intracellular organism whose replication in PMNs and macrophages may be aided by the possession of a polysaccharide capsule. The organism also possesses elements of a type III secretion system that plays a role in its intracellular survival. During infection, there is a florid inflammatory response whose role in disease is unclear.

B. pseudomallei causes a wide spectrum of disease, ranging from asymptomatic infection to abscesses, pneumonia, and disseminated disease. It is a significant cause of fatal community-acquired pneumonia and septicemia in endemic areas, with mortality rates as high as 44% reported in Thailand. Acute pulmonary infection is the most commonly diagnosed form of melioidosis. Pneumonia may be asymptomatic (with routine chest radiographs showing mainly upper-lobe infiltrates) or may present as severe necrotizing disease. *B. pseudomallei* also causes chronic pulmonary infections with systemic manifestations that mimic those of tuberculosis, including chronic cough, fever, hemoptysis, night sweats, and cavitary lung disease. Besides pneumonia, the other principal form of *B. pseudomallei* disease is skin ulceration with associated lymphangitis and regional lymphadenopathy. Spread from the lungs or skin, which is most often documented in debilitated individuals, gives rise to septicemic forms of melioidosis that carry a high mortality rate.

TREATMENT *B. PSEUDOMALLEI* INFECTIONS

B. pseudomallei is susceptible to advanced penicillins and cephalosporins and to carbapenems (Table 189-2). Treatment is divided into two stages: an intensive 2-week phase of therapy with ceftazidime or a carbapenem followed by at least 12 weeks of oral TMP-SMX to eradicate the organism and prevent relapse. The recognition of this bacterium as a potential agent of biologic warfare has stimulated interest in the development of a vaccine.

BURKHOLDERIA MALLEI

B. mallei causes the equine disease glanders in Africa, Asia, and South America. The organism was eradicated from Europe and North America decades ago. The last case seen in the United States occurred in 2001 in a laboratory worker; before that, *B. mallei* had last been seen in this country in 1949. In contrast to the other organisms discussed in this chapter, *B. mallei* is not an environmental organism and does not persist outside its equine hosts. Consequently, *B. mallei* infection is an occupational risk for handlers of horses, equine butchers, and veterinarians in areas of the world where it still exists. The polysaccharide capsule is a critical virulence determinant; diabetics are thought to be especially susceptible to infection by this organism. The organism is transmitted from animals to humans by inoculation into the skin, where it causes local infection with nodules and lymphadenitis. Regional lymphadenopathy is common. Respiratory secretions from infected horses are extremely infectious. Inhalation results in clinical signs of typical pneumonia but may also cause an acute febrile illness with ulceration of the trachea. The organism may disseminate from the skin or lungs to cause septicemia with signs of sepsis. The septicemic form is frequently associated with shock and a high mortality rate. The infection may also enter a chronic phase and present as disseminated abscesses. *B. mallei* infection may present as early as 1–2 days after inhalation or (in cutaneous disease) may not become evident for months.

TREATMENT B. MALLEI INFECTIONS

The antibiotic susceptibility pattern of *B. mallei* is similar to that of *B. pseudomallei*; in addition, the organism is susceptible to the newer macrolides azithromycin and clarithromycin. *B. mallei* infection should be treated with the same drugs and for the same duration as melioidosis.

190 Salmonellosis

David A. Pegues, Samuel I. Miller

Bacteria of the genus *Salmonella* are highly adapted for growth in both humans and animals and cause a wide spectrum of disease. The growth of serotypes *Salmonella typhi* and *Salmonella paratyphi* is restricted to human hosts, in whom these organisms cause enteric (typhoid) fever. The remaining serotypes (nontyphoidal *Salmonella*, or NTS) can colonize the gastrointestinal tracts of a broad range of animals, including mammals, reptiles, birds, and insects. More than 200 serotypes of *Salmonella* are pathogenic to humans, in whom they often cause gastroenteritis and can be associated with localized infections and/or bacteremia.

ETIOLOGY

This large genus of gram-negative bacilli within the family Enterobacteriaceae consists of two species: *Salmonella enterica*, which contains six subspecies, and *Salmonella bongori*. *S. enterica* subspecies I includes almost all the serotypes pathogenic for humans. Members of the seven *Salmonella* subspecies are classified into >2500 serotypes (serovars); for simplicity, *Salmonella* serotypes (most of which are named for the city where they were identified) are often used as the species designation. For example, the full taxonomic designation *S. enterica* subspecies *enterica* serotype Typhimurium can be shortened to *Salmonella* serotype Typhimurium or simply *S. typhimurium*. Serotyping is based on the somatic O antigen (lipopolysaccharide cell-wall components), the surface Vi antigen (restricted to *S. typhi* and *S. paratyphi* C), and the flagellar H antigen.

Salmonellae are gram-negative, non-spore-forming, facultatively anaerobic bacilli that measure 2–3 μm by 0.4–0.6 μm . The initial

identification of salmonellae in the clinical microbiology laboratory is based on growth characteristics. Salmonellae, like other Enterobacteriaceae, produce acid on glucose fermentation, reduce nitrates, and do not produce cytochrome oxidase. In addition, all salmonellae except *Salmonella gallinarum-pullorum* are motile by means of peritrichous flagella, and all but *S. typhi* produce gas (H_2S) on sugar fermentation. Notably, only 1% of clinical isolates ferment lactose; a high level of suspicion must be maintained to detect these rare clinical lactose-fermenting isolates.

Although serotyping of all surface antigens can be used for formal identification, most laboratories perform a few simple agglutination reactions that define specific O-antigen serogroups, designated A, B, C₁, C₂, D, and E. Strains in these six serogroups cause ~99% of *Salmonella* infections in humans and other warm-blooded animals. Molecular typing methods, including pulsed-field gel electrophoresis, polymerase chain reaction fingerprinting, and genomic DNA microarray analysis, are used in epidemiologic investigations to differentiate *Salmonella* strains of a common serotype.

PATHOGENESIS

All *Salmonella* infections begin with ingestion of organisms, most commonly in contaminated food or water. The infectious dose ranges from 200 colony-forming units (CFU) to 10^6 CFU, and the ingested dose is an important determinant of incubation period and disease severity. Conditions that decrease either stomach acidity (an age of <1 year, antacid ingestion, or achlorhydric disease) or intestinal integrity (inflammatory bowel disease, prior gastrointestinal surgery, or alteration of the intestinal flora by antibiotic administration) increase susceptibility to *Salmonella* infection.

Once *S. typhi* and *S. paratyphi* reach the small intestine, they penetrate the mucus layer of the gut and traverse the intestinal layer through phagocytic microfold (M) cells that reside within Peyer's patches. Salmonellae can trigger the formation of membrane ruffles in normally nonphagocytic epithelial cells. These ruffles reach out and enclose adherent bacteria within large vesicles by *bacterial-mediated endocytosis*. This process is dependent on the direct delivery of *Salmonella* proteins into the cytoplasm of epithelial cells by the specialized bacterial type III secretion system. These bacterial proteins mediate alterations in the actin cytoskeleton that are required for *Salmonella* uptake.

After crossing the epithelial layer of the small intestine, *S. typhi* and *S. paratyphi*, which cause enteric (typhoid) fever, are phagocytosed by macrophages. These salmonellae survive the antimicrobial environment of the macrophage by sensing environmental signals that trigger alterations in regulatory systems of the phagocytosed bacteria. For example, PhoP/PhoQ (the best-characterized regulatory system) triggers the expression of outer-membrane proteins and mediates modifications in lipopolysaccharide so that the altered bacterial surface can resist microbicidal activities and potentially alter host cell signaling. In addition, salmonellae encode a second type III secretion system that directly delivers bacterial proteins across the phagosome membrane into the macrophage cytoplasm. This secretion system functions to remodel the *Salmonella*-containing vacuole, promoting bacterial survival and replication.

Once phagocytosed, typhoidal salmonellae disseminate throughout the body in macrophages via the lymphatics and colonize reticuloendothelial tissues (liver, spleen, lymph nodes, and bone marrow). Patients have relatively few or no signs and symptoms during this initial incubation stage. Signs and symptoms, including fever and abdominal pain, probably result from secretion of cytokines by macrophages and epithelial cells in response to bacterial products that are recognized by innate immune receptors when a critical number of organisms have replicated. Over time, the development of hepatosplenomegaly is likely to be related to the recruitment of mononuclear cells and the development of a specific acquired cell-mediated immune response to *S. typhi* colonization. The recruitment of additional mononuclear cells and lymphocytes to Peyer's patches during the several weeks after initial colonization/infection can result in marked enlargement and necrosis

1050 of the Peyer's patches, which may be mediated by bacterial products that promote cell death as well as the inflammatory response.

In contrast to enteric fever, which is characterized by an infiltration of mononuclear cells into the small-bowel mucosa, NTS gastroenteritis is characterized by massive polymorphonuclear leukocyte infiltration into both the large- and small-bowel mucosa. This response appears to depend on the induction of interleukin 8, a strong neutrophil chemotactic factor, which is secreted by intestinal cells as a result of *Salmonella* colonization and translocation of bacterial proteins into host cell cytoplasm. The degranulation and release of toxic substances by neutrophils may result in damage to the intestinal mucosa, causing the inflammatory diarrhea observed with nontyphoidal gastroenteritis. An additional important factor in the persistence of nontyphoidal salmonellae in the intestinal tract and the organisms' capacity to compete with endogenous flora is the ability to utilize the sulfur-containing compound tetrathionate for metabolism in a microaerophilic environment. In the presence of intestinal inflammation, tetrathionate is generated from thiosulfate produced by epithelial cells through inflammatory cell production of reactive oxygen species.

ENTERIC (TYPHOID) FEVER

Enteric (typhoid) fever is a systemic disease characterized by fever and abdominal pain and caused by dissemination of *S. typhi* or *S. paratyphi*. The disease was initially called *typhoid fever* because of its clinical similarity to typhus. In the early 1800s, typhoid fever was clearly defined pathologically as a unique illness on the basis of its association with enlarged Peyer's patches and mesenteric lymph nodes. In 1869, given the anatomic site of infection, the term *enteric fever* was proposed as an alternative designation to distinguish typhoid fever from typhus. However, to this day, the two designations are used interchangeably.

EPIDEMIOLOGY

In contrast to other *Salmonella* serotypes, the etiologic agents of enteric fever—*S. typhi* and *S. paratyphi* serotypes A, B, and C—have no known hosts other than humans. Most commonly, food-borne or waterborne transmission results from fecal contamination by ill or asymptomatic chronic carriers. Sexual transmission between male partners has been described. Health care workers occasionally acquire enteric fever after exposure to infected patients or during processing of clinical specimens and cultures.

With improvements in food handling and water/sewage treatment, enteric fever has become rare in developed nations. Worldwide, however, there are an estimated 27 million cases of enteric fever, with 200,000–600,000 deaths annually. The annual incidence is highest (>100 cases/100,000 population) in south-central and Southeast Asia; medium (10–100 cases/100,000) in the rest of Asia, Africa, Latin America, and Oceania (excluding Australia and New Zealand); and low in other parts of the world (Fig. 190-1). A high incidence of enteric fever correlates with poor sanitation and lack of access to clean drinking water. In endemic regions, enteric fever is more common in urban than rural areas and among young children and adolescents than among other age groups. Risk factors include contaminated water or ice, flooding, food and drinks purchased from street vendors, raw fruits and vegetables grown in fields fertilized with sewage, ill household contacts, lack of hand washing and toilet access, and evidence of prior *Helicobacter pylori* infection (an association probably related to chronically reduced gastric acidity). It is estimated that there is one case of paratyphoid fever for every four cases of typhoid fever, but the incidence of infection associated with *S. paratyphi* A appears to be

increasing, especially in India; this increase may be a result of vaccination for *S. typhi*.

Multidrug-resistant (MDR) strains of *S. typhi* emerged in the 1980s in China and Southeast Asia and have since disseminated widely. These strains contain plasmids encoding resistance to chloramphenicol, ampicillin, and trimethoprim—antibiotics long used to treat enteric fever. With the increased use of fluoroquinolones to treat MDR enteric fever in the 1990s, strains of *S. typhi* and *S. paratyphi* with decreased ciprofloxacin susceptibility (DCS; minimal inhibitory concentration [MIC], 0.125–0.5 µg/mL) or ciprofloxacin resistance (MIC, ≥1 µg/mL) have emerged on the Indian subcontinent, in southern Asia, and (most recently) in sub-Saharan Africa and have been associated with clinical treatment failure. Testing of isolates for resistance to the first-generation quinolone nalidixic acid detects many but not all strains with reduced susceptibility to ciprofloxacin and is no longer recommended. Strains of *S. typhi* and *S. paratyphi* producing extended-spectrum β-lactamases have emerged recently, primarily in India and Nepal.

Approximately 300 cases of typhoid and 150 cases of paratyphoid fever are reported annually in the United States. Of 1902 cases of *S. typhi*-associated enteric fever reported to the Centers for Disease Control and Prevention in 1999–2006, 79% were associated with recent international travel, most commonly to India (47%), Pakistan (10%), Bangladesh (10%), Mexico (7%), and the Philippines (4%). Only 5% of travelers diagnosed with enteric fever had received *S. typhi* vaccine. Overall, 13% of *S. typhi* isolates in the United States were resistant to ampicillin, chloramphenicol, and trimethoprim-sulfamethoxazole (TMP-SMX), and the proportion of DCS isolates increased from 19% in 1999 to 58% in 2006. Infection with DCS *S. typhi* was associated with travel to the Indian subcontinent. Of the 25–30% of reported cases of enteric fever in the United States that are domestically acquired, the majority are sporadic, but outbreaks linked to contaminated food products and previously unrecognized chronic carriers continue to occur.

CLINICAL COURSE

Enteric fever is a misnomer, in that the hallmark features of this disease—fever and abdominal pain—are variable. While fever is documented at presentation in >75% of cases, abdominal pain is reported in only 30–40%. Thus, a high index of suspicion for this potentially fatal systemic illness is necessary when a person presents with fever and a history of recent travel to a developing country.

The incubation period for *S. typhi* averages 10–14 days but ranges from 5 to 21 days, depending on the inoculum size and the host's health and immune status. The most prominent symptom is prolonged fever (38.8°–40.5°C; 101.8°–104.9°F), which can continue for



■ High (>100/100,000/year) ■ Medium (10–100/100,000/year) ■ Low (<10/100,000/year)

FIGURE 190-1 Annual incidence of typhoid fever per 100,000 population. (Adapted from JA Crump et al: The global burden of typhoid fever. *Bull World Health Organ* 82:346, 2004.)



FIGURE 190-2 “Rose spots,” the rash of enteric fever due to *Salmonella typhi* or *Salmonella paratyphi*.

up to 4 weeks if untreated. *S. paratyphi* A is thought to cause milder disease than *S. typhi*, with predominantly gastrointestinal symptoms. However, a prospective study of 669 consecutive cases of enteric fever in Kathmandu, Nepal, found that the infections caused by these organisms were clinically indistinguishable. In this series, symptoms reported on initial medical evaluation included headache (80%), chills (35–45%), cough (30%), sweating (20–25%), myalgias (20%), malaise (10%), and arthralgia (2–4%). Gastrointestinal manifestations included anorexia (55%), abdominal pain (30–40%), nausea (18–24%), vomiting (18%), and diarrhea (22–28%) more commonly than constipation (13–16%). Physical findings included coated tongue (51–56%), splenomegaly (5–6%), and abdominal tenderness (4–5%).

Early physical findings of enteric fever include rash (“rose spots”; 30%), hepatosplenomegaly (3–6%), epistaxis, and relative bradycardia at the peak of high fever (<50%). Rose spots (Fig. 190-2; see also Fig. 25e-9) make up a faint, salmon-colored, blanching, maculopapular rash located primarily on the trunk and chest. The rash is evident in ~30% of patients at the end of the first week and resolves without a trace after 2–5 days. Patients can have two or three crops of lesions, and *Salmonella* can be cultured from punch biopsies of these lesions. The faintness of the rash makes it difficult to detect in highly pigmented patients.

The development of severe disease (which occurs in ~10–15% of patients) depends on host factors (immunosuppression, antacid therapy, previous exposure, and vaccination), strain virulence and inoculum, and choice of antibiotic therapy. Gastrointestinal bleeding (10–20%) and intestinal perforation (1–3%) most commonly occur in the third and fourth weeks of illness and result from hyperplasia, ulceration, and necrosis of the ileocecal Peyer’s patches at the initial site of *Salmonella* infiltration (Fig. 190-3). Both complications are life-threatening and require immediate fluid resuscitation and surgical intervention, with broadened antibiotic coverage for polymicrobial peritonitis (Chap. 159) and treatment of gastrointestinal hemorrhages, including bowel resection. Neurologic manifestations occur in 2–40% of patients and include meningitis, Guillain-Barré syndrome, neuritis, and neuropsychiatric symptoms (described as “muttering delirium” or “coma vigil”), with picking at bedclothes or imaginary objects.

Rare complications whose incidences are reduced by prompt antibiotic treatment include disseminated intravascular coagulation, hematophagocytic syndrome, pancreatitis, hepatic and splenic abscesses and granulomas, endocarditis, pericarditis, myocarditis, orchitis, hepatitis, glomerulonephritis, pyelonephritis and hemolytic-uremic syndrome, severe pneumonia, arthritis, osteomyelitis, endophthalmitis, and parotitis. Up to 10% of patients develop mild relapse, usually within 2–3 weeks of fever resolution and in association with the same strain type and susceptibility profile.

Up to 10% of untreated patients with typhoid fever excrete *S. typhi* in the feces for up to 3 months, and 1–4% develop chronic



FIGURE 190-3 Typical ileal perforation associated with *Salmonella typhi* infection. (From JM Saxe, R Cropsey: Is operative management effective in treatment of perforated typhoid? Am J Surg 189:342, 2005.)


asymptomatic carriage, shedding *S. typhi* in either urine or stool for >1 year. Chronic carriage is more common among women, infants, and persons who have biliary abnormalities or concurrent bladder infection with *Schistosoma haematobium*. The anatomic abnormalities associated with the latter conditions presumably allow prolonged colonization.

DIAGNOSIS

Because the clinical presentation of enteric fever is relatively nonspecific, the diagnosis needs to be considered in any febrile traveler returning from a developing region, especially the Indian subcontinent, the Philippines, or Latin America. Other diagnoses that should be considered in these travelers include malaria, hepatitis, bacterial enteritis, dengue fever, rickettsial infections, leptospirosis, amebic liver abscesses, and acute HIV infection (Chap. 149). Other than a positive culture, no specific laboratory test is diagnostic for enteric fever. In 15–25% of cases, leukopenia and neutropenia are detectable. Leukocytosis is more common among children, during the first 10 days of illness, and in cases complicated by intestinal perforation or secondary infection. Other nonspecific laboratory findings include moderately elevated values in liver function tests and muscle enzyme levels.

The definitive diagnosis of enteric fever requires the isolation of *S. typhi* or *S. paratyphi* from blood, bone marrow, other sterile sites, rose spots, stool, or intestinal secretions. The sensitivity of blood culture is only 40–80%, probably because of high rates of antibiotic use in endemic areas and the small number of *S. typhi* organisms (i.e., <15/mL) typically present in the blood. Because almost all *S. typhi* organisms in blood are associated with the mononuclear cell/platelet fraction, centrifugation of blood and culture of the buffy coat can substantially reduce the time to isolation of the organism but do not increase sensitivity.

Bone marrow culture is 55–90% sensitive, and, unlike that of blood culture, its yield is not reduced by up to 5 days of prior antibiotic therapy. Culture of intestinal secretions (best obtained by a noninvasive duodenal string test) can be positive despite a negative bone marrow culture. If blood, bone marrow, and intestinal secretions are all cultured, the yield is >90%. Stool cultures, although negative in 60–70% of cases during the first week, can become positive during the third week of infection in untreated patients.

 Serologic tests, including the classic Widal test for “febrile agglutinins,” and rapid tests to detect antibodies to outer-membrane proteins or O:9 antigen are available for detection of *S. typhi* in developing countries but have lower positive predictive values than blood culture. More sensitive antigen and nucleic acid amplification tests have been developed to detect *S. typhi* and

1052 *S. paratyphi* in blood but are not yet commercially available and remain impractical in many areas where enteric fever is endemic.

TREATMENT ENTERIC (TYPHOID) FEVER



Prompt administration of appropriate antibiotic therapy prevents severe complications of enteric fever and results in a case-fatality rate of <1%. The initial choice of antibiotics depends on the susceptibility of the *S. typhi* and *S. paratyphi* strains in the area of residence or travel (Table 190-1). For treatment of drug-susceptible typhoid fever, fluoroquinolones are the most effective class of agents, with cure rates of ~98% and relapse and fecal carriage rates of <2%. Experience is most extensive with ciprofloxacin. Short-course ofloxacin therapy is similarly successful against infection caused by quinolone-susceptible strains. However, the increased incidence of DCS *S. typhi* in Asia, which is probably related to the widespread availability of fluoroquinolones over the counter, is now limiting the use of this drug class for empirical therapy. Patients infected with DCS *S. typhi* strains should be treated with ceftriaxone, azithromycin, or high-dose ciprofloxacin. A 7-day course of high-dose fluoroquinolone therapy for DCS enteric fever has been associated with delayed resolution of fever and high rates of fecal carriage during convalescence. Thus, for DCS strains, a 10- to 14-day course of high-dose ciprofloxacin is preferred.

Ceftriaxone, cefotaxime, and (oral) cefixime are effective for treatment of MDR enteric fever, including that caused by DCS and fluoroquinolone-resistant strains. These agents clear fever in ~1 week, with failure rates of ~5–10%, fecal carriage rates of <3%, and relapse rates of 3–6%. Oral azithromycin results in defervescence in 4–6 days, with rates of relapse and convalescent stool carriage of <3%. Against DCS strains, azithromycin is associated with lower rates of treatment failure and shorter durations of hospitalization than are fluoroquinolones. Despite efficient *in vitro* killing of *Salmonella*, first- and second-generation cephalosporins as well as aminoglycosides are ineffective in the treatment of clinical infections.

TABLE 190-1 ANTIBIOTIC THERAPY FOR ENTERIC FEVER IN ADULTS

Indication	Agent	Dosage (Route)	Duration, Days
Empirical Treatment			
	Ceftriaxone ^a	2 g/d (IV)	10–14
	Azithromycin ^b	1 g/d (PO)	5
Fully Susceptible			
Optimal treatment	Ciprofloxacin ^c	500 mg bid (PO) or 400 mg q12h (IV)	5–7
	Azithromycin	1 g/d (PO)	5
Alternative treatment	Amoxicillin	1 g tid (PO) or 2 g q6h (IV)	14
	Chloramphenicol	25 mg/kg tid (PO or IV)	14–21
	Trimethoprim-sulfamethoxazole	160/800 mg bid (PO)	7–14
Multidrug-Resistant			
Optimal treatment	Ceftriaxone ^a	2 g/d (IV)	10–14
	Azithromycin	1 g/d (PO)	5
Alternative treatment	Ciprofloxacin	500 mg bid (PO) or 400 mg q12h (IV)	5–14
Quinolone-Resistant			
Optimal treatment	Ceftriaxone	2 g/d (IV)	10–14
	Azithromycin	1 g/d (PO)	5
Alternative treatment	High-dose ciprofloxacin	750 mg bid (PO) or 400 mg q8h (IV)	10–14

^aOr another third-generation cephalosporin (e.g., cefotaxime, 2 g q8h IV; or cefixime, 400 mg bid PO). ^bOr 1 g on day 1 followed by 500 mg/d PO for 6 days. ^cOr ofloxacin, 400 mg bid PO for 2–5 days.

Most patients with uncomplicated enteric fever can be managed at home with oral antibiotics and antipyretics. Patients with persistent vomiting, diarrhea, and/or abdominal distension should be hospitalized and given supportive therapy as well as a parenteral third-generation cephalosporin or fluoroquinolone, depending on the susceptibility profile. Therapy should be administered for at least 10 days or for 5 days after fever resolution.



In a randomized, prospective, double-blind study of critically ill patients with enteric fever (i.e., those with shock and obtundation) in Indonesia in the early 1980s, the administration of dexamethasone (an initial dose of 3 mg/kg followed by eight doses of 1 mg/kg every 6 h) with chloramphenicol was associated with a substantially lower mortality rate than was treatment with chloramphenicol alone (10% vs 55%). Although this study has not been repeated in the “post-chloramphenicol era,” severe enteric fever remains one of the few indications for glucocorticoid treatment of an acute bacterial infection.

The 1–5% of patients who develop chronic carriage of *Salmonella* can be treated for 4–6 weeks with an appropriate oral antibiotic. Treatment with oral amoxicillin, TMP-SMX, ciprofloxacin, or norfloxacin is ~80% effective in eradicating chronic carriage of susceptible organisms. However, in cases of anatomic abnormality (e.g., biliary or kidney stones), eradication often requires both antibiotic therapy and surgical correction.

PREVENTION AND CONTROL



Theoretically, it is possible to eliminate the salmonellae that cause enteric fever because they survive only in human hosts and are spread by contaminated food and water. However, given the high prevalence of the disease in developing countries that lack adequate sewage disposal and water treatment, this goal is currently unrealistic. Thus, travelers to developing countries should be advised to monitor their food and water intake carefully and to strongly consider immunization against *S. typhi*.

Two typhoid vaccines are commercially available: (1) Ty21a, an oral live attenuated *S. typhi* vaccine (given on days 1, 3, 5, and 7, with a booster every 5 years); and (2) Vi CPS, a parenteral vaccine consisting of purified Vi polysaccharide from the bacterial capsule (given in a single dose, with a booster every 2 years). The old parenteral whole-cell typhoid/paratyphoid A and B vaccine is no longer licensed, largely because of significant side effects, especially fever. An acetone-killed whole-cell vaccine is available only for use by the U.S. military. The minimal age for vaccination is 6 years for Ty21a and 2 years for Vi CPS. In a recent meta-analysis of vaccines for preventing typhoid fever in populations in endemic areas, the cumulative efficacy was 48% for Ty21a at 2.5–3.5 years and 55% for Vi CPS at 3 years. Although data on typhoid vaccines in travelers are limited, some evidence suggests that efficacy rates may be substantially lower than those for local populations in endemic areas. Currently, there is no licensed vaccine for paratyphoid fever.

Vi CPS typhoid vaccine is poorly immunogenic in children <5 years of age because of T cell-independent properties. In the more recently developed Vi-rEPA vaccine, Vi is bound to a nontoxic recombinant protein that is identical to *Pseudomonas aeruginosa* exotoxin A. In 2- to 4-year-olds, two injections of Vi-rEPA induced higher T cell responses and higher levels of serum IgG antibody to Vi than did Vi CPS in 5- to 14-year-olds. In a two-dose trial in 2- to 5-year-old children in Vietnam, Vi-rEPA provided 91% efficacy at 27 months and 89% efficacy at 46 months and was very well tolerated. This vaccine is not yet commercially available in the United States. Efforts to improve the immunogenicity and reduce the number of doses of live attenuated oral vaccines are ongoing.

Typhoid vaccine is not required for international travel, but it is recommended for travelers to areas where there is a moderate to high risk of exposure to *S. typhi*, especially those who are traveling to southern Asia and other developing regions of Asia, Africa, the Caribbean, and Central and South America and who will be exposed to potentially contaminated food and drink. Typhoid vaccine should be considered

even for persons planning <2 weeks of travel to high-risk areas. In addition, laboratory workers who deal with *S. typhi* and household contacts of known *S. typhi* carriers should be vaccinated. Because the protective efficacy of vaccine can be overcome by the high inocula that are commonly encountered in food-borne exposures, immunization is an adjunct and not a substitute for the avoidance of high-risk foods and beverages. Immunization is not recommended for adults residing in typhoid-endemic areas or for the management of persons who may have been exposed in a common-source outbreak.

Enteric fever is a notifiable disease in the United States. Individual health departments have their own guidelines for allowing ill or colonized food handlers or health care workers to return to their jobs. The reporting system enables public health departments to identify potential source patients and to treat chronic carriers in order to prevent further outbreaks. In addition, because 1–4% of patients with *S. typhi* infection become chronic carriers, it is important to monitor patients (especially child-care providers and food handlers) for chronic carriage and to treat this condition if indicated.

NONTYPHOIDAL SALMONELLOSIS

EPIDEMIOLOGY

In the United States, NTS causes ~12 million illnesses annually, and the incidence has remained relatively unchanged during the past two decades. In 2011, the incidence of NTS infection in this country was 16.5/100,000 persons—the highest rate among the 10 food-borne enteric pathogens under active surveillance. Five serotypes accounted for more than half of U.S. infections during the period 1996–2006: *typhimurium* (23%), *enteritidis* (16%), *newport* (10%), *heidelberg* (6%), and *javiana* (5%).

The incidence of nontyphoidal salmonellosis is highest during the rainy season in tropical climates and during the warmer months in temperate climates—a pattern coinciding with the peak in food-borne outbreaks. Rates of morbidity and mortality associated with NTS are highest among the elderly, infants, and immunocompromised individuals, including those with hemoglobinopathies, HIV infection, or infections that cause blockade of the reticuloendothelial system (e.g., bartonellosis, malaria, schistosomiasis, histoplasmosis).

Unlike *S. typhi* and *S. paratyphi*, whose only reservoir is humans, NTS can be acquired from multiple animal reservoirs. Transmission is most commonly associated with food products of animal origin (especially eggs, poultry, undercooked ground meat, and dairy products), fresh produce contaminated with animal waste, and contact with animals or their environments.

S. enteritidis infection associated with chicken eggs emerged as a major cause of food-borne disease during the 1980s and 1990s. *S. enteritidis* infection of the ovaries and upper oviduct tissue of hens results in contamination of egg contents before shell deposition. Infection is spread to egg-laying hens from breeding flocks and through contact with rodents and manure. The percentage of *Salmonella* outbreaks attributed to eggs has declined significantly in the United States, from 33% during 1998–1999 to 15% during 2006–2008. This decrease probably reflects the impact of the coordinated public health response to *S. enteritidis* infection attributed to eggs, including improved on-farm control measures, refrigeration, and education of consumers and food-service workers. Transmission via contaminated eggs can be prevented by cooking eggs until the yolk is solidified and pasteurizing egg products. Despite these control efforts, outbreaks of *S. enteritidis* infection associated with shell eggs continue to occur. In 2010, a national outbreak of *S. enteritidis* infection resulted in more than 1900 reported illnesses and the recall of 500 million eggs.

Centralization of food processing and widespread food distribution have contributed to the increased incidence of NTS in developed countries. Manufactured foods to which recent *Salmonella* outbreaks have been traced include peanut butter; milk products, including infant formula; and various processed foods, including packaged breakfast cereal, salsa, frozen prepared meals, and snack foods. Large outbreaks have also been linked to fresh produce, including alfalfa sprouts, cantaloupe, mangoes, papayas, and tomatoes; these items become

contaminated by manure or water at a single site and then are widely distributed.

An estimated 6% of sporadic *Salmonella* infections in the United States are attributed to contact with reptiles or amphibians, especially iguanas, snakes, turtles, and lizards. Reptile-associated *Salmonella* infection more commonly leads to hospitalization and more frequently involves children, including infants, than do other *Salmonella* infections. Other pets, including African hedgehogs, birds, rodents, baby chicks, ducklings, dogs, and cats, are also potential sources of NTS.



Increasing antibiotic resistance in NTS species is a global problem and has been linked to the widespread use of antimicrobial agents in food animals and especially in animal feed. In the early 1990s, *S. typhimurium* definitive phage type 104 (DT104), characterized by resistance to at least five antibiotics (ampicillin, chloramphenicol, streptomycin, sulfonamides, and tetracyclines; R-type ACSSuT), emerged worldwide. In 2010, resistance to at least ACSSuT was reported in 4.3% of NTS isolates, including 18.6% of *S. typhimurium* isolates. Acquisition is associated with exposure to ill farm animals and to various meat products, including uncooked or undercooked ground beef. Although probably no more virulent than susceptible *S. typhimurium* strains, DT104 strains are associated with an increased risk of bloodstream infection and hospitalization. DCS and trimethoprim-resistant DT104 strains are emerging, especially in the United Kingdom.

Because of increased resistance to conventional antibiotics such as ampicillin and TMP-SMX, extended-spectrum cephalosporins and fluoroquinolones have emerged as the agents of choice for the treatment of MDR NTS infections. In 2010, 2.8% of all NTS strains were resistant to ceftriaxone. Most ceftriaxone-resistant isolates were from children <18 years of age, in whom ceftriaxone is the antibiotic of choice for treatment of invasive NTS infection. These strains contained plasmid-encoded AmpC β -lactamases that were probably acquired by horizontal genetic transfer from *Escherichia coli* strains in food-producing animals—an event linked to the widespread use of the veterinary cephalosporin ceftiofur.

Over the last decade, strains of DCS NTS (MIC, 0.125–1 μ g/mL) have emerged and have been associated with delayed response and treatment failure. In 2009, 2.4% of NTS isolates in the United States were DCS or resistant to ciprofloxacin. These strains have diverse resistance mechanisms, including single and multiple mutations in the DNA gyrase genes *gyrA* and *gyrB* and plasmid-encoded quinolone resistance determinants that may not be reliably detected by nalidixic acid susceptibility testing. In 2012, the U.S. Clinical Laboratory Standards Institute proposed a lower ciprofloxacin susceptibility breakpoint (≥ 0.06 μ g/mL) for all *Salmonella* species to address this issue. Currently, because commercial test systems do not contain ciprofloxacin concentrations low enough to allow use of these breakpoints, laboratories need to determine the ciprofloxacin MIC by Etest or another alternative method.

CLINICAL MANIFESTATIONS

Gastroenteritis Infection with NTS most often results in gastroenteritis indistinguishable from that caused by other enteric pathogens. Nausea, vomiting, and diarrhea occur 6–48 h after the ingestion of contaminated food or water. Patients often experience abdominal cramping and fever (38–39°C; 100.5–102.2°F). Diarrheal stools are usually loose, nonbloody, and of moderate volume. However, large-volume watery stools, bloody stools, or symptoms of dysentery may occur. Rarely, NTS causes pseudoappendicitis or an illness that mimics inflammatory bowel disease.

Gastroenteritis caused by NTS is usually self-limited. Diarrhea resolves within 3–7 days and fever within 72 h. Stool cultures remain positive for 4–5 weeks after infection and—in rare cases of chronic carriage (<1%)—for >1 year. Antibiotic treatment usually is not recommended and may prolong fecal carriage. Neonates, the elderly, and immunosuppressed patients (e.g., transplant recipients, HIV-infected persons) with NTS gastroenteritis are especially susceptible to dehydration and dissemination and may require hospitalization and antibiotic therapy. Acute NTS gastroenteritis was associated with a threefold increased risk of dyspepsia and irritable bowel syndrome at 1 year in a study from Spain.

1054 Bacteremia and Endovascular Infections Up to 8% of patients with NTS gastroenteritis develop bacteremia; of these, 5–10% develop localized infections. Bacteremia and metastatic infection are most common with *Salmonella choleraesuis* and *Salmonella dublin* and among infants, the elderly, and immunocompromised patients, especially those with HIV infection. NTS endovascular infection should be suspected in high-grade or persistent bacteremia, especially with preexisting valvular heart disease, atherosclerotic vascular disease, prosthetic vascular graft, or aortic aneurysm. Arteritis should be suspected in elderly patients with prolonged fever and back, chest, or abdominal pain developing after an episode of gastroenteritis. Endocarditis and arteritis are rare (<1% of cases) but are associated with potentially fatal complications, including valve perforation, endomyocardial abscess, infected mural thrombus, pericarditis, mycotic aneurysms, aneurysm rupture, aortoenteric fistula, and vertebral osteomyelitis.



In some areas of sub-Saharan Africa, NTS may be among the most common causes—or even the most common cause—of bacteremia in children. NTS bacteremia among these children is not associated with diarrhea and has been associated with nutritional status and HIV infection.

Localized Infections • INTRAABDOMINAL INFECTIONS Intraabdominal infections due to NTS are rare and usually manifest as hepatic or splenic abscesses or as cholecystitis. Risk factors include hepatobiliary anatomic abnormalities (e.g., gallstones), abdominal malignancy, and sickle cell disease (especially with splenic abscesses). Eradication of the infection often requires surgical correction of abnormalities and percutaneous drainage of abscesses.

CENTRAL NERVOUS SYSTEM INFECTIONS NTS meningitis most commonly develops in infants 1–4 months of age. It often results in severe sequelae (including seizures, hydrocephalus, brain infarction, and mental retardation), with death in up to 60% of cases. Other rare central nervous system infections include ventriculitis, subdural empyema, and brain abscesses.

PULMONARY INFECTIONS NTS pulmonary infections usually present as lobar pneumonia, and complications include lung abscess, empyema, and bronchopleural fistula formation. The majority of cases occur in patients with lung cancer, structural lung disease, sickle cell disease, or glucocorticoid use.

URINARY AND GENITAL TRACT INFECTIONS Urinary tract infections caused by NTS present as either cystitis or pyelonephritis. Risk factors include malignancy, urolithiasis, structural abnormalities, HIV infection, and renal transplantation. NTS genital infections are rare and include ovarian and testicular abscesses, prostatitis, and epididymitis. Like other focal infections, both genital and urinary tract infections can be complicated by abscess formation.

BONE, JOINT, AND SOFT TISSUE INFECTIONS *Salmonella* osteomyelitis most commonly affects the femur, tibia, humerus, or lumbar vertebrae and is most often seen in association with sickle cell disease, hemoglobinopathies, or preexisting bone disease (e.g., fractures). Prolonged antibiotic treatment is recommended to decrease the risk of relapse and chronic osteomyelitis. Septic arthritis occurs in the same patient population as osteomyelitis and usually involves the knee, hip, or shoulder joints. Reactive arthritis can follow NTS gastroenteritis and is seen most frequently in persons with the HLA-B27 histocompatibility antigen. NTS rarely can cause soft tissue infections, usually at sites of local trauma in immunosuppressed patients.

DIAGNOSIS

The diagnosis of NTS infection is based on isolation of the organism from freshly passed stool or from blood or another ordinarily sterile body fluid. All salmonellae isolated in clinical laboratories should be sent to local public health departments for serotyping. Blood cultures should be done whenever a patient has prolonged or recurrent fever. Endovascular infection should be suspected if there is high-grade bacteremia (>50% of three or more positive blood cultures).

Echocardiography, computed tomography (CT), and indium-labeled white cell scanning are used to identify localized infection. When another localized infection is suspected, joint fluid, abscess drainage, or cerebrospinal fluid should be cultured, as clinically indicated.

TREATMENT NONTYPHOIDAL SALMONELLOSIS

Antibiotics should not be used routinely to treat uncomplicated NTS gastroenteritis. The symptoms are usually self-limited, and the duration of fever and diarrhea is not significantly decreased by antibiotic therapy. In addition, antibiotic treatment has been associated with increased rates of relapse, prolonged gastrointestinal carriage, and adverse drug reactions. Dehydration secondary to diarrhea should be treated with fluid and electrolyte replacement.

Preemptive antibiotic treatment (Table 190-2) should be considered for patients at increased risk for invasive NTS infection, including neonates (probably up to 3 months of age); persons >50 years of age with suspected atherosclerosis; and patients with immunosuppression, cardiac valvular or endovascular abnormalities, or significant joint disease. Treatment should consist of an oral or IV antibiotic administered for 48–72 h or until the patient becomes afebrile. Immunocompromised persons may require up to 7–14 days of therapy. The <1% of persons who develop chronic carriage of NTS should receive a prolonged antibiotic course, as described above for chronic carriage of *S. typhi*.

Because of the increasing prevalence of antibiotic resistance, empirical therapy for life-threatening NTS bacteremia or focal NTS infection should include a third-generation cephalosporin or a fluoroquinolone (Table 190-2). If the bacteremia is low-grade (<50% of positive blood cultures), the patient should be treated for 7–14 days. Patients with HIV/AIDS and NTS bacteremia should receive 1–2 weeks of IV antibiotic therapy followed by 4 weeks of oral therapy with a

TABLE 190-2 ANTIBIOTIC THERAPY FOR NONTYPHOIDAL SALMONELLA INFECTION IN ADULTS

Indication	Agent	Dosage (Route)	Duration, Days
Preemptive Treatment^a			
	Ciprofloxacin ^b	500 mg bid (PO)	2–3
Severe Gastroenteritis^c			
	Ciprofloxacin	500 mg bid (PO) or 400 mg q12h (IV)	3–7
	Trimethoprim-sulfamethoxazole	160/800 mg bid (PO)	
	Amoxicillin	1 g tid (PO)	
	Ceftriaxone	1–2 g/d (IV)	
Bacteremia			
	Ceftriaxone ^d	2 g/d (IV)	7–14
	Ciprofloxacin	400 mg q12h (IV), then 500 mg bid (PO)	
Endocarditis or Arteritis			
	Ceftriaxone	2 g/d (IV)	42
	Ciprofloxacin	400 mg q8h (IV), then 750 mg bid (PO)	
	Ampicillin	2 g q4h (IV)	
Meningitis			
	Ceftriaxone	2 g q12 h (IV)	14–21
	Ampicillin	2 g q4h (IV)	
Other Localized Infection			
	Ceftriaxone	2 g/d (IV)	14–28
	Ciprofloxacin	500 mg bid (PO) or 400 mg q12h (IV)	
	Ampicillin	2 g q6h (IV)	

^aConsider for neonates; persons >50 years of age with possible atherosclerotic vascular disease; and patients with immunosuppression, endovascular graft, or joint prosthesis. ^bOr ofloxacin, 400 mg bid (PO). ^cConsider on an individualized basis for patients with severe diarrhea and high fever who require hospitalization. ^dOr cefotaxime, 2 g q8h (IV).

fluoroquinolone. Patients whose infections relapse after this regimen should receive long-term suppressive therapy with a fluoroquinolone or TMP-SMX, as indicated by bacterial sensitivities.

If the patient has endocarditis or arteritis, treatment for 6 weeks with an IV β -lactam antibiotic (such as ceftriaxone or ampicillin) is indicated. IV ciprofloxacin followed by prolonged oral therapy is an option, but published experience is limited. Early surgical resection of infected aneurysms or other infected endovascular sites is recommended. Patients with infected prosthetic vascular grafts that cannot be resected have been maintained successfully on chronic suppressive oral therapy. For extraintestinal nonvascular infections, a 2- to 4-week course of antibiotic therapy (depending on the infection site) is usually recommended. In chronic osteomyelitis, abscess, or urinary or hepatobiliary infection associated with anatomic abnormalities, surgical resection or drainage may be required in addition to prolonged antibiotic therapy for eradication of infection.

PREVENTION AND CONTROL

Despite widespread efforts to prevent or reduce bacterial contamination of animal-derived food products and to improve food-safety education and training, recent declines in the incidence of NTS in the United States have been modest compared with those of other food-borne pathogens. This observation probably reflects the complex epidemiology of NTS. Identifying effective risk-reduction strategies requires monitoring of every step of food production, from handling of raw animal or plant products to preparation of finished foods. Contaminated food can be made safe for consumption by pasteurization, irradiation, or proper cooking. All cases of NTS infection should be reported to local public health departments because tracking and monitoring of these cases can identify the source(s) of infection and help authorities anticipate large outbreaks. Lastly, the prudent use of antimicrobial agents in both humans and animals is needed to limit the emergence of MDR *Salmonella*.

191 Shigellosis

Philippe J. Sansonetti, Jean Bergounioux

The discovery of *Shigella* as the etiologic agent of dysentery—a clinical syndrome of fever, intestinal cramps, and frequent passage of small, bloody, mucopurulent stools—is attributed to the Japanese microbiologist Kiyoshi Shiga, who isolated the Shiga bacillus (now known as *Shigella dysenteriae* type 1) from patients' stools in 1897 during a large and devastating dysentery epidemic. *Shigella* cannot be distinguished from *Escherichia coli* by DNA hybridization and remains a separate species only on historical and clinical grounds.

DEFINITION

Shigella is a non-spore-forming, gram-negative bacterium that, unlike *E. coli*, is nonmotile and does not produce gas from sugars, decarboxylate lysine, or hydrolyze arginine. Some serovars produce indole, and occasional strains utilize sodium acetate. *Shigella dysenteriae*, *Shigella flexneri*, *Shigella boydii*, and *Shigella sonnei* (serogroups A, B, C, and D, respectively) can be differentiated on the basis of biochemical and serologic characteristics. Genome sequencing of *E. coli* K12, *S. flexneri* 2a, *S. sonnei*, *S. dysenteriae* type 1, and *S. boydii* has revealed that these species have ~93% of genes in common. The three major genomic "signatures" of *Shigella* are (1) a 215-kb virulence plasmid that carries most of the genes required for pathogenicity (particularly invasive capacity); (2) the lack or alteration of genetic sequences encoding products (e.g., lysine decarboxylase) that, if expressed, would attenuate

pathogenicity; and (3) in *S. dysenteriae* type 1, the presence of genes encoding Shiga toxin, a potent cytotoxin.

EPIDEMIOLOGY



The human intestinal tract represents the major reservoir of *Shigella*, which is also found (albeit rarely) in the higher primates. Because excretion of shigellae is greatest in the acute phase of disease, the bacteria are transmitted most efficiently by the fecal-oral route via hand carriage; however, some outbreaks reflect foodborne or waterborne transmission. In impoverished areas, *Shigella* can be transmitted by flies. The high-level infectivity of *Shigella* is reflected by the very small inoculum required for experimental infection of volunteers (100 colony-forming units [CFU]), by the very high attack rates during outbreaks in day-care centers (33–73%), and by the high rates of secondary cases among family members of sick children (26–33%). Shigellosis can also be transmitted sexually.

Throughout history, *Shigella* epidemics have often occurred in settings of human crowding under conditions of poor hygiene—e.g., among soldiers in campaigning armies, inhabitants of besieged cities, groups on pilgrimages, and refugees in camps. Epidemics follow a cyclical pattern in areas such as the Indian subcontinent and sub-Saharan Africa. These devastating epidemics, which are most often caused by *S. dysenteriae* type 1, are characterized by high attack and mortality rates. In Bangladesh, for instance, an epidemic caused by *S. dysenteriae* type 1 was associated with a 42% increase in mortality rate among children 1–4 years of age. Apart from these epidemics, shigellosis is mostly an endemic disease, with 99% of cases occurring in the developing world and the highest prevalences in the most impoverished areas, where personal and general hygiene is below standard. *S. flexneri* isolates predominate in the least developed areas, whereas *S. sonnei* is more prevalent in economically emerging countries and in the industrialized world.

Prevalence in the Developing World In a review published under the auspices of the World Health Organization (WHO), the total annual number of cases in 1966–1997 was estimated at 165 million, and 69% of these cases occurred in children <5 years of age. In this review, the annual number of deaths was calculated to range between 500,000 and 1.1 million. More recent data (2000–2004) from six Asian countries indicate that, even though the incidence of shigellosis remains stable, mortality rates associated with this disease may have decreased significantly, possibly as a result of improved nutritional status. However, extensive and essentially uncontrolled use of antibiotics, which may also account for declining mortality rates, has increased the rate of emergence of multidrug-resistant *Shigella* strains. A 2013 prospective matched case-control study of children <5 years of age emphasizes the importance of *Shigella* in the burden and etiology of diarrheal diseases in developing countries. *Shigella* is one of the top four pathogens associated with moderate to severe diarrhea and is now ranked first among children 12–59 months of age. These moderate to severe cases account for an 8.5-fold increase in mortality incidence over the average diarrheal disease-related mortality. The study's authors conclude that *Shigella* remains a major pathogen to be targeted by health care programs.

An often-overlooked complication of shigellosis is the short- and long-term impairment of the nutritional status of infected children in endemic areas. Combined with anorexia, the exudative enteropathy resulting from mucosal abrasions contributes to rapid deterioration of the patient's nutritional status. Shigellosis is thus a major contributor to stunted growth among children in developing countries.

Peaking in incidence in the pediatric population, endemic shigellosis is rare among young and middle-aged adults, probably because of naturally acquired immunity. Incidence then increases again in the elderly population.

Prevalence in the Industrialized World In pediatric populations, local outbreaks occur when proper and adapted hygiene policies are not implemented in group facilities like day-care centers and institutions for the mentally retarded. In adults, as in children, sporadic cases occur among travelers returning from endemic areas, and rare outbreaks of varying size can follow waterborne or food-borne infections.

Shigella infection occurs essentially through oral contamination via direct fecal-oral transmission, the organism being poorly adapted to survive in the environment. Resistance to low-pH conditions allows shigellae to survive passage through the gastric barrier, an ability that may explain in part why a small inoculum (as few as 100 CFU) is sufficient to cause infection.

The watery diarrhea that usually precedes the dysenteric syndrome is attributable to active secretion and abnormal water reabsorption—a secretory effect at the jejunal level described in experimentally infected rhesus monkeys. This initial purge is probably due to the combined action of an enterotoxin (ShET-1) and mucosal inflammation. The dysenteric syndrome, manifested by bloody and mucopurulent stools, reflects invasion of the mucosa.

The pathogenesis of *Shigella* is essentially determined by a large virulence plasmid of 214 kb comprising ~100 genes, of which 25 encode a type III secretion system that inserts into the membrane of the host cell to allow effectors to transit from the bacterial cytoplasm to the host cell cytoplasm (Fig. 191-1). Bacteria are thereby able to invade intestinal epithelial cells by inducing their own uptake after the initial crossing of the epithelial barrier through M cells (the specialized translocating epithelial cells in the follicle-associated epithelium that covers mucosal lymphoid nodules). The organisms induce apoptosis of subepithelial resident macrophages. Once inside the cytoplasm of intestinal epithelial cells, *Shigella* effectors trigger the cytoskeletal rearrangements necessary to direct uptake of the organism into the epithelial cell. The *Shigella*-containing vacuole is then quickly lysed, releasing bacteria into the cytosol.

Intracellular shigellae next use cytoskeletal components to propel themselves inside the infected cell; when the moving organism and the host cell membrane come into contact, cellular protrusions form and are engulfed by neighboring cells. This series of events permits bacterial cell-to-cell spread.

Cytokines released by a growing number of infected intestinal epithelial cells attract increased numbers of immune cells (particularly polymorphonuclear leukocytes [PMNs]) to the infected site, thus further destabilizing the epithelial barrier, exacerbating inflammation, and leading to the acute colitis that characterizes shigellosis. Evidence indicates that some type III secretion system–injected effectors can control the extent of inflammation, thus facilitating bacterial survival.

Shiga toxin produced by *S. dysenteriae* type 1 increases disease severity. This toxin belongs to a group of A1-B5 protein toxins whose B subunit binds to the receptor globotriaosylceramide on the target cell surface and whose catalytic A subunit is internalized by receptor-mediated endocytosis and interacts with the subcellular machinery to inhibit protein synthesis by expressing RNA N-glycosidase activity on

28S ribosomal RNA. This process leads to inhibition of binding of the amino-acyl-tRNA to the 60S ribosomal subunit and thus to a general shutoff of cell protein biosynthesis. Shiga toxins are translocated from the bowel into the circulation. After binding of the toxins to target cells in the kidney, pathophysiologic alterations may result in hemolytic-uremic syndrome (HUS; see below).

CLINICAL MANIFESTATIONS

The presentation and severity of shigellosis depend to some extent on the infecting serotype but even more on the age and the immunologic and nutritional status of the host. Poverty and poor standards of hygiene are strongly related to the number and severity of diarrheal episodes, especially in children <5 years old who have been weaned.

Shigellosis typically evolves through four phases: incubation, watery diarrhea, dysentery, and the postinfectious phase. The incubation period usually lasts 1–4 days but may be as long as 8 days. Typical initial manifestations are transient fever, limited watery diarrhea, malaise, and anorexia. Signs and symptoms may range from mild abdominal discomfort to severe cramps, diarrhea, fever, vomiting, and tenesmus. The manifestations are usually exacerbated in children, with temperatures up to 40°–41°C (104.0°–105.8°F) and more severe anorexia and watery diarrhea. This initial phase may represent the only clinical manifestation of shigellosis, especially in developed countries. Otherwise, dysentery follows within hours or days and is characterized by uninterrupted excretion of small volumes of bloody mucopurulent stools with increased tenesmus and abdominal cramps. At this stage, *Shigella* produces acute colitis involving mainly the distal colon and the rectum. Unlike most diarrheal syndromes, dysenteric syndromes rarely present with dehydration as a major feature. Endoscopy shows an edematous and hemorrhagic mucosa, with ulcerations and possibly overlying exudates resembling pseudomembranes. The extent of the lesions correlates with the number and frequency of stools and with the degree of protein loss by exudative mechanisms. Most episodes are self-limited and resolve without treatment in 1 week. With appropriate treatment, recovery takes place within a few days to a week, with no sequelae.

Acute life-threatening complications are seen most often in children <5 years of age (particularly those who are malnourished) and in elderly patients. Risk factors for death in a clinically severe case include nonbloody diarrhea, moderate to severe dehydration, bacteremia, absence of fever, abdominal tenderness, and rectal prolapse. Major complications are predominantly intestinal (e.g., toxic megacolon, intestinal perforations, rectal prolapse) or metabolic (e.g., hypoglycemia, hyponatremia, dehydration). Bacteremia is rare and is reported most frequently in severely malnourished and HIV-infected patients. Alterations of consciousness, including seizures, delirium, and coma, may occur, especially in children <5 years old,

and are associated with a poor prognosis; fever and severe metabolic alterations are more often the major causes of altered consciousness than is meningitis or the Ekiri syndrome (toxic encephalopathy associated with bizarre posturing, cerebral edema, and fatty degeneration of viscera), which has been reported mostly in Japanese children. Pneumonia, vaginitis, and keratoconjunctivitis due to *Shigella* are rarely reported. In the absence of serious malnutrition, severe and very unusual clinical manifestations, such as meningitis, may be linked to genetic defects in innate immune functions (i.e., deficiency in interleukin 1 receptor–associated kinase 4 [IRAK-4]) and may require genetic investigation.

Two complications of particular importance are toxic megacolon and HUS. Toxic megacolon is a consequence of severe inflammation extending to the colonic smooth-muscle layer and causing paralysis and dilation. The patient presents with abdominal distention and tenderness, with or without signs of localized or generalized peritonitis. The abdominal x-ray characteristically shows

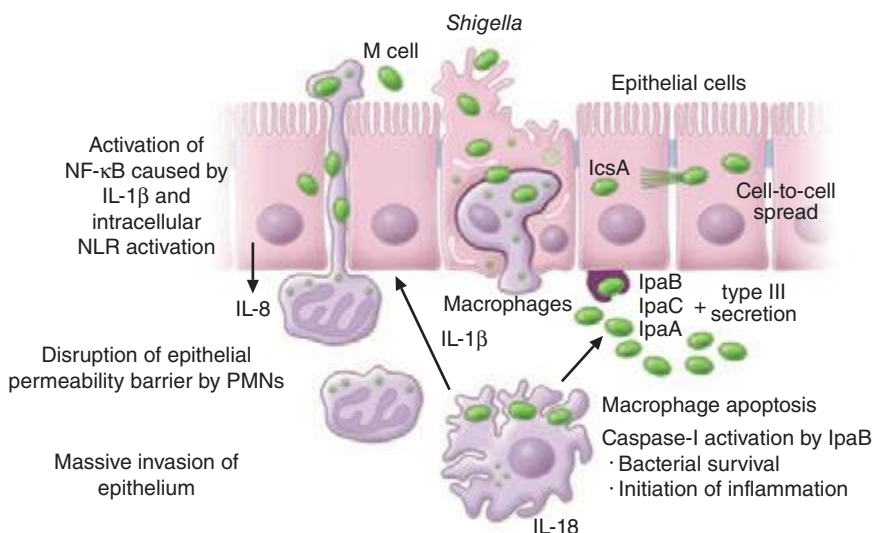


FIGURE 191-1 Invasive strategy of *Shigella flexneri*. IL, interleukin; NF-κB, nuclear factor κB; NLR, NOD-like receptor; PMN, polymorphonuclear leukocyte.

marked dilation of the transverse colon (with the greatest distention in the ascending and descending segments); thumbprinting caused by mucosal inflammatory edema; and loss of the normal haustral pattern associated with pseudopolyps, often extending into the lumen. Pneumatosis coli is an occasional finding. If perforation occurs, radiographic signs of pneumoperitoneum may be apparent. Predisposing factors (e.g., hypokalemia and use of opioids, anticholinergics, loperamide, psyllium seeds, and antidepressants) should be investigated.



Shiga toxin produced by *S. dysenteriae* type 1 has been linked to HUS in developing countries but rarely in industrialized countries, where enterohemorrhagic *E. coli* (EHEC) predominates as the etiologic agent of this syndrome. HUS is an early complication that most often develops after several days of diarrhea. Clinical examination shows pallor, asthenia, and irritability and, in some cases, bleeding of the nose and gums, oliguria, and increasing edema. HUS is a nonimmune (Coombs test–negative) hemolytic anemia defined by a diagnostic triad: microangiopathic hemolytic anemia (hemoglobin level typically <80 g/L [<8 g/dL]), thrombocytopenia (mild to moderate in severity; typically <60,000 platelets/ μ L), and acute renal failure due to thrombosis of the glomerular capillaries (with markedly elevated creatinine levels). Anemia is severe, with fragmented red blood cells (*schizocytes*) in the peripheral smear, high serum concentrations of lactate dehydrogenase and free circulating hemoglobin, and elevated reticulocyte counts. Acute renal failure occurs in 55–70% of cases; however, renal function recovers in most of these cases (up to 70% in various series). Leukemoid reactions, with leukocyte counts of 50,000/ μ L, are sometimes noted in association with HUS.

The postinfectious immunologic complication known as *reactive arthritis* can develop weeks or months after shigellosis, especially in patients expressing the histocompatibility antigen HLA-B27. About 3% of patients infected with *S. flexneri* later develop this syndrome, with arthritis, ocular inflammation, and urethritis—a condition that can last for months or years and can progress to difficult-to-treat chronic arthritis. Postinfectious arthropathy occurs only after infection with *S. flexneri* and not after infection with the other *Shigella* serotypes.

LABORATORY DIAGNOSIS



The differential diagnosis in patients with a dysenteric syndrome depends on the clinical and environmental context. In developing areas, infectious diarrhea caused by other invasive pathogenic bacteria (*Salmonella*, *Campylobacter jejuni*, *Clostridium difficile*, *Yersinia enterocolitica*) or parasites (*Entamoeba histolytica*) should be considered. Only bacteriologic and parasitologic examinations of stool can truly differentiate among these pathogens. A first flare of inflammatory bowel disease, such as Crohn's disease or ulcerative colitis (Chap. 351), should be considered in patients in industrialized countries. Despite the similarity in symptoms, anamnesis discriminates between shigellosis, which usually follows recent travel in an endemic zone, and these other conditions.

Microscopic examination of stool smears shows erythrophagocytic trophozoites with very few PMNs in *E. histolytica* infection, whereas bacterial enteroinvasive infections (particularly shigellosis) are characterized by high PMN counts in each microscopic field. However, because shigellosis often manifests only as watery diarrhea, systematic attempts to isolate *Shigella* are necessary.

The “gold standard” for the diagnosis of *Shigella* infection remains the isolation and identification of the pathogen from fecal material. One major difficulty, particularly in endemic areas where laboratory facilities are not immediately available, is the fragility of *Shigella* and its common disappearance during transport, especially with rapid changes in temperature and pH. In the absence of a reliable enrichment medium, buffered glycerol saline or Cary-Blair medium can be used as a holding medium, but prompt inoculation onto isolation medium is essential. The probability of isolation is higher if the portion of stools that contains bloody and/or mucopurulent material is directly sampled. Rectal swabs can be used, as they offer the highest rate of successful isolation during the acute phase of disease. Blood cultures are positive in fewer than 5% of cases but should be done when a patient presents with a clinical picture of severe sepsis.

In addition to quick processing, the use of several media increases the likelihood of successful isolation: a nonselective medium such as bromocresol-purple agar lactose; a low-selectivity medium such as MacConkey or eosin-methylene blue; and a high-selectivity medium such as Hektoen, *Salmonella-Shigella*, or xylose-lysine-deoxycholate agar. After incubation on these media for 12–18 h at 37°C (98.6°F), shigellae appear as non-lactose-fermenting colonies that measure 0.5–1 mm in diameter and have a convex, translucent, smooth surface. Suspected colonies on nonselective or low-selectivity medium can be subcultured on a high-selectivity medium before being specifically identified or can be identified directly by standard commercial systems on the basis of four major characteristics: glucose positivity (usually without production of gas), lactose negativity, H₂S negativity, and lack of motility. The four *Shigella* serogroups (A–D) can then be differentiated by additional characteristics. This approach adds time and difficulty to the identification process; however, after presumptive diagnosis, the use of serologic methods (e.g., slide agglutination, with group- and then type-specific antisera) should be considered. Group-specific antisera are widely available; in contrast, because of the large number of serotypes and subserotypes, type-specific antisera are rare and more expensive and thus are often restricted to reference laboratories.

TREATMENT SHIGELLOSIS

ANTIBIOTIC SUSCEPTIBILITY OF SHIGELLA



As an enteroinvasive disease, shigellosis requires antibiotic treatment. Since the mid-1960s, however, increasing resistance to multiple drugs has been a dominant factor in treatment decisions. Resistance rates are highly dependent on the geographic area. Clonal spread of particular strains and horizontal transfer of resistance determinants, particularly via plasmids and transposons, contribute to multidrug resistance. The current global status—i.e., high rates of resistance to classic first-line antibiotics such as amoxicillin—has led to a rapid switch to quinolones such as nalidixic acid. However, resistance to such early-generation quinolones has also emerged and spread quickly as a result of chromosomal mutations affecting DNA gyrase and topoisomerase IV; this resistance has necessitated the use of later-generation quinolones as first-line antibiotics in many areas. For instance, a review of the antibiotic resistance history of *Shigella* in India found that, after their introduction in the late 1980s, the second-generation quinolones norfloxacin, ciprofloxacin, and ofloxacin were highly effective in the treatment of shigellosis, including cases caused by multidrug-resistant strains of *S. dysenteriae* type 1. However, investigations of subsequent outbreaks in India and Bangladesh detected resistance to norfloxacin, ciprofloxacin, and ofloxacin in 5% of isolates. The incidence of multidrug resistance parallels the widespread, uncontrolled use of antibiotics and calls for the rational use of effective drugs.

ANTIBIOTIC TREATMENT OF SHIGELLOSIS (TABLE 191-1)

Because of the ready transmissibility of *Shigella*, current public health recommendations in the United States are that every case be treated with antibiotics. Ciprofloxacin is recommended as first-line treatment. A number of other drugs have been tested and shown to be effective, including ceftriaxone, azithromycin, pivmecillinam, and some fifth-generation quinolones. Whereas infections caused by non-*dysenteriae* *Shigella* in immunocompetent individuals are routinely treated with a 3-day course of antibiotics, it is recommended that *S. dysenteriae* type 1 infections be treated for 5 days and that *Shigella* infections in immunocompromised patients be treated for 7–10 days.



Treatment for shigellosis must be adapted to the clinical context, with the recognition that the most fragile patients are children <5 years old, who represent two-thirds of all cases worldwide. There are few data on the use of quinolones in children, but *Shigella*-induced dysentery is a well-recognized indication

TABLE 191-1 RECOMMENDED ANTIMICROBIAL THERAPY FOR SHIGELLOSIS

Antimicrobial Agent	Treatment Schedule		Limitations
	Children	Adults	
First Line			
Ciprofloxacin	15 mg/kg 2 times per day for 3 days, PO	500 mg	
Second Line			
Pivmecillinam	20 mg/kg 4 times per day for 5 days PO	100 mg	Cost No pediatric formulation Frequent administration Emerging resistance
Ceftriaxone	50–100 mg/kg Once a day IM for 2–5 days	–	Efficacy not validated Must be injected
Azithromycin	6–20 mg/kg Once a day for 1–5 days PO	1–1.5 g	Cost Efficacy not validated Minimum inhibitory concentration near serum concentration Rapid emergence of resistance and spread to other bacteria

Source: WHO Library Cataloguing-in-Publication Data: Guidelines for the control of shigellosis, including epidemics due to *Shigella dysenteriae* type 1 (www.who.int/cholera/publications/shigellosis/en/).

for their use. The half-life of ciprofloxacin is longer in infants than in older individuals. The ciprofloxacin dose generally recommended for children is 30 mg/kg per day in two divided doses. Adults living in areas with high standards of hygiene are likely to develop milder, shorter-duration disease, whereas infants in endemic areas can develop severe, sometimes fatal, dysentery. In the former setting, treatment will remain minimal and bacteriologic proof of infection will often come after symptoms have resolved; in the latter setting, antibiotic treatment and more aggressive measures, possibly including resuscitation, are often required.

REHYDRATION AND NUTRITION



Shigella infection rarely causes significant dehydration. Cases requiring aggressive rehydration (particularly in industrialized countries) are uncommon. In developing countries, malnutrition remains the primary indicator for diarrhea-related death, highlighting the importance of nutrition in early management. Rehydration should be oral unless the patient is comatose or presents in shock. Because of the improved effectiveness of reduced-osmolarity oral rehydration solution (especially for children with acute noncholera diarrhea), the WHO and UNICEF now recommend a standard solution of 245 mOsm/L (sodium, 75 mmol/L; chloride, 65 mmol/L; glucose [anhydrous], 75 mmol/L; potassium, 20 mmol/L; citrate, 10 mmol/L). In shigellosis, the coupled transport of sodium to glucose may be variably affected, but oral rehydration therapy remains the easiest and most efficient form of rehydration, especially in severe cases.

Nutrition should be started as soon as possible after completion of initial rehydration. Early refeeding is safe, well tolerated, and clinically beneficial. Because breast-feeding reduces diarrheal losses and the need for oral rehydration in infants, it should be maintained in the absence of contraindications (e.g., maternal HIV infection).

NONSPECIFIC, SYMPTOM-BASED THERAPY

Antimotility agents have been implicated in prolonged fever in volunteers with shigellosis. These agents are suspected of increasing the risk of toxic megacolon and are thought to have been responsible for HUS in children infected by EHEC strains. For safety reasons, it is better to avoid antimotility agents in bloody diarrhea.

TREATMENT OF COMPLICATIONS

There is no consensus regarding the best treatment for toxic megacolon. The patient should be assessed frequently by both medical and surgical teams. Anemia, dehydration, and electrolyte deficits (particularly hypokalemia) may aggravate colonic atony and should be actively treated. Nasogastric aspiration helps to deflate the colon. Parenteral nutrition has not been proven to be beneficial. Fever persisting beyond 48–72 h raises the possibility of local perforation or abscess. Most studies recommend colectomy if, after 48–72 h, colonic distention persists. However, some physicians recommend continuation of medical therapy for up to 7 days if the patient seems to be improving clinically despite persistent megacolon without free perforation. Intestinal perforation, either isolated or complicating toxic megacolon, requires surgical treatment and intensive medical support.

Rectal prolapse must be treated as soon as possible. With the health care provider using surgical gloves or a soft warm wet cloth and the patient in the knee-chest position, the prolapsed rectum is gently pushed back into place. If edema of the rectal mucosa is evident (rendering reintegration difficult), it can be osmotically reduced by the application of gauze impregnated with a warm solution of saturated magnesium sulfate. Rectal prolapse often relapses but usually resolves along with the resolution of dysentery.

HUS must be treated by water restriction, including discontinuation of oral rehydration solution and potassium-rich alimentation. Hemofiltration is usually required.

PREVENTION

Hand washing after defecation or handling of children's feces and before handling of food is recommended. Stool decontamination (e.g., with sodium hypochlorite), together with a cleaning protocol for medical staff as well as for patients, has proven useful in limiting the spread of infection during *Shigella* outbreaks. Ideally, patients should have a negative stool culture before their infection is considered cured. Recurrences are rare if therapeutic and preventive measures are correctly implemented.

Although several live attenuated oral and subunit parenteral vaccine candidates have been produced and are undergoing clinical trials, no vaccine against shigellosis is currently available. Especially given the rapid progression of antibiotic resistance in *Shigella*, a vaccine is urgently needed.

192

Infections Due to *Campylobacter* and Related Organisms

Martin J. Blaser

DEFINITION

Bacteria of the genus *Campylobacter* and of the related genera *Arcobacter* and *Helicobacter* (Chap. 188) cause a variety of inflammatory conditions. Although acute diarrheal illnesses are most common, these organisms may cause infections in virtually all parts of the body, especially in compromised hosts, and these infections may have late nonsuppurative sequelae. The designation *Campylobacter* comes from the Greek for “curved rod” and refers to the organism's vibrio-like morphology.

ETIOLOGY

Campylobacters are motile, non-spore-forming, curved, gram-negative rods. Originally known as *Vibrio fetus*, these bacilli were reclassified as a new genus in 1973 after their dissimilarity to other vibrios

was recognized. More than 15 species have since been identified. These species are currently divided into three genera: *Campylobacter*, *Arcobacter*, and *Helicobacter*. Not all of the species are pathogens of humans. The human pathogens fall into two major groups: those that primarily cause diarrheal disease and those that cause extraintestinal infection. The principal diarrheal pathogen is *Campylobacter jejuni*, which accounts for 80–90% of all cases of recognized illness due to campylobacters and related genera. Other organisms that cause diarrheal disease include *Campylobacter coli*, *Campylobacter upsaliensis*, *Campylobacter lari*, *Campylobacter hyointestinalis*, *Campylobacter fetus*, *Arcobacter butzleri*, *Arcobacter cryaerophilus*, *Helicobacter cinaedi*, and *Helicobacter fennelliae*. The two *Helicobacter* species causing diarrheal disease, *H. cinaedi* and *H. fennelliae*, are intestinal rather than gastric organisms; in terms of the clinical features of the illnesses they cause, these species most closely resemble *Campylobacter* rather than *Helicobacter pylori* (Chap. 188) and thus are considered in this chapter. The pathogenic roles of *Campylobacter concisus*, *Campylobacter ureolyticus*, *Campylobacter troglodytis*, and *Campylobacter pyloridis* are uncertain. A new subspecies—*C. fetus* subspecies *testudinum*—has been described, chiefly in Asian patients; its close resemblance to strains isolated from reptiles suggests a food source.

The major species causing extraintestinal illnesses is *C. fetus*. However, any of the diarrheal agents listed above may cause systemic or localized infection as well, especially in compromised hosts. Neither aerobes nor strict anaerobes, these microaerophilic organisms are adapted for survival in the gastrointestinal mucous layer. This chapter focuses on *C. jejuni* and *C. fetus* as the major pathogens in and prototypes for their groups. The key features of infection are listed by species (excluding *C. jejuni*, described in detail in the text below) in Table 192-1.

EPIDEMIOLOGY



Campylobacters are found in the gastrointestinal tract of many animals used for food (including poultry, cattle, sheep, and swine) and many household pets (including birds, dogs, and cats). These microorganisms usually do not cause illness in their animal hosts. In most cases, campylobacters are transmitted to humans in raw or undercooked food products or through direct contact with infected animals. In the United States and other developed

countries, ingestion of contaminated poultry that has not been sufficiently cooked is the most common mode of acquisition (30–70% of cases). Other modes include ingestion of raw (unpasteurized) milk or untreated water, contact with infected household pets, travel to developing countries (campylobacters being among the leading causes of traveler's diarrhea; Chaps. 149 and 160), oral-anal sexual contact, and (occasionally) contact with an index case who is incontinent of stool (e.g., a baby).

Campylobacter infections are common. Several studies indicate that, in the United States, diarrheal disease due to campylobacters is more common than that due to *Salmonella* and *Shigella* combined. Infections occur throughout the year, but their incidence peaks during summer and early autumn. Persons of all ages are affected; however, attack rates for *C. jejuni* are highest among young children and young adults, whereas those for *C. fetus* are highest at the extremes of age. Systemic infections due to *C. fetus* (and to other *Campylobacter* and related species) are most common among compromised hosts. Persons at increased risk include those with AIDS, hypogammaglobulinemia, neoplasia, liver disease, diabetes mellitus, and generalized atherosclerosis as well as neonates and pregnant women. However, apparently healthy nonpregnant persons occasionally develop transient *Campylobacter* bacteremia as part of a gastrointestinal illness.

In contrast, in many developing countries, *C. jejuni* infections are hyperendemic, with the highest rates among children <2 years old. Infection rates fall with age, as does the illness-to-infection ratio. These observations suggest that frequent exposure to *C. jejuni* leads to the acquisition of immunity.

PATHOLOGY AND PATHOGENESIS



C. jejuni infections may be subclinical, especially in hosts in developing countries who have had multiple prior infections and thus are partially immune. Symptomatic infections mostly occur within 2–4 days (range, 1–7 days) of exposure to the organism in food or water. The sites of tissue injury include the jejunum, ileum, and colon. Biopsies show an acute nonspecific inflammatory reaction, with neutrophils, monocytes, and eosinophils in the lamina propria, as well as damage to the epithelium, including loss of mucus, glandular degeneration, and crypt abscesses. Biopsy findings may be consistent with Crohn's disease or ulcerative colitis, but these "idiopathic"

TABLE 192-1 CLINICAL FEATURES ASSOCIATED WITH INFECTION DUE TO "ATYPICAL" *CAMPYLOBACTER* AND RELATED SPECIES IMPLICATED AS CAUSES OF HUMAN ILLNESS

Species	Common Clinical Features	Less Common Clinical Features	Additional Information
<i>Campylobacter coli</i>	Fever, diarrhea, abdominal pain	Bacteremia ^a	Clinically indistinguishable from <i>C. jejuni</i>
<i>Campylobacter fetus</i>	Bacteremia, ^a sepsis, meningitis, vascular infections	Diarrhea, relapsing fevers	Not usually isolated from media containing cephalothin or incubated at 42°C
<i>Campylobacter upsaliensis</i>	Watery diarrhea, low-grade fever, abdominal pain	Bacteremia, abscesses	Difficult to isolate because of cephalothin susceptibility
<i>Campylobacter lari</i>	Abdominal pain, diarrhea	Colitis, appendicitis	Seagulls frequently colonized; organism often transmitted to humans via contaminated water
<i>Campylobacter hyointestinalis</i>	Watery or bloody diarrhea, vomiting, abdominal pain	Bacteremia	Causes proliferative enteritis in swine
<i>Helicobacter fennelliae</i>	Chronic mild diarrhea, abdominal cramps, proctitis	Bacteremia ^a	Best treated with fluoroquinolones
<i>Helicobacter cinaedi</i>	Chronic mild diarrhea, abdominal cramps, proctitis	Bacteremia ^a	Best treated with fluoroquinolones; identified in healthy hamsters
<i>Campylobacter jejuni</i> subspecies <i>doylei</i>	Diarrhea	Chronic gastritis, bacteremia ^b	Uncertain role as human pathogen
<i>Arcobacter cryaerophilus</i>	Diarrhea	Bacteremia	Cultured under aerobic conditions
<i>Arcobacter butzleri</i>	Fever, diarrhea, abdominal pain, nausea	Bacteremia, appendicitis	Cultured under aerobic conditions; enzootic in nonhuman primates
<i>Campylobacter sputorum</i>	Pulmonary, perianal, groin, and axillary abscesses; diarrhea	Bacteremia	Three clinically relevant biovars: <i>sputorum</i> , <i>faecalis</i> , and <i>paraureolyticus</i>

^aIn immunocompromised hosts, especially HIV-infected persons. ^bIn children.

Source: Adapted from BM Allos, MJ Blaser. Clin Infect Dis 20:1092, 1995.

1060 chronic inflammatory diseases should not be diagnosed unless infectious colitis, *specifically including* that due to infection with *Campylobacter* species and related organisms, has been ruled out.

The high frequency of *C. jejuni* infections and their severity and recurrence among hypogammaglobulinemic patients suggest that antibodies are important in protective immunity. The pathogenesis of infection is uncertain. Both the motility of the strain and its capacity to adhere to host tissues appear to favor disease, but classic enterotoxins and cytotoxins (although they have been described and include cytotoxin distending toxin, or CDT) appear not to play substantial roles in tissue injury or disease production. The organisms have been visualized within the epithelium, albeit in low numbers. The documentation of a significant tissue response and occasionally of *C. jejuni* bacteremia further suggests that tissue invasion is clinically significant, and *in vitro* studies are consistent with this pathogenetic feature.

The pathogenesis of *C. fetus* infections is better defined. Virtually all clinical isolates of *C. fetus* possess a proteinaceous capsule-like structure (an S-layer) that renders the organisms resistant to complement-mediated killing and opsonization. As a result, *C. fetus* can cause bacteremia and can seed sites beyond the intestinal tract. The ability of the organism to switch the S-layer proteins expressed—a phenomenon that results in antigenic variability—may contribute to the chronicity and high rate of recurrence of *C. fetus* infections in compromised hosts.

CLINICAL MANIFESTATIONS

The clinical features of infections due to *Campylobacter* and the related *Arcobacter* and intestinal *Helicobacter* species causing enteric disease appear to be highly similar. *C. jejuni* can be considered the prototype, in part because it is by far the most common enteric pathogen in the group. A prodrome of fever, headache, myalgia, and/or malaise often occurs 12–48 h before the onset of diarrheal symptoms. The most common signs and symptoms of the intestinal phase are diarrhea, abdominal pain, and fever. The degree of diarrhea varies from several loose stools to grossly bloody stools; most patients presenting for medical attention have ≥ 10 bowel movements on the worst day of illness. Abdominal pain usually consists of cramping and may be the most prominent symptom. Pain is usually generalized but may become localized; *C. jejuni* infection may cause pseudoappendicitis. Fever may be the only initial manifestation of *C. jejuni* infection, a situation mimicking the early stages of typhoid fever. Febrile young children may develop convulsions. *Campylobacter* enteritis is generally self-limited; however, symptoms persist for >1 week in 10–20% of patients seeking medical attention, and clinical relapses occur in 5–10% of such untreated patients. Studies of common-source epidemics indicate that milder illnesses or asymptomatic infections may commonly occur.

C. fetus may cause a diarrheal illness similar to that due to *C. jejuni*, especially in normal hosts. This organism also may cause either intermittent diarrhea or nonspecific abdominal pain without localizing signs. Sequelae are uncommon, and the outcome is benign. *C. fetus* may also cause a prolonged relapsing systemic illness (with fever, chills, and myalgias) that has no obvious primary source; this manifestation is especially common among compromised hosts. Secondary seeding of an organ (e.g., meninges, brain, bone, urinary tract, or soft tissue) complicates the course, which may be fulminant. *C. fetus* infections have a tropism for vascular sites: endocarditis, mycotic aneurysm, and septic thrombophlebitis may all occur. Infection during pregnancy often leads to fetal death. A variety of *Campylobacter* species and *H. cinaedi* can cause recurrent cellulitis with fever and bacteremia in immunocompromised hosts.

COMPLICATIONS

Except in infection with *C. fetus*, bacteremia is uncommon, developing most often in immunocompromised hosts and at the extremes of age. Three patterns of extraintestinal infection have been noted: (1) transient bacteremia in a normal host with enteritis (benign course, no specific treatment needed); (2) sustained bacteremia or focal infection in a normal host (bacteremia originating from enteritis, with patients responding well to antimicrobial therapy); and (3) sustained

bacteremia or focal infection in a compromised host. Enteritis may not be clinically apparent. Antimicrobial therapy, possibly prolonged, is necessary for suppression or cure of the infection.

Campylobacter, *Arcobacter*, and intestinal *Helicobacter* infections in patients with AIDS or hypogammaglobulinemia may be severe, persistent, and extraintestinal; relapse after cessation of therapy is common. Hypogammaglobulinemic patients also may develop osteomyelitis and an erysipelas-like rash or cellulitis.

Local suppurative complications of infection include cholecystitis, pancreatitis, and cystitis; distant complications include meningitis, endocarditis, arthritis, peritonitis, cellulitis, and septic abortion. All these complications are rare, except in immunocompromised hosts. Hepatitis, interstitial nephritis, and the hemolytic-uremic syndrome occasionally complicate acute infection. Reactive arthritis and other rheumatologic complaints may develop several weeks after infection, especially in persons with the HLA-B27 phenotype. Guillain-Barré syndrome or its Miller Fisher (cranial polyneuropathy) variant follows *Campylobacter* infections uncommonly—i.e., in 1 of every 1000–2000 cases or, for certain *C. jejuni* serotypes (such as O19), in 1 of every 100–200 cases. Despite the low frequency of this complication, it is now estimated that *Campylobacter* infections, because of their high incidence, may trigger 20–40% of all cases of Guillain-Barré syndrome. The presence of sialylated lipopolysaccharides on *C. jejuni* strains is a form of molecular mimicry that promotes autoimmune recognition of sialylated cell surface molecules on axons. Asymptomatic *Campylobacter* infection also may trigger Guillain-Barré syndrome. Immunoproliferative small-intestinal disease (*alpha chain disease*), a form of lymphoma that originates in small-intestinal mucosa-associated lymphoid tissue, has been associated with *C. jejuni*; antimicrobial therapy has led to marked clinical improvement.

DIAGNOSIS

In patients with *Campylobacter* enteritis, peripheral leukocyte counts reflect the severity of the inflammatory process. However, stools from nearly all *Campylobacter*-infected patients presenting for medical attention in the United States contain leukocytes or erythrocytes. Gram- or Wright-stained fecal smears should be examined in all suspected cases. When the diagnosis of *Campylobacter* enteritis is suspected on the basis of findings indicating inflammatory diarrhea (fever, fecal leukocytes), clinicians can ask the microbiology laboratory to attempt the visualization of organisms with characteristic vibrioid morphology by direct microscopic examination of stools with Gram's staining or to use phase-contrast or dark-field microscopy to identify the organisms' characteristic "darting" motility. Confirmation of the diagnosis of *Campylobacter* infection is based on identification of an isolate from cultures of stool, blood, or another site. *Campylobacter*-specific media should be used to culture stools from all patients with inflammatory or bloody diarrhea. Because all *Campylobacter* species are fastidious, they will not be isolated unless selective media or other selective techniques are used. Not all media are equally useful for isolation of the broad array of campylobacters; therefore, failure to isolate campylobacters from stool does not entirely rule out their presence. Species-specific polymerase chain reaction techniques have been developed to facilitate exact diagnoses. The detection of the organisms in stool almost always implies infection; there is a brief period of post-convalescent fecal carriage and no obvious commensalism in humans. In contrast, *Campylobacter sputorum* and related organisms found in the oral cavity are commensals that only rarely have pathogenic significance. Because of the low levels of metabolic activity of *Campylobacter* species in standard blood culture media, *Campylobacter* bacteremia may be difficult to detect unless laboratorians check for low-positive results in quantitative assays.

DIFFERENTIAL DIAGNOSIS

The symptoms of *Campylobacter* enteritis are not sufficiently unusual to distinguish this illness from that due to *Salmonella*, *Shigella*, *Yersinia*, and other pathogens. The combination of fever and fecal

193 Cholera and Other Vibrioses

Matthew K. Waldor, Edward T. Ryan

leukocytes or erythrocytes is indicative of inflammatory diarrhea, and definitive diagnosis is based on culture or demonstration of the characteristic organisms on stained fecal smears. Similarly, extraintestinal *Campylobacter* illness is diagnosed by culture. Infection due to *Campylobacter* should be suspected in the setting of septic abortion, and that due to *C. fetus* should be suspected specifically in the setting of septic thrombophlebitis. It is important to reiterate that (1) the presentation of *Campylobacter* enteritis may mimic that of ulcerative colitis or Crohn's disease, (2) *Campylobacter* enteritis is much more common than either of the latter (especially among young adults), and (3) biopsy may not distinguish among these entities. Thus a diagnosis of inflammatory bowel disease should not be made until *Campylobacter* infection has been ruled out, especially in persons with a history of foreign travel, significant animal contact, immunodeficiency, or exposure incurring a high risk of transmission.

TREATMENT CAMPYLOBACTER INFECTION

Fluid and electrolyte replacement is central to the treatment of diarrheal illnesses (Chap. 160). Even among patients presenting for medical attention with *Campylobacter* enteritis, not all clearly benefit from specific antimicrobial therapy. Indications for therapy include high fever, bloody diarrhea, severe diarrhea, persistence for >1 week, and worsening of symptoms. A 5- to 7-day course of erythromycin (250 mg orally four times daily or—for children—30–50 mg/kg per day, in divided doses) is the regimen of choice. Both clinical trials and in vitro susceptibility testing indicate that other macrolides, including azithromycin (a 1- or 3-day regimen), also are useful therapeutic agents. An alternative regimen for adults is ciprofloxacin (500 mg orally twice daily) or another fluoroquinolone for 5–7 days, but resistance to this class of agents as well as to tetracyclines is substantial; ~22% of U.S. isolates in 2010 were resistant to ciprofloxacin. Because macrolide resistance usually is much less common (<10%), these drugs are the empirical agents of choice. Patients infected with antibiotic-resistant strains are at increased risk of adverse outcomes. Use of antimotility agents, which may prolong the duration of symptoms and have been associated with toxic megacolon and with death, is not recommended.

For systemic infections, treatment with gentamicin (1.7 mg/kg IV every 8 h after a loading dose of 2 mg/kg), imipenem (500 mg IV every 6 h), or chloramphenicol (50 mg/kg IV each day in three or four divided doses) should be started empirically, but susceptibility testing should then be performed. Ciprofloxacin and amoxicillin-clavulanate are alternative agents for susceptible strains. In the absence of immunocompromise or endovascular infections, therapy should be administered for 14 days. For immunocompromised patients with systemic infections due to *C. fetus* and for patients with endovascular infections, prolonged therapy (for up to 4 weeks) is usually necessary. For recurrent infections in immunocompromised hosts, lifelong therapy/prophylaxis is sometimes necessary.

PROGNOSIS

Nearly all patients recover fully from *Campylobacter* enteritis, either spontaneously or after antimicrobial therapy. Volume depletion probably contributes to the few deaths that are reported. As stated above, occasional patients develop reactive arthritis or Guillain-Barré syndrome or its variants. Systemic infection with *C. fetus* is much more often fatal than that due to related species; this higher mortality rate reflects in part the population affected. Prognosis depends on the rapidity with which appropriate therapy is begun. Otherwise healthy hosts usually survive *C. fetus* infections without sequelae. Compromised hosts often have recurrent and/or life-threatening infections due to a variety of *Campylobacter* species.

Members of the genus *Vibrio* cause a number of important infectious syndromes. Classic among them is cholera, a devastating diarrheal disease caused by *Vibrio cholerae* that has been responsible for seven global pandemics and much suffering over the past two centuries. Epidemic cholera remains a significant public health concern in the developing world today. Other vibrios caused by other *Vibrio* species include syndromes of diarrhea, soft tissue infection, or primary sepsis. All *Vibrio* species are highly motile, facultatively anaerobic, curved gram-negative rods with one or more flagella. In nature, vibrios most commonly reside in tidal rivers and bays under conditions of moderate salinity. They proliferate in the summer months when water temperatures exceed 20°C. As might be expected, the illnesses they cause also increase in frequency during the warm months.

CHOLERA

DEFINITION

Cholera is an acute diarrheal disease that can, in a matter of hours, result in profound, rapidly progressive dehydration and death. Accordingly, *cholera gravis* (the severe form) is a much-feared disease, particularly in its epidemic presentation. Fortunately, prompt aggressive fluid repletion and supportive care can obviate the high mortality that is historically associated with cholera. Although the term *cholera* has occasionally been applied to any severely dehydrating secretory diarrheal illness, whether infectious in etiology or not, it now refers to disease caused by *V. cholerae* serogroup O1 or O139—i.e., the serogroups with epidemic potential.

MICROBIOLOGY AND EPIDEMIOLOGY

The species *V. cholerae* is classified into more than 200 serogroups based on the carbohydrate determinants of their lipopolysaccharide (LPS) O antigens. Although some non-O1 *V. cholerae* serogroups (strains that do not agglutinate in antisera to the O1 group antigen) have occasionally caused sporadic outbreaks of diarrhea, serogroup O1 was, until the emergence of serogroup O139 in 1992, the exclusive cause of epidemic cholera. Two biotypes of *V. cholerae* O1, classical and El Tor, are distinguished. Each biotype is further subdivided into two serotypes, termed *Inaba* and *Ogawa*.

The natural habitat of *V. cholerae* is coastal salt water and brackish estuaries, where the organism lives in close relation to plankton. *V. cholerae* can also exist in freshwater in the presence of adequate nutrients and warmth. Humans become infected incidentally but, once infected, can act as vehicles for spread. Ingestion of water contaminated by human feces is the most common means of acquisition of *V. cholerae*. Consumption of contaminated food also can contribute to spread. There is no known animal reservoir. Although the infectious dose is relatively high, it is markedly reduced in hypochlorhydric persons, in those using antacids, and when gastric acidity is buffered by a meal. Cholera is predominantly a pediatric disease in endemic areas, but it affects adults and children equally when newly introduced into a population. In endemic areas, the burden of disease is often greatest during “cholera seasons” associated with high temperatures, heavy rainfall, and flooding, but cholera can occur year-round. For unexplained reasons, susceptibility to cholera is significantly influenced by ABO blood group status; persons with type O blood are at greatest risk of severe disease if infected, whereas those with type AB are at least risk.



Cholera is native to the Ganges delta in the Indian subcontinent. Since 1817, seven global pandemics have occurred. The current (seventh) pandemic—the first due to the El Tor biotype—began in Indonesia in 1961 and spread in serial waves throughout Asia as *V. cholerae* El Tor displaced the endemic classical biotype, which is thought to have caused the previous six pandemics. In the early 1970s, El Tor cholera erupted in Africa, causing major epidemics

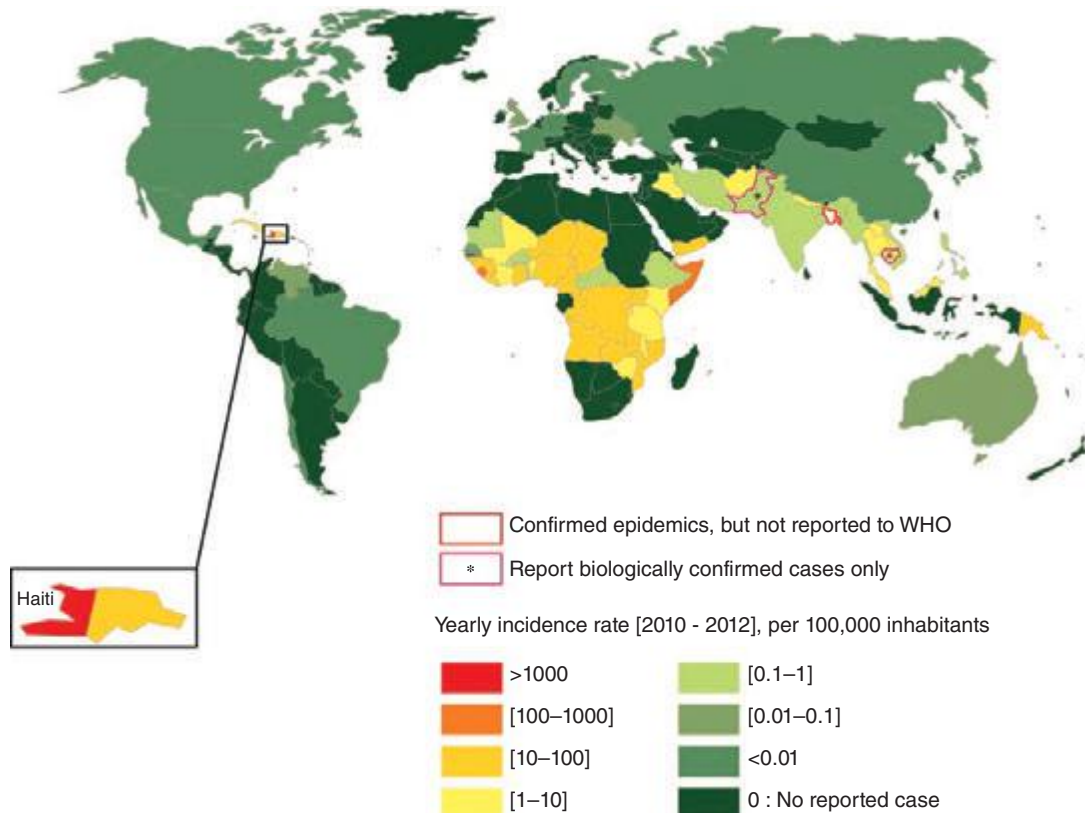


FIGURE 193-1 World distribution of cholera in 2010–2012. WHO, World Health Organization. (Courtesy of Drs. M. and R. Piarroux, Université de la Méditerranée, France; with permission.)

before becoming a persistent endemic problem. Currently, >90% of cholera cases reported annually to the World Health Organization (WHO) are from Africa (Fig. 193-1), but the true burden in Africa as well as in Asia is unknown because diagnosis is often syndromic and because many countries with endemic cholera do not report cholera to the WHO. It is possible that >3 million cases of cholera occur yearly (of which only ~200,000 are reported to the WHO), resulting in >100,000 deaths annually (of which <5000 are reported to the WHO).

After a century without cholera in Latin America, the current cholera pandemic reached Central and South America in 1991. Following an initial explosive spread that affected millions, the burden of disease has markedly decreased in Latin America. In 2010, a severe cholera outbreak began in Haiti, a country with no recorded history of this disease. Several lines of evidence indicate that cholera was likely introduced into Haiti by United Nations security forces from Asia, raising the possibility that asymptomatic carriers of *V. cholerae* play an important role in transmitting cholera over long distances. To date, the outbreak has involved more than 700,000 individuals, resulting in thousands of deaths. The recent history of cholera has been punctuated by such severe outbreaks, especially among impoverished or displaced persons. These outbreaks are often precipitated by war or other circumstances that lead to the breakdown of public health measures. Such was the case in the camps for Rwandan refugees set up in 1994 around Goma, Zaire, and in 2008–2009 in Zimbabwe.

Sporadic endemic infections due to *V. cholerae* O1 strains related to the seventh-pandemic strain have been recognized along the U.S. Gulf Coast of Louisiana and Texas. These infections are typically associated with the consumption of contaminated, locally harvested shellfish. Occasionally, cases in U.S. locations remote from the Gulf Coast have been linked to shipped-in Gulf Coast seafood.

In October 1992, a large-scale outbreak of clinical cholera caused by a new serogroup, O139, occurred in southeastern India. The organism appears to be a derivative of El Tor O1 but has a distinct LPS and an immunologically related O-antigen polysaccharide capsule. (O1 organisms are not encapsulated.) After an initial spread across 11 Asian countries, *V. cholerae* O139 has once again been almost entirely


replaced by O1 strains. The clinical manifestations of disease caused by *V. cholerae* O139 are indistinguishable from those of O1 cholera. Immunity to one, however, is not protective against the other.

PATHOGENESIS

In the final analysis, cholera is a toxin-mediated disease. The watery diarrhea characteristic of cholera is due to the action of cholera toxin, a potent protein enterotoxin elaborated by the organism in the small intestine. The toxin-coregulated pilus (TCP), so named because its synthesis is regulated in parallel with that of cholera toxin, is essential for *V. cholerae* to survive and multiply in (colonize) the small intestine. Cholera toxin, TCP, and several other virulence factors are coordinately regulated by ToxR. This protein modulates the expression of genes coding for virulence factors in response to environmental signals via a cascade of regulatory proteins. Additional regulatory processes, including bacterial responses to the density of the bacterial population (in a phenomenon known as *quorum sensing*), modulate the virulence of *V. cholerae*.

Once established in the human small bowel, the organism produces cholera toxin, which consists of a monomeric enzymatic moiety (the A subunit) and a pentameric binding moiety (the B subunit). The B pentamer binds to GM₁ ganglioside, a glycolipid on the surface of epithelial cells that serves as the toxin receptor and makes possible the delivery of the A subunit to its cytosolic target. The activated A subunit (A₁) irreversibly transfers ADP-ribose from nicotinamide adenine dinucleotide to its specific target protein, the GTP-binding regulatory component of adenylate cyclase. The ADP-ribosylated G protein upregulates the activity of adenylate cyclase; the result is the intracellular accumulation of high levels of cyclic AMP. In intestinal epithelial cells, cyclic AMP inhibits the absorptive sodium transport system in villus cells and activates the secretory chloride transport system in crypt cells, and these events lead to the accumulation of sodium chloride in the intestinal lumen. Because water moves passively to maintain osmolality, isotonic fluid accumulates in the lumen. When the volume of that fluid exceeds the capacity of the rest of the gut to resorb it, watery diarrhea results. Unless the wasted fluid and electrolytes are adequately replaced, shock (due to

profound dehydration) and acidosis (due to loss of bicarbonate) follow. Although perturbation of the adenylate cyclase pathway is the primary mechanism by which cholera toxin causes excess fluid secretion, cholera toxin also enhances intestinal secretion via prostaglandins and/or neural histamine receptors.

 The *V. cholerae* genome comprises two circular chromosomes. Lateral gene transfer has played a key role in the evolution of epidemic *V. cholerae*. The genes encoding cholera toxin (*ctxAB*) are part of the genome of a bacteriophage, CTXΦ. The receptor for this phage on the *V. cholerae* surface is the intestinal colonization factor TCP. Because *ctxAB* is part of a mobile genetic element (CTXΦ), horizontal transfer of this bacteriophage may account for the emergence of new toxigenic *V. cholerae* serogroups. Many of the other genes important for *V. cholerae* pathogenicity, including the genes encoding the biosynthesis of TCP, those encoding accessory colonization factors, and those regulating virulence gene expression, are clustered together in the *V. cholerae* pathogenicity island. Similar clustering of virulence genes is found in other bacterial pathogens. It is believed that pathogenicity islands are acquired by horizontal gene transfer. *V. cholerae* O139 is probably derived from an El Tor O1 strain that acquired the genes for O139 O-antigen synthesis by horizontal gene transfer.

CLINICAL MANIFESTATIONS

Individuals infected with *V. cholerae* O1 or O139 exhibit a range of clinical manifestations. Some individuals are asymptomatic or have only mild diarrhea; others present with the sudden onset of explosive and life-threatening diarrhea (*cholera gravis*). The reasons for the range in signs and symptoms of disease are incompletely understood but include the level of preexisting immunity, blood type, and nutritional status. In a nonimmune individual, after a 24- to 48-h incubation period, cholera characteristically begins with the sudden onset of painless watery diarrhea that may quickly become voluminous. Patients often vomit. In severe cases, volume loss can exceed 250 mL/kg in the first 24 h. If fluids and electrolytes are not replaced, hypovolemic shock and death may ensue. Fever is usually absent. Muscle cramps due to electrolyte disturbances are common. The stool has a characteristic appearance: a nonbilious, gray, slightly cloudy fluid with flecks of mucus, no blood, and a somewhat fishy, inoffensive odor. It has been called “rice-water” stool because of its resemblance to the water in which rice has been washed (Fig. 193-2). Clinical symptoms



FIGURE 193-2 Rice water cholera stool. Note floating mucus and gray watery appearance. (Courtesy of Dr. A. S. G. Faruque, International Centre for Diarrhoeal Disease Research, Dhaka; with permission.)

parallel volume contraction: at losses of <5% of normal body weight, thirst develops; at 5–10%, postural hypotension, weakness, tachycardia, and decreased skin turgor are documented; and at >10%, oliguria, weak or absent pulses, sunken eyes (and, in infants, sunken fontanelles), wrinkled (“washerwoman”) skin, somnolence, and coma are characteristic. Complications derive exclusively from the effects of volume and electrolyte depletion and include renal failure due to acute tubular necrosis. Thus, if the patient is adequately treated with fluid and electrolytes, complications are averted and the process is self-limited, resolving in a few days.

Laboratory data usually reveal an elevated hematocrit (due to hemoconcentration) in nonanemic patients; mild neutrophilic leukocytosis; elevated levels of blood urea nitrogen and creatinine consistent with prerenal azotemia; normal sodium, potassium, and chloride levels; a markedly reduced bicarbonate level (<15 mmol/L); and an elevated anion gap (due to increases in serum lactate, protein, and phosphate). Arterial pH is usually low (~7.2).

DIAGNOSIS

Cholera should be suspected when a patient ≥ 2 years of age develops acute watery diarrhea in an area known to have cholera or when a patient ≥ 5 years of age develops severe dehydration or dies from acute watery diarrhea, even in an area where cholera is not known to be present. The clinical suspicion of cholera can be confirmed by the identification of *V. cholerae* in stool; however, the organism must be specifically sought. With experience, it can be detected directly by dark-field microscopy on a wet mount of fresh stool, and its serotype can be discerned by immobilization with specific antiserum. Laboratory isolation of the organism requires the use of a selective medium such as taurocholate-tellurite-gelatin (TTG) agar or thiosulfate–citrate–bile salts–sucrose (TCBS) agar. If a delay in sample processing is expected, Carey-Blair transport medium and/or alkaline-peptone water-enrichment medium may be used as well. In endemic areas, there is little need for biochemical confirmation and characterization, although these tasks may be worthwhile in places where *V. cholerae* is an uncommon isolate. Standard microbiologic biochemical testing for Enterobacteriaceae will suffice for identification of *V. cholerae*. All vibrios are oxidase-positive. A point-of-care antigen-detection cholera dipstick assay is now commercially available for use in the field or where laboratory facilities are lacking.

TREATMENT CHOLERA

Death from cholera is due to hypovolemic shock; thus treatment of individuals with cholera first and foremost requires fluid resuscitation and management. In light of the level of dehydration (Table 193-1) and the patient’s age and weight, euvoemia should first be rapidly restored, and adequate hydration should then be maintained to replace ongoing fluid losses (Table 193-2). Administration of oral rehydration solution (ORS) takes advantage of the hexose- Na^+ co-transport mechanism to move Na^+ across the gut mucosa together with an actively transported molecule such as glucose (or galactose). Cl^- and water follow. This transport mechanism remains intact even when cholera toxin is active. ORS may be made by adding safe water to prepackaged sachets containing salts and sugar or by adding 0.5 teaspoon of table salt and 6 teaspoons of table sugar to 1 L of safe water. Potassium intake in bananas or green coconut water should be encouraged. A number of ORS formulations are available, and the WHO now recommends “low-osmolarity” ORS for treatment of individuals with dehydrating diarrhea of any cause (Table 193-3). If available, rice-based ORS is considered superior to standard ORS in the treatment of cholera. ORS can be administered via a nasogastric tube to individuals who cannot ingest fluid; however, optimal management of individuals with severe dehydration includes the administration of IV fluid and electrolytes. Because profound acidosis ($\text{pH} < 7.2$) is common in this group, Ringer’s lactate is the best choice among commercial products (Table 193-4); it must be used with additional potassium supplements, preferably given by mouth.

TABLE 193-1 ASSESSING THE DEGREE OF DEHYDRATION IN PATIENTS WITH CHOLERA

Degree of Dehydration	Clinical Findings
None or mild, but diarrhea	Thirst in some cases; <5% loss of total body weight
Moderate	Thirst, postural hypotension, weakness, tachycardia, decreased skin turgor, dry mouth/tongue, no tears; 5–10% loss of total body weight
Severe	Unconsciousness, lethargy, or “floppiness”; weak or absent pulse; inability to drink; sunken eyes (and, in infants, sunken fontanelles); >10% loss of total body weight

TABLE 193-2 TREATMENT OF CHOLERA, BASED ON DEGREE OF DEHYDRATION^a

Degree of Dehydration, Patient's Age (Weight)	Treatment ^b
None or Mild, But Diarrhea^c	
<2 years	1/4–1/2 cup (50–100 mL) of ORS, to a maximum of 0.5 L/d
2–9 years	1/2–1 cup (100–200 mL) of ORS, to a maximum of 1 L/d
≥10 years	As much ORS as desired, to a maximum of 2 L/d
Moderate^{c,d}	
<4 months (<5 kg)	200–400 mL of ORS
4–11 months (5–<8 kg)	400–600 mL of ORS
12–23 months (8–<11 kg)	600–800 mL of ORS
2–4 years (11–<16 kg)	800–1200 mL of ORS
5–14 years (16–<30 kg)	1200–2200 mL of ORS
≥15 years (≥30 kg)	2200–4000 mL of ORS
Severe^e	
All ages and weights	Undertake IV fluid replacement with Ringer's lactate (or, if not available, normal saline). Give 100 mL/kg in the first 3-h period (or the first 6-h period for children <12 months old); start rapidly, then slow down. Give a total of 200 mL/kg in the first 24 h. Continue until the patient is awake, can ingest ORS, and no longer has a weak pulse.

^aAdapted from World Health Organization: First steps for managing an outbreak of acute diarrhoea. Global Task Force on Cholera Control, 2009 (www.who.int/topics/cholera).

^bContinue normal feeding during treatment. ^cReassess regularly; monitor stool and vomit output. ^dVolumes of ORS listed should be given within the first 4 h.

Abbreviation: ORS, oral rehydration solution.

TABLE 193-3 COMPOSITION OF WORLD HEALTH ORGANIZATION REDUCED-OSMOLARITY ORAL REHYDRATION SOLUTION (ORS)^{a,b}

Constituent	Concentration, mmol/L
Na ⁺	75
K ⁺	20
Cl ⁻	65
Citrate ^c	10
Glucose	75
Total osmolarity	245

^aContains (per package, to be added to 1 L of drinking water): NaCl, 2.6 g; Na₃C₆H₅O₇·2H₂O, 2.9 g; KCl, 1.5 g; and glucose (anhydrous), 13.5 g. ^bIf prepackaged ORS is unavailable, a simple homemade alternative can be prepared by combining 3.5 g (~1/2 teaspoon) of NaCl with either 50 g of precooked rice cereal or 6 teaspoons of table sugar (sucrose) in 1 L of drinking water. In that case, potassium must be supplied separately (e.g., in orange juice or coconut water). ^c10 mmol of citrate per liter, which supplies 30 mmol HCO₃⁻/L.

TABLE 193-4 ELECTROLYTE COMPOSITION OF CHOLERA STOOL AND OF INTRAVENOUS REHYDRATION SOLUTION

Substance	Concentration, mmol/L			
	Na ⁺	K ⁺	Cl ⁻	Base
Stool				
Adult	135	15	100	45
Child	100	25	90	30
Ringer's lactate	130	4 ^a	109	28

^aPotassium supplements, preferably administered by mouth, are required to replace the usual potassium losses from stool.

The total fluid deficit in severely dehydrated patients (>10% of body weight) can be replaced safely within the first 3–4 h of therapy, half within the first hour. Transient muscle cramps and tetany are common. Thereafter, oral therapy can usually be initiated, with the goal of maintaining fluid intake equal to fluid output. However, patients with continued large-volume diarrhea may require prolonged IV treatment to match gastrointestinal fluid losses. Severe hypokalemia can develop but will respond to potassium given either IV or orally. In the absence of adequate staff to monitor the patient's progress, the oral route of rehydration and potassium replacement is safer than the IV route.

Although not necessary for cure, the use of an antibiotic to which the organism is susceptible diminishes the duration and volume of fluid loss and hastens clearance of the organism from the stool. Adjunctive antibiotics should therefore be administered to patients with moderate or severe dehydration due to cholera. In many areas, macrolides such as erythromycin (adults, 250 mg orally four times a day for 3 days; children, 12.5 mg/kg per dose four times a day for 3 days) or azithromycin (adults, a single 1-g dose; children, a single 20-mg/kg dose) are the agents of choice. Increasing resistance to tetracyclines is widespread; however, in areas with confirmed susceptibility, tetracycline (nonpregnant adults, 500 mg orally four times a day for 3 days; children >8 years old, 12.5 mg/kg per dose four times a day for 3 days) or doxycycline (nonpregnant adults, a 300-mg single dose; children >8 years old, a single dose of 4–6 mg/kg) may be used. Similarly, increasing resistance to fluoroquinolones is being reported, but in areas with confirmed susceptibility, a fluoroquinolone such as ciprofloxacin may be used (adults, 500 mg twice a day for 3 days; children, 15 mg/kg twice a day for 3 days).

PREVENTION

Provision of safe water and of facilities for sanitary disposal of feces, improved nutrition, and attention to food preparation and storage in the household can significantly reduce the incidence of cholera. In addition, precautions should be taken to prevent the spread of cholera via infected and potentially asymptomatic persons from endemic to nonendemic regions of the world (as was probably the case in the ongoing outbreak in Haiti; see “Microbiology and Epidemiology,” above).

Much effort has been devoted to the development of an effective cholera vaccine over the past few decades, with a particular focus on oral vaccine strains. In an attempt to maximize mucosal responses, two types of oral cholera vaccine have been developed: oral killed vaccines and live attenuated vaccines. Currently, two oral killed cholera vaccines have been prequalified by the WHO and are available internationally. WC-rBS (Dukoral[®]; Crucell, Stockholm, Sweden) contains several biotypes and serotypes of *V. cholerae* O1 supplemented with 1 mg of recombinant cholera toxin B subunit per dose. BivWC (Shanchol[™]; Shantha Biotechnics–Sanofi Pasteur, Mumbai, India) contains several biotypes and serotypes of *V. cholerae* O1 and *V. cholerae* O139 without supplemental cholera toxin B subunit. The vaccines are administered as a two- or three-dose regimen, with

doses usually separated by 14 days. They provide ~60–85% protection for the first few months. Booster immunizations of WC-rBS are recommended after 2 years for individuals ≥ 6 years of age and after 6 months for children 2–5 years of age. For BivWC, which was developed more recently, no formal recommendation regarding booster immunizations exists. However, BivWC was associated with ~60% protection over 5 years among recipients of all ages in a study in Kolkata, India; the rate of protection among children ≤ 5 years of age approximated 40%. Models predict significant herd immunity when vaccination coverage rates exceed 50%. The killed vaccines have been safely administered among populations with high rates of HIV.

Oral live attenuated vaccines for *V. cholerae* are also in development. These strains have in common the fact that they lack the genes encoding cholera toxin. One such vaccine, CVD 103-HgR, was safe and immunogenic in phase 1 and 2 studies but afforded minimal protection in a large field trial in Indonesia. Other live attenuated vaccine candidate strains have been prepared from El Tor and O139 *V. cholerae* and have been tested in studies of volunteers. A possible advantage of live attenuated cholera vaccines is that they may induce protection after a single oral dose. Conjugate and subunit cholera vaccines are also being developed. Recognizing that it may be decades before safe water and adequate sanitation become a reality for those most at risk of cholera, the WHO has now recommended incorporation of cholera vaccination into comprehensive control strategies and has established an international stockpile of oral killed cholera vaccine to assist in outbreak responses. No cholera vaccine is commercially available in the United States.

OTHER VIBRIO SPECIES



The genus *Vibrio* includes several human pathogens that do not cause cholera. Abundant in coastal waters throughout the world, noncholera vibrios can reach high concentrations in the tissues of filter-feeding mollusks. As a result, human infection commonly follows the ingestion of seawater or of raw or undercooked shellfish (Table 193-5). Most noncholera vibrios can be cultured on blood or MacConkey agar, which contains enough salt to support the growth of these halophilic species. In the microbiology laboratory, the species of noncholera vibrios are distinguished by standard biochemical tests. The most important of these organisms are *Vibrio parahaemolyticus* and *Vibrio vulnificus*.

The two major types of syndromes for which these species are responsible are gastrointestinal illness (due to *V. parahaemolyticus*, non-O1/O139 *V. cholerae*, *Vibrio mimicus*, *Vibrio fluvialis*, *Vibrio hollisae*, and *Vibrio furnissii*) and soft tissue infections (due to *V. vulnificus*, *Vibrio alginolyticus*, and *Vibrio damsela*). *V. vulnificus* is also a cause of primary sepsis in some compromised individuals.

SPECIES ASSOCIATED PRIMARILY WITH GASTROINTESTINAL ILLNESS



V. parahaemolyticus Widespread in marine environments, the halophilic *V. parahaemolyticus* causes food-borne enteritis worldwide. This species was originally implicated in enteritis

in Japan in 1953, accounting for 24% of reported cases in one study—a rate that presumably was due to the common practice of eating raw seafood in that country. In the United States, common-source outbreaks of diarrhea caused by this organism have been linked to the consumption of undercooked or improperly handled seafood or of other foods contaminated by seawater. Since the mid-1990s, the incidence of *V. parahaemolyticus* infections has increased in several countries, including the United States. Serotypes O3:K6, O4:K68, and O1:K-untypable, which are genetically related to one another, account in part for this increase. Serotypes O4:K12 and O4:KUT were initially unique to the Pacific Northwest but caused recent outbreaks in the eastern United States and Spain. The enteropathogenicity of *V. parahaemolyticus* is linked to its ability to cause hemolysis on Wagatsuma agar (i.e., the *Kanagawa phenomenon*). Although the mechanisms by which the organism causes diarrhea are not fully defined, the genome sequence of *V. parahaemolyticus* contains two type III secretion systems, which directly inject toxic bacterial proteins into host cells. The activity of one of these secretion systems is required for intestinal colonization and virulence in animal models. *V. parahaemolyticus* should be considered a possible etiologic agent in all cases of diarrhea that can be linked epidemiologically to seafood consumption or to the sea itself.

Infections with *V. parahaemolyticus* can result in two distinct gastrointestinal presentations. The more common of the two presentations (including nearly all cases in North America) is characterized by watery diarrhea, usually occurring in conjunction with abdominal cramps, nausea, and vomiting and accompanied in ~25% of cases by fever and chills. After an incubation period of 4 h to 4 days, symptoms develop and persist for a median of 3 days. Dysentery, the less common presentation, is characterized by severe abdominal cramps, nausea, vomiting, and bloody or mucoid stools. *V. parahaemolyticus* also causes rare cases of wound infection and otitis and very rare cases of sepsis.

Most cases of *V. parahaemolyticus*-associated gastrointestinal illness, regardless of the presentation, are self-limited. Fluid replacement should be stressed. The role of antimicrobials is uncertain, but they may be of benefit in moderate or severe disease. Doxycycline, fluoroquinolones, or macrolides are usually used. Deaths are extremely rare among immunocompetent individuals. Severe infections are associated with underlying diseases, including diabetes, preexisting liver disease, iron-overload states, or immunosuppression.

Non-O1/O139 (Noncholera) *V. cholerae* The heterogeneous non-O1/O139 *V. cholerae* organisms cannot be distinguished from *V. cholerae* O1 or O139 by routine biochemical tests but do not agglutinate in O1 or O139 antiserum. Non-O1/O139 strains have caused several well-studied food-borne outbreaks of gastroenteritis and have also been responsible for sporadic cases of otitis media, wound infection, and bacteremia; although gastroenteritis outbreaks can occur, non-O1/O139 *V. cholerae* strains do not cause epidemics of cholera. Like other vibrios, non-O1/O139 *V. cholerae* organisms are widely distributed in marine environments. In most instances, recognized cases in the United States have been associated with the consumption of raw oysters or with recent travel. The broad clinical spectrum of diarrheal illness caused by these

TABLE 193-5 FEATURES OF SELECTED NONCHOLERA VIBRIOSES

Organism	Vehicle or Activity	Host at Risk	Syndrome
<i>Vibrio parahaemolyticus</i>	Shellfish, seawater	Normal	Gastroenteritis
	Seawater	Normal	Wound infection
Non-O1/O139 <i>Vibrio cholerae</i>	Shellfish, travel	Normal	Gastroenteritis
	Seawater	Normal	Wound infection, otitis media
<i>Vibrio vulnificus</i>	Shellfish	Immunosuppressed ^a	Sepsis, secondary cellulitis
	Seawater	Normal, immunosuppressed ^a	Wound infection, cellulitis
<i>Vibrio alginolyticus</i>	Seawater	Normal	Wound infection, cellulitis, otitis
	Seawater	Burned, other immunosuppressed	Sepsis

^aEspecially with liver disease or hemochromatosis.

Source: Table 161-3 in *Harrisons Principles of Internal Medicine*, 14th edition.

1066 organisms is probably due to the group's heterogeneous virulence attributes.

In the United States, about half of all non-O1/O139 *V. cholerae* isolates are from stool samples. The typical incubation period for gastroenteritis due to these organisms is <2 days, and the illness lasts for ~2–7 days. Patients' stools may be copious and watery or may be partly formed, less voluminous, and bloody or mucoid. Diarrhea can result in severe dehydration. Many cases include abdominal cramps, nausea, vomiting, and fever. Like those with cholera, patients who are seriously dehydrated should receive oral or IV fluids; the value of antibiotics is not clear.

Extraintestinal infections due to non-O1/O139 *V. cholerae* commonly follow occupational or recreational exposure to seawater. Around 10% of non-O1/O139 *V. cholerae* isolates come from cases of wound infection, 10% from cases of otitis media, and 20% from cases of bacteremia (which is particularly likely to develop in patients with liver disease). Extraintestinal infections should be treated with antibiotics. Information to guide antibiotic selection and dosing is limited, but most strains are sensitive in vitro to tetracycline, ciprofloxacin, and third-generation cephalosporins.

SPECIES ASSOCIATED PRIMARILY WITH SOFT TISSUE INFECTION OR BACTEREMIA

(See also Chap. 156)

V. vulnificus Infection with *V. vulnificus* is rare, but this organism is the most common cause of severe *Vibrio* infections in the United States. Like most vibrios, *V. vulnificus* proliferates in the warm summer months and requires a saline environment for growth. In the United States, infections in humans typically occur in coastal states between May and October and most commonly affect men >40 years of age. *V. vulnificus* has been linked to two distinct syndromes: primary sepsis, which usually occurs in patients with underlying liver disease, and primary wound infection, which generally affects people without underlying disease. (*Vulnificus* is Latin for “wound maker.”) Some authors have suggested that *V. vulnificus* also causes gastroenteritis independent of other clinical manifestations. *V. vulnificus* is endowed with a number of virulence attributes, including a capsule that confers resistance to phagocytosis and to the bactericidal activity of human serum as well as a cytotoxin. Measured as the 50% lethal dose in mice, the organism's virulence is considerably increased under conditions of iron overload; this observation is consistent with the propensity of *V. vulnificus* to infect patients who have hemochromatosis.

Primary sepsis most often develops in patients who have cirrhosis or hemochromatosis. However, *V. vulnificus* bacteremia can also affect individuals who have hematopoietic disorders or chronic renal insufficiency, those who are using immunosuppressive medications or alcohol, or (in rare instances) those who have no known underlying disease. After a median incubation period of 16 h, the patient develops malaise, chills, fever, and prostration. One-third of patients develop hypotension, which is often apparent at admission. Cutaneous manifestations develop in most cases (usually within 36 h of onset) and characteristically involve the extremities (the lower more often than the upper). In a common sequence, erythematous patches are followed by ecchymoses, vesicles, and bullae. In fact, sepsis and hemorrhagic bullous skin lesions suggest the diagnosis in appropriate settings. Necrosis and sloughing may also be evident. Laboratory studies reveal leukopenia more often than leukocytosis, thrombocytopenia, or elevated levels of fibrin split products. *V. vulnificus* can be cultured from blood or cutaneous lesions. The mortality rate approaches 50%, with most deaths due to uncontrolled sepsis (Chap. 325). Accordingly, prompt treatment is critical and should include empirical antibiotic administration, aggressive debridement, and general supportive care. *V. vulnificus* is sensitive in vitro to a number of antibiotics, including tetracycline, fluoroquinolones, and third-generation cephalosporins. Data from animal models suggest that either a fluoroquinolone or the combination of a tetracycline and a third-generation cephalosporin should be used in the treatment of *V. vulnificus* septicemia.

V. vulnificus-associated soft tissue infection can complicate either a fresh or an old wound that comes into contact with seawater; the patient may or may not have underlying disease. After a short incubation period (4 h to 4 days; mean, 12 h), the disease begins with swelling, erythema, and (in many cases) intense pain around the wound. These signs and symptoms are followed by cellulitis, which spreads rapidly and is sometimes accompanied by vesicular, bullous, or necrotic lesions. Metastatic events are uncommon. Most patients have fever and leukocytosis. *V. vulnificus* can be cultured from skin lesions and occasionally from the blood. Prompt antibiotic therapy and debridement are usually curative.

V. alginolyticus First identified as a pathogen of humans in 1973, *V. alginolyticus* occasionally causes eye, ear, and wound infections. This species is the most salt-tolerant of the vibrios and can grow in salt concentrations of >10%. Most clinical isolates come from superinfected wounds that presumably become contaminated at the beach. Although its severity varies, *V. alginolyticus* infection tends not to be serious and generally responds well to antibiotic therapy and drainage. A few cases of otitis externa, otitis media, and conjunctivitis due to this pathogen have been described. Tetracycline treatment usually results in cure. *V. alginolyticus* is a rare cause of bacteremia in immunocompromised hosts.

ACKNOWLEDGMENT

The authors gratefully acknowledge the valuable contributions of Drs. ≥Robert Deresiewicz and Gerald T. Keusch, coauthors of this chapter for previous editions.

DEFINITION

Brucellosis is a bacterial zoonosis transmitted directly or indirectly to humans from infected animals, predominantly domesticated ruminants and swine. The disease is known colloquially as *undulant fever* because of its remittent character. Although brucellosis commonly presents as an acute febrile illness, its clinical manifestations vary widely, and definitive signs indicative of the diagnosis may be lacking. Thus the clinical diagnosis usually must be supported by the results of bacteriologic and/or serologic tests.

ETIOLOGIC AGENTS

Human brucellosis is caused by strains of *Brucella*, a bacterial genus that was previously suggested, on genetic grounds, to comprise a single species, *B. melitensis*, with a number of biologic variants exhibiting particular host preferences. This view was challenged on the basis of detailed differences in chromosomal structure and host preference. The traditional classification into nomen species is now favored both because of these differences and because this classification scheme closely reflects the epidemiologic patterns of the infection. The nomen system recognizes *B. melitensis*, which is the most common cause of symptomatic disease in humans and for which the main sources are sheep, goats, and camels; *B. abortus*, which is usually acquired from cattle or buffalo; *B. suis*, which generally is acquired from swine but has one variant enzootic in reindeer and caribou and another in rodents; and *B. canis*, which is acquired most often from dogs. *B. ovis*, which causes reproductive disease in sheep, and *B. neotomae*, which is specific for desert rodents, have not been clearly implicated in human disease. Two new species, *B. ceti* and *B. pinnipedialis*, have recently been identified in marine mammals, including seals and dolphins. At least one case of laboratory-acquired human disease due to one of these species has been described, and apparent cases of natural human infection have been reported. As infections in marine mammals appear to be widespread, more cases of zoonotic infection in humans may be identified. Other newly reported species include *B. microti* (isolated from field voles) and *B. inopinata* (isolated from a patient with a breast implant). Additional novel strains have been described from diverse species, including baboons, foxes, frogs, and various rodents, and the genus likely will expand further in forthcoming years. Moreover, it has become apparent that *Brucella* is closely related to the genus *Ochrobactrum*, which includes environmental bacteria sometimes associated with opportunistic infections. Genomics-based studies are beginning to elucidate the pathway of evolution from free-living soil bacteria to highly successful intracellular pathogens.

All brucellae are small, gram-negative, unencapsulated, nonsporulating, nonmotile rods or coccobacilli. They grow aerobically on peptone-based medium incubated at 37°C; the growth of some types is improved by supplementary CO₂. In vivo, brucellae behave as facultative intracellular parasites. The organisms are sensitive to sunlight, ionizing radiation, and moderate heat; they are killed by boiling and pasteurization but are resistant to freezing and drying. Their resistance to drying renders brucellae stable in aerosol form, facilitating airborne transmission. The organisms can survive for up to 2 months in soft cheeses made from goat's or sheep's milk; for at least 6 weeks in dry soil contaminated with infected urine, vaginal discharge, or placental or fetal tissues; and for at least 6 months in damp soil or liquid manure kept under cool dark conditions. Brucellae are easily killed by a wide range of common disinfectants used under optimal conditions but are likely to be much more resistant at low temperatures or in the presence of heavy organic contamination.

EPIDEMIOLOGY

Brucellosis is a zoonosis whose occurrence and control are closely related to its prevalence in domesticated animals. Its distribution is worldwide apart from the few countries where

it has been eradicated from the animal reservoir. The true global prevalence of human brucellosis is unknown because of the imprecision of diagnosis and the inadequacy of reporting and surveillance systems in many countries. Recently, there has been increased recognition of the high incidence of brucellosis in India and parts of China and of importations to countries in Oceania, such as Fiji. In Europe, the incidence of brucellosis in a country is inversely related to gross domestic product, and, in both developed and less well-resourced settings, human brucellosis is related to rural poverty and inadequate access to medical care. Failure of veterinary control programs due to conflict or for economic reasons contributes further to the emergence and re-emergence of disease, as seen currently in some eastern Mediterranean countries.

Even in well-resourced settings, the true incidence of brucellosis in domesticated animals may be 10–20 times higher than the reported figures. Bovine brucellosis has been the target of control programs in many parts of the world and has been eradicated from the cattle populations of Australia, New Zealand, Bulgaria, Canada, Cyprus, Great Britain (including the Channel Islands), Japan, Luxembourg, Romania, the Scandinavian countries, Switzerland, and the Czech and Slovak Republics, among other nations. Its incidence has been reduced to a low level in the United States and most Western European countries, with a varied picture in other parts of the world. Efforts to eradicate *B. melitensis* infection from sheep and goat populations have been much less successful. These efforts have relied heavily on vaccination programs, which have tended to fluctuate with changing economic and political conditions. In some countries (e.g., Israel), *B. melitensis* has caused serious outbreaks in cattle. Infections with *B. melitensis* still pose a major public health problem in Mediterranean countries; in western, central, and southern Asia; and in parts of Africa and South and Central America.

Human brucellosis is usually associated with occupational or domestic exposure to infected animals or their products. Farmers, shepherds, goatherds, veterinarians, and employees in slaughterhouses and meat-processing plants in endemic areas are occupationally exposed to infection. Family members of individuals involved in animal husbandry may be at risk, although it is often difficult to differentiate food-borne infection from environmental contamination under these circumstances. Laboratory workers who handle cultures or infected samples also are at risk. Travelers and urban residents usually acquire the infection through consumption of contaminated foods. In countries that have eradicated the disease, new cases are most commonly acquired abroad. Dairy products, especially soft cheeses, unpasteurized milk, and ice cream, are the most frequently implicated sources of infection; raw meat and bone marrow may be sources under exceptional circumstances. Infections acquired through cosmetic treatments using materials of fetal origin have been reported. Person-to-person transmission is extremely rare, as is transfer of infection by blood or tissue donation. Although brucellosis is a chronic intracellular infection, there is no evidence for increased prevalence or severity among individuals with HIV infection or with immunodeficiency or immunosuppression of other etiologies.

Brucellosis may be acquired by ingestion, inhalation, or mucosal or percutaneous exposure. Accidental injection of the live vaccine strains of *B. abortus* (S19 and RB51) and *B. melitensis* (Rev 1) can cause disease. *B. melitensis* and *B. suis* have historically been developed as biological weapons by several countries and could be exploited for bioterrorism (Chap. 261e). This possibility should be borne in mind in the event of sudden unexplained outbreaks.

IMMUNITY AND PATHOGENESIS

Exposure to brucellosis elicits both humoral and cell-mediated immune responses. The mechanisms of protective immunity against human brucellosis are presumed to be similar to those documented in laboratory animals, but such generalizations must be interpreted with caution. The response to infection and its outcome are influenced by the virulence, phase, and species of the infecting strain. Differences have been reported between *B. abortus* and *B. suis* in modes of cellular entry and subsequent compartmentalization and processing.

Antibodies promote clearance of extracellular brucellae by bactericidal action and by facilitation of phagocytosis by polymorphonuclear and mononuclear phagocytes; however, antibodies alone cannot eradicate infection. Organisms taken up by macrophages and other cells can establish persistent intracellular infections. The key target cell is the macrophage, and bacterial mechanisms for suppressing intracellular killing and apoptosis result in very large intracellular populations. Opsonized bacteria are actively phagocytosed by neutrophilic granulocytes and by monocytes. In these and other cells, initial attachment takes place via specific receptors, including Fc, C3, fibronectin, and mannose-binding proteins. Opsonized—but not unopsonized—bacteria trigger an oxidative burst inside phagocytes. Unopsonized bacteria are internalized via similar receptors but at much lower efficiency. Smooth strains enter host cells via lipid rafts. Smooth lipopolysaccharide (LPS), β -cyclic glucan, and possibly an invasion-attachment protein (IalB) are involved in this process. Tumor necrosis factor α (TNF- α) produced early in the course of infection stimulates cytotoxic lymphocytes and activates macrophages, which can kill intracellular brucellae (probably mainly through production of reactive oxygen and nitrogen intermediates) and may clear infection. However, virulent *Brucella* cells can suppress the TNF- α response, and control of infection in this situation depends on macrophage activation and interferon γ (IFN- γ) responses. Cytokines such as interleukin (IL) 12 promote production of IFN- γ , which drives T_H1-type responses and stimulates macrophage activation. Inflammatory cytokines, including IL-4, IL-6, and IL-10, downregulate the protective response. As in other types of intracellular infection, it is assumed that initial replication of brucellae takes place within cells of the lymph nodes draining the point of entry. Subsequent hematogenous spread may result in chronic localizing infection at almost any site, although the reticuloendothelial system, musculoskeletal tissues, and genitourinary system are most frequently targeted. Both acute and chronic inflammatory responses develop in brucellosis, and the local tissue response may include granuloma formation with or without necrosis and caseation. Abscesses may also develop, especially in chronic localized infection.

The determinants of pathogenicity of *Brucella* have not been fully characterized, and the mechanisms underlying the manifestations of brucellosis are incompletely understood. The organism is a “stealth” pathogen whose survival strategy is centered on several processes that avoid triggering innate immune responses and that permit survival within monocytic cells. The smooth *Brucella* LPS, which has an unusual O-chain and core-lipid composition, has relatively low endotoxin activity and plays a key role in pyrogenicity and in resistance to phagocytosis and serum killing in the nonimmune host. In addition, LPS is believed to play a role in suppressing phagosome-lysosome fusion and diverting the internalized bacteria into vacuoles located in endoplasmic reticulum, where intracellular replication takes place. Specific exotoxins have not been isolated, but a type IV secretion system (VirB) that regulates intracellular survival and trafficking has been identified. In *B. abortus* this system can be activated extracellularly, but in *B. suis* it is activated (by low pH) only during intracellular growth. Brucellae then produce acid-stable proteins that facilitate the organisms’ survival in phagosomes and may enhance their resistance to reactive oxygen intermediates. A type III secretion system based on modified flagellar structures also has been inferred, although not yet confirmed. Virulent brucellae are resistant to defensins and produce a Cu-Zn superoxide dismutase that increases their resistance to reactive oxygen intermediates. A hemolysin-like protein may trigger the release of brucellae from infected cells.

CLINICAL FEATURES

Brucellosis almost invariably causes fever, which may be associated with profuse sweats, especially at night. In endemic areas, brucellosis may be difficult to distinguish from the many other causes of fever. However, two features recognized in the nineteenth century distinguish brucellosis from other tropical fevers, such as typhoid and malaria: (1) Left untreated, the fever of brucellosis shows an undulating pattern that persists for weeks before the commencement of an afebrile period that may be followed by relapse. (2) The fever of brucellosis is

associated with musculoskeletal symptoms and signs in about one-half of all patients.

The clinical syndromes caused by the different nomen species are similar, although *B. melitensis* tends to be associated with a more acute and aggressive presentation and *B. suis* with focal abscess induction. *B. abortus* infections may be more insidious in onset and more likely to become chronic. *B. canis* infections are reported to present frequently with acute gastrointestinal symptoms.

The incubation period varies from 1 week to several months, and the onset of fever and other symptoms may be abrupt or insidious. In addition to experiencing fever and sweats, patients become increasingly apathetic and fatigued; lose appetite and weight; and have nonspecific myalgia, headache, and chills. Overall, the presentation of brucellosis often fits one of three patterns: febrile illness that resembles typhoid but is less severe; fever and acute monoarthritis, typically of the hip or knee, in a young child; and long-lasting fever, misery, and low-back or hip pain in an older man. In an endemic area (e.g., much of the Middle East), a patient with fever and difficulty walking into the clinic would be regarded as having brucellosis until it was proven otherwise.

Diagnostic clues in the patient’s history include travel to an endemic area, employment in a diagnostic microbiology laboratory, consumption of unpasteurized milk products (including soft cheeses), contact with animals, accidental inoculation with veterinary *Brucella* vaccines, and—in an endemic setting—a history of similar illness in the family (documented in almost 50% of cases). Focal features are present in the majority of patients. The most common are musculoskeletal pain and physical findings in the peripheral and axial skeleton (~40% of cases). Osteomyelitis more commonly involves the lumbar and low thoracic vertebrae than the cervical and high thoracic spine. Individual joints that are most commonly affected by septic arthritis are the knee, hip, sacroiliac, shoulder, and sternoclavicular joints; the pattern may be one of monoarthritis or polyarthritis. Osteomyelitis may also accompany septic arthritis.

In addition to the usual causes of vertebral osteomyelitis or septic arthritis, the most important disease in the differential diagnosis is tuberculosis. This point influences the therapeutic approach as well as the prognosis, given that several antimicrobial agents used to treat brucellosis are also used to treat tuberculosis. Septic arthritis in brucellosis progresses slowly, starting with small pericapsular erosions. In the vertebrae, anterior erosions of the superior end plate are typically the first features to become evident, with eventual involvement and sclerosis of the whole vertebra. Anterior osteophytes eventually develop, but vertebral destruction or impingement on the spinal cord is rare and usually suggests tuberculosis (Table 194e-1).

Other systems may be involved in a manner that resembles typhoid. About one-quarter of patients have a dry cough, usually with few changes visible on the chest x-ray, although pneumonia, empyema,

TABLE 194e-1 RADIOLOGY OF THE SPINE: DIFFERENTIATION OF BRUCELOSIS FROM TUBERCULOSIS

	Brucellosis	Tuberculosis
Site	Lumbar and others	Dorsolumbar
Vertebrae	Multiple or contiguous	Contiguous
Diskitis	Late	Early
Body	Intact until late	Morphology lost early
Canal compression	Rare	Common
Epiphysitis	Anterosuperior (Pom’s sign)	General: upper and lower disk regions, central, subperiosteal
Osteophyte	Anterolateral (parrot beak)	Unusual
Deformity	Wedging uncommon	Anterior wedge, gibbus
Recovery	Sclerosis, whole-body	Variable
Paravertebral abscess	Small, well-localized	Common and discrete loss, transverse process
Psoas abscess	Rare	More likely

intrathoracic adenopathy, or lung abscess can occur. Sputum or pleural effusion cultures are rarely positive in such cases, which respond well to standard brucellosis treatment. One-quarter of patients have hepatosplenomegaly, and 10–20% have significant lymphadenopathy; the differential diagnosis includes glandular fever–like illness such as that caused by Epstein-Barr virus, *Toxoplasma*, cytomegalovirus, HIV, or *Mycobacterium tuberculosis*. Up to 10% of men have acute epididymo-orchitis, which must be distinguished from mumps and from surgical problems such as torsion. Prostatitis, inflammation of the seminal vesicles, salpingitis, and pyelonephritis all occur. There is an increased incidence of fetal loss among infected pregnant women, although teratogenicity has not been described and the tendency toward abortions is much less pronounced in humans than in farm animals.

Neurologic involvement is common, with depression and lethargy whose severity may not be fully appreciated by either the patient or the physician until after treatment. A small proportion of patients develop lymphocytic meningoencephalitis that mimics neurotuberculosis, atypical leptospirosis, or noninfectious conditions and that may be complicated by intracerebral abscess, a variety of cranial nerve deficits, or ruptured mycotic aneurysms.

Endocarditis occurs in ~1% of cases, most often affecting the aortic valve (natural or prosthetic). Any site in the body may be involved in metastatic abscess formation or inflammation; the female breast and the thyroid gland are affected particularly often. Nonspecific maculopapular rashes and other skin manifestations are uncommon and are rarely noticed by the patient even if they develop.

DIAGNOSIS

Because the clinical picture of brucellosis is not distinctive, the diagnosis must be based on a history of potential exposure, a presentation consistent with the disease, and supporting laboratory findings. Results of routine biochemical assays are usually within normal limits, although serum levels of hepatic enzymes and bilirubin may be elevated. Peripheral leukocyte counts are usually normal or low, with relative lymphocytosis. Mild anemia may be documented. Thrombocytopenia and disseminated intravascular coagulation with raised levels of fibrinogen degradation products can develop. The erythrocyte sedimentation rate and C-reactive protein levels are often normal but may be raised.

In body fluids such as cerebrospinal fluid (CSF) or joint fluid, lymphocytosis and low glucose levels are the norm. Elevated CSF levels of adenosine deaminase cannot be used to distinguish tubercular meningitis, as they may also be found in brucellosis. Biopsied samples of tissues such as lymph node or liver may show noncaseating granulomas (Fig. 194e-1) without acid/alcohol-fast bacilli. The radiologic features of bony disease develop late and are much more subtle than those of

tuberculosis or septic arthritis of other etiologies, with less bone and joint destruction. Isotope scanning is more sensitive than plain x-ray and continues to give positive results long after successful treatment.

Isolation of brucellae from blood, CSF, bone marrow, or joint fluid or from a tissue aspirate or biopsy sample is definitive, and attempts at isolation are usually successful in 50–70% of cases. Duplicate cultures should be incubated for up to 6 weeks (in air and 10% CO₂, respectively). Concentration and lysis of buffy coat cells before culture may increase the isolation rate. Cultures in modern nonradiometric or similar signaling systems (e.g., Bactec) usually become positive within 7–10 days but should be maintained for at least 3 weeks before the results are declared negative. All cultures should be handled under containment conditions appropriate for dangerous pathogens. *Brucella* species may be misidentified as *Agrobacterium*, *Ochrobactrum*, or *Psychrobacter (Moraxella) phenylpyruvicus* by the gallery identification strips commonly used in the diagnostic laboratory. In recent years, matrix-assisted laser desorption ionization time-of-flight spectrometry (MALDI-TOF MS) has emerged as a powerful tool in bacterial identification. The relative homogeneity of classical *Brucella* species makes identification beyond the genus level by routine approaches challenging, although further improvements may facilitate discrimination at the species level. The place of this technique in routine diagnostic practice will depend on such refinements. Meanwhile, the authors are aware of cases in which blood culture isolates have been identified incorrectly using MALDI-TOF MS.

The peripheral blood–based polymerase chain reaction has enormous potential to detect bacteremia, to predict relapse, and to exclude “chronic brucellosis.” This method is probably more sensitive and is certainly quicker than blood culture, and it does not carry the attendant biohazard risk posed by culture. Nucleic acid amplification techniques are now quite widely used, although no single standardized procedure has been adopted. Primers for the spacer region between the genes encoding the 16S and 23S ribosomal RNAs (*rrs-rrl*), various outer-membrane protein–encoding genes, the insertion sequence *IS711*, and the protein BCSP31 are sensitive and specific. Blood and other tissues are the most suitable samples for analysis.

Serologic examination often provides the only positive laboratory findings in brucellosis. In acute infection, IgM antibodies appear early and are followed by IgG and IgA. All these antibodies are active in agglutination tests, whether performed by tube, plate, or microagglutination methods. The majority of patients have detectable agglutinins at this stage. As the disease progresses, IgM levels decline, and the avidity and subclass distribution of IgG and IgA change. The result is reduced or undetectable agglutinin titers. However, the antibodies are detectable by alternative tests, including the complement fixation test, Coomb’s antiglobulin test, and enzyme-linked immunosorbent assay. There is no clear cutoff value for a diagnostic titer. Rather, serology results must be interpreted in the context of exposure history and clinical presentation. In endemic areas or in settings of potential occupational exposure, agglutinin titers of 1:320–1:640 or higher are considered diagnostic; in nonendemic areas, a titer of ≥1:160 is considered significant. Repetition of tests after 2–4 weeks may demonstrate a rising titer.

In most centers, the standard agglutination test is still the mainstay of serologic diagnosis, although some investigators rely on the rose bengal test, which has not been fully validated for human diagnostic use. Dipstick assays for anti-*Brucella* IgM are useful for the diagnosis of acute infection but are less sensitive for infection with symptoms of several months’ duration. In an endemic setting, more than 90% of patients with acute bacteremia have standard agglutination titers of at least 1:320. Other screening tests are used in some centers.

Antibody to the *Brucella* LPS O chain—the dominant antigen—is detected by all the conventional tests that employ smooth *B. abortus* cells as antigen. Since *B. abortus* cross-reacts with *B. melitensis* and *B. suis*, there is no advantage in replicating the tests with these antigens. Cross-reactions also occur with the O chains of some other gram-negative bacteria, including *Yersinia enterocolitica* O:9, *Escherichia coli* O157, *Francisella tularensis*, *Salmonella enterica* group N, *Stenotrophomonas maltophilia*, and *Vibrio cholerae*. Cross-reactions

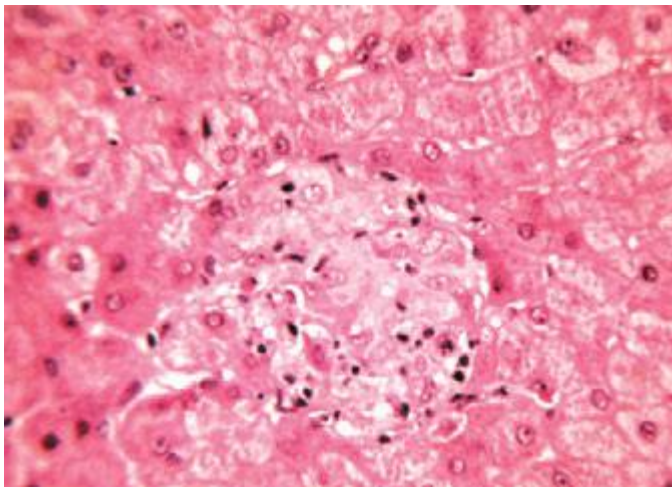


FIGURE 194e-1 Liver biopsy specimen from a patient with brucellosis shows a noncaseating granuloma. (From Mandell’s Atlas of Infectious Diseases, Vol II, in DL Stevens [ed]: Skin, Soft Tissue, Bone and Joint Infections, Fig. 5-9; with permission.)

do not occur with the cell-surface antigens of rough *Brucella* strains such as *B. canis* or *B. ovis*; serologic tests for these nomen species must employ an antigen prepared from either one. The live *B. abortus* vaccine strain RB51 does not elicit antibody responses in serologic tests that use smooth antigens, and this fact must be taken into account if serologic tests are employed in attempts to identify or follow the course of infections in persons accidentally exposed to the vaccine.

TREATMENT BRUCELLOSIS

The broad aims of antimicrobial therapy are to treat and relieve the symptoms of current infection and to prevent relapse. Focal disease presentations may require specific intervention in addition to more prolonged and tailored antibiotic therapy. In addition, tuberculosis must always be excluded, or—to prevent the emergence of resistance—therapy must be tailored to specifically exclude drugs active against tuberculosis (e.g., rifampin used alone) or to include a full antituberculous regimen.

Early experience with streptomycin monotherapy showed that relapse was common; thus dual therapy with tetracyclines became the norm. This is still the most effective combination, but alternatives may be used, with the options depending on local or national policy about the use of rifampin for the treatment of nonmycobacterial infection. For the several antimicrobial agents that are active in vivo, efficacy can usually be predicted by in vitro testing. However, numerous *Brucella* strains show in vitro sensitivity to a whole range of antimicrobials that are therapeutically ineffective, including assorted β -lactams. Moreover, the use of fluoroquinolones remains controversial despite the good in vitro activity and white-cell penetration of most agents of this class. Low intravacuolar pH is probably a factor in the poor performance of these drugs.

For adults with acute nonfocal brucellosis (duration, <1 month), a 6-week course of therapy incorporating at least two antimicrobial agents is required. Complex or focal disease necessitates ≥ 3 months of therapy. Adherence to the therapeutic regimen is very important, and poor adherence underlies almost all cases of apparent treatment failure; such failure is rarely due to the emergence of drug resistance, although increasing resistance to trimethoprim-sulfamethoxazole (TMP-SMX) has been reported at one center. There is good retrospective evidence that a 3-week course of two agents is as effective as a 6-week course for treatment and prevention of relapse in children, but this point has not yet been proven in prospective studies.

The gold standard for the treatment of brucellosis in adults is IM streptomycin (0.75–1 g daily for 14–21 days) together with doxycycline (100 mg twice daily for 6 weeks). In both clinical trials and observational studies, relapse follows such treatment in 5–10% of cases. The usual alternative regimen (and the current World Health Organization recommendation) is rifampin (600–900 mg/d) plus doxycycline (100 mg twice daily) for 6 weeks. The relapse/failure rate is $\sim 10\%$ in trial conditions but rises to $>20\%$ in many non-trial situations, possibly because doxycycline levels are reduced and clearance rates increased by concomitant rifampin administration. Patients who cannot tolerate or receive tetracyclines (children, pregnant women) can be given high-dose TMP-SMX instead (two or three standard-strength tablets twice daily for adults, depending on weight).

Increasing evidence supports the use of an aminoglycoside such as gentamicin (5–6 mg/kg per day for at least 2 weeks) instead of streptomycin, although this regimen is not approved by the U.S. Food and Drug Administration. Shorter courses have been associated with high failure rates in adults. A 5- to 7-day course of therapy with gentamicin and a 3-week course of TMP-SMX may be adequate for children with uncomplicated disease, but prospective trials are still needed to support this recommendation. Early experience with fluoroquinolone monotherapy was disappointing, although it was suggested that ofloxacin or ciprofloxacin, given together with rifampin for 6 weeks, might be an acceptable

alternative to the other 6-week regimens for adults. A substantial meta-analysis did not support the use of fluoroquinolones in first-line treatment regimens, and these drugs are not recommended by an expert consensus group (Ioannina) except in the context of well-designed clinical trials. However, a more recent meta-analysis is more supportive of the efficacy of these drugs, and an adequately powered prospective study will be needed to resolve their role in standard combination therapy. A triple-drug regimen—doxycycline and rifampin combined with an initial course of an aminoglycoside—was superior to double-drug regimens in a meta-analysis. The triple-drug regimen should be considered for all patients with complicated disease and for those for whom treatment adherence is likely to be a problem.

Significant neurologic disease due to *Brucella* species requires prolonged treatment (i.e., for 3–6 months), usually with ceftriaxone supplementation of a standard regimen. *Brucella* endocarditis is treated with at least three drugs (an aminoglycoside, a tetracycline, and rifampin), and many experts add ceftriaxone and/or a fluoroquinolone to reduce the need for valve replacement. Treatment is usually given for at least 6 months, and clinical endpoints for its discontinuation are often difficult to define. Surgery is still required for the majority of cases of infection of prosthetic heart valves and prosthetic joints.

There is no evidence base to guide prophylaxis after exposure to *Brucella* organisms (e.g., in the laboratory), inadvertent immunization with live vaccine intended for use in animals, or exposure to deliberately released brucellae. Most authorities have recommended the administration of rifampin plus doxycycline for 3 weeks after a low-risk exposure (e.g., an unspecified laboratory accident) and for 6 weeks after a major exposure to aerosol or injected material. However, such regimens are poorly tolerated, and doxycycline monotherapy of the same duration may be substituted. (Monotherapy is now the standard recommendation in the United Kingdom but not in the United States.) Rifampin should be omitted after exposure to vaccine strain RB51, which is resistant to rifampin but sensitive to doxycycline. After significant brucellosis exposure, expert consultation is advised for women who are (or may be) pregnant.

PROGNOSIS AND FOLLOW-UP

Relapse occurs in up to 30% of poorly compliant patients. Thus patients should ideally be followed clinically for up to 2 years to detect relapse, which responds to a prolonged course of the same therapy used originally. The general well-being and the body weight of the patient are more useful guides than serology to lack of relapse. IgG antibody levels detected by the standard agglutination test and its variants can remain in the diagnostic range for >2 years after successful treatment. Complement fixation titers usually fall to normal within 1 year of cure. Immunity is not solid; patients can be reinfected after repeated exposures. Fewer than 1% of patients die of brucellosis. When the outcome is fatal, death is usually a consequence of cardiac involvement; more rarely, it results from severe neurologic disease. Despite the low mortality rate, recovery from brucellosis is slow, and the illness can cause prolonged inactivity, with domestic and economic consequences.

The existence of a prolonged chronic brucellosis state after successful treatment remains controversial. Evaluation of patients in whom this state is considered (often those with work-related exposure to brucellae) includes careful exclusion of malingering, nonspecific chronic fatigue syndromes, and other causes of excessive sweating, such as alcohol abuse and obesity. In the future, the availability of more sensitive assays to detect *Brucella* antigen or DNA may help to identify patients with ongoing infection.

PREVENTION

Vaccines based on live attenuated *Brucella* strains, such as *B. abortus* strain 19BA or 104M, have been used in some countries to protect high-risk populations but have displayed only short-term efficacy and

high reactogenicity. Subunit vaccines have been developed but are of uncertain value and cannot be recommended at present. Research in this area has been stimulated by interest in biodefense ([Chap. 261e](#)) and may eventually yield new products. The mainstay of veterinary prevention is a national commitment to testing and slaughter of infected herds/flocks (with compensation for owners), control of animal movement, and active immunization of animals. These measures are usually sufficient to control human disease as well. In their

absence, pasteurization of all milk products before consumption is sufficient to prevent non-occupational animal-to-human transmission. All cases of brucellosis in animals and humans should be reported to the appropriate public health authorities.

ACKNOWLEDGMENT

The authors thank Dr. Adrian M. Whatmore for his review of the manuscript.



195 Tularemia

Richard F. Jacobs, Gordon E. Schutze

Tularemia is a zoonosis caused by *Francisella tularensis*. Humans of any age, sex, or race are universally susceptible to this systemic infection. Tularemia is primarily a disease of wild animals and persists in contaminated environments, ectoparasites, and animal carriers. Human infection is incidental and usually results from interaction with biting or blood-sucking insects, contact with wild or domestic animals, ingestion of contaminated water or food, or inhalation of infective aerosols. The illness is characterized by various clinical

syndromes, the most common of which consists of an ulcerative lesion at the site of inoculation, with regional lymphadenopathy and lymphadenitis. Systemic manifestations, including pneumonia, typhoidal tularemia, meningitis, and fever without localizing findings, pose a greater diagnostic challenge.

ETIOLOGY AND EPIDEMIOLOGY

F. tularensis is a class A bioterrorism agent (Chap. 261e). With rare exceptions, tularemia is the only disease produced by *F. tularensis*—a small (0.2 μm by 0.2–0.7 μm), gram-negative, pleomorphic, nonmotile, non-spore-forming bacillus. Bipolar staining results in a coccoid appearance. The organism is a thinly encapsulated, nonpiliated strict aerobe that invades host cells. In nature, *F. tularensis* is a hardy organism that persists for weeks or months in mud, water, and decaying animal carcasses. Dozens of biting and blood-sucking insects, especially ticks and tabanid flies, serve as vectors. Ticks and wild rabbits are the source for most human cases in endemic areas of the southeastern United States. In Utah, Nevada, and California, tabanid flies are the most common vectors. Animal reservoirs include wild rabbits, squirrels, birds, sheep, beavers, muskrats, and domestic dogs and cats. Person-to-person transmission is rare or nonexistent.

The four subspecies of *F. tularensis* are *tularensis*, *holarctica*, *novicida*, and *mediasiatica*. The first three of these subspecies are found in North America; in fact, subspecies *tularensis* has been isolated only in North America, where it accounts for >70% of cases of tularemia and produces more serious human disease than other subspecies (although, with treatment, the associated fatality rate is <2%). The progression of illness depends on the infecting strain's virulence, the inoculum size, the portal of entry, and the host's immune status.

Ticks pass *F. tularensis* to their offspring transovarially. The organism is found in tick feces but not in large quantities in tick salivary glands. In the United States, the disease is carried by *Dermacentor andersoni* (Rocky Mountain wood tick), *Dermacentor variabilis* (American dog tick), *Dermacentor occidentalis* (Pacific Coast dog tick), and *Amblyomma americanum* (Lone Star tick). *F. tularensis* is transmitted frequently during blood meals taken by embedded ticks after hours of attachment. It is the taking of a blood meal through a fecally contaminated field that transmits the organism. Transmission by ticks and tabanid flies takes place mainly in the spring and summer. However, continued transmission in the winter by trapped or hunted animals has been documented.



Tularemia is most common in the southeastern United States; Arkansas, Missouri, and Oklahoma account for more than half of all reported cases in this country. Small outbreaks in higher-risk populations (e.g., professional landscapers cutting up brush, mowing, and using a leaf blower) have been reported. Although the irregular distribution of cases of tularemia makes worldwide estimates difficult, increasing numbers of cases have been reported between latitudes 30° and 71°N (the Holarctic region) in the Northern Hemisphere. Cases of tularemia have been reported from Europe, Turkey, Canada, Mexico, and Asia. If the disease is caused by subspecies *tularensis*, the clinical manifestations are similar to those in the United States. However, in areas where tularemia is due largely to subspecies *holarctica*, oropharyngeal disease is common. Disease acquisition results from the consumption of water contaminated by live organisms shed by animals (e.g., muskrats, beavers). Subspecies *holarctica* is known to cause milder disease than other subspecies and responds well to fluoroquinolones, especially ciprofloxacin.

PATHOGENESIS AND PATHOLOGY

The most common portal of entry for human infection is through skin or mucous membranes, either directly—through the bite of ticks, other arthropods, or other animals—or via inapparent abrasions. Inhalation or ingestion of *F. tularensis* also can result in infection. *F. tularensis* is extremely infectious: Although >10⁸ organisms are usually required to produce infection via the oral route (oropharyngeal or gastrointestinal tularemia), as few as 10 organisms can result in infection when injected into the skin (ulceroglandular/glandular tularemia) or inhaled (pulmonary tularemia). After inoculation into the skin, the organism

multiplies locally; within 2–5 days (range, 1–10 days), it produces an erythematous, tender, or pruritic papule. The papule rapidly enlarges and forms an ulcer with a black base (chancriform lesion). The bacteria spread to regional lymph nodes, producing lymphadenopathy (buboes). All forms can lead to bacteremia with spread to distant organs, including the central nervous system.

Tularemia is characterized by mononuclear cell infiltration with pyogranulomatous pathology. The histopathologic findings can be quite similar to those in tuberculosis, although tularemia develops more rapidly. As a facultatively intracellular bacterium, *F. tularensis* can parasitize both phagocytic and nonphagocytic host cells and can survive intracellularly for prolonged periods. In the acute phase of infection, the primary organs affected (skin, lymph nodes, liver, and spleen) include areas of focal necrosis, which are initially surrounded by polymorphonuclear leukocytes (PMNs). Subsequently, granulomas form, with epithelioid cells, lymphocytes, and multinucleated giant cells surrounded by areas of necrosis. These areas may resemble caseation necrosis but later coalesce to form abscesses.

Conjunctival inoculation can result in infection of the eye, with regional lymph node enlargement (preauricular lymphadenopathy, Parinaud's complex). Aerosolization and inhalation or hematogenous spread of organisms can result in pneumonia. In the lung, an inflammatory reaction develops, including foci of alveolar necrosis and cell infiltration (initially polymorphonuclear and later mononuclear) with granulomas. Chest roentgenograms usually reveal bilateral patchy infiltrates rather than large areas of consolidation. Pleural effusions are common and may contain blood. Lymphadenopathy occurs in regions draining infected organs. Therefore, in pulmonary infection, mediastinal adenopathy may be evident, whereas patients with oropharyngeal tularemia develop cervical lymphadenopathy. In gastrointestinal or typhoidal tularemia, mesenteric lymphadenopathy may follow the ingestion of large numbers of organisms. (The term *typhoidal tularemia* may be used to describe severe bacteremic disease, irrespective of the mode of transmission or portal of entry.) Meningitis has been reported as a primary or secondary manifestation of bacteremia. Patients may also present with fever and no localizing signs.

IMMUNOLOGY

Although a complete and widely accepted understanding of the protective immune response to *F. tularensis* is lacking, significant advances in the study of natural and protective immunity have been made in recent years and may ultimately result in a vaccine candidate. Complete genomic sequencing and the availability of attenuated *F. tularensis* strains developed through genetic manipulation are facilitating research that will expand our knowledge in this area.

A number of investigators have studied various models and proposed various hypotheses regarding the induction of protective immunity to *F. tularensis*. Although further research is needed, synergy between humoral and cell-mediated immune (CMI) responses appears to be critical in inducing effective immune protection. Elucidation of the molecular mechanisms for the organism's evasion of the host response, pathogen-associated molecular patterns, and effective host immune protection has led to novel vaccination strategies tested in animal models. Antibodies to Fc receptors on antigen-presenting cells have been shown to be protective in animal models of pulmonary tularemia, resulting in both mucosal and CMI responses. This enhanced understanding of mucosal and serum antibodies in combination with a targeted CMI response holds great promise for future vaccine development.

CLINICAL MANIFESTATIONS

Tularemia often starts with a sudden onset of fever, chills, headache, and generalized myalgias and arthralgias (Table 195-1). This onset takes place when the organism penetrates the skin, is ingested, or is inhaled. An incubation period of 2–10 days is followed by the formation of an ulcer at the site of penetration, with local inflammation. The ulcer may persist for several months as organisms are transported via the lymphatics to the regional lymph nodes. These nodes enlarge and may become necrotic and suppurative. If the organism enters the bloodstream, widespread dissemination can result.

TABLE 195-1 CLINICAL PRESENTATION OF TULAREMIA

Sign or Symptom	Rate of Occurrence, %	
	Children	Adults
Lymphadenopathy	96	65
Fever ($\geq 38.3^{\circ}\text{C}$ or $\geq 101^{\circ}\text{F}$)	87	21
Ulcer/eschar/papule	45	51
Myalgias/arthralgias	39	2
Headache	9	5
Cough	9	5
Pharyngitis	43	—
Diarrhea	43	—

Source: Adapted from RF Jacobs, JP Narain: Pediatrics 76:818, 1985; with permission.

In the United States, most patients with tularemia (75–85%) acquire the infection by inoculation of the skin. In adults, the most common localized form is inguinal/femoral lymphadenopathy; in children, it is cervical lymphadenopathy. About 20% of patients develop a generalized maculopapular rash, which occasionally becomes pustular. Erythema nodosum occurs infrequently. The clinical manifestations of tularemia have been divided into various syndromes, which are listed in Table 195-2.

Ulceroglandular/Glandular Tularemia These two forms of tularemia account for ~75–85% of cases. The predominant form in children involves cervical or posterior auricular lymphadenopathy and is usually related to tick bites on the head and neck. In adults, the most common form is inguinal/femoral lymphadenopathy resulting from insect and tick exposures on the lower limbs. In cases related to wild game, the usual portal of entry for *F. tularensis* is either an injury sustained while skinning or cleaning an animal carcass or a bite (usually on the hand). Epitrochlear lymphadenopathy/lymphadenitis is common in patients with bite-related injuries.

In ulceroglandular tularemia, the ulcer is erythematous, indurated, and nonhealing, with a punched-out appearance that lasts 1–3 weeks. The papule may begin as an erythematous lesion that is tender or pruritic; it evolves over several days into an ulcer with sharply demarcated edges and a yellow exudate. The ulcer gradually develops a black base, and simultaneously the regional lymph nodes become tender and severely enlarged (Fig. 195-1). The affected lymph nodes may become fluctuant and drain spontaneously, but the condition usually resolves with effective treatment. Late suppuration of lymph nodes has been described in up to 25% of patients with ulceroglandular/glandular tularemia. Examination of material taken from these late fluctuant nodes after successful antimicrobial treatment reveals sterile necrotic tissue. In 5–10% of patients, the skin lesion may be inapparent, with lymphadenopathy plus systemic signs and symptoms the only physical findings (*glandular tularemia*). Conversely, a tick or deerfly bite on the trunk may result in an ulcer without evident lymphadenopathy.

Oculoglandular Tularemia In ~1% of patients, the portal of entry for *F. tularensis* is the conjunctiva, which the organism usually reaches through contact with contaminated fingers. The inflamed conjunctiva is painful, with numerous yellowish nodules and pinpoint ulcers.

TABLE 195-2 CLINICAL SYNDROMES OF TULAREMIA

Syndrome	Rate of Occurrence, %	
	Children	Adults
Ulceroglandular	45	51
Glandular	25	12
Pulmonary (pneumonia)	14	18
Oropharyngeal	4	—
Oculoglandular	2	—
Typhoidal	2	12
Unclassified	6	11

Source: Adapted from RF Jacobs, JP Narain: Pediatrics 76:818, 1985; with permission.



FIGURE 195-1 An 8-year-old boy with inguinal lymphadenitis and associated tick-bite site characteristic of ulceroglandular tularemia.

Purulent conjunctivitis with regional lymphadenopathy (preauricular, submandibular, or cervical) is evident. Because of debilitating pain, the patient may seek medical attention before regional lymphadenopathy develops. Painful preauricular lymphadenopathy is unique to tularemia and distinguishes it from tuberculosis, sporotrichosis, and syphilis. Corneal perforation may occur.

Oropharyngeal and Gastrointestinal Tularemia Rarely, tularemia follows ingestion of contaminated undercooked meat, oral inoculation of *F. tularensis* from the hands in association with the skinning and cleaning of animal carcasses, or consumption of contaminated food or water. Oral inoculation may result in acute, exudative, or membranous pharyngitis associated with cervical lymphadenopathy or in ulcerative intestinal lesions associated with mesenteric lymphadenopathy, diarrhea, abdominal pain, nausea, vomiting, and gastrointestinal bleeding. Infected tonsils become enlarged and develop a yellowish-white pseudomembrane, which can be confused with that of diphtheria. The clinical severity of gastrointestinal tularemia varies from mild, unexplained, persistent diarrhea with no other symptoms to a fulminant, fatal disease. In fatal cases, the extensive intestinal ulceration found at autopsy suggests an enormous inoculum.

Pulmonary Tularemia Pneumonia due to *F. tularensis* presents as variable parenchymal infiltrates that are unresponsive to treatment with β -lactam antibiotics. Tularemia must be considered in the differential diagnosis of atypical pneumonia in a patient with a history of travel to an endemic area. The disease can result from inhalation of an infectious aerosol or can spread to the lungs and pleura via bacteremia. Inhalation-related pneumonia has been described in laboratory workers after exposure to contaminated materials and, if untreated, can be associated with a relatively high mortality rate. Exposure to *F. tularensis* in aerosols from live domestic animals or dead wildlife (including birds) has been reported to cause pneumonia. Hematogenous dissemination to the lungs occurs in 10–15% of cases of ulceroglandular tularemia and in about half of cases of typhoidal tularemia. Previously, tularemia pneumonia was thought to be a disease of older patients, but as many as 10–15% of children with clinical manifestations of tularemia have parenchymal infiltrates detected by chest roentgenography. Patients with pneumonia usually have a nonproductive cough and may have dyspnea or pleuritic chest pain. Roentgenograms of the chest usually reveal bilateral patchy infiltrates (described as ovoid or lobar densities), lobar parenchymal infiltrates, and cavitory lesions. Pleural effusions may have a predominance of mononuclear leukocytes or PMNs and sometimes red blood cells. Empyema may develop. Blood cultures may be positive for *F. tularensis*.

Typhoidal Tularemia The typhoidal presentation is now considered rare in the United States. The source of infection in typhoidal tularemia is usually associated with pharyngeal and/or gastrointestinal inoculation or bacteremic disease. Fever usually develops without apparent skin lesions or lymphadenopathy. Some patients have cervical and mesenteric lymphadenopathy. In the absence of a history of possible contact with a vector, diagnosis can be extremely difficult. Blood cultures may be positive and patients may present with classic sepsis or septic shock in this acute systemic form of the infection. Typhoidal tularemia is usually associated with a huge inoculum or with a preexisting compromising condition. High continuous fevers, signs of sepsis, and severe headache are common. The patient may be delirious and may develop prostration and shock. If presumptive antibiotic therapy in culture-negative cases does not include an aminoglycoside, the estimated mortality rate is relatively high.

Other Manifestations *F. tularensis* infection has been associated with meningitis, pericarditis, hepatitis, peritonitis, endocarditis, osteomyelitis, and sepsis and septic shock with rhabdomyolysis and acute renal failure. In cases of tularemia meningitis, a mean white blood cell count of 1788/ μL , a predominantly mononuclear cell response (70–100%), a depressed glucose level, an elevated protein concentration, and a negative Gram's stain are typically found on examination of cerebrospinal fluid.

DIFFERENTIAL DIAGNOSIS

When patients in endemic areas present with fever, chronic ulcerative skin lesions, and large tender lymph nodes (Fig. 195-1), a diagnosis of tularemia should be made presumptively, and confirmatory diagnostic testing and appropriate therapy should be undertaken. When the possibility of tularemia is considered in a nonendemic area, an attempt should be made to identify contact with a potential animal vector. The level of suspicion should be especially high in hunters, trappers, game wardens, professional landscapers, veterinarians, laboratory workers, and individuals exposed to an insect or another animal vector. However, up to 40% of patients with tularemia have no known history of epidemiologic contact with an animal vector.

The characteristic presentation of ulceroglandular tularemia does not pose a diagnostic problem, but a less classic progression of regional lymphadenopathy or glandular tularemia must be differentiated from other diseases (Table 195-3). The skin lesion of tularemia may resemble those seen in various other diseases but is generally accompanied by more impressive regional lymphadenopathy. In children, the differentiation of tularemia from cat-scratch disease is made more difficult by the chronic papulovesicular lesion associated with *Bartonella henselae* infection (Chap. 197). Oropharyngeal tularemia can resemble and must be differentiated from pharyngitis due to other bacteria or viruses. Pulmonary tularemia may resemble any atypical pneumonia. Typhoidal tularemia and tularemia meningitis may resemble a variety of other infections.

LABORATORY DIAGNOSIS

The diagnosis of tularemia is most frequently confirmed by agglutination testing. Microagglutination and tube agglutination are the techniques most commonly used to detect antibody to *F. tularensis*. In the standard tube agglutination test, a single titer of $\geq 1:160$ is interpreted as a presumptive positive result. A fourfold increase in titer between paired serum samples collected 2–3 weeks apart is considered diagnostic. False-negative serologic responses are obtained early in infection; up to 30% of patients infected for 3 weeks have sera that test negative. Late in infection, titers into the thousands are common, and titers of 1:20–1:80 may persist for years. Enzyme-linked immunosorbent assays have proved useful for the detection of both antibodies and antigens.

Culture and isolation of *F. tularensis* are difficult. In one study, the organism was isolated in only 10% of more than 1000 human cases, 84% of which were confirmed by serology. The medium of choice is cysteine-glucose-blood agar. *F. tularensis* can be isolated directly from infected ulcer scrapings, lymph node biopsy specimens, gastric washings, sputum, and blood cultures. Colonies are blue-gray, round, smooth, and slightly mucoid. On media containing blood, a small zone of a hemolysis usually surrounds the colony. Slide agglutination tests or direct fluorescent antibody tests with commercially available antisera can be applied directly to culture suspensions for identification. Most clinical laboratories will not attempt to culture *F. tularensis* because of the infectivity of the organism from the culture media and the consequent risk of a laboratory-acquired infection. Although tularemia is not spread from person to person, the organism can be inhaled from culture plates and infect unsuspecting laboratory workers. In most clinical laboratories, biosafety level 2 practices are recommended to handle clinical specimens thought to contain *F. tularensis*; however, biosafety level 3 conditions are required for procedures that produce aerosols or droplets during manipulation of cultures containing or possibly containing this organism.

A variety of polymerase chain reaction (PCR) methods have been used to detect *F. tularensis* DNA in many clinical specimens but mostly in ulceroglandular disease. The majority of these methods target the genes encoding outer-membrane proteins (e.g., *fopA* or *tul4*). A 16S rDNA sequence identification PCR may be helpful when the patient's clinical information does not lead the clinician to suspect a diagnosis of tularemia.

TREATMENT TULAREMIA

Only aminoglycosides, tetracyclines, chloramphenicol, and rifampin are currently approved by the U.S. Food and Drug Administration for the treatment of tularemia. Gentamicin is considered the drug of choice for both adults and children. The dosage for adults and children is 5 mg/kg daily in two divided doses. Gentamicin therapy is typically continued for 7–10 days; however, in mild to moderate

TABLE 195-3 TULAREMIA: DIFFERENTIAL DIAGNOSIS, BY CLINICAL DISEASE CATEGORY

Glandular	Oropharyngeal	Typhoidal	Pulmonary
Pyogenic bacterial infection ^a	Group A streptococcal pharyngitis	Typhoid fever	<i>Mycoplasma pneumoniae</i> pneumonia
Nontuberculous mycobacterial infection	<i>Arcanobacterium haemolyticum</i> pharyngitis	Other <i>Salmonella</i> bacteremias	<i>Chlamydia pneumoniae</i> pneumonia
Sporotrichosis	Diphtheria	Rocky Mountain spotted fever	Psittacosis
Tuberculosis	Infectious mononucleosis	Human monocytotropic ehrlichiosis	<i>Legionella pneumophila</i> pneumonia
Syphilis	Various viral infections ^b	Human granulocytotropic anaplasmosis	Q fever
Anthrax		Infectious mononucleosis	Histoplasmosis
Rat-bite fever		Brucellosis	Blastomycosis
Scrub typhus		Toxoplasmosis	Coccidioidomycosis
Plague		Tuberculosis	Various viral infections ^d
Lymphogranuloma venereum		Sarcoidosis	
Cat-scratch disease		Malignancy ^c	

^a*Staphylococcus aureus*, *Streptococcus pyogenes*. ^bAdenovirus, enteroviruses, parainfluenza virus, influenza viruses A and B, respiratory syncytial virus. ^cHematologic and reticuloendothelial malignancies. ^dInfluenza viruses A and B, parainfluenza virus, respiratory syncytial virus, adenovirus, enteroviruses, hantavirus.

1070 cases of tularemia in which the patient becomes afebrile within the first 48–72 h of gentamicin treatment, a 5- to 7-day course has been successful.

If available, streptomycin given intramuscularly is also effective. The dosage for adults is 2 g/d in two divided doses. For children, the dosage is 30 mg/kg daily in two divided doses (maximal daily dose, 2 g). After a clinical response is demonstrated at 3–5 days, the dosage for children can be reduced to 10–15 mg/kg daily in two divided doses. The total duration of streptomycin therapy in both adults and children is usually 10 days. Unlike streptomycin and gentamicin, tobramycin is ineffective in the treatment of tularemia and should not be used.

Because doxycycline is bacteriostatic against *F. tularensis*, there is a risk of relapse if the patient is not treated for a long enough period. Therefore, if doxycycline is used, it should be given for at least 14 days. The lack of availability of chloramphenicol limits the utility of this agent as a viable treatment option. Fluoroquinolones—specifically, ciprofloxacin and levofloxacin—have been used with good outcomes to treat infections caused by subspecies *holarctica*, which is most often found in Europe. The lack of data on the efficacy of these agents against subspecies *tularensis* limits their use in North America at this time.

F. tularensis cannot be subjected to standardized antimicrobial susceptibility testing because the organism will not grow on the media used. A wide variety of antibiotics, including all β -lactam antibiotics and the cephalosporins, are ineffective for the treatment of tularemia. Several studies indicated that third-generation cephalosporins were active against *F. tularensis* in vitro, but clinical case reports suggested nearly universal failure of ceftriaxone in pediatric patients with tularemia. Although in vitro data indicate that imipenem may be active, therapy with imipenem, sulfanilamides, and macrolides is not presently recommended because of the lack of relevant clinical data.

Virtually all strains of *F. tularensis* are susceptible to streptomycin and gentamicin. Hearing screening should be considered before initiation of streptomycin or gentamicin therapy. In successfully treated patients, defervescence usually occurs within 2 days, but skin lesions and lymph nodes may take 1–2 weeks to heal. When therapy is not initiated within the first several days of illness, defervescence may be delayed. Relapses are uncommon with streptomycin or gentamicin therapy. Late lymph-node suppuration, however, occurs in ~40% of children, regardless of the treatment received. These nodes have typically been found to contain sterile necrotic tissue without evidence of active infection. Patients with fluctuant nodes should receive several days of antibiotic therapy before drainage to minimize the risk to hospital personnel.

PROGNOSIS

If tularemia goes untreated, symptoms usually last 1–4 weeks but may continue for months. The mortality rate from severe untreated infection (including all cases of untreated pulmonary and typhoidal tularemia) can be as high as 30%. However, the overall mortality rate for untreated tularemia is <8%. With appropriate treatment, the mortality rate is <1%. Poor outcomes are often associated with long delays in diagnosis and treatment. Lifelong immunity usually follows tularemia.

PREVENTION

The prevention of tularemia is based on avoidance of exposure to biting and blood-sucking insects, especially ticks and deerflies. A wide range of approaches to vaccine development are being evaluated, but no vaccine against tularemia is yet licensed. Prophylaxis of tularemia has not proved effective in patients with embedded ticks or insect bites. However, in patients who are known to have been exposed to large quantities of organisms (e.g., in the laboratory) and who have incubating infection with *F. tularensis*, early treatment can prevent the development of significant clinical disease.

196 Plague and Other *Yersinia* Infections

Michael B. Prentice

PLAGUE

Plague is a systemic zoonosis caused by *Yersinia pestis*. It predominantly affects small rodents in rural areas of Africa, Asia, and the Americas and is usually transmitted to humans by an arthropod vector (the flea). Less often, infection follows contact with animal tissues or respiratory droplets. Plague is an acute febrile illness that is treatable with antimicrobial agents, but mortality rates among untreated patients are high. Ancient DNA studies have confirmed that the fourteenth-century “Black Death” in Europe was *Y. pestis* infection. Patients can present with the bubonic, septicemic, or pneumonic form of the disease. Although there is concern among the general public about epidemic spread of plague by the respiratory route, this is not the usual route of plague transmission, and established infection-control measures for respiratory plague exist. However, the fatalities associated with plague and the capacity for infection via the respiratory tract mean that *Y. pestis* fits the profile of a potential agent of bioterrorism. Consequently, measures have been taken to restrict access to the organism, including legislation affecting diagnostic and research procedures in some countries (e.g., the United States).

ETIOLOGY

The genus *Yersinia* comprises gram-negative bacteria of the family Enterobacteriaceae (gamma proteobacteria). Overwhelming taxonomic evidence showing *Y. pestis* strains as a clonal group within *Yersinia pseudotuberculosis* suggests recent evolution from the latter organism—an enteric pathogen of mammals that is spread by the fecal-oral route and thus has a phenotype distinctly different from that of *Y. pestis*. When grown in vivo or at 37°C, *Y. pestis* forms an amorphous capsule made from a plasmid-specified fimbrial protein, Caf or fraction 1 (F1) antigen, which is an immunodiagnostic marker of infection.

EPIDEMIOLOGY

Human plague generally follows an outbreak in a host rodent population (epizootic). Mass deaths among the rodent primary hosts lead to a search by fleas for new hosts, with consequent incidental infection of other mammals. The precipitating cause for an epizootic may ultimately be related to climate or other environmental factors. The reservoir for *Y. pestis* causing enzootic plague in natural endemic foci between epizootics (i.e., when the organism may be difficult to detect in rodents or fleas) is a topic of ongoing research and may not be the same in all regions. The enzootic/epizootic pattern may be the result of complex dynamic interactions of host rodents that have different plague susceptibilities and different flea vectors; alternatively, an environmental reservoir may be important.



In general, the enzootic areas for plague are lightly populated regions of Africa, Asia, and the Americas (Fig. 196-1).

Between 2004 and 2009, 12,503 cases of plague, with a global case-fatality rate of 6.7%, were recorded by the World Health Organization (WHO); these figures were obtained by combining cases notified under the International Health Regulations with data from national surveillance programs and publications. More than 97% of these cases were in Africa; the majority of cases were reported from the Democratic Republic of the Congo and the island of Madagascar. The period covered spans a change in the International Health Regulations from a requirement for nations to notify the WHO of all cases of plague to a requirement to report pneumonic plague or any suspected case of plague in an area not known to be endemic for plague. In the past decade, outbreaks of pneumonic plague have been recorded in the Democratic Republic of the Congo, Uganda, Algeria, Madagascar, China, and Peru.

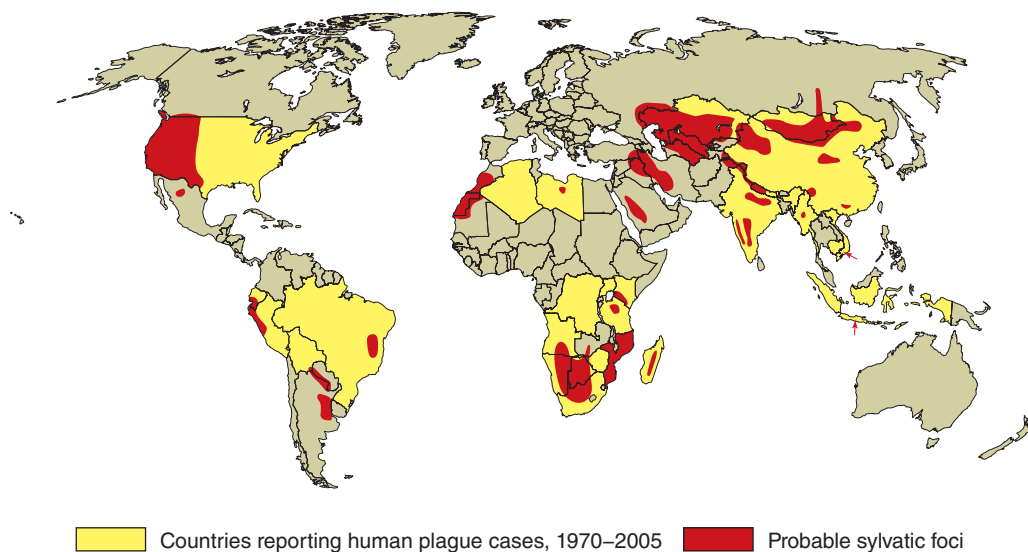


FIGURE 196-1 Approximate global distribution of *Yersinia pestis*. (Compiled from WHO, CDC, and country sources. Reprinted with permission from DT Dennis, GL Campbell: *Plague and other Yersinia infections*, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al [eds]. New York, McGraw-Hill, Chap. 152, 2008.)

Plague was introduced into North America via the port of San Francisco in 1900 as part of the Third Pandemic, which spread around the world from Hong Kong. The disease is presently enzootic on the western side of the continent from southwestern Canada to Mexico. Most human cases in the United States occur in two regions: “Four Corners” (the junction point of New Mexico, Arizona, Colorado, and Utah), especially northern New Mexico, northern Arizona, and southern Colorado; and further west in California, southern Oregon, and western Nevada (<http://www.cdc.gov/plague/maps/index.html>). From 1990 to 2011, 151 cases of plague were reported in the United States, a mean of seven cases per year. Most cases occurred from May to October—the time of year when people are outdoors and rodents and their fleas are most plentiful. The infection is most often acquired by fleabite in peridomestic environments; it can also be acquired through the handling of living or dead small mammals (e.g., rabbits, hares, and prairie dogs) or wild carnivores (e.g., wildcats, coyotes, or mountain lions). Dogs and cats may bring plague-infected fleas into the home, and infected cats may transmit plague directly to humans by the respiratory route. The last recorded case of person-to-person transmission in the United States occurred in 1925.

Plague most often develops in areas with poor sanitary conditions and infestations of rats—in particular, the widely distributed roof rat *Rattus rattus* and the brown rat *Rattus norvegicus* (which serves as a laboratory model of plague). Rat control in warehouses and shipping facilities has been recognized as important in preventing the spread of plague since the early twentieth century and features in the current WHO International Health Regulations. Urban rodents acquire infection from wild rodents, and the proximity of the former to humans increases the risk of transmission. The oriental rat flea *Xenopsylla cheopis* is the most efficient vector for transmission of plague among rats and onward to humans in Asia, Africa, and South America.

Worldwide, bubonic plague is the predominant form reported (80–95% of suspected cases), with mortality rates of 10–20%. The mortality rate is higher (22%) in the small proportion of patients (10–20%) with primary septicemic plague (i.e., systemic *Y. pestis* sepsis with no bubo; see “Clinical Manifestations,” below) and is highest with primary pulmonary plague; in this, the least common of the main plague presentations, the mortality rate approaches 100% without antimicrobial treatment and is >50% even with such treatment. Rare outbreaks of pharyngeal plague following consumption of raw or undercooked camel or goat meat have been reported.

A total of 81 (76%) of the 107 plague cases reported in the United States from 1990 to 2005 were primary bubonic disease, 19 (18%)

were primary septicemic disease, and 5 (5%) were primary pneumonic disease; 2 cases (2%) were not classified. Eleven cases (10%) were fatal.

PATHOGENESIS



As mentioned earlier, genetic evidence suggests that *Y. pestis* is a clone derived from the enteric pathogen *Y. pseudotuberculosis* in the recent evolutionary past (9000–40,000 years ago). The change from infection by the fecal-oral route to a two-stage life cycle, with alternate parasitization of arthropod and mammalian hosts, followed the acquisition of two plasmids (pFra and pPst) and the inactivation of remarkably few *Y. pseudotuberculosis* genes in conjunction with preexisting properties of the *Y. pseudotuberculosis* ancestor (e.g., the presence of a third plasmid, pYV, and the capacity to cause septicemia). In the arthropod-parasitizing portion of its life cycle, *Y. pestis* multiplies and forms biofilm-embedded aggregates in the flea midgut after ingestion of a blood meal containing bacteria. In some fleas, biofilm-embedded bacteria eventually fill the proventriculus (a valve connecting the esophagus to the midgut) and block normal blood feeding. Both “blocked” fleas and those containing masses of biofilm-embedded *Y. pestis* without complete blockage inoculate *Y. pestis* into each bite site. The ability of *Y. pestis* to colonize and multiply in the flea requires phospholipase D encoded by the *ymt* gene on the pFra plasmid, and biofilm synthesis requires the chromosomal *hms* locus shared with *Y. pseudotuberculosis*. However, three *Y. pseudotuberculosis* genes inhibiting biofilm formation or promoting its degradation are inactivated in *Y. pestis*, together with urease, which causes acute flea gastrointestinal toxicity. Blockage takes days or weeks to come about after initial infection of the flea and is followed by the flea’s death. In addition, many flea vectors (including *X. cheopis*) are able to transmit plague in an early-phase unblocked state for up to 1 week after feeding, but 10 fleas in this state are required to infect a mammalian host (mass transmission).

Y. pestis disseminates from the site of inoculation in the mammalian host in a process initially dependent on plasminogen activator Pla, which is encoded by the small pPst plasmid. This surface protease activates mammalian plasminogen, degrades complement, and adheres to the extracellular matrix component laminin. Pla is essential for the high-level virulence of *Y. pestis* in mice by subcutaneous or intradermal injection (laboratory proxies for fleabites) and for the development of primary pneumonic plague. When actual fleabite inoculation is used in mouse models, the fimbrial capsule-forming protein (Ca1 or fraction 1; F1 antigen) encoded on pFra increases the efficiency of transmission, and plasminogen activator is required for the formation

1072 of buboes. Because the antiphagocytic systems in *Y. pestis* are not fully operational at the time of inoculation into the mammalian host, the organism is taken up by macrophages from the inoculation site and transported to regional lymph nodes. After intracellular replication, *Y. pestis* switches to extracellular replication with full expression of its antiphagocytic systems: the type III secretion machines and their effectors encoded by pYV as well as the F1 capsule. Overproduction of the type III secretion substrate and translocation protein LcrV exerts an anti-inflammatory effect, reducing host immune responses. Likewise, *Y. pestis* lipopolysaccharide is modified to minimize stimulation of host Toll-like receptor 4, thereby reducing protective host inflammatory responses during peripheral infection and prolonging host survival with high-grade bacteremia—an effect that probably enhances the pathogen's subsequent transmission by fleabite.

Replication of *Y. pestis* in a regional lymph node results in the local swelling of the lymph node and periglandular region known as a *bubo*. On histology, the node is found to be hemorrhagic or necrotic, with thrombosed blood vessels, and the lymphoid cells and normal architecture are replaced by large numbers of bacteria and fibrin. Periglandular tissues are inflamed and also contain large numbers of bacteria in a serosanguineous, gelatinous exudate.

Continued spread through the lymphatic vessels to contiguous lymph nodes produces second-order primary buboes. Infection is initially contained in the infected regional lymph nodes, although transient bacteremia can be detected. As the infection progresses, spread via efferent lymphatics to the thoracic duct produces high-grade bacteremia. Hematogenous spread to the spleen, liver, and secondary buboes follows, with subsequent uncontrolled septicemia, endotoxic shock, and disseminated intravascular coagulation leading to death. In some patients, this septicemic phase occurs without obvious prior bubo development or lung disease (*septicemic plague*). Hematogenous spread to the lungs results in *secondary plague pneumonia*, with bacteria initially more prominent in the interstitium than in the air spaces (the reverse being the case in *primary plague pneumonia*). Hematogenous spread to other organs, including the meninges, can occur.

CLINICAL MANIFESTATIONS

Bubonic Plague After an incubation period of 2–6 days, the onset of bubonic plague is sudden and is characterized by fever (>38°C), malaise, myalgia, dizziness, and increasing pain due to progressive lymphadenitis in the regional lymph nodes near the fleabite or other inoculation site. Lymphadenitis manifests as a tense, tender swelling (*bubo*) that, when palpated, has a boggy consistency with an underlying hard core. Generally, there is one painful and erythematous bubo with surrounding periganglionic edema. The bubo is most commonly inguinal but can also be crural, axillary (**Fig. 196-2**), cervical, or submaxillary, depending on the site of the bite. Abdominal pain from intraabdominal node involvement can occur without other visible signs. Children are most likely to present with cervical or axillary buboes.

The differential diagnosis includes acute focal lymphadenopathy of other etiologies, such as streptococcal or staphylococcal infection, tularemia, cat-scratch disease, tick typhus, infectious mononucleosis, or lymphatic filariasis. These infections do not progress as rapidly, are not as painful, and are associated with visible cellulitis or ascending lymphangitis—both of which are absent in plague.

Without treatment, *Y. pestis* dissemination occurs and causes serious illness, including pneumonia (*secondary pneumonic plague*) and meningitis. Secondary pneumonic plague can be the source of person-to-person transmission of respiratory infection by productive cough (droplet infection), with the consequent development of primary plague pneumonia. Appropriate treatment of bubonic plague results in fever resolution within 2–5 days, but buboes may remain enlarged for >1 week after initial treatment and can become fluctuant.

Primary Septicemic Plague A minority (10–25%) of infections with *Y. pestis* present as gram-negative septicemia (hypotension, shock) without preceding lymphadenopathy. Septicemic plague occurs in all age groups, but persons older than age 40 years are at elevated risk. Some chronic conditions may predispose to septicemic plague: in



FIGURE 196-2 Plague patient in the southwestern United States with a left axillary bubo and an unusual plague ulcer and eschar at the site of the infective flea bite. (Reprinted with permission from DT Dennis, GL Campbell: *Plague and other Yersinia infections*, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al [eds]. New York, McGraw-Hill, Chap. 152, 2008.)

2009 in the United States, a fatal laboratory-acquired infection with an attenuated *Y. pestis* strain manifested as septicemic plague in a 60-year-old researcher with diabetes mellitus and undiagnosed hemochromatosis. These conditions also carry an increased risk of septicemia with other pathogenic *Yersinia* species. The term *septicemic plague* can be confusing since most patients with buboes have detectable bacteremia at some stage, with or without systemic signs of sepsis. In laboratory experiments, however, septicemic disease without histologic changes in lymph nodes is seen in a minority of mice infected via fleabites.

Pneumonic Plague Primary pneumonic plague results from inhalation of infectious bacteria in droplets expelled from another person or an animal with primary or secondary plague pneumonia. This syndrome has a short incubation period, averaging from a few hours to 2–3 days (range, 1–7 days), and is characterized by a sudden onset of fever, headache, myalgia, weakness, nausea, vomiting, and dizziness. Respiratory signs—cough, dyspnea, chest pain, and sputum production with hemoptysis—typically arise after 24 h. Progression of initial segmental pneumonitis to lobar pneumonia and then to bilateral lung involvement may occur (**Fig. 196-3**). The possible release of aerosolized *Y. pestis* bacteria in a bioterrorist attack, manifesting as an outbreak of primary pneumonic plague in nonendemic regions or in an urban setting where plague is rarely seen, has been a source of public health concern. Secondary pneumonic plague is a consequence of bacteremia occurring in ~10–15% of patients with bubonic plague. Bilateral alveolar infiltrates are seen on chest x-ray, and diffuse interstitial pneumonitis with scanty sputum production is typical.

Meningitis *Meningeal plague* is uncommon, occurring in ≤6% of plague cases reported in the United States. Presentation with headache and fever typically occurs >1 week after the onset of bubonic or septicemic plague and may be associated with suboptimal antimicrobial therapy (delayed therapy, penicillin administration, or low-dose tetracycline treatment) and cervical or axillary buboes.

Pharyngitis Symptomatic *plague pharyngitis* can follow the consumption of contaminated meat from an animal dying of plague or contact with persons or animals with pneumonic plague. This condition can resemble tonsillitis, with peritonsillar abscess and cervical

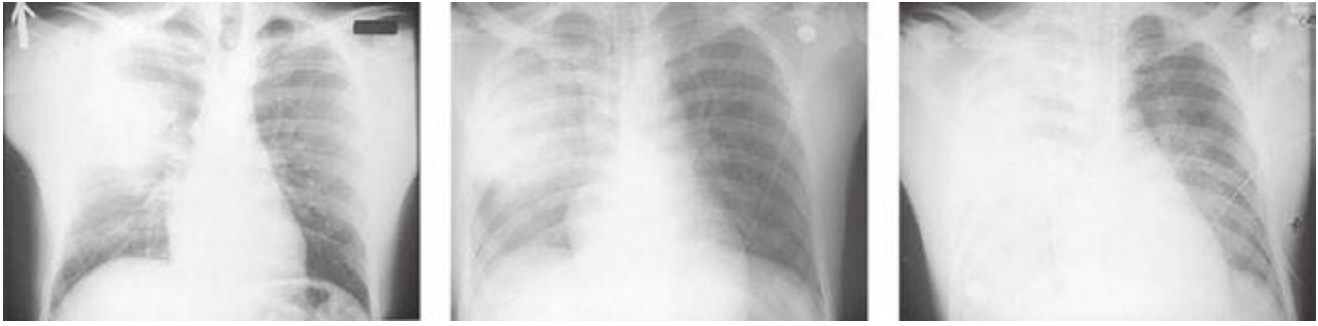


FIGURE 196-3 Sequential chest radiographs of a patient with fatal primary plague pneumonia. **Left:** Upright posteroanterior film taken at admission to the hospital emergency department on the third day of illness, showing segmental consolidation of the right upper lobe. **Center:** Portable anteroposterior film taken 8 h after admission, showing extension of pneumonia to the right middle and right lower lobes. **Right:** Portable anteroposterior film taken 13 h after admission (when the patient had clinical adult respiratory distress syndrome), showing diffuse infiltration throughout the right lung and patchy infiltration of the left lower lung. A cavity later developed at the site of the initial right-upper-lobe consolidation. (Reprinted with permission from DT Dennis, GL Campbell: Plague and other Yersinia infections, in Harrison's Principles of Internal Medicine, 17th ed., AS Fauci et al [eds]. New York, McGraw-Hill, Chap. 152, 2008.)

lymphadenopathy. Asymptomatic pharyngeal carriage of *Y. pestis* can also occur in close contacts of patients with pneumonic plague.

LABORATORY DIAGNOSIS



Because of the scarcity of laboratory facilities in regions where human *Y. pestis* infection is most common, and because of the potential significance of *Y. pestis* isolation in a nonendemic area or an area from which human plague has been absent for many years, the WHO recommends an initial presumptive diagnosis followed by reference laboratory confirmation (Table 196-1). In the United States, comprehensive national diagnostic facilities for plague have been in place since a federal Laboratory Response Network (LRN; www.bt.cdc.gov/lrn/) was set up in 1999 to detect possible use of biological terrorism agents, including *Y. pestis*. Routine diagnostic clinical microbiology laboratories that are included in this network as sentinel-level laboratories use joint protocols from the Centers for Disease Control and Prevention (CDC) and the American Society for

Microbiology to identify suspected *Y. pestis* isolates and to refer these specimens to LRN reference laboratories for confirmatory tests (<http://www.asm.org/index.php/issues/sentinel-laboratory-guidelines>). *Y. pestis* is designated a “Tier 1 select agent” under the Public Health Security and Bioterrorism Preparedness and Response Act of 2002 and subsequent executive orders; the provisions of this act, the Patriot Act of 2001, and related executive orders apply to all U.S. laboratories and individuals working with *Y. pestis*. Details of the applicable regulations are available from the CDC.

Yersinia species are gram-negative coccobacilli (short rods with rounded ends) 1–3 μm in length and 0.5–0.8 μm in diameter. *Y. pestis* in particular appears bipolar (with a “closed safety pin” appearance) and pleomorphic when stained with a polychromatic stain (Wayson or Wright-Giemsa; Fig. 196-4). Its lack of motility distinguishes *Y. pestis* from other *Yersinia* species, which are motile at 25°C and nonmotile at 37°C. Transport medium (e.g., Cary-Blair medium) preserves the viability of *Y. pestis* if transport is delayed.

TABLE 196-1 WORLD HEALTH ORGANIZATION CASE DEFINITIONS OF PLAGUE

Suspected case	Compatible clinical presentation and Consistent epidemiologic features, such as exposure to infected animals or humans and/or evidence of fleabites and/or residence in or travel to a known endemic focus within the previous 10 days
Presumptive case	Meeting the definition of a suspected case plus Putative new or reemerging focus: ≥ 2 of the following tests positive <ul style="list-style-type: none"> • Microscopy: gram-negative coccobacilli in material from bubo, blood, or sputum; bipolar appearance on Wayson or Wright-Giemsa staining • F1 antigen detected in bubo aspirate, blood, or sputum • A single anti-F1 serology without evidence of previous <i>Y. pestis</i> infection or immunization • Polymerase chain reaction (PCR) detection of <i>Y. pestis</i> in bubo aspirate, blood, or sputum Known endemic focus: ≥ 1 of the following tests positive <ul style="list-style-type: none"> • Microscopic evidence of gram-negative or bipolar (Wayson, Wright-Giemsa) coccobacilli from bubo, blood, or sputum sample • A single anti-F1 serology without evidence of previous plague infection or immunization • F1 antigen detected in bubo aspirate, blood, or sputum
Confirmed case	Meeting the definition of a suspected case plus <ul style="list-style-type: none"> • Identification of an isolate from a clinical sample as <i>Y. pestis</i> (colonial morphology and 2 of the 4 following tests positive: phage lysis of cultures at 20–25°C and 37°C; F1 antigen detection; PCR; <i>Y. pestis</i> biochemical profile) or <ul style="list-style-type: none"> • A fourfold rise in anti-F1 antibody titer in paired serum samples or <ul style="list-style-type: none"> • In endemic areas when no other confirmatory test can be performed, a positive rapid diagnostic test with immunochromatography to detect F1 antigen

Source: Interregional Meeting on Prevention and Control of Plague, Antananarivo, Madagascar, 7–11 April 2006 (www.who.int/entity/csr/resources/publications/WHO_HSE_EPR_2008_3w.pdf).

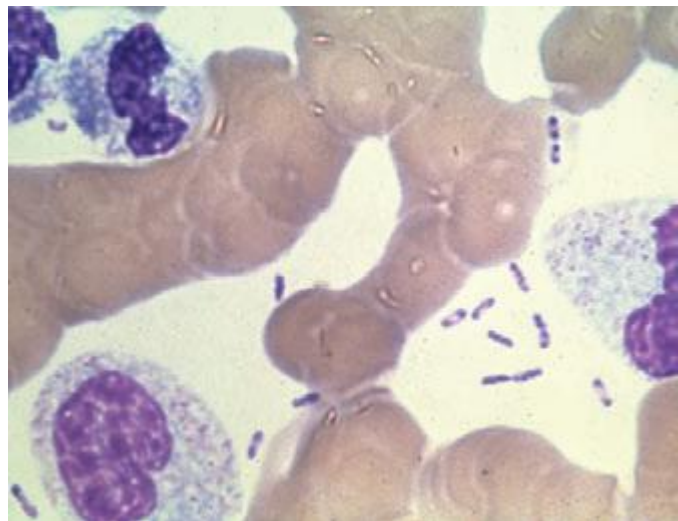


FIGURE 196-4 Peripheral blood smear from a patient with fatal plague septicemia and shock, showing characteristic bipolar-staining *Yersinia pestis* bacilli (Wright's stain, oil immersion). (Reprinted with permission from DT Dennis, GL Campbell: Plague and other *Yersinia* infections, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al [eds]. New York, McGraw-Hill, Chap. 152, 2008.)

The appropriate specimens for diagnosis of bubonic, pneumonic, and septicemic plague are bubo aspirate, bronchoalveolar lavage fluid or sputum, and blood, respectively. Culture of postmortem organ biopsy samples can also be diagnostic. A bubo aspirate is obtained by injection of 1 mL of sterile normal saline into a bubo under local anesthetic and aspiration of a small amount of (usually blood-stained) fluid. Gram's staining of these specimens may reveal gram-negative rods, which are shown by Wayson or Wright-Giemsa staining to be bipolar. These bacteria may even be visible in direct blood smears in septicemic plague (Fig. 196-4); this finding indicates very high numbers of circulating bacteria and a poor prognosis.

Y. pestis grows on nutrient agar and other standard laboratory media but forms smaller colonies than do other Enterobacteriaceae. Specimens should be inoculated onto nutrient-rich media such as sheep blood agar (SBA), into nutrient-rich broth such as brain-heart infusion broth, and onto selective agar such as MacConkey or eosin methylene blue (EMB) agar. *Yersinia*-specific CIN (cefsulodin, tricosan [Irgasan], novobiocin) agar can be useful for culture of contaminated specimens, such as sputum. Blood should be cultured in a standard blood culture system. The optimal growth temperature is 37°C (25–29°C), with pinpoint colonies only on SBA at 24 h. Slower growth occurs at 37°C. *Y. pestis* is oxidase-negative, catalase-positive, urea-negative, indole-negative, and lactose-negative. Automated biochemical identification systems can misidentify *Y. pestis* as *Y. pseudotuberculosis* or other bacterial species.

Reference laboratory tests for definitive identification of isolates include direct immunofluorescence for F1 antigen; specific polymerase chain reaction (PCR) for targets such as F1 antigen, the pesticin gene, and the plasminogen activator gene; and specific bacteriophage lysis. PCR can also be applied to diagnostic specimens, as can direct immunofluorescence for F1 antigen (produced in large amounts by *Y. pestis*) by slide microscopy. An immunochromatographic test strip for F1 antigen detection by monoclonal antibodies in clinical specimens has been devised in Madagascar. This method is effective for both laboratory and near-patient use and is now widely used in endemic countries. A similar test strip for Pla antigen has recently been developed and could be used to detect wild-type or engineered F1-negative virulent strains. Many other rapid diagnostic kits for possible bioterrorism pathogens, including *Y. pestis*, have been described in recent years, but none is widely used for primary or reference laboratory identification, and only one (a field real-time PCR for a range of potential bioterrorism agents) is approved by the U.S. Food

and Drug Administration (FDA). Detailed phylogeographic DNA sequence data based on culture collections have been accumulated to trace plague evolution, and this system could be adapted in the future to determine real-time clinical plague epidemiology.

In the absence of other positive laboratory diagnostic tests, a retrospective serologic diagnosis may be made on the basis of rising titers of hemagglutinating antibody to F1 antigen. Enzyme-linked immunosorbent assays (ELISAs) for IgG and IgM antibodies to F1 antigen are also available.

The white blood cell (WBC) count is generally raised (to 10,000–20,000/ μL) in plague, with neutrophilic leukocytosis and a left shift (numerous immature neutrophils); in some cases, however, the WBC count is normal or leukopenia develops. WBC counts are occasionally very high, especially in children (>100,000/ μL). Levels of fibrinogen degradation products are elevated in a majority of patients, but platelet counts are usually normal or low-normal. However, disseminated intravascular coagulation, with low platelet counts, prolonged prothrombin times, reduced fibrinogen, and elevated fibrinogen degradation product levels, occurs in a significant minority of patients.

TREATMENT PLAGUE

Guidelines for the treatment of plague are given in [Table 196-2](#). A 10-day course of antimicrobial therapy is recommended. Streptomycin has historically been the parenteral treatment of choice for plague and is approved for this indication by the FDA. Although not yet approved by the FDA for plague, gentamicin has proven safe and effective in clinical trials in Tanzania and Madagascar

TABLE 196-2 GUIDELINES FOR THE TREATMENT OF PLAGUE

Drug	Daily Dose	Interval, h	Route
Gentamicin			
Adult	5 mg/kg ^a	24	IM/IV
	3–5 mg/kg	8 (2 mg/kg loading dose followed by 1.7 mg/kg tid, reduced)	IM/IV
Child	5 mg/kg ^a	24	IM/IV
	7.5 mg/kg	8 (2.5 mg/kg tid)	IM/IV
Streptomycin			
Adult	2 g	12	IM
Child	30 mg/kg	12	IM
Levofloxacin			
Adult and child >50 kg	500 mg	24	PO/IV
Child <50 kg and ≥6 months	8 mg/kg (not to exceed 250 mg/dose)	12	PO/IV
Doxycycline			
Adult	200 mg	12 or 24	PO/IV
Child >8 y	4.4 mg/kg	12 or 24	PO/IV
Tetracycline			
Adult	2 g	6	PO/IV
Child >8 y	25–50 mg/kg	6	PO/IV
Chloramphenicol			
Adult	50 mg/kg	6	PO/IV
Child >1 y	50 mg/kg	6	PO/IV

^aAminoglycoside dose should be adjusted in light of renal function. There are no published trial data for once-daily gentamicin as plague therapy in adults or children, but this regimen is efficacious in gram-negative sepsis of other causes and was successful in a recent outbreak of pneumonic plague in the Democratic Republic of the Congo. Neonates up to 1 week of age and premature infants should receive gentamicin (2.5 mg/kg IV bid).

Source: Dennis DT, Campbell GL: Plague and other *Yersinia* infections, in AS Fauci et al (eds): *Harrison's Principles of Internal Medicine*, 17th ed. 2008, p. 980; Inglesby TV et al: Plague as a biological weapon: medical and public health management. Working Group on Civilian Biodefense. *JAMA* 283:2281, 2000; and FDA Product Label Reference ID 3123374 (www.accessdata.fda.gov/drugsatfda_docs/label/2012/020634s061,020635s067,021721s0281/bl.pdf).

and in retrospectively reviewed cases in the United States. In view of streptomycin's adverse-reaction profile and limited availability, some experts now recommend gentamicin over streptomycin. In 2012, the FDA approved levofloxacin for prophylaxis and treatment of plague (including septicemic and pneumonic disease), making it the first antibiotic approved for a new indication under a regulatory approach based on animal studies alone, known as the Animal Rule. An FDA decision on ciprofloxacin is pending. Levofloxacin is more efficacious than ciprofloxacin for postexposure prophylaxis of inhalational anthrax in animal models and also received FDA approval for this indication (**Chap. 261e**); thus it is approved for multiagent prophylaxis in possible bioterrorism exposures.



While systemic chloramphenicol therapy is available in the resource-poor countries primarily affected by plague, it is less likely to be available or used in high-income countries because of its adverse effect profile. Tetracyclines are also effective and can be given by mouth but are not recommended for children under the age of 7 years because of tooth discoloration. Doxycycline is the tetracycline of choice; at an oral dosage of 100 mg twice daily, this drug was as effective as IM gentamicin (2.5 mg/kg twice daily) in a trial in Tanzania.

Although *Y. pestis* is sensitive to β -lactam drugs in vitro and these drugs have been efficacious against plague in some animal models, the response to penicillins has been poor in some clinical cases; thus β -lactams and macrolides are not generally recommended as first-line therapy. Chloramphenicol, alone or in combination, is recommended for some focal complications of plague (e.g., meningitis, endophthalmitis, myocarditis) because of its tissue penetration properties. Fluoroquinolones, effective in vitro and in animal models, are recommended in guidelines for possible bioterrorism-associated pneumonic plague and are increasingly used in therapy, although the only human efficacy data available so far are from a case report. Animal and in vitro studies suggest that fluoroquinolones other than levofloxacin, at doses used in systemic gram-negative sepsis, should be effective as therapy for plague: e.g., ciprofloxacin (400 mg twice daily IV, 500 mg twice daily by mouth), ofloxacin (400 mg twice daily IV or by mouth), or moxifloxacin (400 mg/d IV or by mouth).

PREVENTION



In endemic areas, the control of plague in humans is based on reduction of the likelihood of being bitten by infected fleas or exposed to infected droplets from either humans or animals with plague pneumonia. In the United States, residence and outdoor activity in rural areas of western states where epizootics occur are the main risk factors for infection. To assess potential risks to humans in specific areas, surveillance for *Y. pestis* infection among animal plague hosts and vectors is carried out regularly as well as in response to observed animal die-offs. Personal protective measures include avoidance of areas where a plague epizootic has been identified and publicized (e.g., by warning signs or closure of campsites). Sick or dead animals should not be handled by the general public. Hunters and zoologists should wear gloves when handling wild-animal carcasses in endemic areas. General measures to avoid rodent fleabite during outdoor activity are appropriate and include the use of insect repellent, insecticide, and protective clothing. General measures to reduce peridomestic and occupational human contact with rodents are advised and include rodent-proofing of buildings and food-waste stores and removal of potential rodent habitats (e.g., woodpiles and junk heaps). Flea control by insecticide treatment of wild rodents is an effective means of minimizing human contact with plague if an epizootic is identified in an area close to human habitation. Any attempt to reduce rodent numbers must be preceded by flea suppression to reduce the migration of infected fleas to human hosts. An oral F1-V subunit vaccine using raccoon poxvirus (RCN) as a vector protects prairie dogs against *Y. pestis* injections and is being investigated for efficacy in preventing disease in wild animals, thus potentially reducing human exposure.

Patients in whom pneumonic plague is suspected should be managed in isolation, with droplet precautions observed until pneumonia is excluded or effective antimicrobial therapy has been given for 48 h. Review of the literature published before the advent of antimicrobial agents suggests that the main infective risk is posed by patients in the final stages of disease who are coughing up sputum with plentiful visible blood and/or pus. Cotton and gauze masks were protective in these circumstances. Current surgical masks capable of barrier protection against droplets, including large respiratory particles, are considered protective; a particulate respirator (e.g., N95 or greater) is not required.

Antimicrobial Prophylaxis Postexposure antimicrobial prophylaxis lasting 7 days is recommended following household, hospital, or other close contact with persons with untreated pneumonic plague. (*Close contact* is defined as contact with a patient at <2 m.) In animal aerosol-infection studies, levofloxacin and ciprofloxacin are associated with higher survival rates than doxycycline (**Table 196-3**).

Immunization Studies with candidate plague vaccines in animal models show that neutralizing antibody provides protection against exposure but that cell-mediated immunity is critical for protection and clearance of *Y. pestis* from the host. A killed whole-cell vaccine used in humans required multiple doses, caused significant local and systemic reactions, and was not protective against pneumonic plague; this vaccine is not currently available in the United States. A live attenuated vaccine based on strain EV76 is still used in countries of the former Soviet Union but has significant side effects. The vaccines closest to licensure are subunit vaccines comprising recombinant F1 (rF1) and various recombinant V (rV) proteins produced in *Escherichia coli*, which are combined either as a fusion protein or as a mixture, purified, and adsorbed to aluminum hydroxide for injection. This combination protects mice and various nonhuman primates in laboratory models of bubonic and pneumonic plague and has been evaluated in phase 2 clinical trials. Special ethical considerations with controlled clinical studies involving plague in humans make prelicensure field efficacy studies unlikely. In the United States, the FDA is therefore prepared to assess plague vaccines for human use under the Animal Rule, using efficacy data and other results from animal studies as well as antibodies and other correlates of immunity from human vaccine recipients (www.fda.gov/BiologicsBloodVaccines/ScienceResearch/BiologicsResearchAreas/ucm127288.htm). Live attenuated

TABLE 196-3 GUIDELINES FOR PLAGUE PROPHYLAXIS

Drug	Daily Dose	Interval, h	Route
Doxycycline			
Adult	200 mg	12 or 24	PO
Child ≥ 8 y	If ≥ 45 kg, give adult dosage; if < 45 kg, give 2.2 mg/kg PO bid (maximum, 200 mg)	12	PO
Tetracycline			
Adult	1–2 g	6 or 12	PO
Child ≥ 8 y	25–50 mg/kg	6 or 12	PO
Levofloxacin			
Adult and child > 50 kg	500 mg	24	PO
Child < 50 kg and ≥ 6 months	8 mg/kg (not to exceed 250 mg/dose)	12	PO
Ciprofloxacin			
Adult	1 g	12	PO
Child	40 mg/kg	12	PO
Trimethoprim-Sulfamethoxazole			
Adult	320 mg	12	PO
Child	40 mg/kg	12	PO


Source: Dennis DT, Campbell GL: Plague and other *Yersinia* infections, in AS Fauci et al (eds): *Harrison's Principles of Internal Medicine*. 2008, p. 980; Inglesby TV et al: Plague as a biological weapon: medical and public health management. Working Group on Civilian Biodefense. *JAMA* 283:2281, 2000; and FDA Drug Product Label Reference ID 3123374 (www.accessdata.fda.gov/drugsatfda_docs/label/2012/020634s061,020635s067,021721s028tbl.pdf).

1076 *Y. pseudotuberculosis* and *Salmonella* strains expressing *Y. pestis*-specific antigens have been shown to be protective in laboratory animal models of bubonic and pneumonic plague and could be delivered by the oral route. A wide variety of other delivery mechanisms for *Y. pestis* antigens are being explored. Antigens other than F1 and V that could be added to subunit vaccines are being investigated. Advances providing impetus for exploration of these antigens are (1) the recovery of F1-negative *Y. pestis* strains from natural sources and (2) the observation that F1 antigen is not required for virulence in primate models of pneumonic plague.

YERSINIOSIS

Yersiniosis is a zoonotic infection with an enteropathogenic *Yersinia* species, usually *Yersinia enterocolitica* or *Y. pseudotuberculosis*. The usual hosts for these organisms are pigs and other wild and domestic animals; humans are usually infected by the oral route, and outbreaks from contaminated food occur. Yersiniosis is most common in childhood and in colder climates. Patients present with abdominal pain and sometimes with diarrhea (which may not occur in up to 50% of cases). *Y. enterocolitica* is more closely associated with terminal ileitis and *Y. pseudotuberculosis* with mesenteric adenitis, but both organisms may cause mesenteric adenitis and symptoms of abdominal pain and tenderness that result in pseudoappendicitis, with the surgical removal of a normal appendix. Diagnosis is based on culture of the organism or convalescent serology. *Y. pseudotuberculosis* and some rarer strains of *Y. enterocolitica* are especially likely to cause systemic infection, which is also particularly common among patients with diabetes or iron overload. Systemic sepsis is treatable with antimicrobial agents, but postinfective arthropathy responds poorly to such therapy. Fourteen other *Yersinia* species are now recognized, but all lack the virulence plasmid pYV common to *Y. pestis*, *Y. pseudotuberculosis*, and *Y. enterocolitica* and are generally considered to be, at most, opportunistic pathogens of humans (*Y. aldovae*, *Y. aleksiciae*, *Y. bercovieri*, *Y. entomophaga*, *Y. frederiksenii*, *Y. intermedia*, *Y. kristensenii*, *Y. massiliensis*, *Y. mollaretii*, *Y. nurmii*, *Y. pekkanenii*, *Y. rohdei*, *Y. similis*, and *Y. ruckeri*). Molecular phylogeny shows that *Y. enterocolitica* is more distantly related to *Y. pseudotuberculosis* than these other *Yersinia* species, and the similar virulence plasmid they share has probably been acquired independently by at least one of the two since the species diverged.

EPIDEMIOLOGY

 ***Y. enterocolitica*** *Y. enterocolitica* is found worldwide and has been isolated from a wide variety of wild and domestic animals and environmental samples, including samples of food and water. In vitro, *Y. enterocolitica* is resistant to predation by the protozoan *Acanthamoeba castellanii* and can survive inside it, suggesting a possible mode of environmental persistence. Strains are differentiated by combined biochemical reactions (biovar) and serogroup. Most clinical infections are associated with serogroups O:3, O:9, and O:5,27, with a declining number of O:8 infections in North America. Some O:8 infections, previously confined to North America, have been reported from Europe and Japan in recent years, and serogroup O:8 now causes a high percentage of yersiniosis cases in Poland. Yersiniosis, mostly due to *Y. enterocolitica*, is the third commonest zoonosis reported in Europe; most reports come from northern Europe, especially Germany and Scandinavia. The incidence is highest among children; children under the age of 4 years are more likely to present with diarrhea than are older children. Abdominal pain with mesenteric adenitis and terminal ileitis is more prominent among older children and adults. Septicemia is more likely in patients with preexisting conditions such as diabetes mellitus, liver disease, any condition involving iron overload (including thalassemia and hemochromatosis), advanced age, malignancy, or HIV/AIDS. As in enteritis of other bacterial etiologies, postinfective complications such as reactive arthritis occur mainly in individuals who are HLA-B27 positive. Erythema nodosum (see Fig. 25e-40) following *Yersinia* infection is not associated with HLA-B27 and is more common among women than among men.

Consumption or preparation of raw pork products (such as chitterlings) and some processed pork products is strongly linked with infection

because a high percentage of pigs carry pathogenic *Y. enterocolitica* strains. Outbreaks of *Y. enterocolitica* infection have been associated with consumption of milk (pasteurized, unpasteurized, and chocolate-flavored) and various foods contaminated with springwater. Person-to-person transmission is suspected in a few cases (e.g., in nosocomial and familial outbreaks) but is much less likely with *Y. enterocolitica* than with other causes of gastrointestinal infection, such as *Salmonella*. A multivariate analysis indicates that contact with companion animals is a risk factor for *Y. enterocolitica* infection among children in Sweden, and low-level colonization of dogs and cats with *Y. enterocolitica* has been reported. Transfusion-associated septicemia due to *Y. enterocolitica*, while recognized as a very rare but frequently fatal event for over 30 years, has been difficult to eradicate.



Y. pseudotuberculosis *Y. pseudotuberculosis* is less frequently reported as a cause of human disease than *Y. enterocolitica*, and infection with *Y. pseudotuberculosis* is more likely to present as fever and abdominal pain due to mesenteric lymphadenitis. This organism is associated with wild mammals (rodents, rabbits, and deer), birds, and domestic pigs. Strains are differentiated by combined biochemical reactions (biovar) and serogroup. Although outbreaks are generally rare, several have recently occurred in Finland in association with consumption of lettuce and raw carrots.

PATHOGENESIS

The usual route of infection is oral. Studies with both *Y. enterocolitica* and *Y. pseudotuberculosis* in animal models suggest that initial replication in the small intestine is followed by invasion of Peyer's patches of the distal ileum via M cells, with onward spread to mesenteric lymph nodes. The liver and spleen can also be involved after oral infection. The characteristic histologic appearance of enteropathogenic yersiniae after invasion of host tissues is as extracellular microabscesses surrounded by an epithelioid granulomatous lesion.

Experiments involving oral infection of mice with tagged *Y. enterocolitica* show that only a very small proportion of bacteria in the gut invade tissues. Individual bacterial clones from an orally inoculated pool give rise to each microabscess in a Peyer's patch, and the host restricts the invasion of previously infected Peyer's patches. A prior model positing progressive bacterial spread from Peyer's patches and mesenteric lymph nodes to the liver and spleen appears to be inaccurate: spread of individually tagged clones of *Y. pseudotuberculosis* to the liver and spleen of mice occurs independently of regional lymph node colonization and in mice lacking Peyer's patches.



Invasion requires the expression of several nonfimbrial adhesins, such as invasin (Inv) and—in *Y. pseudotuberculosis*—*Yersinia* adhesin A (YadA). Inv interacts directly with $\beta 1$ integrins, which are expressed on the apical surfaces of M cells but not enterocytes. YadA of *Y. pseudotuberculosis* interacts with extracellular matrix proteins such as collagen and fibronectin to facilitate host cell integrin association and invasion. YadA of *Y. enterocolitica* lacks a crucial N-terminal region and binds collagen and laminin but not fibronectin and does not cause invasion. Inv is chromosomally encoded, whereas YadA is encoded on the virulence plasmid pYV. YadA helps to confer serum resistance by binding host complement regulators such as factor H and C4-binding protein. Another chromosomal gene, *ail* (attachment and invasion locus), encodes the extracellular protein Ail, which also confers serum resistance by binding these complement regulators.

By binding to host cell surfaces, YadA allows targeting of immune effector cells by the pYV plasmid-encoded type III secretion system (*injectisome*). As a consequence, the host's innate immune response is altered; toxins (*Yersinia* outer proteins, or Yops) are injected into host macrophages, neutrophils, and dendritic cells, affecting signal transduction pathways, resulting in reduced phagocytosis and inhibited production of reactive oxygen species by neutrophils, and triggering apoptosis of macrophages. Other factors functional in invasive disease include yersiniabactin (Ybt), a siderophore produced by some strains of *Y. pseudotuberculosis* and *Y. enterocolitica* as well as other Enterobacteriaceae. Yersiniabactin allows bacteria to access iron from saturated lactoferrin during infection and reduces the production of reactive oxygen species by innate immune effector cells, thereby

decreasing bacterial killing. *Y. pseudotuberculosis* and *Y. pestis* make other siderophores in addition to yersiniabactin.

CLINICAL MANIFESTATIONS

Self-limiting diarrhea is the most common reported presentation in infection with pathogenic *Y. enterocolitica*, especially in children under the age of 4, who form the single largest group in most case series. Blood may be detected in diarrheal stool. Older children and adults are more likely than younger children to present with abdominal pain, which can be localized to the right iliac fossa—a situation that often leads to laparotomy for presumed appendicitis (pseudoappendicitis). Appendectomy is not indicated for *Yersinia* infection causing pseudoappendicitis. Thickening of the terminal ileum and cecum is seen on endoscopy and ultrasound, with elevated round or oval lesions that may overlie Peyer's patches. Mesenteric lymph nodes are enlarged. Ulcerations of the mucosa are noted on endoscopy. Gastrointestinal complications include granulomatous appendicitis, a chronic inflammatory condition affecting the appendix that is responsible for $\leq 2\%$ of cases of appendicitis; *Yersinia* is involved in a minority of cases. *Y. enterocolitica* infection can present as acute pharyngitis with or without other gastrointestinal symptoms. Fatal *Y. enterocolitica* pharyngitis has been recorded. Mycotic aneurysm can follow *Y. enterocolitica* bacteremia, as can focal infection (abscess) in many other sites and body compartments (liver, spleen, kidney, bone, meninges, endocardium).



In all age groups, *Y. pseudotuberculosis* infection is more likely to present as abdominal pain and fever than as diarrhea. A superantigenic toxin—*Y. pseudotuberculosis* mitogen (YPM)—is produced by strains seen in eastern Russia in association with Far Eastern scarlet-like fever, a childhood illness with desquamating rash, arthralgia, and toxic shock. A similar illness is recognized in Japan (Izumi fever) and Korea. Similarities have been noted with Kawasaki disease, the idiopathic acute systematic vasculitis of childhood. There is an epidemiologic link between exposure of populations to superantigen-positive *Y. pseudotuberculosis* and an elevated incidence of Kawasaki disease.

Y. enterocolitica or *Y. pseudotuberculosis* septicemia presents as a severe illness with fever and leukocytosis, often without localizing features, and is significantly associated with predisposing conditions such as diabetes mellitus, liver disease, and iron overload. Hemochromatosis combines several of these risk factors. Administration of iron chelators like desferrioxamine, which provide iron accessible to *Yersinia* (and have an inhibitory effect on neutrophil function), may result in *Yersinia* septicemia in patients with iron overload who presumably have an otherwise mild gastrointestinal infection. HIV/AIDS has been associated with *Y. pseudotuberculosis* septicemia. The unusual phenomenon of transfusion-associated septicemia is linked to the ability of *Y. enterocolitica* to multiply at refrigerator temperature (*psychrotrophy*). Typically, the transfused unit has been stored for >20 days, and it is believed that small numbers of yersiniae from an apparently healthy donor with subclinical bacteremia are amplified to very high numbers by growth inside the bag at $\leq 4^{\circ}\text{C}$, with consequent septic shock after transfusion. A method for preventing this very rare event (i.e., a range of 1 case in 500,000 to 1 case in several million transfused units in countries such as the United States and France) without unacceptable restriction in the blood supply has not yet been devised.

POSTINFECTIVE PHENOMENA



Like other invasive infections of intestinal origin (salmonellosis, shigellosis), reactive arthritis (articular arthritis of multiple joints developing within 2–4 weeks of a preceding infection) results from autoimmune activity initiated by the deposition of bacterial components (not viable bacteria) in joints in combination with the immune response to invading bacteria. The majority of individuals affected by reactive arthritis due to *Yersinia* are HLA-B27 positive. Myocarditis with electrocardiographic ST-segment abnormalities may occur with *Yersinia*-associated reactive arthritis. Most *Yersinia*-associated cases follow *Y. enterocolitica* infection (presumably because it is more common than infection with other species), but *Y. pseudotuberculosis*-associated reactive arthritis is also well documented in

Finland, where sporadic and outbreak infections with *Y. pseudotuberculosis* are more common than in other countries. Of infected individuals identified in a recent *Y. pseudotuberculosis* serotype O:3 outbreak in Finland, 12% developed reactive arthritis affecting the small joints of the hands and feet, knees, ankles, and shoulders and lasting >6 months in most cases. Erythema nodosum (see Fig. 25e–40) occurs after *Yersinia* infection (more commonly in women) with no evidence of HLA-B27 linkage.

There is a long-standing association between antithyroid and anti-*Yersinia* antibodies. Antibody evidence of prior *Y. enterocolitica* infection in Graves' disease and increased levels of antithyroid antibody in patients with *Y. enterocolitica* antibodies were first noted in the 1970s. *Y. enterocolitica* contains a thyroid-stimulating hormone (TSH)-binding site that is recognized by anti-TSH antibodies from Graves' disease patients. Raised titers of antibodies to *Y. enterocolitica* whole cells and Yops have been found in some series of Graves' disease patients but not in others. One Danish study of twins found no evidence of an association between asymptomatic *Yersinia* infection (as evidenced by anti-Yop antibody titers) and antithyroid antibodies in euthyroid individuals, while another Danish study of twins with and without Graves' disease found that increased anti-Yop antibody titers were associated with Graves' disease. It remains unclear whether this cross-reactivity is significant in the etiology of Graves' disease.

LABORATORY DIAGNOSIS

Standard laboratory culture methods can be used to isolate enteropathogenic *Yersinia* species from sterile samples, including blood and cerebrospinal fluid. Culture on specific selective media (CIN agar), with or without pre-enrichment in broth or phosphate-buffered saline at either 4°C or 16°C , is the basis of most schema for isolation of yersiniae from stool or other nonsterile samples. Outside known high-incidence areas, specific culture may be carried out by laboratories only upon request. Virulence plasmid-negative strains of *Y. enterocolitica* can be isolated from cultures of stool from asymptomatic individuals, especially after cold enrichment. These strains usually differ in biotype (typically biovar 1a) from virulence plasmid-possessing strains; although some display apparent pathogenicity in a mouse model, virulence plasmid-negative strains are not commonly accepted as human pathogens. Because of the frequency with which the virulence plasmid is lost on laboratory subculture, combined biochemical identification (with biotyping according to a standard schema) and serologic identification are usually required to interpret the significance of an isolate of *Y. enterocolitica* from a nonsterile site. Most pathogenic *Y. enterocolitica* strains currently isolated from humans are of serogroup O:3/biovar 4 or serogroup O:9/biovar 2; this pattern holds even in the United States, where serogroup O:8/biovar 1B strains were previously predominant. Many self-validated multiplex PCR screens for detection of *Y. enterocolitica* in clinical samples—and rather more for its detection in food—have been described, but none of these assays is widely used outside its originating laboratory. Some CE-marked real-time PCR kits are now available in Europe for the diagnosis of yersiniosis in animals; as molecular diagnosis of enteric infection becomes more routine in human disease, it is likely that *Y. enterocolitica* will be included in diagnostic multiplex PCR screens of feces. Because of the presence of Ail in biovar 1a strains, this antigen cannot be used alone in diagnostic assays. A standard for PCR detection in food samples is being prepared by the International Organization for Standardization.

Agglutinating or ELISA antibody titers to specific O-antigen types are used in the retrospective diagnosis of both *Y. enterocolitica* and *Y. pseudotuberculosis* infections. IgA and IgG antibodies persist in patients with reactive arthritis. Serologic cross-reactions between *Y. enterocolitica* serogroup O:9 and *Brucella* are due to the similarity of their lipopolysaccharide structures. Multiple assays are required to cover even the predominant serogroups (*Y. enterocolitica* O:3, O:5, 27, and O:9; *Y. pseudotuberculosis* O:1a, O:1b, and O:3), and these assays are generally available only in reference laboratories. ELISA and western blot tests for antibodies to Yops, which are expressed by all pathogenic strains of *Y. enterocolitica* and *Y. pseudotuberculosis*, are

1078 also available; most of the positivity in these assays probably relates to previous infection with *Y. enterocolitica*.

TREATMENT YERSINIOSIS

Most cases of diarrhea caused by enteropathogenic *Yersinia* are self-limiting. Data from clinical trials do not support antimicrobial treatment for adults or children with *Y. enterocolitica* diarrhea. Systemic infections with bacteremia or focal infections outside the gastrointestinal tract generally require antimicrobial therapy. Infants <3 months of age with documented *Y. enterocolitica* infection may require antimicrobial treatment because of the increased likelihood of bacteremia in this age group. *Y. enterocolitica* strains nearly always express β -lactamases. Because of the relative rarity of systemic *Y. enterocolitica* infection, there are no clinical trial data to guide antimicrobial choice or to suggest the optimal dose and duration of therapy. On the basis of retrospective case series and in vitro sensitivity data, fluoroquinolone therapy is effective for bacteremia in adults; for example, ciprofloxacin is given at a typical dose of 500 mg twice daily by mouth or 400 mg twice daily IV for at least 2 weeks (longer if positive blood cultures persist). A third-generation cephalosporin is an alternative—e.g., cefotaxime (typical dose, 6–8 g/d in three or four divided doses). In children, third-generation cephalosporins are effective; for example, cefotaxime is given to children ≥ 1 month of age at a typical dose of 75–100 mg/kg per day in three or four divided doses, with an increase to 150–200 mg/kg per day in severe cases (maximal daily dose, 8–10 g). Amoxicillin and amoxicillin/clavulanate have shown poor efficacy in case series. Trimethoprim-sulfamethoxazole, gentamicin, and imipenem are all active in vitro. *Y. pseudotuberculosis* strains do not express β -lactamase but are intrinsically resistant to polymyxin. Because human infection with *Y. pseudotuberculosis* is less common than that with *Y. enterocolitica*, less case information is available; however, studies in mice suggest that ampicillin is ineffective. Drugs similar to those used against *Y. enterocolitica* should be used. The best results have been obtained with a quinolone.

Some trials of treatment for reactive arthritis (with a large proportion of cases due to *Yersinia*) found that 3 months of oral ciprofloxacin therapy did not affect outcome. One trial in which the same therapy was given specifically for *Y. enterocolitica*-reactive arthritis found that, while outcome indeed was not affected, there was a trend toward faster remission of symptoms in the treated group. Follow-up 4–7 years after initial antibiotic treatment of reactive arthritis (predominantly following *Salmonella* and *Yersinia* infections) demonstrated apparent efficacy in the prevention of chronic arthritis in HLA-B27-positive individuals. A trial showing that azithromycin therapy did not affect outcome in reactive arthritis included cases believed to follow yersiniosis, although no breakdown of cases was provided. A Cochrane review evaluating the use of antibiotics for reactive arthritis is in progress.

PREVENTION AND CONTROL



Current control measures are similar to those used against other enteric pathogens like *Salmonella* and *Campylobacter*, which colonize the intestine of food animals. The focus is on safe handling and processing of food. No vaccine is effective in preventing intestinal colonization of food animals by enteropathogenic *Yersinia*. Consumption of food made from raw pork (which is popular in Germany and Belgium) should be discouraged at present because it is not possible to eliminate contamination with the enteropathogenic *Yersinia* strains found worldwide in pigs. Exposure of infants to raw pig intestine during domestic preparation of chitterlings is inadvisable. Modification of abattoir technique in Scandinavian countries from the 1990s onward included the removal of pig intestines in a closed plastic bag; levels of carcass contamination with *Y. enterocolitica* were reduced, but such contamination was not eliminated. Experimental pig herds free of pathogenic *Y. enterocolitica* O:3 (and also of *Salmonella*, *Toxoplasma*, and *Trichinella*) have been established in Norway and

may be commercialized in the future because of their enhanced safety. In the food industry, vigilance is required because of the potential for large outbreaks if small numbers of enteropathogenic yersiniae contaminate any ready-to-eat food whose safe preservation is based on refrigeration before consumption.

The rare phenomenon of contamination of blood for transfusion has proved impossible to eradicate. However, leukodepletion is now practiced in most blood transfusion centers, primarily to prevent non-hemolytic febrile transfusion reactions and alloimmunization against HLA antigens. This measure reduces but does not eliminate the risk of *Yersinia* blood contamination.

Notification of yersiniosis is now obligatory in some countries.

197 Bartonella Infections, Including Cat-Scratch Disease

Michael Giladi, Moshe Ephros

Bartonella species are fastidious, facultative intracellular, slow-growing, gram-negative bacteria that cause a broad spectrum of diseases in humans. This genus includes more than 30 distinct species or subspecies, of which at least 16 have been recognized as confirmed or potential human pathogens; *Bartonella bacilliformis*, *Bartonella quintana*, and *Bartonella henselae* are most commonly identified (Table 197-1). Most *Bartonella* species have successfully adapted to survival in specific domestic or wild mammals. Prolonged intraerythrocytic infection in these animals creates a niche where the bacteria are protected from both innate and adaptive immunity and which serves as a reservoir for human infections. *Bartonella* characteristically evades the host immune system by modification of its virulence factors (e.g., lipopolysaccharides or flagella) and by attenuation of the immune response. *B. bacilliformis* and *B. quintana*, which are not zoonotic, are exceptions. Arthropod vectors are often involved. Isolation and characterization of *Bartonella* species are difficult and require special techniques. Clinical presentation generally depends on both the infecting *Bartonella* species and the immune status of the infected individual. *Bartonella* species are susceptible to many antibiotics in vitro; however, clinical responses to therapy and studies in animal models suggest that the minimal inhibitory concentrations of many antimicrobial agents correlate poorly with the drugs' in vivo efficacies in patients with *Bartonella* infections.

CAT-SCRATCH DISEASE

DEFINITION AND ETIOLOGY

Usually a self-limited illness, cat-scratch disease (CSD) has two general clinical presentations. *Typical* CSD, the more common, is characterized by subacute regional lymphadenopathy; *atypical* CSD is the collective designation for numerous extranodal manifestations involving various organs. *B. henselae* is the principal etiologic agent of CSD. Rare cases have been associated with *Afipia felis* and other *Bartonella* species.

EPIDEMIOLOGY

CSD occurs worldwide, favoring warm and humid climates. In temperate climates, incidence peaks during fall and winter; in the tropics, disease occurs year-round. Adults are affected nearly as frequently as children. Intrafamilial clustering is rare, and person-to-person transmission does not occur. Apparently healthy cats constitute the major reservoir of *B. henselae*, and cat fleas (*Ctenocephalides felis*) may be responsible for cat-to-cat transmission. CSD usually follows contact with cats (especially kittens), but other animals (e.g., dogs) have been implicated as possible reservoirs in rare instances. In the United States, the estimated disease incidence is ~10 cases per 100,000 population. About 10% of patients are hospitalized.

TABLE 197-1 BARTONELLA SPECIES KNOWN OR SUSPECTED TO BE HUMAN PATHOGENS

<i>Bartonella</i> Species ^a	Disease ^b	Reservoir Host ^c	Arthropod Vector
<i>B. henselae</i>	Cat-scratch disease, bacillary angiomatosis, bacillary peliosis, bacteremia, endocarditis	Cats, other felines	Cat fleas (<i>Ctenocephalides felis</i>): associated with cat-to-cat, but not with cat-to-human, transmission
<i>B. quintana</i>	Trench fever, chronic bacteremia, bacillary angiomatosis, endocarditis	Humans	Human body lice (<i>Pediculus humanus corporis</i>)
<i>B. bacilliformis</i>	Carrión's disease	Humans	Sandflies (<i>Lutzomyia verrucarum</i>)
<i>B. elizabethae</i>	Endocarditis	Rats, dogs	Unknown
<i>B. grahamii</i> ^d	Lymphadenopathy	Mice, voles	Fleas
<i>B. vinsonii</i> subsp. <i>arupensis</i>	Endocarditis, febrile illness	Mice, dogs	Ticks
<i>B. vinsonii</i> subsp. <i>berkhoffii</i>	Endocarditis	Domestic dogs, coyotes, gray foxes	Ticks
<i>B. washoensis</i>	Myocarditis, meningitis	Squirrels, possibly other rodents	Fleas
<i>B. alsatica</i>	Endocarditis	Rabbits	Unknown
<i>B. koehlerae</i>	Endocarditis	Cats	Unknown
<i>B. clarridgeiae</i>	Possibly cat-scratch disease	Cats	Unknown
<i>B. rochalimae</i>	Bacteremia, fever, splenomegaly	Unknown	Possibly fleas
<i>B. tamiae</i>	Bacteremia, fever, myalgia, rash	Unknown	Unknown
<i>Candidatus B. melophagi</i> ^e	Various clinical manifestations	Unknown	Unknown
<i>Candidatus B. mayotimonensis</i> ^e	Endocarditis	Unknown	Unknown
<i>Candidatus B. ancashi</i> ^e	Verruga peruana–like illness	Unknown	Unknown

^aMany other *Bartonella* species exist but are not recognized as human pathogens. ^b*B. henselae*, *B. vinsonii* subsp. *berkhoffii*, *B. koehlerae*, *Candidatus B. melophagi*, or more than one *Bartonella* spp. (co-infection) were detected in blood samples from patients with extensive arthropod and other animal exposure who presented with various chronic neurologic or neurocognitive syndromes. The role of these pathogens in these patients needs further study. ^cAnimals are implicated when existing evidence supports their infection with *Bartonella* species. Data supporting animal-to-human transmission may be lacking. ^dRetinitis may also be associated with *B. grahamii*. ^e*Candidatus* is a taxonomic status for bacteria that cannot be described in sufficient detail to warrant establishment of a novel taxon or cannot be cultured or propagated in culture media. The phylogenetic relatedness of these bacteria has been determined by gene amplification and sequence analysis.

PATHOGENESIS

Inoculation of *B. henselae*, possibly via contaminated flea feces, usually results from a cat scratch or bite. Infection of mucous membranes or conjunctivae via droplets or licking may occur as well. With lymphatic drainage to one or more regional lymph nodes in immunocompetent hosts, a T_H1 response can result in necrotizing granulomatous lymphadenitis. Dendritic cells, along with their associated chemokines, play a role in the host inflammatory response and granuloma formation.

CLINICAL MANIFESTATIONS AND PROGNOSIS

Of patients with CSD, 85–90% have typical disease. The primary lesion, a small (0.3- to 1-cm) painless erythematous papule or pustule, develops at the inoculation site (usually the site of a scratch or a bite) within days to 2 weeks in about one-third to two-thirds of patients (Fig. 197-1A, B). Lymphadenopathy develops 1–3 weeks or longer after cat contact. The affected lymph node(s) are enlarged and usually painful, sometimes have overlying erythema, and suppurate in 10–15% of cases (Fig. 197-1C, D, and E). Axillary/epitrochlear nodes are most commonly involved; next in frequency are head/neck nodes and then inguinal/femoral nodes. Approximately 50% of patients have fever, malaise, and anorexia. A smaller proportion experience weight loss and night sweats mimicking the presentation of lymphoma. Fever is usually low-grade but infrequently rises to ≥39°C. Resolution is slow, requiring weeks (for fever, pain, and accompanying signs and symptoms) to months (for node shrinkage).

Atypical CSD occurs in 10–15% of patients as extranodal or complicated disease in the absence or presence of lymphadenopathy. Atypical disease includes Parinaud's oculoglandular syndrome (granulomatous conjunctivitis with ipsilateral preauricular lymphadenitis; Fig. 197-1E), granulomatous hepatitis/splenitis, neuroretinitis (often presenting as unilateral deterioration of vision; Fig. 197-1F), and other ophthalmologic manifestations. In addition, neurologic involvement (encephalopathy, seizures, myelitis, radiculitis, cerebellitis, facial and other cranial or peripheral palsies), fever of unknown origin, debilitating myalgia, arthritis or arthralgia (affecting mostly women >20 years old), osteomyelitis (including multifocal disease), tendinitis, neuralgia, and dermatologic manifestations (including erythema nodosum [see Fig. 25e-40], sometimes accompanying arthropathy) occur. Other manifestations and syndromes (pneumonitis, pleural effusion, idiopathic thrombocytopenic purpura, Henoch-Schönlein

purpura, erythema multiforme [see Fig. 25e-25], hypercalcemia, glomerulonephritis, myocarditis) have also been associated with CSD. In elderly patients (>60 years old), lymphadenopathy is more often absent but encephalitis and fever of unknown origin are more common than in younger patients. In immunocompetent individuals, CSD—whether typical or atypical—usually resolves without treatment and without sequelae. Lifelong immunity is the rule.

DIAGNOSIS

Routine laboratory tests usually yield normal or nonspecific results. Histopathology initially shows lymphoid hyperplasia and later demonstrates stellate granulomata with necrosis, coalescing microabscesses, and occasional multinucleated giant cells—findings that, although nonspecific, may narrow the differential diagnosis. Serologic testing (immunofluorescence or enzyme immunoassay) is the most commonly used laboratory diagnostic approach, with variable sensitivity and specificity. Seroconversion may take a few weeks. Other tests are of low sensitivity (culture, Warthin-Starry silver staining), of low specificity (cytology, histopathology), or of limited availability in routine diagnostic laboratories (polymerase chain reaction [PCR], immunohistochemistry). PCR of lymph node tissue, pus, or the primary inoculation lesion is highly sensitive and specific and is particularly useful for definitive and rapid diagnosis in seronegative patients.

APPROACH TO THE PATIENT: Cat-Scratch Disease

A history of cat contact, a primary inoculation lesion, and regional lymphadenopathy are highly suggestive of CSD. A characteristic clinical course and corroborative laboratory tests make the diagnosis very likely. Conversely, when acute- and convalescent-phase sera are negative (as is the case in 10–20% of CSD patients), when spontaneous regression of lymph node size does not occur, and particularly when constitutional symptoms persist, malignancy must be ruled out. Pyogenic lymphadenitis, mycobacterial infection, brucellosis, syphilis, tularemia, plague, toxoplasmosis, sporotrichosis, and histoplasmosis should also be considered. In clinically suspected CSD in a seronegative individual, fine-needle aspiration may be adequate and PCR can confirm the diagnosis. When data are

less supportive of CSD, lymph node biopsy rather than fine-needle aspiration is preferred. In seronegative CSD patients with lymphadenopathy and severe complications (e.g., encephalitis or neuroretinitis), early biopsy is important to establish a specific diagnosis.

TREATMENT CAT-SCRATCH DISEASE

(Table 197-2) Treatment regimens are based on only minimal data. Suppurative nodes should be drained by large-bore needle aspiration and not by incision and drainage in order to avoid chronic draining tracts. Immunocompromised patients must always be treated with systemic antimicrobials.

PREVENTION

Avoiding cats (especially kittens) and instituting flea control are options for immunocompromised patients and for patients with valvular heart disease.

TRENCH FEVER AND CHRONIC BACTEREMIA

DEFINITION AND ETIOLOGY

Trench fever, also known as *5-day fever* or *quintan fever*, is a febrile illness caused by *B. quintana*. It was first described as an epidemic in the trenches of World War I and recently reemerged as chronic bacteremia seen most often in homeless people (also referred to in the latter setting as *urban* or *contemporary trench fever*).



A



B



C

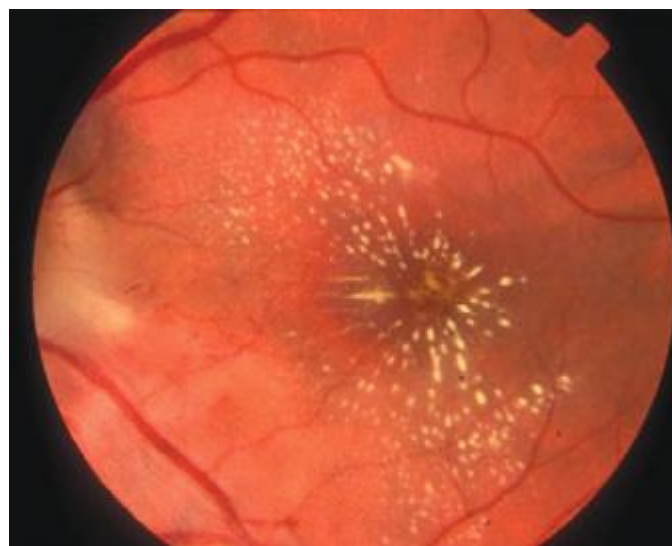


D

FIGURE 197-1 Manifestations of cat-scratch disease. **A.** Primary inoculation lesion. Axillary and epitrochlear lymphadenitis appeared 2 weeks later. **B.** Primary inoculation lesion. Submental lymphadenitis appeared 10 days later. **C.** Axillary lymphadenopathy of 2 weeks' duration. The overlying skin appears normal. **D.** Cervical lymphadenopathy of 6 weeks' duration. The overlying skin is red. Thick, odorless pus (12 mL) was aspirated. **E.** Preauricular lymphadenopathy. **F.** Left-eye neuroretinitis. Note papilledema and stellate macular exudates ("macular star").



E



F

FIGURE 197-1 (Continued)

EPIDEMIOLOGY



In addition to epidemics during World Wars I and II, sporadic outbreaks of trench fever have been reported in many regions of the world. The human body louse has been identified as the vector and humans as the only known reservoir. After a hiatus of several decades during which trench fever was almost forgotten, small clusters of cases of *B. quintana* chronic bacteremia were reported sporadically, primarily from the United States and France, in HIV-uninfected homeless people. Alcoholism and louse infestation were identified as risk factors.

CLINICAL MANIFESTATIONS

The typical incubation period is 15–25 days (range, 3–38 days). “Classical” trench fever, as described in 1919, ranges from a mild febrile illness to a recurrent or protracted and debilitating disease. Onset may be abrupt or preceded by a prodrome of several days. Fever is often periodic, lasting 4–5 days with 5-day (range, 3- to 8-day) intervals between episodes. Other symptoms and signs include headache, back and limb pain, profuse sweating, shivering, myalgia, arthralgia, splenomegaly, a maculopapular rash in occasional cases, and nuchal rigidity in some cases. Untreated, the disease usually lasts

TABLE 197-2 ANTIMICROBIAL THERAPY FOR DISEASE CAUSED BY *BARTONELLA* SPECIES IN ADULTS

Disease	Antimicrobial Therapy
Typical cat-scratch disease	Not routinely indicated; for patients with extensive lymphadenopathy, consider azithromycin (500 mg PO on day 1, then 250 mg PO qd for 4 days)
Cat-scratch disease retinitis	Doxycycline (100 mg PO bid) <i>plus</i> rifampin (300 mg PO bid) for 4–6 weeks
Other atypical cat-scratch disease manifestations ^a	As per retinitis; treatment duration should be individualized
Trench fever or chronic bacteremia with <i>B. quintana</i>	Gentamicin (3 mg/kg IV qd for 14 days) <i>plus</i> doxycycline (200 mg PO qd or 100 mg PO bid for 6 weeks)
Suspected <i>Bartonella</i> endocarditis	Gentamicin ^b (1 mg/kg IV q8h for ≥14 days) <i>plus</i> doxycycline (100 mg PO/IV bid for 6 weeks ^c) <i>plus</i> ceftriaxone (2 g IV qd for 6 weeks)
Confirmed <i>Bartonella</i> endocarditis	As for suspected <i>Bartonella</i> endocarditis <i>minus</i> ceftriaxone
Bacillary angiomatosis	Erythromycin ^d (500 mg PO qid for 3 months) <i>or</i> Doxycycline (100 mg PO bid for 3 months)
Bacillary peliosis	Erythromycin ^d (500 mg PO qid for 4 months) <i>or</i> Doxycycline (100 mg PO bid for 4 months)
Carrión's disease	
Oroya fever	Chloramphenicol (500 mg PO/IV qid for 14 days) <i>plus</i> another antibiotic (β-lactam preferred) <i>or</i> Ciprofloxacin (500 mg PO bid for 10 days)
Verruga peruana	Rifampin (10 mg/kg PO qd, to a maximum of 600 mg, for 14 days) <i>or</i> Streptomycin (15–20 mg/kg IM qd for 10 days)

^aData on treatment efficacy for encephalitis and hepatosplenic CSD are lacking. Therapy similar to that given for retinitis is reasonable. ^bSome experts recommend gentamicin at 3 mg/kg IV qd. If gentamicin is contraindicated, rifampin (300 mg PO bid) can be added to doxycycline for documented *Bartonella* endocarditis. ^cSome experts recommend extending oral doxycycline therapy for 3–6 months. ^dOther macrolides are probably effective and may be substituted for erythromycin or doxycycline.

Source: Recommendations are modified from JM Rolain et al: Antimicrob Agents Chemother 48:1921, 2004.

1082 4–6 weeks. Death is rare. The clinical spectrum of *B. quintana* bacteremia in homeless people ranges from asymptomatic infection to a febrile illness with headache, severe leg pain, and thrombocytopenia. Endocarditis sometimes develops.

DIAGNOSIS


Definitive diagnosis requires isolation of *B. quintana* by blood culture. Some patients have positive blood cultures for several weeks. Patients with acute trench fever typically develop significant titers of antibody to *Bartonella*, whereas those with chronic *B. quintana* bacteremia may be seronegative. Patients with high titers of IgG antibodies should be evaluated for endocarditis. In epidemics, trench fever should be differentiated from epidemic louse-borne typhus and relapsing fever, which occur under similar conditions and share many features.

TREATMENT *B. QUINTANA* BACTEREMIA


(Table 197–2) In a small, randomized, placebo-controlled trial involving homeless people with *B. quintana* bacteremia, therapy with gentamicin and doxycycline was superior to administration of placebo in eradicating bacteremia. Treatment of bacteremia is important even in clinically mild cases to prevent endocarditis. Optimal therapy for trench fever without documented bacteremia is uncertain.

BARTONELLA ENDOCARDITIS

DEFINITION AND ETIOLOGY

 *Coxiella burnetii* (Chap. 211) and *Bartonella* species are the most common pathogens in culture-negative endocarditis (Chap. 155). In France, for example, *Bartonella* species were identified as the etiologic agents in 28% of 348 cases of culture-negative endocarditis. Prevalence, however, varies by geographic location and epidemiologic setting. In addition to *B. quintana* and *B. henselae* (the most common *Bartonella* species implicated in endocarditis, the former more commonly than the latter), other *Bartonella* species have reportedly caused rare cases (Table 197-1).

EPIDEMIOLOGY

 *Bartonella* endocarditis has been reported worldwide. Most patients are adults; more are male than female. Risk factors associated with *B. quintana* endocarditis include homelessness, alcoholism, and body louse infestation; however, individuals with no risk factors have had *Bartonella* endocarditis diagnosed as well. *B. henselae* endocarditis is associated with exposure to cats. Most cases involve native rather than prosthetic valves; the aortic valve accounts for ~60% of cases. Patients with *B. henselae* endocarditis usually have preexisting valvulopathy, whereas *B. quintana* often infects normal valves.

CLINICAL MANIFESTATIONS

Clinical manifestations are usually characteristic of subacute endocarditis of any etiology. However, a substantial number of patients have a prolonged, minimally febrile or even afebrile indolent illness, with mild nonspecific symptoms lasting weeks or months before the diagnosis is made. Initial echocardiography may not show vegetations. Acute, aggressive disease is rare.

DIAGNOSIS

Blood cultures, even with use of special techniques (lysis centrifugation or EDTA-containing tubes), are positive in only ~25% of cases—mostly those caused by *B. quintana* and only rarely those caused by *B. henselae*. Prolonged incubation of cultures (up to 6 weeks) is required. Serologic tests—either immunofluorescence or enzyme immunoassay—usually demonstrate high-titer IgG antibodies to *Bartonella*. Because of cross-antigenicity, serology does not distinguish between *B. quintana* and *B. henselae* and may also be low-titer cross-reactive with other pathogens, such as *C. burnetii* and *Chlamydia* species. Identification of *Bartonella* to the species level is usually accomplished by application of PCR-based methods to valve tissue.

TREATMENT *BARTONELLA* ENDOCARDITIS

(Table 197-2) For patients with culture-negative endocarditis suspected to be due to *Bartonella* species, empirical treatment consists of gentamicin, doxycycline, and ceftriaxone; the major role of ceftriaxone in this regimen is to adequately treat other potential causes of culture-negative endocarditis, including members of the HACEK group (Chap. 183e). Once a diagnosis of *Bartonella* endocarditis has been established, ceftriaxone is discontinued. Aminoglycosides, the only antibiotics known to be bactericidal against *Bartonella*, should be included in the regimen for ≥2 weeks. Indications for valvular surgery are the same as in subacute endocarditis due to other pathogens; however, the proportion of patients who undergo surgery (~60%) is high, probably as a consequence of delayed diagnosis.

BACILLARY ANGIOMATOSIS AND PELIOSIS

DEFINITION AND ETIOLOGY

Bacillary angiomatosis (sometimes called *bacillary epithelioid angiomatosis* or *epithelioid angiomatosis*) is a disease of severely immunocompromised patients, is caused by *B. henselae* or *B. quintana*, and is characterized by neovascular proliferative lesions involving the skin and other organs. Both species cause cutaneous lesions; hepatosplenic lesions are caused only by *B. henselae*, whereas subcutaneous and lytic bone lesions are more frequently associated with *B. quintana*. Bacillary peliosis is a closely related angioproliferative disorder caused by *B. henselae* and involving primarily the liver (peliosis hepatis) but also the spleen and lymph nodes. Bacillary peliosis is characterized by blood-filled cystic structures whose size ranges from microscopic to several millimeters.

EPIDEMIOLOGY

Bacillary angiomatosis and bacillary peliosis occur primarily in HIV-infected persons (Chap. 226) with CD4+ T cell counts <100/μL but also affect other immunosuppressed patients and, in rare instances, immunocompetent patients. The previously reported incidence of ~1 case per 1000 HIV-infected persons is now lower; the decrease is most likely attributable to effective antiretroviral therapy and the routine use of rifabutin and macrolides to prevent *Mycobacterium avium* complex infection in AIDS patients. Contact with cats or cat fleas increases the risk of *B. henselae* infection. Risk factors for *B. quintana* infection are low income, homelessness, and body louse infestation.

CLINICAL MANIFESTATIONS

Bacillary angiomatosis presents most commonly as one or more cutaneous lesions that are not painful and that may be tan, red, or purple in color. Subcutaneous masses or nodules, superficial ulcerated plaques (Fig. 197-2), and verrucous growths are also seen. Nodular forms resemble those seen in fungal or mycobacterial infections. Subcutaneous nodules are often tender. Painful osseous lesions, most often involving long bones, may underlie cutaneous lesions and occasionally develop in their absence. In rare cases, other organs are involved in bacillary angiomatosis. Patients usually have constitutional symptoms, including fever, chills, malaise, headache, anorexia, weight loss, and night sweats. In osseous disease, lytic lesions are generally seen on radiography, and technetium scan shows focal uptake. The differential diagnosis of cutaneous bacillary angiomatosis includes Kaposi's sarcoma, pyogenic granuloma, subcutaneous tumors, and verruga peruana. In bacillary peliosis, hypodense hepatic areas are usually evident on imaging. In patients with advanced immunodeficiency, *B. henselae* and *B. quintana* are important causes of fever of unknown origin. Intermittent bacteremia with positive blood cultures can occur with or without endocarditis.

PATHOLOGY

Bacillary angiomatosis consists of lobular proliferations of small blood vessels lined by enlarged endothelial cells interspersed with mixed infiltrates of neutrophils and lymphocytes, with predominance of the former. Histologic examination of organs with bacillary peliosis reveals small blood-filled cystic lesions partially lined by endothelial

198e

Donovanosis

Nigel O'Farrell

Donovanosis is a chronic, progressive bacterial infection that usually involves the genital region. The condition is generally regarded as a sexually transmitted infection of low infectivity. This infection has been known by many other names, the most common being *granuloma inguinale*.

ETIOLOGY

The causative organism has been reclassified as *Klebsiella granulomatis* *comb nov* on the basis of phylogenetic analysis, although there is ongoing debate about this decision. Some authorities consider the original nomenclature (*Calymmatobacterium granulomatis*) to be more appropriate in light of analysis of 16S rRNA gene sequences.

Donovanosis was first described in Calcutta in 1882, and the causative organism was recognized by Charles Donovan in Madras in 1905. He identified the characteristic Donovan bodies, measuring $1.5 \times 0.7 \mu\text{m}$, in macrophages and the stratum malpighii. The organism was not reproducibly cultured until the mid-1990s, when its isolation in peripheral-blood monocytes and human epithelial cell lines was reported.

EPIDEMIOLOGY



Donovanosis has an unusual geographic distribution that includes Papua New Guinea, parts of southern Africa, India, the Caribbean, French Guyana, Brazil, and aboriginal communities in Australia. In Australia, donovanosis has been almost entirely eliminated through a sustained program backed by strong political commitment and resources at the primary health care level. Although few cases are now reported in the United States, donovanosis was once prevalent in this country, with 5000–10,000 cases recorded in 1947. The largest epidemic recorded was in Dutch South Guinea, where 10,000 cases were identified in a population of 15,000 (the Marind-anim people) between 1922 and 1952.

Donovanosis is associated with poor hygiene and is more common in lower socioeconomic groups than in those who are better off and in men than in women. Infection in sexual partners of index cases occurs to a limited extent. Donovanosis is a risk factor for HIV infection (Chap. 226).

Globally, the incidence of donovanosis has decreased significantly in recent times. This decline probably reflects a greater focus on effective management of genital ulcers because of their role in facilitating HIV transmission.

CLINICAL FEATURES

A lesion starts as a papule or subcutaneous nodule that later ulcerates after trauma. The incubation period is uncertain, but experimental infections in humans indicate a duration of ~50 days. Four types of lesions have been described: (1) the classic ulcerogranulomatous lesion (Fig. 198e-1), a beefy red ulcer that bleeds readily when touched; (2) a hypertrophic or verrucous ulcer with a raised irregular edge; (3) a necrotic, offensive-smelling ulcer causing tissue destruction; and (4) a sclerotic or cicatricial lesion with fibrous and scar tissue.

The genitals are affected in 90% of patients and the inguinal region in 10%. The most common sites of infection are the prepuce, coronal sulcus, frenum, and glans in men and the labia minora and fourchette in women. Cervical lesions may mimic cervical carcinoma. In men, lesions are associated with lack of circumcision. Lymphadenitis is uncommon. Extragenital lesions occur in 6% of cases and may involve the lip, gums, cheek, palate, pharynx, larynx, and chest. Hematogenous spread with involvement of liver and bone has been reported. During pregnancy, lesions tend to develop more quickly and respond more slowly to treatment. Polyarthrititis and osteomyelitis are rare complications. In newborn infants, donovanosis may present with ear infection. Cases in children have been attributed to sitting on the laps of infected adults.



FIGURE 198e-1 Ulcerogranulomatous penile lesion of donovanosis, with some hypertrophic features.

As the incidence of donovanosis has decreased, the number of unusual case reports has appeared to be increasing.

Complications include neoplastic changes, pseudo-elephantiasis, and stenosis of the urethra, vagina, or anus.

DIAGNOSIS

A clinical diagnosis of donovanosis made by an experienced practitioner on the basis of the lesion's appearance usually has a high positive predictive value. The diagnosis is confirmed by microscopic identification of Donovan bodies (Fig. 198e-2) in tissue smears. Preparation of a good-quality smear is important. If donovanosis is suspected on clinical grounds, the smear for Donovan bodies should be taken before swab samples are collected to be tested for other causes of genital ulceration so that enough material can be collected from the ulcer. A swab should be rolled firmly over an ulcer previously cleaned with a dry swab to remove debris. Smears can be examined in a clinical setting by direct microscopy with a rapid Giemsa or Wright's stain. Alternatively, a piece of granulation tissue crushed and spread between two slides can be used. Donovan bodies can be seen in large, mononuclear (Pund) cells as gram-negative intracytoplasmic cysts filled with deeply staining bodies that may have a safety-pin appearance. These cysts eventually rupture and release the infective organisms. Histologic changes

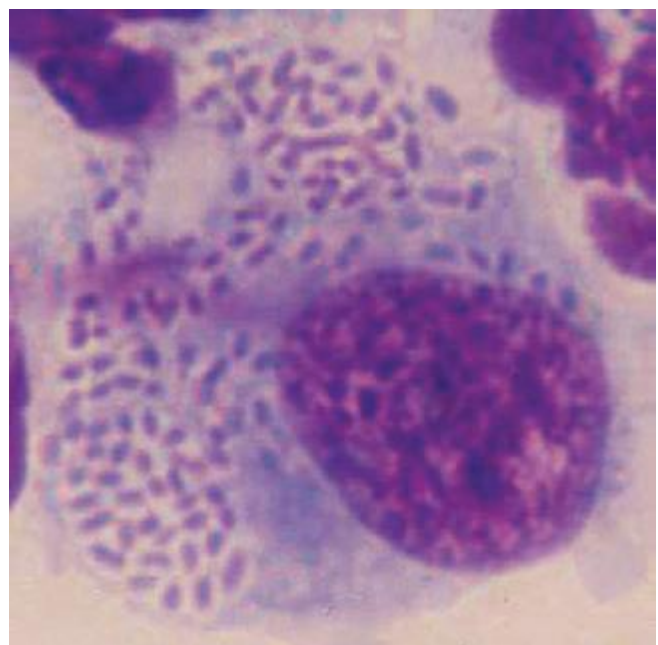


FIGURE 198e-2 Pund cell stained by rapid Giemsa (RapiDiff) technique. Numerous Donovan bodies are visible.

198e-2 include chronic inflammation with infiltration of plasma cells and neutrophils. Epithelial changes include ulceration, microabscesses, and elongation of rete ridges.

A diagnostic polymerase chain reaction (PCR) test was based on the observation that two unique base changes in the *phoE* gene eliminate HaeIII restriction sites, enabling differentiation of *K. granulomatis comb nov* from related *Klebsiella* species. PCR analysis with a colorimetric detection system can now be used in routine diagnostic laboratories. A genital ulcer multiplex PCR that includes *K. granulomatis* has been developed. Serologic tests are only poorly specific and are not currently used.

The differential diagnosis of donovanosis includes primary syphilitic chancres, secondary syphilis (condylomata lata), chancroid, lymphogranuloma venereum, genital herpes, neoplasm, and amebiasis. Mixed infections are common. Histologic appearances should be distinguished from those of rhinoscleroma, leishmaniasis, and histoplasmosis.

TREATMENT DONOVANOSIS

Many patients with donovanosis present quite late with extensive ulceration. They may be embarrassed and have low self-esteem related to their disease. Reassurance that they have a treatable condition is important, as is the need to administer antibiotics and monitor patients for an adequate interval (see below). Epidemiologic treatment of sexual partners and advice about how to improve genital hygiene are recommended.

TABLE 198e-1 EFFECTIVE ANTIBIOTICS FOR THE TREATMENT OF DONOVANOSIS

Antibiotic	Oral Dose
Azithromycin	1 g on day 1, then 500 mg daily for 7 days or 1 g weekly for 4 weeks
Trimethoprim-sulfamethoxazole	960 mg bid for 14 days
Doxycycline	100 mg bid for 14 days
Erythromycin	500 mg qid for 14 days (in pregnant women)
Tetracycline	500 mg qid for 14 days

The recommended drug regimens for donovanosis are shown in **Table 198e-1**. Gentamicin can be added if the response is slow. Ceftriaxone, chloramphenicol, and norfloxacin are also effective. Patients treated for 14 days should be monitored until lesions have healed completely. Those treated with azithromycin probably do not need such rigorous follow-up.

Surgery may be indicated for very advanced lesions.

CONTROL AND PREVENTION

Donovanosis is probably the cause of genital ulceration that is most readily recognizable clinically. Donovanosis is now limited to a few specific locations, and its global eradication is a distinct possibility.



FIGURE 197-2 Nodular lesion of bacillary angiomatosis with superficial ulceration in an AIDS patient with advanced immunodeficiency. (Reprinted with permission from DH Spach and E Darby: *Bartonella Infections, Including Cat-Scratch Disease*, in *Harrison's Principles of Internal Medicine*, 17th ed, AF Fauci et al [eds]. New York, McGraw-Hill, 2008, p 989.)

cells that can be several millimeters in size. Peliotic lesions are surrounded by fibromyxoid stroma containing inflammatory cells, dilated capillaries, and clumps of granular material. Warthin-Starry silver staining of bacillary angiomatosis and peliosis lesions reveals clusters of bacilli. Cultures are usually negative.

DIAGNOSIS

Bacillary angiomatosis and bacillary peliosis are diagnosed on histologic grounds. Blood cultures may be positive.

TREATMENT BACILLARY ANGIOMATOSIS AND PELIOSIS

(Table 197-2) Prolonged therapy with a macrolide or doxycycline is recommended for both bacillary angiomatosis and bacillary peliosis.

PREVENTION

Control of cat-flea infestation and avoidance of cat scratches (for prevention of *B. henselae*) and avoidance and treatment of body louse infestation (for prevention of *B. quintana*) are reasonable strategies for HIV-infected persons. Primary prophylaxis is not recommended, but suppressive therapy with a macrolide or doxycycline is indicated in HIV-infected patients with bacillary angiomatosis or bacillary peliosis until CD4+ T cell counts are >200/ μ L. Relapse may necessitate lifelong suppressive therapy in individual cases.

CARRIÓN'S DISEASE (OROYA FEVER AND VERRUGA PERUANA)

DEFINITION AND ETIOLOGY

Carrión's disease is a biphasic disease caused by *B. bacilliformis*. *Oroya fever* is the initial, bacteremic, systemic form, and *verruca peruana* is its late-onset, eruptive manifestation.

EPIDEMIOLOGY AND PREVENTION



Infection is endemic to the geographically restricted Andes valleys of Peru, Ecuador, and Colombia (~500–3200 m above sea level). Sporadic epidemics occur. The disease is transmitted by the phlebotomine sandfly *Lutzomyia verrucarum*. Humans are the only known reservoir of *B. bacilliformis*. Sandfly control measures (e.g., insecticides) and personal protection measures (e.g., repellents, screening, bed nets) may decrease the risk of infection.

PATHOGENESIS

After inoculation by the sandfly, bacteria invade the blood vessel endothelium and proliferate; the reticuloendothelial system and various organs

may also be involved. Upon reentry into blood vessels, *B. bacilliformis* invades, replicates, and ultimately destroys erythrocytes, with consequent massive hemolysis and sudden, severe anemia. Microvascular thrombosis results in end-organ ischemia. Survivors sometimes develop cutaneous hemangiomas characterized by various inflammatory cells, endothelial proliferation, and the presence of *B. bacilliformis*.

CLINICAL MANIFESTATIONS

The incubation period is 3 weeks (range, 2–14 weeks). Oroya fever may present as a nonspecific bacteremic febrile illness without anemia or as an acute, severe hemolytic anemia with hepatomegaly and jaundice of rapid onset leading to vascular collapse and clouded sensorium. Myalgia, arthralgia, lymphadenopathy, and abdominal pain may develop. Temperature is elevated but not extremely so; high fever may suggest intercurrent infection. Subclinical asymptomatic infection also occurs. In *verruca peruana*, red, hemangioma-like, cutaneous vascular lesions of various sizes appear either weeks to months after systemic illness or with no previous suggestive history. These lesions persist for months up to 1 year. Mucosal and internal lesions may also develop.

DIAGNOSIS AND APPROACH TO THE PATIENT

Systemic illness (with or without anemia) or the development of cutaneous lesions in a person who has been to an endemic area raises the possibility of *B. bacilliformis* infection. Severe anemia with exuberant reticulocytosis—and sometimes thrombocytopenia—can occur. In systemic illness, Giemsa-stained blood films show typical intraerythrocytic bacilli, and blood and bone marrow cultures are positive. Serologic assays may be helpful. Biopsy may be required to confirm the diagnosis of *verruca peruana*. Differential diagnosis includes the spectrum of coendemic systemic febrile illnesses (e.g., typhoid fever, malaria, brucellosis) as well as diseases producing cutaneous vascular lesions (e.g., hemangiomas, bacillary angiomatosis, Kaposi's sarcoma).

TREATMENT CARRIÓN'S DISEASE

(Table 197-2) Antibiotic therapy for systemic *B. bacilliformis* infection usually results in rapid defervescence. Additional antibiotic treatment of intercurrent infection (particularly salmonellosis) is often required. Blood transfusion may be necessary. Treatment of *verruca peruana* usually is not required, although large lesions or those interfering with function may require excision. Patients with numerous lesions, especially lesions that have been present for only a short period, may respond well to antibiotic therapy.

COMPLICATIONS AND PROGNOSIS

Mortality rates associated with Oroya fever have been reported to be as high as 40% without treatment but are considerably lower (~10%) with treatment. Complications such as bacterial superinfection and neurologic and cardiac manifestations occur frequently. Generalized massive edema (anasarca) and petechiae are associated with poor outcome. Permanent immunity usually develops.

198e Donovanosis

Nigel O'Farrell

This is a digital-only chapter. It is available on the DVD that accompanies this book, as well as on Access Medicine/Harrison's Online, and the eBook and "app" editions of HPIIM 19e.

Donovanosis is a chronic, progressive bacterial infection that usually involves the genital region. The condition is generally regarded as a sexually transmitted infection of low infectivity. This infection has been known by many other names, the most common being *granuloma inguinale*.

199 Nocardiosis

Gregory A. Filice

Nocardia, a genus of saprophytic aerobic actinomycetes that are common worldwide, resides in soil, contributing to the decay of organic matter. Nearly 100 species have been identified, mostly on the basis of 16S rRNA gene sequences. Nocardiae are relatively inactive in standard biochemical tests, and speciation is difficult without molecular phylogenetic techniques. Historically, the majority of isolates associated with pneumonia and systemic disease were identified as *Nocardia asteroides*, but the lineage of the type strain was muddled, and it is now clear that human disease is associated with several species. Most clinical laboratories cannot speciate isolates accurately and may identify them simply as *N. asteroides* or *Nocardia* species.

Nine species or species complexes are commonly associated with human disease (Table 199-1). Most systemic disease involves *Nocardia cyriacigeorgica*, *Nocardia farcinica*, *Nocardia pseudobrasiliensis*, and species in the *Nocardia transvalensis* and *Nocardia nova* complexes.

TABLE 199-1 *NOCARDIA* SPECIES MOST COMMONLY ASSOCIATED WITH HUMAN DISEASE AND THEIR IN VITRO SUSCEPTIBILITY PATTERNS

Species	Susceptible to	Resistant to
<i>N. abscessus</i>	Amikacin, amoxicillin/clavulanic acid, ampicillin, ceftriaxone, gentamicin, linezolid, minocycline, TMP-SMX	Ciprofloxacin, clarithromycin, erythromycin, imipenem (v) ^a
<i>N. brevicatena/paucivorans</i> complex (<i>N. brevicatena</i> , <i>N. paucivorans</i> , <i>N. carnea</i> , others)	Amikacin, amoxicillin/clavulanic acid, ampicillin, ceftriaxone, ciprofloxacin, linezolid, minocycline (v), moxifloxacin, tobramycin, TMP-SMX	Ciprofloxacin, clarithromycin, erythromycin, gentamicin, imipenem (v)
<i>N. nova</i> complex (<i>N. nova</i> , <i>N. veterana</i> , <i>N. africana</i> , <i>N. kruczakiae</i> , <i>N. elegans</i> , others)	Amikacin, ampicillin, ceftriaxone, clarithromycin, erythromycin, imipenem, linezolid, minocycline, TMP-SMX	Amoxicillin/clavulanic acid, ciprofloxacin, gentamicin, tobramycin
<i>N. transvalensis</i> complex (<i>N. blacklockiae</i> , <i>N. wallacei</i> , others)	Amoxicillin/clavulanic acid (v), ceftriaxone (v), ciprofloxacin, linezolid, minocycline (v), TMP-SMX	Amikacin, ampicillin, clarithromycin, erythromycin, gentamicin, imipenem (v)
<i>N. farcinica</i>	Amikacin, amoxicillin/clavulanic acid, imipenem (v), linezolid, minocycline (v), TMP-SMX	Ampicillin, ceftriaxone, ciprofloxacin, clarithromycin, erythromycin, gentamicin, tobramycin
<i>N. cyriacigeorgica</i>	Amikacin, ceftriaxone (v), imipenem, linezolid, minocycline (v), TMP-SMX	Amoxicillin/clavulanic acid, ampicillin (v), ciprofloxacin, erythromycin, gentamicin
<i>N. brasiliensis</i>	Amikacin, amoxicillin/clavulanic acid, minocycline, moxifloxacin, TMP-SMX	Ampicillin, ceftriaxone, ciprofloxacin, clarithromycin, imipenem
<i>N. pseudobrasiliensis</i>	Amikacin, ceftriaxone (v), ciprofloxacin, clarithromycin, TMP-SMX	Amoxicillin/clavulanic acid, ampicillin, imipenem, minocycline
<i>N. otitidiscaviarum</i> complex	Amikacin, ciprofloxacin, gentamicin, TMP-SMX	Amoxicillin/clavulanic acid, ampicillin, ceftriaxone, imipenem

^aVariable.

Abbreviation: TMP-SMX, trimethoprim-sulfamethoxazole.

Source: Adapted from multiple sources.

Nocardia brasiliensis is usually associated with disease limited to the skin. Actinomycetoma—an indolent, slowly progressive disease of skin and underlying tissues with nodular swellings and draining sinuses—is often associated with *N. brasiliensis*, *Nocardia otitidiscaviarum*, *N. transvalensis* complex strains, or other actinomycetes.

EPIDEMIOLOGY



Nocardiosis occurs worldwide. The annual incidence, estimated on three continents (North America, Europe, and Australia), is ~0.375 cases per 100,000 persons and may be increasing. The disease is more common among adults than among children and more common among males than among females. Nearly all cases are sporadic, but outbreaks have been associated with contamination of the hospital environment, cosmetic procedures, and parenteral illicit drug use. Person-to-person spread is not well documented. There is no known seasonality.

The majority of cases of pulmonary or disseminated disease occur in people with a host defense defect. Most have deficient cell-mediated immunity, especially that associated with lymphoma, transplantation, glucocorticoid therapy, or AIDS. The incidence is ~140-fold greater among patients with AIDS and ~340-fold greater among bone marrow transplant recipients than in general populations. In AIDS, nocardiosis usually affects persons with <250 CD4+ T lymphocytes/μL. Nocardiosis has also been associated with pulmonary alveolar proteinosis, tuberculosis and other mycobacterial diseases, chronic granulomatous disease, interleukin 12 deficiency, and treatment with monoclonal antibodies that interfere with tumor necrosis factor. Any child with nocardiosis and no known cause of immunosuppression should undergo tests to determine the adequacy of the phagocytic respiratory burst.

Cases of actinomycetoma occur mainly in tropical and subtropical regions, especially those of Mexico, Central and South America, Africa, and India. The most important risk factor is frequent contact with soil or vegetable matter, especially in laborers.

PATHOLOGY AND PATHOGENESIS

Pneumonia and disseminated disease are both thought to follow inhalation of fragmented bacterial mycelia. The characteristic histologic feature of nocardiosis is an abscess with extensive neutrophil infiltration and prominent necrosis. Granulation tissue usually surrounds the lesions, but extensive fibrosis or encapsulation is uncommon.

Actinomycetoma is characterized by suppurative inflammation with sinus tract formation. Granules—microcolonies composed of dense masses of bacterial filaments extending radially from a central core—are occasionally observed in histologic preparations. The granules are frequently found in discharges from lesions of actinomycetoma but almost never in discharges from lesions in other forms of nocardiosis.

Nocardiae have evolved a number of properties that enable them to survive within phagocytes, including neutralization of oxidants, prevention of phagosome-lysosome fusion, and prevention of phagosome acidification. Neutrophils phagocytose the organisms and limit their growth but do not kill them efficiently. Cell-mediated immunity is important for definitive control and elimination of nocardiae.

CLINICAL MANIFESTATIONS

Respiratory Tract Disease Pneumonia, the most common form of nocardial disease in the respiratory tract, is typically subacute; symptoms have usually been present for days or weeks at presentation. The onset is occasionally more acute in immunosuppressed patients. Cough is prominent and produces small amounts of thick, purulent sputum that is not malodorous. Fever, anorexia, weight loss, and malaise are common; dyspnea, pleuritic pain, and hemoptysis are less common. Remissions and exacerbations over several weeks are frequent. Roentgenographic patterns vary, but some are highly suggestive of



FIGURE 199-1 Nocardial pneumonia. A dense infiltrate with a possible cavity and several nodules are apparent in the right lung.

nocardial pneumonia. Infiltrates vary in size and are typically dense. Single or multiple nodules are common (Figs. 199-1 and 199-2), sometimes suggesting tumors or metastases. Infiltrates and nodules tend to cavitate (Fig. 199-2). Empyema is present in one-quarter of cases. Co-infection with *Nocardia* and *Mycobacterium tuberculosis* has been reported from regions where tuberculosis is common.

Nocardiosis may spread directly from the lungs to adjacent tissues. Pericarditis, mediastinitis, and the superior vena cava syndrome have all been reported. Nocardial laryngitis, tracheitis, bronchitis, and sinusitis are much less common than pneumonia. In the major airways, disease often presents as a nodular or granulomatous mass. Nocardiae are sometimes isolated from respiratory secretions of persons without apparent nocardial disease, usually individuals who have underlying lung or airway abnormalities.



FIGURE 199-2 Nocardial pneumonia. A computed tomography scan shows bilateral nodules, with cavitation in the nodule in the left lung.

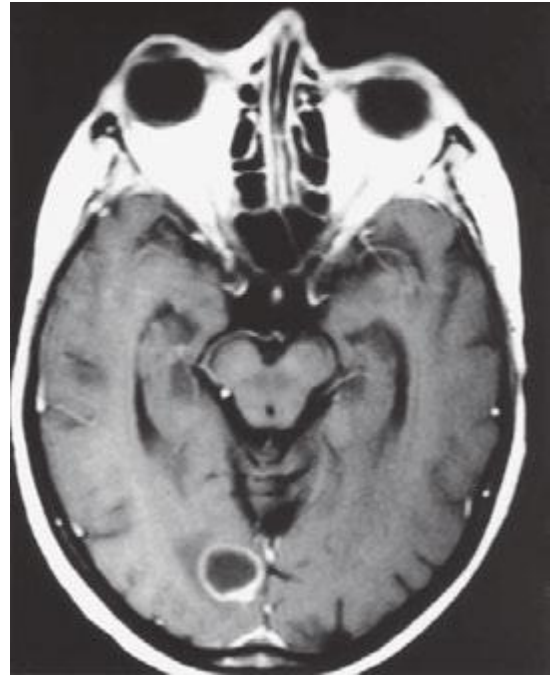


FIGURE 199-3 Nocardial abscesses in the right occipital lobe.

Extrapulmonary Disease In half of all cases of pulmonary nocardiosis, disease appears outside the lungs. In one-fifth of cases of disseminated disease, lung disease is not apparent. The most common site of dissemination is the brain. Other common sites include the skin and supporting structures, kidneys, bone, muscle, and eye, but almost any organ can be involved. Peritonitis has been reported in patients undergoing peritoneal dialysis. Nocardiae have been recovered from blood in a few cases of pneumonia, disseminated disease, or central venous catheter infection. Nocardial endocarditis occurs rarely and can affect either native or prosthetic valves.

The typical manifestation of extrapulmonary dissemination is a subacute abscess. A minority of abscesses outside the lungs or central nervous system (CNS) form fistulas and discharge small amounts of pus. In CNS infections, brain abscesses are usually supratentorial, are often multiloculated, and may be single or multiple (Fig. 199-3). Brain abscesses tend to burrow into the ventricles or extend out into the subarachnoid space. The symptoms and signs are somewhat more indolent than those of other types of bacterial brain abscess. Meningitis is uncommon and is usually due to spread from a nearby brain abscess. Nocardiae are not easily recovered from cerebrospinal fluid (CSF).

Disease Following Transcutaneous Inoculation Disease that follows transcutaneous nocardial inoculation usually takes one of three forms: cellulitis, lymphocutaneous syndrome, or actinomycetoma.

Cellulitis generally begins 1–3 weeks after a recognized breach of the skin, often with soil contamination. Subacute cellulitis, with pain, swelling, erythema, and warmth, develops over days to weeks. The lesions are usually firm and not fluctuant. Disease may progress to involve underlying muscles, tendons, bones, or joints. Dissemination is rare. *N. brasiliensis* and species in the *N. otitidiscaviarum* complex are most common in cellulitis cases.

Lymphocutaneous disease usually begins as a pyodermatous nodule at the site of inoculation, with central ulceration and purulent or honey-colored drainage. Subcutaneous nodules often appear along lymphatics that drain the primary lesion. Most cases of nocardial lymphocutaneous syndrome are associated with *N. brasiliensis*. Similar disease occurs with other pathogens, most notably *Sporothrix schenckii* (Chap. 243) and *Mycobacterium marinum* (Chap. 204).

Actinomycetoma usually begins with a nodular swelling, sometimes at a site of local trauma. Lesions (Fig. 199-4A) typically develop on the feet or hands but may involve the posterior part of the neck, the upper back, the head, and other sites. The nodule eventually breaks down,

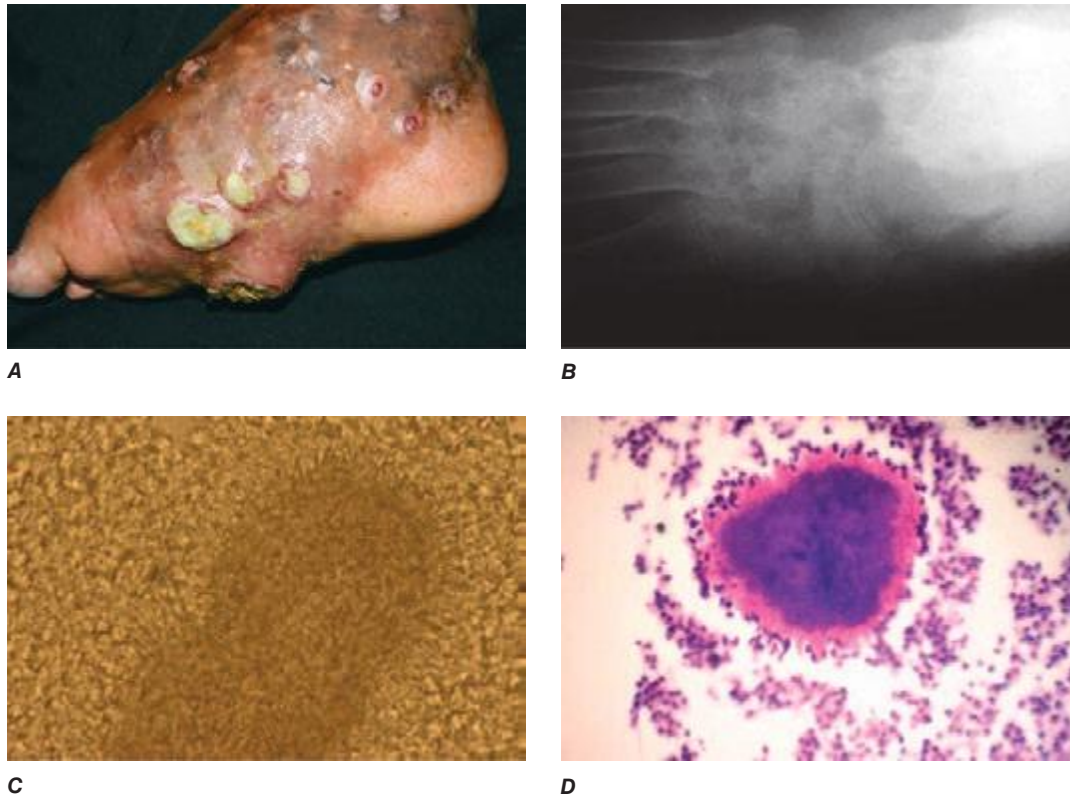


FIGURE 199-4 *Nocardia brasiliensis* mycetoma. **A.** Draining sinuses and giant white grains with a seropurulent discharge. **B.** Radiography of the foot showing marked soft tissue enlargement and bony lytic lesions. **C.** Direct microscopy of grains stained with Lugol's iodine (×40). **D.** Periodic acid–Schiff stain of skin biopsy (×40). (Image provided by Roberto Arenas and Mahreen Ameen, St. John's Institute of Dermatology, Guy's & St Thomas' NHS Trust, London, UK. Reprinted with permission from R Arenas, M Ameen: *Lancet Infect Dis* 10:66, 2010.)

and a fistula appears, typically followed by others. The fistulas tend to come and go, with new ones forming as old ones disappear. The discharge is serous or purulent, may be bloody, and often contains 0.1- to 2-mm white granules consisting of masses of mycelia (Figs. 199-4C and 199-4D). The lesions spread slowly along fascial planes to involve adjacent areas of skin, subcutaneous tissue, and bone. Over months or years, there may be extensive deformation of the affected part. Lesions involving soft tissues are only mildly painful; those affecting bones or joints are more so (Fig. 199-4B). Systemic symptoms are absent or minimal. Infection rarely disseminates from actinomycetoma, and lesions on the hands and feet usually cause only local disability. Lesions on the head, neck, and trunk can invade locally to involve deep organs, with consequent severe disability or death.

Eye Infections *Nocardia* species are uncommon causes of subacute keratitis, usually following eye trauma. Nocardial endophthalmitis can develop after eye surgery. In one series, nocardiae accounted for more than half of culture-proved cases of endophthalmitis after cataract surgery. Endophthalmitis can also occur during disseminated disease. Nocardial infection of lachrymal glands has been reported.

DIAGNOSIS

The first step in diagnosis is examination of sputum or pus for crooked, branching, beaded, gram-positive filaments 1 μm wide and up to 50 μm long (Fig. 199-5). Most nocardiae are acid-fast in direct smears if a weak acid is used for decolorization (e.g., in the modified Kinyoun, Ziehl-Neelsen, and Fite-Faraco methods). The organisms often take up silver stains. Recovery from specimens containing a mixed flora can be improved with selective media (colistin–nalidixic acid agar, modified Thayer-Martin agar, or buffered charcoal–yeast extract agar). Nocardiae grow well on most fungal and mycobacterial media, but procedures used for decontamination of specimens for mycobacterial culture can kill nocardiae and thus should not be used when nocardiae are suspected. Nocardiae grow relatively slowly; colonies may take up to 2 weeks to appear and may not develop their

characteristic appearance—white, yellow, or orange, with aerial mycelia and delicate, dichotomously branched substrate mycelia—for up to 4 weeks. Several blood culture systems support nocardial growth, although nocardiae may not be detected for up to 2 weeks. The growth of nocardiae is so different from that of more common pathogens that the laboratory should be alerted when nocardiosis is suspected in order to maximize the likelihood of isolation.

In nocardial pneumonia, sputum smears are often negative. Unless the diagnosis can be made in smear-negative cases by sampling lesions in more accessible sites, bronchoscopy or lung aspiration is usually necessary. To evaluate the possibility of dissemination in patients

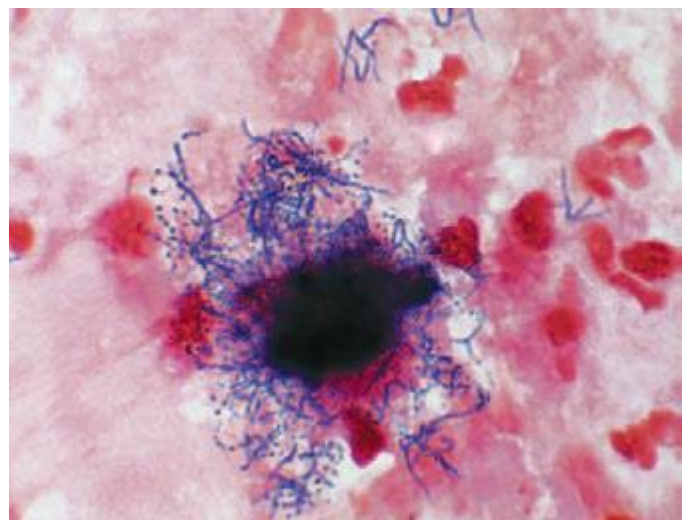


FIGURE 199-5 Gram-stained sputum from a patient with nocardial pneumonia. (Image provided by Charles Cartwright and Susan Nelson, Hennepin County Medical Center, Minneapolis, MN.)

with nocardial pneumonia, a careful history should be obtained and a thorough physical examination performed. Suggestive symptoms or signs should be pursued with further diagnostic tests. Computed tomography (CT) or magnetic resonance imaging (MRI) of the head, with and without contrast material, should be undertaken if signs or symptoms suggest brain involvement. Some authorities recommend brain imaging in all cases of pulmonary or disseminated disease. When clinically indicated, CSF or urine should be concentrated and then cultured. Actinomycetoma, eumycetoma (cases involving fungi; [Chap. 243](#)), and botryomycosis (cases involving cocci or bacilli, often *Staphylococcus aureus*) are difficult to distinguish clinically but are readily distinguished with microbiologic testing or biopsy. Granules should be sought in any discharge. Suspect particles should be washed in saline, examined microscopically, and cultured. Granules in actinomycetoma cases are usually white, pale yellow, pink, or red. Viewed microscopically, they consist of tight masses of fine filaments (0.5–1 μm wide) radiating outward from a central core (Fig. 199-5). Granules from eumycetoma cases are white, yellow, brown, black, or green. Under the microscope, they appear as masses of broader filaments (2–5 μm wide) encased in a matrix. Granules of botryomycosis consist of loose masses of cocci or bacilli. Organisms can also be seen in wound discharge or histologic specimens. The most reliable way to differentiate among the various organisms associated with mycetoma is by culture.

Isolation of nocardiae from sputum or blood occasionally represents colonization, transient infection, or contamination. In typical cases of respiratory tract colonization, Gram-stained specimens are negative and cultures are only intermittently positive. A positive sputum culture in an immunosuppressed patient usually reflects disease. When nocardiae are isolated from sputum of an immunocompetent patient without apparent nocardial disease, the patient should be observed carefully without treatment. A patient with a host-defense defect that increases the risk of nocardiosis should usually receive antimicrobial treatment.

TREATMENT NOCARDIOSIS

For mild or moderate cases, therapy with drugs known to be effective against most isolates is usually adequate. For severe cases or cases that do not respond promptly to antimicrobial therapy, isolates should be sent to a laboratory experienced with *Nocardia* for identification and susceptibility testing. Identification of an isolate to the species level is accomplished with molecular testing, and susceptibility is assessed with a Clinical Laboratory Standards Institute (CLSI)-approved broth dilution test. Nocardial growth is slower than the growth of most clinically important bacteria, and nocardiae tend to clump in suspension so that susceptibility test endpoints are unusual; thus experience is necessary for reliable results. Because nocardiosis is uncommon, data on the relation between susceptibility test results for specific drugs and clinical outcomes in patients treated with these drugs are meager. Careful clinical monitoring is essential, and consultation with clinicians who have experience with nocardiosis is often needed.

Sulfonamides are the drugs of choice ([Tables 199-1 and 199-2](#)). The combination of sulfamethoxazole (SMX) and trimethoprim (TMP) is at least equivalent to a sulfonamide alone and may be slightly more effective, but the combination also poses a modestly greater risk of hematologic toxicity. At the outset, 10–20 mg/kg of TMP and 50–100 mg/kg of SMX are given each day in two divided doses. Later, daily doses can be decreased to as little as 5 mg/kg and 25 mg/kg, respectively. In persons with sulfonamide allergies, desensitization usually allows continuation of therapy with these effective and inexpensive drugs.

Sulfonamide susceptibility testing is difficult. The CLSI standard methodology includes a technique for TMP-SMX but not for a sulfonamide alone. Reported rates of sulfonamide susceptibility have varied widely, and controversy has ensued about the reliability of sulfonamides for therapy. However, clinical responses to

TABLE 199-2 TREATMENT DURATION FOR NOCARDIOSIS

Disease	Duration
Pulmonary or systemic	
Intact host defenses	6–12 months
Deficient host defenses	12 months ^a
CNS disease	12 months ^b
Cellulitis, lymphocutaneous syndrome	2 months
Osteomyelitis, arthritis, laryngitis, sinusitis	4 months
Actinomycetoma	6–12 months after clinical cure
Keratitis	Topical: until apparent cure Systemic: until 2–4 months after apparent cure

^aIn some patients with AIDS and CD4+ T lymphocyte counts of <200/ μL or with chronic granulomatous disease, therapy for pulmonary or systemic disease must be continued indefinitely. ^bIf all apparent CNS disease has been excised, the duration of therapy may be reduced to 6 months.

appropriate sulfonamide treatment are nearly always satisfactory. Sulfonamides remain the drugs of choice in nearly all cases.

Clinical experience with other oral drugs is limited. Minocycline (100–200 mg twice a day) is often effective; other tetracyclines are usually less effective. Linezolid is active against all species in vitro and in vivo, but adverse effects are common with long-term use. Tigecycline appears to be active in vitro against some species, but little clinical experience has been reported. Amoxicillin (875 mg) combined with clavulanic acid (125 mg), given twice a day, has been effective but should be avoided in cases involving strains of the *N. nova* complex, in which clavulanate induces β -lactamase production. Among the quinolones, moxifloxacin and gemifloxacin appear to be most active.

Amikacin, the best-established parenteral drug except in cases involving the *N. transvalensis* complex, is given in doses of 5–7.5 mg/kg every 12 h or 15 mg/kg every 24 h. Serum drug levels should be monitored during prolonged therapy in patients with diminished renal function and in the elderly. Ceftriaxone and imipenem are usually effective except as indicated in Table 199-1.

Patients with severe disease are initially treated with a combination including TMP-SMX, amikacin, and ceftriaxone or imipenem. Clinical improvement is usually noticeable after 1–2 weeks of therapy but may take longer, especially with CNS disease. After definite clinical improvement, therapy can be continued with a single oral drug, usually TMP-SMX. Some experts use two or more drugs for the entire course of therapy, but whether multiple drugs are better than a single agent is not known, and additional drugs increase the risk of toxicity. In patients with nocardiosis who need immunosuppressive therapy for an underlying disease or prevention of transplant rejection, immunosuppressive therapy should be continued.

Use of SMX and TMP in high-risk populations to prevent *Pneumocystis* disease or urinary tract infections appears to reduce but not eliminate the risk of nocardiosis. The incidence of nocardiosis is low enough that prophylaxis solely to prevent this disease is not recommended.

Surgical management of nocardial disease is similar to that of other bacterial diseases. Brain abscesses should be aspirated, drained, or excised if the diagnosis is unclear, if an abscess is large and accessible, or if an abscess fails to respond to chemotherapy. Small or inaccessible brain abscesses should be treated medically; clinical improvement should be noticeable within 1–2 weeks. Brain imaging should be repeated to document the resolution of lesions, although abatement on images often lags behind clinical improvement.

Antimicrobial therapy usually suffices for nocardial actinomycetoma. In deep or extensive cases, drainage or excision of heavily involved tissue may facilitate healing, but structure and function should be preserved whenever possible. Keratitis is treated with topical sulfonamide or amikacin drops plus a sulfonamide or an alternative drug given by mouth.

1088 Nocardial infections tend to relapse (particularly in patients with chronic granulomatous disease), and long courses of antimicrobial therapy are necessary (Table 199-2). If disease is unusually extensive or if the response to therapy is slow, the recommendations in Table 199-2 should be exceeded.

With appropriate treatment, the mortality rate for pulmonary or disseminated nocardiosis outside the CNS should be <5%. CNS disease carries a higher mortality rate. Patients should be followed carefully for at least 6 months after therapy has ended.

200 Actinomycosis and Whipple's Disease

Thomas A. Russo

Actinomycosis and Whipple's disease share characteristics that confound even the skilled diagnostician. Because both diseases are uncommon, the physician's personal experience with their clinical presentations is limited. The laboratory identification of the etiologic agents from the order Actinomycetales is not routine. Thus they remain a diagnostic challenge. However, both of these chronic infections are curable, usually with medical therapy alone. Therefore, an awareness of the full spectrum of these diseases, prompting clinical suspicion, can expedite their diagnosis and treatment and minimize unnecessary surgical interventions (especially with actinomycosis), morbidity, and mortality risk.

ACTINOMYCOSIS

Actinomycosis is an indolent, slowly progressive infection caused by anaerobic or microaerophilic bacteria, primarily of the genus *Actinomyces*, that colonize the mouth, colon, and vagina. Mucosal disruption may lead to infection at virtually any site in the body. In vivo growth of actinomycetes usually results in the formation of characteristic clumps called *grains* or *sulfur granules*. The clinical presentations of actinomycosis are myriad. Common in the preantibiotic era, actinomycosis has diminished in incidence, as has its timely recognition. Actinomycosis has been called the most misdiagnosed disease, and it has been said that no disease is so often missed by experienced clinicians.

Three "classic" clinical presentations that should prompt consideration of this unique infection are (1) the combination of chronicity, progression across tissue boundaries, and mass-like features (mimicking malignancy, with which it is often confused); (2) the development of a sinus tract, which may spontaneously resolve and recur; and (3) a refractory or relapsing infection after a short course of therapy, since cure of established actinomycosis requires prolonged treatment.

ETIOLOGIC AGENTS

Actinomycosis is most commonly caused by *A. israelii*, *A. naeslundii*, *A. odontolyticus*, *A. viscosus*, *A. meyeri*, and *A. gerencseriae*. Most if not all actinomycotic infections are polymicrobial. *Aggregatibacter (Actinobacillus) actinomycetemcomitans*, *Eikenella corrodens*, Enterobacteriaceae, and species of *Fusobacterium*, *Bacteroides*, *Capnocytophaga*, *Staphylococcus*, and *Streptococcus* are commonly isolated with actinomycetes in various combinations, depending on the site of infection. Their contribution to the pathogenesis of actinomycosis is uncertain.

Comparative 16S rRNA gene sequencing has led to the identification of an ever-expanding list of *Actinomyces* species and a reclassification of some species to other genera. At present, 46 species and 2 subspecies have been recognized (www.bacterio.cict.fr/a/actinomyces.html). *A. europaeus*, *A. neuii*, *A. radingae*, *A. graevenitzi*, *A. turicensis*, *A. cardiffensis*, *A. houstonensis*, *A. hongkongensis*, *A. lingnae*, *A. masiliensis*, *A. timonensis*, and *A. funkei* as well as two former *Actinomyces*

species—*Arcanobacterium pyogenes* and *Arcanobacterium bernardiae*—are additional causes of human actinomycosis, albeit not always with a "classic" presentation.

EPIDEMIOLOGY

Actinomycosis has no geographic boundaries and occurs throughout life, with a peak incidence in the middle decades. Males have a threefold higher incidence than females, possibly because of poorer dental hygiene and/or more frequent trauma. Improved dental hygiene and the initiation of antimicrobial treatment before actinomycosis fully develops have probably contributed to a decrease in incidence since the advent of antibiotics. Individuals who do not seek or have access to health care, those who have an intrauterine contraceptive device (IUCD) in place for a prolonged period (see "Pelvic Disease," below), and those who receive bisphosphonate treatment (see "Oral-Cervicofacial Disease," below) are probably at higher risk.

PATHOGENESIS AND PATHOLOGY

The etiologic agents of actinomycosis are members of the normal oral flora and are often cultured from the bronchi, the gastrointestinal tract, and the female genital tract. The critical step in the development of actinomycosis is disruption of the mucosal barrier. Local infection may ensue. Once established, actinomycosis spreads contiguously in a slow, progressive manner, ignoring tissue planes. Although acute inflammation may initially develop at the infection site, the hallmark of actinomycosis is the characteristic chronic, indolent phase manifested by lesions that usually appear as single or multiple indurations. Central necrosis consisting of neutrophils and sulfur granules develops and is virtually diagnostic. The fibrotic walls of the mass are typically described as "wooden." The responsible bacterial and/or host factors have not been identified. Over time, sinus tracts to the skin, adjacent organs, or bone may develop. In rare instances, distant hematogenous seeding may occur. As mentioned above, these unique features of actinomycosis mimic malignancy, with which it is often confused.

Foreign bodies appear to facilitate infection. This association most frequently involves IUCDs. Reports have described an association of actinomycosis with HIV infection; transplantation; common variable immunodeficiency; chronic granulomatous disease; treatment with infliximab, glucocorticoids, or bisphosphonates; and radio- or chemotherapy. Ulcerative mucosal infections (e.g., by herpes simplex virus or cytomegalovirus) may facilitate disease development.

CLINICAL MANIFESTATIONS

Oral-Cervicofacial Disease Actinomycosis occurs most frequently at an oral, cervical, or facial site, usually as a soft tissue swelling, abscess, or mass lesion that is often mistaken for a neoplasm. The angle of the jaw is generally involved, but a diagnosis of actinomycosis should be considered with any mass lesion or relapsing infection in the head and neck (Chap. 44). Radiation therapy and especially bisphosphonate treatment have been recognized as contributing to an increasing incidence of actinomycotic infection of the mandible and maxilla (Fig. 200-1). Canaliculitis (also commonly due to *Propionibacterium propionicum*), otitis, and sinusitis also can develop. Pain, fever, and leukocytosis are variably reported. Contiguous extension to the cranium, cervical spine, or thorax is a potential sequela.

Thoracic Disease Thoracic actinomycosis, which may be facilitated by foreign material, usually follows an indolent progressive course, with involvement of the pulmonary parenchyma and/or the pleural space. Chest pain, fever, and weight loss are common. A cough, when present, is variably productive. The usual radiographic finding is either a mass lesion or pneumonia. On CT, central areas of low attenuation and ring-like rim enhancement may be seen. Cavitory disease or mediastinal or hilar adenopathy may develop. More than 50% of cases include pleural thickening, effusion, or empyema (Fig. 200-2). Rarely, pulmonary nodules or endobronchial lesions occur. Lesions suggestive of actinomycosis include those that cross fissures or pleura; extend into the mediastinum, contiguous bone, or chest wall; or are associated with a sinus tract. In the absence of these findings, thoracic actinomycosis is usually mistaken for a neoplasm or pneumonia due to more usual causes.

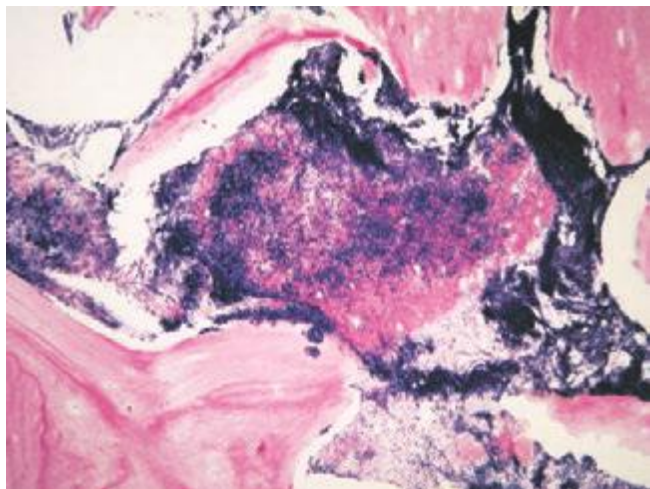


FIGURE 200-1 Bisphosphonate-associated maxillary osteomyelitis due to *A. viscosus*. A sulfur granule is seen within the bone. (Reprinted with permission from NH Naik, TA Russo: Bisphosphonate related osteonecrosis of the jaw: The role of Actinomyces. *Clin Infect Dis* 49:1729, 2009. © 2009 University of Chicago Press.)

Mediastinal infection is uncommon, usually arising from thoracic extension but rarely from perforation of the esophagus, trauma, or extension of head and neck or abdominal disease. The structures within the mediastinum and the heart can be involved in various combinations; consequently, the possible presentations are diverse. Primary endocarditis (in which *A. neuii* has been increasingly described) and isolated disease of the breast occur.

Abdominal Disease Abdominal actinomycosis poses a great diagnostic challenge. Months or years usually pass from the inciting event (e.g., appendicitis, diverticulitis, peptic ulcer disease, spillage of gall stones or bile during laparoscopic cholecystectomy, foreign-body perforation, bowel surgery, or ascension from IUCD-associated pelvic disease) to clinical recognition. Because of the flow of peritoneal fluid and/or the direct extension of primary disease, virtually any abdominal organ, region, or space can be involved. The disease usually presents as an abscess, a mass, or a mixed lesion that is often fixed to underlying tissue and mistaken for a tumor. On CT, enhancement is most often heterogeneous and adjacent bowel is thickened. Sinus tracts to the abdominal wall, to the perianal region, or between the bowel and other organs may develop and mimic inflammatory bowel disease (Chap. 351). Recurrent disease or a wound or fistula that fails to heal suggests actinomycosis.

Hepatic infection usually presents as one or more abscesses or masses (Fig. 200-3). Isolated disease presumably develops via hematogenous seeding from cryptic foci. Imaging and percutaneous techniques have resulted in improved diagnosis and treatment.

All levels of the urogenital tract can be infected. Renal disease usually presents as pyelonephritis and/or renal and perinephric abscess. Bladder involvement, usually due to extension of pelvic disease, may result in ureteral obstruction or fistulas to bowel, skin, or uterus. *Actinomyces* can be detected in urine with appropriate stains and cultures.

Pelvic Disease Actinomycotic involvement of the pelvis occurs most commonly in association with an IUCD. When an IUCD is in place or has recently been removed, pelvic symptoms should prompt consideration of actinomycosis. The risk, although not quantified, appears small. The disease rarely develops when the IUCD has been in place for <1 year, but the risk increases with time. Actinomycosis can also present months after IUCD removal. Symptoms are typically indolent; fever, weight loss, abdominal pain, and abnormal vaginal bleeding or discharge are the most common. The earliest stage of disease—often endometritis—commonly progresses to pelvic masses or a tuboovarian abscess (Fig. 200-4). Unfortunately, because the diagnosis is often delayed, a “frozen pelvis” mimicking malignancy or endometriosis can develop by the time of recognition. Ca125 levels may be elevated, further contributing to misdiagnosis.

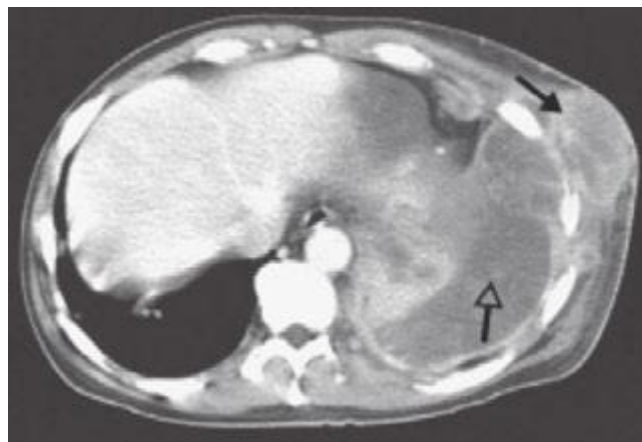
Actinomyces-like organisms (ALOs), which are identified in Papanicolaou-stained specimens in (on average) 7% of women using an IUCD, have a low positive predictive value for diagnosis. Nonetheless, although the risk appears small, the consequences of infection are significant. Therefore, until more quantitative data become available, it seems prudent to remove the IUCD in the presence of symptoms that cannot be accounted for, regardless of whether ALOs are detected, and—if advanced disease is excluded—to initiate a 14-day course of empirical treatment for possible early endometritis. The detection of ALOs in the asymptomatic patient warrants education and close follow-up but not removal of the IUCD unless a suitable contraceptive alternative is agreed on.

Central Nervous System Disease Actinomycosis of the central nervous system (CNS) is rare. Single or multiple brain abscesses are most common. An abscess usually appears on CT as a ring-enhancing lesion with a thick wall that may be irregular or nodular. Magnetic resonance perfusion and spectroscopy findings have also been described, as have primary meningitis, epidural or subdural space infection, and cavernous sinus syndrome.

Musculoskeletal and Soft Tissue Infection Actinomycotic infection of bone and joints is usually due to adjacent soft-tissue infection but may be associated with trauma (e.g., fracture of the mandible), injections,



A



B

FIGURE 200-2 Thoracic actinomycosis. **A.** A chest wall mass from extension of pulmonary infection. **B.** Pulmonary infection is complicated by empyema (open arrow) and extension to the chest wall (closed arrow). (Courtesy of Dr. C. B. Hsiao, Division of Infectious Diseases, Department of Medicine, State University of New York at Buffalo.)

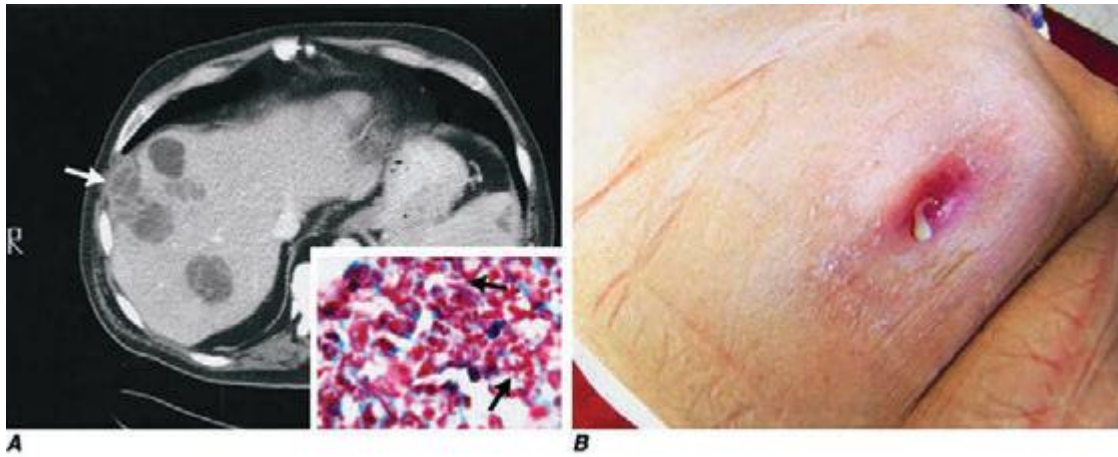


FIGURE 200-3 Hepatic-splenic actinomycosis. **A.** Computed tomogram showing multiple hepatic abscesses and a small splenic lesion due to *A. israelii*. Arrow indicates extension outside the liver. *Inset:* Gram's stain of abscess fluid demonstrating beaded filamentous gram-positive rods. **B.** Subsequent formation of a sinus tract. (Reprinted with permission from Saad M: *Actinomyces hepatic abscess with cutaneous fistula*. *N Engl J Med* 353:e16, 2005. © 2005 Massachusetts Medical Society. All rights reserved.)

surgery, osteoradionecrosis and bisphosphonate osteonecrosis (limited to mandibular and maxillary bones), or hematogenous spread. Because of slow disease progression, new bone formation and bone destruction are seen concomitantly. Infection of an extremity is uncommon and is usually a result of trauma. Skin, subcutaneous tissue, muscle, and bone (with periostitis or acute or chronic osteomyelitis) are involved alone or in various combinations. Cutaneous sinus tracts frequently develop.

Disseminated Disease Hematogenous dissemination of disease from any location rarely results in multiple-organ involvement. *A. meyeri* is most commonly involved. The lungs and liver are most commonly affected, with the presentation of multiple nodules mimicking disseminated malignancy. The clinical presentation may be surprisingly indolent given the extent of disease.

DIAGNOSIS

The diagnosis of actinomycosis is rarely considered. All too often, actinomycosis is first mentioned by the pathologist after extensive surgery. Since medical therapy alone is frequently sufficient for cure, the challenge for the clinician is to consider the possibility of actinomycosis, to diagnose it in the least invasive fashion, and to avoid

unnecessary surgery. The clinical and radiographic presentations that suggest actinomycosis are discussed above. Of note, hypermetabolism has been demonstrated by ^{18}F -fluorodeoxyglucose positron emission tomography (FDG-PET) in actinomycotic disease. Aspirations and biopsies (with or without CT or ultrasound guidance) are being used successfully to obtain clinical material for diagnosis, although surgery may be required. The diagnosis is most commonly made by microscopic identification of sulfur granules (an in vivo matrix of bacteria, calcium phosphate, and host material) in pus or tissues. Occasionally, these granules are identified grossly from draining sinus tracts or pus. Although sulfur granules are a defining characteristic of actinomycosis, granules also are found in mycetoma (Chaps. 199 and 243) and botryomycosis (a chronic suppurative bacterial infection of soft tissue or, in rare cases, visceral tissue that produces clumps of bacteria resembling granules). These entities can easily be differentiated from actinomycosis with appropriate histopathologic and microbiologic studies. Microbiologic identification of actinomycetes is often precluded by prior antimicrobial therapy or failure to perform appropriate microbiologic cultures. For optimal yield, the avoidance of even a single dose of antibiotics is mandatory. Primary isolation usually requires 5–7 days under anaerobic conditions but may take as long as 2–4 weeks. Although not routinely used, 16S rRNA gene amplification and sequencing have been successfully applied to increase diagnostic sensitivity and specificity. Because actinomycetes are components of the normal oral and genital-tract flora, their identification in the absence of sulfur granules in sputum, bronchial washings, and cervicovaginal secretions is of little significance.

TREATMENT ACTINOMYCOSIS

Decisions about treatment are based on the collective clinical experience of the past 65 years. Actinomycosis requires prolonged treatment with high doses of antimicrobial agents; suitable antimicrobial agents and those deemed unreliable are listed in Table 200-1. The need for intensive treatment is presumably due to the drugs' poor penetration of the thick-walled masses common in this infection and/or the sulfur granules themselves, which may represent a biofilm. Although therapy must be individualized, the IV administration of 18–24 million units of penicillin daily for 2–6 weeks, followed by oral therapy with penicillin or amoxicillin (total duration, 6–12 months), is a reasonable guideline for serious infections and bulky disease. Less extensive disease, particularly that involving the oral-cervicofacial region, may be cured with a shorter course. If therapy is extended beyond the resolution of measurable disease, the risk of relapse—a clinical hallmark of this infection—will be minimized; CT and MRI are generally the most sensitive and objective techniques



FIGURE 200-4 Computed tomogram showing pelvic actinomycosis associated with an intrauterine contraceptive device. The device is encased by endometrial fibrosis (solid arrow); also visible are paraendometrial fibrosis (open triangular arrowhead) and an area of suppuration (open arrow).

TABLE 200-1 APPROPRIATE AND INAPPROPRIATE ANTIBIOTIC THERAPY FOR ACTINOMYCOSIS^a

Category	Agent
Extensive successful clinical experience ^b	Penicillin: 3–4 million units IV q4h ^c
	Amoxicillin: 500 mg PO q6h
	Erythromycin: 500–1000 mg IV q6h or 500 mg PO q6h
	Tetracycline: 500 mg PO q6h
	Doxycycline: 100 mg IV or PO q12h
	Minocycline: 100 mg IV or PO q12h
	Clindamycin: 900 mg IV q8h or 300–450 mg PO q6h
Anecdotal successful clinical experience	Ceftriaxone ^c
	Ceftizoxime
	Imipenem-cilastatin
	Piperacillin-tazobactam
Agents that should be avoided	Metronidazole
	Aminoglycosides
	Oxacillin
	Dicloxacillin
	Cephalexin
Agents predicted to be efficacious on the basis of in vitro activity	Moxifloxacin
	Vancomycin
	Linezolid
	Quinupristin-dalfopristin
	Ertapenem ^c
	Azithromycin ^c

^aAdditional coverage for concomitant “companion” bacteria may be required. ^bControlled evaluations have not been performed. Dose and duration require individualization depending on the host, site, and extent of infection. As a general rule, a maximal parenteral antimicrobial dose for 2–6 weeks followed by oral therapy, for a total duration of 6–12 months, is required for serious infections and bulky disease, whereas a shorter course may suffice for less extensive disease, particularly in the oral-cervicofacial region. Monitoring the impact of therapy with CT or MRI is advisable when appropriate. ^cThese agents can be considered for at-home parenteral therapy; penicillin requires a continuous infusion pump.

by which to accomplish this goal. A similar approach is reasonable for immunocompromised patients, although refractory disease has been described in HIV-infected individuals. Although the role played by “companion” microbes in actinomycosis is unclear, many isolates are pathogens in their own right, and a regimen covering these organisms during the initial treatment course is reasonable.

Combined medical-surgical therapy is still advocated in some reports. However, an increasing body of literature now supports an initial attempt at cure with medical therapy alone, even in extensive disease. CT and MRI should be used to monitor the response to therapy. In most cases, either surgery can be avoided or a less extensive procedure can be used. This approach is particularly valuable in sparing critical organs, such as the bladder or the reproductive organs in women of childbearing age. For a well-defined abscess, percutaneous drainage in combination with medical therapy is a reasonable approach. When a critical location is involved (e.g., the epidural space, the CNS), when there is significant hemoptysis, or when suitable medical therapy fails, surgical intervention may be appropriate. In the absence of optimal data, the combination of a prolonged course of antimicrobial therapy and resection—at least of necrotic bone for bisphosphonate-related osteonecrosis of the jaw (BRONJ)—is a reasonable approach.

WHIPPLE'S DISEASE

Whipple's disease, a chronic multiorgan infection caused by *Tropheryma whipplei*, was first described in 1907. The long-held belief that Whipple's disease is an infection was supported by observations on its responsiveness to antimicrobial therapy in the 1950s and the

identification of bacilli via electron microscopy in small-bowel biopsy specimens in the 1960s. This hypothesis was finally confirmed by amplification and sequencing of a partial 16S rRNA polymerase chain reaction (PCR)-generated amplicon from duodenal tissue in 1991. The subsequent successful cultivation of *T. whipplei* enabled whole-genome sequencing and the development of additional diagnostic tests. The development of PCR-based diagnostics has broadened our understanding of both the epidemiology and the clinical syndromes attributable to *T. whipplei*. Exposure to *T. whipplei*, which appears to be much more common than has been appreciated, can be followed by asymptomatic carriage, acute disease, or chronic infection. Chronic infection (Whipple's disease) is a rare development after exposure. “Classic” Whipple's disease is manifested variably by a combination of arthralgias/arthritis, weight loss, chronic diarrhea, abdominal pain, and fever; less commonly, involvement at sites other than the gastrointestinal tract is documented. Acute infection and chronic organ disease in the absence of intestinal involvement (see “Isolated Infection,” below) are described with increasing frequency. Since untreated Whipple's disease is often fatal and delayed diagnosis may lead to irreparable organ damage (e.g., in the CNS), knowledge of the clinical scenarios in which Whipple's should be considered and of an appropriate diagnostic strategy is mandatory.

ETIOLOGIC AGENT



T. whipplei is a weakly staining gram-positive bacillus. Genomic sequence data have revealed that the organism has a small (<1-megabase) chromosome, with many biosynthetic pathways absent or incomplete. This finding is consistent with a host-dependent intracellular pathogen or a pathogen that requires a nutritionally rich extracellular environment. A genotyping scheme based on a variable region has disclosed more than 70 genotypes (GTs) to date. GTs 1 and 3 are most commonly reported, but all GTs appear to be capable of causing similar clinical syndromes.

EPIDEMIOLOGY



Whipple's disease is rare but has been increasingly recognized since the advent of PCR-based diagnostic tools. It occurs in all parts of the globe, with an incidence presently estimated at 1 case per 1 million patient-years. Seroprevalence studies indicate that ~50% of Western Europeans and ~75% of Africans from rural Senegal have been exposed to *T. whipplei*. A predilection for chronic disease has been observed in middle-aged Caucasian men. Males are infected five to eight times more frequently than females. To date, no clear animal or environmental reservoir has been demonstrated. However, the organism has been identified by PCR in sewage water and human feces. Workers with direct exposure to sewage are more likely to be asymptotically colonized than controls, a pattern suggesting fecal-oral spread. Recent data support oral-oral or fecal-oral spread among family members. Further, the development of acute *T. whipplei* pneumonia in children raises the possibility of droplet or airborne transmission.

PATHOGENESIS AND PATHOLOGY

Since rates of exposure to *T. whipplei* appear to be much higher (e.g., ~50% in Western Europe, as stated above) than rates of chronic disease development (0.00001%), it has been hypothesized that chronically infected individuals possess a subtle host-defense abnormality that does not place them at risk for non-*T. whipplei* infection. The HLA alleles DRB1*13 and DQB1*06 may be associated with an increased risk of infection. Chronic infection results in a general state of immunosuppression characterized by low CD4+ T cell counts, high levels of interleukin 10 production, increased activity of regulatory T cells, alternative activation of macrophages with diminished antimicrobial activity (M2 polarization) and ensuing apoptosis, and blunted development of *T. whipplei*-specific T cells. Immunosuppressive glucocorticoid treatment or anti-tumor necrosis factor α therapy appears to accelerate progression of disease. Recently, asymptomatic HIV-infected individuals were found to have significantly higher levels of *T. whipplei* sequence in bronchoalveolar lavage fluid (BALF) than did non-HIV-infected individuals, and these levels decreased

1092 with antiretroviral therapy. A weak humoral response, perhaps due to bacterial glycosylation in patients with chronic disease, appears to differentiate persons who clear the bacillus from asymptomatic carriers. In the initiation of chronic infection, the relative importance of the host's genetic background versus the modulation of the host response by *T. whipplei* is unknown.

T. whipplei has a tropism for myeloid cells, which it invades and in which it can avoid being killed. Infiltration of infected tissue by large numbers of foamy macrophages is a characteristic finding. In the intestine, villi are flat and wide with dilated lacteals. Involvement of lymphatic or hepatic tissue may manifest as noncaseating granulomas that can mimic sarcoid.

CLINICAL MANIFESTATIONS

Asymptomatic Colonization/Carriage Studies using primarily PCR have detected *T. whipplei* sequence in stool, saliva, duodenal tissue, and (rarely) blood in the absence of symptoms. Although prevalence rates are still being defined, in Western European countries, detection in saliva (0.2%) is less common than that in stool (1–11%) and appears to occur only with concomitant fecal carriage. The prevalence of fecal carriage is elevated in individuals with exposure to waste water or sewage (12–26%). However, in rural Senegal, 44% of children age 2–10 had *T. whipplei* detected in fecal samples. The duration of carriage at these sites is still being examined but can be at least 1 year. It is not known how often the carrier state is associated with acute infection, but evolution into chronic disease is uncommon. Bacterial loads are lighter in asymptomatic carriage than in active disease.

Acute Infection *T. whipplei* has been implicated as a cause of acute gastroenteritis in children. It was also detected via PCR in the blood of 6.4% of febrile patients (primarily children) from two villages in Senegal, often with concomitant cough and rhinorrhea. Further, *T. whipplei* has been implicated as a cause of acute pneumonia in the United States and France. These data suggest that primary acquisition can result in symptomatic pulmonary or intestinal infection, which may be more common than has been thought, and only rarely results in chronic disease.

Chronic Infection • "CLASSIC" WHIPPLE'S DISEASE So-called classic Whipple's disease was the initial clinical syndrome recognized, with consequent identification of *T. whipplei*. This chronic infection is defined by involvement of the duodenum and/or jejunum that develops over years. In most individuals, the initial phase of disease manifests primarily as intermittent, occasionally chronic and destructive migratory oligo- or polyarthralgias/seronegative arthritis. Spondylitis, sacroiliitis, and prosthetic hip infection also have been described. This initial stage is often confused with a variety of rheumatologic disorders and, on average, lasts 6–8 years before gastrointestinal symptoms commence. Treatment of presumed inflammatory arthritis with immunosuppressive agents (e.g., glucocorticoids, tumor necrosis factor α antagonists) can accelerate progression of the disease process. Alternatively, antimicrobial therapy used for another indication may reduce symptoms. In fact, the modulation of symptoms in these settings should prompt consideration of Whipple's disease. The intestinal symptoms that develop in the majority of cases are characterized by diarrhea with accompanying weight loss and may be associated with fever and abdominal pain. Diagnostic misdirection can be caused by co-infection with *Giardia lamblia*, which is occasionally identified. Occult gastrointestinal blood loss, hepatosplenomegaly, and ascites are less common. Anemia and hyper eosinophilia may be detected. Rheumatoid factor and antinuclear antibody tests are usually negative. The most common finding on abdominal CT is mesenteric and/or retroperitoneal lymphadenopathy. The endoscopic or video capsule observation of pale, yellow, or shaggy mucosa with erythema or ulceration past the first portion of the duodenum suggests Whipple's disease (Fig. 200-5). In addition to rheumatologic and proximal intestinal disease, neurologic (6–63%), cardiac (17–55%), pulmonary (10–40%), lymphatic (10%), ocular (5–10%), dermal (1–5%), and (in rare instances) other sites are variably involved in classic Whipple's disease.

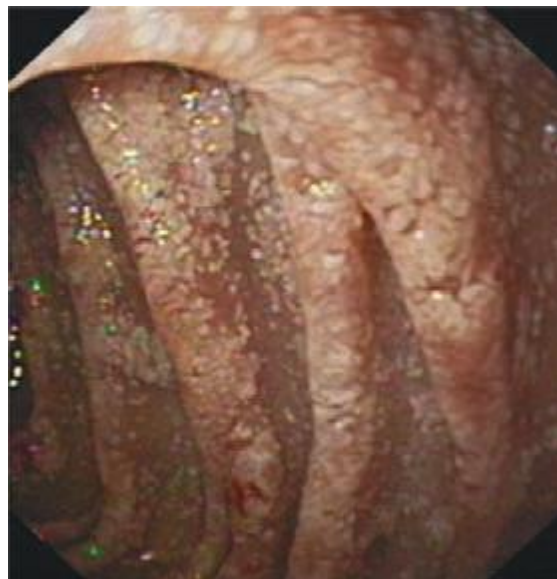


FIGURE 200-5 Endoscopic view of the jejunal mucosa demonstrating a thickened, granular mucosa and "white spots" due to dilated lacteals. (Reprinted with permission from J Bureš et al: Whipple's disease: Our own experience and review of the literature. *Gastroenterol Res Pract*, 2013. <http://dx.doi.org/10.1155/2013/478349>.)

Neurologic disease Asymptomatic neurologic involvement in Whipple's disease has been documented by PCR-based detection in cerebrospinal fluid (CSF). A variety of neurologic manifestations have been reported, the most common of which are cognitive changes progressing to dementia; personality, mood, and sleep-cycle disorders; hypothalamic involvement; and supranuclear ophthalmoplegia. In addition to the latter, neuro-ophthalmologic manifestations of Whipple's disease include supranuclear gaze palsy, oculomasticatory and oculofacial myorhythmia (highly suggestive of Whipple's), nystagmus, and retrobulbar neuritis. Focal neurologic presentations (dependent on lesion location), seizures, ataxia, meningitis, encephalitis, hydrocephalus, myelopathy, and distal polyneuropathy also have been described. Neurologic sequelae occur with CNS disease, and the mortality risk is significant.

MRI results may be normal. Identified lesions (solitary or multifocal) are usually T2 and fluid-attenuated inversion recovery (FLAIR) hyperintense and may enhance with gadolinium. Findings are myriad and not diagnostic, but the limbic system is commonly involved. FDG-PET may reveal increased uptake. CSF analysis may be abnormal; leukocytosis (generally lymphocyte-predominant) and an elevated protein concentration are common. A low CSF glucose level has been reported.

Cardiac disease Endocarditis, which is increasingly recognized in Whipple's disease, presents as culture-negative infection and/or congestive heart failure; hypotension occurs rarely. Embolic events or various arrhythmias may also be noted. Fever is often absent, and Duke clinical criteria are rarely met. Vegetations are identified by echocardiography in 50–75% of cases. All valves, alone or in combination, can be affected; most commonly involved are the aortic and mitral valves. Preexisting valvular disease is found in only a minority of cases, although infection of bioprosthetic valves has been described. Mural, myocardial, or pericardial disease also occurs alone or in combination with valvular involvement. Constrictive pericarditis develops infrequently.

Pulmonary disease Some combination of interstitial disease, nodules, parenchymal infiltrate, and pleural effusion is observed. The clinical significance of *T. whipplei* dominating sequence reads in BALF from HIV-infected individuals is unresolved.

Lymphatic disease Mesenteric and retroperitoneal lymphadenopathy are common with intestinal disease, and mediastinal adenopathy may

be associated with pulmonary infection. Peripheral adenopathy is less common.

Ocular disease (non-neuro-ophthalmologic) Uveitis is the most common form of ocular disease, usually presenting as a change in vision or “floaters.” Anterior (anterior chamber), intermediate (vitreous), and posterior (retina/choroid) uveitis can occur alone or in combination. Postoperative acute or chronic ocular Whipple's disease has been described in association with local or systemic glucocorticoid use; its detection in this setting raises the possibility that asymptomatic or subclinical disease has been unmasked. Keratitis and crystalline keratopathy also have been reported. Patients may be misdiagnosed with sarcoid or Behçet's disease prior to the recognition of Whipple's.

Dermatologic disease Skin hyperpigmentation, particularly in light-exposed areas in the absence of adrenal dysfunction, should be suggestive of Whipple's disease. A variety of other cutaneous manifestations have been described, including erythematous macular lesions, nonthrombocytopenic purpura, subcutaneous nodules, and hyperkeratosis.

Miscellaneous sites Thyroid, renal, testicular, epididymal, gallbladder, skeletal muscle, and bone marrow involvement have all been described. In fact, almost any organ can be involved in classic Whipple's disease, with varying frequency, variable combinations, and myriad signs and symptoms. As a result, Whipple's disease should be considered in the setting of a chronic multisystemic process. Despite its rarity, the combination of rheumatologic and intestinal disease with weight loss, with or without neurologic and cardiac involvement, warrants heightened suspicion.

ISOLATED INFECTION This entity has been defined as infection in the absence of intestinal symptoms, although an occasional small-bowel biopsy may be PCR-positive in this setting. “Isolated infection” is something of a misnomer since multiple nonintestinal sites of *T. whipplei* infection are not uncommon. Infection at the same nonintestinal sites (single or multiple) that are variably involved in classic Whipple's disease may also present as “isolated infection.” Endocarditis, neurologic disease, uveitis, rheumatologic manifestations, and pulmonary involvement are most commonly described. Signs and symptoms are similar to those described for *T. whipplei* infection of these sites in classic Whipple's disease. With enhanced PCR-based diagnostic capabilities, *T. whipplei* infection without concomitant intestinal involvement (of which endocarditis is the best example) will probably be diagnosed increasingly often.

REINFECTION/RELAPSING DISEASE/IMMUNE RECONSTITUTION INFLAMMATORY SYNDROME (IRIS) It has been suggested that, if an underlying host immune defect places an individual at risk for chronic infection, then that person may be at risk for reinfection due to occupational exposure or contact with family members who are asymptotically colonized. One case of apparent relapse that was due to a different genotype supports this contention.

Optimal treatment regimens and durations are still being defined. However, it is clear, especially in the setting of occult or overt CNS disease, that treatment with oral tetracycline or trimethoprim-sulfamethoxazole (TMP-SMX) alone may result in disease relapse.

As in patients treated for HIV or mycobacterial disease, IRIS has been described in patients treated for *T. whipplei* infection. Prior immunosuppressive therapy increases the likelihood of IRIS, in which inflammation recurs after an initial clinical response to treatment and loss of PCR detection of *T. whipplei*. Manifestations include fever, arthritis, skin lesions, pleuritis, uveitis, and orbital and periorbital inflammation.

DIAGNOSIS

Considering *T. whipplei* infection and ensuring that the appropriate tests are performed are the critical steps in making the diagnosis, which otherwise will likely be missed. The clinical presentation will in part dictate which clinical specimens are most likely to enable the diagnosis. In the presence (and perhaps the absence) of gastrointestinal

symptoms, postbulbar duodenal biopsies should be performed. As a general rule, diagnostic yield is greater for tissue specimens than for body fluids. Biopsy of normal-appearing skin may detect *T. whipplei* in the setting of classic Whipple's disease and serve as a minimally invasive means to establish the diagnosis. It is unclear whether CSF should be obtained in the absence of CNS symptoms, but its collection should be considered: the CNS is the most common site for relapse, and thus the information gained by CSF examination could influence the design of the treatment regimen.

The development and implementation of PCR-based diagnostics have significantly increased the sensitivity and specificity of *T. whipplei* identification. PCR can be applied to affected tissues (fixed and nonfixed) and various body fluids (e.g., CSF; aqueous or vitreous humor; joint, pericardial, or pleural fluid; BALF; blood; feces). In some clinical scenarios, a generic 16S rRNA bacterial assay combined with amplicon sequencing can be used to detect and identify *T. whipplei* sequence. Delineation of the *T. whipplei* genomic sequence has enabled the development and broad availability of more sensitive and specific PCR-based assays. The interpretation of a PCR-based diagnostic approach must take into account limitations such as false-positive results due to sample contamination and false-negative results due to organism load, sample quality, and inadequate DNA extraction.

The diagnosis of classic Whipple's disease was originally based on histologic findings in intestinal biopsy specimens, and this diagnostic procedure remains important. Infiltration of the lamina propria with macrophages containing inclusions (representing ingested bacteria) that are positive on periodic acid-Schiff (PAS) staining and resistant to diastase is observed. However, PAS is nonspecific, also yielding positive results with mycobacteria (which can be differentiated with Ziehl-Neelsen stain), *Rhodococcus equi*, *Bacillus cereus*, *Corynebacterium* species, and *Histoplasma* species. *T. whipplei* can also be detected by silver stain, Brown-Brenn (weakly positive), or acridine orange and is not stained by calcofluor. Staining of other tissues or fluids (e.g., ocular aspirations) for PAS-positive inclusions in macrophages can be performed to support the diagnosis. Electron microscopy can be used to identify the trilaminar cell wall of *T. whipplei*.

When available, immunohistochemistry has greater specificity and sensitivity than PAS staining and can be performed on archived fixed tissue. *T. whipplei* has been successfully cultured from blood, CSF, synovial fluid, BALF, valve tissue, duodenal tissue, skeletal muscle, and lymph nodes, but culture is not practical since it takes months to obtain a positive result. Likewise, serology is of limited value for the diagnosis of Whipple's disease because the prevalence of exposure is much higher than that of chronic disease development and the antibody response to *T. whipplei* appears to be blunted in the disease state.

Although histologic or cytologic detection of *T. whipplei* is less specific and sensitive than PCR, a positive result is strongly supportive within the appropriate clinical context and is definitive when combined with a more specific test (e.g., PCR, immunohistochemistry).

TREATMENT WHIPPLE'S DISEASE

Data on treatment are emerging, but questions persist regarding the optimal regimen and duration, which may depend on the site of infection (e.g., CNS and heart valve). Appropriate treatment usually results in a rapid and at times remarkable clinical response (e.g., in CNS disease). Maintenance of a durable response has been more challenging.

Rates of relapse, particularly of CNS disease, were unacceptable with oral tetracycline or TMP-SMX monotherapy. Sequence data now indicate that TMP is not active against *T. whipplei* due to the absence of dihydrofolate reductase, but this drug was used extensively before this fact was known. This information prompted a randomized controlled trial in 40 patients, who received either ceftriaxone (2 g IV q24h) or meropenem (1 g IV q8h) for 2 weeks followed by oral TMP-SMX (160/800 mg) twice a day for 1 year. The efficacy of these regimens was outstanding. The only instance of therapy failure—in a case of asymptomatic CNS infection that was

1094 not eradicated by either regimen—was subsequently cured with oral minocycline and chloroquine (250 mg/d after a loading dose). A follow-up trial reported similar efficacy with a regimen of ceftriaxone (2 g IV q24h) for 2 weeks followed by oral TMP-SMX for 3 months. One issue in these trials was that the CNS doses—and perhaps the duration of ceftriaxone and meropenem treatment as well—were not optimal. Further, investigators have speculated that oral regimens with greater CNS penetrance, such as sulfadiazine (2–4 g/d in 3 or 4 divided doses) and/or doxycycline or minocycline (200 mg/d in 2 divided doses) plus hydroxychloroquine (200 mg three times a day, to raise phagosome pH and increase drug activity in vitro), might render the parenteral phase of treatment unnecessary, given that the one failure of therapy for CNS disease was cured with a similar regimen. Another issue is concern about the potential development of resistance to sulfa drugs. Lastly, it is unclear whether oral sulfa- or tetracycline-based regimens will suffice in endocarditis. Until more data become available, it seems prudent—at least in asymptomatic/symptomatic CNS disease or cardiac infection—to administer CNS-optimized doses of IV ceftriaxone (2 g q12h) or meropenem (2 g q8h) for at least 2 weeks followed by oral doxycycline or minocycline plus hydroxychloroquine or chloroquine for at least 1 year, if tolerated. Although data on the use of PCR to guide therapy do not exist, it seems reasonable that continued *T. whipplei* detection by PCR, especially in the CSF, should dictate at least continuation of therapy and perhaps consideration of an alternative regimen.

The occurrence of a Jarisch-Herxheimer reaction within 24 h of treatment initiation has been described, with rapid resolution. The addition of glucocorticoids may be beneficial in the management of clearly documented IRIS.

Data on certain site-specific treatment issues are even more limited. Anecdotal reports describe successful treatment of uveitis with oral TMP-SMX with or without rifampin, whereas treatment with tetracycline alone has resulted in relapse. Although a role for adjunctive intraocular therapy has been reported, the data are unclear on this point. Surgery may be needed in the setting of endocarditis with significant valve dysfunction; however, timely recognition can result in cure with medical management alone. Although data on the treatment of foreign body-associated infection are virtually nonexistent, medical treatment for a prosthetic hip infection was apparently successful; however, follow-up was limited.

Regardless of the therapeutic regimen chosen, an effort to ensure compliance and close follow-up for potential relapse (or perhaps reinfection), which can occur many years after an apparent cure, will maximize the chances for a good outcome.

cultivating and identifying these organisms in clinical microbiology laboratories continue to leave the anaerobic etiology of an infectious process unproven in many cases. Therefore, an understanding of the types of infections in which anaerobes can play a role is crucial in selecting appropriate microbiologic tools to identify the organisms in clinical specimens and in choosing the most appropriate treatment, including antibiotics and surgical drainage or debridement of the infected site.

This chapter focuses on infections caused by nonsporulating anaerobic bacteria. It does not address clostridial infections and syndromes, which are covered elsewhere ([Chaps. 161 and 179](#)).

DEFINITIONS

Anaerobic bacteria are organisms that require reduced oxygen tension for growth, failing to grow on the surface of solid media in 10% CO₂ in air. (In contrast, *microaerophilic bacteria* can grow in an atmosphere of 10% CO₂ in air or under anaerobic or aerobic conditions, although they grow best in the presence of only a small amount of atmospheric oxygen, and *facultative bacteria* can grow in the presence or absence of air.) Most clinically relevant anaerobes, such as *Bacteroides fragilis*, *Prevotella melaninogenica*, and *Fusobacterium nucleatum*, are relatively aerotolerant. Although they can survive for sustained periods in the presence of up to 2–8% oxygen, they generally do not multiply in this environment. A smaller number of pathogenic anaerobic bacteria (which are also part of the microbiota) die after brief contact with oxygen, even in low concentrations.

ANAEROBES OF THE HUMAN MICROBIOTA

Most human mucocutaneous surfaces harbor a rich indigenous normal microbiota composed of aerobic and anaerobic bacteria. These surfaces are dominated by anaerobic bacteria, which often account for 99.0–99.9% of the culturable microbiota and range in concentration from 10⁹/mL in saliva to 10¹²/mL in gingival scrapings and the colon. It is interesting that anaerobes inhabit many areas of the body that are exposed to air: skin, nose, mouth, and throat. Anaerobes are thought to reside in the portions of these sites that are relatively well protected from oxygen, such as gingival crevices. New technologies based on analyses of microbial DNA have expanded our knowledge of these bacterial populations. For example, in an analysis of 13,555 prokaryotic ribosomal RNA gene sequences from the colon, most bacteria identified were considered uncultivated and novel microorganisms. Two immense projects based on these new technologies, the Human Microbiome Project funded by the U.S. National Institutes of Health and MetaHIT financed by the European Commission, aim to characterize the normal microbiota of healthy individuals.

The major reservoirs of anaerobic bacteria are the mouth, lower gastrointestinal tract, skin, and female genital tract ([Table 201-1](#)). In the oral cavity, the ratio of anaerobic to aerobic bacteria ranges from 1:1 on the surface of a tooth to 1000:1 in the gingival crevices. *Prevotella* and *Porphyromonas* species comprise much of the indigenous oral anaerobic microbiota. *Fusobacterium* and *Bacteroides* (non-*B. fragilis* group) are present in lower numbers. Anaerobic bacteria are not found in appreciable numbers in the normal stomach and upper small intestine. In the distal ileum, the microbiota begins to resemble that of the colon. In the colon, the ratio of anaerobes to facultative species is high; for example, there are 10¹¹–10¹² organisms/g of stool, and >99% of these organisms are anaerobic, with an anaerobe-to-aerobe ratio of ~1000:1. The predominant anaerobes in the human intestine belong to the phyla Bacteroidetes and Firmicutes and include a number of *Bacteroides* species (e.g., members of the *B. fragilis* group, such as *B. fragilis*, *B. thetaiotaomicron*, *B. ovatus*, *B. vulgatus*, *B. uniformis*, and *Parabacteroides distasonis*) as well as various clostridial, peptostreptococcal, and fusobacterial species. In the female genital tract, there are ~10⁹ organisms/mL of secretions, with an anaerobe-to-aerobe ratio of 1:1 to 10:1. The predominant anaerobes in the female genital tract are *Prevotella*, *Bacteroides*, *Fusobacterium*, *Clostridium*, and the anaerobic *Lactobacillus* species.

201 Infections Due to Mixed Anaerobic Organisms

Ronit Cohen-Pradosu, Dennis L. Kasper

Anaerobes comprise the predominant class of bacteria of the normal human microbiota (formerly termed “the normal human flora”) that reside on mucous membranes and predominate in many infectious processes, particularly those arising from mucosal surfaces. These organisms generally cause disease subsequent to the breakdown of mucosal barriers and the leakage of the microbiota into normally sterile sites. Infections resulting from contamination by the microbiota are usually polymicrobial and involve both aerobic and anaerobic bacteria. However, the difficulties encountered in handling specimens in which anaerobes may be important and the technical challenges entailed in

TABLE 201-1 ANAEROBIC HUMAN MICROBIOTA: AN OVERVIEW

Anatomic Site	Total Bacteria ^a	Anaerobic/Aerobic Ratio	Potential Pathogens
Oral cavity			
Saliva	10 ⁸ –10 ⁹	1:1	<i>Fusobacterium nucleatum</i> , <i>Prevotella melaninogenica</i> , <i>Prevotella oralis</i> group, <i>Bacteroides ureolyticus</i> group, <i>Peptostreptococcus</i> spp.
Tooth surface	10 ¹⁰ –10 ¹¹	1:1	
Gingival crevices	10 ¹¹ –10 ¹²	10 ³ :1	
Gastrointestinal tract			
Stomach	0–10 ⁵	1:1	<i>Bacteroides</i> spp. (principally members of the <i>B. fragilis</i> group), <i>Prevotella</i> spp., <i>Clostridium</i> spp., <i>Peptostreptococcus</i> spp.
Jejunum/ileum	10 ⁴ –10 ⁷	1:1	
Terminal ileum and colon	10 ¹¹ –10 ¹²	10 ³ :1	
Female genital tract	10 ⁷ –10 ⁹	10:1	<i>Peptostreptococcus</i> spp., <i>Bacteroides</i> spp., <i>Prevotella bivia</i>

^aPer gram or milliliter.

The skin microbiota contains anaerobes as well, the predominant species being *Propionibacterium acnes* and, in lower numbers, other species of propionibacteria and peptostreptococci.

Commensal bacteria in general and commensal anaerobes in particular have been implicated as crucial mediators of physiologic, metabolic, and immunologic functions in the mammalian host. One of the most important roles that anaerobes serve as components of the normal colonic microbiota is the promotion of resistance to colonization; the presence of anaerobic bacteria effectively interferes with colonization by potentially pathogenic bacterial species through the depletion of oxygen and nutrients, the production of enzymes and toxic end products, and the modulation of the host's intestinal innate immune response. For example, *B. thetaiotaomicron* stimulates Paneth cells to produce RegIIIγ, a bactericidal lectin that can result in killing of gram-positive bacteria. The normal colonic microbiota plays an important role in protection against *Clostridium difficile*-associated diarrhea or colitis—a toxin-mediated, potentially life-threatening disease that results when *C. difficile* spores in the colon transform into toxin-producing vegetative forms after antibiotic elimination of critical components of the competing colonic microbiota.

Bacteroides and other intestinal bacteria ferment carbohydrates and produce volatile fatty acids that are reabsorbed and used by the host as an energy source. The anaerobic intestinal microbiota is also responsible for the production of secreted products that promote human health (e.g., vitamin K and bile acids useful in fat absorption and cholesterol regulation).

Moreover, the anaerobic intestinal microbiota influences the development of an intact mucosa and of mucosa-associated lymphoid tissue. Colonization of germ-free mice with a single species, *B. thetaiotaomicron*, affects the expression of various host genes and corrects deficiencies of nutrient uptake, metabolism, angiogenesis, mucosal barrier function, and enteric nervous system development. The symbiosis factor polysaccharide A (PSA) of *B. fragilis* influences the normal development and function of the mammalian immune system and protects mice against colitis in a model of inflammatory bowel disease. It has also been shown that PSA can confer protection both prophylactically and therapeutically, restraining inflammatory processes at an extraintestinal site (the central nervous system [CNS]) and ameliorating disease in a mouse model of multiple sclerosis. Anaerobes can stimulate specific lymphocyte populations of the small and large intestine and can influence immunologic balance (including T_H1/T_H2 balance) as well as the number of T_H17 and regulatory T cells in gut tissues.

Clearly, the gut microbiota confers many benefits, and its dysregulation may play a role in the pathogenesis of diseases characterized by inflammation and aberrant immune responses, such as inflammatory bowel disease, rheumatoid arthritis, multiple sclerosis, asthma, and type 1 diabetes. Furthermore, the gut microbiota has been associated with obesity and metabolic syndrome. An interesting association between certain microbes found in the microbiota and testosterone production has been suggested as well.

ETIOLOGY

Thousands of species of anaerobic bacteria have been identified as components of the complete human microbiota, with each individual colonized by hundreds of these species. Despite the complex array of bacteria in the normal microbiota, relatively few species are isolated commonly from human infections. Anaerobic infections occur when the harmonious relationship between the host and the host's microbiota is disrupted. Any site in the body is susceptible to infection with these indigenous organisms when a mucosal barrier or the skin is compromised by surgery, trauma, tumor, ischemia, or necrosis, all of which can reduce local tissue redox potentials. Because the sites that are colonized by anaerobes contain many species of bacteria, disruption of anatomic barriers allows contamination of normally sterile sites by many organisms, resulting in mixed infections involving multiple species of anaerobes in combination with synergistically acting facultative or microaerophilic organisms.

Severe mixed infections of the head and neck may arise from an abscessed tooth infected with commensal microbiota of the mouth. Examples of infections arising from an oral source are chronic sinusitis, chronic otitis media, Ludwig's angina, and periodontal abscesses. Brain abscesses and subdural empyema are also commonly associated with the oral microbiota. Oral anaerobes are usually responsible for pleuropulmonary diseases such as aspiration pneumonia, necrotizing pneumonia, lung abscess, and empyema.

Anaerobes from the intestine play an important role in various intraabdominal infections, such as peritonitis and intraabdominal abscesses (Chap. 159). Colonic contents are the source of microorganisms in the case of these infections, which usually follow disruption of intestinal continuity and contamination of the peritoneal cavity. Anaerobic bacteria are isolated frequently in female genital tract infections, such as salpingitis, pelvic peritonitis, tuboovarian abscess, vulvovaginal abscess, septic abortion, and endometritis (Chap. 163). In addition, these bacteria are often found in bacteremia and in infections of the skin, soft tissues, and bones.

Predominant among the anaerobic gram-positive cocci that produce disease are the peptostreptococci; the species of this genus that are most commonly involved in infections are *P. micros*, *P. magnus*, *P. asaccharolyticus*, *P. anaerobius*, and *P. prevotii*. Clostridia (Chap. 179) are anaerobic spore-forming gram-positive rods that are isolated from wounds, abscesses, sites of abdominal infection, and blood. Gram-positive anaerobic non-spore-forming bacilli are uncommon as etiologic agents of human infection. *P. acnes*, a component of the skin microbiota and a rare cause of foreign-body infections, is one of the few nonclostridial gram-positive rods associated with infections. The principal anaerobic gram-negative bacilli found in human infections belong to the *B. fragilis* group and to *Fusobacterium*, *Prevotella*, and *Porphyromonas* species.

The most important potential anaerobic pathogens found in the upper airways and isolated from clinical specimens of oral and

1096 pleuropulmonary infections are the *Fusobacterium* species *F. necrophorum*, *F. nucleatum*, and *F. varium*; *P. melaninogenica*; the *Prevotella* *oralis* group; *Porphyromonas gingivalis*; *Porphyromonas asaccharolytica*; *Peptostreptococcus* species; and the *Bacteroides ureolyticus* group.

The *B. fragilis* group contains the anaerobic pathogens most frequently isolated from clinical infections. Members of this group are part of the normal bowel microbiota; they include several distinct species, such as *B. fragilis*, *B. thetaiotaomicron*, *B. vulgatus*, *B. uniformis*, *B. ovatus*, and *P. distasonis*. *B. fragilis* is the most important clinical isolate, although it is isolated in lower numbers than some other *Bacteroides* species from cultures of the commensal fecal microbiota.

In female genital tract infections, organisms normally colonizing the vagina (e.g., *Prevotella bivia* and *Prevotella disiens*) are the most common isolates. However, *B. fragilis* is not uncommon.

PATHOGENESIS

Anaerobic bacterial infections usually occur when an anatomic barrier is disrupted and constituents of the local microbiota enter a site that was previously sterile. Because of the specific growth requirements of anaerobic organisms and their presence as commensals on mucosal surfaces, conditions must arise that allow these organisms to penetrate mucosal barriers and enter tissue with a lowered oxidation-reduction potential. Therefore, tissue ischemia, trauma, surgery, perforated viscus, shock, and aspiration provide environments conducive to the proliferation of anaerobes. The introduction of many bacterial species into otherwise sterile sites leads to a polymicrobial infection in which certain organisms predominate. Three major factors are involved in the pathogenesis of anaerobic infections: bacterial synergy, bacterial virulence factors, and mechanisms of abscess formation. The ability of different anaerobic bacteria to act synergistically during polymicrobial infection contributes to the pathogenesis of anaerobic infections. It has been postulated that facultative organisms function in part to lower the oxidation-reduction potential in the microenvironment, allowing the propagation of obligate anaerobes. Anaerobes can produce compounds such as succinic acid and short-chain fatty acids that inhibit the ability of phagocytes to clear facultative organisms. In experimental models, facultative and obligate anaerobes synergistically potentiate abscess formation.

Virulence factors associated with anaerobes typically confer the ability to evade host defenses, adhere to cell surfaces, produce toxins and/or enzymes, or display surface structures such as capsular polysaccharides and lipopolysaccharide (LPS) that contribute to pathogenic potential. The ability of an organism to adhere to host tissues is important to the establishment of infection. Some oral species adhere to the epithelium in the oral cavity. *P. melaninogenica* actually attaches to other microorganisms. *P. gingivalis*, a common isolate in periodontal disease, has fimbriae that facilitate attachment. Some *Bacteroides* strains appear to be piliated, a characteristic that may account for their ability to adhere.

The most extensively studied virulence factor of the nonsporulating anaerobes is the capsular polysaccharide complex of *B. fragilis*. This organism is unique among anaerobes in its potential for virulence during growth at normally sterile sites. Although it constitutes only 0.5–1% of the normal colonic microbiota, *B. fragilis* is the anaerobe most commonly isolated from intraabdominal infections and bacteremia. In an animal model of intraabdominal sepsis, the capsular polysaccharide was identified as the major virulence factor of *B. fragilis*; this polymer plays a specific, central role in the induction of abscesses. A series of detailed biologic and molecular studies of this virulence factor showed that *B. fragilis* produces at least eight distinct capsular polysaccharides, far more than the number reported for any other encapsulated bacterium. *B. fragilis* can exhibit distinct surface polysaccharides either alone or in combination by regulating the expression of these different capsules in an on-off manner through a reversible inversion of DNA segments within the promoters for operons containing the genes required for polysaccharide synthesis. Structural analysis of two of these polysaccharides, PSA

and PSB, revealed that each polymer consists of repeating units with positively charged free amino groups and negatively charged groups. This structural feature is rare among bacterial polysaccharides, and the ability of PSA—and, to a lesser extent, PSB—to induce abscesses in animals depends on this zwitterionic charge motif. Intraabdominal abscess induction is related to the capacity of this polysaccharide to stimulate macrophages to release cytokines and chemokines—in particular, interleukin (IL) 8, IL-17, and tumor necrosis factor α (TNF- α)—from resident peritoneal cells through a Toll-like receptor 2-dependent mechanism. The release of cytokines and chemokines results in the chemotaxis of polymorphonuclear neutrophils (PMNs) into the peritoneum, where they adhere to mesothelial cells induced by TNF- α to upregulate their expression of intercellular adhesion molecule 1 (ICAM-1). PMNs adherent to ICAM-1-expressing cells probably represent the nidus for an abscess. PSA also activates T cells to produce certain cytokines, including IL-17 and interferon γ , that are necessary for abscess formation.

B. fragilis produces other virulence factors that allow it to predominate in disease. This organism synthesizes pili, fimbriae, and hemagglutinins that aid in attachment to host cell surfaces. In addition, *Bacteroides* species produce many enzymes and toxins that contribute to pathogenicity. Enzymes such as neuraminidase, protease, glycoside hydrolases, and superoxide dismutases are all produced by *B. fragilis*. Anaerobic bacteria produce a number of exoproteins that can enhance the organisms' virulence. The collagenase produced by *P. gingivalis* may enhance tissue destruction. An association of *B. fragilis* strains positive for the enterotoxin BFT with clinical episodes of diarrhea in children and adults has been suggested. BFT is a metalloprotease that is cytopathic for intestinal epithelial cells and induces fluid secretion and tissue damage in ligated intestinal loops of experimental animals. Recent evidence from mouse models indicates that enterotoxin-producing strains of *B. fragilis* may play a role in colon carcinoma.

Exotoxins produced by clostridial species, including botulinum toxins, tetanus toxin, *C. difficile* toxins A and B, and five toxins produced by *Clostridium perfringens*, are among the most virulent bacterial toxins in mouse lethality assays. Anaerobic gram-negative bacteria such as *B. fragilis* possess LPSs (endotoxins) that are 100–1000 times less biologically potent than endotoxins associated with aerobic gram-negative bacteria. This relative biologic inactivity may account for the lower frequency of disseminated intravascular coagulation and purpura in *Bacteroides* bacteremia than in facultative and aerobic gram-negative bacillary bacteremia. An exception is the LPS from *Fusobacterium*, which may account for the severity of Lemierre's syndrome (see "Complications of Anaerobic Head and Neck Infections," below).

APPROACH TO THE PATIENT: Infection Due to Anaerobic Bacteria

The physician must consider several points when approaching the patient with possible infection due to anaerobic bacteria.

1. Most of the organisms colonizing mucosal sites are commensals; very few cause disease. When these organisms do cause disease, it often occurs in proximity to the mucosal site they colonize.
2. For anaerobes to cause tissue infection, they must spread beyond the normal mucosal barriers.
3. Conditions favoring the propagation of anaerobic bacteria, particularly a lowered oxidation-reduction potential, are necessary. These conditions exist at sites of trauma, tissue destruction, compromised vascular supply, and complications of preexisting infection, which produce necrosis.
4. Frequently, a complex array of infecting microbes can be found. For example, as many as 12 species of organisms can be isolated from a suppurative site.
5. Anaerobic organisms tend to be found in abscess cavities or in necrotic tissue. The failure of an abscess to yield organisms

on routine culture is a clue that the abscess is likely to contain anaerobic bacteria. Often smears of this “sterile pus” are found to be teeming with bacteria when Gram’s stain is applied. Although some facultative organisms (e.g., *Staphylococcus aureus*) are also capable of causing abscesses, abscesses in organs or deeper body tissues should call anaerobic infection to mind.

6. Gas is found in many anaerobic infections of deep tissues but is not diagnostic because it can be produced by aerobic bacteria as well.
7. Although a putrid-smelling infection site or discharge is considered diagnostic for anaerobic infection, this manifestation usually develops late in the course and is present in only 30–50% of cases.
8. Some species (the best example being the *B. fragilis* group) require specific therapy. However, many synergistic infections can be cured with antibiotics directed at some but not all of the organisms involved. Antibiotic therapy, combined with debridement and drainage, disrupts the interdependent relationship among the bacteria, and some species that are resistant to the antibiotic do not survive without the co-infecting organisms.
9. Manifestations of severe sepsis and disseminated intravascular coagulation are unusual in patients with purely anaerobic infection.

EPIDEMIOLOGY

Difficulties in the performance of appropriate cultures, contamination of cultures by components of the normal microbiota, and the lack of readily available, reliable culture techniques have made it impossible to obtain accurate data on incidence or prevalence. However, anaerobic infections are encountered frequently in hospitals with active surgical, trauma, and obstetric and gynecologic services. Depending on the institution, anaerobic bacteria account for 0.5–12% of all cases of bacteremia.

CLINICAL MANIFESTATIONS

Anaerobic Infections of the Mouth, Head, and Neck Anaerobic bacteria are commonly involved in infections of the mouth, head, and neck (Chap. 44). The predominant isolates are components of the normal microbiota of the upper airways—mainly the *Bacteroides oralis* group, pigmented *Prevotella* species, *P. asaccharolytica*, *Fusobacterium* species, peptostreptococci, and microaerophilic streptococci.

Soft tissue infections of the oral-facial area may or may not be odontogenic. Odontogenic infections—primarily dental caries and periodontal disease (gingivitis and periodontitis)—are common and have both local consequences (especially tooth loss) and the potential for life-threatening spread to the deep fascial spaces of the head and neck. Infections of the mouth can arise from either supragingival or subgingival dental plaque composed of bacteria colonizing the tooth surface. Supragingival plaque formation begins with the adherence of gram-positive bacteria to the tooth surface. This form of plaque is influenced by salivary and dietary components, oral hygiene, and local host factors. Supragingival plaque can lead to dental caries and, with further invasion, to pulpitis (endodontic infection) that can further perforate the alveolar bone, causing periapical abscess. Subgingival plaque is associated with periodontal infections (e.g., gingivitis, periodontitis, and periodontal abscess) that can further disseminate to adjacent structures such as the mandible, causing osteomyelitis of the maxillary sinuses. Periodontitis may also result in spreading infection that can involve adjacent bone or soft tissues. In the healthy periodontium, the sparse microbiota consists mainly of gram-positive organisms such as *Streptococcus sanguinis* and *Actinomyces* species. In the presence of gingivitis, there is a shift to a greater proportion of anaerobic gram-negative bacilli in the subgingival microbiota, with predominance of *Prevotella intermedia*. In well-established periodontitis, the complexity of the microbiota increases further. The predominant isolates are *P. gingivalis*, *P. intermedia*, *Aggregatibacter actinomycetemcomitans*, *Tannerella forsythensis*.

Necrotizing Ulcerative Gingivitis Gingivitis may become a necrotizing infection (*trench mouth*, *Vincent’s stomatitis*) (Chap. 44). The onset of disease is usually sudden and is associated with tender bleeding

gums, foul breath, and a bad taste. The gingival mucosa, especially the papillae between the teeth, becomes ulcerated and may be covered by a gray exudate, which is removable with gentle pressure. Patients may become systemically ill, developing fever, cervical lymphadenopathy, and leukocytosis.



Noma (*cancrum oris*) is a necrotizing infection of the oral mucous membranes. It is characterized by destruction of soft tissue and bone and evolves rapidly from gingival inflammation to orofacial gangrene. Noma occurs most frequently in young children (1–4 years of age) with malnutrition or systemic disease. This infection occurs worldwide but is most common in sub-Saharan Africa.

Acute Necrotizing Infections of the Pharynx These infections usually occur in association with ulcerative gingivitis. Symptoms include an extremely sore throat, foul breath, and a bad taste accompanied by fever and a sensation of choking. Examination of the pharynx demonstrates that the tonsillar pillars are swollen, red, ulcerated, and covered with a grayish membrane that peels easily. Lymphadenopathy and leukocytosis are common. The disease may last for only a few days or, if not treated, may persist for weeks. Lesions begin unilaterally but may spread to the other side of the pharynx or the larynx. Aspiration of the infected material by the patient can result in lung abscesses.

Peripharyngeal Space Infections These infections arise from the spread of organisms from the upper airways to potential spaces formed by the fascial planes of the head and neck. The etiology is typically polymicrobial and represents the normal microbiota of the mucosa of the originating site.

Peritonsillar abscess (*quinsy*) is a complication of acute tonsillitis caused mainly by a mixed flora containing anaerobes (e.g., *F. necrophorum* and *Peptostreptococcus* species) and the facultative anaerobe group A *Streptococcus* (Chap. 44). Of cases of submandibular space infection (*Ludwig’s angina*), 80% are caused by infection of the tissues surrounding the second and third molar teeth. This infection results in marked local swelling of tissues, with pain, trismus, and superior and posterior displacement of the tongue. Submandibular swelling of the neck can impair swallowing and cause respiratory obstruction. In some cases, tracheotomy is life-saving. Cervicofacial actinomycosis (Chap. 200) is caused by a branching, gram-positive, non-spore-forming, strict/facultative anaerobe that is a part of the normal oral microbiota. This chronic disease is characterized by abscesses, draining sinus tracts, fistula, bone destruction, and fibrosis. It can easily be mistaken for malignancy or granulomatous disease. Actinomycosis less frequently involves the thorax, abdomen, pelvis, and CNS.

Sinusitis and Otitis Anaerobic bacteria have been implicated in chronic sinusitis but play little role in acute sinusitis. In several studies on chronic sinusitis, anaerobic bacteria were found in 0–52% of cases, depending on the method used to collect specimens. Polymicrobial infection is common, and the predominant anaerobic isolates are pigmented *Prevotella*, *Fusobacterium*, *Peptostreptococcus*, and *P. acnes*. Aerobic gram-negative bacilli and *S. aureus* have also been implicated in chronic sinusitis. Anaerobic bacteria have been isolated in a large percentage of cases of chronic suppurative otitis media in children. The role of anaerobes in acute otitis media is less clear.

Complications of Anaerobic Head and Neck Infections Contiguous cranial spread of these infections can result in osteomyelitis of the skull or mandible or in intracranial infections such as brain abscess and subdural empyema. Caudal spread can produce mediastinitis or pleuropulmonary infection. Hematogenous complications can also result from anaerobic infections of the head and neck. Bacteremia, which occasionally is polymicrobial, can lead to endocarditis or other distant infections. Lemierre’s syndrome (Chap. 44), which has been uncommon in the antimicrobial era, is an acute oropharyngeal infection with secondary septic thrombophlebitis of the internal jugular vein and frequent metastasis, most commonly to the lung. *F. necrophorum* is the usual cause. This infection typically begins with pharyngitis, which is followed by local invasion in the lateral pharyngeal space, with resultant internal jugular vein thrombophlebitis. A typical clinical

1098 triad includes pharyngitis, a tender/swollen neck, and noncavitating pulmonary infiltrates.

CNS Infections CNS infections associated with anaerobic bacteria are brain abscess, epidural abscess, and subdural empyema. Anaerobic meningitis is rare and is usually related to parameningeal collection or shunt infection. If optimal bacteriologic techniques are used, as many as 85% of brain abscesses yield anaerobic bacteria. Most anaerobic brain abscesses arise by direct extension from a site of otorhinolaryngeal infection such as otitis, sinusitis, or tooth infection. Hematogenous dissemination from a distant infected site, usually intraabdominal or pelvic, can occur. Common isolates are *Peptostreptococcus*, *Fusobacterium*, *Bacteroides*, *Prevotella*, *Propionibacterium*, *Eubacterium*, *Veillonella*, and *Actinomyces* species. Facultative or microaerophilic streptococci and coliforms are often part of a mixed infecting flora in brain abscesses.

Pleuropulmonary Infections Anaerobic pleuropulmonary infections result from the aspiration of oropharyngeal contents by patients with predisposing conditions such as dysphagia due to neurologic or esophageal disorders or transiently impaired consciousness due to conditions such as alcohol or drug abuse, seizures, head trauma, and cerebrovascular accidents. Clinical syndromes that are associated with anaerobic pleuropulmonary infection produced by aspiration include aspiration pneumonitis, which can be complicated by necrotizing pneumonia, lung abscess, and empyema. Many of these infections have an indolent course that may serve as a clinical clue differentiating them, for example, from pneumococcal pneumonia, which often presents with abrupt onset, shaking chills, and rapid progression.

The anaerobes most common in pleuropulmonary infections are indigenous to the oral cavity, especially the gingival crevice, and include pigmented and nonpigmented *Prevotella*, *Peptostreptococcus*, and *Bacteroides* species and *F. nucleatum*. Many of these infections are of mixed aerobic-anaerobic etiology, and the predominant aerobes isolated from community-acquired aspiration pneumonias are microaerophilic streptococci such as *Streptococcus milleri*. Studies using in-depth culture techniques in patients with community-acquired lung abscess showed aerobic and microaerophilic streptococci to be the most common pathogens (60% of patients) and anaerobes to be the second most common (26%). In a study on aspiration pneumonia from a long-term care facility, the most common isolates were gram-negative bacilli (49%), anaerobes (16%), and *S. aureus* (12%). Nosocomial aspiration pneumonia commonly involves a mixture of anaerobes and gram-negative bacilli or *S. aureus*.

ASPIRATION PNEUMONITIS Bacterial aspiration pneumonitis must be distinguished from two other clinical syndromes associated with aspiration that are not of bacterial etiology. One syndrome results from aspiration of solids, usually food. Obstruction of major airways typically results in atelectasis and moderate nonspecific inflammation. Therapy consists of removal of the foreign body. The second aspiration syndrome is more easily confused with bacterial aspiration. *Mendelson's syndrome*, a chemical pneumonitis, results from regurgitation of stomach contents and aspiration of chemical material, usually acidic gastric juices. Pulmonary inflammation—including the destruction of the alveolar lining, with transudation of fluid into the alveolar space—occurs with remarkable rapidity. Typically this syndrome develops within hours, often following anesthesia when the gag reflex is depressed. The patient becomes tachypneic, hypoxic, and febrile. The leukocyte count may rise, and the chest x-ray may evolve from normal to a complete bilateral “whiteout” within 8–24 h. Sputum production is minimal. The pulmonary signs and symptoms can resolve quickly with symptom-based therapy or can culminate in respiratory failure, with the subsequent development of bacterial superinfection over a period of days. Antibiotic therapy is not indicated unless bacterial infection supervenes.

In contrast to these syndromes, bacterial aspiration pneumonitis develops over a period of several days or weeks rather than hours. Patients who enter the hospital with this syndrome typically have been ill for several days and generally report low-grade fever, malaise, and

sputum production. In some patients, weight loss and anemia reflect a more chronic process. Usually the history reveals factors predisposing to aspiration, such as alcohol overdose or residence in a nursing home. Examination sometimes yields evidence of periodontal disease. Sputum characteristically is not malodorous unless the process has been under way for at least a week. A mixed bacterial flora with many PMNs is evident on Gram's staining of sputum. Expecterated sputum is unreliable for anaerobic cultures because of inevitable contamination by the normal oral microbiota. Reliable specimens for culture can be obtained by transtracheal or transthoracic aspiration—techniques that are rarely used at present. Culture of protected-brush specimens or bronchoalveolar lavage fluid obtained by bronchoscopy is controversial.

Chest x-rays show consolidation in dependent pulmonary segments: in the basilar segments of the lower lobes if the patient has aspirated while upright and in either the posterior segment of the upper lobe (usually on the right side) or the superior segment of the lower lobe if the patient has aspirated while supine.

NECROTIZING PNEUMONITIS This form of anaerobic pneumonitis is characterized by numerous small abscesses that spread to involve several pulmonary segments. The process can be indolent or fulminating. This syndrome is less common than either aspiration pneumonitis or lung abscess and includes features of both types of infection.

ANAEROBIC LUNG ABSCESSSES (See also Chap. 154) These abscesses result from subacute anaerobic pulmonary infection. The clinical syndrome typically involves a history of constitutional signs and symptoms (including malaise, weight loss, fever, night sweats, and foul-smelling sputum), perhaps over a period of weeks (Chap. 153). Patients who develop lung abscesses characteristically have dental infection and periodontitis, but lung abscesses in edentulous patients have been reported. Abscess cavities may be single or multiple and generally occur in dependent pulmonary segments (Fig. 201-1). Anaerobic abscesses must be distinguished from lesions associated with tuberculosis, neoplasia, and other conditions.

Septic pulmonary emboli may originate from intraabdominal or female genital tract infections and can produce anaerobic pneumonia and abscess.

EMPHYEMA Empyema is a manifestation of long-standing anaerobic pulmonary infection complicated by bronchopleural fistula. The clinical presentation, which includes foul-smelling sputum, resembles that



FIGURE 201-1 Chest radiograph of right-lower-lobe lung abscess in a 60-year-old alcoholic patient. (From GL Mandell [ed]: *Atlas of Infectious Diseases*, Vol VI. Philadelphia, Current Medicine Inc, Churchill Livingstone, 1996; with permission.)

of other anaerobic pulmonary infections. Patients may report pleuritic chest pain and marked chest-wall tenderness.

Empyema may be masked by overlying pneumonitis and should be considered especially in cases of persistent fever despite antibiotic therapy. Diligent physical examination and the use of ultrasound to localize a loculated empyema are important diagnostic tools. The collection of a foul-smelling exudate by thoracentesis is typical. Cultures of infected pleural fluid yield an average of 3.5 anaerobic and 0.6 facultative or aerobic bacterial species. Drainage is required. Defervescence, a return to a feeling of well-being, and resolution of the process may require several months.

Extension from a subdiaphragmatic infection may also result in anaerobic empyema.

Intraabdominal Infections Intraabdominal infections—mainly peritonitis and abscesses—are usually polymicrobial and represent the normal intestinal (especially colonic) microbiota. These infections most often follow a breach in the mucosal barrier resulting from appendicitis, diverticulitis, neoplasm, inflammatory bowel disease, surgery, or trauma. On average, four to six bacterial species are isolated per specimen submitted to the microbiology laboratory, with a predominance of enteric aerobic/facultative gram-negative bacilli, anaerobes, and streptococci/enterococci. The most common isolates are *Escherichia coli* (found in $\geq 50\%$ of patients) and *B. fragilis* (30–50%). Other anaerobes commonly isolated from this type of infection include *Peptostreptococcus*, *Prevotella*, and *Fusobacterium* species. The involvement of clostridia can lead to severe infections. The dominance of four to six bacterial species out of the more than 500 colonic mucosal species is related both to the virulence factors of these species and to the inability of clinical laboratories to culture many other species residing in the colonic mucosa.

Disease originating from proximal-bowel perforation reflects the microbiota of this site, with a predominance of aerobic and anaerobic gram-positive bacteria and *Candida*.

Neutropenic enterocolitis (typhlitis) has been associated with anaerobic infection of the cecum but—in the setting of neutropenia (**Chap. 104**)—may involve the entire bowel. Patients usually present with fever; abdominal pain, tenderness, and distention; and watery diarrhea. The bowel wall is edematous with hemorrhage and necrosis. The primary pathogen is thought by some authorities to be *Clostridium septicum*, but other clostridia and mixed anaerobes have also been implicated. More than 50% of patients developing early clinical signs can benefit from antibiotic therapy and bowel rest. Surgery is sometimes required to remove gangrenous bowel. **See Chap. 159 for a complete discussion of intraabdominal infections.**

Enterotoxigenic *B. fragilis* has been associated with watery diarrhea in a few young children and adults. In case-control studies of children with undiagnosed diarrheal disease, enterotoxigenic *B. fragilis* was isolated from significantly more children with diarrhea than children in the control group.

Pelvic Infections The vagina of a healthy woman is a major reservoir of anaerobic and aerobic bacteria. In the normal microbiota of the female genital tract, anaerobes outnumber aerobes by a ratio of $\sim 10:1$ and include anaerobic gram-positive cocci and *Bacteroides* species (Table 201-1). Anaerobes are isolated from most women with genital tract infections that are not caused by a sexually transmitted pathogen. The major anaerobic pathogens are *B. fragilis*, *P. bivia*, *P. disiens*, *P. melaninogenica*, anaerobic cocci, and *Clostridium* species. Anaerobes are frequently encountered in pelvic inflammatory disease, pelvic abscess, endometritis, tubo-ovarian abscess, septic abortion, and postoperative or postpartum infections. These infections are often of mixed etiology, involving both anaerobes and coliforms; pure anaerobic infections without coliform or other facultative bacterial species occur more often in pelvic than in intraabdominal sites. Septic pelvic thrombophlebitis may complicate the infections and lead to repeated episodes of septic pulmonary emboli. **See Chap. 163 for a complete discussion of pelvic inflammatory disease.**

Anaerobic bacteria have been thought to be contributing factors in the etiology of bacterial vaginosis. This syndrome of unknown etiology is characterized by a profuse malodorous discharge and a change in

the bacterial ecology that results in replacement of the *Lactobacillus*-dominated normal microbiota with an overgrowth of bacterial species including *Gardnerella vaginalis*, *Prevotella* species, *Mobiluncus* species, peptostreptococci, and genital mycoplasmas. A study based on 16S rRNA identification found other anaerobes that were predominant in cases but not in controls: *Atopobium*, *Leptotrichia*, *Megasphaera*, and *Eggerthella*. Pelvic infections due to *Actinomyces* species have been associated with the use of intrauterine devices (**Chap. 200**).

Skin and Soft Tissue Infections Injury to skin, bone, or soft tissue by trauma, ischemia, or surgery creates a suitable environment for anaerobic infections. These infections are most frequently found in sites prone to contamination with feces or with upper airway secretions—e.g., wounds associated with intestinal surgery, decubitus ulcers, or human bites. Moreover, anaerobes have been isolated from cutaneous abscesses, rectal abscesses, and axillary sweat gland infections (*hidradenitis suppurativa*). Anaerobes also are often cultured from foot ulcers of diabetic patients. The deep soft-tissue infections associated with anaerobic bacteria are crepitant cellulitis, synergistic cellulitis, gangrene, and necrotizing fasciitis (**Chaps. 156 and 179**).

These soft tissue or skin infections are usually polymicrobial. A mean of 4.8 bacterial species are isolated, with an anaerobe-to-aerobe ratio of $\sim 3:2$. The most frequently isolated organisms include *Bacteroides*, *Peptostreptococcus*, *Clostridium*, *Enterococcus*, and *Proteus* species. The involvement of anaerobes in these types of infections is associated with a higher frequency of fever, foul-smelling lesions, gas in the tissues, and visible foot ulcer.

Anaerobic bacterial synergistic gangrene (*Meleney's gangrene*), a rare infection of the superficial fascia, is characterized by exquisite pain, redness, and swelling followed by induration. Erythema surrounds a central zone of necrosis. A granulating ulcer forms at the original center as necrosis and erythema extend outward. Symptoms are limited to pain; fever is not typical. These infections usually involve a combination of *Peptostreptococcus* species and *S. aureus*; the usual site of infection is an abdominal surgical wound or the area surrounding an ulcer on an extremity. Treatment includes surgical removal of necrotic tissue and antimicrobial administration.

Necrotizing fasciitis, a rapidly spreading destructive disease of the fascia, is usually attributed to group A streptococci (**Chap. 173**) but can also be a mixed infection involving anaerobes and aerobes, usually occurring after surgeries and in patients with diabetes or peripheral vascular disease. The most frequently isolated anaerobes in these infections are *Peptostreptococcus* and *Bacteroides* species. Gas may be found in the tissues. Similarly, myonecrosis can be associated with mixed anaerobic infection. *Fournier's gangrene* consists of cellulitis involving the scrotum, perineum, and anterior abdominal wall, with mixed anaerobic organisms spreading along deep external fascial planes and causing extensive loss of skin.

Bone and Joint Infections Although actinomycosis (**Chap. 200**) accounts on a worldwide basis for most anaerobic infections in bone, organisms including peptostreptococci or microaerophilic cocci, *Bacteroides* species, *Fusobacterium* species, and *Clostridium* species can also be involved in osteomyelitis (**Chap. 158**). These infections frequently arise adjacent to soft tissue infections. Many patients with osteomyelitis due to anaerobic bacteria have evidence of an anaerobic infection elsewhere in the body; most commonly, infected adjacent soft-tissue sites are the source of the organisms involved. Examples are diabetic foot ulcers and decubitus ulcers that may be complicated by mixed aerobic-anaerobic osteomyelitis. Hematogenous seeding of bone is uncommon. *Prevotella* and *Porphyromonas* species are detected in infections involving the maxilla and mandible, whereas *Clostridium* species have been reported as anaerobic pathogens in cases of osteomyelitis of the long bones following fracture or trauma. *Fusobacteria* have been isolated in pure culture from sites of osteomyelitis adjacent to the perinasal sinuses. Peptostreptococci and microaerophilic cocci have been reported as significant pathogens in infections involving the skull, mastoid, and prosthetic implants placed in bone. In patients with osteomyelitis, the most reliable culture specimen is a bone biopsy sample free of normal uninfected skin and subcutaneous tissue.

In contrast to anaerobic osteomyelitis, most cases of anaerobic arthritis (**Chap. 157**) involve a single isolate, and most cases are secondary to hematogenous spread. The most common isolates are *Fusobacterium* species. Most of the patients involved have uncontrolled peritonissillar infections progressing to septic cervical venous thrombophlebitis (Lemierre's syndrome) and resulting in hematogenous dissemination with a predilection for the joints. Unlike anaerobic osteomyelitis, anaerobic pyoarthritides in most cases is not polymicrobial and may be acquired hematogenously. Anaerobes are important pathogens in infections involving prosthetic joints; in these infections, the causative organisms (such as *Peptostreptococcus* species and *P. acnes*) are part of the normal skin microbiota.

Bacteremia Transient bacteremia is a well-known event in healthy individuals whose anatomic mucosal barriers have been injured (e.g., during dental extractions or dental scaling). These bacteremic episodes, which are often due to anaerobes, have no pathologic consequences. However, anaerobic bacteria are found in cultures of blood from clinically ill patients when proper culture techniques are used. Anaerobes have accounted for 5% (range at various institutions, 0.5–12%) of cases of clinically significant bacteremia. The incidence of anaerobic bacteremia decreased from the 1970s through the early 1990s. This change may have been related to the administration of antibiotic prophylaxis before intestinal surgery, the earlier recognition of localized infections, and the empirical use of broad-spectrum antibiotics for presumed infection. Recent reports present conflicting data regarding rates of anaerobic bacteremia. A study from the Mayo Clinic compared three periods (1993–1996, 1997–2000, and 2001–2004) and found a 74% increase in the mean incidence of anaerobic bacteremia; this finding contrasts with a 45% decrease in incidence from 1977 to 1988 at the same institution. In contrast, a report from Switzerland compared two periods (1997–2001 and 2002–2006) and found decreases in both the number of anaerobe-positive blood cultures and the proportion of all blood culture isolates that were anaerobes.

The majority of anaerobic bacteremias are due to gram-negative bacilli—mainly the *B. fragilis* group, with *B. fragilis* most commonly isolated (60–80% of cases). Other organisms causing bacteremia include *Clostridium* species (10%), *Peptostreptococcus* species (10%), and *Fusobacterium* species (5%).

Once the organism in the blood has been identified, both the portal of bloodstream entry and the underlying problem that probably led to seeding of the bloodstream can often be deduced from an understanding of the organism's normal site of residence. For example, mixed anaerobic bacteremia including *B. fragilis* usually implies a colonic pathology with mucosal disruption from neoplasia, diverticulitis, or some other inflammatory lesion. Debilitating diseases such as malignancies, diabetes, organ transplantation, and abdominal and pelvic surgeries are among the predisposing factors for anaerobic bacteremia. In a retrospective nested case-control study, diabetes was identified as a risk factor for anaerobic bacteremia when the source of bacteremia was unknown. The initial manifestations are determined by the portal of entry and reflect the localized condition. When bloodstream invasion occurs, patients can become extremely ill, with rigors and hectic fevers. The clinical picture may be quite similar to that seen in sepsis involving aerobic gram-negative bacilli. Although complications of anaerobic bacteremia (e.g., septic thrombophlebitis and septic shock) have been reported, their incidence in association with anaerobic bacteremia is low. Anaerobic bacteremia is potentially fatal and requires rapid diagnosis and appropriate therapy. Reported case-fatality rates are high, ranging from 25% to 44%, and appear to increase with the age of the patient (with reported rates of >66% among patients >60 years old), with the isolation of multiple species from the bloodstream, and with the failure to surgically remove a focus of infection. The attributable mortality rate for bacteremia associated with the *B. fragilis* group was examined in a matched case-control study. Patients with *B. fragilis*-group bacteremia had a significantly higher mortality rate (28% vs 8%), with an attributable mortality rate of 19.3% and a mortality risk ratio of 3.2.

Endocarditis and Pericarditis (See also **Chap. 155**) Endocarditis due to anaerobes is uncommon. However, anaerobic streptococci, which are

often classified incorrectly, are responsible for this disease more frequently than is generally appreciated. Gram-negative anaerobes are unusual causes of endocarditis. Signs and symptoms of anaerobic endocarditis are similar to those of endocarditis due to facultative organisms. Mortality rates of 21–43% have been reported for anaerobic endocarditis.

Anaerobes, particularly *B. fragilis* and *Peptostreptococcus* species, are uncommonly found in infected pericardial fluids. Anaerobic pericarditis is associated with a mortality rate of >50%. Anaerobes can reach the pericardial space by hematogenous spread, by spread from a contiguous site of infection (e.g., heart or esophagus), or by direct inoculation arising from trauma or surgery.

DIAGNOSIS

There are three critical steps in the diagnosis of anaerobic infection: (1) proper collection of specimens; (2) rapid transport of the specimens to the microbiology laboratory, preferably in anaerobic transport media; and (3) proper handling of the specimens by the laboratory. Specimens must be collected by meticulous sampling of infected sites, with avoidance of contamination by the normal microbiota. When such contamination is likely, the specimen is unacceptable. Examples of specimens unacceptable for anaerobic culture include sputum collected by expectoration or nasal tracheal suction, bronchoscopy specimens, samples collected directly through the vaginal vault, urine collected by voiding, and feces. Specimens appropriate for anaerobic culture include sterile body fluids such as blood, pleural fluid, peritoneal fluid, cerebrospinal fluid, and aspirates or biopsy samples from normally sterile sites. As a general rule, liquid or tissue specimens are preferred; swab specimens should be avoided.

Because even brief exposure to oxygen may kill some anaerobic organisms and result in failure to isolate them in the laboratory, air must be expelled from the syringe used to aspirate the abscess cavity, and the needle must be capped with a sterile rubber stopper. It is also important to remember that prior antibiotic therapy reduces the cultivability of these bacteria. Specimens can be injected into transport bottles containing a reduced medium or taken immediately in syringes to the laboratory for direct culture on anaerobic media. Delays in transport may lead to a failure to isolate anaerobes due to exposure to oxygen or overgrowth of facultative organisms, which may eliminate or obscure any anaerobes that are present. All clinical specimens from suspected anaerobic infections should be subjected to Gram's staining and examined for organisms with characteristic morphology. It is not unusual for organisms to be observed on Gram's staining but not isolated in culture.

Because of the time and difficulty involved in the isolation of anaerobic bacteria, diagnosis of anaerobic infections must frequently be based on presumptive evidence. There are few clinical clues to the probable presence of anaerobic bacteria at infected sites. The involvement of certain sites with lowered oxidation-reduction potential (e.g., avascular necrotic tissues) and the presence of an abscess favor the diagnosis of an anaerobic infection. When infections occur in proximity to mucosal surfaces normally harboring an anaerobic microbiota, such as the gastrointestinal tract, female genital tract, or oropharynx, anaerobes should be considered as potential etiologic agents. A foul odor is often indicative of anaerobes, which produce certain organic acids as they proliferate in necrotic tissue. Although these odors are nearly pathognomonic for anaerobic infection, the absence of odor does not exclude an anaerobic etiology. The presence of gas in tissues is highly suggestive, but not diagnostic, of anaerobic infection. Because anaerobes often coexist with other bacteria and cause mixed or synergistic infection, Gram's staining of exudate frequently reveals multiple morphotypes suggestive of anaerobes. Sometimes these organisms have morphologic characteristics associated with specific species.

When cultures of obviously infected sites or purulent material yield no growth, streptococci only, or a single aerobic species (such as *E. coli*) and Gram's staining reveals a mixed flora, the involvement of anaerobes should be suspected; the implication is that the anaerobic microorganisms have failed to grow because of inadequate transport and/or culture techniques. Failure of an infection to respond to antibiotics that are not active against anaerobes (e.g., aminoglycosides and—

some circumstances—penicillin, cephalosporins, or tetracyclines) suggests an anaerobic etiology.

TREATMENT ANAEROBIC INFECTIONS

Successful therapy for anaerobic infections requires the administration of a combination of appropriate antibiotics, surgical resection, debridement of devitalized tissues, and drainage either surgically or percutaneously (guided by an imaging technique such as CT, MRI, or ultrasound). Any anatomic breach must be closed promptly, closed spaces drained, tissue compartments decompressed, and an adequate blood supply established. Abscess cavities should be drained as soon as fluctuation or localization occurs.

ANTIBIOTIC THERAPY AND RESISTANCE

The antibiotics used to treat anaerobic infections should be active against both aerobic and anaerobic organisms because many of these infections are of mixed etiology. Antibiotic regimens can usually be selected empirically on the basis of the type of infection, the species of the organisms usually present in such cases, the results of Gram's staining, and a knowledge of antimicrobial resistance patterns (Chap. 170 and Table 201-2). Other factors influencing the selection of antibiotics include need for bactericidal activity and for penetration into certain organs (such as the brain), toxicity, and impact on the normal microbiota. Antibiotics active against clinically relevant anaerobes can be grouped into four categories based on their predicted activity (Table 201-2). Nearly all the drugs listed have toxic side effects, which are described in detail in Chap. 170.

Antibiotic susceptibility testing of anaerobic bacteria has been difficult and controversial. Because of the slow growth rate of many anaerobes, the lack of standardized testing methods and of clinically relevant standards for resistance, and the generally good results obtained with empirical therapy, there has been limited interest in testing these organisms for antibiotic susceptibility. However, one study of antibiotic-treated patients with *Bacteroides* isolates from blood found mortality rates of 45% among those whose isolates were deemed resistant to the agent used and 16% among those whose isolates were deemed sensitive. It is accepted that testing is important for patients with serious or prolonged infections or in cases in which antibiotics have not had an impact. Testing is also helpful in monitoring the activity of new drugs and recording current resistance patterns among anaerobic pathogens. The antibiotics with the greatest activity against nearly all anaerobic bacteria include carbapenems, β -lactam/ β -lactamase inhibitor combinations, metronidazole, and chloramphenicol.



Antibiotic resistance in anaerobic bacteria is an increasing problem. Resistance rates vary with the institution and the geographic region. In recent years, the activity of clindamycin, ceftioxin, cefotetan, and moxifloxacin has decreased against *B. fragilis* and related strains (*B. distasonis*, *B. ovatus*, *B. thetaiotaomicron*, *B. uniformis*, *B. vulgatus*). Multidrug-resistant *B. fragilis* has recently been

reported. Nearly all organisms in the *B. fragilis* group (>97%) are resistant to penicillin G. The cephamycins ceftioxin and cefotetan display greater activity against this group, but rates of resistance have increased, with current figures at ~10% in the United States and higher in Argentina (28%) and Europe (17%). Rates of resistance to β -lactam agents among anaerobes other than *Bacteroides* are lower but are highly variable. β -Lactam/ β -lactamase inhibitor combinations such as ampicillin/sulbactam, ticarcillin/clavulanic acid, and piperacillin/tazobactam are usually good therapeutic options against β -lactamase-producing anaerobes, including the *B. fragilis* group. Although resistance rates reported from most countries are still low, several studies have documented nonsusceptibility to ampicillin/sulbactam in 0.5–3% of isolates in the United States, 3–10% in Europe, and 1–8% in Argentina. Recently, up to 48% of *B. fragilis* isolates in Taiwan were found to be nonsusceptible to ampicillin/sulbactam, and a significant increase in resistance to this combination was also identified among other *Bacteroides*, *Prevotella*, and *Fusobacterium* species.

Carbapenems (ertapenem, doripenem, meropenem, and imipenem) are equally active against anaerobes, with <1% of *B. fragilis* strains showing resistance in the United States and Europe. Higher rates of carbapenem nonsusceptibility are being reported from some countries (5% in Germany, 8% [to doripenem] in Canada, and 7–12% in Taiwan).

Metronidazole is active against gram-negative anaerobes, including the *B. fragilis* group; resistance, although rare (<1%), has been reported in both Europe and the United States. Resistance to metronidazole is more common among gram-positive anaerobes, including *P. acnes*, *Actinomyces* species, lactobacilli, and anaerobic streptococci. Clindamycin is active against many anaerobes. However, rates of resistance to clindamycin among the *B. fragilis* group have increased in the United States from 3% in 1982 to 16% in 1996 and 26% in 2000, with rates as high as 40–50% in some series. Resistance to clindamycin among non-*Bacteroides* anaerobes is much less common (<10%).

Tigecycline is active against some anaerobic bacteria, including *Peptostreptococcus*, *Propionibacterium*, *Prevotella*, *Fusobacterium*, and most *Bacteroides* species. Its efficacy for treatment of intraabdominal infections was comparable to that of imipenem in two phase 3 double-blind clinical trials. This drug is therefore recommended as single-agent treatment for complicated intraabdominal infections, but resistance (~6%) among *Bacteroides* and non-*Bacteroides* species has been reported.

Fluoroquinolones such as moxifloxacin have shown potential in the treatment of mixed aerobic-anaerobic infections. A survey in the United States found a 38% rate of resistance to moxifloxacin among the *B. fragilis* group; in Europe 14–30% of isolates were nonsusceptible to this drug, as were 7–25% of anaerobes isolated from blood cultures in Taiwan. Despite excellent in vitro activity against all clinically important anaerobes, chloramphenicol is less desirable than other active drugs for the treatment of anaerobic infection because of documented clinical failures.

TABLE 201-2 ANTIMICROBIAL THERAPY FOR INFECTIONS INVOLVING COMMONLY ENCOUNTERED ANAEROBIC GRAM-NEGATIVE RODS

Category 1 (Nearly Always Active)	Category 2 (Usually Active)	Category 3 (Variable Resistance)	Category 4 (Resistance)
Carbapenems (imipenem, meropenem, doripenem)	Tigecycline	Cephamycins (ceftioxin, cefotetan)	Aminoglycosides
Metronidazole ^a	High-dose antipseudomonal penicillins	Clindamycin	Monobactams
β -Lactam/ β -lactamase inhibitor combination (ampicillin/sulbactam, ticarcillin/clavulanic acid, piperacillin/tazobactam)		Penicillins	Trimethoprim-sulfamethoxazole
Chloramphenicol ^b		Cephalosporins	
		Tetracycline	
		Vancomycin	
		Erythromycin	
		Moxifloxacin	

^aUsually needs to be given in combination with aerobic bacterial coverage. For infections originating below the diaphragm, aerobic gram-negative coverage is essential. For infections from an oral source, aerobic gram-positive coverage is added. Metronidazole also is not active against *Actinomyces*, *Propionibacterium*, or other gram-positive non-spore-forming bacilli (e.g., *Eubacterium*, *Bifidobacterium*) and is unreliable against peptostreptococci. ^bDespite excellent in vitro activity against all clinically important anaerobes, this drug is less desirable than other active drugs because of documented clinical failures.

1102 If a patient fails to respond to one of the category 1 or category 2 drugs (Table 201-2), consideration should be given to alternative therapy and to determination of the resistance patterns among *Bacteroides* isolates.

INFECTIONS AT SPECIFIC SITES

In clinical situations, specific regimens must be tailored to the initial site of infection. The duration of therapy also depends on the infection site; the reader is referred to specific chapters on sites of infection for recommendations.

Infections above the diaphragm usually reflect the orodental microbiota, which does not include the *B. fragilis* group. β -Lactamase production has been reported in anaerobic strains that are usually isolated from infections originating above the diaphragm. Up to 60% of clinical isolates classified as *Prevotella* or *Porphyromonas* species, non-*B. fragilis* species of *Bacteroides*, or *Fusobacterium* species reportedly produce β -lactamase; thus all β -lactam drugs (penicillins and cephalosporins) are poor options. Because most of these infections have a mixed etiology that includes microaerophilic and aerobic streptococci, antibiotics that cover both aerobic and anaerobic bacteria are recommended. The recommended regimens include clindamycin, a β -lactam/ β -lactamase inhibitor combination, or metronidazole in combination with a drug active against microaerophilic and aerobic streptococci.

Bronchoscopy in lung abscess is indicated only to rule out airway obstruction and does not enhance drainage; in any event, it should be delayed until the antimicrobial regimen has begun to affect the disease process so that the procedure does not spread the infection. Surgery is almost never indicated because of the danger of spilling the abscess contents into the lungs.

Chloramphenicol has been used successfully against anaerobic CNS infections at doses of 30–60 mg/kg per day, with the exact dose depending on the severity of illness. However, penicillin G and metronidazole also cross the blood–brain barrier and are bactericidal for many anaerobic organisms.

Anaerobic infections arising below the diaphragm (e.g., colonic and intraabdominal infections) must be treated specifically with agents active against *Bacteroides* species (Table 201-2). In intraabdominal sepsis (Chap. 159), the use of antibiotics effective against penicillin-resistant anaerobes has clearly reduced the incidence of postoperative infections and serious infectious complications. Specifically, a drug from category 1 (Table 201-2) must be included for broad-spectrum coverage. Single agents suitable for this purpose include the carbapenems, cefoxitin, and β -lactam/ β -lactamase inhibitor combinations. A two-drug regimen is an alternative, with one drug active against coliforms and the other against anaerobes (e.g., a third-generation cephalosporin or a quinolone with metronidazole).

In addition, if the clinician suspects that gram-positive facultative organisms such as enterococci are involved, therapeutic regimens should include ampicillin or vancomycin. Although clindamycin and cefotetan were previously considered acceptable options for intraabdominal infections involving anaerobes, these drugs are no longer recommended because of escalating rates of resistance in the *B. fragilis* group. Ampicillin/sulbactam is not recommended because of high rates of resistance among community-acquired strains of *E. coli* rather than because of resistance in anaerobic bacteria.

A meta-analysis of 40 randomized or quasi-randomized controlled trials of 16 antibiotic regimens for secondary peritonitis showed equivalent clinical success for all regimens.

Cases of anaerobic osteomyelitis in which a mixed flora is isolated from a bone biopsy specimen should be treated with a regimen that covers all isolates. When an anaerobic organism is recognized as a major or sole pathogen infecting a joint, the duration of treatment should be similar to that used for arthritis caused by aerobic bacteria (Chap. 157). Therapy includes the management of underlying disease states, the administration of appropriate antimicrobial agents, temporary joint immobilization, percutaneous drainage of effusions, and (usually) the removal of infected prostheses or internal fixation devices. Surgical drainage and debridement procedures such as sequestrectomy are essential for the removal of necrotic tissue that can sustain anaerobic infections.

The outcome of anaerobic bacteremia is significantly better in patients either initially given or switched to appropriate therapy on the basis of known antibiotic susceptibilities.

FAILURE OF THERAPY

Anaerobic infections that fail to respond to treatment or that relapse should be reassessed. Consideration should be given to additional surgical drainage or debridement. Superinfections with resistant gram-negative facultative or aerobic bacteria should be ruled out. The possibility of drug resistance must be entertained; if resistance is involved, repeat cultures may yield the pathogen.

SUPPORTIVE MEASURES

Other supportive measures in the management of anaerobic infections include careful attention to fluid and electrolyte balance (since extensive local edema may lead to hypoalbuminemia), hemodynamic support for septic shock, immobilization of infected extremities, maintenance of adequate nutrition during chronic infections by parenteral hyperalimentation, relief of pain, and anticoagulation for thrombophlebitis. For patients with severe anaerobic infections of soft tissues, hyperbaric oxygen therapy is advocated by some experts, but its value has not been proven in controlled trials.

SECTION 8 MYCOBACTERIAL DISEASES

202 Tuberculosis

Mario C. Raviglione

Tuberculosis (TB), which is caused by bacteria of the *Mycobacterium tuberculosis* complex, is one of the oldest diseases known to affect humans and a major cause of death worldwide. Recent population genomic studies suggest that *M. tuberculosis* may have emerged ~70,000 years ago in Africa and subsequently disseminated along with anatomically modern humans, expanding globally during the Neolithic Age as human density started to increase. Progenitors of *M. tuberculosis* are likely to have

affected prehumans. This disease most often affects the lungs, although other organs are involved in up to one-third of cases. If properly treated, TB caused by drug-susceptible strains is curable in the vast majority of cases. If untreated, the disease may be fatal within 5 years in 50–65% of cases. Transmission usually takes place through the airborne spread of droplet nuclei produced by patients with infectious pulmonary TB.

ETIOLOGIC AGENT

Mycobacteria belong to the family Mycobacteriaceae and the order Actinomycetales. Of the pathogenic species belonging to the *M. tuberculosis* complex, which comprises eight distinct subgroups, the most common and important agent of human disease is *M. tuberculosis*. The

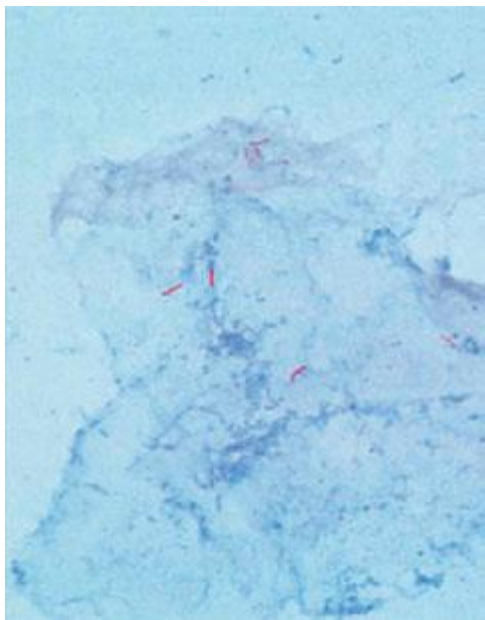


FIGURE 202-1 Acid-fast bacillus smear showing *M. tuberculosis* bacilli. (Courtesy of the Centers for Disease Control and Prevention, Atlanta.)

complex includes *M. bovis* (the bovine tubercle bacillus—characteristically resistant to pyrazinamide, once an important cause of TB transmitted by unpasteurized milk, and currently the cause of a small percentage of human cases worldwide), *M. caprae* (related to *M. bovis*), *M. africanum* (isolated from cases in West, Central, and East Africa), *M. microti* (the “vole” bacillus, a less virulent and rarely encountered organism), *M. pinnipedii* (a bacillus infecting seals and sea lions in the Southern Hemisphere and recently isolated from humans), *M. mungi* (isolated from banded mongooses in southern Africa), *M. orygis* (described recently in oryxes and other Bovidae in Africa and Asia and a potential cause of infection in humans), and *M. canetti* (a rare isolate from East African cases that produces unusual smooth colonies on solid media and is considered closely related to a supposed progenitor type).

M. tuberculosis is a rod-shaped, non-spore-forming, thin aerobic bacterium measuring 0.5 μm by 3 μm . Mycobacteria, including *M. tuberculosis*, are often neutral on Gram’s staining. However, once stained, the bacilli cannot be decolorized by acid alcohol; this characteristic justifies their classification as acid-fast bacilli (AFB; Fig. 202-1). Acid fastness is due mainly to the organisms’ high content of mycolic acids, long-chain cross-linked fatty acids, and other cell-wall lipids. Microorganisms other than mycobacteria that display some acid fastness include species of *Nocardia* and *Rhodococcus*, *Legionella micdadei*, and the protozoa *Isospora* and *Cryptosporidium*. In the mycobacterial cell wall, lipids (e.g., mycolic acids) are linked to underlying arabinogalactan and peptidoglycan. This structure results in very low permeability of the cell wall, thus reducing the effectiveness of most antibiotics. Another molecule in the mycobacterial cell wall, lipoarabinomannan, is involved in the pathogen–host interaction and facilitates the survival of *M. tuberculosis* within macrophages. The complete genome sequence of *M. tuberculosis* comprises 4043 genes encoding 3993 proteins and 50 genes encoding RNAs; its high guanine-plus-cytosine content (65.6%) is indicative of an aerobic “lifestyle.” A large proportion of genes are devoted to the production of enzymes involved in cell wall metabolism.

EPIDEMIOLOGY



More than 5.7 million new cases of TB (all forms, both pulmonary and extrapulmonary) were reported to the World Health Organization (WHO) in 2013; 95% of cases were reported from developing countries. However, because of insufficient case detection and incomplete notification, reported cases may represent only about two-thirds of the total estimated cases. The WHO estimated that

9 million (range, 8.6–9.4 million) new cases of TB occurred worldwide in 2013, 95% of them in developing countries of Asia (5 million), Africa (2.6 million), the Middle East (0.7 million), and Latin America (0.3 million). It is further estimated that 1.49 million (range, 1.32–1.67 million) deaths from TB, including 0.36 million among people living with HIV infection, occurred in 2013, 96% of them in developing countries. Estimates of TB incidence rates (per 100,000 population) and numbers of TB-related deaths in 2013 are depicted in Figs. 202-2 and 202-3, respectively. During the late 1980s and early 1990s, numbers of reported cases of TB increased in industrialized countries. These increases were related largely to immigration from countries with a high incidence of TB; the spread of the HIV epidemic; social problems, such as increased urban poverty, homelessness, and drug abuse; and dismantling of TB services. During the past few years, numbers of reported cases have begun to decline again or have stabilized in most industrialized nations. In the United States, with the re-establishment of stronger control programs, the decline resumed in 1993 and has since been maintained. In 2013, 9582 cases of TB (3.0 cases/100,000 population) were reported to the Centers for Disease Control and Prevention (CDC).

In the United States, TB is uncommon among young adults of European descent, who have only rarely been exposed to *M. tuberculosis* infection during recent decades. In contrast, because of a high risk of transmission in the past, the prevalence of latent *M. tuberculosis* infection (LTBI) is relatively high among elderly whites. In general, adults ≥ 65 years of age have the highest incidence rate per capita (4.9 cases/100,000 population in 2013) and children < 14 years of age the lowest (0.8 case/100,000 population). Blacks account for the highest proportion of cases (37%; 1257 cases in 2013) among U.S.-born persons. TB in the United States is also a disease of adult members of the HIV-infected population, the foreign-born population (64.6% of all cases in 2013), and disadvantaged/marginalized populations. Of the 6193 cases reported among foreign-born persons in 2013, 37% occurred in persons from the Americas and 32% occurred in persons born in the Western Pacific region. Overall, the highest rates per capita were among Asian Americans (18.7 cases/100,000 population). A total of 536 deaths were caused by TB in the United States in 2011. In Canada in 2013, 1638 TB cases were reported (4.7 cases/100,000 population); 70% (1145) of these cases occurred in foreign-born persons and 19% (309 cases) occurred in members of the Canadian aboriginal peoples, whose per capita rate is disproportionately high (23.4 cases/100,000 population) with a peak in the Nunavut territory of 143 cases/100,000 population—a rate similar to that in many highly endemic countries. Similarly, in Europe, TB has reemerged as an important public health problem, mainly as a result of cases among immigrants from high-incidence countries and among marginalized populations, often in large urban settings like London; in 2013, 41% of all cases reported from the United Kingdom occurred in London, and the rate per capita (36 cases/100,000 population) was similar to that in some middle-income countries. In most Western European countries, there are more cases annually among foreign-born than native populations.

Recent data on global trends indicate that in 2013 the TB incidence was stable or falling in most regions; this trend began in the early 2000s and appears to have continued, with an average annual decline of 2% globally. This global decrease is explained largely by the simultaneous reduction in TB incidence in sub-Saharan Africa, where rates had risen steeply since the 1980s as a result of the HIV epidemic and the lack of capacity of health systems and services to deal with the problem effectively, and in Eastern Europe, where incidence increased rapidly during the 1990s because of a deterioration in socioeconomic conditions and the health care infrastructure (although, after peaking in 2001, incidence in Eastern Europe has since declined slowly).

Of the estimated 9 million new cases of TB in 2013, 13% (1.1 million) were associated with HIV infection, and 78% of these HIV-associated cases occurred in Africa. An estimated 0.36 million persons with HIV-associated TB died in 2013. Furthermore, an estimated 480,000 cases (range, 350,000–610,000) of multidrug-resistant TB (MDR-TB)—a form of the disease caused by bacilli resistant at least to isoniazid and rifampin—occurred in 2013. Only 28% of these

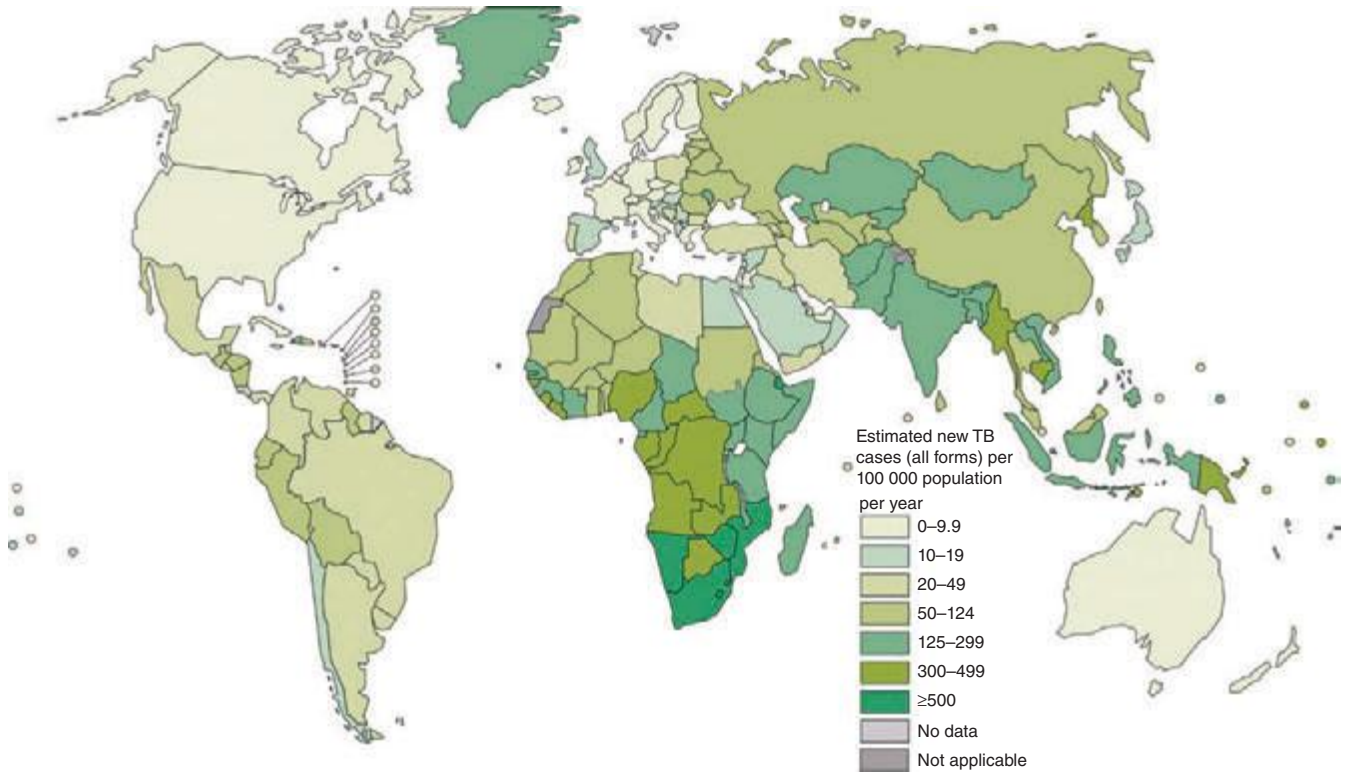


FIGURE 202-2 Estimated tuberculosis (TB) incidence rates (per 100,000 population) in 2013. The designations used and the presentation of material on this map do not imply the expression of any opinion whatsoever on the part of the World Health Organization (WHO) concerning the legal status of any country, territory, city, or area or of its authorities or concerning the delimitation of its frontiers or boundaries. Dotted, dashed, and white lines represent approximate border lines for which there may not yet be full agreement. (Courtesy of the Global TB Programme, WHO; with permission.)

cases were diagnosed because of a lack of culture and drug-susceptibility testing capacity in most settings worldwide. The countries of the former Soviet Union have reported the highest proportions of MDR disease among new TB cases (up to 35–40% in some regions of Russia and Belarus). Overall, 60% of all MDR-TB cases occur in

China, India, the Russian Federation, Pakistan, and Ukraine. Since 2006, 100 countries, including the United States, have reported cases of extensively drug-resistant TB (XDR-TB), in which MDR-TB is compounded by additional resistance to the most powerful second-line anti-TB drugs (fluoroquinolones and at least one of the injectable

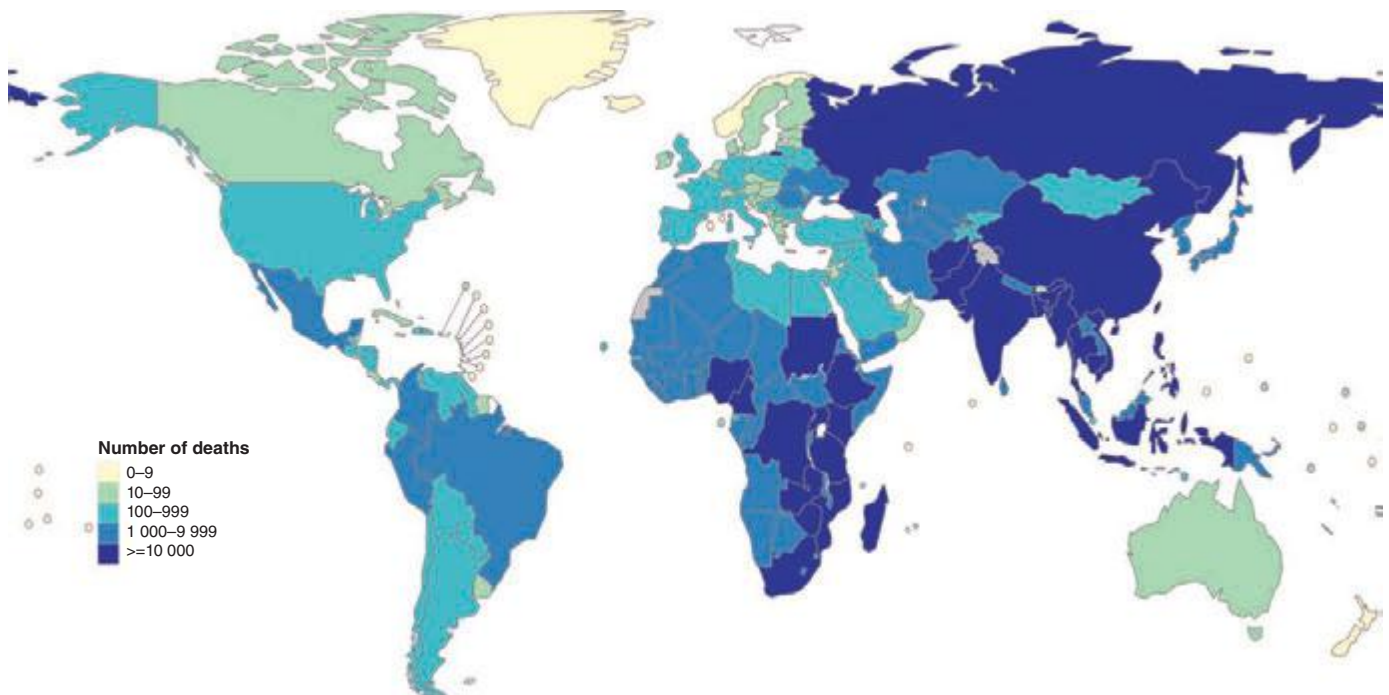


FIGURE 202-3 Estimated numbers of tuberculosis-related deaths in 2013. (See disclaimer in Fig. 202-2. Courtesy of the Global TB Programme, WHO; with permission.)

drugs amikacin, kanamycin, and capreomycin). Up to 10% of the MDR-TB cases worldwide may actually be XDR-TB, but the vast majority of XDR-TB cases remain undiagnosed because reliable methods for drug susceptibility testing are lacking and laboratory capacity is limited. Lately, cases deemed resistant to all anti-TB drugs have been reported from countries such as India, Italy, and Iran; however, this information must be interpreted with caution because drug susceptibility testing for several second-line drugs is neither accurate nor reproducible.

FROM EXPOSURE TO INFECTION

M. tuberculosis is most commonly transmitted from a person with infectious pulmonary TB by droplet nuclei, which are aerosolized by coughing, sneezing, or speaking. The tiny droplets dry rapidly; the smallest (<5–10 µm in diameter) may remain suspended in the air for several hours and may reach the terminal air passages when inhaled. There may be as many as 3000 infectious nuclei per cough. Other routes of transmission of tubercle bacilli (e.g., through the skin or the placenta) are uncommon and of no epidemiologic significance. The probability of contact with a person who has an infectious form of TB, the intimacy and duration of that contact, the degree of infectiousness of the case, and the shared environment in which the contact takes place are all important determinants of the likelihood of transmission. Several studies of close-contact situations have clearly demonstrated that TB patients whose sputum contains AFB visible by microscopy (sputum smear-positive cases) are the most likely to transmit the infection. The most infectious patients have cavitary pulmonary disease or, much less commonly, laryngeal TB and produce sputum containing as many as 10^5 – 10^7 AFB/mL. Patients with sputum smear-negative/culture-positive TB are less infectious, although they have been responsible for up to 20% of transmission in some studies in the United States. Those with culture-negative pulmonary TB and extrapulmonary TB are essentially noninfectious. Because persons with both HIV infection and TB are less likely to have cavitations, they may be less infectious than persons without HIV co-infection. Crowding in poorly ventilated rooms is one of the most important factors in the transmission of tubercle bacilli because it increases the intensity of contact with a case.

The risk of acquiring *M. tuberculosis* infection is determined mainly by exogenous factors. Because of delays in seeking care and in making a diagnosis, it is generally estimated that, in high-prevalence settings, up to 20 contacts may be infected by each AFB-positive case before the index case is diagnosed.

FROM INFECTION TO DISEASE

Unlike the risk of acquiring infection with *M. tuberculosis*, the risk of developing disease after being infected depends largely on endogenous factors, such as the individual's innate immunologic and nonimmunologic defenses and the level at which the individual's cell-mediated immunity (CMI) is functioning. Clinical illness directly following infection is classified as *primary TB* and is common among children in the first few years of life and among immunocompromised persons. Although primary TB may be severe and disseminated, it generally is not associated with high-level transmissibility. When infection is acquired later in life, the chance is greater that the mature immune system will contain it at least temporarily. Bacilli, however, may persist for years before reactivating to produce *secondary* (or *postprimary*) TB, which, because of frequent cavitation, is more often infectious than is primary disease. Overall, it is estimated that up to 10% of infected persons will eventually develop active TB in their lifetime—half of them during the first 18 months after infection. The risk is much higher among HIV-infected persons. Reinfection of a previously infected individual, which is common in areas with high rates of TB transmission, may also favor the development of disease. At the height of the TB resurgence in the United States in the early 1990s, molecular typing and comparison of strains of *M. tuberculosis* suggested that up to one-third of cases of active TB in some inner-city communities were due to recent transmission rather than to reactivation of old latent infection. Age is an important determinant of the risk of disease after infection. Among infected persons, the incidence of TB is highest

TABLE 202-1 RISK FACTORS FOR ACTIVE TUBERCULOSIS IN PERSONS WHO HAVE BEEN INFECTED WITH TUBERCLE BACILLI

Factor	Relative Risk/Odds ^a
Recent infection (<1 year)	12.9
Fibrotic lesions (spontaneously healed)	2–20
Comorbidities and iatrogenic causes	
HIV infection	21–>30
Silicosis	30
Chronic renal failure/hemodialysis	10–25
Diabetes	2–4
IV drug use	10–30
Immunosuppressive treatment	10
Tumor necrosis factor α inhibitors	4–5
Gastrectomy	2–5
Jejunioileal bypass	30–60
Posttransplantation period (renal, cardiac)	20–70
Tobacco smoking	2–3
Malnutrition and severe underweight	2

^aOld infection = 1.

during late adolescence and early adulthood; the reasons are unclear. The incidence among women peaks at 25–34 years of age. In this age group, rates among women may be higher than those among men, whereas at older ages the opposite is true. The risk increases in the elderly, possibly because of waning immunity and comorbidity.

A variety of diseases and conditions favor the development of active TB (Table 202-1). In absolute terms, the most potent risk factor for TB among infected individuals is clearly HIV co-infection, which suppresses cellular immunity. The risk that LTBI will proceed to active disease is directly related to the patient's degree of immunosuppression. In a study of HIV-infected, tuberculin skin test (TST)-positive persons, this risk varied from 2.6 to 13.3 cases/100 person-years and increased as the CD4+ T cell count decreased.

NATURAL HISTORY OF DISEASE

Studies conducted in various countries before the advent of chemotherapy showed that untreated TB is often fatal. About one-third of patients died within 1 year after diagnosis, and more than 50% died within 5 years. The 5-year mortality rate among sputum smear-positive cases was 65%. Of the survivors at 5 years, ~60% had undergone spontaneous remission, while the remainder were still excreting tubercle bacilli. With effective, timely, and proper chemotherapy, patients have a very high chance of being cured. However, improper use of anti-TB drugs, while reducing mortality rates, may also result in large numbers of chronic infectious cases, often with drug-resistant bacilli.

PATHOGENESIS AND IMMUNITY

INFECTION AND MACROPHAGE INVASION

The interaction of *M. tuberculosis* with the human host begins when droplet nuclei containing viable microorganisms propelled into the air by infectious patients are inhaled by a close bystander. Although the majority of inhaled bacilli are trapped in the upper airways and expelled by ciliated mucosal cells, a fraction (usually <10%) reach the alveoli, a unique immunoregulatory environment. There, alveolar macrophages that have not yet been activated (prototypic alternatively activated macrophages) phagocytose the bacilli. Adhesion of mycobacteria to macrophages results largely from binding of the bacterial cell wall to a variety of macrophage cell-surface molecules, including complement receptors, the mannose receptor, the immunoglobulin G Fc γ receptor, and type A scavenger receptors. Phagocytosis is enhanced by complement activation leading to opsonization of bacilli with C3 activation products such as C3b and C3bi. (Bacilli are resistant to complement-mediated lysis.) Binding of certain receptors, such as the mannose receptor, regulates

1106 postphagocytic events such as phagosome–lysosome fusion and inflammatory cytokine production. After a phagosome forms, the survival of *M. tuberculosis* within it seems to depend in part on reduced acidification due to lack of assembly of a complete vesicular proton-adenosine triphosphatase. A complex series of events is generated by the bacterial cell-wall lipoglycan lipoarabinomannan (ManLAM). ManLAM inhibits the intracellular increase of Ca²⁺. Thus, the Ca²⁺/calmodulin pathway (leading to phagosome–lysosome fusion) is impaired, and the bacilli survive within the phagosomes. The *M. tuberculosis* phagosome has been found to inhibit the production of phosphatidylinositol 3-phosphate (PI3P). Normally, PI3P earmarks phagosomes for membrane sorting and maturation, including phagolysosome formation, which would destroy the bacteria. Bacterial factors have also been found to block the host defense of autophagy, in which the cell sequesters the phagosome in a double-membrane vesicle (*autophagosome*) that is destined to fuse with lysosomes. If the bacilli are successful in arresting phagosome maturation, then replication begins and the macrophage eventually ruptures and releases its bacillary contents. Other uninfected phagocytic cells are then recruited to continue the infection cycle by ingesting dying macrophages and their bacillary content, thus in turn becoming infected themselves and expanding the infection.

VIRULENCE OF TUBERCLE BACILLI



M. tuberculosis must be viewed as a complex formed by a multitude of strains that differ in virulence and are capable of producing a variety of manifestations of disease. Since the elucidation of the *M. tuberculosis* genome in 1998, large mutant collections have been generated, and many bacterial genes that contribute to *M. tuberculosis* virulence have been found. Different patterns of virulence defects have been defined in various animal models—predominantly mice but also guinea pigs, rabbits, and nonhuman primates. The *katG* gene encodes for a catalase/oxidase enzyme that protects against oxidative stress and is required for isoniazid activation and subsequent bactericidal activity. Region of difference 1 (RD1) is a 9.5-kb locus that encodes two key small protein antigens—early secretory antigen-6 (ESAT-6) and culture filtrate protein-10 (CFP-10)—as well as a putative secretion apparatus that may facilitate their egress; the absence of this locus in the vaccine strain *M. bovis* bacille Calmette-Guérin (BCG) has been shown to be a key attenuating mutation. The validity of a recent observation in *M. marinum* needs to be confirmed in *M. tuberculosis*; in *M. marinum*, a mutation in the RD1 virulence locus encoding the ESX1 secretion system impairs the capacity of apoptotic macrophages to recruit uninfected cells for further rounds of infection. The results are less replication and fewer new granulomas. Mutants lacking key enzymes of bacterial biosynthesis become auxotrophic for the missing substrate and often are totally unable to proliferate in animals; these include the *leuCD* and *panCD* mutants, which require leucine and pantothenic acid, respectively. The isocitrate lyase gene *icl1* encodes a key step in the glyoxylate shunt that facilitates bacterial growth on fatty acid substrates; this gene is required for long-term persistence of *M. tuberculosis* infection in mice with chronic TB. *M. tuberculosis* mutants in regulatory genes such as sigma factor C and sigma factor H (*sigC* and *sigH*) are associated with normal bacterial growth in mice, but they fail to elicit full tissue pathology. Finally, the mycobacterial protein CarD (expressed by the *carD* gene) seems essential for the control of rRNA transcription that is required for replication and persistence in the host cell. Its loss exposes mycobacteria to oxidative stress, starvation, DNA damage, and ultimately sensitivity to killing by a variety of host mutagens and defensive mechanisms.

INNATE RESISTANCE TO INFECTION



Several observations suggest that genetic factors play a key role in innate nonimmune resistance to infection with *M. tuberculosis* and the development of disease. The existence of this resistance, which is polygenic in nature, is suggested by the differing degrees of susceptibility to TB in different populations. In mice, a gene called *Nramp1* (natural resistance-associated macrophage protein 1) plays a regulatory role in resistance/susceptibility to mycobacteria. The human homologue NRAMP1, which maps to chromosome 2q, may play a role in determining susceptibility to TB, as is suggested by a study

among West Africans. Studies of mouse genetics identified a novel host resistance gene, *ipr1*, which is encoded within the *sst1* locus; *ipr1* encodes an interferon (IFN)-inducible nuclear protein that interacts with other nuclear proteins in macrophages primed with IFNs or infected by *M. tuberculosis*. In addition, polymorphisms in multiple genes, such as those encoding for various major histocompatibility complex (MHC) alleles, IFN- γ , T cell growth factor β , interleukin (IL) 10, mannose-binding protein, IFN- γ receptor, Toll-like receptor 2, vitamin D receptor, and IL-1, have been associated with susceptibility to TB.

THE HOST RESPONSE, GRANULOMA FORMATION, AND “LATENCY”

In the initial stage of host–bacterium interaction, prior to the onset of an acquired CMI response, *M. tuberculosis* disseminates widely through the lymph vessels, spreading to other sites in the lungs and other organs, and undergoes a period of extensive growth within naïve unactivated macrophages; additional naïve macrophages are recruited to the early granuloma. Studies suggest that *M. tuberculosis* uses specific virulence mechanisms to subvert host cellular signaling and to elicit an early regulated proinflammatory response that promotes granuloma expansion and bacterial growth during this key early phase. A study of *M. marinum* infection in zebrafish has delineated one molecular mechanism by which mycobacteria induce granuloma formation. The mycobacterial protein ESAT-6 induces secretion of matrix metalloproteinase 9 (MMP9) by nearby epithelial cells that are in contact with infected macrophages. MMP9 in turn stimulates recruitment of naïve macrophages, thus inducing granuloma maturation and bacterial growth. Disruption of MMP9 function results in reduced bacterial growth. Another study has shown that *M. tuberculosis*-derived cyclic AMP is secreted from the phagosome into host macrophages, subverting the cell's signal transduction pathways and stimulating an elevation in the secretion of tumor necrosis factor α (TNF- α) as well as further proinflammatory cell recruitment. Ultimately, the chemoattractants and bacterial products released during the repeated rounds of cell lysis and infection of newly arriving macrophages enable dendritic cells to access bacilli; these cells migrate to the draining lymph nodes and present mycobacterial antigens to T lymphocytes. At this point, the development of CMI and humoral immunity begins. These initial stages of infection are usually asymptomatic.

About 2–4 weeks after infection, two host responses to *M. tuberculosis* develop: a macrophage-activating CMI response and a tissue-damaging response. The *macrophage-activating response* is a T cell-mediated phenomenon resulting in the activation of macrophages that are capable of killing and digesting tubercle bacilli. The *tissue-damaging response* is the result of a delayed-type hypersensitivity (DTH) reaction to various bacillary antigens; it destroys unactivated macrophages that contain multiplying bacilli but also causes caseous necrosis of the involved tissues (see below). Although both of these responses can inhibit mycobacterial growth, it is the balance between the two that determines the forms of TB that will develop subsequently. With the development of specific immunity and the accumulation of large numbers of activated macrophages at the site of the primary lesion, granulomatous lesions (tubercles) are formed. These lesions consist of accumulations of lymphocytes and activated macrophages that evolve toward epithelioid and giant cell morphologies. Initially, the tissue-damaging response can limit mycobacterial growth within macrophages. As stated above, this response, mediated by various bacterial products, not only destroys macrophages but also produces early solid necrosis in the center of the tubercle. Although *M. tuberculosis* can survive, its growth is inhibited within this necrotic environment by low oxygen tension and low pH. At this point, some lesions may heal by fibrosis, with subsequent calcification, whereas inflammation and necrosis occur in other lesions. Some observations have challenged the traditional view that any encounter between mycobacteria and macrophages results in chronic infection. It is possible that an immune response capable of eradicating early infection may sometimes develop as a consequence, for instance, of disabling mutations in mycobacterial genomes rendering their replication ineffective. Individual granulomas that are formed during this phase of infection can vary in size and cell composition; some can contain the spread of mycobacteria, while others cannot. LTBI ensues as a result of this dynamic balance between the

microorganism and the host. According to recent developments, *latency* may not be an accurate term because bacilli may remain active during this “latent” stage, forming biofilms in necrotic areas within which they temporarily hide. Thus, the term *persist* is probably more accurate to indicate the behavior of the bacilli in this phase. It is important to recognize that latent infection and disease represent not a binary state but rather a continuum along which infection will eventually move in the direction of full containment or disease. The ability to predict, through systemic biomarkers, which affected individuals will progress toward disease would be of immense value in devising prophylactic interventions.

MACROPHAGE-ACTIVATING RESPONSE

CMI is critical at this early stage. In the majority of infected individuals, local macrophages are activated when bacillary antigens processed by macrophages stimulate T lymphocytes to release a variety of lymphokines. These activated macrophages aggregate around the lesion’s center and effectively neutralize tubercle bacilli without causing further tissue destruction. In the central part of the lesion, the necrotic material resembles soft cheese (*caseous necrosis*)—a phenomenon that may also be observed in other conditions, such as neoplasms. Even when healing takes place, viable bacilli may remain dormant within macrophages or in the necrotic material for many years. These “healed” lesions in the lung parenchyma and hilar lymph nodes may later undergo calcification.

DELAYED-TYPE HYPERSENSITIVITY

In a minority of cases, the macrophage-activating response is weak, and mycobacterial growth can be inhibited only by intensified DTH reactions, which lead to lung tissue destruction. The lesion tends to enlarge further, and the surrounding tissue is progressively damaged. At the center of the lesion, the caseous material liquefies. Bronchial walls and blood vessels are invaded and destroyed, and cavities are formed. The liquefied caseous material, containing large numbers of bacilli, is drained through bronchi. Within the cavity, tubercle bacilli multiply, spill into the airways, and are discharged into the environment through expiratory maneuvers such as coughing and talking. In the early stages of infection, bacilli are usually transported by macrophages to regional lymph nodes, from which they gain access to the central venous return; from there they reseed the lungs and may also disseminate beyond the pulmonary vasculature throughout the body via the systemic circulation. The resulting extrapulmonary lesions may undergo the same evolution as those in the lungs, although most tend to heal. In young children with poor natural immunity, hematogenous dissemination may result in highly fatal miliary TB or tuberculous meningitis.

ROLE OF MACROPHAGES AND MONOCYTES

While CMI confers partial protection against *M. tuberculosis*, humoral immunity plays a less well-defined role in protection (although evidence is accumulating on the existence of antibodies to lipoarabinomannan, which may prevent dissemination of infection in children). In the case of CMI, two types of cells are essential: macrophages, which directly phagocytose tubercle bacilli, and T cells (mainly CD4+ T lymphocytes), which induce protection through the production of cytokines, especially IFN- γ . After infection with *M. tuberculosis*, alveolar macrophages secrete various cytokines responsible for a number of events (e.g., the formation of granulomas) as well as systemic effects (e.g., fever and weight loss). However, alternatively activated alveolar macrophages may be particularly susceptible to *M. tuberculosis* growth early on, given their more limited proinflammatory and bactericidal activity, which is related in part to being bathed in surfactant. New monocytes and macrophages attracted to the site are key components of the immune response. Their primary mechanism is probably related to production of oxidants (such as reactive oxygen intermediates or nitric oxide) that have antimycobacterial activity and increase the synthesis of cytokines such as TNF- α and IL-1, which in turn regulate the release of reactive oxygen intermediates and reactive nitrogen intermediates. In addition, macrophages can undergo apoptosis—a defensive mechanism to prevent release of cytokines and bacilli via their sequestration in the apoptotic cell. Recent work also describes the

involvement of neutrophils in the host response, although the timing of their appearance and their effectiveness remain uncertain.

ROLE OF T LYMPHOCYTES

Alveolar macrophages, monocytes, and dendritic cells are also critical in processing and presenting antigens to T lymphocytes, primarily CD4+ and CD8+ T cells; the result is the activation and proliferation of CD4+ T lymphocytes, which are crucial to the host’s defense against *M. tuberculosis*. Qualitative and quantitative defects of CD4+ T cells explain the inability of HIV-infected individuals to contain mycobacterial proliferation. Activated CD4+ T lymphocytes can differentiate into cytokine-producing T_H1 or T_H2 cells. T_H1 cells produce IFN- γ —an activator of macrophages and monocytes—and IL-2. T_H2 cells produce IL-4, IL-5, IL-10, and IL-13 and may also promote humoral immunity. The interplay of these various cytokines and their cross-regulation determine the host’s response. The role of cytokines in promoting intracellular killing of mycobacteria, however, has not been entirely elucidated. IFN- γ may induce the generation of reactive nitrogen intermediates and regulate genes involved in bactericidal effects. TNF- α also seems to be important. Observations made originally in transgenic knockout mice and more recently in humans suggest that other T cell subsets, especially CD8+ T cells, may play an important role. CD8+ T cells have been associated with protective activities via cytotoxic responses and lysis of infected cells as well as with production of IFN- γ and TNF- α . Finally, natural killer cells act as co-regulators of CD8+ T cell lytic activities, and $\gamma\delta$ T cells are increasingly thought to be involved in protective responses in humans.

MYCOBACTERIAL LIPIDS AND PROTEINS

Lipids have been involved in mycobacterial recognition by the innate immune system, and lipoproteins (such as 19-kDa lipoprotein) have been proven to trigger potent signals through Toll-like receptors present in blood dendritic cells. *M. tuberculosis* possesses various protein antigens. Some are present in the cytoplasm and cell wall; others are secreted. That the latter are more important in eliciting a T lymphocyte response is suggested by experiments documenting the appearance of protective immunity in animals after immunization with live, protein-secreting mycobacteria. Among the antigens that may play a protective role are the 30-kDa (or 85B) and ESAT-6 antigens. Protective immunity is probably the result of reactivity to many different mycobacterial antigens. These antigens are being incorporated into newly designed vaccines on various platforms.

SKIN TEST REACTIVITY

Coincident with the appearance of immunity, DTH to *M. tuberculosis* develops. This reactivity is the basis of the TST, which is used primarily for the detection of *M. tuberculosis* infection in persons without symptoms. The cellular mechanisms responsible for TST reactivity are related mainly to previously sensitized CD4+ T lymphocytes, which are attracted to the skin-test site. There, they proliferate and produce cytokines. Although DTH is associated with protective immunity (TST-positive persons are less susceptible to a new *M. tuberculosis* infection than TST-negative persons), it by no means guarantees protection against reactivation. In fact, cases of active TB are often accompanied by strongly positive skin-test reactions. There is also evidence of reinfection with a new strain of *M. tuberculosis* in patients previously treated for active disease. This evidence underscores the fact that previous latent or active TB may not confer fully protective immunity.

CLINICAL MANIFESTATIONS

TB is classified as pulmonary, extrapulmonary, or both. Depending on several factors linked to different populations and bacterial strains, extrapulmonary TB may occur in 10–40% of patients. Furthermore, up to two-thirds of HIV-infected patients with TB may have both pulmonary and extrapulmonary TB or extrapulmonary TB alone.

PULMONARY TB

Pulmonary TB is conventionally categorized as primary or postprimary (adult-type, secondary). This distinction has been challenged

1108 by molecular evidence from TB-endemic areas indicating that a large percentage of cases of adult pulmonary TB result from recent infection (either primary infection or reinfection) and not from reactivation.

Primary Disease Primary pulmonary TB occurs soon after the initial infection with tubercle bacilli. It may be asymptomatic or may present with fever and occasionally pleuritic chest pain. In areas of high TB transmission, this form of disease is often seen in children. Because most inspired air is distributed to the middle and lower lung zones, these areas are most commonly involved in primary TB. The lesion forming after initial infection (*Ghon focus*) is usually peripheral and accompanied by transient hilar or paratracheal lymphadenopathy, which may or may not be visible on standard chest radiography (Fig. 202-4). Some patients develop erythema nodosum on the legs (see Fig. 25e-40) or phlyctenular conjunctivitis. In the majority of cases, the lesion heals spontaneously and becomes evident only as a small calcified nodule. Pleural reaction overlying a subpleural focus is also common. The Ghon focus, with or without overlying pleural reaction, thickening, and regional lymphadenopathy, is referred to as the *Ghon complex*.

In young children with immature CMI and in persons with impaired immunity (e.g., those with malnutrition or HIV infection), primary pulmonary TB may progress rapidly to clinical illness. The initial lesion increases in size and can evolve in different ways. Pleural effusion, which is found in up to two-thirds of cases, results from the penetration of bacilli into the pleural space from an adjacent subpleural focus. In severe cases, the primary site rapidly enlarges, its central portion undergoes necrosis, and cavitation develops (*progressive primary TB*). TB in young children is almost invariably accompanied by hilar or paratracheal lymphadenopathy due to the spread of bacilli from the lung parenchyma through lymphatic vessels. Enlarged lymph nodes may compress bronchi, causing total obstruction with distal collapse, partial obstruction with large-airway wheezing, or a ball-valve effect with segmental/lobar hyperinflation. Lymph nodes may also rupture into the airway with development of pneumonia, often including areas of necrosis and cavitation, distal to the obstruction. Bronchiectasis (Chap. 312) may develop in any segment/lobe damaged by progressive caseating pneumonia. Occult hematogenous dissemination commonly follows primary infection. However, in the absence of a sufficient acquired immune response, which usually contains the infection, disseminated or miliary disease may result (Fig. 202-5). Small granulomatous lesions develop in multiple organs and may cause



FIGURE 202-4 Chest radiograph showing right hilar lymph node enlargement with infiltration into the surrounding lung tissue in a child with primary tuberculosis. (Courtesy of Prof. Robert Gie, Department of Paediatrics and Child Health, Stellenbosch University, South Africa; with permission.)



FIGURE 202-5 Chest radiograph showing bilateral miliary (millet-sized) infiltrates in a child. (Courtesy of Prof. Robert Gie, Department of Paediatrics and Child Health, Stellenbosch University, South Africa; with permission.)

locally progressive disease or result in tuberculous meningitis; this is a particular concern in very young children and immunocompromised persons (e.g., patients with HIV infection).

Postprimary (Adult-Type) Disease Also referred to as *reactivation* or *secondary TB*, postprimary TB is probably most accurately termed *adult-type TB* because it may result from endogenous reactivation of distant LTBI or recent infection (primary infection or reinfection). It is usually localized to the apical and posterior segments of the upper lobes, where the substantially higher mean oxygen tension (compared with that in the lower zones) favors mycobacterial growth. The superior segments of the lower lobes are also more frequently involved. The extent of lung parenchymal involvement varies greatly, from small infiltrates to extensive cavitory disease. With cavity formation, liquefied necrotic contents are ultimately discharged into the airways and may undergo bronchogenic spread, resulting in satellite lesions within the lungs that may in turn undergo cavitation (Figs. 202-6 and 202-7). Massive involvement of pulmonary segments or lobes, with coalescence of lesions, produces caseating pneumonia. While up to one-third of untreated patients reportedly succumb to severe pulmonary TB within a few months after onset (the classic “galloping consumption” of the past), others may undergo a process of spontaneous remission or proceed along a chronic, progressively debilitating course (“consumption” or *phthisis*). Under these circumstances, some pulmonary lesions become fibrotic and may later calcify, but cavities persist in other parts of the lungs. Individuals with such chronic disease continue to discharge tubercle bacilli into the environment. Most patients respond to treatment, with defervescence, decreasing cough, weight gain, and a general improvement in well-being within several weeks.

Early in the course of disease, symptoms and signs are often non-specific and insidious, consisting mainly of diurnal fever and night sweats due to defervescence, weight loss, anorexia, general malaise, and weakness. However, in up to 90% of cases, cough eventually develops—often initially nonproductive and limited to the morning and subsequently accompanied by the production of purulent sputum, sometimes with blood streaking. Hemoptysis develops in 20–30% of cases, and massive hemoptysis may ensue as a consequence of the erosion of a blood vessel in the wall of a cavity. Hemoptysis, however, may also result from rupture of a dilated vessel in a cavity (*Rasmussen’s aneurysm*) or from aspergilloma formation in an old cavity. Pleuritic chest pain sometimes develops in patients with subpleural parenchymal lesions or pleural disease. Extensive disease may produce dyspnea



FIGURE 202-6 Chest radiograph showing a right-upper-lobe infiltrate and a cavity with an air-fluid level in a patient with active tuberculosis. (Courtesy of Dr. Andrea Gori, Department of Infectious Diseases, S. Paolo University Hospital, Milan, Italy; with permission.)

and, in rare instances, adult respiratory distress syndrome. Physical findings are of limited use in pulmonary TB. Many patients have no abnormalities detectable by chest examination, whereas others have detectable rales in the involved areas during inspiration, especially after coughing. Occasionally, rhonchi due to partial bronchial obstruction and classic amphoric breath sounds in areas with large cavities may be heard. Systemic features include fever (often low-grade and intermittent) in up to 80% of cases and wasting. Absence of fever, however, does not exclude TB. In some cases, pallor and finger clubbing develop. The most common hematologic findings are mild anemia, leukocytosis, and thrombocytosis with a slightly elevated erythrocyte sedimentation rate and/or C-reactive protein level. None of these findings is consistent or sufficiently accurate for diagnostic purposes.

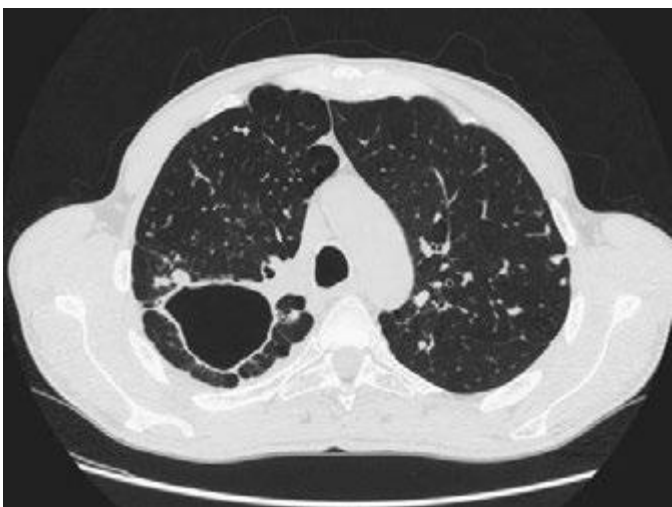


FIGURE 202-7 CT scan showing a large cavity in the right lung of a patient with active tuberculosis. (Courtesy of Dr. Elisa Busi Rizzi, National Institute for Infectious Diseases, Spallanzani Hospital, Rome, Italy; with permission.)

EXTRAPULMONARY TB

In order of frequency, the extrapulmonary sites most commonly involved in TB are the lymph nodes, pleura, genitourinary tract, bones and joints, meninges, peritoneum, and pericardium. However, virtually all organ systems may be affected. As a result of hematogenous dissemination in HIV-infected individuals, extrapulmonary TB is seen more commonly today than in the past in settings of high HIV prevalence.

Lymph Node TB (Tuberculous Lymphadenitis) The most common presentation of extrapulmonary TB in both HIV-seronegative and HIV-infected patients (35% of cases worldwide and more than 40% of cases in the United States in recent series), lymph node disease is particularly frequent among HIV-infected patients and among children (Fig. 202-8). In the United States, besides children, women (particularly non-Caucasians) seem to be especially susceptible. Once caused mainly by *M. bovis*, tuberculous lymphadenitis today is due largely to *M. tuberculosis*. Lymph node TB presents as painless swelling of the lymph nodes, most commonly at posterior cervical and supraclavicular sites (a condition historically referred to as *scrofula*). Lymph nodes are usually discrete in early disease but develop into a matted nontender mass over time and may result in a fistulous tract draining caseous material. Associated pulmonary disease is present in fewer than 50% of cases, and systemic symptoms are uncommon except in HIV-infected patients. The diagnosis is established by fine-needle aspiration biopsy (with a yield of up to 80%) or surgical excision biopsy. Bacteriologic confirmation is achieved in the vast majority of cases, granulomatous lesions with or without visible AFBs are typically seen, and cultures are positive in 70–80% of cases. Among HIV-infected patients, granulomas are less well organized and are frequently absent entirely, but bacterial loads are heavier than in HIV-seronegative patients, with higher yields from microscopy and culture. Differential diagnosis includes a variety of infectious conditions, neoplastic diseases such as lymphomas or metastatic carcinomas, and rare disorders like Kikuchi's disease (necrotizing histiocytic lymphadenitis), Kimura's disease, and Castleman's disease.

Pleural TB Involvement of the pleura accounts for ~20% of extrapulmonary cases in the United States and elsewhere. Isolated pleural effusion usually reflects recent primary infection, and the collection of fluid in the pleural space represents a hypersensitivity response to mycobacterial antigens. Pleural disease may also result from contiguous parenchymal spread, as in many cases of pleurisy accompanying



FIGURE 202-8 Tuberculous lymphadenitis affecting the cervical lymph nodes in a 2-year-old child from Malawi. (Courtesy of Prof. S. Graham, Centre for International Child Health, University of Melbourne, Australia; with permission.)

1110 postprimary disease. Depending on the extent of reactivity, the effusion may be small, remain unnoticed, and resolve spontaneously or may be sufficiently large to cause symptoms such as fever, pleuritic chest pain, and dyspnea. Physical findings are those of pleural effusion: dullness to percussion and absence of breath sounds. A chest radiograph reveals the effusion and, in up to one-third of cases, also shows a parenchymal lesion. Thoracentesis is required to ascertain the nature of the effusion and to differentiate it from manifestations of other etiologies. The fluid is straw colored and at times hemorrhagic; it is an exudate with a protein concentration >50% of that in serum (usually ~4–6 g/dL), a normal to low glucose concentration, a pH of ~7.3 (occasionally <7.2), and detectable white blood cells (usually 500–6000/μL). Neutrophils may predominate in the early stage, but lymphocyte predominance is the typical finding later. Mesothelial cells are generally rare or absent. AFB are rarely seen on direct smear, and cultures often may be falsely negative for *M. tuberculosis*; positive cultures are more common among postprimary cases. Determination of the pleural concentration of adenosine deaminase (ADA) may be a useful screening test, and TB may be excluded if the value is very low. Lysozyme is also present in the pleural effusion. Measurement of IFN-γ, either directly or through stimulation of sensitized T cells with mycobacterial antigens, can be helpful. Needle biopsy of the pleura is often required for diagnosis and is recommended over pleural fluid; it reveals granulomas and/or yields a positive culture in up to 80% of cases. Pleural biopsy can yield a positive result in ~75% of cases when real-time automated nucleic acid amplification is used (the Xpert® MTB/RIF assay [Cepheid, Sunnyvale, CA]; see “Nucleic Acid Amplification Technology,” below), although pleural fluid testing with this assay is not recommended because of low sensitivity. This form of pleural TB responds rapidly to chemotherapy and may resolve spontaneously. Concurrent glucocorticoid administration may reduce the duration of fever and/or chest pain but is not of proven benefit.

Tuberculous empyema is a less common complication of pulmonary TB. It is usually the result of the rupture of a cavity, with spillage of a large number of organisms into the pleural space. This process may create a bronchopleural fistula with evident air in the pleural space. A chest radiograph shows hydropneumothorax with an air-fluid level. The pleural fluid is purulent and thick and contains large numbers of lymphocytes. Acid-fast smears and mycobacterial cultures are often positive. Surgical drainage is usually required as an adjunct to chemotherapy. Tuberculous empyema may result in severe pleural fibrosis and restrictive lung disease. Removal of the thickened visceral pleura (*decortication*) is occasionally necessary to improve lung function.

TB of the Upper Airways Nearly always a complication of advanced cavity pulmonary TB, TB of the upper airways may involve the larynx, pharynx, and epiglottis. Symptoms include hoarseness, dysphonia, and dysphagia in addition to chronic productive cough. Findings depend on the site of involvement, and ulcerations may be seen on laryngoscopy. Acid-fast smear of the sputum is often positive, but biopsy may be necessary in some cases to establish the diagnosis. Carcinoma of the larynx may have similar features but is usually painless.

Genitourinary TB Genitourinary TB, which accounts for ~10–15% of all extrapulmonary cases in the United States and elsewhere, may involve any portion of the genitourinary tract. Local symptoms predominate, and up to 75% of patients have chest radiographic abnormalities suggesting previous or concomitant pulmonary disease. Urinary frequency, dysuria, nocturia, hematuria, and flank or abdominal pain are common presentations. However, patients may be asymptomatic and their disease discovered only after severe destructive lesions of the kidneys have developed. Urinalysis gives abnormal results in 90% of cases, revealing pyuria and hematuria. The documentation of culture-negative pyuria in acidic urine should raise the suspicion of TB. IV pyelography, abdominal computed tomography (CT), or magnetic resonance imaging (MRI) (Fig. 202-9) may show deformities and obstructions; calcifications and ureteral strictures are suggestive findings. Culture of three morning urine specimens yields a definitive diagnosis in nearly 90% of cases. Severe ureteral strictures may lead to hydronephrosis and renal damage. Genital TB is

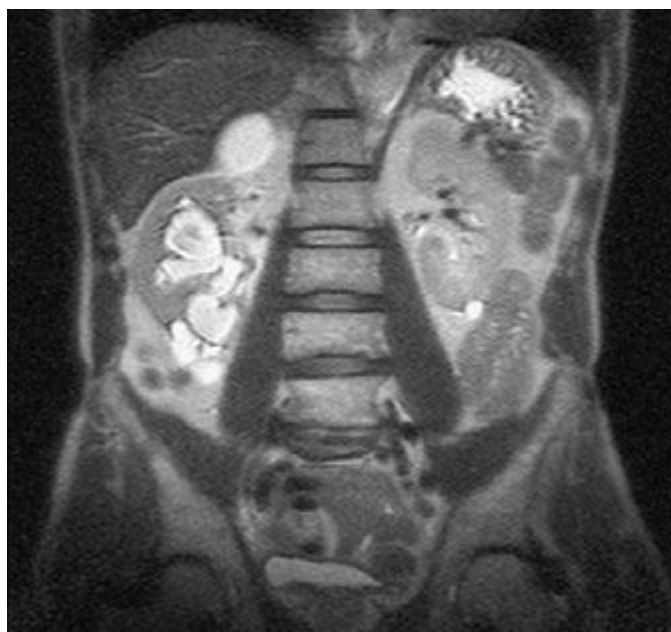


FIGURE 202-9 MRI of culture-confirmed renal tuberculosis. T2-weighted coronal plane: coronal sections showing several renal lesions in both the cortical and the medullary tissues of the right kidney. (Courtesy of Dr. Alberto Matteelli, Department of Infectious Diseases, University of Brescia, Italy; with permission.)

diagnosed more commonly in female than in male patients. In female patients, it affects the fallopian tubes and the endometrium and may cause infertility, pelvic pain, and menstrual abnormalities. Diagnosis requires biopsy or culture of specimens obtained by dilation and curettage. In male patients, genital TB preferentially affects the epididymis, producing a slightly tender mass that may drain externally through a fistulous tract; orchitis and prostatitis may also develop. In almost half of cases of genitourinary TB, urinary tract disease is also present. Genitourinary TB responds well to chemotherapy.

Skeletal TB In the United States, TB of the bones and joints is responsible for ~10% of extrapulmonary cases. In bone and joint disease, pathogenesis is related to reactivation of hematogenous foci or to spread from adjacent paravertebral lymph nodes. Weight-bearing joints (the spine in 40% of cases, the hips in 13%, and the knees in 10%) are most commonly affected. Spinal TB (Pott's disease or tuberculous spondylitis; Fig. 202-10) often involves two or more adjacent vertebral bodies. Whereas the upper thoracic spine is the most common site of spinal TB in children, the lower thoracic and upper lumbar vertebrae are usually affected in adults. From the anterior superior or inferior angle of the vertebral body, the lesion slowly reaches the adjacent body, later affecting the intervertebral disk. With advanced disease, collapse of vertebral bodies results in kyphosis (*gibbus*). A paravertebral “cold” abscess may also form. In the upper spine, this abscess may track to and penetrate the chest wall, presenting as a soft tissue mass; in the lower spine, it may reach the inguinal ligaments or present as a psoas abscess. CT or MRI reveals the characteristic lesion and suggests its etiology. The differential diagnosis includes tumors and other infections. Pyogenic bacterial osteomyelitis, in particular, involves the disk very early and produces rapid sclerosis. Aspiration of the abscess or bone biopsy confirms the tuberculous etiology, as cultures are usually positive and histologic findings highly typical. A catastrophic complication of Pott's disease is paraplegia, which is usually due to an abscess or a lesion compressing the spinal cord. Paraparesis due to a large abscess is a medical emergency and requires rapid drainage. TB of the hip joints, usually involving the head of the femur, causes pain; TB of the knee produces pain and swelling. If the disease goes unrecognized, the joints may be destroyed. Diagnosis requires examination of the synovial fluid, which is thick in appearance, with a high protein concentration and a variable cell count.

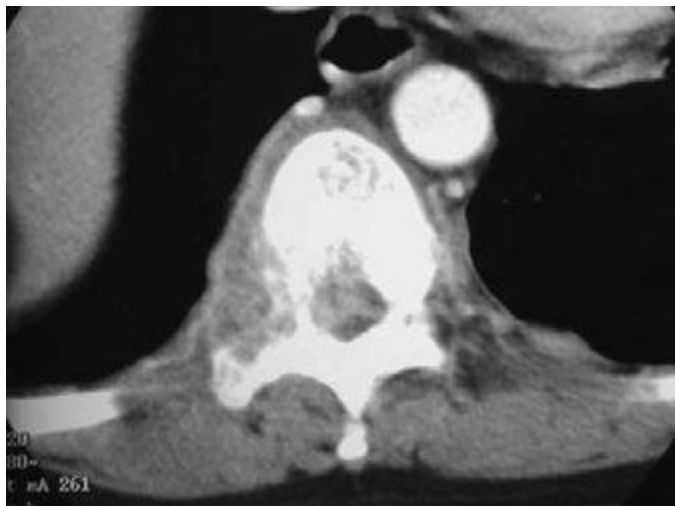


FIGURE 202-10 CT scan demonstrating destruction of the right pedicle of T10 due to Pott's disease. The patient, a 70-year-old Asian woman, presented with back pain and weight loss and had biopsy-proven tuberculosis. (Courtesy of Charles L. Daley, MD, University of California, San Francisco; with permission.)

Although synovial fluid culture is positive in a high percentage of cases, synovial biopsy and tissue culture may be necessary to establish the diagnosis. Skeletal TB responds to chemotherapy, but severe cases may require surgery.

Tuberculous Meningitis and Tuberculoma TB of the central nervous system accounts for ~5% of extrapulmonary cases in the United States. It is seen most often in young children but also develops in adults, especially those infected with HIV. Tuberculous meningitis results from the hematogenous spread of primary or postprimary pulmonary TB or from the rupture of a subependymal tubercle into the subarachnoid space. In more than half of cases, evidence of old pulmonary lesions or a miliary pattern is found on chest radiography. The disease often presents subtly as headache and slight mental changes after a prodrome of weeks of low-grade fever, malaise, anorexia, and irritability. If not recognized, tuberculous meningitis may evolve acutely with severe headache, confusion, lethargy, altered sensorium, and neck rigidity. Typically, the disease evolves over 1–2 weeks, a course longer than that of bacterial meningitis. Because meningeal involvement is pronounced at the base of the brain, paresis of cranial nerves (ocular nerves in particular) is a frequent finding, and the involvement of cerebral arteries may produce focal ischemia. The ultimate evolution is toward coma, with hydrocephalus and intracranial hypertension.

Lumbar puncture is the cornerstone of diagnosis. In general, examination of cerebrospinal fluid (CSF) reveals a high leukocyte count (up to 1000/ μ L), usually with a predominance of lymphocytes but sometimes with a predominance of neutrophils in the early stage; a protein content of 1–8 g/L (100–800 mg/dL); and a low glucose concentration. However, any of these three parameters can be within the normal range. AFBs are infrequently seen on direct smear of CSF sediment, and repeated lumbar punctures increase the yield. Culture of CSF is diagnostic in up to 80% of cases and remains the gold standard. Real-time automated nucleic acid amplification (the Xpert MTB/RIF assay; see “Nucleic Acid Amplification Technology,” below) has a sensitivity of up to 80% and is the preferred initial diagnostic option. Treatment should be initiated immediately upon a positive Xpert MTB/RIF result. A negative result does not exclude a diagnosis of TB and requires further diagnostic workup. Imaging studies (CT and MRI) may show hydrocephalus and abnormal enhancement of basal cisterns or ependyma. If unrecognized, tuberculous meningitis is uniformly fatal. This disease responds to chemotherapy; however, neurologic sequelae are documented in 25% of treated cases, in most of which the diagnosis has been delayed. Clinical trials have demonstrated that patients given adjunctive glucocorticoids may experience faster resolution of CSF

abnormalities and elevated CSF pressure. In one study, adjunctive dexamethasone significantly enhanced the chances of survival among persons >14 years of age but did not reduce the frequency of neurologic sequelae. The dexamethasone schedule was (1) 0.4 mg/kg per day given IV with tapering by 0.1 mg/kg per week until the fourth week, when 0.1 mg/kg per day was administered; followed by (2) 4 mg/d given by mouth with tapering by 1 mg per week until the fourth week, when 1 mg/d was administered.

Tuberculoma, an uncommon manifestation of central nervous system TB, presents as one or more space-occupying lesions and usually causes seizures and focal signs. CT or MRI reveals contrast-enhanced ring lesions, but biopsy is necessary to establish the diagnosis.

Gastrointestinal TB Gastrointestinal TB is uncommon, making up 3.5% of extrapulmonary cases in the United States. Various pathogenetic mechanisms are involved: swallowing of sputum with direct seeding, hematogenous spread, or (largely in developing areas) ingestion of milk from cows affected by bovine TB. Although any portion of the gastrointestinal tract may be affected, the terminal ileum and the cecum are the sites most commonly involved. Abdominal pain (at times similar to that associated with appendicitis) and swelling, obstruction, hematochezia, and a palpable mass in the abdomen are common findings at presentation. Fever, weight loss, anorexia, and night sweats are also common. With intestinal-wall involvement, ulcerations and fistulae may simulate Crohn's disease; the differential diagnosis of this entity is always difficult. Anal fistulae should prompt an evaluation for rectal TB. Because surgery is required in most cases, the diagnosis can be established by histologic examination and culture of specimens obtained intraoperatively.

Tuberculous peritonitis follows either the direct spread of tubercle bacilli from ruptured lymph nodes and intraabdominal organs (e.g., genital TB in women) or hematogenous seeding. Nonspecific abdominal pain, fever, and ascites should raise the suspicion of tuberculous peritonitis. The coexistence of cirrhosis (Chap. 363) in patients with tuberculous peritonitis complicates the diagnosis. In tuberculous peritonitis, paracentesis reveals an exudative fluid with a high protein content and leukocytosis that is usually lymphocytic (although neutrophils occasionally predominate). The yield of direct smear and culture is relatively low; culture of a large volume of ascitic fluid can increase the yield, but peritoneal biopsy (with a specimen best obtained by laparoscopy) is often needed to establish the diagnosis.

Pericardial TB (Tuberculous Pericarditis) Due either to direct extension from adjacent mediastinal or hilar lymph nodes or to hematogenous spread, pericardial TB has often been a disease of the elderly in countries with low TB prevalence. However, it also develops frequently in HIV-infected patients. Case-fatality rates are as high as 40% in some series. The onset may be subacute, although an acute presentation, with dyspnea, fever, dull retrosternal pain, and a pericardial friction rub, is possible. An effusion eventually develops in many cases; cardiovascular symptoms and signs of cardiac tamponade may ultimately appear (Chap. 288). In the presence of effusion, TB must be suspected if the patient belongs to a high-risk population (HIV-infected, originating in a high-prevalence country); if there is evidence of previous TB in other organs; or if echocardiography, CT, or MRI shows effusion and thickness across the pericardial space. A definitive diagnosis can be obtained by pericardiocentesis under echocardiographic guidance. The pericardial fluid must be submitted for biochemical, cytologic, and microbiologic evaluation. The effusion is exudative in nature, with a high count of lymphocytes and monocytes. Hemorrhagic effusion is common. Direct smear examination is very rarely positive. Culture of pericardial fluid reveals *M. tuberculosis* in up to two-thirds of cases, whereas pericardial biopsy has a higher yield. High levels of ADA, lysozyme, and IFN- γ may suggest a tuberculous etiology.

Without treatment, pericardial TB is usually fatal. Even with treatment, complications may develop, including chronic constrictive pericarditis with thickening of the pericardium, fibrosis, and sometimes calcification, which may be visible on a chest radiograph. Systematic reviews and meta-analyses show that adjunctive glucocorticoid treatment remains controversial, with no conclusive evidence of benefits

1112 for all principal outcomes of pericarditis—i.e., no significant impact on resolution of effusion, no significant difference in functional status after treatment, and no significant reduction in the frequency of development of constriction or death. However, in HIV-infected patients, glucocorticoids do improve functional status after treatment.

Caused by direct extension from the pericardium or by retrograde lymphatic extension from affected mediastinal lymph nodes, tuberculous myocarditis is an extremely rare disease. Usually it is fatal and is diagnosed postmortem.

Miliary or Disseminated TB Miliary TB is due to hematogenous spread of tubercle bacilli. Although in children it is often the consequence of primary infection, in adults it may be due to either recent infection or reactivation of old disseminated foci. The lesions are usually yellowish granulomas 1–2 mm in diameter that resemble millet seeds (thus the term *miliary*, coined by nineteenth-century pathologists). Clinical manifestations are nonspecific and protean, depending on the predominant site of involvement. Fever, night sweats, anorexia, weakness, and weight loss are presenting symptoms in the majority of cases. At times, patients have a cough and other respiratory symptoms due to pulmonary involvement as well as abdominal symptoms. Physical findings include hepatomegaly, splenomegaly, and lymphadenopathy. Eye examination may reveal choroidal tubercles, which are pathognomonic of miliary TB, in up to 30% of cases. Meningismus occurs in fewer than 10% of cases.

A high index of suspicion is required for the diagnosis of miliary TB. Frequently, chest radiography (Fig. 202-5) reveals a miliary reticulonodular pattern (more easily seen on underpenetrated film), although no radiographic abnormality may be evident early in the course and among HIV-infected patients. Other radiologic findings include large infiltrates, interstitial infiltrates (especially in HIV-infected patients), and pleural effusion. Sputum-smear microscopy is negative in most cases. Various hematologic abnormalities may be seen, including anemia with leukopenia, lymphopenia, neutrophilic leukocytosis and leukemoid reactions, and polycythemia. Disseminated intravascular coagulation has been reported. Elevation of alkaline phosphatase levels and other abnormal values in liver function tests are detected in patients with severe hepatic involvement. The TST may be negative in up to half of cases, but reactivity may be restored during chemotherapy. Bronchoalveolar lavage and transbronchial biopsy are more likely to provide bacteriologic confirmation, and granulomas are evident in liver or bone-marrow biopsy specimens from many patients. If it goes unrecognized, miliary TB is lethal; with proper early treatment, however, it is amenable to cure. Glucocorticoid therapy has not proved beneficial.

A rare presentation seen in the elderly, *cryptic miliary TB* has a chronic course characterized by mild intermittent fever, anemia, and—ultimately—meningeal involvement preceding death. An acute septicemic form, *nonreactive miliary TB*, occurs very rarely and is due to massive hematogenous dissemination of tubercle bacilli. Pancytopenia is common in this form of disease, which is rapidly fatal. At postmortem examination, multiple necrotic but nongranulomatous (“nonreactive”) lesions are detected.

Less Common Extrapulmonary Forms TB may cause chorioretinitis, uveitis, panophthalmitis, and painful hypersensitivity-related phlyctenular conjunctivitis. Tuberculous otitis is rare and presents as hearing loss, otorrhea, and tympanic membrane perforation. In the nasopharynx, TB may simulate granulomatosis with polyangiitis. Cutaneous manifestations of TB include primary infection due to direct inoculation, abscesses and chronic ulcers, scrofuloderma, lupus vulgaris (a smoldering disease with nodules, plaques, and fissures), miliary lesions, and erythema nodosum. Tuberculous mastitis results from retrograde lymphatic spread, often from the axillary lymph nodes. Adrenal TB is a manifestation of disseminated disease presenting rarely as adrenal insufficiency. Finally, congenital TB results from transplacental spread of tubercle bacilli to the fetus or from ingestion of contaminated amniotic fluid. This rare disease affects the liver, spleen, lymph nodes, and various other organs.

Post-TB Complications TB may cause persisting pulmonary damage in patients whose infection has been considered cured on clinical

grounds. Chronic impairment of lung functions, bronchiectasis, aspergillomas, and chronic pulmonary aspergillosis (CPA) have been associated with TB. CPA may manifest as simple aspergilloma (fungal ball) or chronic cavitary aspergillosis. Early studies revealed that, especially in the presence of large residual cavities, *Aspergillus fumigatus* may colonize the lesion and produce symptoms such as respiratory impairment, hemoptysis, persistent fatigue, and weight loss, often resulting in the erroneous diagnosis of TB recurrence. The detection of *Aspergillus* precipitins (IgG) in the blood suggests CPA, as do radiographic abnormalities such as thickening of the cavitary walls or the presence of a fungal ball inside the cavity. Treatment is difficult. Recent preliminary studies on the use of itraconazole for 6 months suggest that treatment with this agent may be superior to conservative treatment in improving radiologic and clinical manifestations of CPA. Surgical removal of lesions is risky.

HIV-Associated TB (See also Chap. 226) TB is one of the most common diseases among HIV-infected persons worldwide and a major cause of death in this population; more specifically, it is responsible for an estimated 24% of all HIV-related mortality. In certain urban settings in some African countries, the rate of HIV infection among TB patients reaches 70–80%. A person with a positive TST who acquires HIV infection has a 3–13% annual risk of developing active TB. A new TB infection acquired by an HIV-infected individual may evolve to active disease in a matter of weeks rather than months or years. TB can appear at any stage of HIV infection, and its presentation varies with the stage. When CMI is only partially compromised, pulmonary TB presents in a typical manner (Figs. 202-6 and 202-7), with upper-lobe infiltrates and cavitation and without significant lymphadenopathy or pleural effusion. In late stages of HIV infection, when the CD4+ T cell count is <200/μL, a primary TB-like pattern, with diffuse interstitial and subtle infiltrates, little or no cavitation, pleural effusion, and intrathoracic lymphadenopathy, is more common. However, these forms are becoming less common because of the expanded use of antiretroviral treatment (ART). Overall, sputum smears are less frequently positive among TB patients with HIV infection than among those without; thus, the diagnosis of TB may be difficult, especially in view of the variety of HIV-related pulmonary conditions mimicking TB. Extrapulmonary TB is common among HIV-infected patients. In various series, extrapulmonary TB—alone or in association with pulmonary disease—has been documented in 40–60% of all cases in HIV-co-infected individuals. The most common forms are lymphatic, disseminated, pleural, and pericardial. Mycobacteremia and meningitis are also common, particularly in advanced HIV disease. The diagnosis of TB in HIV-infected patients may be complicated not only by the increased frequency of sputum-smear negativity (up to 40% in culture-proven pulmonary cases) but also by atypical radiographic findings, a lack of classic granuloma formation in the late stages, and a negative TST. The Xpert MTB/RIF assay (see “Nucleic Acid Amplification Technology,” below) is the preferred initial diagnostic option, and therapy should be started on the basis of a positive result because treatment delays may be fatal. A negative Xpert MTB/RIF result does not exclude a diagnosis of TB, and culture remains the gold standard.

Exacerbations in systemic (lymphadenopathy) or respiratory symptoms, signs, and laboratory or radiographic manifestations of TB—termed the *immune reconstitution inflammatory syndrome* (IRIS) or *TB immune reconstitution disease* (TB-IRD)—have been associated with the administration of ART and occur in ~10% of HIV-infected TB patients. Usually developing 1–3 months after initiation of ART, IRIS is more common among patients with advanced immunosuppression and extrapulmonary TB. “Unmasking IRIS” may also develop after the initiation of ART in patients with undiagnosed subclinical TB. The earlier ART is started and the lower the baseline CD4+ T cell count, the greater the risk of IRIS. Death due to IRIS is relatively infrequent and occurs mainly among patients who have a high preexisting mortality risk. The presumed pathogenesis of IRIS consists of an immune response that is elicited by antigens released as bacilli are killed during effective chemotherapy and that is temporally associated with improving immune function. There

is no diagnostic test for IRIS, and its confirmation relies heavily upon case definitions incorporating clinical and laboratory data; a variety of case definitions have been suggested. The first priority in the management of a possible case of IRIS is to ensure that the clinical syndrome does not represent a failure of TB treatment or the development of another infection. Mild paradoxical reactions can be managed with symptom-based treatment. Glucocorticoids have been used for more severe reactions, and prednisolone given for 4 weeks at a low dosage (1.5 mg/kg per day for 2 weeks and half that dose for the remaining 2 weeks) has reduced the need for hospitalization and therapeutic procedures and hastened alleviation of symptoms, as reflected by Karnofsky performance scores, quality-of-life assessments, radiographic response, and C-reactive protein levels. The effectiveness of glucocorticoids in alleviating the symptoms of IRIS is probably linked to suppression of proinflammatory cytokine concentrations, as these medications reduce serum concentrations of IL-6, IL-10, IL-12p40, TNF- α , IFN- γ , and IFN- γ -inducible protein 10 (IP-10). Recommendations for the prevention and treatment of TB in HIV-infected individuals are provided below.

DIAGNOSIS

The key to the diagnosis of TB remains a high index of suspicion. Diagnosis is not difficult in persons belonging to high-risk populations who present with typical symptoms and a classic chest radiograph showing upper-lobe infiltrates with cavities (Fig. 202-6). On the other hand, the diagnosis can easily be missed in an elderly nursing-home resident or a teenager with a focal infiltrate. Often, the diagnosis is first entertained when the chest radiograph of a patient being evaluated for respiratory symptoms is abnormal. If the patient has no complicating medical conditions that cause immunosuppression, the chest radiograph may show typical upper-lobe infiltrates with cavitation (Fig. 202-6). The longer the delay between the onset of symptoms and the diagnosis, the more likely is the finding of cavitory disease. In contrast, immunosuppressed patients, including those with HIV infection, may have “atypical” findings on chest radiography—e.g., lower-zone infiltrates without cavity formation.

The several approaches to the diagnosis of TB require, above all, a well-organized laboratory network with an appropriate distribution of tasks at different levels of the health care system. At the peripheral and community levels, screening and referral are the principal tasks—besides clinical assessment and radiography—that can be accomplished through AFB microscopy and/or real-time automated nucleic acid amplification technology (the Xpert MTB/RIF assay; see below). At a secondary level (e.g., a traditional district hospital in a high-incidence setting), additional technology can be adopted, including rapid culture and drug susceptibility testing.

AFB MICROSCOPY

A presumptive diagnosis is commonly based on the finding of AFB on microscopic examination of a diagnostic specimen, such as a smear of expectorated sputum or of tissue (e.g., a lymph node biopsy). Although inexpensive, AFB microscopy has relatively low sensitivity (40–60%) in culture-confirmed cases of pulmonary TB. The traditional method—light microscopy of specimens stained with Ziehl-Neelsen basic fuchsin dyes—is nevertheless satisfactory, although time-consuming. Most modern laboratories processing large numbers of diagnostic specimens use auramine–rhodamine staining and fluorescence microscopy; this approach is more sensitive than the Ziehl-Neelsen method. However, it is expensive because it requires high-cost mercury vapor light sources and a dark room. Less expensive light-emitting diode (LED) fluorescence microscopes are now available. They are as sensitive as—or more sensitive than—traditional fluorescence microscopes. As a result, conventional light and fluorescence microscopes are being replaced with this more recent technology, especially in developing countries. For patients with suspected pulmonary TB, it has been recommended that two or three sputum specimens, preferably collected early in the morning, should be submitted to the laboratory for AFB smear and mycobacterial culture. Two specimens collected on the

same visit may be as effective as three. If tissue is obtained, it is critical that the portion of the specimen intended for culture not be put in formaldehyde. The use of AFB microscopy in examining urine or gastric lavage fluid is limited by the presence of commensal mycobacteria that can cause false-positive results.

NUCLEIC ACID AMPLIFICATION TECHNOLOGY

Several test systems based on amplification of mycobacterial nucleic acid have become available in the past few years. These tests are most useful for the rapid confirmation of TB in persons with AFB-positive specimens, but some also have utility for the diagnosis of AFB-negative pulmonary and extrapulmonary TB. One system that permits rapid diagnosis of TB with high specificity and sensitivity (approaching that of culture) is the fully automated, real-time nucleic acid amplification technology known as the Xpert MTB/RIF assay. Xpert MTB/RIF can simultaneously detect TB and rifampin resistance in <2 h and has minimal biosafety and training requirements. Therefore, it can be housed in nonconventional laboratory settings. The WHO recommends its use worldwide as the initial diagnostic test in adults and children presumed to have MDR-TB or HIV-associated TB. Taking into account the availability of resources, the test may also be used in any adult or child presumed to have TB or as a follow-up test after microscopy in adults presumed to have TB but not at risk of MDR-TB or HIV-associated TB. Xpert MTB/RIF should be the initial test applied to CSF from patients in whom TB meningitis is suspected as well as a replacement test (over conventional microscopy, culture, and histopathology) for selected nonrespiratory specimens—obtained by gastric lavage, fine-needle aspiration, or pleural or other biopsies—from patients in whom extrapulmonary TB is suspected. This test has a sensitivity of 98% among AFB-positive cases and ~70% among AFB-negative specimens. Other tests, such as those based on manual amplification platforms, have not yet been deemed satisfactory for introduction into clinical practice as replacements for existing tests.

MYCOBACTERIAL CULTURE

Definitive diagnosis depends on the isolation and identification of *M. tuberculosis* from a clinical specimen or the identification of specific DNA sequences in a nucleic acid amplification test. Specimens may be inoculated onto egg- or agar-based medium (e.g., Löwenstein-Jensen or Middlebrook 7H10) and incubated at 37°C (under 5% CO₂ for Middlebrook medium). Because most species of mycobacteria, including *M. tuberculosis*, grow slowly, 4–8 weeks may be required before growth is detected. Although *M. tuberculosis* may be identified presumptively on the basis of growth time and colony pigmentation and morphology, a variety of biochemical tests have traditionally been used to speciate mycobacterial isolates. In modern, well-equipped laboratories, liquid culture for isolation and species identification by molecular methods or high-pressure liquid chromatography of mycolic acids has replaced isolation on solid media and identification by biochemical tests. A widely used technology is the mycobacterial growth indicator tube (BBL™ MGIT™; BD, Franklin Lakes, NJ), which uses a fluorescent compound sensitive to the presence of oxygen dissolved in the liquid medium. The appearance of fluorescence detected by fluorometric technology indicates active growth of mycobacteria. A low-cost, rapid immunochromatographic lateral-flow assay based on detection of MTP64 antigen may also be used for species identification of the *M. tuberculosis* complex in culture isolates. These new methods, which are also being introduced in low-income countries, have decreased the time required for bacteriologic confirmation of TB to 2–3 weeks.

DRUG SUSCEPTIBILITY TESTING

Any initial isolate of *M. tuberculosis* should be tested for susceptibility to isoniazid and rifampin in order to detect drug resistance and/or MDR-TB, particularly if one or more risk factors for drug resistance are identified or if the patient either fails to respond to initial therapy or has a relapse after the completion of treatment (see “Treatment Failure and Relapse,” below). In addition, expanded susceptibility testing for second-line anti-TB drugs (especially the fluoroquinolones

1114 and the injectable drugs) is mandatory when MDR-TB is found. Susceptibility testing may be conducted directly (with the clinical specimen) or indirectly (with mycobacterial cultures) on solid or liquid medium. Results are obtained rapidly by direct susceptibility testing on liquid medium, with an average reporting time of 3 weeks. With indirect testing on solid medium, results may be unavailable for ≥ 8 weeks. Highly reliable genotypic methods for the rapid identification of genetic mutations in gene regions known to be associated with resistance to rifampin (such as those in *rpoB*) and isoniazid (such as those in *katG* and *inhA*) have been developed and are being widely implemented for screening of patients at increased risk of drug-resistant TB. Apart from the Xpert MTB/RIF assay, which, as mentioned above, detects rifampin resistance, the most widely used are molecular line probe assays. After extraction of DNA from *M. tuberculosis* isolates or from clinical specimens, the resistance gene regions are amplified by polymerase chain reaction (PCR), and labeled and probe-hybridized PCR products are detected by colorimetric development. This assay reveals the presence of *M. tuberculosis* as well as mutations in target resistance gene regions. A similar approach has been developed for second-line anti-TB drugs such as the fluoroquinolones, the aminoglycosides kanamycin and amikacin, and capreomycin, but the diagnostic accuracy of the current technology is not yet sufficient to recommend its use in clinical practice. Finally, a few noncommercial, inexpensive culture and drug-susceptibility testing methods (e.g., microscopically observed drug susceptibility, or MODS; nitrate reductase assays; and colorimetric redox indicator assays) may be useful in resource-limited settings. Their use is restricted to national reference laboratories with proven proficiency and adequate external quality control as an interim solution while genotypic or automated liquid culture technology is introduced.

RADIOGRAPHIC PROCEDURES

As noted above, the initial suspicion of pulmonary TB is often based on abnormal chest radiographic findings in a patient with respiratory symptoms. Although the “classic” picture is that of upper-lobe disease with infiltrates and cavities (Fig. 202-6), virtually any radiographic pattern—from a normal film or a solitary pulmonary nodule to diffuse alveolar infiltrates in a patient with adult respiratory distress syndrome—may be seen. In the era of AIDS, no radiographic pattern can be considered pathognomonic. CT (Fig. 202-7) may be useful in interpreting questionable findings on plain chest radiography and may be helpful in diagnosing some forms of extrapulmonary TB (e.g., Pott’s disease; Fig. 202-10). MRI is useful in the diagnosis of intracranial TB.

ADDITIONAL DIAGNOSTIC PROCEDURES

Other diagnostic tests may be used when pulmonary TB is suspected. Sputum induction by ultrasonic nebulization of hypertonic saline may be useful for patients who cannot produce a sputum specimen spontaneously. Frequently, patients with radiographic abnormalities that are consistent with other diagnoses (e.g., bronchogenic carcinoma) undergo fiberoptic bronchoscopy with bronchial brushings and endobronchial or transbronchial biopsy of the lesion. Bronchoalveolar lavage of a lung segment containing an abnormality may also be performed. In all cases, it is essential that specimens be submitted for AFB smear, mycobacterial culture, and molecular testing with the Xpert MTB/RIF assay. For the diagnosis of primary pulmonary TB in children, who often do not expectorate sputum, induced sputum specimens and specimens from early-morning gastric lavage may yield positive cultures and/or Xpert MTB/RIF assay results.

Invasive diagnostic procedures are indicated for patients with suspected extrapulmonary TB. In addition to testing of specimens from involved sites (e.g., CSF for tuberculous meningitis, pleural fluid and biopsy samples for pleural disease), biopsy and culture of bone marrow and liver tissue have a good diagnostic yield in disseminated (miliary) TB, particularly in HIV-infected patients, who also have a high frequency of positive blood cultures. In some cases, culture or Xpert MTB/RIF assay results are negative but a clinical diagnosis of TB is supported by consistent epidemiologic evidence (e.g., a history of close contact with an infectious patient) and a compatible clinical

and radiographic response to treatment. In the United States and other industrialized countries with low rates of TB, some patients with limited abnormalities on chest radiographs and sputum positive for AFB are infected with nontuberculous mycobacteria, most commonly organisms of the *M. avium* complex or *M. kansasii* (Chap. 204). Factors favoring the diagnosis of nontuberculous mycobacterial disease over TB include an absence of risk factors for TB and the presence of underlying chronic pulmonary disease.

Patients with HIV-associated TB pose several diagnostic problems (see “HIV-Associated TB,” above). Moreover, HIV-infected patients with sputum culture-positive, AFB-positive TB may present with a normal chest radiograph. The Xpert MTB/RIF assay is the preferred rapid diagnostic test in this population of patients because of simplicity of use and a sensitivity of $\sim 60\%$ among AFB-negative culture-positive cases and of 97% among AFB-positive cases. With the advent of ART, the occurrence of disseminated *M. avium* complex disease that can be confused with TB has become much less common.

SEROLOGIC AND OTHER DIAGNOSTIC TESTS FOR ACTIVE TB

A number of serologic tests based on detection of antibodies to a variety of mycobacterial antigens are marketed in developing countries but not in the United States. Careful independent assessments of these tests suggest that they are not useful as diagnostic aids—especially in persons with a low probability of TB—because of their low sensitivity and specificity and their poor reproducibility. After a rigorous evaluation of the tests, the WHO issued a “negative” recommendation in 2011 in order to prevent their abuse in the private sector of many resource-limited countries. Various methods aimed at detection of mycobacterial antigens in diagnostic specimens are being investigated but are limited at present by low sensitivity. Determinations of ADA and IFN- γ levels in pleural fluid may be useful adjunctive tests in the diagnosis of pleural TB; their utility in the diagnosis of other forms of extrapulmonary TB (e.g., pericardial, peritoneal, and meningeal) is less clear.

DIAGNOSIS OF LATENT *M. TUBERCULOSIS* INFECTION

Tuberculin Skin Testing In 1891, Robert Koch discovered that components of *M. tuberculosis* in a concentrated liquid culture medium, subsequently named “old tuberculin,” were capable of eliciting a skin reaction when injected subcutaneously into patients with TB. In 1932, Seibert and Munday purified this product by ammonium sulfate precipitation to produce an active protein fraction known as *tuberculin purified protein derivative* (PPD). In 1941, PPD-S, developed by Seibert and Glenn, was chosen as the international standard. Later, the WHO and UNICEF sponsored large-scale production of a master batch of PPD (RT23) and made it available for general use. The greatest limitation of PPD is its lack of mycobacterial species specificity, a property due to the large number of proteins in this product that are highly conserved in the various species. In addition, subjectivity of the skin-reaction interpretation, deterioration of the product, and batch-to-batch variations limit the usefulness of PPD.

The skin test with tuberculin PPD (TST) is most widely used in screening for LTBI. It probably measures the response to antigenic stimulation by T cells that reside in the skin rather than the response of recirculating memory T cells. The test is of limited value in the diagnosis of active TB because of its relatively low sensitivity and specificity and its inability to discriminate between LTBI and active disease. False-negative reactions are common in immunosuppressed patients and in those with overwhelming TB. False-positive reactions may be caused by infections with nontuberculous mycobacteria (Chap. 204) and by BCG vaccination. Repeated TST can produce larger reaction sizes due to either boosting or true conversion. The “boosting phenomenon” is a spurious TST conversion resulting from boosting of reactivity on subsequent TST 1–5 weeks after the initial test. Distinguishing boosting from true conversion is difficult yet important and can be based on clinical and epidemiologic considerations. For instance, true conversions are likely after BCG vaccination in a previously TST-negative person or in a close contact of an infectious patient.

IFN- γ Release Assays Two in vitro assays that measure T cell release of IFN- γ in response to stimulation with the highly TB-specific antigens ESAT-6 and CFP-10 are available. The T-SPOT[®] TB test (Oxford Immunotec, Oxford, United Kingdom) is an enzyme-linked immunospot (ELISpot) assay, and the QuantiFERON[®]-TB Gold test (Qiagen GmbH, Hilden, Germany) is a whole-blood enzyme-linked immunosorbent assay (ELISA) for measurement of IFN- γ . The QuantiFERON[®]-TB Gold In-Tube assay, which facilitates blood collection and initial incubation, also contains another specific antigen, TB7.7. These tests likely measure the response of recirculating memory T cells—normally part of a reservoir in the spleen, bone marrow, and lymph nodes—to persisting bacilli producing antigenic signals.

In settings or population groups with low TB and HIV burdens, IFN- γ release assays (IGRAs) have previously been reported to be more specific than the TST as a result of less cross-reactivity due to BCG vaccination and sensitization by nontuberculous mycobacteria. Recent studies, however, suggest that IGRAs may not perform well in serial testing (e.g., among health care workers) and that interpretation of test results is dependent on cutoff values used to define positivity. Potential advantages of IGRAs include logistical convenience, the need for fewer patient visits to complete testing, and the avoidance of somewhat subjective measurements such as skin induration. However, IGRAs require that blood be drawn from the individual and then delivered to the laboratory in a timely fashion. IGRAs also require that testing be performed in a laboratory setting. These requirements pose challenges similar to those faced with the TST, including cold-chain requirements and batch-to-batch variations. Because of higher specificity and other potential advantages, IGRAs have usually replaced the TST for LTBI diagnosis in low-incidence, high-income settings. However, in high-incidence TB and HIV settings and population groups, there is limited and inconclusive evidence about the performance and usefulness of IGRAs. In view of higher costs and increased technical requirements, the WHO does not recommend the replacement of the TST by IGRAs in low- and middle-income countries.

A number of national guidelines on the use of IGRAs for LTBI testing have been issued. In the United States, an IGRA is preferred to the TST for most persons over the age of 5 years who are being screened for LTBI. However, for those at high risk of progression to active TB (e.g., HIV-infected persons), either test—or, to optimize sensitivity, both tests—may be used. Because of the paucity of data on the use of IGRAs in children, the TST is preferred for LTBI testing of children under age 5. In Canada and some European countries, a two-step approach for those with positive TSTs—i.e., initial TST followed by an IGRA—is recommended. However, a TST may boost an IGRA response if the interval between the two tests exceeds 3 days. Similar to the TST, current IGRAs have only modest predictive value for incident active TB, are not useful in identifying patients with the highest risk of progression toward disease, and cannot be used for diagnosis of active TB.

TREATMENT TUBERCULOSIS

The two aims of TB treatment are (1) to prevent morbidity and death by curing TB while preventing the emergence of drug resistance and (2) to interrupt transmission by rendering patients noninfectious. Chemotherapy for TB became possible with the discovery of streptomycin in 1943. Randomized clinical trials clearly indicated that the administration of streptomycin to patients with chronic TB reduced mortality rates and led to cure in the majority of cases. However, monotherapy with streptomycin eventually was associated with the development of resistance to this drug and the resulting failure of treatment. With the introduction into clinical practice of para-aminosalicylic acid (PAS) and isoniazid, it became axiomatic in the early 1950s that cure of TB required the concomitant administration of at least two agents to which the organism was susceptible. Furthermore, early clinical trials demonstrated that a long period of treatment—i.e., 12–24 months—was required to prevent recurrence. The introduction of rifampin (rifampicin) in the early 1970s

heralded the era of effective short-course chemotherapy, with a treatment duration of <12 months. The discovery that pyrazinamide, which was first used in the 1950s, augmented the potency of isoniazid/rifampin regimens led to the use of a 6-month course of this triple-drug regimen as standard therapy.

DRUGS

Four major drugs are considered first-line agents for the treatment of TB: isoniazid, rifampin, pyrazinamide, and ethambutol (**Table 202-2**). These drugs are well absorbed after oral administration, with peak serum levels at 2–4 h and nearly complete elimination within 24 h. These agents are recommended on the basis of their bactericidal activity (i.e., their ability to rapidly reduce the number of viable organisms and render patients noninfectious), their sterilizing activity (i.e., their ability to kill all bacilli and thus sterilize the affected tissues, measured in terms of the ability to prevent relapses), and their low rate of induction of drug resistance by selection of mutant bacilli. Two additional rifamycins, rifapentine and rifabutin, are also available in the United States; however, the level of cross-resistance with rifampin is high. For a detailed discussion of the drugs used for the treatment of TB, see **Chap. 205e**.

Because of a lower degree of efficacy and a higher degree of intolerance and toxicity, six classes of second-line drugs are generally used only for the treatment of patients with TB resistant to first-line drugs: (1) the fluoroquinolone antibiotics; (2) the injectable aminoglycosides kanamycin, amikacin, and streptomycin; (3) the injectable polypeptide capreomycin; and the oral agents (4) ethionamide and prothionamide, (5) cycloserine and terizidone (therizidone), and (6) PAS. Streptomycin, formerly a first-line agent, is now rarely used for drug-resistant TB because resistance levels worldwide are high and it is more toxic than the other drugs in the same class; however, the level of cross-resistance with the other injectables is low. Of the quinolones, later-generation agents such as levofloxacin and moxifloxacin are preferred. Gatifloxacin (no longer marketed in several countries, including the United States, because of previously observed dysglycemia) has recently been tested in a 4-month regimen that produced no detectable major side effects; thus, this drug could be reconsidered as a good alternative. Other drugs (referred to by the WHO as “group 5”) whose efficacy is not clearly defined are used in the treatment of patients with TB resistant to most of the first- and second-line agents; these drugs include clofazimine, linezolid, amoxicillin/clavulanic acid, clarithromycin, and carbapenems such as imipenem/cilastatin and meropenem. Today amithiozone (thiacetazone) is used very rarely because it has been associated with severe and at times fatal skin reactions among HIV-infected patients. Two novel drugs belonging to two new antibiotic classes—the diarylquinoline bedaquiline and the nitroimidazole delamanid—have recently been approved for use in severe cases of MDR-TB by stringent regulatory authorities (the U.S. Food and Drug Administration

TABLE 202-2 RECOMMENDED DOSAGE^a FOR INITIAL TREATMENT OF TUBERCULOSIS IN ADULTS^b

Drug	Dosage	
	Daily Dose	Thrice-Weekly Dose
Isoniazid	5 mg/kg, max 300 mg	10 mg/kg, max 900 mg
Rifampin	10 mg/kg, max 600 mg	10 mg/kg, max 600 mg
Pyrazinamide	25 mg/kg, max 2 g	35 mg/kg, max 3 g
Ethambutol ^c	15 mg/kg	30 mg/kg

^aThe duration of treatment with individual drugs varies by regimen, as detailed in Table 202-3. ^bThe World Health Organization recommends the following dosages for children: isoniazid, 10–15 mg/kg daily, max 300 mg/d; rifampin, 15 (range, 10–20) mg/kg daily, max 600 mg/d; pyrazinamide, 35 (range, 30–40) mg/kg daily; ethambutol, 20 (range, 15–25) mg/kg daily. ^cIn certain settings, streptomycin (15 mg/kg daily, with a maximum dose of 1 g; or 25–30 mg/kg thrice weekly, with a maximum dose of 1.5 g) can replace ethambutol in the initial phase of treatment. However, streptomycin generally is no longer considered a first-line drug.

Source: Based on recommendations of the American Thoracic Society/Infectious Diseases Society of America/Centers for Disease Control and Prevention and the World Health Organization.

1116 [FDA] and the European Medicine Agency [EMA] in the case of bedaquiline; the EMA and the Pharmaceuticals and Medical Devices Agency of Japan in the case of delamanid).

REGIMENS

Standard short-course regimens are divided into an initial, or bactericidal, phase and a continuation, or sterilizing, phase. During the initial phase, the majority of the tubercle bacilli are killed, symptoms resolve, and usually the patient becomes noninfectious. The continuation phase is required to eliminate persisting mycobacteria and prevent relapse. The treatment regimen of choice for virtually all forms of drug-susceptible TB in adults consists of a 2-month initial (or intensive) phase of isoniazid, rifampin, pyrazinamide, and ethambutol followed by a 4-month continuation phase of isoniazid and rifampin (Table 202-3). This regimen can cure TB in more than 90% of patients. In children, most forms of TB in the absence of HIV infection or suspected isoniazid resistance can be safely treated without ethambutol in the intensive phase. Treatment should be given daily throughout the course. However, daily treatment during the intensive phase and intermittently (three times weekly) during the continuation phase is an alternative for patients who can be directly supervised and properly supported. A fully supervised, three-times-weekly regimen throughout the course also can be offered in the absence of HIV infection, although the risk of acquired drug resistance is higher than that among patients treated daily for the full course. In addition, if the infecting strain is resistant to isoniazid, the risks of both acquired resistance and treatment failure are higher with three-times-weekly intensive therapy than with daily treatment in the intensive phase. HIV-infected patients should always receive their initial-phase regimen daily (see below). A continuation phase of once-weekly rifapentine and isoniazid has been shown to be equally effective for HIV-seronegative patients with noncavitary pulmonary TB who have negative sputum cultures at 2 months. Patients with cavitary pulmonary TB and delayed sputum-culture conversion (i.e., those who remain culture-positive at 2 months) should be tested immediately for drug-resistant TB, and a change of regimen should be considered. To prevent isoniazid-related neuropathy, pyridoxine (10–25 mg/d) should be added to the regimen

given to persons at high risk of vitamin B₆ deficiency (e.g., alcoholics; malnourished persons; pregnant and lactating women; and patients with conditions such as chronic renal failure, diabetes, and HIV infection, which are also associated with neuropathy). A full course of therapy (completion of treatment) is defined more accurately by the total number of doses taken than by the duration of treatment, although the course should not include interruptions of longer than 4 weeks. Specific recommendations on the required number of doses for each of the various treatment regimens have been published jointly by the American Thoracic Society, the Infectious Diseases Society of America, and the CDC. In some developing countries where the ability to ensure adherence to treatment is limited, a continuation-phase regimen of daily isoniazid and ethambutol for 6 months was used in the past. However, this regimen is associated with a higher rate of relapse, failure, and death, especially among HIV-infected patients, and is no longer recommended by the WHO.

Lack of adherence to treatment is recognized worldwide as the most important impediment to cure. Moreover, the tubercle bacilli infecting patients who do not fully adhere to the prescribed regimen are likely to become drug resistant. Both patient- and provider-related factors may affect adherence. Patient-related factors include a lack of belief that the illness is significant and/or that treatment will have a beneficial effect; the existence of concomitant medical conditions (notably alcohol or substance abuse); lack of social support; fear of stigma and discrimination associated with TB; and poverty, with attendant joblessness and homelessness. Provider-related factors that may promote adherence include the support, education, and encouragement of patients and the offering of convenient clinic hours. In addition to specific measures promoting adherence, two other strategic approaches are used: direct supervision of treatment with support to the patient, consisting of incentives and enablers such as meals, travel vouchers, cash transfers, and grants to replace income loss; and provision of fixed-drug-combination products that reduce the number of tablets the patient needs to swallow. Because it is difficult to predict which patients will adhere to the recommended treatment for a disease that has important public as well as individual health implications, all patients should have their therapy directly supervised, especially during the initial phase, with

TABLE 202-3 RECOMMENDED ANTITUBERCULOSIS TREATMENT REGIMENS

Indication	Initial Phase		Continuation Phase	
	Duration, Months	Drugs	Duration, Months	Drugs
New smear- or culture-positive cases	2	HRZE ^{a,b}	4	HR ^{a,c,d}
New culture-negative cases	2	HRZE ^a	4	HR ^a
Pregnancy	2	HRE ^e	7	HR
Relapses and treatment default (pending susceptibility testing)	3	HRZES ^f	5	HRE
Failures ^g	← Tailored according to drug susceptibility testing →			
Resistance (or intolerance) to H	Throughout (6–9)	RZE ^h		
Resistance (or intolerance) to R	← Same as for MDR-TB; see below ⁱ →			
MDR-TB (resistance to at least H + R)	Throughout (20 months in most cases)	Q, Inj ^j , Eto/Pto, Z, Cs/PAS		
XDR-TB	← See Table 202-4 →			
Intolerance to Z	2	HRE	7	HR

^aAll drugs can be given daily or intermittently (three times weekly throughout). A twice-weekly regimen after 2–8 weeks of daily therapy during the initial phase is sometimes used, although it is not recommended by the WHO. ^bStreptomycin can be used in place of ethambutol but is no longer considered a first-line drug. ^cSome experts suggest extending the continuation phase to 7 months for patients with cavitary pulmonary tuberculosis who remain sputum culture-positive after the initial phase of treatment. However, treatment in such patients must be guided by drug susceptibility testing to rule out drug-resistant TB. ^dA clinical trial showed that HIV-negative patients with noncavitary pulmonary tuberculosis who have negative sputum AFB smears after the initial phase of treatment can be given once-weekly rifapentine/isoniazid in the continuation phase. ^eThe 6-month regimen with pyrazinamide can probably be used safely during pregnancy and is recommended by the WHO and the International Union Against Tuberculosis and Lung Disease. If pyrazinamide is not included in the initial treatment regimen, the minimal duration of therapy is 9 months. ^fStreptomycin should be discontinued after 2 months. Drug susceptibility results will determine the best regimen option. ^gThe availability of rapid molecular methods to identify drug resistance allows initiation of a proper regimen at the start of treatment. ^hAlthough normally not recommended, a fluoroquinolone may strengthen the regimen for patients with extensive disease. A later-generation agent (such as levofloxacin, moxifloxacin, or possibly gatifloxacin; see text) is preferred. ⁱIsoniazid is added if susceptibility to this agent is confirmed or presumed. ^jAmikacin and kanamycin (aminoglycosides) or capreomycin (polypeptide). Any of these injectable agents is recommended for the first 8 months in most patients, but the duration may be modified according to the clinical response to therapy. Continuation of treatment with the injectable drug for at least 4 months after culture conversion is advised.

Abbreviations: Cs/PAS, cycloserine or para-aminosalicylic acid; E, ethambutol; Eto/Pto, ethionamide or prothionamide; H, isoniazid; Inj, an injectable agent (the aminoglycosides amikacin and kanamycin or the polypeptide capreomycin); MDR-TB, multidrug-resistant tuberculosis; Q, a quinolone antibiotic; R, rifampin; S, streptomycin; WHO, World Health Organization; XDR-TB, extensively drug-resistant tuberculosis; Z, pyrazinamide.

proper social support including education, psychosocial counseling, and material sustainment. In an increasing number of countries, personnel to supervise therapy are usually available through TB control programs of local public health departments and from members of the community who are accepted by the patient to undertake that role and who have been properly educated by health workers. Direct supervision with patient support usually increases the proportion of patients completing treatment in all settings and greatly lessens the chances of failure, relapse, and acquired drug resistance. Fixed-drug-combination products (e.g., isoniazid/rifampin, isoniazid/rifampin/pyrazinamide, and isoniazid/rifampin/pyrazinamide/ethambutol) are available and are strongly recommended as a means of minimizing the likelihood of prescription error and of the development of drug resistance as the result of monotherapy. In some formulations of these combination products, the bioavailability of rifampin has been found to be substandard. Stringent regulatory authorities ensure that combination products are of good quality; however, this type of quality assurance is not always operative in low-income countries. Alternative regimens for patients who exhibit drug intolerance or adverse reactions are listed in Table 202-3. However, severe side effects prompting discontinuation of any of the first-line drugs and use of these alternative regimens are uncommon. The fluoroquinolones moxifloxacin and gatifloxacin have been tested as 4-month treatment-shortening regimens for drug-susceptible TB. Recently published results from these clinical trials failed to show that a 4-month regimen substituting gatifloxacin for ethambutol or moxifloxacin for either ethambutol or isoniazid is noninferior to the standard 6-month regimen. Thus, currently there is no 4-month regimen available for TB treatment.

MONITORING TREATMENT RESPONSE AND DRUG TOXICITY

Bacteriologic evaluation through culture and/or smear microscopy is essential in monitoring the response to treatment for TB. In addition, the patient's weight should be monitored regularly and the drug dosage adjusted with any significant weight change. Patients with pulmonary disease should have their sputum examined monthly until cultures become negative to allow early detection of treatment failure. With the recommended regimen, more than 80% of patients will have negative sputum cultures at the end of the second month of treatment. By the end of the third month, the sputum of virtually all patients should be culture negative. In some patients, especially those with extensive cavitary disease and large numbers of organisms, AFB smear conversion may lag behind culture conversion. This phenomenon is presumably due to the expectoration and microscopic visualization of dead bacilli. As noted above, patients with cavitary disease in whom sputum culture conversion does not occur by 2 months require immediate testing for drug resistance. When a patient's sputum cultures remain positive at ≥ 3 months, treatment failure and drug resistance or poor adherence to the regimen are likely, and testing of drug resistance should guide the choice of the best treatment option (see below). A sputum specimen should be collected by the end of treatment to document cure. If mycobacterial cultures are not practical, then monitoring by AFB smear examination should be undertaken at 2, 5, and 6 months. Smears that are positive after 3 months of treatment when the patient is known to be adherent are indicative of treatment failure and possible drug resistance. Therefore, if not done at the start of treatment, drug susceptibility testing is mandatory at this stage. Bacteriologic monitoring of patients with extrapulmonary TB is more difficult and often is not feasible. In these cases, the response to treatment must be assessed clinically and radiographically.

Monitoring of the response during chemotherapy by nucleic acid amplification technology has not been shown to be suitable. Thus Xpert MTB/RIF should not be used to monitor treatment. Likewise, serial chest radiographs are not recommended because radiographic changes may lag behind bacteriologic response and are not highly sensitive. After the completion of treatment, neither sputum examination nor chest radiography is recommended for routine follow-up purposes. However, a chest radiograph obtained

at the end of treatment may be useful for comparative purposes should the patient develop symptoms of recurrent TB months or years later. Patients should be instructed to report promptly for medical assessment if they develop any such symptoms. In addition, an end-of-treatment chest radiograph may reveal earlier the post-TB complications described above.

During treatment, patients should be monitored for drug toxicity. The most common adverse reaction of significance is hepatitis. Patients should be carefully educated about the signs and symptoms of drug-induced hepatitis (e.g., dark urine, loss of appetite) and should be instructed to discontinue treatment promptly and see their health care provider should these symptoms occur. Although biochemical monitoring is not routinely recommended, all adult patients should undergo baseline assessment of liver function (e.g., measurement of serum levels of hepatic aminotransferases and bilirubin). Older patients, those with concomitant diseases, those with a history of hepatic disease (especially hepatitis C), and those using alcohol daily should be monitored especially closely (i.e., monthly), with repeated measurements of aminotransferases, during the initial phase of treatment. Up to 20% of patients have small increases in aspartate aminotransferase (up to three times the upper limit of normal) that are not accompanied by symptoms and are of no consequence. For patients with symptomatic hepatitis and those with marked (five- to sixfold) elevations in serum levels of aspartate aminotransferase, treatment should be stopped and drugs reintroduced one at a time after liver function has returned to normal. Hypersensitivity reactions usually require the discontinuation of all drugs and rechallenge to determine which agent is the culprit. Because of the variety of regimens available, it usually is not necessary—although it is possible—to desensitize patients. Hyperuricemia and arthralgia caused by pyrazinamide can usually be managed by the administration of acetylsalicylic acid; however, pyrazinamide treatment should be stopped if the patient develops gouty arthritis. Individuals who develop autoimmune thrombocytopenia secondary to rifampin therapy should not receive the drug thereafter. Similarly, the occurrence of optic neuritis with ethambutol is an indication for permanent discontinuation of this drug. Other common manifestations of drug intolerance, such as pruritus and gastrointestinal upset, can generally be managed without the interruption of therapy.

TREATMENT FAILURE AND RELAPSE

As stated above, treatment failure should be suspected when a patient's sputum smears and/or cultures remain positive after 3 months of treatment. In the management of such patients, it is imperative that the current isolate be urgently tested for susceptibility to first- and second-line agents. Initial molecular testing for rifampin resistance should be done if the technology is available. When the results of susceptibility testing are based on molecular methods and are expected to become available within a few days, changes in the regimen can be postponed until that time. However, if the patient's clinical condition is deteriorating, an earlier change in regimen may be indicated. A cardinal rule in the latter situation is always to add more than one drug at a time to a failing regimen: at least two and preferably three drugs that have never been used and to which the bacilli are likely to be susceptible should be added. The patient may continue to take isoniazid and rifampin along with these new agents pending the results of susceptibility tests.

Patients who experience a recurrence after apparently successful treatment (relapse) are less likely to harbor drug-resistant strains (see below) than are patients in whom treatment has failed. Acquired resistance is uncommon among strains from patients in whom relapse follows the completion of a standard short-course regimen. However, pending the results of susceptibility testing, it is prudent to begin the treatment of all patients whose infections have relapsed with a standard regimen containing all four first-line drugs plus streptomycin. In less affluent countries and other settings where facilities for culture and drug susceptibility testing are not yet routinely available and where the prevalence of MDR-TB is low, the WHO recommends that a standard regimen with all four

1118 first-line drugs plus streptomycin be used in all instances of relapse and treatment default. Patients with treatment failure and those relapsing or defaulting with a high likelihood of MDR-TB should receive a regimen that includes second-line agents and is based on their history of anti-TB treatment and the drug resistance patterns in the population (Table 202-3). Once drug susceptibility test results are available, the regimen can be adjusted accordingly.

DRUG-RESISTANT TB

Strains of *M. tuberculosis* resistant to individual drugs arise by spontaneous point mutations in the mycobacterial genome that occur at low but predictable rates (10^{-7} – 10^{-10} for the key drugs). Resistance to rifampin is associated with mutations in the *rpoB* gene in 95% of cases; that to isoniazid with mutations mainly in the *katG* (50–95% of cases) and *inhA* (up to 45%) genes; that to pyrazinamide in the *pncA* gene (up to 98%); that to ethambutol in the *embB* gene (50–65%); that to the fluoroquinolones in the *gyrA*–*gyrB* genes (75–95%); and that to the aminoglycosides mainly in the *rrs* gene (up to 80%). Because there is no cross-resistance among the commonly used drugs, the probability that a strain will be resistant to two drugs is the product of the probabilities of resistance to each drug and thus is low. The development of drug-resistant TB is almost invariably the result of monotherapy—i.e., the failure of the health care provider to prescribe at least two drugs to which tubercle bacilli are susceptible or of the patient to take properly prescribed therapy. In addition, the use of drugs of substandard quality may cause the emergence of drug resistance. Drug-resistant TB may be either primary or acquired. Primary drug resistance is that which develops in a patient infected from the start by a drug-resistant strain. Acquired resistance is that which develops during treatment with an inappropriate regimen. In North America, Western Europe, most of Latin America, and the Persian Gulf States, rates of primary resistance are generally low and isoniazid resistance is most common. In the United States, although rates of primary isoniazid resistance have been stable at ~7–8%, the rate of primary MDR-TB has declined from 2.5% in 1993 to 1% since 2000. As described above, MDR-TB is an increasingly serious problem in some regions, especially in the states of the former Soviet Union and some countries of Asia (Fig. 202-11). Even more serious is the recently described occurrence of XDR-TB due to MDR strains that are also resistant to any fluoroquinolones and to any of three

second-line injectable agents (amikacin, kanamycin, and capreomycin). Creation of drug-resistant TB can be prevented by adherence to the principles of sound treatment: inclusion of at least two quality-assured, bactericidal drugs to which the organism is susceptible; use of fixed-drug-combination products; supervision of treatment with patient support; and verification that patients complete the prescribed course. Transmission of drug-resistant strains can be prevented by implementation of respiratory infection-control measures (see below).

Although the 6-month regimen described in Table 202-3 is generally effective for patients with initial isoniazid-resistant disease, it is prudent to include at least ethambutol and possibly pyrazinamide for the full 6 months and to consider extending the treatment course to 9 months. In such cases, isoniazid probably does not contribute to a successful outcome and could be omitted. In case of documented resistance to both isoniazid and ethambutol, a 9- to 12-month regimen of rifampin, pyrazinamide, and a fluoroquinolone can be used. Any patients whose isolates exhibit resistance to rifampin should be managed as if they had MDR-TB (see below), with the addition of isoniazid if susceptibility to this agent is confirmed via rapid testing or is presumed. MDR-TB, in which bacilli are resistant to (at least) isoniazid and rifampin, is more difficult to manage than is disease caused by drug-susceptible organisms because these two bactericidal drugs are the most potent agents available and because associated resistance to other first-line drugs as well (e.g., ethambutol) is not uncommon. For treatment of MDR-TB, the WHO recommends that in most patients five drugs be used in the initial phase of at least 8 months: a later-generation fluoroquinolone, an injectable agent (the aminoglycosides amikacin or kanamycin or the polypeptide capreomycin), ethionamide (or prothionamide), either cycloserine or PAS, and pyrazinamide. Ethambutol can be added (Table 202-3). Although the optimal duration of treatment is not known, a course of at least 20 months is recommended for previously untreated patients, including the initial phase with an injectable agent, which is usually discontinued at 4 months after culture conversion.

In late 2012, the FDA granted accelerated approval of bedaquiline, a diarylquinoline antibiotic. This new drug, when given for the first 24 weeks (400 mg daily for 2 weeks followed by 200 mg thrice weekly for 22 weeks), has been shown to increase the efficacy of the WHO standard regimen for MDR-TB with faster sputum conversion.

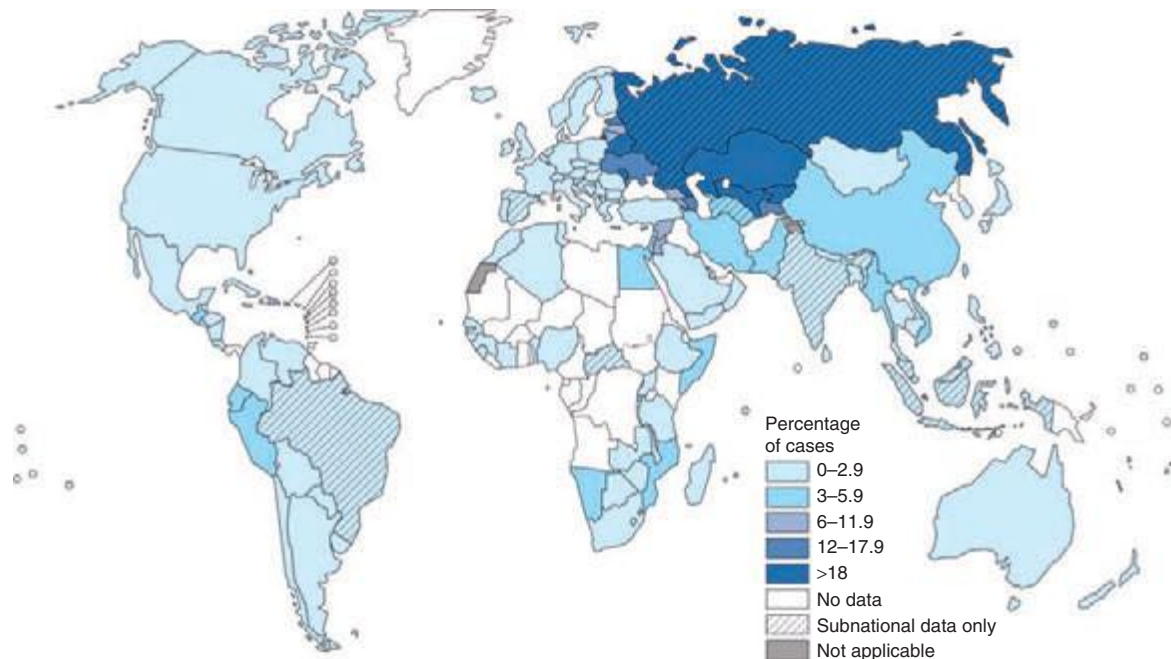


FIGURE 202-11 Percentage of new tuberculosis cases with multidrug resistance in all countries surveyed by the World Health Organization (WHO) Global Drug Resistance Surveillance Project during 1994–2013. (See disclaimer in Fig. 202-2. Courtesy of the Global TB Programme, WHO; with permission.)

Bedaquiline should be used with caution in people >65 years of age and in HIV-infected patients; its use is not advised in children and pregnant women. In early 2014, the European Medical Agency granted accelerated approval of another new agent, the nitroimidazole compound delamanid. Data from a phase 2B clinical trial in which delamanid was added to the WHO-recommended standard MDR-TB regimen have shown increased culture conversion at 2 months. Pending phase 3 trial results and in view of potential side effects of both new drugs (including QT interval prolongation in both cases and hepatotoxicity in the case of bedaquiline), the WHO recommends limiting the use of bedaquiline and delamanid to cases of MDR-TB when an effective WHO-recommended standard MDR-TB regimen cannot be designed because of known resistance, intolerance, or nonavailability of any second-line drugs in the regimen. Patients treated with bedaquiline or delamanid should be counseled, should give informed consent, and should be closely monitored during treatment. In particular, patients with cardiac anomalies such as prolonged QT interval or a history of ventricular arrhythmias should not be given these drugs. Currently, there is no information about simultaneous use of these two agents; therefore, combining them is not recommended.

Finally, a shorter (9-month) regimen consisting of gatifloxacin or moxifloxacin, clofazimine, ethambutol, and pyrazinamide given throughout the treatment period and supplemented by prothionamide, kanamycin, and high-dose isoniazid during an intensive phase of at least 4 months is reportedly effective for MDR-TB in certain settings. Further investigations are necessary to elucidate the role of this shorter regimen in MDR-TB treatment.

Patients with XDR-TB have fewer treatment options and a much poorer prognosis. However, observational studies have shown that aggressive management of cases comprising early drug-susceptibility testing, rational combination of at least five drugs, readjustment of the regimen, strict directly observed therapy, monthly bacteriologic monitoring, and intensive patient support may result in cure and avert death. **Table 202-4** summarizes the management of patients with XDR-TB. Some recently published studies regarding the use of linezolid in patients with XDR-TB suggest that, although it carries a high level of toxicity, this drug increases culture conversion.

For patients with localized disease and sufficient pulmonary reserve, lobectomy or pneumonectomy may be considered. Because the management of patients with MDR- and XDR-TB is complicated by both social and medical factors, care of these patients is ideally provided in specialized centers or, in their absence, in the context of programs with adequate resources and capacity, including community support.

HIV-ASSOCIATED TB

Several observational studies and randomized controlled trials have shown that treatment of HIV-associated TB with anti-TB drugs and simultaneous use of ART are associated with significant reductions in mortality risk and AIDS-related events. Evidence from randomized controlled trials shows that early initiation of ART during anti-TB treatment is associated with a 34–68% reduction in mortality rates, with especially good results in patients with CD4+ T cell counts of <50/μL. Therefore, the main aim in the management of HIV-associated TB is to initiate anti-TB treatment and to immediately consider initiating or continuing ART. All HIV-infected TB patients, regardless of CD4+ T cell count, are candidates for ART, which optimally is initiated as soon as possible after the diagnosis of TB and within the first 8 weeks of anti-TB therapy. However, ART should be started within the first 2 weeks of TB treatment for patients with CD4+ T cell counts of <50/μL. In general, the standard 6-month daily regimen is equally efficacious in HIV-negative and HIV-positive patients for treatment of drug-susceptible TB. As for any other adult living with HIV (**Chap. 226**), first-line ART for TB patients should consist of two nucleoside reverse transcriptase inhibitors (NRTIs) plus a nonnucleoside reverse transcriptase inhibitor (NNRTI). Although TB treatment modalities are similar to those in HIV-negative patients, adverse drug effects may be more pronounced in HIV-infected patients. In this

TABLE 202-4 MANAGEMENT GUIDELINES FOR PATIENTS WITH DOCUMENTED OR STRONGLY SUSPECTED EXTENSIVELY DRUG-RESISTANT TUBERCULOSIS (XDR-TB)

1. Use pyrazinamide and any first-line oral agents that may be effective.
2. Use an injectable agent to which the strain is susceptible, and consider an extended duration of use (12 months or possibly the whole treatment period). If the strain is resistant to all injectable agents, use of one that the patient has not previously received is recommended.^a
3. Use a later-generation fluoroquinolone, such as moxifloxacin, high-dose levofloxacin, or possibly gatifloxacin.^b
4. Use all second-line oral bacteriostatic agents (para-aminosalicylic acid, cycloserine, and ethionamide or prothionamide) that have not been used extensively in a previous regimen or any such agents that are likely to be effective.
5. Add bedaquiline or delamanid and one or more of the following drugs: clofazimine, linezolid, amoxicillin/clavulanic acid, clarithromycin, and carbapenems such as imipenem/cilastatin and meropenem.
6. The simultaneous use of bedaquiline and delamanid is not recommended at the moment in view of the current lack of information on the potential of adverse reactions when these drugs are administered together.
7. Consider treatment with high-dose isoniazid if low-level resistance to this drug is documented.
8. Consider adjuvant surgery if there is localized disease.
9. Enforce strong infection-control measures.
10. Implement strict directly observed therapy and full adherence support as well as comprehensive bacteriologic and clinical monitoring.

^aThis recommendation is made because, although the reproducibility and reliability of susceptibility testing with injectable agents are good, few data are available on the correlation of clinical efficacy with test results. Options with XDR-TB are very limited, and some strains may be affected in vivo by an injectable agent even though they test resistant in vitro. ^bGatifloxacin (no longer marketed in several countries, including the United States, because of previously observed dysglycemia) has recently been tested in a 4-month regimen that produced no detectable major side effects; thus, this drug could be reconsidered as a good alternative. ^cThe number of drugs added is based on how many oral bacteriostatic drugs (see point 4 above) are believed to be effective: the advice is to add one drug if there is confidence in all three bacteriostatic drugs; two if there is confidence in only two bacteriostatic drugs; and three or more if there is confidence in only one bacteriostatic drug or none.

regard, three important considerations are relevant: an increased frequency of paradoxical reactions, interactions between ART components and rifamycins, and development of rifampin mono-resistance with intermittent treatment. IRIS—i.e., the exacerbation of symptoms and signs of TB—has been described above. Rifampin, a potent inducer of enzymes of the cytochrome P450 system, lowers serum levels of many HIV protease inhibitors and some NNRTIs—essential drugs used in ART. In such cases, rifabutin, which has much less enzyme-inducing activity, has been used in place of rifampin. However, dosage adjustments for rifabutin and protease inhibitors are still being assessed. Several clinical trials have found that patients with HIV-associated TB whose degree of immunosuppression is advanced (e.g., CD4+ T cell counts of <100/μL) are prone to treatment failure and relapse with rifampin-resistant organisms when treated with “highly intermittent” (i.e., once- or twice-weekly) rifamycin-containing regimens. Consequently, it is recommended that all TB patients who are infected with HIV receive a rifampin-containing regimen on a daily basis. Because recommendations are frequently updated, consultation of the following websites is advised: www.who.int/hiv, www.who.int/tb, www.cdc.gov/hiv, and www.cdc.gov/tb.

SPECIAL CLINICAL SITUATIONS

Although comparative clinical trials of treatment for extrapulmonary TB are limited, the available evidence indicates that most forms of disease can be treated with the 6-month regimen recommended for patients with pulmonary disease. The WHO and the American Academy of Pediatrics recommend that children with bone and joint TB, tuberculous meningitis, or miliary TB receive up to 12 months of treatment. Treatment for TB may be complicated by underlying medical problems that require special consideration. As a rule, patients with chronic renal failure should not receive aminoglycosides and should receive ethambutol only if serum drug levels

1120 can be monitored. Isoniazid, rifampin, and pyrazinamide may be given in the usual doses in cases of mild to moderate renal failure, but the dosages of isoniazid and pyrazinamide should be reduced for all patients with severe renal failure except those undergoing hemodialysis. Patients with hepatic disease pose a special problem because of the hepatotoxicity of isoniazid, rifampin, and pyrazinamide. Patients with severe hepatic disease may be treated with ethambutol, streptomycin, and possibly another drug (e.g., a fluoroquinolone); if required, isoniazid and rifampin may be administered under close supervision. The use of pyrazinamide by patients with liver failure should be avoided. Silicotuberculosis necessitates the extension of therapy by at least 2 months.

The regimen of choice for pregnant women (Table 202-3) is 9 months of treatment with isoniazid and rifampin supplemented by ethambutol for the first 2 months. Although the WHO has recommended routine use of pyrazinamide for pregnant women, this drug has not been recommended in the United States because of insufficient data documenting its safety in pregnancy. Streptomycin is contraindicated because it is known to cause eighth-cranial-nerve damage in the fetus. Treatment for TB is not a contraindication to breast-feeding; most of the drugs administered will be present in small quantities in breast milk, albeit at concentrations far too low to provide any therapeutic or prophylactic benefit to the child.

Medical consultation on difficult-to-manage cases is provided by the U.S. CDC Regional Training and Medical Consultation Centers (www.cdc.gov/tb/education/rtmc/).

PREVENTION

The best way to prevent TB is to diagnose and isolate infectious cases rapidly and to administer appropriate treatment until patients are rendered noninfectious (usually 2–4 weeks after the start of proper treatment) and the disease is cured. Additional strategies include BCG vaccination and treatment of persons with LTBI who are at high risk of developing active disease.

BCG VACCINATION

BCG was derived from an attenuated strain of *M. bovis* and was first administered to humans in 1921. Many BCG vaccines are available worldwide; all are derived from the original strain, but the vaccines vary in efficacy, ranging from 80% to nil in randomized, placebo-controlled trials. A similar range of efficacy was found in recent observational studies (case-control, historic cohort, and cross-sectional) in areas where infants are vaccinated at birth. These studies and a meta-analysis also found higher rates of efficacy in the protection of infants and young children from serious disseminated forms of childhood TB, such as tuberculous meningitis and miliary TB. BCG vaccine is safe and rarely causes serious complications. The local tissue response begins 2–3 weeks after vaccination, with scar formation and healing within 3 months. Side effects—most commonly, ulceration at the vaccination site and regional lymphadenitis—occur in 1–10% of vaccinated persons. Some vaccine strains have caused osteomyelitis in ~1 case per million doses administered. Disseminated BCG infection (“BCGitis”) and death have occurred in 1–10 cases per 10 million doses administered, although this problem is restricted almost exclusively to persons with impaired immunity, such as children with severe combined immunodeficiency syndrome or adults with HIV infection. BCG vaccination induces TST reactivity, which tends to wane with time. The presence or size of TST reactions after vaccination does not predict the degree of protection afforded.

BCG vaccine is recommended for routine use at birth in countries with high TB prevalence. However, because of the low risk of transmission of TB in the United States and other high-income countries, the unreliable protection afforded by BCG, and its impact on the TST, the vaccine is not recommended for general use. HIV-infected adults and children should not receive BCG vaccine. Moreover, infants whose HIV status is unknown but who have signs and symptoms consistent with HIV infection or who are born to HIV-infected mothers should not receive BCG.

Over the past decade, renewed research and development efforts have been made toward a new TB vaccine. In mid-2014, 16 candidates were in clinical trials and 12 were being field tested. The first new vaccine, for which results of a clinical trial became available in early 2013, is MVA85A/AERAS-485; unfortunately, this viral-vectored vaccine did not show clinical benefit as a booster to BCG.

TREATMENT LATENT TUBERCULOSIS INFECTION

It is estimated that about 2 billion people, or nearly one-third of the human population, have been infected with *M. tuberculosis*. Although only a small fraction of these infections will progress toward active disease, new active cases will continue to emerge from this pool of “latently” infected individuals. Unfortunately, there is no diagnostic test at present that can predict which individuals with LTBI will develop active TB. Treatment of selected persons with LTBI aims at preventing active disease. This intervention (also called *preventive chemotherapy* or *chemoprophylaxis*) is based on the results of a large number of randomized, placebo-controlled clinical trials demonstrating that a 6- to 9-month course of isoniazid reduces the risk of active TB in infected people by up to 90%. Analysis of available data indicates that the optimal duration of treatment is ~9 months. In the absence of reinfection, the protective effect is believed to be lifelong. Clinical trials have shown that isoniazid reduces rates of TB among TST-positive persons with HIV infection. Studies in HIV-infected patients have also demonstrated the effectiveness of shorter courses of rifampin-based treatment.

Candidates for treatment of LTBI are listed in Table 202-5. They can be identified by TST or IGRA of persons in defined high-risk groups. For skin testing, 5 tuberculin units of polysorbate-stabilized PPD should be injected intradermally into the volar surface of the forearm (i.e., the Mantoux method). Multipuncture tests are not recommended. Reactions are read at 48–72 h as the transverse

TABLE 202-5 TUBERCULIN REACTION SIZE AND TREATMENT OF LATENT MYCOBACTERIUM TUBERCULOSIS INFECTION

Risk Group	Tuberculin Reaction Size, mm
HIV-infected persons	≥5
Recent contacts of a patient with TB	≥5 ^a
Organ transplant recipients	≥5
Persons with fibrotic lesions consistent with old TB on chest radiography	≥5
Persons who are immunosuppressed, e.g., due to the use of glucocorticoids or tumor necrosis factor α inhibitors	≥5
Persons with high-risk medical conditions ^b	≥5
Recent immigrants (≤5 years) from high-prevalence countries	≥10
Injection drug users	≥10
Mycobacteriology laboratory personnel; residents and employees of high-risk congregate settings ^c	≥10
Children <5 years of age; children and adolescents exposed to adults in high-risk categories	≥10
Low-risk persons ^d	≥15

^aTuberculin-negative contacts, especially children, should receive prophylaxis for 2–3 months after contact ends and should then undergo repeat TST. Those whose results remain negative should discontinue prophylaxis. HIV-infected contacts should receive a full course of treatment regardless of TST results. ^bThese conditions include silicosis and end-stage renal disease managed by hemodialysis. ^cThese settings include correctional facilities, nursing homes, homeless shelters, and hospitals and other health care facilities. ^dExcept for employment purposes where longitudinal TST screening is anticipated, TST is not indicated for these low-risk persons. A decision to treat should be based on individual risk/benefit considerations.

Source: Adapted from Centers for Disease Control and Prevention: TB elimination—treatment options for latent tuberculosis infection (2011). Available at <http://www.cdc.gov/tb/publications/factsheets/testing/skintestresults.pdf>.

diameter (in millimeters) of induration; the diameter of erythema is not considered. In some persons, TST reactivity wanes with time but can be recalled by a second skin test administered ≥ 1 week after the first (i.e., two-step testing). For persons periodically undergoing the TST, such as health care workers and individuals admitted to long-term-care institutions, initial two-step testing may preclude subsequent misclassification of those who have boosted reactions as TST converters. The cutoff for a positive TST (and thus for treatment) is related both to the probability that the reaction represents true infection and to the likelihood that the individual, if truly infected, will develop TB. Table 202-5 suggests possible cutoff by risk group. Thus, positive reactions for persons with HIV infection, recent close contacts of infectious cases, organ transplant recipients, previously untreated persons whose chest radiograph shows fibrotic lesions consistent with old TB, and persons receiving drugs that suppress the immune system are defined as an area of induration ≥ 5 mm in diameter. A 10-mm cutoff is used to define positive reactions in most other at-risk persons. For persons with a very low risk of developing TB if infected, a cutoff of 15 mm is used. (Except for employment purposes where longitudinal screening is anticipated, the TST is not indicated for these low-risk persons.) A positive IGRA is based on the manufacturers' recommendations; however, good clinical practice requires that epidemiologic and clinical factors also guide the decision to implement treatment for LTBI and that active TB be definitively excluded before the initiation of chemoprophylaxis.

Some TST- and IGRA-negative individuals are also candidates for treatment. Once an appropriate clinical evaluation has excluded active TB, infants and children who have come into contact with infectious cases should be treated for presumed LTBI. HIV-infected persons who have been exposed to an infectious TB patient should receive treatment regardless of the TST result. Any HIV-infected candidate for LTBI treatment must be screened carefully to exclude active TB, which would necessitate full treatment. The use of a clinical algorithm based on four symptoms (current cough, fever, weight loss, and night sweats) helps to define which HIV-infected person is a candidate for LTBI treatment. The absence of all four symptoms tends to exclude active TB. The presence of one of the four symptoms, on the other hand, warrants further investigation for active TB before treatment of LTBI is started. Although administering a TST is prudent, this test is not an absolute requirement—given the logistical challenges—among people living with HIV in high-TB-incidence and low-resource settings.

Before treatment of LTBI begins, it is mandatory to carefully exclude active TB. Several regimens can be used to treat LTBI. The most widely used is that based on isoniazid alone at a daily dose of 5 mg/kg (up to 300 mg/d) for 9 months. On the basis of cost-benefit analyses and concerns about feasibility, a 6-month period of treatment is currently recommended by the WHO, especially in highly TB-endemic countries. Isoniazid can be administered intermittently (twice weekly) at a dose of 15 mg/kg (up to 900 mg) but only as directly observed therapy. An alternative regimen for adults is 4 months of daily rifampin. A 3-month regimen of daily isoniazid and rifampin is used in some countries (e.g., the United Kingdom) for both adults and children who are known not to have HIV infection. A previously recommended regimen of 2 months of rifampin and pyrazinamide has been associated with serious or even fatal hepatotoxicity and now is generally not recommended. The rifampin-containing regimens should be considered for persons who are likely to have been infected with an isoniazid-resistant strain. A recent clinical trial showed that a regimen of isoniazid (900 mg) and rifapentine (900 mg) given once weekly for 12 weeks is as effective as the standard 9-month isoniazid regimen. This regimen was associated with higher treatment completion (82% vs 69%) and less hepatotoxicity (0.4% vs 2.7%) than isoniazid alone, although the rate of permanent discontinuation due to an adverse event was higher (4.9% vs 3.7%).

Currently, the isoniazid-rifapentine regimen is not recommended for children < 2 years of age, people living with HIV infection who are receiving ART, or pregnant women. Rifampin and

rifapentine are contraindicated in HIV-infected individuals receiving protease inhibitors and most NNRTIs. (Efavirenz is the safest agent in this class of antiretrovirals for simultaneous administration with a rifamycin.) Clinical trials to assess the efficacy of long-term isoniazid administration (i.e., for at least 3 years) among people living with HIV in high-TB-transmission settings have shown that this regimen can be more effective than 9 months of isoniazid and is therefore recommended under those circumstances. Isoniazid should not be given to persons with active liver disease. All persons at increased risk of hepatotoxicity (e.g., those abusing alcohol daily and those with a history of liver disease) should undergo baseline and then monthly assessment of liver function. All patients should be carefully educated about hepatitis and instructed to discontinue use of the drug immediately should any symptoms develop. Moreover, patients should be seen and questioned monthly during therapy about adverse reactions and should be given no more than a 1-month supply of drug at each visit. Treatment of LTBI among persons likely to have been infected by a multidrug-resistant strain is a challenge because no regimens have yet been tested in clinical trials. Close observation for early signs of disease is one option; consultation with a TB expert is advised.

It may be more difficult to ensure compliance when treating persons with LTBI than when treating those with active TB. If family members of active cases are being treated, compliance and monitoring may be easier. When feasible, supervised therapy may increase the likelihood of completion. As in active cases, the provision of incentives may also be helpful.

PRINCIPLES OF TB CONTROL

The highest priority in any TB control program is the prompt detection of cases and the provision of short-course chemotherapy to all TB patients under proper case-management conditions, including directly observed therapy. In addition, screening of high-risk groups, including immigrants from high-prevalence countries, migratory workers, prisoners, homeless individuals, substance abusers, and HIV-seropositive persons, is recommended. TST-positive high-risk persons should be treated for LTBI as described above. Contact investigation is an important component of efficient TB control. In the United States and other countries worldwide, a great deal of attention has been given to the transmission of TB (particularly in association with HIV infection) in institutional settings such as hospitals, homeless shelters, and prisons. Measures to limit such transmission include respiratory isolation of persons with suspected TB until they are proven to be noninfectious (at least by sputum AFB smear negativity), proper ventilation in rooms of patients with infectious TB, use of ultraviolet irradiation in areas of increased risk of TB transmission, and periodic screening of personnel who may come into contact with known or unsuspected cases of TB. In the past, radiographic surveys, especially those conducted with portable equipment and miniature films, were advocated for case finding. Today, however, the prevalence of TB in industrialized countries is sufficiently low that "mass miniature radiography" is not cost-effective.

In high-prevalence countries, most TB control programs have made remarkable progress in reducing morbidity and mortality since the mid-1990s by adopting and implementing the strategy promoted by the WHO. Between 2000 and 2013, 37 million lives were saved, and since 1995, 61 million TB cases have been successfully treated. The essential elements of good TB care and control (the DOTS strategy) are (1) political commitment with increased and sustained financing; (2) case detection through quality-assured bacteriology (starting with examination of sputum from patients with cough of > 2 –3 weeks' duration); (3) administration of standardized short-course chemotherapy, with direct supervision and patient support; (4) an effective drug supply and management system; and (5) a monitoring and evaluation system, with impact measurement (including assessment of treatment outcomes—e.g., cure, completion of treatment without bacteriologic proof of cure, death, treatment failure, and default—in all cases

1122 registered and notified). In 2006, the WHO indicated that, although these essential elements remain the fundamental components of any control strategy, additional steps must be undertaken to reach the 2015 international TB control targets set within the United Nations Millennium Development Goals. Thus, a new “Stop TB Strategy” with six components has been promoted since 2006: (1) Pursue high-quality DOTS expansion and enhancement. (2) Address HIV-associated TB, MDR-TB, and the needs of poor and vulnerable populations. (3) Contribute to health system strengthening. (4) Engage all care providers. (5) Empower people with TB and their communities. (6) Enable and promote research. As part of the fourth component, evidence-based International Standards for Tuberculosis Care—focused on diagnosis, treatment, and public health responsibilities—have been introduced for wide adoption by medical and professional societies, academic institutions, and all practitioners worldwide (http://www.who.int/tb/publications/ISTC_3rdEd.pdf?ua=1).

Care and control of HIV-associated TB are particularly challenging in developing countries because existing interventions require collaboration between HIV/AIDS and TB programs as well as standard services. While TB programs must test every patient for HIV in order to provide access to trimethoprim-sulfamethoxazole prophylaxis against common infections and ART, HIV/AIDS programs must regularly screen persons living with HIV/AIDS for active TB, provide treatment for LTBI, and ensure infection control in settings where people living with HIV congregate.

Early and active case detection is considered an important intervention not only among persons living with HIV/AIDS but also among other vulnerable populations, as it reduces transmission in a community and provides early effective care. For TB control efforts to succeed and for elimination to become a realistic target, programs must optimize their performance and include additional interventions as described. Moreover, the approach to TB control and care needs to become holistic and engage beyond dedicated programs. Therefore, the WHO’s “End TB” strategy has been designed and builds on three pillars for the post-2015 era of increased efforts by governments and national programs worldwide: (1) integrated, patient-centered care and prevention; (2) bold policies and supportive systems; and (3) intensified research and innovation. The first pillar incorporates all technological innovations, such as early diagnostic approaches (including universal drug-susceptibility testing and systematic screening of identified, setting-specific, high-risk groups); well-designed treatment regimens for all forms of TB; proper management of HIV-associated TB and other comorbidities; and preventive treatment of persons at high risk. The second pillar is fundamental and is normally beyond the control of dedicated programs, relying on policies forged by the highest-level health and governmental authorities: availability of adequate and well-identified human and financial resources; engagement of civil society organizations and all relevant public and private providers to facilitate care and prevention of all patients; a policy of universal health coverage (which implies avoidance of catastrophic expenditures caused by TB among the poorest); regulatory frameworks for case notifications, vital registration, quality and rational use of medicines, and infection control; social protection mechanisms; poverty alleviation strategies; and interventions on the broader determinants of TB. Finally, the third pillar of the new strategy emphasizes intensification of engagement in research and development of new tools and interventions as well as optimization of implementation and rapid adoption of new tools in endemic countries. In the end, besides specific clinical care and control interventions as described in this chapter, elimination of TB ultimately will require control and attenuation of the multitude of risk factors (e.g., HIV, smoking, and diabetes) and socioeconomic determinants (e.g., extreme poverty, inadequate living conditions and bad housing, alcoholism, malnutrition, and indoor air pollution) with clearly implemented policies within the health sector and other sectors linked to human development and welfare.

ACKNOWLEDGMENT

The contributions of Richard J. O’Brien to this chapter in previous editions are gratefully acknowledged.

Leprosy, first described in ancient Indian texts from the sixth century B.C., is a nonfatal, chronic infectious disease caused by *Mycobacterium leprae*, the clinical manifestations of which are largely confined to the skin, peripheral nervous system, upper respiratory tract, eyes, and testes. The unique tropism of *M. leprae* for peripheral nerves (from large nerve trunks to microscopic dermal nerves) and certain immunologically mediated reactional states are the major causes of morbidity in leprosy. The propensity of the disease, when untreated, to result in characteristic deformities and the recognition in most cultures that the disease is communicable from person to person have resulted historically in a profound social stigma. Today, with early diagnosis and the institution of appropriate and effective antimicrobial therapy, patients can lead productive lives in the community, and deformities and other visible manifestations can largely be prevented.

ETIOLOGY

M. leprae is an obligate intracellular bacillus (0.3–1 μm wide and 1–8 μm long) that is confined to humans, armadillos in certain locales, and sphagnum moss. The organism is acid-fast, indistinguishable microscopically from other mycobacteria, and ideally detected in tissue sections by a modified Fite stain. Strain variability has been documented in this organism. *M. leprae* produces no known toxins and is well adapted to penetrate and reside within macrophages, yet it may survive outside the body for months. In untreated patients, only ~1% of *M. leprae* organisms are viable. The morphologic index (MI), a measure of the number of acid-fast bacilli (AFB) in skin scrapings that stain uniformly bright, correlates with viability. The bacteriologic index (BI), a logarithmic-scaled measure of the density of *M. leprae* in the dermis, may be as high as 4–6+ in untreated patients and falls by 1 unit per year during effective antimicrobial therapy; the rate of decrease is independent of the relative potency of therapy. A rising MI or BI suggests relapse and perhaps—if the patient is being treated—drug resistance. Drug resistance can be confirmed or excluded in the mouse model of leprosy, and resistance to dapsone and rifampin can be documented by the recognition of mutant genes. However, the availability of these technologies is extremely limited.



As a result of reductive evolution, almost half of the *M. leprae* genome contains nonfunctional genes; only 1605 genes encode for proteins, and 1439 genes are shared with *Mycobacterium tuberculosis*. In contrast, *M. tuberculosis* uses 91% of its genome to encode for 4000 proteins. Among the lost genes in *M. leprae* are those for catabolic and respiratory pathways; transport systems; purine, methionine, and glutamine synthesis; and nitrogen regulation. The genome of *M. leprae* provides a metabolic rationale for its obligate intracellular existence and reliance on host biochemical support, a template for targets of drug development, and ultimately a pathway to cultivation. The finding of strain variability among *M. leprae* isolates has provided a powerful tool with which to address anew the organism’s epidemiology and pathobiology and to determine whether relapse represents reactivation or reinfection. The bacterium’s complex cell wall contains large amounts of an *M. leprae*-specific phenolic glycolipid (PGL-1), which is detected in serologic tests. The unique trisaccharide of *M. leprae* binds to the basal lamina of Schwann cells; this interaction is probably relevant to the fact that *M. leprae* is the only bacterium to invade peripheral nerves.

Although it was the first bacterium to be etiologically associated with human disease, *M. leprae* remains one of the few bacterial species that still has not been cultivated on artificial medium or tissue culture. The multiplication of *M. leprae* in mouse footpads (albeit limited, with a doubling time of ~2 weeks) has provided a means to evaluate antimicrobial agents, monitor clinical trials, and screen vaccines. *M. leprae* grows best in cooler tissues (the skin, peripheral nerves, anterior chamber of the eye, upper respiratory tract, and testes), sparing warmer areas of the skin (the axilla, groin, scalp, and midline of the back).

EPIDEMIOLOGY



Demographics Leprosy is almost exclusively a disease of the developing world, affecting areas of Asia, Africa, Latin America, and the Pacific. While Africa has the highest disease prevalence, Asia has the most cases. More than 80% of the world's cases occur in a few countries: India, China, Myanmar, Indonesia, Brazil, Nigeria, Madagascar, and Nepal. Within endemic locales, the distribution of leprosy is quite uneven, with areas of high prevalence bordering on areas with little or no disease. In Brazil the majority of cases occur in the Amazon basin and two western states, while in Mexico leprosy is mostly confined to the Pacific coast. Except as imported cases, leprosy is largely absent from the United States, Canada, and northwestern Europe. In the United States, ~4000 persons have leprosy and 100–200 new cases are reported annually, most of them in California, Texas, New York, and Hawaii among immigrants from Mexico, Southeast Asia, the Philippines, and the Caribbean.



The comparative genomics of single-nucleotide polymorphisms support the likelihood that four distinct strains exist, having originated in East Africa or Central Asia. A mutation spread to Europe and subsequently underwent two separate mutations that were then followed by spread to West Africa and the Americas.

The global prevalence of leprosy is difficult to assess, given that many of the locales with high prevalence lack a significant medical or public health infrastructure. Estimates range from 0.6 to 8 million affected individuals. The lower estimate includes only persons who have not completed chemotherapy, excluding those who may be physically or psychologically damaged from leprosy and who may yet relapse or develop immune-mediated reactions. The higher figure includes patients whose infections probably are already cured and many who have no leprosy-related deformity or disability. Although the figures on the worldwide prevalence of leprosy are debatable, incidence is not falling; there are still an estimated 500,000 new cases annually.

Leprosy is associated with poverty and rural residence. It appears not to be associated with AIDS, perhaps because of leprosy's long incubation period. Most individuals appear to be naturally immune to leprosy and do not develop disease manifestations after exposure. The time of peak onset is in the second and third decades of life.



The most severe lepromatous form of leprosy is twice as common among men as among women and is rarely encountered in children. The frequency of the polar forms of leprosy in different countries varies widely and may in part be genetically determined; certain human leukocyte antigen (HLA) associations are known for

both polar forms of leprosy (see below). Furthermore, variations in immunoregulatory genes are associated with an increased susceptibility to leprosy, particularly the multibacillary form. In India and Africa, 90% of cases are tuberculoid; in Southeast Asia, 50% are tuberculoid and 50% lepromatous; and in Mexico, 90% are lepromatous. (For definitions of disease types, see [Table 203-1](#) and “Clinical, Histologic, and Immunologic Spectrum,” below.)

Transmission The route of transmission of leprosy remains uncertain, and transmission routes may in fact be multiple. Nasal droplet infection, contact with infected soil, and even insect vectors have been considered the prime candidates. Aerosolized *M. leprae* can cause infection in immunosuppressed mice, and a sneeze from an untreated lepromatous patient may contain $>10^{10}$ AFB. Furthermore, both IgA antibody to *M. leprae* and genes of *M. leprae*—demonstrable by polymerase chain reaction (PCR)—have been found in the nose of individuals from endemic areas who have no signs of leprosy and in 19% of occupational contacts of lepromatous patients. Several lines of evidence implicate soil transmission. (1) In endemic countries such as India, leprosy is primarily a rural and not an urban disease. (2) *M. leprae* products reside in soil in endemic locales. (3) Direct dermal inoculation (e.g., during tattooing) may transmit *M. leprae*, and common sites of leprosy in children are the buttocks and thighs, suggesting that microinoculation of infected soil may transmit the disease. Evidence for insect vectors of leprosy includes the demonstration that bedbugs and mosquitoes in the vicinity of leprosia regularly harbor *M. leprae* and that experimentally infected mosquitoes can transmit the infection to mice. Skin-to-skin contact generally is not considered an important route of transmission.

In endemic countries, ~50% of leprosy patients have a history of intimate contact with an infected person (often a household member), while, for unknown reasons, leprosy patients in nonendemic locales can identify such contact only 10% of the time. Moreover, household contact with an infected lepromatous case carries an eventual risk of disease acquisition of ~10% in endemic areas as opposed to only 1% in nonendemic locales. Contact with a tuberculoid case carries a very low risk. Physicians and nurses caring for leprosy patients and the co-workers of these patients are not at risk for leprosy.



Although multilocus variable-number short-nucleotide tandem-repeat (VNTR) analyses have generally demonstrated considerable variability among isolates, highly similar and even identical VNTR results have been obtained with isolates from a limited number

TABLE 203-1 CLINICAL, BACTERIOLOGIC, PATHOLOGIC, AND IMMUNOLOGIC SPECTRUM OF LEPROSY

Feature	Tuberculoid (TT, BT) Leprosy	Borderline (BB, BL) Leprosy	Lepromatous (LL) Leprosy
Skin lesions	One or a few sharply defined annular asymmetric macules or plaques with a tendency toward central clearing, elevated borders	Intermediate between BT- and LL-type lesions; ill-defined plaques with an occasional sharp margin; few or many in number	Symmetric, poorly margined, multiple infiltrated nodules and plaques or diffuse infiltration; xanthoma-like or dermatofibroma papules; leonine facies and eyebrow alopecia
Nerve lesions	Skin lesions anesthetic early; nerve near lesions sometimes enlarged; nerve abscesses most common in BT	Hypesthetic or anesthetic skin lesions; nerve trunk palsies, at times symmetric	Hypesthesia a late sign; nerve palsies variable; acral, distal, symmetric anesthesia common
Acid-fast bacilli (BI) ^a	0–1+	3–5+	4–6+
Lymphocytes	2+	1+	0–1+
Macrophage differentiation	Epithelioid	Epithelioid in BB; usually undifferentiated but may have foamy changes in BL	Foamy change the rule; may be undifferentiated in early lesions
Langerhans giant cells	1–3+	—	—
Lepromin skin test	+++	—	—
Lymphocyte transformation test	Generally positive	1–10%	1–2%
CD4+/CD8+ T cell ratio in lesions	1.2	BB: NT; BL: 0.48	0.50
<i>M. leprae</i> PGL-1 antibodies	60%	85%	95%

^aSee text.

Abbreviations: BB, mid-borderline; BL, borderline lepromatous; BT, borderline tuberculoid; TT, polar tuberculoid; LL, polar lepromatous; BI, bacteriologic index; NT, not tested; PGL-1, phenolic glycolipid 1.

1124 of families with multiple cases. Moreover, VNTR results have been similar for isolates within certain geographic locales and divergent for isolates within others. These findings suggest that genomic analyses may prove useful in the future for defining *M. leprae* transmission patterns.

M. leprae causes disease primarily in humans. However, in Texas and Louisiana, 15% of nine-banded armadillos are infected, and armadillo contact occasionally results in human disease. Armadillos develop disseminated infection after IV inoculation of live *M. leprae*.

CLINICAL, HISTOLOGIC, AND IMMUNOLOGIC SPECTRUM

The incubation period prior to manifestation of clinical disease can vary between 2 and 40 years, although it is generally 5–7 years in duration. This long incubation period is probably, at least in part, a consequence of the extremely long doubling time for *M. leprae* (14 days in mice versus in vitro doubling times of 1 day and 20 min for *M. tuberculosis* and *Escherichia coli*, respectively). Leprosy presents as a spectrum of clinical manifestations that have bacteriologic, pathologic, and immunologic counterparts. The spectrum from polar tuberculoid (TT) to borderline tuberculoid (BT) to mid-borderline (BB, which is rarely encountered) to borderline lepromatous (BL) to polar lepromatous (LL) disease is associated with an evolution from asymmetric localized macules and plaques to nodular and indurated symmetric generalized skin manifestations, an increasing bacterial load, and loss of *M. leprae*-specific cellular immunity (Table 203-1). Distinguishing dermatopathologic characteristics include the number of lymphocytes, giant cells, and AFB as well as the nature of epithelioid cell differentiation. Where a patient presents on the clinical spectrum largely determines prognosis, complications, reactional states, and the intensity of antimicrobial therapy required.

Tuberculoid Leprosy At the less severe end of the spectrum is tuberculoid leprosy, which encompasses TT and BT disease. In general, these forms of leprosy result in symptoms confined to the skin and peripheral nerves. TT leprosy is the most common form of the disease encountered in India and Africa but is virtually absent in Southeast Asia, where BT leprosy is frequent.

The skin lesions of tuberculoid leprosy consist of one or a few hypopigmented macules or plaques (Fig. 203-1) that are sharply demarcated and hypesthetic, often have erythematous or raised borders, and are devoid of the normal skin organs (sweat glands and hair follicles) and thus are dry, scaly, and anhidrotic. AFB are generally absent or few in number. Tuberculoid leprosy patients may have asymmetric enlargement of one or a few peripheral nerves. Indeed, leprosy and certain rare hereditary neuropathies are the only human diseases associated with peripheral-nerve enlargement. Although any peripheral nerve may be enlarged (including small digital and supraclavicular nerves), those most commonly affected are the ulnar, posterior auricular, peroneal, and posterior tibial nerves, with associated hypesthesia and myopathy.



FIGURE 203-1 Tuberculoid (TT) leprosy: a well-defined, hypopigmented, anesthetic macule with anhidrosis and a raised granular margin (arrowhead).

In tuberculoid leprosy, T cells breach the perineurium, and destruction of Schwann cells and axons may be evident, resulting in fibrosis of the epineurium, replacement of the endoneurium with epithelial granulomas, and occasionally caseous necrosis. Such invasion and destruction of nerves in the dermis by T cells are pathognomonic for leprosy.

Circulating lymphocytes from patients with tuberculoid leprosy readily recognize *M. leprae* and its constituent proteins, patients have positive lepromin skin tests (see “Diagnosis,” below), and—owing to a type 1 cytokine pattern in tuberculoid tissues—strong T cell and macrophage activation results in a localized infection. In tuberculoid leprosy tissue, there is a 2:1 predominance of helper CD4+ over CD8+ T lymphocytes. Tuberculoid tissues are rich in the mRNAs of the pro-inflammatory T_H1 family of cytokines: interleukin (IL) 2, interferon γ (IFN- γ), and IL-12; in contrast, IL-4, IL-5, and IL-10 mRNAs are scarce.

Lepromatous Leprosy Lepromatous leprosy patients present with symmetrically distributed skin nodules (Fig. 203-2), raised plaques, or diffuse dermal infiltration, which, when on the face, results in *leonine facies*. Late manifestations include loss of eyebrows (initially the lateral margins only) and eyelashes, pendulous earlobes, and dry scaling skin, particularly on the feet. In LL leprosy, bacilli are numerous in the skin (as many as 10⁹/g), where they are often found in large clumps (*globi*), and in peripheral nerves, where they initially invade Schwann cells, resulting in foamy degenerative myelination and axonal degeneration and later in Wallerian degeneration. In addition, bacilli are plentiful in circulating blood and in all organ systems except the lungs and the central nervous system. Nevertheless, patients are afebrile, and there is no evidence of major organ system dysfunction.

Found almost exclusively in western Mexico and the Caribbean is a form of lepromatous leprosy without visible skin lesions but with diffuse dermal infiltration and a demonstrably thickened dermis, termed *diffuse lepromatosis*.

In lepromatous leprosy, nerve enlargement and damage tend to be symmetric, result from actual bacillary invasion, and are more insidious but ultimately more extensive than in tuberculoid leprosy. Patients with LL leprosy have acral, distal, symmetric peripheral neuropathy and a tendency toward symmetric nerve-trunk enlargement. They may also have signs and symptoms related to involvement of the upper respiratory tract, the anterior chamber of the eye, and the testes.

In untreated LL patients, lymphocytes regularly fail to recognize either *M. leprae* or its protein constituents, and lepromin skin tests are negative (see “Diagnosis,” below). This loss of protective cellular immunity appears to be antigen-specific, as patients are not unusually



FIGURE 203-2 Lepromatous (LL) leprosy: advanced nodular lesions.

susceptible to opportunistic infections, cancer, or AIDS and maintain delayed-type hypersensitivity to *Candida*, *Trichophyton*, mumps virus, tetanus toxoid, and even purified protein derivative of tuberculin. At times, *M. leprae*-specific anergy is reversible with effective chemotherapy. In LL tissues, there is a 2:1 ratio of CD8+ to CD4+ T lymphocytes. LL patients have a predominant T_H2 response and hyperglobulinemia, and LL tissues demonstrate a T_H2 cytokine profile, being rich in mRNAs for IL-4, IL-5, and IL-10 and poor in those for IL-2, IFN- γ , and IL-12. It appears that cytokines mediate a protective tissue response in leprosy, as injection of IFN- γ or IL-2 into lepromatous lesions causes a loss of AFB and histopathologic conversion toward a tuberculoid pattern. Macrophages of lepromatous leprosy patients appear to be functionally intact; circulating monocytes exhibit normal microbicidal function and responsiveness to IFN- γ .

Reactional States Leprea reactions comprise several common immunologically mediated inflammatory states that cause considerable morbidity. Some of these reactions precede diagnosis and the institution of effective antimicrobial therapy; indeed, these reactions may precipitate presentation for medical attention and diagnosis. Other reactions follow the initiation of appropriate chemotherapy; these reactions may cause patients to perceive that their leprosy is worsening and to lose confidence in conventional therapy. Only by warning patients of the potential for these reactions and describing their manifestations can physicians treating leprosy patients ensure continued credibility.

TYPE 1 LEPROA REACTIONS (DOWNGRADING AND REVERSAL REACTIONS) Type 1 lepra reactions occur in almost half of patients with borderline forms of leprosy but not in patients with pure lepromatous disease. Manifestations include classic signs of inflammation within previously involved macules, papules, and plaques and, on occasion, the appearance of new skin lesions, neuritis, and (less commonly) fever—generally low-grade. The nerve trunk most frequently involved in this process is the ulnar nerve at the elbow, which may be painful and exquisitely tender. If patients with affected nerves are not treated promptly with glucocorticoids (see below), irreversible nerve damage may result in as little as 24 h. The most dramatic manifestation is foot-drop, which occurs when the peroneal nerve is involved.

When type 1 lepra reactions precede the initiation of appropriate antimicrobial therapy, they are termed *downgrading reactions*, and the case becomes histologically more lepromatous; when they occur after the initiation of therapy, they are termed *reversal reactions*, and the case becomes more tuberculoid. Reversal reactions often occur in the first months or years after the initiation of therapy but may also develop several years thereafter.

Edema is the most characteristic microscopic feature of type 1 lepra lesions, whose diagnosis is primarily clinical. Reversal reactions are typified by a T_H1 cytokine profile, with an influx of CD4+ T helper cells and increased levels of IFN- γ and IL-2. In addition, type 1 reactions are associated with large numbers of T cells bearing γ/δ receptors—a unique feature of leprosy.

TYPE 2 LEPROA REACTIONS: ERYTHEMA NODOSUM LEPROSUM Erythema nodosum leprosum (ENL) (Fig. 203-3) occurs exclusively in patients near the lepromatous end of the leprosy spectrum (BL/LL), affecting nearly 50% of this group. Although ENL may precede leprosy diagnosis and the initiation of therapy (sometimes, in fact, prompting the diagnosis), in 90% of cases it follows the institution of chemotherapy, generally within 2 years. The most common features of ENL are crops of painful erythematous papules that resolve spontaneously in a few days to a week but may recur; malaise; and fever that can be profound. However, patients may also experience symptoms of neuritis, lymphadenitis, uveitis, orchitis, and glomerulonephritis and may develop anemia, leukocytosis, and abnormal liver function tests (particularly increased aminotransferase levels). Individual patients may have either a single bout of ENL or chronic recurrent manifestations. Bouts may be either mild or severe and generalized; in rare instances, ENL results in death. Skin biopsy of ENL papules reveals vasculitis or panniculitis, sometimes with many lymphocytes but characteristically with polymorphonuclear leukocytes as well.



FIGURE 203-3 Moderately severe skin lesions of erythema nodosum leprosum, some with pustulation and ulceration.

Elevated levels of circulating tumor necrosis factor (TNF) have been demonstrated in ENL; thus, TNF may play a central role in the pathobiology of this syndrome. ENL is thought to be a consequence of immune complex deposition, given its T_H2 cytokine profile and its high levels of IL-6 and IL-8. However, in ENL tissue, the presence of HLA-DR framework antigen of epidermal cells—considered a marker for a delayed-type hypersensitivity response—and evidence of higher levels of IL-2 and IFN- γ than are usually seen in polar lepromatous disease suggest an alternative mechanism.

LUCIO'S PHENOMENON Lucio's phenomenon is an unusual reaction seen exclusively in patients from the Caribbean and Mexico who have the diffuse lepromatous form of lepromatous leprosy, most often those who are untreated. Patients with this reaction develop recurrent crops of large, sharply marginated, ulcerative lesions—particularly on the lower extremities—that may be generalized and, when so, are frequently fatal as a result of secondary infection and consequent septic bacteremia. Histologically, the lesions are characterized by ischemic necrosis of the epidermis and superficial dermis, heavy parasitism of endothelial cells with AFB, and endothelial proliferation and thrombus formation in the larger vessels of the deeper dermis. Like ENL, Lucio's phenomenon is probably mediated by immune complexes.

Complications • THE EXTREMITIES Complications of the extremities in leprosy patients are primarily a consequence of neuropathy leading to insensitivity and myopathy. Insensitivity affects fine touch, pain, and heat receptors but generally spares position and vibration appreciation. The most commonly affected nerve trunk is the ulnar nerve at the elbow, whose involvement results in clawing of the fourth and fifth fingers, loss of dorsal interosseous musculature in the affected hand, and loss of sensation in these distributions. Median nerve involvement in leprosy impairs thumb opposition and grasp; radial nerve dysfunction, although rare in leprosy, leads to wristdrop. Tendon transfers can restore hand function but should not be performed until 6 months after the initiation of antimicrobial therapy and the conclusion of episodes of acute neuritis.

Plantar ulceration, particularly at the metatarsal heads, is probably the most common complication of leprosy neuropathy. Therapy requires careful debridement; administration of appropriate antibiotics; avoidance of weight-bearing until ulcerations are healed, with slowly progressive ambulation thereafter; and wearing of special shoes to prevent recurrence.

Footdrop as a result of peroneal nerve palsy should be treated with a simple nonmetallic brace in the shoe or with surgical correction attained by tendon transfers. Although uncommon, Charcot's joints, particularly of the foot and ankle, may result from leprosy.

The loss of distal digits in leprosy is a consequence of insensitivity, trauma, secondary infection, and—in lepromatous disease—a poorly understood and sometimes profound osteolytic process. Conscientious

1126 protection of the extremities during cooking and work and the early institution of therapy have substantially reduced the frequency and severity of distal digit loss in recent times.

THE NOSE In lepromatous leprosy, bacillary invasion of the nasal mucosa can result in chronic nasal congestion and epistaxis. Saline nose drops may relieve these symptoms. Long-untreated LL leprosy may further result in destruction of the nasal cartilage, with consequent saddle-nose deformity or anosmia (more common in the pre-antibiotic era than at present). Nasal reconstructive procedures can ameliorate significant cosmetic defects.

THE EYE Owing to cranial nerve palsies, lagophthalmos and corneal insensitivity may complicate leprosy, resulting in trauma, secondary infection, and (without treatment) corneal ulcerations and opacities. For patients with these conditions, eyedrops during the day and ointments at night provide some protection from such consequences. Furthermore, in LL leprosy, the anterior chamber of the eye is invaded by bacilli, and ENL may result in uveitis, with consequent cataracts and glaucoma. Thus leprosy is a major cause of blindness in the developing world. Slit-lamp evaluation of LL patients often reveals “corneal beading,” representing globi of *M. leprae*.

THE TESTES *M. leprae* invades the testes, while ENL may cause orchitis. Thus males with lepromatous leprosy often manifest mild to severe testicular dysfunction, with an elevation of luteinizing and follicle-stimulating hormones, decreased testosterone, and aspermia or hypospermia in 85% of LL patients but in only 25% of BL patients. LL patients may become impotent and infertile. Impotence is sometimes responsive to testosterone replacement.

AMYLOIDOSIS Secondary amyloidosis is a complication of LL leprosy and ENL that is encountered infrequently in the antibiotic era. This complication may result in abnormalities of hepatic and particularly renal function.

NERVE ABSCESSSES Patients with various forms of leprosy, but particularly those with the BT form, may develop abscesses of nerves (most commonly the ulnar), with a cellulitic appearance of adjacent skin. In such conditions, the affected nerve is swollen and exquisitely tender. Although glucocorticoids may reduce signs of inflammation, rapid surgical decompression is necessary to prevent irreversible sequelae.

DIAGNOSIS

Leprosy most commonly presents with both characteristic skin lesions and skin histopathology. Thus the disease should be suspected when a patient from an endemic area has suggestive skin lesions or peripheral neuropathy. The diagnosis should be confirmed by histopathology. In tuberculoid leprosy, lesional areas—preferably the advancing edge—must be biopsied because normal-appearing skin does not have pathologic features. In lepromatous leprosy, nodules, plaques, and indurated areas are optimal biopsy sites, but biopsies of normal-appearing skin also are generally diagnostic. Lepromatous leprosy is associated with diffuse hyperglobulinemia, which may result in false-positive serologic tests (e.g., Venereal Disease Research Laboratory, rheumatoid arthritis, and antinuclear antibody tests) and therefore may cause diagnostic confusion. On occasion, tuberculoid lesions may not (1) appear typical, (2) be hypesthetic, and (3) contain granulomas (instead containing only nonspecific lymphocytic infiltrates). In such instances, two of these three characteristics are considered sufficient for a diagnosis. It is preferable to overdiagnose leprosy rather than to allow a patient to remain untreated.

IgM antibodies to PGL-1 are found in 95% of patients with untreated lepromatous leprosy; the titer decreases with effective therapy. However, in tuberculoid leprosy—the form of disease most often associated with diagnostic uncertainty owing to the absence or paucity of AFB—patients have significant antibodies to PGL-1 only 60% of the time; moreover, in endemic locales, exposed individuals without clinical leprosy may harbor antibodies to PGL-1. Thus PGL-1 serology is of little diagnostic utility in tuberculoid leprosy. Heat-killed *M. leprae* (lepromin) has been used as a skin test reagent. It generally elicits a reaction in tuberculoid leprosy patients, may do so in individuals without leprosy, and gives

negative results in lepromatous leprosy patients; consequently, it is likewise of little diagnostic value. Unfortunately, PCR of the skin for *M. leprae*, although positive in LL and BL disease, yields negative results in 50% of tuberculoid cases, again offering little diagnostic assistance.

DIFFERENTIAL DIAGNOSIS

Included in the differential diagnosis of lesions that resemble leprosy are sarcoidosis, leishmaniasis, lupus vulgaris, dermatofibroma, histiocytoma, lymphoma, syphilis, yaws, granuloma annulare, and various other disorders causing hypopigmentation (notably pityriasis alba, tinea, and vitiligo). Sarcoidosis may result in perineural inflammation, but actual granuloma formation within dermal nerves is pathognomonic for leprosy. In lepromatous leprosy, sputum specimens may be loaded with AFB—a finding that can be incorrectly interpreted as representing pulmonary tuberculosis.

TREATMENT LEPROSY

ANTIMICROBIAL THERAPY

Active Agents Established agents used to treat leprosy include dapsone (50–100 mg/d), clofazimine (50–100 mg/d, 100 mg three times weekly, or 300 mg monthly), and rifampin (600 mg daily or monthly; see “Choice of Regimens,” below). Of these drugs, only rifampin is bactericidal. The sulfones (folate antagonists), the foremost of which is dapsone, were the first antimicrobial agents found to be effective for the treatment of leprosy and are still the mainstays of therapy. With sulfone treatment, skin lesions resolve and numbers of viable bacilli in the skin are reduced. Although primarily bacteriostatic, dapsone monotherapy results in a resistance-related relapse rate of only 2.5%; after ≥18 years of therapy and subsequent discontinuation, only another 10% of patients relapse, developing new, usually asymptomatic, shiny, “histoid” nodules. Dapsone is generally safe and inexpensive. Individuals with glucose-6-phosphate dehydrogenase deficiency who are treated with dapsone may develop severe hemolysis; those without this deficiency also have reduced red cell survival and a hemoglobin decrease averaging 1 g/dL. Dapsone’s usefulness is limited occasionally by allergic dermatitis and rarely by the sulfone syndrome (including high fever, anemia, exfoliative dermatitis, and a mononucleosis-type blood picture). It must be remembered that rifampin induces microsomal enzymes, necessitating increased doses of medications such as glucocorticoids and oral birth control regimens. Clofazimine is often cosmetically unacceptable to light-skinned leprosy patients because it causes a red-black skin discoloration that accumulates, particularly in lesional areas, and makes the patient’s diagnosis obvious to members of the community.

Other antimicrobial agents active against *M. leprae* in animal models and at the usual daily doses used in clinical trials include ethionamide/prothionamide; the aminoglycosides streptomycin, kanamycin, and amikacin (but not gentamicin or tobramycin); minocycline; clarithromycin; and several fluoroquinolones, particularly ofloxacin. Next to rifampin, minocycline, clarithromycin, and ofloxacin appear to be most bactericidal for *M. leprae*, but these drugs have not been used extensively in leprosy control programs. Most recently, rifapentine and moxifloxacin have been found to be especially potent against *M. leprae* in mice. In a clinical trial in lepromatous leprosy, moxifloxacin was profoundly bactericidal, matched in potency only by rifampin.

Choice of Regimens Antimicrobial therapy for leprosy must be individualized, depending on the clinical/pathologic form of the disease encountered. Tuberculoid leprosy, which is associated with a low bacterial burden and a protective cellular immune response, is the easiest form to treat and can be cured reliably with a finite course of chemotherapy. In contrast, lepromatous leprosy may have a higher bacillary load than any other human bacterial disease, and the absence of a salutary T cell repertoire requires prolonged or even lifelong chemotherapy. Hence, careful classification of disease prior to therapy is important.

In developed countries, clinical experience with leprosy classification is limited; fortunately, however, the resources needed for skin biopsy are highly accessible, and those for pathologic interpretation are readily available. In developing countries, clinical expertise is greater but is now waning substantially as the care of leprosy patients is integrated into general health services. In addition, access to dermatopathology services is often limited. In such instances, skin smears may prove useful, but in many locales access to the resources needed for their preparation and interpretation also may be unavailable. Use of skin smears is no longer encouraged by the World Health Organization (WHO) and is often replaced by mere counting of lesions, which, together with a lack of capacity for histopathologic assessment, may negatively affect decisions about chemotherapy, increase the potential for reactions, and worsen the ultimate prognosis. A reasoned approach to the treatment of leprosy is confounded by these and several other issues:

1. Even without therapy, TT leprosy may heal spontaneously, and prolonged dapsone monotherapy (even for LL leprosy) is generally curative in 80% of cases.
2. In tuberculoid disease, it is common for no bacilli to be found in the skin prior to therapy, and thus there is no objective measure of therapeutic success. Furthermore, despite adequate treatment, TT and particularly BT lesions often resolve minimally or incompletely, while relapse and late type 1 lepra reactions can be difficult to distinguish.
3. LL leprosy patients commonly harbor viable persistent *M. leprae* organisms after prolonged intensive therapy; the propensity of these organisms to initiate clinical relapse is unclear. Because relapse in LL patients after discontinuation of rifampin-containing regimens usually begins only after 7–10 years, follow-up over the very long term is necessary to assess ultimate clinical outcomes.
4. Even though primary dapsone resistance is exceedingly rare and multidrug therapy is generally recommended (at least for lepromatous leprosy), there is a paucity of information from experimental animals and clinical trials on the optimal combination of antimicrobial agents, dosing schedule, and duration of therapy.

In 1982, the WHO made recommendations for leprosy chemotherapy administered in control programs. These recommendations came on the heels of the demonstration of the relative success of long-term dapsone monotherapy and in the context of concerns about dapsone resistance. Other complicating considerations included the limited resources available for leprosy care in the very areas where it is most prevalent and the frustration and discouragement of patients and program managers with the previous requirement for lifelong therapy for many leprosy patients. Thus, for the first time, the WHO advocated a finite duration of therapy for all forms of leprosy and—given the prohibitive cost of daily rifampin treatment in developing countries—encouraged the monthly administration of this agent as part of a multidrug regimen. Over the ensuing years, the WHO recommendations have been broadly implemented, and the duration of therapy required, particularly for lepromatous leprosy, has been progressively shortened. For treatment purposes, the WHO classifies patients as *paucibacillary* or *multibacillary*. Previously, patients without demonstrable AFB in the dermis were classified as paucibacillary and those with AFB as multibacillary. Currently, in light of the perceived unreliability of skin smears in the field, patients are classified as multibacillary if they have six or more skin lesions and as paucibacillary if they have fewer. (Unfortunately, this classification method has been found wanting, as some patients near the lepromatous pole have only one or a few skin lesions.) The WHO recommends that paucibacillary adults be treated with 100 mg of dapsone daily and 600 mg of rifampin monthly (supervised) for 6 months (Table 203-2). For patients with single-lesion paucibacillary leprosy, the WHO recommends as an alternative a single dose of rifampin (600 mg), ofloxacin (400 mg), and minocycline (100 mg). Multibacillary adults should be treated with 100 mg of dapsone plus 50 mg of clofazimine daily (unsupervised)

TABLE 203-2 ANTIMICROBIAL REGIMENS RECOMMENDED FOR THE TREATMENT OF LEPROSY IN ADULTS

Form of Leprosy	More Intensive Regimen	WHO-Recommended Regimen (1982)
Tuberculoid (paucibacillary)	Dapsone (100 mg/d) for 5 years	Dapsone (100 mg/d, unsupervised) plus rifampin (600 mg/month, supervised) for 6 months
Lepromatous (multibacillary)	Rifampin (600 mg/d) for 3 years plus dapsone (100 mg/d) indefinitely	Dapsone (100 mg/d) plus clofazimine (50 mg/d), unsupervised; and rifampin (600 mg) plus clofazimine (300 mg) monthly (supervised) for 1–2 years

Note: See text for discussion and comparison of the WHO recommendations with the more intensive approach as well as the alternative WHO regimen for single-lesion paucibacillary leprosy.

and with 600 mg of rifampin plus 300 mg of clofazimine monthly (supervised). Originally, the WHO recommended that lepromatous patients be treated for 2 years or until smears became negative (generally in ~5 years); subsequently, the acceptable course was reduced to 1 year—a change that remains especially controversial in the absence of supporting clinical trials.

Several factors have caused many authorities to question the WHO recommendations and to favor a more intensive approach. Among these factors are—for multibacillary patients—a high (double-digit) relapse rate in several locales (reaching 20–40% in one locale, with the rate directly related to the initial bacterial burden) and—for paucibacillary patients—demonstrable lesional activity for years in fully half of patients after the completion of therapy. The more intensive approach (Table 203-2) calls for tuberculoid leprosy to be treated with dapsone (100 mg/d) for 5 years and for lepromatous leprosy to be treated with rifampin (600 mg/d) for 3 years and with dapsone (100 mg/d) throughout life.

With effective antimicrobial therapy, new skin lesions and signs and symptoms of peripheral neuropathy cease appearing. Nodules and plaques of lepromatous leprosy noticeably flatten in 1–2 months and resolve in 1 year or a few years, while tuberculoid skin lesions may disappear, ameliorate, or remain relatively unchanged. Although the peripheral neuropathy of leprosy may improve somewhat in the first few months of therapy, rarely is it significantly alleviated by treatment.

Despite the drawbacks of the WHO's recommendations for multidrug therapy, these regimens have been used almost exclusively worldwide. Although two of the three recommended drugs (dapsone and clofazimine) are only bacteriostatic against *M. leprae* and bactericidal agents have been identified since the WHO formulated its recommendations, significant studies employing the available alternatives in newly designed regimens have not been initiated. Given the recent findings that moxifloxacin, like rifampin, is profoundly bactericidal in leprosy patients and that short-course chemotherapy for tuberculosis is possible only when two or more bactericidal agents are used, a moxifloxacin/rifamycin-based regimen including either minocycline or clarithromycin appears promising; such a regimen may prove to be more reliably curative than WHO-recommended multidrug therapy for lepromatous leprosy and may allow a considerably shorter course of treatment.

THErapy FOR REACTIONS

Type 1 Type 1 lepra reactions are best treated with glucocorticoids (e.g., prednisone, initially at doses of 40–60 mg/d). As inflammation subsides, the glucocorticoid dose can be tapered, but steroid therapy must be continued for at least 3–6 months lest recurrence supervene. Because of the myriad toxicities of prolonged glucocorticoid therapy, the indications for its initiation are strictly limited to lesions whose intense inflammation poses a threat of ulceration; lesions at cosmetically important sites, such as the face; and cases in which neuritis is present. Mild to moderate lepra reactions that do not meet these criteria should be tolerated and glucocorticoid treatment withheld. Thalidomide is ineffective against type 1 lepra

1128 reactions. Clofazimine (200–300 mg/d) is of questionable benefit but in any event is far less efficacious than glucocorticoids.

Type 2 Treatment of ENL must be individualized. If ENL is mild (i.e., if it occurs without fever or other organ involvement and with occasional crops of only a few skin papules), it may be treated with antipyretics alone. However, in cases with many skin lesions, fever, malaise, and other tissue involvement, brief courses (1–2 weeks) of glucocorticoid treatment (initially 40–60 mg/d) are often effective. With or without therapy, individual inflamed papules last for <1 week. Successful therapy is defined by the cessation of skin lesion development and the disappearance of other systemic signs and symptoms. If, despite two courses of glucocorticoid therapy, ENL appears to be recurring and persisting, treatment with thalidomide (100–300 mg nightly) should be initiated, with the dose depending on the initial severity of the reaction. Because even a single dose of thalidomide administered early in pregnancy may result in severe birth defects, including phocomelia, the use of this drug in the United States for the treatment of fertile female patients is tightly regulated and requires informed consent, prior pregnancy testing, and maintenance of birth control measures. Although the mechanism of thalidomide's dramatic action against ENL is not entirely clear, the drug's efficacy is probably attributable to its reduction of TNF levels and IgM synthesis and its slowing of polymorphonuclear leukocyte migration. After the reaction is controlled, lower doses of thalidomide (50–200 mg nightly) are effective in preventing relapses of ENL. Clofazimine in high doses (300 mg nightly) has some efficacy against ENL, but its use permits only a modest reduction of the glucocorticoid dose necessary for ENL control.

Lucio's Phenomenon Neither glucocorticoids nor thalidomide is effective against this syndrome. Optimal wound care and therapy for bacteremia are indicated. Ulcers tend to be chronic and heal poorly. In severe cases, exchange transfusion may prove useful.

PREVENTION AND CONTROL

Vaccination at birth with bacille Calmette-Guérin (BCG) has proved variably effective in preventing leprosy: the results have ranged from total ineffectivity to 80% efficacy. The addition of heat-killed *M. leprae* to BCG does not increase the effectiveness of the vaccine. Because whole mycobacteria contain large amounts of lipids and carbohydrates that have proved in vitro to be immunosuppressive for lymphocytes and macrophages, *M. leprae* proteins may prove to be superior vaccines. Data from a mouse model support this possibility.

Chemoprophylaxis with dapsone may reduce the number of tuberculoid leprosy cases but not the number of lepromatous cases and hence is not recommended, even for household contacts. In addition, single-dose rifampin prophylaxis is of doubtful efficacy. Because leprosy transmission appears to require close prolonged household contact, hospitalized patients need not be isolated.

In 1992, the WHO—on the basis of that organization's treatment recommendations—launched a landmark campaign to eliminate leprosy as a public health problem by the year 2000 (goal, <1 case per 10,000 population). The campaign mobilized and energized nongovernmental organizations and national health services to treat leprosy with multiple drugs and to clean up outdated registries. In these respects, the effort has proved hugely successful, with >6 million patients completing therapy. However, the target of leprosy elimination has not yet been reached. In fact, the success of the WHO campaign in reducing the number of cases worldwide has been largely attributable to the redefinition of what constitutes a case of leprosy. Formerly calculated by disease prevalence, the count is now limited to cases not yet treated with multiple drugs. Worldwide, the annual incidence of leprosy has not fallen. Furthermore, after the completion of therapy, when a patient is no longer considered to represent a "case," half of all patients continue to manifest disease activity for years; relapse rates (at least for multibacillary patients) are unacceptably high; disabilities and deformities go unchecked; and the social stigma of the disease persists.

During most of the twentieth century, nongovernmental organizations, particularly Christian missionaries, provided a medical

infrastructure devoted to the care and treatment of leprosy patients—the envy of those with other medical priorities in the developing world. With the public perception that leprosy is near eradication, resources for patient care are rapidly being diverted, and the burden of patient care is being transferred to nonexistent or overloaded national health services and to health workers who lack the tools and skills needed for the disease's diagnosis and classification and for the selection of nuanced therapy (particularly in cases of reactional neuritis). Thus the prerequisites for a salutary outcome increasingly go unmet.

204 Nontuberculous Mycobacterial Infections

Steven M. Holland

Several terms—nontuberculous mycobacteria (NTM), atypical mycobacteria, mycobacteria other than tuberculosis, and environmental mycobacteria—all refer to mycobacteria other than *Mycobacterium tuberculosis*, its close relatives (*M. bovis*, *M. caprae*, *M. africanum*, *M. pinnipedii*, *M. canetti*), and *M. leprae*. The number of identified species of NTM is growing and will continue to do so because of the use of DNA sequence typing for speciation. The number of known species currently exceeds 150. NTM are highly adaptable and can inhabit hostile environments, including industrial solvents.

EPIDEMIOLOGY

NTM are ubiquitous in soil and water. Specific organisms have recurring niches, such as *M. simiae* in certain aquifers, *M. fortuitum* in pedicure baths, and *M. immunogenum* in metalworking fluids. Most NTM cause disease in humans only rarely unless some aspect of host defense is impaired, as in bronchiectasis, or breached, as by inoculation (e.g., liposuction, trauma). There are few instances of human-to-human transmission of NTM, which occurs almost exclusively in cystic fibrosis. Because infections due to NTM are rarely reported to health agencies and because their identification is sometimes problematic, reliable data on incidence and prevalence are lacking. Disseminated disease denotes significant immune dysfunction (e.g., advanced HIV infection), whereas pulmonary disease, which is much more common, is highly associated with pulmonary epithelial defects but not with systemic immunodeficiency.

In the United States, the incidence and prevalence of pulmonary infection with NTM, mostly in association with bronchiectasis (Chap. 312), have for many years been several-fold higher than the corresponding figures for tuberculosis, and rates of the former are increasing among the elderly. Among patients with cystic fibrosis, who often have bronchiectasis, rates of clinical infection with NTM range from 3% to 15%, with even higher rates among older patients. Although NTM may be recovered from the sputa of many individuals, it is critical to differentiate active disease from commensal harboring of the organisms. A scheme to help with the proper diagnosis of pulmonary infection caused by NTM has been developed by the American Thoracic Society and is widely used. The bulk of nontuberculous mycobacterial disease in North America is due to *M. kansasii*, organisms of the *M. avium* complex (MAC), and *M. abscessus*.



In Europe, Asia, and Australia, the distribution of NTM in clinical specimens is roughly similar to that in North America, with MAC species and rapidly growing organisms such as *M. abscessus* encountered frequently. *M. xenopi* and *M. malmoense* are especially prominent in northern Europe. *M. ulcerans* causes the distinct clinical entity Buruli ulcer, which occurs throughout tropical zones, especially in western Africa. *M. marinum* is a common cause of cutaneous and tendon infections in coastal regions and among individuals exposed to fish tanks or swimming pools.

The true international epidemiology of infections due to NTM is hard to determine because the isolation of these organisms often is not reported and speciation often is not performed for *M. tuberculosis* and NTM. The increasing ease of identification and speciation of these organisms is likely to have a major impact on the description of their international epidemiology in the next few years.

PATHOBIOLOGY

Because exposure to NTM is essentially universal and disease is rare, it can be assumed that normal host defenses against these organisms must be strong and that otherwise healthy individuals in whom significant disease develops are highly likely to have specific susceptibility factors that permit NTM to become established, multiply, and cause disease. At the advent of HIV infection, CD4⁺ T lymphocytes were recognized as key effector cells against NTM; the development of disseminated MAC disease was highly correlated with a decline in CD4⁺ T lymphocyte numbers. Such a decrease has also been implicated in disseminated MAC infection in patients with idiopathic CD4⁺ T lymphocytopenia. Potent inhibitors of tumor necrosis factor α (TNF- α), such as infliximab, adalimumab, certolizumab, golimumab, and etanercept, can neutralize this critical cytokine. The occasional result is severe mycobacterial or fungal infection; these associations indicate that TNF- α is a crucial element in mycobacterial control. However, in cases without the above risk factors, much of the genetic basis of susceptibility to disseminated infection with NTM is accounted for by specific mutations in the interferon γ (IFN- γ)/interleukin 12 (IL-12) synthesis and response pathways.

Mycobacteria are typically phagocytosed by macrophages, which respond with the production of IL-12, a heterodimer composed of IL-12p35 and IL-12p40 moieties that together make up IL-12p70. IL-12 activates T lymphocytes and natural killer cells through binding to its receptor (composed of IL-12R β 1 and IL-12R β 2/IL-23R), with consequent phosphorylation of STAT4. IL-12 stimulation of STAT4 leads to secretion of IFN- γ , which activates neutrophils and macrophages to produce reactive oxidants, to increase expression of the major histocompatibility complex and Fc receptors, and to concentrate certain antibiotics intracellularly. Signaling by IFN- γ through its receptor (composed of IFN- γ R1 and IFN- γ R2) leads to phosphorylation of STAT1, which in turn regulates IFN- γ -responsive genes, such as those coding for IL-12 and TNF- α . TNF- α signals through its own receptor via a downstream complex containing the nuclear factor κ B (NF- κ B) essential modulator (NEMO). Therefore, the positive feedback loop between IFN- γ and IL-12/IL-23 drives the immune response to mycobacteria and other intracellular infections. These genes are known to be the critical ones in the pathway of mycobacterial control: specific Mendelian mutations have been identified in *IFN- γ R1*, *IFN- γ R2*, *STAT1*, *GATA2*, *ISG15*, *IRF8*, *IL-12A*, *IL-12R β 1*, *IL-12R β 2*, *CYBB*, and *NEMO* (Fig. 204-1). Despite the identification of genes associated with disseminated disease, only ~70% of cases of disseminated nontuberculous mycobacterial infections that are not associated with HIV infection have a genetic diagnosis; the implication is that more mycobacterial susceptibility genes and pathways remain to be identified.

In contrast to the recognized genes and mechanisms associated with disseminated nontuberculous mycobacterial infection, the best-recognized underlying condition for pulmonary infection with NTM is bronchiectasis (Chap. 312). Most of the well-characterized forms of bronchiectasis, including cystic fibrosis, primary ciliary dyskinesia, STAT3-deficient hyper-IgE syndrome, and idiopathic bronchiectasis, have high rates of association with nontuberculous mycobacterial infection. The precise mechanism by which bronchiectasis predisposes to locally destructive but not systemic involvement is unknown.

Unlike disseminated or pulmonary infection, “hot-tub lung” represents pulmonary hypersensitivity to NTM—most commonly MAC organisms—growing in underchlorinated, often indoor hot tubs.

CLINICAL MANIFESTATIONS

Disseminated Disease Disseminated MAC or *M. kansasii* infections in patients with advanced HIV infection are now uncommon in North America because of effective antimycobacterial prophylaxis

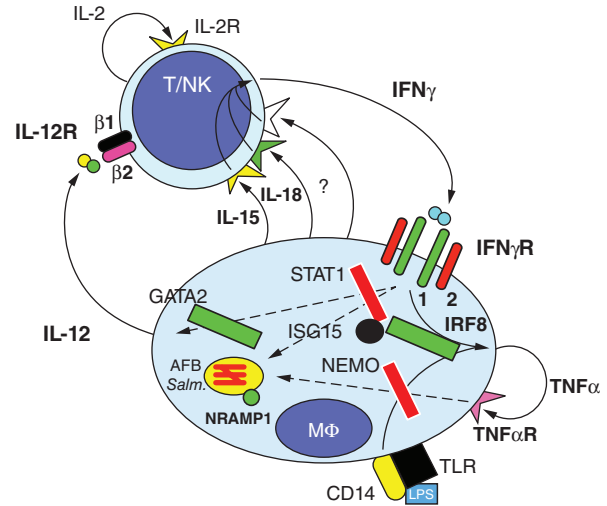


FIGURE 204-1 Cytokine interactions of infected macrophages (M ϕ) with T and natural killer (NK) lymphocytes.

Infection of macrophages by mycobacteria (AFB) leads to the release of heterodimeric interleukin 12 (IL-12). IL-12 acts on its receptor complex (IL-12R), with consequent STAT4 activation and production of homodimeric interferon γ (IFN γ). Through its receptor (IFN γ R), IFN γ activates STAT1, stimulating the production of tumor necrosis factor α (TNF α) and leading to the killing of intracellular organisms such as mycobacteria, salmonellae (Salm), and some fungi. Homotrimeric TNF α acts through its receptor (TNF α R) and requires nuclear factor κ B essential modulator (NEMO) to activate nuclear factor κ B, which also contributes to the killing of intracellular bacteria. Both IFN γ and TNF α lead to upregulation of IL-12. TNF α -blocking antibodies work either by blocking the ligand (infliximab, adalimumab, certolizumab, golimumab) or by providing soluble receptor (etanercept). Mutations in *IFN γ R1*, *IFN γ R2*, *IL-12p40*, *IL-12R β 1*, *IL-12R β 2*, *STAT1*, *GATA2*, *ISG15*, *IRF8*, *CYBB*, and *NEMO* have been associated with a predisposition to mycobacterial infections. Other cytokines, such as IL-15 and IL-18, also contribute to IFN γ production. Signaling through the Toll-like receptor (TLR) complex and CD14 also upregulates TNF α production. LPS, lipopolysaccharide; NRAMP1, natural resistance-associated macrophage protein 1.

and improved treatment of HIV infection. When such mycobacterial disease was common, the portal of entry was the bowel, with spread to bone marrow and the bloodstream. Surprisingly, disseminated infections with rapidly growing NTM (e.g., *M. abscessus*, *M. fortuitum*) are very rare in HIV-infected patients, even those with very advanced HIV infection. Because these organisms are of low intrinsic virulence and disseminate only in conjunction with impaired immunity, disseminated disease can be indolent and progressive over weeks to months. Typical manifestations of malaise, fever, and weight loss are often accompanied by organomegaly, lymphadenopathy, and anemia. Because special cultures or stains are required to identify the organisms, the most critical step in diagnosis is to suspect infection with NTM. Blood cultures may be negative, but involved organs typically have significant organism burdens, sometimes with a grossly impaired granulomatous response. In a child, disseminated involvement (i.e., involvement of two or more organs) without an underlying iatrogenic cause should prompt an investigation of the IFN- γ /IL-12 pathway. Recessive mutations in *IFN- γ R1* and *IFN- γ R2* typically lead to severe infection with NTM. In contrast, dominant negative mutations in *IFN- γ R1*, which lead to overaccumulation of a defective interfering mutant receptor on the cell surface, inhibit normal IFN- γ signaling and thus lead to nontuberculous mycobacterial osteomyelitis. Dominant negative mutations in *STAT1* and recessive mutations in *IL-12R β 1* can produce variable phenotypes consistent with their residual capacities for IFN- γ synthesis and response. Male patients who have disseminated nontuberculous mycobacterial infections along with conical, peg, or missing teeth and an abnormal hair pattern should be evaluated for defects in the pathway

1130 that activates NF- κ B through NEMO. These patients may have associated immune globulin defects as well. Patients with myelodysplasia and mycobacterial disease should be investigated for GATA2 deficiency. A recently recognized group of patients that often develops disseminated infections with rapidly growing NTM (predominantly *M. abscessus*) as well as other opportunistic infections has high-titer neutralizing autoantibodies to IFN- γ . Thus far, this syndrome has been reported most frequently in East Asian female patients.

IV catheters can become infected with NTM, usually as a consequence of contaminated water. *M. abscessus* and *M. fortuitum* sometimes infect deep indwelling lines as well as fluids used in eye surgery, subcutaneous injections, and local anesthetics. Infected catheters should be removed.

Pulmonary Disease Lung disease is by far the most common form of nontuberculous mycobacterial infection in North America and the rest of the industrialized world. The clinical presentation typically consists of months or years of throat clearing, nagging cough, and slowly progressive fatigue. Patients will often have seen physicians multiple times and received symptom-based or transient therapy before the diagnosis is entertained and samples are sent for mycobacterial stains and cultures. Because not all patients can produce sputum, bronchoscopy may be required for diagnosis. The typical lag between onset of symptoms and diagnosis is ~5 years in older women. Predisposing factors include underlying lung diseases such as bronchiectasis (Chap. 312), pneumoconiosis (Chap. 311), chronic obstructive pulmonary disease (Chap. 314), primary ciliary dyskinesia (Chap. 312), α_1 antitrypsin deficiency (Chap. 367e), and cystic fibrosis (Chap. 313). Bronchiectasis and nontuberculous mycobacterial infection often coexist and progress in tandem. This situation makes causality difficult to determine in a given index case, but bronchiectasis is certainly among the most critical predisposing factors that are exacerbated by infection.

MAC organisms are the most common causes of pulmonary nontuberculous mycobacterial infection in North America, but rates vary somewhat by region. MAC infection most commonly develops during the sixth or seventh decade of life in women who have had months or years of nagging intermittent cough and fatigue, with or without sputum production or chest pain. The constellation of pulmonary disease due to NTM in a tall and thin woman who may have chest wall abnormalities is often referred to as Lady Windermere syndrome, after an Oscar Wilde character of the same name. In fact, pulmonary MAC infection does afflict older nonsmoking white women more than men, with onset at ~60 years. Patients tend to be taller and thinner than the general population, with high rates of scoliosis, mitral valve prolapse, and pectus anomalies. Whereas male smokers with upper-lobe cavitory disease tend to carry the same single strain of MAC indefinitely, nonsmoking females with nodular bronchiectasis tend to carry several strains of MAC simultaneously, with changes over the course of their disease.

M. kansasii can cause a clinical syndrome that strongly resembles tuberculosis, consisting of hemoptysis, chest pain, and cavitory lung disease. The rapidly growing NTM, such as *M. abscessus*, have been associated with esophageal motility disorders such as achalasia. Patients with pulmonary alveolar proteinosis are prone to pulmonary nontuberculous mycobacterial and *Nocardia* infections; the underlying mechanism may be inhibition of alveolar macrophage function due to the autoantibodies to granulocyte-macrophage colony-stimulating factor found in these patients.

Cervical Lymph Nodes The most common form of nontuberculous mycobacterial infection among young children in North America is isolated cervical lymphadenopathy, caused most frequently by MAC organisms but also by other NTM. The cervical swelling is typically firm and relatively painless, with a paucity of systemic signs. Because the differential diagnosis of painless adenopathy includes malignancy, many children have infection with NTM diagnosed inadvertently at biopsy; cultures and special stains may not have been requested because mycobacterial disease was not ranked high in the differential. Local fistulae usually resolve completely with resection and/or antibiotic therapy. Likewise, the entity of isolated pediatric intrathoracic nontuberculous mycobacterial infection, which is

probably related to cervical lymph node infection, is usually mistaken for cancer. In neither isolated cervical nor isolated intrathoracic infections with NTM have children with underlying immune defects been identified, nor do the affected children go on to develop other opportunistic infections.

Skin and Soft Tissue Disease Cutaneous involvement with NTM usually requires a break in the skin for introduction of the bacteria. Pedicure bath-associated infection with *M. fortuitum* is more likely if skin abrasion (e.g., during leg shaving) has occurred just before the pedicure. Outbreaks of skin infection are often caused by rapidly growing NTM (especially *M. abscessus*, *M. fortuitum*, and *M. chelonae*) acquired via skin contamination from surgical instruments (especially in cosmetic surgery), injections, and other procedures. These infections are typically accompanied by painful, erythematous, draining subcutaneous nodules, usually without associated fever or systemic symptoms.

M. marinum lives in many water sources and can be acquired from fish tanks, swimming pools, barnacles, and fish scales. This organism typically causes papules or ulcers (“fish-tank granuloma”), but the infection can progress to tendinitis with significant impairment of manual dexterity. Lesions appear days to weeks after inoculation of organisms by a typically minor trauma (e.g., incurred during the cleaning of boats or the handling of fish). Tender nodules due to *M. marinum* can advance up the arm in a pattern also seen with *Sporothrix schenckii* (*sporotricoid spread*). The typical carpal tendon involvement may be the first presenting manifestation and may lead to surgical exploration or steroid injection. The index of suspicion for *M. marinum* infections must be high to ensure that proper specimens obtained during procedures are sent for culture.

M. ulcerans, another waterborne skin pathogen, is found mainly in the tropics, especially in tropical areas of Africa. Infection follows skin trauma or insect bites that allow admission to contaminated water. The skin lesions are typically painless, clean ulcers that slough and can cause osteomyelitis. The toxin mycolactone accounts for the modest host inflammatory response and the painless ulcerations.

DIAGNOSIS

NTM can be detected on acid-fast or fluorochrome smears of sputum or other body fluids. When the organism burden is high, the organisms may appear as gram-positive beaded rods, but this finding is unreliable. (In contrast, nocardiae may appear as gram-positive and beaded but filamentous bacteria.) Again, the requisite and most sensitive step in the diagnosis of any mycobacterial disease is to think of including it in the differential. In almost all laboratories, mycobacterial sample processing, staining, and culture are conducted separately from routine bacteriologic tests; thus many infections go undiagnosed because of the physician’s failure to request the appropriate test. In addition, mycobacteria usually require separate blood culture media. NTM are broadly differentiated into rapidly growing (<7 days) and slowly growing (≥ 7 days) forms. Because *M. tuberculosis* typically takes ≥ 2 weeks to grow, many laboratories refuse to consider culture results final until 6 weeks have elapsed. Newer techniques using liquid culture media permit more rapid isolation of mycobacteria from specimens than is possible with traditional media. Species more readily detected with incubation at 30°C include *M. marinum*, *M. haemophilum*, and *M. ulcerans*. *M. haemophilum* prefers iron supplementation or blood, whereas *M. genavense* requires supplemented medium with the additive mycobactin J. Bacterial formation of pigment in light conditions (*photochromogenicity*) or dark conditions (*scotochromogenicity*) or a lack of bacterial pigment formation (*nonchromogenicity*) has been used to help categorize NTM. In contrast to NTM colonies, *M. tuberculosis* colonies are beige, rough, dry, and flat. Current identification schemes can reliably use biochemical, nucleic acid, or cell wall composition, as assessed by high-performance liquid chromatography or mass spectrometry, for speciation. With the remarkable decline in U.S. cases of tuberculosis over recent decades, NTM have become the mycobacteria most commonly isolated from humans in North America. However, not all isolations of NTM, especially from the lung, reflect pathology and require treatment. Whereas identification of an organism in a blood or organ biopsy specimen in a compatible clinical setting is diagnostic, the

American Thoracic Society recommends that pulmonary infection due to NTM be diagnosed only when disease is clearly demonstrable—i.e., in an appropriate clinical and radiographic setting (nodules, bronchiectasis, cavities) and with repeated isolation of NTM from expectorated sputum or recovery of NTM from bronchoscopy or biopsy specimens. Given the large number of species of NTM and the importance of accurate diagnosis for the implementation of proper therapy, identification of these organisms is ideally taken to the species level.

The purified protein derivative (PPD) of tuberculin is delivered intradermally to evoke a memory T cell response to mycobacterial antigens. This test is variously referred to as the PPD test, the tuberculin skin test, and the Mantoux test, among other designations. Unfortunately, the cutaneous immune response to these tuberculosis-derived filtrate proteins does not differentiate well between infection with NTM and that with *M. tuberculosis*. Because intermediate reactions (~10 mm) to PPD in latent tuberculosis and nontuberculous mycobacterial infections can overlap significantly, the progressive decline in active tuberculosis in the United States means that NTM probably account for increasing proportions of PPD reactivity. In addition, bacille Calmette-Guérin (BCG) can cause some degree of cross-reactivity, posing problems of interpretation for patients who have received BCG vaccine. Assays to measure the elaboration of IFN- γ in response to the relatively tuberculosis-specific proteins ESAT6 and CFP10 form the basis for IFN- γ -release assays (IGRAs). These assays can be performed with whole blood or on membranes. It is important to note that *M. marinum*, *M. kansasii*, and *M. szulgai* also have ESAT6 and CFP10 and may cause false-positive reactions in IGRAs. Despite cross-reactivity with NTM, large PPD reactions (>15 mm) most commonly signify tuberculosis.

Isolation of NTM from blood specimens is clear evidence of disease. Whereas rapidly growing mycobacteria may proliferate in routine blood culture media, slow-growing NTM typically do not; thus it is imperative to suspect the diagnosis and to use the correct bottles for cultures. Isolation of NTM from a biopsy specimen constitutes strong evidence for infection, but cases of laboratory contamination do occur. Identification of organisms on stained sections of biopsy material confirms the authenticity of the culture. Certain NTM require lower incubation temperatures (*M. genavense*) or special additives (*M. haemophilum*) for growth. Some NTM (e.g., *M. tuberculosis*) remain noncultivable but can be identified molecularly in clinical samples.

The radiographic appearance of nontuberculous mycobacterial disease in the lung depends on the underlying disease, the severity of the infection, and the imaging modality used. The advent and increase in the use of computed tomography (CT) has allowed the identification of characteristic changes that are highly consistent with nontuberculous mycobacterial infection, such as the “tree-in-bud” pattern of bronchiolar inflammation (Fig. 204-2). Involvement of the lingual and right-middle lobes is commonly seen on chest CT but is difficult to appreciate on plain film. Severe bronchiectasis and cavity formation are common in more advanced disease. Isolation of NTM from respiratory samples can be confusing. *M. gordonae* is often recovered from respiratory samples but is not usually seen on smear and is almost never a pathogen. Patients with bronchiectasis occasionally have NTM recovered from sputum culture with a negative smear. The American Thoracic Society has developed guidelines for the diagnosis of infection with MAC, *M. abscessus*, and *M. kansasii*. A positive diagnosis requires the growth of NTM from two of three sputum samples, regardless of smear findings; a positive bronchoscopic alveolar sample, regardless of smear findings; or a pulmonary parenchyma biopsy sample with granulomatous inflammation or mycobacteria found on section and NTM found on culture. These guidelines probably apply to other NTM as well.

Although many laboratories use DNA probes to identify *M. tuberculosis*, MAC, *M. gordonae*, and *M. kansasii*, speciation of NTM helps determine the antimycobacterial therapy to be used. Only testing of MAC organisms for susceptibility to clarithromycin and of *M. kansasii* for susceptibility to rifampin is indicated; few data support other *in vitro* susceptibility tests, attractive though they appear. MAC isolates that have not been exposed to macrolides are almost always

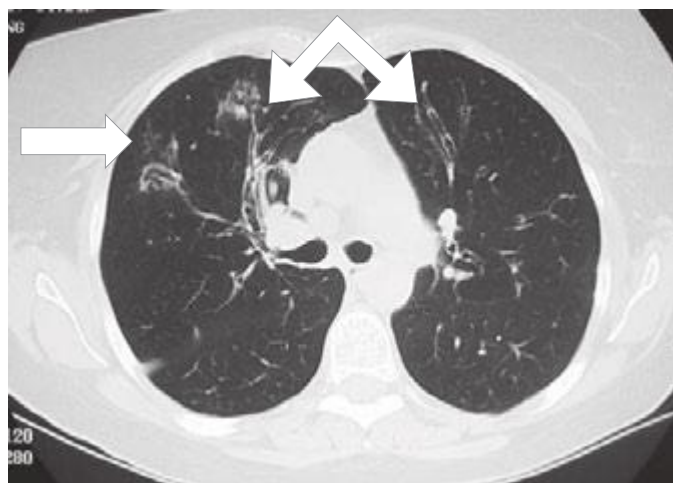


FIGURE 204-2 Chest computed tomography of a patient with pulmonary *Mycobacterium avium* complex infection. Arrows indicate the “tree-in-bud” pattern of bronchiolar inflammation (peripheral right lung) and bronchiectasis (central right and left lungs).

susceptible. NTM that have persisted beyond a course of antimicrobial therapy are often tested for antibiotic susceptibility, but the value and meaning of these tests are undetermined.

PREVENTION

Prophylaxis of MAC disease in patients infected with HIV is started when the CD4⁺ T lymphocyte count falls to <50/ μ L. Azithromycin (1200 mg weekly), clarithromycin (1000 mg daily), or rifabutin (300 mg daily) is effective. Macrolide prophylaxis in immunodeficient patients who are susceptible to NTM (e.g., those with defects in the IFN- γ /IL-12 axis) has not been prospectively validated but seems prudent.

TREATMENT NONTUBERCULOUS MYCOBACTERIA

NTM cause chronic infections that evolve relatively slowly over a period of weeks to years. Therefore, it is rarely necessary to initiate treatment on an emergent basis before the diagnosis is clear and the infecting species is known. Treatment of NTM is complex, often poorly tolerated, and potentially toxic. Just as in tuberculosis, inadequate single-drug therapy is almost always associated with the emergence of antimicrobial resistance and relapse.

MAC infection often requires multidrug therapy, the foundation of which is a macrolide (clarithromycin or azithromycin), ethambutol, and a rifamycin (rifampin or rifabutin). For disseminated nontuberculous mycobacterial disease in HIV-infected patients, the use of rifamycins poses special problems—i.e., rifamycin interactions with protease inhibitors. For pulmonary MAC disease, thrice-weekly administration of a macrolide, a rifamycin, and ethambutol has been successful. Therapy is prolonged, generally continuing for 12 months after culture conversion; typically, a course lasts for at least 18 months. Other drugs with activity against MAC organisms include IV and aerosolized aminoglycosides, fluoroquinolones, and clofazimine. In elderly patients, rifabutin can exert significant toxicity. However, with only modest efforts, most antimycobacterial regimens are well tolerated by most patients. Resection of cavitory lesions or severely bronchiectatic segments has been advocated for some patients, especially those with macrolide-resistant infections. The success of therapy for pulmonary MAC infections depends on whether disease is nodular or cavitory and on whether it is early or advanced, ranging from 20% to 80%.

M. kansasii lung disease is similar to tuberculosis in many ways and is also effectively treated with isoniazid (300 mg/d), rifampin (600 mg/d), and ethambutol (15 mg/kg per day). Other drugs with very high-level activity against *M. kansasii* include clarithromycin, fluoroquinolones, and aminoglycosides. Treatment should continue until

1132 cultures have been negative for at least 1 year. In most instances, *M. kansasii* infection is easily cured.

Rapidly growing mycobacteria pose special therapeutic problems. Extrapulmonary disease in an immunocompetent host is usually due to inoculation (e.g., via surgery, injections, or trauma) or to line infection and is often treated successfully with a macrolide and another drug (with the choice based on in vitro susceptibility), along with removal of the offending focus. In contrast, pulmonary disease, especially that caused by *M. abscessus*, is extremely difficult to cure. Repeated courses of treatment are usually effective in reducing the infectious burden and symptoms. Therapy generally includes a macrolide along with an IV-administered agent such as amikacin, a carbapenem, ceftazidime, or tigecycline. Other oral agents (used according to in vitro susceptibility testing and tolerance) include fluoroquinolones, doxycycline, and linezolid. Because nontuberculous mycobacterial infections are chronic, care must be taken in the long-term use of drugs with neurotoxicities, such as linezolid and ethambutol. Prophylactic pyridoxine has been suggested in these cases. Durations of therapy for *M. abscessus* lung disease are difficult to predict because so many cases are chronic and require intermittent therapy. Expert consultation and management are strongly recommended.

Once recognized, *M. marinum* infection is highly responsive to antimicrobial therapy and is cured relatively easily with any combination of a macrolide, ethambutol, and a rifamycin. Therapy should be continued for 1–2 months after clinical resolution of isolated soft tissue disease; tendon and bone involvement may require longer courses in light of clinical evolution. Other drugs with activity against *M. marinum* include sulfonamides, trimethoprim-sulfamethoxazole, doxycycline, and minocycline.

Treatment of the other NTM is less well defined, but macrolides and aminoglycosides are usually effective, with other agents added as indicated. Expert consultation is strongly encouraged for difficult or unusual infections due to NTM.

PROGNOSIS

The outcomes of nontuberculous mycobacterial infections are closely tied to the underlying condition (e.g., IFN- γ /IL-12 pathway defect, cystic fibrosis) and can range from recovery to death. With no or inadequate treatment, symptoms and signs can be debilitating, including persistent cough, fever, anorexia, and severe lung destruction. With treatment, patients typically regain strength and energy. The optimal

duration of therapy when NTM persist in sputum is unknown, but treatment in this situation can be prolonged.

GLOBAL CONSIDERATIONS



In many countries, pulmonary tuberculosis is diagnosed by smear alone, which is also the method used for monitoring of response and relapse. However, examination of mycobacteria from the affected patients shows that a significant proportion of isolates are actually NTM. Overall, as rates of tuberculosis decline, the proportion of positive smears caused by NTM will increase. Advances in speciation will distinguish tuberculosis from nontuberculous mycobacterial infections and thereby affect rates of assumed relapse and resistance, leading to more targeted and appropriate therapy.



Agents used for the treatment of mycobacterial infections, including tuberculosis (TB), leprosy, and infections due to nontuberculous mycobacteria (NTM), are administered in multiple-drug regimens for prolonged courses. Currently, more than 160 species of mycobacteria have been identified, the majority of which do not cause disease in humans. While the incidence of disease caused by *Mycobacterium tuberculosis* has been declining in the United States, TB remains a leading cause of morbidity and mortality in developing countries—particularly in sub-Saharan Africa, where the HIV epidemic rages. Effective drug regimens are not all that is needed; without a well-organized infrastructure for diagnosis and treatment of TB, therapeutic and control efforts are severely hampered (Chaps. 2 and 13e). Infections with NTM have gained in clinical prominence in the United States and other developed countries. These largely environmental organisms often establish infection in immunocompromised patients or in persons with structural lung disease.

TUBERCULOSIS

GENERAL PRINCIPLES

The earliest recorded human case of TB dates back 9000 years. Early treatment modalities, such as bloodletting, were replaced by sanatorium regimens in the late nineteenth century. The discovery of streptomycin in 1943 launched the era of antibiotic treatment for TB. Over subsequent decades, the discovery of additional agents and the use of multiple-drug regimens allowed progressive shortening of the treatment course from years to as little as 6 months with the regimen for drug-susceptible TB. Latent TB infection (LTBI) and active TB disease are diagnosed by history, physical examination, radiographic imaging, tuberculin skin test, interferon γ release assays, acid-fast staining, mycobacterial cultures, and/or new molecular diagnostics. LTBI is treated with isoniazid (optimally given daily or twice weekly for 9 months), rifampin (daily for 4 months), or isoniazid plus rifapentine (weekly for 3 months) (Table 205e-1).

For active or suspected TB disease, clinical factors, including HIV co-infection, symptom duration, radiographic appearance, and public health concerns about TB transmission, drive diagnostic testing and treatment initiation. Multiple-drug regimens are used for the treatment of TB disease (Table 205e-2). Initially, an intensive phase consisting of four drugs—isoniazid, rifampin, pyrazinamide, and ethambutol—given for 2 months is followed by a continuation phase of isoniazid and rifampin for 4 months, for a total treatment duration of 6 months. The continuation phase is extended to 7 months (for a total treatment duration of 9 months) if the 2-month course of pyrazinamide is not completed or, for patients with cavitary pulmonary TB, if sputum cultures remain positive beyond 2 months of treatment (delayed culture conversion).

Treatment of TB in individuals co-infected with HIV poses significant challenges, but some progress is being made. Recent data show improved survival when antiretroviral therapy (ART) is initiated early during TB treatment. Interactions of rifampin with protease inhibitors

or non-nucleoside reverse transcriptase inhibitors are significant and require close monitoring and dose adjustments. The TB immune reconstitution inflammatory syndrome (IRIS) may appear as early as 1 week after initiation of ART and manifests as paradoxical worsening or unmasking of existing TB infection. Conservative management consists of continued administration of ART and TB medications. However, severe or debilitating IRIS has been anecdotally treated with varying doses of glucocorticoids. Intermittent therapy in patients co-infected with HIV and *M. tuberculosis* has been associated with low plasma levels of several key TB drugs and with higher rates of treatment failure or relapse; therefore, intermittent twice-weekly therapy for TB in HIV-co-infected individuals is not recommended.

Adherence to medications is critical in achieving a cure with antimycobacterial therapy. Consequently, directly observed therapy (DOT) by trained staff, either in the clinic or at home, is recommended to ensure adherence. In addition, monthly dispensing of TB medicines is recommended because monthly clinical monitoring for hepatotoxicity due to these medications is essential for all patients. Discontinuation of suspected offending agents at the onset of hepatitis symptoms reduces the risk of progression to fatal hepatitis. Clinical monitoring includes at least monthly assessment for symptoms (nausea, vomiting, abdominal discomfort, and unexplained fatigue) and signs (jaundice, dark urine, light stools, diffuse pruritus) of hepatotoxicity, although the latter represent comparatively late manifestations (Table 205e-3). The presence of such symptoms and signs mandates provisional discontinuation of potentially hepatotoxic agents. Biochemical testing of at least serum alanine aminotransferase and total bilirubin levels and exclusion of other causes of these abnormalities are also indicated during treatment for those at risk for hepatotoxicity (Table 205e-3). For patients with active TB, monthly mycobacterial cultures of sputum are recommended until it is certain that the organisms have been cleared and the patient has responded to therapy or until no sputum is available for culture.

If significant clinical improvement does not occur or the patient's condition deteriorates over the course of therapy, possibilities include treatment failure due to nonadherence, poor medication absorption, or the development of resistance. For patients co-infected with HIV and *M. tuberculosis*, IRIS, which is a diagnosis of exclusion, should also be a consideration. Drug susceptibility testing should be repeated at this point. If resistance is documented or strongly suspected, at least two efficacious drugs to which the isolate is susceptible or which the patient has not already taken should be added to the therapeutic regimen.

Multidrug-resistant TB (MDR-TB) is defined as disease caused by a strain of *M. tuberculosis* that is resistant to both isoniazid and rifampin—the most efficacious of the first-line TB drugs. The risk of MDR-TB is elevated in patients presenting from geographic areas in which $\geq 5\%$ of incident cases are MDR-TB and in patients previously treated for TB. Treatment regimens for MDR-TB generally include a late-generation fluoroquinolone and an injectable second-line agent (such as capreomycin, amikacin, or kanamycin). Regimens of at least five drugs are recommended for the treatment of MDR-TB. Both standardized and optimized/customized regimens are in use around the world. Extensively drug-resistant TB (XDR-TB) is defined as MDR-TB with additional resistance to any fluoroquinolone and at least one of the second-line injectable agents. Treatment of XDR-TB is individualized

TABLE 205e-1 REGIMENS FOR THE TREATMENT OF LATENT TUBERCULOSIS INFECTION IN ADULTS

Regimen	Schedule	Duration	Comments
Isoniazid	300 mg/d (5 mg/kg) Alternative: 900 mg twice weekly (15 mg/kg)	9 months (6 months acceptable)	Supplement with pyridoxine (25–50 mg daily). Twice-weekly regimens require directly observed therapy.
Rifampin	600 mg/d (10 mg/kg)	4 months	Broader efficacy studies are needed.
Isoniazid plus rifapentine	900 mg (15 mg/kg) weekly + 900 mg weekly	3 months	Directly observed therapy is recommended for once-weekly treatment. This regimen may be supplemented with pyridoxine (25–50 mg/d).

Sources: D Menzies et al: Ann Intern Med 149:689, 2008; American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America: Am J Respir Crit Care Med 167:603, 2003; T Sterling et al: N Engl J Med 365:2155, 2011.

TABLE 205e-2 SIMPLIFIED APPROACH TO TREATMENT OF ACTIVE TUBERCULOSIS (TB) IN ADULTS

Culture Results	Intensive Phase	Continuation Phase	Extension of Total Treatment
Culture positive	HRZE for 2 months, daily or intermittent (with dose adjustment)	HR for 4 months, daily or 5 d/wk or HR for 4 months, intermittent (with dose adjustment)	To 9 months, if 2 months of Z is not completed or culture conversion is prolonged and cavitation is evident on plain radiograph ^a
Culture negative	HRZE for 2 months	2 months	To 6 months, if patient is infected with HIV
Extrapulmonary	HRZE for 2 months	HR for 4–7 months, daily or 5 d/wk ^b	To 9–12 months in TB meningitis. Some recommend 9 months for bone/joint TB.
Resistant to H	QRZE ^c or, less often, RZES for 6 months	...	Prolonged culture conversion, cavitation
Resistant to R	HZEQ ^c (IA ^d) for 2 months	HEQ(S) for 10–16 months	Prolonged culture conversion, delayed response
Resistant to HR ^e	ZEQ ^c (IA ^d) ± alternative agents ^f for 18–24 months	...	Prolonged culture conversion

^aCulture conversion is prolonged if it occurs beyond 2 months. Some providers extend the continuation phase to 7 months if there is *either* prolonged culture conversion or cavitation. ^bMany experts recommend a continuation phase of 7 months for all extrapulmonary TB, including miliary disease. For TB pericarditis and meningitis, the addition of glucocorticoids is recommended. ^cLevofloxacin and moxifloxacin are the preferred fluoroquinolones. Gatifloxacin is associated with dysglycemia but may be an acceptable alternative; in a recent trial of TB treatment, this drug did not cause dysglycemia in patients who received it thrice weekly for 4 months. Ofloxacin and ciprofloxacin should generally be avoided because of resistance. ^dInjectable agents: streptomycin, amikacin, kanamycin, and capreomycin. ^eMultidrug-resistant TB should be managed by or in close consultation with an expert TB clinician. Surgical management should be considered. ^fAlternative agents: cycloserine, ethionamide, para-aminosalicylic acid, clarithromycin, linezolid, and amoxicillin-clavulanate.

Abbreviations: E, ethambutol; H, isoniazid; IA, injectable agent; Q, fluoroquinolone; R, rifampin; S, streptomycin; Z, pyrazinamide.

Sources: American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America: *Am J Respir Crit Care Med* 167:603, 2003; C Mitnick et al: *N Engl J Med* 359:563, 2008; World Health Organization 2011 update: Guidelines for the programmatic management of drug-resistant tuberculosis (www.who.int/tb/challenges/mdr/programmatic_guidelines_for_mdr/tb/en/index.html).

on the basis of complete phenotypic and, if possible, genotypic antimicrobial susceptibility testing. Therapeutic regimens for either MDR-TB or XDR-TB should be constructed with input from experienced clinicians who should continue the management of the disease.

FIRST-LINE ANTITUBERCULOSIS DRUGS

The following discussion of individual anti-TB agents focuses on treatment of TB in adults, unless otherwise noted. Several agents are being

actively investigated during the current remarkable period of drug development for TB treatment.

Isoniazid Isoniazid is a critical drug for treatment of both TB disease and LTBI. Isoniazid has excellent bactericidal activity against both intracellular *M. tuberculosis* and extracellular, actively dividing organisms. This drug is bacteriostatic against slowly dividing organisms. In treatment of LTBI, isoniazid is considered the first-line agent because it is generally well tolerated, has well-established efficacy, and

TABLE 205e-3 MONITORING AND CLINICAL MANAGEMENT OF TUBERCULOSIS TREATMENT IN ADULTS^a

Drug	Assessment	Management
LTBI Treatment		
With hepatic risk factors ^b , check ALT and bilirubin at baseline. If ALT is $\geq 3 \times$ ULN or total bilirubin is $> 2 \times$ ULN, defer treatment and reevaluate.		
Isoniazid	Determine whether hepatic risk factors are present. If so, obtain baseline and periodic ALT and bilirubin values.	If ALT is $5 \times$ ULN (or $3 \times$ ULN with symptoms) ^c or if bilirubin reaches jaundice levels (usually $> 2 \times$ ULN), interrupt treatment. With normalization, consider an alternative agent.
Rifampin	Same as above	Same as above
TB Treatment		
Check ALT, bilirubin, platelets, creatinine, and hepatitis panel for all patients at baseline. If hepatic risk factors are present, check ALT and bilirubin monthly.		
Isoniazid	If ALT is $> 5 \times$ ULN (or $> 3 \times$ ULN with hepatitis symptoms) ^c	Obtain history of alcohol consumption and concomitant drug use. In most instances, discontinue isoniazid, pyrazinamide, rifampin, and other hepatotoxic drugs. Consider alternative agents. Obtain viral hepatitis serologies. Rechallenge: With normalization of liver enzymes, rifampin and isoniazid may be sequentially reintroduced. With no recurrence of hepatotoxicity, pyrazinamide is not resumed in many cases. Alternative rechallenge protocols have been used.
Rifampin	If primary elevation is in bilirubin and alkaline phosphatase, most likely due to rifampin	Discontinue rifampin if total bilirubin reaches jaundice levels (usually $> 2 \times$ ULN). May try to reintroduce; if not tolerated, may substitute fluoroquinolone
Ethambutol	Decrease in visual acuity or color vision or appearance on monthly screening	Discontinue ethambutol and repeat ocular exam. Peripheral neuropathy may be a precursor of ocular toxicity; if it occurs, consider repeat ocular exam.
Pyrazinamide	If ALT is $> 5 \times$ ULN (or $> 3 \times$ ULN with symptoms) ^c	Same as for isoniazid
Fluoroquinolone	If QTc prolongation is discovered incidentally on ECG	Check audiometry and at least BUN and creatinine monthly.
Aminoglycoside	Abnormal results on audiometry testing, BUN, creatinine, electrolytes at baseline or on monthly check	Discontinue aminoglycoside if not MDR-TB. As appropriate, assess renal function, correct electrolytes, or seek ENT consultation.

^aAll regimens require monthly clinical monitoring. ^bHepatic risk factors: chronic alcohol use, viral hepatitis, preexisting liver disease, pregnancy or ≤ 3 months postpartum, hepatotoxic medications. ^cRelevant manifestations include nausea, vomiting, abdominal pain, jaundice, or unexplained fatigue.

Abbreviations: ALT, alanine aminotransferase; BUN, blood urea nitrogen; ECG, electrocardiogram; ENT, ear, nose, and throat; LTBI, latent tuberculosis infection; MDR-TB, multidrug-resistant tuberculosis; ULN, upper limit of normal.

Sources: JJ Saukkonen et al: *Am J Respir Crit Care Med* 174:935, 2006; American Thoracic Society/Centers for Disease Control and Prevention/Infectious Diseases Society of America: *Am J Respir Crit Care Med* 167:603, 2003.

is inexpensive. In this setting, the drug is taken daily or intermittently (i.e., twice weekly) as DOT for 9 months. The 9-month course is more efficacious than the 6-month course (75–90% vs ≤65%), but extension of treatment to 12 months is not likely to provide further protection. A 6-month course of daily or intermittent isoniazid is considered second-line, but acceptable, therapy. A recent large open-label, multicenter, randomized, controlled trial showed that weekly DOT with isoniazid and rifampin, administered over 3 months, was not inferior to daily isoniazid given for 9 months and had a higher treatment completion rate than the single-drug regimen.

For treatment of TB disease, isoniazid is used in combination with other agents to ensure killing of both actively dividing *M. tuberculosis* and slowly growing "persister" organisms. Unless the organism is resistant, the standard regimen includes isoniazid, rifampin, ethambutol, and pyrazinamide (Table 205e-2). Isoniazid is often given together with 25–50 mg of pyridoxine daily to prevent drug-related peripheral neuropathy.

MECHANISM OF ACTION Isoniazid is a prodrug activated by the mycobacterial KatG catalase-peroxidase; isoniazid is coupled with reduced nicotinamide adenine dinucleotide (NADH). The resulting isonicotinic acyl-NADH complex blocks the mycobacterial ketoenoylreductase known as InhA, binding to its substrate and inhibiting fatty acid synthase and ultimately mycolic acid synthesis. Mycolic acids are essential components of the mycobacterial cell wall. KatG activation of isoniazid also results in the release of free radicals that have antimycobacterial activity, including nitric oxide.

The minimal inhibitory concentrations (MICs) of isoniazid for wild-type (untreated) susceptible strains are <0.1 µg/mL for *M. tuberculosis* and 0.5–2 µg/mL for *Mycobacterium kansasii*.

PHARMACOLOGY Isoniazid is the hydrazide of isonicotinic acid, a small, water-soluble molecule. The usual adult oral daily dose of 300 mg results in peak serum levels of 3–5 µg/mL within 30 min to 2 h after ingestion—well in excess of the MICs for most susceptible strains of *M. tuberculosis*. Both oral and IM preparations of isoniazid reach effective levels in the body, although antacids and high-carbohydrate meals may interfere with oral absorption. Isoniazid diffuses well throughout the body, reaching therapeutic concentrations in body cavities and fluids, with concentrations in cerebrospinal fluid (CSF) comparable to those in serum.

Isoniazid is metabolized in the liver via acetylation by *N*-acetyltransferase 2 (NAT2) and hydrolysis. Both fast- and slow-acetylation phenotypes occur; patients who are "fast acetylators" may have lower serum levels of isoniazid, whereas "slow acetylators" may have higher levels and experience more toxicity. Satisfactory isoniazid levels are attained in the majority of homozygous fast NAT2 acetylators given a dose of 6 mg/kg and in the majority of homozygous slow acetylators given only 3 mg/kg. Genotyping is increasingly being used to characterize isoniazid-related pharmacogenomic responses.

Isoniazid's interactions with other drugs are due primarily to its inhibition of the cytochrome P450 system. Among the drugs with significant isoniazid interactions are warfarin, carbamazepine, benzodiazepines, acetaminophen, clopidogrel, maraviroc, dronedarone, salmeterol, tamoxifen, eplerenone, and phenytoin.

DOSING The recommended daily dose for the treatment of TB in the United States is 5 mg/kg for adults and 10–20 mg/kg for children, with a maximal daily dose of 300 mg for both. For intermittent therapy in adults (usually twice per week), the dose is 15 mg/kg, with a maximal daily dose of 900 mg. Isoniazid does not require dosage adjustment in patients with renal disease. When the 12-dose, 3-month weekly LTBI regimen is used, the dose of isoniazid is 15 mg/kg, with a maximal dose of 900 mg, and the drug is coadministered with rifampin.

RESISTANCE Although isoniazid, along with rifampin, is the mainstay of TB treatment regimens, ~7% of clinical *M. tuberculosis* isolates in the United States are resistant. Rates of primary isoniazid resistance among untreated patients are significantly higher in many populations born outside the United States. Five separate pathways for isoniazid

resistance have been elucidated. Most strains have amino acid changes in either the catalase-peroxidase gene (*katG*) or the mycobacterial ketoenoylreductase gene (*inhA*). Less frequently, alterations in *kasA*, the gene for an enzyme involved in mycolic acid elongation, and loss of NADH dehydrogenase 2 activity confer isoniazid resistance. In 20–30% of isoniazid-resistant *M. tuberculosis* isolates, increased expression of efflux pump genes, such as *effA*, *mmpL7*, *mmr*, *p55*, and the Tap-like gene *Rv1258c*, has been implicated as the underlying mechanism of resistance.

ADVERSE EFFECTS Although isoniazid is generally well tolerated, drug-induced liver injury and peripheral neuropathy are significant adverse effects associated with this agent. Isoniazid may cause asymptomatic transient elevation of aminotransferase levels (often termed *hepatic adaptation*) in up to 20% of recipients. Other adverse reactions include rash (2%), fever (1.2%), anemia, acne, arthritic symptoms, a systemic lupus erythematosus–like syndrome, optic atrophy, seizures, and psychiatric symptoms. Symptomatic hepatitis occurs in fewer than 0.1% of persons treated with isoniazid alone for LTBI, and fulminant hepatitis with hepatic failure occurs in fewer than 0.01%. Isoniazid-associated hepatitis is idiosyncratic, but its incidence increases with age, with daily alcohol consumption, and in women who are within 3 months postpartum.

In patients who have liver disorders or HIV infection, who are pregnant or in the 3-month postpartum period, who have a history of liver disease (e.g., hepatitis B or C, alcoholic hepatitis, or cirrhosis), who use alcohol regularly, who have multiple medical problems, or who have other risk factors for chronic liver disease, the risks and benefits of treatment for LTBI should be weighed. If treatment is undertaken, these patients should have serum concentrations of alanine aminotransferase (ALT) determined at baseline. Routine baseline hepatic ALT testing based solely on an age of >35 years is optional and depends on individual concerns. Monthly biochemical monitoring during isoniazid treatment is indicated for patients whose baseline liver function tests yield abnormal results and for persons at risk for hepatic disease, including the groups just mentioned. Guidelines recommend that isoniazid be discontinued in the presence of hepatitis symptoms or jaundice and an ALT level three times the upper limit of normal or in the absence of symptoms with an ALT level five times the upper limit of normal (Table 205e-3).

Peripheral neuropathy associated with isoniazid occurs in up to 2% of patients given 5 mg/kg. Isoniazid appears to interfere with pyridoxine (vitamin B₆) metabolism. The risk of isoniazid-related neurotoxicity is greatest for patients with preexisting disorders that also pose a risk of neuropathy, such as HIV infection; for those with diabetes mellitus, alcohol abuse, or malnutrition; and for those simultaneously receiving other potentially neuropathic medications, such as stavudine. These patients should be given prophylactic pyridoxine (25–50 mg/d).

Rifampin Rifampin is a semisynthetic derivative of *Amycolatopsis rifamycinica* (formerly known as *Streptomyces mediterranei*). The most active antimycobacterial agent available, rifampin is the keystone of first-line treatment for TB. Introduced in 1968, this drug eventually permitted dramatic shortening of the TB treatment course. Rifampin has both sterilizing and bactericidal activity against dividing and nondividing *M. tuberculosis*. The drug is also active against an array of other organisms, including some gram-positive and gram-negative bacteria, *Legionella*, *M. kansasii*, and *Mycobacterium marinum*.

Rifampin, administered for 4 months, is also an alternative agent to isoniazid for the treatment of LTBI, although efficacy data are scant at this time. A 3-month course of rifampin alone has been found to be similar in efficacy to a 6-month course of isoniazid. Although the efficacy of the 4-month regimen of rifampin is under study, comparison of this regimen with 9 months of isoniazid in randomized safety and tolerability studies suggests fewer adverse events, including hepatotoxicity; less treatment interruption; a higher completion rate; and greater cost-effectiveness.

205e-4 **MECHANISM OF ACTION** Rifampin exerts both intracellular and extracellular bactericidal activity. Like other rifamycins, rifampin specifically binds to and inhibits mycobacterial DNA-dependent RNA polymerase, blocking RNA synthesis. Susceptible strains of *M. tuberculosis* as well as *M. kansasii* and *M. marinum* are inhibited by rifampin concentrations of 1 µg/mL.

PHARMACOLOGY Rifampin is a fat-soluble, complex macrocyclic molecule readily absorbed after oral administration. Serum levels of 10–20 µg/mL are achieved 2.5 h after the usual adult oral dose of 10 mg/kg (given without food). Rifampin has a half-life of 1.5–5 h. The drug distributes well throughout most body tissues, including CSF. Rifampin turns body fluids such as urine, saliva, sputum, and tears a reddish-orange color—an effect that offers a simple means of assessing patients' adherence to this medication. Rifampin is excreted primarily through the bile and enters the enterohepatic circulation; <30% of a dose is renally excreted.

As a potent inducer of the hepatic cytochrome P450 system, rifampin can decrease the half-life of some drugs, such as digoxin, warfarin, phenytoin, prednisone, cyclosporine, methadone, oral contraceptives, clarithromycin, azole antifungal agents, quinidine, antiretroviral protease inhibitors, and non-nucleoside reverse transcriptase inhibitors. The Centers for Disease Control and Prevention has issued guidelines for the management of drug interactions during treatment of HIV and *M. tuberculosis* co-infection (www.cdc.gov/tb/publications/guidelines/TB_HIV_Drugs/default.htm).

DOSING The daily dosage of rifampin is 10 mg/kg for adults and 10–20 mg/kg for children, with a maximum of 600 mg/d for both. The drug is given once daily, twice weekly, or three times weekly. No adjustments of dose or frequency are necessary in patients with renal insufficiency.

RESISTANCE Resistance to rifampin in *M. tuberculosis*, *Mycobacterium leprae*, and other organisms is the consequence of spontaneous, mostly missense point mutations in a core region of the bacterial gene coding for the β subunit of RNA polymerase (*rpoB*). RNA polymerase altered in this manner is no longer subject to inhibition by rifampin. Most rapidly and slowly growing NTM harbor intrinsic resistance to rifampin, for which the mechanism has yet to be determined.

ADVERSE EFFECTS Adverse events associated with rifampin are infrequent and generally mild. Hepatotoxicity due to rifampin alone is uncommon in the absence of preexisting liver disease and often consists of isolated hyperbilirubinemia rather than aminotransferase elevation. Other adverse reactions include rash, pruritus, gastrointestinal symptoms, and pancytopenia. Rarely, a hypersensitivity reaction may occur with intermittent therapy, manifesting as fever, chills, malaise, rash, and—in some instances—renal and hepatic failure.

Ethambutol Ethambutol is a bacteriostatic antimycobacterial agent first synthesized in 1961. A component of the standard first-line regimen, ethambutol provides synergy with the other drugs in the regimen and is generally well tolerated. Susceptible species include *M. tuberculosis*, *M. marinum*, *M. kansasii*, and organisms of the *Mycobacterium avium* complex (MAC); however, among first-line drugs, ethambutol is the least potent against *M. tuberculosis*. This agent is also used in combination with other agents in the continuation phase of treatment when patients cannot tolerate isoniazid or rifampin or are infected with organisms resistant to either of the latter drugs.

MECHANISM OF ACTION Ethambutol is bacteriostatic against *M. tuberculosis*. Its primary mechanism of action is the inhibition of the arabinosyltransferases involved in cell wall synthesis, which probably inhibits the formation of arabinogalactan and lipoarabinomannan. The MIC of ethambutol for susceptible strains of *M. tuberculosis* is 0.5–2 µg/mL.

PHARMACOLOGY AND DOSING From a single dose of ethambutol, 75–80% is absorbed within 2–4 h of administration. Serum levels peak at 2–4 µg/mL after the standard adult daily dose of 15 mg/kg. Ethambutol is well distributed throughout the body except in the CSF; a dosage of 25 mg/kg is necessary for attainment of a CSF level half of that in serum. For intermittent therapy, the dosage is 50 mg/kg twice weekly. To prevent

toxicity, the dosage must be lowered and the frequency of administration reduced for patients with renal insufficiency.

ADVERSE EFFECTS Ethambutol is usually well tolerated and has no significant interactions with other drugs. Optic neuritis, the most serious adverse effect reported, typically presents as reduced visual acuity, central scotoma, and loss of the ability to see green (or, less commonly, red). The cause of this neuritis is unknown, but it may be due to an effect of ethambutol on the amacrine and bipolar cells of the retina. Symptoms typically develop several months after initiation of therapy, but ocular toxicity soon after initiation of ethambutol has been described. The risk of ocular toxicity is dose dependent, occurring in 1–5% of patients, and can be increased by renal insufficiency. The routine use of ethambutol in younger children is not recommended because monitoring for visual complications can be difficult. If drug-resistant TB is suspected, ethambutol can be used for treatment of children.

All patients starting therapy with ethambutol should have a baseline test for visual acuity, visual fields, and color vision and should undergo an examination of the optic fundus. Visual acuity and color vision should be monitored monthly or less often as needed. Cessation of ethambutol in response to early symptoms of ocular toxicity usually results in reversal of the deficit within several months. Recovery of all visual function may take up to 1 year. In the elderly and in patients whose symptoms are not recognized early, deficits may be permanent. Some experts think that supplementation with hydroxocobalamin (vitamin B₁₂) is beneficial for patients with ethambutol-related ocular toxicity. Other adverse effects of ethambutol are rare. Peripheral sensory neuropathy occurs in rare instances.

RESISTANCE Ethambutol resistance in *M. tuberculosis* and NTM is associated primarily with missense mutations in the *embB* gene that encodes for arabinosyltransferase. Mutations have been found in resistant strains at codon 306 in 50–70% of cases. Mutations at *embB306* can cause significantly increased MICs of ethambutol, resulting in clinical resistance.

Pyrazinamide A nicotinamide analog, pyrazinamide is an important bactericidal drug used in the initial phase of TB treatment. Its administration for the first 2 months of therapy with rifampin and isoniazid allows treatment duration to be shortened from 9 months to 6 months and decreases rates of relapse.

MECHANISM OF ACTION Pyrazinamide's antimycobacterial activity is essentially limited to *M. tuberculosis*. The drug is more active against slowly replicating organisms than against actively replicating organisms. Pyrazinamide is a prodrug that is converted by the mycobacterial pyrimidase to the active form, pyrazinoic acid (POA). This agent is active only in acidic environments (pH <6.0), as are found within phagocytes or granulomas. The exact mechanism of action of POA is unclear, but fatty acid synthetase I may be the primary target in *M. tuberculosis*. Susceptible strains of *M. tuberculosis* are inhibited by pyrazinamide concentrations of 16–50 µg/mL at pH 5.5.

PHARMACOLOGY AND DOSING Pyrazinamide is well absorbed after oral administration, with peak serum concentrations of 20–60 µg/mL at 1–2 h after ingestion of the recommended adult daily dose of 15–30 mg/kg (maximum, 2 g/d). It distributes well to various body compartments, including CSF, and is an important component of treatment for tuberculous meningitis. The serum half-life of the drug is 9–11 h with normal renal and hepatic function. Pyrazinamide is metabolized in the liver to POA, 5-hydroxypyrazinamide, and 5-hydroxy-POA. A high proportion of pyrazinamide and its metabolites (~70%) is excreted in the urine. The dosage must be adjusted according to the level of renal function in patients with reduced creatinine clearance.

ADVERSE EFFECTS At the higher dosages used previously, hepatotoxicity was seen in as many as 15% of patients treated with pyrazinamide. However, at the currently recommended dosages, hepatotoxicity now occurs less commonly when this drug is administered with isoniazid and rifampin during the treatment of TB. Older age, active liver disease, HIV infection, and low albumin levels may increase the risk of

hepatotoxicity. The use of pyrazinamide with rifampin for the treatment of LTBI is no longer recommended because of unacceptable rates of hepatotoxicity and death in this setting. Hyperuricemia is a common adverse effect of pyrazinamide therapy that usually can be managed conservatively. Clinical gout is rare.



Although pyrazinamide is recommended by international TB organizations for routine use in pregnancy, it is not recommended in the United States because of inadequate teratogenicity data.

RESISTANCE The basis of pyrazinamide resistance in *M. tuberculosis* is a mutation in the *pncA* gene coding for pyrazinamidase, the enzyme that converts the prodrug to active POA. Resistance to pyrazinamide is associated with loss of pyrazinamidase activity, which prevents conversion of pyrazinamide to POA. Of pyrazinamide-resistant *M. tuberculosis* isolates, 72–98% have mutations in *pncA*. Conventional methods of testing for susceptibility to pyrazinamide may produce both false-negative and false-positive results because the high-acidity environment required for the drug's activation also inhibits the growth of *M. tuberculosis*. There is some controversy as to the clinical significance of in vitro pyrazinamide resistance.

OTHER FIRST-LINE DRUGS

Rifabutin Rifabutin, a semisynthetic derivative of rifamycin S, inhibits mycobacterial DNA-dependent RNA polymerase. Rifabutin is recommended in place of rifampin for the treatment of HIV-co-infected individuals who are taking protease inhibitors or non-nucleoside reverse transcriptase inhibitors, particularly nevirapine. Rifabutin's effect on hepatic enzyme induction is less pronounced than that of rifampin. Protease inhibitors may cause significant increases in rifabutin levels through inhibition of hepatic metabolism. Rifabutin is more active in vitro than rifampin against MAC organisms and other NTM, but its clinical superiority has not been established.

PHARMACOLOGY Like rifampin, rifabutin is lipophilic and is absorbed rapidly after oral administration, reaching peak serum levels 2–4 h after ingestion. Rifabutin distributes best to tissues, reaching levels 5–10 times higher than those in plasma. Unlike rifampin, rifabutin and its metabolites are partially cleared by the hepatic microsomal system. Rifabutin's slow clearance results in a mean serum half-life of 45 h—much longer than the 3- to 5-h half-life of rifampin. Clarithromycin (but not azithromycin) and fluconazole appear to increase rifabutin levels by inhibiting hepatic metabolism.

ADVERSE EFFECTS Rifabutin is generally well tolerated, with adverse effects occurring at higher doses. The most common adverse events are gastrointestinal; other reactions include rash, headache, asthenia, chest pain, myalgia, and insomnia. Less common adverse reactions include fever, chills, a flulike syndrome, anterior uveitis, hepatitis, *Clostridium difficile*-associated diarrhea, a diffuse polymyalgia syndrome, and yellow skin discoloration (“pseudo-jaundice”). Laboratory abnormalities include neutropenia, leukopenia, thrombocytopenia, and increased levels of liver enzymes. Approximately 80% of patients who develop rifampin-related adverse events are able to complete TB treatment with rifabutin.

RESISTANCE Similar to rifampin resistance, resistance to rifabutin is mediated by some mutations in *rpoB*.

Rifapentine Rifapentine is a semisynthetic cyclopentyl rifamycin, sharing a mechanism of action with rifampin. Rifapentine is lipophilic and has a prolonged half-life that permits weekly or twice-weekly dosing. Therefore, this drug is the subject of intensive clinical investigation aimed at determining optimal dosing and frequency of administration. Currently, rifapentine is an alternative to rifampin in the continuation phase of treatment for noncavitary drug-susceptible pulmonary TB in HIV-seronegative patients who have negative sputum smears at completion of the initial phase of treatment. When administered in these specific circumstances, rifapentine (10 mg/kg, up to 600 mg) is given once weekly with isoniazid. Because of higher rates of relapse, this regimen is not recommended for patients with TB disease and HIV coinfection. A large randomized controlled trial recently demonstrated

that, for latent TB, a 12-dose (3-month) regimen of weekly DOT with a weight-based dose of isoniazid and rifapentine was noninferior to daily isoniazid for 9 months. Although the rate of permanent drug discontinuation due to adverse events was higher with rifapentine/isoniazid, this regimen had a higher treatment completion rate than daily isoniazid in this study. The efficacy of this combination regimen in HIV-infected individuals not receiving ART and in children <12 years of age is under study. The regimen is not recommended for pregnant women, for persons with hypersensitivity reactions to isoniazid or rifampin, or for HIV-infected individuals taking ART.

PHARMACOLOGY Rifapentine's absorption is improved when the drug is taken with food. After oral administration, rifapentine reaches peak serum concentrations in 5–6 h and achieves a steady state in 10 days. The half-life of rifapentine and its active metabolite, 25-desacetyl rifapentine, is ~13 h. The administered dose is excreted via the liver (70%).

ADVERSE EFFECTS The adverse-effects profile of rifapentine is similar to that of other rifamycins. Rifapentine is teratogenic in animal models and is relatively contraindicated in pregnancy.

RESISTANCE Rifapentine resistance is mediated by mutations in *rpoB*. Mutations that cause resistance to rifampin also cause resistance to rifapentine.



Streptomycin Streptomycin was the first antimycobacterial agent used for the treatment of TB. Derived from *Streptomyces griseus*, streptomycin is bactericidal against dividing *M. tuberculosis* organisms but has only low-level early bactericidal activity. This drug is administered only by the IM and IV routes. In developed nations, streptomycin is used infrequently because of its toxicity, the inconvenience of injections, and drug resistance. In developing countries, however, streptomycin is used because of its low cost.

MECHANISM OF ACTION Streptomycin inhibits protein synthesis by binding at a site on the 30S mycobacterial ribosome.

PHARMACOLOGY AND DOSING Serum levels of streptomycin peak at 25–45 µg/mL after a 1-g dose. This agent penetrates poorly into the CSF, reaching levels that are only 20% of serum levels. The usual daily dose of streptomycin (given IM either daily or 5 days per week) is 15 mg/kg for adults and 20–40 mg/kg for children, with a maximum of 1 g/d for both. For patients ≥60 years of age, 10 mg/kg is the recommended daily dose, with a maximum of 750 mg/d. Because streptomycin is eliminated almost exclusively by the kidneys, its use in patients with renal impairment should be avoided or implemented with caution, with lower doses and less frequent administration.

ADVERSE EFFECTS Adverse reactions occur frequently with streptomycin (10–20% of patients). Ototoxicity (primarily vestibulotoxicity), neuropathy, and renal toxicity are the most common and the most serious. Renal toxicity, usually manifested as nonoliguric renal failure, is less common with streptomycin than with other frequently used aminoglycosides, such as gentamicin. Manifestations of vestibular toxicity include loss of balance, vertigo, and tinnitus. Patients receiving streptomycin must be monitored carefully for these adverse effects, undergoing audiometry at baseline and monthly thereafter.

RESISTANCE Spontaneous mutations conferring resistance to streptomycin are relatively common, occurring in 1 in 10⁶ organisms. In the two-thirds of streptomycin-resistant *M. tuberculosis* strains exhibiting high-level resistance, mutations have been identified in one of two genes: a 16S rRNA gene (*rrs*) or the gene encoding ribosomal protein S12 (*rpsL*). Both targets are believed to be involved in streptomycin ribosomal binding. However, low-level resistance, which is seen in about one-third of resistant isolates, has no associated resistance mutation. A gene (*gidB*) that confers low-level resistance to streptomycin has recently been identified. Strains of *M. tuberculosis* resistant to streptomycin generally are not cross-resistant to capreomycin or amikacin. Streptomycin is not used for the treatment of MDR-TB or XDR-TB because of (1) the high prevalence of streptomycin resistance among strains resistant to isoniazid and (2) the unreliability of drug susceptibility testing.

Second-line antituberculosis agents are indicated for treatment of drug-resistant TB, for patients who are intolerant or allergic to first-line agents, and when first-line supplemental agents are unavailable.

Fluoroquinolones Fluoroquinolones inhibit mycobacterial DNA gyrase and topoisomerase IV, preventing cell replication and protein synthesis, and are bactericidal. The later-generation fluoroquinolones levofloxacin and moxifloxacin are the most active against *M. tuberculosis* and are recommended for the treatment of MDR-TB. They are also being investigated for their potential to shorten the course of treatment for TB. In a recent trial, gatifloxacin, which had been withdrawn from the market because of significant dysglycemia, was assessed for treatment shortening; although its inclusion in the TB treatment regimen did not shorten the duration of therapy from 6 to 4 months, the drug did not cause dysglycemia in TB patients who took it thrice weekly for 4 months. Ciprofloxacin and ofloxacin are no longer recommended for the treatment of TB because of poor efficacy. Despite the documented resistance of the infecting strains to these and other early-generation fluoroquinolones, use of a later-generation fluoroquinolone in patients with XDR-TB has been associated with favorable outcomes. Fluoroquinolones are also considered safe alternatives for patients who develop treatment-limiting adverse effects due to first-line agents. Levofloxacin and moxifloxacin have both been used effectively in the treatment of MDR-TB. The optimal dose of levofloxacin for this indication is being actively studied, but doses of at least 750 mg are commonly used.

The fluoroquinolones are well absorbed orally, reach high serum levels, and distribute well into body tissues and fluids. Their absorption is decreased by co-ingestion with products containing multivalent cations, such as antacids. Adverse effects are relatively infrequent (0.5–10% of patients) and include gastrointestinal intolerance, rashes, dizziness, and headache. Most studies of fluoroquinolone side effects have been based on relatively short-term administration for bacterial infections, but trials have now shown the relative safety and tolerability of fluoroquinolones administered for months during TB treatment in adults. Although the potential to prolong the QTc interval, leading to cardiac arrhythmias, has been a source of concern with fluoroquinolones, cessation of treatment due to this adverse effect is rare. There is increasing interest in the use of fluoroquinolones in children, which has traditionally been avoided because of the risks of tendon rupture and cartilage damage, because the benefits in treatment of drug-resistant TB may outweigh the risks.

Mycobacterial resistance can develop rapidly when a fluoroquinolone is inadvertently administered alone. Empirical fluoroquinolone therapy for presumed community-acquired pneumonia is associated with increased fluoroquinolone resistance in *M. tuberculosis*. Mutations in the genes encoding for DNA gyrase (*gyrA* and *gyrB*) are implicated in the majority of cases—but not all cases—of clinical resistance to fluoroquinolones.

Injectable Agents • CAPREOMYCIN Capreomycin, a cyclic peptide antibiotic derived from *Streptomyces capreolus*, is an important first-choice second-line agent used for treatment of MDR-TB, particularly when additional resistance to aminoglycosides is documented. Capreomycin is administered by the IM route; an inhaled preparation is under study. A dose of 15 mg/kg per day is given five to seven times per week (maximal daily dose, 1 g) and results in peak blood levels of 20–40 µg/mL. The dosage may be reduced to 1 g two or three times per week 2–4 months after mycobacterial cultures become negative. For individuals ≥60 years of age, the dose should be reduced to 10 mg/kg per day (maximal daily dose, 750 mg). For patients with renal insufficiency, the drug should be given intermittently and at lower dosage (12–15 mg/kg two or three times per week). A minimal duration of 3 months is recommended for MDR-TB treatment. Penetration of capreomycin into the CSF is believed to be poor.

The mechanism of capreomycin's action is not well understood but involves interference with the mycobacterial ribosome and inhibition of

protein synthesis. Resistance to capreomycin is associated with mutations that inactivate a ribosomal methylase (*tlyA*) or that encode genes for the 16S ribosomal subunit (*rrs*). Cross-resistance to kanamycin and amikacin is common with *rrs* but not always with *tlyA* mutations. However, some strains that are resistant to streptomycin, kanamycin, and amikacin generally remain susceptible to capreomycin.

Adverse effects of capreomycin are relatively common. Significant hypokalemia and hypomagnesemia as well as oto- and renal toxicity have been reported.

AMIKACIN AND KANAMYCIN Amikacin and kanamycin are aminoglycosides that exert mycobactericidal activity by binding to the 16S ribosomal subunit. The spectrum of antibiotic activity for amikacin and kanamycin includes *M. tuberculosis*, several NTM species, and aerobic gram-negative and gram-positive bacteria. Although amikacin is highly active against *M. tuberculosis*, it is used only infrequently because of its significant side effects. The usual daily adult dosage of both amikacin and kanamycin is 15–30 mg/kg given IM or IV (maximal daily dose, 1 g), with a reduction to 10 mg/kg for patients ≥60 years old. For patients with renal insufficiency, the dose and frequency should be reduced (12–15 mg/kg two or three times per week). Mycobacterial resistance is due to mutations in the genes encoding the 16S ribosomal RNA gene. Cross-resistance among kanamycin, amikacin, and capreomycin is common. Isolates resistant to streptomycin are frequently susceptible to amikacin or kanamycin. Adverse effects of amikacin include ototoxicity (in up to 10% of recipients, with auditory dysfunction occurring more commonly than vestibulotoxicity), nephrotoxicity, and neurotoxicity. Kanamycin has a similar side-effects profile, but adverse reactions are thought to be less frequent and less severe.

Other Second-Line Agents • ETHIONAMIDE Ethionamide is a derivative of isonicotinic acid. Its mechanism of action is through inhibition of the *inhA* gene product enoyl-acyl carrier protein (acp) reductase, which is involved in mycolic acid synthesis. Ethionamide is bacteriostatic against metabolically active *M. tuberculosis* and some NTM. It is used in the treatment of drug-resistant TB, but its use is limited by severe gastrointestinal reactions (including abdominal pain, nausea, and vomiting) as well as significant central and peripheral neurologic side effects, reversible hepatitis (in ~5% of recipients), hypersensitivity reactions, and hypothyroidism. Ethionamide should be taken with food to reduce gastrointestinal effects and with pyridoxine (50–100 mg/d) to limit neuropathic side effects.

CYCLOSERINE Cycloserine is an analog of the amino acid D-alanine and prevents cell wall synthesis. It inhibits the action of enzymes, including alanine racemase, that are involved in the production of peptidoglycans. Cycloserine is active against a range of bacteria, including *M. tuberculosis*. Mechanisms of mycobacterial resistance are not well understood, but overexpression of alanine racemase can confer resistance in *Mycobacterium smegmatis*. Cycloserine is well absorbed after oral administration and is widely distributed throughout body fluids, including CSF. The usual adult dosage is 250 mg two or three times per day. Serious potential side effects include seizures and psychosis (with suicide in some cases), peripheral neuropathy, headache, somnolence, and allergic reactions. Drug levels are monitored to achieve optimal dosing and to reduce the risk of adverse effects, especially in patients with renal failure. Cycloserine should be administered as DOT only with caution and with support from experienced TB physicians to patients with epilepsy, active alcohol abuse, severe renal insufficiency, or a history of depression or psychosis.

PARA-AMINOSALICYLIC ACID Para-aminosalicylic acid (PAS, 4-aminosalicylic acid) is an oral agent used in the treatment of MDR- and XDR-TB. Its bacteriostatic activity is due to inhibition of folate synthesis and of iron uptake. PAS has relatively little activity as an anti-TB agent. Adverse effects may include high-level nausea, vomiting, and diarrhea. PAS may cause hemolysis in patients with glucose-6-phosphate dehydrogenase deficiency. The drug should be taken with acidic foods to improve absorption. Enteric-coated PAS granules (4 g orally

every 8 h) appear to be better tolerated than other formulations and produce higher therapeutic blood levels. PAS has a short half-life (1 h), and 80% of the dose is excreted in the urine.

CLOFAZIMINE Clofazimine is a fat-soluble riminophenazine dye used primarily in the treatment of leprosy worldwide. It is currently gaining popularity in the management of MDR- and XDR-TB because of its low cost and intracellular and extracellular activity. By increasing reactive oxygen species and causing membrane destabilization, clofazimine may promote killing of antibiotic-tolerant *M. tuberculosis* persister organisms. In addition to antimicrobial activity, the drug has other pharmacologic properties—e.g., anti-inflammatory, pro-oxidative, and immunopharmacologic. Clofazimine has a half-life of ~70 days in humans, and average steady-state concentrations are achieved at ~1 month. Ingestion with fatty meals can improve its low and variable rates of absorption (45–62%). Common side effects include gastrointestinal intolerance and reversible orange-to-brownish discoloration of the skin, bodily fluids, and secretions. Dose adjustment may be necessary in patients with severe hepatic impairment. Clofazimine is being studied as part of a regimen developed in Bangladesh for potential shortening of the MDR-TB treatment course. A recent meta-analysis suggested that inclusion of clofazimine in a multidrug regimen for treatment of MDR-TB was associated with a favorable outcome. Newer analogues with improved pharmacokinetics and alternative formulations of clofazimine (liposomal, nanosuspension, inhalational) are being studied.

NEWER ANTITUBERCULOSIS DRUGS

Oxazolidinones Linezolid is an oxazolidinone used primarily for the treatment of drug-resistant gram-positive bacterial infections. However, this drug is active in vitro against *M. tuberculosis* and NTM. Several case series have suggested that linezolid may help clear mycobacteria relatively rapidly when included in a regimen for the treatment of complex cases of MDR- and XDR-TB. Linezolid's mechanism of action is disruption of protein synthesis by binding to the 50S bacterial ribosome. Linezolid has nearly 100% oral bioavailability, with good penetration into tissues and fluids, including CSF. Clinical resistance to linezolid has been reported, but the mechanism is unclear. Adverse effects may include optic and peripheral neuropathy, pancytopenia, and lactic acidosis. Linezolid is a weak monoamine oxidase inhibitor and can be associated with the serotonin syndrome when given concomitantly with serotonergic drugs (primarily antidepressants such as selective serotonin-reuptake inhibitors). A recent meta-analysis showed that ~80% of patients with MDR- or XDR-TB can be successfully treated with linezolid-containing anti-TB regimens; however, significant adverse events attributed to linezolid were reported. For MDR-TB treatment, linezolid is usually administered at a dose of 600 mg (or less in some cases) once daily, which appears to be effective. The single daily dose is associated with fewer adverse events than twice-a-day dosing.

PNU 100480 and AZD 5847, modified versions of oxazolidinones and protein synthesis inhibitors, are undergoing phase 1 trials and appear to have greater efficacy than linezolid against *M. tuberculosis*. However, the adverse effect profiles of these compounds compared with that of linezolid need further investigation.

Amoxicillin-Clavulanate and Carbapenems β -Lactam agents are largely ineffective for the treatment of *M. tuberculosis* because of resistance conferred by a hydrolyzing class A β -lactamase. Because clavulanate may theoretically inhibit the β -lactamase, amoxicillin-clavulanate has been used in the treatment of MDR-TB; however, it is a comparatively weak agent. Carbapenems are poor substrates for class A β -lactamases found in *M. tuberculosis*. Accordingly, meropenem and imipenem have in vitro activity against *M. tuberculosis*, and their use to treat MDR- and XDR-TB has been reported anecdotally. Nevertheless, the need to administer carbapenems by the IV route and lack of information on the drugs' long-term side effects have restricted their use to certain severe cases only.

Diarylquinolines Bedaquiline (TMC207 or R207910) is a new diarylquinoline with a novel mechanism of action: inhibition of the mycobacterial

ATP synthetase proton pump. TMC207 is bactericidal for drug-susceptible and MDR strains of *M. tuberculosis*. Resistance has been reported and is due to point mutations in the *atpE* gene encoding for subunit c of ATP synthetase. A phase 2 randomized controlled clinical trial in MDR-TB patients demonstrated substantial improvement in 2-month culture-conversion rates as well as a reduction in acquired resistance to companion drugs. This drug is metabolized by the hepatic cytochrome CYP3A4. Rifampin lowers TMC207 levels by 50%, and protease inhibitors also interact significantly with this drug. The oral bioavailability of TMC207 appears to be excellent. The dosage is 400 mg/d for the first 2 weeks and then 200 mg thrice weekly. The elimination half-life is long (>14 days). A single dose of this drug can inhibit the growth of *M. tuberculosis* for up to 1 week through a combination of long plasma half-life, high-level tissue penetration, and long tissue half-life. Bedaquiline added to a background regimen improved the 2-month sputum culture conversion rate in multicenter, randomized placebo-controlled trials, and these results led to approval by the U.S. Food and Drug Administration (FDA). However, a higher death rate in one trial was observed in the bedaquiline arm than in the control arm (11.4% vs 2.5%); the result was a “black box” warning from the FDA, which also included QT prolongation. The Centers for Disease Control and Prevention has made a provisional recommendation for the use of bedaquiline for 24 weeks in adults with laboratory-confirmed pulmonary MDR-TB when no other effective treatment regimen can be provided.

Nitroimidazoles The prodrugs delamanid (OPC-67683) and PA 824 are novel nitro-dihydro-imidazooxazole derivatives that are activated by *M. tuberculosis*-specific flavin-dependent nitroreductases whose antimycobacterial activity is attributable to inhibition of mycolic acid biosynthesis. These drugs are currently in phase 2 clinical trials and show potential in shortening treatment duration through their activity against nonreplicating drug-susceptible and drug-resistant mycobacteria. Delamanid was shown in a randomized, placebo-controlled, multinational clinical trial to significantly improve the culture conversion rate at 2 months. QT prolongation occurred significantly more often in delamanid-treated patients, but no clinically relevant events were reported.

Diamines SQ109, an ethambutol analogue with a 1,2-diamine pharmacophore, is the most promising of the diamines for TB treatment. It is activated by mycobacterial cytochrome enzymes and inhibits mycobacterial cell-wall synthesis by an unknown mechanism. It has a high tissue protein-binding capacity with a very long half-life (~61 h) in humans. In vitro studies have demonstrated that SQ109 has low MICs against both susceptible and resistant *M. tuberculosis* strains as well as a synergistic effect when administered with isoniazid and rifampin. The drug is under study in clinical trials for TB treatment.

Pyrroles LL3858, a pyrrole derivative, has entered clinical trials examining its utility in the treatment of drug-susceptible and drug-resistant TB. The drug's mechanism of action is unknown. However, because it is active against *M. tuberculosis* strains that are resistant to available anti-TB drugs, its target is thought to differ from those of currently used agents.

NONTUBERCULOUS MYCOBACTERIA

More than 150 species of NTM have been identified. Only a minority of these environmental organisms, which are found in soil and water, are important human pathogens. NTM cause extensive disease, primarily in persons with preexisting pulmonary disease or immunocompromise, but also can cause nodular/bronchiectatic disease in otherwise seemingly healthy hosts. NTM are also important causes of infections in surgical settings. The two major classes of NTM are the slow-growing and rapidly growing species; subcultures of the latter grow within 1 week. The growth characteristics of NTM have diagnostic, therapeutic, and prognostic implications. The rate of growth can provide useful preliminary information within a specific clinical context, in that growth within 2–3 weeks is much more likely to indicate an NTM than *M. tuberculosis*. When NTM do grow from cultures, colonization should be distinguished from active disease in order to optimize the

risk and benefit of prolonged treatment with multiple medications. According to the recommendations of the American Thoracic Society and the Infectious Diseases Society of America, significant clinical manifestations and/or sputum radiographic evidence of progressive disease consistent with NTM infection as well as either reproducible sputum culture results or a single positive culture are required for the diagnosis of NTM pulmonary disease. Isolation of NTM from blood or from an infected-appearing extrapulmonary site, such as soft tissue or bone, is usually indicative of disseminated or local NTM infection (Chap. 204). Treatment of NTM disease is prolonged and requires multiple medications. Side effects of the regimens employed are common, and intermittent therapy is often used to mitigate these adverse events. Treatment regimens depend on the NTM species, the extent or type of disease, and—to some degree—drug susceptibility test results. The nodular bronchiectatic form of MAC infection is generally treated three times per week, whereas fibrocavitary or disseminated MAC infection is treated daily.

THERAPEUTIC CONSIDERATIONS FOR SPECIFIC NTM

M. avium Complex Among the NTM, MAC organisms most commonly cause human disease. In immunocompetent hosts, MAC species are most often found in conjunction with underlying significant lung disease, such as chronic obstructive pulmonary disease or bronchiectasis. For patients with nodular or bronchiectatic MAC lung disease, an initial regimen consisting of clarithromycin or azithromycin, rifampin or rifabutin, and ethambutol is given three times per week. Routine initial testing for macrolide resistance is recommended, as is testing at 6 months with a failing regimen (i.e., with cultures persistently positive for NTM).

In immunocompromised individuals, disseminated MAC infection is generally treated with clarithromycin, ethambutol, and rifabutin. Azithromycin may be substituted in patients unable to tolerate clarithromycin. Amikacin and fluoroquinolones are often used in salvage regimens. Treatment for disseminated MAC infection in AIDS patients may be lifelong in the absence of immune reconstitution. At least 12 months of MAC therapy and 6 months of effective immune reconstitution may be adequate.

M. kansasii *M. kansasii* is the second most common NTM causing human disease. It is also the second most common cause of NTM pulmonary disease in the United States, where it is most often reported in the southeastern region. *M. kansasii* infection can be treated with isoniazid, rifampin, and ethambutol; therapy continues for 12 months after culture conversion. Rifampin-resistant *M. kansasii* has been treated with clarithromycin, trimethoprim-sulfamethoxazole, and streptomycin.

Rapidly Growing Mycobacteria Rapidly growing mycobacteria causing human disease include *Mycobacterium abscessus*, *Mycobacterium fortuitum*, and *Mycobacterium chelonae*. Treatment of these mycobacteria is complex and should be undertaken with input from experienced clinicians. Testing for macrolide resistance is recommended. However, in rapidly growing mycobacteria, an inducible *erm* gene may confer in vivo macrolide resistance to isolates that are susceptible in vitro.

M. marinum *M. marinum* is an NTM found in salt water and freshwater, including swimming pools and fish tanks. It is a cause of localized soft-tissue infections, which may require surgical management. Combination regimens include clarithromycin and either ethambutol or rifampin. Other agents with activity against *M. marinum* include doxycycline, minocycline, and trimethoprim-sulfamethoxazole.

DRUGS FOR THE TREATMENT OF NTM

Clarithromycin

Clarithromycin is a macrolide antibiotic with broad activity against many gram-positive and gram-negative bacteria as well as NTM. This drug is active against MAC organisms and many other NTM species, inhibiting protein synthesis by binding to the 50S mycobacterial ribosomal subunit. NTM resistance to macrolides is probably caused by overexpression of the gene *ermB*, with consequent methylation of the binding site. Clarithromycin is well absorbed orally and distributes well to tissues. It is cleared both hepatically and renally; the dosage should be reduced in renal insufficiency. Clarithromycin is a substrate for and inhibits cytochrome 3A4 and should not be administered with cisapride, pimozide, or terfenadine because cardiac arrhythmias may occur. Numerous drugs interact with clarithromycin through the CYP3A4 metabolic pathway. Rifampin lowers clarithromycin levels; conversely, rifampin levels are increased by clarithromycin. However, the clinical relevance of this interaction does not appear to be great.

For patients with nodular/bronchiectatic MAC infection, the dosage of clarithromycin is 500 mg, given morning and evening three times a week. For the treatment of fibrocavitary or severe nodular/bronchiectatic MAC infection, a dose of 500–1000 mg is given daily. Disseminated MAC infection is treated with 1000 mg daily. Clarithromycin is used in combination regimens that typically include ethambutol and a rifamycin in order to avoid the development of macrolide resistance. Adverse effects include frequent gastrointestinal intolerance, hepatotoxicity, headache, rash, and rare instances of hypoglycemia. Clarithromycin is contraindicated during pregnancy because of its teratogenicity in animal models.

Azithromycin Azithromycin is a derivative of erythromycin. Although technically an azalide and not a macrolide, it works similarly to macrolides, inhibiting protein synthesis through binding to the 50S ribosomal subunit. Resistance to azithromycin is almost always associated with complete cross-resistance to clarithromycin. Azithromycin is well absorbed orally, with good tissue penetration and a prolonged half-life (~48 h). The usual dosage for treatment of MAC infection is 250 mg/d or 500 mg three times per week. Azithromycin is used in combination with other agents to avoid the development of resistance. For prophylaxis against disseminated MAC infection in immunocompromised individuals, a dose of 1200 mg once per week is given. Because azithromycin is not metabolized by cytochrome P450, it interacts with few drugs. Adjustment of the dosage on the basis of renal function is not necessary.

Cefoxitin Cefoxitin is a second-generation parenteral cephalosporin with activity against rapidly growing NTM, particularly *M. abscessus*, *M. marinum*, and *M. chelonae*. Its mechanism of action against NTM is unknown but may involve inactivation of cell wall synthesis enzymes. High doses are used for treatment of NTM: 200 mg/kg IV three or four times per day, with a maximal daily dose of 12 g. The half-life of cefoxitin is ~1 h, with primarily renal clearance that requires adjustment in renal insufficiency. Adverse effects are uncommon but include gastrointestinal manifestations, rash, eosinophilia, fever, and neutropenia.

CONCLUSION

Treatment of mycobacterial infections requires multiple-drug regimens that often exert significant side effects with the potential to limit tolerability. The prolonged duration of treatment has vastly improved results over those obtained in past decades, but drugs and regimens that will shorten treatment duration and limit adverse drug effects and interactions are needed.

SECTION 9 SPIROCHETAL DISEASES

206 Syphilis

Sheila A. Lukehart

DEFINITION

Syphilis, a chronic systemic infection caused by *Treponema pallidum* subspecies *pallidum*, is usually sexually transmitted and is characterized by episodes of active disease interrupted by periods of latency. After an incubation period averaging 2–6 weeks, a primary lesion appears—often associated with regional lymphadenopathy—that resolves without treatment. The secondary stage, associated with generalized mucocutaneous lesions and generalized lymphadenopathy, is followed by a latent period of subclinical infection lasting years or decades. Central nervous system (CNS) involvement may occur early in infection and may be symptomatic or asymptomatic. In the preantibiotic era, about one-third of patients

with untreated cases developed the tertiary stage, characterized by progressive destructive mucocutaneous, musculoskeletal, or parenchymal lesions; aortitis; or late CNS manifestations.

ETIOLOGY

The Spirochaetales include four genera that are pathogenic for humans and for a variety of other animals: *Leptospira* species, which cause leptospirosis (Chap. 208); *Borrelia* species, which cause relapsing fever and Lyme disease (Chaps. 209 and 210); *Brachyspira* species, which cause intestinal infections; and *Treponema* species, which cause the diseases known collectively as *treponematoses* (see also Chap. 207e). The *Treponema* species include *T. pallidum* subspecies *pallidum*, which causes venereal syphilis; *T. pallidum* subspecies *pertenue*, which causes yaws; *T. pallidum* subspecies *endemicum*, which causes endemic syphilis or bejel; and *T. carateum*, which causes pinta. Until recently, the subspecies were distinguished primarily by the clinical syndromes they produce. Researchers have now identified molecular signatures

that can differentiate the three subspecies of *T. pallidum* by culture-independent methods based on polymerase chain reaction (PCR), but other sequence signatures cross subspecies boundaries in certain strains. Other *Treponema* species found in the human mouth, genital mucosa, and gastrointestinal tract have been associated with disease (e.g., periodontitis), but their role as primary etiologic agents is unclear.

T. pallidum subspecies *pallidum* (referred to hereafter as *T. pallidum*), a thin spiral organism, has a cell body surrounded by a trilaminar cytoplasmic membrane, a delicate peptidoglycan layer providing some structural rigidity, and a lipid-rich outer membrane containing relatively few integral membrane proteins. Endoflagella wind around the cell body in the periplasmic space and are responsible for motility.



T. pallidum cannot be cultured in vitro, and little was known about its metabolism until the genome was sequenced in 1998. This spirochete possesses severely limited metabolic capabilities, lacking the genes required for de novo synthesis of most amino acids, nucleotides, and lipids. In addition, *T. pallidum* lacks genes encoding the enzymes of the Krebs cycle and oxidative phosphorylation. The organism contains numerous compensatory genes predicted to encode transporters of amino acids, carbohydrates, and lipids. In addition, genome analyses and other studies have revealed the existence of a 12-member gene family (*tpr*) that bears similarities to variable outer-membrane antigens of other spirochetes. One member, *TprK*, has discrete variable (V) regions that undergo antigenic variation during infection, providing a mechanism for immune evasion.

The only known natural host for *T. pallidum* is the human. *T. pallidum* can infect many mammals, but only humans, higher apes, and a few laboratory animals regularly develop syphilitic lesions. Rabbits are used to propagate virulent strains of *T. pallidum* and serve as the animal model that best reflects human disease and immunopathology.

TRANSMISSION AND EPIDEMIOLOGY

Nearly all cases of syphilis are acquired by sexual contact with infectious lesions (i.e., the chancre, mucous patch, skin rash, or condylomata lata; see Fig. 25e-20). Less common modes of transmission include nonsexual personal contact, infection in utero, blood transfusion, and organ transplantation.

SYPHILIS IN THE UNITED STATES

With the advent of penicillin therapy, the total number of cases of syphilis reported annually in the United States declined significantly to a low of 31,575 cases in 2000—a 95% decrease from 1943—with <6000 reported cases of infectious primary and secondary syphilis (the latter being a better indicator of disease activity than total syphilis cases). Since 2000, the number of cases of primary and secondary syphilis has more than doubled, with more than 14,000 cases reported in 2012 (Fig. 206-1). Approximately 70% of these cases were in men who have

sex with men (MSM), 20–70% of whom are co-infected with HIV (depending on geographic location). The number of primary and secondary cases among women in the United States increased from 2004 to 2008 but has since been declining in conjunction with a decline in congenital syphilis. Surveillance of the number of new cases of primary and secondary syphilis has revealed multiple 7- to 10-year cycles, which may be attributed to herd immunity in at-risk populations, changing sexual behaviors, and changes in control efforts.

The populations at highest risk for acquiring syphilis have changed over time, with outbreaks among MSM in the pre-HIV era of the late 1970s and early 1980s as well as at present. It is speculated that recent increases in syphilis and other sexually transmitted infections in MSM may be due to unprotected sex between persons who are HIV concordant and to disinhibition caused by highly effective antiretroviral therapies. The syphilis epidemic that peaked in 1990 predominantly affected African-American heterosexual men and women and occurred largely in urban areas, where infectious syphilis was correlated with the exchange of sex for crack cocaine. The rate of primary and secondary syphilis among African Americans nearly doubled between 2003 and 2009, remains higher than rates for other racial/ethnic groups, but has since declined somewhat (Fig. 206-1).

The incidence of congenital syphilis roughly parallels that of infectious syphilis in women. In 2011, 360 cases in infants <1 year of age were reported, for a decline of 20% since 2008. The case definition for congenital syphilis was broadened in 1989 and now includes all live or stillborn infants delivered to women with untreated or inadequately treated syphilis.

One-third to one-half of individuals named as sexual contacts of persons with infectious syphilis become infected. Many have already developed manifestations of syphilis when they are first seen, and ~30% of asymptomatic contacts examined within 30 days of exposure actually have incubating infection and will later develop infectious syphilis if not treated. Thus, identification and treatment of all recently exposed sexual contacts continue to be important aspects of syphilis control.

GLOBAL SYPHILIS



Syphilis remains a significant health problem globally; the number of new infections is estimated at 11 million per year. The regions that are most affected include sub-Saharan Africa, South America, China, and Southeast Asia. During the past decade, the incidence rate in China has increased by approximately eightfold, and higher rates of infectious syphilis have been reported among MSM in many European countries. Worldwide, there are estimated to be 1.4 million cases of syphilis among pregnant women, with 500,000 adverse pregnancy outcomes annually (e.g., stillbirth, neonatal and early fetal death, prematurity/low birth weight, and infection in newborns). Congenital syphilis rates in China are ~150 cases per 100,000 live births.

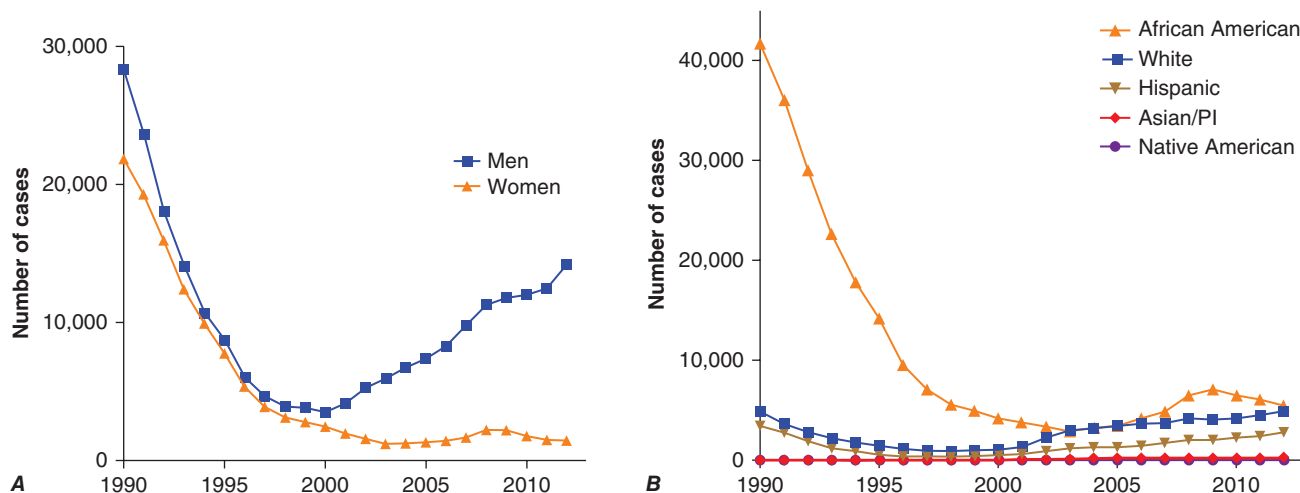


FIGURE 206-1 Primary and secondary syphilis in the United States, 1990–2012, by sex (A) and by race or ethnicity (B). (Data from the Centers for Disease Control and Prevention.)

T. pallidum rapidly penetrates intact mucous membranes or microscopic abrasions in skin and, within a few hours, enters the lymphatics and blood to produce systemic infection and metastatic foci long before the appearance of a primary lesion. Blood from a patient with incubating or early syphilis is infectious. The generation time of *T. pallidum* during early active disease in vivo is estimated to be ~30 h, and the incubation period of syphilis is inversely proportional to the number of organisms inoculated. The 50% infectious dose for intradermal inoculation in humans has been calculated to be 57 organisms, and the treponeme concentration generally reaches 10^7 /g of tissue before a clinical lesion appears. The median incubation period in humans (~21 days) suggests an average inoculum of 500–1000 infectious organisms for naturally acquired disease; the incubation period rarely exceeds 6 weeks.

The primary lesion appears at the site of inoculation, usually persists for 4–6 weeks, and then heals spontaneously. Histopathologic examination shows perivascular infiltration, chiefly by CD4+ and CD8+ T lymphocytes, plasma cells, and macrophages, with capillary endothelial proliferation and subsequent obliteration of small blood vessels. The cellular infiltration displays a T_H1 -type cytokine profile consistent with the activation of macrophages. Phagocytosis of opsonized organisms by activated macrophages ultimately causes their destruction, resulting in spontaneous resolution of the chancre.

The generalized parenchymal, constitutional, and mucocutaneous manifestations of secondary syphilis usually appear ~6–8 weeks after the chancre heals, although primary and secondary manifestations may overlap. In contrast, some patients may enter the latent stage without ever recognizing secondary lesions. The histopathologic features of secondary maculopapular skin lesions include hyperkeratosis of the epidermis, capillary proliferation with endothelial swelling in the superficial dermis, dermal papillae with transmigration of polymorphonuclear leukocytes, and—in the deeper dermis—perivascular infiltration by CD8+ T lymphocytes, CD4+ T lymphocytes, macrophages, and plasma cells. Treponemes are found in many tissues, including the aqueous humor of the eye and the cerebrospinal fluid (CSF). *T. pallidum* invades the CNS during the first weeks or months of infection, and CSF abnormalities are detected in as many as 40% of patients during the secondary stage. Clinical hepatitis and immune complex–induced glomerulonephritis are relatively rare but recognized manifestations of secondary syphilis; liver function tests may yield abnormal results in up to one-quarter of patients with early syphilis. Generalized nontender lymphadenopathy is noted in 85% of patients with secondary syphilis. The paradoxical appearance of secondary manifestations despite high titers of antibody (including immobilizing antibody) to *T. pallidum* may result from immune evasion due to antigenic variation or changes in expression of surface antigens. Secondary lesions generally subside within 2–6 weeks, and the infection enters the latent stage, which is detectable only by serologic testing. In the preantibiotic era, up to 25% of untreated patients experienced at least one generalized or localized mucocutaneous relapse, usually during the first year. Therefore, identification and examination of sexual contacts are most important for patients with syphilis of <1 year's duration.

As stated earlier, about one-third of patients with untreated latent syphilis developed clinically apparent tertiary disease in the preantibiotic era, when the most common types of tertiary disease were the gumma (a usually benign granulomatous lesion); cardiovascular syphilis (usually involving the vasa vasorum of the ascending aorta and resulting in aneurysm); and late symptomatic neurosyphilis (tabes dorsalis and paresis). In Western countries today, specific treatment for early and latent syphilis and coincidental therapy (i.e., therapy with antibiotics that are given for other conditions but are active against treponemes) have nearly eliminated tertiary syphilis. Asymptomatic CNS involvement, however, is still demonstrable in up to 40% of persons with early syphilis and 25% of patients with late latent syphilis, and cases of general paresis and tabes dorsalis are being reported from China. The factors that contribute to the development and progression of tertiary disease are unknown.

The course of untreated syphilis was studied retrospectively in a group of nearly 2000 patients with primary or secondary disease diagnosed

clinically (the Oslo Study, 1891–1951) and was assessed prospectively in 431 African-American men with seropositive latent syphilis of ≥ 3 years' duration (the notorious Tuskegee Study, 1932–1972). In the Oslo Study, 24% of patients developed relapsing secondary lesions within 4 years, and 28% eventually developed one or more manifestations of tertiary syphilis. Cardiovascular syphilis, including aortitis, was detected in 10% of patients; 7% of patients developed symptomatic neurosyphilis, and 16% developed benign tertiary gummatous syphilis. Syphilis was the primary cause of death in 15% of men and 8% of women. Cardiovascular syphilis was documented in 35% of men and 22% of women who eventually came to autopsy. In general, serious late complications were nearly twice as common among men as among women.

The Tuskegee Study showed that the death rate among untreated African-American men with syphilis (25–50 years old) was 17% higher than the rate among uninfected subjects and that 30% of all deaths were attributable to cardiovascular or, to a lesser extent, CNS syphilis. Anatomic evidence of aortitis was found in 40–60% of autopsied subjects with syphilis (vs 15% of control subjects), whereas CNS syphilis was found in only 4%. Rates of hypertension were also higher among the infected subjects. The ethical issues eventually raised by this study, begun in the preantibiotic era but continuing into the early 1970s, had a major influence on the development of current guidelines for human medical experimentation, and the history of the study may still contribute to a reluctance of some African Americans to participate as subjects in clinical research.

CLINICAL MANIFESTATIONS

Primary Syphilis The typical primary chancre usually begins as a single painless papule that rapidly becomes eroded and usually becomes indurated, with a characteristic cartilaginous consistency on palpation of the edge and base of the ulcer. Multiple primary lesions are seen in a minority of patients. In heterosexual men the chancre is usually located on the penis (Fig. 206-2; see also Fig. 25e-17), whereas in MSM it may be found in the anal canal or rectum, in the mouth, or on the external genitalia. Oral sex has been identified as the source of infection in some MSM. In women, common primary sites are the cervix and labia. Consequently, primary syphilis goes unrecognized in women and homosexual men more often than in heterosexual men.



FIGURE 206-2 Primary syphilis with a firm, nontender chancre.

Atypical primary lesions are common. The clinical appearance depends on the number of treponemes inoculated and on the immunologic status of the patient. A large inoculum produces a dark-field-positive ulcerative lesion in nonimmune volunteers but may produce a small dark-field-negative papule, an asymptomatic but seropositive latent infection, or no response at all in some individuals with a history of syphilis. A small inoculum may produce only a papular lesion, even in nonimmune individuals. Therefore, syphilis should be considered even in the evaluation of trivial or atypical dark-field-negative genital lesions. The genital lesions that most commonly must be differentiated from those of primary syphilis include those caused by herpes simplex virus infection (Chap. 216), chancroid (Chap. 182), traumatic injury, and donovanosis (Chap. 198e). Regional (usually inguinal) lymphadenopathy accompanies the primary syphilitic lesion, appearing within 1 week of lesion onset. The nodes are firm, nonsuppurative, and painless. Inguinal lymphadenopathy is bilateral and may occur with anal as well as with external genital chancres. The chancre generally heals within 4–6 weeks (range, 2–12 weeks), but lymphadenopathy may persist for months.

Secondary Syphilis The protean manifestations of the secondary stage usually include mucocutaneous lesions and generalized nontender lymphadenopathy. The healing primary chancre may still be present in ~15% of cases, and the stages may overlap more frequently in persons with concurrent HIV infection. The skin rash consists of macular, papular, papulosquamous, and occasionally pustular syphilides; often more than one form is present simultaneously. The eruption may be very subtle, and 25% of patients with a discernible rash may be unaware that they have dermatologic manifestations. Initial lesions are pale red or pink, nonpruritic, discrete macules distributed on the trunk and proximal extremities; these macules progress to papular lesions that are distributed widely and that frequently involve the palms and soles (Fig. 206-3; see also Figs. 25e-18 and 25e-19). Rarely, severe necrotic lesions (*lues maligna*) may appear; they are more commonly reported in HIV-infected individuals. Involvement of the hair follicles may result in patchy alopecia of the scalp hair, eyebrows, or beard in up to 5% of cases.

In warm, moist, intertriginous areas (commonly the perianal region, vulva, and scrotum), papules can enlarge to produce broad, moist, pink or gray-white, highly infectious lesions (*condylomata lata*; see Fig. 25e-20) in 10% of patients with secondary syphilis. Superficial mucosal erosions (*mucous patches*) occur in 10–15% of patients and commonly involve the oral or genital mucosa (see Fig. 25e-21). The typical mucous patch is a painless silver-gray erosion surrounded by a red periphery.

Constitutional signs and symptoms that may accompany or precede secondary syphilis include sore throat (15–30%), fever (5–8%), weight loss (2–20%), malaise (25%), anorexia (2–10%), headache (10%), and meningismus (5%). *Acute meningitis* occurs in only 1–2% of cases, but CSF cell and protein concentrations are increased in up to 40% of cases, and viable *T. pallidum* organisms have been recovered from CSF during

primary and secondary syphilis in 30% of cases; the latter finding is often but not always associated with other CSF abnormalities.

Less common complications of secondary syphilis include hepatitis, nephropathy, gastrointestinal involvement (hypertrophic gastritis, patchy proctitis, or a rectosigmoid mass), arthritis, and periostitis. Ocular findings associated with secondary syphilis include pupillary abnormalities and optic neuritis as well as the classic iritis or uveitis. The diagnosis of ocular syphilis is often considered in affected patients only after they fail to respond to steroid therapy. Anterior uveitis has been reported in 5–10% of patients with secondary syphilis, and *T. pallidum* has been demonstrated in aqueous humor from such patients. Hepatic involvement is common in syphilis; although it is usually asymptomatic, up to 25% of patients may have abnormal liver function tests. Frank syphilitic hepatitis may be seen. Renal involvement usually results from immune complex deposition and produces proteinuria associated with an acute nephrotic syndrome. Like those of primary syphilis, the manifestations of the secondary stage resolve spontaneously, usually within 1–6 months.

Latent Syphilis Positive serologic tests for syphilis, together with a normal CSF examination and the absence of clinical manifestations of syphilis, indicate a diagnosis of latent syphilis in an untreated person. The diagnosis is often suspected on the basis of a history of primary or secondary lesions, a history of exposure to syphilis, or the delivery of an infant with congenital syphilis. A previous negative serologic test or a history of lesions or exposure may help establish the duration of latent infection, which is an important factor in the selection of appropriate therapy. *Early latent* syphilis is limited to the first year after infection, whereas *late latent* syphilis is defined as that of ≥ 1 year's duration (or of unknown duration). *T. pallidum* may still seed the bloodstream intermittently during the latent stage, and pregnant women with latent syphilis may infect the fetus in utero. Moreover, syphilis has been transmitted through blood transfusion or organ donation from patients with latent syphilis. It was previously thought that untreated late latent syphilis had three possible outcomes: (1) persistent lifelong infection; (2) development of late syphilis; or (3) spontaneous cure, with reversion of serologic tests to negative. It is now apparent, however, that the more sensitive treponemal antibody tests rarely, if ever, become nonreactive without treatment. Although progression to clinically evident late syphilis is very rare today, the occurrence of spontaneous cure is in doubt.

Involvement of the CNS Traditionally, neurosyphilis has been considered a late manifestation of syphilis, but this view is inaccurate. CNS syphilis represents a continuum encompassing early invasion (usually within the first weeks or months of infection), months to years of asymptomatic involvement, and, in some cases, development of early or late neurologic manifestations.

ASYMPTOMATIC NEUROSYPHILIS The diagnosis of asymptomatic neurosyphilis is made in patients who lack neurologic symptoms and signs but who have CSF abnormalities including mononuclear pleocytosis, increased protein concentrations, or CSF reactivity in the Venereal Disease Research Laboratory (VDRL) test. CSF abnormalities are demonstrated in up to 40% of cases of primary or secondary syphilis and in 25% of cases of latent syphilis. *T. pallidum* has been recovered by inoculation into rabbits of CSF from up to 30% of patients with primary or secondary syphilis but less frequently by inoculation of CSF from patients with latent syphilis. The presence of *T. pallidum* in CSF is often associated with other CSF abnormalities, but organisms can be recovered from patients with otherwise normal CSF. Although the prognostic implications of these findings in early syphilis are uncertain, it may be appropriate to conclude that even patients with



FIGURE 206-3 Secondary syphilis. **Left:** Maculopapular truncal eruption. **Middle:** Papules on the palms. **Right:** Papules on the soles. (Courtesy of Jill McKenzie and Christina Marra.)

1136 early syphilis who have such findings do indeed have asymptomatic neurosyphilis and should be treated for neurosyphilis; such treatment is particularly important in patients with concurrent HIV infection. Before the advent of penicillin, the risk of development of clinical neurosyphilis in untreated asymptomatic persons was roughly proportional to the intensity of CSF changes, with the overall cumulative probability of progression to clinical neurosyphilis ~20% in the first 10 years but increasing with time. Most experts agree that neurosyphilis is more common in HIV-infected persons, while immunocompetent patients with untreated latent syphilis and normal CSF probably run a very low risk of subsequent neurosyphilis. In several recent studies, neurosyphilis was associated with a rapid plasma reagin (RPR) titer of $\geq 1:32$, regardless of clinical stage or HIV infection status.

SYMPTOMATIC NEUROSYPHILIS The major clinical categories of symptomatic neurosyphilis include meningeal, meningovascular, and parenchymatous syphilis. The last category includes general paresis and tabes dorsalis. The onset of symptoms usually occurs <1 year after infection for meningeal syphilis, up to 10 years after infection for meningovascular syphilis, at ~20 years for general paresis, and at 25–30 years for tabes dorsalis. Neurosyphilis is more frequently symptomatic in patients who are co-infected with HIV, particularly in the setting of a low CD4+ T lymphocyte count. In addition, recent evidence suggests that syphilitic infection worsens the cognitive impairment seen in HIV-infected persons and that this effect persists even after treatment for syphilis.

Meningeal syphilis may present as headache, nausea, vomiting, neck stiffness, cranial nerve involvement, seizures, and changes in mental status. This condition may be concurrent with or may follow the secondary stage. Patients presenting with uveitis, iritis, or hearing loss often have meningeal syphilis, but these clinical findings can also be seen in patients with normal CSF.

Meningovascular syphilis reflects meningitis together with inflammatory vasculitis of small, medium, or large vessels. The most common presentation is a stroke syndrome involving the middle cerebral artery of a relatively young adult. However, unlike the usual thrombotic or embolic stroke syndrome of sudden onset, meningovascular syphilis often becomes manifest after a subacute encephalitic prodrome (with headaches, vertigo, insomnia, and psychological abnormalities), which is followed by a gradually progressive vascular syndrome.

The manifestations of *general paresis* reflect widespread late parenchymal damage and include abnormalities corresponding to the mnemonic *paresis*: personality, affect, reflexes (hyperactive), eye (e.g., Argyll Robertson pupils), sensorium (illusions, delusions, hallucinations), intellect (a decrease in recent memory and in the capacity for orientation, calculations, judgment, and insight), and speech. *Tabes dorsalis* is a late manifestation of syphilis that presents as symptoms and signs of demyelination of the posterior columns, dorsal roots, and dorsal root ganglia. Symptoms include ataxic wide-based gait and foot drop; paresthesia; bladder disturbances; impotence; areflexia; and loss of positional, deep-pain, and temperature sensations. Trophic joint degeneration (Charcot's joints) and perforating ulceration of the feet can result from loss of pain sensation. The small, irregular Argyll Robertson pupil, a feature of both tabes dorsalis and paresis, reacts to accommodation but not to light. *Optic atrophy* also occurs frequently in association with tabes.


Other Manifestations of Late Syphilis The slowly progressive inflammatory process leading to tertiary disease begins early during infection, although these manifestations may not become clinically apparent for years or decades. Early syphilitic aortitis becomes evident soon after secondary lesions subside, and treponemes that trigger the development of gummas may have seeded the tissue years earlier.

CARDIOVASCULAR SYPHILIS Cardiovascular manifestations, usually appearing 10–40 years after infection, are attributable to endarteritis obliterans of the vasa vasorum, which provide the blood supply to large vessels; *T. pallidum* DNA has been detected by PCR in aortic tissue. Cardiovascular involvement results in uncomplicated aortitis, aortic regurgitation, saccular aneurysm (usually of the ascending aorta), or coronary ostial stenosis. In the preantibiotic era, symptomatic

cardiovascular complications developed in ~10% of persons with late untreated syphilis. Today, this form of late syphilis is rarely seen in the developed world. Linear calcification of the ascending aorta on chest x-ray films suggests asymptomatic syphilitic aortitis, as arteriosclerosis seldom produces this sign. Only 1 in 10 aortic aneurysms of syphilitic origin involves the abdominal aorta.

LATE BENIGN SYPHILIS (GUMMA) Gummas are usually solitary lesions ranging from microscopic to several centimeters in diameter. Histologic examination shows a granulomatous inflammation, with a central area of necrosis due to endarteritis obliterans. Although rarely demonstrated microscopically, *T. pallidum* has been detected by PCR or recovered from these lesions, and penicillin treatment results in rapid resolution, confirming the treponemal stimulus for the inflammation. Common sites include the skin and skeletal system; however, any organ (including the brain) may be involved. Gummas of the skin produce indolent, painless, indurated nodular or ulcerative lesions that may resemble other chronic granulomatous conditions, including tuberculosis, sarcoidosis, leprosy, and deep fungal infections. Skeletal gummas most frequently involve the long bones, although any bone may be affected. Upper respiratory gummas can lead to perforation of the nasal septum or palate.

Congenital Syphilis Transmission of *T. pallidum* across the placenta from a syphilitic woman to her fetus may occur at any stage of pregnancy, but fetal damage generally does not occur until after the fourth month of gestation, when fetal immunologic competence begins to develop. This timing suggests that the pathogenesis of congenital syphilis, like that of adult syphilis, depends on the host immune response rather than on a direct toxic effect of *T. pallidum*. The risk of fetal infection during untreated early maternal syphilis is ~75–95%, decreasing to ~35% for maternal syphilis of >2 years' duration. Adequate treatment of the woman before the 16th week of pregnancy should prevent fetal damage, and treatment before the third trimester should adequately treat the infected fetus. Untreated maternal infection may result in a rate of fetal loss of up to 40% (with stillbirth more common than abortion because of the late onset of fetal pathology), prematurity, neonatal death, or nonfatal congenital syphilis. Among infants born alive, only fulminant congenital syphilis is clinically apparent at birth, and these babies have a very poor prognosis. The most common clinical problem is the healthy-appearing baby born to a mother with a positive serologic test

 Routine serologic testing in early pregnancy is considered cost-effective in virtually all populations, even in areas with a low prenatal prevalence of syphilis. Low-tech point-of-care tests have been developed and are being widely implemented to facilitate antenatal testing in resource-poor settings. A recent study demonstrated the high cost-effectiveness of using these tests for screening (with subsequent treatment) in sub-Saharan Africa. Adverse outcomes were reduced, with 64,000 fewer stillbirths, 25,000 fewer neonatal deaths, and up to 25,000 fewer live births of infants with syphilis. The intervention would remain cost-effective even if the current syphilis seroprevalence among pregnant women declined from its present 3.1% to 0.4%. Where the prevalence of syphilis is high or when the patient is at high risk of reinfection, serologic testing should be repeated in the third trimester and at delivery. Neonatal congenital syphilis must be differentiated from other generalized congenital infections, including rubella, cytomegalovirus or herpes simplex virus infection, and toxoplasmosis, as well as from erythroblastosis fetalis.

The manifestations of congenital syphilis include (1) early manifestations, which appear within the first 2 years of life (often at 2–10 weeks of age), are infectious, and resemble the manifestations of secondary syphilis in the adult; (2) late manifestations, which appear after 2 years and are noninfectious; and (3) residual stigmata. The earliest manifestations of congenital syphilis include rhinitis, or “snuffles” (23%); mucocutaneous lesions (35–41%); bone changes (61%), including osteochondritis, osteitis, and periostitis detectable by x-ray examination of long bones; hepatosplenomegaly (50%); lymphadenopathy (32%); anemia (34%); jaundice (30%); thrombocytopenia; and leukocytosis. CNS invasion by *T. pallidum* is detectable in 22% of infected

neonates. Neonatal death is usually due to pulmonary hemorrhage, secondary bacterial infection, or severe hepatitis.

Late congenital syphilis (untreated after 2 years of age) is subclinical in 60% of cases; the clinical spectrum in the remainder of cases may include interstitial keratitis (which occurs at 5–25 years of age), eighth-nerve deafness, and recurrent arthropathy. Bilateral knee effusions are known as *Clutton's joints*. Neurosyphilis was present in about one-quarter of untreated patients with late congenital syphilis in the preantibiotic era. Gummatous periostitis occurs at 5–20 years of age and, as in nonvenereal endemic syphilis, tends to cause destructive lesions of the palate and nasal septum.

Classic stigmata include *Hutchinson's teeth* (centrally notched, widely spaced, peg-shaped upper central incisors), “mulberry” molars (sixth-year molars with multiple, poorly developed cusps), saddle nose, and saber shins.

LABORATORY EXAMINATIONS

Demonstration of the Organism *T. pallidum* cannot be detected by culture. Historically, dark-field microscopy and immunofluorescence antibody staining have been used to identify this spirochete in samples from moist lesions such as chancres or condylomata lata, but these tests are rarely available outside of research laboratories. Sensitive and specific PCR tests have been developed but are not commercially available, although a number of laboratories perform in-house validated PCR testing.

T. pallidum can be found in tissue with appropriate silver stains, but these results should be interpreted with caution because artifacts resembling *T. pallidum* are often seen. Tissue treponemes can be demonstrated more reliably in research laboratories by PCR or by immunofluorescence or immunohistochemical methods using specific monoclonal or polyclonal antibodies to *T. pallidum*.

Serologic Tests for Syphilis There are two types of serologic test for syphilis: nontreponemal and treponemal. Both are reactive in persons with any treponemal infection, including yaws, pinta, and endemic syphilis.

The most widely used nontreponemal antibody tests for syphilis are the RPR and VDRL tests, which measure IgG and IgM directed against a cardiolipin-lecithin-cholesterol antigen complex. The RPR test is easier to perform and uses unheated serum or plasma; it is the test of choice for rapid serologic diagnosis in a clinical setting. The VDRL test remains the standard for examining CSF and is superior to the RPR for this purpose. The RPR and VDRL tests are recommended for screening or for quantitation of serum antibody. The titer reflects disease activity, rising during the evolution of early syphilis, often exceeding 1:32 in secondary syphilis, and declining thereafter without therapy. After treatment for early syphilis, a persistent fall by fourfold or more (e.g., a decline from 1:32 to 1:8) is considered an adequate response. VDRL titers do not correspond directly to RPR titers, and sequential quantitative testing (as for response to therapy) must employ a single test. As will be discussed (see “Evaluation for Neurosyphilis,” below), the RPR titer may be useful in determining which patients will benefit from CSF examination.

Treponemal tests measure antibodies to native or recombinant *T. pallidum* antigens and include the fluorescent treponemal antibody–absorbed (FTA–ABS) test and the *T. pallidum* particle agglutination (TPPA) test, both of which are more sensitive for primary syphilis than the previously used hemagglutination tests. The *T. pallidum* hemagglutination (TPHA) test is widely used in Europe but is not available in the United States. When used to confirm positive nontreponemal test results, treponemal tests have a very high positive predictive value for diagnosis of syphilis. Treponemal enzyme or chemiluminescence immunoassays (EIAs/CIAs), based largely on reactivity to recombinant antigens, have also been developed and are now widely used as screening tests by large laboratories. In a screening setting, however, treponemal tests give false-positive results at rates as high as 1–2%, and the rate is higher with the EIA/CIA tests. Treponemal tests are likely to remain reactive even after adequate treatment and cannot differentiate

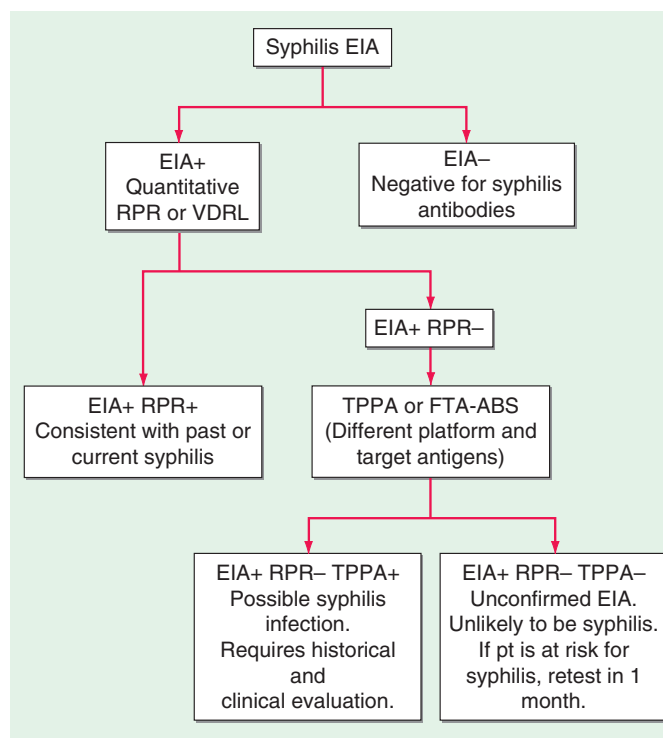


FIGURE 206-4 Algorithm for interpretation of results from syphilis enzyme immunoassays (EIAs) used for screening. FTA–ABS, fluorescent treponemal antibody–absorbed; RPR, rapid plasma reagin; TPPA, *Treponema pallidum* particle agglutination; VDRL, Venereal Disease Research Laboratory. (Based on the 2010 Sexually Transmitted Diseases Treatment Guidelines from the Centers for Disease Control and Prevention.)

past from current *T. pallidum* infection. **Figure 206-4** provides a suggested algorithm for management of such cases.

Both nontreponemal and treponemal tests may be nonreactive in early primary syphilis, although treponemal tests are slightly more sensitive (85–90%) during this stage than nontreponemal tests (~80%). All tests are reactive during secondary syphilis. (Fewer than 1% of patients with high titers have a nontreponemal test that is nonreactive or weakly reactive with undiluted serum but is reactive with diluted serum—the *prozone phenomenon*.) VDRL and RPR sensitivity and titers may decline in untreated persons with late latent syphilis, but treponemal tests remain sensitive in these stages. After treatment for early syphilis, nontreponemal test titers will generally decline or the tests will become nonreactive, whereas treponemal tests often remain reactive after therapy and are not helpful in determining the infection status of persons with past syphilis.

For practical purposes, most clinicians need to be familiar with three uses of serologic tests for syphilis recommended by the Centers for Disease Control and Prevention (CDC): (1) screening or diagnosis (RPR or VDRL), (2) quantitative measurement of antibody to assess clinical syphilis activity or to monitor response to therapy (RPR or VDRL), and (3) confirmation of a syphilis diagnosis in a patient with a reactive RPR or VDRL test (FTA–ABS, TPPA, EIA/CIA). Studies have not demonstrated the utility of IgM testing for adult syphilis. Whereas IgM titers appear to decline after therapy, the presence or absence of specific IgM does not strictly correlate with *T. pallidum* infection. Moreover, no commercially available IgM test is recommended, even for evaluation of infants with suspected congenital syphilis.

False-Positive Serologic Tests for Syphilis The lipid antigens of nontreponemal tests are similar to those found in human tissues, and the tests may be reactive (usually with titers $\leq 1:8$) in persons without treponemal infection. Among patients being screened for syphilis because of risk factors, clinical suspicion, or history of exposure, ~1% of reactive tests are falsely positive. Modern VDRL and RPR tests are highly specific, and false-positive reactions are largely limited to persons with

1138 autoimmune conditions or injection drug use. The prevalence of false-positive results increases with advancing age, approaching 10% among persons >70 years old. In a patient with a false-positive nontreponemal test, syphilis is excluded by a nonreactive treponemal test.

False-positive reactions may also occur with the treponemal tests, particularly the new, very sensitive EIA/CIA tests. When a low-prevalence population for syphilis is screened, the number of false-positive reactions may outnumber true positives, leading to unnecessary treatment. Although the precise reason is not known, it has been shown that sera from patients with periodontal disease react with antigens used in the EIA/CIA tests, presumably as a result of cross-reactive epitopes in the many treponemes that infect the gingival crevices during periodontal disease.

Evaluation for Neurosyphilis Involvement of the CNS is detected by examination of CSF for pleocytosis (>5 white blood cells/ μ L), increased protein concentration (>45 mg/dL), or VDRL reactivity. Elevated CSF cell counts and protein concentrations are not specific for neurosyphilis and may be confounded by HIV co-infection. Because CSF pleocytosis may also be due to HIV, some studies have suggested using a CSF white-cell cutoff of 20 cells/ μ L as diagnostic of neurosyphilis in HIV-infected patients with syphilis. The CSF VDRL test is highly specific and, when reactive, is considered diagnostic of neurosyphilis; however, this test is insensitive and may be nonreactive even in cases of symptomatic neurosyphilis. The FTA-ABS test on CSF is reactive far more often than the VDRL test on CSF in all stages of syphilis, but reactivity may reflect passive transfer of serum antibody into the CSF. A nonreactive FTA-ABS test on CSF, however, may be used to rule out asymptomatic neurosyphilis. The utility of measuring CXCL13 in CSF to distinguish between neurosyphilis and HIV-related CSF abnormalities has been demonstrated.

Clearly, all *T. pallidum*-infected patients who have signs or symptoms consistent with neurologic disease (e.g., meningitis, hearing loss) or ophthalmic disease (e.g., uveitis, iritis) should have a CSF examination, regardless of disease stage. The appropriate management of asymptomatic persons is less clear. Lumbar puncture on all asymptomatic patients with untreated syphilis is impractical and unnecessary. Because standard therapy with penicillin G benzathine fails to result in treponemical drug levels in CSF, however, it is important to identify those persons at higher risk for having or developing neurosyphilis so that appropriate therapy may be given. Viable *T. pallidum* has been isolated from the CSF of several patients (with and without concurrent HIV infection) after penicillin G benzathine therapy for early syphilis. Large-scale prospective studies have now provided evidence-based guidelines for determining which syphilis patients may benefit most from CSF examination for evidence of neurosyphilis. Specifically, patients with RPR titers of $\geq 1:32$ are at higher risk of having neurosyphilis (11-fold and 6-fold higher in HIV-infected and HIV-uninfected persons, respectively), as are HIV-infected patients with CD4+ T cell counts of $\leq 350/\mu$ L. Guidelines for CSF examination are shown in [Table 206-1](#).


Evaluation of HIV-Infected Patients for Syphilis Because persons at highest risk for syphilis are also at increased risk for HIV infection, these

two infections frequently coexist. There is evidence that syphilis and other genital ulcer diseases are important risk factors for acquisition and transmission of HIV infection. Some manifestations of syphilis may be altered in patients with concurrent HIV infection, and multiple cases of neurologic relapse after standard therapy have been reported in these patients.

Persons with newly diagnosed HIV infection should be tested for syphilis; conversely, all patients with newly diagnosed syphilis should be tested for HIV infection. Some authorities, persuaded by reports of persistent *T. pallidum* in CSF of HIV-infected persons after standard therapy for early syphilis, recommend CSF examination for evidence of neurosyphilis for all co-infected patients, regardless of the stage of syphilis, with treatment for neurosyphilis if CSF abnormalities are found. Others, on the basis of their own clinical experience, believe that standard therapy—without CSF examination—is sufficient for all cases of early syphilis in HIV-infected patients without neurologic signs or symptoms. As described above, RPR titer and CD4+ T cell count can be used to identify patients at higher risk of neurosyphilis for lumbar puncture, although some cases of neurosyphilis will be missed, even when these criteria are used. [Table 206-1](#) summarizes guidelines suggested by published studies. Serologic testing after treatment is important for all patients with syphilis, particularly for those also infected with HIV.

TREATMENT SYPHILIS

TREATMENT OF ACQUIRED SYPHILIS

 The CDC's 2010 guidelines for the treatment of syphilis are summarized in [Table 206-2](#) and are discussed below. Penicillin G is the drug of choice for all stages of syphilis. *T. pallidum* is killed by very low concentrations of penicillin G, although a long period of exposure to penicillin is required because of the unusually slow rate of multiplication of the organism. The efficacy of penicillin against syphilis remains undiminished after 60 years of use, and there is no evidence of penicillin resistance in *T. pallidum*. Other antibiotics effective in syphilis include the tetracyclines and the cephalosporins. Aminoglycosides and spectinomycin inhibit *T. pallidum* only in very large doses, and the sulfonamides and the quinolones are inactive. Azithromycin has shown significant promise as an effective oral agent against *T. pallidum*; however, strains harboring 23S rRNA mutations that confer macrolide resistance are widespread; such strains represent >80% of recent isolates from Seattle and San Francisco and have now been identified in multiple North American and European sites. Macrolide resistance mutations have been identified in nearly all samples reported from some regions of China. In contrast, a study based in Madagascar documented the equivalence of benzathine penicillin and azithromycin for treatment of early syphilis, although a sample from one azithromycin clinical failure in that study showed the presence of a 23S rRNA resistance mutation. A more recent survey from South Africa showed a very low (1%) frequency of known 23s rRNA resistance mutations. In short, the prevalence of resistant strains varies widely by geographic location, and routine treatment of syphilis with azithromycin is not recommended. In all cases, careful follow-up of any patient treated for syphilis with azithromycin must be ensured.

Early Syphilis Patients and Their Contacts Penicillin G benzathine is the most widely used agent for the treatment of early syphilis; a single dose of 2.4 million units is recommended. Preventive treatment is also recommended for individuals who have been exposed to infectious syphilis within the previous 3 months. *The regimens recommended for prevention are the same as those recommended for early syphilis.* Penicillin G benzathine cures >95% of cases of early syphilis, although clinical relapse can follow treatment, particularly in patients with concurrent HIV infection. Because the risk of neurologic relapse may be higher in HIV-infected patients, CSF examination is recommended in HIV-seropositive individuals with syphilis of any stage, particularly those with a serum RPR titer of $\geq 1:32$ or a

TABLE 206-1 INDICATIONS FOR CEREBROSPINAL FLUID EXAMINATION IN ADULTS WITH ALL STAGES OF SYPHILIS

All Patients

Signs or symptoms of nervous system involvement (e.g., meningitis, hearing loss, cranial nerve dysfunction, altered mental status, ophthalmic disease [e.g., uveitis, iritis, pupillary abnormalities], ataxia, loss of vibration sense), or RPR or VDRL titer $\geq 1:32$, or Active tertiary syphilis, or Suspected treatment failure

Additional Indications in HIV-Infected Persons

CD4+ T cell count $\leq 350/\mu$ L, or All HIV-infected persons (recommended by some experts)

Source: Adapted from the 2010 Sexually Transmitted Diseases Treatment Guidelines from the Centers for Disease Control and Prevention.

TABLE 206-2 RECOMMENDATIONS FOR THE TREATMENT OF SYPHILIS^a

Stage of Syphilis	Patients without Penicillin Allergy	Patients with Confirmed Penicillin Allergy ^b
Primary, secondary, or early latent	CSF normal or not examined: Penicillin G benzathine (single dose of 2.4 mU IM) CSF abnormal: Treat as neurosyphilis	CSF normal or not examined: Tetracycline HCl (500 mg PO qid) or doxycycline (100 mg PO bid) for 2 weeks CSF abnormal: Treat as neurosyphilis
Late latent (or latent of uncertain duration), cardiovascular, or benign tertiary	CSF normal or not examined: Penicillin G benzathine (2.4 mU IM weekly for 3 weeks) CSF abnormal: Treat as neurosyphilis	CSF normal and patient not infected with HIV: Tetracycline HCl (500 mg PO qid) or doxycycline (100 mg PO bid) for 4 weeks CSF normal and patient infected with HIV: Desensitization and treatment with penicillin if compliance cannot be ensured CSF abnormal: Treat as neurosyphilis
Neurosyphilis (asymptomatic or symptomatic)	Aqueous crystalline penicillin G (18–24 mU/d IV, given as 3–4 mU q4h or continuous infusion) for 10–14 days or Aqueous procaine penicillin G (2.4 mU/d IM) plus oral probenecid (500 mg qid), both for 10–14 days	Desensitization and treatment with penicillin ^c
Syphilis in pregnancy	According to stage	Desensitization and treatment with penicillin

^aSee Table 206-1 and text for indications for CSF examination. ^bBecause of the documented presence of macrolide resistance in many *T. pallidum* strains in North America, Europe, and China, azithromycin or other macrolides should be used with caution only when treatment with penicillin or doxycycline is not feasible. Azithromycin should not be used for men who have sex with men or for pregnant women. ^cLimited data suggest that ceftriaxone (2 g/d either IM or IV for 10–14 days) can be used; however, cross-reactivity between penicillin and ceftriaxone is possible.

Abbreviations: CSF, cerebrospinal fluid; mU, million units.

Source: Adapted from the 2010 Sexually Transmitted Diseases Treatment Guidelines from the Centers for Disease Control and Prevention.

CD4+ T cell count of $\leq 350/\mu\text{L}$. Therapy appropriate for neurosyphilis should be given if there is any evidence of CNS infection.

Late Latent Syphilis or Syphilis of Unknown Duration If the CSF is normal or is not examined, the recommended treatment is penicillin G benzathine (7.2 million units total; Table 206-2). If CSF abnormalities are found, the patient should be treated for neurosyphilis.

Tertiary Syphilis CSF examination should be performed. If the CSF is normal, the recommended treatment is penicillin G benzathine (7.2 million units total; Table 206-2). If CSF abnormalities are found, the patient should be treated for neurosyphilis. The clinical response to treatment for benign tertiary syphilis is usually impressive. However, responses to therapy for cardiovascular syphilis are not dramatic because aortic aneurysm and aortic regurgitation cannot be reversed by antibiotics.

Syphilis in Penicillin-Allergic Patients For penicillin-allergic patients with syphilis, a 2-week (early syphilis) or 4-week (late or late latent syphilis) course of therapy with doxycycline or tetracycline is recommended (Table 206-2). These regimens appear to be effective in early syphilis but have not been tested for late or late latent syphilis, and compliance may be problematic. Limited studies suggest that ceftriaxone (1 g/d, given IM or IV for 8–10 days) is effective for early syphilis. These nonpenicillin regimens have not been carefully evaluated in HIV-infected individuals and should be used with caution. If compliance and follow-up cannot be ensured, penicillin-allergic HIV-infected persons with late latent or late syphilis should be desensitized and treated with penicillin.

Neurosyphilis Penicillin G benzathine, given in total doses of up to 7.2 million units, does not produce detectable concentrations of penicillin G in CSF and should not be used for treatment of neurosyphilis. Asymptomatic neurosyphilis may relapse as symptomatic disease after treatment with benzathine penicillin, and the risk of relapse may be higher in HIV-infected patients. Both symptomatic and asymptomatic neurosyphilis should be treated with aqueous penicillin (Table 206-2). Administration either of IV aqueous crystalline penicillin G or of IM aqueous procaine penicillin G plus oral probenecid in recommended doses is thought to ensure treponemidal concentrations of penicillin G in CSF. The clinical response to penicillin therapy for meningeal syphilis is dramatic, but treatment of neurosyphilis with existing parenchymal damage may only arrest disease progression. No data suggest

that additional therapy (e.g., penicillin G benzathine for 3 weeks) is beneficial after treatment for neurosyphilis.

The use of antibiotics other than penicillin G for the treatment of neurosyphilis has not been studied, although very limited data suggest that ceftriaxone may be used. In patients with penicillin allergy demonstrated by skin testing, desensitization and treatment with penicillin are recommended.

Management of Syphilis in Pregnancy Every pregnant woman should undergo a nontreponemal test at her first prenatal visit and, if at high risk of exposure, again in the third trimester and at delivery. In the untreated pregnant patient with presumed syphilis, expeditious treatment appropriate to the stage of the disease is essential. Patients should be warned of the risk of a Jarisch-Herxheimer reaction, which may be associated with mild premature contractions but rarely results in premature delivery.

Penicillin is the only recommended agent for the treatment of syphilis in pregnancy. If the patient has a documented penicillin allergy, desensitization and penicillin therapy should be undertaken according to the CDC's 2010 guidelines. After treatment, a quantitative nontreponemal test should be repeated monthly throughout pregnancy to assess therapeutic efficacy. Treated women whose antibody titers rise by fourfold or whose titers do not decrease by fourfold over a 3-month period should be re-treated.

EVALUATION AND MANAGEMENT OF CONGENITAL SYPHILIS

Whether or not they are infected, newborn infants of mothers with reactive serologic tests may themselves have reactive tests because of transplacental transfer of maternal IgG antibody. For asymptomatic infants born to women treated adequately with penicillin during the first or second trimester of pregnancy, monthly quantitative nontreponemal tests may be performed to monitor for appropriate reduction in antibody titers. Rising or persistent titers indicate infection, and the infant should be treated. Detection of neonatal IgM antibody may be useful, but no commercially available test is currently recommended.

An infant should be treated at birth if the treatment status of the seropositive mother is unknown; if the mother has received inadequate or nonpenicillin therapy; if the mother received penicillin therapy in the third trimester; or if the infant may be difficult to follow. The CSF should be examined to obtain baseline values before treatment. Penicillin is the only recommended drug for the treatment of syphilis in infants. Specific recommendations for the treatment of infants and older children are included in the CDC's 2010 treatment guidelines.

1140 JARISCH-HERXHEIMER REACTION

A dramatic although usually mild reaction consisting of fever, chills, myalgias, headache, tachycardia, increased respiratory rate, increased circulating neutrophil count, and vasodilation with mild hypotension may follow the initiation of treatment for syphilis. This reaction is thought to be a response to lipoproteins released by dying *T. pallidum* organisms. The Jarisch-Herxheimer reaction occurs in 50% of patients with primary syphilis, 90% of those with secondary syphilis, and a lower proportion of persons with later-stage disease. Defervescence takes place within 12–24 h. In patients with secondary syphilis, erythema and edema of the mucocutaneous lesions may increase. Patients should be warned to expect such symptoms, which can be managed with symptom-based treatment. Steroid or other anti-inflammatory therapy is not required for this mild transient reaction.

FOLLOW-UP EVALUATION OF RESPONSES TO THERAPY

Efficacy of treatment should be assessed by clinical evaluation and monitoring of the quantitative VDRL or RPR titer for a fourfold decline (e.g., from 1:32 to 1:8). Patients with primary or secondary syphilis should be examined 6 and 12 months after treatment and persons with latent or late syphilis at 6, 12, and 24 months. More frequent clinical and serologic examination (3, 6, 9, 12, and 24 months) is recommended for patients concurrently infected with HIV, regardless of the stage of syphilis.

After successful treatment of seropositive first-episode primary or secondary syphilis, the VDRL or RPR titer progressively declines, becoming negative by 12 months in 40–75% of seropositive primary cases and in 20–40% of secondary cases. Patients with HIV infection or a history of prior syphilis are less likely to become nonreactive in the VDRL or RPR test. Rates of decline of serologic titers appear to be slower, and serologically defined treatment failures more common, among HIV-infected patients than among those without HIV co-infection; however, effective antiretroviral therapy may reduce these differences. Re-treatment should be considered if serologic responses are not adequate or if clinical signs persist or recur. Because it is difficult to differentiate treatment failure from reinfection, the CSF should be examined, with treatment for neurosyphilis if CSF is abnormal and treatment for late latent syphilis if CSF is normal. A minority of patients treated for early syphilis may experience a one-dilution titer increase within 14 days after treatment; however, this early elevation does not significantly affect the serologic outcome at 6 months after treatment. Patients treated for late latent syphilis frequently have low initial VDRL or RPR titers and may not have a fourfold decline after therapy with penicillin. In such patients, re-treatment is not warranted unless the titer rises or signs and symptoms of syphilis appear. Because treponemal tests may remain reactive despite treatment for seropositive syphilis, these tests are not useful in following the response to therapy.

The activity of neurosyphilis (symptomatic or asymptomatic) correlates best with CSF pleocytosis, and this measure provides the most sensitive index of response to treatment. Repeat CSF examinations should be performed every 6 months until the cell count is normal. An elevated CSF cell count falls to normal in 3–12 months in adequately treated HIV-uninfected patients. The persistence of mild pleocytosis in HIV-infected patients may be due to the presence of HIV in CSF; this scenario may be difficult to distinguish from treatment failure. Elevated levels of CSF protein fall more slowly, and the CSF VDRL titer declines gradually over several years. In patients treated for neurosyphilis, a fourfold reduction in serum RPR titer has been positively correlated with normalization of CSF abnormalities; this correlation is stronger in HIV-uninfected patients and in HIV-infected patients receiving effective antiretroviral therapy.

IMMUNITY TO SYPHILIS

The rate of development of acquired resistance to *T. pallidum* after natural or experimental infection is related to the size of the antigenic stimulus, which depends on both the size of the infecting inoculum

and the duration of infection before treatment. Both humoral and cellular responses are considered to be of major importance in immunity and in the healing of early lesions. Cellular infiltration, predominantly by T lymphocytes and macrophages, produces a T_H1 cytokine milieu consistent with the clearance of organisms by activated macrophages. Specific antibody enhances phagocytosis and is required for macrophage-mediated killing of *T. pallidum*. Recent studies demonstrate antigenic variation of the TprK protein, which may lead to persistence of infection and determine susceptibility to reinfection with another strain. Comparative genomic studies have revealed some sequence variations among *T. pallidum* strains, which can be differentiated by molecular typing methods. Possible correlations between molecular type and clinical manifestations are being examined.

207e Endemic Treponematoses

Sheila A. Lukehart

The endemic treponematoses are chronic diseases that are transmitted by direct contact, usually during childhood, and, like syphilis, can cause severe late manifestations years after initial infection. These diseases are caused by very close relatives of *Treponema pallidum* subspecies *pallidum*, the etiologic agent of venereal syphilis (Chap. 206). Yaws, pinta, and endemic syphilis are traditionally distinguished from venereal syphilis by mode of transmission, age of acquisition, geographic distribution, and clinical features; however, there is some overlap for each of these factors. Generally, yaws flourishes in moist tropical areas of several regions, endemic syphilis is found primarily in arid climates, and pinta is found in temperate foci in the Americas (Fig. 207e-1). These infections are usually limited to rural areas of developing nations and are seen in developed countries only among recent immigrants from endemic regions. Our “knowledge” about the endemic treponematoses is based on observations by health care workers who have visited endemic areas; virtually no well-designed studies of the natural history, diagnosis, or treatment of these infections have been conducted. The treponemal infections are compared and contrasted in Table 207e-1.

EPIDEMIOLOGY



In a World Health Organization (WHO)-sponsored mass eradication campaign from 1952 to 1969, more than 160 million people in Africa, Asia, and South America were examined for treponemal infections, and more than 50 million cases, contacts, and persons with latent infections were treated. This campaign reduced the prevalence of active yaws from >20% to <1% in many areas. In recent decades, lack of focused surveillance and diversion of resources have resulted in documented resurgence of these infections in some regions. The most recent WHO global estimate (1995) suggested that there are 460,000 new cases per year (mostly yaws) and a prevalence of 2.5 million infected persons; during subsequent years, an increased incidence was documented in some countries. Recent areas of resurgent yaws morbidity include West Africa (Ivory Coast, Ghana, Togo, Benin), the Central African Republic, Nigeria, and rural Democratic Republic of the Congo. The prevalence of endemic syphilis is estimated to be >10% in some regions of northern Ghana, Mali, Niger, Burkina Faso, and Senegal. In Asia and the Pacific Islands, reports suggest active outbreaks of yaws in Indonesia, Papua New Guinea, the Solomon Islands, East Timor, Vanuatu, Laos, and Kampuchea. India actively renewed its focus on yaws control in 1996, achieved zero-case status in 2003, and declared elimination in 2006. In the Americas, foci of yaws

are thought to persist in Haiti and other Caribbean islands, Peru, Colombia, Ecuador, Brazil, Guyana, and Surinam, although recent data are lacking. Pinta is limited to Central America and northern South America, where it is found rarely and only in very remote villages. Evidence of yaws-like and venereal diseases, with treponemal seroreactivity, in wild gorillas and baboons in Africa has led to speculation that there may be an animal reservoir for yaws.

MICROBIOLOGY



The etiologic agents of the endemic treponematoses are listed in Table 207e-1. These little-studied organisms are morphologically identical to *T. pallidum* subspecies *pallidum* (the agent of venereal syphilis), and no definitive antigenic differences among them have been identified to date. A controversy has existed about whether the pathogenic treponemes are truly separate organisms, as genome sequencing indicates that yaws and syphilis treponemes are 99.8% identical. Three of the four organisms are classified as subspecies of *T. pallidum*; the fourth (*T. carateum*) remains a separate species simply because no organisms have been available for genetic studies. Based on analysis of the small number of strains currently available, molecular signatures—assessed by polymerase chain reaction (PCR) amplification of *tpr* genes and restriction digestion—have been identified that can differentiate the *T. pallidum* subspecies. Whether these genetic differences are related to distinct clinical characteristics of these diseases has not been determined. Full genome sequencing of an unclassified strain (Fribourg-Blanc) isolated from a baboon in 1966 shows a very high degree of homology with available strains of *T. pallidum* subspecies *pertenue*. This observation is consistent with an earlier report that the Fribourg-Blanc strain can cause experimental infection of humans. Molecular analyses of additional samples from affected baboons suggests that the nonhuman primate samples diverge from the evolutionary tree prior to the clade that contains the human isolates, but uncertainty remains about the importance of the nonhuman primate reservoir for human infection.

CLINICAL FEATURES

All of the treponemal infections, including syphilis, are chronic and are characterized by defined disease stages, with a localized primary lesion, disseminated secondary lesions, periods of latency, and possible late lesions. Primary and secondary stages are more frequently overlapping in yaws and endemic syphilis than in venereal syphilis, and the late manifestations of pinta are very mild relative to the destructive lesions of the other treponematoses. The current preference is to divide the clinical course of the endemic treponematoses into “early” and “late” stages.

The major clinical distinctions made between venereal syphilis and the nonvenereal infections are the apparent lack of congenital transmission and of central nervous system (CNS) involvement in the

nonvenereal infections. It is not known whether these distinctions are entirely accurate. Because of the high degree of genetic relatedness among the organisms, there is little biological reason to think that *T. pallidum* subspecies *endemicum* and *T. pallidum* subspecies *pertenue* would be unable to cross the blood-brain barrier or to invade the placenta. These organisms are like *T. pallidum* subspecies *pallidum* in that they obviously disseminate from the site of initial infection and can persist for decades. The lack of recognized congenital infection may be due to the fact that childhood infections often reach the latent stage (low bacterial load) before girls reach sexual maturity. Neurologic involvement may go unrecognized because of the lack of trained medical personnel in endemic regions, the delay of many years between infection

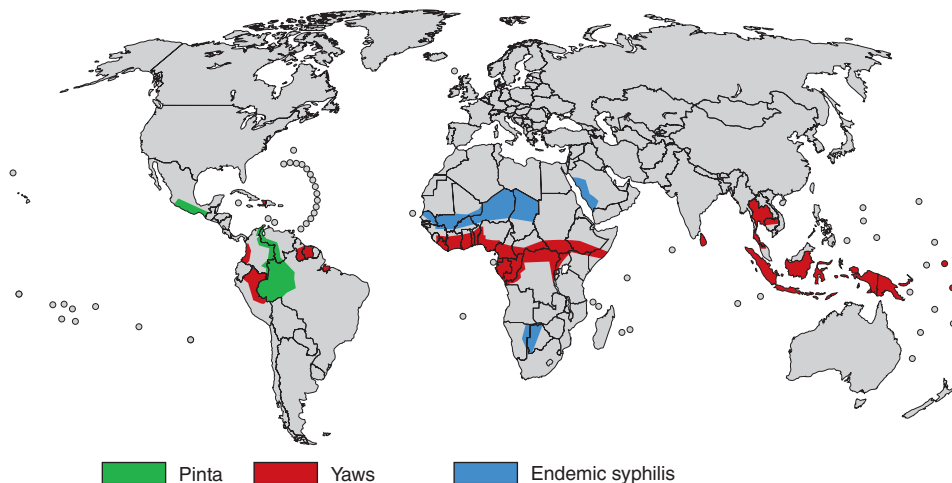


FIGURE 207e-1 Geographic distribution of endemic treponematoses. (Courtesy of the World Health Organization; updated from www.who.int/yaws/epidemiology/Map_yaws_90s.jpg.)

TABLE 207e-1 COMPARISON OF THE TREPONEMES AND ASSOCIATED DISEASES

Feature	Venereal Syphilis	Yaws	Endemic Syphilis	Pinta
Organism	<i>T. pallidum</i> subsp. <i>pallidum</i>	<i>T. pallidum</i> subsp. <i>pertenue</i>	<i>T. pallidum</i> subsp. <i>endemicum</i>	<i>T. carateum</i>
Common modes of transmission	Sexual, transplacental	Skin-to-skin	Mouth-to-mouth or via shared drinking/eating utensils	Skin-to-skin
Usual age of acquisition	Sexual maturity or in utero	Early childhood	Early childhood	Late childhood
Primary lesion	Cutaneous ulcer (chancre)	Papilloma, often ulcerative	Mucosal papule, rarely seen	Nonulcerating papule with satellites, pruritic
Common location	Genital, oral, anal	Extremities	Oral	Extremities, face
Secondary lesions	Mucocutaneous lesions; condylomata lata	Cutaneous papulosquamous lesions; condylomata lata, osteoperiostitis	Mucocutaneous lesions (mucous patch, split papule, condylomata lata); osteoperiostitis	Pintides, pigmented, pruritic
Infectious relapses	~25%	Common	Unknown	Unknown
Late complications	Gummas, cardiovascular and central nervous system involvement ^a	Destructive gummas of skin, bone, cartilage	Destructive gummas of skin, bone, cartilage	Nondestructive, dyschromic, achromic macules

^aCentral nervous system involvement and congenital infection in the endemic treponematoses have been postulated by some investigators (see text).

and possible CNS manifestations, or a low rate of symptomatic CNS disease. Some published evidence supports congenital transmission as well as cardiovascular, ophthalmologic, and CNS involvement in yaws and endemic syphilis. Although the reported studies have been small, have failed to control for other causes of CNS abnormalities, and in some instances have not included serologic confirmation, it may be erroneous to accept unquestioningly the frequently repeated belief that these organisms fail to cause such manifestations.

Yaws Also known as *pian*, *framboesia*, or *bouba*, yaws is characterized by the development of one or several primary lesions (“mother yaw”) followed by multiple disseminated skin lesions. All early skin lesions are infectious and may persist for many months; cutaneous relapses are common during the first 5 years. Late manifestations, affecting ~10% of untreated persons, are destructive and can involve skin, bone, and joints.

The infection is transmitted by direct contact with infectious lesions, often during play or group sleeping, and may be enhanced by disruption of the skin by insect bites or abrasions. After an average of 3–4 weeks, the first lesion begins as a papule—usually on an extremity—and then enlarges (particularly during moist warm weather) to become papillomatous or “raspberry-like” (thus the name “framboesia”) (Fig. 207e-2A). Regional lymphadenopathy develops, and the lesion usually heals within 6 months; dissemination is thought to occur during the early weeks of infection. A generalized secondary eruption (Fig. 207e-2B), accompanied by generalized lymphadenopathy, appears either concurrent with or after the primary lesion; may take several forms (macular, papular, or papillomatous); and may become

secondarily infected with other bacteria. Painful papillomatous lesions on the soles of the feet result in a crablike gait (“crab yaws”), and periostitis may result in nocturnal bone pain and polydactylitis. Late yaws is manifested by gummas of the skin and long bones, hyperkeratoses of the palms and soles, osteitis and periostitis, and hydrarthrosis. The late gummatous lesions are characteristically extensive. Destruction of the nose, maxilla, palate, and pharynx is termed *gangosa* and is similar to the destructive lesions seen in leprosy and leishmaniasis.

Endemic Syphilis The early lesions of endemic syphilis (*bejel*, *siti*, *dichuchwa*, *njovera*, *skerljevo*) are localized primarily to mucocutaneous and mucosal surfaces. The infection is reportedly transmitted by direct contact, by kissing, by pre-mastication of food, or by sharing of drinking and eating utensils. A role for insects in transmission has been suggested but is unproven. The initial lesion, usually an intraoral papule, often goes unrecognized and is followed by mucous patches (Fig. 207e-2C) on the oral mucosa and mucocutaneous lesions resembling the condylomata lata of secondary syphilis. This eruption may last for months or even years, and treponemes can readily be demonstrated in early lesions. Periostitis and regional lymphadenopathy are common. After a variable period of latency, late manifestations may appear, including osseous and cutaneous gummas. Destructive gummas, osteitis, and gangosa are more common in endemic syphilis than in yaws.

Pinta Pinta (*mal del pinto*, *carate*, *azul*, *purupuru*) is the most benign of the treponemal infections. This disease has three stages that are characterized by marked changes in skin color (Fig. 207e-2D), but pinta does not appear to cause destructive lesions or to involve



FIGURE 207e-2 Clinical manifestations of endemic treponematoses. A. Papillomatous initial lesion of early yaws. **B.** Disseminated lesions of early yaws. **C.** Mucous patches of endemic syphilis. **D.** Pigmented macules of pinta. (Photos published with permission from Dr. David Fegan, Brisbane, Australia [A and B]; and from PL Perine et al: *Handbook of Endemic Treponematoses*, Geneva, World Health Organization, 1984 [C and D].)

tissues other than the skin. The initial papule is most often located on the extremities or face and is pruritic. After one to many months of infection, numerous disseminated secondary lesions (*pintides*) appear. These lesions are initially red but become deeply pigmented, ultimately turning a dark slate blue. The secondary lesions are infectious and highly pruritic and may persist for years. Late pigmented lesions are called *dyschromic macules* and contain treponemes. Over time, most pigmented lesions show varying degrees of depigmentation, becoming brown and eventually white and giving the skin a mottled appearance. White achromic lesions are characteristic of the late stage.

DIAGNOSIS

Diagnosis of the endemic treponematoses is based on clinical manifestations and, when available, dark-field microscopy and serologic testing. The same serologic tests that are used for venereal syphilis ([Chap. 206](#)) become reactive during all treponemal infections. Although several targets have been evaluated for specific serodiagnosis, to date there is no antibody test that can discriminate among the different infections. The nonvenereal treponemal infections should be considered in the evaluation of a reactive syphilis serology in any person who has emigrated from an endemic area. Sensitive PCR assays can be used to confirm treponemal infection and to identify the etiologic agent in research laboratories.

TREATMENT ENDEMIC TREPONEMATOSES

The WHO-recommended therapy for patients and their contacts is benzathine penicillin G (1.2 million units IM for adults; 600,000 units for children <10 years old). This dose is half of that recommended for early venereal syphilis, and no controlled efficacy studies have been conducted. Definitive evidence of resistance to penicillin is lacking, although relapsing lesions have been reported after penicillin treatment in Papua New Guinea. A recent study in that nation demonstrated equivalence between IM benzathine penicillin G and a single

oral dose of azithromycin (30 mg/kg, up to a maximum of 2 g). This finding provided the WHO's revitalized yaws eradication program with a much easier regimen for use in mass treatment. Although macrolide resistance mutations are common in circulating strains of *T. pallidum* subspecies *pallidum* in many parts of the world, analysis of a limited number of yaws samples from Papua New Guinea has yielded no evidence of resistance mutations to date. Limited data suggest the efficacy of tetracycline for treatment of yaws, but no data exist for other endemic treponematoses. Solely on the basis of experience with venereal syphilis, it is thought that doxycycline or tetracycline (at doses appropriate for syphilis; [Chap. 206](#)) are alternatives for patients allergic to penicillin. A Jarisch-Herxheimer reaction ([Chap. 206](#)) may follow treatment of endemic treponematoses. Nontreponemal serologic titers (in the Venereal Disease Research Laboratory [VDRL] slide test or the rapid plasma reagin [RPR] test) usually decline after effective therapy, but patients may not become seronegative.

CONTROL

Buoyed by the successful elimination of yaws in India in 2006 and the availability of an inexpensive, single-dose oral drug for treatment, in 2012 the WHO renewed its efforts to eradicate yaws globally by 2020. Enthusiasm is high; several planning meetings have been held to develop country-specific plans of action; and resources are being sought. Some caution is warranted: The possible animal reservoir will need to be evaluated. There may be only a window of time during which countries can successfully use azithromycin for yaws eradication before resistance begins to appear in yaws organisms. Given the ongoing lower-dose azithromycin mass treatment campaigns against trachoma, often in populations also at high risk for yaws, development of macrolide resistance is likely at some point. Complete drug coverage and continued careful surveillance by health centers (the weak link in prior control efforts) will be essential for success.

A decorative sunburst graphic in shades of yellow and orange, positioned behind the chapter number 208.

208 Leptospirosis

Rudy A. Hartskeerl, Jiří F. P. Wagenaar

Leptospirosis is a globally important zoonotic disease whose apparent reemergence is illustrated by recent outbreaks on virtually all continents. The disease is caused by pathogenic *Leptospira* species and is characterized by a broad spectrum of clinical manifestations, varying from asymptomatic infection to fulminant, fatal disease. In its mild form, leptospirosis may present as nonspecific symptoms such as fever, headache, and myalgia. Severe leptospirosis, characterized by jaundice, renal dysfunction, and hemorrhagic diathesis, is often referred to as *Weil's syndrome*. With or without jaundice, severe pulmonary hemorrhage is increasingly recognized as an important presentation of severe disease.

ETIOLOGIC AGENT

Leptospira species are spirochetes belonging to the order Spirochaetales and the family Leptospiraceae. Traditionally, the genus *Leptospira* comprised two species: the pathogenic *L. interrogans* and the free-living *L. biflexa*, now designated *L. interrogans* sensu lato and *L. biflexa* sensu lato, respectively. Twenty-two *Leptospira* species with pathogenic (10 species), intermediate (5 species), and nonpathogenic (7 species) status have now been described on the basis of phylogenetic and virulence

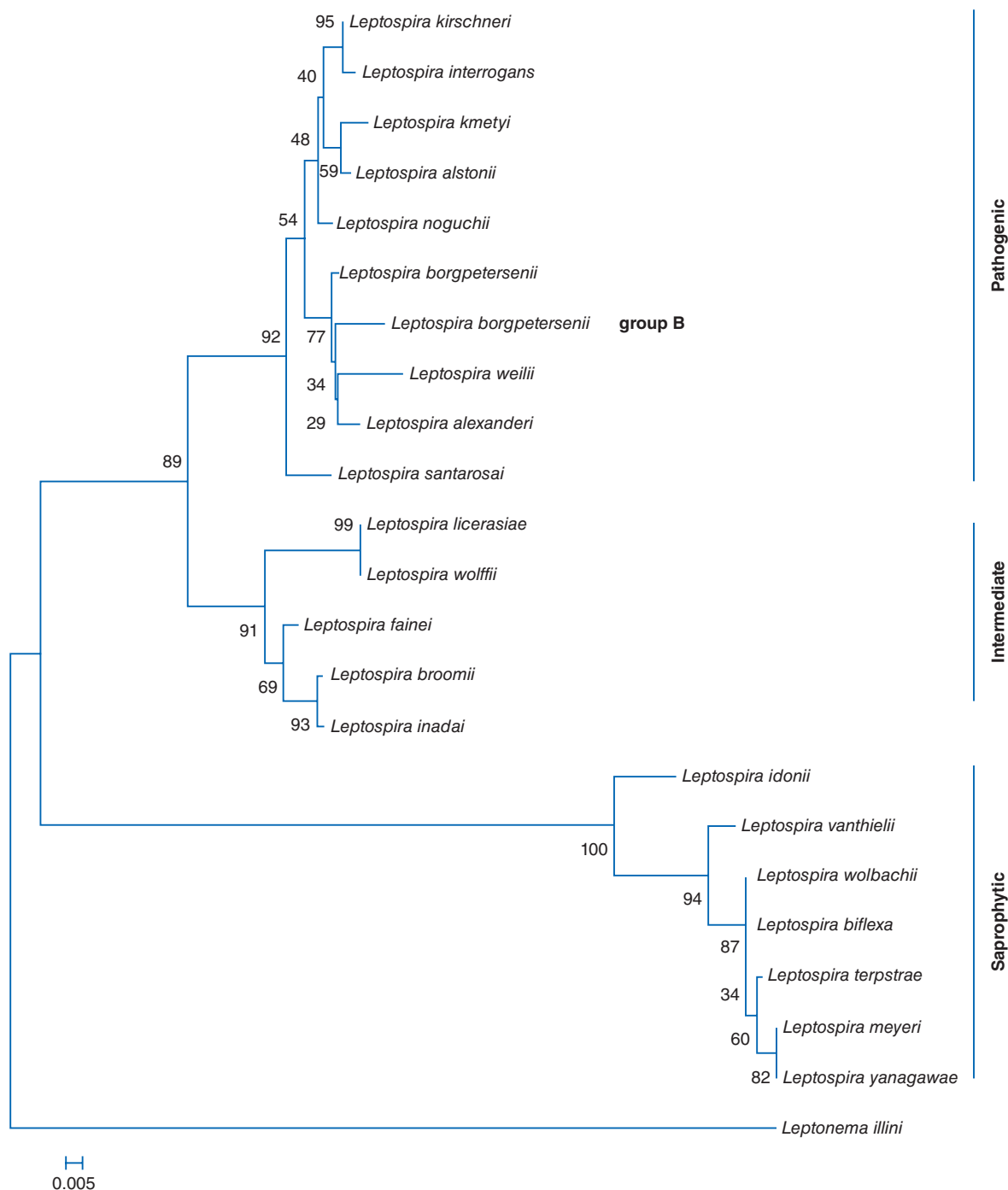



FIGURE 208-1 Differentiation of pathogenic, intermediate, and nonpathogenic (saprophytic) *Leptospira* species by molecular phylogenetic analysis using the *rrs* gene and including the potentially new pathogenic species *Leptospira borgpetersenii* group B and the saprophytic species *Leptospira idonii*. Scale bar indicates the rate of nucleotide substitutions per base pair. (Figure prepared and provided by Dr. A. Ahmed, KIT Biomedical Research, Amsterdam, The Netherlands.)

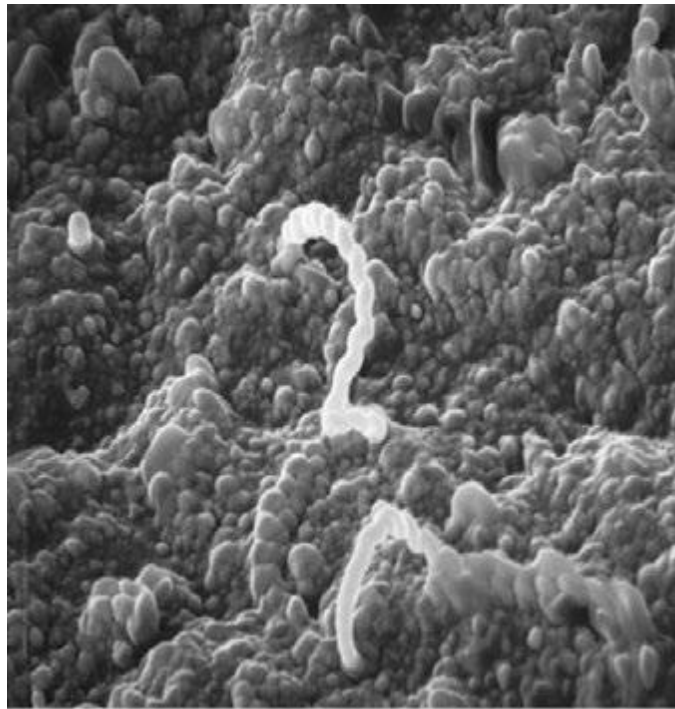
analyses (Fig. 208-1). Genome sequences of five *Leptospira* species (*L. biflexa*, *L. interrogans*, *L. santarosai*, *L. borgpetersenii*, and *L. licerasiae*) have been published, and the availability of genome sequences of a wide variety of *Leptospira* strains will undoubtedly lead to a better understanding of the pathogenesis of leptospirosis. However, classification based on serologic differences better serves clinical, diagnostic, and epidemiologic purposes. Pathogenic *Leptospira* species are divided into serovars according to their antigenic composition. More than 250 serovars make up the 26 serogroups.

Leptospires are coiled, thin, highly motile organisms that have hooked ends and two periplasmic flagella, with polar extrusions from the cytoplasmic membrane that are responsible for motility (Fig. 208-2). These organisms are 6–20 μm long and $\sim 0.1 \mu\text{m}$ wide;

they stain poorly but can be seen microscopically by dark-field examination and after silver impregnation staining of tissues. Leptospires require special media and conditions for growth; it may take weeks to months for cultures to become positive.

EPIDEMIOLOGY

 Leptospirosis has a worldwide distribution but occurs most commonly in the tropics and subtropics because the climate and occasionally poor hygienic conditions favor the pathogen's survival and distribution. In most countries, leptospirosis is an underappreciated problem. Most cases occur in men, with a peak incidence during the summer and fall in both the Northern and Southern Hemispheres and during the rainy season in the tropics.



1
0.3 μm

FIGURE 208-2 Transmission electron microscopic image of *Leptospira interrogans* invading equine conjunctival tissue. (Image kindly provided by Dr. JE Nally, National Animal Disease Center, U.S. Department of Agriculture, Ames, IA. This image appears on the homepage of the European Leptospirosis Society website [<http://eurolepto.ucd.ie/>].)

Reliable data on morbidity and mortality from leptospirosis have gradually started to appear. Current information on global human leptospirosis varies but indicates that approximately 1 million severe cases occur per year, with a mean case–fatality rate of nearly 10%.

As a zoonosis, leptospirosis affects almost all mammalian species and represents a significant veterinary burden. Rodents, especially rats, are the most important reservoir, although other wild mammals as well as domestic and farm animals may also harbor these microorganisms. Leptospire establish a symbiotic relationship with their host and can persist in the urogenital tract for years. Some serovars are generally associated with particular animals—e.g., Icterohaemorrhagiae and Copenhageni with rats, Grippityphosa with voles, Hardjo with cattle, Canicola with dogs, and Pomona with pigs—but may occur in other animals as well.

Leptospirosis presents as both an endemic and an epidemic disease. Transmission of leptospire may follow direct contact with urine, blood, or tissue from an infected animal or, more commonly, exposure to environmental contamination. The dogma that human-to-human transmission is very rare is challenged by recent findings on household clustering, asymptomatic renal colonization, and prolonged excretion of leptospire. (Both of the latter features imply human infection sources that are not recognized.) Because leptospire can survive in a humid environment for many months, water is an important vehicle in their transmission. Epidemics of leptospirosis are not well understood. Outbreaks may result from exposure to flood waters contaminated by urine from infected animals, as has been reported from several countries. However, it is also true that outbreaks may occur without floods, and floods often occur without outbreaks.

The vast majority of infections with *Leptospira* cause no or only mild disease in humans. A small percentage of infections (~1%) lead to severe, potentially fatal complications. The proportion of leptospirosis cases that are mild is unknown because patients either do not seek or do not have access to medical care or because the nonspecific symptoms are interpreted as an influenza-like illness. Reported cases surely

represent a significant underestimation of the total number. Certain occupational groups are at especially high risk, including veterinarians, agricultural workers, sewage workers, slaughterhouse employees, and workers in the fishing industry. Risk factors include direct or indirect contact with animals, including exposure to water and soil contaminated with animal urine. Leptospirosis has also been recognized in deteriorating inner cities and suburban areas where rat populations are expanding.

Recreational exposure and domestic-animal contact are prominent sources of leptospirosis. Recreational freshwater activities, such as canoeing, windsurfing, swimming, and waterskiing, place persons at risk for infection. Several outbreaks have followed sporting events. For example, an outbreak took place in 1998 among athletes after a triathlon in Springfield, Illinois. Ingestion of one or more swallows of lake water during the swimming leg of the triathlon was a prominent risk factor for illness. Heavy rains that preceded the triathlon, with consequent agricultural runoff, are likely to have increased the level of leptospiral contamination in the lake water. In another outbreak, 42% of participants contracted leptospirosis during the 2000 Eco-Challenge-Sabah multisport endurance race in Malaysian Borneo. Swimming in the Segama River was shown to be an independent risk factor.

In addition, leptospirosis is a traveler's disease. Large proportions of patients acquire the infection while traveling in tropical countries, usually during adventurous activities such as whitewater rafting, jungle trekking, and caving. Transmission via laboratory accidents has been reported but is rare. New data indicate that leptospirosis may develop after unanticipated immersion in contaminated water (e.g., in an automobile accident) more frequently than has generally been thought and can also result from an animal bite.

PATHOGENESIS

Transmission occurs through cuts, abraded skin, or mucous membranes, especially the conjunctival and oral mucosa. After entry, the organisms proliferate, cross tissue barriers, and disseminate hematogenously to all organs (*leptospiremic phase*). During this initial incubation period, leptospire can be isolated from the bloodstream (**Fig. 208-3**). The organisms are able to survive in the nonimmune host: they evade complement-mediated killing by binding factor H, a strong inhibitor of the complement system, on their surface. Moreover, pathogenic leptospire resist ingestion and killing by neutrophils, monocytes, and macrophages. During the immune phase, the appearance of antibodies coincides with the disappearance of leptospire from the blood. However, the bacteria persist in various organs, including liver, lung, kidney, heart, and brain. Autopsy findings illustrate the involvement of multiple organ systems in severe disease. Renal pathology shows both acute tubular damage and interstitial nephritis. Acute tubular lesions progress in time to interstitial edema and acute tubular necrosis. Severe nephritis is observed in patients who survive long enough to develop it and seems to be a secondary response to acute epithelial damage. The reported deregulation of the expression of several transporters along the nephron, including the proximal sodium-hydrogen exchanger 3 (NHE3), aquaporins 1 and 2 (AQP1 and AQP2), Na⁺-K⁺ ATPase, and the Na-K-2Cl cotransporter NKCC2, contributes to tubular potassium wasting, hypokalemia, and polyuria. Histopathology of the liver shows focal necrosis, foci of inflammation, and plugging of bile canaliculi. Widespread hepatocellular necrosis is not found. Petechiae and hemorrhages are observed in the heart, lungs (**Fig. 208-4**), kidneys (and adrenals), pancreas, liver, gastrointestinal tract (including retroperitoneal fat, mesentery, and omentum), muscles, prostate, testis, and brain (subarachnoid bleeding). Several studies show an association between hemorrhage and thrombocytopenia. Although the underlying mechanisms of thrombocytopenia have not been elucidated, it seems likely that platelet consumption plays an important role. A consumptive coagulopathy may occur, with elevated markers of coagulation activation (thrombin–antithrombin complexes, prothrombin fragments 1 and 2, D-dimer), diminished anticoagulant markers (antithrombin, protein C), and deregulated fibrinolytic activity. Overt disseminated intravascular coagulation (DIC) has been documented in patients from Thailand and Indonesia. Elevated plasma levels of soluble E-selectin and

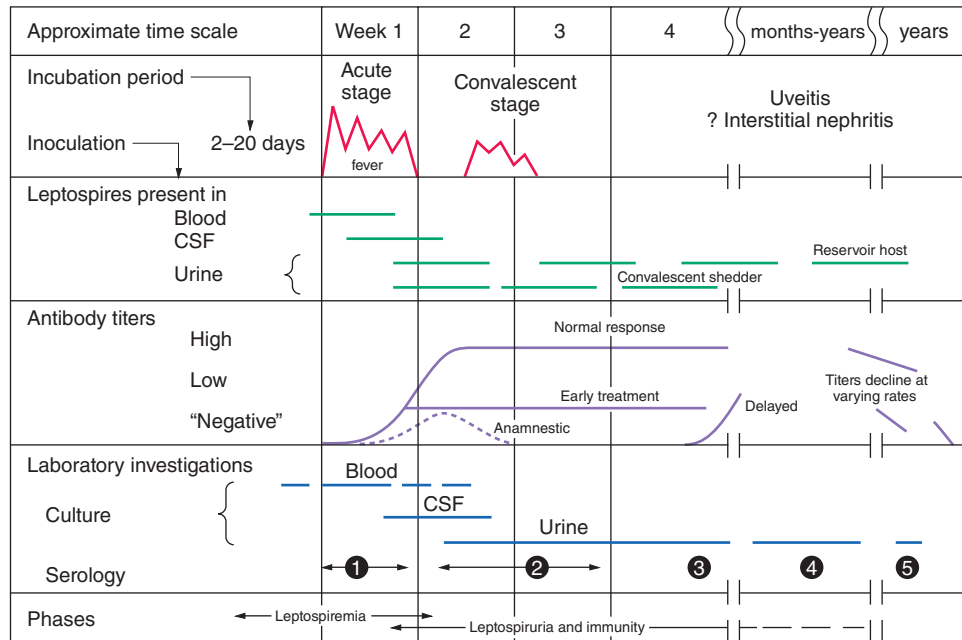


FIGURE 208-3 Biphasic nature of leptospirosis and relevant investigations at different stages of disease. Note that an incubation period of up to 1 month has now been documented. Specimens 1 and 2 for serology are acute-phase serum samples; specimen 3 is a convalescent-phase serum sample that may facilitate detection of a delayed immune response; and specimens 4 and 5 are follow-up serum samples that can provide epidemiologic information, such as the presumptive infecting serogroup. CSF, cerebrospinal fluid. (Reprinted as adapted by PN Levett: *Clin Microbiol Rev* 14:296, 2001 [from LH Turner: *Leptospirosis*. *BMJ* 1:231, 1969] with permission from the American Society for Microbiology and the BMJ Publishing Group.)

von Willebrand factor in patients with leptospirosis reflect endothelial cell activation. Experimental models show that pathogenic leptospires or leptospiral proteins are able to activate endothelial cells in vitro and to disrupt endothelial-cell barrier function, promoting dissemination. Platelets have been shown to aggregate on activated endothelium in the human lung, whereas histology reveals swelling of activated endothelial cells but no evident vasculitis or necrosis. Immunoglobulin and complement deposition have been demonstrated in lung tissue involved in pulmonary hemorrhage.

Leptospira species have a typical double-membrane cell wall structure harboring a variety of membrane-associated proteins, including an unusually high number of lipoproteins. The peptidoglycan layer is located close to the cytoplasmic membrane. The lipopolysaccharide (LPS) in the outer membrane has an unusual structure with a relatively low endotoxic potency. Pathogenic leptospires contain a variety of genes coding for proteins involved in motility and in cell and tissue adhesion and invasion that represent potential virulence factors. Many of these are surface-exposed outer-membrane proteins

(OMPs). To date, the only leptospiral virulence factor shown to satisfy Koch's molecular postulates is *loa22* encoding a surface-exposed protein with an unknown function. However, the gene is not confined to pathogenic *Leptospira* species.

Immunity to *Leptospira* depends on the production of circulating antibodies to serovar-specific LPS. It is unclear whether other antigens play a significant role in protective humoral immunity. Moreover, immunity may not be confined to antibody responses; involvement of the innate-immune Toll-like receptor 2 (TLR2) and TLR4 activation pathways in controlling infection has been demonstrated, whereas in vaccinated cattle a cell-mediated immune response is correlated with protection.

It is likely that several surface-exposed proteins mediate leptospire–host cell interactions, and these proteins may represent candidate vaccine components. Although animal-model studies have shown various degrees of vaccine efficacy for various putative virulence-associated OMPs, it is not yet clear whether such proteins elicit acceptable levels of sterilizing immunity.

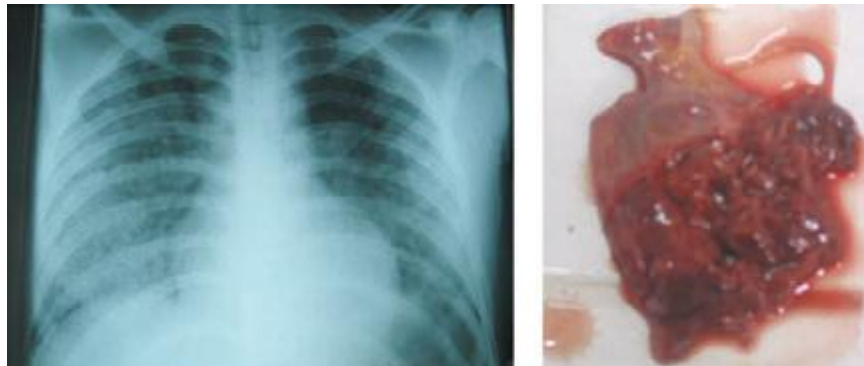


FIGURE 208-4 Severe pulmonary hemorrhage in leptospirosis. **Left panel:** Chest x-ray. **Right panel:** Gross appearance of right lower lobes of lung at autopsy. This patient, a 15-year-old from the Peruvian Amazonian city of Iqitos, died several days after presentation with acute illness, jaundice, and hemoptysis. Blood culture yielded *Leptospira interrogans* serovar Copenhageni/Icterohaemorrhagiae. (Adapted with permission from E Segura et al: *Clin Infect Dis* 40:343, 2005. © 2005 by the Infectious Diseases Society of America.)

Although leptospirosis is a potentially fatal disease with bleeding and multiorgan failure as its clinical hallmarks, the majority of cases are thought to be relatively mild, presenting as the sudden onset of a febrile illness. The incubation period is usually 1–2 weeks but ranges from 1 to 30 days. (Figure 208-3 indicates a slightly different range, but an incubation period of up to 1 month has now been documented.) Leptospirosis is classically described as biphasic. The acute *leptospiremic phase* is characterized by fever of 3–10 days' duration, during which time the organism can be cultured from blood. During the *immune phase*, resolution of symptoms may coincide with the appearance of antibodies, and leptospire can be cultured from the urine. The distinction between the first and second phases is not always clear: milder cases do not always include the second phase, and severe disease may be monophasic and fulminant. The idea that distinct clinical syndromes are associated with specific serogroups has been refuted, although some serovars tend to cause more severe disease than others.

Mild Leptospirosis Most patients are asymptomatic or only mildly ill and do not seek medical attention. Serologic evidence of past inapparent infection is frequently found in persons who have been exposed but have not become ill. Mild symptomatic leptospirosis usually presents as a flu-like illness of sudden onset, with fever, chills, headache, nausea, vomiting, abdominal pain, conjunctival suffusion (redness without exudate), and myalgia. Muscle pain is intense and especially affects the calves, back, and abdomen. The headache is intense, localized to the frontal or retroorbital region (resembling that occurring in dengue), and sometimes accompanied by photophobia. Aseptic meningitis may be present and is more common among children than among adults. Although *Leptospira* can be cultured from the cerebrospinal fluid (CSF) in the early phase, the majority of cases follow a benign course with regard to the central nervous system; symptoms disappear within a few days but may persist for weeks.

Physical examination may include any of the following findings, none of which is pathognomonic for leptospirosis: fever, conjunctival suffusion, pharyngeal injection, muscle tenderness, lymphadenopathy, rash, meningismus, hepatomegaly, and splenomegaly. If present, the rash is often transient; may be macular, maculopapular, erythematous, or hemorrhagic (petechial or ecchymotic); and may be misdiagnosed as due to scrub typhus or viral infection. Lung auscultation may reveal crackles, and mild jaundice may be present.

The natural course of mild leptospirosis usually involves spontaneous resolution within 7–10 days, but persistent symptoms have been documented. In the absence of a clinical diagnosis and antimicrobial therapy, the mortality rate in mild leptospirosis is low.

Severe Leptospirosis Although the onset of severe leptospirosis may be no different from that of mild leptospirosis, severe disease is often rapidly progressive and is associated with a case–fatality rate ranging from 1 to 50%. Higher mortality rates are associated with an age >40, altered mental status, acute renal failure, respiratory insufficiency, hypotension, and arrhythmias. The classic presentation, often referred to as *Weill's syndrome*, encompasses the triad of hemorrhage, jaundice, and acute kidney injury.

Patients die of septic shock with multiorgan failure and/or severe bleeding complications that most commonly involve the lungs (pulmonary hemorrhage), gastrointestinal tract (melena, hemoptysis), urogenital tract (hematuria), and skin (petechiae, ecchymosis, and bleeding from venipuncture sites). Pulmonary hemorrhage (with or without jaundice) is now recognized as a widespread public health problem, presenting with cough, chest pain, respiratory distress, and hemoptysis that may not be apparent until patients are intubated.

Jaundice occurs in 5–10% of all patients with leptospirosis; it can be profound and give an orange cast to the skin but usually is not associated with fulminant hepatic necrosis. Physical examination may reveal an enlarged and tender liver.

Acute kidney injury is common in severe disease, presenting after several days of illness, and can be either nonoliguric or oliguric. Typical electrolyte abnormalities include hypokalemia and

hyponatremia. Loss of magnesium in the urine is uniquely associated with leptospiral nephropathy. Hypotension is associated with acute tubular necrosis, oliguria, or anuria, requiring fluid resuscitation and sometimes vasopressor therapy. Hemodialysis can be life-saving, with renal function typically returning to normal in survivors.

Other syndromes include (necrotizing) pancreatitis, cholecystitis, skeletal muscle involvement, rhabdomyolysis (with moderately elevated serum creatine kinase levels), and neurologic manifestations including aseptic meningitis. Cardiac involvement is commonly reflected on the electrocardiogram as nonspecific ST- and T-wave changes. Repolarization abnormalities and arrhythmias are considered poor prognostic factors. Myocarditis has been described. Rare hematologic complications include hemolysis, thrombotic thrombocytopenic purpura, and hemolytic-uremic syndrome.

Long-term symptoms following severe leptospirosis include fatigue, myalgia, malaise, and headache and may persist for years. Autoimmune-associated uveitis, a potentially chronic condition, is a recognized sequela of leptospirosis.

DIAGNOSIS

The clinical diagnosis of leptospirosis should be based on an appropriate exposure history combined with any of the protean manifestations of the disease. Returning travelers from endemic areas usually have a history of recreational freshwater activities or other mucosal or percutaneous contact with contaminated surface waters or soil. For nontravelers, recreational water contact and occupational hazards that involve direct or indirect animal contact should be explored (see “Epidemiology,” above).

Although biochemical, hematologic, and urinalysis findings in acute leptospirosis are nonspecific, certain patterns may suggest the diagnosis. Laboratory results usually show signs of a bacterial infection, including leukocytosis with a left shift and elevated markers of inflammation (C-reactive protein level and erythrocyte sedimentation rate). Thrombocytopenia (platelet count $\leq 100 \times 10^9/L$) is common and is associated with bleeding and renal failure. In severe disease, signs of coagulation activation may be present, varying from borderline abnormalities to a serious derangement compatible with DIC as defined by international criteria. The kidneys are invariably involved in leptospirosis. Related findings range from urinary sediment changes (leukocytes, erythrocytes, and hyaline or granular casts) and mild proteinuria in mild disease to renal failure and azotemia in severe leptospirosis. Nonoliguric hypokalemic renal insufficiency (see “Clinical Manifestations,” above) is characteristic of early leptospirosis. Serum bilirubin levels may be high, whereas rises in aminotransferase and alkaline phosphatase levels are usually moderate. Although clinical symptoms of pancreatitis are not a common finding, amylase levels are often elevated. When symptoms of aseptic meningitis develop, examination of the CSF shows pleocytosis that can range from a few cells to >1000 cells/ μL , with a polymorphonuclear cell predominance. The protein concentration in the CSF may be elevated; CSF glucose levels are normal.

In severe leptospirosis, pulmonary radiographic abnormalities are more common than would be expected on the basis of physical examination (Fig. 208-4). The most common radiographic finding is a patchy bilateral alveolar pattern that corresponds to scattered alveolar hemorrhage. These abnormalities predominantly affect the lower lobes. Other findings include pleura-based densities (representing areas of hemorrhage) and diffuse ground-glass attenuation typical of acute respiratory distress syndrome (ARDS).

A definitive diagnosis of leptospirosis is based on isolation of the organism from the patient, on a positive result in the polymerase chain reaction (PCR), or on seroconversion or a rise in antibody titer. In cases with strong clinical evidence of infection, a single antibody titer of 1:200–1:800 (depending on whether the case occurs in a low- or high-endemic area) in the microscopic agglutination test (MAT) is required. Preferably, a fourfold or greater rise in titer is detected between acute- and convalescent-phase serum specimens. Antibodies generally do not reach detectable levels until the second week of illness. The antibody response can be affected by early treatment with antibiotics.

The MAT, which uses a battery of live leptospiral strains, and the enzyme-linked immunosorbent assay (ELISA), which uses a broadly reacting antigen, are the standard serologic procedures. The MAT usually is available only in specialized laboratories and is used for determination of the antibody titer and for tentative identification of the involved leptospiral serogroup—and, when epidemiologic background information is available, the putative serovar. This point underscores the importance of testing antigens representative of the serovars prevalent in the particular geographic area. However, cross-reactions occur frequently, and thus definitive identification of the infecting serovar or serogroup is not possible without isolation of the causative organism. Because serologic testing lacks sensitivity in the early acute phase of the disease (up to day 5), it cannot be used as the basis for a timely decision about whether to start treatment.

In addition to the MAT and the ELISA, various rapid tests with diagnostic value have been developed, and some of these are commercially available. These rapid tests mainly apply lateral flow, (latex) agglutination, or ELISA methodology and are reasonably sensitive and specific, although results reported in the literature vary, probably as a consequence of differences in test interpretation, (re)exposure risks, serovar distribution, and the use of biased serum panels. These methods do not require culture or MAT facilities and are useful in settings that lack a strong medical infrastructure. PCR methodologies, notably real-time PCR, have become increasingly widely implemented. Compared with serology, PCR offers a great advantage: the capacity to confirm the diagnosis of leptospirosis with a high degree of accuracy during the first 5 days of illness.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of leptospirosis is broad, reflecting the diverse clinical presentations of the disease. Although leptospirosis transmission is more common in tropical and subtropical regions, the absence of a travel history does not exclude the diagnosis. When fever, headache, and myalgia predominate, influenza and other common and less common (e.g., dengue and chikungunya) viral infections should be considered. Malaria, typhoid fever, ehrlichiosis, viral hepatitis, and acute HIV infection may mimic the early stages of leptospirosis and are important to recognize. Rickettsial diseases, hantavirus infections (hemorrhagic fever with renal syndrome or hantavirus cardiopulmonary syndrome), and dengue share epidemiologic and clinical features with leptospirosis. Dual infections have been reported. In this light, it is advisable to conduct serologic testing for hantavirus, rickettsiae, and dengue virus when leptospirosis is suspected. When bleeding is detected, dengue hemorrhagic fever and other viral hemorrhagic fevers, including hantavirus infection, yellow fever, Rift Valley fever, filovirus infections, and Lassa fever, should be considered.

TREATMENT LEPTOSPIROSIS

Severe leptospirosis should be treated with IV penicillin (Table 208-1) as soon as the diagnosis is considered. Leptospire are highly susceptible to a broad range of antibiotics, and early intervention may prevent the development of major organ system failure or lessen its severity. Although studies supporting antibiotic therapy have produced conflicting results, clinical trials are difficult to perform in settings where patients frequently present for medical care with late stages of disease. Antibiotics are less likely to benefit patients in whom organ damage has already occurred. Two open-label randomized studies comparing penicillin with parenteral cefotaxime, parenteral ceftriaxone, and doxycycline showed no significant differences among the antibiotics with regard to complications and mortality risk. Thus ceftriaxone, cefotaxime, or doxycycline is a satisfactory alternative to penicillin for the treatment of severe leptospirosis.

In mild cases, oral treatment with doxycycline, azithromycin, ampicillin, or amoxicillin is recommended. In regions where rickettsial diseases are coendemic, doxycycline or azithromycin is the drug of choice. In rare instances, a Jarisch-Herxheimer reaction develops within hours after the initiation of antimicrobial therapy.

TABLE 208-1 TREATMENT AND CHEMOPROPHYLAXIS OF LEPTOSPIROSIS IN ADULTS^a

Indication	Regimen
Treatment	
Mild leptospirosis	Doxycycline ^b (100 mg PO bid) <i>or</i> Amoxicillin (500 mg PO tid) <i>or</i> Ampicillin (500 mg PO tid)
Moderate/severe leptospirosis	Penicillin (1.5 million units IV or IM q6h) <i>or</i> Ceftriaxone (2 g/d IV) <i>or</i> Cefotaxime (1 g IV q6h) <i>or</i> Doxycycline (loading dose of 200 mg IV, then 100 mg IV q12h)
Chemoprophylaxis^c	
	Doxycycline ^b (200 mg PO once a week) <i>or</i> Azithromycin (250 mg PO once or twice a week)

^aAll regimens are given for 7 days. ^bDoxycycline should not be given to pregnant women or children. ^cThe efficacy of doxycycline prophylaxis in endemic or epidemic settings remains unclear. Experiments in animal models and a cost-effectiveness model indicate that azithromycin has a number of characteristics that may make it efficacious in treatment and prophylaxis.


Aggressive supportive care for leptospirosis is essential and can be life-saving. Patients with nonoliguric renal dysfunction require aggressive fluid and electrolyte resuscitation to prevent dehydration and precipitation of oliguric renal failure. Peritoneal dialysis or hemodialysis should be provided to patients with oliguric renal failure. Rapid initiation of hemodialysis has been shown to reduce mortality risk and typically is necessary only for short periods. Patients with pulmonary hemorrhage may have reduced pulmonary compliance (as seen in ARDS) and may benefit from mechanical ventilation with low tidal volumes to avoid high ventilation pressures. Evidence is contradictory for the use of glucocorticoids and desmopressin as adjunct therapy for pulmonary involvement associated with severe leptospirosis.

PROGNOSIS

Most patients with leptospirosis recover. However, post-leptospirosis symptoms, mainly of a depression-like nature, may occur and persist for years after the acute disease. Mortality rates are highest among patients who are elderly and those who have severe disease (pulmonary hemorrhage, Weil's syndrome). Leptospirosis during pregnancy is associated with high fetal mortality rates. Long-term follow-up of patients with renal failure and hepatic dysfunction has documented good recovery of renal and hepatic function.

PREVENTION

Individuals who may be exposed to *Leptospira* through their occupations or their involvement in recreational freshwater activities should be informed about the risks. Measures for controlling leptospirosis include avoidance of exposure to urine and tissues from infected animals through proper eyewear, footwear, and other protective equipment. Targeted rodent control strategies could be considered.

 Vaccines for agricultural and companion animals are generally available, and their use should be encouraged. The veterinary vaccine used in a given area should contain the serovars known to be present in that area. Unfortunately, some vaccinated animals still excrete leptospire in their urine. Vaccination of humans against a specific serovar prevalent in an area has been undertaken in some European and Asian countries and has proved effective. Although a large-scale trial of vaccine in humans has been reported from Cuba, no conclusions can be drawn about efficacy and adverse reactions because of insufficient details on study design. The efficacy of chemoprophylaxis with doxycycline (200 mg once a week) or azithromycin (in pregnant women and children) is being disputed, but focused pre- and postexposure administration is indicated in instances of well-defined short-term exposure (Table 208-1).

209 Relapsing Fever

Alan G. Barbour

Relapsing fever is caused by infection with any of several species of *Borrelia* spirochetes. Physicians in ancient Greece distinguished relapsing fever from other febrile disorders by its characteristic clinical presentation: two or more fever episodes separated by varying periods of well-being. In the nineteenth century, relapsing fever was one of the first diseases to be associated with a specific microbe by virtue of its characteristic laboratory finding: the presence of large numbers of spirochetes of the genus *Borrelia* in the blood.

The host responds with systemic inflammation that results in an illness ranging from a flulike syndrome to sepsis. Other manifestations are the consequences of central nervous system (CNS) involvement and coagulopathy. Antigenic variation of the spirochetes' surface proteins accounts for the infection's relapsing course. Acquired immunity follows the serial development of antibodies to each of the several variants appearing during an infection. Treatment with antibiotics results in rapid cure but at the risk of a moderate to severe Jarisch-Herxheimer reaction.



Louse-borne relapsing fever caused large epidemics well into the twentieth century and currently occurs in northeastern Africa. At present, however, most cases of relapsing fever are tick-borne in origin. Sporadic cases and small outbreaks are focally distributed on most continents, with Africa most affected. In North America, the majority of reports of relapsing fever have been from the western United States and Canada. Nevertheless, the recent discovery that another species in the relapsing fever group causes human disease in the same geographic distribution as Lyme disease (Chap. 210) confounds epidemiologic distinctions between the two major types of *Borrelia* infection.

ETIOLOGIC AGENT

Coiled, thin microscopic filaments that swim in one direction and then coil up before heading in another were first observed in the blood of patients with relapsing fever in the 1880s (www.youtube.com/watch?v=VxDPV2lBd9U). These microbes were categorized as spirochetes and grouped as several species in the genus *Borrelia*. It was not until the 1960s that the organisms were isolated in pure culture. The breakthrough cultivation medium and its derivatives are rich in their ingredients, ranging from simple (e.g., amino acids and *N*-acetylglucosamine) to more complex (e.g., serum and protein hydrolysates). The limited biosynthetic capacity of *Borrelia* cells is accounted for by a genome content one-quarter that of *Escherichia coli*.

Like other spirochetes, the helix-shaped *Borrelia* cells have two membranes, the outer of which is more loosely secured than in other double-membrane bacteria, such as *E. coli*. As a consequence, fixed organisms with damaged membranes can assume a variety of morphologies in smears and histologic preparations. The flagella of spirochetes run between the two membranes and are not on the cell surface. Although technically gram-negative in their staining properties, the

10- to 20- μ m-long *Borrelia* cells, with a diameter of 0.1–0.2 μ m, are too narrow to be seen by bright-field microscopy of Gram-stained specimens.

EPIDEMIOLOGY



The several species of *Borrelia* that cause relapsing fever have restricted geographic distributions (Table 209-1). The exception is *Borrelia recurrentis*, which is also the only species transmitted by the louse. Louse-borne relapsing fever (LBRF) is usually acquired from a body louse (*Pediculus humanus corporis*), with humans serving as the reservoir. Acquisition occurs not from the bite itself but from either rubbing the insect's feces into the bite site with the fingers in response to irritation or inoculation of feces into the conjunctivae or an open wound. Although LBRF transmission is currently limited to Ethiopia and adjacent countries, the disease has had a global distribution in the past, and that potential remains. Epidemics with thousands of cases of LBRF can occur under circumstances of famine, natural disaster, refugee migration, and war.

All other known species of relapsing fever agents are tick-borne—in most cases, by soft ticks of the genus *Ornithodoros* (Fig. 209-1). Tick-borne relapsing fever (TBRF) is found on most continents but is absent or rare in tropical, low-desert, arctic, or alpine environments. For most species, the reservoirs of infection are small to medium-sized mammals, usually rodents but sometimes pigs and other domestic animals living in or around human habitats. However, one species, *Borrelia duttonii* in sub-Saharan Africa, is largely maintained by tick transmission between human hosts. In North America, TBRF occurs as single cases or small case clusters through transient exposure of persons to infested buildings or caves in less populated areas where the rodent reservoirs have nests. The two main *Borrelia* species involved in North America are *Borrelia hermsii* (in the mountainous west) and *Borrelia turicatae* (in the southwestern and south-central regions). The soft tick vectors typically feed for no more than 30 min, usually without being noticed, while the victim is sleeping. Transovarial transmission from one generation of ticks to the next means that infection risk may persist in an area long after incriminated mammalian reservoirs have been eradicated.

A newly recognized pathogen, *Borrelia miyamotoi*, belongs to the clade of relapsing fever species but is transmitted to humans from other mammals by hard ticks (e.g., *Ixodes scapularis* in the eastern United States) that also transmit Lyme disease, babesiosis, anaplasmosis, ehrlichiosis, and arboviral encephalitis. *B. miyamotoi* is acquired through outdoor activities and through contact with ticks in forested and shrubby areas during recreation, work, or activities around the home. In residents of areas where *B. miyamotoi* and *Borrelia burgdorferi* coexist, the prevalence of antibodies to the former is about one-third of that to the latter.

PATHOGENESIS AND IMMUNITY

Unlike LBRF spirochetes, TBRF spirochetes enter the body in the tick's saliva with the onset of feeding. From an inoculum of a few cells, the spirochetes proliferate in the blood, doubling every 6 h to numbers of 10^6 – 10^7 /mL or more. *Borrelia* species are extracellular pathogens; their

TABLE 209-1 RELAPSING FEVER BORRELIA SPECIES, BY GEOGRAPHIC REGION, VECTOR, AND PRIMARY RESERVOIR

Species	Region(s)	Arthropod Vector(s)	Primary Reservoir
<i>B. crocidurae</i>	Africa	<i>Ornithodoros erraticus</i> , <i>O. sonrai</i> (soft ticks)	Mammals
<i>B. duttonii</i>	Africa	<i>O. moubata</i>	Humans
<i>B. hermsii</i>	North America	<i>O. hermsii</i>	Mammals
<i>B. hispanica</i>	Europe, North Africa	<i>O. erraticus</i> complex	Mammals
<i>B. miyamotoi</i>	Eurasia, North America	<i>Ixodes</i> species (hard ticks)	Mammals
<i>B. persica</i>	Eurasia	<i>O. tholozani</i>	Mammals
<i>B. recurrentis</i>	Africa, global ^a	<i>Pediculus humanus corporis</i> (human body louse)	Humans
<i>B. turicatae</i>	North America	<i>O. turicata</i>	Mammals
<i>B. venezuelensis</i>	Central and South America	<i>O. rudis</i>	Mammals

^aAlthough transmission is currently limited to Ethiopia and adjacent countries, *B. recurrentis* infection has had a global distribution in the past, and that potential remains.

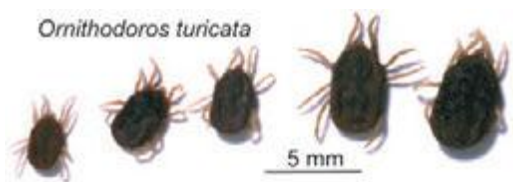


FIGURE 209-1 *Ornithodoros turicata* soft ticks of different ages.

presence inside cells likely represents a dead end for the bacteria after phagocytosis. Binding of the spirochetes to erythrocytes leads to aggregation of red blood cells, their sequestration in the spleen and liver, and hepatosplenomegaly and anemia. A bleeding disorder is probably the consequence of thrombocytopenia, impaired hepatic production of clotting factors, and/or blockage of small vessels by aggregates of spirochetes, erythrocytes, and platelets. Some species are neurotropic and frequently enter the brain, where they are comparatively sheltered from host immunity. Relapsing fever spirochetes can cross the maternal-fetal barrier and cause placental damage and inflammation, leading to intrauterine growth retardation and congenital infection.

Although *Borrelia* species do not have potent exotoxins or a lipopolysaccharide endotoxin, they have abundant membrane-associated lipoproteins whose recognition and binding by Toll-like receptors on host cells can lead to a proinflammatory process similar to that in endotoxemia, with elevations of tumor necrosis factor α , interleukin 6, and interleukin 8 concentrations.

IgM antibodies specific for the serotype-defining surface lipoprotein appear after a few days of infection and soon reach a concentration that causes lysis of bacteria in the blood through either direct bactericidal action or opsonization. The release of large amounts of lipoproteins and other bacterial products from dying bacteria provokes a “crisis,” during which there can be an increase in temperature, hypotension, and other signs of shock. A similar phenomenon occurring in some patients soon after the initiation of antibiotic treatment is characterized by the abrupt worsening of the condition, which is called a Jarisch-Herxheimer reaction (JHR).

CLINICAL MANIFESTATIONS

Relapsing fever presents with the sudden onset of fever. Febrile periods are punctuated by intervening afebrile periods of a few days; this pattern occurs at least twice. The patient’s temperature is $\geq 39^{\circ}\text{C}$ and may be as high as 43°C . The first fever episode often ends in a crisis lasting ~15–30 min and consisting of rigors, a further elevation in temperature, and increases in pulse and blood pressure. The crisis phase is followed by profuse diaphoresis, falling temperature, and hypotension, which usually persist for several hours. In LBRF, the first episode of fever is unremitting for 3–6 days; it is usually followed by a single milder episode. In TBRF, multiple febrile periods last 1–3 days each. In both forms, the interval between fevers ranges from 4 to 14 days, sometimes with symptoms of malaise and fatigue.

The symptoms that accompany the fevers are usually nonspecific. Headache, neck stiffness, arthralgia, myalgia, and nausea may accompany the first and subsequent febrile episodes. An enlarging spleen and liver cause abdominal pain. A nonproductive cough is common during LBRF and—in combination with fever and myalgias—may suggest influenza. Acute respiratory distress syndrome may occur during TBRF.

On physical examination, the patient may be delirious or apathetic. There may be body lice in the patient’s clothes or signs of insect bites. In regions with *B. miyamotoi* infection, a hard tick may be embedded in the skin. Epistaxis, petechiae, and ecchymoses are common during LBRF but not in TBRF. Splenomegaly or spleen tenderness is common in both forms of relapsing fever. The majority of patients with LBRF and ~10% of patients with TBRF have discernible hepatomegaly.

Localizing neurologic findings are more common in TBRF than in LBRF. In North America, *B. turicatae* infection has neurologic manifestations more often than *B. hermsii* infection. Meningoencephalitis can result in residual hemiplegia or aphasia. Myelitis and radiculopathy

may develop. Unilateral or bilateral Bell’s palsy and deafness from seventh or eighth cranial nerve involvement are the most common forms of cranial neuritis and typically present in the second or third febrile episode, not the first. Visual impairment from unilateral or bilateral iridocyclitis or panophthalmitis may be permanent. In LBRF, neurologic manifestations such as altered mental state or stiff neck are thought to be secondary to spirochetemia and systemic inflammation rather than to direct invasion of the nervous system.

Myocarditis appears to be common in both forms of relapsing fever and accounts for some deaths. Most commonly, myocarditis is evidenced by gallops on cardiac auscultation, a prolonged QT_c interval, and cardiomegaly and pulmonary edema on chest radiography.

General laboratory studies are not specific. Mild to moderate normocytic anemia is common, but frank hemolysis and hemoglobinuria do not develop. Leukocyte counts are usually in the normal range or only slightly elevated, and leukopenia can occur during the crisis. Platelet counts can fall below 50,000/ μL . Laboratory evidence of hepatitis can be found, with elevated serum concentrations of unconjugated bilirubin and aminotransferases; the prothrombin and partial thromboplastin times may be moderately prolonged.

Analysis of the cerebrospinal fluid (CSF) is indicated in cases of suspected relapsing fever with signs of meningitis or meningoencephalitis. The presence of mononuclear pleocytosis and mildly to moderately elevated protein levels justifies intravenous antibiotic therapy in TBRF.

The manifestations and course of *B. miyamotoi* infection remain incompletely characterized, but reports to date indicate that the sign most often reported by patients at presentation is fever without respiratory symptoms starting 1–2 weeks after a tick bite and recurring once or twice in some cases. Meningoencephalitis with spirochetes in the CSF was documented in one immunodeficient adult.

DIAGNOSIS

Relapsing fever should be considered in a patient with the characteristic fever pattern and a history of recent exposure—i.e., within 1–2 weeks before illness onset—to body lice or soft-bodied ticks in geographic areas with documented current or past transmission. Because of the longevity of the ticks and the transovarial transmission of the pathogen in the ticks, a case of relapsing fever may be diagnosed many years after the last case reported in that locale.

The bedrock for laboratory diagnosis remains the same as it has been for a century: direct detection of the spirochetes by microscopy of the blood. Manual differential counts of white blood cells by Wright or Giemsa stain usually reveal spirochetes in thin blood smears if their concentration is $\geq 10^5/\text{mL}$ and several oil-immersion fields are examined (Fig. 209-2). The preferred time to obtain a blood specimen is between the fever’s onset and its peak. Lower concentrations of spirochetes may be revealed by a thick blood smear that is either directly stained with acridine orange and then examined by fluorescence microscopy or treated with 0.5% acetic acid before Giemsa or Wright staining. An alternative to a fixed blood smear is a wet mount of anticoagulated blood mixed with saline and examined by phase-contrast or dark-field microscopy for motile spirochetes.

Polymerase chain reaction (PCR) and similar DNA amplification procedures are increasingly used for examination of an extract of blood. PCR may reveal spirochetes between febrile episodes, since there are already escape variants in the population when the first wave of bacteria is neutralized.

Culture of blood or CSF in Barbour-Stoenner-Kelly broth medium is an option for isolation of *Borrelia* species except for *B. miyamotoi*, which is noncultivable or poorly cultivable. However, few laboratories offer this service. An alternative for tick-borne *Borrelia* species, including *B. miyamotoi*, is inoculation of blood or CSF into immunodeficient mice and examination of the animal’s blood after a few days.

Options for serologic confirmation of infection are limited. Most assays that are available commercially or in reference laboratories are based on whole cells of a single *Borrelia* species. These assays may not detect the major variant antigens to which the patient is mainly responding or may yield false-positive results due to antibodies to cross-reactive antigens of related bacteria, including *B. burgdorferi*. The most

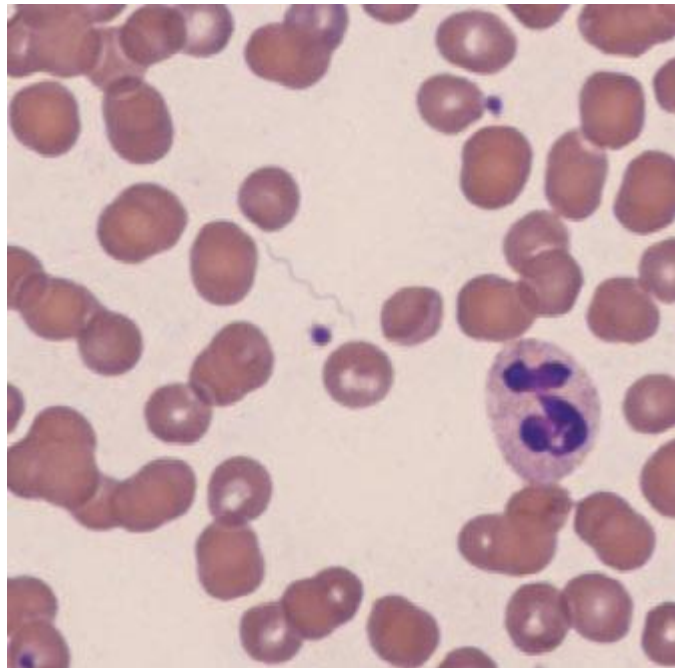


FIGURE 209-2 Photomicrograph of tick-borne relapsing fever spirochete (*Borrelia turicatae*) in a Wright-Giemsa-stained thin blood smear. Included in the figure are a polymorphonuclear leukocyte and two platelets.

promising new assays under development are based on recombinant antigens such as GlpQ, a protein antigen of all relapsing fever *Borrelia* species (including *B. miyamotoi*) but not of any Lyme disease species.

DIFFERENTIAL DIAGNOSIS

Depending on the patient's history of residential, occupational, travel, and recreational exposures, the differential diagnosis of relapsing fever includes one or more of the following infections that feature either periodicity in the fever pattern or an extended single febrile period with nonspecific constitutional symptoms: Colorado tick fever (which, along with dengue, can have a "saddleback" fever course), Rocky Mountain spotted fever and other rickettsioses, ehrlichiosis, anaplasmosis, tick-borne arbovirus infection, and babesiosis in North America, Europe, Russia, and northeastern Asia. Elsewhere in the Americas and Asia and in most of Africa, malaria, typhoid fever, typhus and other rickettsioses, dengue, brucellosis, and leptospirosis may also be considered. Depending on the geographic area and types of exposure, malaria, louse-borne typhus, typhoid fever, or Lyme disease may complicate relapsing fever.

TREATMENT RELAPSING FEVER

Penicillins and tetracyclines have been the antibiotics of choice for relapsing fever for several decades. Erythromycin has been a long-standing second choice. There is no evidence of acquired resistance to these antibiotics. *Borrelia* species are also susceptible to most cephalosporins and chloramphenicol, but there is less clinical experience with these drugs. Borreliae are relatively resistant to rifampin, sulfonamides, fluoroquinolones, and aminoglycosides. Spirochetes are no longer detectable in the blood within a few hours after the first dose of an effective antibiotic.

A single dose of antibiotic is usually sufficient for the treatment of LBRF (Fig. 209-3). The recurrence rate after antibiotic treatment is $\leq 5\%$. For adults, a single dose of oral tetracycline (500 mg), oral doxycycline (200 mg), or intramuscular penicillin G procaine (400,000–800,000 units) is effective. The corresponding doses for children are oral tetracycline at 12.5 mg/kg, oral doxycycline at 5 mg/kg, and intramuscular penicillin G procaine at 200,000–400,000 units.

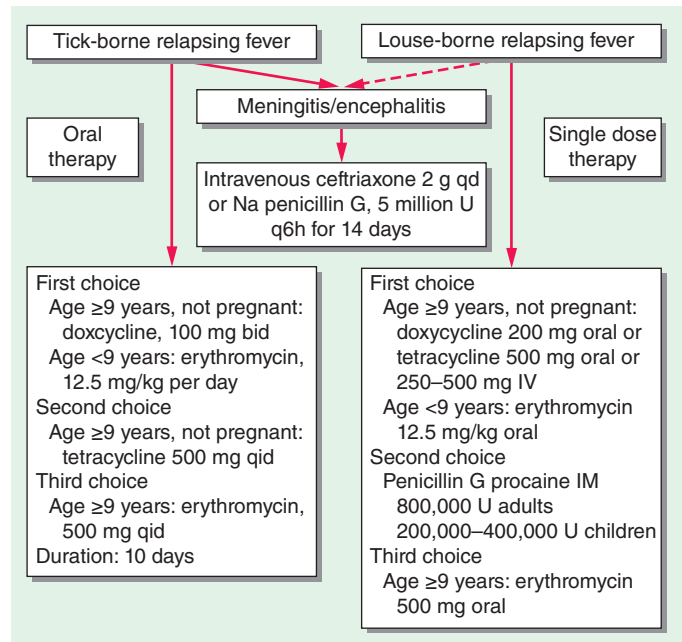


FIGURE 209-3 Algorithm for treatment of relapsing fever. If it is not known whether the patient has tick-borne or louse-borne relapsing fever, the patient should be treated for the tick-borne form. The dashed line indicates that central nervous system invasion in louse-borne relapsing fever is uncommon.

When an adult patient is stuporous or nauseated, the intravenous dose is 250–500 mg. Tetracycline is contraindicated in pregnant and nursing women and in children <9 years old; for individuals in these groups who are allergic to penicillin, oral erythromycin (500 mg for adults and 12.5 mg/kg for children) is an alternative. Tetracycline is marginally superior to penicillin G in terms of time to fever clearance and relapse rate.

The accumulated anecdotal reports on TBRF therapy indicate a recurrence rate of $\geq 20\%$ after single-dose treatment. This high rate of recurrence plausibly is due to the greater propensity of tick-borne species than of *B. recurrentis* to invade the CNS, from which they can reinvade the bloodstream after antibiotic levels decline. Accordingly, multiple antibiotic doses are recommended. The preferred treatment for adults is a 10-day course of tetracycline (500 mg or 12.5 mg/kg orally every 6 h) or doxycycline (100 mg twice daily). When tetracyclines are contraindicated, the alternative is erythromycin (500 mg or 12.5 mg/kg orally every 6 h) for 10 days. If a β -lactam antibiotic is given, it should be administered intravenously rather than orally, especially if CNS involvement is confirmed or suspected. For adults, the regimen is penicillin G (5 million units IV every 6 h) or ceftriaxone (2 g IV daily) for 10–14 days.

Experience with the treatment of *B. miyamotoi* infection is limited, but this organism likely has the same antibiotic susceptibilities as other *Borrelia* species. Until more is known about treatment efficacy, therapy for *B. miyamotoi* infection can follow the guidelines for Lyme disease—including parenteral therapy for CNS involvement—because it may be difficult to rule out co-infection.

The JHR during treatment of relapsing fever can be severe and can even end in death if precautions are not in place for close monitoring and provision of cardiovascular and volume support as needed. Rigors, fever, and hypotension occur within 2–3 h of initiation of antibiotic treatment. The incidence of the JHR is $\sim 80\%$ in LBRF and $\sim 50\%$ in TBRF. Both penicillin and tetracycline can elicit the JHR.

PROGNOSIS

The mortality rates for untreated LBRF and TBRF are in the ranges of 10–70% and 4–10%, respectively, and are largely determined by coexisting conditions, such as malnutrition and dehydration.

Death from untreated relapsing fever is most common during the first fever episode. With prompt antibiotic treatment, the mortality rate is 2–5% for LBRF and <2% for TBRF. Features associated with a poor prognosis include concurrence with malaria, typhus, or typhoid; pregnancy; stupor or coma on admission; diffuse bleeding; poor liver function; myocarditis; and bronchopneumonia. The mortality rate from the JHR in LBRF, in the absence of adequate monitoring and resuscitation measures, is ~5%. Some patients have survived the crisis or the JHR only to die suddenly either later that day or on the next day. Relapsing fever during pregnancy frequently leads to abortion or stillbirth, but congenital malformations have not been reported. Although it is possible that spirochetes may persist in the CNS or other sequestered sites after bacteremia has resolved, chronic disease or disability from a persistent infection has not been attributed to relapsing fever. Partial immunity against reinfection seems to develop in residents of endemic areas.

PREVENTION

There is no vaccine for either LBRF or TBRF. Reduction of exposure to lice and ticks is the key strategy for prevention. LBRF can be prevented through improved personal hygiene, reduction of crowding, better access to washing facilities, and selected use of pesticides. Infested clothing is an important factor in maintaining body lice. The risk of TBRF can be reduced by construction of houses with concrete or sealed plank floors and without thatched roofs or mud walls. Log cabins pose a particular risk in North America when rodents nest in the roof or beneath the house or porch. Interiors of buildings infested with *Ornithodoros* ticks can be treated with pesticides. If residing in a high-risk environment, individuals should not sleep on the floor, and beds should be moved away from the wall. With an exposure to TBRF, postexposure treatment with doxycycline (200 mg on day 1 followed by 100 mg/d for 4 days) was efficacious in preventing infection in a placebo-controlled trial.

210 Lyme Borreliosis

Allen C. Steere

DEFINITION

Lyme borreliosis is caused by a spirochete, *Borrelia burgdorferi sensu lato*, that is transmitted by ticks of the *Ixodes ricinus* complex. The infection usually begins with a characteristic expanding skin lesion, erythema migrans (EM; stage 1, localized infection). After several days or weeks, the spirochete may spread to many different sites (stage 2, disseminated infection). Possible manifestations of disseminated infection include secondary annular skin lesions, meningitis, cranial neuritis, radiculoneuritis, peripheral neuritis, carditis, atrioventricular nodal block, or migratory musculoskeletal pain. Months or years later (usually after periods of latent infection), intermittent or persistent arthritis, chronic encephalopathy or polyneuropathy, or acrodermatitis may develop (stage 3, persistent infection). Most patients experience early symptoms of the illness during the summer, but the infection may not become symptomatic until it progresses to stage 2 or 3.

Lyme disease was recognized as a separate entity in 1976 because of a geographic cluster of children in Lyme, Connecticut, who were thought to have juvenile rheumatoid arthritis. It became apparent that Lyme disease was a multisystemic illness that affected primarily the skin, nervous system, heart, and joints. Epidemiologic studies of patients with EM implicated certain *Ixodes* ticks as vectors of the disease. Early in the twentieth century, EM had been described in Europe and attributed to *I. ricinus* tick bites. In 1982, a previously unrecognized spirochete, now called *Borrelia burgdorferi*, was recovered from *Ixodes scapularis* ticks and then from patients with Lyme disease. The entity is now called Lyme disease or Lyme borreliosis.

ETIOLOGIC AGENT



B. burgdorferi, the causative agent of Lyme disease, is a fastidious microaerophilic bacterium. The spirochete's genome is quite small (~1.5 Mb) and consists of a highly unusual linear chromosome of 950 kb as well as 17–21 linear and circular plasmids. The most remarkable aspect of the *B. burgdorferi* genome is that there are sequences for more than 100 known or predicted lipoproteins—a larger number than in any other organism. The spirochete has few proteins with biosynthetic activity and depends on its host for most of its nutritional requirements. It has no sequences for recognizable toxins.



Currently, 13 closely related borrelial species are collectively referred to as *Borrelia burgdorferi sensu lato* (i.e., “*B. burgdorferi* in the general sense”). The human infection Lyme borreliosis is caused primarily by three pathogenic genospecies: *B. burgdorferi sensu stricto* (“*B. burgdorferi* in the strict sense,” hereafter referred to simply as *B. burgdorferi*), *Borrelia garinii*, and *Borrelia afzelii*. *B. burgdorferi* is the sole cause of the infection in the United States; all three genospecies are found in Europe, and the latter two species occur in Asia.

Strains of *B. burgdorferi* have been subdivided according to several typing schemes: one based on sequence variation of outer-surface protein C (OspC), a second based on differences in the 16S–23S rRNA intergenic spacer region (RST or IGS), and a third called *multilocus sequence typing*. From these typing systems, it is apparent that strains of *B. burgdorferi* differ in pathogenicity. OspC type A (RST1) strains seem to be particularly virulent and may have played a role in the emergence of Lyme disease in epidemic form in the northeastern United States in the late twentieth century.

EPIDEMIOLOGY



The 13 known genospecies of *B. burgdorferi sensu lato* live in nature in enzootic cycles involving 14 species of ticks that are part of the *I. ricinus* complex. *I. scapularis* (Fig. 475-1) is the principal vector in the eastern United States from Maine to Georgia and in the midwestern states of Wisconsin, Minnesota, and Michigan. *I. pacificus* is the vector in the western states of California and Oregon. The disease is acquired throughout Europe (from Great Britain to Scandinavia to European Russia), where *I. ricinus* is the vector, and in Asian Russia, China, and Japan, where *I. persulcatus* is the vector. These ticks may transmit other agents as well. In the United States, *I. scapularis* also transmits *Babesia microti*; *Anaplasma phagocytophilum*; *Ehrlichia* species Wisconsin; *Borrelia miyamotoi*; and, in rare instances, Powassan encephalitis virus (the deer tick virus) (see “Differential Diagnosis,” below). In Europe and Asia, *I. ricinus* and *I. persulcatus* also transmit tick-borne encephalitis virus.

Ticks of the *I. ricinus* complex have larval, nymphal, and adult stages. They require a blood meal at each stage. The risk of infection in a given area depends largely on the density of these ticks as well as their feeding habits and animal hosts, which have evolved differently in different locations. For *I. scapularis* in the northeastern United States, the white-footed mouse and certain other rodents are the preferred hosts of the immature larvae and nymphs. It is critical that both of the tick's immature stages feed on the same host because the life cycle of the spirochete depends on horizontal transmission: in early summer from infected nymphs to mice and in late summer from infected mice to larvae, which then molt to become the infected nymphs that will begin the cycle again the following year. It is the tiny nymphal tick that is primarily responsible for transmission of the disease to humans during the early summer months. White-tailed deer, which are not involved in the life cycle of the spirochete, are the preferred host for the adult stage of *I. scapularis* and seem to be critical to the tick's survival.

Lyme disease is now the most common vector-borne infection in the United States and Europe. Since surveillance was begun by the Centers for Disease Control and Prevention (CDC) in 1982, the number of cases in the United States has increased dramatically. More than 30,000 new cases are now reported each summer, but the actual number of new cases is probably closer to 300,000 annually. In Europe, reported frequencies of the disease are highest in the middle of the continent and in Scandinavia.

To maintain its complex enzootic cycle, *B. burgdorferi* must adapt to two markedly different environments: the tick and the mammalian host. The spirochete expresses outer-surface protein A (OspA) in the midgut of the tick, whereas OspC is upregulated as the organism travels to the tick's salivary gland. There, OspC binds a tick salivary-gland protein (Salp15), which is required for infection of the mammalian host. The tick usually must be attached for at least 24 h for transmission of *B. burgdorferi*.

After injection into the human skin, the spirochete downregulates OspC and upregulates the VlsE lipoprotein. This protein undergoes extensive antigenic variation, which is necessary for spirochetal survival. After several days to weeks, *B. burgdorferi* may migrate outward in the skin, producing EM, and may spread hematogenously or in the lymph to other organs. The only known virulence factors of *B. burgdorferi* are surface proteins that allow the spirochete to attach to mammalian proteins, integrins, glycosaminoglycans, or glycoproteins. For example, spread through the skin and other tissue matrices may be facilitated by the binding of human plasminogen and its activators to the surface of the spirochete. Some *Borrelia* strains bind complement regulator-acquiring surface proteins (FHL-1/reconnectin, or factor H), which help to protect spirochetes from complement-mediated lysis. Dissemination of the organism in the blood is facilitated by binding to the fibrinogen receptor on activated platelets ($\alpha_{IIb}\beta_3$) and the vitronectin receptor ($\alpha_v\beta_3$) on endothelial cells. As the name indicates, spirochetal decorin-binding proteins A and B bind decorin, a glycosaminoglycan on collagen fibrils; this binding may explain why the organism is commonly aligned with collagen fibrils in the extracellular matrix in the heart, nervous system, or joints.

To control and eradicate *B. burgdorferi*, the host mounts both innate and adaptive immune responses, resulting in macrophage- and antibody-mediated killing of the spirochete. As part of the innate immune response, complement may lyse the spirochete in the skin. Cells at affected sites release potent proinflammatory cytokines, including interleukin 6, tumor necrosis factor α , interleukin 1 β , and interferon γ . Patients who are homozygous for a Toll-like receptor 1 polymorphism (1805GG), particularly when infected with highly inflammatory *B. burgdorferi* RST1 strains, have exceptionally high levels of proinflammatory cytokines. The purpose of the adaptive immune response appears to be the production of specific antibodies, which opsonize the organism—a step necessary for optimal spirochetal killing. Studies with protein arrays expressing ~1200 *B. burgdorferi* proteins detected antibody responses to a total of 120 spirochetal proteins (particularly outer-surface lipoproteins) in a population of patients with Lyme arthritis. Histologic examination of all affected tissues reveals an infiltration of lymphocytes, macrophages, and plasma cells with some degree of vascular damage, including mild vasculitis or hypervascular occlusion. These findings suggest that the spirochete may have been present in or around blood vessels.

In enzootic infection, *B. burgdorferi* spirochetes must survive this immune assault only during the summer months before returning to larval ticks to begin the cycle again the following year. In contrast, infection of humans is a dead-end event for the spirochete. Within several weeks or months, innate and adaptive immune mechanisms—even without antibiotic treatment—control widely disseminated infection, and generalized systemic symptoms wane. However, without antibiotic therapy, spirochetes may survive in localized niches for several more years. For example, *B. burgdorferi* infection in the United States may cause persistent arthritis or, in rare cases, subtle encephalopathy or polyneuropathy. Thus, immune mechanisms seem to succeed eventually in the near or total eradication of *B. burgdorferi* from selected niches, including the joints or nervous system, and symptoms resolve in most patients.

CLINICAL MANIFESTATIONS

Early Infection: Stage 1 (Localized Infection) Because of the small size of nymphal ixodid ticks, most patients do not remember the preceding tick bite. After an incubation period of 3–32 days, EM usually begins as a red macule or papule at the site of the tick bite that expands slowly to



FIGURE 210-1 A classic erythema migrans lesion (9 cm in diameter) is shown near the right axilla. The lesion has partial central clearing, a bright red outer border, and a target center. (Courtesy of Vijay K. Sikand, MD; with permission.)

form a large annular lesion (Fig. 210-1). As the lesion increases in size, it often develops a bright red outer border and partial central clearing. The center of the lesion sometimes becomes intensely erythematous and indurated, vesicular, or necrotic. In other instances, the expanding lesion remains an even, intense red; several red rings are found within an outside ring; or the central area turns blue before the lesion clears. Although EM can be located anywhere, the thigh, groin, and axilla are particularly common sites. The lesion is warm but not often painful. Approximately 20% of patients do not exhibit this characteristic skin manifestation.

Early Infection: Stage 2 (Disseminated Infection) In cases in the United States, *B. burgdorferi* often spreads hematogenously to many sites within days or weeks after the onset of EM. In these cases, patients may develop secondary annular skin lesions similar in appearance to the initial lesion. Skin involvement is commonly accompanied by severe headache, mild stiffness of the neck, fever, chills, migratory musculoskeletal pain, arthralgias, and profound malaise and fatigue. Less common manifestations include generalized lymphadenopathy or splenomegaly, hepatitis, sore throat, nonproductive cough, conjunctivitis, iritis, or testicular swelling. Except for fatigue and lethargy, which are often constant, the early signs and symptoms of Lyme disease are typically intermittent and changing. Even in untreated patients, the early symptoms usually become less severe or disappear within several weeks. In ~15% of patients, the infection presents with these nonspecific systemic symptoms.

Symptoms suggestive of meningeal irritation may develop early in Lyme disease when EM is present but usually are not associated with cerebrospinal fluid (CSF) pleocytosis or an objective neurologic deficit. After several weeks or months, ~15% of untreated patients develop frank neurologic abnormalities, including meningitis, subtle encephalitic signs, cranial neuritis (including bilateral facial palsy), motor or sensory radiculoneuropathy, peripheral neuropathy, mononeuritis multiplex, cerebellar ataxia, or myelitis—alone or in various combinations. In children, the optic nerve may be affected because of inflammation or increased intracranial pressure, and these effects may lead to blindness. In the United States, the usual pattern consists of fluctuating symptoms of meningitis accompanied by facial palsy and peripheral radiculoneuropathy. Lymphocytic pleocytosis (~100 cells/ μ L) is found in CSF, often along with elevated protein levels and normal or slightly low glucose concentrations. In Europe and Asia, the first neurologic sign is characteristically radicular pain, which is followed by the development of CSF pleocytosis (meningopolyneuritis or *Bannwarth's syndrome*); meningeal or encephalitic signs are frequently absent. These early neurologic abnormalities usually resolve completely within months, but in rare cases chronic neurologic disease may occur later.

Within several weeks after the onset of illness, ~8% of patients develop cardiac involvement. The most common abnormality is a

fluctuating degree of atrioventricular block (first-degree, Wenckebach, or complete heart block). Some patients have more diffuse cardiac involvement, including electrocardiographic changes indicative of acute myopericarditis, left ventricular dysfunction evident on radionuclide scans, or (in rare cases) cardiomegaly or fatal pancarditis. Cardiac involvement lasts for only a few weeks in most patients but may recur in untreated patients. Chronic cardiomyopathy caused by *B. burgdorferi* has been reported in Europe.

During this stage, musculoskeletal pain is common. The typical pattern consists of migratory pain in joints, tendons, bursae, muscles, or bones (usually without joint swelling) lasting for hours or days and affecting one or two locations at a time.

Late Infection: Stage 3 (Persistent Infection) Months after the onset of infection, ~60% of patients in the United States who have received no antibiotic treatment develop frank arthritis. The typical pattern comprises intermittent attacks of oligoarticular arthritis in large joints (especially the knees), lasting for weeks or months in a given joint. A few small joints or periarticular sites also may be affected, primarily during early attacks. The number of patients who continue to have recurrent attacks decreases each year. However, in a small percentage of cases, involvement of large joints—usually one or both knees—is persistent and may lead to erosion of cartilage and bone.

White cell counts in joint fluid range from 500 to 110,000/ μ L (average, 25,000/ μ L); most of these cells are polymorphonuclear leukocytes. Tests for rheumatoid factor or antinuclear antibodies usually give negative results. Examination of synovial biopsy samples reveals fibrin deposits, villous hypertrophy, vascular proliferation, microangiopathic lesions, and a heavy infiltration of lymphocytes and plasma cells.

Although most patients with Lyme arthritis respond well to antibiotic therapy, a small percentage in the northeastern United States have persistent (*antibiotic-refractory*) arthritis for months or even for several years after receiving oral and IV antibiotic therapy for 2 or 3 months. Although more often these patients are initially infected with RST1 strains of *B. burgdorferi*, this complication is not thought to result from persistent infection. Results of culture and polymerase chain reaction (PCR) for *B. burgdorferi* in synovial tissue obtained in the postantibiotic period have been uniformly negative. Rather, infection-induced autoimmunity, retained spirochetal antigens, or both may play a role in this outcome. Antibiotic-refractory arthritis is associated with a higher frequency of certain class II major histocompatibility complex molecules (particularly HLA-DRB1*0401 or -*0101 molecules); the Toll-like receptor 1 polymorphism 1805GG, which leads to exceptionally high levels of cytokines and chemokines in affected joints; and low frequencies of FoxP3+ T regulatory cells in synovial fluid, which correlate with longer posttreatment durations of arthritis. The recent identification of a novel human autoantigen, endothelial cell growth factor, as a target of T and B cell responses in patients with Lyme disease provided the first direct evidence of autoimmune T and B cell responses in this illness. However, multiple spirochetal or additional yet-to-be identified autoantigens may have a role in antibiotic-refractory arthritis.

Although rare, chronic neurologic involvement also may become apparent from months to several years after the onset of infection, sometimes after long periods of latent infection. The most common form of chronic central nervous system involvement is subtle encephalopathy affecting memory, mood, or sleep, and the most common form of peripheral neuropathy is an axonal polyneuropathy manifested as either distal paresthesia or spinal radicular pain. Patients with encephalopathy frequently have evidence of memory impairment in neuropsychological tests and abnormal results in CSF analyses. In cases of polyneuropathy, electromyography generally shows extensive abnormalities of proximal and distal nerve segments. Encephalomyelitis or leukoencephalitis, a rare manifestation of Lyme borreliosis associated primarily with *B. garinii* infection in Europe, is a severe neurologic disorder that may include spastic paraparesis, upper motor-neuron bladder dysfunction, and, rarely, lesions in the periventricular white matter.



Acrodermatitis chronica atrophicans, the late skin manifestation of Lyme borreliosis, has been associated primarily with *B. afzelii* infection in Europe and Asia. It has been observed

especially often in elderly women. The skin lesions, which are usually found on the acral surface of an arm or leg, begin insidiously with reddish-violaceous discoloration; they become sclerotic or atrophic over a period of years.

The basic patterns of Lyme borreliosis are similar worldwide, but there are regional variations, primarily between the illness found in North America, which is caused exclusively by *B. burgdorferi*, and that found in Europe, which is caused primarily by *B. afzelii* and *B. garinii*. With each of the *Borrelia* species, the infection usually begins with EM. However, *B. burgdorferi* strains in the eastern United States often disseminate widely; they are particularly arthritogenic, and they may cause antibiotic-refractory arthritis. *B. garinii* typically disseminates less widely, but it is especially neurotropic and may cause borreliac encephalomyelitis. *B. afzelii* often infects only the skin but may persist in that site, where it may cause several different dermatoborrelioses, including acrodermatitis chronica atrophicans.

Post-Lyme Syndrome (Chronic Lyme Disease) Despite resolution of the objective manifestations of the infection with antibiotic therapy, ~10% of patients (although the reported percentages vary widely) continue to have subjective pain, neurocognitive manifestations, or fatigue symptoms. These symptoms usually improve and resolve within months but may last for years. At the far end of the spectrum, the symptoms may be similar to or indistinguishable from chronic fatigue syndrome ([Chap. 464e](#)) and fibromyalgia ([Chap. 396](#)). Compared with symptoms of active Lyme disease, post-Lyme symptoms tend to be more generalized or disabling. They include marked fatigue, severe headache, diffuse musculoskeletal pain, multiple symmetric tender points in characteristic locations, pain and stiffness in many joints, diffuse paresthesias, difficulty with concentration, and sleep disturbances. Patients with this condition lack evidence of joint inflammation, have normal neurologic test results, and may exhibit anxiety and depression. In contrast, late manifestations of Lyme disease, including arthritis, encephalopathy, and neuropathy, are usually associated with minimal systemic symptoms. Currently, no evidence indicates that persistent subjective symptoms after recommended courses of antibiotic therapy are caused by active infection.

DIAGNOSIS

The culture of *B. burgdorferi* in Barbour-Stoenner-Kelly (BSK) medium permits definitive diagnosis, but this method has been used primarily in research studies. Moreover, with a few exceptions, positive cultures have been obtained only early in the illness—particularly from biopsy samples of EM skin lesions, less often from plasma samples, and occasionally from CSF samples. Later in the infection, PCR is greatly superior to culture for the detection of *B. burgdorferi* DNA in joint fluid; this is the major use for PCR testing in Lyme disease. However, because *B. burgdorferi* DNA may persist for at least weeks after spirochetal killing with antibiotics, detection of spirochetal DNA in joint fluid is not an accurate test of active joint infection in Lyme disease and cannot be used reliably to determine the adequacy of antibiotic therapy. The sensitivity of PCR determinations in CSF from patients with neuroborreliosis has been much lower than that in joint fluid. There seems to be little if any role for PCR in the detection of *B. burgdorferi* DNA in blood or urine samples. Moreover, this procedure must be carefully controlled to prevent contamination.

Because of the problems associated with direct detection of *B. burgdorferi*, Lyme disease is usually diagnosed by the recognition of a characteristic clinical picture accompanied by serologic confirmation. Although serologic testing may yield negative results during the first several weeks of infection, almost all patients have a positive antibody response to *B. burgdorferi* after that time. The limitation of serologic tests is that they do not clearly distinguish between active and inactive infection. Patients with previous Lyme disease—particularly in cases progressing to late stages—often remain seropositive for years, even after adequate antibiotic treatment. In addition, ~10% of patients are seropositive because of asymptomatic infection. If these individuals subsequently develop another illness, the positive serologic test for

TABLE 210-1 ALGORITHM FOR TESTING FOR AND TREATING LYME DISEASE

Pretest Probability	Example	Recommendation
High	Patients with erythema migrans	Empirical antibiotic treatment without serologic testing
Intermediate	Patients with oligoarticular arthritis	Serologic testing and antibiotic treatment if test results are positive
Low	Patients with nonspecific symptoms (myalgias, arthralgias, fatigue)	Neither serologic testing nor antibiotic treatment

Source: Adapted from the recommendations of the American College of Physicians (G Nichol et al: *Ann Intern Med* 128:37, 1998, with permission).

Lyme disease may cause diagnostic confusion. According to an algorithm published by the American College of Physicians (Table 210-1), serologic testing for Lyme disease is recommended only for patients with at least an intermediate pretest probability of Lyme disease, such as those with oligoarticular arthritis. It should not be used as a screening procedure in patients with pain or fatigue syndromes. In such patients, the probability of a false-positive serologic result is higher than that of a true-positive result.

For serologic analysis of Lyme disease in the United States, the CDC recommends a two-step approach in which samples are first tested by enzyme-linked immunosorbent assay (ELISA) and equivocal or positive results are then tested by western blotting. During the first weeks of infection, both IgM and IgG responses to the spirochete should be determined, preferably in both acute- and convalescent-phase serum samples. Approximately 20–30% of patients have a positive response detectable in acute-phase samples, whereas ~70–80% have a positive response during convalescence (2–4 weeks later). After 4–8 weeks of infection (by which time most patients with active Lyme disease have disseminated infection), the sensitivity and specificity of the IgG response to the spirochete are both very high—in the range of 99%—as determined by the two-test approach of ELISA and western blot. At this point and thereafter, a single test (that for IgG) is usually sufficient. In persons with illness of >2 months' duration, a positive IgM test result alone is likely to be false-positive and therefore should not be used to support the diagnosis.

According to current criteria adopted by the CDC, an IgM western blot is considered positive if two of the following three bands are present: 23, 39, and 41 kDa. However, the combination of two such bands may still represent a false-positive result. Misuse or misinterpretation of IgM blots has been a factor in the incorrect diagnosis of Lyme disease in patients with other illnesses. An IgG blot is considered positive if 5 of the following 10 bands are present: 18, 23, 28, 30, 39, 41, 45, 58, 66, and 93 kDa. In European cases, no single set of criteria for the interpretation of immunoblots results in high levels of sensitivity and specificity in all countries.

The most promising second-generation serologic test is the VlsE C6 peptide IgG ELISA, which employs a 26-mer of the sixth invariant region of the VlsE lipoprotein of *B. burgdorferi*. The results achieved with this test are similar to those obtained with the standard two-test approach (sonicate IgM and IgG ELISA and western blot). The principal advantage of the C6 peptide ELISA is the early detection of an IgG response, which renders an IgM test unnecessary. However, not all patients with late Lyme disease have a response to the C6 peptide, and this test is not quite as specific as sonicate western blot. Thus, at present, a two-test approach that includes western blot is still recommended. Blotting can also be helpful in assessing the duration of current or past disease.

After successful antibiotic treatment, antibody titers decline slowly but responses (including that to the VlsE C6 peptide) may persist for years. Moreover, not only the IgG but also the IgM response may persist for years after therapy. Therefore, even a positive IgM response cannot be interpreted as confirmation of recent infection or reinfection unless the clinical picture is appropriate.

DIFFERENTIAL DIAGNOSIS

Classic EM is a slowly expanding erythema, often with partial central clearing. If the lesion expands little, it may represent the red papule of an uninfected tick bite. If the lesion expands rapidly, it may represent cellulitis (e.g., streptococcal cellulitis) or an allergic reaction, perhaps to tick saliva. Patients with secondary annular lesions may be thought to have erythema multiforme, but neither the development of blistering mucosal lesions nor the involvement of the palms or soles is a feature of *B. burgdorferi* infection. In the eastern United States, an EM-like skin lesion, sometimes with mild systemic symptoms, may be associated with *Amblyomma americanum* tick bites. However, the cause of this Southern tick-associated rash illness (STARI) has not yet been identified. This tick may also transmit *Ehrlichia chaffeensis*, a rickettsial agent (Chap. 211).

As stated above, *I. scapularis* ticks in the United States may transmit not only *B. burgdorferi* but also *B. microti*, a red blood cell parasite (Chap. 249); *A. phagocytophilum*, the agent of human granulocytotropic anaplasmosis (Chap. 211); *Ehrlichia* species Wisconsin; *B. miyamotoi*, a relapsing fever spirochete (Chap. 209); or (rarely) Powassan encephalitis virus (the deer tick virus, which is closely related to European tick-borne encephalitis virus) (Chap. 233). Although babesiosis and anaplasmosis are most often asymptomatic, infection with any of these agents may cause nonspecific systemic symptoms, particularly in the young or elderly, and co-infected patients may have more severe or persistent symptoms than patients infected with a single agent. Standard blood counts may yield clues regarding the presence of co-infection with *Anaplasma* or *Babesia*. Anaplasmosis may cause leukopenia or thrombocytopenia, and babesiosis may cause thrombocytopenia or (in severe cases) hemolytic anemia. IgM serologic responses may confuse the diagnosis. For example, *A. phagocytophilum* may elicit a positive IgM response to *B. burgdorferi*. The frequency of co-infection in different studies has been variable. In one prospective study, 4% of patients with EM had evidence of co-infection.

Facial palsy caused by *B. burgdorferi*, which occurs in the early disseminated phase of the infection (often in July, August, or September), is usually recognized by its association with EM. However, in rare cases, facial palsy without EM may be the presenting manifestation of Lyme disease. In such cases, both the IgM and the IgG responses to the spirochete are usually positive. The most common infectious agents that cause facial palsy are herpes simplex virus type 1 (Bell's palsy; Chap. 216) and varicella-zoster virus (Ramsay Hunt syndrome; Chap. 217).

Later in the infection, oligoarticular Lyme arthritis most resembles reactive arthritis in an adult or the pauciarticular form of juvenile idiopathic arthritis in a child. Patients with Lyme arthritis usually have the strongest IgG antibody responses seen in Lyme borreliosis, with reactivity to many spirochetal proteins.

The most common problem in diagnosis is to mistake Lyme disease for chronic fatigue syndrome (Chap. 464e) or fibromyalgia (Chap. 396). This difficulty is compounded by the fact that a small percentage of patients do in fact develop these chronic pain or fatigue syndromes in association with or soon after Lyme disease. Moreover, a counter culture has emerged that ascribes pain and fatigue syndromes to chronic Lyme disease when there is little or no evidence of *B. burgdorferi* infection. In such cases, the term *chronic Lyme disease*, which is equated with chronic *B. burgdorferi* infection, is a misnomer, and the use of prolonged, dangerous, and expensive antibiotic treatment is not warranted.

TREATMENT LYME BORRELIOSIS

ANTIBIOTIC TREATMENT

As outlined in the algorithm in Fig. 210-2, the various manifestations of Lyme disease can usually be treated successfully with orally administered antibiotics; the exceptions are objective neurologic abnormalities and third-degree atrioventricular heart block, which are generally treated with IV antibiotics, and arthritis that does not

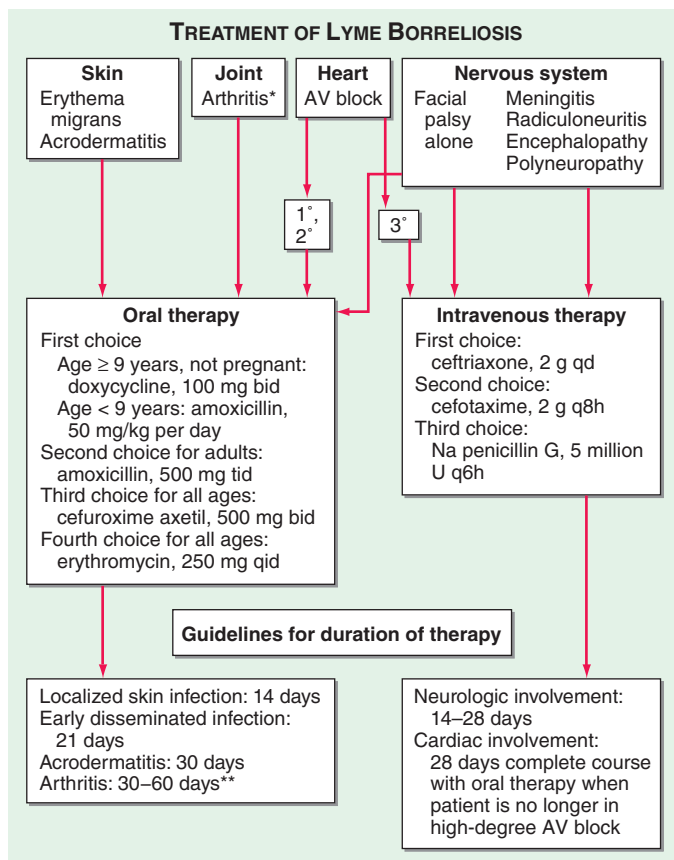


FIGURE 210-2 Algorithm for the treatment of the various early or late manifestations of Lyme borreliosis. AV, atrioventricular. *For arthritis, oral therapy should be tried first; if arthritis is unresponsive, IV therapy should be administered. **For Lyme arthritis, IV ceftriaxone (2 g given once a day for 14–28 days) also is effective and is necessary for a small percentage of patients; however, compared with oral treatment, this regimen is less convenient to administer, has more side effects, and is more expensive.

respond to therapy. For early Lyme disease, doxycycline is effective and can be administered to men and nonpregnant women. An advantage of this regimen is that it is also effective against *A. phagocytophilum*, which is transmitted by the same tick that transmits the Lyme disease agent. Amoxicillin, cefuroxime axetil, and erythromycin or its congeners are second-, third-, and fourth-choice alternatives, respectively. In children, amoxicillin is effective (not more than 2 g/d); in cases of penicillin allergy, cefuroxime axetil or erythromycin may be used. In contrast to second- or third-generation cephalosporin antibiotics, first-generation cephalosporins, such as cephalexin, are not effective. For patients with infection localized to the skin, a 14-day course of therapy is generally sufficient; in contrast, for patients with disseminated infection, a 21-day course is recommended. Approximately 15% of patients experience a Jarisch-Herxheimer-like reaction during the first 24 h of therapy. In multicenter studies, more than 90% of patients whose early Lyme disease was treated with these regimens had satisfactory outcomes. Although some patients reported symptoms after treatment, objective evidence of persistent infection or relapse was rare, and re-treatment was usually unnecessary.

Oral administration of doxycycline or amoxicillin for 30 days is recommended for the initial treatment of Lyme arthritis in patients who do not have concomitant neurologic involvement. Among patients with arthritis who do not respond to oral antibiotics, re-treatment with IV ceftriaxone for 28 days is appropriate. In patients with arthritis in whom joint inflammation persists for months or even several years after both oral and IV antibiotics, treatment

with nonsteroidal anti-inflammatory agents, therapy with disease-modifying antirheumatic drugs, or synovectomy may be successful.

In the United States, parenteral antibiotic therapy is usually used for objective neurologic abnormalities (with the exception of facial palsy alone). Patients with neurologic involvement are most commonly treated with IV ceftriaxone for 14–28 days, but IV cefotaxime or IV penicillin G for the same duration also may be effective. In Europe, similar results have been obtained with oral doxycycline and IV antibiotics in the treatment of acute neuroborreliosis. In patients with high-degree atrioventricular block or a PR interval of >0.3 s, IV therapy for at least part of the course and cardiac monitoring are recommended, but the insertion of a permanent pacemaker is not necessary.

It is unclear how and whether asymptomatic infection should be treated, but patients with such infection are often given a course of oral antibiotics. Because maternal-fetal transmission of *B. burgdorferi* seems to occur rarely (if at all), standard therapy for the manifestations of the illness is recommended for pregnant women. Long-term persistence of *B. burgdorferi* has not been documented in any large series of patients after treatment with currently recommended regimens. Although an occasional patient requires a second course of antibiotics, there is no indication for multiple, repeated antibiotic courses in the treatment of Lyme disease.

CHRONIC LYME DISEASE

After appropriately treated Lyme disease, a small percentage of patients continue to have subjective symptoms, primarily musculoskeletal pain, neurocognitive difficulties, or fatigue. This *chronic Lyme disease* or *post-Lyme syndrome* is sometimes a disabling condition that is similar to chronic fatigue syndrome or fibromyalgia. In a large study, one group of patients with post-Lyme syndrome received IV ceftriaxone for 30 days followed by oral doxycycline for 60 days, while another group received IV and oral placebo preparations for the same durations. No significant differences were found between groups in the numbers of patients reporting that their symptoms had improved, become worse, or stayed the same. Such patients are best treated for the relief of symptoms rather than with prolonged courses of antibiotics.

PROPHYLAXIS AFTER A TICK BITE

The risk of infection with *B. burgdorferi* after a recognized tick bite is so low that antibiotic prophylaxis is not routinely indicated. However, if an attached, engorged *I. scapularis* nymph is found or if follow-up is anticipated to be difficult, a single 200-mg dose of doxycycline, which usually prevents Lyme disease when given within 72 h after the tick bite, may be administered.

PROGNOSIS

The response to treatment is best early in the disease. Later treatment of Lyme borreliosis is still effective, but the period of convalescence may be longer. Eventually, most patients recover with minimal or no residual deficits.

REINFECTION

Reinfection may occur after EM when patients are treated with antimicrobial agents. In such cases, the immune response is not adequate to provide protection from subsequent infection. However, patients who develop an expanded immune response to the spirochete over a period of months (e.g., those with Lyme arthritis) have protective immunity for a period of years and rarely, if ever, acquire the infection again.

PREVENTION

Protective measures for the prevention of Lyme disease may include the avoidance of tick-infested areas, the use of repellents and acaricides, tick checks, and modification of landscapes in or near residential areas. Although a vaccine for Lyme disease used to be available, the manufacturer has discontinued its production. Therefore, no vaccine is now commercially available for the prevention of this infection.

211

Rickettsial Diseases

David H. Walker, J. Stephen Dumler, Thomas Marrie

The rickettsiae are a heterogeneous group of small, obligately intracellular, gram-negative coccobacilli and short bacilli, most of which are transmitted by a tick, mite, flea, or louse vector. Except in the case of louse-borne typhus, humans are incidental hosts. Among rickettsiae, *Coxiella burnetii*, *Rickettsia prowazekii*, and *R. typhi* have the well-documented ability to survive for an extended period outside the reservoir or vector and to be extremely infectious: inhalation of a single *Coxiella* microorganism can cause pneumonia. High-level infectivity and severe illness after inhalation make *R. prowazekii*, *R. rickettsii*, *R. typhi*, *R. conorii*, and *C. burnetii* bioterrorism threats.

Clinical infections with rickettsiae can be classified according to (1) the taxonomy and diverse microbial characteristics of the agents, which belong to seven genera (*Rickettsia*, *Orientia*, *Ehrlichia*, *Anaplasma*, *Neorickettsia*, *Candidatus Neoehrlichia*, and *Coxiella*); (2) epidemiology; or (3) clinical manifestations. The clinical manifestations of all the acute presentations are similar during the first 5 days: fever, headache, and myalgias with or without nausea, vomiting, and cough. As the course progresses, clinical manifestations—including occurrence of a macular, maculopapular, or vesicular rash; eschar; pneumonitis; and meningoenzephalitis—vary from one disease to another. Given the 15 etiologic agents with varied mechanisms of transmission, geographic distributions, and associated disease manifestations, the consideration of rickettsial diseases as a single entity poses complex challenges (Table 211-1).

Establishing the etiologic diagnosis of rickettsioses is very difficult during the acute stage of illness, and definitive diagnosis usually requires the examination of paired serum samples after convalescence. Heightened clinical suspicion is based on epidemiologic data, history of exposure to vectors or reservoir animals, travel to endemic locations, clinical manifestations (sometimes including rash or eschar), and characteristic laboratory findings (including thrombocytopenia, normal or low white blood cell [WBC] counts, elevated hepatic enzyme levels, and hyponatremia). Such suspicion should prompt empirical treatment. Doxycycline is the drug of choice for most of these infections. Only one agent, *C. burnetii*, has been documented to cause chronic illness. No other species, *R. prowazekii*, causes recrudescent illness (Brill-Zinsser disease) when latent infection is reactivated years after resolution of the acute illness.

Rickettsial infections dominated by fever may resolve without further clinical evolution. However, after nonspecific early manifestations, the illnesses can also evolve along one or more of several principal clinical lines: (1) development of a macular or maculopapular rash; (2) development of an eschar at the site of tick or mite feeding; (3) development of a vesicular rash (often in rickettsialpox and African tick-bite fever); (4) development of pneumonitis with chest radiographic opacities and/or rales (Q fever and severe cases of Rocky Mountain spotted fever [RMSF], Mediterranean spotted fever [MSF], louse-borne typhus, human monocytotropic ehrlichiosis [HME], human granulocytotropic anaplasmosis [HGA], scrub typhus, and murine typhus); (5) development of meningoenzephalitis (louse-borne typhus and severe cases of RMSF, scrub typhus, HME, murine typhus, MSF, and [rarely] Q fever); and (6) progressive hypotension and multiorgan failure as seen with sepsis or toxic shock syndromes (RMSF, MSF, louse-borne typhus, murine typhus, scrub typhus, HME, HGA, and neohrlichiosis).

Epidemiologic clues to the transmission of a particular pathogen include (1) environmental exposure to ticks, fleas, or mites during the season of activity of the vector species for the disease in the appropriate geographic region (spotted fever and typhus rickettsioses, scrub typhus, ehrlichioses, anaplasmosis); (2) travel to or residence in an endemic geographic region during the incubation period (Table 211-1); (3) exposure to parturient ruminants, cats, and dogs (Q fever); (4) exposure to flying squirrels (*R. prowazekii* infection); and (5) history of previous louse-borne typhus (recrudescent typhus).

Clinical laboratory findings, such as thrombocytopenia (particularly in spotted fever and typhus rickettsioses, ehrlichioses, anaplasmosis, and scrub typhus), normal or low WBC counts, mild to moderate serum elevations of hepatic aminotransferases, and hyponatremia, suggest some common pathophysiologic mechanisms.

Application of these clinical, epidemiologic, and laboratory principles requires consideration of a rickettsial diagnosis and knowledge of the individual diseases.

TICK-, MITE-, LOUSE-, AND FLEA-BORNE RICKETTSIOSES

These diseases, caused by organisms of the genera *Rickettsia* and *Orientia* in the family Rickettsiaceae, result from endothelial cell infection and increased vascular permeability. Pathogenic rickettsial species are very closely related, have small genomes (as a result of reductive evolution, which eliminated many genes for biosynthesis of intracellularly available molecules), and are traditionally separated into typhus and spotted fever groups on the basis of lipopolysaccharide antigens. Some diseases and their agents (e.g., *R. africae*, *R. parkeri*, and *R. sibirica*) are too similar to require separate descriptions. Indeed, the similarities among MSF (*R. conorii* [all strains] and *R. massiliae*), North Asian tick typhus (*R. sibirica*), Japanese spotted fever (*R. japonica*), and Flinders Island spotted fever (*R. honei*) far outweigh the minor variations. The Rickettsiaceae that cause life-threatening infections are, in order of decreasing case-fatality rate, *R. rickettsii* (RMSF); *R. prowazekii* (louse-borne typhus); *Orientia tsutsugamushi* (scrub typhus); *R. conorii* (MSF); *R. typhi* (murine typhus); and, in rare cases, other spotted fever-group organisms. Some agents (e.g., *R. parkeri*, *R. africae*, *R. akari*, *R. slovacca*, *R. honei*, *R. felis*, *R. massiliae*, *R. helvetica*, *R. heilongjiangensis*, *R. aeschlimannii*, and *R. monacensis*) have never been documented to cause a fatal illness.

ROCKY MOUNTAIN SPOTTED FEVER



Epidemiology RMSF occurs in 47 states (with the highest prevalence in the south-central and southeastern states) as well as in Canada, Mexico, and Central and South America.

The infection is transmitted by *Dermacentor variabilis*, the American dog tick, in the eastern two-thirds of the United States and California; by *D. andersoni*, the Rocky Mountain wood tick, in the western United States; by *Rhipicephalus sanguineus* in Mexico, Arizona, and probably Brazil; and by *Amblyomma cajennense* and *A. aureolatum* in Central and/or South America. Maintained principally by transovarian transmission from one generation of ticks to the next, *R. rickettsii* can be acquired by uninfected ticks through the ingestion of a blood meal from rickettsemic small mammals.

Humans become infected during tick season (in the Northern Hemisphere, from May to September), although some cases occur in winter. The mortality rate was 20–25% in the preantibiotic era and remains at ~3–5%, principally because of delayed diagnosis and treatment. The case-fatality ratio increases with each decade of life above age 20.

Pathogenesis *R. rickettsii* organisms are inoculated into the dermis along with secretions of the tick's salivary glands after ≥6 h of feeding.

TABLE 211-1 FEATURES OF SELECTED RICKETTSIAL INFECTIONS

Disease	Organism	Transmission	Geographic Range	Incubation Period, Days	Duration, Days	Rash, %	Eschar, %	Lymphadenopathy ^a
Rocky Mountain spotted fever	<i>Rickettsia rickettsii</i>	Tick bite: <i>Dermacentor andersoni</i> , <i>D. variabilis</i> <i>Amblyomma cajennense</i> , <i>A. aureolatum</i> <i>Rhipicephalus sanguineus</i>	United States Central/South America Mexico, Brazil, United States	2–14	10–20	90	<1	+
Mediterranean spotted fever	<i>R. conorii</i>	Tick bite: <i>R. sanguineus</i> , <i>Rhipicephalus pumilio</i>	Southern Europe, Africa, Middle East, Central Asia	5–7	7–14	97	50	+
African tick-bite fever	<i>R. africae</i>	Tick bite: <i>A. hebraeum</i> , <i>A. variegatum</i>	Sub-Saharan Africa, West Indies	4–10	4–19	50	90	++++
Maculatum disease	<i>R. parkeri</i>	Tick bite: <i>A. maculatum</i>	United States, South America	2–10	6–16	88	94	++
Rickettsialpox	<i>R. akari</i>	Mite bite: <i>Liponyssoides sanguineus</i>	United States, Ukraine, Turkey, Mexico, Croatia	10–17	3–11	100	90	+++
Tick-borne lymphadenopathy	<i>R. slovaca</i>	Tick bite: <i>D. marginatus</i> , <i>D. reticularis</i>	Europe	7–9	17–180	5	100	++++
Flea-borne spotted fever	<i>R. felis</i>	Flea (mechanism undetermined): <i>Ctenocephalides felis</i>	Worldwide	8–16	8–16	80	15	—
Epidemic typhus	<i>R. prowazekii</i>	Louse feces: <i>Pediculus humanus corporis</i> , fleas and lice of flying squirrels, or recrudescence	Worldwide	7–14	10–18	80	None	—
Murine typhus	<i>R. typhi</i>	Flea feces: <i>Xenopsylla cheopis</i> , <i>C. felis</i> , others	Worldwide	8–16	9–18	80	None	—
Human monocytotropic ehrlichiosis	<i>Ehrlichia chaffeensis</i>	Tick bite: <i>A. americanum</i> , <i>D. variabilis</i>	United States	1–21	3–21	26	None	++
Ewingii ehrlichiosis	<i>E. ewingii</i>	Tick bite: <i>A. americanum</i>	United States	1–21	4–21	0	None	—
Unnamed ehrlichiosis	<i>E. muris</i> -like agent	Tick bite: <i>Ixodes scapularis</i>	United States	Unknown	3–14	None	None	—
Human granulocytotropic anaplasmosis	<i>Anaplasma phagocytophilum</i>	Tick bite: <i>I. scapularis</i> , <i>I. ricinus</i> , <i>I. pacificus</i> , <i>I. persulcatus</i>	United States, Europe, Asia	4–8	3–14	Rare	None	—
Unnamed disease	<i>Candidatus Neoehrlichia mikurensis</i>	Tick bite: <i>I. ricinus</i> , <i>I. persulcatus</i> , <i>Haemaphysalis concinna</i>	Europe, China	≥8	11–75	10	None	—
Scrub typhus	<i>Orientia tsutsugamushi</i>	Mite bite: <i>Leptotrombidium deliense</i> , others	Asia, Australia, Pacific and Indian Ocean islands	9–18	6–21	50	35	+++
Q fever	<i>Coxiella burnetii</i>	Inhalation of aerosols of infected parturition material (goats, sheep, cattle, cats, others), ingestion of infected milk or milk products	Worldwide except New Zealand, Antarctica	3–30	5–57	<1	None	—

^a++++, severe; +++, marked; ++, moderate; +, present in a small proportion of cases; —, not a noted feature.

The rickettsiae spread lymphohematogenously throughout the body and infect numerous foci of contiguous endothelial cells. The dose-dependent incubation period is ~1 week (range, 2–14 days). Occlusive thrombosis and ischemic necrosis are not the fundamental pathologic bases for tissue and organ injury. Instead, increased vascular permeability, with resulting edema, hypovolemia, and ischemia, is responsible. Consumption of platelets results in thrombocytopenia in 32–52% of patients, but disseminated intravascular coagulation with hypofibrinogenemia is rare. Activation of platelets, generation of thrombin, and activation of the fibrinolytic system all appear to be homeostatic physiologic responses to endothelial injury.

Clinical Manifestations Early in the illness, when medical attention usually is first sought, RMSF is difficult to distinguish from many self-limiting viral illnesses. Fever, headache, malaise, myalgia, nausea, vomiting, and anorexia are the most common symptoms during the first 3 days. The patient becomes progressively more ill as vascular infection and injury advance. In one large series, only one-third of patients were diagnosed with presumptive RMSF early in the clinical

course and treated appropriately as outpatients. In the tertiary-care setting, RMSF is all too often recognized only when late severe manifestations, developing at the end of the first week or during the second week of illness in patients without appropriate treatment, prompt return to a physician or hospital and admission to an intensive care unit.

The progressive nature of the infection is clearly manifested in the skin. Rash is evident in only 14% of patients on the first day of illness and in only 49% during the first 3 days. Macules (1–5 mm) appear first on the wrists and ankles and then on the remainder of the extremities and the trunk. Later, more severe vascular damage results in frank hemorrhage at the center of the maculopapule, producing a petechia that does not disappear upon compression (Fig. 211-1). This sequence of events is sometimes delayed or aborted by effective treatment. However, the rash is a variable manifestation, appearing on day 6 or later in 20% of cases and not appearing at all in 9–16% of cases. Petechiae occur in 41–59% of cases, appearing on or after day 6 in 74% of cases that manifest a rash. Involvement of the palms and soles, often considered diagnostically important, usually develops relatively late in

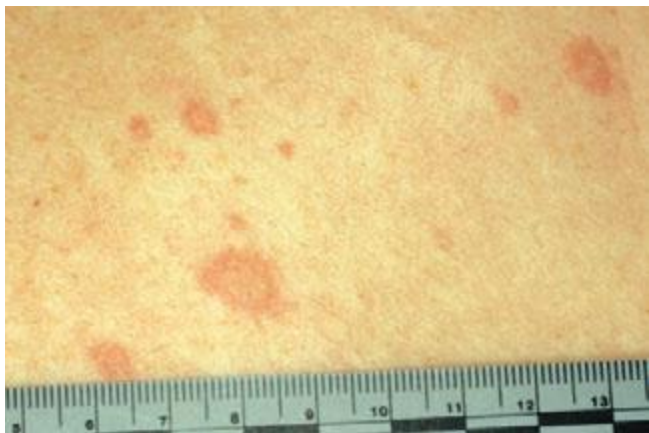


FIGURE 211-1 *Top:* Petechial lesions of Rocky Mountain spotted fever on the lower legs and soles of a young, previously healthy patient. *Bottom:* Close-up of lesions from the same patient. (Photos courtesy of Dr. Lindsey Baden; with permission.)

the course (after day 5 in 43% of cases) and does not develop at all in 18–64% of cases.

Hypovolemia leads to prerenal azotemia and (in 17% of cases) hypotension. Infection of the pulmonary microcirculation leads to noncardiogenic pulmonary edema; 12% of patients have severe respiratory disease, and 8% require mechanical ventilation. Cardiac involvement manifests as dysrhythmia in 7–16% of cases.

Besides respiratory failure, central nervous system (CNS) involvement is the other important determinant of the outcome of RMSF. Encephalitis, presenting as confusion or lethargy, is apparent in 26–28% of cases. Progressively severe encephalitis manifests as stupor or delirium in 21–26% of cases, ataxia in 18%, coma in 10%, and seizures in 8%. Numerous focal neurologic deficits have been reported. Meningoencephalitis results in cerebrospinal fluid (CSF) pleocytosis in 34–38% of cases; usually there are 10–100 cells/ μL and a mononuclear predominance, but occasionally there are >100 cells/ μL and a polymorphonuclear predominance. The CSF protein concentration is increased in 30–35% of cases, but the CSF glucose concentration is usually normal.

Renal failure, often reversible with rehydration, is caused by acute tubular necrosis in severe cases with shock. Hepatic injury with increased serum aminotransferase concentrations (38% of cases) is due to focal death of individual hepatocytes without hepatic failure. Jaundice is recognized in 9% of cases and an elevated serum bilirubin concentration in 18–30%.

Life-threatening bleeding is rare. Anemia develops in 30% of cases and is severe enough to require transfusions in 11%. Blood is detected in the stool or vomitus of 10% of patients, and death has followed massive upper-gastrointestinal hemorrhage.

Other characteristic clinical laboratory findings include increased plasma levels of proteins of the acute-phase response (C-reactive

protein, fibrinogen, ferritin, and others), hypoalbuminemia, and hyponatremia (in 56% of cases) due to the appropriate secretion of antidiuretic hormone in response to the hypovolemic state. Myositis occurs occasionally, with marked elevations in serum creatine kinase levels and multifocal rhabdomyonecrosis. Ocular involvement includes conjunctivitis in 30% of cases and retinal vein engorgement, flame hemorrhages, arterial occlusion, and papilledema with normal CSF pressure in some instances.

In untreated cases, the patient usually dies 8–15 days after onset. A rare presentation, fulminant RMSF, is fatal within 5 days after onset. This fulminant presentation is seen most often in male black patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency and may be related to an undefined effect of hemolysis on the rickettsial infection. Although survivors of RMSF usually return to their previous state of health, permanent sequelae, including neurologic deficits and gangrene necessitating amputation of extremities, may follow severe illness.

Diagnosis The diagnosis of RMSF during the acute stage is more difficult than is generally appreciated. The most important epidemiologic factor is a history of exposure to a potentially tick-infested environment within the 14 days preceding disease onset during a season of possible tick activity. However, only 60% of patients actually recall being bitten by a tick during the incubation period.

The differential diagnosis for early clinical manifestations of RMSF (fever, headache, and myalgia without a rash) includes influenza, enteroviral infection, infectious mononucleosis, viral hepatitis, leptospirosis, typhoid fever, gram-negative or gram-positive bacterial sepsis, HME, HGA, murine typhus, sylvatic flying-squirrel typhus, and rickettsialpox. Enterocolitis may be suggested by nausea, vomiting, and abdominal pain; prominence of abdominal tenderness has resulted in exploratory laparotomy. CNS involvement can masquerade as bacterial or viral meningoencephalitis. Cough, pulmonary signs, and chest radiographic opacities can lead to a diagnostic consideration of bronchitis or pneumonia.

At presentation during the first 3 days of illness, only 3% of patients exhibit the classic triad of fever, rash, and history of tick exposure. When a rash appears, a diagnosis of RMSF should be considered. However, many illnesses considered in the differential diagnosis also can be associated with a rash, including rubeola, rubella, meningococemia, disseminated gonococcal infection, secondary syphilis, toxic shock syndrome, drug hypersensitivity, idiopathic thrombocytopenic purpura, thrombotic thrombocytopenic purpura, Kawasaki syndrome, and immune complex vasculitis. Conversely, any person in an endemic area with a provisional diagnosis of one of the above illnesses could have RMSF. Thus, if a viral infection is suspected during RMSF season in an endemic area, it should always be kept in mind that RMSF can mimic viral infection early in the course; if the illness worsens over the next couple of days after initial presentation, the patient should return for reevaluation.

The most common serologic test for confirmation of the diagnosis is the indirect immunofluorescence assay. Not until 7–10 days after onset is a diagnostic titer of ≥ 64 usually detectable. The sensitivity and specificity of the indirect immunofluorescence IgG assay are 89–100% and 99–100%, respectively. It is important to understand that serologic tests for RMSF are usually negative at the time of presentation for medical care and that treatment should not be delayed while a positive serologic result is awaited.

The only diagnostic test that has proven useful during the acute illness is immunohistologic examination of a cutaneous biopsy sample from a rash lesion for *R. rickettsii*. Examination of a 3-mm punch biopsy from such a lesion is 70% sensitive and 100% specific. The sensitivity of polymerase chain reaction (PCR) amplification and detection of *R. rickettsii* DNA in peripheral blood is improving. However, although rickettsiae are present in large quantities in heavily infected foci of endothelial cells, there are relatively low quantities in the circulation. Cultivation of rickettsiae in cell culture is feasible but is seldom undertaken because of biohazard concerns. The recent dramatic increase in the reported incidence of RMSF correlates with

the use of single-titer spotted fever–group cross-reactive enzyme immunoassay serology. Few cases are specifically determined to be caused by *R. rickettsii*. Currently, many febrile persons who do not have RMSF present with cross-reactive antibodies, possibly because of previous exposure to the highly prevalent spotted fever–group rickettsia *R. amblyomii*.

TREATMENT ROCKY MOUNTAIN SPOTTED FEVER

The drug of choice for the treatment of both children and adults with RMSF is doxycycline, except when the patient is pregnant or allergic to this drug (see below). Because of the severity of RMSF, immediate empirical administration of doxycycline should be strongly considered for any patient with a consistent clinical presentation in the appropriate epidemiologic setting. Doxycycline is administered orally (or, in the presence of coma or vomiting, intravenously) at 200 mg/d in two divided doses. For children with suspected RMSF, up to five courses of doxycycline may be administered with minimal risk of dental staining. Other regimens include oral tetracycline (25–50 mg/kg per day) in four divided doses. Treatment with chloramphenicol, a less effective drug, is advised only for patients who are pregnant or allergic to doxycycline. The antirickettsial drug should be administered until the patient has been afebrile and improving clinically for 2–3 days. β -Lactam antibiotics, erythromycin, and aminoglycosides have no role in the treatment of RMSF, and sulfa-containing drugs are associated with more adverse outcomes than no treatment at all. There is little clinical experience with fluoroquinolones, clarithromycin, and azithromycin, which are not recommended. The most seriously ill patients are managed in intensive care units, with careful administration of fluids to achieve optimal tissue perfusion without precipitating noncardiogenic pulmonary edema. In some severely ill patients, hypoxemia requires intubation and mechanical ventilation; oliguric or anuric acute renal failure requires hemodialysis; seizures necessitate the use of antiseizure medication; anemia or severe hemorrhage necessitates transfusions of packed red blood cells; or bleeding with severe thrombocytopenia requires platelet transfusions. Heparin is not a useful component of treatment, and there is no evidence that glucocorticoids affect outcome.

Prevention Avoidance of tick bites is the only available preventive approach. Use of protective clothing and tick repellents, inspection of the body once or twice a day, and removal of ticks before they inoculate rickettsiae reduce the risk of infection. Prophylactic doxycycline treatment of tick bites has no proven role in preventing RMSF.

MEDITERRANEAN SPOTTED FEVER (BOUTONNEUSE FEVER), AFRICAN TICK-BITE FEVER, AND OTHER TICK-BORNE SPOTTED FEVERS

Epidemiology *R. conorii* is prevalent in southern Europe, Africa, and southwestern and south-central Asia. Regional names for the disease caused by this organism include Mediterranean spotted fever, Kenya tick typhus, Indian tick typhus, Israeli spotted fever, and Astrakhan spotted fever. The disease is characterized by high fever, rash, and—in most geographic locales—an inoculation eschar (*tâche noire*) at the site of the tick bite. A severe form of the disease (mortality rate, 50%) occurs in patients with diabetes, alcoholism, or heart failure.

African tick-bite fever, caused by *R. africae*, occurs in rural areas of sub-Saharan Africa and in the Caribbean islands and is transmitted by *Amblyomma hebraeum* and *A. variegatum* ticks. The average incubation period is 4–10 days. The mild illness consists of headache, fever, eschar, and regional lymphadenopathy. *Amblyomma* ticks often feed in groups, with the consequent development of multiple eschars. Rash may be vesicular, sparse, or absent altogether. Because of tourism in sub-Saharan Africa, African tick-bite fever is the rickettsiosis most frequently imported into Europe and North America. A similar

disease caused by the closely related species *R. parkeri* is transmitted by *A. maculatum* in the United States and by *A. triste* in South America.

R. japonica causes Japanese spotted fever, which also occurs in Korea. Similar diseases in northern Asia are caused by *R. sibirica* and *R. heilongjiangensis*. Queensland tick typhus due to *R. australis* is transmitted by *Ixodes holocyclus* ticks. Flinders Island spotted fever, found on the island for which it is named as well as in Tasmania, mainland Australia, and Asia, is caused by *R. honei*. In Europe, patients infected with *R. slovaca* after a wintertime *Dermacentor* tick bite manifest an afebrile illness with an eschar (usually on the scalp) and painful regional lymphadenopathy.

Diagnosis Diagnosis of these tick-borne spotted fevers is based on clinical and epidemiologic findings and is confirmed by serology, immunohistochemical demonstration of rickettsiae in skin biopsy specimens, cell-culture isolation of rickettsiae, or PCR of skin biopsy, eschar, or blood samples. Serologic diagnosis detects antibodies to antigens shared among spotted fever–group rickettsiae, hindering identification of the etiologic species. In an endemic area, a possible diagnosis of rickettsial spotted fevers should be considered when patients present with fever, rash, and/or a skin lesion consisting of a black necrotic area or a crust surrounded by erythema.

TREATMENT TICK-BORNE SPOTTED FEVERS

Successful therapeutic agents include doxycycline (100 mg bid orally for 1–5 days) and chloramphenicol (500 mg qid orally for 7–10 days). Pregnant patients may be treated with josamycin (3 g/d orally for 5 days). Data on the efficacy of treatment of mildly ill children with clarithromycin or azithromycin should not be extrapolated to adults or to patients with moderate or severe illness.

RICKETTSIALPOX

R. akari infects mice and their mites (*Liponyssoides sanguineus*), which maintain the organisms by transovarial transmission.

Epidemiology Rickettsialpox is recognized principally in New York City, but cases have also been reported in other urban and rural locations in the United States and in Ukraine, Croatia, Mexico, and Turkey. Investigation of eschars suspected of representing bioterrorism-associated cutaneous anthrax revealed that rickettsialpox occurs more frequently than previously realized.

Clinical Manifestations A papule forms at the site of the mite's feeding, develops a central vesicle, and becomes a 1- to 2.5-cm painless black crusted eschar surrounded by an erythematous halo (Fig. 211-2). Enlargement of the regional lymph nodes draining the eschar suggests initial lymphogenous spread. After an incubation period of



FIGURE 211-2 Eschar at the site of the mite bite in a patient with rickettsialpox. (Reprinted from A Krusell et al: *Emerg Infect Dis* 8:727, 2002. Photo obtained by Dr. Kenneth Kaye.)



FIGURE 211-3 **Top:** Papulovesicular lesions on the trunk of the patient with rickettsialpox shown in Fig. 211-2. **Bottom:** Close-up of lesions from the same patient. (Reprinted from A Krusell et al: *Emerg Infect Dis* 8:727, 2002. Photos obtained by Dr. Kenneth Kaye.)

10–17 days, during which the eschar and regional lymphadenopathy frequently go unnoticed, disease onset is marked by malaise, chills, fever, headache, and myalgia. A macular rash appears 2–6 days after onset and usually evolves sequentially into papules, vesicles, and crusts that heal without scarring (Fig. 211-3); in some cases, the rash remains macular or maculopapular. Some patients develop nausea, vomiting, abdominal pain, cough, conjunctivitis, or photophobia. Without treatment, fever lasts 6–10 days.

Diagnosis and Treatment Clinical, epidemiologic, and convalescent serologic data establish the diagnosis of a spotted fever–group rickettsiosis that is seldom pursued further. Doxycycline is the drug of choice for treatment.

FLEA-BORNE SPOTTED FEVER



An emerging rickettsiosis caused by *R. felis* occurs worldwide. Maintained transovarially in the geographically widespread cat flea *Ctenocephalides felis*, the infection has been described as moderately severe, with fever, rash, and headache as well as CNS, gastrointestinal, and pulmonary symptoms.

EPIDEMIC (LOUSE-BORNE) TYPHUS

Epidemiology The human body louse (*Pediculus humanus corporis*) lives in clothing under poor hygienic conditions and usually in impoverished cold areas. Lice acquire *R. prowazekii* when they ingest blood from a rickettsemic patient. The rickettsiae multiply in the louse's mid-gut epithelial cells and are shed in its feces. The infected louse leaves a febrile person and deposits infected feces on its subsequent host during its blood meal; the patient autoinoculates the organisms by scratching. The louse is killed by the rickettsiae and does not pass *R. prowazekii* to its offspring.



Epidemic typhus haunts regions afflicted by wars and disasters. An outbreak involved 100,000 people in refugee camps in Burundi in 1997. A small focus was documented in Russia in 1998; sporadic cases were reported from Algeria, and frequent outbreaks occurred in Peru. Eastern flying squirrels (*Glaucomys volans*) and their lice and fleas maintain *R. prowazekii* in a zoonotic cycle.

Brill-Zinsser disease is a recrudescence illness occurring years after acute epidemic typhus, probably as a result of waning immunity. *R. prowazekii* remains latent for years; its reactivation results in sporadic cases of disease in louse-free populations or in epidemics in louse-infested populations.

Rickettsiae are potential agents of bioterrorism (Chap. 261e). Infections with *R. prowazekii* and *R. rickettsii* have high case–fatality ratios. These organisms cause difficult-to-diagnose diseases and are highly infectious when inhaled as aerosols. Organisms resistant to tetracycline or chloramphenicol have been developed in the laboratory.

Clinical Manifestations After an incubation period of ~1–2 weeks, the onset of illness is abrupt, with prostration, severe headache, and fever rising rapidly to 38.8°–40.0°C (102°–104°F). Cough is prominent, developing in 70% of patients. Myalgias are usually severe. A rash begins on the upper trunk, usually on the fifth day, and then becomes generalized, involving the entire body except the face, palms, and soles. Initially, this rash is macular; without treatment, it becomes maculopapular, petechial, and confluent. The rash often goes undetected in black skin; 60% of African patients have spotless epidemic typhus. Photophobia, with considerable conjunctival injection and eye pain, is common. The tongue may be dry, brown, and furred. Confusion and coma are common. Skin necrosis and gangrene of the digits as well as interstitial pneumonia may occur in severe cases. Untreated disease is fatal in 7–40% of cases, with outcome depending primarily on the condition of the host. Patients with untreated infections develop renal insufficiency and multiorgan involvement in which neurologic manifestations are frequently prominent. Overall, 12% of patients with epidemic typhus have neurologic involvement. Infection associated with North American flying squirrels is a milder illness; whether this milder disease is due to host factors (e.g., better health status) or attenuated virulence is unknown.

Diagnosis and Treatment Epidemic typhus is sometimes misdiagnosed as typhoid fever in tropical countries (Chap. 190). The means even for serologic studies are often unavailable in settings of louse-borne typhus. Epidemics can be recognized by the serologic or immunohistochemical diagnosis of a single case or by detection of *R. prowazekii* in a louse found on a patient. Doxycycline (200 mg/d, given in two divided doses) is administered orally or—if the patient is comatose or vomiting—intravenously. Although under epidemic conditions a single 200-mg oral dose is effective, treatment is generally continued until 2–3 days after defervescence. Pregnant patients should be evaluated individually and treated with chloramphenicol early in pregnancy or, if necessary, with doxycycline late in pregnancy.

Prevention Prevention of epidemic typhus involves control of body lice. Clothes should be changed regularly, and insecticides should be used every 6 weeks to control the louse population.

ENDEMIC MURINE TYPHUS

Epidemiology *R. typhi* is maintained in mammalian host/flea cycles, with rats (*Rattus rattus* and *R. norvegicus*) and the Oriental rat flea (*Xenopsylla cheopis*) as the classic zoonotic niche. Fleas acquire *R. typhi* from rickettsemic rats and carry the organism throughout their life span. Nonimmune rats and humans are infected when rickettsia-laden flea feces contaminate pruritic bite lesions; less frequently, the flea bite transmits the organisms. Transmission can also occur via inhalation of aerosolized rickettsiae from flea feces. Infected rats appear healthy, although they are rickettsemic for ~2 weeks.



Murine typhus occurs mainly in Texas and southern California, where the classic rat/flea cycle is absent and an opossum/cat flea (*C. felis*) cycle is prominent. Globally, endemic typhus occurs mainly in warm (often coastal) areas

throughout the tropics and subtropics, where it is highly prevalent though often unrecognized. The incidence peaks from April through June in southern Texas and during the warm months of summer and early fall in other geographic locations. Patients seldom recall exposure to fleas, although exposure to animals such as cats, opossums, and rats is reported in nearly 40% of cases.

Clinical Manifestations The incubation period of experimental murine typhus averages 11 days (range, 8–16 days). Headache, myalgia, arthralgia, nausea, and malaise develop 1–3 days before onset of chills and fever. Nearly all patients experience nausea and vomiting early in the illness.

The duration of untreated illness averages 12 days (range, 9–18 days). Rash is present in only 13% of patients at presentation for medical care (usually ~4 days after onset of fever), appearing an average of 2 days later in half of the remaining patients and never appearing in the others. The initial macular rash is often detected by careful inspection of the axilla or the inner surface of the arm. Subsequently, the rash becomes maculopapular, involving the trunk more often than the extremities; it is seldom petechial and rarely involves the face, palms, or soles. A rash is detected in only 20% of patients with darkly pigmented skin.

Pulmonary involvement is frequently prominent; 35% of patients have a hacking, nonproductive cough, and 23% of patients who undergo chest radiography have pulmonary densities due to interstitial pneumonia, pulmonary edema, and pleural effusions. Bibasilar rales are the most common pulmonary sign. Less common clinical manifestations include abdominal pain, confusion, stupor, seizures, ataxia, coma, and jaundice. Clinical laboratory studies frequently reveal anemia and leukopenia early in the course, leukocytosis late in the course, thrombocytopenia, hyponatremia, hypoalbuminemia, mildly increased serum hepatic aminotransferases, and prerenal azotemia. Complications can include respiratory failure, hematemesis, cerebral hemorrhage, and hemolysis. Severe illness necessitates the admission of 10% of hospitalized patients to an intensive care unit. Greater severity is generally associated with old age, underlying disease, and treatment with a sulfonamide; the case-fatality rate is 1%. In a study of children with murine typhus, 50% suffered only nocturnal fevers, feeling well enough for active daytime play.

Diagnosis and Treatment Serologic studies of acute- and convalescent-phase sera can provide a diagnosis, and an immunohistochemical method for identification of typhus group-specific antigens in biopsy samples has been developed. Cultivation and PCR are used only infrequently and are not widely available. Nevertheless, most patients are treated empirically with doxycycline (100 mg bid orally for 7–15 days) on the basis of clinical suspicion. Ciprofloxacin provides an alternative if doxycycline is contraindicated.

SCRUB TYPHUS

Epidemiology *O. tsutsugamushi* differs substantially from *Rickettsia* species both genetically and in cell wall composition (i.e., it lacks lipopolysaccharide). *O. tsutsugamushi* is maintained by transovarial transmission in trombiculid mites. After hatching, infected larval mites (chiggers, the only stage that feeds on a host) inoculate organisms into the skin. Infected chiggers are particularly likely to be found in areas of heavy scrub vegetation during the wet season, when mites lay eggs.

Scrub typhus is endemic and reemerging in eastern and southern Asia, northern Australia, and islands of the western Pacific and Indian Oceans. Infections are prevalent in these regions; in some areas, >3% of the population is infected or reinfected each month. Immunity wanes over 1–3 years, and the organism exhibits remarkable antigenic diversity.

Clinical Manifestations Illness varies from mild and self-limiting to fatal. After an incubation period of 6–21 days, onset is characterized by fever, headache, myalgia, cough, and gastrointestinal symptoms. Some patients recover spontaneously after a few days. The classic case description includes an eschar where the chigger has fed, regional lymphadenopathy, and a maculopapular rash—signs that are seldom seen in indigenous patients. Fewer than 50% of Westerners develop an

eschar, and fewer than 40% develop a rash (on day 4–6 of illness). Severe cases typically manifest with encephalitis and interstitial pneumonia due to vascular injury. The case-fatality rate for untreated classic cases is 7% but would probably be lower if all mild cases were diagnosed.

Diagnosis and Treatment Serologic assays (indirect fluorescent antibody, indirect immunoperoxidase, and enzyme immunoassays) are the mainstays of laboratory diagnosis. PCR amplification of *Orientia* genes from eschars and blood also is effective. Patients are treated with doxycycline (100 mg bid orally for 7–15 days), azithromycin (500 mg orally for 3 days), or chloramphenicol (500 mg qid orally for 7–15 days). Some cases of scrub typhus in Thailand are caused by strains that have high doxycycline or chloramphenicol minimal inhibitory concentrations (MICs) but that are susceptible to azithromycin and rifampin.

EHRlichIOSES AND ANAPLASMOSIS

Ehrlichioses are acute febrile infections caused by members of the family Anaplasmataceae, which is made up of obligately intracellular organisms of five genera: *Ehrlichia*, *Anaplasma*, *Wolbachia*, *Candidatus Neoehrlichia*, and *Neorickettsia*. The bacteria reside in vertebrate reservoirs and target vacuoles of hematopoietic cells (Fig. 211-4). Three *Ehrlichia* species and one *Anaplasma* species are transmitted by ticks to humans and cause infection that can be severe and prevalent. *E. chaffeensis*, the agent of HME, and an *E. muris*-like agent (EMLA) infect predominantly mononuclear phagocytes; *E. ewingii* and *A. phagocytophilum* infect neutrophils. Infection with *Candidatus Neoehrlichia mikurensis* is less well characterized, but the agent has been identified in human blood neutrophils.

Ehrlichia, *Candidatus Neoehrlichia*, and *Anaplasma* are maintained by horizontal tick-mammal-tick transmission, and humans are only inadvertently infected. *Wolbachiae* are associated with human filariasis, since they are important for filarial viability and pathogenicity; antibiotic treatment targeting *wolbachiae* is a strategy for filariasis control. *Neorickettsiae* parasitize flukes (trematodes) that in turn parasitize aquatic snails, fish, and insects. Only a single human *neorickettsiosis* has been described: sennetsu fever, an infectious mononucleosis-like illness that was first identified in 1953 and is associated with the ingestion of raw fish containing *N. sennetsu*-infected flukes.

HUMAN MONOCYTOTROPIC EHRlichIOSIS

Epidemiology More than 8404 cases of *E. chaffeensis* infection had been reported to the Centers for Disease Control and Prevention (CDC) as of April 2013. However, active prospective surveillance has documented an incidence as high as 414 cases per 100,000 population

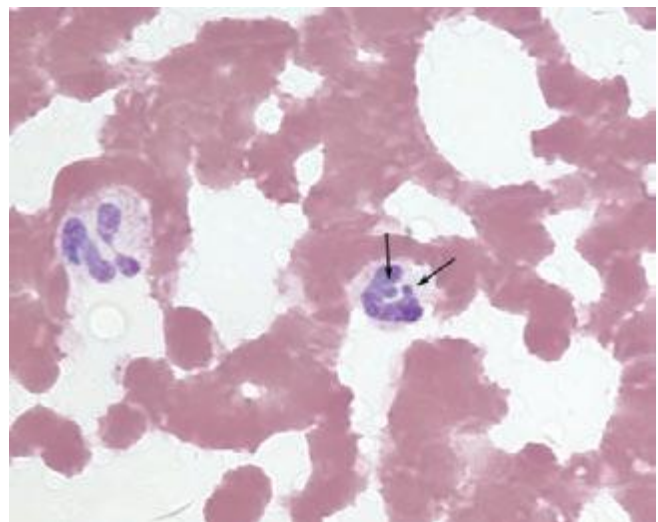


FIGURE 211-4 Peripheral-blood smear from a patient with human granulocytotropic anaplasmosis. A neutrophil contains two morulae (vacuoles filled with *A. phagocytophilum*). (Photo courtesy of Dr. J. Stephen Dumler.)

1160 in some U.S. regions. Most *E. chaffeensis* infections are identified in the south-central, southeastern, and mid-Atlantic states, but cases have also been recognized in California and New York. All stages of the Lone Star tick (*A. americanum*) feed on white-tailed deer—a major reservoir. Dogs and coyotes also serve as reservoirs and often lack clinical signs. Tick bites and exposures are frequently reported by patients in rural areas, especially in May through July. The median age of HME patients is 52 years; however, severe and fatal infections in children also are well recognized. Of patients with HME, 60% are male.



E. chaffeensis has been detected in South America, Africa, and Asia.

Clinical Manifestations *E. chaffeensis* disseminates hematogenously from the dermal blood pool created by the feeding tick. After a median incubation period of 8 days, illness develops. Clinical manifestations are undifferentiated and include fever (96% of cases), headache (72%), myalgia (68%), and malaise (77%). Less frequently observed are nausea, vomiting, and diarrhea (25–57%); cough (28%); rash (26% overall, 6% at presentation); and confusion (20%). HME can be severe: 49% of patients with documented cases are hospitalized, and ~2% die. Severe manifestations include a toxic shock–like or septic shock–like syndrome, adult respiratory distress syndrome, cardiac failure, hepatitis, meningoencephalitis, hemorrhage, and—in immunocompromised patients—overwhelming ehrlichial infection. Laboratory findings are valuable in the differential diagnosis of HME; 61% of patients have leukopenia (initially lymphopenia, later neutropenia), 73% have thrombocytopenia, and 84% have elevated serum levels of hepatic aminotransferases. Despite low blood cell counts, the bone marrow is hypercellular, and noncaseating granulomas can be present. Vasculitis is not a component of HME.

Diagnosis HME can be fatal. Early empirical antibiotic therapy based on clinical diagnosis diminishes adverse outcomes. This diagnosis is suggested by fever with a known tick exposure during the preceding 3 weeks, thrombocytopenia and/or leukopenia, and increased serum aminotransferase levels. Morulae are demonstrated in <10% of peripheral-blood smears. HME can be confirmed during active infection by PCR amplification of *E. chaffeensis* nucleic acids in blood obtained before the start of doxycycline therapy. Retrospective serodiagnosis requires a consistent clinical picture and a fourfold increase in *E. chaffeensis* antibody titer to ≥ 64 in paired sera obtained ~3 weeks apart. Separate specific diagnostic tests are necessary for HME and HGA.

EWINGII EHRLICHIOSIS AND EHRLICHIA MURIS–LIKE INFECTIONS

Ehrlichia ewingii, originally a neutrophil pathogen causing fever and lameness in dogs, resembles *E. chaffeensis* in its tick vector (*A. americanum*) and vertebrate reservoirs (white-tailed deer and dogs). An *E. muris*–like agent (EMLA) has been discovered and identified as the cause of human infections in Wisconsin and Minnesota. *E. ewingii* and EMLA illnesses are similar to but less severe than HME. Many cases occur in immunocompromised patients. No specific serologic diagnostic tests for ewingii or EMLA ehrlichiosis are readily available.

CANDIDATUS NEOEHRLICHIA MIKURENSIS INFECTION



Candidatus Neoehrlichia mikurensis, a bacterium in a phylogenetic clade between *Ehrlichia* and *Anaplasma*, was originally identified in *Ixodes ricinus* ticks from the Netherlands and in mice and *Ixodes ovatus* ticks from Japan. By means of broad-range 16S rRNA gene amplification and sequence analysis, this organism was identified as the cause of severe and sometimes prolonged febrile illnesses in European immunocompromised patients with tick bites or exposures and in Chinese patients with a mild febrile illness after being bitten by *Ixodes persulcatus* and *Haemaphysalis concinna* ticks. The clinical presentation is similar to those of HME and HGA. Specific diagnostic methods have been developed but are not widely available.

TREATMENT EHRLICHIOSES

Doxycycline is effective for HME as well as for ewingii and EMLA ehrlichioses; the use of this drug in *Candidatus N. mikurensis* infection is associated with disease resolution. Therapy with doxycycline (100 mg given PO or IV twice daily) or tetracycline (250–500 mg given PO every 6 h) lowers hospitalization rates and shortens fever duration. *E. chaffeensis* is not susceptible to chloramphenicol *in vitro*, and the use of this drug is controversial. While a few reports document *E. chaffeensis* persistence in humans, this finding is rare; most infections are cured by short courses of doxycycline (continuing for 3–5 days after defervescence). Although poorly studied, rifampin may be suitable when doxycycline is contraindicated.

PREVENTION

HME, ewingii ehrlichiosis, EMLA infection, and *Candidatus N. mikurensis* infection can be prevented by the avoidance of ticks in endemic areas. The use of protective clothing and tick repellents, careful post-exposure tick searches, and prompt removal of attached ticks probably diminish infection risk.

HUMAN GRANULOCYTOTROPIC ANAPLASMOSIS



Epidemiology As of April 2013, 10,181 cases of HGA had been reported to the CDC, most in the upper midwestern and northeastern United States; the geographic distribution is similar to that for Lyme disease because of the shared *I. scapularis* tick vector. White-footed mice, squirrels, and white-tailed deer in the United States and red deer in Europe are natural reservoirs for *A. phagocytophilum*. HGA incidence peaks in May through July, but the disease can occur throughout the year with exposure to *Ixodes* ticks. HGA often affects males (59%) and older persons (median age, 51 years).

Clinical Manifestations Seroprevalence rates are high in endemic regions; thus it seems likely that most individuals develop subclinical infections. The incubation period for HGA is 4–8 days, after which the disease manifests as fever (75–100% of cases), myalgia (77%), headache (82%), and malaise (97%). A minority of patients develop nausea, vomiting, or diarrhea (22–39%); cough (27%); or confusion (17%). Rash (6%) is almost invariably concurrent erythema migrans attributable to Lyme disease. Most patients develop thrombocytopenia (75%) and/or leukopenia (55%) with increased serum hepatic aminotransferase levels (83%).

Severe complications occur most often in the elderly and include adult respiratory distress syndrome, a toxic shock–like syndrome, and life-threatening opportunistic infections. Meningoencephalitis is rarely documented with HGA, but brachial plexopathy, cranial nerve involvement, and demyelinating polyneuropathy are reported. For HGA, 7% of patients require intensive care, and the case-fatality rate is 0.6%. Neither vasculitis nor granulomas are components of HGA. While co-infections with *Borrelia burgdorferi* and *Babesia microti* (transmitted by the same tick vector[s]) occur, there is little evidence of comorbidity or persistence. HGA is rarely acquired via transfusion.

Diagnosis HGA should be included in the differential diagnosis of influenza-like illnesses during seasons with *Ixodes* tick activity (May through December), especially with known tick bite or exposure. Concurrent thrombocytopenia, leukopenia, or elevated serum levels of alanine or aspartate aminotransferase further increase the likelihood of HGA. Many HGA patients develop Lyme disease antibodies in the absence of clinical findings consistent with that diagnosis. Thus, HGA should be considered in the differential diagnosis of atypical severe Lyme disease presentations. Peripheral-blood film examination for neutrophil morulae can yield a diagnosis in 20–75% of infections. PCR testing of blood from patients with active disease before doxycycline therapy is sensitive and specific. Serodiagnosis is retrospective, requiring a fourfold increase in *A. phagocytophilum* antibody titer (to ≥ 160) in paired serum samples obtained 1 month apart. Since seroprevalence is high in some regions, a single acute-phase titer should not be used for diagnosis.

No prospective studies of therapy for HGA have been conducted. However, doxycycline (100 mg PO twice daily) is effective. Rifampin therapy is associated with improvement of HGA in pregnant women and children. Most treated patients defervesce within 24–48 h.

Prevention HGA prevention requires tick avoidance. Transmission can be documented as few as 4 h after a tick bite.

Q FEVER

The agent of Q fever is *Coxiella burnetii*, a small intracellular prokaryote that only recently was grown in cell-free medium. *C. burnetii*, a pleomorphic coccobacillus with a gram-negative cell wall, survives in harsh environments; it escapes intracellular killing in macrophages by inhibiting the final step in phagosome maturation (cathepsin fusion) and has adapted to the acidic phagolysosome by producing superoxide dismutase. Infection with *C. burnetii* induces a range of immunomodulatory responses, from immunosuppression in chronic Q fever to the production of autoantibodies, particularly those to smooth muscle and cardiac muscle.

Q fever encompasses two broad clinical syndromes: acute and chronic infection. The host's immune response (rather than the particular strain) most likely determines whether chronic Q fever develops. *C. burnetii* survives in monocytes from patients with chronic Q fever but not in monocytes from patients with acute Q fever or from uninfected subjects. Impairment of the bactericidal activity of the *C. burnetii*-infected monocyte is associated with overproduction of interleukin 10. The CD4+/CD8+ ratio is decreased in Q fever endocarditis. Very few organisms and a strong cellular response are observed in patients with acute Q fever, while many organisms and a moderate cellular response occur in chronic Q fever. Immune control of *C. burnetii* is T cell-dependent, but 80–90% of bone marrow aspirates obtained years after recovery from Q fever contain *C. burnetii* DNA. *C. burnetii*'s ready multiplication within trophoblasts accounts for the high concentrations it can reach in the placenta.

Epidemiology Q fever is a zoonosis. The primary sources of human infection are infected cattle, sheep, and goats. However, cats, rabbits, pigeons, and dogs also serve as sources for transmission of *C. burnetii* to humans. The wildlife reservoir is extensive and includes ticks, coyotes, gray foxes, skunks, raccoons, rabbits, deer, mice, bears, birds, and opossums. In female animals *C. burnetii* localizes to the uterus and mammary glands. Infection is reactivated during pregnancy and after radiotherapy in mouse models. High concentrations of *C. burnetii* are found in the placenta. At the time of parturition, the bacteria are released into the air, and infection follows inhalation of aerosolized organisms by a susceptible host. Windstorms can generate *C. burnetii* aerosols months after soil contamination during parturition. Individuals up to 18 km from the source have been infected. Because it is easily dispersed as an aerosol, *C. burnetii* is a potential agent of bioterrorism (Chap. 261e), with a high infectivity rate and pneumonia as the major manifestation.

Determining the source of an outbreak of Q fever can be challenging. An outbreak of Q fever at a horse-boarding ranch in Colorado in 2005 was due to spread of infection from two herds of goats that had been acquired by the owners. PCR testing confirmed the presence of *C. burnetii* in the soil and among the goats. Of 138 persons who lived within 1 mile of the ranch and who were also tested, 11 (8%) had evidence of *C. burnetii* infection, and 8 of these 11 individuals had no direct contact with the ranch.

Persons at risk for Q fever include abattoir workers, veterinarians, farmers, and other individuals who have contact with infected animals (particularly newborn animals) or products of conception. The organism is shed in milk for weeks to months after parturition. The ingestion of contaminated milk in some geographic areas probably represents a major route of transmission to humans. A recent outbreak of Q fever associated with ingestion of raw milk confirms the oral route of transmission. In rare instances, person-to-person transmission follows labor and childbirth in an infected woman, autopsy of an infected

individual, or blood transfusion. Some evidence suggests that *C. burnetii* can be sexually transmitted among humans. Crushing an infected tick between the fingers has resulted in Q fever; the implication is that percutaneous transmission can occur.



Infections due to *C. burnetii* occur in most geographic locations except New Zealand and Antarctica. Thus Q fever can be associated with travel. The number of reported cases of Q fever in the United States ranges from 28 to 54 per year. More than 70% of these cases occur in males, and April, May, and June are the most common months for acquisition. Q fever continues to be common in Australia, with 30 cases per 1 million population per year. Cases among abattoir workers in Australia declined dramatically as a result of a vaccination program. An outbreak of Q fever began in the Netherlands in 2007, and by 2010 more than 4000 cases had been reported. Pneumonia was a common manifestation in this outbreak. The outbreak was due to a combination of high-density goat farming in areas abutting large urban populations and environmental factors. Farms where spread did not occur had high vegetation densities and lower groundwater concentrations.

The primary manifestations of acute Q fever differ geographically (e.g., pneumonia in Nova Scotia and granulomatous hepatitis in Marseille). These differences could reflect the route of infection (i.e., ingestion of contaminated milk for hepatitis and inhalation of contaminated aerosols for pneumonia) or strain differences. In the Netherlands outbreak, sequelae of infection in pregnant women were rare; this was not the case among pregnant women elsewhere.

Young age seems to be protective against disease caused by *C. burnetii*. In a large outbreak in Switzerland, symptomatic infection occurred five times more often among persons >15 years of age than among younger individuals. In many outbreaks, men are affected more commonly than women; the proposed explanation is that female hormones are partially protective.

Clinical Manifestations • ACUTE Q FEVER The symptoms of acute Q fever are nonspecific; common among them are fever, extreme fatigue, photophobia, and severe headache that is frequently retro-orbital. Other symptoms include chills, sweats, nausea, vomiting, and diarrhea, each occurring in 5–20% of cases. Cough develops in about half of patients with Q fever pneumonia. Neurologic manifestations of acute Q fever are uncommon; however, in one outbreak in the United Kingdom, 23% of 102 patients had neurologic signs and symptoms as the major manifestation. A nonspecific rash may be evident in 4–18% of patients. The WBC count is usually normal. Thrombocytopenia occurs in ~25% of patients, and reactive thrombocytosis (with platelet counts exceeding 10⁹/μL) frequently develops during recovery. Chest radiography can show opacities similar to those seen in pneumonia caused by other pathogens, but multiple rounded opacities in patients in endemic areas suggest a diagnosis of Q fever pneumonia.

Acute Q fever occasionally complicates pregnancy. In one series, it resulted in premature birth in 35% of cases and in abortion or neonatal death in 43%. Neonatal death (previous or current) and lower infant birth weight are three times more likely among women seropositive for *C. burnetii*.



After the usual incubation period of 3–30 days, 1070 patients with acute Q fever in southern France presented with hepatitis (40%), both pneumonia and hepatitis (20%), pneumonia (17%), isolated fever (14%), CNS involvement (2%), and pericarditis or myocarditis (1%). Acalculous cholecystitis, pancreatitis, lymphadenopathy, spontaneous rupture of the spleen, transient hypoplastic anemia, bone marrow necrosis, hemolytic anemia, histiocytic hemophagocytosis, optic neuritis, and erythema nodosum were less common manifestations.

POST-Q FEVER FATIGUE SYNDROME Prolonged fatigue can follow Q fever and can be accompanied by a constellation of symptoms including headaches, sweats, arthralgia, myalgias, blurred vision, muscle fasciculations, and enlarged and painful lymph nodes. Long-term persistence of a noninfective, nonbiodegraded complex of *Coxiella* cell components, with its antigens and specific lipopolysaccharide, has been detected in the affected persons. Patients who develop this syndrome have a higher frequency of carriage of HLA-DRB1*11 and of the 2/2 genotype of the interferon γ intron 1 microsatellite.

1162 CHRONIC Q FEVER Chronic Q fever almost always implies endocarditis and usually occurs in patients with previous valvular heart disease, immunosuppression, or chronic renal insufficiency. Fever is usually absent or low grade. Valvular vegetations are detected in only 12% of patients by transthoracic echocardiography, but the rate of detection is higher (21–50%) with transesophageal echocardiography. The vegetations in chronic Q fever endocarditis differ from those in bacterial endocarditis, manifesting as endothelium-covered nodules on the valves. A high index of suspicion is necessary for timely diagnosis. Patients with chronic Q fever are often ill for >1 year before the diagnosis is made. The disease should be suspected in all patients with culture-negative endocarditis. In addition, all patients with valvular heart disease and an unexplained purpuric eruption, renal insufficiency, stroke, and/or progressive heart failure should be tested for *C. burnetii* infection. Patients with chronic Q fever have hepatomegaly and/or splenomegaly, which—in combination with rheumatoid factor, elevated erythrocyte sedimentation rate, high C-reactive protein level, and/or increased γ -globulin concentrations (up to 60–70 g/L)—suggests this diagnosis. Other manifestations of chronic Q fever include infection of vascular prostheses, aneurysms, and bone as well as chronic sternal wound infection. Unusual manifestations include chronic thrombocytopenia, mixed cryoglobulinemia, and livedo reticularis.

Diagnosis Isolation of *C. burnetii* from buffy-coat blood samples or tissue specimens by a shell-vial technique is easy but requires a biosafety level 3 laboratory. PCR detects *C. burnetii* DNA in tissue specimens, including paraffin-embedded samples. Serology is the most commonly used diagnostic tool. Indirect immunofluorescence is sensitive and specific and is the method of choice. Rheumatoid factor should be adsorbed from the specimen before testing. With chronic infection, the titer to phase I antigen is usually much higher than that to phase II antigen (i.e., *C. burnetii* that has truncated lipopolysaccharide associated with gene deletions during laboratory passages), and the diagnosis should not be based on serology alone. Rather, the entire clinical setting must be taken into consideration. An anti-phase I IgG titer of ≥ 6400 would be considered a major criterion for the diagnosis of chronic Q fever, while a titer of ≥ 800 but ≤ 6400 would be a minor criterion. In acute Q fever, a fourfold rise in titer can be demonstrated between acute- and convalescent-phase serum samples.

Fluorodeoxyglucose positron emission tomography combined with CT (FDG-PET/CT) can be useful because it can detect not only valvular infection but also intravascular infection elsewhere as well as osteomyelitis.

serum doxycycline levels. A serum level-to-doxycycline MIC ratio of ≥ 1 is associated with a rapid decline in phase I antibodies with the doxycycline-hydroxychloroquine regimen. Patients treated with this regimen must be advised about photosensitivity and retinal toxicity risks. The doxycycline-hydroxychloroquine regimen was successful in one patient with HIV infection and Q fever endocarditis. The Jarisch-Herxheimer reaction occasionally complicates the treatment of chronic Q fever. Treatment of *C. burnetii*-infected aortic aneurysms is the same as that for Q fever endocarditis. Surgical intervention is often required.

If doxycycline-hydroxychloroquine cannot be used, the regimen chosen should include at least two antibiotics active against *C. burnetii*. Rifampin (300 mg once daily) combined with doxycycline (100 mg twice daily) or ciprofloxacin (750 mg twice daily) has been used successfully. The management of patients with Q fever endocarditis is complex and should preferably be undertaken by individuals with experience in managing this illness. Monitoring of antibody titers on a quarterly basis is an essential part of the management of these patients. Thus the laboratory should be contacted and asked to save all serum samples from such patients so that the current sample can be run with the previous one. There is incomplete agreement on the antibody titer at which therapy can be stopped. However, it is reasonable to discontinue treatment if IgG antibody levels have decreased by fourfold at 1 year, if IgM antibody to phase II has disappeared, and if the patient is clinically stable.

Patients with acute Q fever and lesions of native heart valves (e.g., bicuspid aortic valve), prosthetic valves, or prosthetic intravascular material should undergo serologic monitoring every 4 months for 2 years. If the phase I IgG titer is >800 , further investigation is warranted. Some authorities recommend that patients with valvulopathy and acute Q fever receive doxycycline and hydroxychloroquine to prevent chronic Q fever. For women who exhibit a serologic profile of chronic Q fever after childbirth, hydroxychloroquine and doxycycline should be given for 1 year.

BIOLOGIC MODIFYING AGENTS

Interferon γ was successful in the treatment of a 3-year-old boy with prolonged fever, abdominal pain, and thrombocytopenia due to *C. burnetii* that had not been eradicated with conventional antibiotic therapy. Many patients with granulomatous hepatitis due to Q fever have a prolonged febrile illness that is unresponsive to antibiotics. For these individuals, treatment with prednisone (0.5 mg/kg) has resulted in defervescence within 2–15 days. After defervescence, the glucocorticoid dose is tapered over the next month.

TREATMENT Q FEVER

ANTIBIOTICS

Treatment of acute Q fever with doxycycline (100 mg twice daily for 14 days) is usually successful. Quinolones also are effective. When Q fever is diagnosed during pregnancy, treatment with trimethoprim-sulfamethoxazole (TMP-SMX) is recommended for the duration of the pregnancy. One study showed no intrauterine fetal deaths and substantial reduction of obstetric complications in a group of Q fever patients treated with TMP-SMX.

The treatment of chronic Q fever is difficult and requires careful follow-up. Addition of hydroxychloroquine (to alkalinize the phagolysosome) renders doxycycline bactericidal against *C. burnetii*, and this combination is currently the favored regimen. Treatment with doxycycline (100 mg bid) and hydroxychloroquine (200 mg tid; plasma concentration maintained at 0.8–1.2 $\mu\text{g/mL}$) for 18 months is superior to a regimen of doxycycline and ofloxacin. Among 21 patients who received doxycycline and hydroxychloroquine, 1 died of a surgical complication, 2 were still being treated at the end of the study, 1 was still being evaluated, and 17 were cured. The mean duration of treatment was 31 months. In the ofloxacin and doxycycline group of 14 patients, 1 had died, 1 was still being treated, 7 had relapsed, and 5 had been cured by the end of the study. Optimal management of Q fever endocarditis entails determining the MIC of doxycycline for the patient's isolate and measuring

Prevention A whole-cell vaccine (Q-Vax) licensed in Australia effectively prevents Q fever in abattoir workers. Before administration of the vaccine, skin testing with intradermal diluted *C. burnetii* vaccine is performed, serologic testing is undertaken, and a history of possible Q fever is sought. Vaccine is given only to patients with no history of Q fever and negative results in serologic and skin testing.

Good animal-husbandry practices are important in preventing widespread contamination of the environment by *C. burnetii*. These practices include isolating aborting animals for up to 14 days, raising feed bunks to prevent contamination of feed by excreta, destroying aborted materials (by burning and burying fetal membranes and still-born animals), and wearing masks and gloves when handling aborted materials. Vaccination of sheep and goats and a culling program were effective in the Netherlands outbreak. Only seronegative pregnant animals should be used in research settings, and only seronegative animals should be permitted in petting zoos.

During an outbreak of Q fever and for 4 weeks after it ceases, blood donations should not be accepted from individuals who live in the affected area.

ACKNOWLEDGMENT

The contributions of Didier Raoult, MD, to this chapter in previous editions are gratefully acknowledged.

Mycoplasmas are prokaryotes of the class Mollicutes. Their size (150–350 nm) is closer to that of viruses than to that of bacteria. Unlike viruses, however, mycoplasmas grow in cell-free culture media; in fact, they are the smallest organisms capable of independent replication.



The entire genomes of many *Mycoplasma* species have been sequenced and have been found to be among the smallest of all prokaryotic genomes. Sequencing information for these genomes has helped define the minimal set of genes necessary for cellular life. The absence of genes related to the synthesis of amino acids, fatty acid metabolism, and cholesterol dictates the mycoplasmas' parasitic or saprophytic dependence on a host for exogenous nutrients and necessitates the use of complex fastidious media to culture these organisms. Mycoplasmas lack a cell wall and are bound only by a cell membrane. The absence of a cell wall explains the inactivity of β -lactam antibiotics (penicillins and cephalosporins) against infections caused by these organisms.

At least 13 *Mycoplasma* species, two *Acholeplasma* species, and two *Ureaplasma* species have been isolated from humans. Most of these species are thought to be normal inhabitants of oral and urogenital mucous membranes. Only four species—*M. pneumoniae*, *M. hominis*, *U. urealyticum*, and *U. parvum*—have been shown conclusively to be pathogenic in immunocompetent humans. *M. pneumoniae* primarily infects the respiratory tract, while *M. hominis*, *U. urealyticum*, and *U. parvum* are associated with a variety of genitourinary tract disorders and neonatal infections. Some data indicate that *M. genitalium* may be a cause of disease in humans. Other mycoplasmas may cause disease in immunocompromised persons.

MYCOPLASMA PNEUMONIAE

PATHOGENESIS

M. pneumoniae is generally thought to act as an extracellular pathogen. Although the organism has been shown to exist and replicate within human cells, it is not known whether these intracellular events contribute to the pathogenesis of disease. *M. pneumoniae* attaches to ciliated respiratory epithelial cells by means of a complex terminal organelle at the tip of one end of the organism. Cytoadherence is mediated by interactive adhesins and accessory proteins clustered on this organelle. After extracellular attachment, *M. pneumoniae* causes injury to host respiratory tissue. The mechanism of injury is thought to be mediated by the production of hydrogen peroxide and of a recently identified ADP-ribosylating and vacuolating cytotoxin of *M. pneumoniae* that has many similarities to pertussis toxin. Because mycoplasmas lack a cell wall, they also lack cell wall–derived stimulators of the innate immune system, such as lipopolysaccharide, lipoteichoic acid, and murein (peptidoglycan) fragments. However, lipoproteins from the mycoplasma cell membrane appear to have inflammatory properties, probably acting through Toll-like receptors (primarily TLR2) on macrophages and other cells. Lung biopsy specimens from patients with *M. pneumoniae* respiratory tract infection reveal an inflammatory process involving the trachea, bronchioles, and peribronchial tissue, with a monocytic infiltrate coinciding with a luminal exudate of polymorphonuclear leukocytes.

Experimental evidence indicates that innate immunity provides most of the host's defense against mycoplasma infection in the lungs, whereas cellular immunity may actually play an immunopathogenic role, exacerbating mycoplasma lung disease. Humoral immunity appears to provide protection against dissemination of *M. pneumoniae* infection; patients with humoral immunodeficiencies do not have more severe lung disease than do immunocompetent patients in the early stages of infection but more often develop disseminated infection resulting in syndromes such as arthritis, meningitis, and osteomyelitis. The immunity that follows severe *M. pneumoniae* infections is more protective and longer-lasting than that following mild infections.

Genuine second attacks of *M. pneumoniae* pneumonia have been reported infrequently.

EPIDEMIOLOGY

M. pneumoniae infection occurs worldwide. It is likely that the incidence of upper respiratory illness due to *M. pneumoniae* is up to 20 times that of pneumonia caused by this organism. Infection is spread from one person to another by respiratory droplets expectorated during coughing and results in clinically apparent disease in an estimated 80% of cases. The incubation period for *M. pneumoniae* is 2–4 weeks; therefore, the time-course of infection in a specific population may be several weeks long. Intrafamilial attack rates are as high as 84% among children and 41% among adults. Outbreaks of *M. pneumoniae* illness often occur in institutional settings such as military bases, boarding schools, and summer camps. Infections tend to be endemic, with sporadic epidemics every 4–7 years. There is no seasonal pattern.

Most significantly, *M. pneumoniae* is a major cause of community-acquired respiratory illness in both children and adults and is often grouped with *Chlamydia pneumoniae* and *Legionella* species as being among the most important bacterial causes of “atypical” community-acquired pneumonia. For community-acquired pneumonia in adults, *M. pneumoniae* is the most frequently detected “atypical” organism. Analysis of 13 studies of community-acquired pneumonia published since 1995 (which included 6207 ambulatory and hospitalized adults) showed that the overall prevalence of *M. pneumoniae* was 22.7%; by comparison, the prevalence of *C. pneumoniae* was 11.7%, and that of *Legionella* species was 4.6%. *M. pneumoniae* pneumonia is also referred to as Eaton agent pneumonia (the organism having first been isolated in the early 1940s by Monroe Eaton), primary atypical pneumonia, and “walking” pneumonia.

CLINICAL MANIFESTATIONS

Upper Respiratory Tract Infections and Pneumonia Acute *M. pneumoniae* infections generally manifest as pharyngitis, tracheobronchitis, reactive airway disease/wheezing, or a nonspecific upper respiratory syndrome. Little evidence supports the commonly held belief that this organism is an important cause of otitis media, with or without bullous myringitis. Pneumonia develops in 3–13% of infected individuals; its onset is usually gradual, occurring over several days, but may be more abrupt. Although *Mycoplasma pneumoniae* may begin with a sore throat, the most common presenting symptom is cough. The cough is typically nonproductive, but some patients produce sputum. Headache, malaise, chills, and fever are noted in the majority of patients.

On physical examination, wheezes or rales are detected in ~80% of patients with *M. pneumoniae* pneumonia. In many patients, however, pneumonia can be diagnosed only by chest radiography. The most common radiographic pattern is that of peribronchial pneumonia with thickened bronchial markings, streaks of interstitial infiltration, and areas of subsegmental atelectasis. Segmental or lobar consolidation is not uncommon. While clinically evident pleural effusions are infrequent, lateral decubitus views reveal that up to 20% of patients have pleural effusions.

Overall, the clinical presentation of pneumonia in an individual patient is not useful for differentiating *M. pneumoniae* pneumonia from other types of community-acquired pneumonia. The possibility of *M. pneumoniae* infection deserves particular consideration when community-acquired pneumonia fails to respond to treatment with a penicillin or a cephalosporin—antibiotics that are ineffective against mycoplasmas. Symptoms usually resolve within 2–3 weeks after the onset of illness. Although *M. pneumoniae* pneumonia is generally self-limited, appropriate antimicrobial therapy significantly shortens the duration of clinical illness. Infection uncommonly results in critical illness and only rarely in death. In some patients, long-term recurrent wheezing or reactive airway disease may follow the resolution of acute pneumonia. The significance of chronic infection, especially as it relates to asthma, is an area of active investigation.

Extrapulmonary Manifestations An array of extrapulmonary manifestations may develop during *M. pneumoniae* infection. The most significant are neurologic, dermatologic, cardiac, rheumatologic, and

1164 hematologic in nature. Extrapulmonary manifestations can be a result of disseminated infection, especially in patients with humoral immunodeficiencies (e.g., septic arthritis); postinfectious autoimmune phenomena (e.g., Guillain-Barré syndrome); or possibly ADP-ribosylating toxin. Overall, these manifestations are uncommon, given the frequency of *M. pneumoniae* infection. Notably, many patients with extrapulmonary *M. pneumoniae* disease do not have respiratory disease.

Skin eruptions described with *M. pneumoniae* infection include erythematous (macular or maculopapular), vesicular, bullous, petechial, and urticarial rashes. In some reports, 17% of patients with *M. pneumoniae* pneumonia have had an exanthem. Erythema multiforme major (Stevens-Johnson syndrome) is the most clinically significant skin eruption associated with *M. pneumoniae* infection; it appears to occur more commonly with *M. pneumoniae* than with other infectious agents.

A wide spectrum of neurologic manifestations has been reported with *M. pneumoniae* infection. The most common are meningoencephalitis, encephalitis, Guillain-Barré syndrome, and aseptic meningitis. *M. pneumoniae* has been implicated as a likely etiologic agent in 5–7% of cases of encephalitis. Other neurologic manifestations may include cranial neuropathy, acute psychosis, cerebellar ataxia, acute demyelinating encephalomyelitis, cerebrovascular thromboembolic events, and transverse myelitis.

Hematologic manifestations of *M. pneumoniae* infection include hemolytic anemia, aplastic anemia, cold agglutinins, disseminated intravascular coagulation, and hypercoagulopathy. When anemia does occur, it generally develops in the second or third week of illness.

In addition, hepatitis, glomerulonephritis, pancreatitis, myocarditis, pericarditis, rhabdomyolysis, and arthritis (septic and reactive) have been convincingly ascribed to *M. pneumoniae* infection. Septic arthritis has been described most commonly in hypogammaglobulinemic patients.

DIAGNOSIS

Clinical findings, nonmicrobiologic laboratory tests, and chest radiography are not useful for differentiating *M. pneumoniae* pneumonia from other types of community-acquired pneumonia. In addition, since *M. pneumoniae* lacks a cell wall, it is not visible on Gram's stain. Although of historical interest, the measurement of cold agglutinin titers is no longer recommended for the diagnosis of *M. pneumoniae* infection because the findings are nonspecific and assays specific for *M. pneumoniae* are now available.

Acute *M. pneumoniae* infection can be diagnosed by polymerase chain reaction (PCR) detection of the organism in respiratory tract secretions or by isolation of the organism in culture (Table 212-1). Oropharyngeal, nasopharyngeal, and pulmonary specimens are all acceptable for diagnosing *M. pneumoniae* pneumonia. Other bodily fluids, such as cerebrospinal fluid, are acceptable for extrapulmonary infection. *M. pneumoniae* culture (which requires special media) is not recommended for routine diagnosis because the organism may take weeks to grow and is often difficult to isolate from clinical specimens. In contrast, PCR allows rapid, specific diagnosis earlier in the course of clinical illness.


The diagnosis can also be established by serologic tests for IgM and IgG antibodies to *M. pneumoniae* in paired (acute- and convalescent-phase) serum samples; enzyme-linked immunoassay is

the recommended serologic method. An acute-phase sample alone is not adequate for diagnosis, as antibodies to *M. pneumoniae* may not develop until 2 weeks into the illness; therefore, it is important to test paired samples. In addition, IgM antibody to *M. pneumoniae* can persist for up to 1 year after acute infection. Thus its presence may indicate recent rather than acute infection.

The combination of PCR of respiratory tract secretions and serologic testing constitutes the most sensitive and rapid approach to the diagnosis of *M. pneumoniae* infection.

TREATMENT MYCOPLASMA PNEUMONIAE INFECTIONS

Although in the majority of untreated cases symptoms resolve within 2–3 weeks without significant associated morbidity, *M. pneumoniae* pneumonia can be a serious illness that responds to appropriate antimicrobial therapy (Table 212-2). Randomized, double-blind, placebo-controlled trials in adults have demonstrated that antimicrobial treatment significantly decreases the duration of fever, cough, malaise, hospitalization, and radiologic abnormalities in *M. pneumoniae* pneumonia. Treatment options for acute *M. pneumoniae* infection include macrolides (e.g., oral azithromycin, 500 mg on day 1, then 250 mg/d on days 2–5), tetracyclines (e.g., oral doxycycline, 100 mg twice daily for 10–14 days), and respiratory fluoroquinolones. However, ciprofloxacin and ofloxacin are not recommended because of their high minimal inhibitory concentrations against *M. pneumoniae* isolates and their poor performance in experimental studies. A 10- to 14-day course of quinolone therapy appears adequate.

 In Japan and China, very high levels (up to ≥90%) of *M. pneumoniae* resistance to macrolides have been reported. In Europe and to a lesser degree in the United States, macrolide-resistant *M. pneumoniae* is emerging. In investigated outbreaks of respiratory illness due to *M. pneumoniae* in the United States, macrolide resistance has been reported in 8–27% of isolates. Clinical studies have demonstrated that, when treated with macrolides, patients with community-acquired pneumonia due to macrolide-resistant *M. pneumoniae* experience a significantly longer duration of symptoms than do patients infected with macrolide-sensitive organisms; thus macrolide resistance in *M. pneumoniae* does appear to have clinical significance. If macrolide resistance is prominent in a particular geographic locale or is suspected, then a nonmacrolide antibiotic should be considered for treatment; in addition, culture of *M. pneumoniae* may prove useful in these instances, providing an isolate for susceptibility testing.

Clinical observations and experimental data suggest that the addition of glucocorticoids to an antibiotic regimen may be of value for the treatment of severe or refractory *M. pneumoniae* pneumonia. However, relevant clinical experience is limited. Even though appropriate antibiotic therapy significantly reduces the duration of respiratory illness, it does not appear to shorten the duration of detection of *M. pneumoniae* by culture or PCR; therefore, a test of cure or eradication is not suggested.

The roles of antimicrobial drugs, glucocorticoids, and IV immunoglobulin in the treatment of neurologic disease due to *M. pneumoniae* remain unknown.

TABLE 212-2 ANTIMICROBIAL AGENTS OF CHOICE FOR MYCOPLASMA INFECTIONS^a

Organism	Drug(s)
<i>M. pneumoniae</i>	Azithromycin, clarithromycin, erythromycin, doxycycline, levofloxacin, moxifloxacin, gemifloxacin (not ciprofloxacin or ofloxacin)
<i>U. urealyticum</i> , <i>U. parvum</i>	Azithromycin, clarithromycin, erythromycin, doxycycline
<i>M. hominis</i>	Doxycycline, clindamycin
<i>M. genitalium</i>	Azithromycin

^aAntimicrobial resistance has been reported in mycoplasmas, as described in the text.

TABLE 212-1 DIAGNOSTIC TESTS FOR RESPIRATORY MYCOPLASMA PNEUMONIAE INFECTION^a

Test	Sensitivity, %	Specificity, %
Respiratory culture	≤60	100
Respiratory PCR	65–90	90–100
Serologic studies ^b	55–100	55–100

^aA combination of PCR and serology is suggested for routine diagnosis. If macrolide resistance is suspected, *M. pneumoniae* culture may prove useful, providing an isolate for susceptibility testing. ^bAcute- and convalescent-phase serum samples are recommended.

Abbreviation: PCR, polymerase chain reaction.

EPIDEMIOLOGY

M. hominis, *M. genitalium*, *U. urealyticum*, and *U. parvum* can cause urogenital tract disease. The significance of isolation of these organisms in a variety of other syndromes is unknown and in some cases is being investigated. *M. fermentans* has not been shown convincingly to cause human disease.

While urogenital mycoplasmas may be transmitted to a fetus during passage through a colonized birth canal, sexual contact is the major mode of transmission, and the risk of colonization increases dramatically with increasing numbers of sexual partners. In asymptomatic women, these mycoplasmas may be found throughout the lower urogenital tract. The vagina yields the largest number of organisms; next most densely colonized are the periurethral area and the cervix. Ureaplasmas are isolated less often from urine than from the cervix, but *M. hominis* is found with approximately the same frequency at these two sites. Ureaplasmas are isolated from the vagina of 40–80% of sexually active, asymptomatic women and *M. hominis* from 21–70%. The two microorganisms are found concurrently in 31–60% of women. In men, colonization with each organism is less prevalent. Mycoplasmas have been isolated from urine, semen, and the distal urethra of asymptomatic men.

CLINICAL MANIFESTATIONS

Urethritis, Pyelonephritis, and Urinary Calculi In many episodes of *Chlamydia*-negative nongonococcal urethritis, ureaplasmas may be the causative agent. These organisms may also cause chronic voiding symptoms in women. The common presence of ureaplasmas in the urethra of asymptomatic men suggests either that only certain serovars are pathogenic or that predisposing factors, such as lack of immunity, must exist in persons who develop symptomatic infection. Alternatively, disease may develop only upon initial exposure to ureaplasmas. Ureaplasmas have been implicated in epididymitis. *M. genitalium* also appears to cause urethritis. *M. genitalium* and ureaplasmas do not have a known role in prostatitis. *M. hominis* does not appear to play a primary etiologic role in urethritis, epididymitis, or prostatitis.

Evidence suggests that *M. hominis* causes up to 5% of cases of acute pyelonephritis. Ureaplasmas have not been associated with this disease.

Ureaplasmas play a limited role in the production of urinary calculi. The frequency with which ureaplasmas reach the kidney, the predisposing factors that allow them to do so, and the relative frequency of urinary tract calculi induced by this organism (compared with other organisms) are not known.

Pelvic Inflammatory Disease *M. hominis* can cause pelvic inflammatory disease. In most episodes, *M. hominis* occurs as part of a polymicrobial infection, but the organism may play an independent role in a limited number of cases. Some data also support an association of *M. genitalium* with pelvic inflammatory disease. Ureaplasmas are not thought to cause pelvic inflammatory disease.

Postpartum and Postabortal Infection Studies implicate *M. hominis* as the primary pathogen in ~5–10% of women who have postpartum or postabortal fever; ureaplasmas have been implicated to a lesser degree. These infections are generally self-limited; however, if symptoms persist, specific antimicrobial therapy should be given. Ureaplasmas also appear to play a role in occasional postcesarean wound infections.

Non-urogenital Infection In rare instances, *M. hominis* causes non-urogenital infections, such as brain abscess, wound infection, post-sternotomy mediastinitis, endocarditis, and neonatal meningitis. These infections are most common among immunocompromised and hypogammaglobulinemic patients. Ureaplasmas and *M. hominis* can cause septic arthritis in immunodeficient patients. Ureaplasmas probably cause neonatal pneumonitis; their significant role in the development of bronchopulmonary dysplasia—the chronic lung disease of premature infants—has been documented in a number of studies. It is unclear whether ureaplasmas and *M. hominis* cause

infertility, spontaneous abortion, premature labor, low birth weight, or chorioamnionitis. 1165

DIAGNOSIS

Culture and PCR are both appropriate methods for the isolation of urogenital mycoplasmas. Culture of these organisms, however, requires special techniques and media that generally are available only at larger medical centers and reference laboratories. Serologic testing is not recommended for the clinical diagnosis of urogenital *Mycoplasma* infections.

TREATMENT UROGENITAL MYCOPLASMA INFECTIONS

Because colonization with urogenital mycoplasmas is common, it appears at present that their isolation from the urogenital tract in the absence of disease generally does not warrant treatment. Macrolides and doxycycline are considered the antimicrobial agents of choice for *Ureaplasma* infections (Table 212-2). *Ureaplasma* resistance to macrolides, doxycycline, quinolones, and chloramphenicol has been reported. *M. hominis* is resistant to macrolides. Doxycycline is generally the drug of choice for *M. hominis* infections, although resistance has been reported. Clindamycin is generally active against *M. hominis*. Quinolones are active in vitro against *M. hominis*. For *M. genitalium*, the agent of choice appears to be azithromycin; treatment failures have been reported with other macrolides as well as with quinolones.

213 Chlamydial Infections

Charlotte A. Gaydos, Thomas C. Quinn

Chlamydiae are obligate intracellular bacteria that cause a wide variety of diseases in humans and animals.

ETIOLOGIC AGENTS

The chlamydiae were originally classified as four species in the genus *Chlamydia*: *C. trachomatis*, *C. pneumoniae*, *C. psittaci*, and *C. pecorum* (the last species being found in ruminants). The *C. psittaci* group has been separated into three species: *C. psittaci*, *C. felis*, and *C. abortus*. The mouse pneumonitis strain (MoPn) is now classified as *C. muridarum*, and the guinea pig inclusion conjunctivitis strain (GPIC) is now designated *C. caviae*.

C. trachomatis is divided into two biovars: trachoma and LGV (lymphogranuloma venereum). The trachoma biovar causes two major types of disease in humans: ocular trachoma, the leading infectious cause of preventable blindness in the developing world; and urogenital infections, which are sexually or neonatally transmitted. The 18 serovars of *C. trachomatis* fall into three groups: the trachoma serovars A, B, Ba, and C; the oculogenital serovars D–K; and the LGV serovars L₁–L₃. Serovars can be distinguished by serologic typing with monoclonal antibodies or by molecular gene typing. However, serovar identification usually is not important clinically since the antibiotic susceptibility pattern is the same for all three groups. The one exception applies when LGV is suspected on clinical grounds; in this situation, serovar determination is important because a longer treatment duration is required for LGV strains.

BIOLOGY, GROWTH CYCLE, AND PATHOGENESIS

BIOLOGY

During their intracellular growth, chlamydiae produce characteristic intracytoplasmic inclusions that can be visualized by direct fluorescent antibody (DFA) or Giemsa staining of infected clinical material, such as conjunctival scrapings or cervical or urethral epithelial

1166 cells. Chlamydiae are nonmotile, gram-negative, obligate intracellular bacteria that replicate within the cytoplasm of host cells, forming the characteristic membrane-bound inclusions that are the basis for some diagnostic tests. Originally considered to be large viruses, chlamydiae differ from viruses in possessing RNA and DNA as well as a cell wall that is quite similar in structure to the cell wall of typical gram-negative bacteria. However, chlamydiae lack peptidoglycan; their structural integrity depends on disulfide binding of outer-membrane proteins.

GROWTH CYCLE

Among the defining characteristics of chlamydiae is a unique growth cycle that involves alternation between two highly specialized morphologic forms (Figs. 213-1 and 213-2): the elementary body (EB), which is the infectious form and is specifically adapted for extracellular survival, and the metabolically active and replicating reticulate body (RB), which is not infectious, is adapted for an intracellular environment, and does not survive well outside the host cell. The biphasic growth cycle begins with attachment of the EB (diameter, 0.25–0.35 μm) at specific sites on the surface of the host cell. The EB enters the cell through a process similar to receptor-mediated endocytosis and resides in an inclusion, where the entire growth cycle is completed. The chlamydiae prevent phagosome-lysosome fusion. The inclusion membrane is modified by insertion of chlamydial antigens. Once the EB has entered the cell, it reorganizes into an RB, which is larger (0.5–1 μm) and contains more RNA. After ~8 h, the RB starts to divide by binary fission. The intracytoplasmic, membrane-bound inclusion body containing the RBs increases in size as the RBs multiply. Approximately 18–24 h after infection of the cell, these RBs begin to become EBs by a reorganization or condensation process that is poorly understood. After rupture of the inclusion body, the EBs are released to initiate another cycle of infection.

Chlamydiae are susceptible to many broad-spectrum antibiotics and possess a number of enzymes, but they have a very restricted

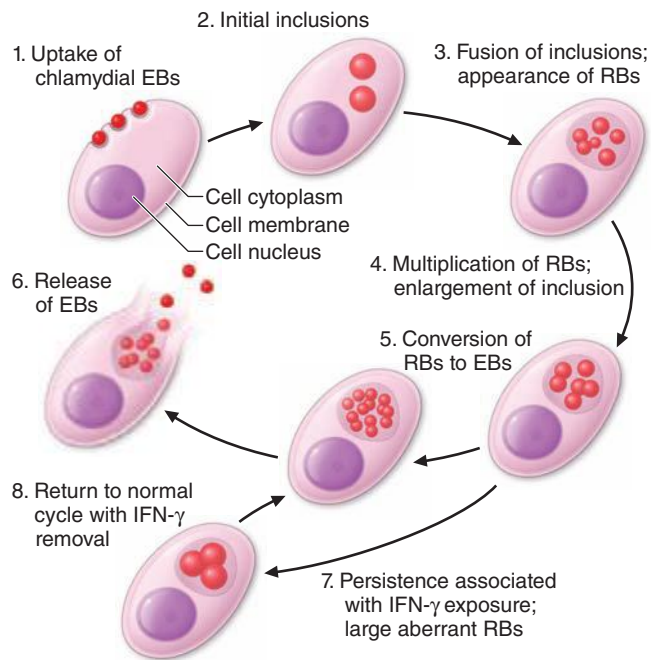


FIGURE 213-2 Chlamydial life cycle. EBs, elementary bodies; RBs, reticulate bodies; IFN- γ , interferon γ . (Reprinted with permission from WE Stamm: Chlamydial infections, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al [eds]. New York, McGraw-Hill, 2008, p 1071.)

metabolic capacity. None of these metabolic reactions results in the production of energy. Chlamydiae have thus been considered to be energy parasites that use the ATP produced by the host cell for their own metabolic functions. Many aspects of chlamydial molecular biology are not well understood, but the sequencing of several chlamydial genomes and new proteomics research have provided researchers with many relevant tools for elucidating the biology of the life cycle.

PATHOGENESIS

Genital infections are mostly caused by *C. trachomatis* serovars D–K, with serovars D, E, and F involved most often. Molecular typing of the major outer-membrane protein gene (*omp1*) from which serovar differences arise has been used to demonstrate that polymorphisms can occur in isolates from patients who are exposed frequently to multiple infections, while less variation is observed in isolates from less sexually active populations. Polymorphisms in the major outer-membrane protein may provide antigenic variation, and the different forms allow persistence in the community because immunity to one is not protective against the others.

The trachoma biovar is essentially a parasite of squamocolumnar epithelial cells; the LGV biovar is more invasive and involves lymphoid cells. As is typical of chlamydiae, *C. trachomatis* strains are capable of causing chronic, clinically inapparent, asymptomatic infections. Because the duration of the chlamydial growth cycle is ~48–72 h, the incubation period of sexually transmitted chlamydial infections is relatively long—generally 1–3 weeks. *C. trachomatis* causes cell death as a result of its replicative cycle and can induce cell damage whenever it persists. However, few toxic effects are demonstrated, and cell death because of chlamydial replication is not sufficient to account for disease manifestations, the majority of which are due to immunopathologic mechanisms or nonspecific host responses to the organism or its byproducts.

In recent years, the entire genomes of various chlamydial species have been sequenced, the field of proteomics has become established, host innate immunity has been more precisely delineated, and innovative host cell–chlamydial interaction studies have been conducted. As a result, many insights have been gained into how chlamydiae adapt and replicate in their intracellular environment and produce disease. These insights into pathogenesis include information

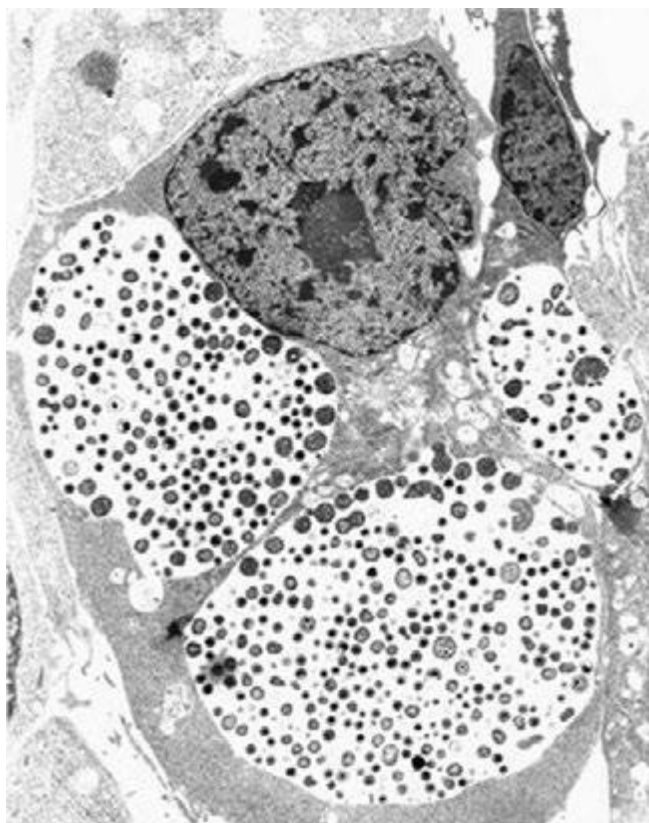


FIGURE 213-1 Chlamydial intracellular inclusions filled with smaller dense elementary bodies and larger reticulate bodies. (Reprinted with permission from WE Stamm: Chlamydial infections, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al [eds]. New York, McGraw-Hill, 2008, p 1070.)

on the regulation of gene expression, protein localization, the type III secretion system, the roles of CD4+ and CD8+ T lymphocytes in the host response, and T lymphocyte trafficking.

The chlamydial heat-shock protein, which shares antigenic epitopes with similar proteins of other bacteria and with human heat-shock protein, may sensitize the host, and repeated infections may cause host cell damage. Persistent or recurrent chlamydial infections are associated with fibrosis, scarring, and complications following simple epithelial infections. A common endpoint of these late consequences is scarring of mucous membranes. Genital complications can lead to pelvic inflammatory disease (PID) and its late consequences of infertility, ectopic pregnancy, and chronic pelvic pain, while ocular infections may lead to blinding trachoma. High levels of antibody to human heat-shock protein have been associated with tubal factor infertility and ectopic pregnancy. Without adequate therapy, chlamydial infections may persist for several years, although symptoms—if present—usually abate.

The pathogenic mechanisms of *C. pneumoniae* have yet to be completely elucidated. The same is true for *C. psittaci*, except that this agent infects cells very efficiently and causes disease that may reflect direct cytopathic effects.

CHLAMYDIA TRACHOMATIS INFECTIONS

GENITAL INFECTIONS

Spectrum Although chlamydiae cause a number of human diseases, localized lower genital tract infections caused by *C. trachomatis* and the sequelae of such infections are the most important in terms of medical and economic impact. Oculogenital infections due to *C. trachomatis* serovars D–K are transmitted during sexual contact or from mother to baby during childbirth and are associated with many syndromes, including cervicitis, salpingitis, acute urethral syndrome, endometritis, ectopic pregnancy, infertility, and PID in female patients; urethritis, proctitis, and epididymitis in male patients; and conjunctivitis and pneumonia in infants. Women bear the greatest burden of morbidity because of the serious sequelae of these infections. Untreated infections lead to PID, and multiple episodes of PID can lead to tubal factor infertility and chronic pelvic pain. Studies estimate that up to 80–90% of women and >50% of men with *C. trachomatis* genital infections lack symptoms; other patients have very mild symptoms. Thus a large reservoir of infected persons continues to transmit infection to sexual partners.

As their designations reflect, the LGV serovars (L_1 , L_2 , and L_3) cause LGV, an invasive sexually transmitted disease (STD) characterized by acute lymphadenitis with bubo formation and/or acute hemorrhagic proctitis (see “Lymphogranuloma Venereum,” below).



Epidemiology *C. trachomatis* genital infections are global in distribution. The World Health Organization (WHO) estimated in 2008 that >106.4 million cases occur annually worldwide. This figure makes chlamydial infection the most prevalent sexually transmitted bacterial infection in the world. The associated morbidity is substantial, and economic costs are high.

In the United States, chlamydial infections are the most commonly reported of all infectious diseases. In 2012, 1.3 million cases were reported to the U.S. Centers for Disease Control and Prevention (CDC); however, the CDC estimates that 2–3 million new cases occur per year, with substantial underreporting due to lack of screening in some populations. Rates of infection have increased every year; higher rates among women than among men reflect the focus on expansion of screening programs for women during the past 20 years, the use of increasingly sensitive diagnostic tests, an increased emphasis on case reporting, and improvements in the information systems used for reporting. The CDC and other professional organizations recommend annual screening of all sexually active women ≤ 25 years of age as well as rescreening of previously infected individuals at 3 months. Young women have the highest infection rates; in 2012, the figures were 3416.5 and 3722.5 cases per 100,000 population at 15–19 and 20–24 years of age, respectively. Age-specific rates among men, while much lower than those among women, were highest in the 20- to

24-year-old age group, at 1343.3 cases per 100,000. In 2012, rates increased for all racial and ethnic groups, with the highest rates among African Americans. For example, the rate of chlamydial infection among African-American girls 15–19 years of age was 7507.1 cases per 100,000—almost six times the rate among Caucasian girls in the same age group (1301.5/100,000). The rate among African-American women 20–24 years old was 4.8 times the rate among Caucasian women in the same age group. Similar racial disparities in reported rates of chlamydial infection exist among men. For boys 15–19 years of age, the rate among African Americans was 11.1 times the rate among Caucasians. The rate among Native Americans/Alaska Natives was more than four times the rate among Caucasians (648.3), and the rate among Latinos (383.6) was two times higher than that among Caucasians. These disparities are important reflections of health inequities in the United States.

The above statistics are based on case reporting. Studies based on screening surveys estimate that the U.S. prevalence of *C. trachomatis* cervical infection is 5% among asymptomatic female college students and prenatal patients, >10% among women seen in family planning clinics, and >20% among women seen in STD clinics. The prevalence of genital *C. trachomatis* infections varies substantially by geographic locale, with the highest rates in the southeastern United States. However, asymptomatic infections have been detected in >8–10% of young female military recruits from all parts of the country. The prevalence of *C. trachomatis* in the cervix of pregnant women is 5–10 times higher than that of *Neisseria gonorrhoeae*. The prevalence of genital infection with either agent is highest among women who are between the ages of 18 and 24, single, and non-Caucasian (e.g., African-American, Latina, Asian, Pacific Islander). Infections recur frequently in these same risk groups and are often acquired from untreated sexual partners. The use of oral contraception and the presence of cervical ectopy also confer an increased risk. The proportion of infections that are asymptomatic appears to be higher for *C. trachomatis* than for *N. gonorrhoeae*, and symptomatic *C. trachomatis* infections are clinically less severe. Mild or asymptomatic *C. trachomatis* infections of the fallopian tubes nonetheless cause ongoing tubal damage and infertility. The costs of *C. trachomatis* infections and their complications to the U.S. health care system have recently been estimated to exceed \$516.7 million annually.

Clinical Manifestations • NONGONOCOCCAL AND POSTGONOCOCCAL URETHRITIS

C. trachomatis is the most common cause of nongonococcal urethritis (NGU) and postgonococcal urethritis (PGU). The designation PGU refers to NGU developing in men 2–3 weeks after treatment of gonococcal urethritis with single doses of agents such as penicillin or cephalosporins, which lack antimicrobial activity against chlamydiae. Since current treatment regimens for gonorrhea have evolved and now include combination therapy with tetracycline, doxycycline, or azithromycin—all of which are effective against concomitant chlamydial infection—both the incidence of PGU and the causative role of *C. trachomatis* in this syndrome have declined.

In the United States, most of the estimated 2 million cases of acute urethritis are NGU, and *C. trachomatis* is implicated in 30–50% of these cases. The cause of most of the remaining cases of NGU is uncertain, but recent evidence suggests that *Ureaplasma urealyticum*, *Mycoplasma genitalium*, *Trichomonas vaginalis*, and herpes simplex virus (HSV) cause some cases. The rate of involvement of *C. trachomatis* in urethral infection ranges from 3–7% among asymptomatic men to 15–20% among symptomatic men attending STD clinics. A multisite study of men in Baltimore, Seattle, Denver, and San Francisco reported an overall chlamydial prevalence of 7% in urine samples assessed by nucleic acid amplification tests (NAATs). As in women, infection in men is age related, with young age as the greatest risk factor for chlamydial urethritis. The prevalence among men is highest at 20–24 years of age. In STD clinics, urethritis is usually less prevalent among men who have sex with men (MSM) than among heterosexual men and is almost always much more common among African-American men than among Caucasian men. One study reported prevalences of 19% and 9% among nonwhite and white heterosexual men, respectively.

NGU is diagnosed by documentation of a leukocytic urethral exudate and by exclusion of gonorrhea by Gram's staining or culture. *C. trachomatis* urethritis is generally less severe than gonococcal urethritis, although in any individual patient these two forms of urethritis cannot reliably be differentiated solely on clinical grounds. Symptoms include urethral discharge (often whitish and mucoid rather than frankly purulent), dysuria, and urethral itching. Physical examination may reveal meatal erythema and tenderness as well as a urethral exudate that is often demonstrable only by stripping of the urethra.

At least one-third of male patients with *C. trachomatis* urethral infection have no evident signs or symptoms of urethritis. The availability of NAATs for first-void urine specimens has facilitated broader-based testing for asymptomatic infection in male patients. As a result, asymptomatic chlamydial urethritis has been demonstrated in 5–10% of sexually active male adolescents screened at school-based clinics or community centers. Such patients generally have pyuria (≥ 15 leukocytes per 400 \times microscopic field in the sediment of first-void urine), a positive leukocyte esterase test, or an increased number of leukocytes on a Gram-stained smear prepared from a urogenital swab inserted 1–2 cm into the anterior urethra. To differentiate between true urethritis and functional symptoms in symptomatic patients or to make a presumptive diagnosis of *C. trachomatis* infection in high-risk but asymptomatic men (e.g., male patients in STD clinics, sex partners of women with nongonococcal salpingitis or mucopurulent cervicitis, fathers of children with inclusion conjunctivitis), the examination of an endourethral specimen for increased leukocytes is useful if specific diagnostic tests for chlamydiae are not available. Alternatively, urethritis can be assayed noninvasively by examination of a first-void urine sample for pyuria, either by microscopy or by the leukocyte esterase test. Urine (or a urethral swab) can also be tested directly for chlamydiae by DNA amplification methods, as described below (see "Detection Methods").

EPIDIDYMITIS Chlamydial urethritis may be followed by acute epididymitis, but this condition is rare, generally occurring in sexually active patients <35 years of age; in older men, epididymitis is usually associated with gram-negative bacterial infection and/or instrumentation procedures. It is estimated that 50–70% of cases of acute epididymitis are caused by *C. trachomatis*. The condition usually presents as unilateral scrotal pain with tenderness, swelling, and fever in a young man, often occurring in association with chlamydial urethritis. The illness may be mild enough to treat with oral antibiotics on an outpatient basis or severe enough to require hospitalization and parenteral therapy. Testicular torsion should be excluded promptly by radionuclide scan, Doppler flow study, or surgical exploration in a teenager or young adult who presents with acute unilateral testicular pain without urethritis. The possibility of testicular tumor or chronic infection (e.g., tuberculosis) should be excluded when a patient with unilateral intrascrotal pain and swelling does not respond to appropriate antimicrobial therapy.

REACTIVE ARTHRITIS Reactive arthritis consists of conjunctivitis, urethritis (or, in female patients, cervicitis), arthritis, and characteristic mucocutaneous lesions. It may develop in 1–2% of cases of NGU and is thought to be the most common type of peripheral inflammatory arthritis in young men. *C. trachomatis* has been recovered from the urethra of 16–44% of patients with reactive arthritis and from 69% of men who have signs of urogenital inflammation at the time of examination. Antibodies to *C. trachomatis* have also been detected in 46–67% of patients with reactive arthritis, and *Chlamydia*-specific cell-mediated immunity has been documented in 72%. In addition, *C. trachomatis* has been isolated from synovial biopsy samples from 15 of 29 patients in a number of small series and from a smaller proportion of synovial fluid specimens. Chlamydial nucleic acids have been identified in synovial membranes and chlamydial EBs in joint fluid. The pathogenesis of reactive arthritis is unclear, but this condition probably represents an abnormal host response to a number of infectious agents, including those associated with bacterial gastroenteritis (e.g., *Salmonella*, *Shigella*, *Yersinia*, or *Campylobacter*), or to infection with *C. trachomatis* or *N. gonorrhoeae*. Since >80% of affected patients have

the HLA-B27 phenotype and since other mucosal infections produce an identical syndrome, chlamydial infection is thought to initiate an aberrant hyperactive immune response that produces inflammation of the involved target organs in these genetically predisposed individuals. Evidence of exaggerated cell-mediated and humoral immune responses to chlamydial antigens in reactive arthritis supports this hypothesis. The finding of chlamydial EBs and DNA in joint fluid and synovial tissue from patients with reactive arthritis suggests that chlamydiae may actually spread from genital to joint tissues in these patients—perhaps in macrophages.

NGU is the initial manifestation of reactive arthritis in 80% of patients, typically occurring within 14 days after sexual exposure. The urethritis may be mild and may even go unnoticed by the patient. Similarly, gonococcal urethritis may precede reactive arthritis, but coinfection with an agent of NGU is difficult to rule out. The urethral discharge may be purulent or mucopurulent, and patients may or may not report dysuria. Accompanying prostatitis, usually asymptomatic, has been described. Arthritis usually begins ~4 weeks after the onset of urethritis but may develop sooner or, in a small percentage of cases, may actually precede urethritis. The knees are most frequently involved; next most commonly affected are the ankles and small joints of the feet. Sacroiliitis, either symmetrical or asymmetrical, is documented in two-thirds of patients. Mild bilateral conjunctivitis, iritis, keratitis, or uveitis is sometimes present but lasts for only a few days. Finally, dermatologic manifestations occur in up to 50% of patients. The initial lesions—usually papules with a central yellow spot—most often involve the soles and palms and, in ~25% of patients, eventually epithelialize and thicken to produce keratoderma blenorrhagicum. Circinate balanitis is usually painless and occurs in fewer than half of patients. The initial episode of reactive arthritis usually lasts 2–6 months.

PROCTITIS Primary anal or rectal infections with *C. trachomatis* have been described in women and MSM who practice anal intercourse. In these infections, rectal involvement is initially characterized by severe anorectal pain, a bloody mucopurulent discharge, and tenesmus. Oculogenital serovars D–K and LGV serovars L₁, L₂, and L₃ have been found to cause proctitis. The LGV serovars are far more invasive and cause much more severely symptomatic disease, including severe ulcerative proctocolitis that can be clinically confused with HSV proctitis. Histologically, LGV proctitis may resemble Crohn's disease in that giant cell formation and granulomas are detected. In the United States and Europe, cases of LGV proctitis occur almost exclusively in MSM, many of whom are positive for HIV infection.

The less invasive non-LGV serovars of *C. trachomatis* cause mild proctitis. Many infected individuals are asymptomatic, and in these cases infection is diagnosed only by routine culture or NAAT of rectal swabs. The number of fecal leukocytes is usually abnormal in both asymptomatic and symptomatic cases. Sigmoidoscopy may yield normal findings or may reveal mild inflammatory changes or small erosions or follicles in the lower 10 cm of the rectum. Histologic examination of rectal biopsies generally shows anal crypts and prominent follicles as well as neutrophilic infiltration of the lamina propria. Chlamydial proctitis is best diagnosed by isolation of *C. trachomatis* from the rectum and documentation of a response to appropriate therapy. NAATs are reportedly more sensitive than culture for diagnosis and are also specific.

MUCOPURULENT CERVICITIS Although many women with chlamydial infections of the cervix have no symptoms, almost half generally have local signs of infection on examination. Cervicitis is usually characterized by the presence of a mucopurulent discharge, with >20 neutrophils per microscopic field visible in strands of cervical mucus in a thinly smeared, gram-stained preparation of endocervical exudate. Hypertrophic ectopy of the cervix may also be evident as an edematous area near the cervical os that is congested and bleeds easily on minor trauma (e.g., when a specimen is collected with a swab). A Papanicolaou smear shows increased numbers of neutrophils as well as a characteristic pattern of mononuclear inflammatory cells including plasma cells, transformed lymphocytes, and histiocytes. Cervical biopsy shows a predominantly mononuclear cell infiltrate of

the subepithelial stroma. Clinical experience and collaborative studies indicate that a cutoff of >30 polymorphonuclear neutrophils (PMNs) per 1000× field in a gram-stained smear of cervical mucus correlates best with chlamydial or gonococcal cervicitis.

Clinical recognition of chlamydial cervicitis depends on a high index of suspicion and careful cervical examination. No genital symptoms are specifically correlated with chlamydial cervical infection. The differential diagnosis of a mucopurulent discharge from the endocervical canal in a young, sexually active woman includes gonococcal endocervicitis, salpingitis, endometritis, and intrauterine contraceptive device–induced inflammation. Diagnosis of cervicitis is based on the presence of PMNs on a cervical swab as noted above; the presence of chlamydiae is confirmed by either culture or NAAT.

PELVIC INFLAMMATORY DISEASE Inflammation of sections of the fallopian tube is often referred to as salpingitis or PID. The proportion of acute salpingitis cases caused by *C. trachomatis* varies geographically and with the population studied. It has been estimated that *C. trachomatis* causes up to 50% of PID cases in the United States. PID occurs via ascending intraluminal spread of *C. trachomatis* or *N. gonorrhoeae* from the lower genital tract. Mucopurulent cervicitis is often followed by endometritis, endosalpingitis, and finally pelvic peritonitis. Evidence of mucopurulent cervicitis is often found in women with laparoscopically verified salpingitis. Similarly, endometritis, demonstrated by an endometrial biopsy showing plasma cell infiltration of the endometrial epithelium, is documented in most women with laparoscopy-verified chlamydial (or gonococcal) salpingitis. Chlamydial endometritis can also occur in the absence of clinical evidence of salpingitis. Histologic evidence of endometritis has been correlated with a syndrome consisting of vaginal bleeding, lower abdominal pain, and uterine tenderness in the absence of adnexal tenderness. Chlamydial salpingitis produces milder symptoms than gonococcal salpingitis and may be associated with less marked adnexal tenderness. Thus, mild adnexal or uterine tenderness in a sexually active woman with cervicitis suggests chlamydial PID.

Chronic untreated endometrial and tubal inflammation can result in tubal scarring, impaired tubal function, tubal occlusion, and infertility, even among women who report no prior treatment for PID. *C. trachomatis* has been implicated particularly often in “subclinical” PID on the basis of (1) a lack of history of PID among *Chlamydia*-seropositive women with tubal damage or (2) detection of chlamydial DNA or antigen among asymptomatic women with tubal infertility. These data suggest that the best method to prevent PID and its sequelae is surveillance and control of lower genital tract infections along with diagnosis and treatment of sex partners and prevention of reinfections. Promotion of early symptom recognition and health care presentation may reduce the frequency and severity of sequelae of PID.

PERIHEPATITIS The Fitz-Hugh–Curtis syndrome was originally described as a complication of gonococcal PID. However, studies over the past several decades have suggested that chlamydial infection is more commonly associated with perihepatitis than is *N. gonorrhoeae*. Perihepatitis should be suspected in young, sexually active women who develop right-upper-quadrant pain, fever, or nausea. Evidence of salpingitis may or may not be found on examination. Frequently, perihepatitis is strongly associated with extensive tubal scarring, adhesions, and inflammation observed at laparoscopy, and high titers of antibody to the 57-kDa chlamydial heat-shock protein have been documented. Culture and/or serologic evidence of *C. trachomatis* is found in three-fourths of women with this syndrome.

URETHRAL SYNDROME IN WOMEN In the absence of infection with uropathogens such as coliforms or *Staphylococcus saprophyticus*, *C. trachomatis* is the pathogen most commonly isolated from college women with dysuria, frequency, and pyuria. Screening studies can recover *C. trachomatis* from both the cervix and the urethra; in up to 25% of infected women, the organism is isolated only from the urethra. The urethral syndrome in women consists of dysuria and frequency in conjunction with chlamydial urethritis, pyuria, and no bacteriuria or urinary pathogens. Although symptoms of the urethral syndrome

may develop in some women with chlamydial infection, the majority of women attending STD clinics for urethral chlamydial infection do not have dysuria or frequency. Even in women with chlamydial urethritis causing the acute urethral syndrome, signs of urethritis such as urethral discharge, meatal redness, and swelling are uncommon. However, mucopurulent cervicitis in a woman presenting with dysuria and frequency strongly suggests *C. trachomatis* urethritis. Other correlates of chlamydial urethral syndrome include a duration of dysuria of >7–10 days, lack of hematuria, and lack of suprapubic tenderness. Abnormal urethral Gram’s stains showing >10 PMNs per 1000× field in women with dysuria but without coliform bacteriuria support the diagnosis of chlamydial urethritis. Other possible diagnoses include gonococcal or trichomonal infection of the urethra.

INFECTION IN PREGNANCY AND THE NEONATAL PERIOD Infections during pregnancy can be transmitted to infants during delivery. Approximately 20–30% of infants exposed to *C. trachomatis* in the birth canal develop conjunctivitis, and 10–15% subsequently develop pneumonia. Consequently, all newborn infants receive ocular prophylaxis at birth to prevent ophthalmia neonatorum. Without treatment, conjunctivitis usually develops at 5–19 days of life and often results in a profuse mucopurulent discharge. Roughly half of infected infants develop clinical evidence of inclusion conjunctivitis. However, it is impossible to differentiate chlamydial conjunctivitis from other forms of neonatal conjunctivitis (e.g., that due to *N. gonorrhoeae*, *Haemophilus influenzae*, *Streptococcus pneumoniae*, or HSV) on clinical grounds; thus laboratory diagnosis is required. Inclusions within epithelial cells are often detected in Giemsa-stained conjunctival smears, but these smears are considerably less sensitive than cultures or NAATs for chlamydiae. Gram-stained smears may show gonococci or occasional small gram-negative coccobacilli in *Haemophilus* conjunctivitis, but smears should be accompanied by cultures or NAATs for these agents.

C. trachomatis has also been isolated frequently and persistently from the nasopharynx, rectum, and vagina of infected infants—occasionally for >1 year in the absence of treatment. In some cases, otitis media results from perinatally acquired chlamydial infection. Pneumonia may develop in infants from 2 weeks to 4 months of age. *C. trachomatis* is estimated to cause 20–30% of pneumonia cases in infants <6 months of age. Epidemiologic studies have linked chlamydial pulmonary infection in infants with increased occurrence of subacute lung disease (bronchitis, asthma, wheezing) in later childhood.



LYMPHOGRANULOMA VENEREUM *C. trachomatis* serovars L₁, L₂, and L₃ cause LGV, an invasive systemic STD. The peak incidence of LGV corresponds with the age of greatest sexual activity: the second and third decades of life. The worldwide incidence of LGV is falling, but the disease is still endemic and a major cause of morbidity in parts of Asia, Africa, South America, and the Caribbean. LGV is rare in industrialized countries; for more than a decade, the reported incidence in the United States has been only 0.1 case per 100,000 population. In the Bahamas, an apparent outbreak of LGV was described in association with a concurrent increase in heterosexual infection with HIV. Reports of outbreaks with the newly identified variant L_{2b} in Europe, Australia, and the United States indicate that LGV is becoming more prevalent among MSM. These cases have usually presented as hemorrhagic proctocolitis in HIV-positive men. More widespread use of NAATs for identification of rectal infections may have enhanced case recognition.

LGV begins as a small painless papule that tends to ulcerate at the site of inoculation, often escaping attention. This primary lesion heals in a few days without scarring and is usually recognized as LGV only in retrospect. LGV strains of *C. trachomatis* have occasionally been recovered from genital ulcers and from the urethra of men and the endocervix of women who present with inguinal adenopathy; these areas may be the primary sites of infection in some cases. Proctitis is more common among people who practice receptive anal intercourse, and an elevated white blood cell count in anorectal smears may predict LGV in these patients. Ulcer formation may facilitate transmission of HIV infection and other sexually transmitted and blood-borne diseases.

As NAATs for *C. trachomatis* are being used more often, increasing numbers of cases of LGV proctitis are being recognized in MSM. Such patients present with anorectal pain and mucopurulent, bloody rectal discharge. Sigmoidoscopy reveals ulcerative proctitis or proctocolitis, with purulent exudate and mucosal bleeding. Histopathologic findings in the rectal mucosa include granulomas with giant cells, crypt abscesses, and extensive inflammation. These clinical, sigmoidoscopic, and histopathologic findings may closely resemble those of Crohn's disease of the rectum.

The most common presenting picture in heterosexual men and women is the *inguinal syndrome*, which is characterized by painful inguinal lymphadenopathy beginning 2–6 weeks after presumed exposure; in rare instances, the onset comes after a few months. The inguinal adenopathy is unilateral in two-thirds of cases, and palpable enlargement of the iliac and femoral nodes is often evident on the same side as the enlarged inguinal nodes. The nodes are initially discrete, but progressive periadenitis results in a matted mass of nodes that becomes fluctuant and suppurative. The overlying skin becomes fixed, inflamed, and thin, and multiple draining fistulas finally develop. Extensive enlargement of chains of inguinal nodes above and below the inguinal ligament (“the sign of the groove”) is not specific and, although not uncommon, is documented in only a minority of cases. Spontaneous healing usually takes place after several months; inguinal scars or granulomatous masses of various sizes persist for life. Massive pelvic lymphadenopathy may lead to exploratory laparotomy.

Constitutional symptoms are common during the stage of regional lymphadenopathy and, in cases of proctitis, may include fever, chills, headache, meningismus, anorexia, myalgias, and arthralgias. Other systemic complications are infrequent but include arthritis with sterile effusion, aseptic meningitis, meningoencephalitis, conjunctivitis, hepatitis, and erythema nodosum (Fig. 25e-40). Complications of untreated anorectal infection include perirectal abscess; anal fistulas; and rectovaginal, rectovesical, and ischiorectal fistulas. Secondary bacterial infection probably contributes to these complications. Rectal stricture is a late complication of anorectal infection and usually develops 2–6 cm from the anal orifice—i.e., at a site within reach on digital rectal examination. A small percentage of cases of LGV in men present as chronic progressive infiltrative, ulcerative, or fistular lesions of the penis, urethra, or scrotum. Associated lymphatic obstruction may produce elephantiasis. When urethral stricture occurs, it usually involves the posterior urethra and causes incontinence or difficulty with urination.

Diagnosis • DETECTION METHODS Historically, chlamydiae were cultivated in the yolk sac of embryonated eggs. The organisms can be grown more easily in tissue culture, but cell culture—once considered the diagnostic gold standard—has been replaced by nonculture assays (Table 213-1). In general, culture for chlamydiae in clinical specimens is now performed only in specialized laboratories. The first nonculture assays, such as DFA staining of clinical material and enzyme immunoassay (EIA), have been replaced by NAATs, which are molecular

TABLE 213-1 DIAGNOSTIC TESTS FOR SEXUALLY TRANSMITTED AND PERINATAL *CHLAMYDIA TRACHOMATIS* INFECTION

Infection	Suggestive Signs/Symptoms	Presumptive Diagnosis ^a	Confirmatory Test of Choice
Men			
NGU, PGU	Discharge, dysuria	Gram's stain with >4 neutrophils per oil-immersion field; no gonococci	Urine or urethral NAAT for <i>C. trachomatis</i>
Epididymitis	Unilateral intrascrotal swelling, pain, tenderness; fever; NGU	Gram's stain with >4 neutrophils per oil-immersion field; no gonococci; urinalysis with pyuria	Urine or urethral NAAT for <i>C. trachomatis</i>
Women			
Cervicitis	Mucopurulent cervical discharge, bleeding and edema of the zone of cervical ectopy	Cervical Gram's stain with ≥20 neutrophils per oil-immersion field in cervical mucus	Urine, cervical, or vaginal NAAT for <i>C. trachomatis</i>
Salpingitis	Lower abdominal pain, cervical motion tenderness, adnexal tenderness or masses	<i>C. trachomatis</i> always potentially present in salpingitis	Urine, cervical, or vaginal NAAT for <i>C. trachomatis</i>
Urethritis	Dysuria and frequency without hematuria	MPC; sterile pyuria; negative routine urine culture	Urine or urethral NAAT for <i>C. trachomatis</i>
Adults of Either Sex			
Proctitis	Rectal pain, discharge, tenesmus, bleeding; history of receptive anorectal intercourse	Negative gonococcal culture and Gram's stain; at least 1 neutrophil per oil-immersion field in rectal Gram's stain	Rectal NAAT for <i>C. trachomatis</i> or culture
Reactive arthritis	NGU, arthritis, conjunctivitis, typical skin lesions	Gram's stain with >4 neutrophils per oil-immersion field; lack of gonococci indicative of NGU	Urine or urethral NAAT for <i>C. trachomatis</i>
LGV	Regional adenopathy, primary lesion, proctitis, systemic symptoms	None	Culture of LGV strain from node or rectum, occasionally from urethra or cervix; NAAT for <i>C. trachomatis</i> from these sites; LGV CF titer, ≥1:64; micro-IF titer, ≥1:512
Neonates			
Conjunctivitis	Purulent conjunctival discharge 6–18 days after delivery	Negative culture and Gram's stain for gonococci, <i>Haemophilus</i> spp., pneumococci, staphylococci	Conjunctival NAAT for <i>C. trachomatis</i> ; FA-stained scraping of conjunctival material
Infant pneumonia	Afebrile; staccato cough, diffuse rales, bilateral hyperinflation, interstitial infiltrates	None	Chlamydial culture or NAAT of sputum, pharynx, eye, rectum; micro-IF antibody to <i>C. trachomatis</i> —fourfold change in IgG or IgM antibody titer

^aA presumptive diagnosis of chlamydial infection is often made in the syndromes listed when gonococci are not found. A positive test for *Neisseria gonorrhoeae* does not exclude the involvement of *C. trachomatis*, which often is present in patients with gonorrhea.

Abbreviations: CF, complement-fixing; FA, fluorescent antibody; LGV, lymphogranuloma venereum; micro-IF, microimmunofluorescence; MPC, mucopurulent cervicitis; NAAT, nucleic acid amplification test; NGU, nongonococcal urethritis; PGU, postgonococcal urethritis.

Source: Reprinted with permission from WE Stamm: Chlamydial infections, in *Harrison's Principles of Internal Medicine*, 17th ed, AS Fauci et al (eds). New York, McGraw-Hill, 2008, p 1075.

tests that amplify the nucleic acids in clinical specimens. NAATs are currently recommended by the CDC as the diagnostic assays of choice; four or five NAAT assays approved by the U.S. Food and Drug Administration (FDA) are commercially available, some as high-throughput robotic platforms. Point-of-care diagnostic assays (including NAATs), by which patients can be treated before leaving the clinic, are of increasing interest and are becoming available.

CHOICE OF SPECIMEN Cervical and urethral swabs have traditionally been used for the diagnosis of STDs in female and male patients, respectively. However, given the greatly increased sensitivity and specificity of NAATs, less invasive samples (e.g., urine for both sexes and vaginal swabs for women) can be used. For screening of asymptomatic women, the CDC now recommends that self-collected or clinician-collected vaginal swabs, which are slightly more sensitive than urine, be used. Urine screening tests are often used in outreach screening programs, however. For symptomatic women undergoing a pelvic examination, cervical swab samples are desirable because they have slightly higher chlamydial counts. For male patients, a urine specimen is the sample of choice, but self-collected penile-meatal swabs have been explored.

ALTERNATIVE SPECIMEN TYPES Ocular samples from babies and adults can be assessed by NAATs. However, since commercial NAATs for this purpose have not yet been approved by the FDA, laboratories must perform their own verification studies. Samples from rectal and pharyngeal sites have been used successfully to detect chlamydiae, but laboratories must verify test performance.

OTHER DIAGNOSTIC ISSUES Because NAATs detect nucleic acids instead of live organisms, they should be used with caution as test-of-cure assays. Residual nucleic acid from cells rendered noninfective by antibiotics may continue to yield a positive result in NAATs until as long as 3 weeks after therapy, when viable organisms have actually been eradicated. Therefore, clinicians should not use NAATs for test of cure until after 3 weeks. The CDC currently does not recommend a test of cure after treatment for infection with *C. trachomatis*. However, because incidence studies have demonstrated that previous chlamydial infection increases the probability of becoming reinfected, the CDC does recommend that previously infected individuals be rescreened 3 months after treatment.

SEROLOGY Serologic testing may be helpful in the diagnosis of LGV and neonatal pneumonia caused by *C. trachomatis*. The serologic test of choice is the microimmunofluorescence (MIF) test, in which high-titer purified EBs mixed with embryonated chicken yolk-sac material are affixed to a glass microscope slide to which dilutions of serum are applied. After incubation and washing, fluorescein-conjugated IgG or IgM antibody is applied. The test is read with an epifluorescence microscope, with the highest dilution of serum producing visible fluorescence designated as the titer. The MIF test is not widely available and is highly labor intensive. Although the complement fixation (CF) test also can be used, it employs only lipopolysaccharide (LPS) as the antigen and therefore identifies the pathogen only to the genus level. Single-point titers of >1:64 support a diagnosis of LGV, in which it is difficult to demonstrate rising antibody titers; i.e., paired serum samples are difficult to obtain since, by its very nature, the disease results in the patient's being seen by the physician after the acute stage. Any antibody titer of >1:16 is considered significant evidence of exposure to chlamydiae. However, serologic testing is never recommended for diagnosis of uncomplicated genital infections of the cervix, urethra, and lower genital tract or for *C. trachomatis* screening of asymptomatic individuals.

TREATMENT *C. TRACHOMATIS* GENITAL INFECTIONS

A 7-day course of tetracycline (500 mg four times daily), doxycycline (100 mg twice daily), erythromycin (500 mg four times daily), or a fluoroquinolone (ofloxacin, 300 mg twice daily; or levofloxacin, 500 mg/d) can be used for treatment of uncomplicated chlamydial infections. A single 1-g oral dose of azithromycin is as effective as a 7-day

course of doxycycline for the treatment of uncomplicated genital *C. trachomatis* infections in adults. Azithromycin causes fewer adverse gastrointestinal reactions than do older macrolides such as erythromycin. The single-dose regimen of azithromycin has great appeal for the treatment of patients with uncomplicated chlamydial infection (especially those without symptoms and those with a likelihood of poor compliance) and of the sexual partners of infected patients. These advantages must be weighed against the considerably greater cost of azithromycin. Whenever possible, the single 1-g dose should be given as directly observed therapy. Although not approved by the FDA for use in pregnancy, this regimen appears to be safe and effective for this purpose. However, amoxicillin (500 mg three times daily for 7 days) also can be given to pregnant women. The fluoroquinolones are contraindicated in pregnancy. A 2-week course of treatment is recommended for complicated chlamydial infections (e.g., PID, epididymitis) and at least a 3-week course of doxycycline (100 mg orally twice daily) or erythromycin base (500 mg orally four times daily) for LGV. Failure of treatment with a tetracycline in genital infections usually indicates poor compliance or reinfection rather than involvement of a drug-resistant strain. To date, clinically significant drug resistance has not been observed in *C. trachomatis*.

Treatment or testing for chlamydiae should be considered among *N. gonorrhoeae*-infected patients because of the frequency of co-infection. Systemic treatment with erythromycin has been recommended for ophthalmia neonatorum and for *C. trachomatis* pneumonia in infants. For the treatment of adult inclusion conjunctivitis, a single 1-g dose of azithromycin is as effective as standard 10-day treatment with doxycycline. Recommended treatment regimens for both bubonic and anogenital LGV include tetracycline, doxycycline, or erythromycin for 21 days.

SEX PARTNERS

The continued high prevalence of chlamydial infections in most parts of the United States is due primarily to the failure to diagnose—and therefore treat—patients with symptomatic or asymptomatic infection and their sex partners. Urethral or cervical infection with *C. trachomatis* has been well documented in a high proportion of the sex partners of patients with NGU, epididymitis, reactive arthritis, salpingitis, and endocervicitis. If possible, confirmatory laboratory tests for chlamydiae should be undertaken in these individuals, but even those without positive tests or evidence of clinical disease who have recently been exposed to proven or possible chlamydial infection (e.g., NGU) should be offered therapy. A novel approach is partner-delivered therapy, in which infected patients receive treatment and are also provided with single-dose azithromycin to give to their sex partner(s).

NEONATES AND INFANTS

In neonates with conjunctivitis or infants with pneumonia, erythromycin ethylsuccinate or estolate can be given orally at a dosage of 50 mg/kg per day, preferably in four divided doses, for 2 weeks. Careful attention must be given to compliance with therapy—a frequent problem. Relapses of eye infection are common after topical treatment with erythromycin or tetracycline ophthalmic ointment and may also follow oral erythromycin therapy. Thus follow-up cultures should be performed after treatment. Both parents should be examined for *C. trachomatis* infection and, if diagnostic testing is not readily available, should be treated with doxycycline or azithromycin.

Prevention Since many chlamydial infections are asymptomatic, effective control and prevention must involve periodic screening of individuals at risk. Selective cost-effective screening criteria have been developed. Among women, young age (generally <25 years) is a critical risk factor for chlamydial infections in nearly all studies. Other risk factors include mucopurulent cervicitis; multiple, new, or symptomatic male sex partners; and lack of barrier contraceptive use. In some settings, screening based on young age may be as sensitive

1172 as criteria that incorporate behavioral and clinical measures. Another strategy is universal testing of all patients in high-prevalence clinic populations (e.g., STD clinics, juvenile detention facilities, and family planning clinics).

The effectiveness of selective screening in reducing the prevalence of chlamydial infection among women has been demonstrated in several studies. In the Pacific Northwest, where extensive screening began in family planning clinics in 1998 and in STD clinics in 1993, the prevalence declined from 10% in the 1980s to <5% in 2000. Similar trends have occurred in association with screening programs elsewhere. In addition, screening can effect a reduction in upper genital tract disease. In Seattle, women at a large health maintenance organization who were screened for chlamydial infection on a routine basis had a lower incidence of symptomatic PID than did women who received standard care and underwent more selective screening.

In settings with low to moderate prevalence, the prevalence at which selective screening becomes more cost-effective than universal screening must be defined. Most studies have concluded that universal screening is preferable in settings with a chlamydial prevalence of >3–7%. Depending on the criteria used, selective screening is likely to be more cost-effective when prevalence falls below 3%. Nearly all regions of the United States have now initiated screening programs, particularly in family planning and STD clinics. Along with single-dose therapy, the availability of highly sensitive and specific diagnostic NAATs using urine specimens and self-obtained vaginal swabs makes it feasible to mount an effective nationwide *Chlamydia* control program, with screening of high-risk individuals in traditional health-care settings and in novel outreach and community-based settings. The U.S. Preventive Services Task Force has given *C. trachomatis* screening a Grade A recommendation, which means that private insurance and Medicare will cover its cost under the Affordable Care Act.

TRACHOMA



Epidemiology Trachoma—a sequela of ocular disease in developing countries—continues to be a leading cause of preventable infectious blindness worldwide. The WHO estimates that ~6 million people have been blinded by trachoma and that ~1.3 million people in developing countries still suffer from preventable blindness due to trachoma; certainly hundreds of millions live in trachoma-endemic areas. Foci of trachoma persist in Australia, the South Pacific, and Latin America. Serovars A, B, Ba, and C are isolated from patients with clinical trachoma in areas of endemicity in developing countries in Africa, the Middle East, Asia, and South America.

The trachoma-hyperendemic areas of the world are in northern and sub-Saharan Africa, the Middle East, drier regions of the Indian subcontinent, and Southeast Asia. In hyperendemic areas, the prevalence of trachoma is essentially 100% by the second or third year of life. Active disease is most common among young children, who are the reservoir for trachoma. By adulthood, active infection is infrequent but sequelae result in blindness. In such areas, trachoma constitutes the major cause of blindness.

Trachoma is transmitted through contact with discharges from the eyes of infected patients. Transmission is most common under poor hygienic conditions and most often takes place between family members or between families with shared facilities. Flies can also transfer the mucopurulent ocular discharges, carrying the organisms on their legs from one person to another. The International Trachoma Initiative founded by the WHO in 1998 aims to eliminate blinding trachoma globally by 2020.

Clinical Manifestations Both endemic trachoma and adult inclusion conjunctivitis present initially as conjunctivitis characterized by small lymphoid follicles in the conjunctiva. In regions with hyperendemic classic blinding trachoma, the disease usually starts insidiously before the age of 2 years. Reinfection is common and probably contributes to the pathogenesis of trachoma. Studies using polymerase chain reaction (PCR) or other NAATs indicate that chlamydial DNA is often present in the ocular secretions of patients with trachoma, even in the absence

of positive cultures. Thus persistent infection may be more common than was previously thought.

The cornea becomes involved, with inflammatory leukocytic infiltrations and superficial vascularization (*pannus formation*). As the inflammation continues, conjunctival scarring eventually distorts the eyelids, causing them to turn inward so that the lashes constantly abrade the eyeball (*trichiasis and entropion*); eventually the corneal epithelium is abraded and may ulcerate, with subsequent corneal scarring and blindness. Destruction of the conjunctival goblet cells, lacrimal ducts, and lacrimal gland may produce a “dry-eye” syndrome, with resultant corneal opacity due to drying (*xerosis*) or secondary bacterial corneal ulcers.

Communities with blinding trachoma often experience seasonal epidemics of conjunctivitis due to *H. influenzae* that contribute to the intensity of the inflammatory process. In such areas, the active infectious process usually resolves spontaneously in affected persons at 10–15 years of age, but conjunctival scars continue to shrink, producing trichiasis and entropion with subsequent corneal scarring in adults. In areas with milder and less prevalent disease, the process may be much slower, with active disease continuing into adulthood; blindness is rare in these cases.

Eye infection with oculogenital *C. trachomatis* strains in sexually active young adults presents as an acute onset of unilateral follicular conjunctivitis and preauricular lymphadenopathy similar to that seen in acute conjunctivitis caused by adenovirus or HSV. If untreated, the disease may persist for 6 weeks to 2 years. It is frequently associated with corneal inflammation in the form of discrete opacities (“infiltrates”), punctate epithelial erosions, and minor degrees of superficial corneal vascularization. Very rarely, conjunctival scarring and eyelid distortion occur, particularly in patients treated for many months with topical glucocorticoids. Recurrent eye infections develop most often in patients whose sexual partners are not treated with antimicrobial agents.

Diagnosis

The clinical diagnosis of classic trachoma can be made if two of the following signs are present: (1) lymphoid follicles on the upper tarsal conjunctiva; (2) typical conjunctival scarring; (3) vascular pannus; or (4) limbal follicles or their sequelae, Herbert’s pits. The clinical diagnosis of endemic trachoma should be confirmed by laboratory tests in children with relatively marked degrees of inflammation. Intracytoplasmic chlamydial inclusions are found in 10–60% of Giemsa-stained conjunctival smears in such populations, but chlamydial NAATs are more sensitive and are often positive when smears or cultures are negative. Follicular conjunctivitis in European or American adults living in trachomatous regions is rarely due to trachoma.

TREATMENT TRACHOMA

Adult inclusion conjunctivitis responds well to treatment with the same regimens used in uncomplicated genital infections—namely, azithromycin (a 1-g single oral dose) or doxycycline (100 mg twice daily for 7 days). Simultaneous treatment of all sexual partners is necessary to prevent ocular reinfection and chlamydial genital disease. Topical antibiotic treatment is not required for patients who receive systemic antibiotics.

PSITTACOSIS

Psittacine birds and many other avian species act as natural reservoirs for *C. psittaci*-type organisms, which are common pathogens in domestic mammals and birds. The species *C. psittaci*, which now includes only avian strains, affects humans only as a zoonosis. (The other strains previously included in this species have been placed into different species that generally reflect the animals they infect: *C. abortus*, *C. muridarum*, *C. suis*, *C. felis*, and *C. caviae*.) Although all birds are susceptible, pet birds (parrots, parakeets, macaws, and cockatiels)

and poultry (turkeys and ducks) are most frequently involved in transmission of *C. psittaci* to humans. Exposure is greatest among poultry-processing workers and owners of pet birds. Infectious forms of the organisms are shed from both symptomatic and apparently healthy birds and may remain viable for several months. *C. psittaci* can be transmitted to humans by direct contact with infected birds or by inhalation of aerosols from avian nasal discharges and from infectious avian fecal or feather dust. Transmission from person to person has never been demonstrated.

The diagnosis is usually established serologically. Psittacosis in humans may present as acute primary atypical pneumonia (which can be fatal in up to 10% of untreated cases); as severe chronic pneumonia; or as a mild illness or asymptomatic infection in persons exposed to infected birds.

EPIDEMIOLOGY

Fewer than 50 confirmed cases of psittacosis are reported in the United States each year, although many more cases probably occur than are reported. Control of psittacosis depends on control of avian sources of infection. A pandemic of psittacosis was once stopped by banning shipment or importation of psittacine birds. Birds can receive prophylaxis in the form of a tetracycline-containing feed. Imported birds are currently quarantined for 30 days of treatment.

CLINICAL MANIFESTATIONS

Typical symptoms include fever, chills, muscular aches and pains, severe headache, hepato- and/or splenomegaly, and gastrointestinal symptoms. Cardiac complications may involve endocarditis and myocarditis. Fatal cases were common in the preantibiotic era. As a result of quarantine of imported birds and improved veterinary-hygienic measures, outbreaks and sporadic cases of psittacosis are now rare. Severe pneumonia requiring management in an intensive care unit may develop. Endocarditis, hepatitis, and neurologic complications may occur, and fatal cases have been reported. The incubation period is usually 5–19 days but can last as long as 28 days.

DIAGNOSIS

Previously, the most widely used serologic test for diagnosing chlamydial infections was the genus-specific CF test, in which assay of paired serum specimens often shows fourfold or greater increases in antibody titer. The CF test remains useful, but the gold standard of serologic tests is now the MIF test, which is not widely available (see section on diagnosis of *C. trachomatis* genital infection, above). Any antibody titer above 1:16 is considered significant evidence of exposure to chlamydiae, and a fourfold titer rise in paired sera in combination with a clinically compatible syndrome can be used to diagnose psittacosis. Some commercially available serologic tests based on measurement of antibodies to LPS can be useful when the clinical diagnosis is consistent with bird exposure; however, since these tests are reactive for all chlamydiae (i.e., all chlamydiae contain LPS), caution must be used in their interpretation.

TREATMENT PSITTACOSIS


The antibiotic of choice is tetracycline; the dosage for adults is 250 mg four times a day, continued for at least 3 weeks to avoid relapse. Severely ill patients may need cardiovascular and respiratory support. Erythromycin (500 mg four times a day by mouth) is an alternative therapy.

CHLAMYDIA PNEUMONIAE INFECTIONS

C. pneumoniae is a common cause of human respiratory diseases, such as pneumonia and bronchitis. This organism has been reported to account for as many as 10% of cases of community-acquired pneumonia, most of which are diagnosed by serology. Serologic studies

have linked *C. pneumoniae* to atherosclerosis; isolation and PCR detection in cardiovascular tissues have also been reported. These findings suggest an expanded range of diseases and syndromes for *C. pneumoniae*. The role of *C. pneumoniae* in the etiology of atherosclerosis has been discussed since 1988, when Finnish researchers presented serologic evidence of an association of this organism with coronary heart disease and acute myocardial infarction. Subsequently, the organism was identified in atherosclerotic lesions by culture, PCR, immunohistochemistry, and transmission electron microscopy; however, discrepant study results (including those of animal studies) and failure of large-scale treatment studies have raised doubts as to the etiologic role of *C. pneumoniae* in atherosclerosis. Large-scale case-cohort studies have demonstrated some association of *C. pneumoniae* with lung cancer, as evaluated by serology.

EPIDEMIOLOGY

 Primary infection occurs mainly in school-aged children and reinfection in adults. Seroprevalence rates of 40–70% show that *C. pneumoniae* is widespread in both industrialized and developing countries. Seropositivity usually is first detected at school age, and rates generally increase by ~10% per decade. About 50% of individuals have detectable antibody at 30 years of age, and most have detectable antibody by the eighth decade of life. Although serologic evidence suggests that *C. pneumoniae* may be associated with up to 10% of cases of community-acquired pneumonia, most of this evidence is based not on paired serum samples but rather on a single high IgG titer. Some doubt exists about the true prevalence and etiologic role of *C. pneumoniae* in atypical pneumonia, especially since reports of cross-reactivity have raised questions about the specificity of serology when only a single serum sample is used for diagnosis.

PATHOGENESIS

Little is known about the pathogenesis of *C. pneumoniae* infection. It begins in the upper respiratory tract and, in many persons, persists as a prolonged asymptomatic condition of the upper respiratory mucosal surfaces. However, evidence of replication within vascular endothelium and synovial membranes of joints shows that, in at least some individuals, the organism is transported to distant sites, perhaps within macrophages. A *C. pneumoniae* outer-membrane protein may induce host immune responses whose cross-reactivity with human proteins results in an autoimmune reaction.

As mentioned above, epidemiologic studies have demonstrated an association between serologic evidence of *C. pneumoniae* infection and atherosclerotic disease of the coronary and other arteries. In addition, *C. pneumoniae* has been identified in atherosclerotic plaques by electron microscopy, DNA hybridization, and immunocytochemistry. The organism has been recovered in culture from atheromatous plaque—a result indicating the presence of viable replicating bacteria in vessels. Evidence from animal models supports the hypothesis that *C. pneumoniae* infection of the upper respiratory tract is followed by recovery of the organism from atheromatous lesions in the aorta and that the infection accelerates the process of atherosclerosis, especially in hypercholesterolemic animals. Antimicrobial treatment of the infected animals reverses the increased risk of atherosclerosis. In humans, two small trials in patients with unstable angina or recent myocardial infarction suggested that antibiotics reduce the likelihood of subsequent untoward cardiac events. However, larger-scale trials have not documented an effect of various antichlamydial regimens on the risk of these events.

CLINICAL MANIFESTATIONS

C. pneumoniae was first reported as the etiologic agent of mild atypical pneumonia in military recruits and college students. The clinical spectrum of *C. pneumoniae* infection includes acute pharyngitis, sinusitis, bronchitis, and pneumonitis, primarily in young adults. The clinical manifestations of primary infection appear to be more severe

1174 and prolonged than those of reinfection. The pneumonitis of *C. pneumoniae* pneumonia resembles that of *Mycoplasma pneumoniae* in that leukocytosis is frequently lacking and patients often have prominent antecedent upper respiratory tract symptoms, fever, nonproductive cough, mild to moderate illness, minimal findings on chest auscultation, and small segmental infiltrates on chest x-ray. In elderly patients, pneumonia due to *C. pneumoniae* can be especially severe and may necessitate hospitalization and respiratory support.

Chronic infection with *C. pneumoniae* has been reported among patients with chronic obstructive pulmonary disease and may also play a role in the natural history of asthma, including exacerbations. The clinical symptoms of respiratory infections caused by *C. pneumoniae* are nonspecific and do not differ from those caused by other agents of atypical pneumonia, such as *Mycoplasma pneumoniae*.

DIAGNOSIS

Serology, PCR amplification, and culture can be used to diagnose *C. pneumoniae* infection. Serology has been the traditional method of diagnosing infection by *C. pneumoniae*. The gold standard serologic test is the MIF test (see section on diagnosis of *C. trachomatis* genital infection, above). Any antibody titer above 1:16 is considered significant evidence of exposure to chlamydiae. According to a CDC-sponsored expert working group, the diagnosis of acute *C. pneumoniae* infection requires demonstration of a fourfold rise in titer in paired serum samples. There are no official recommendations for diagnosis of chronic infections, although many research studies have used high titers of IgA as an indicator. The older CF tests and EIAs for LPS are not

recommended, as they are not specific for *C. pneumoniae* but identify the chlamydiae only to the genus level. The organism is very difficult to grow in tissue culture but has been cultivated in HeLa cells, HEp-2 cells, and HL cells. Although NAATs are commercially available for *C. trachomatis*, only research-based PCR assays are available for *C. pneumoniae*.

TREATMENT *C. PNEUMONIAE* INFECTIONS

Although few controlled trials of treatment have been reported, *C. pneumoniae* is inhibited in vitro by erythromycin, tetracycline, azithromycin, clarithromycin, gatifloxacin, and gemifloxacin. Recommended therapy consists of 2 g/d of either tetracycline or erythromycin for 10–14 days. Other macrolides (e.g., azithromycin) and some fluoroquinolones (e.g., levofloxacin and gatifloxacin) also appear to be effective.

ACKNOWLEDGMENT

The authors wish to acknowledge the late Walter E. Stamm, MD, for his significant contributions to the field of Chlamydia research. Dr. Stamm wrote the chapters on chlamydiae for previous editions of Harrison's Principles of Internal Medicine, and we thank the editors for permission to reproduce Figs. 213-1 and 213-2 as well as Table 213-1 from his chapter in the 17th edition. Dr. Stamm died on December 14, 2009, and this chapter is dedicated to him.

SECTION 11

VIRAL DISEASES: GENERAL CONSIDERATIONS

214e Medical Virology

Fred Wang, Elliott Kieff

DEFINING A VIRUS

Viruses are obligate intracellular parasites. They consist of a DNA or RNA genome surrounded by protein. They may also have an outer-membrane lipoprotein envelope. Viruses can replicate only within cells because their nucleic acid does not encode many enzymes necessary for the metabolism of proteins, carbohydrates, or lipids or for the generation of high-energy phosphates. Typically, viral nucleic acids encode messenger RNA (mRNA) and proteins necessary for replicating, packaging, and releasing progeny virus from infected cells.

Viruses differ from virusoids, viroids, and prions. *Virusoids* are nucleic acids that depend on cells and helper viruses for packaging their nucleic acids into virus-like particles. *Viroids* are naked, cyclical, mostly double-strand small RNAs that appear to be restricted to plants, spread from cell to cell, and are replicated by cellular RNA polymerase II. Prions (Chap. 453e) are abnormal proteins that propagate and cause disease by altering the structure of a normal cell protein. Prions cause neurodegenerative diseases such as Creutzfeldt-Jakob disease, Gerstmann-Sträussler disease, kuru, and human or bovine spongiform encephalopathy (“mad cow disease”).

VIRUS STRUCTURE

Viral genomes may consist of single- or double-strand DNA, single- or double-strand RNA, single-strand or segmented antisense RNA, or double-strand segmented RNA. Viral nucleic acids may encode only a few genes or more than 100. Sense-strand viral RNA genomes can be translated directly into protein, whereas antisense RNAs must be copied into translatable RNA. Sense and antisense genomes are also referred to as *positive-strand* and *negative-strand genomes*, respectively. Viral nucleic acid is usually associated with virus-encoded nucleoprotein(s) in the virus core. Viral nucleic acids and nucleoproteins are almost always enclosed in a protein *capsid*. Because of the limited genetic complexity of viruses, their capsids are usually composed of multimers of identical *capsomeres* made up of one or a few proteins. Capsids have icosahedral or helical symmetry. Icosahedral capsid structures approximate spheres and have two-, three-, or fivefold axes of symmetry, whereas helical capsid structures have only a twofold axis of symmetry. The nucleic acid, nucleoprotein(s), and protein capsid together are called a *nucleocapsid*.

Many viruses are composed of a nucleic acid core and a capsid. For these viruses, the outer capsid surface mediates contact with uninfected cells' plasma membranes. Other viruses are more complex and have an outer phospholipid, cholesterol, glycoprotein, and glycolipid envelope that is derived from virus-modified infected cell membranes. Cell nuclear, endoplasmic reticulum, Golgi, or plasma membranes that become parts of the viral envelope have usually been modified during infection by the insertion of virus-encoded glycoproteins, which mediate contact of enveloped virus with uninfected cell surfaces. Matrix or tegument proteins may fill the space between the nucleocapsid and the outer envelope of the virus.

Enveloped viruses are usually sensitive to lipid solvents or detergents that can dissolve the envelope, whereas viruses with protein nucleocapsid exteriors may be somewhat detergent resistant. A schematic diagram of large and complex herpesviruses is shown in Fig. 214e-1. Structures of prototypical pathogenic human viruses are described in Table 214e-1. The relative sizes and structures of typical pathogenic human viruses are shown in Fig. 214e-2.

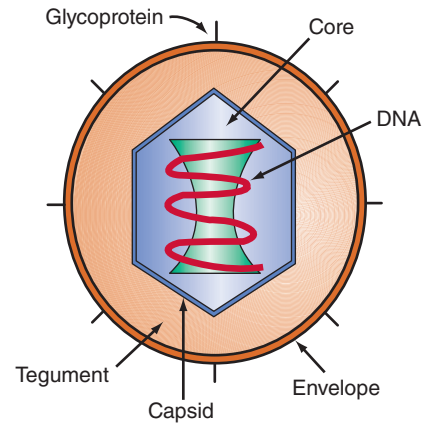


FIGURE 214e-1 Schematic diagram of an enveloped herpesvirus with an icosahedral nucleocapsid. The approximate respective dimensions of the nucleocapsid and the enveloped particles are 110 and 180 nm. The capsid is composed of 162 capsomeres: 150 with sixfold and 12 with fivefold axes of symmetry.

TAXONOMY OF PATHOGENIC HUMAN VIRUSES

As is apparent from Table 214e-1 and Fig. 214e-2, the classification of viruses into orders and families is based on nucleic acid composition, nucleocapsid size and symmetry, and presence or absence of an envelope. Viruses of a single family have similar structures and may be morphologically indistinguishable in electron micrographs. Subclassification into genera depends on similarity in epidemiology, biologic effects, and nucleic acid sequence.

Most viruses that infect humans have a common name related to their pathologic effects or the circumstances of their discovery. In addition, formal species names—consisting of the name of the host followed by the family or genus of the virus and a number—have been assigned by the International Committee on Taxonomy of Viruses. This dual terminology can cause confusion when viruses are referred to by either name—e.g., varicella-zoster virus (VZV) or human herpesvirus 3 (HHV-3).

VIRAL INFECTION IN VITRO

STAGES OF VIRAL INFECTION OF CELLS IN CULTURE

Viral Interactions with Cell Surfaces and Cell Entry To deliver its nucleic acid payload to the cell cytoplasm or nucleoplasm, a virus must overcome barriers posed by the cell's plasma and cytoplasmic membranes. Infection is frequently initiated by weak electrostatic or hydrophobic interactions with the cell surface. Subsequent stronger, more specific attachment to a cell plasma membrane protein, carbohydrate, glycolipid, heparan sulfate proteoglycan, or sialic acid enables stable binding to a specific cell surface “receptor” that mediates fusion with the cell plasma membrane (see Table 145e-1). Receptor binding is often augmented by a viral surface protein interaction with more than one cell surface protein or co-receptor. Receptors and co-receptors are important determinants of the species and cell type that a virus can infect. For example, the HIV envelope glycoprotein binds to the T cell surface protein CD4 and then engages a chemokine receptor that is the definitive co-receptor for the virus and mediates entry into the cell cytoplasm. The Epstein-Barr virus (EBV) glycoprotein gp350 binds to the B lymphocyte complement receptor CD21 and then uses a major histocompatibility complex (MHC) class II molecule as a co-receptor and an integrin for definitive entry.

TABLE 214e-1 VIRUS FAMILIES PATHOGENIC FOR HUMANS

Family	Representative Viruses	Type of RNA/DNA	Lipid Envelope		
RNA Viruses					
Picornaviridae	Poliovirus	(+) RNA	No		
	Coxsackievirus				
	Echovirus				
	Enterovirus				
	Rhinovirus				
Caliciviridae	Hepatitis A virus	(+) RNA	No		
	Norovirus				
Togaviridae	Hepatitis E virus	(+) RNA	Yes		
	Rubella virus				
Flaviviridae	Eastern equine encephalitis virus	(+) RNA	Yes		
	Western equine encephalitis virus				
	Yellow fever virus				
	Dengue virus				
	St. Louis encephalitis virus				
Coronaviridae	West Nile virus	(+) RNA	Yes		
	Hepatitis C virus				
Rhabdoviridae	Hepatitis G virus	(+) RNA	Yes		
	Coronaviruses ^a				
Filoviridae	Rabies virus	(–) RNA	Yes		
	Vesicular stomatitis virus				
Paramyxoviridae	Marburg virus	(–) RNA	Yes		
	Ebola virus				
	Parainfluenza virus				
	Respiratory syncytial virus				
	Newcastle disease virus				
Orthomyxoviridae	Mumps virus	(–) RNA, 8 segments	Yes		
	Rubeola (measles) virus				
Bunyaviridae	Influenza A, B, and C viruses	(–) RNA, 3 circular segments	Yes		
	Hantavirus				
	California encephalitis virus				
Arenaviridae	Sandfly fever virus	(–) RNA, 2 circular segments	Yes		
	Lymphocytic choriomeningitis virus				
	Lassa fever virus				
Reoviridae	South American hemorrhagic fever virus	ds RNA, 10–12 segments	No		
	Rotavirus				
Retroviridae	Reovirus	(+) RNA, 2 identical segments	Yes		
	Colorado tick fever virus				
	Human T lymphotropic virus types 1 and 2				
Hepadnaviridae	Human immunodeficiency virus types 1 and 2	ds DNA with ss portions	Yes		
	Hepatitis B virus				
	Parvoviridae			ss DNA	No
	Papillomaviridae				
	Polyomaviridae				
Adenoviridae	JC virus	ds DNA	No		
	BK virus				
Herpesviridae	Merkel cell polyoma virus	ds DNA	Yes		
	Human adenoviruses				
	Herpes simplex virus types 1 and 2 ^b				
	Varicella-zoster virus ^c				
	Epstein-Barr virus ^d				
	Cytomegalovirus ^e				
	Human herpesvirus 6				
	Human herpesvirus 7				
	Kaposi's sarcoma-associated herpesvirus ^f				
	Poxviridae			Variola (smallpox) virus	ds DNA
Orf virus					
Molluscum contagiosum virus					

^aIncluding the coronaviruses causing severe acute respiratory syndrome (SARS) and Middle Eastern respiratory syndrome (MERS). ^bAlso called human herpesvirus 1 (HHV-1) and HHV-2, respectively. ^cAlso called HHV-3. ^dAlso called HHV-4. ^eAlso called HHV-5. ^fAlso called HHV-8.

Abbreviations: ds, double-strand; ss, single-strand.

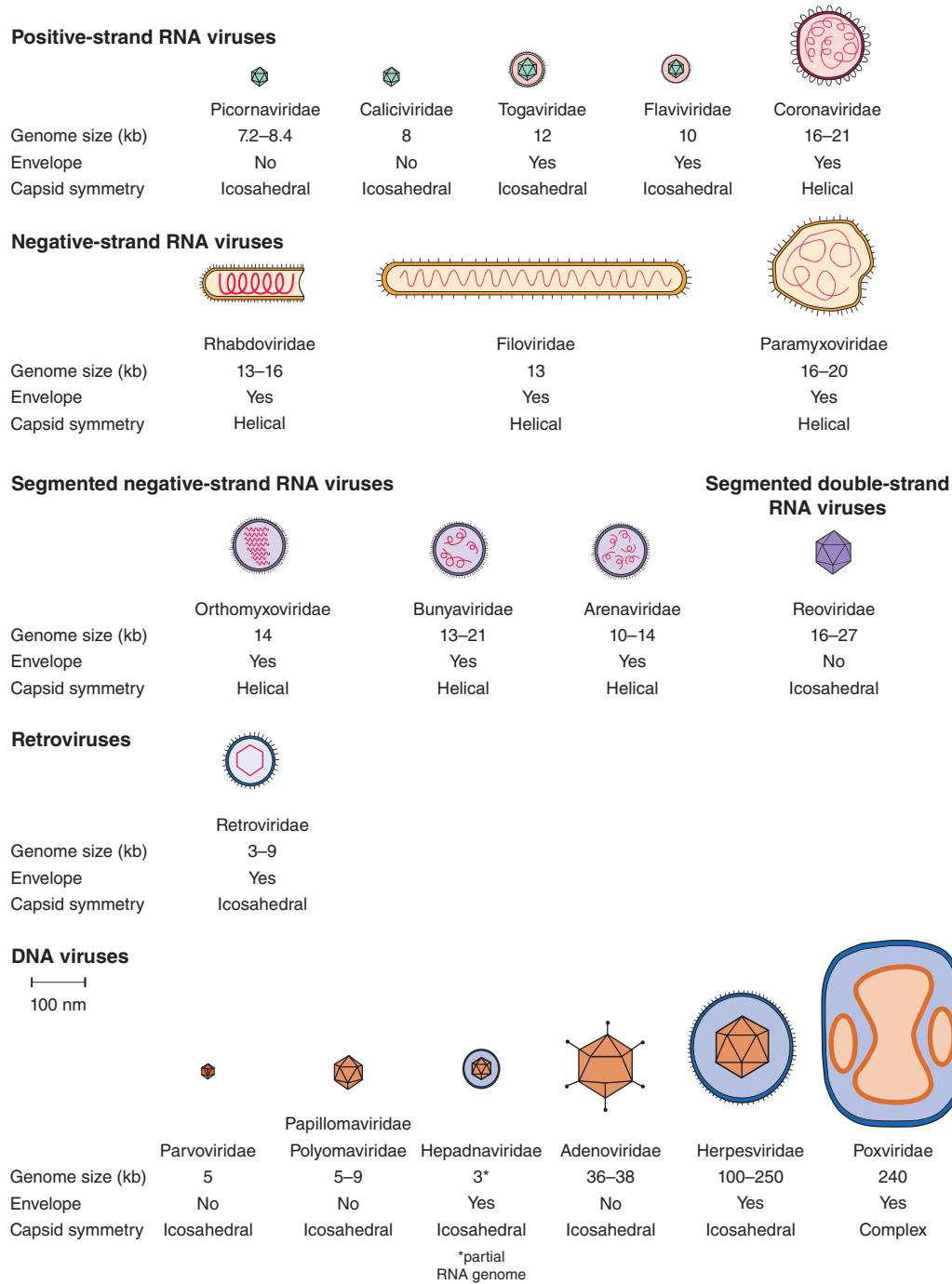


FIGURE 214e-2 Schematic diagrams of the major virus families including species that infect humans. The viruses are grouped by genome type and are drawn approximately to scale. Prototype viruses of each family that cause human disease are listed in Table 214e-1.

Viruses have evolved a wide range of strategies to enter cells. Influenza virus has an outer-membrane hemagglutinin glycoprotein that binds to sialic acid on respiratory tract cell plasma membranes. The hemagglutinin mediates adsorption to cell membranes, receptor aggregation, and endocytosis. As the endosome pH decreases in the cell cytoplasm, the influenza hemagglutinin conformation changes, enabling hydrophobic helices, which are initially at the base of the hemagglutinin, to extend, interacting and fusing with the endosome membrane and thereby releasing the viral genome into the cell cytoplasm. The influenza virus M2 membrane channel protein has a key role in lowering endosome pH and permitting virus and cell membrane fusion.

Nonenveloped viruses (e.g., human papillomaviruses [HPVs]) and some enveloped viruses have evolved to partially fuse with cell plasma

membrane receptors and be internalized into endosomes. The low pH in an endosome can then trigger virus membrane or capsid fusion with the endocytic membrane, releasing viral DNA into the cytoplasm to initiate infection.

Hydrophobic interactions required for fusion can be susceptible to chemical inhibition or blockade. The HIV envelope glycoprotein gp120 is associated with gp41 on the viral surface. HIV gp120 binding to CD4 and then to specific chemokine receptors results in conformational changes that allow gp41 to initiate cell membrane fusion. The anti-HIV drug enfuvirtide is a small peptide derived from the gp41 structure. Enfuvirtide binds to gp41 and prevents conformational changes required for fusion. In contrast, maraviroc prevents virus entry by binding to the CCR5 receptor, thereby blocking gp120 binding to CCR5 and preventing gp120 fusion with CCR5.

After uncoating and release of viral nucleoprotein into the cytoplasm, the viral genome is transported to sites of expression and replication. To produce infectious progeny, viruses must produce proteins necessary for replicating their nucleic acids as well as structural proteins necessary for coating their nucleic acids and for assembling nucleic acids and proteins into progeny virus. Different viruses use different strategies and gene repertoires to accomplish these goals. Most DNA viruses, except for poxviruses, replicate their nucleic acid and assemble into nucleocapsids in the cell nucleus. RNA viruses, except for influenza viruses, transcribe and replicate their RNA and assemble in the cytoplasm before envelopment at the cell plasma membrane. The replication strategies of DNA and RNA viruses and of positive- and negative-strand RNA viruses are presented and discussed separately below. Medically important viruses of each group are used for illustrative purposes.

POSITIVE-STRAND RNA VIRUSES RNA viruses of medical importance include positive-strand picornaviruses, flaviviruses, togaviruses, caliciviruses, and coronaviruses. Genome RNA from positive-strand RNA viruses is released into the cytoplasm without associated enzymes. Cell ribosomes recognize and associate with the viral genome's internal ribosome entry sequence and translate a virus-encoded polyprotein. Proteases within the polyprotein cleave out the viral RNA polymerase and other viral proteins necessary for replication. Antigenomic RNA is next transcribed from the genome RNA template. Positive-strand genomes and mRNAs are then transcribed from the antigenome RNA by the viral RNA polymerase and are translated into capsid proteins. Genomic RNA is encapsidated in the cytoplasm and released as the infected cell undergoes lysis.

NEGATIVE-STRAND RNA VIRUSES Medically important negative-strand RNA viruses include rhabdoviruses, filoviruses, paramyxoviruses, orthomyxoviruses, and bunyaviruses. The genomes of negative-strand viruses are frequently segmented. Negative-strand RNA viral genomes are released into the cytoplasm with an associated RNA polymerase and one or more polymerase accessory proteins. The viral RNA polymerase transcribes mRNAs as well as full-length antigenome RNA, which is the template for genome RNA replication. Viral mRNAs encode the viral RNA polymerase and accessory factors as well as viral structural proteins. Except for influenza virus, which transcribes its mRNAs and antigenome RNAs in the cell nucleus, negative-strand RNA viruses replicate entirely in the cytoplasm. All negative-strand RNA viruses, including influenza viruses, assemble in the cytoplasm.

DOUBLE-STRAND SEGMENTED RNA VIRUSES Double-strand RNA viruses are taxonomically grouped in the family Reoviridae. The medically important viruses in this group are rotaviruses and Colorado tick fever virus. Reovirus genomes have 10–12 RNA segments. Reovirus particles contain an RNA polymerase complex. These viruses replicate and assemble in the cell cytoplasm.

DNA VIRUSES Medically important DNA viruses include parvoviruses, which have small single-strand DNA genomes and cause transient arthritis, and polyomaviruses, including the smaller polyomaviruses such as JC virus, which causes progressive multifocal leukoencephalopathy in immunocompromised patients; BK virus; and Merkel cell polyomavirus. The larger HPVs cause warts as well as cervical, penile, and oral carcinomas. The next larger DNA viruses are adenoviruses, which mostly cause transient respiratory tract and ocular inflammatory disease. The herpesviruses include eight viruses that cause a wide range of inflammatory and malignant diseases in humans. EBV is an important cause of lymphomas and Hodgkin's disease in both immunocompromised and immunocompetent people and of nasopharyngeal carcinoma in southern Chinese and northern African populations. Cytomegalovirus (CMV) is an important cause of transplacental infections and neonatal neurologic impairment. Poxviruses, the largest DNA viruses and the largest viruses that infect humans (barely visible by light microscopy), cause smallpox, monkeypox, and molluscum contagiosum. Aside from those of poxviruses, other DNA virus genomes enter the cell nucleus and are transcribed by cellular RNA polymerase II.

After receptor binding and fusion with plasma membranes or endocytic vesicle membranes, herpesvirus nucleocapsids are released into the cytoplasm with tegument proteins and are transported along microtubules to a nuclear pore. Capsids then release DNA into the nucleus.

DNA virus transcription and mRNA processing depend on both viral and cellular proteins. For herpes simplex virus (HSV), a viral tegument protein enters the nucleus and activates immediate-early genes, the first genes expressed after infection. Transcription of immediate-early genes requires the viral tegument protein and cell transcription factors. HSV becomes nonreplicating, or latent, in neurons because essential cell transcription factors for expression of viral immediate-early genes are docked in the cytoplasm in neurons. Heat shock or other cell stresses can cause these cell factors to enter the nucleus, activate viral gene expression, and initiate replication. This information explains HSV-1 latency in neurons and activation of replicative infection.

For adenoviruses and herpesviruses, transcription of immediate-early genes results in expression of early proteins necessary for viral DNA replication. Viral DNA synthesis is required to turn on late-gene expression and production of viral structural components. The HPVs, polyomaviruses, and parvoviruses are not dependent on transactivators encoded from the viral genome for early-gene transcription. Instead, their early genes have upstream enhancing elements that bind cell transcription factors. The early genes encode proteins that are necessary for viral DNA synthesis and late-gene transcription. DNA virus late genes encode structural proteins necessary for viral assembly and for viral egress from the infected cell. Late-gene transcription is continuously dependent on DNA replication. Therefore, inhibitors of DNA replication also stop late-gene transcription.

Each DNA virus family uses unique mechanisms for replicating its DNA. Adenovirus and herpesvirus DNAs are linear in the virion. Adenovirus DNA remains linear in infected cells and replicates as a linear genome, using an initiator protein–DNA complex. In contrast, herpesvirus DNA circularizes in the infected cell, and genomes replicate into linear concatemers through a “rolling-circle” mechanism. Full-length DNA genomes are cleaved and packaged into virus. Herpesviruses encode a DNA polymerase and at least six other viral proteins necessary for viral DNA replication. Acyclovir and ganciclovir prevent viral DNA synthesis when they are phosphorylated and incorporated into DNA by the viral polymerase. Herpesviruses also encode enzymes that increase the deoxynucleotide triphosphate pools. HPV and polyomavirus DNAs are circular both within the virus and in infected cells. These genomes are reproduced by cellular DNA replication enzymes and remain circular through replication and packaging. HPV and polyomavirus early proteins are necessary for DNA replication in both latent and viral replicative phases. Early viral proteins stimulate cells to remain in cycle, facilitating viral DNA replication.

Parvoviruses have negative single-strand DNA genomes and are the smallest DNA viruses. Their genomes are half the size of HPV genomes and include only two genes. The replication of autonomous parvoviruses, such as B19, depends on cellular DNA replication and requires the virus-encoded Rep protein. Other parvoviruses, such as adeno-associated virus (AAV), are not autonomous and require helper viruses of the adenovirus or herpesvirus family for their replication. AAV is being used as a potentially safe human gene therapy vector because its replication protein causes integration at a single chromosome site. The small genome size limits the range of proteins that can be expressed from AAV vectors.

As stated above, poxviruses are the largest DNA viruses. They are unique among DNA viruses in replicating and assembling in the cytoplasm. To accomplish cytoplasmic replication, poxviruses encode transcription factors, an RNA polymerase II orthologue, enzymes for RNA capping, enzymes for RNA polyadenylation, and enzymes for viral DNA synthesis. Poxvirus DNA also has a unique structure. The double-strand linear DNA is covalently linked at the ends, making a covalently closed double-strand circular genome. Replication of the circular genomes is initiated by nicking in inverted repeats at the ends of the linear DNA. During DNA replication, the genome is cleaved

within the terminal inverted repeats, and the inverted repeats self-prime complementary-strand synthesis by the virus-encoded DNA polymerase. Like herpesviruses, poxviruses encode several enzymes that increase deoxynucleotide triphosphate precursor levels and thus facilitate viral DNA synthesis.

VIRUSES THAT USE BOTH RNA AND DNA GENOMES IN THEIR LIFE CYCLE Retroviruses, including HIV, are RNA viruses that use a DNA intermediate to replicate their genomes. In contrast, hepatitis B virus (HBV) is a DNA virus that uses an RNA intermediate to replicate its genome. Thus these viruses are not purely RNA or DNA viruses. Retroviruses are RNA viruses with two identical sense-strand genomes and associated reverse transcriptase and integrase enzymes. Retroviruses differ from all other viruses in that they reverse-transcribe themselves into partially duplicated double-strand DNA copies and then routinely integrate into the host genome as part of their persistence and replication strategies. Inhibitors of reverse transcriptase (e.g., zidovudine) or integrase (e.g., raltegravir) are now commonly used as antiviral treatments for HIV infection. Integration of remnants and even complete copies of simple retrovirus DNAs into the human genome raises the possibility of replication-competent simple human retroviruses. However, endogenous human retrovirus replication has not been documented or associated with any disease. Integrated, replication-competent retroviral DNAs are also present in many animal species, such as pigs. These porcine retroviruses are a potential cause for concern in xenotransplantation because retrovirus replication could cause disease in humans.

Cellular RNA polymerase II and transcription factors regulate transcription from the integrated provirus DNA genome. Some retroviruses also encode regulators of transcription and RNA processing, such as Tax and Rex in human T lymphotropic virus (HTLV) types 1 and 2. HIV-1 and HIV-2 have orthologous Tat and Rev genes as well as the additional accessory proteins Vpr, Vpu, and Vif, which are important for efficient infection and immune escape. Full-length proviral transcripts are made from a promoter in the viral terminal repeat and serve as both genome RNAs that are packaged in the nucleocapsids and differentially spliced mRNAs that encode for the virus Gag protein, polymerase/integrase protein, and envelope glycoprotein. The Gag protein includes a protease that cleaves it into several components, including a viral matrix protein that coats the viral RNA. Viral RNA polymerase/integrase, matrix protein, and cellular tRNAs are key components in the viral nucleocapsid. Protease inhibitors have been developed as effective agents against infections caused by HIV (e.g., saquinavir) or hepatitis C virus (HCV) (e.g., telaprevir).

HBV replication is unique in several respects. The HBV genome is a partially double-strand DNA genome that is repaired in infected cells to a fully double-strand circular DNA by the virion polymerase. Viral mRNAs are transcribed from the closed circular viral episome by the cellular RNA polymerase II and are translated to yield HBV proteins, including core protein, surface antigen, and polymerase. In addition, a full-genome-length mRNA is packaged into viral core particles in the cytoplasm of infected cells as an intermediate for viral DNA replication. This RNA associates with the viral polymerase, which also has reverse transcriptase activity and converts the full-length encapsidated RNA genome into partially double-strand DNA. Thus, nucleos(t)ide analogs that inhibit reverse transcription (e.g., tenofovir) are commonly used to treat HBV infection. HBV is believed to mature by budding through the cell's plasma membrane, which has been modified by the insertion of viral surface antigen protein.

Viral Assembly and Egress For most viruses, nucleic acid and structural protein synthesis is accompanied by the assembly of protein and nucleic acid complexes. The assembly and egress of mature infectious virus mark the end of the *eclipse phase* of infection, during which infectious virus cannot be recovered from the infected cell. Nucleic acids from RNA viruses and poxviruses assemble into nucleocapsids in the cytoplasm. For all DNA viruses except poxviruses, viral DNA assembles into nucleocapsids in the nucleus. In general, the capsid proteins of viruses with icosahedral nucleocapsids can self-assemble into densely packed and highly ordered capsid structures. Herpesviruses require an assembly protein as a scaffold for capsid assembly. Viral nucleic acid then

spools into the assembled capsid. For herpesviruses, a full unit of the viral DNA genome is packaged into the capsid, and a capsid-associated nuclease cleaves the viral DNA at both ends. In the case of viruses with helical nucleocapsids, the protein component appears to assemble around the nucleic acid, which contributes to capsid organization.

Viruses must egress from the infected cell and not bind back to their receptor(s) on the outer surface of the plasma membrane. Viruses can acquire envelopes from cytoplasmic membranes or by budding through the cell's plasma membrane. Excess viral membrane glycoproteins are synthesized to saturate cell receptors and facilitate separation of the virus from the infected cell. Some viruses encode membrane proteins with enzymatic activity for receptor destruction. Influenza virus, for example, encodes a glycoprotein with neuraminidase activity. Neuraminidase destroys sialic acid on the infected cell's plasma membrane so that newly released virus does not get stuck to the dying cell. Oseltamivir and zanamivir are neuraminidase inhibitors that are used to treat or provide prophylaxis for influenza virus infection. Herpesvirus nucleocapsids acquire an initial envelope by assembling in the nucleus and then budding through the nuclear membrane into the endoplasmic reticular space. The initially enveloped herpesvirus is then de-enveloped and released from the cell either by exocytosis or by re-envelopment at the plasma membrane. Nonenveloped viruses depend on the death and dissolution of the infected cell for their release.

FIDELITY OF VIRAL REPLICATION

Hundreds or thousands of progeny may be produced from a single virus-infected cell. Many particles partially assemble and never mature into virions. Many mature-appearing virions are imperfect and have only incomplete or nonfunctional genomes. Despite the inefficiency of assembly, a typical virus-infected cell releases 10–1000 infectious progeny. Some of these progeny may contain genomes that differ from those of the virus that infected the cell. Smaller, “defective” viral genomes have been noted with the replication of many RNA and DNA viruses. Virions with defective genomes can be produced in large numbers through packaging of incompletely synthesized nucleic acid. Adenovirus packaging is notoriously inefficient, and a high ratio of particle to infectious virus may limit the amount of recombinant adenovirus that can be administered for gene therapy since the immunogenicity of defective particles may contribute to adverse effects.

Changes in viral genomes can lead to mutant viruses of medical significance. In general, viral nucleic acid replication is more error-prone than cellular nucleic acid replication. RNA polymerases and reverse transcriptases are significantly more error-prone than DNA polymerases. Mutations can also be introduced into the HIV genome by APOBEC3G, a cellular protein that is packaged in the virion. APOBEC3G deaminates cytidine in the virion RNA to uridine. When reverse transcriptase subsequently uses the altered virion RNA as a template in the infected cell, a guanosine-to-adenosine mutation is introduced into the proviral DNA. Mutations resulting in less efficient viral growth, or fitness, may be detrimental to the virus. HIV-encoded Vif blocks APOBEC3G activity in the virion, inhibiting the debilitating effects of hypermutation on genetic integrity. Nevertheless, mutations resulting in evasion of the host immune response or resistance to antiviral drugs are preferentially selected in patients, with the consequent perpetuation of infection. Viral genomes can also be altered by recombination or reassortment between two related viruses in a single infected cell. Although this occurrence is unusual under most circumstances of natural infection, the genome changes can be substantial and can significantly alter virulence or epidemiology. Reassortment of the avian or mammalian influenza A hemagglutinin gene into a human influenza background can result in the emergence of new epidemic or pandemic influenza A strains.

VIRAL GENES NOT REQUIRED FOR VIRAL REPLICATION

Viruses frequently have genes encoding proteins that are not directly involved in replication or packaging of the viral nucleic acid, in virion assembly, or in regulation of the transcription of viral genes involved in those processes. Most of these proteins fall into five classes: (1) proteins that directly or indirectly alter cell growth; (2) proteins that inhibit cellular RNA or protein synthesis so that viral mRNA

can be efficiently transcribed or translated; (3) proteins that promote cell survival or inhibit apoptosis so that progeny virus can mature and escape from the infected cell; (4) proteins that inhibit the host interferon response; and (5) proteins that downregulate host inflammatory or immune responses so that viral infection can proceed in an infected person to the extent consistent with the survival of the virus and its efficient transmission to a new host. More complex viruses of the poxvirus or herpesvirus family encode many proteins that serve these functions. Some of these viral proteins have motifs similar to those of cellular proteins, while others are quite novel. Virology has increasingly focused on these more sophisticated strategies evolved by viruses to permit the establishment of long-term infection in humans and other animals. These strategies often provide unique insights into the control of cell growth, cell survival, macromolecular synthesis, proteolytic processing, immune or inflammatory suppression, immune resistance, cytokine mimicry, or cytokine blockade.

MicroRNAs (miRNAs) are small noncoding RNAs that can regulate gene expression at the posttranscriptional level by targeting—and usually silencing—mRNAs. miRNAs were initially discovered in plants and plant viruses, where they alter expression of cell defenses. Herpesviruses are especially rich in miRNAs; for example, at least 23 miRNAs have been identified in EBV and 11 in CMV. Adenovirus and polyomavirus miRNAs have also been described. Increasing data indicate that animal viruses encode miRNAs to alter the growth and survival of host cells and the innate and acquired immune responses.

HOST RANGE

The concept of host range was originally based on the cell types in which a virus replicates in tissue culture. For the most part, the host range is limited by specific cell-surface proteins required for viral adsorption or penetration—i.e., to the cell types that express receptors or co-receptors for a specific virus. Another common basis for host-range limitation is the degree of transcriptional activity from viral promoters in different cell types. Most DNA viruses depend not only on cellular RNA polymerase II and the basal components of the cellular transcription complex but also on activated components and transcriptional accessory factors, both of which differ among differentiated tissues, among cells at various phases of the cell cycle, and between resting and cycling cells.

The importance of host range factors is illustrated by the effects of specific host determinants that limit the replication of influenza virus with avian or porcine hemagglutinins in humans. These viral proteins have adapted to bind avian or porcine sialic acids, and spread of avian or porcine influenza viruses in human populations is limited by their ability to infect human cells.

VIRAL CYTOPATHIC EFFECTS AND INHIBITORS OF APOPTOSIS

The replication of almost all viruses has adverse effects on the infected cell, inhibiting cellular synthesis of DNA, RNA, or proteins through efficient competition for key substrates and enzymatic processes. These general inhibitory effects enable viruses to nonspecifically limit components of innate host resistance, such as interferon (IFN) production. Viruses can specifically inhibit host protein synthesis by attacking a component of the translational initiation complex—frequently, a component that is not required for efficient translation of viral RNAs. Poliovirus protease 2A, for example, cleaves a cellular component of the complex that ordinarily facilitates translation of cellular mRNAs by interacting with their cap structure. Poliovirus RNA is efficiently translated without a cap because it has an internal ribosome entry sequence. Influenza virus inhibits the processing of mRNA by snatching cap structures from nascent cellular RNAs and using them as primers in the synthesis of viral mRNA. HSV has a virion tegument protein that inhibits cellular mRNA translation.

Apoptosis is the expected consequence of virus-induced inhibition of cellular macromolecular synthesis and viral nucleic acid replication. Although the induction of apoptosis may be important for the release of some viruses (particularly nonenveloped viruses), many viruses have acquired genes or parts of genes that enable them to

forestall infected-cell death. This delay increases the yield from viral replication. Adenoviruses and herpesviruses encode analogues of the cellular Bcl2 protein, which block mitochondrial enhancement of proapoptotic stimuli. Poxviruses and some herpesviruses also encode caspase inhibitors. Many viruses, including HPVs and adenoviruses, encode proteins that inhibit p53 or its downstream proapoptotic effects.

VIRAL INFECTION IN VIVO

TRANSMISSION

The capsid and envelope of a virus protect the genome and enable efficient transmission of the virus from cell to cell and to new prospective hosts. Most common viral infections are spread by direct contact, by ingestion of contaminated water or food, or by inhalation of aerosolized particles. In all these situations, infection begins on an epithelial or mucosal surface and spreads along the mucosa and into deeper tissues. Infection may spread to cells that can enter blood vessels, lymphatics, or neural circuits. HBV, HCV, HTLV, and HIV are dependent on transmission by parenteral inoculation. Some viruses are transmitted only between humans. The dependence of smallpox virus and poliovirus infections on interhuman transmission makes it feasible to eliminate these viruses from human circulation by mass vaccination. Herpesviruses also survive by interhuman transmission but may be more difficult to eliminate because they establish persistent latent infection in humans and continuously reactivate to infect new and naïve generations.

Animals are also important reservoirs and vectors for transmission of viruses causing human disease. Insect vectors can mediate parenteral transfer of viruses that reach high titers in animal or human hosts. Arboviruses are parenterally transmitted from mammalian species to humans by mosquito vectors. Herpes B, monkeypox, rabies, and viral hemorrhagic fevers are other examples of zoonotic infections caused by direct contact with animals, animal tissues, or arthropod vectors.

PRIMARY INFECTION

Initial viral infections usually last for several days or weeks. During this period, the concentration of virus at sites of infection rises and then falls, usually to unmeasurable levels. The rise and fall of viral replication at a given site depend on local innate immune responses and the access of systemic antibody and cell immune effectors to the virus. Typically, primary infections with enteroviruses, mumps virus, measles virus, rubella virus, rotavirus, influenza virus, AAV, adenovirus, HSV, and VZV are cleared from almost all sites within 3–4 weeks. Some viruses are especially proficient in altering or evading innate and acquired immune responses. Primary infection with AAV, EBV, or CMV can last for several months. Characteristically, primary infections due to HBV, HCV, hepatitis D virus (HDV), HIV, HPV, and molluscum contagiosum virus (MCV) extend beyond several weeks. For some of these viruses (e.g., HPV, HBV, HCV, HDV, and MCV), the manifestations of primary infection are almost indistinguishable from the persistent phase.

Disease manifestations usually arise as a consequence of viral replication, infected-cell injury or death, and local inflammatory and innate immune responses. Disease severity may not necessarily correlate with the level of viral replication alone. For example, the clinical manifestations of intense primary infection with poliovirus, enterovirus, rabies virus, measles virus, mumps virus, or HSV at mucosal surfaces may be inapparent or relatively mild, whereas limited replication in neural cells can have dramatic consequences. Similarly, rubella virus or CMV infections in utero or neonatal HSV infections may have much more devastating effects than infections in adults.

Primary infections are cleared by nonspecific innate and specific adaptive immune responses. Thereafter, an immunocompetent host is usually immune to the disease manifestations of reinfection by the same virus. Immunity frequently does not prevent transient surface colonization on reexposure, persistent colonization, or even limited deeper infection.

PERSISTENT AND LATENT INFECTIONS

Relatively few viruses cause persistent or latent infections. HBV, HCV, rabies virus, measles virus, HIV, HTLV, HPV, HHVs, and MCV are notable exceptions. The mechanisms for persistent infection vary. HCV RNA polymerase and HIV reverse transcriptase are error-prone and generate variant genomes. Genome variation can be sufficient to permit evasion of host immune responses, thereby allowing persistent infection. HIV is also directly immunosuppressive, depleting CD4+ T lymphocytes and compromising CD8+ cytotoxic T cell immune responsiveness. Moreover, HIV encodes the Nef protein, which downmodulates MHC class I expression, rendering HIV-infected cells partially resistant to immune CD8+ T cell lysis.

DNA viruses have low mutation rates. Their persistence in human populations usually depends on their ability to establish latent infection in some cells, to reactivate from latency, and then to replicate at epithelial surfaces. *Latency* is defined as a state of infection in which virus is not replicating, viral genes associated with lytic infection are not expressed, and infectious virus is not made. The complete viral genome is present and may be replicated by cellular DNA polymerase in conjunction with replication of the cell's genome. HPVs establish latent infection in basal epithelial cells. The latently infected basal cell replicates, along with the HPV episome, by using cellular DNA polymerase. Some of the progeny cells provide new latently infected basal cells, whereas others go on to squamous differentiation. Infected cells that differentiate to squamous cells become permissive for lytic viral infection. Herpesviruses establish latent infection in nonreplicating neural cells (HSV and VZV) or in replicating cells of hematopoietic lineages (EBV, CMV, HHV-6, HHV-7, and Kaposi's sarcoma-associated herpesvirus [KSHV, also known as HHV-8]). In their latent stage, HPV and herpesvirus genomes are largely hidden from the normal immune response. Reactivated HPV and herpesvirus infections escape immediate and effective immune responses in highly immune hosts by inhibiting host innate immune and inflammatory responses. In addition, HPV, HSV, and VZV are somewhat protected because they replicate in the middle and upper layers of the squamous epithelium—sites not routinely visited by cells that mediate or amplify immune and inflammatory responses. HSV and CMV are also known to encode proteins that downregulate MHC class I expression and antigenic peptide presentation, enabling infected cells to escape recognition by and cytotoxic effects of CD8+ T lymphocytes.

Like other poxviruses, MCV cannot establish latent infection. This virus causes persistent infection in hypertrophic skin lesions that last for months or years. MCV encodes a chemokine homologue that probably blocks inflammatory responses, an MHC class I analogue that blocks cytotoxic T lymphocyte attack, and inhibitors of cell death that prolong infected-cell viability.

PERSISTENT VIRAL INFECTIONS AND CANCER

Persistent viral infection is estimated to be the root cause of as many as 20% of human malignancies. Cancer is an accidental and highly unusual or long-term effect of oncogenic human viral infection. With most “oncogenic viruses,” infection is a critical and ultimately determinative early step in carcinogenesis. Latent HPV infection can block cell death and cause cervical cells to proliferate. A virus-infected cell with an integrated HPV genome overexpressing E6 and E7 undergoes subsequent cellular genetic changes that enhance autonomous malignant cell growth.

Most hepatocellular carcinoma is believed to be caused by chronic inflammatory, immune, and regenerative responses to HBV or HCV infection. Epidemiologic data firmly link HBV and HCV infections to hepatocellular carcinoma. These infections elicit repetitive cycles of virus-induced liver injury followed by tissue repair and regeneration. Over decades, chronic viral infection, repetitive tissue regeneration, and acquired chromosomal changes can result in proliferative nodules. Further chromosomal mutations can lead to the degeneration of cells in a proliferating nodule into hepatocellular carcinoma. In rare instances, HBV DNA integrates into cellular DNA, promoting overexpression of a cell gene that can also contribute to oncogenesis.

Most cervical carcinoma is caused by persistent infection with “high-risk” HPV type 16 or 18. In contrast to HBV and HCV infections, which

stimulate cell growth as a consequence of virus-induced cell death, HPV type 16 or 18 proteins E6 and E7 destroy p53 and pRB, respectively. Elimination of these key tumor-suppressive cell proteins increases cell growth, cell survival, and cell genome instability. However, like HBV and HCV infections, HPV infection alone is not sufficient for carcinogenesis. Cervical carcinoma is inevitably associated with persistent HPV infection and integration of the HPV genome into chromosomal DNA. Integrations that result in overexpression of E6 and E7 from HPV type 16 or 18 cause more profound changes in cell growth and survival and permit subsequent chromosomal changes that result in cervical carcinoma.

EBV is the most unusual oncogenic virus in that normal B cell infection results in latency with expression of viral proteins that can cause endless B lymphocyte growth. In almost all humans, strong CD4+ and CD8+ T cell immune responses to the antigenic EBV latent-infection nuclear proteins prevent uncontrolled B cell lymphoproliferation. However, when humans are severely immunosuppressed by transplantation-associated medication, HIV infection, or genetic immune deficiencies, EBV-induced B cell malignancies can emerge.

EBV infection also has a role in the long-term development of B lymphocyte and epithelial cell malignancies. Persistent EBV infection with expression of an EBV latency-associated integral membrane protein (LMP1) in latently infected epithelial cells appears to be a critical early step in the evolution of anaplastic nasopharyngeal carcinoma, a common malignancy in populations in southern China and northern Africa. Genomic instability and chromosomal abnormalities also contribute to the development of EBV-associated nasopharyngeal carcinoma. EBV is an important cause of Hodgkin's lymphoma. High-level expression of LMP1 or LMP2 in Reed-Sternberg cells is a hallmark in up to 50% of Hodgkin's lymphoma cases. LMP1-induced nuclear factor- κ B (NF- κ B) activity may prolong the survival of defective B cells that are normally eliminated by apoptosis, thereby allowing other genetic changes leading to the development of malignant Reed-Sternberg cells.

The HTLV-1 Tax and Rex proteins are critical to the initiation of cutaneous adult T cell lymphoma/leukemias that occur long after primary HTLV-1 infection. Tax-induced NF- κ B activation may contribute to cytokine production, infected-cell survival, and eventual outgrowth of malignant cells.

Molecular data confirm the presence of KSHV DNA in all Kaposi's tumors, including those associated with HIV infection, transplantation, and familial transmission. KSHV infection is also etiologically implicated in pleural-effusion lymphomas and multicentric Castlemans disease, which are more common among HIV-infected than among HIV-uninfected people. KSHV also has a virus-encoded cyclin, an IFN regulatory factor, and a latency-associated nuclear antigen that are implicated in increased-cell proliferation and survival.

Evidence supporting a causal role for viral infection in all of these malignancies includes (1) epidemiologic data, (2) the presence of viral DNA in all tumor cells, (3) the ability of the viruses to transform human cells in culture, (4) the results of in vitro cell culture-based assays that reveal transforming effects of specific viral genes on cell growth or survival, (5) pathologic data indicating the expression of transforming viral genes in premalignant or malignant cells in vivo, (6) the demonstration in animal models that these viral genes can cause malignant cell growth, and (7) the ability of virus-specific vaccines to reduce the incidence of virus-associated malignancy.

Virus-related malignancies provide an opportunity to expand our understanding of the biologic mechanisms important in the development of cancer. They also offer unique opportunities to develop diagnostics, vaccines, or therapeutics that could prevent or specifically treat cancers associated with viral infection. Widespread immunization against hepatitis B has resulted in a decreased prevalence of HBV-associated hepatitis and will probably prevent most HBV-related liver cancers. Current HPV vaccines can reduce rates of colonization with high-risk HPV strains and thereby decrease the risk of cervical cancer. The successful use of in vitro-expanded EBV-specific T cell populations to treat or prevent EBV-associated posttransplantation lymphoproliferative disease demonstrates the potential of immunoprevention or immunotherapy against virus-associated cancers.

Resistance to viral infections is initially provided by factors that are not virus-specific. Physical protection is afforded by the cornified layers of the skin and by mucous secretions that continuously sweep over mucosal surfaces. Once the first cell is infected, IFNs are induced and confer resistance to RNA virus replication. Viral infection may also trigger the release of other cytokines from infected cells. These cytokines may be chemotactic to inflammatory and immune cells. Viral protein epitopes expressed on the cell surface in the context of MHC class I and II proteins can stimulate the expansion of T cell populations with receptors that can recognize virus-encoded peptides presented on the cell surface by MHC class I proteins. Cytokines and antigens released by virus-induced cell death further attract inflammatory cells, dendritic cells, granulocytes, natural killer (NK) cells, and B lymphocytes to sites of infection and to draining lymph nodes. IFNs and NK cells are particularly important in containing viral infection for the first several days. Granulocytes and macrophages are also important in the phagocytosis and degradation of viruses, especially after an initial antibody response.

By 7–10 days after infection, virus-specific antibody responses, virus-specific human leukocyte antigen (HLA) class II-restricted CD4+ helper T lymphocyte responses, and virus-specific HLA class I-restricted CD8+ cytotoxic T lymphocyte responses develop. These responses, whose magnitude typically increases over the second and third weeks of infection, are important for rapid recovery. Also between the second and third weeks, the antibody type usually changes from IgM to IgG; IgG or IgA antibody can then be detected at infected mucosal surfaces. Antibody may directly neutralize virus by binding to its surface and preventing cell attachment or penetration. Complement can significantly enhance antibody-mediated virus neutralization. Antibody and complement can also lyse virus-infected cells that express viral membrane proteins on the cell surface. Cells infected with a replicating enveloped virus usually express the virus-envelope glycoproteins on the cell plasma membrane. Specific antibodies can bind to the glycoproteins, fix complement, and lyse the infected cell.

Antibody and CD4+/CD8+ T lymphocyte responses to viral infection can remain at high levels for several months after primary infection but usually wane over time. Low-level persistence of antibody-producing B lymphocytes and CD4+ or CD8+ T lymphocyte responses as memory cells can provide a rapid response to a second infection or an early barrier to reinfection with the same virus. Redevelopment of T cell immunity may take longer than secondary antibody responses, particularly when many years have elapsed between primary infection and reexposure. However, persistent infections or frequent reactivations from latency can result in sustained high-level T cell responses. EBV and CMV typically induce high-level CD4+ and CD8+ T cell responses that are maintained for decades after primary infection.

Some viruses have genes that alter innate and acquired host defenses. Adenoviruses encode small RNAs that inhibit IFN-induced, protein kinase R (PKR)-mediated shutoff of infected-cell protein synthesis. Adenovirus E1A can also directly inhibit IFN-mediated changes in cell gene transcription. Moreover, adenovirus E3 proteins prevent tumor necrosis factor (TNF)-induced cytolysis and block HLA class I synthesis by the infected cell. HSV ICP47 and CMV US11 also block class I antigen presentation. EBV encodes an interleukin (IL) 10 homologue that inhibits NK and T cell responses. Vaccinia virus encodes a soluble receptor for IFN- α and binding proteins for IFN- γ , IL-1, IL-18, and TNF, which inhibit host innate and adaptive immune responses. Vaccinia virus also encodes a caspase inhibitor that inhibits the ability of CD8+ cytotoxic T cells to kill virus-infected cells. Some poxviruses and herpesviruses encode chemokine-binding proteins that inhibit cell inflammatory responses. The adoption of these strategies by viruses highlights the importance of the corresponding host resistance factors in containing viral infection and the importance of redundancy in host resistance.

The host inflammatory and immune responses to viral infection do not come without a price. These responses contribute to the symptoms, signs, and other pathophysiologic manifestations of viral infection. Inflammation at sites of viral infection can subvert an

effective immune response and induce tissue death and dysfunction. Moreover, immune responses to viral infection could, in principle, result in immune attack upon cross-reactive epitopes on normal cells, with consequent autoimmunity.

INTERFERONS

All human cells can synthesize IFN- α or IFN- β in response to viral infection. These IFN responses are usually induced by the presence of double-strand viral RNA, which can be made by both RNA and DNA viruses and sensed by double-strand RNA binding proteins (e.g., PKR and RIG-I) in the cell cytoplasm. IFN- γ is not closely related to IFN- α or IFN- β and is produced mainly by NK cells and by immune T lymphocytes responding to IL-12. IFN- α and - β bind to the IFN- α receptor, whereas IFN- γ binds to a different but related receptor. Both receptors signal through receptor-associated JAK kinases and other cytoplasmic proteins, including “STAT” proteins, which are tyrosine-phosphorylated by JAK kinases, translocate to the nucleus, and activate promoters for specific cell genes. Three types of antiviral effects are induced by IFN at the transcriptional level. The first effect is attributable to the induction of 2'-5' oligo(A) synthetases, which require double-strand RNA for their activation. Activated synthetase polymerizes oligo(A) and thereby activates RNase L, which in turn degrades single-strand RNA. A second effect results from the induction of PKR, a serine and threonine kinase that is also activated by double-strand RNA. PKR phosphorylates and negatively regulates the translational initiation factor eIF2 α , shutting down protein synthesis in the infected cell. A third effect is initiated through the induction of Mx proteins, a family of GTPases that is particularly important in inhibiting the replication of influenza virus and vesicular stomatitis virus. These IFN effects are mostly directed against the infected cell, causing virus and cell dysfunction and thereby limiting viral replication.

DIAGNOSTIC VIROLOGY

A wide variety of methods are used to diagnose viral infection. Serology and virus isolation in tissue culture remain important standards. Acute- and convalescent-phase sera with rising titers of antibody to virus-specific antigens and a shift from IgM to IgG antibodies are generally accepted as diagnostic of acute viral infection. Serologic diagnosis is based on a more than fourfold rise in IgG antibody concentration when acute- and convalescent-phase serum samples are analyzed at the same time.

Immunofluorescence, hemadsorption, and hemagglutination assays for antiviral antibodies are labor-intensive and have been replaced by enzyme-linked immunosorbent assays (ELISAs), which generally use the specific viral proteins most frequently targeted by the antibody response. The proteins are purified from virus-infected cells or produced by recombinant DNA technology and are attached to a solid phase, where they can be incubated with serum, washed to eliminate nonspecific antibodies, and allowed to react with an enzyme-linked reagent to detect human IgG or IgM antibody specifically adhering to the viral antigen. The amount of antibody can then be quantitated by the intensity of a color reaction mediated by the linked enzyme. ELISAs can be sensitive and automated. Western blots can simultaneously confirm the presence of antibody to multiple specific viral proteins. The proteins are separated by size and transferred to an inert membrane, where they are incubated with serum antibodies. Western blots have an internal specificity control because the level of reactivity for viral proteins can be compared with that for cellular proteins in the same sample. Western blots require individual evaluation and are inherently difficult to quantitate or automate.

Isolation of virus in tissue culture depends on infection and replication in susceptible cells. Growth of virus in cell cultures can frequently be identified by effects on cell morphology under light microscopy. For example, HSV produces a typical cytopathic effect in rabbit kidney cells within 3 days. Other viral cytopathic effects may not be as diagnostically distinctive. Identification usually requires confirmation by staining with virus-specific monoclonal antibodies. The efficiency and speed of virus identification can be enhanced by combining short-term culture with immune detection. In assays with “shell vials”

of tissue culture cells growing on a coverslip, viral infection can be detected by staining with a monoclonal antibody to a specific viral protein expressed early in viral replication. Thus, virus-infected cells can be detected within hours or days of inoculation, whereas several rounds of infection would be required to produce visible cytopathic effects.

Isolation of virus in tissue culture also depends on the collection of specimens from appropriate sites and the rapid transport of these specimens in appropriate medium to the virology laboratory (Chap. 150e). Rapid transport maintains viral viability and limits bacterial and fungal overgrowth. Enveloped viruses are generally more sensitive to freezing and thawing than nonenveloped viruses. The most appropriate site for culture depends on the pathogenesis of the virus in question. Nasopharyngeal, tracheal, or endobronchial aspirates are most appropriate for the identification of respiratory viruses. Sputum cultures generally are less appropriate because bacterial contamination and viscosity threaten tissue-culture cell viability. Aspirates of vesicular fluid are useful for isolation of HSV and VZV. Nasopharyngeal aspirates and stool specimens may be useful when the patient has fever and a rash and an enteroviral infection is suspected. Adenoviruses can be cultured from the urine of patients with hemorrhagic cystitis. CMV can frequently be isolated from cultures of urine or buffy coat. Biopsy material can be effectively cultured when viruses infect major organs, as in HSV encephalitis or adenovirus pneumonia.

The isolation of a virus does not necessarily establish disease causality. Viruses can persistently or intermittently colonize normal human mucosal surfaces. Saliva can be positive for herpesviruses, and normal urine samples can be positive for CMV. Isolations from blood, cerebrospinal fluid (CSF), or tissue are more often diagnostic of significant viral infection.

Another method aimed at increasing the speed of viral diagnosis is direct testing for antigen or cytopathic effects. Virus-infected cells from the patient may be detected by staining with virus-specific monoclonal antibodies. For example, epithelial cells obtained by nasopharyngeal aspiration can be stained with a variety of specific monoclonal antibodies to identify the specific infecting respiratory virus. Antigen and serologic assays can be multiplexed to detect multiple analytes simultaneously by coupling of reagents to color-coded beads for each analyte and detection by flow cytometry.

Nucleic acid amplification techniques bring speed, sensitivity, and specificity to diagnostic virology. The ability to directly amplify minute amounts of viral nucleic acids in specimens means that detection no longer depends on viable virus and its replication. For example, amplification and detection of HSV nucleic acids in the CSF of patients with HSV encephalitis is a more sensitive detection method than culture of virus from CSF. The extreme sensitivity of these tests can be a problem, because subclinical infection or contamination can lead to false-positive results. Detection of viral nucleic acids does not necessarily indicate virus-induced disease.

Measurement of the amount of viral RNA or DNA in peripheral blood is an important means for determining whether a patient is at increased risk for virus-induced disease and for evaluating clinical responses to antiviral chemotherapy. Nucleic acid technologies for RNA quantification are routinely used in AIDS patients to evaluate responses to antiviral agents and to detect viral resistance or noncompliance with therapy. Virus-load measurements are also useful for evaluating the treatment of patients with HBV and HCV infections. Nucleic acid testing or direct staining with CMV-specific monoclonal antibodies to quantitate virus-infected cells in the peripheral blood (CMV antigenemia) is useful for identifying immunosuppressed patients who may be at risk for CMV-induced disease.

DRUG TREATMENT FOR VIRAL INFECTIONS

Multiple steps in the life cycles of viruses can be effectively targeted by antiviral drugs (Chaps. 215e and 216). Nucleoside and nonnucleoside reverse transcriptase inhibitors prevent HIV provirus synthesis, whereas protease inhibitors block maturation of the HIV and HCV polyprotein after infection of the cell. Enfuvirtide is a small peptide derived from HIV gp41 that acts before cell infection by preventing a conformational change required for initial fusion of the virus with the cell membrane. Raltegravir is an integrase inhibitor that is approved

for use with other anti-HIV drugs. Amantadine and rimantadine inhibit the influenza M2 protein, preventing release of viral RNA early during infection, whereas zanamivir and oseltamivir inhibit the influenza neuraminidase, which is necessary for the efficient release of mature virions from infected cells.

Viral genomes can evolve resistance to drugs by mutation and selection, by recombination with a drug-resistant virus, or (in the case of influenza virus and other segmented RNA viral genomes) by reassortment. The emergence of drug-resistant strains can limit the efficacy of antiviral therapy. As in antibacterial therapy, excessive and inappropriate use of antiviral therapy can select for the emergence of drug-resistant strains. HIV genotyping is a rapid method for identifying drug-resistant viruses. Resistance to reverse transcriptase or protease inhibitors has been associated with specific mutations in the reverse transcriptase or protease genes. Identification of these mutations by polymerase chain reaction amplification and nucleic acid sequencing can be clinically useful for determining which antiviral agents may still be effective. Drug resistance also can arise in herpesviruses but is a less common clinical problem.

IMMUNIZATION FOR THE PREVENTION OF VIRAL INFECTIONS

Viral vaccines are among the outstanding accomplishments of medical science. Smallpox has been eradicated except as a potential weapon of biological warfare or bioterrorism (Chap. 261e). Poliovirus eradication may soon follow. Measles can be contained or eliminated. Excess mortality due to influenza virus epidemics can be prevented, and the threat of influenza pandemics can be decreased by contemporary killed or live attenuated influenza vaccines. Mumps, rubella, and chickenpox are well controlled by childhood vaccination in the developed world. Reimmunization of mature adults can be used to control herpes zoster. New rotavirus vaccines can have a major impact on this leading cause of gastroenteritis and prominent cause of childhood death worldwide. Widespread HBV vaccination has dramatically lowered the frequency of acute and chronic hepatitis and is expected to lead to a dramatic decrease in the incidence of hepatocellular carcinoma. The HPV vaccine was the first vaccine specifically licensed to prevent virus-induced cancer. Use of purified proteins, genetically engineered live-virus vaccines, and recombinant DNA-based strategies will make it possible to immunize against severe infections with other viruses. The development of effective HIV and HCV vaccines is complicated by the high mutation rate of viral RNA polymerase and reverse transcriptase, the population-based and individual divergence of HIV or HCV genomes, and repeated high-level exposure in some populations. Concerns about the use of smallpox and other viruses as weapons necessitate maintenance of immunity to agents that are not encountered naturally.

VIRUSES AS NOVEL THERAPEUTIC TOOLS OR AGENTS

Viruses are being used experimentally to deliver biotherapeutic agents or novel vaccines. Foreign genes can be inserted into viral nucleic acids, and the recombinant virus vectors can be used to infect the patient or the patient's cells *ex vivo*. Retrovirus integration into the cell genome has been used to functionally replace the abnormal gene in T cells of patients with severe combined immunodeficiency, thereby restoring immune function. Recombinant adenovirus, AAV, and retroviruses are being explored for use in diseases due to single-gene defects, such as cystic fibrosis and hemophilia. AAV carrying a lipoprotein lipase gene is now being used in Europe to treat a rare lipid-processing disease and is the first gene therapy approved for clinical use. Recombinant poxviruses, adenoviruses, and influenza viruses are also being used experimentally as vaccine vectors. Viral vectors are being tested experimentally for the expression of cytokines that can enhance immunity against tumor cells or for the expression of proteins that can increase the sensitivity of tumor cells to chemotherapy. HSV deficient for replication in resting cells is being used to selectively kill proliferating glioblastoma cells after injections into CNS tumors. For improved safety, nonreplicating viruses are frequently used in clinical trials. Potential adverse events associated with virus-mediated gene transfer include the induction of inflammatory and antiviral immune responses. Instances of retrovirus-induced human malignancies have raised concerns about the safety of retroviral gene therapy vectors.

The field of antiviral therapy—both the number of antiviral drugs and our understanding of their optimal use—historically has lagged behind that of antibacterial treatment, but significant progress has been made in recent years on new drugs for several viral infections. The development of antiviral drugs poses several challenges. Viruses replicate intracellularly and often use host cell enzymes, macromolecules, and organelles for synthesis of viral particles. Therefore, useful antiviral compounds must discriminate between host and viral functions with a high degree of specificity; agents without such selectivity are likely to be too toxic for clinical use.

Significant progress has also been made in the development of laboratory assays to assist clinicians in the appropriate use of antiviral drugs. Phenotypic and genotypic assays for resistance to antiviral drugs are becoming more widely available, and correlations of laboratory results with clinical outcomes are being better defined. Of particular note has been the development of highly sensitive and specific methods that measure the concentration of virus in blood (*virus load*) and permit direct assessment of the antiviral effect of a given drug regimen in that host site. Virus load measurements have been useful in recognizing the risk of disease progression in patients with viral infections and in identifying patients for whom antiviral chemotherapy might be of greatest benefit. As with any *in vitro* laboratory test, results are highly dependent on and likely vary with the laboratory techniques used.

Information regarding the pharmacodynamics of antiviral drugs, and particularly the relationship of concentration effects to efficacy, has been slow to develop but is also expanding. However, assays to measure concentrations of antiviral drugs, especially of their active moieties within cells, are still primarily research procedures not widely available to clinicians. Thus, there are limited guidelines for adjusting dosages of antiviral agents to maximize antiviral activity and minimize toxicity. Consequently, clinical use of antiviral drugs must be accompanied by particular vigilance for unanticipated adverse effects.

Like that of other infections, the course of viral infections is profoundly affected by interplay between the pathogen and a complex set of host defenses. The presence or absence of preexisting immunity, the ability to mount humoral and/or cell-mediated immune responses, and the stimulation of innate immunity are important determinants of the outcome of viral infections. The state of the host's defenses needs to be considered when antiviral agents are used or evaluated.

As with any therapy, the optimal use of antiviral compounds requires a specific and timely diagnosis. For some viral infections, such as herpes zoster, the clinical manifestations are so characteristic that a diagnosis can be made on clinical grounds alone. For other viral infections, such as influenza A, epidemiologic information (e.g., the documentation of a community-wide influenza outbreak) can be used to make a presumptive diagnosis with a high degree of accuracy. However, for most of the remaining viral infections, including herpes simplex encephalitis, cytomegaloviral infections other than retinitis, and enteroviral infections, diagnosis on clinical grounds alone cannot be accomplished with certainty. For such infections, rapid viral diagnostic techniques are of great importance. Considerable progress has also been made in recent years in the development of such tests, which are now widely available for a number of viral infections.

Despite these complexities, the efficacy of a number of antiviral compounds has been clearly established in rigorously conducted and controlled studies. As summarized in [Table 215e-1](#), this chapter reviews the antiviral drugs that are currently approved or are likely to be considered for approval in the near future for use against viral infections other than those caused by HIV. [Antiretroviral drugs are reviewed in Chap. 226.](#)

ZANAMIVIR, OSELTAMIVIR, PERAMIVIR, AND LANINAMIVIR

Zanamivir and oseltamivir are inhibitors of the influenza viral neuraminidase enzyme, which is essential for release of virus from infected cells and for its subsequent spread throughout the respiratory tract of the infected host. The enzyme cleaves terminal sialic acid residues and thus destroys the cellular receptors to which the viral hemagglutinin attaches. Zanamivir and oseltamivir are sialic acid transition-state analogues and are highly active and specific inhibitors of the neuraminidases of both influenza A and B viruses. The antineuraminidase activity of the two drugs is similar, although zanamivir has somewhat greater *in vitro* activity against influenza B virus. Zanamivir may also be active against certain strains of influenza A virus that are resistant to oseltamivir. Both zanamivir and oseltamivir act through competitive and reversible inhibition of the active site of influenza A and B viral neuraminidases and have relatively little effect on mammalian cell enzymes.

Oseltamivir phosphate is an ethyl ester prodrug that is converted to oseltamivir carboxylate by esterases in the liver. Orally administered oseltamivir has a bioavailability of >60% and a plasma half-life of 7–9 h. The drug is excreted unmetabolized, primarily by the kidneys. Zanamivir has low oral bioavailability and is administered orally via a hand-held inhaler. By this route, ~15% of the dose is deposited in the lower respiratory tract, and low plasma levels of the drug are detected. The toxicities most frequently encountered with orally administered oseltamivir are nausea, gastrointestinal discomfort, and (less commonly) vomiting. Gastrointestinal discomfort is usually transient and is less likely if the drug is administered with food. Neuropsychiatric events (delirium, self-injury) have been reported in children who have been taking oseltamivir, primarily in Japan. Zanamivir is orally inhaled and is generally well tolerated, although exacerbations of asthma may occur. An IV formulation of zanamivir is under development and is available from GlaxoSmithKline as part of clinical trials.

Inhaled zanamivir and orally administered oseltamivir have been effective in the treatment of naturally occurring, uncomplicated influenza A or B in otherwise healthy adults. In placebo-controlled studies, illness has been shortened by 1.0–1.5 days of therapy with either of these drugs when treatment is administered within 2 days of onset of symptoms. Pooled analyses of clinical studies of oseltamivir suggest that treatment may reduce the likelihood of hospitalizations and of certain respiratory tract complications associated with influenza, and observational studies suggest that oseltamivir may reduce mortality rates associated with influenza A outbreaks ([Chap. 224](#)). Once-daily inhaled zanamivir or once-daily orally administered oseltamivir can provide prophylaxis against laboratory-documented influenza A- and influenza B-associated illness.

Resistance to the neuraminidase inhibitors may develop by changes in the viral neuraminidase enzyme, by changes in the hemagglutinin that make it more resistant to the actions of the neuraminidase, or by both mechanisms. Isolates that are resistant to oseltamivir—most commonly through the H275Y mutation, which leads to a change from histidine to tyrosine at that residue in the neuraminidase—remain sensitive to zanamivir. Certain mutations impart resistance to both oseltamivir and zanamivir (e.g., I223R, which leads to a change from isoleucine to arginine). Because the mechanisms of action of the neuraminidase inhibitors differ from those of the adamantanes (see below), zanamivir and oseltamivir are active against strains of influenza A virus that are resistant to amantadine and rimantadine.

Appropriate use of antiviral agents against influenza viruses depends on a knowledge of the resistance patterns of circulating viruses. As of this writing, currently circulating influenza A/H1N1 and H3N2 viruses (2013–2014) were sensitive to zanamivir and oseltamivir, with a few exceptions for oseltamivir. Up-to-date information on patterns of resistance to antiviral drugs is available from the Centers for Disease Control and Prevention (CDC) at www.cdc.gov/flu.

TABLE 215e-1 ANTIVIRAL CHEMOTHERAPY AND CHEMOPROPHYLAXIS

Infection	Drug	Route	Dosage	Comment
Influenza A and B: treatment	Oseltamivir	Oral	Adults: 75 mg bid × 5 d Children 1–12 years: 30–75 mg bid, depending on weight, ^a × 5 d	When started within 2 d of onset in uncomplicated disease, zanamivir and oseltamivir reduce symptom duration by 1.0–1.5 and 1.3 d, respectively. Their effectiveness in prevention or treatment of complications is unclear, although some analyses suggest that oseltamivir may reduce the frequency of respiratory tract complications and hospitalizations. Oseltamivir's side effects of nausea and vomiting can be reduced in frequency by drug administration with food. Zanamivir may exacerbate bronchospasm in patients with asthma. Amantadine and rimantadine are not recommended for routine use unless antiviral susceptibilities are known because of widespread resistance in A/H3N2 viruses since 2005–2006 and in pandemic A/H1N1 viruses in 2009–2010. Their efficacy in treatment of uncomplicated disease caused by sensitive viruses has been similar to that of neuraminidase inhibitors.
	Zanamivir	Inhaled orally	Adults and children ≥7 years: 10 mg bid × 5 d	
Influenza A: treatment	Amantadine ^b	Oral	Adults: 100 mg qd or bid × 5–7 d Children 1–9 years: 5 mg/kg per day (maximum, 150 mg/d) × 5–7 d	
	Rimantadine ^b	Oral	100 mg qd or bid × 5–7 d in adults	
Influenza A and B: prophylaxis	Oseltamivir	Oral	Adults: 75 mg/d Children ≥1 year: 30–75 mg/d, depending on weight ^a	Prophylaxis must be continued for the duration of exposure and can be administered simultaneously with inactivated vaccine. Unless the sensitivity of isolates is known, neither amantadine nor rimantadine is currently recommended for prophylaxis or therapy.
	Zanamivir	Inhaled orally	Adults and children ≥5 years: 10 mg/d	
Influenza A: prophylaxis	Amantadine ^b or rimantadine ^b	Oral	Adults: 200 mg/d Children 1–9 years: 5 mg/kg per day (maximum, 150 mg/d)	
RSV infection	Ribavirin	Small-particle aerosol	Administered 12–18 h/d from a reservoir containing 20 mg/mL × 3–6 d	Use of ribavirin is to be considered for treatment of infants and young children hospitalized with RSV pneumonia and bronchiolitis, according to the American Academy of Pediatrics.
CMV disease	Ganciclovir	IV	5 mg/kg bid × 14–21 d; then 5 mg/kg per day as maintenance dose	Ganciclovir, valganciclovir, foscarnet, and cidofovir are approved for treatment of CMV retinitis in patients with AIDS. They are also used for colitis, pneumonia, or “wasting” syndrome associated with CMV and for prevention of CMV disease in transplant recipients.
	Valganciclovir	Oral	900 mg bid × 21 d; then 900 mg/d as maintenance dose	Valganciclovir has largely supplanted oral ganciclovir and is frequently used in place of IV ganciclovir.
	Foscarnet	IV	60 mg/kg q8h × 14–21 d; then 90–120 mg/kg per day as maintenance dose	Foscarnet is not myelosuppressive and is active against acyclovir- and ganciclovir-resistant herpesviruses.
	Cidofovir	IV	5 mg/kg once weekly × 2 weeks, then once every other week; given with probenecid and hydration	
	Fomivirsen	Intravitreal	330 mg on days 1 and 15 followed by 330 mg monthly as maintenance	Fomivirsen has reduced the rate of progression of CMV retinitis in patients in whom other regimens have failed or have not been well tolerated. The major form of toxicity is ocular inflammation.
Varicella: immunocompetent host	Acyclovir	Oral	20 mg/kg (maximum, 800 mg) 4 or 5 times daily × 5 d	Treatment confers modest clinical benefit when administered within 24 h of rash onset.
	Valacyclovir	Oral	Children 2–18 years: 20 mg/kg tid (not to exceed 1 g tid) × 5 d	
Varicella: immunocompromised host	Acyclovir	IV	10 mg/kg q8h × 7 d	A change to oral valacyclovir can be considered once fever has subsided if there is no evidence of visceral involvement.
Herpes simplex encephalitis	Acyclovir	IV	10 mg/kg q8h × 14–21 d	Results are optimal when therapy is initiated early. Some authorities recommend treatment for 21 d to prevent relapses.
Neonatal herpes simplex	Acyclovir	IV	20 mg/kg q8h × 14–21 d	Serious morbidity is common despite therapy. Prolonged oral administration after initial IV therapy has been suggested because of long-term sequelae associated with cutaneous recurrences of HSV infection.

(Continued)

TABLE 215e-1 ANTIVIRAL CHEMOTHERAPY AND CHEMOPROPHYLAXIS (CONTINUED)

Infection	Drug	Route	Dosage	Comment
Genital herpes simplex, primary: treatment	Acyclovir	IV	5 mg/kg q8h × 5–10 d	The IV route is preferred for infections severe enough to warrant hospitalization or with neurologic complications.
		Oral	400 mg tid or 200 mg 5 times daily × 7–10 d	The oral route is preferred for patients whose condition does not warrant hospitalization. Adequate hydration must be maintained.
		Topical	5% ointment; 4–6 applications daily × 7–10 d	Topical use—largely supplemented by oral therapy—may obviate systemic administration to pregnant women. Systemic symptoms and untreated areas are not affected.
	Valacyclovir	Oral	1 g bid × 7–10 d	Valacyclovir appears to be as effective as acyclovir but can be administered less frequently.
Genital herpes simplex, recurrent: treatment	Famciclovir	Oral	250 mg tid × 7–10 d ^c	Famciclovir appears to be similar in effectiveness to acyclovir.
	Acyclovir	Oral	400 mg tid × 5 d or 800 mg tid × 2 d	The clinical effect is modest and is enhanced if therapy is initiated early. Treatment does not affect recurrence rates.
Genital herpes simplex, recurrent: suppression	Famciclovir	Oral	125 mg bid × 5 d, 1000 mg bid × 1 d, or 500 mg once, then 250 mg PO bid × 3 doses	
	Valacyclovir	Oral	500 mg bid × 3 d or 1 g once a day × 5 d	
	Acyclovir	Oral	400 mg bid	Suppressive therapy is recommended only for patients with at least 6–10 recurrences per year. “Breakthrough” occasionally takes place, and asymptomatic shedding of virus occurs. The need for suppressive therapy should be reevaluated after 1 year. Suppression with valacyclovir reduces transmission of genital HSV among virus-discordant couples.
Mucocutaneous herpes simplex in immunocompromised host: treatment	Acyclovir	Oral	500–1000 mg/d or 250–500 mg bid	
		IV	5 mg/kg q8h × 7–14 d	
		Topical	5% ointment; 4–6 applications daily × 7 d or until healed	The choice of the IV or oral route and the duration of therapy depend on the severity of infection and the patient’s ability to take oral medication. Oral or IV treatment has supplanted topical therapy except for small, easily accessible lesions. Foscarnet is used for acyclovir-resistant viruses.
	Valacyclovir	Oral	1 g tid × 7–10 d ^c	
Mucocutaneous herpes simplex in immunocompromised host: prevention of recurrence during intense immunosuppression	Acyclovir	Oral	400 mg 2–5 times daily or 800 mg bid	Treatment is administered during periods when intense immunosuppression is expected—e.g., during antitumor chemotherapy or after transplantation—and is usually continued for 2–3 months.
		IV	5 mg/kg q12h	
	Valacyclovir	Oral	500 mg to 1 g bid or tid	
	Famciclovir	Oral	500 mg bid ^c	
Herpes simplex orolabialis, recurrent ^e	Penciclovir	Topical	1.0% cream applied q2h during waking hours × 4 d	Treatment shortens healing time and symptom duration by 0.5–1.0 d (versus placebo).
	Valacyclovir	Oral	2 g q12h × 1 d	Therapy begun at earliest symptom reduces disease duration by 1 d.
	Famciclovir ^c	Oral	1500 mg once or 750 mg bid × 1 d	Therapy begun within 1 h of prodrome decreases time to healing by 1.8–2.2 d.
	Docosanol ^f	Topical	10% cream 5 times daily until healed	Application at initial symptoms reduces healing time by 1 d.
Herpes simplex keratitis	Trifluridine	Topical	1 drop of 1% ophthalmic solution q2h while awake (maximum, 9 drops daily)	Therapy should be undertaken in consultation with an ophthalmologist.
	Vidarabine	Topical	0.5-in. ribbon of 3% ophthalmic ointment 5 times daily	
Herpes zoster: immunocompetent host	Valacyclovir	Oral	1 g tid × 7 d	Valacyclovir may be more effective than acyclovir for pain relief; otherwise, it has a similar effect on cutaneous lesions and should be given within 72 h of rash onset.
	Famciclovir	Oral	500 mg q8h × 7 d	The duration of postherpetic neuralgia is shorter than with placebo. Famciclovir showed overall efficacy similar to that of acyclovir in a comparative trial. It should be given ≤72 h after rash onset.
	Acyclovir	Oral	800 mg 5 times daily × 7–10 d	Acyclovir causes faster resolution of skin lesions than placebo and provides some relief of acute symptoms if given within 72 h of rash onset. Combined with tapering doses of prednisone, acyclovir improves quality-of-life outcomes.

(Continued)

TABLE 215e-1 ANTIVIRAL CHEMOTHERAPY AND CHEMOPROPHYLAXIS (CONTINUED)

Infection	Drug	Route	Dosage	Comment
Herpes zoster: immunocompromised host	Acyclovir	IV	10 mg/kg q8h × 7 d	Effectiveness in localized zoster is most marked when treatment is given early. Foscarnet may be used for acyclovir-resistant VZV infections.
	Valacyclovir	Oral	800 mg 5 times daily × 7 d	
	Famciclovir	Oral	1 g tid × 7 d ^c	
Herpes zoster ophthalmicus	Acyclovir	Oral	600–800 mg 5 times daily × 10 d	Treatment reduces ocular complications, including ocular keratitis and uveitis.
	Valacyclovir	Oral	1 g tid × 7 d	
	Famciclovir	Oral	500 mg tid × 7 d	
Condyloma acuminatum	IFN-α2b	Intralesional	1 million units per wart (maximum of 5) thrice weekly × 3 weeks	Intralesional treatment frequently results in regression of warts, but lesions often recur. Parenteral administration may be useful if lesions are numerous.
	IFN-αn3	Intralesional	250,000 units per wart (maximum of 10) twice weekly × up to 8 weeks	
Chronic hepatitis B	IFN-α2b	SC	5 million units daily or 10 million units thrice weekly × 16–24 weeks	HBeAg and DNA are eliminated in 33–37% of cases. Histopathologic improvement is also seen.
	Pegylated IFN-α2a	SC	180 µg weekly × 48 weeks	ALT levels return to normal in 39% of patients, and histologic improvement occurs in 38%.
	Lamivudine	Oral	100 mg/d × 12–18 months; 150 mg bid as part of therapy for HIV infection	Lamivudine monotherapy is well tolerated and effective in reduction of HBV DNA levels, normalization of ALT levels, and improvement in histopathology. However, resistance develops in 24% of recipients when lamivudine is used as monotherapy for 1 year.
	Adefovir dipivoxil	Oral	10 mg/d × 48 weeks	A return of ALT levels to normal is documented in 48–72% of recipients and improved liver histopathology in 53–64%. Adefovir is effective in lamivudine-resistant hepatitis B. Renal function should be monitored.
	Entecavir	Oral	0.5 mg/d × 48 weeks (1 mg/d if HBV is resistant to lamivudine)	Normalization of ALT is seen in 68–78% of recipients and loss of HBeAg in 21%. Entecavir is active against lamivudine-resistant HBV.
	Telbivudine	Oral	600 mg/d × 52 weeks	HBV DNA is reduced by >5 log ₁₀ copies/mL along with normalization of ALT levels in 74–77% of patients and improved histopathology in 65–67%. Resistance develops in 9–22% of patients after 2 years of therapy. Elevated CPK levels and myopathy may occur.
	Tenofovir	Oral	300 mg/d × 48 weeks	ALT levels return to normal in 68–76% of patients, and liver histopathology improves in 72–74%. Resistance is uncommon with up to 2 years of therapy.
Chronic hepatitis C	IFN-α2a or IFN-α2b	SC	3 million units thrice weekly × 12–24 months	SVRs are noted in 20–30% of patients. Normalization of ALT levels and improvements in liver histopathology are also seen.
	IFN-α2b/ribavirin	SC (IFN)/oral (ribavirin)	3 million units thrice weekly (IFN)/1000–1200 mg daily (ribavirin) × 6–12 months	Combination therapy results in SVR in up to 40–50% of recipients.
	Pegylated IFN-α2b	SC	1.5 µg weekly × 48 weeks	The slower clearance of pegylated IFNs than of standard IFNs permits once-weekly administration. Pegylated formulations appear to be superior to standard IFNs in efficacy, both as monotherapy and in combination with ribavirin, and have largely supplanted standard IFNs in treatment of hepatitis C. SVRs were seen in 42–51% of patients infected with HCV genotype 1 and in 76–82% of those infected with genotype 2 or 3.
	Pegylated IFN-α2a	SC	180 µg weekly × 48 weeks	
	Pegylated IFN-α2b/ribavirin	SC (IFN)/oral (ribavirin)	1.5 µg/kg weekly (IFN)/800–1400 mg daily (ribavirin) × 24–48 weeks	
	Pegylated IFN-α2a/ribavirin	SC (IFN)/oral (ribavirin)	180 µg weekly (IFN)/800–1200 mg daily (ribavirin) × 24–48 weeks	
IFN-alfacon	SC	9–15 µg thrice weekly × 6–12 months	Doses of 9 and 15 µg are equivalent to IFN-α2a and IFN-α2b doses of 3 million units and 5 million units, respectively.	

(Continued)

TABLE 215e-1 ANTIVIRAL CHEMOTHERAPY AND CHEMOPROPHYLAXIS (CONTINUED)

Infection	Drug	Route	Dosage	Comment
	Sofosbuvir ^a	Oral	HCV genotypes 1, 4, 5, and 6: 400 mg qd with daily weight-based ribavirin (1000 mg [<75 kg] to 1200 mg [>75 kg]) and weekly pegylated IFN for 12 weeks. Genotypes 2 and 3: 400 mg qd with daily weight-based ribavirin for 12 and 24 weeks, respectively	Sofosbuvir is generally well tolerated, and most common side effects have been attributable to concomitantly administered IFN and ribavirin. Sofosbuvir is recommended in triple combination with pegylated IFN and ribavirin as first-line therapy for genotypes 1, 4, 5, and 6, with SVRs in 89–97% of treatment-naïve patients, and in double combination with ribavirin for genotypes 2 and 3.
	Simeprevir ^a	Oral	Alternative regimen for genotypes 1 and 4: 150 mg qd for 12 weeks plus daily ribavirin and weekly pegylated IFN for 24 weeks and for 24–48 weeks, respectively	Simeprevir has supplanted the first-generation protease inhibitors boceprevir and telaprevir. Its metabolism by cytochrome CYP3A can result in interactions with other drugs. Photosensitivity and reversible hyperbilirubinemia are associated toxicities. Testing for the Q80K-resistant variant should be carried out since this variant is present in one-third of HCV genotype 1a infections. Triple combinations with pegylated IFN and ribavirin result in SVRs in 80% of genotype 1 infections without Q80K.
Chronic hepatitis D	IFN- α 2a or IFN- α 2b	SC	9 million units thrice weekly \times 12 months	The overall efficacy and the optimal regimen and duration of therapy are not fully established. Sustained SVRs have been seen in 25–30% of patients for IFN- α and in 17–43% for pegylated IFN- α .
	Pegylated IFN- α 2b	SC	1.5 μ g weekly \times 48 weeks	
	Pegylated IFN- α 2a	SC	180 μ g weekly \times 48 weeks	

^aFor detailed weight recommendations and for children <1 year of age, see www.cdc.gov/flu/professionals/antivirals/summary-clinicians.htm. ^bAmantadine and rimantadine are not recommended for routine use because of widespread resistance in currently circulating A/H3N2 and pandemic A/H1N1 viruses. Their use may be considered if sensitivities become reestablished. ^cNot approved for this indication by the U.S. Food and Drug Administration (FDA). ^dApproved by the FDA for treatment of HIV-infected individuals. ^eAcyclovir suspension (15 mg/kg PO, to a maximum of 200 mg per dose) given for 7 d has been reported to be effective in treatment of primary herpetic gingivostomatitis in children. ^fActive ingredient: benzyl alcohol. Available without prescription. ^gConsult www.hcvguidelines.org for recommendations regarding treatment of null or partial responders to IFN regimens or of patients ineligible to receive IFN.

Abbreviations: ALT, alanine aminotransferase; CMV, cytomegalovirus; CPK, creatine phosphokinase; HBeAg, hepatitis B e antigen; HBV, hepatitis B virus; HCV, hepatitis C virus; HSV, herpes simplex virus; IFN, interferon; RGT, response-guided therapy; RSV, respiratory syncytial virus; SVR, sustained virologic response; VZV, varicella-zoster virus.

Zanamivir and oseltamivir have been approved by the U.S. Food and Drug Administration (FDA) for treatment of influenza in adults and in children (those ≥ 7 years old for zanamivir and those ≥ 1 year old for oseltamivir) who have been symptomatic for ≤ 2 days. Oseltamivir is approved for prophylaxis of influenza in individuals ≥ 1 year of age and zanamivir for those ≥ 5 years of age (Table 215e-1). Guidelines for the use of oseltamivir in children <1 year of age can be accessed through the CDC website, as noted in the footnote to Table 215e-1.

Peramivir is an investigational neuraminidase inhibitor that can be administered intravenously to patients for whom such an intervention is considered necessary. It has been approved in Japan, China, and South Korea but not in the United States, where it has been available in clinical trials through BioCryst Pharmaceuticals. Oseltamivir-resistant viruses generally exhibit reduced sensitivity to peramivir.

Laninamivir octonate is an investigational neuraminidase that has been approved in Japan. It is the prodrug of laninamivir, which is administered by oral inhalation and has a prolonged half-life of ~ 3 days. In limited studies, it has been investigated as single-dose therapy for influenza; its effects were similar to those obtained with multiple dosing of zanamivir or oseltamivir.

AMANTADINE AND RIMANTADINE

Amantadine and the closely related compound rimantadine are primary symmetric amines that have antiviral activity limited to influenza A viruses. Amantadine and rimantadine have a long history of efficacy in the prophylaxis and treatment of influenza A infections in humans. However, high frequencies of resistance to these drugs were noted among influenza A/H3N2 viruses in the 2005–2006 influenza season and continued to be seen in 2013–2014. The pandemic A/H1N1 viruses that circulated in 2009–2010 were also resistant to amantadine and rimantadine, and circulating influenza A/H1N1 viruses in the 2013–2014 season were largely resistant. Therefore, these agents are no

longer recommended unless the sensitivity of the particular isolate of influenza A virus is known, in which case their use may be considered. Amantadine and rimantadine act through inhibition of the ion channel function of the influenza A M2 matrix protein, on which uncoating of the virus depends. A substitution of a single amino acid at critical sites in the M2 protein can result in a virus that is resistant to amantadine and rimantadine.

Amantadine and rimantadine have been shown to be effective in the prophylaxis of influenza A in large-scale studies of young adults and in less extensive studies of children and elderly persons. In such studies, efficacy rates of 55–80% in the prevention of influenza-like illness were noted, and even higher rates were reported when virus-specific attack rates were calculated. Amantadine and rimantadine have also been found to be effective in the treatment of influenza A infection in studies involving predominantly young adults and, to a lesser extent, children. Administration of these compounds within 24–72 h after the onset of illness has resulted in a reduction of the duration of signs and symptoms by $\sim 50\%$ compared with that in placebo recipients. The effect on signs and symptoms of illness is superior to that of commonly used antipyretic-analgesic agents. Only anecdotal reports are available concerning the efficacy of amantadine or rimantadine in the prevention or treatment of complications of influenza (e.g., pneumonia).

Amantadine and rimantadine are available only in oral formulations and are ordinarily administered to adults once or twice daily, with a dosage of 100–200 mg/d. Despite their structural similarities, the two compounds have different pharmacokinetics. Amantadine is not metabolized and is excreted almost entirely by the kidneys, with a half-life of 12–17 h and peak plasma concentrations of 0.4 μ g/mL. In contrast, rimantadine is extensively metabolized to hydroxylated derivatives and has a half-life of 30 h. Only 30–40% of an orally administered dose of rimantadine is recovered in the urine. The peak plasma levels of rimantadine are approximately half those of amantadine,

but rimantadine is concentrated in respiratory secretions to a greater extent than amantadine. For prophylaxis, the compounds must be administered daily for the period at risk (i.e., duration of the exposure). For therapy, amantadine or rimantadine is generally administered for 5–7 days.

Although these compounds are generally well tolerated, 5–10% of amantadine recipients experience mild central nervous system side effects consisting primarily of dizziness, anxiety, insomnia, and difficulty in concentrating. These effects are rapidly reversible upon cessation of the drug's administration. At a dose of 200 mg/d, rimantadine is better tolerated than amantadine; in a large-scale study of young adults, adverse effects were no more frequent among rimantadine recipients than among placebo recipients. Seizures and worsening of congestive heart failure have also been reported in patients treated with amantadine, although a causal relationship has not been established. The dosage of amantadine should be reduced to 100 mg/d in patients with renal insufficiency—i.e., a creatinine clearance rate (Cr_{cl}) of <50 mL/min—and in the elderly. A rimantadine dose of 100 mg/d should be used for patients with a Cr_{cl} of <10 mL/min and for the elderly.

RIBAVIRIN

Ribavirin is a synthetic nucleoside analogue that inhibits a wide range of RNA and DNA viruses. The mechanism of action of ribavirin is not completely defined and may be different for different groups of viruses. Ribavirin-5'-monophosphate blocks the conversion of inosine-5'-monophosphate to xanthosine-5'-monophosphate and interferes with the synthesis of guanine nucleotides as well as with that of both RNA and DNA. Ribavirin-5'-monophosphate also inhibits capping of virus-specific messenger RNA in certain viral systems.

Ribavirin administered as a small-particle aerosol to young children hospitalized with respiratory syncytial virus (RSV) infection has been clinically beneficial and has improved oxygenation in some studies (7 of 11). Although ribavirin has been approved for treatment of infants hospitalized with RSV infection, the American Academy of Pediatrics has recommended that it be considered on an individual basis rather than used routinely in that setting. Aerosolized ribavirin has also been administered to older children and adults (including immunosuppressed patients) with severe RSV and parainfluenza virus infections and to older children and adults with influenza A or B infection, but the benefit of this treatment, if any, is unclear. In RSV infections in immunosuppressed patients, ribavirin has been given in combination with anti-RSV immunoglobulins.

Orally administered ribavirin has not been effective in the treatment of influenza A virus infections. IV or oral ribavirin has reduced mortality rates among patients with Lassa fever; it has been particularly effective in this regard when given within the first 6 days of illness. IV ribavirin has been reported to be of clinical benefit in the treatment of hemorrhagic fever with renal syndrome caused by Hantaan virus and as therapy for Argentinean hemorrhagic fever. Oral ribavirin has also been recommended for the treatment and prophylaxis of Congo-Crimean hemorrhagic fever. Use of IV ribavirin in patients with hantavirus pulmonary syndrome in the United States has not been associated with clear-cut benefits.

Oral administration of ribavirin reduces serum aminotransferase levels in patients with chronic hepatitis C virus (HCV) infection; because it appears not to reduce serum HCV RNA levels, the mechanism of this effect is unclear. The drug provides added benefit when given by mouth in doses of 800–1200 mg/d in combination with interferon (IFN) α 2b or α 2a (see below), and the triple combination of ribavirin, IFN, and sofosbuvir or simeprevir has been approved for the treatment of patients with chronic HCV infection (see below). Recent data suggest that oral ribavirin may be beneficial in resolution of chronic hepatitis E infection associated with organ transplantation. Large oral doses of ribavirin (800–1000 mg/d) have been associated with reversible hematopoietic toxicity. This effect has not been observed with aerosolized ribavirin, apparently because little drug is absorbed systemically. Aerosolized administration of ribavirin is generally well tolerated but occasionally is associated with bronchospasm, rash, or conjunctival irritation. It should be administered under

close supervision—particularly in the setting of mechanical ventilation, where precipitation of the drug is possible. Health care workers exposed to the drug have experienced minor toxicity, including eye and respiratory tract irritation. Because ribavirin is mutagenic, teratogenic, and embryotoxic, its use is generally contraindicated in pregnancy. Its administration as an aerosol poses a risk to pregnant health care workers. Because clearance of ribavirin is primarily renal, dose reduction is required in the setting of significant renal dysfunction.

AGENTS OF INVESTIGATIVE INTEREST

DAS181 is an investigational antiviral agent with activity against influenza A and B and parainfluenza viruses. It is a sialidase linked to a respiratory epithelium-anchoring domain. This agent cleaves the terminal sialic acid residues on human respiratory cells, reducing the binding of the aforementioned respiratory viruses. DAS181 is administered by oral inhalation and is being evaluated in the treatment of parainfluenza type 3 infections in recipients of lung and stem cell transplants.

ANTIVIRAL DRUGS ACTIVE AGAINST HERPESVIRUS INFECTIONS

ACYCLOVIR AND VALACYCLOVIR

Acyclovir is a highly potent and selective inhibitor of the replication of certain herpesviruses, including herpes simplex virus (HSV) types 1 and 2, varicella-zoster virus (VZV), and Epstein-Barr virus (EBV). It is relatively ineffective in the treatment of human cytomegalovirus (CMV) infections; however, some studies have indicated effectiveness in the prevention of CMV-associated disease in immunosuppressed patients. Valacyclovir, the L-valyl ester of acyclovir, is converted almost entirely to acyclovir by intestinal and hepatic hydrolysis after oral administration. Valacyclovir offers pharmacokinetic advantages over orally administered acyclovir: it exhibits significantly greater oral bioavailability, results in higher blood levels, and can be given less frequently than acyclovir (two or three rather than five times daily).

The high degree of selectivity of acyclovir is related to its mechanism of action, which requires that the compound first be phosphorylated to acyclovir monophosphate. This phosphorylation occurs efficiently in herpesvirus-infected cells by means of a virus-coded thymidine kinase. In uninfected mammalian cells, little phosphorylation of acyclovir occurs, and the drug is therefore concentrated in herpesvirus-infected cells. Acyclovir monophosphate is subsequently converted by host cell kinases to a triphosphate that is a potent inhibitor of virus-induced DNA polymerase but has relatively little effect on host cell DNA polymerase. Acyclovir triphosphate can also be incorporated into viral DNA, with early chain termination.

Acyclovir is available in IV, oral, and topical forms, while valacyclovir is available in an oral formulation. IV acyclovir is effective in the treatment of mucocutaneous HSV infections in immunocompromised hosts, in whom it reduces time to healing, duration of pain, and virus shedding. When administered prophylactically during periods of intense immunosuppression (e.g., related to chemotherapy for leukemia or transplantation) and before the development of lesions, IV acyclovir reduces the frequency of HSV-associated disease. After prophylaxis is discontinued, HSV lesions recur. IV acyclovir is also effective in the treatment of HSV encephalitis.

Because VZV is generally less sensitive to acyclovir than is HSV, higher doses of acyclovir must be used to treat VZV infections. In immunocompromised patients with herpes zoster, IV acyclovir reduces the frequency of cutaneous dissemination and visceral complications and—in one comparative trial—was more effective than vidarabine. Acyclovir, administered at oral doses of 800 mg five times a day, had a modest beneficial effect on localized herpes zoster lesions in both immunocompromised and immunocompetent patients. Combination of acyclovir with a tapering regimen of prednisone appeared to be more effective than acyclovir alone in terms of quality-of-life outcomes in immunocompetent patients over age 50 with herpes zoster. A comparative study of acyclovir (800 mg PO five times daily) and valacyclovir (1 g PO three times daily) in immunocompetent patients with herpes zoster indicated that the latter drug may be more effective in eliciting the resolution of zoster-associated pain. Orally administered

acyclovir (600 mg five times a day) reduced complications of herpes zoster ophthalmicus in a placebo-controlled trial.

In chickenpox, a modest overall clinical benefit is attained when oral acyclovir therapy is begun within 24 h of the onset of rash in otherwise healthy children (20 mg/kg, up to a maximum of 800 mg, four times a day) or adults (800 mg five times a day). IV acyclovir has also been reported to be effective in the treatment of immunocompromised children with chickenpox.

The most widespread use of acyclovir is in the treatment of genital HSV infections. IV or oral acyclovir or oral valacyclovir has shortened the duration of symptoms, reduced virus shedding, and accelerated healing when used for the treatment of primary genital HSV infections. Oral acyclovir and valacyclovir have also had a modest effect in treatment of recurrent genital HSV infections. However, the failure of treatment of either primary or recurrent disease to reduce the frequency of subsequent recurrences has indicated that acyclovir is ineffective in eliminating latent infection. Documented chronic oral administration of acyclovir for up to 6 years or of valacyclovir for up to 1 year has reduced the frequency of recurrences markedly during therapy; once the drug is discontinued, lesions recur. In one study, suppressive therapy with valacyclovir (500 mg once daily for 8 months) reduced transmission of HSV-2 genital infections among discordant couples by 50%. A modest effect on herpes labialis (i.e., a reduction of disease duration by 1 day) was seen when valacyclovir was administered upon detection of the first symptom of a lesion at a dose of 2 g every 12 h for 1 day. In AIDS patients, chronic or intermittent administration of acyclovir has been associated with the development of HSV and VZV strains resistant to the action of the drug and with clinical failures. The most common mechanism of resistance is a deficiency of the virus-induced thymidine kinase. Patients with HSV or VZV infections resistant to acyclovir have frequently responded to foscarnet.

With the availability of the oral and IV forms, there are few indications for topical acyclovir, although treatment with this formulation has been modestly beneficial in primary genital HSV infections and in mucocutaneous HSV infections in immunocompromised hosts.

Overall, acyclovir is remarkably well tolerated and is generally free of toxicity. The most frequently encountered form of toxicity is renal dysfunction because of drug crystallization, particularly after rapid IV administration or with inadequate hydration. Central nervous system changes, including lethargy and tremors, are occasionally reported, primarily in immunosuppressed patients. However, whether these changes are related to acyclovir, to concurrent administration of other therapy, or to underlying infection remains unclear. Acyclovir is excreted primarily unmetabolized by the kidneys via both glomerular filtration and tubular secretion. Approximately 15% of a dose of acyclovir is metabolized to 9-[(carboxymethoxy)methyl]guanine or other minor metabolites. Reduction in dosage is indicated in patients with a Cr_{cl} of <50 mL/min. The half-life of acyclovir is ~3 h in normal adults, and the peak plasma concentration after a 1-h infusion of a dose of 5 mg/kg is 9.8 μ g/mL. Approximately 22% of an orally administered acyclovir dose is absorbed, and peak plasma concentrations of 0.3–0.9 μ g/mL are attained after administration of a 200-mg dose. Acyclovir penetrates relatively well into the cerebrospinal fluid (CSF), with concentrations approaching half of those found in plasma.

Acyclovir causes chromosomal breakage at high doses, but its administration to pregnant women has not been associated with fetal abnormalities. Nonetheless, the potential risks and benefits of acyclovir should be carefully assessed before the drug is used in pregnancy.

Valacyclovir exhibits three to five times greater bioavailability than acyclovir. The concentration-time curve for valacyclovir, given as 1 g PO three times daily, is similar to that for acyclovir, given as 5 mg/kg IV every 8 h. The safety profiles of valacyclovir and acyclovir are similar, although thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome has been reported in immunocompromised patients who have received high doses of valacyclovir (8 g/d). Valacyclovir is approved for the treatment of herpes zoster, of initial and recurrent episodes of genital HSV infection, and of herpes labialis in immunocompetent adults as well as for suppressive treatment of genital herpes. Although it has not been extensively studied in other clinical settings involving HSV

or VZV infections, many consultants use valacyclovir rather than oral acyclovir in settings where only the latter has been approved because of valacyclovir's superior pharmacokinetics and more convenient dosing schedule.

CIDOFIVIR

Cidofovir is a phosphonate nucleotide analogue of cytosine. Its major use is in CMV infections, but it is active against a broad range of herpesviruses, including HSV, human herpesvirus (HHV) types 6A and 6B, HHV-8, and certain other DNA viruses such as polyomaviruses, papillomaviruses, adenoviruses, and poxviruses, including variola (smallpox) and vaccinia. Cidofovir does not require initial phosphorylation by virus-induced kinases; the drug is phosphorylated by host cell enzymes to cidofovir diphosphate, which is a competitive inhibitor of viral DNA polymerases and, to a lesser extent, of host cell DNA polymerases. Incorporation of cidofovir diphosphate slows or terminates nascent DNA chain elongation. Cidofovir is active against HSV isolates that are resistant to acyclovir because of absent or altered thymidine kinase and against CMV isolates that are resistant to ganciclovir because of UL97 phosphotransferase mutations. CMV isolates resistant to ganciclovir on the basis of UL54 mutations are usually resistant to cidofovir as well. Cidofovir is usually active against foscarnet-resistant CMV, although cross-resistance to foscarnet has been described.

Cidofovir has poor oral availability and is administered intravenously. It is excreted primarily by the kidney and has a plasma half-life of 2.6 h. Cidofovir diphosphate's intracellular half-life of >48 h is the basis for the recommended dosing regimen of 5 mg/kg once a week for the initial 2 weeks and then 5 mg/kg every other week. The major toxic effect of cidofovir is proximal renal tubular injury, as manifested by elevated serum creatinine levels and proteinuria. The risk of nephrotoxicity can be reduced by vigorous saline hydration and by concomitant oral administration of probenecid. Neutropenia, rashes, and gastrointestinal tolerance may also occur.

IV cidofovir has been approved for the treatment of CMV retinitis in AIDS patients who are intolerant of ganciclovir or foscarnet or in whom those drugs have failed. In a controlled study, a maintenance dosage of 5 mg/kg per week administered to AIDS patients reduced the progression of CMV retinitis from that seen at 3 mg/kg. Intravitreal cidofovir has been used to treat CMV retinitis but has been associated with significant toxicity. IV cidofovir has been reported anecdotally to be effective for treatment of acyclovir-resistant mucocutaneous HSV infections. Likewise, topically administered cidofovir is reportedly beneficial against mucocutaneous HSV infections in HIV-infected patients. Anecdotal use of IV cidofovir has been described in disseminated adenoviral infections in immunosuppressed patients and in genitourinary infections with BK virus in renal transplant recipients; however, its efficacy, if any, in these circumstances is not established.

CMX-001 (brincidofovir) is an ester prodrug of cidofovir that can be administered orally and may be less nephrotoxic than IV cidofovir. It is being evaluated for prevention of CMV infection in stem cell transplant recipients and for treatment of BK nephropathy and adenovirus infections.

FOMIVIRSEN

Fomivirsen is the first antisense oligonucleotide approved by the FDA for therapy in humans. This phosphorothioate oligonucleotide, 21 nucleotides in length, inhibits CMV replication through interaction with CMV messenger RNA. Fomivirsen is complementary to messenger transcripts of the major immediate early region 2 (IE2) of CMV, which codes for proteins regulating viral gene expression. In addition to its antisense mechanism of action, fomivirsen may exert activity against CMV through inhibition of viral adsorption to cells as well as direct inhibition of viral replication. Because of its different mechanism of action, fomivirsen is active against CMV isolates that are resistant to nucleoside or nucleotide analogues, such as ganciclovir, foscarnet, or cidofovir.

Fomivirsen has been approved for intravitreal administration in the treatment of CMV retinitis in AIDS patients who have failed to respond to other treatments or cannot tolerate them. Injections of 330 mg for two doses 2 weeks apart, followed by maintenance doses of 330 mg monthly, significantly reduce the rate of progression of CMV retinitis. The major toxicity is ocular inflammation, including vitritis and iritis, which usually responds to topically administered glucocorticoids.

GANCICLOVIR AND VALGANCICLOVIR

An analogue of acyclovir, ganciclovir is active against HSV and VZV and is markedly more active than acyclovir against CMV. Ganciclovir triphosphate inhibits CMV DNA polymerase and can be incorporated into CMV DNA, whose elongation it eventually terminates. In HSV- and VZV-infected cells, ganciclovir is phosphorylated by virus-encoded thymidine kinases; in CMV-infected cells, it is phosphorylated by a viral kinase encoded by the UL97 gene. Ganciclovir triphosphate is present in tenfold higher concentrations in CMV-infected cells than in uninfected cells. Ganciclovir is approved for the treatment of CMV retinitis in immunosuppressed patients and for the prevention of CMV disease in transplant recipients. It is widely used for the treatment of other CMV-associated syndromes, including pneumonia, esophagogastrointestinal infections, hepatitis, and “wasting” illness.

Ganciclovir is available for IV or oral administration. Because its oral bioavailability is low (5–9%), relatively large doses (1 g three times daily) must be administered by this route. Oral ganciclovir has largely been supplanted by valganciclovir, which is the L-valyl ester of ganciclovir. Valganciclovir is well absorbed orally, with a bioavailability of 60%, and is rapidly hydrolyzed to ganciclovir in the intestine and liver. The area under the curve for a 900-mg dose of valganciclovir is equivalent to that for 5 mg/kg of IV ganciclovir, although peak serum concentrations are ~40% lower for valganciclovir. The serum half-life is 3.5 h after IV administration of ganciclovir and 4.0 h after PO administration of valganciclovir. Ganciclovir is excreted primarily by the kidneys in an unmetabolized form, and its dosage should be reduced in cases of renal failure. Ganciclovir therapy at the most commonly used initial IV dosage—i.e., 5 mg/kg every 12 h for 14–21 days—can be changed to valganciclovir (900 mg PO twice daily) when the patient can tolerate oral therapy. The maintenance dose is 5 mg/kg IV daily or five times per week for ganciclovir and 900 mg by mouth once a day for valganciclovir. Dose adjustment in patients with renal dysfunction is required. Intraocular ganciclovir, given by either intravitreal injection or intraocular implantation, has also been used to treat CMV retinitis.

Ganciclovir is effective as prophylaxis against CMV-associated disease in organ and bone marrow transplant recipients. Oral ganciclovir administered prophylactically to AIDS patients with CD4+ T cell counts of <100/μL has provided protection against the development of CMV retinitis. However, the long-term benefits of this approach to prophylaxis in AIDS patients have not been established, and most experts do not recommend the use of oral ganciclovir for this purpose. As already mentioned, valganciclovir has supplanted oral ganciclovir in settings where oral prophylaxis or therapy is considered.

The administration of ganciclovir has been associated with profound bone marrow suppression, particularly neutropenia, which significantly limits the drug's use in many patients. Bone marrow toxicity is potentiated in the setting of renal dysfunction and when other bone marrow suppressants, such as zidovudine or mycophenolate mofetil, are used concomitantly.

Resistance has been noted in CMV isolates obtained after therapy with ganciclovir, especially those from patients with AIDS or from patients receiving prolonged ganciclovir therapy after organ transplantation. Such resistance may develop through a mutation in either the viral UL97 gene or the viral DNA polymerase. Ganciclovir-resistant isolates are usually sensitive to foscarnet (see below) or may be sensitive to cidofovir, depending on the mechanism of resistance (see above).

FAMCICLOVIR AND PENCICLOVIR

Famciclovir is the diacetyl 6-deoxyester of the guanosine analogue penciclovir. This agent is well absorbed orally, has a bioavailability of 77%, and is rapidly converted to penciclovir by deacetylation

and oxidation in the intestine and liver. Penciclovir's spectrum of activity and mechanism of action are similar to those of acyclovir. Thus, penciclovir usually is not active against acyclovir-resistant viruses. However, some acyclovir-resistant viruses with altered thymidine kinase or DNA polymerase substrate specificity may be sensitive to penciclovir. This drug is phosphorylated initially by a virus-encoded thymidine kinase and subsequently by cellular kinases to penciclovir triphosphate, which inhibits HSV-1, HSV-2, VZV, and EBV as well as hepatitis B virus (HBV). The serum half-life of penciclovir is 2 h, but the intracellular half-life of penciclovir triphosphate is 7–20 h—markedly longer than that of acyclovir triphosphate. The latter is the basis for the less frequent (twice-daily) dosing schedule for famciclovir than for acyclovir. Penciclovir is eliminated primarily in the urine by both glomerular filtration and tubular secretion. The usually recommended dosage interval should be adjusted for renal insufficiency.

Clinical trials involving immunocompetent adults with herpes zoster showed that famciclovir was superior to placebo in eliciting the resolution of skin lesions and virus shedding and in shortening the duration of postherpetic neuralgia; moreover, administered at 500 mg every 8 h, famciclovir was at least as effective as acyclovir administered at an oral dose of 800 mg five times daily. Famciclovir was also effective in the treatment of herpes zoster in immunosuppressed patients. Clinical trials have demonstrated its effectiveness in the suppression of genital HSV infections for up to 1 year and in the treatment of initial and recurrent episodes of genital herpes. Famciclovir is effective as therapy for mucocutaneous HSV infections in HIV-infected patients. Application of a 1% penciclovir cream reduces the duration of signs and symptoms of herpes labialis in immunocompetent patients (by 0.5–1 day) and has been approved for that purpose by the FDA. Famciclovir is generally well tolerated, with occasional headache, nausea, and diarrhea reported in frequencies similar to those among placebo recipients. The administration of high doses of famciclovir for 2 years was associated with an increased incidence of mammary adenocarcinomas in female rats, but the clinical significance of this effect is unknown.

FOSCARNET

Foscarnet (phosphonoformic acid) is a pyrophosphate-containing compound that potentially inhibits herpesviruses, including CMV. This drug inhibits DNA polymerases at the pyrophosphate binding site at concentrations that have relatively little effect on cellular polymerases. Foscarnet does not require phosphorylation to exert its antiviral activity and is therefore active against HSV and VZV isolates that are resistant to acyclovir because of deficiencies in thymidine kinase as well as against most ganciclovir-resistant strains of CMV. Foscarnet also inhibits the reverse transcriptase of HIV and is active against HIV *in vivo*.

Foscarnet is poorly soluble and must be administered intravenously via an infusion pump in a dilute solution over 1–2 h. The plasma half-life of foscarnet is 3–5 h and increases with decreasing renal function because the drug is eliminated primarily by the kidneys. It has been estimated that 10–28% of a dose may be deposited in bone, where it can persist for months. The most common initial dosage of foscarnet—60 mg/kg every 8 h for 14–21 days—is followed by a maintenance dose of 90–120 mg/kg once a day.

Foscarnet is approved for the treatment of CMV retinitis in patients with AIDS and of acyclovir-resistant mucocutaneous HSV infections. In a comparative clinical trial, the drug appeared to be about as efficacious as ganciclovir against CMV retinitis but was associated with a longer survival period, possibly because of its activity against HIV. Intraocular foscarnet has been used to treat CMV retinitis. In addition, foscarnet has been employed to treat acyclovir-resistant HSV and VZV infections as well as ganciclovir-resistant CMV infections, although resistance to foscarnet has been reported in CMV isolates obtained during therapy. Foscarnet has also been used to treat HHV-6 infections in immunosuppressed patients.

The major form of toxicity associated with foscarnet is renal impairment. Thus renal function should be monitored closely, particularly during the initial phase of therapy. Because foscarnet binds divalent metal ions, hypocalcemia, hypomagnesemia, hypokalemia,

and hypo- or hyperphosphatemia can develop. Saline hydration and slow infusion appear to protect the patient against nephrotoxicity and electrolyte disturbances. Although hematologic abnormalities have been documented (most commonly anemia), foscarnet is not generally myelosuppressive and can be administered concomitantly with myelosuppressive medications.

TRIFLURIDINE

Trifluridine is a pyrimidine nucleoside active against HSV-1, HSV-2, and CMV. Trifluridine monophosphate irreversibly inhibits thymidylate synthetase, and trifluridine triphosphate inhibits viral and, to a lesser extent, cellular DNA polymerases. Because of systemic toxicity, trifluridine's use is limited to topical therapy. Trifluridine is approved for treatment of HSV keratitis, against which trials have shown that it is more effective than topical idoxuridine but similar in efficacy to topical vidarabine. The drug has benefited some patients with HSV keratitis who have failed to respond to idoxuridine or vidarabine. Topical application of trifluridine to sites of acyclovir-resistant HSV mucocutaneous infection has also been beneficial in some cases.

VIDARABINE

Vidarabine is a purine nucleoside analogue with activity against HSV-1, HSV-2, VZV, and EBV. Vidarabine inhibits viral DNA synthesis through its 5'-triphosphorylated metabolite, although its precise molecular mechanisms of action are not completely understood. IV-administered vidarabine has been shown to be effective in the treatment of herpes simplex encephalitis, mucocutaneous HSV infections, herpes zoster in immunocompromised patients, and neonatal HSV infections. Its use has been supplanted by that of IV acyclovir, which is more effective and easier to administer. Production of the IV preparation has been discontinued by the manufacturer, but vidarabine is available as an ophthalmic ointment, which is effective in the treatment of HSV keratitis.

AGENTS OF INVESTIGATIVE INTEREST

Maribavir is a benzimidazole that inhibits CMV and EBV. This drug inhibits the CMV UL97 kinase and does not require intracellular phosphorylation for its antiviral activity. Its mechanism of action involves blocking viral DNA synthesis and virion egress. Maribavir is orally administered and has been associated with taste disturbance and diarrhea. In phase 3 studies, it was not efficacious in the prevention of CMV infection in recipients of hematopoietic stem cell and adult liver transplants. However, when used at somewhat higher doses, it may be efficacious for the treatment of refractory or resistant CMV infections in transplant recipients.

Letermovir is an investigational drug with activity against CMV. It is a dihydroquinoxaline that acts through inhibition of the viral terminase enzyme complex. This mechanism of action differs from that of ganciclovir, foscarnet, and cidofovir, which inhibit viral DNA polymerase; therefore, letermovir is active against CMV isolates that are resistant to those drugs. It is orally administered and is reportedly well tolerated. Letermovir is being evaluated as prophylaxis against CMV in hematopoietic stem cell recipients.

Inhibition of the helicase-primase heterotrimeric complex of HSV-1 and HSV-2 represents a novel mechanism of action of amenamevir and pritelivir. These drugs are being assessed for prevention and treatment of HSV genital infection. The efficacy of amenamevir, administered as a single oral dose of 1200 mg for recurrent genital herpes, was comparable to that of valacyclovir given for 3 days. Pritelivir has a long half-life (up to 80 h) and was studied in a placebo-controlled trial of suppression of genital HSV infections. Compared with placebo, pritelivir—a loading dose followed by either a daily oral dose of 75 mg for 4 weeks or a weekly dose of 400 mg for 4 weeks—reduced HSV shedding and days of genital lesions. Additional clinical studies of the helicase-primase inhibitors of HSV are planned.

ANTIVIRAL DRUGS ACTIVE AGAINST HEPATITIS VIRUSES

LAMIVUDINE

Lamivudine is a pyrimidine nucleoside analogue that is used primarily in combination therapy against HIV infection ([Chap. 226](#)). Its activity

against HBV is attributable to inhibition of the viral DNA polymerase. This drug has also been approved for the treatment of chronic HBV infection. At doses of 100 mg/d given for 1 year to patients positive for hepatitis B e antigen (HBeAg), lamivudine is well tolerated and results in suppression of HBV DNA levels, normalization of serum aminotransferase levels in 40–75% of patients, and reduction of hepatic inflammation and fibrosis in 50–60% of patients. Loss of HBeAg occurs in 30% of patients. Lamivudine also appears to be useful in the prevention or suppression of HBV infection associated with liver transplantation. Resistance to lamivudine develops in 24% of patients treated for 1 year and is associated with changes in the YMDD motif of HBV DNA polymerase. Because of the frequency of development of resistance, lamivudine has been largely supplanted by less-resistance-prone drugs for the treatment of HBV infection.

ADEFOVIR DIPIVOXIL

Adefovir dipivoxil is the oral prodrug of adefovir, an acyclic nucleotide analogue of adenosine monophosphate that is active against HBV, HIV, HSV, CMV, and poxviruses. It is phosphorylated by cellular kinases to the active triphosphate moiety, which is a competitive inhibitor of HBV DNA polymerase and results in chain termination after incorporation into nascent viral DNA. Adefovir is administered orally and is eliminated primarily by the kidneys, with a plasma half-life of 5–7.5 h. In clinical studies, therapy with adefovir at a dose of 10 mg/d for 48 weeks resulted in normalization of serum alanine aminotransferase (ALT) levels in 48–72% of patients and improved liver histology in 53–64%; it also resulted in a 3.5- to 3.9- \log_{10} reduction in the number of HBV DNA copies per milliliter of plasma. Adefovir was effective in treatment-naïve patients as well as in those infected with lamivudine-resistant HBV. Resistance to adefovir appears to develop less readily than that to lamivudine, but adefovir resistance rates of 15–18% have been reported after 192 weeks of treatment and may reach 30% after 5 years. This agent is generally well tolerated. Significant nephrotoxicity attributable to adefovir is uncommon at the dose used in the treatment of HBV infections (10 mg/d) but is a treatment-limiting adverse effect at the higher doses used in therapy for HIV infections (30–120 mg/d). In any case, renal function should be monitored in patients taking adefovir, even at the lower dose. Adefovir is approved only for treatment of chronic HBV infection.

TENOFOVIR DISOPROXIL FUMARATE

Tenofovir disoproxil fumarate is a prodrug of tenofovir, a nucleotide analogue of adenosine monophosphate with activity against both retroviruses and hepadnaviruses. In both immunocompetent and immunocompromised patients (including those co-infected with HIV and HBV), tenofovir given at a dose of 300 mg/d for 48 weeks reduced HBV replication by 4.6–6 \log_{10} , normalized ALT levels in 68–76% of patients, and improved liver histopathology in 72–74% of patients. Resistance develops uncommonly during ≥ 2 years of therapy, and tenofovir is active against lamivudine-resistant HBV. The safety profile of tenofovir is similar to that of adefovir, but nephrotoxicity has not been encountered at the dose used for HBV therapy. Tenofovir is approved for the treatment of HIV and chronic HBV infections. [For a more detailed discussion of tenofovir, see Chap. 226.](#)

ENTECAVIR

Entecavir is a cyclopentyl 2'-deoxyguanosine analogue that inhibits HBV through interaction of entecavir triphosphate with several HBV DNA polymerase functions. At a dose of 0.5 mg/d given for 48 weeks, entecavir reduced HBV DNA copies by 5.0–6.9 \log_{10} , normalized serum aminotransferase levels in 68–78% of patients, and improved histopathology in 70–72% of patients. Entecavir inhibits lamivudine-resistant viruses that have M550I or M550V/L526M mutations but only at serum concentrations 20- or 30-fold higher than those obtained with the 0.5-mg/d dose. Thus, higher doses of entecavir (1 mg/d) are recommended for the treatment of patients infected with lamivudine-resistant HBV. Development of resistance to entecavir is uncommon in treatment-naïve patients but does occur at unacceptably high rates

215e-10 (43% after 4 years) in patients previously infected with lamivudine-resistant virus. Entecavir-resistant strains appear to be sensitive to adefovir and tenofovir.

Entecavir is highly bioavailable but should be taken on an empty stomach because food interferes with its absorption. The drug is eliminated primarily in unchanged form by the kidneys, and its dosage should be adjusted for patients with Cr_{Cl} values of <50 mL/min. Overall, entecavir is well tolerated, with a safety profile similar to that of lamivudine. As with other anti-HBV treatments, exacerbation of hepatitis may occur when entecavir therapy is stopped. Entecavir is approved for treatment of chronic hepatitis B, including infection with lamivudine-resistant viruses, in adults. Entecavir has some activity against HIV-1 (median effective concentration, 0.026 to >10 μ M) but should not be used as monotherapy in HIV-positive patients because of the potential for development of HIV resistance due to the M184V mutation.

TELBIVUDINE

Telbivudine is a β -L enantiomer of thymidine and is a potent, selective inhibitor of HBV. Its active form is telbivudine triphosphate, which inhibits HBV DNA polymerase and causes chain termination but has little or no activity against human DNA polymerase. Administration of telbivudine at an oral dose of 600 mg/d for 52 weeks to patients with chronic hepatitis B resulted in reduction of HBV DNA by 5.2–6.4 \log_{10} copies/mL along with normalization of ALT levels in 74–77% of recipients and improved histopathology in 65–67% of patients. Telbivudine-resistant HBV is generally cross-resistant with lamivudine-resistant virus but is usually susceptible to adefovir. After 2 years of therapy, resistance to telbivudine was noted in isolates from 22% of HBeAg-positive patients and in those from 9% of HBeAg-negative patients.

Orally administered telbivudine is rapidly absorbed; because it is eliminated primarily by the kidneys, its dosage should be reduced in patients with a Cr_{Cl} value of <50 mL/min. Telbivudine is generally well tolerated, but increases in serum levels of creatinine kinases as well as fatigue and myalgias have been observed. As with other anti-HBV drugs, hepatitis may be exacerbated in patients who discontinue telbivudine therapy. Telbivudine has been approved for the treatment of adults with chronic hepatitis B who have evidence of viral replication and either persistently elevated serum aminotransferase levels or histopathologically active disease, but it has not been widely used because of the frequency of development of resistance, as noted above.

INTERFERONS

IFNs are cytokines that exhibit a broad spectrum of antiviral activities as well as immunomodulating and antiproliferative properties. IFNs are not available for oral administration but must be given IM, SC, or IV. Early studies with human leukocyte IFN demonstrated an effect in the prophylaxis of experimentally induced rhinovirus infections in humans and in the treatment of VZV infections in immunosuppressed patients. DNA recombinant technology has made available highly purified α , β , γ , and λ IFNs that have been evaluated in a variety of viral infections. Results of such trials have confirmed the effectiveness of intranasally administered IFN in the prophylaxis of rhinovirus infections, although its use has been associated with nasal mucosal irritation. Studies have also demonstrated a beneficial effect of intralesionally or systemically administered IFNs on genital warts. The effect of systemic administration consists primarily of a reduction in the size of the warts, and this mode of therapy may be useful in persons who have numerous warts that cannot easily be treated by individual intralesional injections. However, lesions frequently recur after either intralesional or systemic IFN therapy is discontinued.

IFNs have undergone extensive study in the treatment of chronic HBV infection. The administration of standard IFN- α 2b (5 million units daily or 10 million units three times a week for 16–24 weeks) to patients with stable chronic HBV infection resulted in loss of markers of HBV replication, such as HBeAg and HBV DNA, in 33–37% of cases; 8% of patients also became negative for hepatitis B surface antigen. In most patients who lose HBeAg and HBV DNA markers, serum aminotransferases return to normal levels, and both short- and long-term improvements in liver histopathology have been described.

Predictors of a favorable response to standard IFN therapy include low pretherapy levels of HBV DNA, high pretherapy serum levels of ALT, a short duration of chronic HBV infection, and active inflammation in liver histopathology. Poor responses are seen in immunosuppressed patients, including those with HIV infection.

In pegylated IFNs, IFN alphas are linked to polyethylene glycol. This linkage results in slower absorption, decreased clearance, and more sustained serum concentrations, thereby permitting a more convenient, once-weekly dosing schedule; in many instances, pegylated IFN has supplanted standard IFN. After 48 weeks of treatment with 180 μ g of pegylated IFN- α 2a, HBV DNA was reduced by 4.1–4.5 \log_{10} copies/mL, with normalization of serum ALT levels in 39% of patients and improved histology in 38%. Response rates were somewhat higher when lamivudine was administered with pegylated IFN- α 2a. Adverse effects of IFN are common and include fever, chills, myalgia, fatigue, neurotoxicity (manifested primarily as somnolence, depression, anxiety, and confusion), and leukopenia. Autoantibodies (e.g., antithyroid antibodies) can also develop. IFN- α 2b and pegylated IFN- α 2a are approved for the treatment of patients with chronic hepatitis B. Data supporting the therapeutic efficacy of pegylated interferon- α 2b in HBV infection have been published; the drug has not been approved for this indication in the United States but has been approved for treatment of chronic HBV infection in other countries.

Several IFN preparations, including IFN- α 2a, IFN- α 2b, IFN- α 1, and IFN- α m1 (lymphoblastoid), have been studied as therapy for chronic HCV infections. A variety of monotherapy regimens have been studied, of which the most common for standard IFN is IFN- α 2b or - α 2a at 3 million units three times per week for 12–18 months. The addition of oral ribavirin to IFN- α 2b—either as initial therapy or after failure of IFN therapy alone—results in significantly higher rates of sustained virologic and/or serum ALT responses (40–50%) than are obtained with monotherapy. Comparative studies indicate that pegylated IFN- α 2b or - α 2a therapy is more effective than standard IFN treatment against chronic HCV infection. The combination of SC pegylated IFN and oral ribavirin results in sustained virologic responses (SVRs) in 42–51% of patients with HCV genotype 1 infection and in 76–82% of patients with genotype 2 or 3 infection. Ribavirin appears to have a small antiviral effect in HCV infection but may also be working through an immunomodulatory effect in combination with IFN. Optimal results with ribavirin appear to be associated with weight-based dosing. Prognostic factors for a favorable response include an age of <40 years, a short duration of infection, low levels of HCV RNA, a lesser degree of liver histopathology, and infection with HCV genotypes other than 1. IFN- α 1, a synthetic “consensus” α interferon, appears to produce response rates similar to those elicited by standard IFN- α 2a or - α 2b alone. In 2014, the approval of a polymerase inhibitor, sofosbuvir, and a second-generation protease inhibitor, simeprevir, led to revised recommendations for treatment of hepatitis C with triple combinations of pegylated IFN, ribavirin, and one of these two drugs, depending on the viral genotype (see below and Table 215e-1).

IFN- α and pegylated IFN- α are active against hepatitis D, but high doses are required (9 million units three times per week for 48 weeks). IFN- α elicited an SVR in 25–30% of patients, whereas pegylated IFN- α had a variable effect, evoking an SVR in 17–43% of patients. However, long-term biochemical and histologic improvements have been seen, even in the absence of sustained inhibition of viral replication.

POLYMERASE INHIBITORS

SOFSBUVIR

Sofosbuvir is the prodrug of a uridine nucleoside inhibitor of the HCV RNA NS5B polymerase. Its metabolism to the active uridine nucleoside triphosphate results in chain termination. Sofosbuvir is active against all HCV genotypes (1–6) and has a median effective concentration (EC_{50}) of 0.7–2.6 μ M against NS5B. Resistance to sofosbuvir is conferred by an S282T substitution in NS5B, but clinically expressed resistance to treatment has only rarely been encountered in patients who receive sofosbuvir.

Sofosbuvir is administered orally and is unaffected by food. After oral administration, plasma concentrations of sofosbuvir and of its active metabolite peak in 0.5–2 h and 2–4 h, respectively. Approximately 61–65% of sofosbuvir is bound in plasma proteins, but very little of the active metabolite is bound. Both sofosbuvir and its active metabolite are cleared renally, with $t_{1/2}$ values of 0.4 h and 27 h, respectively. Sofosbuvir is relatively free from clinically significant drug interactions, although P-glycoprotein inducers can reduce sofosbuvir concentrations.

Sofosbuvir is generally well tolerated and has not been associated with significant toxicity. The most common side effects in recipients of sofosbuvir have been attributable to concomitant administration of IFN and ribavirin in combination clinical trials (see below).

Sofosbuvir has been studied in a variety of controlled and open-label clinical trials. In late 2013, the results of these trials led to its recommendation—in triple combination with pegylated IFN and ribavirin—as first-line treatment for chronic hepatitis due to HCV genotypes 1, 4, 5, and 6, in which SVRs among treatment-naïve patients were 89–97%. For HCV genotypes 2 and 3, IFN-free regimens consisting of sofosbuvir and ribavirin have been recommended, with SVRs among treatment-naïve patients of 93% for genotype 2 and 61% for genotype 3.

PROTEASE INHIBITORS

BOCEPREVIR, TELAPREVIR

This drug class is specifically designed to inhibit the 3/4A (NS3/4A) HCV protease. These agents resemble the HCV polypeptide and, when processed by the viral protease, form a covalent bond with the catalytic NS3 serine residues, block further activity, and prevent proteolytic cleavage of the HCV polyprotein into NS4A, NS4B, NS5A, and NS5B proteins. Boceprevir and telaprevir are linear ketoamide compounds that are active against HCV genotype 1 (1b > 1a) and much less so against genotypes 2 and 3. These first-generation protease inhibitors received approval for combination therapy (with IFN and ribavirin) for genotype 1 infection. Neither boceprevir nor telaprevir is now recommended for the treatment of hepatitis C. These drugs have been supplanted by sofosbuvir and by simeprevir, a second-generation protease inhibitor with improved pharmacokinetic properties, fewer drug–drug interactions, and less overall toxicity (see below).

SIMEPREVIR

Simeprevir is a second-generation NS3/4A protease inhibitor with antiviral activity against genotype 1 (1b > 1a); the EC_{50} is 9.4 nM in an HCV genotype 1b replicon. The NS3 polymorphism Q80K, which is present in approximately one-third of patients carrying HCV genotype 1b, increases the EC_{50} by elevenfold and results in clinical resistance to

simeprevir. Thus, testing for Q80K should be carried out if treatment with simeprevir is being considered. Cross-resistance occurs between simeprevir and the first-generation protease inhibitors boceprevir and telaprevir.

Simeprevir is orally administered as a 150-mg capsule, and its bioavailability is increased by administration with food. The serum concentration peaks 4–6 h after oral administration. The drug's elimination half-life is 10–13 h in healthy individuals and 41 h in patients with hepatitis C. Simeprevir is nearly entirely bound by plasma proteins and cleared by biliary excretion. Because there is no renal excretion, dose adjustments are not required in the presence of renal dysfunction. Simeprevir is metabolized by hepatic CYP3A and therefore should not be used in patients with decompensated liver function.

Because of its metabolism by cytochrome P450 3A (CYP3A), simeprevir interacts with drugs that induce or inhibit CYP3A, and these interactions may concomitantly increase or reduce plasma concentrations of simeprevir. Administration of simeprevir may also increase plasma concentrations of drugs that are substrates for hepatic organic anion-transporting polypeptide 1B1 or 1B3 or for P glycoprotein transporters.

Toxicity observed during clinical trials with simeprevir included photosensitivity (usually mild or moderate) in 28% of recipients and reversible hyperbilirubinemia (both conjugated and unconjugated), which was generally mild to moderate. Most of the other adverse effects that were seen in clinical trials with simeprevir were attributable to concomitant administration of IFN and ribavirin.

Simeprevir has been recommended as a component of alternative treatment—in combination with pegylated IFN and ribavirin—of chronic infection with HCV genotypes 1 and 4. Daily simeprevir, daily ribavirin, and weekly pegylated IFN for 12 weeks followed by another 12 weeks of pegylated IFN and ribavirin resulted in an SVR of 80% in the absence of the Q80K variant. In general, simeprevir-based triple therapy appeared to be 10% less likely to yield an SVR than sofosbuvir-based therapy and more likely to cause adverse effects. However, for prior nonresponders or partial responders to pegylated IFN, the IFN-free regimen of simeprevir, sofosbuvir, and ribavirin shows promise.

INVESTIGATIVE AGENTS OF INTEREST

Next-generation direct-acting antivirals against HCV inhibitors are under active development. These agents include second-generation inhibitors of NS3/4, NS5B polymerase inhibitors, and inhibitors of NS5A (a membrane-associated phosphoprotein that is part of the HCV RNA replication complex). These investigational agents are making progress toward IFN-free regimens, shorter courses of therapy, improved tolerability, and reduction of resistance. For updated information, readers should consult <http://www.hcvguidelines.org/>.

SECTION 12 INFECTIONS DUE TO DNA VIRUSES

216 Herpes Simplex Virus Infections

Lawrence Corey

DEFINITION

Herpes simplex viruses (HSV-1, HSV-2; *Herpesvirus hominis*) produce a variety of infections involving mucocutaneous surfaces, the central nervous system (CNS), and—on occasion—visceral organs. Prompt recognition and treatment reduce the morbidity and mortality rates associated with HSV infections.

ETIOLOGIC AGENT



The genome of HSV is a linear, double-strand DNA molecule (molecular weight, $\sim 100 \times 10^6$) that encodes >90 transcription units with 84 identified proteins. The genomic structures of the two HSV subtypes are similar. The overall genomic sequence homology between HSV-1 and HSV-2 is $\sim 50\%$, whereas the proteome homology is $>80\%$. The homologous sequences are distributed over the entire genome map, and most of the polypeptides specified by one viral type are antigenically related to polypeptides of the other viral type. Many type-specific regions unique to HSV-1 and HSV-2 proteins do exist, however, and a number of them appear to be important in host immunity. These type-specific regions have been used to develop serologic assays that distinguish between the two viral subtypes. Either restriction endonuclease analysis or sequencing of viral DNA can be used to distinguish between the two subtypes and among strains of each subtype. The variability of nucleotide sequences from clinical strains of HSV-1 and HSV-2 is such that HSV isolates obtained from two individuals can be differentiated by restriction enzyme patterns or genomic sequences. Moreover, epidemiologically related sources, such as sexual partners, mother-infant pairs, or persons involved in a common-source outbreak, can be inferred from such patterns.

The viral genome is packaged in a regular icosahedral protein shell (capsid) composed of 162 capsomeres (see Fig. 214e-1). The outer covering of the virus is a lipid-containing membrane (envelope) acquired as the DNA-containing capsid buds through the inner nuclear membrane of the host cell. Between the capsid and lipid bilayer of the envelope is the tegument. Viral replication has both nuclear and cytoplasmic phases. Initial attachment to the cell membrane involves interactions of viral glycoproteins C and B with several cellular heparan sulfate-like surface receptors. Subsequently, viral glycoprotein D binds to cellular co-receptors that belong to the tumor necrosis factor receptor family of proteins, the immunoglobulin superfamily (nectin family), or both. The ubiquity of these receptors contributes to the wide host range of herpesviruses. HSV replication is highly regulated. After fusion and entry, the nucleocapsid enters the cytoplasm and several viral proteins are released from the virion. Some of these viral proteins shut off host protein synthesis (by increasing cellular RNA degradation), whereas others “turn on” the transcription of early genes of HSV replication. These early gene products, designated α genes, are required for synthesis of the subsequent polypeptide group, the β polypeptides, many of which are regulatory proteins and enzymes required for DNA replication. Most current antiviral drugs interfere with β proteins, such as viral DNA polymerase. The third (γ) class of HSV genes requires viral DNA replication for expression and encodes most structural proteins specified by the virus.

After viral genome replication and structural protein synthesis, nucleocapsids are assembled in the cell's nucleus. Envelopment occurs as the nucleocapsids bud through the inner nuclear membrane into the perinuclear space. In some cells, viral replication in the nucleus forms two types of inclusion bodies: type A basophilic Feulgen-positive bodies that contain viral DNA and eosinophilic inclusion bodies that are devoid of viral nucleic acid or protein and represent a “scar” of viral infection.

Enveloped virions are then transported via the endoplasmic reticulum and the Golgi apparatus to the cell surface.

Viral genomes are maintained by some neuronal cells in a repressed state called *latency*. Latency, which is associated with transcription of only a limited number of virus-encoded RNAs, accounts for the presence of viral DNA and RNA in neural tissue at times when infectious virus cannot be isolated. Maintenance and growth of neural cells from latently infected ganglia in tissue culture result in production of infectious virions (*explantation*) and in subsequent permissive infection of susceptible cells (*co-cultivation*). Activation of the viral genome may then occur, resulting in *reactivation*—the normal pattern of regulated viral gene expression and replication and HSV release. The release of virions from the neuron follows a complex process of anterograde transport down the length of neuronal axons. In experimental animals, ultraviolet light, systemic and local immunosuppression, and trauma to the skin or ganglia are associated with reactivation.

Three noncoding RNA latency-associated transcripts (LATs) are found in the nuclei of latently infected neurons. Deletion mutants of the LAT region exhibit reduced efficiency in their later reactivation. Substitution of HSV-1 LATs for HSV-2 LATs induces an HSV-1 reactivation pattern. These data indicate that LATs apparently maintain—rather than establish—latency. HSV-1 LATs promote the survival of acutely infected neurons, perhaps by inhibiting apoptotic pathways. LAT transcript abundance and low genome-copy number correlate with subnuclear positioning of HSV genomes around the centromere. Indeed, chromatization of HSV DNA appears to play a vital role in silencing expression of lytic replication genes. Highly expressed during latency, LAT-derived micro-RNA appears to silence expression of the key neurovirulence factor infected-cell protein 34.5 (ICP34.5) and to bind in an antisense configuration to the immediate-early protein ICP0 messenger RNA to prevent expression, which is vital to HSV reactivation. Although certain viral transcripts are known to be necessary for reactivation from latency, the molecular mechanisms of HSV latency are not fully understood, and strategies to interrupt or maintain latency in neurons are in developmental stages.

While latency is the predominant state of virus on a per-neuron basis, the high frequency of oral and genital tract reactivation for HSV-1 and HSV-2 suggests that the viruses are rarely quiescent within the entire biomass of ganglionic tissue. Recent data indicate that HSV-2 antigen is often shed: most persons infected with HSV-2 have frequent subclinical bursts of reactivation lasting 2–4 h, and the host mucosal immune system can contain viral reactivation in the mucosa before the development of clinical reactivation. Supporting this clinical observation, recent work using microdissection plus real-time polymerase chain reaction (PCR) of individual neurons from cadaveric trigeminal ganglia explants revealed that many more neurons (2–10%) harbor HSV than would be predicted by *in situ* hybridization studies for LATs. Viral copy number is highly variable between neurons, with extremely high levels in certain neurons, and HSV DNA copy numbers are similar in LAT-positive and LAT-negative neurons; these findings add to the uncertainty about the role that LATs play in preventing reactivation.

PATHOGENESIS

Exposure to HSV at mucosal surfaces or abraded skin sites permits entry of the virus into cells of the epidermis and dermis and initiation of viral replication therein. HSV infections are usually acquired subclinically. Whether clinical or subclinical, HSV acquisition is associated with sufficient viral replication to permit infection of either sensory or autonomic nerve endings. On entry into the neuronal cell, the virus—or, more likely, the nucleocapsid—is transported intra-axonally to the nerve cell bodies in ganglia. In humans, the transit interval of spread to the ganglia after virus inoculation into peripheral tissue is unknown. During the initial phase of infection, viral replication occurs in ganglia

1176 and contiguous neural tissue. Virus then spreads to other mucocutaneous surfaces through centrifugal migration of infectious virions via peripheral sensory nerves. This mode of spread helps explain the large surface area involved, the high frequency of new lesions distant from the initial crop of vesicles that is characteristic in patients with primary genital or oral-labial HSV infection, and the ability to recover virus from neural tissue distant from neurons innervating the inoculation site. Contiguous spread of locally inoculated virus also may take place and allow further mucosal extension of disease. Recent studies have demonstrated HSV viremia—another mechanism for extension of infection throughout the body—in ~30–40% of persons with primary HSV-2 infection. Latent infection with both viral subtypes in both sensory and autonomic ganglia has been demonstrated. For HSV-1 infection, trigeminal ganglia are most commonly infected, although extension to the inferior and superior cervical ganglia also occurs. With genital infection, sacral nerve root ganglia (S2–S5) are most commonly affected.

After resolution of primary disease, infectious HSV can no longer be cultured from the ganglia; however, latent infection, as defined by the presence of viral DNA, persists in 2–11% of ganglionic cells in the anatomic region of the initial infection. The mechanism of reactivation from latency is unknown. Increasingly, studies indicate that host T cell responses at the ganglionic and peripheral mucosal levels influence the frequency and severity of HSV reactivation. HSV-specific T cells have been recovered from peripheral-nerve root ganglia. Many of these resident CD8+ T cells are juxtaposed with latently HSV-1-infected neurons in the trigeminal ganglia and can block reactivation with both interferon (IFN) γ release and granzyme B degradation of the immediate-early protein ICP4. In addition, there appears to be a latent viral load in the ganglia that correlates positively with the number of neurons infected and the rate of reactivation but inversely with the number of CD8+ cells present. It is not known whether reactivating stimuli transiently suppress these immune cells, independently upregulate transcription of lytic genes, or both. Moreover, host containment in the mucosa has been demonstrated. Once virus reaches the dermal-epidermal junction, there are three possible outcomes: rapid host containment of infection near the site of reactivation; spread of virus into the epidermis, with a micro-ulceration associated with low-titer subclinical shedding; and subsequent rapid (within hours) containment of virus with widespread replication and necrosis of epithelial cells and subsequent clinical recurrence (the latter defined clinically by a skin blister and ulceration). Histologically, herpetic lesions involve a thin-walled vesicle or ulceration in the basal region, multinucleated cells that may include intranuclear inclusions, necrosis, and an acute inflammatory infection. Re-epithelialization occurs once viral replication is restricted, almost always in the absence of a scar.

Analysis of the DNA from sequential isolates of HSV or from isolates from multiple infected ganglia in any one individual has revealed similar, if not identical, restriction endonuclease or DNA sequence patterns in most persons. As more sensitive genomic technologies are developed, evidence of multiple strains of the same subtype is increasingly being reported. For example, infection of individual neurons with multiple strains of drug-susceptible and drug-resistant virus in severely immunosuppressed patients indicates that ganglia can be reseeded during chronic infection. Because exposure to mucosal shedding is relatively common during a person's lifetime, current data suggest that exogenous infection with different strains of the same subtype, while possible, is uncommon.

IMMUNITY


Host responses influence the acquisition of HSV disease, the severity of infection, resistance to the development of latency, the maintenance of latency, and the frequency of recurrences. Both antibody-mediated and cell-mediated reactions are clinically important. Immunocompromised patients with defects in cell-mediated immunity experience more severe and more extensive HSV infections than those with deficits in humoral immunity, such as agammaglobulinemia. Experimental ablation of lymphocytes indicates that T cells play a major role in preventing lethal disseminated disease, although antibodies help reduce titers

of virus in neural tissue. Some clinical manifestations of HSV appear to be related to the host immune response (e.g., stromal opacities associated with recurrent herpetic keratitis). The surface viral glycoproteins have been shown to be targets of antibodies that mediate neutralization and immune-mediated cytolysis (antibody-dependent cell-mediated cytotoxicity). Monoclonal antibodies specific for each of the known viral glycoproteins have, in experimental infections, conferred protection against subsequent neurologic disease or ganglionic latency. In humans, however, subunit glycoprotein vaccines have been largely ineffective in reducing acquisition of infection. Multiple cell populations, including natural killer cells, macrophages, and a variety of T lymphocytes, play a role in host defenses against HSV infections, as do lymphokines generated by T lymphocytes. In animals, passive transfer of primed lymphocytes confers protection from subsequent HSV challenge. Maximal protection usually requires the activation of multiple T cell subpopulations, including cytotoxic T cells and T cells responsible for delayed hypersensitivity. The latter may confer protection by the antigen-stimulated release of lymphokines (e.g., IFNs), which in turn have a direct antiviral effect and both activate and enhance a variety of specific and nonspecific effector cells. The HSV virion contains a variety of genes that are directed at the inhibition of host responses. These include gene no. 12 (*US-12*), which can bind to the cellular transporter-activating protein TAP-1 and reduce the ability of this protein to bind HSV peptides to human leukocyte antigen (HLA) class I, thereby reducing recognition of viral proteins by cytotoxic T cells of the host. This effect can be overcome by the addition of IFN- γ , but this reversal requires 24–48 h; thus, the virus has time to replicate and invade other host cells. Entry of infectious HSV-1 and HSV-2 inhibits several signaling pathways of both CD4+ and CD8+ T cells, leading to their functional impairment in killing and influencing the spectrum of their cytokine secretion.

Increasing evidence suggests that HSV-specific CD8+ T cell responses are critical for clearance of virus from lesions. Immunosuppressed patients with frequent and prolonged HSV lesions have fewer functional CD8+ T cells directed at HSV. HSV-specific CD8+ T cells have been shown to persist in the genital skin at the dermal-epidermal junction contiguous to neuronal endings. Even during clinical quiescence, these CD8+ T cells make both antiviral and cytotoxic proteins indicative of immune surveillance. These resident memory CD8+ T cells appear to be “first responders” capable of controlling viral reactivation at the site of viral release into the dermis. This rapid “on and off” interplay between the virus and host helps explain the variability in clinical disease severity between episodes in any single individual. Differences of 30–60 min in host responses can result in 100- to 1000-fold differences in viral levels and can determine whether an episode of disease is subclinical or clinical.

There is a strong association between the magnitude of the CD8+ T lymphocyte response and the clearance of virus from genital lesions. The location, effectiveness, and longevity of the T lymphocytes (and perhaps of other immune effector cells) may be important in the expression of disease and the likelihood of transmission over time.

EPIDEMIOLOGY

 Seroepidemiologic studies have documented HSV infections worldwide. The past 15 years have shown that the prevalence of HSV-2 is even higher in the developing than in the developed world. In sub-Saharan Africa, HSV-2 seroprevalence among pregnant women may approach 60%, and annual acquisition rates among teenage girls may verge on 20%. The global incidence has been estimated at ~23.6 million infections per year. As in the developed world, the rate of HSV-2 coital acquisition as well as the serologic prevalence is higher among women than among men. Most of this HSV-2 acquisition is preceded by acquisition of HSV-1; the frequency of genital HSV-1 in the developing world is low at present.

Infection with HSV-1 is acquired more frequently and earlier than infection with HSV-2. More than 90% of adults have antibodies to HSV-1 by the fifth decade of life. In populations of low socioeconomic status, most persons acquire HSV-1 infection before the third decade of life. Antibodies to HSV-2 are not detected routinely until puberty.

Antibody prevalence rates correlate with past sexual activity and vary greatly among different population groups. There is evidence that the prevalence of HSV-2 has decreased slightly over the past decade in the United States. Serosurveys indicate that 15–20% of the U.S. population has antibodies to HSV-2. In most routine obstetric and family planning clinics, 25% of women have HSV-2 antibodies, although only 10% of those who are seropositive for HSV-2 report a history of genital lesions. As many as 50% of heterosexual adults attending sexually transmitted disease clinics have antibodies to HSV-2.

Many studies continue to show that both incident and—more important—prevalent HSV-2 infection enhances the acquisition rate of HIV-1. More specifically, HSV-2 infection is associated with a two- to fourfold increase in HIV-1 acquisition. This association has been amply demonstrated in heterosexual men and women in both the developed and the developing worlds. Epidemiologically, regions of the world with high HSV-2 prevalence and selected populations within such regions have a higher population-based incidence of HIV-1. One study indicated that approximately one-quarter of HIV infections in the high-prevalence city of Kisumu, Kenya, were directly attributable to HSV-2.

In addition, HSV-2 facilitates the spread of HIV into low-risk populations on a per-coital basis, and prevalent HSV-2 appears to increase the risk of HIV infection by seven- to ninefold. Mathematical models suggest that ~33–50% of HIV-1 infections may be attributable to HSV-2 both in men who have sex with men (MSM) and in sub-Saharan Africa. In addition, HSV-2 is more frequently reactivated in and transmitted by persons co-infected with HIV-1 as opposed to persons not co-infected. Thus, most areas of the world with a high HIV-1 prevalence also have a high HSV-2 prevalence. A wide variety of serologic surveys have indicated a similar or even higher seroprevalence of HSV-2 in most parts of Central America, South America, and Africa. In Africa, HSV-2 seroprevalence has ranged from 40% to 70% in obstetric and other sexually experienced populations. Antibody prevalence rates average ~5–10% higher among women than among men.

Several studies suggest that many cases of “asymptomatic” genital HSV-2 infection are, in fact, simply unrecognized or confined to anatomic regions of the genital tract that are not easily visualized. Asymptomatic seropositive persons shed virus on mucosal surfaces almost as frequently as do those with symptomatic disease. This large reservoir of unidentified carriers of HSV-2 and the frequent asymptomatic reactivation of the virus from the genital tract have fostered the continued spread of genital herpes throughout the world. HSV-2 infection is an independent risk factor for the acquisition and transmission of infection with HIV-1. Among co-infected persons, HIV-1 virions can be shed from herpetic lesions of the genital region. This shedding may facilitate the spread of HIV through sexual contact. HSV-2 reactivation is associated with a localized persistent inflammatory response consisting of high concentrations of CCR5-enriched CD4+ T cells as well as inflammatory dendritic cells in the submucosa of the genital skin. These cells can support HIV infection and replication and hence are likely to account for the almost threefold increase in HIV acquisition among persons with genital herpes. Unfortunately, antiviral therapy does not reduce this subclinical postreactivation inflammation, probably because of the inability of current antiviral agents to prevent the release of small amounts of HSV antigen into the genital mucosa.

HSV infections occur throughout the year. Transmission can result from contact with persons who have active ulcerative lesions or with persons who have no clinical manifestations of infection but who are shedding HSV from mucocutaneous surfaces. HSV reactivation on genital skin and mucosal surfaces is common. The frequency of sampling influences the frequency of detection. Recent studies indicate that most HSV-1 and HSV-2 episodes last <4–6 h; thus, replication of virus and clearance by the host are rapid. Even with once-daily sampling, HSV DNA can be detected on 20–30% of days by PCR. Corresponding figures for HSV-1 in oral secretions are similar. Rates of shedding are highest during the initial years after acquisition, with viral shedding occurring on as many as 30–50% of days during this period. Immunosuppressed patients shed HSV from mucosal sites at an even

higher frequency (20–80% of days). These high rates of mucocutaneous reactivation suggest that exposure to HSV from sexual or other close contact (kissing, sharing of glasses or silverware) is common and help explain the continuing spread and high seroprevalence of HSV infections worldwide. Reactivation rates vary widely among individuals. Among HIV-positive patients, a low CD4+ T cell count and a high HIV-1 load are associated with increased rates of HSV reactivation. Daily antiviral chemotherapy for HSV-2 infection can reduce shedding rates but does not eliminate shedding, as measured by PCR or culture.

CLINICAL SPECTRUM

HSV has been isolated from nearly all visceral and mucocutaneous sites. The clinical manifestations and course of HSV infection depend on the anatomic site involved, the age and immune status of the host, and the antigenic type of the virus. Primary HSV infections (i.e., first infections with either HSV-1 or HSV-2 in which the host lacks HSV antibodies in acute-phase serum) are frequently accompanied by systemic signs and symptoms. Compared with recurrent episodes, primary infections, which involve both mucosal and extramucosal sites, are characterized by a longer duration of symptoms and virus isolation from lesions. The incubation period ranges from 1 to 26 days (median, 6–8 days). Both viral subtypes can cause genital and oral-facial infections, and the infections caused by the two subtypes are clinically indistinguishable. However, the frequency of reactivation of infection is influenced by anatomic site and virus type. Genital HSV-2 infection is twice as likely to reactivate and recurs 8–10 times more frequently than genital HSV-1 infection. Conversely, oral-labial HSV-1 infection recurs more frequently than oral-labial HSV-2 infection. Asymptomatic shedding rates follow the same pattern.

Oral-Facial Infections Gingivostomatitis and pharyngitis are the most common clinical manifestations of first-episode HSV-1 infection, whereas recurrent herpes labialis is the most common clinical manifestation of reactivation HSV-1 infection. HSV pharyngitis and gingivostomatitis usually result from primary infection and are most common among children and young adults. Clinical symptoms and signs, which include fever, malaise, myalgias, inability to eat, irritability, and cervical adenopathy, may last 3–14 days. Lesions may involve the hard and soft palate, gingiva, tongue, lip, and facial area. HSV-1 or HSV-2 infection of the pharynx usually results in exudative or ulcerative lesions of the posterior pharynx and/or tonsillar pillars. Lesions of the tongue, buccal mucosa, or gingiva may occur later in the course in one-third of cases. Fever lasting 2–7 days and cervical adenopathy are common. It can be difficult to differentiate HSV pharyngitis clinically from bacterial pharyngitis, *Mycoplasma pneumoniae* infections, and pharyngeal ulcerations of noninfectious etiologies (e.g., Stevens-Johnson syndrome). No substantial evidence suggests that reactivation of oral-labial HSV infection is associated with symptomatic recurrent pharyngitis.

Reactivation of HSV from the trigeminal ganglia may be associated with asymptomatic virus excretion in the saliva, development of intraoral mucosal ulcerations, or herpetic ulcerations on the vermilion border of the lip or external facial skin. About 50–70% of seropositive patients undergoing trigeminal nerve-root decompression and 10–15% of those undergoing dental extraction develop oral-labial HSV infection a median of 3 days after these procedures. Clinical differentiation of intraoral mucosal ulcerations due to HSV from aphthous, traumatic, or drug-induced ulcerations is difficult.

In immunosuppressed patients, HSV infection may extend into mucosal and deep cutaneous layers. Friability, necrosis, bleeding, severe pain, and inability to eat or drink may result. The lesions of HSV mucositis are clinically similar to mucosal lesions caused by cytotoxic drug therapy, trauma, or fungal or bacterial infections. Persistent ulcerative HSV infections are among the most common infections in patients with AIDS. HSV and *Candida* infections often occur concurrently. Systemic antiviral therapy speeds the rate of healing and relieves the pain of mucosal HSV infections in immunosuppressed patients. The frequency of HSV reactivation during the early phases of transplantation or induction chemotherapy is high (50–90%), and prophylactic systemic antiviral agents such as IV acyclovir and penciclovir or

1178 the oral congeners of these drugs are used to reduce reactivation rates. Patients with atopic eczema may also develop severe oral-facial HSV infections (*eczema herpeticum*), which may rapidly involve extensive areas of skin and occasionally disseminate to visceral organs. Extensive eczema herpeticum has resolved promptly with the administration of IV acyclovir. Erythema multiforme may also be associated with HSV infections (see Figs. 70-9 and 25e-25); some evidence suggests that HSV infection is the precipitating event in ~75% of cases of cutaneous erythema multiforme. HSV antigen has been demonstrated both in circulatory immune complexes and in skin lesion biopsy samples from these cases. Patients with severe HSV-associated erythema multiforme are candidates for chronic suppressive oral antiviral therapy.

HSV-1 and varicella-zoster virus (VZV) have been implicated in the etiology of Bell's palsy (flaccid paralysis of the mandibular portion of the facial nerve). Some but not all trials have documented quicker resolution of facial paralysis with the prompt initiation of antiviral therapy, with or without glucocorticoids. However, other trials have shown little benefit. Thus there is no consensus on the relative value of antiviral drugs alone, glucocorticoids alone, and the two modalities combined for the treatment of Bell's palsy.

Genital Infections First-episode primary genital herpes is characterized by fever, headache, malaise, and myalgias. Pain, itching, dysuria, vaginal and urethral discharge, and tender inguinal lymphadenopathy are the predominant local symptoms. Widely spaced bilateral lesions of the external genitalia are characteristic (Fig. 216-1). Lesions may be present in varying stages, including vesicles, pustules, or painful erythematous ulcers. The cervix and urethra are involved in >80% of women with first-episode infections. First episodes of genital herpes in patients who have had prior HSV-1 infection are associated with systemic symptoms in a few patients and with faster healing than primary genital herpes. Subclinical DNAemia has been found in ~30% of cases of true primary genital herpes. The clinical courses of acute first-episode genital herpes are similar for HSV-1 and HSV-2 infection. However, the recurrence rates of genital disease differ with the viral subtype: the 12-month recurrence rates among patients with first-episode HSV-2 and HSV-1 infections are ~90% and ~55%, respectively (median number of recurrences, 4 and <1, respectively). Recurrence rates for genital HSV-2 infections vary greatly among individuals and



FIGURE 216-1 Genital herpes: primary vulvar infection, with multiple, extremely painful, punched-out, confluent, shallow ulcers on the edematous vulva and perineum. Micturition is often very painful. Associated inguinal lymphadenopathy is common. (Reprinted with permission from K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.)

over time within the same individual. HSV has been isolated from the urethra and urine of men and women without external genital lesions. A clear mucoid discharge and dysuria are characteristics of symptomatic HSV urethritis. HSV has been isolated from the urethra of 5% of women with the dysuria-frequency syndrome. Occasionally, HSV genital tract disease is manifested by endometritis and salpingitis in women and by prostatitis in men. About 15% of cases of HSV-2 acquisition are associated with nonlesional clinical syndromes, such as aseptic meningitis, cervicitis, or urethritis. A more complete discussion of the differential diagnosis of genital herpes is presented in **Chap. 163**.

Both HSV-1 and HSV-2 can cause symptomatic or asymptomatic rectal and perianal infections. HSV proctitis is usually associated with rectal intercourse. However, subclinical perianal shedding of HSV is detected in women and men who report no rectal intercourse. This phenomenon is due to the establishment of latency in the sacral dermatome from prior genital tract infection, with subsequent reactivation in epithelial cells in the perianal region. Such reactivations are often subclinical. Symptoms of HSV proctitis include anorectal pain, anorectal discharge, tenesmus, and constipation. Sigmoidoscopy reveals ulcerative lesions of the distal 10 cm of the rectal mucosa. Rectal biopsies show mucosal ulceration, necrosis, polymorphonuclear and lymphocytic infiltration of the lamina propria, and (in occasional cases) multinucleated intranuclear inclusion-bearing cells. Perianal herpetic lesions are also found in immunosuppressed patients receiving cytotoxic therapy. Extensive perianal herpetic lesions and/or HSV proctitis is common among patients with HIV infection.

Herpetic Whitlow Herpetic whitlow—HSV infection of the finger—may occur as a complication of primary oral or genital herpes by inoculation of virus through a break in the epidermal surface or by direct introduction of virus into the hand through occupational or some other type of exposure. Clinical signs and symptoms include abrupt-onset edema, erythema, and localized tenderness of the infected finger. Vesicular or pustular lesions of the fingertip that are indistinguishable from lesions of pyogenic bacterial infection are seen. Fever, lymphadenitis, and epitrochlear and axillary lymphadenopathy are common. The infection may recur. Prompt diagnosis (to avoid unnecessary and potentially exacerbating surgical therapy and/or transmission) is essential. Antiviral chemotherapy is usually recommended (see below).


Herpes Gladiatorum HSV may infect almost any area of skin. Mucocutaneous HSV infections of the thorax, ears, face, and hands have been described among wrestlers. Transmission of these infections is facilitated by trauma to the skin sustained during wrestling. Several recent outbreaks have illustrated the importance of prompt diagnosis and therapy to contain the spread of this infection.

Eye Infections HSV infection of the eye is the most common cause of corneal blindness in the United States. HSV keratitis presents as an acute onset of pain, blurred vision, chemosis, conjunctivitis, and characteristic dendritic lesions of the cornea. Use of topical glucocorticoids may exacerbate symptoms and lead to involvement of deep structures of the eye. Debridement, topical antiviral treatment, and/or IFN therapy hastens healing. However, recurrences are common, and the deeper structures of the eye may sustain immunopathologic injury. Stromal keratitis due to HSV appears to be related to T cell-dependent destruction of deep corneal tissue. An HSV-1 epitope that is autoreactive with T cell-targeting corneal antigens has been postulated to be a factor in this infection. Chorioretinitis, usually a manifestation of disseminated HSV infection, may occur in neonates or in patients with HIV infection. HSV and VZV can cause acute necrotizing retinitis as an uncommon but severe manifestation.

Central and Peripheral Nervous System Infections HSV accounts for 10–20% of all cases of sporadic viral encephalitis in the United States. The estimated incidence is ~2.3 cases per 1 million persons per year. Cases are distributed throughout the year, and the age distribution appears to be biphasic, with peaks at 5–30 and >50 years of age. HSV-1 causes >95% of cases.

The pathogenesis of HSV encephalitis varies. In children and young adults, primary HSV infection may result in encephalitis; presumably,

exogenously acquired virus enters the CNS by neurotropic spread from the periphery via the olfactory bulb. However, most adults with HSV encephalitis have clinical or serologic evidence of mucocutaneous HSV-1 infection before the onset of CNS symptoms. In ~25% of the cases examined, the HSV-1 strains from the oropharynx and brain tissue of the same patient differ; thus some cases may result from reinfection with another strain of HSV-1 that reaches the CNS. Two theories have been proposed to explain the development of actively replicating HSV in localized areas of the CNS in persons whose ganglionic and CNS isolates are similar. Reactivation of latent HSV-1 infection in trigeminal or autonomic nerve roots may be associated with extension of virus into the CNS via nerves innervating the middle cranial fossa. HSV DNA has been demonstrated by DNA hybridization in brain tissue obtained at autopsy—even from healthy adults. Thus, reactivation of long-standing latent CNS infection may be another mechanism for the development of HSV encephalitis.

 Recent studies have identified genetic polymorphisms in two separate genes among families with a high frequency of HSV encephalitis. Peripheral-blood mononuclear cells from these patients (predominantly children) appear to secrete reduced levels of IFN in response to HSV. These observations suggest that some cases of sporadic HSV encephalitis may be related to host genetic determinants.

The clinical hallmark of HSV encephalitis has been the acute onset of fever and focal neurologic symptoms and signs, especially in the temporal lobe (Fig. 216-2). Clinical differentiation of HSV encephalitis from other viral encephalitides, focal infections, or noninfectious processes is difficult. Elevated cerebrospinal fluid (CSF) protein levels, leukocytosis (predominantly lymphocytes), and red blood cell counts due to hemorrhagic necrosis are common. While brain biopsy has been the gold standard for defining HSV encephalitis, a highly sensitive and specific PCR for detection of HSV DNA in CSF has largely replaced biopsy for defining CNS infection. Although titers of antibody to HSV in CSF and serum increase in most cases of HSV encephalitis, they rarely do so earlier than 10 days into the illness and therefore, although useful in retrospect, generally are not helpful in establishing an early clinical diagnosis. In rare cases, demonstration of HSV antigen, HSV DNA, or HSV replication in brain tissue obtained by biopsy is highly sensitive; examination of such tissue also provides the opportunity to identify alternative, potentially treatable causes of encephalitis. Antiviral chemotherapy with acyclovir reduces the rate of death from HSV encephalitis. Most authorities recommend the administration of IV acyclovir to patients with presumed HSV encephalitis until the

diagnosis is confirmed or an alternative diagnosis is made. All confirmed cases should be treated with IV acyclovir (30 mg/kg per day in three divided doses for 14–21 days). After the completion of therapy, the clinical recurrence of encephalitis requiring more treatment has been reported. For this reason, some authorities prefer to treat initially for 21 days, and many continue therapy until HSV DNA has been eliminated from the CSF. Even with therapy, neurologic sequelae are common, especially among persons >50 years of age.

HSV DNA has been detected in CSF from 3–15% of persons presenting to the hospital with aseptic meningitis. HSV meningitis, which is usually seen in association with primary genital HSV infection, is an acute, self-limited disease manifested by headache, fever, and mild photophobia and lasting 2–7 days. Lymphocytic pleocytosis in the CSF is characteristic. Neurologic sequelae of HSV meningitis are rare. HSV is the most commonly identified cause of recurrent lymphocytic meningitis (*Mollaret's meningitis*). Demonstration of HSV antibodies in CSF or persistence of HSV DNA in CSF can establish the diagnosis. For persons with frequent recurrences of HSV meningitis, daily antiviral therapy has reduced the occurrence of such episodes.

Autonomic nervous system dysfunction, especially of the sacral region, has been reported in association with both HSV and VZV infections. Numbness, tingling of the buttocks or perineal areas, urinary retention, constipation, CSF pleocytosis, and (in males) impotence may occur. Symptoms appear to resolve slowly over days or weeks. Occasionally, hypoesthesia and/or weakness of the lower extremities persists for many months. Transitory hypoesthesia of the area of skin innervated by the trigeminal nerve and vestibular system dysfunction (as measured by electronystagmography) are the predominant signs of disease. Whether antiviral chemotherapy can abort these signs or reduce their frequency and severity is not yet known. Rarely, transverse myelitis, manifested by a rapidly progressive symmetric paralysis of the lower extremities or Guillain-Barré syndrome, follows HSV infection. Similarly, peripheral nervous system involvement (Bell's palsy) or cranial polyneuritis may be related to reactivation of HSV-1 infection.

Visceral Infections HSV infection of visceral organs usually results from viremia, and multiple-organ involvement is common. Occasionally, however, the clinical manifestations of HSV infection involve only the esophagus, lung, or liver. HSV esophagitis may result from direct extension of oral-pharyngeal HSV infection into the esophagus or may occur de novo by reactivation and spread of HSV to the esophageal mucosa via the vagus nerve. The predominant symptoms of HSV esophagitis are odynophagia, dysphagia, substernal pain, and weight loss. Multiple oval ulcerations appear on an erythematous base with or without a patchy white pseudomembrane. The distal esophagus is most commonly involved. With extensive disease, diffuse friability may spread to the entire esophagus. Neither endoscopic nor barium examination can reliably differentiate HSV esophagitis from *Candida* esophagitis or from esophageal ulcerations due to thermal injury, radiation, or corrosives. Endoscopically obtained secretions for cytologic examination and culture or DNA detection by PCR provide the most useful material for diagnosis. Systemic antiviral chemotherapy usually reduces the severity and duration of symptoms and heals esophageal ulcerations.

HSV pneumonitis is uncommon except in severely immunosuppressed patients and may result from extension of herpetic tracheobronchitis into lung parenchyma. Focal necrotizing pneumonitis usually ensues. Hematogenous dissemination of virus from sites of oral or genital mucocutaneous

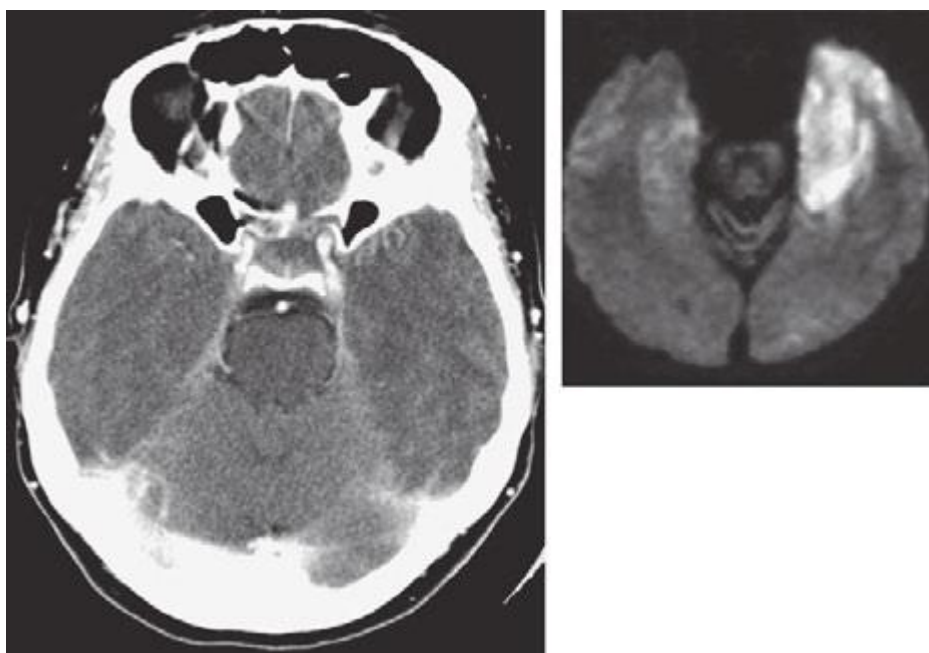


FIGURE 216-2 Computed tomography and diffusion-weighted magnetic resonance imaging scans of the brain of a patient with left-temporal-lobe herpes simplex virus encephalitis.

1180 disease may also occur, producing bilateral interstitial pneumonitis. Bacterial, fungal, and parasitic pathogens are commonly present in HSV pneumonitis. The mortality rate from untreated HSV pneumonia in immunosuppressed patients is high (>80%). HSV has also been isolated from the lower respiratory tract of persons with acute respiratory distress syndrome and prolonged intubation. Most authorities believe that the presence of HSV in tracheal aspirates in such settings is due to reactivation of HSV in the tracheal region and localized tracheitis in persons with long-term intubation. Such patients should be evaluated for extension of HSV infection into the lung parenchyma. Controlled trials assessing the role of antiviral agents used against HSV in morbidity and mortality associated with acute respiratory distress syndrome have not been conducted. The role of lower respiratory tract HSV infection in overall rates of morbidity and mortality associated with these conditions is unclear. HSV is an uncommon cause of hepatitis in immunocompetent patients. HSV infection of the liver is associated with fever, abrupt elevations of bilirubin and serum aminotransferase levels, and leukopenia (<4000 white blood cells/ μ L). Disseminated intravascular coagulation may also develop.

Other reported complications of HSV infection include monarticular arthritis, adrenal necrosis, idiopathic thrombocytopenia, and glomerulonephritis. Disseminated HSV infection in immunocompetent patients is rare. In immunocompromised patients, burn patients, or malnourished individuals, HSV occasionally disseminates to other visceral organs, such as the adrenal glands, pancreas, small and large intestines, and bone marrow. Rarely, primary HSV infection in pregnancy disseminates and may be associated with the death of both mother and fetus. This uncommon event is usually related to the acquisition of primary infection in the third trimester. Disseminated HSV infection is best detected by the presence of HSV DNA in plasma or blood.

Neonatal HSV Infections Of all HSV-infected populations, neonates (infants younger than 6 weeks) have the highest frequency of visceral and/or CNS infection. Without therapy, the overall rate of death from neonatal herpes is 65%; <10% of neonates with CNS infection develop normally. Although skin lesions are the most commonly recognized features of disease, many infants do not develop lesions at all or do so only well into the course of disease. Neonatal infection is usually acquired perinatally from contact with infected genital secretions at delivery. Congenitally infected infants have been reported. Of neonatal HSV infections, 30–50% are due to HSV-1 and 50–70% to HSV-2. The risk of developing neonatal HSV infection is 10 times higher for an infant born to a mother who has recently acquired HSV than for other infants. Neonatal HSV-1 infections may also be acquired through postnatal contact with immediate family members who have symptomatic or asymptomatic oral-labial HSV-1 infection or through nosocomial transmission within the hospital. All neonates with presumed herpes should be treated with IV acyclovir and then placed on maintenance oral antiviral therapy for the first 6–12 months of life. Antiviral chemotherapy with high-dose IV acyclovir (60 mg/kg per day) has reduced the mortality rate from neonatal herpes to ~15%. However, rates of morbidity, especially among infants with HSV-2 infection involving the CNS, are still very high.

HSV in Pregnancy In the United States, 22% of all pregnant women and 55% of non-Hispanic black pregnant women are seropositive for HSV-2. However, the risk of mother-to-child transmission of HSV in the perinatal period is highest when the infection is acquired near the time of labor—that is, in previously HSV-seronegative women. The clinical manifestations of recurrent genital herpes—including the frequency of subclinical versus clinical infection, duration of lesions, pain, and constitutional symptoms—are similar in pregnant and nonpregnant women. Recurrences increase in frequency over the course of pregnancy. However, when women are seropositive for HSV-2 at the outset of pregnancy, no effect on neonatal outcomes (including birth weight and gestational age) is seen. First-episode infections in pregnancy have more severe consequences for mother and infant. Maternal visceral dissemination during the third trimester occasionally occurs, as does premature birth or intrauterine growth retardation.

The acquisition of primary disease in pregnancy, whether related to HSV-1 or HSV-2, carries the risk of transplacental transmission of virus to the neonate and can result in spontaneous abortion, although this outcome is relatively uncommon. For newly acquired genital HSV infection during pregnancy, most authorities recommend treatment with acyclovir (400 mg three times daily) or valacyclovir (500–1000 mg twice daily) for 7–10 days. However, the impact of this intervention on transmission is unknown. The high HSV-2 prevalence rate in pregnancy and the low incidence of neonatal disease (1 case per 6000–20,000 live births) indicate that only a few infants are at risk of acquiring HSV. Therefore, cesarean section is not warranted for all women with recurrent genital disease. Because intrapartum transmission of infection accounts for the majority of cases, abdominal delivery need be considered only for women who are shedding HSV at delivery. Several studies have shown no correlation between recurrence of viral shedding before delivery and viral shedding at term. Hence, weekly virologic monitoring and amniocentesis are not recommended.

The frequency of transmission from mother to infant is markedly higher among women who acquire HSV near term (30–50%) than among those in whom HSV-2 infection is reactivated at delivery (<1%). Although maternal antibody to HSV-2 is protective, antibody to HSV-1 offers little or no protection against neonatal HSV-2 infection. Primary genital infection with HSV-1 leads to a particularly high risk of transmission during pregnancy and accounts for an increasing proportion of neonatal HSV cases. Moreover, during reactivation, HSV-1 appears more transmissible to the neonate than HSV-2. Only 2% of women who are seropositive for HSV-2 have HSV-2 isolated from cervical secretions at delivery, and only 1% of infants exposed in this manner develop infection, presumably because of the protective effects of maternally transferred antibodies and perhaps lower viral titers during reactivation. Despite the low frequency of transmission of HSV in this setting, 30–50% of infants with neonatal HSV are born to mothers with established genital herpes.

Isolation of HSV by cervicovaginal swab at the time of delivery is the greatest risk factor for intrapartum HSV transmission (relative risk = 346); however, culture-negative, PCR-positive cases of intrapartum transmission are well described. New acquisition of HSV (odds ratio [OR] = 49), isolation of HSV-1 versus HSV-2 (OR = 35), cervical versus vulvar HSV detection (OR = 15), use of fetal scalp electrodes (OR = 3.5), and young age confer further risk of transmission, whereas abdominal delivery is protective (OR = 0.14). Physical examination poorly predicts the absence of shedding, and PCR far exceeds culture in terms of sensitivity and speed. Therefore, PCR detection at the onset of labor should be used to aid clinical decision-making for women with HSV-2 antibody. Because cesarean section appears to be an effective means of reducing maternal-fetal transmission, patients with recurrent genital herpes should be encouraged to come to the hospital early at the time of delivery for careful examination of the external genitalia and cervix as well as collection of a swab sample for viral isolation. Women who have no evidence of lesions can have a vaginal delivery. The presence of active lesions on the cervix or external genitalia is an indication for cesarean delivery.

If first-episode exposure has occurred (e.g., if HSV serologies show that the mother is seronegative or if the mother is HSV-1-seropositive and the isolate at delivery is found to be HSV-2), many authorities would initiate antiviral therapy for the infant with IV acyclovir. At a minimum, samples for viral cultures and PCR should be obtained from the throat, nasopharynx, eyes, and rectum of these infants immediately and at 5- to 10-day intervals. Lethargy, skin lesions, or fever should be evaluated promptly. All infants from whom HSV is isolated 24 h after delivery should be treated with IV acyclovir at recommended doses.

DIAGNOSIS

Both clinical and laboratory criteria are useful for diagnosing HSV infections. A clinical diagnosis can be made accurately when characteristic multiple vesicular lesions on an erythematous base are present. However, herpetic ulcerations may resemble skin ulcerations of other etiologies. Mucosal HSV infections may also present as urethritis or pharyngitis without cutaneous lesions. Thus, laboratory studies to

confirm the diagnosis and to guide therapy are recommended. While staining of scrapings from the base of the lesions with Wright's, Giemsa's (Tzanck preparation), or Papanicolaou's stain to detect giant cells or intranuclear inclusions of *Herpesvirus* infection is a well-described procedure, few clinicians are skilled in this technique, the sensitivity of staining is low (<30% for mucosal swabs), and these cytologic methods do not differentiate between HSV and VZV infections.

HSV infection is best confirmed in the laboratory by detection of virus, viral antigen, or viral DNA in scrapings from lesions. HSV DNA detection by PCR is the most sensitive laboratory technique for detecting mucosal or visceral HSV infections and should be used when available. HSV causes a discernible cytopathic effect in a variety of cell culture systems, and this effect can be identified within 48–96 h after inoculation. Spin-amplified culture with subsequent staining for HSV antigen has shortened the time needed to identify HSV to <24 h. The sensitivity of all detection methods depends on the stage of the lesions (with higher sensitivity for vesicular than for ulcerative lesions), on whether the patient has a first or a recurrent episode of the disease (with higher sensitivity in first than in recurrent episodes), and on whether the sample is from an immunosuppressed or an immunocompetent patient (with more antigen or DNA in immunosuppressed patients). Laboratory confirmation permits subtyping of the virus; information on subtype may be useful epidemiologically and may help to predict the frequency of reactivation after first-episode oral-labial or genital HSV infection.

Serologic assays with whole-virus antigen preparations, such as complement fixation, neutralization, indirect immunofluorescence, passive hemagglutination, radioimmunoassay, and enzyme-linked immunosorbent assay, are useful for differentiating uninfected (seronegative) persons from those with past HSV-1 or HSV-2 infection, but they do not reliably distinguish between the two viral subtypes. Serologic assays that identify antibodies to type-specific surface proteins (epitopes) of the two viral subtypes have been developed and can distinguish reliably between the human antibody responses to HSV-1 and HSV-2. The most commonly used assays are those that measure antibodies to glycoprotein G of HSV-1 (gG1) and HSV-2 (gG2). A western blot assay that can detect several HSV type-specific proteins can also be used.

Acute- and convalescent-phase serum samples can be useful in demonstrating seroconversion during primary HSV-1 or HSV-2 infection. However, few available tests report titers, and increases in index values do not reflect first episodes in all patients. Serologic assays based on type-specific proteins should be used to identify asymptomatic carriers of HSV-1 or HSV-2. No reliable IgM method for defining acute HSV infection is available.

Several studies have shown that persons with previously unrecognized HSV-2 infection can be taught to identify symptomatic reactivations. Individuals seropositive for HSV-2 should be told about the high frequency of subclinical reactivation on mucosal surfaces that are not visible to the eye (e.g., cervix, urethra, perianal skin) or in microscopic ulcerations that may not be clinically symptomatic. Transmission of infection during such episodes is well established. HSV-2-seropositive persons should be educated about the high likelihood of subclinical shedding and the role condoms (male or female) may play in reducing transmission. Antiviral therapy with valacyclovir (500 mg once daily) has been shown to reduce the transmission of HSV-2 between sexual partners.

TREATMENT HERPES SIMPLEX VIRUS INFECTIONS

Many aspects of mucocutaneous and visceral HSV infections are amenable to antiviral chemotherapy. For mucocutaneous infections, acyclovir and its congeners famciclovir and valacyclovir have been the mainstays of therapy. Several antiviral agents are available for topical use in HSV eye infections: idoxuridine, trifluorothymidine, topical vidarabine, and cidofovir. For HSV encephalitis and neonatal herpes, IV acyclovir is the treatment of choice.

All licensed antiviral agents for use against HSV inhibit the viral DNA polymerase. One class of drugs, typified by the drug acyclovir, is made up of substrates for the HSV enzyme thymidine kinase (TK).

Acyclovir, ganciclovir, famciclovir, and valacyclovir are all selectively phosphorylated to the monophosphate form in virus-infected cells. Cellular enzymes convert the monophosphate form of the drug to the triphosphate, which is then incorporated into the viral DNA chain. Acyclovir is the agent most frequently used for the treatment of HSV infections and is available in IV, oral, and topical formulations. Valacyclovir, the valyl ester of acyclovir, offers greater bioavailability than acyclovir and thus can be administered less frequently. Famciclovir, the oral formulation of penciclovir, is clinically effective in the treatment of a variety of HSV-1 and HSV-2 infections. Ganciclovir is active against both HSV-1 and HSV-2; however, it is more toxic than acyclovir, valacyclovir, and famciclovir and generally is not recommended for the treatment of HSV infections. Anecdotal case reports suggest that ganciclovir may also be less effective than acyclovir for treatment of HSV infections. All three recommended compounds—acyclovir, valacyclovir, and famciclovir—have proved effective in shortening the duration of symptoms and lesions of mucocutaneous HSV infections in both immunocompromised and immunocompetent patients (Table 216-1). IV and oral formulations prevent reactivation of HSV in seropositive immunocompromised patients during induction chemotherapy or in the period immediately after bone marrow or solid organ transplantation. Chronic daily suppressive therapy reduces the frequency of reactivation disease among patients with frequent genital or oral-labial herpes. Only valacyclovir has been subjected to clinical trials that demonstrated reduced transmission of HSV-2 infection between sexual partners. IV acyclovir (30 mg/kg per day, given as a 10-mg/kg infusion over 1 h at 8-h intervals) is effective in reducing rates of death and morbidity from HSV encephalitis. Early initiation of therapy is a critical factor in outcome. The major side effect associated with IV acyclovir is transient renal insufficiency, usually due to crystallization of the compound in the renal parenchyma. This adverse reaction can be avoided if the medication is given slowly over 1 h and the patient is well hydrated. Because CSF levels of acyclovir average only 30–50% of plasma levels, the dosage of acyclovir used for treatment of CNS infection (30 mg/kg per day) is double that used for treatment of mucocutaneous or visceral disease (15 mg/kg per day). Even higher doses of IV acyclovir are used for neonatal HSV infection (60 mg/kg per day in three divided doses).

Increasingly, shorter courses of therapy are being used for recurrent mucocutaneous infection with HSV-1 or HSV-2 in immunocompetent patients. One-day courses of famciclovir and valacyclovir are clinically effective, more convenient, and generally less costly than longer courses of therapy (Table 216-1). These short-course regimens should be reserved for immunocompetent hosts.

SUPPRESSION OF MUCOCUTANEOUS HERPES

Recognition of the high frequency of subclinical reactivation provides a well-accepted rationale for the use of daily antiviral therapy to suppress reactivations of HSV, especially in persons with frequent clinical reactivations (e.g., those with recently acquired genital HSV infection). Immunosuppressed persons, including those with HIV infection, may also benefit from daily antiviral therapy. Recent studies have shown the efficacy of daily acyclovir and valacyclovir in reducing the frequency of HSV reactivations among HIV-positive persons. Regimens used include acyclovir (400–800 mg twice daily), famciclovir (500 mg twice daily), and valacyclovir (500 mg twice daily); valacyclovir at a dose of 4 g/d was associated with thrombotic thrombocytopenic purpura in one study of HIV-infected persons. In addition, daily treatment of HSV-2 reduces the titer of HIV RNA in plasma (0.5-log reduction) and in genital mucosa (0.33-log reduction).

REDUCED HSV TRANSMISSION TO SEXUAL PARTNERS

Once-daily valacyclovir (500 mg) has been shown to reduce transmission of HSV-2 between sexual partners. Transmission rates are higher from males to females and among persons with frequent HSV-2 reactivation. Serologic screening can be used to identify at-risk couples. Daily valacyclovir appears to be more effective at reducing subclinical shedding than daily famciclovir.

I. Mucocutaneous HSV infections**A. Infections in immunosuppressed patients**

1. *Acute symptomatic first or recurrent episodes:* IV acyclovir (5 mg/kg q8h) or oral acyclovir (400 mg qid), famciclovir (500 mg bid or tid), or valacyclovir (500 mg bid) is effective. Treatment duration may vary from 7 to 14 days.
2. *Suppression of reactivation disease (genital or oral-labial):* IV acyclovir (5 mg/kg q8h) or oral valacyclovir (500 mg bid) or acyclovir (400–800 mg 3–5 times per day) prevents recurrences during the 30-day period immediately after transplantation. Longer-term HSV suppression is often used for persons with continued immunosuppression. In bone marrow and renal transplant recipients, oral valacyclovir (2 g/d) is also effective in reducing cytomegalovirus infection. Oral valacyclovir at a dose of 4 g/d has been associated with thrombotic thrombocytopenic purpura after extended use in HIV-positive persons. In HIV-infected persons, oral acyclovir (400–800 mg bid), valacyclovir (500 mg bid), or famciclovir (500 mg bid) is effective in reducing clinical and subclinical reactivations of HSV-1 and HSV-2.

B. Infections in immunocompetent patients**1. Genital herpes**

- a. *First episodes:* Oral acyclovir (200 mg 5 times per day or 400 mg tid), valacyclovir (1 g bid), or famciclovir (250 mg bid) for 7–14 days is effective. IV acyclovir (5 mg/kg q8h for 5 days) is given for severe disease or neurologic complications such as aseptic meningitis.
- b. *Symptomatic recurrent genital herpes:* Short-course (1- to 3-day) regimens are preferred because of low cost, likelihood of adherence, and convenience. Oral acyclovir (800 mg tid for 2 days), valacyclovir (500 mg bid for 3 days), or famciclovir (750 or 1000 mg bid for 1 day, a 1500-mg single dose, or 500 mg stat followed by 250 mg q12h for 3 days) effectively shortens lesion duration. Other options include oral acyclovir (200 mg 5 times per day), valacyclovir (500 mg bid), and famciclovir (125 mg bid for 5 days).
- c. *Suppression of recurrent genital herpes:* Oral acyclovir (400–800 mg bid) or valacyclovir (500 mg daily) is given. Patients with >9 episodes per year should take oral valacyclovir (1 g daily or 500 mg bid) or famciclovir (250 mg bid or 500 mg bid).

2. Oral-labial HSV infections

- a. *First episode:* Oral acyclovir is given (200 mg 5 times per day or 400 mg tid); an oral acyclovir suspension can be used (600 mg/m² qid). Oral famciclovir (250 mg bid) or valacyclovir (1 g bid) has been used clinically. The duration of therapy is 5–10 days.
 - b. *Recurrent episodes:* If initiated at the onset of the prodrome, single-dose or 1-day therapy effectively reduces pain and speeds healing. Regimens include oral famciclovir (a 1500-mg single dose or 750 mg bid for 1 day) or valacyclovir (a 2-g single dose or 2 g bid for 1 day). Self-initiated therapy with 6-times-daily topical penciclovir cream effectively speeds healing of oral-labial HSV. Topical acyclovir cream has also been shown to speed healing.
 - c. *Suppression of reactivation of oral-labial HSV:* If started before exposure and continued for the duration of exposure (usually 5–10 days), oral acyclovir (400 mg bid) prevents reactivation of recurrent oral-labial HSV infection associated with severe sun exposure.
3. *Surgical prophylaxis of oral or genital HSV infection:* Several surgical procedures, such as laser skin resurfacing, trigeminal nerve-root decompression, and lumbar disk surgery, have been associated with HSV reactivation. IV acyclovir (3–5 mg/kg q8h) or oral acyclovir (800 mg bid), valacyclovir (500 mg bid), or famciclovir (250 mg bid) effectively reduces reactivation. Therapy should be initiated 48 h before surgery and continued for 3–7 days.
 4. *Herpetic whitlow:* Oral acyclovir (200 mg) is given 5 times daily (alternative: 400 mg tid) for 7–10 days.
 5. *HSV proctitis:* Oral acyclovir (400 mg 5 times per day) is useful in shortening the course of infection. In immunosuppressed patients or in patients with severe infection, IV acyclovir (5 mg/kg q8h) may be useful.
 6. *Herpetic eye infections:* In acute keratitis, topical trifluorothymidine, vidarabine, idoxuridine, acyclovir, penciclovir, and interferon are all beneficial. Debridement may be required. Topical steroids may worsen disease.

II. Central nervous system HSV infections

- A. *HSV encephalitis:* IV acyclovir (10 mg/kg q8h; 30 mg/kg per day) is given for 10 days or until HSV DNA is no longer detected in cerebrospinal fluid.
- B. *HSV aseptic meningitis:* No studies of systemic antiviral chemotherapy exist. If therapy is to be given, IV acyclovir (15–30 mg/kg per day) should be used.
- C. *Autonomic radiculopathy:* No studies are available. Most authorities recommend a trial of IV acyclovir.

III. Neonatal HSV infections: Oral acyclovir (60 mg/kg per day, divided into 3 doses) is given. The recommended duration of IV treatment is 21 days. Monitoring for relapse should be undertaken. Continued suppression with oral acyclovir suspension should be given for 3–4 months.

IV. Visceral HSV infections

- A. *HSV esophagitis:* IV acyclovir (15 mg/kg per day) is given. In some patients with milder forms of immunosuppression, oral therapy with valacyclovir or famciclovir is effective.
- B. *HSV pneumonitis:* No controlled studies exist. IV acyclovir (15 mg/kg per day) should be considered.

V. Disseminated HSV infections: No controlled studies exist. IV acyclovir (5 mg/kg q8h) should be tried. Adjustments for renal insufficiency may be needed. No definite evidence indicates that therapy will decrease the risk of death.

VI. Erythema multiforme associated with HSV: Anecdotal observations suggest that oral acyclovir (400 mg bid or tid) or valacyclovir (500 mg bid) will suppress erythema multiforme.

VII. Infections due to acyclovir-resistant HSV: IV foscarnet (40 mg/kg IV q8h) should be given until lesions heal. The optimal duration of therapy and the usefulness of its continuation to suppress lesions are unclear. Some patients may benefit from cutaneous application of trifluorothymidine or 5% cidofovir gel.

ACYCLOVIR RESISTANCE

Acyclovir-resistant strains of HSV have been identified. Most of these strains have an altered substrate specificity for phosphorylating acyclovir. Thus, cross-resistance to famciclovir and valacyclovir is usually found. Occasionally, an isolate with altered TK specificity arises and is sensitive to famciclovir but not to acyclovir. In some patients infected with TK-deficient virus, higher doses of acyclovir are associated with clearing of lesions. In others, clinical disease progresses despite high-dose therapy. Almost all clinically significant acyclovir resistance has been seen in immunocompromised patients, and HSV-2 isolates are more often resistant than HSV-1 strains. A study by the Centers for Disease Control and Prevention indicated that

~5% of HSV-2 isolates from HIV-positive persons exhibit some degree of *in vitro* resistance to acyclovir. Of HSV-2 isolates from immunocompetent patients attending sexually transmitted disease clinics, <0.5% show reduced *in vitro* sensitivity to acyclovir. The lack of appreciable change in the frequency of detection of such isolates in the past 20 years probably reflects the reduced transmission of TK-deficient mutants. Isolation of HSV from lesions persisting despite adequate dosages and blood levels of acyclovir should raise the suspicion of acyclovir resistance. Therapy with the antiviral drug foscarnet is useful in acyclovir-resistant cases (Chap. 215e). Because of its toxicity and cost, this drug is usually reserved for patients with extensive mucocutaneous infections. Cidofovir is a nucleotide

analogue and exists as a phosphonate or monophosphate form. Most TK-deficient strains of HSV are sensitive to cidofovir. Cidofovir ointment speeds healing of acyclovir-resistant lesions. No well-controlled trials of systemic cidofovir have been reported. True TK-negative variants of HSV appear to have a reduced capacity to spread because of altered neurovirulence—a feature important in the relatively infrequent presence of such strains in immunocompetent populations, even with increasing use of antiviral drugs.

ACYCLOVIR EFFICACY IN THE DEVELOPING WORLD



Early studies of acyclovir-like drugs were performed solely in the developed world. Recent studies have shown that, although acyclovir-like drugs are effective in the developing world, their clinical and virologic benefits seem reduced from those in European and U.S. populations. The mechanism of this phenomenon is uncertain. Acyclovir therapy does not reduce the rate of HIV acquisition; however, HIV load among MSM in the United States decreased by 1.3 log₁₀ in contrast to 0.9 log₁₀ among Peruvian MSM and 0.5 log₁₀ among African women.

PREVENTION

The success of efforts to control HSV disease on a population basis through suppressive antiviral chemotherapy and/or educational programs will be limited. Barrier forms of contraception (especially condoms) decrease the likelihood of transmission of HSV infection, particularly during periods of asymptomatic viral excretion. When lesions are present, HSV infection may be transmitted by skin-to-skin contact despite the use of a condom. Nevertheless, the available data suggest that consistent condom use is an effective means of reducing the risk of genital HSV-2 transmission. Chronic daily antiviral therapy with valacyclovir can also be partially effective in reducing acquisition of HSV-2, especially among susceptible women. There are no comparative efficacy studies of valacyclovir versus condom use. Most authorities suggest both approaches. The need for a vaccine to prevent acquisition of HSV infection is great, especially in light of the role HSV-2 plays in enhancing the acquisition and transmission of HIV-1.

A substantial portion of neonatal HSV cases could be prevented by reducing the acquisition of HSV by women in the third trimester of pregnancy. Neonatal HSV infection can result from either the acquisition of maternal infection near term or the reactivation of infection at delivery in the already-infected mother. Thus strategies for reducing neonatal HSV are complex. Some authorities have recommended that antiviral therapy with acyclovir or valacyclovir be given to HSV-2-infected women in late pregnancy as a means of reducing reactivation of HSV-2 at term. Data are not available to support the efficacy of this approach. Moreover, the high treatment-to-prevention ratio makes this a dubious public health approach, even though it can reduce the frequency of HSV-associated cesarean delivery.

demonstrated. Viral isolates from patients with chickenpox and herpes zoster produced similar alterations in tissue culture—specifically, the appearance of eosinophilic intranuclear inclusions and multinucleated giant cells. These results suggested that the viruses were biologically similar. Restriction endonuclease analyses of viral DNA from a patient with chickenpox who subsequently developed herpes zoster verified the molecular identity of the two viruses responsible for these different clinical presentations.

VZV is a member of the family Herpesviridae, sharing with other members such structural characteristics as a lipid envelope surrounding a nucleocapsid with icosahedral symmetry, a total diameter of ~180–200 nm, and centrally located double-stranded DNA that is ~125,000 bp in length.

PATHOGENESIS AND PATHOLOGY

Primary Infection Transmission occurs readily by the respiratory route; the subsequent localized replication of the virus at an undefined site (presumably the nasopharynx) leads to seeding of the lymphatic/reticuloendothelial system and ultimately to the development of viremia. Viremia in patients with chickenpox is reflected in the diffuse and scattered nature of the skin lesions and can be confirmed in selected cases by the recovery of VZV from the blood or routinely by the detection of viral DNA in either blood or lesions by polymerase chain reaction (PCR). Vesicles involve the corium and dermis, with degenerative changes characterized by ballooning, the presence of multinucleated giant cells, and eosinophilic intranuclear inclusions. Infection may involve localized blood vessels of the skin, resulting in necrosis and epidermal hemorrhage. With the evolution of disease, the vesicular fluid becomes cloudy because of the recruitment of polymorphonuclear leukocytes and the presence of degenerated cells and fibrin. Ultimately, the vesicles either rupture and release their fluid (which includes infectious virus) or are gradually reabsorbed.

Recurrent Infection The mechanism of reactivation of VZV that results in herpes zoster is unknown. Presumably, the virus infects dorsal root ganglia during chickenpox, where it remains latent until reactivated. Histopathologic examination of representative dorsal root ganglia during active herpes zoster demonstrates hemorrhage, edema, and lymphocytic infiltration.

Active replication of VZV in other organs, such as the lung or the brain, can occur during either chickenpox or herpes zoster but is uncommon in the immunocompetent host. Pulmonary involvement is characterized by interstitial pneumonitis, multinucleated giant cell formation, intranuclear inclusions, and pulmonary hemorrhage. Central nervous system (CNS) infection leads to histopathologic evidence of perivascular cuffing similar to that encountered in measles and other viral encephalitides. Focal hemorrhagic necrosis of the brain, characteristic of herpes simplex virus (HSV) encephalitis, develops infrequently in VZV infection.

EPIDEMIOLOGY AND CLINICAL MANIFESTATIONS

Chickenpox Humans are the only known reservoir for VZV. Chickenpox is highly contagious, with an attack rate of at least 90% among susceptible (seronegative) individuals. Persons of both sexes and all races are infected equally. The virus is endemic in the population at large; however, it becomes epidemic among susceptible individuals during seasonal peaks—namely, late winter and early spring in the temperate zone. Much of our knowledge of the disease's natural history and incidence predates the licensure of the chickenpox vaccine in 1995. Historically, children 5–9 years old are most commonly affected and account for 50% of all cases. Most other cases involve children 1–4 and 10–14 years old. Approximately 10% of the population of the United States over the age of 15 is susceptible to infection. VZV vaccination during the second year of life has dramatically changed the epidemiology of infection, causing a significant decrease in the annualized incidence of chickenpox.

The incubation period of chickenpox ranges from 10 to 21 days but is usually 14–17 days. Secondary attack rates in susceptible siblings

217 Varicella-Zoster Virus Infections

Richard J. Whitley

DEFINITION

Varicella-zoster virus (VZV) causes two distinct clinical entities: varicella (chickenpox) and herpes zoster (shingles). Chickenpox, a ubiquitous and extremely contagious infection, is usually a benign illness of childhood characterized by an exanthematous vesicular rash. With reactivation of latent VZV (which is most common after the sixth decade of life), herpes zoster presents as a dermatomal vesicular rash, usually associated with severe pain.

ETIOLOGY

Early in the twentieth century, similarities in the histopathologic features of skin lesions resulting from varicella and herpes zoster were



FIGURE 217-1 Varicella lesions at various stages of evolution: vesicles on an erythematous base, umbilical vesicles, and crusts.

within a household are 70–90%. Patients are infectious ~48 h before onset of the vesicular rash, during the period of vesicle formation (which generally lasts 4–5 days), and until all vesicles are crusted.

Clinically, chickenpox presents as a rash, low-grade fever, and malaise, although a few patients develop a prodrome 1–2 days before onset of the exanthem. In the immunocompetent patient, chickenpox is usually a benign illness associated with lassitude and with body temperatures of 37.8°–39.4°C (100°–103°F) of 3–5 days' duration. The skin lesions—the hallmark of the infection—include maculopapules, vesicles, and scabs in various stages of evolution (Fig. 217-1). These lesions, which evolve from maculopapules to vesicles over hours to days, appear on the trunk and face and rapidly spread to involve other areas of the body. Most are small and have an erythematous base with a diameter of 5–10 mm. Successive crops appear over a 2- to 4-day period. Lesions can also be found on the mucosa of the pharynx and/or the vagina. Their severity varies from one person to another. Some individuals have very few lesions, while others have as many as 2000. Younger children tend to have fewer vesicles than older individuals. Secondary and tertiary cases within families are associated with a relatively large number of vesicles. Immunocompromised patients—both children and adults, particularly those with leukemia—have lesions (often with a hemorrhagic base) that are more numerous and take longer to heal than those of immunocompetent patients. Immunocompromised individuals are also at greater risk for visceral complications, which occur in 30–50% of cases and are fatal 15% of the time in the absence of antiviral therapy.

The most common infectious complication of varicella is secondary bacterial superinfection of the skin, which is usually caused by *Streptococcus pyogenes* or *Staphylococcus aureus*, including strains that are methicillin-resistant. Skin infection results from excoriation of lesions after scratching. Gram's staining of skin lesions should help clarify the etiology of unusually erythematous and pustulated lesions.

The most common extracutaneous site of involvement in children is the CNS. The syndrome of acute cerebellar ataxia and meningeal inflammation generally appears ~21 days after onset of the rash and rarely develops in the pre-eruptive phase. The cerebrospinal fluid (CSF) contains lymphocytes and elevated levels of protein. CNS involvement is a benign complication of VZV infection in children and generally does not require hospitalization. Aseptic meningitis, encephalitis, transverse myelitis, and Guillain-Barré syndrome can also occur. Reye's syndrome has been reported in children concomitantly treated with aspirin. Encephalitis is reported in 0.1–0.2% of children with chickenpox. Other than supportive care, no specific therapy (e.g., acyclovir administration) has proved efficacious for patients with CNS involvement.

Varicella pneumonia, the most serious complication following chickenpox, develops more often in adults (up to 20% of cases) than in children and is particularly severe in pregnant women. Pneumonia due to VZV usually has its onset 3–5 days into the illness and is associated with tachypnea, cough, dyspnea, and fever. Cyanosis, pleuritic chest pain, and hemoptysis are frequently noted. Roentgenographic evidence of disease consists of nodular infiltrates and interstitial pneumonitis. Resolution of pneumonitis parallels improvement of the skin rash; however, patients may have persistent fever and compromised pulmonary function for weeks.

Other complications of chickenpox include myocarditis, corneal lesions, nephritis, arthritis, bleeding diatheses, acute glomerulonephritis, and hepatitis. Hepatic involvement, distinct from Reye's syndrome and usually asymptomatic, is common in chickenpox and is generally characterized by elevated levels of liver enzymes, particularly aspartate and alanine aminotransferases.

Perinatal varicella is associated with mortality rates as high as 30% when maternal disease develops within 5 days before delivery or within 48 h thereafter. Illness in this setting is unusually severe because the newborn does not receive protective transplacental antibodies and has an immature immune system. *Congenital varicella*, with clinical manifestations of limb hypoplasia, cicatricial skin lesions, and microcephaly at birth, is extremely uncommon.

Herpes Zoster Herpes zoster (shingles) is a sporadic disease that results from reactivation of latent VZV from dorsal root ganglia. Most patients with shingles have no history of recent exposure to other individuals with VZV infection. Herpes zoster occurs at all ages, but its incidence is highest (5–10 cases per 1000 persons) among individuals in the sixth decade of life and beyond. Data suggest that 1.2 million cases occur annually in the United States. Recurrent herpes zoster is exceedingly rare except in immunocompromised hosts, especially those with AIDS.

Herpes zoster is characterized by a unilateral vesicular dermatomal eruption, often associated with severe pain. The dermatomes from T3 to L3 are most frequently involved. If the ophthalmic branch of the trigeminal nerve is involved, *zoster ophthalmicus* results. The factors responsible for the reactivation of VZV are not known. In children, reactivation is usually benign; in adults, it can be debilitating because of pain. The onset of disease is heralded by pain within the dermatome, which may precede lesions by 48–72 h; an erythematous maculopapular rash evolves rapidly into vesicular lesions (Fig. 217-2). In the normal host, these lesions may remain few in number and continue to form for



FIGURE 217-2 Close-up of lesions of disseminated zoster. Note lesions at different stages of evolution, including pustules and crusting. (Photo courtesy of Lindsey Baden; with permission.)

only 3–5 days. The total duration of disease is generally 7–10 days; however, it may take as long as 2–4 weeks for the skin to return to normal. Patients with herpes zoster can transmit infection to seronegative individuals, with consequent chickenpox. In a few patients, characteristic localization of pain to a dermatome with serologic evidence of herpes zoster has been reported in the absence of skin lesions, an entity known as *zoster sine herpetica*. When branches of the trigeminal nerve are involved, lesions may appear on the face, in the mouth, in the eye, or on the tongue. *Zoster ophthalmicus* is usually a debilitating condition that can result in blindness in the absence of antiviral therapy. In *Ramsay Hunt syndrome*, pain and vesicles appear in the external auditory canal, and patients lose their sense of taste in the anterior two-thirds of the tongue while developing ipsilateral facial palsy. The geniculate ganglion of the sensory branch of the facial nerve is involved.

In both normal and immunocompromised hosts, the most debilitating complication of herpes zoster is pain associated with acute neuritis and postherpetic neuralgia. Postherpetic neuralgia is uncommon in young individuals; however, at least 50% of zoster patients over age 50 report some degree of pain in the involved dermatome for months after the resolution of cutaneous disease. Changes in sensation in the dermatome, resulting in either hypo- or hyperesthesia, are common.

CNS involvement may follow localized herpes zoster. Many patients without signs of meningeal irritation have CSF pleocytosis and moderately elevated levels of CSF protein. Symptomatic meningoencephalitis is characterized by headache, fever, photophobia, meningitis, and vomiting. A rare manifestation of CNS involvement is granulomatous angiitis with contralateral hemiplegia, which can be diagnosed by cerebral arteriography. Other neurologic manifestations include transverse myelitis with or without motor paralysis.

Like chickenpox, herpes zoster is more severe in immunocompromised than immunocompetent individuals. Lesions continue to form for >1 week, and scabbing is not complete in most cases until 3 weeks into the illness. Patients with Hodgkin's disease and non-Hodgkin's lymphoma are at greatest risk for progressive herpes zoster. Cutaneous dissemination (**Fig. 217-3**) develops in ~40% of these patients. Among patients with cutaneous dissemination, the risk of pneumonitis, meningoencephalitis, hepatitis, and other serious complications is increased by 5–10%. However, even in immunocompromised patients, disseminated zoster is rarely fatal.

Recipients of hematopoietic stem cell transplants are at particularly high risk of VZV infection. Of all cases of posttransplantation VZV infection, 30% occur within 1 year (50% of these within 9 months); 45% of the patients involved have cutaneous or visceral dissemination. The mortality rate in this situation is 10%. Postherpetic neuralgia,



FIGURE 217-3 Herpes zoster in an HIV-infected patient is seen as hemorrhagic vesicles and pustules on an erythematous base grouped in a dermatomal distribution.

scarring, and bacterial superinfection are especially common in VZV infections occurring within 9 months of transplantation. Among infected patients, concomitant graft-versus-host disease increases the chance of dissemination and/or death.

DIFFERENTIAL DIAGNOSIS

(See also **Chap. 25e**) The diagnosis of chickenpox is not difficult. The characteristic rash and a history of recent exposure should lead to a prompt diagnosis. Other viral infections that can mimic chickenpox include disseminated HSV infection in patients with atopic dermatitis and the disseminated vesiculopapular lesions sometimes associated with coxsackievirus infection, echovirus infection, or atypical measles. However, these rashes are more commonly morbilliform with a hemorrhagic component rather than vesicular or vesiculopustular. Rickettsialpox (**Chap. 211**) is sometimes confused with chickenpox; however, rickettsialpox can be distinguished easily by detection of the “herald spot” at the site of the mite bite and the development of a more pronounced headache. Serologic testing is also useful in differentiating rickettsialpox from varicella and can confirm susceptibility in adults unsure of their chickenpox history. Concern about smallpox has recently increased because of the threat of bioterrorism (**Chap. 261e**). The lesions of smallpox are larger than those of chickenpox and are all at the same stage of evolution at any given time.

Unilateral vesicular lesions in a dermatomal pattern should lead rapidly to the diagnosis of herpes zoster, although the occurrence of shingles without a rash has been reported. Both HSV and coxsackievirus infections can cause dermatomal vesicular lesions. Supportive diagnostic virology and fluorescent staining of skin scrapings with monoclonal antibodies are helpful in ensuring the proper diagnosis. In the prodromal stage of herpes zoster, the diagnosis can be exceedingly difficult and may be made only after lesions have appeared or by retrospective serologic assessment.

LABORATORY FINDINGS

Unequivocal confirmation of the diagnosis is possible only through the isolation of VZV in susceptible tissue-culture cell lines, the demonstration of either seroconversion or a fourfold or greater rise in antibody titer between acute-phase and convalescent-phase serum specimens, or the detection of VZV DNA by PCR. A rapid impression can be obtained by a Tzanck smear, with scraping of the base of the lesions in an attempt to demonstrate multinucleated giant cells; however, the sensitivity of this method is low (~60%). PCR technology for the detection of viral DNA in vesicular fluid is available in a limited number of diagnostic laboratories. Direct immunofluorescent staining of cells from the lesion base or detection of viral antigens by other assays (such as the immunoperoxidase assay) is also useful, although these tests are not commercially available. The most frequently employed serologic tools for assessing host response are the immunofluorescent detection of antibodies to VZV membrane antigens, the fluorescent antibody to membrane antigen (FAMA) test, immune adherence hemagglutination, and enzyme-linked immunosorbent assay (ELISA). The FAMA test and the ELISA appear to be most sensitive.

TREATMENT VARICELLA-ZOSTER VIRUS INFECTIONS

Medical management of chickenpox in the immunologically normal host is directed toward the prevention of avoidable complications. Obviously, good hygiene includes daily bathing and soaks. Secondary bacterial infection of the skin can be avoided by meticulous skin care, particularly with close cropping of fingernails. Pruritus can be decreased with topical dressings or the administration of antipruritic drugs. Tepid water baths and wet compresses are better than drying lotions for the relief of itching. Administration of aspirin to children with chickenpox should be avoided because of the association of aspirin derivatives with the development of Reye's syndrome. Acyclovir (800 mg by mouth five times daily), valacyclovir (1 g three times daily), or famciclovir (250 mg three times daily) for 5–7 days is recommended for adolescents and adults with chickenpox of

1186 ≤24 h duration. (Valacyclovir is licensed for use in children and adolescents. Famciclovir is recommended but not licensed for varicella.) Likewise, acyclovir therapy may be of benefit to children <12 years of age if initiated early in the disease (<24 h) at a dose of 20 mg/kg every 6 h. The advantages (i.e., pharmacokinetics) of the second-generation agents valacyclovir and famciclovir are described in [Chap. 215e](#).

Aluminum acetate soaks for the management of herpes zoster can be both soothing and cleansing. Patients with herpes zoster benefit from oral antiviral therapy, as evidenced by accelerated healing of lesions and resolution of zoster-associated pain with acyclovir, valacyclovir, or famciclovir. Acyclovir is administered at a dosage of 800 mg five times daily for 7–10 days. However, valacyclovir and famciclovir are superior in terms of pharmacokinetics and pharmacodynamics and should be used preferentially. Famciclovir, the prodrug of penciclovir, is at least as effective as acyclovir and perhaps more so; the dose is 500 mg by mouth three times daily for 7 days. Valacyclovir, the prodrug of acyclovir, accelerates healing and resolution of zoster-associated pain more promptly than acyclovir. The dose is 1 g by mouth three times daily for 5–7 days. Compared with acyclovir, both famciclovir and valacyclovir offer the advantage of less frequent administration. All three of these drugs are now off patent.

In severely immunocompromised hosts (e.g., transplant recipients, patients with lymphoproliferative malignancies), both chickenpox and herpes zoster (including disseminated disease) should be treated, at least at the outset, with IV acyclovir, which reduces the occurrence of visceral complications but has no effect on healing of skin lesions or pain. The dose is 10 mg/kg every 8 h for 7 days. For low-risk immunocompromised hosts, oral therapy with valacyclovir or famciclovir appears beneficial. If medically feasible, it is desirable to decrease immunosuppressive treatment concomitant with the administration of IV acyclovir.

Patients with varicella pneumonia often require ventilatory support. Persons with zoster ophthalmicus should be referred immediately to an ophthalmologist. Therapy for this condition consists of the administration of analgesics for severe pain and the use of atropine. Acyclovir, valacyclovir, and famciclovir all accelerate healing. Decisions about the use of glucocorticoids should be made by the ophthalmologist.

The management of acute neuritis and/or postherpetic neuralgia can be particularly difficult. In addition to the judicious use of analgesics ranging from nonnarcotics to narcotic derivatives, drugs such as gabapentin, pregabalin, amitriptyline hydrochloride, lidocaine (patches), and fluphenazine hydrochloride are reportedly beneficial for pain relief. In one study, glucocorticoid therapy administered early in the course of localized herpes zoster significantly accelerated such quality-of-life improvements as a return to usual activity and termination of analgesic medications. The dose of prednisone administered orally was 60 mg/d on days 1–7, 30 mg/d on days 8–14, and 15 mg/d on days 15–21. This regimen is appropriate only for relatively healthy elderly persons with moderate or severe pain at presentation. Patients with osteoporosis, diabetes mellitus, glycosuria, or hypertension may not be appropriate candidates. Glucocorticoids should not be used without concomitant antiviral therapy.

PREVENTION

Three methods are used for the prevention of VZV infections. First, a live attenuated varicella vaccine (Oka) is recommended for all children >1 year of age (up to 12 years of age) who have not had chickenpox and for adults known to be seronegative for VZV. Two doses are recommended for all children: the first at 12–15 months of age and the second at ~4–6 years of age. VZV-seronegative persons >13 years of age should receive two doses of vaccine at least 1 month apart. The vaccine is both safe and efficacious. Breakthrough cases are mild and may result in spread of the vaccine virus to susceptible contacts. The universal vaccination of children is resulting in a decreased incidence of chickenpox in sentinel communities. Furthermore, inactivation of the vaccine virus significantly decreases the occurrence of herpes zoster after hematopoietic stem-cell transplantation.

TABLE 217-1 RECOMMENDATIONS FOR VZIG ADMINISTRATION

Exposure Criteria
1. Exposure to a person with chickenpox or zoster <ol style="list-style-type: none"> Household: residence in the same household Playmate: face-to-face indoor play Hospital <p>Varicella: same 2- to 4-bed room or adjacent beds in large ward, face-to-face contact with infectious staff member or patient, visit by a person deemed contagious</p> <p>Zoster: intimate contact (e.g., touching or hugging) with a person deemed contagious</p> Newborn infant: onset of varicella in the mother ≤5 days before delivery or ≤48 h after delivery; VZIG not indicated if the mother has zoster
2. Patient should receive VZIG as soon as possible but not >96 h after exposure
Candidates (Provided They Have Significant Exposure) Include:
1. Immunocompromised susceptible children without a history of varicella or varicella immunization
2. Susceptible pregnant women
3. Newborn infants whose mother had onset of chickenpox within 5 days before or within 48 h after delivery
4. Hospitalized premature infant (≥28 weeks of gestation) whose mother lacks a reliable history of chickenpox or serologic evidence of protection against varicella
5. Hospitalized premature infant (<28 weeks of gestation or ≤1000-g birth weight), regardless of maternal history of varicella or VZV serologic status

In individuals >50 years of age, a VZV vaccine with 18 times the viral content of the Oka vaccine decreased the incidence of shingles by 51%, the burden of illness by 61%, and the incidence of postherpetic neuralgia by 66%. The Advisory Committee on Immunization Practices has therefore recommended that persons in this age group be offered this vaccine in order to reduce the frequency of shingles and the severity of postherpetic neuralgia.

A second approach is to administer varicella-zoster immune globulin (VZIG) to individuals who are susceptible, are at high risk for developing complications of varicella, and have had a significant exposure. This product should be given within 96 h (preferably within 72 h) of the exposure. Indications for administration of VZIG appear in [Table 217-1](#).

Lastly, antiviral therapy can be given as prophylaxis to individuals at high risk who are ineligible for vaccine or who are beyond the 96-h window after direct contact. While the initial studies have used acyclovir, similar benefit can be anticipated with either valacyclovir or famciclovir. Therapy is instituted 7 days after intense exposure. At this time, the host is midway into the incubation period. This approach significantly decreases disease severity, if not totally preventing disease.

218 Epstein-Barr Virus Infections, Including Infectious Mononucleosis

Jeffrey I. Cohen

DEFINITION

Epstein-Barr virus (EBV) is the cause of heterophile-positive infectious mononucleosis (IM), which is characterized by fever, sore throat, lymphadenopathy, and atypical lymphocytosis. EBV is also associated with several tumors, including nasopharyngeal and gastric carcinoma, Burkitt's lymphoma, Hodgkin's disease, and (in patients

with immunodeficiencies) B cell lymphoma. The virus is a member of the family Herpesviridae. The two types of EBV that are widely prevalent in nature are not distinguishable by conventional serologic tests.

EPIDEMIOLOGY



EBV infections occur worldwide. These infections are most common in early childhood, with a second peak during late adolescence. By adulthood, more than 90% of individuals have been infected and have antibodies to the virus. IM is usually a disease of young adults. In lower socioeconomic groups and in areas of the world with deficient standards of hygiene (e.g., developing regions), EBV tends to infect children at an early age, and IM is uncommon. In areas with higher standards of hygiene, infection with EBV is often delayed until adulthood, and IM is more prevalent.

EBV is spread by contact with oral secretions. The virus is frequently transmitted from asymptomatic adults to infants and among young adults by transfer of saliva during kissing. Transmission by less intimate contact is rare. EBV has been transmitted by blood transfusion and by bone marrow transplantation. More than 90% of asymptomatic seropositive individuals shed the virus in oropharyngeal secretions. Shedding is increased in immunocompromised patients and those with IM.

PATHOGENESIS

EBV is transmitted by salivary secretions. The virus infects the epithelium of the oropharynx and the salivary glands and is shed from these cells. While B cells may become infected after contact with epithelial cells, studies suggest that lymphocytes in the tonsillar crypts can be infected directly. The virus then spreads through the bloodstream. The proliferation and expansion of EBV-infected B cells along with reactive T cells during IM result in enlargement of lymphoid tissue. Polyclonal activation of B cells leads to the production of antibodies to host-cell and viral proteins. During the acute phase of IM, up to 1 in every 100 B cells in the peripheral blood is infected by EBV; after recovery, 1–50 in every 1 million B cells is infected. During IM, there is an inverted CD4+/CD8+ T cell ratio. The percentage of CD4+ T cells decreases, while there are large clonal expansions of CD8+ T cells; up to 40% of CD8+ T cells are directed against EBV antigens during acute infection. Memory B cells, not epithelial cells, are the reservoir for EBV in the body. When patients are treated with acyclovir, shedding of EBV from the oropharynx stops but the virus persists in B cells.

The EBV receptor (CD21) on the surface of B cells is also the receptor for the C3d component of complement. EBV infection of epithelial cells results in viral replication and production of virions. When B cells are infected by EBV in vitro, they become transformed and can proliferate indefinitely. During latent infection of B cells, only the EBV nuclear antigens (EBNAs), latent membrane proteins (LMPs), and small EBV RNAs (EBERs) are expressed in vitro. EBV-transformed B cells secrete immunoglobulin; only a small fraction of these cells produce virus.

Cellular immunity is more important than humoral immunity in controlling EBV infection. In the initial phase of infection, suppressor T cells, natural killer cells, and nonspecific cytotoxic T cells are important in controlling the proliferation of EBV-infected B cells. Levels of markers of T cell activation and serum interferon γ are elevated. Later in infection, human leukocyte antigen-restricted cytotoxic T cells that recognize EBNAs and LMPs and destroy EBV-infected cells are generated.

If T cell immunity is compromised, EBV-infected B cells may begin to proliferate. When EBV is associated with lymphoma in immunocompetent persons, virus-induced proliferation is but one step in a multistep process of neoplastic transformation. In many EBV-containing tumors, LMP-1 mimics members of the tumor necrosis factor receptor family (e.g., CD40), transmitting growth-proliferating signals.

CLINICAL MANIFESTATIONS

Signs and Symptoms Most EBV infections in infants and young children either are asymptomatic or present as mild pharyngitis with or without tonsillitis. In contrast, ~75% of infections in adolescents present as IM. IM in the elderly often presents with nonspecific symptoms, including prolonged fever, fatigue, myalgia, and malaise. In contrast, pharyngitis, lymphadenopathy, splenomegaly, and atypical lymphocytes are relatively rare in elderly patients.

TABLE 218-1 SIGNS AND SYMPTOMS OF INFECTIOUS MONONUCLEOSIS

Manifestation	Median Percentage of Patients (Range)
Symptoms	
Sore throat	75 (50–87)
Malaise	47 (42–76)
Headache	38 (22–67)
Abdominal pain, nausea, or vomiting	17 (5–25)
Chills	10 (9–11)
Signs	
Lymphadenopathy	95 (83–100)
Fever	93 (60–100)
Pharyngitis or tonsillitis	82 (68–90)
Splenomegaly	51 (43–64)
Hepatomegaly	11 (6–15)
Rash	10 (0–25)
Periorbital edema	13 (2–34)
Palatal enanthem	7 (3–13)
Jaundice	5 (2–10)

The incubation period for IM in young adults is ~4–6 weeks. A prodrome of fatigue, malaise, and myalgia may last for 1–2 weeks before the onset of fever, sore throat, and lymphadenopathy. Fever is usually low-grade and is most common in the first 2 weeks of the illness; however, it may persist for >1 month. Common signs and symptoms are listed along with their frequencies in [Table 218-1](#). Lymphadenopathy and pharyngitis are most prominent during the first 2 weeks of the illness, while splenomegaly is more prominent during the second and third weeks. Lymphadenopathy most often affects the posterior cervical nodes but may be generalized. Enlarged lymph nodes are frequently tender and symmetric but are not fixed in place. Pharyngitis, often the most prominent sign, can be accompanied by enlargement of the tonsils with an exudate resembling that of streptococcal pharyngitis. A morbilliform or papular rash, usually on the arms or trunk, develops in ~5% of cases ([Fig. 218-1](#)). Many patients treated with ampicillin develop a macular rash; this rash is not predictive of future adverse reactions to penicillins. Erythema nodosum and erythema multiforme also have been described ([Chap. 72](#)). The severity of the disease correlates with the levels of CD8+ T cells and EBV DNA in the blood. Most patients have symptoms for 2–4 weeks, but nearly 10% have fatigue that persists for ≥ 6 months.

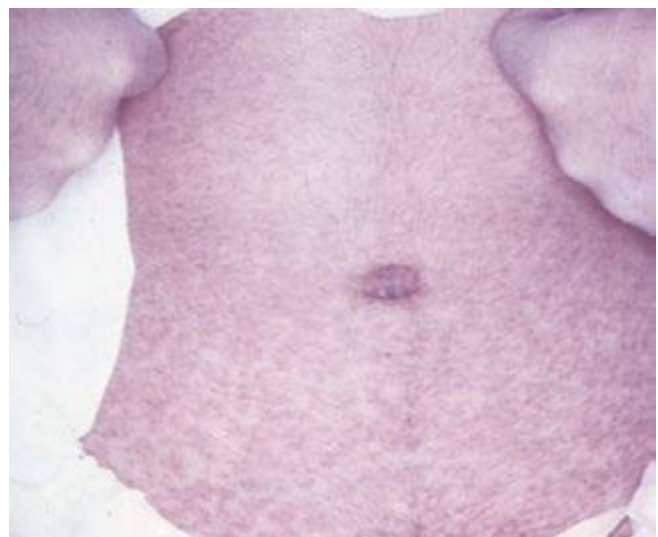


FIGURE 218-1 Rash in a patient with infectious mononucleosis due to Epstein-Barr virus. (Courtesy of Maria Turner, MD; with permission.)

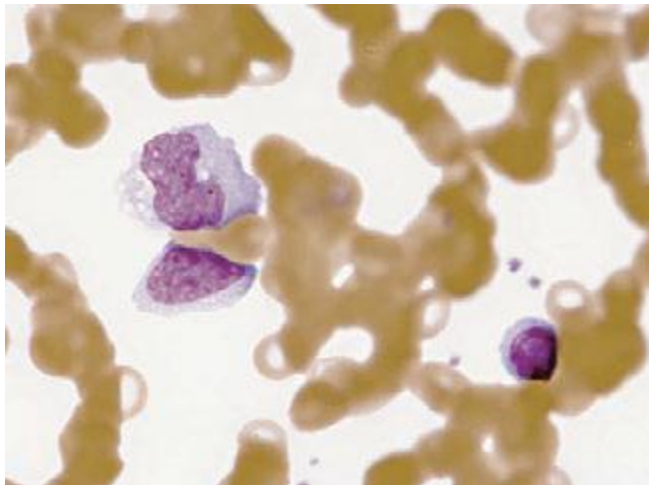


FIGURE 218-2 Atypical lymphocytes from a patient with infectious mononucleosis due to Epstein-Barr virus.

Laboratory Findings The white blood cell count is usually elevated and peaks at 10,000–20,000/ μL during the second or third week of illness. Lymphocytosis is usually demonstrable, with >10% atypical lymphocytes. The latter cells are enlarged lymphocytes that have abundant cytoplasm, vacuoles, and indentations of the cell membrane (Fig. 218-2). CD8⁺ cells predominate among the atypical lymphocytes. Low-grade neutropenia and thrombocytopenia are common during the first month of illness. Liver function is abnormal in >90% of cases. Serum levels of aminotransferases and alkaline phosphatase are usually mildly elevated. The serum concentration of bilirubin is elevated in ~40% of cases.

Complications Most cases of IM are self-limited. Deaths are very rare and are most often due to central nervous system (CNS) complications, splenic rupture, upper airway obstruction, or bacterial superinfection.

When CNS complications develop, they usually do so during the first 2 weeks of EBV infection; in some patients, especially children, they are the only clinical manifestations of IM. Heterophile antibodies and atypical lymphocytes may be absent. Meningitis and encephalitis are the most common neurologic abnormalities, and patients may present with headache, meningismus, or cerebellar ataxia. Acute hemiplegia and psychosis also have been described. The cerebrospinal fluid contains mainly lymphocytes, with occasional atypical lymphocytes. Most cases resolve without neurologic sequelae. Acute EBV infection has also been associated with cranial nerve palsies (especially those involving cranial nerve VII), Guillain-Barré syndrome, acute transverse myelitis, and peripheral neuritis.

Autoimmune hemolytic anemia occurs in ~2% of cases during the first 2 weeks. In most cases, the anemia is Coombs-positive, with cold agglutinins directed against the red blood cell antigen. Most patients with hemolysis have mild anemia that lasts for 1–2 months, but some patients have severe disease with hemoglobinuria and jaundice. Nonspecific antibody responses may also include rheumatoid factor, antinuclear antibodies, anti-smooth muscle antibodies, antiplatelet antibodies, and cryoglobulins. IM has been associated with red-cell aplasia, severe granulocytopenia, thrombocytopenia, pancytopenia, and hemophagocytic lymphohistiocytosis. The spleen ruptures in <0.5% of cases. Splenic rupture is more common among male than female patients and may manifest as abdominal pain, referred shoulder pain, or hemodynamic compromise.

Hypertrophy of lymphoid tissue in the tonsils or adenoids can result in upper airway obstruction, as can inflammation and edema of the epiglottis, pharynx, or uvula. About 10% of patients with IM develop streptococcal pharyngitis after their initial sore throat resolves.

Other rare complications associated with acute EBV infection include hepatitis (which can be fulminant), myocarditis or pericarditis, pneumonia with pleural effusion, interstitial nephritis, genital ulcerations, and vasculitis.

EBV-Associated Diseases Other Than IM EBV-associated lymphoproliferative disease has been described in patients with congenital or acquired immunodeficiency, including those with severe combined immunodeficiency, patients with AIDS, and recipients of bone marrow or organ transplants who are receiving immunosuppressive drugs (especially cyclosporine). Proliferating EBV-infected B cells infiltrate lymph nodes and multiple organs, and patients present with fever and lymphadenopathy or gastrointestinal symptoms. Pathologic studies show B cell hyperplasia or poly- or monoclonal lymphoma.

X-linked lymphoproliferative disease is a recessive disorder of young boys who have a normal response to childhood infections but develop fatal lymphoproliferative disorders after infection with EBV. The protein associated with most cases of this syndrome (SAP) binds to a protein that mediates interactions of B and T cells. Most patients with this syndrome die of acute IM. Others develop hypogammaglobulinemia, malignant B cell lymphomas, aplastic anemia, or agranulocytosis. Disease resembling X-linked lymphoproliferative disease has also been associated with mutations in XIAP. Mutations in ITK, MagT1, or CD27 are associated with inability to control EBV and lymphoma. Moreover, IM has proved fatal to some patients with no obvious preexisting immune abnormality.

Oral hairy leukoplakia (Fig. 218-3) is an early manifestation of infection with HIV in adults (Chap. 226). Most patients present with raised, white corrugated lesions on the tongue (and occasionally on the buccal mucosa) that contain EBV DNA. Children infected with HIV can develop lymphoid interstitial pneumonitis; EBV DNA is often found in lung tissue from these patients.

Patients with chronic fatigue syndrome may have titers of antibody to EBV that are elevated but are not significantly different from those in healthy EBV-seropositive adults. While some patients have malaise and fatigue that persist for weeks or months after IM, persistent EBV infection is not a cause of chronic fatigue syndrome. Chronic active EBV infection is very rare and is distinct from chronic fatigue syndrome. The affected patients have an illness lasting >6 months, with elevated levels of EBV DNA in the blood, high titers of antibody to EBV, and evidence of organ involvement, including hepatosplenomegaly, lymphadenopathy, and pneumonitis, uveitis, or neurologic disease.

EBV is associated with several malignancies. About 15% of cases of Burkitt's lymphoma in the United States and ~90% of those in Africa are associated with EBV (Chap. 134). African patients with Burkitt's lymphoma have high levels of antibody to EBV, and their tumor tissue usually contains viral DNA. Malaria in African patients may impair cellular immunity to EBV and induce polyclonal B cell activation with an expansion of EBV-infected B cells. These changes may enhance the proliferation of B cells with elevated EBV DNA in the bloodstream, thereby increasing the likelihood of a *c-myc* translocation—the hallmark of Burkitt's lymphoma. EBV-containing Burkitt's lymphoma also occurs in patients with AIDS.

Anaplastic nasopharyngeal carcinoma is common in southern China and is uniformly associated with EBV; the affected tissues



FIGURE 218-3 Oral hairy leukoplakia often presents as white plaques on the lateral surface of the tongue and is associated with Epstein-Barr virus infection.

contain viral DNA and antigens. Patients with nasopharyngeal carcinoma often have elevated titers of antibody to EBV (Chap. 106). High levels of EBV plasma DNA before treatment or detectable levels of EBV DNA after radiation therapy correlate with lower rates of overall survival and relapse-free survival among patients with nasopharyngeal carcinoma.

Worldwide, the most common EBV-associated malignancy is gastric carcinoma. About 9% of these tumors are EBV-positive.

EBV has been associated with Hodgkin's disease, especially the mixed-cellularity type (Chap. 134). Patients with Hodgkin's disease often have elevated titers of antibody to EBV. In about half of cases in the United States, viral DNA and antigens are found in Reed-Sternberg cells. The risk of EBV-positive Hodgkin's disease is significantly increased in young adults for several years after EBV-seropositive IM. About 50% of non-Hodgkin's lymphomas in patients with AIDS are EBV-positive.

EBV is present in B cells of lesions from patients with lymphomatoid granulomatosis. In some cases, EBV DNA has been detected in tumors from immunocompetent patients with angiocentric nasal NK/T cell lymphoma, T cell lymphoma, and CNS lymphoma. Studies have demonstrated viral DNA in leiomyosarcomas from AIDS patients and in smooth-muscle tumors from organ transplant recipients. Virtually all CNS lymphomas in AIDS patients are associated with EBV. Studies have found that a history of IM and higher levels of antibodies to EBV before the onset of disease is more common in persons with multiple sclerosis than in the general population; additional research on a possible causal relationship is needed.

DIAGNOSIS

Serologic Testing (Fig. 218-4) The heterophile test is used for the diagnosis of IM in children and adults. In the test for this antibody, human serum is absorbed with guinea pig kidney, and the heterophile titer is defined as the greatest serum dilution that agglutinates sheep, horse, or cow erythrocytes. The heterophile antibody does not interact with EBV proteins. A titer of ≥ 40 is diagnostic of acute EBV infection in a patient who has symptoms compatible with IM and atypical lymphocytes. Tests for heterophile antibodies are positive in 40% of patients with IM during the first week of illness and in 80–90% during the third week. Therefore, repeated testing may be necessary, especially if the initial test is performed early. Tests usually remain positive for 3 months after the onset of illness, but heterophile antibodies can persist for up to 1 year. These antibodies usually are not detectable in children < 5 years of age, in the elderly, or in patients presenting with symptoms not typical of IM. The commercially available monospot test for heterophile antibodies is somewhat more sensitive than the classic heterophile test. The monospot test is $\sim 75\%$ sensitive and $\sim 90\%$ specific compared with EBV-specific serologies (see below). False-positive monospot results are more common among persons with connective tissue disease, lymphoma, viral hepatitis, and malaria.

EBV-specific antibody testing is used for patients with suspected acute EBV infection who lack heterophile antibodies and for patients with atypical infections. Titers of IgM and IgG antibodies to viral capsid antigen (VCA) are elevated in the serum of more than 90% of patients at the onset of disease. IgM antibody to VCA is most useful

for the diagnosis of acute IM because it is present at elevated titers only during the first 2–3 months of the disease; in contrast, IgG antibody to VCA is usually not useful for diagnosis of IM but is often used to assess past exposure to EBV because it persists for life. Seroconversion to EBNA positivity also is useful for the diagnosis of acute infection with EBV. Antibodies to EBNA become detectable relatively late (3–6 weeks after the onset of symptoms) in nearly all cases of acute EBV infection and persist for the lifetime of the patient. These antibodies may be lacking in immunodeficient patients and in those with chronic active EBV infection.

Titers of other antibodies also may be elevated in IM; however, these elevations are less useful for diagnosis. Antibodies to early antigens are detectable 3–4 weeks after the onset of symptoms in patients with IM. About 70% of individuals with IM have early antigen diffuse (EA-D) antibodies during the illness; the presence of EA-D antibodies is especially likely in patients with relatively severe disease. These antibodies usually persist for only 3–6 months. Levels of EA-D antibodies are also elevated in patients with nasopharyngeal carcinoma or chronic active EBV infection. Early antigen restricted (EA-R) antibodies are only occasionally detected in patients with IM but are often found at elevated titers in patients with African Burkitt's lymphoma or chronic active EBV infection. IgA antibodies to EBV antigens have proved useful for the identification of patients with nasopharyngeal carcinoma and of persons at high risk for the disease.

Other Studies Detection of EBV DNA, RNA, or proteins has been valuable in demonstrating the association of the virus with various malignancies. The polymerase chain reaction has been used to detect EBV DNA in the cerebrospinal fluid of some AIDS patients with lymphomas and to monitor the amount of EBV DNA in the blood of patients with lymphoproliferative disease. Detection of high levels of EBV DNA in blood for a few days to several weeks after the onset of IM may be useful if serologic studies yield equivocal results. Culture of EBV from throat washings or blood is not helpful in the diagnosis of acute infection, since EBV persists in the oropharynx and in B cells for the lifetime of the infected individual.

Differential Diagnosis Whereas $\sim 90\%$ of cases of IM are due to EBV, 5–10% of cases are due to cytomegalovirus (CMV) (Chap. 219). CMV is the most common cause of heterophile-negative mononucleosis; less common causes of IM and differences from IM due to EBV are shown in Table 218-2.

TREATMENT EBV-ASSOCIATED DISEASE

Therapy for IM consists of supportive measures, with rest and analgesia. Excessive physical activity during the first month should be avoided to reduce the possibility of splenic rupture, which often necessitates splenectomy. Glucocorticoid therapy is not indicated for uncomplicated IM and in fact may predispose to bacterial superinfection. Prednisone (40–60 mg/d for 2–3 days, with subsequent tapering of the dose over 1–2 weeks) has been used for the prevention of airway obstruction in patients with severe tonsillar hypertrophy, for

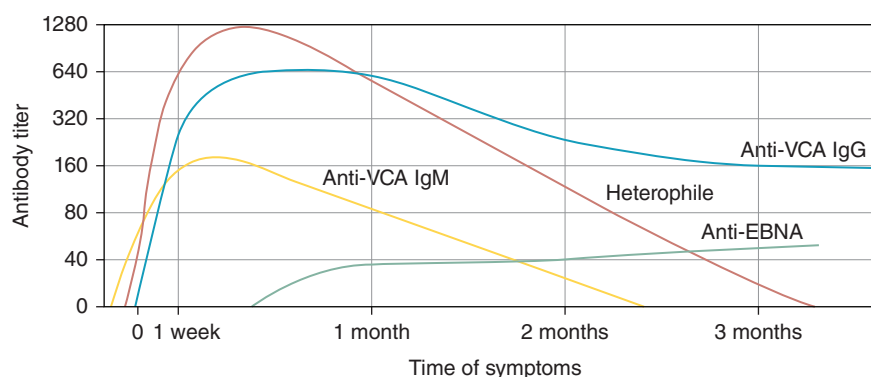


FIGURE 218-4 Pattern of Epstein-Barr virus (EBV) serology during acute infection. EBNA, Epstein-Barr nuclear antigen; VCA, viral capsid antigen. (From JI Cohen, in NS Young et al [eds]: *Clinical Hematology*. Philadelphia, Mosby, 2006.)

TABLE 218-2 DIFFERENTIAL DIAGNOSIS OF INFECTIOUS MONONUCLEOSIS

Etiology	Sign or Symptom				Differences from EBV Mononucleosis
	Fever	Adenopathy	Sore Throat	Atypical Lymphocytes	
EBV infection	+	+	+	+	—
CMV infection	+	±	±	+	Older age at presentation, longer duration of fever
HIV infection	+	+	+	±	Diffuse rash, oral/genital ulcers, aseptic meningitis
Toxoplasmosis	+	+	±	±	Less splenomegaly, exposure to cats or raw meat
HHV-6	+	+	+	+	Older age at presentation
Streptococcal pharyngitis	+	+	+	—	No splenomegaly, less fatigue
Viral hepatitis	+	±	—	±	Higher aminotransferase levels
Rubella	+	+	±	±	Maculopapular rash, no splenomegaly
Lymphoma	+	+	+	+	Fixed, nontender lymph nodes
Drugs ^a	+	+	—	±	Occurs at any age

^aMost commonly phenytoin, carbamazepine, sulfonamides, or minocycline.

Abbreviations: CMV, cytomegalovirus; EBV, Epstein-Barr virus; HHV, human herpesvirus.

autoimmune hemolytic anemia, for hemophagocytic lymphohistiocytosis, and for severe thrombocytopenia. Glucocorticoids have also been administered to rare patients with severe malaise and fever and to patients with severe CNS or cardiac disease.

Acyclovir has had no significant clinical impact on IM in controlled trials. In one study, the combination of acyclovir and prednisolone had no significant effect on the duration of symptoms of IM.

Acyclovir, at a dosage of 400–800 mg five times daily, has been effective for the treatment of oral hairy leukoplakia (despite common relapses). The posttransplantation EBV lymphoproliferative syndrome (Chap. 169) generally does not respond to antiviral therapy. When possible, therapy should be directed toward reduction of immunosuppression. Antibody to CD20 (rituximab) has been effective in some cases. Infusions of donor lymphocytes are often effective for stem cell transplant recipients, although graft-versus-host disease can occur. Infusions of EBV-specific cytotoxic T cells have been used to prevent EBV lymphoproliferative disease in high-risk settings as well as to treat the disease. Interferon α administration, cytotoxic chemotherapy, and radiation therapy (especially for CNS lesions) also have been used. Infusion of autologous EBV-specific cytotoxic T lymphocytes has shown promise in small studies of patients with nasopharyngeal carcinoma and Hodgkin's disease. Treatment of several cases of X-linked lymphoproliferative disease with antibody to CD20 resulted in a successful outcome of what otherwise would probably have been fatal acute EBV infection.

PREVENTION

The isolation of patients with IM is unnecessary. A vaccine directed against the major EBV glycoprotein reduced the frequency of IM but did not affect the rate of asymptomatic infection in a phase 2 trial.

219 Cytomegalovirus and Human Herpesvirus Types 6, 7, and 8

Camille Nelson Kotton, Martin S. Hirsch

CYTOMEGALOVIRUS


DEFINITION

Cytomegalovirus (CMV), which was initially isolated from patients with congenital cytomegalic inclusion disease, is now recognized as an important pathogen in all age groups. In addition to inducing severe birth defects, CMV causes a wide spectrum of disorders in older children and adults, ranging from an asymptomatic subclinical infection

to a mononucleosis syndrome in healthy individuals to disseminated disease in immunocompromised patients. Human CMV is one of several related species-specific viruses that cause similar diseases in various animals. All are associated with the production of characteristic enlarged cells—hence the name *cytomegalovirus*.

CMV, a β -herpesvirus, has double-stranded DNA, four species of mRNA, a protein capsid, and a lipoprotein envelope. Like other herpesviruses, CMV demonstrates icosahedral symmetry, replicates in the cell nucleus, and can cause either a lytic and productive or a latent infection. CMV can be distinguished from other herpesviruses by certain biologic properties, such as host range and type of cytopathology. Viral replication is associated with the production of large intranuclear inclusions and smaller cytoplasmic inclusions. CMV appears to replicate in a variety of cell types in vivo; in tissue culture it grows preferentially in fibroblasts. Although there is little evidence that CMV is oncogenic in vivo, it does transform fibroblasts in rare instances, and genomic transforming fragments have been identified.

EPIDEMIOLOGY

 CMV has a worldwide distribution. In many regions of the world, the vast majority of adults are seropositive for CMV, whereas only half of adults in the United States and Canada are seropositive. In regions where the prevalence of CMV antibody is high, immunocompromised adults are more likely to undergo reactivation disease rather than primary infection. Data generated in specific regions should be considered in the context of local seropositivity rates, when appropriate.

Of newborns in the United States, ~1% are infected with CMV; the percentages are higher in many less-developed countries. Communal living and poor personal hygiene facilitate spread. Perinatal and early childhood infections are common. CMV may be present in breast milk, saliva, feces, and urine. Transmission has occurred among young children in day-care centers and has been traced from infected toddler to pregnant mother to developing fetus. When an infected child introduces CMV into a household, 50% of susceptible family members seroconvert within 6 months.

CMV is not readily spread by casual contact but rather requires repeated or prolonged intimate exposure for transmission. In late adolescence and young adulthood, CMV is often transmitted sexually, and asymptomatic carriage in semen or cervical secretions is common. Antibody to CMV is present at detectable levels in a high proportion of sexually active men and women, who may harbor several strains simultaneously. Transfusion of blood products containing viable leukocytes may transmit CMV, with a frequency of 0.14–10% per unit transfused. Transfusion of leukocyte-reduced or CMV-seronegative blood significantly decreases the risk of CMV transmission.

Once infected, an individual generally carries CMV for life. The infection usually remains silent. CMV reactivation syndromes develop more frequently, however, when T lymphocyte-mediated immunity

is compromised—for example, after organ transplantation, with lymphoid neoplasms and certain acquired immunodeficiencies (in particular, HIV infection; **Chap. 226**), or during critical illness in intensive care units. Most primary CMV infections in organ transplant recipients (**Chap. 169**) result from transmission via the graft. In CMV-seropositive transplant recipients, infection results from reactivation of latent virus or from infection by a new strain. CMV infection may also be associated with diseases as diverse as coronary artery stenosis and malignant gliomas, but these associations require further validation.

PATHOGENESIS

Congenital CMV infection can result from either primary or reactivation infection of the mother. However, clinical disease in the fetus or newborn is related almost exclusively to primary maternal infection (**Table 219-1**). The factors determining the severity of congenital infection are unknown; a deficient capacity to produce precipitating antibodies and to mount T cell responses to CMV is associated with relatively severe disease.

Primary infection with CMV in late childhood or adulthood is often associated with a vigorous T lymphocyte response that may contribute to the development of a mononucleosis syndrome similar to that which follows infection with Epstein-Barr virus (**Chap. 218**). The hallmark of such infection is the appearance of atypical lymphocytes in the peripheral blood; these cells are predominantly activated CD8+ T lymphocytes. Polyclonal activation of B cells by CMV contributes to the development of rheumatoid factors and other autoantibodies during mononucleosis.

Once acquired, CMV persists indefinitely in host tissues. The sites of persistent infection probably include multiple cell types and various organs. Transmission via blood transfusion or organ transplantation is due primarily to latent infections in these tissues. If the host's T cell responses become compromised by disease or by iatrogenic immunosuppression, latent virus can reactivate to cause a variety of syndromes. Chronic antigenic stimulation in the presence of immunosuppression (for example, after organ transplantation) appears to be an ideal setting for CMV activation and CMV disease. Certain particularly potent suppressants of T cell immunity (e.g., antithymocyte globulin, alemtuzumab) are associated with a high rate of clinical CMV syndromes. CMV may itself contribute to further T lymphocyte hyporesponsiveness, which often precedes superinfection with other opportunistic pathogens such as bacteria, molds, and *Pneumocystis*.

PATHOLOGY

Cytomegalic cells in vivo (presumed to be infected epithelial cells) are two to four times larger than surrounding cells and often contain an 8- to 10- μ m intranuclear inclusion that is eccentrically placed and is surrounded by a clear halo, producing an "owl's eye" appearance. Smaller granular cytoplasmic inclusions are demonstrated occasionally. Cytomegalic cells are found in a wide variety of organs, including the salivary gland, lung, liver, kidney, intestine, pancreas, adrenal gland, and central nervous system.

The cellular inflammatory response to infection consists of plasma cells, lymphocytes, and monocyte-macrophages. Granulomatous reactions occasionally develop, particularly in the liver. Immunopathologic reactions may contribute to CMV disease. Immune complexes have been detected in infected infants, sometimes in association with CMV-related glomerulopathies. Immune-complex glomerulopathy has also been observed in some CMV-infected patients after renal transplantation.

CLINICAL MANIFESTATIONS

Congenital CMV Infection Fetal infections range from subclinical to severe and disseminated. Cytomegalic inclusion disease develops in ~5% of infected fetuses and is seen almost exclusively in infants born to mothers who develop primary infections during pregnancy. Petechiae, hepatosplenomegaly, and jaundice are the most common presenting features (60–80% of cases). Microcephaly with or without cerebral calcifications, intrauterine growth retardation, and prematurity are reported in 30–50% of cases. Inguinal hernias and chorioretinitis are less common. Laboratory abnormalities include elevated alanine aminotransferase levels in serum, thrombocytopenia, conjugated hyperbilirubinemia, hemolysis, and elevated protein levels in cerebrospinal fluid. The prognosis for severely infected infants is poor; the mortality rate is 20–30%, and few survivors escape intellectual or hearing difficulties later in childhood. The differential diagnosis of cytomegalic inclusion disease in infants includes syphilis, rubella, toxoplasmosis, infection with herpes simplex virus or enterovirus, and bacterial sepsis.

Most congenital CMV infections are clinically inapparent at birth. Of asymptotically infected infants, 5–25% develop significant psychomotor, hearing, ocular, or dental abnormalities over the next several years.

Perinatal CMV Infection The newborn may acquire CMV at delivery by passage through an infected birth canal or by postnatal contact with infected breast milk or other maternal secretions. Of infants who are breast-fed for >1 month by seropositive mothers, 40–60% become infected. Iatrogenic transmission can result from blood transfusion; use of leukocyte-reduced or CMV-seronegative blood products for transfusion into low-birth-weight seronegative infants or seronegative pregnant women decreases risk.

The great majority of infants infected at or after delivery remain asymptomatic. However, protracted interstitial pneumonitis has been associated with perinatally acquired CMV infection, particularly in premature infants, and occasionally has been accompanied by infection with *Chlamydia trachomatis*, *Pneumocystis*, or *Ureaplasma urealyticum*. Poor weight gain, adenopathy, rash, hepatitis, anemia, and atypical lymphocytosis may also be found, and CMV excretion often persists for months or years.

CMV Mononucleosis The most common clinical manifestation of CMV infection in immunocompetent hosts beyond the neonatal period is a heterophile antibody-negative mononucleosis syndrome, which may develop spontaneously or follow transfusion of leukocyte-containing blood products. Although the syndrome occurs at all ages, it most

TABLE 219-1 CMV DISEASE IN THE IMMUNOCOMPROMISED HOST

Population	Risk Factors	Principal Syndromes	Treatment	Prevention
Fetus	Primary maternal infection/early pregnancy	Cytomegalic inclusion disease	Ganciclovir for symptomatic neonates	Avoidance of exposure; possibly, maternal treatment with CMV immunoglobulin during pregnancy
Organ transplant recipient	Seropositivity of donor and/or recipient; potent immunosuppressive regimen; treatment of rejection	Febrile leukopenia; gastrointestinal disease; pneumonia	Ganciclovir or valganciclovir	Prophylaxis or preemptive therapy with ganciclovir or valganciclovir
Hematopoietic stem cell transplant recipient	Graft-vs.-host disease; older age of recipient; seropositive recipient; viremia	Pneumonia; gastrointestinal disease	Ganciclovir or valganciclovir or foscarnet, \pm CMV immunoglobulin	Prophylaxis or preemptive therapy with ganciclovir or valganciclovir
Person with AIDS	<100 CD4+ T cells/ μ L; CMV seropositivity	Retinitis; gastrointestinal disease; neurologic disease	Ganciclovir, valganciclovir, foscarnet, or cidofovir	Oral valganciclovir

1192 often involves sexually active young adults. With incubation periods of 20–60 days, the illness generally lasts for 2–6 weeks. Prolonged high fevers, sometimes with chills, profound fatigue, and malaise, characterize this disorder. Myalgias, headache, and splenomegaly are common, but in CMV (as opposed to Epstein-Barr virus) mononucleosis, exudative pharyngitis and cervical lymphadenopathy are rare. Occasional patients develop rubelliform rashes, often after exposure to ampicillin or certain other antibiotics. Less common are interstitial or segmental pneumonia, myocarditis, pleuritis, arthritis, and encephalitis. In rare cases, Guillain-Barré syndrome complicates CMV mononucleosis. The characteristic laboratory abnormality is relative lymphocytosis in peripheral blood, with >10% atypical lymphocytes. Total leukocyte counts may be low, normal, or markedly elevated. Although significant jaundice is uncommon, serum aminotransferase and alkaline phosphatase levels are often moderately elevated. Heterophile antibodies are absent; however, transient immunologic abnormalities are common and may include the presence of cryoglobulins, rheumatoid factors, cold agglutinins, and antinuclear antibodies. Hemolytic anemia, thrombocytopenia, and granulocytopenia complicate recovery in rare instances.

Most patients recover without sequelae, although postviral asthenia may persist for months. The excretion of CMV in urine, genital secretions, and/or saliva often continues for months or years. Rarely, CMV infection is fatal in immunocompetent hosts; survivors can have recurrent episodes of fever and malaise, sometimes associated with autonomic nervous system dysfunction (e.g., attacks of sweating or flushing).

CMV Infection in the Immunocompromised Host (Table 219-1) CMV is the viral pathogen most commonly complicating organ transplantation (Chap. 169). In recipients of kidney, heart, lung, liver, pancreas, and vascularized composite (hand, face, other) transplants, CMV induces a variety of syndromes, including fever and leukopenia, hepatitis, colitis, pneumonitis, esophagitis, gastritis, and retinitis. CMV disease is an independent risk factor for both graft loss and death. Without prophylaxis, the period of maximal risk is between 1 and 4 months after transplantation. Disease likelihood and viral replication levels generally are greater after primary infection than after reactivation. Molecular studies indicate that seropositive transplant recipients are susceptible to infection with donor-derived, genotypically variant CMV, and such infection often results in disease. Reactivation infection, although common, is less likely than primary infection to be important clinically. The risk of clinical disease is related to various factors, such as degree of immunosuppression, use of antilymphocyte antibodies, lack of anti-CMV prophylaxis, and co-infection with other pathogens. The transplanted organ is particularly vulnerable as a target for CMV infection; thus there is a tendency for CMV hepatitis to follow liver transplantation and for CMV pneumonitis to follow lung transplantation.

CMV viremia occurs in roughly one-third of hematopoietic stem cell transplant recipients; the risk of severe disease may be reduced by prophylaxis or preemptive therapy with antiviral drugs. The risk is greatest 5–13 weeks after transplantation, and identified risk factors include certain types of immunosuppressive therapy, an allogeneic (rather than an autologous) graft, acute graft-versus-host disease, older age, and pretransplantation recipient seropositivity.

CMV is an important pathogen in patients with advanced HIV infection (Chap. 226), in whom it may cause retinitis or disseminated disease, particularly when peripheral-blood CD4+ T cell counts fall below 50–100/ μ L. As treatment for underlying HIV infection has improved, the incidence of serious CMV infections (e.g., retinitis) has decreased. However, during the first few weeks after institution of highly active antiretroviral therapy, acute flare-ups of CMV retinitis may occur secondary to an immune reconstitution inflammatory syndrome.

Syndromes produced by CMV in immunocompromised hosts often begin with prolonged fatigue, fever, malaise, anorexia, night sweats, and arthralgias or myalgias. Liver function abnormalities, leukopenia, thrombocytopenia, and atypical lymphocytosis may be observed during these episodes. The development of tachypnea, hypoxemia,

and unproductive cough signals respiratory involvement. Radiologic examination of the lung often shows bilateral interstitial or reticulonodular infiltrates that begin in the periphery of the lower lobes and spread centrally and superiorly; localized segmental, nodular, or alveolar patterns are less common. The differential diagnosis includes *Pneumocystis* infection; other viral, bacterial, or fungal infections; pulmonary hemorrhage; and injury secondary to irradiation or to treatment with cytotoxic drugs.

Gastrointestinal CMV involvement may be localized or extensive and almost exclusively affects immunocompromised hosts. Colitis is the most common clinical manifestation in organ transplant recipients. Ulcers of the esophagus, stomach, small intestine, or colon may result in bleeding or perforation. CMV infection may lead to exacerbations of underlying ulcerative colitis. Hepatitis occurs frequently, particularly after liver transplantation. Acalculous cholecystitis and adrenalitis also have been described.

CMV rarely causes meningoencephalitis in otherwise healthy individuals. Two forms of CMV encephalitis are seen in patients with AIDS. One resembles HIV encephalitis and presents as progressive dementia; the other is a ventriculoencephalitis characterized by cranial-nerve deficits, nystagmus, disorientation, lethargy, and ventriculomegaly. In immunocompromised patients, CMV can also cause subacute progressive polyradiculopathy, which is often reversible if recognized and treated promptly.

CMV retinitis is an important cause of blindness in immunocompromised patients, particularly patients with advanced AIDS (Chap. 226). Early lesions consist of small, opaque, white areas of granular retinal necrosis that spread in a centrifugal manner and are later accompanied by hemorrhages, vessel sheathing, and retinal edema (Fig. 219-1). CMV retinopathy must be distinguished from that due to other conditions, including toxoplasmosis, candidiasis, and herpes simplex virus infection.

Fatal CMV infections are often associated with persistent viremia and the involvement of multiple organ systems. Progressive pulmonary infiltrates, pancytopenia, hyperamylasemia, and hypotension are characteristic features that are frequently found in conjunction with a terminal bacterial, fungal, or protozoan superinfection. Extensive adrenal necrosis with CMV inclusions is often documented at autopsy, as is CMV involvement of many other organs.

DIAGNOSIS

CMV infection usually cannot be diagnosed reliably on clinical grounds alone. Isolation of CMV or detection of its antigens or DNA in appropriate clinical specimens is the preferred approach. The most common method of detection is quantitative nucleic acid testing (QNAT) for CMV by polymerase chain reaction (PCR) technology,



FIGURE 219-1 Cytomegalovirus infection in a patient with AIDS may appear as an arcuate zone of retinitis with hemorrhages and optic disk swelling. Often CMV is confined to the retinal periphery, beyond view of the direct ophthalmoscope.

for which blood or other specimens can be used; some centers use a CMV antigenemia test, an immunofluorescence assay that detects CMV antigens (pp65) in peripheral-blood leukocytes. Such assays may yield a positive result several days earlier than culture methods. QNAT may predict the risk for disease progression, particularly in immunocompromised hosts. CMV DNA in cerebrospinal fluid is useful in the diagnosis of CMV encephalitis or polyradiculopathy. Considerable variation exists among assays and laboratories; a recently introduced international testing standard should help reduce variation in PCR test results.

Virus excretion or viremia is readily detected by culture of appropriate specimens on human fibroblast monolayers. If CMV titers are high, as is common in congenital disseminated infection and in AIDS, characteristic cytopathic effects may be detected within a few days. However, in some situations (e.g., CMV mononucleosis), viral titers are low, and cytopathic effects may take several weeks to appear. Many laboratories expedite diagnosis with an overnight tissue-culture method (shell vial assay) involving centrifugation and an immunocytochemical detection technique employing monoclonal antibodies to an immediate-early CMV antigen. Isolation of virus from urine or saliva does not, by itself, constitute proof of acute infection, since excretion from these sites may continue for months or years after illness. Detection of viremia is a better predictor of acute infection.

A variety of serologic assays detect antibody to CMV. An increased level of IgG antibody to CMV may not be detectable for up to 4 weeks after primary infection. Detection of CMV-specific IgM is sometimes useful in the diagnosis of recent or active infection; however, circulating rheumatoid factors may result in occasional false-positive IgM tests. Serology is especially helpful when used to predict risk of CMV infection and disease in transplant recipients.

PREVENTION

Prevention of CMV in organ and hematopoietic stem cell transplant recipients is usually based on one of two methods: universal prophylaxis or preemptive therapy. With universal prophylaxis, antiviral drugs are used for a defined period, often 3 or 6 months. One clinical trial demonstrated that, in CMV-seronegative recipients with seropositive donors, prophylaxis was more effective at prevention when given for 200 days rather than 100 days. With preemptive therapy, patients are monitored weekly for CMV viremia, and antiviral treatment is initiated once viremia is detected. Because of the bone marrow-suppressive effects of universal prophylaxis, preemptive therapy is more commonly employed in hematopoietic stem cell transplant recipients. For patients with advanced HIV infection (CD4+ T cell counts of <50/μL), some experts have advocated prophylaxis with valganciclovir (see below). However, side effects, lack of proven benefit, possible induction of viral resistance, and high cost have precluded the wide acceptance of this practice. Preemptive therapy is under study in HIV-infected patients.

Several additional measures are useful for the prevention of CMV transmission to CMV-naïve, high-risk patients. The use of CMV-seronegative or leukocyte-depleted blood greatly decreases the rate of transfusion-associated transmission. In a placebo-controlled trial, a CMV glycoprotein B vaccine reduced infection rates among 464 CMV-seronegative women; this outcome raises the possibility that this experimental vaccine will reduce rates of congenital infection, but further studies must validate this approach. A CMV glycoprotein B vaccine with MF59 adjuvant appeared effective in reducing the risk and duration of viremia in both seropositive and seronegative renal transplant recipients at risk for CMV infection. CMV immune globulin has been reported to prevent congenital CMV infection in infants of women with primary infection during pregnancy. Studies in hematopoietic stem cell transplant recipients have produced conflicting results.

Prophylactic acyclovir or valacyclovir may reduce rates of CMV infection and disease in renal transplant recipients, although neither drug is effective in the treatment of active CMV disease.

Ganciclovir is a guanosine derivative that has considerably more activity against CMV than its congener acyclovir. After intracellular conversion by a viral phosphotransferase encoded by CMV gene region UL97, ganciclovir triphosphate is a selective inhibitor of CMV DNA polymerase. Several clinical studies have indicated response rates of 70–90% among patients with AIDS who are given ganciclovir for the treatment of CMV retinitis or colitis. In severe infections (e.g., CMV pneumonia in hematopoietic stem cell transplant recipients), ganciclovir is often combined with CMV immune globulin. Prophylactic or suppressive ganciclovir may be useful in high-risk hematopoietic stem cell or organ transplant recipients (e.g., those who are CMV-seropositive before transplantation). In many patients with AIDS, persistently low CD4+ T cell counts, and CMV disease, clinical and virologic relapses occur promptly if treatment with ganciclovir is discontinued. Therefore, prolonged maintenance regimens are recommended for such patients. Resistance to ganciclovir is more common among patients treated for >3 months and is usually related to mutations in the CMV UL97 gene (or, less commonly, the UL54 gene).

Valganciclovir is an orally bioavailable prodrug that is rapidly metabolized to ganciclovir in intestinal tissues and the liver. Approximately 60–70% of an oral dose of valganciclovir is absorbed. An oral valganciclovir dose of 900 mg results in ganciclovir blood levels similar to those obtained with an IV ganciclovir dose of 5 mg/kg. Valganciclovir appears to be as effective as IV ganciclovir for both CMV induction (treatment) and maintenance regimens, while providing the ease of oral dosing. Furthermore, the adverse-event profiles and rates of resistance for the two drugs are similar.

Ganciclovir or valganciclovir therapy for CMV disease consists of a 14- to 21-day induction course (5 mg/kg IV twice daily for ganciclovir or 900 mg PO twice daily for valganciclovir), sometimes followed by maintenance therapy (e.g., valganciclovir, 900 mg/d). Peripheral-blood neutropenia develops in roughly one-quarter of treated patients but may be ameliorated by granulocyte colony-stimulating factor or granulocyte-macrophage colony-stimulating factor. Whether to use maintenance therapy should depend on the overall level of immunocompromise and the risk of recurrent disease. Discontinuation of maintenance therapy should be considered in patients with AIDS who, while receiving antiretroviral therapy, have a sustained (3- to 6-month) increase in CD4+ T cell counts to >100/μL.

For treatment of CMV retinitis, ganciclovir may also be administered via a slow-release pellet sutured into the eye. Although this intraocular device provides good local protection, contralateral eye disease and disseminated disease are not affected, and early retinal detachment is possible. A combination of intraocular and systemic therapy may be better than the intraocular implant alone.

Foscarnet (sodium phosphonoformate) inhibits CMV DNA polymerase. Because this agent does not require phosphorylation to be active, it is also effective against most ganciclovir-resistant isolates. Foscarnet is less well tolerated than ganciclovir and causes considerable toxicity, including renal dysfunction, hypomagnesemia, hypokalemia, hypocalcemia, genital ulcers, dysuria, nausea, and paresthesia. Moreover, foscarnet administration requires the use of an infusion pump and close clinical monitoring. With aggressive hydration and dose adjustments for renal dysfunction, the toxicity of foscarnet can be reduced. The use of foscarnet should be avoided when a saline load cannot be tolerated (e.g., in cardiomyopathy). The approved induction regimen is 60 mg/kg every 8 h for 2 weeks, although 90 mg/kg every 12 h is equally effective and no more toxic. Maintenance infusions should deliver 90–120 mg/kg once daily. No oral preparation is available. Foscarnet-resistant virus may emerge during extended therapy. This drug is used more frequently after hematopoietic stem cell transplantation than in other situations to avoid the myelosuppressive effects of ganciclovir; in

1194 general, foscarnet is also the first choice for infections with ganciclovir-resistant CMV.

Cidofovir is a nucleotide analogue with a long intracellular half-life that allows intermittent IV administration. Induction regimens of 5 mg/kg weekly for 2 weeks are followed by maintenance regimens of 3–5 mg/kg every 2 weeks. Cidofovir can cause severe nephrotoxicity through dose-dependent proximal tubular cell injury; however, this adverse effect can be tempered somewhat by saline hydration and probenecid. Cidofovir is used primarily for ganciclovir-resistant virus.

HUMAN HERPESVIRUS (HHV) TYPES 6, 7, AND 8

HHV-6 AND HHV-7



HHV-6 and -7 seropositivity rates are generally high throughout the world. HHV-6 was first isolated in 1986 from peripheral-blood leukocytes of six persons with various lymphoproliferative disorders. Two genetically distinct variants (HHV-6A and HHV-6B) are now recognized. HHV-6 appears to be transmitted by saliva and possibly by genital secretions.

Infection with HHV-6 frequently occurs during infancy as maternal antibody wanes. The peak age of acquisition is 9–21 months; by 24 months, seropositivity rates approach 80%. Older siblings appear to serve as a source of transmission. Congenital infection also may occur, and 1% of newborns are infected with HHV-6; placental infection with HHV-6 has been described. Most postnatally infected children develop symptoms (fever, fussiness, and diarrhea). A minority develop exanthem subitum (roseola infantum; see Fig. 25e-5), a common illness characterized by fever with subsequent rash. In addition, ~10–20% of febrile seizures without rash during infancy are caused by HHV-6. After initial infection, HHV-6 persists in peripheral-blood mononuclear cells as well as in the central nervous system, salivary glands, and female genital tract.

In older age groups, HHV-6 has been associated with mononucleosis syndromes; in immunocompromised hosts, encephalitis, pneumonitis, syncytial giant-cell hepatitis, and disseminated disease are seen. In transplant recipients, HHV-6 infection may also be associated with graft dysfunction. Acute HHV-6-associated limbic encephalitis has been reported in hematopoietic stem cell transplant recipients and is characterized by memory loss, confusion, seizures, hyponatremia, and abnormal electroencephalographic and MRI results. High plasma loads of HHV-6 DNA in hematopoietic stem cell transplant recipients are associated with allelic-mismatched donors, use of glucocorticoids, delayed monocyte and platelet engraftment, development of limbic encephalitis, and increased all-cause mortality rates. Like many other viruses, HHV-6 has been implicated in the pathogenesis of multiple sclerosis, although further study is needed to distinguish between association and etiology.

HHV-7 was isolated in 1990 from T lymphocytes from the peripheral blood of a healthy 26-year-old man. The virus is frequently acquired during childhood, albeit at a later age than HHV-6. HHV-7 is commonly present in saliva, which is presumed to be the principal source of infection; breast milk also can carry the virus. Viremia can be associated with either primary or reactivation infection. The most common clinical manifestations of childhood HHV-7 infections are fever and seizures. Some children present with respiratory or gastrointestinal signs and symptoms. An association has been made between HHV-7 and pityriasis rosea, but evidence is insufficient to indicate a causal relationship.

Clustering of HHV-6, HHV-7, and CMV infections in transplant recipients can make it difficult to sort out the roles of the various agents in individual clinical syndromes. HHV-6 and HHV-7 appear to be susceptible to ganciclovir and foscarnet, although definitive evidence of clinical response is lacking.

HHV-8

Unique herpesvirus-like DNA sequences were reported during 1994 and 1995 in tissues derived from Kaposi's sarcoma (KS) and body cavity-based lymphoma occurring in patients with AIDS. The virus

from which these sequences were derived is designated HHV-8 or Kaposi's sarcoma-associated herpesvirus (KSHV). HHV-8, which infects B lymphocytes, macrophages, and both endothelial and epithelial cells, appears to be causally related not only to KS and a subgroup of AIDS-related B cell body cavity-based lymphomas (primary effusion lymphomas) but also to multicentric Castleman's disease, a lymphoproliferative disorder of B cells. The association of HHV-8 with several other diseases has been reported but not confirmed.



HHV-8 seropositivity occurs worldwide, with areas of high endemicity influencing rates of disease. Unlike other herpesvirus infections, HHV-8 infection is much more common in some geographic areas (e.g., central and southern Africa) than in others (North America, Asia, northern Europe). In high-prevalence areas, infection occurs in childhood, seropositivity is associated with having a seropositive mother or (to a lesser extent) older sibling, and HHV-8 may be transmitted in saliva. In low-prevalence areas, infections typically occur in adults, probably with sexual transmission. Concurrent epidemics of HIV-1 and HHV-8 infections among certain populations (e.g., men who have sex with men) in the late 1970s and early 1980s appear to have resulted in the frequent association of AIDS and KS. Transmission of HHV-8 may also be associated with organ transplantation, injection drug use, and blood transfusion; however, transmission via blood transfusion in the United States appears to be quite rare.

Primary HHV-8 infection in immunocompetent children may manifest as fever and maculopapular rash. Among individuals with intact immunity, chronic asymptomatic infection is the rule, and neoplastic disorders generally develop only after subsequent immunocompromise. Immunocompromised persons with primary infection may present with fever, splenomegaly, lymphoid hyperplasia, pancytopenia, or rapid-onset KS. Quantitative analysis of HHV-8 DNA suggests a predominance of latently infected cells in KS lesions and frequent lytic replication in multicentric Castleman's disease.

Effective antiretroviral therapy for HIV-infected individuals has led to a marked reduction in rates of KS among persons dually infected with HHV-8 and HIV in resource-rich areas. HHV-8 itself is susceptible in vitro to ganciclovir, foscarnet, and cidofovir. A small, randomized, double-blind, placebo-controlled, crossover trial suggested that oral valganciclovir administered once daily reduced HHV-8 replication. However, clinical benefits of valganciclovir or other drugs in HHV-8 infection have not yet been demonstrated. Sirolimus has been shown to inhibit the progression of dermal KS in kidney transplant recipients while providing effective immunosuppression.

220e Molluscum Contagiosum, Monkeypox, and Other Poxvirus Infections

Fred Wang

The poxvirus family includes a large number of related DNA viruses that infect various vertebrate hosts. The poxviruses responsible for infections in humans, the geographic locations in which these infections are found, the host reservoirs, and the main manifestations are listed in [Table 220e-1](#). Infections with orthopoxviruses—e.g., smallpox (variola major) virus ([Chap. 261e](#)) or the zoonotic monkeypox virus—can result in systemic, potentially lethal human disease. Other poxvirus infections cause primarily localized skin disease in humans.

MOLLUSCUM CONTAGIOSUM

Molluscum contagiosum virus is an obligate human pathogen that causes distinctive proliferative skin lesions. These lesions measure 2–5 mm in diameter and are pearly, flesh-colored, and umbilicated, with a characteristic dimple at the center ([Fig. 220e-1](#)). A relative lack of inflammation and necrosis distinguishes these proliferative lesions from other poxvirus lesions. Lesions may be found—singly or in clusters—anywhere on the body except on the palms and soles and may be associated with an eczematous rash.

Molluscum contagiosum is highly prevalent among children and is the most common human disease resulting from poxvirus infection. Swimming pools are a common vector for transmission. Atopy and compromise of skin integrity increase the risk of infection. Genital lesions are more common in adults, to whom the virus may be transmitted by sexual contact. The incubation period ranges from 2 weeks to 6 months, with an average of 2–7 weeks. In most cases, the disease is self-limited and regresses spontaneously after 3–4 months in immunocompetent hosts. There are no systemic complications, but skin lesions may persist for 3–5 years. Molluscum contagiosum can be associated with immunosuppression and is frequently seen among HIV-infected patients ([Chap. 226](#)). The disease can be more generalized, severe, and persistent in AIDS patients than in other groups. Moreover, molluscum contagiosum can be exacerbated in the immune reconstitution inflammatory syndrome (IRIS) associated with the initiation of antiretroviral therapy.

The diagnosis of molluscum contagiosum is typically based on its clinical presentation and can be confirmed by histologic demonstration of the cytoplasmic eosinophilic inclusions (*molluscum bodies*) that are characteristic of poxvirus replication. Molluscum contagiosum



FIGURE 220e-1 Molluscum contagiosum is a cutaneous poxvirus infection characterized by multiple umbilicated flesh-colored or hypopigmented papules.

virus cannot be propagated *in vitro*, but electron microscopy and molecular studies can be used for its identification.

There is no specific systemic treatment for molluscum contagiosum, but a variety of techniques for physical ablation have been used. Cidofovir displays *in vitro* activity against many poxviruses, and case reports suggest that parenteral or topical cidofovir may have some efficacy in the treatment of recalcitrant molluscum contagiosum in immunosuppressed hosts.

MONKEYPOX



Although monkeypox virus was named after the animal from which it was originally isolated, rodents are the primary viral reservoir. Human infections with monkeypox virus typically occur in Africa when humans come into direct contact with infected animals. Human-to-human propagation of monkeypox infection is rare. Human disease is characterized by a systemic illness and vesicular rash similar to those of variola. The clinical presentation of monkeypox can be confused with that of the more common varicella-zoster virus infection ([Chap. 217](#)). Compared with the lesions of this herpesvirus infection, monkeypox lesions tend to be more uniform (i.e., in the same stage of development), diffuse, and peripheral in distribution. Lymphadenopathy is a prominent feature of monkeypox infection.

The first outbreak of human monkeypox infection in the Western Hemisphere occurred during 2003, when more than 70 cases were reported in the midwestern United States. The outbreak was linked to contact with pet prairie dogs that had become infected while being housed with rodents imported from Ghana. Patients presented most

TABLE 220e-1 POXVIRUSES AND HUMAN INFECTIONS

Genus	Species	Geographic Location	Host Reservoir	Human Disease
<i>Orthopoxvirus</i>	Variola ^a	Extinct	Humans	Smallpox, systemic
	Monkeypox	Africa	Rodents	Smallpox-like, systemic
	Cowpox	Europe	Rodents	Local pox lesion, occasionally systemic
	Buffalopox	Indian subcontinent	Water buffalo	Local pox lesion, mild illness
	Cantagalo and Araçatuba	South America	Cattle	Local pox lesion, mild illness
	Vaccinia	—	—	Local pox lesions (smallpox vaccine)
<i>Molluscipoxvirus</i>	Molluscum contagiosum	Worldwide	Humans	Multiple cutaneous lesions
<i>Parapoxvirus</i>	Orf	Worldwide	Sheep, goats	Local pox lesions (contagious pustular dermatitis)
	Pseudocowpox (paravaccinia)	Worldwide	Cattle	Local pox lesions (milker's nodule)
	Bovine papular stomatitis	Worldwide	Cattle	Local pox lesions
	Deerpox	Deer herds	Deer	Local pox lesions
	Sealpox	Seal colonies	Seals	Local pox lesions
<i>Yatapoxvirus</i>	Tanapox	Africa	Monkeys	Local pox lesions

^aSee [Chap. 261e](#).

220e-2 frequently with fever, rash, and lymphadenopathy ~12 days after exposure. Nine patients were hospitalized, but there were no deaths. Smallpox vaccination can provide cross-reactive immunity to monkeypox infection; studies of people exposed in the outbreak detected subclinical infection in a few vaccinated individuals—an observation suggesting the possibility of long-term vaccine protection. The risk of human disease from animal orthopoxvirus infections may increase as smallpox immunity wanes in the general population and the popularity of exotic animals as household pets grows.

OTHER ZONOTIC POXVIRUS INFECTIONS

Cowpox and buffalopox are rare zoonotic infections characterized by cutaneous poxlike lesions and mild systemic illness. Outbreaks of

similar poxlike lesions among cattle and farm workers in Brazil have been due to Cantagalo and Araçatuba viruses, which are virtually identical to vaccinia virus and may have become established in cattle during smallpox vaccination programs.

Parapoxviruses are widely scattered among animal species, but only a few are known to cause human disease via direct contact with infected animals. Parapoxviruses are antigenically distinct from orthopoxviruses and share no cross-immunity. *Tanapox* virus belongs to a separate, antigenically distinct genus and usually causes a single nodular lesion on the exposed area after contact with infected monkeys.

221 Parvovirus Infections

Kevin E. Brown

Parvoviruses, members of the family Parvoviridae, are small (diameter, ~22 nm), nonenveloped, icosahedral viruses with a linear single-strand DNA genome of ~5000 nucleotides. These viruses are dependent on either rapidly dividing host cells or helper viruses for replication. At least four groups of parvoviruses infect humans: parvovirus B19 (B19V), dependoparvoviruses (adeno-associated viruses; AAVs), PARV4/5 virus, and human bocaviruses (HBoVs). Human dependoparvoviruses are nonpathogenic and will not be considered further in this chapter.

PARVOVIRUS B19

DEFINITION

B19V is the type member of the genus *Erythroparvovirus*. On the basis of viral sequence, B19V is divided into three genotypes (designated 1, 2, and 3), but only a single B19V antigenic type has been described. Genotype 1 is predominant in most parts of the world; genotype 2 is rarely associated with active infection; and genotype 3 appears to predominate in parts of western Africa.

EPIDEMIOLOGY

B19V exclusively infects humans, and infection is endemic in virtually all parts of the world. Transmission occurs predominantly via the respiratory route and is followed by the onset of rash and arthralgia. By the age of 15 years, ~50% of children have detectable IgG; this figure rises to >90% among the elderly. In pregnant

women, the estimated annual seroconversion rate is ~1%. Within households, secondary infection rates approach 50%.

Detection of high-titer B19V in blood is not unusual (see “Pathogenesis,” below). Transmission can occur as a result of transfusion, most commonly of pooled components. To reduce the risk of transmission, plasma pools are screened by nucleic acid amplification technology, and high-titer pools are discarded. B19V is resistant to both heat and solvent-detergent inactivation.

PATHOGENESIS

B19V replicates primarily in erythroid progenitors. This specificity is due in part to the limited tissue distribution of the primary B19V receptor, blood group P antigen (globoside). Infection leads to high-titer viremia, with >10¹² virus particles (or IU)/mL detectable in the blood at the apex (Fig. 221-1), and virus-induced cytotoxicity results in cessation of red cell production. In immunocompetent individuals, viremia and arrest of erythropoiesis are transient and resolve as the IgM and IgG antibody response is mounted. In individuals with normal erythropoiesis, there is only a minimal drop in hemoglobin levels; however, in those with increased erythropoiesis (especially with hemolytic anemia), this cessation of red cell production can induce a transient crisis with severe anemia (Fig. 221-1). Similarly, if an individual (or, after maternal infection, a fetus) does not mount a neutralizing antibody response and halt the lytic infection, erythroid production is compromised and chronic anemia develops (Fig. 221-1).

The immune-mediated phase of illness, which begins 2–3 weeks after infection as the IgM response peaks, manifests as the rash of fifth disease together with arthralgia and/or frank arthritis. Low-level B19V DNA can be detected by polymerase chain reaction (PCR) in blood and tissues for months to years after acute infection. The B19V

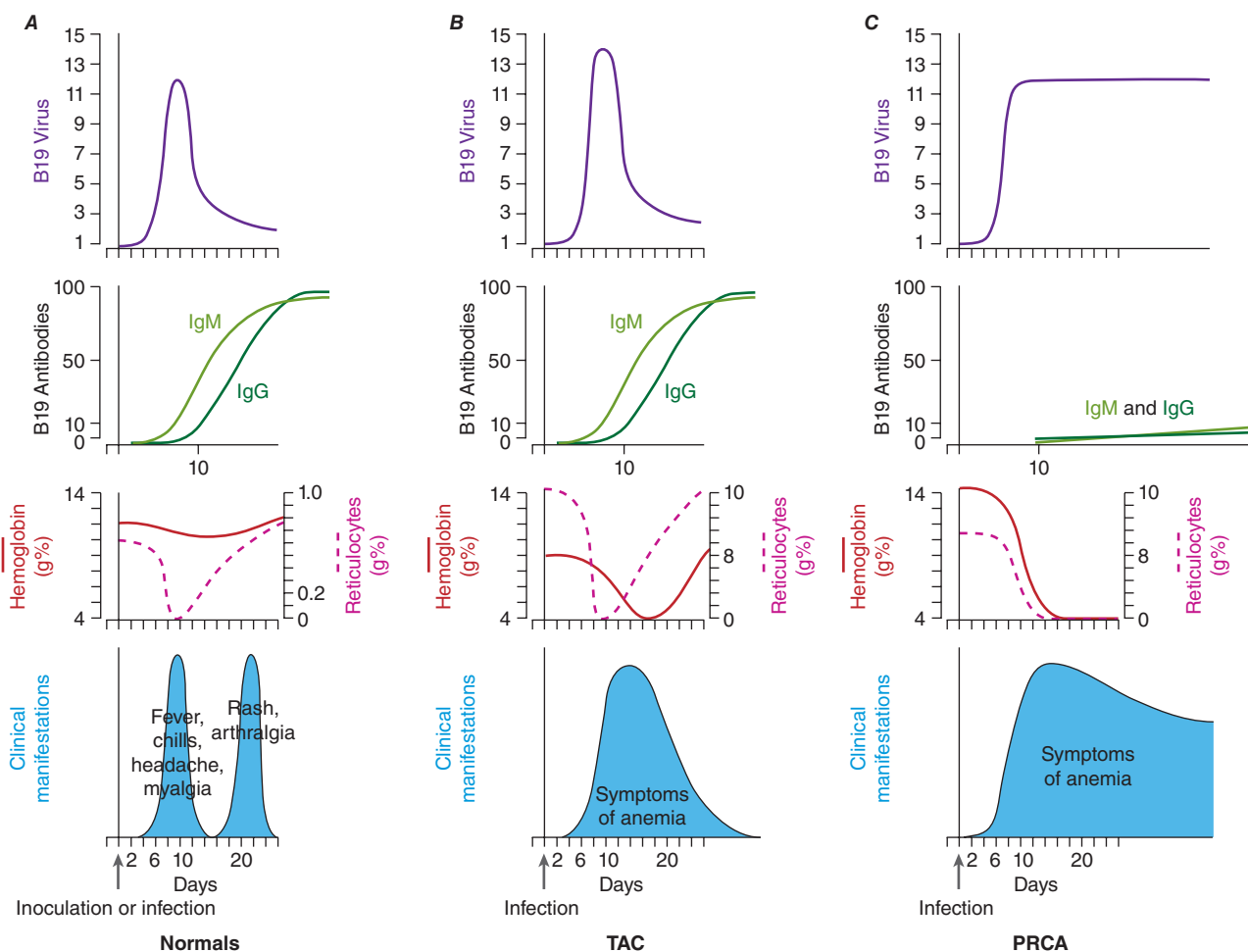


FIGURE 221-1 Schematic of the time course of parvovirus B19V infection in (A) normals (erythema infectiosum), (B) transient aplastic crisis (TAC), and (C) chronic anemia/pure red-cell aplasia (PRCA). (Reprinted with permission from NS Young, KE Brown: *N Engl J Med* 350:586, 2004. © 2004 Massachusetts Medical Society. All rights reserved.)



FIGURE 221-2 Young child with erythema infectiosum, or fifth disease, showing typical “slapped-cheek” appearance.

receptor is found in a variety of other cells and tissues, including megakaryocytes, endothelial cells, placenta, myocardium, and liver. Infection of these tissues by B19V may be responsible for some of the more unusual presentations of the infection. Rare individuals who lack P antigen are naturally resistant to B19V infection.

CLINICAL MANIFESTATIONS

Erythema Infectiosum Most B19V infections are asymptomatic or are associated with only a mild nonspecific illness. The main manifestation of symptomatic B19V infection is erythema infectiosum, also known as *fifth disease* or *slapped-cheek disease* (Fig. 221-2 and Fig. 25e-1A). Infection begins with a minor febrile prodrome ~7–10 days after exposure, and the classic facial rash develops several days later; after 2–3 days, the erythematous macular rash may spread to the extremities in a lacy reticular pattern. However, its intensity and distribution vary, and B19V-induced rash is difficult to distinguish from other viral exanthems. Adults typically do not exhibit the “slapped-cheek” phenomenon but present with arthralgia, with or without the macular rash.

Polyarthropathy Syndrome Although uncommon among children, arthropathy occurs in ~50% of adults and is more common among women than among men. The distribution of the affected joints is often symmetrical, with arthralgia affecting the small joints of the hands and occasionally the ankles, knees, and wrists. Resolution usually occurs

within a few weeks, but recurring symptoms can continue for months. The illness may mimic rheumatoid arthritis, and rheumatoid factor can often be detected in serum. B19V infection may trigger rheumatoid disease in some patients and has been associated with juvenile idiopathic arthritis.

Transient Aplastic Crisis Asymptomatic transient reticulocytopenia occurs in most individuals with B19V infection. However, in patients who depend on continual rapid production of red cells, infection can cause transient aplastic crisis (TAC). Affected individuals include those with hemolytic disorders, hemoglobinopathies, red cell enzymopathies, and autoimmune hemolytic anemias. Patients present with symptoms of severe anemia (sometimes life-threatening) and a low reticulocyte count, and bone marrow examination reveals an absence of erythroid precursors and characteristic giant pronormoblasts. As its name indicates, the illness is transient, and anemia resolves with the cessation of cytopathic infection in the erythroid progenitors.

Pure Red-Cell Aplasia/Chronic Anemia Chronic B19V infection has been reported in a wide range of immunosuppressed patients, including those with congenital immunodeficiency, AIDS (Chap. 226), lymphoproliferative disorders (especially acute lymphocytic leukemia), and transplantation (Chap. 169). Patients have persistent anemia with reticulocytopenia, absent or low levels of B19V IgG, high titers of B19V DNA in serum, and—in many cases—scattered giant pronormoblasts in bone marrow. Rarely, nonerythroid hematologic lineages are also affected. Transient neutropenia, lymphopenia, and thrombocytopenia (including idiopathic thrombocytopenic purpura) have been observed. B19V occasionally causes a hemophagocytic syndrome.



Recent studies in Papua New Guinea and Ghana, where malaria is endemic, suggest that co-infection with *Plasmodium* and B19V plays a major role in the development of severe anemia in young children. Further studies must determine whether B19V infection contributes to severe anemia in other malarial regions.

Hydrops Fetalis B19V infection during pregnancy can lead to hydrops fetalis and/or fetal loss. The risk of transplacental fetal infection is ~30%, and the risk of fetal loss (predominantly early in the second trimester) is ~9%. The risk of congenital infection is <1%. Although B19V does not appear to be teratogenic, anecdotal cases of eye damage and central nervous system (CNS) abnormalities have been reported. Cases of congenital anemia have also been described. B19V probably causes 10–20% of all cases of nonimmune hydrops.

Unusual Manifestations B19V infection may rarely cause hepatitis, vasculitis, myocarditis, glomerulosclerosis, or meningitis. A variety of other cardiac manifestations, CNS diseases, and autoimmune infections have also been reported. However, B19V DNA can be detected by PCR for years in many tissues; this finding is of no known clinical significance, but its interpretation may cause confusion regarding B19V disease association.

DIAGNOSIS

Diagnosis of B19V infection in immunocompetent individuals is generally based on detection of B19V IgM antibodies (Table 221-1). IgM can be detected at the time of rash in erythema infectiosum and

TABLE 221-1 DISEASES ASSOCIATED WITH HUMAN PARVOVIRUS B19 INFECTION AND METHODS OF DIAGNOSIS

Disease	Host(s)	IgM	IgG	PCR	Quantitative PCR
Fifth disease	Healthy children	Positive	Positive	Positive	>10 ⁴ IU/mL
Polyarthropathy syndrome	Healthy adults (more often women)	Positive within 3 months of onset	Positive	Positive	>10 ⁴ IU/mL
Transient aplastic crisis	Patients with increased erythropoiesis	Negative/positive	Negative/positive	Positive	Often >10 ¹² IU/mL, but rapidly decreases
Persistent anemia/pure red-cell aplasia	Immunodeficient or immunocompetent patients	Negative/weakly positive	Negative/weakly positive	Positive	Often >10 ¹² IU/mL, but should be >10 ⁶ in the absence of treatment
Hydrops fetalis/congenital anemia	Fetus (<20 weeks)	Negative/positive	Positive	Positive amniotic fluid or tissue	n/a

Abbreviations: IU, international units (1 IU equals ~1 genome); n/a, not applicable; PCR, polymerase chain reaction.

by the third day of TAC in patients with hematologic disorders; these antibodies remain detectable for ~3 months. B19V IgG is detectable by the seventh day of illness and persists throughout life. Quantitative detection of B19V DNA should be used for the diagnosis of early TAC or chronic anemia. Although B19V levels fall rapidly with the development of the immune response, DNA can be detectable by PCR for months or even years after infection, even in healthy individuals; therefore, quantitative PCR should be used. In acute infection at the height of viremia, $>10^{12}$ B19V DNA IU/mL of serum can be detected; however, titers fall rapidly within 2 days. Patients with aplastic crisis or B19V-induced chronic anemia generally have $>10^5$ B19V DNA IU/mL.

TREATMENT PARVOVIRUS B19 INFECTION

No antiviral drug effective against B19V is available, and treatment of B19V infection often targets symptoms only. TAC precipitated by B19V infection frequently necessitates symptom-based treatment with blood transfusions. In patients receiving chemotherapy, temporary cessation of treatment may result in an immune response and resolution. If this approach is unsuccessful or not applicable, commercial immune globulin (IVIg; Gammagard, Sandoglobulin) from healthy blood donors can cure or ameliorate persistent B19V infection in immunosuppressed patients. Generally, the dose used is 400 mg/kg daily for 5–10 days. Like patients with TAC, immunosuppressed patients with persistent B19V infection should be considered infectious. Administration of IVIg is not beneficial for erythema infectiosum or B19V-associated polyarthropathy. Intrauterine blood transfusion can prevent fetal loss in some cases of fetal hydrops.

PREVENTION

No vaccine has been approved for the prevention of B19V infection, although vaccines based on B19V virus-like particles expressed in insect cells are known to be highly immunogenic. Phase 1 trials of a putative vaccine were discontinued because of adverse side effects.

PARV4/5

DEFINITION

The PARV4 viral sequence was initially detected in a patient with an acute viral syndrome. Similar sequences, including the related PARV5 sequence, have been detected in pooled plasma collections. The DNA sequence of PARV4/5 is distinctly different from that of all other parvoviruses, and this virus is now classified as a member of the newly described genus *Tetraparvovirus*.

EPIDEMIOLOGY



PARV4 DNA is commonly found in plasma pools but at lower concentrations than levels of B19V DNA found before in plasma pools prior to screening. The higher levels of PARV4 DNA and IgG antibody in tissues (bone marrow and lymphoid tissue) and sera from IV drug users than in the corresponding specimens from control patients suggest that the virus is transmitted predominantly by parenteral means in the United States and Europe. Evidence for non-parenteral transmission in other parts of the world is limited.

CLINICAL MANIFESTATIONS

To date, PARV4/5 infection has been associated only with mild clinical disease (rash and/or transient aminotransferase elevation).

HUMAN BOCAVIRUSES

DEFINITION

Animal bocaviruses are associated with mild respiratory symptoms and enteritis in young animals. HBoV1 was originally identified in the respiratory tract of young children with lower respiratory tract infections. More recently, HBoV1 and the related viruses HBoV2, HBoV3, and HBoV4 have all been identified in human fecal samples.

EPIDEMIOLOGY



Seroepidemiologic studies with HBoV virus-like particles suggest that human bocavirus infection is common. Worldwide, most individuals are infected before the age of 5 years.

CLINICAL MANIFESTATIONS

HBoV1 DNA is found in respiratory secretions from 2–20% of children with acute respiratory infection, often in the presence of other pathogens; in these circumstances, the role of HBoV1 in disease pathogenesis is unknown. Clinical disease due to HBoV1 is associated with evidence of primary infection (IgG seroconversion or the presence of IgM), HBoV1 DNA in serum, or high-titer HBoV1 DNA ($>10^4$ genome copies/mL) in respiratory secretions. Symptoms are not dissimilar from those of other viral respiratory infections, and cough and wheezing are commonly reported. There is no specific treatment for bocavirus infection. The role of human bocaviruses in childhood gastroenteritis remains to be established.

222

Human Papillomavirus Infections

Aaron C. Ermel, Darron R. Brown

Investigation of human papillomavirus (HPV) infection began in earnest in the 1980s after Harold zur Hausen postulated that infection with these viruses was associated with cervical cancer. It is now recognized that HPV infection of the human genital tract is extremely common and causes clinical states ranging from asymptomatic infection to genital warts (*condylomata acuminata*); dysplastic lesions or invasive cancers of the anus, penis, vulva, vagina, and cervix; and a subset of oropharyngeal cancers. This chapter describes the epidemiology of HPV in general and as a pathogen, the natural history of HPV infections and associated cancers, strategies to prevent HPV infection and HPV-associated disease, and treatment modalities.

PATHOGENESIS



Molecular Overview HPV is an icosahedral, nonenveloped, 8000-base-pair, double-stranded DNA virus with a diameter of 55 nm. Like those of other papillomaviruses, HPV's genome consists of an early (E) gene region, a late (L) gene region, and a noncoding region that contains regulatory elements. The E1, E2, E5, E6, and E7 proteins are expressed early in the growth cycle and are necessary for viral replication and cellular transformation. The E6 and E7 proteins cause malignant transformation by targeting the human cell cycle-regulatory molecules p53 and Rb (retinoblastoma protein), respectively, for degradation. Translation of the L1 and L2 transcripts and splicing of an E1[^]E4 transcript occur later. The L1 gene encodes the 54-kDa major capsid protein that makes up the majority of the virus shell; the 77-kDa L2 minor protein constitutes a smaller percentage of the capsid mass.

More than 125 HPV types have been identified and are numerically designated according to a unique L1 gene sequence. Approximately 40 HPV types are regularly found in the anogenital tract and are subdivided into high-risk and low-risk categories on the basis of the associated risk of cervical cancer. For example, HPV-6 and HPV-11 cause genital warts and ~10% of low-grade cervical lesions and are thus designated low risk. HPV-16 and HPV-18 cause dysplastic lesions and invasive cancers of the cervix and are considered high risk.

HPV targets basal keratinocytes after microtrauma has exposed these cells to the virus. The HPV replication cycle is completed as keratinocytes undergo differentiation. Virions are assembled in the nuclei of differentiated keratinocytes and can be detected by electron microscopy. Infection is transmitted by contact with virus contained in these desquamated keratinocytes (or with free virus) from an infected individual.

1198 Immune Response to HPV Infection A cell-mediated immune response plays an important role in controlling the progression of natural HPV infection. Histologic examination of lesions in individuals who experience regression of genital warts demonstrates infiltration of T cells and macrophages. CD4+ T cell regulation is particularly important in controlling HPV infections, as evidenced by the higher rates of infection and disease among immunosuppressed individuals, particularly those who are infected with HIV. Specific T-cell responses may be measured against HPV proteins, the most important of which appear to be the E2 and E6 proteins. In women with HPV-16 cervical infection, a strong T-cell response to HPV-16-derived E2 protein is associated with a lack of progression of cervical disease.

Natural HPV infection of the genital tract gives rise to a serum antibody response in only 60–70% of individuals because there is no viremic phase during infection. Significant, although incomplete, protection against type-specific reinfection is associated with the presence of neutralizing antibodies. Serum antibodies likely reach the cervical epithelium and secretions by transudation and exudation. Therefore, protection against infection is related to the amount of neutralizing antibody at the site of infection and lasts as long as levels of neutralizing antibodies are sufficient.

EPIDEMIOLOGY AND NATURAL HISTORY OF HPV-ASSOCIATED MALIGNANCY

General Population HPV is transmitted by sexual intercourse, by oral sex, and possibly by touching of a partner's genitalia. In cross-sectional and longitudinal studies, ~40% of young women have evidence of HPV infection, with peaks during the teens and early twenties—soon after first coitus. The number of lifetime sexual partners correlates with the likelihood of HPV infection and the subsequent risk of HPV-associated malignancy. HPV infection may develop in a monogamous person whose partner is infected. Most HPV infections become undetectable after 6–9 months. However, with prolonged follow-up and frequent sampling, the same HPV types may again be detected weeks or months after becoming undetectable. Whether such episodic detection indicates viral latency followed by reactivation or reinfection with an identical HPV type is still debated.



Although HPV is the causative agent of several cancers, most attention has focused on cervical cancer—the second most common cancer among women worldwide, which affects more than 500,000 women and kills more than 275,000 women annually. More than 85% of all cervical cancer cases and deaths occur in women living in low-income countries, especially in sub-Saharan Africa, Asia, and South and Central America. A quarter-century of evidence shows that HPV causes nearly 100% of cervical cancers. HPV infection is the most significant risk factor for cervical cancer; relative risks range from 10 to 20 and exceed 100 in prospective and case-control studies, respectively. The time from HPV infection to cervical cancer diagnosis may exceed 20 years. Cervical cancer peaks in the fifth and sixth decades of life among women living in developed countries but as much as a decade earlier among women living in resource-poor countries. Persistent carriers of oncogenic HPV types are at greatest risk for high-grade cervical dysplasia and cancer. Why only certain HPV infections eventually lead to malignancy is not clear. Biomarkers that can predict which women will develop cervical cancer are not available. Immunosuppression in general plays a significant role in re-detection/reactivation of HPV infections, while other factors such as smoking, hormonal changes, *Chlamydia* infection, and nutritional deficits promote viral persistence and cancer.

The International Agency for Research on Cancer concludes that HPV types 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, and 59 are carcinogenic in the uterine cervix. HPV-16 is particularly virulent and causes 50% of cervical cancers. Worldwide, HPV-16 and HPV-18 cause 70% of cervical squamous cell carcinomas and 85% of cervical adenocarcinomas. Oncogenic types other than HPV-16 and HPV-18 cause the remaining 30% of cervical cancers. HPV-16 and HPV-18 also cause nearly 90% of anal cancers worldwide. Although oncogenic HPV infection is necessary for the development of cervical malignancy, only

~3–5% of infected women will ever develop this cancer, even in the absence of cytologic screening.

In addition to cervical and anal cancer, other HPV-associated cancers include vulvar and vaginal cancer, which are associated with HPV in 50–70% of cases; penile cancer, which is caused by HPV in 50% of cases; and oropharyngeal squamous cell carcinoma (OPSCC). Over the past two decades, an epidemic of OPSCC related to oncogenic infection with HPV (primarily HPV-16) has developed. Annual rates of OPSCC among men in the United States have been increasing from a low of 0.27 case/100,000 in 1973 to 0.57 case/100,000 as of 2004; rates in women have remained relatively stable at ~0.17 case/100,000 per year. The increase in the incidence of OPSCC is greatest among white men 40–50 years of age. Nearly 14,000 new cases were diagnosed in the United States in 2013. Annual rates of OPSCCs of the base of the tongue and the tonsil have increased dramatically—i.e., by 1.3% and 0.6%, respectively. Fewer data are available from developing countries about OPSCCs.

Effects of HIV on HPV-Associated Disease HIV infection accelerates the natural progression of HPV infections. HIV-infected persons are more likely than other individuals to develop genital warts and to have lesions that are more recalcitrant to treatment. HIV infection has been consistently associated with precancerous cervical lesions, including low-grade cervical intraepithelial neoplasia (CIN) and CIN 3, the immediate precursor to cervical cancer. Women with HIV/AIDS have significantly higher rates of cervical cancer and of subsets of some vulvar, vaginal, and oropharyngeal tumors than women in the general population. Studies indicate a direct relation between low CD4+ T lymphocyte counts and the risk of cervical cancer. Some studies show a reduced likelihood of HPV infection and precancerous lesions of the cervix in HIV-infected women receiving antiretroviral therapy (ART). The incidence of cervical cancer among HIV-infected women has not changed significantly since ART was introduced, possibly because of preexisting oncogenic HPV infections.



The burden of HPV-related cancers is expected to increase in HIV-infected patients, given the prolonged life expectancies made possible by ART. For women living in developing countries where cervical cancer screening is not widely available, this situation may have significant consequences. Thus, elucidating the interactions of HIV infection and cervical cancer with cofactors such as diet, other sexually transmitted infections, and environmental exposures is a research focus with potentially enormous implications for women living in low- and middle-income countries.

Similar to that of cervical cancer, the incidence of anal cancer is strongly influenced by HIV infection. HIV-infected men who have sex with men (MSM) and HIV-infected women have much higher rates of anal cancer than HIV-uninfected populations. Specifically, the incidence has been found to be as high as 130 cases/100,000 among HIV-positive MSM as opposed to only 5 cases/100,000 among HIV-negative MSM. The advent of ART has not impacted the incidence of anal cancer and high-grade anal intraepithelial neoplasia in the HIV-infected population.

More information on screening, prevention, and treatment in the HIV-infected population can be found at the Department of Health and Human Services website (aidsinfo.nih.gov/guidelines).

CLINICAL MANIFESTATIONS OF HPV INFECTION

HPV infects the female vulva, vagina, and cervix and the male urethra, penis, and scrotum. Perianal, anal, and oropharyngeal infections occur in both genders. **Figures 222-1, 222-2, and 222-3** show vulvar, penile, and perianal warts, respectively. Genital warts are caused primarily by HPV-6 or HPV-11; their surface is either smooth or rough. Penile genital warts are usually 2–5 mm in diameter and often occur in groups. A second type of penile lesion, keratotic plaques, is slightly raised above the normal epithelium and has a rough, often pigmented surface. Vulvar warts are soft, whitish papules that either are sessile or have multiple fine, finger-like projections. These lesions are most often located in the introitus and on the labia. In nonmucosal areas, lesions



FIGURE 222-1 Vulvar warts. (Downloaded from <http://www2a.cdc.gov/stdtraining/ready-to-use/Manuals/HPV/hpv-slides-2013.pdf>.)

are similar in appearance to those in men: dry and keratotic. Vulvar lesions can appear as smooth, sometimes pigmented papules that may coalesce. Vaginal lesions appear as multiple areas of elongated papillae. Biopsy of vulvar or vaginal lesions may reveal malignancy, which is not always reliably identified by clinical examination.

Subclinical cervical HPV infections are common, and the cervix may appear normal on examination. Cervical lesions often appear as papillary proliferations near the transformation zone. Irregular vascular loops are present beneath the surface epithelium. Patients who develop cervical cancer arising from HPV infection may present with a variety of symptoms. Early carcinomas appear eroded and bleed easily. More advanced carcinomas present as ulcerated lesions or as an exophytic cervical mass. Some cervical carcinomas are located in the cervical canal and may be difficult to see. Bleeding, symptoms of a mass lesion in late stages, and metastatic disease that may manifest as bowel or bladder obstruction due to direct extension of the tumor have also been described.



FIGURE 222-2 Condyloma acuminata of the shaft of the penis.



FIGURE 222-3 Perianal warts. (Reprinted from K Wolff, RA Johnson, AP Saavedra: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 7th ed. New York, McGraw-Hill, 2013.)

Patients with squamous cell cancer of the anus have more variable presentations. The most common presentations include rectal bleeding and pain or a mass sensation. Of patients who are diagnosed with anal cancer, 20% may present with no specific symptoms at the time of diagnosis; rather, the lesion is found fortuitously.

PREVENTION OF HPV INFECTION: HPV VACCINES

Vaccines effective in preventing HPV infection and HPV-associated disease represent a major development in the last decade. The vaccines use virus-like particles (VLPs) that consist of the HPV L1 major capsid protein. The L1 protein self-assembles into VLPs when expressed in eukaryotic cells (i.e., yeast for the Merck vaccine or insect cells for the GlaxoSmithKline vaccine; see below). These VLPs contain the same epitopes as the HPV virion. However, they do not contain genetic material and cannot transmit infection. The immunogenicity of HPV vaccines relies on the development of conformational neutralizing antibodies to epitopes displayed on viral capsids.

Several large trials have demonstrated the high degree of safety and efficacy of HPV vaccines. The evidence to date has shown high and sustained efficacy against disease caused by those HPV types represented in the vaccines (HPV-6, -11, -16, and -18 in the Merck vaccine and HPV-16 and -18 in the GlaxoSmithKline vaccine). However, no therapeutic effect against active infection or disease has been found for either vaccine.

Bivalent Vaccine (Cervarix) A bivalent L1 VLP vaccine (HPV-16 and -18), marketed under the name Cervarix (GlaxoSmithKline), is administered by IM injection at months 0, 1, and 6. This vaccine was tested in 18,644 women 15–25 years of age who were residing in the United States, South America, Europe, and Asia. The primary endpoints of the study included vaccine efficacy against persistent infections with HPV-16 and -18. Investigators also assessed the vaccine's efficacy against CIN of grade 2 or higher due to HPV-16 and -18 in women who had no evidence of infection with these HPV types at baseline; in these women, vaccine efficacy was 94.9% (95% confidence interval [CI], 87.7 to 98.4) against CIN ≥ 2 related to HPV-16 or HPV-18, 91.7% (95% CI, 66.6 to 99.1) against CIN ≥ 3 , and 100% (95% CI, -8.6 to 100) against adenocarcinoma in situ.

Adverse events were evaluated in phase 3 trials in a subset of 3077 women who received the bivalent vaccine and 3080 women (controls) who received hepatitis A vaccine. Injection-site adverse events (pain, redness, and swelling) and systemic adverse events (fatigue, headache, and myalgia) were reported more frequently in the HPV vaccine group than in the control group. Serious adverse events (mainly injection-site reactions), new-onset chronic disease, or medically significant conditions occurred in 3.5% of HPV vaccine recipients and in 3.5% of women receiving the control vaccine.

The bivalent vaccine is approved in the United States for prevention of cervical cancer, CIN ≥ 2 , adenocarcinoma in situ, and CIN 1 caused by HPV-16 and -18. This vaccine is approved for administration to girls and women 9–25 years of age.

Quadrivalent Vaccine (Gardasil) A quadrivalent L1 VLP (HPV-6, -11, -16, and -18) vaccine, marketed under the name Gardasil (Merck), is administered IM at months 0, 2, and 6. A combined efficacy analysis based on data from four randomized double-blind clinical studies including more than 20,000 participants demonstrated that the vaccine's efficacy against external genital warts was 98.9% (95% CI, 93.7 to 100). Its efficacy was 95.2% (95% CI, 87.2 to 98.7) in protecting against CIN, 100% (95% CI, 92.9 to 100) against HPV-16- or HPV-18-related CIN 2/3 or adenocarcinoma in situ, and 100% (95% CI, 55.5 to 100.0) against HPV-16- or HPV-18-related vulvar intraepithelial neoplasia grades 2 and 3 (VIN 2/3) and vaginal intraepithelial neoplasia grades 2 and 3 (VaIN 2/3).

Safety data on the quadrivalent HPV vaccine are available from seven clinical trials including nearly 12,000 girls and women 9–26 years of age who received the vaccine and ~10,000 who received placebo. A larger proportion of young women reported injection-site adverse events in the vaccine groups than in the aluminum-containing or saline placebo groups. Systemic adverse events were reported by similar proportions of vaccine and placebo recipients and were described as mild or moderate by most participants. The types of serious adverse events reported were similar for the two groups. Ten persons who received the quadrivalent vaccine and seven persons who received placebo died during the course of the trials; no deaths were considered to be vaccine related.

During the course of the quadrivalent vaccine trials, surveillance data on the development of new medical conditions were collected for up to 4 years after vaccination. No statistically significant differences in the incidence of any medical conditions between vaccine and placebo recipients were found; this result indicated a very good safety profile. A recent safety review by the U.S. Food and Drug Administration and the Centers for Disease Control and Prevention (CDC) examined events related to Gardasil that had been reported to the Vaccine Adverse Events Reporting System. Adverse events were consistent with those seen in previous safety studies of the vaccine. It is noteworthy that rates of syncope and venous thrombotic events were higher with Gardasil than those that have usually been documented for other vaccines.

The quadrivalent vaccine is approved for (1) vaccination of girls and women 9–26 years of age to prevent genital warts and cervical cancer caused by HPV-6, -11, -16, and -18; (2) vaccination of the same population to prevent precancerous or dysplastic lesions, including cervical adenocarcinoma in situ, CIN 2/3, VIN 2/3, VaIN 2/3, and CIN 1; (3) vaccination of boys and men 9–26 years of age to prevent genital warts caused by HPV-6 and -11; and (4) vaccination of individuals 9–26 years of age to prevent anal cancer and associated precancerous lesions due to HPV-6, -11, -16, and -18.

Cross-Protection of HPV Vaccines Women vaccinated with either of the available vaccines produce neutralizing antibodies against types related to HPV-16 or -18. Analyses of data from clinical trials suggest that both vaccines may offer cross-protection against nonvaccine types. The bivalent vaccine appears more efficacious against HPV-31, -33, and -45 than the quadrivalent vaccine, but differences in study design make direct comparisons difficult. In addition, vaccine efficacy against persistent infections with HPV-31 and -45 appeared to wane over time in the bivalent vaccine trials, whereas efficacy against persistent infection with HPV-16 or -18 remained stable.



Second-Generation Vaccines While HPV-16 and -18 cause the majority of cervical cancers worldwide, global data have shown that HPV-31, -33, -35, -45, -52, and -58 are the next most frequently detected types in invasive cervical cancers. Second-generation vaccines that are in development incorporate VLPs of additional oncogenic HPV types (other than HPV-16 and -18), including HPV-31, -33, -45, -52, and -58; efficacy studies are ongoing.

If vaccines with these five additional oncogenic types prove to be effective, mathematical models estimate that the level of protection could be raised to 90% of all squamous cell cervical cancers worldwide.

Recommendations for Vaccination The CDC's Advisory Committee for Immunization Practice recommends administration of the quadrivalent HPV vaccine—with the schedule used in the vaccine trials—to all boys and girls 11–12 years of age as well as to boys/men and girls/women 13–26 years of age who have not previously been vaccinated or who have not completed the full series. For women, Papanicolaou (Pap) smear testing and screening for HPV DNA are not recommended before vaccination. After vaccination, Pap testing is recommended to detect disease caused by other oncogenic HPV types.

PREVENTION OF HPV-ASSOCIATED DISEASE



After HPV infection occurs, prevention of HPV-associated disease relies on screening. Women residing in developing countries who lack access to cervical screening programs have a higher rate of cervical cancer and a lower rate of cancer-specific survival. Approximately 75% of women living in developed countries have been screened in the past 5 years, whereas the figure is only ~5% among women living in developing countries. Economic and logistic obstacles likely impede routine screening of these populations for cervical cancer.

The primary method used for cancer screening is cervical cytology via Pap smear. The guidelines of the American Society of Colposcopy and Cervical Pathology recommend initiation of cervical cancer screening at age 21, regardless of the age of sexual debut. Women 21–29 years old with a normal Pap smear should have the test repeated every 3 years. Although adolescent and young women often test positive for HPV DNA, they are at very low risk of cervical cancer. Co-testing, or testing for HPV DNA at the time of the Pap smear, is not recommended for women in this age group because the presence of HPV DNA does not correlate with the presence of high-grade squamous intraepithelial neoplasia. Women 30–65 years of age should have a Pap smear every 3 years if testing for HPV DNA is not performed. The screening interval for women in this age group can be extended to every 5 years if co-testing results are negative. HPV testing is not recommended for partners of women with HPV or for screening for conditions other than cervical cancer.

Currently, there is no clear consensus regarding screening for anal cancer and its precursors, including high-grade anal intraepithelial lesions. This lack of clarity is due to an inadequate understanding of optimal treatment for low- or high-grade anal dysplasia found during cytologic screening. The current HIV treatment guidelines suggest that there may be a benefit to screening, but an effect on the associated morbidity and mortality of anal squamous cell cancer has not been consistently demonstrated.

TREATMENT HPV-ASSOCIATED LESIONS

OVERVIEW AND GENERAL RECOMMENDATIONS

A variety of treatment modalities are available for various HPV infections, but none has been proven to eliminate HPV from tissue adjacent to the destroyed and infected tissue. Treatment efficacies are limited by frequent recurrences (presumably due to reinfection acquired from an infected partner), reactivation of latent virus, or autoinoculation from nearby infected cells. The goals of treatment include prevention of virus transmission, eradication of premalignant lesions, and reduction of symptoms.

Treatment is generally successful in eliminating visible lesions and grossly diseased tissue. Different therapies are indicated for genital warts, vaginal and cervical disease, and perianal and anal disease.

An optimal therapy for HPV-related genital tract disease that combines high efficacy, low toxicity, low cost, and low recurrence rate is not available. For genital warts of the penis or vulva, cryotherapy (see below) is safest, least expensive, and most effective. Guidelines for the treatment of genital warts can be found on the CDC website (www.cdc.gov/std/treatment/2010/genital-warts.htm). Women with vaginal

lesions should be referred to a gynecologist experienced in colposcopy and treatment of these lesions. Treatment of cervical disease involves careful inspection, biopsy, and histopathologic grading to determine the severity and extent of disease. Women with evidence of cervical HPV infection should be referred to a gynecologist familiar with HPV and experienced in colposcopy. Optimal follow-up of these patients includes colposcopic examination of the cervix and vagina on a yearly basis. Guidelines from the American College of Gynecology and Obstetrics are available for the treatment of cervical dysplasia and cancer.

For anal or perianal lesions, cryotherapy or surgical removal is safest and most effective. Anoscopy and/or sigmoidoscopy should be performed when patients have perianal lesions, and suspicious lesions should be biopsied to rule out malignancy.

THERAPEUTIC OPTIONS

Tables 222-1 and 222-2 list the available patient-administered and physician-administered treatments, respectively.

Podophyllotoxin Podophyllotoxin (0.05% solution or gel and 0.15% cream) induces necrosis of genital wart tissue that heals within a few days. It is relatively effective and can be self-administered. Podophyllotoxin is applied twice daily on three consecutive days of the week for a maximum of 4 weeks. Adverse effects are common and include pain, inflammation, erosion, and burning or itching. Podophyllotoxin should not be used to treat vaginal, cervical, or anal lesions. The safety of podophyllotoxin during pregnancy has not been established.

Sinecatechins Sinecatechins (15% ointment) is used to treat genital warts but should not be used to treat vaginal, cervical, or anal lesions. Sinecatechins causes an inflammatory response when applied topically three times a day for up to 4 months. Clearance rates approach 60% in some studies, and recurrence rates are 6–9%. Adverse effects (redness, burning, itching, and pain at the site of application) are generally mild. The safety of sinecatechins during pregnancy is unknown.

Imiquimod Imiquimod (5% or 3.75% cream) is a patient-applied topical immunomodulatory agent thought to activate immune cells by binding to a Toll-like receptor—an event that leads to an inflammatory response. Imiquimod 5% cream is applied to genital warts at bedtime three times per week for up to 16 weeks. Warts are cleared in ~56% of patients, more often in women than in men; recurrence rates approach 13%. Local inflammatory side effects are common. Rates of clearance of genital warts with the 3.75% formulation are not as high, but the duration of treatment is shorter (i.e., daily application for a maximum of 8 weeks), and fewer local and systemic adverse reactions occur. Imiquimod should not be used to treat vaginal, cervical, or anal lesions. The safety of imiquimod during pregnancy has not been established.

Cryotherapy Cryotherapy (liquid nitrogen) for HPV-associated lesions causes cellular death. Genital warts usually disappear after two or three weekly sessions but often recur. Cryotherapy, which

TABLE 222-2 RECOMMENDED TREATMENTS THAT MUST BE ADMINISTERED BY A CLINICIAN FOR GENITAL WARTS AND OTHER HUMAN PAPILLOMAVIRUS-ASSOCIATED LESIONS

Variable	Cryotherapy	Surgical Removal	Laser	Interferon
Effectiveness	Good	Excellent	Excellent	Good
Recurrence	Frequent	Frequent	Frequent	Frequent
Adverse effects	Mild, well tolerated	Mild, well tolerated	Mild to moderate, well tolerated	Frequent, moderately severe
Availability	Good	Good	Fair	Fair
Cost	Inexpensive	Moderately expensive	Very expensive	Very expensive

is nontoxic and is not associated with significant adverse reactions, can also be used for diseased cervical tissue. Local pain occurs frequently.

Surgical Methods Exophytic lesions can be surgically removed after intradermal injection of 1% lidocaine. This treatment is well tolerated but can cause scarring and requires hemostasis. Genital warts can also be destroyed by electrocautery, in which no additional hemostasis is required.

Laser Therapy Laser treatment affords destruction of exophytic lesions and other HPV-infected tissue while preserving normal tissue. Local anesthetics are generally adequate. Efficacy for genital lesions is at least equal to that of other therapies (60–90%), with low recurrence rates (5–10%). Complications include local pain, vaginal discharge, periurethral swelling, and penile or vulvar swelling. Laser therapy has also been used successfully to treat cervical dysplasia and anal disease caused by HPV.

Interferon (IFN) Recombinant IFN- α is used for intralesional treatment of genital warts, including perianal lesions. The recommended dosage is 1.0×10^6 IU of IFN injected into each lesion three times weekly for 3 weeks. IFN therapy causes clearance of infected cells by immune-boosting effects. Adverse events include headache, nausea, vomiting, fatigue, and myalgia. IFN therapy is costly and should be reserved for severe cases that do not respond to cheaper treatments. IFN should not be used to treat vaginal, cervical, or anal lesions.

Other Therapies Both trichloroacetic acid and bichloroacetic acid are caustic agents that destroy warts by coagulation of proteins. Neither of these agents is recommended for treatment.

COUNSELING

Most sexually active adults will be infected with HPV during their lives. For all patients (vaccinated or unvaccinated), certain behavioral interventions can reduce the risk of acquiring HPV. Physicians can provide their patients with measures that can reduce this risk. The only way to avoid acquiring an HPV infection is to abstain from sexual activity, including intimate touching and oral sex. Practicing safe sex (partner reduction, condom use) may lower the likelihood of HPV transmission. Most HPV infections are controlled by the immune system and cause no symptoms or disease. Some infections lead to genital warts and cervical precancers. Genital warts can be treated for cosmetic reasons and to prevent spread of infection to others. Even after resolution of genital warts, latent virus can persist in normal-appearing skin or mucosa and thus theoretically can be transmitted to uninfected partners. Precancerous cervical lesions should be treated to prevent progression to cancer.

TABLE 222-1 RECOMMENDED TREATMENTS THAT CAN BE SELF-ADMINISTERED FOR GENITAL WARTS

Variable	Podophyllotoxin	Sinecatechins	Imiquimod
Effectiveness	Good	Good	Good
Recurrence	Frequent	Frequent	Frequent
Adverse effects	Frequent, can be severe	Frequent, mild	Frequent, mild to moderate
Availability	Good	Fair	Fair
Cost	Inexpensive	Inexpensive	Expensive

223 Common Viral Respiratory Infections

Raphael Dolin

GENERAL CONSIDERATIONS

Acute viral respiratory illnesses are among the most common of human diseases, accounting for one-half or more of all acute illnesses. The incidence of acute respiratory disease in the United States is 3–5.6 cases per person per year. The rates are highest among children <1 year old (6.1–8.3 cases per year) and remain high until age 6, when a progressive decrease begins. Adults have 3–4 cases per person per year. Morbidity from acute respiratory illnesses accounts for 30–50% of time lost from work by adults and for 60–80% of time lost from school by children. The use of antibacterial agents to treat viral respiratory infections represents a major source of abuse of that category of drugs.

It has been estimated that two-thirds to three-fourths of cases of acute respiratory illness are caused by viruses. More than 200 antigenically distinct viruses from 10 genera have been reported to cause acute respiratory illness, and it is likely that additional agents will be described in the future. The vast majority of these viral infections involve the upper respiratory tract, but lower respiratory tract disease can also develop, particularly in younger age groups, in the elderly, and in certain epidemiologic settings.

The illnesses caused by respiratory viruses traditionally have been divided into multiple distinct syndromes, such as the “common cold,” pharyngitis, croup (laryngotracheobronchitis), tracheitis, bronchiolitis, bronchitis, and pneumonia. Each of these general categories of illness has a certain epidemiologic and clinical profile; for example, croup occurs exclusively in very young children and has a characteristic clinical course. Some types of respiratory illness are more likely to be associated with certain viruses (e.g., the common cold with rhinoviruses), whereas others occupy characteristic epidemiologic niches (e.g., adenovirus

infections in military recruits). The syndromes most commonly associated with infections with the major respiratory virus groups are summarized in [Table 223-1](#). Most respiratory viruses clearly have the potential to cause more than one type of respiratory illness, and features of several types of illness may be found in the same patient. Moreover, the clinical illnesses induced by these viruses are rarely sufficiently distinctive to permit an etiologic diagnosis on clinical grounds alone, although the epidemiologic setting increases the likelihood that one group of viruses rather than another is involved. In general, laboratory methods must be relied on to establish a specific viral diagnosis.

This chapter reviews viral infections caused by six of the major groups of respiratory viruses: rhinoviruses, coronaviruses, respiratory syncytial viruses, metapneumoviruses, parainfluenza viruses, and adenoviruses. The extraordinary outbreaks of lower respiratory tract disease associated with coronaviruses (severe acute respiratory syndrome [SARS] in 2002–2003 and Middle East respiratory syndrome [MERS] in 2012–2013) are also discussed. Influenza viruses, which are a major cause of death as well as morbidity, are reviewed in [Chap. 224](#). Herpesviruses, which occasionally cause pharyngitis and which also cause lower respiratory tract disease in immunosuppressed patients, are reviewed in [Chap. 216](#). Enteroviruses, which account for occasional respiratory illnesses during the summer months, are reviewed in [Chap. 228](#).

RHINOVIRUS INFECTIONS

ETIOLOGIC AGENT

Rhinoviruses are members of the Picornaviridae family—small (15- to 30-nm) nonenveloped viruses that contain a single-stranded RNA genome. Human rhinoviruses were first classified by immunologic serotype and are now divided into three genetic species: HRV-A, HRV-B, and HRV-C. The 102 serotypes described initially are encompassed by HRV-A and HRV-B species, whereas HRV-C encompasses more than 60 previously unrecognized serotypes. In contrast to other members of the picornavirus family, such as enteroviruses, rhinoviruses are acid-labile and are almost completely inactivated at pH ≤3. HRV-A and HRV-B species grow preferentially at 33°–34°C (the temperature of the human nasal passages) rather

TABLE 223-1 ILLNESSES ASSOCIATED WITH RESPIRATORY VIRUSES

Virus	Most Frequent	Frequency of Respiratory Syndromes	
		Occasional	Infrequent
Rhinoviruses	Common cold	Exacerbation of chronic bronchitis and asthma	Pneumonia in children
Coronaviruses ^{a,b}	Common cold	Exacerbation of chronic bronchitis and asthma	Pneumonia and bronchiolitis
Human respiratory syncytial virus	Pneumonia and bronchiolitis in young children	Common cold in adults	Pneumonia in elderly and immunosuppressed patients
Parainfluenza viruses	Croup and lower respiratory tract disease in young children	Pharyngitis and common cold	Tracheobronchitis in adults; lower respiratory tract disease in immunosuppressed patients
Adenoviruses	Common cold and pharyngitis in children	Outbreaks of acute respiratory disease in military recruits ^c	Pneumonia in children; lower respiratory tract and disseminated disease in immunosuppressed patients
Influenza A viruses	Influenza ^d	Pneumonia and excess mortality in high-risk patients	Pneumonia in healthy individuals
Influenza B viruses	Influenza ^d	Rhinitis or pharyngitis alone	Pneumonia
Enteroviruses	Acute undifferentiated febrile illnesses ^e	Rhinitis or pharyngitis alone	Pneumonia
Herpes simplex viruses	Gingivostomatitis in children; pharyngotonsillitis in adults	Tracheitis and pneumonia in immunocompromised patients	Disseminated infection in immunocompromised patients
Human metapneumoviruses	Upper and lower respiratory tract disease in children	Upper respiratory tract illness in adults	Pneumonia in elderly and immunosuppressed patients

^aSevere acute respiratory syndrome–associated coronavirus (SARS-CoV) caused epidemics of pneumonia from November 2002 to July 2003 (see text). ^bMiddle East respiratory syndrome coronavirus (MERS-CoV) has caused severe respiratory illnesses from 2012 to the time of this writing (2014); see text. ^cSerotypes 4 and 7 most commonly; also serotypes 14 and 21. ^dFever, cough, myalgia, malaise. ^eMay or may not have a respiratory component.

than at 37°C (the temperature of the lower respiratory tract), whereas HRV-C viruses replicate well at either temperature. Of the 101 initially recognized serotypes of rhinovirus, 88 use intercellular adhesion molecule 1 (ICAM-1) as a cellular receptor and constitute the “major” receptor group, 12 use the low-density lipoprotein receptor (LDLR) and constitute the “minor” receptor group, and 1 uses decay-accelerating factor.

EPIDEMIOLOGY



Rhinovirus infections are worldwide in distribution. They are a prominent cause of the common cold and have been detected in up to 50% of common cold-like illnesses by tissue culture and polymerase chain reaction (PCR) techniques. Overall rates of rhinovirus infection are higher among infants and young children and decrease with increasing age. Rhinovirus infections occur throughout the year, with seasonal peaks in early fall and spring in temperate climates. These infections are most often introduced into families by preschool or grade-school children <6 years old. Of initial illnesses in family settings, 25–70% are followed by secondary cases, with the highest attack rates among the youngest siblings at home. Attack rates also increase with family size.

Rhinoviruses appear to spread through direct contact with infected secretions, usually respiratory droplets. In some studies of volunteers, transmission was most efficient by hand-to-hand contact, with subsequent self-inoculation of the conjunctival or nasal mucosa. Other studies demonstrated transmission by large- or small-particle aerosol. Virus can be recovered from plastic surfaces inoculated 1–3 h previously; this observation suggests that environmental surfaces contribute to transmission. In studies of married couples in which neither partner had detectable serum antibody, transmission was associated with prolonged contact (≥ 122 h) during a 7-day period. Transmission was infrequent unless (1) virus was recoverable from the donor’s hands and nasal mucosa, (2) at least 1000 TCID₅₀ (50% tissue culture infectious dose) of virus was present in nasal washes from the donor, and (3) the donor was at least moderately symptomatic with the “cold.” Despite anecdotal observations, exposure to cold temperatures, fatigue, and sleep deprivation have not been associated with increased rates of rhinovirus-induced illness in volunteers, although some studies have suggested that psychologically defined “stress” may contribute to development of symptoms.

By adulthood, nearly all individuals have neutralizing antibodies to multiple rhinovirus serotypes, although the prevalence of antibody to any one serotype varies widely. Multiple serotypes circulate simultaneously, and generally no single serotype or group of serotypes has been more prevalent than the others.

PATHOGENESIS

Rhinoviruses infect cells through attachment to specific cellular receptors; as mentioned above, most serotypes attach to ICAM-1, while a few use LDLR. Relatively limited information is available on the histopathology and pathogenesis of acute rhinovirus infections in humans. Examination of biopsy specimens obtained during experimentally induced and naturally occurring illness indicates that the nasal mucosa is edematous, is often hyperemic, and—during acute illness—is covered by a mucoid discharge. There is a mild infiltrate with inflammatory cells, including neutrophils, lymphocytes, plasma cells, and eosinophils. Mucus-secreting glands in the submucosa appear hyperactive; the nasal turbinates are engorged, a condition that may lead to obstruction of nearby openings of sinus cavities. Several mediators—e.g., bradykinin; lysylbradykinin; prostaglandins; histamine; interleukins 1 β , 6, and 8; interferon γ -induced protein 10; and tumor necrosis factor α —have been linked to the development of signs and symptoms in rhinovirus-induced colds.

The incubation period for rhinovirus illness is short, generally 1–2 days. Virus shedding coincides with the onset of illness or may begin shortly before symptoms develop. The mechanisms of immunity to rhinovirus infection are not well worked out. In some studies, the presence of homotypic antibody has been associated with significantly reduced rates of subsequent infection and illness, but data conflict regarding the relative importance of serum and local antibody in protection from rhinovirus infection.

CLINICAL MANIFESTATIONS

The most common clinical manifestations of rhinovirus infections are those of the common cold. Illness usually begins with rhinorrhea and sneezing accompanied by nasal congestion. The throat is frequently sore, and in some cases sore throat is the initial complaint. Systemic signs and symptoms, such as malaise and headache, are mild or absent, and fever is unusual in adults but may occur in up to one-third of children. Illness generally lasts for 4–9 days and resolves spontaneously without sequelae. In children, bronchitis, bronchiolitis, and bronchopneumonia have been reported; nevertheless, it appears that rhinoviruses are not major causes of lower respiratory tract disease in children. Rhinoviruses may cause exacerbations of asthma and chronic pulmonary disease in adults. The vast majority of rhinovirus infections resolve without sequelae, but complications related to obstruction of the eustachian tubes or sinus ostia, including otitis media or acute sinusitis, can develop. In immunosuppressed patients, particularly bone marrow transplant recipients, severe and even fatal pneumonias have been associated with rhinovirus infections.

DIAGNOSIS

Although rhinoviruses are the most frequently recognized cause of the common cold, similar illnesses are caused by a variety of other viruses, and a specific viral etiologic diagnosis cannot be made on clinical grounds alone. Rather, rhinovirus infection is diagnosed by isolation of the virus from nasal washes or nasal secretions in tissue culture. In practice, this procedure is rarely undertaken because of the benign, self-limited nature of the illness. In most settings, detection of rhinovirus RNA is more sensitive and easier by PCR than by tissue culture. Accordingly, PCR has generally become the standard for detection of rhinoviruses in clinical specimens. Given the many serotypes of rhinovirus, diagnosis by serum antibody tests is currently impractical. Likewise, common laboratory tests, such as white blood cell count and erythrocyte sedimentation rate, are not helpful.

TREATMENT RHINOVIRUS INFECTIONS

Because rhinovirus infections are generally mild and self-limited, treatment is not usually necessary. Therapy in the form of first-generation antihistamines and nonsteroidal anti-inflammatory drugs may be beneficial in patients with particularly pronounced symptoms, and an oral decongestant may be added if nasal obstruction is particularly troublesome. Reduction of activity is prudent in instances of significant discomfort or fatigability. Antibacterial agents should be used only if bacterial complications such as otitis media or sinusitis develop. Specific antiviral therapy is not available.

PREVENTION

Intranasal application of interferon sprays has been effective in the prophylaxis of rhinovirus infections but is also associated with local irritation of the nasal mucosa. Studies of prevention of rhinovirus infection by blocking of ICAM-1 or by binding of drug (pleconaril) to parts of the viral capsid have yielded mixed results. Experimental vaccines to certain rhinovirus serotypes have been generated, but their usefulness is questionable because of the myriad serotypes involved and the uncertainty about mechanisms of immunity. Thorough hand washing, environmental decontamination, and protection against autoinoculation may help to reduce rates of transmission of infection.

CORONAVIRUS INFECTIONS

ETIOLOGIC AGENT

Coronaviruses are pleomorphic, single-stranded RNA viruses that measure 100–160 nm in diameter. The name derives from the crown-like appearance produced by the club-shaped projections that stud the viral envelope. Coronaviruses infect a wide variety of animal species and have been divided into four genera. Coronaviruses that infect humans (HCoVs) fall into two genera: *Alphacoronavirus*

1204 and *Betacoronavirus*. Severe acute respiratory syndrome coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV) are betacoronaviruses.

In general, human coronaviruses have been difficult to cultivate *in vitro*, and some strains grow only in human tracheal organ cultures rather than in tissue culture. SARS-CoV and MERS-CoV are exceptions whose ready growth in African green monkey kidney (Vero E6) cells greatly facilitates their study.

EPIDEMIOLOGY



Human coronavirus infections are present throughout the world. Seroprevalence studies of strains HCoV-229E and HCoV-OC43 have demonstrated that serum antibodies are acquired early in life and increase in prevalence with advancing age, so that >80% of adult populations have antibodies detectable by enzyme-linked immunosorbent assay (ELISA). Overall, coronaviruses account for 10–35% of common colds, depending on the season. Coronavirus infections appear to be particularly prevalent in late fall, winter, and early spring—times when rhinovirus infections are less common.

An extraordinary outbreak of the coronavirus-associated illness known as SARS occurred in 2002–2003. The outbreak apparently began in southern China and eventually resulted in 8096 recognized cases in 28 countries in Asia, Europe, and North and South America; ~90% of cases occurred in China and Hong Kong. The natural reservoir of SARS-CoV appeared to be the horseshoe bat, and the outbreak may have originated from human contact with infected semidomesticated animals such as the palm civet. In most cases, however, the infection was transmitted from human to human. Case–fatality rates varied among outbreaks, with an overall figure of ~9.5%. The disease appeared to be somewhat milder in cases in the United States and was clearly less severe among children. The outbreak ceased in 2003; 17 cases were detected in 2004, mostly in laboratory-associated settings, and no cases have been reported subsequently.

The mechanisms of transmission of SARS are incompletely understood. Clusters of cases suggest that spread may occur via both large- and small-droplet aerosols and perhaps via the fecal–oral route as well. The outbreak of illness in a large apartment complex in Hong Kong suggested that environmental sources, such as sewage or water, may also play a role in transmission. Some ill individuals (“super-spreaders”) appeared to be hyperinfectious and were capable of transmitting infection to 10–40 contacts, although most infections resulted in spread either to no one or to three or fewer individuals.

Since it began in June 2012, another extraordinary outbreak of serious respiratory illness, MERS, has been linked with a coronavirus (MERS-CoV). Through May 2014, a total of 536 cases and 145 deaths (27%) have been reported. All cases have been associated with contact or travel to six countries in or near the Arabian Peninsula: Jordan, Kuwait, Oman, Qatar, Saudi Arabia, and the United Arab Emirates. Cases have also been reported in France, Italy, Tunisia, Germany, Spain, and the United Kingdom. Person-to-person transmission has been documented, but sustained spread in communities has not. The source of MERS-CoV has not been established, but it is suspected that bats may be the animal reservoir and that camels serve as an intermediate host.

PATHOGENESIS

Coronaviruses that cause the common cold (e.g., strains HCoV-229E and HCoV-OC43) infect ciliated epithelial cells in the nasopharynx via the aminopeptidase N receptor (group 1) or a sialic acid receptor (group 2). Viral replication leads to damage of ciliated cells and induction of chemokines and interleukins, with consequent common-cold symptoms similar to those induced by rhinoviruses.

SARS-CoV infects cells of the respiratory tract via the angiotensin-converting enzyme 2 receptor. The result is a systemic illness in which virus is also found in the bloodstream, in the urine, and (for up to 2 months) in the stool. Virus persists in the respiratory tract for 2–3 weeks, and titers peak ~10 days after the onset of systemic illness. Pulmonary pathology consists of hyaline membrane formation, desquamation of pneumocytes in alveolar spaces, and an interstitial

infiltrate made up of lymphocytes and mononuclear cells. Giant cells are frequently seen, and coronavirus particles have been detected in type II pneumocytes. Elevated levels of proinflammatory cytokines and chemokines have been detected in sera from patients with SARS.

Because MERS-CoV was so recently detected, little is known at present about its pathogenesis. However, it may well be similar to that of SARS-CoV.

CLINICAL MANIFESTATIONS

After an incubation period that generally lasts 2–7 days (range, 1–14 days), SARS usually begins as a systemic illness marked by the onset of fever, which is often accompanied by malaise, headache, and myalgias and is followed in 1–2 days by a nonproductive cough and dyspnea. Approximately 25% of patients have diarrhea. Chest x-rays can show a variety of infiltrates, including patchy areas of consolidation—most frequently in peripheral and lower lung fields—or interstitial infiltrates, which can progress to diffuse involvement. In severe cases, respiratory function may worsen during the second week of illness and progress to frank adult respiratory distress syndrome accompanied by multiorgan dysfunction. Risk factors for severe disease include an age of >50 years and comorbidities such as cardiovascular disease, diabetes, and hepatitis. Illness in pregnant women may be particularly severe, but SARS-CoV infection appears to be milder in children than in adults.

Information regarding the clinical manifestations of MERS-CoV is limited. The case–fatality rate has been high in the initial cases, but this may represent an ascertainment bias, and it is clear that mild cases occur as well. The median incubation period has been estimated to be 5.2 days, and a secondary case was estimated to have an incubation period of 9–12 days. Cases have been reported that begin with cough and fever and progress to acute respiratory distress and respiratory failure within a week. Other cases have manifested as mild upper respiratory symptoms only. Renal failure has been noted, and DPP-4, the host cell receptor for MERS-CoV, is expressed at high levels in the kidney; these findings suggest that direct viral infection of the kidney may lead to renal dysfunction. Diarrhea and vomiting are also common in MERS, and pericarditis has been reported.

The clinical features of common colds caused by human coronaviruses are similar to those of illness caused by rhinoviruses. In studies of volunteers, the mean incubation period of colds induced by coronaviruses (3 days) is somewhat longer than that of illness caused by rhinoviruses, and the duration of illness is somewhat shorter (mean, 6–7 days). In some studies, the amount of nasal discharge was greater in colds induced by coronaviruses than in those induced by rhinoviruses. Coronaviruses other than SARS-CoV have been recovered occasionally from infants with pneumonia and from military recruits with lower respiratory tract disease and have been associated with worsening of chronic bronchitis. Two novel coronaviruses, HCoV-NL63 and HCoV-HKU1, have been isolated from patients hospitalized with acute respiratory illness. Their overall role as causes of human respiratory disease remains to be determined.

LABORATORY FINDINGS AND DIAGNOSIS

Laboratory abnormalities in SARS include lymphopenia, which is documented in ~50% of cases and which mostly affects CD4+ T cells but also involves CD8+ T cells and natural killer cells. Total white blood cell counts are normal or slightly low, and thrombocytopenia may develop as the illness progresses. Elevated serum levels of aminotransferases, creatine kinase, and lactate dehydrogenase have been reported.

A rapid diagnosis of SARS-CoV infection can be made by reverse-transcription PCR (RT-PCR) of respiratory tract samples and plasma early in the illness and of urine and stool later on. SARS-CoV can also be grown from respiratory tract samples by inoculation into Vero E6 tissue culture cells, in which a cytopathic effect is seen within days. RT-PCR appears to be more sensitive than tissue culture, but only around one-third of cases are positive by PCR at initial presentation. Serum antibodies can be detected by ELISA or immunofluorescence, and nearly all patients develop detectable serum antibodies within 28 days after the onset of illness.

Laboratory abnormalities in MERS-CoV infection include lymphopenia with or without neutropenia, thrombocytopenia, and elevated levels of lactate dehydrogenase. MERS-CoV can be isolated in tissue culture in Vero and LLC-MK2 cells, but PCR techniques are more sensitive and rapid and are the standard for laboratory diagnosis. Serologic tests using ELISA and immunofluorescence techniques have also been developed.

Laboratory diagnosis of coronavirus-induced colds is rarely required. Coronaviruses that cause those illnesses are frequently difficult to cultivate *in vitro* but can be detected in clinical samples by ELISA or immunofluorescence assays or by RT-PCR for viral RNA. These research procedures can be used to detect coronaviruses in unusual clinical settings.

TREATMENT CORONAVIRUS INFECTIONS

There is no specific therapy for SARS with established efficacy. Although ribavirin has frequently been used, it has little if any activity against SARS-CoV *in vitro*, and no beneficial effect on the course of illness has been demonstrated. Because of suggestions that immunopathology may contribute to the disease, glucocorticoids have also been widely used, but their benefit, if any, likewise remains to be established. Supportive care to maintain pulmonary and other organ-system functions remains the mainstay of therapy. Similarly, there is no established antiviral therapy for MERS. Interferon $\alpha 2b$ and ribavirin have displayed activity against MERS-CoV *in vitro* and in a rhesus macaque model, but data are not available on its use in human cases of MERS. The approach to the treatment of common colds caused by coronaviruses is similar to that discussed above for rhinovirus-induced illnesses.

PREVENTION



The recognition of SARS led to a worldwide mobilization of public health resources to apply infection control practices to contain the disease. Case definitions were established, travel advisories were proposed, and quarantines were imposed in certain locales. As of this writing, no additional cases of SARS have been reported since 2004. However, it remains unknown whether the disappearance of cases is a result of control measures, whether it is part of a seasonal or otherwise unexplained epidemiologic pattern of SARS, or when or whether SARS might reemerge. The U.S. Centers for Disease Control and Prevention (CDC) and the World Health Organization (WHO) maintain recommendations for surveillance and assessment of potential cases of SARS (www.cdc.gov/sars/). The frequent transmission of the disease to health care workers makes it mandatory that strict infection-control practices be employed by health care facilities to prevent airborne, droplet, and contact transmission from any suspected cases of SARS. Health care workers who enter areas in which patients with SARS may be present should don gowns, gloves, and eye and respiratory protective equipment (e.g., an N95 filtering facepiece respirator certified by the National Institute for Occupational Safety and Health).

Similarly, the WHO and the CDC have issued recommendations for identification, prevention, and control of MERS-CoV infections (www.cdc.gov/coronavirus/mers/index.html). Isolation precautions against airborne spread of infection should be instituted for patients hospitalized for suspected MERS, as described above for SARS.

Vaccines have been developed against several animal coronaviruses but not against known human coronaviruses. The emergence of SARS-CoV and MERS-CoV has stimulated interest in the development of vaccines against such agents.

HUMAN RESPIRATORY SYNCYTIAL VIRUS INFECTIONS

ETIOLOGIC AGENT

Human respiratory syncytial virus (HRSV) is a member of the Paramyxoviridae family (genus *Pneumovirus*). It is an enveloped virus ~150–350 nm in diameter and is so named because its replication *in vitro* leads to the fusion of neighboring cells into large multinucleated syncytia. The single-stranded RNA genome codes for 11 virus-specific proteins. Viral RNA is contained in a helical nucleocapsid surrounded by a lipid envelope bearing two glycoproteins: the G protein, by which the virus attaches to cells, and the F (fusion) protein, which facilitates entry of the virus into the cell by fusing host and viral membranes. HRSV is considered to be of a single antigenic type, but two distinct subgroups (A and B) and multiple subtypes within each subgroup have now been described. Antigenic diversity is reflected by differences in the G protein, whereas the F protein is relatively conserved. Both antigenic groups can circulate simultaneously in outbreaks, although there are typically alternating patterns in which one subgroup predominates over 1- to 2-year periods.

EPIDEMIOLOGY



HRSV is a major respiratory pathogen of young children and the foremost cause of lower respiratory disease in infants. Infection with HRSV is seen throughout the world in annual epidemics that occur in late fall, winter, or spring and last up to 5 months. The virus is rarely encountered during the summer. Rates of illness are highest among infants 1–6 months of age, peaking at 2–3 months of age. The attack rates among susceptible infants and children are extraordinarily high, approaching 100% in settings such as day-care centers where large numbers of susceptible infants are present. By age 2, virtually all children will have been infected with HRSV. HRSV accounts for 20–25% of hospital admissions of young infants and children for pneumonia and for up to 75% of cases of bronchiolitis in this age group. It has been estimated that more than half of infants who are at risk will become infected during an HRSV epidemic.

In older children and adults, reinfection with HRSV is frequent, but disease is milder than in infancy. A common cold–like syndrome is the illness most commonly associated with HRSV infection in adults. It has been increasingly appreciated that severe lower respiratory tract disease with pneumonitis can occur in elderly (often institutionalized) adults, in individuals with cardiopulmonary disease, and in patients with immunocompromising disorders or treatment, including recipients of hematopoietic stem cell transplants (HSCTs) and solid-organ transplants (SOTs). HRSV is also an important nosocomial pathogen; during an outbreak, it can infect pediatric patients and up to 25–50% of the staff on pediatric wards. The spread of HRSV among families is efficient: up to 40% of siblings may become infected when the virus is introduced into the family setting.

HRSV is transmitted primarily by close contact with contaminated fingers or fomites and by self-inoculation of the conjunctiva or anterior nares. Virus may also be spread by coarse aerosols produced by coughing or sneezing, but it is inefficiently spread by fine-particle aerosols. The incubation period is ~4–6 days, and virus shedding may last for ≥ 2 weeks in children and for shorter periods in adults. In immunosuppressed patients, shedding can continue for weeks.

PATHOGENESIS

Little is known about the histopathology of minor HRSV infection. Severe bronchiolitis or pneumonia is characterized by necrosis of the bronchiolar epithelium and a peribronchiolar infiltrate of lymphocytes and mononuclear cells. Inter-alveolar thickening and filling of alveolar spaces with fluid can also be found. The correlates of protective immunity to HRSV are incompletely understood. Because reinfection occurs frequently and is often associated with illness, the immunity that develops after single episodes of infection clearly is not complete or long-lasting. However, the cumulative effect of multiple reinfections is to temper subsequent disease and to provide some temporary measure of protection against infection. Studies of experimentally induced disease in healthy volunteers indicate that the presence of nasal IgA neutralizing antibody correlates more closely with protection than does the presence of serum antibody. Studies in infants, however, suggest that maternally acquired antibody provides some protection from lower respiratory tract disease, although illness can be severe even in infants who have moderate levels of maternally derived serum antibody. The relatively severe disease observed in immunosuppressed

1206 patients and experimental animal models indicates that cell-mediated immunity is an important mechanism of host defense against HRSV. Evidence suggests that major histocompatibility class I-restricted cytotoxic T cells may be particularly important in this regard.

CLINICAL MANIFESTATIONS

HRSV infection leads to a wide spectrum of respiratory illnesses. In infants, 25–40% of infections result in lower respiratory tract involvement, including pneumonia, bronchiolitis, and tracheobronchitis. In this age group, illness begins most frequently with rhinorrhea, low-grade fever, and mild systemic symptoms, often accompanied by cough and wheezing. Most patients recover gradually over 1–2 weeks. In more severe illness, tachypnea and dyspnea develop, and eventually frank hypoxia, cyanosis, and apnea can ensue. Physical examination may reveal diffuse wheezing, rhonchi, and rales. Chest radiography shows hyperexpansion, peribronchial thickening, and variable infiltrates ranging from diffuse interstitial infiltrates to segmental or lobar consolidation. Illness may be particularly severe in children born prematurely and in those with congenital cardiac disease, bronchopulmonary dysplasia, nephrotic syndrome, or immunosuppression. One study documented a 37% mortality rate among infants with HRSV pneumonia and congenital cardiac disease.

In adults, the most common symptoms of HRSV infection are those of the common cold, with rhinorrhea, sore throat, and cough. Illness is occasionally associated with moderate systemic symptoms such as malaise, headache, and fever. HRSV has also been reported to cause lower respiratory tract disease with fever in adults, including severe pneumonia in the elderly—particularly in nursing-home residents, among whom its impact can rival that of influenza. HRSV pneumonia can be a significant cause of morbidity and death among patients undergoing stem cell and solid organ transplantation, in whom case-fatality rates of 20–80% have been reported. Sinusitis, otitis media, and worsening of chronic obstructive and reactive airway disease have also been associated with HRSV infection.

LABORATORY FINDINGS AND DIAGNOSIS

The diagnosis of HRSV infection can be suspected on the basis of a suggestive epidemiologic setting—that is, severe illness among infants during an outbreak of HRSV in the community. Infections in older children and adults cannot be differentiated with certainty from those caused by other respiratory viruses. The specific diagnosis is established by detection of HRSV in respiratory secretions, such as sputum, throat swabs, or nasopharyngeal washes. Virus can be isolated in tissue culture, but this method has been largely supplanted by rapid viral diagnostic techniques consisting of immunofluorescence or ELISA of nasopharyngeal washes, aspirates, and (less satisfactorily) nasopharyngeal swabs. With specimens from children, these techniques have sensitivities and specificities of 80–95%; they are somewhat less sensitive with specimens from adults. RT-PCR detection techniques have shown even higher rates of sensitivity and specificity, particularly in adults. Serologic diagnosis may be made by comparison of acute- and convalescent-phase serum specimens by ELISA or by neutralization or complement-fixation tests. These tests may be useful in older children and adults but are less sensitive in children <4 months of age.

TREATMENT

HUMAN RESPIRATORY SYNCYTIAL VIRUS INFECTIONS

Treatment of upper respiratory tract HRSV infection is aimed primarily at the alleviation of symptoms and is similar to that for other viral infections of the upper respiratory tract. For lower respiratory tract infections, respiratory therapy, including hydration, suctioning of secretions, and administration of humidified oxygen and anticholinergic agents, is given as needed. In severe hypoxia, intubation and ventilatory assistance may be required. Studies of infants with HRSV infection who were given aerosolized ribavirin, a nucleoside analogue active in vitro against HRSV, demonstrated a modest beneficial effect on the resolution of lower respiratory tract illness,

including alleviation of blood-gas abnormalities, in some studies. The American Academy of Pediatrics does not recommend routine use of ribavirin but states that treatment with aerosolized ribavirin “may be considered” for infants who are severely ill or who are at high risk for complications of HRSV infection; included are premature infants and those with bronchopulmonary dysplasia, congenital heart disease, or immunosuppression. The efficacy of ribavirin against HRSV pneumonia in older children and adults, including those with immunosuppression, has not been established. No benefit has been found in the treatment of HRSV pneumonia with standard immunoglobulin; immunoglobulin with high titers of antibody to HRSV (RSVlg), which is no longer available; or chimeric mouse–human monoclonal IgG antibody to HRSV (palivizumab). Combined therapy with aerosolized ribavirin and palivizumab is being evaluated in immunosuppressed patients with HRSV pneumonia.

PREVENTION

Monthly administration of RSVlg (no longer available) or palivizumab has been approved as prophylaxis against HRSV for children <2 years of age who have bronchopulmonary dysplasia or cyanotic heart disease or who were born prematurely. Considerable interest exists in the development of vaccines against HRSV. Inactivated whole-virus vaccines have been ineffective; in one study, they actually potentiated disease in infants. Other approaches include immunization with purified F and G surface glycoproteins of HRSV or generation of stable live attenuated virus vaccines. In settings where rates of transmission are high (e.g., pediatric wards), barrier methods for the protection of hands and conjunctivae may be useful in reducing the spread of virus.

HUMAN METAPNEUMOVIRUS INFECTIONS

ETIOLOGIC AGENT

Human metapneumovirus (HMPV) is a viral respiratory pathogen that has been assigned to the Paramyxoviridae family (genus *Metapneumovirus*). Its morphology and genomic organization are similar to those of avian metapneumoviruses, which are recognized respiratory pathogens of turkeys. HMPV particles may be spherical, filamentous, or pleomorphic in shape and measure 150–600 nm in diameter. Particles contain 15-nm projections from the surface that are similar in appearance to those of other Paramyxoviridae. The single-stranded RNA genome codes for nine proteins that, except for the absence of nonstructural proteins, generally correspond to those of HRSV. HMPV is of only one antigenic type; two closely related genotypes (A and B), four subgroups, and two sublineages have been described.

EPIDEMIOLOGY

HMPV infections are worldwide in distribution, are most frequent during the winter in temperate climates, and occur early in life, so that serum antibodies to the virus are present in 50% of children by age 2 and in nearly all children by age 5. HMPV infections have been detected in older age groups, including elderly adults, and in both immunocompetent and immunosuppressed hosts. This virus accounts for 1–5% of childhood upper respiratory tract infections and for 10–15% of respiratory tract illnesses requiring hospitalization of children. In addition, HMPV causes 2–4% of acute respiratory illnesses in ambulatory adults and elderly patients. HMPV has been detected in a few cases of SARS, but its role (if any) in these illnesses has not been established.

CLINICAL MANIFESTATIONS

The spectrum of clinical illnesses associated with HMPV is similar to that associated with HRSV and includes both upper and lower respiratory tract illnesses, such as bronchiolitis, croup, and pneumonia. Reinfection with HMPV is common among older children and adults and has manifestations ranging from subclinical infections to common cold syndromes and occasionally pneumonia, which is seen primarily in elderly patients and those with cardiopulmonary diseases. Serious HMPV infections occur in



immunocompromised patients, including those with neoplasia, recipients of HSCTs, and children with HIV infection.

DIAGNOSIS

HMPV can be detected in nasal aspirates and respiratory secretions by immunofluorescence, by PCR (the most sensitive technique), or by growth in rhesus monkey kidney (LLC-MK2) tissue cultures. A serologic diagnosis can be made by ELISA, which uses HMPV-infected tissue culture lysates as sources of antigens.

TREATMENT HUMAN METAPNEUMOVIRUS INFECTIONS

Treatment for HMPV infections is primarily supportive and symptom-based. Ribavirin is active against HMPV *in vitro*, but its efficacy *in vivo* is unknown.

PREVENTION

Vaccines against HMPV are in the early stages of development.

PARAINFLUENZA VIRUS INFECTIONS

ETIOLOGIC AGENT

Parainfluenza viruses belong to the Paramyxoviridae family (genera *Respirovirus* and *Rubulavirus*). They are 150–200 nm in diameter, are enveloped, and contain a single-stranded RNA genome. The envelope is studded with two glycoproteins: one possesses both hemagglutinin and neuraminidase activity, and the other contains fusion activity. The viral RNA genome is enclosed in a helical nucleocapsid and codes for six structural and several accessory proteins. All types of parainfluenza virus (1, 2, 3, 4A, and 4B) share certain antigens with other members of the Paramyxoviridae family, including mumps and Newcastle disease viruses.

EPIDEMIOLOGY



Parainfluenza viruses are distributed throughout the world; infection with serotypes 4A and 4B has been reported less widely, probably because these types are more difficult than the other three to grow in tissue culture. Infection is acquired in early childhood; by 5 years of age, most children have antibodies to serotypes 1, 2, and 3. Types 1 and 2 cause epidemics during the fall, often occurring in an alternate-year pattern. Type 3 infection has been detected during all seasons, but epidemics have occurred annually in the spring.

The contribution of parainfluenza infections to respiratory disease varies with both the location and the year. In studies conducted in the United States, parainfluenza virus infections have accounted for 4.3–22% of respiratory illnesses in children. The major importance of these viruses is as a cause of lower respiratory illness in young children, in whom they rank second only to HRSV in that regard. Parainfluenza virus type 1 is the most common cause of croup (laryngotracheobronchitis) in children, whereas serotype 2 causes similar, although generally less severe, disease. Type 3 is an important cause of bronchiolitis and pneumonia in infants, whereas illnesses associated with types 4A and 4B have generally been mild. Unlike types 1 and 2, type 3 frequently causes illness during the first month of life, when passively acquired maternal antibody is still present. Parainfluenza viruses are spread through infected respiratory secretions, primarily by person-to-person contact and/or by large droplets, and by contact with fomites contaminated with respiratory secretions. The incubation period has varied from 3 to 6 days in experimental infections but may be somewhat shorter for naturally occurring disease in children.

In adults, parainfluenza virus infections are generally mild and account for fewer than 10% of respiratory illnesses. The advent of contemporary laboratory methods for diagnosis has increased awareness of the impact of parainfluenza infections in adults. In a recent study, parainfluenza virus was the third most common viral isolate from patients 16–64 years old who required hospitalization (0.7 isolate/1000 population). In the 2009 influenza pandemic, parainfluenza virus type 3 was the second most common cause of illness after influenza virus.

PATHOGENESIS

Immunity to parainfluenza viruses is incompletely understood, but evidence suggests that immunity to infections with serotypes 1 and 2 is mediated by local IgA antibodies in the respiratory tract. Passively acquired serum neutralizing antibodies also confer some protection against infection with types 1, 2, and (to a lesser degree) 3. Studies in experimental animal models and in immunosuppressed patients suggest that T cell-mediated immunity may also be important in parainfluenza virus infections. Lack of cellular immune responses is associated with an increased risk of progressive and fatal disease in HSCT recipients.

CLINICAL MANIFESTATIONS

Parainfluenza virus infections occur most frequently among children, in whom initial infection with serotype 1, 2, or 3 is associated with an acute febrile illness in 50–80% of cases. Children may present with coryza, sore throat, hoarseness, and cough that may or may not be croupy. In severe croup, fever persists, with worsening coryza and sore throat. A brassy or barking cough may progress to frank stridor. Most children recover over the next 1 or 2 days, although progressive airway obstruction and hypoxia ensue occasionally. If bronchiolitis or pneumonia develops, progressive cough accompanied by wheezing, tachypnea, and intercostal retractions may occur. In this setting, sputum production increases modestly. Physical examination documents nasopharyngeal discharge and oropharyngeal injection, along with rhonchi, wheezes, or coarse breath sounds. Chest x-rays can show air trapping and occasionally interstitial infiltrates.

In older children and adults, parainfluenza infections tend to be milder, presenting most frequently as a common cold or as hoarseness, with or without cough. Lower respiratory tract involvement in older children and adults is uncommon, although tracheobronchitis and community-acquired pneumonia have been reported in adults.

Parainfluenza viruses, most frequently type 3, are important pathogens in immunosuppressed patients—particularly in HSCT recipients but also in SOT recipients (especially recipients of lung transplants). Patients receiving cancer chemotherapy are also at risk for severe parainfluenza infection. Severe, prolonged, and even fatal parainfluenza-associated respiratory illnesses have been reported in children and adults with severe immunosuppression.

LABORATORY FINDINGS AND DIAGNOSIS

The clinical syndromes caused by parainfluenza viruses (with the possible exception of croup in young children) are not sufficiently distinctive to be diagnosed on clinical grounds alone. A specific diagnosis is established by detection of virus in respiratory tract secretions, throat swabs, or nasopharyngeal washings. Growth of the virus in tissue culture is detected either by hemagglutination or by a cytopathic effect. A rapid diagnosis may be made by identification of parainfluenza antigens in exfoliated cells from the respiratory tract with immunofluorescence or ELISA, although these techniques appear to be less sensitive than tissue culture. Highly specific and sensitive PCR assays have also been developed and have now become the standard for viral diagnosis. Serologic diagnosis can be established by hemagglutination-inhibition, complement-fixation, or neutralization testing of acute- and convalescent-phase specimens. However, because frequent heterotypic responses occur among the parainfluenza serotypes, the serotype causing illness often cannot be identified by serologic techniques alone.

Acute epiglottitis caused by *Haemophilus influenzae* type b must be differentiated from viral croup. Influenza A virus is also a common cause of croup during epidemic periods.

TREATMENT PARAINFLUENZA VIRUS INFECTIONS

For upper respiratory tract illness, symptoms can be treated as discussed for other viral respiratory tract illnesses. If complications such as sinusitis, otitis, or superimposed bacterial bronchitis develop, appropriate antibacterial drugs should be administered. Mild cases of croup should be treated with bed rest and moist air generated by vaporizers. More severe cases require hospitalization

1208 and close observation for the development of respiratory distress. If acute respiratory distress develops, humidified oxygen and intermittent racemic epinephrine are usually administered. Aerosolized or systemically administered glucocorticoids are beneficial; systemic administration has a more profound effect. No specific antiviral therapy has been established. Ribavirin is active against parainfluenza viruses in vitro, and anecdotal reports describe its use clinically, particularly in immunosuppressed patients, but its efficacy, if any, is unclear. DAS181, a sialidase with activity against parainfluenza viruses, is undergoing evaluation in immunosuppressed patients.

PREVENTION

Vaccines against parainfluenza viruses are under development.

ADENOVIRUS INFECTIONS

ETIOLOGIC AGENT

Adenoviruses are complex DNA viruses that measure 70–80 nm in diameter. Human adenoviruses belong to the genus *Mastadenovirus*, which includes 51 serotypes. Adenoviruses have a characteristic morphology consisting of an icosahedral shell composed of 20 equilateral triangular faces and 12 vertices. The protein coat (capsid) consists of hexon subunits with group-specific and type-specific antigenic determinants and penton subunits at each vertex primarily containing group-specific antigens. A fiber with a knob at the end projects from each penton; this fiber contains type-specific and some group-specific antigens. Human adenoviruses have been divided into seven subgroups (A through G) on the basis of the homology of DNA genomes and other properties. Revised criteria for classifying human adenoviruses have been proposed; reflecting recent approaches to the characterization of novel adenoviruses, the revised criteria include genome sequence and computational analysis in addition to traditional serologic criteria. The adenovirus genome is a linear double-stranded DNA that codes for structural and nonstructural polypeptides. The replicative cycle of adenovirus may result either in lytic infection of cells or in the establishment of a latent infection (primarily involving lymphoid cells). Some adenovirus types can induce oncogenic transformation, and tumor formation has been observed in rodents; however, despite intensive investigation, adenoviruses have not been associated with tumors in humans.

EPIDEMIOLOGY

Adenovirus infections most frequently affect infants and children. Infections occur throughout the year but are most common from fall to spring. In the United States, adenoviruses account for ~10% of acute respiratory infections in children but for <2% of respiratory illnesses in civilian adults. Nearly 100% of adults have serum antibody to multiple serotypes—a finding indicating that infection is common in childhood. Types 1, 2, 3, and 5 are the most common isolates from children. Certain adenovirus serotypes—particularly 4 and 7 but also 3, 14, and 21—are associated with outbreaks of acute respiratory disease in military recruits. Clusters of particularly severe disease have been seen with adenovirus 14.

Adenovirus infection can be transmitted by inhalation of aerosolized virus, by inoculation of virus into conjunctival sacs, and probably by the fecal-oral route as well. Type-specific antibody generally develops after infection and is associated with protection—albeit incomplete—against infection with the same serotype.

CLINICAL MANIFESTATIONS

In children, adenoviruses cause a variety of clinical syndromes. The most common is an acute upper respiratory tract infection, with prominent rhinitis. On occasion, lower respiratory tract disease, including bronchiolitis and pneumonia, also develops. Adenoviruses, particularly types 3 and 7, cause pharyngoconjunctival fever, a characteristic acute febrile illness of children that occurs in outbreaks, most often in summer camps. The syndrome is marked by bilateral conjunctivitis in which the bulbar and palpebral conjunctivae have a granular appearance. Low-grade fever is frequently present for the first

3–5 days, and rhinitis, sore throat, and cervical adenopathy develop. The illness generally lasts for 1–2 weeks and resolves spontaneously. Febrile pharyngitis without conjunctivitis has also been associated with adenovirus infection. Adenoviruses have been isolated from cases of whooping cough with or without *Bordetella pertussis*; the significance of adenovirus in that disease is unknown.

In adults, the most frequently reported illness has been acute respiratory disease caused by adenovirus types 4 and 7 in military recruits. This illness is marked by a prominent sore throat and the gradual onset of fever, which often reaches 39°C (102.2°F) on the second or third day of illness. Cough is almost always present, and coryza and regional lymphadenopathy are frequently seen. Physical examination may show pharyngeal edema, injection, and tonsillar enlargement with little or no exudate. If pneumonia has developed, auscultation and x-ray of the chest may indicate areas of patchy infiltration.

Adenoviruses have been associated with a number of non-respiratory tract diseases, including acute diarrheal illness caused by types 40 and 41 in young children and hemorrhagic cystitis caused by types 11 and 21. Epidemic keratoconjunctivitis, caused most frequently by types 8, 19, and 37, has been associated with contaminated common sources such as ophthalmic solutions and roller towels. Adenoviruses have also been implicated in disseminated disease and pneumonia in immunosuppressed patients, including recipients of SOTs or HSCTs. In HSCT recipients, adenovirus infections have manifested as pneumonia, hepatitis, nephritis, colitis, encephalitis, and hemorrhagic cystitis. In SOT recipients, adenovirus infection may involve the organ transplanted (e.g., hepatitis in liver transplants, nephritis in renal transplants) but can disseminate to other organs as well. In patients with AIDS, high-numbered and intermediate adenovirus serotypes have been isolated, usually in the setting of low CD4+ T cell counts, but their isolation often has not been clearly linked to disease manifestations. Adenovirus nucleic acids have been detected in myocardial cells from patients with “idiopathic” cardiomyopathies, and adenoviruses have been suggested as causative agents in some cases.

LABORATORY FINDINGS AND DIAGNOSIS

Adenovirus infection should be suspected in the epidemiologic setting of acute respiratory disease in military recruits and in certain clinical syndromes (such as pharyngoconjunctival fever or epidemic keratoconjunctivitis) in which outbreaks of characteristic illnesses occur. In most cases, however, illnesses caused by adenovirus infection cannot be differentiated from those caused by a number of other viral respiratory agents and *Mycoplasma pneumoniae*. A definitive diagnosis of adenovirus infection is established by detection of the virus in tissue culture (as evidenced by cytopathic changes) and by specific identification with immunofluorescence or other immunologic techniques. Rapid viral diagnosis can be established by immunofluorescence or ELISA of nasopharyngeal aspirates, conjunctival or respiratory secretions, urine, or stool. Highly sensitive and specific PCR assays and nucleic acid hybridization are available and have become the standard for diagnosis based on clinical specimens. Adenovirus types 40 and 41, which have been associated with diarrheal disease in children, require special tissue-culture cells for isolation, and these serotypes are most commonly detected by direct ELISA of stool or by PCR. Serum antibody rises can be demonstrated by complement-fixation or neutralization tests, ELISA, radioimmunoassay, or (for those adenoviruses that hemagglutinate red cells) hemagglutination-inhibition tests.

TREATMENT ADENOVIRUS INFECTIONS

Only symptom-based treatment and supportive therapy are available for adenovirus infections, and clinically useful antiviral therapy has not been established. Ribavirin and cidofovir are active in vitro against certain adenoviruses. Retrospective studies and anecdotes describe the use of these agents in disseminated adenovirus infections, but definitive efficacy data from controlled studies are not available. An oral liposomal form of cidofovir (CMX001) is being evaluated for adenovirus infections in immunosuppressed patients.

PREVENTION

Live vaccines have been developed against adenovirus types 4 and 7 and have been highly efficacious in control of acute respiratory disease among military recruits. These vaccines consist of live, unattenuated virus administered in enteric-coated capsules. Infection of the gastrointestinal tract with types 4 and 7 does not cause disease but stimulates local and systemic antibodies that are protective against subsequent acute respiratory disease due to those serotypes. These vaccines were not produced from 1999 to 2011 but are now available again and are being used effectively in military recruits. Adenoviruses are also being studied as live-virus vectors for the delivery of vaccine antigens and for gene therapy.

224 Influenza

Yehuda Z. Cohen, Raphael Dolin

DEFINITION

Influenza is an acute respiratory illness caused by infection with influenza viruses. The illness affects the upper and/or lower respiratory tract and is often accompanied by systemic signs and symptoms such as fever, headache, myalgia, and weakness. Outbreaks of illness of variable extent and severity occur nearly every year. Such outbreaks result in significant morbidity rates in the general population and in increased mortality rates among certain high-risk patients, mainly as a result of pulmonary complications.

ETIOLOGIC AGENT

Influenza viruses are members of the Orthomyxoviridae family, of which influenza A, B, and C viruses constitute three separate genera. The designation of influenza viruses as type A, B, or C is based on antigenic characteristics of the nucleoprotein (NP) and matrix (M) protein antigens. Influenza A viruses are further subdivided (subtyped) on the basis of the surface hemagglutinin (H) and neuraminidase (N) antigens; individual strains are designated according to the site of origin, isolate number, year of isolation, and subtype—for example, influenza A/California/07/2009 (H1N1). Influenza A has 18 distinct H subtypes and 11 distinct N subtypes, of which only H1, H2, H3, N1, and N2 have been associated with epidemics of disease in humans. Avian influenza A viruses have been associated with small outbreaks and sporadic cases in humans (see below). Influenza B and C viruses are designated similarly to influenza A viruses, but H and N antigens from these viruses do not receive subtype designations because intratypic variations in influenza B antigens are less extensive than those in influenza A viruses and may not occur with influenza C virus.

Influenza A and B viruses are major human pathogens and the most extensively studied of the Orthomyxoviridae. Type A and type B viruses are morphologically similar. The virions are irregularly shaped spherical particles, measure 80–120 nm in diameter, and have a lipid envelope from the surface of which the H and N glycoproteins project (Fig. 224-1). The hemagglutinin is the site by which the virus binds to sialic acid cell receptors, whereas the neuraminidase degrades the receptor and plays a role in the release of the virus from infected cells after replication has taken place. Influenza viruses enter cells by receptor-mediated endocytosis, forming a virus-containing endosome. The viral hemagglutinin mediates fusion of the endosomal membrane with the virus envelope, and viral nucleocapsids are subsequently released into the cytoplasm. Immune responses to the H antigen are the major determinants of protection against infection with influenza virus, whereas those to the N antigen limit viral spread and contribute to reduction of the infection. The lipid envelope of influenza A virus also contains the M proteins M1 and M2, which are involved in stabilization of the lipid envelope and in virus assembly. The virion also contains the NP antigen, which is associated with the viral genome, as well as three polymerase (P) proteins that are essential for transcription

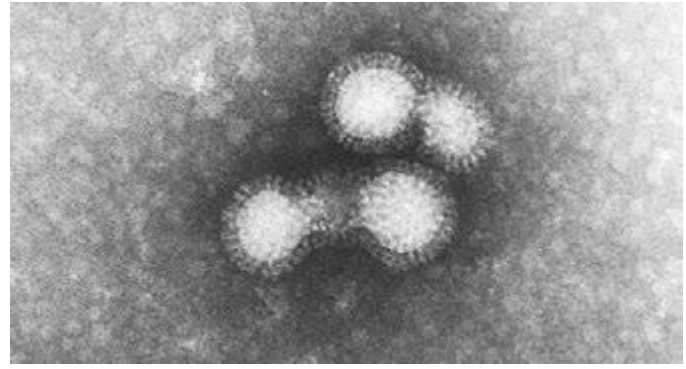



FIGURE 224-1 An electron micrograph of influenza A virus (×40,000).

and synthesis of viral RNA. Two nonstructural proteins function as an interferon antagonist and posttranscriptional regulator (NS1) and a nuclear export factor (NS2 or NEP).

The genomes of influenza A and B viruses consist of eight single-strand RNA segments, which code for the structural and nonstructural proteins. Because the genome is segmented, the opportunity for gene reassortment during infection is high; reassortment often takes place during infection of cells with more than one influenza A virus.

EPIDEMIOLOGY

 Influenza outbreaks occur virtually every year, although their extent and severity vary widely. Localized outbreaks take place at variable intervals, usually every 1–3 years. Global pandemics have occurred at variable intervals, but much less frequently than inter-pandemic outbreaks (Table 224-1). The most recent pandemic emerged in March of 2009 and was caused by an influenza A/H1N1 virus that rapidly spread worldwide over the next several months.

Influenza A Virus • ANTIGENIC VARIATION AND INFLUENZA OUTBREAKS AND PANDEMICS The most extensive and severe outbreaks of influenza are caused by influenza A viruses, in part because of the remarkable propensity of the H and N antigens of these viruses to undergo periodic antigenic variation. Major antigenic variations, called *antigenic shifts*, are seen only with influenza A viruses and may be associated with pandemics. Minor variations are called *antigenic drifts*. Antigenic variation may involve the hemagglutinin alone or both the hemagglutinin and the neuraminidase. An example of an antigenic shift involving both the hemagglutinin and the neuraminidase is that of 1957, when the predominant influenza A virus subtype shifted from H1N1 to H2N2; this shift resulted in a severe pandemic, with an estimated 70,000 excess deaths (i.e., deaths in excess of the number expected without an influenza epidemic) in the United States alone. This excess mortality was significantly greater than that during inter-pandemic influenza seasons.

TABLE 224-1 EMERGENCE OF ANTIGENIC SUBTYPES OF INFLUENZA A VIRUS ASSOCIATED WITH PANDEMIC OR EPIDEMIC DISEASE

Years	Subtype	Extent of Outbreak
1889–1890	H2N8 ^a	Severe pandemic
1900–1903	H3N8 ^a	?Moderate epidemic
1918–1919	H1N1 ^b (formerly HswN1)	Severe pandemic
1933–1935	H1N1 ^b (formerly HON1)	Mild epidemic
1946–1947	H1N1	Mild epidemic
1957–1958	H2N2	Severe pandemic
1968–1969	H3N2	Moderate pandemic
1977–1978 ^c	H1N1	Mild pandemic
2009–2010 ^d	H1N1	Pandemic

^a As determined by retrospective serologic survey of individuals alive during those years (“seroarchaeology”). ^b Hemagglutinins formerly designated as Hsw and H0 are now classified as variants of H1. ^c From this time until 2008–2009, viruses of the H1N1 and H3N2 subtypes circulated either in alternating years or concurrently. ^d A novel influenza A/H1N1 virus emerged to cause this pandemic.


1210 In 1968, an antigenic shift involving only the hemagglutinin occurred (H2N2 to H3N2); the subsequent pandemic was less severe than that of 1957. In 1977, an H1N1 virus emerged and caused a pandemic that primarily affected younger individuals (i.e., those born after 1957). As shown in Table 224-1, H1N1 viruses circulated from 1918 to 1956; thus, individuals born prior to 1957 would be expected to have some degree of immunity to H1N1 viruses. The pandemic of 2009–2010 was caused by an A/H1N1 virus against which little immunity was present in the general population, although approximately one-third of individuals born before 1950 had some apparent immunity to related H1N1 strains.

During most outbreaks of influenza A, a single subtype has circulated at a time. However, since 1977, H1N1 and H3N2 viruses have circulated simultaneously, resulting in outbreaks of varying severity. In some outbreaks, influenza B viruses have also circulated simultaneously with influenza A viruses. In 2009–2010, the pandemic A/H1N1 virus appeared to circulate nearly exclusively.

FEATURES OF PANDEMIC AND INTERPANDEMIC INFLUENZA A Pandemics provide the most dramatic evidence of the impact of influenza A. However, illnesses occurring between pandemics (interpandemic disease) also account for extensive mortality and morbidity, albeit over a longer period. In the United States, influenza was associated with an average of 23,000 excess deaths per season in 1976–2007 and with a maximum of 48,600 excess deaths during the 2003–2004 season.

Influenza A viruses that circulate between pandemics demonstrate antigenic drifts in the H antigen. These antigenic drifts result from point mutations in the RNA segment that codes for the hemagglutinin and occur most frequently in five hypervariable regions. Epidemiologically significant strains—that is, those with the potential to cause widespread outbreaks—exhibit changes in amino acids in at least two of the major antigenic sites in the hemagglutinin molecule. Because two point mutations are unlikely to occur simultaneously, it is believed that antigenic drifts result from point mutations occurring sequentially during the spread of virus from person to person. Antigenic drifts have been reported nearly annually since 1977 for H1N1 viruses and since 1968 for H3N2 viruses.

Interpandemic influenza A outbreaks usually begin abruptly, peak over a 2- to 3-week period, generally last for 2–3 months, and often subside almost as rapidly as they began. In contrast, pandemic influenza may begin with rapid transmission at multiple locations, have high attack rates, and extend beyond the usual seasonality, with multiple waves of attack before or after the main outbreak. In interpandemic outbreaks, the first indication of influenza activity is an increase in the number of children with febrile respiratory illnesses who present for medical attention. This increase is followed by increases in rates of influenza-like illnesses among adults and eventually by an increase in hospital admissions for patients with pneumonia, worsening of congestive heart failure, and exacerbations of chronic pulmonary disease. Rates of absence from work and school also rise at this time. An increase in the number of deaths caused by pneumonia and influenza is generally a late observation in an outbreak. Attack rates have been highly variable from outbreak to outbreak in interpandemic influenza but most commonly are in the range of 10–20% of the general population.

 Although pandemic influenza may occur throughout the year, interpandemic influenza occurs almost exclusively during the winter months in the temperate zones of the Northern and Southern hemispheres. In those locations, it is highly unusual to detect influenza A virus at other times, although rises in serum antibody titer or even outbreaks have been noted rarely during warm-weather months. In contrast, influenza virus infections occur throughout the year in the tropics. Where or how influenza A viruses persist between outbreaks in temperate zones is unknown. It is possible that the viruses are maintained in the human population on a worldwide basis by person-to-person transmission and that large population clusters support a low level of interepidemic transmission. Alternatively, human strains may persist in animal reservoirs. Convincing evidence to support either explanation is not available. In the modern era, rapid transportation may contribute to the transmission of viruses among widespread geographic locales.

The factors that result in the inception and termination of outbreaks of influenza A are incompletely understood. A major determinant of the extent and severity of an outbreak is the level of immunity in the population at risk. With the emergence of an antigenically novel influenza virus to which little or no immunity is present in a community, extensive outbreaks may occur. When the absence of immunity is worldwide, epidemic disease may spread around the globe, resulting in a pandemic. Such pandemic waves can continue for several years, until immunity in the population reaches a high level. In the years following pandemic influenza, antigenic drifts among influenza viruses result in outbreaks of variable severity in populations with high levels of immunity to the pandemic strain that circulated earlier. This situation persists until another antigenically novel pandemic strain emerges. On the other hand, outbreaks sometimes end despite the persistence of a large pool of susceptible individuals in the population. It has been suggested that certain influenza A viruses may be intrinsically less virulent and cause less severe disease than other variants, even in immunologically virgin subjects. If so, then other (undefined) factors besides the level of preexisting immunity must play a role in the epidemiology of influenza.

Avian and Swine Influenza Viruses Aquatic birds are the largest reservoir of influenza A viruses, harboring 16 hemagglutinin (H1–H16) and nine neuraminidase (N1–N9) subtypes. (In addition, H17N10 and H18N11 viruses are found in bats.) Influenza A pandemic strains in 1957 (A/H2N2) and in 1968 (A/H3N2) resulted from reassortment of gene segments between human and avian viruses. The influenza A/H1N1 virus that caused the most severe pandemic of modern times (1918–1919) appears to have been an adaptation of an avian virus to human infection. Thus, there is concern that avian influenza viruses with novel hemagglutinin and neuraminidase antigens have the potential to emerge as pandemic strains.



Avian influenza A viruses have been reported to cause sporadic cases and small outbreaks in humans, usually after direct contact with birds (most commonly poultry). Sustained person-to-person transmission in the community has not been observed. Avian influenza A/H5N1 virus has been noted to cause illness in humans since 1997, with 648 cases reported to the World Health Organization as of January 2014. It is not clear whether the high observed case–fatality rate (59%) reflects preferential detection of severe cases. A/H7N7 infections have been noted in poultry industry workers; conjunctivitis was the most prominent feature, although a minority of individuals also had respiratory illness. More than 333 cases of avian A/H7N9 infection have been reported in China, with case–fatality rates of 36% among the infected patients admitted to the hospital. Most H7N9 isolates are sensitive to neuraminidase inhibitors, but a few isolates have exhibited high-level resistance to oseltamivir and diminished sensitivity to zanamivir. Infections with avian H9N2 viruses have been reported primarily among children in Hong Kong and have consisted largely of mild respiratory illnesses. Mild cases of illness due to influenza H10N7 virus in Egypt and Australia have also been reported. In 2013, the first cases of human infection with avian A/H10N8 and H6N1 viruses were described.

Influenza A viruses also circulate in swine but rarely infect humans. Whereas humans primarily have α -2,6-galactose receptors for hemagglutinins and birds primarily have α -2,3-galactose receptors, swine have both types of receptors. Thus, swine hosts efficiently sustain simultaneous infection with both human and avian viruses, thereby facilitating reassortment of genetic segments between viruses of both species. The pandemic A/H1N1 strain of 2009–2010 was a quadruple reassortant among swine, avian, and human influenza viruses. The influenza A virus subtypes that circulate most commonly in swine are H1N1, H1N2, and H3N2. When a predominantly swine virus causes infections in humans, it is designated a variant virus by the addition of “v” after the subtype. For example, influenza A/H3N2v virus was responsible for 321 cases of human infection reported in the United States in 2011 and 2012 and for 18 cases in 2013. Almost all of the affected patients had had close contact with swine. Only limited person-to-person transmission of swine influenza virus has been

noted. Since 2005, 16 human cases caused by A/H1N1v virus and 5 caused by A/H1N2v virus have been detected in the United States.

Influenza B and C Viruses Influenza B virus causes outbreaks that are generally less extensive and are associated with less severe disease than those caused by influenza A virus, although the disease may occasionally be severe. The hemagglutinin and neuraminidase of influenza B viruses undergo less frequent and less extensive variation than those of influenza A viruses; this characteristic may account, in part, for the lesser severity of influenza B. Outbreaks of influenza B occur most frequently in schools and military camps, although outbreaks in institutions in which elderly individuals reside have also been noted on occasion. Since the 1980s, two antigenically distinct “lineages” of influenza B virus have circulated: Victoria and Yamagata.

In contrast to influenza A and B viruses, influenza C virus appears to be a relatively minor cause of disease in humans. It has been associated with common cold-like symptoms and occasionally with lower respiratory tract illness. The widespread prevalence of serum antibody to this virus indicates that asymptomatic infection may be common.

Influenza-Associated Morbidity and Mortality Rates Rates of morbidity and mortality caused by influenza outbreaks continue to be substantial. Most individuals who die in this setting have underlying diseases that place them at high risk for complications of influenza (Table 224-2). On average, there were 226,000 influenza-associated hospitalizations per year in the United States in 1979–2001. Recently, the moderately severe influenza season in 2012–2013 was associated with 381,500 hospitalizations (42 per 100,000 persons). Excess annual hospitalizations for groups of adults and children with high-risk medical conditions ranged from 40 to 1900 per 100,000 during outbreaks of influenza in 1973–2004. The most prominent high-risk conditions are chronic cardiac and pulmonary diseases and old age. Mortality rates among individuals with chronic metabolic or renal diseases or certain immunosuppressive diseases have also been elevated, although they remain lower than mortality rates among patients with chronic cardiopulmonary diseases. In the pandemic of 2009–2010, increased risk of severe disease was noted in children from birth to 4 years of age and in pregnant women. The morbidity rate attributable to influenza in the general population is considerable. It is estimated that interepidemic outbreaks of influenza currently incur annual economic costs of more than \$87 billion in the United States. For pandemics, it is estimated that annual economic costs would range from \$89.7 to \$209.4 billion for attack rates of 15–35%.

PATHOGENESIS AND IMMUNITY

The initial event in influenza is infection of the respiratory epithelium with influenza virus acquired from respiratory secretions of acutely infected individuals. In all likelihood, the virus is transmitted via aerosols generated by coughs and sneezes, although transmission through hand-to-hand contact, other personal contact, and even fomites may take place. Experimental evidence suggests that infection by a small-particle aerosol (particle diameter <10 μm) is more efficient than

that by larger droplets. Initially, viral infection involves the ciliated columnar epithelial cells, but it may also involve other respiratory tract cells, including alveolar cells, mucous gland cells, and macrophages. In infected cells, virus replicates within 4–6 h, after which infectious virus is released to infect adjacent or nearby cells. In this way, infection spreads from a few foci to a large number of respiratory cells over several hours. In experimentally induced infection, the incubation period of illness has ranged from 18 to 72 h, depending on the size of the viral inoculum. Histopathologic study reveals degenerative changes, including granulation, vacuolization, swelling, and pyknotic nuclei in infected ciliated cells. The cells eventually become necrotic and desquamate; in some areas, previously columnar epithelium is replaced by flattened and metaplastic epithelial cells. The severity of illness is correlated with the quantity of virus shed in secretions; thus, the degree of viral replication itself may be an important factor in pathogenesis. Despite the frequent development of systemic signs and symptoms such as fever, headache, and myalgias, influenza virus has only rarely been detected in extrapulmonary sites (including the bloodstream). Evidence suggests that the pathogenesis of systemic symptoms in influenza may be related to the induction of certain cytokines, particularly tumor necrosis factor α , interferon α , interleukin 6, and interleukin 8, in respiratory secretions and in the bloodstream.

The host response to influenza infections involves a complex interplay of humoral antibody, local antibody, cell-mediated immunity, interferon, and other host defenses. Serum antibody responses, which can be detected by the second week after primary infection, are measured by a variety of techniques: hemagglutination inhibition (HI), complement fixation (CF), neutralization, enzyme-linked immunosorbent assay (ELISA), and antineuraminidase antibody assay. Antibodies to the hemagglutinin appear to be the most important mediators of immunity; in several studies, HI titers of ≥ 40 have been associated with protection from infection. Secretory antibodies produced in the respiratory tract are predominantly of the IgA class, and secretory antibody neutralization titers of ≥ 4 have also been associated with protection. A variety of cell-mediated immune responses, both antigen-specific and antigen-nonspecific, can be detected early after infection and depend on the prior immune status of the host. These responses include T cell proliferative, T cell cytotoxic, and natural killer cell activity. In humans, CD8+ as well as CD4+ T lymphocytes are directed at conserved regions of internal proteins (NP, M, and P) as well as at the surface proteins H and N. Interferons can be detected in respiratory secretions shortly after the shedding of virus has begun, and rises in interferon titers coincide with decreases in virus shedding.

The host defense factors responsible for cessation of virus shedding and resolution of illness have not been defined specifically. Virus shedding generally stops within 2–5 days after symptoms first appear, at a time when serum and local antibody responses often are not detectable by conventional techniques, although antibody rises may be detected earlier by use of highly sensitive techniques, particularly in individuals with previous immunity to the virus. It has been suggested that interferon, cell-mediated immune responses, and/or nonspecific inflammatory responses all contribute to the resolution of illness. CD8+ cytotoxic T lymphocyte responses may be particularly important in this regard.

CLINICAL MANIFESTATIONS

Influenza is most frequently described as a respiratory illness characterized by systemic symptoms, such as headache, feverishness, chills, myalgia, and malaise, as well as accompanying respiratory tract signs and symptoms, particularly cough and sore throat. In some cases, the onset is so abrupt that patients can recall the precise time they became ill. However, the spectrum of clinical presentations is wide, ranging from a mild, afebrile respiratory illness similar to the common cold (with either a gradual or an abrupt onset) to severe prostration with relatively few respiratory signs and symptoms. In most of the cases that come to a physician's attention, the patient has a fever, with temperatures of 38°–41°C (100.4°–105.8°F). A rapid temperature rise within the first 24 h of illness is generally followed by gradual defervescence over 2–3 days, although, on occasion, fever may last as long as 1 week.

TABLE 224-2 PERSONS AT HIGHER RISK FOR COMPLICATIONS OF INFLUENZA OR FOR INFLUENZA-RELATED VISITS TO HEALTH CARE FACILITIES

All children from birth to <5 years, especially <2 years
All persons ≥ 50 years old
Pregnant women
Adults and children who have chronic pulmonary (including asthma) or cardiovascular (except isolated hypertension), renal, hepatic, neurologic, hematologic, or metabolic disorders (including diabetes mellitus)
Persons who have immunosuppression (including that caused by medications or by HIV infection)
Children and adolescents (6 months to 18 years old) who are receiving long-term aspirin therapy and who might be at risk for Reye's syndrome after influenza virus infection
Residents of nursing homes and other long-term care facilities
Native Americans/Alaska Natives
Persons who are morbidly obese (body mass index ≥ 40 kg/m ²)

1212 Patients report a feverish feeling and chilliness, but true rigors are rare. Headache, either generalized or frontal, is often particularly troublesome. Myalgias may involve any part of the body but are most common in the legs and lumbosacral area. Arthralgias may also develop.

Respiratory symptoms often become more prominent as systemic symptoms subside. Many patients have a sore throat or persistent cough, which may last for ≥ 1 week and which is often accompanied by substernal discomfort. Ocular signs and symptoms include pain on motion of the eyes, photophobia, and burning of the eyes.

In the elderly, influenza may have a relatively subtle presentation. Typical features such as sore throat, myalgia, and even fever may be absent, and general symptoms such as anorexia, malaise, weakness, and dizziness may predominate.

Physical findings are usually minimal in uncomplicated influenza. Early in the illness, the patient appears flushed, and the skin is hot and dry, although diaphoresis and mottled extremities are sometimes evident, particularly in older patients. Examination of the pharynx may yield surprisingly unremarkable results despite a severe sore throat, but injection of the mucous membranes and postnasal discharge are apparent in some cases. Mild cervical lymphadenopathy may be noted, especially in younger individuals. The results of chest examination are largely negative in uncomplicated influenza, although rhonchi, wheezes, and scattered rales have been reported with variable frequency in different outbreaks. Frank dyspnea, hyperpnea, cyanosis, diffuse rales, and signs of consolidation are indicative of pulmonary complications. Patients with apparently uncomplicated influenza have been reported to have a variety of mild ventilatory defects and increased alveolar-capillary diffusion gradients; thus, subclinical pulmonary involvement may be more common than is appreciated.

In uncomplicated influenza, the acute illness generally resolves over 2–5 days, and most patients have largely recovered in 1 week, although cough may persist 1–2 weeks longer. In a significant minority (particularly the elderly), however, symptoms of weakness or lassitude (postinfluenza asthenia) may persist for several weeks and may prove troublesome for persons who wish to resume their full level of activity promptly. The pathogenetic basis for this asthenia is unknown, although pulmonary function abnormalities may persist for several weeks after uncomplicated influenza.

COMPLICATIONS

Complications of influenza occur most frequently in patients >65 years old and in those with certain chronic disorders, including cardiac or pulmonary diseases, diabetes mellitus, hemoglobinopathies, renal dysfunction, and immunosuppression. Pregnancy in the second or third trimester predisposes to complications with influenza. Children <5 years old (especially infants) are also at high risk for complications (Table 224-2).

Pulmonary Complications • PNEUMONIA The most significant complication of influenza is pneumonia: “primary” influenza viral pneumonia, secondary bacterial pneumonia, or mixed viral and bacterial pneumonia (discussed below).

Primary influenza viral pneumonia Primary influenza viral pneumonia is the least common but most severe of the pneumonic complications. It presents as acute influenza that does not resolve but instead progresses relentlessly, with persistent fever, dyspnea, and eventual cyanosis. Sputum production is generally scanty, but the sputum can contain blood. Few physical signs may be evident early in the illness. In more advanced cases, diffuse rales may be noted, and imaging findings consistent with diffuse interstitial infiltrates and/or acute respiratory distress syndrome may be present. In such cases, arterial blood-gas determinations show marked hypoxia. Viral cultures of respiratory secretions and lung parenchyma, especially if samples are taken early in illness, yield high titers of virus. In fatal cases of primary viral pneumonia, histopathologic examination reveals a marked inflammatory reaction in the alveolar septa, with edema and infiltration by lymphocytes, macrophages, occasional plasma cells, and variable numbers of neutrophils. Fibrin thrombi in alveolar capillaries, along with necrosis and hemorrhage, have also been noted. Eosinophilic hyaline membranes can be found lining alveoli and alveolar ducts.

Primary influenza viral pneumonia has a predilection for individuals with cardiac disease, particularly those with mitral stenosis, but has also been reported in otherwise-healthy young adults as well as in older individuals with chronic pulmonary disorders. In some pandemics of influenza (notably those of 1918 and 1957), pregnancy increased the risk of primary influenza pneumonia. Subsequent epidemics of influenza have been associated with increased rates of hospitalization among pregnant women, which were also noted in the pandemic of 2009–2010.

Secondary bacterial pneumonia Secondary bacterial pneumonia follows acute influenza. Improvement of the patient’s condition over 2–3 days is followed by a reappearance of fever along with clinical signs and symptoms of bacterial pneumonia, including cough, production of purulent sputum, and physical and x-ray signs of consolidation. The most common bacterial pathogens in this setting are *Streptococcus pneumoniae*, *Staphylococcus aureus*, and *Haemophilus influenzae*—organisms that can colonize the nasopharynx and that cause infection in the wake of changes in bronchopulmonary defenses. Secondary bacterial pneumonia occurs most frequently in high-risk individuals with chronic pulmonary and cardiac disease and in elderly individuals. Patients with secondary bacterial pneumonia often respond to appropriate antibiotic therapy when it is instituted promptly.

Mixed viral and bacterial pneumonia Perhaps the most common pneumonic complications during outbreaks of influenza have mixed features of viral and bacterial pneumonia. Patients may experience a gradual progression of their acute illness or may show transient improvement followed by clinical exacerbation, with eventual manifestation of the clinical features of bacterial pneumonia. Sputum cultures may contain both influenza A virus and one of the bacterial pathogens described above. Patchy infiltrates or areas of consolidation may be detected by physical examination and chest x-ray. Patients with mixed viral and bacterial pneumonia generally have less widespread involvement of the lung than those with primary viral pneumonia, and their bacterial infections may respond to appropriate antibacterial drugs. Mixed viral and bacterial pneumonia occurs primarily in patients with chronic cardiovascular and pulmonary diseases.

OTHER PULMONARY COMPLICATIONS Other pulmonary complications associated with influenza include worsening of chronic obstructive pulmonary disease and exacerbation of chronic bronchitis and asthma. In children, influenza infection may present as croup. Sinusitis and otitis media (the latter occurring particularly often in children) may also be associated with influenza.

Extrapulmonary Complications Myositis, rhabdomyolysis, and myoglobinuria are occasional complications of influenza infection. Although myalgias are exceedingly common in influenza, true myositis is rare. Patients with acute myositis have exquisite tenderness of the affected muscles, most commonly in the legs, and may not be able to tolerate even the slightest pressure, such as the touch of bedsheets. In the most severe cases, there is frank swelling and boggy muscles. Serum levels of creatine phosphokinase and aldolase are markedly elevated, and an occasional patient develops renal failure from myoglobinuria. The pathogenesis of influenza-associated myositis is also unclear, although the presence of influenza virus in affected muscles has been reported.

Myocarditis and pericarditis were reported in association with influenza virus infection during the 1918–1919 pandemic; these reports were based largely on histopathologic findings, and these complications have been reported only infrequently since that time. Electrocardiographic changes during acute influenza are common among patients who have cardiac disease but have been ascribed most often to exacerbations of the underlying cardiac disease rather than to direct involvement of the myocardium with influenza virus. Epidemiologic data have shown an association between influenza outbreaks and increased cardiovascular-associated hospitalizations.

Central nervous system (CNS) complications such as encephalitis and transverse myelitis have been associated with influenza. Encephalitis is a rare but potentially serious complication that has been reported with influenza A and B virus infections. Children <5 years of age appear to be at greatest risk. The pathogenetic mechanisms by which influenza causes CNS disease are unclear. Guillain-Barré

syndrome has been reported following influenza infection and, uncommonly, after influenza vaccination (see “Prophylaxis,” below).

Toxic shock syndrome associated with *S. aureus* or group A streptococcal infection following acute influenza infection has been described (Chaps. 172 and 173).

Reye’s syndrome is a serious complication in children that is associated with influenza B and—to a lesser extent—influenza A virus infection as well as with varicella-zoster virus and other viral infections. An epidemiologic association between Reye’s syndrome and aspirin therapy for the antecedent viral infection has been noted; the syndrome’s incidence has decreased markedly with widespread warnings regarding aspirin use by children with acute viral respiratory infections.

In addition to complications involving the specific organ systems described above, influenza outbreaks include cases in which elderly and other high-risk individuals develop influenza and subsequently experience a gradual deterioration of underlying cardiovascular, pulmonary, or renal function—changes that occasionally are irreversible and lead to death. These deaths contribute to the overall excess mortality associated with influenza outbreaks.

LABORATORY FINDINGS AND DIAGNOSIS

During acute influenza, virus may be detected in throat swabs, nasopharyngeal swabs or washes, or sputum. Reverse-transcriptase polymerase chain reaction (RT-PCR) is the most sensitive and specific technique for detection of influenza viruses. RT-PCR can differentiate among influenza subtypes and is used for detection of avian influenza viruses. Rapid influenza diagnostic tests (RIDTs) detect influenza virus antigens by immunologic or enzymatic techniques. RIDTs yield results quickly, and some tests can distinguish between influenza A and B viruses. Although relatively specific, RIDTs vary in sensitivity with the technique and the virus to be detected.

Influenza virus may be isolated from tissue culture or chick embryos, but these labor-intensive procedures generally are no longer used for diagnostic purposes. Serologic methods for diagnosis require comparison of antibody titers in sera obtained during the acute illness with those in sera obtained 10–14 days after the onset of illness and are useful primarily in retrospect and for epidemiologic studies.

Other laboratory tests generally are not helpful in the specific diagnosis of influenza virus infection. Leukocyte counts are variable, frequently being low early in illness and normal or slightly elevated later. Severe leukopenia has been described in overwhelming viral or bacterial infection, whereas leukocytosis with $>15,000$ cells/ μL raises the suspicion of secondary bacterial infection.

DIFFERENTIAL DIAGNOSIS

During a community-wide outbreak, a clinical diagnosis of influenza can be made with a high degree of certainty in patients who present to a physician’s office with the typical febrile respiratory illness described above. In the absence of an outbreak (i.e., in sporadic or isolated cases), influenza may be difficult to differentiate on clinical grounds alone from an acute respiratory illness caused by any of a variety of respiratory viruses or by *Mycoplasma pneumoniae*. Severe streptococcal pharyngitis or early bacterial pneumonia may mimic acute influenza, although bacterial pneumonias generally do not run a self-limited course. Purulent sputum in which a bacterial pathogen can be detected by Gram’s staining is an important diagnostic feature in bacterial pneumonia.

TREATMENT INFLUENZA

(See also Chap. 215e) Specific antiviral therapy is available for influenza (Table 224-3): the neuraminidase inhibitors zanamivir and oseltamivir for both influenza A and influenza B and the adamantane agents amantadine and rimantadine for influenza A. The epidemiologic patterns of resistance to the influenza antiviral drugs are crucial elements in the selection of treatment. Up-to-date information on patterns of resistance to influenza antiviral drugs is available through www.cdc.gov/flu.

A 5-day course of oseltamivir or zanamivir reduces the duration of signs and symptoms of uncomplicated influenza by 1–1.5 days if treatment is started within 2 days of the onset of illness and may be effective if started up to 5 days after onset of symptoms. Zanamivir is administered via an oral inhalation device and may exacerbate bronchospasm in asthmatic patients. Oseltamivir has been associated with nausea and vomiting, whose frequency can be reduced by administration of the drug with food. Oseltamivir has also been associated with neuropsychiatric side effects in children. Peramivir, an investigational neuraminidase inhibitor that can be administered intravenously, is being evaluated in clinical trials, as is an intravenous form of zanamivir.

Amantadine and rimantadine are active only against influenza A, and widespread resistance exists among influenza A/H1N1 and A/H3N2 viruses that are circulating currently; thus, the use of these drugs is not recommended unless influenza isolates are known to be sensitive. Amantadine or rimantadine treatment of illness caused by sensitive strains of influenza A virus reduces the duration of symptoms of uncomplicated influenza by ~50% if begun within 48 h after onset of illness—an effect similar to that of the neuraminidase inhibitors. Of amantadine recipients, 5–10% experience mild

TABLE 224-3 ANTIVIRAL MEDICATIONS FOR TREATMENT AND PROPHYLAXIS OF INFLUENZA

Antiviral Drug	Age Group (Years)		
	Children (≤ 12)	13–64	≥ 65
Oseltamivir			
Treatment, influenza A and B	Age 1–12, dose varies by weight ^a	75 mg PO bid	75 mg PO bid
Prophylaxis, influenza A and B	Age 1–12, dose varies by weight ^a	75 mg PO qd	75 mg PO qd
Zanamivir			
Treatment, influenza A and B	Age 7–12, 10 mg bid by inhalation	10 mg bid by inhalation	10 mg bid by inhalation
Prophylaxis, influenza A and B	Age 5–12, 10 mg qd by inhalation	10 mg qd by inhalation	10 mg qd by inhalation
Amantadine^c			
Treatment, influenza A	Age 1–9, 5 mg/kg in 2 divided doses, up to 150 mg/d	Age ≥ 10 , 100 mg PO bid	≤ 100 mg/d
Prophylaxis, influenza A	Age 1–9, 5 mg/kg in 2 divided doses, up to 150 mg/d	Age ≥ 10 , 100 mg PO bid	≤ 100 mg/d
Rimantadine^c			
Treatment, influenza A	Not approved	100 mg PO bid	100–200 mg/d
Prophylaxis, influenza A	Age 1–9, 5 mg/kg in 2 divided doses, up to 150 mg/d	Age ≥ 10 , 100 mg PO bid	100–200 mg/d

^a <15 kg: 30 mg bid; >15 –23 kg: 45 mg bid; >23 –40 kg: 60 mg bid; >40 kg: 75 mg bid. For children <1 year of age, see www.cdc.gov/h1n1flu/recommendations.htm. ^b <15 kg: 30 mg qd; >15 –23 kg: 45 mg qd; >23 –40 kg: 60 mg qd; >40 kg: 75 mg qd. For children <1 year of age, see www.cdc.gov/h1n1flu/recommendations.htm. ^cAmantadine and rimantadine are not currently recommended (2013–2014) because of widespread resistance in influenza A viruses. Their use may be reconsidered if viral susceptibility is reestablished.

1214 CNS side effects, primarily jitteriness, anxiety, insomnia, or difficulty concentrating. These side effects disappear promptly upon cessation of therapy. Rimantadine appears to be equally efficacious and is associated with less frequent CNS side effects than is amantadine.

Ribavirin is a nucleoside analogue with activity against influenza A and B viruses in vitro. Its efficacy against influenza when administered as an aerosol is reportedly variable, and it is ineffective when administered orally. Its efficacy in the treatment of influenza A or B has not been established.

The therapeutic efficacy of antiviral compounds in influenza has been demonstrated primarily in studies of young adults with uncomplicated disease. The effectiveness of these drugs in the treatment or prevention of complications of influenza is unclear. Pooled analyses of observational investigations and some efficacy studies have suggested that treatment with oseltamivir may reduce the frequency of lower respiratory complications and hospitalization. Therapy for primary influenza pneumonia is directed at maintaining oxygenation and is most appropriately undertaken in an intensive care unit, with aggressive respiratory and hemodynamic support as needed.

Antibacterial drugs should be reserved for the treatment of bacterial complications of acute influenza, such as secondary bacterial pneumonia. The choice of antibiotics should be guided by Gram's staining and culture of appropriate specimens of respiratory secretions, such as sputum. If the etiology of a case of bacterial pneumonia is unclear from an examination of respiratory secretions, empirical antibiotics effective against the most common bacterial pathogens in this setting (*S. pneumoniae*, *S. aureus*, and *H. influenzae*) should be selected (**Chaps. 171, 172, and 182**).

For uncomplicated influenza in individuals at low risk of complications, symptom-based rather than antiviral therapy may be considered. Acetaminophen or nonsteroidal anti-inflammatory agents can be used for relief of headache, myalgia, and fever, but salicylates should be avoided in children <18 years of age because of the possible association with Reye's syndrome (see "Extrapulmonary Complications," above). Because cough is ordinarily self-limited, treatment with cough suppressants generally is not indicated; codeine-containing compounds may be used if the cough is particularly troublesome. Patients should be advised to rest and maintain hydration during acute illness and to return to full activity only gradually after illness has resolved, especially if it has been severe.

PROPHYLAXIS

The major public health measure for prevention of influenza is vaccination. Both inactivated (killed) and live attenuated vaccines are available and are generated from isolates of influenza A and B viruses that circulated in the previous influenza seasons and are anticipated to circulate in the upcoming season. For inactivated vaccines, 50–80% protection against influenza is expected if the vaccine virus and the currently circulating viruses are closely related. Available inactivated vaccines have been highly purified and are associated with few reactions. Up to 5% of individuals experience low-grade fever and mild systemic symptoms 8–24 h after vaccination, and up to one-third develop mild redness or tenderness at the vaccination site. Although the 1976 swine influenza vaccine appears to have been associated with an increased frequency of Guillain-Barré syndrome, influenza vaccines administered since 1976 generally have not been. Possible exceptions were noted during the 1992–1993 and 1993–1994 influenza seasons, when there may have been an excess risk of this syndrome (slightly more than 1 case per 1 million vaccine recipients). Large-scale studies of vaccination with the 2009 pandemic H1N1 vaccine also suggested a possible increased risk of Guillain-Barré syndrome (1 case per 1 million vaccinees). However, the overall health risk following influenza substantially outweighs the potential risk associated with vaccination.

A live attenuated influenza vaccine administered by intranasal spray is available. The vaccine is generated by reassortment between currently circulating strains of influenza A and B viruses and a cold-adapted, attenuated master strain. The cold-adapted vaccine is well

tolerated and highly efficacious (>90% protective) in young children; in one study, it provided protection against a circulating influenza virus that had drifted antigenically away from the vaccine strain. Live attenuated vaccine is approved for use in healthy nonpregnant persons 2–49 years of age.

Since 1975, influenza vaccines have been trivalent—i.e., they have contained two influenza A subtypes (H3N2 and H1N1) and one influenza B component. However, two antigenically distinct lineages of influenza B virus have circulated since the 1980s, and a quadrivalent vaccine that includes both B lineages is now available (2013–2014) as well. Quadrivalent vaccines are available in both inactivated and live-attenuated vaccine formulations.

Inactivated influenza vaccines have been noted to be less immunogenic in the elderly. A higher-dose trivalent vaccine containing 60 µg of each antigen and a lower-dose, intradermally administered trivalent vaccine containing 9 µg of each antigen have been approved for use in individuals ≥65 years of age and individuals 18–64 years of age, respectively.

The influenza vaccines discussed above are manufactured in eggs and should not be administered to persons with true hypersensitivity to eggs. For use in this situation, an egg-free vaccine manufactured in cells through recombinant DNA techniques (Flublok®; Protein Sciences Corporation, Meriden, CT) has been approved. Active research is under way to develop vaccines with broad activity against antigenically distinct subtypes ("universal influenza vaccines").

Historically, the U.S. Public Health Service has recommended influenza vaccination for certain groups at high risk for complications of influenza on the basis of age or underlying disease (Table 224-2) or for their close contacts. Although such individuals will continue to be the focus of vaccination programs, the recommendations have been progressively expanded, and immunization of the entire population above the age of 6 months has been recommended since 2010–2011. (Approved influenza vaccines are not available for infants <6 months of age.) This expanded recommendation reflects increased recognition of previously unappreciated risk factors (e.g., obesity, postpartum conditions, and racial or ethnic influences) as well as an appreciation that more widespread use of vaccine is required for influenza control. Inactivated vaccines may be administered safely to immunocompromised patients. Influenza vaccination is not associated with exacerbations of chronic nervous system diseases such as multiple sclerosis. Vaccine should be administered early in the autumn before influenza outbreaks occur and should then be given annually to maintain immunity against the most current influenza virus strains.

Although antiviral drugs provide chemoprophylaxis against influenza, their use for that purpose has been limited because of concern about current and future development of resistance. Chemoprophylaxis with oseltamivir or zanamivir has been 84–89% efficacious against influenza A and B (Table 224-3). Chemoprophylaxis with amantadine or rimantadine is no longer recommended because of widespread resistance to these drugs. In earlier studies with sensitive viruses, prophylaxis with amantadine or rimantadine was 70–100% effective against illness associated with influenza A virus.

Chemoprophylaxis for healthy persons after community exposure generally is not recommended but may be considered for individuals at high risk of complications who have had close contact with an acutely ill person with influenza. During an outbreak, antiviral chemoprophylaxis can be administered simultaneously with inactivated vaccine because the drugs do not interfere with an immune response to the vaccine. However, concurrent administration of chemoprophylaxis and live attenuated vaccine may interfere with the immune response to the latter. Antiviral drugs should not be administered until at least 2 weeks after administration of live vaccine, and administration of live vaccine should not begin until at least 48 h after antiviral drug administration has been stopped. Chemoprophylaxis may also be considered to control nosocomial outbreaks of influenza. For that purpose, prophylaxis should be instituted promptly when influenza activity is detected and must be continued daily for the duration of the outbreak.

SECTION 14

INFECTIONS DUE TO HUMAN IMMUNODEFICIENCY VIRUS AND OTHER HUMAN RETROVIRUSES

225e The Human Retroviruses

Dan L. Longo, Anthony S. Fauci

The retroviruses, which make up a large family (Retroviridae), infect mainly vertebrates. These viruses have a unique replication cycle whereby their genetic information is encoded by RNA rather than DNA. Retroviruses contain an RNA-dependent DNA polymerase (a reverse transcriptase) that directs the synthesis of a DNA form of the viral genome after infection of a host cell. The designation *retrovirus* denotes that information in the form of RNA is transcribed into DNA in the host cell—a sequence that overturned a central dogma of molecular biology: that information passes unidirectionally from DNA to RNA to protein. The observation that RNA was the source of genetic information in the causative agents of certain animal tumors led to a number of paradigm-shifting biologic insights regarding not only the direction of genetic information passage but also the viral etiology of certain cancers and the concept of oncogenes as normal host genes scavenged and altered by a viral vector.

The family Retroviridae includes seven subfamilies (Table 225e-1). Members of two of the families infect humans with pathologic consequences: the deltaretroviruses, of which human T cell lymphotropic virus (HTLV) type 1 is the most important in humans; and lentiviruses, of which HIV is the most important in humans.

The wide variety of interactions of a retrovirus with its host range from completely benign events (e.g., silent carriage of endogenous retroviral sequences in the germline genome of many animal species) to rapidly fatal infections (e.g., exogenous infection with an oncogenic virus such as Rous sarcoma virus in chickens). The ability of retroviruses to acquire and alter the structure and function of host cell sequences has revolutionized our understanding of molecular carcinogenesis. The viruses can insert into the germline genome of the host cell and behave as a transposable or movable genetic element. They can activate or inactivate genes near the site of integration into the genome. They can rapidly alter their own genome by recombination and mutation under selective environmental stimuli.

Most human viral diseases occur as a consequence of tissue destruction either directly by the virus itself or indirectly by the host's response to the virus. Although these mechanisms are operative in retroviral infections, retroviruses have additional mechanisms of inducing disease, including the malignant transformation of an infected cell and the induction of an immunodeficiency state that renders the host susceptible to opportunistic diseases (infections and neoplasms; Chap. 226).

TABLE 225e-1 CLASSIFICATION OF RETROVIRUSES: THE FAMILY RETROVIRIDAE

Genus	Example(s)	Feature
<i>Alpharetrovirus</i>	Rous sarcoma virus	Contains <i>src</i> oncogene
<i>Betaretrovirus</i>	Mouse mammary tumor virus	Exogenous or endogenous
<i>Gammaretrovirus</i>	Abelson murine leukemia virus	Contains <i>abl</i> oncogene
<i>Deltaretrovirus</i>	HTLV-1	Causes T cell lymphoma and neurologic disease
<i>Epsilonretrovirus</i>	Walleye dermal sarcoma virus	Not known to be pathogenic in humans
<i>Lentivirus</i>	HIV-1, -2	Causes AIDS
<i>Spumavirus</i>	Simian foamy virus	Not known to be pathogenic in humans

STRUCTURE AND LIFE CYCLE

All retroviruses are similar in structure, genome organization, and mode of replication. Retroviruses are 70–130 nm in diameter and have a lipid-containing envelope surrounding an icosahedral capsid with a dense inner core. The core contains two identical copies of the single-strand RNA genome. The RNA molecules are 8–10 kb long and are complexed with reverse transcriptase and tRNA. Other viral proteins, such as integrase, are also components of the virion particle. The RNA has features usually found in mRNA: a cap site at the 5' end of the molecule, which is important in the initiation of mRNA translation, and a polyadenylation site at the 3' end, which influences mRNA turnover (i.e., messages with shorter polyA tails turn over faster than messages with longer polyA tails). However, the retroviral RNA is not translated; instead it is transcribed into DNA. The DNA form of the retroviral genome is called a *provirus*.

The replication cycle of retroviruses proceeds in two phases (Fig. 225e-1). In the first phase, the virus enters the cytoplasm after binding to one or more specific cell-surface receptors; the viral RNA and reverse transcriptase synthesize a double-strand DNA version of the RNA template; and the provirus moves into the nucleus and integrates into the host cell genome. This proviral integration is permanent. Although some animal retroviruses integrate into a single specific site of the host genome in every infected cell, the human retroviruses integrate randomly. This first phase of replication depends entirely on gene products in the virus. The second phase includes the synthesis and processing of viral genomes, mRNAs, and proteins using host cell machinery, often under the influence of viral gene products. Virions are assembled and released from the cell by budding from the membrane; host cell membrane proteins are frequently incorporated into the envelope of the virus. Proviral integration occurs during the S-phase of the cell cycle; thus, in general, nondividing cells are resistant to retroviral infection. Only the lentiviruses are able to infect nondividing cells. Once a host cell is infected, it is infected for the life of the cell.



Retroviral genomes include both coding and noncoding sequences (Fig. 225e-2). In general, noncoding sequences are important recognition signals for DNA or RNA synthesis or processing events and are located in the 5' and 3' terminal regions of the genome. All retroviral genomes are terminally redundant, containing identical sequences called *long terminal repeats* (LTRs). The ends of the retroviral RNA genome differ slightly in sequence from the integrated retroviral DNA. In the latter, the LTR sequences are repeated in both the 5' and the 3' terminus of the virus. The LTRs contain sequences involved in initiating the expression of the viral proteins, the integration of the provirus, and the polyadenylation of viral RNAs. The primer binding site, which is critical for the initiation of reverse transcription, and the viral packaging sequences are located outside the LTR sequences. The coding regions include the *gag* (group-specific antigen, core protein), *pol* (RNA-dependent DNA polymerase), and *env* (envelope) genes. The *gag* gene encodes a precursor polyprotein that is cleaved to form three to five capsid proteins; a fraction of the Gag precursor proteins also contain a protease responsible for cleaving the Gag and Pol polyproteins. A Gag-Pol polyprotein gives rise to the protease that is responsible for cleaving the Gag-Pol polyprotein. The *pol* gene encodes three proteins: the reverse transcriptase, the integrase, and the protease. The reverse transcriptase copies the viral RNA into the double-strand DNA provirus, which inserts itself into the host cell DNA via the action of integrase. The protease cleaves the Gag-Pol polyprotein into smaller protein products. The *env* gene encodes the envelope glycoproteins: one protein that binds to specific surface receptors and determines what cell types can be infected and a smaller transmembrane protein that anchors the complex to the envelope. Fig. 225e-3 shows how the retroviral gene products make up the virus structure.

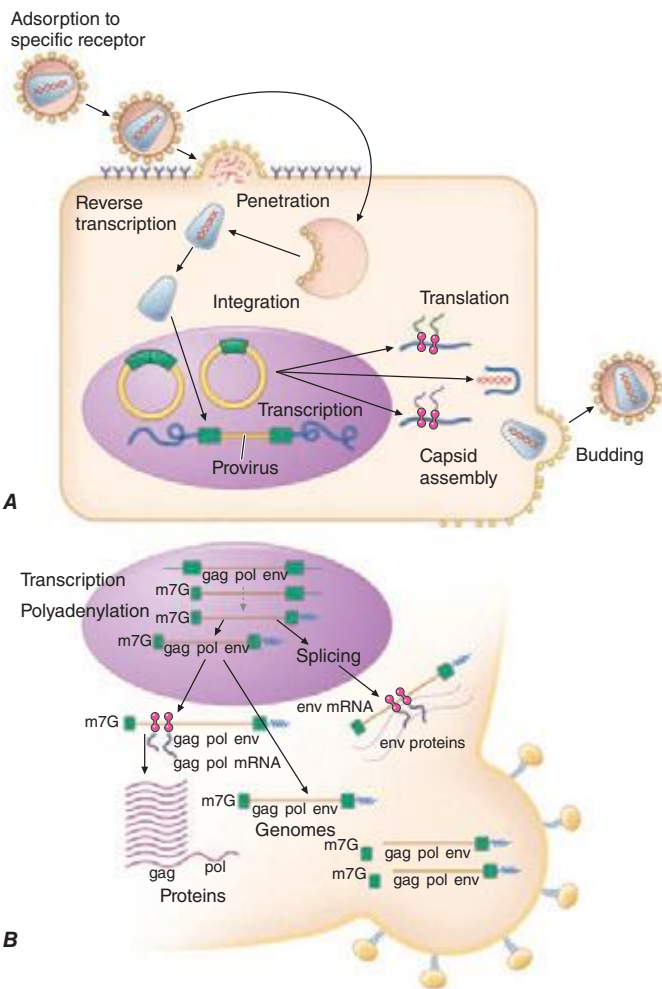


FIGURE 225e-1 The life cycle of retroviruses. **A.** Overview of virus replication. The retrovirus enters a target cell by binding to a specific cell-surface receptor; once the virus is internalized, its RNA is released from the nucleocapsid and is reverse-transcribed into proviral DNA. The provirus is inserted into the genome and then transcribed into RNA; the RNA is translated; and virions assemble and are extruded from the cell membrane by budding. **B.** Overview of retroviral gene expression. The provirus is transcribed, capped, and polyadenylated. Viral RNA molecules then have one of three fates: they are exported to the cytoplasm, where they are packaged as the viral RNA in infectious viral particles; they are spliced to form the message for the envelope polyprotein; or they are translated into Gag and Pol proteins. Most of the messages for the Pol protein fail to initiate Pol translation because of a stop codon before its initiation; however, in a fraction of the messages, the stop codon is missed and the Pol proteins are translated. (Modified from JM Coffin, in BN Fields, DM Knipe [eds]: *Fields Virology*. New York, Raven, 1990; with permission.)

HTLVs have a region between *env* and the 3' LTR that encodes several proteins and transcripts in overlapping reading frames (Fig. 225e-2). Tax is a 40-kDa protein that does not bind to DNA but induces the expression of host cell transcription factors that alter host cell gene expression and is capable of inducing cell transformation under certain circumstances. Rex is a 27-kDa protein that regulates the expression of viral mRNAs. Other transcripts from this region (p12, p13, p30) tend to restrict expression of viral genes and diminish the immunogenicity of infected cells. The protein of *HBZ*, a product of the complementary proviral DNA strand, interacts with many cellular transcription factors and signaling proteins. It stimulates proliferation of infected cells and is the only viral product universally expressed in HTLV-1-infected tumor cells. These proteins are produced from messages that are similar but that are spliced differently from overlapping but distinct exons.

The lentiviruses in general, and HIV-1 and -2 in particular, contain a larger genome than other pathogenic retroviruses. They contain an untranslated region between *pol* and *env* that encodes portions of several proteins, varying with the reading frame into which the mRNA is spliced. Tat is a 14-kDa protein that augments the expression of virus from the LTR. The Rev protein of HIV-1, similar to the Rex protein of HTLV, regulates RNA splicing and/or RNA transport. The Nef protein downregulates CD4, the cellular receptor for HIV; alters host T cell-activation pathways; and enhances viral infectivity. The Vif protein is necessary for the proper assembly of the HIV nucleoprotein core in many types of cells; without Vif, proviral DNA is not efficiently produced in these infected cells. In addition, the Vif protein targets APOBEC (apolipoprotein B mRNA-editing enzyme catalytic polypeptide, a cytidine deaminase that mutates the viral sequence) for proteasomal degradation, thus blocking its virus-suppressing effect. Vpr, Vpu (HIV-1 only), and Vpx (HIV-2 only) are viral proteins encoded by translation of the same message in different reading frames. As noted above, oncogenic retroviruses depend on cell proliferation for their replication; lentiviruses can infect nondividing cells, largely through effects mediated by Vpr. Vpr facilitates transport of the provirus into the nucleus and can induce other cellular changes, such as G₂ growth arrest and differentiation of some target cells. Vpx is structurally related to Vpr, but its functions are not fully defined. Vpu promotes the degradation of CD4 in the endoplasmic reticulum and stimulates the release of virions from infected cells.

Retroviruses can be either exogenously acquired (by infection with an infected cell or a free virion capable of replication) or transmitted in the germline as endogenous virus. Endogenous retroviruses are often replication defective. The human genome contains endogenous retroviral sequences, but there are no known replication-competent endogenous retroviruses in humans.

In general, viruses that contain only the *gag*, *pol*, and *env* genes either are not pathogenic or take a long time to induce disease; these observations indicate the importance of the other regulatory genes in viral disease pathogenesis. The pathogenesis of neoplastic transformation by retroviruses relies on the chance integration of the provirus at a spot in the genome resulting in the expression of a cellular gene (proto-oncogene) that becomes transforming by virtue of its unregulated expression. For example, avian leukosis virus causes B cell leukemia by inducing the expression of *myc*. Some retroviruses possess captured and altered cellular genes near their integration site, and these viral oncogenes can transform the infected host cell. Viruses that have oncogenes often have lost a portion of their genome that is required for replication. Such viruses need helper viruses to reproduce, a feature that may explain why these acute transforming retroviruses are rare in nature. All human retroviruses identified to date are exogenous and are not acutely transforming (i.e., they lack a transforming oncogene).

These remarkable properties of retroviruses have led to experimental efforts to use them as vectors to insert specific genes into particular cell types, a process known as *gene therapy* or *gene transfer*. The process could be used to repair a genetic defect or to introduce a new property that could be used therapeutically; for example, a gene (e.g., thymidine kinase) that would make a tumor cell susceptible to killing by a drug (e.g., ganciclovir) could be inserted. One source of concern about the use of retroviral vectors in humans is that replication-competent viruses might rescue endogenous retroviral replication, with unpredictable results. This concern is not merely hypothetical: the detection of proteins encoded by endogenous retroviral sequences on the surface of cancer cells implies that the genetic events leading to the cancer were able to activate the synthesis of these usually silent genes.

HUMAN T CELL LYMPHOTROPIC VIRUS

HTLV-1 was isolated in 1980 from a T cell lymphoma cell line from a patient originally thought to have cutaneous T cell lymphoma. Later it became clear that the patient had a distinct form of lymphoma (originally reported in Japan) called *adult T cell leukemia/lymphoma* (ATL). Serologic data have determined that HTLV-1 is the cause of at least two important diseases: ATL and tropical spastic paraparesis, also

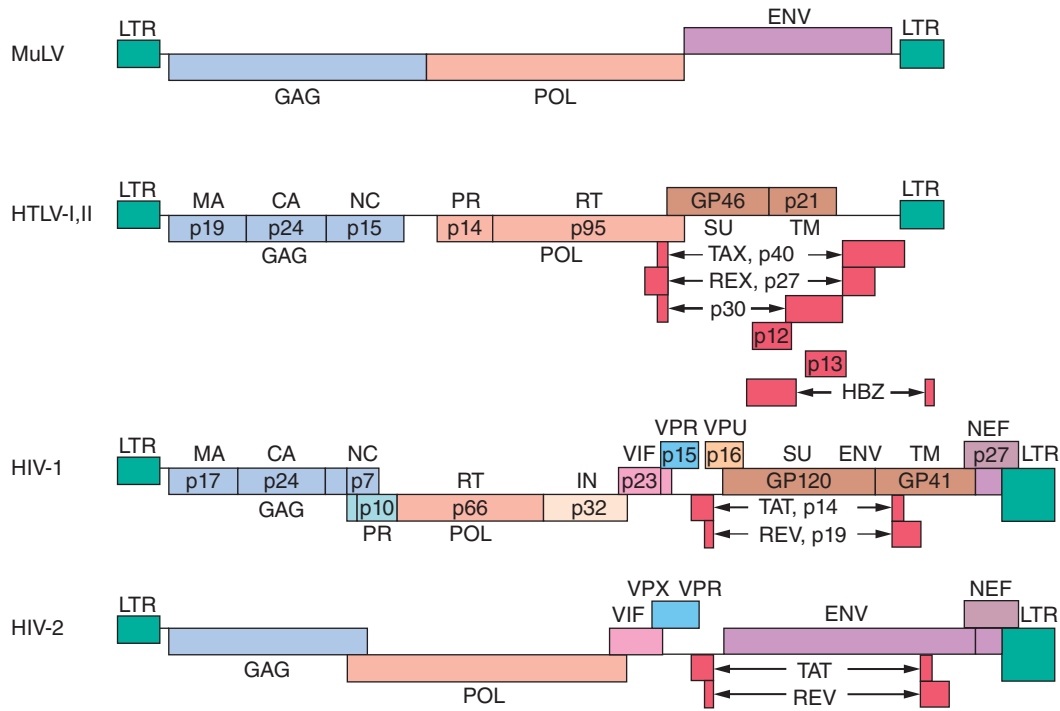


FIGURE 225e-2 Genomic structure of retroviruses. The murine leukemia virus MuLV has the typical three structural genes: *gag*, *pol*, and *env*. The *gag* region gives rise to three proteins: matrix (MA), capsid (CA), and nucleic acid-binding (NC) proteins. The *pol* region encodes both a protease (PR) responsible for cleaving the viral polyproteins and a reverse transcriptase (RT). In addition, HIV *pol* encodes an integrase (IN). The *env* region encodes a surface protein (SU) and a small transmembrane protein (TM). The human retroviruses have additional gene products translated in each of the three possible reading frames. HTLV-1 and HTLV-2 have *tax* and *rex* genes with exons on either side of the *env* gene. HIV-1 and HIV-2 have six accessory gene products: *tat*, *rev*, *vif*, *nef*, *vpr*, and either *vpu* (in HIV-1) or *vpx* (in HIV-2). The genes for these proteins are located mainly between the *pol* and *env* genes. GP, glycoprotein; HBZ, HTLV-1 basic leucine zipper domain-containing protein; LTR, long terminal repeat.

called *HTLV-1-associated myelopathy* (HAM). HTLV-1 may also play a role in infective dermatitis, arthritis, uveitis, and Sjögren’s syndrome.

Two years after the isolation of HTLV-1, HTLV-2 was isolated from a patient with an unusual form of hairy cell leukemia that affected

T cells. Epidemiologic studies of HTLV-2 failed to reveal a consistent disease association. Similarly, HTLV-3 and HTLV-4 have been identified but have no known disease association.

BIOLOGY AND MOLECULAR BIOLOGY

Because the biology of HTLV-1 and that of HTLV-2 are similar, the following discussion will focus on HTLV-1.

Human glucose transporter protein 1 (GLUT-1) functions as a receptor for HTLV-1, probably acting together with neuropilin-1 (NRP1) and heparan sulfate proteoglycans. Generally, only T cells are productively infected, but infection of B cells and other cell types is occasionally detected. The most common outcome of HTLV-1 infection is latent carriage of randomly integrated provirus in CD4+ T cells. HTLV-1 does not contain an oncogene and does not insert into a unique site in the genome. Indeed, most infected cells express no viral gene products. The only viral gene product that is routinely expressed in tumor cells transformed by HTLV-1 *in vivo* is *hbz*. The *tax* gene is thought to be critical to the transformation process but is not expressed in the tumor cells of many ATL patients, possibly because of the immunogenicity of *tax*-expressing cells. Cells transformed *in vitro*, by contrast, actively transcribe HTLV-1 RNA and produce infectious virions. Most HTLV-1-transformed cell lines are the result of the infection of a normal host T cell *in vitro*. It is difficult to establish cell lines derived from authentic ATL cells.

Although *tax* does not itself bind to DNA, it does induce the expression of a wide range of host cell gene products, including transcription factors (especially *c-rel/NF-κB*, *ets-1* and *-2*, and members of the *fos/jun* family), cytokines (e.g., interleukin [IL] 2, granulocyte-macrophage colony-stimulating factor, and tumor necrosis factor), membrane proteins and receptors (major histocompatibility [MHC] molecules and IL-2 receptor α), and chromatin remodeling complexes. The genes activated by *tax* are generally controlled by transcription factors of the *c-rel/NF-κB* and cyclic AMP response element binding (CREB) protein families. It is unclear how this induction of host

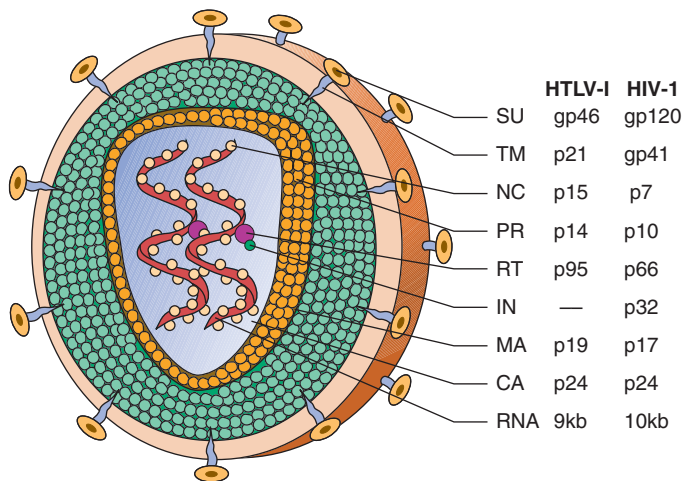


FIGURE 225e-3 Schematic structure of human retroviruses. The surface glycoprotein (SU) is responsible for binding to receptors of host cells. The transmembrane protein (TM) anchors SU to the virus. NC is a nucleic acid-binding protein found in association with the viral RNA. A protease (PR) cleaves the polyproteins encoded by the *gag*, *pol*, and *env* genes into their functional components. RT is reverse transcriptase, and IN is an integrase present in some retroviruses (e.g., HIV-1) that facilitates insertion of the provirus into the host genome. The matrix protein (MA) is a Gag protein closely associated with the lipid of the envelope. The capsid protein (CA) forms the major internal structure of the virus, the core shell.

gene expression leads to neoplastic transformation; *tax* can interfere with G_1 and mitotic cell-cycle checkpoints, block apoptosis, inhibit DNA repair, and promote antigen-independent T cell proliferation. Induction of a cytokine–autocrine loop has been proposed; however, IL-2 is not the crucial cytokine. The involvement of IL-4, IL-7, and IL-15 has been proposed.

In light of the irregular expression of *tax* in ATL cells, it has been suggested that *tax* is important in the early phases of transformation but is not essential for the maintenance of the transformed state. The maintenance role is thought to be due to *hbz* expression. As is clear from the epidemiology of HTLV-1 infection, transformation of an infected cell is a rare event and may depend on heterogeneous second, third, or fourth genetic hits. No consistent chromosomal abnormalities have been described in ATL; however, aneuploidy is common and individual cases with p53 mutations and translocations involving the T cell receptor genes on chromosome 14 have been reported. *Tax* may repress certain DNA repair enzymes, permitting the accumulation of genetic damage that would normally be repaired. However, the molecular pathogenesis of HTLV-1-induced neoplasia is not fully understood.

FEATURES OF HTLV-1 INFECTION

Epidemiology HTLV-1 infection is transmitted in at least three ways: from mother to child, especially via breast milk; through sexual activity, more commonly from men to women; and through the blood—via contaminated transfusions or contaminated needles. The virus is most commonly transmitted perinatally. Compared with HIV, which can be transmitted in cell-free form, HTLV-1 is less infectious, and its transmission usually requires cell-to-cell contact.



HTLV-1 is endemic in southwestern Japan and Okinawa, where >1 million persons are infected. Antibodies to HTLV-1 are present in the serum of up to 35% of Okinawans, 10% of residents of the Japanese island of Kyushu, and <1% of persons in nonendemic regions of Japan. Despite this high prevalence of infection, only ~500 cases of ATL are diagnosed in this area each year. Clusters of infection have been noted in other areas of the Orient, such as Taiwan; in the Caribbean basin, including northeastern South America; in northwestern South America; in central and southern Africa; in Italy, Israel, Iran, and Papua New Guinea; in the Arctic; and in the southeastern part of the United States (Fig. 225e-4). An estimated 5–10 million persons have HTLV-1 infection worldwide.

Progressive spastic or ataxic myelopathy developing in an individual who is HTLV-1 positive (i.e., who has serum antibodies to HTLV-1) may be due to direct infection of the nervous system with the virus, but destruction of the pyramidal tracts appears to involve HTLV-1-infected CD4+ T cells; a similar disorder may result from infection with HIV or HTLV-2. In rare instances, patients with HAM are seronegative but have detectable antibody to HTLV-1 in cerebrospinal fluid (CSF).

The cumulative lifetime risk of developing ATL is 3% among HTLV-1-infected patients, with a threefold greater risk among men than among women; a similar cumulative risk is projected for HAM (4%), but with women more commonly affected than men. The distribution of these two diseases overlaps the distribution of HTLV-1, with >95% of affected patients showing serologic evidence of HTLV-1 infection. The latency period between infection and the emergence of disease is 20–30 years for ATL. For HAM, the median latency period is ~3.3 years (range, 4 months to 30 years). The development of ATL is rare among persons infected by blood products; however, ~20% of patients with HAM acquire HTLV-1 from contaminated blood. ATL is more common among perinatally infected individuals, whereas HAM is more common among persons infected via sexual transmission.

Associated Diseases • ATL Four clinical types of HTLV-1-induced neoplasia have been described: acute, lymphomatous, chronic, and smoldering. All of these tumors are monoclonal proliferations of CD4+ postthymic T cells with clonal proviral integrations and clonal T cell receptor gene rearrangements.

ACUTE ATL About 60% of patients who develop malignancy have classic acute ATL, which is characterized by a short clinical prodrome (~2 weeks between the first symptoms and the diagnosis) and an aggressive natural history (median survival period, 6 months). The clinical picture is dominated by rapidly progressive skin lesions, pulmonary involvement, hypercalcemia, and lymphocytosis with cells containing lobulated or “flower-shaped” nuclei (see Fig. 134-10). The malignant cells have monoclonal proviral integrations and express CD4, CD3, and CD25 (low-affinity IL-2 receptors) on their surface. Serum levels of CD25 can be used as a tumor marker. Anemia and thrombocytopenia are rare. The skin lesions may be difficult to distinguish from those in mycosis fungoides. Lytic bone lesions, which are common, do not contain tumor cells but rather are composed of osteolytic cells, usually without osteoblastic activity. Despite the leukemic picture, bone marrow involvement is patchy in most cases.

The hypercalcemia of ATL is multifactorial; the tumor cells produce osteoclast-activating factors (tumor necrosis factor α , IL-1, lymphotoxin) and can also produce a parathyroid hormone–like molecule. Affected patients have an underlying immunodeficiency that makes them susceptible to opportunistic infections similar to those seen in patients with AIDS (Chap. 226). The pathogenesis of the immunodeficiency is unclear. Pulmonary infiltrates in ATL patients reflect leukemic infiltration half the time and opportunistic infections with organisms such as *Pneumocystis* and other fungi the other half. Gastrointestinal symptoms are nearly always related to opportunistic infection. *Strongyloides stercoralis* is a gastrointestinal parasite that has a pattern of endemic distribution similar to that of HTLV-1. HTLV-1-infected persons also infected with this parasite may develop ATL more often or more rapidly than those without *Strongyloides* infections. Serum concentrations of lactate dehydrogenase and alkaline

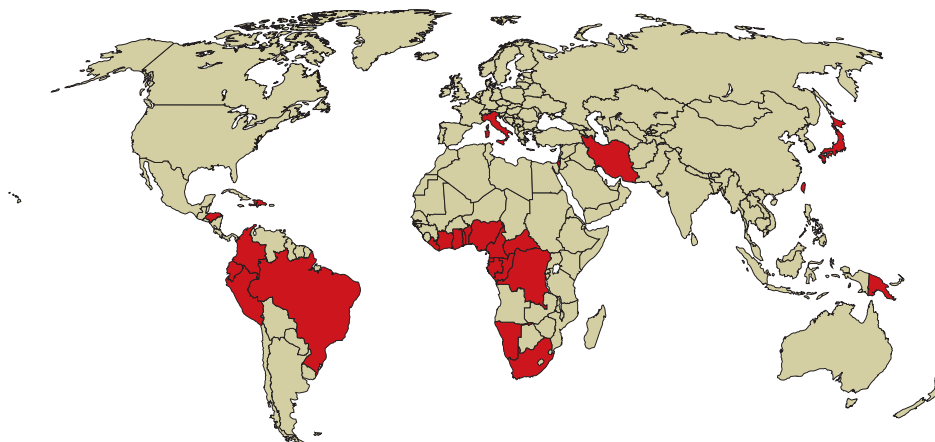


FIGURE 225e-4 Global distribution of HTLV-1 infection. Countries with a prevalence of HTLV-1 infection of 1–5% are shaded darkly. Note that the distribution of infected patients is not uniform in endemic countries. For example, the people of southwestern Japan and northeastern Brazil are more commonly affected than those in other regions of those countries.

phosphatase are often elevated in ATL. About 10% of patients have leptomeningeal involvement leading to weakness, altered mental status, paresthesia, and/or headache. Unlike other forms of central nervous system (CNS) lymphoma, ATL may be accompanied by normal CSF protein levels. The diagnosis depends on finding ATL cells in the CSF (Chap. 134).

LYMPHOMATOUS ATL The lymphomatous type of ATL occurs in ~20% of patients and is similar to the acute form in its natural history and clinical course, except that circulating abnormal cells are rare and lymphadenopathy is evident. The histology of the lymphoma is variable but does not influence the natural history. In general, the diagnosis is suspected on the basis of the patient's birthplace (see "Epidemiology," above) and the presence of skin lesions and hypercalcemia. The diagnosis is confirmed by the detection of antibodies to HTLV-1 in serum.

CHRONIC ATL Patients with the chronic form of ATL generally have normal serum levels of calcium and lactate dehydrogenase and no involvement of the CNS, bone, or gastrointestinal tract. The median duration of survival for these patients is 2 years. In some cases, chronic ATL progresses to the acute form of the disease.

SMOLDERING ATL Fewer than 5% of patients have the smoldering form of ATL. In this form, the malignant cells have monoclonal proviral integration; <5% of peripheral blood cells exhibit typical morphologic abnormalities; hypercalcemia, adenopathy, and hepatosplenomegaly do not develop; the CNS, the bones, and the gastrointestinal tract are not involved; and skin lesions and pulmonary lesions may be present. The median survival period for this small subset of patients appears to be ≥ 5 years.

HAM (TROPICAL SPASTIC PARAPARESIS) In contrast to ATL, in which there is a slight predominance of male patients, HAM affects female patients disproportionately. HAM resembles multiple sclerosis in certain ways (Chap. 458). The onset is insidious. Symptoms include weakness or stiffness in one or both legs, back pain, and urinary incontinence. Sensory changes are usually mild, but peripheral neuropathy may develop. The disease generally takes the form of slowly progressive and unremitting thoracic myelopathy; one-third of patients are bedridden within 10 years of diagnosis, and one-half are unable to walk unassisted by this point. Patients display spastic paraparesis or paraplegia with hyperreflexia, ankle clonus, and extensor plantar responses. Cognitive function is usually spared; cranial nerve abnormalities are unusual.

Magnetic resonance imaging (MRI) reveals lesions in both the white matter and the paraventricular regions of the brain as well as in the spinal cord. Pathologic examination of the spinal cord shows symmetric degeneration of the lateral columns, including the corticospinal tracts; some cases involve the posterior columns as well. The spinal meninges and cord parenchyma contain an inflammatory infiltrate with myelin destruction.

HTLV-1 is not usually found in cells of the CNS but may be detected in a small population of lymphocytes present in the CSF. In general, HTLV-1 replication is greater in HAM than in ATL, and patients with HAM have a stronger immune response to the virus. Antibodies to HTLV-1 are present in the serum and appear to be produced in the CSF of HAM patients, where titers are often higher than in the serum. The pathophysiology of HAM may involve the induction of autoimmune destruction of neural cells by T cells with specificity for viral components such as Tax or Env proteins. One theory is that susceptibility to HAM may be related to the presence of human leukocyte antigen (HLA) alleles capable of presenting viral antigens in a fashion that leads to autoimmunity. Insufficient data are available to confirm an HLA association. However, antibodies in the sera of HAM patients have been shown to bind a neuron-specific antigen (heteronuclear ribonuclear protein A1 [hnRNP A1]) and to interfere with neurotransmission *in vitro*.

It is unclear what factors influence whether HTLV-1 infection will cause disease and, if it does, whether it will induce a neoplasm (ATL) or an autoimmune disorder (HAM). Differences in viral strains, the susceptibility of particular MHC haplotypes, the route of HTLV-1

infection, the viral load, and the nature of the HTLV-1-related immune response are putative factors, but few definitive data are available.

OTHER PUTATIVE HTLV-1-RELATED DISEASES Even in the absence of the full clinical picture of HAM, bladder dysfunction is common in HTLV-1-infected women. In areas where HTLV-1 is endemic, diverse inflammatory and autoimmune diseases have been attributed to the virus, including uveitis, dermatitis, pneumonitis, rheumatoid arthritis, and polymyositis. However, a causal relationship between HTLV-1 and these illnesses has not been established.

Prevention Women in endemic areas should not breast-feed their children, and blood donors should be screened for serum antibodies to HTLV-1. As in the prevention of HIV infection, the practice of safe sex and the avoidance of needle sharing are important.

TREATMENT HTLV-1 INFECTION

For the small number of patients who develop HTLV-1-related disease, therapies are not curative. In patients with the acute and lymphomatous types of ATL, the disease progresses rapidly. Hypercalcemia is generally controlled by glucocorticoid administration and cytotoxic therapy directed against the neoplasm. The tumor is highly responsive to combination chemotherapy that is used against other forms of lymphoma; however, patients are susceptible to overwhelming bacterial and opportunistic infections, and ATL relapses within 4–10 months after remission in most cases. The combination of interferon α and zidovudine may extend survival. Because viral replication is not clearly associated with ATL progression, zidovudine is probably effective through its cytotoxic effects (as a chain-terminating thymidine analogue) rather than its antiviral effects. LSG15, a multidrug chemotherapy program developed in Japan, induces complete responses in about one-third of patients, about half of whom survive more than 2 years; however, but the median survival time is about 13 months. A pilot trial suggested that mogamulizumab, an antibody to CCR4 (a receptor for a number of chemokines, including RANTES and TARC), improved response rates when added to chemotherapy. An experimental approach using an yttrium 90–labeled or toxin-conjugated antibody to the IL-2 receptor appears promising but is not widely available. Patients with the chronic or smoldering form of ATL may be managed with an expectant approach: treat any infections, and watch and wait for signs of progression to acute disease.

Patients with HAM may obtain some benefit from the use of glucocorticoids to reduce inflammation. Antiretroviral regimens have not been effective. In one study, danazol (200 mg three times daily) produced significant neurologic improvement in five of six treated patients, with resolution of urinary incontinence in two cases, decreased spasticity in three, and restoration of the ability to walk after confinement to a wheelchair in two. Antibody to IL-15 receptor β chain has been tested with some promising clinical effects in small numbers of patients. Physical therapy and rehabilitation are important components of management.

FEATURES OF HTLV-2 INFECTION

Epidemiology HTLV-2 is endemic in certain Native American tribes and in Africa. It is generally considered to be a New World virus that was brought from Asia to the Americas 10,000–40,000 years ago during the migration of infected populations across the Bering land bridge. The mode of transmission of HTLV-2 is probably the same as that of HTLV-1 (see above). HTLV-2 may be less readily transmitted sexually than HTLV-1.

Studies of large cohorts of injection drug users with serologic assays that reliably distinguish HTLV-1 from HTLV-2 indicated that the vast majority of HTLV-positive cohort members were infected with HTLV-2. The seroprevalence of HTLV in a cohort of 7841 injection drug users from drug treatment centers in Baltimore, Chicago, Los Angeles, New Jersey (Asbury Park and Trenton), New York City (Brooklyn and Harlem), Philadelphia, and San Antonio was 20.9%,

225e-6 with >97% of cases due to HTLV-2. The seroprevalence of HTLV-2 was higher in the Southwest and the Midwest than in the Northeast. In contrast, the seroprevalence of HIV-1 was higher in the Northeast than in the Southwest or the Midwest. Approximately 3% of the cohort members were infected with both HTLV-2 and HIV-1. The seroprevalence of HTLV-2 increased linearly with age. Women were significantly more likely to be infected with HTLV-2 than were men; the virus is thought to be more efficiently transmitted from male to female than from female to male.

Associated Diseases Although HTLV-2 was isolated from a patient with a T cell variant of hairy cell leukemia, this virus has not been consistently associated with a particular disease and in fact has been thought of as “a virus searching for a disease.” However, evidence is accumulating that HTLV-2 may play a role in certain neurologic, hematologic, and dermatologic diseases. These data require confirmation, particularly in light of the previous confusion regarding the relative prevalences of HTLV-1 and HTLV-2 among injection drug users.

Prevention Avoidance of needle sharing, adherence to safe-sex practices, screening of blood (by assays for HTLV-1, which also detect HTLV-2), and avoidance of breast-feeding by infected women are important principles in the prevention of spread of HTLV-2.

HUMAN IMMUNODEFICIENCY VIRUS

HIV-1 and HIV-2 are members of the lentivirus subfamily of Retroviridae and are the only lentiviruses known to infect humans. The lentiviruses are slow-acting by comparison with viruses that cause acute infection (e.g., influenza virus) but not by comparison with other retroviruses. The features of acute primary infection with HIV resemble those of more classic acute infections. The characteristic chronicity of HIV disease is consistent with the designation *lentivirus*.

For a detailed discussion of HIV, see Chap. 226.

226 Human Immunodeficiency Virus Disease: AIDS and Related Disorders

Anthony S. Fauci, H. Clifford Lane

AIDS was first recognized in the United States in the summer of 1981, when the U.S. Centers for Disease Control and Prevention (CDC) reported the unexplained occurrence of *Pneumocystis jirovecii* (formerly *P. carinii*) pneumonia in five previously healthy homosexual men in Los Angeles and of Kaposi's sarcoma (KS) with or without *P. jirovecii* pneumonia and other opportunistic infections in 26 previously healthy homosexual men in New York, San Francisco, and Los Angeles. The disease was soon recognized in male and female injection drug users; in hemophiliacs and blood transfusion recipients; among female sexual partners of men with AIDS; and among infants born to mothers with AIDS. In 1983, human immunodeficiency virus (HIV) was isolated from a patient with lymphadenopathy, and by 1984 it was demonstrated clearly to be the causative agent of AIDS. In 1985, a sensitive enzyme-linked immunosorbent assay (ELISA) was developed; this led to an appreciation of the scope and evolution of the HIV epidemic at first in the United States and other developed nations and ultimately among developing nations throughout the world (see "HIV Infection and AIDS Worldwide," below). The staggering worldwide evolution of the HIV pandemic has been matched by an explosion of information in the areas of HIV virology, pathogenesis (both immunologic and virologic), treatment of HIV disease, treatment and prophylaxis of the opportunistic diseases associated with HIV infection, and prevention of HIV infection. The information flow related to HIV disease is enormous and continues to expand, and it has become almost impossible for the health care generalist to stay abreast of the literature.

The purpose of this chapter is to present the most current information available on the scope of the epidemic; on its pathogenesis, treatment, and prevention; and on prospects for vaccine development. Above all, the aim is to provide a solid scientific basis and practical clinical guidelines for a state-of-the-art approach to the HIV-infected patient.

DEFINITION

The current U.S. CDC classification system for HIV infection and AIDS categorizes people on the basis of clinical conditions associated with HIV infection and CD4+ T lymphocyte measurement. A confirmed HIV case can be classified in one of five HIV infection stages (0, 1, 2, 3, or unknown). If there was a negative HIV test within 6 months of the first HIV infection diagnosis, the stage is 0, and remains 0 until 6 months after diagnosis. Advanced HIV disease (AIDS) is classified as stage 3 if one or more specific opportunistic illness has been diagnosed (Table 226-1). Otherwise, the stage is determined by CD4 test results and immunologic criteria (Table 226-2). If none of these criteria apply (e.g., because of missing information on CD4 test results), the stage is U (unknown).

The definition and staging criteria of AIDS are complex and comprehensive and were established for surveillance purposes rather than for the practical care of patients. Thus, the clinician should not focus

TABLE 226-1 CDC STAGE 3 (AIDS)-DEFINING OPPORTUNISTIC ILLNESSES IN HIV INFECTION

Bacterial infections, multiple or recurrent ^a
Candidiasis of bronchi, trachea, or lungs
Candidiasis of esophagus
Cervical cancer, invasive ^b
Coccidioidomycosis, disseminated or extrapulmonary
Cryptococcosis, extrapulmonary
Cryptosporidiosis, chronic intestinal (>1 month's duration)
Cytomegalovirus disease (other than liver, spleen, or nodes), onset at age >1 month
Cytomegalovirus retinitis (with loss of vision)
Encephalopathy attributed to HIV
Herpes simplex: chronic ulcers (>1 month's duration) or bronchitis, pneumonitis, or esophagitis (onset at age >1 month)
Histoplasmosis, disseminated or extrapulmonary
Isosporiasis, chronic intestinal (>1 month's duration)
Kaposi's sarcoma
Lymphoma, Burkitt's (or equivalent term)
Lymphoma, immunoblastic (or equivalent term)
Lymphoma, primary, of brain
<i>Mycobacterium avium</i> complex or <i>Mycobacterium kansasii</i> , disseminated or extrapulmonary
<i>Mycobacterium tuberculosis</i> of any site, pulmonary; ^b disseminated, or extrapulmonary
<i>Mycobacterium</i> , other species or unidentified species, disseminated or extrapulmonary
<i>Pneumocystis jirovecii</i> (previously known as <i>Pneumocystis carinii</i>) pneumonia
Pneumonia, recurrent ^a
Progressive multifocal leukoencephalopathy
<i>Salmonella</i> septicemia, recurrent
Toxoplasmosis of brain, onset at age >1 month
Wasting syndrome attributed to HIV

^aOnly among children age <6 years. ^bOnly among adults, adolescents, and children age ≥6 years.

Source: MMWR 63(No. RR-03), April 11, 2014.

TABLE 226-2 CDC HIV INFECTION STAGES 1–3 BASED ON AGE-SPECIFIC CD4+ T LYMPHOCYTE COUNT OR CD4+ T LYMPHOCYTE PERCENTAGE OF TOTAL LYMPHOCYTES^a

Stage ^a	Age on Date of CD4 T+ Lymphocyte Test					
	<1 Year		1–5 Years		6 Years through Adult	
	Cells/ μ L	%	Cells/ μ L	%	Cells/ μ L	%
1	$\geq 1,500$	≥ 34	$\geq 1,000$	≥ 30	≥ 500	≥ 26
2	750–1,499	26–33	500–999	22–29	200–499	14–25
3	<750	<26	<500	<22	<200	<14

^aThe stage is based primarily on the CD4+ T-lymphocyte count; the CD4+ T-lymphocyte count takes precedence over the CD4 T-lymphocyte percentage, and the percentage is considered only if the count is missing.

Source: MMWR 63(No. RR-03), April 11, 2014.

on whether the patient fulfills the strict definition of AIDS, but should view HIV disease as a spectrum ranging from primary infection, with or without the acute syndrome, to the asymptomatic stage, to advanced stages associated with opportunistic diseases (see “Pathophysiology and Pathogenesis,” below).

ETIOLOGIC AGENT

HIV is the etiologic agent of AIDS; it belongs to the family of human retroviruses (Retroviridae) and the subfamily of lentiviruses (Chap. 225e). Nononcogenic lentiviruses cause disease in other animal species, including sheep, horses, goats, cattle, cats, and monkeys. The four retroviruses known to cause human disease belong to two distinct groups: the human T lymphotropic viruses (HTLV)-1 and HTLV-2, which are transforming retroviruses; and the human immunodeficiency viruses, HIV-1 and HIV-2, which cause cytopathic effects either directly or indirectly (Chap. 225e). The most common cause of HIV disease throughout the world, and certainly in the United States, is HIV-1, which comprises several subtypes with different geographic distributions (see “Molecular Heterogeneity of HIV-1,” below). HIV-2 was first identified in 1986 in West African patients and was originally confined to West Africa. However, a number of cases that generally can be traced to West Africa or to sexual contacts with West Africans have been identified throughout the world. The currently defined groups of HIV-1 (M, N, O, P) and the HIV-2 groups A through H each are likely derived from a separate transfer to humans from a nonhuman primate reservoir. HIV-1 viruses likely came from chimpanzees and/or gorillas, and HIV-2 from sooty mangabeys. The AIDS pandemic is primarily caused by the HIV-1 M group viruses. Although HIV-1 group O and HIV-2 viruses have been found in numerous countries, including those in the developed world, they have caused much more localized epidemics. The taxonomic relationship between primate lentiviruses is shown in Fig. 226-1.

MORPHOLOGY OF HIV

Electron microscopy shows that the HIV virion is an icosahedral structure (Fig. 226-2) containing numerous external spikes formed by the two major envelope proteins, the external gp120 and the transmembrane gp41. The HIV envelope exists as a trimeric heterodimer. The virion buds from the surface of the infected cell and incorporates a variety of host proteins into its lipid bilayer. The structure of HIV-1 is schematically diagrammed in Fig. 226-2B.

REPLICATION CYCLE OF HIV

HIV is an RNA virus whose hallmark is the reverse transcription of its genomic RNA to DNA by the enzyme *reverse transcriptase*. The replication cycle of HIV begins with the high-affinity binding of the gp120 protein via a portion of its V1 region near the N terminus to its receptor on the host cell surface, the CD4 molecule (Fig. 226-3). The CD4 molecule is a 55-kDa protein found predominantly on a subset of T lymphocytes that are responsible for helper function in the immune system (Chap. 372e). It is also expressed on the surface of monocytes/macrophages and dendritic/Langerhans cells. Once it binds to CD4, the gp120 protein undergoes a conformational change that facilitates binding to one of two major co-receptors. The two major co-receptors for HIV-1 are CCR5 and CXCR4. Both receptors belong to the family of seven-transmembrane-domain G protein-coupled cellular receptors, and the use of one or the other or both receptors by the virus for entry into the cell is an important determinant of the cellular tropism of the virus. Certain dendritic cells (DCs) express a diversity of C-type lectin receptors on their surface—one of which is called *DC-SIGN*—that also bind with high affinity to the HIV gp120 envelope protein, allowing DCs to facilitate virus spread to CD4+ T cells. Following binding of the envelope protein to the CD4 molecule associated with the above-mentioned conformational change in the viral envelope gp120, fusion with the host cell membrane occurs via the newly exposed gp41 molecule penetrating the plasma membrane of the target cell and then coiling upon itself to bring the virion and target cell together (Fig. 226-4). Following fusion, uncoating of the capsid protein shell is initiated—a step that facilitates reverse transcription and leads to formation of the preintegration complex, composed of viral RNA, enzymes, and accessory proteins and surrounded by capsid and matrix proteins (Fig. 226-3). As the preintegration complex traverses

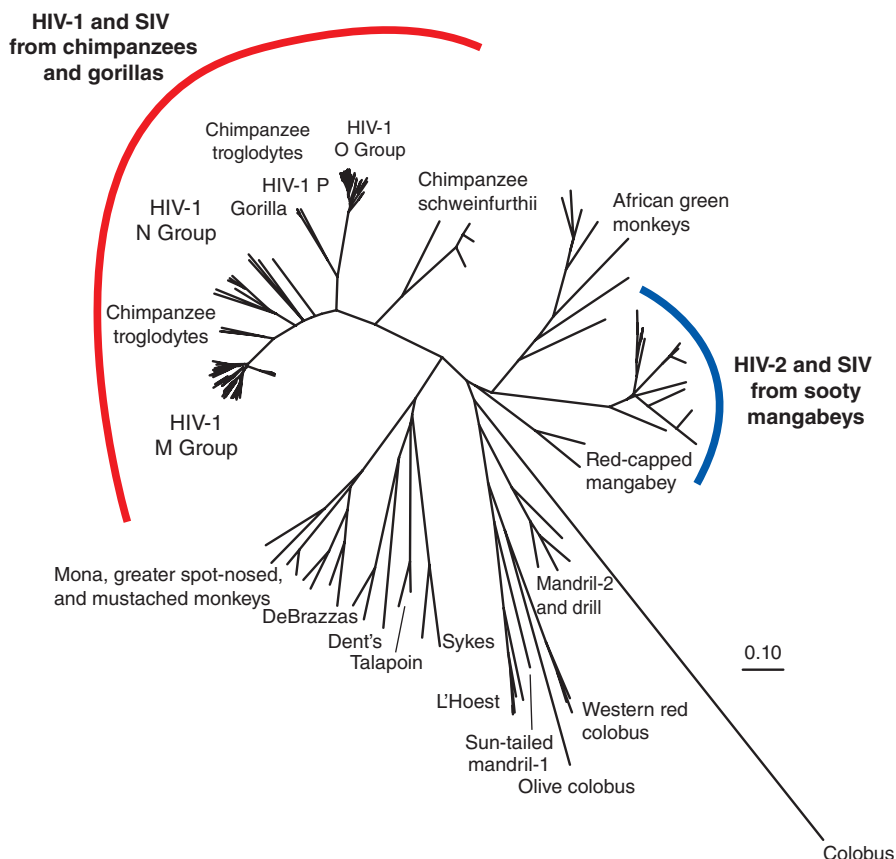


FIGURE 226-1 A phylogenetic tree based on the complete genomes of primate immunodeficiency viruses. The scale (0.10) indicates a 10% difference at the nucleotide level. (Prepared by Brian Foley, PhD, of the HIV Sequence Database, Theoretical Biology and Biophysics Group, Los Alamos National Laboratory; additional information at www.hiv.lanl.gov/content/sequence/HelpDocs/subtypes.html.)

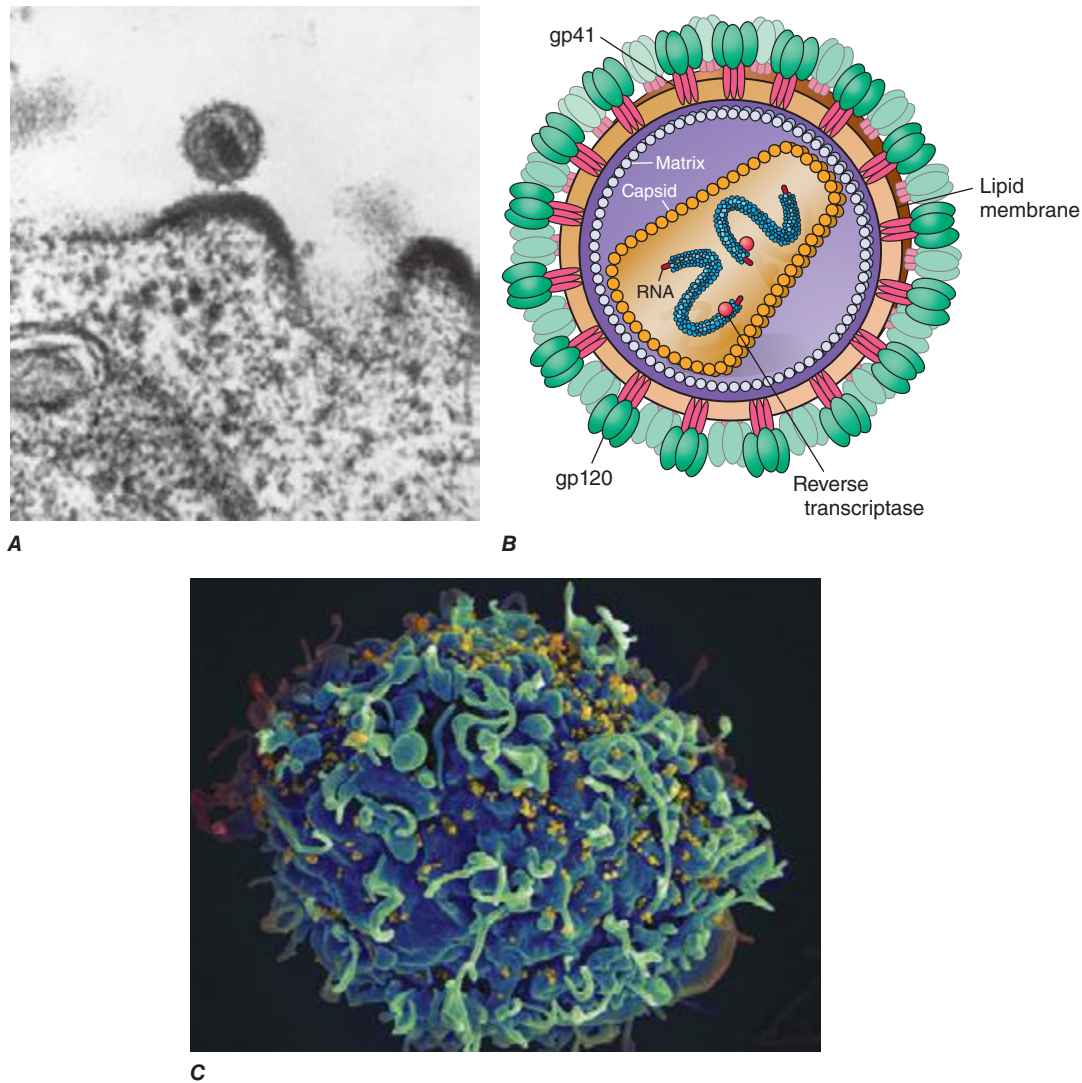


FIGURE 226-2 **A.** Electron micrograph of HIV. Figure illustrates a typical virion following budding from the surface of a CD4⁺ T lymphocyte, together with two additional incomplete virions in the process of budding from the cell membrane. **B.** Structure of HIV-1, including the gp120 envelope, gp41 transmembrane components of the envelope, genomic RNA, enzyme reverse transcriptase, p18(17) inner membrane (matrix), and p24 core protein (capsid). (Copyright by George V. Kelvin.) (Adapted from RC Gallo: *Sci Am* 256:46, 1987.) **C.** Scanning electron micrograph of HIV-1 virions infecting a human CD4⁺ T lymphocyte. The original photograph was imaged at 8000 \times magnification. (Courtesy of Elizabeth R. Fischer, Rocky Mountain Laboratories, National Institute of Allergy and Infectious Diseases; with permission.)

the cytoplasm to reach the nucleus, the viral reverse transcriptase enzyme catalyzes the reverse transcription of the genomic RNA into DNA, resulting in the formation of double-stranded proviral HIV-DNA. At the preintegration steps of the replication cycle, the viral genome is vulnerable to cellular factors that can block the progression of infection. In particular, the cytoplasmic tripartite motif-containing protein 5- α (TRIM5- α) is a host restriction factor that interacts with retroviral capsids (Fig. 226-3). Although the exact mechanisms of action of TRIM5- α remain unclear, the HIV-1 capsid is not recognized by the human form of TRIM5- α . Thus this host factor is not effective in restricting HIV-1 replication in human cells. The apolipoprotein B mRNA editing enzyme (catalytic polypeptide-like 3 [APOBEC3]) family of cellular proteins also inhibits progression of virus infection after virus has entered the cell and prior to entering the nucleus (Fig. 226-3). APOBEC3 proteins, which are incorporated into virions and released into the cytoplasm of a newly infected cell, bind to the single minus-strand DNA intermediate and deaminate viral cytidine, causing hypermutation of retroviral genomes. HIV has evolved a powerful strategy to protect itself from APOBEC. The viral protein Vif targets APOBEC3 for proteasomal degradation.

With activation of the cell, the viral DNA accesses the nuclear pore and is exported from the cytoplasm to the nucleus, where it

is integrated into the host cell chromosomes through the action of another virally encoded enzyme, *integrase* (Fig. 226-3). HIV provirus (DNA) integrates into the nuclear DNA preferentially within introns of active genes and regional hotspots. This provirus may remain transcriptionally inactive (latent) or it may manifest varying levels of gene expression, up to active production of virus.

Cellular activation plays an important role in the replication cycle of HIV and is critical to the pathogenesis of HIV disease (see “Pathogenesis and Pathophysiology,” below). Following initial binding, fusion, and internalization of the nucleic acid contents of virions into the target cell, incompletely reverse-transcribed DNA intermediates are labile in quiescent cells and do not integrate efficiently into the host cell genome unless cellular activation occurs shortly after infection. Furthermore, some degree of activation of the host cell is required for the initiation of transcription of the integrated proviral DNA into either genomic RNA or mRNA. This latter process may not necessarily be associated with the detectable expression of the classic cell-surface markers of activation. In this regard, activation of HIV expression from the latent state depends on the interaction of a number of cellular and viral factors. Following transcription, HIV mRNA is translated into proteins that undergo modification through glycosylation, myristoylation, phosphorylation,

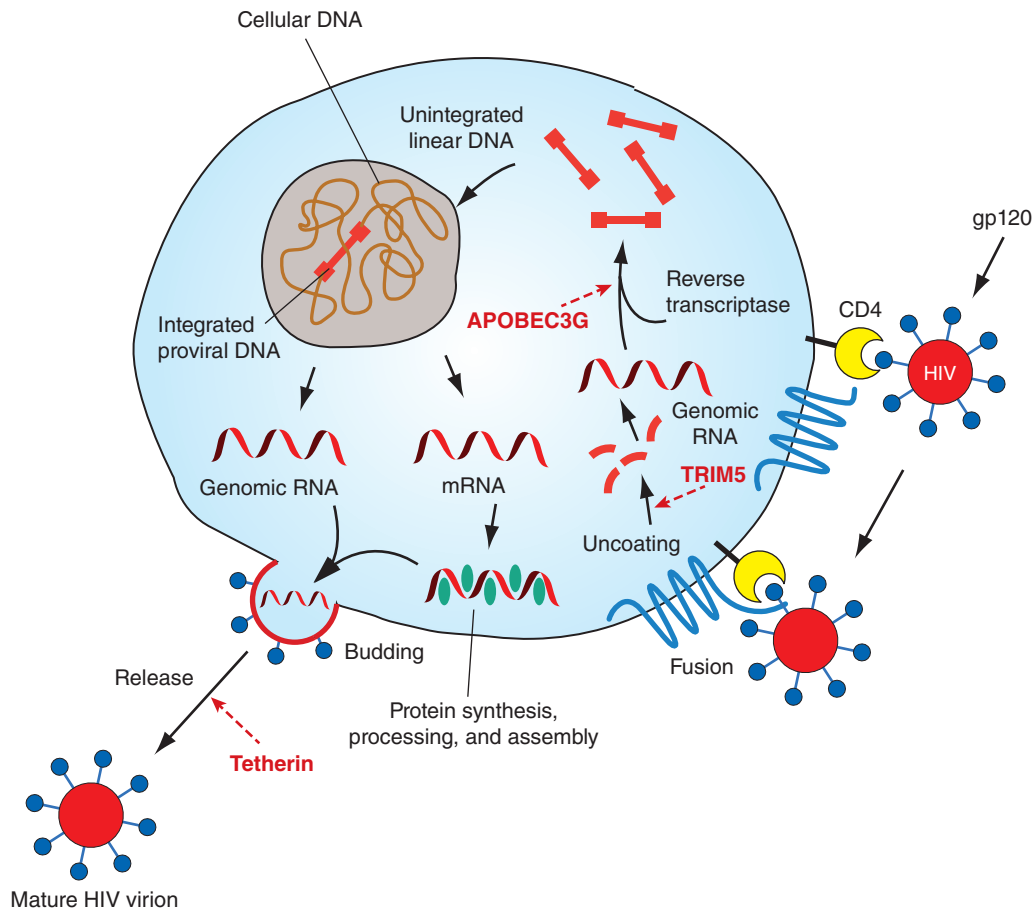


FIGURE 226-3 The replication cycle of HIV. See text for description. (Adapted from AS Fauci: *Nature* 384:529, 1996.)

and cleavage. The viral particle is formed by the assembly of HIV proteins, enzymes, and genomic RNA at the plasma membrane of the cells. Budding of the progeny virion through the lipid bilayer of the host cell membrane is the point at which the core acquires its external envelope and where the host restriction factor tetherin can inhibit the release of budding particles (Fig. 226-3). Tetherin is an interferon (IFN)-induced type II transmembrane protein that interferes

with virion detachment, although the HIV accessory protein Vpu counteracts the effect through direct interactions with tetherin. During or soon after budding, the virally encoded protease catalyzes the cleavage of the gag-pol precursor to yield the mature virion. Progression through the virus replication cycle is profoundly influenced by a variety of viral regulatory gene products. Likewise, each point in the replication cycle of HIV is a real or potential target for therapeutic intervention. Thus far, the reverse transcriptase, protease, and integrase enzymes as well as the process of virus–target cell binding and fusion have proved clinically to be susceptible to pharmacologic disruption.

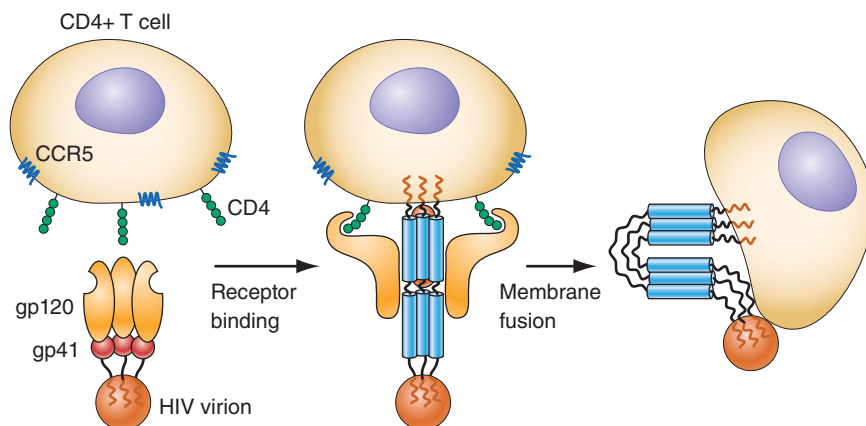


FIGURE 226-4 Binding and fusion of HIV-1 with its target cell. HIV-1 binds to its target cell via the CD4 molecule, leading to a conformational change in the gp120 molecule that allows it to bind to the co-receptor CCR5 (for R5-using viruses). The virus then firmly attaches to the host cell membrane in a coiled-spring fashion via the newly exposed gp41 molecule. Virus–cell fusion occurs as the transitional intermediate of gp41 undergoes further changes to form a hairpin structure that draws the two membranes into close proximity (see text for details). (Adapted from D Montefiori, JP Moore: *Science* 283:336, 1999; with permission.)

HIV GENOME

Figure 226-5 illustrates schematically the arrangement of the HIV genome. Like other retroviruses, HIV-1 has genes that encode the structural proteins of the virus: *gag* encodes the proteins that form the core of the virion (including p24 antigen); *pol* encodes the enzymes responsible for protease processing of viral proteins, reverse transcription, and integration; and *env* encodes the envelope glycoproteins. However, HIV-1 is more complex than other retroviruses, particularly those of the nonprimate group, in that it also contains at least six other genes (*tat*, *rev*, *nef*, *vif*, *vpr*, and *vpu*), which code for proteins involved in the modification of the host cell to enhance virus growth and the regulation of viral gene expression. Several of these proteins are thought to play a role in the pathogenesis of HIV disease; their various functions are listed in Fig. 226-5. Flanking these genes are the long terminal repeats (LTRs), which contain regulatory elements

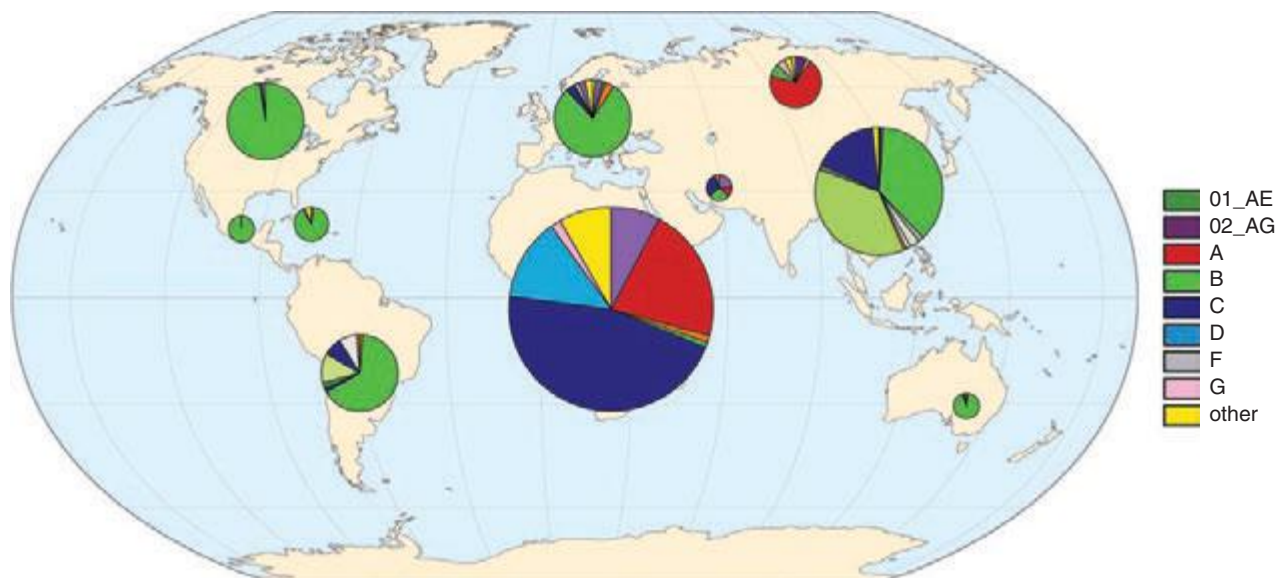


FIGURE 226-6 Global geographic distribution of HIV-1 subtypes and recombinant forms. Distributions derived from relative frequency of subtypes among >500,000 HIV genomic sequences in the Los Alamos National Laboratory HIV Sequence Database. (Additional information available at www.hiv.lanl.gov/components/sequence/HIV/geo/geo.comp.)

subtype C, with smaller proportions of infections caused by subtype A, subtype G, CRF02_AG, and other subtypes and recombinants. In South Africa, the country with the largest number of prevalent infections (6.3 million in 2013), >97% of the HIV-1 isolates sequenced are of subtype C. In Asia, HIV-1 isolates of the CRF01_AE lineage and subtypes C and B predominate. CRF01_AE accounts for most infections in south and southeast Asia, while >95% of infections in India, home to an estimated 2.1 million HIV-infected individuals, are of subtype C (see “HIV Infection and AIDS Worldwide,” below). Subtype B viruses are the overwhelmingly predominant viruses seen in the United States, Canada, certain countries in South America, western Europe, and Australia. It is thought that, purely by chance, subtype B was seeded into the United States and Europe in the late 1970s, thereby establishing an overwhelming founder effect. Many countries have co-circulating viral subtypes that are giving rise to new CRFs. Sequence analyses of HIV-1 isolates from infected individuals indicate that recombination among viruses of different clades likely occurs as a result of infection of an individual with viruses of more than one subtype, particularly in geographic areas where subtypes overlap, and more often in sub-epidemics driven by IV drug use than in those driven by sexual transmission.

The extraordinary diversity of HIV, reflected by the presence of multiple subtypes, circulating recombinant forms, and continuous viral evolution, has implications for possible differential rates of transmission, rates of disease progression, responses to therapy, and the development of resistance to antiretroviral drugs. This diversity is also a formidable obstacle to HIV vaccine development, as a broadly useful vaccine would need to induce protective responses against a wide range of viral strains.

TRANSMISSION

HIV is transmitted primarily by sexual contact (both heterosexual and male to male); by blood and blood products; and by infected mothers to infants intrapartum, perinatally, or via breast milk. After more than 30 years of experience and observations regarding other potential modalities of transmission, there is no evidence that HIV is transmitted by casual contact or that the virus can be spread by insects, such as by a mosquito bite. **Table 226-3** lists the estimated risk of HIV transmission for various types of exposures.

SEXUAL TRANSMISSION

HIV infection is predominantly a sexually transmitted infection (STI) worldwide. By far the most common mode of infection, particularly in

developing countries, is heterosexual transmission, although in many western countries a resurgence of male-to-male sexual transmission has occurred. Although a wide variety of factors including viral load and the presence of ulcerative genital diseases influence the efficiency of heterosexual transmission of HIV, such transmission is generally inefficient. A recent systemic review found a low per-act risk of heterosexual transmission in the absence of antiretrovirals: 0.04% for female-to-male transmission and 0.08% for male-to-female transmission during vaginal intercourse in the absence of antiretroviral therapy or condom use (Table 226-3).

HIV has been demonstrated in seminal fluid both within infected mononuclear cells and in cell-free material. The virus appears to concentrate in the seminal fluid, particularly in situations where there are increased numbers of lymphocytes and monocytes in the fluid,

TABLE 226-3 ESTIMATED PER-ACT PROBABILITY OF ACQUIRING HIV FROM AN INFECTED SOURCE, BY EXPOSURE ACT

Type of Exposure	Risk per 10,000 Exposures
Parenteral	
Blood transfusion	9250
Needle-sharing during injection drug use	63
Percutaneous (needle-stick)	23
Sexual	
Receptive anal intercourse	138
Insertive anal intercourse	11
Receptive penile-vaginal intercourse	8
Insertive penile-vaginal intercourse	4
Receptive oral intercourse	Low
Insertive oral intercourse	Low
Other^a	
Biting	Negligible
Spitting	Negligible
Throwing body fluids (including semen or saliva)	Negligible
Sharing sex toys	Negligible

^aHIV transmission through these exposure routes is technically possible but unlikely and not well documented.

Sources: CDC, www.cdc.gov/hiv/policies/law/risk.html; P Patel: AIDS 28:1509, 2014.

as in genital inflammatory states such as urethritis and epididymitis, conditions closely associated with other STIs. The virus has also been demonstrated in cervical smears and vaginal fluid. There is an elevated risk of HIV transmission associated with unprotected receptive anal intercourse (URAI) among both men and women compared to the risk associated with receptive vaginal intercourse. Although data are limited, the per-act risk for HIV transmission via URAI has been estimated to be ~1.4% (Table 226-3). The risk of HIV acquisition associated with URAI is probably higher than that seen in penile-vaginal intercourse because only a thin, fragile rectal mucosal membrane separates the deposited semen from potentially susceptible cells in and beneath the mucosa, and micro-trauma of the mucosal membrane may be associated with anal intercourse. Anal douching and sexual practices that traumatize the rectal mucosa also increase the likelihood of infection. It is likely that anal intercourse provides at least two modalities of infection: (1) direct inoculation into blood in cases of traumatic tears in the mucosa; and (2) infection of susceptible target cells, such as Langerhans cells, in the mucosal layer in the absence of trauma. Insertive anal intercourse also confers an increased risk of HIV acquisition compared to insertive vaginal intercourse. Although the vaginal mucosa is several layers thicker than the rectal mucosa and less likely to be traumatized during intercourse, the virus can be transmitted to either partner through vaginal intercourse. As noted in Table 226-3, male-to-female HIV transmission is usually more efficient than female-to-male transmission. The differences in reported transmission rates between men and women may be due in part to the prolonged exposure to infected seminal fluid of the vaginal and cervical mucosa, as well as the endometrium (when semen enters through the cervical os). By comparison, the penis and urethral orifice are exposed relatively briefly to infected vaginal fluid. Among various cofactors examined in studies of heterosexual HIV transmission, the presence of other STIs has been strongly associated with HIV transmission. In this regard, there is a close association between genital ulcerations and transmission, owing to both susceptibility to infection and infectivity. Infections with microorganisms such as *Treponema pallidum* (Chap. 206), *Haemophilus ducreyi* (Chap. 182), and herpes simplex virus (HSV; Chap. 216) are important causes of genital ulcerations linked to transmission of HIV. In addition, pathogens responsible for non-ulcerative inflammatory STIs such as those caused by *Chlamydia trachomatis* (Chap. 213), *Neisseria gonorrhoeae* (Chap. 181), and *Trichomonas vaginalis* (Chap. 254) also are associated with an increased risk of transmission of HIV infection. Bacterial vaginosis, an infection related to sexual behavior, but not strictly an STI, also may be linked to an increased risk of transmission of HIV infection. Several studies suggest that treating other STIs and genital tract syndromes may help prevent transmission of HIV. This effect is most prominent in populations in which the prevalence of HIV infection is relatively low. It is noteworthy that this principle may not apply to the treatment of HSV infections since it has been shown that even following anti-HSV therapy with resulting healing of HSV-related genital ulcers, HIV acquisition is not reduced. Biopsy studies revealed the likely explanation is that HIV receptor-positive inflammatory cells persisted in the genital tissue despite the healing of ulcers, and so HIV-susceptible targets remained at the site.

The quantity of HIV-1 in plasma is a primary determinant of the risk of HIV-1 transmission. In a cohort of heterosexual couples in Uganda discordant for HIV infection and not receiving antiretroviral therapy, the mean serum HIV RNA level was significantly higher among HIV-infected subjects whose partners seroconverted than among those whose partners did not seroconvert. In fact, transmission was rare when the infected partner had a plasma level of <1700 copies of HIV RNA per milliliter, even when genital ulcer disease was present (Fig. 226-7). The rate of HIV transmission per coital act was highest during the early stage of HIV infection when plasma HIV RNA levels were high and in advanced disease as the viral set point increased.

Antiretroviral therapy dramatically reduces plasma viremia in most HIV-infected individuals (see “Treatment,” below) and is associated with a reduction in risk of transmission. In a large study of serodiscordant couples, earlier treatment of the HIV-infected partner with

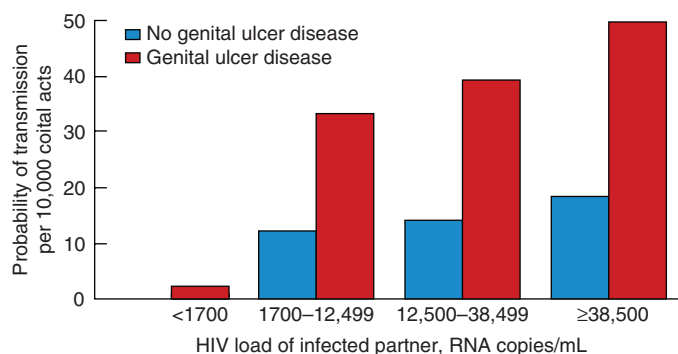


FIGURE 226-7 Probability of HIV transmission per coital act among monogamous, heterosexual, HIV-serodiscordant couples in Uganda. (From RH Gray et al: *Lancet* 357:1149, 2001.)

antiretroviral therapy rather than treatment delayed until the CD4+ T cells count fell below 250 cells per μL was associated with a 96% reduction in HIV transmission to the uninfected partner. This approach has been widely referred to as *treatment as prevention* or TasP. Several studies also have suggested a beneficial effect of antiretroviral treatment at the community level.

A number of studies including large, randomized, controlled trials clearly have indicated that male *circumcision* is associated with a lower risk of acquisition of HIV infection for heterosexual men. Studies are conflicting as to whether circumcision protects against HIV acquisition among men who have sex with men, but data suggest that circumcision is protective in those men who have sex with men who are insertive only. The benefit of circumcision may be due to increased susceptibility of uncircumcised men to ulcerative STIs, as well as to other factors such as microtrauma to the foreskin and glans penis. In addition, the highly vascularized inner foreskin tissue contains a high density of Langerhans cells as well as increased numbers of CD4+ T cells, macrophages, and other cellular targets for HIV. Finally, the moist environment under the foreskin may promote the presence or persistence of microbial flora that, via inflammatory changes, may lead to even higher concentrations of target cells for HIV in the foreskin. In addition, randomized trials have demonstrated that male circumcision also reduces hepatitis C virus (HCV) type 2, human papillomavirus virus (HPV), and genital ulcer disease in men as well as HPV, genital ulcer disease, bacterial vaginosis, and *Trichomonas vaginalis* infections among female partners of circumcised men. Thus, there may be an added benefit of diminution of risk for HIV acquisition to the female sexual partners of circumcised men.

In some studies the use of oral contraceptives was associated with an increase in incidence of HIV infection over and above that which might be expected by not using a condom for birth control. This phenomenon may be due to drug-induced changes in the cervical mucosa, rendering it more vulnerable to penetration by the virus. Adolescent girls might also be more susceptible to infection upon exposure due to the properties of an immature genital tract with increased cervical ectopy or exposed columnar epithelium.

Oral sex is a much less efficient mode of transmission of HIV than is anal intercourse or vaginal intercourse (Table 226-3). A number of studies have reported that the incidence of transmission of infection by oral sex among couples discordant for HIV was extremely low. However, there have been well-documented reports of HIV transmission that likely resulted from fellatio or cunnilingus. Therefore, the assumption that oral sex is completely safe is not warranted.

The association of alcohol consumption and illicit drug use with unsafe sexual behavior, both homosexual and heterosexual, leads to an increased risk of sexual transmission of HIV. Methamphetamine and other so-called club drugs (e.g., ecstasy, ketamine, and gamma hydroxybutyrate), sometimes taken in conjunction with PDE-5 inhibitors such as sildenafil (Viagra), tadalafil (Cialis), or vardenafil (Levitra), have been associated with risky sexual practices and increased risk of HIV infection, particularly among men who have sex with men.

1222 TRANSMISSION THROUGH INJECTION DRUG USE

HIV can be transmitted to injection drug users (IDUs) who are exposed to HIV while sharing injection paraphernalia such as needles, syringes, the water in which drugs are mixed, or the cotton through which drugs are filtered. Parenteral transmission of HIV during injection drug use does not require IV puncture; SC (“skin popping”) or IM (“muscling”) injections can transmit HIV as well, even though these behaviors are sometimes erroneously perceived as low-risk. Among IDUs, the risk of HIV infection increases with the duration of injection drug use; the frequency of needle sharing; the number of partners with whom paraphernalia are shared, particularly in the setting of “shooting galleries” where drugs are sold and large numbers of IDUs may share a limited number of “works”; comorbid psychiatric conditions such as antisocial personality disorder; the use of cocaine in injectable form or smoked as “crack”; and the use of injection drugs in a geographic location with a high prevalence of HIV infection, such as certain inner-city areas in the United States. As noted in Table 226-3, the per-act risk of transmission from injection drug use with a contaminated needle has been estimated to be approximately 0.6%.

TRANSMISSION BY TRANSFUSED BLOOD AND BLOOD PRODUCTS

HIV can be transmitted to individuals who receive HIV-tainted blood transfusions, blood products, or transplanted tissue. The first cases of AIDS among transfusion recipients and individuals with hemophilia or other clotting disorders were reported in 1982. The vast majority of HIV infections acquired via contaminated blood transfusions, blood components, or transplanted tissue in resource-rich countries occurred prior to the spring of 1985, when mandatory testing of donated blood for HIV-1 was initiated. It is estimated that >90% of individuals exposed to HIV-contaminated blood products become infected (Table 226-3). Although blood screening for HIV is becoming more universal even in the developing world, unfortunately, in some resource-poor countries, HIV continues to be transmitted by blood, blood products, and tissues due to inadequate screening. Transfusions of whole blood, packed red blood cells, platelets, leukocytes, and plasma are all capable of transmitting HIV infection. In contrast, hyperimmune gamma globulin, hepatitis B immune globulin, plasma-derived hepatitis B vaccine, and Rh₀ immune globulin have not been associated with transmission of HIV infection. The procedures involved in processing these products either inactivate or remove the virus.

Currently, in the United States and in most developed countries, the following measures have made the risk of transmission of HIV infection by transfused blood or blood products extremely small: the screening of blood donations for antibodies to HIV-1 and HIV-2 and determination of the presence of HIV nucleic acid usually in mini-pools of several specimens; the careful selection of potential blood donors with health history questionnaires to exclude individuals with risk behavior; and opportunities for self-deferral and the screening out of HIV-negative individuals with serologic testing for infections that have shared risk factors with HIV, such as hepatitis B and C and syphilis. The chance of infection of a hemophiliac via clotting factor concentrates has essentially been eliminated because of the added layer of safety resulting from heat treatment of the concentrates. It is currently estimated that the risk of infection with HIV in the United States via transfused screened blood is approximately 1 in 2 million units. Therefore, since ~16 million donations are collected in the United States each year, despite the best efforts of science, one cannot completely eliminate the risk of transfusion-related transmission of HIV. In this regard, a case of transfusion-related transmission of HIV was reported in the United States in 2010, which was tracked to a blood donation in 2008; this was the first such reported case since 2002. Transmission of HIV (both HIV-1 and HIV-2) by blood or blood products is still an ongoing threat in certain developing countries, particularly in sub-Saharan Africa, where routine screening of blood is not universally practiced. In other countries, there have been reports of sporadic breakdowns in routinely available screening procedures in which contaminated blood was allowed to be transfused, resulting in small clusters of patients becoming infected. For example, in China in the 1990s, a disturbingly large number of persons became infected

by selling blood in situations where the collectors reused needles that were contaminated and, in some instances, mixed blood products from a number of individuals, separated the plasma, and reinfused mixed red blood cells back into the individual donors.

OCCUPATIONAL TRANSMISSION OF HIV: HEALTH CARE WORKERS, LABORATORY WORKERS, AND THE HEALTH CARE SETTING

There is a small but definite occupational risk of HIV transmission to health care workers and laboratory personnel and potentially others who work with HIV-containing materials, particularly when sharp objects are used. An estimated 600,000 to 800,000 health care workers are stuck with needles or other sharp medical instruments in the United States each year. The global number of HIV infections among health care workers attributable to sharps injuries has been estimated to be 1000 cases (range, 200–5000) per year. As of 2010, there had been 57 documented cases of occupational HIV transmission to health care workers in the United States and 143 possible transmissions. There have been no confirmed cases reported since 1999.

Exposures that place a health care worker at potential risk of HIV infection are percutaneous injuries (e.g., a needle stick or cut with a sharp object) or contact of mucous membrane or nonintact skin (e.g., exposed skin that is chapped, abraded, or afflicted with dermatitis) with blood, tissue, or other potentially infectious body fluids. Large, multi-institutional studies have indicated that the risk of HIV transmission following skin puncture from a needle or a sharp object that was contaminated with blood from a person with documented HIV infection is ~0.3% and after a mucous membrane exposure it is 0.09% (see “HIV and the Health Care Worker,” below) if the injured and/or exposed person is not treated within 24 h with antiretroviral drugs. The risk of hepatitis B virus (HBV) infection following a similar type of exposure is ~6–30% in nonimmune individuals; if a susceptible worker is exposed to HBV, postexposure prophylaxis with hepatitis B immune globulin and initiation of HBV vaccine is >90% effective in preventing HBV infection. The risk of HCV infection following percutaneous injury is ~1.8% (Chap. 360).

Rare HIV transmission after nonintact skin exposure has been documented, but the average risk for transmission by this route has not been precisely determined; however, it is estimated to be less than the risk for mucous membrane exposure. Transmission of HIV through intact skin has not been documented. Currently in developed countries, virtually all puncture wounds and mucous membrane exposures in health care workers involving blood from a patient with documented HIV infection are treated prophylactically with combination antiretroviral therapy (cART). This practice, referred to as *postexposure prophylaxis* or PEP, has dramatically reduced the occurrence of puncture-related transmissions of HIV to health care workers.

In addition to blood and visibly bloody body fluids, semen and vaginal secretions also are considered potentially infectious; however, they have not been implicated in occupational transmission from patients to health care workers. The following fluids also are considered potentially infectious: cerebrospinal fluid, synovial fluid, pleural fluid, peritoneal fluid, pericardial fluid, and amniotic fluid. The risk for transmission after exposure to fluids or tissues other than HIV-infected blood also has not been quantified, but it is probably considerably lower than the risk after blood exposures. Feces, nasal secretions, saliva, sputum, sweat, tears, urine, and vomitus are not considered potentially infectious for HIV unless they are visibly bloody. Rare cases of HIV transmission via human bites have been reported, but not in the setting of occupational exposure.

An increased risk for HIV infection following percutaneous exposures to HIV-infected blood is associated with exposures involving a relatively large quantity of blood, as in the case of a device visibly contaminated with the patient’s blood, a procedure that involves a hollow-bore needle placed directly in a vein or artery, or a deep injury. Factors that might be associated with mucocutaneous transmission of HIV include exposure to an unusually large volume of blood and prolonged contact. In addition, the risk increases for exposures to blood from untreated patients with advanced-stage disease or those patients in the acute stage of HIV infection, owing to the higher levels of HIV in the blood under those

circumstances. Since the beginning of the HIV epidemic, there have been rare instances where transmission of infection from a health care worker to patients seemed highly probable. Despite these small number of documented cases, the risk of HIV transmission involving health care workers (infected or not) to patients is extremely low in developed countries—in fact, too low to be measured accurately. In this regard, several epidemiologic studies have been performed tracing thousands of patients of HIV-infected dentists, physicians, surgeons, obstetricians, and gynecologists, and no other cases of HIV transmission that could be linked to the health care providers were identified.

Breaches in infection control and the reuse of contaminated syringes, failure to properly sterilize surgical instruments, and/or hemodialysis equipment have also resulted rarely in the transmission of HIV from patient to patient in hospitals, nursing homes, and outpatient settings. Finally, these very rare occurrences of transmission of HIV as well as HBV and HCV to and from health care workers in the workplace underscore the importance of the use of universal precautions when caring for all patients (see below and [Chap. 168](#)).

MOTHER-TO-CHILD TRANSMISSION OF HIV

HIV infection can be transmitted from an infected mother to her fetus during pregnancy, during delivery, or by breast-feeding. This remains an important form of transmission of HIV infection in certain developing countries, where the proportion of infected women to infected men is ~1:1. Virologic analyses of aborted fetuses indicate that HIV can be transmitted to the fetus during the first or second trimesters of pregnancy. However, maternal transmission to the fetus occurs most commonly in the perinatal period. Two studies performed in Rwanda and the Democratic Republic of Congo (then called Zaire) indicated that among all mother-to-child transmissions of HIV, the relative proportions were 23–30% before birth, 50–65% during birth, and 12–20% via breast-feeding.

In the absence of prophylactic antiretroviral therapy to the mother during pregnancy, labor, and delivery, and to the fetus following birth, the probability of transmission of HIV from mother to infant/fetus ranges from 15% to 25% in industrialized countries and from 25% to 35% in developing countries. These differences may relate to the adequacy of prenatal care as well as to the stage of HIV disease and the general health of the mother during pregnancy. Higher rates of transmission have been reported to be associated with many factors—the best documented of which is the presence of high maternal levels of plasma viremia, with the risk increasing linearly with the level of maternal plasma viremia. It is very unlikely that mother-to-child transmission will occur if the mother's level of plasma viremia is <1000 copies of HIV RNA/mL of blood and extremely unlikely if the level is undetectable (i.e., <50 copies/mL). However, there may not be a lower “threshold” below which transmission never occurs, since certain studies have reported rare transmission by women with viral RNA levels <50 copies/mL. Increased mother-to-child transmission is also correlated with closer human leukocyte antigen (HLA) match between mother and child. A prolonged interval between membrane rupture and delivery is another well-documented risk factor for transmission. Other conditions that are potential risk factors, but that have not been consistently demonstrated, are the presence of chorioamnionitis at delivery; STIs during pregnancy; illicit drug use during pregnancy; cigarette smoking; preterm delivery; and obstetric procedures such as amniocentesis, amniocopy, fetal scalp electrodes, and episiotomy. In a seminal study conducted in the United States and France in the 1990s, zidovudine treatment of HIV-infected pregnant women from the beginning of the second trimester through delivery and of the infant for 6 weeks following birth dramatically decreased the rate of intrapartum and perinatal transmission of HIV infection from 22.6% in the untreated group to <5%. Today, the rate of mother-to-child transmission has fallen to 1% or less in pregnant women who are receiving combination antiretroviral therapy (cART) for their HIV infection. Such treatment, combined with cesarean section delivery, has rendered mother-to-child transmission of HIV an unusual event in the United States and other developed nations. In this regard, both the United States Public Health Service and the World Health

Organization guidelines recommend that all pregnant HIV-infected women should receive cART for the health of the mother and to prevent perinatal transmission regardless of plasma HIV RNA copy number or CD4+ T cell counts.

Breast-feeding is an important modality of transmission of HIV infection in developing countries, particularly where mothers continue to breast-feed for prolonged periods. The risk factors for mother-to-child transmission of HIV via breast-feeding are not fully understood; factors that increase the likelihood of transmission include detectable levels of HIV in breast milk, the presence of mastitis, low maternal CD4+ T cell counts, and maternal vitamin A deficiency. The risk of HIV infection via breast-feeding is highest in the early months of breast-feeding. In addition, exclusive breast-feeding has been reported to carry a lower risk of HIV transmission than mixed feeding. In developed countries, breast feeding of babies by an HIV-infected mother is contraindicated since alternative forms of adequate nutrition, i.e., formulas, are readily available. In developing countries, where breast-feeding may be essential for the overall health of the infant, the continuation of cART in the infected mother during the period of breastfeeding markedly diminishes the risk of transmission of HIV to the infant. In fact, once cART has been initiated in a pregnant woman, many experts recommend that therapy be continued for life.

TRANSMISSION OF HIV BY OTHER BODY FLUIDS

Although HIV can be isolated typically in low titers from saliva of a small proportion of infected individuals, there is no convincing evidence that saliva can transmit HIV infection, either through kissing or through other exposures, such as occupationally to health care workers. Saliva contains endogenous antiviral factors; among these factors, HIV-specific immunoglobulins of IgA, IgG, and IgM isotypes are detected readily in salivary secretions of infected individuals. It has been suggested that large glycoproteins such as mucins and thrombospondin 1 sequester HIV into aggregates for clearance by the host. In addition, a number of soluble salivary factors inhibit HIV to various degrees in vitro, probably by targeting host cell receptors rather than the virus itself. Perhaps the best studied of these, secretory leukocyte protease inhibitor (SLPI), blocks HIV infection in several cell culture systems, and it is found in saliva at levels that approximate those required for inhibition of HIV in vitro. In this regard, higher salivary levels of SLPI in breast-fed infants were associated with a decreased risk of HIV transmission through breast milk. It has also been suggested that submandibular saliva reduces HIV infectivity by stripping gp120 from the surface of virions, and that saliva-mediated disruption and lysis of HIV-infected cells occurs because of the hypotonicity of oral secretions. There have been outlier cases of suspected transmission by saliva, but these have probably been blood-to-blood transmissions. Transmission of HIV by a human bite can occur but is a rare event. Although virus can be identified, if not isolated, from virtually any body fluid, there is no evidence that HIV transmission can occur as a result of exposure to tears, sweat, or urine. However, there have been isolated cases of transmission of HIV infection by body fluids that may or may not have been contaminated with blood. Most of these situations occurred in the setting of a close relative providing intensive nursing care for an HIV-infected person without observing universal precautions, underscoring the importance of adhering to such precautions in the handling of body fluids and wastes from HIV-infected individuals.

EPIDEMIOLOGY

HIV INFECTION AND AIDS WORLDWIDE


 HIV infection/AIDS is a global pandemic, with cases reported from virtually every country. At the end of 2013, an estimated 35.0 million individuals were living with HIV infection, according to the Joint United Nations Programme on HIV/AIDS (UNAIDS). An estimated 95% of people living with HIV/AIDS reside in low- and middle-income countries; ~50% are female, and 3.2 million are children <15 years. The distribution of these cases is illustrated in [Fig. 226-8](#). The estimated number of people living with HIV—i.e.,



FIGURE 226-8 Estimated number of adults and children living with HIV infection as of December, 2013. Total: 35.0 (33.2 million–37.2 million). (From Joint United Nations Programme on HIV/AIDS [UNAIDS].)

the global prevalence—has increased more than fourfold since 1990, reflecting the combined effects of continued high rates of new HIV infections and the life-prolonging impact of antiretroviral therapy (Fig. 226-9). In 2013, the global prevalence rate among persons age 15–49 years was 0.8%, with rates varying widely by country and region as illustrated in Fig. 226-10.

In 2013, an estimated 2.1 million new cases of HIV infection occurred worldwide, including 240,000 among children <15 years; about 40% of new infections were among persons under age 25. Between 2001 and 2013, the estimated number of new HIV infections globally fell by 38% (Fig. 226-9). Recent reductions in global HIV incidence likely reflect progress with HIV prevention efforts and the increased provision to HIV-infected people of antiretroviral therapy, which makes them much less likely to transmit the virus to sexual partners. In 2013, global AIDS deaths totaled 1.5 million (including 190,000 children <15 years), a 35% decrease since 2005 that coincides with a rapid expansion of access to antiretroviral therapy (Fig. 226-9). Since the beginning of the pandemic, an estimated 39 million people have died of an AIDS-related illness.

The HIV epidemic has occurred in “waves” in different regions of the world, each wave having somewhat different characteristics depending on the demographics of the country and region in question and the timing of the introduction of HIV into the population. Although the AIDS epidemic was first recognized in the United States and shortly thereafter in Western Europe, it very likely began in sub-Saharan Africa (see above), which has been particularly devastated by the epidemic. More than 70% of all people with HIV infection (~25 million), and nearly 90% of all HIV-infected children live in that region, even though sub-Saharan Africa is home to just 12% of the world’s population (Fig. 226-8). Within the region, southern Africa is worst-affected. In nine southern African countries, seroprevalence data indicate that >10% of the adult population age 15–49 is HIV-infected (Fig. 226-10). In addition, among high-risk individuals (e.g., commercial sex workers, patients attending STI clinics) who live in urban areas of sub-Saharan Africa, seroprevalence is now >50% in some places. Recent data offer promising signs of declining HIV incidence and prevalence in many countries in the region, although frequently at levels that remain high. Heterosexual exposure is

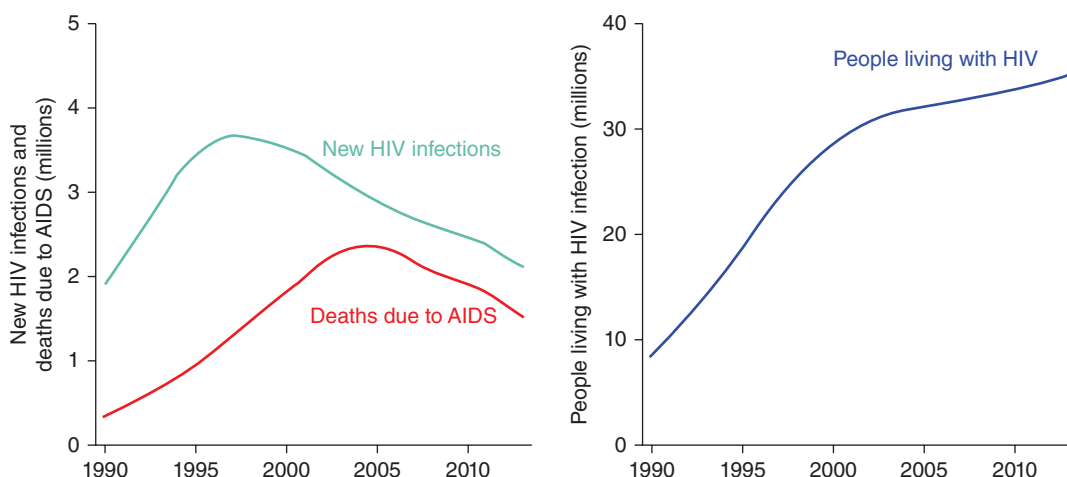


FIGURE 226-9 Global estimates of HIV incidence and AIDS deaths (left) and, HIV prevalence (right), 1990–2013. (From UNAIDS.)

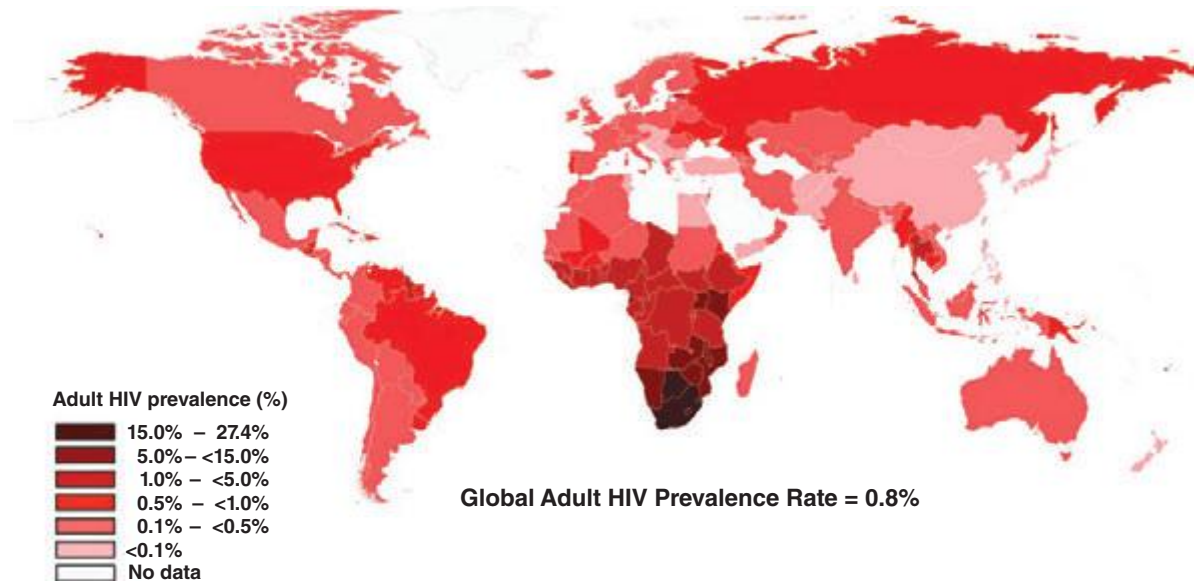


FIGURE 226-10 Global adult HIV prevalence rate, 2013. Data are estimates for adults aged 15–49 years. (From UNAIDS.)

the primary mode of HIV transmission in sub-Saharan Africa, with women and girls disproportionately affected, accounting for ~60 percent of all HIV infections in that region. In 2013, an estimated 230,000 people were living with HIV in the Middle East/North Africa region. Cases are largely concentrated among IDUs, men who have sex with men, and sex workers and their clients.

In Asia and the Pacific, an estimated 4.8 million people were living with HIV at the end of 2013. In this region of the world, HIV prevalence is highest in southeast Asian countries, with wide variation in trends between different countries. Among countries in Asia, only Thailand has an adult seroprevalence rate of >1%. However, the populations of many Asian nations are so large (especially India and China) that even low infection and seroprevalence rates result in large numbers of people living with HIV. Although Asia's epidemic has been concentrated for some time among specific populations—sex workers and their clients, men who have sex with men, and IDUs—it is expanding to the heterosexual partners of those most at risk.

The epidemic is expanding in Eastern Europe and Central Asia, where ~1.1 million people were living with HIV at the end of 2013. The Russian Federation and Ukraine account for the majority of HIV cases in the region. Driven initially by injection drug use and increasingly by heterosexual transmission, the number of new infections in this region has increased dramatically over the past decade.

Approximately 1.9 million people are living with HIV/AIDS in Central and South America and the Caribbean. Brazil is home to the largest number of HIV-infected people in the region. However, the epidemic has been slowed in that country due to successful treatment and prevention efforts. Men who have sex with men account for the largest proportion of HIV infections in Central and South America. The Caribbean region has the highest regional adult seroprevalence rate after Africa. Heterosexual transmission, often tied to sex work, is the main driver of transmission in the region.

Approximately 2.3 million people are living with HIV/AIDS in North America and western and central Europe. The number of new infections among men who have sex with men has increased over the past decade in these mostly high-income areas, while rates of new infections among heterosexuals have stabilized and infections among women and IDUs have fallen.

HIV INFECTION AND AIDS IN THE UNITED STATES

About 1.7 million people have been infected with HIV in the United States since the beginning of the epidemic, of whom >630,000 have died. Approximately 1.1 million individuals in the United States are living with HIV infection, ~16–18% of whom are unaware of their infection, according to recent estimates. As illustrated in Fig. 226-11,

only a fraction of HIV-infected people are able to negotiate the steps in the HIV “care continuum,” from diagnosis, to entering into and staying in care, to receiving antiretroviral therapy, and ultimately to achieving a suppressed viral load (see “Treatment,” below).

More than 60% of those living with HIV/AIDS are Black/African-American or Hispanic/Latino, and more than half are men who have sex with men. The estimated HIV seroprevalence rate among all individuals age 13 years or older in the United States is ~0.5%. Approximately 2% of Black/African-American adults are HIV-infected in the United States, higher than any other group.

The number of new HIV infections in the United States, *HIV incidence*, peaked at about 130,000 per year in the late 1980s, followed by declines. For more than a decade, HIV incidence has remained stable at approximately 50,000 per year, with the proportion of new infections increasing in recent years among men who have sex with men and falling among women and IDUs. Among adults and adolescents newly diagnosed with HIV infection in 2011 (regardless of stage of infection), ~79% were males and ~21% were women. Of new HIV diagnoses among men, ~79% were attributed to male-to-male sexual contact, ~12% to heterosexual contact, ~6% to injection drug use, and ~4% to a combination of male-to-male sexual contact and injection drug use. Of new HIV diagnoses among women, ~86% were due to heterosexual contact and ~14% to injection drug use

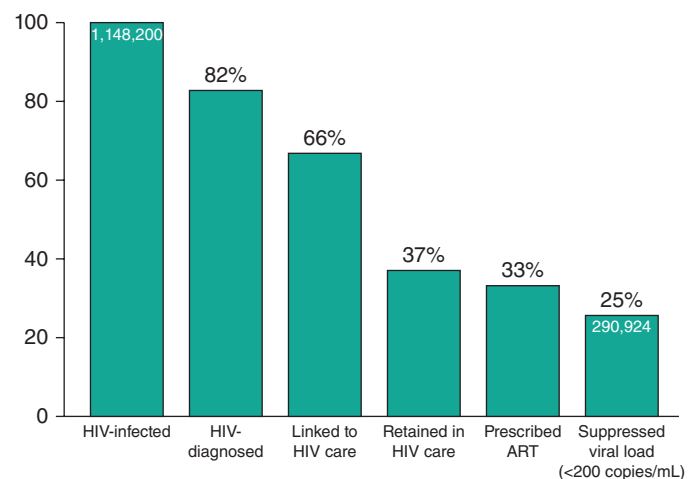


FIGURE 226-11 Estimated percentage of HIV-infected people engaged in selected stages of the continuum of HIV care in the United States. (Adapted from HI Hall et al: *JAMA Intern Med* 173:1337, 2013.)

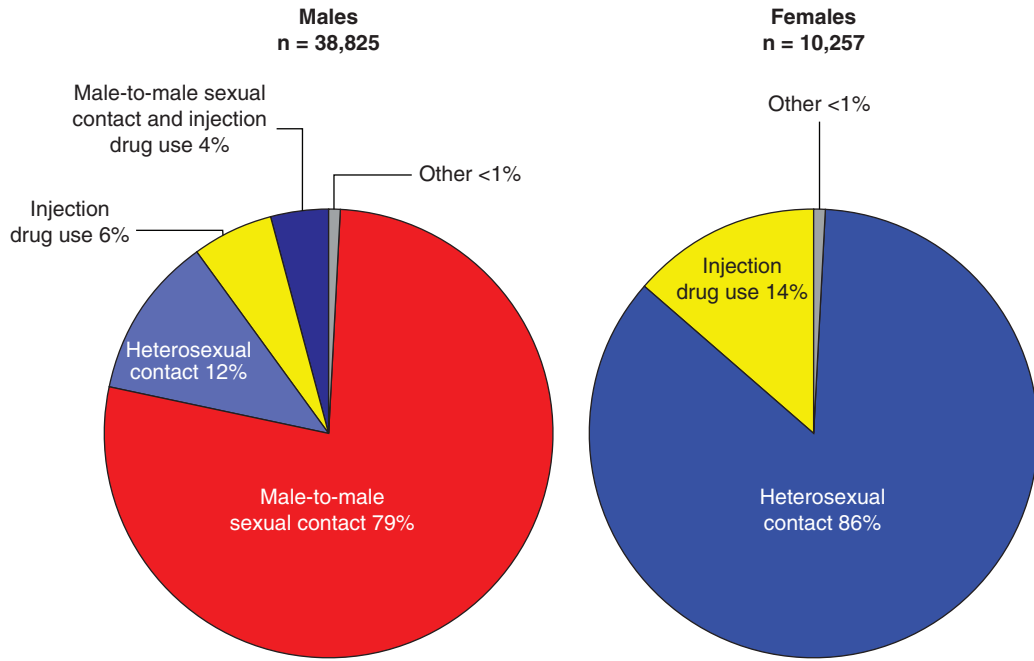


FIGURE 226-12 Transmission categories of adults and adolescents diagnosed with HIV infection (regardless of stage) in 2011, United States. (From CDC.)

(Fig. 226-12). The estimated numbers of new HIV infections in 2011 in the United States for the 10 most affected subpopulations are shown in Fig. 226-13.

Perinatal HIV transmission, from an HIV-infected mother to her baby, has declined significantly in the United States, largely due to

the implementation of guidelines for the universal counseling and voluntary HIV testing of pregnant women and the use of antiretroviral therapy for pregnant women and newborn infants to prevent infection (Fig. 226-14). In 2011, fewer than 200 children were diagnosed with HIV infection in the United States.

HIV infection and AIDS have disproportionately affected minority populations in the United States. Among those diagnosed with HIV (regardless of stage of infection) in 2011, 47% percent were Blacks/African Americans, a group that constitutes only 12% of the U.S. population (Fig. 226-15A). The estimated rate of new HIV diagnoses in 2011 by race/ethnicity per 100,000 population in the United States is shown in Fig. 226-15B.

The number of individuals diagnosed with AIDS and deaths among persons with AIDS in the United States rose steadily through the 1980s; AIDS cases peaked in 1993 and deaths in 1995 (Fig. 226-16). Since then, the annual numbers of AIDS-related deaths in the United States have fallen ~70%. This trend is due to several factors, including improved prophylaxis and treatment of opportunistic infections, growing experience among the health professions in caring for HIV-infected individuals, improved access to health care, and a decrease in new infections due to saturational effects and prevention efforts.

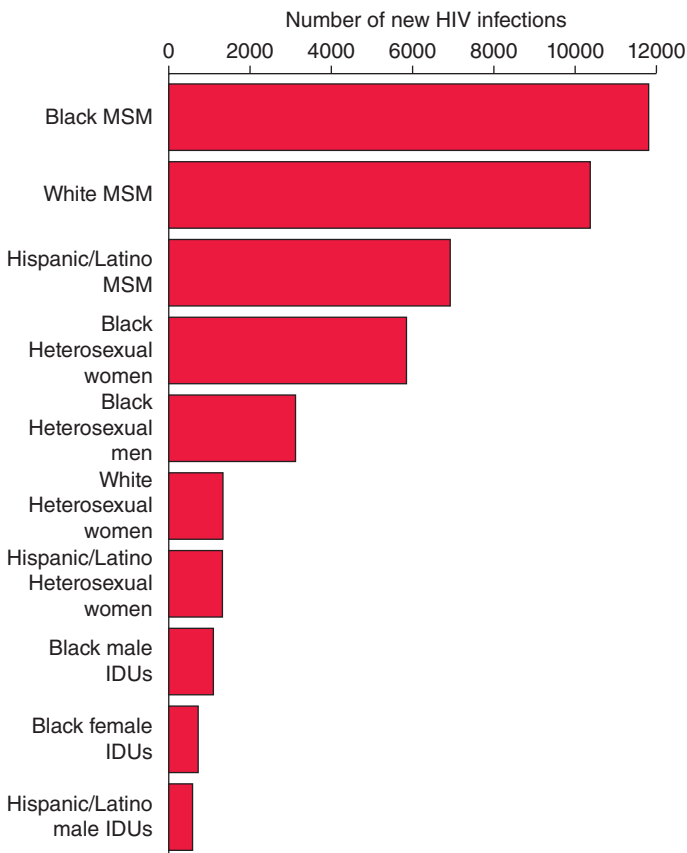


FIGURE 226-13 Estimated number of new HIV infections in the United States, 2011, for the most affected subpopulations. (From CDC.)

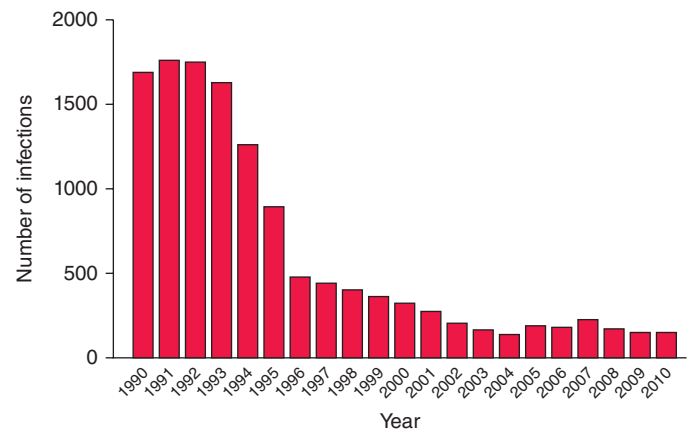


FIGURE 226-14 Estimated number of HIV-infected infants, United States, 1990–2010. (From CDC.)

However, the most influential factor clearly has been the increased use of potent antiretroviral drugs, generally administered in a combination of three or four agents.

Although the HIV/AIDS epidemic on the whole is plateauing in the United States, it is spreading rapidly among certain populations, stabilizing in others, and decreasing in others. Similar to other STIs, HIV infection will not spread homogeneously throughout the population of the United States. However, it is clear that anyone who practices high-risk behavior is at risk for HIV infection. In addition, recent increases in infections and AIDS cases among young men who have sex with men as well as the spread in pockets of poverty in both urban and rural regions (particularly among underserved minority populations in the southern United States with inadequate access to health care) testify that the epidemic of HIV infection in the United States remains a public health problem of major proportion.

PATHOPHYSIOLOGY AND PATHOGENESIS

The hallmark of HIV disease is a profound immunodeficiency resulting primarily from a progressive quantitative and qualitative deficiency of the subset of T lymphocytes referred to as *helper T cells* occurring in a setting of polyclonal immune activation. The *helper* subset of T cells is defined phenotypically by the presence on its surface of the CD4 molecule (Chap. 372e), which serves as the primary cellular receptor for HIV. A co-receptor must also be present together with CD4 for efficient binding, fusion, and entry of HIV-1 into its target cells (Figs. 226-3 and 226-4). HIV uses two major co-receptors, CCR5 and CXCR4, for fusion and entry; these co-receptors are also the primary receptors for certain chemoattractive cytokines termed *chemokines* and belong to the seven-transmembrane-domain G protein-coupled family of receptors. A number of mechanisms responsible for cellular depletion and/or immune dysfunction of CD4+ T cells have been demonstrated in vitro; these include direct infection and destruction of these cells by HIV, as well as indirect effects such as immune clearance of infected cells, cell death associated with aberrant immune activation, and immune exhaustion due to aberrant cellular activation

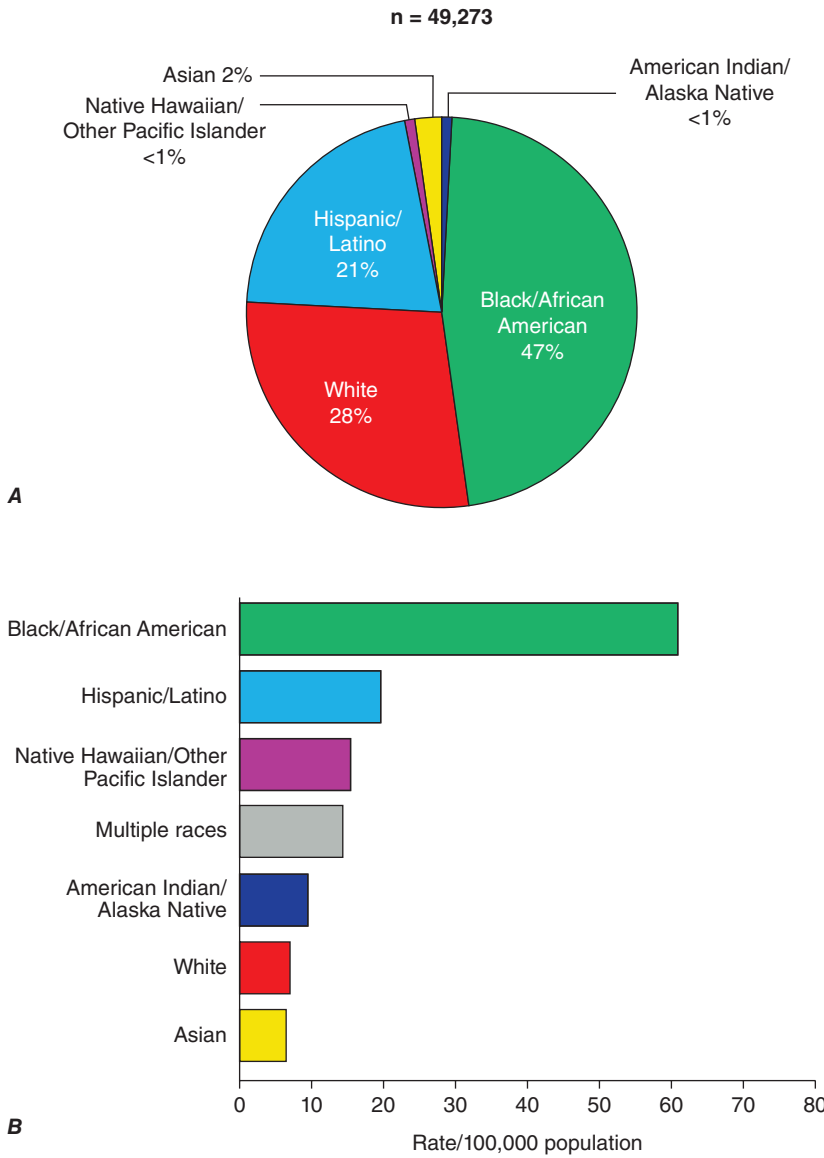


FIGURE 226-15 Race/ethnicity of persons (including children) diagnosed with HIV infection (regardless of stage) during 2011 in the United States. **A.** Estimated proportion of new infections by race/ethnicity. **B.** Estimated rate of new infections by race/ethnicity (per 100,000 population). (From CDC.)

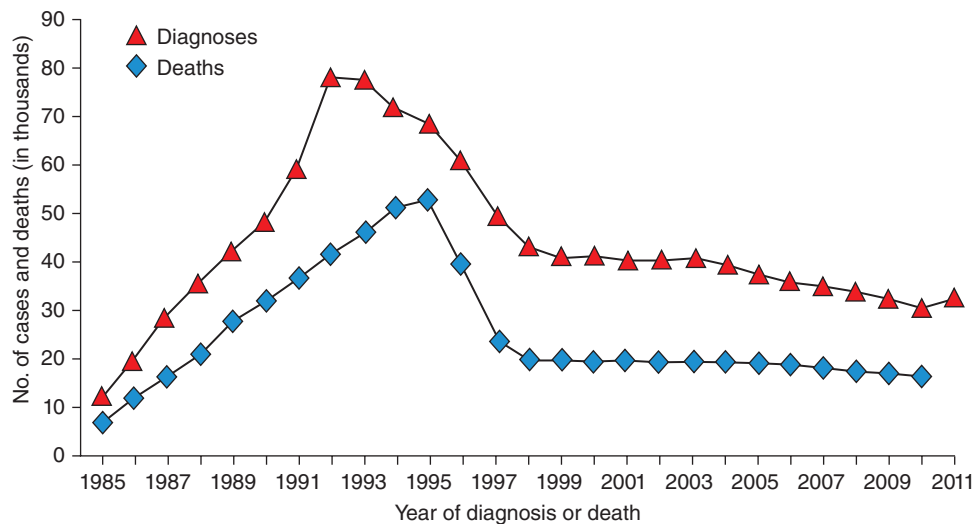


FIGURE 226-16 Estimated number of AIDS cases and AIDS deaths, United States, 1985–2011. (From CDC.)

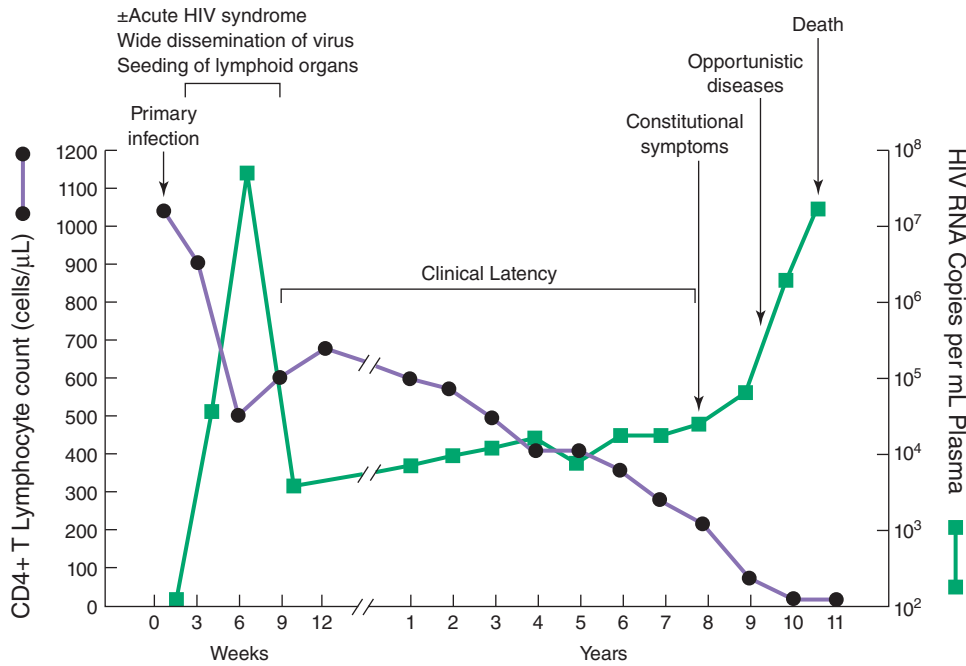


FIGURE 226-17 Typical course of an untreated HIV-infected individual. See text for detailed description. (From G Pantaleo et al: *N Engl J Med* 328:327, 1993. Copyright 1993 Massachusetts Medical Society. All rights reserved.)

with resulting cellular dysfunction. Patients with CD4+ T cell levels below certain thresholds are at high risk of developing a variety of opportunistic diseases, particularly the infections and neoplasms that are AIDS-defining illnesses. Some features of AIDS, such as Kaposi's sarcoma and certain neurologic abnormalities, cannot be explained completely by the immunodeficiency caused by HIV infection, since these complications may occur prior to the development of severe immunologic impairment.

The combination of viral pathogenic and immunopathogenic events that occurs during the course of HIV disease from the moment of initial (primary) infection through the development of advanced-stage disease is complex and varied. It is important to appreciate that the pathogenic mechanisms of HIV disease are multifactorial and multiphasic and are different at different stages of the disease. Therefore, it is essential to consider the typical clinical course of an untreated HIV-infected individual in order to more fully appreciate these pathogenic events (Fig. 226-17).

EARLY EVENTS IN HIV INFECTION: PRIMARY INFECTION AND INITIAL DISSEMINATION OF VIRUS

Using mucosal transmission as a model, the earliest events (within hours) that occur following exposure of HIV to the mucosal surface determine whether an infection will be established as well as the subsequent course of events following infection. Although the mucosal barrier is relatively effective in limiting access of HIV to susceptible targets in the lamina propria, the virus can cross the barrier by transport on Langerhans cells, an epidermal type of DC, just beneath the surface or through microscopic rents in the mucosa. Significant disruptions in the mucosal barrier as seen in ulcerative genital disease facilitate viral entry and increase the efficiency of infection. Virus then seeks susceptible targets, which are primarily CD4+ T cells that are spatially dispersed in the mucosa. This spatial dispersion of targets provides a significant obstacle to the establishment of infection. Such obstacles account for the low efficiency of sexual transmission of HIV (see "Sexual Transmission," above). Both "partially" resting CD4+ T cells and activated CD4+ T cells serve as early amplifiers of infection. Resting CD4+ T cells are more abundant; however, activated CD4+ T cells produce larger amounts of virus. In order for infection to become established, the basic reproductive rate (R_0) must become equal to or greater than 1, i.e., each infected cell would infect at least

one other cell. Once infection is established, the virus replicates in lymphoid cells in the mucosa, the submucosa, and to some extent the lymphoreticular tissues that drain the gut tissues. For a variable period of time ranging from a few to several days, the virus cannot yet be detected in the plasma. This period is referred to as the "eclipse" phase of infection. As more virus is produced within several days to weeks, it is disseminated, first to the draining lymph nodes and then to other lymphoid compartments where it has easy access to dense concentrations of CD4+ T cell targets, allowing for a burst of high-level viremia that is readily detectable by currently available assays (Fig. 226-18). An important lymphoid organ, the gut-associated lymphoid tissue (GALT), is a major target of HIV infection and the location where large numbers of CD4+ T cells (usually memory cells) are infected and depleted, both by direct viral effects and by activation-associated apoptosis. Once virus replication reaches this threshold and virus is widely disseminated, infection is firmly established and the process is irreversible. It is important to point out

that the initial infection of susceptible cells may vary somewhat with the route of infection. Virus that enters directly into the bloodstream via infected blood or blood products (i.e., transfusions, use of contaminated needles for injection drugs, sharp-object injuries, maternal-to-fetal transmission either intrapartum or perinatally, or sexual intercourse where there is enough trauma to cause bleeding) is likely first cleared from the circulation to the spleen and other lymphoid organs, where primary focal infections begin, followed by wider dissemination throughout other lymphoid tissues as described above.

It has been demonstrated that sexual transmission of HIV is the result of a single infectious event and that a viral genetic bottleneck exists for transmission. In this regard, certain characteristics of the HIV envelope glycoprotein have a major influence on transmission, at least in subtype A and C viruses. Transmitting viruses, often referred to as "founder viruses," are usually underrepresented in the circulating viremia of the transmitting partner and are less-diverged viruses with signature sequences including shorter V1-V2 loop sequences and fewer predicted N-linked glycosylation sites relative to the major circulating variants. These viruses are almost exclusively R5 strains and are usually sensitive to neutralization by antibody from the transmitting partner. Once replication proceeds in the newly infected partner, the founder virus diverges and accumulates glycosylation sites, becoming progressively more resistant to neutralization (Fig. 226-19).

The acute burst of viremia and wide dissemination of virus in primary HIV infection may be associated with an *acute HIV syndrome*, which occurs to varying degrees in ~50% of individuals with primary infection (see below). This syndrome is usually associated with high levels of viremia measured in millions of copies of HIV RNA per milliliter of plasma that last for several weeks. Acute mononucleosis-like symptoms are well correlated with the presence of viremia. Virtually all patients develop some degree of viremia during primary infection, which contributes to virus dissemination throughout the lymphoid tissue, even though they may remain asymptomatic or not recall experiencing symptoms. It appears that the initial level of plasma viremia in primary HIV infection does not necessarily determine the rate of disease progression; however, the set point of the level of steady-state plasma viremia after ~1 year does seem to correlate with the slope of disease progression in the untreated patient. The strikingly high levels of viremia observed in many patients with acute HIV infection is felt to be associated with a higher likelihood of transmission of the virus

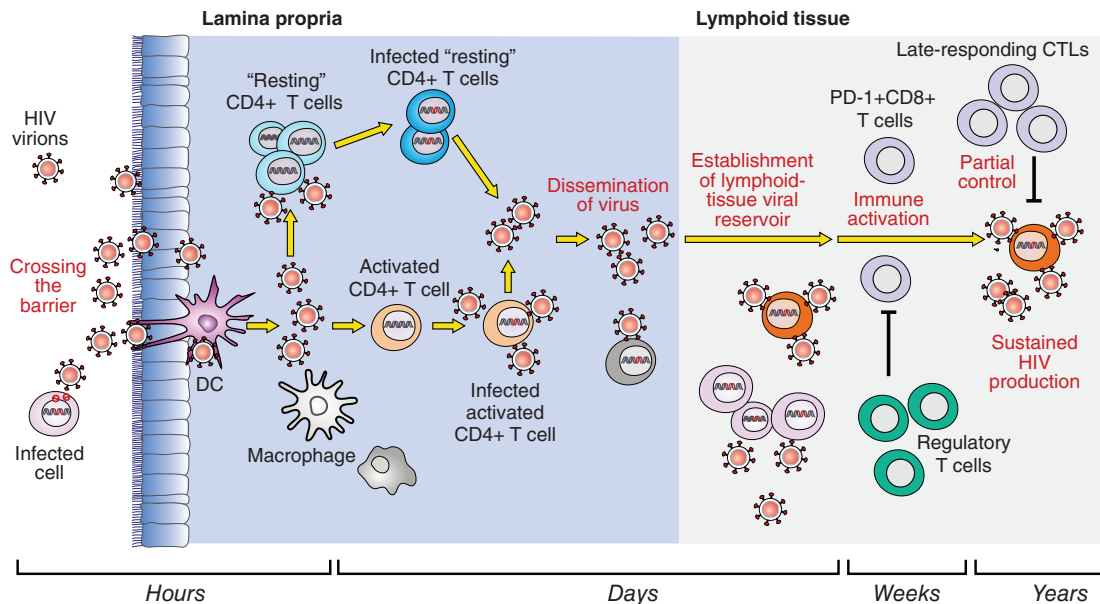


FIGURE 226-18 Summary of early events in HIV infection. See text for detailed description. CTLs, cytolytic T lymphocytes; HIV, human immunodeficiency virus. (Adapted from AT Haase: *Nat Rev Immunol* 5:783, 2005.)

to others by a variety of routes including sexual transmission, shared needles and syringes, and mother-to-child transmission intrapartum, perinatally, or via breast milk.

ESTABLISHMENT OF CHRONIC AND PERSISTENT INFECTION

Persistence of Virus Replication HIV infection is unique among human viral infections. Despite the robust cellular and humoral immune responses that are mounted following primary infection (see “Immune Response to HIV,” below), once infection has been established the virus succeeds in escaping complete immune-mediated clearance, paradoxically seems to thrive on immune activation, and is never eliminated completely from the body. Rather, a chronic infection develops and persists with varying degrees of continual virus replication in the untreated patient for a median of ~10 years before the patient becomes clinically ill (see “Advanced HIV Disease,” below). It is this establishment of a chronic, persistent infection that is the hallmark of HIV disease. Throughout the often protracted course of chronic infection, virus replication can invariably be detected in untreated patients by widely available assays that measure copies of HIV RNA per milliliter of plasma. Levels of virus vary greatly in most untreated patients, ranging from several thousand to a few million copies of HIV RNA per milliliter of plasma. Studies using highly sensitive molecular techniques have demonstrated that even in certain

patients in whom plasma viremia is suppressed to below detection (lower limit, 20–50 copies of HIV RNA/mL depending on manufacturer) by cART, there is a continual low level of virus replication. In other human viral infections, with very few exceptions, if the host survives, the virus is completely cleared from the body and a state of immunity against subsequent infection develops. HIV infection very rarely kills the host during primary infection. Certain viruses, such as HSV (Chap. 216), are not completely cleared from the body after infection, but instead enter a latent state; in these cases, clinical latency is accompanied by microbiologic latency. This is not the case with HIV infection as described above. Chronicity associated with persistent virus replication can also be seen in certain cases of HBV and HCV infections (Chap. 362); however, in these infections the immune system is not a target of the virus.

Escape of HIV from Effective Immune System Control Inherent to the establishment of chronicity of HIV infection is the ability of the virus to evade adequate control and elimination by both the cellular and humoral limbs of the immune system. There are a number of mechanisms whereby the virus accomplishes this evasion. Paramount among these is the establishment of a sustained level of replication associated with the generation of viral diversity via mutation and recombination. The selection of mutants that escape control by CD8⁺ cytolytic T lymphocytes (CTLs) is critical to the propagation and progression of HIV infection. The high rate of virus replication associated with inevitable mutations also contributes to the inability of antibody to neutralize the autologous virus and thus to contain the virus quasispecies present in an individual at any given time. Extensive analyses of sequential HIV isolates and host responses have demonstrated that viral escape from B cell and CD8⁺ T cell epitopes occurs early after infection and allows the virus to stay one step ahead of effective immune responses. Virus-specific CD8⁺ CTLs expand greatly during primary HIV infection, and likely represent the high-affinity responses that would be expected to be most efficient in eliminating virus-infected cells; however, the restriction is generally incomplete as viral replication persists at relatively high levels in the majority of individuals. In addition to viral escape from CTLs through high rates of mutation, it is thought that the initially strong response becomes qualitatively dysfunctional owing to the overwhelming immune activation resulting from persistent viral replication, similar to the exhaustion of CD8⁺ CTLs that has been reported in the murine model of lymphocytic choriomeningitis virus (LCMV) infection. Several studies have indicated that exhaustion of HIV-specific CD8⁺ T cells during prolonged immune activation is associated with expression of inhibitory receptors, such

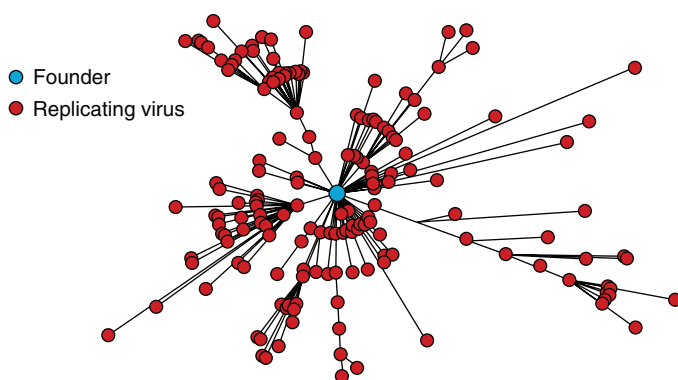


FIGURE 226-19 As HIV diverges from founder to chronically replicating virus, it accumulates N-linked glycosylation sites. See text for detailed description. (Adapted from CA Derdeyn et al: *Science* 303:2019, 2004; B Chohan et al: *J Virol* 79:6528, 2005; and BF Keele et al: *Proc Natl Acad Sci USA* 105:7552, 2008.)

1230 as programmed death (PD) 1 molecule (of the B7-CD28 family of molecules), as well as loss of polyreactivity and proliferative capacity. Another mechanism contributing to the evasion by HIV of immune system control is the downregulation of HLA class I molecules on the surface of HIV-infected cells by the viral proteins Nef, Tat, and Vpu, resulting in the lack of ability of the CD8+ CTL to recognize and kill the infected target cell. Although this downregulation of HLA class I molecules would favor elimination of HIV-infected cells by natural killer (NK) cells, this latter mechanism does not seem to remove HIV-infected cells effectively (see below).

The principal targets of neutralizing antibodies against HIV are the envelope proteins gp120 and gp41. HIV employs at least three mechanisms to evade neutralizing responses: hypervariability in the primary sequence of the envelope, extensive glycosylation of the envelope, and conformational masking of neutralizing epitopes. Several studies that have followed the humoral immune response to HIV from the earliest points after primary infection indicate that the virus continually mutates to escape the emerging antibody response such that the sequential antibodies that are induced do not neutralize the autologous virus. *Broadly neutralizing antibodies* capable of neutralizing a wide range of primary HIV isolates in vitro occur in only about 20% of HIV-infected individuals, and, when they do occur, it generally requires 2 to 3 years of infection with continual virus replication to drive the affinity maturation of the antibodies. Unfortunately, by the time these broadly neutralizing antibodies are formed, they are ineffective in containing the virus replication in the patient. Persistent viremia also results in exhaustion of B cells similar to the exhaustion reported for CD4+ T cells, adding to the defects in the humoral response to HIV.

CD4+ T cell help is essential for the integrity of antigen-specific immune responses, both humoral and cell-mediated. HIV preferentially infects activated CD4+ T cells including HIV-specific CD4+ T cells, and so this loss of viral-specific helper T cell responses has profound negative consequences for the immunologic control of HIV replication. Furthermore, this loss occurs early in the course of infection, and animal studies indicate that 40–70% of all memory CD4+ T cells in the GALT are eliminated during acute infection. Another potential means of escape of HIV-infected cells from elimination by CD8+ CTLs is the sequestration of infected cells in immunologically privileged sites such as the central nervous system (CNS).

Finally, the escape of HIV from immune-mediated elimination during primary infection allows the formation of a pool of latently infected cells that may not be recognized or completely eliminated by virus-specific CTLs or by ART (see below). Thus, despite a potent immune response and the marked downregulation of virus replication following primary HIV infection, HIV succeeds in establishing a state of chronic infection with a variable degree of persistent virus replication. During this period most patients make the clinical transition from acute primary infection to variable periods of clinical latency or smoldering disease activity (see below).

The HIV Reservoir: Obstacles to the Eradication of Virus A pool of latently infected, resting CD4+ T cells that serves as at least one component of the persistent reservoir of virus exists in virtually all HIV-infected individuals, including those who are receiving cART. Such cells carry an integrated form of HIV DNA in the genome of the host and can remain in this state until an activation signal drives the expression of HIV transcripts and ultimately replication-competent virus. This form of latency is to be distinguished from preintegration latency, in which HIV enters a resting CD4+ T cell and, in the absence of an activation signal, reverse transcription of the HIV genome occurs to a certain extent but the resulting proviral DNA fails to integrate into the host genome. This period of preintegration latency may last hours to days, and if no activation signal is delivered to the cell, the proviral DNA loses its capacity to initiate a productive infection. If these cells do become activated prior to decay of the preintegration complex, reverse transcription proceeds to completion and the virus continues along its replication cycle (see above and Fig. 226-20). The pool of cells that are in the postintegration state of latency is established early during the

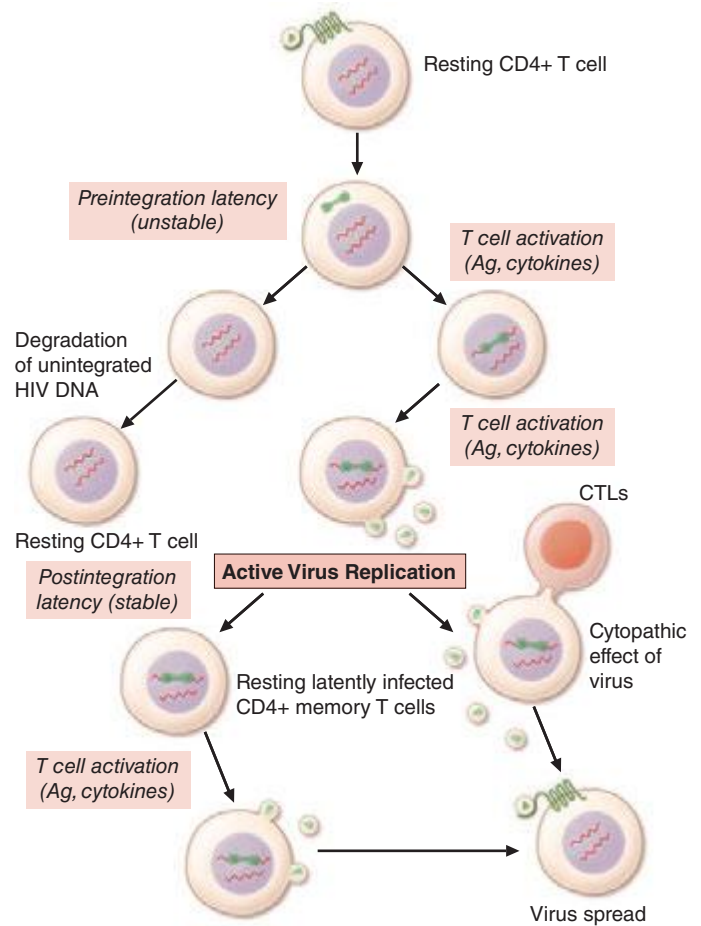


FIGURE 226-20 Generation of latently infected, resting CD4+ T cells in HIV-infected individuals. See text for details. Ag, antigen; CTLs, cytolytic T lymphocytes. (Courtesy of TW Chun; with permission.)

course of primary HIV infection. Despite the suppression of plasma viremia to <50 copies of HIV RNA per milliliter by potent regimens of cART administered over several years, this pool of latently infected cells persists and can give rise to replication-competent virus upon cellular activation. Modeling studies built on projections of decay curves have estimated that in such a setting of prolonged suppression, it would require a few to several years for the pool of latently infected cells to be completely eliminated. This has not been documented to occur spontaneously in any patients very likely because the latent viral reservoir is continually replenished by the low levels of persistent virus replication that may remain below the limits of detection of current assays (see below) (Fig. 226-20), even in patients who for the most part are treated successfully. Reservoirs of HIV-infected cells, latent or otherwise, can exist in a number of compartments including the lymphoid tissue, peripheral blood, and the CNS (likely in cells of the monocyte/macrophage lineage) as well as in other unidentified locations. Over the past several years attempts have been made to eliminate HIV in the latent viral reservoir using agents that stimulate resting CD4+ T cells during the course of cART; however, such attempts have been unsuccessful. Thus, this persistent reservoir of infected cells and/or low levels of persistent virus replication are major obstacles to the goal of eradication of virus from infected individuals and hence a “cure,” despite the favorable clinical outcomes that have resulted from cART.

Viral Dynamics The dynamics of viral production and turnover have been quantified using mathematical modeling in the setting of the administration of reverse transcriptase and protease inhibitors to HIV-infected individuals in clinical studies. Treatment with these drugs resulted in a precipitous decline in the level of plasma viremia, which typically fell by well over 90% within 2 weeks. The number of CD4+ T cells in the blood increased concurrently, which suggested that the

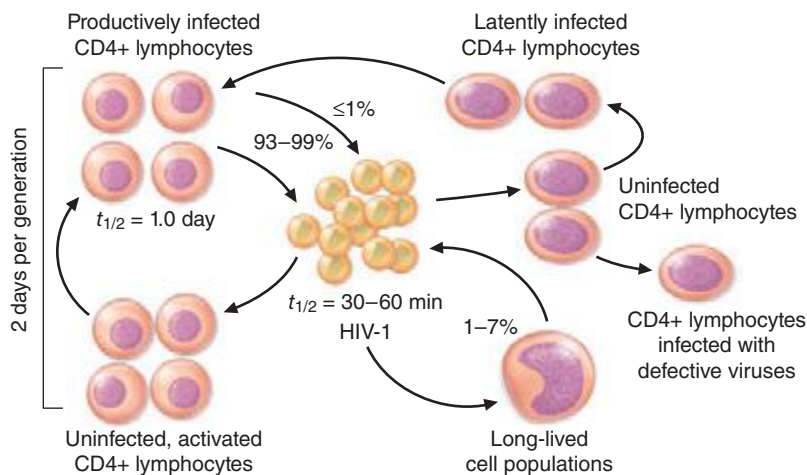


FIGURE 226-21 Dynamics of HIV infection in vivo. See text for detailed description. (From AS Perelson et al: *Science* 271:1582, 1996.)

killing of CD4+ T cells was linked directly to the levels of replicating virus. However, a significant component of the early rise in CD4+ T cell numbers following the initiation of therapy may be due to the redistribution of cells into the peripheral blood from other tissue compartments throughout the body as a consequence of therapy-related diminution in viremia-associated immune system activation. It was determined on the basis of modeling the kinetics of viral decline and the emergence of resistant mutants during therapy that 93–99% of the circulating virus originated from recently infected, rapidly turning over CD4+ T cells and that ~1–7% of circulating virus originated from longer-lived cells, likely monocytes/macrophages. A negligible amount of circulating virus originated from the pool of latently infected cells (Fig. 226-21). It was also determined that the half-life of a circulating virion was ~30–60 min and that of productively infected cells was 1 day. Given the relatively steady level of plasma viremia and of infected cells, it appears that extremely large amounts of virus (~ 10^{10} – 10^{11} virions) are produced and cleared from the circulation each day. In addition, data suggest that the minimal duration of the HIV-1 replication cycle in vivo is ~2 days. Other studies have demonstrated that the decrease in plasma viremia that results from cART correlates closely with a decrease in virus replication in lymph nodes, further confirming that lymphoid tissue is the main site of HIV replication and the main source of plasma viremia.

The level of steady-state viremia, called the viral *set point*, at ~1 year following acquisition of HIV infection has important prognostic implications for the progression of HIV disease in the untreated patient. It has been demonstrated that as a group untreated HIV-infected individuals who have a low set point at 6 months to 1 year following infection progress to AIDS much more slowly than individuals whose set point is very high at that time (Fig. 226-22).

Clinical Latency versus Microbiologic Latency With the exception of long-term nonprogressors (see “Long-Term Survivors and Long-Term Nonprogressors,” below), the level of CD4+ T cells in the blood decreases progressively in HIV-infected individuals in the absence of cART. The decline in CD4+ T cells may be gradual or abrupt, the latter usually reflecting a significant spike in the level of plasma viremia. Most patients are relatively asymptomatic while this progressive decline is taking place (see below) and are often described as being in a state of *clinical latency*. However, this term is misleading; it does not mean disease latency, since progression, although slow in many cases, is generally relentless during this period. Furthermore, clinical latency should not be confused with microbiologic latency, since varying levels of virus replication inevitably occur during this period of clinical latency. Even in those rare patients who have <50 copies of HIV RNA

per milliliter in the absence of therapy, there is virtually always some degree of ongoing virus replication.

ADVANCED HIV DISEASE

In untreated patients or in patients in whom therapy has not adequately controlled virus replication, after a variable period, usually measured in years, the CD4+ T cell count falls below a critical level (<200/ μ L) and the patient becomes highly susceptible to opportunistic disease (Fig. 226-17). For this reason, the CDC case definition of AIDS includes all HIV-infected individuals over 5 years of age with CD4+ T cell counts below this level (Table 226-2). Patients may experience constitutional signs and symptoms or may develop an opportunistic disease abruptly without any prior symptoms, although the latter scenario is unusual. The depletion of CD4+ T cells continues to be progressive and unrelenting in this phase. It is not uncommon for CD4+ T cell counts in the untreated patient to drop to as low as 10/ μ L or even to zero. In countries where cART and prophylaxis and treatment for opportunistic infections are readily accessible to such patients, survival is increased dramatically even in those patients with advanced HIV disease. In contrast, untreated patients who progress to this severest form of immunodeficiency usually succumb to opportunistic infections or neoplasms (see below).

LONG-TERM SURVIVORS AND LONG-TERM NONPROGRESSORS

It is important to distinguish between the terms *long-term survivor* and *long-term nonprogressor*. Long-term nonprogressors are by definition long-term survivors; however, the reverse is not always true. Predictions from one study that antedated the availability of effective cART estimated that ~13% of homosexual/bisexual men who were infected at an early age may remain free of clinical AIDS for >20 years. Many of these individuals may have progressed in their degree of immune deficiency; however, they certainly survived for a considerable period of time. With the advent of effective cART, the survival of HIV-infected individuals has dramatically increased. Early in the AIDS epidemic, prior to the availability of therapy, if a patient presented with a life-threatening opportunistic infection, the median survival was 26 weeks from the time of presentation. Currently, an HIV-infected 20-year-old individual in a high-income country who is appropriately treated with cART can expect to live at least 50 years according to mathematical model projections. In the face of cART, long-term survival is becoming commonplace. Definitions of

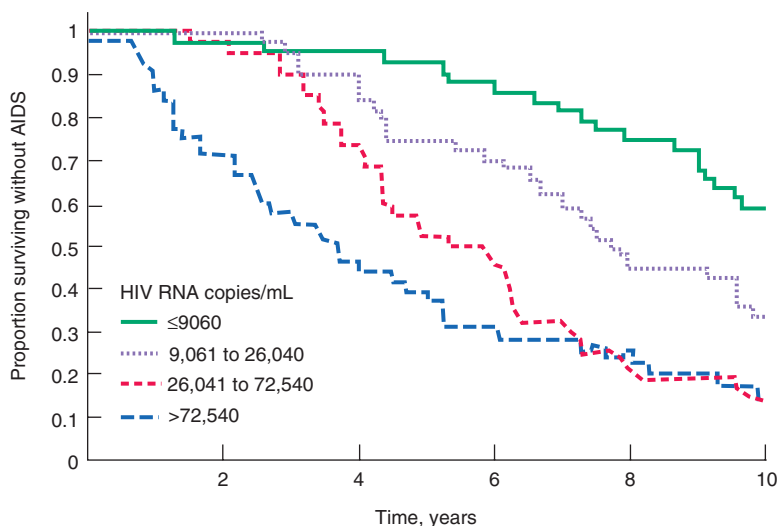


FIGURE 226-22 Relationship between levels of virus and rates of disease progression. Kaplan-Meier curves for AIDS-free survival stratified by baseline HIV-1 RNA categories (copies per milliliter). (From JW Mellors et al: *Science* 272:1167, 1996.)

1232 long-term nonprogressors have varied considerably over the years, and so such individuals constitute a heterogeneous group. Long-term nonprogressors were first described in the 1990s. Originally, individuals were considered to be long-term nonprogressors if they had been infected with HIV for a long period (≥ 10 years), their CD4+ T cell counts were in the normal range, and they remained stable over years without receiving cART. Approximately 5–15% of HIV-infected individuals fell into this broader nonprogressor category. However, this group was rather heterogeneous and over time a significant proportion of these individuals progressed and ultimately required therapy. From this broader group, a much smaller subgroup of “elite” controllers or nonprogressors was identified, and they constituted less than 1% of HIV-infected individuals. These elite controllers, by definition, have extremely low levels of plasma viremia and normal CD4+ T cell counts. It is noteworthy that certain of their HIV-specific immune responses are robust and clearly superior to those of HIV-infected progressors. In this group of elite controllers certain HLA class I haplotypes are overrepresented, particularly HLA-B57-01 and HLA-B27-05. Outside of the subgroup of elite controllers, a number of other genetic factors have been shown to be involved to a greater or lesser degree in the control of virus replication and thus in the rate of HIV disease progression (see “Genetic Factors in HIV-1 and AIDS Pathogenesis,” below).

LYMPHOID ORGANS AND HIV PATHOGENESIS

Regardless of the portal of entry of HIV, lymphoid tissues are the major anatomic sites for the establishment and propagation of HIV infection. Despite the use of measurements of plasma viremia to determine the level of disease activity, virus replication occurs mainly in lymphoid tissue and not in blood; indeed, the level of plasma viremia directly reflects virus production in lymphoid tissue.

Some patients experience progressive generalized lymphadenopathy early in the course of the infection; others experience varying degrees of transient lymphadenopathy. Lymphadenopathy reflects the cellular activation and immune response to the virus in the lymphoid tissue, which is generally characterized by follicular or germinal center hyperplasia. Lymphoid tissue involvement is a common denominator of virtually all patients with HIV infection, even those without easily detectable lymphadenopathy.

Simultaneous examinations of lymph tissue and peripheral blood in patients and monkeys during various stages of HIV and SIV infection, respectively, have led to substantial insight into the pathogenesis of HIV disease. In most of the original human studies, peripheral lymph nodes have been used predominantly as the source of lymphoid tissue. More recent studies in monkeys and humans have also focused on the GALT, where the earliest burst of virus replication occurs associated with marked depletion of CD4+ T cells. In detailed studies of peripheral lymph node tissue, using a combination of polymerase chain reaction (PCR) techniques for HIV DNA and HIV RNA in tissue and HIV RNA in plasma, in situ hybridization for HIV RNA, and light and electron microscopy, the following picture has emerged. During acute HIV infection resulting from mucosal transmission, virus replication progressively amplifies from scattered lymphoid cells in the lamina propria to draining lymphoid tissue, leading to high levels of plasma viremia. The GALT plays a major role in the amplification of virus replication, and virus is disseminated from replication in the GALT to peripheral lymphoid tissue. A profound degree of cellular activation occurs (see below) and is reflected in follicular or germinal center hyperplasia. At this time copious amounts of extracellular virions (both infectious and defective) are trapped on the processes of the follicular dendritic cells (FDCs) in the germinal centers of the lymph nodes. Virions that have bound complement components on their surfaces attach to the surface of FDCs via interactions with complement receptors and likely via Fc receptors that bind to antibodies that are attached to the virions. In situ hybridization reveals expression of virus in individual cells of the paracortical area and, to a lesser extent, the germinal center (Fig. 226-23). The persistence of trapped virus after the transition from acute to chronic infection likely reflects a steady state whereby trapped virus turns over and is replaced by fresh

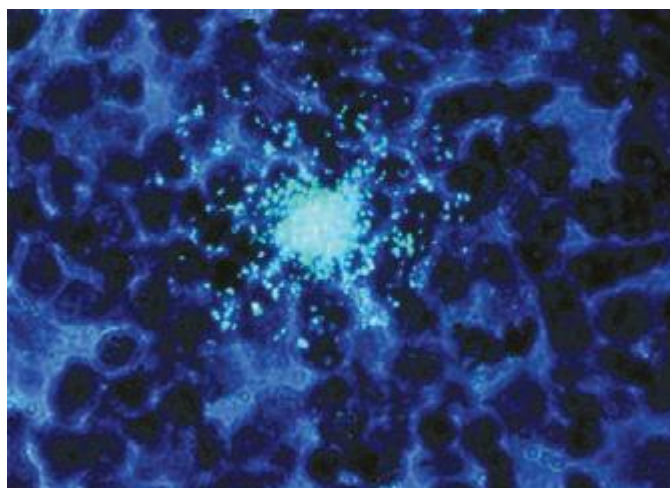


FIGURE 226-23 HIV in the lymph node of an HIV-infected individual. An individual cell infected with HIV shown expressing HIV RNA by in situ hybridization using a radiolabeled molecular probe. Original $\times 500$. (Adapted from G Pantaleo, AS Fauci et al: *Nature* 362:355, 1993.)

virions that are continually produced. The trapped virus, either as whole virion or shed envelope, serves as a continual activator of CD4+ T cells, thus driving further virus replication.

During early and chronic/asymptomatic stages of HIV disease, the architecture of lymphoid tissues is generally preserved and may even be hyperplastic owing to an increased presence of B cells and specialized CD4+ T cells called follicular helper CD4+ T cells (TF_H) in prominent germinal centers. Extracellular virions can be seen by electron microscopy attached to FDC processes. The trapping of antigen is a physiologically normal function for the FDCs, which present antigen to B cells and, along with the action of TF_H cells, contribute to the generation of B cell memory. However, in the case of HIV, persistent cellular activation, resulting in the secretion of proinflammatory cytokines such as interleukin (IL) 1β , tumor necrosis factor (TNF) α , IFN- γ , and IL-6, which can induce viral replication (see below) and diminish the effectiveness of the immune response against the virus. In addition, the CD4+ TF_H cells that are recruited into the germinal center to provide help to B cells in the generation of an HIV-specific immune response are highly susceptible to infection by virus either trapped on FDC or propagated locally. Thus, in HIV infection, a normal physiologic function of the immune system that contributes to the clearance of virus, as well as to the generation of a specific immune response, can also have deleterious consequences.

As the disease progresses, the architecture of lymphoid tissues begins to show disruption. Confocal microscopy reveals destruction of the fibroblastic reticular cell (FRC) and FDC networks in the T cell zone and B cell follicles, respectively, and an incapacity to replenish naïve cells. The mechanisms of destruction are not completely understood, but they are thought to be associated with collagen deposition causing fibrosis and loss of production of cytokines such as IL-7 and lymphotoxin- α , which are critical to the maintenance of lymphoid tissues and their lymphocyte constituents. As the disease progresses to an advanced stage, there is complete disruption of the architecture of the lymphoid tissues, accompanied by dissolution of the FRC and FDC networks. At this point, the lymph nodes are “burnt out.” This destruction of lymphoid tissue compounds the immunodeficiency of HIV disease and contributes both to the inability to control HIV replication (leading usually to high levels of plasma viremia in the untreated or inadequately treated patient) and to the inability to mount adequate immune responses against opportunistic pathogens. The events from primary infection to the ultimate destruction of the immune system are illustrated in Fig. 226-24. Recently, nonhuman primate studies and some human studies have examined GALT at various stages of HIV disease. Within the GALT, the basal level of activation combined with virus-mediated cellular activation results in the infection and elimination of an estimated 50–90% of CD4+ T cells in the gut. The extent of

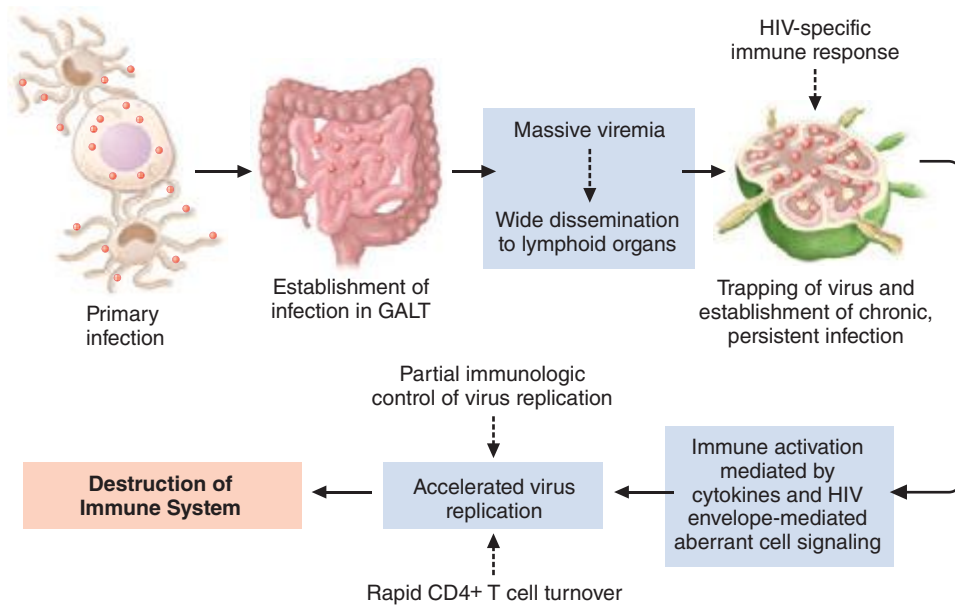


FIGURE 226-24 Events that transpire from primary HIV infection through the establishment of chronic persistent infection to the ultimate destruction of the immune system. See text for details. CTLs, cytolytic T lymphocytes; GALT, gut-associated lymphoid tissue.

this early damage to GALT, which constitutes a major component of lymphoid tissue in the body, may play a role in determining the potential for immunologic recovery of the memory cell subset.

IMMUNE ACTIVATION, INFLAMMATION, AND HIV PATHOGENESIS

Activation of the immune system and variable degrees of inflammation are essential components of any appropriate immune response to a foreign antigen. However, immune activation and inflammation, which can be considered aberrant in HIV-infected individuals, play a critical role in the pathogenesis of HIV disease and other chronic conditions associated with HIV disease. Immune activation and inflammation in the HIV-infected individual contribute substantially to (1) the replication of HIV, (2) the induction of immune dysfunction, and (3) the increased incidence of chronic conditions associated with persistent immune activation and inflammation (Table 226-4).

INDUCTION OF HIV REPLICATION BY ABERRANT IMMUNE ACTIVATION The immune system is normally in a state of homeostasis, awaiting perturbation by foreign antigenic stimuli. Once the immune response deals with and clears the antigen, the system returns to relative quiescence (Chap. 372e). This is generally not the case in HIV infection where, in the untreated patient, virus replication is invariably persistent with very few exceptions and immune activation is persistent. HIV replicates most efficiently in activated CD4+ T cells; in HIV infection, chronic activation provides the cell substrates necessary for persistent virus replication throughout the course of HIV disease, particularly in the untreated patient and to variable degrees even in certain patients receiving cART whose levels of plasma

viremia are suppressed to below the level of detection by standard assays. From a virologic standpoint, although quiescent CD4+ T cells can be infected with HIV, reverse transcription, integration, and virus spread are much more efficient in activated cells. Furthermore, cellular activation induces expression of virus in cells latently infected with HIV. In essence, immune activation and inflammation provide the engine that drives HIV replication. In addition to endogenous factors such as cytokines, a number of exogenous factors such as other microbes that are associated with heightened cellular activation can enhance HIV replication and thus may have important effects on HIV pathogenesis. Co-infection in vivo or in vitro with a range of viruses, such as HSV types 1 and 2, cytomegalovirus (CMV), human herpesvirus (HHV) 6, Epstein-Barr virus (EBV), HBV, adenovirus, and HTLV-1 have been shown to upregulate HIV expression. In addition, infestation with nematodes has been shown to be associated with a heightened state of immune

activation that facilitates HIV replication; in certain studies deworming of the infected host has resulted in a decrease in plasma viremia. Two diseases of extraordinary global health significance, malaria and tuberculosis (TB), have been shown to increase HIV viral load in dually infected individuals. Globally, *Mycobacterium tuberculosis* is the most common opportunistic infection in HIV-infected individuals (Chap. 202). In addition to the fact that HIV-infected individuals are more likely to develop active TB after exposure, it has been demonstrated that active TB can accelerate the course of HIV infection. It has also been shown that levels of plasma viremia are greatly elevated in HIV-infected individuals with active TB who are not on cART, compared with pre-TB levels and levels of viremia after successful treatment of the active TB. The situation is similar in the interaction between HIV and malaria parasites (Chap. 248). Acute infection of HIV-infected individuals with *Plasmodium falciparum* increases HIV viral load, and the increased viral load is reversed by effective malaria treatment.

MICROBIAL TRANSLOCATION AND PERSISTENT IMMUNE ACTIVATION One proposed mechanism of persistent immune activation involves the disruption of the mucosal barrier in the gut due to HIV replication in and disruption of submucosal lymphoid tissue. As a result of this disruption, there is an increase in the products, particularly lipopolysaccharide (LPS), of bacteria that translocate from the bowel lumen through the damaged mucosa to the circulation, leading to persistent systemic immune activation and inflammation. This effect can persist even after the HIV viral load is brought to <50 copies/mL by cART. Depletion in the GALT of IL-17–producing T cells, which are responsible for defense against extracellular bacteria and fungi, also is thought to contribute to HIV pathogenesis.

PERSISTENT IMMUNE ACTIVATION AND INFLAMMATION INDUCE IMMUNE DYSFUNCTION The activated state in HIV infection is reflected by hyperactivation of B cells leading to hypergammaglobulinemia; increased lymphocyte turnover; activation of monocytes; expression of activation markers on CD4+ and CD8+ T cells; increased activation-associated cellular apoptosis; lymph node hyperplasia, particularly early in the course of disease; increased secretion of proinflammatory cytokines, particularly IL-6; elevated levels of high-sensitivity C-reactive protein, fibrinogen, D-dimer, neopterin, β_2 -microglobulin, acid-labile interferon, soluble (s) IL-2 receptors (R), sTNFR, sCD27, and sCD40L; and autoimmune phenomena (see “Autoimmune Phenomena,” below). Even in the absence of direct infection of a target cell, HIV envelope proteins can interact

TABLE 226-4 CONDITIONS ASSOCIATED WITH PERSISTENT IMMUNE ACTIVATION AND INFLAMMATION IN PATIENTS WITH HIV INFECTION

- Accelerated aging syndrome
- Bone fragility
- Cancers
- Cardiovascular disease
- Diabetes
- Kidney disease
- Liver disease
- Neurocognitive dysfunction

1234 with cellular receptors (CD4 molecules and chemokine receptors) to deliver potent activation signals resulting in calcium flux, the phosphorylation of certain proteins involved in signal transduction, colocalization of cytoplasmic proteins including those involved in cell trafficking, immune dysfunction, and, under certain circumstances, apoptosis. From an immunologic standpoint, chronic exposure of the immune system to a particular antigen over an extended period may ultimately lead to an inability to sustain an adequate immune response to the antigen in question. In many chronic viral infections, including HIV infection, persistent viremia is associated with “functional exhaustion” of virus-specific T cells, decreasing their capacity to proliferate and perform effector functions. It has been demonstrated that this phenomenon may be mediated, at least in part, by the upregulation of inhibitory receptors on HIV-specific T cells, such as PD-1 shared by both CD4+ and CD8+ T cells, as well as CTLA-4 on CD4+ and Tim-3, 2B4, and CD106 on CD8+ T cells. Furthermore, the ability of the immune system to respond to a broad spectrum of antigens may be compromised if immunocompetent cells are maintained in a state of chronic activation.

The deleterious effects of chronic immune activation on the progression of HIV disease are well established. As in most conditions of persistent antigen exposure, the host must maintain sufficient activation of antigen (HIV)-specific responses but must also prevent excessive activation and potential immune-mediated damage to tissues. Certain studies suggest that normal immunosuppressive mechanisms that act to keep hyperimmune activation in check, particularly CD4+, FoxP3+, CD25+ regulatory T cells (T-regs), may be dysfunctional or depleted in the context of advanced HIV disease.

MEDICAL CONDITIONS ASSOCIATED WITH PERSISTENT IMMUNE ACTIVATION AND INFLAMMATION IN HIV DISEASE It has become clear, as the survival of HIV-infected individuals has increased, that a number of previously unrecognized medical complications are associated with HIV disease—and that these complications relate to chronic immune activation and inflammation (Table 226-4). These complications can appear even after patients have experienced years of adequate control of viral replication (plasma viremia below detectable levels) for several years. Of particular note are endothelial cell dysfunction and its relationship to cardiovascular disease. Other chronic conditions that have been reported include bone fragility, certain cancers, persistent immune dysfunction, diabetes, kidney and liver disease, and neurocognitive dysfunction, thus presenting an overall picture of accelerated aging.

Apoptosis Apoptosis is a form of programmed cell death that is a normal mechanism for the elimination of effete cells in organogenesis as well as in the cellular proliferation that occurs during a normal immune response (Chap. 372e). Apoptosis is largely dependent on cellular activation, and the aberrant cellular activation associated with HIV disease is correlated with a heightened state of apoptosis. HIV can trigger both Fas-dependent and Fas-independent pathways of apoptosis, the former of which is generally referred to as activation-induced cell death through an extrinsic pathway and involves the upregulation of the death receptor Fas and Fas ligand. Fas-independent pathways can be either extrinsic with different death receptors or intrinsic due to the downregulation of the antiapoptotic proteins such as Bcl-2. More recently, the phenomenon of *pyroptosis*, an inflammatory form of cell death involving the upregulation of the proinflammatory enzyme caspase-1 and release of the proinflammatory cytokine IL-1 β , has been linked to a bystander effect of HIV replication on CD4+ T cells. Certain viral gene products have been associated with enhanced susceptibility to apoptosis; these include Env, Tat, and Vpr. In contrast, Nef has been shown to possess antiapoptotic properties. A number of studies, including those examining lymphoid tissue, have demonstrated that the rate of apoptosis is elevated in HIV infection and that apoptosis is seen in “bystander” cells such as CD8+ T cells and B cells as well as in uninfected CD4+ T cells. The intensity of apoptosis correlates with the general state of activation of the immune system and not with the stage of disease or with viral burden. It is likely that nonspecific apoptosis of immunocompetent cells related to immune activation contributes to the immune abnormalities in HIV disease.

Autoimmune Phenomena The autoimmune phenomena that are common in HIV-infected individuals reflect, at least in part, chronic immune activation and the dysregulation of B and T cells. Although these phenomena usually occur in the absence of autoimmune disease, a wide spectrum of clinical manifestations that may be associated with autoimmunity have been described (see “Immunologic and Rheumatologic Diseases,” below). Autoimmune phenomena include antibodies against autoantigens expressed on intact lymphocytes and other cells, or against proteins released from dying cells. Antiplatelet antibodies have some clinical relevance in that they may contribute to the thrombocytopenia of HIV disease (see below). Antibodies to nuclear and cytoplasmic components of cells have been reported, as have antibodies to cardiolipin and phospholipids; CD4 molecules; CD43 molecules; C1q-A; variable regions of the T cell receptor α , β , and γ chains; Fas; denatured collagen; and IL-2. In addition, autoantibodies to a range of serum proteins, including albumin, immunoglobulin, and thyroglobulin, have been reported. Molecular mimicry, either from opportunistic pathogens or from HIV itself, is also a trigger or co-factor in autoimmunity. Antibodies against the HIV envelope proteins, especially gp41, often cross-react with host proteins; the best known examples are antibodies directed against the membrane-proximal external region of gp41 that also react with phospholipids and cardiolipin. The phenomenon of polyreactive HIV-specific antibodies may be beneficial to the host (see “Immune Response to HIV,” below).

The increased occurrence and/or exacerbation of certain autoimmune diseases have been reported in HIV infection; these diseases include psoriasis, idiopathic thrombocytopenic purpura, Graves’ disease, antiphospholipid syndrome, and primary biliary cirrhosis. The majority of these manifestations were described prior to the advent of cART and have decreased in frequency since its widespread use. However, with increasing availability of cART, an *immune reconstitution inflammatory syndrome* (IRIS) has become increasingly common in infected individuals, particularly those with low CD4+ T cell counts. IRIS is an autoimmune-like phenomenon characterized by a paradoxical deterioration of clinical condition, which is usually compartmentalized to a particular organ system, in individuals in whom cART has recently been initiated. It is associated with a decrease in viral load and at least partial recovery of immune competence, which is usually associated with increases in CD4+ T cell counts. The immunopathogenesis is felt to be related to an increase in immune response against the presence of residual antigens that are usually microbial and is commonly seen with underlying *Mycobacterium tuberculosis* and cryptococcosis. This syndrome is discussed in more detail below.

CYTOKINES AND OTHER SOLUBLE FACTORS IN HIV PATHOGENESIS

The immune system is homeostatically regulated by a complex network of immunoregulatory cytokines, which are pleiotropic and redundant and operate in an autocrine and paracrine manner. They are expressed continuously, even during periods of apparent quiescence of the immune system. On perturbation of the immune system by antigenic challenge, the expression of cytokines increases to varying degrees (Chap. 372e). Cytokines that are important components of this immunoregulatory network are thought to play major roles in HIV disease, during both the early and chronic phases of infection. A potent pro-inflammatory “cytokine storm” is induced during the acute phase of HIV infection, likely a response by inflammatory cells recruited to mucosal tissues where the virus initially replicates at very high levels. Cytokines that are induced during this early phase include IFN- α , IL-15, and the CXC chemokine IP-10 (CXCL10), followed by IL-6, IL-12, and TNF- α , and a delayed peak of the anti-inflammatory cytokine IL-10. Soluble factors of innate immunity are also induced shortly after infection, including neopterin and β -microglobulin. Several of these early-expressed cytokines and factors are not downregulated following the early phase of HIV infection, as seen in self-resolving viral infections, and persist during the chronic phase of infection and contribute to maintaining high levels of immune activation. Among the cytokines and factors associated with early innate immune responses, they are intended to contain viral replication, although most are potent inducers of HIV expression/replication

because of their ability to induce immune activation that leads to enhanced viral production and an increase in readily available target cells for HIV (activated CD4+ T cells). The induction of IFN- α , one of the first cytokines induced during primary HIV infection, is thought to play a particularly important role in HIV pathogenesis by inducing a large number of IFN-associated genes that activate the immune system and alter the homeostasis of CD4+ T cells. Other cytokines that are elevated during the chronic phase of HIV infection and linked to immune activation include IFN- γ , the CC-chemokine RANTES (CCL5), macrophage inflammatory protein (MIP)-1 β (CCL4), and IL-18.

Several specific cytokines and soluble factors have been associated with HIV pathogenesis at various stages of disease, in various tissues or organs, and in the regulation of HIV replication. Plasma levels of IP-10 are predictive of disease progression, whereas the proinflammatory cytokine IL-6, soluble CD14 (sCD14), and coagulation marker D-dimer are associated with increased risk of all-cause mortality in HIV-infected individuals. In particular, IL-6, sCD14, and D-dimer are associated with increased risk of cardiovascular disease and other causes of death, even in individuals receiving cART. IL-18 has also been shown to play a role in the development of the HIV-associated lipodystrophy syndrome, whereas increased levels of transforming growth factor (TGF)- β are associated with the induction of collagen deposition in lymph nodes (see above). Elevated levels of TNF- α and IL-6 have been demonstrated in plasma and cerebrospinal fluid (CSF), and increased expression of TNF- α , IL-1 β , IFN- γ , and IL-6 has been demonstrated in the lymph nodes of HIV-infected individuals. RANTES, MIP-1 α (CCL3), and MIP-1 β (CCL4) (Chap. 372e) inhibit infection by and spread of R5 HIV-1 strains, while *stromal cell-derived factor* (SDF) 1 inhibits infection by and spread of X4 strains. The mechanisms whereby the CC-chemokines RANTES (CCL5), MIP-1 α (CCL3), and MIP-1 β (CCL4) inhibit infection of R5 strains of HIV, or SDF-1 blocks X4 strains of HIV, involve blocking of the binding of the virus to its co-receptors, the CC-chemokine receptor CCR5 and the CXCR4, respectively. Other soluble factors that have not yet been fully characterized have also been shown to suppress HIV replication, independent of co-receptor usage.

LYMPHOCYTE TURNOVER IN HIV INFECTION

The immune systems of patients with HIV infection are characterized by a profound increase in lymphocyte turnover that is immediately reduced with effective cART. Studies utilizing in vivo or in vitro labeling of lymphocytes in the S-phase of the cell cycle have demonstrated a tight correlation between the degree of lymphocyte turnover and plasma levels of HIV RNA. This increase in turnover is seen in CD4+ and CD8+ T lymphocytes as well as B lymphocytes and can be observed in peripheral blood and lymphoid tissue. Mathematical models derived from these data suggest that one can view the lymphoid pool as consisting of dynamically distinct subpopulations of cells that are differentially affected by HIV infection. A major consequence of HIV infection appears to be a shift in cells from a more quiescent pool to a pool with a higher turnover rate. It is likely that a consequence of a higher rate of turnover is a higher rate of cell death. The role of the thymus in adult human T cell homeostasis and HIV pathogenesis is an area of controversy. While some data point to an important role for the thymus in maintaining T cell numbers and suggest that impairment of thymic function may be responsible for the declines in CD4+ T cells seen in the setting of HIV infection, other studies have concluded that the thymus plays a minor role in HIV pathogenesis. More recently, it has been suggested that the more rapid decline in CD4+ compared to CD8+ T cells may be linked to alterations in inflammatory and homeostatic cytokines that caused increased activation-induced death of CD4+ but not CD8+ T cells (see Table 226-5 for additional mechanisms of depletion).

THE ROLE OF VIRAL RECEPTORS AND CO-RECEPTORS IN HIV PATHOGENESIS

As mentioned above, HIV-1 utilizes two major co-receptors along with CD4 to bind to, fuse with, and enter target cells; these co-receptors are CCR5 and CXCR4, which are also receptors for certain endogenous

TABLE 226-5 PROPOSED MECHANISMS OF CD4+ T CELL DYSFUNCTION AND DEPLETION

Direct Mechanisms	Indirect Mechanisms
Loss of plasma membrane integrity due to viral budding	Aberrant intracellular signaling events
Accumulation of unintegrated viral DNA	Autoimmunity
Activation of DNA-dependent protein kinase during viral integration into host genome	
Interference with cellular RNA processing	Innocent bystander killing of viral antigen-coated cells
Intracellular gp120-CD4 autofusion events	Apoptosis, pyroptosis (caspase-1 associated inflammation), autophagy
Syncytia formation	Inhibition of lymphopoiesis from reduced survival cytokines and lymphoid tissue integrity Activation-induced cell death Elimination of HIV-infected cells by virus-specific immune responses

chemokines. Strains of HIV that utilize CCR5 as a co-receptor are referred to as *R5 viruses*. Strains of HIV that utilize CXCR4 are referred to as *X4 viruses*. Many virus strains are *dual tropic* in that they utilize both CCR5 and CXCR4; these are referred to as *R5X4 viruses*.

The natural chemokine ligands for the major HIV co-receptors can readily block entry of HIV. For example, the CC-chemokines RANTES (CCL5), MIP-1 α (CCL3), and MIP-1 β (CCL4), which are the natural ligands for CCR5, block entry of R5 viruses, whereas SDF-1, the natural ligand for CXCR4, blocks entry of X4 viruses. The mechanism of inhibition of viral entry is a steric inhibition of binding that is not dependent on signal transduction (Fig. 226-25).

The transmitting virus is almost invariably an R5 virus that predominates during the early stages of HIV disease. In ~40% of HIV-infected individuals, there is a transition to a predominantly X4 virus that is associated with a relatively rapid progression of disease. However, at least 60% of infected individuals progress in their disease while maintaining predominance of an R5 virus. It should be pointed out that clade C viruses, unlike other subgroups, almost never switch from CCR5 tropism to CXCR4 tropism; the reason for this difference is unclear.

The basis for the tropism of different envelope glycoproteins for either CCR5 or CXCR4 relates to the ability of the HIV envelope, including the third variable region (V3 loop) of gp120, to interact with these co-receptors. In this regard, binding of gp120 to CD4 induces a conformational change in gp120 that increases its affinity for CCR5 (see above). Finally, R5 viruses are more efficient in infecting monocytes/macrophages and microglial cells of the brain (see “Neuropathogenesis in HIV Disease,” below).

THE INTEGRIN $\alpha 4\beta 7$ AND MUCOSAL TRANSMISSION OF HIV Several “accessory receptors” for HIV have been reported over the years, although only a few have withstood the test of time. These receptors are not necessary for virus binding and fusion to its target CD4+ T cell or for virus replication. However, the integrin $\alpha 4\beta 7$ is an accessory receptor for HIV and it likely plays an important role in the transmission of HIV at mucosal surfaces such as the genital tract and gut. The integrin $\alpha 4\beta 7$, which is the gut homing receptor for peripheral T cells, binds in its activated form to a specific tripeptide in the V2 loop of gp120, resulting in rapid activation of leukocyte function-associated antigen 1 (LFA-1), the central integrin in the establishment of virologic synapses, which facilitate efficient cell-to-cell spread of HIV. It has been demonstrated that $\alpha 4\beta 7^{\text{high}}$ CD4+ T cells are more susceptible to productive infection than are $\alpha 4\beta 7^{\text{low-neg}}$ CD4+ T cells because this cellular subset is enriched with metabolically active CD4+ T cells that are CCR5 $^{\text{high}}$. These cells are present at the gut and genital tract mucosal surfaces. Importantly, it has been demonstrated that the virus that is transmitted during sexual exposure binds much more efficiently to $\alpha 4\beta 7$ than does the virus that diversifies from the transmitting virus over time by mutation,

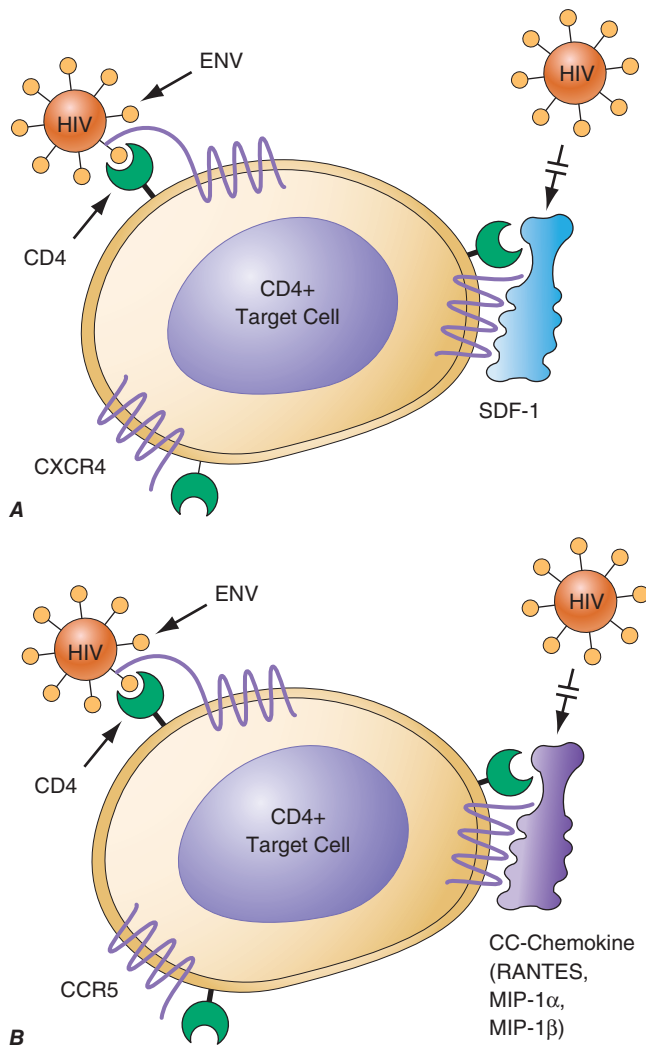


FIGURE 226-25 Model for the role of co-receptors CXCR4 and CCR5 in the efficient binding and entry of X4 (A) and R5 (B) strains of HIV-1, respectively, into CD4+ target cells. Blocking of this initial event in the virus life cycle can be accomplished by inhibition of binding to the co-receptor by the normal ligand for the receptor in question. The ligand for CXCR4 is stromal cell-derived factor (SDF-1); the ligands for CCR5 are RANTES, MIP-1 α , and MIP-1 β .

particularly involving the accumulation of glycosylation sites (see “Early Events in HIV Infection: Primary Infection and Initial Dissemination of Virus,” above).

CELLULAR TARGETS OF HIV

Although the CD4+ T lymphocytes and to a lesser extent CD4+ cells of monocyte lineage are the principal targets of HIV, virtually any cell that expresses the CD4 molecule together with co-receptor molecules (see above and below) can potentially be infected with HIV. Circulating DCs have been reported to express low levels of CD4, and, depending on their stage of maturation, these cells can be infected with HIV. Epidermal Langerhans cells express CD4 and have been infected by HIV *in vivo*, although, as has been shown *in vivo* for DCs, FDCs, and B cells, these cells are more likely to bind and transfer virus to activated CD4+ T cells than to be productively infected themselves.

In vitro, HIV has been reported also to infect a wide range of cells and cell lines that express low levels of CD4, no detectable CD4, or only CD4 mRNA. However, since the only cells that have been shown unequivocally to be infected with HIV and to support replication of the virus are CD4+ T lymphocytes and cells of monocyte/macrophage lineage, the physiologic relevance of the *in vitro* infection of these other cell types is unclear.

Of potentially important clinical relevance is the demonstration that thymic precursor cells, which were assumed to be negative for CD3, CD4, and CD8 molecules, actually do express low levels of CD4 and can be infected with HIV *in vitro*. In addition, human thymic epithelial cells transplanted into an immunodeficient mouse can be infected with HIV by direct inoculation of virus into the thymus. Since these cells may play a role in the normal regeneration of CD4+ T cells, it is possible that their infection and depletion contribute, at least in part, to the impaired ability of the CD4+ T cell pool to completely reconstitute itself in certain infected individuals in whom cART has suppressed viral replication to <50 copies of HIV RNA per milliliter (see below). In addition, CD34+ monocyte precursor cells have been shown to be infected *in vivo* in patients with advanced HIV disease. It is likely that these cells express low levels of CD4, and therefore it is not essential to invoke CD4-independent mechanisms to explain the infection.

ABNORMALITIES OF MONONUCLEAR CELLS

CD4+ T Cells The primary immunopathogenic lesion in HIV infection involves CD4+ T cells, and the range of CD4+ T cell abnormalities in advanced HIV infection is broad. The defects are both quantitative and qualitative and ultimately impact virtually every limb of the immune system, indicating the critical dependence of the integrity of the immune system on the inducer/helper function of CD4+ T cells. In advanced HIV disease, most of the observed immune defects can ultimately be explained by the quantitative depletion of CD4+ T cells. However, T cell dysfunction can be demonstrated in patients early in the course of infection, even when the CD4+ T cell count is in the low-normal range. The degree and spectrum of dysfunctions increase as the disease progresses, reflecting the range of CD4+ T cell functional heterogeneity, especially in lymphoid tissues. One of the first sites of intense HIV replication is in the GALT where CD4+ T_H17 cells reside; they are important for host defense against extracellular pathogens in the intestinal mucosa and help maintain the integrity of the gut epithelium. In HIV infection, they are depleted by direct and indirect effects of viral replication and cause loss of gut homeostasis and integrity, as well as a shift to a more T_H1 phenotype. Studies have shown that even after many years of cART, normalization of the CD4+ T cells in the GALT remains incomplete. In lymph nodes, HIV perturbs another important subset of the CD4+ helper T lineage, namely TF_H cells (see “Lymphoid Organs and HIV Pathogenesis,” above). TF_H cells, which are either derived directly from naïve CD4+ T cells or other T_H precursors, migrate into B follicles during germinal center reactions and provide help to antigen-specific B cells through cell-cell interactions and secretion of cytokines to which B cells respond, the most important of which is IL-21. As with T_H17 cells, TF_H cells are highly susceptible to HIV infection. However, in contrast to T_H17 and most other CD4+ T cell subsets, the number of TF_H cells is increased in lymph nodes of HIV-infected individuals, especially those who are viremic. It is unclear whether this increase is helpful to responding B cells, although the likely outcome is that the increase in numbers is detrimental to the quality of the humoral immune response against HIV (see “Immune Response to HIV,” below). In addition, defects of central memory cells are a critical component of HIV immunopathogenesis. The progressive loss of antigen-specific CD4+ T cells has important implications for the control of HIV infection. In this regard, there is a correlation between the maintenance of HIV-specific CD4+ T cell proliferative responses and improved control of infection. Essentially every T cell function has been reported to be abnormal at some stage of HIV infection. Loss of polyfunctional HIV-specific CD4+ T cells, especially those that produce IL-2, occurs early in disease, whereas IFN-producing CD4+ T cells are maintained longer and do not correlate with control of HIV viremia. Other abnormalities include impaired expression of IL-2 receptors, defective IL-2 production, reduced expression of the IL-7 receptor (CD127), and a decreased proportion of CD4+ T cells that express CD28, a major co-stimulatory molecule necessary for the normal activation of T cells, which is also depleted as a result of aging. Cells lacking expression of CD28 do not respond normally to activation signals and may express markers of terminal activation including HLA-DR, CD38, and CD45RO. As mentioned

above (“Immune Activation, Inflammation, and HIV Pathogenesis”), a subset of CD4+ T cells referred to as *T regulatory cells*, or T-regs, may be involved in damping aberrant immune activation that propagates HIV replication. The presence of these T-reg cells correlates with lower viral loads and higher CD4+/CD8+ T cell ratios. A loss of this T-reg capability with advanced disease may be detrimental to the control of virus replication.

It is difficult to explain completely the profound immunodeficiency noted in HIV-infected individuals solely on the basis of direct infection and quantitative depletion of CD4+ T cells. This is particularly apparent during the early stages of HIV disease, when CD4+ T cell numbers may be only marginally decreased. In this regard, it is likely that CD4+ T cell dysfunction results from a combination of depletion of cells due to direct infection of the cell and a number of virus-related but indirect effects on the cell (Table 226-5). Several of these effects have been demonstrated *ex vivo* and/or by the analysis of cells isolated from the peripheral blood. However, as explained above, many of the defects are related to specialized CD4+ T cells that reside in lymphoid tissues. Furthermore, since the main targets of HIV infection are immunocompetent cells, these responses may contribute to immune cell depletion and immunologic dysfunction by eliminating both infected cells and “innocent bystander” cells. Soluble viral proteins, particularly gp120, can bind with high affinity to the CD4 molecules on uninfected T cells and monocytes; in addition, virus and/or viral proteins can bind to DCs or FDCs. HIV-specific antibody can recognize these bound molecules and potentially collaborate in the elimination of the cells by ADCC. HIV envelope glycoproteins gp120 and gp160 manifest high-affinity binding to the CD4 molecule as well as to various chemokine receptors. Intracellular signals transduced by gp120 through both CD4 and CCR5/CXCR4 have been associated with a number of immunopathogenic processes including anergy, apoptosis, and abnormalities of cell trafficking. The molecular mechanisms responsible for these abnormalities include dysregulation of the T cell receptor–phosphoinositide pathway, p56lck activation, phosphorylation of focal adhesion kinase, activation of the MAP kinase and ras signaling pathways, and downregulation of the co-stimulatory molecules CD40 ligand and CD80.

The inexorable decline in CD4+ T cell counts that occurs in most HIV-infected individuals may result in part from the inability of the immune system to regenerate over an extended period of time the rapidly turning over CD4+ T cell pool efficiently enough to compensate for both HIV-mediated and naturally occurring attrition of cells. In this regard, the degree and duration of decline of CD4+ T cells at the time of initiation of therapy is an important predictor of the restoration of these cells. A person who maintains a very low CD4+ T cell count for a considerable period of time before the initiation of cART almost invariably has an incomplete reconstitution of such cells. At least two major mechanisms may contribute to the failure of the CD4+ T cell pool to reconstitute itself adequately over the course of HIV infection. The first is the destruction of lymphoid precursor cells, including thymic and bone marrow progenitor cells; the other is the gradual disruption of the lymphoid tissue microenvironment, which is essential for efficient regeneration of immunocompetent cells. Finally, during the advanced stages of CD4+ T lymphopenia, there are increased serum levels of the homeostatic cytokine IL-7. It was initially felt that this elevation was a homeostatic response to the lymphopenia; however, recent findings suggest that the increase in serum IL-7 was a result of reduced utilization of the cytokine related to the loss of cells expressing the IL-7 receptor, CD127, which serves as a normal physiologic regulator of IL-7 production.

CD8+ T Cells A relative CD8+ T lymphocytosis is generally associated with high levels of HIV plasma viremia and likely reflects an immune response to the virus as well as dysregulated homeostasis associated with generalized immune activation. During the late stages of HIV infection, there may be a significant reduction in the numbers of CD8+ T cells despite the presence of high levels of viremia. HIV-specific CD8+ CTLs have been demonstrated in HIV-infected individuals early in the course of disease, and their emergence often coincides

with a decrease in plasma viremia—an observation that is a factor in the proposal that virus-specific CTLs can control HIV disease for a finite period of time in a certain percentage of infected individuals. However, emergence of HIV escape mutants that ultimately evade these HIV-specific CD8+ T cells has been described in the majority of HIV-infected individuals who are not receiving cART. In addition, as the disease progresses, the functional capability of these cells gradually decreases, at least in part due to the persistent nature of HIV infection that causes functional exhaustion via the upregulation of inhibitory receptors such as PD-1 on HIV-specific CD8+ T cells (see “Immune Activation, Inflammation, and HIV Pathogenesis,” above). As chronic immune activation persists, there are also systemic effects on CD8+ T cells, such that as a population they assume an abnormal phenotype characterized by expression of activation markers such as HLA-DR and CD38 with an absence of expression of the IL-2 receptor (CD25) and a reduced expression of the IL-7 receptor (CD127). In addition, CD8+ T cells lacking CD28 expression are increased in HIV disease, reflecting a skewed expansion of a less differentiated CD8+ T cell subset. This skewing of subsets is also associated with diminished polyfunctionality, a qualitative difference that distinguishes nonprogressors from progressors. It has been reported that nonprogressors can also be distinguished from progressors by the maintenance in the former of a high proliferative capacity of their HIV-specific CD8+ T cells coupled to increases in perforin expression, characteristics that are markedly diminished in advanced HIV disease. It has been reported that the phenotype of CD8+ T cells in HIV-infected individuals may be of prognostic significance. Those individuals whose CD8+ T cells developed a phenotype of HLA-DR+/CD38– following seroconversion had stabilization of their CD4+ T cell counts, whereas those whose CD8+ T cells developed a phenotype of HLA-DR+/CD38+ had a more aggressive course and a poorer prognosis. In addition to the defects in HIV-specific CD8+ CTLs, functional defects in other MHC-restricted CTLs, such as those directed against influenza and CMV, have been demonstrated. CD8+ T cells secrete a variety of soluble factors that inhibit HIV replication, including the CC-chemokines RANTES (CCL5), MIP-1 α (CCL3), and MIP-1 β (CCL4) as well as potentially a number of unidentified factors. The presence of high levels of HIV viremia *in vivo* as well as exposure of CD8+ T cells *in vitro* to HIV envelope, both of which are associated with aberrant immune activation, have been shown to be associated with a variety of cellular functional abnormalities. Furthermore, since the integrity of CD8+ T cell function depends in part on adequate inductive signals from CD4+ T cells, the defect in CD8+ CTLs is likely compounded by the quantitative loss and qualitative dysfunction of CD4+ T cells.

B Cells The predominant defect in B cells from HIV-infected individuals is one of aberrant cellular activation, which is reflected by increased propensity to terminal differentiation and immunoglobulin secretion and increased expression of markers of activation and exhaustion. As a result of activation and differentiation *in vivo*, B cells from HIV viremic patients manifest a decreased capacity to mount a proliferative response to ligation of the B cell antigen receptor and other B cell stimuli *in vitro*. B cells from HIV-infected individuals manifest enhanced spontaneous secretion of immunoglobulins *in vitro*, a process that reflects their highly differentiated state *in vivo*. There is also an increased incidence of EBV-related B cell lymphomas in HIV-infected individuals that are likely due to combined effects of defective T cell immune surveillance and increased turnover that increases the risk of oncogenesis. Untransformed B cells cannot be infected with HIV, although HIV or its products can activate B cells directly. B cells from patients with high levels of viremia bind virions to their surface via the CD21 complement receptor. It is likely that *in vivo* activation of B cells by replication-competent or defective virus as well as viral products during the viremic state accounts at least in part for their activated phenotype. B cell subpopulations from HIV-infected individuals undergo a number of changes over the course of HIV disease, including the attrition of resting memory B cells and replacement with several aberrant memory and differentiated B cell subpopulations that collectively express reduced levels of CD21

and either increased expression of activation markers or inhibitory receptors associated with functional exhaustion. The more activated and differentiated B cells are also responsible for increased secretion of immunoglobulins and increased susceptibility to Fas-mediated apoptosis. In more advanced disease, there is also the appearance of immature B cells associated with CD4+ T cell lymphopenia. Cognate B cell–CD4+ T cell interactions are abnormal in viremic HIV-infected individuals in that B cells respond poorly to CD4+ T cell help and CD4+ T cells receive inadequate co-stimulatory signals from activated B cells. In vivo, the aberrant activated state of B cells manifests itself by hypergammaglobulinemia and by the presence of circulating immune complexes and autoantibodies. HIV-infected individuals respond poorly to primary and secondary immunizations with protein and polysaccharide antigens. Using immunization with influenza vaccine, it has been demonstrated that there is a memory B cell defect in HIV-infected individuals, particularly those with high levels of HIV viremia. There is also evidence that responses to HIV and non-HIV antigens in infected individuals, especially those who remain viremic, are enriched in abnormal subsets of B cells that either are highly prone to apoptosis or show signs of functional exhaustion. Taken together, these B cell defects are likely responsible in part for the inadequate response to HIV as well as to decreased response to vaccinations and the increase in certain bacterial infections seen in advanced HIV disease in adults, as well as for the important role of bacterial infections in the morbidity and mortality rates of HIV-infected children, who cannot mount an adequate humoral response to common bacterial pathogens. The absolute number of circulating B cells may be depressed in HIV infection; this phenomenon likely reflects increased activation-induced apoptosis as well as a redistribution of cells out of the circulation and into the lymphoid tissue—phenomena that are associated with ongoing viral replication.

Monocytes/Macrophages Circulating monocytes are generally normal in number in HIV-infected individuals; however, there is evidence of increased activation within this lineage. The increased level of sCD14 and other biomarkers (see above) reported in HIV-infected individuals is an indirect marker of monocyte activation in vivo. A number of other abnormalities of circulating monocytes have been reported in HIV-infected individuals, many of which may be related directly or indirectly to aberrant in vivo immune activation. In this regard, increased levels of lipopolysaccharide (LPS) are found in the sera of HIV-infected individuals due, at least in part, to translocation across the gut mucosal barrier (see above). LPS is a highly inflammatory bacterial product that preferentially binds to macrophages through CD14 and Toll-like receptors, resulting in cellular activation. Functional abnormalities of monocyte/macrophages in HIV disease include decreased secretion of IL-1 and IL-12; increased secretion of IL-10 and IL-18; defects in antigen presentation and induction of T cell responses due to decreased MHC class II expression; and abnormalities of Fc receptor function, C3 receptor-mediated clearance, oxidative burst responses, and certain cytotoxic functions such as ADCC, possibly related to low levels of expression of Fc and complement receptors. Monocytes express the CD4 molecule and several co-receptors for HIV on their surface, including CCR5, CXCR4, and CCR3, and thus are potential targets of HIV infection. The degree of cytopathicity of HIV for cells of the monocyte lineage is low, and HIV can replicate in cells of the monocyte lineage with relatively little cytopathic effect. Hence, monocyte-lineage cells may play a role in the dissemination of HIV in the body and can serve as reservoirs of HIV infection, thus representing an obstacle to the eradication of HIV by antiretroviral drugs. In vivo infection of circulating monocytes is difficult to demonstrate; however, infection of tissue macrophages and macrophage-lineage cells in the brain (infiltrating macrophages or resident microglial cells) and lung (pulmonary alveolar macrophages) can be demonstrated easily. Tissue macrophages are an important source of HIV during the inflammatory response associated with opportunistic infections. Infection of monocyte precursors in the bone marrow may directly or indirectly be responsible for certain of the hematologic abnormalities in HIV-infected individuals. However, as with DCs, monocytes and

macrophages express high levels of host restriction factors that likely help explain the low contribution of myeloid cells to the overall viral burden in HIV-infected individuals.

Dendritic and Langerhans Cells DCs and Langerhans cells are thought to play an important role in the initiation of HIV infection by virtue of the ability of HIV to bind to cell-surface C-type lectin receptors, particularly DC-SIGN (see above) and Langerin. This allows efficient presentation of virus to CD4+ T cell targets that become infected; complexes of infected CD4+ T cells and DCs provide an optimal microenvironment for virus replication. There was once considerable disagreement regarding the HIV infectibility and hence the depletion as well as the dysfunction of DCs themselves. However, since the recognition of myeloid (mDC) and plasmacytoid (pDC) subsets, there has been a better appreciation of specific DC dysfunction in HIV disease. pDCs are an important component of the innate immune system and secrete large amounts of IFN- α in response to viral infections. The numbers of circulating pDCs are decreased in HIV infection through mechanisms that remain unclear, and there are conflicting reports regarding the frequency of pDCs in lymphoid tissues, with some studies suggesting that their increased tissue presence and secretion of inflammatory cytokines such as IFN- α contributes to lymphoid hyperplasia. The mDCs or conventional DCs are involved in the initiation of adaptive immunity in draining lymph nodes by presenting antigen to T cells and B cells, as well as by secreting cytokines such as IL-12, IL-15, and IL-18 that activate other immune cells. There are also indications that the relatively low infectibility of DCs may be associated with the expression of host restriction factors, including APOBEC3G (see above).

Natural Killer Cells The role of NK cells is to provide immunosurveillance against virus-infected cells, certain tumor cells, and allogeneic cells (Chap. 372e). There are no convincing data that HIV productively infects NK cells in vivo; however, functional abnormalities in NK cells have been observed throughout the course of HIV disease, and the severity of these abnormalities increases as disease progresses. NK cells are part of the innate immune system and act by direct killing of infected cells and secretion of antiviral cytokines. In early HIV infection there is an increase in the activation of NK cells, and the capacity to secrete IFN- γ is maintained, although they manifest reduced cytotoxic function. During chronic HIV infection, both NK cell cytotoxicity and cytokine secretion become impaired. Given that HIV infection of target cells downregulates HLA-A and -B, but not HLA-C and -D molecules, this may explain in part the relative inability of NK cells to kill HIV-infected target cells. However, the NK cell impairments, especially in patients with high levels of virus replication, are associated with an expansion of an “anergic” CD56–/CD16+ NK cell subset. This abnormal subset of NK cells manifests an increased expression of inhibitory NK cell receptors (iNKR) and a substantial decrease in expression of natural cytotoxicity receptors (NCR) and shows a markedly impaired lytic activity. The overrepresentation of this abnormal subset of NK cells may explain in part the observed defects in NK cell function in HIV-infected individuals and likely begins to occur during primary infection. NK cells also serve as important sources of HIV-inhibitory CC-chemokines. NK cells isolated from HIV-infected individuals constitutively produce high levels of MIP-1 α (CCL3), MIP-1 β (CCL4), and RANTES (CCL5), although the impact of these chemokines on HIV replication in vivo is unclear. Finally, NK cell–DC interactions are important for normal immune function. NK cells and DCs reciprocally modulate each other’s activation and maturation. These interactions are markedly impaired in HIV-infected individuals with high levels of plasma viremia.

GENETIC FACTORS IN HIV-1 AND AIDS PATHOGENESIS

PHENOTYPES OF SUSCEPTIBILITY AND RESPONSE TO HIV INFECTION It is well known that individuals vary in their susceptibility to acquiring HIV infection and that there is wide variation in both the steady-state level of HIV that is established soon after infection (virologic setpoint) as well as the rate at which HIV-infected patients progress to AIDS. Some striking examples include sex workers who remain uninfected despite repeated exposure to HIV; HIV-infected individuals who

spontaneously control viral replication in the absence of cART (HIV controllers); patients who resist disease progression for at least 8–10 years, despite viremia; and those progressing to AIDS within 3 years. Investigators have hypothesized that genetic differences may partly explain this interindividual variation in risk of acquiring HIV infection and disease progression rates. In addition to these phenotypes, it has been hypothesized that genetic variation may partly underpin the risk of developing specific AIDS-defining illnesses (e.g., renal and neurologic diseases) and non-AIDS comorbidities (e.g., cardiovascular disease), as well as the variable recovery in CD4+ T cell counts observed while receiving cART.

Candidate gene approaches and genome-wide association studies (GWAS) have demonstrated associations between gene variations and the above-mentioned phenotypes. A list of some of these associations is shown in [Table 226-6](#). While *in vitro* genome-wide functional scanning using RNA interference has identified hundreds of host factors that may be involved in the HIV life cycle, the association of these genes with HIV susceptibility and/or disease progression remains largely undefined. Below is a discussion of a few key genes with strong associations and their implications for improving clinical care.

ASSOCIATIONS WITH CCR5 AND TRANSLATION OF GENETIC FINDINGS TO THE CLINIC Possibly, the most dramatic example of the importance of genetic studies for identifying host factors that influence HIV-AIDS pathogenesis is from studies related to the gene that encodes for CC chemokine receptor 5 (CCR5). While *in vitro* studies established that CCR5 is the major HIV co-receptor for the cell entry of HIV-1 into the host, it was genetic studies that established the seminal *in vivo* role of this receptor for the initial entry of HIV and AIDS pathogenesis.

Genetic analysis revealed that the *in vitro* resistance to CCR5-using R5 strains of HIV is in some instances due to carriage of two defective CCR5 alleles. This defect is a 32-bp deletion in the coding sequence (designated as the $\Delta 32$ allele). The CCR5 $\Delta 32$ allele encodes a truncated protein that is not expressed on the cell surface.

Approximately 1% of individuals of European ancestry are homozygous for the CCR5 $\Delta 32$ allele. Depending on the geographic region in Europe, up to 20% of individuals are heterozygous for the CCR5 $\Delta 32$ allele. The CCR5 $\Delta 32$ allele is either absent or extremely rare in other populations. The evolutionary pressure that resulted in the emergence of the CCR5 $\Delta 32$ allele in the European population remains unknown and has been speculated to be secondary to an ancestral pandemic such as the plague.

Individuals homozygous for the CCR5 $\Delta 32$ allele ($\Delta 32/\Delta 32$) lack CCR5 surface expression and are highly resistant to acquiring HIV infection. Heterozygosity for the CCR5 $\Delta 32$ allele is also associated with a reduced risk of acquiring HIV. Consequently, the frequency of the CCR5 $\Delta 32$ allele is enriched in individuals of European descent who remain uninfected despite exposure to the virus. Although the CCR5 $\Delta 32/\Delta 32$ genotype is associated with profound resistance to acquiring HIV, a few individuals with this genotype have become infected with the X4 HIV strain and, in some instances, experienced an accelerated disease course. In general, CCR5 $\Delta 32$ heterozygosity is associated with a slower HIV disease course.

Subsequent studies identified single nucleotide polymorphisms (SNPs) in the promoter (regulatory region) of CCR5 that influence its expression levels. Alleles bearing specific cassettes of linked polymorphisms (haplotypes) were identified and designated as human haplogroups A to G*2 (HHA to HHG*2). The CCR5 $\Delta 32$ is found on the HHG*2 haplotype. The CCR5 HHE haplotype was associated with higher CCR5 expression, and genetic association studies have shown that homozygosity for the CCR5 HHE haplotype is associated with an increased risk of acquiring HIV, progressing rapidly to AIDS, and reduced immune recovery on cART. The CCR2-64I-bearing HHF*2 haplotype is associated with a slower HIV disease course. The CCR5 HHA haplotype is the ancestral CCR5 haplotype and is associated with a lower CCR5 expression. The HHA haplotype was associated with slower disease progression in African populations and has been speculated to be a basis for why SIV-infected chimpanzees (who all carry the ancestral CCR5 HHA haplotype) may resist disease progression. The

CCR5 haplotypes also influence cell-mediated immunity and immune recovery on cART.

The association of variations in the CCR5 gene with HIV-AIDS phenotypes is also an example of how discoveries made in the laboratory (bench) have been translated to improve health outcomes (bedside). The discovery that the CCR5 $\Delta 32/\Delta 32$ genotype is associated with strong resistance to HIV infection, and that uninfected Caucasians bearing this genotype did not appear to have impaired immunity, led to the development of two kinds of therapies. First, it spurred the development of a new class of FDA-approved therapies, entry inhibitors (e.g., maraviroc), that block the interaction of CCR5 with the HIV envelope. Second, it led to the development of novel experimental cellular therapies. An HIV-infected patient with acute myelogenous leukemia was given an allogeneic stem-cell transplantation from an HLA-compatible person whose cells lacked expression of CCR5 due to the $\Delta 32/\Delta 32$ genotype. There has been no evidence of HIV-1 infection in the transplanted patient thus far (6 years). This observation spurred the hope of an HIV cure and led to the development of additional novel cellular therapies involving autologous transplantation of CD4+ T-cells in which the CCR5 gene is inactivated *ex vivo* using new gene editing procedures.

DISCOVERY OF HLA CLASS I ALLELES THAT ASSOCIATE WITH VIROLOGIC CONTROL OF HIV INFECTION There is a strong association between variations within the HLA-B gene with protective (e.g., HLA-B*57 and -B*27 alleles) or detrimental (e.g., HLA-B*35 allele) outcomes during HIV infection.

Carriage of the HLA-B*57 and/or HLA-B*27 alleles is associated with slower disease progression. The beneficial effects of these alleles may relate in part to their consistent associations with a lower virologic setpoint as well as to higher cell-mediated immunity. The protective effect of the HLA-B*57 and -B*27 alleles on HIV disease course is underscored by the finding that the prevalence of these alleles is higher among long-term nonprogressors and HIV elite controllers (see above). On the other hand, the HLA-B*35 allele has been associated with faster progression to AIDS and higher viral load. The prevalence of the HLA-B alleles differs between populations. HLA-B*57:01 in Europeans and HLA-B*57:03 in African Americans are the protective alleles. In some populations (e.g., Japanese) where the HLA-B*57/-B*27 alleles are absent, HLA-B*51 is associated with a protective phenotype.

Possession of the protective HLA-B alleles is associated with broader and stronger CD8+ T cell responses to HIV epitopes. The mechanisms underlying the differential effects of the HLA-B alleles on HIV disease course may relate to differences in the ability of antigen-presenting cells to present immunodominant HIV epitopes to T helper or cytotoxic T lymphocytes in the context of MHC-encoded molecules. This may result in differential immune responses that influence viral replication. In this regard, the HLA-B alleles that impact HIV disease course differ in their amino acid residues in the HLA-B peptide-binding groove—and this may play a critical role in virologic control.

Investigators have also examined the influence of extended HLA haplotypes (linked alleles) on HIV disease course. The extended HLA ancestral haplotype (AH) 8.1 is defined by the presence of HLA-A1, HLA-B8, and HLA-DR3 alleles. AH 8.1 is the most common ancestral haplotype in Caucasians (present in 10%) and is associated with multiple autoimmune diseases in HIV-uninfected persons. These associations of AH 8.1 are thought to be due to a genetically determined hyperresponsiveness characterized by high TNF- α production and lack of complement C4A. Strong epidemiologic data indicate that carriage of AH 8.1 in HIV-infected persons is associated with a rapid decline in CD4+ T cells and faster progression to AIDS development. Gene-gene interactions between HLA alleles and other genes (e.g., killer cell immunoglobulin-like receptors) also may influence HIV disease progression rates.

POLYMORPHISMS IDENTIFIED BY GWAS THAT ASSOCIATE WITH VIROLOGIC CONTROL Large-scale GWASs have been conducted for the phenotype of viral load, including in a large group of HIV controllers. GWAS in HIV-infected persons of European ancestry identified four SNPs in genes in the HLA class I loci that associated with virologic control.

TABLE 226-6 HOST GENETIC FACTORS THAT INFLUENCE RISK OF HIV-1 ACQUISITION AND RATES OF HIV-1 DISEASE PROGRESSION

Gene ^a	Genetic Variation	Mechanisms ^b	Genetic Effect on HIV-AIDS ^c
Genes in MHC Locus			
<i>HLA-B</i>	B*27 and B*57 B*35	Presentation of specific HIV antigens Restriction of specific HIV peptide presentation	Slower progression to AIDS; lower viral load Faster progression to AIDS; higher viral load
<i>HLA class I allele</i>	HLA-Bw4	Providing ligands for activating KIR	Slower progression to AIDS
	Homozygosity of <i>HLA</i> -class I alleles	Reduced repertoire for epitope recognition	Faster progression to AIDS; increased risk of mother-to-child transmission
<i>HLA class II allele</i>	Shared donor-recipient <i>HLA</i> alleles	Preadaptation of HIV strains	Faster disease progression
	Rare <i>HLA</i> alleles	Limited adaptation of HIV strains; less frequent escape mutants	Protection against HIV infection
<i>HLA class II allele</i>	<i>HLA-DRB1</i> alleles	Influences protein specificity of CD4+ T cell responses to HIV Gag and Nef proteins	<i>HLA-DRB1*15:02</i> —lower viral load; <i>HLA-DRB1*03:01</i> —higher viral load
<i>HLA</i> extended haplotype	A1-B8-DR3-DQ2 (AH 8.1)	Increased proinflammatory responses; higher TNF- α production	Faster progression to AIDS
<i>HLA-C</i>	35 kb upstream, rs9264942-C	Increased expression of <i>HLA-C</i>	Decreased viral load set point
<i>HCP5</i>	rs2395029-G	Linkage disequilibrium with <i>HLA-B*57:01</i>	Lower viral load
<i>MICA</i>	Noncoding SNP near <i>MICA</i> , rs4418214-T	May affect <i>HLA</i> class I peptide presentation—linkage with protective <i>HLA-B</i> alleles	Enriched in HIV-1 controllers
<i>PSORS1C3</i>	rs3131018-A	May affect <i>HLA</i> class I peptide presentation	Enriched in HIV-1 controllers
<i>ZNRD1</i>	rs9261174-C	Possible interference in processing of HIV transcripts; influences <i>ZNRD1</i> expression; linkage disequilibrium with <i>HLA-A10</i>	AIDS disease retardation
Chemokine Receptors			
<i>CCR5</i>	32-bp deletion in the ORF (Δ 32), rs333	Truncated <i>CCR5</i> protein	Δ 32/ Δ 32: resistance to acquiring HIV infection Δ 32/wild type: delays AIDS onset; improves immune reconstitution during ART
<i>CCR2</i>	Promoter SNPs, haplotypes (HHA to HHG*2)	Altered <i>CCR5</i> expression, e.g., HHE allele correlates with high <i>CCR5</i> expression	HHE/HHE: increased HIV/AIDS susceptibility
	SNP in ORF (64 V \rightarrow I), rs1799864	Possibly due to linkage with polymorphism in <i>CCR5</i> promoter	64I: delayed AIDS onset
<i>CXCR6</i>	rs2234358 G \rightarrow T in the 3'UTR	Trafficking of effector T cells and activation of NK T cells Minor HIV co-receptor	Prevalence of rs2234358-T was lower in long-term nonprogressors
<i>CX3CR1</i>	SNPs in ORF (249 V \rightarrow I); and 280 T \rightarrow M, rs3732378)	280M reduces receptor expression and binding of fractalkine, the CX3CR1 ligand	249I and 280M are associated with faster AIDS onset in some Caucasian cohorts; inconsistent effects were detected in other cohorts
<i>DARC</i>	African-specific promoter SNP (-46T \rightarrow C), rs2814778	-46C/C associates with low neutrophil counts; influences circulating chemokine levels; alters HIV binding to RBCs and transfection of HIV-1	-46C/C: increased risk of acquiring HIV but slower HIV disease progression; Duffy-null-associated low neutrophil trait associated with increased HIV risk in Africans
Chemokines			
<i>CCL3L</i> , <i>CCL4L</i>	Gene copy number of <i>CCL3L</i> and <i>CCL4L</i>	High numbers of <i>CCL3L</i> and <i>CCL4L</i> gene-containing segmental duplications correlate with high <i>CCL3L</i> and <i>CCL4L</i> levels	Gene copy number lower than population median associates with increased HIV/AIDS susceptibility and reduced immune reconstitution during ART
<i>CCL5</i>	Promoter SNPs	Altered gene expression	Altered HIV-AIDS susceptibility
<i>CCL2</i>	Promoter SNP (-2578 T \rightarrow G), rs1024611	-2578G allele: increased <i>CCL2</i> expression and monocyte recruitment	-2578G/G associates with increased risk of developing HIV-1-associated dementia and a rapid AIDS onset
Cytokines			
<i>IL-6</i>	Promoter SNP (-174 G \rightarrow C), rs1800795	-174G/G associates with increased IL-6 and CRP levels	-174G/G associates with high risk of KS development and variable recovery of CD4 cells during ART
<i>IL-10</i>	Promoter SNP (-592 C \rightarrow A), rs1800872	-592A results in decreased IL-10 levels	-592A associates with increased HIV-AIDS susceptibility
Innate Immunity Genes			
<i>MBL</i>	Coding alleles (O)	Low plasma concentration and structural variation of <i>MBL</i> protein	Slow progression to AIDS in heterozygous subjects (A/O)
	X allele (promoter SNP -221)	Decreased levels of <i>MBL</i> protein	Faster progression to AIDS in homozygous X/X subjects
<i>Apobec-3G</i>	ORF SNP (186 H \rightarrow R), rs8177832	Reduced anti-HIV-1 activity	186R associates with rapid AIDS onset in African Americans
<i>TLR7</i>	ORF SNP (Gln11Leu), rs179008	Decreased expression of <i>TLR7</i> leading to lack of recognition of HIV-infected cells	The Leu-containing protein associated with higher viral load and faster progression to AIDS

(Continued)

TABLE 226-6 HOST GENETIC FACTORS THAT INFLUENCE RISK OF HIV-1 ACQUISITION AND RATES OF HIV-1 DISEASE PROGRESSION (CONTINUED)

Gene ^a	Genetic Variation	Mechanisms ^b	Genetic Effect on HIV-AIDS ^c
<i>PARD3B</i>	rs11884476 (C→G), near exon 20	Direct interaction with HIV, signaling through SMAD family of proteins	rs11884476-G associated with slower progression to AIDS
Others			
<i>ApoE</i>	E4 allele	E4 enhances HIV cell entry in vitro	ApoE4/E4 associates with rapid AIDS onset and dementia
<i>ApoL1/MYH9</i>	Several risk haplotypes including G1	Unknown	Increased risk for HIV-associated nephropathy
<i>RYR3</i>	ORF SNP (A →G), rs2229116	Unknown, potential impact on calcium signaling and homeostasis	rs2229116-G associated with subclinical atherosclerosis
<i>PROX1</i>	rs17762192-G, 36kb upstream of <i>PROX1</i>	Unknown, presumably due to its impact on <i>PROX1</i> expression, which is a negative regulator of IFN-γ	rs17762192-G: reduced rate of disease progression
Gene–Gene Interaction			
<i>KIR+HLA</i>	<i>KIR3DS1</i> + <i>HLA-Bw4-80Ile</i>	Altered NK cell activity required to eliminate HIV-infected cells	<i>KIR3DS1</i> with <i>HLA Bw4-80I</i> +: delayed AIDS onset
	<i>HLA-C1</i> + <i>KIR2DL3</i> ,	Reduction of inhibitory <i>KIR</i> likely results in increased immune activation; impaired killing of latently infected cells; and a higher proviral burden	<i>HLA-C1</i> +/ <i>KIR2DL3</i> +: better immune recovery after viral load suppression on ART
<i>LILRB2+HLA</i>	<i>LILRB2</i> + <i>HLA class I</i>	Regulation of dendritic cells by <i>LILRB2</i> - <i>HLA</i> engagement	Control of HIV-1
<i>CCL3L1+CCR5</i>	Low <i>CCL3L1</i> gene copies + detrimental <i>CCR5</i> genotypes	Low <i>CCL3L1</i> and high <i>CCR5</i> expression	Increased HIV/AIDS susceptibility and reduced immune reconstitution during ART

^aRepresentative genes and polymorphisms and ^bpossible mechanisms are listed. ^cSome of the associations are population specific and may display cohort-specific effects.

Note: Apobec, apolipoprotein B mRNA editing enzyme, catalytic polypeptide-like; ApoE, apolipoprotein E; ART, antiretroviral therapy; CCL, CC ligand; CCL3L, CCL3-like; CCR5, CC chemokine receptor 5; CRP, C-reactive protein; CXCR6, chemokine (C-X-C motif) receptor 6; DARC, Duffy antigen receptor for chemokines; HCP5, HLA class I histocompatibility antigen protein P5; HHE, human haplogroup E; HLA, human leukocyte antigen; IFN, interferon; IL, interleukin; LILRB2, leukocyte immunoglobulin-like receptor B2; KIR, killer cell immunoglobulin-like receptors; KS, Kaposi's sarcoma; MBL, mannose-binding lectin; MHC, major histocompatibility complex; MICA, MHC class I polypeptide-related sequence A; ORF, open reading frame; *PARD3B*, par-3 family cell polarity regulator beta; *PROX1*, prospero homeobox 1; *PSORS1C3*, psoriasis susceptibility 1 candidate 3; SNP, single nucleotide polymorphism; rs#, SNP identification number in SNP database from NCBI; UTR, untranslated region; ZNRD1, zinc ribbon domain containing 1; +, present, –, absent.

Sources: Sunil K Ahuja, MD, Weijing He, MD, and Gabriel Catano, MD. Reviews for additional information: P An et al: *Trends Genet* 26:119, 2010; J Fellay: *Antivir Ther* 14:731, 2009; RA Kaslow et al: *J Infect Dis* 191:568, 2005; D van Manen et al: *Retrovirology* 9:70, 2012; MP Martin et al: *Immunol Rev* 254:245, 2013; S Limou et al: *Front Immunol* 4:118, 2013.

These SNPs are within or in the vicinity of *HCP5*, *HLA-C*, *MICA*, and *PSORS1C3* genes. The protective effects of the SNPs in *HCP5* and *MICA* may relate to their linkage with known protective *HLA-B* alleles. The protective *HCP5* allele is in linkage disequilibrium with the *HLA-B*57:01*, and the protective *MICA* allele tags with the *HLA-B*57:01* and *B*27:05* alleles. The protective *HLA-C* SNP is associated with higher *HLA-C* expression, and this effect is thought to be due to the altered binding of a microRNA to the *HLA-C* mRNA. Higher *HLA-C* expression has been associated with beneficial HIV phenotypes. The mechanism associated with the SNP in *PSORS1C3* is unknown. GWAS in African Americans identified a SNP that tags the *HLA-B*57:03* allele that is known to associate with lower virologic setpoint and slower disease course. Together these GWAS data underscore the importance of variations in *HLA class I* loci in control of viral replication.

GENETIC ASSOCIATIONS WITH SPECIFIC AIDS AND NON-AIDS CONDITIONS • Carotid artery disease Many of the non-AIDS events in HIV-infected individuals resemble those related to immune senescence and those found in the HIV-uninfected aging population. A functional SNP in the ryanodine receptor 3 (*RYR3*) gene was found to be associated with an increased risk of common carotid intima-media thickness (cIMT), which is a surrogate for subclinical atherosclerosis. Functional studies on *RYR3* and its isoforms demonstrate a major role of these receptors in modulating endothelial function and atherogenesis via calcium signaling pathways, providing a biologically plausible mechanism by which the SNP in *RYR3* may associate with increased cIMT risk.

Renal disease HIV-1-associated nephropathy (HIVAN) is a form of focal sclerosing glomerulonephritis caused by direct infection of kidney epithelial cells with HIV. HIVAN is more common in persons of African descent. There is evidence that polymorphisms in the *MYH9* gene and in the neighboring *APOL1* gene are a strong determinant of susceptibility to HIVAN in African Americans. The effect of carrying two *APOL1* risk alleles explains nearly 35% of HIVAN.

The mechanisms by which *MYH9/APOL1* variants predispose to HIVAN are currently unknown.

HIV-associated neurocognitive disorder HIV-associated neurocognitive disorder (HAND) comprises a spectrum of neurocognitive deficits due to HIV infection. Variations in the apolipoprotein E (*ApoE*) gene have strong associations with Alzheimer's disease in the HIV-uninfected population. In HIV-infected persons, possession of the *ApoE4* allele has been associated with several cognitive outcomes including dementia, peripheral neuropathy, and impairment in cognition and immediate and delayed verbal memory. Macrophage recruitment and activation plays a central role in the development of many of the HAND syndromes. Variations in chemokines that play an influential role in macrophage activation and recruitment, namely *CCL2* (MCP-1) and *CCL3* (MIP-1α), have been shown to alter the risk of developing HAND. Variations in mitochondrial genes also have been associated with risk of AIDS and HAND.

ASSOCIATIONS WITH ART-RELATED ADVERSE EVENTS Abacavir, an effective antiretroviral agent, is associated with significant risk of hypersensitivity reactions (2–9% of cases). Interestingly, while the *HLA-B*57:01* allele is associated with a slower HIV disease course, possession of this allele is associated with a higher risk of abacavir-associated hypersensitivity. Pharmacogenetic screening for the *HLA-B*57:01* allele is recommended before initiation of abacavir treatment.

NEUROPATHOGENESIS IN HIV DISEASE

While there has been a remarkable decrease in the incidence in the severe forms of HIV encephalopathy among those with access to treatment in the era of effective cART, HIV-infected individuals can still experience milder forms of neurocognitive impairment despite adequate cART. A variety of factors may contribute to the neurocognitive decline, which includes lack of complete control of HIV replication in the brain, production of HIV proteins that may be neurotoxic, low CD4+ T cell nadir, chronic immune activation, comorbidities such as

1242 drug abuse, and the potential for neurotoxicity of certain of the anti-retroviral drugs. HIV has been demonstrated in the brain and CSF of infected individuals with and without neuropsychiatric abnormalities. As opposed to lymphoid tissues, there are no resident lymphocytes in the brain. The main cell types that are infected in the brain in vivo are the perivascular macrophages and the microglial cells; low-level viral replication is also seen in perivascular astrocytes. Monocytes that have already been infected in the blood can migrate into the brain, where they then reside as macrophages, or macrophages can be directly infected within the brain. The precise mechanisms whereby HIV enters the brain are unclear; however, they are thought to relate, at least in part, to the ability of virus-infected and immune-activated macrophages to induce adhesion molecules such as E-selectin and vascular cell adhesion molecule 1 (VCAM-1) on brain endothelium. Other studies have demonstrated that HIV gp120 enhances the expression of intercellular adhesion molecule 1 (ICAM-1) in glial cells; this effect may facilitate entry of HIV-infected cells into the CNS. Virus isolates from the brain are preferentially R5 strains as opposed to X4 strains; in this regard, HIV-infected individuals who are heterozygous for *CCR5-Δ32* appear to be relatively protected against the development of HIV encephalopathy. Once HIV enters the brain due to pressures of the local environment, it evolves to develop distinct sequences in the *env*, *tat*, and *LTR* genes. These unique sequences have been associated with neurocognitive dysfunction; however, it is unclear if they are causal (see below).

HIV-infected individuals may manifest white matter lesions as well as neuronal loss. The white matter lesions are due to axonal injury and a disruption of the blood-brain barrier and not due to demyelination. Given the absence of evidence of HIV infection of neurons either in vivo or in vitro, it is highly unlikely that direct infection of these cells accounts for their loss. Rather, the HIV-mediated effects on neurons are thought to involve indirect pathways whereby viral proteins, particularly gp120 and Tat, trigger the release of endogenous neurotoxins from macrophages and to a lesser extent from astrocytes. In addition, it has been demonstrated that both HIV-1 Nef and Tat can induce chemotaxis of leukocytes, including monocytes, into the CNS. Neurotoxins can be released from monocytes as a consequence of infection and/or immune activation. Monocyte-derived neurotoxic factors have been reported to kill neurons via the *N*-methyl-D-aspartate (NMDA) receptor. In addition, HIV gp120 shed by virus-infected monocytes could cause neurotoxicity by antagonizing the function of vasoactive intestinal peptide (VIP), by elevating intracellular calcium levels, and by decreasing nerve growth factor levels in the cerebral cortex. A variety of monocyte-derived cytokines can contribute directly or indirectly to the neurotoxic effects in HIV infection; these include TNF- α , IL-1, IL-6, TGF- β , IFN- γ , platelet-activating factor, and endothelin. Furthermore, among the CC-chemokines, elevated levels of monocyte chemoattractant protein-1 (MCP-1 or CCL-2) in the brain and CSF have been shown to correlate best with the presence and degree of HIV encephalopathy. In addition, infection and/or activation of monocyte-lineage cells can result in increased production of eicosanoids, quinolinic acid, nitric oxide, excitatory amino acids such as L-cysteine and glutamate, arachidonic acid, platelet activating factor, free radicals, TNF- α , and TGF- β , which may contribute to neurotoxicity. Astrocytes may play diverse roles in HIV neuropathogenesis. Reactive gliosis or astrocytosis has been demonstrated in the brains of HIV-infected individuals, and TNF- α and IL-6 have been shown to induce astrocyte proliferation. In addition, astrocyte-derived IL-6 can induce HIV expression in infected cells in vitro. Furthermore, it has been suggested that astrocytes may downregulate macrophage-produced neurotoxins. Treatment with cART leads to improvement in neuropsychiatric manifestations and a decrease in these cytokine levels in CSF, suggesting that they are driven by the virus or by its products. However, even in patients on long-term cART, there may be evidence of persistently activated lymphocytes in the CSF. It is unclear if these lymphocytes may contribute to neuronal injury in the brain or are critical for controlling the CNS viral reservoir. The contribution of host genetic factors to development of neuropsychiatric manifestations of HIV infection has not been well studied. However, evidence supports

the role of the E4 allele for apoE in an increased risk of HIV-associated neurocognitive disorders and peripheral neuropathy.

It has also been suggested that the CNS may serve as a relatively sequestered site for a reservoir of latently infected cells that might be a barrier for the eradication of virus by cART (see “Reservoirs of HIV-Infected Cells: Obstacles to the Eradication of Virus,” above).

PATHOGENESIS OF KAPOSI'S SARCOMA

There are at least four distinct epidemiologic forms of KS: (1) the classic form that occurs in older men of predominantly Mediterranean or eastern European Jewish backgrounds with no recognized contributing factors; (2) the equatorial African form that occurs in all ages, also without any recognized precipitating factors; (3) the form associated with organ transplantation and its attendant iatrogenic immunosuppressed state; and (4) the form associated with HIV-1 infection. In the latter two forms, KS is an opportunistic disease; in HIV-infected individuals, unlike typical opportunistic infections, its occurrence is not strictly related to the level of depression of CD4+ T cell counts. The pathogenesis of KS is complex; fundamentally, it is an angioproliferative disease that is not a true neoplastic sarcoma, at least not in its early stages. It is a manifestation of excessive proliferation of spindle cells that are believed to be of vascular origin and have features in common with endothelial and smooth-muscle cells. In HIV disease the development of KS is dependent on the interplay of a variety of factors including HIV-1 itself, human herpes virus 8 (HHV-8), immune activation, and cytokine secretion. A number of epidemiologic and virologic studies have clearly linked HHV-8, which is also referred to as *Kaposi's sarcoma-associated herpesvirus* (KSHV), to KS not only in HIV-infected individuals but also in individuals with the other forms of KS. HHV-8 is a γ -herpesvirus related to EBV and *herpesvirus saimiri*. It encodes a homologue to human IL-6 and, in addition to KS, has been implicated in the pathogenesis of body cavity lymphoma, multiple myeloma, and monoclonal gammopathy of undetermined significance. Sequences of HHV-8 are found universally in the lesions of KS, and patients with KS are virtually all seropositive for HHV-8. HHV-8 DNA sequences can be found in the B cells of 30–50% of patients with KS and 7% of patients with AIDS without clinically apparent KS.

Between 1 and 2% of eligible blood donors are positive for antibodies to HHV-8, while the prevalence of HHV-8 seropositivity in HIV-infected men is 30–35%. The prevalence of HHV-8 seropositivity in HIV-infected women is ~4%. This finding is reflective of the lower incidence of KS in women. It has been debated whether HHV-8 is actually the transforming agent in KS; the bulk of the cells in the tumor lesions of KS are not neoplastic cells. However, it has been demonstrated that endothelial cells can be transformed in vitro by HHV-8. In this regard, HHV-8 possesses a number of genes, including homologues of the IL-8 receptor, Bcl-2, and cyclin D, that can potentially transform the host cell. Despite the complexity of the pathogenic events associated with the development of KS in HIV-infected individuals, HHV-8 is the etiologic agent of this disease. The initiation and/or propagation of KS requires an activated state and is mediated, at least in part, by cytokines. A number of factors, including TNF- α , IL-1 β , IL-6, granulocyte-macrophage colony-stimulating factor (GM-CSF), basic fibroblast growth factor, and oncostatin M, function in an autocrine and paracrine manner to sustain the growth and chemotaxis of the KS spindle cells. In this regard, KSHV-derived IL-6 has been demonstrated to induce proliferation of lymphoma cells and to inhibit the cytostatic effects of IFN- α on KSHV-infected lymphoma cells.

IMMUNE RESPONSE TO HIV

As detailed above and below, following the initial burst of viremia during primary infection, HIV-infected individuals mount robust immune responses that in most cases substantially curtail the levels of plasma viremia and likely contribute to delaying the ultimate development of clinically apparent disease for a median of 10 years in untreated individuals. This immune response contains elements of both humoral and cell-mediated immunity involving both innate and adaptive immune responses (Table 226-7; Fig. 226-26). It is directed against multiple antigenic determinants of the HIV virion as

TABLE 226-7 ELEMENTS OF THE IMMUNE RESPONSE TO HIV

Humoral immunity
Binding antibodies
Neutralizing antibodies
Type specific
Group specific
Broadly neutralizing
Antibodies participating in antibody-dependent cellular cytotoxicity (ADCC)
Protective
Pathogenic (bystander killing)
Enhancing antibodies
Complement
Cell-mediated immunity
Helper CD4+ T lymphocytes
Class I MHC-restricted cytotoxic CD8+ T lymphocytes
CD8+ T cell-mediated inhibition (noncytolytic)
ADCC
Natural killer cells

Abbreviation: MHC, major histocompatibility complex.

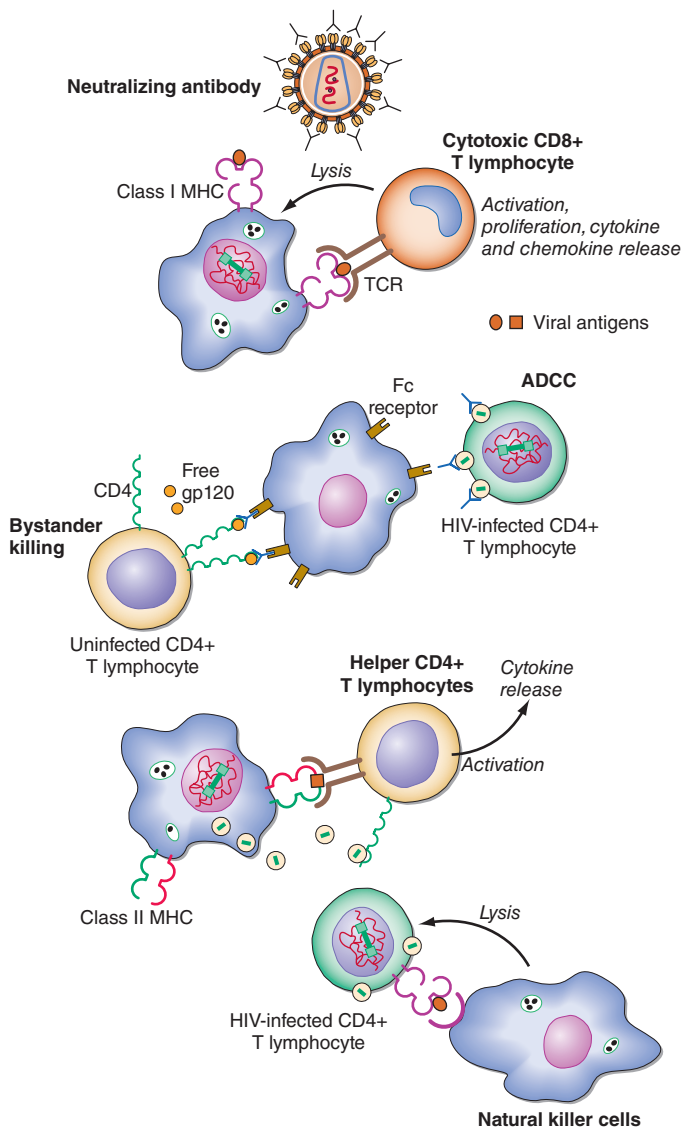


FIGURE 226-26 Schematic representation of the different immunologic effector mechanisms thought to be active in the setting of HIV infection. Detailed descriptions are given in the text. ADCC, antibody-dependent cellular cytotoxicity; MHC, major histocompatibility complex; TCR, T cell receptor.

well as against viral proteins expressed on the surface of infected cells. Ironically, those CD4+ T cells with T cell receptors specific for HIV are theoretically those CD4+ T cells most likely to be activated—and thus to serve as early targets for productive HIV infection and the cell death or dysfunction associated with infection. Thus, an early consequence of HIV infection is interference with and decrease of the helper T cell population needed to generate an effective immune response.

Although a great deal of investigation has been directed toward delineating and better understanding the components of this immune response, it remains unclear which immunologic effector mechanisms are most important in delaying progression of infection and which, if any, play a role in the pathogenesis of HIV disease. This lack of knowledge has also hampered the ability to develop an effective vaccine for HIV disease.

HUMORAL IMMUNE RESPONSE

Antibodies to HIV usually appear within 3–6 weeks and almost invariably within 12 weeks of primary infection (**Fig. 226-27**); rare exceptions are in individuals who have defects in the ability to produce HIV-specific antibodies. Detection of these antibodies forms the basis of most diagnostic screening tests for HIV infection. The appearance of HIV-binding antibodies detected by ELISA and Western blot assays occurs prior to the appearance of neutralizing antibodies; the latter generally appear following the initial decreases in plasma viremia and are more closely related to the appearance of HIV-specific CD8+ T lymphocytes. The first antibodies detected are those directed against the immunodominant region of the envelope gp41, followed by the appearance of antibodies to the structural or gag protein p24 and the gag precursor p55. Antibodies to p24 gag are followed by the appearance of antibodies to the outer envelope glycoprotein (gp120), the gag protein p17, and the products of the *pol* gene (p31 and p66). In addition, one may see antibodies to the low-molecular-weight regulatory proteins encoded by the HIV genes *vpr*, *vpu*, *vif*, *rev*, *tat*, and *nef*. On rare occasion, levels of HIV-specific antibodies may decline during treatment of acute HIV infection.

While antibodies to multiple antigens of HIV are produced, the precise functional significance of these different antibodies is unclear. The only viral proteins that elicit neutralizing antibodies are the envelope proteins gp120 and gp41. Antibodies directed toward the envelope proteins of HIV have been characterized both as being protective and as possibly contributing to the pathogenesis of HIV disease. Among the protective antibodies are those that function to neutralize HIV directly and prevent the spread of infection to additional cells, as well as those that participate in ADCC. The first neutralizing antibodies are directed against the autologous infecting

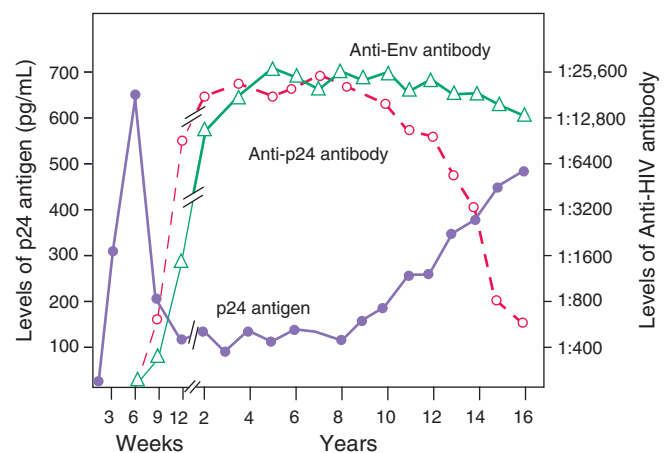


FIGURE 226-27 Relationship between antigenemia and the development of antibodies to HIV. Levels of plasma HIV parallel those of p24 antigen. Antibodies to HIV proteins are generally seen 6–12 weeks following infection and 3–6 weeks after the development of plasma viremia. Late in the course of illness, antibody levels to p24 decline, generally in association with a rising titer of p24 antigen.

1244 virus and appear after approximately 12 weeks of infection. Due to its high rate of mutation the virus is usually able to quickly escape these (and subsequent) neutralizing antibodies. One important mechanism of immune escape is the addition of N-linked glycosylation sites, forming a glycan shield that interferes with envelope recognition by these initial antibodies.

A number of broad and potent HIV-neutralizing envelope-specific antibodies have been isolated from HIV-infected individuals in studies designed to better understand the host response to HIV infection. Approximately 20% of patients develop antibodies capable of neutralizing highly diverse strains. These studies have revealed at least five major sites within the HIV envelope that are able to elicit broadly-neutralizing antibodies. These sites include antibodies directed toward the CD4 binding site (CD4bs) of gp120, those binding glycan-dependent epitopes in the V1/V2 region of gp120, those near the base of the V3 region of gp120, those binding to the gp120/gp41 bridge, and those binding to the membrane-proximal region of gp41 (Fig. 226-28). Several of these antibodies contain unique features including high levels of somatic hypermutation, selective germline gene usage (especially for CD4bs antibodies) and long heavy chain complementary determining regions (especially CDRH3). Of note, while these antibodies are broadly neutralizing in vitro, their precise in vivo significance is unclear and the patients from whom they were derived demonstrate evidence of ongoing viral replication unless treated with cART.

The other major class of protective antibodies are those that participate in ADCC, a form of cell-mediated immunity (Chap. 372e) in which NK cells that bear Fc receptors are armed with specific anti-HIV antibodies that bind to the NK cells via their Fc portion. These armed NK cells then bind to and destroy cells expressing HIV antigens. The levels of anti-envelope antibodies capable of mediating ADCC are highest in the earlier stages of HIV infection. Antibodies to both gp120 and gp41 have been shown to participate in ADCC-mediated killing of HIV-infected cells. In vitro, IL-2 can augment ADCC-mediated killing.

In addition to playing a role in host defense, HIV-specific antibodies have also been implicated in disease pathogenesis. Antibodies directed to gp41, when present in low titer, have been shown in vitro to be capable of facilitating infection of cells through an Fc receptor-mediated mechanism known as *antibody enhancement*. Thus, the same regions of the envelope protein of HIV that give rise to antibodies capable of mediating ADCC can also elicit the production of antibodies that can facilitate infection of cells in vitro. In addition, it has been postulated that anti-gp120 antibodies that participate in the ADCC killing of HIV-infected cells might also kill uninfected CD4+ T cells if the uninfected cells had bound free gp120, a phenomenon referred to as *bystander killing*.

One of the most primitive components of the humoral immune system is the complement system (Chap. 372e). This element of innate immunity consists of ~30 proteins that are found circulating in blood or associated with cell membranes. While HIV alone is capable of directly activating the complement cascade, the resulting lysis is

weak due to the presence of host cell regulatory proteins captured in the virion envelope during budding. It is possible that complement-opsonized HIV virions have increased infectivity in a manner analogous to antibody-mediated enhancement.

CELLULAR IMMUNE RESPONSE

Given that T cell-mediated immunity is known to play a major role in host defense against most viral infections (Chap. 372e), it is generally thought to be an important component of the host immune response to HIV. T cell immunity can be divided into two major categories: that mediated by *helper/inducer CD4+ T cells* and that mediated by *cytotoxic/immunoregulatory CD8+ T cells*.

HIV-specific CD4+ T cells can be detected in the majority of HIV-infected patients through the use of flow cytometry to measure intracellular cytokine production in response to MHC class II tetramers pulsed with HIV peptides or through lymphocyte proliferation assays utilizing HIV antigens such as p24. These cells likely play a critical role in the orchestration of the immune response to HIV by providing help to HIV-specific B cells and CD8+ T cells. They may also be capable of directly killing HIV-infected cells. HIV-specific CD4+ T cells may be preferential targets of HIV infection by HIV-infected antigen-presenting cells during the generation of an immune response to HIV (Fig. 226-26). However, they also are likely to undergo clonal expansions in response to HIV antigens and thus survive as a population of cells. No clear correlations exist between levels of HIV-specific CD4+ T lymphocytes and plasma HIV RNA levels; however, in the setting of high viral loads, CD4+ T cell responses to HIV antigens appear to shift from one of proliferation and IL-2 production to one of IFN- γ production. Thus, while a reverse correlation exists between the level of p24-specific proliferation and levels of plasma HIV viremia, the nature of the causal relationship between these parameters is unclear.

MHC class I-restricted, HIV-specific CD8+ T cells have been identified in the peripheral blood of patients with HIV-1 infection. These cells include CTLs that produce perforins and T cells that can be induced by HIV antigens to express an array of cytokines such as IFN- γ , IL-2, MIP-1 β , and TNF- α . CTLs have been identified in the peripheral blood of patients within weeks of HIV infection and prior to the appearance of plasma virus. The selective pressure they exert on the evolution of the population of circulating viruses reflects their potential role in control of HIV infection. These CD8+ T lymphocytes, through their HIV-specific antigen receptors, bind to and cause the lytic destruction of target cells bearing autologous MHC class I molecules presenting HIV antigens. Two types of CTL activity can be demonstrated in the peripheral blood or lymph node mononuclear cells of HIV-infected individuals. The first type directly lyses appropriate target cells in culture without prior in vitro stimulation (*spontaneous CTL activity*). The other type of CTL activity reflects the *precursor frequency of CTLs* (CTLp); this type of CTL activity can be demonstrated by stimulation of CD8+ T cells in vitro with a mitogen such as phytohemagglutinin or anti-CD3 antibody.

In addition to CTLs, CD8+ T cells capable of being induced by HIV antigens to express cytokines such as IFN- γ also appear in the setting of HIV-1 infection. It is not clear whether these are the same or different effector pools compared with those cells mediating cytotoxicity; in addition, the relative roles of each in host defense against HIV are not fully understood. It does appear that these CD8+ T cells are driven to in vivo expansion by HIV antigen. There is a direct correlation between levels of CD8+ T cells capable of producing IFN- γ in response to HIV antigens and plasma levels of HIV-1 RNA. Thus, while these cells are clearly induced by HIV-1 infection, their overall ability to control infection remains unclear. Multiple HIV antigens, including Gag, Env, Pol, Tat, Rev, and Nef, can elicit CD8+ T cell responses. Among patients who control viral replication in the absence of antiretroviral drugs are a subset of patients referred to as elite nonprogressors (see "Long-Term Survivors and Long-Term Nonprogressors," above) whose peripheral blood contains a population of CD8+ T cells that undergo substantial proliferation and perforin expression in response to HIV antigens. It is possible that these cells play an important role in HIV-specific host defense.

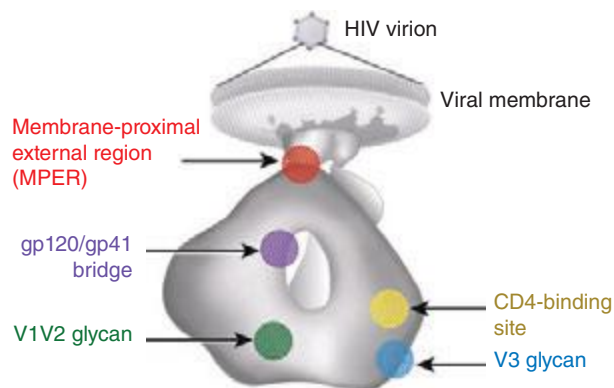


FIGURE 226-28 Known targets of broadly neutralizing antibodies against HIV-1. (Adapted from PD Kwong, JR Mascola: *Immunity* 37:412, 2012.)

At least three other forms of cell-mediated immunity to HIV have been described: non-cytolytic CD8+ T cell-mediated suppression of HIV replication, ADCC, and NK cell activity. *Non-cytolytic CD8+ T cell-mediated suppression of HIV replication* refers to the ability of CD8+ T cells from an HIV-infected patient to inhibit the replication of HIV in tissue culture without killing infected targets. There is no requirement for HLA compatibility between the CD8+ T cells and the HIV-infected cells. This effector mechanism is thus nonspecific and appears to be mediated by soluble factor(s) including the CC-chemokines RANTES (CCL5), MIP-1 α (CCL3), and MIP-1 β (CCL4). These CC-chemokines are potent suppressors of HIV replication and operate at least in part via blockade of the HIV co-receptor (CCR5) for R5 (macrophage-tropic) strains of HIV-1 (see above). ADCC, as described above in relation to humoral immunity, involves the killing of HIV-expressing cells by NK cells armed with specific antibodies directed against HIV antigens. Finally, *NK cells* alone have been shown to be capable of killing HIV-infected target cells in tissue culture. This primitive cytotoxic mechanism of host defense is directed toward nonspecific surveillance for neoplastic transformation and viral infection through recognition of altered class I MHC molecules.

DIAGNOSIS AND LABORATORY MONITORING OF HIV INFECTION

The establishment of HIV as the causative agent of AIDS and related syndromes early in 1984 was followed by the rapid development of sensitive screening tests for HIV infection. By March 1985, blood donors in the United States were routinely screened for antibodies to HIV. In 1996, blood banks in the United States added the p24 antigen capture assay to the screening process to help identify the rare infected individuals who were donating blood in the time (up to 3 months) between infection and the development of antibodies. In 2002, the ability to detect early infection with HIV was further enhanced by the licensure of nucleic acid testing (NAT) as a routine part of blood donor screening. These refinements decreased the interval between infection and detection (window period) from 22 days for antibody testing to 16 days with p24 antigen testing and subsequently to 12 days with NAT. The development of sensitive assays for monitoring levels of plasma viremia ushered in a new era of being able to monitor the progression of HIV disease more closely. Utilization of these tests, coupled with the measurement of levels of CD4+ T lymphocytes in peripheral blood, is essential in the management of patients with HIV infection.

DIAGNOSIS OF HIV INFECTION

The CDC has recommended that screening for HIV infection be performed as a matter of routine health care. The diagnosis of HIV infection depends on the demonstration of antibodies to HIV and/or the direct detection of HIV or one of its components. As noted above, antibodies to HIV generally appear in the circulation 3–12 weeks following infection.

The standard blood screening tests for HIV infection are based on the detection of antibodies to HIV. A common platform is the ELISA, also referred to as an *enzyme immunoassay* (EIA). This solid-phase assay is an extremely good screening test with a sensitivity of >99.5%. Most diagnostic laboratories use commercial kits that contain antigens from both HIV-1 and HIV-2 and thus are able to detect antibodies to either. These kits use both natural and recombinant antigens and are continuously updated to increase their sensitivity to newly discovered species, such as group O viruses (Fig. 226-1). The fourth-generation EIA tests combine detection of antibodies to HIV with detection of the p24 antigen of HIV. EIA tests are generally scored as positive (highly reactive), negative (nonreactive), or indeterminate (partially reactive). While the EIA is an extremely sensitive test, it is not optimal with regard to specificity. This is particularly true in studies of low-risk individuals, such as volunteer blood donors. In this latter population, only 10% of EIA-positive individuals are subsequently confirmed to have HIV infection. Among the factors associated with false-positive EIA tests are antibodies to class II antigens (such as may be seen following pregnancy, blood transfusion, or transplantation), autoantibodies, hepatic disease, recent influenza vaccination, and acute viral infections. For these reasons, anyone suspected of having

HIV infection based on a positive or inconclusive EIA result should ideally have the result confirmed with a more specific assay such as the Western blot. One can estimate whether an individual has a recent infection with HIV-1 by comparing the results on a standard EIA test that will score positive for all infected individuals with the results on an assay modified to be less sensitive (“detuned assay”) that will score positive for individuals with established HIV infection and negative for individuals with recent infection. In rare instances, an HIV-infected individual treated early in the course of infection may revert to a negative EIA. This does *not* indicate clearing of infection; rather, it signifies levels of ongoing exposure to virus or viral proteins insufficient to maintain a measurable antibody response. When these individuals have discontinued therapy, viruses and antibodies have reappeared.

The most commonly used confirmatory test is the Western blot (Fig. 226-29). This assay takes advantage of the fact that multiple HIV antigens of different, well-characterized molecular weights elicit the production of specific antibodies. These antigens can be separated on the basis of molecular weight, and antibodies to each component can be detected as discrete bands on the Western blot. A negative Western blot is one in which no bands are present at molecular weights corresponding to HIV gene products. In a patient with a positive or indeterminate EIA and a negative Western blot, one can conclude with certainty that the EIA reactivity was a false positive. On the other hand, a Western blot demonstrating antibodies to products of all three of the major genes of HIV (*gag*, *pol*, and *env*) is conclusive evidence of infection with HIV. Criteria established by the U.S. Food and Drug Administration (FDA) in 1993 for a positive Western blot state that a result is considered positive if antibodies exist to two of the three HIV proteins: p24, gp41, and gp120/160. Using these criteria, ~10% of all blood donors deemed positive for HIV-1 infection lacked an antibody band to the *pol* gene product p31. Some 50% of these blood donors were subsequently found to be false positives. Thus, the absence of the p31 band should increase the suspicion that one may be dealing with a false-positive test result. In this setting it is prudent to obtain additional confirmation with an RNA-based test for HIV-1 and/or a follow-up Western blot. By definition, Western blot patterns of reactivity that do not fall into the positive or negative categories are considered “indeterminate.” There are two possible explanations for an indeterminate Western blot result. The most likely explanation in a low-risk individual is that the patient being tested has antibodies that cross-react with one of the proteins of HIV. The most common patterns of cross-reactivity are antibodies that react with p24 and/or p55. The least likely explanation in this setting is that the individual is infected with HIV and is in the process of mounting a classic antibody response. In either instance, the Western blot should be repeated in 1 month to determine whether the indeterminate pattern is a pattern in evolution. In addition, one may attempt to confirm a diagnosis of HIV infection with the p24 antigen capture assay or one of the tests for HIV RNA (discussed below). While the Western blot is an excellent confirmatory test for HIV infection in patients with a positive or indeterminate EIA, it is a poor screening test. Among individuals with a negative EIA and PCR for HIV, 20–30% may show one or more bands on Western blot. While these bands are usually faint and represent cross-reactivity, their presence creates a situation in which other diagnostic modalities (such as DNA PCR, RT-PCR, or p24 antigen capture) must be employed to ensure that the bands do not indicate early HIV infection.

A guideline for the use of these serologic tests in attempting to make a diagnosis of HIV infection is depicted in Fig. 226-30. In patients in whom HIV infection is suspected, the appropriate initial test is the EIA. If the result is negative, unless there is strong reason to suspect early HIV infection (as in a patient exposed within the previous 3 months), the diagnosis is ruled out and retesting should be performed only as clinically indicated. If the EIA is indeterminate or positive, the test should be repeated. If the repeat is negative on two occasions, one can assume that the initial positive reading was due to a technical error in the performance of the assay and that the patient is negative. If the repeat is indeterminate or positive, one should proceed to the HIV-1 Western blot. If the Western blot is positive, the diagnosis is HIV-1

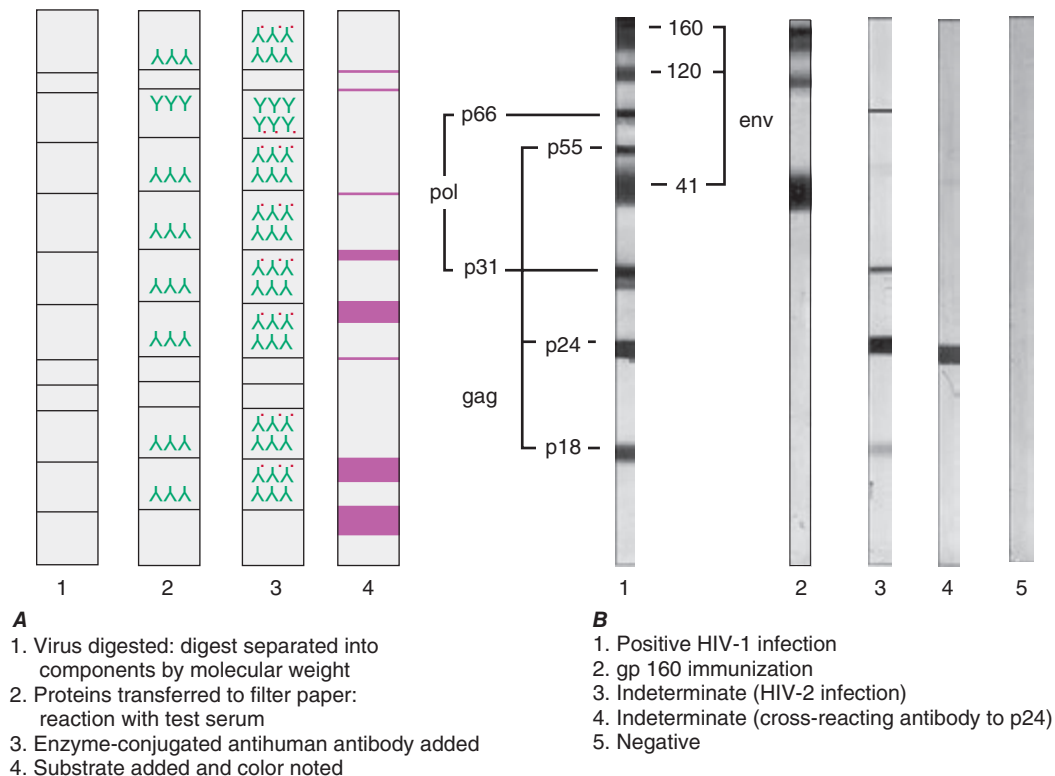


FIGURE 226-29 Western blot assay for detection of antibodies to HIV. **A.** Schematic representation of how a Western blot is performed. **B.** Examples of patterns of Western blot reactivity. In each instance the Western blot strip contains antigens to HIV-1. The serum from the patient immunized to the HIV-1 envelope gp160 contains only antibodies to the HIV-1 envelope proteins. The serum from the patient with HIV-2 infection cross-reacts with both *reverse transcriptase* and *gag* gene products of HIV-1.

infection. If the Western blot is negative, the EIA can be assumed to have been a false positive for HIV-1 and the diagnosis of HIV-1 infection is ruled out. It would also be prudent at this point to perform specific serologic testing for HIV-2 following the same type of algorithm. If the Western blot for HIV-1 is indeterminate, it should be repeated in 4–6 weeks; in addition, one may proceed to a p24 antigen capture assay, HIV-1 RNA assay, or HIV-1 DNA PCR and specific serologic testing for HIV-2. If the p24 and HIV RNA assays are negative and

there is no progression in the Western blot, a diagnosis of HIV-1 is ruled out. If either the p24 or HIV-1 RNA assay is positive and/or the HIV-1 Western blot shows progression, a tentative diagnosis of HIV-1 infection can be made and later confirmed with a follow-up Western blot demonstrating a positive pattern. In addition to these standard laboratory-based assays for detecting antibodies to HIV, an series of point-of-care tests can provide results in 1–60 min. Among the most popular of these is the OraQuick Rapid HIV-1 antibody test that can be run on blood, plasma, or saliva. The sensitivity and specificity of this test is ~99% when run on whole blood. Specificity remains the same but sensitivity drops to 98% when the test is run on saliva. While negative results from this test are adequate to rule out a diagnosis of HIV infection, a positive finding should be considered preliminary and confirmed with standard serologic testing, as described above. Two rapid test kits are licensed for home use. They are the OraQuick In-Home HIV test and the Home Access HIV-1 test system.

A variety of laboratory tests are available for the direct detection of HIV or its components (Table 226-8). These tests may be of considerable help in making a diagnosis of HIV infection when the Western blot results are indeterminate. In addition, the tests detecting levels of HIV RNA can be used to determine prognosis and to assess the response to antiretroviral therapies. The simplest of the direct detection tests is the *p24 antigen capture assay*. This is an EIA-type assay in which the solid phase consists of antibodies to the p24 antigen of HIV. It detects the viral protein p24 in the blood of HIV-infected individuals where it exists either as free antigen or complexed to anti-p24 antibodies. Overall, ~30% of individuals with untreated HIV infection have detectable levels of free p24 antigen. This increases to ~50% when samples are treated with a weak acid to dissociate antigen-antibody complexes. Throughout the course of HIV infection, an equilibrium exists between p24 antigen and anti-p24 antibodies. During the first few weeks of infection, before an immune response develops, there is a brisk rise in p24 antigen levels (Fig. 226-27). After the development of anti-p24 antibodies, these levels decline. Late in the course of infection, when circulating levels of virus are high, p24 antigen levels

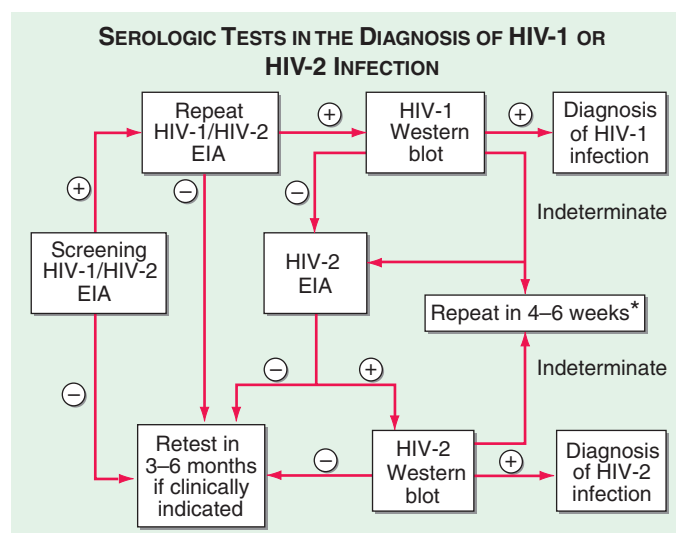


FIGURE 226-30 Algorithm for the use of serologic tests in the diagnosis of HIV-1 or HIV-2 infection. *Stable indeterminate Western blot 4–6 weeks later makes HIV infection unlikely. However, it should be repeated twice at 3-month intervals to rule out HIV infection. Alternatively, one may test for HIV-1 p24 antigen or HIV RNA. EIA, enzyme immunoassay.

TABLE 226-8 CHARACTERISTICS OF TESTS FOR DIRECT DETECTION OF HIV

Test	Technique	Sensitivity ^a	Cost/Test ^b
Immune complex–dissociated p24 antigen capture assay	Measurement of levels of HIV-1 core protein in an EIA-based format following dissociation of antigen-antibody complexes by weak acid treatment	Positive in 50% of patients; detects down to 15 pg/mL of p24 protein	\$1–2
HIV RNA by PCR	Target amplification of HIV-1 RNA via reverse transcription followed by PCR	Reliable to 40 copies/mL of HIV RNA	\$75–150
HIV RNA by bDNA	Measurement of levels of particle-associated HIV RNA in a nucleic acid capture assay employing signal amplification	Reliable to 50 copies/mL of HIV RNA	\$75–150
HIV RNA by TMA	Target amplification of HIV-1 RNA via reverse transcription followed by T7 RNA polymerase	Reliable to 100 copies/mL of HIV RNA	\$225
HIV RNA by NASBA	Isothermal nucleic acid amplification with internal controls	Reliable to 80 copies/mL of HIV RNA	\$75–150

^aSensitivity figures refer to those approved by the U.S. Food and Drug Administration. ^bPrices may be lower in large-volume settings.

Abbreviations: bDNA, branched DNA; cDNA, complementary DNA; EIA, enzyme immunoassay; TMA, transcription-mediated amplification; NASBA, nucleic acid sequence–based amplification; PCR, polymerase chain reaction.

also increase, particularly when detected by techniques involving dissociation of antigen-antibody complexes. The p24 antigen capture assay has its greatest use as a screening test for HIV infection in patients suspected of having the acute HIV syndrome, as high levels of p24 antigen are present prior to the development of antibodies. Its use as a stand-alone test for routine blood donor screening for HIV infection has been replaced by use of NAT or “fourth-generation” assays that combine antigen and antibody testing. The ability to measure and monitor levels of HIV RNA in the plasma of patients with HIV infection has been of extraordinary value in furthering our understanding of the pathogenesis of HIV infection, in monitoring the response to cART, and in providing a diagnostic tool in settings where measurements of anti-HIV antibodies may be misleading, such as in acute infection and neonatal infection. Four assays are predominantly used for this purpose. They are reverse transcriptase PCR (RT-PCR; Amplicor); branched DNA (bDNA; VERSANT); transcription-mediated amplification (TMA; APTIMA); and nucleic acid sequence–based amplification (NASBA; NucliSENS). These tests are of value in making a diagnosis of HIV infection, in establishing initial prognosis, and in monitoring the effects of therapy. In addition to these four commercially available tests, the DNA PCR also is employed by research laboratories for making a diagnosis of HIV infection by amplifying HIV proviral DNA from peripheral blood mononuclear cells. The commercially available RNA detection tests have a sensitivity of 40–8100 copies of HIV RNA per milliliter of plasma. Research laboratory–based RNA assays can detect as few as one HIV RNA copy per milliliter, while the DNA PCR tests can detect proviral DNA at a frequency of one copy per 10,000–100,000 cells. Thus, these tests are extremely sensitive. One frequent consequence of a high degree of sensitivity is some loss of specificity, and false-positive results have been reported with each of these techniques. For this reason, a positive EIA with a confirmatory Western blot remains the “gold standard” for a diagnosis of HIV infection, and the interpretation of other test results must be done with this in mind.

In the RT-PCR technique, following DNase treatment, a cDNA copy is made of all RNA species present in plasma. Because HIV is an RNA virus, this will result in the production of DNA copies of the HIV genome in amounts proportional to the amount of HIV RNA present in plasma. This cDNA is then amplified and characterized using standard PCR techniques, employing primer pairs that can distinguish genomic cDNA from messenger cDNA. The bDNA assay involves the use of a solid-phase nucleic acid capture system and signal amplification through successive nucleic acid hybridizations to detect small quantities of HIV RNA. Both tests can achieve a tenfold increase in sensitivity to 40–50 copies of HIV RNA per milliliter with a preconcentration step in which plasma undergoes ultracentrifugation to pellet the viral particles. In the TMA assay, a cDNA copy of viral RNA is made using primers that contain a promoter sequence for T7 RNA polymerase. T7 polymerase is then added to produce multiple copies of RNA amplicon from the DNA template. It is qualified at 100 copies/mL. The NASBA technique involves the isothermal amplification of a sequence within the gag region of HIV in the presence of

internal standards and employs the production of multiple RNA copies through the action of T7-RNA polymerase. The resulting RNA species are quantitated through hybridization with a molecular beacon DNA probe that is quenched in the absence of hybridization. The lower limit of detection for the NucliSENS assay is 80 copies/mL.

In addition to being a diagnostic and prognostic tool, RT-PCR and DNA-PCR are also useful for amplifying defined areas of the HIV genome for sequence analysis and have become an important technique for studies of sequence diversity and microbial resistance to antiretroviral agents. In patients with a positive or indeterminate EIA test and an indeterminate Western blot, and in patients in whom serologic testing may be unreliable (such as patients with hypogammaglobulinemia or advanced HIV disease), these tests for quantitating HIV RNA in plasma or detecting proviral DNA in peripheral blood mononuclear cells are valuable tools for making a diagnosis of HIV infection; however, they should be used for diagnosis only when standard serologic testing has failed to provide a definitive result.

LABORATORY MONITORING OF PATIENTS WITH HIV INFECTION

The epidemic of HIV infection and AIDS has provided the clinician with new challenges for integrating clinical and laboratory data to effect optimal patient management. The close relationship between clinical manifestations of HIV infection and CD4+ T cell count has made measurement of CD4+ T cell numbers a routine part of the evaluation of HIV-infected individuals. The discovery of HIV as the cause of AIDS led to the development of sensitive tests that allow one to monitor the levels of HIV in the blood. Determinations of peripheral blood CD4+ T cell counts and measurements of the plasma levels of HIV RNA provide a powerful set of tools for determining prognosis and monitoring response to therapy.

CD4+ T Cell Counts The CD4+ T cell count is the laboratory test generally accepted as the best indicator of the immediate state of immunologic competence of the patient with HIV infection. This measurement, which can be made directly or calculated as the product of the percentage of CD4+ T cells (determined by flow cytometry) and the total lymphocyte count (determined by the white blood cell count [WBC] multiplied by the lymphocyte differential percentage), has been shown to correlate very well with the level of immunologic competence. Patients with CD4+ T cell counts <200/μL are at high risk of disease from *P. jiroveci*, while patients with CD4+ T cell counts <50/μL are at high risk of disease from CMV, mycobacteria of the *M. avium* complex (MAC), and/or *T. gondii* (Fig. 226-31). Once the CD4+ T cell count is <200/μL, patients should be placed on a regimen for *P. jiroveci* prophylaxis, and once the count is <50/μL, primary prophylaxis for MAC infection is indicated. As with any laboratory measurement, one may wish to obtain two determinations prior to any significant changes in patient management based on CD4+ T cell count alone. Patients with HIV infection should have CD4+ T cell measurements performed at the time of diagnosis and every 3–6 months thereafter. More frequent measurements should be made if a declining trend is noted. For patients who have been on cART for at least 2 years

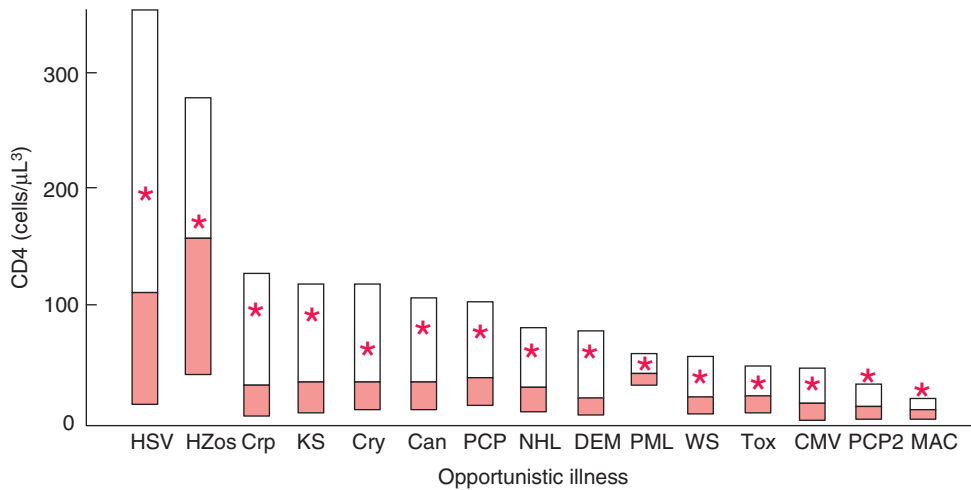


FIGURE 226-31 Relationship between CD4+ T cell counts and the development of opportunistic diseases. Boxplot of the median (line inside the box), first quartile (bottom of the box), third quartile (top of the box), and mean (asterisk) CD4+ lymphocyte count at the time of the development of opportunistic disease. Can, candidal esophagitis; CMV, cytomegalovirus infection; Crp, cryptosporidiosis; Cry, cryptococcal meningitis; DEM, AIDS dementia complex; HSV, herpes simplex virus infection; HZos, herpes zoster; KS, Kaposi's sarcoma; MAC, *Mycobacterium avium* complex bacteremia; NHL, non-Hodgkin's lymphoma; PCP, primary *Pneumocystis jiroveci* pneumonia; PCP2, secondary *P. jiroveci* pneumonia; PML, progressive multifocal leukoencephalopathy; Tox, *Toxoplasma gondii* encephalitis; WS, wasting syndrome. (From RD Moore, RE Chaisson: *Ann Intern Med* 124:633, 1996.)

with HIV RNA levels persistently <50 copies/mL, the monitoring of the CD4 count is felt by many to be optional. There are a handful of clinical situations in which the CD4+ T cell count may be misleading. Patients with HTLV-1/HIV co-infection may have elevated CD4+ T cell counts that do not accurately reflect their degree of immune competence. In patients with hypersplenism or those who have undergone splenectomy, and in patients receiving medications that suppress the bone marrow such as IFN- α , the CD4+ T cell percentage may be a more reliable indication of immune function than the CD4+ T cell count. A CD4+ T cell percentage of 15 is comparable to a CD4+ T cell count of 200/ μ L.

HIV RNA Determinations Facilitated by highly sensitive techniques for the precise quantitation of small amounts of nucleic acids, the measurement of serum or plasma levels of HIV RNA has become an essential component in the monitoring of patients with HIV infection. As discussed in "Diagnosis of HIV Infection," above, the most commonly used technique is the RT-PCR assay. This assay generates data in the form of number of copies of HIV RNA per milliliter of serum or plasma and can reliably detect as few as 40 copies of HIV RNA per milliliter of plasma. Research-based assays can detect down to one copy per milliliter. While it is common practice to describe levels of HIV RNA below these cut-offs as "undetectable," this is a term that should be avoided as it is imprecise and leaves the false impression that the level of virus is 0. By utilizing more sensitive, nested PCR techniques and by studying tissue levels of virus as well as plasma levels, HIV RNA can be detected in virtually every patient with HIV infection. The one notable exception to this is a patient who underwent cytoreductive therapy followed by a bone marrow transplant from a CCR5 Δ 32 homozygous donor.

Measurements of changes in HIV RNA levels over time have been of great value in delineating the relationship between levels of virus and rates of disease progression (Fig. 226-22), the rates of viral turnover, the relationship between immune system activation and viral replication, and the time to development of drug resistance. HIV RNA measurements are greatly influenced by the state of activation of the immune system and may fluctuate greatly in the setting of secondary infections or immunization. For these reasons, decisions based on HIV RNA levels should never be made on a single determination. Measurements of plasma HIV RNA levels should be made at the time of HIV diagnosis and every 3–6 months thereafter in the

untreated patient. Following the initiation of therapy or any change in therapy, plasma HIV RNA levels should be monitored approximately every 4 weeks until the effectiveness of the therapeutic regimen is determined by the development of a new steady-state level of HIV RNA. In most instances of effective antiretroviral therapy the plasma level of HIV RNA will drop to <50 copies/mL within 6 months of the initiation of treatment. During therapy, levels of HIV RNA should be monitored every 3–6 months to evaluate the continuing effectiveness of therapy.

HIV Resistance Testing The availability of multiple antiretroviral drugs as treatment options has generated a great deal of interest in the potential for measuring the sensitivity of an individual's HIV virus(es) to different antiretroviral agents. HIV resistance testing can be done through either genotypic or phenotypic measurements. In the genotypic assays, sequence analyses of the HIV genomes obtained from patients are compared with sequences

of viruses with known antiretroviral resistance profiles. In the phenotypic assays, the in vivo growth of viral isolates obtained from the patient is compared to the growth of reference strains of the virus in the presence or absence of different antiretroviral drugs. A modification of this phenotypic approach utilizes a comparison of the enzymatic activities of the reverse transcriptase, protease, or integrase genes obtained by molecular cloning of patients' isolates to the enzymatic activities of genes obtained from reference strains of HIV in the presence or absence of different drugs targeted to these genes. These tests are quite good in identifying those antiretroviral agents that have been utilized in the past and suggesting agents that may be of future value in a given patient. Resistance testing is recommended at the time of initial diagnosis and, if therapy is not initiated at that time, at the time of initiation of cART. Drug resistance testing is also indicated in the setting of virologic failure and should be performed while the patient is still on the failing regimen because of the propensity for the pool of HIV quasiespecies to rapidly revert to wild-type in the absence of the selective pressures of cART. In the hands of experts, resistance testing enhances the short-term ability to decrease viral load by ~0.5 log compared with changing drugs merely on the basis of drug history. In addition to the use of resistance testing to help in the selection of new drugs in patients with virologic failure, it may also be of value in selecting an initial regimen for treatment of therapy-naïve individuals. This is particularly true in geographic areas with a high level of background resistance. The patient needs to have an HIV-1 RNA level above 500–1000 copies/mL for an accurate resistance determination. Resistance assays lose their consistency at lower levels of plasma viremia.

Co-Receptor Tropism Assays Following the licensure of maraviroc as the first CCR5 antagonist for the treatment of HIV infection (see below), it became necessary to be able to determine whether a patient's virus was likely to respond to this treatment. Patients tend to have CCR5-tropic virus early in the course of infection, with a trend toward CXCR4 viruses later in disease. The antiretroviral agent maraviroc is effective only against CCR5-tropic viruses. Because the genotypic determinants of cellular tropism are poorly defined, a phenotypic assay is necessary to determine this property of HIV. Two commercial assays, the Trofile assay (Monogram Biosciences) and the Phenoscript assay (VIRalliance), are available to make this determination. These assays clone the envelope regions of the patient's virus into an indicator virus that is then used to infect target cells expressing either CCR5 or CXCR4 as their

TABLE 226-9 ASSOCIATION BETWEEN HIGH-SENSITIVITY CRP, IL-6, AND D-DIMER WITH ALL-CAUSE MORTALITY IN PATIENTS WITH HIV INFECTION

Marker	Unadjusted		Adjusted	
	Odds Ratio (Fourth/First)	p	Odds Ratio (Fourth/First)	p
Hs-CRP	2.0	.05	2.8	.03
IL-6	8.3	<.0001	11.8	<.0001
D-dimer	12.4	<.0001	26.5	<.0001

Abbreviations: Hs-CRP, high-sensitivity C-reactive protein; IL-6, interleukin 6.

Source: From LH Kuller et al: *PLoS Med* 5:e203, 2008.

co-receptor. These assays take weeks to perform and are expensive. Another, less costly option is to obtain a genotypic assay of the V3 region of HIV-1 and then employ a computer algorithm to predict viral tropism from the sequence. While this approach is less expensive than the classic phenotypic assay, there are fewer data to validate its predictive value.

Other Tests A variety of other laboratory tests have been studied as potential markers of HIV disease activity. Among these are quantitative culture of replication-competent HIV from plasma, peripheral blood mononuclear cells, or resting CD4+ T cells; circulating levels of β_2 -microglobulin, soluble IL-2 receptor, IgA, acid-labile endogenous IFN, or TNF- α ; and the presence or absence of activation markers such as CD38, HLA-DR, and PD-1 on CD4+ or CD8+ T cells. Nonspecific serologic markers of inflammation and/or coagulation such as IL-6, D-dimer, and sCD14 have been shown to have a high correlation with all-cause mortality (Table 226-9). While these measurements have value as markers of disease activity and help to increase our understanding of the pathogenesis of HIV disease, they do not currently play a major role in the monitoring of patients with HIV infection.

CLINICAL MANIFESTATIONS

The clinical consequences of HIV infection encompass a spectrum ranging from an acute syndrome associated with primary infection to a prolonged asymptomatic state to advanced disease. It is best to regard HIV disease as beginning at the time of primary infection and progressing through various stages. As mentioned above, active virus replication and progressive immunologic impairment occur throughout the course of HIV infection in most patients. With the exception of the rare, true, “elite” virus controllers or long-term nonprogressors (see “Long-Term Survivors and Long-Term Nonprogressors,” above), HIV disease in untreated patients inexorably progresses even during the clinically latent stage. Since the mid-1990s, cART has had a major impact on preventing and reversing the progression of disease over extended periods of time in a substantial proportion of adequately treated patients.

ACUTE HIV INFECTION

It is estimated that 50–70% of individuals with HIV infection experience an acute clinical syndrome ~3–6 weeks after primary infection (Fig. 226-32). Varying degrees of clinical severity have been reported, and although it has been suggested that symptomatic seroconversion leading to the seeking of medical attention indicates an increased risk for an accelerated course of disease, there does not appear to be a correlation between the level of the initial burst of viremia in acute HIV infection and the subsequent course of disease. The typical clinical findings in the acute HIV syndrome are listed in Table 226-10; they occur along with a burst of plasma viremia. It has been reported that several symptoms of the acute HIV syndrome (fever, skin rash, pharyngitis, and myalgia) occur less frequently in those infected by injection drug use compared with those infected by sexual contact. The syndrome is typical of an acute viral syndrome and has been likened to acute infectious mononucleosis. Symptoms usually persist for one to several weeks and gradually subside as an immune response to HIV develops and the levels of plasma viremia decrease. Opportunistic infections have been reported during this stage of infection, reflecting the immunodeficiency that results from reduced numbers of CD4+

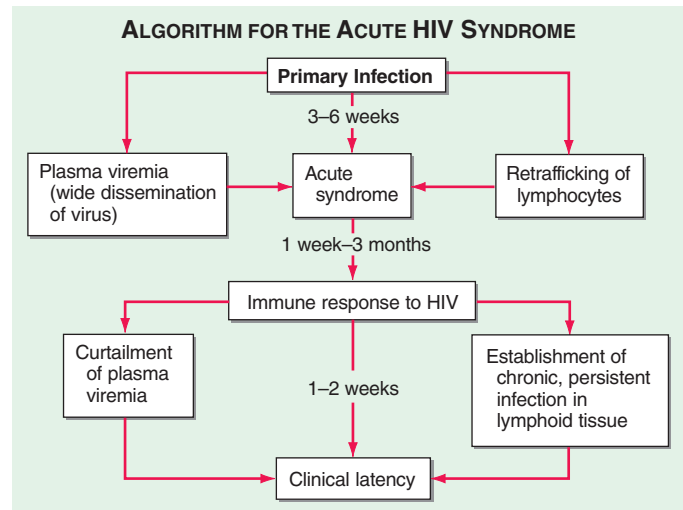


FIGURE 226-32 The acute HIV syndrome. See text for detailed description. (Adapted from G Pantaleo et al: *N Engl J Med* 328:327, 1993. Copyright 1993 Massachusetts Medical Society. All rights reserved.)

T cells and likely also from the dysfunction of CD4+ T cells owing to viral protein and endogenous cytokine-induced perturbations of cells (Table 226-5) associated with the extremely high levels of plasma viremia. A number of immunologic abnormalities accompany the acute HIV syndrome, including multiphasic perturbations of the numbers of circulating lymphocyte subsets. The number of total lymphocytes and T cell subsets (CD4+ and CD8+) are initially reduced. An inversion of the CD4+/CD8+ T cell ratio occurs later because of a rise in the number of CD8+ T cells. In fact, there may be a selective and transient expansion of CD8+ T cell subsets, as determined by T cell receptor analysis (see above). The total circulating CD8+ T cell count may remain elevated or return to normal; however, CD4+ T cell levels usually remain somewhat depressed, although there may be a slight rebound toward normal. Lymphadenopathy occurs in ~70% of individuals with primary HIV infection. Most patients recover spontaneously from this syndrome and many are left with only a mildly depressed CD4+ T cell count that remains stable for a variable period before beginning its progressive decline; in some individuals, the CD4+ T cell count returns to the normal range. Approximately 10% of patients manifest a fulminant course of immunologic and clinical deterioration after primary infection, even after the disappearance of initial symptoms. In most patients, primary infection with or without the acute syndrome is followed by a prolonged period of clinical latency or smoldering low disease activity.

THE ASYMPTOMATIC STAGE—CLINICAL LATENCY

Although the length of time from initial infection to the development of clinical disease varies greatly, the median time for untreated patients is ~10 years. As emphasized above, HIV disease with active virus replication is ongoing and progressive during this asymptomatic period. The rate of disease progression is directly correlated with HIV

TABLE 226-10 CLINICAL FINDINGS IN THE ACUTE HIV SYNDROME

General	Neurologic
Fever	Meningitis
Pharyngitis	Encephalitis
Lymphadenopathy	Peripheral neuropathy
Headache/retroorbital pain	Myelopathy
Arthralgias/myalgias	Dermatologic
Lethargy/malaise	Erythematous maculopapular rash
Anorexia/weight loss	Mucocutaneous ulceration
Nausea/vomiting/diarrhea	

Source: From B Tindall, DA Cooper: *AIDS* 5:1, 1991.

1250 RNA levels. Patients with high levels of HIV RNA in plasma progress to symptomatic disease faster than do patients with low levels of HIV RNA (Fig. 226-22). Some patients referred to as *long-term nonprogressors* show little if any decline in CD4+ T cell counts over extended periods of time. These patients generally have extremely low levels of HIV RNA; a subset, referred to as *elite nonprogressors*, exhibits HIV RNA levels <50 copies/mL. Certain other patients remain entirely asymptomatic despite the fact that their CD4+ T cell counts show a steady progressive decline to extremely low levels. In these patients, the appearance of an opportunistic disease may be the first manifestation of HIV infection. During the asymptomatic period of HIV infection, the average rate of CD4+ T cell decline is ~50/μL per year. When the CD4+ T cell count falls to <200/μL, the resulting state of immunodeficiency is severe enough to place the patient at high risk for opportunistic infection and neoplasms and, hence, for clinically apparent disease.

SYMPTOMATIC DISEASE

Symptoms of HIV disease can appear at any time during the course of HIV infection. Generally speaking, the spectrum of illnesses that one observes changes as the CD4+ T cell count declines. The more severe and life-threatening complications of HIV infection occur in patients with CD4+ T cell counts <200/μL. A diagnosis of AIDS is made in individuals age 6 years and older with HIV infection and a CD4+ T cell count <200/μL (Stage 3, Table 226-2) and in anyone with HIV infection who develops one of the HIV-associated diseases considered to be indicative of a severe defect in cell-mediated immunity (Table 226-1). While the causative agents of the secondary infections are characteristically opportunistic organisms such as *P. jiroveci*, atypical mycobacteria, CMV, and other organisms that do not ordinarily cause disease in the absence of a compromised immune system, they also include common bacterial and mycobacterial pathogens. Following the widespread use of cART and implementation of guidelines for the prevention of opportunistic infections (Table 226-11), the incidence of these secondary infections has decreased dramatically (Fig. 226-33). Overall, the clinical spectrum of HIV disease is constantly changing as patients live longer and new and better approaches to treatment and prophylaxis are developed. In addition to the classic AIDS-defining illnesses, patients with HIV infection also have an increase in serious non-AIDS illnesses, including non-AIDS related cancers and cardiovascular, renal, and hepatic disease. Non-AIDS events dominate the disease burden for patients with HIV infection receiving cART (Table 226-4). While AIDS-related illnesses are the leading cause of death in patients with HIV infection, they account for fewer than 50% of deaths. Non-AIDS-defining malignancies, liver disease, and cardiovascular disease each account for 10–15% of deaths in patients with HIV infection. The physician providing care to a patient with HIV infection must be well versed in general internal medicine as well as HIV-related opportunistic diseases. In general, it should be stressed that a key element of treatment of symptomatic complications of HIV disease, whether they are primary or secondary, is achieving good control of HIV replication through the use of cART and instituting primary and secondary prophylaxis for opportunistic infections as indicated.

Diseases of the Respiratory System Acute bronchitis and sinusitis are prevalent during all stages of HIV infection. The most severe cases tend to occur in patients with lower CD4+ T cell counts. Sinusitis presents as fever, nasal congestion, and headache. The diagnosis is made by CT or MRI. The maxillary sinuses are most commonly involved; however, disease is also frequently seen in the ethmoid, sphenoid, and frontal sinuses. While some patients may improve without antibiotic therapy, radiographic improvement is quicker and more pronounced in patients who have received antimicrobial therapy. It is postulated that this high incidence of sinusitis results from an increased frequency of infection with encapsulated organisms such as *H. influenzae* and *Streptococcus pneumoniae*. In patients with low CD4+ T cell counts one may see mucormycosis infections of the sinuses. In contrast to the course of this infection in other patient populations, mucormycosis of the sinuses in patients with HIV infection may progress more slowly. In this setting aggressive, frequent

local debridement in addition to local and systemic amphotericin B may result in effective treatment.

Pulmonary disease is one of the most frequent complications of HIV infection. The most common manifestation of pulmonary disease is pneumonia. Three of the 10 most common AIDS-defining illnesses are recurrent bacterial pneumonia, tuberculosis, and pneumonia due to the unicellular fungus *P. jiroveci*. Other major causes of pulmonary infiltrates include other mycobacterial infections, other fungal infections, nonspecific interstitial pneumonitis, KS, and lymphoma.

Bacterial pneumonia is seen with an increased frequency in patients with HIV infection, with 0.8–2.0 cases per 100 person-years. Patients with HIV infection are particularly prone to infections with encapsulated organisms. *S. pneumoniae* (Chap. 171) and *H. influenzae* (Chap. 182) are responsible for most cases of bacterial pneumonia in patients with AIDS. This may be a consequence of altered B cell function and/or defects in neutrophil function that may be secondary to HIV disease (see above). Pneumonias due to *S. aureus* (Chap. 172) and *P. aeruginosa* (Chap. 189) also are reported to occur with an increased frequency in patients with HIV infection. *S. pneumoniae* (pneumococcal) infection may be the earliest serious infection to occur in patients with HIV disease. This can present as pneumonia, sinusitis, and/or bacteremia. Patients with untreated HIV infection have a sixfold increase in the incidence of pneumococcal pneumonia and a 100-fold increase in the incidence of pneumococcal bacteremia. Pneumococcal disease may be seen in patients with relatively intact immune systems. In one study, the baseline CD4+ T cell count at the time of a first episode of pneumococcal pneumonia was ~300/μL. Of interest is the fact that the inflammatory response to pneumococcal infection appears proportional to the CD4+ T cell count. Due to this high risk of pneumococcal disease, immunization with the conjugated pneumococcal vaccine followed by booster immunization with the 23-valent pneumococcal polysaccharide vaccine is one of the generally recommended prophylactic measures for patients with HIV infection. This is likely most effective if given while the CD4+ T cell count is >200/μL and, if given to patients with lower CD4+ T cell counts, should be repeated once the count has been above 200 for 6 months. Although clear guidelines do not exist, it also makes sense to repeat immunization every 5 years. The incidence of bacterial pneumonia is cut in half when patients quit smoking.

Pneumocystis pneumonia (PCP), once the hallmark of AIDS, has dramatically declined in incidence following the development of effective prophylactic regimens and the widespread use of cART. It is, however, still the single most common cause of pneumonia in patients with HIV infection in the United States and can be identified as a likely etiologic agent in 25% of cases of pneumonia in patients with HIV infection, with an incidence in the range of 2–3 cases per 100 person-years. Approximately 50% of cases of HIV-associated PCP occur in patients who are unaware of their HIV status. The risk of PCP is greatest among those who have experienced a previous bout of PCP and those who have CD4+ T cell counts of <200/μL. Overall, 79% of patients with PCP have CD4+ T cell counts <100/μL and 95% of patients have CD4+ T cell counts <200/μL. Recurrent fever, night sweats, thrush, and unexplained weight loss also are associated with an increased incidence of PCP. For these reasons, it is strongly recommended that all patients with CD4+ T cell counts <200/μL (or a CD4 percentage <15) receive some form of PCP prophylaxis. The incidence of PCP is approaching zero in patients with known HIV infection receiving appropriate cART and prophylaxis. In the United States, primary PCP is now occurring at a median CD4+ T cell count of 36/μL, while secondary PCP is occurring at a median CD4+ T cell count of 10/μL. Patients with PCP generally present with fever and a cough that is usually nonproductive or productive of only scant amounts of white sputum. They may complain of a characteristic retrosternal chest pain that is worse on inspiration and is described as sharp or burning. HIV-associated PCP may have an indolent course characterized by weeks of vague symptoms and should be included in the differential diagnosis of fever, pulmonary complaints, or unexplained weight loss in any patient with HIV infection and <200 CD4+ T cells/μL. The most

TABLE 226-11 NIH/CDC/IDSA 2013 GUIDELINES FOR THE PREVENTION OF OPPORTUNISTIC INFECTIONS IN PERSONS INFECTED WITH HIV

Pathogen	Indications	First Choice(s)	Alternatives			
Recommended as Standard of Care for Primary and Secondary Prophylaxis						
<i>Pneumocystis jiroveci</i>	CD4+ T cell count <200/ μ L	Trimethoprim/sulfamethoxazole (TMP/SMX), 1 DS tablet qd PO	Dapsone 50 mg bid PO or 100 mg/d PO			
	or		or			
	Oropharyngeal candidiasis	or	Dapsone 50 mg/d PO + Pyrimethamine 50 mg/week PO + Leucovorin 25 mg/week PO			
	or		or			
	Prior bout of PCP		Leucovorin 25 mg/week PO			
	May stop prophylaxis if CD4+ T cell count >200/ μ L for \geq 3 months	TMP/SMX, 1 SS tablet qd PO	or (Dapsone 200 mg PO + Pyrimethamine 75 mg PO + Leucovorin 25 mg) weekly PO			
			or Aerosolized pentamidine, 300 mg via Respigard II nebulizer every month			
			or Atovaquone 1500 mg/d PO			
			or TMP/SMX 1 DS tablet 3 \times /week PO			
<i>Mycobacterium tuberculosis</i>	Isoniazid sensitive	Skin test >5 mm or Positive IFN- γ release assay or Prior positive test without treatment or Close contact with case of active pulmonary TB	(Isoniazid 300 mg PO + Pyridoxine 25 mg PO) qd \times 9 months or Isoniazid 900 mg PO twice weekly + Pyridoxine 25 mg PO daily \times 9 months			
				Drug resistant	Same with high probability of exposure to drug-resistant TB	Consult local public health authorities
<i>Mycobacterium-avium</i> complex	CD4+ T cell count <50/ μ L	Azithromycin 1200 mg weekly PO or 600 mg twice weekly PO	Rifabutin (dose adjusted based upon cART regimen)			
	or	Clarithromycin 500 mg bid PO				
	Prior documented disseminated disease	Clarithromycin 500 mg bid PO + Ethambutol 15 (mg/kg)/d PO	Azithromycin 500–600 mg/d PO + Ethambutol 15 (mg/kg)/d PO			
	May stop prophylaxis if CD4+ T cell count >100/ μ L for \geq 6 months					
<i>Toxoplasma gondii</i>	TOXO IgG antibody positive and CD4+ T cell count <100/ μ L	TMP/SMX 1 DS tablet PO qd	TMP/SMX 1 DS 3 \times weekly PO			
			or			
			TMP/SMX, 1 SS PO daily			
			or			
			Dapsone 50 mg/d PO + Pyrimethamine 50 mg weekly PO + Leucovorin 25 mg weekly PO			
			or			
			(Dapsone 200 mg PO + Pyrimethamine 75 mg PO + Leucovorin 25 mg PO) weekly			
			or			
			Atovaquone 1500 mg PO daily \pm (Pyrimethamine 25 mg PO + Leucovorin 10 mg PO) daily			
			or			
	Prior toxoplasmic encephalitis and CD4+ T cell count <200/ μ L	Sulfadiazine 2000–4000 mg in 2–4 divided doses daily PO + Pyrimethamine 25–50 mg/d PO + Leucovorin 10–25 mg/d PO	Clindamycin 600 mg q8h PO + Pyrimethamine 25–50 mg/d PO + Leucovorin 10–25 mg/d PO			
			or			

(Continued)

Pathogen	Indications	First Choice(s)	Alternatives
<i>Toxoplasma gondii</i>	May stop prophylaxis if CD4+ T cell count >200/μL for ≥3 months		TMP/SMX 1 DS tablet bid or Atovaquone 750–1500 mg PO bid ± (Pyrimethamine 25 mg/d PO + Leucovorin 10 mg/d PO) or Sulfadiazine 2000–4000 mg/d (in 2–4 divided doses) PO
Varicella zoster virus	Significant exposure to chickenpox or shingles in a patient with no history of immunization or prior exposure to either	Varicella zoster immune globulin, IM, within 10 d of exposure (1-800-843-7477)	Acyclovir 800 mg PO 5 × day for 5–7 days or Valacyclovir 1 g PO tid for 5–7 days
<i>Cryptococcus neoformans</i>	Prior documented disease May stop prophylaxis if CD4+ T cell count >100/μL, no evidence of active fungal infection, and HIV RNA levels <500 copies/mL for >3 months	Fluconazole 200 mg/d PO	Itraconazole 200 mg/d PO
<i>Histoplasma capsulatum</i>	Prior documented disease or CD4+ T cell count <150/μL and high risk (endemic area or occupational exposure) May stop prophylaxis after 1 year if CD4+ T cell count >150/μL and patient on cART for ≥6 months	Itraconazole 200 mg bid PO	Fluconazole 400 mg/d PO
<i>Coccidioides immitis</i>	Prior documented disease or positive serology and CD4+ T cell count <250/μL if from a disease endemic area. (For this indication prophylaxis can be stopped if CD4+ T cell count ≥250 for 6 months.)	Fluconazole 400 mg/d PO	
<i>Penicillium marneffei</i>	Prior documented disease Patients with CD4+T cell counts <100 who live or stay in northern Thailand, Southern China, or Vietnam May stop secondary prophylaxis in patients on ARV therapy with CD4+ T cell count >100/μL for ≥6 months	Itraconazole 200 mg/d PO	Fluconazole 400 mg PO once weekly
<i>Salmonella</i> species	Prior recurrent bacteremia	Ciprofloxacin 500 mg bid PO for ≥6 months	
<i>Bartonella</i>	Prior infection May stop if CD4+ T cell count >200/μL for >3 months	Doxycycline 200 mg/d PO or Azithromycin 1200 mg weekly PO or Clarithromycin 500 mg bid PO	
Cytomegalovirus	Prior end-organ disease May stop prophylaxis if CD4+ T cell count >100/μL for 6 months and no evidence of active CMV disease Restart if prior retinitis and CD4+ T cells <100/μL	Valganciclovir 900 mg bid PO	Cidofovir 5 mg/kg every other week IV + Probenecid or Foscarnet 90–120 (mg/kg)/d IV
Immunizations Generally Recommended			
Hepatitis B virus	All susceptible (anti-HBc- and anti-HBs-negative) patients	Hepatitis B vaccine: 3 doses	
Hepatitis A virus	All susceptible (anti-HAV-negative) patients	Hepatitis A vaccine: 2 doses	
Influenza virus	All patients annually	Inactivated trivalent influenza virus vaccine 1 dose yearly	Oseltamivir 75 mg PO qd or Rimantadine or amantadine 100 mg PO bid (influenza A only)
<i>Streptococcus pneumoniae</i>	All patients, preferably before CD4+ T cell count ≤200/μL	Pneumococcal conjugated vaccine (13) 0.5 mL IM × 1 followed in 8 weeks or more by pneumococcal polysaccharide vaccine (23) if CD4+ T cell count >200/μL	

(Continued)

TABLE 226-11 NIH/CDC/IDSA 2013 GUIDELINES FOR THE PREVENTION OF OPPORTUNISTIC INFECTIONS IN PERSONS INFECTED WITH HIV (CONTINUED)

Pathogen	Indications	First Choice(s)	Alternatives
<i>Streptococcus pneumoniae</i>		Reimmunize patients initially immunized at a CD4+ T cell count <100/ μ L whose CD4+ T cell count then increases to >200/ μ L	
Human papillomavirus	All patients 13–26 years of age	HPV vaccine; 3 doses	
Recommended for Prevention of Severe or Frequent Recurrences			
Herpes simplex	Frequent/severe recurrences	Valacyclovir 500 mg bid PO or Acyclovir 400 mg bid PO or Famciclovir 500 mg bid PO	
<i>Candida</i>	Frequent/severe recurrences	Fluconazole 100–200 mg/d PO	Posaconazole 400 mg bid PO

Abbreviations: ARV, antiretroviral; bid, twice daily; DS, double-strength; PCP, *Pneumocystis jiroveci* pneumonia; PO, by mouth; SS, single-strength; TB, tuberculosis.

common finding on chest x-ray is either a normal film, if the disease is suspected early, or a faint bilateral interstitial infiltrate. The classic finding of a dense perihilar infiltrate is unusual in patients with AIDS. In patients with PCP who have been receiving aerosolized pentamidine for prophylaxis, one may see an x-ray picture of upper lobe cavitory disease, reminiscent of TB. Other less common findings on chest x-ray include lobar infiltrates and pleural effusions. Thin-section CT

may demonstrate a patchy ground-glass appearance. Routine laboratory evaluation is usually of little help in the differential diagnosis of PCP. A mild leukocytosis is common, although this may not be obvious in patients with prior neutropenia. Elevation of lactate dehydrogenase is common. Arterial blood-gases may indicate hypoxemia with a decline in $P_{a_{O_2}}$ and an increase in the arterial-alveolar (a–A) gradient. Arterial blood-gas measurements not only aid in making the diagnosis of PCP but also provide important information for staging the severity of the disease and directing treatment (see below). A definitive diagnosis of PCP requires demonstration of the organism in samples obtained from induced sputum, bronchoalveolar lavage, transbronchial biopsy, or open-lung biopsy. PCR has been used to detect specific DNA sequences for *P. jiroveci* in clinical specimens where histologic examinations have failed to make a diagnosis.

In addition to pneumonia, a number of other clinical problems have been reported in HIV-infected patients as a result of infection with *P. jiroveci*. Otic involvement may be seen as a primary infection, presenting as a polypoid mass involving the external auditory canal. In patients receiving aerosolized pentamidine for prophylaxis against PCP, one may see a variety of extrapulmonary manifestations of *P. jiroveci*. These include ophthalmic lesions of the choroid, a necrotizing vasculitis that resembles Burger's disease, bone marrow hypoplasia, and intestinal obstruction. Other organs that have been involved include lymph nodes, spleen, liver, kidney, pancreas, pericardium, heart, thyroid, and adrenals. Organ infection may be associated with cystic lesions that may appear calcified on CT or ultrasound.

The standard treatment for PCP or disseminated pneumocystosis is trimethoprim/sulfamethoxazole (TMP/SMX). A high (20–85%) incidence of side effects, particularly skin rash and bone marrow suppression, is seen with TMP/SMX in patients with HIV infection. Alternative treatments for mild to moderate PCP include dapsone/trimethoprim, clindamycin/primaquine, and atovaquone. IV pentamidine is the treatment of choice for severe disease in the patient unable to tolerate TMP/SMX. For patients with a $P_{a_{O_2}}$ <70 mmHg or with an a–A gradient >35 mmHg, adjunct glucocorticoid therapy should be used in addition to specific antimicrobials. Overall, treatment should be continued for 21 days and followed by secondary prophylaxis. Prophylaxis for PCP is indicated for any HIV-infected individual who has experienced a prior bout of PCP, any patient with a CD4+ T cell count of <200/ μ L or a CD4 percentage <15, any patient with unexplained fever for >2 weeks, and any patient with a recent history of oropharyngeal candidiasis. The preferred regimen for prophylaxis is TMP/SMX, one double-strength tablet daily. This regimen also provides protection against toxoplasmosis and some bacterial respiratory pathogens. For patients who cannot tolerate TMP/SMX, alternatives for prophylaxis include dapsone plus pyrimethamine plus leucovorin, aerosolized pentamidine administered by the Respigard II nebulizer, and atovaquone. Primary or secondary prophylaxis for PCP can be

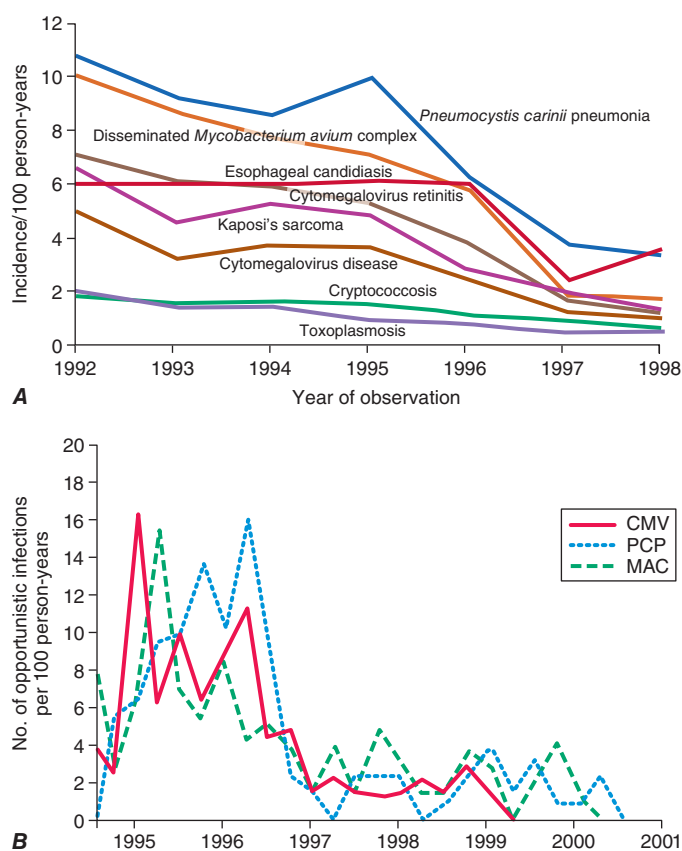


FIGURE 226-33 **A.** Decrease in the incidence of opportunistic infections and Kaposi's sarcoma in HIV-infected individuals with CD4+ T cell counts <100/ μ L from 1992 through 1998. (Adapted and updated from FJ Palella et al: *N Engl J Med* 338:853, 1998, and JE Kaplan et al: *Clin Infect Dis* 30[S1]:S5, 2000, with permission.) **B.** Quarterly incidence rates of cytomegalovirus (CMV), *Pneumocystis jiroveci* pneumonia (PCP), and *Mycobacterium avium* complex (MAC) from 1995 to 2001. (From FJ Palella et al: *AIDS* 16:1617, 2002.)

1254 discontinued in those patients treated with cART who maintain good suppression of HIV (<50 copies/mL) and CD4+ T cell counts >200/μL for at least 3 months.

M. tuberculosis, once thought to be on its way to extinction in the United States, experienced a resurgence associated with the HIV epidemic (Chap. 202). Worldwide, approximately one-third of all AIDS-related deaths are associated with TB, and TB is the primary cause of death for 10–15% of patients with HIV infection. In the United States ~5% of AIDS patients have active TB. Patients with HIV infection are more likely to have active TB by a factor of 100 when compared with an HIV-negative population. For an asymptomatic HIV-negative person with a positive purified protein derivative (PPD) skin test, the risk of reactivation TB is around 1% per year. For the patient with untreated HIV infection, a positive PPD skin test, and no signs or symptoms of TB, the rate of reactivation TB is 7–10% per year. Untreated TB can accelerate the course of HIV infection. Levels of plasma HIV RNA increase in the setting of active TB and decline in the setting of successful TB treatment. Active TB is most common in patients 25–44 years of age, in African Americans and Hispanics, in patients in New York City and Miami, and in patients in developing countries. In these demographic groups, 20–70% of the new cases of active TB are in patients with HIV infection. The epidemic of TB embedded in the epidemic of HIV infection probably represents the greatest health risk to the general public and the health care profession associated with the HIV epidemic. In contrast to infection with atypical mycobacteria such as MAC, active TB often develops relatively early in the course of HIV infection and may be an early clinical sign of HIV disease. In one study, the median CD4+ T cell count at presentation of TB was 326/μL. The clinical manifestations of TB in HIV-infected patients are quite varied and generally show different patterns as a function of the CD4+ T cell count. In patients with relatively high CD4+ T cell counts, the typical pattern of pulmonary reactivation occurs: patients present with fever, cough, dyspnea on exertion, weight loss, night sweats, and a chest x-ray revealing cavitory apical disease of the upper lobes. In patients with lower CD4+ T cell counts, disseminated disease is more common. In these patients the chest x-ray may reveal diffuse or lower-lobe bilateral reticulonodular infiltrates consistent with miliary spread, pleural effusions, and hilar and/or mediastinal adenopathy. Infection may be present in bone, brain, meninges, GI tract, lymph nodes (particularly cervical lymph nodes), and viscera. Some patients with advanced HIV infection and active TB may have no symptoms of illness, and thus screening for TB should be part of the initial evaluation of every patient with HIV infection. Approximately 60–80% of HIV-infected patients with TB have pulmonary disease, and 30–40% have extrapulmonary disease. Respiratory isolation and a negative-pressure room should be used for patients in whom a diagnosis of pulmonary TB is being considered. This approach is critical to limit nosocomial and community spread of infection. Culture of the organism from an involved site provides a definitive diagnosis. Blood cultures are positive in 15% of patients. This figure is higher in patients with lower CD4+ T cell counts. In the setting of fulminant disease one cannot rely on the accuracy of a negative PPD skin test to rule out a diagnosis of TB. In addition, IFN-γ release assays may be difficult to interpret due to high backgrounds as a consequence of HIV-associated immune activation. TB is one of the conditions associated with HIV infection for which cure is possible with appropriate therapy. Therapy for TB is generally the same in the HIV-infected patient as in the HIV-negative patient (Chap. 202). Due to the possibility of multidrug-resistant or extensively drug-resistant TB, drug susceptibility testing should be performed to guide therapy. Due to pharmacokinetic interactions, adjusted doses of rifabutin should be substituted for rifampin in patients receiving the HIV protease inhibitors or nonnucleoside reverse transcriptase inhibitors. Treatment is most effective in programs that involve directly observed therapy. Initiation of cART and/or anti-TB therapy may be associated with clinical deterioration due to immune reconstitution inflammatory syndrome (IRIS) reactions. These are most common in patients initiating both treatments at the same time, may occur as early as 1 week after initiation of cART therapy, and are seen more frequently in patients

with advanced HIV disease. For these reasons it is recommended that initiation of cART be delayed in antiretroviral-naïve patients with CD4 counts >50 cells/μL until 2–4 weeks following the initiation of treatment for TB. For patients with lower CD4 counts the benefits of more immediate cART outweigh the risks of IRIS, and cART should be started as soon as possible in those patients. Effective prevention of active TB can be a reality if the health care professional is aggressive in looking for evidence of latent or active TB by making sure that all patients with HIV infection receive a PPD skin test or evaluation with an IFN-γ release assay. Anergy testing is not of value in this setting. Since these tests rely on the host mounting an immune response to *M. tuberculosis*, patients with CD4+ T cell counts <200 cells/μL should be retested if their CD4+ T cell counts rise to persistently above 200. Patients at risk of continued exposure to TB should be tested annually. HIV-infected individuals with a skin-test reaction of >5 mm, those with a positive IFN-γ release assay, or those who are close household contacts of persons with active TB should receive treatment with 9 months of isoniazid and pyridoxine.

Atypical mycobacterial infections are also seen with an increased frequency in patients with HIV infection. Infections with at least 12 different mycobacteria have been reported, including *M. bovis* and representatives of all four Runyon groups. The most common atypical mycobacterial infection is with *M. avium* or *M. intracellulare* species—the *Mycobacterium avium* complex (MAC). Infections with MAC are seen mainly in patients in the United States and are rare in Africa. It has been suggested that prior infection with *M. tuberculosis* decreases the risk of MAC infection. MAC infections probably arise from organisms that are ubiquitous in the environment, including both soil and water. There is little evidence for person-to-person transmission of MAC infection. The presumed portals of entry are the respiratory and GI tracts. MAC infection is a late complication of HIV infection, occurring predominantly in patients with CD4+ T cell counts of <50/μL. The average CD4+ T cell count at the time of diagnosis is 10/μL. The most common presentation is disseminated disease with fever, weight loss, and night sweats. At least 85% of patients with MAC infection are mycobacteremic, and large numbers of organisms can often be demonstrated on bone marrow biopsy. The chest x-ray is abnormal in ~25% of patients, with the most common pattern being that of a bilateral, lower-lobe infiltrate suggestive of miliary spread. Alveolar or nodular infiltrates and hilar and/or mediastinal adenopathy can also occur. Other clinical findings include endobronchial lesions, abdominal pain, diarrhea, and lymphadenopathy. Anemia and elevated liver alkaline phosphatase are common. The diagnosis is made by the culture of blood or involved tissue. The finding of two consecutive sputum samples positive for MAC is highly suggestive of pulmonary infection. Cultures may take 2 weeks to turn positive. Therapy consists of a macrolide, usually clarithromycin, with ethambutol. Some physicians elect to add a third drug from among rifabutin, ciprofloxacin, or amikacin in patients with extensive disease. Therapy was generally for life; however, with the use of cART it is possible to discontinue therapy in patients with sustained suppression of HIV replication and CD4+ T cell counts >100/μL for 3–6 months. Primary prophylaxis for MAC is indicated in patients with HIV infection and CD4+ T cell counts <50/μL (Table 226-11). This may be discontinued in patients in whom cART induces a sustained suppression of viral replication and an increase in CD4+ T cell count to >100/μL for ≥6 months.

Rhodococcus equi is a gram-positive, pleomorphic, acid-fast, non-spore-forming bacillus that can cause pulmonary and/or disseminated infection in patients with advanced HIV infection. Fever and cough are the most common presenting signs. Radiographically one may see cavitory lesions and consolidation. Blood cultures are often positive. Treatment is based on antimicrobial sensitivity testing.

Fungal infections of the lung, in addition to PCP, can be seen in patients with AIDS. Patients with pulmonary cryptococcal disease present with fever, cough, dyspnea, and, in some cases, hemoptysis. A focal or diffuse interstitial infiltrate is seen on chest x-ray in >90% of patients. In addition, one may see lobar disease, cavitory disease, pleural effusions, and hilar or mediastinal adenopathy. More than half of patients are fungemic, and 90% of patients have concomitant CNS

infection. *Coccidioides immitis* is a mold that is endemic in the southwest United States. It can cause a reactivation pulmonary syndrome in patients with HIV infection. Most patients with this condition will have CD4+ T cell counts <250/ μ L. Patients present with fever, weight loss, cough, and extensive, diffuse reticulonodular infiltrates on chest x-ray. One may also see nodules, cavities, pleural effusions, and hilar adenopathy. While serologic testing is of value in the immunocompetent host, serologies are negative in 25% of HIV-infected patients with coccidioidal infection. Invasive aspergillosis is not an AIDS-defining illness and is generally not seen in patients with AIDS in the absence of neutropenia or administration of glucocorticoids. When it does occur, *Aspergillus* infection may have an unusual presentation in the respiratory tract of patients with AIDS, where it gives the appearance of a pseudomembranous tracheobronchitis. Primary pulmonary infection of the lung may be seen with *histoplasmosis*. The most common pulmonary manifestation of histoplasmosis, however, is in the setting of disseminated disease, presumably due to reactivation. In this setting respiratory symptoms are usually minimal, with cough and dyspnea occurring in 10–30% of patients. The chest x-ray is abnormal in ~50% of patients, showing either a diffuse interstitial infiltrate or diffuse small nodules, and the urine will often be positive for *Histoplasma* antigen.

Two forms of *idiopathic interstitial pneumonia* have been identified in patients with HIV infection: lymphoid interstitial pneumonitis (LIP) and nonspecific interstitial pneumonitis (NIP). LIP, a common finding in children, is seen in about 1% of adult patients with untreated HIV infection. This disorder is characterized by a benign infiltrate of the lung and is thought to be part of the polyclonal activation of lymphocytes seen in the context of HIV and EBV infections. Transbronchial biopsy is diagnostic in 50% of the cases, with an open-lung biopsy required for diagnosis in the remainder of cases. This condition is generally self-limited and no specific treatment is necessary. Severe cases have been managed with brief courses of glucocorticoids. Although rarely a clinical problem since the use of cART, evidence of NIP may be seen in up to half of all patients with untreated HIV infection. Histologically, interstitial infiltrates of lymphocytes and plasma cells in a perivascular and peribronchial distribution are present. When symptomatic, patients present with fever and nonproductive cough occasionally accompanied by mild chest discomfort. Chest x-ray is usually normal or may reveal a faint interstitial pattern. Similar to LIP, NIP is a self-limited process for which no therapy is indicated other than appropriate management of the underlying HIV infection. HIV-related pulmonary arterial hypertension (HIV-PAH) is seen in ~0.5% of HIV-infected individuals. Patients may present with an array of symptoms including shortness of breath, fatigue, syncope, chest pain, and signs of right-sided heart failure. Chest x-ray reveals dilated pulmonary vessels and right-sided cardiomegaly with right ventricular hypertrophy seen on electrocardiogram. cART does not appear to be of clear benefit, and the prognosis is quite poor with a median survival in the range of 2 years.

Neoplastic diseases of the lung including KS and lymphoma are discussed below in the section on neoplastic diseases.

Diseases of the Cardiovascular System Heart disease is a relatively common postmortem finding in HIV-infected patients (25–75% in autopsy series). The most common form of heart disease is coronary heart disease. In one large series the overall rate of myocardial infarction (MI) was 3.5/1000 patient-years, 28% of these events were fatal, and MI was responsible for 7% of all deaths in the cohort. In patients with HIV infection, cardiovascular disease may be associated with classic risk factors such as smoking, a direct consequence of HIV infection, or a complication of cART. Patients with HIV infection have higher levels of triglycerides, lower levels of high-density lipoprotein cholesterol, and a higher prevalence of smoking than cohorts of individuals without HIV infection. The finding that the rate of cardiovascular disease events was lower in patients on antiretroviral therapy than in those randomized to undergo a treatment interruption identified a clear association between HIV replication and risk of cardiovascular disease. In one study, a baseline CD4+ T cell count of <500/ μ L was found to

be an independent risk factor for cardiovascular disease comparable in magnitude to that attributable to smoking. While the precise pathogenesis of this association remains unclear, it is likely related to the immune activation and increased propensity for coagulation seen as a consequence of HIV replication. Exposure to HIV protease inhibitors and certain reverse transcriptase inhibitors has been associated with increases in total cholesterol and/or risk of MI. Any increases in the risk of death from MI resulting from the use of certain antiretrovirals must be balanced against the marked increases in overall survival brought about by these drugs.

Another form of heart disease associated with HIV infection is a dilated cardiomyopathy associated with congestive heart failure (CHF) referred to as *HIV-associated cardiomyopathy*. This generally occurs as a late complication of HIV infection and, histologically, displays elements of myocarditis. For this reason some have advocated treatment with IV immunoglobulin (IVIg). HIV can be directly demonstrated in cardiac tissue in this setting, and there is debate over whether it plays a direct role in this condition. Patients present with typical findings of CHF including edema and shortness of breath. Patients with HIV infection may also develop cardiomyopathy as side effects of IFN- α or nucleoside analogue therapy. These are reversible once therapy is stopped. KS, cryptococcosis, Chagas' disease, and toxoplasmosis can involve the myocardium, leading to cardiomyopathy. In one series, most patients with HIV infection and a treatable myocarditis were found to have myocarditis associated with toxoplasmosis. Most of these patients also had evidence of CNS toxoplasmosis. Thus, MRI or double-dose contrast CT scan of the brain should be included in the workup of any patient with advanced HIV infection and cardiomyopathy.

A variety of other cardiovascular problems are found in patients with HIV infection. Pericardial effusions may be seen in the setting of advanced HIV infection. Predisposing factors include TB, CHF, mycobacterial infection, cryptococcal infection, pulmonary infection, lymphoma, and KS. While pericarditis is quite rare, in one series 5% of patients with HIV disease had pericardial effusions that were considered to be moderate or severe. Tamponade and death have occurred in association with pericardial KS, presumably owing to acute hemorrhage. Nonbacterial thrombotic endocarditis has been reported and should be considered in patients with unexplained embolic phenomena. IV pentamidine, when given rapidly, can result in hypotension as a consequence of cardiovascular collapse.

Diseases of the Oropharynx and Gastrointestinal System Oropharyngeal and GI diseases are common features of HIV infection. They are most frequently due to secondary infections. In addition, oral and GI lesions may occur with KS and lymphoma.

Oral lesions, including *thrush*, *hairy leukoplakia*, and *aphthous ulcers* (Fig. 226-34), are particularly common in patients with untreated HIV infection. Thrush, due to *Candida* infection, and oral hairy leukoplakia, presumed due to EBV, are usually indicative of fairly advanced immunologic decline; they generally occur in patients with CD4+ T cell counts of <300/ μ L. In one study, 59% of patients with oral candidiasis went on to develop AIDS in the next year. Thrush appears as a white, cheesy exudate, often on an erythematous mucosa in the posterior oropharynx. While most commonly seen on the soft palate, early lesions are often found along the gingival border. The diagnosis is made by direct examination of a scraping for pseudohyphal elements. Culturing is of no diagnostic value, as patients with HIV infection may have a positive throat culture for *Candida* in the absence of thrush. Oral hairy leukoplakia presents as white, frondlike lesions, generally along the lateral borders of the tongue and sometimes on the adjacent buccal mucosa (Fig. 226-34). Despite its name, oral hairy leukoplakia is not considered a premalignant condition. Lesions are associated with florid replication of EBV. While usually more disconcerting as a sign of HIV-associated immunodeficiency than a clinical problem in need of treatment, severe cases have been reported to respond to topical podophyllin or systemic therapy with anti-herpesvirus agents. Aphthous ulcers of the posterior oropharynx also are seen with regularity in patients with untreated HIV infection



FIGURE 226-34 Various oral lesions in HIV-infected individuals. **A.** Thrush. **B.** Hairy leukoplakia. **C.** Aphthous ulcer. **D.** Kaposi's sarcoma.

(Fig. 226-34). These lesions are of unknown etiology and can be quite painful and interfere with swallowing. Topical anesthetics provide immediate symptomatic relief of short duration. The fact that thalidomide is an effective treatment for this condition suggests that the pathogenesis may involve the action of tissue-destructive cytokines. Palatal, glossal, or gingival ulcers may also result from cryptococcal disease or histoplasmosis.

Esophagitis (Fig. 226-35) may present with odynophagia and retrosternal pain. Upper endoscopy is generally required to make an accurate diagnosis. Esophagitis may be due to *Candida*, CMV, or HSV. While CMV tends to be associated with a single large ulcer, HSV infection is more often associated with multiple small ulcers. The esophagus may also be the site of KS and lymphoma. Like the oral mucosa, the esophageal mucosa may have large, painful ulcers of unclear etiology that may respond to thalidomide. While achlorhydria is a common problem in patients with HIV infection, other gastric problems are generally rare. Among the neoplastic conditions involving the stomach are KS and lymphoma.

Infections of the small and large intestine leading to diarrhea, abdominal pain, and occasionally fever are among the most significant GI problems in HIV-infected patients. They include infections with bacteria, protozoa, and viruses.

Bacteria may be responsible for secondary infections of the GI tract. Infections with enteric pathogens such as *Salmonella*, *Shigella*, and *Campylobacter* are more common in men who have sex with men and are often more severe and more apt to relapse in patients with HIV infection. Patients with untreated HIV have approximately a 20-fold increased risk of infection with *S. typhimurium*.



FIGURE 226-35 Barium swallow of a patient with *Candida* esophagitis. The flow of barium along the mucosal surface is grossly irregular.

They may present with a variety of nonspecific symptoms including fever, anorexia, fatigue, and malaise of several weeks' duration. Diarrhea is common but may be absent. Diagnosis is made by culture of blood and stool. Long-term therapy with ciprofloxacin is the recommended treatment. HIV-infected patients also have an increased incidence of *S. typhi* infection in areas of the world where typhoid is a problem. *Shigella* spp., particularly *S. flexneri*, can cause severe intestinal disease in HIV-infected individuals. Up to 50% of patients will develop bacteremia. *Campylobacter* infections occur with an increased frequency in patients with HIV infection. While *C. jejuni* is the strain most frequently isolated, infections with many other strains have been reported. Patients usually present with crampy abdominal pain, fever, and bloody diarrhea. Infection may also present as proctitis. Stool examination reveals the presence of fecal leukocytes. Systemic infection can occur, with up to 10% of infected patients exhibiting bacteremia. Most strains are sensitive to erythromycin. Abdominal pain and diarrhea may be seen with MAC infection.

Fungal infections may also be a cause of diarrhea in patients with HIV infection. Histoplasmosis, coccidioidomycosis, and penicilliosis have all been identified as a cause of fever and diarrhea in patients with HIV infection. Peritonitis has been seen with *C. immitis*.

Cryptosporidia, microsporidia, and *Isospora belli* (Chap. 254) are the most common opportunistic protozoa that infect the GI tract and cause diarrhea in HIV-infected patients. Cryptosporidial infection may present in a variety of ways, ranging from a self-limited or intermittent diarrheal illness in patients in the early stages of HIV infection to a severe, life-threatening diarrhea in severely immunodeficient individuals. In patients with untreated HIV infection and CD4+ T cell counts of <300/ μ L, the incidence of cryptosporidiosis is ~1% per year. In 75% of cases the diarrhea is accompanied by crampy abdominal pain, and 25% of patients have nausea and/or vomiting. Cryptosporidia may also cause biliary tract disease in the HIV-infected patient, leading to cholecystitis with or without accompanying cholangitis and pancreatitis secondary to papillary stenosis. The diagnosis of cryptosporidial diarrhea is made by stool examination or biopsy of the small intestine. The diarrhea is noninflammatory, and the characteristic finding is the presence of oocysts that stain with acid-fast dyes. Therapy is predominantly supportive, and marked improvements have been reported in the setting of effective cART. Treatment with up to 2000 mg/d of nitazoxanide (NTZ) is associated with improvement in symptoms or a decrease in shedding of organisms in about half of patients. Its overall role in the management of this condition remains unclear. Patients can minimize their risk of developing cryptosporidiosis by avoiding contact with human and animal feces, by not drinking untreated water from lakes or rivers, and by not eating raw shellfish.

Microsporidia are small, unicellular, obligate intracellular parasites that reside in the cytoplasm of enteric cells (Chap. 254). The main species causing disease in humans is *Enterocytozoon bieneusi*. The clinical manifestations are similar to those described for cryptosporidia and include abdominal pain, malabsorption, diarrhea, and cholangitis. The small size of the organism may make it difficult to detect; however, with the use of chromotrope-based stains, organisms can be identified in stool samples by light microscopy. Definitive diagnosis generally depends on electron-microscopic examination of a stool specimen, intestinal aspirate, or intestinal biopsy specimen. In contrast to cryptosporidia, microsporidia have been noted in a variety of extraintestinal locations, including the eye, brain, sinuses, muscle, and liver, and they have been associated with conjunctivitis and hepatitis. The most effective way to deal with microsporidia in a patient with HIV infection is to restore the immune system by treating the HIV infection with cART. Albendazole, 400 mg bid, has been reported to be of benefit in some patients.

I. belli is a coccidian parasite (Chap. 254) most commonly found as a cause of diarrhea in patients from tropical and subtropical regions. Its cysts appear in the stool as large, acid-fast structures that can be differentiated from those of cryptosporidia on the basis of size, shape, and number of sporocysts. The clinical syndromes of *Isospora* infection are identical to those caused by cryptosporidia. The important distinction

is that infection with *Isospora* is generally relatively easy to treat with TMP/SMX. While relapses are common, a thrice-weekly regimen of TMP/SMX appears adequate to prevent recurrence.

CMV colitis was once seen as a consequence of advanced immunodeficiency in 5–10% of patients with AIDS. It is much less common with the advent of cART. CMV colitis presents as diarrhea, abdominal pain, weight loss, and anorexia. The diarrhea is usually nonbloody, and the diagnosis is achieved through endoscopy and biopsy. Multiple mucosal ulcerations are seen at endoscopy, and biopsies reveal characteristic intranuclear and cytoplasmic inclusion bodies. Secondary bacteremias may result as a consequence of thinning of the bowel wall. Treatment is with either ganciclovir or foscarnet for 3–6 weeks. Relapses are common, and maintenance therapy is typically necessary in patients whose HIV infection is poorly controlled. Patients with CMV disease of the GI tract should be carefully monitored for evidence of CMV retinitis.

In addition to disease caused by specific secondary infections, patients with HIV infection may also experience a chronic diarrheal syndrome for which no etiologic agent other than HIV can be identified. This entity is referred to as *AIDS enteropathy* or *HIV enteropathy*. It is most likely a direct result of HIV infection in the GI tract. Histologic examination of the small bowel in these patients reveals low-grade mucosal atrophy with a decrease in mitotic figures, suggesting a hyporegenerative state. Patients often have decreased or absent small-bowel lactase and malabsorption with accompanying weight loss.

The initial evaluation of a patient with HIV infection and diarrhea should include a set of stool examinations, including culture, examination for ova and parasites, and examination for *Clostridium difficile* toxin. Approximately 50% of the time this workup will demonstrate infection with pathogenic bacteria, mycobacteria, or protozoa. If the initial stool examinations are negative, additional evaluation, including upper and/or lower endoscopy with biopsy, will yield a diagnosis of microsporidial or mycobacterial infection of the small intestine ~30% of the time. In patients for whom this diagnostic evaluation is nonrevealing, a presumptive diagnosis of HIV enteropathy can be made if the diarrhea has persisted for >1 month. An algorithm for the evaluation of diarrhea in patients with HIV infection is given in Fig. 226-36.

Rectal lesions are common in HIV-infected patients, particularly the perirectal ulcers and erosions due to the reactivation of HSV

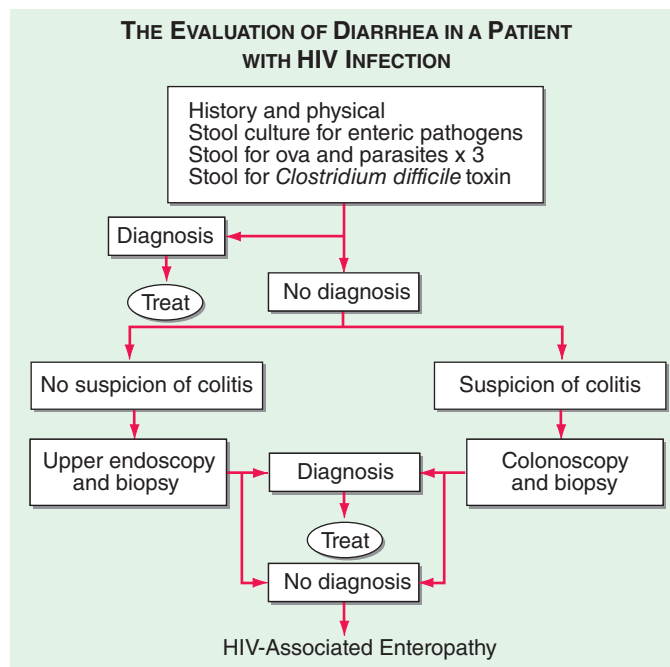


FIGURE 226-36 Algorithm for the evaluation of diarrhea in a patient with HIV infection. HIV-associated enteropathy is a diagnosis of exclusion and can be made only after other, generally treatable, forms of diarrheal illness have been ruled out.



FIGURE 226-37 Severe, erosive perirectal herpes simplex in a patient with AIDS.

(Fig. 226-37). These lesions may appear quite atypical, as denuded skin without vesicles. They typically respond well to treatment with acyclovir, famciclovir, or foscarnet. Other rectal lesions encountered in patients with HIV infection include condylomata acuminata, KS, and intraepithelial neoplasia (see below).

Hepatobiliary Diseases Diseases of the hepatobiliary system are a major problem in patients with HIV infection. It has been estimated that approximately 15% of the deaths of patients with HIV infection are related to liver disease. While this is predominantly a reflection of the problems encountered in the setting of co-infection with hepatitis B or C, it is also a reflection of the hepatic injury, ranging from hepatic steatosis to hypersensitivity reactions to immune reconstitution, that can be seen in the context of cART.

The prevalence of co-infection with HIV and hepatitis viruses varies by geographic region. In the United States, ~90% of HIV-infected individuals have evidence of infection with HBV; 6–14% have chronic HBV infection; 5–50% of patients are co-infected with HCV; and co-infections with hepatitis D, E, and/or G viruses are common. Among IV drug users with HIV infection, rates of HCV infection range from 70% to 95%. HIV infection has a significant impact on the course of hepatitis virus infection. It is associated with approximately a threefold increase in the development of persistent hepatitis B surface antigenemia. Patients infected with both HBV and HIV have decreased evidence of inflammatory liver disease. The presumption that this is due to the immunosuppressive effects of HIV infection is supported by the observations that this situation can be reversed, and one may see the development of more severe hepatitis following the initiation of effective cART. In studies of the impact of HIV on HBV infection, four- to tenfold increases in liver-related mortality rates have been noted in patients with HIV and active HBV infection compared to rates in patients with either infection alone. There is, however, only a slight increase in overall mortality rate in HIV-infected individuals who are also hepatitis B surface antigen (HBsAg)-positive. IFN- α is less successful as treatment for HBV in patients with HIV co-infection. Lamivudine, emtricitabine, adefovir/tenofovir/entecavir, and telbivudine alone or in combination are useful in the treatment of hepatitis B in patients with HIV infection. It is important to remember that all the above-mentioned drugs also have activity

against HIV and should not be used alone in patients with HIV infection, in order to avoid the emergence of quasispecies of HIV resistant to these drugs. For this reason, the need to treat hepatitis B infection in a patient with HIV infection is an indication to treat HIV infection in that same patient, regardless of CD4+ T cell count. HCV infection is more severe in the patient with HIV infection; it does not appear to affect overall mortality rates in HIV-infected individuals when other variables such as age, baseline CD4+ T cell count, and use of cART are taken into account. In the setting of HIV and HCV co-infection, levels of HCV are approximately tenfold higher than in the HIV-negative patient with HCV infection. There is a 50% higher overall mortality rate with a five-fold increased risk of death due to liver disease in patients chronically infected with both HCV and HIV. Use of directly acting agents for the treatment of HCV leads to cure rates approaching 100%, even in patients with HIV co-infection. Successful treatment of HCV in HIV-infected patients decreases mortality. Hepatitis A virus infection is not seen with an increased frequency in patients with HIV infection. It is recommended that all patients with HIV infection who have not experienced natural infection be immunized with hepatitis A and/or hepatitis B vaccines. Infection with hepatitis G virus, also known as GB virus C, is seen in ~50% of patients with HIV infection. For reasons that are currently unclear, there are data to suggest that patients with HIV infection co-infected with this virus have a decreased rate of progression to AIDS.

A variety of other infections also may involve the liver. Granulomatous hepatitis may be seen as a consequence of mycobacterial or fungal infections, particularly MAC infection. Hepatic masses may be seen in the context of TB, peliosis hepatis, or fungal infection. Among the fungal opportunistic infections, *C. immitis* and *Histoplasma capsulatum* are those most likely to involve the liver. Biliary tract disease in the form of papillary stenosis or sclerosing cholangitis has been reported in the context of cryptosporidiosis, CMV infection, and KS. When no diagnosis can be made, the term AIDS cholangiopathy is used. Hemophagocytic lymphohistiocytosis of the liver has been seen in the setting of Hodgkin's disease.

Many of the drugs used to treat HIV infection are metabolized by the liver and can cause liver injury. Fatal hepatic reactions have been reported with a wide array of antiretrovirals including nucleoside analogues, nonnucleoside analogues, and protease inhibitors. Nucleoside analogues work by inhibiting DNA synthesis. This can result in toxicity to mitochondria, which can lead to disturbances in oxidative metabolism. This may manifest as hepatic steatosis and, in severe cases, lactic acidosis and fulminant liver failure. It is important to be aware of this condition and to watch for it in patients with HIV infection receiving nucleoside analogues. It is reversible if diagnosed early and the offending agent(s) discontinued. Nevirapine has been associated with at times fatal fulminant and cholestatic hepatitis, hepatic necrosis, and hepatic failure. Indinavir may cause mild to moderate elevations in serum bilirubin in 10–15% of patients in a syndrome similar to Gilbert's syndrome. A similar pattern of hepatic injury may be seen with atazanavir. In the patient receiving cART with an unexplained increase in hepatic transaminases, strong consideration should be given to drug toxicity.

Pancreatic injury is most commonly a consequence of drug toxicity, notably that secondary to pentamidine or dideoxynucleosides. While up to half of patients in some series have biochemical evidence of pancreatic injury, <5% of patients show any clinical evidence of pancreatitis that is not linked to a drug toxicity.

Diseases of the Kidney and Genitourinary Tract Diseases of the kidney or genitourinary tract may be a direct consequence of HIV infection, due to an opportunistic infection or neoplasm, or related to drug toxicity. Overall, microalbuminuria is seen in ~20% of untreated HIV-infected patients; significant proteinuria is seen in closer to 2%. The presence of microalbuminuria has been associated with an increase in all-cause mortality rate. *HIV-associated nephropathy* (HIVAN) was first described in IDUs and was initially thought to be IDU nephropathy in patients with HIV infection; it is now recognized as a true direct complication of HIV infection. Although the majority of patients with this condition have CD4+ T cell counts <200/ μ L,

HIV-associated nephropathy can be an early manifestation of HIV infection and is also seen in children. Over 90% of reported cases have been in African-American or Hispanic individuals; the disease is not only more prevalent in these populations but also more severe and is the third leading cause of end-stage renal failure among African Americans age 20–64 in the United States. Proteinuria is the hallmark of this disorder. Edema and hypertension are rare. Ultrasound examination reveals enlarged, hyperechogenic kidneys. A definitive diagnosis is obtained through renal biopsy. Histologically, focal segmental glomerulosclerosis is present in 80%, and mesangial proliferation in 10–15% of cases. Prior to effective antiretroviral therapy, this disease was characterized by relatively rapid progression to end-stage renal disease. Patients with HIV-associated nephropathy should be treated for their HIV infection regardless of CD4+ T cell count. Treatment with angiotensin-converting enzyme (ACE) inhibitors and/or prednisone, 60 mg/d, also has been reported to be of benefit in some cases. The incidence of this disease in patients receiving adequate cART has not been well defined; however, the impression is that it has decreased in frequency and severity. It is the leading cause of end-stage renal disease in patients with HIV infection.

Among the drugs commonly associated with renal damage in patients with HIV disease are pentamidine, amphotericin, adefovir, didanosine, tenofovir, and foscarnet. TMP/SMX may compete for tubular secretion with creatinine and cause an increase in the serum creatinine level. Sulfadiazine may crystallize in the kidney and result in an easily reversible form of renal shutdown, while indinavir or atazanavir may form renal calculi. Adequate hydration is the mainstay of treatment and prevention for these latter two conditions.

Genitourinary tract infections are seen with a high frequency in patients with HIV infection; they present with skin lesions, dysuria, hematuria, and/or pyuria and are managed in the same fashion as in patients without HIV infection. Infections with HSV are covered below (“Dermatologic Diseases”). Infections with *T. pallidum*, the etiologic agent of syphilis, play an important role in the HIV epidemic. In HIV-negative individuals, genital syphilitic ulcers as well as the ulcers of chancroid are major predisposing factors for heterosexual transmission of HIV infection. While most HIV-infected individuals with syphilis have a typical presentation, a variety of formerly rare clinical problems may be encountered in the setting of dual infection. Among them are *lues maligna*, an ulcerating lesion of the skin due to a necrotizing vasculitis; unexplained fever; nephrotic syndrome; and neurosyphilis. The most common presentation of syphilis in the HIV-infected patient is that of *condylomata lata*, a form of secondary syphilis. Neurosyphilis may be asymptomatic or may present as acute meningitis, neuroretinitis, deafness, or stroke. The rate of neurosyphilis may be as high as 1% in patients with HIV infection, and one should consider a lumbar puncture to look for neurosyphilis in all patients with HIV infection and secondary syphilis. As a consequence of the immunologic abnormalities seen in the setting of HIV infection, diagnosis of syphilis through standard serologic testing may be challenging. On the one hand, a significant number of patients have false-positive Venereal Disease Research Laboratory (VDRL) tests due to polyclonal B cell activation. On the other hand, the development of a new positive VDRL may be delayed in patients with new infections, and the anti-fluorescent treponemal antibody (anti-FTA) test may be negative due to immunodeficiency. Thus, dark-field examination of appropriate specimens should be performed in any patient in whom syphilis is suspected, even if the patient has a negative VDRL. Similarly, any patient with a positive serum VDRL test, neurologic findings, and an abnormal spinal fluid examination should be considered to have neurosyphilis and treated accordingly, regardless of the CSF VDRL result. In any setting, patients treated for syphilis need to be carefully monitored to ensure adequate therapy. Approximately one-third of patients with HIV infection will experience a Jarisch-Herxheimer reaction upon initiation of therapy for syphilis.

Vulvovaginal candidiasis is a common problem in women with HIV infection. Symptoms include pruritus, discomfort, dyspareunia, and dysuria. Vulvar infection may present as a morbilliform rash that

may extend to the thighs. Vaginal infection is usually associated with a white discharge, and plaques may be seen along an erythematous vaginal wall. Diagnosis is made by microscopic examination of the discharge for pseudohyphal elements in a 10% potassium hydroxide solution. Mild disease can be treated with topical therapy. More serious disease can be treated with fluconazole. Other causes of vaginitis include *Trichomonas* and mixed bacteria.

Diseases of the Endocrine System and Metabolic Disorders A variety of endocrine and metabolic disorders are seen in the context of HIV infection. These may be a direct consequence of HIV infection, secondary to opportunistic infections or neoplasms, or related to medication side effects. Between 33% and 75% of patients with HIV infection receiving thymidine analogues or protease inhibitors as a component of cART develop a syndrome often referred to as *lipodystrophy*, consisting of elevations in plasma triglycerides, total cholesterol, and apolipoprotein B, as well as hyperinsulinemia and hyperglycemia. Many of the patients have been noted to have a characteristic set of body habitus changes associated with fat redistribution, consisting of truncal obesity coupled with peripheral wasting (Fig. 226-38). Truncal obesity is apparent as an increase in abdominal girth related to increases in mesenteric fat, a dorsocervical fat pad (“buffalo hump”) reminiscent of patients with Cushing’s syndrome, and enlargement of the breasts. The peripheral wasting, or lipoatrophy, is particularly noticeable in the face and buttocks and by the prominence of the veins in the legs. These changes may develop at any time ranging from ~6 weeks to several years following the initiation of cART. Approximately 20% of the patients with HIV-associated lipodystrophy meet the criteria for the *metabolic syndrome* as defined by The International Diabetes Federation or The U.S. National Cholesterol Education Program Adult Treatment Panel III. The lipodystrophy syndrome has been reported in association with regimens containing a variety of different drugs, and while initially reported in the setting of protease inhibitor therapy, it appears that similar changes can also be induced by protease-sparing regimens. It has been suggested that the lipoatrophy changes are particularly severe in patients receiving the thymidine analogues stavudine and zidovudine. National Cholesterol Education Program (NCEP) guidelines should be followed in the management of these lipid abnormalities (Chap. 291e), and consideration should be given to changing the components of cART with avoidance of thymidine analogues (azidothymidine and stavudine) and protease inhibitors. Due to concerns regarding drug interactions, the most commonly utilized lipid-lowering agents in this setting are gemfibrozil and atorvastatin. In addition, lactic acidosis is associated with cART. This is most commonly seen with nucleoside analogue reverse transcriptase inhibitors and can be fatal (see below).

Patients with advanced HIV disease may develop hyponatremia due to the syndrome of inappropriate antidiuretic hormone (vasopressin) secretion (SIADH) as a consequence of increased free-water intake and decreased free-water excretion. SIADH is usually seen in conjunction with pulmonary or CNS disease. Low serum sodium may also be due to adrenal insufficiency; a concomitant high serum potassium should alert one to this possibility. Hyperkalemia may be secondary to adrenal insufficiency; HIV nephropathy; or medications, particularly trimethoprim and pentamidine. Hypokalemia may be seen in the setting of tenofovir or amphotericin therapy. Adrenal gland disease may be due to mycobacterial infections, CMV disease, cryptococcal disease, histoplasmosis, or ketoconazole toxicity. Iatrogenic Cushing’s syndrome with suppression of the hypothalamic-pituitary-adrenal axis may be seen with the use of local glucocorticoids (injected or inhaled) in patients receiving ritonavir. This is due to inhibition of the hepatic enzyme CYP3A4 by ritonavir leading to prolongation of the glucocorticoid half-life.

Thyroid function may be altered in 10–15% of patients with HIV infection. Both hypo- and hyperthyroidism may be seen. The predominant abnormality is subclinical hypothyroidism. In the setting of cART, up to 10% of patients have been noted to have elevated thyroid-stimulating hormone levels, suggesting that this may be

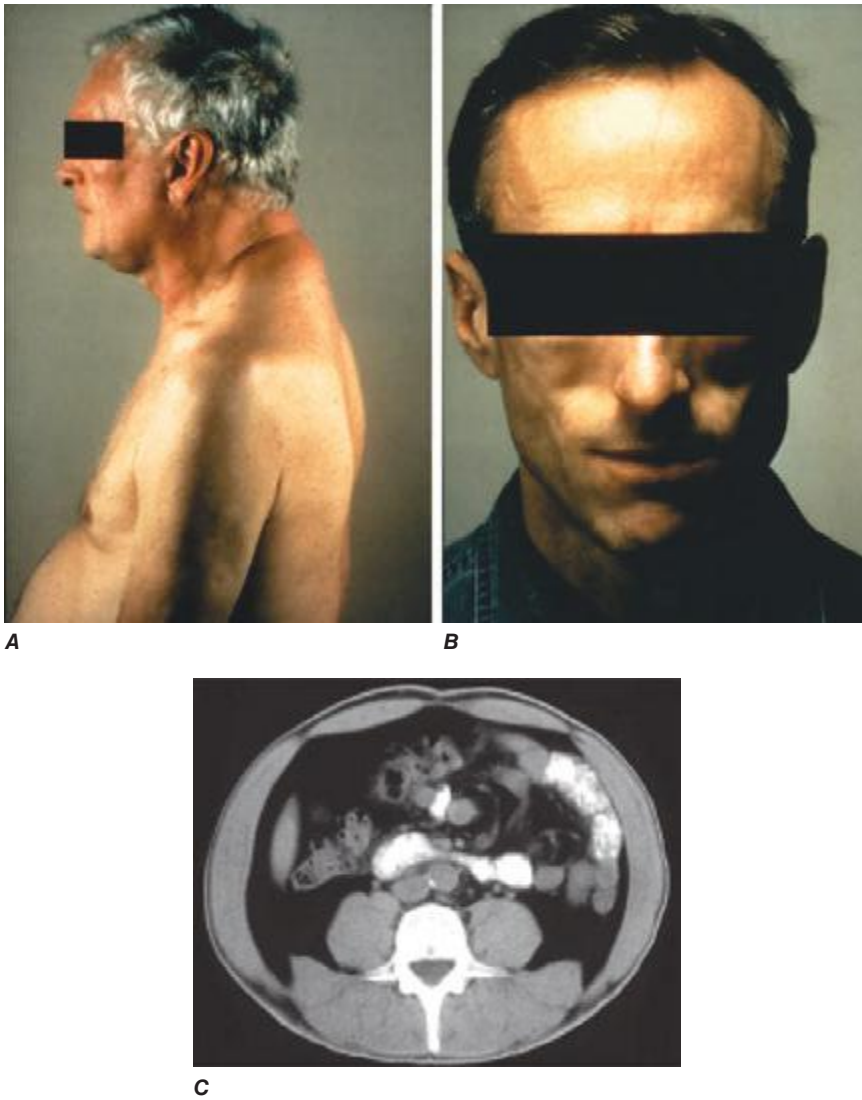


FIGURE 226-38 Characteristics of lipodystrophy. **A.** Truncal obesity and buffalo hump. **B.** Facial wasting. **C.** Accumulation of intraabdominal fat on CT scan.

a manifestation of immune reconstitution. Immune-reconstitution Graves' disease may occur as a late (9–48 months) complication of cART. In advanced HIV disease, infection of the thyroid gland may occur with opportunistic pathogens, including *P. jiroveci*, CMV, mycobacteria, *Toxoplasma gondii*, and *Cryptococcus neoformans*. These infections are generally associated with a nontender, diffuse enlargement of the thyroid gland. Thyroid function is usually normal. Diagnosis is made by fine-needle aspirate or open biopsy.

Depending on the severity of disease, HIV infection is associated with *hypogonadism* in 20–50% of men. While this is generally a complication of underlying illness, testicular dysfunction may also be a side effect of ganciclovir therapy. In some surveys, up to two-thirds of patients report decreased libido and one-third complain of erectile dysfunction. Androgen-replacement therapy should be considered in patients with symptomatic hypogonadism. HIV infection does not seem to have a significant effect on the menstrual cycle outside the setting of advanced disease.

Immunologic and Rheumatologic Diseases Immunologic and rheumatologic disorders are common in patients with HIV infection and range from excessive immediate-type hypersensitivity reactions (Chap. 376) to an increase in the incidence of reactive arthritis (Chap. 384) to conditions characterized by a diffuse infiltrative lymphocytosis. The occurrence of these phenomena is an apparent paradox in the setting of the profound immunodeficiency and immunosuppression that

characterizes HIV infection and reflects the complex nature of the immune system and its regulatory mechanisms.

Drug allergies are the most significant allergic reactions occurring in HIV-infected patients and appear to become more common as the disease progresses. They occur in up to 65% of patients who receive therapy with TMP/SMX for PCP. In general, these drug reactions are characterized by erythematous, morbilliform eruptions that are pruritic, tend to coalesce, and are often associated with fever. Nonetheless, ~33% of patients can be maintained on the offending therapy, and thus these reactions are not an immediate indication to stop the drug. Anaphylaxis is extremely rare in patients with HIV infection, and patients who have a cutaneous reaction during a single course of therapy can still be considered candidates for future treatment or prophylaxis with the same agent. The one exception to this is the nucleoside analogue abacavir, where fatal hypersensitivity reactions have been reported with rechallenge. This hypersensitivity is strongly associated with the HLA-B5701 haplotype, and a hypersensitivity reaction to abacavir is an absolute contraindication to future therapy. For other agents, including TMP/SMX, desensitization regimens are moderately successful. While the mechanisms underlying these allergic-type reactions remain unknown, patients with HIV infection have been noted to have elevated IgE levels that increase as the CD4+ T cell count declines. The numerous examples of patients with multiple drug reactions suggest that a common pathway is involved.

HIV infection shares many similarities with a variety of autoimmune diseases, including a substantial polyclonal B cell activation that is associated with a high incidence of antiphospholipid antibodies, such as anticardiolipin antibodies, VDRL antibodies, and lupus-like anticoagulants. In addition, HIV-infected individuals have an increased incidence of antinuclear antibodies. Despite these serologic findings, there is no evidence that HIV-infected individuals have an

increase in two of the more common autoimmune diseases, i.e., systemic lupus erythematosus and rheumatoid arthritis. In fact, it has been observed that these diseases may be somewhat ameliorated by the concomitant presence of HIV infection, suggesting that an intact CD4+ T cell limb of the immune response plays an integral role in the pathogenesis of these conditions. Similarly, there are anecdotal reports of patients with common variable immunodeficiency (Chap. 374), characterized by hypogammaglobulinemia, who have had a normalization of Ig levels following the development of HIV infection, suggesting a possible role for overactive CD4+ T cell immunity in certain forms of that syndrome. The one autoimmune disease that may occur with an increased frequency in patients with HIV infection is a variant of primary Sjögren's syndrome (Chap. 383). Patients with HIV infection may develop a syndrome consisting of parotid gland enlargement, dry eyes, and dry mouth that is associated with lymphocytic infiltrates of the salivary gland and lung. One also can see peripheral neuropathy, polymyositis, renal tubular acidosis, and hepatitis. In contrast to Sjögren's syndrome, in which the lymphocytic infiltrates are composed predominantly of CD4+ T cells, in patients with HIV infection the infiltrates are composed predominantly of CD8+ T cells. In addition, while patients with Sjögren's syndrome are mainly women who have autoantibodies to Ro and La and who frequently have HLA-DR3 or -B8 MHC haplotypes, HIV-infected individuals with this syndrome are usually African-American men who do not have anti-Ro or anti-La and who

most often are HLA-DR5. This syndrome appears to be less common with the increased use of effective cART. The term *diffuse infiltrative lymphocytosis syndrome* (DILS) is used to describe this entity and to distinguish it from Sjögren's syndrome.

Approximately one-third of HIV-infected individuals experience arthralgias; furthermore, 5–10% are diagnosed as having some form of reactive arthritis, such as Reiter's syndrome or psoriatic arthritis as well as undifferentiated spondyloarthropathy (Chap. 384). These syndromes occur with increasing frequency as the competency of the immune system declines. This association may be related to an increase in the number of infections with organisms that may trigger a reactive arthritis with progressive immunodeficiency or to a loss of important regulatory T cells. Reactive arthritides in HIV-infected individuals generally respond well to standard treatment; however, therapy with methotrexate has been associated with an increase in the incidence of opportunistic infections and should be used with caution and only in severe cases.

HIV-infected individuals also experience a variety of joint problems without obvious cause that are referred to generically as *HIV- or AIDS-associated arthropathy*. This syndrome is characterized by subacute oligoarticular arthritis developing over a period of 1–6 weeks and lasting 6 weeks to 6 months. It generally involves the large joints, predominantly the knees and ankles, and is nonerosive with only a mild inflammatory response. X-rays are nonrevealing. Nonsteroidal anti-inflammatory drugs are only marginally helpful; however, relief has been noted with the use of intraarticular glucocorticoids. A second form of arthritis also thought to be secondary to HIV infection is called *painful articular syndrome*. This condition, reported as occurring in as many as 10% of AIDS patients, presents as an acute, severe, sharp pain in the affected joint. It affects primarily the knees, elbows, and shoulders; lasts 2–24 h; and may be severe enough to require narcotic analgesics. The cause of this arthropathy is unclear; however, it is thought to result from a direct effect of HIV on the joint. This condition is reminiscent of the fact that other lentiviruses, in particular the caprine arthritis-encephalitis virus, are capable of directly causing arthritis.

A variety of other immunologic or rheumatologic diseases have been reported in HIV-infected individuals, either de novo or in association with opportunistic infections or drugs. Using the criteria of widespread musculoskeletal pain of at least 3 months' duration and the presence of at least 11 of 18 possible tender points by digital palpation, 11% of an HIV-infected cohort containing 55% IDUs were diagnosed as having *fibromyalgia* (Chap. 396). While the incidence of frank arthritis was less in this population than in other studied populations that consisted predominantly of men who have sex with men, these data support the concept that there are musculoskeletal problems that occur as a direct result of HIV infection. In addition there have been reports of leukocytoclastic vasculitis in the setting of zidovudine therapy. CNS angitis and polymyositis also have been reported in HIV-infected individuals. Septic arthritis is surprisingly rare, especially given the increased incidence of staphylococcal bacteremias seen in this population. When septic arthritis has been reported, it has usually been due to *Staphylococcus aureus*, systemic fungal infection with *C. neoformans*, *Sporothrix schenckii*, or *H. capsulatum* or to systemic mycobacterial infection with *M. tuberculosis*, *M. haemophilum*, *M. avium*, or *M. kansasii*.

Patients with HIV infection treated with cART have been found to have an increased incidence of osteonecrosis or avascular necrosis of the hip and shoulders. In a study of asymptomatic patients, 4.4% were found to have evidence of osteonecrosis on MRI. While precise cause-and-effect relationships have been difficult to establish, this complication has been associated with the use of lipid-lowering agents, systemic glucocorticoids, and testosterone; bodybuilding exercise; alcohol consumption; and the presence of anticardiolipin antibodies. Osteoporosis has been reported in 7% of women with HIV infection, with 41% of women demonstrating some degree of osteopenia. Several studies have documented decreases in bone mineral density of 2–6% in the first 2 years following the initiation of cART. This may be particularly apparent with tenofovir-containing regimens.

TABLE 226-12 CHARACTERISTICS OF IMMUNE RECONSTITUTION INFLAMMATORY SYNDROME (IRIS)

- Paradoxical worsening of an existing clinical condition or abrupt appearance of a new clinical finding (unmasking) is seen following the initiation of antiretroviral therapy
- Occurs weeks to months following the initiation of antiretroviral therapy
- Is most common in patients starting therapy with a CD4+ T cell count <50/μL who experience a precipitous drop in viral load
- Is frequently seen in the setting of tuberculosis; particularly when cART is starting soon after initiation of anti-TB therapy
- Can be fatal

Immune Reconstitution Inflammatory Syndrome (IRIS)

Following the initiation of effective cART, a paradoxical worsening of preexisting, untreated, or partially treated opportunistic infections may be noted. One may also see exacerbations of pre-existing or the development of new autoimmune conditions following the initiation of antiretrovirals (Table 226-12). IRIS related to a known pre-existing infection or neoplasm is referred to as *paradoxical IRIS*, while IRIS associated with a previously undiagnosed condition is referred to as *unmasking IRIS*. The term *immune reconstitution disease (IRD)* is sometimes used to distinguish IRIS manifestations related to opportunistic diseases from IRIS manifestations related to autoimmune diseases. IRD is particularly common in patients with underlying untreated mycobacterial or fungal infections. IRIS is seen in 10–30% of patients, depending on the clinical setting, and is most common in patients starting therapy with CD4+ T cell counts <50 cells/μL who have a precipitous drop in HIV RNA levels following the initiation of cART. Signs and symptoms may appear anywhere from 2 weeks to 2 years after the initiation of cART and can include localized lymphadenitis, prolonged fever, pulmonary infiltrates, hepatitis, increased intracranial pressure, uveitis, sarcoidosis, and Graves' disease. The clinical course can be protracted, and severe cases can be fatal. The underlying mechanism appears to be related to a phenomenon similar to type IV hypersensitivity reactions and reflects the immediate improvements in immune function that occur as levels of HIV RNA drop and the immunosuppressive effects of HIV infection are controlled. In severe cases, the use of immunosuppressive drugs such as glucocorticoids may be required to blunt the inflammatory component of these reactions while specific antimicrobial therapy takes effect.

Diseases of the Hematopoietic System Disorders of the hematopoietic system including lymphadenopathy, anemia, leukopenia, and/or thrombocytopenia are common throughout the course of HIV infection and may be the direct result of HIV, manifestations of secondary infections and neoplasms, or side effects of therapy (Table 226-13). Direct histologic examination and culture of lymph node or bone marrow tissue are often diagnostic. A significant percentage of bone marrow aspirates from patients with HIV infection have been reported to contain lymphoid aggregates, the precise significance of which is unknown. Initiation of cART will lead to reversal of most hematologic complications that are the direct result of HIV infection.

TABLE 226-13 CAUSES OF BONE MARROW SUPPRESSION IN PATIENTS WITH HIV INFECTION

HIV infection	Medications
Mycobacterial infections	Zidovudine
Fungal infections	Dapsone
B19 parvovirus infection	Trimethoprim/sulfamethoxazole
Lymphoma	Pyrimethamine
	5-Flucytosine
	Ganciclovir
	Interferon α
	Trimetrexate
	Foscarnet

Some patients, otherwise asymptomatic, may develop *persistent generalized lymphadenopathy* as an early clinical manifestation of HIV infection. This condition is defined as the presence of enlarged lymph nodes (>1 cm) in two or more extralingual sites for >3 months without an obvious cause. The lymphadenopathy is due to marked follicular hyperplasia in the node in response to HIV infection. The nodes are generally discrete and freely movable. This feature of HIV disease may be seen at any point in the spectrum of immune dysfunction and is not associated with an increased likelihood of developing AIDS. Paradoxically, a loss in lymphadenopathy or a decrease in lymph node size outside the setting of cART may be a prognostic marker of disease progression. In patients with CD4+ T cell counts >200/ μL , the differential diagnosis of lymphadenopathy includes KS, TB, Castleman's disease, and lymphoma. In patients with more advanced disease, lymphadenopathy may also be due to atypical mycobacterial infection, toxoplasmosis, systemic fungal infection, or bacillary angiomatosis. While indicated in patients with CD4+ T cell counts <200/ μL , lymph node biopsy is not indicated in patients with early-stage disease unless there are signs and symptoms of systemic illness, such as fever and weight loss, or unless the nodes begin to enlarge, become fixed, or coalesce. Monoclonal gammopathy of unknown significance (MGUS) (Chap. 136), defined as the presence of a serum monoclonal IgG, IgA, or IgM in the absence of a clear cause, has been reported in 3% of patients with HIV infection. The overall clinical significance of this finding in patients with HIV infection is unclear, although it has been associated with other viral infections, non-Hodgkin's lymphoma, and plasma cell malignancy.

Anemia is the most common hematologic abnormality in HIV-infected patients and, in the absence of a specific treatable cause, is independently associated with a poor prognosis. While generally mild, anemia can be quite severe and require chronic blood transfusions. Among the specific reversible causes of anemia in the setting of HIV infection are drug toxicity, systemic fungal and mycobacterial infections, nutritional deficiencies, and parvovirus B19 infections. Zidovudine may block erythroid maturation prior to its effects on other marrow elements. A characteristic feature of zidovudine therapy is an elevated mean corpuscular volume (MCV). Another drug used in patients with HIV infection that has a selective effect on the erythroid series is dapson. This drug can cause a serious hemolytic anemia in patients who are deficient in glucose-6-phosphate dehydrogenase and can create a functional anemia in others through induction of methemoglobinemia. Folate levels are usually normal in HIV-infected individuals; however, vitamin B₁₂ levels may be depressed as a consequence of achlorhydria or malabsorption. True autoimmune hemolytic anemia is rare, although ~20% of patients with HIV infection may have a positive direct antiglobulin test as a consequence of polyclonal B cell activation. Infection with parvovirus B19 may also cause anemia. It is important to recognize this possibility given the fact that it responds well to treatment with IVIg. Erythropoietin levels in patients with HIV infection and anemia are generally lower than expected given the degree of anemia. Treatment with erythropoietin may result in an increase in hemoglobin levels. An exception to this is a subset of patients with zidovudine-associated anemia in whom erythropoietin levels may be quite high.

During the course of HIV infection, neutropenia may be seen in approximately half of patients. In most instances it is mild; however, it can be severe and can put patients at risk of spontaneous bacterial infections. This is most frequently seen in patients with severely advanced HIV disease and in patients receiving any of a number of potentially myelosuppressive therapies. In the setting of neutropenia, diseases that are not commonly seen in HIV-infected patients, such as aspergillosis or mucormycosis, may occur. Both granulocyte colony-stimulating factor (G-CSF) and GM-CSF increase neutrophil counts in patients with HIV infection regardless of the cause of the neutropenia. Earlier concerns about the potential of these agents to also increase levels of HIV were not confirmed in controlled clinical trials.

Thrombocytopenia may be an early consequence of HIV infection. Approximately 3% of patients with untreated HIV infection and CD4+ T cell counts $\geq 400/\mu\text{L}$ have platelet counts <150,000/ μL . For untreated

patients with CD4+ T cell counts <400/ μL , this incidence increases to 10%. In patients receiving antiretrovirals, thrombocytopenia is associated with hepatitis C, cirrhosis, and ongoing high-level HIV replication. Thrombocytopenia is rarely a serious clinical problem in patients with HIV infection and generally responds well to successful cART. Clinically, it resembles the thrombocytopenia seen in patients with idiopathic thrombocytopenic purpura (Chap. 140). Immune complexes containing anti-gp120 antibodies and anti-anti-gp120 antibodies have been noted in the circulation and on the surface of platelets in patients with HIV infection. Patients with HIV infection have also been noted to have a platelet-specific antibody directed toward a 25-kDa component of the surface of the platelet. Other data suggest that the thrombocytopenia in patients with HIV infection may be due to a direct effect of HIV on megakaryocytes. Whatever the cause, it is very clear that the most effective medical approach to this problem has been the use of cART. For patients with platelet counts <20,000/ μL , a more aggressive approach combining IVIg or anti-Rh Ig for an immediate response and cART for a more lasting response is appropriate. Rituximab has been used with some success in otherwise refractory cases. Splenectomy is a rarely needed option and is reserved for patients refractory to medical management. Because of the risk of serious infection with encapsulated organisms, all patients with HIV infection about to undergo splenectomy should be immunized with pneumococcal polysaccharide. It should be noted that, in addition to causing an increase in the platelet count, removal of the spleen will result in an increase in the peripheral blood lymphocyte count, making CD4+ T cell counts unreliable markers of immunocompetence. In this setting, the clinician should rely on the CD4+ T cell percentage for making diagnostic decisions with respect to the likelihood of opportunistic infections. A CD4+ T cell percentage of 15 is approximately equivalent to a CD4+ T cell count of 200/ μL . In patients with early HIV infection, thrombocytopenia has also been reported as a consequence of classic thrombotic thrombocytopenic purpura (Chap. 140). This clinical syndrome, consisting of fever, thrombocytopenia, hemolytic anemia, and neurologic and renal dysfunction, is a rare complication of early HIV infection. As in other settings, the appropriate management is the use of salicylates and plasma exchange. Other causes of thrombocytopenia include lymphoma, mycobacterial infections, and fungal infections.

The incidence of venous thromboembolic disease such as deep-vein thrombosis or pulmonary embolus is approximately 1% per year in patients with HIV infection. This is approximately 10 times higher than that seen in an age-matched population. Factors associated with an increased risk of clinical thrombosis include age over 45, history of an opportunistic infection, lower CD4 count, and estrogen use. Abnormalities of the coagulation cascade including decreased protein S activity, increases in factor VIII, anticardiolipin antibodies, or lupus-like anticoagulant have been reported in more than 50% of patients with HIV infection. The clinical significance of this increased propensity toward thromboembolic disease is likely reflected in the observation that elevations in D-dimer are strongly associated with all-cause mortality in patients with HIV infection (Table 226-9).

Dermatologic Diseases Dermatologic problems occur in >90% of patients with HIV infection. From the macular, roseola-like rash seen with the acute seroconversion syndrome to extensive end-stage KS, cutaneous manifestations of HIV disease can be seen throughout the course of HIV infection. Among the more common nonneoplastic problems are seborrheic dermatitis, folliculitis, and opportunistic infections. Extrapulmonary pneumocystosis may cause a necrotizing vasculitis. Neoplastic conditions are covered below.

Seborrheic dermatitis occurs in 3% of the general population and in up to 50% of patients with HIV infection. Seborrheic dermatitis increases in prevalence and severity as the CD4+ T cell count declines. In HIV-infected patients, seborrheic dermatitis may be aggravated by concomitant infection with *Pityrosporum*, a yeastlike fungus; use of topical antifungal agents has been recommended in cases refractory to standard topical treatment.

Folliculitis is among the most prevalent dermatologic disorders in patients with HIV infection and is seen in ~20% of patients. It is

more common in patients with CD4+ T cell counts <200 cells/ μ L. Pruritic papular eruption is one of the most common pruritic rashes in patients with HIV infection. It appears as multiple papules on the face, trunk, and extensor surfaces and may improve with cART. *Eosinophilic pustular folliculitis* is a rare form of folliculitis that is seen with increased frequency in patients with HIV infection. It presents as multiple, urticarial perifollicular papules that may coalesce into plaquelike lesions. Skin biopsy reveals an eosinophilic infiltrate of the hair follicle, which in certain cases has been associated with the presence of a mite. Patients typically have an elevated serum IgE level and may respond to treatment with topical anthelmintics. Pruritus is a common symptom in patients with HIV infection and can lead to prurigo nodularis. Patients with HIV infection have also been reported to develop a severe form of *Norwegian scabies* with hyperkeratotic psoriasiform lesions.

Both *psoriasis* and *ichthyosis*, although they are not reported to be increased in frequency, may be particularly severe when they occur in patients with HIV infection. Preexisting psoriasis may become guttate in appearance and more refractory to treatment in the setting of HIV infection.

Reactivation herpes zoster (shingles) is seen in 10–20% of patients with HIV infection. This reactivation syndrome of varicella-zoster virus indicates a modest decline in immune function and may be the first indication of clinical immunodeficiency. In one series, patients who developed shingles did so an average of 5 years after HIV infection. In a cohort of patients with HIV infection and localized zoster, the subsequent rate of the development of AIDS was 1% per month. In that study, AIDS was more likely to develop if the outbreak of zoster was associated with severe pain, extensive skin involvement, or involvement of cranial or cervical dermatomes. The clinical manifestations of reactivation zoster in HIV-infected patients, although indicative of immunologic compromise, are not as severe as those seen in other immunodeficient conditions. Thus, while lesions may extend over several dermatomes, involve the spinal cord, and/or be associated with frank cutaneous dissemination, visceral involvement has not been reported. In contrast to patients without a known underlying immunodeficiency state, patients with HIV infection tend to have recurrences of zoster with a relapse rate of ~20%. Valacyclovir, acyclovir, or famciclovir is the treatment of choice. Foscarnet may be of value in patients with acyclovir-resistant virus.

Infection with *herpes simplex virus* in HIV-infected individuals is associated with recurrent orolabial, genital, and perianal lesions as part of recurrent reactivation syndromes (Chap. 216). As HIV disease progresses and the CD4+ T cell count declines, these infections become more frequent and severe. Lesions often appear as beefy red, are exquisitely painful, and have a tendency to occur high in the gluteal cleft (Fig. 226-37). Perirectal HSV may be associated with proctitis and anal fissures. HSV should be high in the differential diagnosis of any HIV-infected patient with a poorly healing, painful perirectal lesion. In addition to recurrent mucosal ulcers, recurrent HSV infection in the form of *herpetic whitlow* can be a problem in patients with HIV infection, presenting with painful vesicles or extensive cutaneous erosion. Valacyclovir, acyclovir or famciclovir is the treatment of choice in these settings. It is noteworthy that even subclinical reactivation of herpes simplex may be associated with increases in plasma HIV RNA levels.

Diffuse skin eruptions due to *Molluscum contagiosum* may be seen in patients with advanced HIV infection. These flesh-colored, umbilicated lesions may be treated with local therapy. They tend to regress with effective cART. Similarly, *condyloma acuminatum* lesions may be more severe and more widely distributed in patients with low CD4+ T cell counts. Imiquimod cream may be helpful in some cases. Atypical mycobacterial infections may present as erythematous cutaneous nodules, as may fungal infections, *Bartonella*, *Acanthamoeba*, and KS. Cutaneous infections with *Aspergillus* have been noted at the site of IV catheter placement.

The skin of patients with HIV infection is often a target organ for drug reactions (Chap. 74). Although most skin reactions are mild and not necessarily an indication to discontinue therapy, patients may

have particularly severe cutaneous reactions, including erythroderma, *Stevens-Johnson syndrome*, and toxic epidermal necrolysis, as a reaction to drugs—particularly sulfa drugs, nonnucleoside reverse transcriptase inhibitors, abacavir, amprenavir, darunavir, fosamprenavir, and tipranavir. Similarly, patients with HIV infection are often quite photosensitive and burn easily following exposure to sunlight or as a side effect of radiation therapy (Chap. 75).

HIV infection and its treatment may be accompanied by cosmetic changes of the skin that are not of great clinical importance but may be troubling to patients. Yellowing of the nails and straightening of the hair, particularly in African-American patients, have been reported as a consequence of HIV infection. Zidovudine therapy has been associated with elongation of the eyelashes and the development of a bluish discoloration to the nails, again more common in African-American patients. Therapy with clofazimine may cause a yellow-orange discoloration of the skin and urine.

Neurologic Diseases Clinical disease of the nervous system accounts for a significant degree of morbidity in a high percentage of patients with HIV infection (Table 226-14). The neurologic problems that occur in HIV-infected individuals may be either primary to the pathogenic processes of HIV infection or secondary to opportunistic infections or neoplasms. Among the more frequent opportunistic diseases that involve the CNS are toxoplasmosis, cryptococcosis, progressive multifocal leukoencephalopathy, and primary CNS lymphoma. Other less common problems include mycobacterial infections; syphilis; and infection with CMV, HTLV-1, *Trypanosoma cruzi*, or *Acanthamoeba*. Overall, secondary diseases of the CNS have been reported to occur in approximately one-third of patients with AIDS. These data antedate the widespread use of cART, and this frequency is considerably lower in patients receiving effective antiretroviral drugs. Primary processes related to HIV infection of the nervous system are reminiscent of those seen with other lentiviruses, such as the Visna-Maedi virus of sheep.

Neurologic problems directly attributable to HIV occur throughout the course of infection and may be inflammatory, demyelinating, or degenerative in nature. The term *HIV-associated neurocognitive disorders* (HAND) is used to describe a spectrum of disorders that range from asymptomatic neurocognitive impairment (ANI) to minor neurocognitive disorder (MND) to clinically severe dementia. The most severe form, *HIV-associated dementia* (HAD), also referred to as the *AIDS dementia complex*, or *HIV encephalopathy*, is considered an AIDS-defining illness. Most HIV-infected patients have some neurologic problem during the course of their disease. Even in the setting of suppressive cART, approximately 50% of HIV-infected individuals can be shown to have mild to moderate neurocognitive impairment using sensitive neuropsychiatric testing. As noted in the section on

TABLE 226-14 NEUROLOGIC DISEASES IN PATIENTS WITH HIV INFECTION

Opportunistic infections	HIV-1 infection
Toxoplasmosis	Aseptic meningitis
Cryptococcosis	HIV-associated neurocognitive disorders (HAND), including HIV encephalopathy/AIDS dementia complex
Progressive multifocal leukoencephalopathy	Myelopathy
Cytomegalovirus	Vacuolar myelopathy
Syphilis	Pure sensory ataxia
<i>Mycobacterium tuberculosis</i>	Paresthesia/dysesthesia
HTLV-1 infection	Peripheral neuropathy
Amebiasis	Acute inflammatory demyelinating polyneuropathy (Guillain-Barré syndrome)
Neoplasms	Chronic inflammatory demyelinating polyneuropathy (CIDP)
Primary CNS lymphoma	Mononeuritis multiplex
Kaposi's sarcoma	Distal symmetric polyneuropathy
	Myopathy

1264 pathogenesis, damage to the CNS may be a direct result of viral infection of the CNS macrophages or glial cells or may be secondary to the release of neurotoxins and potentially toxic cytokines such as IL-1 β , TNF- α , IL-6, and TGF- β . It has been reported that HIV-infected individuals with the E4 allele for apoE are at increased risk for AIDS encephalopathy and peripheral neuropathy. Virtually all patients with HIV infection have some degree of nervous system involvement with the virus. This is evidenced by the fact that CSF findings are abnormal in ~90% of patients, even during the asymptomatic phase of HIV infection. CSF abnormalities include pleocytosis (50–65% of patients), detection of viral RNA (~75%), elevated CSF protein (35%), and evidence of intrathecal synthesis of anti-HIV antibodies (90%). It is important to point out that evidence of infection of the CNS with HIV does not imply impairment of cognitive function. The neurologic function of an HIV-infected individual should be considered normal unless clinical signs and symptoms suggest otherwise.

Aseptic meningitis may be seen in any but the very late stages of HIV infection. In the setting of acute primary infection, patients may experience a syndrome of headache, photophobia, and meningismus. Rarely, an acute encephalopathy due to encephalitis may occur. Cranial nerve involvement may be seen, predominantly cranial nerve VII but occasionally V and/or VIII. CSF findings include a lymphocytic pleocytosis, elevated protein level, and normal glucose level. This syndrome, which cannot be clinically differentiated from other viral meningitides (Chap. 165), usually resolves spontaneously within 2–4 weeks; however, in some patients, signs and symptoms may become chronic. Aseptic meningitis may occur any time in the course of HIV infection; however, it is rare following the development of AIDS. This suggests that clinical aseptic meningitis in the context of HIV infection is an immune-mediated disease.

Cryptococcus is the leading infectious cause of meningitis in patients with AIDS (Chap. 239). While the vast majority of these are due to *C. neoformans*, up to 12% may be due to *C. gattii*. Cryptococcal meningitis is the initial AIDS-defining illness in ~2% of patients and generally occurs in patients with CD4+ T cell counts <100/ μ L. Cryptococcal meningitis is particularly common in untreated patients with AIDS in Africa, occurring in ~5% of patients. Most patients present with a picture of subacute meningoencephalitis with fever, nausea, vomiting, altered mental status, headache, and meningeal signs. The incidence of seizures and focal neurologic deficits is low. The CSF profile may be normal or may show only modest elevations in WBC or protein levels and decreases in glucose. The opening pressure in the CSF is usually elevated. In addition to meningitis, patients may develop cryptococomas and cranial nerve involvement. Approximately one-third of patients also have pulmonary disease. Uncommon manifestations of cryptococcal infection include skin lesions that resemble *molluscum contagiosum*, lymphadenopathy, palatal and glossal ulcers, arthritis, gastroenteritis, myocarditis, and prostatitis. The prostate gland may serve as a reservoir for smoldering cryptococcal infection. The diagnosis of cryptococcal meningitis is made by identification of organisms in spinal fluid with india ink examination or by the detection of cryptococcal antigen. Blood cultures for fungus are often positive. A biopsy may be needed to make a diagnosis of CNS cryptococcoma. Treatment is with IV amphotericin B 0.7 mg/kg daily, or liposomal amphotericin 4–6 mg/kg daily, with flucytosine 25 mg/kg qid for at least 2 weeks if possible, continuing with amphotericin alone ideally until the CSF culture turns negative. Decreases in renal function in association with amphotericin can lead to increases in flucytosine levels and subsequent bone marrow suppression. Amphotericin is followed by fluconazole 400 mg/d PO for 8 weeks, and then fluconazole 200 mg/d until the CD4+ T cell count has increased to >200 cells/ μ L for 6 months in response to cART. Repeated lumbar puncture may be required to manage increased intracranial pressure. Symptoms may recur with initiation of cART as an immune reconstitution syndrome (see above). Other fungi that may cause meningitis in patients with HIV infection are *C. immitis* and *H. capsulatum*. Meningoencephalitis has also been reported due to *Acanthamoeba* or *Naegleria*.

HIV-associated dementia consists of a constellation of signs and symptoms of CNS disease. While this is generally a late complication

of HIV infection that progresses slowly over months, it can be seen in patients with CD4+ T cell counts >350 cells/ μ L. A major feature of this entity is the development of dementia, defined as a decline in cognitive ability from a previous level. It may present as impaired ability to concentrate, increased forgetfulness, difficulty reading, or increased difficulty performing complex tasks. Initially these symptoms may be indistinguishable from findings of situational depression or fatigue. In contrast to “cortical” dementia (such as Alzheimer’s disease), aphasia, apraxia, and agnosia are uncommon, leading some investigators to classify HIV encephalopathy as a “subcortical dementia” characterized by defects in short-term memory and executive function (see below). In addition to dementia, patients with HIV encephalopathy may also have motor and behavioral abnormalities. Among the motor problems are unsteady gait, poor balance, tremor, and difficulty with rapid alternating movements. Increased tone and deep tendon reflexes may be found in patients with spinal cord involvement. Late stages may be complicated by bowel and/or bladder incontinence. Behavioral problems include apathy, irritability, and lack of initiative, with progression to a vegetative state in some instances. Some patients develop a state of agitation or mild mania. These changes usually occur without significant changes in level of alertness. This is in contrast to the finding of somnolence in patients with dementia due to toxic/metabolic encephalopathies.

HIV-associated dementia is the initial AIDS-defining illness in ~3% of patients with HIV infection and thus only rarely precedes clinical evidence of immunodeficiency. Clinically significant encephalopathy eventually develops in ~25% of untreated patients with AIDS. As immunologic function declines, the risk and severity of HIV-associated dementia increases. Autopsy series suggest that 80–90% of patients with HIV infection have histologic evidence of CNS involvement. Several classification schemes have been developed for grading HIV encephalopathy; a commonly used clinical staging system is outlined in Table 226-15.

The precise cause of HIV-associated dementia remains unclear, although the condition is thought to be a result of a combination of direct effects of HIV on the CNS and associated immune activation. HIV has been found in the brains of patients with HIV encephalopathy by Southern blot, in situ hybridization, PCR, and electron microscopy. Multinucleated giant cells, macrophages, and microglial cells appear to be the main cell types harboring virus in the CNS. Histologically, the major changes are seen in the subcortical areas of the brain and include pallor and gliosis, multinucleated giant cell encephalitis, and vacuolar myelopathy. Less commonly, diffuse or focal spongiform changes occur in the white matter. Areas of the brain involved in motor function, language, and judgment are most severely affected.

There are no specific criteria for a diagnosis of HIV-associated dementia, and this syndrome must be differentiated from a number of other diseases that affect the CNS of HIV-infected patients (Table 226-14). The diagnosis of dementia depends on demonstrating a decline in

TABLE 226-15 CLINICAL STAGING OF HAND ACCORDING TO FRASCATI CRITERIA

Stage	Neurocognitive Status ^a	Functional Status ^b
Asymptomatic	1 SD below mean in 2 cognitive domains	No impairments in activities of daily living
Mild neurocognitive disorder	1 SD below mean in 2 cognitive domains	Impairments in activities of daily living
HIV-associated dementia	2 SD below mean in 2 cognitive domains	Notable impairments in activities of daily living

^aNeurocognitive testing should include assessment of at least 5 domains, including attention-information processing, language, abstraction-executive, complex perceptual motor skills, memory (including learning and recall), simple motor skills, or sensory perceptual skills. Appropriate norms must be available to establish the number of domains in which performance is below 1 SD. ^bFunctional status is typically assessed by self-reporting but might be corroborated by a collateral source. No agreed measures exist for HIV-associated neurocognitive disorder criteria. Note that, for diagnosis of HIV-associated neurocognitive disorder, other causes of dementia must be ruled out and potential confounding effects of substance use or psychiatric illness should be considered.

Source: Adapted from A Antinori et al: Neurology 69:1789, 2007.



FIGURE 226-39 AIDS dementia complex. Postcontrast CT scan through the lateral ventricles of a 47-year-old man with AIDS, altered mental status, and dementia. The lateral and third ventricles and the cerebral sulci are abnormally prominent. Mild white matter hypodensity is seen adjacent to the frontal horns of the lateral ventricles.

cognitive function. This can be accomplished objectively with the use of a Mini-Mental Status Examination (MMSE) in patients for whom prior scores are available. For this reason, it is advisable for all patients with a diagnosis of HIV infection to have a baseline MMSE. However, changes in MMSE scores may be absent in patients with mild HIV encephalopathy. Imaging studies of the CNS, by either MRI or CT, often demonstrate evidence of cerebral atrophy (Fig. 226-39). MRI may also reveal small areas of increased density on T2-weighted images. Lumbar puncture is an important element of the evaluation of patients with HIV infection and neurologic abnormalities. It is generally most helpful in ruling out or making a diagnosis of opportunistic infections. In HIV encephalopathy, patients may have the nonspecific findings of an increase in CSF cells and protein level. While HIV RNA can often be detected in the spinal fluid and HIV can be cultured from the CSF, this finding is not specific for HIV encephalopathy. There appears to be no correlation between the presence of HIV in the CSF and the presence of HIV encephalopathy. Elevated levels of macrophage chemoattractant protein (MCP-1), β_2 -microglobulin, neopterin, and quinolinic acid (a metabolite of tryptophan reported to cause CNS injury) have been noted in the CSF of patients with HIV encephalopathy. These findings suggest that these factors as well as inflammatory cytokines may be involved in the pathogenesis of this syndrome.

Combination antiretroviral therapy is of benefit in patients with HIV-associated dementia. Improvement in neuropsychiatric test scores has been noted for both adult and pediatric patients treated with antiretrovirals. The rapid improvement in cognitive function noted with the initiation of cART suggests that at least some component of this problem is quickly reversible, again supporting at least a partial role of soluble mediators in the pathogenesis. It should also be noted that these patients have an increased sensitivity to the side effects of neuroleptic drugs. The use of these drugs for symptomatic treatment is associated with an increased risk of extrapyramidal side effects; therefore, patients with HIV encephalopathy who receive these agents must be monitored carefully. It is felt by many physicians that the decrease in the prevalence of severe cases of HAND brought about by cART has resulted in an increase in the prevalence of milder forms of this disorder.

Seizures may be a consequence of opportunistic infections, neoplasms, or HIV encephalopathy (Table 226-16). The seizure threshold is often lower than normal in patients with advanced HIV infection due in part to the frequent presence of electrolyte abnormalities. Seizures are seen in 15–40% of patients with cerebral toxoplasmosis,

TABLE 226-16 CAUSES OF SEIZURES IN PATIENTS WITH HIV INFECTION

Disease	Overall Contribution to First Seizure, %	Fraction of Patients Who Have Seizures, %
HIV encephalopathy	24–47	7–50
Cerebral toxoplasmosis	28	15–40
Cryptococcal meningitis	13	8
Primary central nervous system lymphoma	4	15–30
Progressive multifocal leukoencephalopathy	1	20

Source: From DM Holtzman et al: Am J Med 87:173, 1989.

15–35% of patients with primary CNS lymphoma, 8% of patients with cryptococcal meningitis, and 7–50% of patients with HIV encephalopathy. Seizures may also be seen in patients with CNS tuberculosis, aseptic meningitis, and progressive multifocal leukoencephalopathy. Seizures may be the presenting clinical symptom of HIV disease. In one study of 100 patients with HIV infection presenting with a first seizure, cerebral mass lesions were the most common cause, responsible for 32 of the 100 new-onset seizures. Of these 32 cases, 28 were due to toxoplasmosis and 4 to lymphoma. HIV encephalopathy accounted for an additional 24 new-onset seizures. Cryptococcal meningitis was the third most common diagnosis, responsible for 13 of the 100 seizures. In 23 cases, no cause could be found, and it is possible that these cases represent a subcategory of HIV encephalopathy. Of these 23 cases, 16 (70%) had 2 or more seizures, suggesting that anticonvulsant therapy is indicated in all patients with HIV infection and seizures unless a rapidly correctable cause is found. While phenytoin remains the initial treatment of choice, hypersensitivity reactions to this drug have been reported in >10% of patients with AIDS, and therefore the use of phenobarbital or valproic acid should be considered as alternatives. Due to a variety of drug-drug interactions between antiseizure medications and antiretrovirals, drug levels need to be monitored carefully.

Patients with HIV infection may present with *focal neurologic deficits* from a variety of causes. The most common causes are toxoplasmosis, progressive multifocal leukoencephalopathy, and CNS lymphoma. Other causes include cryptococcal infections (discussed above; also Chap. 239), stroke, and reactivation of Chagas' disease.

Toxoplasmosis has been one of the most common causes of secondary CNS infections in patients with AIDS, but its incidence is decreasing in the era of cART. It is most common in patients from the Caribbean and from France, where the seroprevalence of *T. gondii* is around 50%. This figure is closer to 15% in the United States. Toxoplasmosis is generally a late complication of HIV infection and usually occurs in patients with CD4+ T cell counts <200/ μ L. Cerebral toxoplasmosis is thought to represent a reactivation of latent tissue cysts. It is 10 times more common in patients with antibodies to the organism than in patients who are seronegative. Patients diagnosed with HIV infection should be screened for IgG antibodies to *T. gondii* during the time of their initial workup. Those who are seronegative should be counseled about ways to minimize the risk of primary infection including avoiding the consumption of undercooked meat and careful hand washing after contact with soil or changing the cat litter box. The most common clinical presentation of cerebral toxoplasmosis in patients with HIV infection is fever, headache, and focal neurologic deficits. Patients may present with seizure, hemiparesis, or aphasia as a manifestation of these focal deficits or with a picture more influenced by the accompanying cerebral edema and characterized by confusion, dementia, and lethargy, which can progress to coma. The diagnosis is usually suspected on the basis of MRI findings of multiple lesions in multiple locations, although in some cases only a single lesion is seen. Pathologically, these lesions generally exhibit inflammation and central necrosis and, as a result, demonstrate ring enhancement on contrast MRI (Fig. 226-40) or, if MRI is unavailable or contraindicated, on double-dose contrast CT. There is usually evidence of surrounding edema. In addition to toxoplasmosis, the differential diagnosis of single or multiple enhancing mass lesions in

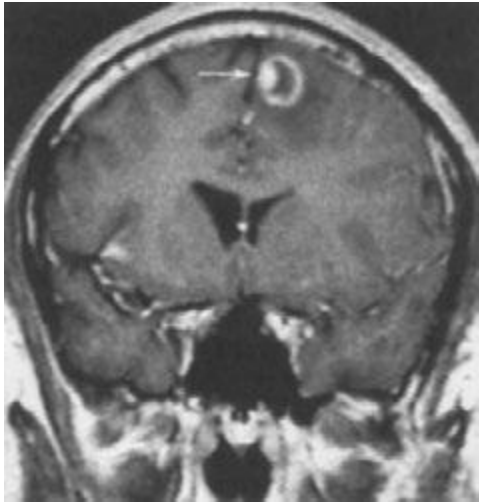


FIGURE 226-40 Central nervous system toxoplasmosis. A coronal postcontrast T1-weighted MRI scan demonstrates a peripheral enhancing lesion in the left frontal lobe, associated with an eccentric nodular area of enhancement (arrow); this so-called eccentric target sign is typical of toxoplasmosis.

the HIV-infected patient includes primary CNS lymphoma and, less commonly, TB or fungal or bacterial abscesses. The definitive diagnostic procedure is brain biopsy. However, given the morbidity rate that can accompany this procedure, it is usually reserved for the patient who has failed 2–4 weeks of empiric therapy for toxoplasmosis. If the patient is seronegative for *T. gondii*, the likelihood that a mass lesion is due to toxoplasmosis is <10%. In that setting, one may choose to be more aggressive and perform a brain biopsy sooner. Standard treatment is sulfadiazine and pyrimethamine with leucovorin as needed for a minimum of 4–6 weeks. Alternative therapeutic regimens include clindamycin in combination with pyrimethamine; atovaquone plus pyrimethamine; and azithromycin plus pyrimethamine plus rifabutin. Relapses are common, and it is recommended that patients with a history of prior toxoplasmic encephalitis receive maintenance therapy with sulfadiazine, pyrimethamine, and leucovorin as long as their CD4+ T cell counts remain <200 cells/ μ L. Patients with CD4+ T cell counts <100/ μ L and IgG antibody to *Toxoplasma* should receive primary prophylaxis for toxoplasmosis. Fortunately, the same daily regimen of a single double-strength tablet of TMP/SMX used for *P. jiroveci* prophylaxis provides adequate primary protection against toxoplasmosis. Secondary prophylaxis/maintenance therapy for toxoplasmosis may be discontinued in the setting of effective cART and increases in CD4+ T cell counts to >200/ μ L for 6 months.

JC virus, a human polyomavirus that is the etiologic agent of *progressive multifocal leukoencephalopathy* (PML), is an important opportunistic pathogen in patients with AIDS (Chap. 164). While ~80% of the general adult population has antibodies to JC virus, indicative of prior infection, <10% of healthy adults show any evidence of ongoing viral replication. PML is the only known clinical manifestation of JC virus infection. It is a late manifestation of AIDS and is seen in ~1–4% of patients with AIDS. The lesions of PML begin as small foci of demyelination in subcortical white matter that eventually coalesce. The cerebral hemispheres, cerebellum, and brainstem may all be involved. Patients typically have a protracted course with multifocal neurologic deficits, with or without changes in mental status. Approximately 20% of patients experience seizures. Ataxia, hemiparesis, visual field defects, aphasia, and sensory defects may occur. Headache, fever, nausea, and vomiting are rarely seen. Their presence should suggest another diagnosis. MRI typically reveals multiple, nonenhancing white matter lesions that may coalesce and have a predilection for the occipital and parietal lobes. The lesions show signal hyperintensity on T2-weighted images and diminished signal on T1-weighted images. The measurement of JC virus DNA levels in CSF has a diagnostic sensitivity of 76% and a specificity of close to 100%. Prior to the availability of cART,

the majority of patients with PML died within 3–6 months of the onset of symptoms. Paradoxical worsening of PML has been seen with initiation of cART as an immune reconstitution syndrome. There is no specific treatment for PML; however, a median survival of 2 years and survival of >15 years have been reported in patients with PML treated with cART for their HIV disease. Despite having a significant impact on survival, only ~50% of patients with HIV infection and PML show neurologic improvement with cART. Studies with other antiviral agents such as cidofovir have failed to show clear benefit. Factors influencing a favorable prognosis for PML in the setting of HIV infection include a CD4+ T cell count >100/ μ L at baseline and the ability to maintain an HIV viral load of <500 copies/mL. Baseline HIV-1 viral load does not have independent predictive value of survival. PML is one of the few opportunistic infections that continues to occur with some frequency despite the widespread use of cART.

Reactivation American trypanosomiasis may present as acute meningoencephalitis with focal neurologic signs, fever, headache, vomiting, and seizures. Accompanying cardiac disease in the form of arrhythmias or heart failure should increase the index of suspicion. The presence of antibodies to *T. cruzi* supports the diagnosis. In South America, reactivation of *Chagas' disease* is considered to be an AIDS-defining condition and may be the initial AIDS-defining condition. The majority of cases occur in patients with CD4+ T cell counts <200 cells/ μ L. Lesions appear radiographically as single or multiple hypodense areas, typically with ring enhancement and edema. They are found predominantly in the subcortical areas, a feature that differentiates them from the deeper lesions of toxoplasmosis. *T. cruzi* amastigotes, or trypanosomes, can be identified from biopsy specimens or CSF. Other CSF findings include elevated protein and a mild (<100 cells/ μ L) lymphocytic pleocytosis. Organisms can also be identified by direct examination of the blood. Treatment consists of benzimidazole (2.5 mg/kg bid) or nifurtimox (2 mg/kg qid) for at least 60 days, followed by maintenance therapy for the duration of immunodeficiency with either drug at a dose of 5 mg/kg three times a week. As is the case with cerebral toxoplasmosis, successful therapy with antiretrovirals may allow discontinuation of therapy for Chagas' disease.

Stroke may occur in patients with HIV infection. In contrast to the other causes of focal neurologic deficits in patients with HIV infection, the symptoms of a stroke are sudden in onset. Patients with HIV infection have an increased prevalence of many classic risk factors associated with stroke, including smoking and diabetes. It has been reported that HIV infection itself can lead to an increase in carotid artery stiffness. The relative increase in risk for stroke as a consequence of HIV infection is more pronounced in women and in individuals between the ages of 18 and 29. Among the secondary infectious diseases in patients with HIV infection that may be associated with stroke are vasculitis due to cerebral varicella zoster or neurosyphilis and septic embolism in association with fungal infection. Other elements of the differential diagnosis of stroke in the patient with HIV infection include atherosclerotic cerebral vascular disease, thrombotic thrombocytopenic purpura, and cocaine or amphetamine use.

Primary CNS lymphoma is discussed below in the section on neoplastic diseases.

Spinal cord disease, or myelopathy, is present in ~20% of patients with AIDS, often as part of HIV-associated neurocognitive disorder. In fact, 90% of the patients with HIV-associated myelopathy have some evidence of dementia, suggesting that similar pathologic processes may be responsible for both conditions. Three main types of spinal cord disease are seen in patients with AIDS. The first of these is a vacuolar myelopathy, as mentioned above. This condition is pathologically similar to subacute combined degeneration of the cord, such as that occurring with pernicious anemia. Although vitamin B₁₂ deficiency can be seen in patients with AIDS as a primary complication of HIV infection, it does not appear to be responsible for the myelopathy seen in the majority of patients. Vacuolar myelopathy is characterized by a subacute onset and often presents with gait disturbances, predominantly ataxia and spasticity; it may progress to include bladder and bowel dysfunction. Physical findings include evidence of increased deep tendon reflexes and extensor plantar responses. The second

form of spinal cord disease involves the dorsal columns and presents as a pure sensory ataxia. The third form is also sensory in nature and presents with paresthesias and dysesthesias of the lower extremities. In contrast to the cognitive problems seen in patients with HIV encephalopathy, these spinal cord syndromes do not respond well to antiretroviral drugs, and therapy is mainly supportive.

One important disease of the spinal cord that also involves the peripheral nerves is a *myelopathy* and *polyradiculopathy* seen in association with CMV infection. This entity is generally seen late in the course of HIV infection and is fulminant in onset, with lower extremity and sacral paresthesias, difficulty in walking, areflexia, ascending sensory loss, and urinary retention. The clinical course is rapidly progressive over a period of weeks. CSF examination reveals a predominantly neutrophilic pleocytosis, and CMV DNA can be detected by CSF PCR. Therapy with ganciclovir or foscarnet can lead to rapid improvement, and prompt initiation of foscarnet or ganciclovir therapy is important in minimizing the degree of permanent neurologic damage. Combination therapy with both drugs should be considered in patients who have been previously treated for CMV disease. Other diseases involving the spinal cord in patients with HIV infection include HTLV-1-associated myelopathy (HAM) (Chap. 225e), neurosyphilis (Chap. 206), infection with herpes simplex (Chap. 216) or varicella-zoster (Chap. 217), TB (Chap. 202), and lymphoma (Chap. 134).

Peripheral neuropathies are common in patients with HIV infection. They occur at all stages of illness and take a variety of forms. Early in the course of HIV infection, an acute inflammatory demyelinating polyneuropathy resembling Guillain-Barré syndrome may occur (Chap. 460). In other patients, a progressive or relapsing-remitting inflammatory neuropathy resembling chronic inflammatory demyelinating polyneuropathy (CIPD) has been noted. Patients commonly present with progressive weakness, areflexia, and minimal sensory changes. CSF examination often reveals a mononuclear pleocytosis, and peripheral nerve biopsy demonstrates a perivascular infiltrate suggesting an autoimmune etiology. Plasma exchange or IVIg has been tried with variable success. Because of the immunosuppressive effects of glucocorticoids, they should be reserved for severe cases of CIPD refractory to other measures. Another autoimmune peripheral neuropathy seen in patients with AIDS is mononeuritis multiplex (Chaps. 460 and 385) due to a necrotizing arteritis of peripheral nerves. The most common peripheral neuropathy in patients with HIV infection is a *distal sensory polyneuropathy* (DSPN) also referred to as painful sensory neuropathy (HIV-SN), predominantly sensory neuropathy, or distal symmetric peripheral neuropathy. This condition may be a direct consequence of HIV infection or a side effect of dideoxynucleoside therapy. It is more common in taller individuals, older individuals, and those with lower CD4 counts. Two-thirds of patients with AIDS may be shown by electrophysiologic studies to have some evidence of peripheral nerve disease. Presenting symptoms are usually painful burning sensations in the feet and lower extremities. Findings on examination include a stocking-type sensory loss to pinprick, temperature, and touch sensation and a loss of ankle reflexes. Motor changes are mild and are usually limited to weakness of the intrinsic foot muscles. Response of this condition to antiretrovirals has been variable, perhaps because antiretrovirals are responsible for the problem in some instances. When due to dideoxynucleoside therapy, patients with lower extremity peripheral neuropathy may complain of a sensation that they are walking on ice. Other entities in the differential diagnosis of peripheral neuropathy include diabetes mellitus, vitamin B₁₂ deficiency, and side effects from metronidazole or dapsone. For distal symmetric polyneuropathy that fails to resolve following the discontinuation of dideoxynucleosides, therapy is symptomatic; gabapentin, carbamazepine, tricyclics, or analgesics may be effective for dysesthesias. Treatment-naïve patients may respond to cART.

Myopathy may complicate the course of HIV infection; causes include HIV infection itself, zidovudine, and the generalized wasting syndrome. HIV-associated myopathy may range in severity from an asymptomatic elevation in creatine kinase levels to a subacute syndrome characterized by proximal muscle weakness and myalgias. Quite pronounced elevations in creatine kinase may occur in asymptomatic

patients, particularly after exercise. The clinical significance of this as an isolated laboratory finding is unclear. A variety of both inflammatory and noninflammatory pathologic processes have been noted in patients with more severe myopathy, including myofiber necrosis with inflammatory cells, nemaline rod bodies, cytoplasmic bodies, and mitochondrial abnormalities. Profound muscle wasting, often with muscle pain, may be seen after prolonged zidovudine therapy. This toxic side effect of the drug is dose-dependent and is related to its ability to interfere with the function of mitochondrial polymerases. It is reversible following discontinuation of the drug. Red ragged fibers are a histologic hallmark of zidovudine-induced myopathy.

Ophthalmologic Diseases Ophthalmologic problems occur in ~50% of patients with advanced HIV infection. The most common abnormal findings on funduscopic examination are cotton-wool spots. These are hard white spots that appear on the surface of the retina and often have an irregular edge. They represent areas of retinal ischemia secondary to microvascular disease. At times they are associated with small areas of hemorrhage and thus can be difficult to distinguish from CMV retinitis. In contrast to CMV retinitis, however, these lesions are not associated with visual loss and tend to remain stable or improve over time.

One of the most devastating consequences of HIV infection is CMV retinitis. Patients at high risk of CMV retinitis (CD4+ T cell count <100/μL) should undergo an ophthalmologic examination every 3–6 months. The majority of cases of CMV retinitis occur in patients with a CD4+ T cell count <50/μL. Prior to the availability of cART, this CMV reactivation syndrome was seen in 25–30% of patients with AIDS. In the cART era this has dropped to close to 2%. CMV retinitis usually presents as a painless, progressive loss of vision. Patients may also complain of blurred vision, “floaters,” and scintillations. The disease is usually bilateral, although typically it affects one eye more than the other. The diagnosis is made on clinical grounds by an experienced ophthalmologist. The characteristic retinal appearance is that of perivascular hemorrhage and exudate. In situations where the diagnosis is in doubt due to an atypical presentation or an unexpected lack of response to therapy, vitreous or aqueous humor sampling with molecular diagnostic techniques may be of value. CMV infection of the retina results in a necrotic inflammatory process, and the visual loss that develops is irreversible. CMV retinitis may be complicated by rhegmatogenous retinal detachment as a consequence of retinal atrophy in areas of prior inflammation. Therapy for CMV retinitis consists of oral valganciclovir, IV ganciclovir, or IV foscarnet, with cidofovir as an alternative. Combination therapy with ganciclovir and foscarnet has been shown to be slightly more effective than either ganciclovir or foscarnet alone in the patient with relapsed CMV retinitis. A 3-week induction course is followed by maintenance therapy with oral valganciclovir. If CMV disease is limited to the eye, intravitreal injections of ganciclovir or foscarnet may be considered. Intravitreal injections of cidofovir are generally avoided due to the increased risk of uveitis and hypotony. Maintenance therapy is continued until the CD4+ T cell count remains >100 μL for >6 months. The majority of patients with HIV infection and CMV disease develop some degree of uveitis with the initiation of cART. The etiology of this is unknown; however, it has been suggested that this may be due to the generation of an enhanced immune response to CMV as an IRIS (see above). In some instances this has required the use of topical glucocorticoids.

Both HSV and varicella zoster virus can cause a rapidly progressing, bilateral necrotizing retinitis referred to as the *acute retinal necrosis syndrome*, or *progressive outer retinal necrosis* (PORN). This syndrome, in contrast to CMV retinitis, is associated with pain, keratitis, and iritis. It is often associated with orolabial HSV or trigeminal zoster. Ophthalmologic examination reveals widespread pale gray peripheral lesions. This condition is often complicated by retinal detachment. It is important to recognize and treat this condition with IV acyclovir as quickly as possible to minimize the loss of vision.

Several other secondary infections may cause ocular problems in HIV-infected patients. *P. jiroveci* can cause a lesion of the choroid that may be detected as an incidental finding on ophthalmologic examination. These lesions are typically bilateral, are from half to twice the disc

1268 diameter in size, and appear as slightly elevated yellow-white plaques. They are usually asymptomatic and may be confused with cotton-wool spots. Chorioretinitis due to toxoplasmosis can be seen alone or, more commonly, in association with CNS toxoplasmosis. KS may involve the eyelid or conjunctiva, while lymphoma may involve the retina. Syphilis may lead to a uveitis that is highly associated with the presence of neurosyphilis.

Additional Disseminated Infections and Wasting Syndrome Infections with species of the small, gram-negative, *Rickettsia*-like organism *Bartonella* (Chap. 197) are seen with increased frequency in patients with HIV infection. While it is not considered an AIDS-defining illness by the CDC, many experts view infection with *Bartonella* as indicative of a severe defect in cell-mediated immunity. It is usually seen in patients with CD4+ T cell counts <100/μL and is a significant cause of unexplained fever in patients with advanced HIV infection. Among the clinical manifestations of *Bartonella* infection are bacillary angiomatosis, cat-scratch disease, and trench fever. *Bacillary angiomatosis* is usually due to infection with *B. henselae* and is linked to exposure to flea-infested cats. It is characterized by a vascular proliferation that leads to a variety of skin lesions that have been confused with the skin lesions of KS. In contrast to the lesions of KS, the lesions of bacillary angiomatosis generally blanch, are painful, and typically occur in the setting of systemic symptoms. Infection can extend to the lymph nodes, liver (peliosis hepatis), spleen, bone, heart, CNS, respiratory tract, and GI tract. *Cat-scratch disease* also is due to *B. henselae* and generally begins with a papule at the site of inoculation. This is followed several weeks later by the development of regional adenopathy and malaise. Infection with *B. quintana* is transmitted by lice and has been associated with case reports of trench fever, endocarditis, adenopathy, and bacillary angiomatosis. The organism is quite difficult to culture, and diagnosis often relies on identifying the organism in biopsy specimens using the Warthin-Starry or similar stains. Treatment is with either doxycycline or erythromycin for at least 3 months.

Histoplasmosis is an opportunistic infection that is seen most frequently in patients in the Mississippi and Ohio River valleys, Puerto Rico, the Dominican Republic, and South America. These are all areas in which infection with *H. capsulatum* is endemic (Chap. 236). Because of this limited geographic distribution, the percentage of AIDS cases in the United States with histoplasmosis is only ~0.5. Histoplasmosis is generally a late manifestation of HIV infection; however, it may be the initial AIDS-defining condition. In one study, the median CD4+ T cell count for patients with histoplasmosis and AIDS was 33/μL. While disease due to *H. capsulatum* may present as a primary infection of the lung, disseminated disease, presumably due to reactivation, is the most common presentation in HIV-infected patients. Patients usually present with a 4- to 8-week history of fever and weight loss. Hepatosplenomegaly and lymphadenopathy are each seen in about 25% of patients. CNS disease, either meningitis or a mass lesion, is seen in 15% of patients. Bone marrow involvement is common, with thrombocytopenia, neutropenia, and anemia occurring in 33% of patients. Approximately 7% of patients have mucocutaneous lesions consisting of a maculopapular rash and skin or oral ulcers. Respiratory symptoms are usually mild, with chest x-ray showing a diffuse infiltrate or diffuse small nodules in ~50% of cases. The gastrointestinal tract may be involved. Diagnosis is made by silver staining of tissue, by culturing the organisms from blood, bone marrow, or tissue, or by detecting antigen in blood or urine. Treatment is typically with liposomal amphotericin B followed by maintenance therapy with oral itraconazole until the serum histoplasma antigen is <2 units, the patient has been on antiretrovirals for at least 6 months, and the CD4 count is >150 cells/μL. In the setting of mild infection, it may be appropriate to initiate therapy with itraconazole alone.

Following the spread of HIV infection to southeast Asia, disseminated infection with the fungus *Penicillium marneffeii* was recognized as a complication of HIV infection and is considered an AIDS-defining condition in those parts of the world where it occurs. *P. marneffeii* is the third most common AIDS-defining illness in Thailand, following TB and cryptococcosis. It is more frequently diagnosed in the rainy than the dry season. Clinical features include fever, generalized

lymphadenopathy, hepatosplenomegaly, anemia, thrombocytopenia, and papular skin lesions with central umbilication. Treatment is with amphotericin B followed by itraconazole until the CD4+ T cell count is >100 cells/μL for at least 6 months.

Visceral leishmaniasis (Chap. 251) is recognized with increasing frequency in patients with HIV infection who live in or travel to areas endemic for this protozoal infection transmitted by sandflies. The clinical presentation is one of hepatosplenomegaly, fever, and hematologic abnormalities. Lymphadenopathy and other constitutional symptoms may be present. A chronic, relapsing course is seen in two-thirds of co-infected patients. Organisms can be isolated from cultures of bone marrow aspirates. Histologic stains may be negative, and antibody titers are of little help. Patients with HIV infection usually respond well initially to standard therapy with amphotericin B or pentavalent antimony compounds. Eradication of the organism is difficult, however, and relapses are common.

Patients with HIV infection are at a slightly increased risk of clinical malaria. This is particularly true for patients from nonendemic areas with presumed primary infection and in patients with lower CD4+ T cell counts. HIV-positive individuals with CD4+ T cell counts <300 cells/μL have a poorer response to malaria treatment than others. Co-infection with malaria is associated with a modest increase in HIV viral load. The risk of malaria may be decreased with TMP/SMX prophylaxis.

Generalized wasting is an AIDS-defining condition; it is defined as involuntary weight loss of >10% associated with intermittent or constant fever and chronic diarrhea or fatigue lasting >30 days in the absence of a defined cause other than HIV infection. Prior to the widespread use of cART it was the initial AIDS-defining condition in ~10% of patients with AIDS in the United States and is an indication for initiation of cART. Generalized wasting is rarely seen today with the earlier initiation of antiretrovirals. A constant feature of this syndrome is severe muscle wasting with scattered myofiber degeneration and occasional evidence of myositis. Glucocorticoids may be of some benefit; however, this approach must be carefully weighed against the risk of compounding the immunodeficiency of HIV infection. Androgenic steroids, growth hormone, and total parenteral nutrition have been used as therapeutic interventions with variable success.

Neoplastic Diseases The neoplastic diseases considered to be AIDS-defining conditions are Kaposi's sarcoma, non-Hodgkin's lymphoma, and invasive cervical carcinoma. In addition, there is also an increase in the incidence of a variety of non-AIDS-defining malignancies including Hodgkin's disease; multiple myeloma; leukemia; melanoma; and cervical, brain, testicular, oral, lung, gastric, liver, renal, and anal cancers. Since the introduction of potent cART, there has been a marked reduction in the incidence of KS (Fig. 226-33) and CNS lymphoma, such that the non-AIDS-defining malignancies now account for more morbidity and mortality in patients with HIV infection than the AIDS-defining malignancies. Rates of non-Hodgkin's lymphoma have declined as well; however, this decline has not been as dramatic as the decline in rates of KS. In contrast, cART has had little effect on human papillomavirus (HPV)-associated malignancies. As patients with HIV infection live longer, a wider array of cancers is seen in this population. While some may only reflect known risk factors (e.g., smoking, alcohol consumption, co-infection with other viruses such as hepatitis B) that are increased in patients with HIV infection, some may be a direct consequence of HIV and are clearly increased in patients with lower CD4+ T cell counts.

Kaposi's sarcoma is a multicentric neoplasm consisting of multiple vascular nodules appearing in the skin, mucous membranes, and viscera. The course ranges from indolent, with only minor skin or lymph node involvement, to fulminant, with extensive cutaneous and visceral involvement. In the initial period of the AIDS epidemic, KS was a prominent clinical feature of the first cases of AIDS, occurring in 79% of the patients diagnosed in 1981. By 1989 it was seen in only 25% of cases, by 1992 the number had decreased to 9%, and by 1997 the number was <1%. HHV-8 (KSHV) has been strongly implicated as a viral cofactor in the pathogenesis of KS.

Clinically, KS has varied presentations and may be seen at any stage of HIV infection, even in the presence of a normal CD4+ T cell



FIGURE 226-41 Kaposi's sarcoma in three patients with AIDS demonstrating (A) periorbital edema and bruising; (B) classic truncal distribution of lesions; and (C) upper extremity lesions.

count. The initial lesion may be a small, raised reddish-purple nodule on the skin (Fig. 226-41), a discoloration on the oral mucosa (Fig. 226-34D), or a swollen lymph node. Lesions often appear in sun-exposed areas, particularly the tip of the nose, and have a propensity to occur in areas of trauma (Koebner phenomenon). Because of the vascular nature of the tumors and the presence of extravasated red blood cells in the lesions, their colors range from reddish to purple to brown and often take the appearance of a bruise, with yellowish discoloration and tattooing. Lesions range in size from a few millimeters to several centimeters in diameter and may be either discrete or confluent. KS lesions most commonly appear as raised macules; however, they can also be papular, particularly in patients with higher CD4+ T cell counts. Confluent lesions may give rise to surrounding lymphedema and may be disfiguring when they involve the face and disabling when they involve the lower extremities or the surfaces of joints. Apart from skin, the lymph nodes, GI tract, and lung are the organ systems most commonly affected by KS. Lesions have been reported in virtually every organ, including the heart and the CNS. In contrast to most malignancies, in which lymph node involvement implies metastatic spread and a poor prognosis, lymph node involvement may be seen very early in KS and is of no special clinical significance. In fact, some patients may present with disease limited to the lymph nodes. These are generally patients with relatively intact immune function and thus the patients with the best prognosis. Pulmonary involvement with KS generally presents with shortness of breath. Some 80% of patients with pulmonary KS also have cutaneous lesions. The chest x-ray characteristically shows bilateral lower lobe infiltrates that obscure the margins of the mediastinum and diaphragm (Fig. 226-42). Pleural effusions



FIGURE 226-42 Chest x-ray of a patient with AIDS and pulmonary Kaposi's sarcoma. The characteristic findings include dense bilateral lower lobe infiltrates obscuring the heart borders and pleural effusions.

are seen in 70% of cases of pulmonary KS, a fact that is often helpful in the differential diagnosis. GI involvement is seen in 50% of patients with KS and usually takes one of two forms: (1) mucosal involvement, which may lead to bleeding that can be severe; these patients sometimes also develop symptoms of GI obstruction if lesions become large; and (2) biliary tract involvement. KS lesions may infiltrate the gallbladder and biliary tree, leading to a clinical picture of obstructive jaundice similar to that seen with sclerosing cholangitis. Several staging systems have been proposed for KS. One in common use was developed by the National Institute of Allergy and Infectious Diseases AIDS Clinical Trials Group; it distinguishes patients on the basis of tumor extent, immunologic function, and presence or absence of systemic disease (Table 226-17).

A diagnosis of KS is based on biopsy of a suspicious lesion. Histologically one sees a proliferation of spindle cells and endothelial cells, extravasation of red blood cells, hemosiderin-laden macrophages, and, in early cases, an inflammatory cell infiltrate. Included in the differential diagnosis are lymphoma (particularly for oral lesions), bacillary angiomatosis, and cutaneous mycobacterial infections.

Management of KS (Table 226-18) should be carried out in consultation with an expert since definitive treatment guidelines do not exist. In the majority of cases, effective cART will go a long way in achieving control. Antiretroviral therapy has been associated with the spontaneous regression of KS lesions. Paradoxically, it has also been associated with the initial appearance of KS as a form of IRIS. For patients in whom tumor persists or is compromising vital functions or in whom control of HIV replication is not possible, a variety of options exist. In some cases, lesions remain quite indolent, and many of these patients can be managed with no specific treatment. Fewer than 10% of AIDS patients with KS die as a consequence of their malignancy,

TABLE 226-17 NATIONAL INSTITUTE OF ALLERGY AND INFECTIOUS DISEASES AIDS CLINICAL TRIALS GROUP TIS STAGING SYSTEM FOR KAPOSI'S SARCOMA

Parameter	Good Risk (Stage 0): All of the Following	Poor Risk (Stage 1): Any of the Following
Tumor (T)	Confined to skin and/or lymph nodes and/or minimal oral disease	Tumor-associated edema or ulceration Extensive oral lesions GI lesions Nonnodal visceral lesions
Immune system (I)	CD4+ T cell count $\geq 200/\mu\text{L}$	CD4+ T cell count $< 200/\mu\text{L}$
Systemic illness (S)	No B symptoms ^a Karnofsky performance status ≥ 70 No history of opportunistic infection, neurologic disease, lymphoma, or thrush	B symptoms ^a present Karnofsky performance status < 70 History of opportunistic infection, neurologic disease, lymphoma, or thrush

^aDefined as unexplained fever, night sweats, $>10\%$ involuntary weight loss, or diarrhea persisting for more than 2 weeks.

TABLE 226-18 MANAGEMENT OF AIDS-ASSOCIATED KAPOSI'S SARCOMA

Observation and optimization of antiretroviral therapy
Single or limited number of lesions
Radiation
Intralesional vinblastine
Cryotherapy
Extensive disease
Initial therapy
Interferon α (if CD4+ T cells >150/ μ L)
Liposomal daunorubicin
Subsequent therapy
Liposomal doxorubicin
Paclitaxel
Combination chemotherapy with low-dose doxorubicin, bleomycin, and vinblastine (ABV)
Targeted radiation

and death from secondary infections is considerably more common. Thus, whenever possible one should avoid treatment regimens that may further suppress the immune system and increase susceptibility to opportunistic infections. Treatment is indicated under two main circumstances. The first is when a single lesion or a limited number of lesions are causing significant discomfort or cosmetic problems, such as with prominent facial lesions, lesions overlying a joint, or lesions in the oropharynx that interfere with swallowing or breathing. Under these circumstances, treatment with localized radiation, intralesional vinblastine, topical 9-*cis*-retinoic acid, or cryotherapy may be helpful. It should be noted that patients with HIV infection are particularly sensitive to the side effects of radiation therapy. This is especially true with respect to the development of radiation-induced mucositis; doses of radiation directed at mucosal surfaces, particularly in the head and neck region, should be adjusted accordingly. The use of systemic therapy, either IFN- α or chemotherapy, should be considered in patients with a large number of lesions or in patients with visceral involvement. The single most important determinant of response appears to be the CD4+ T cell count. This relationship between response rate and baseline CD4+ T cell count is particularly true for IFN- α . The response rate to IFN- α for patients with CD4+ T cell counts >600/ μ L is ~80%, while the response rate for patients with counts <150/ μ L is <10%. In contrast to the other systemic therapies, IFN- α provides an added advantage of having antiretroviral activity; thus, it may be the appropriate first choice for single-agent systemic therapy for early patients with disseminated disease. A variety of chemotherapeutic agents also have been shown to have activity against KS. Four of them—liposomal daunorubicin, liposomal doxorubicin, vinblastine, and paclitaxel—have been approved by the FDA for this indication. Liposomal daunorubicin is approved as first-line therapy for patients with advanced KS. It has fewer side effects than conventional chemotherapy. In contrast, liposomal doxorubicin and paclitaxel are approved only for KS patients who have failed standard chemotherapy. Response rates vary from 23 to 88%, appear to be comparable to what had been achieved earlier with combination chemotherapy regimens, and are greatly influenced by CD4+ T cell count.

Lymphomas occur with an increased frequency in patients with congenital or acquired T cell immunodeficiencies (Chap. 374). AIDS is no exception; at least 6% of all patients with AIDS develop lymphoma at some time during the course of their illness. This is a 120-fold increase in incidence compared with the general population. In contrast to the situation with KS, primary CNS lymphoma, and most opportunistic infections, the incidence of AIDS-associated systemic lymphomas has not experienced a dramatic decrease as a consequence of the widespread use of effective cART. Lymphoma occurs in all risk groups, with the highest incidence in patients with hemophilia and the lowest incidence in patients from the Caribbean or Africa with heterosexually acquired infection. Lymphoma is a late manifestation of HIV infection, generally occurring in patients with CD4+ T cell counts <200/ μ L. As

HIV disease progresses, the risk of lymphoma increases. The attack rate for lymphoma increases exponentially with increasing duration of HIV infection and decreasing level of immunologic function. At 3 years following a diagnosis of HIV infection, the risk of lymphoma is 0.8% per year; by 8 years after infection, it is 2.6% per year. As individuals with HIV infection live longer as a consequence of improved cART and better treatment and prophylaxis of opportunistic infections, it is anticipated that the incidence of lymphomas may increase.

Three main categories of lymphoma are seen in patients with HIV infection: grade III or IV immunoblastic lymphoma, Burkitt's lymphoma, and primary CNS lymphoma. Approximately 90% of these lymphomas are B cell in phenotype; more than half contain EBV DNA. Some are associated with KSHV. These tumors may be either monoclonal or oligoclonal in nature and are probably in some way related to the pronounced polyclonal B cell activation seen in patients with AIDS.

Immunoblastic lymphomas account for ~60% of the cases of lymphoma in patients with AIDS. The majority of these are diffuse large B cell lymphomas (DLBCL). They are generally high grade and would have been classified as diffuse histiocytic lymphomas in earlier classification schemes. This tumor is more common in older patients, increasing in incidence from 0% in HIV-infected individuals <1 year old to >3% in those >50 years of age. Two variants of immunoblastic lymphoma that are seen primarily in HIV-infected patients are primary effusion lymphoma (PEL) and its solid variant, plasmacytic lymphoma of the oral cavity. PEL, also referred to as body cavity lymphoma, presents with lymphomatous pleural, pericardial, and/or peritoneal effusions in the absence of discrete nodal or extranodal masses. The tumor cells do not express surface markers for B cells or T cells and are felt to represent a preplasmacytic stage of differentiation. While both HHV-8 and EBV DNA sequences have been found in the genomes of the malignant cells from patients with body cavity lymphoma, KSHV is felt to be the driving force behind the oncogenesis (see above).

Small noncleaved cell lymphoma (Burkitt's lymphoma) accounts for ~20% of the cases of lymphoma in patients with AIDS. It is most frequent in patients 10–19 years old and usually demonstrates characteristic *c-myc* translocations from chromosome 8 to chromosomes 14 or 22. Burkitt's lymphoma is not commonly seen in the setting of immunodeficiency other than HIV-associated immunodeficiency, and the incidence of this particular tumor is more than 1000-fold higher in the setting of HIV infection than in the general population. In contrast to African Burkitt's lymphoma, where 97% of the cases contain EBV genome, only 50% of HIV-associated Burkitt's lymphomas are EBV-positive.

Primary CNS lymphoma accounts for ~20% of the cases of lymphoma in patients with HIV infection. In contrast to HIV-associated Burkitt's lymphoma, primary CNS lymphomas are usually positive for EBV. In one study, the incidence of Epstein-Barr positivity was 100%. This malignancy does not have a predilection for any particular age group. The median CD4+ T cell count at the time of diagnosis is ~50/ μ L. Thus, CNS lymphoma generally presents at a later stage of HIV infection than does systemic lymphoma. This may explain, at least in part, the poorer prognosis for this subset of patients.

The clinical presentation of lymphoma in patients with HIV infection is quite varied, ranging from focal seizures to rapidly growing mass lesions in the oral mucosa (Fig. 226-43) to persistent unexplained fever. At least 80% of patients present with extranodal disease, and a similar percentage have B-type symptoms of fever, night sweats, or weight loss. Virtually any site in the body may be involved. The most common extranodal site is the CNS, which is involved in approximately one-third of all patients with lymphoma. Approximately 60% of these cases are primary CNS lymphoma. Primary CNS lymphoma generally presents with focal neurologic deficits, including cranial nerve findings, headaches, and/or seizures. MRI or CT generally reveals a limited number (one to three) of 3- to 5-cm lesions (Fig. 226-44). The lesions often show ring enhancement on contrast administration and may occur in any location. Contrast enhancement is usually less pronounced than that seen with toxoplasmosis.



FIGURE 226-43 Immunoblastic lymphoma involving the hard palate of a patient with AIDS.

Locations that are most commonly involved with CNS lymphoma are deep in the white matter. The main diseases in the differential diagnosis are cerebral toxoplasmosis and cerebral Chagas' disease. In addition to the 20% of lymphomas in HIV-infected individuals that are primary CNS lymphomas, CNS disease is also seen in HIV-infected patients with systemic lymphoma. Approximately 20% of patients with systemic lymphoma have CNS disease in the form of leptomeningeal involvement. This fact underscores the importance of lumbar puncture in the staging evaluation of patients with systemic lymphoma.

Systemic lymphoma is seen at earlier stages of HIV infection than primary CNS lymphoma. In one series the mean CD4+ T cell count was 226/ μ L. In addition to lymph node involvement, systemic lymphoma may commonly involve the GI tract, bone marrow, liver, and lung. GI tract involvement is seen in ~25% of patients. Any site in the GI tract may be involved, and patients may complain of difficulty swallowing or abdominal pain. The diagnosis is usually suspected on the basis of CT or MRI of the abdomen. Bone marrow involvement is seen in ~20% of patients and may lead to pancytopenia. Liver and lung involvement are each seen in ~10% of patients. Pulmonary disease may present as a mass lesion, multiple nodules, or an interstitial infiltrate.

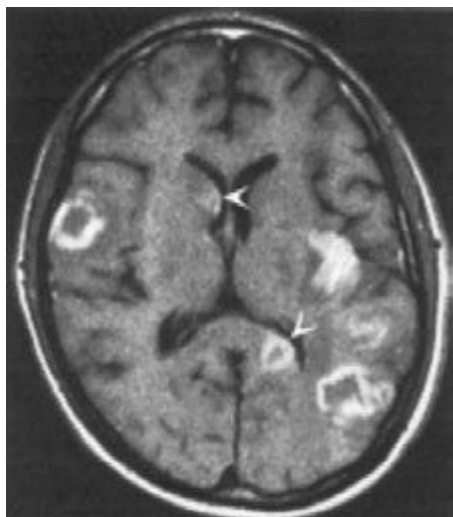


FIGURE 226-44 Central nervous system lymphoma. Postcontrast T1-weighted MRI scan in a patient with AIDS, an altered mental status, and hemiparesis. Multiple enhancing lesions, some ring-enhancing, are present. The left sylvian lesion shows gyral and subcortical enhancement, and the lesions in the caudate and splenium (arrowheads) show enhancement of adjacent ependymal surfaces.

Both conventional and unconventional approaches have been employed in an attempt to treat HIV-related lymphomas. Systemic lymphoma is generally treated by the oncologist with combination chemotherapy. Earlier disappointing figures are being replaced with more optimistic results for the treatment of systemic lymphoma following the availability of more effective cART and the use of rituximab in CD20+ tumors. While there is some controversy regarding the use of antiretrovirals during chemotherapy, there is no question that their use overall in patients with HIV lymphoma has improved survival. Concerns regarding synergistic bone marrow toxicities with chemotherapy and cART are mitigated with the use of cART regimens that avoid bone marrow-toxic antiretrovirals. As in most situations in patients with HIV disease, those with higher CD4+ T cell counts tend to fare better. Response rates as high as 72% with a median survival of 33 months and disease-free intervals up to 9 years have been reported. Treatment of primary CNS lymphoma remains a significant challenge. Treatment is complicated by the fact that this illness usually occurs in patients with advanced HIV disease. Palliative measures such as radiation therapy provide some relief. The prognosis remains poor in this group, with a 2-year survival of 29%.

Multicentric Castleman's disease is a KSHV-associated lymphoproliferative disorder that is seen with an increased frequency in patients with HIV infection. While not a true malignancy, it shares many features with lymphoma including generalized lymphadenopathy, hepatosplenomegaly, and systemic symptoms of fever, fatigue, and weight loss. Pulmonary symptoms may be seen in ~50% of patients. KS is present in 75–82% of cases. Lymph node biopsies reveal a predominance of interfollicular plasma cells and/or germinal centers with vascularization and an "onionskin" (hyaline vascular) appearance. Prior to the availability of cART, HIV-infected patients with multicentric Castleman's disease had a 15-fold increased risk of developing non-Hodgkin's lymphoma compared with HIV-infected patients in general. Treatment typically involves chemotherapy. Anecdotal reports of success with rituximab suggest that more specific treatment may be successful, although, in one series treatment with rituximab was associated with worsening of coexisting KS. The median survival of patients with treated multicentric Castleman's disease pre-cART was initially reported as 14 months. This has increased to a 2-year survival of more than 90% in the era of cART.

Evidence of infection with *human papillomavirus* (HPV), associated with *intraepithelial dysplasia of the cervix* or *anus*, is approximately twice as common in HIV-infected individuals as in the general population and can lead to intraepithelial neoplasia and eventually invasive cancer. In a series of studies, HIV-infected men were examined for evidence of anal dysplasia, and Papanicolaou (Pap) smears were found to be abnormal in 20–80%. These changes tend to persist and are generally not affected by cART, raising the possibility of a subsequent transition to a more malignant condition. While the incidence of an abnormal Pap smear of the cervix is ~5% in otherwise healthy women, the incidence of abnormal cervical smears in women with HIV infection is 30–60%, and *invasive cervical cancer* is included as an AIDS-defining condition. While only small increases in the absolute numbers of cervical or anal cancers have been seen as a consequence of HIV infection, the relative risk of these conditions when one compares HIV-infected to -noninfected men and women is on the order of 10- to 100-fold. Given the high rates of dysplasia and relative risks for cervical and anal cancer, a comprehensive gynecologic and rectal examination, including Pap smear, is indicated at the initial evaluation and 6 months later for all patients with HIV infection. If these examinations are negative at both time points, the patient should be followed with yearly evaluations. If an initial or repeat Pap smear shows evidence of severe inflammation with reactive squamous changes, the next Pap smear should be performed at 3 months. If, at any time, a Pap smear shows evidence of squamous intraepithelial lesions, colposcopic examination with biopsies as indicated should be performed. The 2-year survival rate for HIV-infected patients with invasive cervical cancer is 64% compared with 79% in non-HIV-infected patients. In addition to rectal and cervical lesions, HPV can also lead to head and

1272 neck cancers. In one study of men who have sex with men, 25% were found to have oral HPV; high-risk HPV genotypes were three times more common in the HIV-infected men. The most common HPV genotypes in the general population and the genotypes upon which current HPV vaccines are based are 6, 11, 16, and 18. This is not the case in the HIV-infected population, where other genotypes such as 58 and 53 also are prominent. This raises concerns about the level of effectiveness of the current HPV vaccines for HIV-infected patients. Despite this, it is recommended that patients with HIV infection be vaccinated against HPV.

IDIOPATHIC CD4+ T LYMPHOCYTOPENIA

A syndrome was recognized in 1992 characterized by an absolute CD4+ T cell count of $<300/\mu\text{L}$ or $<20\%$ of total T cells on a minimum of two occasions at least 6 weeks apart; no evidence of HIV-1, HIV-2, HTLV-1, or HTLV-2 on testing; and the absence of any defined immunodeficiency or therapy associated with decreased levels of CD4+ T cells. By mid-1993, ~100 patients had been described. After extensive multicenter investigations, a series of reports were published in early 1993, which together allowed a number of conclusions. Idiopathic CD4+ lymphocytopenia (ICL) is a very rare syndrome, as determined by studies of blood donors and cohorts of HIV-seronegative men who have sex with men. Cases were clearly identified as early as 1983 and were remarkably similar to the clinical features of ICL that had been identified decades earlier. The definition of ICL based on CD4+ T cell counts coincided with the ready availability of testing for CD4+ T cells in patients suspected of being immunodeficient. Although, as a result of immune deficiency, certain patients with ICL develop some of the opportunistic diseases (particularly cryptococcosis, nontuberculous mycobacterial infections, and cervical dysplasia) seen in HIV-infected patients, the syndrome is demographically, clinically, and immunologically unlike HIV infection and AIDS. Fewer than half of the reported ICL patients had risk factors for HIV infection, and there were wide geographic and age distributions. The fact that a significant proportion of patients did have risk factors probably reflects a selection bias, in that physicians who take care of HIV-infected patients are more likely to monitor CD4+ T cells. Approximately half of the patients are women, compared with approximately one-third among HIV-infected individuals in the United States. Many patients with ICL remained clinically stable, and their condition did not deteriorate progressively as is common with seriously immunodeficient HIV-infected patients. Approximately 15% of patients with ICL experience spontaneous reversal of the CD4+ T lymphocytopenia. Immunologic abnormalities in ICL are somewhat different from those of HIV infection. ICL patients often have increases in CD4+ T cell activation with decreases in CD8+ T cells and B cells. Furthermore, immunoglobulin levels are either normal or, more commonly, decreased in patients with ICL, compared with the usual hypergammaglobulinemia of HIV-infected individuals. Virologic studies of these patients have revealed no evidence of HIV-1, HIV-2, HTLV-1, or HTLV-2 or of any other mononuclear cell-tropic virus. Furthermore, there has been no epidemiologic evidence to suggest that a transmissible microbe was involved. The cases of ICL have been widely dispersed, with no clustering. Close contacts and sexual partners who were studied were clinically well and were serologically, immunologically, and virologically negative for HIV. ICL is a heterogeneous syndrome, and it is highly likely that there is no common cause; however, there may be common causes among subgroups of patients that are currently unrecognized.

Patients who present with laboratory data consistent with ICL should be worked up for underlying diseases that could be responsible for the immune deficiency. If no underlying cause is detected, no specific therapy should be initiated. However, if opportunistic diseases occur, they should be treated appropriately (see above). Depending on the level of the CD4+ T cell count, patients should receive prophylaxis for the commonly encountered opportunistic infections.

GENERAL PRINCIPLES OF PATIENT MANAGEMENT

The CDC guidelines call for the testing for HIV infection to be a part of routine medical care. It is recommended that the patient be informed of the intention to test, as is the case with other routine laboratory determinations, and be given the opportunity to “opt out.” Such an approach is critical to the goal of identifying as many infected individuals as possible since 16–18% of the >1.1 million individuals in the United States who are HIV-infected are not aware of their status. Under these circumstances of routine testing, although it is desirable, pretest counseling may not always be built into the testing process. However, no matter how well prepared a patient is for adversity, the discovery of a diagnosis of HIV infection is a devastating event. Thus, physicians should be sensitive to this fact and, where possible, execute some degree of pretest counseling to at least partially prepare the patient should the results demonstrate the presence of HIV infection. Following a diagnosis of HIV infection, the health care provider should be prepared to immediately activate support systems for the newly diagnosed patient. These should include an experienced social worker or nurse who can spend time talking to the person and ensuring that he or she is emotionally stable. Most communities have HIV support centers that can be of great help in these difficult situations.

The treatment of patients with HIV infection requires not only a comprehensive knowledge of the possible disease processes that may occur and up-to-date knowledge of and experience with cART, but also the ability to deal with the problems of a chronic, potentially life-threatening illness. A comprehensive knowledge of internal medicine is required to deal with the changing spectrum of illnesses associated with HIV infection, many of which are similar to a state of accelerated aging. Great advances have been made in the treatment of patients with HIV infection. The appropriate use of potent cART and other treatment and prophylactic interventions are of critical importance in providing each patient with the best opportunity to live a long and healthy life despite the presence of HIV infection. In contrast to the earlier days of this epidemic, a diagnosis of HIV infection need no longer be equated with having an inevitably fatal disease. In addition to medical interventions, the health care provider has a responsibility to provide each patient with appropriate counseling and education concerning their disease as part of a comprehensive care plan. Patients must be educated about the potential transmissibility of their infection and about the fact that while health care providers may refer to levels of the virus as “undetectable,” this is more a reflection of the sensitivity of the assay being used to measure the virus than a comment on the presence or absence of the virus. It is important for patients to be aware that the virus is still present and capable of being transmitted at all stages of HIV disease. Thus, there must be frank discussions concerning sexual practices and the sharing of syringes and other paraphernalia used in illicit drug use. The treating physician not only must be aware of the latest medications available for patients with HIV infection but also must educate patients concerning the natural history of their illness and listen and be sensitive to their fears and concerns. As with other diseases, therapeutic decisions should be made in consultation with the patient, when possible, and with the patient’s proxy if the patient is incapable of making decisions. In this regard, it is recommended that all patients with HIV infection, and in particular those with CD4+ T cell counts $<200/\mu\text{L}$, designate a trusted individual with durable power of attorney to make medical decisions on their behalf, if necessary.

Following a diagnosis of HIV infection, there are several examinations and laboratory studies that should be performed to help determine the extent of disease and provide baseline standards for future reference (Table 226-19). In addition to routine chemistry, fasting lipid profile, aspartate aminotransferase, alanine aminotransferase, total and direct bilirubin, fasting glucose and hematology screening panels, Pap smear, urinalysis, and chest x-ray, one should also obtain a CD4+ T cell count, two separate plasma HIV

TABLE 226-19 INITIAL EVALUATION OF THE PATIENT WITH HIV INFECTION

History and physical examination
Routine chemistry and hematology
AST, ALT, direct and indirect bilirubin
Lipid profile and fasting glucose
CD4+ T lymphocyte count
Two plasma HIV RNA levels
HIV resistance testing
HLA-B5701 screening
RPR or VDRL test
Anti- <i>Toxoplasma</i> antibody titer
PPD skin test or IFN- γ release assay
Mini-Mental Status Examination
Serologies for hepatitis A, hepatitis B, and hepatitis C
Immunization with pneumococcal polysaccharide; influenza as indicated
Immunization with hepatitis A and hepatitis B if seronegative
Counseling regarding natural history and transmission
Help contacting others who might be infected

Abbreviations: ALT, alanine aminotransferase; AST, aspartate aminotransferase; PPD, purified protein derivative; RPR, rapid plasma reagin; VDRL, Venereal Disease Research Laboratory.

RNA levels, an HIV resistance test, a rapid plasma reagin or VDRL test, an anti-*Toxoplasma* antibody titer, and serologies for hepatitis A, B, and C. A PPD test or IFN- γ release assay should be done and an MMSE performed and recorded. A pregnancy test should be done in women in whom the drug efavirenz is being considered, and HLA-B5701 testing should be done in all patients in whom the drug abacavir is being considered. Patients should be immunized with pneumococcal polysaccharide, with annual influenza shots, and, if seronegative for these viruses, with HPV, hepatitis A, and hepatitis B vaccines. The status of hepatitis C infection should be determined. In addition, patients should be counseled with regard to sexual practices and needle sharing, and counseling should be offered to those whom the patient knows or suspects may also be infected. Once these baseline activities are performed, short- and long-term medical management strategies should be developed based on the most recent information available and modified as new information becomes available. The field of HIV medicine is changing rapidly, and it is difficult to remain fully up to date. Fortunately there are a series of excellent sites on the Internet that are frequently updated, and they provide the most recent information on a variety of topics, including consensus panel reports on treatment (**Table 226-20**).

ANTIRETROVIRAL THERAPY

Combination antiretroviral therapy (cART), also referred to as highly active antiretroviral therapy (HAART), is the cornerstone of management of patients with HIV infection. Following the initiation of widespread use of cART in the United States in 1995–1996, marked declines were noted in the incidence of most AIDS-defining conditions (Fig. 226-33). Suppression of HIV replication is an important component in prolonging life as well as in improving the quality of life in patients with HIV infection. Adequate suppression requires

TABLE 226-20 RESOURCES AVAILABLE ON THE WORLD WIDE WEB ON HIV DISEASE

www.aidsinfo.nih.gov	AIDSinfo, a service of the U.S. Department of Health and Human Services, posts federally approved treatment guidelines for HIV and AIDS; provides information on federally funded and privately funded clinical trials and CDC publications and data
www.cdcnpi.org	Updates on epidemiologic data and prevention information from the CDC

Abbreviation: CDC, Centers for Disease Control and Prevention.

strict adherence to prescribed regimens of antiretroviral drugs. This has been facilitated by the coformulations of antiretrovirals and the development of once-daily regimens. Unfortunately, many of the most important questions related to the treatment of HIV disease currently lack definitive answers. Among them are the questions of when therapy should be started, what the best initial regimen is, when a given regimen should be changed, and what it should be changed to when a change is made. Notwithstanding these uncertainties, the physician and patient must come to a mutually agreeable plan based on the best available data. In an effort to facilitate this process, the U.S. Department of Health and Human Services makes available on the Internet (www.aidsinfo.nih.gov) a series of periodically updated guidelines, including “Guidelines for the Use of Antiretroviral Agents in HIV-Infected Adults and Adolescents” and “Guidelines for the Prevention of Opportunistic Infections in Persons Infected with Human Immunodeficiency Virus.” At present, an extensive clinical trials network, involving both clinical investigators and patient advocates, is in place attempting to develop improved approaches to therapy. Consortia comprising representatives of academia, industry, independent foundations, and the federal government are involved in the process of drug development, including a wide-ranging series of clinical trials. As a result, new therapies and new therapeutic strategies are continually emerging. New drugs are often available through expanded-access programs prior to official licensure. Given the complexity of this field, decisions regarding cART are best made in consultation with experts.

Currently available drugs for the treatment of HIV infection as part of a combination regimen fall into four categories: those that inhibit the viral reverse transcriptase enzyme (nucleoside and nucleotide reverse transcriptase inhibitors; nonnucleoside reverse transcriptase inhibitors), those that inhibit the viral protease enzyme (protease inhibitors), those that inhibit the viral integrase enzyme (integrase inhibitors), and those that interfere with viral entry (fusion inhibitors; CCR5 antagonists) (**Table 226-21**; **Fig. 226-45**).

The FDA-approved reverse transcriptase inhibitors include the *nucleoside analogues* zidovudine, didanosine, zalcitabine, stavudine, lamivudine, abacavir, and emtricitabine; the *nucleotide analogue* tenofovir; and the *nonnucleoside reverse transcriptase inhibitors* nevirapine, delavirdine, efavirenz, and etravirine (**Table 226-21**; **Fig. 226-45**). These represent the first class of drugs licensed for the treatment of HIV infection. They are indicated for this use as part of combination regimens. It should be stressed that none of these drugs should be used as monotherapy for HIV infection due to the relative ease with which drug resistance may develop under such circumstances. Thus, when lamivudine, emtricitabine, or tenofovir is used to treat hepatitis B infection in the setting of HIV infection, one should ensure that the patient is also on additional antiretroviral medication. The reverse transcriptase inhibitors block the HIV replication cycle at the point of RNA-dependent DNA synthesis, the reverse transcription step. While the nonnucleoside reverse transcriptase inhibitors are quite selective for the HIV-1 reverse transcriptase, the nucleoside and nucleotide analogues inhibit a variety of DNA polymerases in addition to those of the HIV-1 reverse transcriptase. For this reason, serious side effects are more varied with the nucleoside analogues and include mitochondrial damage that can lead to hepatic steatosis and lactic acidosis as well as peripheral neuropathy and pancreatitis. The use of either of the thymidine analogues zidovudine and stavudine has been associated with a syndrome of hyperlipidemia, glucose intolerance/insulin resistance, and fat redistribution often referred to as *lipodystrophy syndrome* (discussed in “Diseases of the Endocrine System and Metabolic Disorders,” above). The reverse transcriptase inhibitors preferred for use according to the DHHS Panel on the use of antiretroviral drugs are lamivudine, emtricitabine, abacavir, tenofovir, and rilpivirine.

Lamivudine (3TC; 2',3'-dideoxy-3'-thiacytidine) is the fifth of the nucleoside analogues to be licensed in the United States. It is the negative enantiomer of a dideoxy analogue of cytidine. In actual practice, lamivudine or the closely related drug emtricitabine (see

TABLE 226-21 ANTIRETROVIRAL DRUGS USED IN THE TREATMENT OF HIV INFECTION

Drug	Status	Indication	Dose in Combination	Supporting Data	Toxicity
Nucleoside or Nucleotide Reverse Transcriptase Inhibitors					
Zidovudine (AZT, azidothymidine, Retrovir, 3'azido-3'-deoxythymidine)	Licensed	Treatment of HIV infection in combination with other antiretroviral agents Prevention of maternal-fetal HIV transmission	200 mg q8h or 300 mg bid	19 vs 1 death in original placebo-controlled trial in 281 patients with AIDS or ARC In pregnant women with CD4+ T cell count $\geq 200/\mu\text{L}$, AZT PO beginning at weeks 14–34 of gestation plus IV drug during labor and delivery plus PO AZT to infant for 6 weeks decreased transmission of HIV by 67.5% (from 25.5% to 8.3%), n = 363	Anemia, granulocytopenia, myopathy, lactic acidosis, hepatomegaly with steatosis, headache, nausea, nail pigmentation, lipid abnormalities, lipoatrophy, hyperglycemia
Lamivudine (Epivir, 2'3'-dideoxy-3'-thiacytidine, 3TC)	Licensed	In combination with other antiretroviral agents for the treatment of HIV infection	150 mg bid 300 mg qd	In combination with AZT superior to AZT alone with respect to changes in CD4+ T cell counts in 495 patients who were zidovudine-naïve and 477 patients who were zidovudine-experienced; overall CD4+ T cell counts for the zidovudine group were at baseline by 24 weeks, while in the group treated with zidovudine plus lamivudine, they were 10–50 cells/ μL above baseline; 54% decrease in progression to AIDS/death compared with AZT alone	Flare of hepatitis in HBV-co-infected patients who discontinue drug
Emtricitabine (FTC, Emtriva)	Licensed	In combination with other antiretroviral agents for the treatment of HIV infection	200 mg qd	Comparable to stavudine in combination with didanosine and efavirenz in 571 treatment-naïve patients; similar to 3TC in combination with AZT or stavudine + NNRTI or PI in 440 patients doing well for ≥ 12 weeks on a 3TC regimen	Hepatotoxicity in HBV-co-infected patients who discontinue drug, skin discoloration
Abacavir (Ziagen)	Licensed	For treatment of HIV infection in combination with other antiretroviral agents	300 mg bid	Abacavir + AZT + 3TC equivalent to indinavir + AZT + 3TC with regard to viral load suppression ($\sim 60\%$ in each group with < 400 HIV RNA copies/mL plasma) and CD4+ T cell increase ($\sim 100/\mu\text{L}$ in each group) at 24 weeks	Hypersensitivity reaction In HLA-B5701+ individuals (can be fatal); fever, rash, nausea, vomiting, malaise or fatigue, and loss of appetite
Tenofovir (Viread)	Licensed	For use in combination with other antiretroviral agents when treatment is indicated	300 mg qd	Reduction of ~ 0.6 log in HIV-1 RNA levels when added to background regimen in treatment-experienced patients	Renal, osteomalacia, flare of hepatitis in HBV-co-infected patients who discontinue drug
Non-Nucleoside Reverse Transcriptase Inhibitors					
Nevirapine (Viramune)	Licensed	In combination with other antiretroviral agents for treatment of progressive HIV infection	200 mg/d \times 14 days then 200 mg bid or 400 mg extended release qd	Increase in CD4+ T cell count, decrease in HIV RNA when used in combination with nucleosides	Skin rash, hepatotoxicity
Efavirenz (Sustiva)	Licensed	For treatment of HIV infection in combination with other antiretroviral agents	600 mg qhs	Efavirenz + AZT + 3TC comparable to indinavir + AZT + 3TC with regard to viral load suppression (a higher percentage of the efavirenz group achieved viral load < 50 copies/mL, but the discontinuation rate in the indinavir group was unexpectedly high, accounting for most treatment "failures"); CD4 cell increase ($\sim 140/\mu\text{L}$ in each group) at 24 weeks	Rash, dysphoria, elevated liver function tests, drowsiness, abnormal dreams, depression, lipid abnormalities, potentially teratogenic
Etravirine (Intelence)	Licensed	In combination with other antiretroviral agents in treatment-experienced patients whose HIV is resistant to nonnucleoside reverse transcriptase inhibitors and other antiretroviral medications	200 mg bid	Higher rates of HIV RNA suppression to < 50 copies/mL (56% vs 39%); greater increases in CD4+ T cell count (89 vs 64 cells) compared to placebo when given in combination with an optimized background regimen	Rash, nausea, hypersensitivity reactions

(Continued)

TABLE 226-21 ANTIRETROVIRAL DRUGS USED IN THE TREATMENT OF HIV INFECTION (CONTINUED)

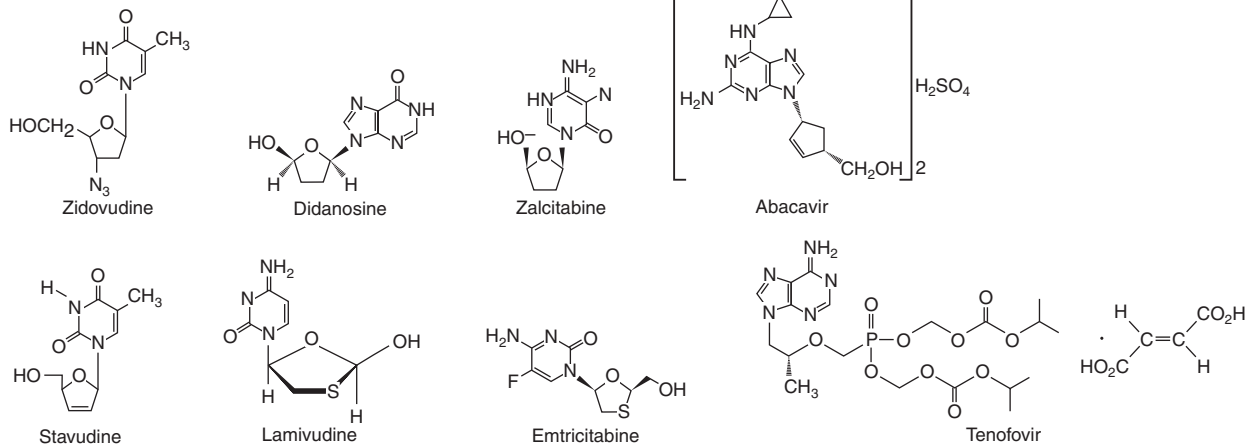
Drug	Status	Indication	Dose in Combination	Supporting Data	Toxicity
Rilpivirine (Edurant)	Licensed	In combination with other drugs in previously untreated patients when treatment is indicated.	25 mg qd	Noninferior to efavirenz with respect to suppression at week 48 in 1368 treatment-naïve individuals except in patients with pretherapy HIV RNA levels >100,000 where it was inferior	Nausea, dizziness, somnolence, vertigo, less CNS toxicity and rash than Efavirenz
Protease Inhibitors					
Ritonavir (Norvir)	Licensed	In combination with other antiretroviral agents for treatment of HIV infection when treatment is warranted	600 mg bid (also used in lower doses as pharmacokinetic booster)	Reduction in the cumulative incidence of clinical progression or death from 34% to 17% in patients with CD4+ T cell count <100/μL treated for a median of 6 months	Nausea, abdominal pain, hyperglycemia, fat redistribution, lipid abnormalities, may alter levels of many other drugs, paresthesias, hepatitis
Atazanavir (Reyataz)	Licensed	For treatment of HIV infection in combination with other antiretroviral agents	400 mg qd or 300 mg qd + ritonavir 100 mg qd when given with efavirenz	Comparable to efavirenz when given in combination with AZT + 3TC in a study of 810 treatment-naïve patients; comparable to nevirapin when given in combination with stavudine + 3TC in a study of 467 treatment-naïve patients	Hyperbilirubinemia, PR prolongation, nausea, vomiting, hyperglycemia, fat maldistribution, rash transaminase elevations, renal stones
Darunavir (Prezista)	Licensed	In combination with 100 mg ritonavir for combination therapy in treatment-experienced adults	600 mg + 100 mg ritonavir twice daily with food	At 24 weeks, patients with prior extensive exposure to antiretrovirals treated with a new combination including darunavir showed a -1.89-log change in HIV RNA levels and a 92-cell increase in CD4+ T cells compared with -0.48 log and 17 cells in the control arm	Diarrhea, nausea, headache, skin rash, hepatotoxicity, hyperlipidemia, hyperglycemia
Entry Inhibitors					
Enfuvirtide (Fuzeon)	Licensed	In combination with other agents in treatment-experienced patients with evidence of HIV-1 replication despite ongoing antiretroviral therapy	90 mg SC bid	In treatment of experienced patients, superior to placebo when added to new optimized background (37% vs 16% with <400 HIV RNA copies/mL at 24 weeks; + 71 vs + 35 CD4+ T cells at 24 weeks)	Local injection reactions, hypersensitivity reactions, increased rate of bacterial pneumonia
Maraviroc (Selzentry)	Licensed	In combination with other antiretroviral agents in adults infected with only CCR5-tropic HIV-1	150–600 mg bid depending on concomitant medications (see text)	At 24 weeks, among 635 patients with CCR5-tropic virus and HIV-1 RNA >5000 copies/mL despite at least 6 months of prior therapy with at least 1 agent from 3 of the 4 antiretroviral drug classes, 61% of patients randomized to maraviroc achieved HIV RNA levels <400 copies/mL compared with 28% of patients randomized to placebo	Hepatotoxicity, nasopharyngitis, fever, cough, rash, abdominal pain, dizziness, musculoskeletal symptoms
Integrase Inhibitor					
Raltegravir (Isentress)	Licensed	In combination with other antiretroviral agents	400 mg bid	At 24 weeks, among 436 patients with 3-class drug resistance, 76% of patients randomized to receive raltegravir achieved HIV RNA levels <400 copies/mL compared with 41% of patients randomized to receive placebo	Nausea, headache, diarrhea, CPK elevation, muscle weakness, rhabdomyolysis
Elvitegravir (Available only in combination with cobicistat, tenofovir, and emtricitabine [Stribild])	Licensed	Fixed-dose combination	1 tablet daily	Noninferior to raltegravir or atazanavir/ritonavir in treatment-experienced patients.	Diarrhea, nausea, upper respiratory infections, headache
Dolutegravir (Tivicay)	Licensed	In combination with other antiretroviral agents	50 mg daily for treatment-naïve patients 50 mg twice daily for treatment-experienced patients or those also receiving efavirenz or rifampin	Noninferior to raltegravir, superior to efavirenz or darunavir/ritonavir	Insomnia, headache, hypersensitivity reactions, hepatotoxicity

Abbreviations: ARC, AIDS-related complex; NRTIs, nonnucleoside reverse transcriptase inhibitors.

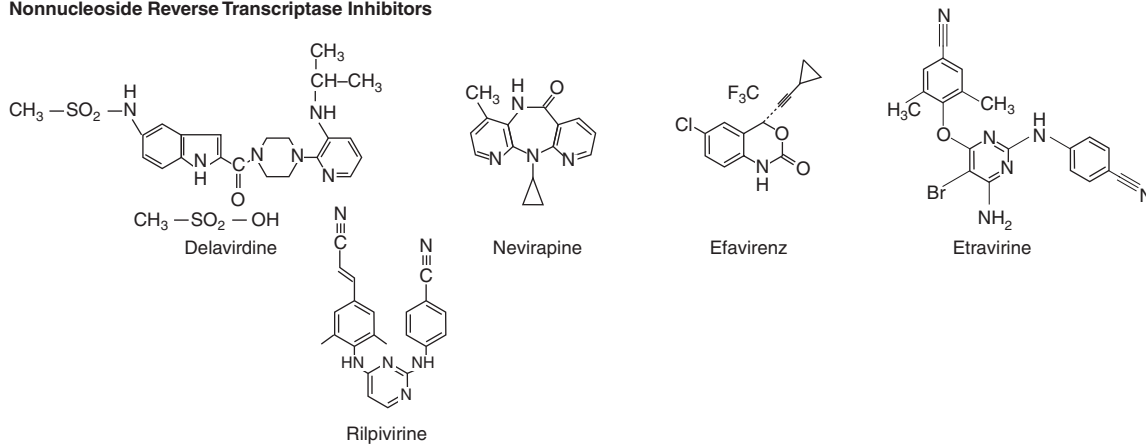
1276 below) is a frequent element of many different combination regimens currently in use. These two drugs and the nucleotide reverse transcriptase inhibitor tenofovir (see below) also have activity against hepatitis B virus. For this reason flares of hepatitis may be seen in co-infected patients starting and/or or stopping any of these three agents due to the confounding issues of direct effects

on hepatitis B, direct effects on HIV, and immune reconstitution (see above). To prevent the development of resistant strains of HIV, these drugs should never be used on their own for the treatment of hepatitis B in the patient with HIV infection. Lamivudine is available either alone or in coformulations including zidovudine and/or abacavir (Table 226-22). One reason behind the excellent synergy

Nucleoside or Nucleotide Reverse Transcriptase Inhibitors



Nonnucleoside Reverse Transcriptase Inhibitors



Protease Inhibitors

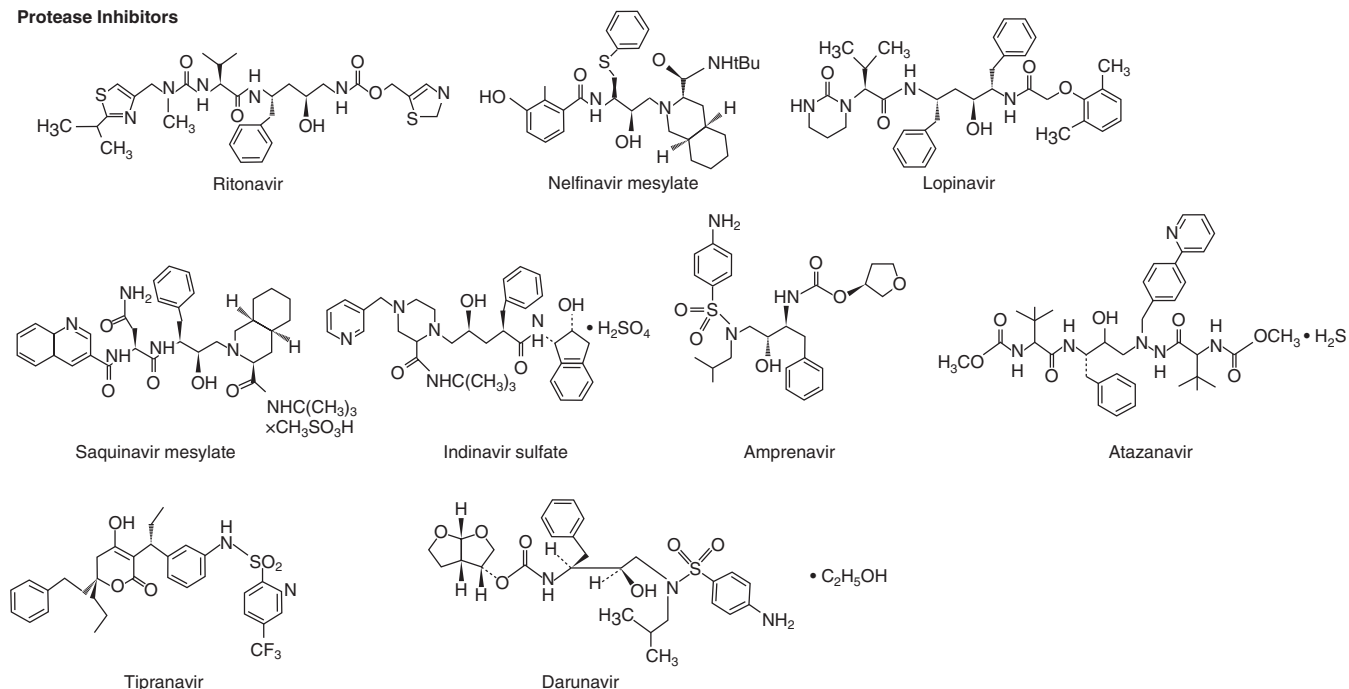


FIGURE 226-45 Molecular structures of antiretroviral agents.

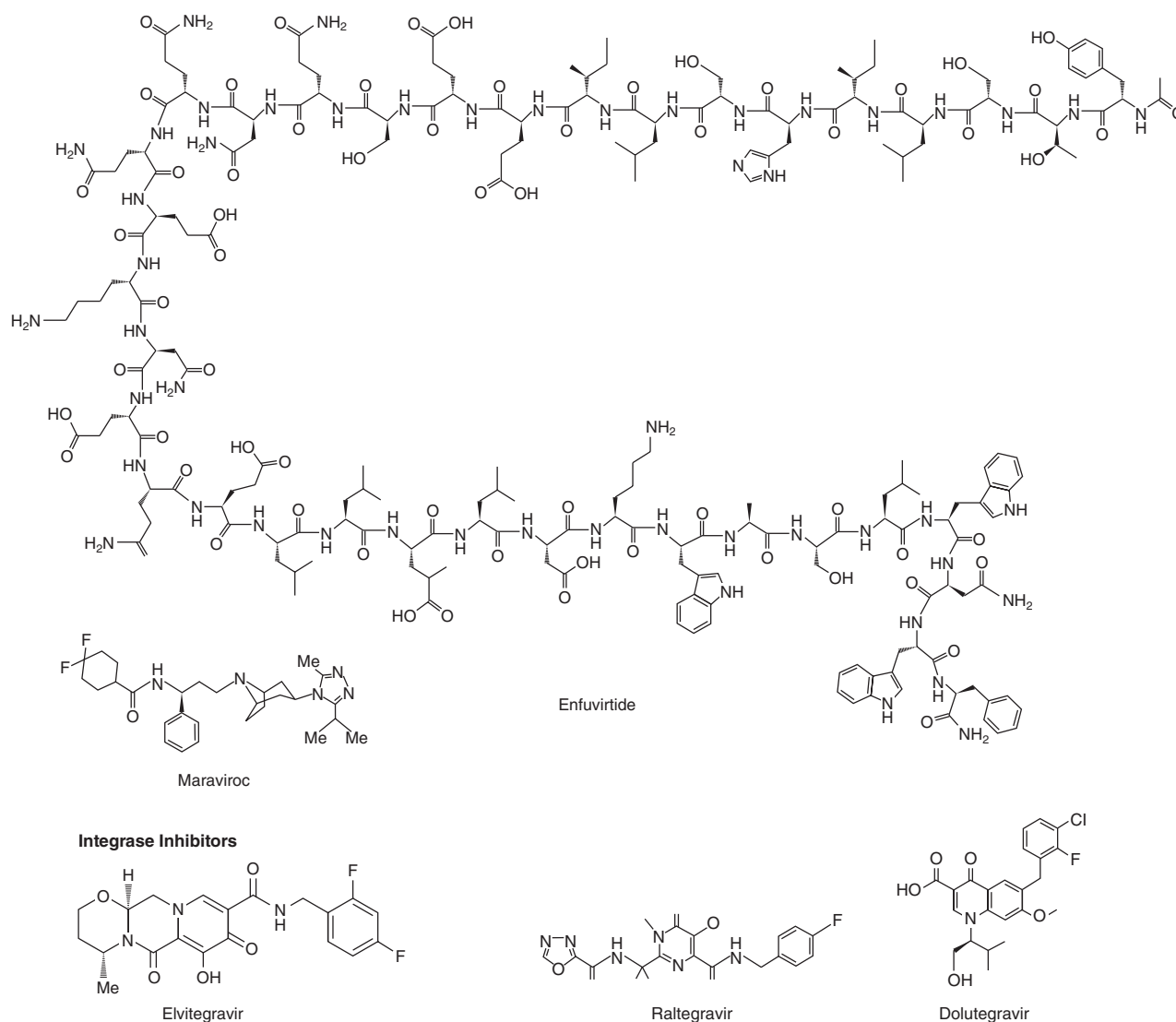


FIGURE 226-45 (Continued)

seen between lamivudine and the other nucleoside analogues may be that strains of HIV resistant to lamivudine (M184V substitution) appear to have enhanced sensitivity to other nucleosides, and thus development of dual resistance is more difficult. In addition, there is a suggestion that 3TC-resistant strains of HIV may be less virulent and are less able to generate new mutants than are strains of HIV that are 3TC-sensitive. Lamivudine is among the best tolerated and least toxic of the nucleoside analogues.

TABLE 226-22 COMBINATION FORMULATIONS OF ANTIRETROVIRAL DRUGS

Name	Combination
Combivir	Zidovudine + lamivudine
Complera	Tenofovir + emtricitabine + rilpivirine
Epzicom	Abacavir + lamivudine
Stribild	Tenofovir + emtricitabine + elvitegravir + cobicistat
Triumeq	Abacavir + lamivudine + dolutegravir
Trizivir	Zidovudine + lamivudine + abacavir
Truvada	Tenofovir + emtricitabine
Atripla	Tenofovir + emtricitabine + efavirenz
Triomune ^a	Stavudine + lamivudine + nevirapine

^aNot licensed in the United States.

Emtricitabine (FTC; 5-fluoro-1-(2R,5S)-[2-(hydroxymethyl)-1,3-oxathiolan-5-yl]cytosine) is the negative enantiomer of a thio analogue of cytidine with a fluorine in the 5 position. It is licensed for use in combination with other antiretroviral agents for treatment of HIV-1 infection in adults. Compared with lamivudine, it is similar in activity and has a longer half-life. It is available either alone or coformulated with tenofovir or tenofovir and efavirenz (Table 226-22). As with lamivudine, resistance to emtricitabine is associated with the M184V mutation in reverse transcriptase. Viruses showing the K65R mutation in reverse transcriptase may have reduced susceptibility to emtricitabine.

Abacavir {(1*S,cis*)-4-[2-amino-6-(cyclopropylamino)-9H-purin-9-yl]-2-cyclopentene-1-methanol sulfate (salt)(2:1)} is a synthetic carbocyclic analogue of the nucleoside guanosine. It is licensed to be used in combination with other antiretroviral agents for the treatment of HIV-1 infection. Hypersensitivity reactions that may occur with initial therapy or rechallenge have been reported in ~4% of patients treated with this drug, and patients developing signs or symptoms of hypersensitivity such as fever, skin rash, fatigue, and GI symptoms should discontinue the drug and not restart it. Fatal hypersensitivity reactions have been reported with rechallenge. Abacavir hypersensitivity occurs with a higher frequency in patients who are HLA-B5701-positive. It is recommended that patients be screened for HLA-B5701 prior to initiation of abacavir and that

1278 abacavir only be used as a last resort and with close monitoring in patients who are HLA-B5701-positive. Abacavir-resistant strains of HIV are typically also resistant to lamivudine, emtricitabine, didanosine, and zalcitabine. In randomized trials abacavir was found to be inferior to tenofovir in patients with baseline HIV RNA levels >100,000 copies/mL. Abacavir is formulated alone as well as in combination with lamivudine, with zidovudine and lamivudine or with lamivudine and dolutegravir.

Tenofovir disoproxil fumarate (9-[(R)-2-[[bis[[[(isopropoxycarbonyl)oxy]methoxy]phosphinyloxy]methoxy]propyl]adenine fumarate [1:1]) is an acyclic nucleoside phosphonate diester analogue of adenosine monophosphate. It undergoes diester hydrolysis to form the nucleoside monophosphate (nucleotide) tenofovir and is the first nucleotide analogue to be licensed for treatment of HIV infection. It is indicated in combination with other antiretroviral agents for the treatment of HIV-1 infection and in combination with emtricitabine for pre-exposure prophylaxis for HIV-1 prevention in populations at high risk of HIV infection. HIV isolates with increased resistance typically express a K65R mutation in reverse transcriptase and a three- to fourfold reduction in sensitivity to tenofovir. Tenofovir is primarily eliminated by the kidneys, and renal impairment including a Fanconi-like syndrome with hypophosphatemia may occur. Tenofovir is contraindicated in patients with renal impairment. An investigational prodrug analogue with less renal toxicity, tenofovir alafenamide fumarate is currently in clinical trials. Small but statistically significant decreases in bone mineral density have been noted in patients receiving tenofovir. Coadministration of tenofovir with didanosine leads to a 60% increase in didanosine levels, and thus doses of didanosine need to be adjusted and patients monitored carefully if these two drugs are used in combination. In addition, CD4+ T cell increases may be blunted in patients on this combination. Coadministration of tenofovir with atazanavir leads to a decrease in atazanavir levels, and thus low-dose ritonavir (see below) needs to be added when these drugs are used in combination. Tenofovir is available alone and coformulated with emtricitabine, emtricitabine and efavirenz, emtricitabine and rilpivirine, or emtricitabine, elvitegravir and cobicistat.

Nevirapine, delavirdine, efavirenz, etravirine, and rilpivirine are nonnucleoside inhibitors of the HIV-1 reverse transcriptase and are licensed for use in combination with nucleoside analogues for the treatment of HIV-infected adults. Coformulations that include efavirenz or nevirapine are available (Table 226-22). These agents inhibit reverse transcriptase by binding to regions of the enzyme outside the active site and causing conformational changes in the enzyme that render it inactive. Although these agents are active in the nanomolar range, they are also very selective for the reverse transcriptase of HIV-1, have no activity against HIV-2, and, when used as monotherapy, are associated with the rapid emergence of drug-resistant mutants (Table 226-21; Fig. 226-46). Efavirenz and rilpivirine are administered once a day, nevirapine and etravirine twice a day, and delavirdine three times a day. All are associated with the development of a maculopapular rash, generally seen within the first few weeks of therapy. While it is possible to treat through this rash, it is important to be sure that one is not dealing with a more severe eruption such as Stevens-Johnson syndrome by looking carefully for signs of mucosal involvement, significant fever, or painful lesions with desquamation. Severe, life-threatening, and in some cases fatal hepatotoxicity, including fulminant and cholestatic hepatitis, hepatic necrosis, and hepatic failure, have been reported in patients treated with nevirapine. There is a suggestion that this is more common in women with higher CD4+ T cell counts. Many patients treated with efavirenz note a feeling of light-headedness, dizziness, or out of sorts following the initiation of therapy. Some complain of vivid dreams. These symptoms tend to disappear after several weeks of therapy. Aside from difficulties with dreams, taking efavirenz at bedtime may minimize the side effects. Efavirenz may cause fetal harm when administered during the first trimester to a pregnant woman. Women of childbearing potential should undergo pregnancy testing prior to initiation of efavirenz. Efavirenz is

commonly used in combination with two nucleoside analogues as part of initial treatment regimens. Etravirine is a diarylpyrimidine derivative currently licensed for treatment of HIV infection in combination with other agents. In contrast to the other nonnucleoside reverse transcriptase inhibitors, which all exhibit cross-resistance, etravirine may be active against strains of HIV that are resistant to other nonnucleoside reverse transcriptase inhibitors. Among its side effects are rash, headache, nausea, and diarrhea. Rilpivirine is effective across a broad range of NNRTI-resistant viruses and shares cross-resistance with etravirine. It is better tolerated and has a higher rate of virologic failure than efavirenz, particularly in those with HIV RNA >100,000. It is only available as part of a combination regimen with tenofovir and emtricitabine.

The HIV-1 protease inhibitors (saquinavir, indinavir, ritonavir, nelfinavir, amprenavir, fosamprenavir, lopinavir/ritonavir, atazanavir, tipranavir, and darunavir) are a major part of the therapeutic armamentarium of antiretrovirals. When used as part of initial regimens in combination with reverse transcriptase inhibitors, these agents have been shown to be capable of suppressing levels of HIV replication to under 50 copies/mL in the majority of patients for a minimum of 5 years. As in the case of reverse transcriptase inhibitors, resistance to protease inhibitors can develop rapidly in the setting of monotherapy, and thus these agents should be used only as part of combination therapeutic regimens. A summary of known resistance mutations for protease inhibitors is shown in Fig. 226-46. The protease inhibitors preferred for use according to the DHHS Panel on the use of antiretroviral drugs are ritonavir (only as a pharmacokinetic enhancer), atazanavir, and darunavir.

Ritonavir was the first protease inhibitor for which clinical efficacy was demonstrated. In a study of 1090 patients with CD4+ T cell counts <100/ μ L who were randomized to receive either placebo or ritonavir in addition to any other licensed medications, patients receiving ritonavir had a reduction in the cumulative incidence of clinical progression or death from 34% to 17%. Mortality decreased from 10.1% to 5.8%. At full doses, ritonavir is poorly tolerated. Among the main side effects are nausea, diarrhea, abdominal pain, hyperlipidemia, and circumoral paresthesia. Ritonavir has a high affinity for several isoforms of cytochrome P450 (3A4, 2D6), and its use can result in large increases in the plasma concentrations of drugs metabolized by these pathways. Among the agents affected in this manner are most other protease inhibitors, macrolide antibiotics, *R*-warfarin, ondansetron, rifabutin, most calcium channel blockers, glucocorticoids, and some of the chemotherapeutic agents used to treat KS and/or lymphomas. In addition, ritonavir may increase the activity of glucuronyltransferases, thus decreasing the levels of drugs metabolized by this pathway. Overall, great care must be taken when prescribing additional drugs to patients taking protease inhibitors in general and ritonavir in particular. As mentioned above, the pharmacodynamic boosting property of ritonavir, seen with doses as low as 100–200 mg once or twice a day, is often used in the setting of cART for HIV infection to derive more convenient regimens. For example, when given with low-dose ritonavir, saquinavir and indinavir can be given on twice-a-day schedules and taken with food.

Atazanavir is an azapeptide inhibitor of the HIV-1 protease that was licensed in 2003. An advantage of atazanavir is that total cholesterol and triglyceride levels do not increase as much with atazanavir as with other protease inhibitors. This coupled with the fact that it can be given on a once-daily schedule made atazanavir a popular component of initial treatment regimens following its licensure. Atazanavir is associated with increases in serum bilirubin, renal stones, and prolongations of the ECG PR interval. Atazanavir-resistant isolates emerging in previously treatment-naïve individuals frequently harbor an I50L substitution. This mutation in some instances is associated with increased sensitivity to other protease inhibitors. Atazanavir requires an acidic gastric pH for absorption, and its use in combination with a proton pump inhibitor is contraindicated due to concerns about absorption. Atazanavir is an inhibitor of cytochrome P3A, and its use may be associated with

Mutations in the Protease Gene Associated with Resistance to Protease Inhibitors

Atazanavir +/- ritonavir	L 10	G 16	K 20	L 24	V 32	L 33	E 34	M 36	M 46	G 48	I 50	F 53	I 54	D 60	I 62	I 64	A 71	G 73	V 82	I 84	I 85	N 88	L 90	I 93
	I	E	R	I	I	I	Q	I	I	V	L	L	L	E	V	L	V	C	A	V	V	S	M	L
	F	M	I		F	L		L	L		Y	V				M	I	S	T				M	L
	V	I			V	V		V								V	T	T	F					
	C	T	V								M	T	A			L	A	I						
Darunavir/ ritonavir	V 11				V 32	L 33			I 47	I 50	I 54						T 74	L 76		I 84			L 89	
	I				I	F			V	V	M	L					P	V		V			V	
Fosamprenavir/ ritonavir	L 10				V 32				M 46	I 47	I 50	I 54					G 73	L 76	V 82	I 84			L 90	
	F				I				I	V	V	L					S	V	A	V			M	
	I								L			V							F					
	R											M							S					
	V											T							T					
Indinavir/ ritonavir	L 10	K 20	L 24		V 32			M 36	M 46		I 54						A 71	G 73	L 76	V 77	V 82	I 84	L 90	
	I	M	I		I			I	I		V						V	S	V	I	A	V	M	
	R	R						L									T	A			F			
	V																				T			
Lopinavir/ ritonavir	L 10	K 20	L 24		V 32	L 33			M 46	I 47	I 50	F 53	I 54			L 63	A 71	G 73	L 76	V 82	I 84		L 90	
	F	M	I		I	F			I	V	V	L	V			P	V	S	V	A	V		M	
	I	R			I				L	A		L	A						T		F			
	R											L	A								T			
	V											M	T	S							S			
Nelfinavir	L 10				D 30			M 36	M 46								A 71		V 77	V 82	I 84	N 88	L 90	
	F				N			I	I								V		I	A	V	D	M	
	I								L								T			F		S		
																				T				
																				S				
Saquinavir/ ritonavir	L 10		L 24						G 48		I 54			I 62			A 71	G 73	V 77	V 82	I 84		L 90	
	I		I						V		V			V			V	S	I	A	V		M	
	R										L						T			F				
	V																			T				
																				S				
Tipranavir ritonavir	L 10				L 33		M 36	K 43	M 46	I 47	I 54	Q 58		H 69		T 74	V 82	N 83	I 84				L 89	
	V				F		L	T	L	V	A	E		K		P	L	D	V				M	
							V				M	V		R			T						I	
											V													

Mutations in the Envelope Gene Associated with Resistance to Entry Inhibitors

Enfuvirtide	G 36	I 37	V 38	Q 39	Q 40	N 42	N 43
	D	V	A	R	H	T	D
	S		M				
			E				
Maraviroc	Activity limited to patients with R5 viruses						

Mutations in the Integrase Gene Associated with Resistance to Integrase Strand Transfer Inhibitors

Dolutegravir	F 121	E 138	G 140	Q 148					
	Y	A	S	H					
		K	A						
Elvitegravir	T 66	E 92	T 97	F 121	S 147	Q 148	N 155		
	I	Q	A	Y	G	H	H		
	A	G			H				
	K				K				
Raltegravir	L 74	E 92	T 97	F 121	E 138	G 140	Y 143	Q 148	N 155
	M	Q	A	Y	A	A	R	H	H
					K	S	H	K	R
							C	R	

FIGURE 226-46 (Continued)

increased levels of calcium channel blockers, macrolide antibiotics, HMG-CoA reductase inhibitors, and sildenafil. Levels of atazanavir are lower in the presence of tenofovir or efavirenz. In these settings, levels of atazanavir should be boosted with the use of low-dose ritonavir. In a head-to-head comparison, more patients discontinued atazanavir than either darunavir or raltegravir. The

main reasons for discontinuation were bilirubin elevations and gastrointestinal side effects.

Darunavir is a nonpeptidic HIV protease inhibitor initially licensed in 2006. It is indicated for coadministration with 100 mg of ritonavir and other antiretroviral agents for the treatment of HIV infection. In initial studies in treatment-experienced subjects, 46% of

patients achieved a reduction in HIV RNA viral loads to <50 copies/mL. Studies in treatment-naïve patients demonstrated efficacy superior to lopinavir/ritonavir-containing regimens but inferior to dolutegravir. Skin rash, which may be severe, is seen in 7% of patients and may be related to the sulfonamide moiety contained in the molecule. GI intolerance and headache are the other most frequent side effects.

Entry inhibitors act by interfering with the binding of HIV to its receptor or co-receptor or by interfering with the process of fusion (see above). The first drug in this class to be licensed was the fusion inhibitor *enfuvirtide*, or T-20, followed by the CCR5 antagonist *maraviroc*. A variety of additional small molecules that bind to HIV-1 co-receptors are currently in clinical trials.

Enfuvirtide is a linear 36-amino-acid synthetic peptide with the N terminus acetylated and the C terminus a carboxamide. It is composed of naturally occurring L-amino acid residues and interferes with the fusion of the viral and cellular membranes by binding to the HR1 region in the gp41 subunit of the HIV-1 envelope. This binding interferes with the coil-coil interaction required to approximate the viral envelope and the host cell membrane during the process of viral fusion. *Enfuvirtide* was licensed in 2003 for treatment of HIV-1 infection in combination with other antiretroviral agents in treatment-experienced patients with ongoing viral replication despite antiretroviral therapy. *Enfuvirtide* is not active against HIV-2. *Enfuvirtide*-resistant isolates of HIV exhibit amino acid changes in positions 36–45 of gp41. In two independent studies, patients who had persistent viremia despite prior treatment with agents from all three available classes of drugs were randomized to receive an individualized regimen (based on prior treatment history and resistance profile) with or without *enfuvirtide*. The change in plasma HIV-1 RNA from baseline was ~1 log greater (–1.53 vs –0.68) in patients randomized to receive *enfuvirtide*. Among the drawbacks of this agent are the requirement for twice-a-day injection, the occurrence of injection-site reactions in close to 100% of patients, and an increase in bacterial pneumonia in the *enfuvirtide*-treated patients compared with the control patients (4.68 vs 0.61 events per 100 patient-years) in the phase III studies.

Maraviroc is a CCR5 antagonist that interferes with HIV binding at the stage of co-receptor engagement. It was licensed in 2007 for treatment of HIV infection in combination with other agents in treatment-experienced patients infected with only CCR5-tropic (R5) virus resistant to multiple agents. The license was extended in 2009 to include treatment-naïve patients with R5 virus. A co-receptor tropism assay should be performed if one is considering the use of *maraviroc* to ensure that the potential patient is only harboring R5 viruses. In phase III trials of treatment-experienced patients randomized to receive optimal therapy plus *maraviroc* or placebo, 61% of patients randomized to *maraviroc* achieved HIV RNA levels <400 copies/mL compared with 28% of patients randomized to placebo. An allergic reaction-associated hepatotoxicity has been reported with *maraviroc*. Among the most common side effects of *maraviroc* are dizziness due to postural hypotension, cough, fever, colds, rash, muscle and joint pain, and stomach pain. *Maraviroc* is a substrate of CYP3A and Pgp, and the recommend dose varies depending on concomitant medications. In combination with nucleoside analogues, tipranavir/ritonavir, *enfuvirtide*, and/or nevirapine, the dose is 300 mg twice daily. In the presence of CYP3A inhibitors, such as most protease inhibitors, the dose is 150 mg twice daily. In the presence of CYP3A inducers such as efavirenz, the dose is 600 mg twice daily.

Integrase inhibitors act by blocking the action of the HIV integrase enzyme and thus preventing integration of the HIV provirus into the host cell genome. They are among the most potent and safest of the antiretroviral drugs and frequently part of initial combination regimens. The three licensed integrase inhibitors are *raltegravir*, *elvitegravir* and *dolutegravir*.

Raltegravir is an inhibitor of the viral enzyme integrase and the first of this class to be approved. It acts by interfering with the binding of the preintegration complex to host DNA and as such is

referred to as an integrase strand transfer inhibitor (INSTI). *Raltegravir* was approved in 2007 for treatment of HIV infection in combination with other agents in treatment-experienced patients, and the approval was extended in 2009 to include treatment-naïve patients. *Raltegravir* exhibits a wide range of activity against HIV-1 and HIV-2, including viruses with multiple resistance mutations to other classes of drugs. As with several other compounds, resistance to *raltegravir* comes at the expense of replicative fitness. In two phase III studies in which 436 patients with 3-class antiretroviral drug resistance were randomized to an optimized background regimen with *raltegravir* or placebo, 76% of patients receiving *raltegravir* achieved HIV RNA levels <400 copies/mL compared with 41% of patients randomized to the placebo arm. In contrast to many other antiretroviral drugs the side-effect profile of *raltegravir* is minimal, with similar side-effect profiles noted for the *raltegravir* and placebo groups.

Elvitegravir is an integrase inhibitor that was approved in 2012 as part of a fixed-dose combination tablet also containing tenofovir, emtricitabine, and cobicistat (*Stribild*). The cobicistat acts much in the same way as low-dose ritonavir to boost the concentrations of *elvitegravir* by inhibiting CYP3A such that once-a-day dosing of *Stribild* is sufficient. *Elvitegravir* demonstrates cross-resistance with *raltegravir*. In two randomized, controlled trials, *elvitegravir* was found to be noninferior to efavirenz in one study and noninferior to atazanavir/ritonavir in the other. The most common side effects experienced with *elvitegravir* are diarrhea, nausea, upper respiratory infection, and headache. The cobicistat component of the fixed-dose tablet inhibits tubular secretion of creatinine, resulting in increases in serum creatinine, and is not recommended for patients with estimated creatinine clearances <70 mL/min.

Dolutegravir was approved in 2013 for use as part of a combination regimen in either treatment-naïve or -experienced patients. It comes as a 50-mg tablet and is given once daily in treatment-naïve patients and twice daily in treatment-experienced patients. Isolates of HIV that have developed resistance to *raltegravir* or *elvitegravir* may still be sensitive to *dolutegravir*. Its main side effects are insomnia and headache. In two randomized, controlled trials it has been shown to be superior to either efavirenz (n = 833) or darunavir/ritonavir (n = 484) in combination with nucleos(t)ide analogues due to lower rates of discontinuation. In a third trial of 822 patients it was shown to be noninferior to *raltegravir*.

PRINCIPLES OF THERAPY

The principles of therapy for HIV infection have been articulated by a panel sponsored by the U.S. Department of Health and Human Services as a working group of the NIH Office of AIDS Research Advisory Council. These principles are summarized in [Table 226-23](#). As noted in these guidelines, cART of HIV infection does not lead to eradication or cure of HIV. The single possible exception to this is an individual with HIV infection who received an allogeneic stem cell transplant for treatment of acute myelogenous leukemia. His conditioning regimen included cytotoxic chemotherapy, total-body irradiation, and antithymocyte immunoglobulin. The donor cells were homozygous for the CCR5Δ32 mutation (see above) and thus resistant to HIV infection. Despite cART being stopped the day of the transplant, the patient has exhibited no signs of active HIV infection for more than 8 years.

Treatment decisions must take into account the fact that one is dealing with a chronic infection that requires daily therapy. While early therapy is generally the rule in infectious diseases, immediate treatment of every HIV-infected individual upon diagnosis may not be prudent, and therapeutic decisions must take into account the balance between risks and benefits. Patients initiating antiretroviral therapy must be willing to commit to life-long treatment and understand the importance of adherence to their prescribed regimen. The importance of adherence is illustrated by the observation that treatment interruption is associated with rapid increases in HIV RNA levels, rapid declines in CD4+ T cell counts, and an increased risk of clinical progression. While it seems reasonable to assume that the complications associated with cART could be minimized by

TABLE 226-23 PRINCIPLES OF THERAPY OF HIV INFECTION

1. Ongoing HIV replication leads to immune system damage, progression to AIDS, and systemic immune activation.
2. Plasma HIV RNA levels indicate the magnitude of HIV replication and the rate of CD4+ T cell destruction. CD4+ T cell counts indicate the current level of competence of the immune system.
3. Maximal suppression of viral replication is a goal of therapy; the greater the suppression the less likely the appearance of drug-resistant quasiespecies.
4. The most effective therapeutic strategies involve the simultaneous initiation of combinations of effective anti-HIV drugs with which the patient has not been previously treated and that are not cross-resistant with antiretroviral agents that the patient has already received.
5. The antiretroviral drugs used in combination regimens should be used according to optimum schedules and dosages.
6. The number of available drugs is limited. Any decisions on antiretroviral therapy have a long-term impact on future options for the patient.
7. Women should receive optimal antiretroviral therapy regardless of pregnancy status.
8. The same principles apply to children and adults. The treatment of HIV-infected children involves unique pharmacologic, virologic, and immunologic considerations.
9. Compliance is an important part of ensuring maximal effect from a given regimen. The simpler the regimen, the easier it is for the patient to be compliant.

Source: Modified from *Principles of Therapy of HIV Infection*, USPHS, and the Henry J. Kaiser Family Foundation.

intermittent treatment regimens designed to minimize exposure to the drugs in question, all efforts to do so have paradoxically been associated with an increase in serious adverse events in the patients randomized to intermittent therapy, suggesting that some “non-AIDS-associated” serious adverse events such as heart attack and stroke may be linked to HIV replication. Thus, unless contraindicated for reasons of toxicity, patients started on cART should remain on cART.

At present, the U.S. Department of Health and Human Services Guidelines panel recommends that everyone with HIV infection be treated with cART. The evidence for this is strongest for patients with CD4+ T cell counts <350/μL. Clinical trials are underway to more carefully determine the benefit of initiating therapy in patients with CD4+ T cell counts ≥350/μL. In addition, one may wish to administer a 6-week course of therapy to uninfected individuals immediately following a high-risk exposure to HIV. The combination of tenofovir and emtricitabine is also indicated for pre-exposure prophylaxis in individuals at high risk of HIV infection. For patients diagnosed with an opportunistic infection and HIV infection at the same time, one may consider a 2- to 4-week delay in the initiation of antiretroviral therapy during which time treatment is focused on the opportunistic infection. This delay may decrease the severity of any subsequent immune reconstitution inflammatory syndrome by lowering the antigenic burden of the opportunistic infection. For patients with advanced HIV infection (CD4+ <50 cells/μL), however, cART should be initiated as soon as possible.

Once the decision has been made to initiate therapy, the health care provider must decide which drugs to use as the first regimen. The decision regarding choice of drugs not only will affect the immediate response to therapy but also will have implications regarding options for future therapeutic regimens. The initial regimen is usually the most effective insofar as the virus has yet to develop significant resistance. HIV is capable of rapidly developing resistance to any single agent, and therapy must be given as a multidrug combination. Given that patients can be infected with viruses that harbor drug resistance mutations, it is recommended that a viral genotype be done prior to the initiation of therapy to optimize the selection of antiretroviral agents. The combination regimens currently recommended for initial therapy in any treatment-naïve patient are listed in **Table 226-24**. Other regimens containing abacavir and rilpivirine may be appropriate for patients with HIV RNA levels <100,000 copies/mL. It is currently unclear whether treatment-naïve individuals with

TABLE 226-24 INITIAL COMBINATION REGIMENS FOR ANY TREATMENT-NAÏVE PATIENT REGARDLESS OF HIV RNA LEVEL OR CD4 COUNT

- I. Non-Nucleoside Reverse Transcriptase Inhibitor Based:
Efavirenz + tenofovir + emtricitabine
- II. Protease Inhibitor Based:
Atazanavir/ritonavir + tenofovir + emtricitabine
Darunavir/ritonavir + tenofovir + emtricitabine
- III. Integrase Inhibitor Based:
Dolutegravir + tenofovir + emtricitabine
Raltegravir + tenofovir + emtricitabine

Source: *Guidelines for the Use of Antiretroviral Agents in HIV-Infected Adults and Adolescents*, USPHS.

<50 copies/mL of HIV RNA benefit from cART. Following the initiation of therapy one should expect a rapid, at least 1-log (tenfold) reduction in plasma HIV RNA levels within 1–2 months and then a slower decline in plasma HIV RNA levels to <50 copies/mL within 6 months. During this same time there should be a rise in the CD4+ T cell count of 100–150/μL that is also particularly brisk during the first month of therapy. Subsequently, one should anticipate a CD4+ T cell count increase of 50–100 cells/year until numbers approach normal. Many clinicians feel that failure to achieve these endpoints is an indication for a change in therapy. Other reasons for a change in therapy include a persistently declining CD4+ T cell count, a consistent increase in HIV RNA levels to >200 copies/mL, clinical deterioration, or drug toxicity (**Table 226-25**). As in the case of initiating therapy, changing therapy may have a lasting impact on future therapeutic options. When changing therapy because of treatment failure (clinical progression or worsening laboratory parameters), it is important to attempt to provide a regimen with at least two new active drugs. This decision can be guided by resistance testing (see below). In the patient in whom a change is made for reasons of drug toxicity, a simple replacement of one drug is reasonable. It should be stressed that in attempting to sort out a drug toxicity it may be advisable to hold all therapy for a period of time to distinguish between drug toxicity and disease progression. Drug toxicity will usually begin to show signs of reversal within 1–2 weeks. Prior to changing a treatment regimen because of drug failure, it is important to ensure that the patient has been adherent to the prescribed regimen. As in the case of initial therapy, the simpler the new therapeutic regimen, the easier it is for the patient to be compliant. Plasma HIV RNA levels should be monitored every 3–6 months during therapy and more frequently if one is contemplating a change in regimen due to an increase in viral load or immediately following a change in regimen.

In order to determine an optimal therapeutic regimen for initial therapy or for a patient on a failing regimen, one may attempt to measure antiretroviral drug susceptibility through genotyping or phenotyping of HIV quasiespecies and to determine adequacy of dosing through measurement of drug levels. Genotyping may be done through cDNA sequencing. Phenotypic assays typically measure the

TABLE 226-25 INDICATIONS FOR CHANGING ANTIRETROVIRAL THERAPY IN PATIENTS WITH HIV INFECTION^a

- Less than a 1-log drop in plasma HIV RNA by 4 weeks following the initiation of therapy
- A reproducible significant increase (defined as threefold or greater) from the nadir of plasma HIV RNA level not attributable to intercurrent infection, vaccination, or test methodology
- Persistently declining CD4+ T cell numbers
- Clinical deterioration
- Side effects

^aGenerally speaking, a change should involve the initiation of at least two drugs felt to be effective in the given patient. The exception to this is when change is being made to manage toxicity, in which case a single substitution is reasonable.

Source: *Guidelines for the Use of Antiretroviral Agents in HIV-Infected Adults and Adolescents*, USPHS.

enzymatic activity of viral enzymes in the presence or absence of different concentrations of different drugs and have also been used to determine co-receptor tropism. These assays will generally detect quasispecies present at a frequency of $\geq 10\%$. NextGen sequencing may allow detection of quasispecies at frequencies down to 1%. It is generally recommended that resistance testing be used in selecting initial therapy in settings where the risk of transmission of resistant virus is high (such as the United States and Europe) and in determining new regimens for patients experiencing virologic failure while on therapy. Resistance testing may be of particular value in distinguishing drug-resistant virus from poor patient compliance. Due to the rapid rate at which drug-resistant viruses revert to wild-type, it is recommended that resistance testing performed in the setting of drug failure be carried out while the patient is still on the failing regimen. Measurement of plasma drug levels can also be used to tailor an individual treatment. The inhibitory quotient, defined as the trough blood level/ IC_{50} of the patient's virus, is used by some to determine the adequacy of dosing of a given treatment regimen. Despite the best of efforts there will still be patients with ongoing high levels of HIV replication while receiving the best available therapy. These patients will receive benefit from remaining on antiretroviral therapy even though it is not fully suppressive.

In addition to the licensed medications discussed above, a large number of experimental agents are being evaluated as possible therapies for HIV infection. Therapeutic strategies are being developed to interfere with virtually every step of the replication cycle of the virus (Fig. 226-3). In addition, as more is discovered about the role of the immune system in controlling viral replication, additional strategies, generically referred to as "immune-based therapies," are being developed as a complement to antiviral therapy. Among the antiviral agents in early clinical trials are additional nucleoside and nucleotide analogues, protease inhibitors, fusion inhibitors, receptor and co-receptor antagonists, and integrase inhibitors as well as new antiviral strategies including antisense nucleic acids and maturation inhibitors. Among the immune-based therapies being evaluated are IFN- α , bone marrow transplantation, adoptive transfer of lymphocytes genetically modified to resist infection or enhance HIV-specific immunity, active immunotherapy with inactivated HIV or its components, IL-7, and IL-15.

HIV AND THE HEALTH CARE WORKER

Health care workers, especially those who deal with large numbers of HIV-infected patients, have a small but definite risk of becoming infected with HIV as a result of professional activities (see "Occupational Transmission of HIV: Health Care Workers, Laboratory Workers, and the Health Care Setting," above). The first case of HIV transmission from a patient to health care worker was reported in 1984. Occupational transmission of HIV has been reported in most countries; as noted above, the global number of HIV infections among health care workers attributable to punctures/cuts has been estimated to be 1000 cases (range, 200–5000) per year.

In the United States 57 health care workers for whom case investigations were completed as of 2010 had documented seroconversions to HIV following occupational exposures. The routes of exposure resulting in infection were as follows: 48 percutaneous (puncture/cut injury); 5 mucocutaneous (mucous membrane and/or skin); 2 both percutaneous and mucocutaneous; and 2 of unknown route. Of the 57 health care personnel, 49 were exposed to HIV-infected blood; 3 to concentrated virus in a laboratory; 1 to visibly bloody fluid; and 4 to an unspecified fluid. The individuals with documented seroconversions included 19 laboratory workers (16 of whom were clinical laboratory workers), 24 nurses, 6 physicians, 2 surgical technicians, 1 dialysis technician, 1 respiratory therapist, 1 health aide, 1 embalmer/morgue technician, and 2 housekeeper/maintenance workers. In addition, more than 140 possible cases of occupationally acquired HIV infection have been reported among health care personnel in the United States. The number of these workers who actually acquired their infection through occupational exposures is not known. Taken

together, data from several large studies suggest that the risk of HIV infection following a percutaneous exposure to HIV-contaminated blood is $\sim 0.3\%$, and after a mucous membrane exposure, $\sim 0.09\%$. Although episodes of HIV transmission after nonintact skin exposure have been documented, the average risk for transmission by this route has not been precisely quantified but is estimated to be less than the risk for mucous membrane exposures. The risk for transmission after exposure to fluids or tissues other than HIV-infected blood also has not been quantified but is probably considerably lower than for blood exposures. A seroprevalence survey of 3420 orthopedic surgeons, 75% of whom practiced in an area with a relatively high prevalence of HIV infection and 39% of whom reported percutaneous exposure to patient blood, usually through an accident involving a suture needle, failed to reveal any cases of possible occupational infection, suggesting that the risk of infection with a suture needle may be considerably less than that with a blood-drawing (hollow-bore) needle.

Most cases of health care worker seroconversion occur as a result of needle-stick injuries. When one considers the circumstances that result in needle-stick injuries, it is immediately obvious that adhering to the standard guidelines for dealing with sharp objects would result in a significant decrease in this type of accident. In one study, 27% of needle-stick injuries resulted from improper disposal of the needle (over half of these were due to recapping the needle), 23% occurred during attempts to start an IV line, 22% occurred during blood drawing, 16% were associated with an IM or SC injection, and 12% were associated with giving an IV infusion.

Clinicians should consider potential occupational exposures to HIV as urgent medical concerns to ensure timely postexposure management and possible administration of postexposure antiretroviral prophylaxis (PEP). Recommendations regarding PEP must take into account that a variety of circumstances determine the risk of transmission of HIV following occupational exposure. In this regard, several factors have been associated with an increased risk for occupational transmission of HIV infection, including deep injury, the presence of visible blood on the instrument causing the exposure, injury with a device that had been placed in the vein or artery of the source patient, terminal illness in the source patient, and lack of postexposure cART in the exposed health care worker. Other important considerations when considering PEP in the health care worker include known or suspected pregnancy or breast-feeding, the possibility of exposure to drug-resistant virus, and toxicities of PEP regimens. Regardless of the decision to use PEP, the wound should be cleansed immediately and antiseptic applied. If a decision is made to offer PEP, U.S. Public Health Service guidelines recommend (1) a combination of two nucleoside analogue reverse transcriptase inhibitors given for 4 weeks for less severe exposures, or (2) a combination of two nucleoside analogue reverse transcriptase inhibitors plus a third drug given for 4 weeks for more severe exposures. Most clinicians administer the latter regimen in all cases in which a decision is made to treat. Detailed guidelines are available from the *Updated U.S. Public Health Service Guidelines for the Management of Occupational Exposures to HIV and Recommendations for Postexposure Prophylaxis* (CDC, 2005). The report emphasizes the importance of adherence to PEP when it is indicated, follow-up of exposed workers to improve PEP adherence, monitoring for adverse events (including seroconversion), and expert consultation in the management of exposures.

For consultation on the treatment of occupational exposures to HIV and other bloodborne pathogens, the clinician managing the exposed patient can call the National Clinicians' Post-Exposure Prophylaxis Hotline (PEpline) at 888-448-4911. This service is available 24 hours a day at no charge. (Additional information on the Internet is available at www.nccc.ucsf.edu.) PEpline support may be especially useful in challenging situations, such as when drug-resistant HIV strains are suspected or the health care worker is pregnant.

Health care workers can minimize their risk of occupational HIV infection by following the CDC guidelines of July 1991, which include adherence to universal precautions, refraining from direct patient care if one has exudative lesions or weeping dermatitis, and disinfecting and sterilizing reusable devices employed in invasive procedures.

1284 The premise of universal precautions is that every specimen should be handled as if it came from someone infected with a bloodborne pathogen. All samples should be double-bagged, gloves should be worn when drawing blood, and spills should be immediately disinfected with bleach.

In attempting to put this small but definite risk to the health care worker in perspective, it is important to point out that ~200 health care workers die each year as a result of occupationally acquired hepatitis B infection. The tragedy in this instance is that these infections and deaths due to HBV could be greatly decreased by more extended use of the HBV vaccine. The risk of HBV infection following a needle-stick injury from a hepatitis antigen-positive patient is much higher than the risk of HIV infection (see “Transmission,” above). There are multiple examples of needle-stick injuries where the patient was positive for both HBV and HIV and the health care worker became infected only with HBV. For these reasons, it is advisable, given the high prevalence of HBV infection in HIV-infected individuals, that all health care workers dealing with HIV-infected patients be immunized with the HBV vaccine.

TB is another infection common to HIV-infected patients that can be transmitted to the health care worker. For this reason, all health care workers should know their PPD status, have it checked yearly, and receive 6 months of isoniazid treatment if their skin test converts to positive. In addition, all patients in whom a diagnosis of TB is being entertained should be placed immediately in respiratory isolation, pending results of the diagnostic evaluation. The emergence of drug-resistant organisms, including the extensively drug-resistant TB strains that have been identified in Africa, has made TB an increasing problem for health care workers. This is particularly true for the health care worker with preexisting HIV infection.

One of the most charged issues ever to come between health care workers and patients is that of transmission of infection from HIV-infected health care workers to their patients. This is discussed in “Occupational Transmission of HIV: Health Care Workers, Laboratory Workers, and the Health Care Setting,” above. Theoretically, the same universal precautions that are used to protect the health care worker from the HIV-infected patient will also protect the patient from the HIV-infected health care worker.

▲ PREVENTIVE VACCINE AGAINST HIV INFECTION

Given that human behavior, especially human sexual behavior, is extremely difficult to change, a critical modality for preventing the spread of HIV infection is the development of a safe and effective vaccine. Historically, vaccines have provided a safe, cost-effective, and efficient means of preventing illness, disability, and death from infectious diseases. Successful vaccines for the most part are predicated on the assumptions that the body can mount an adequate immune response to the microbe or virus in question during natural infection and that the vaccine will mimic the natural response to infection. Even with serious diseases, such as smallpox, poliomyelitis, measles, and influenza among others, the body in the vast majority of cases clears the infectious agent and provides protection, which is usually life-long against future exposure. Unfortunately, this is not the case with HIV infection since the natural immune response to HIV infection is unable to clear the virus from the body and cases of superinfection are not uncommon. Some of the factors that contribute to the problematic nature of development of a preventive HIV vaccine are the high mutability of the virus, the fact that the infection can be transmitted by cell-free or cell-associated virus, the fact that the HIV provirus integrates itself into the genome of the target cell and may remain in a latent form unexposed to the immune system, the likely need for the development of effective mucosal immunity, and the fact that it has been difficult to establish the precise correlates of protective immunity to HIV infection. A fraction of a percent of HIV-infected individuals are “elite controllers” in that they maintain extremely low and even undetectable levels of viremia in the absence of cART, and a number of individuals have been exposed to HIV multiple times but remain uninfected; these facts suggest that there are elements

of host defense or an HIV-specific immune response that have the potential to be protective. Early attempts to develop a vaccine with the envelope protein gp120 aimed at inducing neutralizing antibodies in humans were unsuccessful in that the elicited antisera failed to neutralize primary isolates of HIV cultured and tested in fresh peripheral blood mononuclear cells. In this regard, two phase 3 trials were undertaken in the United States and Thailand using soluble gp120, and the vaccines failed to protect human volunteers from HIV infection. In addition, two separate vaccine trials aimed at eliciting CD8+ T cell responses to prevent infection and, if unsuccessful in preventing infection to control postinfection viremia, also failed at both goals. Recently, a vaccine using a poxvirus vector prime expressing various viral proteins followed by an envelope protein boost was tested in a 16,000-person clinical trial (RV144) conducted in Thailand among predominantly low-prevalence heterosexuals. The vaccine provided the first positive, albeit very modest, signal ever reported in an HIV vaccine trial, showing 31% protection against acquisition of infection. Such a result is certainly not sufficient justification for clinical use of the vaccine, but it served as an important first step in the direction of the development of a safe and effective vaccine against HIV infection. Follow-up studies of RV144 indicate that nonneutralizing or weakly neutralizing antibody responses against certain constant epitopes in the otherwise highly variable V1-V2 region of the HIV envelope may be associated with the modest degree of protection observed in that clinical trial. Additional studies are planned in attempts to improve on the results of RV144 by a variety of approaches, including increasing the number of vaccine boosts with envelope protein.

An area of HIV vaccine research that is currently being actively pursued is the attempt to induce broadly neutralizing antibodies by developing as immunogens for vaccination certain epitopes on the HIV envelope that are the targets of naturally occurring broadly neutralizing antibodies during HIV infection. It is curious that only about 20% of HIV-infected individuals develop broadly neutralizing antibodies in response to natural infection and they do so only after 2 to 3 years of ongoing infection. By the time these antibodies appear, they can neutralize a broad range of primary HIV isolates, but they appear to be ineffective against the autologous virus in the infected subject. Upon close examination, these broadly neutralizing antibodies manifest a high degree of somatic mutations that were accumulated over time and are responsible for their affinity maturation and broadly neutralizing capacity. The goal of current efforts is to develop the conformationally correct HIV envelope epitopes that, when used as immunogens, would direct the immune response of an uninfected individual to the production of broadly neutralizing antibodies over a reasonable time frame by sequential immunizations. It remains to be seen whether this approach will be feasible.

PREVENTION

Education, counseling, and behavior modification are the cornerstones of any HIV prevention strategy. A major problem in the United States and elsewhere is that many infections are passed on by those who do not know that they are infected. Of the ~1.1 million persons in the United States who are HIV-infected, it is estimated that ~16–18% do not know their HIV status and approximately 49% of all new infections are transmitted by those people who are not aware that they are infected. In this regard, the CDC has recommended that HIV testing become part of routine medical care and that all individuals between the ages of 13 and 64 years be tested at least one time. These individuals should be informed of the testing and be tested without the need for written informed consent. Each individual could “opt out” of testing, but testing would otherwise be routinely administered. Individuals who are practicing high-risk behavior should be tested more often. In addition to identifying individuals who might benefit from cART, information gathered from such an approach should serve as the basis for behavior-modification programs, both for infected individuals who may be unaware of their HIV status and who could infect others and for uninfected individuals practicing high-risk behavior. The practice of “safer sex” is the most effective way for sexually active uninfected

individuals to avoid contracting HIV infection and for infected individuals to avoid spreading infection. Abstinence from sexual relations is the only absolute way to prevent sexual transmission of HIV infection. However, for many individuals this may not be feasible, and there are a number of relatively safe practices that can markedly decrease the chances of transmission of HIV infection. Partners engaged in monogamous sexual relationships who wish to be assured of safety should both be tested for HIV antibody. If both are negative, it must be understood that any divergence from monogamy puts both partners at risk; open discussion of the importance of honesty in such relationships should be encouraged. When the HIV status of either partner is not known, or when one partner is positive, there are a number of options. Use of condoms can markedly decrease the chance of HIV transmission. It should be remembered that condoms are not 100% effective in preventing transmission of HIV infection, and there is a ~10% failure rate of condoms used for contraceptive purposes. Most condom failures result from breakage or improper usage, such as not wearing the condom for the entire period of intercourse. Latex condoms are preferable, since virus has been shown to leak through natural skin condoms. Petroleum-based gels should never be used for lubrication of the condom, since they increase the likelihood of condom rupture. Some men who have sex with men practice fellatio as a “minimal risk” activity compared with anal intercourse. It should be emphasized that receptive fellatio is definitely not safe sex, and although the incidence of transmission via fellatio is considerably less than that of rectal or vaginal intercourse, there has been documentation of transmission of HIV where receptive fellatio was the only sexual act performed (see “Transmission,” above). Topical microbicides composed of gels containing antiretroviral drugs have been shown to be efficacious in preventing acquisition of HIV infection in women engaging in vaginal intercourse. However, there has been a considerable degree of variability in efficacy related to the variable adherence of participants to the use of the intervention. In general, it is felt that microbicides can

be quite efficacious; however, adherence is a major stumbling block to their broad effectiveness. Pre-exposure prophylaxis (PreP) using oral antiretroviral drugs on a daily basis in uninfected men who have sex with men and transgender women has been shown to be efficacious in preventing acquisition of HIV infection. The degree of efficacy can be very high (>90%) if subjects adhere strictly to the regimen. However, adherence has proven to be a problem in maximizing the overall effectiveness of this approach.

Adult male circumcision has been shown to result in a 50% to 65% reduction in HIV acquisition in the circumcised subject. Clearly, this approach has considerable potential as a preventive strategy for HIV infection and is currently being pursued, particularly in developing nations, as a component of HIV prevention. The most effective way to prevent transmission of HIV infection among IDUs is to stop the use of injectable drugs. Unfortunately, that is extremely difficult to accomplish unless the individual enters a treatment program. For those who will not or cannot participate in a drug treatment program and who will continue to inject drugs, the avoidance of sharing of needles and other paraphernalia (“works”) is the next best way to avoid transmission of infection. However, the cultural and social factors that contribute to the sharing of paraphernalia are complex and difficult to overcome. In addition, needles and syringes may be in short supply. Under these circumstances, paraphernalia should be cleaned after each usage with a virucidal solution, such as undiluted sodium hypochlorite (household bleach). Programs that provide sterile needles to addicts in exchange for used needles have resulted in a marked decrease in HIV transmission without increasing the use of injection drugs. It is important for IDUs to be tested for HIV infection and counseled to avoid transmission to their sexual partners. Oral PreP also is effective in preventing acquisition of HIV infections among IDUs. Prevention of transmission through blood or blood products and prevention of mother-to-child transmission are discussed in “Transmission,” above.

SECTION 15 INFECTIONS DUE TO RNA VIRUSES

227 Viral Gastroenteritis

Umesh D. Parashar, Roger I. Glass

Acute infectious gastroenteritis is a common illness that affects persons of all ages worldwide. It is a leading cause of mortality among children in developing countries, accounting for an estimated 0.7 million deaths each year, and is responsible for up to 10–12% of all hospitalizations among children in industrialized countries, including the United States. Elderly persons, especially those with debilitating health conditions, also are at risk of severe complications and death from acute gastroenteritis. Among healthy young adults, acute gastroenteritis is rarely fatal but incurs substantial medical and social costs, including those of time lost from work.

Several enteric viruses have been recognized as important etiologic agents of acute infectious gastroenteritis (Table 227-1, Fig. 227-1). Although most viral gastroenteritis is caused by RNA viruses, the DNA viruses that are occasionally involved (e.g., adenovirus types 40 and 41) are included in this chapter. Illness caused by these viruses is characterized by the acute onset of vomiting and/or diarrhea, which may be accompanied by fever, nausea, abdominal cramps, anorexia, and malaise. As shown in Table 227-2, several features can help distinguish gastroenteritis caused by viruses from that caused by bacterial agents. However, the distinction based on clinical and epidemiologic parameters alone is often difficult, and laboratory tests are required to confirm the diagnosis.

HUMAN CALICIVIRUSES

Etiologic Agent The Norwalk virus is the prototype strain of a group of small (27–40 nm), nonenveloped, round, icosahedral viruses with relatively amorphous surface features on visualization by electron microscopy. These viruses have been difficult to classify because they have not been adapted to growth in cell culture and no animal models are available. Molecular cloning and characterization have demonstrated that the viruses have a single, positive-strand RNA genome ~7.5 kb in length and possess a single virion-associated protein—similar to that of typical caliciviruses—with a molecular mass of 60 kDa. On the basis of these molecular characteristics, these viruses are presently classified in two genera belonging to the family Calciviridae: the *noroviruses* and the *sapoviruses* (previously called Norwalk-like viruses and Sapporo-like viruses, respectively).



Epidemiology Infections with the Norwalk and related human caliciviruses are common worldwide, and most adults have antibodies to these viruses. Antibody is acquired at an earlier age in developing countries—a pattern consistent with the presumed fecal-oral mode of transmission. Infections occur year-round, although, in temperate climates, a distinct increase has been noted in cold-weather months. Noroviruses may be the most common infectious agents of mild gastroenteritis in the community and affect all age groups, whereas sapoviruses primarily cause gastroenteritis in children. Noroviruses also cause traveler’s diarrhea, and outbreaks have occurred among military personnel deployed to various parts of the world. The limited data available indicate that norovirus may be the second most common viral agent (after rotavirus) among young

TABLE 227-1 VIRAL CAUSES OF GASTROENTERITIS AMONG HUMANS

Virus	Family	Genome	Primary Age Group at Risk	Clinical Severity	Detection Assays
Group A rotavirus	Reoviridae	Double-strand segmented RNA	Children <5 years	+++	EM, EIA (commercial), PAGE, RT-PCR
Norovirus	Caliciviridae	Positive-sense single-strand RNA	All ages	++	EM, RT-PCR
Sapovirus	Caliciviridae	Positive-sense single-strand RNA	Children <5 years	+	EM, RT-PCR
Astrovirus	Astroviridae	Positive-sense single-strand RNA	Children <5 years	+	EM, EIA, RT-PCR
Adenovirus (mainly types 40 and 41)	Adenoviridae	Double-strand DNA	Children <5 years	+/+ +	EM, EIA (commercial), PCR

Abbreviations: EIA, enzyme immunoassay; EM, electron microscopy; PAGE, polyacrylamide gel electrophoresis; PCR, polymerase chain reaction; RT-PCR, reverse-transcription PCR.

children and the most common agent among older children and adults. In the United States, with the decline in severe rotavirus disease following implementation of rotavirus vaccines, norovirus has become the leading cause of medically attended gastroenteritis in young children. Noroviruses are also recognized as the major cause of epidemics of gastroenteritis worldwide. In the United States, >90% of outbreaks of nonbacterial gastroenteritis are caused by noroviruses.

Virus is transmitted predominantly by the fecal-oral route but is also present in vomitus. Because an inoculum with very few viruses can be infectious, transmission can occur by aerosolization, by contact with contaminated fomites, and by person-to-person contact. Viral shedding and infectivity are greatest during the acute illness, but challenge studies with Norwalk virus in volunteers indicate that viral antigen may be shed by asymptotically infected persons and also by symptomatic persons before the onset of symptoms and for several weeks after the resolution of illness. Viral shedding can be prolonged in immunocompromised individuals.

Pathogenesis The exact sites and cellular receptors for attachment of viral particles have not been determined. Data suggest that carbohydrates that are similar to human histo-blood group antigens and are present on the gastroduodenal epithelium of individuals with the secretor phenotype may serve as ligands for the attachment of Norwalk virus. Additional studies must more fully elucidate norovirus-carbohydrate interactions, including potential strain-specific variations. After the infection of volunteers, reversible lesions are noted in the upper jejunum, with broadening and blunting of the villi, shortening of the microvilli, vacuolization of the lining epithelium, crypt hyperplasia, and infiltration of the lamina propria by polymorphonuclear neutrophils and lymphocytes. The lesions persist for at least 4 days after the resolution of symptoms and are associated with malabsorption of carbohydrates and fats and a decreased level of brush-border enzymes.

Adenylate cyclase activity is not altered. No histopathologic changes are seen in the stomach or colon, but gastric motor function is delayed, and this alteration is believed to contribute to the nausea and vomiting that are typical of this illness.

Clinical Manifestations Gastroenteritis caused by Norwalk and related human caliciviruses has a sudden onset following an average incubation period of 24 h (range, 12–72 h). The illness generally lasts 12–60 h and is characterized by one or more of the following symptoms: nausea, vomiting, abdominal cramps, and diarrhea. Vomiting is more prevalent among children, whereas a greater proportion of adults develop diarrhea. Constitutional symptoms are common, including headache, fever, chills, and myalgias. The stools are characteristically loose and watery, without blood, mucus, or leukocytes. White cell counts are generally normal; rarely, leukocytosis with relative lymphopenia may be observed. Death is a rare outcome and usually results from severe dehydration in vulnerable persons (e.g., elderly patients with debilitating health conditions).

Immunity Approximately 50% of persons challenged with Norwalk virus become ill and acquire short-term immunity against the infecting strain. Immunity to Norwalk virus appears to correlate inversely with level of antibody; i.e., persons with higher levels of preexisting antibody to Norwalk virus are more susceptible to illness. This observation suggests that some individuals have a genetic predisposition to illness. Specific ABO, Lewis, and secretor blood group phenotypes may influence susceptibility to norovirus infection.

Diagnosis Cloning and sequencing of the genomes of Norwalk and several other human caliciviruses have allowed the development of assays based on polymerase chain reaction (PCR) for detection of virus in stool and vomitus. Virus-like particles produced by expression of capsid proteins in a recombinant baculovirus vector have been

used to develop enzyme immunoassays (EIAs) for detection of virus in stool or a serologic response to a specific viral antigen. These newer diagnostic techniques are considerably more sensitive than previous detection methods, such as electron microscopy, immune electron microscopy, and EIAs based on reagents derived from humans. However, no currently available single assay can detect all human caliciviruses because of their great genetic and antigenic diversity. In addition, the assays are still cumbersome and are available primarily in research laboratories, although they are increasingly being adopted by public health laboratories for routine screening of fecal specimens from patients affected by outbreaks of gastroenteritis. Commercial EIA kits have limited sensitivity and usefulness in clinical practice and are of greatest

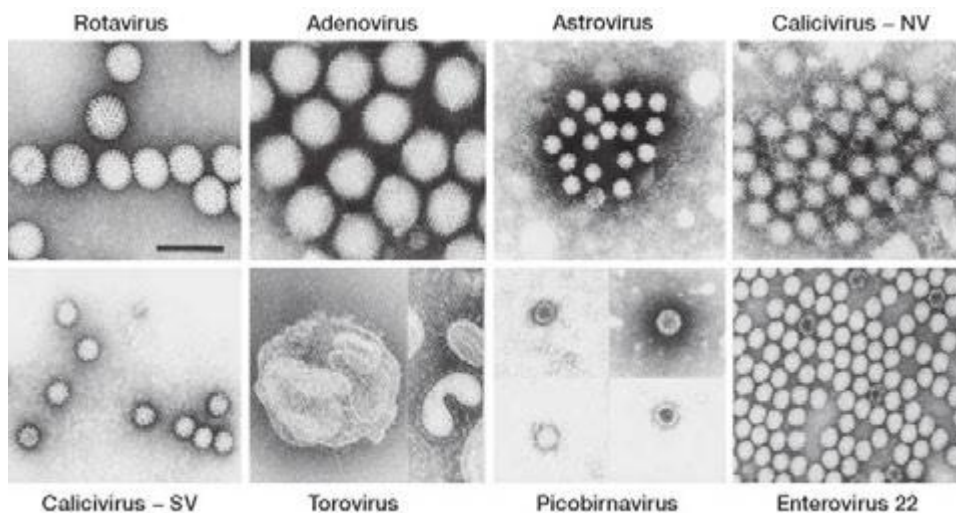


FIGURE 227-1 Viral agents of gastroenteritis. NV, norovirus; SV, sapovirus.

TABLE 227-2 CHARACTERISTICS OF GASTROENTERITIS CAUSED BY VIRAL AND BACTERIAL AGENTS

Feature	Viral Gastroenteritis	Bacterial Gastroenteritis
Setting	Incidence similar in developing and developed countries	More common in settings with poor hygiene and sanitation
Infectious dose	Low (10–100 viral particles) for most agents	High (>10 ⁵ bacteria) for <i>Escherichia coli</i> , <i>Salmonella</i> , <i>Vibrio</i> ; medium (10 ² –10 ⁵ bacteria) for <i>Campylobacter jejuni</i> ; low (10–100 bacteria) for <i>Shigella</i>
Seasonality	In temperate climates, winter seasonality for most agents; year-round occurrence in tropical areas	More common in summer or rainy months, particularly in developing countries with a high disease burden
Incubation period	1–3 days for most agents; can be shorter for norovirus	1–7 days for common agents (e.g., <i>Campylobacter</i> , <i>E. coli</i> , <i>Shigella</i> , <i>Salmonella</i>); a few hours for bacteria producing preformed toxins (e.g., <i>Staphylococcus aureus</i> , <i>Bacillus cereus</i>)
Reservoir	Primarily humans	Depending on species, human (e.g., <i>Shigella</i> , <i>Salmonella</i>), animal (e.g., <i>Campylobacter</i> , <i>Salmonella</i> , <i>E. coli</i>), and water (e.g., <i>Vibrio</i>) reservoirs exist
Fever	Common with rotavirus and norovirus; uncommon with other agents	Common with agents causing inflammatory diarrhea (e.g., <i>Salmonella</i> , <i>Shigella</i>)
Vomiting	Prominent and can be the only presenting feature, especially in children	Common with bacteria producing preformed toxins; less prominent in diarrhea due to other agents
Diarrhea	Common; nonbloody in almost all cases	Prominent and occasionally bloody with agents causing inflammatory diarrhea
Duration	1–3 days for norovirus and sapovirus; 2–8 days for other viruses	1–2 days for bacteria producing preformed toxins; 2–8 days for most other bacteria
Diagnosis	This is often a diagnosis of exclusion in clinical practice. Commercial enzyme immunoassays are available for detection of rotavirus and adenovirus, but identification of other agents is limited to research and public health laboratories.	Fecal examination for leukocytes and blood is helpful in differential diagnosis. Culture of stool specimens, sometimes on special media, can identify several pathogens. Molecular techniques are useful epidemiologic tools but are not routinely used in most laboratories.
Treatment	Supportive therapy to maintain adequate hydration and nutrition should be given. Antibiotics and antimotility agents are contraindicated.	Supportive hydration therapy is adequate for most patients. Antibiotics are recommended for patients with dysentery caused by <i>Shigella</i> or diarrhea caused by <i>Vibrio cholerae</i> and for some patients with <i>Clostridium difficile</i> colitis.

utility in outbreaks, in which many specimens are tested and only a few need be positive to identify norovirus as the cause.

TREATMENT INFECTIONS WITH NORWALK AND RELATED HUMAN CALICIVIRUSES

The disease is self-limited, and oral rehydration therapy is generally adequate. If severe dehydration develops, IV fluid therapy is indicated. No specific antiviral therapy is available.

Prevention Epidemic prevention relies on situation-specific measures, such as control of contamination of food and water, exclusion of ill food handlers, and reduction of person-to-person spread through good personal hygiene and disinfection of contaminated fomites. The role of immunoprophylaxis is not clear, given the lack of long-term immunity from natural disease, but efforts to develop norovirus vaccines are ongoing. In a clinical study, a candidate virus-like particle norovirus vaccine was shown to protect against homologous viral challenge.

ROTA VIRUS

Etiologic Agent Rotaviruses are members of the family Reoviridae. The viral genome consists of 11 segments of double-strand RNA that are enclosed in a triple-layered, nonenveloped, icosahedral capsid 75 nm in diameter. Viral protein 6 (VP6), the major structural protein, is the target of commercial immunoassays and determines the group specificity of rotaviruses. There are seven major groups of rotavirus (A through G); human illness is caused primarily by group A and, to a much lesser extent, by groups B and C. Two outer-capsid proteins, VP7 (G-protein) and VP4 (P-protein), determine serotype specificity, induce neutralizing antibodies, and form the basis for binary classification of rotaviruses (G and P types). The segmented genome of rotavirus allows genetic reassortment (i.e., exchange of genome segments between viruses) during co-infection—a property that may play a role in viral evolution and that has been utilized in the development of reassortant animal-human rotavirus-based vaccines.



Epidemiology Worldwide, nearly all children are infected with rotavirus by 3–5 years of age. Neonatal infections are common but are often asymptomatic or mild, presumably because of protection by maternal antibody or breast milk. Compared with rotavirus disease in industrialized countries, disease in developing countries occurs at a younger age, is less seasonal, and is more frequently caused by uncommon rotavirus strains. Moreover, because of suboptimal access to hydration therapy, rotavirus is a leading cause of diarrheal death among children in the developing world, with the highest mortality rates among children in sub-Saharan Africa and South Asia (Fig. 227-2).

First infections after 3 months of age are likely to be symptomatic, and the incidence of disease peaks among children 4–23 months of age. Reinfections are common, but the severity of disease decreases with each repeat infection. Therefore, severe rotavirus infections are less common among older children and adults than among younger individuals. Nevertheless, rotavirus can cause illness in parents and caretakers of children with rotavirus diarrhea, immunocompromised persons, travelers, and elderly individuals and should be considered in the differential diagnosis of gastroenteritis among adults.

In tropical settings, rotavirus disease occurs year-round, with less pronounced seasonal peaks than in temperate settings, where rotavirus disease occurs predominantly during the cooler fall and winter months. Before the introduction of rotavirus vaccine in the United States, the rotavirus season each year began in the Southwest during the autumn and early winter (October through December) and migrated across the continent, peaking in the Northeast during late winter and spring (March through May). The reasons for this characteristic pattern are not clear but may be correlated with state-specific differences in birth rates, which could influence the rate of accumulation of susceptible infants after each rotavirus season. After the implementation of routine vaccination of U.S. infants against rotavirus in 2006, the characteristic prevaccine geotemporal pattern of U.S. rotavirus was dramatically altered, and these changes were accompanied by substantial declines in rotavirus detections by a national network of sentinel laboratories (Fig. 227-3). During the latest two seasons with available data (spanning 2010–2012), the number of rotavirus detections declined by 74–90% from the prevaccine baseline, and the

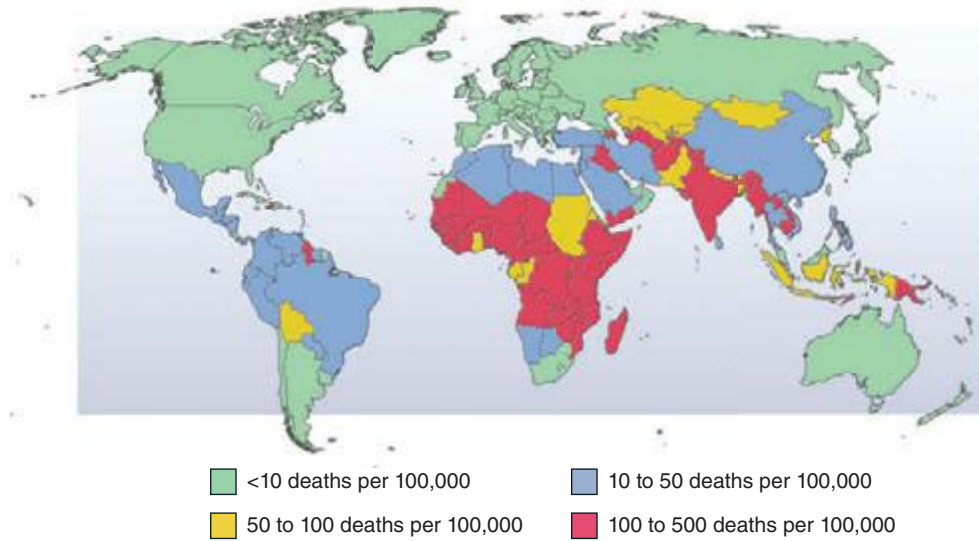


FIGURE 227-2 Rotavirus mortality rates by country, per 100,000 children <5 years of age. (Reproduced with permission from UD Parashar et al: *J Infect Dis* 200:59, 2009.)

annual proportion of rotavirus tests that were positive was below 10% in both seasons (compared with a prevaccine baseline median of 26%). A pattern of biennial increases in rotavirus activity has emerged during the five postvaccine seasons (2007–2012), but activity has remained substantially below prevaccine levels in each season.

During episodes of rotavirus-associated diarrhea, virus is shed in large quantities in stool (10^7 – 10^{12} /g). Viral shedding detectable by EIA usually subsides within 1 week but may persist for >30 days in immunocompromised individuals; it may be detected for longer periods by sensitive molecular assays, such as PCR. The virus is transmitted predominantly through the fecal-oral route. Spread through respiratory secretions, person-to-person contact, or contaminated environmental surfaces has been postulated to explain the rapid acquisition of antibody in the first 3 years of life, regardless of sanitary conditions.

At least 10 different G serotypes of group A rotavirus have been identified in humans, but only 5 types (G1 through G4 and G9) are

border enzymes characteristic of differentiated cells are reduced, and this change leads to the accumulation of unmetabolized disaccharides and consequent osmotic diarrhea. Studies in mice indicate that a nonstructural rotavirus protein, NSP4, functions as an enterotoxin and contributes to secretory diarrhea by altering epithelial cell function and permeability. In addition, rotavirus may evoke fluid secretion through activation of the enteric nervous system in the intestinal wall. Data indicate that rotavirus antigenemia and viremia are common among children with acute rotavirus infection, although the antigen and RNA levels in serum are substantially lower than those in stool.

Clinical Manifestations The clinical spectrum of rotavirus infection ranges from subclinical infection to severe gastroenteritis leading to life-threatening dehydration. After an incubation period of 1–3 days, the illness has an abrupt onset, with vomiting frequently preceding the onset of diarrhea. Up to one-third of patients may have a temperature of >39°C. The stools are characteristically loose and watery and only infrequently contain red or white cells. Gastrointestinal symptoms generally resolve in 3–7 days.

Respiratory and neurologic features in children with rotavirus infection have been reported, but causal associations have not been proven. Moreover, rotavirus infection has been associated with a variety of other clinical conditions (e.g., sudden infant death syndrome, necrotizing enterocolitis, intussusception, Kawasaki's disease, and type 1 diabetes), but no causal relationship has been confirmed with any of these syndromes.

Rotavirus does not appear to be a major opportunistic pathogen in children with HIV infection. In severely immunodeficient children, rotavirus can cause protracted diarrhea with prolonged viral excretion and, in rare instances, can disseminate systemically. Persons who are immunosuppressed for bone marrow transplantation also are at risk for severe or even fatal rotavirus disease.

Immunity Protection against rotavirus disease is correlated with the presence of virus-specific secretory IgA antibodies in the intestine and, to some extent, the serum. Because

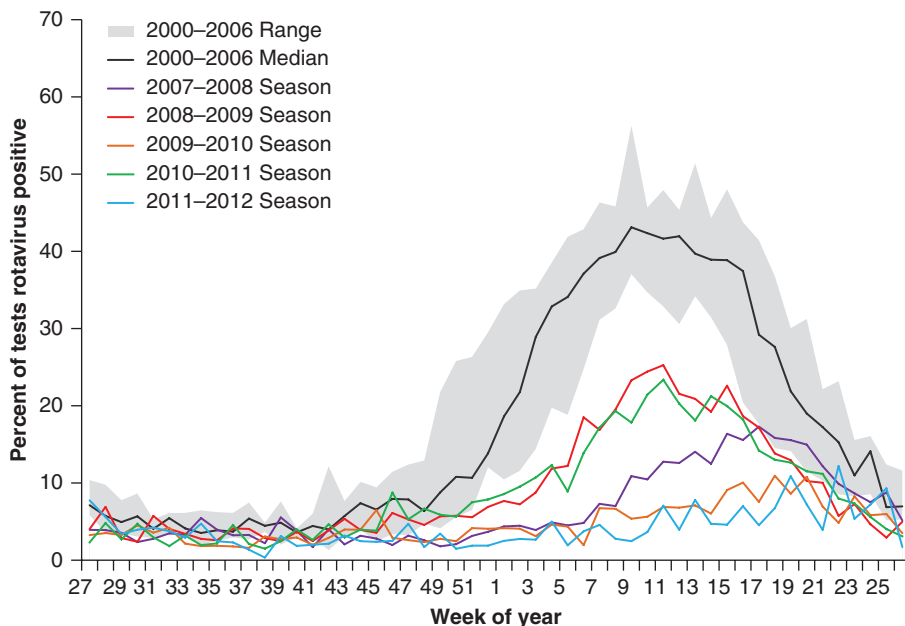


FIGURE 227-3 Percentage of rotavirus tests with positive results, by week of year, July–June, 2000–2012. The maximal or minimal percentage of rotavirus-positive tests for 2000–2006 may have occurred during any of the six baseline seasons. Data are from the National Respiratory and Enteric Virus Surveillance System. (Adapted from Centers for Disease Control and Prevention, 2012.)

virus-specific IgA production at the intestinal surface is short lived, complete protection against disease is only temporary. However, each infection and subsequent reinfection confers progressively greater immunity; thus severe disease is most common among young children with first or second infections. Immunologic memory is believed to be important in the attenuation of disease severity upon reinfection.

Diagnosis Illness caused by rotavirus is difficult to distinguish clinically from that caused by other enteric viruses. Because large quantities of virus are shed in feces, the diagnosis can usually be confirmed by a wide variety of commercially available EIAs or by techniques for detecting viral RNA, such as gel electrophoresis, probe hybridization, or PCR.

TREATMENT ROTAVIRUS INFECTIONS

Rotavirus gastroenteritis can lead to severe dehydration. Thus appropriate treatment should be instituted early. Standard oral rehydration therapy is successful for most children who can take fluids by mouth, but IV fluid replacement may be required for patients who are severely dehydrated or are unable to tolerate oral therapy because of frequent vomiting. The therapeutic roles of probiotics, bismuth subsalicylate, enkephalinase inhibitors, and nitazoxanide have been evaluated in clinical studies but are not clearly defined. Antibiotics and antimotility agents should be avoided. In immunocompromised children with chronic symptomatic rotavirus disease, orally administered immunoglobulins or colostrum may result in the resolution of symptoms, but the best choices regarding agents and their doses have not been well studied, and treatment decisions are often empirical.



Prevention Efforts to develop rotavirus vaccines were pursued because it was apparent—given the similar rates in less developed and industrialized nations—that improvements in hygiene and sanitation were unlikely to reduce disease incidence. The first rotavirus vaccine licensed in the United States in 1998 was withdrawn from the market within 1 year because it was linked with a low incidence of intussusception, a severe bowel obstruction.

In 2006, promising safety and efficacy results for two new rotavirus vaccines were reported from large clinical trials conducted in North America, Europe, and Latin America. Both vaccines are now recommended for routine immunization of all U.S. infants, and their use has rapidly led to a >70–80% decline in rotavirus hospitalizations and emergency department visits at hospitals across the United States. Indirect benefits from vaccination (i.e., herd immunity) have also been documented in many settings. In April 2009, the World Health Organization recommended the use of rotavirus vaccines in all countries worldwide. As of May 2013, a total of 42 countries, including 5 low-income countries in Africa and Asia, have incorporated rotavirus vaccine into their national childhood immunization programs. In Mexico and in Brazil, a decline in deaths from childhood diarrhea following introduction of rotavirus vaccines has been documented. Postmarketing surveillance has identified a low risk of intussusception in some countries; however, the benefits of vaccination exceed the risks, and no changes in vaccine administration policy have been implemented.



The different epidemiology of rotavirus disease and the greater prevalence of co-infection with other enteric pathogens, of comorbidities, and of malnutrition in developing countries may adversely affect the performance of oral rotavirus vaccines, as is the case with oral vaccines against poliomyelitis, cholera, and typhoid in these regions. Therefore, evaluation of the efficacy of rotavirus vaccines in resource-poor settings of Africa and Asia was specifically recommended, and these trials have now been completed. As anticipated, the efficacy of rotavirus vaccines was moderate (50–65%) in these settings when compared with that in industrialized countries. Nevertheless, even a moderately efficacious rotavirus vaccine would be likely to have substantial public health benefits in these areas with a high disease burden.

OTHER VIRAL AGENTS OF GASTROENTERITIS

Enteric *adenoviruses* of serotypes 40 and 41 belonging to subgroup F are 70- to 80-nm viruses with double-strand DNA that cause ~2–12% of all diarrhea episodes in young children. Unlike adenoviruses that cause respiratory illness, enteric adenoviruses are difficult to cultivate in cell lines, but they can be detected with commercially available EIAs. Adenovirus types 31 and 42–49 have been linked to diarrhea in HIV-infected and other immunocompromised persons.

Astroviruses are 28- to 30-nm viruses with a characteristic icosahedral structure and a positive-sense, single-strand RNA. At least seven serotypes have been identified, of which serotype 1 is most common. Astroviruses are primarily pediatric pathogens, causing ~2–10% of cases of mild to moderate gastroenteritis in children. The availability of simple immunoassays to detect virus in fecal specimens and of molecular methods to confirm and characterize strains will permit more comprehensive assessment of the etiologic role of these agents.

Toroviruses are 100- to 140-nm, enveloped, positive-strand RNA viruses that are recognized as causes of gastroenteritis in horses (Berne virus) and cattle (Breda virus). Their role as a cause of diarrhea in humans is still unclear, but studies from Canada have demonstrated associations between torovirus excretion and both nosocomial gastroenteritis and necrotizing enterocolitis in neonates. These associations require further evaluation.

Picobirnaviruses are small, bisegmented, double-strand RNA viruses that cause gastroenteritis in a variety of animals. Their role as primary causes of gastroenteritis in humans remains unclear, but several studies have found an association between picobirnaviruses and gastroenteritis in HIV-infected adults.

Several other viruses (e.g., enteroviruses, reoviruses, pestiviruses, and parvovirus B) have been identified in the feces of patients with diarrhea, but their etiologic role in gastroenteritis has not been proven. Diarrhea has also been noted as a manifestation of infection with recently recognized viruses that primarily cause severe respiratory illness: the severe acute respiratory syndrome–associated coronavirus (SARS-CoV), influenza A/H5N1 virus, and the current pandemic strain of influenza A/H1N1 virus.

228

Enterovirus, Parechovirus, and Reovirus Infections

Jeffrey I. Cohen

ENTEROVIRUSES

CLASSIFICATION AND CHARACTERIZATION

Enteroviruses, members of the family Picornaviridae, are so designated because of their ability to multiply in the gastrointestinal tract. Despite their name, these viruses are not a prominent cause of gastroenteritis. Enteroviruses encompass more than 100 human serotypes: 3 serotypes of poliovirus, 21 serotypes of coxsackievirus A, 6 serotypes of coxsackievirus B, 28 serotypes of echovirus, enteroviruses 68–71, and multiple new enteroviruses (beginning with enterovirus 73) that have been identified by molecular techniques. Human enteroviruses have been reclassified into four species designated A–D. Echoviruses 22 and 23 have been reclassified as parechoviruses 1 and 2 on the basis of low nucleotide homology and differences in viral proteins. Enterovirus surveillance conducted in the United States by the Centers for Disease Control and Prevention (CDC) in 2007–2008 showed that the most common enterovirus serotype, coxsackievirus B1, was followed in frequency by echoviruses 18, 9, and 6; together, these four viruses accounted for 52% of all isolates.

Human enteroviruses contain a single-stranded RNA genome surrounded by an icosahedral capsid comprising four viral proteins. These viruses have no lipid envelope and are stable in acidic environments,

1290 including the stomach. They are susceptible to chlorine-containing cleansers but resistant to inactivation by standard disinfectants (e.g., alcohol, detergents) and can persist for days at room temperature.

PATHOGENESIS AND IMMUNITY

Much of what is known about the pathogenesis of enteroviruses has been derived from studies of poliovirus infection. After ingestion, poliovirus is thought to infect epithelial cells in the mucosa of the gastrointestinal tract and then to spread to and replicate in the submucosal lymphoid tissue of the tonsils and Peyer's patches. The virus next spreads to the regional lymph nodes, a viremic phase ensues, and the virus replicates in organs of the reticuloendothelial system. In some cases, a second episode of viremia occurs and the virus replicates further in various tissues, sometimes causing symptomatic disease.

It is uncertain whether poliovirus reaches the central nervous system (CNS) during viremia or whether it also spreads via peripheral nerves. Since viremia precedes the onset of neurologic disease in humans, it has been assumed that the virus enters the CNS via the bloodstream. The poliovirus receptor is a member of the immunoglobulin superfamily. Poliovirus infection is limited to primates, largely because their cells express the viral receptor. Studies demonstrating the poliovirus receptor in the end-plate region of muscle at the neuromuscular junction suggest that, if the virus enters the muscle during viremia, it could travel across the neuromuscular junction up the axon to the anterior horn cells. Studies of monkeys and of transgenic mice expressing the poliovirus receptor show that, after IM injection, poliovirus does not reach the spinal cord if the sciatic nerve is cut. Taken together, these findings suggest that poliovirus can spread directly from muscle to the CNS by neural pathways.

Poliovirus can usually be cultured from the blood 3–5 days after infection, before the development of neutralizing antibodies. While viral replication at secondary sites begins to slow 1 week after infection, it continues in the gastrointestinal tract. Poliovirus is shed from the oropharynx for up to 3 weeks after infection and from the gastrointestinal tract for as long as 12 weeks; hypogammaglobulinemic patients can shed poliovirus for >20 years. During replication in the gastrointestinal tract, attenuated oral poliovirus can mutate, reverting to a more neurovirulent phenotype within a few days; however, additional mutations are probably required for full neurovirulence. One patient with hypogammaglobulinemia who had been infected 12 years earlier and was receiving IV immune globulin suddenly developed quadriplegia and respiratory muscle paralysis and died; analysis showed that the virus had reverted to a more wild-type sequence.

Humoral and secretory immunity in the gastrointestinal tract is important for the control of enterovirus infections. Enteroviruses induce specific IgM, which usually persists for <6 months, and specific IgG, which persists for life. Capsid protein VP1 is the predominant target of neutralizing antibody, which generally confers lifelong protection against subsequent disease caused by the same serotype but does not prevent infection or virus shedding. Enteroviruses also induce cellular immunity whose significance is uncertain. Patients with impaired cellular immunity are not known to develop unusually severe disease when infected with enteroviruses. In contrast, the severe infections in patients with agammaglobulinemia emphasize the importance of humoral immunity in controlling enterovirus infections. Disseminated enterovirus infections have occurred in hematopoietic cell transplant recipients. IgA antibodies are instrumental in reducing poliovirus replication in and shedding from the gastrointestinal tract. Breast milk contains IgA specific for enteroviruses and can protect humans from infection.

EPIDEMIOLOGY



Enteroviruses have a worldwide distribution. More than 50% of nonpoliovirus enterovirus infections and more than 90% of poliovirus infections are subclinical. When symptoms do develop, they are usually nonspecific and occur in conjunction with fever; only a minority of infections are associated with specific clinical syndromes. The incubation period for most enterovirus infections ranges from 2 to 14 days but usually is <1 week.

Enterovirus infection is more common in socioeconomically disadvantaged areas, especially in those where conditions are crowded and in tropical areas where hygiene is poor. Infection is most common among infants and young children; serious illness develops most often during the first few days of life and in older children and adults. In developing countries, where children are infected at an early age, poliovirus infection has less often been associated with paralysis; in countries with better hygiene, older children and adults are more likely to be seronegative, become infected, and develop paralysis. Passively acquired maternal antibody reduces the risk of symptomatic infection in neonates. Young children are the most frequent shedders of enteroviruses and are usually the index cases in family outbreaks. In temperate climates, enterovirus infections occur most often in the summer and fall; no seasonal pattern is apparent in the tropics.

Most enteroviruses are transmitted primarily by the fecal-oral or oral-oral route. Patients are most infectious shortly before and after the onset of symptomatic disease, when virus is present in the stool and throat. The ingestion of virus-contaminated food or water also can cause disease. Certain enteroviruses (such as enterovirus 70, which causes acute hemorrhagic conjunctivitis) can be transmitted by direct inoculation from the fingers to the eye. Airborne transmission is important for some viruses that cause respiratory tract disease, such as coxsackievirus A21. Enteroviruses can be transmitted across the placenta from mother to fetus, causing severe disease in the newborn. The transmission of enteroviruses through blood transfusions or insect bites has not been documented. Nosocomial spread of coxsackievirus and echovirus has taken place in hospital nurseries.

CLINICAL FEATURES

Poliovirus Infection Most infections with poliovirus are asymptomatic. After an incubation period of 3–6 days, ~5% of patients present with a minor illness (abortive poliomyelitis) manifested by fever, malaise, sore throat, anorexia, myalgias, and headache. This condition usually resolves in 3 days. About 1% of patients present with aseptic meningitis (nonparalytic poliomyelitis). Examination of cerebrospinal fluid (CSF) reveals lymphocytic pleocytosis, a normal glucose level, and a normal or slightly elevated protein level; CSF polymorphonuclear leukocytes may be present early. In some patients, especially children, malaise and fever precede the onset of aseptic meningitis.

PARALYTIC POLIOMYELITIS The least common presentation is that of paralytic disease. After one or several days, signs of aseptic meningitis are followed by severe back, neck, and muscle pain and by the rapid or gradual development of motor weakness. In some cases the disease appears to be biphasic, with aseptic meningitis followed first by apparent recovery but then (1–2 days later) by the return of fever and the development of paralysis; this form is more common among children than among adults. Weakness is generally asymmetric, is proximal more than distal, and may involve the legs (most commonly); the arms; or the abdominal, thoracic, or bulbar muscles. Paralysis develops during the febrile phase of the illness and usually does not progress after defervescence. Urinary retention may also occur. Examination reveals weakness, fasciculations, decreased muscle tone, and reduced or absent reflexes in affected areas. Transient hyperreflexia sometimes precedes the loss of reflexes. Patients frequently report sensory symptoms, but objective sensory testing usually yields normal results. Bulbar paralysis may lead to dysphagia, difficulty in handling secretions, or dysphonia. Respiratory insufficiency due to aspiration, involvement of the respiratory center in the medulla, or paralysis of the phrenic or intercostal nerves may develop, and severe medullary involvement may lead to circulatory collapse. Most patients with paralysis recover some function weeks to months after infection. About two-thirds of patients have residual neurologic sequelae.

Paralytic disease is more common among older individuals, pregnant women, and persons exercising strenuously or undergoing trauma at the time of CNS symptoms. Tonsillectomy predisposes to bulbar poliomyelitis, and IM injections increase the risk of paralysis in the involved limb(s).

VACCINE-ASSOCIATED POLIOMYELITIS The risk of developing poliomyelitis after oral vaccination is estimated at 1 case per 2.5 million doses. The risk is ~2000 times higher among immunodeficient persons, especially in persons with hypo- or agammaglobulinemia. Before 1997, an average of eight cases of vaccine-associated poliomyelitis occurred—in both vaccinees and their contacts—in the United States each year. With the change in recommendations first to a sequential regimen of inactivated poliovirus vaccine (IPV) and oral poliovirus vaccine (OPV) in 1997 and then to an all-IPV regimen in 2000, the number of cases of vaccine-associated polio declined. From 1997 to 1999, six such cases were reported in the United States; no cases have been reported since 1999.

POSTPOLIO SYNDROME The *postpolio syndrome* presents as a new onset of weakness, fatigue, fasciculations, and pain with additional atrophy of the muscle group involved during the initial paralytic disease 20–40 years earlier. The syndrome is more common among women and with increasing time after acute disease. The onset is usually insidious, and weakness occasionally extends to muscles that were not involved during the initial illness. The prognosis is generally good; progression to further weakness is usually slow, with plateau periods of 1–10 years. The postpolio syndrome is thought to be due to progressive dysfunction and loss of motor neurons that compensated for the neurons lost during the original infection and not to persistent or reactivated poliovirus infection.

Other Enteroviruses An estimated 5–10 million cases of symptomatic disease due to enteroviruses other than poliovirus occur in the United States each year. Among neonates, enteroviruses are the most common cause of aseptic meningitis and nonspecific febrile illnesses. Certain clinical syndromes are more likely to be caused by certain serotypes (Table 228-1).

NONSPECIFIC FEBRILE ILLNESS (SUMMER GRIPPE) The most common clinical manifestation of enterovirus infection is a nonspecific febrile illness. After an incubation period of 3–6 days, patients present with an acute onset of fever, malaise, and headache. Occasional cases are associated with upper respiratory symptoms, and some cases include nausea and vomiting. Symptoms often last for 3–4 days, and most cases resolve in a week. While infections with other respiratory viruses occur more often from late fall to early spring, febrile illness due to enteroviruses frequently occurs in the summer and early fall.

TABLE 228-1 MANIFESTATIONS COMMONLY ASSOCIATED WITH ENTEROVIRUS SEROTYPES

Manifestation	Serotype(s) of Indicated Virus	
	Coxsackievirus	Echovirus (E) and Enterovirus (Ent)
Acute hemorrhagic conjunctivitis	A24	E70
Aseptic meningitis	A2, 4, 7, 9, 10; B1–5	E4, 6, 7, 9, 11, 13, 16, 18, 19, 30, 33; Ent70, 71
Encephalitis	A9; B1–5	E3, 4, 6, 7, 9, 11, 18, 25, 30; Ent71
Exanthem	A4, 5, 9, 10, 16; B1, 3–5	E4–7, 9, 11, 16–19, 25, 30; Ent71
Generalized disease of the newborn	B1–5	E4–7, 9, 11, 14, 16, 18, 19
Hand-foot-and-mouth disease	A5–7, 9, 10, 16; B1, 2, 5	Ent71
Herpangina	A1–10, 16, 22; B1–5	E6, 9, 11, 16, 17, 25, 30; Ent71
Myocarditis, pericarditis	A4, 9, 16; B1–5	E6, 9, 11, 22
Paralysis	A4, 7, 9; B1–5	E2–4, 6, 7, 9, 11, 18, 30; Ent70, 71
Pleurodynia	A1, 2, 4, 6, 9, 10, 16; B1–6	E1–3, 6, 7, 9, 11, 12, 14, 16, 19, 24, 25, 30
Pneumonia	A9, 16; B1–5	E6, 7, 9, 11, 12, 19, 20, 30; Ent-D68, 71

GENERALIZED DISEASE OF THE NEWBORN Most serious enterovirus infections in infants develop during the first week of life, although severe disease can occur up to 3 months of age. Neonates often present with an illness resembling bacterial sepsis, with fever, irritability, and lethargy. Laboratory abnormalities include leukocytosis with a left shift, thrombocytopenia, elevated values in liver function tests, and CSF pleocytosis. The illness can be complicated by myocarditis and hypotension, fulminant hepatitis and disseminated intravascular coagulation, meningitis or meningoencephalitis, or pneumonia. It may be difficult to distinguish neonatal enterovirus infection from bacterial sepsis, although a history of a recent virus-like illness in the mother provides a clue.

ASEPTIC MENINGITIS AND ENCEPHALITIS In children and young adults, enteroviruses are the cause of up to 90% of cases of aseptic meningitis in which an etiologic agent can be identified. Patients with aseptic meningitis typically present with an acute onset of fever, chills, headache, photophobia, and pain on eye movement. Nausea and vomiting also are common. Examination reveals meningismus without localizing neurologic signs; drowsiness or irritability may also be apparent. In some cases, a febrile illness may be reported that remits but returns several days later in conjunction with signs of meningitis. Other systemic manifestations may provide clues to an enteroviral cause, including diarrhea, myalgias, rash, pleurodynia, myocarditis, and herpangina. Examination of the CSF invariably reveals pleocytosis; the CSF cell count shows a shift from neutrophil to lymphocyte predominance within 1 day of presentation, and the total cell count does not exceed 1000/μL. The CSF glucose level is usually normal (in contrast to the low CSF glucose level in mumps), with a normal or slightly elevated protein concentration. Partially treated bacterial meningitis may be particularly difficult to exclude in some instances. Enteroviral meningitis is more common in summer and fall in temperate climates, while viral meningitis of other etiologies is more common in winter and spring. Symptoms ordinarily resolve within a week, although CSF abnormalities can persist for several weeks. Enteroviral meningitis is often more severe in adults than in children. Neurologic sequelae are rare, and most patients have an excellent prognosis.

Enteroviral encephalitis is much less common than enteroviral aseptic meningitis. Occasional highly inflammatory cases of enteroviral meningitis may be complicated by a mild form of encephalitis that is recognized on the basis of progressive lethargy, disorientation, and sometimes seizures. Less commonly, severe primary encephalitis may develop. An estimated 10–35% of cases of viral encephalitis are due to enteroviruses. Immunocompetent patients generally have a good prognosis.

Patients with hypogammaglobulinemia, agammaglobulinemia, or severe combined immunodeficiency may develop chronic meningitis or encephalitis; about half of these patients have a dermatomyositis-like syndrome, with peripheral edema, rash, and myositis. They may also have chronic hepatitis. Patients may develop neurologic disease while receiving immunoglobulin replacement therapy. Echoviruses (especially echovirus 11) are the most common pathogens in this situation.

Paralytic disease due to enteroviruses other than poliovirus occurs sporadically and is usually less severe than poliomyelitis. Most cases are due to enterovirus 70 or 71 or to coxsackievirus A7 or A9. Guillain-Barré syndrome is also associated with enterovirus infection. While earlier studies suggested a link between enteroviruses and chronic fatigue syndrome, most recent studies have not demonstrated such an association.

PLEURODYNIA (BORNHOLM DISEASE) Patients with pleurodynia present with an acute onset of fever and spasms of pleuritic chest or upper abdominal pain. Chest pain is more common in adults, and abdominal pain is more common in children. Paroxysms of severe, knifelike pain usually last 15–30 min and are associated with diaphoresis and tachypnea. Fever peaks within an hour after the onset of paroxysms and subsides when pain resolves. The involved muscles are tender to palpation, and a pleural rub may be detected. The white blood cell count and chest x-ray results are usually normal. Most cases are due to coxsackievirus B and occur during epidemics. Symptoms resolve in a few days, and recurrences are rare. Treatment includes the administration of nonsteroidal anti-inflammatory agents or the application of heat to the affected muscles.

1292 MYOCARDITIS AND PERICARDITIS Enteroviruses are estimated to cause up to one-third of cases of acute myocarditis. Coxsackievirus B and its RNA have been detected in pericardial fluid and myocardial tissue in some cases of acute myocarditis and pericarditis. Most cases of enteroviral myocarditis or pericarditis occur in newborns, adolescents, or young adults. More than two-thirds of patients are male. Patients often present with an upper respiratory tract infection that is followed by fever, chest pain, dyspnea, arrhythmias, and occasionally heart failure. A pericardial friction rub is documented in half of cases, and the electrocardiogram shows ST-segment elevations or ST- and T-wave abnormalities. Serum levels of myocardial enzymes are often elevated. Neonates commonly have severe disease, while most older children and adults recover completely. Up to 10% of cases progress to chronic dilated cardiomyopathy. Chronic constrictive pericarditis may also be a sequela.

EXANTHEMS Enterovirus infection is the leading cause of exanthems in children in the summer and fall. While exanthems are associated with many enteroviruses, certain types have been linked to specific syndromes. Echoviruses 9 and 16 have frequently been associated with exanthem and fever. Rashes may be discrete or confluent, beginning on the face and spreading to the trunk and extremities. Echovirus 9 is the most common cause of a rubelliform (discrete) rash. Unlike the rash of rubella, the enteroviral rash occurs in the summer and is not associated with lymphadenopathy. Roseola-like rashes develop after defervescence, with macules and papules on the face and trunk. The Boston exanthem, caused by echovirus 16, is a roseola-like rash. A variety of other rashes have been associated with enteroviruses, including erythema multiforme (see Fig. 25e-25) and vesicular, urticarial, petechial, or purpuric lesions. Enanthems also occur, including lesions that resemble the Koplik's spots seen with measles (see Fig. 25e-2).

HAND-FOOT-AND-MOUTH DISEASE (Fig. 228-1) After an incubation period of 4–6 days, patients with hand-foot-and-mouth disease present with fever, anorexia, and malaise; these manifestations are followed by the development of sore throat and vesicles (see Fig. 25e-23) on the

buccal mucosa and often on the tongue and then by the appearance of tender vesicular lesions on the dorsum of the hands, sometimes with involvement of the palms. The vesicles may form bullae and quickly ulcerate. About one-third of patients also have lesions on the palate, uvula, or tonsillar pillars, and one-third have a rash on the feet (including the soles) or on the buttocks. The disease is highly infectious, with attack rates of close to 100% among young children. The lesions usually resolve in 1 week. Most cases are due to coxsackievirus A16 or enterovirus 71.



An epidemic of enterovirus 71 infection in Taiwan in 1998 resulted in thousands of cases of hand-foot-and-mouth disease or herpangina (see below). Severe complications included CNS disease, myocarditis, and pulmonary hemorrhage. About 90% of those who died were children ≤ 5 years old, and death was associated with pulmonary edema or pulmonary hemorrhage. CNS disease included aseptic meningitis, flaccid paralysis (similar to that seen in poliomyelitis), and rhombencephalitis with myoclonus and tremor or ataxia. The mean age of patients with CNS complications was 2.5 years, and MRI in cases with encephalitis usually showed brain-stem lesions. Follow-up of children at 6 months showed persistent dysphagia, cranial nerve palsies, hypoventilation, limb weakness, and atrophy; at 3 years, persistent neurologic sequelae were documented, with delayed development and impaired cognitive function.

Another epidemic of enterovirus 71 infection occurred in China in 2008–2010, with nearly 500,000 infections and 126 deaths. Infections were associated with fever, rash, brain-stem encephalitis with myoclonic jerks, and limb trembling; some cases progressed to seizures and coma. Lung findings included pulmonary edema and hemorrhage; while the level of creatine kinase MB was sometimes elevated, myocardial necrosis was generally not found.

Cyclic epidemics occur every 2–3 years in other Asian countries. However, the virus circulates at lower rates in the United States, Europe, and Africa. In the United States, hand-foot-and-mouth disease is most commonly associated with coxsackievirus A16. Between November 2011 and February 2012, outbreaks of hand-foot-and-mouth disease

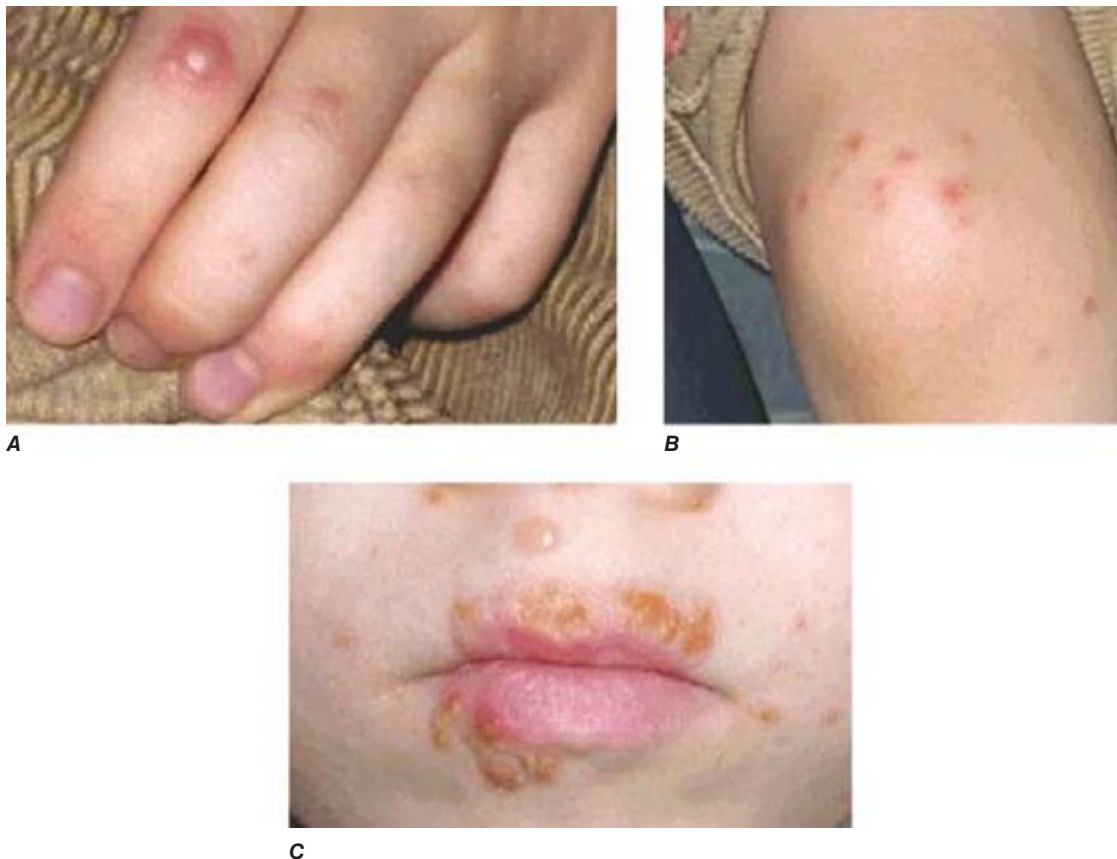


FIGURE 228-1 Vesicular eruptions of the hand (A), foot (B), and mouth (C) of a 6-year-old boy with coxsackievirus A6 infection. (Images reprinted courtesy of Centers for Disease Control and Prevention/Emerging Infectious Diseases.)

due to coxsackievirus A6 occurred in several U.S. states, and 19% of the affected persons were hospitalized.

HERPANGINA Herpangina is usually caused by coxsackievirus A and presents as acute-onset fever, sore throat, odynophagia, and grayish-white papulovesicular lesions on an erythematous base that ulcerate. The lesions can persist for weeks; are present on the soft palate, anterior pillars of the tonsils, and uvula; and are concentrated in the posterior portion of the mouth. In contrast to herpes stomatitis, enteroviral herpangina is not associated with gingivitis. Acute lymphonodular pharyngitis associated with coxsackievirus A10 presents as white or yellow nodules surrounded by erythema in the posterior oropharynx. The lesions do not ulcerate.

ACUTE HEMORRHAGIC CONJUNCTIVITIS Patients with acute hemorrhagic conjunctivitis present with an acute onset of severe eye pain, blurred vision, photophobia, and watery discharge from the eye. Examination reveals edema, chemosis, and subconjunctival hemorrhage and often shows punctate keratitis and conjunctival follicles as well (Fig. 228-2). Preauricular adenopathy is often found. Epidemics and nosocomial spread have been associated with enterovirus 70 and coxsackievirus A24. Systemic symptoms, including headache and fever, develop in 20% of cases, and recovery is usually complete in 10 days. The sudden onset and short duration of the illness help to distinguish acute hemorrhagic conjunctivitis from other ocular infections, such as those due to adenovirus and *Chlamydia trachomatis*. Paralysis has been associated with some cases of acute hemorrhagic conjunctivitis due to enterovirus 70 during epidemics.

OTHER MANIFESTATIONS Enteroviruses are an infrequent cause of childhood pneumonia and the common cold. In the fall of 2014, enterovirus D68 infection was confirmed in more than 500 persons with mild to severe respiratory illnesses in 43 U.S. states. Nearly all reported cases were in children, many of whom had asthma. Enterovirus D68 was detected in upper respiratory tract specimens from some patients with unexplained acute neurologic disease during outbreaks of infection with this virus; however, the virus was not detected in the CSF, and at present the link between the virus and neurologic disease is uncertain. Coxsackievirus B has been isolated at autopsy from the pancreas of a few children presenting with type 1 diabetes mellitus; however, most attempts to isolate the virus have been unsuccessful. Other diseases that have been associated with enterovirus infection include parotitis, bronchitis, bronchiolitis, croup, infectious lymphocytosis, polymyositis, acute arthritis, and acute nephritis.

DIAGNOSIS

Isolation of enterovirus in cell culture is the traditional diagnostic procedure. While cultures of stool, nasopharyngeal, or throat samples from patients with enterovirus diseases are often positive, isolation of the virus from these sites does not prove that it is directly associated with disease because these sites are frequently colonized for weeks in

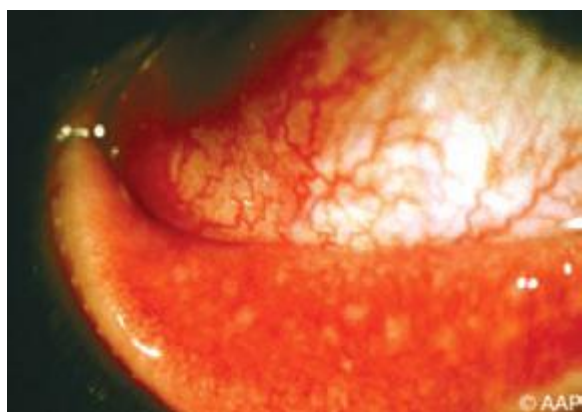


FIGURE 228-2 Acute hemorrhagic conjunctivitis due to enterovirus 70. (Image reprinted with permission from Red Book 2012: Committee on Infectious Diseases, 29th ed. Used with permission of the American Academy of Pediatrics.)

patients with subclinical infections. Isolation of virus from the throat is more likely to be associated with disease than is isolation from the stool since virus is shed for shorter periods from the throat. Cultures of CSF, serum, fluid from body cavities, or tissues are positive less frequently, but a positive result is indicative of disease caused by enterovirus. In some cases, the virus is isolated only from the blood or only from the CSF; therefore, it is important to culture multiple sites. Cultures are more likely to be positive earlier than later in the course of infection. Most human enteroviruses can be detected within a week after inoculation of cell cultures. Cultures may be negative because of the presence of neutralizing antibody, lack of susceptibility of the cells used, or inappropriate handling of the specimen. Coxsackievirus A may require inoculation into special cell-culture lines or into suckling mice.

Identification of the enterovirus serotype is useful primarily for epidemiologic studies and, with a few exceptions, has little clinical utility. It is important to identify serious infections with enterovirus during epidemics and to distinguish the vaccine strain of poliovirus from the other enteroviruses in the throat or in the feces. Stool and throat samples for culture as well as acute- and convalescent-phase serum specimens should be obtained from all patients with suspected poliomyelitis. In the absence of a positive CSF culture, a positive culture of stool obtained within the first 2 weeks after the onset of symptoms is most often used to confirm the diagnosis of poliomyelitis. If poliovirus infection is suspected, two or more fecal and throat swab samples should be obtained at least 1 day apart and cultured for enterovirus as soon as possible. If poliovirus is isolated, it should be sent to the CDC for identification as either wild-type or vaccine virus.

Reverse-transcriptase polymerase chain reaction (PCR) has been used to amplify viral nucleic acid from CSF, serum, urine, stool, conjunctiva, throat swabs, and tissues. A pan-enterovirus PCR assay can detect all human enteroviruses. With the proper controls, PCR of the CSF is highly sensitive (70–100%) and specific (>80%) and is more rapid than culture. PCR of the CSF is less likely to be positive when patients present ≥ 3 days after the onset of meningitis or with enterovirus 71 infection; in these cases, PCR of throat or rectal swabs—although less specific than PCR of CSF—should be considered.

PCR of serum is also highly sensitive and specific in the diagnosis of disseminated disease. PCR may be particularly helpful for the diagnosis and follow-up of enterovirus disease in immunodeficient patients receiving immunoglobulin therapy, whose viral cultures may be negative. Antigen detection is less sensitive than PCR.

Serologic diagnosis of enterovirus infection is limited by the large number of serotypes and the lack of a common antigen. Demonstration of seroconversion may be useful in rare cases for confirmation of culture results, but serologic testing is usually limited to epidemiologic studies. Serum should be collected and frozen soon after the onset of disease and again ~ 4 weeks later. Measurement of neutralizing titers is the most accurate method for antibody determination; measurement of complement-fixation titers is usually less sensitive. Titers of virus-specific IgM are elevated in both acute and chronic infection.

TREATMENT ENTEROVIRUS INFECTIONS

Most enterovirus infections are mild and resolve spontaneously; however, intensive supportive care may be needed for cardiac, hepatic, or CNS disease. IV, intrathecal, or intraventricular immunoglobulin has been used with apparent success in some cases for the treatment of chronic enterovirus meningoencephalitis and dermatomyositis in patients with hypogammaglobulinemia or agammaglobulinemia. The disease may stabilize or resolve during therapy; however, some patients decline inexorably despite therapy. IV immunoglobulin often prevents severe enterovirus disease in these patients. IV administration of immunoglobulin with high titers of antibody to the infecting virus has been used in some cases of life-threatening infection in neonates, who may not have maternally acquired antibody. In one trial involving neonates with enterovirus infections, immunoglobulin containing very high titers of antibody to the infecting virus reduced rates of viremia; however, the study was too small to show a substantial clinical benefit. The level of

1294 enteroviral antibodies varies with the immunoglobulin preparation. A phase 2 trial of pleconaril for severe neonatal enterovirus disease has been completed; however, as of this writing, the results have not been reported and the drug is not available on a compassionate-use basis. Glucocorticoids are contraindicated.

Good hand-washing practices and the use of gowns and gloves are important in limiting nosocomial transmission of enteroviruses during epidemics. Enteric precautions are indicated for 7 days after the onset of enterovirus infections. Enterovirus 71 vaccine candidates are under development.

PREVENTION AND ERADICATION OF POLIOVIRUS



(See also Chap. 148) After a peak of 57,879 cases of poliomyelitis in the United States in 1952, the introduction of IPV in 1955 and of OPV in 1961 ultimately eradicated disease due to wild-type poliovirus in the Western Hemisphere. Such disease has not been documented in the United States since 1979, when cases occurred among religious groups who had declined immunization. In the Western Hemisphere, paralysis due to wild-type poliovirus was last documented in 1991.

In 1988, the World Health Organization adopted a resolution to eradicate poliomyelitis by the year 2000. From 1988 to 2001, the number of cases worldwide decreased by >99%, with only 496 confirmed cases reported in 2001. Wild-type poliovirus type 2 has not been detected in the world since 1999. The Americas were certified free of indigenous wild-type poliovirus transmission in 1994, the Western Pacific Region in 2000, and the European Region in 2002. However, in 2002, there were 1922 cases of polio, with 1600 cases reported in India. In fact, after the nadir of 496 cases in 2001, 21 countries that had previously been free of polio reported cases imported from 6 polio-endemic countries in 2002–2005. By 2006, polio transmission had been reduced in most of these 21 countries. In 2012, 293 cases of polio were reported (the lowest number ever in a 1-year period); 85% were from Nigeria, Pakistan, and Afghanistan, the only countries where polio remains endemic (Table 228-2). As of November 2013, there had been 390 cases of polio in 2013 compared with 293 cases in 2012. The increase was associated with a marked rise in imported cases, including more than 180 cases in Somalia, more than 10 cases each in Kenya and Syria, and cases in Cameroon and Ethiopia. Also in 2013, wild-type poliovirus was detected in sewage in Israel, prompting a massive vaccination campaign with OPV. As of November 2013, India had not reported a case of polio since January 2011. Polio is a source of concern for unimmunized or partially immunized travelers. Importation of poliovirus accounted for ~50% of cases in 2013. Clearly, global eradication of polio is necessary to eliminate the risk of importation of wild-type virus. Outbreaks are thought to have been facilitated by suboptimal rates of vaccination, isolated pockets of unvaccinated children, poor sanitation and crowding, improper vaccine-storage conditions, and a reduced level of response to one of the serotypes in the vaccine.

TABLE 228-2 LABORATORY-CONFIRMED CASES OF POLIOMYELITIS IN 2012

Country	Type of Transmission	No. of Cases
Nigeria	Endemic	130 ^a
Pakistan	Endemic	74 ^b
Afghanistan	Endemic	46 ^c
Chad	Imported	17 ^d
Democratic Republic of the Congo	Vaccine-derived	17
Kenya	Vaccine-derived	3
Yemen	Vaccine-derived	2
China	Vaccine-derived	2
Niger	Imported	1
Somalia	Vaccine-derived	1
Total		293

^aOf these cases, 8 were vaccine-derived. ^bOf these cases, 16 were vaccine-derived.

^cOf these cases, 9 were vaccine-derived. ^dOf these cases, 12 were vaccine-derived.

Source: World Health Organization.

While the global eradication campaign has markedly reduced the number of cases of endemic polio, doubts have been raised as to whether eradication is a realistic goal, given the large number of asymptomatic infections and the political instability in developing countries.

The occurrence of outbreaks of poliomyelitis due to circulating vaccine-derived poliovirus of all three types has been increasing, especially in areas with low vaccination rates. In Egypt, 32 cases of vaccine-derived polio occurred in 1983–1993; in the Dominican Republic and Haiti, 21 cases occurred in 2000–2001; in Indonesia, 46 cases were reported in 2005; in Nigeria, 385 cases occurred in 2005–2012; in the Democratic Republic of the Congo, 64 cases were reported in 2008–2012; in Pakistan, 16 cases occurred in 2012, and at least 30 cases occurred in 2013. These OPV-derived viruses reverted to a more neurovirulent phenotype after undetected circulation (probably for >2 years). The epidemic in Hispaniola was rapidly terminated after intensive vaccination with OPV. In 2005, a case of vaccine-derived polio occurred in an unvaccinated U.S. woman returning from a visit to Central and South America. In the same year, an unvaccinated immunocompromised infant in Minnesota was found to be shedding vaccine-derived poliovirus; further investigation identified 4 of 22 infants in the same community who were shedding the virus. All 5 infants were asymptomatic. These outbreaks emphasize the need for maintaining high levels of vaccine coverage and continued surveillance for circulating virus.

IPV is used in most industrialized countries and OPV in most developing countries, including those in which polio still is or recently was endemic. While IM injections of other vaccines (live or attenuated) can be given concurrently with OPV, unnecessary IM injections should be avoided during the first month after OPV vaccination because they increase the risk of vaccine-associated paralysis. Since 1988, an enhanced-potency inactivated poliovirus vaccine has been available in the United States.

After several doses of OPV alone, the seropositivity rate for individual poliovirus serotypes may still be suboptimal for children in developing countries; one or more supplemental doses of IPV can increase the rate of seropositivity for these serotypes. Against a given serotype, monovalent OPV containing only that serotype is more immunogenic than trivalent vaccine because of a lack of interference from other serotypes. With eradication of wild-type poliovirus type 2, bivalent OPV (types 1 and 3), which was shown to be superior to trivalent OPV, has been the vaccine of choice to eliminate polio and has markedly reduced rates of polio in Nigeria. As the frequency of wild-type polio declines and reports of polio associated with circulating vaccine-derived viruses increase, the World Health Organization is investigating whether IPV can be produced from OPV strains that require less biocontainment, ultimately replacing OPV.

OPV and IPV induce antibodies that persist for at least 5 years. Both vaccines induce IgG and IgA antibodies. Compared with recipients of IPV, recipients of OPV shed less virus and less frequently develop reinfection with wild-type virus after exposure to poliovirus. Although IPV is safe and efficacious, OPV offers the advantages of ease of administration, lower cost, and induction of intestinal immunity resulting in a reduction in the risk of community transmission of wild-type virus. Because of progress toward global eradication of polio and the continued occurrence of cases of vaccine-associated polio, an all-IPV regimen was recommended in 2000 for childhood poliovirus vaccination in the United States, with vaccine administration at 2, 4, and 6–18 months and 4–6 years of age. The risk of vaccine-associated polio should be discussed before OPV is administered. Recommendations for vaccination of adults are listed in Table 228-3.

There are concerns about discontinuing vaccination in the event that endemic spread of poliovirus is eliminated. Among the reasons for these concerns are that poliovirus is shed from some immunocompromised persons for >10 years, that vaccine-derived poliovirus can circulate and cause disease, and that wild-type poliovirus is present in research laboratories.

PARECHOVIRUSES

Human parechoviruses (HPeVs), like enteroviruses, are members of the family Picornaviridae. The 16 serotypes of HPeV commonly cause

TABLE 228-3 RECOMMENDATIONS FOR POLIOVIRUS VACCINATION OF ADULTS

- Most adults in the United States have been vaccinated during childhood and are at little risk of exposure to wild-type virus in the United States. Immunization is recommended for those with a higher risk of exposure than the general population, including:
 - travelers to areas where poliovirus is or may be epidemic or endemic;
 - members of communities or population groups with disease caused by wild-type polioviruses;
 - laboratory workers handling specimens that may contain wild-type polioviruses; and
 - health care workers in close contact with patients who may be excreting wild-type polioviruses.
- Three doses of IPV are recommended for adults who need to be immunized. The second dose should be given 1–2 months after the first dose; the third dose should be given 6–12 months after the second dose.
- Adults who are at increased risk of exposure to wild-type poliovirus and who have previously completed primary immunization should receive a single dose of IPV. Adults who did not complete primary immunization should receive the remaining required doses of IPV.

Abbreviation: IPV, inactivated poliovirus vaccine.

Source: Modified from Pickering LK, ed. Red Book 2012: Committee on Infectious Diseases, 29th ed.

infections in early childhood. HPeV-1 infections occur throughout the year, while other parechovirus infections occur more commonly in summer and fall. Infections with HPeVs present similarly to those due to enteroviruses and may cause generalized disease of the newborn, aseptic meningitis, encephalitis, transient paralysis, exanthems, respiratory tract disease, and gastroenteritis. While HPeV-1 is the most common serotype and generally causes mild disease, deaths of infants in the United States have been associated with HPeV-1, HPeV-3, and HPeV-6. HPeVs can be isolated from the same sites as enteroviruses, including the nasopharynx, stool, and respiratory tract secretions. PCR using pan-enterovirus primers does not detect HPeVs, and while PCR assays are performed by the CDC and research laboratories, many commercial laboratories do not perform the test.

REOVIRUSES

Reoviruses are double-stranded RNA viruses encompassing three serotypes. Serologic studies indicate that most humans are infected with reoviruses during childhood. Most infections either are asymptomatic or cause mild upper respiratory tract symptoms. Reovirus is considered a rare cause of mild gastroenteritis or meningitis in infants and children. Speculation regarding an association of reovirus type 3 with idiopathic neonatal hepatitis and extrahepatic biliary atresia is based on an elevated prevalence of antibody to reovirus in some affected patients and the detection of viral RNA by PCR in hepatobiliary tissues in some studies. New orthoreoviruses have been associated with human disease—e.g., Melaka and Kampar viruses with fever and acute respiratory disease in Malaysia, and Nelson Bay virus with acute respiratory disease in a traveler from Bali.

229 Measles (Rubeola)

Kaitlin Rainwater-Lovett, William J. Moss

DEFINITION

Measles is a highly contagious viral disease that is characterized by a prodromal illness of fever, cough, coryza, and conjunctivitis followed by the appearance of a generalized maculopapular rash. Before the widespread use of measles vaccines, it was estimated that measles caused between 5 million and 8 million deaths worldwide each year.

GLOBAL CONSIDERATIONS



Remarkable progress has been made in reducing global measles incidence and mortality rates through measles vaccination. In the Americas, intensive vaccination and surveillance efforts—based in part on the successful Pan American Health Organization strategy of periodic nationwide measles vaccination campaigns (supplementary immunization activities, or SIAs)—and high levels of routine measles vaccine coverage interrupted endemic transmission of measles virus. In the United States, high-level coverage with two doses of measles vaccine eliminated endemic measles virus transmission in 2000. More recently, progress has been made in reducing measles incidence and mortality rates in sub-Saharan Africa and Asia as a consequence of increasing routine measles vaccine coverage and provision of a second dose of measles vaccine through mass measles vaccination campaigns and childhood immunization programs.

In 2003, the World Health Assembly endorsed a resolution urging member countries to reduce the number of deaths attributed to measles by 50% (compared with 1999 estimates) by the end of 2005. This target was met. Global measles mortality rates were further reduced in 2008; during that year, there were an estimated 164,000 deaths due to measles (uncertainty bounds: 115,000 and 222,000 deaths). These achievements attest to the enormous public-health significance of measles vaccination. However, recent large outbreaks of measles in Europe and Africa illustrate the challenges faced in sustaining measles control: in these outbreaks, measles was imported into countries that had eliminated indigenous transmission of measles virus.

The Measles and Rubella Initiative, a partnership led by the American Red Cross, the United Nations Foundation, UNICEF, the U.S. Centers for Disease Control and Prevention (CDC), and the World Health Organization (WHO), is playing an important role in reducing global measles incidence and mortality rates. Since its inception in 2001, the Initiative has provided governments and communities in more than 80 countries with technical and financial support for routine immunization activities, mass vaccination campaigns, and disease surveillance systems. Through its 2012–2020 Global Measles and Rubella Strategic Plan, the Initiative aims to reduce measles deaths by 95% (compared with year 2000 estimates) by 2015 and to eliminate measles from at least five of the six WHO regions by 2020. As regional goals for measles elimination are set, global measles eradication is likely to become a public health goal in the near future.

ETIOLOGY

Measles virus is a spherical, nonsegmented, single-stranded, negative-sense RNA virus and a member of the *Morbillivirus* genus in the family Paramyxoviridae. Measles was originally a zoonotic infection, arising from animal-to-human transmission of an ancestral morbillivirus ~10,000 years ago, when human populations had attained sufficient size to sustain virus transmission. Although RNA viruses typically have high mutation rates, measles virus is considered to be an antigenically monotypic virus; i.e., the surface proteins responsible for inducing protective immunity have retained their antigenic structure across time and distance. The public health significance of this stability is that measles vaccines developed decades ago from a single strain of measles virus remain protective worldwide. Measles virus is killed by ultraviolet light and heat, and attenuated measles vaccine viruses retain these characteristics, necessitating a cold chain for vaccine transport and storage.

EPIDEMIOLOGY

Measles virus is one of the most highly contagious directly transmitted pathogens. Outbreaks can occur in populations in which <10% of persons are susceptible. Chains of transmission are common among household contacts, school-age children, and health care workers. There are no latent or persistent measles virus infections that result in prolonged contagiousness, nor are there animal reservoirs for the virus. Thus, measles virus can be maintained in human populations only by an unbroken chain of acute infections, which requires

1296 a continuous supply of susceptible individuals. Newborns become susceptible to measles virus infection when passively acquired maternal antibody is lost; when not vaccinated, these infants account for the bulk of new susceptible individuals.



Endemic measles has a typical temporal pattern characterized by yearly seasonal epidemics superimposed on longer epidemic cycles of 2–5 years or more. In temperate climates, annual measles outbreaks typically occur in the late winter and early spring. These annual outbreaks are probably attributable to social networks facilitating transmission (e.g., congregation of children at school) and environmental factors favoring the viability and transmission of measles virus. Measles cases continue to occur during interepidemic periods in large populations, but at low incidence. The longer epidemic cycles occurring every several years result from the accumulation of susceptible persons over successive birth cohorts and the subsequent decline in the number of susceptibles following an outbreak.

Secondary attack rates among susceptible household and institutional contacts generally exceed 90%. The average age at which measles occurs depends on rates of contact with infected persons, protective maternal antibody decline, and vaccine coverage. In densely populated urban settings with low-level vaccination coverage, measles is a disease of infants and young children. The cumulative distribution can reach 50% by 1 year of age, with a significant proportion of children acquiring measles before 9 months—the age of routine vaccination in many countries, in line with the schedule recommended by the WHO's Expanded Programme on Immunization. As measles vaccine coverage increases or population density decreases, the age distribution shifts toward older children. In such situations, measles cases predominate in school-age children. Infants and young children, although susceptible if not protected by vaccination, are not exposed to measles virus at a rate sufficient to cause a large disease burden in this age group. As vaccination coverage increases further, the age distribution of cases may be shifted into adolescence and adulthood; this distribution is seen in measles outbreaks in the United States and necessitates targeted measles vaccination programs for these older age groups.

Persons with measles are infectious for several days before and after the onset of rash, when levels of measles virus in blood and body fluids are highest and when cough, coryza, and sneezing, which facilitate virus spread, are most severe. The contagiousness of measles before the onset of recognizable disease hinders the effectiveness of quarantine measures. Viral shedding by children with impaired cell-mediated immunity can be prolonged.

Medical settings are well-recognized sites of measles virus transmission. Children may present to health care facilities during the prodrome, when the diagnosis is not obvious although the child is infectious and is likely to infect susceptible contacts. Health care workers can acquire measles from infected children and transmit measles virus to others. Nosocomial transmission can be reduced by maintenance of a high index of clinical suspicion, use of appropriate isolation precautions when measles is suspected, administration of measles vaccine to susceptible children and health care workers, and documentation of health care workers' immunity to measles (i.e., proof of receipt of two doses of measles vaccine or detection of antibodies to measles virus).

As efforts at measles control are increasingly successful, public perceptions of the risk of measles as a disease diminish and are replaced by concerns about possible adverse events associated with measles vaccine. As a consequence, numerous measles outbreaks have occurred because of opposition to vaccination on religious or philosophical grounds or unfounded fears of serious adverse events (see "Active Immunization," below).

PATHOGENESIS

Measles virus is transmitted primarily by respiratory droplets over short distances and, less commonly, by small-particle aerosols that remain suspended in the air for long periods. Airborne transmission appears to be important in certain settings, including schools, physicians' offices, hospitals, and enclosed public places. The virus can be

transmitted by direct contact with infected secretions but does not survive for long on fomites.

The incubation period for measles is ~10 days to fever onset and 14 days to rash onset. This period may be shorter in infants and longer (up to 3 weeks) in adults. Infection is initiated when measles virus is deposited on epithelial cells in the respiratory tract, oropharynx, or conjunctivae (Fig. 229-1A). During the first 2–4 days after infection, measles virus proliferates locally in the respiratory mucosa and spreads to draining lymph nodes. Virus then enters the bloodstream in infected leukocytes (primarily monocytes), producing the primary viremia that disseminates infection throughout the reticuloendothelial system. Further replication results in secondary viremia that begins 5–7 days after infection and disseminates measles virus throughout the body. Replication of measles virus in these target organs, together with the host's immune response, is responsible for the signs and symptoms of measles that occur 8–12 days after infection and mark the end of the incubation period (Fig. 229-1B).

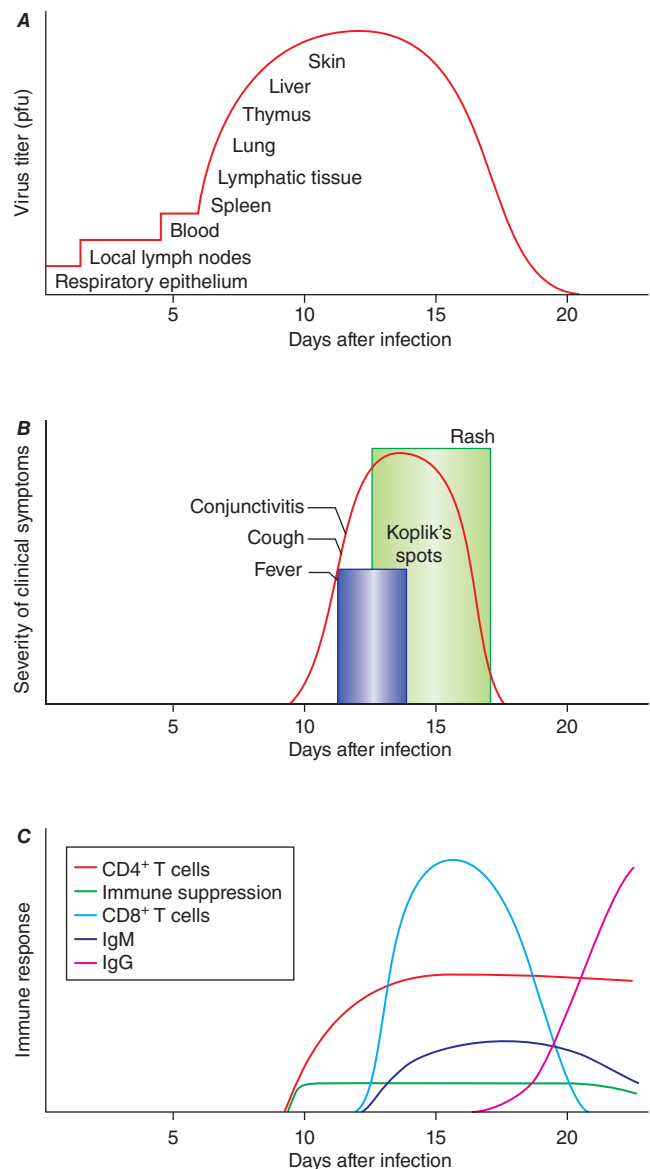


FIGURE 229-1 Measles virus infection: pathogenesis, clinical features, and immune responses. **A.** Spread of measles virus, from initial infection of the respiratory tract through dissemination to the skin. **B.** Appearance of clinical signs and symptoms, including Koplik's spots and rash. **C.** Antibody and T cell responses to measles virus. The signs and symptoms of measles arise coincident with the host immune response. (Source: Modified from WJ Moss, DE Griffin: *Nat Rev Microbiol* 4:900, 2006.)

IMMUNE RESPONSES

Host immune responses to measles virus are essential for viral clearance, clinical recovery, and the establishment of long-term immunity (Fig. 229-1C). Early nonspecific (innate) immune responses during the prodromal phase include activation of natural killer cells and increased production of antiviral proteins. The adaptive immune responses consist of measles virus-specific antibody and cellular responses. The protective efficacy of antibodies to measles virus is illustrated by the immunity conferred to infants from passively acquired maternal antibodies and the protection of exposed, susceptible individuals after administration of anti-measles virus immunoglobulin. The first measles virus-specific antibodies produced after infection are of the IgM subtype, with a subsequent switch to predominantly IgG1 and IgG4 isotypes. The IgM antibody response is typically absent following reexposure or revaccination and serves as a marker of primary infection.

The importance of cellular immunity to measles virus is demonstrated by the ability of children with agammaglobulinemia (congenital inability to produce antibodies) to recover fully from measles and the contrasting picture for children with severe defects in T lymphocyte function, who often develop severe or fatal disease (Chap. 374). The initial predominant T_H1 response (characterized by interferon γ) is essential for viral clearance, and the later T_H2 response (characterized by interleukin 4) promotes the development of measles virus-specific antibodies that are critical for protection against reinfection.

The duration of protective immunity following wild-type measles virus infection is generally thought to be lifelong. Immunologic memory to measles virus includes both continued production of measles virus-specific antibodies and circulation of measles virus-specific CD4⁺ and CD8⁺ T lymphocytes.

However, the intense immune responses induced by measles virus infection are paradoxically associated with depressed responses to unrelated (non-measles virus) antigens, which persist for several weeks to months beyond resolution of the acute illness. This state of immune suppression enhances susceptibility to secondary infections with bacteria and viruses that cause pneumonia and diarrhea and is responsible for a substantial proportion of measles-related morbidity and deaths. Delayed-type hypersensitivity responses to recall antigens, such as tuberculin, are suppressed, and cellular and humoral responses to new antigens are impaired. Reactivation of tuberculosis and remission of autoimmune diseases after measles have been described and are attributed to this period of immune suppression.

APPROACH TO THE PATIENT:

Measles

Clinicians should consider measles in persons presenting with fever and generalized erythematous rash, particularly when measles virus is known to be circulating or the patient has a history of travel to endemic areas. Appropriate precautions must be taken to prevent nosocomial transmission. The diagnosis requires laboratory confirmation except during large outbreaks in which an epidemiologic link to a confirmed case can be established. Care is largely supportive and consists of the administration of vitamin A and antibiotics (see “Treatment,” below). Complications of measles, including secondary bacterial infections and encephalitis, may occur after acute illness and require careful monitoring, particularly in immunocompromised persons.

CLINICAL MANIFESTATIONS

In most persons, the signs and symptoms of measles are highly characteristic (Fig. 229-1B). Fever and malaise beginning ~10 days after exposure are followed by cough, coryza, and conjunctivitis. These signs and symptoms increase in severity over 4 days. Koplik's spots (see Fig. 25e-2) develop on the buccal mucosa ~2 days before the rash appears. The characteristic rash of measles (see Fig. 25e-3) begins 2 weeks after infection, when the clinical manifestations are most

severe, and signal the host's immune response to the replicating virus. Headache, abdominal pain, vomiting, diarrhea, and myalgia may be present.

Koplik's spots (see Fig. 25e-2) are pathognomonic of measles and consist of bluish white dots ~1 mm in diameter surrounded by erythema. The lesions appear first on the buccal mucosa opposite the lower molars but rapidly increase in number to involve the entire buccal mucosa. They fade with the onset of rash.

The rash of measles begins as erythematous macules behind the ears and on the neck and hairline. The rash progresses to involve the face, trunk, and arms (see Fig. 25e-3), with involvement of the legs and feet by the end of the second day. Areas of confluent rash appear on the trunk and extremities, and petechiae may be present. The rash fades slowly in the same order of progression as it appeared, usually beginning on the third or fourth day after onset. Resolution of the rash may be followed by desquamation, particularly in undernourished children.

Because the characteristic rash of measles is a consequence of the cellular immune response, it may not develop in persons with impaired cellular immunity (e.g., those with AIDS; Chap. 226). These persons have a high case-fatality rate and frequently develop giant-cell pneumonitis caused by measles virus. T lymphocyte defects due to causes other than HIV-1 infection (e.g., cancer chemotherapy) also are associated with increased severity of measles.

A severe atypical measles syndrome was observed in recipients of a formalin-inactivated measles vaccine (used in the United States from 1963 to 1967 and in Canada until 1970) who were subsequently exposed to wild-type measles virus. The atypical rash began on the palms and soles and spread centripetally to the proximal extremities and trunk, sparing the face. The rash was initially erythematous and maculopapular but frequently progressed to vesicular, petechial, or purpuric lesions (see Fig. 25e-22).

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of measles includes other causes of fever, rash, and conjunctivitis, including rubella, Kawasaki disease, infectious mononucleosis, roseola, scarlet fever, Rocky Mountain spotted fever, enterovirus or adenovirus infection, and drug sensitivity. Rubella is a milder illness without cough and with distinctive lymphadenopathy. The rash of roseola (exanthem subitum) (see Fig. 25e-5) appears after fever has subsided. The atypical lymphocytosis in infectious mononucleosis contrasts with the leukopenia commonly observed in children with measles.

DIAGNOSIS

Measles is readily diagnosed on clinical grounds by clinicians familiar with the disease, particularly during outbreaks. Koplik's spots (see Fig. 25e-2) are especially helpful because they appear early and are pathognomonic. Clinical diagnosis is more difficult (1) during the prodromal illness; (2) when the rash is attenuated by passively acquired antibodies or prior immunization; (3) when the rash is absent or delayed in immunocompromised children or severely undernourished children with impaired cellular immunity; and (4) in regions where the incidence of measles is low and other pathogens are responsible for the majority of illnesses with fever and rash. The CDC case definition for measles requires (1) a generalized maculopapular rash of at least 3 days' duration; (2) fever of at least 38.3°C (101°F); and (3) cough, coryza, or conjunctivitis.

Serology is the most common method of laboratory diagnosis. The detection of measles virus-specific IgM in a single specimen of serum or oral fluid is considered diagnostic of acute infection, as is a four-fold or greater increase in measles virus-specific IgG antibody levels between acute- and convalescent-phase serum specimens. Primary infection in the immunocompetent host results in antibodies that are detectable within 1–3 days of rash onset and reach peak levels in 2–4 weeks. Measles virus-specific IgM antibodies may not be detectable until 4–5 days or more after rash onset and usually fall to undetectable levels within 4–8 weeks of rash onset.

Several methods for measurement of antibodies to measles virus are available. Neutralization tests are sensitive and specific, and the results are highly correlated with protective immunity; however, these tests

1298 require propagation of measles virus in cell culture and thus are expensive and laborious. Commercially available enzyme immunoassays are most frequently used. Measles can also be diagnosed by isolation of the virus in cell culture from respiratory secretions, nasopharyngeal or conjunctival swabs, blood, or urine. Direct detection of giant cells in respiratory secretions, urine, or tissue obtained by biopsy provides another method of diagnosis.

For detection of measles virus RNA by reverse-transcriptase polymerase chain reaction amplification of RNA extracted from clinical specimens, primers targeted to highly conserved regions of measles virus genes are used. Extremely sensitive and specific, this assay may also permit identification and characterization of measles virus genotypes for molecular epidemiologic studies and can distinguish wild-type from vaccine virus strains.

TREATMENT MEASLES

There is no specific antiviral therapy for measles. Treatment consists of general supportive measures, such as hydration and administration of antipyretic agents. Because secondary bacterial infections are a major cause of morbidity and death attributable to measles, effective case management involves prompt antibiotic treatment for patients who have clinical evidence of bacterial infection, including pneumonia and otitis media. *Streptococcus pneumoniae* and *Haemophilus influenzae* type b are common causes of bacterial pneumonia following measles; vaccines against these pathogens probably lower the incidence of secondary bacterial infections following measles.

Vitamin A is effective for the treatment of measles and can markedly reduce rates of morbidity and mortality. The WHO recommends administration of once-daily doses of 200,000 IU of vitamin A for 2 consecutive days to all children with measles who are ≥ 12 months of age. Lower doses are recommended for younger children: 100,000 IU per day for children 6–12 months of age and 50,000 IU per day for children < 6 months old. A third dose is recommended 2–4 weeks later for children with evidence of vitamin A deficiency. While such deficiency is not a widely recognized problem in the United States, many American children with measles do, in fact, have low serum levels of vitamin A, and these children experience increased measles-associated morbidity. The Committee on Infectious Diseases of the American Academy of Pediatrics recommends that the administration of two consecutive daily doses of vitamin A be considered for children who are hospitalized with measles and its complications as well as for children with measles who are immunodeficient; who have ophthalmologic evidence of vitamin A deficiency, impaired intestinal absorption, or moderate to severe malnutrition; or who have recently immigrated from areas with high measles mortality rates. Parenteral and oral formulations of vitamin A are available.

Anecdotal reports have described the recovery of previously healthy pregnant and immunocompromised patients with measles pneumonia and of immunocompromised patients with measles encephalitis after treatment with aerosolized and IV ribavirin. However, the clinical benefits of ribavirin in measles have not been conclusively demonstrated in clinical trials.

COMPLICATIONS

Most complications of measles involve the respiratory tract and include the effects of measles virus replication itself and secondary bacterial infections. Acute laryngotracheobronchitis (croup) can occur during measles and may result in airway obstruction, particularly in young children. Giant-cell pneumonitis due to replication of measles virus in the lungs can develop in immunocompromised children, including those with HIV-1 infection. Many children with measles develop diarrhea, which contributes to undernutrition.

Most complications of measles result from secondary bacterial infections of the respiratory tract that are attributable to a state of immune suppression lasting for several weeks to months after acute measles. Otitis media and bronchopneumonia are most common and

may be caused by *S. pneumoniae*, *H. influenzae* type b, or staphylococci. Recurrence of fever or failure of fever to subside with the rash suggests secondary bacterial infection.

Rare but serious complications of measles involve the central nervous system (CNS). Postmeasles encephalomyelitis complicates ~ 1 in 1000 cases, affecting mainly older children and adults. Encephalomyelitis occurs within 2 weeks of rash onset and is characterized by fever, seizures, and a variety of neurologic abnormalities. The finding of periventricular demyelination, the induction of immune responses to myelin basic protein, and the absence of measles virus in the brain suggest that postmeasles encephalomyelitis is an autoimmune disorder triggered by measles virus infection. Other CNS complications that occur months to years after acute infection are measles inclusion body encephalitis (MIBE) and subacute sclerosing panencephalitis (SSPE). In contrast to postmeasles encephalomyelitis, MIBE and SSPE are caused by persistent measles virus infection. MIBE is a rare but fatal complication that affects individuals with defective cellular immunity and typically occurs months after infection. SSPE is a slowly progressive disease characterized by seizures and progressive deterioration of cognitive and motor functions, with death occurring 5–15 years after measles virus infection. SSPE most often develops in persons infected with measles virus at < 2 years of age.

PROGNOSIS



Most persons with measles recover and develop long-term protective immunity to reinfection. Measles case-fatality proportions vary with the average age of infection, the nutritional and immunologic status of the population, measles vaccine coverage, and access to health care. Among previously vaccinated persons who do become infected, disease is less severe and mortality rates are significantly lower. In developed countries, < 1 in 1000 children with measles die. In endemic areas of sub-Saharan Africa, the measles case-fatality proportion may be 5–10% or even higher. Measles is a major cause of childhood deaths in refugee camps and in internally displaced populations, where case-fatality proportions have been as high as 20–30%.

PREVENTION

Passive Immunization Human immunoglobulin given shortly after exposure can attenuate the clinical course of measles. In immunocompetent persons, administration of immunoglobulin within 72 h of exposure usually prevents measles virus infection and almost always prevents clinical measles. Administered up to 6 days after exposure, immunoglobulin will still prevent or modify the disease. Prophylaxis with immunoglobulin is recommended for susceptible household and nosocomial contacts who are at risk of developing severe measles, particularly children < 1 year of age, immunocompromised persons (including HIV-infected persons previously immunized with live attenuated measles vaccine), and pregnant women. Except for premature infants, children < 6 months of age usually will be partially or completely protected by passively acquired maternal antibody. If measles is diagnosed in a household member, all unimmunized children in the household should receive immunoglobulin. The recommended dose is 0.25 mL/kg given intramuscularly. Immunocompromised persons should receive 0.5 mL/kg. The maximum total dose is 15 mL. IV immunoglobulin contains antibodies to measles virus; the usual dose of 100–400 mg/kg generally provides adequate prophylaxis for measles exposures occurring as long as 3 weeks or more after IV immunoglobulin administration.

Active Immunization The first live attenuated measles vaccine was developed by passage of the Edmonston strain in chick embryo fibroblasts to produce the Edmonston B virus, which was licensed in 1963 in the United States. Further passage of Edmonston B virus produced the more attenuated Schwarz vaccine that currently serves as the standard in much of the world. The Moraten (“more attenuated Enders”) strain, which was licensed in 1968 and is used in the United States, is genetically closely related to the Schwarz strain.

Lyophilized measles vaccines are relatively stable, but reconstituted vaccine rapidly loses potency. Live attenuated measles vaccines are inactivated by light and heat and lose about half their potency at 20°C

and almost all their potency at 37°C within 1 h after reconstitution. Therefore, a cold chain must be maintained before and after reconstitution. Antibodies first appear 12–15 days after vaccination, and titers peak at 1–3 months. Measles vaccines are often combined with other live attenuated virus vaccines, such as those for mumps and rubella (MMR) and for mumps, rubella, and varicella (MMR-V).

The recommended age of first vaccination varies from 6 to 15 months and represents a balance between the optimal age for seroconversion and the probability of acquiring measles before that age. The proportions of children who develop protective levels of antibody after measles vaccination approximate 85% at 9 months of age and 95% at 12 months. Common childhood illnesses concomitant with vaccination may reduce the level of immune response, but such illness is not a valid reason to withhold vaccination. Measles vaccines have been well tolerated and immunogenic in HIV-1-infected children and adults, although antibody levels may wane. Because of the potential severity of wild-type measles virus infection in HIV-1-infected children, routine measles vaccination is recommended except for those who are severely immunocompromised. Measles vaccination is contraindicated in individuals with other severe deficiencies of cellular immunity because of the possibility of disease due to progressive pulmonary or CNS infection with the vaccine virus.

The duration of vaccine-induced immunity is at least several decades if not longer. Rates of secondary vaccine failure 10–15 years after immunization have been estimated at ~5% but are probably lower when vaccination takes place after 12 months of age. Decreasing antibody concentrations do not necessarily imply a complete loss of protective immunity: a secondary immune response usually develops after reexposure to measles virus, with a rapid rise in antibody titers in the absence of overt clinical disease.

Standard doses of currently licensed measles vaccines are safe for immunocompetent children and adults. Fever to 39.4°C (103°F) occurs in ~5% of seronegative vaccine recipients, and 2% of vaccine recipients develop a transient rash. Mild transient thrombocytopenia has been reported, with an incidence of ~1 case per 40,000 doses of MMR vaccine.

Since the publication of a report in 1998 hypothesizing that MMR vaccine may cause a syndrome of autism and intestinal inflammation, much public attention has focused on this purported association. The events that followed publication of this report led to diminished vaccine coverage in the United Kingdom and provide important lessons in the misinterpretation of epidemiologic evidence and the communication of scientific results to the public. The publication that incited the concern was a case series describing 12 children with a regressive developmental disorder and chronic enterocolitis; 9 of these children had autism. In 8 of the 12 cases, the parents associated onset of the developmental delay with MMR vaccination. This simple temporal association was misinterpreted and misrepresented as a possible causal relationship, first by the lead author of the study and then by elements of the media and the public. Subsequently, several comprehensive reviews and additional epidemiologic studies refuted evidence of a causal relationship between MMR vaccination and autism.

PROSPECTS FOR MEASLES ERADICATION

Progress in global measles control has renewed discussion of measles eradication. In contrast to poliovirus eradication, the eradication of measles virus will not entail challenges posed by prolonged shedding of potentially virulent vaccine viruses and environmental viral reservoirs. However, in comparison with smallpox eradication, higher levels of population immunity will be necessary to interrupt measles virus transmission, more highly skilled health care workers will be required to administer measles vaccines, and containment through case detection and ring vaccination will be more difficult for measles virus because of infectivity before rash onset. New tools, such as aerosol administration of measles vaccines, will facilitate mass vaccination campaigns. Despite enormous progress, measles remains a leading vaccine-preventable cause of childhood mortality worldwide and continues to cause outbreaks in communities with low vaccination coverage rates in industrialized nations.

230e Rubella (German Measles)

Laura A. Zimmerman, Susan E. Reef

Rubella was historically viewed as a variant of measles or scarlet fever. Not until 1962 was a separate viral agent for rubella isolated. After an epidemic of rubella in Australia in the early 1940s, the ophthalmologist Norman Gregg noticed the occurrence of congenital cataracts among infants whose mothers had reported rubella infection during early pregnancy, and congenital rubella syndrome (CRS; see “Clinical Manifestations,” below) was first described.

ETIOLOGY

Rubella virus is a member of the *Togaviridae* family and the only member of the genus *Rubivirus*. This single-stranded RNA enveloped virus measures 50–70 nm in diameter. Its core protein is surrounded by a single-layer lipoprotein envelope with spike-like projections containing two glycoproteins, E1 and E2. There is only one antigenic type of rubella virus, and humans are its only known reservoir.

PATHOGENESIS AND PATHOLOGY

Although the pathogenesis of postnatal (acquired) rubella has been well documented, data on pathology are limited because of the mildness of the disease. Rubella virus is spread from person to person via respiratory droplets. Primary implantation and replication in the nasopharynx are followed by spread to the lymph nodes. Subsequent viremia occurs, which in pregnant women often results in infection of the placenta. Placental virus replication may lead to infection of fetal organs. The pathology of CRS in the infected fetus is well defined, with almost all organs found to be infected; however, the pathogenesis of CRS is only poorly delineated. In tissue, infections with rubella virus have diverse effects, ranging from no obvious impact to cell destruction. The hallmark of fetal infection is chronicity, with persistence throughout fetal development in utero and for up to 1 year after birth.

Individuals with acquired rubella may shed virus from 7 days before rash onset to ~5–7 days thereafter. Both clinical and subclinical infections are considered contagious. Infants with CRS may shed large quantities of virus from bodily secretions, particularly from the throat and in the urine, up to 1 year of age. Outbreaks of rubella, including some in nosocomial settings, have originated with index cases of CRS. Thus only individuals immune to rubella should have contact with infants who have CRS or who are congenitally infected with rubella virus but are not showing signs of CRS.

EPIDEMIOLOGY

The largest recent rubella epidemic in the United States took place in 1964–1965, when an estimated 12.5 million cases occurred, resulting in ~20,000 cases of CRS. Since the introduction of the routine rubella vaccination program in the United States in 1969, the number of rubella cases reported each year has dropped by >99%; the rate of vaccination coverage with rubella-containing vaccine has been >90% among children 19–35 months old since 1995 and >95% for kindergarten and first-grade entrants since 1980. In 1989 a goal for the elimination of rubella and CRS in the United States was set, and in 2004 a panel of experts agreed unanimously that rubella was no longer an endemic disease in this country. The criteria used to document lack of endemic transmission included low disease incidence, high nationwide rubella antibody seroprevalence, outbreaks that were few and contained (i.e., small numbers of cases), and lack of endemic virus transmission (as assessed by genetic sequencing). In the United States, interruption of endemic transmission of rubella virus has been sustained since 2001; in 2012, however, three cases of CRS were reported in infants whose mothers had acquired rubella infection abroad. Thus health care providers should remain vigilant, considering the possibility of rubella infection in patients emigrating or returning from countries without rubella control programs and the accompanying potential for CRS among their infants.



FIGURE 230e-1 Mild maculopapular rash of rubella in a child.



Although rubella and CRS are no longer endemic in the United States, they remain important public health problems globally. The number of rubella cases reported worldwide in 1999 was ~900,000; this figure declined steadily to 94,030 in 2012. However, numbers of rubella cases are substantially underestimated because cases in many countries are identified through measles surveillance systems that are not specific for rubella. In 2010, it was estimated that 103,000 cases of CRS occur globally.

CLINICAL FEATURES

Acquired Rubella Acquired rubella commonly presents with a generalized maculopapular rash that usually lasts for up to 3 days (**Fig. 230e-1**), although as many as 50% of cases may be subclinical or without rash. When it occurs, the rash is usually mild and may be difficult to detect in persons with darker skin. In children, rash is usually the first sign of illness. However, in older children and adults, a 1- to 5-day prodrome often precedes the rash and may include low-grade fever, malaise, and upper respiratory symptoms. The incubation period is 14 days (range, 12–23 days).

Lymphadenopathy, particularly occipital and postauricular, may be noted during the second week after exposure. Although acquired rubella is usually thought of as a benign disease, arthralgia and arthritis are common in infected adults, particularly women. Thrombocytopenia and encephalitis are less common complications.

Congenital Rubella Syndrome The most serious consequence of rubella virus infection can develop when a woman becomes infected during pregnancy, particularly during the first trimester. The resulting complications may include miscarriage, fetal death, premature delivery, or live birth with congenital defects. Infants infected with rubella virus in utero may have myriad physical defects (**Table 230e-1**), which most commonly relate to the eyes, ears, and heart. This constellation of severe birth defects is known as *congenital rubella syndrome*. In addition to permanent manifestations, there are a host of transient physical manifestations, including thrombocytopenia with purpura/ptechiae (e.g., dermal erythroptosis, “blueberry muffin syndrome”). Some infants may be born with congenital rubella virus infection but have no apparent signs or symptoms of CRS and are referred to as “infants with congenital rubella infection only.”

DIAGNOSIS

Acquired Rubella Clinical diagnosis of acquired rubella is difficult because of the mimicry of many illnesses with rashes, the varied

TABLE 230e-1 COMMON TRANSIENT AND PERMANENT MANIFESTATIONS IN INFANTS WITH CONGENITAL RUBELLA SYNDROME

Transient Manifestations	Permanent Manifestations
Hepatosplenomegaly	Hearing impairment/deafness
Interstitial pneumonitis	Congenital heart defects (patent ductus arteriosus, pulmonary arterial stenosis)
Thrombocytopenia with purpura/ petechiae (e.g., dermal erythrocytosis, or "blueberry muffin syndrome")	Eye defects (cataracts, cloudy cornea, microphthalmos, pigmentary retinopathy, congenital glaucoma)
Hemolytic anemia	Microcephaly
Bony radiolucencies	Central nervous system sequelae (mental and motor delay, autism)
Intrauterine growth retardation	
Adenopathy	
Meningoencephalitis	

clinical presentations, and the high rates of subclinical and mild disease. Illnesses that may be similar to rubella in presentation include scarlet fever, roseola, toxoplasmosis, fifth disease, measles, and illnesses with suboccipital and postauricular lymphadenopathy. Thus laboratory documentation of rubella virus infection is considered the only reliable way to confirm acute disease.

Laboratory assessment of rubella infection is conducted by serologic and virologic methods. For acquired rubella, serologic diagnosis is most common and depends on the demonstration of IgM antibodies in an acute-phase serum specimen or a fourfold rise in IgG antibody titer between acute- and convalescent-phase specimens. The enzyme-linked immunosorbent assay IgM capture technique is considered most accurate for serologic diagnosis, but the indirect IgM assay also is acceptable. After rubella virus infection, IgM antibody may be detectable for up to 6 weeks. In case of a negative result for IgM in specimens taken earlier than day 5 after rash onset, serologic testing should be repeated. Although uncommon, reinfection with rubella virus is possible, and IgM antibodies may be present. To detect a rise in IgG antibody titer indicative of acute disease, the acute-phase serum specimen should be collected within 7–10 days after onset of illness and the convalescent-phase specimen ~14–21 days after the first specimen.

IgG avidity testing is used in conjunction with IgG testing. Low-avidity antibodies indicate recent infection. Mature (high-avidity) IgG antibodies most likely indicate an infection occurring at least 2 months previously. This test helps distinguish primary infection from reinfection. Avidity testing may be particularly useful in diagnosing rubella in pregnant women and assessing the risk of CRS.

Rubella virus can be isolated from the blood and nasopharynx during the prodromal period and for as long as 2 weeks after rash onset. However, as the secretion of virus in individuals with acquired rubella is maximal just before or up to 4 days after rash onset, this is the optimal time frame for collecting specimens for viral cultures. Rubella can also be diagnosed by viral RNA detection in a reverse-transcriptase polymerase chain reaction (RT-PCR) assay.

Congenital Rubella Syndrome A clinical diagnosis of CRS is reasonable when an infant presents with a combination of cataracts, hearing impairment, and heart defects; this pattern is seen in ~10% of infants with CRS. Infants may present with different combinations of defects depending on when infection occurs during gestation. Hearing impairment is the most common single defect of CRS. However, as with acquired rubella, laboratory diagnosis of congenital infection is highly recommended, particularly because most features of the clinical presentation are nonspecific and may be associated with other intrauterine infections. Early diagnosis of CRS facilitates appropriate medical intervention for specific disabilities and prompts implementation of infection control measures.

Diagnostic tests used to confirm CRS include serologic assays and virus detection. In an infant with congenital infection, serum IgM antibodies are normally present for up to 6 months but may be detectable for up to 1 year after birth. In some instances, IgM may not be detectable until 1 month of age; thus infants who have symptoms consistent with CRS but who test negative shortly after birth should be retested at 1 month. A

rubella serum IgG titer persisting beyond the time expected after passive transfer of maternal IgG antibody (i.e., a rubella titer that does not decline at the expected rate of a twofold dilution per month) is another serologic criterion used to confirm CRS.

In congenital infection, rubella virus is isolated most commonly from throat swabs and less commonly from urine and cerebrospinal fluid. Infants with congenital rubella may excrete virus for up to 1 year, but specimens for virus isolation are most likely to be positive if obtained within the first 6 months after birth. Rubella virus in infants with CRS can also be detected by RT-PCR.

Rubella Diagnosis in Pregnant Women In the United States, screening for rubella IgG antibodies is recommended as part of routine prenatal care. Pregnant women with a positive IgG antibody serologic test are considered immune. Susceptible pregnant women should be vaccinated postpartum.

A susceptible pregnant woman exposed to rubella virus should be tested for IgM antibodies and/or a fourfold rise in IgG antibody titer between acute- and convalescent-phase serum specimens to determine whether she was infected during pregnancy. Pregnant women with evidence of acute infection must be clinically monitored, and gestational age at the time of maternal infection must be determined to assess the possibility of risk to the fetus. Of women infected with rubella virus during the first 11 weeks of gestation, up to 90% deliver an infant with CRS; for maternal infection during the first 20 weeks of pregnancy, the CRS rate is 20%.

TREATMENT RUBELLA

No specific therapy is available for rubella virus infection. Symptom-based treatment for various manifestations, such as fever and arthralgia, is appropriate. Immunoglobulin does not prevent rubella virus infection after exposure and therefore is not recommended as routine postexposure prophylaxis. Although immunoglobulin may modify or suppress symptoms, it can create an unwarranted sense of security: infants with congenital rubella have been born to women who received immunoglobulin shortly after exposure. Administration of immunoglobulin should be considered only if a pregnant woman who has been exposed to rubella will not consider termination of the pregnancy under any circumstances. In such cases, IM administration of 20 mL of immunoglobulin within 72 h of rubella exposure may reduce—but does not eliminate—the risk of rubella.

PREVENTION

After the isolation of rubella virus in the early 1960s and the occurrence of a devastating pandemic, a vaccine for rubella was developed and licensed in 1969. Currently, the majority of rubella-containing vaccines (RCVs) used worldwide are combined measles and rubella (MR) or measles, mumps, and rubella (MMR) formulations. A tetravalent measles, mumps, rubella, and varicella (MMRV) vaccine is available but is not widely used.

The public health burden of rubella infection is measured primarily through the resulting CRS cases. The 1964–1965 rubella epidemic in the United States encompassed >30,000 infections during pregnancy. CRS occurred in ~20,000 infants born alive, including >11,000 infants who were deaf, >3500 infants who were blind, and almost 2000 infants who were mentally retarded. The cost of this epidemic exceeded \$1.5 billion. In 1983, the cost per child with CRS was estimated at \$200,000.

In most countries, there is little documented evidence to illuminate the epidemiology of CRS. Clusters of CRS cases have been reported in developing countries. Before the introduction of an immunization program, the incidence of CRS is 0.1–0.2 per 1000 live births during endemic periods and 1–4 per 1000 live births during epidemic periods. Where rubella virus is circulating and women of childbearing age are susceptible, CRS cases will continue to occur.

The most effective method of preventing acquired rubella and CRS is through vaccination with an RCV. One dose induces seroconversion in ≥95% of persons ≥1 year of age. Immunity is considered long-term and

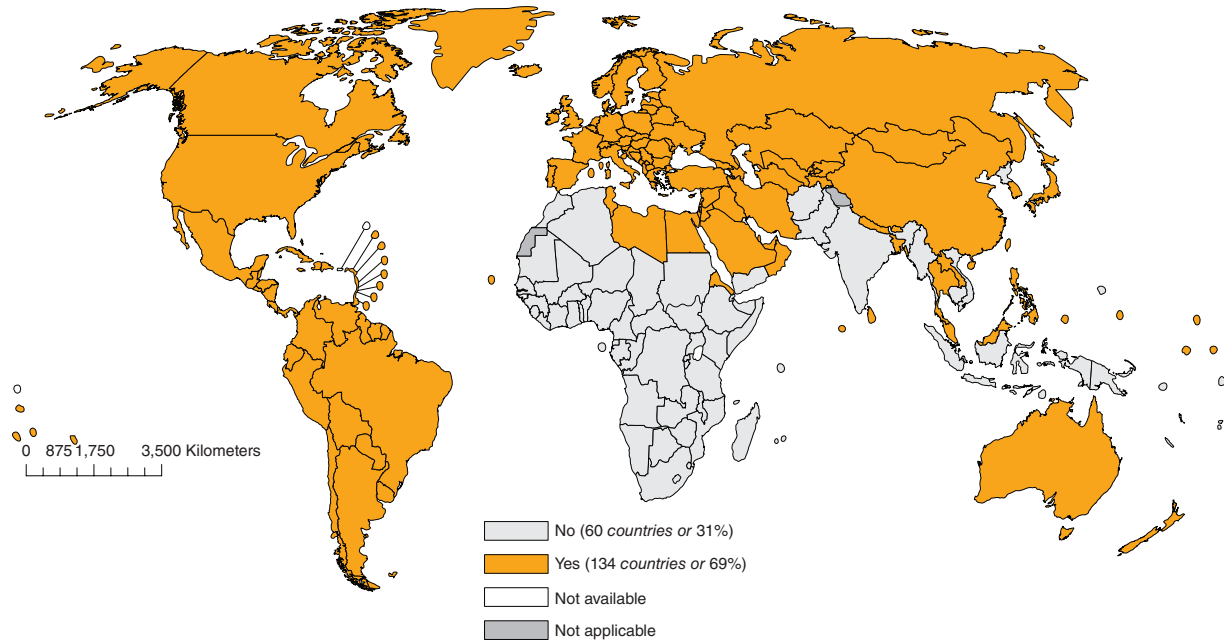


FIGURE 230e-2 Countries using rubella vaccine in their national immunization schedule, 2012. (From the World Health Organization.)

is probably lifelong. The most commonly used vaccine globally is the RA27/3 virus strain. The current recommendation for routine rubella vaccination in the United States is a first dose of MMR vaccine at 12–15 months of age and a second dose at 4–6 years. Target groups for rubella vaccine include children ≥ 1 year of age, adolescents and adults without documented evidence of immunity, individuals in congregate settings (e.g., college students, military personnel, child care and health care workers), and susceptible women before and after pregnancy.

Because of the theoretical risk of transmission of live attenuated rubella vaccine virus to the developing fetus, women known to be pregnant should not receive an RCV. In addition, pregnancy should

be avoided for 28 days after receipt of an RCV. In follow-up studies of 680 unknowingly pregnant women who received rubella vaccine, no infant was born with CRS. Receipt of an RCV during pregnancy is not ordinarily a reason to consider termination of the pregnancy.

As of 2012, 134 (69%) of the 194 member countries of the World Health Organization recommended inclusion of an RCV in the routine childhood vaccination schedule (Fig. 230e-2). Goals for control or elimination of rubella and CRS have been established in the American Region, the European Region, the South-East Asia Region, and the Western Pacific Region. The other two regions (Eastern Mediterranean and African) have not yet set such goals.

DEFINITION

Mumps is an illness characterized by acute-onset unilateral or bilateral tender, self-limited swelling of the parotid or other salivary gland(s) that lasts at least 2 days and has no other apparent cause.

ETIOLOGIC AGENT

Mumps is caused by a paramyxovirus with a negative-strand, nonsegmented RNA genome of 15,384 bases encoding at least 8 proteins: the nucleo- (N), phospho- (P), V, matrix (M), fusion (F), small hydrophobic (SH), hemagglutinin-neuraminidase (HN), and large (L) proteins. The N, P, and L proteins together provide the polymerase activity responsible for genome transcription and replication. The viral genome is surrounded by a host cell-derived lipid bilayer envelope containing the M, F, SH, and HN proteins. The M protein is involved in viral assembly, whereas the HN and F proteins are responsible for cell attachment and entry and are the major targets of virus-neutralizing antibody. The V and SH proteins are accessory proteins, acting as antagonists of the host antiviral response; the former interferes with the interferon response and the latter with the tumor necrosis factor α (TNF- α)-mediated apoptotic signaling pathway. Because of the hypervariability of the SH gene, its nucleotide sequence is used to “genotype” the virus for molecular epidemiologic purposes. Thus far, 12 mumps virus genotypes have been assigned by SH gene sequence and are designated A–N (with the exclusion of E and M, which have been merged with genotypes C and K, respectively).

Nucleotide sequencing of clinical isolates shows that virus genotypes D and G circulate predominantly in the Western Hemisphere; genotypes F, C, and I in the Asia-Pacific region; and genotypes B, H, J, and K in the Southern Hemisphere (Fig. 231e-1). Although numerous mumps virus genotypes have been identified and some vary antigenically from others, only one serotype exists, and there is no evidence to suggest that certain circulating virus strains are more virulent or contagious than others.

EPIDEMIOLOGY



Mumps is endemic worldwide, with epidemics every 3–5 years in unvaccinated populations. These epidemics typically occur in locations where children and young adults congregate, such as schools, military barracks, and other institutions. In countries without national mumps vaccination programs, the estimated annual global incidence is 100–1000 cases per 100,000 population. After the introduction of mumps vaccine in the United States in 1967, the number of reported cases declined dramatically. By 2001, fewer than 300 cases were reported, representing a 99.8% reduction from prevaccine-era levels. Mumps incidence remained at historic lows in the United States until 2006, when 6584 cases were reported—the largest outbreak since 1987. At the time of the 2006 outbreak, the disease was resurging globally, even in populations with high-level vaccination coverage. The number of reported U.S. cases declined precipitously in the 2 years that followed but then spiked in 2009–2010, with focal outbreaks in New York and New Jersey, and again in 2011, with a focal outbreak in California. A recent study by the Centers for Disease Control and Prevention (CDC) showed that two-dose coverage with measles-mumps-rubella (MMR) vaccine in major U.S. cities (94.8%) remains at or very near the level needed to contain these childhood infections; however, focal areas with inadequate vaccination coverage still leave some children at risk. Sporadic, large-scale mumps outbreaks continue to be reported worldwide, sometimes in countries where the disease was once under control.

Although historically a disease of unvaccinated children, with the largest proportion of cases occurring in children 5–9 years of age during the prevaccine era, mumps now frequently occurs in older age groups—primarily college students, most of whom were vaccinated in early childhood. This shift in age distribution and the occurrence of mumps in vaccinated populations are probably the result of several coincident circumstances, including (1) situations promoting the spread of respiratory viruses among young adults (e.g., residence in college dormitories), (2) waning of vaccine immunity with time, (3) lack of endemically circulating wild-type virus to periodically boost vaccine-induced immune responses, and (4) continuing global epidemics (due to either lack of mumps vaccination programs or, where such programs do exist, low rates of mumps vaccination). The notable

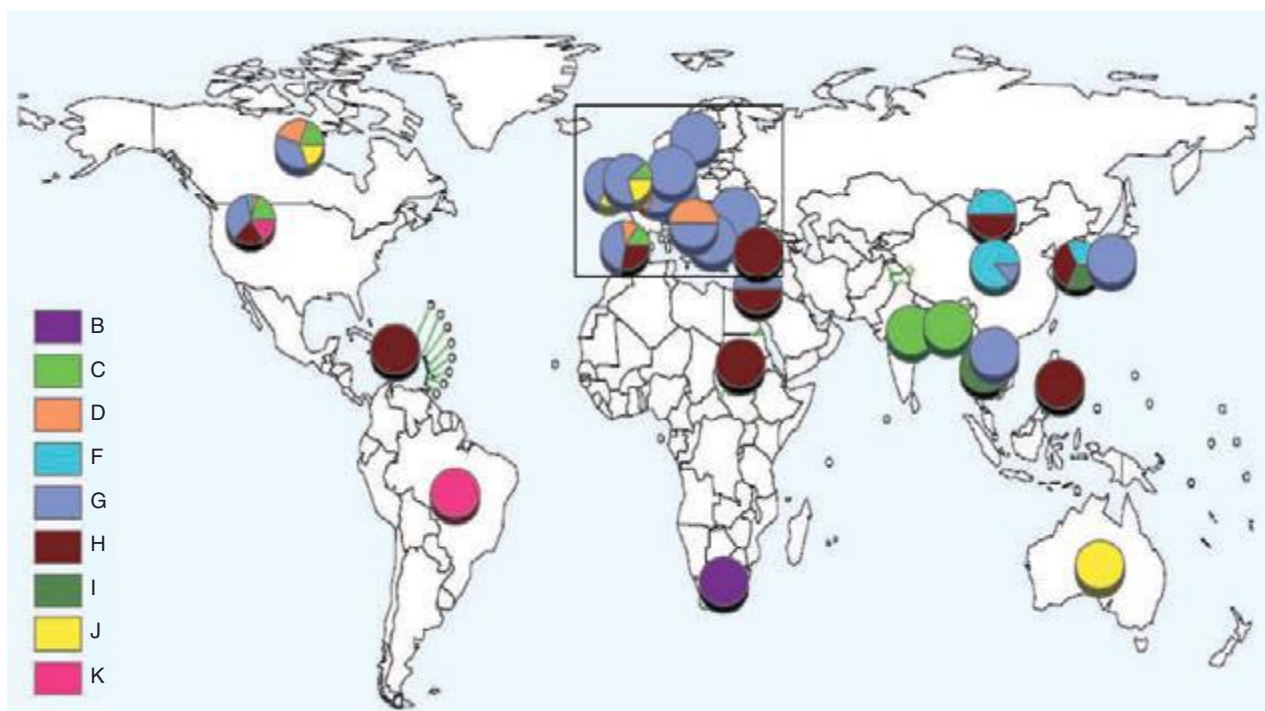


FIGURE 231e-1 Distribution of reported mumps genotypes, 2005–2011 (data as of April 20, 2012). Pie-slice size is proportional to the number of years each genotype was reported. (Figure courtesy of WHO, with permission; http://www.who.int/immunization_monitoring/diseases/mumps/en/index.html; accessed September 11, 2012.)

231e-2 decline in mumps vaccination–induced immunity with time may be due to both declining titers and decreasing avidity of antibodies. The waning of mumps immunity over time is supported by studies suggesting that a third dose of MMR vaccine significantly reduces the mumps attack rate; however, these studies were not adequately controlled to rule out the possibility that the observed declines in mumps incidence were unrelated to the intervention. Therefore, the effectiveness of a third dose of MMR vaccine remains to be demonstrated.

PATHOGENESIS

Humans are the only natural hosts for mumps virus infection. The incubation period of mumps is ~19 days (range, 7–23 days). The virus is transmitted by the respiratory route via droplets, saliva, and fomites. Mumps virus is typically shed from 1 week before to 1 week after symptom onset, although this window appears to be narrower in vaccinated individuals. Persons are most contagious 1–2 days before onset of clinical symptoms. Inference from related respiratory diseases and animal studies indicates that primary replication likely occurs in the nasal mucosa or upper respiratory mucosal epithelium. Mononuclear cells and cells within regional lymph nodes can become infected; such infection facilitates the development of viremia and poses a risk for a wide array of acute inflammatory reactions. Classic sites of mumps virus replication include the salivary glands, testes, pancreas, ovaries, mammary glands, and central nervous system (CNS).

Little is known about the pathology of mumps since the disease is rarely fatal. The virus replicates well in glandular epithelium, but classic parotitis is not a necessary component of mumps infection. Affected glands contain perivascular and interstitial mononuclear cell infiltrates and exhibit hemorrhage with prominent edema. Necrosis of acinar and epithelial duct cells is evident in the salivary glands and in the germinal epithelium of the seminiferous tubules of the testes. The virus probably enters cerebrospinal fluid (CSF) through the choroid plexus or via transiting mononuclear cells during plasma viremia. Although relevant data are limited, typical mumps encephalitis appears to be secondary to respiratory spread and is probably a parainfectious process, as suggested by perivenous demyelination, perivascular mononuclear cell inflammation, and relative sparing of neurons. Although rare, presumed primary encephalitis has been associated with mumps virus isolation from brain tissue. Evidence of placental and intrauterine spread in pregnancy has been found in both early and late gestation.

CLINICAL MANIFESTATIONS

Up to half of mumps virus infections are asymptomatic or lead to nonspecific respiratory symptoms. Inapparent infections are more common in adults than in children. The prodrome of mumps consists of low-grade fever, malaise, myalgia, headache, and anorexia. Mumps parotitis—acute-onset unilateral or bilateral swelling of the parotid or other salivary glands that lasts >2 days and has no other apparent cause—develops in 70–90% of symptomatic infections, usually within 24 h of prodromal symptoms but sometimes as long as 1 week thereafter. Parotitis is generally bilateral, although the two sides may not be involved synchronously. Unilateral involvement is documented in about one-third of cases. Swelling of the parotid is accompanied by tenderness and obliteration of the space between the earlobe and the angle of the mandible (Figs. 231e-2 and 231e-3). The patient frequently reports an earache and finds it difficult to eat, swallow, or talk. The orifice of the parotid duct is commonly red and swollen. The submaxillary and sublingual glands are involved less often than the parotid gland and are almost never involved alone. Glandular swelling increases for a few days and then gradually subsides, disappearing within 1 week. Recurrent sialadenitis is a rare sequela of mumps parotitis. In ~6% of mumps cases, obstruction of lymphatic drainage secondary to bilateral salivary gland swelling may lead to presternal pitting edema, associated often with submandibular adenitis and rarely with the more life-threatening supraglottic edema.



FIGURE 231e-2 A comparison of a person before acquiring mumps (A) and on day 3 (B) of acute bilateral parotitis. (Courtesy of patient C.M. From Shanley JD. The resurgence of mumps in young adults and adolescents. *Cleve Clin J Med* 2007; 74:42-48. Reprinted with permission. Copyright © 2007 Cleveland Clinic Foundation. All rights reserved.)

Epididymo-orchitis is the next most common manifestation of mumps, developing in 15–30% of cases in postpubertal males, with bilateral involvement in 10–30% of those cases. Orchitis, accompanied by fever, typically occurs during the first week of parotitis but can develop up to 6 weeks after parotitis or in its absence. The testis is painful and tender and can be enlarged to several times its normal size; this condition usually resolves within 1 week. Testicular atrophy develops in one-half of affected men. Sterility after mumps is rare, although subfertility is estimated to occur in 13% of cases of unilateral orchitis and in 30–87% of cases of bilateral orchitis. Oophoritis occurs in ~5% of women with mumps and may be associated with lower abdominal pain and vomiting but has only rarely been associated with sterility or premature menopause. Mumps infection in postpubertal women may also present with mastitis.

Documented CSF pleocytosis indicates that mumps virus invades the CNS in ~50% of cases; however, symptomatic CNS disease, typically in the form of aseptic meningitis, occurs in <10% of cases, with a male predominance. CNS symptoms of aseptic meningitis (e.g., stiff neck, headache, and drowsiness) appear ~5 days after parotitis and also occur often in the absence of parotid involvement. Within the first 24 h polymorphonuclear leukocytes may predominate in CSF (1000–2000 cells/ μ L), but by the second day nearly all the cells are lymphocytes. The glucose level in CSF may be low and the protein

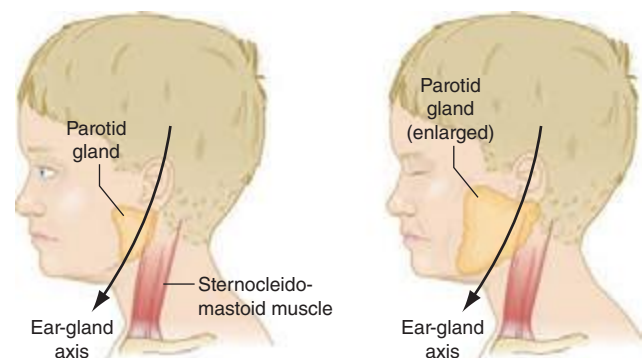


FIGURE 231e-3 Schematic drawing of a parotid gland infected with mumps virus (right) compared with a normal gland (left). An enlarged cervical lymph node is usually posterior to the imaginary line. (Reprinted with permission from Gershon A et al: *Mumps*, in *Krugman's Infectious Diseases of Children*, 11th ed. Philadelphia, Elsevier, 2004, p 392.)

concentration high, a pattern reminiscent of bacterial meningitis. Mumps meningitis is a self-limited manifestation without significant risk of death or long-term sequelae. Cranial nerve palsies have occasionally led to permanent sequelae, particularly deafness. The reported incidence of mumps-associated hearing loss varies between 1 in 1000 and 1 in 100,000. In ~0.1% of infections, mumps virus may cause encephalitis, which presents as high fever with marked changes in the level of consciousness, seizures, and focal neurologic symptoms. Electroencephalographic abnormalities may be seen. Permanent sequelae are sometimes identified in survivors, and adult infections more commonly have poor outcomes than do pediatric infections. The mortality rate associated with mumps encephalitis is ~1.5%. Other CNS problems occasionally associated with mumps include cerebellar ataxia, facial palsy, transverse myelitis, hydrocephalus, Guillain-Barré syndrome, flaccid paralysis, and behavioral changes.

Mumps pancreatitis, which may present as abdominal pain, occurs in ~4% of infections but is difficult to diagnose because an elevated serum amylase level can be associated with either parotitis or pancreatitis. An etiologic association of mumps virus and juvenile diabetes mellitus remains controversial. Myocarditis and endocardial fibroelastosis are rare and self-limited but may represent severe complications of mumps infection; however, mumps-associated electrocardiographic abnormalities have been reported in up to 15% of cases. Other unusual complications include thyroiditis, nephritis, arthritis, hepatic disease, keratouveitis, and thrombocytopenic purpura. Abnormal renal function is common, but severe, life-threatening nephritis is rare. It remains at issue whether an excessive number of spontaneous abortions are associated with gestational mumps. Mumps in pregnancy does not appear to lead to premature birth, low birth weight, or fetal malformations.

DIFFERENTIAL DIAGNOSIS

During a mumps outbreak, the diagnosis is made easily in patients with parotitis and a history of recent exposure; however, when disease incidence is low, other causes of parotitis should be considered and laboratory testing is required for case confirmation. Infectious causes of parotitis include other viruses (e.g., HIV, coxsackievirus, parainfluenza virus type 3, influenza A virus, Epstein-Barr virus, adenovirus, parvovirus B19, lymphocytic choriomeningitis virus, human herpesvirus 6), gram-positive bacteria, atypical mycobacteria, and *Bartonella* species. Rarely, other gram-negative or anaerobic bacteria are associated with parotitis. Parotitis can also develop in the setting of sarcoidosis, Sjögren's syndrome, Mikulicz's syndrome, Parinaud's syndrome, uremia, diabetes mellitus, laundry starch ingestion, malnutrition, cirrhosis, and some drug treatments. Unilateral parotitis can be caused by ductal obstruction, cysts, and tumors. In the absence of parotitis or other salivary gland enlargement, symptoms of other visceral organ and/or CNS involvement may predominate, and a laboratory diagnosis is required. Other entities should be considered when manifestations consistent with mumps appear in organs other than the parotid. Testicular torsion may produce a painful scrotal mass resembling that seen in mumps orchitis. Other viruses (e.g., enteroviruses) may cause aseptic meningitis that is clinically indistinguishable from that due to mumps virus.

LABORATORY DIAGNOSIS

Laboratory diagnosis is primarily based on detection of viral RNA by reverse-transcriptase polymerase chain reaction (RT-PCR) or on serology. Detection of viral antigens (e.g., via mumps virus-specific immunofluorescent staining of cultured clinical specimens) is comparatively inefficient and is no longer commonly performed.

For RT-PCR-based testing, viral RNA can be extracted either directly from clinical samples or from cell cultures incubated with clinical samples. Buccal swabs appear to be the best specimens for virus detection, particularly when obtained within 2 days of clinical onset; however, mumps virus can also be detected readily in throat swabs and saliva and, in cases of meningitis, in CSF. Despite the apparent high frequency of viremia during mumps, mumps virus has rarely been detected in blood. The ability to detect viral RNA in clinical samples

rapidly diminishes beyond the first week after symptom onset, and in several studies rates of virus detection were substantially lower in recipients of two vaccine doses than in unvaccinated persons or recipients of one dose. The rate of false-negative RT-PCR findings can be quite high, approaching 70% in some studies.

A serologic diagnosis is typically made by enzyme-linked immunosorbent assay (ELISA). The data must be interpreted with caution. In vaccinated persons with mumps, IgM is typically absent; thus, a negative IgM result in a vaccinated person does not rule out mumps. In addition, regardless of vaccination status, IgM may not be detectable if serum is assayed too early (prior to day 3 of symptom onset) or too late (beyond 6 weeks after symptom onset) in the course of disease. Reliance on a rise in IgG titer in paired acute- and convalescent-phase sera also is problematic: IgG titers in convalescent-phase sera may be only nominally greater than those in acute-phase sera. Thus, at present, the capacity of RNA or viral antigen detection to confirm cases is much greater than that of serologic testing. Traditional and labor-intensive serologic tests such as complement fixation, hemagglutination inhibition, and virus neutralization are now performed only rarely. The main downside to replacement of these functional serologic assays with the more rapid ELISA method is the latter's detection of all virus-specific antibodies, including those that are nonneutralizing (i.e., nonprotective). Thus, an individual who is seropositive by ELISA may lack protective levels of antibody. While there is a strong association between the presence of mumps virus neutralizing antibody and protection from disease, an absolute antibody titer predictive of serologic protection is lacking; in this respect, mumps differs from other respiratory infections, such as measles.

TREATMENT MUMPS

Mumps is generally a benign, self-resolving illness. Therapy for parotitis and other clinical manifestations is symptom based and supportive. The administration of analgesics and the application of warm or cold compresses to the parotid area may be helpful. Testicular pain may be minimized by the local application of cold compresses and gentle support for the scrotum. Anesthetic blocks also may be used. Neither the administration of glucocorticoids nor incision of the tunica albuginea is of proven value in severe orchitis. Anecdotal information on a small number of patients with orchitis suggests that SC administration of interferon $\alpha 2b$ may help preserve the organ and fertility. Lumbar puncture is occasionally performed to relieve headache associated with meningitis. Mumps immune globulin has not been consistently shown to be effective in preventing mumps and is not recommended for treatment or postexposure prophylaxis.

PREVENTION



Vaccination is the only practical control measure. Nearly all developed countries use mumps-containing vaccines, but in many countries mumps is not a notifiable disease and vaccination is often voluntary. However, where used, mumps vaccination has had a tremendous impact, with reductions in incidence and morbidity typically exceeding 90%. Despite the tremendous success of mumps vaccination programs, large mumps outbreaks continue to occur globally, even in settings of high-level two-dose vaccine coverage. Whereas outbreaks historically involved young (often unvaccinated) children in primary and secondary schools, more recent outbreaks have predominantly involved young adults, particularly on college and university campuses. While primary and secondary (waning-immunity) vaccine failures have been hypothesized to be factors in mumps outbreaks in several countries, in some countries other factors may have played a role, such as lack of compliance with the recommended vaccine schedule, changes to vaccination schedules resulting in missed cohorts, or changes in population demographics, such as large-scale immigration.

In the United States, the benefit-cost ratios for mumps vaccination alone are >13 for direct costs (e.g., medical expenses) and >24 for societal costs (including productivity losses for patients and caregivers).

231e-4 Several mumps virus vaccines are used throughout the world; in the United States, only the live attenuated Jeryl Lynn strain is used. Current recommendations are that mumps vaccine be administered as part of the combined trivalent measles-mumps-rubella vaccine (M-M-R® II) or the quadrivalent measles-mumps-rubella-varicella vaccine (ProQuad®). Monovalent vaccine is no longer produced for the U.S. market but is available in other countries.

Before administering mumps-containing vaccine, physicians should always consult the latest recommendations from the Advisory Committee on Immunization Practices (ACIP). Current recommendations for children specify two doses of mumps-containing vaccine: the first dose given on or after the first birthday and the second dose administered no earlier than 28 days after the first. In the United States, children often receive the second dose between the ages of 4 and 6 years.

In 2009, the ACIP revised its recommendations for evidence of mumps immunity in health care personnel to include (1) documented administration of two doses of a preparation containing live mumps vaccine, (2) laboratory evidence of immunity or laboratory confirmation of disease, or (3) birth date before 1957. For unvaccinated health care personnel born before 1957 who lack laboratory evidence of mumps immunity or laboratory confirmation of mumps, health care facilities should consider two doses of MMR vaccine separated by the appropriate interval; during a mumps outbreak, vaccination of these individuals is recommended.

Mumps vaccine contains live attenuated virus. It is not recommended for pregnant women, for individuals who have had a

life-threatening allergic reaction to components of the vaccine, or for people in settings of clinically significant primary or secondary immunosuppression. (For details, see the ACIP guidelines on the CDC's website [www.cdc.gov/vaccines/pubs/acip-list.htm].) Occasionally, febrile reactions and parotitis have been reported soon after mumps vaccination. Allergic reactions after vaccination (e.g., rash and pruritus) are uncommon and are usually mild and self-limited. More serious complications, such as aseptic meningitis, have been causally associated with certain vaccine strains but not with the Jeryl Lynn strain.

Immunity to mumps is associated with the development of neutralizing antibody, although a specific correlate of protection has not been established. Seroconversion occurs in ~95% of recipients of the Jeryl Lynn strain; however, vaccine efficacy is ~80% for one dose and 90% for two doses. Recent data indicate declining seropositivity rates with time since vaccination. Studies are under way to assess the value of including a third dose in the immunization schedule. Although it is generally accepted that mumps virus is serologically monotypic, antigenic differences between virus isolates have been detected. It is unclear whether such differences can lead to immune escape. The role of the cellular arm of the immune response is unclear, but there is evidence that it may help limit virus spread and complications.

ACKNOWLEDGMENT

The authors thank and acknowledge Dr. Anne Gershon, the author of this chapter in earlier editions.

232 Rabies and Other Rhabdovirus Infections

Alan C. Jackson

RABIES

Rabies is a rapidly progressive, acute infectious disease of the central nervous system (CNS) in humans and animals that is caused by infection with rabies virus. The infection is normally transmitted from animal vectors. Rabies has encephalitic and paralytic forms that progress to death.

ETIOLOGIC AGENT

Rabies virus is a member of the family Rhabdoviridae. Two genera in this family, *Lyssavirus* and *Vesiculovirus*, contain species that cause human disease. Rabies virus is a lyssavirus that infects a broad range of animals and causes serious neurologic disease when transmitted to humans. This single-strand RNA virus has a nonsegmented, negative-sense (antisense) genome that consists of 11,932 nucleotides and encodes 5 proteins: nucleocapsid protein, phosphoprotein, matrix protein, glycoprotein, and a large polymerase protein. Rabies virus variants, which can be characterized by distinctive nucleotide sequences, are associated with specific animal reservoirs. Six other non-rabies virus species in the *Lyssavirus* genus have been reported to cause a clinical picture similar to rabies. Vesicular stomatitis virus, a vesiculovirus, causes vesiculation and ulceration in cattle, horses, and other animals and causes a self-limited, mild, systemic illness in humans (see “Other Rhabdoviruses,” below).



Rabies is a zoonotic infection that occurs in a variety of mammals throughout the world except in Antarctica and on some islands. Rabies virus is usually transmitted to humans by the bite of an infected animal. Worldwide, endemic canine rabies is estimated to cause 55,000 human deaths annually. Most of these deaths occur in Asia and Africa, with rural populations and children most frequently affected. Thus, in many resource-poor and resource-limited countries, canine rabies continues to be a threat to humans. However, in Latin America, rabies control efforts in dogs have been quite successful in recent years. Endemic canine rabies has been eliminated from the United States and most other resource-rich countries. Rabies is endemic in wildlife species, and a variety of animal reservoirs have been identified in different countries. Surveillance data from 2012 identified 6162 confirmed animal cases of rabies in the United States (including Puerto Rico). Only 8% of these cases were in domestic animals, including 257 cases in cats, 84 in dogs, and 115 in cattle. In North American wildlife reservoirs, including bats, raccoons, skunks, and foxes, the infection is endemic, with involvement of one or more rabies virus variants in each reservoir species (Fig. 232-1). “Spillover” of rabies to other wildlife species and to domestic animals occurs. Bat rabies virus variants are present in every state except Hawaii and are responsible for most indigenously acquired human rabies cases in the United States. Raccoon rabies is endemic along the entire eastern coast of the United States. Skunk rabies is present in the midwestern states, with another focus in California. Rabies in foxes occurs in Texas, New Mexico, Arizona, and Alaska.

In Canada and Europe, epizootics of rabies in red foxes have been well controlled with the use of baits containing rabies vaccine. A similar approach is used in Canada to control raccoon rabies.

Rabies virus variants isolated from humans or other mammalian species can be identified by reverse-transcription polymerase chain reaction (RT-PCR) amplification and sequencing or by characterization with monoclonal antibodies. These techniques are helpful in human cases with no known history of exposure. Worldwide, most human rabies is transmitted from dogs in countries with endemic canine rabies and dog-to-dog transmission, and human cases can be imported

by travelers returning from these regions. In North America, human disease is usually associated with transmission from bats; there may be no known history of bat bite or other bat exposure in these cases. Most human cases are due to a bat rabies virus variant associated with silver-haired and tricolored bats. These are small bats whose bite may not be recognized, and the virus has adapted for replication at skin temperature and in cell types that are present in the skin.

Transmission from nonbite exposures is relatively uncommon. Aerosols generated in the laboratory or in caves containing millions of Brazilian free-tail bats have rarely caused human rabies. Transmission has resulted from corneal transplantation and also from solid organ transplantation and a vascular conduit (for a liver transplant) from undiagnosed donors with rabies in Texas, Florida, and Germany. Human-to-human transmission is extremely rare, although hypothetical concern about transmission to health care workers has prompted the implementation of barrier techniques to prevent exposures.

PATHOGENESIS

The incubation period of rabies (defined as the interval between exposure and the onset of clinical disease) is usually 20–90 days but in rare cases is as short as a few days or >1 year. During most of the incubation period, rabies virus is thought to be present at or close to the site of inoculation (Fig. 232-2). In muscles, the virus is known to bind to nicotinic acetylcholine receptors on postsynaptic membranes at neuromuscular junctions, but the exact details of viral entry into the skin and SC tissues have not yet been clarified. Rabies virus spreads centripetally along peripheral nerves toward the CNS at a rate of up to ~250 mm/d via retrograde fast axonal transport to the spinal cord or brainstem. Once the virus enters the CNS, it rapidly disseminates to other regions of the CNS via fast axonal transport along neuroanatomic connections. Neurons are prominently infected in rabies; infection of astrocytes is unusual. After CNS infection becomes established, there is centrifugal spread along sensory and autonomic nerves to other tissues, including the salivary glands, heart, adrenal glands, and skin. Rabies virus replicates in acinar cells of the salivary glands and is secreted in the saliva of rabid animals that serve as vectors of

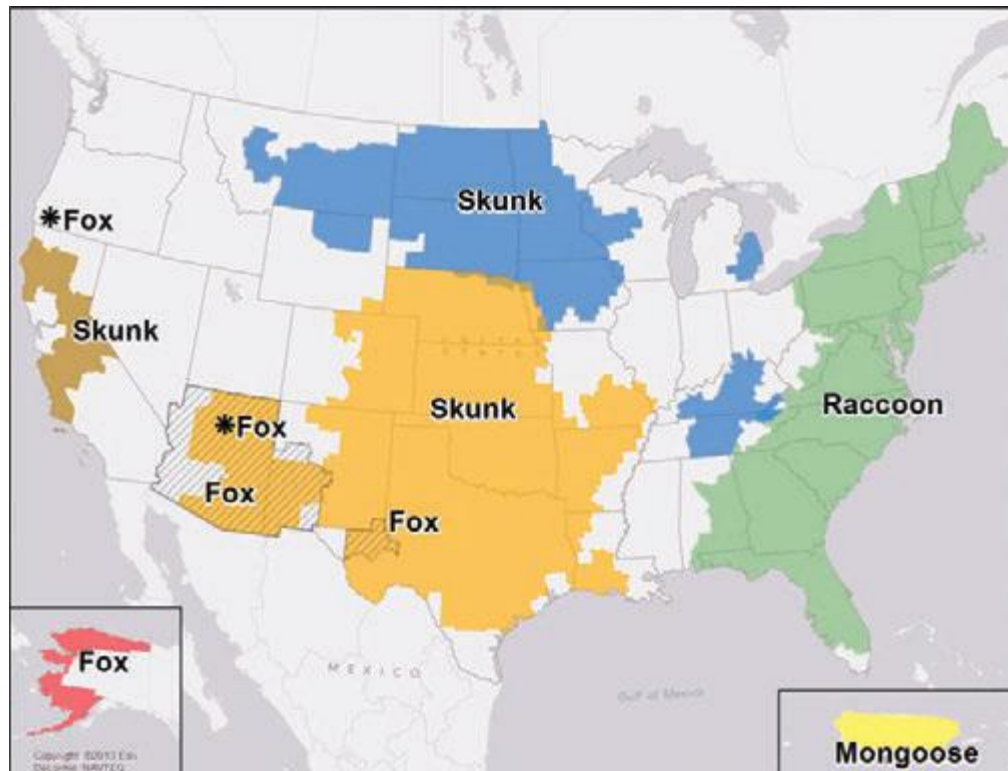


FIGURE 232-1 Distribution of the major rabies virus variants among wild terrestrial reservoirs in the United States and Puerto Rico, 2008–2012. *Potential host-shift event. (From JL Dyer et al: *J Am Vet Med Assoc* 243:805, 2013.)

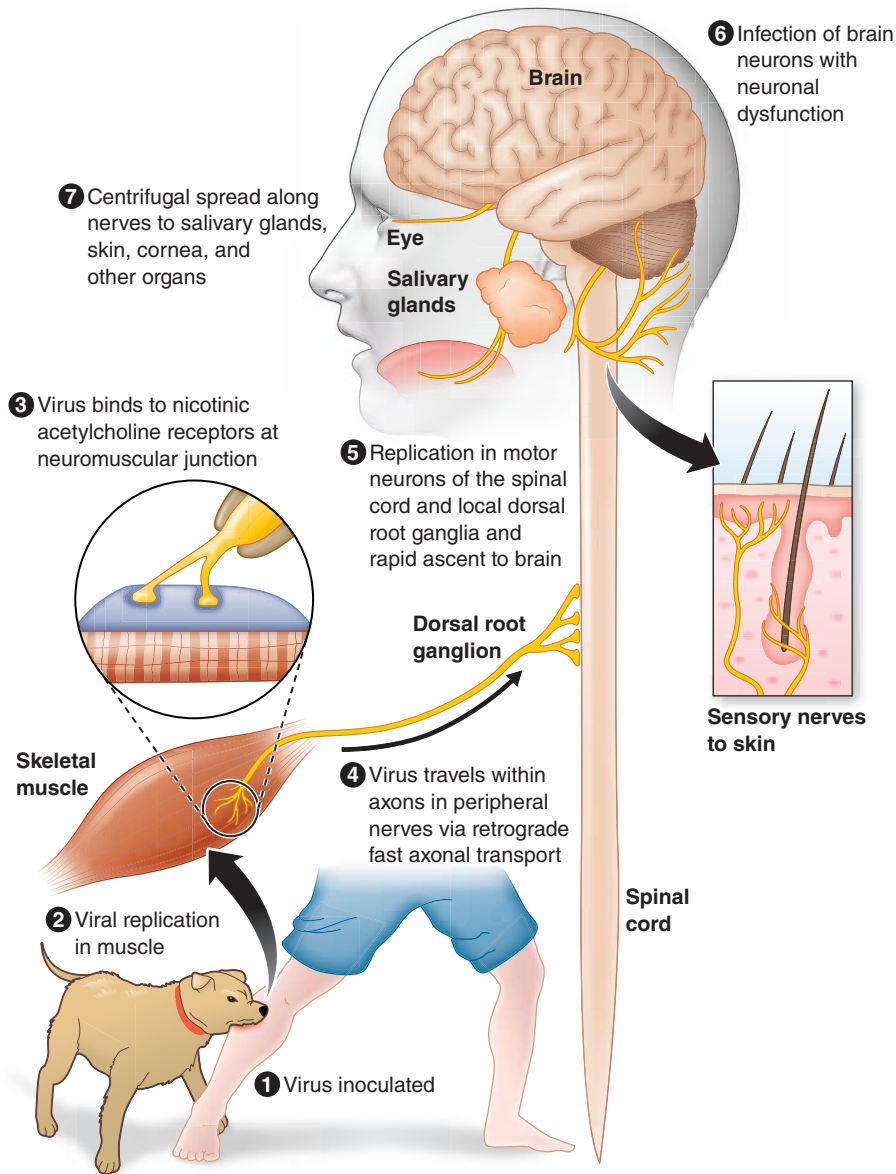


FIGURE 232-2 Schematic representation of the pathogenetic events following peripheral inoculation of rabies virus by an animal bite. (Adapted from Jackson AC: *Human disease, in Rabies: scientific basis of the disease and its management*, 3rd ed., AC Jackson [ed], Oxford, UK, Elsevier Academic Press, 2013, pp 269–298; with permission.)

the disease. There is no well-documented evidence for hematogenous spread of rabies virus.

Pathologic studies show mild inflammatory changes in the CNS in rabies, with mononuclear inflammatory infiltration in the leptomeninges, perivascular regions, and parenchyma, including microglial nodules called *Babes nodules*. Degenerative neuronal changes usually are not prominent, and there is little evidence of neuronal death; neuronophagia is observed occasionally. The pathologic changes are surprisingly mild in light of the clinical severity and fatal outcome of the disease. The most characteristic pathologic finding in rabies is the *Negri body* (Fig. 232-3). Negri bodies are eosinophilic cytoplasmic inclusions in brain neurons that are composed of rabies virus proteins and viral RNA. These inclusions occur in a minority of infected neurons, are commonly observed in Purkinje cells of the cerebellum and in pyramidal neurons of the hippocampus, and are less frequently seen in cortical and brainstem neurons. Negri bodies are not observed in all cases of rabies. The lack of prominent degenerative neuronal changes has led to the concept that neuronal dysfunction—rather than neuronal death—is responsible for clinical disease in rabies. The basis for behavioral changes, including the aggressive behavior of rabid animals, is not well understood.

CLINICAL MANIFESTATIONS

In rabies, the emphasis must be on postexposure prophylaxis (PEP) initiated before any symptoms or signs develop. Rabies should usually be suspected on the basis of the clinical presentation. The disease usually presents as atypical encephalitis with relative preservation of consciousness. Rabies may be difficult to recognize late in the clinical course when progression to coma has occurred. A minority of patients present with acute flaccid paralysis. There are prodromal, acute neurologic, and comatose phases that usually progress to death despite aggressive therapy (Table 232-1).

Prodromal Features The clinical features of rabies begin with nonspecific prodromal manifestations, including fever, malaise, headache, nausea, and vomiting. Anxiety or agitation may also occur. The earliest specific neurologic symptoms of rabies include paresthesias, pain, or pruritus near the site of the exposure, one or more of which occur in 50–80% of patients and strongly suggest rabies. The wound has usually healed by this point, and these symptoms probably reflect infection with associated inflammatory changes in local dorsal root or cranial sensory ganglia.

Encephalitic Rabies Two acute neurologic forms of rabies are seen in humans: encephalitic (furious) in 80% and paralytic in 20%. Some of the manifestations of encephalitic rabies, including fever, confusion, hallucinations, combativeness, and seizures, may be seen in other viral encephalitides as well. Autonomic dysfunction is common and may result in hypersalivation, gooseflesh, cardiac arrhythmia, and priapism. In encephalitic rabies, episodes of hyperexcitability are typically followed by periods of complete lucidity that become shorter as the disease progresses. Rabies encephalitis is distinguished by early brainstem involvement, which results in the classic features of *hydrophobia* (involuntary, painful contraction of the diaphragm and accessory respiratory, laryngeal, and pharyngeal muscles in response to swallowing liquids) and *aerophobia* (the same features caused by stimulation from a draft of air). These symptoms are probably due to dysfunction of infected brainstem neurons that normally inhibit inspiratory neurons near the nucleus ambiguus, resulting in exaggerated defense reflexes that protect the respiratory tract. The combination of hypersalivation and pharyngeal dysfunction is also responsible for the classic appearance of “foaming at the mouth” (Fig. 232-4). Brainstem dysfunction progresses rapidly, and coma—followed within days by death—is the rule unless the course is prolonged by supportive measures. With such measures, late complications can include cardiac and/or respiratory failure, disturbances of water balance (syndrome of inappropriate antidiuretic hormone secretion or diabetes insipidus), noncardiogenic pulmonary edema, and gastrointestinal hemorrhage. Cardiac arrhythmias may be due to dysfunction affecting vital centers in the brainstem or to myocarditis. Multiple-organ failure is common in patients treated aggressively in critical care units.

Paralytic Rabies About 20% of patients have paralytic rabies in which muscle weakness predominates and cardinal features of encephalitic rabies (hyperexcitability, hydrophobia, and aerophobia) are lacking. There is early and prominent flaccid muscle weakness, often beginning in the bitten extremity and spreading to produce quadriplegia and facial

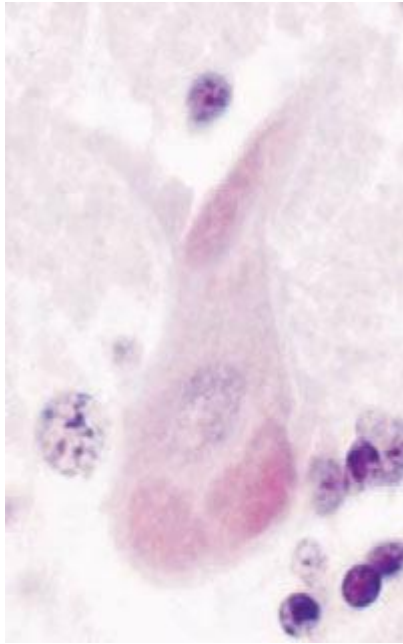


FIGURE 232-3 Three large Negri bodies in the cytoplasm of a cerebellar Purkinje cell from an 8-year-old boy who died of rabies after being bitten by a rabid dog in Mexico. (From AC Jackson, E Lopez-Corella: *N Engl J Med* 335:568, 1996. © Massachusetts Medical Society.)

weakness. Sphincter involvement is common, sensory involvement is usually mild, and these cases are commonly misdiagnosed as Guillain-Barré syndrome. Patients with paralytic rabies generally survive a few days longer than those with encephalitic rabies, but multiple-organ failure nevertheless ensues.

LABORATORY INVESTIGATIONS

Most routine laboratory tests in rabies yield normal results or show nonspecific abnormalities. Complete blood counts are usually normal. Examination of cerebrospinal fluid (CSF) often reveals mild mononuclear cell pleocytosis with a mildly elevated protein level. Severe pleocytosis (>1000 white cells/ μ L) is unusual and should prompt a search for an alternative diagnosis. CT head scans are usually normal in rabies. MRI brain scans may show signal abnormalities in the brainstem or other gray-matter areas, but these findings are variable and nonspecific. Electroencephalograms show only nonspecific abnormalities. Of course, important tests in suspected cases of rabies include those that may identify an alternative, potentially treatable diagnosis (see “Differential Diagnosis,” below).

TABLE 232-1 CLINICAL STAGES OF RABIES		
Phase	Typical Duration	Symptoms and Signs
Incubation period	20–90 days	None
Prodrome	2–10 days	Fever, malaise, anorexia, nausea, vomiting; paresthesias, pain, or pruritus at the wound site
Acute neurologic disease		
Encephalitic (80%)	2–7 days	Anxiety, agitation, hyperactivity, bizarre behavior, hallucinations, autonomic dysfunction, hydrophobia
Paralytic (20%)	2–10 days	Flaccid paralysis in limb(s) progressing to quadriplegia with facial paralysis
Coma, death ^a	0–14 days	

^aRecovery is rare.

Source: MAW Hattwick: Rabies virus, in *Principles and Practice of Infectious Diseases*, GL Mandell et al (eds). New York, Wiley, 1979, pp 1217–1228. Adapted with permission from Elsevier.



FIGURE 232-4 Hydrophobic spasm of inspiratory muscles associated with terror in a patient with encephalitic (furious) rabies who is attempting to swallow water. (Copyright DA Warrell, Oxford, UK; with permission.)

DIAGNOSIS

In North America, a diagnosis of rabies often is not considered until relatively late in the clinical course, even with a typical clinical presentation. This diagnosis should be considered in patients presenting with acute atypical encephalitis or acute flaccid paralysis, including those in whom Guillain-Barré syndrome is suspected. The absence of an animal-bite history is common in North America. The lack of hydrophobia is not unusual in rabies. Once rabies is suspected, rabies-specific laboratory tests should be performed to confirm the diagnosis. Diagnostically useful specimens include serum, CSF, fresh saliva, skin biopsy samples from the neck, and brain tissue (rarely obtained before death). Because skin biopsy relies on the demonstration of rabies virus antigen in cutaneous nerves at the base of hair follicles, samples are usually taken from hairy skin at the nape of the neck. Corneal impression smears are of low diagnostic yield and are generally not performed. Negative antemortem rabies-specific laboratory tests never exclude a diagnosis of rabies, and tests may need to be repeated after an interval for diagnostic confirmation.

Rabies Virus–Specific Antibodies In a previously unimmunized patient, serum neutralizing antibodies to rabies virus are diagnostic. However, because rabies virus infects immunologically privileged neuronal tissues, serum antibodies may not develop until late in the disease. Antibodies may be detected within a few days after the onset of symptoms, but some patients die without detectable antibodies. The presence of rabies virus–specific antibodies in the CSF suggests rabies encephalitis, regardless of immunization status. A diagnosis of rabies is questionable in patients who recover from rabies without developing serum neutralizing antibodies to rabies virus.

RT-PCR Amplification Detection of rabies virus RNA by RT-PCR is highly sensitive and specific. This technique can detect virus in fresh saliva samples, skin, CSF, and brain tissues. In addition, RT-PCR with genetic sequencing can distinguish among rabies virus variants, permitting identification of the probable source of an infection.

Direct Fluorescent Antibody Testing Direct fluorescent antibody (DFA) testing with rabies virus antibodies conjugated to fluorescent dyes is highly sensitive and specific and can be performed quickly and applied to skin biopsies and brain tissue. In skin biopsies, rabies virus antigen may be detected in cutaneous nerves at the base of hair follicles.

DIFFERENTIAL DIAGNOSIS

The diagnosis of rabies may be difficult without a history of animal exposure, and no exposure to an animal (e.g., a bat) may be recalled. The presentation of rabies is usually quite different from that of acute viral encephalitis due to most other causes, including herpes simplex encephalitis and arboviral (e.g., West Nile) encephalitis. Early neurologic symptoms may occur at the site of the bite, and there may be early features of brainstem involvement with preservation of consciousness. Anti-*N*-methyl-*D*-aspartate receptor (anti-NMDA) encephalitis occurs in young patients (especially females) and is characterized by behavioral changes, autonomic instability, hypoventilation, and seizures. Postinfectious (immune-mediated) encephalomyelitis may follow influenza, measles, mumps, and other infections; it may also occur as a sequela of immunization with rabies vaccine derived from neural tissues, which are used only in resource-limited and resource-poor countries. Rabies may present with unusual neuropsychiatric symptoms and may be misdiagnosed as a psychiatric disorder. Rabies hysteria may occur as a psychological response to the fear of rabies and is often characterized by a shorter incubation period than rabies, aggressive behavior, inability to communicate, and a long course with recovery.

As previously mentioned, paralytic rabies may mimic Guillain-Barré syndrome. In these cases, fever, bladder dysfunction, a normal sensory examination, and CSF pleocytosis favor a diagnosis of rabies. Conversely, Guillain-Barré syndrome may occur as a complication of rabies vaccination with a neural tissue–derived product (e.g., suckling mouse brain vaccine) and may be mistaken for paralytic rabies (i.e., vaccine failure).

TREATMENT RABIES

There is no established treatment for rabies. There have been many recent treatment failures with the combination of antiviral drugs, ketamine, and therapeutic (induced) coma—measures that were used in a healthy survivor in whom antibodies to rabies virus were detected at presentation. Expert opinion should be sought before a course of experimental therapy is embarked upon. A palliative approach may be appropriate for some patients.

PROGNOSIS

Rabies is an almost uniformly fatal disease but is nearly always preventable with appropriate postexposure therapy during the early incubation period (see below). There are seven well-documented cases of survival from rabies. All but one of these patients had received rabies vaccine before disease onset. The single survivor who had not received vaccine had neutralizing antibodies to rabies virus in serum and CSF at clinical presentation. Most patients with rabies die within several days of the onset of illness, despite aggressive care in a critical care unit.

PREVENTION

Postexposure Prophylaxis Since there is no effective therapy for rabies, it is extremely important to prevent the disease after an animal exposure. **Figure 232-5** shows the steps involved in making decisions about PEP. On the basis of the history of the exposure and local epidemiologic information, the physician must decide whether initiation of PEP is warranted. Healthy dogs, cats, or ferrets may be confined and observed for 10 days. PEP is not necessary if the animal remains healthy. If the animal develops signs of rabies during the observation period, it should be euthanized immediately; the head should be transported to the laboratory under refrigeration, rabies virus should be sought by DFA testing, and viral isolation should be attempted by cell culture and/or mouse inoculation. Any animal other than a dog, cat, or ferret should be euthanized immediately and the head submitted for laboratory examination. In high-risk exposures and in areas where canine rabies is endemic, rabies prophylaxis should be initiated without waiting for laboratory results. If the laboratory results prove to be negative, it may safely be concluded that the animal's saliva did not

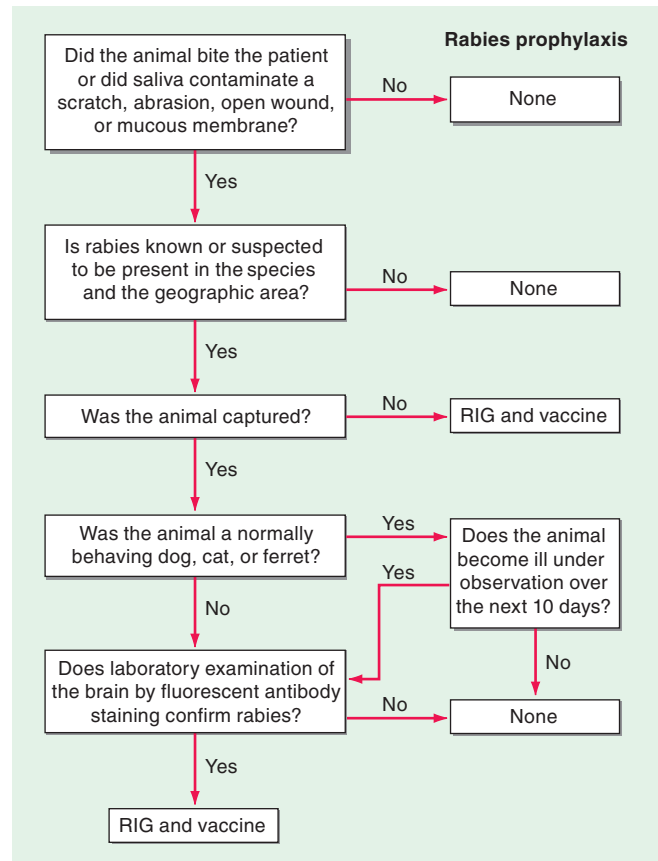


FIGURE 232-5 Algorithm for rabies postexposure prophylaxis. RIG, rabies immune globulin. (From L Corey, in *Harrison's Principles of Internal Medicine*, 15th ed. E Braunwald et al [eds]; New York, McGraw-Hill, 2001; adapted with permission.)

contain rabies virus, and immunization should be discontinued. If an animal escapes after an exposure, it must be considered rabid, and PEP must be initiated unless information from public health officials indicates otherwise (i.e., there is no endemic rabies in the area). Although controversial, the use of PEP may be warranted when a person (e.g., a small child or a sleeping adult) has been present in the same space as a bat and an unrecognized bite cannot be reliably excluded.

PEP includes local wound care and both active and passive immunization. Local wound care is essential and may greatly decrease the risk of rabies virus infection. Wound care should not be delayed, even if the initiation of immunization is postponed pending the results of the 10-day observation period. All bite wounds and scratches should be washed thoroughly with soap and water. Devitalized tissues should be debrided, tetanus prophylaxis given, and antibiotic treatment initiated whenever indicated.

All previously unvaccinated persons (but not those who have previously been immunized) should be passively immunized with rabies immune globulin (RIG). If RIG is not immediately available, it should be administered no later than 7 days after the first vaccine dose. After day 7, endogenous antibodies are being produced, and passive immunization may actually be counterproductive. If anatomically feasible, the entire dose of RIG (20 IU/kg) should be infiltrated at the site of the bite; otherwise, any RIG remaining after infiltration of the bite site should be administered IM at a distant site. With multiple or large wounds, the RIG preparation may need to be diluted in order to obtain a sufficient volume for adequate infiltration of all wound sites. If the exposure involves a mucous membrane, the entire dose should be administered IM. Rabies vaccine and RIG should never be administered at the same site or with the same syringe. Commercially available RIG in the United States is purified from the serum of hyperimmunized human donors. These human RIG preparations are much better tolerated than are the equine-derived preparations still in use in

1304 some countries (see below). Serious adverse effects of human RIG are uncommon. Local pain and low-grade fever may occur.

Two purified inactivated rabies vaccines are available for rabies PEP in the United States. They are highly immunogenic and remarkably safe compared with earlier vaccines. Four 1-mL doses of rabies vaccine should be given IM in the deltoid area. (The anterolateral aspect of the thigh also is acceptable in children.) Gluteal injections, which may not always reach muscle, should not be given and have been associated with rare vaccine failures. Ideally, the first dose should be given as soon as possible after exposure; failing that, it should be given without further delay. The three additional doses should be given on days 3, 7, and 14; a fifth dose on day 28 is no longer recommended. Pregnancy is not a contraindication for immunization. Glucocorticoids and other immunosuppressive medications may interfere with the development of active immunity and should not be administered during PEP unless they are essential. Routine measurement of serum neutralizing antibody titers is not required, but titers should be measured 2–4 weeks after immunization in immunocompromised persons. Local reactions (pain, erythema, edema, and pruritus) and mild systemic reactions (fever, myalgias, headache, and nausea) are common; anti-inflammatory and antipyretic medications may be used, but immunization should not be discontinued. Systemic allergic reactions are uncommon, but anaphylaxis does occur rarely and can be treated with epinephrine and antihistamines. The risk of rabies development should be carefully considered before the decision is made to discontinue vaccination because of an adverse reaction.



Most of the burden of rabies PEP is borne by persons with the fewest resources. In addition to the rabies vaccines discussed above, vaccines grown in either primary cell lines (hamster or dog kidney) or continuous cell lines (Vero cells) are satisfactory and are available in many countries outside the United States. Less expensive vaccines derived from neural tissues are still used in a diminishing number of developing countries; however, these vaccines are associated with serious neuroparalytic complications, including postinfectious encephalomyelitis and Guillain-Barré syndrome. The use of these vaccines should be discontinued as soon as possible, and progress has been made in this regard. Worldwide, >10 million individuals receive postexposure rabies vaccine each year.

If human RIG is unavailable, purified equine RIG can be used in the same manner at a dose of 40 IU/kg. Before the administration of equine RIG, hypersensitivity should be assessed by intradermal testing with a 1:10 dilution. The incidence of anaphylactic reactions and serum sickness has been low with recent equine RIG products.

Preexposure Rabies Vaccination Preexposure rabies prophylaxis should be considered for people with an occupational or recreational risk of rabies exposures, including certain travelers to rabies-endemic areas. The primary schedule consists of three doses of rabies vaccine given on days 0, 7, and 21 or 28. Serum neutralizing antibody tests help determine the need for subsequent booster doses. When a previously immunized individual is exposed to rabies, two booster doses of vaccine should be administered on days 0 and 3. Wound care remains essential. As stated above, RIG should not be administered to previously vaccinated persons.

OTHER RHABDOVIRUSES

OTHER LYSSAVIRUSES



A growing number of lyssaviruses other than rabies virus have been discovered to infect bat populations in Europe, Africa, Asia, and Australia. Six of these viruses have produced a very small number of cases of a human disease indistinguishable from rabies: European bat lyssaviruses 1 and 2, Australian bat lyssavirus, Irkut virus, and Duvenhage virus. Mokola virus, a lyssavirus that has been isolated from shrews with an unknown reservoir species in Africa, may also produce human disease indistinguishable from rabies.

VESICULAR STOMATITIS VIRUS

Vesicular stomatitis is a viral disease of cattle, horses, pigs, and some wild mammals. Vesicular stomatitis virus is a member of the genus

Vesiculovirus in the family Rhabdoviridae. Outbreaks of vesicular stomatitis in horses and cattle occur sporadically in the southwestern United States. The animal infection is associated with severe vesiculation and ulceration of oral tissues, teats, and feet and may be clinically indistinguishable from the more dangerous foot-and-mouth disease. Epidemics are usually seasonal, typically beginning in the late spring, and are probably due to arthropod vectors. Direct animal-to-animal spread can also occur, although the virus cannot penetrate intact skin. Transmission to humans usually results from direct contact with infected animals (particularly cattle) and occasionally follows laboratory exposure. In human disease, early conjunctivitis is followed by an acute influenza-like illness with fever, chills, nausea, vomiting, headache, retrobulbar pain, myalgias, substernal pain, malaise, pharyngitis, and lymphadenitis. Small vesicular lesions may be present on the buccal mucosa or on the fingers. Encephalitis is very rare. The illness usually lasts 3–6 days, with complete recovery. Subclinical infections are common. A serologic diagnosis can be made on the basis of a rise in titer of complement-fixing or neutralizing antibodies. Therapy is symptom-based.

233

Arthropod-Borne and Rodent-Borne Virus Infections

Jens H. Kuhn, Clarence J. Peters

This chapter summarizes the major features of selected arthropod-borne and rodent-borne viruses. Numerous viruses of this category are transmitted in nature among animals without ever infecting humans. Other viruses incidentally infect humans, but only a proportion of these viruses induce human disease. In addition, certain viral agents are regularly introduced into human populations or spread among humans by certain arthropods (specifically, insects and ticks) or by chronically infected rodents. These zoonotic viruses are taxonomically diverse and therefore differ fundamentally from one another in terms of virion morphology, replication strategies, genomic organization, and genome sequence. While a virus's classification in a taxon is enlightening with regard to natural maintenance strategies, sensitivity to antiviral agents, and particular aspects of pathogenesis, the classification does not necessarily predict which clinical signs and symptoms (if any) the virus will cause in humans. Zoonotic viruses are evolving, and "new" zoonotic viruses are regularly discovered. The epizootiology and epidemiology of zoonotic viruses continue to change as a result of environmental alterations affecting vectors, reservoirs, wildlife, livestock, and humans. Zoonotic viruses are most numerous in the tropics but are also found in temperate and even frigid climates. The distribution and seasonal activity of a zoonotic virus may vary, and the rate at which it changes is likely to depend largely on ecologic conditions (e.g., rainfall and temperature), which can affect the density of virus vectors and reservoirs and the development of infection.

Arthropod-borne viruses (arboviruses) infect their vectors after ingestion of a blood meal from a viremic, usually nonhuman vertebrate; some arthropods may also become infected by saliva-activated transmission. The arthropod vectors then develop chronic, systemic infection as the viruses penetrate the gut and spread throughout the body to the salivary glands; such virus dissemination, referred to as *extrinsic incubation*, typically lasts 1–3 weeks in mosquitoes. At this point, if the salivary glands become involved, the arthropod vector is competent to continue the chain of transmission by infecting a vertebrate during a subsequent blood meal. An alternative mechanism for virus maintenance in its arthropod vector is *transovarial transmission*. The arthropod generally is unharmed by the infection, and the natural vertebrate partner usually has only transient viremia with no overt disease.

Rodent-borne viruses are maintained in nature by transmission between rodents, which become chronically infected. Usually a high degree of rodent–virus specificity is observed, and overt disease in the reservoir host is rare.

ETIOLOGY

Arthropod-borne and rodent-borne zoonotic viruses belong to at least seven families: Arenaviridae, Bunyaviridae, Flaviviridae, Orthomyxoviridae, Reoviridae, Rhabdoviridae, and Togaviridae (Table 233-1).

ARENAVIRIDAE

The members of the family Arenaviridae that infect humans are all assigned to the genus *Arenavirus*. The members of this genus are divided into two main phylogenetic branches: Old World viruses (the Lassa–lymphocytic choriomeningitis serocomplex) and New World viruses (the Tacaribe serocomplex). Human arenaviruses form spherical, oval, or pleomorphic enveloped and spiked virions (~50–300 nm in diameter) that bud from the infected cell's plasma membrane. The particles contain two genomic single-stranded RNAs (S, ~3.5 kb; and L, ~7.5 kb) encoding structural proteins in an ambisense orientation. Most arenaviruses persist in nature by chronically infecting rodents. The Old World viruses are maintained by murid rodents that often are persistently viremic and commonly transmit viruses vertically and horizontally. New World viruses are found in cricetid rodents; horizontal transmission is typical, vertical infection may occur, and persistent viremia may be observed. Strikingly, each arenavirus is predominantly adapted to one particular type of rodent. Humans usually become infected through inhalation of or direct contact with infected rodent excreta or secretions (e.g., aerosols of rodents in harvesting machines; aerosolized dried rodent urine or feces in barns or houses; direct contact with rodents in traps). Person-to-person transmission of arenaviruses is uncommon.

BUNYAVIRIDAE

The family Bunyaviridae includes four medically significant genera: *Hantavirus*, *Nairovirus*, *Orthobunyavirus*, and *Phlebovirus*. The members of all these genera form spherical to pleomorphic enveloped virions containing three genomic single-stranded RNAs (S, ~1–2 kb; M, 3.6–5.3 kb; and L, 6.4–12.3 kb) of negative (hantaviruses, nairoviruses, orthobunyaviruses) or ambisense (nairoviruses) polarity. Bunyaviruses mature into particles ~80–120 nm in diameter in the Golgi complex of infected cells and exit these cells by exocytosis.

Hantaviruses are unique among the bunyaviruses in that they are not transmitted by arthropods but instead are maintained in nature by rodents that chronically shed virions. Old World hantaviruses are harbored by murid and cricetid rodents, and New World hantaviruses are maintained by cricetid rodents. As with arenaviruses, individual hantaviruses usually are specifically adapted to a particular type of rodent. However, hantaviruses do not cause chronic viremia in their rodent hosts and are transmitted only horizontally from rodent to rodent. Similar to arenaviruses, hantaviruses infect humans primarily through inhalation of or direct contact with rodent excreta or secretions, and person-to-person transmission is not a common event (with the notable exception of Andes virus). Although there is overlap, the human Old World hantaviruses usually are the etiologic agents of hemorrhagic fever with renal syndrome, whereas the New World viruses usually cause hantavirus pulmonary syndrome.

Nairoviruses are maintained by ixodid ticks, which vertically (transovarially and transstadially) transmit these viruses to progeny tick generations and horizontally spread them through viremic vertebrate hosts. Humans are usually infected via a tick bite or during handling of infected vertebrates.

Orthobunyaviruses are largely mosquito-borne and rarely midge-borne and have viremic vertebrate intermediate hosts. Many orthobunyaviruses are also transovarially transmitted in their mosquito host. Numerous orthobunyaviruses have been associated with human infection and disease. They have been considered to be members of ~19 serogroups based on antigenic cross-reactions, but this grouping is

currently undergoing revision with the accumulation of new genomic data and phylogenetic analyses. Human viruses are found in at least nine serogroups.

Phleboviruses are transmitted vertically (transovarially) in their arthropod hosts and horizontally through viremic vertebrate hosts. Phleboviruses are divided into two groups: the phlebotomus group viruses are transmitted by sandflies and the Uukuniemi group viruses by ticks. Phleboviruses are assigned to at least 10 serocomplexes; human pathogens are found in at least four of these serocomplexes.

FLAVIVIRIDAE

The family Flaviviridae currently includes four genera, one of which (*Flavivirus*) comprises arthropod-borne human viruses. Flaviviruses sensu stricto have single-stranded positive-sense RNA genomes (~11 kb) and form spherical enveloped particles 40–60 nm in diameter. The flaviviruses discussed here belong to two phylogenetically and antigenically distinct groups and are transmitted among vertebrates by mosquitoes and ixodid ticks, respectively. Vectors are usually infected when they feed on viremic hosts; as in the case of most other viruses discussed here, humans are accidental hosts who usually are infected by arthropod bites. Arthropods maintain flavivirus infections horizontally, although transovarial transmission has been documented. Under certain circumstances, flaviviruses can also be transmitted by aerosols or via contaminated food products; in particular, raw milk can transmit tick-borne encephalitis virus.

ORTHOMYXOVIRIDAE

The family Orthomyxoviridae includes two genera of medically relevant arthropod-borne viruses: *Quarantavirus* and *Thogotovirus*. Quarantaviruses are transmitted among birds by ixodid ticks, whereas thogotoviruses have a predilection for mammalian host reservoirs and can be transmitted by both ixodid ticks and mosquitoes.

REOVIRIDAE

The family Reoviridae contains viruses with linear, multisegmented, double-stranded RNA genomes (~16–29 kb in total). These viruses produce particles that have icosahedral symmetry and are 60–80 nm in diameter. In contrast to all other virions discussed here, reovirions are not enveloped and thus are insensitive to detergent inactivation. Fifteen genera of reoviruses are currently recognized. Human arthropod-borne viruses are found among the genera *Coltivirus* (subfamily Spinareovirinae), *Orbivirus*, and *Seadornavirus* (subfamily Sedoreovirinae). Arthropod-borne coltiviruses possess 12 genome segments. Coltiviruses are transmitted by numerous tick types transstadially but not transovarially. Overall maintenance of the transmission cycle, therefore, involves viremic mammalian hosts infected by tick bites. Arthropod-borne orbiviruses have 10 genome segments and are transmitted by mosquitoes or ixodid ticks, whereas relevant seadornaviruses have 12 genome segments and are transmitted exclusively by mosquitoes.

RHABDOVIRIDAE

The family Rhabdoviridae is included in the order *Mononegavirales*. Viruses of the nine rhabdovirus genera have linear, nonsegmented, single-stranded RNA genomes of negative polarity (~11–15 kb) and form bullet-shaped to pleomorphic enveloped particles (100–430 nm long and 45–100 nm wide). Only the genus *Vesiculovirus* includes human arthropod-borne viruses, all of which are transmitted by insects (biting midges, mosquitoes, and sandflies). **The general properties of rhabdoviruses are discussed in more detail in Chap. 232.**

TOGAVIRIDAE

The members of the family Togaviridae have linear, single- and positive-stranded RNA genomes (~9.7–11.8 kb) and form enveloped icosahedral virions (~60–70 nm in diameter) that bud from the plasma membrane of the infected cell. The togaviruses discussed here are all members of the genus *Alphavirus* and are transmitted among vertebrates by mosquitoes.

TABLE 233-1 ZOOBOTIC ARTHROPOD- AND RODENT-BORNE VIRUSES THAT INFECT HUMANS

Virus Group	Virus (Abbreviation)	Principal Reservoir Host(s)	Vector(s)	Syndrome ^a
Alphaviruses (Barmah Forest serocomplex)	Barmah Forest virus (BFV)	Horses, marsupials	Biting midges (<i>Culicoides marksii</i>), mosquitoes (<i>Aedes camptorhynchus</i> , <i>A. normanensis</i> , <i>A. notoscriptus</i> , <i>A. vigilax</i> , <i>Culex annulirostris</i>)	A/R
Alphaviruses (Semliki Forest serocomplex)	Chikungunya virus (CHIKV)	Bats, nonhuman primates	Mosquitoes (<i>Aedes</i> , <i>Culex</i> spp.)	A/R ^b
	Mayaro virus (MAYV)	Nonhuman primates, possums, rodents	Mosquitoes (predominantly <i>Haemagogus</i> spp.)	A/R
	O'nyong-nyong virus (ONNV)	Unknown	Mosquitoes (in particular <i>Anopheles gambiae</i> , <i>A. funestus</i> , <i>Mansonia</i> spp.)	A/R
	Una virus (UNAV)	Birds, horses, rodents	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Coquillettidia</i> , <i>Culex</i> , <i>Ochlerotatus</i> , <i>Psorophora</i> spp.)	F/M
	Ross River virus (RRV)	Macropods, rodents	Mosquitoes (<i>Aedes normanensis</i> , <i>A. vigilax</i> , <i>Culex annulirostris</i>)	A/R
Alphaviruses (eastern equine encephalitis serocomplex)	Semliki Forest virus (SFV)	Birds, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> spp.)	A/R
	Eastern equine encephalitis virus (EEEV)	Freshwater swamp birds	Mosquitoes (<i>Aedes</i> , <i>Coquillettidia</i> , <i>Culex</i> spp.; <i>Culiseta melanura</i> , <i>Mansonia perturbans</i> , <i>Psorophora</i> spp.)	E
	Everglades virus (EVEV)	Hispid cotton rats (<i>Sigmodon hispidus</i>)	Mosquitoes (<i>Culex cedecei</i>)	F/M, E
	Mucambo virus (MUCV)	Nonhuman primates, rodents	Mosquitoes (<i>Culex</i> , <i>Ochlerotatus</i> spp.)	F/M, E
Alphaviruses (Venezuelan equine encephalitis serocomplex)	Tonate virus (TONV)	Suriname crested oropendolas (<i>Psarocolius decumanus</i>)	Mosquitoes (<i>Culex portesi</i>)	F/M, E
	Venezuelan equine encephalitis virus (VEEV)	Horses, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> spp.; <i>Psorophora confinnis</i>)	F/M, E
	Sindbis virus (SINV)	Birds	Mosquitoes (<i>Culex</i> , <i>Culiseta</i> spp.)	A/R
Alphaviruses (western equine encephalitis serocomplex)	Western equine encephalitis virus (WEEV)	Lagomorphs, passerine birds	Mosquitoes (<i>Culex tarsalis</i>)	E
	Lassa virus (LASV)	Natal multimammate mice (<i>Mastomys natalensis</i>)	None	F/M, VHF
Arenaviruses (Old World)	Lujó virus (LUJV)	Unknown	None	VHF
	Lymphocytic choriomeningitis virus (LCMV)	House mice (<i>Mus musculus</i>)	None	E, F/M, (VHF)
	Chapare virus (CHPV)	Unknown	None	VHF
Arenaviruses (New World)	Guanarito virus (GTOV)	Short-tailed zygodonts (<i>Zygodontomys brevicauda</i>)	None	VHF
	Junín virus (JUNV)	Drylands lauchas (<i>Calomys musculinus</i>)	None	VHF
	Machupo virus (MACV)	Big lauchas (<i>Calomys callosus</i>)	None	VHF
	Sabiá virus (SABV)	Unknown	None	VHF
	Whitewater Arroyo virus (WWAV) ^c	White-throated woodrats (<i>Neotoma albigula</i>)	None	(E)
	Bunyaviruses (genus undetermined)	Bangui virus (BGIV)	Unknown	Unknown
Bunyaviruses (genus undetermined)	Gan Gan virus (GGV)	Unknown	Mosquitoes (<i>Aedes</i> , <i>Culex</i> spp.)	A/R
	Tataguine virus (TATV)	Unknown	Mosquitoes (<i>Anopheles</i> spp.)	F/M
	Trubanaman virus (TRUV)	Unknown	Mosquitoes (<i>Anopheles</i> , <i>Culex</i> spp.)	(A/R)
	Colorado tick fever virus (CTFV)	Bushy-tailed woodrats (<i>Neotoma cinerea</i>), Columbian ground squirrels (<i>Spermophilus columbianus</i>), deer mice (<i>Peromyscus maniculatus</i>), golden-mantled ground squirrels (<i>Spermophilus lateralis</i>), least chipmunks (<i>Tamias minimus</i>), North American porcupines (<i>Erethizon dorsata</i>), yellow pine chipmunks (<i>Tamias amoenus</i>)	Ixodid ticks (predominantly <i>Dermacentor andersoni</i>)	E, F/M
Coltivirus	Eyach virus (EYAV)	Lagomorphs, rodents	Ixodid ticks (<i>Ixodes ricinus</i> , <i>I. ventralloii</i>)	E, F/M
	Salmon River virus (SRV)	Unknown	Ixodid ticks (<i>Ixodes</i> spp.)	E, F/M
	Dengue viruses 1–4 (DENV 1–4)	Nonhuman primates	Mosquitoes (predominantly <i>Aedes aegypti</i> , <i>A. albopictus</i>)	F/M, VHF
Flaviviruses (mosquito-borne)	Japanese encephalitis virus (JEV)	Ardeid wading birds (in particular herons), horses, pigs	Mosquitoes (<i>Culex</i> spp., in particular <i>C. tritaeniorhynchus</i>)	E
	Kokobera virus (KOKV)	Macropods, horses	Mosquitoes (<i>Culex</i> spp.)	A/R
	Murray Valley encephalitis virus (MVEV)	Birds	Mosquitoes (predominantly <i>Culex annulirostris</i>)	E

(Continued)

TABLE 233-1 ZOOONOTIC ARTHROPOD- AND RODENT-BORNE VIRUSES THAT INFECT HUMANS (CONTINUED)

Virus Group	Virus (Abbreviation)	Principal Reservoir Host(s)	Vector(s)	Syndrome ^a
	Rocio virus (ROCV)	Rufous-collared sparrows (<i>Zonotrichia capensis</i>)	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Psorophora</i> spp.)	E
	St. Louis encephalitis virus (SLEV)	Columbiform and passeriform birds (finches, sparrows)	Mosquitoes (predominantly <i>Culex</i> spp., in particular <i>C. nigripalpus</i> , <i>C. pipiens</i> , <i>C. quinquefasciatus</i> , <i>C. tarsalis</i>)	E
	Usutu virus (USUV)	Passerine birds	Mosquitoes (<i>Culex</i> spp., in particular <i>C. pipiens</i>)	(E)
	West Nile virus (WNV) ^d	Passerine birds (blackbirds, crows, finches, sparrows), small mammals, horses	Mosquitoes (<i>Culex</i> spp., in particular <i>C. pipiens</i> , <i>C. quinquefasciatus</i> , <i>C. restuans</i> , <i>C. tarsalis</i>)	E
	Yellow fever virus (YFV)	Nonhuman primates (<i>Alouatta</i> , <i>Ateles</i> , <i>Cebus</i> , <i>Cercopithecus</i> , <i>Colobus</i> spp.)	Mosquitoes (<i>Aedes</i> spp., in particular <i>A. aegypti</i>)	VHF
	Zika virus (ZIKV)	Nonhuman primates (<i>Macaca</i> , <i>Pongo</i> spp.)	Mosquitoes (<i>Aedes</i> spp.)	A/R, F/M
Flaviviruses (tick-borne)	Kyasanur Forest disease virus (KFDV) ^e	Indomalayan vandeleurias (<i>Vandeleuria oleracea</i>), roof rats (<i>Rattus rattus</i>)	Ixodid ticks (predominantly <i>Haemaphysalis spinigera</i>); sand tamps (<i>Ornithodoros savignyi</i>)	VHF
	Omsk hemorrhagic fever virus (OHFV)	Migratory birds, rodents	Ixodid ticks (predominantly <i>Dermacentor</i> spp.)	VHF
	Powassan virus (POWV)	Red squirrels (<i>Tamiasciurus hudsonicus</i>), white-footed deer mice (<i>Peromyscus leucopus</i>), woodchucks (<i>Marmota monax</i>), other small mammals	Ixodid ticks (in particular <i>Ixodes cookei</i> , other <i>Ixodes</i> spp., <i>Dermacentor</i> spp.)	E
	Tick-borne encephalitis virus (TBEV)	Passerine birds, deer, eulipotyphla, goats, grouse, small mammals, rodents, sheep	Ixodid ticks (<i>Ixodes gibbosus</i> , <i>I. persulcatus</i> , <i>I. ricinus</i> ; sporadically <i>Dermacentor</i> , <i>Haemaphysalis</i> , <i>Hyalomma</i> spp.)	E, F/M, (VHF)
Hantaviruses (Old World)	Amur/Soochong virus (ASV)	Korean field mice (<i>Apodemus peninsulae</i>)	None	VHF
	Dobrava-Belgrade virus (DOBV)	Caucasus field mice (<i>Apodemus ponticus</i>), striped field mice (<i>Apodemus agrarius</i>), yellow-necked field mice (<i>Apodemus flavicollis</i>)	None	VHF
	Gou virus (GOUV)	Brown rats (<i>Rattus norvegicus</i>), roof rats (<i>R. rattus</i>), Oriental house rats (<i>Rattus tanezumi</i>)	None	VHF
	Hantaan virus (HTNV)	Striped field mice (<i>A. agrarius</i>)	None	VHF
	Kurkino virus	Striped field mice (<i>A. agrarius</i>)	None	VHF
	Muju virus (MUJV)	Korean red-backed voles (<i>Myodes regulus</i>)	None	VHF
	Puumala virus (PUUV)	Bank voles (<i>Myodes glareolus</i>)	None	(P), VHF
	Saaremaa virus (SAAV)	Striped field mice (<i>A. agrarius</i>)	None	VHF
	Seoul virus (SEOV)	Brown rats (<i>R. norvegicus</i>), roof rats (<i>R. rattus</i>)	None	VHF
	Sochi virus	Caucasus field mice (<i>A. ponticus</i>)	None	VHF
	Tula virus (TULV)	Common voles (<i>Microtus arvalis</i>), East European voles (<i>Microtus levis</i>), field voles (<i>Microtus agrestis</i>)	None	(P), VHF
Hantaviruses (New World)	Anajatuba virus (ANJV)	Fornes' colilargos (<i>Oligoryzomys fornesi</i>)	None	P
	Andes virus (ANDV)	Long-tailed colilargos (<i>Oligoryzomys longicaudatus</i>)	None	P
	Araraquara virus (ARAV)	Hairy-tailed akodonts (<i>Necomys lasiurus</i>)	None	P
	Araucária virus (ARAUV)	Black-footed colilargos (<i>Oligoryzomys nigripes</i>)	None	P
	Bayou virus (BAYV)	Marsh rice rats (<i>Oryzomys palustris</i>)	None	P
	Bermejo virus (BMJV)	Chacoan colilargos (<i>Oligoryzomys chacoensis</i>)	None	P
	Black Creek Canal virus (BCCV)	Hispid cotton rats (<i>S. hispidus</i>)	None	P
	Blue River virus (BRV)	White-footed deer mice (<i>P. leucopus</i>)	None	P
	Castelo dos Sonhos virus (CASV)	Brazilian colilargos (<i>Oligoryzomys eliurus</i>)	None	P
	Choclo virus (CHOV)	Fulvous colilargos (<i>Oligoryzomys fulvescens</i>)	None	F/M

(Continued)

TABLE 233-1 ZOOBOTIC ARTHROPOD- AND RODENT-BORNE VIRUSES THAT INFECT HUMANS (CONTINUED)

Virus Group	Virus (Abbreviation)	Principal Reservoir Host(s)	Vector(s)	Syndrome ^a
	El Moro Canyon virus (ELMCV)	Sumichrast's harvest mice (<i>Reithrodontomys sumichrasti</i>), western harvest mice (<i>Reithrodontomys megalotis</i>)	None	P
	Juquitiba virus (JUQV)	Black-footed colilargos (<i>O. nigripes</i>)	None	P
	Laguna Negra virus (LANV)	Little lauchas (<i>Calomys laucha</i>)	None	P
	Lechiguanas virus (LECV)	Flavescent colilargos (<i>Oligoryzomys flavescens</i>)	None	P
	Maciel virus (MCLV)	Dark-furred akodonts (<i>Necromys obscurus</i>)	None	P
	Maripa virus	Unknown	None	P
	Monongahela virus (MGLV)	North American deer mice (<i>P. maniculatus</i>)	None	P
	Muleshoe virus (MULV)	Hispid cotton rat (<i>S. hispidus</i>)	None	P
	New York virus (NYV)	White-footed deer mice (<i>P. leucopus</i>)	None	P
	Orán virus (ORNV)	Long-tailed colilargos (<i>O. longicaudatus</i>)	None	P
	Paranoá virus	Unknown	None	P
	Pergamino virus	Azara's akodonts (<i>Akodon azarae</i>)	None	P
	Río Mamoré virus (RIOMV)	Common bristly mice (<i>Neacomys spinosus</i>)	None	P
	Sin Nombre virus (SNV)	North American deer mice (<i>P. maniculatus</i>)	None	P
	Tunari virus (TUNV)	Unknown	None	P
Nairoviruses (Crimean-Congo hemorrhagic fever virus group)	Crimean-Congo hemorrhagic fever virus (CCHFV)	Cattle, dogs, goats, hares, hedgehogs, mice, ostriches, sheep	Predominantly ixodid ticks (<i>Hyalomma</i> spp.)	VHF
Nairoviruses (Dugbe virus group)	Dugbe virus (DUGV)	Northern giant pouched rats (<i>Cricetomys gambianus</i>), Zébu cattle (<i>Bos primigenius</i>)	Biting midges (<i>Culicoides</i> spp.), ixodid ticks (<i>Amblyomma</i> , <i>Hyalomma</i> , <i>Rhipicephalus</i> spp.)	F/M
	Nairobi sheep disease virus ^f (NSDV)	Sheep	Ixodid ticks (<i>Haemaphysalis</i> , <i>Rhipicephalus</i> spp.), mosquitoes (<i>Culex</i> spp.)	F/M
Nairoviruses (Sakhalin virus group)	Avalon virus (AVAV)	European herring gulls (<i>Larus argentatus</i>)	Ixodid ticks (<i>Ixodes uriae</i>)	(Polyradiculoneuritis?)
Nairoviruses (Thiafora virus group)	Erve virus (ERVEV)	Greater white-toothed shrews (<i>Crocidura russula</i>)	?	(Thunderclap headache?)
Orbiviruses	Kemerovo virus (KEMV)	Birds, rodents	Ixodid ticks (<i>Ixodes persulcatus</i>)	E, F/M
	Lebombo virus (LEBV)	Unknown	Mosquitoes (<i>Aedes</i> , <i>Mansonia</i> spp.)	F/M
	Orungo virus (ORUV)	Camels, cattle, goats, nonhuman primates, sheep	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Culex</i> spp.)	E, F/M
	Tribeč virus (TRBV) ^g	Bank voles (<i>M. glareolus</i>), birds, common pine voles (<i>Microtus subterraneus</i>), goats, hares	Ixodid ticks (<i>Ixodes persulcatus</i> , <i>I. ricinus</i>)	F/M
Orthobunyaviruses (Anopheles A serogroup)	Tacaiuma virus (TCMV)	Nonhuman primates	Mosquitoes (<i>Anopheles</i> , <i>Haemagogus</i> spp.)	F/M
Orthobunyaviruses (Bunyamwera serogroup)	Batai virus (BATV) ^h	Birds, camels, cattle, goats, rodents, sheep	Mosquitoes (<i>Aedes abnormalis</i> , <i>A. curtipes</i> , <i>Anopheles barbirostris</i> , <i>Culex gelidus</i> , other spp.)	F/M
	Bunyamwera virus (BUNV)	Birds, cows, goats, horses, sheep	Mosquitoes (<i>Aedes</i> spp.)	F/M
	Cache Valley virus (CVV)	Cattle, deer, foxes, horses, nonhuman primates, raccoons	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Culiseta</i> spp.)	F/M
	Fort Sherman virus (FSV)	Unknown	Mosquitoes?	F/M
	Germiston virus (GERV)	Rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Guaroa virus (GROV)	Unknown	Mosquitoes (<i>Anopheles</i> spp.)	F/M
	Ilesha virus (ILEV)	Unknown	Mosquitoes (<i>Anopheles gambiae</i>)	F/M, (VHF)
	Ngari virus (NRIV)	Unknown	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> spp.)	F/M, VHF
	Shokwe virus (SHOV)	Rodents	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Mansonia</i> spp.)	F/M
	Xingu virus (XINV)	Unknown	Unknown	F/M
Orthobunyaviruses (Bwamba serogroup)	Bwamba virus (BWAV)	Unknown	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Mansonia</i> spp.)	F/M
	Pongola virus (PGAV)	Cattle, donkeys, goats, sheep	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Mansonia</i> spp.)	F/M

(Continued)

TABLE 233-1 ZONOTIC ARTHROPOD- AND RODENT-BORNE VIRUSES THAT INFECT HUMANS (CONTINUED)

Virus Group	Virus (Abbreviation)	Principal Reservoir Host(s)	Vector(s)	Syndrome ^a
Orthobunyaviruses (California serogroup)	California encephalitis virus (CEV)	Lagomorphs, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Culiseta</i> , <i>Psorophora</i> spp.)	E, F/M
	Inkoo virus (INKV)	Cattle, foxes, hares, moose, rodents	Mosquitoes (<i>Aedes</i> spp.)	E, F/M
	Jamestown Canyon virus (JCV)	Bison, deer, elk, moose	Mosquitoes (<i>Aedes</i> , <i>Culiseta</i> , <i>Ochlerotatus</i> spp.)	E, F/M
	La Crosse virus (LACV)	Chipmunks, squirrels	Mosquitoes (<i>Ochlerotatus triseriatus</i>)	E, F/M
	Lumbo virus (LUMV)	Unknown	Mosquitoes (<i>Aedes pembaensis</i>)	E, F/M
	Snowshoe hare virus (SSHV)	Snowshoe hares, squirrels, other small mammals	Mosquitoes (<i>Aedes</i> , <i>Culiseta</i> , <i>Ochlerotatus</i> spp.)	E, F/M
	Tahyña virus (TAHV)	Cattle, dogs, eulipotyphla, foxes, hares, horses, pigs, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Culiseta</i> spp.)	E, F/M
Orthobunyaviruses (group C serogroup)	Apeú virus (APEUV)	Bare-tailed woolly opossums (<i>Caluromys philander</i>) and other opossums; rodents; tufted capuchins (<i>Cebus apella</i>)	Mosquitoes (<i>Aedes</i> , <i>Culex</i> spp.)	F/M
	Caraparú virus (CARV)	Rodents, tufted capuchins (<i>C. apella</i>)	Mosquitoes (<i>Culex</i> spp.)	F/M
	Itaqui virus (ITQV)	Capuchins (<i>Cebus</i> spp.), opossums, rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Madrid virus (MADV)	Capuchins (<i>Cebus</i> spp.), opossums, rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Marituba virus (MTBV)	Capuchins (<i>Cebus</i> spp.), opossums, rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Murutucú virus (MURV)	Capuchins (<i>Cebus</i> spp.), opossums, pale-throated sloths (<i>Bradypus tridactylus</i>), rodents	Mosquitoes (<i>Coquillettidia</i> , <i>Culex</i> spp.)	F/M
	Nepuyo virus (NEPV)	Bats (<i>Artibeus</i> spp.), rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Oriboca virus (ORIV)	Capuchins (<i>Cebus</i> spp.), opossums, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Mansonia</i> , <i>Psorophora</i> spp.)	F/M
	Ossa virus (OSSAV)	Rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Restan virus (RESV)	Unknown	Mosquitoes (<i>Culex</i> spp.)	F/M
Zungarococha virus (ZUNV)	Unknown	Unknown	F/M	
Orthobunyaviruses (Guama serogroup)	Catu virus (CATUV)	Bats, capuchins (<i>Cebus</i> spp.), opossums, rodents	Mosquitoes (<i>Culex</i> spp.)	F/M
	Guama virus (GMAV)	Bats, capuchins (<i>Cebus</i> spp.), howlers (<i>Alouatta</i> spp.), marsupials, rodents	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Limatus</i> , <i>Mansonia</i> , <i>Psorophora</i> , <i>Trichoprosopon</i> spp.)	F/M
Orthobunyaviruses (Nyando serogroup)	Nyando virus (NDV)	Unknown	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> spp.), sandflies (<i>Lutzomyia</i> spp.)	F/M
Orthobunyaviruses (Simbu serogroup)	Iquitos virus (IQTV)	Unknown	Unknown	F/M
	Oropouche virus (OROV)	Marmosets (<i>Callithrix</i> spp.), pale-throated sloth (<i>B. tridactylus</i>)	Biting midges (<i>Culicoides paraensis</i>), mosquitoes (<i>Coquillettidia venezuelensis</i> , <i>Culex quinquefasciatus</i> , <i>Mansonia</i> spp., <i>Ochlerotatus serratus</i>)	F/M
Orthobunyaviruses (Wyeomyia serogroup)	Wyeomyia virus (WYOV)	Unknown	Mosquitoes (<i>Wyeomyia</i> spp.)	F/M
Phleboviruses (Bhanja serocomplex)	Bhanja virus ¹ (BHAV)	Cattle, four-toed hedgehog (<i>Atelerix albiventris</i>), goats, sheep, striped ground squirrels (<i>Xerus erythropus</i>)	Ixodid ticks (<i>Amblyomma</i> , <i>Dermacentor</i> , <i>Haemaphysalis</i> , <i>Hyalomma</i> , <i>Rhipicephalus</i> spp.)	E, F/M
	Heartland virus	Cattle, deer, elk, goats, sheep?	Ixodid ticks (<i>Amblyomma americanum</i>)	F/M
Phleboviruses (Candiru serocomplex)	"Severe fever with thrombocytopenia syndrome virus" ("SFTSV")	Cattle, chicken, dogs, goats, rodents, sheep?	Ixodid ticks (<i>Haemaphysalis longicornis</i> , <i>Rhipicephalus microplus</i>)	F/M, VHF
	Alenquer virus (ALEV)	Unknown	Unknown	F/M
	Candiru virus (CDUV)	Unknown	Unknown	F/M
	Escharate virus (ESCV)	Unknown	Unknown	F/M
	Maldonado virus (MLOV)	Unknown	Unknown	F/M
	Morumbi virus (MRBV)	Unknown	Unknown	F/M
Phleboviruses (Punta Toro serocomplex)	Serra Norte virus (SRNV)	Unknown	Unknown	F/M
	Punta Toro virus (PTV)	Unknown	Sandflies (<i>Lutzomyia</i> spp.)	F/M
Phleboviruses (sandfly fever serocomplex)	Chagres virus (CHGV)	Unknown	Sandflies (<i>Lutzomyia</i> spp.)	F/M
	Chios virus	Unknown	Unknown	E
	Rift Valley fever virus (RVFV)	Cattle, sheep	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Coquillettidia</i> , <i>Culex</i> , <i>Eretmapodites</i> , <i>Mansonia</i> spp.)	E, F/M, VHF

(Continued)

TABLE 233-1 ZOOBOTIC ARTHROPOD- AND RODENT-BORNE VIRUSES THAT INFECT HUMANS (CONTINUED)

Virus Group	Virus (Abbreviation)	Principal Reservoir Host(s)	Vector(s)	Syndrome ^a
	Sandfly fever Cyprus virus (SFCV)	Unknown	Unknown	F/M
	Sandfly fever Naples virus (SFNV)	Unknown	Sandflies (<i>Phlebotomus papatasi</i> , <i>P. perfiliewi</i> , <i>P. perniciosus</i>)	F/M
	Sandfly fever Sicilian virus (SFSV)	Eulipotyphla, least weasles (<i>Mustela nivalis</i>), rodents	Sandflies (particularly <i>Phlebotomus papatasi</i>)	F/M
	Sandfly fever Turkey virus (SFTV)	Unknown	Sandflies (<i>Phlebotomus</i> spp.)	F/M
	Toscana virus (TOSV)	Unknown	Sandflies (<i>Phlebotomus papatasi</i> , <i>P. perfiliewi</i>)	E, F/M
Phleboviruses (Uukuniemi serocomplex)	Uukuniemi virus (UUKV)	Birds, cattle, rodents	Ixodid ticks (<i>Ixodes</i> spp.)	F/M
Quarantaviruses	Quaranfil virus (QRFV)	Birds	Ixodid ticks (<i>Argas arboreus</i>)	F/M
Seadornaviruses	Banna virus (BAV)	Cattle, pigs	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Culiseta</i> spp.)	E
Thogotoviruses	Dhori virus (DHOV) ^k	Bats, camels, horses	Mosquitoes (<i>Aedes</i> , <i>Anopheles</i> , <i>Culex</i> spp.), ixodid ticks (<i>Dermacentor</i> , <i>Hyalomma</i> , <i>Ornithodoros</i> spp.)	E, F/M
	Thogoto virus (THOV)	Camels, cattle	Ixodid ticks (<i>Amblyomma</i> , <i>Hyalomma</i> , <i>Rhipicephalus</i> spp.)	E, F/M
Vesiculoviruses	Chandipura virus (CHPV)	Hedgehogs	Mosquitoes (<i>Aedes aegypti</i>), sandflies (<i>Phlebotomus</i> , <i>Sergentomyia</i> spp.)	E, F/M
	Isfahan virus (ISFV)	Great gerbils (<i>Rhombomys opimus</i>)	Sandflies (<i>Phlebotomus papatasi</i>)	F/M
	Piry virus (PIRYV)	Gray four-eyed opossums (<i>Philander opossum</i>)	Mosquitoes (<i>Aedes</i> , <i>Culex</i> , <i>Toxorhynchites</i> spp.)	F/M
	Vesicular stomatitis Indiana virus (VSIV)	Cattle, horses, pigs	Sandflies (<i>Lutzomyia</i> spp.)	F/M
	Vesicular stomatitis New Jersey virus (VSNJV)	Cattle, horses, pigs	Biting midges (<i>Culicoides</i> spp.), chloropid flies, mosquitoes (<i>Culex</i> , <i>Mansonia</i> spp.), muscoid flies (<i>Musca</i> spp.), simuliid flies	F/M

^aAbbreviations refer to the syndrome most commonly associated with the virus: A/R, arthritis/rash; E, encephalitis; F/M, fever/myalgia; P, pulmonary; VHF, viral hemorrhagic fever. Abbreviations are placed in parentheses when cases are either extremely rare or controversial. ^bIn the older literature, chikungunya virus often is also listed as a causative agent of VHF. However, later studies revealed that, in most cases, people with "chikungunya hemorrhagic fever" were co-infected with one or more dengue viruses, an observation suggesting that the VHF was actually severe dengue. ^cWhitewater Arroyo virus is often listed as a causative agent of VHF in the literature, but convincing data associating this virus with VHF have not been published. ^dAlso includes Kunjin virus. ^eIncludes the recently described Alkhurma/Alkhurma variant of Kyasanur Forest disease virus. ^fAlso known as Ganjam virus. ^gAlso known as Brezová virus, Cvilin virus, Kharagsh virus, Koliba virus, or Lipovnik virus. ^hAlso known as Čalovo virus or Chittoor virus. ⁱAlso known as Palma virus. ^jThe final virus name has not yet been decided. Alternatives used in the literature are Huaiyangshan virus (HYSV) and Henan fever virus (HNFV). ^kAlso known as Astra virus and Batken virus.

EPIDEMIOLOGY

The distributions of arthropod-borne and rodent-borne viruses are restricted by the areas inhabited by their reservoir hosts and/or vectors. Consequently, a patient's geographic origin or travel history can provide important clues in the differential diagnosis. **Table 233-2** lists the approximate geographic distribution of most arthropod-borne and rodent-borne infections. Many of these diseases can be acquired in either rural or urban settings; these diseases include yellow fever, dengue (previously called dengue fever), severe dengue (previously called dengue hemorrhagic fever and dengue shock syndrome), chikungunya virus disease, hemorrhagic fever with renal syndrome caused by Seoul virus, sandfly fever caused by sandfly fever Naples and Sicilian viruses, and Oropouche virus disease.

DIAGNOSIS

In patients with suspected viral infection, a recognized history of mosquito bite has little diagnostic significance, but a history of tick bite is more diagnostically useful. Exposure to rodents is sometimes reported by persons infected with arenaviruses or hantaviruses. Laboratory diagnosis is required in all cases, although epidemics occasionally provide enough clinical and epidemiologic clues for a presumptive etiologic diagnosis. For most arthropod-borne and rodent-borne viruses, acute-phase serum samples (collected within 3 or 4 days of onset) have yielded isolates. Paired serum samples have been used to demonstrate rising antibody titers. Intensive efforts to develop rapid tests for viral hemorrhagic fevers have resulted in reliable antigen-detection enzyme-linked immunosorbent assays (ELISAs), IgM-capture ELISAs,

and multiplex polymerase chain reaction (PCR) assays. These tests can provide a diagnosis based on a single serum sample within a few hours and are particularly useful in patients with severe disease. More sensitive reverse-transcription PCR (RT-PCR) assays may yield diagnoses based on samples without detectable antigen and may also provide useful genetic information about the etiologic agent.

Hantavirus infections differ from other viral infections discussed here in that severe acute disease is immunopathologic; patients present with serum IgM that serves as the basis for a sensitive and specific test. At diagnosis, patients with encephalitides generally are no longer viremic or antigenemic and usually do not have virions in cerebrospinal fluid (CSF). In this situation, the value of serologic methods for IgM determination and RT-PCR is high. IgM-capture ELISA is increasingly being used for the simultaneous testing of serum and CSF. IgG ELISA or classic serology is useful in the evaluation of past exposure to viruses, many of which circulate in areas with minimal medical infrastructures and sometimes cause only mild or subclinical infection.

SYNDROMES

The spectrum of possible human responses to infection with arthropod- or rodent-borne viruses is wide, and knowledge of the outcome of most of these infections is limited. People infected with these viruses may not develop signs of illness. If viral disease is recognized, it can usually be grouped into one of five broad categories: arthritis and rash, encephalitis, fever and myalgia, pulmonary disease, or viral hemorrhagic fever (VHF) (**Table 233-3**). These categories often overlap. For example, infections with West Nile and Venezuelan equine

TABLE 233-2 GEOGRAPHIC DISTRIBUTION (UNITED NATIONS GEOScheme) OF ZOO NOTIC ARTHROPOD-BORNE OR RODENT-BORNE VIRAL DISEASES

Area	Type of Disease ^a						
	Arenaviral	Bunyaviral	Flaviviral	Orthomyxoviral	Reoviral	Rhabdoviral	Togaviral
Africa	Lassa fever; Lujo virus infection	Bangui, Batai, Bhanja, Bunyamwera, and Bwamba virus infections; Crimean-Congo hemorrhagic fever; Dugbe, Germiston, Ilesha virus infections; Nairobi sheep disease virus infection; Ngari, Nyando, and Pongola virus infections; Rift Valley fever, sandfly fever/Pappataci fever/phlebotomus fever; Shokwe, Tatuquine virus infections	Dengue/severe dengue; (Usutu virus infection); West Nile virus infection; yellow fever; Zika virus infection	Dhori, Quarantil, and Thogoto virus infections	Lebombo, Orungo, and Tribeč virus infections	—	Chikungunya virus disease; o'nyong-nyong fever; Semliki Forest and Sindbis virus infections
Central Asia	—	Bhanja virus infection; Crimean-Congo hemorrhagic fever	Tick-borne viral encephalitis	Dhori virus infection	—	Isfahan virus infection	—
Eastern Asia	—	Crimean-Congo hemorrhagic fever; hemorrhagic fever with renal syndrome; sandfly fever/Pappataci fever/phlebotomus fever; severe fever with thrombocytopenia syndrome	Dengue/severe dengue; Japanese encephalitis; Kyasanur Forest disease; tick-borne viral encephalitis	—	Banna virus infection	—	—
Southern Asia	—	Batai and Bhanja virus infections; Crimean-Congo hemorrhagic fever; hemorrhagic fever with renal syndrome; Nairobi sheep disease virus infection; sandfly fever/Pappataci fever/phlebotomus fever	Dengue/severe dengue; Japanese encephalitis; Kyasanur Forest disease; West Nile virus infection; Zika virus infection	Dhori, Quarantil, and Thogoto virus infections	—	Chandipura and Isfahan virus infections	Chikungunya virus disease
South-Eastern Asia	—	Batai virus infection; hemorrhagic fever with renal syndrome	Dengue/severe dengue; Japanese encephalitis; West Nile virus infection; Zika virus infection	—	—	—	Chikungunya virus disease
Western Asia	—	Batai and Bhanja virus infections; Crimean-Congo hemorrhagic fever; hemorrhagic fever with renal syndrome; sandfly fever/Pappataci fever/phlebotomus fever	Dengue/severe dengue; Kyasanur Forest disease; tick-borne viral encephalitis; West Nile virus infection	Dhori and Quarantil virus infections	—	—	Chikungunya virus disease
Latin America and the Caribbean	"Brazilian hemorrhagic fever"; Chapare virus infection; Junin/Argentinian hemorrhagic fever; lymphocytic choriomeningitis/meningoencephalitis; Machupo/Bolivian hemorrhagic fever; "Venezuelan hemorrhagic fever"	Alenquer, Apeú, Bunyamwera, Cache Valley, Candiru, Caraparú, Catu, Chagres, Escharate, Fort Sherman, Guama, and Guaroa virus infections; hantavirus pulmonary syndrome; Itaqui, Jucituba, Madrid, Maldonado, Marituba, Mayaro, Morumbi, Murutucú, Nepuyo, and Oriboca virus infections; Oropouche virus disease; Ossa, Punta Toro, Restan, Serra Norte, Tacaiuma, Trinidad, Wyeomyia, Xingu, and Zungarococha virus infections	Dengue/severe dengue; Rocio virus disease; yellow fever	—	—	Piry virus disease; vesicular stomatitis virus disease/Indiana fever	Chikungunya virus disease; Mayaro virus infection; Mucambo, Tonate, and Una virus infections; Venezuelan equine fever
Northern America	Lymphocytic choriomeningitis/meningoencephalitis; (Whitewater Arroyo virus infection)	(Avalon) and Cache Valley virus infections; California (meningo)encephalitis; hantavirus pulmonary syndrome; Heartland virus and Nepuyo virus infections	Dengue/severe dengue; Powassan virus disease; St. Louis encephalitis; West Nile virus infection	—	Colorado tick fever; Salmon River virus infection	Vesicular stomatitis virus disease/Indiana fever	Eastern equine encephalitis; Everglades virus infection; western equine encephalitis

(Continued)

Area	Type of Disease ^a						
	Arenaviral	Bunyaviral	Flaviviral	Orthomyxoviral	Reoviral	Rhabdoviral	Togaviral
Europe	Lymphocytic choriomeningitis/meningoencephalitis	(Avalon) and Bhanja virus infections; California (meningo)encephalitis; Crimean-Congo hemorrhagic fever; (Erve virus infection); hemorrhagic fever with renal syndrome; Inkoo virus infection; sandfly fever/Pappataci fever/phlebotomus fever; Uukuniemi virus infection	Dengue/severe dengue; tick-borne viral encephalitis; Omsk hemorrhagic fever; (Usutu virus infection); West Nile virus infection	Dhori and Thogoto virus infections	Eyach, Kemerovo, and Tribeč virus infections	—	Chikungunya virus disease; Sindbis virus infection
Oceania	—	Gan Gan and (Trubanaman virus) infections	Australian encephalitis; dengue/severe dengue; Japanese encephalitis; Kokobera virus infection; Murray Valley encephalitis; West Nile virus infection; Zika virus infection	—	—	—	Barmah Forest virus infection; Ross River disease; Sindbis virus infection

^aQuotation marks indicate common usage in the absence of International Classification of Disease version 10 (ICD-10) recognition. Diseases not acknowledged by the ICD-10 are designated as "virus infection." Disease names are placed in parentheses when cases are either extremely rare or controversial.

TABLE 233-3 CLINICAL SYNDROMES CAUSED BY ZONOTIC ARTHROPOD-BORNE OR RODENT-BORNE VIRUSES

Syndrome	Virus ^a
Arthritis and rash (A/R)	Bunyaviridae: Gan Gan and (Trubanaman) viruses Flaviviridae: Kokobera and Zika viruses Togaviridae: Barmah Forest, chikungunya, Mayaro, o'nyong-nyong, Ross River, Semliki Forest, and Sindbis viruses
Encephalitis (E)	Arenaviridae: lymphocytic choriomeningitis and (Whitewater Arroyo) viruses Bunyaviridae: Bhanja, California encephalitis, Chios, Inkoo, Jamestown Canyon, La Crosse, Lumbo, Rift Valley fever, snowshoe hare, Tahyña, and Toscana viruses Flaviviridae: Japanese encephalitis, Murray Valley encephalitis, Powassan, Rocio, St. Louis encephalitis, tick-borne encephalitis, (Usutu), and West Nile viruses Orthomyxoviridae: Dhori and Thogoto viruses Reoviridae: Banna, Colorado tick fever, Eyach, Kemerovo, Orungo, and Salmon River viruses Rhabdoviridae: Chandipura virus Togaviridae: eastern equine encephalitis, Everglades, Mucambo, Tonate, Venezuelan equine encephalitis, and western equine encephalitis viruses
Fever and myalgia (F/M)	Arenaviridae: Lassa and lymphocytic choriomeningitis viruses Bunyaviridae: Alenquer, Apeú, Bangui, Batai, Bhanja, Bunyamwera, Bwamba, Cache Valley, California encephalitis, Candiru, Caraparú, Catu, Chagres, Choclo, Dugbe, Escharate, Fort Sherman, Germiston, Guama, Guaroa, Heartland, Ilesha, Inkoo, Iquitos, Itaqui, Jamestown Canyon, La Crosse, Lumbo, Madrid, Maldonado, Marituba, Morumbi, Nairobi sheep disease, Nepuyo, Ngari, Nyando, Oriboca, Oropouche, Ossa, Pongola, Punta Toro, Restan, Rift Valley fever, sandfly fever Cyprus, sandfly fever Naples, sandfly fever Sicilian, sandfly fever Turkey, Serra Norte, "severe fever with thrombocytopenia syndrome," Shokwe, snowshoe hare, Tacaiuma, Tahyña, Tataguine, Thogoto, Toscana, Uukuniemi, Wyeomyia, Xingu, and Zungarococha viruses Flaviviridae: dengue 1–4, tick-borne encephalitis, and Zika viruses Orthomyxoviridae: Dhori and Quaranfil viruses Reoviridae: Colorado tick fever, Eyach, Kemerovo, Lebombo, Orungo, Salmon River, and Tribeč viruses Rhabdoviridae: Chandipura, Isfahan, Piry, vesicular stomatitis Indiana, and vesicular stomatitis New Jersey viruses Togaviridae: Everglades, Mucambo, Tonate, Una, and Venezuelan equine encephalitis viruses
Pulmonary disease (P)	Bunyaviridae: Anajatuba, Andes, Araucária, Bayou, Bermejo, Black Creek Canal, Blue River, Castelo dos Sonhos, El Moro Canyon, Jucituba, Laguna Negra, Lechiguana, Maciel, Monongahela, Muleshoe, New York, Orán, Paranoá, Pergamino, (Puumala), Río Mamoré, Sin Nombre, (Tula), and Tunari viruses
Viral hemorrhagic fever (VHF)	Arenaviridae: Chapare, Guanarito, Junín, Lassa, Lujo, (lymphocytic choriomeningitis), Machupo, and Sabiá viruses Bunyaviridae: Amur/Soochong, Crimean-Congo hemorrhagic fever, Dobrava-Belgrade, Gou, Hantaan, (Ilesha), Kurkino, Muju, Ngari, Puumala, Rift Valley fever, Saaremaa, Seoul, "severe fever with thrombocytopenia syndrome," Sochi, and Tula viruses Flaviviridae: dengue 1–4, Kyasanur Forest disease, Omsk hemorrhagic fever, (tick-borne encephalitis), and yellow fever viruses

^aVirus names are placed in parentheses when human infections are either extremely rare or controversial.

encephalitis viruses are discussed here as encephalitides, but during epidemics many patients present with much milder febrile syndromes. Similarly, Rift Valley fever virus is best known as a cause of VHF, but the attack rates for febrile disease are far higher, and encephalitis and blindness occasionally occur as well. Lymphocytic choriomeningitis virus is classified here as a cause of fever and myalgia because this syndrome is the most common disease manifestation; even when central nervous system (CNS) disease evolves during infection with this virus, neural manifestations are usually mild and are preceded by fever and myalgia. Infection with any dengue virus type (1, 2, 3, or 4) is considered as a cause of fever and myalgia because this syndrome is by far the most common manifestation worldwide. However, severe dengue is a VHF with a complicated pathogenesis that is of tremendous importance in pediatric practice in certain areas of the world. Unfortunately, most of the known arthropod- or rodent-borne viral diseases have not been studied in detail with modern medical approaches; thus available data may be incomplete or biased. The reader must be aware that data on geographic distribution are often fuzzy: the literature frequently is not clear as to whether the data pertain to the distribution of a particular virus or the areas where human disease has been observed. In addition, the designations for viruses and viral diseases have changed multiple times over decades. Here, virus and taxon names are in line with the latest reports of the International Committee on Taxonomy of Viruses, and disease names are largely in accordance with the World Health Organization's International Classification of Disease version 10 (ICD-10) and more recent updates.

ARTHRITIS AND RASH

Arthritides are common accompaniments of several viral diseases, such as hepatitis B, parvovirus B19 infection, and rubella, and occasionally accompany infection due to adenoviruses, enteroviruses, herpesviruses, and mumps virus. Two ungrouped bunyaviruses, Gan Gan virus and Trubanaman virus, and the flavivirus Kokobera virus have been associated with single cases of polyarthritic disease. Arthropod-borne alphaviruses are also common causes of arthritides—usually acute febrile diseases accompanied by the development of a maculopapular rash. Rheumatic involvement includes arthralgia alone, periarticular swelling, and (less commonly) joint effusions. Most alphavirus infections are less severe and have fewer articular manifestations in children than in adults. In temperate climates, these ailments are summer diseases. No specific therapies or licensed vaccines exist. The most important alphavirus arthritides are Barmah Forest virus infection, chikungunya virus disease, Ross River disease, and Sindbis virus infection. A large (>2 million cases), albeit isolated, epidemic was caused by o'nyong nyong virus in 1959–1961 (o'nyong nyong fever). Mayaro, Semliki Forest, and Una viruses have caused isolated cases or limited and infrequent epidemics (30 to several hundred cases per year) in the past. Signs and symptoms of infections with these viruses often are similar to those observed with chikungunya virus disease.

Chikungunya Virus Disease Disease caused by chikungunya virus is endemic in rural areas of Africa. Intermittent epidemics take place in towns and cities of both Africa and Asia. *Aedes aegypti* mosquitoes are the usual vectors for the disease in urban areas. In 2004, a massive epidemic began in the Indian Ocean region (in particular on the islands of Réunion and Mauritius) and was most likely spread by travelers; *Aedes albopictus* was identified as the major vector of chikungunya virus during that epidemic. Between 2013 and 2014, several thousand chikungunya virus infections were reported (and several tens of thousands of cases were suspected) from Caribbean islands. The virus was imported to Italy, France, and the United States by travelers from the Caribbean. Chikungunya virus poses a threat to the continental United States as suitable vector mosquitoes are present in the southern states. The disease is most common among adults, in whom the clinical presentation may be dramatic. The abrupt onset of chikungunya virus disease follows an incubation period of 2–10 days. Fever (often severe) with a saddleback pattern and severe arthralgia are accompanied by chills and constitutional symptoms and signs, such as abdominal pain, anorexia, conjunctival injection, headache, nausea,

and photophobia. Migratory polyarthritis mainly affects the small joints of the ankles, feet, hands, and wrists, but the larger joints are not necessarily spared. Rash may appear at the outset or several days into the illness; its development often coincides with defervescence, which occurs around day 2 or 3 of the disease. The rash is most intense on the trunk and limbs and may desquamate. Young children develop less prominent signs and are therefore less frequently hospitalized. Children also often develop a bullous rather than a maculopapular/ petechial rash. Maternal–fetal transmission has been reported and in some cases has led to fetal death. Recovery may require weeks, and some elderly patients may continue to experience joint pain, recurrent effusions, or stiffness for several years. This persistence of signs and symptoms may be especially common in HLA-B27-positive patients. In addition to arthritis, petechiae are occasionally seen and epistaxis is not uncommon, but chikungunya virus should not be considered a VHF agent. A few patients develop leukopenia. Elevated concentrations of aspartate aminotransferase (AST) and C-reactive protein have been described, as have mildly decreased platelet counts. Treatment of chikungunya virus disease relies on nonsteroidal anti-inflammatory drugs and sometimes chloroquine for refractory arthritis.

Barmah Forest Virus Infection and Ross River Disease Barmah Forest virus and Ross River virus cause diseases that are indistinguishable on clinical grounds alone (hence the previously common disease designation *epidemic polyarthritis* for both infections). Ross River virus has caused epidemics in Australia, Papua New Guinea, and the South Pacific since the beginning of the twentieth century and continues to be responsible for ~4800 cases of disease in rural and suburban areas annually. In 1979–1980, the virus swept through the Pacific Islands, causing more than 500,000 infections. Ross River virus is predominantly transmitted by *Aedes normanensis*, *Aedes vigilax*, and *Culex annulirostris*. Wallabies and rodents are probably the main vertebrate hosts. Barmah Forest virus infections have been on the rise in recent years. In 2005–2006, roughly 2000 cases were recorded in Australia. Barmah Forest virus is transmitted by both *Aedes* and *Culex* mosquitoes and has been isolated from biting midges. The vertebrate hosts remain to be determined, but serologic studies implicate horses and possums.

Of the human Barmah Forest and Ross River virus infections surveyed, 55–75% were asymptomatic; however, these viral diseases can be debilitating. The incubation period is 7–9 days; the onset of illness is sudden, and disease is usually ushered in by disabling symmetrical joint pain. A nonitchy, diffuse, maculopapular rash (more common in Barmah Forest virus infection) generally develops coincidentally or follows shortly, but in some patients it can precede joint pains by several days. Constitutional symptoms such as low-grade fever, asthenia, headache, myalgia, and nausea are not prominent or are absent in many patients. Most patients are incapacitated for considerable periods (≥6 months) by joint involvement, which interferes with grasping, sleeping, and walking. Ankle, interphalangeal, knee, metacarpophalangeal, and wrist joints are most often involved, although elbows, shoulders, and toes may also be affected. Periarticular swelling and tenosynovitis are common, and one-third of patients have true arthritis (more common in Ross River disease). Myalgia and nuchal stiffness may accompany joint pains. Only half of all patients with arthritis can resume normal activities within 4 weeks, and 10% still must limit their activity after 3 months. Occasional patients are symptomatic for 1–3 years but without progressive arthropathy.

In the diagnosis of either infection, clinical laboratory values are normal or variable. Tests for rheumatoid factor and antinuclear antibodies are negative, and the erythrocyte sedimentation rate is acutely elevated. Joint fluid contains 1000–60,000 mononuclear cells/ μ L, and viral antigen can usually be detected in macrophages. IgM antibodies are valuable in the diagnosis of this infection, although such antibodies occasionally persist for years. Isolation of the virus from blood after mosquito inoculation or growth of the virus in cell culture is possible early in the illness. Because of the great economic impact of annual epidemics in Australia, an inactivated Ross River virus vaccine is under development. Nonsteroidal anti-inflammatory drugs such as naproxen or acetylsalicylic acid are effective for treatment.

1314 Sindbis Virus Infection Sindbis virus is transmitted among birds by infected mosquitoes. Infections with northern European or southern African variants are particularly likely in rural environments. After an incubation period of <1 week, Sindbis virus infection begins with rash and arthralgia. Constitutional clinical signs are not marked, and fever is modest or lacking altogether. The rash, which lasts ~1 week, begins on the trunk, spreads to the extremities, and evolves from macules to papules that often vesiculate. The arthritis is multiarticular, migratory, and incapacitating, with resolution of the acute phase in a few days; the ankles, elbows, knees, phalangeal joints, wrists, and—to a much lesser extent—proximal and axial joints are involved. Persistence of joint pain and occasionally of arthritis is a major problem and may continue for months or even years despite lack of deformities.

Zika Virus Infection Zika virus is an emerging pathogen that is transmitted among nonhuman primates and humans by *Aedes* mosquitoes. Human infections are usually benign and are most likely misdiagnosed as dengue or influenza. Zika virus infection is characterized by influenza-like clinical signs, including fever, headaches, and malaise. A maculopapular rash, conjunctivitis, myalgia, and arthralgia usually accompany or follow those manifestations. Zika virus infection was first documented in Africa in 1947 and was later recognized in southeastern and southern Asia. In recent years, the number of Zika virus infections reported from Micronesia and Polynesia has increased steadily.

ENCEPHALITIS

The major encephalitis viruses are found in the families Bunyaviridae, Flaviviridae, Rhabdoviridae, and Togaviridae. However, individual agents of other families, including Dhori virus and thogotovirus (Orthomyxoviridae) as well as Banna virus (Reoviridae), have been known to cause isolated cases of encephalitis as well. Arboviral encephalitides are seasonal diseases, commonly occurring in the warmer months. Their incidence varies markedly with time and place, depending on ecologic factors. The causative viruses differ substantially in terms of case–infection ratio (i.e., the ratio of clinical to sub-clinical infections), lethality rate, and residual disease. Humans are not important amplifiers of these viruses.

All the viral encephalitides discussed in this section have a similar pathogenesis. An infected arthropod ingests blood from a human and thereby initiates infection. The initial viremia is thought to originate from the lymphoid system. Viremia leads to multifocal entry into the CNS, presumably through infection of olfactory neuroepithelium, with passage through the cribriform plate; “Trojan horse” entry with infected macrophages; or infection of brain capillaries. During the viremic phase, there may be little or no recognizable disease except in tick-borne flavivirus encephalitides, which may manifest with clearly delineated phases of fever and systemic illness.

CNS lesions arise partly from direct neuronal infection and subsequent damage and partly from edema, inflammation, and other indirect effects. The usual pathologic features of arboviral encephalitides are focal necroses of neurons, inflammatory glial nodules, and perivascular lymphoid cuffing. Involved areas display the “luxury perfusion” phenomenon, with normal or increased total blood flow and low oxygen extraction. The typical patient presents with a prodrome of nonspecific constitutional signs and symptoms, including fever, abdominal pain, sore throat, and respiratory signs. Headache, meningeal signs, photophobia, and vomiting follow quickly. The severity of human infection varies from an absence of signs/symptoms to febrile headache, aseptic meningitis, and full-blown encephalitis. The proportions and severity of these manifestations vary with the infecting virus. Involvement of deeper brain structures in less severe cases may be signaled by lethargy, somnolence, and intellectual deficit (as disclosed by the mental status examination). More severely affected patients are obviously disoriented and may become comatose. Tremors, loss of abdominal reflexes, cranial nerve palsies, hemiparesis, monoparesis, difficulty swallowing, limb-girdle syndrome, and frontal lobe signs are all common. Spinal and motor neuron diseases are documented after West Nile and Japanese encephalitis virus infections. Seizures and focal signs may be evident early or may appear during the course of the

disease. Some patients present with an abrupt onset of fever, convulsions, and other signs of CNS involvement. The acute encephalitis usually lasts from a few days to as long as 2–3 weeks. The infections may be fatal, or recovery may be slow, with weeks or months required for the return of maximal recoupable function, or incomplete, with persisting long-term deficits. Difficulty concentrating, fatigability, tremors, and personality changes are common during recovery.

The diagnosis of arboviral encephalitides depends on the careful evaluation of a febrile patient with CNS disease and the performance of laboratory studies to determine etiology. Clinicians should (1) consider empirical acyclovir treatment for herpesvirus meningoencephalitis and antibiotic treatment for bacterial meningitis until test results are received; (2) exclude intoxication and metabolic or oncologic causes, including paraneoplastic syndromes, hyperammonemia, liver failure, and anti-NMDA receptor encephalitis; and (3) rule out a brain abscess or a stroke. Leptospirosis, neurosyphilis, Lyme disease, cat-scratch disease, and more recently described viral encephalitides (e.g., Nipah virus infection) should be considered if epidemiologically relevant. CSF examination usually shows a modest increase in leukocyte counts—in the tens or hundreds or perhaps a few thousand. Early in the process, a significant proportion of these leukocytes may be polymorphonuclear, but mononuclear cells are usually predominant later. CSF glucose concentrations are generally normal. There are exceptions to this pattern of findings: in eastern equine encephalitis, for example, polymorphonuclear leukocytes may predominate during the first 72 h of disease and hypoglycorrhachia may be detected. In lymphocytic choriomeningitis/meningoencephalitis, lymphocyte counts may be in the thousands, and the glucose concentration may be diminished. A humoral immune response is usually detectable at or near the onset of disease. Both serum (acute- or convalescent-phase) and CSF should be examined for IgM antibodies and viruses by plaque-reduction neutralization assay and/or (RT)-PCR. Virus generally cannot be isolated from blood or CSF, although Japanese encephalitis virus has been recovered from CSF of patients with severe disease. RT-PCR analysis of CSF may yield positive results. Viral antigen is present in brain tissue, although its distribution may be focal. Electroencephalography usually shows diffuse abnormalities and is not directly helpful.

Experience with medical imaging is still evolving. Both computed tomography (CT) and magnetic resonance imaging (MRI) scans may be normal except for evidence of preexisting conditions or occasional diffuse edema. Imaging is generally nonspecific in that most patients do not present with pathognomonic lesions, but it can be used to rule out other suspected causes of disease. It is important to remember that imaging may yield negative results if done early in the disease course but later may detect lesions. For example, eastern equine encephalitis (focal abnormalities) and severe Japanese encephalitis (hemorrhagic bilateral thalamic lesions) have caused lesions detectable by medical imaging.

Comatose patients may require management of intracranial pressure elevations, inappropriate secretion of antidiuretic hormone, respiratory failure, or seizures. Specific therapies for these viral encephalitides are not available. The only practical preventive measures are vector management and personal protection against the arthropod transmitting the virus. For Japanese encephalitis or tick-borne viral encephalitis, vaccination should be considered in certain circumstances (see relevant sections below).

Bunyaviruses: California (Meningo)encephalitis The isolation of California encephalitis virus established California serogroup orthobunyaviruses as causes of encephalitides. However, California encephalitis virus has been implicated in only a very few cases of encephalitis, whereas its close relative, La Crosse virus, is the major cause of encephalitis in this serogroup (~70 cases per year in the United States). California (meningo)encephalitis due to La Crosse virus infection is most commonly reported from the upper Midwest of the United States but is also found in other areas of the central and eastern parts of the country, most often in West Virginia, Tennessee, North Carolina, and Georgia. The serogroup includes 13 other viruses, some of which (e.g., Inkoo, Jamestown Canyon, Lumbo, snowshoe hare, and Tahyña

viruses) also cause human disease. Transovarial transmission is a strong component of transmission of the California serogroup viruses in *Aedes* and *Ochlerotatus* mosquitoes. The mosquito vector of La Crosse virus is *Ochlerotatus triseriatus*. In addition to transovarial transmission, acquisition through feeding on viremic chipmunks and other mammals as well as venereal transmission can result in infection of this mosquito. *O. triseriatus* breeds in sites such as tree holes and abandoned tires and bites during daylight hours. The habits of this mosquito correlate with the risk factors for human cases: recreation in forested areas, residence at a forest's edge, and the presence of water-containing abandoned tires around the home. Intensive environmental modification based on these findings has reduced the incidence of disease in a highly endemic area in the U.S. Midwest.

Most humans are infected from July through September. The Asian tiger mosquito (*A. albopictus*) efficiently transmits La Crosse virus to mice and also transmits the agent transovarially in the laboratory; this aggressive anthropophilic mosquito has the capacity to urbanize, and its possible impact on transmission of virus to humans is of concern. The prevalence of antibody to La Crosse virus in humans is $\geq 20\%$ in endemic areas, a figure indicating that infection is common but often asymptomatic. CNS disease has been recognized primarily in children <15 years of age.

The illness from La Crosse virus varies from aseptic meningitis accompanied by confusion to severe and occasionally fatal encephalitis (lethality rate, <0.5%). The incubation period is ~3–7 days. Although there may be prodromal symptoms/signs, the onset of CNS disease is sudden, with fever, headache, and lethargy often joined by nausea and vomiting, convulsions (in one-half of patients), and coma (in one-third of patients). Focal seizures, hemiparesis, tremor, aphasia, chorea, Babinski signs, and other evidence of significant neurologic dysfunction are common, but residual disease is not. Approximately 10% of patients have recurrent seizures in the succeeding months. Other serious sequelae of La Crosse virus infection are rare, although a decrease in scholastic standing in children has been reported and mild personality change has occasionally been suggested.

The blood leukocyte count is commonly elevated in patients with La Crosse virus infection, sometimes reaching 20,000/ μL , and is usually accompanied by a left shift. CSF leukocyte counts are typically 30–500/ μL with a mononuclear cell predominance (although 25–90% of cells are polymorphonuclear in some patients). The blood protein concentration is normal or slightly increased, and the glucose concentration is normal. Specific virologic diagnosis based on IgM-capture assays of serum and CSF is efficient. The only human anatomic site from which virus has been isolated is the brain.

Treatment is supportive over a 1- to 2-week acute phase during which status epilepticus, cerebral edema, and inappropriate secretion of antidiuretic hormone are important concerns. A phase 2B clinical trial of IV ribavirin in children with La Crosse virus infection was discontinued during dose escalation because of adverse effects.

Jamestown Canyon virus has been implicated in several cases of encephalitis in adults, usually with a significant respiratory illness at onset. Human infection with this virus has been documented in New York, Wisconsin, Ohio, Michigan, Ontario, and other areas of North America where the vector mosquito, *Aedes stimulans*, feeds on its main host, the white-tailed deer. Tahyña virus can be found in central Europe, Russia, China, and Africa. The virus is a prominent cause of febrile disease but can also cause pharyngitis, pulmonary syndromes, aseptic meningitis, or meningoencephalitis.

Flaviviruses The most important flavivirus encephalitides are Japanese encephalitis, St. Louis encephalitis, tick-borne encephalitis, and West Nile virus infection. Australian encephalitis (Murray Valley encephalitis) and Rocio virus infection resemble Japanese encephalitis but are documented only occasionally in Australia and Brazil, respectively. Powassan virus has caused ~50 cases of often-severe disease (lethality rate, ~10%), frequently occurring among children in eastern Canada and the United States. Usutu virus has caused only individual cases of human infection, but such infections may be underdiagnosed.

JAPANESE ENCEPHALITIS Japanese encephalitis is the most important viral encephalitis in Asia. Each year 35,000–50,000 cases and more

than 15,000 deaths are reported. Japanese encephalitis virus is found throughout Asia, including far eastern Russia, Japan, China, India, Pakistan, and southeastern Asia, and causes occasional epidemics on western Pacific islands. The virus has been detected in the Torres Strait islands, and five human encephalitis cases have been identified on the nearby Australian mainland. The virus is particularly common in areas where irrigated rice fields attract the natural avian vertebrate hosts and provide abundant breeding sites for mosquitoes such as *Culex tritaeniorhynchus*, which transmit the virus to humans. Additional amplification by pigs, which suffer abortion, and horses, which develop encephalitis, may be significant as well. Vaccination of these additional amplifying hosts may reduce the transmission of the virus.

Clinical signs of Japanese encephalitis emerge after an incubation period of 5–15 days and range from an unspecific febrile presentation (nausea, vomiting, diarrhea, cough) to aseptic meningitis, meningoencephalitis, acute flaccid paralysis, and severe encephalitis. Common findings are cerebellar signs, cranial nerve palsies, and cognitive and speech impairments. A Parkinsonian presentation and seizures are typical in severe cases. Effective vaccines are available. Vaccination is indicated for summer travelers to rural Asia, where the risk of acquiring Japanese encephalitis is considered to be about 1 per 5000 to 1 per 20,000 travelers per week if travel duration exceeds 3 weeks. Usually two intramuscular doses of the vaccine are given 28 days apart, with the second dose administered at least 1 week prior to travel.

ST. LOUIS ENCEPHALITIS St. Louis encephalitis virus is transmitted between mosquitoes and birds. This virus causes a low-level endemic infection among rural residents of the western and central United States, where *Culex tarsalis* is the vector (see “Western Equine Encephalitis,” below). The more urbanized mosquitoes *Culex pipiens* and *Culex quinquefasciatus* have been responsible for epidemics resulting in hundreds or even thousands of cases in cities of the central and eastern United States. Most cases occur in June through October. The urban mosquitoes breed in accumulations of stagnant water and sewage with high organic content and readily feed on humans in and around houses at dusk. The elimination of open sewers and trash-filled drainage systems is expensive and may not be possible, but screening of houses and implementation of personal protective measures may be an effective approach to the prevention of infection. The rural mosquito vector is most active at dusk and outdoors; its bites can be avoided by modification of activities and use of repellents.

Disease severity increases with age. St. Louis encephalitis virus infections that result in aseptic meningitis or mild encephalitis are concentrated among children and young adults, while severe and fatal cases primarily affect the elderly. Infection rates are similar in all age groups; thus the greater susceptibility of older persons to disease is a biologic consequence of aging. St. Louis encephalitis has an abrupt onset after an incubation period of 4–21 days, sometimes following a prodrome, and begins with fever, lethargy, confusion, and headache. In addition, nuchal rigidity, hypotonia, hyperreflexia, myoclonus, and tremors are common. Severe cases can include cranial nerve palsies, hemiparesis, and seizures. Patients often report dysuria and may have viral antigen in urine as well as pyuria. The overall rate of lethality is generally ~7% but may reach 20% among patients >60 years of age. Recovery is slow. Emotional lability, difficulties with concentration and memory, asthenia, and tremors are commonly prolonged in older convalescent patients. The CSF of patients with St. Louis encephalitis usually contains tens to hundreds of leukocytes, with a lymphocytic predominance and a left shift. The CSF glucose concentration is normal in these patients.

TICK-BORNE VIRAL ENCEPHALITIS Tick-borne encephalitis viruses are currently subdivided into four groups: the western/European subtype (previously called central European encephalitis virus), the (Ural-) Siberian subtype (previously called Russian spring-summer encephalitis virus), the Far Eastern subtype, and the louping ill subtype (previously called louping ill virus or, in Japan, Negishi virus). Small mammals and grouse, deer, and sheep are the vertebrate amplifiers for these viruses, which are transmitted by ticks. The risk of infection varies by geographic area and can be highly localized within a given area; human

1316 infections usually follow either outdoor activities resulting in tick bites or consumption of raw (unpasteurized) milk from infected goats or, less commonly, from other infected animals (cows, sheep). Milk seems to represent the main transmission route for louping ill viruses, which cause disease only very rarely. The western/European subtype viruses are transmitted mainly by *Ixodes ricinus* from Scandinavia to the Ural Mountains. (Ural-)Siberian viruses are transmitted predominantly by *Ixodes persulcatus* from Europe across the Ural Mountains to the Pacific Ocean; louping ill viruses seem to be confined primarily to Great Britain. Several thousand infections with tick-borne encephalitis virus are recorded each year among people of all ages. Human tick-borne viral encephalitis occurs between April and October, with a peak in June and July.

Western/European viruses classically caused bimodal disease. After an incubation period of 7–14 days, the illness begins with a *fever-myalgia phase* (arthralgia, fever, headaches, myalgia, nausea) that lasts for 2–4 days and is thought to correlate with viremia. A subsequent remission for several days is followed by the recurrence of fever and the onset of meningeal signs. The *CNS phase* (7–10 days before onset of improvement) varies from mild aseptic meningitis, which is more common among younger patients, to severe (meningo-)encephalitis with coma, seizures, tremors, and motor signs. Spinal and medullary involvement can lead to typical limb-girdle paralysis and respiratory paralysis. Most patients with western/European virus infections recover (lethality rate, 1%), and only a minority of patients have significant deficits. However, the lethality rate from (Ural-)Siberian virus infections reaches 7–8%.

Infections with Far Eastern viruses generally run a more abrupt course. The encephalitic syndrome caused by these viruses sometimes begins without a remission from the fever-myalgia phase and has more severe manifestations than the western/European syndrome. The lethality rate is high (20–40%), and major sequelae—most notably, lower motor neuron paralyzes of the proximal muscles of the extremities, trunk, and neck—are common, developing in approximately one-half of patients. Thrombocytopenia sometimes develops during the initial febrile illness, resembling the early hemorrhagic phase of some other tick-borne flavivirus infections, such as Kyasanur Forest disease. In the early stage of the illness, virus may be isolated from the blood. In the CNS phase, IgM antibodies are detectable in serum and/or CSF.

Diagnosis of tick-borne viral encephalitis primarily relies on serology and detection of viral genomes by RT-PCR. There is no specific therapy for infection. However, effective alum-adjuvanted, formalin-inactivated virus vaccines are produced in Austria, Germany, and Russia in chicken embryo cells (FSME-Immun® and Encepur®). Two doses of the Austrian vaccine separated by an interval of 1–3 months appear to be effective in the field, and antibody responses are similar when vaccine is given on days 0 and 14. Because rare cases of postvaccination Guillain-Barré syndrome have been reported, vaccination should be reserved for persons likely to experience rural exposure in an endemic area during the season of transmission. Cross-neutralization for the western/European and Far Eastern variants has been established, but there are no published field studies on cross-protection among formalin-inactivated vaccines.

Because 0.2–4% of ticks in endemic areas may be infected, the use of immunoglobulin prophylaxis of tick-borne viral encephalitis has been raised. Prompt administration of high-titered specific antibody preparations should probably be undertaken, although no controlled data are available to prove the efficacy of this measure. Immunoglobulins should be considered because of the risk of antibody-mediated enhancement of infection or antigen-antibody complex deposition in tissues.

WEST NILE VIRUS INFECTION West Nile virus is now the primary cause of arboviral encephalitis in the United States. In 2012, 2873 cases of neuroinvasive disease (e.g., meningitis, encephalitis, acute flaccid paralysis), with 270 deaths, and 2801 cases of non-neuroinvasive infection were reported. West Nile virus was initially described as being transmitted among wild birds by *Culex* mosquitoes in Africa, Asia, and southern Europe. In addition, the virus has been implicated in severe and fatal

hepatic necrosis in Africa. West Nile virus was introduced into New York City in 1999 and subsequently spread to other areas of the northeastern United States, causing die-offs among crows, exotic zoo birds, and other birds. The virus has continued to spread and is now found in almost all states as well as in Canada, Mexico, South America, and the Caribbean islands. *C. pipiens* remains the major vector in the northeastern United States, but several other *Culex* species and *A. albopictus* are also involved. Jays compete with crows and other corvids as amplifiers and lethal targets in other areas of the country.

West Nile virus is a common cause of febrile disease without CNS involvement (incubation period, 3–14 days), but it occasionally causes aseptic meningitis and severe encephalitis, particularly among the elderly. The fever-myalgia syndrome caused by West Nile virus differs from that caused by other viruses in terms of the frequent—rather than occasional—appearance of a maculopapular rash concentrated on the trunk (especially in children) and the development of lymphadenopathy. Back pain, fatigue, headache, myalgia, retroorbital pain, sore throat, nausea and vomiting, and arthralgia (but not arthritis) are common accompaniments that may persist for several weeks. Encephalitis, sequelae, and death are all more common among elderly, diabetic, and hypertensive patients and among patients with previous CNS insults. In addition to the more severe motor and cognitive sequelae, milder findings may include tremor, slight abnormalities in motor skills, and loss of executive functions. Intense clinical interest and the availability of laboratory diagnostic methods have made it possible to define a number of unusual clinical features. Such features include chorioretinitis, flaccid paralysis with histologic lesions resembling poliomyelitis, and initial presentation with fever and focal neurologic deficits in the absence of diffuse encephalitis. Immunosuppressed patients may have fulminant courses or develop persistent CNS infection. Virus transmission through both transplantation and blood transfusion has necessitated screening of blood and organ donors by nucleic acid-based tests. Occasionally, pregnant women infect their fetuses with West Nile virus.

Rhabdoviruses: Chandipura Virus Infection Chandipura virus seems to be an emerging and increasingly important human virus in India, where it is transmitted among hedgehogs by mosquitoes and sandflies. In humans, the disease begins as an influenza-like illness, with fever, headache, abdominal pain, nausea, and vomiting; these manifestations are followed by neurologic impairment and infection-related or autoimmune-mediated encephalitis. Chandipura virus infection is characterized by high lethality in children. Several hundred cases of infection are recorded in India every year. Infections with other arthropod-borne rhabdoviruses (Isfahan, Piry, vesicular stomatitis Indiana, vesicular stomatitis New Jersey) may imitate the early febrile stage of Chandipura virus infection.

Togaviruses • EASTERN EQUINE ENCEPHALITIS This disease is encountered primarily in swampy foci along the eastern coast of the United States, with a few inland foci as far removed as Michigan. Infected humans present for medical care from June through October. During this period, the bird-*Culiseta* mosquito cycle spills over into other mosquitoes such as *Aedes sollicitans* or *Aedes vexans*, which are more likely to feed on mammals. There is concern over the potential role of the introduced anthropophilic mosquito species *A. albopictus*, which has been found to be infected with eastern equine encephalitis virus and is an effective experimental vector in the laboratory. Horses are a common target for the virus. Contact with unvaccinated horses may be associated with human disease, but horses probably do not play a significant role in amplification of the virus.

Eastern equine encephalitis is one of the most destructive of the arboviral diseases, with a sudden onset after an incubation period of ~5–10 days, rapid progression, 50–75% lethality, and frequent sequelae in survivors. This severity is reflected in the extensive necrotic lesions and polymorphonuclear infiltrates found at postmortem examination of the brain. Acute polymorphonuclear CSF pleocytosis, often occurring during the first 1–3 days of disease, is another indication of severity. In addition, leukocytosis with a left shift is a common feature. A formalin-inactivated vaccine has been used to protect laboratory workers but is not generally available or applicable.

VENEZUELAN EQUINE ENCEPHALITIS Venezuelan equine encephalitis viruses are separated into epizootic viruses (subtypes IA/B and IC) and enzootic viruses (subtypes ID, IE, and IF). Closely related enzootic viruses are Everglades virus, Mucambo virus, and Tonate virus. Epizootic viruses are found primarily in humid tropical-forest habitats and are maintained between culicoid mosquitoes and rodents. These viruses cause human disease but are not pathogenic for horses and do not cause epizootics. Enzootic viruses are common causes of acute febrile disease. Everglades virus has caused encephalitis in humans in Florida. Extrapolation from the rate of genetic change suggests that Everglades virus may have been introduced into Florida <200 years ago. Everglades virus is most closely related to the ID subtype viruses that appear to have given evolutionary rise to the epizootic variants active in South America.

Epizootic viruses have an unknown natural cycle but periodically cause extensive epizootics/epidemics in equids and humans in the Americas. These epizootics/epidemics are the result of high-level viremia in horses and mules, which transmit the infection to several types of mosquitoes. Infected mosquitoes in turn infect humans and perpetuate virus transmission. Humans also have high-level viremia, but their role in virus transmission is unclear. Epizootics of Venezuelan equine fever occurred repeatedly in South America at intervals of ≤10 years from the 1930s until 1969, when a massive epizootic spread throughout Central America and Mexico, reaching southern Texas in 1971. Genetic sequencing suggested that the virus from that outbreak originated from residual “un-inactivated” IA/B subtype virus in veterinary vaccines. The outbreak was terminated in Texas with a live attenuated vaccine (TC-83) originally developed for human use by the U.S. Army; the epizootic virus was then used for further production of inactivated veterinary vaccines. No further epizootic disease was identified until 1995, when additional epizootics took place in Colombia, Venezuela, and Mexico. The viruses involved in these epizootics as well as previously epizootic IC viruses are close phylogenetic relatives of known enzootic ID viruses. This finding suggests that active evolution and selection of epizootic viruses are under way in South America.

During epizootics, extensive human infection is the rule, with clinical disease in 10–60% of infected individuals. Most infections result in notable acute febrile disease, while relatively few infections (5–15%) result in neurologic disease. A low rate of CNS invasion is supported by the absence of encephalitis among the many infections resulting from exposure to aerosols in the laboratory setting or from vaccination accidents. The most recent large epizootic of Venezuelan equine fever occurred in Colombia and Venezuela in 1995; of the more than 85,000 clinical cases, 4% (with a higher proportion among children than adults) included neurologic symptoms/signs, and 300 cases ended in death.

The prevention of epizootic Venezuelan equine fever depends on vaccination of horses with the attenuated TC-83 vaccine or with an inactivated vaccine prepared from that variant. Enzootic viruses are genetically and antigenically different from epizootic viruses, and protection against the former with vaccines prepared from the latter is relatively ineffective. Humans can be protected by immunization with similar vaccines prepared from Everglades virus, Mucambo virus, and Venezuelan equine encephalitis virus, but the use of the vaccines is restricted to laboratory personnel because of reactogenicity, possible fetal pathogenicity, and limited availability.

WESTERN EQUINE ENCEPHALITIS The primary maintenance cycle of western equine encephalitis virus in the United States is between *C. tarsalis* and birds, principally sparrows and finches. Equids and humans become infected, and both suffer encephalitis without amplifying the virus in nature. St. Louis encephalitis virus is transmitted in a similar cycle in the same regions harboring western equine encephalitis virus; disease caused by the former occurs about a month earlier than that caused by the latter (July through October). Large epidemics of western equine encephalitis occurred in the western and central United States and Canada during the 1930s through 1950s, but in recent years the disease has been uncommon. From 1964 through 2010, only 640 cases were reported in the United States. This decline in incidence may reflect in part the integrated approach to mosquito management that

has been employed in irrigation projects and the increasing use of agricultural pesticides. The decreased incidence of western equine encephalitis almost certainly reflects the increased tendency for humans to be indoors behind closed windows at dusk—the peak biting period by the major vector.

After an incubation period of ~5–10 days, western equine encephalitis virus causes a typical diffuse viral encephalitis, with an increased attack rate and increased morbidity among the young, particularly children <2 years old. In addition, the lethality rate is high among the young and the very elderly (3–7% overall). One-third of individuals who have convulsions during the acute illness have subsequent seizure activity. Infants <1 year old—particularly those in the first months of life—are at serious risk of motor and intellectual damage. Twice as many males as females develop clinical encephalitis after 5–9 years of age. This difference in incidence may be related to greater outdoor exposure of boys to the vector but may also be due in part to biologic differences. A formalin-inactivated vaccine has been used to protect laboratory workers but is not generally available.

FEVER AND MYALGIA

The fever and myalgia syndrome is most commonly associated with zoonotic virus infection. Many of the numerous viruses listed in Table 233-1 probably cause at least a few cases of this syndrome, but only some of these viruses have prominent associations with the syndrome and are of biomedical importance. The fever and myalgia syndrome typically begins with the abrupt onset of fever, chills, intense myalgia, and malaise. Patients may also report joint or muscle pains, but true arthritis is not found. Anorexia is characteristic and may be accompanied by nausea or even vomiting. Headache is common and may be severe, with photophobia and retroorbital pain. Physical findings are minimal and are usually confined to conjunctival injection with pain on palpation of muscles or the epigastrium. The duration of symptoms/signs is quite variable (generally 2–5 days), with a biphasic course in some instances. The spectrum of disease varies from subclinical to temporarily incapacitating. Less constant findings include a nonpruritic maculopapular rash. Epistaxis may occur but does not necessarily indicate a bleeding diathesis. A minority of the patients may develop aseptic meningitis. This diagnosis is difficult to make in remote areas, given patients' photophobia and myalgia as well as the lack of opportunity to examine the CSF. Although pharyngitis or radiographic evidence of pulmonary infiltrates is found in some patients, the agents causing this syndrome are not primary respiratory pathogens.

The differential diagnosis includes anicteric leptospirosis, rickettsial diseases, and the early stages of other syndromes discussed in this chapter. The fever and myalgia syndrome is often described as “influenza-like,” but the usual absence of cough and coryza makes influenza an unlikely confounder except at the earliest stages. Treatment is supportive, but acetylsalicylic acid is avoided because of the potential for exacerbated bleeding or Reye's syndrome. Complete recovery is the general outcome for people with this syndrome, although prolonged asthenia and nonspecific symptoms have been described in some patients, particularly after infection with lymphocytic choriomeningitis virus or dengue virus types 1–4.

Efforts at prevention of viral infection are best based on vector control, which, however, may be expensive or impossible. For mosquito control, destruction of breeding sites is generally the most economically and environmentally sound approach. Emerging containment technologies include the release of genetically modified mosquitoes and the spread of *Wolbachia* bacteria to limit mosquito multiplication rates. Depending on the vector and its habits, other possible approaches include the use of screens or other barriers (e.g., permethrin-impregnated bed nets) to prevent the vector from entering dwellings, judicious application of arthropod repellents such as *N,N*-diethyltoluamide (DEET) to the skin, wearing of long-sleeved and ideally permethrin-impregnated clothing, and avoidance of the vectors' habitats and times of peak activity.

Arenaviruses Lymphocytic choriomeningitis/meningoencephalitis is the only human arenavirus infection resulting predominantly in fever

1318 and myalgia. Lymphocytic choriomeningitis virus is transmitted to humans from the common house mouse (*Mus musculus*) by aerosols of excreta and secreta. The virus is maintained in the mouse mainly by vertical transmission from infected dams. The vertically infected mouse remains viremic and sheds virus for life, with high concentrations of virus in all tissues. Infected colonies of pet hamsters also can serve as a link to humans. Infections among scientists and animal caretakers can occur because the virus is widely used in immunology laboratories as a model of T cell function and can silently infect cell cultures and passaged tumor lines. In addition, patients may have a history of residence in rodent-infested housing or other exposure to rodents. An antibody prevalence of ~5–10% has been reported among adults from Argentina, Germany, and the United States.

Lymphocytic choriomeningitis/meningoencephalitis differs from the general syndrome of fever and myalgia in that the onset is gradual. Conditions occasionally associated with the disease are orchitis, transient alopecia, arthritis, pharyngitis, cough, and maculopapular rash. An estimated one-fourth of patients (or fewer) experience a febrile phase of 3–6 days. After a brief remission, many develop renewed fever accompanied by severe headache, nausea and vomiting, and meningeal signs lasting for ~1 week (the CNS phase). These patients virtually always recover fully, as do the rare patients with clear-cut signs of encephalitis. Recovery may be delayed by transient hydrocephalus. During the initial febrile phase, leukopenia and thrombocytopenia are common, and virus can usually be isolated from blood. During the CNS phase, the virus may be found in the CSF, and antibodies are present in the blood. The pathogenesis of lymphocytic choriomeningitis/meningoencephalitis is thought to resemble that following direct intracranial inoculation of the virus into adult mice. The onset of the immune response leads to T cell–mediated immunopathologic meningitis. During the meningeal phase, CSF mononuclear-cell counts range from the hundreds to the low thousands per microliter, and hypoglycorrhachia is found in one-third of patients.

IgM-capture ELISA, immunochemistry, and RT-PCR are used in the diagnosis of lymphocytic choriomeningitis/meningoencephalitis. IgM-capture ELISA of serum and CSF usually yields positive results; RT-PCR assays have been developed for probing CSF. Because patients who have fulminant infections transmitted by recent organ transplantation do not mount an immune response, immunohistochemistry or RT-PCR is required for diagnosis. Infection should be suspected in acutely ill febrile patients with marked leukopenia and thrombocytopenia. In patients with aseptic meningitis, any of the following suggests lymphocytic choriomeningitis/meningoencephalitis: a well-marked febrile prodrome, adult age, occurrence in the autumn, low CSF glucose levels, or CSF mononuclear-cell counts of >1000/μL. In pregnant women, infection may lead to fetal invasion with consequent congenital hydrocephalus and chorioretinitis. Because the maternal infection may be mild, causing only a short febrile illness, antibodies to the virus should be sought in both the mother and the fetus under suspicious circumstances, particularly in TORCH (toxoplasmosis, rubella, cytomegalovirus, herpes simplex, and HIV)–negative neonatal hydrocephalus.

Bunyaviruses Numerous bunyaviruses cause fever and myalgia. Many of these viruses cause individual infections and usually do not result in epidemics—e.g., the viruses of the orthobunyavirus Anopheles A serogroup (e.g., Tacaiuma virus), Bwamba serogroup (Bwamba virus, Pongola virus), Guama serogroup (Catu virus, Guama virus), Nyando serogroup (Nyando virus), and Wyeomyia serogroup (Wyeomyia virus); the unclassified bunyavirus Tataguine virus; the phlebovirus Bhanja complex (Bhanja virus, Heartland virus) and Candiru complex (Alenquer, Candiru, Escharate, Maldonado, Morumbi, and Serra Norte viruses); the hantavirus Choclo virus; and the Dugbe and Nairobi sheep disease naiviruses. In the relevant orthobunyaviral Bunyamwera serogroup (Bunyamwera, Batai, Cache Valley, Fort Sherman, Germiston, Guaroa, Ilesha, Ngari, Shokwe, and Xingu viruses), Ngari virus recently has been implicated in a large epidemic in Africa.

ORTHOBUNYAVIRUS GROUP C SEROGROUP Apeú, Caraparú, Itaqui, Madrid, Marituba, Murutucú, Nepuyo, Oriboca, Ossa, Restan, and Zungarococha

viruses are among the most common causes of arboviral infection in humans entering South American jungles. These viruses cause acute febrile disease and are transmitted by mosquitoes in neotropical forests.

ORTHOBUNYAVIRUS SIMBU SEROGROUP Oropouche virus is transmitted in Central and South America by a biting midge, *Culicoides paraensis*, which often breeds to high density in cacao husks and other vegetable detritus found in towns and cities. Explosive epidemics involving thousands of patients have been reported from several towns in Brazil and Peru. Rash and aseptic meningitis have been detected in a number of patients. Iquitos virus, a recently discovered reassortant and close relative of Oropouche virus, causes disease that is easily mistaken for Oropouche virus disease; its overall epidemiologic significance remains to be determined.

PHLEBOVIRUS SANDFLY FEVER SEROGROUP A previous designation for sandfly fever, “3-day fever,” instructively describes the brief debilitating course associated with this essentially benign infection. There is neither a rash nor CNS involvement, and complete recovery is the rule. Sandfly fever is caused by at least six distinct phleboviruses of the phlebovirus sandfly fever serocomplex (Chagres virus, sandfly fever Cyprus virus, sandfly fever Naples virus, sandfly fever Sicilian virus, sandfly fever Turkey virus, and Toscana virus). Sandfly fever Naples virus, sandfly fever Sicilian virus, and Toscana viruses are the most important human pathogens of this group. *Phlebotomus* sandflies transmit the viruses, probably among small mammals, and infect humans by bites. Female sandflies may be infected by the oral route as they take a blood meal and may transmit the virus to offspring when they lay their eggs after a second blood meal. This prominent transovarial transmission confounds virus control.

Sandfly fever is found in the circum-Mediterranean area, extending to the east through the Balkans into parts of China as well as into western Asia. Chagres virus is endemic in Panama. Sandflies are found in both rural and urban settings and are known for their short flight ranges and their small sizes; the latter enables them to penetrate standard mosquito screens and netting. Epidemics have been described in the wake of natural disasters and wars. After World War II, extensive spraying in parts of Europe to control malaria greatly reduced sandfly populations and sandfly fever Naples virus transmission; the incidence of sandfly fever continues to be low.

A common pattern of disease in endemic areas consists of high attack rates among travelers and military personnel and little or no disease in the local population, who are protected after childhood infection. Toscana virus infection is common during the summer among rural residents and vacationers, particularly in Italy, Spain, and Portugal; a number of cases have been identified in travelers returning to Germany and Scandinavia. The disease may manifest as an uncomplicated febrile illness but is often associated with aseptic meningitis, with virus isolated from the CSF.

Punta Toro virus is a phlebovirus that is not part of the sandfly fever serocomplex but that, like the members of this complex, is transmitted by sandflies. Punta Toro virus causes a sandfly fever–like disease in the Latin American tropical forest, where the vectors rest on tree buttresses. Epidemics have not been reported, but antibody prevalence among inhabitants of villages in endemic areas indicates a cumulative lifetime exposure rate of >50%.

Flaviviruses The most clinically important flaviviruses that cause the fever and myalgia syndrome are dengue viruses 1–4. In fact, dengue is probably the most important arthropod-borne viral disease worldwide, with 50–100 million infections occurring per year. Year-round transmission of dengue viruses 1–4 occurs between latitudes of 25°N and 25°S, but seasonal forays of the viruses into the United States and Europe have been documented. All four viruses have *A. aegypti* as their principal vector. Through increasing spread of mosquitoes throughout the tropics and subtropics and international travel of infected humans, large areas of the world have become vulnerable to the introduction of dengue viruses. Thus, dengue and severe dengue (see “Viral Hemorrhagic Fevers,” below) are becoming increasingly common. For instance, conditions favorable to dengue virus 1–4 transmission via

A. aegypti exist in Hawaii and the southern United States. The range of a lesser dengue virus vector, *A. albopictus*, now extends from Asia to the United States, the Indian Ocean, parts of Europe, and Hawaii. *A. aegypti* typically breeds near human habitation, using relatively fresh water from sources such as water jars, vases, discarded containers, coconut husks, and old tires. The mosquito usually inhabits dwellings and bites during the day. Bursts of dengue cases are to be expected in the southern United States, particularly along the Mexican border, where containers of water may be infested with *A. aegypti*. Closed habitations with air-conditioning may inhibit transmission of many arboviruses, including dengue viruses 1–4.

Dengue begins after an incubation period averaging 4–7 days, when the typical patient experiences the sudden onset of fever, frontal headache, retroorbital pain, and back pain along with severe myalgias. These symptoms gave rise to the colloquial designation of dengue as “break-bone fever.” Often a transient macular rash appears on the first day, as do adenopathy, palatal vesicles, and scleral injection. The illness may last a week, with additional symptoms and clinical signs usually including anorexia, nausea or vomiting, and marked cutaneous hypersensitivity. Near the time of defervescence on days 3–5, a maculopapular rash begins on the trunk and spreads to the extremities and the face. Epistaxis and scattered petechiae are often noted in uncomplicated dengue, and preexisting gastrointestinal lesions may bleed during the acute illness.

Laboratory findings of dengue include leukopenia, thrombocytopenia, and, in many cases, elevations of serum aminotransferase concentrations. The diagnosis is made by IgM ELISA or paired serology during recovery or by antigen-detection ELISA or RT-PCR during the acute phase. Virus is readily isolated from blood in the acute phase if mosquito inoculation or mosquito cell culture is used.

Reoviruses Several orbiviruses (Lebombo, Kemerovo, Orungo, and Tribeč viruses) and coltivirus (Colorado tick fever, Eyach, and Salmon River viruses) can cause fever and myalgia in humans. With the exception of Lebombo and Orungo viruses, all of these viruses are transmitted by ticks. The most important reoviral arthropod-borne disease is Colorado tick fever. Several hundred patients with this disease are reported annually in the United States. The infection is acquired between March and November through the bite of an infected ixodid tick, the Rocky Mountain wood tick (*Dermacentor andersoni*), in mountainous western regions at altitudes of 1200–3000 m. Small mammals serve as amplifying hosts. The most common presentation is fever and myalgia; meningoencephalitis is not uncommon, and hemorrhagic disease, pericarditis, myocarditis, orchitis, and pulmonary presentations have also been reported. Rash develops in a minority of patients. Leukopenia and thrombocytopenia are also noted. The disease usually lasts 7–10 days and is often biphasic. The most important differential diagnostic considerations since the beginning of the twentieth century have been Rocky Mountain spotted fever (although Colorado tick fever is much more common in Colorado) and tularemia. Colorado tick fever virus replicates for several weeks in erythropoietic cells and can be found in erythrocytes. This feature, detected in erythroid smears stained by immunofluorescence, can be diagnostically helpful and is important during screening of blood donors.

PULMONARY DISEASE

Hantavirus pulmonary syndrome (HPS) was first described in 1993, but retrospective identification of cases by immunohistochemistry (1978) and serology (1959) support the idea that HPS is a recently discovered rather than a truly new disease. The causative agents are hantaviruses of a distinct phylogenetic lineage that is associated with the cricetid rodent subfamily Sigmodontinae. Sin Nombre virus, which chronically infects North American deer mice (*Peromyscus maniculatus*), is the most important agent of HPS in the United States. Several other related viruses (Anajatuba, Andes, Araraquara, Araucária, Bayou, Bermejo, Black Creek Canal, Blue River, Castelo dos Sonhos, El Moro Canyon, Juquitiba, Laguna Negra, Lechiguana, Maciel, Monongahela, Muleshoe, New York, Orán, Paranoá, Pergamino, Río

Mamoré, and Tunari) cause the disease in North and South America, but Andes virus is unusual in that it has been implicated in human-to-human transmission. HPS particularly affects rural residents living in dwellings permeable to rodent entry or working in occupations that pose a risk of rodent exposure. Each type of rodent has its own particular habits; in the case of deer mice, these behaviors include living in and around human habitation.

HPS begins with a prodrome of ~3–4 days (range, 1–11 days) comprising fever, malaise, myalgia, and—in many cases—gastrointestinal disturbances such as abdominal pain, nausea, and vomiting. Dizziness is common and vertigo occasional. Severe prodromal symptoms/signs may bring some patients to medical attention, but most cases are recognized as the pulmonary phase begins. Typical signs are slightly lowered blood pressure, tachycardia, tachypnea, mild hypoxemia, thrombocytopenia, and early radiographic signs of pulmonary edema. Physical findings in the chest are often surprisingly scant. The conjunctival and cutaneous signs of vascular involvement seen in hantavirus VHF (see below) are uncommon. During the next few hours, decompensation may progress rapidly to severe hypoxemia and respiratory failure.

The HPS differential diagnosis includes abdominal surgical conditions and pyelonephritis as well as rickettsial disease, sepsis, meningococemia, plague, tularemia, influenza, and relapsing fever. A specific diagnosis is best made by IgM antibody testing of acute-phase serum, which has yielded positive results even in the prodrome. Tests using a Sin Nombre virus antigen detect antibodies to the related HPS-causing hantaviruses. Occasionally, heterotypic viruses will react only in the IgG ELISA, but such a finding is highly suspicious given the very low seroprevalence of these viruses in normal populations. RT-PCR is usually positive when used to test blood clots obtained in the first 7–9 days of illness and when used to test tissues; this assay is useful in identifying the infecting virus in areas outside the home range of deer mice and in atypical cases.

During the prodrome, the differential diagnosis of HPS is difficult, but by the time of presentation or within 24 h thereafter, a number of diagnostically helpful clinical features become apparent. Cough usually is not present at the outset. Interstitial edema is evident on a chest x-ray. Later, bilateral alveolar edema with a central distribution develops in the setting of a normal-sized heart; occasionally, the edema is initially unilateral. Pleural effusions are often seen. Thrombocytopenia, circulating atypical lymphocytes, and a left shift (often with leukocytosis) are almost always evident; thrombocytopenia is a particularly important early clue. Hemoconcentration, hypoalbuminemia, and proteinuria should also be sought for diagnosis. Although thrombocytopenia virtually always develops and prolongation of the partial thromboplastin time is the rule, clinical evidence for coagulopathy or laboratory indications of disseminated intravascular coagulation (DIC) are found in only a minority of severely ill patients. Patients with severe illness also have acidosis and elevated serum lactate concentrations. Mildly increased values in renal function tests are common, but patients with severe HPS often have markedly elevated serum creatinine concentrations. Some New World hantaviruses other than Sin Nombre virus (e.g., Andes virus) have been associated with more kidney involvement, but few such cases have been studied.

Management of HPS during the first few hours after presentation is critical. The goal is to prevent severe hypoxemia by oxygen therapy, with intubation and intensive respiratory management if needed. During this period, hypotension and shock with increasing hematocrit invite aggressive fluid administration, but this intervention should be undertaken with great caution. Because of low cardiac output with myocardial depression and increased pulmonary vascular permeability, shock should be managed expectantly with pressors and modest infusion of fluid guided by pulmonary capillary wedge pressure. Mild cases can be managed by frequent monitoring and oxygen administration without intubation. Many patients require intubation to manage hypoxemia and developing shock. Extracorporeal membrane oxygenation is instituted in severe cases, ideally before the onset of shock. The procedure is indicated in patients who have a cardiac index of 2.3 L/min/m² or an arterial oxygen tension/fractional inspired oxygen (Pa_{O₂}/F_{I_{O₂}) ratio of}

1320 <50 and who are unresponsive to conventional support. Lethality rates remain at ~30–40% even with good management, but most patients surviving the first 48 h of hospitalization are extubated and discharged within a few days with no apparent long-term residua. The antiviral drug ribavirin inhibits hantaviruses in vitro but did not have a marked effect on patients treated in an open-label study.

VIRAL HEMORRHAGIC FEVER

VHF is a constellation of findings based on vascular instability and decreased vascular integrity. An assault, direct or indirect, on the microvasculature leads to increased permeability and (particularly when platelet function is decreased) to actual disruption and local hemorrhage (a positive tourniquet sign). Blood pressure is decreased, and in severe cases shock supervenes. Cutaneous flushing and conjunctival suffusion are examples of common, observable abnormalities in the control of local circulation. Hemorrhage occurs infrequently. In most patients, hemorrhage is an indication of widespread vascular damage rather than a life-threatening loss of blood volume. In some VHFs, specific organs may be particularly impaired. For instance, the kidneys are primary targets in hemorrhagic fever with renal syndrome (HFRS), and the liver is a primary target in yellow fever and filovirus diseases. However, in all of these diseases, generalized circulatory disturbance is critically important. The pathogenesis of VHF is poorly understood and varies among the viruses regularly implicated in the syndrome. In some viral infections, direct damage to the vascular system or even to parenchymal cells of target organs is an important factor; in other viral infections, soluble mediators are thought to play a major role in the development of hemorrhage or fluid redistribution.

The acute phase in most cases of VHF is associated with ongoing virus replication and viremia. VHFs begin with fever and myalgia, usually of abrupt onset. (Arenavirus infections are the exceptions as they often develop gradually.) Within a few days, the patient presents for medical attention because of increasing prostration that is often accompanied by abdominal or chest pain, anorexia, dizziness, severe headache, hyperesthesia, photophobia, and nausea or vomiting and other gastrointestinal disturbances. Initial examination often reveals only an acutely ill patient with conjunctival suffusion, tenderness to palpation of muscles or abdomen, and borderline hypotension or postural hypotension, perhaps with tachycardia. Petechiae (often best visualized in the axillae), flushing of the head and thorax, periorbital edema, and proteinuria are common. AST concentrations are usually elevated at presentation or within a day or two thereafter. Hemoconcentration from vascular leakage, which is usually evident, is most marked in HFRS and in severe dengue. The seriously ill patient progresses to more severe clinical signs and develops shock and other findings typical of the causative virus. Shock, multifocal bleeding, and CNS involvement (encephalopathy, coma, seizures) are all poor prognostic signs.

One of the major diagnostic clues to VHF is travel to an endemic area within the incubation period for a given syndrome. Except in infections with Seoul, dengue, and yellow fever viruses, which have urban hosts/vectors, travel to a rural setting is especially suggestive of a diagnosis of VHF. In addition, several diseases considered in the differential diagnosis—falciparum malaria, shigellosis, typhoid fever, leptospirosis, relapsing fever, and rickettsial diseases—are treatable and potentially lethal.

Early recognition of VHF is important because of the need for virus-specific therapy and supportive measures. Such measures include prompt, atraumatic hospitalization; judicious fluid therapy that takes into account the patient's increased capillary permeability; administration of cardiotoxic drugs; use of pressors to maintain blood pressure at levels that will support renal perfusion; treatment of the relatively common secondary bacterial (and the more rare fungal) infections; replacement of clotting factors and platelets as indicated; and the usual precautionary measures used in the treatment of patients with hemorrhagic diatheses. DIC should be treated only if clear laboratory evidence of its existence is found and if laboratory monitoring of therapy is feasible; there is no proven benefit of such therapy. The available evidence suggests that VHF patients have decreased cardiac output

and will respond poorly to fluid loading as it is often practiced in the treatment of shock associated with bacterial sepsis. Specific therapy is available for several of the VHFs. Strict barrier nursing and other precautions against infection of medical staff and visitors are indicated when VHFs are encountered except when the illness is due to dengue viruses, hantaviruses, Rift Valley fever virus, or yellow fever virus.

Novel VHF-causing agents are still being discovered. Besides the viruses listed below, the latest addition may be the unclassified rhabdovirus Bas-Congo virus, which has been associated with three cases of VHF in the Democratic Republic of the Congo. However, Koch's postulates have not yet been fulfilled to prove cause and effect.

Arenaviruses The most important arenaviruses causing VHF are Junín virus, Lassa virus, and Machupo virus. Chapare, Guanarito, Lujo, and Sabiá viruses have caused limited and/or infrequent outbreaks or individual cases.

JUNÍN/ARGENTINIAN AND MACHUPO/BOLIVIAN HEMORRHAGIC FEVERS These severe diseases (with fetal lethality rates reaching 15–30%) are caused by Junín virus and Machupo virus, respectively. Their clinical presentations are similar, but their epidemiology differs because of the distribution and behavior of the viruses' rodent reservoirs. Junín/Argentinian hemorrhagic fever has thus far been recorded only in rural areas of Argentina, whereas Machupo/Bolivian hemorrhagic fever seems to be confined to rural Bolivia. Infection with the causative agents almost always results in disease, and all ages and both sexes are affected. Person-to-person or nosocomial transmission is rare but has occurred. The transmission of Junín/Argentinian hemorrhagic fever from convalescing men to their wives suggests the need for counseling of patients with arenavirus hemorrhagic fever concerning the avoidance of intimate contacts for several weeks after recovery. Compared with the pattern in Lassa fever (see below), thrombocytopenia—often marked—is the rule, hemorrhage is common, and CNS dysfunction (e.g., marked confusion, tremors of the upper extremities and tongue, and cerebellar signs) is much more common in disease caused by Junín virus and Machupo virus. Some cases follow a predominantly neurologic course, with a poor prognosis.

The clinical laboratory is helpful in diagnosis since thrombocytopenia, leukopenia, and proteinuria are typical findings. Junín/Argentinian hemorrhagic fever is readily treated with convalescent-phase plasma given within the first 8 days of illness. In the absence of passive antibody therapy, IV ribavirin in the dose recommended for Lassa fever is likely to be effective in all the South American VHFs caused by arenaviruses. A safe, effective, live attenuated vaccine exists for Junín/Argentinian hemorrhagic fever. After vaccination of more than 250,000 high-risk persons in the endemic area, the incidence of this VHF decreased markedly. In experimental animals, this vaccine is cross-protective against Machupo/Bolivian hemorrhagic fever.

LASSA FEVER Lassa virus is known to cause endemic and epidemic disease in Nigeria, Sierra Leone, Guinea, and Liberia, although it is probably more widely distributed in western Africa. In countries where Lassa virus is endemic, Lassa fever can be a prominent cause of febrile disease. For example, in one hospital in Sierra Leone, laboratory-confirmed Lassa fever is consistently responsible for one-fifth of admissions to the medical wards. In western Africa alone, probably tens of thousands of Lassa virus infections occur annually. Lassa virus can be transmitted by close person-to-person contact. The virus is often present in urine during convalescence and is suspected to be present in seminal fluid early in recovery. Nosocomial spread has occurred but is uncommon if proper sterile parenteral techniques are used. All ages and both sexes are affected; the incidence of disease is highest in the dry season, but transmission takes place year-round.

Among the VHF agents, only arenaviruses are typically associated with a gradual onset of illness, which begins after an incubation period of 5–16 days. Hemorrhage is seen in only ~15–30% of Lassa fever patients; a maculopapular rash is often noted in light-skinned patients. Effusions are common, and male-dominant pericarditis may develop late. Maternal lethality is higher than the usual 15–30% and is especially increased during the last trimester. The fetal death rate

reaches 90%. Excavation of the uterus may increase survival rates of pregnant women, but data on Lassa fever and pregnancy are still sparse. These figures suggest that interruption of the pregnancy of Lassa virus–infected women should be considered. White blood cell counts are normal or slightly elevated, and platelet counts are normal or somewhat low. Deafness coincides with clinical improvement in ~20% of patients and is permanent and bilateral in some patients. Reinfection may occur but has not been associated with severe disease.

High-level viremia or a high serum AST concentration statistically predicts a fatal outcome. Thus, patients with an AST concentration of >150 IU/mL should be treated with IV ribavirin. This antiviral nucleoside analogue appears to be effective in reducing case–fatality rates from those documented among retrospective controls. However, possible side effects, such as reversible anemia (which usually does not require transfusion), dependent hemolytic anemia, and bone marrow suppression, need to be kept in mind. Ribavirin should be given by slow IV infusion in a dose of 32 mg/kg; this dose should be followed by 16 mg/kg every 6 h for 4 days and then by 8 mg/kg every 8 h for 6 days. Inactivated Lassa virus vaccines failed in preclinical studies.

Bunyaviruses The most important VHF-causing bunyaviruses are Crimean-Congo hemorrhagic fever virus, hantaviruses, Rift Valley fever virus, and “severe fever with thrombocytopenia syndrome virus.” Other bunyaviruses—e.g., the Garissa variant of Ngari virus and Ilesha virus—have caused sporadic VHF outbreaks in Africa.

CRIMEAN-CONGO HEMORRHAGIC FEVER (CCHF) This severe VHF has a wide geographic distribution, potentially emerging wherever virus-bearing ticks occur. Because of the propensity of CCHF virus–transmitting ticks to feed on domestic livestock and certain wild mammals, veterinary serosurveys are the most effective mechanism for the monitoring of virus circulation in a particular region. Human infections are acquired via tick bites or during the crushing of infected ticks. Domestic animals do not become ill but do develop viremia. Thus, there is risk of acquiring CCHF during sheep shearing, slaughter, and contact with infected hides or carcasses from recently slaughtered infected animals. Nosocomial epidemics are common and are usually related to extensive blood exposure or needlesticks.

Although generally similar to other VHFs, CCHF causes extensive liver damage, resulting in jaundice in some patients. Clinical laboratory values indicate DIC and show elevations in concentrations of AST, creatine phosphokinase, and bilirubin. Patients who do not survive generally have more distinct changes than survivors in the concentrations of these markers, even in the early days of illness, and also develop leukocytosis rather than leukopenia. In addition, thrombocytopenia is more marked and develops earlier in patients who do not survive than in survivors. The benefit of IV ribavirin for treatment remains hotly debated, but clinical experience and retrospective comparison of patients with ominous clinical laboratory values suggest that ribavirin may be efficacious. No human or veterinary vaccines are recommended.

HEMORRHAGIC FEVER WITH RENAL SYNDROME HFRS is the most important VHF today, with more than 100,000 cases of severe disease in Asia annually and milder infections numbering in the thousands in Europe. The disease is widely distributed in Eurasia. The major causative viruses are Puumala virus (Europe), Dobrava-Belgrade virus (the Balkans), and Hantaan virus (eastern Asia). Amur/Soochong, Gou, Kurkino, Muju, Saaremaa, Sochi, and Tula viruses also cause HFRS but much less frequently and in more geographically confined areas determined by the distribution of reservoir hosts. Seoul virus is exceptional in that it is associated with brown rats (*Rattus norvegicus*); therefore, the virus has a worldwide distribution because of the migration of these rodents on ships. Despite the wide distribution of Seoul virus, only mild or moderate HFRS occurs in Asia, and human disease has been difficult to identify in many areas of the world. Most cases of HFRS occur in rural residents or vacationers; the exception is Seoul virus infection, which may be acquired in an urban or rural setting or from contaminated laboratory-rat colonies. Classic Hantaan virus infection

in Korea and in rural China is most common in the spring and fall and is related to rodent density and agricultural practices. Human infection is acquired primarily through aerosols of rodent urine, although virus is also present in rodent saliva and feces. Patients with HFRS are not infectious.

Severe cases of HFRS evolve in four identifiable stages. The *febrile stage* lasts 3 or 4 days and is identified by the abrupt onset of fever, headache, severe myalgia, thirst, anorexia, and often nausea and vomiting. Photophobia, retroorbital pain, and pain on ocular movement are common, and the vision may become blurred with ciliary body inflammation. Flushing over the face, the V area of the neck, and the back is characteristic, as are pharyngeal injection, periorbital edema, and conjunctival suffusion. Petechiae often develop in areas of pressure, the conjunctivae, and the axillae. Back pain and tenderness to percussion at the costovertebral angle reflect massive retroperitoneal edema. Laboratory evidence of mild to moderate DIC is present. Other laboratory findings of HFRS include proteinuria and active urinary sediment. The *hypotensive stage* lasts from a few hours to 48 h and begins with falling blood pressure and sometimes shock. The relative bradycardia typical of the febrile phase is replaced by tachycardia. Kinin activation is marked. The rising hematocrit reflects increasing vascular leakage. Leukocytosis with a left shift develops, and thrombocytopenia continues. Atypical lymphocytes—which in fact are activated CD8+ and, to a lesser extent, CD4+ T cells—circulate. Proteinuria is marked, and the urine’s specific gravity falls to 1.010. Renal circulation is congested and compromised from local and systemic circulatory changes resulting in necrosis of tubules, particularly at the corticomedullary junction, and oliguria. During the *oliguric stage*, hemorrhagic tendencies continue, probably in large part because of uremic bleeding defects. Oliguria persists for 3–10 days before the return of renal function marks the onset of the *polyuric stage* (diuresis and hyposthenuria), which carries the danger of dehydration and electrolyte abnormalities.

Mild cases of HFRS may be much less stereotypical. The presentation may include only fever, gastrointestinal abnormalities, and transient oliguria followed by hyposthenuria. Infections with Puumala virus, the most common cause of HFRS in Europe (*nephropathia epidemica*), result in a much-attenuated picture but the same general presentation. Bleeding manifestations are found in only 10% of patients, hypotension rather than shock is usually documented, and oliguria is present in only about half of patients. The dominant features may be fever, abdominal pain, proteinuria, mild oliguria, and sometimes blurred vision or glaucoma followed by polyuria and hyposthenuria in recovery. The lethality rate is <1%.

HFRS should be suspected in patients with rural exposure in an endemic area. Prompt recognition of the disease permits rapid hospitalization and expectant management of shock and renal failure. Useful clinical laboratory parameters include leukocytosis, which may be leukemoid and is associated with a left shift; thrombocytopenia; and proteinuria. HFRS is readily diagnosed by an IgM-capture ELISA that is positive at admission or within 24–48 h thereafter. The isolation of hantaviruses is difficult, but RT-PCR of a blood clot collected early in the clinical course or of tissues obtained postmortem should give positive results. Such testing is usually undertaken if definitive identification of the infecting virus is required.

Mainstays of therapy are management of shock, reliance on vasopressors, modest crystalloid infusion, IV human serum albumin administration, and treatment of renal failure with prompt dialysis to prevent overhydration that may result in pulmonary edema and to control hypertension that increases the possibility of intracranial hemorrhage. Use of IV ribavirin has reduced lethality and morbidity in severe cases, provided treatment is begun within the first 4 days of illness. Lethality may be as high as 15% but with proper therapy should be <5%. Sequelae have not been definitively established.

RIFT VALLEY FEVER The natural range of Rift Valley fever virus was previously confined to sub-Saharan Africa, with circulation of the virus markedly enhanced by substantial rainfall. The El Niño Southern Oscillation phenomenon of 1997 facilitated subsequent spread of Rift

1322 Valley fever to the Arabian Peninsula, with epidemic disease in 2000. The virus has also been found in Madagascar and has been introduced into Egypt, where it caused major epidemics in 1977–1979, 1993, and thereafter. Rift Valley fever virus is maintained in nature by transovarial transmission in floodwater *Aedes* mosquitoes and presumably also has a vertebrate amplifier. Increased transmission during particularly heavy rains leads to epizootics characterized by high-level viremia in cattle, goats, or sheep. Numerous types of mosquitoes then feed on these animals and become infected, thereby increasing the possibility of human infections. Remote sensing via satellite can detect the ecological changes associated with high rainfall that predict the likelihood of Rift Valley fever virus transmission. High-resolution satellites can also detect the special depressions in floodwaters from which the mosquitoes emerge. In addition, the virus can be transmitted by contact with blood or aerosols from domestic animals. Transmission risk is therefore high during birthing, and both abortuses and placentas need to be handled with caution. Slaughtered animals are not infectious because anaerobic glycolysis in postmortem tissues results in an acidic environment that rapidly inactivates bunyaviruses. Neither person-to-person nor nosocomial transmission of Rift Valley fever has been documented.

Rift Valley fever virus is unusual in that it causes several clinical syndromes. Most infections are manifested as the fever–myalgia syndrome. A small proportion of infections result in VHF with especially prominent liver involvement. Renal failure and DIC are also common features. Perhaps 10% of otherwise mild infections lead to retinal vasculitis, and some patients have permanently impaired vision. Funduscopic examination reveals edema, hemorrhages, and infarction of the retina as well as optic nerve degeneration. In a small proportion of patients (<1 in 200), retinal vasculitis is followed by viral encephalitis.

No proven therapy exists for Rift Valley fever. Both retinal disease and encephalitis occur after the acute febrile syndrome has resolved and serum neutralizing antibody has developed—events suggesting that only supportive care need be given. Epidemic disease is best prevented by vaccination of livestock. The ability of this virus to propagate after introduction into Egypt suggests that other potentially receptive areas, including the United States, should develop response plans. Rift Valley fever, like Venezuelan equine encephalitis, is likely to be controlled only with adequate stocks of an effective live attenuated vaccine, but such global stocks are unavailable. A formalin-inactivated vaccine confers immunity in humans, but quantities are limited and three injections are required; this vaccine is recommended for potentially exposed laboratory workers and for veterinarians working in sub-Saharan Africa. A new live attenuated vaccine, MP-12, is being tested in humans and may soon become available for general use. The vaccine is safe and licensed for use in sheep and cattle.

SEVERE FEVER WITH THROMBOCYTOPENIA SYNDROME This is a recently described tick-borne disease caused by a previously unknown and still-unclassified phlebovirus. Numerous human infections have been reported during the past few years from China, and several cases have also been detected in Japan and South Korea. The clinical presentation ranges from mild nonspecific fever to severe VHF with a high (>12%) lethality rate.

Flaviviruses The most important flaviviruses that cause VHF are the mosquito-borne dengue viruses 1–4 and yellow fever viruses. These viruses are widely distributed and cause tens to hundreds of thousands of infections each year. Kyasanur Forest disease virus and Omsk hemorrhagic fever virus are geographically very restricted but important tick-borne flaviviruses that cause VHF, sometimes with subsequent viral encephalitis. Tick-borne encephalitis virus has caused VHF in a few patients. There is currently no therapy for these VHFs, but an inactivated vaccine has been used in India to prevent Kyasanur Forest disease.

SEVERE DENGUE Several weeks after convalescence from infection with dengue virus 1, 2, 3, or 4, the transient protection conferred by that infection against reinfection with a heterotypic dengue virus usually

wanes. Heterotypic reinfection may result in classic dengue or, less commonly, in severe dengue. In the past 20 years, *A. aegypti* has progressively reinvaded Latin America and other areas, and frequent travel by infected individuals has introduced multiple variants of dengue viruses 1–4 from many geographic areas. Thus the pattern of hyperendemic transmission of multiple dengue virus serotypes established in the Americas and the Caribbean has led to the emergence of severe dengue as a major problem. Among the millions of dengue virus 1–4 infections, ~500,000 cases of severe dengue occur annually, with a lethality rate of ~2.5%. The induction of vascular permeability and shock depends on multiple factors, such as the presence or absence of enhancing and nonneutralizing antibodies, age (susceptibility to severe dengue drops considerably after 12 years of age), sex (females are more often affected than males), race (whites are more often affected than blacks), nutritional status (malnutrition is protective), or sequence of infections (e.g., dengue virus 1 infection followed by dengue virus 2 infection seems to be more dangerous than dengue virus 4 infection followed by dengue virus 2 infection). In addition, considerable heterogeneity exists among each dengue virus population. For instance, Southeast Asian dengue virus 2 variants have more potential to cause severe dengue than do other variants.

Severe dengue is identified by the detection of bleeding tendencies (tourniquet test, petechiae) or overt bleeding in the absence of underlying causes, such as preexisting gastrointestinal lesions. Shock may result from increased vascular permeability. In milder cases of severe dengue, restlessness, lethargy, thrombocytopenia (<100,000/ μ L), and hemoconcentration are detected 2–5 days after the onset of typical dengue, usually at the time of defervescence. The maculopapular rash that often develops in dengue may also appear in severe dengue. In more severe cases, frank shock is apparent, with low pulse pressure, cyanosis, hepatomegaly, pleural effusions, and ascites; in some patients, severe ecchymoses and gastrointestinal bleeding develop. The period of shock lasts only 1 or 2 days.

A virologic diagnosis of severe dengue can be made by the usual means. However, multiple flavivirus infections result in broad immune responses to several members of the genus, and this situation may result in a lack of virus specificity of the IgM and IgG immune responses. A secondary antibody response can be sought with tests against several flavivirus antigens to demonstrate the characteristic wide spectrum of reactivity.

Most patients with shock respond promptly to close monitoring, oxygen administration, and infusion of crystalloid or—in severe cases—colloid. The case–fatality rates reported vary greatly with case ascertainment and quality of treatment; however, most patients with severe dengue respond well to supportive therapy, and the overall lethality rate at an experienced center in the tropics is probably as low as 1%.

The key to control of both dengue and severe dengue is the control of *A. aegypti*, which also reduces the risk of urban yellow fever and chikungunya virus circulation. Control efforts have been handicapped by the presence of nondegradable tires and long-lived plastic containers in trash repositories (perfect mosquito breeding grounds when filled with water during rainfall) and by insecticide resistance. Urban poverty and an inability of the public health community to mobilize the populace to respond to the need to eliminate mosquito breeding sites are also factors in lack of mosquito control. A tetravalent live attenuated dengue vaccine based on the attenuated yellow fever virus 17D platform is currently being evaluated in phase 3 clinical trials in Latin America, Asia, and Australia. At least two other live attenuated candidate vaccines based on modified recombinant dengue viruses have been evaluated in phase 1 clinical studies, but the results have not been promising.

YELLOW FEVER Yellow fever virus had caused major epidemics in Africa and Europe before its transmission by *A. aegypti* mosquitoes was discovered in 1900. Urban yellow fever became established in the New World as a result of colonization with *A. aegypti*, originally an African mosquito. Subsequently, different types of mosquitoes and nonhuman primates were found to maintain yellow fever virus in Africa and also in Central and South American jungles. Transmission to humans

is incidental, occurring via bites from mosquitoes that have fed on viremic monkeys. After the identification of *A. aegypti* mosquitoes as vectors of yellow fever, containment strategies were aimed at increased mosquito control. Today, urban yellow fever transmission occurs only in some African cities, but the threat exists in the great cities of South America, where reinfestation by *A. aegypti* has taken place and dengue virus 1–4 transmission by the same mosquito is common. Despite the existence of a highly effective and safe vaccine, several hundred jungle yellow fever cases occur annually in South America, and thousands of jungle and urban cases occur each year in Africa (29,000–60,000 estimated for 2013).

Yellow fever is a typical VHF accompanied by prominent hepatic necrosis. A period of viremia, typically lasting 3 or 4 days, is followed by a period of “intoxication.” During the latter phase in severe cases, characteristic jaundice, hemorrhages, black vomit, anuria, and terminal delirium occur, perhaps related in part to extensive hepatic involvement. Blood leukocyte counts may be normal or reduced and are often high in terminal stages. Albuminuria is usually noted and may be marked. As renal function fails in terminal or severe cases, the concentration of blood urea nitrogen rises proportionately. Abnormalities detected in liver function tests range from modest elevations of AST concentrations in mild cases to severe derangement.

Urban yellow fever can be prevented by the control of *A. aegypti*. The continuing sylvatic cycles require vaccination of all visitors to areas of potential transmission with live attenuated variant 17D vaccine virus, which cannot be transmitted by mosquitoes. With few exceptions, reactions to the vaccine are minimal; immunity is provided within 10 days and lasts for at least 25–35 years. An egg allergy mandates caution in vaccine administration. Although there are no documented harmful effects of the vaccine on fetuses, pregnant women should be immunized only if they are definitely at risk of exposure to yellow fever virus. Because vaccination has been associated with several cases of encephalitis in children <6 months of age, it is contraindicated in this age group, nor is it recommended for infants 6–8 months of age unless the risk of exposure is very high. Rare, serious, multisystemic adverse reactions (occasionally fatal) have been reported, particularly affecting the elderly, and risk-to-benefit should be weighed prior to vaccine administration to individuals ≥60 years of age. Nevertheless, the number of deaths of unvaccinated travelers with yellow fever exceeds the number of deaths from vaccination, and a liberal vaccination policy for travelers to involved areas should be pursued. Timely information on changes in yellow fever distribution and yellow fever vaccine requirements can be obtained from the U.S. Centers for Disease Control and Prevention (<http://www.cdc.gov/vaccines/vpd-vac/yf/default.htm>).

234 Ebola virus and Marburgvirus Infections

Jens H. Kuhn

Several viruses of the family Filoviridae cause severe and frequently fatal viral hemorrhagic fevers in humans. Introduction of filoviruses into human populations is an extremely rare event that most likely occurs by direct or indirect contact with healthy mammalian filovirus hosts or by contact with infected, sick, or deceased nonhuman primates. Filoviruses are highly infectious but not very contagious. Natural human-to-human transmission takes place through direct person-to-person (usually skin-to-skin) contact or exposure to infected bodily fluids and tissues; there is no evidence of such transmission by aerosol or respiratory droplets. Infections progress rapidly from influenza-like to hemorrhagic manifestations and typically culminate in multiple-organ dysfunction syndrome and shock. Treatment of

filovirus infections is of necessity entirely supportive because no specific efficacious antiviral agents or vaccines are yet available.

Filoviruses are categorized as World Health Organization (WHO) Risk Group 4 Pathogens. Consequently, all work with material suspected of containing filoviruses should be conducted only in maximal containment (biosafety level 4) laboratories. Experienced personnel handling these viruses must wear appropriate personal protective gear (see “Prevention,” below) and follow rigorous standard operating procedures. The proper authorities and WHO reference laboratories should be contacted immediately when filovirus infections are suspected.

ETIOLOGY

The family Filoviridae includes three genera: *Cuevavirus*, *Ebolavirus*, and *Marburgvirus* (Table 234-1 and Fig. 234-1). The available data suggest that the only known cuevavirus, Lloviu virus (LLOV), and one ebolavirus, Reston virus (RESTV), are not pathogenic for humans. The remaining four ebolaviruses—Bundibugyo virus (BDBV), Ebola virus (EBOV), Sudan virus (SUDV), and Tai Forest virus (TAFV)—cause Ebola virus disease (EVD; International Classification of Disease, Tenth Revision [ICD-10], code A98.4). The two marburgviruses, Marburg virus (MARV) and Ravn virus (RAVV), are the etiologic agents of Marburg virus disease (MVD; ICD-10 code A98.3).



Filoviruses have linear, nonsegmented, single-stranded, negative-sense RNA genomes that are ~19 kb in length. These genomes contain six or seven genes that encode the following seven structural proteins: nucleoprotein, polymerase cofactor (VP35), matrix protein (VP40), glycoprotein (GP_{1,2}), transcriptional cofactor (VP30), secondary matrix protein (VP24), and RNA-dependent RNA polymerase (L protein). Cuevaviruses and ebolaviruses, but not marburgviruses, also encode three nonstructural proteins of unknown function (sGP, ssGP, and Δ-peptide). Filovirions are unique among human virus particles in that they are predominantly pleomorphic filaments but also assume torus- or 6-like shapes (width, ~80 nm; average length, ≥790 nm). These enveloped virions contain helical ribonucleocapsids and are covered with GP_{1,2} spikes (Fig. 234-2).

TABLE 234-1 FILOVIRUS TAXONOMY

Current Taxonomy and Nomenclature	Previous Taxonomy and Nomenclature
Order Mononegavirales	Order Mononegavirales
Family Filoviridae	Family Filoviridae
Genus <i>Marburgvirus</i>	Genus <i>Marburgvirus</i>
Species <i>Marburg marburgvirus</i>	Species <i>Lake Victoria marburgvirus</i>
Virus 1: Marburg virus (MARV)	Virus: Lake Victoria marburgvirus (MARV)
Virus 2: Ravn virus (RAVV)	
Genus <i>Ebolavirus</i>	Genus <i>Ebolavirus</i>
Species <i>Tai Forest ebolavirus</i>	Species <i>Cote d'Ivoire ebolavirus</i> [sic ^a]
Virus: Tai Forest virus (TAFV)	Virus: Cote d'Ivoire ebolavirus [sic] (CIEBOV)
Species <i>Reston ebolavirus</i>	Species <i>Reston ebolavirus</i>
Virus: Reston virus (RESTV)	Virus: Reston ebolavirus (REBOV)
Species <i>Sudan ebolavirus</i>	Species <i>Sudan ebolavirus</i>
Virus: Sudan virus (SUDV)	Virus: Sudan ebolavirus (SEBOV)
Species <i>Zaire ebolavirus</i>	Species <i>Zaire ebolavirus</i>
Virus: Ebola virus (EBOV)	Virus: Zaire ebolavirus (ZEBOV)
Species <i>Bundibugyo ebolavirus</i>	
Virus: Bundibugyo virus (BDBV)	
Genus <i>Cuevavirus</i>	
Species <i>Lloviu cuevavirus</i>	
Virus: Lloviu virus (LLOV)	

^aThe correct spelling of the country for which this virus is named is Côte d'Ivoire. The lack of a circumflex in “Cote” in the virus designation produced a false country name. This fact is denoted by “[sic].”

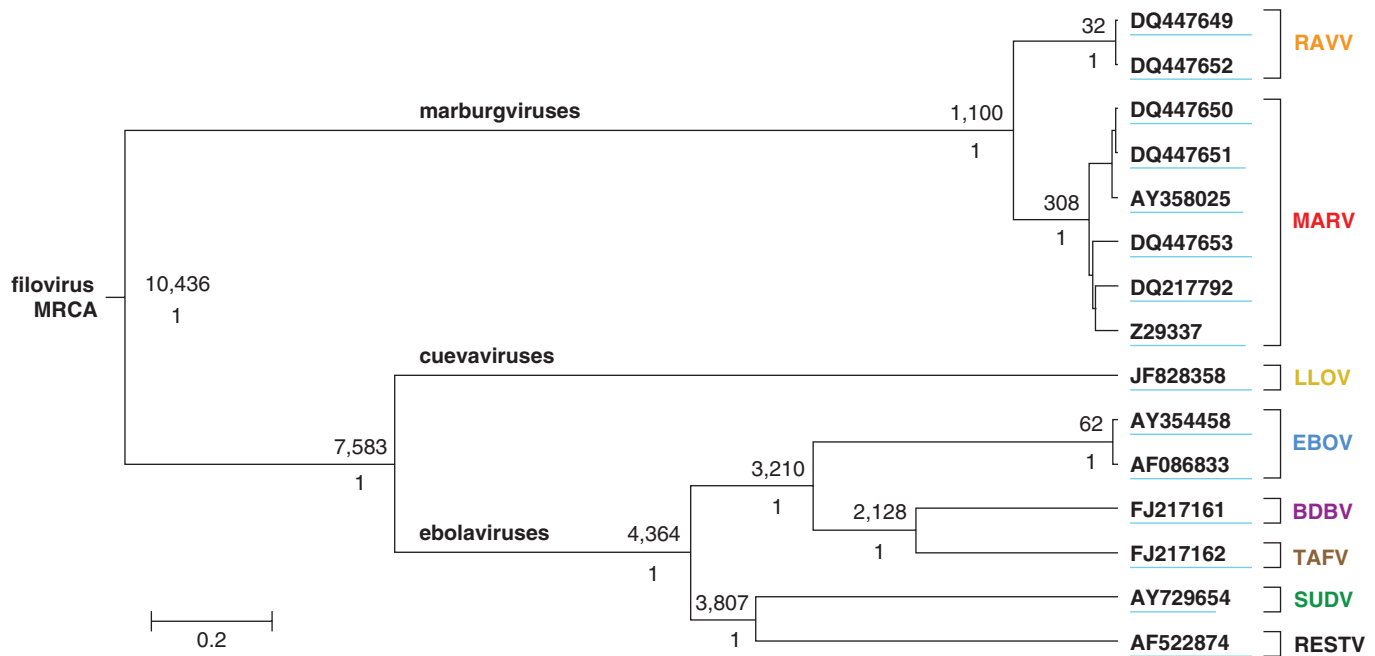


FIGURE 234-1 Filovirus phylogeny/evolution. Bayesian coalescent analysis of representative variants of all known filovirus clades (represented by underlined GenBank accession numbers). The maximal clade credibility tree is shown with the most recent common ancestor (MRCA) at each node. Posterior probability values are shown beneath MRCA estimates in years. Scale is in substitutions/site based on an analysis performed by Dr. Serena Carroll, Centers for Disease Control and Prevention. BDBV, Bundibugyo virus; EBOV, Ebola virus; LLOV, Lloviu virus; MARV, Marburg virus; RAVV, Ravn virus; RESTV, Reston virus; SUDV, Sudan virus; TAFV, Tai Forest virus.

EPIDEMIOLOGY



To date (i.e., as of December 3, 2014), a total of 20,012 human filovirus infections and 8058 fatalities have been recorded (Fig. 234-3). These numbers emphasize both the high degree of lethality (number of deaths per number of sick people; 40.3%) and the overall low mortality (impact on healthy population) of filovirus infections. At least for the moment, natural filovirus infections do not

pose a global threat. Filoviruses pathogenic for humans appear to be exclusively endemic to Equatorial Africa, although this distribution may change if natural or artificial environmental alterations lead to filovirus host migration and increased contacts between nonhuman hosts and humans (Fig. 234-4). The majority of recorded EVD and MVD outbreaks can be traced back to single index cases who transmitted the infection to others. These chains of contacts suggest that only around 50 natural host-to-human spillover events have occurred since the discovery of filoviruses in 1967. Outbreak frequency, case numbers, and overall lethality probably depend on the particular etiologic agent, the geographic location and socioeconomic conditions of the affected country, and local customs. In particular, the availability of personal protective gear and reusable medical equipment, such as syringes and needles, has affected overall case numbers in the past, and outbreaks have been contained when local burial practices, such as ritual washing, have been either prevented or altered by the use of gloves. The incidence of EVD and MVD may have increased over the past two decades (Figs. 234-3 and 234-4), but researchers debate whether the observed change is due to increased filovirus activity, more frequent contact between filovirus hosts and humans, or continuous improvement in surveillance capabilities.

EVD and MVD outbreaks are associated with distinct meteorologic and geographic conditions and are probably associated with distinct hosts or reservoirs. The four ebolaviruses that cause disease in humans are endemic in humid rainforests. EVD outbreaks have often been linked to hunting or contact with bush meat (i.e., meat from apes, other nonhuman primates, duikers, or bush pigs) in forests. Ecologic studies indicate that EBOV may be the etiologic agent of extensive and frequently fatal epizootics among wild chimpanzee and gorilla populations. However, replicating isolates of ebolaviruses from wild nonhuman primates have thus far been obtained only in the case of TAFV, which was isolated from a succumbed western chimpanzee in Côte d'Ivoire in 1994. The marburgviruses MARV and RAVV, on the other hand, seem to infect hosts inhabiting arid woodlands. MVD outbreaks have almost always been epidemiologically linked to visits to or work in natural or artificial caves or mines. A pteropid (fruit) bat, the cave-dwelling Egyptian rousette (*Rousettus aegyptiacus*), serves as a natural and subclinically infected reservoir for both MARV and RAVV. Although bats are suspected to be the hosts for ebolaviruses as



FIGURE 234-2 Ebola virus particle: the first transmission electron micrograph of an Ebola virion in a culture of Vero cells inoculated with a blood sample from a patient from the 1976 Zaire outbreak of Ebola virus disease. Shown is the typical and unique filamentous and pleomorphic structure of filovirions. (PHIL ID#1833, taken by Dr. Fredrick A. Murphy, Centers for Disease Control and Prevention.)

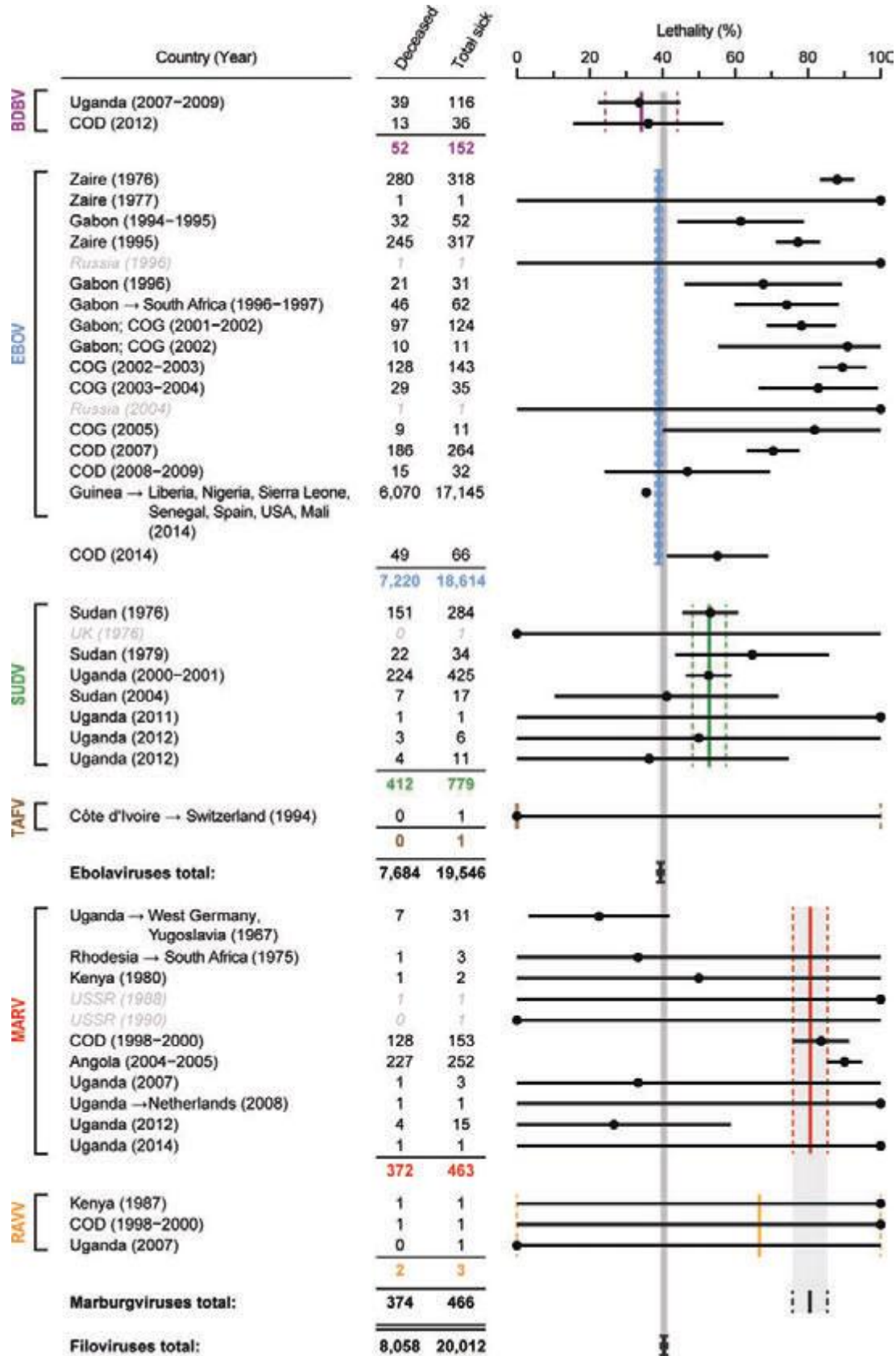


FIGURE 234-3 Characteristics of outbreaks of human filovirus disease. Six of eight known filoviruses have caused disease in humans in the past. Outbreaks are listed by virus in chronological order. Laboratory infections are shaded gray and italicized. Arrows indicate international case exportation. Total number of cases and total number of lethal cases are summarized in the middle column (2014 EBOV infections as of December 3). The lethality/case–fatality rate (black dots) for each outbreak is plotted on a 0–100% scale along with 99% confidence intervals (black horizontal lines). The overall case–fatality rate for disease caused by a particular virus is delineated by vertical bold-colored lines, with vertical bold-colored dashed lines indicating the corresponding 99% confidence intervals; the overall case–fatality rate for all ebolavirus infections, all marburgvirus infections, and all filovirus infections are shown by vertical gray bars. BDBV, Bundibugyo virus; COD, Democratic Republic of the Congo (formerly Zaire); COG, Republic of the Congo; EBOV, Ebola virus; MARV, Marburg virus; RAVV, Ravn virus; SUDV, Sudan virus; TAFV, Tai Forest virus; UK, United Kingdom; USSR, Union of Soviet Socialist Republics (today Russia).

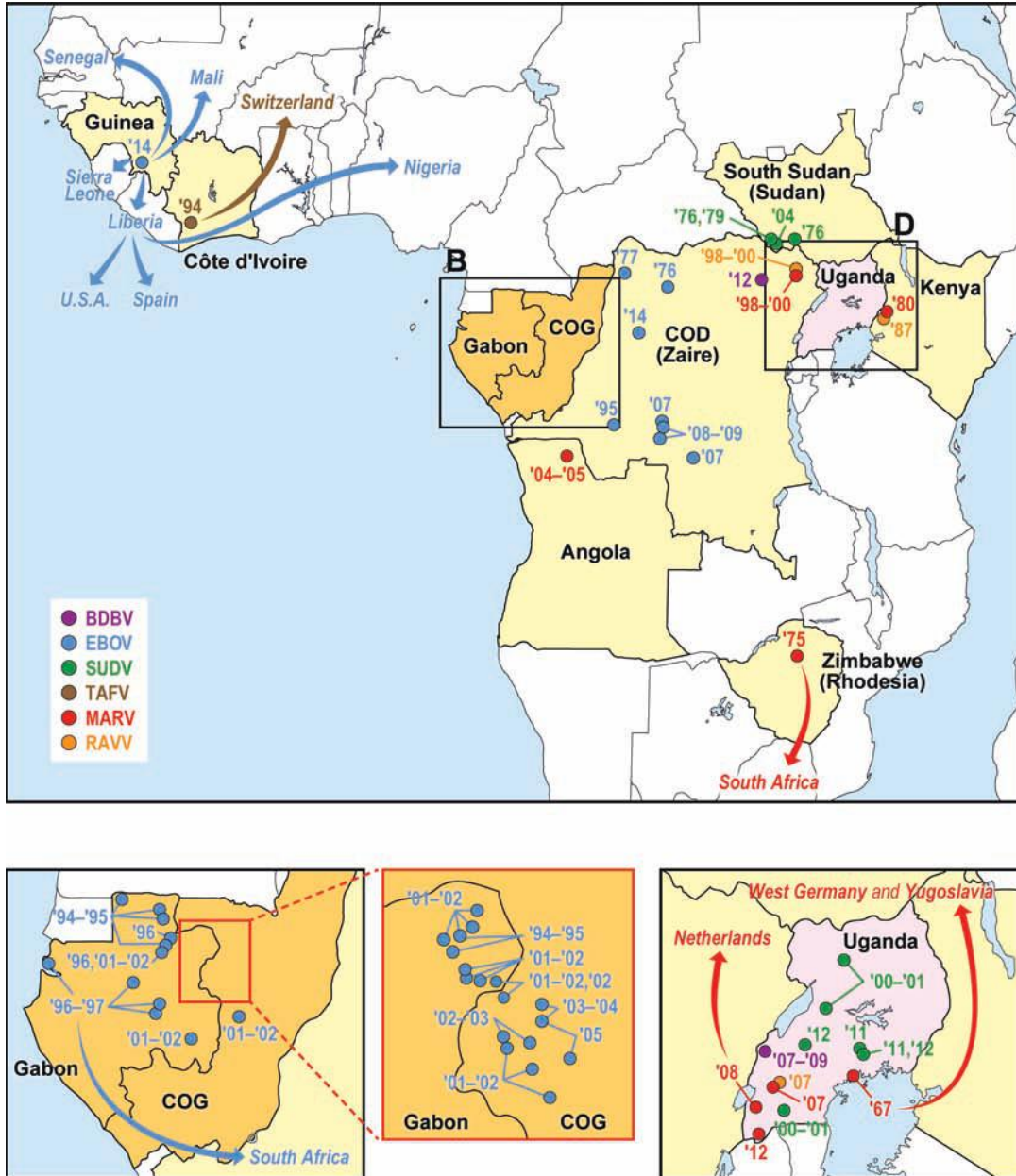


FIGURE 234-4 Geographic distribution of human filovirus disease outbreaks and years of occurrence. Arrows indicate international case exportation. BDBV, Bundibugyo virus; COD, Democratic Republic of the Congo (formerly Zaire); COG, Republic of the Congo; EBOV, Ebola virus; MARV, Marburg virus; RAVV, Ravn virus; SUDV, Sudan virus; TAFV, Tai Forest virus.

well, definitive proof is still lacking. In fact, thus far, only EBOV and RESTV have been loosely connected to frugivorous and insectivorous bats by means of antibody or genome fragment detection, whereas the hosts of BDBV, SUDV, and TAFV remain unclear.

PATHOGENESIS

Human infections typically occur through direct exposure of skin lesions or mucosal surfaces to contaminated bodily fluids or material or by parenteral inoculation (e.g., via accidental needlesticks or reuse of needles in poorly equipped hospitals). Numerous studies, both in vitro and in vivo (in several animal models of human disease), have shed light on key pathogenetic events that evolve subsequent to filovirion exposure. The GP_{1,2} spikes on the surface of filovirions determine their cell and tissue tropism by engaging yet-unidentified cell-surface molecules and the intracellular receptor Niemann-Pick C1.

One of the pathogenetic hallmarks of filovirus infection is a pronounced suppression of the immune system. The first targets of filovirions are local macrophages, monocytes, and dendritic cells.

Several structural proteins of filovirions, in particular VP35, VP40, and VP24, then suppress cellular innate immune responses by, for instance, inhibiting the interferon pathway and thereby enabling a productive filovirus infection. The result is the secretion of copious numbers of progeny virions, as evidenced by high titers in the bloodstream (>10⁶ plaque-forming units [pfu]/mL of serum in humans) and the lymphatics, and dissemination to most tissues. Filovirions then infect additional phagocytic cells, such as other macrophages (alveolar, peritoneal, pleural), Kupffer cells in the liver, and microglia, as well as other targets, such as adrenal cortical cells, fibroblasts, hepatocytes, endothelial cells, and a variety of epithelial cells. Infection leads to the secretion of soluble signaling molecules (varying with the cell type) that most likely are crucial factors in immune response modulation and development of multiorgan dysfunction syndrome. For instance, infected macrophages react by secreting proinflammatory cytokines, a response that leads to further recruitment of macrophages to the site of infection. In contrast, infected dendritic cells are not activated to secrete cytokines, and expression of major histocompatibility class II

antigens is partially suppressed. Immunosuppression occurs in part by massive lymphoid depletion in lymph nodes, spleen, and thymus in the absence of reactive inflammatory cellular responses. Results from animal studies suggest that depletion is a direct consequence of considerable bystander apoptosis of lymphocytes; this explanation would also account for the severe lymphopenia that develops in patients. The consequence of these events is not only florid filovirus dissemination but also a proclivity of the patient for secondary bacterial and fungal infections.

Other pathogenetic hallmarks of filovirus infections are a severe disturbance of the clotting system and the impairment of vascular integrity. Disseminated intravascular coagulation is the cause of the severe imbalance in the clotting system of filovirus-infected patients. Thrombocytopenia, increased concentrations of tissue factor, consumption of clotting factors, increased concentrations of fibrin degradation products (D-dimers), and declining concentrations of protein C are typical features of infection. Consequently, the occlusion of small vessels by widely distributed microthrombi leads to extensive necroses/hypoxic infarcts in target tissues (particularly the gonads,

kidneys, liver, and spleen) in the absence of marked inflammatory responses. In addition, petechiae, ecchymoses, extensive visceral effusions, and other hemorrhagic signs are observed in internal organs, mucous membranes, and skin. Actual severe blood loss, however, is a rare event. Aberrance in cytokines or other factors such as nitric oxide and direct infection and activation of endothelial cells most likely are responsible for upregulated permeability of the endothelia of blood vessels. This upregulation leads to fluid redistribution (*third spacing*); interstitial and myocardial edema and hypovolemic shock are common developments. Clinical improvement is rare and is usually characterized by falling viral titers during the development of a virus-specific immune response.

CLINICAL MANIFESTATIONS

MVD and EVD cannot be differentiated by mere observation of clinical manifestations. The incidence of clinical signs does not differ significantly among infections caused by disparate filoviruses (Table 234-2). The incubation period ranges from 3 to 25 days, after which infected people develop a biphasic syndrome with a 1- to 2-day

TABLE 234-2 DISTRIBUTION OF CLINICAL SIGNS/SYMPTOMS OF FILOVIRUS-INFECTED PATIENTS IN THREE REPRESENTATIVE OUTBREAKS

Sign/Symptom	Frequency (%) Among Survivors			Frequency (%) Among Fatal Cases		
	BDBV	EBOV	MARV	BDBV	EBOV	MARV
Abdominal pain	88	68	59	93	62	57
Abortion	NR	5	NR	NR	2	NR
Anorexia	83	47	77	80	43	72
Anuria	NR	0	NR	NR	7	NR
Arthralgia or myalgia	83	79	55	86	50	55
Asthenia	NR	95	NR	NR	85	NR
Bleeding from puncture sites	NR	5	0	NR	8	7
Bleeding from the gums	NR	0	23	NR	15	36
Bleeding from any site	NR	NR	59	NR	NR	71
Bloody stools	NR	5	NR	NR	7	NR
Chest pain	NR	5	18	NR	10	4
Conjunctival injection	NR	47	14	NR	42	42
Convulsions	NR	0	NR	NR	2	NR
Cough	NR	26	9	NR	7	5
Diarrhea	92	84	59	87	86	56
Difficulty breathing	26	NR	36	57	NR	58
Dysesthesia	NR	5	NR	NR	0	NR
Epistaxis	NR	0	18	NR	2	34
Fever	100	95	100	100	93	92
Headaches	84	74	73	93	52	79
Hearing loss	NR	11	NR	NR	5	NR
Hematemesis	NR	0	68	NR	13	76
Hematoma	NR	0	0	NR	2	3
Hematuria	NR	16	NR	NR	7	NR
Hemoptysis	NR	11	9	NR	0	4
Hepatomegaly (without jaundice)	NR	5	NR	NR	2	NR
Hiccups	17	5	18	40	17	44
Lumbar pain	NR	26	5	NR	12	8
Maculopapular rash	35	16	NR	33	14	NR
Malaise or fatigue	96	NR	86	100	NR	83
Melena	NR	16	41	NR	8	58
Nausea and vomiting	92	68	77	87	73	76
Petechiae	NR	0	9	NR	8	7
Sore throat, odynophagia, or dysphagia	43	58	43	60	56	43
Splenomegaly	NR	5	NR	NR	2	NR
Tachypnea	NR	0	NR	NR	31	NR
Tinnitus	NR	11	NR	NR	1	NR

Abbreviations: BDBV, Bundibugyo virus; EBOV, Ebola virus; MARV, Marburg virus; NR, not reported.

1328 relative remission separating the two phases. The first phase (disease onset until around day 5–7) resembles influenza and is characterized by sudden onset of fever and chills, severe headaches, cough, myalgia, pharyngitis, arthralgia of the larger joints, development of a maculopapular rash, and other signs/symptoms (Table 234-2). The second phase (approximately 5–7 days after disease onset and thereafter) involves the gastrointestinal tract (abdominal pain with vomiting and/or diarrhea), respiratory tract (chest pain, cough), vascular system (postural hypotension, edema), and central nervous system (confusion, coma, headache). Hemorrhagic manifestations such as subconjunctival injection, nosebleeds, hematemesis, hematuria, and melena are typical (Table 234-2).

Typical laboratory findings are leukopenia (with cell counts as low as 1000/ μ L) with a left shift prior to leukocytosis, thrombocytopenia (with counts as low as 50,000/ μ L), increased concentrations of liver and pancreatic enzymes (aspartate aminotransferase > alanine aminotransferase, γ -glutamyltransferase, serum amylase), hypokalemia, hypoproteinemia, increased creatinine and urea concentrations with proteinuria, and prolonged prothrombin and partial thromboplastin times.

Patients usually succumb to disease 4–14 days after infection. Patients who survive experience prolonged and sometimes incapacitating sequelae such as arthralgia, asthenia, iridocyclitis, hearing loss, myalgia, orchitis, parotitis, psychosis, recurrent hepatitis, transverse myelitis, or uveitis. Temporary hair loss and desquamation of skin areas previously affected by a typical maculopapular rash are visible consequences of the disease. Rarely, filoviruses can persist in the liver, eyes, or testicles of survivors and may cause recurrent disease months after convalescence.

DIAGNOSIS

Filovirus infections cannot be diagnosed on the basis of clinical presentation alone. Numerous diseases typical for Equatorial Africa need to be considered in the differential diagnosis of a febrile patient. Almost all of these diseases occur at a much higher incidence than filovirus infections and are therefore the more likely candidates during differential diagnostic deliberations. The most important of the infectious diseases that closely mimic EVD and MVD are falciparum malaria and typhoid fever; also important are enterohemorrhagic *Escherichia coli* enteritis, gram-negative septicemia (including shigellosis), meningococcal septicemia, rickettsial infections, fulminant viral hepatitis, leptospirosis, measles, and all other viral hemorrhagic fevers (in particular, yellow fever). Other ailments, such as venomous snakebites, warfarin intoxication, and the many transient or inherited platelet and vascular disorders, also must be considered. Visits to caves or mines and direct contact with bats, nonhuman primates (especially apes), or bush meat should raise suspicion of filovirus infection, as should admission to or treatment in rural hospitals or direct contact with severely ill local residents.

If EVD or MVD is suspected on the basis of epidemiologic history, exposure history, and/or clinical manifestations, infectious disease specialists and the proper public health authorities, including the WHO, should be notified immediately. Laboratory diagnosis of EVD and MVD is relatively straightforward but requires maximal containment (biosafety level 4), which usually is not available in filovirus-endemic countries, or the involvement of on-site personnel trained in the use of diagnostic assays adapted for field use. Consequently, diagnostic samples should be collected with great caution and with use of proper personal protective equipment and strict barrier nursing techniques. With adherence to established biosafety precautionary measures, samples should be sent in suitable transport media to national or international WHO reference laboratories. Acute-phase blood/serum is the preferred diagnostic specimen because it usually contains high titers of filovirions and filovirion-specific antibodies.

The current methods of choice for the diagnosis of filovirus infection are reverse-transcription polymerase chain reaction (detection limit, 1000–2000 virus genome copies per milliliter of serum) and antigen capture enzyme-linked immunosorbent assay (ELISA) for the detection of filovirus genomes and filovirion components,

respectively. Direct IgM and IgG or IgM capture ELISA is used for the detection of filovirion-targeting antibodies from patients in later stages of disease—i.e., those who have been able to mount a detectable immune response, including survivors. All these assays can be conducted on samples treated with guanidinium isothiocyanate (for polymerase chain reaction) or cobalt-60 irradiation (for ELISA) or subjected to other effective measures that render filoviruses noninfectious. Virus isolation in cell culture and plaque assays for quantification or diagnostic confirmation is relatively easy but must be performed in maximal-containment laboratories. If available, electron microscopic examination of properly inactivated samples or cultures can confirm the diagnosis because filovirions have unique filamentous shapes (Fig. 234-2). Formalin-fixed skin biopsies can be useful for safe postmortem diagnoses.

TREATMENT FILOVIRUS INFECTIONS

Any treatment of patients with suspected or confirmed filovirus infection must be administered under increased safety precautions by experienced specialists using appropriate personal protective equipment (see “Prevention,” below). Treatment of EVD and MVD is entirely supportive because no accepted/approved, efficacious, specific antiviral agents or vaccines are yet available. The one exception is hyperimmune equine immunoglobulin, which has been approved in Russia—in the absence of convincing efficacy data—for emergency treatment of laboratory infections. Given the extraordinarily high lethality of filoviruses, special protocols may be established by ad hoc expert groups to outline treatment of exposed individuals with one of several regimens that have shown promise in experimental nonhuman primates. Current options include postexposure vaccination with filovirus GP_{1,2}-expressing recombinant replicating vesicular stomatitis Indiana virus; administration of specific filovirus genome- or transcript-targeting small interfering RNAs or phosphorodiamidate morpholino oligomers; administration of filovirus-specific antibodies or antibody cocktails (convalescent sera have not yet been proven effective); and use of a synthetic adenosine analog (BCX4430) that acts as a non-obligate RNA chain terminator. In the absence of these candidate treatments, measures to stabilize patients include those generally recommended for severe septicemia/sepsis/shock. Countermeasures should address hypotension and hypoperfusion, vascular leakage in the systemic and pulmonary circulatory system, disseminated intravascular coagulation and overt hemorrhaging, acute kidney failure, and electrolyte (especially potassium) imbalances. Pain management and administration of antipyretics and antiemetics should always be considered.

COMPLICATIONS

Given the severe immunosuppression induced by filovirus infection, secondary infections should be kept in mind and appropriately treated as early as possible. Pregnancy and labor cause severe and frequently fatal complications in filovirus infections due to clotting factor consumption, fetal loss, and/or severe blood loss during birth.

PROGNOSIS

The prognosis of filovirus infections is generally poor, although outcome probably depends somewhat on which particular virus causes the infection (Fig. 234-3). Convalescence may take months, with skin peeling, alopecia, prostration, weight loss, orchitis, amnesia, confusion, and anxiety as typical sequelae. Rarely, filoviruses persist in apparently healthy survivors and are either reactivated by unknown means at a later point or transmitted sexually. Condom use or abstinence from sexual activity for at least 3 months after disappearance of clinical signs is therefore recommended for survivors.

CONTROL AND PREVENTION

Currently, filovirus vaccines are not available. Prevention of filovirus infection in nature is difficult because the ecology of the viruses is not completely understood. As stated above, frugivorous cave-dwelling

pteropid bats (Egyptian rousettes) have been identified as healthy carriers of MARV and RAVV. Avoidance of direct or indirect contact with these bats is therefore useful advice to people entering or living in areas where the animals can be found. Prevention seems to be more difficult in the case of ebolaviruses, for which definite reservoirs have not yet been pinpointed. EVD outbreaks have been associated not with bats but rather with hunting or consumption of nonhuman primates. The mechanism of introduction of ebolaviruses into nonhuman primate populations is unclear. Therefore, the best advice to locals and travelers is to avoid contact with bush meat, nonhuman primates, and bats.

Relatively simple barrier nursing techniques, vigilant use of proper personal protective equipment, and quarantine measures usually suffice to terminate or at least contain filovirus disease outbreaks. Isolation of filovirus-infected people and avoidance of direct person-to-person contact without proper personal protective equipment

usually suffice to prevent further spread as the pathogens are not transmitted through droplets or aerosols under natural conditions. Typical protective gear sufficient to prevent filovirus infections consists of disposable gloves, gowns, and shoe covers and a face shield and/or goggles. If available, N-95/N-100 respirators may be used to further limit infection risk. Positive air pressure respirators should be considered for high-risk medical procedures such as intubation or suctioning. Medical equipment used in the care of a filovirus-infected patient, such as gloves or syringes, should never be reused unless safety-tested sterilization or disinfection methods are properly applied. Because filovirions are enveloped, disinfection with detergents, such as 1% sodium deoxycholate, diethyl ether, or phenolic compounds, is relatively straightforward. Bleach solutions of 1:100 and 1:10 are recommended for surface disinfection and application to excreta/corpses, respectively. Whenever possible, potentially contaminated materials should be autoclaved, irradiated, or destroyed.

SECTION 16 FUNGAL INFECTIONS

235 Diagnosis and Treatment of Fungal Infections

John E. Edwards, Jr.

TERMINOLOGY AND MICROBIOLOGY

Traditionally, fungal infections have been classified into specific categories based on both anatomic location and epidemiology. The most common general anatomic categories are mucocutaneous and deep organ infection; the most common general epidemiologic categories are endemic and opportunistic infection. Although *mucocutaneous infections* can cause serious morbidity, they are rarely fatal. *Deep organ infections* also cause severe illness in many cases and, in contrast to mucocutaneous infections, are often fatal. The *endemic mycoses* (e.g., coccidioidomycosis) are caused by fungal organisms that are not part of the normal human microbiota but rather are acquired from environmental sources. In contrast, *opportunistic mycoses* are caused by organisms (e.g., *Candida* and *Aspergillus*) that commonly are components of the normal human microbiota and whose ubiquity in nature renders them easily acquired by the immunocompromised host (Table 235-1). Opportunistic fungi cause serious infections when the immunologic response of the host becomes ineffective, allowing the organisms to transition from harmless commensals to invasive pathogens. Frequently, the diminished effectiveness of the immune system is a result of advanced modern therapies that coincidentally either cause an imbalance in the host's microbiota or directly interfere with immunologic responses. Endemic mycoses cause more severe

illness in immunocompromised patients than in immunocompetent individuals.

Patients acquire deep organ infection with endemic fungi almost exclusively by inhalation. Cutaneous infections result either from hematogenous dissemination or, more often, from direct contact with soil—the natural reservoir for the vast majority of endemic mycoses. The dermatophytic fungi may be acquired by human-to-human transmission, but the majority of infections result from environmental contact. In contrast, the opportunistic fungus *Candida* invades the host from normal sites of colonization, usually the mucous membranes of the gastrointestinal tract. In general, innate immunity is the primary defense mechanism against fungi. Although antibodies are formed during many fungal infections (and even during commensalism), they generally do not constitute the primary mode of host defense. Nevertheless, in selected infections, as discussed below, measurement of antibody titers may be a useful diagnostic test.

Three other terms frequently used in clinical discussions of fungal infections are *yeast*, *mold*, and *dimorphic fungus*. *Yeasts* are seen as rounded single cells or as budding organisms. *Candida* and *Cryptococcus* are traditionally classified as yeasts. *Molds* grow as filamentous forms called *hyphae* both at room temperature and in invaded tissue. *Aspergillus*, *Rhizopus* (the genus that causes mucormycosis, also known as zygomycosis), and fungi commonly infecting the skin to cause ringworm and related cutaneous conditions are classified as molds. Variations occur within this classification of yeasts and molds. For instance, when *Candida* infects tissue, both yeasts and filamentous forms may be present (except with *C. glabrata*, which forms only yeasts in tissue); in contrast, *Cryptococcus* exists only in yeast form. *Dimorphic* is the term used to describe fungi that grow as yeasts or large spherical structures in tissue but as filamentous forms at room temperature in the environment. Classified in this group are the organisms causing blastomycosis, paracoccidioidomycosis, coccidioidomycosis, histoplasmosis, and sporotrichosis.

The incidence of nearly all fungal infections has risen substantially. Opportunistic infections have increased in frequency as a consequence of intentional immunosuppression in organ and stem cell transplantation and other disorders, the administration of cytotoxic chemotherapy for cancers, the liberal use of antibacterial agents, and, more recently, the increasing use of monoclonal antibodies.


 Within a global context, the incidence of endemic mycoses has increased in geographic locations where there has been substantial population growth. When advances in medical care (e.g., more aggressive treatment of cancer or organ transplantation) are introduced into a given area, the opportunistic mycoses increase in incidence.

TABLE 235-1 ENDEMIC AND OPPORTUNISTIC MYCOSES

Endemic Mycoses ^a	Opportunistic Mycoses
Coccidioidomycosis	Candidiasis
Histoplasmosis	Aspergillosis
Blastomycosis	Cryptococcosis
Phaeohiphomyces	Mucormycosis (zygomycosis)
Penicilliosis	Scedosporiosis
Sporotrichosis	Trichosporonosis
Paracoccidioidomycosis	Fusariosis
	Pneumocystosis

^aThe endemic mycoses can also occur as opportunistic infections.

The definitive diagnosis of any fungal infection requires histopathologic identification of the fungus invading tissue and accompanying evidence of an inflammatory response. The identification of an inflammatory response has been especially important with regard to *Aspergillus* infection. *Aspergillus* is ubiquitous and can float in the air onto biopsy material. Therefore, in rare but important instances, this fungus is an ex vivo contaminant during processing of a specimen for microscopy, with a consequent incorrect diagnosis. The stains most commonly used to identify fungi are periodic acid–Schiff and Gomori methenamine silver. *Candida*, unlike other fungi, is visible on gram-stained tissue smears. Hematoxylin and eosin stain is not sufficient to identify *Candida* in tissue specimens. When positive, an india ink preparation of cerebrospinal fluid (CSF) is diagnostic for cryptococcosis. Most laboratories now use calcofluor white staining coupled with fluorescent microscopy to identify fungi in fluid specimens.

Extensive investigations of the diagnosis of deep organ fungal infections have yielded a variety of tests with different degrees of specificity and sensitivity. The most reliable tests are the detection of antibody to *Coccidioides immitis* in serum and CSF; of *Histoplasma capsulatum* antigen in urine, serum, and CSF; and of cryptococcal polysaccharide antigen in serum and CSF. These tests have a general sensitivity and specificity of 90%; however, because of variability among laboratories, testing on multiple occasions is advisable. The test for galactomannan has been used extensively in Europe and is now approved in the United States for diagnosis of aspergillosis. Sources of concern regarding galactomannan are the incidence of false-negative results and the need for multiple serial tests to reduce this incidence. The β -glucan test for *Candida* is also under evaluation but, like the galactomannan test, still requires additional validation; this test has a negative predictive value of ~90%. Both of these tests are being used with increasing frequency, especially for guiding the timing of initiation and duration of therapy. The galactomannan test is being evaluated in both serum and bronchoalveolar lavage fluid. Numerous polymerase chain reaction assays to detect antigens are in the developmental stages, as are nucleic acid hybridization techniques; currently, these tests are not widely available.

Of the fungal organisms, *Candida* is by far the most frequently recovered from blood. Although *Candida* species can be detected with any of the automated blood culture systems widely used at present, the lysis-centrifugation technique increases the sensitivity of blood cultures for *Candida* and for less common organisms (e.g., *H. capsulatum*). Lysis-centrifugation should be used when disseminated fungal infection is suspected.

Except in the cases of coccidioidomycosis, cryptococcosis, and histoplasmosis, there are no fully validated and widely used tests for serodiagnosis of disseminated fungal infection. Skin tests for the endemic mycoses are no longer available.

TREATMENT FUNGAL INFECTIONS

This discussion is intended as a brief overview of general strategies for the use of antifungal agents in the treatment of fungal infections. Regimens, schedules, and strategies are detailed in the chapters on specific mycoses that follow in this section. The doses cited here are standard doses for adults with invasive infection.

Since fungal organisms are eukaryotic cells that contain most of the same organelles (with many of the same physiologic functions) as human cells, the identification of drugs that selectively kill or inhibit fungi but are not toxic to human cells has been highly problematic. Far fewer antifungal than antibacterial agents have been introduced into clinical medicine.

AMPHOTERICIN B

The introduction of amphotericin B (AmB) in the late 1950s revolutionized the treatment of fungal infections in deep organs. Before AmB became available, cryptococcal meningitis and other disseminated fungal infections were nearly always fatal. For nearly a decade after AmB was introduced, it was the only effective agent

for the treatment of life-threatening fungal infections. AmB remains the broadest-spectrum antifungal agent but carries several disadvantages, including significant nephrotoxicity, lack of an oral preparation, and unpleasant side effects (fever, chills, and nausea) during treatment. To circumvent nephrotoxicity and infusion side effects, lipid formulations of AmB were developed and have virtually replaced the original colloidal deoxycholate formulation in clinical use (although the older formulation is still available). The lipid formulations include liposomal AmB (L-AmB; 3–5 mg/kg per day) and AmB lipid complex (ABLC; 5 mg/kg per day). A third preparation, AmB colloidal dispersion (ABCD; 3–4 mg/kg per day), is rarely used because of the high incidence of side effects associated with infusion.

The lipid formulations of AmB have the disadvantage of being considerably more expensive than the deoxycholate formulation. Experience is still accumulating on the comparative efficacy, toxicity, and advantages of the different formulations for specific clinical fungal infections, including central nervous system (CNS) infection. Whether there is a clinically significant difference in these drugs with respect to CNS penetration or nephrotoxicity remains controversial. Despite these issues and despite the expense, the lipid formulations are now much more commonly used than AmB deoxycholate in developed countries. In developing countries, AmB deoxycholate is still preferred because of the expense of the lipid formulations.

AZOLES

This class of antifungal drugs offers important advantages over AmB: the azoles cause little or no nephrotoxicity and are available in oral formulations. Early azoles included ketoconazole and miconazole, which have been replaced by newer agents for the treatment of deep organ fungal infections. The azoles' mechanism of action is inhibition of ergosterol synthesis in the fungal cell wall. Unlike AmB, these drugs are considered fungistatic, not fungicidal.

Fluconazole Since its introduction, fluconazole has played an extremely important role in the treatment of a wide variety of serious fungal infections. Its major advantages are the availability of both oral and IV formulations, a long half-life, satisfactory penetration of most body fluids (including ocular fluid and CSF), and minimal toxicity (especially relative to that of AmB). Its disadvantages include (usually reversible) hepatotoxicity and—at high doses—alopecia, muscle weakness, and dry mouth with a metallic taste. Fluconazole is not effective for the treatment of aspergillosis, mucormycosis, or *Scedosporium apiospermum* infections. It is less effective than the newer azoles against *Candida glabrata* and *Candida krusei*.

Fluconazole has become the agent of choice for the treatment of coccidioidal meningitis, although relapses have followed therapy with this drug. In addition, fluconazole is useful as both consolidation and maintenance therapy for cryptococcal meningitis. This agent has been shown to be as efficacious as AmB in the treatment of candidemia. The effectiveness of fluconazole in candidemia and the drug's relatively minimal toxicity, in conjunction with the inadequacy of diagnostic tests for widespread hematogenously disseminated candidiasis, have led to a change in the paradigm for candidemia management. The standard of care is now to treat all candidemic patients with an antifungal agent and to change all their intravascular lines, if feasible, rather than merely removing a singular suspect intravascular line and then observing the patient. The usual fluconazole regimen for treatment of candidemia is 400 mg/d given until 2 weeks after the last positive blood culture.

Fluconazole is considered effective as fungal prophylaxis in bone marrow transplant recipients and high-risk liver transplant patients. Its general use for prophylaxis in patients with leukemia, in AIDS patients with low CD4+ T cell counts, and in patients on surgical intensive care units remains controversial. Because of concerns about the possibility of infection due to resistant *Candida* species and of infection with *Aspergillus* species, many clinicians are initiating therapy with an echinocandin, which is then replaced by fluconazole once a susceptible *Candida* species is recovered and concern about *Aspergillus* is diminished.

Voriconazole Voriconazole, which is available in both oral and IV formulations, has a broader spectrum than fluconazole against *Candida* species (including *C. glabrata* and *C. krusei*) and is active against *Aspergillus*, *Scedosporium*, and *Fusarium*. It is generally considered the first-line drug of choice for treatment of aspergillosis. A few case reports have shown voriconazole to be effective in individual patients with coccidioidomycosis, blastomycosis, and histoplasmosis, but because of limited data this agent is not recommended for primary treatment of the endemic mycoses. Among the disadvantages of voriconazole (compared with fluconazole) are its more numerous interactions with many of the drugs used in patients predisposed to fungal infections. Hepatotoxicity, skin rashes (including photosensitivity), and visual disturbances are relatively common. Skin cancer surveillance is now recommended for patients taking voriconazole. In addition, voriconazole is considerably more expensive than fluconazole. Moreover, it is advisable to monitor voriconazole levels in certain patients since (1) this drug is completely metabolized in the liver by CYP2C9, CYP3A4, and CYP2C19; and (2) human genetic variability in CYP2C19 activity exists. Dosages should be reduced accordingly in patients with liver failure. Dose adjustments for renal insufficiency are not necessary; however, because the IV formulation is prepared in cyclodextrin, it should not be given to patients with severe renal insufficiency.

Itraconazole Itraconazole is available in IV and oral (capsule and suspension) formulations. Varying blood levels among patients taking oral itraconazole reflect a disadvantage compared with the other azoles. Itraconazole is the drug of choice for mild to moderate histoplasmosis and blastomycosis and has often been used for chronic mucocutaneous candidiasis. It has been approved by the U.S. Food and Drug Administration (FDA) for use in febrile neutropenic patients. Itraconazole has also proved useful for the treatment of chronic coccidioidomycosis, sporotrichosis, and *S. apiospermum* infection. The mucocutaneous and cutaneous fungal infections that have been treated successfully with itraconazole include oropharyngeal candidiasis (especially in AIDS patients), tinea versicolor, tinea capitis, and onychomycosis. Disadvantages of itraconazole include its poor penetration into CSF, the use of cyclodextrin in both the oral suspension and the IV formulation, the variable absorption of the drug in capsule form, and the need for monitoring of blood levels in patients taking capsules for disseminated mycoses. Reported cases of severe congestive heart failure in patients taking itraconazole have been a source of concern. Like the other azoles, itraconazole can cause hepatic toxicity.

Posaconazole Posaconazole is approved by the FDA for prophylaxis of aspergillosis and candidiasis in patients at high risk for developing these infections because of severe immunocompromise. It has also been approved for the treatment of oropharyngeal candidiasis and has been evaluated as therapy for zygomycosis, fusariosis, aspergillosis, cryptococcosis, and various other forms of candidal infection. The relevant studies of posaconazole in zygomycosis, fusariosis, and aspergillosis have examined salvage therapy. A study of more than 90 patients whose zygomycosis was refractory to other therapy yielded encouraging results. No trials of posaconazole for the treatment of candidemia have yet been reported. Case reports have described the drug's efficacy in coccidioidomycosis and histoplasmosis. Controlled trials have shown its effectiveness as a prophylactic agent in patients with acute leukemia and in bone marrow transplant recipients. In addition, posaconazole has been found to be effective against fluconazole-resistant *Candida* species. The results of a large-scale study of the use of posaconazole as salvage therapy for aspergillosis indicated that it is an alternative to other agents for salvage therapy; however, that study predated the use of voriconazole and the echinocandins.

ECHINOCANDINS

The echinocandins, including the FDA-approved drugs caspofungin, anidulafungin, and micafungin, have added considerably to the antifungal armamentarium. All three of these agents inhibit

β -1,3-glucan synthase, which is necessary for cell wall synthesis in fungi and is not a component of human cells. None of these agents is currently available in an oral formulation. The echinocandins are considered fungicidal for *Candida* and fungistatic for *Aspergillus*. Their greatest use to date is against candidal infections. They offer two advantages: broad-spectrum activity against all *Candida* species and relatively low toxicity. The minimal inhibitory concentrations (MICs) of all the echinocandins are highest against *Candida parapsilosis*; it is not clear whether these higher MIC values represent less clinical effectiveness against this species. The echinocandins are among the safest antifungal agents.

In controlled trials, *caspofungin* has been at least as efficacious as AmB for the treatment of candidemia and invasive candidiasis and as efficacious as fluconazole for the treatment of candidal esophagitis. In addition, caspofungin has been efficacious as salvage therapy for aspergillosis. *Anidulafungin* has been approved by the FDA as therapy for candidemia in nonneutropenic patients and for *Candida* esophagitis, intraabdominal infection, and peritonitis. In controlled trials, anidulafungin has been shown to be noninferior and possibly superior to fluconazole against candidemia and invasive candidiasis. It is as efficacious as fluconazole against candidal esophagitis. When anidulafungin is used with cyclosporine, tacrolimus, or voriconazole, no dosage adjustment is required for either drug in the combination. *Micafungin* has been approved for the treatment of esophageal candidiasis and candidemia and for prophylaxis in patients receiving stem cell transplants. In a head-to-head trial, micafungin was noninferior to caspofungin for the treatment of candidemia. Studies thus far have shown that coadministration of micafungin and cyclosporine does not require dose adjustments for either drug. When micafungin is given with sirolimus, the area under the plasma drug concentration–time curve rises for sirolimus, usually necessitating a reduction in its dose. In open-label trials, favorable results have been obtained with micafungin for the treatment of deep-seated *Aspergillus* and *Candida* infections.

FLUCYTOSINE (5-FLUOROCYTOSINE)

The use of flucytosine has diminished as newer antifungal drugs have been developed. This agent is now used most commonly in combination with AmB (deoxycholate or lipid formulations) for the initial treatment of cryptococcal meningitis. Flucytosine has a unique mechanism of action based on intrafungal conversion to 5-fluorouracil, which is toxic to the fungal cell. Development of resistance to the compound has limited its use as a single agent. Flucytosine is nearly always used in combination with AmB. Its good penetration into the CSF makes it attractive for use with AmB for treatment of cryptococcal meningitis. Flucytosine has also been recommended for the treatment of candidal meningitis in combination with AmB; comparative trials with AmB alone have not been done. Significant and frequent bone marrow depression is seen with flucytosine when this drug is used with AmB.

GRISEOFULVIN AND TERBINAFINE

Historically, griseofulvin has been useful primarily for ringworm infection. This agent is usually given for relatively long periods. Terbinafine has been used primarily for onychomycosis but also for ringworm. In comparative studies, terbinafine has been as effective as itraconazole and more effective than griseofulvin for both conditions.

TOPICAL ANTIFUNGAL AGENTS

A detailed discussion of the agents used for the treatment of cutaneous fungal infections and onychomycosis is beyond the scope of this chapter; the reader is referred to the dermatology literature. Many classes of compounds have been used to treat the common fungal infections of the skin. Among the azoles used are clotrimazole, econazole, miconazole, oxiconazole, sulconazole, ketoconazole, tioconazole, butoconazole, and terconazole. In general, topical treatment of vaginal candidiasis has been successful. Since little difference is thought to exist in the efficacy of the various vaginal preparations, the choice of agent is made by the physician and/or

1332 the patient on the basis of preference and availability. Fluconazole given orally at 150 mg has the advantage of not requiring repeated intravaginal application. Nystatin is a polyene that has been used for both oropharyngeal thrush and vaginal candidiasis. Useful agents in other classes include ciclopirox olamine, haloprogin, terbinafine, naftifine, tolnaftate, and undecylenic acid.

236 Histoplasmosis

Chadi A. Hage, L. Joseph Wheat

ETIOLOGY



Histoplasma capsulatum, a thermal dimorphic fungus, is the etiologic agent of histoplasmosis. In most endemic areas, *H. capsulatum* var. *capsulatum* is the causative agent. In Africa, *H. capsulatum* var. *duboisii* also is found; var. *duboisii* can be differentiated from var. *capsulatum* as the *duboisii* yeasts are larger. In Central and South America, histoplasmosis is common and is caused by clades of *H. capsulatum* var. *capsulatum* that differ genetically from those involved elsewhere.

Mycelia—the naturally infectious form of *Histoplasma*—have a characteristic appearance, with microconidial and macroconidial forms. Microconidia are oval and are small enough (2–4 μm) to reach the terminal bronchioles and alveoli. Shortly after infecting the host, mycelia transform into the yeasts that are found inside macrophages and other phagocytes. The yeast forms are characteristically small (2–5 μm), with occasional narrow budding. In the laboratory, mycelia are best grown at room temperature, whereas yeasts are grown at 37°C on enriched media.

EPIDEMIOLOGY



Histoplasmosis is the most prevalent endemic mycosis in North America. Although this fungal disease has been reported throughout the world, its endemicity is particularly notable in certain parts of North, Central, and South America; Africa; and Asia. In Europe, histoplasmosis is diagnosed fairly often, mostly in emigrants from or travelers to endemic areas on other continents. In the United States, the endemic areas spread over the Ohio and Mississippi river valleys. This pattern is related to the humid and acidic nature of the soil in these areas. Soil enriched with bird or bat droppings promotes the growth and sporulation of *Histoplasma*. Disruption of soil containing the organism leads to aerosolization of the microconidia and exposure of humans nearby. Activities associated with high-level exposure include spelunking, excavation, cleaning of chicken coops, demolition and remodeling of old buildings, and cutting of dead trees. Most cases seen outside of highly endemic areas represent imported disease—e.g., cases reported in Europe after travel to the Americas, Africa, or Asia.

PATHOGENESIS AND PATHOLOGY

Infection follows inhalation of microconidia (Fig. 236-1). Once they reach the alveolar spaces, microconidia are rapidly recognized and engulfed by alveolar macrophages. At this point, the microconidia transform into budding yeasts (Fig. 236-2), a process that is integral to the pathogenesis of histoplasmosis and is dependent on the availability of calcium and iron inside the phagocytes. The yeasts are capable of growing and multiplying inside resting macrophages. Neutrophils and then lymphocytes are attracted to the site of infection. Before the development of cellular immunity, yeasts use the phagosomes as a vehicle for translocation to local draining lymph nodes, whence they spread hematogenously throughout the reticuloendothelial system. Adequate cellular immunity develops ~2 weeks after infection. T cells produce interferon γ to assist the macrophages in killing the organism

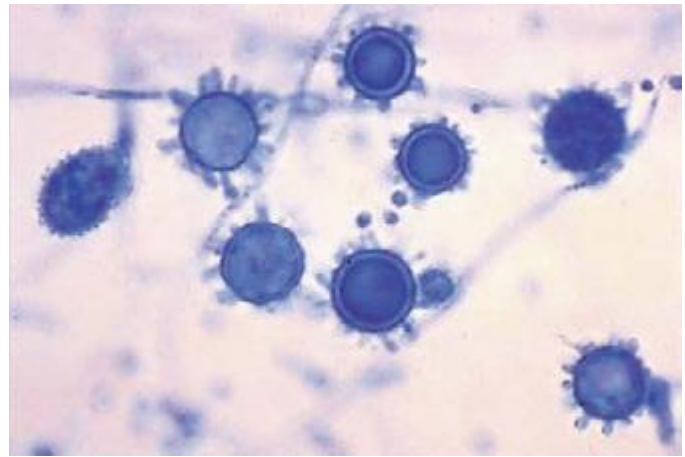


FIGURE 236-1 Spiked spherical conidia of *H. capsulatum* (lactophenol cotton blue stain).

and controlling the progression of disease. Interleukin 12 and tumor necrosis factor α (TNF- α) play an essential role in cellular immunity to *H. capsulatum*. In the immunocompetent host, macrophages, lymphocytes, and epithelial cells eventually organize and form granulomas that contain the organisms. These granulomas typically fibrose and calcify; calcified mediastinal lymph nodes and hepatosplenic calcifications are frequently found in healthy individuals from endemic areas. In immunocompetent hosts, infection with *H. capsulatum* confers some immunity to reinfection. In patients with impaired cellular immunity, the infection is not contained and can disseminate. Progressive disseminated histoplasmosis (PDH) can involve multiple organs, most commonly the bone marrow, spleen, liver (Fig. 236-3), adrenal glands, and mucocutaneous membranes. Unlike latent tuberculosis, latent histoplasmosis rarely reactivates.

Structural lung disease (e.g., emphysema) impairs the clearance of pulmonary histoplasmosis, and chronic pulmonary disease can result. This chronic process is characterized by progressive inflammation, tissue necrosis, and fibrosis mimicking cavitary tuberculosis.

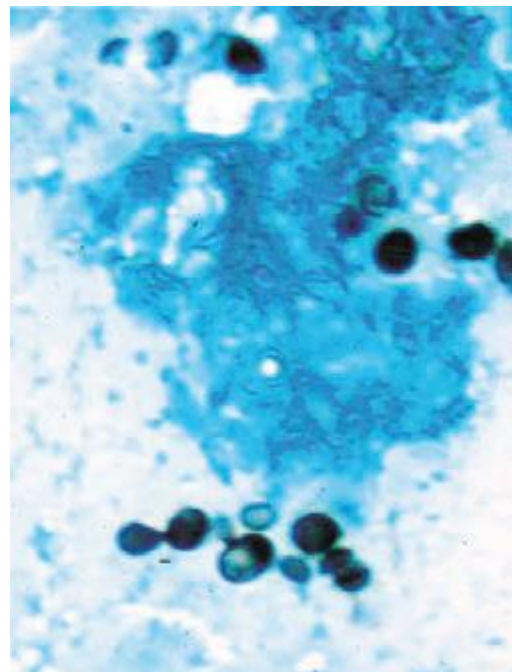


FIGURE 236-2 Small (2–5 μm) narrow budding yeasts of *H. capsulatum* from bronchoalveolar lavage fluid (Grocott's methenamine silver stain).

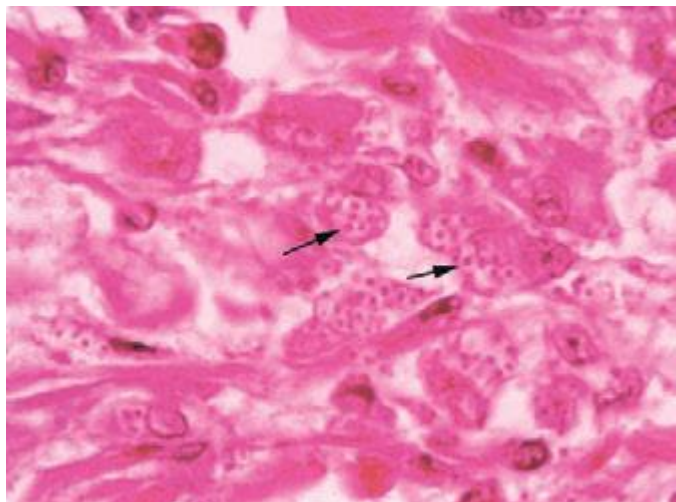


FIGURE 236-3 Intracellular yeasts (arrows) of *H. capsulatum* in a liver biopsy specimen (hematoxylin and eosin stain).

CLINICAL MANIFESTATIONS

The clinical spectrum of histoplasmosis ranges from asymptomatic infection to life-threatening illness. The attack rate and the extent and severity of the disease depend on the intensity of exposure, the immune status of the exposed individual, and the underlying lung architecture of the host.

In immunocompetent individuals with low-level exposure, most *Histoplasma* infections are either asymptomatic or mild and self-limited. Of adults residing in endemic areas, 50–80% have skin-test and/or radiographic evidence of previous infection without clinical manifestations. When symptoms do develop, they usually appear 1–4 weeks after exposure. Heavy exposure leads to a flulike illness with fever, chills, sweats, headache, myalgia, anorexia, cough, dyspnea, and chest pain. Chest radiographs usually show signs of pneumonitis with prominent hilar or mediastinal adenopathy. Pulmonary infiltrates may be focal with light exposure or diffuse with heavy exposure. Rheumatologic symptoms of arthralgia or arthritis, often associated with erythema nodosum, occur in 5–10% of patients with acute histoplasmosis. Pericarditis may also develop. These manifestations represent inflammatory responses to the acute infection rather than its direct effects. Hilar or mediastinal lymph nodes may undergo necrosis and coalesce to form large mediastinal masses that can cause compression of great vessels, proximal airways, and the esophagus. These necrotic lymph nodes may also rupture and create fistulas between mediastinal structures (e.g., bronchoesophageal fistulas).

PDH is typically seen in immunocompromised individuals, who account for ~70% of cases. Common risk factors include AIDS (CD4+ T cell count, <200/μL), extremes of age, immunosuppressive medications administered for prevention or treatment of rejection following transplantation (e.g., prednisone, mycophenolate, calcineurin inhibitors, and biologic response modifiers), and methotrexate, anti-TNF-α agents, or other biologic response modifiers given for inflammatory arthritis or Crohn's disease.

The spectrum of PDH ranges from an acute, rapidly fatal course—with diffuse interstitial or reticulonodular lung infiltrates causing respiratory failure, shock, coagulopathy, and multiorgan failure—to a more subacute course with a focal organ distribution. Common manifestations include fever and weight loss. Hepatosplenomegaly also is common. Other findings may include meningitis or focal brain lesions, ulcerations of the oral mucosa, gastrointestinal ulcerations, and adrenal insufficiency. Prompt recognition of this devastating illness is of paramount importance in patients with more severe manifestations or with underlying immunosuppression, especially AIDS (**Chap. 226**).

Chronic cavitary histoplasmosis is seen in smokers who have structural lung disease (e.g., bullous emphysema). This chronic illness is characterized by productive cough, dyspnea, low-grade fever, night sweats, and weight loss. Chest radiographs usually show upper-lobe infiltrates,

cavitation, and pleural thickening—findings resembling those of tuberculosis. Without treatment, the course is slowly progressive.

Fibrosing mediastinitis is an uncommon and serious complication of histoplasmosis. In certain patients, acute infection is followed for unknown reasons by progressive fibrosis around the hilar and mediastinal lymph nodes. Involvement may be unilateral or bilateral; bilateral involvement carries a worse prognosis. Major manifestations include superior vena cava syndrome, obstruction of pulmonary vessels, and airway obstruction. Patients may experience recurrent pneumonia, hemoptysis, or respiratory failure. Fibrosing mediastinitis is fatal in up to one-third of cases.

In healed histoplasmosis, calcified mediastinal nodes or lung parenchyma may erode through the walls of the airways and cause hemoptysis. This condition is called *broncholithiasis*.



The clinical features and management of histoplasmosis caused by the genetically different clades in Central and South America are similar to those of the disease in North America. African histoplasmosis caused by var. *duboisii* is clinically distinct and is characterized by frequent skin and bone involvement.

DIAGNOSIS

Fungal culture remains the gold standard diagnostic test for histoplasmosis. However, culture results may not be known for up to 1 month, and cultures are often negative in less severe cases. Cultures are positive in ~75% of cases of PDH and chronic pulmonary histoplasmosis. Cultures of bronchoalveolar lavage (BAL) fluid are positive in about half of cases that include acute pulmonary histoplasmosis causing diffuse infiltrates with hypoxemia. In PDH, the culture yield is highest for BAL fluid, bone marrow aspirate, and blood. Cultures of sputum or bronchial washings are usually positive in chronic pulmonary histoplasmosis. Cultures are typically negative, however, in other forms of histoplasmosis.

Fungal stains of cytopathology or biopsy materials showing structures resembling *Histoplasma* yeasts are helpful in the diagnosis of PDH, yielding positive results in about half of cases. Yeasts can be seen in BAL fluid (Fig. 236-2) from patients with diffuse pulmonary infiltrates, in bone marrow biopsy samples, and in biopsy specimens of other involved organs (e.g., the adrenal glands). Occasionally, yeasts are seen in blood smears from patients with severe PDH. However, artifacts and other fungal elements sometimes stain positive and may be misidentified as *Histoplasma* yeasts.

The detection of *Histoplasma* antigen in body fluids is extremely useful in the diagnosis of PDH and acute diffuse pulmonary histoplasmosis. The sensitivity of this technique is >95% in patients with PDH and >80% in patients with acute pulmonary histoplasmosis if both urine and serum are tested. Antigen level correlates with severity of illness in PDH and can be used to follow disease progression as levels predictably decrease with effective therapy. An increase in antigen levels also predicts relapse. Antigen can be detected in cerebrospinal fluid from patients with meningitis and in BAL fluid from those with pneumonia. Cross-reactivity occurs with African histoplasmosis, blastomycosis, coccidioidomycosis, paracoccidioidomycosis, and *Penicillium marneffei* infection.

Serologic tests, including immunodiffusion and complement fixation, are useful for the diagnosis of histoplasmosis in immunocompetent patients. Serum antibody titers may rise fourfold in patients with acute histoplasmosis. Serologic tests are especially useful for the diagnosis of chronic pulmonary histoplasmosis. The limitations of serology, however, include insensitivity early in the course of infection (at least 1 month is required for the production of antibodies), insensitivity in immunosuppressed patients, and persistence of detectable antibody for several years after infection. Positive results from past infection may lead to a misdiagnosis of active histoplasmosis in a patient with another disease process.

TREATMENT HISTOPLASMOSIS

Treatment recommendations for histoplasmosis are summarized in **Table 236-1**. Treatment is indicated for all patients with PDH or chronic pulmonary histoplasmosis as well as for symptomatic

TABLE 236-1 RECOMMENDATIONS FOR THE TREATMENT OF HISTOPLASMOVIS

Type of Histoplasmosis	Treatment Recommendations	Comments
Acute pulmonary, moderate to severe illness with diffuse infiltrates and/or hypoxemia	Lipid AmB (3–5 mg/kg per day) ± glucocorticoids for 1–2 weeks; then itraconazole (200 mg bid) for 12 weeks. Monitor renal and hepatic function.	Patients with mild cases usually recover without therapy, but itraconazole should be considered if the patient's condition has not improved after 1 month.
Chronic/cavitary pulmonary	Itraconazole (200 mg qd or bid) for at least 12 months. Monitor hepatic function.	Continue treatment until radiographic findings show no further improvement. Monitor for relapse after treatment is stopped.
Progressive disseminated	Lipid AmB (3–5 mg/kg per day) for 1–2 weeks; then itraconazole (200 mg bid) for at least 12 months. Monitor renal and hepatic function.	Liposomal AmB is preferred, but the AmB lipid complex may be used because of cost. Chronic maintenance therapy may be necessary if the degree of immunosuppression cannot be reduced.
Central nervous system	Liposomal AmB (5 mg/kg per day) for 4–6 weeks; then itraconazole (200 mg bid or tid) for at least 12 months. Monitor renal and hepatic function.	A longer course of lipid AmB is recommended because of the high risk of relapse. Itraconazole should be continued until cerebrospinal fluid or CT abnormalities clear.

Abbreviation: AmB, amphotericin B.

patients with acute pulmonary histoplasmosis causing diffuse infiltrates, especially with hypoxemia. In most cases of pulmonary histoplasmosis, treatment is not recommended because the degree of exposure is not heavy; the infection is asymptomatic or symptoms are mild, subacute, and not progressive; and the illness resolves without therapy.

The preferred treatments for histoplasmosis include the lipid formulations of amphotericin B (AmB) in more severe cases and itraconazole in others. Liposomal AmB has been more effective than the deoxycholate formulation for treatment of PDH in patients with AIDS. The deoxycholate formulation is an alternative to lipid formulations for patients at low risk for nephrotoxicity. Posaconazole, voriconazole, and fluconazole are alternatives for patients who cannot take itraconazole.

In severe cases requiring hospitalization, a lipid formulation of AmB is followed by itraconazole. In patients with meningitis, a lipid formulation of AmB should be given for 4–6 weeks before the switch to itraconazole. In immunosuppressed patients, the degree of immunosuppression should be reduced if possible, although immune reconstitution inflammatory syndrome (IRIS) may ensue. Antiretroviral treatment improves the outcome of PDH in patients with AIDS and is recommended; however, whether antiretroviral treatment should be delayed to avoid IRIS is unknown.

Blood levels of itraconazole should be monitored to ensure adequate drug exposure, with target concentrations of the parent drug and its hydroxy metabolites of 1–5 µg/mL as measured by high-performance liquid chromatography and 2–10 µg/mL as measured by microbiologic assay. Drug interactions should be carefully assessed: itraconazole not only is cleared by cytochrome P450 metabolism but also inhibits cytochrome P450. This profile causes interactions with many other medications.

The duration of treatment for acute pulmonary histoplasmosis is 6–12 weeks, while that for PDH and chronic pulmonary histoplasmosis is ≥1 year. Antigen levels in urine and serum should be monitored during and for at least 1 year after therapy for PDH. Stable or rising antigen levels suggest treatment failure or relapse.

Previously, lifelong itraconazole maintenance therapy was recommended for patients with AIDS once histoplasmosis was diagnosed. Today, however, maintenance therapy is not required for patients who respond well to antiretroviral therapy, with CD4+ T cell counts of at least 150/µL (preferably >250/µL); who complete at least 1 year of itraconazole therapy; and who exhibit neither clinical evidence of active histoplasmosis nor an antigenuria level of >4 ng/mL. Maintenance therapy also appears to be unnecessary in patients receiving immunosuppressive treatment if the degree of immunosuppression can be reduced through an approach similar to that used for patients with AIDS.

Fibrosing mediastinitis, which represents a chronic fibrotic reaction to past mediastinal histoplasmosis rather than an active infection, does not respond to antifungal therapy. While treatment is often prescribed for patients with pulmonary histoplasmosis who have not recovered within 1 month and for those with persistent mediastinal lymphadenopathy, the effectiveness of antifungal therapy in these situations is unknown.


237 Coccidioidomycosis

Neil M. Ampel

DEFINITION AND ETIOLOGY

Coccidioidomycosis, commonly known as Valley fever (see “Epidemiology,” below), is caused by dimorphic soil-dwelling fungi of the genus *Coccidioides*. Genetic analysis has demonstrated the existence of two species, *C. immitis* and *C. posadasii*. These species are indistinguishable with regard to the clinical disease they cause and their appearance on routine laboratory media. Thus, the organisms will be referred to simply as *Coccidioides* for the remainder of this chapter.

EPIDEMIOLOGY

 Coccidioidomycosis is confined to the Western Hemisphere between the latitudes of 40°N and 40°S. In the United States, areas of high endemicity include the southern portion of the San Joaquin Valley of California and the south-central region of Arizona. However, infection may be acquired in other areas of the southwestern United States, including the southern coastal counties in California, southern Nevada, southwestern Utah, southern New Mexico, and western Texas, including the Rio Grande Valley. Outside the United States, coccidioidomycosis is endemic to northern Mexico as well as to localized regions of Central America. In South America, there are endemic foci in Colombia, Venezuela, northeastern Brazil, Paraguay, Bolivia, and north-central Argentina.

The risk of infection is increased by direct exposure to soil harboring *Coccidioides*. Because of difficulty in isolating *Coccidioides* from the soil, the precise characteristics of potentially infectious soil are not known. In the United States, several outbreaks of coccidioidomycosis have been associated with soil from archaeological excavations of Amerindian sites both within and outside of the recognized endemic region. These cases often involved alluvial soils in regions of relative aridity with moderate temperature ranges. *Coccidioides* was isolated at depths of 2–20 cm below the surface. The recent identification of three cases of coccidioidomycosis in eastern Washington State may suggest that the endemic region is expanding.

In endemic areas, many cases of *Coccidioides* infection occur without obvious soil or dust exposure. Climatic factors appear to increase the infection rate in these regions. In particular, periods of aridity following rainy seasons have been associated with marked increases in the number of symptomatic cases. Overall, the incidence within the United States has increased substantially over the past decade, with nearly 43 cases per 100,000 residents of the endemic region in

2011. Most of that increase has occurred in south-central Arizona, where most of that state's population resides, and in the southern San Joaquin Valley of California, a much less populated region. The factors causing this increase have not been fully elucidated; however, an influx of older individuals without prior coccidioidal infection appears to be involved. Other variables, such as climate change, construction activity, and increased awareness and reporting, may also be factors. Health care providers should consider coccidioidomycosis when evaluating persons with pneumonia who live in or have traveled to endemic areas.

PATHOGENESIS, PATHOLOGY, AND IMMUNE RESPONSE

On agar media and in the soil, *Coccidioides* organisms exist as filamentous molds. Within this mycelial structure, individual filaments (*hyphae*) elongate and branch, some growing upward. Alternating cells within the hyphae degenerate, leaving barrel-shaped viable elements called *arthroconidia*. Measuring ~2 by 5 μm , arthroconidia may become airborne for extended periods. Their small size allows them to evade initial mechanical mucosal defenses and reach deep into the bronchial tree, where infection is initiated in the nonimmune host.

Once in a susceptible host, the arthroconidia enlarge, become rounded, and develop internal septations. The resulting structures, called *spherules* (Fig. 237-1), may attain sizes of 200 μm and are unique to *Coccidioides*. The septations encompass uninuclear elements called *endospores*. Spherules may rupture and release packets of endospores that can themselves develop into spherules, thus propagating infection locally. If returned to artificial media or the soil, the fungus reverts to its mycelial stage.

Clinical observations and data from studies of animals strongly support the critical role of a robust cellular immune response in the host's control of coccidioidomycosis. Necrotizing granulomas containing

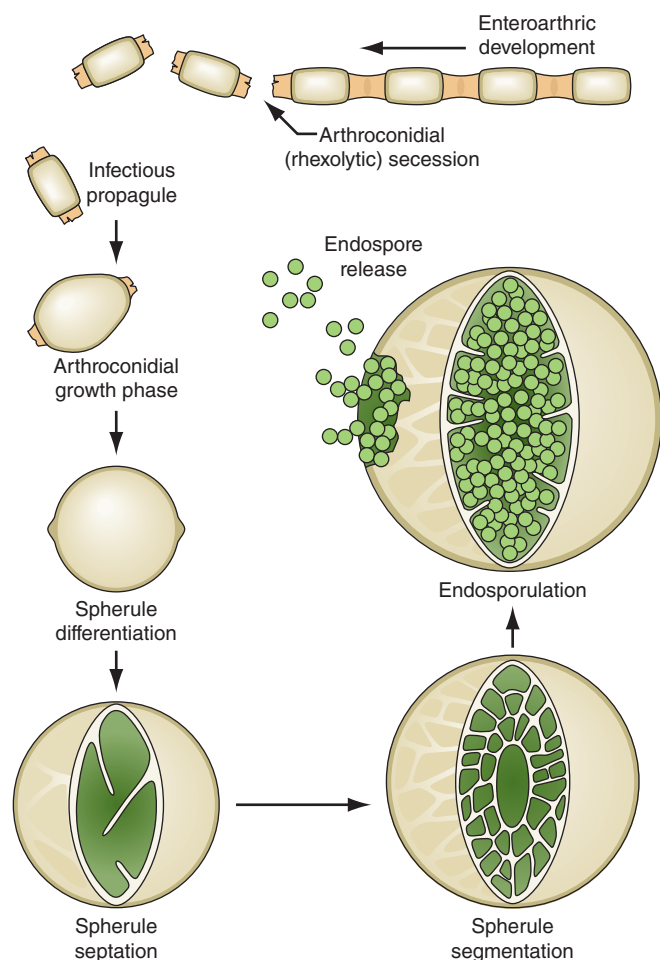


FIGURE 237-1 Life cycle of *Coccidioides*. (From TN Kirkland, J Fierer: *Emerg Infect Dis* 2:192, 1996.)

spherules are typically identified in patients with resolved pulmonary infection. In disseminated disease, granulomas are generally poorly formed or do not develop at all, and a polymorphonuclear leukocyte response occurs frequently. In patients who are asymptomatic or in whom the initial pulmonary infection resolves, delayed-type hypersensitivity to coccidioidal antigens has been routinely documented.

CLINICAL AND LABORATORY MANIFESTATIONS

Of infected individuals, 60% are completely asymptomatic, and the remaining 40% have symptoms that are related principally to pulmonary infection, including fever, cough, and pleuritic chest pain. The risk of symptomatic illness increases with age. Coccidioidomycosis is commonly misdiagnosed as community-acquired bacterial pneumonia.

There are several cutaneous manifestations of primary pulmonary coccidioidomycosis. Toxic erythema consisting of a maculopapular rash has been noted in some cases. Erythema nodosum (see Fig. 25e-40)—typically over the lower extremities—or erythema multiforme (see Fig. 25e-25)—usually in a necklace distribution—may occur; these manifestations are seen particularly often in women. Arthralgias and arthritis may develop. The diagnosis of primary pulmonary coccidioidomycosis is suggested by a history of night sweats or profound fatigue as well as by peripheral-blood eosinophilia and hilar or mediastinal lymphadenopathy on chest radiography. While pleuritic chest pain is common, pleural effusions occur in fewer than 10% of cases. Such effusions are invariably associated with a pulmonary infiltrate on the same side. The cellular content of these effusions is mononuclear in nature; *Coccidioides* is rarely grown from effusions.

In most patients, primary pulmonary coccidioidomycosis usually resolves without sequelae in weeks. However, several pneumonic complications may arise. Pulmonary nodules are residua of primary pneumonia. Generally single, frequently located in the upper lobes, and ≤ 4 cm in diameter, nodules are often discovered on a routine chest radiograph in an asymptomatic patient. Calcification is uncommon. Coccidioidal pulmonary nodules can be difficult to distinguish radiographically from pulmonary malignancies. Like malignancies, coccidioidal nodules often enhance on positron emission tomography. However, routine CT often demonstrates multiple nodules in coccidioidomycosis. Biopsy is often required to distinguish between these two conditions.

Pulmonary cavities occur when a nodule extrudes its contents into the bronchus, resulting in a thin-walled shell. These cavities can be associated with persistent cough, hemoptysis, and pleuritic chest pain. Rarely, a cavity may rupture into the pleural space, causing pyopneumothorax. In such cases, patients present with acute dyspnea, and the chest radiograph reveals a collapsed lung with a pleural air-fluid level. Chronic or persistent pulmonary coccidioidomycosis manifests with prolonged symptoms of fever, cough, and weight loss and is radiographically associated with pulmonary scarring, fibrosis, and cavities. It occurs most commonly in patients who already have chronic lung disease due to other etiologies.

In some cases, primary pneumonia presents as a diffuse reticulonodular pulmonary process (detected by plain chest radiography) in association with dyspnea and fever. Primary diffuse coccidioidal pneumonia may occur in settings of intense environmental exposure or profoundly suppressed cellular immunity (e.g., in patients with AIDS), with unrestrained fungal growth that is frequently associated with fungemia.

Clinical dissemination outside the thoracic cavity occurs in fewer than 1% of infected individuals. Dissemination is more likely to occur in male patients, particularly those of African-American or Filipino ancestry, and in persons with depressed cellular immunity, including patients with HIV infection and peripheral-blood CD4⁺ T cell counts of $<250/\mu\text{L}$; those receiving chronic glucocorticoid therapy; those with allogeneic solid-organ transplants; and those being treated with tumor necrosis factor α antagonists. Women who acquire infection during the second or third trimester of pregnancy also are at risk for disseminated disease. Common sites for dissemination include the skin, bone, joints, soft tissues, and meninges. Dissemination may follow symptomatic or asymptomatic pulmonary infection and may involve

1336 only one site or multiple anatomic foci. When it occurs, clinical dissemination is usually evident within the first few months after primary pulmonary infection.

Coccidioidal meningitis, if untreated, is uniformly fatal. Patients usually present with a persistent headache, which is sometimes accompanied by lethargy and confusion. Nuchal rigidity, if present, is not severe. Examination of cerebrospinal fluid (CSF) demonstrates lymphocytic pleocytosis with profound hypoglycorrhachia and elevated protein levels. CSF eosinophilia is occasionally documented. With or without appropriate therapy, patients may develop hydrocephalus, which presents clinically as a marked decline in mental status, often with gait disturbances.

DIAGNOSIS

As mentioned above, coccidioidomycosis is often misdiagnosed as community-acquired bacterial pneumonia. Clues that suggest a diagnosis of coccidioidomycosis include peripheral-blood eosinophilia, hilar or mediastinal adenopathy on radiographic imaging, marked fatigue, and failure to improve with antibiotic therapy.

Serology plays an important role in establishing a diagnosis of coccidioidomycosis. Several techniques are available, including the traditional tube-precipitin (TP) and complement-fixation (CF) assays, immunodiffusion (IDTP and IDCF), and enzyme immunoassay (EIA) to detect IgM and IgG antibodies. TP and IgM antibodies are found in serum soon after infection and persist for weeks. They are not useful for gauging disease progression and are not found in the CSF. The CF and IgG antibodies occur later in the course of the disease and persist longer than TP and IgM antibodies. Rising CF titers are associated with clinical progression, and the presence of CF antibody in CSF is indicative of coccidioidal meningitis. Antibodies disappear over time in persons whose clinical illness resolves.

Because of its commercial availability, the coccidioidal EIA is frequently used as a screening tool for coccidioidal serology. There has been concern that the IgM EIA is occasionally falsely positive, particularly in asymptomatic individuals. In addition, while the sensitivity and specificity of the IgG EIA appear to be higher than those of the CF and IDCF assays, the optical density obtained in the EIA does not correlate with the serologic titer of either of the latter tests.

Coccidioides grows within 3–7 days at 37°C on a variety of artificial media, including blood agar. Therefore, it is always useful to obtain samples of sputum or other respiratory fluids and tissues for culture in suspected cases of coccidioidomycosis. The clinical laboratory should be alerted to the possibility of this diagnosis, since *Coccidioides* poses a significant laboratory hazard if it is inadvertently inhaled. The organism can also be identified directly. While treatment of samples with potassium hydroxide is rarely fruitful in establishing the diagnosis, examination of sputum or other respiratory fluids after Papanicolaou or Gomori methenamine silver staining reveals spherules in a significant proportion of patients with pulmonary coccidioidomycosis. For fixed tissues (e.g., those obtained from biopsy specimens), spherules with surrounding inflammation can be demonstrated with hematoxylin-eosin or Gomori methenamine silver staining.

A commercially available test for coccidioidal antigenuria and antigenemia has been developed and appears to be particularly useful in immunosuppressed patients with severe or disseminated disease. False-positive results may occur in cases of histoplasmosis or blastomycosis. Some laboratories offer genomic detection by polymerase chain reaction.

TREATMENT COCCIDIOIDOMYCOSIS

Currently, two main classes of antifungal agents are useful for the treatment of coccidioidomycosis (Table 237-1). While once prescribed routinely, amphotericin B in all its formulations is now reserved for only the most severe cases of dissemination and for intrathecal or intraventricular administration to patients with coccidioidal meningitis in whom triazole antifungal therapy has failed. The original formulation of amphotericin B, which is dispersed with deoxycholate, is usually administered intravenously in doses

TABLE 237-1 CLINICAL PRESENTATIONS OF COCCIDIOIDOMYCOSIS, THEIR FREQUENCY, AND RECOMMENDED INITIAL THERAPY FOR THE IMMUNOCOMPETENT HOST

Clinical Presentation	Frequency, %	Recommended Therapy
Asymptomatic infection	60	None
Primary pneumonia (focal)	40	In most cases, none ^a
Diffuse pneumonia	<1	Amphotericin B followed by prolonged oral triazole therapy
Pulmonary sequelae	5	
Nodule	—	None
Cavity	—	In most cases, none ^b
Chronic pneumonia	—	Prolonged triazole therapy
Disseminated disease	≤1	
Skin, bone, joint, soft tissue disease	—	Prolonged triazole therapy ^c
Meningitis	—	Lifelong triazole therapy ^d

^aTreatment is indicated for hosts with depressed cellular immunity as well as for those with prolonged symptoms and signs of increased severity, including night sweats for >3 weeks, weight loss of >10%, a complement-fixation titer of >1:16, and extensive pulmonary involvement on chest radiography. ^bTreatment (usually with the oral triazoles fluconazole and itraconazole) is recommended for persistent symptoms. ^cIn severe cases, some clinicians would use amphotericin B as initial therapy. ^dIntraventricular or intrathecal amphotericin B is recommended in cases of triazole failure. Hydrocephalus may occur, requiring a CSF shunt.

Note: See text for dosages and durations.

of 0.7–1.0 mg/kg either daily or three times per week. The newer lipid-based formulations—amphotericin B lipid complex (ABLC), amphotericin B colloidal dispersion (ABCD), and amphotericin B liposomal complex (L-Amb)—are associated with less renal toxicity. The lipid dispersions are administered intravenously at doses of 5 mg/kg daily or three times per week.

Triazole antifungals are the principal drugs now used to treat most cases of coccidioidomycosis. Clinical trials have demonstrated the usefulness of both fluconazole and itraconazole. Evidence indicates that itraconazole may be more efficacious against bone and joint disease. Because of its demonstrated penetration into CSF, fluconazole is the azole of choice for the treatment of coccidioidal meningitis, but itraconazole also is effective. For both drugs, a minimal oral adult dosage of 400 mg/d should be used. The maximal dose of itraconazole is 200 mg three times daily, but higher doses of fluconazole may be given. Two newer triazole antifungals, posaconazole and voriconazole, are now available. Data suggest that both drugs may be useful against infections, including meningitis, in which prior fluconazole therapy has failed. High-dose triazole therapy may be teratogenic during the first trimester of pregnancy; thus, amphotericin B should be considered as therapy for coccidioidomycosis in pregnant women during this period.

Most patients with focal primary pulmonary coccidioidomycosis require no therapy. Patients for whom antifungal therapy should be considered include those with underlying cellular immunodeficiencies and those with prolonged symptoms and signs of extensive disease. Specific criteria include symptoms persisting for ≥2 months, night sweats occurring for >3 weeks, weight loss of >10%, a serum CF antibody titer of >1:16, and extensive pulmonary involvement apparent on chest radiography.

Diffuse pulmonary coccidioidomycosis represents a special situation. Because most patients with this form of disease are profoundly hypoxicemic and critically ill, many clinicians favor beginning therapy with amphotericin B and switching to an oral triazole antifungal once clinical improvement occurs.

The nodules that may follow primary pulmonary coccidioidomycosis do not require treatment. As noted above, these nodules are not easily distinguished from pulmonary malignancies by means of radiographic imaging. Close clinical follow-up and biopsy may be required to distinguish between these two entities. Most pulmonary cavities do not require therapy. Antifungal treatment should be considered in

patients with persistent cough, pleuritic chest pain, and hemoptysis. Occasionally, pulmonary coccidioidal cavities become secondarily infected. This development is usually manifested by an air-fluid level within the cavity. Bacterial flora or *Aspergillus* species are commonly involved, and therapy directed at these organisms should be considered. Surgery is rarely required except in cases of persistent hemoptysis or pyopneumothorax. For chronic pulmonary coccidioidomycosis, prolonged antifungal therapy—lasting for at least 1 year—is usually required, with monitoring of symptoms, radiographic changes, sputum cultures, and serologic titers.

Most cases of disseminated coccidioidomycosis require prolonged antifungal therapy. Duration of treatment is based on resolution of the signs and symptoms of the lesions in conjunction with a significant decline in serum CF antibody titer. Such therapy routinely is continued for at least several years. Relapse occurs in 15–30% of individuals once therapy is discontinued.

Coccidioidal meningitis poses a special challenge. While most patients with this form of disease respond to treatment with oral triazoles, 80% experience relapse when therapy is stopped. Thus, lifelong therapy is recommended. In cases of triazole failure, intrathecal or intraventricular amphotericin B may be used. Installation requires considerable expertise and should be undertaken only by an experienced health care provider. Shunting of CSF in addition to appropriate antifungal therapy is required in cases of meningitis complicated by hydrocephalus. It is prudent to obtain expert consultation in all cases of coccidioidal meningitis.

PREVENTION

There are no proven methods to reduce the risk of acquiring coccidioidomycosis among residents of an endemic region, but avoidance of direct contact with uncultivated soil or with visible dust containing soil is reasonable. For individuals with suppressed cellular immunity, the risk of developing symptomatic coccidioidomycosis is greater than that in the general population. Among those about to undergo allogeneic solid-organ transplantation, antifungal therapy is appropriate when there is evidence of active or recent coccidioidomycosis. Several cases of donor-transmitted coccidioidomycosis have occurred during transplantation. If possible, donors from an endemic region should be screened for coccidioidomycosis before transplantation. Data on the use of antifungal agents for prophylaxis in other situations are limited. The administration of an antifungal drug to prevent symptomatic coccidioidomycosis is not recommended for HIV-1-infected patients who live in an endemic region. Most experts would administer a triazole to patients with a history of active coccidioidomycosis or a positive coccidioidal serology in whom therapy with tumor necrosis factor α antagonists is being initiated.

238 Blastomycosis

Donna C. Sullivan, Rathel L. Nolan, III

Blastomycosis is a systemic pyogranulomatous infection, involving primarily the lungs, that follows inhalation of the conidia of *Blastomyces dermatitidis*. Pulmonary blastomycosis varies from an asymptomatic infection to acute or chronic pneumonia. Hematogenous dissemination to skin, bones, and the genitourinary system is common; however, almost any organ can be involved.

ETIOLOGIC AGENT

B. dermatitidis is the asexual state of *Ajellomyces dermatitidis*. Two serotypes have been identified on the basis of the presence or absence of the A antigen. Distinct genotypic groups have been differentiated by rDNA polymerase chain reaction restriction fragment length

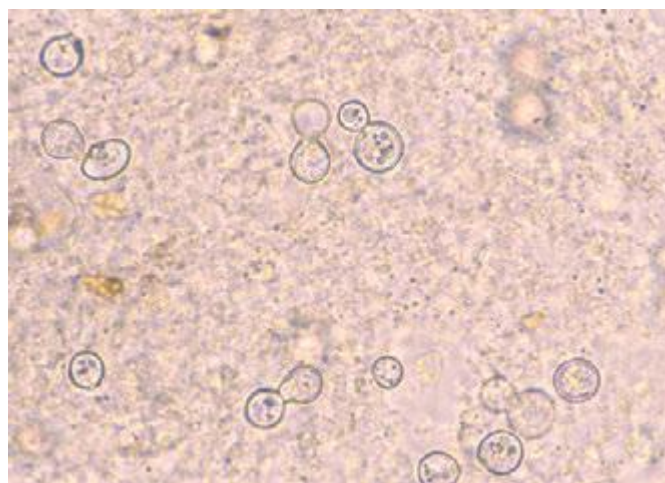


FIGURE 238-1 *Blastomyces dermatitidis* broad-based budding yeast in the aspirate of a chest wall abscess. Note the presence of multiple nuclei, the thickened cell wall, and the broad-based bud.

polymorphisms and microsatellite markers. *B. dermatitidis* exhibits thermal dimorphism, growing as the mycelial phase at room temperature and as the yeast phase at 37°C. Primary isolation in the laboratory is most dependable for the mycelial phase incubated at 30°C. Definitive identification usually requires conversion to the yeast phase at 37°C or—now more commonly—the use of nucleic acid amplification techniques that detect mycelial-phase growth. Under the microscope, the yeast cells are usually 8–15 μm in diameter, have thick refractile cell walls, are multinucleate, and exhibit a single, large, broad-based bud (Fig. 238-1).

EPIDEMIOLOGY

Most cases of blastomycosis have been reported in North America. Endemic areas include the southeastern and south-central states bordering the Mississippi and Ohio river basins, the midwestern states, and the Canadian provinces bordering the Great Lakes. A small endemic area exists in New York and Canada along the St. Lawrence River. Acute blastomycosis is typically found only in North America, and the clinical presentation of blastomycosis in nonendemic areas is as a chronic disease.



Outside North America, blastomycosis occurs sporadically in Nigeria, Zimbabwe, Tunisia, Saudi Arabia, Israel, Lebanon, and India. The disease has been reported most frequently in Africa.

Early studies indicated that middle-aged men with outdoor occupations were at greatest risk. Reported outbreaks, however, do not suggest a predilection according to sex, age, race, occupation, or season. The specific niche in nature in which the organism resides remains uncertain; *B. dermatitidis* probably grows as microfoci in the warm, moist soil of wooded areas rich in organic debris. Inhalation of conidia following exposure to soil, whether related to work or recreation, appears to be the common factor associated with infection. Outbreaks of human disease may be preceded by the occurrence of disease in simultaneously exposed dogs. Zoonotic transmission is rare but has been reported in association with dog bites, pet kinkajou bites, cat scratches, and animal necropsies.

PATHOGENESIS

Alveolar macrophages and polymorphonuclear leukocytes are critical for phagocytosis and killing of the inhaled conidia of *B. dermatitidis*. The interaction of these mediators of the innate immune response with local host factors, such as lung surfactant, plays a significant role in inhibiting conversion to the pathogenic yeast form. This inhibition prevents the establishment of symptomatic disease and may account for the high frequency of asymptomatic infections in outbreaks. Once conversion to the thick-walled yeast form has occurred, phagocytosis and killing are much more difficult, and the development of clinically

1338 apparent infection is much more likely. Ultimately, the T lymphocyte response—specifically, a T_H1 response—is the primary factor in limiting infection and dissemination. Moreover, yeast-phase conversion results in the expression of yeast phase-specific proteins such as the 120-kDa glycoprotein adhesin BAD-1 and the *Blastomyces* yeast phase-specific protein 1 (BYS1). BAD-1 has been well characterized as a virulence factor and is the major epitope for humoral and cellular immunity. The role of BYS1, putatively identified as a signal peptide, has not been determined.

APPROACH TO THE PATIENT: Blastomycosis

Blastomycosis most commonly presents as acute or chronic pneumonia that has been refractory to therapy with antibacterial drugs. Whether acute or chronic, blastomycosis may mimic many other disease processes. For example, acute pulmonary blastomycosis may present with signs and symptoms indistinguishable from those of bacterial pneumonia or influenza, and chronic pulmonary blastomycosis may mimic malignancy or tuberculosis. Skin lesions are often misdiagnosed as basal cell or squamous cell carcinoma, pyoderma gangrenosum, or keratoacanthoma. Laryngeal lesions are frequently mistaken for squamous cell carcinoma. Thus, the clinician must maintain a high index of suspicion and ensure that secretions or biopsy materials from patients who live in or have visited regions endemic for blastomycosis are subjected to careful histologic evaluation. This diligence is especially important in caring for individuals with pneumonia who fail to respond to treatment with antibacterial agents.

CLINICAL MANIFESTATIONS

Acute pulmonary infection is often diagnosed in association with point-source outbreaks. Typical symptoms include the abrupt onset of fever, chills, pleuritic chest pain, arthralgias, and myalgias. Cough is initially nonproductive but frequently becomes purulent as disease progresses. Chest radiographs usually reveal alveolar infiltrates with consolidation. Pleural effusions and hilar adenopathy are uncommon. Most patients diagnosed with pulmonary blastomycosis have chronic indolent pneumonia with signs and symptoms of fever, weight loss, productive cough, and hemoptysis. The most common radiologic findings are alveolar infiltrates with or without cavitation, mass lesions that mimic bronchogenic carcinoma, and fibronodular infiltrates. Hematogenous dissemination to the skin, bones, and genitourinary tract occurs most often in association with chronic pulmonary disease. Although blastomycosis is not considered an opportunistic infection, immunosuppression has been recognized as a risk factor for more serious pulmonary involvement, including respiratory failure (adult respiratory distress syndrome) associated with miliary disease or diffuse pulmonary infiltrates. In the late stages of AIDS, mortality rates of $\geq 50\%$ have been documented. Most deaths occur within the first few days of therapy. Solid-organ transplant recipients with endemic fungal infections, including both histoplasmosis and blastomycosis, frequently have more severe pulmonary disease as well as dissemination. Blastomycosis has been associated with a mortality rate of 36% in these patients.



In Africa, pulmonary cases typically include bony involvement (frequently of the vertebrae), with subcutaneous abscesses of the chest wall or legs. All of the manifestations seen in African patients fall within the spectrum of blastomycosis observed in North America. The increased prevalence of chronic and disseminated bone disease in these patients may reflect a delay in diagnosis in regions where spinal disease is often treated empirically as tuberculosis.

Skin disease is the most common extrapulmonary manifestation of blastomycosis. Two types of skin lesions occur: verrucous (more common) and ulcerative. Osteomyelitis occurs in as many as one-fourth of *B. dermatitidis* infections. The vertebrae, pelvis, sacrum,

skull, ribs, and long bones are most frequently involved. Patients with *B. dermatitidis* osteomyelitis often present with contiguous soft-tissue abscesses or chronic draining sinuses. In men, blastomycosis may involve the prostate and epididymis. Central nervous system (CNS) disease occurs in fewer than 5% of immunocompetent patients with blastomycosis. A recent multicenter review identified 22 patients with CNS disease, of whom 12 (54%) met at least one criterion for immunosuppression; although most cases of CNS blastomycosis are associated with infection at other sites, 22.7% of the reviewed cases had only CNS involvement. CNS disease, usually presenting as a brain abscess, has been reported in $\sim 40\%$ of cases in patients with AIDS. Less common forms of CNS disease are cranial or spinal epidural abscess and meningitis.

DIAGNOSIS

Definitive diagnosis of blastomycosis requires growth of the organism from sputum, bronchial washings, pus, or biopsy material. Specimens should be inoculated onto a fungal medium such as Sabouraud dextrose agar, with or without chloramphenicol. *B. dermatitidis* is generally visible in 5–10 days but may require incubation for up to 30 days if only a few organisms are present in the specimen. A presumptive diagnosis may be based on demonstration of the characteristic broad-based budding yeast by microscopic examination of wet preps of sputum in pneumonia or of skin-lesion scrapings. Serologic testing for antibodies to *B. dermatitidis* by complement fixation, immunodiffusion, or enzyme immunoassay is of little value for diagnosis because of limited sensitivity and specificity as well as cross-reactivity with other fungal antigens.

A *Blastomyces* antigen assay that detects antigen in urine and serum is commercially available and is reasonably sensitive and specific (MiraVista Diagnostics, Indianapolis, IN). Antigen detection appears to be more sensitive in urine than in serum. This antigen test may be useful for monitoring of patients during therapy or for early detection of relapse. Chemiluminescent DNA probes (AccuProbe; GenProbe Inc., San Diego, CA) are commonly used to confirm identification of *B. dermatitidis* once growth has been detected in culture. Repetitive sequence-based PCR is available (DiversiLab System; bioMérieux, Durham, NC). Molecular identification techniques are currently used only to supplement traditional diagnostic methods.

TREATMENT BLASTOMYCOSIS

The Infectious Diseases Society of America has published guidelines for the treatment of blastomycosis. Selection of an appropriate therapeutic regimen must be based on the clinical form and severity of the disease, the immune status of the patient, and the toxicity of the antifungal agent (Table 238-1). Although spontaneous cures of acute pulmonary infection are well documented, there are no criteria by which to distinguish patients whose disease will progress or resolve without treatment. Thus all patients with blastomycosis should be treated.

Itraconazole is the agent of choice for immunocompetent patients with mild to moderate pulmonary or non-CNS extrapulmonary disease. Therapy is continued for 6–12 months. Amphotericin B (AmB) is the preferred initial treatment for patients who are severely immunocompromised, who have life-threatening disease or CNS disease, or whose disease progresses during treatment with itraconazole. Although not rigorously studied, lipid formulations of AmB provide an alternative for patients who cannot tolerate AmB deoxycholate. Most patients with non-CNS disease whose clinical condition improves after an initial course of AmB (usually 2 weeks in duration) can be switched to itraconazole to complete 6–12 months of therapy. Fluconazole, because of its excellent penetration of the CNS, is useful in the treatment of patients with brain abscess or meningitis after an initial course of AmB.

Voriconazole has been used successfully to treat refractory blastomycosis, blastomycosis in immunosuppressed patients, and—given

TABLE 238-1 TREATMENT OF BLASTOMYCOSIS^a

Disease	Primary Therapy	Alternative Therapy
Immunocompetent Patient/Life-Threatening Disease		
Pulmonary	Lipid AmB, 3–5 mg/kg qd, or AmB deoxycholate, 0.7–1.0 mg/kg qd (total dose: 1.5–2.5 g)	Itraconazole, 200–400 mg/d (once patient's condition has stabilized)
Disseminated		
CNS	Lipid AmB, 3–5 mg/kg qd, or AmB deoxycholate, 0.7–1.0 mg/kg qd (total dose: at least 2 g)	Fluconazole, 800 mg/d (if patient is intolerant to full course of AmB)
Non-CNS	Lipid AmB, 3–5 mg/kg qd, or AmB deoxycholate, 0.7–1.0 mg/kg qd (total dose: 1.5–2.5 g)	Itraconazole, 200–400 mg/d (once patient's condition has stabilized)
Immunocompetent Patient/Non-Life-Threatening Disease		
Pulmonary or disseminated (non-CNS)	Itraconazole, 200–400 mg/d, or Lipid AmB, 3–5 mg/kg qd, or AmB deoxycholate, 0.5–0.7 mg/kg qd (in patients intolerant to itraconazole or whose disease progresses despite therapy)	Fluconazole, 400–800 mg/d, or Ketoconazole, 400–800 mg/d
Immunocompromised Patient^b		
All infections	Lipid AmB, 3–5 mg/kg qd, or AmB deoxycholate, 0.7–1.0 mg/kg qd (total dose: 1.5–2.5 g)	Itraconazole, 200–400 mg/d (non-CNS disease, once clinically improved)

^aTherapy is generally given for 6–12 months. For bone and joint disease, a 12-month course is usually necessary. ^bSuppressive therapy with itraconazole may be considered for patients whose immunocompromised state continues. Fluconazole (800 mg/d) may be useful for patients who have CNS disease or cannot tolerate itraconazole.

Abbreviations: AmB, amphotericin B; CNS, central nervous system.

its good penetration of cerebrospinal fluid—CNS disease. Posaconazole has also been used for refractory pulmonary disease. The echinocandins have variable activity against *B. dermatitidis* and therefore are not used in the treatment of blastomycosis.

PROGNOSIS

Cure rates are 90–95% among compliant immunocompetent patients given itraconazole for mild to moderate pulmonary and extrapulmonary disease without CNS involvement. Bone and joint disease usually requires 12 months of therapy. The fewer than 5% of infections that relapse after an initial course of itraconazole usually respond well to a second treatment course.

ACKNOWLEDGMENT

The authors thank Dr. Stanley W. Chapman, Professor Emeritus, University of Mississippi, for his continued help and support and for his contributions to this chapter in an earlier edition.

239 Cryptococcosis

Arturo Casadevall

DEFINITION AND ETIOLOGY

Cryptococcus, a genus of yeast-like fungi, is the etiologic agent of cryptococcosis. Both species, *C. neoformans* and *C. gattii*, can cause cryptococcosis in humans. The two varieties of *C. neoformans*—*grubii* and *neoformans*—correlate with serotypes A and D, respectively. *C. gattii*, although not divided into varieties, also is antigenically diverse, encompassing serotypes B and C. Most clinical microbiology laboratories do not routinely distinguish between *C. neoformans* and *C. gattii*, or among varieties, but rather identify and report all isolates simply as *C. neoformans*.

EPIDEMIOLOGY

Cryptococcosis was first described in the 1890s but remained relatively rare until the mid-twentieth century, when advances in diagnosis and increases in the number of immunosuppressed individuals markedly raised its reported prevalence. Although serologic evidence of cryptococcal infection is common among immunocompetent individuals, cryptococcal disease (cryptococcosis) is relatively rare in the absence of impaired immunity. Individuals at high risk for disease due to *C. neoformans* include patients with hematologic malignancies, recipients of solid organ transplants who require ongoing immunosuppressive therapy, persons whose medical conditions necessitate glucocorticoid therapy, and patients with advanced HIV infection and CD4+ T lymphocyte counts of <200/μL. In contrast, *C. gattii*-related disease is not associated with specific immune deficits and often occurs in immunocompetent individuals.



Cryptococcal infection is acquired from the environment. *C. neoformans* and *C. gattii* inhabit different ecologic niches. *C. neoformans* is frequently found in soils contaminated with avian excreta and can easily be recovered from shaded and humid soils contaminated with pigeon droppings. In contrast, *C. gattii* is not found in bird feces. Instead, it inhabits a variety of arboreal species, including several types of eucalyptus tree. *C. neoformans* strains are found throughout the world; however, var. *grubii* (serotype A) strains are far more common than var. *neoformans* (serotype D) strains among both clinical and environmental isolates. The geographic distribution of *C. gattii* was thought to be largely limited to tropical regions until an outbreak of cryptococcosis caused by a new serotype B strain began in Vancouver in 1999. This outbreak has extended into the United States, and *C. gattii* infections are being encountered increasingly in several states in the Pacific Northwest.

The global burden of cryptococcosis was recently estimated at ~1 million cases, with >600,000 deaths annually. Thus cryptococci are important human pathogens. Since the onset of the HIV pandemic in the early 1980s, the overwhelming majority of cryptococcosis cases have occurred in patients with AIDS (Chap. 226). To comprehend the impact of HIV infection on the epidemiology of cryptococcosis, it is instructive to note that in the early 1990s there were >1000 cases of cryptococcal meningitis each year in New York City—a figure far exceeding that for all cases of bacterial meningitis. With the advent of effective antiretroviral therapy, the incidence of AIDS-related cryptococcosis has been sharply reduced among treated individuals. Thus most cases of cryptococcosis now occur in resource-limited regions of the world. The disease remains distressingly common in regions where antiretroviral therapy is not readily available (e.g., parts of Africa and Asia); in these regions, up to one-third of patients with AIDS have cryptococcosis. Among HIV-infected persons, those with a decreased percentage of memory B cells expressing IgM may be at greater risk for cryptococcosis.

Cryptococcal infection is acquired by inhalation of aerosolized infectious particles. The exact nature of these particles is not known; the two leading candidate forms are small desiccated yeast cells and basidiospores. Little is known about the pathogenesis of initial infection. Serologic studies have shown that cryptococcal infection is acquired in childhood, but it is not known whether the initial infection is symptomatic. Given that cryptococcal infection is common while disease is rare, the consensus is that pulmonary defense mechanisms in immunologically intact individuals are highly effective at containing this fungus. It is not clear whether initial infection leads to a state of immunity or whether most individuals are subject throughout life to frequent and recurrent infections that resolve without clinical disease. However, evidence indicates that some human cryptococcal infections lead to a state of latency in which viable organisms are harbored for prolonged periods, possibly in granulomas. Thus the inhalation of cryptococcal cells and/or spores can be followed by either clearance or establishment of the latent state. The consequences of prolonged harboring of cryptococcal cells in the lung are not known, but evidence from animal studies indicates that the organisms' prolonged presence could alter the immunologic milieu in the lung and predispose to allergic airway disease.

Cryptococcosis usually presents clinically as chronic meningoencephalitis. The mechanisms by which the fungus undergoes extrapulmonary dissemination and enters the central nervous system (CNS) remain poorly understood. The mechanism by which cryptococcal cells cross the blood–brain barrier is a subject of intensive study. Current evidence suggests that both direct fungal-cell migration across the endothelium and fungal-cell carriage inside macrophages as “Trojan horse” invaders can occur. *Cryptococcus* species have well-defined virulence factors that include the expression of the polysaccharide capsule, the ability to make melanin, and the elaboration of enzymes (e.g., phospholipase and urease) that enhance the survival of fungal cells in tissue. Among these virulence factors, the capsule and melanin production have been most extensively studied. The cryptococcal capsule is antiphagocytic, and the capsular polysaccharide has been associated with numerous deleterious effects on host immune function. Cryptococcal infections can elicit little or no tissue inflammatory response. The immune dysfunction seen in cryptococcosis has been attributed to the release of copious amounts of capsular polysaccharide into tissues, where it probably interferes with local immune responses (Fig. 239-1). In clinical practice, the capsular polysaccharide is the antigen that is measured as a diagnostic marker of cryptococcal infection.

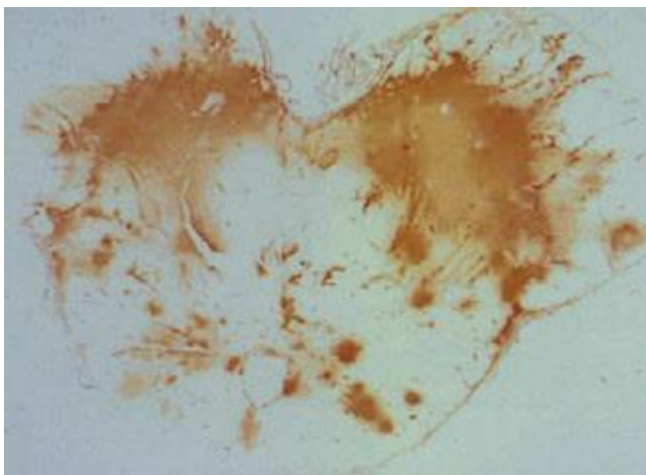


FIGURE 239-1 Cryptococcal antigen in human brain tissue, as revealed by immunohistochemical staining. Brown areas show polysaccharide deposits in the midbrain of a patient who died of cryptococcal meningitis. (Reprinted with permission from SC Lee et al: *Hum Pathol* 27:839, 1996.)

APPROACH TO THE PATIENT:

Cryptococcosis

Cryptococcosis should be included in the differential diagnosis when any patient presents with findings suggestive of chronic meningitis. Concern about cryptococcosis is heightened by a history of headache and neurologic symptoms in a patient with an underlying immunosuppressive disorder or state that is associated with an increased incidence of cryptococcosis, such as advanced HIV infection or solid organ transplantation.

CLINICAL MANIFESTATIONS

The clinical manifestations of cryptococcosis reflect the site of fungal infection. The spectrum of disease caused by *Cryptococcus* species consists predominantly of meningoencephalitis and pneumonia, but skin and soft tissue infections also occur; in fact, cryptococcosis can affect any tissue or organ. CNS involvement usually presents as signs and symptoms of chronic meningitis, such as headache, fever, lethargy, sensory deficits, memory deficits, cranial nerve palsy, vision deficits, and meningismus. Cryptococcal meningitis differs from bacterial meningitis in that many *Cryptococcus*-infected patients present with symptoms of several weeks' duration. In addition, classic characteristics of meningeal irritation, such as meningismus, may be absent in cryptococcal meningitis. Indolent cases can present as subacute dementia. Meningeal cryptococcosis can lead to sudden catastrophic vision loss.

Pulmonary cryptococcosis usually presents as cough, increased sputum production, and chest pain. Patients infected with *C. gattii* can present with granulomatous pulmonary masses known as *cryptococcomas*. Fever develops in a minority of cases. Like CNS disease, pulmonary cryptococcosis can follow an indolent course, and the majority of cases probably do not come to clinical attention. In fact, many cases are discovered incidentally during the workup of an abnormal chest radiograph obtained for other diagnostic purposes. Pulmonary cryptococcosis can be associated with antecedent diseases such as malignancy, diabetes, and tuberculosis.

Skin lesions are common in patients with disseminated cryptococcosis and can be highly variable, including papules, plaques, purpura, vesicles, tumor-like lesions, and rashes. The spectrum of cryptococcosis in HIV-infected patients is so varied and has changed so much since the advent of antiretroviral therapy that a distinction between HIV-related and HIV-unrelated cryptococcosis is no longer pertinent. In patients with AIDS and solid organ transplant recipients, the lesions of cutaneous cryptococcosis often resemble those of molluscum contagiosum (Fig. 239-2; Chap. 220e).

DIAGNOSIS

A diagnosis of cryptococcosis requires the demonstration of yeast cells in normally sterile tissues. Visualization of the capsule of fungal cells in cerebrospinal fluid (CSF) mixed with India ink is a useful rapid diagnostic technique. Cryptococcal cells in India ink have a distinctive appearance because their capsules exclude ink particles. However, the CSF India ink examination may yield negative results in patients with a low fungal burden. This examination should be performed by a trained individual, since leukocytes and fat globules can sometimes be mistaken for fungal cells. Cultures of CSF and blood that are positive for cryptococcal cells are diagnostic for cryptococcosis. In cryptococcal meningitis, CSF examination usually reveals evidence of chronic meningitis with mononuclear cell pleocytosis and increased protein levels. A particularly useful test is cryptococcal antigen (CRAG) detection in CSF and blood. The assay is based on serologic detection of cryptococcal polysaccharide and is both sensitive and specific. A positive CRAG test provides strong presumptive evidence for cryptococcosis; however, because the result is often negative in pulmonary cryptococcosis, the test is less useful in the diagnosis of pulmonary disease and is of only limited usefulness in monitoring the response to therapy.



FIGURE 239-2 Disseminated fungal infection. A liver transplant recipient developed six cutaneous lesions similar to the one shown. Biopsy and serum antigen testing demonstrated *Cryptococcus*. Important features of the lesion include a benign-appearing fleshy papule with central umbilication resembling molluscum contagiosum. (Photo courtesy of Dr. Lindsey Baden; with permission.)



In areas of Africa where there is a high prevalence of HIV infection, routine screening of blood for CRAg in HIV-infected patients with low CD4+ T lymphocyte counts may identify individuals at high risk of cryptococcal disease who are candidates for antifungal therapy. Similarly, CRAg screening has shown that a significant proportion of HIV-infected patients hospitalized with pneumonia in Thailand harbor cryptococcal infection. Inexpensive point-of-care CRAg tests that are under development could be of great diagnostic benefit in resource-limited regions.

TREATMENT CRYPTOCOCCOSIS

Both the site of infection and the immune status of the host must be considered in the selection of therapy for cryptococcosis. The disease has two general patterns of manifestation: (1) pulmonary cryptococcosis, with no evidence of extrapulmonary dissemination; and (2) extrapulmonary (systemic) cryptococcosis, with or without meningoencephalitis. Pulmonary cryptococcosis in an immunocompetent host sometimes resolves without therapy. However, given the propensity of *Cryptococcus* species to disseminate from the lung, the inability to gauge the host's immune status precisely, and the availability of low-toxicity therapy in the form of fluconazole, the current recommendation is for pulmonary cryptococcosis in an immunocompetent individual to be treated with fluconazole (200–400 mg/d for 3–6 months). Extrapulmonary cryptococcosis without CNS involvement in an immunocompetent host can be treated with the same regimen, although amphotericin B (AmB; 0.5–1 mg/kg daily for 4–6 weeks) may be required for more severe cases. In general, extrapulmonary cryptococcosis without CNS involvement requires less intensive therapy, with the caveat that morbidity and death in cryptococcosis are associated with meningeal involvement. Thus the decision to categorize cryptococcosis as “extrapulmonary without CNS involvement” should be made only after careful evaluation of the CSF reveals no evidence of cryptococcal infection. For CNS involvement in a host without AIDS or obvious immune impairment, most authorities recommend initial therapy with AmB (0.5–1 mg/kg daily) during an induction phase, which is followed by prolonged therapy with fluconazole (400 mg/d) during a consolidation phase. For cryptococcal meningoencephalitis without a concomitant immunosuppressive condition, the recommended regimen is AmB (0.5–1 mg/kg) plus flucytosine (100 mg/kg) daily for

6–10 weeks. Alternatively, patients can be treated with AmB (0.5–1 mg/kg) plus flucytosine (100 mg/kg) daily for 2 weeks and then with fluconazole (400 mg/d) for at least 10 weeks. Patients with immunosuppression are treated with the same initial regimens except that consolidation therapy with fluconazole is given for a prolonged period to prevent relapse.

Cryptococcosis in patients with HIV infection always requires aggressive therapy and is considered incurable unless immune function improves. Consequently, therapy for cryptococcosis in the setting of AIDS has two phases: induction therapy (intended to reduce the fungal burden and alleviate symptoms) and lifelong maintenance therapy (to prevent a symptomatic clinical relapse). Pulmonary and extrapulmonary cryptococcosis without evidence of CNS involvement can be treated with fluconazole (200–400 mg/d). In patients who have more extensive disease, flucytosine (100 mg/kg per day) may be added to the fluconazole regimen for 10 weeks, with lifelong fluconazole maintenance therapy thereafter. For HIV-infected patients with evidence of CNS involvement, most authorities recommend induction therapy with AmB. An acceptable regimen is AmB (0.7–1 mg/kg) plus flucytosine (100 mg/kg) daily for 2 weeks followed by fluconazole (400 mg/d) for at least 10 weeks and then by lifelong maintenance therapy with fluconazole (200 mg/d). Fluconazole (400–800 mg/d) plus flucytosine (100 mg/kg per day) for 6–10 weeks followed by fluconazole (200 mg/d) as maintenance therapy is an alternative. Newer triazoles like voriconazole and posaconazole are highly active against cryptococcal strains and appear effective clinically, but clinical experience with these agents in the treatment of cryptococcosis is limited. Lipid formulations of AmB can be substituted for AmB deoxycholate in patients with renal impairment. Neither caspofungin nor micafungin is effective against *Cryptococcus* species; consequently, neither drug has a role in the treatment of cryptococcosis. Cryptococcal meningoencephalitis is often associated with increased intracranial pressure, which is believed to be responsible for damage to the brain and cranial nerves. Appropriate management of CNS cryptococcosis requires careful attention to the management of intracranial pressure, including the reduction of pressure by repeated therapeutic lumbar puncture and the placement of shunts. Recent studies suggest that the addition of a short course of interferon γ to antifungal therapy in patients with HIV infection increases clearance of cryptococci from the CSF.

In HIV-infected patients with previously treated cryptococcosis who are receiving fluconazole maintenance therapy, it may be possible to discontinue antifungal drug treatment if antiretroviral therapy results in immunologic improvement. However, certain recipients of maintenance therapy who have a history of successfully treated cryptococcosis can develop a troublesome immune reconstitution syndrome when antiretroviral therapy produces a rebound in immunologic function.

PROGNOSIS AND COMPLICATIONS

Even with antifungal therapy, cryptococcosis is associated with high rates of morbidity and death. For the majority of patients with cryptococcosis, the most important prognostic factors are the extent and the duration of the underlying immunologic deficits that predisposed them to develop the disease. Therefore, cryptococcosis is often curable with antifungal therapy in individuals with no apparent immunologic dysfunction, but, in patients with severe immunosuppression (e.g., those with AIDS), the best that can be hoped for is that antifungal therapy will induce remission, which can then be maintained with lifelong suppressive therapy. Before the advent of antiretroviral therapy, the median overall survival period for AIDS patients with cryptococcosis was <1 year. Cryptococcosis in patients with underlying neoplastic disease has a particularly poor prognosis. For CNS cryptococcosis, poor prognostic markers are a CSF assay positive for yeast cells on initial India ink examination (evidence of a heavy fungal burden), high CSF pressure, low CSF glucose levels, low CSF pleocytosis (<2/ μ L), recovery of yeast cells from extraneural sites, absence of antibody to capsular

1342 polysaccharide, a CSF or serum cryptococcal antigen level of $\geq 1:32$, and concomitant glucocorticoid therapy or hematologic malignancy. A response to treatment does not guarantee cure since relapse of cryptococcosis is common even among patients with relatively intact immune systems. Complications of CNS cryptococcosis include cranial nerve deficits, vision loss, and cognitive impairment.

PREVENTION

No vaccine is available for cryptococcosis. In patients at high risk (e.g., those with advanced HIV infection and CD4+ T lymphocyte counts of $<200/\mu\text{L}$), primary prophylaxis with fluconazole (200 mg/d) is effective in reducing the prevalence of disease. Since antiretroviral therapy raises the CD4+ T lymphocyte count, it constitutes an immunologic form of prophylaxis. However, cryptococcosis in the setting of immune reconstitution has been reported in patients with HIV infection and in recipients of solid organ transplants.

240 Candidiasis

John E. Edwards, Jr.

The genus *Candida* encompasses more than 150 species, only a few of which cause disease in humans. With rare exceptions (although the exceptions are increasing in number), the human pathogens are *C. albicans*, *C. guilliermondii*, *C. krusei*, *C. parapsilosis*, *C. tropicalis*, *C. kefyr*, *C. lusitaniae*, *C. dubliniensis*, and *C. glabrata*. Ubiquitous in nature, they inhabit the gastrointestinal tract (including the mouth and oropharynx), the female genital tract, and the skin. Although cases of candidiasis have been described since antiquity in debilitated patients, the advent of *Candida* species as common human pathogens dates to the introduction of modern therapeutic approaches that suppress normal host defense mechanisms. Of these relatively recent advances, the most important is the use of antibacterial agents that alter the normal human microbiota and allow nonbacterial species to become more prevalent in the commensal flora. With the introduction of antifungal agents, the causes of *Candida* infections shifted from an almost complete dominance of *C. albicans* to the common involvement of *C. glabrata* and the other species listed above. The non-*albicans* species now account for approximately half of all cases of candidemia and hematogenously disseminated candidiasis. Recognition of this change is clinically important, since the various species differ in susceptibility to the newer antifungal agents. In developed countries, where medical therapeutics are commonly used, *Candida* species are now among the most common nosocomial pathogens.

Candida is a small, thin-walled, ovoid yeast that measures 4–6 μm in diameter and reproduces by budding. Organisms of this genus occur in three forms in tissue: blastospores, pseudohyphae, and hyphae. *Candida* grows readily on simple medium; lysis centrifugation enhances its recovery from blood. Species are identified by biochemical testing (currently with automated devices) or on special agar (e.g., CHROMagar).

EPIDEMIOLOGY

Candida organisms are ubiquitous in nature; worldwide, these fungi are present in humans as commensals, in animals, in foods, and on inanimate objects. In developed countries, where advanced medical therapeutics are commonly used (see “Treatment,” below), *Candida* species are now among the most common health care–associated pathogens. In the United States, these species are the fourth most common isolates from the blood of hospitalized patients. In countries where advanced medical care is rarely available, mucocutaneous *Candida* infections, such as thrush, are more common than deep organ infections, which rarely occur; however, the

incidence of deep organ candidiasis increases steadily as advances in health care—such as therapy with broad-spectrum antibiotics, more aggressive treatment of cancer, and the use of immunosuppression for sustaining organ transplants—are introduced and implemented. In recent decades, as a result of the HIV epidemic, the incidence of thrush and *Candida* esophagitis has increased substantially. In aggregate, the global incidence of infections due to *Candida* species has risen steadily over the past few decades.

PATHOGENESIS

In the most serious form of *Candida* infection, the organisms disseminate hematogenously and form microabscesses and small macroabscesses in major organs. Although the exact mechanism is not known, *Candida* probably enters the bloodstream from mucosal surfaces after growing to large numbers as a consequence of bacterial suppression by antibacterial drugs; alternatively, in some instances, the organism may enter from the skin. A change from the blastospore stage to the pseudohyphal and hyphal stages is generally considered integral to the organism’s penetration into tissue. However, *C. glabrata* can cause extensive infection even though it does not transform into pseudohyphae or hyphae. Adherence to both epithelial and endothelial cells, thought to be the first step in invasion and infection, has been studied extensively, and several adhesins have been identified. Biofilm formation also is considered important in pathogenesis. Numerous reviews of cases of hematogenously disseminated candidiasis have identified the predisposing factors or conditions associated with disseminated disease (Table 240-1). Women who receive antibacterial agents may develop vaginal candidiasis.

Innate immunity is the most important defense mechanism against hematogenously disseminated candidiasis, and the neutrophil is the most important component of this defense. Macrophages also play an important defensive role. STAT1, Dectin-1, CARD9, and T_H1 and T_H17 lymphocytes contribute significantly to innate defense (see “Clinical Manifestations,” below). Although many immunocompetent individuals have antibodies to *Candida*, the role of these antibodies in defense against the organism is not clear. Multiple genetic polymorphisms that predispose to disseminated candidiasis will most likely be identified in future studies.

CLINICAL MANIFESTATIONS

Mucocutaneous Candidiasis Thrush is characterized by white, adherent, painless, discrete or confluent patches in the mouth, on the tongue, or in the esophagus, occasionally with fissuring at the corners of the mouth. This form of disease caused by *Candida* can also occur at points of contact with dentures. Organisms are identifiable in gram-stained scrapings from lesions. The occurrence of thrush in a young, otherwise healthy-appearing person should prompt an investigation for underlying HIV infection. More commonly, thrush is seen as a nonspecific manifestation of severe debilitating illness. Vulvovaginal candidiasis is accompanied by pruritus, pain, and vaginal discharge which is usually thin but may contain whitish “curds” in severe cases. A subset of patients with recurrent vulvovaginitis have a deficiency in the surface expression of Dectin-1, a major recognition factor for β -glucan on *Candida*. This deficiency leads to suboptimal functioning of the CARD9 pathway, which ultimately increases the propensity for recurrent vaginal infections.

TABLE 240-1 WELL-RECOGNIZED FACTORS AND CONDITIONS PREDISPOSING TO HEMATOGENOUSLY DISSEMINATED CANDIDIASIS

Antibacterial agents	Abdominal and thoracic surgery
Indwelling intravenous catheters	Cytotoxic chemotherapy
Hyperalimentation fluids	Immunosuppressive agents for organ transplantation
Indwelling urinary catheters	Respirators
Parenteral glucocorticoids	Neutropenia
Severe burns	Low birth weight (neonates)
HIV-associated low CD4+ T cell counts	Diabetes



FIGURE 240-1 Macronodular skin lesions associated with hematogenously disseminated candidiasis. *Candida* organisms are usually but not always visible on histopathologic examination. The fungi grow when a portion of the biopsied specimen is cultured. Therefore, for optimal identification, both histopathology and culture should be performed. (Image courtesy of Dr. Noah Craft and the Victor Newcomer collection at UCLA, archived by Logical Images, Inc.; with permission.)

Other *Candida* skin infections include *paronychia*, a painful swelling at the nail-skin interface; *onychomycosis*, a fungal nail infection rarely caused by this genus; *intertrigo*, an erythematous irritation with redness and pustules in the skin folds; *balanitis*, an erythematous-pustular infection of the glans penis; *erosio interdigitalis blastomycetica*, an infection between the digits of the hands or toes; *folliculitis*, with pustules developing most frequently in the area of the beard; *perianal candidiasis*, a pruritic, erythematous, pustular infection surrounding the anus; and *diaper rash*, a common erythematous-pustular perineal infection in infants. *Generalized disseminated cutaneous candidiasis*, another form of infection that occurs primarily in infants, is characterized by widespread eruptions over the trunk, thorax, and extremities. The diagnostic macronodular lesions of hematogenously disseminated candidiasis (Fig. 240-1) indicate a high probability of dissemination to multiple organs as well as the skin. While the lesions are seen predominantly in immunocompromised patients treated with cytotoxic drugs, they may also develop in patients without neutropenia.

Chronic mucocutaneous candidiasis is a heterogeneous infection of the hair, nails, skin, and mucous membranes that persists despite intermittent therapy. The onset of disease usually comes in infancy or within the first two decades of life but in rare cases comes in later life. The condition may be mild and limited to a specific area of the skin or nails, or it may take a severely disfiguring form (*Candida* granuloma) characterized by exophytic outgrowths on the skin. Chronic mucocutaneous candidiasis is usually associated with specific immunologic dysfunction; most frequently reported is a failure of T lymphocytes to proliferate or to excrete cytokines in response to stimulation by *Candida* antigens in vitro. A subset of the affected patients have mutations in the STAT1 gene resulting in an insufficiency of interferon γ , interleukin 17, and interleukin 22.

Approximately half of patients with chronic mucocutaneous candidiasis have associated endocrine abnormalities that together are designated the autoimmune polyendocrinopathy–candidiasis–ectodermal dystrophy (APECED) syndrome. This syndrome is due to mutations in the autoimmune regulator (*AIRE*) gene and is most prevalent among Finns, Iranian Jews, Sardinians, northern Italians, and Swedes. Conditions that usually follow the onset of the disease include hypoparathyroidism, adrenal insufficiency, autoimmune thyroiditis, Graves' disease, chronic active hepatitis, alopecia, juvenile-onset pernicious anemia, malabsorption, and primary hypogonadism. In addition, dental enamel dysplasia, vitiligo, pitted nail dystrophy, and calcification of the tympanic membranes may occur. Patients with

chronic mucocutaneous candidiasis rarely develop hematogenously disseminated candidiasis, probably because their neutrophil function remains intact.

Deeply Invasive Candidiasis Deeply invasive *Candida* infections may or may not be due to hematogenous seeding. Deep esophageal infection may result from penetration by organisms from superficial esophageal erosions; joint or deep wound infection from contiguous spread of organisms from the skin; kidney infection from catheter-initiated spread of organisms through the urinary tract; infection of intraabdominal organs and the peritoneum from perforation of the gastrointestinal tract; and gallbladder infection from retrograde migration of organisms from the gastrointestinal tract into the biliary drainage system.

However, far more commonly, deeply invasive candidiasis results from hematogenous seeding of various organs as a complication of candidemia. Once the organism gains access to the intravascular compartment (either from the gastrointestinal tract or, less often, from the skin through the site of an indwelling intravascular catheter), it may spread hematogenously to a variety of deep organs. The brain, chorioretina (Fig. 240-2), heart, and kidneys are most commonly infected and the liver and spleen less commonly so (most often in neutropenic patients). In fact, nearly any organ can become involved, including the endocrine glands, pancreas, heart valves (native or prosthetic), skeletal muscle, joints (native or prosthetic), bone, and meninges. *Candida* organisms can also spread hematogenously to the skin and cause classic macronodular lesions (Fig. 240-1). Frequently, painful muscular involvement also is evident beneath the area of affected skin. Chorioretinal involvement and skin involvement are highly significant, since both findings are associated with a very high probability of abscess formation in multiple deep organs as a result of generalized hematogenous seeding. Ocular involvement (Fig. 240-2) may require specific treatment (e.g., partial vitrectomy or intraocular injection of antifungal agents) to prevent permanent blindness. An ocular examination is indicated for all patients with candidemia, whether or not they have ocular manifestations.

DIAGNOSIS

The diagnosis of *Candida* infection is established by visualization of pseudohyphae or hyphae on wet mount (saline and 10% KOH), tissue Gram's stain, periodic acid–Schiff stain, or methenamine silver stain in the presence of inflammation. Absence of organisms on hematoxylin-eosin staining does not reliably exclude *Candida* infection. The most



FIGURE 240-2 Hematogenous *Candida* endophthalmitis. A classic off-white lesion projecting from the chorioretina into the vitreous causes the surrounding haze. The lesion is composed primarily of inflammatory cells rather than organisms. Lesions of this type may progress to cause extensive vitreal inflammation and eventual loss of the eye. Partial vitrectomy, combined with IV and possibly intravitreal antifungal therapy, may be helpful in controlling the lesions. (Image courtesy of Dr. Gary Holland; with permission.)

TABLE 240-2 TREATMENT OF MUCOCUTANEOUS CANDIDAL INFECTIONS

Disease	Preferred Treatment	Alternatives
Cutaneous	Topical azole	Topical nystatin
Vulvovaginal	Oral fluconazole (150 mg) or azole cream or suppository	Nystatin suppository
Thrush	Clotrimazole troches	Nystatin, fluconazole
Esophageal	Fluconazole tablets (100–200 mg/d) or itraconazole solution (200 mg/d)	Caspofungin, micafungin, or amphotericin B

challenging aspect of diagnosis is determining which patients with *Candida* isolates have hematogenously disseminated candidiasis. For instance, recovery of *Candida* from sputum, urine, or peritoneal catheters may indicate mere colonization rather than deep-seated infection, and *Candida* isolation from the blood of patients with indwelling intravascular catheters may reflect inconsequential seeding of the blood from or growth of the organisms on the catheter. Despite extensive research into both antigen and antibody detection systems, there is currently no widely available and validated diagnostic test to distinguish patients with inconsequential seeding of the blood from those whose positive blood cultures represent hematogenous dissemination to multiple organs. Many studies are under way to establish the utility of the β -glucan test; at present, its greatest utility is its negative predictive value (~90%). Meanwhile, the presence of ocular or macronodular skin lesions is highly suggestive of widespread infection of multiple deep organs. Although extensive research is being conducted on other tests for infection, such as PCR, none of these tests is fully validated or widely available at present.

TREATMENT CANDIDA INFECTIONS

MUCOCUTANEOUS CANDIDA INFECTION

The treatment of mucocutaneous candidiasis is summarized in [Table 240-2](#).

CANDIDEMIA AND SUSPECTED HEMATOGENOUSLY DISSEMINATED CANDIDIASIS

All patients with candidemia are treated with a systemic antifungal agent. A certain percentage of patients, including many of those

who have candidemia associated with an indwelling intravascular catheter, probably have “benign” candidemia rather than deep-organ seeding. However, because there is no reliable way to distinguish benign candidemia from deep-organ infection, and because antifungal drugs less toxic than amphotericin B are available, antifungal treatment for candidemia—with or without clinical evidence of deep-organ involvement—has become the standard of practice. In addition, if an indwelling intravascular catheter is present, it is best to remove or replace the device whenever feasible.

The drugs used for the treatment of candidemia and suspected disseminated candidiasis are listed in [Table 240-3](#). Various lipid formulations of amphotericin B, three echinocandins, and the azoles fluconazole and voriconazole are used; no agent within a given class has been clearly identified as superior to the others. Most institutions choose an agent from each class on the basis of their own specific microbial epidemiology, strategies to minimize toxicities, and cost considerations. Unless azole resistance is considered likely, fluconazole is the agent of choice for the treatment of candidemia and suspected disseminated candidiasis in nonneutropenic, hemodynamically stable patients. Initial treatment in the context of likely azole resistance depends, as mentioned above, on the epidemiology of the individual hospital. For example, certain hospitals have a high rate of recovery of *C. glabrata*, while others do not. At institutions where non-*albicans* *Candida* species are frequently recovered, therapy with an echinocandin is typically started while the results of sensitivity testing are awaited. For hemodynamically unstable or neutropenic patients, initial treatment with broader-spectrum agents is desirable; these drugs include polyenes, echinocandins, or later-generation azoles such as voriconazole. Once the clinical response has been assessed and the pathogen specifically identified, the regimen can be altered accordingly. At present, the vast majority of *C. albicans* isolates are sensitive to fluconazole. Isolates of *C. glabrata* and *C. krusei* are less sensitive to fluconazole and more sensitive to polyenes and echinocandins. *C. parapsilosis* is less sensitive to echinocandins in vitro; however, this lesser sensitivity is considered nonsignificant.

Some generalizations exist regarding the management of specific *Candida* infections. Recovery of *Candida* from sputum is almost never indicative of underlying pulmonary candidiasis and does not by itself warrant antifungal treatment. Similarly, *Candida* in the urine of a patient with an indwelling bladder catheter may represent colonization only rather than bladder or kidney infection; however, the threshold for systemic treatment is lower in severely ill patients in

TABLE 240-3 AGENTS FOR THE TREATMENT OF DISSEMINATED CANDIDIASIS

Agent	Route of Administration	Dose ^a	Comment
Amphotericin B deoxycholate	IV only	0.5–1.0 mg/kg daily	Being replaced by lipid formulations
Amphotericin B lipid formulations			Not FDA approved as primary therapy, but used commonly because less toxic than amphotericin B deoxycholate
Liposomal (AmBisome, Abelcet)	IV only	3.0–5.0 mg/kg daily	
Lipid complex (ABLX)	IV only	3.0–5.0 mg/kg daily	
Colloidal dispersion (ABCD)	IV only	3.0–5.0 mg/kg daily	Associated with frequent infusion reactions
Azoles ^b			
Fluconazole	IV and oral	400 mg/d	Most commonly used
Voriconazole	IV and oral	400 mg/d	Multiple drug interactions
			Approved for candidemia in nonneutropenic patients
Echinocandins			Broad spectrum against <i>Candida</i> species; approved for disseminated candidiasis
Caspofungin	IV only	50 mg/d	
Anidulafungin	IV only	100 mg/d	
Micafungin	IV only	100 mg/d	

^aFor loading doses and adjustments in renal failure, see Pappas PG et al: Clinical practice guidelines for the management of candidiasis: 2009 update by the Infectious Diseases Society of America. *Clin Infect Dis* 48:503, 2009. The recommended duration of therapy is 2 weeks beyond the last positive blood culture and the resolution of signs and symptoms of infection. ^bAlthough ketoconazole is approved for the treatment of disseminated candidiasis, it has been replaced by the newer agents listed in this table. Posaconazole has been approved for prophylaxis in neutropenic patients and for oropharyngeal candidiasis. FDA, U.S. Food and Drug Administration.

241 Aspergillosis

David W. Denning

this category since it is impossible to distinguish colonization from lower or upper urinary tract infection. If the isolate is *C. albicans*, most clinicians use oral fluconazole rather than a bladder washout with amphotericin B, which was more commonly used in the past. Caspofungin has been used with success; although echinocandins are poorly excreted into the urine, they may be an option, especially for non-*albicans* isolates. The doses and duration are the same as for disseminated candidiasis. The significance of the recovery of *Candida* from abdominal drains in postoperative patients is unclear, but again the threshold for treatment is generally low because most of the affected patients have been subjected to factors predisposing to disseminated candidiasis.

Removal of the infected valve and long-term antifungal therapy constitute appropriate treatment for *Candida* endocarditis. Although definitive studies are not available, patients usually are treated for weeks with a systemic antifungal agent (Table 240-2) and then given chronic suppressive therapy for months or years (sometimes indefinitely) with an oral azole (usually fluconazole at 400–800 mg/d).

Hematogenous *Candida* endophthalmitis is a special problem requiring ophthalmologic consultation. When lesions are expanding or are threatening the macula, an IV polyene combined with flucytosine (25 mg/kg four times daily) has been the regimen of choice, although comparative studies with other regimens have not yet been reported. As more data on the azoles and echinocandins become available, new strategies involving these agents are developing. Of paramount importance is the decision to perform a partial vitrectomy. This procedure debulks the infection and can preserve sight, which may otherwise be lost as a result of vitreal scarring. All patients with candidemia should undergo ophthalmologic examination because of the relatively high frequency of this ocular complication. Not only can this examination detect a developing eye lesion early in its course; in addition, identification of a lesion signifies a probability of ~90% of deep-organ abscesses and may prompt prolongation of therapy for candidemia beyond the recommended 2 weeks after the last positive blood culture. Although the basis for the consensus is a very small data set, the recommended treatment for *Candida* meningitis is a polyene (Table 240-3) plus flucytosine (25 mg/kg four times daily). Successful treatment of *Candida*-infected prosthetic material (e.g., an artificial joint) nearly always requires removal of the infected material followed by long-term administration of an antifungal agent selected on the basis of the isolate's sensitivity and the logistics of administration.

PROPHYLAXIS

The use of antifungal agents to prevent *Candida* infections has been controversial, but some general principles have emerged. Most centers administer prophylactic fluconazole (400 mg/d) to recipients of allogeneic stem cell transplants. High-risk liver transplant recipients also are given fluconazole prophylaxis in most centers. The use of prophylaxis for neutropenic patients has varied considerably from center to center; many centers that elect to give prophylaxis to this population use either fluconazole (200–400 mg/d) or a lipid formulation of amphotericin B (AmBiSome, 1–2 mg/d). Caspofungin (50 mg/d) also has been recommended. Some centers have used itraconazole suspension (200 mg/d). Posaconazole (200 mg three times daily) also has been approved by the FDA for prophylaxis in neutropenic patients and is gaining in popularity.

Prophylaxis is sometimes given to surgical patients at very high risk. The widespread use of prophylaxis for nearly all patients in general surgical or medical intensive care units is not—and should not be—a common practice for three reasons: (1) the incidence of disseminated candidiasis is relatively low, (2) the cost-benefit ratio is suboptimal, and (3) increased resistance with widespread prophylaxis is a valid concern.


Prophylaxis for oropharyngeal or esophageal candidiasis in HIV-infected patients is not recommended unless there are frequent recurrences.

Aspergillosis is the collective term used to describe all disease entities caused by any one of ~50 pathogenic and allergenic species of *Aspergillus*. Only those species that grow at 37°C can cause invasive infection, although some species without this ability can cause allergic syndromes. *A. fumigatus* is responsible for most cases of invasive aspergillosis, almost all cases of chronic aspergillosis, and most allergic syndromes. *A. flavus* is more prevalent in some hospitals and causes a higher proportion of cases of sinus infections, cutaneous infections, and keratitis than *A. fumigatus*. *A. niger* can cause invasive infection but more commonly colonizes the respiratory tract and causes external otitis. *A. terreus* causes only invasive disease, usually with a poor prognosis. *A. nidulans* occasionally causes invasive infection, primarily in patients with chronic granulomatous disease.

EPIDEMIOLOGY AND ECOLOGY

Aspergillus has a worldwide distribution, most commonly growing in decomposing plant materials (i.e., compost) and in bedding. This hyaline (nonpigmented), septate, branching mold produces vast numbers of conidia (spores) on stalks above the surface of mycelial growth. Aspergilli are found in indoor and outdoor air, on surfaces, and in water from surface reservoirs. Daily exposures vary from a few to many millions of conidia; the latter high numbers of conidia are encountered in hay barns and other very dusty environments. The required size of the infecting inoculum is uncertain; however, only intense exposures (e.g., during construction work, handling of moldy bark or hay, or composting) are sufficient to cause disease in healthy immunocompetent individuals. Allergic syndromes may be exacerbated by continuous antigenic exposure arising from sinus or airway colonization or from nail infection. High-efficiency particulate air (HEPA) filtration is often protective against infection; thus HEPA filters should be installed and monitored for efficiency in operating rooms and in areas of the hospital that house high-risk patients.

The incubation period of invasive aspergillosis after exposure is highly variable, extending in documented cases from 2 to 90 days. Thus community acquisition of an infecting strain frequently manifests as invasive infection during hospitalization, although nosocomial acquisition is also common. Outbreaks usually are directly related to a contaminated air source in the hospital.

 Global aspergillosis incidence and prevalence have been estimated (Table 241-1). However, given the inadequate diagnostic capability in almost all low- and middle-income countries, the accuracy of these estimates is uncertain. The frequency of different manifestations of aspergillosis varies considerably with geographic location; most notably, chronic granulomatous sinusitis is rare outside the Middle East and India, and fungal keratitis is particularly common in Nepal, Myanmar, Bhutan, and India (800 and 113 cases/100,000 population, respectively). The potential effects of chronic pulmonary aspergillosis after pulmonary tuberculosis have only recently been appreciated and include life-threatening hemoptysis, misdiagnosis of smear-negative tuberculosis, and general exacerbation of posttuberculosis morbidity.

RISK FACTORS AND PATHOGENESIS

The primary risk factors for invasive aspergillosis are profound neutropenia and glucocorticoid use; risk increases with longer duration of these conditions. Higher doses of glucocorticoids increase the risk of both acquisition of invasive aspergillosis and death from the infection. Neutrophil and/or phagocyte dysfunction also is an important risk factor, as evidenced by aspergillosis in chronic granulomatous disease, advanced HIV infection, and relapsed leukemia. An increasing incidence of invasive aspergillosis in medical intensive care units suggests that, in patients who are not immunocompromised, temporary abrogation of protective responses as a result of glucocorticoid use or a general anti-inflammatory state is a significant risk factor. Many

TABLE 241-1 DISEASE FREQUENCY AND DIAGNOSTIC SENSITIVITY FOR DIFFERENT MANIFESTATIONS OF ASPERGILLOSIS


Parameter	Type of Disease		
	Invasive	Chronic	Allergic
Incidence/100,000 ^a	8.6	10.4	7 ^b
Prevalence/100,000 ^a	—	32.8	286 ^c
Global burden ^d	~200,000	~3,000,000	~10,000,000
Mortality rate without treatment	~100%	~50%	<1%
Respiratory Diagnostic Sensitivity^e			
Culture	✓	✓	✓
Microscopy	✓	?	?
Antigen	✓✓✓	?	✓✓✓
Real-time PCR	✓✓✓	✓✓✓	✓✓✓
Blood Diagnostic Sensitivity^e			
Culture	✗	✗	✗
Antigen	✓✓	✓	✗
β-D-glucan	✓✓	✓	?
Real-time PCR	✓✓	?	?
IgG antibody	✓	✓✓✓✓	✓✓
IgE antibody	✗	✓✓	✓✓✓✓

^aIncidence and prevalence figures are for Europe. From www.ecdc.europa.eu/en/publications/publications/risk-assessment-impact-environmental-usage-of-triazoles-on-aspergillus-spp-resistance-to-medical-triazoles.pdf. ^bPeople are not born with allergic fungal disease; the annual frequency with which it occurs is not known. ^cAllergic bronchopulmonary aspergillosis and severe asthma with fungal sensitization. ^dGD Brown et al: Human fungal infections: the hidden killers. *Sci Transl Med* 2012;4:165rv13. ^eKey for sensitivity: 1 check = limited (as the text indicates, 10–30% for culture); 2 checks = higher; 3 checks = >80%; and 4 checks = ~95%.

Abbreviation: PCR, polymerase chain reaction.

patients have some evidence of prior pulmonary disease—typically, a history of pneumonia or chronic obstructive pulmonary disease. Therapy with infliximab, adalimumab, alemtuzumab, daclizumab, rituximab, and possibly bevacizumab therapy also carries an increased risk of invasive aspergillosis, as do severe liver disease and high levels of stored iron in bone marrow.

Patients with chronic pulmonary aspergillosis have a wide spectrum of underlying pulmonary disease, including tuberculosis, prior pneumothorax, or chronic obstructive pulmonary disease. These patients are immunocompetent except for some cytokine regulation defects, most of which are consistent with an inability to mount an inflammatory immune (T_H1-like) response or to control it adequately. Glucocorticoids accelerate disease progression.

 Allergic bronchopulmonary aspergillosis (ABPA) is associated with polymorphisms of interleukin (IL) 4Ra, IL-10, and SPA2 genes (and others) and with heterozygosity of the cystic fibrosis transmembrane conductance regulator (*CFTR*) gene. These associations

suggest a strong genetic basis for the development of a T_H2-like and “allergic” response to *A. fumigatus*.

CD4+CD25+ T (T_{reg}) cells also appear to be pivotal in determining disease phenotype. Remarkably, high-dose glucocorticoid treatment for exacerbations of ABPA almost never leads to invasive aspergillosis.

CLINICAL FEATURES AND APPROACH TO THE PATIENT (Table 241-2)

Invasive Pulmonary Aspergillosis Both the frequency of invasive disease and the pace of its progression increase with greater degrees of immunocompromise. Invasive aspergillosis is arbitrarily classified as acute and subacute, with courses of ≤1 month and 1–3 months, respectively. More than 80% of cases of invasive aspergillosis involve the lungs. The most common clinical features are no symptoms at all, fever, cough (sometimes productive), nondescript chest discomfort, trivial hemoptysis, and shortness of breath. Although the fever often responds to glucocorticoids, the disease progresses. The keys to early diagnosis in at-risk patients are a high index of suspicion, screening for circulating antigen (in leukemia), and urgent CT of the thorax. Invasive aspergillosis is one of the most common diagnostic errors revealed at autopsy.

Invasive Sinusitis The sinuses are involved in 5–10% of cases of invasive aspergillosis, especially affecting patients with leukemia and recipients of hematopoietic stem cell transplants. In addition to fever, the most common features are nasal or facial discomfort, blocked nose, and nasal discharge (sometimes bloody). Endoscopic examination of the nose reveals pale, dusky or necrotic-looking tissue in any location. CT or MRI of the sinuses is essential but does not distinguish invasive *Aspergillus* sinusitis from preexisting allergic or bacterial sinusitis early in the disease process.

Tracheobronchitis Occasionally, only the airways are infected by *Aspergillus*. The resulting manifestations range from acute or chronic bronchitis to ulcerative or pseudomembranous tracheobronchitis. These entities are particularly common among lung transplant recipients. Obstruction with mucous plugs occurs in normal individuals; in persons with ABPA, cystic fibrosis, and/or bronchiectasis; and in immunocompromised patients.

Disseminated Aspergillosis In the most severely immunocompromised patients, *Aspergillus* disseminates from the lungs to multiple organs—most often to the brain but also to the skin, thyroid, bone, kidney, liver, gastrointestinal tract, eye (endophthalmitis), and heart valve. Aside from cutaneous lesions, the most common features are gradual clinical deterioration over 1–3 days, with low-grade fever and features of mild sepsis, and nonspecific abnormalities in laboratory tests. In most cases, at least one localization becomes apparent before death. Blood cultures are almost always negative.

Cerebral Aspergillosis Hematogenous dissemination to the brain is a devastating complication of invasive aspergillosis. Single or multiple lesions may develop. In acute disease, hemorrhagic infarction is most

TABLE 241-2 MAJOR MANIFESTATIONS OF ASPERGILLOSIS

Organ	Type of Disease			
	Invasive (Acute and Subacute)	Chronic	Saprophytic	Allergic
Lung	Angioinvasive (in neutropenia), non-angioinvasive, granulomatous	Chronic cavitary, chronic fibrosing	Aspergilloma (single), airway colonization	Allergic bronchopulmonary, severe asthma with fungal sensitization, extrinsic allergic alveolitis
Sinus	Acute invasive	Chronic invasive, chronic granulomatous	Maxillary fungal ball	Allergic fungal sinusitis, eosinophilic fungal rhinosinusitis
Brain	Abscess, hemorrhagic infarction, meningitis	Granulomatous, meningitis	None	None
Skin	Acute disseminated, locally invasive (trauma, burns, IV access)	External otitis, onychomycosis	None	None
Heart	Endocarditis (native or prosthetic), pericarditis	None	None	None
Eye	Keratitis, endophthalmitis	None	None	None described

typical, and cerebral abscess is common. Rarer manifestations include meningitis, mycotic aneurysm, and cerebral granuloma (mimicking a brain tumor). Local spread from cranial sinuses also occurs. Postoperative infection develops rarely and is exacerbated by glucocorticoids, often given after neurosurgery. The presentation can be either acute or subacute, with mood changes, focal signs, seizures, and decline in mental status. MRI is the most useful immediate investigation; unenhanced CT of the brain is usually nonspecific, and contrast is often contraindicated because of poor renal function.

Endocarditis Most cases of *Aspergillus* endocarditis are prosthetic valve infections resulting from contamination during surgery. Native valve disease is reported, especially as a feature of disseminated infection and in persons using illicit IV drugs. Culture-negative endocarditis with large vegetations is the most common presentation; embolelectomy occasionally reveals the diagnosis.

Cutaneous Aspergillosis Dissemination of *Aspergillus* occasionally results in cutaneous features, usually an erythematous or purplish nontender area that progresses to a necrotic eschar. Direct invasion of the skin occurs in neutropenic patients at the site of IV catheter insertion and in burn patients; such invasion may also follow trauma. Wounds may become infected with *Aspergillus* (especially *A. flavus*) after surgery.

Chronic Pulmonary Aspergillosis The hallmark of chronic cavitary pulmonary aspergillosis (also called semi-invasive aspergillosis, chronic necrotizing aspergillosis, or complex aspergilloma) (Fig. 241-1) is one or more pulmonary cavities expanding over a period of months or years in association with pulmonary symptoms and systemic manifestations such as fatigue and weight loss. (Pulmonary aspergillosis developing over <3 months is better classified as subacute invasive aspergillosis.) Often mistaken initially for tuberculosis, almost all cases occur in patients with prior pulmonary disease (e.g., tuberculosis, atypical mycobacterial infection, sarcoidosis, rheumatoid lung disease, pneumothorax, bullae) or lung surgery. The onset is insidious, and systemic features may be more prominent than pulmonary symptoms. Cavities may have a fluid level or a well-formed fungal ball, but pericavitary infiltrates and multiple cavities—with or without pleural thickening—are typical. An irregular internal cavity surface and thickened cavity walls are indicative of disease activity.

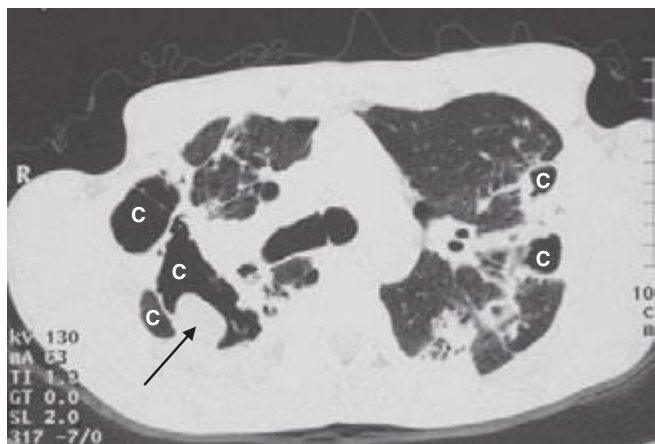


FIGURE 241-1 CT scan image of the chest in a patient with long-standing bilateral chronic cavitary pulmonary aspergillosis. This patient had a history of several bilateral pneumothoraces and had required bilateral pleurodesis in 1990. CT scan then demonstrated multiple bullae, and sputum cultures grew *A. fumigatus*. The patient had initially weakly and later strongly positive serum *Aspergillus* antibody tests (precipitins). This scan (2003) shows a mixture of thick- and thin-walled cavities in both lungs (each marked with C), with a probable fungal ball (black arrow) protruding into the large cavity on the patient's right side (R). There is also considerable pleural thickening bilaterally.

IgG antibodies (usually precipitating) to *Aspergillus* are almost always detectable in blood, and levels fall slowly with successful therapy. Some patients have concurrent infections—even without a fungal ball—with atypical mycobacteria and/or other bacterial pathogens. One or more *Aspergillus* nodules that resemble early lung carcinoma and may cavitate have been recognized. If untreated, chronic pulmonary aspergillosis typically progresses (sometimes relatively rapidly) to unilateral or upper-lobe fibrosis. This end-stage entity is termed *chronic fibrosing pulmonary aspergillosis*.

Aspergilloma Aspergilloma (fungal ball) occurs in up to 20% of residual pulmonary cavities ≥ 2.5 cm in diameter. Signs and symptoms associated with single (simple) aspergillomas are minor, including cough (sometimes productive), hemoptysis, wheezing, and mild fatigue. More significant signs and symptoms are associated with chronic cavitary pulmonary aspergillosis and should be treated as such. The vast majority of fungal balls are caused by *A. fumigatus*, but *A. niger* has been implicated, particularly in diabetic patients; aspergillomas due to *A. niger* can lead to oxalosis with renal dysfunction. The most significant complication of aspergilloma is life-threatening hemoptysis, which may be the presenting manifestation. Some fungal balls resolve spontaneously, but the cavity may still be infected.

Chronic Aspergillus Sinusitis Three entities are subsumed under this broad designation: fungal ball of the sinus, chronic invasive sinusitis, and chronic granulomatous sinusitis. *Fungal ball of the sinus* is limited to the maxillary sinus (except in rare cases involving the sphenoid sinus) and consists of a chronic saprophytic entity in which the sinus cavity is filled with a fungal ball. Maxillary disease is associated with prior upper-jaw root canal work and chronic (bacterial) sinusitis. About 90% of CT scans show focal hyperattenuation related to concretions; on MRI scans, the T2-weighted signal is decreased, whereas it is increased in bacterial sinusitis. Removal of the fungal ball is curative. No tissue invasion is demonstrable histologically or radiologically.

In contrast, *chronic invasive sinusitis* is a slowly destructive process that most commonly affects the ethmoid and sphenoid sinuses but can involve any sinus. Patients are usually but not always immunocompromised to some degree (e.g., as a result of diabetes or HIV infection). Imaging of the cranial sinuses shows opacification of one or more sinuses, local bone destruction, and invasion of local structures. The differential diagnosis is wide, including infections caused by numerous other fungi; sphenoid sinusitis is often caused by bacteria. Apart from a history of chronic nasal discharge and blockage, loss of the sense of smell, and persistent headache, the usual presenting features are related to local involvement of critical structures. The orbital apex syndrome (blindness and proptosis) is characteristic. Facial swelling, cavernous sinus thrombosis, carotid artery occlusion, pituitary fossa, and brain and skull base invasion have been described.

Chronic granulomatous sinusitis due to *Aspergillus* is most commonly seen in the Middle East and India and is often caused by *A. flavus*. It typically presents late, with facial swelling and unilateral proptosis. The prominent granulomatous reaction histologically distinguishes this disease from chronic invasive sinusitis, in which tissue necrosis with a low-grade mixed-cell infiltrate is typical. IgG antibodies to *A. flavus* are usually detectable.

Allergic Bronchopulmonary Aspergillosis In almost all cases, ABPA represents a hypersensitivity reaction to *A. fumigatus*; rare cases are due to other aspergilli and other fungi. ABPA occurs in ~2.5% of patients with asthma who are referred to secondary care and in up to 15% of adults with cystic fibrosis. Episodes of bronchial obstruction with mucous plugs leading to coughing fits, “pneumonia,” consolidation, and breathlessness are typical. Many patients report coughing up thick sputum casts. Eosinophilia commonly develops before systemic glucocorticoids are given. The cardinal diagnostic tests include an elevated serum level of total IgE (usually >1000 IU/mL), a positive skin-prick test in response to *A. fumigatus* extract, or detection of *Aspergillus*-specific IgE and IgG antibodies. The presence of hyperattenuated mucus in airways is highly specific. Central bronchiectasis is characteristic, and some patients develop chronic cavitary pulmonary aspergillosis.

1348 Severe Asthma with Fungal Sensitization Many adults with severe asthma do not fulfill the criteria for ABPA and yet are allergic to fungi. Although *A. fumigatus* is a common allergen, numerous other fungi (e.g., *Cladosporium* and *Alternaria* species) are implicated by skin-prick testing and/or specific IgE radioallergosorbent testing. Serum total IgE concentrations are <1000 IU/mL, and bronchiectasis is moderately common.

Allergic Sinusitis Like the lungs, the sinuses manifest allergic responses to *Aspergillus* and other fungi. The affected patients present with chronic (i.e., perennial) sinusitis that is relatively unresponsive to antibiotics. Many of these patients have nasal polyps, and all have congested nasal mucosae and sinuses full of mucoid material. The histologic hallmarks of allergic fungal sinusitis are local eosinophilia and Charcot-Leyden crystals. Removal of abnormal mucus and polyps, with local and occasionally systemic administration of glucocorticoids, usually leads to resolution. Persistent or recurrent signs and symptoms may require more extensive surgery (ethmoidectomy) and possibly local antifungal therapy. Recurrence is common, often after another bacterial or viral infection.

Superficial Aspergillosis *Aspergillus* can cause keratitis and otitis externa. The former may be difficult to diagnose early enough to save the patient's sight. Treatment requires local surgical debridement as well as intensive topical antifungal therapy. Otitis externa usually resolves with debridement and local application of antifungal agents.

DIAGNOSIS

Several techniques are required to establish the diagnosis of any form of aspergillosis with confidence (Table 241-1). Patients with acute invasive aspergillosis have a relatively heavy load of fungus in the affected organ; thus culture, molecular diagnosis, antigen detection, and histopathology usually confirm the diagnosis. However, the pace of progression leaves only a narrow window for making the diagnosis without losing the patient, and some invasive procedures are not possible because of coagulopathy, respiratory compromise, and other factors. Currently, ~40% of cases of invasive aspergillosis are missed clinically and are diagnosed only at autopsy. Histologic examination of affected tissue reveals either infarction, with invasion of blood vessels by many fungal hyphae, or acute necrosis, with limited inflammation and fewer hyphae. *Aspergillus* hyphae are hyaline, narrow, and septate, with branching at 45°; no yeast forms are present in infected tissue. Hyphae can be seen in cytology or microscopy preparations, which therefore provide a rapid means of presumptive diagnosis.

Culture is important in confirming the diagnosis, given that multiple other (rarer) fungi can mimic *Aspergillus* species histologically. Bacterial agar is less sensitive than fungal media for culture. Thus, if physicians do not request fungal culture, the diagnosis may be missed. Culture may be falsely positive (e.g., in patients whose airways are colonized by *Aspergillus*) or falsely negative. Only 10–30% of patients with invasive aspergillosis have a positive culture at any time. Both antigen detection and real-time polymerase chain reaction (PCR) are faster and much more sensitive than culture of respiratory samples and blood.

The *Aspergillus* antigen test relies on detection of galactomannan release from *Aspergillus* organisms during growth. Positive serum antigen results usually precede clinical or radiologic features by several days. The sensitivity of antigen detection is reduced by antifungal prophylaxis and empirical therapy.

Definitive confirmation of the diagnosis requires (1) a positive culture of a sample taken directly from an ordinarily sterile site (e.g., a brain abscess) or (2) positive results of both histologic testing and culture of a sample taken from an affected organ (e.g., sinuses or skin). Most diagnoses of invasive aspergillosis are inferred from fewer data, including the presence of the *halo sign* on a high-resolution thoracic CT scan, in which a localized ground-glass appearance representing hemorrhagic infarction surrounds a nodule. While a halo sign may be

produced by other fungi, *Aspergillus* species are by far the most common cause. Halo signs are present for ~7 days early in the course of infection in neutropenic patients and are a good prognostic feature, reflecting an early diagnosis. Thick CT sections can give the false appearance of a halo sign, as can other technical factors. Other common radiologic features of invasive pulmonary aspergillosis include nodules and pleural-based infarction or cavitation, with pleural fluid apparent in 10% of patients.

For chronic invasive aspergillosis, *Aspergillus* antibody testing is invaluable although relatively imprecise. Biopsy of new nodules reveals hyphae surrounded by cells of chronic inflammation and sometimes granulomas. Antibody titers fall with successful therapy. Cultures are infrequently positive but are important in checking for azole resistance. Real-time PCR of sputum is often strongly positive. Some patients with chronic pulmonary aspergillosis also have elevated titers of total serum IgE and *Aspergillus*-specific IgE.

ABPA and severe asthma with fungal sensitization are diagnosed serologically with elevated total and specific serum IgE levels and with skin-prick tests. Allergic *Aspergillus* sinusitis is usually diagnosed histologically, although precipitating antibodies in blood also may be useful.

TREATMENT ASPERGILLOSIS

Antifungal drugs active against *Aspergillus* include voriconazole, itraconazole, posaconazole, caspofungin, micafungin, and amphotericin B (AmB). Drug interactions with azoles must be considered before these agents are prescribed. In addition, plasma azole concentrations vary substantially from one patient to another, and many authorities recommend monitoring to ensure that drug concentrations are adequate but not excessive. Initial IV administration is preferred for acute invasive aspergillosis and oral administration for all other disease that requires antifungal therapy. Current recommendations are shown in Table 241-3.

Voriconazole is the preferred agent for invasive aspergillosis; caspofungin, posaconazole, and lipid-associated AmB are second-line agents. AmB is not active against *A. terreus* or *A. nidulans*. An infectious disease consultation is advised for patients with invasive disease, given the complexity of management. Combination therapy (voriconazole plus an echinocandin) for acute invasive aspergillosis may be beneficial for non-neutropenic patients. Immune reconstitution can complicate recovery. The duration of therapy for invasive aspergillosis varies from ~3 months to several years, depending on the patient's immune status and response to therapy. Relapse occurs if the response is suboptimal and immune reconstitution is not complete.

Itraconazole is currently the preferred oral agent for chronic and allergic forms of aspergillosis. Voriconazole or posaconazole can be substituted when failure, emergence of resistance, or adverse events occur. An itraconazole dose of 200 mg twice daily is recommended, with monitoring of drug concentrations in the blood. Chronic cavitary pulmonary aspergillosis probably requires life-long therapy, whereas the duration of treatment for other forms of chronic and allergic aspergillosis requires case-by-case evaluation.



Resistance to one or more azoles, although uncommon, is present in isolates from the environment in many regions, including northern Europe, India, China, and North America. Resistance may be derived from azole fungicide use for crops. In addition, resistance arising from multiple mechanisms may develop during long-term treatment, and a positive culture during antifungal therapy is an indication for susceptibility testing. Combined resistance to itraconazole and voriconazole is the most common type of cross-resistance. Glucocorticoids should be used in chronic cavitary pulmonary aspergillosis only if covered by adequate antifungal therapy.

Surgical treatment is important in several forms of aspergillosis, including fungal ball of the sinus and single aspergillomas, in which surgery is curative; invasive aspergillosis involving bone,

TABLE 241-3 TREATMENT OF ASPERGILLOSIS^a

Indication	Primary Treatment	Evidence Level ^b	Precautions	Secondary Treatment	Comments
Invasive ^c	Voriconazole	AI	Drug interactions (especially with rifampin), renal failure (IV only)	AmB, caspofungin, posaconazole, micafungin	As primary therapy, voriconazole carries 20% more responses than AmB. Consider initial combination therapy with an echinocandin in non-neutropenic patients.
Prophylaxis	Posaconazole, itraconazole solution	AI	Diarrhea and vomiting with itraconazole, vincristine interaction	Micafungin, aerosolized AmB	Some centers monitor plasma levels of itraconazole and posaconazole.
Single aspergilloma	Surgery	BII	Multicavity disease: poor outcome of surgery, medical therapy preferable	Itraconazole, voriconazole, intracavity AmB	Single large cavities with an aspergilloma are best resected.
Chronic pulmonary ^c	Itraconazole, voriconazole	BII	Poor absorption of itraconazole capsules with proton pump inhibitors or H ₂ blockers	Posaconazole, IV AmB, IV micafungin	Resistance may emerge during treatment, especially if plasma drug levels are subtherapeutic.
ABPA/SAFS	Itraconazole	AI	Some glucocorticoid interactions, including with inhaled formulations	Voriconazole, posaconazole	Long-term therapy is helpful in most cases. No evidence indicates whether therapy modifies progression to bronchiectasis/fibrosis.

^aFor information on duration of therapy, see text. ^bEvidence levels are those used in treatment guidelines (TJ Walsh et al: Treatment of aspergillosis: Clinical practice guidelines of the Infectious Diseases Society of America [IDSA]. Clin Infect Dis 46:327, 2008). ^cAn infectious disease consultation is appropriate for these patients.

Note: The oral dose is usually 200 mg bid for voriconazole and itraconazole and 400 mg bid for posaconazole suspension. The IV dose of voriconazole for adults is 6 mg/kg twice at 12-h intervals (loading doses) followed by 4 mg/kg q12h; a larger dose is required for children and teenagers. Plasma monitoring is helpful in optimizing the dosage. Caspofungin is given as a single loading dose of 70 mg and then at 50 mg/d; some authorities use 70 mg/d for patients weighing >80 kg, and lower doses are required with hepatic dysfunction. Micafungin is given as 50 mg/d for prophylaxis and as at least 150 mg/d for treatment; this drug has not yet been approved by the U.S. Food and Drug Administration (FDA) for this indication. AmB deoxycholate is given at a daily dose of 1 mg/kg if tolerated. Several strategies are available for minimizing renal dysfunction. Lipid-associated AmB is given at 3 mg/kg (AmBisome) or 5 mg/kg (Abelcet). Different regimens are available for aerosolized AmB, but none is FDA approved. Other considerations that may alter dose selection or route include age; concomitant medications; renal, hepatic, or intestinal dysfunction; and drug tolerability.

Abbreviations: AmB, amphotericin B; ABPA, allergic bronchopulmonary aspergillosis; SAFS, severe asthma with fungal sensitization.

heart valve, sinuses, and proximal areas of the lung; brain abscess; keratitis; and endophthalmitis. In allergic fungal sinusitis, removal of abnormal mucus and polyps, with local and occasionally systemic glucocorticoid treatment, usually leads to resolution. Persistent or recurrent signs and symptoms may require more extensive surgery (ethmoidectomy) and possibly local antifungal therapy. Surgery is problematic in chronic pulmonary aspergillosis, usually resulting in serious complications. Bronchial artery embolization is preferred for problematic hemoptysis.

PROPHYLAXIS

In situations in which moderate or high risk is predicted (e.g., after induction therapy for acute myeloid leukemia), the need for antifungal prophylaxis for superficial and systemic candidiasis and for invasive aspergillosis is generally accepted. Fluconazole is commonly used in these situations but has no activity against *Aspergillus* species. Itraconazole capsules are ineffective, and itraconazole solution offers only modest efficacy. Posaconazole solution is more effective. Some data support the use of IV micafungin. No prophylactic regimen is completely successful.

OUTCOME

Invasive aspergillosis is curable if immune reconstitution occurs, whereas allergic and chronic forms are not. The mortality rate for invasive aspergillosis is ~50% if the infection is treated but is 100% if the diagnosis is missed. Infection with a voriconazole-resistant strain carries a mortality rate of 88%. Cerebral aspergillosis, *Aspergillus* endocarditis, and bilateral extensive invasive pulmonary aspergillosis have very poor outcomes, as does invasive infection in persons with late-stage AIDS or relapsed uncontrolled leukemia and in recipients of allogeneic hematopoietic stem cell transplants.

The mortality rate for chronic cavitary pulmonary aspergillosis is ~30% 6 months after presentation, falling to ~15% annually thereafter. After 12 months with no antifungal therapy, 70% of patients have deteriorated and 30% are stable (Fig. 241-2). Therapy fails in ~30% of recipients of antifungal therapy and still more often if azole resistance is present.

Both ABPA and SAFS patients respond to antifungal therapy; ~60% respond to itraconazole and ~80% to voriconazole and posaconazole (if tolerated). If the severity of asthma declines, the inhaled glucocorticoid dose can be reduced and oral glucocorticoids can be stopped.

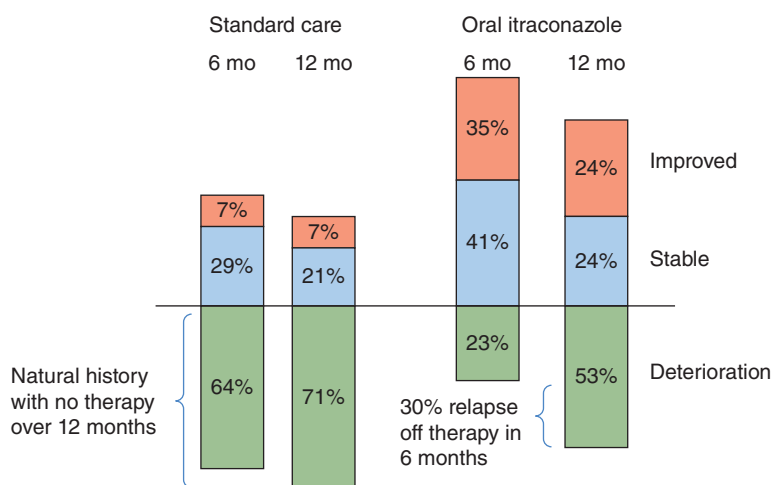


FIGURE 241-2 Comparison of the impact of itraconazole therapy (400 mg/d) and standard care on chronic cavitary pulmonary aspergillosis at 6 and 12 months. (After R Agarwal et al: Itraconazole in chronic cavitary pulmonary aspergillosis: a randomised controlled trial and systematic review of literature. Mycoses 56:559, 2013.)

Mucormycosis represents a group of life-threatening infections caused by fungi of the order Mucorales of the subphylum Mucoromycotina (formerly known as the class Zygomycetes). Infection caused by the Mucorales is most accurately referred to as *mucormycosis*, although the term *zygomycosis* may still be used by some sources. Mucormycosis is highly invasive and relentlessly progressive, resulting in higher rates of morbidity and mortality than many other infections. However, recent studies have suggested that mortality rates from mucormycosis have declined with newer therapies. A high index of suspicion is critical for diagnosis, and early initiation of therapy—often before confirmation of the diagnosis—is necessary to optimize outcomes.

ETIOLOGY



Fungi of the order Mucorales belong to seven medically relevant families (Table 242-1), all of which can cause mucormycosis. Among the Mucorales, *Rhizopus oryzae* (in the family Mucoraceae) is by far the most common cause of infection in the Western Hemisphere. Less frequently isolated species of the Mucoraceae that cause a similar spectrum of infections include *Rhizopus microsporus*, *Rhizomucor pusillus*, *Lichtheimia corymbifera* (formerly *Absidia corymbifera*), *Apophysomyces elegans*, and *Mucor* species (which, despite its name, only rarely causes mucormycosis). Increasing numbers of cases of mucormycosis due to infection with *Cunninghamella* species (family Cunninghamellaceae) have also been reported, particularly in highly immunocompromised patients. Rare case reports have demonstrated the ability of fungi in the remaining families of the Mucorales to cause mucormycosis, although other Mucorales can be the major cause of disease in certain geographic areas (e.g., *A. elegans* in India and *Mucor irregularis* in China).

PATHOGENESIS

The Mucorales are ubiquitous environmental fungi to which humans are constantly exposed. These fungi cause infection primarily in patients with diabetes or defects in phagocytic function (e.g., those associated with neutropenia or glucocorticoid treatment). Patients with elevated levels of free iron, which supports fungal growth in serum and tissues, are likewise at increased risk for mucormycosis. In iron-overloaded patients with end-stage renal failure, treatment with deferoxamine predisposes to the development of rapidly fatal disseminated mucormycosis; this agent, an iron chelator for the human host, serves as a fungal siderophore, directly delivering iron to the Mucorales. Furthermore, patients with diabetic ketoacidosis (DKA) are at high risk of developing rhinocerebral mucormycosis. The acidosis causes dissociation of iron from sequestering proteins in serum,

resulting in enhanced fungal survival and virulence. Nevertheless, the majority of diabetic patients who present with mucormycosis are not acidotic, and, even absent acidosis, hyperglycemia directly contributes to the risk of mucormycosis by at least three likely mechanisms: (1) hyperglycation of iron-sequestering proteins, disrupting normal iron sequestration; (2) upregulation of a mammalian cell receptor (GRP78) that binds to Mucorales, enabling tissue penetration (due to both a direct effect of hyperglycemia and increasing levels of free iron, which independently enhances GRP78 expression); and (3) induction of poorly characterized defects in phagocytic function.

EPIDEMIOLOGY

Mucormycosis typically occurs in patients with diabetes mellitus, solid organ or hematopoietic stem cell transplantation (HSCT), prolonged neutropenia, or malignancy. The majority of diabetic patients are not acidotic on presentation with mucormycosis. Furthermore, patients often have no previously recognized history of diabetes mellitus when they present with mucormycosis. In these instances, presentation for mucormycosis may result in the first clinical recognition of hyperglycemia, which may have been unmasked by recent glucocorticoid use. Thus a high index of suspicion of mucormycosis must be maintained, even in the absence of a known history of diabetes, if hyperglycemia is present. In patients undergoing HSCT, mucormycosis develops at least as commonly during nonneutropenic as during neutropenic periods, probably because of glucocorticoid treatment of graft-versus-host disease. Mucormycosis can occur as isolated cutaneous or subcutaneous infection in immunologically normal individuals after traumatic implantation of soil or vegetation (e.g., due to natural disasters or motor vehicle accidents) or in nosocomial settings via direct access through IV catheters, SC injections, or maceration of the skin by a moist dressing.

Patients receiving antifungal prophylaxis with either itraconazole or voriconazole may be at increased risk of mucormycosis. These patients typically present with disseminated mucormycosis, the most lethal form of disease. Breakthrough mucormycosis also has been described in patients receiving posaconazole or echinocandin prophylaxis.

CLINICAL MANIFESTATIONS

Mucormycosis can be divided into at least six clinical categories based on clinical presentation and the involvement of a particular anatomic site: rhino-orbital-cerebral, pulmonary, cutaneous, gastrointestinal, disseminated, and miscellaneous. These categories of invasive mucormycosis tend to affect patients with specific defects in host defense. For example, patients with DKA typically develop the rhino-orbital-cerebral form and much more rarely develop pulmonary or disseminated disease. In contrast, pulmonary mucormycosis occurs most commonly in leukemic patients who are receiving chemotherapy and in patients undergoing HSCT.

Rhino-Orbital-Cerebral Disease Rhino-orbital-cerebral mucormycosis continues to be the most common form of the disease. Most cases occur in patients with diabetes, although such cases (probably due to glucocorticoid use) are increasingly being described in the transplantation setting, often along with glucocorticoid-induced diabetes mellitus. The initial symptoms of rhino-orbital-cerebral mucormycosis are nonspecific and include eye or facial pain and facial numbness followed by the onset of conjunctival suffusion and blurry vision. Fever may be absent in up to half of cases. White blood cell counts are typically elevated as long as the patient has functioning bone marrow. If untreated, infection usually spreads from the ethmoid sinus to the orbit, resulting in compromise of extraocular muscle function and proptosis, typically with chemosis. Onset of signs and symptoms in the contralateral eye, with resulting bilateral proptosis, chemosis, vision loss, and ophthalmoplegia, is ominous, suggesting the development of cavernous sinus thrombosis.

Upon visual inspection, infected tissue may appear to be normal during the earliest stages of fungal spread, then progressing through an erythematous phase, with or without edema, before the onset of a violaceous appearance and finally the development of a black necrotic eschar. Infection can sometimes extend from the sinuses into the

TABLE 242-1 TAXONOMY OF FUNGI CAUSING MUCORMYCOSIS (SUBPHYLUM MUCOROMYCOTINA, ORDER MUCORALES)

Family	Genus (Species Listed for Some)
Mucoraceae	<i>Rhizopus oryzae</i> <i>Rhizopus microsporus</i> <i>Rhizomucor</i> <i>Mucor</i> <i>Actinomucor</i>
Lichtheimiaceae	<i>Lichtheimia</i> (formerly <i>Mycocladus</i> , formerly <i>Absidia</i>)
Cunninghamellaceae	<i>Cunninghamella</i>
Thamniaceae	<i>Cokeromyces</i>
Mortierellaceae	<i>Mortierella</i>
Saksenaceae	<i>Saksenaea</i> <i>Apophysomyces</i>
Syncephalastraceae	<i>Syncephalastrum</i>

mouth and produce painful necrotic ulcerations of the hard palate, but this is a late finding that suggests extensive, well-established infection.

Pulmonary Disease Pulmonary mucormycosis is the second most common manifestation. Symptoms include dyspnea, cough, and chest pain; fever is often but not invariably present. Angioinvasion results in necrosis, cavitation, and/or hemoptysis. Lobar consolidation, isolated masses, nodular disease, cavities, or wedge-shaped infarcts may be seen on chest radiography. High-resolution chest CT is the best method for determining the extent of pulmonary mucormycosis and may demonstrate evidence of infection before it is seen on chest x-ray. In the setting of cancer, where mucormycosis may be difficult to differentiate from aspergillosis, the presence of ≥ 10 pulmonary nodules, pleural effusion, or concomitant sinusitis makes mucormycosis more likely. It is critical to distinguish mucormycosis from aspergillosis as rapidly as possible because treatments for these infections differ. Indeed, voriconazole—the first-line treatment for aspergillosis—exacerbates mucormycosis in mouse and fly models of infection.

Cutaneous Disease Cutaneous mucormycosis may result from external implantation of the fungus or conversely from hematogenous dissemination. External implantation-related infection has been described in the setting of soil exposure from trauma (e.g., in a motor vehicle accident or natural disaster), penetrating injury with plant material (e.g., a thorn), injections of medications (e.g., insulin), catheter insertion, contamination of surgical dressings, and use of tape to secure endotracheal tubes. Cutaneous disease can be highly invasive, penetrating into muscle, fascia, and even bone. In mucormycosis, necrotizing fasciitis carries a mortality rate approaching 80%. Necrotic cutaneous lesions in the setting of hematogenous dissemination also are associated with an extremely high mortality rate. However, with prompt, aggressive surgical debridement, isolated cutaneous mucormycosis has a favorable prognosis and a low mortality rate.

Gastrointestinal Disease In the past, gastrointestinal mucormycosis occurred primarily in premature neonates in association with disseminated disease and necrotizing enterocolitis. However, there has been a marked increase in case reports describing adults with neutropenia or other immunocompromising conditions. In addition, gastrointestinal disease has been reported as a nosocomial process following administration of medications mixed with contaminated wooden applicator sticks. Nonspecific abdominal pain and distention associated with nausea and vomiting are the most common symptoms. Gastrointestinal bleeding is common, and fungating masses may be seen in the stomach at endoscopy. The disease may progress to visceral perforation, with extremely high mortality rates.

Disseminated and Miscellaneous Forms of Disease Hematogenously disseminated mucormycosis may originate from any primary site of infection. The most common site of dissemination is the brain, but metastatic lesions may also be found in any other organ. The mortality rate associated with dissemination to the brain approaches 100%. Even without central nervous system (CNS) involvement, mortality rates for disseminated mucormycosis exceed 90%. Miscellaneous forms of mucormycosis may affect any body site, including bones, mediastinum, trachea, kidneys, and (in association with dialysis) peritoneum.

DIAGNOSIS

A high index of suspicion is required for diagnosis of mucormycosis. Unfortunately, autopsy series have shown that up to half of cases are diagnosed only post-mortem. Because the Mucorales are environmental isolates, definitive diagnosis requires a positive culture from a sterile site (e.g., a needle aspirate, a tissue biopsy specimen, or pleural fluid) or histopathologic evidence of invasive mucormycosis. A probable diagnosis of mucormycosis can be established by culture

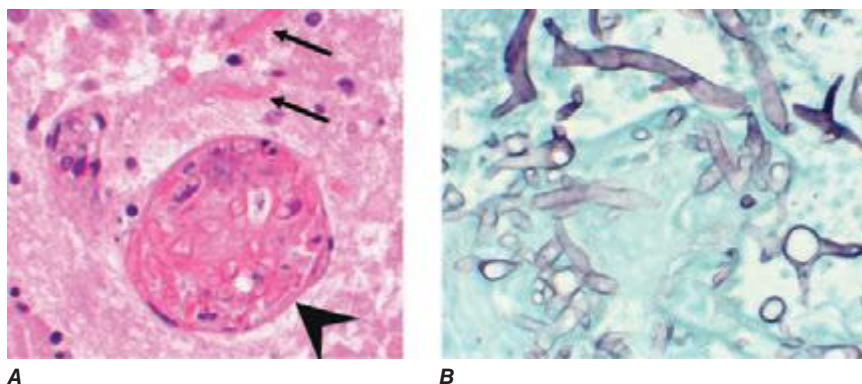


FIGURE 242-1 Histopathology sections of *Rhizopus oryzae* in infected brain. **A.** Broad, ribbon-like, nonseptate hyphae in the parenchyma (arrows) and a thrombosed blood vessel with extensive intravascular hyphae (arrowhead) (hematoxylin and eosin). **B.** Extensive, broad, ribbon-like hyphae invading the parenchyma (Gomori methenamine silver).

from a nonsterile site (e.g., sputum or bronchoalveolar lavage) when a patient has appropriate risk factors as well as clinical and radiographic evidence of disease. However, given the urgency of administering therapy early, the patient should be treated while confirmation of the diagnosis is awaited.

Biopsy with histopathologic examination remains the most sensitive and specific modality for definitive diagnosis (Fig. 242-1). Biopsy reveals characteristic wide (≥ 6 - to $30\text{-}\mu\text{m}$), thick-walled, ribbon-like, aseptate hyphal elements that branch at right angles. Other fungi, including *Aspergillus*, *Fusarium*, and *Scedosporium* species, have septa, are thinner, and branch at acute angles. Because artificial septa may result from folding of tissue during processing (which may also alter the appearance of the angle of branching), the width and the ribbon-like form of the fungus are the most reliable features distinguishing mucormycosis. The Mucorales are visualized most effectively with periodic acid-Schiff or methenamine silver stain or, if the organism burden is high, with hematoxylin and eosin. While histopathology can identify the Mucorales, specific species can be identified only by culture. Polymerase chain reaction (PCR) is being investigated as a diagnostic tool for mucormycosis but is not yet approved by the U.S. Food and Drug Administration (FDA) for this purpose and is not generally available.

Unfortunately, cultures are positive in fewer than half of cases of mucormycosis. Nevertheless, the Mucorales are not fastidious organisms and tend to grow quickly (i.e., within 48 h) on culture media. The likely explanation for the low sensitivity of culture is that the Mucorales form long filamentous structures that are killed by tissue homogenization—the standard method for preparing tissue cultures in the clinical microbiology laboratory. Thus the laboratory should be advised when a diagnosis of mucormycosis is suspected, and the tissue should be cut into sections and placed in the center of culture dishes rather than homogenized. There is also substantial variability among isolates in optimal growth temperature, so growth at both room temperature and 37°C is advisable.

Imaging techniques often yield subtle findings that underestimate the extent of disease. For example, the most common finding on CT or MRI of the head or sinuses of a patient with rhino-orbital mucormycosis is sinusitis that is indistinguishable from bacterial sinusitis. It is also common to detect no abnormalities in sinus bones despite clinical evidence of progressive disease. MRI is more sensitive ($\sim 80\%$) for detecting orbital and CNS disease than is CT. High-risk patients should always undergo endoscopy and/or surgical exploration, with biopsy of the areas of suspected infection. If mucormycosis is suspected, initial empirical therapy with a polyene antifungal agent should be initiated while the diagnosis is being confirmed.

DIFFERENTIAL DIAGNOSIS

Other mold infections, including aspergillosis, scedosporiosis, fusariosis, and infections caused by the dematiaceous fungi (brown-pigmented

1352 soil organisms), can cause clinical syndromes identical to mucormycosis. Histopathologic examination usually allows distinction of the Mucorales from these other organisms, and a positive culture permits definitive species identification. As stated above, it is important to distinguish the Mucorales from these other fungi, as the preferred antifungal treatments differ (i.e., polyenes for the Mucorales vs. expanded-spectrum triazoles for most septate molds). The entomophthoromycoses caused by *Basidiobolus* and *Conidiobolus* also can cause identical clinical syndromes. These fungi may appear similar to the Mucorales on histopathology and can be reliably distinguished from the latter only by culture.

In a patient with sinusitis and proptosis, orbital cellulitis and cavernous sinus thrombosis caused by bacterial pathogens (most commonly *Staphylococcus aureus*, but also streptococcal and gram-negative species) must be excluded. *Klebsiella rhinoscleromatis* is a rare cause of an indolent facial rhinoscleroma syndrome that may appear similar to mucormycosis. Finally, the Tolosa-Hunt syndrome causes painful ophthalmoplegia, ptosis, headache, and cavernous sinus inflammation; biopsies and clinical follow-up may be needed to distinguish the Tolosa-Hunt syndrome from mucormycosis by the lack of progression of the former entity.

TREATMENT MUCORMYCOSIS

GENERAL PRINCIPLES

The successful treatment of mucormycosis requires four steps: (1) early diagnosis; (2) reversal of underlying predisposing risk factors, if possible; (3) surgical debridement; and (4) prompt antifungal therapy. Early diagnosis of mucormycosis is critical, since early initiation of therapy is associated with improved survival rates. It is also crucial to reverse (or prevent) underlying defects in host defense during treatment (e.g., by stopping or reducing the dosage of immunosuppressive medications or by rapidly restoring

euglycemia and normal acid-base status). Finally, iron administration to patients with active mucormycosis should be avoided, as iron exacerbates infection in animal models. Blood transfusion typically results in some liberation of free iron due to hemolysis, so a conservative approach to red blood cell transfusions is advisable.

Blood vessel thrombosis and resulting tissue necrosis during mucormycosis can result in poor penetration of antifungal agents to the site of infection. Therefore, debridement of all necrotic tissues is critical for eradication of disease. Surgery has been found (by logistic regression and in multiple case series) to be an independent variable for favorable outcome in patients with mucormycosis. Limited data from a retrospective study support the use of intraoperative frozen sections to delineate the margins of infected tissues, with sparing of tissues lacking evidence of infection. A multidisciplinary team, including an internist, an infectious disease specialist, and surgical specialists whose expertise is relevant to the sites of infection, is typically required for the management of mucormycosis.

ANTIFUNGAL THERAPY

Primary therapy for mucormycosis should be based on a polyene antifungal agent (Table 242-2), except perhaps for mild localized infection (e.g., isolated suprafacial cutaneous infection) that has been eradicated surgically in an immunocompetent patient. Amphotericin B (AmB) deoxycholate remains the only licensed antifungal agent for the treatment of mucormycosis. However, lipid formulations of AmB are significantly less nephrotoxic, can be administered at higher doses, and are probably more effective than AmB deoxycholate for this purpose. Liposomal amphotericin B (LAmB) is preferred to amphotericin B lipid complex (ABLC) for management of CNS infection on the basis of retrospective survival data and superior brain penetration; there is no clear advantage of either agent for non-CNS infections.

TABLE 242-2 FIRST-LINE ANTIFUNGAL OPTIONS FOR THE TREATMENT OF MUCORMYCOSIS^a

Drug	Recommended Dosage	Advantages and Supporting Studies	Disadvantages
Primary Antifungal Therapy			
AmB deoxycholate	1.0–1.5 mg/kg qd	<ul style="list-style-type: none"> >5 decades of clinical experience Inexpensive Only licensed agent for treatment of mucormycosis 	<ul style="list-style-type: none"> Highly toxic Poor CNS penetration
LAmB	5–10 mg/kg qd	<ul style="list-style-type: none"> Less nephrotoxic than AmB deoxycholate Better CNS penetration than AmB deoxycholate or ABLC Better outcomes than with AmB deoxycholate in murine models and a retrospective clinical review 	<ul style="list-style-type: none"> Expensive
ABLC	5 mg/kg qd	<ul style="list-style-type: none"> Less nephrotoxic than AmB deoxycholate Murine and retrospective clinical data suggest benefit of combination therapy with echinocandins 	<ul style="list-style-type: none"> Expensive Possibly less efficacious than LAmB for CNS infection
Primary Combination Therapy^b			
Caspofungin plus lipid polyene	70-mg IV loading dose, then 50 mg/d for ≥2 weeks 50 mg/m ² IV in children	<ul style="list-style-type: none"> Favorable toxicity profile Synergistic in murine disseminated mucormycosis Retrospective clinical data suggest superior outcomes for rhino-orbital-cerebral mucormycosis. 	<ul style="list-style-type: none"> Very limited clinical data on combination therapy
Micafungin or anidulafungin plus lipid polyene	100 mg/d for ≥2 weeks Micafungin: 4 mg/kg qd in children Micafungin: 10 mg/kg qd in low-birth-weight infants Anidulafungin: 1.5 mg/kg qd in children	<ul style="list-style-type: none"> Favorable toxicity profile Synergistic with LAmB in murine model of disseminated mucormycosis 	<ul style="list-style-type: none"> No clinical data

^aPrimary therapy should generally include a polyene. Non-polyene-based regimens may be appropriate for patients who refuse or are intolerant of polyene therapy or for relatively immunocompetent patients with mild disease (e.g., isolated suprafacial cutaneous infection) that can be surgically eradicated. ^bProspective randomized trials are necessary to confirm the suggested benefit (from animal and small retrospective human studies) of combination therapy for mucormycosis. Dose escalation of any echinocandin is not recommended because of a paradoxical loss of benefit of combination therapy at echinocandin doses of ≥3 mg/kg qd.

Abbreviations: ABLC, AmB lipid complex; AmB, amphotericin B; CNS, central nervous system; LAmB, liposomal AmB.

Source: Modified from B Spellberg et al: Clin Infect Dis 48:1743, 2009.

243 Superficial Mycoses and Less Common Systemic Mycoses

Carol A. Kauffman

The optimal dosages for antifungal treatment of mucormycosis are not known. Starting dosages of 1 mg/kg per day for AmB deoxycholate and 5 mg/kg per day for LAmB and ABLC are commonly given to adults and children. Dose escalation of LAmB to 7.5 or 10 mg/kg per day for CNS mucormycosis may be considered in light of the limited penetration of polyenes into the brain. Because of auto-induction of metabolism, which results in paradoxically lower drug levels, there is no advantage to escalating the LAmB dose above 10 mg/kg per day, and doses of 5 mg/kg per day are probably adequate for non-CNS infections. ABLC dose escalation above 5 mg/kg per day is not advisable given the lack of relevant data and the drug's potential toxicity.

Echinocandin-lipid polyene combinations improved survival rates among mice with disseminated mucormycosis (including CNS disease) and were associated with significantly better outcomes than polyene monotherapy in a small retrospective clinical study involving patients with rhino-orbital-cerebral mucormycosis. Although combination therapy may be considered on the basis of these limited data sets, definitive clinical trials are needed to establish whether it offers any real advantage over monotherapy for mucormycosis. Echinocandins should be administered at standard, FDA-approved doses, since dose escalation has resulted in paradoxical loss of efficacy in preclinical models.

In contrast to deferoxamine, the iron chelator deferasirox is fungicidal against clinical isolates of the Mucorales. In mice with DKA and disseminated mucormycosis, combination deferasirox-LAmB therapy resulted in synergistic improvement of survival rates and reduced the fungal burden in brain. Unfortunately, a randomized, double-blind, phase 2 safety clinical trial of adjunctive therapy with deferasirox (plus LAmB) documented excess mortality in the patients treated with deferasirox. It is noteworthy that the study population included primarily patients with active malignancy, and few patients in the study had diabetes mellitus as their only risk factor. Deferasirox is therefore contraindicated as therapy in patients with active malignancy, but its role in patients who have diabetes mellitus without malignancy (the setting in which its preclinical efficacy was optimal) remains uncertain.

Posaconazole is the only FDA-approved azole with in vitro activity against the Mucorales. However, pharmacokinetic/pharmacodynamic data raise concerns about the reliability with which adequate in vivo levels of orally administered posaconazole are attained. Furthermore, posaconazole is inferior in efficacy to AmB for the treatment of murine mucormycosis and is not superior to placebo for treatment of murine infection with *R. oryzae*. Moreover, posaconazole-polyene combination therapy is not superior to polyene monotherapy for mucormycosis in mice, and no comparative data are available for combination therapy in humans.

The roles of recombinant cytokines and neutrophil transfusions in the primary treatment of mucormycosis are not clear, although it is intuitive that earlier recovery of neutrophil counts should improve survival rates. Limited data indicate that hyperbaric oxygen may be useful in centers with the appropriate technical expertise and facilities.

In general, antifungal therapy for mucormycosis should be continued until resolution of clinical signs and symptoms of infection and resolution of underlying immunosuppression. For patients with mucormycosis who are receiving immunosuppressive medications, secondary antifungal prophylaxis is typically continued for as long as the immunosuppressive regimen is administered.

The role of radiographic follow-up in determining prognosis and therapeutic duration is being studied. Analysis of data from the phase 2 DEFEAT Mucor study indicated that early radiographic progression (within the first 2 weeks) did not predict long-term mortality risk, nor did early radiographic stability/regression predict long-term survival. Therefore, caution should be used in reacting to short-term, serial radiographic results, and greater emphasis should be placed on clinical response, particularly within the first 2–4 weeks after initiation of therapy.

ENDEMIC MYCOSES (DIMORPHIC FUNGI)

Dimorphic fungi exist in discrete environmental niches as molds that produce conidia, which are their infectious form. In tissues and at temperatures of >35°C, the mold converts to the yeast form.

Other endemic mycoses—histoplasmosis, coccidioidomycosis, and blastomycosis—are discussed in Chaps. 236, 237, and 238, respectively.

SPOROTRICHOSIS



Etiologic Agent *Sporothrix schenckii* is a thermally dimorphic fungus that is found worldwide in sphagnum moss, decaying vegetation, and soil.

Epidemiology and Pathogenesis Sporotrichosis most commonly infects persons who participate in outdoor activities such as landscaping, gardening, and tree farming. Infected animals can transmit *S. schenckii* to humans. An outbreak of sporotrichosis in Rio de Janeiro that began in 1998 and that has involved >2000 people has been traced to cats, which are highly susceptible to this infection. Sporotrichosis is primarily a localized infection of skin and subcutaneous tissues that follows traumatic inoculation of conidia. Osteoarticular sporotrichosis is uncommon, occurring most often in middle-aged men who abuse alcohol, and pulmonary sporotrichosis occurs almost exclusively in persons with chronic obstructive pulmonary disease who have inhaled the organism from the environment. Dissemination occurs rarely, almost always in markedly immunocompromised patients, especially those with AIDS.

Clinical Manifestations Days or weeks after inoculation, a papule develops at the site and then usually ulcerates but is not very painful. Similar lesions develop sequentially along the lymphatic channels proximal to the original lesion (Fig. 243-1). Some patients develop a fixed cutaneous lesion that can be verrucous or ulcerative and that remains localized without lymphatic extension. The differential diagnosis of lymphocutaneous sporotrichosis includes nocardiosis, tularemia, nontuberculous mycobacterial infection (especially that due to *Mycobacterium marinum*), and leishmaniasis. Osteoarticular sporotrichosis can present as chronic synovitis or septic arthritis. Pulmonary sporotrichosis must be differentiated from tuberculosis and from other fungal pneumonias.



FIGURE 243-1 Several nodular lesions that developed after a young boy pricked his index finger with a thorn. A culture yielded *S. schenckii*. (Courtesy of Dr. Angela Restrepo.)

1354 Numerous ulcerated skin lesions, with or without spread to visceral organs (including the central nervous system [CNS]), are characteristic of disseminated sporotrichosis.

Diagnosis *S. schenckii* usually grows readily as a mold when material from a cutaneous lesion is incubated at room temperature. Histopathologic examination of biopsy material shows a mixed granulomatous and pyogenic reaction, and tiny oval or cigar-shaped yeasts are sometimes visualized with special stains. Serologic testing is not useful.

Treatment and Prognosis Guidelines for the management of the various forms of sporotrichosis have been published by the Infectious Diseases Society of America (Table 243-1). Itraconazole is the drug of choice for lymphocutaneous sporotrichosis. Fluconazole is less effective; voriconazole and posaconazole have not been used for sporotrichosis. Saturated solution of potassium iodide (SSKI) also is effective for lymphocutaneous infection and costs much less than itraconazole. However, SSKI is poorly tolerated because of adverse reactions, including metallic taste, salivary gland swelling, rash, and fever. Terbinafine appears to be effective but has been used in few patients. Treatment for lymphocutaneous sporotrichosis is continued for 2–4 weeks after all lesions have resolved, usually for a total of 3–6 months. Pulmonary and osteoarticular forms of sporotrichosis are treated with itraconazole for at least 1 year. Severe pulmonary infection and disseminated sporotrichosis, including that involving the CNS, are treated initially with amphotericin B (AmB), which is followed by itraconazole after improvement has been noted. Lifelong suppressive therapy with itraconazole is required for AIDS patients. The success rate for treatment of lymphocutaneous sporotrichosis is 90–100%, but other forms of the disease respond poorly to antifungal therapy.

TABLE 243-1 SUGGESTED TREATMENT FOR ENDEMIC MYCOSES		
Disease	First-Line Therapy	Alternatives/Comments
Sporotrichosis		
Cutaneous, lymphocutaneous	Itraconazole, 200 mg/d until 2–4 weeks after lesions resolve	SSKI, increasing doses ^a Terbinafine, 500 mg bid
Pulmonary, osteoarticular	Itraconazole, 200 mg bid for 12 months	Lipid AmB ^b for severe pulmonary disease until stable; then itraconazole
Disseminated, central nervous system	Lipid AmB ^b for 4–6 weeks	Itraconazole, 200 mg bid after AmB for 12 months Itraconazole maintenance for AIDS patients: 200 mg/d until CD4+ T cell count has been >200/μL for 12 months
Paracoccidioidomycosis		
Chronic (adult form)	Itraconazole, 100–200 mg/d for 6–12 months	TMP-SMX, 160/800 mg bid for 12–36 months
Acute (juvenile form)	AmB ^c until improvement	Itraconazole, 200 mg bid after AmB for 12 months
Penicilliosis		
Mild or moderate	Itraconazole, 200 mg bid for 12 weeks	Itraconazole maintenance for AIDS patients: 200 mg/d until CD4+ T cell count has been >100/μL for 6 months
Severe	AmB ^c until improvement	Itraconazole, 200 mg bid after AmB for 12 weeks Itraconazole maintenance: as for mild or moderate disease

^aThe starting dosage is 5–10 drops tid in water or juice. The dosage is increased weekly by 10 drops per dose, as tolerated, up to 40–50 drops tid. ^bThe dosage of lipid AmB is 3–5 mg/kg daily; the higher dosage should be used when the central nervous system is involved. ^cThe dosage of AmB deoxycholate is 0.6–1.0 mg/kg daily.

Abbreviations: AmB, amphotericin B; SSKI, saturated solution of potassium iodide; TMP-SMX, trimethoprim-sulfamethoxazole.

PARACOCIDIOMYCOSIS



Etiologic Agent, Epidemiology, and Pathogenesis *Paracoccidioides brasiliensis* is a thermally dimorphic fungus that is endemic in humid areas of Central and South America, especially in Brazil. A striking male-to-female ratio varies from 14:1 to as high as 70:1 (in rural Brazil). Most patients are middle-aged or elderly men from rural areas. Paracoccidioidomycosis develops after the inhalation of aerosolized conidia encountered in the environment. For most patients, disease rarely develops at the time of the initial infection but appears years later, presumably after reactivation of a latent infection.

Clinical Manifestations Two major syndromes are associated with paracoccidioidomycosis: the acute or juvenile form and the chronic or adult form. The acute form is uncommon, occurs mostly in persons <30 years old, and manifests as disseminated infection of the reticuloendothelial system. Immunocompromised individuals also manifest this type of rapidly progressive disease. The chronic form of paracoccidioidomycosis accounts for ~90% of cases and predominantly affects older men. The primary manifestation is progressive pulmonary disease, primarily in the lower lobes, with fibrosis. Ulcerative and nodular mucocutaneous lesions in the nares and mouth—another common manifestation of chronic paracoccidioidomycosis—must be differentiated from leishmaniasis (Chap. 251) and squamous cell carcinoma (Chap. 105).

Diagnosis The diagnosis is established by growth of the organism in culture. A presumptive diagnosis can be made by detection of the distinctive thick-walled yeast, with multiple narrow-necked buds attached circumferentially, in purulent material or tissue biopsies.

Treatment and Prognosis Itraconazole is the treatment of choice for paracoccidioidomycosis (Table 243-1). Ketoconazole is also effective but more toxic; voriconazole and posaconazole have been used with success in a few cases. Sulfonamides also are effective and are the least costly agents, but the response is slower and the relapse rate higher. Seriously ill patients should be treated with AmB initially. Patients with paracoccidioidomycosis have an excellent response to therapy, but pulmonary fibrosis is often progressive in those with chronic disease.

PENICILLIOSIS



Etiologic Agent, Epidemiology, and Pathogenesis *Penicillium marneffeii* is a thermally dimorphic fungus that is endemic in the soil in certain areas of Vietnam, Thailand, and several other southeastern Asian countries. The epidemiology of penicilliosis is linked to bamboo rats, which are infected with the fungus but rarely manifest disease. The disease occurs most often among persons living in rural areas in which the rats are found, but there is no evidence for transmission of the infection directly from rats to humans. Infection is rare in immunocompetent hosts, and most cases are reported in persons who have advanced AIDS. Infection results from the inhalation of conidia from the environment. The organism converts to the yeast phase in the lungs and then spreads hematogenously to the reticuloendothelial system.

Clinical Manifestations The clinical manifestations of penicilliosis mimic those of disseminated histoplasmosis and include fever, fatigue, weight loss, dyspnea, diarrhea (in some cases), lymphadenopathy, hepatosplenomegaly, and skin lesions, which appear as papules that often umbilicate and resemble molluscum contagiosum (Chap. 220e).

Diagnosis Penicilliosis is diagnosed by culture of *P. marneffeii* from blood or from biopsy samples of skin, bone marrow, or lymph node. The organism usually grows within 1 week as a mold that produces a distinctive red pigment. Histopathologic examination of tissues and smears of blood or material from skin lesions shows oval or elliptical yeast-like organisms with central septation and can quickly establish a presumptive diagnosis.

Treatment and Prognosis Patients who have severe disease should be treated initially with AmB until their condition improves; therapy can

then be changed to itraconazole (Table 243-1). Patients who have mild symptoms can be treated from the start with itraconazole. For patients with AIDS, suppressive therapy with itraconazole is recommended until immune reconstitution (related to successful therapy for HIV infection with antiretroviral drugs) is evident. Disseminated penicilliosis is usually fatal if not treated. With treatment, the mortality rate is ~10%.

PHAEOHYPHOMYCOSES

In these common soil organisms (also called *dematiaceous* fungi), melanin causes the hyphae and/or conidia to be darkly pigmented. The term *phaeohyphomycosis* is used to describe any infection with a pigmented mold. This definition encompasses two specific syndromes—eumycetoma and chromoblastomycosis—as well as all other types of infections caused by these organisms. It is important to note that eumycetomas can be caused by hyaline molds as well as brown-black molds and that only about half of all mycetomas are due to fungi. Actinomycetes cause the remainder (Chap. 199). Most of the involved fungi cause localized subcutaneous infections after direct inoculation, but disseminated infection and serious focal visceral infections also occur, especially in immunocompromised patients.

Etiologic Agents A large number of pigmented molds can cause human infection. All are found in the soil or on plants, and some cause economically important plant diseases. The most common cause of eumycetoma is *Madurella* species. *Fonsecaea* and *Cladophialophora* species are responsible for most cases of chromoblastomycosis. Disseminated infection and focal visceral infections are caused by a variety of dematiaceous fungi; *Alternaria*, *Exophiala*, *Curvularia*, and *Wangiella* species are among the more common molds reported to cause human infection. Recently, *Exserohilum* species have caused a large outbreak of severe, sometimes fatal CNS and osteoarticular infections following the injection of methylprednisolone contaminated with this fungus.

Epidemiology and Pathogenesis Eumycetoma and chromoblastomycosis are acquired by inoculation through the skin. These two syndromes are seen almost entirely in tropical and subtropical areas and occur mostly in rural laborers who are frequently exposed to the organisms. Other infections with dematiaceous molds are acquired by inhalation, by traumatic inoculation into the eye or through the skin, or by injection of contaminated medication. Melanin is a virulence factor for all the pigmented molds. Several organisms, specifically *Cladophialophora bantiana* and *Rhinocladiella mackenziei*, are neurotropic and likely to cause CNS infection. In an immunocompromised patient or when a pigmented mold is injected directly into a deep structure, these organisms become opportunists, invading blood vessels and mimicking better-known opportunistic infections, such as aspergillosis.

Clinical Manifestations Eumycetoma is a chronic subcutaneous and cutaneous infection that usually occurs on the lower extremities and that is characterized by swelling, development of sinus tracts, and the appearance of grains that are actually colonies of fungi discharged from the sinus tract. As the infection progresses, adjacent fascia and bony structures become involved. The disease is indolent and disfiguring, progressing slowly over years. Complications include fractures of infected bone and bacterial superinfection.

Chromoblastomycosis is an indolent subcutaneous infection characterized by nodular, verrucous, or plaque-like painless lesions that occur predominantly on the lower extremities and grow slowly over months to years. There is hardly ever extension to adjacent structures, as is seen with eumycetoma. Long-term consequences include bacterial superinfection, chronic lymphedema, and (rarely) the development of squamous cell carcinoma.

Dematiaceous molds are the most common cause of allergic fungal sinusitis and a less common cause of invasive fungal sinusitis. Keratitis occurs with traumatic corneal inoculation. Even in many immunocompromised patients, inoculation through the skin generally produces localized cyst-like, nodular lesions at the entry site.

However, other immunocompromised patients develop pneumonia, brain abscess, or disseminated infection. Epidural injection of *Exserohilum*-contaminated steroids has led to meningitis, basilar stroke, epidural abscess or phlegmon, vertebral osteomyelitis, and arachnoiditis.

Diagnosis The specific diagnosis of infection with a pigmented mold is established by growth of the organism in culture. However, in eumycetoma, a tentative clinical diagnosis can be made when a patient presents with a lesion characterized by swelling, sinus tracts, and grains. Histopathologic examination and culture are necessary to confirm that the etiologic agent is a mold and not an actinomycete. In chromoblastomycosis, the diagnosis rests on the histologic demonstration of sclerotic bodies (dark brown, thick-walled, septate fungal forms that resemble large yeasts) in the tissues; culture establishes which pigmented mold is causing the infection. For other infections, growth of the organism is essential to differentiate infection with a hyaline mold (e.g., *Aspergillus* or *Fusarium*) from that due to a pigmented mold. No serologic assays for pigmented molds are available. Polymerase chain reaction (PCR) assays are increasingly used in the diagnosis of infection due to pigmented molds but are available only through fungal reference laboratories.

Treatment and Prognosis Treatment of eumycetoma and chromoblastomycosis involves both surgical extirpation of the lesion and use of antifungal agents. Surgical removal of the lesions of both eumycetoma and chromoblastomycosis is most effective if performed before extensive spread has occurred. In chromoblastomycosis, cryosurgery and laser therapy have been used with variable success. The antifungal agents of choice are itraconazole, voriconazole, and posaconazole. The most experience has accrued with itraconazole; less experience has been gained with the newer azoles, which are active in vitro and have been reported to be effective in a few patients. Flucytosine and terbinafine also have been used to treat chromoblastomycosis. Chromoblastomycosis and eumycetoma are chronic indolent infections that are difficult to cure but are not life-threatening.

Disseminated and focal visceral infections are treated with the appropriate antifungal agent; the choice of agent is based on the location and extent of the infection, in vitro testing, and clinical experience with the specific infecting organism. AmB is not effective against many of these organisms but has been used successfully against others. The most experience has accrued with itraconazole in the treatment of localized infections. Voriconazole is increasingly used when infections are disseminated or involve the CNS because this drug reaches adequate concentrations within the CNS and because both IV and well-absorbed oral formulations are available. The role of posaconazole has not been established but will likely expand. Disseminated and focal visceral infections, especially those involving the CNS, are associated with high mortality rates.

OPPORTUNISTIC FUNGAL INFECTIONS

Two genera of hyaline (nonpigmented) molds, *Fusarium* and *Scedosporium*, and one yeast-like genus, *Trichosporon*, have become prominent pathogens among immunocompromised patients. Infections caused by *Fusarium* and *Scedosporium* species overlap with invasive aspergillosis in their clinical manifestations, and, when seen in tissues, these organisms appear similar to *Aspergillus*. In the immunocompetent host, these fungi cause localized infections of skin, skin structures, and subcutaneous tissues, but their role as causes of infection in immunocompromised patients will be emphasized in this section.

FUSARIOSIS



Etiologic Agent, Epidemiology, and Pathogenesis *Fusarium* species, which are found worldwide in soil and on plants, have emerged as major opportunists in markedly immunocompromised patients. Most human infections follow inhalation of conidia, but ingestion and direct inoculation also can lead to disease. An outbreak of severe *Fusarium* keratitis among soft contact lens wearers was traced back to a particular brand of contact lens solution and

1356 individual contact lens cases that had been contaminated. Disseminated infection is reported most often in patients who have a hematologic malignancy, are neutropenic, have received a stem cell or solid organ transplant, or have a severe burn.

Clinical Manifestations In immunocompetent persons, *Fusarium* species cause localized infections of various organs. These organisms commonly cause fungal keratitis, which can extend into the anterior chamber of the eye; cause loss of vision; and require corneal transplantation. Onychomycosis due to *Fusarium* species, while basically an annoyance in immunocompetent patients, is a source of subsequent hematogenous dissemination and should be aggressively sought and treated in neutropenic patients. In profoundly immunocompromised patients, fusariosis is angioinvasive, and clinical manifestations mimic those of aspergillosis. Pulmonary infection is characterized by multiple nodular lesions. Sinus infection is likely to lead to invasion of adjacent structures. Disseminated fusariosis occurs primarily in neutropenic patients with hematologic malignancies and in allogeneic stem cell transplant recipients, especially those with graft-versus-host disease. Disseminated fusariosis differs from disseminated aspergillosis in that skin lesions are extremely common with fusariosis; the lesions are nodular or necrotic, are usually painful, and appear over time in different locations (Fig. 243-2).

Diagnosis The diagnostic approach usually includes both documentation of the growth of *Fusarium* species from involved tissue and demonstration of invasion by histopathologic techniques that show septate hyphae in tissues. The organism is difficult to differentiate from *Aspergillus* species in tissues; thus, identification with culture is imperative. An extremely helpful diagnostic clue is growth in blood cultures, which are positive in as many as 50% of patients with disseminated fusariosis. There are no serologic assays for *Fusarium*. PCR techniques have proved useful but are available only through fungal reference laboratories.

Treatment and Prognosis *Fusarium* species are resistant to many antifungal agents. A lipid formulation of AmB (at least 5 mg/kg daily), voriconazole (200–400 mg twice daily), or posaconazole (300 mg daily) is recommended. Many physicians use both a lipid formulation of AmB and either voriconazole or posaconazole because susceptibility information is not available when therapy must be initiated. Serum drug levels should be monitored with either azole to ensure that absorption is adequate and with voriconazole to avoid toxicity. Mortality rates for disseminated fusariosis have been as high as 85%. With the improved antifungal therapy now available, mortality rates

have fallen to ~50%. However, if neutropenia persists, the mortality rate approaches 100%.

SCEDOSPORIOSIS

Etiologic Agent The genus *Scedosporium* includes several pathogens. The major causes of human infections are *Scedosporium apiospermum*, which in its sexual state is termed *Pseudallescheria boydii*, and *S. prolificans*. The *S. apiospermum* complex encompasses several species but will be referred to here simply as *S. apiospermum*.

Epidemiology and Pathogenesis *S. apiospermum* is found worldwide in temperate climates in tidal flats, swamps, ponds, manure, and soil. This organism is known as a cause of pneumonia, disseminated infection, and brain abscess in near-drowning victims. *S. prolificans* is also found in soil but is more geographically restricted. Infection occurs predominantly through inhalation of conidia, but direct inoculation through the skin or into the eye also can occur.

Clinical Manifestations Among immunocompetent persons, *Scedosporium* species are a prominent cause of eumycetoma. Keratitis as a result of accidental corneal inoculation is a sight-threatening infection. In patients who have hematologic malignancies (especially acute leukemia with neutropenia), recipients of solid organ or stem cell transplants, and patients receiving glucocorticoids, *Scedosporium* species are angioinvasive, causing pneumonia and widespread dissemination with abscesses. Pulmonary infection mimics aspergillosis; nodules, cavities, and lobar infiltrates are common. Disseminated infection involves the skin, heart, brain, and many other organs. Skin lesions are not as common or as painful as those of fusariosis.

Diagnosis Diagnosis depends on the growth of *Scedosporium* species from involved tissue and the demonstration of invasion by histopathologic techniques that show septate hyphae in tissues. Culture evidence is essential because *Scedosporium* species are difficult to differentiate from *Aspergillus* in tissues. Demonstration of tissue invasion is essential because these ubiquitous environmental molds can be mere contaminants or colonizers. *S. prolificans* can grow in blood cultures, but *S. apiospermum* usually does not. There are no serologic assays for *Scedosporium*. PCR techniques have proved useful but are available only through fungal reference laboratories.

Treatment and Prognosis *Scedosporium* species are resistant to AmB, echinocandins, and some azoles. Voriconazole is the agent of choice for *S. apiospermum*, and posaconazole also has been used for this infection. *S. prolificans* is resistant in vitro to almost every available antifungal agent; the addition of agents such as terbinafine to a voriconazole regimen has been attempted because in vitro data suggest possible synergy against some strains of *S. prolificans*. Mortality rates for invasive *S. apiospermum* infection are ~50%, but those for invasive *S. prolificans* infection remain as high as 85–100%.

TRICHOSPORONOSIS

Etiologic Agent The genus *Trichosporon* contains many species, some of which cause localized infection of hair and nails. The major pathogen responsible for invasive infection is *Trichosporon asahii*. *Trichosporon* species grow as yeast-like colonies in vitro; in vivo, however, hyphae, pseudohyphae, and arthroconidia can also be seen.

Epidemiology and Pathogenesis These yeasts are commonly found in soil, sewage, and water and in rare instances can colonize human skin and the human gastrointestinal tract. Most infections follow fungal inhalation or entry via central venous catheters. Systemic infection occurs almost exclusively in immunocompromised hosts, including those who have hematologic malignancies, are neutropenic, have received a solid organ transplant, or are receiving glucocorticoids.

Clinical Manifestations Disseminated trichosporonosis resembles invasive candidiasis, and fungemia is often the initial manifestation of infection. Pneumonia, skin lesions, and sepsis syndrome are common. The skin lesions begin as papules or nodules surrounded by erythema and progress to central necrosis. A chronic form of infection mimics hepatosplenic candidiasis (chronic disseminated candidiasis).



FIGURE 243-2 Painful necrotic foot lesion that developed over a week in a woman who had acute leukemia and who had been neutropenic for 2 months. *Fusarium* species were grown from a punch biopsy. (Courtesy of Dr. Nessorine Ktaich.)

Diagnosis The diagnosis of systemic *Trichosporon* infection is established by growth of the organism from involved tissues or from blood. Histopathologic examination of a skin lesion showing a mixture of yeast forms, arthroconidia, and hyphae can lead to an early presumptive diagnosis of trichosporonosis. The serum cryptococcal antigen latex agglutination test may be positive in patients with disseminated trichosporonosis because *T. asahii* and *Cryptococcus neoformans* share polysaccharide antigens.

Treatment and Prognosis Rates of response to AmB have been disappointing, and many *Trichosporon* isolates are resistant in vitro. Voriconazole appears to be the antifungal agent of choice and is used at a dosage of 200–400 mg twice daily. The mortality rates for disseminated *Trichosporon* infection have been as high as 70% but are decreasing with the use of newer azoles, such as voriconazole; however, patients who remain neutropenic are likely to succumb to this infection.

SUPERFICIAL CUTANEOUS INFECTIONS

Fungal infections of the skin and skin structures are caused by molds and yeasts that do not invade deeper tissues but rather cause disease merely by inhabiting the superficial layers of skin, hair follicles, and nails. These agents are the most common cause of fungal infections of humans but only rarely cause serious infections.

YEAST INFECTIONS

Etiologic Agents The lipophilic yeast *Malassezia* is dimorphic in that it lives on the skin in the yeast phase but transforms to the mold phase as it causes disease. Most species require exogenous lipids for growth.

Epidemiology and Pathogenesis *Malassezia* species are part of the indigenous human flora found in the stratum corneum of the back, chest, scalp, and face—areas rich in sebaceous glands. Disease is more common in humid areas. The organisms do not invade below the stratum corneum and generally elicit little if any inflammatory response.

Clinical Manifestations *Malassezia* species cause tinea versicolor (also called *pityriasis versicolor*), folliculitis, and seborrheic dermatitis. Tinea versicolor presents as flat round scaly patches of hypo- or hyperpigmented skin on the neck, chest, or upper arms. The lesions are usually asymptomatic but can be pruritic. They can be mistaken for vitiligo, but the latter is not scaly. Folliculitis occurs over the back and chest and mimics bacterial folliculitis. Seborrheic dermatitis manifests as erythematous pruritic scaly lesions in the eyebrows, moustache, nasolabial folds, and scalp. The scalp lesions are termed *cradle cap* in babies and *dandruff* in adults. Seborrheic dermatitis can be severe in patients with advanced AIDS. Fungemia and disseminated infection occur rarely with *Malassezia* species—almost always in premature neonates receiving parenteral lipid preparations through a central venous catheter.

Diagnosis *Malassezia* infections are diagnosed clinically in most cases. If scrapings are collected on a microscope slide on which a drop of potassium hydroxide has been placed, a mixture of budding yeasts and short septate hyphae is seen. In order to culture *M. furfur* from those patients in whom disseminated infection is suspected, sterile olive oil must be added to the medium.

Treatment and Prognosis Topical creams and lotions, including selenium sulfide shampoo, ketoconazole shampoo or cream, terbinafine cream, and ciclopirox cream, are effective in treating *Malassezia* infections and are usually given for 2 weeks. Mild topical steroid creams are sometimes used to treat seborrheic dermatitis. For extensive disease, itraconazole (200 mg/d) or fluconazole (200 mg/d) can be used for 5–7 days. The rare cases of fungemia caused by *Malassezia* species are treated with AmB or fluconazole, prompt removal of the catheter, and discontinuance of parenteral lipid infusions. *Malassezia* skin infections are benign and self-limited, although recurrences are the rule. The outcome of systemic infection depends on the host's underlying conditions, but most infected infants do well.

DERMATOPHYTE (MOLD) INFECTIONS

Etiologic Agents The molds that cause skin infections in humans include the genera *Trichophyton*, *Microsporum*, and *Epidermophyton*.

These organisms, which are not components of the normal skin flora, can live within the keratinized structures of the skin—hence the term *dermatophytes*.



Epidemiology and Pathogenesis Dermatophytes occur worldwide, and infections with these organisms are extremely common. Some organisms cause disease only in humans and can be transmitted by person-to-person contact and by fomites, such as hairbrushes or wet floors, that have been contaminated by infected individuals. Several species cause infections in cats and dogs and can readily be transmitted from these animals to humans. Finally, some dermatophytes are spread from contact with soil. The characteristic ring shape of cutaneous lesions is the result of the organisms' outward growth in a centrifugal pattern in the stratum corneum. Fungal invasion of the nail usually occurs through the lateral or superficial nail plates and then spreads throughout the nail; when hair shafts are invaded, the organisms can be found either within the shaft or surrounding it. Symptoms are caused by the inflammatory reaction elicited by fungal antigens and not by tissue invasion. Dermatophyte infections occur more commonly in male than in female patients, and progesterone has been shown to inhibit dermatophyte growth.

Clinical Manifestations Dermatophyte infection of the skin is often called *ringworm*. This term is confusing because worms are not involved. *Tinea*, the Latin word for *worm*, describes the serpentine nature of the skin lesions and is a less confusing designation that is used in conjunction with the name of the body part affected—e.g., tinea capitis (head), tinea pedis (feet), tinea corporis (body), tinea cruris (crotch), and tinea unguium (nails, although infection at this site is more often termed *onychomycosis*).

Tinea capitis occurs most commonly in children 3–7 years old. Children with tinea capitis usually present with well-demarcated scaly patches in which hair shafts are broken off right above the skin; alopecia can result. Tinea corporis is manifested by well-demarcated, annular, pruritic, scaly lesions that undergo central clearing. Usually one or several small lesions are present. In some cases, tinea corporis can involve much of the trunk or manifest as folliculitis with pustule formation. The rash should be differentiated from contact dermatitis, eczema, and psoriasis. Tinea cruris is seen almost exclusively in men. The perineal rash is erythematous and pustular, has a discrete scaly border, is without satellite lesions, and is usually pruritic. The rash must be differentiated from intertriginous candidiasis, erythrasma, and psoriasis.

Tinea pedis also is more common among men than among women. It usually starts in the web spaces of the toes; peeling, maceration, and pruritus are followed by development of a scaly pruritic rash along the lateral and plantar surfaces of the feet. Hyperkeratosis of the soles of the feet often ensues. Tinea pedis has been implicated in lower-extremity cellulitis, as streptococci and staphylococci can gain entrance to the tissues through fissures between the toes. Onychomycosis affects toenails more often than fingernails and is most common among persons who have tinea pedis. The nail becomes thickened and discolored and may crumble; onycholysis almost always occurs. Onychomycosis is more common in older adults and in persons with vascular disease, diabetes mellitus, and trauma to the nails. Fungal infection must be differentiated from psoriasis, which can mimic onychomycosis but usually has associated skin lesions.

Diagnosis Many dermatophyte infections are diagnosed by their clinical appearance. If the diagnosis is in doubt, as is often the case in children with tinea capitis, scrapings should be taken from the edge of a lesion with a scalpel blade, transferred to a slide to which a drop of potassium hydroxide is added, and examined under a microscope for the presence of hyphae. Cultures are indicated if an outbreak is suspected or the patient does not respond to therapy. Culture of the nail is especially useful as an aid to decisions about both diagnosis and treatment.

Treatment and Prognosis Dermatophyte infections usually respond to topical therapy. Lotions or sprays are easier than creams to apply to large or hairy areas. Particularly for tinea cruris, the affected

TABLE 243-2 SUGGESTED ORAL TREATMENT FOR EXTENSIVE TINEA INFECTIONS AND ONYCHOMYCOSIS

Antifungal Agent	Suggested Dosage	Comments
Extensive Tinea Skin Infection		
Terbinafine	250 mg/d for 1–2 weeks	Adverse reactions minimal with short treatment period
Itraconazole ^a	200 mg/d for 1–2 weeks	Adverse reactions minimal with short treatment period except for drug interactions
Onychomycosis		
Terbinafine	250 mg/d for 3 months	Slightly superior to itraconazole; monitor for hepatotoxicity
Itraconazole ^a	200 mg/d for 3 months or 200 mg bid for 1 week each month for 3 months	Drug interactions frequent; monitor for hepatotoxicity; rarely causes hypokalemia, hypertension, edema; use with caution in patients with congestive heart failure

^aItraconazole capsules require food and gastric acid for absorption, whereas itraconazole solution is taken on an empty stomach.

area should be kept as dry as possible. When patients have extensive skin lesions, oral itraconazole or terbinafine can hasten resolution (Table 243-2). Terbinafine interacts with fewer drugs than itraconazole and is generally the first-line agent. Onychomycosis does not respond to topical therapy, although ciclopirox nail lacquer applied daily for a year is occasionally beneficial. Itraconazole and terbinafine both accumulate in the nail plate and can be used to treat onychomycosis (Table 243-2). These agents are more effective and better tolerated than griseofulvin and ketoconazole. The major decision to be made with regard to therapy is whether the extent of nail involvement justifies the use of systemic antifungal agents that have adverse effects, may interact with other drugs, and are costly. Treating for cosmetic reasons alone is discouraged. Relapses of tinea cruris and tinea pedis are common and should be treated early with topical creams to avoid development of more extensive disease. Relapses of onychomycosis follow treatment in 25–30% of cases.

immunosuppression. These cases were subsequently recognized as the first cases of what came to be known as the acquired immunodeficiency syndrome (AIDS) (Chap. 226). The incidence of PCP increased dramatically as the AIDS epidemic grew: without chemoprophylaxis or antiretroviral therapy (ART), 80–90% of patients with HIV/AIDS in North America and Western Europe ultimately develop one or more episodes of PCP. While its incidence declined with the introduction of anti-*Pneumocystis* prophylaxis and combination ART, PCP has continued to be a leading cause of AIDS-associated morbidity in the United States and Western Europe, particularly in individuals who do not know they are infected with HIV until they are profoundly immunosuppressed and in HIV-infected patients who are not receiving ART or PCP prophylaxis.

PCP also develops in HIV-uninfected patients who are immunocompromised secondary to hematologic or malignant neoplasms, stem cell or solid organ transplantation, and immunosuppressive medications. The incidence of PCP depends on the degree of immunosuppression. PCP is increasingly reported among individuals receiving tumor necrosis factor α inhibitors and antilymphocyte monoclonal antibodies for rheumatologic or other diseases. While clinical PCP in immunocompetent hosts has not been clearly documented, studies have shown that *Pneumocystis* organisms can colonize the airways of children and adults who are not overtly immunocompromised. The relevance of these organisms to acute or chronic syndromes, such as chronic obstructive pulmonary disease (COPD), in immunocompetent patients is being investigated.



In some developing countries, the incidence of PCP among HIV-infected individuals has been found to be lower than that in industrialized countries. This lower incidence may be due to competing mortality from infectious diseases such as tuberculosis and bacterial pneumonia, which typically occur before patients become immunosuppressed enough to develop PCP. Geographic variations in *Pneumocystis* exposure and underdiagnosis also may explain the apparent lower frequency of PCP in some countries.

PATHOGENESIS AND PATHOLOGY

Life Cycle and Transmission The life cycle of *Pneumocystis* involves both sexual and asexual reproduction, and the organism exists as a trophic form, a cyst, and a precyst at various points. Serologic and molecular studies have demonstrated that most humans are exposed to *Pneumocystis* early in life. It was historically thought that *Pneumocystis* developed from reactivation of latent infection, but *de novo* infections from environmental sources and person-to-person transmission occur as well. Outbreaks of PCP suggest that nosocomial transmission can take place, and studies with rodents show that immunocompetent animals can serve as reservoirs for transmission of *P. carinii* (the infecting species in rodents) to immunocompetent and immunosuppressed animals. However, *Pneumocystis* organisms are species specific, and thus humans are infected only by other humans who transmit *P. jirovecii*; humans cannot be infected with animal species of *Pneumocystis* such as *P. murina* (mice) or *P. oryctolagi* (rabbits). The utility of respiratory isolation in preventing transmission from patients with PCP to other immunosuppressed individuals has been debated; no clear evidence exists, although it seems prudent to isolate patients with active PCP from other immunosuppressed patients.

Role of Immunity Defects in cellular and/or humoral immunity predispose to development of PCP. CD4+ T cells are critical in host defense against *Pneumocystis*. For HIV-infected patients, the incidence is inversely related to the CD4+ T cell count: at least 80% of cases occur at counts of <200 cells/ μ L, and most of these cases develop at counts of <100 cells/ μ L. CD4+ T cell counts are less specific and thus less useful in predicting the risk of PCP in HIV-uninfected, immunosuppressed patients.

Lung Pathology *Pneumocystis* has a unique tropism for the lung. It is presumably inhaled into the alveolar space. Clinically apparent pneumonia occurs only if an individual is immunocompromised. *Pneumocystis* proliferates in the lung, provoking a mononuclear cell

244 Pneumocystis Infections

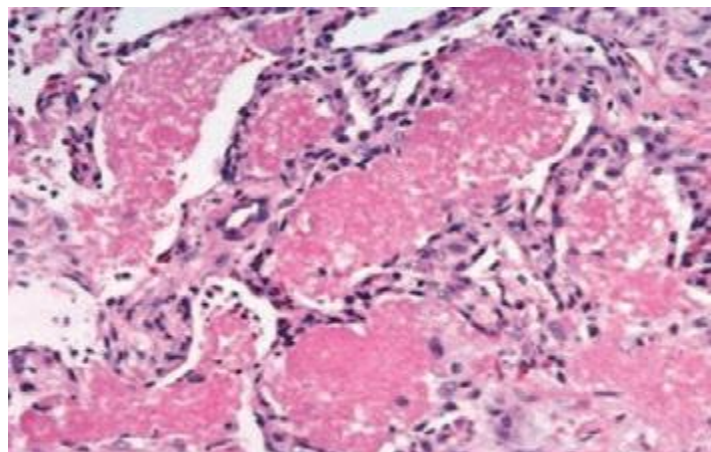
Henry Masur, Alison Morris

DEFINITION AND DESCRIPTION

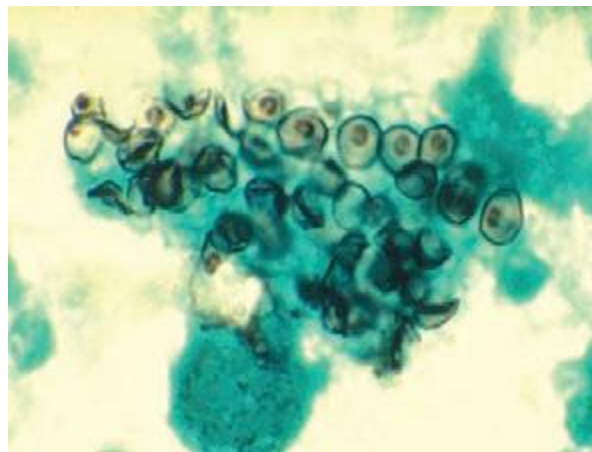
Pneumocystis is an opportunistic pathogen that is an important cause of pneumonia in immunocompromised hosts, particularly those with HIV infection (Chap. 226), and in individuals with organ transplants, those with hematologic malignancies, and those receiving immunosuppressive therapy. The organism was discovered in rodents in 1906 and was initially believed to be a protozoan. Because *Pneumocystis* cannot be cultured, our understanding of its biology has been limited, but molecular techniques have demonstrated that the organism is actually a fungus. Formerly known as *Pneumocystis carinii*, the species infecting humans has been renamed *Pneumocystis jirovecii*.

EPIDEMIOLOGY

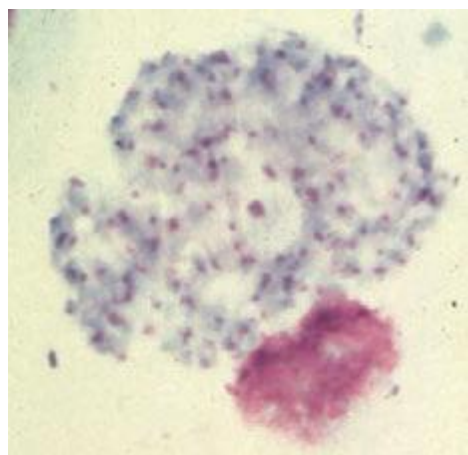
P. jirovecii pneumonia (PCP) came to medical attention when cases were reported in malnourished orphans in Europe during World War II. The disease was later recognized in other immunosuppressed populations but was rare in the era before HIV/AIDS and before intensive immunosuppressive therapy for organ transplantation and autoimmune disorders. In 1981, PCP was first reported in men who had sex with men and in IV drug users who had no obvious cause of



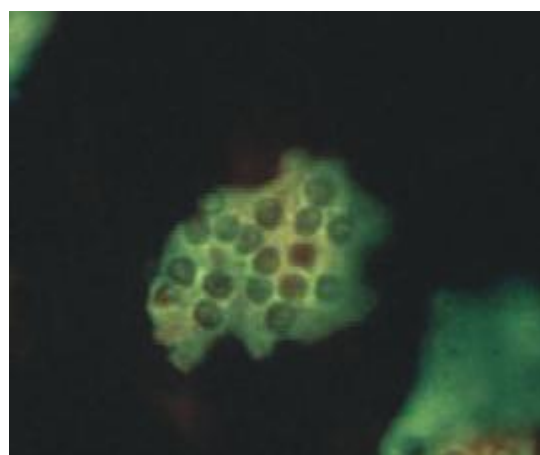
A



B



C



D

FIGURE 244-1 Direct microscopy of *Pneumocystis pneumonia*. **A.** Transbronchial lung biopsy stained with hematoxylin and eosin shows eosinophilic alveolar filling. **B.** Methenamine silver–stained bronchoalveolar lavage (BAL) fluid. **C.** Giemsa-stained BAL fluid. **D.** Immunofluorescent stain of BAL fluid.

response. The alveoli become filled with proteinaceous material, and alveolar damage results in increased alveolar-capillary injury and surfactant abnormalities. Stained lung sections typically show foamy, vacuolated alveolar exudates composed largely of viable and nonviable organisms (Fig. 244-1A). Interstitial edema and fibrosis may develop, and organisms can be seen in the alveolar space with silver or other stains. Moreover, the organisms can be seen when tissue is subjected to colorimetric or immunofluorescent staining (Fig. 244-1B–D).

CLINICAL FEATURES

Clinical Presentation PCP presents as acute or subacute pneumonia that may initially be characterized by a vague sense of dyspnea alone but that subsequently manifests as fever and nonproductive cough with progressive shortness of breath ultimately resulting in respiratory failure and death. Extrapulmonary manifestations of PCP are rare but can include involvement of almost any organ, most notably lymph nodes, spleen, and liver.

Physical Examination The physical examination findings in PCP are nonspecific. Patients have decreased oxygen saturation—at rest or with exertion—that, without treatment, progresses to severe hypoxemia. Patients may initially have a normal chest examination and no adventitious sounds but later, without treatment, develop diffuse rales and signs of consolidation. Oral thrush in a patient with HIV infection indicates an increased risk for PCP.

Laboratory Findings The results of routine laboratory tests are nonspecific in PCP. Serum levels of lactate dehydrogenase (LDH) are often elevated due to pulmonary damage; however, a normal LDH level does not rule out PCP, nor is an elevated LDH value specific for PCP. The peripheral white blood cell count may be elevated, but the increase is usually modest. Hepatic and renal function are typically normal.

Radiographic Findings Although the initial chest radiograph may be normal when patients have mild symptoms, the classic radiographic appearance of PCP consists of diffuse bilateral interstitial infiltrates that are perihilar and symmetric (Fig. 244-2A)—yet another finding that is not specific for PCP. The interstitial infiltrates can progress to alveolar filling (Fig. 244-2B). High-resolution chest CT shows diffuse ground-glass opacities in virtually all patients with PCP (Fig. 244-2C). A normal chest CT essentially rules out the diagnosis of PCP. Cysts and pneumothoraces are common chest radiographic findings (Fig. 244-2D). A wide variety of atypical radiographic findings have been described, including asymmetric patterns, upper lobe infiltrates, mediastinal adenopathy, nodules, cavities, and effusions.

DIAGNOSIS

The optimal sample for diagnostic examination depends on how ill the patient is and what resources are available. Before the 1990s, diagnoses of PCP were usually established by open lung biopsy; later, transbronchial lung biopsy was employed. Hematoxylin and eosin staining of pulmonary

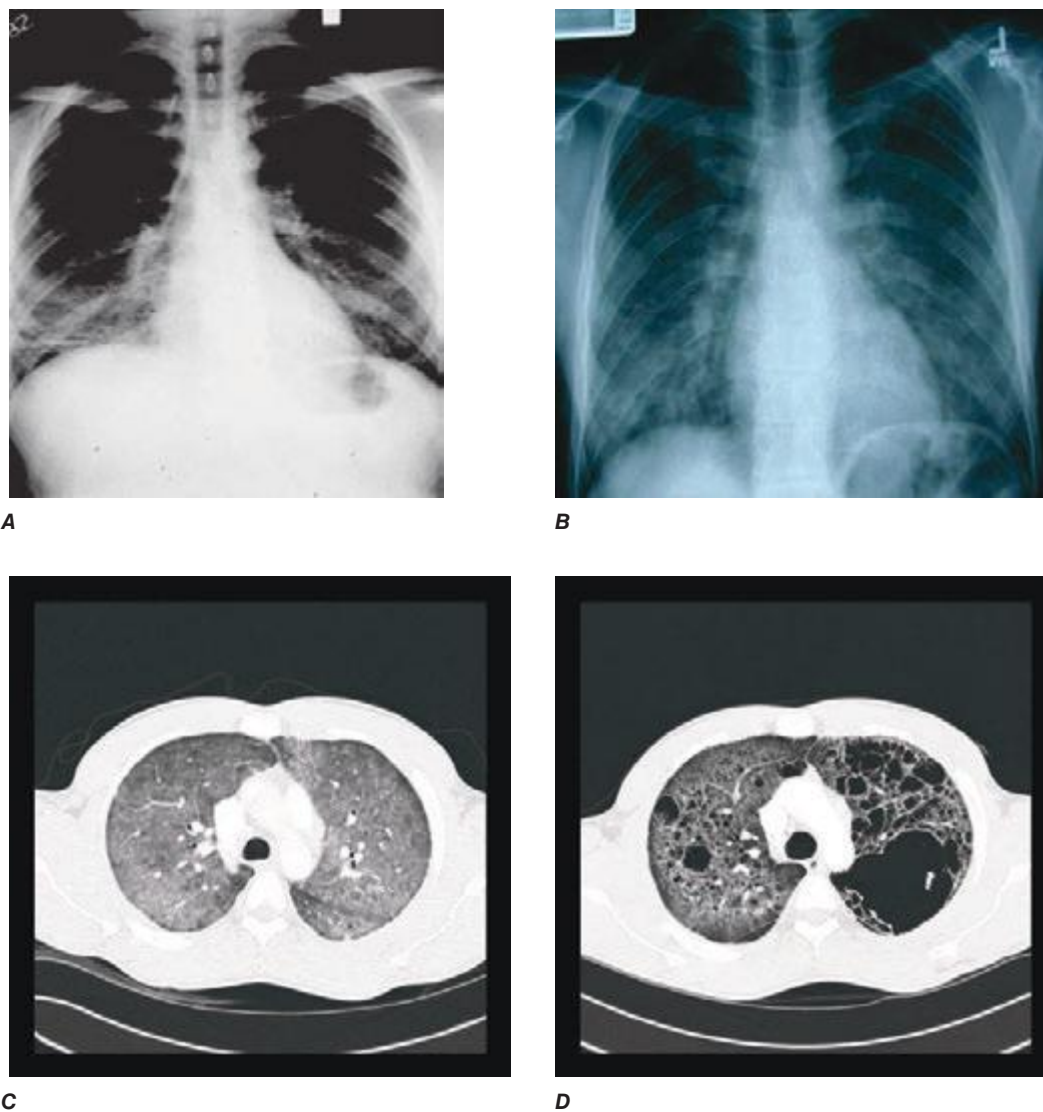


FIGURE 244-2 Radiographs in *Pneumocystis pneumonia*. **A.** Posterior-anterior chest radiograph showing symmetric interstitial infiltrates. **B.** Posterior-anterior chest radiograph showing symmetric alveolar infiltrates (courtesy of Alison Morris). **C.** CT image demonstrating symmetric interstitial infiltrates and ground-glass opacities. **D.** CT image showing symmetric interstitial infiltrates, ground-glass opacities, and pneumatoceles.

tissue demonstrates a foamy alveolar infiltrate and a mononuclear interstitial infiltrate (Fig. 244-2A). This appearance is pathognomonic for PCP even though the organisms cannot be specifically identified with this stain. The diagnosis is typically established in lung tissue or pulmonary secretions by highly specific staining of the cyst—e.g., with methenamine silver (Fig. 244-2B), toluidine blue O, or Giemsa (Fig. 244-2C)—or by staining with a specific immunofluorescent antibody (Fig. 244-2D).

The demonstration of organisms in bronchoalveolar lavage fluid is almost 100% sensitive and specific for PCP in patients with either HIV infection or immunosuppression of other etiologies. The organisms are identified with the specific stains indicated above for lung biopsy. While expectorated sputum or throat swabs have very low sensitivity, an induced sputum sample obtained and interpreted by an experienced provider can be highly sensitive and specific. The reported sensitivity of induced sputum for PCP is widely variable (55–90%), however, and is dependent on both the characteristics of the patient and the experience of the center conducting the test.

Recently, many laboratories have offered polymerase chain reaction (PCR) testing of respiratory specimens for *Pneumocystis*. However, these PCR tests are so sensitive that it is difficult to distinguish patients with colonization (i.e., those whose acute lung disease is due to some other process but who have low levels of *Pneumocystis* DNA in the lungs) from those with acute pneumonia due to *Pneumocystis*. Such

PCR tests on appropriate samples may be more useful for ruling out a diagnosis of PCP if they are negative than for definitively attributing the disease to *Pneumocystis*.

There has been considerable interest in serologic tests such as assays for (1→3)- β -D-glucan, levels of which are frequently elevated in patients with PCP. However, no serologic assays developed to date offer substantial sensitivity or specificity.

COURSE AND PROGNOSIS

Untreated, PCP is invariably fatal. Patients with HIV infection often have an indolent course that presents as mild exercise intolerance or chest tightness without fever or cough and a normal or nearly normal posterior-anterior chest radiograph, with progression over days, weeks, or even a few months to fever, cough, diffuse alveolar infiltrates, and profound hypoxemia. Some patients with HIV infection and most patients with other types of immunosuppression have more acute disease that progresses over a few days to respiratory failure. Rare patients also develop distributive shock. A few unusual patients present with extrapulmonary manifestations in the skin or soft tissue, retina, brain, liver, kidney, or spleen that are nonspecific in presentation and can be diagnosed only by histology.

Factors that influence mortality risk include the patient's age and degree of immunosuppression as well as the presence of preexisting

lung disease, a low serum albumin level, the need for mechanical ventilation, and the development of pneumothorax. With advances in supportive critical care, the prognosis for patients with PCP who require intubation and respiratory support has improved and now depends to a large extent on comorbidities and the prognosis of the underlying disease. Since patients typically do not respond to therapy for 4–8 days, supportive care for a minimum of 10 days is a reasonable consideration if such support is compatible with the patient's wishes and the prognosis of comorbidities. Patients whose condition continues to deteriorate after 3 or 4 days or has not improved after 7–10 days should be reevaluated to determine whether other infectious processes are present (either having been missed on initial evaluation or having developed during treatment) or whether noninfectious processes (e.g., congestive heart failure, pulmonary emboli, pulmonary hypertension, drug toxicity, or a neoplastic process) are causing pulmonary dysfunction.

TREATMENT *P. JIROVECI* PNEUMONIA

The treatment of choice for PCP is trimethoprim-sulfamethoxazole (TMP-SMX), given either IV or PO for 14–21 days (Table 244-1). TMP-SMX, which interferes with the organism's folate metabolism, is at least as effective as alternative agents and is better tolerated. However, TMP-SMX can cause leukopenia, hepatitis, rash, and fever as well as anaphylactic and anaphylactoid reactions, and patients with HIV infection have an unusually high incidence of hypersensitivity to TMP-SMX. Monitoring of serum drug levels is useful if renal function or toxicities are issues. Maintenance of a 2-h post-dose sulfamethoxazole level of 100–150 µg/mL has been associated with a successful outcome. Resistance to TMP-SMX cannot be measured by organism growth inhibition in the laboratory because *Pneumocystis* cannot be cultured. However, mutations in the target gene for sulfamethoxazole that confer in vitro sulfa resistance to other organisms have been found in *Pneumocystis*. The clinical relevance of these mutations for the response to therapy is unknown. Sulfadiazine plus pyrimethamine, an oral regimen more often used

for treatment of toxoplasmosis, also is highly effective. Dapsone plus pyrimethamine or dapsone plus trimethoprim also can be used.

Intravenous pentamidine or the combination of clindamycin plus primaquine is an option for patients who cannot tolerate TMP-SMX and for patients in whose treatment TMP-SMX appears to be failing. Pentamidine must be given IV over at least 60 min to avoid potentially lethal hypotension. Adverse effects can be severe and irreversible and include renal dysfunction, dysglycemia (life-threatening hypoglycemia that can occur days or weeks after initial infusion and be followed by hyperglycemia), neutropenia, and torsades des pointes. Clindamycin plus primaquine is effective, but primaquine can be given only by the oral route—a disadvantage for patients who cannot ingest or absorb oral drugs.

A major advance in therapy for PCP was the recognition that glucocorticoids could improve survival rates among HIV-infected patients with moderate to severe disease (room air PO₂ <70 mmHg; or alveolar-arterial oxygen gradient, ≥35 mmHg). Glucocorticoids appear to reduce the pulmonary inflammation that occurs after specific therapy is started and organisms begin to die, eliciting inflammation. Therapy with glucocorticoids should be the standard of care for patients with HIV infection and probably is also effective for patients with other immunodeficiencies. This treatment should be started for moderate or severe disease when therapy for PCP is initiated, even if the diagnosis has not yet been confirmed. If HIV-infected or HIV-uninfected patients are receiving high-dose glucocorticoids when they develop PCP, there are theoretical advantages to increasing or decreasing the steroid dose, but there is no convincing evidence on which to base any specific strategy.

No definitive trials have defined the best therapeutic algorithm for patients in whom TMP-SMX treatment for PCP is failing. If no other treatable infectious or noninfectious processes are detected and pulmonary dysfunction appears to be due to PCP alone, many authorities would switch from TMP-SMX to either IV pentamidine or IV clindamycin plus oral primaquine. Some authorities would add the second drug or drug combination to TMP-SMX rather than switching regimens. If patients are not already receiving them, glucocorticoids should be added to the regimen; the dosage and regimen, which are usually chosen empirically, depend on what glucocorticoid regimen (if any) the patient was receiving when PCP therapy was begun.

For patients with HIV infection who present with PCP before the initiation of ART, ART should be started within the first 2 weeks of therapy for PCP in most cases. Immune reconstitution inflammatory syndrome (IRIS) can occur, however, and the decision to initiate ART thus requires considerable expertise in terms of optimal timing relative to PCP recovery as well as in the other factors that are relevant when ART is initiated in any patient.

PREVENTION

The most effective method for preventing PCP is to eliminate the cause of immunosuppression by withdrawing immunosuppressive therapy or treating the underlying cause, e.g., HIV infection. Patients who are susceptible to PCP benefit from chemoprophylaxis during the period of susceptibility. For patients with HIV infection, CD4+ T cell counts are a reliable marker of susceptibility, and counts below 200 cells/µL are an indication to start prophylaxis (Table 244-2). For patients with HIV infection who are not receiving ART, oral candidiasis or prior PCP also is an indication for chemoprophylaxis, regardless of CD4+ T cell count. For such patients not receiving ART, any prior episode of an AIDS-defining illness or pneumonia should encourage the use of chemoprophylaxis. However, patients who are not adherent to ART are not likely to take PCP chemoprophylaxis.

For patients without HIV infection, there is no laboratory parameter, including the CD4+ T cell count, that predicts susceptibility to PCP with adequate positive and negative accuracy. The period of susceptibility is usually estimated on the basis of experience with the

TABLE 244-1 TREATMENT OF PNEUMOCYSTOSIS (14–21 DAYS)

Drug(s)	Dose, Route	Adverse Effects
First-Choice Agent		
TMP-SMX	TMP (5 mg/kg) plus SMX (25 mg/kg) q6–8h PO or IV (2 double-strength tablets tid or qid)	Fever, rash, cytopenias, hepatitis, hyperkalemia
Alternative Agents		
TMP <i>plus</i> Dapsone	5 mg/kg q6–8h PO 100 mg qd PO	Hemolysis (G6PD deficiency), methemoglobinemia, rash, fever, gastrointestinal disturbances
Atovaquone	750 mg bid PO	Rash, fever, hepatitis
Clindamycin <i>plus</i> Primaquine	300–450 mg q6h PO or 600 mg q6–8h IV 15–30 mg qd PO	Hemolysis (G6PD deficiency), methemoglobinemia, neutropenia, rash
Pentamidine	3–4 mg/kg qd IV	Hypotension, azotemia, cardiac arrhythmias (torsades des pointes), pancreatitis, dysglycemia, hypocalcemia, neutropenia, hepatitis
Adjunctive Agent		
Prednisone or methylprednisolone	40 mg bid × 5 d, 40 mg qd × 5 d, 20 mg qd × 11 d; PO or IV	Peptic ulcer disease, hyperglycemia, mood alteration, hypertension

Abbreviations: G6PD, glucose-6-phosphate dehydrogenase; TMP-SMX, trimethoprim-sulfamethoxazole.

TABLE 244-2 PROPHYLAXIS OF PNEUMOCYSTOSIS

Drug(s)	Dose, Route	Comments
First-Choice Agent		
TMP-SMX	1 tablet (double- or single-strength) qd PO	Incidence of hypersensitivity is high. Rechallenge for non-life-threatening hypersensitivity; consider dose-escalation protocol.
Alternative Agents		
Dapsone	50 mg bid or 100 mg qd PO	Hemolysis is associated with G6PD deficiency.
Dapsone <i>plus</i>	50 mg qd PO	Leucovorin ameliorates cytopenias due to pyrimethamine.
Pyrimethamine <i>plus</i>	50 mg weekly PO	
Leucovorin	25 mg weekly PO	
Dapsone <i>plus</i>	200 mg weekly PO	Leucovorin ameliorates cytopenias due to pyrimethamine.
Pyrimethamine <i>plus</i>	75 mg weekly PO	
Leucovorin	25 mg weekly PO	
Pentamidine	300 mg monthly via Respigard II nebulizer	Aerosol may cause bronchospasm. Pentamidine is probably less effective than TMP-SMX or dapsone regimens.
Atovaquone	1500 mg qd PO	Requires fatty meal for optimal absorption.

Abbreviations: G6PD, glucose-6-phosphate dehydrogenase; TMP-SMX, trimethoprim-sulfamethoxazole.

underlying disease and immunosuppressive regimen. Patients receiving a prolonged course of high-dose glucocorticoids appear to be particularly susceptible to PCP. The glucocorticoid exposure threshold that warrants chemoprophylaxis is controversial, but such preventive therapy should be strongly considered for any patient receiving more than the equivalent of 20 mg of prednisone daily for 30 days.

TMP-SMX is the most effective prophylactic drug: few patients experience a PCP breakthrough when they are reliably taking a recommended TMP-SMX chemoprophylactic regimen. Several TMP-SMX regimens have been used successfully. One double-strength tablet daily is the regimen with which there is the most experience, but either one single-strength tablet daily or one double-strength tablet two or three times weekly also has been recommended for various populations of patients.

For patients who cannot tolerate TMP-SMX (usually because of hypersensitivity or bone marrow suppression), alternative drugs include daily dapsone, weekly dapsone-pyrimethamine, and monthly aerosol pentamidine. Patients who develop hypersensitivity to TMP-SMX can sometimes tolerate the drug if a gradual dose-escalation protocol is used. Dapsone cross-reacts with sulfonamides in a substantial fraction of patients and therefore is rarely useful in patients with a history of life-threatening reactions to TMP-SMX. Aerosolized pentamidine is highly effective, but it is not as effective as TMP-SMX and may not provide protection in areas of the lung that are not well ventilated. Atovaquone is also effective and well tolerated; however, this drug is available only as an oral preparation, and gastrointestinal absorption is unpredictable in patients with abnormal gastrointestinal motility or function.

245e Laboratory Diagnosis of Parasitic Infections

Sharon L. Reed, Charles E. Davis

The cornerstone for the diagnosis of parasitic infections is a thorough history of the patient's illness. Epidemiologic aspects of the illness are especially important because the risks of acquiring many parasites are closely related to occupation, recreation, or travel to areas of high endemicity. Without a basic knowledge of the epidemiology and life cycles of the major parasites, it is difficult to approach the diagnosis of parasitic infections systematically. Accordingly, the medical classification of important human parasites in this chapter emphasizes their geographic distribution, their transmission, and the anatomic location and stages of their life cycle in humans. The text and tables are intended to serve as a guide to the correct diagnostic procedures for the major parasitic infections; in addition, the reader is referred to other chapters that contain more comprehensive information about each infection (Chaps. 247–260). Tables 245e-1, 245e-2, and 245e-3 summarize the geographic distributions, the anatomic locations, and the methods employed for the diagnosis of flatworm, roundworm, and protozoal infections, respectively.

In addition to selecting the correct diagnostic procedures, physicians must counsel their patients to ensure that specimens are collected properly and arrive at the laboratory promptly. For example, the diagnosis of bancroftian filariasis is unlikely to be confirmed by the laboratory unless blood is drawn near midnight, when the nocturnal microfilariae are active. Laboratory personnel and surgical pathologists should be notified in advance when a parasitic infection is suspected. Continuing interaction with the laboratory staff and the surgical pathologists increases the likelihood that parasites in body fluids or biopsy specimens will be examined carefully by the most capable individuals.

INTESTINAL PARASITES

Most helminths and protozoa exit the body in the fecal stream. Many laboratories now use a stool collection kit with instructions for patients to transfer portions of the sample directly into bacterial carrier medium and fixative, which increases yield. If kits are not available, the patient should be instructed to collect feces in a clean waxed or cardboard container and to record the time of collection on the container. Refrigeration will preserve trophozoites for a few hours and protozoal cysts and helminthic ova for several days. Contamination with water (which could contain free-living protozoa) or with urine (which can damage trophozoites) should be avoided. Fecal samples should be collected before ingestion of barium or other contrast agents for radiologic procedures and before treatment with antidiarrheal agents and antacids; these substances change the consistency of the feces and interfere with microscopic detection of parasites. Because of the cyclic shedding of most parasites in the feces, a minimum of three samples collected on alternate days should be examined. Examination of a single sample can be up to 50% less sensitive.

Analysis of fecal samples entails both macroscopic and microscopic examination. Watery or loose stools are more likely to contain protozoal trophozoites, but protozoal cysts and all stages of helminths may be found in formed feces. If adult worms or tapeworm segments are observed, they should be transported promptly to the laboratory or washed and preserved in fixative for later examination. The only tapeworm with motile segments is *Taenia saginata*, the beef tapeworm, which patients sometimes bring to the physician. Because the ova of

T. saginata are morphologically indistinguishable from those of *Taenia solium* (the cause of cysticercosis), motility is an important distinguishing characteristic.

Microscopic examination of feces is not complete until direct wet mounts have been evaluated and concentration techniques as well as permanent stains have been applied. Before accepting a report of negativity for ova and parasites as final, the physician should insist that the laboratory undertake each of these procedures. Some intestinal parasites are more readily detected in material other than feces. For example, examination of duodenal contents is sometimes necessary to detect *Giardia lamblia*, *Cryptosporidium*, and *Strongyloides* larvae. Use of the “cellophane tape” technique to detect pinworm ova on the perianal skin sometimes also reveals ova of *T. saginata* deposited perianally when the motile segments disintegrate (Table 245e-4).

Two routine solutions are used to make wet mounts for identification of the various life stages of helminths and protozoa: physiologic saline for trophozoites, cysts, ova, and larvae and dilute iodine solution for protozoal cysts and ova. Iodine solution must never be used to examine specimens for trophozoites because it kills the parasites and thus eliminates their characteristic motility.

The two most common concentration procedures for detecting small numbers of cysts and ova are formalin-ether sedimentation and zinc sulfate flotation. The formalin-ether technique is preferable because all parasites sediment but not all float. Slides permanently stained for trophozoites should be prepared before concentration. Additional slides stained for cysts and ova may be made from the concentrate.

In many instances, especially in the differentiation of *Entamoeba histolytica* from other amebas, identification of parasites from wet mounts or concentrates must be considered tentative. Permanently stained smears allow study of the cellular detail necessary for definitive identification. The iron-hematoxylin stain is excellent for critical work, but trichrome staining, which can be completed in 1 h, is a satisfactory alternative that also reveals parasites in specimens preserved in polyvinyl alcohol fixative. Modified acid-fast staining and fluorescent auramine microscopy are useful adjuncts for detection and identification of several intestinal protozoa, including *Cryptosporidium* and *Cyclospora*, while direct fluorescent antibody testing is in common use for *Cryptosporidium* and *Giardia*. Microsporidia, which cause chronic diarrhea in HIV-infected patients, may be missed unless a special modified trichrome stain or fluorescent screening with calcofluor white is requested (Table 245e-3).

BLOOD AND TISSUE PARASITES

Invasion of tissue by protozoa and helminths renders the choice of diagnostic techniques more difficult. For example, physicians must understand that aspiration of an amebic liver abscess rarely reveals *E. histolytica* because the trophozoites are located primarily in the abscess wall. They must remember that the urine sediment offers the best opportunity to detect *Schistosoma haematobium* in the young Ethiopian immigrant or the American traveler who returns from Africa with hematuria. Tables 245e-1, 245e-2, and 245e-3, which offer a quick guide to the geographic distribution and anatomic locations of the major tissue parasites, should help the physician to select the appropriate body fluid or biopsy site for microscopic examination. Tables 245e-5 and 245e-6 provide additional information about the identification of parasites in samples from specific anatomic locations. The laboratory procedures for detection of parasites in other body fluids are similar to those used in the examination of feces. The physician should insist on wet mounts, concentration techniques, and permanent stains for all body fluids. The trichrome or iron-hematoxylin stain is satisfactory for all tissue helminths in body fluids other than blood,

TABLE 245e-1 FLATWORM INFECTIONS

Parasite	Geographic Distribution	Life-Cycle Hosts		Diagnosis			
		Intermediate (Transmission)	Definitive	Parasite Stage	Body Fluid or Tissue	Serologic Tests	Other
Tapeworms (Cestodes)							
Intestinal tapeworms							
<i>Taenia saginata</i> (beef tapeworm)	Worldwide	Beef	Humans	Ova, segments	Feces	—	Motile segments
<i>Hymenolepis nana</i> (dwarf tapeworm)	Worldwide	Grain beetles	Humans, mice ^a	Ova	Feces	—	—
<i>Diphyllobothrium latum</i> (fish tapeworm)	Worldwide	Copepods–fish ^b	Humans, other mammals	Ova, segments	Feces	—	Megaloblastic anemia in 1%
<i>T. solium</i> ^c (pork tapeworm)	Worldwide	Swine	Humans	Ova, segments	Feces	WB	Especially Mexico, Central and South America, Africa
Somatic tapeworms							
<i>Echinococcus granulosus</i> (hydatid disease)	Sheep-raising and hunting areas	Sheep, camels, humans, others	Dogs	Hydatid	Lung, liver	WB, EIA	Chest radiography, CT, MRI
<i>E. multilocularis</i> (hydatid disease)	Subarctic areas	Rodents, humans	Foxes, dogs, cats	Hydatid	Liver	—	May resemble cholangiocellular carcinoma
<i>T. solium</i> ^c (pork tapeworm)	Worldwide	Swine, humans	Humans	Cysticercus	Muscles, CNS	WB	CT, MRI, radiography
Flukes (Trematodes)							
Intestinal flukes							
<i>Fasciolopsis buski</i>	China, India	Snails–water chestnuts	Humans	Ova	Feces	—	—
<i>Heterophyes heterophyes</i>	Far East, India	Snails–fish	Humans	Ova	Feces	—	—
<i>Metagonimus yokogawai</i>	Far East, Balkans, North Africa	Snails–fish	Humans	Ova	Feces	—	—
Liver flukes							
<i>Clonorchis sinensis</i>	China, Southeast Asia	Snails–fish	Humans	Ova	Feces, bile	—	Recurrent bacterial cholangitis
<i>Fasciola hepatica</i>	Sheep-raising areas	Snails–watercress	Humans, sheep	Ova	Feces, ^d bile	EIA	Cirrhosis, portal hypertension
Lung flukes							
<i>Paragonimus</i> spp.	Orient, Africa, the Americas	Snails–crabs/crayfish	Humans, other mammals	Adults, ova	Lung, sputum, feces	WB, EIA	Chest radiography, CT, MRI
Blood flukes							
<i>Schistosoma mansoni</i>	Africa, Central and South America, West Indies	Snails	Humans	Ova, adults	Feces	EIA, WB	Rectal snips, liver biopsy
<i>S. haematobium</i>	Africa	Snails	Humans	Ova, adults	Urine	WB	Liver, urine, or bladder biopsy
<i>S. japonicum</i>	Far East	Snails	Humans	Ova, adults	Feces	WB	Liver biopsy

^aLarvae also can mature in intestinal villi of humans and mice. ^bWhen there are two intermediate hosts, the first is separated from the second by a dash. Definitive hosts are infected by the second intermediate host. ^c*T. solium* can cause either intestinal infections or cysticercosis. Its ova are identical to those of *T. saginata*; scolices and segments of the two species differ. ^dOva seldom reach the fecal stream during acute disease.

Note: CNS, central nervous system; EIA, enzyme immunoassay; WB, western blot. Serologic tests listed in Tables 245e-1, 245e-2, and 245e-3 are available commercially or from the Centers for Disease Control and Prevention, Atlanta, GA.

but microfilarial worms and blood protozoa are more easily visualized with Giemsa or Wright's staining.

The parasites most commonly detected in Giemsa-stained blood smears are the plasmodia, microfilariae, and African trypanosomes (Table 245e-5). Most patients with Chagas' disease present in the chronic phase, when *Trypanosoma cruzi* is no longer microscopically detectable in blood smears. Wet mounts are sometimes more sensitive than stained smears for the detection of microfilariae and African trypanosomes because these active parasites cause noticeable movement of the erythrocytes in the microscopic field. Filtration of blood through

a polycarbonate filter (pore size, 3–5 μm) facilitates the detection of microfilariae. The intracellular amastigote forms of *Leishmania* species and *T. cruzi* can sometimes be visualized in stained smears of peripheral blood, but aspirates of the bone marrow, liver, and spleen are the best sources for microscopic detection and culture of *Leishmania* in kala-azar and of *T. cruzi* in chronic Chagas' disease. The diagnosis of malaria and the critical distinction among the various *Plasmodium* species are made by microscopic examination of stained thick and thin blood films (Chap. 248). Because the lab infrastructure and technical expertise may not be available in many rural areas with high levels of

TABLE 245e-2 ROUNDWORM INFECTIONS

Parasite	Geographic Distribution	Life-Cycle Hosts		Diagnosis			
		Intermediate (Transmission)	Definitive	Parasite Stage	Body Fluid or Tissue	Serologic Tests	Other
Intestinal Roundworms							
<i>Enterobius vermicularis</i> (pinworm)	Temperate and tropical zones	Fecal-oral	Humans	Ova	Perianal skin	—	“Cellophane tape” test
<i>Trichuris trichiura</i> (whipworm)	Temperate and tropical zones	Soil, fecal-oral	Humans	Ova	Feces	—	Rectal prolapse
<i>Ascaris lumbricoides</i> (roundworm of humans)	Temperate and tropical zones	Soil, fecal-oral	Humans	Ova	Feces	—	Sx of pulmonary migration
<i>Ancylostoma duodenale</i> (Old World hookworm)	Eurasia, Africa, Pacific	Soil to skin	Humans	Ova/larvae	Feces	—	Sx of pulmonary migration, anemia
<i>Necator americanus</i> (New World hookworm)	U.S., Africa, worldwide	Soil to skin	Humans	Ova/larvae	Feces	—	Sx of pulmonary migration, anemia
<i>Strongyloides stercoralis</i> (strongyloidiasis)	Moist tropics and subtropics	Soil to skin	Humans	Larvae	Feces, sputum, duodenal fluid	EIA	Dissemination in immunodeficiency
<i>Capillaria philippinensis</i>	Southeast Asia, Taiwan, Egypt	Raw fish	Birds	Ova, larvae, adults	Feces	—	Malabsorption/autoinfection, biopsy
Tissue Roundworms							
<i>Trichinella spiralis</i> (trichinellosis)	Worldwide	Swine/humans	Swine/humans	Larvae	Muscle	EIA	Muscle biopsy
<i>Wuchereria bancrofti</i> (filariasis)	Coastal areas in tropics and subtropics	Mosquitoes	Humans	Microfilariae	Blood, lymph nodes	EIA, RAPID, PCR ^b	Nocturnal periodicity ^a
<i>Brugia malayi</i> (filariasis)	Asia, Indian subcontinent	Mosquitoes	Humans	Microfilariae	Blood	EIA, RAPID, PCR ^b	Nocturnal
<i>Loa loa</i> (African eye worm)	West and Central Africa	Mango flies (<i>Chrysops</i>)	Humans	Microfilariae	Blood	LIPS ^b , PCR ^b	May be visible in eye, diurnal
<i>Onchocerca volvulus</i> (river blindness)	Africa, Mexico, Central and South America	Blackflies	Humans	Adults/larvae	Skin/eye	LIPS ^b , PCR ^b	Examine nodules or skin snips
<i>Dracunculus medinensis</i> (guinea worm)	Africa	<i>Cyclops</i>	Humans	Adults/larvae	Skin	—	May be visible in lesion
<i>Angiostrongylus cantonensis</i>	Southeast Asia, Pacific, Caribbean	Snails/slugs, shrimp/fish	Rats	Larvae	CSF (rarely found)	—	Eosinophilic meningitis
Larva Migrans Syndromes							
<i>Ancylostoma braziliense</i> (creeping eruption)	Tropical and temperate zones	Soil to skin	Dogs/cats, humans	Larvae	Skin	—	Dog and cat hookworm
<i>Toxocara canis</i> and <i>T. cati</i> (visceral larva migrans), <i>Baylisascaris</i>	Tropical and temperate zones	Soil, fecal-oral	Dogs/cats, raccoons, humans	Larvae	Viscera, CNS, eye	EIA	Also caused by roundworms of other species

^aExcept for infection acquired in the South Pacific, blood should be drawn at midnight. ^bLIPS (the luciferase immunoprecipitation system) for serology and PCR (polymerase chain reaction) are available from the Laboratory of Parasitic Diseases at the National Institutes of Health: 301-496-5398.

Abbreviations: CNS, central nervous system; CSF, cerebrospinal fluid; EIA, enzyme immunoassay; RAPID, rapid immunographic assay (available internationally); Sx, signs/symptoms.

malaria, rapid detection tests (RDTs) are increasingly being used to fill this gap. These are immunochromatographic capture assays with monoclonal antibodies to species-specific antigens (histidine-rich protein 2 [PfHRP2] or aldolase of *Plasmodium falciparum*) or conserved *Plasmodium* antigens (lactate dehydrogenase). The World Health Organization sponsored a major testing program evaluating the different RDTs. Lower performance rates have been reported in a variety of field sites, especially in areas where deletions of the *pfhrp2* gene have been detected. Subpatent infection and identification of *Plasmodium* species can also be confirmed by polymerase chain reaction (PCR), but for this purpose PCR is primarily a research tool. *P. knowlesi*, a simian parasite, has been identified as the cause of an increasing number of infections in Malaysian Borneo and other areas of Southeast Asia; PCR

or another molecular method is required to differentiate *P. knowlesi* from *P. malariae*.

Although most tissue parasites stain with the traditional hematoxylin and eosin, appropriate special stains should also be applied to surgical biopsy specimens. The surgical pathologist who is accustomed to applying silver stains for *Pneumocystis* to induced sputum and transbronchial biopsies may need to be reminded to examine wet mounts and iron-hematoxylin-stained preparations of pulmonary specimens for helminthic ova and *E. histolytica*. The clinician should also be able to advise the surgeon and pathologist about optimal techniques for the identification of parasites in specimens obtained by certain specialized minor procedures (Table 245e-6). For example, the excision of skin snips for the diagnosis of onchocerciasis, the collection of rectal snips

TABLE 245e-3 PROTOZOAL INFECTIONS

Parasite	Geographic Distribution	Life-Cycle Hosts		Parasite Stage	Diagnosis		
		Intermediate (Transmission)	Definitive		Body Fluid or Tissue	Serologic Tests	Other
Intestinal Protozoans							
<i>Entamoeba histolytica</i> (amebiasis)	Worldwide, especially tropics	Fecal-oral	Humans	Troph, cyst	Feces, liver	EIA, antigen detection	Ultrasound, liver CT, PCR
<i>Giardia lamblia</i> (giardiasis)	Worldwide	Fecal-oral	Humans	Troph, cyst	Feces	Antigen detection	DFA, PCR
<i>Isospora belli</i>	Worldwide	Fecal-oral	Humans	Oocyst	Feces	—	Acid-fast ^a
<i>Cryptosporidium</i>	Worldwide	Fecal-oral	Humans, other animals	Oocyst	Feces	Antigen detection	Acid-fast, ^a DFA, biopsy, PCR
<i>Cyclospora cayentanensis</i>	Worldwide?	Fecal-oral	Humans, other animals?	Oocyst	Feces	—	Acid-fast, ^a modified safranin, autofluorescence, biopsy, PCR
Microsporidia: <i>Enterocytozoon bieneusi</i> , <i>Encephalitozoon</i> spp. (microsporidiosis)	Worldwide?	?	Animals, humans	Spore	Feces	—	Modified trichrome, biopsy, PCR
Free-Living Amebas							
<i>Naegleria</i>	Worldwide	Warm water	Humans	Troph, cyst	CNS, nares	DFA	Biopsy, nasal swab, culture
<i>Acanthamoeba</i>	Worldwide	Soil, water	Humans	Troph, cyst	CNS, skin, cornea	DFA	Biopsy, scrapings, culture
<i>Balamuthia</i>	The Americas	Soil?	Humans, other animals	Troph, cyst	Brain	DFA	Biopsy, PCR
Blood and Tissue Protozoans							
<i>Plasmodium</i> spp. (malaria)	Subtropics and tropics	Mosquitoes	Humans	Asexual	Blood	RDT	PCR
<i>Babesia microti</i> (babesiosis)	U.S., especially New England	Ticks	Rodents, humans	Asexual	Blood	IIF	Risk in asplenia, PCR
<i>Trypanosoma rhodesiense</i> (African sleeping sickness)	Sub-Saharan East Africa	Tsetse flies	Humans, herbivores	Tryp	Blood, CSF	IIF ^b	Also chancre, lymph nodes
<i>T. gambiense</i> (African sleeping sickness)	Sub-Saharan West Africa	Tsetse flies	Humans, swine	Tryp	Blood, CSF	Card agglutination, ^c IIF ^b	Also chancre, lymph nodes
<i>T. cruzi</i> (Chagas' disease)	Mexico to South America	Reduviid bugs (triatomes)	Humans, dogs, wild animals	Amastigote, trypan	Multiple organs/ blood	IIF, EIA	Reactivation in immunosuppression
<i>Leishmania tropica</i> , etc.	Widespread in tropics and subtropics	Sandflies (<i>Phlebotomus</i>)	Humans, dogs, rodents	Amastigote	Skin	IFA, EIA ^d	Biopsy, scrapings, culture
<i>L. braziliensis</i> (mucocutaneous)	Mexico to South America	Sandflies (<i>Lutzomyia</i>)	Humans, dogs, rodents	Amastigote	Skin, mucous membranes	IFA ^b , EIA	Biopsy, scrapings, culture
<i>L. donovani</i> (kala-azar)	Widespread in tropics and subtropics	Sandflies (<i>Phlebotomus</i>)	Humans, dogs, wild animals	Amastigote	RE system	IFA ^b , EIA	Biopsy, culture, PCR
<i>Toxoplasma gondii</i> (toxoplasmosis)	Worldwide	Humans, other mammals	Cats	Cyst, troph	CNS, eye, muscles, other	EIA, IIF	PCR

^aAcid-fastness is best demonstrated by auramine fluorescence or modified acid-fast stain. ^bContact the CDC at 404-718-4100. ^cCard agglutination is provided to endemic countries by the World Health Organization. ^dLimited specificity; most sensitive for *L. donovani*.

Abbreviations: CNS, central nervous system; CSF, cerebrospinal fluid; DFA, direct fluorescent antibody; EIA, enzyme immunoassay; IFA, indirect fluorescent antibody; IIF, indirect immunofluorescence; PCR, polymerase chain reaction; RDT, rapid detection test; RE, reticuloendothelial; troph, trophozoite; trypan, trypanomastigote form.

for the diagnosis of schistosomiasis, and punch biopsy of skin lesions for the identification and culture of cutaneous and mucocutaneous species of *Leishmania* are simple procedures, but the diagnosis can be missed if the specimens are improperly obtained or processed.

NONSPECIFIC TESTS

Eosinophilia (>500/ μ L) commonly accompanies infections with most of the tissue helminths; absolute numbers of eosinophils may be high in trichinellosis and the migratory phases of filariasis (Table 245e-7).

Intestinal helminths provoke eosinophilia only during pulmonary migration of the larval stages. Eosinophilia is not a manifestation of protozoal infections. Parasitic causes of eosinophilia in cerebrospinal fluid include nematodes (e.g., *Angiostrongylus*, *Gnathostoma*, *Toxocara*, and *Baylisascaris* species) as well as flatworms (e.g., *T. solium* and *Schistosoma* species).

Like the hypochromic microcytic anemia of heavy hookworm infections, other nonspecific laboratory abnormalities may suggest parasitic infection in patients with appropriate geographic and/or environmental exposures. Biochemical evidence of cirrhosis or an

TABLE 245e-4 ALTERNATIVE PROCEDURES FOR LABORATORY DIAGNOSIS OF PARASITES FOUND IN FECES^a

Parasites and Fecal Stages	Alternative Diagnostic Procedures
Tapeworms (Cestodes)	
<i>Taenia saginata</i> ova and segments	Perianal "cellophane tape" test for ova
<i>T. solium</i> ova and segments	Serology; brain biopsy for neurocysticercosis
Flukes (Trematodes)	
<i>Clonorchis (Opisthorchis) sinensis</i> ova	Examination of bile for ova and adults in cholangitis
<i>Fasciola hepatica</i> ova	Examination of bile for ova and adults in cholangitis
<i>Paragonimus</i> ova	Serology; sputum; biopsy of lung or brain for larvae
<i>Schistosoma</i> ova	Serology for all; rectal snips (especially for <i>S. mansoni</i>), urine (<i>S. haematobium</i>), liver biopsy and liver ultrasound
Roundworms	
<i>Enterobius vermicularis</i> ova and adults	Perianal "cellophane tape" test for ova and adults
<i>Trichuris trichiura</i> ova	None
<i>Ascaris lumbricoides</i> ova and adults	Examination of sputum for larvae in lung disease
Hookworm ova and occasional larvae	Examination of sputum for larvae in lung disease
<i>Strongyloides</i> larvae	Duodenal aspirate or jejunal biopsy; serology; sputum or lung biopsy for filariform larvae in disseminated disease
Protozoans	
<i>Entamoeba histolytica</i> trophozoites and cysts	Serology; liver biopsy for trophozoites and cysts
<i>Giardia lamblia</i> trophozoites and cysts	Duodenal aspirate or jejunal biopsy
<i>Isoospora belli</i> oocysts	Duodenal aspirate or jejunal biopsy ^b
<i>Cryptosporidium</i> oocysts	Duodenal aspirate or jejunal biopsy ^b
Microsporidian spores	Duodenal aspirate or jejunal biopsy

^aStains and concentration techniques are discussed in the text. ^b*Isoospora* and *Cryptosporidium* are acid-fast.

TABLE 245e-5 IDENTIFICATION OF PARASITES IN BLOOD AND OTHER BODY FLUIDS

Body Fluid, Parasite	Enrichment/Stain	Culture Technique
Blood		
<i>Plasmodium</i> spp.	Thick and thin smears/Giemsa or Wright's	Not useful for diagnosis
<i>Leishmania</i> spp.	Buffy coat/Giemsa	Media available from CDC
African trypanosomes ^a	Buffy coat, anion column/wet mount and Giemsa	Mouse or rat inoculation ^b
<i>Trypanosoma cruzi</i> ^c	As for African species	As above and xenodiagnosis
<i>Toxoplasma gondii</i>	Buffy coat/Giemsa	Fibroblast cell lines
Microfilariae ^d	Filtration/wet mount and Giemsa	None
Urine		
<i>Schistosoma haematobium</i>	Centrifugation/wet mount	None
Microfilariae (in chyluria)	As for blood	None
Spinal Fluid		
African trypanosomes	Centrifugation, anion column/wet mount and Giemsa	As for blood
<i>Naegleria fowleri</i>	Centrifugation/wet mount and Giemsa or trichrome	Nonnutrient agar overlaid with <i>Escherichia coli</i>

^a*Trypanosoma rhodesiense* and *T. gambiense*. ^bInject mice intraperitoneally with 0.2 mL of whole heparinized blood (0.5 mL for rats). After 5 days, check tail blood daily for trypanosomes as described above. Call the CDC (770-488-7775) for information on diagnosis and treatment. ^cDetectable in blood by conventional techniques only during acute disease. Xenodiagnosis is successful in ~50% of patients with chronic Chagas' disease. ^dDaytime (1000–1400 h) and nighttime (2200–0200 h) blood samples should be drawn to maximize the chance of detecting *Wuchereria* (nocturnal except for Pacific strains), *Brugia* (nocturnal), and *Loa loa* (diurnal).

TABLE 245e-6 MINOR PROCEDURES FOR DIAGNOSIS OF PARASITIC INFECTIONS

Parasite(s) and Stage	Procedure
<i>Onchocerca volvulus</i> and <i>Mansonella streptocerca</i> microfilariae	Skin snips: Lift skin with a needle and excise ~1 mg to a depth of 0.5 mm from several sites. Weigh each sample, place it in 0.5 mL of saline for 4 h, and examine wet mounts and Giemsa stains of the saline either directly or after filtration. Count microfilariae. ^a
<i>Loa loa</i> adults and <i>O. volvulus</i> adults and microfilariae	Biopsies of subcutaneous nodules: Stain routine histopathologic sections and impression smears with Giemsa.
<i>Trichinella spiralis</i> larvae (and perhaps <i>Taenia solium</i> cysticerci)	Muscle biopsies: Excise ~1.0 g of deltoid or gastrocnemius muscle and squash between two glass slides for direct microscopic examination.
<i>Schistosoma</i> ova of all species, but especially <i>S. mansoni</i>	Rectal snips: From four areas of mucosa, take 2-mg snips, tease onto a glass slide, and flatten with a second slide before examining directly at 10x. Preparations may be fixed in alcohol or stained.
<i>Trypanosoma gambiense</i> and <i>T. rhodesiense</i> trypomastigotes	Aspirate of chancre or lymph node: ^b Aspirate center with an 18-gauge needle, place a drop on a slide, and examine for motile forms. An otherwise insufficient volume of material may be stained with Giemsa.
<i>Acanthamoeba</i> spp. trophozoites or cysts	Corneal scrapings: Have an ophthalmologist obtain a sample for immediate Giemsa staining and culture on nutrient agar overlaid with <i>Escherichia coli</i> .
Cutaneous and mucocutaneous <i>Leishmania</i> spp.	Swabs, aspirates, or punch biopsies of skin lesions: Obtain a specimen from the margin of a lesion for Giemsa staining of impression smears; section and culture on special media from the CDC.

^aCounts of >100/mg are associated with a significant risk of complications. ^bLymph node aspiration is contraindicated in some infections and should be used judiciously.

TABLE 245e-7 PARASITES FREQUENTLY ASSOCIATED WITH EOSINOPHILIA^a

Parasite	Comment
Tapeworms (Cestodes)	
<i>Echinococcus granulosus</i>	When hydatid cyst leaks
<i>Taenia solium</i>	During muscle encystation and in cerebrospinal fluid with neurocysticercosis
Flukes (Trematodes)	
<i>Paragonimus</i> spp.	Uniformly high in acute stage
<i>Fasciola hepatica</i>	May be high in acute stage
<i>Clonorchis (Opisthorchis) sinensis</i>	Variable
<i>Schistosoma mansoni</i>	50% of infected travelers
<i>S. haematobium</i>	25% of infected travelers
<i>S. japonicum</i>	Up to 6000/μL in acute infection
Roundworms	
<i>Ascaris lumbricoides</i>	During larval migration
Hookworm species	During larval migration
<i>Strongyloides stercoralis</i>	Profound during migration and early years of infection
<i>Trichinella spiralis</i>	Up to 7000/μL
Filarial species ^b	Varies but can reach 5000–8000/μL
<i>Toxocara</i> spp.	>3000/μL
<i>Ancylostoma braziliense</i>	With extensive cutaneous eruption
<i>Gnathostoma spinigerum</i>	In visceral larva migrans and eosinophilic meningitis
<i>Angiostrongylus cantonensis</i>	In eosinophilic meningitis
<i>A. costaricensis</i>	During larval migration in mesenteric vessels

^aVirtually every helminth has been associated with eosinophilia. This table includes both common and uncommon parasites that frequently elicit eosinophilia during infection. ^b*Wuchereria bancrofti*, *Brugia* spp., *Loa loa*, and *Onchocerca volvulus*.

abnormal urine sediment in an African immigrant certainly raises the possibility of schistosomiasis, and anemia and thrombocytopenia in a febrile traveler or immigrant are among the hallmarks of malaria. CT and MRI also contribute to the diagnosis of infections with many tissue parasites and have become invaluable adjuncts in the diagnosis of neurocysticercosis and cerebral toxoplasmosis.

ANTIBODY AND ANTIGEN DETECTION

Useful antibody assays for many of the important tissue parasites are available; most of those listed in **Table 245e-8** can be obtained from the Centers for Disease Control and Prevention (CDC) in Atlanta. The results of serologic tests not listed in the tables should be interpreted with caution.

The value of antibody assays is limited by several factors. For example, the preparation of thick and thin blood smears remains the procedure of choice for the diagnosis of malaria in individual patients because diagnostic titers to plasmodia develop slowly and do not differentiate species—a critical step in patient management. Filarial antigens cross-react with those from other nematodes; as in assays for antibody to most parasites, the presence of antibody in the filarial assay fails to distinguish between past and current infection. Despite these specific limitations, the restricted geographic distribution of many tropical parasites increases the diagnostic usefulness of both the presence and the absence of antibody in travelers from industrialized countries. In contrast, a large proportion of the world's population has been exposed to *Toxoplasma gondii*, and the presence of IgG antibody to *T. gondii* does not constitute proof of active disease.

Fewer antibody assays are available for the diagnosis of infection with intestinal parasites. *E. histolytica* is the major exception. Sensitive,

TABLE 245e-8 SEROLOGIC AND MOLECULAR TESTS FOR PARASITIC INFECTIONS^a

Parasite, Infection	Antibody	Antigen or DNA/RNA
Tapeworms		
Echinococcosis	WB, EIA	
Cysticercosis	WB	
Flukes		
Paragonimiasis	WB, EIA ^b	
Schistosomiasis	EIA, WB	
Fascioliasis	EIA ^b	
Roundworms		
Strongyloidiasis	EIA	
Trichinellosis	EIA	
Toxocariasis	EIA	
Filariasis	EIA ^c	RAPID ^c
Protozoans		
Amebiasis	EIA	EIA, ^b RAPID; ^b PCR
Giardiasis		EIA, ^b RAPID; ^b DFA, PCR
Cryptosporidiosis		EIA, ^b DFA, RAPID; ^b PCR
Malaria (all species)	IIF ^d	RAPID, PCR
Babesiosis	IIF	PCR
Chagas' disease	IIF, EIA	PCR
Leishmaniasis	IIF, EIA	PCR ^b
Toxoplasmosis	IIF, EIA (IgM) ^e	PCR ^b
Microsporidiosis		PCR
Cyclosporiasis		PCR
Acanthamebiasis		DFA, PCR
Naegleriasis		DFA, PCR
Balamuthiasis		DFA

^aUnless otherwise noted, all tests are available at the CDC. ^bResearch or commercial laboratories only. ^cAvailable at the NIH (301-496-5398) and commercially. ^dOf limited use for management of acute disease. ^eDetermination of infection within the last 3 months may require additional tests by a research laboratory.

Note: DFA, direct fluorescent antibody; EIA, enzyme immunoassay; IIF, indirect immunofluorescence; PCR, polymerase chain reaction; RAPID, rapid immunographic assay; WB, western blot. Most antigen and antibody parasite detection kits are available commercially. Most PCRs listed are now available at the CDC and in commercial or research laboratories. Contact the CDC (404-718-4100).

specific serologic tests are invaluable in the diagnosis of amebiasis. Commercial kits for the detection of antigen by enzyme-linked immunosorbent assay or of whole organisms by fluorescent antibody assay are now available for several protozoan parasites (Table 245e-8).

MOLECULAR TECHNIQUES

DNA hybridization with probes that are repeated many times in the genome of a specific parasite and amplification of a specific DNA fragment by PCR have now been established as useful techniques for the diagnosis of several protozoan infections (Table 245e-8). Although PCR is very sensitive, it is an adjunct to conventional techniques for parasite detection and should be requested only when microscopic and immunodiagnostic procedures fail to establish the probable diagnosis. For example, only multiple negative blood smears or the failure to identify the infecting species justifies PCR for the diagnosis or proper management of malaria. In addition to PCR of anticoagulated blood, the CDC (www.cdc.gov/dpdx/) and several commercial laboratories now perform PCRs for detection of certain specific parasites in stool samples, biopsy specimens, and bronchoalveolar lavage fluid (Table 245e-8). Although PCRs are now used primarily for the detection of protozoans, active research efforts are likely to establish their feasibility for the detection of several helminths.

246e Agents Used to Treat Parasitic Infections

Thomas A. Moore

Parasitic infections afflict more than half of the world's population and impose a substantial health burden, particularly in underdeveloped nations, where they are most prevalent. The reach of some parasitic diseases, including malaria, has expanded over the past few decades as a result of factors such as deforestation, population shifts, global warming, and other climatic events. Despite major efforts at vaccine development and vector control, chemotherapy remains the single most effective means of controlling parasitic infections. Efforts to combat the spread of some diseases are hindered by the development and spread of drug resistance, the limited introduction of new antiparasitic agents, and the proliferation of counterfeit medications. However, there are good reasons to be optimistic. Ambitious global initiatives aimed at controlling or eliminating threats such as AIDS, tuberculosis, and malaria have demonstrated some early successes. Recognition of the substantial burden imposed by the "neglected" tropical diseases has generated multinational partnerships to develop and deploy effective antiparasitic agents. Vaccines against several tropical diseases are being developed, and clinical trials for vaccines against parasites continue.

This chapter deals exclusively with the agents used to treat infections due to parasites. Specific treatment recommendations for the parasitic diseases of humans are listed in subsequent chapters. Many of the agents discussed herein are approved by the U.S. Food and Drug Administration (FDA) but are considered investigational for the treatment of certain infections. Drugs marked in the text with an asterisk (*) are available through the Centers for Disease Control and Prevention (CDC) Drug Service (telephone: 404-639-3670 or 404-639-2888; www.cdc.gov/ncpcid/dsr/). Drugs marked with a dagger (†) are available only through their manufacturers; contact information for these manufacturers may be available from the CDC.

Table 246e-1 presents a brief overview of each agent (including some drugs that are covered in other chapters), along with major

toxicities, spectrum of activity, and safety for use during pregnancy and lactation.

Albendazole Like all benzimidazoles, albendazole acts by selectively binding to free β -tubulin in nematodes, inhibiting the polymerization of tubulin and the microtubule-dependent uptake of glucose. Irreversible damage occurs in gastrointestinal (GI) cells of the nematodes, resulting in starvation, death, and expulsion by the host. This fundamental disruption of cellular metabolism offers treatment for a wide range of parasitic diseases.

Albendazole is poorly absorbed from the GI tract. Administration with a fatty meal increases its absorption by two- to sixfold. Poor absorption may be advantageous for the treatment of intestinal helminths, but successful treatment of tissue helminth infections (e.g., hydatid disease and neurocysticercosis) requires that a sufficient amount of active drug reach the site of infection. The metabolite albendazole sulfoxide is responsible for the drug's therapeutic effect outside the gut lumen. Albendazole sulfoxide crosses the blood-brain barrier, reaching a level significantly higher than that achieved in plasma. The high concentrations of albendazole sulfoxide attained in cerebrospinal fluid (CSF) may explain the efficacy of albendazole in the treatment of neurocysticercosis.

Albendazole is extensively metabolized in the liver, but there are few data regarding the drug's use in patients with hepatic disease. Single-dose albendazole therapy in humans is largely without side effects (overall frequency, $\leq 1\%$). More prolonged courses (e.g., as administered for cystic and alveolar echinococcal disease) have been associated with liver function abnormalities and bone marrow toxicity. Thus, when prolonged use is anticipated, the drug should be administered in treatment cycles of 28 days interrupted by 14-day intervals off therapy. Prolonged therapy with full-dose albendazole (800 mg/d) should be approached cautiously in patients also receiving drugs with known effects on the cytochrome P450 system.

Amodiaquine Amodiaquine has been widely used in the treatment of malaria for >40 years. Like chloroquine (the other major 4-aminoquinoline), amodiaquine is now of limited use because of the spread of resistance. Amodiaquine interferes with hemozoin formation

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
4-Aminoquinolines					
Amodiaquine	Malaria ^b	Agranulocytosis, hepatotoxicity	No information	Not assigned	No information
Chloroquine	Malaria ^b	<i>Occasional:</i> pruritus, nausea, vomiting, headache, hair depigmentation, exfoliative dermatitis, reversible corneal opacity. <i>Rare:</i> irreversible retinal injury, nail discoloration, blood dyscrasias	Antacids and kaolin: reduced absorption of chloroquine Ampicillin: bioavailability reduced by chloroquine Cimetidine: increased serum levels of chloroquine Cyclosporine: serum levels increased by chloroquine	Not assigned ^c	Yes
8-Aminoquinolines					
Primaquine	Malaria ^b	<i>Frequent:</i> hemolysis in patients with G6PD deficiency <i>Occasional:</i> methemoglobinemia, GI disturbances. <i>Rare:</i> CNS symptoms	Quinacrine: potentiated toxicity of primaquine	Contraindicated	No information
Tafenoquine	Malaria ^b	<i>Frequent:</i> hemolysis in patients with G6PD deficiency, mild GI upset. <i>Occasional:</i> methemoglobinemia, headaches	No information	Not assigned	No information

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Aminoalcohols					
Halofantrine	Malaria ^b	<i>Frequent:</i> abdominal pain, diarrhea. <i>Occasional:</i> ECG disturbances (dose-related prolongation of QTc and PR interval), nausea, pruritus. Contraindicated in persons who have cardiac disease or who have taken mefloquine in the preceding 3 weeks	Concomitant use of agents that prolong QTc interval contraindicated	C	No information
Lumefantrine	Malaria ^b	<i>Occasional:</i> nausea, vomiting, diarrhea, abdominal pain, anorexia, headache, dizziness	No major interactions	Not assigned	No information
Aminoglycosides					
Paromomycin	Amebiasis, ^b infection with <i>Dientamoeba fragilis</i> , giardiasis, cryptosporidiosis, leishmaniasis	<i>Frequent:</i> GI disturbances (oral dosing only). <i>Occasional:</i> nephrotoxicity, ototoxicity, vestibular toxicity (parenteral dosing only)	No major interactions	Not assigned ^c	No information
Amphotericin B					
Amphotericin B deoxycholate Amphotec (InterMune) Amphotericin B lipid complex, ABLC (Abelcet) Amphotericin B, liposomal (AmBisome)	Leishmaniasis, ^d amebic meningoencephalitis	<i>Frequent:</i> fever, chills, hypokalemia, hypomagnesemia, nephrotoxicity. <i>Occasional:</i> vomiting, dyspnea, hypotension	Antineoplastic agents: renal toxicity, bronchospasm, hypotension Glucocorticoids, ACTH, digitalis: hypokalemia Zidovudine: increased myelo- and nephrotoxicity	B	No information
Antimonials					
Pentavalent antimony ^e	Leishmaniasis	<i>Frequent:</i> arthralgias/myalgias, pancreatitis, ECG changes (QT prolongation, T wave flattening or inversion)	No major interactions	Not assigned	Yes
Meglumine antimoniate		<i>Frequent:</i> arthralgias/myalgias, pancreatitis, ECG changes (QT prolongation, T wave flattening or inversion)	Antiarrhythmics and tricyclic antidepressants: increased risk of cardiotoxicity	Not assigned	No information
Artemisinin and derivatives					
Arteether Artemether Artesunate ^e Dihydroartemisinin	Malaria ^f	<i>Occasional:</i> neurotoxicity (ataxia, convulsions), nausea, vomiting, anorexia, contact dermatitis	No information No clinically significant interactions Mefloquine: levels decreased and clearance accelerated by artesunate Mefloquine: increased absorption	Not assigned C C Not assigned	Yes ^g Yes ^g Yes ^g Yes ^g
Atovaquone	Malaria, ^b babesiosis	<i>Frequent:</i> nausea, vomiting. <i>Occasional:</i> abdominal pain, headache	Plasma levels decreased by rifampin, tetracycline; bioavailability decreased by metoclopramide	C	No information

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Azoles Fluconazole Itraconazole Ketoconazole	Leishmaniasis	<i>Serious:</i> hepatotoxicity. <i>Rare:</i> exfoliative skin disorders, anaphylaxis	Warfarin, oral hypoglycemics, phenytoin, cyclosporine, theophylline, digoxin, dofetilide, quinidine, carbamazepine, rifabutin, busulfan, docetaxel, vinca alkaloids, pimozone, alprazolam, diazepam, midazolam, triazolam, verapamil, atorvastatin, cerivastatin, lovastatin, simvastatin, tacrolimus, sirolimus, indinavir, ritonavir, saquinavir, alfentanil, buspirone, methylprednisolone, trimetrexate: plasma levels increased by azoles Carbamazepine, phenobarbital, phenytoin, isoniazid, rifabutin, rifampin, antacids, H ₂ -receptor antagonists, proton pump inhibitors, nevirapine: decreased plasma levels of azoles Clarithromycin, erythromycin, indinavir, ritonavir: increased plasma levels of azoles	C	Yes
Benzimidazoles Albendazole	Ascariasis, capillariasis, clonorchiasis, cutaneous larva migrans, cysticercosis, ^b echinococcosis, ^b enterobiasis, eosinophilic enterocolitis, gnathostomiasis, hookworm, lymphatic filariasis, microsporidiosis, strongyloidiasis, trichinellosis, trichostrongyliasis, trichuriasis, visceral larva migrans	<i>Occasional:</i> nausea, vomiting, abdominal pain, headache, reversible alopecia, elevated aminotransferases. <i>Rare:</i> leukopenia, rash	Dexamethasone, praziquantel: plasma level of albendazole sulfoxide increased by ~50%	C	Yes ^d
Mebendazole	Ascariasis, ^b capillariasis, eosinophilic enterocolitis, enterobiasis, ^b hookworm, ^b trichinellosis, trichostrongyliasis, trichuriasis, ^b visceral larva migrans	<i>Occasional:</i> diarrhea, abdominal pain, elevated aminotransferases. <i>Rare:</i> agranulocytosis, thrombocytopenia, alopecia	Cimetidine: inhibited mebendazole metabolism	C	No information
Thiabendazole	Strongyloidiasis, ^b cutaneous larva migrans, ^b visceral larva migrans ^b	<i>Frequent:</i> anorexia, nausea, vomiting, diarrhea, headache, dizziness, asparagus-like urine odor. <i>Occasional:</i> drowsiness, giddiness, crystalluria, elevated aminotransferases, psychosis. <i>Rare:</i> hepatitis, seizures, angioneurotic edema, Stevens-Johnson syndrome, tinnitus	Theophylline: serum levels increased by thiabendazole	C	No information
Triclabendazole	Fascioliasis, paragonimiasis	<i>Occasional:</i> abdominal cramps, diarrhea, biliary colic, transient headache	No information	Not assigned	Yes
Benznidazole	Chagas' disease	<i>Frequent:</i> rash, pruritus, nausea, leukopenia, paresthesias	No major interactions	Not assigned	No information
Clindamycin	Babesiosis, malaria, toxoplasmosis	<i>Occasional:</i> pseudomembranous colitis, abdominal pain, diarrhea, nausea/vomiting. <i>Rare:</i> pruritus, skin rashes	No major interactions	B	Yes ^d
Diloxanide furoate	Amebiasis	<i>Frequent:</i> flatulence. <i>Occasional:</i> nausea, vomiting, diarrhea. <i>Rare:</i> pruritus	None reported	Contraindicated	No information
Eflornithine ^b (difluoromethylornithine, DFMO)	Trypanosomiasis	<i>Frequent:</i> pancytopenia. <i>Occasional:</i> diarrhea, seizures. <i>Rare:</i> transient hearing loss	No major interactions	Contraindicated	No information

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Emetine and dehydroemetine ^e	Amebiasis, fascioliasis	<i>Severe:</i> cardiotoxicity. <i>Frequent:</i> pain at injection site. <i>Occasional:</i> dizziness, headache, GI symptoms	None reported	X	No information
Folate antagonists					
Dihydrofolate reductase inhibitors					
Pyrimethamine	Malaria, ^b isosporiasis, toxoplasmosis ^b	<i>Occasional:</i> folate deficiency. <i>Rare:</i> rash, seizures, severe skin reactions (toxic epidermal necrolysis, erythema multiforme, Stevens-Johnson syndrome)	Sulfonamides, proguanil, zidovudine: increased risk of bone marrow suppression when used concomitantly	C	Yes
Proguanil and chlorproguanil	Malaria	<i>Occasional:</i> urticaria. <i>Rare:</i> hematuria, GI disturbances	No major interactions	C	Yes
Trimethoprim	Cyclosporiasis, isosporiasis	Hyperkalemia, GI upset, mild stomatitis	Methotrexate: reduced clearance Warfarin: effect prolonged Phenytoin: hepatic metabolism increased	C	Yes
Dihydropteroate synthetase inhibitors: sulfonamides					
Sulfadiazine	Malaria, ^b toxoplasmosis ^b	<i>Frequent:</i> GI disturbances, allergic skin reactions, crystalluria. <i>Rare:</i> severe skin reactions (toxic epidermal necrolysis, erythema multiforme, Stevens-Johnson syndrome), agranulocytosis, aplastic anemia, hypersensitivity of the respiratory tract, hepatitis, interstitial nephritis, hypoglycemia, aseptic meningitis	Thiazide diuretics: increased risk of thrombocytopenia in elderly patients Warfarin: effect prolonged by sulfonamides Methotrexate: levels increased by sulfonamides Phenytoin: metabolism impaired by sulfonamides Sulfonylureas: effect prolonged by sulfonamides	B	Yes
Sulfamethoxazole					
Sulfadoxine					
Dihydropteroate synthetase inhibitors: sulfones					
Dapsone	Leishmaniasis, malaria, toxoplasmosis	<i>Frequent:</i> rash, anorexia. <i>Occasional:</i> hemolysis, methemoglobinemia, neuropathy, allergic dermatitis, anorexia, nausea, vomiting, tachycardia, headache, insomnia, psychosis, hepatitis. <i>Rare:</i> agranulocytosis	Rifampin: lowered plasma levels of dapsone	C	Yes
Fumagillin	Microsporidiosis	<i>Rare:</i> neutropenia, thrombocytopenia	None reported	No information	No information
Furazolidone	Giardiasis	<i>Frequent:</i> nausea/vomiting, brown urine. <i>Occasional:</i> rectal itching, headache. <i>Rare:</i> hemolytic anemia, disulfiram-like reactions, MAO inhibitor interactions	Risk of hypertensive crisis when administered for >5 days with MAO inhibitors	C	No information
Iodoquinol	Amebiasis, ^b balantidiasis, <i>D. fragilis</i> infection	<i>Occasional:</i> headache, rash, pruritus, thyrotoxicosis, nausea, vomiting, abdominal pain, diarrhea. <i>Rare:</i> optic neuritis, peripheral neuropathy, seizures, encephalopathy	No major interactions	C	No information
Ivermectin	Ascariasis, cutaneous larva migrans, gnathostomiasis, loiasis, lymphatic filariasis, onchocerciasis, ^b scabies, strongyloidiasis, ^b trichuriasis	<i>Occasional:</i> fever, pruritus, headache, myalgias. <i>Rare:</i> hypotension	No major interactions	C	Yes ^d
Macrolides					
Azithromycin	Babesiosis	<i>Occasional:</i> nausea, vomiting, diarrhea, abdominal pain. <i>Rare:</i> angioedema, cholestatic jaundice	Cyclosporine and digoxin: levels increased by azithromycin Nelfinavir: increased levels of azithromycin	B	Yes

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Spiramycin ^b	Toxoplasmosis	<i>Occasional:</i> GI disturbances, transient skin eruptions. <i>Rare:</i> thrombocytopenia, QT prolongation in an infant, cholestatic hepatitis	No major interactions	Not assigned ^c	Yes ^d
Mefloquine	Malaria ^b	<i>Frequent:</i> lightheadedness, nausea, headache. <i>Occasional:</i> confusion; nightmares; insomnia; visual disturbance; transient and clinically silent ECG abnormalities, including sinus bradycardia, sinus arrhythmia, first-degree AV block, prolongation of QTc interval, and abnormal T waves. <i>Rare:</i> psychosis, convulsions, hypotension	Administration of halofantrine <3 weeks after mefloquine use may produce fatal QTc prolongation. Mefloquine may lower plasma levels of anticonvulsants. Levels are decreased and clearance is accelerated by artesunate.	C	Yes
Melarsoprol ^e	Trypanosomiasis	<i>Frequent:</i> myocardial injury, encephalopathy, peripheral neuropathy, hypertension. <i>Occasional:</i> G6PD-induced hemolysis, erythema nodosum leprosum. <i>Rare:</i> hypotension	No major interactions	Not assigned	No information
Metrifonate	Schistosomiasis	<i>Frequent:</i> abdominal pain, nausea, vomiting, diarrhea, headache, vertigo, bronchospasm. <i>Rare:</i> cholinergic symptoms	No major interactions	B	No
Miltefosine	Leishmaniasis ^b , primary amebic meningoencephalitis	<i>Frequent:</i> mild and transient (1–2 days) GI disturbances within first 2 weeks of therapy (resolve after treatment completion); motion sickness. <i>Occasional:</i> reversible elevations of creatinine and aminotransferases	No major interactions	Not assigned	No information
Niclosamide	Intestinal cestodes ^b	<i>Occasional:</i> nausea, vomiting, dizziness, pruritus	No major interactions	B	No information
Nifurtimox ^e	Chagas' disease	<i>Frequent:</i> nausea, vomiting, abdominal pain, insomnia, paresthesias, weakness, tremors. <i>Rare:</i> seizures (all reversible and dose-related)	No major interactions	Not assigned	No information
Nitazoxanide	Cryptosporidiosis, ^b giardiasis ^b	<i>Occasional:</i> abdominal pain, diarrhea. <i>Rare:</i> vomiting, headache	No major interactions	B	No information
Nitroimidazoles Metronidazole	Amebiasis, ^b balantidiasis, dracunculiasis, giardiasis, trichomoniasis, ^b <i>D. fragilis</i> infection	<i>Frequent:</i> nausea, headache, anorexia, metallic after-taste. <i>Occasional:</i> vomiting, insomnia, vertigo, paresthesias, disulfiram-like effects. <i>Rare:</i> seizures, peripheral neuropathy	Warfarin: effect enhanced by metronidazole Disulfiram: psychotic reaction Phenobarbital, phenytoin: accelerate elimination of metronidazole Lithium: serum levels elevated by metronidazole Cimetidine: prolonged half-life of metronidazole	B	Yes
Tinidazole	Amebiasis, ^b giardiasis, trichomoniasis	<i>Occasional:</i> nausea, vomiting, metallic taste	See metronidazole.	C	Yes
Oxamniquine	Schistosomiasis	<i>Occasional:</i> dizziness, drowsiness, headache, orange urine, elevated aminotransferases. <i>Rare:</i> seizures	No major interactions	C	No information
Paromomycin	Amebiasis, ^b <i>D. fragilis</i> infection, giardiasis, cryptosporidiosis, leishmaniasis	<i>Frequent:</i> GI disturbances (oral dosing only). <i>Occasional:</i> nephrotoxicity, ototoxicity, vestibular toxicity (parenteral dosing only)	No major interactions	Oral: B Parenteral: not assigned ^c	No information

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Pentamidine isethionate	Leishmaniasis, trypanosomiasis	<i>Frequent:</i> hypotension, hypoglycemia, pancreatitis, sterile abscesses at IM injection sites, GI disturbances, reversible renal failure. <i>Occasional:</i> hepatotoxicity, cardiotoxicity, delirium. <i>Rare:</i> anaphylaxis	No major interactions	C	No information
Piperazine and derivatives					
Piperazine	Ascariasis, enterobiasis	<i>Occasional:</i> nausea, vomiting, diarrhea, abdominal pain, headache. <i>Rare:</i> neurotoxicity, seizures	None reported	C	No information
Diethylcarbamazine ^e	Lymphatic filariasis, loiasis, tropical pulmonary eosinophilia	<i>Frequent:</i> dose-related nausea, vomiting. <i>Rare:</i> fever, chills, arthralgias, headaches	None reported	Not assigned ^d	No information
Praziquantel	Clonorchiasis, ^b cysticercosis, diphyllorbothriasis, hymenolepiasis, taeniasis, opisthorchiasis, intestinal trematodes, paragonimiasis, schistosomiasis ^b	<i>Frequent:</i> abdominal pain, diarrhea, dizziness, headache, malaise. <i>Occasional:</i> fever, nausea. <i>Rare:</i> pruritus, singultus	No major interactions	B	Yes
Pyrantel pamoate	Ascariasis, eosinophilic enterocolitis, enterobiasis, ^b hookworm, trichostrongyliasis	<i>Occasional:</i> GI disturbances, headache, dizziness, elevated aminotransferases	No major interactions	C	No information
Pyronaridine	Malaria	<i>Occasional:</i> headache, nausea	None reported to date	B	Yes
Quinacrine ^h	Giardiasis ^b	<i>Frequent:</i> headache, nausea, vomiting, bitter taste. <i>Occasional:</i> yellow-orange discoloration of skin, sclerae, urine; begins after 1 week of treatment and lasts up to 4 months after drug discontinuation. <i>Rare:</i> psychosis, exfoliative dermatitis, retinopathy, G6PD-induced hemolysis, exacerbation of psoriasis, disulfiram-like effects	Primaquine: toxicity potentiated by quinacrine	C	No information
Quinine and quinidine	Malaria, babesiosis	<i>Frequent:</i> cinchonism (tinnitus, high-tone deafness, headache, dysphoria, nausea, vomiting, abdominal pain, visual disturbances, postural hypotension), hyperinsulinemia resulting in life-threatening hypoglycemia. <i>Occasional:</i> deafness, hemolytic anemia, arrhythmias, hypotension due to rapid IV infusion	Carbonic anhydrase inhibitors, thiazide diuretics: reduced renal elimination of quinidine Amiodarone, cimetidine: increased quinidine levels Nifedipine: decreased quinidine levels; quinidine slows metabolism of nifedipine Phenobarbital, phenytoin, rifampin: accelerated hepatic elimination of quinidine Verapamil: reduced hepatic clearance of quinidine Diltiazem: decreased clearance of quinidine	X	Yes ^g
Quinolones					
Ciprofloxacin	Cyclosporiasis, isosporiasis	<i>Occasional:</i> nausea, diarrhea, vomiting, abdominal pain/discomfort, headache, restlessness, rash. <i>Rare:</i> myalgias/arthralgias, tendon rupture, CNS symptoms (nervousness, agitation, insomnia, anxiety, nightmares or paranoia); convulsions	Probenecid: increased serum levels of ciprofloxacin Theophylline, warfarin: serum levels increased by ciprofloxacin	C	Yes

(Continued)

TABLE 246e-1 OVERVIEW OF AGENTS USED FOR THE TREATMENT OF PARASITIC INFECTIONS (CONTINUED)

Drugs by Class	Parasitic Infection(s)	Adverse Effects	Major Drug-Drug Interactions	Pregnancy Class ^a	Breast Milk
Suramin ^f	Trypanosomiasis	<i>Frequent:</i> immediate: fever, urticaria, nausea, vomiting, hypotension; delayed (up to 24 h): exfoliative dermatitis, stomatitis, paresthesias, photophobia, renal dysfunction. <i>Occasional:</i> nephrotoxicity, adrenal toxicity, optic atrophy, anaphylaxis	No major interactions	Not assigned	No information
Tetracyclines	Balantidiasis, <i>D. fragilis</i> infection, malaria; lymphatic filariasis (doxycycline)	<i>Frequent:</i> GI disturbances. <i>Occasional:</i> photosensitivity dermatitis. <i>Rare:</i> exfoliative dermatitis, esophagitis, hepatotoxicity	Warfarin: effect prolonged by tetracyclines	D	Yes

^aBased on U.S. Food and Drug Administration pregnancy categories of A–D, X. ^bApproved by the FDA for this indication. ^cUse in pregnancy is recommended by international organizations outside the United States. ^dOnly AmBisome has been approved by the FDA for this indication. ^eAvailable through the CDC. ^fOnly artemether (in combination with lumefantrine) and artesunate have been approved by the FDA for this indication. ^gNot believed to be harmful. ^hAvailable through the manufacturer.

Abbreviations: ACTH, adrenocorticotropic hormone; AV, atrioventricular; CNS, central nervous system; ECG, electrocardiogram; G6PD, glucose 6-phosphate dehydrogenase; GI, gastrointestinal; MAO, monoamine oxidase.

through complexation with heme. It is rapidly absorbed and acts as a prodrug after oral administration; the principal plasma metabolite monodesethylamodiaquine is the predominant antimalarial agent. Amodiaquine and its metabolites are all excreted in urine, but there are no recommendations concerning dosage adjustment in patients with impaired renal function. Agranulocytosis and hepatotoxicity can develop with repeated use; therefore, this drug should not be used for prophylaxis. Despite widespread resistance, amodiaquine has been shown to be effective in some areas when combined with other antimalarial drugs (e.g., artesunate, sulfadoxine-pyrimethamine), particularly in children. Although licensed, amodiaquine is not yet available in the United States.

Amphotericin B See Table 246e-1 and Chap. 235.

Antimonials^g Despite associated adverse reactions and the need for prolonged parenteral treatment, the pentavalent antimonial compounds (designated Sb⁵⁺) have remained the first-line therapy for all forms of leishmaniasis throughout the world, primarily because they are affordable and effective and have survived the test of time. Pentavalent antimonials are active only after bio-reduction to the trivalent Sb(III) form, which inhibits trypanothione reductase, a critical enzyme involved in the oxidative stress management of *Leishmania* species. The fact that *Leishmania* species use trypanothione rather than glutathione (which is used by mammalian cells) may explain the parasite-specific activity of antimonials. The drugs are taken up by the reticuloendothelial system, and their activity against *Leishmania* species may be enhanced by this localization. Sodium stibogluconate is the only pentavalent antimonial available in the United States; meglumine antimoniate is used principally in francophone countries.

Resistance is a major problem in some areas. Although low-level unresponsiveness to Sb⁵⁺ was identified in India in the 1970s, incremental increases in both the recommended daily dosage (to 20 mg/kg) and the duration of treatment (to 28 days) satisfactorily compensated for the growing resistance until around 1990. There has since been steady erosion in the capacity of Sb⁵⁺ to induce long-term cure in patients with kala-azar who live in eastern India. Co-infection with HIV impairs the treatment response.

Sodium stibogluconate is available in aqueous solution and is administered parenterally. Antimony appears to have two elimination phases. When the drug is administered IV, the mean half-life of the first phase is <2 h; the mean half-life of the terminal elimination phase is nearly 36 h. This slower phase may be due to conversion of pentavalent antimony to a trivalent form that is the likely cause of the side effects often seen with prolonged therapy.

Artemisinin Derivatives^h Artesunate, artemether, arteether, and the parent compound artemisinin are sesquiterpene lactones derived from the

wormwood plant *Artemisia annua*. These agents are at least tenfold more potent in vivo than other antimalarial drugs and presently show no cross-resistance with known antimalarial drugs; thus, they have become first-line agents for the treatment of severe falciparum malaria. The artemisinin compounds are rapidly effective against the asexual blood forms of *Plasmodium* species but are not active against intrahepatic forms. Artemisinin and its derivatives are highly lipid soluble and readily cross both host and parasite cell membranes. One factor that explains the drugs' highly selective toxicity against malaria is that parasitized erythrocytes concentrate artemisinin and its derivatives to concentrations 100-fold higher than those in uninfected erythrocytes. The antimalarial effect of these agents results primarily from dihydroartemisinin, a compound to which artemether and artesunate are both converted. In the presence of heme or molecular iron, the endoperoxide moiety of dihydroartemisinin decomposes, generating free radicals and other metabolites that damage parasite proteins. The compounds are available for oral, rectal, IV, or IM administration, depending on the derivative. In the United States, IV artesunate is available for the treatment of severe, quinidine-unresponsive malaria through the CDC malaria hotline (770-488-7788, M–F, 0800–1630 EST; 770-488-7100 after hours). Artemisinin and its derivatives are cleared rapidly from the circulation. Their short half-lives limit their value for prophylaxis and monotherapy. These agents should be used only in combination with another, longer-acting agent (e.g., artesunate-mefloquine, dihydroartemisinin-piperaquine). A combined formulation of artemether and lumefantrine is available for the treatment of acute uncomplicated falciparum malaria acquired in areas where *Plasmodium falciparum* is resistant to chloroquine and antifolates.

Atovaquone Atovaquone is a hydroxynaphthoquinone that exerts broad-spectrum antiprotozoal activity via selective inhibition of parasite mitochondrial electron transport. This agent exhibits potent activity against toxoplasmosis and babesiosis when used with pyrimethamine and azithromycin, respectively. Atovaquone possesses a novel mode of action against *Plasmodium* species, inhibiting the electron transport system at the level of the cytochrome bc1 complex. The drug is active against both the erythrocytic and the exoerythrocytic stages of *Plasmodium* species; however, because it does not eradicate hypnozoites from the liver, patients with *Plasmodium vivax* or *Plasmodium ovale* infections must be given radical prophylaxis.

Malarone[®] is a fixed-dose combination of atovaquone and proguanil used for malaria prophylaxis as well as for the treatment of acute, uncomplicated *P. falciparum* malaria. Malarone has been shown to be effective in regions with multidrug-resistant *P. falciparum*. Resistance to atovaquone has yet to be reported.

The bioavailability of atovaquone varies considerably. Absorption after a single oral dose is slow, increases two- to threefold with a fatty

meal, and is dose-limited above 750 mg. The elimination half-life is increased in patients with moderate hepatic impairment. Because of the potential for drug accumulation, the use of atovaquone is generally contraindicated in persons with a creatinine clearance rate <30 mL/min. No dosage adjustments are needed in patients with mild to moderate renal impairment.

Azithromycin See Table 246e-1 and Chap. 170.

Azoles See Table 246e-1 and Chap. 235.

Benznidazole* This oral nitroimidazole derivative is used to treat Chagas' disease, with cure rates of 80–90% recorded in acute infections. Benznidazole is believed to exert its trypanocidal effects by generating oxygen radicals to which the parasites are more sensitive than mammalian cells because of a relative deficiency in antioxidant enzymes. Benznidazole also appears to alter the balance between pro- and anti-inflammatory mediators by downregulating the synthesis of nitrite, interleukin (IL) 6, and IL-10 in macrophages. Benznidazole is highly lipophilic and readily absorbed. The drug is extensively metabolized; only 5% of the dose is excreted unchanged in the urine. Benznidazole is well tolerated; adverse effects are rare and usually manifest as GI upset or pruritic rash.

Chloroquine This 4-aminoquinoline has marked, rapid schizonticidal and gametocidal activity against blood forms of *P. ovale* and *Plasmodium malariae* and against susceptible strains of *P. vivax* and *P. falciparum*. It is not active against intrahepatic forms (*P. vivax* and *P. ovale*). Parasitized erythrocytes accumulate chloroquine in significantly greater concentrations than do normal erythrocytes. Chloroquine, a weak base, concentrates in the food vacuoles of intraerythrocytic parasites because of a relative pH gradient between the extracellular space and the acidic food vacuole. Once it enters the acidic food vacuole, chloroquine is rapidly converted to a membrane-impermeable protonated form and is trapped. Continued accumulation of chloroquine in the parasite's acidic food vacuoles results in drug levels that are 600-fold higher at this site than in plasma. The high accumulation of chloroquine results in an increase in pH within the food vacuole to a level above that required for the acid proteases' optimal activity, inhibiting parasite heme polymerase; as a result, the parasite is effectively killed with its own metabolic waste. Compared with susceptible strains, chloroquine-resistant plasmodia transport chloroquine out of intraparasitic compartments more rapidly and maintain lower chloroquine concentrations in their acid vesicles. Hydroxychloroquine, a congener of chloroquine, is equivalent to chloroquine in its antimalarial efficacy but is preferred to chloroquine for the treatment of autoimmune disorders because it produces less ocular toxicity when used in high doses.

Chloroquine is well absorbed. However, because it exhibits extensive tissue binding, a loading dose is required to yield effective plasma concentrations. A therapeutic drug level in plasma is reached 2–3 h after oral administration (the preferred route). Chloroquine can be administered IV, but excessively rapid parenteral administration can result in seizures and death from cardiovascular collapse. The mean half-life of chloroquine is 4 days, but the rate of excretion decreases as plasma levels decline, making once-weekly administration possible for prophylaxis in areas with sensitive strains. About one-half of the parent drug is excreted in urine, but the dose should not be reduced for persons with acute malaria and renal insufficiency.

Ciprofloxacin See Table 246e-1 and Chap. 170.

Clindamycin See Table 246e-1 and Chap. 170.

Dapsone See Table 246e-1 and Chap. 205e.

Dehydroemetine Emetine is an alkaloid derived from ipecac; dehydroemetine is synthetically derived from emetine and is considered less toxic. Both agents are active against *Entamoeba histolytica* and appear to work by blocking peptide elongation and thus inhibiting protein synthesis. Emetine is rapidly absorbed after parenteral

administration, rapidly distributed throughout the body, and slowly excreted in the urine in unchanged form. Both agents are contraindicated in patients with renal disease.

Diethylcarbamazine* A derivative of the antihelminthic agent piperazine with a long history of successful use, diethylcarbamazine (DEC) remains the treatment of choice for lymphatic filariasis and loiasis and has also been used for visceral larva migrans. Although piperazine itself has no antifilarial activity, the piperazine ring of DEC is essential for the drug's activity. DEC's mechanism of action remains to be fully defined. Proposed mechanisms include immobilization due to inhibition of parasite cholinergic muscle receptors, disruption of microtubule formation, and alteration of helminthic surface membranes resulting in enhanced killing by the host's immune system. DEC enhances adherence properties of eosinophils. The development of resistance under drug pressure (i.e., a progressive decrease in efficacy when the drug is used widely in human populations) has not been observed, although DEC has variable effects when administered to persons with filariasis. Monthly administration provides effective prophylaxis against both bancroftian filariasis and loiasis.

DEC is well absorbed after oral administration, with peak plasma concentrations reached within 1–2 h. No parenteral form is available. The drug is eliminated largely by renal excretion, with <5% found in feces. If more than one dose is to be administered to an individual with renal dysfunction, the dose should be reduced commensurate with the reduction in creatinine clearance rate. Alkalinization of the urine prevents renal excretion and increases the half-life of DEC. Use in patients with onchocerciasis can precipitate a Mazzotti reaction, with pruritus, fever, and arthralgias. Like other piperazines, DEC is active against *Ascaris* species. Patients co-infected with this nematode may expel live worms after treatment.

Diloxanide Furoate Diloxanide furoate, a substituted acetanilide, is a luminally active agent used to eradicate the cysts of *E. histolytica*. After ingestion, diloxanide furoate is hydrolyzed by enzymes in the lumen or mucosa of the intestine, releasing furoic acid and the ester diloxanide; the latter acts directly as an amebicide.

Diloxanide furoate is given alone to asymptomatic cyst passers. For patients with active amebic infections, diloxanide is generally administered in combination with a 5-nitroimidazole such as metronidazole or tinidazole. Diloxanide furoate is rapidly absorbed after oral administration. When coadministered with a 5-nitroimidazole, only diloxanide appears in the systemic circulation; levels peak within 1 h and disappear within 6 h. About 90% of an oral dose is excreted in the urine within 48 h, chiefly as the glucuronide metabolite. Diloxanide furoate is contraindicated in pregnant and breast-feeding women and in children <2 years of age.

Eflornithine* Eflornithine (difluoromethylornithine, or DFMO) is a fluorinated analogue of the amino acid ornithine. Although originally designed as an antineoplastic agent, eflornithine has proven effective against some trypanosomatids. At one point, the production of this effective agent ceased despite the increasing incidence of human African trypanosomiasis; however, production resumed after eflornithine was discovered to be an effective cosmetic depilatory agent.

Eflornithine has specific activity against all stages of infection with *Trypanosoma brucei gambiense*; however, it is inactive against *Trypanosoma brucei rhodesiense*. The drug acts as an irreversible suicide inhibitor of ornithine decarboxylase, the first enzyme in the biosynthesis of the polyamines putrescine and spermidine. Polyamines are essential for the synthesis of trypanothione, an enzyme required for the maintenance of intracellular thiols in the correct redox state and for the removal of reactive oxygen metabolites. However, polyamines are also essential for cell division in eukaryotes, and ornithine decarboxylase is similar in trypanosomes and mammals. The selective antiparasitic activity of eflornithine is partly explained by the structure of the trypanosomal enzyme, which lacks a 36-amino-acid C-terminal sequence found on mammalian ornithine decarboxylase. This difference results in a lower turnover of ornithine decarboxylase and a more rapid decrease of polyamines in trypanosomes than in the mammalian host.

The diminished effectiveness of eflornithine against *T. b. rhodesiense* appears to be due to the parasite's ability to replace the inhibited enzyme more rapidly than *T. b. gambiense*.

Eflornithine is less toxic but more costly than conventional therapy. It can be administered IV or PO. The dose should be reduced in renal failure. Eflornithine readily crosses the blood-brain barrier; CSF levels are highest in persons with the most severe central nervous system (CNS) involvement.

Fumagillin^f Fumagillin is a water-insoluble antibiotic that is derived from the fungus *Aspergillus fumigatus* and is active against microsporidia. This drug is effective when used topically to treat ocular infections due to *Encephalitozoon* species. When given systemically, fumagillin was effective but caused thrombocytopenia in all recipients in the second week of treatment; this side effect was readily reversed when administration of the drug was stopped. The mechanisms by which fumagillin inhibits microsporidial replication are poorly understood, although the drug may inhibit methionine aminopeptidase 2 by irreversibly blocking the active site.

Furazolidone This nitrofurantoin derivative is an effective alternative agent for the treatment of giardiasis and also exhibits activity against *Isoospora belli*. Because it is the only agent active against *Giardia* that is available in liquid form, it is most often used to treat young children. Furazolidone undergoes reductive activation in *Giardia lamblia* trophozoites—an event that, unlike the reductive activation of metronidazole, involves an NADH oxidase. The killing effect correlates with the toxicity of reduced products, which damage important cellular components, including DNA. Although furazolidone had been thought to be largely unabsorbed when administered orally, the occurrence of systemic adverse reactions indicates that this is not the case. More than 65% of the drug dose can be recovered from the urine as colored metabolites. Omeprazole reduces the oral bioavailability of furazolidone.

Furazolidone is a monoamine oxidase (MAO) inhibitor; thus, caution should be used in its concomitant administration with other drugs (especially indirectly acting sympathomimetic amines) and in the consumption of food and drink containing tyramine during treatment. However, hypertensive crises have not been reported in patients receiving furazolidone, and it has been suggested that—because furazolidone inhibits MAOs gradually over several days—the risks are small if treatment is limited to a 5-day course. Because hemolytic anemia can occur in patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency and glutathione instability, furazolidone treatment is contraindicated in mothers who are breast-feeding and in neonates.

Halofantrine This 9-phenanthrenemethanol is one of three classes of arylaminoalcohols first identified as potential antimalarial agents by the World War II Malaria Chemotherapy Program. Its activity is believed to be similar to that of chloroquine, although it is an oral alternative for the treatment of malaria due to chloroquine-resistant *P. falciparum*.

Although the mechanism of action is poorly understood, halofantrine is thought to share mechanism(s) with the 4-aminoquinolines, forming a complex with ferriprotoporphyrin IX and interfering with the degradation of hemoglobin.

Halofantrine exhibits erratic bioavailability, but its absorption is significantly enhanced when it is taken with a fatty meal. The elimination half-life of halofantrine is 1–2 days; it is excreted mainly in feces. Halofantrine is metabolized into *N*-debutyl-halofantrine by the cytochrome P450 enzyme CYP3A4. Grapefruit juice should be avoided during treatment because it increases both halofantrine's bioavailability and halofantrine-induced QT interval prolongation by inhibiting CYP3A4 at the enterocyte level.

Iodoquinol Iodoquinol (diiodohydroxyquin), a hydroxyquinoline, is an effective luminal agent for the treatment of amebiasis, balantidiasis, and infection with *Dientamoeba fragilis*. Its mechanism of action is unknown. It is poorly absorbed. Because the drug contains 64% organically bound iodine, it should be used with caution in patients

with thyroid disease. Iodine dermatitis occurs occasionally during iodoquinol treatment. Protein-bound serum iodine levels may be increased during treatment and can interfere with certain tests of thyroid function. These effects may persist for as long as 6 months after discontinuation of therapy. Iodoquinol is contraindicated in patients with liver disease. Most serious are the reactions related to prolonged high-dose therapy (optic neuritis, peripheral neuropathy), which should not occur if the recommended dosage regimens are followed.

Ivermectin Ivermectin (22,23-dihydroavermectin) is a derivative of the macrocyclic lactone avermectin produced by the soil-dwelling actinomycete *Streptomyces avermitilis*. Ivermectin is active at low doses against a wide range of helminths and ectoparasites. It is the drug of choice for the treatment of onchocerciasis, strongyloidiasis, cutaneous larva migrans, and scabies. Ivermectin is highly active against microfilariiae of the lymphatic filariases but has no macrofilaricidal activity. When ivermectin is used in combination with other agents such as DEC or albendazole for treatment of lymphatic filariasis, synergistic activity is seen. Although active against the intestinal helminths *Ascaris lumbricoides* and *Enterobius vermicularis*, ivermectin is only variably effective in trichuriasis and is ineffective against hookworms. Widespread use of ivermectin for treatment of intestinal nematode infections in sheep and goats has led to the emergence of drug resistance in veterinary practice; this development may portend problems in human medical use.

Data suggest that ivermectin acts by opening the neuromuscular membrane-associated, glutamate-dependent chloride channels. The influx of chloride ions results in hyperpolarization and muscle paralysis—particularly of the nematode pharynx, with consequent blockage of the oral ingestion of nutrients. As these chloride channels are present only in invertebrates, paralysis is seen only in the parasite.

Ivermectin is available for administration to humans only as an oral formulation. The drug is highly protein bound; it is almost completely excreted in feces. Both food and beer increase the bioavailability of ivermectin significantly. Ivermectin is distributed widely throughout the body; animal studies indicate that it accumulates at the highest concentration in adipose tissue and liver, with little accumulation in the brain. Few data exist to guide therapy in hosts with conditions that may influence drug pharmacokinetics.

Ivermectin is generally administered as a single dose of 150–200 µg/kg. In the absence of parasitic infection, the adverse effects of ivermectin in therapeutic doses are minimal. Adverse effects in patients with filarial infections include fever, myalgia, malaise, lightheadedness, and (occasionally) postural hypotension. The severity of such side effects is related to the intensity of parasite infection, with more symptoms in individuals with a heavy parasite burden. In onchocerciasis, skin edema, pruritus, and mild eye irritation may also occur. The adverse effects are generally self-limiting and only occasionally require symptom-based treatment with antipyretics or antihistamines. More severe complications of ivermectin therapy for onchocerciasis include encephalopathy in patients heavily infected with *Loa loa*.

Lumefantrine Lumefantrine (benflumetol), a fluorene arylaminoalcohol derivative synthesized in the 1970s by the Chinese Academy of Military Medical Sciences (Beijing), has marked blood schizonticidal activity against a wide range of plasmodia. This agent conforms structurally and in mode of action to other arylaminoalcohols (quinine, mefloquine, and halofantrine). Lumefantrine exerts its antimalarial effect as a consequence of its interaction with heme, a degradation product of hemoglobin metabolism. Although its antimalarial activity is slower than that of the artemisinin-based drugs, the recrudescence rate with the recommended lumefantrine regimen is lower. The pharmacokinetic properties of lumefantrine are reminiscent of those of halofantrine, with variable oral bioavailability, considerable augmentation of oral bioavailability by concomitant fat intake, and a terminal elimination half-life of ~4–5 days in patients with malaria.

Artemether and lumefantrine have synergistic activity, and the combined formulation of artemether and lumefantrine is effective for the treatment of falciparum malaria in areas where *P. falciparum* is resistant to chloroquine and antifolates.

246e-10 Mebendazole This benzimidazole is a broad-spectrum antiparasitic agent widely used to treat intestinal helminthiases. Its mechanism of action is similar to that of albendazole; however, it is a more potent inhibitor of parasite malic dehydrogenase and exhibits a more specific and selective effect against intestinal nematodes than the other benzimidazoles.

Mebendazole is available only in oral form but is poorly absorbed from the GI tract; only 5–10% of a standard dose is measurable in plasma. The proportion absorbed from the GI tract is extensively metabolized in the liver. Metabolites appear in the urine and bile; impaired liver or biliary function results in higher plasma mebendazole levels in treated patients. No dose reduction is warranted in patients with renal function impairment. Because mebendazole is poorly absorbed, its incidence of side effects is low. Transient abdominal pain and diarrhea sometimes occur, usually in persons with massive parasite burdens.

Mefloquine Mefloquine is the preferred drug for prophylaxis of chloroquine-resistant malaria; high doses can be used for treatment. Despite the development of drug-resistant strains of *P. falciparum* in parts of Africa and Southeast Asia, mefloquine is an effective drug throughout most of the world. Cross-resistance of mefloquine with halofantrine and with quinine has been documented in limited areas. Like quinine and chloroquine, this quinoline is active only against the asexual erythrocytic stages of malarial parasites. Unlike quinine, however, mefloquine has a relatively poor affinity for DNA and, as a result, does not inhibit the synthesis of parasitic nucleic acids and proteins. Although both mefloquine and chloroquine inhibit hemozoin formation and heme degradation, mefloquine differs in that it forms a complex with heme that may be toxic to the parasite.

Mefloquine HCl is poorly water soluble and intensely irritating when given parenterally; thus it is available only in tablet form. Its absorption is adversely affected by vomiting and diarrhea but is significantly enhanced when the drug is administered with or after food. About 98% of the drug binds to protein. Mefloquine is excreted mainly in the bile and feces; therefore, no dose adjustment is needed in persons with renal insufficiency. The drug and its main metabolite are not appreciably removed by hemodialysis. No special chemoprophylactic dosage adjustments are indicated for the achievement of plasma concentrations in dialysis patients that are similar to those in healthy persons. Pharmacokinetic differences have been detected among various ethnic populations. In practice, however, these distinctions are of minor importance compared with host immune status and parasite sensitivity. In patients with impaired liver function, the elimination of mefloquine may be prolonged, leading to higher plasma levels.

Mefloquine should be used with caution by individuals participating in activities requiring alertness and fine-motor coordination. A recent FDA review found that dizziness, vertigo, or tinnitus can persist or become permanent as a result of treatment with the drug; thus, a boxed warning was mandated. If the drug is to be administered for a prolonged period, periodic evaluations are recommended, including liver function tests and ophthalmic examinations. Sleep abnormalities (insomnia, abnormal dreams) have occasionally been reported. Psychosis and seizures occur rarely; mefloquine should not be prescribed to patients with neuropsychiatric conditions, including depression, generalized anxiety disorder, psychosis, schizophrenia, and seizure disorder. If acute anxiety, depression, restlessness, or confusion develops during prophylaxis, these psychiatric symptoms may be considered prodromal to a more serious event, and the drug should be discontinued.

Concomitant use of quinine, quinidine, or drugs producing β -adrenergic blockade may cause significant electrocardiographic abnormalities or cardiac arrest. Halofantrine must not be given simultaneously with or <3 weeks after mefloquine because a potentially fatal prolongation of the QTc interval on electrocardiography may occur. No data exist on mefloquine use after halofantrine use. Administration of mefloquine with quinine or chloroquine may increase the risk of convulsions. Mefloquine may lower plasma levels of anticonvulsants. Caution should be exercised with regard to concomitant antiretroviral

therapy, since mefloquine has been shown to exert variable effects on ritonavir pharmacokinetics that are not explained by hepatic CYP3A4 activity or ritonavir protein binding. Vaccinations with attenuated live bacteria should be completed at least 3 days before the first dose of mefloquine.

Women of childbearing age who are traveling to areas where malaria is endemic should be warned against becoming pregnant and encouraged to practice contraception during malaria prophylaxis with mefloquine and for up to 3 months thereafter. However, in the case of unplanned pregnancy, use of mefloquine is not considered an indication for pregnancy termination. Analysis of prospectively monitored cases demonstrates a prevalence of birth defects and fetal loss comparable to background rates.

Melarsoprol Melarsoprol has been used since 1949 for the treatment of human African trypanosomiasis. This trivalent arsenical compound is indicated for the treatment of African trypanosomiasis with neurologic involvement and for the treatment of early disease that is resistant to suramin or pentamidine. Melarsoprol, like other drugs containing heavy metals, interacts with thiol groups of several different proteins; however, its antiparasitic effects appear to be more specific. Trypanothione reductase is a key enzyme involved in the oxidative stress management of both *Trypanosoma* and *Leishmania* species, helping to maintain an intracellular reducing environment by reduction of disulfide trypanothione to its dithiol derivative dihydrotrypanothione. Melarsoprol sequesters dihydrotrypanothione, depriving the parasite of its main sulfhydryl antioxidant, and inhibits trypanothione reductase, depriving the parasite of the essential enzyme system that is responsible for keeping trypanothione reduced. These effects are synergistic. The selectivity of arsenical action against trypanosomes is due at least in part to the greater melarsoprol affinity of reduced trypanothione than of other monothiols (e.g., cysteine) on which the mammalian host depends for maintenance of high thiol levels. Melarsoprol enters the parasite via an adenosine transporter; drug-resistant strains lack this transport system.

Melarsoprol is always administered IV. A small but therapeutically significant amount of the drug enters the CSF. The compound is excreted rapidly, with ~80% of the arsenic found in feces.

Melarsoprol is highly toxic. The most serious adverse reaction is reactive encephalopathy, which affects 6% of treated individuals and usually develops within 4 days of the start of therapy, with an average case-fatality rate of 50%. Glucocorticoids are administered with melarsoprol to prevent this development. Because melarsoprol is intensely irritating, care must be taken to avoid infiltration of the drug.

Metrifonate Metrifonate has selective activity against *Schistosoma haematobium*. This organophosphorous compound is a prodrug that is converted nonenzymatically to dichlorvos (2,2-dichlorovinyl dimethylphosphate, DDVP), a highly active chemical that irreversibly inhibits the acetylcholinesterase enzyme. Schistosomal cholinesterase is more susceptible to dichlorvos than is the corresponding human enzyme. The exact mechanism of action of metrifonate is uncertain, but the drug is believed to inhibit tegumental acetylcholine receptors that mediate glucose transport.

Metrifonate is administered in a series of three doses at 2-week intervals. After a single oral dose, metrifonate produces a 95% decrease in plasma cholinesterase activity within 6 h, with a fairly rapid return to normal. However, 2.5 months are required for erythrocyte cholinesterase levels to return to normal. Treated persons should not be exposed to neuromuscular blocking agents or organophosphate insecticides for at least 48 h after treatment.

Metronidazole and Other Nitroimidazoles See Table 246e-1 and Chap. 170.

Miltefosine In the early 1990s, miltefosine (hexadecylphosphocholine), originally developed as an antineoplastic agent, was discovered to have significant antiproliferative activity against *Leishmania* species, *Trypanosoma cruzi*, and *T. brucei* parasites in vitro and in experimental animal models. Miltefosine is the first oral drug that has proved to be highly effective and comparable to amphotericin B against visceral

leishmaniasis in India, where antimonial-resistant cases are prevalent. Miltefosine is also effective in previously untreated visceral infections. Cure rates in cutaneous leishmaniasis are comparable to those obtained with antimony. Miltefosine has also been shown to be effective against the free-living amoeba *Naegleria fowleri*.

The activity of miltefosine is attributed to interaction with cell signal transduction pathways and inhibition of phospholipid and sterol biosynthesis. Resistance to miltefosine has not been observed clinically. The drug is readily absorbed from the GI tract, is widely distributed, and accumulates in several tissues. The efficacy of a 28-day treatment course in Indian visceral leishmaniasis is equivalent to that of amphotericin B therapy; however, it appears that a shortened course of 21 days may be equally efficacious.

General recommendations for the use of miltefosine are limited by the exclusion of specific groups from the published clinical trials: persons <12 or >65 years of age, persons with the most advanced disease, breast-feeding women, HIV-infected patients, and individuals with significant renal or hepatic insufficiency.

Niclosamide[†] Niclosamide is active against a wide variety of adult tapeworms but not against tissue cestodes. It is also a molluscicide and is used in snail-control programs. The drug uncouples oxidative phosphorylation in parasite mitochondria, thereby blocking the uptake of glucose by the intestinal tapeworm and resulting in the parasite's death. Niclosamide rapidly causes spastic paralysis of intestinal cestodes in vitro. Its use is limited by its side effects, the necessarily long duration of therapy, the recommended use of purgatives, and—most important—limited availability (i.e., on a named-patient basis from the manufacturer).

Niclosamide is poorly absorbed. Tablets are given on an empty stomach in the morning after a liquid meal the night before, and this dose is followed by another 1 h later. For treatment of hymenolepiasis, the drug is administered for 7 days. A second course is often prescribed. The scolex and proximal segments of the tapeworms are killed on contact with niclosamide and may be digested in the gut. However, disintegration of the adult tapeworm results in the release of viable ova, which theoretically can result in autoinfection. Although fears of the development of cysticercosis in patients with *Taenia solium* infections have proved unfounded, it is still recommended that a brisk purgative be given 2 h after the first dose.

Nifurtimox^{*} This nitrofurantoin compound is an inexpensive and effective oral agent for the treatment of acute Chagas' disease. Trypanosomes lack catalase and have very low levels of peroxidase; as a result, they are very vulnerable to by-products of oxygen reduction. When nifurtimox is reduced in the trypanosome, a nitro anion radical is formed and undergoes autooxidation, resulting in the generation of the superoxide anion O_2^- , hydrogen peroxide (H_2O_2), hydroperoxyl radical (HO_2), and other highly reactive and cytotoxic molecules. Despite the abundance of catalases, peroxidases, and superoxide dismutases that neutralize these destructive radicals in mammalian cells, nifurtimox has a poor therapeutic index. Prolonged use is required, but the course may have to be interrupted because of drug toxicity, which develops in 40–70% of recipients. Nifurtimox is well absorbed and undergoes rapid and extensive biotransformation; <0.5% of the original drug is excreted in urine.

Nitazoxanide Nitazoxanide is a 5-nitrothiazole compound used for the treatment of cryptosporidiosis and giardiasis; it is active against other intestinal protozoa as well. The drug is approved for use in children 1–11 years of age.

The antiprotozoal activity of nitazoxanide is believed to be due to interference with the pyruvate-ferredoxin oxidoreductase (PFOR) enzyme-dependent electron transfer reaction that is essential to anaerobic energy metabolism. Studies have shown that the PFOR enzyme from *G. lamblia* directly reduces nitazoxanide by transfer of electrons in the absence of ferredoxin. The DNA-derived PFOR protein sequence of *Cryptosporidium parvum* appears to be similar to that of *G. lamblia*. Interference with the PFOR enzyme-dependent electron

transfer reaction may not be the only pathway by which nitazoxanide exerts antiprotozoal activity.

After oral administration, nitazoxanide is rapidly hydrolyzed to an active metabolite, tizoxanide (desacetyl-nitazoxanide). Tizoxanide then undergoes conjugation, primarily by glucuronidation. It is recommended that nitazoxanide be taken with food; however, no studies have been conducted to determine whether the pharmacokinetics of tizoxanide and tizoxanide glucuronide differ in fasted versus fed subjects. Tizoxanide is excreted in urine, bile, and feces, and tizoxanide glucuronide is excreted in urine and bile. The pharmacokinetics of nitazoxanide in patients with impaired hepatic and/or renal function have not been studied. Tizoxanide is highly bound to plasma protein (>99.9%). Therefore, caution should be used when administering this agent concurrently with other highly plasma protein-bound drugs with narrow therapeutic indices, as competition for binding sites may occur.

Oxamniquine This tetrahydroquinoline derivative is an effective alternative agent for the treatment of *Schistosoma mansoni*, although susceptibility to this drug exhibits regional variation. Oxamniquine exhibits anticholinergic properties, but its primary mode of action seems to rely on ATP-dependent enzymatic drug activation generating an intermediate that alkylates essential macromolecules, including DNA. In treated adult schistosomes, oxamniquine produces marked tegumental alterations that are similar to those seen with praziquantel but that develop less rapidly, becoming evident 4–8 days after treatment.

Oxamniquine is administered orally as a single dose and is well absorbed. Food retards absorption and reduces bioavailability. About 70% of an administered dose is excreted in urine as a mixture of pharmacologically inactive metabolites. Patients should be warned that their urine might have an intense orange-red color. Side effects are uncommon and usually mild, although hallucinations and seizures have been reported.

Paromomycin (Aminosidine) First isolated in 1956, this aminoglycoside is an effective oral agent for the treatment of infections due to intestinal protozoa. Parenteral paromomycin appears to be effective against visceral leishmaniasis in India.

Paromomycin inhibits protozoan protein synthesis by binding to the 30S ribosomal RNA in the aminoacyl-tRNA site, causing misreading of mRNA codons. Paromomycin is less active against *G. lamblia* than standard agents; however, like other aminoglycosides, paromomycin is poorly absorbed from the intestinal lumen, and the high levels of drug in the gut compensate for this relatively weak activity. If absorbed or administered systemically, paromomycin can cause ototoxicity and nephrotoxicity. However, systemic absorption is very limited, and toxicity should not be a concern in persons with normal kidneys. Topical formulations are not generally available.

Pentamidine isethionate This diamidine is an effective alternative agent for some forms of leishmaniasis and trypanosomiasis. It is available for parenteral and aerosolized administration. Although its mechanism of action remains undefined, it is known to exert a wide range of effects, including interaction with trypanosomal kinetoplast DNA; interference with polyamine synthesis by a decrease in the activity of ornithine decarboxylase; and inhibition of RNA polymerase, ribosomal function, and the synthesis of nucleic acids and proteins.

Pentamidine isethionate is well absorbed, is highly tissue bound, and is excreted slowly over several weeks, with an elimination half-life of 12 days. No steady-state plasma concentration is attained in persons given daily injections; the result is extensive accumulation of pentamidine in tissues, primarily the liver, kidney, adrenal, and spleen. Pentamidine does not penetrate well into the CNS. Pulmonary concentrations of pentamidine are increased when the drug is delivered in aerosolized form.

Piperazine The antihelminthic activity of piperazine is confined to ascariasis and enterobiasis. Piperazine acts as an agonist at extrasynaptic γ -aminobutyric acid (GABA) receptors, causing an influx of chloride ions in the nematode somatic musculature. Although the initial result is hyperpolarization of the muscle fibers, the ultimate effect is flaccid

246e-12 paralysis, leading to the expulsion of live worms. Patients should be warned, as this occurrence can be unsettling.

Praziquantel This heterocyclic pyrazinoisoquinoline derivative is highly active against a broad spectrum of trematodes and cestodes. It is the mainstay of treatment for schistosomiasis and is a critical part of community-based control programs.

All of the effects of praziquantel can be attributed either directly or indirectly to an alteration of intracellular calcium concentrations. Although the exact mechanism of action remains unclear, the major mechanism is disruption of the parasite tegument, causing tetanic contractures with loss of adherence to host tissues and, ultimately, disintegration or expulsion. Praziquantel induces changes in the antigenicity of the parasite by causing the exposure of concealed antigens. Praziquantel also produces alterations in schistosomal glucose metabolism, including decreases in glucose uptake, lactate release, glycogen content, and ATP levels.

Praziquantel exerts its parasitic effects directly and does not need to be metabolized to be effective. It is well absorbed but undergoes extensive first-pass hepatic clearance. Levels of the drug are increased when it is taken with food, particularly carbohydrates, or with cimetidine. Serum levels are reduced by glucocorticoids, chloroquine, carbamazepine, and phenytoin. Praziquantel is completely metabolized in humans, with 80% of the dose recovered as metabolites in urine within 4 days. It is not known to what extent praziquantel crosses the placenta, but retrospective studies suggest that it is safe in pregnancy.

Patients with schistosomiasis who have heavy parasite burdens may develop abdominal discomfort, nausea, headache, dizziness, and drowsiness. Symptoms begin 30 min after ingestion, may require spasmolytics for relief, and usually disappear spontaneously after a few hours.

Primaquine Phosphate Primaquine, an 8-aminoquinoline, has a broad spectrum of activity against all stages of plasmodial development in humans but has been used most effectively for eradication of the hepatic stage of these parasites. Despite its toxicity, it remains the drug of choice for radical cure of *P. vivax* infections. Primaquine must be metabolized by the host to be effective. It is, in fact, rapidly metabolized; only a small fraction of the dose of the parent drug is excreted unchanged. Although the parasitocidal activity of the three oxidative metabolites remains unclear, they are believed to affect both pyrimidine synthesis and the mitochondrial electron transport chain. The metabolites appear to have significantly less antimalarial activity than primaquine; however, their hemolytic activity is greater than that of the parent drug.

Primaquine causes marked hypotension after parenteral administration and therefore is given only by the oral route. It is rapidly and almost completely absorbed from the GI tract.

Patients should be tested for G6PD deficiency before they receive primaquine. The drug may induce the oxidation of hemoglobin into methemoglobin, irrespective of the G6PD status of the patient. Primaquine is otherwise well tolerated.

Proguanil (Chloroguanide) Proguanil inhibits plasmodial dihydrofolate reductase and is used with atovaquone for oral treatment of uncomplicated malaria or with chloroquine for malaria prophylaxis in parts of Africa without widespread chloroquine-resistant *P. falciparum*.

Proguanil exerts its effect primarily by means of the metabolite cycloguanil, whose inhibition of dihydrofolate reductase in the parasite disrupts deoxythymidylate synthesis, thus interfering with a key pathway involved in the biosynthesis of pyrimidines required for nucleic acid replication. There are no clinical data indicating that folate supplementation diminishes drug efficacy; women of childbearing age for whom atovaquone/proguanil is prescribed should continue taking folate supplements to prevent neural tube birth defects.

Proguanil is extensively absorbed regardless of food intake. The drug is 75% protein bound. The main routes of elimination are hepatic biotransformation and renal excretion; 40–60% of the proguanil dose is excreted by the kidneys. Drug levels are increased and elimination is impaired in patients with hepatic insufficiency.

Pyrantel Pamoate Pyrantel is a tetrahydropyrimidine formulated as pamoate. This safe, well-tolerated, inexpensive drug is used to treat a variety of intestinal nematode infections but is ineffective in trichuriasis. Pyrantel pamoate is usually effective in a single dose. Its target is the nicotinic acetylcholine receptor on the surface of nematode somatic muscle. Pyrantel depolarizes the neuromuscular junction of the nematode, resulting in its irreversible paralysis and allowing the natural expulsion of the worm.

Pyrantel pamoate is poorly absorbed from the intestine; >85% of the dose is passed unaltered in feces. The absorbed portion is metabolized and excreted in urine. Piperazine is antagonistic to pyrantel pamoate and should not be used concomitantly.

Pyrantel pamoate has minimal toxicity at the oral doses used to treat intestinal helminthic infection. It is not recommended for pregnant women or for children <12 months old.

Pyrimethamine When combined with short-acting sulfonamides, this diaminopyrimidine is effective in malaria, toxoplasmosis, and isosporiasis. Unlike mammalian cells, the parasites that cause these infections cannot use preformed pyrimidines obtained through salvage pathways but rather rely completely on de novo pyrimidine synthesis, for which folate derivatives are essential cofactors. The efficacy of pyrimethamine is increasingly limited by the development of resistant strains of *P. falciparum* and *P. vivax*. Single amino acid substitutions to parasite dihydrofolate reductase confer resistance to pyrimethamine by decreasing the enzyme's binding affinity for the drug.

Pyrimethamine is well absorbed; the drug is 87% bound to human plasma proteins. In healthy volunteers, drug concentrations remain at therapeutic levels for up to 2 weeks; drug levels are lower in patients with malaria.

At the usual dosage, pyrimethamine alone causes little toxicity except for occasional skin rashes and, more rarely, blood dyscrasias. Bone marrow suppression sometimes occurs at the higher doses used for toxoplasmosis; at these doses, the drug should be administered with folic acid.

Pyronaridine This potent antimalarial is a benzonaphthyridine derivative first synthesized by Chinese researchers in 1970. Like chloroquine, pyronaridine targets heme formation, inhibiting the production of β -heme by forming complexes with it, with consequent enhancement of heme-induced hemolysis. However, this drug is more potent than chloroquine: for complete lysis, pyronaridine is required at only 1/100th of the concentration needed with chloroquine. It also inhibits glutathione-dependent heme degradation. Despite its similar mode of action, pyronaridine remains effective against chloroquine-resistant strains. When combined with artesunate, it has been shown to be effective for the treatment of acute, uncomplicated infection caused by *P. falciparum* or *P. vivax* in areas of low transmission with evidence of artemisinin resistance.

Pyronaridine is readily absorbed, widely distributed throughout the body, metabolized by the liver, and excreted in urine and stool. Its use is contraindicated in patients with severe liver or kidney impairment. Pyronaridine has been shown in vitro to be an inhibitor of both CYP2D6 and P-glycoprotein, and these effects may have clinical relevance for patients taking medications for cardiac disease (e.g., metoprolol and digoxin).

Quinacrine* Quinacrine is the only drug approved by the FDA for the treatment of giardiasis. Although its production was discontinued in 1992, quinacrine can be obtained from alternative sources through the CDC Drug Service. The antiprotozoal mechanism of quinacrine has not been fully elucidated. The drug inhibits NADH oxidase—the same enzyme that activates furazolidone. The differing relative quinacrine uptake rate between human cells and *G. lamblia* may explain the selective toxicity of the drug. Resistance correlates with decreased drug uptake.

Quinacrine is rapidly absorbed from the intestinal tract and is widely distributed in body tissues. Alcohol is best avoided due to a disulfiram-like effect.

Quinine and Quinidine When combined with another agent, the cinchona alkaloid quinine is effective for the oral treatment of both uncomplicated, chloroquine-resistant malaria and babesiosis. Quinine acts rapidly against the asexual blood stages of all forms of the human malaria parasites. For severe malaria, only quinidine (the dextroisomer of quinine) is available in the United States. Quinine concentrates in the acidic food vacuoles of *Plasmodium* species. The drug inhibits the nonenzymatic polymerization of the highly reactive, toxic heme molecule into the nontoxic polymer pigment hemozoin.

Quinine is readily absorbed when given orally. In patients with malaria, the elimination half-life of quinine increases according to the severity of the infection. However, toxicity is avoided by an increase in the concentration of plasma glycoproteins. The cinchona alkaloids are extensively metabolized, particularly by CYP3A4; only 20% of the dose is excreted unchanged in urine. The drug's metabolites are also excreted in urine and may be responsible for toxicity in patients with renal failure. Renal excretion of quinine is decreased when cimetidine is taken and increased when the urine is acidic. The drug readily crosses the placenta.

Quinidine is both more potent as an antimalarial and more toxic than quinine. Its use requires cardiac monitoring. Dose reduction is necessary in persons with severe renal impairment.

Spiramycin[†] This macrolide is used to treat acute toxoplasmosis in pregnancy and congenital toxoplasmosis. While the mechanism of action is similar to that of other macrolides, the efficacy of spiramycin in toxoplasmosis appears to stem from its rapid and extensive intracellular penetration, which results in macrophage drug concentrations 10–20 times greater than serum concentrations.

Spiramycin is rapidly and widely distributed throughout the body and reaches concentrations in the placenta up to five times those in serum. This agent is excreted mainly in bile. Indeed, in humans, the urinary excretion of active compounds represents only 20% of the administered dose.

Serious reactions to spiramycin are rare. Of the available macrolides, spiramycin appears to have the lowest risk of drug interactions. Complications of treatment are rare but, in neonates, can include life-threatening ventricular arrhythmias that disappear with drug discontinuation.

Sulfonamides See Table 246e-1 and Chap. 170.

Suramin[†] This derivative of urea is the drug of choice for the early stage of African trypanosomiasis. The drug is polyanionic and acts by forming stable complexes with proteins, thus inhibiting multiple enzymes essential to parasite energy metabolism. Suramin appears to inhibit all trypanosome glycolytic enzymes more effectively than it inhibits the corresponding host enzymes.

Suramin is parenterally administered. It binds to plasma proteins and persists at low levels for several weeks after infusion. Its metabolism is negligible. This drug does not penetrate the CNS.

Tafenoquine Tafenoquine is an 8-aminoquinoline with causal prophylactic activity. Its prolonged half-life (2–3 weeks) allows longer dosing intervals when the drug is used for prophylaxis. Tafenoquine has been well tolerated in clinical trials. When tafenoquine is taken with food, its absorption is increased by 50% and the most commonly reported adverse event—mild GI upset—is diminished. Like primaquine, tafenoquine is a potent oxidizing agent, causing hemolysis in patients with G6PD deficiency as well as methemoglobinemia.

Tetracyclines See Table 246e-1 and Chap. 170.

Thiabendazole Discovered in 1961, thiabendazole remains one of the most potent of the numerous benzimidazole derivatives. However, its

use has declined significantly because of a higher frequency of adverse effects than is seen with other, equally effective agents.

Thiabendazole is active against most intestinal nematodes that infect humans. Although the exact mechanism of its antihelminthic activity has not been fully elucidated, it is likely to be similar to that of other benzimidazole drugs: namely, inhibition of polymerization of parasite β -tubulin. The drug also inhibits the helminth-specific enzyme fumarate reductase. In animals, thiabendazole has anti-inflammatory, antipyretic, and analgesic effects, which may explain its usefulness in dracunculiasis and trichinellosis. Thiabendazole also suppresses egg and/or larval production by some nematodes and may inhibit the subsequent development of eggs or larvae passed in feces. Despite the emergence and global spread of thiabendazole-resistant trichostrongyliasis among sheep, there have been no reports of drug resistance in humans.

Thiabendazole is available in tablet form and as an oral suspension. The drug is rapidly absorbed from the GI tract but can also be absorbed through the skin. Thiabendazole should be taken after meals. This agent is extensively metabolized in the liver before ultimately being excreted; most of the dose is excreted within the first 24 h. The usual dose of thiabendazole is determined by the patient's weight, but some treatment regimens are parasite specific. No particular adjustments are recommended in patients with renal or hepatic failure; only cautious use is advised.

Coadministration of thiabendazole to patients taking theophylline can result in an increase in theophylline levels by >50%. Therefore, serum levels of theophylline should be monitored closely in this situation.

Tinidazole This nitroimidazole is effective for the treatment of amebiasis, giardiasis, and trichomoniasis. Like metronidazole, tinidazole must undergo reductive activation by the parasite's metabolic system before it can act on protozoal targets. Tinidazole inhibits the synthesis of new DNA in the parasite and causes degradation of existing DNA. The reduced free-radical derivatives alkylate DNA, with consequent cytotoxic damage to the parasite. This damage appears to be produced by short-lived reduction intermediates, resulting in helix destabilization and strand breakage of DNA. The mechanism of action and side effects of tinidazole are similar to those of metronidazole, but adverse events appear to be less frequent and severe with tinidazole. In addition, the significantly longer half-life of tinidazole (>12 h) offers potential cure with a single dose.

Triclabendazole While most benzimidazoles have broad-spectrum antihelminthic activity, they exhibit minimal or no activity against *F. hepatica*. In contrast, the antihelminthic activity of triclabendazole is highly specific for *Fasciola* and *Paragonimus* species, with little activity against nematodes, cestodes, and other trematodes. Triclabendazole is effective against all stages of *Fasciola* species. The active sulfoxide metabolite of triclabendazole binds to fluke tubulin by assuming a unique nonplanar configuration and disrupts microtubule-based processes. Resistance to triclabendazole in veterinary use has been reported in Australia and Europe; however, no resistance has been documented in humans.

Triclabendazole is rapidly absorbed after oral ingestion; administration with food enhances its absorption and shortens the elimination half-life of the active metabolite. Both the sulfoxide and the sulfone metabolites are highly protein bound (>99%). Treatment with triclabendazole is typically given in one or two doses. No clinical data are available regarding dose adjustment in renal or hepatic insufficiency; however, given the short course of therapy and extensive hepatic metabolism of triclabendazole, dose adjustment is unlikely to be necessary. No information exists on drug interactions.

Trimethoprim-Sulfamethoxazole See Table 246e-1 and Chap. 170.

SECTION 18 PROTOZOAL INFECTIONS

247 Amebiasis and Infection with Free-Living Amebas

Rosa M. Andrade, Sharon L. Reed

AMEBIASIS


DEFINITION

Amebiasis is an infection with the intestinal protozoan *Entamoeba histolytica*. About 90% of infections are asymptomatic, and the remaining 10% produce a spectrum of clinical syndromes ranging from dysentery to abscesses of the liver or other organs.

LIFE CYCLE AND TRANSMISSION

E. histolytica is acquired by ingestion of viable cysts from fecally contaminated water, food, or hands. Food-borne exposure is most prevalent and is particularly likely when food handlers are shedding cysts or food is being grown with feces-contaminated soil, fertilizer, or water. Besides the drinking of contaminated water, less common means of transmission include oral and anal sexual practices and—in rare instances—direct rectal inoculation through colonic irrigation devices. Motile trophozoites are released from cysts in the small intestine and, in most patients, remain as harmless commensals in the large bowel. After encystation, infectious cysts are shed in the stool and can survive for several weeks in a moist environment. In some patients, the trophozoites invade either the bowel mucosa, causing symptomatic colitis, or the bloodstream, causing distant abscesses of the liver, lungs, or brain. The trophozoites may not encyst in patients with active dysentery, and motile hematophagous trophozoites are frequently present in fresh stools. Trophozoites are rapidly killed by exposure to air or stomach acid, however, and therefore cannot transmit infection.

EPIDEMIOLOGY

 About 10% of the world's population is infected with *Entamoeba*, the majority with noninvasive *Entamoeba dispar*. Amebiasis results from infection with *E. histolytica* and is the third most common cause of death from parasitic disease (after schistosomiasis and malaria). Invasive colitis and liver abscesses are sevenfold more common among men than among women; this difference has been attributed to a disparity in complement-mediated killing. The

wide spectrum of clinical disease caused by *Entamoeba* is due in part to the differences between these two infecting species. *E. histolytica* has unique isoenzymes, surface antigens, DNA markers, and virulence properties that distinguish it from other genetically related and morphologically identical species, such as *E. dispar* and *E. moshkovskii*.

Most asymptomatic carriers, including men who have sex with men (MSM) and patients with AIDS, harbor *E. dispar* and have self-limited infections. In this respect, *E. dispar* is dissimilar to other enteric pathogens such as *Cryptosporidium* and *Cystoisospora belli*, which can cause self-limited illnesses in immunocompetent hosts but devastating diarrhea in patients with AIDS. These observations indicate that *E. dispar* is incapable of causing invasive disease. Unlike *E. dispar*, *E. histolytica* can cause invasive disease, as demonstrated in recent reports from Korea, China, and India that suggest higher prevalences of amebic seroconversion, invasive amebiasis, and amebic liver abscesses among HIV-positive than HIV-negative patients. In another study, 10% of asymptomatic patients who were colonized with *E. histolytica* went on to develop amebic colitis, while the rest remained asymptomatic and cleared the infection within 1 year.

The potential of *E. moshkovskii* to cause diarrhea, weight loss, and colitis was recently demonstrated in a mouse model of cecal infection. However, the pathogenic potential of this species is not clear. A prospective evaluation of children from the Mirpur community of Dhaka, Bangladesh, found that most children who had diarrheal diseases associated with *E. moshkovskii* were simultaneously infected with at least one other enteric pathogen.

Areas of highest incidence of *Entamoeba* infection (due to inadequate sanitation and crowding) include most developing countries in the tropics, particularly Mexico, India, and nations of Central and South America, tropical Asia, and Africa. In a 4-year follow-up study of preschool children in a highly endemic area of Bangladesh, 80% of children had at least one episode of *E. histolytica* infection and 53% had more than one episode. Naturally acquired immunity did develop but was usually short-lived and correlated with the presence in the stool of secretory IgA antibody to the major adherence lectin galactose *N*-acetylgalactosamine (Gal/GalNAc). The main groups at risk for amebiasis in developed countries are returned travelers, recent immigrants, MSM, military personnel, and inmates of institutions. Data from the GeoSentinel Surveillance Network, which come from tropical medicine clinics on six continents, showed that, among long-term travelers (trip duration, >6 months), diarrhea due to *E. histolytica* was among the most common diagnoses.

Both trophozoites (Fig. 247-1) and cysts (Fig. 247-2) are found in the intestinal lumen, but only trophozoites of *E. histolytica* invade tissue. The trophozoite is 20–60 μm in diameter and contains vacuoles and a nucleus with a characteristic central nucleolus. In animals, depletion of intestinal mucus, diffuse inflammation, and disruption of the epithelial barrier precede trophozoite contact with the colonic mucosa. Trophozoites attach to colonic mucus and epithelial cells by their Gal/GalNAc lectin. The earliest intestinal lesions are microulcerations of the mucosa of the cecum, sigmoid colon, or rectum that release erythrocytes, inflammatory cells, and epithelial cells. Proctoscopy reveals small ulcers with heaped-up margins and normal intervening mucosa (Fig. 247-3A). Submucosal extension of ulcerations under viable-appearing surface mucosa causes the classic “flask-shaped” ulcer containing trophozoites at the margins of dead and viable tissues. Although neutrophilic infiltrates may accompany the early lesions in animals, human intestinal infection is marked by a paucity of inflammatory cells, probably in part because of the killing of neutrophils by trophozoites (Fig. 247-3B). Treated ulcers characteristically heal with little or no scarring. Occasionally, however, full-thickness necrosis and perforation occur.

Rarely, intestinal infection results in the formation of a mass lesion, or *ameboma*, in the bowel lumen. The overlying mucosa is usually thin and ulcerated, while other layers of the wall are thickened, edematous, and hemorrhagic; this condition results in exuberant formation of granulation tissue with little fibrous-tissue response.

A number of virulence factors have been linked to the ability of *E. histolytica* to invade through the interglandular epithelium. One factor consists of the extracellular cysteine proteinases that degrade collagen, elastin, IgA, IgG, and the anaphylatoxins C3a and C5a. Other enzymes may disrupt glycoprotein bonds between mucosal epithelial cells in the gut. Amebas can lyse neutrophils, monocytes, lymphocytes, and cells of colonic and hepatic lines. The cytolytic effect of amebas appears to require direct contact with target cells and may be linked to the release of phospholipase A and pore-forming peptides. *E. histolytica* trophozoites also cause apoptosis of human cells. Phagocytosis is a virulence factor that leads to defective parasite proliferation if inhibited. This process is potentially modulated by calmodulin-like calcium-binding protein 3, which pairs with actin and myosin during initiation and formation of phagosomes. Another virulence factor is the ability to resist reactive oxygen species, reactive nitrogen species such as nitric oxide, or S-nitrosothiols such as S-nitrosoglutathione (GSNO) and S-nitrosocysteine (CysNO). *E. histolytica* trophozoites are constantly exposed to reactive oxygen and nitrogen species from their own metabolism and host defenses during tissue invasion. Overexpression of hydrogen peroxide regulatory motif-binding protein appears to increase *E. histolytica* cytotoxicity. Since *E. histolytica* lacks glutathione and glutathione reductase, it relies on its thioredoxin/thioredoxin reductase system to prevent, regulate, and repair the damage caused by oxidative stress. This antioxidant system is versatile in that it can reduce

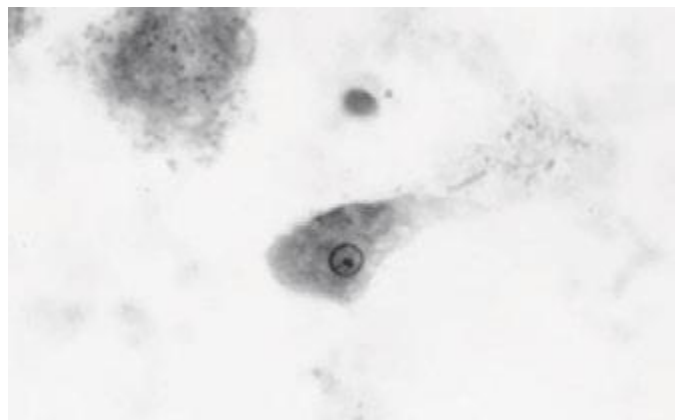


FIGURE 247-1 Trophozoite of *E. histolytica*. A single nucleus with a central, dot-like nucleolus is seen (trichrome stain).

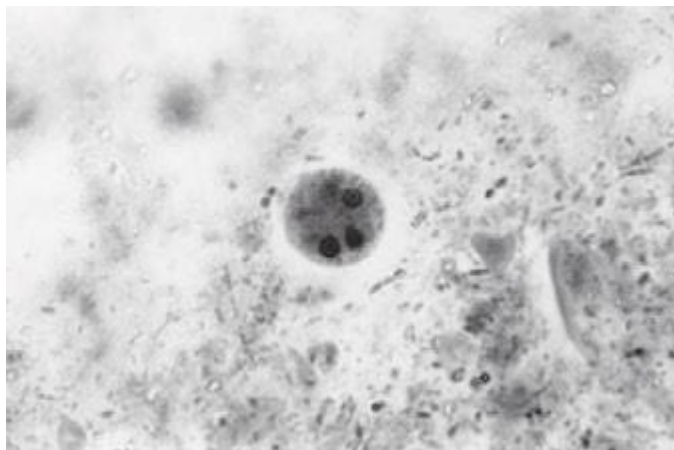
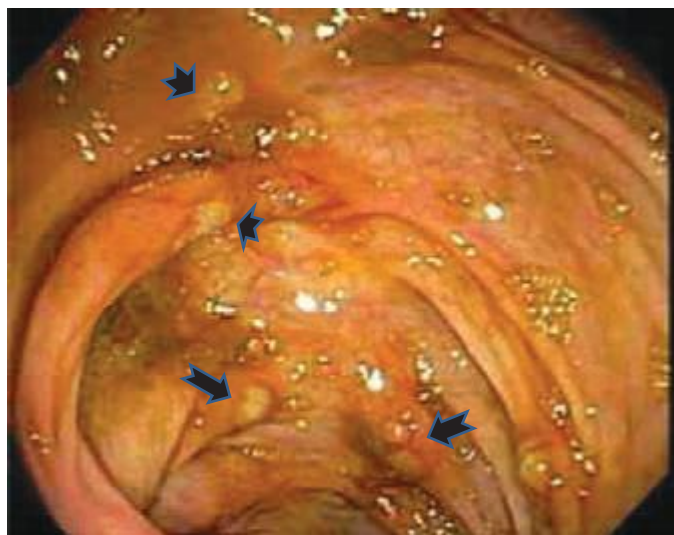
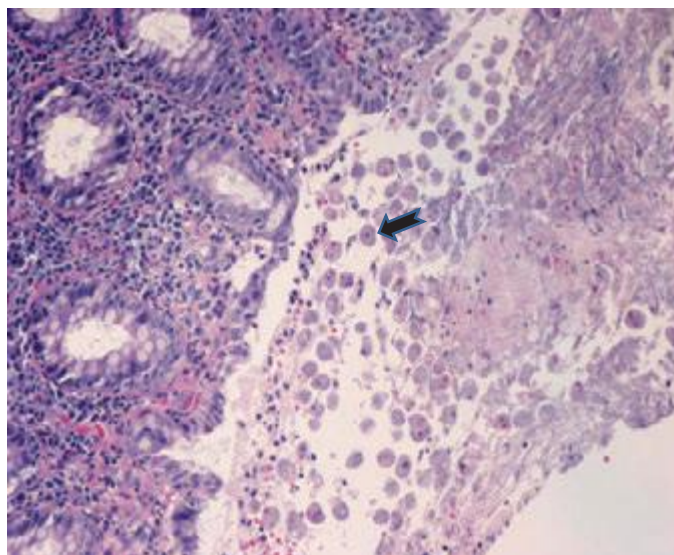


FIGURE 247-2 Cyst of *E. histolytica*. Three of the four nuclei are visible (trichrome stain).



A



B

FIGURE 247-3 Endoscopic and histopathologic features of intestinal amebiasis. **A.** Appearance of ulcers on colonoscopy (arrows). **B.** Inflammatory infiltrate and *E. histolytica* trophozoites (arrow) in invasive amebic colitis (hematoxylin and eosin). (Courtesy of the Department of Pathology and Gastroenterology, VA San Diego Medical Center.)

reactive nitrogen species and use an alternative electron donor such as the reduced form of nicotinamide adenine dinucleotide. Metronidazole, the current standard of therapy for amebiasis, seems to exert its anti-parasitic effect through the inhibition of this antioxidant system. Newer therapeutic candidates targeting this system, such as auranofin, also have demonstrated *in vitro* and *in vivo* efficacy against this parasite.

Liver abscesses are always preceded by intestinal colonization, which may be asymptomatic. Blood vessels may be compromised early by wall lysis and thrombus formation. Trophozoites invade veins to reach the liver through the portal venous system. *E. histolytica* is resistant to complement-mediated lysis—a property critical to survival in the bloodstream. In contrast, *E. dispar* is rapidly lysed by complement and is thus restricted to the bowel lumen. Inoculation of amebas into the portal system of hamsters results in an acute cellular infiltrate consisting predominantly of neutrophils. Later, the neutrophils are lysed by contact with amebas, and the release of neutrophil toxins may contribute to necrosis of hepatocytes. The liver parenchyma is replaced by necrotic material that is surrounded by a thin rim of congested liver tissue. The necrotic contents of a liver abscess are classically described as “anchovy paste,” although the fluid is variable in color and is composed of bacteriologically sterile granular debris with few or no cells. Amebas, if seen, tend to be found near the capsule of the abscess.

Host innate and adaptive immunity are important factors that determine susceptibility to invasive disease and its clinical outcome. While neutrophils were thought to contribute to tissue damage in intestinal and liver amebiasis due to their cytotoxic effects on host epithelial cells, a recent report suggests that they may exert a protective effect in susceptible mice. Neutropenia, induced with an antibody to Gr-1 (i.e., to peripheral neutrophils), led to death in C3H/HeJ mice and to severe disease in CBA mice (both of which are relatively susceptible to *E. histolytica* infection), while it had no effect on C57BL/6 mice, which are known for their intrinsic resistance to infection with this parasite.

Antimicrobial peptides, such as cathelicidins, are an important part of innate immunity and are induced by *E. histolytica* upon intestinal invasion in a mouse model. In this model, cecal cathelicidin-related antimicrobial peptide (CRAMP) mRNA increased more than fourfold by 3 days and more than 100-fold at 7 days. However, *E. histolytica* remained resistant to cathelicidin-mediated killing, probably because the antimicrobial peptide was digested by amebic cysteine proteinases.



IgA plays a critical role in acquired immunity to *E. histolytica*. A study in Bangladeshi schoolchildren revealed that an intestinal IgA response to Gal/GalNAc reduced the risk of new *E. histolytica* infection by 64%. Serum IgG antibody is not protective; titers correlate with the duration of illness rather than with the severity of disease. Indeed, Bangladeshi children with a serum IgG response were more likely than those without such a response to develop new *E. histolytica* infection. In infants from this same Bangladeshi community, passive immunity conferred by maternal parasite-specific IgA via breastfeeding resulted in a 39% reduction in risk of infection and a 64% reduction in risk of diarrheal disease from *E. histolytica* during the first year of life.

A link between nutrition and immunity is demonstrated by the elevated rate of infections due to protozoan parasites, including *E. histolytica*, among undernourished children in developing countries. Resistance to amebiasis is associated with a polymorphism in the receptor for the adipocytokine leptin. Children in a Bangladeshi cohort with a mutant R223 leptin receptor allele were nearly four times more likely to be infected with *E. histolytica* than those carrying the ancestral Q223 allele. This mutant allele is overrepresented in many geographic areas with a high prevalence of amebiasis, such as Bangladesh and India.

CLINICAL SYNDROMES

Intestinal Amebiasis The most common type of amebic infection is asymptomatic cyst passage. Even in highly endemic areas, most patients harbor *E. dispar*.

Symptomatic amebic colitis develops 2–6 weeks after the ingestion of infectious *E. histolytica* cysts. A gradual onset of lower abdominal pain and mild diarrhea is followed by malaise, weight loss, and diffuse

lower abdominal or back pain. Cecal involvement may mimic acute appendicitis. Patients with full-blown dysentery may pass 10–12 stools per day. The stools contain little fecal material and consist mainly of blood and mucus. In contrast to those with bacterial diarrhea, fewer than 40% of patients with amebic dysentery are febrile. Virtually all patients have heme-positive stools.

More fulminant intestinal infection, with severe abdominal pain, high fever, and profuse diarrhea, is rare and occurs predominantly in children. Patients may develop toxic megacolon, in which there is severe bowel dilation with intramural air. Patients receiving glucocorticoids are at risk for severe amebiasis. Uncommonly, patients develop a chronic form of amebic colitis, which can be confused with inflammatory bowel disease. The association between severe amebiasis complications and glucocorticoid therapy emphasizes the importance of excluding amebiasis when inflammatory bowel disease is suspected. An occasional patient presents with only an asymptomatic or tender abdominal mass caused by an ameboma, which is easily confused with cancer on barium studies. A positive serologic test or biopsy can prevent unnecessary surgery in this setting. The syndrome of post-amebic colitis—i.e., persistent diarrhea following documented cure of amebic colitis—is controversial; no evidence of recurrent amebic infection can be found, and re-treatment usually has no effect.

Amebic Liver Abscess Extraintestinal infection by *E. histolytica* most often involves the liver. Of travelers who develop an amebic liver abscess after leaving an endemic area, 95% do so within 5 months. Young patients with an amebic liver abscess are more likely than older patients to present in the acute phase with prominent symptoms of <10 days' duration. Most patients are febrile and have right-upper-quadrant pain, which may be dull or pleuritic in nature and may radiate to the shoulder. Point tenderness over the liver and right-sided pleural effusion are common. Jaundice is rare. Although the initial site of infection is the colon, fewer than one-third of patients with an amebic abscess have active diarrhea. Older patients from endemic areas are more likely to have a subacute course lasting 6 months, with weight loss and hepatomegaly. About one-third of patients with chronic presentations are febrile. Thus, the clinical diagnosis of an amebic liver abscess may be difficult to establish because the symptoms and signs are often nonspecific. Since 10–15% of patients present only with fever, amebic liver abscess must be considered in the differential diagnosis of fever of unknown origin ([Chap. 26](#)).

Complications of Amebic Liver Abscess Pleuropulmonary involvement, which is reported in 20–30% of patients, is the most frequent complication of amebic liver abscess. Manifestations include sterile effusions, contiguous spread from the liver, and rupture into the pleural space. Sterile effusions and contiguous spread usually resolve with medical therapy, but frank rupture into the pleural space requires drainage. A hepatobronchial fistula may cause cough productive of large amounts of necrotic material that may contain amebas. This dramatic complication carries a good prognosis. Abscesses that rupture into the peritoneum may present as an indolent leak or an acute abdomen and require both percutaneous catheter drainage and medical therapy. Rupture into the pericardium, usually from abscesses of the left lobe of the liver, carries the gravest prognosis; it can occur during medical therapy and requires surgical drainage.

Other Extraintestinal Sites The genitourinary tract may become involved by direct extension of amebiasis from the colon or by hematogenous spread of the infection. Painful genital ulcers, characterized by a punched-out appearance and profuse discharge, may develop secondary to extension from either the intestine or the liver. Both of these conditions respond well to medical therapy. Cerebral involvement has been reported in fewer than 0.1% of patients in large clinical series. Symptoms and prognosis depend on the size and location of the lesion.

DIAGNOSTIC TESTS

Laboratory Diagnosis Stool examinations, serologic tests, and non-invasive imaging of the liver are the most important procedures in the diagnosis of amebiasis. Fecal findings suggestive of amebic colitis

1366 include a positive test for heme, a paucity of neutrophils, and amebic cysts or trophozoites. The definitive diagnosis of amebic colitis is made by the demonstration of hematophagous trophozoites of *E. histolytica* (Fig. 247-1). Because trophozoites are killed rapidly by water, drying, or barium, it is important to examine at least three fresh stool specimens. Examination of a combination of wet mounts, iodine-stained concentrates, and trichrome-stained preparations of fresh stool and concentrates for cysts (Fig. 247-2) or trophozoites (Fig. 247-1) confirms the diagnosis in 75–95% of cases. Culture of amebas is more sensitive, but this diagnostic method is not routinely available. If stool examinations are negative, sigmoidoscopy with biopsy of the edge of ulcers may increase the yield, but this procedure is dangerous during fulminant colitis because of the risk of perforation. Trophozoites in a biopsy specimen from a colonic mass confirm the diagnosis of ameboma, but trophozoites are rare in liver aspirates because they are found in the abscess capsule and not in the readily aspirated necrotic center. Accurate diagnosis requires experience, since the trophozoites may be confused with neutrophils and the cysts must be differentiated morphologically from those of *Entamoeba hartmanni*, *Entamoeba coli*, and *Endolimax nana*, which do not cause clinical disease and do not warrant therapy. Unfortunately, the cysts of *E. histolytica* cannot be distinguished microscopically from those of *E. dispar* or *E. moshkovskii*. Therefore, the microscopic diagnosis of *E. histolytica* can be made only by the detection of *Entamoeba* trophozoites that have ingested erythrocytes. In terms of sensitivity, stool diagnostic tests based on the detection of the Gal/GalNAc lectin of *E. histolytica* compare favorably with the polymerase chain reaction and with isolation in culture followed by isoenzyme analysis.

Serology is an important addition to the methods used for parasitologic diagnosis of invasive amebiasis. Enzyme-linked immunosorbent assays and agar gel diffusion assays are positive in more than 90% of patients with colitis, amebomas, or liver abscess. Positive results in conjunction with the appropriate clinical syndrome suggest active disease because serologic findings usually revert to negative within 6–12 months. Even in highly endemic areas such as South Africa, fewer than 10% of asymptomatic individuals have a positive amebic serology. The interpretation of the indirect hemagglutination test is more difficult because titers may remain positive for as long as 10 years.

Up to 10% of patients with acute amebic liver abscess may have negative serologic findings; in suspected cases with an initially negative result, testing should be repeated in 1 week. In contrast to carriers of *E. dispar*, most asymptomatic carriers of *E. histolytica* develop antibodies. Thus, serologic tests are helpful in assessing the risk of invasive amebiasis in asymptomatic, cyst-passing individuals in nonendemic areas. Serologic tests also should be performed in patients with ulcerative colitis before the institution of glucocorticoid therapy to prevent the development of severe colitis or toxic megacolon owing to unsuspected amebiasis.

Routine hematology and chemistry tests usually are not very helpful in the diagnosis of invasive amebiasis. About three-fourths of patients with an amebic liver abscess have leukocytosis ($>10,000$ cells/ μL); this condition is particularly likely if symptoms are acute or complications have developed. Invasive amebiasis does not elicit eosinophilia. Anemia, if present, is usually multifactorial. Even with large liver abscesses, liver enzyme levels are normal or minimally elevated. The alkaline phosphatase level is most often elevated and may remain so for months. Aminotransferase elevations suggest acute disease or a complication.

Radiographic Studies Radiographic barium studies are potentially dangerous in acute amebic colitis. Amebomas are usually identified first by a barium enema, but biopsy is necessary for differentiation from carcinoma.

Radiographic techniques such as ultrasonography, CT, and MRI are all useful for detection of the round or oval hypoechoic cyst of an amebic liver abscess. More than 80% of patients who have had symptoms for >10 days have a single abscess of the right lobe of the liver (Fig. 247-4). Approximately 50% of patients who have had symptoms for <10 days have multiple abscesses. Findings associated

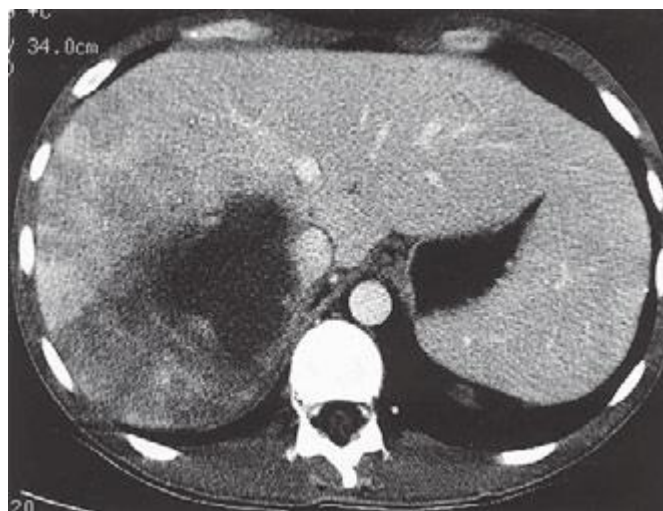


FIGURE 247-4 Abdominal CT scan of a large amebic abscess of the right lobe of the liver. (Courtesy of the Department of Radiology, UCSD Medical Center, San Diego; with permission.)

with complications include large abscesses (>10 cm) in the superior part of the right lobe, which may rupture into the pleural space; multiple lesions, which must be differentiated from pyogenic abscesses; and lesions of the left lobe, which may rupture into the pericardium. Because abscesses resolve slowly and may increase in size in patients who are responding clinically to therapy, frequent follow-up ultrasonography may prove confusing. Complete resolution of a liver abscess within 6 months can be anticipated in two-thirds of patients, but 10% may have persistent abnormalities for a year.

Differential Diagnosis The differential diagnosis of intestinal amebiasis includes bacterial diarrheas (Chap. 160) caused by *Campylobacter* (Chap. 192); enteroinvasive *Escherichia coli* (Chap. 186); and species of *Shigella* (Chap. 191), *Salmonella* (Chap. 190), and *Vibrio* (Chap. 193). Although the typical patient with amebic colitis has less prominent fever than in these other conditions as well as heme-positive stools with few neutrophils, correct diagnosis requires bacterial cultures, microscopic examination of stools, and amebic serologic testing. As has already been mentioned, amebiasis must be ruled out in any patient thought to have inflammatory bowel disease.

Because of the variety of presenting signs and symptoms, amebic liver abscess can easily be confused with pulmonary or gallbladder disease or with any febrile illness with few localizing signs, such as malaria (Chap. 248) or typhoid fever (Chap. 190). The diagnosis should be considered in members of high-risk groups who have recently traveled outside the United States (Chap. 149) and in inmates of institutions. Once radiographic studies have identified an abscess in the liver, the most important differential diagnosis is between amebic and pyogenic abscess. Patients with pyogenic abscess typically are older and have a history of underlying bowel disease or recent surgery. Amebic serology is helpful, but aspiration of the abscess, with Gram's staining and culture of the material, may be required for differentiation of the two diseases.

TREATMENT AMEBIASIS

INTESTINAL DISEASE

The drugs used to treat amebiasis can be classified according to their primary site of action (Table 247-1). Luminal amebicides are poorly absorbed; they reach high concentrations in the bowel, but their activity is limited to cysts and trophozoites close to the mucosa. Only two luminal drugs are available in the United States: iodoquinol and paromomycin. Indications for the use of luminal agents include eradication of cysts in patients with colitis or a liver abscess and treatment of asymptomatic carriers. The majority of

TABLE 247-1 DRUG THERAPY FOR AMEBIASIS

Indication	Therapy
Asymptomatic carriage	Luminal agent: iodoquinol (650-mg tablets), 650 mg tid for 20 days; or paromomycin (250-mg tablets), 500 mg tid for 10 days
Acute colitis	Metronidazole (250- or 500-mg tablets), 750 mg PO or IV tid for 5–10 days; or tinidazole, 2 g/d PO for 3 days plus Luminal agent as above
Amebic liver abscess	Metronidazole, 750 mg PO or IV for 5–10 days; or tinidazole, 2 g PO once; or ornidazole, ^a 2 g PO once plus Luminal agent as above

^aNot available in the United States.

asymptomatic individuals who pass cysts are colonized with *E. dispar*, which does not warrant specific therapy. However, it is prudent to treat asymptomatic individuals who pass cysts unless *E. dispar* colonization can be definitively demonstrated by specific antigen-detection tests.

Tissue amebicides reach high concentrations in the blood and tissue after oral or parenteral administration. The development of nitroimidazole compounds, especially metronidazole, was a major advance in the treatment of invasive amebiasis. Patients with amebic colitis should be treated with IV or oral metronidazole. Side effects include nausea, vomiting, abdominal discomfort, and a disulfiram-like reaction. Another longer-acting imidazole compound, tinidazole, is also effective and available in the United States. All patients should also receive a full course of therapy with a luminal agent, since metronidazole does not eradicate cysts. Resistance to metronidazole has been selected in the laboratory but has not been found in clinical isolates. Relapses are not uncommon and probably represent reinfection or failure to eradicate amebas from the bowel because of an inadequate dosage or duration of therapy.

AMEBIC LIVER ABSCESS

Metronidazole is the drug of choice for amebic liver abscess. Longer-acting nitroimidazoles (tinidazole and ornidazole) have been effective as single-dose therapy in developing countries. With early diagnosis and therapy, mortality rates from uncomplicated amebic liver abscess are <1%. There is no evidence that combined therapy with two drugs is more effective than the single-drug regimen. Studies of South Africans with liver abscesses demonstrated that 72% of patients without intestinal symptoms had bowel infection with *E. histolytica*; thus, all treatment regimens should include a luminal agent to eradicate cysts and prevent further transmission. Amebic liver abscess recurs rarely.

More than 90% of patients respond dramatically to metronidazole therapy with decreases in both pain and fever within 72 h. Indications for aspiration of liver abscesses are (1) the need to rule out a pyogenic abscess, particularly in patients with multiple lesions; (2) the lack of a clinical response in 3–5 days; (3) the threat of imminent rupture; and (4) the need to prevent rupture of left-lobe abscesses into the pericardium. There is no evidence that aspiration, even of large abscesses (up to 10 cm), accelerates healing. Percutaneous drainage may be successful even if the liver abscess has already ruptured. Surgery should be reserved for instances of bowel perforation and rupture into the pericardium.


PREVENTION

Amebic infection is spread by ingestion of food or water contaminated with cysts. Since an asymptomatic carrier may excrete up to 15 million cysts per day, prevention of infection requires adequate sanitation and eradication of cyst carriage. In high-risk areas, infection can be minimized by the avoidance of unpeeled fruits and vegetables

and the use of bottled water. Because cysts are resistant to readily attainable levels of chlorine, disinfection by iodination (tetraglycine hydroperiodide) is recommended. There is no effective prophylaxis.

INFECTION WITH FREE-LIVING AMEBAS

EPIDEMIOLOGY

 Free-living amebas of the genera *Acanthamoeba* and *Naegleria* are distributed throughout the world and have been isolated from a wide variety of fresh and brackish water, including that from lakes, taps, hot springs, swimming pools, and heating and air-conditioning units, and even from the nasal passages of healthy children. Encystation may protect the protozoa from desiccation and food deprivation. The persistence of *Legionella pneumophila* in water supplies may be attributable in part to chronic infection of free-living amebas, particularly *Naegleria*. Free-living amebas of the genus *Balamuthia* have been isolated from soil samples, including a sample from a flowerpot linked to a fatal infection in a child.

NAEGLERIA INFECTIONS

Primary amebic meningoencephalitis caused by *Naegleria fowleri* follows the aspiration of water contaminated with trophozoites or cysts or the inhalation of contaminated dust, leading to invasion of the olfactory neuroepithelium. Infection is most common among otherwise healthy children or young adults, who often report recent swimming in lakes or heated swimming pools. Rarely, some cases occur when contaminated water is used for nasal irrigation. After an incubation period of 2–15 days, severe headache, high fever, nausea, vomiting, and meningismus develop. Photophobia and palsies of the third, fourth, and sixth cranial nerves are common. Rapid progression to seizures and coma may follow. The prognosis is uniformly poor: most patients die within a week. Recently, two surviving children were treated with miltefosine, an investigational drug that is available through the Centers for Disease Control and Prevention (CDC) for the treatment of *Naegleria* infections.

The diagnosis of *Naegleria* infection should be considered in any patient who has purulent meningitis without evidence of bacteria on Gram's staining, antigen detection assay, and culture. Other laboratory findings resemble those for fulminant bacterial meningitis, with elevated intracranial pressure, high white blood cell counts (up to 20,000/ μ L), and elevated protein concentrations and low glucose levels in cerebrospinal fluid (CSF). Diagnosis depends on the detection of motile trophozoites in wet mounts of fresh spinal fluid. Antibodies to *Naegleria* species have been detected in healthy adults; serologic testing is not useful in the diagnosis of acute infection.

ACANTHAMOEBA INFECTIONS

Granulomatous Amebic Encephalitis Infection with *Acanthamoeba* species follows a more indolent course and typically occurs in chronically ill or debilitated patients. Risk factors include lymphoproliferative disorders, chemotherapy, glucocorticoid therapy, lupus erythematosus, and AIDS. Infection usually reaches the central nervous system hematogenously from a primary focus in the sinuses, skin, or lungs. In the central nervous system, the onset is insidious, and the syndrome often mimics a space-occupying lesion. Altered mental status, headache, and stiff neck may be accompanied by focal findings such as cranial nerve palsies, ataxia, and hemiparesis. Cutaneous ulcers or hard nodules containing amebas are frequently detected in AIDS patients with disseminated *Acanthamoeba* infection and can be an important diagnostic site.

Examination of the CSF for trophozoites may be diagnostically helpful, but lumbar puncture may be contraindicated because of increased intracerebral pressure. CT frequently reveals cortical and subcortical lesions of decreased density consistent with embolic infarcts. In other patients, multiple enhancing lesions with edema may mimic the computed tomographic appearance of toxoplasmosis (Chap. 253). Demonstration of the trophozoites and cysts of *Acanthamoeba* on wet mounts or in biopsy specimens establishes the diagnosis. Culture on nonnutrient agar plates seeded with *Escherichia coli* also may be helpful. Fluorescein-labeled antiserum is available

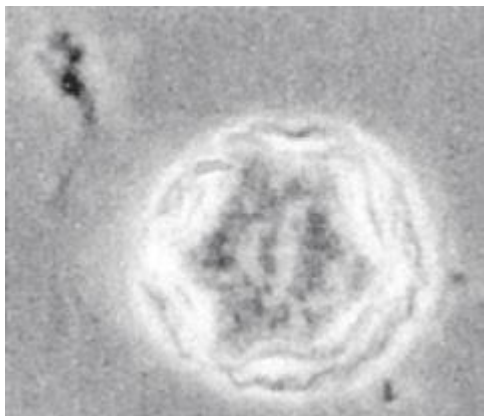


FIGURE 247-5 Double-walled cyst of *Acanthamoeba castellanii*, as seen by phase-contrast microscopy. (From DJ Krogstad et al, in A Balows et al [eds]: *Manual of Clinical Microbiology*, 5th ed. Washington, DC, American Society for Microbiology, 1991.)

from the CDC for the detection of protozoa in biopsy specimens. Granulomatous amebic encephalitis in patients with AIDS may have an accelerated course (with survival for only 3–40 days) because of poor granuloma formation in these individuals. Various antimicrobial agents have been used to treat *Acanthamoeba* infection, but the infection is almost uniformly fatal. The CDC has now made miltefosine available because of improved survival rates when the drug is included in treatment regimens.

Keratitis The incidence of keratitis caused by *Acanthamoeba* has increased in the past 30 years, in part as a result of improved diagnosis. Earlier infections were associated with trauma to the eye and exposure to contaminated water. At present, most infections are linked to extended-wear contact lenses, and rare cases are associated with laser-assisted in situ keratomileusis (LASIK). Risk factors include the use of homemade saline, the wearing of lenses while swimming, and inadequate disinfection. Since contact lenses presumably cause microscopic trauma, the early corneal findings may be nonspecific. The first symptoms usually include tearing and the painful sensation of a foreign body. Once infection is established, progression is rapid; the characteristic clinical sign is an annular, paracentral corneal ring representing a corneal abscess. Deeper corneal invasion and loss of vision may follow.

The differential diagnosis includes bacterial, mycobacterial, and herpetic infection. The irregular polygonal cysts of *Acanthamoeba* (Fig. 247-5) may be identified in corneal scrapings or biopsy material, and trophozoites can be grown on special media. Cysts are resistant to available drugs, and the results of medical therapy have been disappointing. Some reports have suggested partial responses to propamidine isethionate eyedrops. Severe infections usually require keratoplasty.

BALAMUTHIA INFECTIONS

Balamuthia mandrillaris, a free-living amoeba previously referred to as a leptomyxid amoeba, is an important etiologic agent of amebic meningoencephalitis in immunocompetent hosts. The course is typically subacute, with focal neurologic signs, fever, seizures, and headaches leading to death within 1 week to several months after onset. Examination of CSF reveals mononuclear or neutrophilic pleocytosis, elevated protein levels, and normal to low glucose concentrations. Multiple hypodense lesions are usually detected with imaging studies (Fig. 247-6). This mixed picture of space-occupying lesions with CSF pleocytosis is suggestive of *Balamuthia*. Fluorescent antibody is available from the CDC for brain biopsy specimens. The variety of drugs used to treat the few surviving patients (i.e., fewer than five reported in the United States) includes pentamidine, flucytosine, sulfadiazine, and macrolides. The CDC recommends that miltefosine now be included, as for the other free-living amoebas. The differential diagnosis includes tuberculomas (Chap. 202) and neurocysticercosis (Chap. 260).

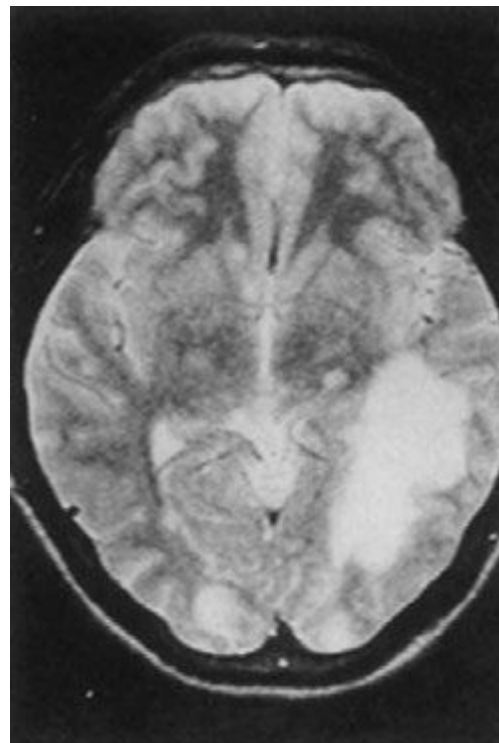


FIGURE 247-6 Brain MRI of amebic meningoencephalitis due to *Balamuthia mandrillaris*. A large lesion in the parieto-occipital lobe and other smaller lesions are seen. (Courtesy of the Department of Radiology, UCSD Medical Center, San Diego.)

248 Malaria

Nicholas J. White, Joel G. Breman

Humanity has but three great enemies: Fever, famine, and war; of these by far the greatest, by far the most terrible, is fever.

—William Osler

Malaria is a protozoan disease transmitted by the bite of infected *Anopheles* mosquitoes. The most important of the parasitic diseases of humans, it is transmitted in 106 countries containing 3 billion people and causes approximately 2000 deaths each day; mortality rates are decreasing as a result of highly effective control programs in several countries. Malaria has been eliminated from the United States, Canada, Europe, and Russia; in the late twentieth and early twenty-first centuries, however, its prevalence rose in many parts of the tropics. Increases in the drug resistance of the parasite, the insecticide resistance of its vectors, and human travel and migration have contributed to this resurgence. Occasional local transmission after importation of malaria has occurred in several southern and eastern areas of the United States and in Europe, indicating the continual danger to non-malarious countries. Although there are many successful new control initiatives as well as promising research initiatives, malaria remains today, as it has been for centuries, a heavy burden on tropical communities, a threat to nonendemic countries, and a danger to travelers.

ETIOLOGY AND PATHOGENESIS

Six species of the genus *Plasmodium* cause nearly all malarial infections in humans. These are *P. falciparum*, *P. vivax*, two morphologically identical sympatric species of *P. ovale* (as suggested by recent evidence), *P. malariae*, and—in Southeast Asia—the monkey malaria parasite *P. knowlesi* (Table 248-1). While almost all deaths are caused

TABLE 248-1 CHARACTERISTICS OF *PLASMODIUM* SPECIES INFECTING HUMANS

Characteristic	Finding for Indicated Species ^a			
	<i>P. falciparum</i>	<i>P. vivax</i>	<i>P. ovale</i>	<i>P. malariae</i>
Duration of intrahepatic phase (days)	5.5	8	9	15
Number of merozoites released per infected hepatocyte	30,000	10,000	15,000	15,000
Duration of erythrocytic cycle (hours)	48	48	50	72
Red cell preference	Younger cells (but can invade cells of all ages)	Reticulocytes and cells up to 2 weeks old	Reticulocytes	Older cells
Morphology	Usually only ring forms ^b ; banana-shaped gametocytes	Irregularly shaped large rings and trophozoites; enlarged erythrocytes; Schüffner's dots	Infected erythrocytes, enlarged and oval with tufted ends; Schüffner's dots	Band or rectangular forms of trophozoites common
Pigment color	Black	Yellow-brown	Dark brown	Brown-black
Ability to cause relapses	No	Yes	Yes	No

^aIn Southeast Asia, the monkey malaria parasite *P. knowlesi* also causes disease in humans. Young ring forms resemble those of *P. falciparum*, while older trophozoites resemble those of *P. malariae*. Reliable identification requires molecular genotyping. ^bParasitemias of >2% are suggestive of *P. falciparum* infection.

by *falciparum* malaria, *P. knowlesi* and occasionally *P. vivax* also can cause severe illness. Human infection begins when a female anopheline mosquito inoculates plasmodial *sporozoites* from its salivary gland during a blood meal (Fig. 248-1). These microscopic motile forms of the malaria parasite are carried rapidly via the bloodstream to the liver, where they invade hepatic parenchymal cells and begin a period of asexual reproduction. By this amplification process (known as *intrahepatic* or *preerythrocytic schizogony* or *merogony*), a single sporozoite eventually may produce from 10,000 to >30,000 daughter merozoites. The swollen infected liver cells eventually burst, discharging motile *merozoites* into the bloodstream. These merozoites then invade the red blood cells (RBCs) and multiply six- to twentyfold every 48 h (*P. knowlesi*, 24 h; *P. malariae*, 72 h). When the parasites reach densities of ~50/μL of blood (~100 million parasites in the blood of an adult), the symptomatic stage of the infection begins. In *P. vivax* and *P. ovale* infections, a proportion of the intrahepatic forms do not divide immediately but remain inert for a period ranging from 3 weeks to ≥1 year before reproduction begins. These dormant forms, or *hypnozoites*, are the cause of the relapses that characterize infection with these two species.

After entry into the bloodstream, merozoites rapidly invade erythrocytes and become *trophozoites*. Attachment is mediated via

a specific erythrocyte surface receptor. For *P. falciparum*, the reticulocyte-binding protein homologue 5 (PfRh5) is indispensable for erythrocyte invasion. Basigin (CD147, EMMPRIN) is the erythrocyte receptor of PfRh5. In the case of *P. vivax*, this receptor is related to the Duffy blood-group antigen Fy^a or Fy^b. Most West Africans and people with origins in that region carry the Duffy-negative FyFy phenotype and are therefore resistant to *P. vivax* malaria. During the early stage of intraerythrocytic development, the small “ring forms” of the different parasitic species appear similar under light microscopy. As the trophozoites enlarge, species-specific characteristics become evident, pigment becomes visible, and the parasite assumes an irregular or ameboid shape. By the end of the intraerythrocytic life cycle, the parasite has consumed two-thirds of the RBC's hemoglobin and has grown to occupy most of the cell. It is now called a *schizont*. Multiple nuclear divisions have taken place (*schizogony* or *merogony*). The RBC then ruptures to release 6–30 daughter merozoites, each potentially capable of invading a new RBC and repeating the cycle. The disease in human beings is caused by the direct effects of the asexual parasite—RBC invasion and destruction—and by the host's reaction. After release from the liver (*P. vivax*, *P. ovale*, *P. malariae*, *P. knowlesi*), some of the blood-stage parasites develop into morphologically distinct,

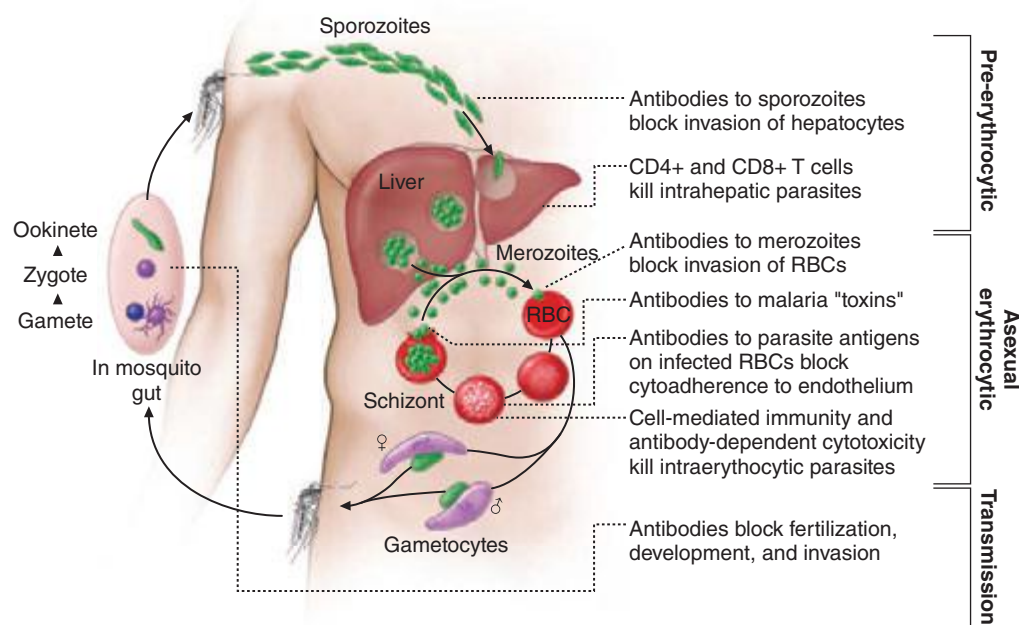


FIGURE 248-1 The malaria transmission cycle from mosquito to human and targets of immunity. RBC, red blood cell.

1370 longer-lived sexual forms (*gametocytes*) that can transmit malaria. In falciparum malaria, a delay of several asexual cycles precedes this switch to gametocytogenesis.

After being ingested in the blood meal of a biting female anopheline mosquito, the male and female gametocytes form a zygote in the insect's midgut. This zygote matures into an ookinete, which penetrates and encysts in the mosquito's gut wall. The resulting oocyst expands by asexual division until it bursts to liberate myriad motile sporozoites, which then migrate in the hemolymph to the salivary gland of the mosquito to await inoculation into another human at the next feeding.

EPIDEMIOLOGY

Malaria occurs throughout most of the tropical regions of the world (Fig. 248-2). *P. falciparum* predominates in Africa, New Guinea, and Hispaniola (i.e., the Dominican Republic and Haiti); *P. vivax* is more common in Central America. The prevalence of these two species is approximately equal in South America, the Indian subcontinent, eastern Asia, and Oceania. *P. malariae* is found in most endemic areas, especially throughout sub-Saharan Africa, but is much less common. *P. ovale* is relatively unusual outside of Africa and, where it is found, comprises <1% of isolates. Patients infected with *P. knowlesi* have been



FIGURE 248-2 Malaria-endemic countries in the Americas (*bottom*) and in Africa, the Middle East, Asia, and the South Pacific (*top*), 2007. CAR, Central African Republic; DROC, Democratic Republic of the Congo; UAE, United Arab Emirates. Several countries in the Americas, the Middle East, and North Africa are close to eliminating malaria.

identified on the island of Borneo and, to a lesser extent, elsewhere in Southeast Asia, where the main hosts, long-tailed and pig-tailed macaques, are found.

The epidemiology of malaria is complex and may vary considerably even within relatively small geographic areas. Endemicity traditionally has been defined in terms of parasitemia rates or palpable-spleen rates in children 2–9 years of age and classified as hypoendemic (<10%), mesoendemic (11–50%), hyperendemic (51–75%), and holoendemic (>75%). Until recently, it was uncommon to use these indices for planning control programs; however, many countries are now conducting national surveys to assess program progress. In holo- and hyperendemic areas (e.g., certain regions of tropical Africa or coastal New Guinea) where there is intense *P. falciparum* transmission, people may sustain more than one infectious mosquito bite per day and are infected repeatedly throughout their lives. In such settings, rates of morbidity and mortality due to malaria are considerable during early childhood. Immunity against disease is hard won in these areas, and the burden of disease in young children is high; by adulthood, however, most malarial infections are asymptomatic. As control measures progress and urbanization expands, environmental conditions become less conducive to transmission, and all age groups may lose protective immunity and become susceptible to illness. Constant, frequent, year-round infection is termed *stable transmission*. In areas where transmission is low, erratic, or focal, full protective immunity is not acquired, and symptomatic disease may occur at all ages. This situation usually exists in hypoendemic areas and is termed *unstable transmission*. Even in stable-transmission areas, there is often an increased incidence of symptomatic malaria coinciding with increased mosquito breeding and transmission during the rainy season. Malaria can behave like an epidemic disease in some areas, particularly those with unstable malaria, such as northern India (the Punjab region), the horn of Africa, Rwanda, Burundi, southern Africa, and Madagascar. An epidemic can develop when there are changes in environmental, economic, or social conditions, such as heavy rains following drought or migrations (usually of refugees or workers) from a nonmalarious region to an area of high transmission, along with failure to invest in national programs; a breakdown in malaria control and prevention services caused by war or civil disorder can intensify epidemic conditions. This situation usually results in considerable mortality among all age groups.

The principal determinants of the epidemiology of malaria are the number (density), the human-biting habits, and the longevity of the anopheline mosquito vectors. More than 100 of the >400 anopheline species can transmit malaria, but the ~40 species that do so commonly vary considerably in their efficiency as malaria vectors. More specifically, the transmission of malaria is directly proportional to the density of the vector, the square of the number of human bites per day per mosquito, and the tenth power of the probability of the mosquito's surviving for 1 day. Mosquito longevity is particularly important because the portion of the parasite's life cycle that takes place within the mosquito—from gametocyte ingestion to subsequent inoculation (*sporogony*)—lasts 8–30 days, depending on ambient temperature; thus, to transmit malaria, the mosquito must survive for >7 days. Sporogony is not completed at cooler temperatures—i.e., <16°C (60.8°F) for *P. vivax* and <21°C (69.8°F) for *P. falciparum*; thus transmission does not occur below these temperatures or at high altitudes, although malaria outbreaks and transmission have occurred in the highlands (>1500 m) of eastern Africa, which were previously free of vectors. The most effective mosquito vectors of malaria are those, such as *Anopheles gambiae* in Africa, that are long-lived, occur in high densities in tropical climates, breed readily, and bite humans in preference to other animals. The entomologic inoculation rate (i.e., the number of sporozoite-positive mosquito bites per person per year) is the most common measure of malaria transmission and varies from <1 in some parts of Latin America and Southeast Asia to >300 in parts of tropical Africa.

ERYTHROCYTE CHANGES IN MALARIA

After invading an erythrocyte, the growing malarial parasite progressively consumes and degrades intracellular proteins, principally

hemoglobin. The potentially toxic heme is detoxified by lipid-mediated crystallization to biologically inert hemozoin (malaria pigment). The parasite also alters the RBC membrane by changing its transport properties, exposing cryptic surface antigens, and inserting new parasite-derived proteins. The RBC becomes more irregular in shape, more antigenic, and less deformable.

In *P. falciparum* infections, membrane protuberances appear on the erythrocyte's surface 12–15 h after the cell's invasion. These “knobs” extrude a high-molecular-weight, antigenically variant, strain-specific erythrocyte membrane adhesive protein (PfEMP1) that mediates attachment to receptors on venular and capillary endothelium—an event termed *cytoadherence*. Several vascular receptors have been identified, of which intercellular adhesion molecule 1 is probably the most important in the brain, chondroitin sulfate B in the placenta, and CD36 in most other organs. Thus, the infected erythrocytes stick inside and eventually block capillaries and venules. At the same stage, these *P. falciparum*-infected RBCs may also adhere to uninfected RBCs (to form rosettes) and to other parasitized erythrocytes (agglutination). The processes of cytoadherence, rosetting, and agglutination are central to the pathogenesis of falciparum malaria. They result in the sequestration of RBCs containing mature forms of the parasite in vital organs (particularly the brain), where they interfere with microcirculatory flow and metabolism. Sequestered parasites continue to develop out of reach of the principal host defense mechanism: splenic processing and filtration. As a consequence, only the younger ring forms of the asexual parasites are seen circulating in the peripheral blood in falciparum malaria, and the level of peripheral parasitemia underestimates the true number of parasites within the body. Severe malaria is also associated with reduced deformability of the uninfected erythrocytes, which compromises their passage through the partially obstructed capillaries and venules and shortens RBC survival.

In the other human malarias, sequestration does not occur, and all stages of the parasite's development are evident on peripheral-blood smears. Whereas *P. vivax*, *P. ovale*, and *P. malariae* show a marked predilection for either young RBCs (*P. vivax*, *P. ovale*) or old cells (*P. malariae*) and produce a level of parasitemia that is seldom >2%, *P. falciparum* can invade erythrocytes of all ages and may be associated with very high levels of parasitemia.

HOST RESPONSE

Initially, the host responds to plasmodial infection by activating nonspecific defense mechanisms. Splenic immunologic and filtrative clearance functions are augmented in malaria, and the removal of both parasitized and uninfected erythrocytes is accelerated. The spleen is able to remove damaged ring-form parasites and return the once-infected erythrocytes to the circulation, where their survival period is shortened. The parasitized cells escaping splenic removal are destroyed when the schizont ruptures. The material released induces the activation of macrophages and the release of proinflammatory cytokines, which cause fever and exert other pathologic effects. Temperatures of ≥40°C (104°F) damage mature parasites; in untreated infections, the effect of such temperatures is to further synchronize the parasitic cycle, with eventual production of the regular fever spikes and rigors that originally served to characterize the different malarias. These regular fever patterns (quotidian, daily; tertian, every 2 days; quartan, every 3 days) are seldom seen today in patients who receive prompt and effective antimalarial treatment.

The geographic distributions of sickle cell disease, hemoglobins C and E, hereditary ovalocytosis, the thalassemias, and glucose-6-phosphate dehydrogenase (G6PD) deficiency closely resemble that of falciparum malaria before the introduction of control measures. This similarity suggests that these genetic disorders confer protection against death from falciparum malaria. For example, HbA/S heterozygotes (sickle cell trait) have a sixfold reduction in the risk of dying from severe falciparum malaria. Hemoglobin S-containing RBCs impair parasite growth at low oxygen tensions, and *P. falciparum*-infected RBCs containing hemoglobins S and C exhibit reduced cytoadherence because of reduced surface presentation of the adhesin PfEMP1. Parasite multiplication in HbA/E heterozygotes is reduced at high

1372 parasite densities. In Melanesia, children with α -thalassemia appear to have more frequent malaria (both vivax and falciparum) in the early years of life, and this pattern of infection appears to protect them against severe disease. In Melanesian ovalocytosis, rigid erythrocytes resist merozoite invasion, and the intraerythrocytic milieu is hostile.

Nonspecific host defense mechanisms stop the infection's expansion, and the subsequent strain-specific immune response then controls the infection. Eventually, exposure to sufficient strains confers protection from high-level parasitemia and disease but not from infection. As a result of this state of infection without illness (*premunition*), asymptomatic parasitemia is common among adults and older children living in regions with stable and intense transmission (i.e., holo- or hyperendemic areas) and also in parts of low-transmission areas. Immunity is mainly specific for both the species and the strain of infecting malarial parasite. Both humoral immunity and cellular immunity are necessary for protection, but the mechanisms of each are incompletely understood (Fig. 248-1). Immune individuals have a polyclonal increase in serum levels of IgM, IgG, and IgA, although much of this antibody is unrelated to protection. Antibodies to a variety of parasitic antigens presumably act in concert to limit in vivo replication of the parasite. In the case of falciparum malaria, the most important of these antigens is the surface adhesin—the variant protein PfEMP1. Passively transferred IgG from immune adults has been shown to reduce levels of parasitemia in children. Passive transfer of maternal antibody contributes to the relative (but not complete) protection of infants from severe malaria in the first months of life. This complex immunity to disease declines when a person lives outside an endemic area for several months or longer.

Several factors retard the development of cellular immunity to malaria. These factors include the absence of major histocompatibility antigens on the surface of infected RBCs, which precludes direct T cell recognition; malaria antigen-specific immune unresponsiveness; and the enormous strain diversity of malarial parasites, along with the ability of the parasites to express variant immunodominant antigens on the erythrocyte surface that change during the course of infection. Parasites may persist in the blood for months or years (or, in the case of *P. malariae*, for decades) if treatment is not given. The complexity of the immune response in malaria, the sophistication of the parasites' evasion mechanisms, and the lack of a good in vitro correlate with clinical immunity have all slowed progress toward an effective vaccine.

CLINICAL FEATURES

Malaria is a very common cause of fever in tropical countries. The first symptoms of malaria are nonspecific; the lack of a sense of well-being, headache, fatigue, abdominal discomfort, and muscle aches followed by fever are all similar to the symptoms of a minor viral illness. In some instances, a prominence of headache, chest pain, abdominal pain, cough, arthralgia, myalgia, or diarrhea may suggest another diagnosis. Although headache may be severe in malaria, the neck stiffness and photophobia seen in meningitis do not occur. While myalgia may be prominent, it is not usually as severe as in dengue fever, and the muscles are not tender as in leptospirosis or typhus. Nausea, vomiting, and orthostatic hypotension are common. The classic malarial paroxysms, in which fever spikes, chills, and rigors occur at regular intervals, are relatively unusual and suggest infection with *P. vivax* or *P. ovale*. The fever is usually irregular at first (that of falciparum malaria may never become regular); the temperature of nonimmune individuals and children often rises above 40°C (104°F) in conjunction with tachycardia and sometimes delirium. Although childhood febrile convulsions may occur with any of the malarials, generalized seizures are specifically associated with falciparum malaria and may herald the development of encephalopathy (cerebral malaria). Many clinical abnormalities have been described in acute malaria, but most patients with uncomplicated infections have few abnormal physical findings other than fever, malaise, mild anemia, and (in some cases) a palpable spleen. Anemia is common among young children living in areas with stable transmission, particularly where resistance has compromised the efficacy

of antimalarial drugs. In nonimmune individuals with acute malaria, the spleen takes several days to become palpable, but splenic enlargement is found in a high proportion of otherwise healthy individuals in malaria-endemic areas and reflects repeated infections. Slight enlargement of the liver is also common, particularly among young children. Mild jaundice is common among adults; it may develop in patients with otherwise uncomplicated malaria and usually resolves over 1–3 weeks. Malaria is not associated with a rash like those seen in meningococcal septicemia, typhus, enteric fever, viral exanthems, and drug reactions. Petechial hemorrhages in the skin or mucous membranes—features of viral hemorrhagic fevers and leptospirosis—develop only very rarely in severe falciparum malaria.

SEVERE FALCIPARUM MALARIA

Appropriately and promptly treated, uncomplicated falciparum malaria (i.e., the patient can swallow medicines and food) carries a mortality rate of <0.1%. However, once vital-organ dysfunction occurs or the total proportion of erythrocytes infected increases to >2% (a level corresponding to >10¹² parasites in an adult), mortality risk rises steeply. The major manifestations of severe falciparum malaria are shown in Table 248-2, and features indicating a poor prognosis are listed in Table 248-3.

TABLE 248-2 MANIFESTATIONS OF SEVERE FALCIPARUM MALARIA

Signs	Manifestations
Major	
Unarousable coma/ cerebral malaria	Failure to localize or respond appropriately to noxious stimuli; coma persisting for >30 min after generalized convulsion
Acidemia/acidosis	Arterial pH of <7.25 or plasma bicarbonate level of <15 mmol/L; venous lactate level of >5 mmol/L; manifests as labored deep breathing, often termed “respiratory distress”
Severe normochromic, normocytic anemia	Hematocrit of <15% or hemoglobin level of <50 g/L (<5 g/dL) with parasitemia <10,000/ μ L
Renal failure	Serum or plasma creatinine level of >265 μ mol/L (>3 mg/dL); urine output (24 h) of <400 mL in adults or <12 mL/kg in children; no improvement with rehydration
Pulmonary edema/adult respiratory distress syndrome	Noncardiogenic pulmonary edema, often aggravated by overhydration
Hypoglycemia	Plasma glucose level of <2.2 mmol/L (<40 mg/dL)
Hypotension/shock	Systolic blood pressure of <50 mmHg in children 1–5 years or <80 mmHg in adults; core/skin temperature difference of >10°C; capillary refill >2 s
Bleeding/disseminated intravascular coagulation	Significant bleeding and hemorrhage from the gums, nose, and gastrointestinal tract and/or evidence of disseminated intravascular coagulation
Convulsions	More than two generalized seizures in 24 h; signs of continued seizure activity, sometimes subtle (e.g., tonic-clonic eye movements without limb or face movement)
Other	
Hemoglobinuria ^a	Macroscopic black, brown, or red urine; not associated with effects of oxidant drugs and red blood cell enzyme defects (such as G6PD deficiency)
Extreme weakness	Prostration; inability to sit unaided ^b
Hyperparasitemia	Parasitemia level of >5% in nonimmune patients (>10% in any patient)
Jaundice	Serum bilirubin level of >50 mmol/L (>3 mg/dL) if combined with a parasite density of 100,000/ μ L or other evidence of vital-organ dysfunction

^aHemoglobinuria may also occur in uncomplicated malaria and in patients with G6PD deficiency who take primaquine. ^bIn children who are normally able to sit.

Abbreviation: G6PD, glucose-6-phosphate dehydrogenase.

TABLE 248-3 FEATURES INDICATING A POOR PROGNOSIS IN SEVERE FALCIPARUM MALARIA

Clinical	
Marked agitation	
Hyperventilation (respiratory distress)	
Hypothermia (<36.5°C; <97.7°F)	
Bleeding	
Deep coma	
Repeated convulsions	
Anuria	
Shock	
Laboratory	
Biochemistry	
Hypoglycemia (<2.2 mmol/L)	
Hyperlactatemia (>5 mmol/L)	
Acidosis (arterial pH <7.3, serum HCO ₃ <15 mmol/L)	
Elevated serum creatinine (>265 μmol/L)	
Elevated total bilirubin (>50 μmol/L)	
Elevated liver enzymes (AST/ALT 3 times upper limit of normal)	
Elevated muscle enzymes (CPK ↑, myoglobin ↑)	
Elevated urate (>600 μmol/L)	
Hematology	
Leukocytosis (>12,000/μL)	
Severe anemia (PCV <15%)	
Coagulopathy	
Decreased platelet count (<50,000/μL)	
Prolonged prothrombin time (>3 s)	
Prolonged partial thromboplastin time	
Decreased fibrinogen (<200 mg/dL)	
Parasitology	
Hyperparasitemia	
Increased mortality at >100,000/μL	
High mortality at >500,000/μL	
>20% of parasites identified as pigment-containing trophozoites and schizonts	
>5% of neutrophils with visible pigment	

Abbreviations: ALT, alanine aminotransferase; AST, aspartate aminotransferase; CPK, creatine phosphokinase; PCV, packed cell volume.

Cerebral Malaria Coma is a characteristic and ominous feature of falciparum malaria and, despite treatment, is associated with death rates of ~20% among adults and 15% among children. Any obtundation, delirium, or abnormal behavior should be taken very seriously. The onset may be gradual or sudden following a convulsion.

Cerebral malaria manifests as diffuse symmetric encephalopathy; focal neurologic signs are unusual. Although some passive resistance to head flexion may be detected, signs of meningeal irritation are absent. The eyes may be divergent and a pout reflex is common, but other primitive reflexes are usually absent. The corneal reflexes are preserved, except in deep coma. Muscle tone may be either increased or decreased. The tendon reflexes are variable, and the plantar reflexes may be flexor or extensor; the abdominal and cremasteric reflexes are absent. Flexor or extensor posturing may be seen. On routine funduscopy, ~15% of patients have retinal hemorrhages; with pupillary dilation and indirect ophthalmoscopy, this figure increases to 30–40%. Other fundoscopic abnormalities (Fig. 248-3) include discrete spots of retinal opacification (30–60%), papilledema (8% among children, rare among adults), cotton wool spots (<5%), and decolorization of a retinal vessel or segment of vessel (occasional cases). Convulsions, usually generalized and often repeated, occur in ~10% of adults and up to 50% of children with cerebral malaria. More covert seizure activity also is common, particularly among children, and may manifest as repetitive tonic-clonic eye movements or even hypersalivation. Whereas adults



FIGURE 248-3 The eye in cerebral malaria: perimacular whitening and pale-centered retinal hemorrhages. (Courtesy of N. Beare, T. Taylor, S. Harding, S. Lewallen, and M. Molyneux; with permission.)

rarely (i.e., in <3% of cases) suffer neurologic sequelae, ~10% of children surviving cerebral malaria—especially those with hypoglycemia, severe anemia, repeated seizures, and deep coma—have residual neurologic deficits when they regain consciousness; hemiplegia, cerebral palsy, cortical blindness, deafness, and impaired cognition have been reported. The majority of these deficits improve markedly or resolve completely within 6 months. However, the prevalence of some other deficits increases over time; ~10% of children surviving cerebral malaria have a persistent language deficit. There may also be deficits in learning, planning and executive functions, attention, memory, and nonverbal functioning. The incidence of epilepsy is increased and life expectancy decreased among these children.

Hypoglycemia Hypoglycemia, an important and common complication of severe malaria, is associated with a poor prognosis and is particularly problematic in children and pregnant women. Hypoglycemia in malaria results from a failure of hepatic gluconeogenesis and an increase in the consumption of glucose by both the host and, to a much lesser extent, the malaria parasites. To compound the situation, quinine, which is still widely used for the treatment of both severe and uncomplicated falciparum malaria, is a powerful stimulant of pancreatic insulin secretion. Hyperinsulinemic hypoglycemia is especially troublesome in pregnant women receiving quinine treatment. In severe disease, the clinical diagnosis of hypoglycemia is difficult: the usual physical signs (sweating, gooseflesh, tachycardia) are absent, and the neurologic impairment caused by hypoglycemia cannot be distinguished from that caused by malaria.

Acidosis Acidosis, an important cause of death from severe malaria, results from accumulation of organic acids. Hyperlactatemia commonly coexists with hypoglycemia. In adults, coexisting renal impairment often compounds the acidosis; in children, ketoacidosis also may contribute. Other, still-unidentified organic acids are major contributors to acidosis. Acidotic breathing, sometimes called “respiratory distress,” is a sign of poor prognosis. It is followed often by circulatory failure refractory to volume expansion or inotropic drug treatment and ultimately by respiratory arrest. The plasma concentrations of bicarbonate or lactate are the best biochemical prognosticators in severe malaria. Hypovolemia is not a major contributor to acidosis. Lactic acidosis is caused by the combination of anaerobic glycolysis in tissues where sequestered parasites interfere with microcirculatory flow,

1374 lactate production by the parasites, and a failure of hepatic and renal lactate clearance. The prognosis of severe acidosis is poor.

Noncardiogenic Pulmonary Edema Adults with severe falciparum malaria may develop noncardiogenic pulmonary edema even after several days of antimalarial therapy. The pathogenesis of this variant of the adult respiratory distress syndrome is unclear. The mortality rate is >80%. This condition can be aggravated by overly vigorous administration of IV fluid. Noncardiogenic pulmonary edema can also develop in otherwise uncomplicated vivax malaria, where recovery is usual.

Renal Impairment Acute kidney injury is common in severe falciparum malaria, but oliguric renal failure is rare among children. The pathogenesis of renal failure is unclear but may be related to erythrocyte sequestration and agglutination interfering with renal microcirculatory flow and metabolism. Clinically and pathologically, this syndrome manifests as acute tubular necrosis. Renal cortical necrosis never develops. Acute renal failure may occur simultaneously with other vital-organ dysfunction (in which case the mortality risk is high) or may progress as other disease manifestations resolve. In survivors, urine flow resumes in a median of 4 days, and serum creatinine levels return to normal in a mean of 17 days (Chap. 334). Early dialysis or hemofiltration considerably enhances the likelihood of a patient's survival, particularly in acute hypercatabolic renal failure.

Hematologic Abnormalities Anemia results from accelerated RBC removal by the spleen, obligatory RBC destruction at parasite schizogony, and ineffective erythropoiesis. In severe malaria, both infected and uninfected RBCs show reduced deformability, which correlates with prognosis and development of anemia. Splenic clearance of all RBCs is increased. In nonimmune individuals and in areas with unstable transmission, anemia can develop rapidly and transfusion is often required. As a consequence of repeated malarial infections, children in many areas of Africa and on the island of New Guinea may develop severe anemia resulting from both shortened survival of uninfected RBCs and marked dyserythropoiesis. Anemia is a common consequence of antimalarial drug resistance, which results in repeated or continued infection.

Slight coagulation abnormalities are common in falciparum malaria, and mild thrombocytopenia is usual (a normal platelet count should raise questions about the diagnosis of malaria). Of patients with severe malaria, <5% have significant bleeding with evidence of disseminated intravascular coagulation. Hematemesis from stress ulceration or acute gastric erosions also may occur rarely.

Liver Dysfunction Mild hemolytic jaundice is common in malaria. Severe jaundice is associated with *P. falciparum* infections; is more common among adults than among children; and results from hemolysis, hepatocyte injury, and cholestasis. When accompanied by other vital-organ dysfunction (often renal impairment), liver dysfunction carries a poor prognosis. Hepatic dysfunction contributes to hypoglycemia, lactic acidosis, and impaired drug metabolism. Occasional patients with falciparum malaria may develop deep jaundice (with hemolytic, hepatic, and cholestatic components) without evidence of other vital-organ dysfunction, in which case the prognosis is good.

Other Complications HIV/AIDS and malnutrition predispose to more severe malaria in nonimmune individuals; malaria anemia is worsened by concurrent infections with intestinal helminths, hookworm in particular. Septicemia may complicate severe malaria, particularly in children. Differentiating severe malaria from sepsis with incidental parasitemia in childhood is very difficult. In endemic areas, *Salmonella* bacteremia has been associated specifically with *P. falciparum* infections. Chest infections and catheter-induced urinary tract infections are common among patients who are unconscious for >3 days. Aspiration pneumonia may follow generalized convulsions. The frequencies of complications of severe falciparum malaria are summarized in Table 248-4.

MALARIA IN PREGNANCY

Malaria in early pregnancy causes abortion. In areas of high malaria transmission, falciparum malaria in primi- and secundigravid women

TABLE 248-4 RELATIVE INCIDENCE OF SEVERE COMPLICATIONS OF FALCIPARUM MALARIA

Complication	Nonpregnant Adults	Pregnant Women	Children
Anemia	+	++	+++
Convulsions	+	+	+++
Hypoglycemia	+	+++	+++
Jaundice	+++	+++	+
Renal failure	+++	+++	-
Pulmonary edema	++	+++	+

Note: -, rare; +, infrequent; ++, frequent; +++, very frequent.

is associated with low birth weight (average reduction, ~170 g) and consequently increased infant mortality rates. In general, infected mothers in areas of stable transmission remain asymptomatic despite intense accumulation of parasitized erythrocytes in the placental microcirculation. Maternal HIV infection predisposes pregnant women to more frequent and higher-density malaria infections, predisposes their newborns to congenital malarial infection, and exacerbates the reduction in birth weight associated with malaria.

In areas with unstable transmission of malaria, pregnant women are prone to severe infections and are particularly vulnerable to high parasitemias with anemia, hypoglycemia, and acute pulmonary edema. Fetal distress, premature labor, and stillbirth or low birth weight are common results. Fetal death is usual in severe malaria. Congenital malaria occurs in <5% of newborns whose mothers are infected; its frequency and the level of parasitemia are related directly to the parasite density in maternal blood and in the placenta. *P. vivax* malaria in pregnancy is also associated with a reduction in birth weight (average, 110 g), but, in contrast to the situation in falciparum malaria, this effect is more pronounced in multigravid than in primigravid women. About 350,000 women die in childbirth yearly, with most deaths occurring in low-income countries; maternal death from hemorrhage at childbirth is correlated with malaria-induced anemia.

MALARIA IN CHILDREN

Most of the 660,000 persons who die of falciparum malaria each year are young African children. Convulsions, coma, hypoglycemia, metabolic acidosis, and severe anemia are relatively common among children with severe malaria, whereas deep jaundice, oliguric acute kidney injury, and acute pulmonary edema are unusual. Severely anemic children may present with labored deep breathing, which in the past has been attributed incorrectly to "anemic congestive cardiac failure" but in fact is usually caused by metabolic acidosis, often compounded by hypovolemia. In general, children tolerate antimalarial drugs well and respond rapidly to treatment.

TRANSFUSION MALARIA

Malaria can be transmitted by blood transfusion, needle-stick injury, sharing of needles by infected injection drug users, or organ transplantation. The incubation period in these settings is often short because there is no preerythrocytic stage of development. The clinical features and management of these cases are the same as for naturally acquired infections. Radical chemotherapy with primaquine is unnecessary for transfusion-transmitted *P. vivax* and *P. ovale* infections.

CHRONIC COMPLICATIONS OF MALARIA

TROPICAL SPLENOMEGALY (HYPERREACTIVE MALARIAL SPLENOMEGALY)

Chronic or repeated malarial infections produce hypergammaglobulinemia; normochromic, normocytic anemia; and, in certain situations, splenomegaly. Some residents of malaria-endemic areas in tropical Africa and Asia exhibit an abnormal immunologic response to repeated infections that is characterized by massive splenomegaly, hepatomegaly, marked elevations in serum titers of IgM and malarial antibody, hepatic sinusoidal lymphocytosis, and (in Africa) peripheral B cell lymphocytosis. This syndrome has been associated with

the production of cytotoxic IgM antibodies to CD8+ T lymphocytes, antibodies to CD5+ T lymphocytes, and an increase in the ratio of CD4+ to CD8+ T cells. These events may lead to uninhibited B cell production of IgM and the formation of cryoglobulins (IgM aggregates and immune complexes). This immunologic process stimulates reticuloendothelial hyperplasia and clearance activity and eventually produces splenomegaly. Patients with hyperreactive malarial splenomegaly present with an abdominal mass or a dragging sensation in the abdomen and occasional sharp abdominal pains suggesting perisplenitis. Anemia and some degree of pancytopenia are usually evident, and in some cases malarial parasites cannot be found in peripheral-blood smears. Vulnerability to respiratory and skin infections is increased; many patients die of overwhelming sepsis. Persons with hyperreactive malarial splenomegaly who are living in endemic areas should receive antimalarial chemoprophylaxis; the results are usually good. In nonendemic areas, antimalarial treatment is advised. In some cases refractory to therapy, clonal lymphoproliferation may develop and can then evolve into a malignant lymphoproliferative disorder.

QUARTAN MALARIAL NEPHROPATHY

Chronic or repeated infections with *P. malariae* (and possibly with other malarial species) may cause soluble immune complex injury to the renal glomeruli, resulting in the nephrotic syndrome. Other unidentified factors must contribute to this process since only a very small proportion of infected patients develop renal disease. The histologic appearance is that of focal or segmental glomerulonephritis with splitting of the capillary basement membrane. Subendothelial dense deposits are seen on electron microscopy, and immunofluorescence reveals deposits of complement and immunoglobulins; in samples of renal tissue from children, *P. malariae* antigens are often visible. A coarse-granular pattern of basement membrane immunofluorescent deposits (predominantly IgG3) with selective proteinuria carries a better prognosis than a fine-granular, predominantly IgG2 pattern with nonselective proteinuria. Quartan nephropathy usually responds poorly to treatment with either antimalarial agents or glucocorticoids and cytotoxic drugs.

BURKITT'S LYMPHOMA AND EPSTEIN-BARR VIRUS INFECTION

It is possible that malaria-related immune dysregulation provokes infection with lymphoma viruses. Burkitt's lymphoma is strongly associated with Epstein-Barr virus. The prevalence of this childhood tumor is high in malarious areas of Africa.

DIAGNOSIS

DEMONSTRATION OF THE PARASITE

The diagnosis of malaria rests on the demonstration of asexual forms of the parasite in stained peripheral-blood smears. After a negative blood smear, repeat smears should be made if there is a high degree of suspicion. Of the Romanowsky stains, Giemsa at pH 7.2 is preferred; Field's, Wright's, or Leishman's stain can also be used. Both thin (Figs. 248-4 and 248-5; see also Figs. 250e-3 and 250e-4) and thick (Figs. 248-6, 248-7, 248-8, and 248-9) blood smears should be examined. The thin blood smear should be rapidly air-dried, fixed in anhydrous methanol, and stained; the RBCs in the tail of the film should then be examined under oil immersion ($\times 1000$ magnification). The level of parasitemia is expressed as the number of parasitized erythrocytes per 1000 RBCs. The thick blood film should be of uneven thickness. The smear should be dried thoroughly and stained without fixing. As many layers of erythrocytes overlies one another and are lysed during the staining procedure, the thick film has the advantage of concentrating the parasites (by 40- to 100-fold compared with a thin blood film) and thus increasing diagnostic sensitivity. Both parasites and white blood cells (WBCs) are counted, and the number of parasites per unit volume is calculated from the total leukocyte count. Alternatively, a WBC count of 8000/ μL is assumed. This figure is converted to the number of parasitized erythrocytes per microliter. A minimum of 200 WBCs should be counted under oil immersion. Interpretation of blood smear films requires some experience because artifacts are common. Before a thick smear is judged to be negative, 100–200 fields should be examined under oil immersion. In high-transmission areas, the presence of up to 10,000 parasites/ μL of blood may be tolerated without symptoms or signs in partially immune individuals. Thus in these areas the detection of malaria parasites is sensitive but has low specificity in identifying

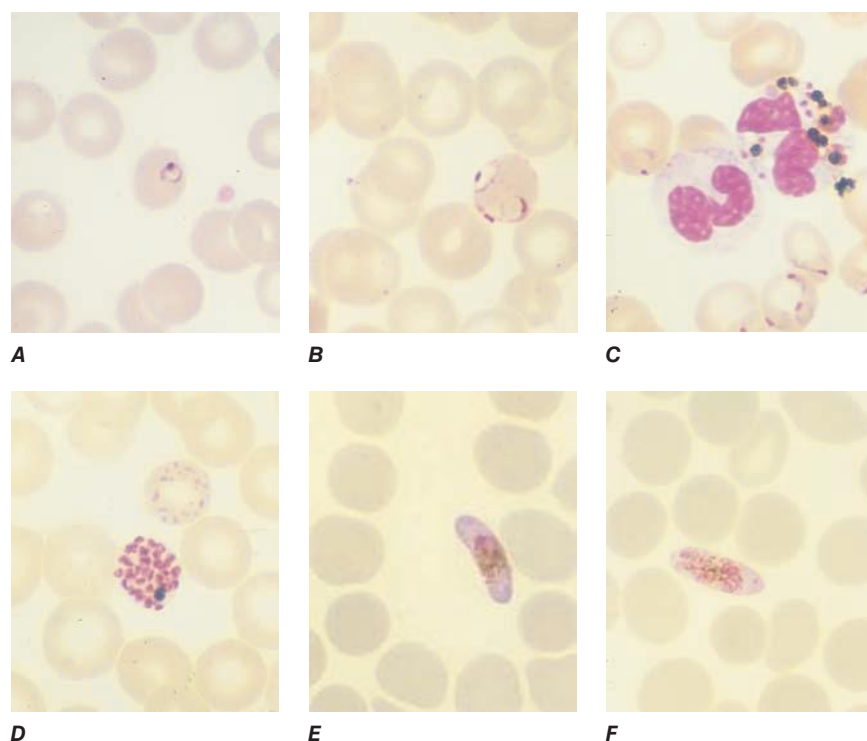


FIGURE 248-4 Thin blood films of *Plasmodium falciparum*. **A.** Young trophozoites. **B.** Old trophozoites. **C.** Pigment in polymorphonuclear cells and trophozoites. **D.** Mature schizonts. **E.** Female gametocytes. **F.** Male gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

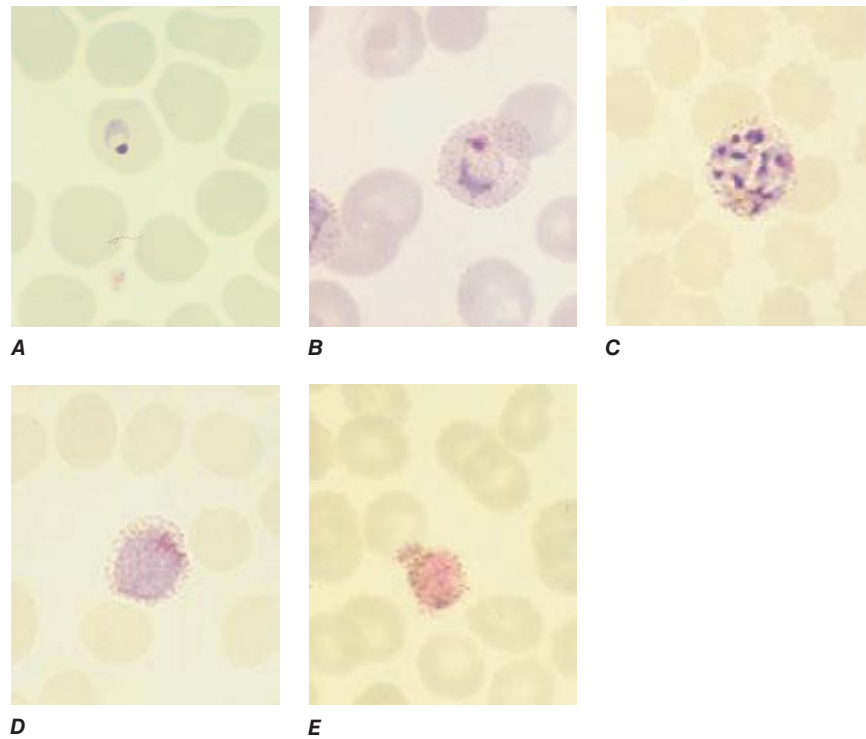


FIGURE 248-5 Thin blood films of *Plasmodium vivax*. **A.** Young trophozoites. **B.** Old trophozoites. **C.** Mature schizonts. **D.** Female gametocytes. **E.** Male gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

malaria as the cause of illness. Low-density parasitemia is common in other conditions causing fever.

Rapid, simple, sensitive, and specific antibody-based diagnostic stick or card tests that detect *P. falciparum*-specific, histidine-rich protein 2 (PfHRP2), lactate dehydrogenase, or aldolase antigens in finger-prick blood samples are now being used widely in control programs (Table 248-5). Some of these rapid diagnostic tests carry a second antibody, which allows falciparum malaria to be distinguished from the less dangerous malarias. PfHRP2-based tests may remain positive for several weeks after acute infection. This feature is a disadvantage in high-transmission areas where infections are frequent, but it is of value in the diagnosis of severe malaria in patients who have taken antimalarial drugs and cleared peripheral parasitemia (but in whom the PfHRP2 test remains strongly positive). Rapid diagnostic tests are replacing microscopy in many areas because of their simplicity and speed. Their disadvantage is that they do not quantify parasitemia.

The relationship between parasitemia and prognosis is complex; in general, patients with $>10^5$ parasites/ μL are at increased risk of dying, but nonimmune patients may die with much lower counts, and partially immune persons may tolerate parasitemia levels many times higher with only minor symptoms. In severe malaria, a poor prognosis is indicated by a predominance of more mature *P. falciparum*

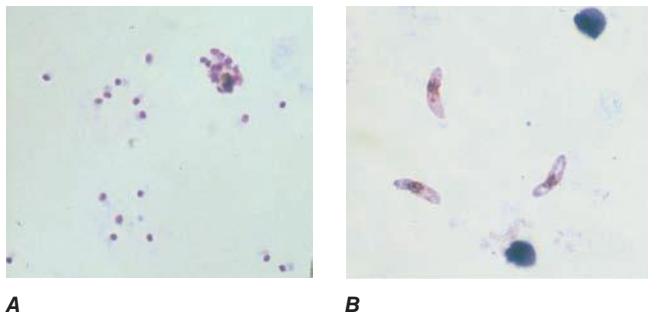


FIGURE 248-6 Thick blood films of *Plasmodium falciparum*. **A.** Trophozoites. **B.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

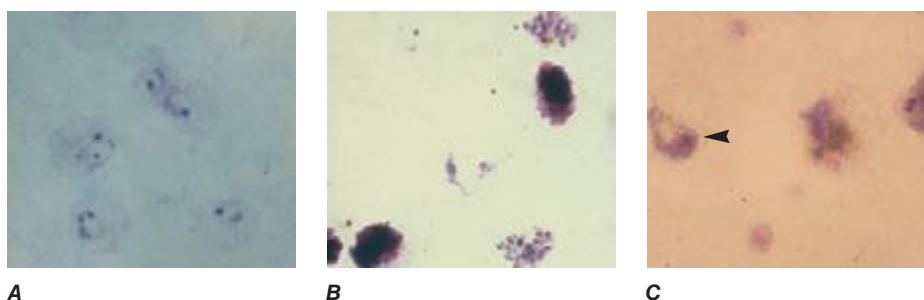


FIGURE 248-7 Thick blood films of *Plasmodium vivax*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

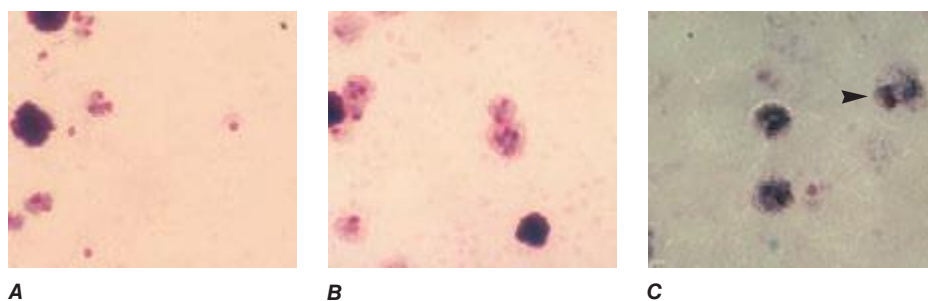


FIGURE 248-8 Thick blood films of *Plasmodium ovale*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections, 2nd ed*, with the permission of the World Health Organization.)

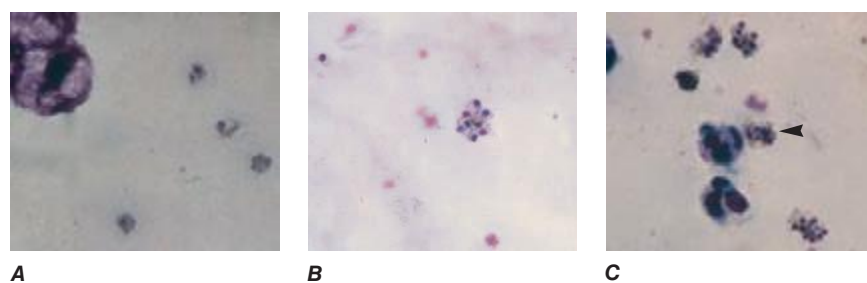


FIGURE 248-9 Thick blood films of *Plasmodium malariae*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections, 2nd ed*, with the permission of the World Health Organization.)

TABLE 248-5 STANDARD METHODS FOR THE DIAGNOSIS OF MALARIA^a

Method	Procedure	Advantages	Disadvantages
Thick blood film ^b	Blood should be uneven in thickness but thin enough that the hands of a watch can be read through part of the spot. Stain dried, unfixed blood spot with Giemsa, Field's, or another Romanowsky stain. Count number of asexual parasites per 200 WBCs (or per 500 at low densities). Count gametocytes separately. ^c	Sensitive (0.001% parasitemia); species specific; inexpensive	Requires experience (artifacts may be misinterpreted as low-level parasitemia); underestimates true count
Thin blood film ^d	Stain fixed smear with Giemsa, Field's, or another Romanowsky stain. Count number of RBCs containing asexual parasites per 1000 RBCs. In severe malaria, assess stage of parasite development and count neutrophils containing malaria pigment. ^e Count gametocytes separately. ^c	Rapid; species specific; inexpensive; in severe malaria, provides prognostic information ^f	Insensitive (<0.05% parasitemia); uneven distribution of <i>P. vivax</i> , as enlarged infected red cells concentrate at leading edge
PfHRP2 dipstick or card test	A drop of blood is placed on the stick or card, which is then immersed in washing solutions. Monoclonal antibody capture of parasitic antigens reads out as a colored band.	Robust and relatively inexpensive; rapid; sensitivity similar to or slightly lower than that of thick films (~0.001% parasitemia)	Detects only <i>Plasmodium falciparum</i> ; remains positive for weeks after infection ^g ; does not quantitate <i>P. falciparum</i> parasitemia
<i>Plasmodium</i> LDH dipstick or card test	A drop of blood is placed on the stick or card, which is then immersed in washing solutions. Monoclonal antibody capture of parasitic antigens reads out as two colored bands. One band is genus specific (all malarias), and the other is specific for <i>P. falciparum</i> .	Rapid; sensitivity similar to or slightly lower than that of thick films for <i>P. falciparum</i> (~0.001% parasitemia)	Slightly more difficult preparation than PfHRP2 tests; may miss low-level parasitemia with <i>P. vivax</i> , <i>P. ovale</i> , and <i>P. malariae</i> and may not speciate these organisms; does not quantitate <i>P. falciparum</i> parasitemia
Microtube concentration methods with acridine orange staining	Blood is collected in a specialized tube containing acridine orange, anticoagulant, and a float. After centrifugation, which concentrates the parasitized cells around the float, fluorescence microscopy is performed.	Sensitivity similar or superior to that of thick films (~0.001% parasitemia); ideal for processing large numbers of samples rapidly	Does not speciate or quantitate; requires fluorescence microscopy

^aMalaria cannot be diagnosed clinically with accuracy, but treatment should be started on clinical grounds if laboratory confirmation is likely to be delayed. In areas of the world where malaria is endemic and transmission is high, low-level asymptomatic parasitemia is common in otherwise healthy people. Thus malaria may not be the cause of a fever, although in this context the presence of >10,000 parasites/ μ L (~0.2% parasitemia) does indicate that malaria is the cause. Antibody and polymerase chain reaction tests have no role in the diagnosis of malaria except that PCR is increasingly used for genotyping and speciation in mixed infections and for detection of low-level parasitemias in asymptomatic residents of endemic areas. ^bAsexual parasites/200 WBCs \times 40 = parasite count/ μ L (assumes a WBC count of 8000/ μ L). See Figs. 248-6 through 248-9. ^c*P. falciparum* gametocytemia may persist for days or weeks after clearance of asexual parasites. Gametocytemia without asexual parasitemia does not indicate active infection. ^dParasitized RBCs (%) \times hematocrit \times 1256 = parasite count/ μ L. See Figs. 248-4 and 248-5. ^eThe presence of >100,000 parasites/ μ L (~2% parasitemia) is associated with an increased risk of severe malaria, but some patients have severe malaria with lower counts. At any level of parasitemia, the finding that >50% of parasites are tiny rings (cytoplasm thickness less than half of nucleus width) carries a relatively good prognosis. The presence of visible pigment in >20% of parasites or of phagocytosed pigment in >5% of polymorphonuclear leukocytes (indicating massive recent schizogony) carries a worse prognosis. ^fPersistence of PfHRP2 is a disadvantage in high-transmission settings, where many asymptomatic people have positive tests, but can be used to diagnostic advantage in low-transmission settings when a sick patient has previously received unknown treatment (which, in endemic areas, often consists of antimalarial drugs). A positive PfHRP2 test indicates that the illness is falciparum malaria, even if the blood smear is negative.

Abbreviations: LDH, lactate dehydrogenase; PfHRP2, *P. falciparum* histidine-rich protein 2; RBCs, red blood cells; WBCs, white blood cells.

1378 parasites (i.e., >20% of parasites with visible pigment) in the peripheral-blood film or by the presence of phagocytosed malarial pigment in >5% of neutrophils. In *P. falciparum* infections, gametocytemia peaks 1 week after the peak of asexual parasites. Because the mature gametocytes of *P. falciparum* (unlike those of other plasmodia) are not affected by most antimalarial drugs, their persistence does not constitute evidence of drug resistance. Phagocytosed malarial pigment is sometimes seen inside peripheral-blood monocytes or polymorphonuclear leukocytes and may provide a clue to recent infection if malaria parasites are not detectable. After the clearance of the parasites, this intraphagocytic malarial pigment is often evident for several days in the peripheral blood films or for longer in bone marrow aspirates or smears of fluid expressed after intradermal puncture. Staining of parasites with the fluorescent dye acridine orange allows more rapid diagnosis of malaria (but not speciation of the infection) in patients with low-level parasitemia.

Molecular diagnosis by polymerase chain reaction (PCR) amplification of parasite nucleic acid is more sensitive than microscopy or rapid diagnostic tests for detecting malaria parasites and defining malarial species. While currently impractical in the standard clinical setting, PCR is used in reference centers in endemic areas. In epidemiologic surveys, sensitive PCR detection may prove very useful in identifying asymptomatic infections as control and eradication programs drive parasite prevalence down to very low levels. Serologic diagnosis with either indirect fluorescent antibody or enzyme-linked immunosorbent assays may prove useful as measures of transmission intensity in future epidemiologic studies. Serology has no place in the diagnosis of acute illness.

LABORATORY FINDINGS

Normochromic, normocytic anemia is usual. The leukocyte count is generally normal, although it may be raised in very severe infections. There is slight monocytosis, lymphopenia, and eosinopenia, with reactive lymphocytosis and eosinophilia in the weeks after the acute infection. The erythrocyte sedimentation rate, plasma viscosity, and levels of C-reactive protein and other acute-phase proteins are high. The platelet count is usually reduced to $\sim 10^5/\mu\text{L}$. Severe infections may be accompanied by prolonged prothrombin and partial thromboplastin times and by more severe thrombocytopenia. Levels of antithrombin III are reduced even in mild infection. In uncomplicated malaria, plasma concentrations of electrolytes, blood urea nitrogen (BUN), and creatinine are usually normal. Findings in severe malaria may include metabolic acidosis, with low plasma concentrations of glucose, sodium, bicarbonate, calcium, phosphate, and albumin together with elevations in lactate, BUN, creatinine, urate, muscle and liver enzymes, and conjugated and unconjugated bilirubin. Hypergammaglobulinemia is usual in immune and semi-immune subjects. Urinalysis generally gives normal results. In adults and children with cerebral malaria, the mean cerebrospinal fluid (CSF) opening pressure at lumbar puncture is ~ 160 mm; usually the CSF content is normal or there is a slight elevation of total protein level (<1.0 g/L [<100 mg/dL]) and cell count ($<20/\mu\text{L}$).

TREATMENT MALARIA

(Table 248-6) When a patient in or from a malarious area presents with fever, thick and thin blood smears should be prepared and examined immediately to confirm the diagnosis and identify the species of infecting parasite (Figs. 248-4 through 248-9). Repeat blood smears should be performed at least every 12–24 h for 2 days if the first smears are negative and malaria is strongly suspected. Alternatively, a rapid antigen detection card or stick test should be performed. Patients with severe malaria or those unable to take oral drugs should receive parenteral antimalarial therapy. If there is any doubt about the resistance status of the infecting organism, it should be considered resistant. Antimalarial drug susceptibility testing can be performed but is rarely available, has poor predictive value in an individual case, and yields results too slowly to influence the choice of treatment. Several drugs are available for oral treatment. The choice of drug depends on the likely sensitivity of

TABLE 248-6 REGIMENS FOR THE TREATMENT OF MALARIA^a

Type of Disease or Treatment	Regimen(s)
Uncomplicated Malaria	
Known chloroquine-sensitive strains of <i>Plasmodium vivax</i> , <i>P. malariae</i> , <i>P. ovale</i> , <i>P. knowlesi</i> , <i>P. falciparum</i> ^b	Chloroquine (10 mg of base/kg stat followed by 5 mg/kg at 12, 24, and 36 h or by 10 mg/kg at 24 h and 5 mg/kg at 48 h) or Amodiaquine (10–12 mg of base/kg qd for 3 days)
Radical treatment for <i>P. vivax</i> or <i>P. ovale</i> infection	In addition to chloroquine or amodiaquine as detailed above, primaquine (0.5 mg of base/kg qd in tropical regions and 0.25 mg/kg for temperate-origin <i>P. vivax</i>) should be given for 14 days to prevent relapse. In mild G6PD deficiency, 0.75 mg of base/kg should be given once weekly for 8 weeks. Primaquine should not be given in severe G6PD deficiency.
Sensitive <i>P. falciparum</i> malaria ^c	Artesunate ^d (4 mg/kg qd for 3 days) plus sulfadoxine (25 mg/kg)/pyrimethamine (1.25 mg/kg) as a single dose or Artesunate ^d (4 mg/kg qd for 3 days) plus amodiaquine (10 mg of base/kg qd for 3 days) ^e
Multidrug-resistant <i>P. falciparum</i> malaria	Either artemether-lumefantrine ^d (1.5/9 mg/kg bid for 3 days with food) or Artesunate ^d (4 mg/kg qd for 3 days) plus mefloquine (24–25 mg of base/kg—either 8 mg/kg qd for 3 days or 15 mg/kg on day 2 and then 10 mg/kg on day 3) ^e or Dihydroartemisinin-piperaquine ^d (2.5/20 mg/kg qd for 3 days)
Second-line treatment/treatment of imported malaria	Either artesunate ^d (2 mg/kg qd for 7 days) or quinine (10 mg of salt/kg tid for 7 days) plus 1 of the following 3: 1. Tetracycline ^f (4 mg/kg qid for 7 days) 2. Doxycycline ^f (3 mg/kg qd for 7 days) 3. Clindamycin (10 mg/kg bid for 7 days) or Atovaquone-proguanil (20/8 mg/kg qd for 3 days with food)
Severe Falciparum Malaria^g	
	Artesunate ^d (2.4 mg/kg stat IV followed by 2.4 mg/kg at 12 and 24 h and then daily if necessary) ^h or, if unavailable, Artemether ^d (3.2 mg/kg stat IM followed by 1.6 mg/kg qd) or, if unavailable, Quinine dihydrochloride (20 mg of salt/kg ⁱ infused over 4 h, followed by 10 mg of salt/kg infused over 2–8 h q8h) or, if unavailable, Quinidine (10 mg of base/kg ^j infused over 1–2 h, followed by 1.2 mg of base/kg per hour ^k with electrocardiographic monitoring)

^aIn endemic areas, except in pregnant women and infants, a single dose of primaquine (0.25 mg of base/kg) should be added as a gametocytocidal to all falciparum malaria treatments to prevent transmission. This addition is considered safe even in G6PD deficiency. ^bVery few areas now have chloroquine-sensitive *P. falciparum* malaria (Fig. 248-2). ^cIn areas where the partner drug to artesunate is known to be effective. ^dArtemisinin derivatives are not readily available in some temperate countries. ^eFixed-dose coformulated combinations are available. The World Health Organization now recommends artemisinin combination regimens as first-line therapy for falciparum malaria in all tropical countries and advocates use of fixed-dose combinations. ^fTetracycline and doxycycline should not be given to pregnant women or to children <8 years of age. ^gOral treatment should be substituted as soon as the patient recovers sufficiently to take fluids by mouth. ^hArtesunate is the drug of choice when available. The doses in children weighing <20 kg should be 3 mg/kg. The data from large studies in Southeast Asia showed a 35% lower mortality rate than with quinine, and very large studies in Africa showed a 22.5% reduction in mortality rate compared with quinine. ⁱA loading dose should not be given if therapeutic doses of quinine or quinidine have definitely been administered in the previous 24 h. Some authorities recommend a lower dose of quinidine. ^jInfusions can be given in 0.9% saline and 5–10% dextrose in water. Infusion rates for quinine and quinidine should be carefully controlled.

Abbreviation: G6PD, glucose-6-phosphate dehydrogenase.

the infecting parasites. Despite increasing evidence of chloroquine resistance in *P. vivax* (from parts of Indonesia, Oceania, eastern and southern Asia, and Central and South America), chloroquine remains a first-line treatment for the non-falciparum malarias (*P. vivax*, *P. ovale*, *P. malariae*, *P. knowlesi*) except in Indonesia and Papua New Guinea, where high levels of resistance in *P. vivax* are prevalent.

The treatment of falciparum malaria has changed radically in recent years. In all endemic areas, the World Health Organization (WHO) now recommends artemisinin-based combinations (ACTs) as first-line treatment for uncomplicated falciparum malaria. These combinations are also highly effective for the other malarias. These rapidly and reliably effective drugs are sometimes unavailable in temperate countries, where treatment recommendations are limited by the registered available drugs. Fake or substandard antimalarials are commonly sold in many Asian and African countries. Thus, careful attention is required at the time of purchase and later, especially if the patient fails to respond as expected. Characteristics of antimalarial drugs are shown in [Table 248-7](#).

SEVERE MALARIA

In large studies, parenteral artesunate, a water-soluble artemisinin derivative, has reduced mortality rates in severe falciparum malaria among Asian adults and children by 35% and among African children by 22.5% compared with mortality rates with quinine treatment. Artesunate has therefore become the drug of choice for all patients with severe malaria everywhere. Artesunate is given by IV injection but can also be given by IM injection. Artemether and the closely related drug artemotil (arteether) are oil-based formulations given by IM injection; they are erratically absorbed and do not confer the same survival benefit as artesunate. A rectal formulation of artesunate has been developed as a community-based pre-referral treatment for patients in the rural tropics who cannot take oral medications. Pre-referral administration of rectal artesunate has been shown to decrease mortality risk among severely ill children in communities without access to immediate parenteral treatment. Although the artemisinin compounds are safer than quinine and considerably safer than quinidine, only one formulation is available in the United States. IV artesunate has been approved by the U.S. Food and Drug Administration for emergency use against severe malaria and can be obtained through the Centers for Disease Control and Prevention (CDC) Drug Service (see end of chapter for contact information). The antiarrhythmic quinidine gluconate is as effective as quinine and, as it was more readily available, replaced quinine for the treatment of malaria in the United States. The administration of quinidine must be closely monitored if dysrhythmias and hypotension are to be avoided. If total plasma levels exceed 8 µg/mL or the QT_c interval exceeds 0.6 s or the QRS complex widens by more than 25% of baseline, then infusion rates should be slowed or infusion stopped temporarily. If arrhythmia or saline-unresponsive hypotension develops, treatment with this drug should be discontinued. Quinine is safer than quinidine; cardiovascular monitoring is not required except when the recipient has cardiac disease.

Severe falciparum malaria constitutes a medical emergency requiring intensive nursing care and careful management. The patient should be weighed and, if comatose, placed on his or her side. Frequent evaluation of the patient's condition is essential. Adjunctive treatments such as high-dose glucocorticoids, urea, heparin, dextran, desferrioxamine, antibody to tumor necrosis factor α, high-dose phenobarbital (20 mg/kg), mannitol, or large-volume fluid or albumin boluses have proved either ineffective or harmful in clinical trials and should not be used. In acute renal failure or severe metabolic acidosis, hemofiltration or hemodialysis should be started as early as possible.

In severe malaria, parenteral antimalarial treatment should be started immediately. Artesunate, given by either IV or IM injection, is the agent of choice; it is simple to administer, safe, and rapidly effective. It does not require dose adjustments in liver dysfunction or renal failure, and it should be used in pregnant women with severe malaria. If artesunate is unavailable and artemether, quinine,

or quinidine is used, an initial loading dose must be given so that therapeutic concentrations are reached as soon as possible. Both quinine and quinidine will cause dangerous hypotension if injected rapidly; when given IV, they must be administered carefully by rate-controlled infusion only. If this approach is not possible, quinine may be given by deep IM injections into the anterior thigh. The optimal therapeutic range for quinine and quinidine in severe malaria is not known with certainty, but total plasma concentrations of 8–15 mg/L for quinine and 3.5–8.0 mg/L for quinidine are effective and do not cause serious toxicity. The systemic clearance and apparent volume of distribution of these alkaloids are markedly reduced and plasma protein binding is increased in severe malaria, so that the blood concentrations attained with a given dose are higher. If the patient remains seriously ill or in acute renal failure for >2 days, maintenance doses of quinine or quinidine should be reduced by 30–50% to prevent toxic accumulation of the drug. The initial doses should never be reduced. If safe and feasible, exchange transfusion may be considered for patients with severe malaria, although the precise indications for this procedure have not been agreed upon and there is no clear evidence that this measure is beneficial, particularly with artesunate treatment. Convulsions should be treated promptly with IV (or rectal) benzodiazepines. The role of prophylactic anticonvulsants in children is uncertain. If respiratory support is not available, then a full loading dose of phenobarbital (20 mg/kg) to prevent convulsions should not be given as it may cause respiratory arrest.

When the patient is unconscious, the blood glucose level should be measured every 4–6 h. All patients should receive a continuous infusion of dextrose, and blood concentrations ideally should be maintained above 4 mmol/L. Hypoglycemia (<2.2 mmol/L or 40 mg/dL) should be treated immediately with bolus glucose. The parasite count and hematocrit level should be measured every 6–12 h. Anemia develops rapidly; if the hematocrit falls to <20%, then whole blood (preferably fresh) or packed cells should be transfused slowly, with careful attention to circulatory status. Renal function should be checked daily. Children presenting with severe anemia and acidotic breathing require immediate blood transfusion. Accurate assessment is vital. Management of fluid balance is difficult in severe malaria, particularly in adults, because of the thin dividing line between overhydration (leading to pulmonary edema) and underhydration (contributing to renal impairment). As soon as the patient can take fluids, oral therapy should be substituted for parenteral treatment.

UNCOMPLICATED MALARIA

Infections due to sensitive strains of *P. vivax*, *P. knowlesi*, *P. malariae*, and *P. ovale* should be treated with oral chloroquine (total dose, 25 mg of base/kg) or with an ATC known to be efficacious. In much of the tropics, drug-resistant *P. falciparum* has been increasing in distribution, frequency, and intensity. It is now accepted that, to prevent resistance, falciparum malaria should be treated with drug combinations and not with single drugs in endemic areas; the same rationale has been applied successfully to the treatment of tuberculosis, HIV/AIDS, and cancers. This combination strategy is based on simultaneous use of two or more drugs with different modes of action. ACT regimens are now recommended as first-line treatment for falciparum malaria throughout the malaria-affected world. These regimens are safe and effective in adults, children, and after the first trimester of pregnancy (uncertainty regarding safety currently precludes their use in the first trimester). The rapidly eliminated artemisinin component is usually an artemisinin derivative (artesunate, artemether, or dihydroartemisinin) given for 3 days, and the partner drug is usually a more slowly eliminated antimalarial to which *P. falciparum* is sensitive. Five ACT regimens are currently recommended by the WHO. In areas with multidrug-resistant falciparum malaria (parts of Asia and South America, including those with mefloquine-resistant parasites; [Fig. 248-10](#)), artemether-lumefantrine, artesunate-mefloquine, or dihydroartemisinin-piperazine should be used; these regimens provide cure rates of >90%. In areas with sensitive parasites, the aforementioned combinations, artesunate-sulfadoxine-pyrimethamine, or artesunate-amodiaquine also may be

TABLE 248-7 PROPERTIES OF ANTIMALARIAL DRUGS

Drug(s)	Pharmacokinetic Properties	Antimalarial Activity	Minor Toxicity	Major Toxicity
Quinine, quinidine	Good oral and IM absorption (quinine); Cl and V_d reduced, but plasma protein binding (principally to ∞ 1 acid glycoprotein) increased (90%) in malaria; quinine $t_{1/2}$: 16 h in malaria, 11 h in healthy persons; quinidine $t_{1/2}$: 13 h in malaria, 8 h in healthy persons	Acts mainly on trophozoite blood stage; kills gametocytes of <i>P. vivax</i> , <i>P. ovale</i> , and <i>P. malariae</i> (but not <i>P. falciparum</i>); no action on liver stages	Common: "Cinchonism": tinnitus, high-tone hearing loss, nausea, vomiting, dysphoria, postural hypotension; ECG QT_c interval prolongation (quinine usually by <10% but quinidine by up to 25%) Rare: Diarrhea, visual disturbance, rashes Note: Very bitter taste	Common: Hypoglycemia Rare: Hypotension, blindness, deafness, cardiac arrhythmias, thrombocytopenia, hemolysis, hemolytic-uremic syndrome, vasculitis, cholestatic hepatitis, neuromuscular paralysis Note: Quinidine more cardiotoxic
Chloroquine	Good oral absorption, very rapid IM and SC absorption; complex pharmacokinetics; enormous Cl and V_d (unaffected by malaria); blood concentration profile determined by distribution processes in malaria; $t_{1/2}$: 1–2 months	As for quinine but acts slightly earlier in asexual cycle	Common: Nausea, dysphoria, pruritus in dark-skinned patients, postural hypotension, slight ECG QT_c prolongation Rare: Accommodation difficulties, keratopathy, rash Note: Bitter taste, well tolerated	Acute: Hypotensive shock (parenteral), cardiac arrhythmias, neuropsychiatric reactions Chronic: Retinopathy (cumulative dose, >100 g), skeletal and cardiac myopathy
Piperaquine	Adequate oral absorption, may be enhanced by fats; similar pharmacokinetics to chloroquine; $t_{1/2}$: 21–28 days	As for chloroquine, but retains activity against multi-drug-resistant <i>P. falciparum</i>	Epigastric pain, diarrhea, slight ECG QT_c prolongation	None identified
Amodiaquine	Good oral absorption; largely converted to active metabolite desethylamodiaquine	As for chloroquine	Nausea (tastes better than chloroquine)	Agranulocytosis; hepatitis, mainly with prophylactic use; should not be used with efavirenz
Primaquine	Complete oral absorption; active metabolite not known; $t_{1/2}$: 5–7 h	Radical cure; eradicates hepatic forms of <i>P. vivax</i> and <i>P. ovale</i> ; kills all stages of gametocyte development of <i>P. falciparum</i>	Nausea, vomiting, diarrhea, abdominal pain, hemolysis, methemoglobinemia	Massive hemolysis in subjects with severe G6PD deficiency
Mefloquine	Adequate oral absorption; no parenteral preparation; $t_{1/2}$: 14–20 days (shorter in malaria)	As for quinine	Nausea, giddiness, dysphoria, fuzzy thinking, sleeplessness, nightmares, sense of dissociation	Neuropsychiatric reactions, convulsions, encephalopathy
Halofantrine ^a	Highly variable absorption related to fat intake; $t_{1/2}$: 1–3 days (active desbutyl metabolite $t_{1/2}$: 3–7 days)	As for quinine	Diarrhea	Cardiac conduction disturbances; atrioventricular block; marked ECG QT_c interval prolongation; potentially lethal ventricular tachyarrhythmias
Lumefantrine	Highly variable absorption related to fat intake; $t_{1/2}$: 3–4 days	As for quinine	None identified	None identified
Artemisinin and derivatives (artemether, artesunate)	Good oral absorption, slow and variable absorption of IM artemether; artesunate and artemether biotransformed to active metabolite dihydroartemisinin; all drugs eliminated very rapidly; $t_{1/2}$: <1 h	Broader stage specificity and more rapid than other drugs; no action on liver stages; kills all but fully mature gametocytes of <i>P. falciparum</i>	Reduction in reticulocyte count (but not anemia); neutropenia at high doses; in some cases, delayed anemia after treatment of severe malaria with hyperparasitemia	Anaphylaxis, urticaria, fever
Pyrimethamine	Good oral absorption, variable IM absorption; $t_{1/2}$: 4 days	For blood stages, acts mainly on mature forms; causal prophylactic	Well tolerated	Megaloblastic anemia, pancytopenia, pulmonary infiltration
Proguanil (chloroguanide)	Good oral absorption; biotransformed to active metabolite cycloguanil; $t_{1/2}$: 16 h; biotransformation reduced by oral contraceptive use and in pregnancy	Causal prophylactic; not used alone for treatment	Well tolerated; mouth ulcers and rare alopecia	Megaloblastic anemia in renal failure
Atovaquone	Highly variable absorption related to fat intake; $t_{1/2}$: 30–70 h	Acts mainly on trophozoite blood stage	None identified	None identified
Tetracycline, doxycycline ^b	Excellent absorption; $t_{1/2}$: 8 h for tetracycline, 18 h for doxycycline	Weak antimalarial activity; should not be used alone for treatment	Gastrointestinal intolerance, deposition in growing bones and teeth, photosensitivity, moniliasis, benign intracranial hypertension	Renal failure in patients with impaired renal function (tetracycline)

^aHalofantrine should not be used by patients with long ECG QT_c intervals or known conduction disturbances or by those taking drugs that may affect ventricular repolarization—e.g., quinidine, quinine, mefloquine, chloroquine, neuroleptics, antiarrhythmics, tricyclic antidepressants, and some antihistamines. ^bTetracycline and doxycycline should not be given to pregnant women or to children <8 years of age.

Abbreviations: Cl , systemic clearance; ECG, electrocardiogram; G6PD, glucose-6-phosphate dehydrogenase; V_d , total apparent volume of distribution.

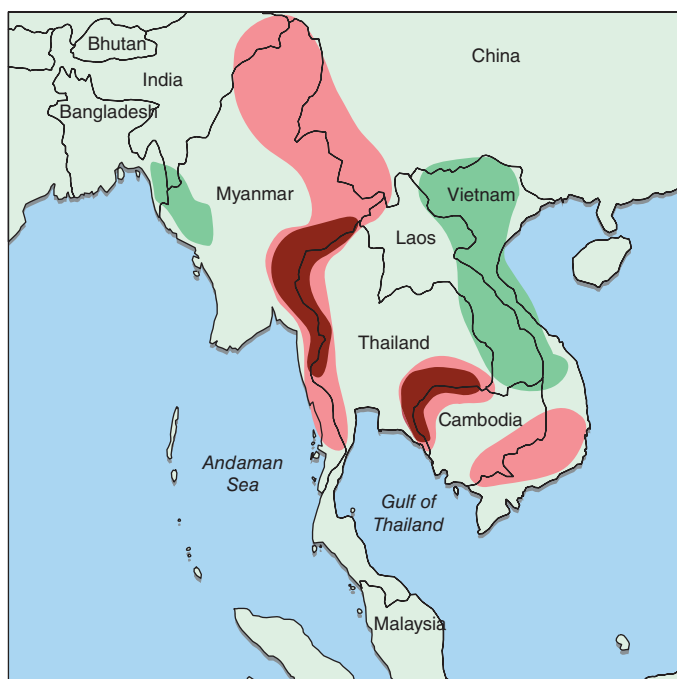


FIGURE 248-10 Mefloquine and artemisinin resistance in *Plasmodium falciparum* in Southeast Asia: high-level mefloquine resistance (dark red), low-level mefloquine resistance (pink), and mefloquine sensitivity (failure rate, <20%; green). There is insufficient information for other areas. Artemisinin resistance is now prevalent in areas where mefloquine resistance has been reported (pink areas).

used. Pyronaridine-artesunate is still under evaluation. Atovaquone-proguanil is highly effective everywhere, although it is seldom used in endemic areas because of its high cost and the propensity for rapid emergence of resistance. Of great concern is the emergence of artemisinin-resistant *P. falciparum* in western Cambodia and eastern Myanmar. Infections with these parasites are cleared slowly from the blood, with clearance times typically exceeding 3 days, and cure rates with ACTs are reduced.

The 3-day ACT regimens are all well tolerated, although mefloquine is associated with increased rates of vomiting and dizziness. As second-line treatments for recrudescence following first-line therapy, a different ACT regimen may be given; another alternative is a 7-day course of either artesunate or quinine plus tetracycline, doxycycline, or clindamycin. Tetracycline and doxycycline cannot be given to pregnant women or to children <8 years of age. Oral quinine is extremely bitter and regularly produces cinchonism comprising tinnitus, high-tone deafness, nausea, vomiting, and dysphoria. Adherence is poor with the required 7-day regimens of quinine.

Patients should be monitored for vomiting for 1 h after the administration of any oral antimalarial drug. If there is vomiting, the dose should be repeated. Symptom-based treatment, with tepid sponging and acetaminophen administration, lowers fever and thereby reduces the patient's propensity to vomit these drugs. Minor central nervous system reactions (nausea, dizziness, sleep disturbances) are common. The incidence of serious adverse neuropsychiatric reactions to mefloquine treatment is ~1 in 1000 in Asia but may be as high as 1 in 200 among Africans and Caucasians. All the antimalarial quinolines (chloroquine, mefloquine, and quinine) exacerbate the orthostatic hypotension associated with malaria, and all are tolerated better by children than by adults. Pregnant women, young children, patients unable to tolerate oral therapy, and nonimmune individuals (e.g., travelers) with suspected malaria should be evaluated carefully and hospitalization considered. If there is any doubt as to the identity of the infecting malarial species, treatment for *falciparum* malaria should be given. A negative blood smear makes malaria unlikely but does not rule it out

completely; thick blood films should be checked again 1 and 2 days later to exclude the diagnosis. Nonimmune patients receiving treatment for malaria should have daily parasite counts performed until the thick films are negative. If the level of parasitemia does not fall below 25% of the admission value in 48 h or if parasitemia has not cleared by 7 days (and adherence is assured), drug resistance is likely and the regimen should be changed.

To eradicate persistent liver stages and prevent relapse (radical treatment), primaquine (0.5 mg of base/kg or, in infections acquired in temperate areas, 0.25 mg/kg) should be given daily for 14 days to patients with *P. vivax* or *P. ovale* infections after laboratory tests for G6PD deficiency have proved negative. If the patient has a mild variant of G6PD deficiency, primaquine can be given in a dose of 0.75 mg of base/kg (45 mg maximum) once weekly for 8 weeks. Pregnant women with vivax or ovale malaria should not be given primaquine but should receive suppressive prophylaxis with chloroquine (5 mg of base/kg per week) until delivery, after which radical treatment can be given.

COMPLICATIONS

Acute Renal Failure If the plasma level of BUN or creatinine rises despite adequate rehydration, fluid administration should be restricted to prevent volume overload. As in other forms of hypercatabolic acute renal failure, renal replacement therapy is best performed early (Chap. 334). Hemofiltration and hemodialysis are more effective than peritoneal dialysis and are associated with lower mortality risk. Some patients with renal impairment pass small volumes of urine sufficient to allow control of fluid balance; these cases can be managed conservatively if other indications for dialysis do not arise. Renal function usually improves within days, but full recovery may take weeks.

Acute Pulmonary Edema (Acute Respiratory Distress Syndrome) Patients should be positioned with the head of the bed at a 45° elevation and given oxygen and IV diuretics. Pulmonary artery occlusion pressures may be normal, indicating increased pulmonary capillary permeability. Positive-pressure ventilation should be started early if the immediate measures fail (Chap. 326).

Hypoglycemia An initial slow injection of 50% dextrose (0.5 g/kg) should be followed by an infusion of 10% dextrose (0.10 g/kg per hour). The blood glucose level should be checked regularly thereafter as recurrent hypoglycemia is common, particularly among patients receiving quinine or quinidine. In severely ill patients, hypoglycemia commonly occurs together with metabolic (lactic) acidosis and carries a poor prognosis.

Other Complications Patients who develop spontaneous bleeding should be given fresh blood and IV vitamin K. Convulsions should be treated with IV or rectal benzodiazepines and, if necessary, respiratory support. Aspiration pneumonia should be suspected in any unconscious patient with convulsions, particularly with persistent hyperventilation; IV antimicrobial agents and oxygen should be administered, and pulmonary toilet should be undertaken. Hypoglycemia or gram-negative septicemia should be suspected when the condition of any patient suddenly deteriorates for no obvious reason during antimalarial treatment. In malaria-endemic areas where a high proportion of children are parasitemic, it is usually impossible to distinguish severe malaria from bacterial sepsis with confidence. These children should be treated with both antimalarials and broad-spectrum antibiotics from the outset. Because nontyphoidal *Salmonella* infections are particularly common, empirical antibiotics should be selected to cover these organisms. Antibiotics should be considered for severely ill patients of any age who are not responding to antimalarial treatment.

PREVENTION

In recent years, considerable progress has been made in malaria prevention, control, and research. Distribution of insecticide-treated

1382 bed-nets (ITNs) has been shown to reduce all-cause mortality in African children by 20%. New drugs have been discovered and developed, and one vaccine candidate (the RTS,S vaccine) will soon be considered for registration. Highly effective drugs, long-lasting ITNs, and insecticides for spraying dwellings are being purchased for endemic countries by the Global Fund to Fight AIDS, Tuberculosis, and Malaria; the President's Malaria Initiative (funded by the U.S. Agency for International Development and managed by the CDC in partnership with endemic countries); UNICEF; and other organizations. Malaria research and control are being strongly supported by the National Institute of Allergy and Infectious Diseases, the CDC, the Wellcome Trust, the Bill & Melinda Gates Foundation, the Multilateral Initiative on Malaria, the Roll Back Malaria Partnership, and the WHO among others. While a laudable goal, the global eradication of malaria is not feasible in the immediate future because of the widespread distribution of *Anopheles* breeding sites; the great number of infected persons; the continued use of ineffective antimalarial drugs; and inadequacies in human and material resources, infrastructure, and control programs. The call for and commitment to ultimate eradication of malaria by the Gates Foundation in 2007—seconded by Margaret Chan, Director General of the WHO—added great impetus to all malaria initiatives, especially those aimed at discovery and implementation of new interventions. Malaria may be contained by judicious use of insecticides to kill the mosquito vector, rapid diagnosis, patient management, and—where effective and feasible—administration of intermittent preventive treatment, seasonal malaria chemoprevention, or chemoprophylaxis to high-risk groups such as pregnant women, young children, and travelers from nonendemic regions. Malaria researchers are intensifying their efforts to gain a better understanding of parasite-human-mosquito interactions and to develop more effective control and prevention interventions. Despite the enormous investment in efforts to develop a malaria vaccine and the 30–60% efficacy in African children of a recombinant protein sporozoite-targeted adjuvanted vaccine (RTS,S) in field trials, no safe, highly effective, long-lasting vaccine is likely to be available for general use in the near future (Chap. 148). Indeed, protection from RTS,S in the very youngest recipients dropped to 16% only 4 years after vaccination. While there is great promise for one or several malaria vaccines on the more distant horizon, prevention and control measures continue to rely on antivector and drug-use strategies. Furthermore, recent gains are threatened by increasing insecticide resistance and behavioral changes (to avoid ITN contact) in anopheline mosquito vectors and by spreading artemisinin resistance in *P. falciparum*.

PERSONAL PROTECTION AGAINST MALARIA

Simple measures to reduce the frequency of infected-mosquito bites in malarious areas are very important. These measures include the avoidance of exposure to mosquitoes at their peak feeding times (usually dusk to dawn) and the use of insect repellents containing 10–35% DEET (or, if DEET is unacceptable, 7% picaridin), suitable clothing, and ITNs or other insecticide-impregnated materials. Widespread use of bed nets treated with residual pyrethroids reduces the incidence of malaria in areas where vectors bite indoors at night.

CHEMOPROPHYLAXIS

(Table 248-8; wwwnc.cdc.gov/travel/yellowbook/2014/chapter-3-infectious-diseases-related-to-travel/malaria) Recommendations for prophylaxis depend on knowledge of local patterns of *Plasmodium* species drug sensitivity and the likelihood of acquiring malarial infection. When there is uncertainty, drugs effective against resistant *P. falciparum* should be used (atovaquone-proguanil [Malarone], doxycycline, or mefloquine). Chemoprophylaxis is never entirely reliable, and malaria should always be considered in the differential diagnosis of fever in patients who have traveled to endemic areas, even if they are taking prophylactic antimalarial drugs.

Pregnant women traveling to malarious areas should be warned about the potential risks. All pregnant women at risk in endemic areas should be encouraged to attend regular antenatal clinics. Mefloquine

is the only drug advised for pregnant women traveling to areas with drug-resistant malaria; this drug is generally considered safe in the second and third trimesters of pregnancy, and the data on first-trimester exposure, although limited, are reassuring. Chloroquine and proguanil are regarded as safe. The safety of other prophylactic antimalarial agents in pregnancy has not been established. Antimalarial prophylaxis has been shown to reduce mortality rates among children between the ages of 3 months and 4 years in malaria-endemic areas; however, it is not a logistically or economically feasible option in many countries. The alternative—to give intermittent preventive treatment or seasonal malaria chemoprevention—shows promise for more widespread use in infants, young children, and pregnant women. Children born to nonimmune mothers in endemic areas (usually expatriates moving to malaria-endemic areas) should receive prophylaxis from birth.

Travelers should start taking antimalarial drugs 2 days to 2 weeks before departure so that any untoward reactions can be detected and so that therapeutic antimalarial blood concentrations will be present when needed (Table 248-8). Antimalarial prophylaxis should continue for 4 weeks after the traveler has left the endemic area, except if atovaquone-proguanil or primaquine has been taken; these drugs have significant activities against the liver stage of the infection (causal prophylaxis) and can be discontinued 1 week after departure from the endemic area. If suspected malaria develops while a traveler is abroad, obtaining a reliable diagnosis and antimalarial treatment locally is a top priority. Presumptive self-treatment for malaria with atovaquone-proguanil (for 3 consecutive days) or another drug can be considered under special circumstances; medical advice on self-treatment should be sought before departure for malarious areas and as soon as possible after illness begins. Every effort should be made to confirm the diagnosis by parasitologic studies.

Atovaquone-proguanil (Malarone; 3.75/1.5 mg/kg or 250/100 mg, daily adult dose) is a fixed-combination, once-daily prophylactic agent that is very well tolerated by adults and children, with fewer adverse gastrointestinal effects than chloroquine-proguanil and fewer adverse central nervous system effects than mefloquine. It is proguanil itself, rather than the antifolate metabolite cycloguanil, that acts synergistically with atovaquone. This combination is effective against all types of malaria, including multidrug-resistant falciparum malaria. Atovaquone-proguanil is best taken with food or a milky drink to optimize absorption. There are insufficient data on the safety of this regimen in pregnancy.

Mefloquine (250 mg of salt weekly, adult dose) has been widely used for malarial prophylaxis because it is usually effective against multidrug-resistant falciparum malaria and is reasonably well tolerated. The drug has been associated with rare episodes of psychosis and seizures at prophylactic doses; these reactions are more frequent at the higher doses used for treatment. More common side effects with prophylactic doses of mefloquine include mild nausea, dizziness, fuzzy thinking, disturbed sleep patterns, vivid dreams, and malaise. The drug is contraindicated for use by travelers with known hypersensitivity to mefloquine or related compounds (e.g., quinine, quinidine) and by persons with active or recent depression, anxiety disorder, psychosis, schizophrenia, another major psychiatric disorder, or seizures; mefloquine is not recommended for persons with cardiac conduction abnormalities although the evidence that it is cardiotoxic is very weak. Confidence is increasing with regard to the safety of mefloquine prophylaxis during pregnancy; in studies in Africa, mefloquine prophylaxis was found to be effective and safe during pregnancy. However, in one study from Thailand, treatment of malaria with mefloquine was associated with an increased risk of stillbirth; this effect was not seen subsequently.

Daily administration of doxycycline (100 mg daily, adult dose) is an effective alternative to atovaquone-proguanil or mefloquine. Doxycycline is generally well tolerated but may cause vulvovaginal thrush, diarrhea, and photosensitivity and cannot be used by children <8 years old or by pregnant women.

Chloroquine can no longer be relied upon to prevent *P. falciparum* infections in most areas but is used to prevent and treat malaria due to the other human *Plasmodium* species and for *P. falciparum* malaria in Central American countries west and north of the Panama

TABLE 248-8 DRUGS USED IN THE PROPHYLAXIS OF MALARIA

Drug	Usage	Adult Dose	Pediatric Dose	Comments
Atovaquone-proguanil (Malarone)	Prophylaxis in areas with chloroquine- or mefloquine-resistant <i>Plasmodium falciparum</i>	1 adult tablet PO ^a	5–8 kg: ½ pediatric tablet ^b daily ≥8–10 kg: ¾ pediatric tablet daily ≥10–20 kg: 1 pediatric tablet daily ≥20–30 kg: 2 pediatric tablets daily ≥30–40 kg: 3 pediatric tablets daily ≥40 kg: 1 adult tablet daily	Begin 1–2 days before travel to malarious areas. Take daily at the same time each day while in the malarious areas and for 7 days after leaving such areas. Atovaquone-proguanil is contraindicated in persons with severe renal impairment (creatinine clearance rate <30 mL/min). In the absence of data, it is not recommended for children weighing <5 kg, pregnant women, or women breast-feeding infants weighing <5 kg. Atovaquone-proguanil should be taken with food or a milky drink.
Chloroquine phosphate (Aralen and generic)	Prophylaxis only in areas with chloroquine-sensitive <i>P. falciparum</i> ^c or areas with <i>P. vivax</i> only	300 mg of base (500 mg of salt) PO once weekly	5 mg/kg of base (8.3 mg of salt/kg) PO once weekly, up to maximum adult dose of 300 mg of base	Begin 1–2 weeks before travel to malarious areas. Take weekly on the same day of the week while in the malarious areas and for 4 weeks after leaving such areas. Chloroquine phosphate may exacerbate psoriasis.
Doxycycline (many brand names and generic)	Prophylaxis in areas with chloroquine- or mefloquine-resistant <i>P. falciparum</i> ^c	100 mg PO qd (except in pregnant women; see Comments)	≥8 years of age: 2 mg/kg, up to adult dose	Begin 1–2 days before travel to malarious areas. Take daily at the same time each day while in the malarious areas and for 4 weeks after leaving such areas. Doxycycline is contraindicated in children <8 years of age and in pregnant women.
Hydroxychloroquine sulfate (Plaquenil)	An alternative to chloroquine for primary prophylaxis only in areas with chloroquine-sensitive <i>P. falciparum</i> ^c or areas with <i>P. vivax</i> only	310 mg of base (400 mg of salt) PO once weekly	5 mg of base/kg (6.5 mg of salt/kg) PO once weekly, up to maximum adult dose of 310 mg of base	Begin 1–2 weeks before travel to malarious areas. Take weekly on the same day of the week while in the malarious areas and for 4 weeks after leaving such areas. Hydroxychloroquine may exacerbate psoriasis.
Mefloquine (Lariam and generic)	Prophylaxis in areas with chloroquine-resistant <i>P. falciparum</i> ^c	228 mg of base (250 mg of salt) PO once weekly	≤9 kg: 4.6 mg of base/kg (5 mg of salt/kg) PO once weekly 10–19 kg: ¼ tablet once weekly 20–30 kg: ½ tablet once weekly 31–45 kg: ¾ tablet once weekly ≥46 kg: 1 tablet once weekly	Begin 1–2 weeks before travel to malarious areas. Take weekly on the same day of the week while in the malarious areas and for 4 weeks after leaving such areas. Mefloquine is contraindicated in persons allergic to this drug or related compounds (e.g., quinine and quinidine) and in persons with active or recent depression, generalized anxiety disorder, psychosis, schizophrenia, other major psychiatric disorders, or seizures. Use with caution in persons with psychiatric disturbances or a history of depression. Mefloquine is not recommended for persons with cardiac conduction abnormalities.
Primaquine	For prevention of malaria in areas with mainly <i>P. vivax</i>	30 mg of base (52.6 mg of salt) PO qd	0.5 mg of base/kg (0.8 mg of salt/kg) PO qd, up to adult dose; should be taken with food	Begin 1–2 days before travel to malarious areas. Take daily at the same time each day while in the malarious areas and for 7 days after leaving such areas. Primaquine is contraindicated in persons with G6PD deficiency. It is also contraindicated during pregnancy and in lactation unless the infant being breast-fed has a documented normal G6PD level.
Primaquine	Used for presumptive antirelapse therapy (terminal prophylaxis) to decrease risk of relapses of <i>P. vivax</i> and <i>P. ovale</i>	30 mg of base (52.6 mg of salt) PO qd for 14 days after departure from the malarious area	0.5 mg of base/kg (0.8 mg of salt/kg), up to adult dose, PO qd for 14 days after departure from the malarious area	This therapy is indicated for persons who have had prolonged exposure to <i>P. vivax</i> and/or <i>P. ovale</i> . It is contraindicated in persons with G6PD deficiency as well as during pregnancy and in lactation unless the infant being breast-fed has a documented normal G6PD level.

^aAn adult tablet contains 250 mg of atovaquone and 100 mg of proguanil hydrochloride. ^bA pediatric tablet contains 62.5 mg of atovaquone and 25 mg of proguanil hydrochloride.

^cVery few areas now have chloroquine-sensitive malaria (Fig. 248-2).

Source: CDC: www.cdc.gov/malaria/travelers/drugs.html.

Canal, Caribbean countries, and some countries in the Middle East. Chloroquine-resistant *P. vivax* has been reported from parts of eastern Asia, Oceania, and Central and South America. This drug is generally well tolerated, although some patients cannot take it because of malaise, headache, visual symptoms (due to reversible keratopathy), gastrointestinal intolerance, or pruritus. Chloroquine is considered safe in pregnancy. With chronic administration for >5 years, a characteristic dose-related retinopathy may develop, but this condition is rare at the doses used for antimalarial prophylaxis. Idiosyncratic or allergic reactions are also rare. Skeletal and/or cardiac myopathy is a

potential problem with protracted prophylactic use; such myopathy is more likely to occur at the high doses used in the treatment of rheumatoid arthritis. Neuropsychiatric reactions and skin rashes are unusual. When used continuously, amodiaquine, a related aminoquinoline, is associated with a high risk of agranulocytosis (~1 person in 2000) and hepatotoxicity (~1 person in 16,000); thus this agent should not be used for prophylaxis.

Primaquine (daily adult dose, 0.5 mg of base/kg or 30 mg taken with food), an 8-aminoquinoline compound, has proved safe and effective in the prevention of drug-resistant falciparum and vivax

1384 malaria in adults. This drug can be considered for persons who are traveling to areas with or without drug-resistant *P. falciparum* and who are intolerant to other recommended drugs. Abdominal pain and oxidant hemolysis—the principal adverse effects—are not common as long as the drug is taken with food and is not given to G6PD-deficient persons, in whom it can cause serious hemolysis. Travelers must be tested for G6PD deficiency and be shown to have a level in the normal range before receiving primaquine. Primaquine should not be given to pregnant women or neonates. Primaquine, given in a single dose of 0.25 mg/kg as a gametocytocide, together with an ACT is recommended in falciparum malaria treatment regimens in malaria elimination programs.

In the past, the dihydrofolate reductase inhibitors pyrimethamine and proguanil (chloroguanide) were administered widely, but the rapid selection of resistance in both *P. falciparum* and *P. vivax* has limited their use. Whereas antimalarial quinolines such as chloroquine (a 4-aminoquinoline) act on the erythrocyte stage of parasitic development, the dihydrofolate reductase inhibitors also inhibit preerythrocytic growth in the liver (causal prophylaxis) and development in the mosquito (sporontocidal activity). Proguanil is safe and well tolerated, although mouth ulceration occurs in ~8% of persons using this drug; it is considered safe for antimalarial prophylaxis in pregnancy. The prophylactic use of the combination of pyrimethamine and sulfadoxine is not recommended because of an unacceptable incidence of severe toxicity, principally exfoliative dermatitis and other skin rashes, agranulocytosis, hepatitis, and pulmonary eosinophilia (incidence, 1:7000; fatal reactions, 1:18,000). The combination of pyrimethamine with dapsone (0.2/1.5 mg/kg weekly; 12.5/100 mg, adult dose) has been used in some countries. Dapsone may cause methemoglobinemia and allergic reactions and (at higher doses) may pose a significant risk of agranulocytosis. Proguanil and the pyrimethamine-dapsone combination are not available in the United States.

Because of the increasing spread and intensity of antimalarial drug resistance (Figs. 248-2 and 248-10), the CDC recommends that travelers and their providers consider their destination, type of travel, and current medications and health risks when choosing antimalarial chemoprophylaxis. There is an increasingly appreciated problem of counterfeit and substandard antimalarial drugs (and other medicines) on the shelves of pharmacies in Southeast Asia and sub-Saharan Africa; hence, travelers should purchase their preventive drugs from a reputable source before going to a malarious country. Consultation for the evaluation of prophylaxis failures or treatment of malaria can be obtained from state and local health departments and the CDC Malaria Hotline (770-488-7788) or the CDC Emergency Operations Center (770-488-7100).

babesiosis and is endemic in the northeastern and upper midwestern United States. Seven states in these two regions (Connecticut, Massachusetts, Minnesota, New Jersey, New York, Rhode Island, and Wisconsin) account for >95% of the reported cases. Other etiologic agents include *Babesia duncani* and *B. duncani*-type organisms on the West Coast and *Babesia divergens*-like organisms in Kentucky, Missouri, and Washington State.

The primary causative agent of human babesiosis in Europe is *B. divergens*, but *Babesia venatorum* and *B. microti* also have been reported. In Asia, cases due to *B. microti*-like organisms have been documented in Japan, Taiwan, and the People's Republic of China. A case caused by *B. venatorum* also has been reported from the People's Republic of China. A case of *B. microti* infection was described in Australia. Sporadic cases due to uncharacterized species have been reported in Colombia, Egypt, India, Mozambique, and South Africa.

Incidence More than 1100 cases were reported in the United States in 2011, the year the disease became nationally notifiable. This figure represents a fivefold increase in incidence over the past decade. The incidence of babesiosis is markedly underestimated because symptoms are nonspecific and because young healthy individuals typically experience a mild or asymptomatic infection and may not seek medical attention. Fewer than 50 cases of *B. divergens*, *B. divergens*-like, and *B. venatorum* infections have been reported. Babesiosis caused by *B. duncani* and *B. duncani*-type organisms has also been sporadic, with fewer than 10 reported cases.



Modes of Transmission In the United States, *B. microti* is transmitted to humans primarily by the nymphal stage of the deer tick (*Ixodes scapularis*), the same tick that transmits the causative agents of Lyme disease (Chap. 210) and human granulocytotropic anaplasmosis (Chap. 211). Transmission generally occurs from May through October, with three-fourths of cases presenting in July and August. The vectors for transmission of *B. duncani* and *B. divergens*-like organisms are thought to be *Ixodes pacificus* and *Ixodes dentatus*, respectively. In Europe, *Ixodes ricinus* is the vector for *B. divergens* and *B. venatorum*. In Japan, *B. microti*-like organisms have been found in *Ixodes ovatus* ticks.

Babesiosis occasionally is acquired through transfusion of blood or blood products. *B. microti* is the most common transfusion-transmitted pathogen reported to the U.S. Food and Drug Administration, and more than 170 such cases have been identified. Three transfusion-transmitted cases caused by *B. duncani* have been documented. Transfusion-transmitted cases occur year-round, although most cases occur from June through November. More than 80% of cases occur in endemic areas. Transfusion-transmitted babesiosis occurs in nonendemic areas when unrecognized *Babesia*-contaminated blood products are imported from endemic areas: asymptotically infected residents of endemic areas donate blood in nonendemic areas, or residents of nonendemic areas travel to endemic areas, become infected, and donate blood after they return home. Approximately three-quarters of the transfusion-transmitted babesiosis cases reported between 1979 and 2009 occurred in the last decade of this period, and about one-fifth of patients died.

Seven cases of probable or confirmed congenital *B. microti* infection have been described. Other cases of neonatal babesiosis have been acquired by transfusion or tick bite.

CLINICAL MANIFESTATIONS

Asymptomatic *B. microti* Infection At least 20% of adults and 40% of children do not experience symptoms following *B. microti* infection. Asymptomatic infection, whether treated or not, may persist for >1 year after acute babesial illness. There is no evidence of long-term complications following asymptomatic infection; however, people who are asymptotically infected may transmit the infection when they donate blood.

Mild to Moderate *B. microti* Illness Symptoms typically develop following an incubation period of 1–4 weeks after tick bite and 1–9 weeks (but as long as 6 months) after transfusion of blood products. Patients experience a gradual onset of malaise, fatigue, and weakness. Fever

249 Babesiosis

Edouard G. Vannier, Peter J. Krause

Babesiosis is an emerging tick-borne infectious disease caused by protozoan parasites of the genus *Babesia* that invade and eventually lyse red blood cells (RBCs). Most cases are due to *Babesia microti* and occur in the United States, particularly in the Northeast and upper Midwest. The infection typically is mild in young and otherwise healthy individuals but can be severe and sometimes fatal in persons >50 years of age and in immunocompromised patients. Sporadic cases have been reported in Europe and the rest of the world.

ETIOLOGY AND EPIDEMIOLOGY



Geographic Distribution More than 100 *Babesia* species are found in wild and domestic animals; a few of these species cause infection in humans (Fig. 249-1). *B. microti*, a parasite of small rodents, is the most common etiologic agent of human

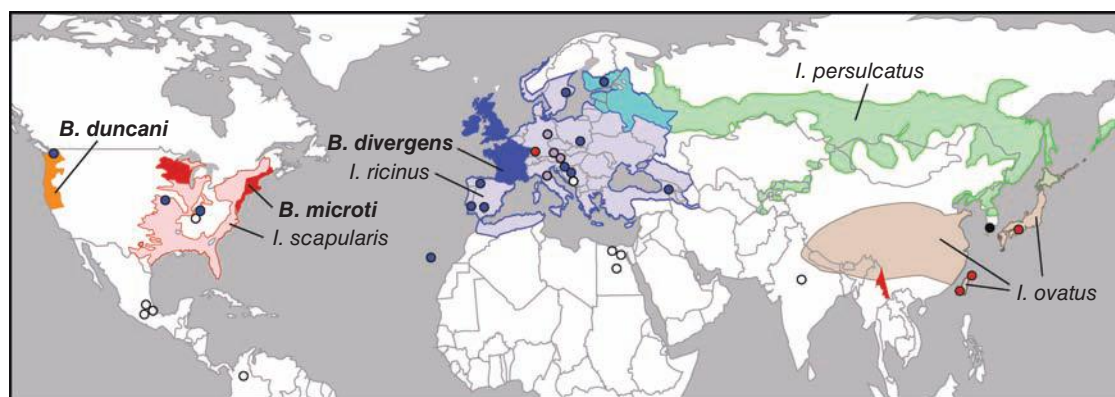


FIGURE 249-1 Worldwide distribution of human babesiosis. Dark colors designate areas where human babesiosis is either endemic or sporadic (as defined by more than three tick-borne cases reported in a country or state). Isolated cases of babesiosis are denoted by circles. Colors designate causative *Babesia* species: *Babesia microti* and *B. microti*-like organisms in red, *Babesia duncani* and *B. duncani*-type organisms in orange, *Babesia divergens* and *B. divergens*-like organisms in blue, *Babesia venatorum* in purple, KO1 in black, and unspicied *Babesia* organisms in white. Due to space constraints, the 10 cases reported from Montenegro are denoted by a single white circle, and those from Australia, Mozambique, and South Africa are not shown. Light colors denote areas that are enzootic for *Ixodes* tick species known to transmit one or several *Babesia* species but where human babesiosis has yet to be documented. (Adapted from E Vannier, PJ Krause: *N Engl J Med* 366:2397, 2012.)

can reach 40.9°C (105.6°F) and is accompanied by one or more of the following: chills, sweats, headache, myalgia, arthralgia, nausea, anorexia, and dry cough. Less common symptoms include sore throat, photophobia, abdominal pain, vomiting, weight loss, shortness of breath, and depression. On physical examination, fever is the salient feature. Ecchymoses and petechiae have been reported. An erythema migrans rash signifies concurrent Lyme disease (Chap. 210). Splenomegaly and hepatomegaly occasionally are noted. Lymphadenopathy is absent. Jaundice, slight pharyngeal erythema, and retinopathy with splinter hemorrhages and retinal infarcts rarely occur. Symptoms typically last 1–2 weeks, but fatigue may persist for several months.

Severe *B. microti* Illness Severe babesiosis requires hospital admission and typically occurs in patients with one or more of the following: age of >50 years, neonatal prematurity, male gender, asplenia, HIV/AIDS, malignancy, hemoglobinopathy, and immunosuppressive therapy. More than one-third of hospitalized patients develop complications, including acute respiratory distress syndrome, disseminated intravascular coagulation, congestive heart failure, renal failure, splenic infarcts, and splenic rupture. Patients who develop complications tend to have severe anemia (hemoglobin, ≤ 10 g/L). Laboratory prognostic factors for severe outcome—defined by hospitalization for >14 days, an intensive care unit stay of >2 days, or death—include an elevated alkaline phosphatase level (>125 U/L) and parasitemia of >4%. The fatality rate is 5–9% among hospitalized patients but is ~20% among immunocompromised patients and patients with transfusion-transmitted babesiosis.



Other *Babesia* Infections Cases of *B. duncani* infection range in severity from asymptomatic to fatal. Clinical manifestations are similar to those reported for *B. microti* infection. All three patients infected with *B. divergens*-like organisms in the United States required hospitalization; one died. Most cases of *B. divergens* infection in Europe have occurred in people lacking a spleen. The incubation period is 1–3 weeks. Symptoms appear suddenly and consist of fever (>41°C or 105.8°F), shaking chills, drenching sweats, headache, myalgia, and lumbar and abdominal pain. Hemoglobinuria and jaundice are commonly noted, and mild hepatomegaly may occur. If the infection is not treated, patients often develop pulmonary edema and renal failure. All four patients infected with *B. venatorum* in Europe had been splenectomized; their illness ranged from mild to severe, and none died. A child in China who developed *B. venatorum* illness had an intact spleen and survived the infection.

PATHOGENESIS

Anemia is a key feature of the pathogenesis of babesiosis. Hemolytic anemia caused by rupture of infected RBCs generates cell debris

that may accumulate in the kidney and cause renal failure. Anemia also results from the clearance of intact RBCs as they pass through the splenic red pulp and encounter resident macrophages. *Babesia* antigens expressed at the RBC membrane promote opsonization and facilitate uptake by splenic macrophages. In addition, RBCs are poorly deformable as a result of oxidation generated by the parasite and the host immune response and are filtered out as they attempt to squeeze across the venous vasculature. Bone marrow suppression due to cytokine production may also contribute to anemia.

An appropriate immune response is necessary for the control and clearance of *Babesia*. However, several lines of evidence suggest that an excessive response contributes to pathogenesis. Studies using laboratory mice have clearly established that CD4⁺ T cells are critical for resistance to and resolution of *B. microti* infection. CD4⁺ T cells are a major source of interferon γ (IFN- γ), and lack of this cytokine causes resistant mice to become highly susceptible to *B. microti*. IFN- γ is central to host resistance in *B. duncani* infection, but natural killer cells are its main source. *B. duncani* infection is more severe than *B. microti* infection in rodents and is characterized by pulmonary inflammation. Tumor necrosis factor α is expressed around alveolar septa, whereas IFN- γ is detected around pulmonary vessels. Blockade of either cytokine promotes the survival of mice infected with *B. duncani*.

DIAGNOSIS

A diagnosis of babesiosis should be considered for any patient who lives or travels in a *Babesia*-endemic area and presents with a febrile illness in the late spring, summer, or early autumn or within 6 months after a blood transfusion. Co-infection with *Babesia* should be considered in cases of Lyme disease or human granulocytotropic anaplasmosis when symptoms are more severe or prolonged than usual.

Screening laboratory tests can help support the diagnosis of babesiosis. A complete blood count often shows anemia and thrombocytopenia. Low hematocrit, hemoglobin, and haptoglobin levels and elevated reticulocyte counts and lactate dehydrogenase levels are consistent with hemolytic anemia. Liver enzyme tests often reveal elevated levels of alkaline phosphatase, aspartate and alanine aminotransferases, and bilirubin. Urinalysis may show hemoglobinuria, excess urobilinogen, and proteinuria. Elevated levels of blood urea nitrogen and serum creatinine indicate renal compromise.

A specific diagnosis usually is established by microscopic examination of Giemsa-stained thin blood smears (Fig. 249-2). *Babesia* trophozoites appear round, pear-shaped, or ameboid. The ring form is most common and lacks the central brownish deposit (hemozoin) typical of *Plasmodium falciparum* trophozoites (see Fig. 250e-1C). Other distinguishing features are the absence of schizonts and gametocytes and the occasional presence of tetrads (“Maltese cross”).

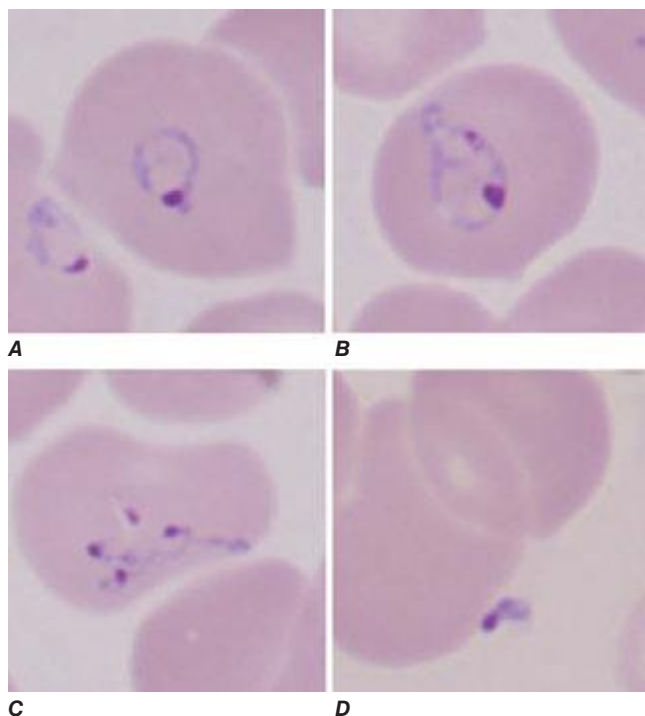


FIGURE 249-2 Giemsa-stained thin blood films showing *Babesia microti* parasites. *B. microti* are obligate parasites of erythrocytes. Trophozoites may appear as ring forms (A) or as ameboid forms (B). Merozoites can be arranged in tetrads and are pathognomonic (C). Extracellular parasites can be noted, particularly when parasitemia is high (D). (Adapted from E Vannier, PJ Krause: *N Engl J Med* 366:2397, 2012.)

Tetrads are characteristic of *B. microti*, *B. duncani*, and *B. divergens*-like organisms in human erythrocytes. Because the number of parasitized RBCs may be low, particularly at the onset of symptoms, identification of the parasite may require multiple blood smears over several days. Parasitemia levels can range from 1% to 20% in immunocompetent hosts and can be as high as 85% in immunocompromised patients. If parasites cannot be identified by microscopy and the disease is still suspected, amplification of the babesial 18S rRNA gene by polymerase chain reaction (PCR) is recommended. Quantitative PCR has greatly lowered the threshold for detection of *B. microti* DNA.

Serology can suggest or confirm the diagnosis of babesiosis. An indirect immunofluorescent antibody test for *B. microti* is most commonly used. IgM titers of $\geq 1:64$ and IgG titers of $\geq 1:1024$ suggest active or recent infection. Titers typically decline over 6–12 months. Antibodies to *B. microti* do not cross-react with *B. duncani* or *B. divergens* antigen. In *B. divergens* infection, serology is of limited use because symptoms often appear before antibodies can be detected. Sera from patients infected with *B. divergens*-like organisms or *B. venatorum* are reactive against *B. divergens* antigen.

TREATMENT BABESIOSIS

ASYMPTOMATIC *B. MICROTI* INFECTION

People who experience asymptomatic *B. microti* infection often are not diagnosed and treated. Current guidelines recommend antibiotic therapy for asymptomatic carriers only if parasitemia persists for >3 months. Laboratory-based tests are being developed for the purpose of screening the blood supply and will result in the identification of a greater number of asymptomatic *B. microti* carriers, raising the question of whether they should be treated.

MILD TO MODERATE *B. MICROTI* ILLNESS

Atovaquone plus azithromycin, given orally for 7–10 days, is the recommended antibiotic treatment combination for mild to moderate

TABLE 249-1 TREATMENT OF HUMAN BABESIOSIS

Adults	Children
<i>B. microti</i> Infection (Mild to Moderate Illness^{a,b})	
Atovaquone (750 mg q12h PO)	Atovaquone (20 mg/kg q12h PO; maximum, 750 mg/dose)
plus	plus
Azithromycin (500 mg/d PO on day 1, 250 mg/d PO thereafter)	Azithromycin (10 mg/kg qd PO on day 1 [maximum, 500 mg/dose], 5 mg/kg qd PO thereafter [maximum, 250 mg/dose])
<i>B. microti</i> Infection (Severe Illness^{c,d})	
Clindamycin (300–600 mg q6h IV or 600 mg q8h PO)	Clindamycin (7–10 mg/kg q6–8h IV or 7–10 mg/kg q6–8h PO; maximum, 600 mg/dose)
plus	plus
Quinine (650 mg q6–8h PO)	Quinine (8 mg/kg q8h PO; maximum, 650 mg/dose)
plus	plus
Consider exchange transfusion	Consider exchange transfusion
<i>B. divergens</i> Infection	
Immediate complete exchange transfusion	Immediate complete exchange transfusion
plus	plus
Clindamycin (600 mg q6–8h IV)	Clindamycin (7–10 mg/kg q6–8h IV; maximum, 600 mg/dose)
plus	plus
Quinine (650 mg q8h PO)	Quinine (8 mg/kg q8h PO; maximum, 650 mg/dose)

^aTreatment duration, 7–10 days. ^bA high dose of azithromycin (600–1000 mg) combined with atovaquone has been recommended for immunocompromised hosts. ^cTreatment typically is given for 7–10 days, but its duration may vary. In severely immunocompromised patients, therapy should be continued for at least 6 weeks, including 2 weeks after parasites are no longer detected on blood smear. ^dSeveral alternative regimens have been used in a limited number of cases of *B. microti* infection, and their efficacy is uncertain. These regimens include atovaquone, azithromycin, and clindamycin; atovaquone, azithromycin, and doxycycline; atovaquone, clindamycin, and doxycycline; atovaquone, doxycycline, and artemisinin; atovaquone-proguanil; azithromycin and quinine; and azithromycin, clindamycin, and doxycycline.

Sources: (1) ME Falagas, MS Klemperer: *Clin Infect Dis* 22:809, 1996. (2) PJ Krause et al: *N Engl J Med* 343:1454, 2000. (3) PJ Krause et al: *Clin Infect Dis* 46:370, 2008. (4) CM Shih, CC Wang: *Am J Trop Med Hyg* 59:509, 1998. (5) CP Stowell et al: *N Engl J Med* 356:2313, 2007. (6) JM Vyas et al: *Clin Infect Dis* 45:1588, 2007. (7) GP Wormser et al: *Clin Infect Dis* 50:381, 2010.

babesiosis (Table 249-1). Clindamycin plus quinine is a second choice. Symptoms usually begin to resolve within 48 h of therapy initiation, but complete resolution may take weeks to months. An atypical or poor response to therapy should raise concern about the possibility of concurrent Lyme disease (Chap. 210) or human granulocytotropic anaplasmosis (Chap. 211). In the first prospective trial of antibabesial therapy, the combination of atovaquone plus azithromycin was compared with clindamycin plus quinine in adults. These two drug combinations were equally effective in resolving symptoms and clearing parasitemia. Adverse effects were reported in 15% of trial participants who received atovaquone plus azithromycin but in 72% of those who received clindamycin plus quinine. Adverse reactions were so severe that treatment had to be stopped in about one-third of participants taking clindamycin plus quinine but in only 2% of those taking atovaquone plus azithromycin.

SEVERE *B. MICROTI* ILLNESS

Clindamycin given intravenously plus quinine given orally for 7–10 days constitute the recommended treatment for severe babesiosis. Intravenous quinidine may be used instead of oral quinine but requires cardiac monitoring because of the risk of QT prolongation and polymorphic ventricular tachycardia.

Standard antimicrobial therapy is sometimes insufficient to resolve symptoms and clear parasitemia, especially in patients with

marked immunosuppression due to splenectomy, HIV/AIDS, malignancy, and/or immunosuppressive therapy (including rituximab for B cell lymphomas). In such patients, antimicrobial therapy should be administered for at least 6 weeks, including 2 weeks after parasites are no longer observed on blood smear. High-dose azithromycin (600–1000 mg/d) plus atovaquone have been successfully used in immunocompromised patients. Resistance to atovaquone plus azithromycin has occurred in a few cases. In patients who are unresponsive to atovaquone plus azithromycin or who do not tolerate clindamycin plus quinine, alternative regimens have been used (Table 249-1).

Partial or complete RBC exchange transfusion is recommended in patients with high-grade parasitemia (>10%), severe anemia (hemoglobin, <10 g/dL), or pulmonary, hepatic, or renal compromise. Parasitemia and hematocrit should be monitored daily until symptoms recede and the parasitemia level is <5%.

OTHER *BABESIA* INFECTIONS

The regimen for *B. duncani* infections typically consists of intravenous clindamycin (600 mg tid/qid or 1200 mg bid) plus oral quinine (600–650 mg tid) for 7–10 days. A regimen often used for *B. divergens*-like infections is intravenous clindamycin (600 mg tid/qid, 900 mg tid, or 1200 mg bid) plus oral quinine or quinidine (650 mg tid).



In Europe, *B. divergens* infection is considered a medical emergency. The recommended treatment is immediate complete blood exchange transfusion and therapy with intravenous clindamycin plus either oral quinine or intravenous quinidine. Some cases have been cured with exchange transfusion and clindamycin monotherapy. Anemia may persist for >1 month and require additional transfusion.

PREVENTION

No vaccine is available for human use. There is no role for antibiotic prophylaxis. Individuals who reside in endemic areas, especially those at risk for severe babesiosis, should wear clothing that covers the lower part of the body, apply tick repellents (such as DEET) to clothing, and limit outdoor activities where ticks may abound from May through October. The skin should be thoroughly examined after outdoor activities, and ticks should be removed with tweezers. Individuals with a history of symptomatic babesiosis or with positive *Babesia* serology are indefinitely deferred from donating blood.

250e Atlas of Blood Smears of Malaria and Babesiosis

Nicholas J. White, Joel G. Breman

Six species of blood protozoan parasites cause human malaria (Chap. 248): the potentially lethal and often drug-resistant *Plasmodium falciparum*; the relapsing parasites *Plasmodium vivax* and *Plasmodium ovale* (with what appear to be two morphologically identical sympatric species of *P. ovale*); *Plasmodium malariae*, which can persist at low densities for years; and, in infections in individuals living in or close to tropical forests in Southeast Asia, *Plasmodium knowlesi*, a monkey parasite that microscopically resembles *P. falciparum* (young forms) and *P. malariae* (older forms) but is identified definitively by molecular methods.

The malaria parasites are readily seen under the microscope ($\times 1000$ magnification) in thick and thin blood smears stained with supravital dyes (e.g., Giemsa's, Field's, Wright's, Leishman's). The morphologic characteristics of the parasites are summarized in Table 250e-1. In the thick film, lysis of red blood cells by water leaves the stained white cells and parasites, allowing detection of densities as low as 50 parasites/ μL . This degree of sensitivity is up to 100 times greater than that of the thin film, in which the cells are fixed and the malaria parasites are seen inside the red cells. The thin film is better for speciation and provides useful prognostic information in severe falciparum malaria. Several findings are associated with increased mortality risk: high parasite counts, more mature parasites ($>20\%$ containing visible malaria pigment), and phagocytosed malaria pigment in $>5\%$ of neutrophils.

Babesia microti (Chap. 249) appears as a small ring form resembling *P. falciparum*. Unlike *Plasmodium*, *Babesia* does not cause the production of pigment in parasites, nor are schizonts or gametocytes formed.

TABLE 250e-1 MORPHOLOGIC CHARACTERISTICS OF HUMAN MALARIA PARASITES^a

	<i>Plasmodium falciparum</i>	<i>Plasmodium vivax</i>	<i>Plasmodium ovale</i> ^b	<i>Plasmodium malariae</i>
Asexual parasites	Usually only fine blue ring forms (some resembling stereo headsets) are seen. Parasitemia level may exceed 2%.	Irregular, large, fairly thick rings become highly pleomorphic as the parasite grows. Parasitemia level is low.	Regular, dense ring enlarges to compact, blue, mature trophozoite (rectangular or band-form). Parasitemia level is low.	Dense, thick rings mature to dense, round trophozoites. Parasitemia level is low.
Schizonts	Rare in peripheral blood; 8–32 merozoites, dark brown-black pigment	Common; 12–18 merozoites, orange-brown pigment	8–14 merozoites, brown or black pigment	8–10 merozoites, dark brown or black pigment
Gametocytes	Banana-shaped; male: light blue; female: darker blue; a few scattered blue-black pigment granules in cytoplasm	Round or oval; male: round, pale blue; female: oval, dark blue; triangular nucleus, a few orange-brown pigment granules	Large, round, dense, and blue (like <i>P. malariae</i>), but prominent James's dots; brown pigment	Large, oval; male: pale blue; female: dense blue; large black pigment granules
RBC changes	RBCs are normal in size. As the parasite matures, the RBC cytoplasm becomes pale, the cells become crenated, and a few small red dots may appear over the cytoplasm (Maurer's clefts).	RBCs are enlarged. Pale red Schüffner's dots increase in number as the parasite matures.	RBCs become oval with tufted ends. Red James's dots are prominent.	RBCs are normal in size and shape. No red dots are seen.

^aThe early trophozoites of *Plasmodium knowlesi* resemble those of *P. falciparum*. The late and mature trophozoites and schizonts of *P. knowlesi* appear very similar to those of *P. malariae*. The differences are that (1) *P. knowlesi* trophozoites may have double chromatin dots, with two or three parasites per RBC, and may cause higher-level parasitemia; and (2) *P. knowlesi* mature schizonts have 16 merozoites rather than the 8–10 found with *P. malariae*. ^bTwo morphologically identical sympatric species, according to recent evidence.

Abbreviation: RBC, red blood cell.

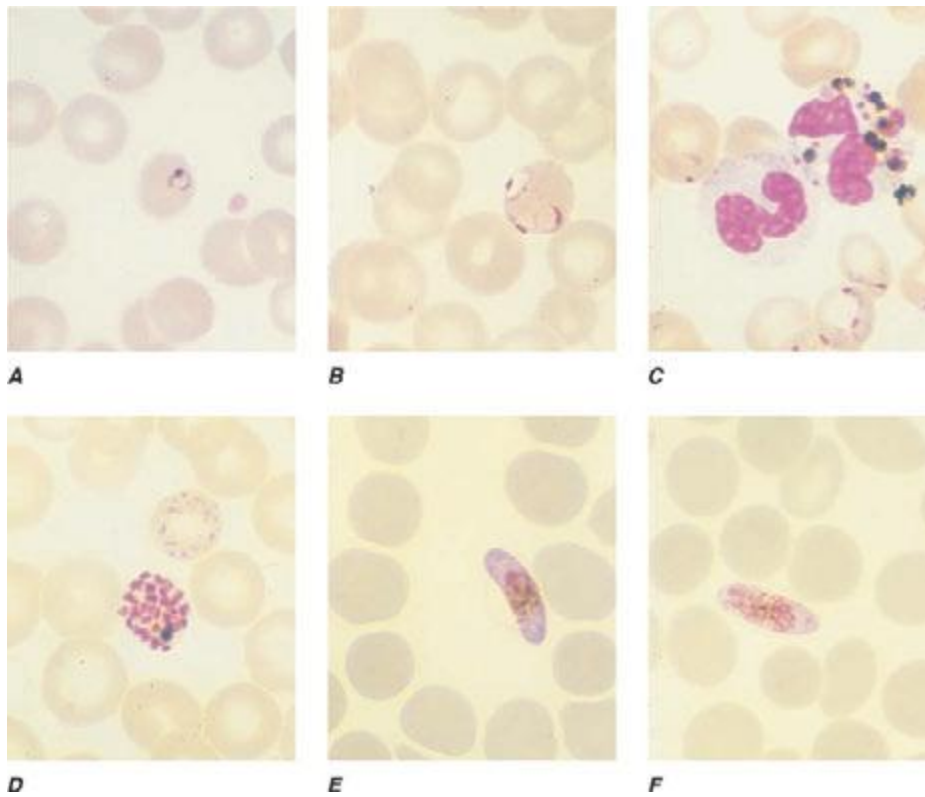


FIGURE 250e-1 Thin blood films of *Plasmodium falciparum*. **A.** Young trophozoites. **B.** Old trophozoites. **C.** Pigment in polymorphonuclear cells and trophozoites. **D.** Mature schizonts. **E.** Female gametocytes. **F.** Male gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

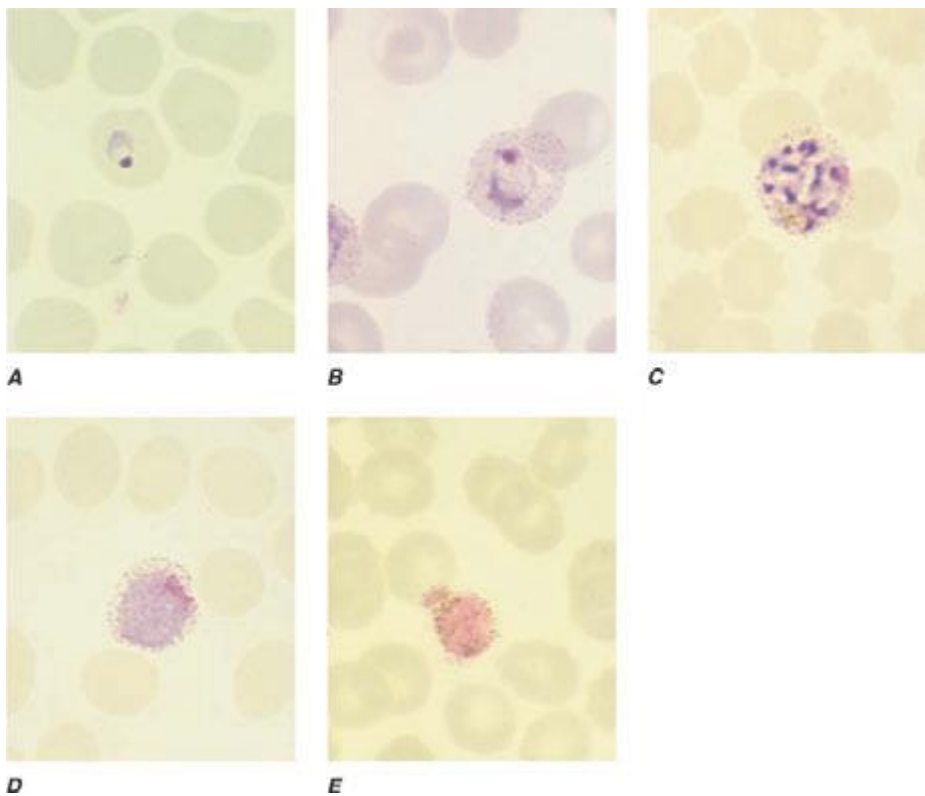


FIGURE 250e-2 Thin blood films of *Plasmodium vivax*. **A.** Young trophozoites. **B.** Old trophozoites. **C.** Mature schizonts. **D.** Female gametocytes. **E.** Male gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

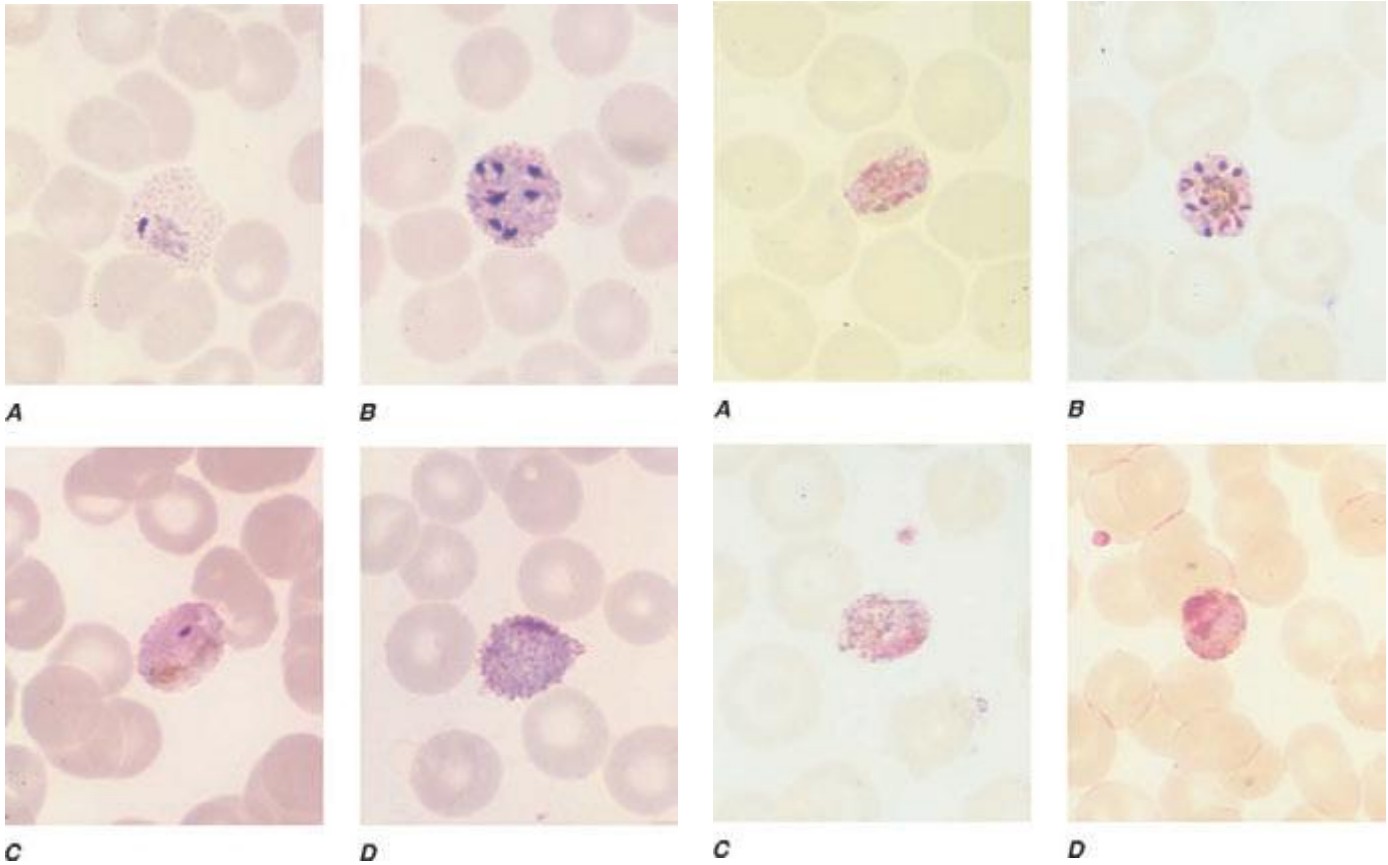


FIGURE 250e-3 Thin blood films of *Plasmodium ovale*. **A.** Old trophozoites. **B.** Mature schizonts. **C.** Male gametocytes. **D.** Female gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

FIGURE 250e-4 Thin blood films of *Plasmodium malariae*. **A.** Old trophozoites. **B.** Mature schizonts. **C.** Male gametocytes. **D.** Female gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

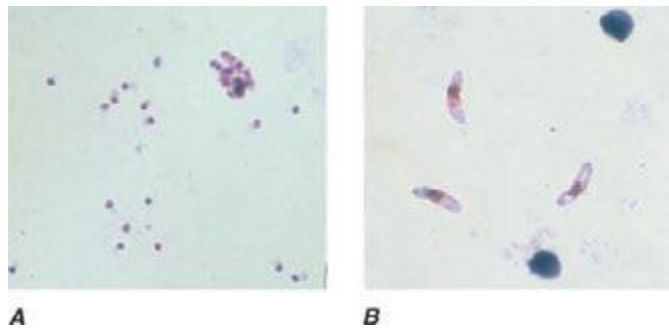


FIGURE 250e-5 Thick blood films of *Plasmodium falciparum*. **A.** Trophozoites. **B.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

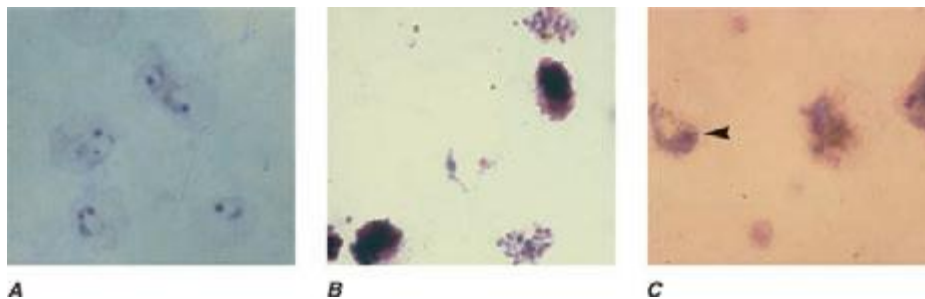


FIGURE 250e-6 Thick blood films of *Plasmodium vivax*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

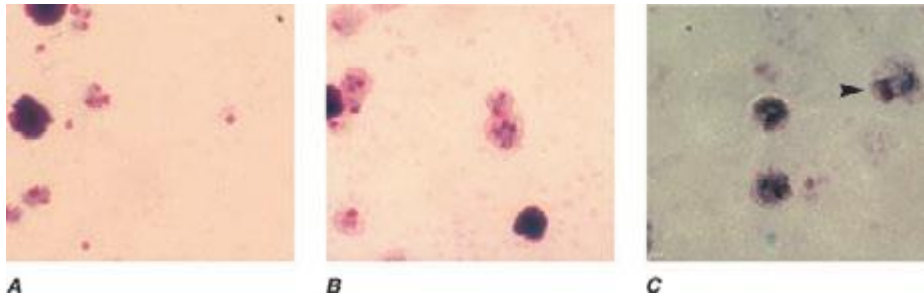


FIGURE 250e-7 Thick blood films of *Plasmodium ovale*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

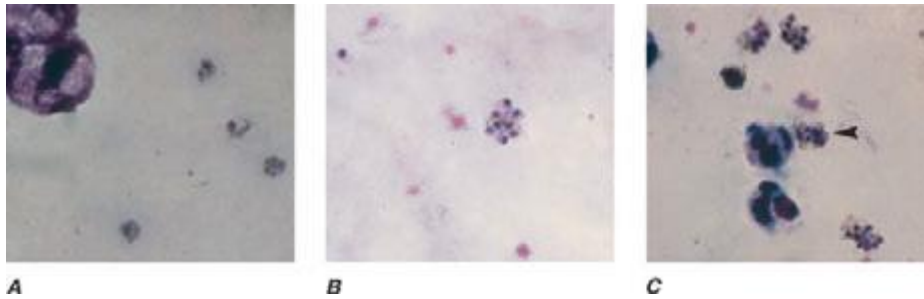


FIGURE 250e-8 Thick blood films of *Plasmodium malariae*. **A.** Trophozoites. **B.** Schizonts. **C.** Gametocytes. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

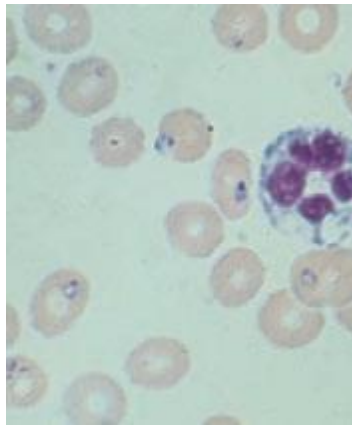


FIGURE 250e-9 Thin blood film showing trophozoites of *Babesia*. (Reproduced from *Bench Aids for the Diagnosis of Malaria Infections*, 2nd ed, with the permission of the World Health Organization.)

251 Leishmaniasis

Shyam Sundar

DEFINITION

Encompassing a complex group of disorders, leishmaniasis is caused by unicellular eukaryotic obligatory intracellular protozoa of the genus *Leishmania* and primarily affects the host's reticuloendothelial system. *Leishmania* species produce widely varying clinical syndromes ranging from self-healing cutaneous ulcers to fatal visceral disease. These syndromes fall into three broad categories: visceral leishmaniasis (VL), cutaneous leishmaniasis (CL), and mucosal leishmaniasis (ML).

ETIOLOGY AND LIFE CYCLE

Leishmaniasis is caused by ~20 species of the genus *Leishmania* in the order Kinetoplastida and the family Trypanosomatidae (Table 251-1). Several clinically important species are of the subspecies *Viannia*. The organisms are transmitted by phlebotomine sandflies of the genus *Phlebotomus* in the “Old World” (Asia, Africa, and Europe) and the genus *Lutzomyia* in the “New World” (the Americas). Transmission may be anthroponotic (i.e., the vector transmits the infection from infected humans to healthy humans) or zoonotic (i.e., the vector transmits the infection from an animal reservoir to humans). Human-to-human transmission via shared infected needles has been documented in IV drug users in the Mediterranean region. In utero transmission to the fetus occurs rarely.

Leishmania organisms occur in two forms: extracellular, flagellate promastigotes (length, 10–20 μm) in the sandfly vector and intracellular, nonflagellate amastigotes (length, 2–4 μm ; Fig. 251-1) in vertebrate hosts, including humans. Promastigotes are introduced through the proboscis of the female sandfly into the skin of the vertebrate host. Neutrophils predominate among the host cells that first encounter and take up promastigotes at the site of parasite delivery. The infected neutrophils may undergo apoptosis and release viable parasites that are taken up by macrophages, or the apoptotic cells may themselves be taken up by macrophages and dendritic cells. The parasites multiply as amastigotes inside macrophages, causing cell rupture with subsequent invasion of other macrophages. While feeding on infected hosts, sandflies pick up amastigotes, which transform into the flagellate form in the flies' posterior midgut and multiply by binary fission; the promastigotes then migrate to the anterior midgut and can infect a new host when flies take another blood meal.

EPIDEMIOLOGY



Leishmaniasis occurs in 98 countries—most of them developing—in tropical and temperate regions (Fig. 251-2). More than 1.5 million cases occur annually, of which 0.7–1.2 million are CL (and its variations) and 200,000–400,000 are VL. More than 350 million people are at risk, with an overall prevalence of 12 million. Although the distribution of *Leishmania* is limited by the distribution of sandfly vectors, human leishmaniasis is on the increase worldwide.

VISCERAL LEISHMANIASIS

VL (also known as *kala-azar*, a Hindi term meaning “black fever”) is caused by the *Leishmania donovani* complex, which includes *L. donovani* and *Leishmania infantum* (the latter designated *Leishmania chagasi* in the New World); these species are responsible for anthroponotic and zoonotic transmission, respectively. India and neighboring Bangladesh, Sudan, South Sudan, Ethiopia, and Brazil are the four largest foci of VL and account for 90% of the world's VL burden, with India being the worst affected. Zoonotic VL is reported from all countries in the Middle East, Pakistan, and other countries from western Asia to China. Endemic foci also exist in the independent states of the former Soviet Union, mainly Georgia and Azerbaijan. In the Horn of Africa, Sudan, South Sudan, Ethiopia, Kenya, Uganda, and Somalia report VL. In Sudan and South Sudan, large outbreaks are thought to be anthroponotic, although zoonotic transmission also occurs. VL is rare in West and sub-Saharan Africa.

TABLE 251-1 GEOGRAPHIC DISTRIBUTION AND CHARACTERISTIC EPIDEMIOLOGY OF LEISHMANIASES

Organism, Endemic Region	Clinical Syndrome	Species	Vector	Reservoir	Transmission	Setting
<i>Leishmania donovani</i> Complex						
South Asia	VL, PKDL	<i>L. donovani</i>	<i>Phlebotomus argentipes</i>	Humans	Anthroponotic	Rural, domestic
Sudan, South Sudan, Somalia, Ethiopia, Kenya, Uganda	VL, PKDL	<i>L. donovani</i>	<i>P. orientalis</i> , <i>P. martini</i>	Humans, rodents in Sudan, canines	Anthroponotic, occasionally zoonotic	Majority peridomestic, occasionally sylvatic
Mediterranean basin, Middle East, central Asia, China	VL, CL	<i>L. infantum</i>	<i>P. perniciosus</i> , <i>P. ariasi</i>	Dogs, foxes, jackals	Zoonotic	Domestic, peridomestic
Middle East, Saudi Arabia, Yemen	VL	<i>L. donovani</i>	<i>P. perniciosus</i> , <i>P. ariasi</i>	Dogs, foxes, jackals	Zoonotic	Domestic, peridomestic
Central and South America	VL, CL	<i>L. infantum</i> ^a	<i>Lutzomyia longipalpis</i>	Foxes, dogs, opossums	Zoonotic	Domestic, peridomestic, periurban
Azerbaijan, Armenia, Georgia, Kazakhstan, Kyrgyzstan, Tajikistan, Turkmenistan, Uzbekistan	VL	<i>L. infantum</i>	<i>P. turanicus</i>	Humans, dogs, foxes	Anthroponotic, zoonotic	Domestic
<i>L. tropica</i>						
Western India to Turkey, parts of North and East Africa	CL, leishmaniasis recidivans	<i>L. tropica</i>	<i>P. sergenti</i>	Humans	Anthroponotic	Urban domestic, peridomestic
<i>L. major</i>						
Western and Central Asia, North and sub-Saharan Africa	CL	<i>L. major</i>	<i>P. papatasi</i> , <i>P. duboscqi</i>	Nile rats, rodents	Zoonotic	Sylvatic, peridomestic
Kazakhstan, Turkmenistan, Uzbekistan	CL	<i>L. major</i>	<i>P. papatasi</i> , <i>P. duboscqi</i>	Gerbils	Zoonotic	Rural
<i>L. aethiopia</i>						
Ethiopia, Uganda, Kenya	CL, DCL	<i>L. aethiopia</i>	<i>P. longipes</i> , <i>P. pedifer</i>	Hyraxes	Zoonotic	Sylvatic, peridomestic
Subspecies <i>Viannia</i>						
Peru, Ecuador	CL, ML	<i>L. (V.) peruviana</i>	<i>Lutzomyia verrucarum</i> , <i>L. peruensis</i>	Wild rodents	Zoonotic	Andean Valleys
Guyana, Surinam, French Guyana, Ecuador, Brazil, Colombia, Bolivia	CL, ML	<i>L. (V.) guyanensis</i>	<i>L. umbratilis</i>	Sloths, arboreal anteaters, opossums	Zoonotic	Tropical forest
Central America, Ecuador, Colombia	CL, ML	<i>L. (V.) panamensis</i>	<i>L. trapidoi</i>	Sloths	Zoonotic	Tropical forest and deforested areas
South and Central America	CL, ML	<i>L. (V.) braziliensis</i>	<i>Lutzomyia</i> spp., <i>L. umbratilis</i> , <i>Psychodopygus wellcomei</i>	Forest rodents, peridomestic animals	Zoonotic	Tropical forest and deforested areas
<i>L. mexicana</i> Complex						
Central America and northern parts of South America	CL, ML, DCL	<i>L. amazonensis</i>	<i>L. flaviscutellata</i>	Forest rodents	Zoonotic	Tropical forest and deforested areas
	CL, ML, DCL	<i>L. mexicana</i>	<i>L. olmeca</i>	Variety of forest rodents and marsupials	Zoonotic	Tropical forest and deforested areas
	CL, DCL	<i>L. pifanoi</i>	<i>L. olmeca</i>	Variety of forest rodents and marsupials	Zoonotic	Tropical forest and deforested areas

^a*L. infantum* is designated *L. chagasi* in the New World.

Abbreviations: CL, cutaneous leishmaniasis; DCL, diffuse cutaneous leishmaniasis; ML, mucosal leishmaniasis; PKDL, post-kala-azar dermal leishmaniasis; VL, visceral leishmaniasis.

Mediterranean VL, long an established endemic disease due to *L. infantum*, has a large canine reservoir and was seen primarily in infants before the advent of HIV infection. In Mediterranean Europe, 70% of adult VL cases are associated with HIV co-infection. The combination is deadly because of the impact of the two infections together

on the immune system. IV drug users are at particular risk. Other forms of immunosuppression (e.g., that associated with organ transplantation) also predispose to VL. In the Americas, disease caused by *L. infantum* is endemic from Mexico to Argentina, but 90% of cases in the New World are reported from northeastern Brazil.

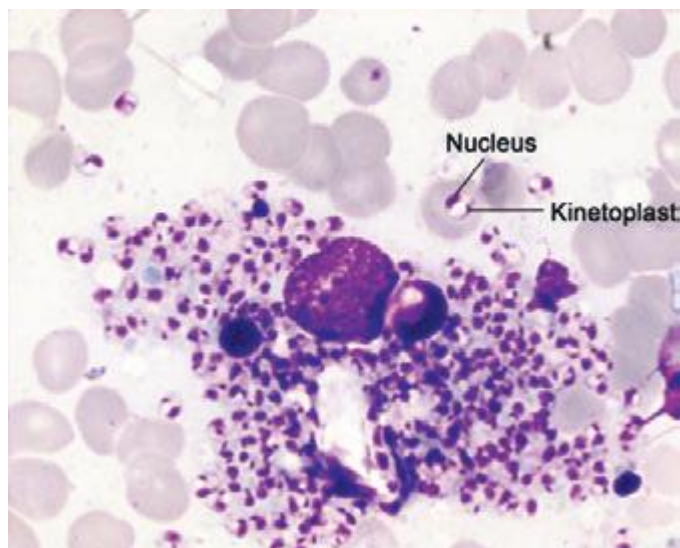


FIGURE 251-1 A macrophage with numerous intracellular amastigotes (2–4 μm) in a Giemsa-stained splenic smear from a patient with visceral leishmaniasis. Each amastigote contains a nucleus and a characteristic kinetoplast consisting of multiple copies of mitochondrial DNA. A few extracellular parasites are also visible.

Immunopathogenesis The majority of individuals infected by *L. donovani* or *L. infantum* mount a successful immune response and control the infection, never developing symptomatic disease. Forty-eight hours after intradermal injection of killed promastigotes, these individuals exhibit delayed-type hypersensitivity (DTH) to leishmanial antigens in the leishmanin skin test (also called the Montenegro skin test). Results in mouse models indicate that the development of acquired resistance to leishmanial infection is controlled by the production of interleukin (IL) 12 by antigen-presenting cells and the subsequent secretion of interferon (IFN) γ , tumor necrosis factor (TNF) α , and other proinflammatory cytokines by the T helper 1 (T_H1) subset of T lymphocytes. The immune response in patients developing active VL is complex; in addition to increased production of multiple proinflammatory cytokines and chemokines, patients with active disease have markedly elevated levels of IL-10 in serum as well as enhanced IL-10 mRNA expression in lesional tissues. The main disease-promoting activity of IL-10 in VL may be to condition host

macrophages for enhanced survival and growth of the parasite. IL-10 can render macrophages unresponsive to activation signals and inhibit killing of amastigotes by downregulating the production of TNF- α and nitric oxide. Multiple antigen-presentation functions of dendritic cells and macrophages are also suppressed by IL-10. Patients with such suppression do not have positive leishmanin skin tests, nor do their peripheral-blood mononuclear cells respond to leishmanial antigens in vitro. Organs of the reticuloendothelial system are predominantly affected, with remarkable enlargement of the spleen, liver, and lymph nodes in some regions. The tonsils and intestinal submucosa are also heavily infiltrated with parasites. Bone marrow dysfunction results in pancytopenia.

Clinical Features On the Indian subcontinent and in the Horn of Africa, persons of all ages are affected by VL. In endemic areas of the Americas and the Mediterranean basin, immunocompetent infants and small children as well as immunodeficient adults are affected especially often. The most common presentation of VL is an abrupt onset of moderate- to high-grade fever associated with rigor and chills. Fever may continue for several weeks with decreasing intensity, and the patient may become afebrile for a short period before experiencing another bout of fever. The spleen may be palpable by the second week of illness and, depending on the duration of illness, may become hugely enlarged (Fig. 251-3). Hepatomegaly (usually moderate in degree) soon follows. Lymphadenopathy is common in most endemic regions of the world except the Indian subcontinent, where it is rare. Patients lose weight and feel weak, and the skin gradually develops dark discoloration due to hyperpigmentation that is most easily seen in brown-skinned individuals. In advanced illness, hypoalbuminemia may manifest as pedal edema and ascites. Anemia appears early and may become severe enough to cause congestive heart failure. Epistaxis, retinal hemorrhages, and gastrointestinal bleeding are associated with thrombocytopenia. Secondary infections such as measles, pneumonia, tuberculosis, bacillary or amebic dysentery, and gastroenteritis are common. Herpes zoster, chickenpox, boils in the skin, and scabies may also occur. Untreated, the disease is fatal in most patients, including 100% of those with HIV co-infection.

Leukopenia and anemia occur early and are followed by thrombocytopenia. There is a marked polyclonal increase in serum immunoglobulins. Serum levels of hepatic aminotransferases are raised in a significant proportion of patients, and serum bilirubin levels are elevated occasionally. Renal dysfunction is uncommon.

Laboratory Diagnosis Demonstration of amastigotes in smears of tissue aspirates is the gold standard for the diagnosis of VL (Fig. 251-1). The

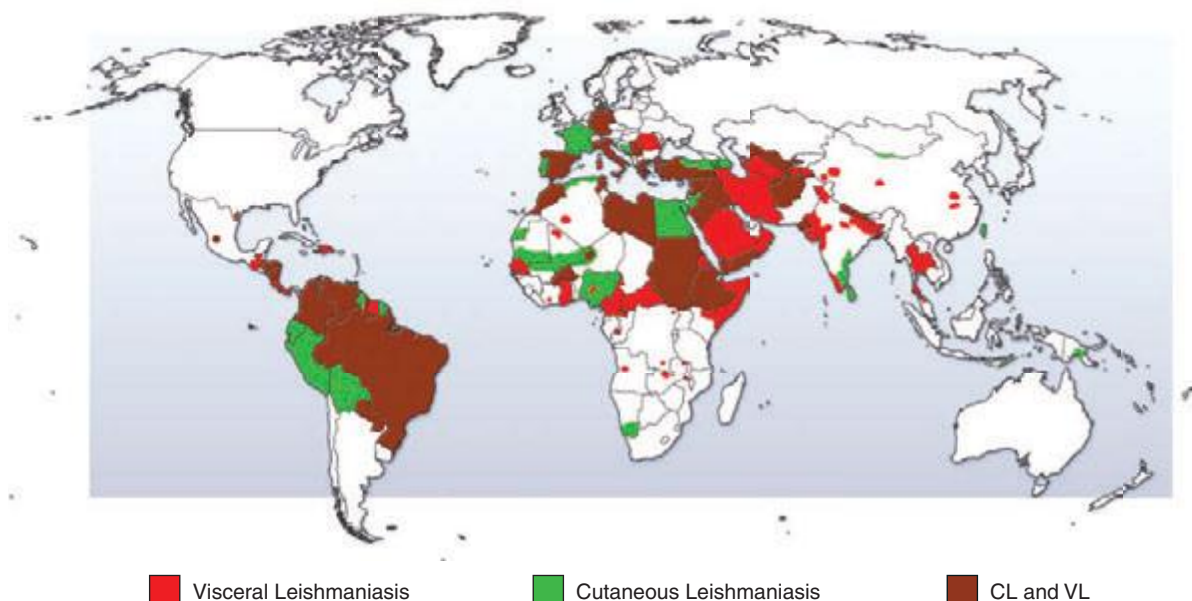



FIGURE 251-2 Worldwide distribution of human leishmaniasis. CL, cutaneous leishmaniasis; VL, visceral leishmaniasis.



FIGURE 251-3 A patient with visceral leishmaniasis has a hugely enlarged spleen visible through the surface of the abdomen. Splenomegaly is the most important feature of visceral leishmaniasis.

sensitivity of splenic smears is >95%, whereas smears of bone marrow (60–85%) and lymph node aspirates (50%) are less sensitive. Culture of tissue aspirates increases sensitivity. Splenic aspiration is invasive and may be dangerous in untrained hands. Several serologic techniques are currently used to detect antibodies to *Leishmania*. An enzyme-linked immunosorbent assay (ELISA) and the indirect immunofluorescent antibody test (IFAT) are used in sophisticated laboratories.

 In the field, however, a rapid immunochromatographic test based on the detection of antibodies to a recombinant antigen (rK39) consisting of 39 amino acids conserved in the kinesin region of *L. infantum* is used worldwide. The test requires only a drop of fingerprick blood or serum, and the result can be read within 15 min. Except in East Africa (where both its sensitivity and its specificity are lower), the sensitivity of the rK39 rapid diagnostic test (RDT) in immunocompetent individuals is ~98% and its specificity is 90%. In Sudan, an RDT based on a new synthetic polypeptide, rK28, was more sensitive (96.8%) and specific (96.2%) than rK39-based RDTs. Qualitative detection of leishmanial nucleic acid by polymerase chain reaction (PCR) and quantitative detection by real-time PCR are confined to specialized laboratories and have yet to be used for routine diagnosis of VL in endemic areas. PCR can distinguish among the major species of *Leishmania* infecting humans.

Differential Diagnosis VL is easily mistaken for malaria. Other febrile illnesses that may mimic VL include typhoid fever, tuberculosis, brucellosis, schistosomiasis, and histoplasmosis. Splenomegaly due to portal hypertension, chronic myeloid leukemia, tropical splenomegaly syndrome, and (in Africa) schistosomiasis may also be confused with VL. Fever with neutropenia or pancytopenia in patients from

an endemic region strongly suggests a diagnosis of VL; hypergammaglobulinemia in patients with long-standing illness strengthens the diagnosis. In nonendemic countries, a careful travel history is essential when any patient presents with fever.

TREATMENT VISCERAL LEISHMANIASIS

GENERAL CONSIDERATIONS

Severe anemia should be corrected by blood transfusion, and other comorbid conditions should be managed promptly. Treatment of VL is complex because the optimal drug, dosage, and duration vary with the endemic region. Despite completing recommended treatment, some patients experience relapse (most often within 6 months), and prolonged follow-up is recommended. A pentavalent antimonial is the drug of choice in most endemic regions of the world, but there is widespread resistance to antimony in the Indian state of Bihar, where either amphotericin B (AmB) deoxycholate or miltefosine is preferred. Dose requirements for AmB are lower in India than in the Americas, Africa, or the Mediterranean region. In Mediterranean countries, where cost is seldom an issue, liposomal AmB is the drug of choice. In immunocompetent patients, relapses are uncommon with AmB in its deoxycholate and lipid formulations. Antileishmanial therapy has recently evolved as new drugs and delivery systems have become available and resistance to antimonial compounds has emerged.

Except for AmB (deoxycholate and lipid formulations), antileishmanial drugs are available in the United States only from the Centers for Disease Control and Prevention.

PENTAVALENT ANTIMONIAL COMPOUNDS

Two pentavalent antimonial (Sb^V) preparations are available: sodium stibogluconate (100 mg of Sb^V/mL) and meglumine antimoniate (85 mg of Sb^V/mL). The daily dose is 20 mg/kg by IV infusion or IM injection, and therapy continues for 28–30 days. Cure rates exceed 90% in Africa, the Americas, and most of the Old World but are <50% in Bihar, India, as a result of resistance. Adverse reactions to Sb^V treatment are common and include arthralgia, myalgia, and elevated serum levels of aminotransferases. Electrocardiographic changes are common. Concave ST-segment elevation is not significant, but prolongation of QT_c to >0.5 s may herald ventricular arrhythmia and sudden death. Chemical pancreatitis is common but usually does not require discontinuation of treatment; severe clinical pancreatitis occurs in immunosuppressed patients.

AMPHOTERICIN B

AmB is currently used as a first-line drug in Bihar, India. In other parts of the world, it is used when initial antimonial treatment fails. Conventional AmB deoxycholate is administered in doses of 0.75–1.0 mg/kg on alternate days for a total of 15 infusions. Fever with chills is an almost universal adverse reaction to AmB infusions. Nausea and vomiting are also common, as is thrombophlebitis in the infused veins. Acute toxicities can be minimized by administration of antihistamines like chlorpheniramine and antipyretic agents like acetaminophen before each infusion. AmB can cause renal dysfunction and hypokalemia and, in rare instances, elicits hypersensitivity reactions, bone marrow suppression, and myocarditis, all of which can be fatal.

The several lipid formulations of AmB developed to replace the deoxycholate formulation are preferentially taken up by reticuloendothelial tissues. Because very little free drug is available to cause toxicity, a large amount of drug can be delivered over a short period. Liposomal AmB has been used extensively to treat VL in all parts of the world. With a terminal half-life of ~150 h, liposomal AmB can be detected in the liver and spleen of animals for several weeks after a single dose. This is the only drug approved by the U.S. Food and Drug Administration (FDA) for the treatment of VL; the regimen is 3 mg/kg daily on days 1–5, 14, and 21 (total dose, 21 mg/kg). However, the total-dose requirement for different regions of the world varies widely. In Asia, it is 10–15 mg/kg; in Africa, ~18 mg/kg;

and in Mediterranean/American regions, ≥ 20 mg/kg. The daily dose is flexible (1–10 mg/kg). In a study in India, a single dose of 10 mg/kg cured infection in 96% of patients. Adverse effects of liposomal AmB are usually mild and include infusion reactions, backache, and occasional reversible nephrotoxicity.

PAROMOMYCIN

Paromomycin (aminosidine) is an aminocyclitol-aminoglycoside antibiotic with antileishmanial activity. Its mechanism of action against *Leishmania* has yet to be established. Paromomycin is approved in India for the treatment of VL at an IM dose of 11 mg of base/kg daily for 21 days; this regimen produces a cure rate of 95%. However, the optimal dose has not been established in other endemic regions. Paromomycin is a relatively safe drug, but some patients develop hepatotoxicity, reversible ototoxicity, and (in rare instances) nephrotoxicity and tetany.

MILTEFOSINE

Miltefosine, an alkylphosphocholine, is the first oral compound approved for the treatment of leishmaniasis. This drug has a long half-life (150–200 h); its mechanism of action is not clearly understood. The recommended therapeutic regimens for patients on the Indian subcontinent are a daily dose of 50 mg for 28 days for patients weighing < 25 kg, a twice-daily dose of 50 mg for 28 days for patients weighing ≥ 25 kg, and 2.5 mg/kg for 28 days for children 2–11 years of age. These regimens have resulted in a cure rate of 94% in India. However, recent studies from the Indian subcontinent indicate a decline in the cure rate. Doses in other regions remain to be established. Because of its long half-life, miltefosine is prone to induce resistance in *Leishmania*. Its adverse effects include mild to moderate vomiting and diarrhea in 40% and 20% of patients, respectively; these reactions usually clear spontaneously after a few days. Rare cases of severe allergic dermatitis, hepatotoxicity, and nephrotoxicity have been reported. Because miltefosine is expensive and is associated with significant adverse events, it is best administered as directly observed therapy to ensure completion of treatment and to minimize the risk of resistance induction. Because miltefosine is teratogenic in rats, its use is contraindicated during pregnancy and (unless contraceptive measures are strictly adhered to for at least 3 months after treatment) in women of childbearing age.

MULTIDRUG THERAPY

Multidrug therapy for leishmaniasis is likely to be preferred in the future. Its potential advantages in VL include (1) better compliance and lower costs associated with shorter treatment courses and decreased hospitalization, (2) less toxicity due to lower drug doses and/or shorter duration of treatment, and (3) a reduced likelihood that resistance to either agent will develop. In a study from India, one dose of liposomal AmB (5 mg/kg) followed by miltefosine for 7 days, paromomycin for 10 days, or both miltefosine and paromomycin simultaneously for 10 days (in their usual daily doses) produced a cure rate of $> 97\%$ (all three combinations). In Africa, a combination of Sb^v and paromomycin given for 17 days was as effective and safe as Sb^v given for 30 days.

PROGNOSIS OF TREATED VL PATIENTS

Recovery from VL is quick. Within a week after the start of treatment, defervescence, regression of splenomegaly, weight gain, and recovery of hematologic parameters are evident. With effective treatment, no parasites are recovered from tissue aspirates at the posttreatment evaluation. Continued clinical improvement over 6–12 months is suggestive of cure. A small percentage of patients (with the exact figure depending on the regimen used) relapse but respond well to treatment with AmB deoxycholate or lipid formulations.

VL IN THE IMMUNOCOMPROMISED HOST

HIV/VL co-infection has been reported from 35 countries. Where both infections are endemic, VL behaves as an opportunistic infection in HIV-1-infected patients. HIV infection can increase the risk of developing VL by severalfold in endemic areas. Co-infected patients usually show the classic signs of VL, but they may present with atypical features due to loss of immunity and involvement of unusual anatomic locations, with, for example, infiltration of the skin, oral mucosa, gastrointestinal tract, lungs, and other organs. Serodiagnostic tests are commonly negative. Parasites can be recovered from unusual sites such as bronchoalveolar lavage fluid and buffy coat. Liposomal AmB is the drug of choice for HIV/VL co-infection—both for primary treatment and for treatment of relapses. A total dose of 40 mg/kg, administered as 4 mg/kg on days 1–5, 10, 17, 24, 31, and 38, is considered optimal and is approved by the FDA, but most patients experience a relapse within 1 year. Pentavalent antimonials and AmB deoxycholate can also be used where liposomal AmB is not accessible. Reconstitution of patients' immunity by antiretroviral therapy has led to a dramatic decline in the incidence of co-infection in the Mediterranean basin. In contrast, HIV/VL co-infection is on the rise in African and Asian countries. Ethiopia is worst affected: up to 30% of VL patients are also infected with HIV. Because restoration of the CD4+ T cell count to $> 200/\mu\text{L}$ does decrease the frequency of relapse, antiretroviral therapy (in addition to antileishmanial therapy) is a cornerstone for the management of HIV/VL co-infection. Secondary prophylaxis with liposomal AmB has been shown to delay relapses, but no regimen has been established as optimal.

POST-KALA-AZAR DERMAL LEISHMANIASIS

On the Indian subcontinent and in Sudan and other East African countries, 2–50% of patients develop skin lesions concurrent with or after the cure of VL. Most common are hypopigmented macules, papules, and/or nodules or diffuse infiltration of the skin and sometimes of the oral mucosa. The African and Indian diseases differ in several respects; important features of post-kala-azar dermal leishmaniasis (PKDL) in these two regions are listed in [Table 251-2](#), and disease in an Indian patient is depicted in [Fig. 251-4](#).

In PKDL, parasites are scanty in hypopigmented macules but may be seen and cultured more easily from nodular lesions. Cellular infiltrates are heavier in nodules than in macules. Lymphocytes are the dominant cells; next most common are histiocytes and plasma

TABLE 251-2 CLINICAL, EPIDEMIOLOGIC, AND THERAPEUTIC FEATURES OF POST-KALA-AZAR DERMAL LEISHMANIASIS: EAST AFRICA AND THE INDIAN SUBCONTINENT

Feature	East Africa	Indian Subcontinent
Most affected country	Sudan and South Sudan	Bangladesh
Incidence among patients with VL	~50%	~2–17%
Interval between VL and PKDL	During VL to 6 months	6 months to 3 years
Age distribution	Mainly children	Any age
History of prior VL	Yes	Not necessarily
Rashes of PKDL in presence of active VL	Yes	No
Treatment with sodium stibogluconate	2–3 months	2–4 months
Natural course	Spontaneous cure in majority of patients	Spontaneous cure in minority of patients

Abbreviations: PKDL, post-kala-azar dermal leishmaniasis; VL, visceral leishmaniasis.



FIGURE 251-4 Post-kala-azar dermal leishmaniasis in an Indian patient. Note nodules of varying size involving the entire face. The face is erythematous, and the surface of some of the large nodules is discolored.

cells. In about half of cases, epithelioid cells—scattered individually or forming compact granulomas—are seen. The diagnosis is based on history and clinical findings, but rK39 and other serologic tests are positive in most cases. Indian PKDL is treated with pentavalent antimonials for 60–120 days. This prolonged course frequently leads to noncompliance. The alternative—several courses of AmB spread over several months—is expensive and unacceptable for most patients. Oral miltefosine for 12 weeks, in the usual daily doses, cures most patients with Indian PKDL. In East Africa, a majority of patients experience spontaneous healing. In those with persistent lesions, the response to 60 days of treatment with a pentavalent antimonial is good.

CUTANEOUS LEISHMANIASIS

CL can be broadly divided into Old World and New World forms. Old World CL caused by *Leishmania tropica* is anthroponotic and is confined to urban or suburban areas throughout its range. Zoonotic CL is most commonly due to *Leishmania major*, which naturally parasitizes several species of desert rodents that act as reservoirs over wide areas of the Middle East, Africa, and central Asia. Local outbreaks of human disease are common. Major outbreaks currently affect Afghanistan and Pakistan in association with refugees and population movement. CL is increasingly seen in tourists and military personnel on mission in CL-endemic regions of countries like Afghanistan and as a coinfection in HIV-infected patients. *Leishmania aethiopica* is restricted to the highlands of Ethiopia, Kenya, and Uganda, where it is a natural parasite of hyraxes. New World CL is mainly zoonotic and is most often caused by *Leishmania mexicana*, *Leishmania (Viannia) panamensis*, and *Leishmania amazonensis*. A wide range of forest animals act as reservoirs, and human infections with these species are predominantly rural. As a result of extensive urbanization and deforestation, *Leishmania (Viannia) braziliensis* has adapted to peridomestic and urban animals, and CL due to this organism is increasingly becoming

an urban disease. In the United States, a few cases of CL have been acquired indigenously in Texas.

Immunopathogenesis As in VL, the proinflammatory (T_H1) response in CL may result in either asymptomatic or subclinical infection. However, in some individuals, the immune response causes ulcerative skin lesions, the majority of which heal spontaneously, leaving a scar. Healing is usually followed by immunity to reinfection with that species of parasite.

Clinical Features A few days or weeks after the bite of a sandfly, a papule develops and grows into a nodule that ulcerates over some weeks or months. The base of the ulcer, which is usually painless, consists of necrotic tissue and crusted serum, but secondary bacterial infection sometimes occurs. The margins of the ulcer are raised and indurated. Lesions may be single or multiple and vary in size from 0.5 to >3 cm (Fig. 251-5). Lymphatic spread and lymph gland involvement may be palpable and may precede the appearance of the skin lesion. There may be satellite lesions, especially in *L. major* and *L. tropica* infections. The lesions usually heal spontaneously after 2–15 months. Lesions due to *L. major* and *L. mexicana* tend to heal rapidly, whereas those due to *L. tropica* and parasites of subspecies *Viannia* heal more slowly. In CL caused by *L. tropica*, new lesions—usually scaly, erythematous papules and nodules—develop in the center or periphery of a healed sore, a condition known as *leishmaniasis recidivans*. Lesions of *L. mexicana* and *Leishmania (Viannia) peruviana* closely resemble those seen in the Old World; however, lesions on the pinna of the ear are common, chronic, and destructive in the former infections. *L. mexicana* is responsible for chilero's ulcer, the so-called self-healing sore of Mexico. CL lesions on exposed body parts (e.g., the face and hands), permanent scar formation, and social stigmatization may cause anxiety and depression and may affect the quality of life of CL patients.

Differential Diagnosis A typical history (an insect bite followed by the events leading to ulceration) in a resident of or a traveler to an endemic focus strongly suggests CL. Cutaneous tuberculosis, fungal infections, leprosy, sarcoidosis, and malignant ulcers are sometimes mistaken for CL.

Laboratory Diagnosis Demonstration of amastigotes in material obtained from a lesion remains the diagnostic gold standard. Microscopic examination of slit skin smears, aspirates, or biopsies of the lesion is used for detection of parasites. Culture of smear or biopsy material may yield *Leishmania*. PCR is more sensitive than microscopy and culture and allows identification of *Leishmania* to the species level. This information is important in decisions about therapy because responses to treatment can vary with the species. Isoenzyme profiling is used to determine species for research purposes.



FIGURE 251-5 Cutaneous leishmaniasis in a Bolivian child. There are multiple ulcers resulting from several sandfly bites. The edges of the ulcers are raised. (Courtesy of P. Desjeux, Retired Medical Officer, World Health Organization, Geneva, Switzerland.)

Although lesions heal spontaneously in the majority of cases, their spread or persistence indicates that treatment may be needed. One or a few small lesions due to “self-healing species” can be treated with topical agents. Systemic treatment is required for lesions over the face, hands, or joints; multiple lesions; large ulcers; lymphatic spread; New World CL with the potential for development of ML; and CL in HIV co-infected patients.

A pentavalent antimonial is the first-line drug for all forms of CL and is used in a dose of 20 mg/kg for 20 days, as for VL. The exceptions to this rule are CL caused by *Leishmania (Viannia) guyanensis*, for which pentamidine isethionate is the drug of choice (two injections of 4 mg of salt/kg separated by a 48-h interval), and CL due to *L. aethiopicum*, which responds to paromomycin (16 mg/kg daily) but not to antimonials. Relapses usually respond to a second course of treatment. In Peru, topical imiquimod (5–7.5%) plus parenteral antimonials have been shown to cure CL more rapidly than antimonials alone. Azoles and triazoles have been used with mixed responses in both Old and New World CL but have not been adequately assessed for this indication in clinical trials. In *L. major* infection, oral fluconazole (200 mg/d for 6 weeks) resulted in a higher rate of cure than placebo (79% vs 34%) and also cured infection faster. Adverse effects include gastrointestinal symptoms and hepatotoxicity. Ketoconazole (600 mg/d for 28 days) is 76–90% effective in CL due to *L. (V.) panamensis* and *L. mexicana* in Panama and Guatemala. Miltefosine has been used in CL in doses of 2.5 mg/kg for 28 days. This agent is effective against *L. major* infections. In Colombia, where CL is due to *L. (V.) panamensis*, miltefosine was also effective, with a cure rate of 91%. For *L. (V.) braziliensis* infections, however, the results with miltefosine are less consistent. In Brazil, miltefosine cured 71% of patients with *L. (V.) guyanensis* infection. Other drugs, such as dapsone, allopurinol, rifampin, azithromycin, and pentoxifylline, have been used either alone or in combinations, but most of the relevant studies have had design limitations that preclude meaningful conclusions.

Small lesions (≤ 3 cm in diameter) may conveniently be treated weekly until cure with an intralesional injection of a pentavalent antimonial at a dose adequate to blanch the lesion (0.2–2.0 mL). An ointment containing 15% paromomycin sulfate, either alone or with 0.5% gentamicin or 12% methylbenzoniun chloride, cured 70–82% of lesions due to *L. major* in 20 days and may be suitable for lesions caused by other species. Heat therapy with an FDA-approved radio-frequency generator and cryotherapy with liquid nitrogen have also been used successfully.

Diffuse Cutaneous Leishmaniasis (DCL) DCL is a rare form of leishmaniasis caused by *L. amazonensis* and *L. mexicana* in South and Central America and by *L. aethiopicum* in Ethiopia and Kenya. DCL is characterized by the lack of a cell-mediated immune response to the parasite, the uncontrolled multiplication of which thus continues unabated. The DTH response does not develop, and lymphocytes do not respond to leishmanial antigens in vitro. DCL patients have a polarized immune response with high levels of immunosuppressive cytokines, including IL-10, transforming growth factor (TGF) β , and IL-4, and low concentrations of IFN- γ . Profound immunosuppression leads to widespread cutaneous disease. Lesions may initially be confined to the face or a limb but spread over months or years to other areas of the skin. They may be symmetrically or asymmetrically distributed and include papules, nodules, plaques, and areas of diffuse infiltration. These lesions do not ulcerate. The overlying skin is usually erythematous in pale-skinned patients. The lesions are teeming with parasites, which are therefore easy to recover. DCL does not heal spontaneously and is difficult to treat. If relapse and drug resistance are to be prevented, treatment should be continued for some time after lesions have healed and parasites can no longer be isolated. In the New World, repeated 20-day courses of pentavalent antimonials are given, with an intervening drug-free period of 10 days. Miltefosine has been used for several

months with a good initial response. Combinations should be tried. In Ethiopia, a combination of paromomycin (14 mg/kg per day) and sodium stibogluconate (10 mg/kg per day) is effective.

MUCOSAL LEISHMANIASIS

The subgenus *Viannia* is widespread from the Amazon basin to Paraguay and Costa Rica and is responsible for deep sores and for ML (Table 251-1). In *L. (V.) braziliensis* infections, cutaneous lesions may be simultaneously accompanied by mucosal spread of the disease or followed by spread years later. ML is typically caused by *L. (V.) braziliensis* and rarely by *L. amazonensis*, *L. (V.) guyanensis*, and *L. (V.) panamensis*. Young men with chronic lesions of CL are at particular risk. Overall, ~3% of infected persons develop ML. Not every patient with ML has a history of prior CL. ML is almost entirely confined to the Americas. In rare cases, ML may also be caused by Old World species like *L. major*, *L. infantum (L. chagasi)*, or *L. donovani*.

Immunopathogenesis and Clinical Features The immune response is polarized toward a T_H1 response, with marked increases of IFN- γ and TNF- α and varying levels of T_H2 cytokines (IL-10 and TGF- β). Patients have a stronger DTH response with ML than with CL, and their peripheral-blood mononuclear cells respond strongly to leishmanial antigens. The parasite spreads via the lymphatics or the bloodstream to mucosal tissues of the upper respiratory tract. Intense inflammation leads to destruction, and severe disability ensues. Lesions in or around the nose or mouth (espundia; Fig. 251-6) are the typical presentation of ML. Patients usually provide a history of self-healed CL preceding ML by 1–5 years. Typically, ML presents as nasal stuffiness and bleeding followed by destruction of nasal cartilage, perforation of the nasal septum, and collapse of the nasal bridge. Subsequent involvement of the pharynx and larynx leads to difficulty in swallowing and phonation.



FIGURE 251-6 Mucosal leishmaniasis in a Brazilian patient. There is extensive inflammation around the nose and mouth, destruction of the nasal mucosa, ulceration of the upper lip and nose, and destruction of the nasal septum. (Courtesy of R. Dietz, Universidade Federal do Espírito Santo, Vitória, Brazil.)

1394 The lips, cheeks, and soft palate may also be affected. Secondary bacterial infection is common, and aspiration pneumonia may be fatal. Despite the high degree of T_{H1} immunity and the strong DTH response, ML does not heal spontaneously.

Laboratory Diagnosis Tissue biopsy is essential for identification of parasites, but the rate of detection is poor unless PCR techniques are used. The strongly positive DTH response fails to distinguish between past and present infection.

TREATMENT MUCOSAL LEISHMANIASIS

The regimen of choice is a pentavalent antimonial agent administered at a dose of 20 mg of Sb^V/kg for 30 days. Patients with ML require long-term follow-up with repeated oropharyngeal and nasal examination. With failure of therapy or relapse, patients may receive another course of an antimonial but then become unresponsive, presumably because of resistance in the parasite. In this situation, AmB should be used. An AmB deoxycholate dose totaling 25–45 mg/kg is appropriate. There are no controlled trials of liposomal AmB, but administration of 2–3 mg/kg for 20 days is considered adequate. Miltefosine (2.5 mg/kg for 28 days) cured 71% of ML patients in Bolivia. The more extensive the disease, the worse the prognosis; thus prompt, effective treatment and regular follow-up are essential.

PREVENTION OF LEISHMANIASIS

No vaccine is available for any form of leishmaniasis. Inoculation with live *L. major* (“leishmanization”) is practiced in Iran. Anthroponotic leishmaniasis is controlled by case finding, treatment, and vector control with insecticide-impregnated bed nets and curtains and residual insecticide spraying. Control of zoonotic leishmaniasis is more difficult. Use of insecticide-impregnated collars for dogs, treatment of infected domestic dogs, and culling of street dogs are measures that have been used with uncertain efficacy to prevent transmission of *L. infantum*. In Brazil, a canine vaccine has been found to promote a decrease in the human and canine incidence of zoonotic VL. Two vaccines, Leishmune[®] and Leish-Tec[®], are licensed in Brazil; Leishmune provides significant protection to vaccinated dogs. CaniLeish[®] is the first licensed canine vaccine developed in Europe. Personal prophylaxis with bed nets and repellants may reduce the risk of CL infections in the New World.

In 10–30% of chronically infected patients, cardiac and/or gastrointestinal symptoms develop that can lead to serious morbidity and even death.

LIFE CYCLE AND TRANSMISSION

T. cruzi is transmitted among its mammalian hosts by hematophagous triatomine insects, often called reduviid bugs. The insects become infected by sucking blood from animals or humans with circulating parasites. Ingested organisms multiply in the gut of the triatomines, and infective forms are discharged with the feces at the time of subsequent blood meals. Transmission to a second vertebrate host occurs when breaks in the skin, mucous membranes, or conjunctivae become contaminated with bug feces that contain infective parasites. *T. cruzi* can also be transmitted by transfusion of blood donated by infected persons, by organ transplantation, from mother to unborn child, by ingestion of contaminated food or drink, and in laboratory accidents.

PATHOLOGY

Initial infection at the site of parasite entry is characterized by local histologic changes that include the presence of parasites within leukocytes and cells of subcutaneous tissues and the development of interstitial edema, lymphocytic infiltration, and reactive hyperplasia of adjacent lymph nodes. After dissemination of the organisms through the lymphatics and the bloodstream, primarily muscles (including the myocardium) (Fig. 252-1) and ganglion cells may become heavily parasitized. The characteristic pseudocysts present in sections of infected tissues are intracellular aggregates of multiplying parasites.

In persons with chronic *T. cruzi* infections who develop related clinical manifestations, the heart is the organ most commonly affected. Changes include thinning of the ventricular walls, biventricular enlargement, apical aneurysms, and mural thrombi. Widespread lymphocytic infiltration, diffuse interstitial fibrosis, and atrophy of myocardial cells are often apparent. Although parasites are difficult to find in myocardial tissue by conventional histologic methods, more sensitive techniques of parasite detection, such as immunohistochemistry and polymerase chain reaction (PCR), have more frequently demonstrated *T. cruzi* antigens and parasite DNA in chronic lesions. Conduction-system abnormalities often affect the right branch and the left anterior branch of the bundle of His. In chronic Chagas disease of the gastrointestinal tract (*megadisease*), the esophagus and colon may exhibit varying degrees of dilation. On microscopic examination, focal inflammatory lesions with lymphocytic infiltration are seen, and the number of neurons in the myenteric plexus may be markedly reduced. Accumulating evidence implicates the persistence of parasites and the accompanying chronic inflammation—rather than autoimmune mechanisms—as the basis for the pathology in patients with chronic *T. cruzi* infection.

252 Chagas Disease and African Trypanosomiasis

Louis V. Kirchhoff, Anis Rassi Jr.

Although the genus *Trypanosoma* contains many species of protozoans, only *T. cruzi*, *T. brucei gambiense*, and *T. brucei rhodesiense* cause disease in humans. *T. cruzi* is the etiologic agent of Chagas disease in the Americas; *T. b. gambiense* and *T. b. rhodesiense* cause African trypanosomiasis.

CHAGAS DISEASE (AMERICAN TRYPANOSOMIASIS)

DEFINITION

Chagas disease, or American trypanosomiasis, is a zoonosis caused by the protozoan parasite *T. cruzi*. Acute Chagas disease is usually a mild febrile illness that results from initial infection with the organism. After spontaneous resolution of the acute illness, most infected persons remain for life in the indeterminate phase of chronic Chagas disease, which is characterized by subpatent parasitemia, easily detectable IgG antibodies to *T. cruzi*, and an absence of associated signs and symptoms.

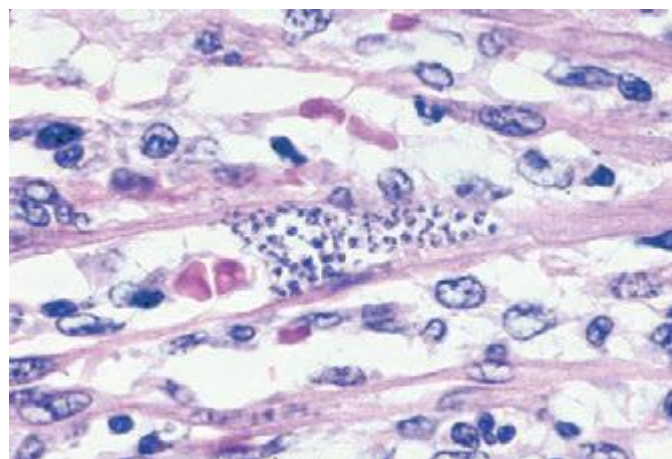


FIGURE 252-1 *Trypanosoma cruzi* in the heart muscle of a child who died of acute Chagas myocarditis. An infected myocyte containing several dozen *T. cruzi* amastigotes is in the center of the field (hematoxylin and eosin, 900 ×).

EPIDEMIOLOGY



T. cruzi is found only in the Americas. Wild and domestic mammals harboring *T. cruzi* and infected triatomines are found in spotty distributions from the southern United States to southern Argentina. Humans become involved in the cycle of transmission when infected vectors take up residence in the primitive wood, adobe, and stone houses common in much of Latin America. Thus human *T. cruzi* infection is a health problem primarily among the poor in rural areas of Mexico and Central and South America. Most new *T. cruzi* infections in rural settings occur in children, but the incidence is unknown because most cases go undiagnosed. Historically, transfusion-associated transmission of *T. cruzi* was a serious public health problem in many endemic countries. Transmission by this route has been largely eliminated, however, as effective programs for serologic screening of donated blood have been implemented. Several dozen patients with HIV and chronic *T. cruzi* infections who underwent acute recrudescence of the latter have been described. These patients generally presented with *T. cruzi* brain abscesses, a manifestation of the illness that does not occur in immunocompetent persons. Currently, it is estimated that 8 million people are chronically infected with *T. cruzi* and that 14,000 deaths due to the illness occur each year. The resulting morbidity and mortality make Chagas disease the most important parasitic disease burden in Latin America.

In recent years, the rate of *T. cruzi* transmission has decreased markedly in several endemic countries as a result of successful programs involving vector control, screening of donated blood, and education of at-risk populations. A major program, which began in 1991 in the “southern cone” nations of South America (Uruguay, Paraguay, Bolivia, Brazil, Chile, and Argentina), has provided the framework for much of this progress. Uruguay and Chile were certified free of transmission by the main domiciliary vector species (*Triatoma infestans*) in the late 1990s, and Brazil was declared transmission-free in 2006. Transmission has been reduced markedly in Argentina as well. Similar control programs have been initiated in the countries of northern South America and in the Central American nations.

Acute Chagas disease is rare in the United States, where 22 cases of autochthonous transmission and seven instances of transmission by blood transfusion have been reported. Moreover, *T. cruzi* was transmitted to five recipients of organs from three *T. cruzi*-infected donors, two of whom became infected through cardiac transplants. Acute Chagas disease has been reported in only one tourist returning to the United States from Latin America, although three such instances have been reported in Europe as well as one in Canada. In contrast, the prevalence of chronic *T. cruzi* infections in the United States has increased considerably in recent years. An estimated 23 million immigrants from Chagas-endemic countries currently live in the United States, ~17 million of whom are Mexicans. The total number of *T. cruzi*-infected persons living in the United States is estimated to be 300,000. Screening of the U.S. blood supply for *T. cruzi* infection began in January 2007. The overall prevalence of *T. cruzi* infection among donors is ~1 in 13,300, and to date nearly 3000 infected donors have been identified and deferred permanently (see “Diagnosis,” below).

CLINICAL COURSE

The first signs of acute Chagas disease develop at least 1 week after invasion by the parasites. When the organisms enter through a break in the skin, an indurated area of erythema and swelling (the *chagoma*), accompanied by local lymphadenopathy, may appear. *Romaña sign*—the classic finding in acute Chagas disease, which consists of unilateral painless edema of the palpebrae and periocular tissues—can result when the conjunctiva is the portal of entry. These initial local signs may be followed by malaise, fever, anorexia, and edema of the face and lower extremities. Generalized lymphadenopathy and hepatosplenomegaly may develop. Severe myocarditis develops rarely; most deaths in acute Chagas disease are due to heart failure. Neurologic signs are not common, but meningoencephalitis occurs occasionally, especially in children <2 years old. Usually within 4–8 weeks, acute signs and symptoms resolve spontaneously in virtually all patients, with commencement of the asymptomatic or indeterminate form of chronic *T. cruzi* infection.

Symptomatic chronic Chagas disease becomes apparent years or even decades after the initial infection. The heart is commonly involved, and symptoms are caused by rhythm disturbances, segmental or dilated cardiomyopathy, and thromboembolism. Right bundle branch block is a common electrocardiographic abnormality, but other types of intraventricular and atrioventricular blocks, premature ventricular contractions, and tachy- and bradyarrhythmias occur frequently. Cardiomyopathy often results in biventricular heart failure, with a predominance of right-sided failure at advanced stages. Embolization of mural thrombi to the brain or other areas may take place. Sudden death is the main cause of death in Chagas heart disease; congestive heart failure and stroke are next most common. Patients with megaesophagus suffer from dysphagia, odynophagia, chest pain, and regurgitation. Aspiration can occur (especially during sleep) in patients with severe esophageal dysfunction, and repeated episodes of aspiration pneumonitis are common. Weight loss, cachexia, and pulmonary infection can result in death. Patients with megacolon are plagued by abdominal pain and chronic constipation, which predisposes to fecaloma formation. Advanced megacolon can cause obstruction, volvulus, septicemia, and death.

DIAGNOSIS

The diagnosis of acute Chagas disease requires the detection of parasites. Microscopic examination of fresh anticoagulated blood or the buffy coat is the simplest way to see the motile organisms. Parasites also can be seen in Giemsa-stained thin and thick blood smears. Microhematocrit tubes containing acridine orange as a stain can be used for the same purpose. When used repeatedly by experienced personnel, all of these methods yield positive results in a high proportion of cases of acute Chagas disease. Serologic testing does not play a major role in diagnosing acute Chagas disease. PCR assays often give positive results in infected patients in whom traditional parasitologic tests are negative, including infants with congenital Chagas disease.

Chronic Chagas disease is diagnosed by the detection of specific IgG antibodies that bind to *T. cruzi* antigens. Demonstration of the parasite is not of primary importance. In Latin America, ~30 assays are commercially available, including several based on recombinant antigens. Although these tests usually show good sensitivity and reasonable specificity, false-positive reactions may occur—typically with samples from patients who have other infectious and parasitic diseases or autoimmune disorders. In addition, confirmatory testing has presented a persistent challenge. For these reasons, the World Health Organization recommends that specimens be tested in at least two assays and that well-characterized positive and negative comparison samples be included in each run. The Chagas radioimmuno precipitation assay (RIPA), a highly sensitive and specific confirmatory method for detecting antibodies to *T. cruzi*, is approved under the Clinical Laboratory Improvement Amendment and available in the laboratory of one of the authors (L.V.K.). In 2006, the U.S. Food and Drug Administration (FDA) approved a test to screen blood and organ donors for *T. cruzi* infection (Ortho *T. cruzi* ELISA Test System; Ortho-Clinical Diagnostics, Raritan, NJ). Since January 2007, the vast majority of U.S. blood donors have been screened, and positive units have undergone confirmatory testing with the Chagas RIPA. A second test for donor screening was approved by the FDA in 2010 (Abbott PRISM® Chagas Assay; Abbott Laboratories, Abbott Park, IL), as was an enzyme strip assay (Abbott ESA Chagas) in 2011. The use of PCR assays to detect *T. cruzi* DNA in chronically infected persons has been studied extensively; unfortunately, the sensitivity of this approach has not been shown to be reliably greater than that of serology.


TREATMENT CHAGAS DISEASE

Therapy for Chagas disease is still unsatisfactory. For many years now, only two drugs—nifurtimox and benznidazole—have been available for this purpose. Regrettably, both drugs lack efficacy and may cause bothersome side effects.

In acute Chagas disease, nifurtimox markedly reduces the duration of symptoms and parasitemia and decreases the mortality rate.

1396 Nevertheless, limited studies have shown that only ~70% of acute infections are cured by a full course of treatment. Common adverse effects of nifurtimox include anorexia, nausea, vomiting, weight loss, and abdominal pain. Neurologic reactions to the drug may include restlessness, disorientation, insomnia, twitching, paresthesia, polyneuritis, and seizures. These symptoms usually disappear when the dosage is reduced or treatment is discontinued. The recommended daily dosage is 8–10 mg/kg for adults, 12.5–15 mg/kg for adolescents, and 15–20 mg/kg for children 1–10 years of age. The drug should be given orally in four divided doses each day, and therapy should be continued for 90–120 days. Nifurtimox is available from the Drug Service of the Centers for Disease Control and Prevention (CDC) in Atlanta (telephone number, 404-639-3670).


The efficacy of benznidazole is similar or even superior to that of nifurtimox. A cure rate of >90% among congenitally infected infants treated before their first birthday has been reported. Adverse effects include rash, peripheral neuropathy, and rare instances of granulocytopenia. The recommended oral dosage is 5 mg/kg per day for 60 days for adults and 5–10 mg/kg per day for 60 days for children, with administration of two or three divided doses. Benznidazole is generally considered the drug of choice in Latin America.

 The question of whether adults in the indeterminate or chronic symptomatic phase of Chagas disease should be treated with nifurtimox or benznidazole has been debated for years. The fact that cure rates in persons with long-established chronic infection are notably inferior to those in patients with acute or recent chronic infection is central to this controversy. No convincing evidence from randomized controlled trials indicates that nifurtimox or benznidazole treatment of adults in the indeterminate or chronic symptomatic phase reduces either the appearance or progression of symptoms or mortality rates. On the basis of results of some observational studies, a panel of experts convened by the CDC in 2006 recommended that adults <50 years old with presumably long-standing indeterminate *T. cruzi* infections—or even with mild to moderate disease—be offered treatment. A large randomized clinical trial (the BENEFIT multicenter trial) designed to assess the parasitologic and clinical efficacy of benznidazole in 2856 adults (18–75 years old) with chronic Chagas heart disease (without advanced lesions) is being performed in Brazil, Argentina, Colombia, Bolivia, and El Salvador, but results will not be available until 2015. In contrast, randomized studies have shown that treatment of children is useful, and the current consensus of Latin American authorities and the CDC panel of experts is that all *T. cruzi*-infected persons up to 18 years old and all adults known to have become infected recently should be given benznidazole or nifurtimox.

The usefulness of antifungal azoles for the treatment of Chagas disease has been studied in laboratory animals and to a lesser extent in humans. To date, none of these drugs has exhibited a level of anti-*T. cruzi* activity that would justify its use in humans. Several newer drugs in this class have shown promise in animal studies and are currently being evaluated in phase 2 clinical trials.

Patients who develop cardiac and/or gastrointestinal disease in association with *T. cruzi* infection should be referred to appropriate subspecialists for further evaluation and treatment. Pacemakers can be useful in patients with ominous arrhythmias. The usefulness of implantable cardioverter defibrillators in persons with Chagas heart disease has not been established and currently is being studied in a prospective randomized trial. Cardiac transplantation is an option for patients with end-stage chagasic cardiomyopathy; more than 150 such transplantations have been done in Brazil and the United States. The survival rate among Chagas disease cardiac transplant recipients seems to be higher than that among persons receiving cardiac transplants for other reasons. This better outcome may be due to the fact that lesions are limited to the heart in most patients with symptomatic chronic Chagas disease.

PREVENTION

 Because drug therapy has limitations and vaccines are not available, the control of *T. cruzi* transmission in endemic countries depends on the reduction of domiciliary vector populations by spraying of insecticides, improvements in housing, and education of at-risk persons. As noted above, these measures, coupled with serologic screening of blood donors, have markedly reduced transmission of the parasite in many endemic countries. Tourists would be wise to avoid sleeping in dilapidated houses in rural areas of endemic countries. Mosquito nets and insect repellent can provide additional protection.

In view of the possibly serious consequences of chronic *T. cruzi* infection, it would be prudent for all immigrants from endemic regions who are living in the United States to be tested for evidence of infection. Identification of persons harboring the parasite would permit periodic electrocardiographic monitoring, which is important to detect incipient heart disease and guide further diagnostic studies and treatment. The possibility of congenital transmission is yet another justification for screening. *T. cruzi* is classified as a Risk Group 2 agent in the United States and a Risk Group 3 agent in some European countries. Laboratory staff should work with the parasite or infected vectors and mammals at containment levels consistent with the risk group designation in their areas.

SLEEPING SICKNESS (AFRICAN TRYPANOSOMIASIS)

DEFINITION

Sleeping sickness, or human African trypanosomiasis (HAT), is caused by flagellated protozoan parasites that belong to the *T. brucei* complex and are transmitted to humans by tsetse flies. In untreated patients, the trypanosomes first cause a febrile illness that is followed months or years later by progressive neurologic impairment and death.

THE PARASITES AND THEIR TRANSMISSION

The East African (*rhodesiense*) and the West African (*gambiense*) forms of sleeping sickness are caused, respectively, by two trypanosome subspecies: *T. b. rhodesiense* and *T. b. gambiense*. These subspecies are morphologically indistinguishable but cause illnesses that are epidemiologically and clinically distinct (Table 252-1). The parasites are transmitted by blood-sucking tsetse flies of the genus *Glossina*. The insects acquire the infection when they ingest blood from infected mammalian hosts. After many cycles of multiplication in the midgut of the vector, the parasites migrate to the salivary glands. Their transmission takes place when they are inoculated into a mammalian host during a subsequent blood meal. The injected trypanosomes multiply in the blood (Fig. 252-2) and other extracellular spaces and evade immune destruction for long periods by undergoing antigenic variation, a process driven by gene switching in which the antigenic structure of the organisms' surface coat of glycoproteins changes periodically.

TABLE 252-1 COMPARISON OF WEST AFRICAN AND EAST AFRICAN TRYPANOSOMIASSES

Point of Comparison	West African (<i>gambiense</i>)	East African (<i>rhodesiense</i>)
Organism	<i>T. b. gambiense</i>	<i>T. b. rhodesiense</i>
Vectors	Tsetse flies (palpalis group)	Tsetse flies (morsitans group)
Primary reservoir	Humans	Antelope and cattle
Human illness	Chronic (late CNS disease)	Acute (early CNS disease)
Duration of illness	Months to years	<9 months
Lymphadenopathy	Prominent	Minimal
Parasitemia	Low	High
Epidemiology	Rural populations	Workers in wild areas, rural populations, tourists in game parks

Abbreviation: CNS, central nervous system.

Source: Reprinted with permission from LV Kirchhoff, in GL Mandell et al (eds): *Principles and Practice of Infectious Diseases*, 7th ed. Philadelphia, Elsevier Churchill Livingstone, 2010.

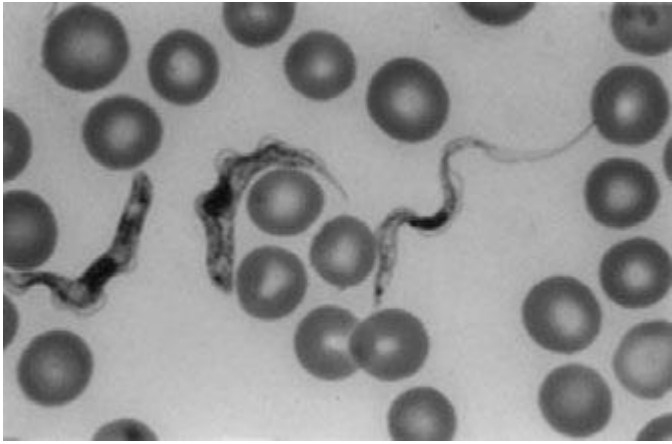


FIGURE 252-2 *Trypanosoma brucei rhodesiense* parasites in rat blood. The slender parasite is thought to be the form that multiplies in mammalian hosts, whereas the stumpy forms are nondividing and are capable of infecting insect vectors (Giemsa, 1200 \times). (Courtesy of Dr. G. A. Cook, Madison, WI; with permission.)

PATHOGENESIS AND PATHOLOGY

A self-limited inflammatory lesion (*trypanosomal chancre*) may appear a week or so after the bite of an infected tsetse fly. A systemic febrile illness then evolves as the parasites are disseminated through the lymphatics and bloodstream. Systemic HAT without central nervous system (CNS) involvement is generally referred to as *stage 1 disease*. In this stage, widespread lymphadenopathy and splenomegaly reflect marked lymphocytic and histiocytic proliferation and invasion of morular cells, which are plasmacytes that may be involved in the production of IgM. Endarteritis, with perivascular infiltration of both parasites and lymphocytes, may develop in lymph nodes and the spleen. Myocarditis develops frequently in patients with stage 1 disease and is especially common in *T. b. rhodesiense* infections.

Hematologic manifestations that accompany stage 1 HAT include moderate leukocytosis, thrombocytopenia, and anemia. High levels of immunoglobulins, consisting primarily of polyclonal IgM, are a constant feature, and heterophile antibodies, antibodies to DNA, and rheumatoid factor are often detected. High levels of antigen-antibody complexes may play a role in the tissue damage and increased vascular permeability that facilitate dissemination of the parasites.

Stage 2 disease involves invasion of the CNS. The presence of trypanosomes in perivascular areas is accompanied by intense infiltration of mononuclear cells. Abnormalities in cerebrospinal fluid (CSF) include increased pressure, elevated total protein concentration, and pleocytosis. In addition, trypanosomes are frequently found in CSF.

EPIDEMIOLOGY



The trypanosomes that cause sleeping sickness are found only in sub-Saharan Africa. After its near-eradication in the mid-1960s, sleeping sickness underwent a resurgence in the 1990s, primarily in Uganda, Sudan, the Central African Republic, the Democratic Republic of the Congo, and Angola. A subsequent increase in control activities reduced the incidence in many endemic areas, however, and in 2009 fewer than 10,000 cases were reported to the World Health Organization. Although underreporting is a persistent problem, the level of control achieved to date was the basis for convening a panel of experts in 2009 to develop a vision for eradication of HAT.

Humans are the only reservoir of *T. b. gambiense*, which occurs in widely distributed foci in tropical rain forests of Central and West Africa. *Gambiense* trypanosomiasis is primarily a problem in rural populations; tourists rarely become infected. Trypanotolerant antelope species in savanna and woodland areas of Central and East Africa are the principal reservoir of *T. b. rhodesiense*. Cattle can also be infected with this and other trypanosome species but generally succumb to the infection. Because risk results from contact with tsetse flies that feed on

wild animals, humans acquire *T. b. rhodesiense* infection only incidentally, usually while visiting or working in areas where infected game and vectors are present. Roughly one or two imported cases of HAT acquired in East African parks are reported to the CDC each year.

CLINICAL COURSE

A painful trypanosomal chancre appears in some patients at the site of inoculation of the parasite. Hematogenous and lymphatic dissemination (stage 1 disease) is marked by the onset of fever. Typically, bouts of high temperatures lasting several days are separated by afebrile periods. Lymphadenopathy is prominent in *T. b. gambiense* trypanosomiasis. The nodes are discrete, movable, rubbery, and nontender. Cervical nodes are often visible, and enlargement of the nodes of the posterior cervical triangle, or *Winterbottom's sign*, is a classic finding. Pruritus and maculopapular rashes are common. Inconstant findings include malaise, headache, arthralgias, weight loss, edema, hepatosplenomegaly, and tachycardia. The differential diagnosis of stage 1 HAT includes many diseases that are common in the tropics and are associated with fevers. HIV infection, malaria, and typhoid fever are common in populations at risk for HAT and need to be considered.

CNS invasion (stage 2 disease) is characterized by the insidious development of protean neurologic manifestations that are accompanied by progressive abnormalities in the CSF. A picture of progressive indifference and daytime somnolence develops (hence the designation "sleeping sickness"), sometimes alternating with restlessness and insomnia at night. A listless gaze accompanies a loss of spontaneity, and speech may become halting and indistinct. Extrapyramidal signs may include choreiform movements, tremors, and fasciculations. Ataxia is frequent, and the patient may appear to have Parkinson's disease, with a shuffling gait, hypertonia, and tremors. In the final phase, progressive neurologic impairment ends in coma and death.

The most striking difference between the *gambiense* and *rhodesiense* forms of HAT is that the latter illness tends to follow a more acute course. Typically, in tourists with *T. b. rhodesiense* disease, systemic signs of infection, such as fever, malaise, and headache, appear before the end of the trip or shortly after the return home. Persistent tachycardia unrelated to fever is common early in the course of *T. b. rhodesiense* trypanosomiasis, and death may result from arrhythmias and congestive heart failure before CNS disease develops. In general, untreated *T. b. rhodesiense* trypanosomiasis leads to death in a matter of weeks to months, often without a clear distinction between the hemolymphatic and CNS stages. In contrast, *T. b. gambiense* disease can smolder for many months or even for years.

DIAGNOSIS

A definitive diagnosis of HAT requires detection of the parasite. If a chancre is present, fluid should be expressed and examined directly by light microscopy for the highly motile trypanosomes. The fluid also should be fixed and stained with Giemsa. Material obtained by needle aspiration of lymph nodes early in the illness should be examined similarly. Examination of wet preparations and Giemsa-stained thin and thick films of serial blood samples is also useful. If parasites are not seen initially in blood, efforts should be made to concentrate the organisms, which can be done in microhematocrit tubes containing acridine orange. Alternatively, the buffy coat from 10–15 mL of anticoagulated blood can be examined directly under a microscope. The likelihood of finding parasites in blood is higher in stage 1 than in stage 2 disease and in patients infected with *T. b. rhodesiense* rather than *T. b. gambiense*. Trypanosomes may also be seen in material aspirated from the bone marrow; the aspirate can be inoculated into liquid culture medium, as can blood, buffy coat, lymph node aspirates, and CSF. It is essential to examine CSF from all patients in whom HAT is suspected. Abnormalities in the CSF that may be associated with stage 2 disease include an increase in the CSF cell count as well as increases in opening pressure and in levels of total protein and IgM. Trypanosomes may be seen in the sediment of centrifuged CSF. Any CSF abnormality in a patient in whom trypanosomes have been found at other sites must be viewed as pathognomonic for CNS involvement and thus must prompt specific treatment for CNS disease. In patients

1398 with CSF pleocytosis in whom parasites are not found, tuberculous meningitis and HIV-associated CNS infections such as cryptococcosis should be considered in the differential diagnosis.

A number of serologic assays, such as the card agglutination test for trypanosomes (CATT) for *T. b. gambiense*, are available to aid in the diagnosis of HAT. Their ease of use makes them valuable for epidemiologic surveys, but their variable sensitivity and specificity mandate that decisions about treatment be based on demonstration of the parasite. Accurate PCR assays for detecting African trypanosomes in humans have been developed, but the lack of the necessary technical and human resources in most endemic areas stands in the way of their widespread use.

TREATMENT SLEEPING SICKNESS

The drugs used for treatment of HAT are suramin, pentamidine, eflornithine, and the organic arsenical melarsoprol. In the United States, these drugs can be obtained from the CDC. Therapy for HAT must be individualized on the basis of the infecting subspecies, the presence or absence of CNS disease, adverse reactions, and occasionally drug resistance. The choices of drugs for the treatment of HAT are summarized in [Table 252-2](#).

Suramin is highly effective against stage 1 *rhodesiense* HAT. However, it can cause serious adverse effects and must be administered under the close supervision of a physician. A 100- to 200-mg IV test dose should be given to detect hypersensitivity. The dosage for adults is 20 mg/kg on days 1, 5, 12, 18, and 26. The drug is given by slow IV infusion of a freshly prepared 10% aqueous solution. Approximately 1 patient in 20,000 has an immediate, severe, and potentially fatal reaction to the drug, developing nausea, vomiting, shock, and seizures. Less severe reactions include fever, photophobia, pruritus, arthralgias, and skin eruptions. Renal damage is the most common important adverse effect of suramin. Transient proteinuria often appears during treatment. A urinalysis should be done before each dose, and treatment should be discontinued if proteinuria increases or if casts and red cells appear in the sediment. Suramin should not be given to patients with renal insufficiency.

Pentamidine is the first-line drug for treatment of stage 1 *gambiense* HAT. The dose for both adults and children is 4 mg/kg per day, given IM or IV for 7–10 days. Frequent, immediate adverse reactions include nausea, vomiting, tachycardia, and hypotension. These reactions are usually transient and do not warrant cessation of therapy. Other adverse reactions include nephrotoxicity, abnormal liver function tests, neutropenia, rashes, hypoglycemia, and sterile abscesses. Suramin is an alternative agent for stage 1 *T. b. gambiense* disease.

Eflornithine is highly effective for treatment of both stages of *gambiense* sleeping sickness. In the trials on which the FDA based its approval, this agent cured >90% of 600 patients with stage 2 disease. The recommended treatment schedule is 400 mg/kg per day, given IV in four divided doses, for 2 weeks. Adverse reactions include diarrhea, anemia, thrombocytopenia, seizures, and hearing loss. The high dosage and duration of therapy required are disadvantages that make widespread use of eflornithine difficult. A randomized trial

comparing the standard eflornithine regimen (400 mg/kg per day infused over 6 h for 14 days) with nifurtimox-eflornithine combination therapy (NECT; oral nifurtimox, 15 mg/kg per day in three divided doses, plus IV eflornithine, 200 mg/kg per day in two divided doses, both for 7 days) in adults with stage 2 *gambiense* HAT showed improved efficacy and reduced adverse effects with combination therapy, making this drug suitable for first-line use.

The arsenical melarsoprol is the drug of choice for the treatment of *rhodesiense* HAT with CNS involvement and is an alternative agent for stage 2 *gambiense* disease. The “short course” of melarsoprol that is currently recommended has been shown to be noninferior to the decades-old treatment course for *T. b. rhodesiense*, which was administered over several weeks and was much more toxic. The short-course regimen consists of 10 daily doses of 2.2 mg/kg IV, each given with prednisolone (1 mg/kg).

Melarsoprol is highly toxic and should be administered with great care. As noted, all patients receiving melarsoprol should be given prednisolone to reduce the likelihood of drug-induced encephalopathy. Without prednisolone prophylaxis, the incidence of reactive encephalopathy has been as high as 18% in some series. Clinical manifestations of reactive encephalopathy include high fever, headache, tremor, impaired speech, seizures, and even coma and death. Treatment with melarsoprol should be discontinued at the first sign of encephalopathy but may be restarted cautiously at lower doses a few days after signs have resolved. Extravasation of the drug results in intense local reactions. Vomiting, abdominal pain, nephrotoxicity, and myocardial damage can occur.

PREVENTION

HAT poses complex public-health and epizootic problems in Africa. Considerable progress has been made in many areas through control programs that focus on eradication of vectors and drug treatment of infected humans. People can reduce their risk of acquiring trypanosomiasis by avoiding areas known to harbor infected insects, by wearing protective clothing, and by using insect repellent. Chemoprophylaxis is not recommended, and no vaccine is available to prevent transmission of the parasites.

253 Toxoplasma Infections

Kami Kim, Lloyd H. Kasper

DEFINITION

Toxoplasmosis is caused by infection with the obligate intracellular parasite *Toxoplasma gondii*. Acute infection acquired after birth may be asymptomatic but is thought to result in the lifelong chronic persistence of cysts in the host’s tissues. In both acute and chronic toxoplasmosis, the parasite is responsible for clinically evident disease, including lymphadenopathy, encephalitis, myocarditis, and pneumonitis. *Congenital* toxoplasmosis is an infection of newborns that results from the transplacental passage of parasites from an infected mother to the fetus. These infants may be asymptomatic at birth, but most later manifest a wide range of signs and symptoms, including chorioretinitis, strabismus, epilepsy, and psychomotor retardation. In immunocompetent individuals, toxoplasmosis can also present as acute disease (typically chorioretinitis) associated with food- or waterborne sources.

ETIOLOGY

T. gondii is an intracellular coccidian that infects both birds and mammals. There are two distinct stages in the life cycle of *T. gondii* that yield transmissible forms of the parasite ([Fig. 253-1](#)). In the *asexual* stages, tissue cysts that contain bradyzoites or sporulated oocysts that contain sporozoites are ingested by an intermediate host (e.g., a human, mouse, sheep, pig, or bird). The cyst is rapidly digested by the

TABLE 252-2 TREATMENT OF HUMAN AFRICAN TRYPANOSOMIASIS^a

Causative Organism	Clinical Stage	
	1 (Normal CSF)	2 (Abnormal CSF)
<i>T. b. gambiense</i> (West African)	Pentamidine Alternative: Suramin	Eflornithine Alternatives: NECT Melarsoprol ^b
<i>T. b. rhodesiense</i> (East African)	Suramin Alternative: Pentamidine	Melarsoprol ^b

^aFor doses and duration, see text. ^bShort course.

Abbreviations: CSF, cerebrospinal fluid; NECT, nifurtimox-eflornithine combination therapy.



T. gondii infects a wide range of mammals and birds. Its seroprevalence depends on the locale and the age of the population. Generally, hot arid climatic conditions are associated with a low prevalence of infection. In the United States and most European countries, the seroprevalence increases with age and exposure. For example, in the United States, 5–30% of individuals 10–19 years old and 10–67% of those >50 years old have serologic evidence of exposure. In Central America, France, Turkey, and Brazil, the seroprevalence is higher. Because of increased awareness of food-borne infections, the prevalence of seropositivity has decreased worldwide.

TRANSMISSION

Oral Transmission Most cases of human *Toxoplasma* infection are thought to be acquired by the oral route. Transmission can be attributable to ingestion of sporulated oocysts from contaminated soil, food, or water. During acute feline infection, a cat may excrete as many as 100 million parasites per day. These very stable sporozoite-containing oocysts are highly infectious and may remain viable for many years in soil or water. Humans infected during an oocyst-transmitted infection develop stage-specific antibodies to the oocyst/sporozoite.

Children and adults also can acquire infection from tissue cysts containing bradyzoites. The ingestion of a single cyst is all that is required for human infection. Undercooking or insufficient freezing of meat is an important source of infection in the developed world. In the United States, lamb products and pork products may yield evidence of cysts that contain bradyzoites, but the overall prevalence of *T. gondii* has been gradually decreasing. The incidence in beef is much lower—perhaps as low as 1%. Direct ingestion of bradyzoite cysts in these various meat products leads to acute infection.

Transmission via Blood or Organs In addition to being transmitted orally, *T. gondii* can be transmitted directly from a seropositive donor to a seronegative recipient in a transplanted heart, heart-lung, kidney, liver, or pancreas. Viable parasites can be cultured from refrigerated anticoagulated blood, which may be a source of infection in individuals receiving blood transfusions. *T. gondii* reactivation has been reported in bone marrow, hematopoietic stem cell, and liver transplant recipients as well as in individuals with AIDS. Although antibody titers generally are not useful in monitoring *T. gondii* infection, individuals with higher antibody titers may be at relatively high risk for reactivation after hematopoietic stem cell transplantation; thus routine polymerase chain reaction (PCR) screening of blood from these patients may be in order. Finally, laboratory personnel can be infected after contact with contaminated needles or glassware or with infected tissue.

Transplacental Transmission On average, about one-third of all women who acquire infection with *T. gondii* during pregnancy transmit the parasite to the fetus; the remainder give birth to normal, uninfected babies. Of the various factors that influence fetal outcome, gestational age at the time of infection is the most critical (see below). Few data support a role for recrudescence of maternal infection as the source of congenital disease, although rare cases of transmission by immunocompromised women (e.g., those infected with HIV or those receiving high-dose glucocorticoids) have been reported. Thus, women who are seropositive before pregnancy usually are protected against acute infection and do not give birth to congenitally infected neonates.

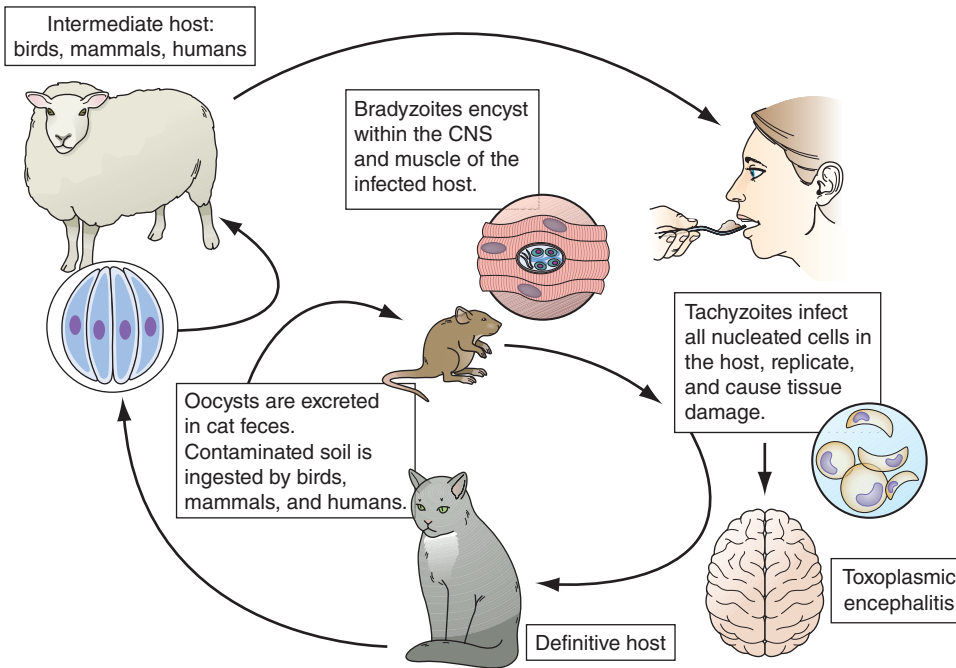


FIGURE 253-1 Life cycle of *Toxoplasma gondii*. The cat is the definitive host in which the sexual phase of the cycle is completed. Oocysts shed in cat feces can infect a wide range of animals, including birds, rodents, grazing domestic animals, and humans. The bradyzoites found in the muscle of food animals may infect humans who eat insufficiently cooked meat products, particularly lamb and pork. Although human disease can take many forms, congenital infection and encephalitis from reactivation of latent infection in the brains of immunosuppressed persons are the most important manifestations. CNS, central nervous system. (Courtesy of Dominique Buzoni-Gatel, Institut Pasteur, Paris; with permission.)

acidic-pH gastric secretions. Bradyzoites or sporozoites are released, enter the small-intestinal epithelium, and transform into rapidly dividing tachyzoites. The tachyzoites can infect and replicate in all mammalian cells except red blood cells. The parasite actively penetrates the cell and forms a parasitophorous vacuole. Parasite replication continues within the vacuole. After the parasites reach a critical mass, intracellular signaling within the host and the parasite, including calcium fluxes, result in parasite egress from the vacuole. The host cell is destroyed, and the released tachyzoites infect adjoining cells. The tachyzoite replication cycle within an infected organ causes cytopathology. Most tachyzoites are eliminated by the host's humoral and cell-mediated immune responses. Tissue cysts containing many bradyzoites develop 7–10 days after systemic tachyzoite infection. These tissue cysts occur in various host organs but persist principally within the central nervous system (CNS) and muscle. The development of this chronic stage completes the asexual portion of the life cycle. Active infection in the immunocompromised host is most likely to be due to the spontaneous release of encysted parasites that undergo rapid transformation into tachyzoites within the CNS and are not contained by the immune system.

The *sexual* stage in the life cycle takes place in the cat (the definitive host). The parasite's sexual phase is defined by the formation of oocysts within the feline host. This enteroepithelial cycle begins with the ingestion of the bradyzoite tissue cysts and, after several intermediate stages, culminates in the production of gametes. Gamete fusion produces a zygote, which envelops itself in a rigid wall and is secreted in the feces as an unsporulated oocyst. After 2–3 days of exposure to air at ambient temperature, the noninfectious oocyst sporulates to produce eight sporozoite progeny. The sporulated oocyst can be ingested by an intermediate host, such as a person emptying a cat's litter box or a pig rummaging in a barnyard. It is in the intermediate host that *T. gondii* completes its life cycle.



Sporulated oocysts are environmentally hardy and very infectious; they are thought to be sources of waterborne outbreaks such as those reported in Victoria (British Columbia, Canada) and in South America.

The following general guidelines can be used to evaluate congenital infection. There is essentially no risk if the mother becomes infected ≥ 6 months before conception. If infection is acquired < 6 months before conception, the likelihood of transplacental infection increases as the interval between infection and conception decreases. Women with documented acute toxoplasmosis should be counseled to use appropriate measures to prevent pregnancy for 6 months after infection. In pregnancy, if the mother becomes infected during the first trimester, the incidence of transplacental infection is lowest ($\sim 15\%$), but the disease in the neonate is most severe. If maternal infection occurs during the third trimester, the incidence of transplacental infection is greatest (65%), but the infant is usually asymptomatic at birth. Infected infants who are normal at birth may have a higher incidence of learning disabilities and chronic neurologic sequelae than uninfected children. Only a small proportion (20%) of women infected with *T. gondii* develop clinical signs of infection. Often the diagnosis is first appreciated when routine postconception serologic tests show evidence of specific antibody.

PATHOGENESIS

Upon the host's ingestion of either tissue cysts containing bradyzoites or oocysts containing sporozoites, the parasites are released from the cysts by the digestive process. Bradyzoites are resistant to the effect of pepsin and invade the host's gastrointestinal tract. Within enterocytes (or other gut-associated cells), the parasites undergo morphologic transformation, giving rise to invasive tachyzoites. These tachyzoites induce a parasite-specific secretory IgA response. From the gastrointestinal tract, parasites disseminate to a variety of organs, particularly lymphatic tissue, skeletal muscle, myocardium, retina, placenta, and the CNS. At these sites, the parasite infects host cells, replicates, and invades the adjoining cells. In this fashion, the hallmarks of the infection develop: cell death and focal necrosis surrounded by an acute inflammatory response.

In the immunocompetent host, both the humoral and the cellular immune responses control infection; parasite virulence and tissue tropism may be strain specific. Tachyzoites are sequestered by a variety of immune mechanisms, including induction of parasitocidal antibody, activation of macrophages with radical intermediates, production of interferon γ (IFN- γ), and stimulation of CD8+ cytotoxic T lymphocytes. These antigen-specific lymphocytes are capable of killing both extracellular parasites and target cells infected with parasites. As tachyzoites are cleared from the acutely infected host, tissue cysts containing bradyzoites begin to appear, usually within the CNS and the retina. Studies indicate that *Toxoplasma* secretes signaling molecules into infected host cells and that these molecules modulate host gene expression, host metabolism, and host immune response. While it was initially thought that cysts with bradyzoites are not eliminated by the immune system, recent studies in the murine model indicate that both CD8+ T cells and alternatively activated macrophages are able to kill cysts in vivo; some cysts persist, however, and the ability to eliminate cysts may depend on the genetic background of the infected host.

In the immunocompromised or fetal host, the immune factors necessary to control the spread of tachyzoite infection are lacking. This altered immune state allows the persistence of tachyzoites and gives rise to progressive focal destruction that results in organ failure (i.e., necrotizing encephalitis, pneumonia, and myocarditis).

It is thought that all infected individuals have persistent infection with cysts containing bradyzoites, but this lifelong infection usually remains subclinical. Although bradyzoites are in a slow metabolic phase, cysts do degenerate and rupture within the CNS. This degenerative process, with the development of new bradyzoite-containing cysts, is the most probable source of recrudescence in immunocompromised individuals and the most likely stimulus for the persistence of antibody titers in the immunocompetent host. Although the concept is controversial, the persistence of toxoplasmosis has been hypothesized to be a contributing factor to a variety of neuropsychiatric conditions, including schizophrenia and bipolar disease. In rodents, infection clearly has significant effects on behavior, increasing predation.

PATHOLOGY

Cell death and focal necrosis due to replicating tachyzoites induce an intense mononuclear inflammatory response in any tissue or cell type infected. Tachyzoites rarely can be visualized by routine histopathologic staining of these inflammatory lesions. However, immunofluorescent staining with parasitic antigen-specific antibodies can reveal either the organism itself or evidence of antigen. In contrast to this inflammatory process caused by tachyzoites, bradyzoite-containing cysts cause inflammation only at the early stages of development, and even this inflammation may be a response to the presence of tachyzoite antigens. Once the cysts reach maturity, the inflammatory process can no longer be detected, and the cysts remain immunologically quiescent within the brain matrix until they rupture.

Lymph Nodes During acute infection, lymph node biopsy demonstrates characteristic findings, including follicular hyperplasia and irregular clusters of tissue macrophages with eosinophilic cytoplasm. Granulomas rarely are evident in these specimens. Although tachyzoites are not usually visible, they can be sought either by subinoculation of infected tissue into mice, with resultant disease, or by PCR. PCR amplification of DNA fragments of *Toxoplasma* genes is effective and sensitive in establishing lymph node infection by tachyzoites.

Eyes In the eye, infiltrates of monocytes, lymphocytes, and plasma cells may produce uni- or multifocal lesions. Granulomatous lesions and chorioretinitis can be observed in the posterior chamber after acute necrotizing retinitis. Other ocular complications include iridocyclitis, cataracts, and glaucoma.

Central Nervous System During CNS involvement, both focal and diffuse meningoencephalitis can be documented, with evidence of necrosis and microglial nodules. Necrotizing encephalitis in patients without AIDS is characterized by small diffuse lesions with perivascular cuffing in contiguous areas. In the AIDS population, polymorphonuclear leukocytes may be present in addition to monocytes, lymphocytes, and plasma cells. Cysts containing bradyzoites frequently are found contiguous with the necrotic tissue border. As a consequence of combined antiretroviral therapy (cART) for AIDS, the incidence of toxoplasmosis has decreased in the developed world. Its incidence in under-resourced settings is not known.

Lungs and Heart Among patients with AIDS who die of toxoplasmosis, 40–70% have involvement of the lungs and heart. Interstitial pneumonitis can develop in neonates and immunocompromised patients. Thickened and edematous alveolar septa infiltrated with mononuclear and plasma cells are apparent. This inflammation may extend to the endothelial walls. Tachyzoites and bradyzoite-containing cysts have been observed within the alveolar membrane. Superimposed bronchopneumonia can be caused by other microbial agents. Cysts and aggregates of parasites in cardiac muscle tissue are evident in patients with AIDS who die of toxoplasmosis. Focal necrosis surrounded by inflammatory cells is associated with hyaline necrosis and disrupted myocardial cells. Pericarditis is associated with toxoplasmosis in some patients.

Gastrointestinal Tract Rare cases of human gastrointestinal tract infection with *T. gondii* have presented as ulcerations in the mucosa. Acute infection in certain strains of inbred mice (C57BL/6) results in lethal ileitis within 7–9 days. This inflammatory bowel disease has been recognized in several other mammalian species, including pigs and nonhuman primates. Although the association between human inflammatory bowel disease and either acute or recurrent *Toxoplasma* infection has not been established, studies have demonstrated recognition of the infection by human intestinal epithelial cells, as evidenced by mitogen-activated protein kinase phosphorylation, nuclear factor κ B translocation, and interleukin (IL) 8 secretion.

Other Sites Pathologic changes during disseminated infection are similar to those described for the lymph nodes, eyes, and CNS. In patients with AIDS, the skeletal muscle, pancreas, stomach, and kidneys can be involved, with necrosis, invasion by inflammatory cells, and (rarely) tachyzoites detectable by routine staining. Large necrotic

lesions may cause direct tissue destruction. In addition, secondary effects from acute infection of these various organs, including pancreatitis, myositis, and glomerulonephritis, have been reported.

HOST IMMUNE RESPONSE

Acute *Toxoplasma* infection evokes a cascade of protective immune responses in the immunocompetent host. *Toxoplasma* enters the host at the gut mucosal level and evokes a mucosal immune response that includes the production of antigen-specific secretory IgA. Titers of serum IgA antibody directed at p30 (SAG1) are a useful marker for congenital and acute toxoplasmosis. Milk-whey IgA from acutely infected mothers contains a high titer of antibody to *T. gondii* and can block infection of enterocytes in vitro. In mice, IgA intestinal secretions directed at the parasite are abundant and are associated with the induction of mucosal T cells.

Within the host, *T. gondii* rapidly induces detectable levels of both IgM and IgG serum antibodies. Monoclonal gammopathy of the IgG class can occur in congenitally infected infants. IgM levels may be increased in newborns with congenital infection. The polyclonal IgG antibodies evoked by infection are parasitocidal in vitro in the presence of serum complement and are the basis for the Sabin-Feldman dye test. However, cell-mediated immunity is the major protective response evoked by the parasite during host infection. Macrophages are activated after phagocytosis of antibody-opsonized parasites. This activation can lead to death of the parasite by either an oxygen-dependent or an oxygen-independent process. If the parasite is not phagocytosed and enters the macrophage by active penetration, it continues to replicate, and this replication may represent the mechanism for transport and dissemination to distant organs. *Toxoplasma* stimulates a robust IL-12 response by human dendritic cells. The requirement for costimulation via CD40/154 has been established. The CD4+ and CD8+ T cell responses are antigen-specific and further stimulate the production of a variety of important lymphokines that expand the T cell and natural killer cell repertoire. *T. gondii* is a potent inducer of a T_H1 phenotype, with IL-12 and IFN- γ playing an essential role in the control of the parasites' growth in the host. Regulation of the inflammatory response is at least partially under the control of a T_H2 response that includes the production of IL-4 and IL-10 in seropositive individuals. Both asymptomatic patients and those with active infection may have a depressed CD4+ to-CD8+ ratio. This shift may be correlated with a disease syndrome but is not necessarily correlated with disease outcome. Human T cell clones of both the CD4+ and the CD8+ phenotypes are cytolytic against parasite-infected macrophages. These T cell clones produce cytokines that are "microbistatic." IL-18, IL-7, and IL-15 upregulate the production of IFN- γ and may be important during acute and chronic infection. The effect of IFN- γ may be paradoxical, with stimulation of a host down-regulatory response as well.

Although *T. gondii* infection is believed to be recrudescence in patients with AIDS or other immunocompromised states, antibody titers are not useful in establishing reactivation or in following the activity of infection. An absence of positive serologies suggests an alternative diagnosis, although AIDS patients may have borderline positive or low serologies. T cells from AIDS patients with reactivation of toxoplasmosis fail to secrete both IFN- γ and IL-2. This alteration in the production of these critical immune cytokines contributes to the persistence of infection. *Toxoplasma* infection frequently develops late in the course of AIDS, when the loss of T cell-dependent protective mechanisms, particularly CD8+ T cells, becomes most pronounced.

CLINICAL MANIFESTATIONS

In persons whose immune systems are intact, acute toxoplasmosis is usually asymptomatic and self-limited. This condition can go unrecognized in 80–90% of adults and children with acquired infection. The asymptomatic nature of this infection makes diagnosis difficult in mothers infected during pregnancy. In contrast, the wide range of clinical manifestations in congenitally infected children includes severe neurologic complications such as hydrocephalus, microcephaly, mental retardation, and chorioretinitis. If prenatal infection is severe, multiorgan failure and subsequent intrauterine fetal death can occur.

In children and adults, chronic infection can persist throughout life, with little consequence to the immunocompetent host.

Toxoplasmosis in Immunocompetent Patients The most common manifestation of acute toxoplasmosis is cervical lymphadenopathy. The nodes may be single or multiple, are usually nontender, are discrete, and vary in firmness. Lymphadenopathy also may be found in suboccipital, supraclavicular, inguinal, and mediastinal areas. Generalized lymphadenopathy occurs in 20–30% of symptomatic patients. Between 20% and 40% of patients with lymphadenopathy also have headache, malaise, fatigue, and fever (usually with a temperature of <40°C [$<104^{\circ}\text{F}$]). A smaller proportion of symptomatic individuals have myalgia, sore throat, abdominal pain, maculopapular rash, meningoencephalitis, and confusion. Rare complications associated with infection in the normal immune host include pneumonia, myocarditis, encephalopathy, pericarditis, and polymyositis. Signs and symptoms associated with acute infection usually resolve within several weeks, although the lymphadenopathy may persist for some months. In one epidemic, toxoplasmosis was diagnosed correctly in only 3 of the 25 patients who consulted physicians. If toxoplasmosis is considered in the differential diagnosis, routine laboratory and serologic screening should precede node biopsy.



It is now appreciated that genotypes of *T. gondii* prevalent in South America may be more virulent than those typically seen in North America or Europe. These genotypes may be associated with acute or recurrent ocular disease in immunocompetent individuals and have also been associated with pneumonitis and a fulminant sepsis picture in immunologically normal individuals. Thus a detailed history is critical for establishing a diagnosis.

The results of routine laboratory studies are usually unremarkable except for minimal lymphocytosis, an elevated erythrocyte sedimentation rate, and a nominal increase in serum aminotransferase levels. Evaluation of cerebrospinal fluid (CSF) in cases with evidence of encephalopathy or meningoencephalitis shows an elevation of intracranial pressure, mononuclear pleocytosis (10–50 cells/mL), a slight increase in protein concentration, and (occasionally) an increase in the gamma globulin level. PCR amplification of the *Toxoplasma* DNA target sequence in CSF may be beneficial. The CSF of chronically infected individuals is normal.

Infection of Immunocompromised Patients Patients with AIDS and those receiving immunosuppressive therapy for lymphoproliferative disorders are at greatest risk for developing acute toxoplasmosis. Toxoplasmosis has also been reported after treatment with antibodies to tumor necrosis factor. The infection may be due either to reactivation of latent infection or to acquisition of parasites from exogenous sources such as blood or transplanted organs. In individuals with AIDS, >95% of cases of *Toxoplasma* encephalitis (TE) are believed to be due to recrudescence infection. In most of these cases, encephalitis develops when the CD4+ T cell count falls below 100/ μL . In immunocompromised hosts, the disease may be rapidly fatal if untreated. Thus, accurate diagnosis and initiation of appropriate therapy are necessary to prevent fulminant infection.

Toxoplasmosis is a principal opportunistic infection of the CNS in persons with AIDS. Although geographic origin may be related to frequency of infection, it has no correlation with the severity of disease in immunocompromised hosts. Individuals with AIDS who are seropositive for *T. gondii* are at high risk for encephalitis. Before the advent of current cART, about one-third of the 15–40% of adult AIDS patients in the United States who were latently infected with *T. gondii* developed TE. TE may still be a presenting infection in individuals who are unaware of their positive HIV status.

The signs and symptoms of acute toxoplasmosis in immunocompromised patients principally involve the CNS (Fig. 253-2). More than 50% of patients with clinical manifestations have intracerebral involvement. Clinical findings at presentation range from nonfocal to focal dysfunction. CNS findings include encephalopathy, meningoencephalitis, and mass lesions. Patients may present with altered mental status (75%), fever (10–72%), seizures (33%), headaches (56%), and focal neurologic findings (60%), including motor deficits, cranial nerve

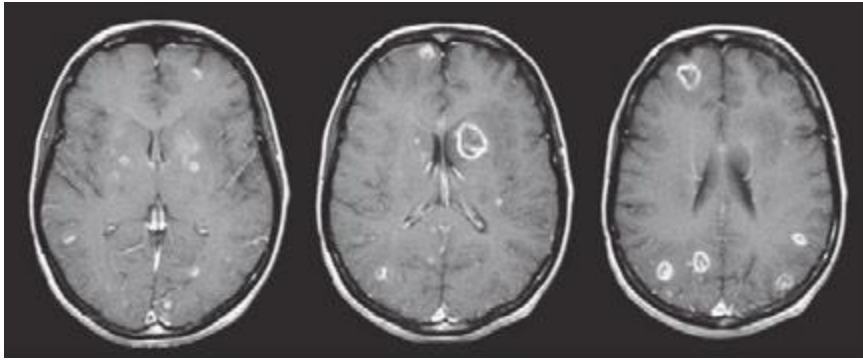


FIGURE 253-2 Toxoplasmic encephalitis in a 36-year-old patient with AIDS. The multiple lesions are demonstrated by MRI scanning (T1-weighted with gadolinium enhancement). (Courtesy of Clifford Eskey, Dartmouth Hitchcock Medical Center, Hanover, NH; with permission.)

palsies, movement disorders, dysmetria, visual-field loss, and aphasia. Patients who present with evidence of diffuse cortical dysfunction develop evidence of focal neurologic disease as infection progresses. This altered condition is due not only to the necrotizing encephalitis caused by direct invasion by the parasite but also to secondary effects, including vasculitis, edema, and hemorrhage. The onset of infection can range from an insidious process over several weeks to an acute presentation with fulminant focal deficits, including hemiparesis, hemiplegia, visual-field defects, localized headache, and focal seizures.

Although lesions can occur anywhere in the CNS, the areas most often involved appear to be the brainstem, basal ganglia, pituitary gland, and corticomedullary junction. Brainstem involvement gives rise to a variety of neurologic dysfunctions, including cranial nerve palsy, dysmetria, and ataxia. With basal ganglionic infection, patients may develop hydrocephalus, choreiform movements, and choreoathetosis. *Toxoplasma* usually causes encephalitis, and meningeal involvement is uncommon. CSF findings may be unremarkable or may include a modest increase in cell count and in protein—but not glucose—concentration. Nonetheless, the parasite may be detected by PCR in CSF from many patients with TE.

Cerebral toxoplasmosis must be differentiated from other opportunistic infections or tumors in the CNS of AIDS patients. The differential diagnosis includes herpes simplex encephalitis, cryptococcal meningitis, progressive multifocal leukoencephalopathy, and primary CNS lymphoma. Involvement of the pituitary gland can give rise to panhypopituitarism and hyponatremia from inappropriate secretion of vasopressin (antidiuretic hormone). HIV-associated neurocognitive disorder (HAND) may present as cognitive impairment, attention loss, and altered memory. Brain biopsy in patients who have been treated for TE but who continue to exhibit neurologic dysfunction often fails to identify organisms.

Autopsies of *Toxoplasma*-infected patients have demonstrated the involvement of multiple organs, including the lungs, gastrointestinal tract, pancreas, skin, eyes, heart, and liver. *Toxoplasma* pneumonia can be confused with *Pneumocystis* pneumonia (PcP). Respiratory involvement usually presents as dyspnea, fever, and a nonproductive cough and may rapidly progress to acute respiratory failure with hemoptysis, metabolic acidosis, hypotension, and (occasionally) disseminated intravascular coagulation. Histopathologic studies demonstrate necrosis and a mixed cellular infiltrate. The presence of organisms is a helpful diagnostic indicator, but organisms can also be found in healthy tissue. Infection of the heart is usually asymptomatic but can be associated with cardiac tamponade or biventricular failure. Infections of the gastrointestinal tract and the liver have been documented.

Congenital Toxoplasmosis Between 400 and 4000 infants born each year in the United States are affected by congenital toxoplasmosis. Acute infection in mothers acquiring *T. gondii* during pregnancy is usually asymptomatic; most such women are diagnosed via prenatal serologic screening. Infection of the placenta leads to

hematogenous infection of the fetus. As gestation proceeds, the proportion of fetuses that become infected increases, but the clinical severity of the infection declines. Although infected children may initially be asymptomatic, the persistence of *T. gondii* can result in reactivation and clinical disease—most frequently chorioretinitis—decades later. Factors associated with relatively severe disabilities include delays in diagnosis and in initiation of therapy, neonatal hypoxia and hypoglycemia, profound visual impairment (see “Ocular Infection,” below), uncorrected hydrocephalus, and increased intracranial pressure. If treated appropriately, upwards of 70% of children have normal developmental, neurologic, and ophthalmologic findings at follow-up evaluations. Treatment for 1 year with pyrimethamine, a sulfonamide, and folinic acid is tolerated with minimal toxicity (see “Treatment,” below).



Ocular Infection Infection with *T. gondii* is estimated to cause 35% of all cases of chorioretinitis in the United States and Europe. It was formerly thought that the majority of cases of ocular disease were due to congenital infection. New ocular toxoplasmosis in immunocompetent individuals occurs more commonly than was previously appreciated and has been associated with outbreaks in Victoria (British Columbia) and in South America. A variety of ocular manifestations are documented, including blurred vision, scotoma, photophobia, and eye pain. Macular involvement occurs, with loss of central vision, and nystagmus is secondary to poor fixation. Involvement of the extraocular muscles may lead to disorders of convergence and to strabismus. Ophthalmologic examination should be undertaken in newborns with suspected congenital infection. As the inflammation resolves, vision improves, but episodic flare-ups of chorioretinitis, which progressively destroy retinal tissue and lead to glaucoma, are common. The ophthalmologic examination reveals yellow-white, cotton-like patches with indistinct margins of hyperemia. As the lesions age, white plaques with distinct borders and black spots within the retinal pigment become more apparent. Lesions usually are located near the posterior pole of the retina; they may be single but are more commonly multiple. Congenital lesions may be unilateral or bilateral and show evidence of massive chorioretinal degeneration with extensive fibrosis. Surrounding these areas of involvement are a normal retina and vasculature. In patients with AIDS, retinal lesions are often large, with diffuse retinal necrosis, and include both free tachyzoites and cysts containing bradyzoites. Toxoplasmic chorioretinitis may be a prodrome to the development of encephalitis.

DIAGNOSIS

Tissue and Body Fluids The differential diagnosis of acute toxoplasmosis can be made by appropriate culture, serologic testing, and PCR (Table 253-1). Although available only at specialized laboratories, the isolation of *T. gondii* from blood or other body fluids can be accomplished after subinoculation of the sample into the peritoneal cavity of mice. If no parasites are found in the mouse’s peritoneal fluid 6–10 days after inoculation, its anti-*Toxoplasma* serum titer can be evaluated 4–6 weeks after inoculation. Isolation of *T. gondii* from the patient’s body fluids reflects acute infection, whereas isolation from biopsied tissue is an indication only of the presence of tissue cysts and should not be misinterpreted as evidence of acute toxoplasmosis. Persistent parasitemia in patients with latent, asymptomatic infection is rare. Histologic examination of lymph nodes may suggest the characteristic changes described above. Demonstration of tachyzoites in lymph nodes establishes the diagnosis of acute toxoplasmosis. Like subinoculation into mice, histologic demonstration of cysts containing bradyzoites confirms prior infection with *T. gondii* but is nondiagnostic for acute infection.

Serology The procedures mentioned above have great diagnostic value but are limited by difficulties encountered either in the growth

TABLE 253-1 DIFFERENTIAL LABORATORY DIAGNOSIS OF TOXOPLASMOSIS

Clinical Setting	Alternative Diagnosis	Distinguishing Characteristics
Mononucleosis syndrome	Epstein-Barr virus infection	Serology
	Cytomegalovirus infection	Serology/PCR or culture
	HIV infection	Serology/viral load
	<i>Bartonella</i> infection (cat-scratch disease)	Biopsy (PCR or culture)/serology
	Lymphoma	Biopsy
Congenital infection	Cytomegalovirus infection	Viral culture/PCR
	Herpes simplex virus infection	Viral culture/PCR
	Rubella virus infection	Viral culture/serology
	Syphilis	Serology
	Listeriosis	Bacterial culture
Chorioretinitis in immunocompetent individual	Tuberculosis	Bacterial culture
	Syphilis	Serology
	Histoplasmosis	Serology/culture
Chorioretinitis in AIDS patient	Cytomegalovirus infection	Viral culture/PCR
	Syphilis	Serology
	Herpes simplex virus infection	Viral culture/PCR
	Varicella-zoster virus infection	Viral culture/PCR
	Fungal infection	Culture
CNS lesions in AIDS patient	Lymphoma or metastatic tumor	Tissue biopsy
	Brain abscess	Biopsy and culture
	Progressive multifocal leukoencephalopathy	PCR for JC virus
	Fungal infection	Biopsy and culture
	Mycobacterial infection	Biopsy and culture

Abbreviations: CNS, central nervous system; PCR, polymerase chain reaction.

Source: Adapted from JD Schwartzman: *Toxoplasmosis, in Principles and Practice of Clinical Parasitology*. Hoboken, Wiley, 2001.

of parasites *in vivo* or in the identification of tachyzoites by histochemical methods. Serologic testing has become the routine method of diagnosis.

Diagnosis of acute infection with *T. gondii* can be established by detection of the simultaneous presence of IgG and IgM antibodies to *Toxoplasma* in serum. The presence of circulating IgA favors the diagnosis of an acute infection. The Sabin-Feldman dye test, the indirect fluorescent antibody test, and the enzyme-linked immunosorbent assay (ELISA) all satisfactorily measure circulating IgG antibody to *Toxoplasma*. Positive IgG titers (>1:10) can be detected as early as 2–3 weeks after infection. These titers usually peak at 6–8 weeks and decline slowly to a new baseline level that persists for life. Antibody avidity increases with time and can be useful in difficult cases during pregnancy for establishing when infection may have occurred. The serum IgM titer should be measured in concert with the IgG titer to better establish the time of infection; either the double-sandwich IgM-ELISA or the IgM-immunosorbent assay (IgM-ISAGA) should be used. Both assays are specific and sensitive, with fewer false-positive results than other commercial tests. The double-sandwich IgA-ELISA is more sensitive than the IgM-ELISA for detecting congenital infection in the fetus and newborn. Although a negative IgM result with a positive IgG titer indicates distant infection, IgM can persist for >1 year and should not necessarily be considered a reflection of acute disease. If acute toxoplasmosis is suspected, a more extensive panel of serologic tests can be performed. In the United States, testing is

available at the *Toxoplasma* Serology Laboratory at Palo Alto Medical Foundation (<http://www.pamf.org/serology/clinicianguide.html>).

Molecular Diagnostics Molecular approaches can directly detect *T. gondii* in biologic samples independent of the serologic response. Results obtained with PCR have suggested high sensitivity, specificity, and clinical utility in the diagnosis of TE, and PCR technology may be becoming more readily available in resource-poor settings. Real-time PCR is a promising technique that can provide quantitative results. Isolates can be genotyped and polymorphic sequences can be obtained, with consequent identification of the precise strain. Molecular epidemiologic studies with polymorphic markers have been useful in correlating clinical signs and symptoms of disease with different *T. gondii* genotypes.

The Immunocompetent Adult or Child For the patient who presents with lymphadenopathy only, a positive IgM titer is an indication of acute infection—and an indication for therapy, if clinically warranted (see “Treatment,” below). The serum IgM titer should be determined again in 3 weeks. An elevation in the IgG titer without an increase in the IgM titer suggests that infection is present but is not acute. If there is a borderline increase in either IgG or IgM, the titers should be reassessed in 3–4 weeks.

The Immunocompromised Host A presumptive clinical diagnosis of TE in patients with AIDS is based on clinical presentation, history of exposure (as evidenced by positive serology), and radiologic evaluation. To detect latent infection with *T. gondii*, HIV-infected persons should be tested for IgG antibody to *Toxoplasma* soon after HIV infection is diagnosed. When these criteria are used, the predictive value is as high as 80%. More than 97% of patients with AIDS and toxoplasmosis have IgG antibody to *T. gondii* in serum. IgM serum antibody usually is not detectable. Although IgG titers do not correlate with active infection, serologic evidence of infection virtually always precedes the development of TE. It is therefore important to determine the *Toxoplasma* antibody status of all patients infected with HIV. Antibody titers may range from negative to 1:1024 in patients with AIDS and TE. Fewer than 3% of patients have no demonstrable antibody to *Toxoplasma* at diagnosis of TE.

Patients with TE have focal or multifocal abnormalities demonstrable by CT or MRI. Neuroradiologic evaluation should include double-dose contrast CT of the head. By this test, single and frequently multiple contrast-enhancing lesions (<2 cm) may be identified. MRI usually demonstrates multiple lesions located in both hemispheres, with the basal ganglia and corticomedullary junction most commonly involved; MRI provides a more sensitive evaluation of the efficacy of therapy than does CT (Fig. 253-2). These findings are not pathognomonic of *Toxoplasma* infection, because 40% of CNS lymphomas are multifocal and 50% are ring-enhancing. For both MRI and CT scans, the rate of false-negative results is ~10%. The finding of a single lesion on an MRI scan increases the likelihood of primary CNS lymphoma (in which solitary lesions are four times more likely than in TE) and strengthens the argument for the performance of a brain biopsy. A therapeutic trial of anti-*Toxoplasma* medications is frequently used to assess the diagnosis. Treatment of presumptive TE with pyrimethamine plus sulfadiazine or clindamycin results in quantifiable clinical improvement in >50% of patients by day 3. Leucovorin is administered to prevent bone marrow toxicity. By day 7, >90% of treated patients show evidence of improvement. In contrast, if patients fail to respond or have lymphoma, clinical signs and symptoms worsen by day 7. Patients in this category require brain biopsy with or without a change in therapy. This procedure can now be performed by a stereotactic CT-guided method that reduces the potential for complications. Brain biopsy for *T. gondii* identifies organisms in 50–75% of cases. PCR amplification of CSF may also confirm toxoplasmosis or suggest alternative diagnoses (Table 253-1), such as progressive multifocal leukoencephalopathy (JC virus positive) or primary CNS lymphoma (Epstein-Barr virus positive).

CT and MRI with contrast are currently the standard diagnostic imaging tests for TE. As in other conditions, the radiologic response may lag behind the clinical response. Resolution of lesions may take

1404 from 3 weeks to 6 months. Some patients show clinical improvement despite worsening radiographic findings.

Congenital Infection The issue of concern when a pregnant woman has evidence of recent *T. gondii* infection is whether the fetus is infected. PCR analysis of the amniotic fluid for the B1 gene of *T. gondii* has replaced fetal blood sampling. Serologic diagnosis is based on the persistence of IgG antibody or a positive IgM titer after the first week of life (a time frame that excludes placental leak). The IgG determination should be repeated every 2 months. An increase in IgM beyond the first week of life is indicative of acute infection. Up to 25% of infected newborns may be seronegative and have normal routine physical examinations. Thus assessment of the eye and the brain, with ophthalmologic testing, CSF evaluation, and radiologic studies, is important in establishing the diagnosis.

Ocular Toxoplasmosis The serum antibody titer may not correlate with the presence of active lesions in the fundus, particularly in cases of congenital toxoplasmosis. In general, a positive IgG titer (measured in undiluted serum if necessary) in conjunction with typical lesions establishes the diagnosis. Antibody production in ocular fluids, expressed in terms of the Goldmann-Witmer coefficient, has been described for diagnosis of ocular disease but does not always correlate with PCR results. If lesions are atypical and the serum antibody titer is in the low-positive range, the diagnosis is presumptive. The parasitic antigen-specific polyclonal IgG assay as well as parasite-specific PCR may facilitate the diagnosis. Accordingly, the clinical diagnosis of ocular toxoplasmosis can be supported in 60–90% of cases by laboratory tests, depending on the time of anterior chamber puncture and the panel of antibody analyses used. In the remaining cases, the possibility of a falsely negative laboratory diagnosis or of an incorrect clinical diagnosis cannot be clarified further.

TREATMENT TOXOPLASMOSIS

CONGENITAL INFECTION

Congenitally infected neonates are treated with daily oral pyrimethamine (1 mg/kg) and sulfadiazine (100 mg/kg) with folinic acid for 1 year. Depending on the signs and symptoms, prednisone (1 mg/kg per day) may be used for congenital infection. Some U.S. states and some countries routinely screen pregnant women (France, Austria) and/or newborns (Denmark, Massachusetts). Management and treatment regimens vary with the country and the treatment center. Most experts use spiramycin to treat pregnant women who have acute toxoplasmosis early in pregnancy and use pyrimethamine/sulfadiazine/folinic acid to treat women who seroconvert after 18 weeks of pregnancy or in cases of documented fetal infection. This treatment is somewhat controversial: clinical studies, which have included few untreated women, have not proven the efficacy of such therapy in preventing congenital toxoplasmosis. However, studies do suggest that treatment during pregnancy decreases the severity of infection. Many women who are infected in the first trimester elect termination of pregnancy. Those who do not terminate pregnancy are offered prenatal antibiotic therapy to reduce the frequency and severity of *Toxoplasma* infection in the infant. The optimal duration of treatment for a child with asymptomatic congenital toxoplasmosis is not clear, although most clinicians in the United States would treat the child for 1 year in light of cohort investigations conducted by the National Collaborative Chicago-Based, Congenital Toxoplasmosis Study.

INFECTION IN IMMUNOCOMPETENT PATIENTS

Immunologically competent adults and older children who have only lymphadenopathy do not require specific therapy unless they have persistent, severe symptoms. Patients with ocular toxoplasmosis are usually treated for 1 month with pyrimethamine plus either sulfadiazine or clindamycin and sometimes with prednisone. Treatment should be supervised by an ophthalmologist familiar with *Toxoplasma* disease. Ocular disease can be self-limited without treatment, but therapy is typically considered for lesions that are severe or close to the fovea or optic disc.

INFECTION IN IMMUNOCOMPROMISED PATIENTS

Primary Prophylaxis Patients with AIDS should be treated for acute toxoplasmosis; in immunocompromised patients, toxoplasmosis is rapidly fatal if untreated. Before the introduction of cART, the median survival time was >1 year for patients who could tolerate treatment for TE. Despite their toxicity, the drugs used to treat TE were required for survival prior to cART. The incidence of TE has declined as the survival of patients with HIV infection has increased through the use of cART.



In Africa, many patients are diagnosed with HIV infection only after developing opportunistic infections. Hence, the optimal management of these opportunistic infections is important if the benefits of subsequent cART are to be realized. The incidence of TE in under-resourced settings is not clear because of a lack of facilities for serologic testing and imaging. AIDS patients who are seropositive for *T. gondii* and who have a CD4+ T lymphocyte count of <100/μL should receive prophylaxis against TE.

Of the currently available agents, trimethoprim-sulfamethoxazole (TMP-SMX) appears to be an effective alternative for treatment of TE in resource-poor settings where the preferred combination of pyrimethamine plus sulfadiazine is not available. The daily dose of TMP-SMX (one double-strength tablet) that is recommended as the preferred regimen for prophylaxis of PcP is effective against TE. If patients cannot tolerate TMP-SMX, the recommended alternative is dapsone-pyrimethamine, which likewise is effective against PcP. Atovaquone with or without pyrimethamine also can be considered. Prophylactic monotherapy with dapsone, pyrimethamine, azithromycin, clarithromycin, or aerosolized pentamidine is probably insufficient. AIDS patients who are seronegative for *Toxoplasma* and are not receiving prophylaxis for PcP should be retested for IgG antibody to *Toxoplasma* if their CD4+ T cell count drops to <100/μL. If seroconversion has taken place, then the patient should be given prophylaxis as described above.

Discontinuing Primary Prophylaxis Current studies indicate that prophylaxis against TE can be discontinued in patients who have responded to cART and whose CD4+ T lymphocyte count has been >200/μL for 3 months. Although patients with CD4+ T lymphocyte counts of <100/μL are at greatest risk for developing TE, the risk that this condition will develop when the count has increased to 100–200/μL has not been established. Thus, prophylaxis should be discontinued when the count has increased to >200/μL. Discontinuation of therapy reduces the pill burden; the potential for drug toxicity, drug interaction, or selection of drug-resistant pathogens; and cost. Prophylaxis should be recommenced if the CD4+ T lymphocyte count again decreases to <100–200/μL.

Individuals who have completed initial therapy for TE should receive treatment indefinitely unless immune reconstitution, with a CD4+ T cell count of >200/μL, occurs as a consequence of cART. Combination therapy with pyrimethamine plus sulfadiazine plus leucovorin is effective for this purpose. An alternative to sulfadiazine in this regimen is clindamycin.

Discontinuing Secondary Prophylaxis (Long-Term Maintenance Therapy)

Patients receiving secondary prophylaxis for TE are at low risk for recurrence when they have completed initial therapy for TE, remain asymptomatic, and have evidence of restored immune function. Individuals with HIV infection should have a CD4+ T lymphocyte count of >200/μL for at least 6 months after cART. This recommendation is consistent with more extensive data indicating the safety of discontinuing secondary prophylaxis for other opportunistic infections during advanced HIV disease. A repeat MRI brain scan is recommended. Secondary prophylaxis should be reinstituted if the CD4+ T lymphocyte count decreases to <200/μL.

PREVENTION

All HIV-infected persons should be counseled regarding sources of *Toxoplasma* infection. The chances of primary infection with *Toxoplasma* can be reduced by not eating undercooked meat and

by avoiding oocyst-contaminated material (i.e., a cat's litter box). Specifically, lamb, beef, and pork should be cooked to an internal temperature of 165°–170°F; from a more practical perspective, meat cooked until it is no longer pink inside usually satisfies this requirement. Hands should be washed thoroughly after work in the garden, and all fruits and vegetables should be washed. Ingestion of raw shellfish is a risk factor for toxoplasmosis, given that the filter-feeding mechanism of clams and mussels concentrates oocysts.

If the patient owns a cat, the litter box should be cleaned or changed daily, preferably by an HIV-negative, nonpregnant person; alternatively, patients should wash their hands thoroughly after changing the litter box. Litter boxes should be changed daily if possible, as freshly excreted oocysts will not have sporulated and will not be infectious. Patients should be encouraged to keep their cats inside and not to adopt or handle stray cats. Cats should be fed only canned or dried commercial food or well-cooked table food, not raw or undercooked meats. Patients need not be advised to part with their cats or to have their cats tested for toxoplasmosis. Blood intended for transfusion into *Toxoplasma*-seronegative immunocompromised individuals should be screened for antibody to *T. gondii*. Although such serologic screening is not routinely performed, seronegative women should be screened for evidence of infection several times during pregnancy if they are exposed to environmental conditions that put them at risk for infection with *T. gondii*. HIV-positive individuals should adhere closely to these preventive measures.

254 Protozoal Intestinal Infections and Trichomoniasis

Peter F. Weller

PROTOZOAL INFECTIONS

GIARDIASIS



Giardia intestinalis (also known as *G. lamblia* or *G. duodenalis*) is a cosmopolitan protozoal parasite that inhabits the small intestines of humans and other mammals. Giardiasis is one of the most common parasitic diseases in both developed and developing countries worldwide, causing both endemic and epidemic intestinal disease and diarrhea.

Life Cycle and Epidemiology (Fig. 254-1) Infection follows the ingestion of environmentally hardy cysts, which excyst in the small intestine, releasing flagellated trophozoites (Fig. 254-2) that multiply by binary fission. *Giardia* remains a pathogen of the proximal small bowel and does not disseminate hematogenously. Trophozoites remain free in the lumen or attach to the mucosal epithelium by means of a ventral sucking disk. As a trophozoite encounters altered conditions, it forms a morphologically distinct cyst, which is the stage of the parasite usually found in the feces. Trophozoites may be present and even predominate in loose or watery stools, but it is the resistant cyst that survives outside the body and is responsible for transmission. Cysts do not tolerate heating or desiccation, but they do remain viable for months in cold fresh water. The number of cysts excreted varies widely but can approach 10^7 per gram of stool.

Ingestion of as few as 10 cysts is sufficient to cause infection in humans. Because cysts are infectious when excreted, person-to-person transmission occurs where fecal hygiene is poor. Giardiasis is especially prevalent in day-care centers; person-to-person spread also takes place in other institutional settings with poor fecal hygiene and during anal-oral contact. If food is contaminated with *Giardia* cysts after cooking or preparation, food-borne transmission can occur. Waterborne transmission accounts for episodic infections (e.g., in campers and travelers) and for major epidemics in metropolitan areas.

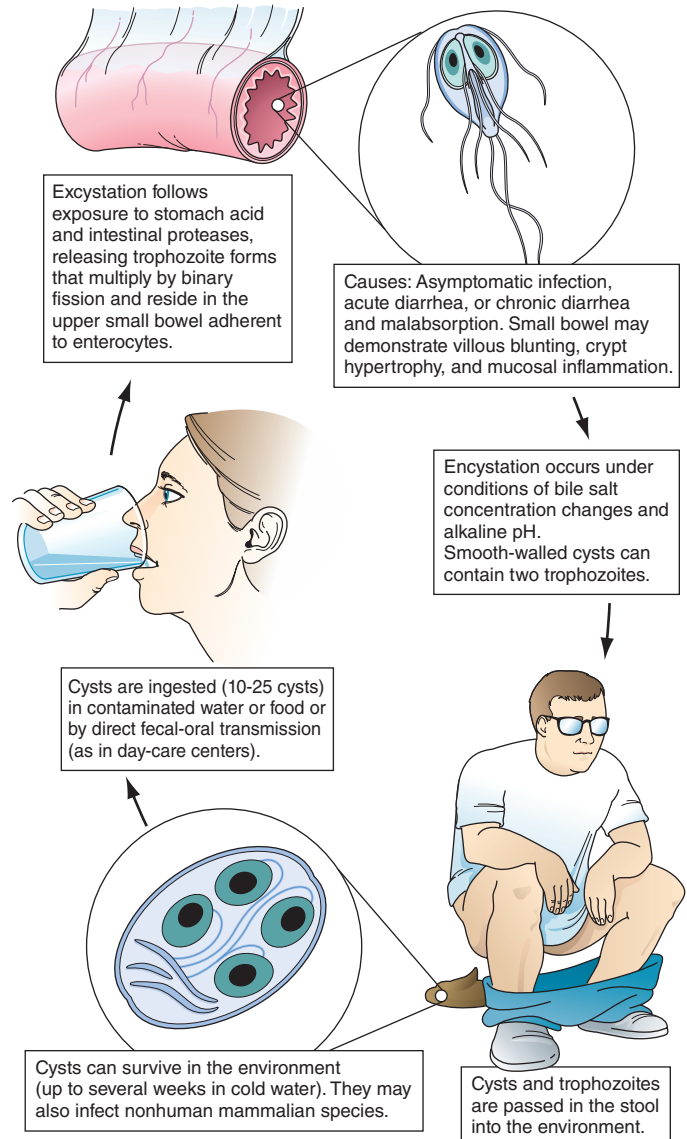


FIGURE 254-1 Life cycle of *Giardia*. (Reprinted with permission from RL Guerrant et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 987. © 2006, with permission from Elsevier Science.)

Surface water, ranging from mountain streams to large municipal reservoirs, can become contaminated with fecally derived *Giardia* cysts. The efficacy of water as a means of transmission is enhanced by the small infectious inoculum of *Giardia*, the prolonged survival of cysts



FIGURE 254-2 Flagellated, binucleate *Giardia* trophozoites.

1406 in cold water, and the resistance of cysts to killing by routine chlorination methods that are adequate for controlling bacteria. Viable cysts can be eradicated from water by either boiling or filtration.



In the United States, *Giardia* (like *Cryptosporidium*; see below) is a common cause of waterborne epidemics of gastroenteritis. *Giardia* is common in developing countries, and infections may be acquired by travelers.

There are several recognized genotypes or assemblages of *G. intestinalis*. Human infections are due to assemblages A and B, whereas other assemblages are more common in other animals, including cats and dogs. Like beavers from reservoirs implicated in epidemics, dogs and cats have been found to be infected with assemblages A and B, an observation suggesting that these animals might be sources of human infection.

Giardiasis, like cryptosporidiosis, creates a significant economic burden because of the costs incurred in the installation of water filtration systems required to prevent waterborne epidemics, in the management of epidemics that involve large communities, and in the evaluation and treatment of endemic infections.

Pathophysiology The reasons that some, but not all, infected patients develop clinical manifestations and the mechanisms by which *Giardia* causes alterations in small-bowel function are largely unknown. Although trophozoites adhere to the epithelium, they are not invasive but may elicit apoptosis of enterocytes, epithelial barrier dysfunction, and epithelial cell malabsorption and secretion. Consequent lactose intolerance and, in a minority of infected adults and children, significant malabsorption are clinical signs of the loss of brush-border enzyme activities. In most infections, the morphology of the bowel is unaltered; however, in chronically infected, symptomatic patients, the histopathologic findings (including flattened villi) and the clinical manifestations at times resemble those of tropical sprue and gluten-sensitive enteropathy. The pathogenesis of diarrhea in giardiasis is not known.

The natural history of *Giardia* infection varies markedly. Infections may be aborted, transient, recurrent, or chronic. *G. intestinalis* parasites vary genotypically, and such variations might contribute to different courses of infection. Parasite as well as host factors may be important in determining the course of infection and disease. Both cellular and humoral responses develop in human infections, but their precise roles in disease pathogenesis and/or control of infection are unknown. Because patients with hypogammaglobulinemia suffer from prolonged, severe infections that are poorly responsive to treatment, humoral immune responses appear to be important. The greater susceptibilities of the young than of the old and of newly exposed persons than of chronically exposed populations suggest that at least partial protective immunity may develop.

Clinical Manifestations Disease manifestations of giardiasis range from asymptomatic carriage to fulminant diarrhea and malabsorption. Most infected persons are asymptomatic, but in epidemics the proportion of symptomatic cases may be higher. Symptoms may develop suddenly or gradually. In persons with acute giardiasis, symptoms develop after an incubation period that lasts at least 5–6 days and usually 1–3 weeks. Prominent early symptoms include diarrhea, abdominal pain, bloating, belching, flatus, nausea, and vomiting. Although diarrhea is common, upper intestinal manifestations such as nausea, vomiting, bloating, and abdominal pain may predominate. The duration of acute giardiasis is usually >1 week, although diarrhea often subsides. Individuals with chronic giardiasis may present with or without having experienced an antecedent acute symptomatic episode. Diarrhea is not necessarily prominent, but increased flatus, loose stools, sulfurous belching, and (in some instances) weight loss occur. Symptoms may be continual or episodic and may persist for years. Some persons who have relatively mild symptoms for long periods recognize the extent of their discomfort only in retrospect. Fever, the presence of blood and/or mucus in

TABLE 254-1 DIAGNOSIS OF INTESTINAL PROTOZOAL INFECTIONS

Parasite	Stool O+P ^a	Fecal Acid-Fast Stain	Fecal Antigen Immunoassays	Fecal NAATs ^b	Other
<i>Giardia</i>	+	–	+	+	
<i>Cryptosporidium</i>	–	+	+	+	
<i>Isospora</i>	–	+	–	+	
<i>Cyclospora</i>	–	+	–	+	
Microsporidia	–	–	–	+	Special fecal stains, tissue biopsies

^aO+P, ova and parasites. ^bNucleic acid amplification tests.

the stools, and other signs and symptoms of colitis are uncommon and suggest a different diagnosis or a concomitant illness. Symptoms tend to be intermittent yet recurring and gradually debilitating, in contrast with the acute disabling symptoms associated with many enteric bacterial infections. Because of the less severe illness and the propensity for chronic infections, patients may seek medical advice late in the course of the illness; however, disease can be severe, resulting in malabsorption, weight loss, growth retardation, and dehydration. A number of extraintestinal manifestations have been described, such as urticaria, anterior uveitis, and arthritis; whether these are caused by giardiasis or concomitant processes is unclear.

Giardiasis can be severe in patients with hypogammaglobulinemia and can complicate other preexisting intestinal diseases, such as that occurring in cystic fibrosis. In patients with AIDS, *Giardia* can cause enteric illness that is refractory to treatment.

Diagnosis (Table 254-1) Giardiasis is diagnosed by detection of parasite antigens in the feces, by identification of cysts in the feces or of trophozoites in the feces or small intestines, or by nucleic acid amplification tests (NAATs). Cysts are oval, measure 8–12 μm × 7–10 μm, and characteristically contain four nuclei. Trophozoites are pear-shaped, dorsally convex, flattened parasites with two nuclei and four pairs of flagella (Fig. 254-2). The diagnosis is sometimes difficult to establish. Direct examination of fresh or properly preserved stools as well as concentration methods should be used. Because cyst excretion is variable and may be undetectable at times, repeated examination of stool, sampling of duodenal fluid, and biopsy of the small intestine may be required to detect the parasite. Tests for parasitic antigens in stool are at least as sensitive and specific as good microscopic examinations and are easier to perform. Newer NAATs are highly sensitive but are not always available for clinical use at present.

TREATMENT GIARDIASIS

Cure rates with metronidazole (250 mg thrice daily for 5 days) are usually >90%. Tinidazole (2 g once by mouth) may be more effective than metronidazole. Albendazole (400 mg daily for 5–10 days) is as effective as metronidazole and is associated with fewer side effects. Nitazoxanide (500 mg twice daily for 3 days) is an alternative agent for treatment of giardiasis. Paromomycin, an oral aminoglycoside that is not well absorbed, can be given to symptomatic pregnant patients, although information is limited on how effectively this agent eradicates infection.

Almost all patients respond to therapy and are cured, although some with chronic giardiasis experience delayed resolution of symptoms after eradication of *Giardia*. For many of the latter patients, residual symptoms probably reflect delayed regeneration of intestinal brush-border enzymes. Continued infection should be documented by stool examinations before treatment is repeated. Patients who remain infected after repeated treatments should be evaluated for reinfection through family members, close personal contacts, and environmental sources as well as for hypogammaglobulinemia. In cases refractory to multiple treatment courses, prolonged therapy with metronidazole (750 mg thrice daily for 21 days) or therapy with varied combinations of multiple agents has been successful.

Prevention Giardiasis can be prevented by consumption of uncontaminated food and water and by personal hygiene during the provision of care for infected children. Boiling or filtering potentially contaminated water prevents infection.

CRYPTOSPORIDIOSIS



The coccidian parasite *Cryptosporidium* causes diarrheal disease that is self-limited in immunocompetent human hosts but can be severe in persons with AIDS or other forms of immunodeficiency. Two species of *Cryptosporidium*, *C. hominis* (especially in the United States, sub-Saharan Africa, and Asia) and *C. parvum* (in Europe), cause most human infections.

Life Cycle and Epidemiology *Cryptosporidium* species are widely distributed in the world. Cryptosporidiosis is acquired by the consumption of oocysts (50% infectious dose: ~132 *C. parvum* oocysts in nonimmune individuals), which excyst to liberate sporozoites that in turn enter and infect intestinal epithelial cells. The parasite's further development involves both asexual and sexual cycles, which produce forms capable of infecting other epithelial cells and of generating oocysts that are passed in the feces. *Cryptosporidium* species infect a number of animals, and *C. parvum* can spread from infected animals to humans. Since oocysts are immediately infectious when passed in feces, person-to-person transmission takes place in day-care centers and among household contacts and medical providers. Waterborne transmission (especially that of *C. hominis*) accounts for infections in travelers and for common-source epidemics. Oocysts are quite hardy and resist killing by routine chlorination. Both drinking water and recreational water (e.g., pools, waterslides) have been increasingly recognized as sources of infection.

Pathophysiology Although intestinal epithelial cells harbor cryptosporidia in an intracellular vacuole, the means by which secretory diarrhea is elicited remain uncertain. No characteristic pathologic changes are found by biopsy. The distribution of infection can be spotty within the principal site of infection, the small bowel. Cryptosporidia are found in the pharynx, stomach, and large bowel of some patients and at times in the respiratory tract. Especially in patients with AIDS, involvement of the biliary tract can cause papillary stenosis, sclerosing cholangitis, or cholecystitis.

Clinical Manifestations Asymptomatic infections can occur in both immunocompetent and immunocompromised hosts. In immunocompetent persons, symptoms develop after an incubation period of ~1 week and consist principally of watery nonbloody diarrhea, sometimes in conjunction with abdominal pain, nausea, anorexia, fever, and/or weight loss. In these hosts, the illness usually subsides after 1–2 weeks. In contrast, in immunocompromised hosts (especially those with AIDS and CD4+ T cell counts <100/μL), diarrhea can be chronic, persistent, and remarkably profuse, causing clinically significant fluid and electrolyte depletion. Stool volumes may range from 1 to 25 L/d. Weight loss, wasting, and abdominal pain may be severe. Biliary tract involvement can manifest as mid-epigastric or right-upper-quadrant pain.

Diagnosis (Table 254-1) Evaluation starts with fecal examination for small oocysts, which are smaller (4–5 μm in diameter) than the fecal stages of most other parasites. Because conventional stool examination for ova and parasites (O+P) does not detect *Cryptosporidium*, specific testing must be requested. Detection is enhanced by evaluation of stools (obtained on multiple days) by several techniques, including modified acid-fast and direct immunofluorescent stains and enzyme immunoassays. Newer NAATs are being employed. Cryptosporidia can also be identified by light and electron microscopy at the apical surfaces of intestinal epithelium from biopsy specimens of the small bowel and, less frequently, the large bowel.

TREATMENT CRYPTOSPORIDIOSIS

Nitazoxanide, approved by the U.S. Food and Drug Administration (FDA) for the treatment of cryptosporidiosis, is available in tablet form for adults (500 mg twice daily for 3 days) and as an elixir for

children. To date, however, this agent has not been effective for the treatment of HIV-infected patients, in whom improved immune status due to antiretroviral therapy can lead to amelioration of cryptosporidiosis. Otherwise, treatment includes supportive care with replacement of fluids and electrolytes and administration of antidiarrheal agents. Biliary tract obstruction may require papillotomy or T-tube placement. Prevention requires minimizing exposure to infectious oocysts in human or animal feces. Use of submicron water filters may minimize acquisition of infection from drinking water.

CYSTOISOSPORIASIS

The coccidian parasite *Cystoisospora belli* causes human intestinal disease. Infection is acquired by the consumption of oocysts, after which the parasite invades intestinal epithelial cells and undergoes both sexual and asexual cycles of development. Oocysts excreted in stool are not immediately infectious but must undergo further maturation.

Although *C. belli* infects many animals, little is known about the epidemiology or prevalence of this parasite in humans. It is most common in tropical and subtropical countries. Acute infections can begin abruptly with fever, abdominal pain, and watery nonbloody diarrhea and can last for weeks or months. In patients who have AIDS or are immunocompromised for other reasons, infections often are not self-limited but rather resemble cryptosporidiosis, with chronic, profuse watery diarrhea. Eosinophilia, which is not found in other enteric protozoan infections, may be detectable. The diagnosis (Table 254-1) is usually made by detection of the large (~25-μm) oocysts in stool by modified acid-fast staining. Oocyst excretion may be low-level and intermittent; if repeated stool examinations are unrevealing, sampling of duodenal contents by aspiration or small-bowel biopsy (often with electron microscopic examination) may be necessary. NAATs are promising newer diagnostic tools.

TREATMENT CYSTOISOSPORIASIS

Trimethoprim-sulfamethoxazole (TMP-SMX, 160/800 mg four times daily for 10 days; and for HIV-infected patients, then continuing three times daily for 3 weeks) is effective. For patients intolerant of sulfonamides, pyrimethamine (50–75 mg/d) can be used. Relapses can occur in persons with AIDS and necessitate maintenance therapy with TMP-SMX (160/800 mg three times per week).

CYCLOSPORIASIS



Cyclospora cayetanensis, a cause of diarrheal illness, is globally distributed; illness due to *C. cayetanensis* has been reported in the United States, Asia, Africa, Latin America, and Europe. The epidemiology of this parasite has not yet been fully defined, but waterborne transmission and food-borne transmission (e.g., by basil, sweet peas, and imported raspberries) have been recognized. The full spectrum of illness attributable to *Cyclospora* has not been delineated. Some infected patients may be without symptoms, but many have diarrhea, flulike symptoms, and flatulence and belching. The illness can be self-limited, can wax and wane, or, in many cases, can involve prolonged diarrhea, anorexia, and upper gastrointestinal symptoms, with sustained fatigue and weight loss in some instances. Diarrheal illness may persist for >1 month. *Cyclospora* can cause enteric illness in patients infected with HIV.

The parasite is detectable in epithelial cells of small-bowel biopsy samples and elicits secretory diarrhea by unknown means. The absence of fecal blood and leukocytes indicates that disease due to *Cyclospora* is not caused by destruction of the small-bowel mucosa. The diagnosis (Table 254-1) can be made by detection of spherical 8- to 10-μm oocysts in the stool, although routine stool O+P examinations are not sufficient. Specific fecal examinations must be requested to detect the oocysts, which are variably acid-fast and are fluorescent when viewed with ultraviolet light microscopy. Newer NAATs are proving to be sensitive. Cyclosporiasis should be considered in the differential diagnosis of prolonged diarrhea, with or without a history of travel by the patient to other countries.

Cyclosporiasis is treated with TMP-SMX (160/800 mg twice daily for 7–10 days). HIV-infected patients may experience relapses after such treatment and thus may require longer-term suppressive maintenance therapy.

MICROSPORIDIOSIS

Microsporidia are obligate intracellular spore-forming protozoa that infect many animals and cause disease in humans, especially as opportunistic pathogens in AIDS. Microsporidia are members of a distinct phylum, Microspora, which contains dozens of genera and hundreds of species. The various microsporidia are differentiated by their developmental life cycles, ultrastructural features, and molecular taxonomy based on ribosomal RNA. The complex life cycles of the organisms result in the production of infectious spores (Fig. 254-3). Currently, eight genera of microsporidia—*Encephalitozoon*, *Pleistophora*, *Nosema*, *Vittaforma*, *Trachipleistophora*, *Anncalia*, *Microsporidium*, and *Enterocytozoon*—are recognized as causes of human disease. Although some microsporidia are probably prevalent causes of self-limited or asymptomatic infections in immunocompetent patients, little is known about how microsporidiosis is acquired.

Microsporidiosis is most common among patients with AIDS, less common among patients with other types of immunocompromise, and

rare among immunocompetent hosts. In patients with AIDS, intestinal infections with *Enterocytozoon bieneusi* and *Encephalitozoon* (formerly *Septata*) *intestinalis* are recognized to contribute to chronic diarrhea and wasting; these infections had been found in 10–40% of patients with chronic diarrhea. Both organisms have been found in the biliary tracts of patients with cholecystitis. *E. intestinalis* may also disseminate to cause fever, diarrhea, sinusitis, cholangitis, and bronchiolitis. In patients with AIDS, *Encephalitozoon hellem* has caused superficial keratoconjunctivitis as well as sinusitis, respiratory tract disease, and disseminated infection. Myositis due to *Pleistophora* has been documented. *Nosema*, *Vittaforma*, and *Microsporidium* have caused stromal keratitis associated with trauma in immunocompetent patients.

Microsporidia are small gram-positive organisms with mature spores measuring 0.5–2 $\mu\text{m} \times 1\text{--}4 \mu\text{m}$. Diagnosis of microsporidial infections in tissue often requires electron microscopy, although intracellular spores can be visualized by light microscopy with hematoxylin and eosin, Giemsa, or tissue Gram's stain. For the diagnosis of intestinal microsporidiosis, modified trichrome or chromotrope 2R-based staining and Uvitex 2B or calcofluor fluorescent staining reveal spores in smears of feces or duodenal aspirates. Definitive therapies for microsporidial infections remain to be established. For superficial keratoconjunctivitis due to *E. hellem*, topical therapy with fumagillin suspension has shown promise (Chap. 246e). For enteric infections with *E. bieneusi* and *E. intestinalis* in HIV-infected patients, therapy with albendazole may be efficacious (Chap. 246e).

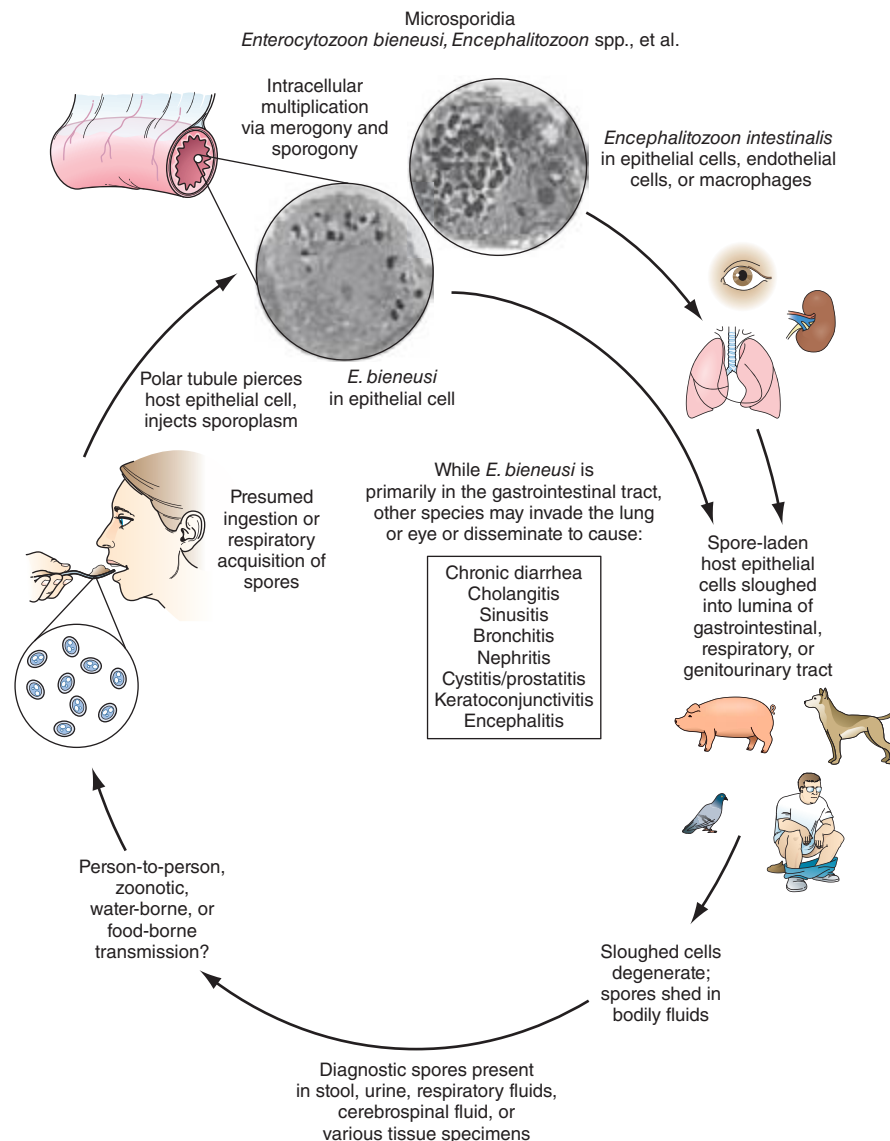


FIGURE 254-3 Life cycle of microsporidia. (Reprinted with permission from RL Guerrant et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 1128. © 2006, with permission from Elsevier Science.)

OTHER INTESTINAL PROTOZOA



Balantidiasis *Balantidium coli* is a large ciliated protozoal parasite that can produce a spectrum of large-intestinal disease analogous to amebiasis. The parasite is widely distributed in the world. Since it infects pigs, cases in humans are more common where pigs are raised. Infective cysts can be transmitted from person to person and through water, but many cases are due to the ingestion of cysts derived from porcine feces in association with slaughtering, with use of pig feces for fertilizer, or with contamination of water supplies by pig feces.

Ingested cysts liberate trophozoites, which reside and replicate in the large bowel. Many patients remain asymptomatic, but some have persisting intermittent diarrhea, and a few develop more fulminant dysentery. In symptomatic individuals, the pathology in the bowel—both gross and microscopic—is similar to that seen in amebiasis, with varying degrees of mucosal invasion, focal necrosis, and ulceration. Balantidiasis, unlike amebiasis, only rarely spreads hematogenously to other organs. The diagnosis is made by detection of the trophozoite stage in stool or sampled colonic tissue. Tetracycline (500 mg four times daily for 10 days) is an effective therapeutic agent.

Blastocystosis *Blastocystis hominis* remains an organism of uncertain pathogenicity. Some patients who pass *B. hominis* in their stools are asymptomatic, whereas others have diarrhea and associated intestinal symptoms. Diligent evaluation reveals other potential bacterial, viral, or protozoal causes of diarrhea in some but not all patients with symptoms. Because the pathogenicity of *B. hominis* is uncertain and because therapy for *Blastocystis* infection is neither specific nor uniformly effective, patients with prominent intestinal symptoms should be fully evaluated for other infectious causes of diarrhea. If diarrheal symptoms associated with *Blastocystis* are prominent, either metronidazole (750 mg thrice daily for 10 days) or TMP-SMX (160 mg/800 mg twice daily for 7 days) can be used.

Dientamoebiasis *Dientamoeba fragilis* is unique among intestinal protozoa in that it has a trophozoite stage but not a cyst stage. How trophozoites survive to transmit infection is not known. When symptoms develop in patients with *D. fragilis* infection, they are generally mild and include intermittent diarrhea, abdominal pain, and anorexia. The diagnosis is made by the detection of trophozoites in stool; the lability of these forms accounts for the greater yield when fecal samples are preserved immediately after collection. Since fecal excretion rates vary, examination of several samples obtained on alternate days increases the rate of detection. Iodoquinol (650 mg three times daily for 20 days) or paromomycin (25–35 mg/kg per day in three doses for 7 days) is appropriate for treatment.

TRICHOMONIASIS

Various species of trichomonads can be found in the mouth (in association with periodontitis) and occasionally in the gastrointestinal tract.

Trichomonas vaginalis—one of the most prevalent protozoal parasites in the United States—is a pathogen of the genitourinary tract and a major cause of symptomatic vaginitis (**Chap. 163**).

Life Cycle and Epidemiology *T. vaginalis* is a pear-shaped, actively motile organism that measures about 10×7 μm , replicates by binary fission, and inhabits the lower genital tract of females and the urethra and prostate of males. In the United States, it accounts for ~3 million infections per year in women. While the organism can survive for a few hours in moist environments and could be acquired by direct contact, person-to-person venereal transmission accounts for virtually all cases of trichomoniasis. Its prevalence is greatest among persons with multiple sexual partners and among those with other sexually transmitted diseases (**Chap. 163**).

Clinical Manifestations Many men infected with *T. vaginalis* are asymptomatic, although some develop urethritis and a few have epididymitis or prostatitis. In contrast, infection in women, which has an incubation period of 5–28 days, is usually symptomatic and manifests with malodorous vaginal discharge (often yellow), vulvar erythema and itching, dysuria or urinary frequency (in 30–50% of patients), and dyspareunia. These manifestations, however, do not clearly distinguish trichomoniasis from other types of infectious vaginitis.

Diagnosis Detection of motile trichomonads by microscopic examination of wet mounts of vaginal or prostatic secretions has been the conventional means of diagnosis. Although this approach provides an immediate diagnosis, its sensitivity for the detection of *T. vaginalis* is only ~50–60% in routine evaluations of vaginal secretions. Direct immunofluorescent antibody staining is more sensitive (70–90%) than wet-mount examinations. *T. vaginalis* can be recovered from the urethra of both males and females and is detectable in males after prostatic massage. A new NAAT, APTIMA, is FDA approved and is highly sensitive and specific for urine and for endocervical and vaginal swabs from women.

TREATMENT TRICHOMONIASIS

Metronidazole (either a single 2-g dose or 500-mg doses twice daily for 7 days) or tinidazole (a single 2-g dose) is effective. All sexual partners must be treated concurrently to prevent reinfection, especially from asymptomatic males. In males with persistent symptomatic urethritis after therapy for nongonococcal urethritis, metronidazole therapy should be considered for possible trichomoniasis. Alternatives to metronidazole for treatment during pregnancy are not readily available. Reinfection often accounts for apparent treatment failures, but strains of *T. vaginalis* exhibiting high-level resistance to metronidazole have been encountered. Treatment of these resistant infections with higher oral doses, parenteral doses, or concurrent oral and vaginal doses of metronidazole or with tinidazole has been successful.

SECTION 19 HELMINTHIC INFECTIONS

255e Introduction to Helminthic Infections

Peter F. Weller

The word *helminth* is derived from the Greek *helmins* (“parasitic worm”). Helminthic worms are highly prevalent and, depending on the species, may exist as free-living organisms or as parasites of plant or animal hosts. The parasitic helminths have co-evolved with specific mammalian and other host species. Accordingly, most helminthic infections are restricted to nonhuman hosts, and only rarely do these zoonotic helminths accidentally cause human infections.

Helminthic parasites of humans belong to two phyla: Nematelminthes, which includes nematodes (roundworms), and Platyhelminthes, which includes cestodes (tapeworms) and trematodes (flukes). Helminthic parasites of humans reside within the human body and hence are the cause of true infections. In contrast, parasites of other genera that reside only on mucocutaneous surfaces of humans (e.g., the parasites causing myiasis and scabies) are considered to represent infestations rather than infections.

Helminthic parasites differ substantially from protozoan parasites in several respects. First, protozoan parasites are unicellular organisms, whereas helminthic parasites are multicellular worms that possess differentiated organ systems. Second, helminthic parasites have complex life cycles that require sequential stages of development outside the human host. Thus, most helminths do not complete their replication within the human host; rather, they develop to a certain stage within the mammalian host and, as part of their obligatory life cycle, must mature further outside that host. During the “extra-human” stages of their life cycle, helminths exist either as free-living organisms or as parasites within another host species and thereafter mature into new developmental stages capable of infecting humans. Thus, with only two exceptions (*Strongyloides stercoralis* and *Capillaria philippinensis*, which are capable of internal reinfection), increases in the number of adult helminths (i.e., the “worm burden”) within the human host require repeated exogenous reinfections. In the case of protozoan parasites, a brief, even singular exposure (e.g., a single mosquito bite transmitting malaria) may lead rapidly to intense parasite loads and overwhelming infections; in contrast, for all but the two helminths noted above, increases in worm burden require multiple and usually ongoing exposures to infectious forms, such as ingestion of eggs of intestinal helminths or waterborne exposures to infectious cercariae of *Schistosoma mansoni*. This requirement is germane both to the consideration of helminthic infections in individuals and to ongoing global efforts to interrupt and/or minimize the acquisition of helminthic infections by humans.

Third, helminthic infections have a predilection toward stimulation of host immune responses that elicit eosinophilia within human tissues and blood. The many protozoan infections characteristically do not elicit eosinophilia in infected humans, with only three exceptions (two intestinal protozoan parasites, *Cystoisospora belli* and *Dientamoeba fragilis*, and tissue-borne *Sarcocystis* species). The magnitude of helminth-elicited eosinophilia tends to correlate with the extent of tissue invasion by larvae or adult helminths. For example, in several helminthic infections, including acute schistosomiasis (Katayama syndrome), paragonimiasis, and hookworm and *Ascaris* infections, eosinophilia is most pronounced during the early phases of infection, when migrations of infecting larvae and progression of subsequent developmental stages through the tissues are greatest. In established infections, local eosinophilia is often present around helminths in tissues, but blood eosinophilia may be intermittent, mild, or absent. In helminthic infections in which parasites are well contained within tissues (e.g.,

echinococcal cysts) or confined within the lumen of the intestinal tract (e.g., adult *Ascaris* or tapeworms), eosinophilia is usually absent.

NEMATODES

Nematodes are nonsegmented roundworms. Species of nematodes are remarkably diverse and abundant in nature. Among the many thousands of nematode species, few are parasites of humans. Most nematodes are free-living, and these species have variably evolved to survive in diverse ecologic niches, including saltwater, freshwater, or soil. The well-studied organism *Caenorhabditis elegans* is a free-living nematode. Nematodes can be either beneficial or deleterious parasites of plants. Parasitic nematodes have co-evolved with specific mammalian hosts and have no capacity to live their full life cycles in other hosts. Uncommonly, humans are exposed to infectious stages of nonhuman nematode parasites, and the resultant zoonotic nematode infections can elicit inflammatory and immune responses as larval forms migrate and die in the unsuitable human host. Examples include pulmonary coin lesions due to mosquito-transmitted infections with the dog heartworm *Dirofilaria immitis*; eosinophilic meningoencephalitis due to ingested eggs of the raccoon ascarid *Baylisascaris procyonis*; and eosinophilic meningitis due to ingestion of larvae of the rat lungworm *Angiostrongylus cantonensis*.

Nematode parasites of humans include worms that reside in the intestinal tract or localize in extraintestinal vascular or tissue sites. Roundworms are bisexual, with separate male and female forms (except for *S. stercoralis*, whose adult females are hermaphroditic in the human intestinal tract). Depending on the species, fertilized females release either larvae or eggs containing larvae. Nematodes have five developmental stages: an adult stage and four sequential larval stages. These parasites characteristically are surrounded by a durable outer cuticular layer. Nematodes have a nervous system; a muscular system, including muscle cells under the cuticle; and a developed intestinal tract, including an oral cavity and an elongated gut that ends in an anal pore. Adults may range in size from minute to >1 meter in length (with *Dracunculus medinensis*, for example, at the long end of this spectrum).

Humans acquire infections with nematode parasites by various routes, depending on the parasitic species. Ingestion of eggs passed in human feces is a major global health problem with many of the intestinal helminths (e.g., *Ascaris lumbricoides*). In other species, infecting larvae penetrate skin exposed to fecally contaminated soil (e.g., *S. stercoralis*) or traverse the skin after the bite of infected insect vectors (e.g., filariae). Some nematode infections are acquired by consumption of specific animal-derived foods (e.g., trichinellosis from raw or undercooked pork or wild carnivorous mammals). As noted above, only two nematodes, *S. stercoralis* and *C. philippinensis*, can internally reinfect humans; thus, for all other nematodes, any increases in worm burden must be due to continued exogenous reinfections.

CESTODES

Tapeworms are the cestode parasites of humans. Adult tapeworms are elongated, segmented, hermaphroditic flatworms that reside in the intestinal lumen or, in their larval forms, may live in extraintestinal tissues. Tapeworms include a head (*scolex*) and a number of attached segments (*proglottids*). The worms attach to the intestinal tract via their scolices, which may possess suckers, hooks, or grooves. The scolex is the site of formation of new proglottids. Tapeworms do not have a functional gut tract; rather, each tapeworm segment passively and actively obtains nutrients through its specialized surface tegument. Mature proglottids possess both male and female sex organs, but insemination usually occurs between adjacent proglottids. Fertilized proglottids release eggs that are passed in the feces. When ingested by an intermediate host, an egg releases an oncosphere that penetrates the

255e-2 gut and develops further in tissues as a cysticercus. Humans acquire infection by ingesting animal tissues that contain cysticerci, and the resultant tapeworms develop and reside in the proximal small bowel (e.g., *Taenia solium*, *T. saginata*). Alternatively, if humans ingest eggs of these cestodes that have been passed in human or animal feces, oncospheres develop and can cause space-occupying extraintestinal cystic lesions in tissues; examples include cysticercosis due to *T. solium* and hydatid disease due to species of *Echinococcus*.

TREMATODES

Trematodes of medical importance include blood flukes, intestinal flukes, and tissue flukes. Adult flukes are often leaf-shaped flatworms. Oral and/or ventral suckers help adult flukes maintain their positions in situ. Flukes have an oral cavity but no distal anal pore. Nutrients are obtained both through their integument and by ingestion into the blind intestinal tract. Flukes are hermaphroditic except for blood flukes (schistosomes), which are bisexual. Eggs are passed in human feces (*Fasciola*, *Fasciolopsis*, *Clonorchis*, *Schistosoma japonicum*, *S. mansoni*), urine (*S. haematobium*), or sputum and feces (*Paragonimus*).

Expelled eggs release miracidia—usually in water—that infect specific snail species. Within snails, parasites multiply and cercariae are released. Depending on the species, cercariae can penetrate the skin (schistosomes) or can develop into metacercariae that can be ingested with plants (e.g., watercress for *Fasciola*) or with fish (*Clonorchis*) or crabs (*Paragonimus*).

CONCLUSION



Many of the so-called neglected tropical diseases are due to helminthic infections. The health impacts of many helminthic infections are varied and are based on the frequent need for repeated exposures to increase the worm burdens in infected humans. In global regions where exposures to specific helminths occur even in childhood (e.g., fecally derived intestinal nematodes, mosquito-transmitted filariae, or waterborne snail-transmitted schistosome infections), the morbidities in infected individuals can include nutritional, developmental, cognitive, and functional impairments. Ongoing global mass-treatment programs are currently aimed at diminishing the local prevalences of specific helminths and their consequent impacts on the health of local populations.

256 Trichinellosis and Other Tissue Nematode Infections

Peter F. Weller

Nematodes are elongated, symmetric roundworms. Parasitic nematodes of medical significance may be broadly classified as either predominantly intestinal or tissue nematodes. This chapter covers the tissue nematodes that cause trichinellosis, visceral and ocular larva migrans, cutaneous larva migrans, cerebral angiostrongyliasis, and gnathostomiasis. All of these zoonotic infections result from incidental exposure to infectious nematodes. The clinical symptoms of these infections are due largely to invasive larval stages that (except in the case of *Trichinella*) do not reach maturity in humans.

TRICHINELLOSIS

Trichinellosis develops after the ingestion of meat containing cysts of *Trichinella* (e.g., pork or other meat from a carnivore). Although most infections are mild and asymptomatic, heavy infections can cause severe enteritis, periorbital edema, myositis, and (infrequently) death.

Life Cycle and Epidemiology Eight species of *Trichinella* are recognized as causes of infection in humans. Two species are distributed worldwide: *T. spiralis*, which is found in a great variety of carnivorous and omnivorous animals, and *T. pseudospiralis*, which is found in mammals and birds. *T. nativa* is present in Arctic regions and infects bears; *T. nelsoni* is found in equatorial eastern Africa, where it is common among felid predators and scavengers such as hyenas and bush pigs; and *T. britovi* is found in Europe, western Africa, and western Asia among carnivores but not among domestic swine. *T. murrelli* is present in North American game animals.

After human consumption of trichinose meat, encysted larvae are liberated by digestive acid and proteases (Fig. 256-1). The larvae invade the small-bowel mucosa and mature into adult worms. After

~1 week, female worms release newborn larvae that migrate via the circulation to striated muscle. The larvae of all species except *T. pseudospiralis*, *T. papuae*, and *T. zimbabwensis* then encyst by inducing a radical transformation in the muscle cell architecture. Although host immune responses may help to expel intestinal adult worms, they have few deleterious effects on muscle-dwelling larvae.

Human trichinellosis is often caused by the ingestion of infected pork products and thus can occur in almost any location where the meat of domestic or wild swine is eaten. Human trichinellosis may also be acquired from the meat of other animals, including dogs (in parts of Asia and Africa), horses (in Italy and France), and bears and walrus (in northern regions). Although cattle (being herbivores) are not natural hosts of *Trichinella*, beef has been implicated in outbreaks when contaminated or adulterated with trichinose pork. Laws that prohibit the feeding of uncooked garbage to pigs have greatly reduced the transmission of trichinellosis in the United States. About 12 cases of trichinellosis are reported annually in this country, but most mild cases probably remain undiagnosed. Recent U.S. and Canadian outbreaks have been attributable to consumption of wild game (especially bear meat) and, less frequently, of pork.

Pathogenesis and Clinical Features Clinical symptoms of trichinellosis arise from the successive phases of parasite enteric invasion, larval migration, and muscle encystment (Fig. 256-1). Most light infections (those with <10 larvae per gram of muscle) are asymptomatic, whereas heavy infections (which can involve >50 larvae per gram of muscle) can be life-threatening. Invasion of the gut by large numbers of parasites occasionally provokes diarrhea during the first week after infection. Abdominal pain, constipation, nausea, or vomiting also may be prominent.

Symptoms due to larval migration and muscle invasion begin to appear in the second week after infection. The migrating *Trichinella* larvae provoke a marked local and systemic hypersensitivity reaction, with fever and hypereosinophilia. Periorbital and facial edema is common, as are hemorrhages in the subconjunctivae, retina, and nail beds ("splinter" hemorrhages). A maculopapular rash, headache, cough, dyspnea, or dysphagia sometimes develops. Myocarditis with tachyarrhythmias or heart failure—and, less commonly, encephalitis or pneumonitis—may develop and accounts for most deaths of patients with trichinellosis.

Upon onset of larval encystment in muscle 2–3 weeks after infection, symptoms of myositis with myalgias, muscle edema, and weakness develop, usually overlapping with the inflammatory reactions to migrating larvae. The most commonly involved muscle groups include the extraocular muscles; the biceps; and the muscles of the jaw, neck, lower back, and diaphragm. Peaking ~3 weeks after infection, symptoms subside only gradually during a prolonged convalescence. Uncommon infections with *T. pseudospiralis*, whose larvae do not encapsulate in muscles, elicit prolonged polymyositis-like illness.

Laboratory Findings and Diagnosis Blood eosinophilia develops in >90% of patients with symptomatic trichinellosis and may peak at a level of >50% 2–4 weeks after infection. Serum levels of muscle enzymes, including creatine phosphokinase, are elevated in most symptomatic patients. Patients should be questioned thoroughly about their consumption of pork or wild animal meat and about illness in other individuals who ate the same meat. A presumptive clinical diagnosis can be based on fevers, eosinophilia, periorbital edema, and myalgias after a suspect meal. A rise in the titer of parasite-specific antibody, which usually does not occur until after the third week

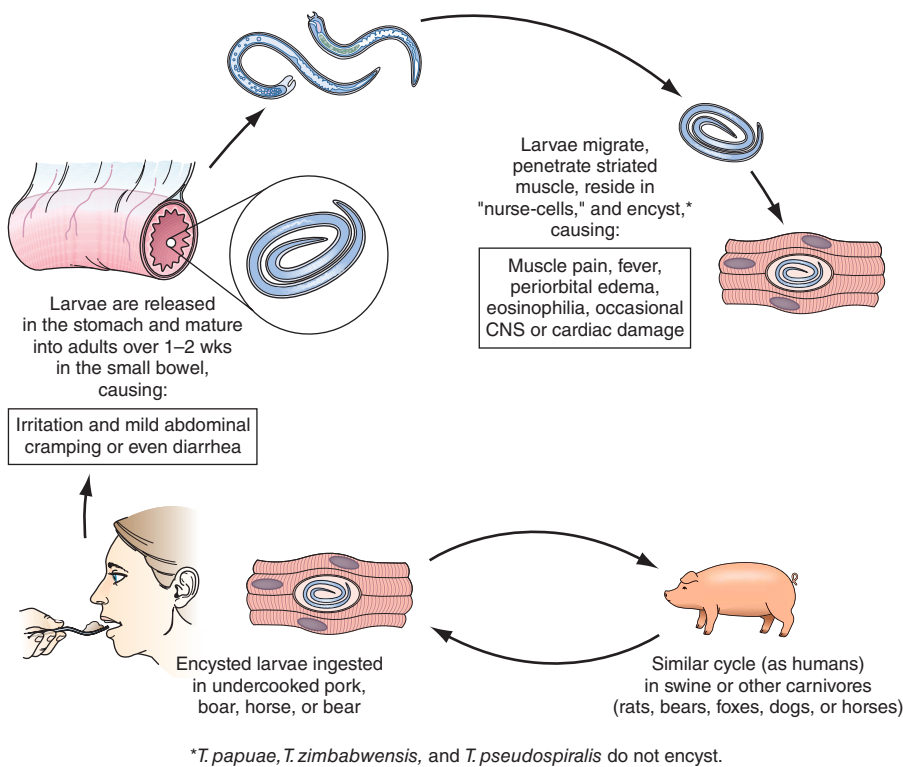


FIGURE 256-1 Life cycle of *Trichinella spiralis* (cosmopolitan); *nelsoni* (equatorial Africa); *britovi* (Europe, western Africa, western Asia); *nativa* (Arctic); *murrelli* (North America); *papuae* (Papua New Guinea); *zimbabwensis* (Tanzania); and *pseudospiralis* (cosmopolitan). CNS, central nervous system. (Reprinted from RL Guerrant et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 1218. © 2006, with permission from Elsevier Science.)

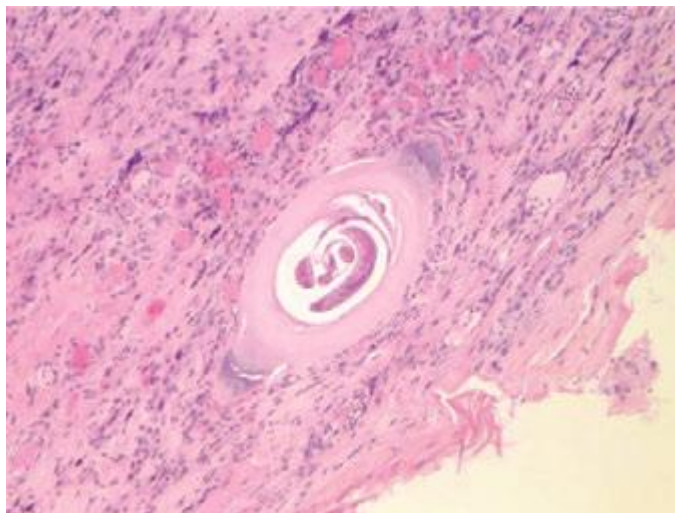


FIGURE 256-2 *Trichinella larva* encysted in a characteristic hyalinized capsule in striated muscle tissue. (Photo/Wadsworth Center, New York State Department of Health. Reprinted from *MMWR* 53:606, 2004; public domain.)

of infection, confirms the diagnosis. Alternatively, a definitive diagnosis requires surgical biopsy of at least 1 g of involved muscle; the yields are highest near tendon insertions. The fresh muscle tissue should be compressed between glass slides and examined microscopically (Fig. 256-2), because larvae may be missed by examination of routine histopathologic sections alone.

TREATMENT TRICHINELLOSIS

Most lightly infected patients recover uneventfully with bed rest, antipyretics, and analgesics. Glucocorticoids like prednisone (Table 256-1) are beneficial for severe myositis and myocarditis. Mebendazole and albendazole are active against enteric stages of the parasite, but their efficacy against encysted larvae has not been conclusively demonstrated.

Prevention Larvae may be killed by cooking pork until it is no longer pink or by freezing it at -15°C for 3 weeks. However, Arctic *T. nativa*

TABLE 256-1 THERAPY FOR TISSUE NEMATODE INFECTIONS

Infection	Severity	Treatment
Trichinellosis	Mild	Supportive
	Moderate	Albendazole (400 mg bid \times 8–14 days) or Mebendazole (200–400 mg tid \times 3 days, then 400 mg tid \times 8–14 days)
	Severe	Add glucocorticoids (e.g., prednisone, 1 mg/kg qd \times 5 days)
Visceral larva migrans	Mild to moderate	Supportive
	Severe	Glucocorticoids (as above)
	Ocular	Not fully defined; albendazole (800 mg bid for adults, 400 mg bid for children) with glucocorticoids \times 5–20 days has been effective
Cutaneous larva migrans		Ivermectin (single dose, 200 $\mu\text{g}/\text{kg}$) or Albendazole (200 mg bid \times 3 days)
Angiostrongyliasis	Mild to moderate	Supportive
	Severe	Glucocorticoids (as above)
Gnathostomiasis		Ivermectin (200 $\mu\text{g}/\text{kg}$ per day \times 2 days) or Albendazole (400 mg bid \times 21 days)

larvae in walrus or bear meat are relatively resistant and may remain viable despite freezing.

VISCERAL AND OCULAR LARVA MIGRANS

Visceral larva migrans is a syndrome caused by nematodes that are normally parasitic for nonhuman host species. In humans, these nematode larvae do not develop into adult worms but instead migrate through host tissues and elicit eosinophilic inflammation. The most common form of visceral larva migrans is toxocariasis due to larvae of the canine ascarid *Toxocara canis*; the syndrome is due less commonly to the feline ascarid *T. cati* and even less commonly to the pig ascarid *Ascaris suum*. Rare cases with eosinophilic meningoencephalitis have been caused by the raccoon ascarid *Baylisascaris procyonis*.



Life Cycle and Epidemiology The canine roundworm *T. canis* is distributed among dogs worldwide. Ingestion of infective eggs by dogs is followed by liberation of *Toxocara* larvae, which penetrate the gut wall and migrate intravascularly into canine tissues, where most remain in a developmentally arrested state. During pregnancy, some larvae resume migration in bitches and infect puppies prenatally (through transplacental transmission) or after birth (through suckling). Thus, in lactating bitches and puppies, larvae return to the intestinal tract and develop into adult worms, which produce eggs that are released in the feces. Eggs must undergo embryonation over several weeks to become infectious. Humans acquire toxocariasis mainly by eating soil contaminated by puppy feces that contains infective *T. canis* eggs. Visceral larva migrans is most common among children who habitually eat dirt.

Pathogenesis and Clinical Features Clinical disease most commonly afflicts preschool children. After humans ingest *Toxocara* eggs, the larvae hatch and penetrate the intestinal mucosa, from which they are carried by the circulation to a wide variety of organs and tissues. The larvae invade the liver, lungs, central nervous system (CNS), and other sites, provoking intense local eosinophilic granulomatous responses. The degree of clinical illness depends on larval number and tissue distribution, reinfection, and host immune responses. Most light infections are asymptomatic and may be manifest only by blood eosinophilia. Characteristic symptoms of visceral larva migrans include fever, malaise, anorexia and weight loss, cough, wheezing, and rashes. Hepatosplenomegaly is common. These features may be accompanied by extraordinary peripheral eosinophilia, which may approach 90%. Uncommonly, seizures or behavioral disorders develop. Rare deaths are due to severe neurologic, pneumonic, or myocardial involvement.

The ocular form of the larva migrans syndrome occurs when *Toxocara* larvae invade the eye. An eosinophilic granulomatous mass, most commonly in the posterior pole of the retina, develops around the entrapped larva. The retinal lesion can mimic retinoblastoma in appearance, and mistaken diagnosis of the latter condition can lead to unnecessary enucleation. The spectrum of eye involvement also includes endophthalmitis, uveitis, and chorioretinitis. Unilateral visual disturbances, strabismus, and eye pain are the most common presenting symptoms. In contrast to visceral larva migrans, ocular toxocariasis usually develops in older children or young adults with no history of pica; these patients seldom have eosinophilia or visceral manifestations.

Diagnosis In addition to eosinophilia, leukocytosis and hypergammaglobulinemia may be evident. Transient pulmonary infiltrates are apparent on chest x-rays of about one-half of patients with symptoms of pneumonitis. The clinical diagnosis can be confirmed by an enzyme-linked immunosorbent assay for toxocaral antibodies. Stool examination for parasite eggs is worthless in toxocariasis, since the larvae do not develop into egg-producing adults in humans.

TREATMENT VISCERAL AND OCULAR LARVA MIGRANS

The vast majority of *Toxocara* infections are self-limited and resolve without specific therapy. In patients with severe myocardial, CNS, or pulmonary involvement, glucocorticoids may be employed to reduce inflammatory complications. Available anthelmintic drugs,

1412 including mebendazole and albendazole, have not been shown conclusively to alter the course of larva migrans. Control measures include prohibiting dog excreta in public parks and playgrounds, deworming dogs, and preventing pica in children. Treatment of ocular disease is not fully defined, but the administration of albendazole in conjunction with glucocorticoids has been effective (Table 256-1).

CUTANEOUS LARVA MIGRANS

Cutaneous larva migrans (“creeping eruption”) is a serpiginous skin eruption caused by burrowing larvae of animal hookworms, usually the dog and cat hookworm *Ancylostoma braziliense*. The larvae hatch from eggs passed in dog and cat feces and mature in the soil. Humans become infected after skin contact with soil in areas frequented by dogs and cats, such as areas underneath house porches. Cutaneous larva migrans is prevalent among children and travelers in regions with warm humid climates, including the southeastern United States.

After larvae penetrate the skin, erythematous lesions form along the tortuous tracks of their migration through the dermal-epidermal junction; the larvae advance several centimeters in a day. The intensely pruritic lesions may occur anywhere on the body and can be numerous if the patient has lain on the ground. Vesicles and bullae may form later. The animal hookworm larvae do not mature in humans and, without treatment, will die after an interval ranging from weeks to a couple of months, with resolution of skin lesions. The diagnosis is made on clinical grounds. Skin biopsies only rarely detect diagnostic larvae. Symptoms can be alleviated by ivermectin or albendazole (Table 256-1).

ANGIOSTRONGYLIASIS

Angiostrongylus cantonensis, the rat lung worm, is the most common cause of human eosinophilic meningitis (Fig. 256-3).

Life Cycle and Epidemiology This infection occurs principally in Southeast Asia and the Pacific Basin but has spread to other areas of the world, including the Caribbean islands, countries in Central and South America, and the southern United States. *A. cantonensis* larvae produced by adult worms in the rat lung migrate to the gastrointestinal tract and are expelled with the feces. They develop into infective larvae in land snails and slugs. Humans acquire the infection by ingesting raw infected mollusks; vegetables contaminated by mollusk

slime; or crabs, freshwater shrimp, and certain marine fish that have themselves eaten infected mollusks. The larvae then migrate to the brain.

Pathogenesis and Clinical Features The parasites eventually die in the CNS, but not before initiating pathologic consequences that, in heavy infections, can result in permanent neurologic sequelae or death. Migrating larvae cause marked local eosinophilic inflammation and hemorrhage, with subsequent necrosis and granuloma formation around dying worms. Clinical symptoms develop 2–35 days after the ingestion of larvae. Patients usually present with an insidious or abrupt excruciating frontal, occipital, or bitemporal headache. Neck stiffness, nausea and vomiting, and paresthesias are also common. Fever, cranial and extraocular nerve palsies, seizures, paralysis, and lethargy are uncommon.

Laboratory Findings Examination of cerebrospinal fluid (CSF) is mandatory in suspected cases and usually reveals an elevated opening pressure, a white blood cell count of 150–2000/μL, and an eosinophilic pleocytosis of >20%. The protein concentration is usually elevated and the glucose level normal. The larvae of *A. cantonensis* are only rarely seen in CSF. Peripheral-blood eosinophilia may be mild. The diagnosis is generally based on the clinical presentation of eosinophilic meningitis together with a compatible epidemiologic history.

TREATMENT ANGIOSTRONGYLIASIS

Specific chemotherapy is not of benefit in angiostrongyliasis; larvicidal agents may exacerbate inflammatory brain lesions. Management consists of supportive measures, including the administration of analgesics, sedatives, and—in severe cases—glucocorticoids (Table 256-1). Repeated lumbar punctures with removal of CSF can relieve symptoms. In most patients, cerebral angiostrongyliasis has a self-limited course, and recovery is complete. The infection may be prevented by adequately cooking snails, crabs, and prawns and inspecting vegetables for mollusk infestation. Other parasitic or fungal causes of eosinophilic meningitis in endemic areas may include gnathostomiasis (see below), paragonimiasis (Chap. 259), schistosomiasis (Chap. 259), neurocysticercosis (Chap. 260), and coccidioidomycosis (Chap. 237).

GNATHOSTOMIASIS

Infection of human tissues with larvae of *Gnathostoma spinigerum* can cause eosinophilic meningoencephalitis, migratory cutaneous swellings, or invasive masses of the eye and visceral organs.

Life Cycle and Epidemiology Human gnathostomiasis occurs in many countries and is notably endemic in Southeast Asia and parts of China and Japan. In nature, the mature adult worms parasitize the gastrointestinal tract of dogs and cats. First-stage larvae hatch from eggs passed into water and are ingested by *Cyclops* species (water fleas). Infective third-stage larvae develop in the flesh of many animal species (including fish, frogs, eels, snakes, chickens, and ducks) that have eaten either infected *Cyclops* or another infected second intermediate host. Humans typically acquire the infection by eating raw or undercooked fish or poultry. Raw fish dishes, such as *som fak* in Thailand and *sashimi* in Japan, account for many cases of human gnathostomiasis. Some cases in Thailand result from the local practice of applying frog or snake flesh as a poultice.

Pathogenesis and Clinical Features Clinical symptoms are due to the aberrant migration of a single larva into cutaneous, visceral, neural, or ocular tissues. After invasion, larval migration may cause local inflammation, with pain, cough, or hematuria accompanied by fever and eosinophilia. Painful, itchy, migratory swellings may develop in the skin, particularly in the distal extremities

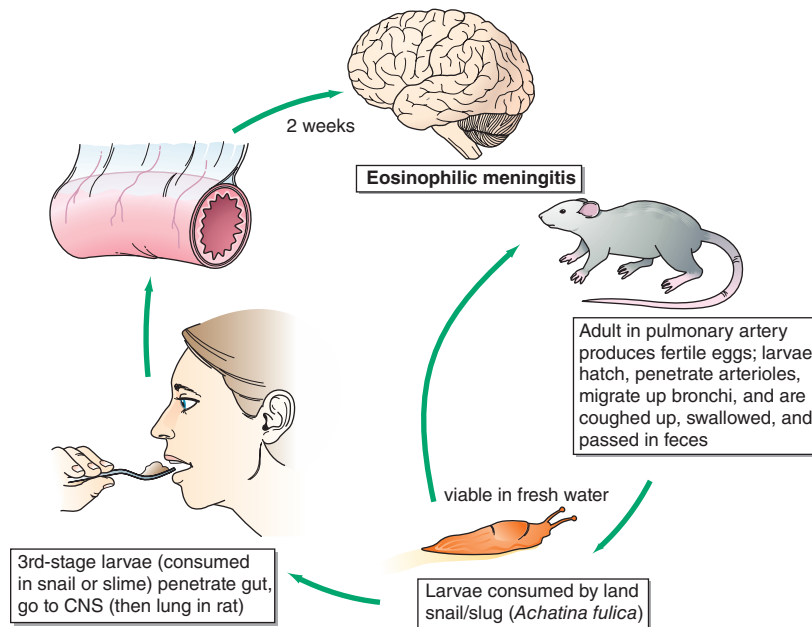


FIGURE 256-3 Life cycle of *Angiostrongylus cantonensis* (rat lung worm), found in Southeast Asia, Pacific Islands, Cuba, Australia, Japan, China, Mauritius, and U.S. ports. CNS, central nervous system. (Reprinted from RL Guerrant et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 1225. © 2006, with permission from Elsevier Science.)

or periorbital area. Cutaneous swellings usually last ~1 week but often recur intermittently over many years. Larval invasion of the eye can provoke a sight-threatening inflammatory response. Invasion of the CNS results in eosinophilic meningitis with myeloencephalitis, a serious complication due to ascending larval migration along a large nerve track. Patients characteristically present with agonizing radicular pain and paresthesias in the trunk or a limb, which are followed shortly by paraplegia. Cerebral involvement, with focal hemorrhages and tissue destruction, is often fatal.

Diagnosis and Treatment Cutaneous migratory swellings with marked peripheral eosinophilia, supported by an appropriate geographic and dietary history, generally constitute an adequate basis for a clinical diagnosis of gnathostomiasis. However, patients may present with ocular or cerebrospinal involvement without antecedent cutaneous swellings. In the latter case, eosinophilic pleocytosis is demonstrable (usually along with hemorrhagic or xanthochromic CSF), but worms are almost never recovered from CSF. Surgical removal of the parasite from subcutaneous or ocular tissue, though rarely feasible, is both diagnostic and therapeutic. Albendazole or ivermectin may be helpful (Table 256-1). At present, cerebrospinal involvement is managed with supportive measures and generally with a course of glucocorticoids. Gnathostomiasis can be prevented by adequate cooking of fish and poultry in endemic areas.

257 Intestinal Nematode Infections

Peter F. Weller, Thomas B. Nutman

More than a billion persons worldwide are infected with one or more species of intestinal nematodes. Table 257-1 summarizes biologic and clinical features of infections due to the major intestinal parasitic nematodes. These parasites are most common in regions with poor fecal sanitation, particularly in resource-poor countries in the tropics and subtropics, but they have also been seen with increasing frequency among immigrants and refugees to resource-rich countries. Although nematode infections are not usually fatal, they contribute to malnutrition and diminished work capacity. It is interesting that these helminth infections may protect some individuals from allergic disease. Humans may on occasion be infected with nematode parasites that ordinarily infect animals; these zoonotic infections produce diseases such as trichostrongyliasis, anisakiasis, capillariasis, and abdominal angiostrongyliasis.

Intestinal nematodes are roundworms; they range in length from 1 mm to many centimeters when mature (Table 257-1). Their life cycles are complex and highly varied; some species, including *Strongyloides stercoralis* and *Enterobius vermicularis*, can be transmitted directly from person to person, while others, such as *Ascaris lumbricoides*, *Necator americanus*, and *Ancylostoma duodenale*, require a soil phase for development. Because most helminth parasites do not self-replicate, the acquisition of a heavy burden of adult worms requires repeated exposure to the parasite in its infectious stage, whether larva or egg. Hence, clinical disease, as opposed to asymptomatic infection, generally develops only with prolonged residence in an endemic area and is typically related to infection intensity. In persons with marginal nutrition, intestinal helminth infections may impair growth and development. Eosinophilia and elevated serum IgE levels are features of many helminth infections and, when unexplained, should always prompt a search for intestinal helminths. Significant protective immunity to intestinal nematodes appears not to develop in humans, although mechanisms of parasite immune evasion and host immune responses to these infections have not been elucidated in detail.

ASCARIASIS

A. lumbricoides is the largest intestinal nematode parasite of humans, reaching up to 40 cm in length. Most infected individuals have low

worm burdens and are asymptomatic. Clinical disease arises from larval migration in the lungs or effects of the adult worms in the intestines.

Life Cycle Adult worms live in the lumen of the small intestine. Mature female *Ascaris* worms are extraordinarily fecund, each producing up to 240,000 eggs a day, which pass with the feces. Ascarid eggs, which are remarkably resistant to environmental stresses, become infective after several weeks of maturation in the soil and can remain infective for years. After infective eggs are swallowed, larvae hatched in the intestine invade the mucosa, migrate through the circulation to the lungs, break into the alveoli, ascend the bronchial tree, and return—through swallowing—to the small intestine, where they develop into adult worms. Between 2 and 3 months elapse between initial infection and egg production. Adult worms live for 1–2 years.



Epidemiology *Ascaris* is widely distributed in tropical and subtropical regions as well as in other humid areas, including the rural southeastern United States. Transmission typically occurs through fecally contaminated soil and is due either to a lack of sanitary facilities or to the use of human feces as fertilizer. With their propensity for hand-to-mouth fecal carriage, younger children are most affected. Infection outside endemic areas, though uncommon, can occur when eggs on transported vegetables are ingested.

Clinical Features During the lung phase of larval migration, ~9–12 days after egg ingestion, patients may develop an irritating nonproductive cough and burning substernal discomfort that is aggravated by coughing or deep inspiration. Dyspnea and blood-tinged sputum are less common. Fever is usually reported. Eosinophilia develops during this symptomatic phase and subsides slowly over weeks. Chest x-rays may reveal evidence of eosinophilic pneumonitis (Löfller's syndrome), with rounded infiltrates a few millimeters to several centimeters in size. These infiltrates may be transient and intermittent, clearing after several weeks. Where there is seasonal transmission of the parasite, seasonal pneumonitis with eosinophilia may develop in previously infected and sensitized hosts.

In established infections, adult worms in the small intestine usually cause no symptoms. In heavy infections, particularly in children, a large bolus of entangled worms can cause pain and small-bowel obstruction, sometimes complicated by perforation, intussusception, or volvulus. Single worms may cause disease when they migrate into aberrant sites. A large worm can enter and occlude the biliary tree, causing biliary colic, cholecystitis, cholangitis, pancreatitis, or (rarely) intrahepatic abscesses. Migration of an adult worm up the esophagus can provoke coughing and oral expulsion of the worm. In highly endemic areas, intestinal and biliary ascariasis can rival acute appendicitis and gallstones as causes of surgical acute abdomen.

Laboratory Findings Most cases of ascariasis can be diagnosed by microscopic detection of characteristic *Ascaris* eggs (65 by 45 μm) in fecal samples. Occasionally, patients present after passing an adult worm—identifiable by its large size and smooth cream-colored surface—in the stool or, much less commonly, through the mouth or nose. During the early transpulmonary migratory phase, when eosinophilic pneumonitis occurs, larvae can be found in sputum or gastric aspirates before diagnostic eggs appear in the stool. The eosinophilia that is prominent during this early stage usually decreases to minimal levels in established infection. Adult worms may be visualized, occasionally serendipitously, on contrast studies of the gastrointestinal tract. A plain abdominal film may reveal masses of worms in gas-filled loops of bowel in patients with intestinal obstruction. Pancreaticobiliary worms can be detected by ultrasound and endoscopic retrograde cholangiopancreatography; the latter method also has been used to extract biliary *Ascaris* worms.

TREATMENT ASCARIASIS

Ascariasis should always be treated to prevent potentially serious complications. Albendazole (400 mg once), mebendazole (100 mg twice daily for 3 days or 500 mg once), or ivermectin (150–200 $\mu\text{g}/\text{kg}$

Feature	Parasitic Nematode				
	<i>Ascaris lumbricoides</i> (Roundworm)	<i>Necator americanus</i> , <i>Ancylostoma duodenale</i> (Hookworm)	<i>Strongyloides stercoralis</i>	<i>Trichuris trichiura</i> (Whipworm)	<i>Enterobius vermicularis</i> (Pinworm)
Global prevalence in humans (millions)	807	576	100	604	209
Endemic areas	Worldwide	Hot, humid regions	Hot, humid regions	Worldwide	Worldwide
Infective stage	Egg	Filariform larva	Filariform larva	Egg	Egg
Route of infection	Oral	Percutaneous	Percutaneous or autoinfection	Oral	Oral
Gastrointestinal location of worms	Jejunal lumen	Jejunal mucosa	Small-bowel mucosa	Cecum, colonic mucosa	Cecum, appendix
Adult worm size	15–40 cm	7–12 mm	2 mm	30–50 mm	8–13 mm (female)
Pulmonary passage of larvae	Yes	Yes	Yes	No	No
Incubation period ^a (days)	60–75	40–100	17–28	70–90	35–45
Longevity	1 year	<i>N. americanus</i> : 2–5 years <i>A. duodenale</i> : 6–8 years	Decades (owing to autoinfection)	5 years	2 months
Fecundity (eggs/day/worm)	240,000	<i>N. americanus</i> : 4000–10,000 <i>A. duodenale</i> : 10,000–25,000	5000–10,000	3000–7000	2000
Principal symptoms	Rarely, biliary obstruction or, in heavy infections, gastrointestinal obstruction	Iron-deficiency anemia in heavy infection	Gastrointestinal symptoms; malabsorption or sepsis in hyperinfection	Gastrointestinal symptoms or anemia in heavy infection	Perianal pruritus
Diagnostic stage	Eggs in stool	Eggs in fresh stool, larvae in old stool	Larvae in stool or duodenal aspirate; sputum in hyperinfection	Eggs in stool	Eggs from perianal skin on cellulose acetate tape
Treatment	Mebendazole Albendazole Ivermectin	Mebendazole Albendazole	Ivermectin Albendazole	Mebendazole Albendazole Ivermectin	Mebendazole Albendazole

^aTime from infection to egg production by mature female worm.

once) is effective. These medications are contraindicated in pregnancy, however. Mild diarrhea and abdominal pain are uncommon side effects of these agents. Partial intestinal obstruction should be managed with nasogastric suction, IV fluid administration, and instillation of piperazine through the nasogastric tube, but complete obstruction and its severe complications require immediate surgical intervention.

HOOKWORM

Two hookworm species (*A. duodenale* and *N. americanus*) are responsible for human infections. Most infected individuals are asymptomatic. Hookworm disease develops from a combination of factors—a heavy worm burden, a prolonged duration of infection, and an inadequate iron intake—and results in iron-deficiency anemia and, on occasion, hypoproteinemia.

Life Cycle Adult hookworms, which are ~1 cm long, use buccal teeth (*Ancylostoma*) or cutting plates (*Necator*) to attach to the small-bowel mucosa and suck blood (0.2 mL/d per *Ancylostoma* adult) and interstitial fluid. The adult hookworms produce thousands of eggs daily. The eggs are deposited with feces in soil, where rhabditiform larvae hatch and develop over a 1-week period into infectious filariform larvae. Infective larvae penetrate the skin and reach the lungs by way of the bloodstream. There they invade alveoli and ascend the airways before being swallowed and reaching the small intestine. The prepatent period from skin invasion to appearance of eggs in the feces is ~6–8 weeks, but it may be longer with *A. duodenale*. Larvae of *A. duodenale*, if swallowed, can survive and develop directly in the intestinal mucosa. Adult hookworms may survive over a decade but usually live ~6–8 years for *A. duodenale* and 2–5 years for *N. americanus*.



Epidemiology *A. duodenale* is prevalent in southern Europe, North Africa, and northern Asia, and *N. americanus* is the predominant species in the Western Hemisphere and equatorial Africa. The two species overlap in many tropical regions, particularly Southeast Asia. In most areas, older children have the highest incidence and greatest intensity of hookworm infection. In rural areas where fields are fertilized with human feces, older working adults also may be heavily infected.

Clinical Features Most hookworm infections are asymptomatic. Infective larvae may provoke pruritic maculopapular dermatitis (“ground itch”) at the site of skin penetration as well as serpiginous tracks of subcutaneous migration (similar to those of cutaneous larva migrans; [Chap. 256](#)) in previously sensitized hosts. Larvae migrating through the lungs occasionally cause mild transient pneumonitis, but this condition develops less frequently in hookworm infection than in ascariasis. In the early intestinal phase, infected persons may develop epigastric pain (often with postprandial accentuation), inflammatory diarrhea, or other abdominal symptoms accompanied by eosinophilia. The major consequence of chronic hookworm infection is iron deficiency. Symptoms are minimal if iron intake is adequate, but marginally nourished individuals develop symptoms of progressive iron-deficiency anemia and hypoproteinemia, including weakness and shortness of breath.

Laboratory Findings The diagnosis is established by the finding of characteristic 40- by 60- μ m oval hookworm eggs in the feces. Stool-concentration procedures may be required to detect light infections. Eggs of the two species are indistinguishable by light microscopy. In a stool sample that is not fresh, the eggs may have hatched to release rhabditiform larvae, which need to be differentiated from those of *S. stercoralis*. Hypochromic microcytic anemia, occasionally with eosinophilia or hypoalbuminemia, is characteristic of hookworm disease.

Hookworm infection can be eradicated with several safe and highly effective anthelmintic drugs, including albendazole (400 mg once) and mebendazole (500 mg once). Mild iron-deficiency anemia can often be treated with oral iron alone. Severe hookworm disease with protein loss and malabsorption necessitates nutritional support and oral iron replacement along with deworming. There is some concern that the benzimidazoles (mebendazole and albendazole) are becoming less effective against human hookworms.



Ancylostoma caninum* and *Ancylostoma braziliense *A. caninum*, the canine hookworm, has been identified as a cause of human eosinophilic enteritis, especially in northeastern Australia. In this zoonotic infection, adult hookworms attach to the small intestine (where they may be visualized by endoscopy) and elicit abdominal pain and intense local eosinophilia. Treatment with mebendazole (100 mg twice daily for 3 days) or albendazole (400 mg once) or endoscopic removal is effective. Both of these animal hookworm species can cause cutaneous larva migrans (“creeping eruption”; Chap. 256).

STRONGYLOIDIASIS

S. stercoralis is distinguished by its ability—unique among helminths (except for *Capillaria*; see below)—to replicate in the human host. This capacity permits ongoing cycles of autoinfection as infective larvae are internally produced. Strongyloidiasis can thus persist for decades without further exposure of the host to exogenous infective larvae. In immunocompromised hosts, large numbers of invasive *Strongyloides* larvae can disseminate widely and can be fatal.

Life Cycle In addition to a parasitic cycle of development, *Strongyloides* can undergo a free-living cycle of development in the soil (Fig. 257-1). This adaptability facilitates the parasite’s survival in the absence of mammalian hosts. Rhabditiform larvae passed in feces can transform into infectious filariform larvae either directly or after a free-living phase of development. Humans acquire strongyloidiasis when filariform larvae in fecally contaminated soil penetrate the skin or mucous membranes. The larvae then travel through the bloodstream to the lungs, where they break into the alveolar spaces, ascend the bronchial tree, are swallowed, and thereby reach the small intestine. There the larvae mature into adult worms that penetrate the mucosa of the proximal small bowel. The minute (2-mm-long) parasitic adult female worms reproduce by parthenogenesis; adult males do not exist. Eggs hatch in the intestinal mucosa, releasing rhabditiform larvae that migrate to the lumen and pass with the feces into soil. Alternatively, rhabditiform larvae in the bowel can develop directly into filariform larvae that penetrate the colonic wall or perianal skin and enter the circulation to repeat the migration that establishes ongoing internal reinfection. This autoinfection cycle allows strongyloidiasis to persist for decades.



Epidemiology *S. stercoralis* is spottily distributed in tropical areas and other hot, humid regions and is particularly common in Southeast Asia, sub-Saharan Africa, and Brazil. In the United States, the parasite is endemic in parts of the Southeast and is found in immigrants, refugees, travelers, and military personnel who have lived in endemic areas.

Clinical Features In uncomplicated strongyloidiasis, many patients are asymptomatic or have mild

cutaneous and/or abdominal symptoms. Recurrent urticaria, often involving the buttocks and wrists, is the most common cutaneous manifestation. Migrating larvae can elicit a pathognomonic serpiginous eruption, *larva currens* (“running larva”). This pruritic, raised, erythematous lesion advances as rapidly as 10 cm/h along the course of larval migration. Adult parasites burrow into the duodenojejunal mucosa and can cause abdominal (usually midepigastic) pain, which resembles peptic ulcer pain except that it is aggravated by food ingestion. Nausea, diarrhea, gastrointestinal bleeding, mild chronic colitis, and weight loss can occur. Small-bowel obstruction may develop with early, heavy infection. Pulmonary symptoms are rare in uncomplicated strongyloidiasis. Eosinophilia is common, with levels fluctuating over time.

The ongoing autoinfection cycle of strongyloidiasis is normally constrained by unknown factors of the host’s immune system. Abrogation of host immunity, especially with glucocorticoid therapy and much less commonly with other immunosuppressive medications, leads to hyperinfection, with the generation of large numbers of filariform larvae. Colitis, enteritis, or malabsorption may develop. In disseminated strongyloidiasis, larvae may invade not only gastrointestinal tissues and the lungs but also the central nervous system, peritoneum, liver, and kidneys. Moreover, bacteremia may develop because of the passage of enteric flora through disrupted mucosal barriers. Gram-negative sepsis, pneumonia, or meningitis may complicate or dominate the clinical course. Eosinophilia is often absent in severely infected patients. Disseminated strongyloidiasis, particularly in patients with unsuspected infection who are given glucocorticoids, can be fatal. Strongyloidiasis is a frequent complication of infection with human T cell lymphotropic virus type 1, but disseminated strongyloidiasis is not common among patients infected with HIV-1.

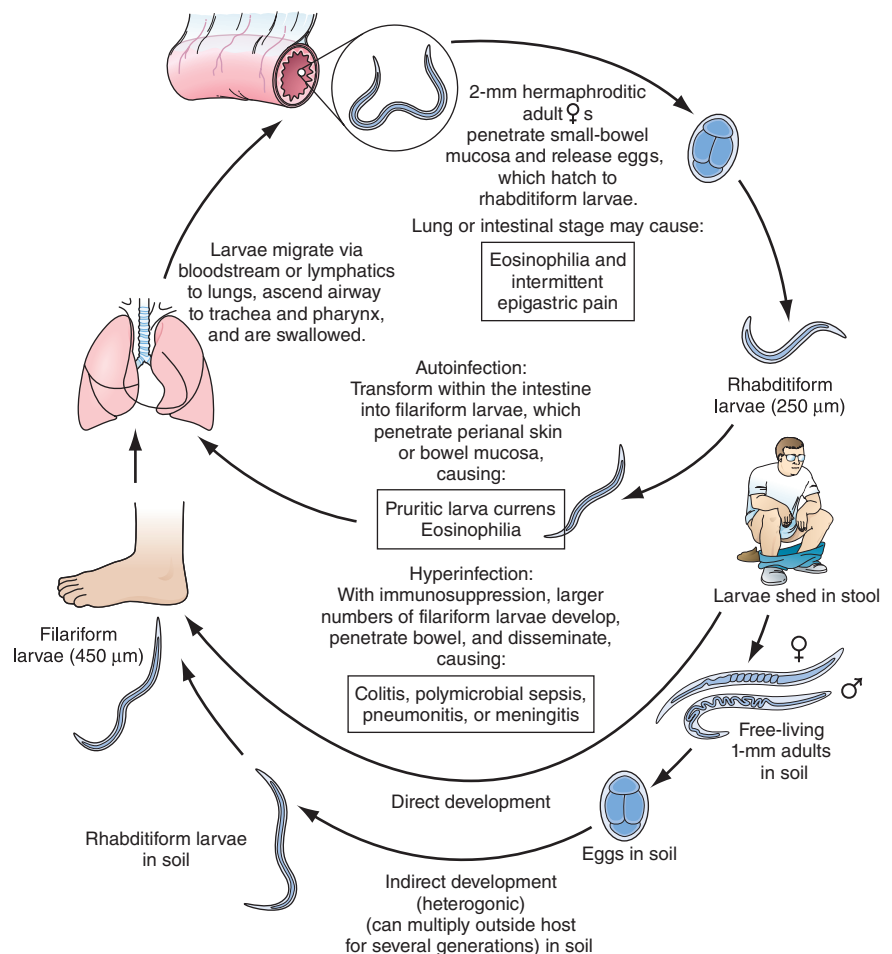


FIGURE 257-1 Life cycle of *Strongyloides stercoralis*. (Adapted from Guerrant RL et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 1276. © 2006, with permission from Elsevier Science.)

1416 Diagnosis In uncomplicated strongyloidiasis, the finding of rhabditiform larvae in feces is diagnostic. Rhabditiform larvae are ~250 µm long, with a short buccal cavity that distinguishes them from hookworm larvae. In uncomplicated infections, few larvae are passed and single stool examinations detect only about one-third of cases. Serial examinations and the use of the agar plate detection method improve the sensitivity of stool diagnosis. In uncomplicated strongyloidiasis (but not in hyperinfection), stool examinations may be repeatedly negative. *Strongyloides* larvae may also be found by sampling of the duodenojejunal contents by aspiration or biopsy. An enzyme-linked immunosorbent assay for serum antibodies to antigens of *Strongyloides* is a sensitive method for diagnosing uncomplicated infections. Such serologic testing should be performed for patients whose geographic histories indicate potential exposure, especially those who exhibit eosinophilia and/or are candidates for glucocorticoid treatment of other conditions. In disseminated strongyloidiasis, filariform larvae should be sought in stool as well as in samples obtained from sites of potential larval migration, including sputum, bronchoalveolar lavage fluid, or surgical drainage fluid.

TREATMENT STRONGYLOIDIASIS

Even in the asymptomatic state, strongyloidiasis must be treated because of the potential for subsequent dissemination and fatal hyperinfection. Ivermectin (200 µg/kg daily for 2 days) is consistently more effective than albendazole (400 mg daily for 3 days). For disseminated strongyloidiasis, treatment with ivermectin should be extended for at least 5–7 days or until the parasites have been eradicated. In immunocompromised hosts, the course of ivermectin should be repeated 2 weeks after initial treatment.

TRICHURIASIS



Most infections with *Trichuris trichiura* are asymptomatic, but heavy infections may cause gastrointestinal symptoms. Like the other soil-transmitted helminths, whipworm is distributed globally in the tropics and subtropics and is most common among poor children from resource-poor regions of the world.

Life Cycle Adult *Trichuris* worms reside in the colon and cecum, the anterior portions threaded into the superficial mucosa. Thousands of eggs laid daily by adult female worms pass with the feces and mature in the soil. After ingestion, infective eggs hatch in the duodenum, releasing larvae that mature before migrating to the large bowel. The entire cycle takes ~3 months, and adult worms may live for several years.

Clinical Features Tissue reactions to *Trichuris* are mild. Most infected individuals have no symptoms or eosinophilia. Heavy infections may result in anemia, abdominal pain, anorexia, and bloody or mucoid diarrhea resembling inflammatory bowel disease. Rectal prolapse can result from massive infections in children, who often suffer from malnourishment and other diarrheal illnesses. Moderately heavy *Trichuris* burdens also contribute to growth retardation.

Diagnosis and Treatment The characteristic 50- by 20-µm lemon-shaped *Trichuris* eggs are readily detected on stool examination. Adult worms, which are 3–5 cm long, are occasionally seen on proctoscopy. Mebendazole (500 mg once) or albendazole (400 mg daily for 3 doses) is safe and moderately effective for treatment, with cure rates of 70–90%. Ivermectin (200 µg/kg daily for 3 doses) is also safe but is not quite as efficacious as the benzimidazoles.

ENTEROBIASIS (PINWORM)



E. vermicularis is more common in temperate countries than in the tropics. In the United States, ~40 million persons are infected with pinworms, with a disproportionate number of cases among children.

Life Cycle and Epidemiology *Enterobius* adult worms are ~1 cm long and dwell in the cecum. Gravid female worms migrate nocturnally into the perianal region and release up to 2000 immature eggs each. The eggs become infective within hours and are transmitted by hand-to-mouth

passage. From ingested eggs, larvae hatch and mature into adults. This life cycle takes ~1 month, and adult worms survive for ~2 months. Self-infection results from perianal scratching and transport of infective eggs on the hands or under the nails to the mouth. Because of the ease of person-to-person spread, pinworm infections are common among family members.

Clinical Features Most pinworm infections are asymptomatic. Perianal pruritus is the cardinal symptom. The itching, which is often worse at night as a result of the nocturnal migration of the female worms, may lead to excoriation and bacterial superinfection. Heavy infections have been alleged to cause abdominal pain and weight loss. On rare occasions, pinworms invade the female genital tract, causing vulvovaginitis and pelvic or peritoneal granulomas. Eosinophilia is uncommon.

Diagnosis Since pinworm eggs are not released in feces, the diagnosis cannot be made by conventional fecal ova and parasite tests. Instead, eggs are detected by the application of clear cellulose acetate tape to the perianal region in the morning. After the tape is transferred to a slide, microscopic examination will detect pinworm eggs, which are oval, measure 55 by 25 µm, and are flattened along one side.

TREATMENT ENTEROBIASIS

Infected children and adults should be treated with mebendazole (100 mg once) or albendazole (400 mg once), with the same treatment repeated after 2 weeks. Treatment of household members is advocated to eliminate asymptomatic reservoirs of potential reinfection.

TRICHOSTRONGYLIASIS



Trichostrongylus species, which are normally parasites of herbivorous animals, occasionally infect humans, particularly in Asia and Africa. Humans acquire the infection by accidentally ingesting *Trichostrongylus* larvae on contaminated leafy vegetables. The larvae do not migrate in humans but mature directly into adult worms in the small bowel. These worms ingest far less blood than hookworms; most infected persons are asymptomatic, but heavy infections may give rise to mild anemia and eosinophilia. In stool examinations, *Trichostrongylus* eggs resemble hookworm eggs but are larger (85 by 115 µm). Treatment consists of mebendazole or albendazole (**Chap. 246e**).

ANISAKIASIS



Anisakiasis is a gastrointestinal infection caused by the accidental ingestion in uncooked saltwater fish of nematode larvae belonging to the family Anisakidae. The incidence of anisakiasis in the United States has increased as a result of the growing popularity of raw fish dishes. Most cases occur in Japan, the Netherlands, and Chile, where raw fish—sashimi, pickled green herring, and ceviche, respectively—are national culinary staples. Anisakid nematodes parasitize large sea mammals such as whales, dolphins, and seals. As part of a complex parasitic life cycle involving marine food chains, infectious larvae migrate to the musculature of a variety of fish. Both *Anisakis simplex* and *Pseudoterranova decipiens* have been implicated in human anisakiasis, but an identical gastric syndrome may be caused by the red larvae of eustrongylid parasites of fish-eating birds.

When humans consume infected raw fish, live larvae may be coughed up within 48 h. Alternatively, larvae may immediately penetrate the mucosa of the stomach. Within hours, violent upper abdominal pain accompanied by nausea and occasionally vomiting ensues, mimicking an acute abdomen. The diagnosis can be established by direct visualization on upper endoscopy, outlining of the worm by contrast radiographic studies, or histopathologic examination of extracted tissue. Extraction of the burrowing larvae during endoscopy is curative. In addition, larvae may pass to the small bowel, where they penetrate the mucosa and provoke a vigorous eosinophilic granulomatous response. Symptoms may appear 1–2 weeks after the infective meal, with intermittent abdominal pain, diarrhea, nausea, and fever resembling the manifestations of Crohn's disease. The diagnosis may be suggested by barium studies and confirmed by curative

surgical resection of a granuloma in which the worm is embedded. Anisakid eggs are not found in the stool, since the larvae do not mature in humans. Serologic tests have been developed but are not widely available.

Anisakid larvae in saltwater fish are killed by cooking to 60°C, freezing at -20°C for 3 days, or commercial blast freezing, but usually not by salting, marinating, or cold smoking. No medical treatment is available; surgical or endoscopic removal should be undertaken.

CAPILLARIASIS



Intestinal capillariasis is caused by ingestion of raw fish infected with *Capillaria philippinensis*. Subsequent autoinfection can lead to a severe wasting syndrome. The disease occurs in the Philippines and Thailand and, on occasion, elsewhere in Asia. The natural cycle of *C. philippinensis* involves fish from fresh and brackish water. When humans eat infected raw fish, the larvae mature in the intestine into adult worms, which produce invasive larvae that cause intestinal inflammation and villus loss. Capillariasis has an insidious onset with nonspecific abdominal pain and watery diarrhea. If untreated, progressive autoinfection can lead to protein-losing enteropathy, severe malabsorption, and ultimately death from cachexia, cardiac failure, or superinfection. The diagnosis is established by identification of the characteristic peanut-shaped (20- by 40- μ m) eggs on stool examination. Severely ill patients require hospitalization and supportive therapy in addition to prolonged anthelmintic treatment with albendazole (200 mg twice daily for 10 days; [Chap. 246e](#)).

ABDOMINAL ANGIOSTRONGYLIASIS

Abdominal angiostrongyliasis is found in Latin America and Africa. The zoonotic parasite *Angiostrongylus costaricensis* causes eosinophilic ileocolitis after the ingestion of contaminated vegetation. *A. costaricensis* normally parasitizes the cotton rat and other rodents, with slugs and snails serving as intermediate hosts. Humans become infected by accidentally ingesting infective larvae in mollusk slime deposited on fruits and vegetables; children are at highest risk. The larvae penetrate the gut wall and migrate to the mesenteric artery, where they develop into adult worms. Eggs deposited in the gut wall provoke an intense eosinophilic granulomatous reaction, and adult worms may cause mesenteric arteritis, thrombosis, or frank bowel infarction. Symptoms may mimic those of appendicitis, including abdominal pain and tenderness, fever, vomiting, and a palpable mass in the right iliac fossa. Leukocytosis and eosinophilia are prominent. CT with contrast medium typically shows inflamed bowel, often with concomitant obstruction, but a definitive diagnosis is usually made surgically with partial bowel resection. Pathologic study reveals a thickened bowel wall with eosinophilic granulomas surrounding the *Angiostrongylus* eggs. In nonsurgical cases, the diagnosis rests solely on clinical grounds because larvae and eggs cannot be detected in the stool. Medical therapy for abdominal angiostrongyliasis is of uncertain efficacy. Careful observation and surgical resection for severe symptoms are the mainstays of treatment.

subcutaneous tissues of humans. The offspring of adults are microfilariae, which, depending on their species, are 200–250 μ m long and 5–7 μ m wide, may or may not be enveloped in a loose sheath, and either circulate in the blood or migrate through the skin (Table 258-1). To complete the life cycle, microfilariae are ingested by the arthropod vector and develop over 1–2 weeks into new infective larvae. Adult worms live for many years, whereas microfilariae survive for 3–36 months. The bacterial endosymbiont *Wolbachia* has been found intracellularly in all stages of *Brugia*, *Wuchereria*, *Mansonella*, and *Onchocerca* species and has become a target for antifilarial chemotherapy.

Usually, infection is established only with repeated, prolonged exposures to infective larvae. Since the clinical manifestations of filarial diseases develop relatively slowly, these infections should be considered to induce chronic infections with possible long-term debilitating effects. In terms of the nature, severity, and timing of clinical manifestations, patients with filarial infections who are native to endemic areas and have lifelong exposure may differ significantly from those who are travelers or who have recently moved to these areas. Characteristically, filarial disease is more acute and intense in newly exposed individuals than in natives of endemic areas.

LYMPHATIC FILARIASIS

Lymphatic filariasis is caused by *W. bancrofti*, *B. malayi*, or *B. timori*. The threadlike adult parasites reside in lymphatic channels or lymph nodes, where they may remain viable for more than two decades.

EPIDEMIOLOGY



W. bancrofti, the most widely distributed filarial parasite of humans, affects an estimated 110 million people and is found throughout the tropics and subtropics, including Asia and the Pacific Islands, Africa, areas of South America, and the Caribbean basin. Humans are the only definitive host for the parasite. Generally, the subperiodic form is found only in the Pacific Islands; elsewhere, *W. bancrofti* is nocturnally periodic. Nocturnally periodic forms of microfilariae are scarce in peripheral blood by day and increase at night, whereas subperiodic forms are present in peripheral blood at all times and reach maximal levels in the afternoon. Natural vectors for *W. bancrofti* are *Culex fatigans* mosquitoes in urban settings and *Anopheles* or *Aedes* mosquitoes in rural areas.

Brugian filariasis due to *B. malayi* occurs primarily in eastern India, Indonesia, Malaysia, and the Philippines. *B. malayi* also has two forms distinguished by the periodicity of microfilaremia. The more common nocturnal form is transmitted in areas of coastal rice fields, while the subperiodic form is found in forests. *B. malayi* naturally infects cats as well as humans. The distribution of *B. timori* is limited to the islands of southeastern Indonesia.

PATHOLOGY

The principal pathologic changes result from inflammatory damage to the lymphatics, which is typically caused by adult worms and not by microfilariae. Adult worms live in afferent lymphatics or sinuses of lymph nodes and cause lymphatic dilation and thickening of the vessel walls. The infiltration of plasma cells, eosinophils, and macrophages in and around the infected vessels, along with endothelial and connective tissue proliferation, leads to tortuosity of the lymphatics and damaged or incompetent lymph valves. Lymphedema and chronic stasis changes with hard or brawny edema develop in the overlying skin. These consequences of filarial infection are due both to the direct effects of the worms and to the host's inflammatory response to the parasite. Inflammatory responses are believed to cause the granulomatous and proliferative processes that precede total lymphatic obstruction. It is thought that the lymphatic vessel remains patent as long as the worm remains viable and that the death of the worm leads to enhanced granulomatous reactions and fibrosis. Lymphatic obstruction results, and, despite collateralization, lymphatic function is compromised.

CLINICAL FEATURES

The most common presentations of the lymphatic filariases are asymptomatic (or subclinical) microfilaremia, hydrocele ([Fig. 258-1](#)), acute

258 Filarial and Related Infections

Thomas B. Nutman, Peter F. Weller

Filarial worms are nematodes that dwell in the subcutaneous tissues and the lymphatics. Eight filarial species infect humans ([Table 258-1](#)); of these, four—*Wuchereria bancrofti*, *Brugia malayi*, *Onchocerca volvulus*, and *Loa loa*—are responsible for most serious filarial infections. Filarial parasites, which infect an estimated 170 million persons worldwide, are transmitted by specific species of mosquitoes or other arthropods and have a complex life cycle, including infective larval stages carried by insects and adult worms that reside in either lymphatic or

TABLE 258-1 CHARACTERISTICS OF THE FILARIAE

Organism	Periodicity	Distribution	Vector	Location of Adult	Microfilarial Location	Sheath
<i>Wuchereria bancrofti</i>	Nocturnal	Cosmopolitan areas worldwide, including South America, Africa, southern Asia, Papua New Guinea, China, Indonesia	<i>Culex, Anopheles</i> (mosquitoes)	Lymphatic tissue	Blood	+
	Subperiodic	Eastern Pacific	<i>Aedes</i> (mosquitoes)	Lymphatic tissue	Blood	+
<i>Brugia malayi</i>	Nocturnal	Southeast Asia, Indonesia, India	<i>Mansonia, Anopheles</i> (mosquitoes)	Lymphatic tissue	Blood	+
	Subperiodic	Indonesia, Southeast Asia	<i>Coquillettidia, Mansonia</i> (mosquitoes)	Lymphatic tissue	Blood	+
<i>B. timori</i>	Nocturnal	Indonesia	<i>Anopheles</i> (mosquitoes)	Lymphatic tissue	Blood	+
<i>Loa loa</i>	Diurnal	West and Central Africa	<i>Chrysops</i> (deerflies)	Subcutaneous tissue	Blood	+
<i>Onchocerca volvulus</i>	None	South and Central America, Africa	<i>Simulium</i> (blackflies)	Subcutaneous tissue	Skin, eye	–
<i>Mansonella ozzardi</i>	None	South and Central America	<i>Culicoides</i> (midges)	Undetermined site	Blood	–
		Caribbean	<i>Simulium</i> (blackflies)			
<i>M. perstans</i>	None	South and Central America, Africa	<i>Culicoides</i> (midges)	Body cavities, mesentery, perirenal tissue	Blood	–
<i>M. streptocerca</i>	None	West and Central Africa	<i>Culicoides</i> (midges)	Subcutaneous tissue	Skin	–

adenolymphangitis (ADL), and chronic lymphatic disease. In areas where *W. bancrofti* or *B. malayi* is endemic, the overwhelming majority of infected individuals have few overt clinical manifestations of filarial infection despite large numbers of circulating microfilariae in the peripheral blood. Although they may be clinically asymptomatic, virtually all persons with *W. bancrofti* or *B. malayi* microfilaremia have some degree of subclinical disease that includes microscopic hematuria and/or proteinuria, dilated (and tortuous) lymphatics (visualized by imaging), and—in men with *W. bancrofti* infection—scrotal lymphangiectasia (detectable by ultrasound). Despite these findings, the majority of individuals appear to remain clinically asymptomatic for years; in relatively few does the infection progress to either acute or chronic disease.

ADL is characterized by high fever, lymphatic inflammation (lymphangitis and lymphadenitis), and transient local edema. The lymphangitis is retrograde, extending peripherally from the lymph node

draining the area where the adult parasites reside. Regional lymph nodes are often enlarged, and the entire lymphatic channel can become indurated and inflamed. Concomitant local thrombophlebitis can occur as well. In brugian filariasis, a single local abscess may form along the involved lymphatic tract and subsequently rupture to the surface. The lymphadenitis and lymphangitis can involve both the upper and lower extremities in both bancroftian and brugian filariasis, but involvement of the genital lymphatics occurs almost exclusively with *W. bancrofti* infection. This genital involvement can be manifested by funiculitis, epididymitis, and scrotal pain and tenderness. In endemic areas, another type of acute disease—dermatolymphangioadenitis (DLA)—is recognized as a syndrome that includes high fever, chills, myalgias, and headache. Edematous inflammatory plaques clearly demarcated from normal skin are seen. Vesicles, ulcers, and hyperpigmentation may also be noted. There is often a history of trauma, burns, irradiation, insect bites, punctiform lesions, or chemical injury. Entry lesions, especially in the interdigital area, are common. DLA is often diagnosed as cellulitis.

If lymphatic damage progresses, transient lymphedema can develop into lymphatic obstruction and the permanent changes associated with elephantiasis (Fig. 258-2). Brawny edema follows early pitting edema, the subcutaneous tissues thicken, and hyperkeratosis occurs. Fissuring of the skin develops, as do hyperplastic changes. Superinfection of these poorly vascularized tissues becomes a problem. In bancroftian filariasis, in which genital involvement is common, hydroceles may develop (Fig. 258-1); in advanced stages, this condition may evolve into scrotal lymphedema and scrotal elephantiasis. Furthermore, if there is obstruction of the retroperitoneal lymphatics, increased renal lymphatic pressure leads to rupture of the renal lymphatics and the development of chyluria, which is usually intermittent and most prominent in the morning.

The clinical manifestations of filarial infections in travelers or transmigrants who have recently entered an endemic region are distinctive. Given a sufficient number of bites by infected vectors, usually over a 3- to 6-month period, recently exposed patients can develop acute lymphatic or scrotal inflammation with or without urticaria and localized angioedema. Lymphadenitis of epitrochlear, axillary, femoral, or inguinal lymph nodes is often followed by retrogradely evolving lymphangitis. Acute attacks are short-lived and are not usually accompanied by fever. With prolonged exposure to infected mosquitoes, these attacks, if untreated, become more severe and lead to permanent lymphatic inflammation and obstruction.

DIAGNOSIS

A definitive diagnosis can be made only by detection of the parasites and hence can be difficult. Adult worms localized in lymphatic



FIGURE 258-1 Hydrocele associated with *Wuchereria bancrofti* infection.



FIGURE 258-2 Elephantiasis of the lower extremity associated with *Wuchereria bancrofti* infection.

vessels or nodes are largely inaccessible. Microfilariae can be found in blood, in hydrocele fluid, or (occasionally) in other body fluids. Such fluids can be examined microscopically, either directly or—for greater sensitivity—after concentration of the parasites by the passage of fluid through a polycarbonate cylindrical-pore filter (pore size, 3 μm) or by the centrifugation of fluid fixed in 2% formalin (Knott's concentration technique). The timing of blood collection is critical and should be based on the periodicity of the microfilariae in the endemic region involved. Many infected individuals do not have microfilaremia, and definitive diagnosis in such cases can be difficult. Assays for circulating antigens of *W. bancrofti* permit the diagnosis of microfilaremic and cryptic (amicrofilaremic) infection. Two tests are commercially available: an enzyme-linked immunosorbent assay (ELISA) and a rapid-format immunochromatographic card test. Both assays have sensitivities of 93–100% and specificities approaching 100%. There are currently no tests for circulating antigens in brugian filariasis.

Polymerase chain reaction (PCR)-based assays for DNA of *W. bancrofti* and *B. malayi* in blood have been developed. A number of studies indicate that the sensitivity of this diagnostic method is equivalent to or greater than that of parasitologic methods.

In cases of suspected lymphatic filariasis, examination of the scrotum, the lymph nodes, or (in female patients) the breast by means of high-frequency ultrasound in conjunction with Doppler techniques may result in the identification of motile adult worms within dilated lymphatics. Worms may be visualized in the lymphatics of the spermatic cord in up to 80% of men infected with *W. bancrofti*. Live adult worms have a distinctive pattern of movement within the lymphatic vessels (termed the *filarial dance sign*). Radionuclide lymphoscintigraphic imaging of the limbs reliably demonstrates widespread lymphatic abnormalities in both subclinical microfilaremic persons and those with clinical manifestations of lymphatic pathology. Although of potential utility in the delineation of anatomic changes associated with infection, lymphoscintigraphy is unlikely to assume primacy in the diagnostic evaluation of individuals with suspected infection; it is principally a research tool, although it has been used more widely for assessment of lymphedema of any cause. Eosinophilia and elevated serum concentrations of IgE and antifilarial antibody

support the diagnosis of lymphatic filariasis. There is, however, extensive cross-reactivity between filarial antigens and antigens of other helminths, including the common intestinal roundworms; thus, interpretations of serologic findings can be difficult. In addition, residents of endemic areas can become sensitized to filarial antigens (and thus be serologically positive) through exposure to infected mosquitoes without having patent filarial infections.

The ADL associated with lymphatic filariasis must be distinguished from thrombophlebitis, infection, and trauma. Retrograde evolution is a characteristic feature that helps distinguish filarial lymphangitis from ascending bacterial lymphangitis. Chronic filarial lymphedema must also be distinguished from the lymphedema of malignancy, postoperative scarring, trauma, chronic edematous states, and congenital lymphatic system abnormalities.

TREATMENT LYMPHATIC FILARIASIS

With newer definitions of clinical syndromes in lymphatic filariasis and new tools to assess clinical status (e.g., ultrasound, lymphoscintigraphy, circulating filarial antigen assays, PCR), approaches to treatment based on infection status can be considered.

Orally administered diethylcarbamazine (DEC; 6 mg/kg daily for 12 days), which has both macro- and microfilaricidal properties, remains the drug of choice for the treatment of active lymphatic filariasis (defined by microfilaremia, antigen positivity, or adult worms on ultrasound), although albendazole (400 mg twice daily by mouth for 21 days) has also demonstrated macrofilaricidal efficacy. A 4- to 6-week course of oral doxycycline (targeting the intracellular *Wolbachia*) also has significant macrofilaricidal activity, as does DEC/albendazole used daily for 7 days. The addition of DEC to a 3-week course of doxycycline is efficacious in lymphatic filariasis.

Regimens that combine single doses of albendazole (400 mg) with either DEC (6 mg/kg) or ivermectin (200 $\mu\text{g}/\text{kg}$) all have a sustained microfilaricidal effect and are the mainstay of programs for the eradication of lymphatic filariasis in Africa (albendazole/ivermectin) and elsewhere (albendazole/DEC) (see "Prevention and Control," below).

As has already been mentioned, a growing body of evidence indicates that, although they may be asymptomatic, virtually all persons with *W. bancrofti* or *B. malayi* microfilaremia have some degree of subclinical disease (hematuria, proteinuria, abnormalities on lymphoscintigraphy). Thus, early treatment of asymptomatic persons who have microfilaremia is recommended to prevent further lymphatic damage. For ADL, supportive treatment (including the administration of antipyretics and analgesics) is recommended, as is antibiotic therapy if secondary bacterial infection is likely. Similarly, because lymphatic disease is associated with the presence of adult worms, treatment with DEC is recommended for microfilaria-negative carriers of adult worms.


In persons with chronic manifestations of lymphatic filariasis, treatment regimens that emphasize hygiene, prevention of secondary bacterial infections, and physiotherapy have gained wide acceptance for morbidity control. These regimens are similar to those recommended for lymphedema of most nonfilarial causes and are known by a variety of names, including *complex decongestive physiotherapy* and *complex lymphedema therapy*. Hydroceles (Fig. 258-1) can be managed surgically. With chronic manifestations of lymphatic filariasis, drug treatment should be reserved for individuals who have evidence of active infection; however, a 6-week course of doxycycline has been shown to provide improvement in filarial lymphedema irrespective of disease activity.

Side effects of DEC treatment include fever, chills, arthralgias, headaches, nausea, and vomiting. Both the development and the severity of these reactions are directly related to the number of microfilariae circulating in the bloodstream. The adverse reactions may represent either an acute hypersensitivity reaction to the antigens being released by dead and dying parasites or an inflammatory reaction induced by the intracellular *Wolbachia* endosymbionts freed from their intracellular niche.


1420 Ivermectin has a side effect profile similar to that of DEC when used in lymphatic filariasis. In patients infected with *L. loa* who have high levels of microfilaremia, DEC—like ivermectin (see “Loiasis,” below)—can elicit severe encephalopathic complications. When used in single-dose regimens for the treatment of lymphatic filariasis, albendazole is associated with relatively few side effects.

PREVENTION AND CONTROL

To protect themselves against filarial infection, individuals must avoid contact with infected mosquitoes by using personal protective measures, including bed nets, particularly those impregnated with insecticides such as permethrin. Community-based intervention is the current approach to elimination of lymphatic filariasis as a public health problem. The underlying tenet of this approach is that mass annual distribution of antimicrofilarial chemotherapy—albendazole with either DEC (for all areas except those where onchocerciasis is coendemic; see section on onchocerciasis treatment, below) or ivermectin—will profoundly suppress microfilaremia. If the suppression is sustained, then transmission can be interrupted.

 Created by the World Health Organization in 1997, the Global Programme to Eliminate Lymphatic Filariasis is based on mass administration of single annual doses of DEC plus albendazole in non-African regions and of albendazole plus ivermectin in Africa. Available information from late 2013 indicated that more than 792 million persons in 53 countries had thus far participated. Not only has lymphatic filariasis been eliminated in some defined areas, but collateral benefits—avoidance of disability and treatment of intestinal helminths and other conditions (e.g., scabies and louse infestation)—have also been noted. The strategy of the global program is being refined, and attempts are being made to integrate this effort with other mass-treatment strategies (e.g., deworming programs, malaria control, and trachoma control) in an integrated control strategy.

TROPICAL PULMONARY EOSINOPHILIA

 Tropical pulmonary eosinophilia (TPE) is a distinct syndrome that develops in some individuals infected with the lymphatic-dwelling filarial species. This syndrome affects males and females in a ratio of 4:1, often during the third decade of life. The majority of cases have been reported from India, Pakistan, Sri Lanka, Brazil, Guyana, and Southeast Asia.

Clinical Features The main features include a history of residence in filarial-endemic regions, paroxysmal cough and wheezing (usually nocturnal and probably related to the nocturnal periodicity of microfilariae), weight loss, low-grade fever, lymphadenopathy, and pronounced blood eosinophilia (>3000 eosinophils/ μ L). Chest x-rays or CT scans may be normal but generally show increased bronchovascular markings. Diffuse miliary lesions or mottled opacities may be present in the middle and lower lung fields. Tests of pulmonary function show restrictive abnormalities in most cases and obstructive defects in half. Characteristically, total serum IgE levels (4–40 KIU/mL) and antifilarial antibody titers are markedly elevated.

Pathology In TPE, microfilariae and parasite antigens are rapidly cleared from the bloodstream by the lungs. The clinical symptoms result from allergic and inflammatory reactions elicited by the cleared parasites. In some patients, trapping of microfilariae in other reticuloendothelial organs can cause hepatomegaly, splenomegaly, or lymphadenopathy. A prominent, eosinophil-enriched, intraalveolar infiltrate is often reported, and with it comes the release of cytotoxic proinflammatory eosinophil granule proteins that may mediate some of the pathology seen in TPE. In the absence of successful treatment, interstitial fibrosis can lead to progressive pulmonary damage.


Differential Diagnosis TPE must be distinguished from asthma, Löffler’s syndrome, allergic bronchopulmonary aspergillosis, allergic granulomatosis with angiitis (Churg-Strauss syndrome), the systemic vasculitides (most notably, periarteritis nodosa and granulomatosis with polyangiitis), chronic eosinophilic pneumonia, and the idiopathic hypereosinophilic syndrome.

TREATMENT TROPICAL PULMONARY EOSINOPHILIA

DEC is used at a daily dosage of 4–6 mg/kg for 14 days. Symptoms usually resolve within 3–7 days after the initiation of therapy. Relapse, which occurs in ~12–25% of cases (sometimes after an interval of years), requires re-treatment.

ONCHOCERCIASIS

EPIDEMIOLOGY

 Onchocerciasis (“river blindness”) is caused by the filarial nematode *O. volvulus*, which infects an estimated 37 million individuals in 35 countries worldwide. The majority of individuals infected with *O. volvulus* live in the equatorial region of Africa extending from the Atlantic coast to the Red Sea. In the Americas, isolated foci were identified in Mexico, Guatemala, Colombia, Ecuador, Venezuela, and Brazil. The infection is also found in Yemen.

ETIOLOGY

Infection in humans begins with the deposition of infective larvae on the skin by the bite of an infected blackfly. The larvae develop into adults, which are typically found in subcutaneous nodules. About 7 months to 3 years after infection, the gravid female releases microfilariae that migrate out of the nodule and throughout the tissues, concentrating in the dermis. Infection is transmitted to other persons when a female fly ingests microfilariae from the host’s skin and these microfilariae then develop into infective larvae. Adult *O. volvulus* females and males are ~40–60 cm and ~3–6 cm in length, respectively. The life span of adults can be as long as 18 years, with an average of ~9 years. Because the blackfly vector breeds along free-flowing rivers and streams (particularly in rapids) and generally restricts its flight to an area within several kilometers of these breeding sites, both biting and disease transmission are most intense in these locations.

PATHOLOGY

Onchocerciasis primarily affects the skin, eyes, and lymph nodes. In contrast to the pathology in lymphatic filariasis, the damage in onchocerciasis is elicited by microfilariae and not by adult parasites. In the skin, there are mild but chronic inflammatory changes that can result in loss of elastic fibers, atrophy, and fibrosis. The subcutaneous nodules (*onchocercomata*) consist primarily of fibrous tissues surrounding the adult worm, often with a peripheral ring of inflammatory cells (characterized as lymphatic in origin) surrounded by an endothelial layer. In the eye, neovascularization and corneal scarring lead to corneal opacities and blindness. Inflammation in the anterior and posterior chambers frequently results in anterior uveitis, chorioretinitis, and optic atrophy. Although punctate opacities are due to an inflammatory reaction surrounding dead or dying microfilariae, the pathogenesis of most manifestations of onchocerciasis is still unclear.

CLINICAL FEATURES

Skin Pruritus and rash are the most common manifestations of onchocerciasis. The pruritus can be incapacitating; the rash is typically a papular eruption (Fig. 258-3) that is generalized rather than localized to a particular region of the body. Long-term infection results in exaggerated and premature wrinkling of the skin, loss of elastic fibers, and epidermal atrophy that can lead to loose, redundant skin and hypo- or hyperpigmentation. Localized eczematoid dermatitis can cause hyperkeratosis, scaling, and pigmentary changes. In an immunologically hyperreactive form of onchodermatitis (commonly termed *sowdah* or *localized onchodermatitis*), the affected skin darkens as a consequence of the profound inflammation that occurs as microfilariae in the skin are cleared.



 **Onchocercomata** These subcutaneous nodules, which can be palpable and/or visible, contain the adult worm. In African patients, they are common over the coccyx and sacrum, the trochanter of the femur, the lateral anterior crest, and other bony prominences; in patients from South and Central America, nodules



FIGURE 258-3 Papular eruption as a consequence of onchocerciasis.

tend to develop preferentially in the upper part of the body, particularly on the head, neck, and shoulders. Nodules vary in size and characteristically are firm and not tender. It has been estimated that, for every palpable nodule, there are four deeper nonpalpable ones.

Ocular Tissue Visual impairment is the most serious complication of onchocerciasis and usually affects only those persons with moderate or heavy infections. Lesions may develop in all parts of the eye. The most common early finding is conjunctivitis with photophobia. Punctate keratitis—acute inflammatory reactions surrounding dying microfilariae and manifested as “snowflake” opacities—is common among younger patients and resolves without apparent complications.

 Sclerosing keratitis occurs in 1–5% of infected persons and is the leading cause of onchocercal blindness in Africa. Anterior uveitis and iridocyclitis develop in ~5% of infected persons in Africa. In Latin America, complications of the anterior uveal tract (pupillary deformity) may cause secondary glaucoma. Characteristic chorioretinal lesions develop as a result of atrophy and hyperpigmentation of the retinal pigment epithelium. Constriction of the visual fields and overt optic atrophy may occur.

Lymph Nodes Mild to moderate lymphadenopathy is common, particularly in the inguinal and femoral areas, where the enlarged nodes may hang down in response to gravity (“hanging groin”), sometimes predisposing to inguinal and femoral hernias.

Systemic Manifestations Some heavily infected individuals develop cachexia with loss of adipose tissue and muscle mass. Among adults who become blind, there is a three- to fourfold increase in the mortality rate.

DIAGNOSIS

Definitive diagnosis depends on the detection of an adult worm in an excised nodule or, more commonly, of microfilariae in a skin snip. Skin snips are obtained with a corneal-scleral punch, which collects a blood-free skin biopsy sample extending to just below the epidermis, or by lifting of the skin with the tip of a needle and excision of a small (1- to 3-mm) piece with a sterile scalpel blade. The biopsy tissue is incubated in tissue culture medium or in saline on a glass slide or flat-bottomed microtiter plate. After incubation for 2–4 h (or occasionally overnight in light infections), microfilariae emergent from the skin can be seen by low-power microscopy.

Eosinophilia and elevated serum IgE levels are common but, because they occur in many parasitic infections, are not diagnostic

in themselves. Assays to detect specific antibodies to *Onchocerca* and PCR to detect onchocercal DNA in skin snips are used in specialized laboratories and are highly sensitive and specific.

TREATMENT ONCHOCERCIASIS

The main goals of therapy are to prevent the development of irreversible lesions and to alleviate symptoms. Surgical excision is recommended when nodules are located on the head (because of the proximity of microfilaria-producing adult worms to the eye), but chemotherapy is the mainstay of management. Ivermectin, a semisynthetic macrocyclic lactone active against microfilariae, is the first-line agent for the treatment of onchocerciasis. It is given orally in a single dose of 150 µg/kg, either yearly or semiannually. More frequent ivermectin administration (every 3 months) has been suggested to ameliorate pruritus and skin disease.



After treatment, most individuals have few or no reactions. Pruritus, cutaneous edema, and/or maculopapular rash occurs in ~1–10% of treated individuals. In areas of Africa coendemic for *O. volvulus* and *L. loa*, however, ivermectin is contraindicated (as it is for pregnant or breast-feeding women) because of severe posttreatment encephalopathy, especially in patients who are heavily microfilaremic for *L. loa* (>8000 microfilariae/mL). Although ivermectin treatment results in a marked drop in microfilarial density, its effect can be short-lived (<3 months in some cases). Thus, it is occasionally necessary to give ivermectin more frequently for persistent symptoms.

A 6-week course of doxycycline is macrofilaristatic, rendering female adult worms sterile for long periods.

PREVENTION

Vector control has been beneficial in highly endemic areas in which breeding sites are vulnerable to insecticide spraying, but most areas endemic for onchocerciasis are not suited to this type of control. Community-based administration of ivermectin every 6–12 months is being used to interrupt transmission in endemic areas. This measure, in conjunction with vector control, has already helped eliminate the infection in most of Latin America and has reduced the prevalence of disease in many endemic foci in Africa. No drug has proved useful for prophylaxis of *O. volvulus* infection.

LOIASIS

ETIOLOGY AND EPIDEMIOLOGY

Loiasis is caused by *L. loa* (the African eye worm), which is present in the rainforests of West and Central Africa. Adult parasites (females, 50–70 mm long and 0.5 mm wide; males, 25–35 mm long and 0.25 mm wide) live in subcutaneous tissues. Microfilariae circulate in the blood with a diurnal periodicity that peaks between 12:00 noon and 2:00 P.M.

CLINICAL FEATURES

Manifestations of loiasis in natives of endemic areas may differ from those in temporary residents or visitors. Among the indigenous population, loiasis is often an asymptomatic infection with microfilaremia. Infection may be recognized only after subconjunctival migration of an adult worm (Fig. 258-4) or may be manifested by episodic Calabar swellings—evanescent localized areas of angioedema and erythema developing on the extremities and less frequently at other sites. Nephropathy, encephalopathy, and cardiomyopathy can occur but are rare. In patients who are not residents of endemic areas, allergic symptoms predominate, episodes of Calabar swelling tend to be more frequent and debilitating, microfilaremia is less common, and eosinophilia and increased levels of antifilarial antibodies are characteristic.

PATHOLOGY

The pathogenesis of the manifestations of loiasis is poorly understood. Calabar swellings are thought to result from a hypersensitivity reaction to adult worm antigens.



FIGURE 258-4 Adult *Loa loa* worm being surgically removed after its subconjunctival migration.

DIAGNOSIS

Definitive diagnosis of loiasis requires the detection of microfilariae in the peripheral blood or the isolation of the adult worm from the eye (Fig. 258-4) or from a subcutaneous biopsy specimen collected from a site of swelling developing after treatment. PCR-based assays for the detection of *L. loa* DNA in blood are available in specialized laboratories and are highly sensitive and specific, as are some newer recombinant antigen-based serologic techniques. In practice, the diagnosis must often be based on a characteristic history and clinical presentation, blood eosinophilia, and elevated levels of antifilarial antibodies, particularly in travelers to an endemic region, who are usually amicrofilaremic. Other clinical findings in travelers include hypergammaglobulinemia, elevated levels of serum IgE, and elevated leukocyte and eosinophil counts.

TREATMENT LOIASIS

DEC (8–10 mg/kg per day administered orally for 21 days) is effective against both the adult and the microfilarial forms of *L. loa*, but multiple courses are frequently necessary before loiasis resolves completely. In cases of heavy microfilaremia, allergic or other inflammatory reactions can take place during treatment, including central nervous system involvement with coma and encephalitis. Heavy infections can be treated initially with apheresis to remove the microfilariae and with glucocorticoids (40–60 mg of prednisone per day) followed by doses of DEC (0.5 mg/kg per day). If antifilarial treatment has no adverse effects, the prednisone dose can be rapidly tapered and the dose of DEC gradually increased to 8–10 mg/kg per day.

Albendazole or ivermectin is effective in reducing microfilarial loads, although neither is approved for this purpose by the U.S. Food and Drug Administration. Moreover, ivermectin is contraindicated in patients with >8000 microfilariae/mL because this drug has been associated with severe adverse events (including encephalopathy and death) in heavily infected patients with loiasis in West and Central Africa. DEC (300 mg weekly) is an effective prophylactic regimen for loiasis.

STREPTOCERCIASIS



Mansonella streptocerca, found mainly in the tropical forest belt of Africa from Ghana to the Democratic Republic of the Congo, is transmitted by biting midges. The major clinical manifestations involve the skin and include pruritus, papular rashes, and pigmentation changes. Many infected individuals have inguinal adenopathy, although most are asymptomatic. The diagnosis is made by

detection of the characteristic microfilariae in skin snips. Ivermectin at a single dose of 150 µg/kg leads to sustained suppression of microfilariae in the skin and is probably the treatment of choice for streptocerciasis.

MANSONELLA PERSTANS INFECTION



M. perstans, distributed across the center of Africa and in northeastern South America, is transmitted by midges. Adult worms reside in serous cavities—pericardial, pleural, and peritoneal—as well as in the mesentery and the perirenal and retroperitoneal tissues. Microfilariae circulate in the blood without periodicity. The clinical and pathologic features of the infection are poorly defined. Most patients appear to be asymptomatic, but manifestations may include transient angioedema and pruritus of the arms, face, or other parts of the body (analogous to the Calabar swellings of loiasis); fever; headache; arthralgias; and right-upper-quadrant pain. Occasionally, pericarditis and hepatitis occur. The diagnosis is based on the demonstration of microfilariae in blood or serosal effusions. Perstans filariasis is often associated with peripheral-blood eosinophilia and antifilarial antibody elevations.

With the identification of a *Wolbachia* endosymbiont in *M. perstans*, doxycycline (200 mg twice a day) for 6 weeks has been established as the first effective treatment for this infection.

MANSONELLA OZZARDI INFECTION



The distribution of *M. ozzardi* is restricted to Central and South America and certain Caribbean islands. Adult worms are rarely recovered from humans. Microfilariae circulate in the blood without periodicity. Although this organism has often been considered nonpathogenic, headache, articular pain, fever, pulmonary symptoms, adenopathy, hepatomegaly, pruritus, and eosinophilia have been ascribed to *M. ozzardi* infection. The diagnosis is made by detection of microfilariae in peripheral blood. Ivermectin is effective in treating this infection.

DRACUNCULIASIS (GUINEA WORM INFECTION)

ETIOLOGY AND EPIDEMIOLOGY



The incidence of dracunculiasis, caused by *Dracunculus medinensis*, has declined dramatically because of global eradication efforts. In 2012, only 542 cases worldwide had been identified. The infection is currently endemic only in Chad, Ethiopia, Mali, and South Sudan.

Humans acquire *D. medinensis* when they ingest water containing infective larvae derived from *Cyclops*, a crustacean that is the intermediate host. Larvae penetrate the stomach or intestinal wall, mate, and mature. The adult male probably dies; the female worm develops over a year and migrates to subcutaneous tissues, usually in the lower extremity. As the thin female worm, ranging in length from 30 cm to 1 m, approaches the skin, a blister forms that, over days, breaks down and forms an ulcer. When the blister opens, large numbers of motile, rhabditiform larvae can be released into stagnant water; ingestion by *Cyclops* completes the life cycle.

CLINICAL FEATURES

Few or no clinical manifestations of dracunculiasis are evident until just before the blister forms, when there is an onset of fever and generalized allergic symptoms, including periorbital edema, wheezing, and urticaria. The emergence of the worm is associated with local pain and swelling. When the blister ruptures (usually as a result of immersion in water) and the adult worm releases larva-rich fluid, symptoms are relieved. The shallow ulcer surrounding the emerging adult worm heals over weeks to months. Such ulcers, however, can become secondarily infected, the result being cellulitis, local inflammation, abscess formation, or (uncommonly) tetanus. Occasionally, the adult worm does not emerge but becomes encapsulated and calcified.

DIAGNOSIS

The diagnosis is based on the findings developing with the emergence of the adult worm, as described above.

TREATMENT DRACUNCULIASIS

Gradual extraction of the worm by winding of a few centimeters on a stick each day remains the common and effective practice. Worms may be excised surgically. No drug is effective in treating dracunculiasis.

PREVENTION

Prevention, which remains the only real control measure, depends on the provision of safe drinking water.

ZOONOTIC FILARIAL INFECTIONS

Dirofilariae that affect primarily dogs, cats, and raccoons occasionally infect humans incidentally, as do *Brugia* and *Onchocerca* parasites that affect small mammals. Because humans are an abnormal host, the parasites never develop fully. Pulmonary dirofilarial infection caused by the canine heartworm *Dirofilaria immitis* generally presents in humans as a solitary pulmonary nodule. Chest pain, hemoptysis, and cough are uncommon. Infections with *D. repens* (from dogs) or *D. tenuis* (from raccoons) can cause local subcutaneous nodules in humans. Zoonotic *Brugia* infection can produce isolated lymph node enlargement, whereas zoonotic *Onchocerca* can cause subconjunctival masses. Eosinophilia levels and antifilarial antibody titers are not commonly elevated. Excisional biopsy is both diagnostic and curative. These infections usually do not respond to chemotherapy.

259 Schistosomiasis and Other Trematode Infections

Charles H. King, Adel A. F. Mahmoud

Trematodes, or flatworms, are a group of morphologically and biologically heterogeneous organisms that belong to the phylum Platyhelminthes. Human infection with trematodes occurs in many geographic areas and can cause considerable morbidity and mortality. The dependence on one drug—praziquantel—for treatment of most infections caused by trematodes raises the specter of developing resistance in these worms; several instances of reduced drug efficacy have already been reported. The widespread use of oxamniquine in the 1970s to reduce the impact of schistosomiasis resulted in the development of significant resistance. Recently, a single quantitative trait locus on schistosomal chromosome 6 was identified as the genetic basis for resistance.

ETIOLOGIC AGENTS AND THEIR LIFE CYCLES

For clinical purposes, significant trematode infections of humans may be divided according to the tissues invaded by the adult stage of the fluke, whether bloodstream, biliary tree, intestines, or lungs (Table 259-1). Trematodes share some common morphologic features, including macroscopic size (from one to several centimeters); dorsoventral, flattened, bilaterally symmetric bodies (adult worms); and the prominence of two suckers. Except for schistosomes, all human parasitic trematodes are hermaphroditic. Their life cycles involve a definitive host (mammalian/human), in which adult worms initiate sexual reproduction, and an intermediate host (snail), in which asexual multiplication of larvae occurs. More than one intermediate host may be necessary for some species of trematodes. Human infection is initiated either by direct penetration of intact skin or by ingestion. Upon maturation within humans, adult flukes initiate sexual reproduction and egg production. Helminth ova leave the definitive host in excreta or sputum and, upon reaching suitable environmental

TABLE 259-1 MAJOR HUMAN TREMATODE INFECTIONS

Trematode	Transmission	Endemic Area(s)
Blood Flukes		
<i>Schistosoma mansoni</i>	Skin penetration by cercariae released from snails	Africa, South America, Middle East
<i>S. japonicum</i>	Skin penetration by cercariae released from snails	China, Philippines, Indonesia
<i>S. intercalatum</i>	Skin penetration by cercariae released from snails	West Africa
<i>S. mekongi</i>	Skin penetration by cercariae released from snails	Southeast Asia
<i>S. haematobium</i>	Skin penetration by cercariae released from snails	Africa, Middle East
Biliary (Hepatic) Flukes		
<i>Clonorchis sinensis</i>	Ingestion of metacercariae in freshwater fish	Eastern Asia
<i>Opisthorchis viverrini</i>	Ingestion of metacercariae in freshwater fish	Eastern Asia, Thailand
<i>O. felinus</i>	Ingestion of metacercariae in freshwater fish	Eastern Asia, Europe
<i>Fasciola hepatica</i>	Ingestion of metacercariae on aquatic plants or in water	Worldwide
<i>F. gigantica</i>	Ingestion of metacercariae on aquatic plants or in water	Sporadic, Africa
Intestinal Flukes		
<i>Fasciolopsis buski</i>	Ingestion of metacercariae on aquatic plants	Southeast Asia
<i>Heterophyes heterophyes</i>	Ingestion of metacercariae in freshwater or brackish-water fish	Eastern Asia, North Africa
Lung Flukes		
<i>Paragonimus westermani</i> and related species	Ingestion of metacercariae in crayfish or crabs	Global except North America and Europe

conditions, they hatch, releasing free-living miracidia that seek specific snail intermediate hosts. After asexual reproduction, cercariae are released from infected snails. In certain species, these organisms infect humans; in others, they find a second intermediate host to allow encystment into metacercariae—the infective stage for humans.

The host-parasite relationship in trematode infections is a product of certain biologic features of these organisms: they are multicellular, undergo several developmental changes within the host, and usually result in chronic infections. In general, the distribution of worm infections in human populations is *overdispersed*; i.e., it follows a negative binomial statistical distribution in which most infected individuals harbor low worm burdens while a small percentage are heavily infected. It is the heavily infected minority who are particularly prone to disease sequelae and who constitute an epidemiologically significant reservoir of infection in endemic areas. Recent evidence indicates that the prevalence of morbidity in infected populations is greater than was previously thought. Morbidity and death due to trematode infections reflect a multifactorial process that results from the tipping of a delicate balance between intensity of infection and host reactions, which initiate and modulate immunologic and pathologic outcome. Furthermore, the genetics of the parasite and of the human host contribute to the outcome of infection and disease. Infections with trematodes that migrate through or reside in host tissues are associated with a moderate to high degree of peripheral-blood eosinophilia; this association is of significance in protective and immunopathologic sequelae and is a useful clinical indicator of infection.

APPROACH TO THE PATIENT: Trematode Infection

The approach to individuals with suspected trematode infection begins with a question: Where have you been? Details of geographic history, exposure to freshwater bodies, and indulgence in local eating habits (without ensuring safety of food and drink) are all essential elements in eliciting the history of the present illness. The workup plan must include a detailed physical examination and tests appropriate for suspected infection. Diagnosis is based either on detection of the relevant stage of the parasite in excreta, sputum, or (rarely) tissue samples or on sensitive and specific serologic tests. Consultation with physicians familiar with these infections or with the U.S. Centers for Disease Control and Prevention (CDC) is helpful in guiding diagnosis and selecting therapy.

GLOBAL CONSIDERATIONS: EPIDEMIOLOGY OF TREMATODE INFECTIONS



Except among international travelers, trematode infections are quite rare in high-income countries because good sanitation and hygiene block trematode transmission and because transmission is tied to the distribution of the specific snail species that serve as intermediate hosts during the parasites' life cycle. In contrast, parasitic fluke infections are quite common in underdeveloped areas of Africa, Asia, and South America, with an estimated 440 million people affected by past or present *Schistosoma* infection and another 60 million people affected by the other foodborne trematodes. These infections are not benign; they result in multiyear chronic inflammatory disorders that significantly affect performance status and health-related quality of life. Global disease burden estimates indicate that at least 5 million years of healthy life are lost each year in the more than 90 endemic countries around the world.

BLOOD FLUKES: SCHISTOSOMIASIS

Human schistosomiasis is caused by five species of the parasitic trematode genus *Schistosoma*: *S. mansoni*, *S. japonicum*, *S. mekongi*, and *S. intercalatum* cause intestinal and hepatic schistosomiasis, and *S. haematobium* causes urogenital schistosomiasis. Infection may cause considerable morbidity in the intestines, liver, or urinary tract, and a small proportion of affected individuals die. Other schistosomes (e.g., avian species) may invade human skin but then die in subcutaneous tissue, producing only self-limiting cutaneous manifestations.

ETIOLOGY

Human infection is initiated by penetration of intact skin with infective cercariae. These organisms, which are released from infected snails in freshwater bodies, measure ~2 mm in length and possess an anterior and a ventral sucker that attach to the skin and facilitate penetration. Once in subcutaneous tissue, cercariae transform into schistosomula, with morphologic, membrane, and immunologic changes. The cercarial outer membrane changes from a trilaminar to a heptalaminar structure that is then maintained throughout the organism's life span in humans. This transformation is thought to be the schistosome's main adaptive mechanism for survival in humans. Schistosomula begin their migration within 2–4 days via venous or lymphatic vessels, reaching the lungs and finally the liver parenchyma. Sexually mature worms descend into the venous system at specific anatomic locations: intestinal veins (*S. mansoni*, *S. japonicum*, *S. mekongi*, and *S. intercalatum*) and vesical and other pelvic veins (*S. haematobium*). After mating, adult gravid females travel against venous blood flow to small tributaries, where they deposit their ova intravascularly. Schistosome ova (Fig. 259-1) have specific morphologic features that vary with the species. Aided by enzymatic secretions through minipores in eggshells, ova move through the venous wall, traversing host tissues to reach the lumen of the intestinal or urinary tract, and are voided with stools or urine. Approximately 50% of ova are retained in host tissues locally (intestines or urinary tract) or are carried by venous blood flow to the liver and other organs. Schistosome ova that reach freshwater bodies hatch, releasing free-living miracidia that seek the snail intermediate host and undergo several cycles of asexual multiplication. Finally, infective cercariae are shed from snails to complete the transmission cycle.

Adult schistosomes are ~1–2 cm long. Males are slightly shorter than females, with flattened bodies and anteriorly curved edges forming the gynecophoral canal, in which mature adult females are usually held. Females are longer, slender, and rounded in cross-section. The precise nature of biochemical and reproductive exchanges between the two sexes is unknown, as are the regulatory mechanisms for pairing. Adult schistosomes parasitize specific sites in the host venous system. What guides adult intestinal schistosomes to branches of the superior or inferior mesenteric veins or adult *S. haematobium* worms to the vesical plexus is unknown. In addition, adult worms inhibit the coagulation cascade and evade the effector arms of the host immune responses by still-undetermined mechanisms. The genome of schistosomes is relatively large (~270 Mb) and is arrayed on seven pairs of autosomes and one pair of sex chromosomes. Sequencing of the *S. japonicum*, *S. mansoni*, and *S. haematobium* genomes has provided insight into the worms' genomic and proteomic features, offering an opportunity to

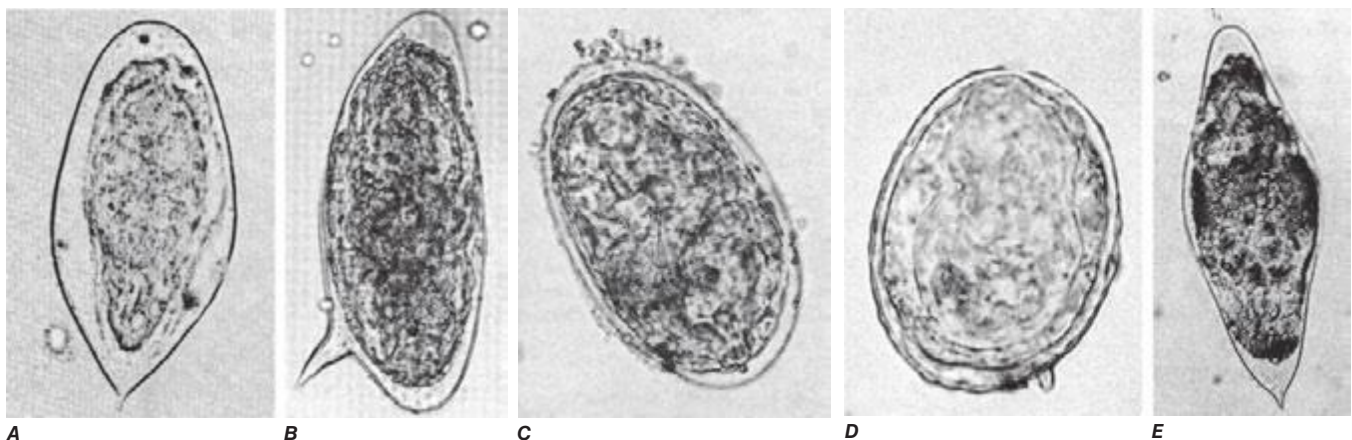


FIGURE 259-1 Morphology of schistosome eggs, the diagnostic stage of the parasite's life cycle. **A.** *Schistosoma haematobium* egg (in a urine sample) is large (~140 μ m long), with a terminal spine. **B.** *S. mansoni* egg (in a fecal sample) is large (~150 μ m long), with a thin shell and lateral spine. **C.** *S. japonicum* egg (fecal) is smaller than that of *S. mansoni* (~90 μ m long), with a small spine or hooklike structure. **D.** *S. mekongi* egg (fecal) is similar to that of *S. japonicum* but smaller (~65 μ m long). **E.** *S. intercalatum* egg (fecal) is larger than that of *S. haematobium* (~190 μ m long), with a longer, sharply pointed spine. (From LR Ash, TC Orihel: *Atlas of Human Parasitology*, 3rd ed. Chicago, ASCP Press, 1990; with permission.)

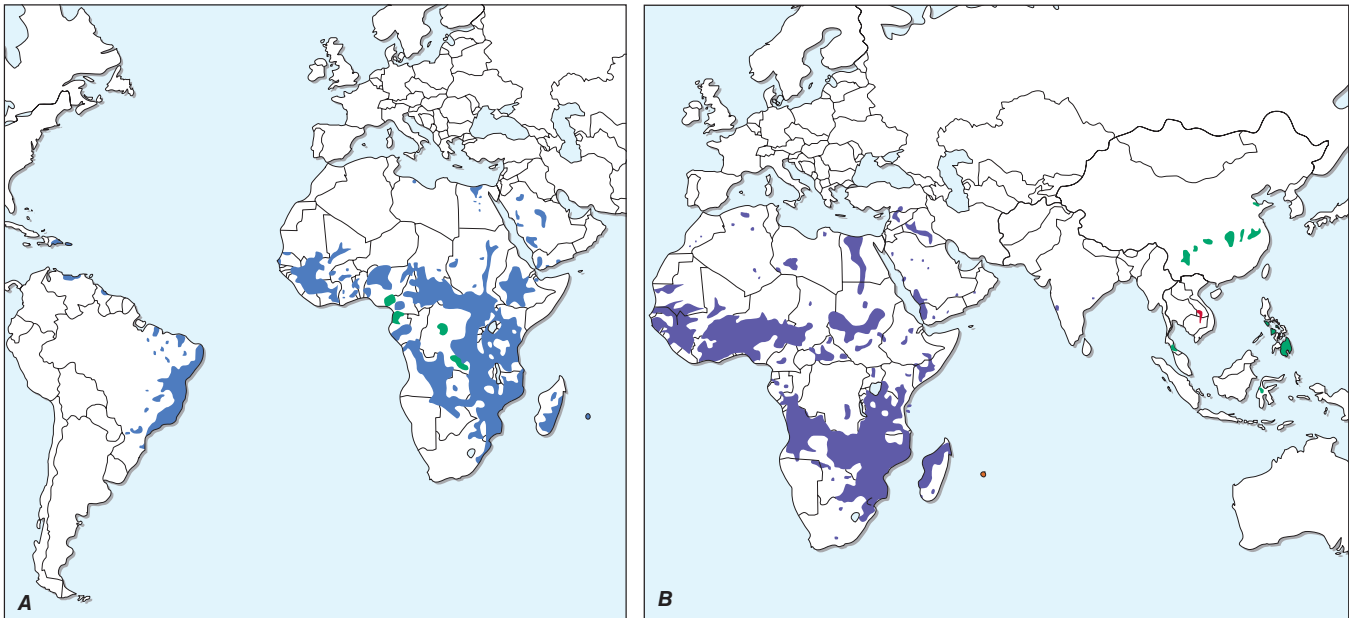


FIGURE 259-2 Global distribution of schistosomiasis. **A.** *Schistosoma mansoni* infection (dark blue) is endemic in Africa, the Middle East, South America, and a few Caribbean countries. *S. intercalatum* infection (green) is endemic in sporadic foci in West and Central Africa. **B.** *S. haematobium* infection (purple) is endemic in Africa and the Middle East. The major endemic countries for *S. japonicum* infection (green) are China, the Philippines, and Indonesia. *S. mekongi* infection (red) is endemic in sporadic foci in Southeast Asia.

discover new drug targets and to understand the molecular basis of pathogenesis.

EPIDEMIOLOGY



The global distribution of schistosome infection in human populations (Fig. 259-2) is dependent on both parasite and host factors. Information on prevalence and global distribution is inexact. At present, the five *Schistosoma* species are estimated to infect 200–300 million individuals (mostly children and young adults) in South America, the Caribbean, Africa, the Middle East, and Southeast Asia. Notably, parasite-related disease persists after active infection resolves, leaving a substantial health burden among adult populations. Thus, the overall number of humans likely to be affected by *Schistosoma*-related disease is now ~440 million. The total population living under conditions favoring transmission risk numbers ~700 million—a fact reflecting the global public health significance of schistosomiasis.

In endemic areas, the rate of yearly onset of new infection (incidence) is generally low. Prevalence, on the other hand, starts to be appreciable by the age of 3–4 years and builds to a maximum that varies by endemic region (up to 100%) in the 12- to 20-year age group. Prevalence then stabilizes or decreases slightly in older age groups (>40 years). Intensity of infection (as measured by fecal or urinary egg counts, which correlate with adult worm burdens in most circumstances) follows the increase in prevalence up to the age of 12–20 years and then declines markedly in older age groups. This decline may reflect acquisition of resistance or may be due to changes in water contact patterns, since older people have less exposure. Infection with schistosomes in human populations has a peculiar pattern. Most infected individuals harbor low worm burdens, and only a small proportion suffer from high-intensity infection. This pattern may be due to differences in worm infectivity or to a spectrum of genetic susceptibilities in human populations.

Disease due to schistosome infection is the consequence of parasitologic, host, and associated viral infections and of nutritional and environmental factors. Most disease syndromes relate to the presence of one or more of the parasite stages in humans. Disease manifestations in the populations of endemic areas correlate, in general, with intensity and duration of infection as well as with age and genetic susceptibility of the host. Overall, severe *Schistosoma*-specific disease

manifestations are relatively rare among persons infected with any of the intestinal schistosomes. In contrast, symptoms of urogenital schistosomiasis manifest clinically in most *S. haematobium*-infected individuals. In addition, all forms of *Schistosoma* infection are associated with subclinical systemic morbidities that can significantly affect physical and cognitive performance, causing, for example, growth stunting, undernutrition, and anemia of chronic inflammation. New estimates of total morbidity due to chronic schistosomiasis indicate a significantly greater burden than was previously appreciated.

Schistosomiasis appears to be a cofactor in the spread and progression of HIV/AIDS in areas where both diseases are endemic. Increased emphasis should be placed on the treatment of schistosome infections in persons at risk of HIV/AIDS.

PATHOGENESIS AND IMMUNITY

Cercarial invasion is associated with dermatitis arising from dermal and subdermal inflammatory responses, both humoral and cell-mediated. As the parasites approach sexual maturity in the liver of infected individuals and as oviposition commences, acute schistosomiasis or Katayama syndrome (a serum sickness–like illness; see “Clinical Features,” below) may occur. The associated antigen excess results in formation of soluble immune complexes, which may be deposited in several tissues, initiating multiple pathologic events. In chronic schistosomiasis, most disease manifestations are due to eggs retained in host tissues. The granulomatous response around these ova is cell-mediated and is regulated both positively and negatively by a cascade of cytokine, cellular, and humoral responses. Granuloma formation begins with recruitment of a host of inflammatory cells in response to antigens secreted by the living organism within the ova. Cells recruited initially include phagocytes, antigen-specific T cells, and eosinophils. Fibroblasts, giant cells, and B lymphocytes predominate later. Over time, these cumulative lesions reach a size many times that of parasite eggs, thus inducing organomegaly and obstruction. Immunomodulation or downregulation of host responses to schistosome eggs plays a significant role in limiting the extent of the granulomatous lesions—and consequently disease—in chronically infected experimental animals or humans. The underlying mechanisms involve another cascade of regulatory cytokines and idiotypic antibodies. Subsequent to the granulomatous response, fibrosis sets in, resulting in more permanent disease sequelae. Because schistosomiasis is also a



FIGURE 259-3 Chronic hepatosplenomegaly caused by schistosomiasis mansoni.

Liver and spleen enlargement, ascites, and wasting are characteristically seen in patients with chronic *Schistosoma mansoni* infection.

chronic infection, the accumulation of antigen–antibody complexes results in deposits in renal glomeruli and may cause significant kidney disease.

The better-studied pathologic sequelae in schistosomiasis are those observed in liver disease. Ova that are carried by portal blood embolize to the liver. Because of their size (~150 × 60 μm in the case of *S. mansoni*), they lodge at presinusoidal sites, where granulomas are formed. These granulomas contribute to the hepatomegaly observed in infected individuals (Fig. 259-3). Schistosomal liver enlargement is also associated with certain class I and class II human leukocyte antigen (HLA) haplotypes and markers; its genetic basis appears to be polygenic. Presinusoidal portal blockage causes several hemodynamic changes, including portal hypertension and associated development of portosystemic collaterals at the esophagogastric junction and other sites. Esophageal varices are most likely to break and cause repeated episodes of hematemesis. Because changes in hepatic portal blood flow occur slowly, compensatory arterialization of the blood flow through the liver is established. Although this compensatory mechanism may be associated with certain metabolic side effects, retention of hepatocyte perfusion permits maintenance of normal liver function for several years.

The second most significant pathologic change in the liver relates to fibrosis. It is characteristically periportal (Symmers' clay pipe–stem fibrosis) but may be diffuse. Fibrosis, when diffuse, may be seen in areas of egg deposition and granuloma formation but is also seen in distant locations such as portal tracts. Schistosomiasis results in pure fibrotic lesions in the liver; cirrhosis occurs only when other toxic factors or infectious agents (e.g., hepatitis B or C virus) are involved. Deposition of fibrotic tissue in the extracellular matrix results from the interaction of T lymphocytes with cells of the fibroblast series; several cytokines, such as interleukin (IL) 2, IL-4, IL-1, and transforming growth factor β, are known to stimulate fibrogenesis. The process may be dependent on the genetic constitution of the host. Furthermore, regulatory cytokines that can suppress T cell responses and fibrogenesis, such as IL-10, interferon γ, or IL-12, may play a role in modulating the response.

Although the above description focuses on granuloma formation and fibrosis of the liver, similar processes occur in urogenital schistosomiasis. Granuloma formation at the lower end of the ureters obstructs urinary flow, with subsequent development of hydroureter and hydronephrosis. Similar lesions in the urinary bladder cause the protrusion of papillomatous structures into its cavity; these may ulcerate and/or bleed. The chronic stage of infection is associated with scarring and deposition of calcium in the bladder wall. Among women,

involvement of the birth canal can cause cervical or vaginal wall polyps and friability leading to contact bleeding, with an apparently increased risk of HIV transmission. Secondary infertility or subfertility can also result from female genital schistosomiasis involving the uterus, fallopian tubes, or ovaries. Among men, *S. haematobium* infection can result in prostatic and testicular lesions with hematospermia. Superficial cutaneous lesions of the perineum can occur in both sexes.

Studies on immunity to schistosomiasis, whether innate or adaptive, have expanded our knowledge of the components of these responses and target antigens. The critical question, however, is whether humans acquire immunity to schistosomes. Epidemiologic data suggest the onset of acquired immunity during the course of infection in young adults. Curative treatment of infected populations in endemic areas is followed by differentiation in the pattern of reinfection. Some (susceptible) individuals acquire reinfection rapidly, whereas other (resistant) individuals are reinfected slowly. This difference may be explained by differences in transmission, immunologic response, or genetic susceptibility. The

mechanism of acquired immunity involves antibodies, complement, and several effector cells, particularly eosinophils. Furthermore, the intensity of schistosome infection has been correlated with a region in chromosome 5. In several studies, a few protective schistosome antigens have been identified as vaccine candidates, but none has been fully evaluated in human populations to date.

CLINICAL FEATURES

In general, disease manifestations of schistosomiasis occur in three stages, which vary not only by species but also by intensity of infection and other host factors, such as age and genetics of the human host. During the phase of cercarial invasion, a form of dermatitis may be observed. This so-called swimmers' itch occurs most often with *S. mansoni* and *S. japonicum* infections, manifesting 2 or 3 days after invasion as an itchy maculopapular rash on the affected areas of the skin. The condition is particularly severe when humans are exposed to avian schistosomes. This form of cercarial dermatitis is also seen around freshwater lakes in the northern United States, particularly in the spring and summer months. Cercarial dermatitis is a self-limiting clinical entity. During worm maturation and at the beginning of oviposition (i.e., 4–8 weeks after skin invasion), acute schistosomiasis or Katayama syndrome—a serum sickness–like illness with fever, generalized lymphadenopathy, and hepatosplenomegaly—may develop. Individuals with acute schistosomiasis have a high degree of peripheral-blood eosinophilia. Parasite-specific antibodies may be detected before schistosome eggs are identified in excreta.



Acute schistosomiasis has become an important clinical entity worldwide because of increased travel to endemic areas. Travelers are exposed to parasites while swimming or wading in freshwater bodies and upon their return present with acute manifestations. The course of acute schistosomiasis is generally benign, but central nervous system (CNS) schistosomiasis and even deaths are occasionally reported in association with heavy exposure to schistosomes among travelers and migrants.

The main clinical manifestations of chronic schistosomiasis are species-dependent. Intestinal species (*S. mansoni*, *S. japonicum*, *S. mekongi*, and *S. intercalatum*) cause intestinal and hepatosplenic disease as well as several manifestations associated with portal hypertension. During the intestinal phase, which may begin a few months after infection and may last for years, symptomatic patients characteristically have colicky abdominal pain, bloody diarrhea, and anemia. Patients may also report fatigue and an inability to perform daily routine functions and may show evidence of growth retardation and anemia. This more subtle form of schistosomiasis morbidity is generally

underappreciated. The severity of intestinal schistosomiasis is often related to the intensity of the worm burden. The disease runs a chronic course and may result in colonic polyposis, which has been reported from some endemic areas, such as Egypt and Uganda.

The hepatosplenic phase of disease manifests early (during the first year of infection, particularly in children) with liver enlargement due to parasite-induced granulomatous lesions. Hepatomegaly is seen in ~15–20% of infected individuals; it correlates roughly with intensity of infection, occurs more often in children, and may be related to specific HLA haplotypes. In subsequent phases of infection, presinusoidal blockage of blood flow leads to portal hypertension and splenomegaly (Fig. 259-3). Moreover, portal hypertension may lead to varices at the lower end of the esophagus and at other sites. Patients with schistosomiasis may have right-upper-quadrant “dragging” pain during the hepatomegaly phase, and this pain may move to the left upper quadrant as splenomegaly progresses. Bleeding from esophageal varices may, however, be the first clinical manifestation of this phase. Patients may experience repeated bleeding but seem to tolerate its impact, because an adequate total hepatic blood flow permits normal liver function for a considerable period. In late-stage disease, typical fibrotic changes occur along with liver function deterioration and the onset of ascites, hypoalbuminemia, and defects in coagulation. Intercurrent viral infections of the liver (especially hepatitis B and C), toxic insults (excessive ethanol ingestion or exposure to organic poisons or aflatoxin), or nutritional deficiencies may well accelerate or exacerbate the deterioration of hepatic function.

The extent and severity of intestinal and hepatic disease in schistosomiasis *mansoni* and *japonica* have been well described. Although it was originally thought that *S. japonicum* might induce more severe disease manifestations because the adult worms can produce 10 times more eggs than *S. mansoni*, subsequent field studies have not supported this claim. Clinical observations of individuals infected with *S. mekongi* or *S. intercalatum* have been less detailed, partly because of the limited geographic distribution of these organisms.

The clinical manifestations of *S. haematobium* infection occur relatively early and involve a high percentage of infected individuals. Up to 80% of children infected with *S. haematobium* have dysuria, frequency, and hematuria. Hematuria may sometimes occur only at the end of voiding. Urine examination reveals blood and albumin as well as an unusually high frequency of bacterial urinary tract infections and urinary sediment cellular metaplasia. These manifestations correlate with the intensity of infection, the presence of urinary bladder granulomas, and subsequent ulceration. Along with local effects of granuloma formation in the urinary bladder, obstruction of the lower end of the ureters results in hydronephrosis and hydronephrosis, which may be seen in 25–50% of infected children. As infection progresses, bladder granulomas undergo fibrosis, which results in typical sandy patches visible on cystoscopy. In many endemic areas, an association between squamous cell carcinoma of the bladder and *S. haematobium* infection has been observed. Such malignancy is detected in a younger age group than is transitional cell carcinoma. In fact, *S. haematobium* has now been classified as a human carcinogen. Genital schistosomiasis (described in the previous section) is a common presenting symptom among adults of both sexes.

Significant disease may occur in other organs during chronic schistosomiasis. Lung and CNS disease have been documented; other sites, such as the skin and the genital organs, are less frequently affected. In pulmonary schistosomiasis, embolized eggs lodge in small arterioles, producing acute necrotizing arteriolitis and granuloma formation. During *S. mansoni* and *S. japonicum* infection, schistosome eggs reach the lungs after the development of portosystemic collateral circulation; in *S. haematobium* infection, ova may reach the lungs directly via connections between the vesical and systemic circulation. Subsequent fibrous tissue deposition leads to endarteritis obliterans, pulmonary hypertension, and cor pulmonale. The most common symptoms are cough, fever, and dyspnea. Cor pulmonale may be diagnosed radiologically on the basis of prominence of the right side of the heart and dilation of the pulmonary artery. Frank evidence of right-sided heart failure may be seen in late cases.

Although less common than pulmonary manifestations, CNS schistosomiasis is important, characteristically occurring in association with *S. japonicum* infection. Migratory worms deposit eggs in the brain and induce a granulomatous response. The frequency of this manifestation among infected individuals in some endemic areas (e.g., the Philippines) is calculated at 2–4%. Jacksonian epilepsy due to *S. japonicum* infection is the second most common cause of epilepsy in these areas. *S. mansoni* and *S. haematobium* infections have been associated with transverse myelitis. This syndrome is thought to be due to eggs traveling to the venous plexus around the spinal cord. In schistosomiasis *mansoni*, transverse myelitis is usually seen in the chronic stage after the development of portal hypertension and portosystemic shunts, which allow ova to travel to the spinal cord veins. This proposed sequence of events has been challenged because of a few reports of transverse myelitis occurring early in the course of *S. mansoni* infection. More information is needed to confirm these observations. During schistosomiasis caused by *Schistosoma haematobium*, ova may travel through communication between vesical and systemic veins, resulting in spinal cord disease that may be detected at any stage of infection. Pathologic study of lesions in schistosomal transverse myelitis may reveal eggs along with necrotic or granulomatous lesions. Patients usually present with acute or rapidly progressing lower-leg weakness accompanied by sphincter dysfunction.

DIAGNOSIS

Physicians in areas not endemic for schistosomiasis face considerable diagnostic challenges. In the most common clinical presentation, a traveler returns with symptoms and signs of acute syndromes of schistosomiasis—namely, cercarial dermatitis or Katayama syndrome. Central to a correct diagnosis is a thorough inquiry into the patient's history of travel and exposure to freshwater bodies—whether slow- or fast-running—in an endemic area. Differential diagnosis of fever in returned travelers includes a spectrum of infections whose etiologies are viral (e.g., dengue fever), bacterial (e.g., enteric fever, leptospirosis), rickettsial, or protozoal (e.g., malaria). In cases of Katayama syndrome, prompt diagnosis is essential and is based on clinical presentation, high-level peripheral-blood eosinophilia, and a positive serologic assay for schistosomal antibodies. Two tests are available at the CDC: the Falcon assay screening test/enzyme-linked immunosorbent assay (FAST-ELISA) and the confirmatory enzyme-linked immunoelectrotransfer blot (EITB). Both tests are highly sensitive and ~96% specific. In some instances, examination of stool or urine for ova may yield positive results.


Individuals with established infection are diagnosed by a combination of geographic history, characteristic clinical presentation, and presence of schistosome ova in excreta. The diagnosis may also be established with the serologic assays mentioned above or with those that detect circulating schistosome antigens. These assays can be applied to blood, urine, or other body fluids (e.g., cerebrospinal fluid). For suspected schistosome infection, stool examination by the Kato thick smear or any other concentration method generally identifies most patients with heavy infection but does not identify all lightly infected individuals. For the latter patients, a point-of-care test to detect parasite circulating cathodic antigen in urine may prove very useful in establishing the presence of active *S. mansoni* infection and in monitoring the clearance of infection after treatment. For *S. haematobium*, urine may be examined by microscopy of sediment or by filtration of a known volume through Nuclepore filters. Sensitivity can be further improved by testing for parasite DNA in urine sediment. The Kato thick smear and Nuclepore filtration provide quantitative data on the intensity of infection, which is of value in assessing the degree of tissue damage and in monitoring the effect of chemotherapy. Schistosome infection may also be diagnosed by examination of tissue specimens, typically rectal biopsy samples; except in rare circumstances, other biopsy procedures (e.g., liver biopsy) are not needed.

The differential diagnosis of schistosomal hepatomegaly must include viral hepatitis of all etiologies, miliary tuberculosis, malaria, visceral leishmaniasis, ethanol abuse, and causes of hepatic and portal vein obstruction. The differential diagnosis of hematuria in *S. haematobium*

TREATMENT SCHISTOSOMIASIS


Treatment of schistosomiasis depends on the stage of infection and the clinical presentation. Other than topical dermatologic applications for relief of itching, no specific treatment is indicated for cercarial dermatitis caused by avian schistosomes. Therapy for acute schistosomiasis or Katayama syndrome needs to be adjusted appropriately for each case. Although antischistosomal chemotherapy may be used, it does not have a significant impact on maturing worms. In severe acute schistosomiasis, management in an acute-care setting is necessary, with supportive measures and consideration of glucocorticoid treatment to reduce inflammation. Once the acute critical phase is over, specific chemotherapy is indicated for parasite elimination. For all individuals with established infection, treatment to eradicate the parasite should be administered. The drug of choice is praziquantel, which—depending on the infecting species (Table 259-2)—is administered PO as a total of 40 or 60 mg/kg in two or three doses over a single day. Praziquantel treatment results in parasitologic cure in ~85% of cases and reduces egg counts by >90%. Efficacy rates among children <5 years old have been reported to be lower. These children are more likely to need re-treatment to effect a cure. Few side effects have been encountered, and those that do develop usually do not interfere with completion of treatment. Dependence on a single chemotherapeutic agent has raised the possibility of development of resistance in schistosomes; to date, such resistance does not seem to be clinically significant. The effect of antischistosomal treatment on disease manifestations varies by stage. Early hepatomegaly and bladder lesions are known to resolve after chemotherapy, but the late established manifestations, such as fibrosis, do not recede. Additional management modalities are needed for individuals with other manifestations, such as hepatocellular failure or recurrent hematemesis. The use of these interventions is guided by general medical and surgical principles.

PREVENTION AND CONTROL


 Transmission of schistosomiasis is dependent on human behavior. Because the geographic distribution of infections in endemic regions of the world is not clearly demarcated, it is prudent for travelers to endemic areas to avoid contact with *all* freshwater bodies, irrespective of the speed of water flow or unsubstantiated claims of safety. Some topical agents, when applied to the skin, may inhibit cercarial penetration, but none is currently available. If exposure occurs, a

follow-up visit with a health care provider is strongly recommended. Prevention of infection in inhabitants of endemic areas is a significant challenge. Residents of these regions use freshwater bodies for sanitary, domestic, recreational, and agricultural purposes. Several control measures have been used, including application of molluscicides, provision of sanitary water and sewage disposal, chemotherapy, and health education to effect behavioral change in terms of water-contact activities. Current recommendations to countries endemic for schistosomiasis emphasize the use of multiple approaches. With the advent of an oral, safe, and effective broad-spectrum antischistosomal agent (praziquantel), chemotherapy has been most successful in reducing the intensity of infection and reversing disease. The duration of this positive impact depends on the transmission dynamics of the parasite in any specific endemic region. The ultimate goal of research on prevention and control is the development of a vaccine. Although there are a few promising leads, this goal probably is not within reach during the next decade.

LIVER (BILIARY) FLUKES

 Several species of biliary fluke infecting humans are particularly common in Southeast Asia and Russia. Other species are transmitted in Europe, Africa, and the Americas. On the basis of their migratory pathway in humans, these infections may be divided into the *Clonorchis* and *Fasciola* groups (Table 259-1).


CLONORCHIASIS AND OPISTHORCHIASIS

 Infection with *Clonorchis sinensis*, the Chinese or oriental fluke, is endemic among fish-eating mammals in Southeast Asia. Humans are an incidental host; the prevalence of human infection is highest in China, Vietnam, and Korea. Infection with *Opisthorchis viverrini* and *O. felineus* is zoonotic in cats and dogs. Transmission to humans occurs occasionally, particularly in Thailand (*O. viverrini*) and in Southeast Asia and eastern Europe (*O. felineus*). Data on the exact geographic distribution of these infectious agents in human populations are rudimentary.

Infection with any of these three species is established by ingestion of raw or inadequately cooked freshwater fish harboring metacercariae. These organisms excyst in the duodenum, releasing larvae that travel through the ampulla of Vater and mature into adult worms in bile canaliculi. Mature flukes are flat and elongated, measuring 1–2 cm in length. The hermaphroditic worms reproduce by releasing small operculated eggs, which pass with bile into the intestines and are voided with stools. The life cycle is completed in the environment in specific freshwater snails (the first intermediate host) along with later encystment of snail-derived cercariae as infectious metacercariae in freshwater fish.

Except for late sequelae, the exact clinical syndromes caused by clonorchiasis and opisthorchiasis are not well defined. Because most infected individuals harbor a low worm burden, many are minimally symptomatic. Moderate to heavy infection may be associated with vague right-upper-quadrant pain. In contrast, chronic or repeated infection is associated with manifestations such as cholangitis, cholangiohepatitis, and biliary obstruction. Cholangiocarcinoma is epidemiologically related to *C. sinensis* infection in China and to *O. viverrini* infection in northeastern Thailand. This association has resulted in classification of these infectious agents as human carcinogens.

FASCIOLIASIS

 Infections with *Fasciola hepatica* and *F. gigantica* are worldwide zoonoses that are particularly endemic in sheep-raising countries. Human cases have been reported in South America, Europe, Africa, and Australia. Recent estimates indicate a worldwide prevalence of 17 million cases. High endemicity has been reported in certain areas of Peru and Bolivia. In most endemic areas the predominant species is *F. hepatica*, but in Asia and Africa a varying degree of overlap with *F. gigantica* has been observed.

Humans acquire fascioliasis by ingestion of metacercariae attached to certain aquatic plants, such as watercress, water caltrop, and water chestnuts. Infection may also be acquired by consumption of contaminated

TABLE 259-2 DRUG THERAPY FOR HUMAN TREMATODE INFECTIONS

Infection	Drug of Choice	Adult Dose and Duration
Blood Flukes		
<i>S. mansoni</i> , <i>S. intercalatum</i> , <i>S. haematobium</i>	Praziquantel	20 mg/kg, 2 doses in 1 day
<i>S. japonicum</i> , <i>S. mekongi</i>	Praziquantel	20 mg/kg, 3 doses in 1 day
Biliary (Hepatic) Flukes		
<i>Clonorchis sinensis</i> , <i>Opisthorchis viverrini</i> , <i>O. felineus</i>	Praziquantel	25 mg/kg, 3 doses in 1 day
<i>Fasciola hepatica</i> , <i>F. gigantica</i>	Triclabendazole	10 mg/kg once
Intestinal Flukes		
<i>Fasciolopsis buski</i> , <i>Heterophyes heterophyes</i>	Praziquantel ^a	25 mg/kg, 3 doses in 1 day
Lung Flukes		
<i>Paragonimus westermani</i>	Praziquantel ^a	25 mg/kg, 3 doses per day for 2 days

^aNot approved by the U.S. Food and Drug Administration for this indication.

water or ingestion of food items washed with such water. Acquisition of human infection through consumption of freshly prepared raw liver containing immature flukes has been reported. Infection is initiated when metacercariae excyst, penetrate the gut wall, and travel through the peritoneal cavity to invade the liver capsule. Adult worms migrate through the liver parenchyma and finally reach bile ducts, where they produce large operculated eggs that are voided in bile through the gastrointestinal tract to the outside environment. The flukes' life cycle is completed in specific snails (the first intermediate host) followed by encystment on aquatic plants.

Clinical features of fascioliasis relate to the stage and intensity of infection. Acute disease develops during parasite migration (1–2 weeks after infection) and includes fever, right-upper-quadrant pain, hepatomegaly, and eosinophilia. Computed tomography (CT) of the liver may show multiple parenchymal holes/or migratory tracks. Symptoms and signs usually subside as the parasites reach their final habitat. In individuals with chronic infection, bile duct obstruction and biliary cirrhosis are infrequently demonstrated. No relation to hepatic malignancy has been ascribed to fascioliasis.

DIAGNOSIS

Diagnosis of infection with any of the biliary flukes depends on a high degree of suspicion, elicitation of an appropriate geographic history, and stool examination for characteristically shaped parasite ova. Additional evidence may be obtained by documenting peripheral-blood eosinophilia or imaging the liver. Serologic testing is helpful, particularly in lightly infected individuals.

TREATMENT BILIARY FLUKES

Drug therapy (praziquantel or triclabendazole) is summarized in Table 259-2. Patients with anatomic lesions in the biliary tract or malignancy are managed according to general medical guidelines.

INTESTINAL FLUKES



Two species of intestinal flukes cause human infection in defined geographic areas worldwide (Table 259-1). The large *Fasciolopsis buski* (adults measure 2 × 7 cm) is endemic in Southeast Asia, whereas the smaller *Heterophyes heterophyes* is found in the Nile Delta of Egypt. Infection is initiated by ingestion of metacercariae attached to aquatic plants (*F. buski*) or encysted in freshwater or brackish-water fish (*H. heterophyes*). Flukes mature in human intestines, and eggs are passed with stools. Most individuals infected with intestinal flukes are asymptomatic. In heavy *F. buski* infection, diarrhea, abdominal pain, and malabsorption may be encountered. Heavy infection with *H. heterophyes* may be associated with abdominal pain and mucous diarrhea. The diagnosis is established by detection of characteristically shaped ova in stool samples. The drug of choice for treatment is praziquantel (Table 259-2).

LUNG FLUKES



Infection with the lung fluke *Paragonimus westermani* (Table 259-1) and related species (e.g., *P. africanus*) is endemic in many parts of the world, excluding North America and Europe. Endemicity is particularly noticeable in West Africa, Central and South America, and Asia. In nature, the reservoir hosts of *P. westermani* are wild and domestic felines. In Africa, *P. africanus* has been found in other species, such as dogs. Adult lung flukes, which are 7–12 mm in length, are found encapsulated in the lungs of infected persons. In rare circumstances, flukes are found encysted in the CNS (cerebral paragonimiasis) or the abdominal cavity. Humans acquire lung fluke infection by ingesting infective metacercariae encysted in the muscles

and viscera of crayfish and freshwater crabs. In endemic areas, these crustaceans are consumed raw, marinated, or pickled. Once the organisms reach the duodenum, they excyst, penetrate the gut wall, and travel through the peritoneal cavity, diaphragm, and pleural space to reach the lungs. Mature flukes are found in the bronchioles surrounded by cystic lesions. Parasite eggs are either expectorated with sputum or swallowed and passed to the outside environment with feces. The life cycle is completed in snails and freshwater crustaceans.

When maturing flukes lodge in lung tissues, they cause hemorrhage and necrosis, resulting in cyst formation. The adjacent lung parenchyma shows evidence of inflammatory infiltration, predominantly by eosinophils. Cysts usually measure 1–2 cm in diameter and may contain one or two worms each. With the onset of oviposition, cysts usually rupture in adjacent bronchioles—an event allowing ova to exit the human host. Older cysts develop thickened walls, which may undergo calcification. During the active phase of paragonimiasis, lung tissues surrounding parasite cysts may show evidence of pneumonia, bronchitis, bronchiectasis, and fibrosis.

Pulmonary paragonimiasis is particularly symptomatic in persons with moderate to heavy infection. Productive cough with brownish sputum or frank hemoptysis associated with peripheral-blood eosinophilia is usually the presenting feature. Chest examination may reveal signs of pleurisy. In chronic cases, bronchitis or bronchiectasis may predominate, but these conditions rarely proceed to lung abscess. Imaging of the lungs demonstrates characteristic features, including patchy densities, cavities, pleural effusion, and ring shadows. Cerebral paragonimiasis presents as either space-occupying lesions or epilepsy.

DIAGNOSIS

Pulmonary paragonimiasis is diagnosed by detection of parasite ova in sputum and/or stools. Serology is of considerable help in egg-negative cases and in cerebral paragonimiasis. The differential diagnosis includes active tuberculosis, bacterial lung abscess, and lung carcinoma.

TREATMENT LUNG FLUKES

The drug of choice for treatment is praziquantel (Table 259-2). Other medical or surgical management may be needed for pulmonary or cerebral lesions.

CONTROL AND PREVENTION OF TISSUE FLUKES



For residents of nonendemic areas who are visiting an endemic region, the only effective preventive measure is to avoid ingestion of local plants, fish, or crustaceans; if their ingestion is necessary, these items should be washed and cooked thoroughly. Instruction on water and food preparation and consumption should be included in physicians' advice to travelers (Chap. 149). Interruption of transmission among residents of endemic areas depends on avoiding ingestion of infective stages and disposing of feces and sputum appropriately to prevent hatching of eggs in the environment. These two approaches rely greatly on socioeconomic development, health education, and significant behavioral change. In countries where economic progress has resulted in financial and social improvements, transmission has decreased. The third approach to control in endemic communities entails selective use of chemotherapy for individuals posing the highest risk of transmission (i.e., those with heavy infections). The availability of praziquantel—a broad-spectrum, safe, and effective anthelmintic agent—provides a means for reducing the reservoirs of infection in human populations. However, the existence of most of these helminthic infections as zoonoses in several animal species complicates control efforts.

Cestodes, or tapeworms, are segmented worms. The adults reside in the gastrointestinal tract, but the larvae can be found in almost any organ. Human tapeworm infections can be divided into two major clinical groups. In one group, humans are the definitive hosts, with the adult tapeworms living in the gastrointestinal tract (*Taenia saginata*, *Diphyllobothrium*, *Hymenolepis*, and *Dipylidium caninum*). In the other, humans are intermediate hosts, with larval-stage parasites present in the tissues; diseases in this category include echinococcosis, sparganosis, and coenurosis. Humans may be either the definitive or the intermediate hosts for *Taenia solium*. Both stages of *Hymenolepis nana* are found simultaneously in the human intestines.

The ribbon-shaped tapeworm attaches to the intestinal mucosa by means of sucking cups or hooks located on the scolex. Behind the scolex is a short, narrow neck from which proglottids (segments) form. As each proglottid matures, it is displaced further back from the neck by the formation of new, less mature segments. The progressively elongating chain of attached proglottids, called the *strobila*, constitutes the bulk of the tapeworm. The length varies among species. In some, the tapeworm may consist of more than 1000 proglottids and may be several meters long. The mature proglottids are hermaphroditic and produce eggs, which are subsequently released. Because eggs of the different *Taenia* species are morphologically identical, differences in the morphology of the scolex or proglottids provide the basis for diagnostic identification to the species level.

Most human tapeworms require at least one intermediate host for complete larval development. After ingestion of the eggs or proglottids by an intermediate host, the larval oncospheres are activated, escape the egg, and penetrate the intestinal mucosa. The oncosphere migrates to tissues and develops into an encysted form known as a *cysticercus* (single scolex), a *coenurus* (multiple scolices), or a *hydatid* (cyst with daughter cysts, each containing several protoscolices). The definitive host's ingestion of tissues containing a cyst enables a scolex to develop into a tapeworm.

TAENIASIS SAGINATA AND TAENIASIS ASIATICA



The beef tapeworm *T. saginata* occurs in all countries where raw or undercooked beef is eaten. It is most prevalent in sub-Saharan African and Middle Eastern countries. *T. asiatica* is closely related to *T. saginata* and is found in Asia, with pigs as intermediate hosts. The clinical manifestations and morphology of these two species are very similar and are therefore discussed together.

Etiology and Pathogenesis Humans are the only definitive host for the adult stage of *T. saginata* and *T. asiatica*. The tapeworms, which can reach 8 m in length with 1000–2000 proglottids, inhabit the upper jejunum. The scolex of *T. saginata* has four prominent suckers, whereas *T. asiatica* has an unarmed rostellum. Each gravid segment has 15–30 uterine branches (in contrast to 8–12 for *T. solium*). The eggs are indistinguishable from those of *T. solium*; they measure 30–40 μm , contain the oncosphere, and have a thick brown striated shell. Eggs deposited on vegetation can live for months or years until they are ingested by cattle or other herbivores (*T. saginata*) or pigs (*T. asiatica*). The embryo released after ingestion invades the intestinal wall and is carried to striated muscle or viscera, where it transforms into the cysticercus. When ingested in raw or undercooked meat, this form can infect humans. After the cysticercus is ingested, it takes ~2 months for the mature adult worm to develop.

Clinical Manifestations Patients become aware of the infection most commonly by noting passage of proglottids in their feces. The proglottids are often motile, and patients may experience perianal discomfort when proglottids are discharged. Mild abdominal pain or discomfort, nausea, change in appetite, weakness, and weight loss can occur.

Diagnosis The diagnosis is made by the detection of eggs or proglottids in the stool. Eggs may also be present in the perianal area; thus, if proglottids or eggs are not found in the stool, the perianal region should be examined with use of a cellophane-tape swab (as in pinworm infection; Chap. 257). Distinguishing *T. saginata* or *T. asiatica* from *T. solium* requires examination of mature proglottids. All three species can be distinguished by examining the scolex. Available serologic tests are not helpful diagnostically. Eosinophilia and elevated levels of serum IgE may be detected.

TREATMENT TAENIASIS SAGINATA AND TAENIASIS ASIATICA

A single dose of praziquantel (10 mg/kg) is highly effective.

Prevention The major method of preventing infection is the adequate cooking of beef or pork viscera; exposure to temperatures as low as 56°C for 5 min will destroy cysticerci. Refrigeration or salting for long periods or freezing at –10°C for 9 days also kills cysticerci in beef. General preventive measures include inspection of beef and proper disposal of human feces.

TAENIASIS SOLIUM AND CYSTICERCOSIS

The pork tapeworm *T. solium* can cause two distinct forms of infection in humans: adult tapeworms in the intestine or larval forms in the tissues (cysticercosis). Humans are the only definitive hosts for *T. solium*; pigs are the usual intermediate hosts, although other animals may harbor the larval forms.



T. solium is found worldwide in areas where pigs are raised and have access to human feces. However, it is most prevalent in Latin America, sub-Saharan Africa, China, India, and Southeast Asia. Cysticercosis occurs in industrialized nations largely as a result of the immigration of infected persons from endemic areas.

Etiology and Pathogenesis The adult tapeworm generally resides in the upper jejunum. The scolex attaches by both sucking disks and two rows of hooklets. The adult worm usually lives for a few years. The tapeworm, usually ~3 m in length, may have as many as 1000 proglottids, each of which produces up to 50,000 eggs. Proglottids are released and excreted into the feces, and the eggs in these proglottids are infective for both humans and animals. The eggs may survive in the environment for several months. After ingestion of eggs by the pig intermediate host, the larvae are activated, escape the egg, penetrate the intestinal wall, and are carried to many tissues; they are most frequently identified in striated muscle of the neck, tongue, and trunk. Within 60–90 days, the encysted larval stage develops. These cysticerci can survive for months to years. By ingesting undercooked pork containing cysticerci, humans acquire infections that lead to intestinal tapeworms. Infections that cause human cysticercosis follow the ingestion of *T. solium* eggs, usually from close contact with a tapeworm carrier. Autoinfection may occur if an individual with an egg-producing tapeworm ingests eggs derived from his or her own feces.

Clinical Manifestations Intestinal infections with *T. solium* may be asymptomatic. Fecal passage of proglottids may be noted by patients. Other symptoms are infrequent.

In cysticercosis, the clinical manifestations are variable. Cysticerci can be found anywhere in the body but are most commonly detected in the brain, cerebrospinal fluid (CSF), skeletal muscle, subcutaneous tissue, or eye. The clinical presentation of cysticercosis depends on the number and location of cysticerci as well as on the extent of associated inflammatory responses or scarring. Neurologic manifestations are the most common (Fig. 260-1). Seizures are associated with inflammation surrounding cysticerci in the brain parenchyma. These seizures may be generalized, focal, or Jacksonian. Hydrocephalus results from CSF flow obstruction by cysticerci and accompanying inflammation or by CSF outflow obstruction from arachnoiditis. Symptoms of increased intracranial pressure, including headache, nausea, vomiting, changes in

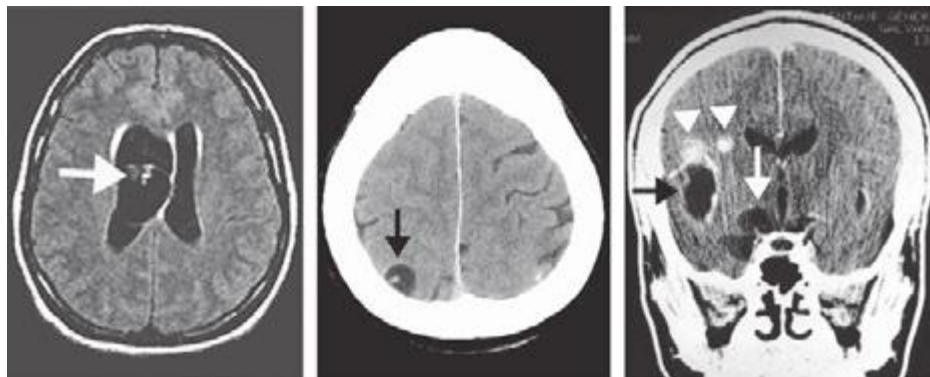


FIGURE 260-1 Neurocysticercosis is caused by *Taenia solium*. Neurologic infection can be classified on the basis of the location and viability of the parasites. When the parasites are in the ventricles, they often cause obstructive hydrocephalus. **Left:** Magnetic resonance imaging showing a cysticercus in the lateral ventricle, with resultant hydrocephalus. The *arrow* points to the scolex within the cystic parasite. **Center:** CT showing a parenchymal cysticercus, with enhancement of the cyst wall and an internal scolex (*arrow*). **Right:** Multiple cysticerci, including calcified lesions from prior infection (*arrowheads*), viable cysticerci in the basilar cisterns (*white arrow*), and a large degenerating cysticercus in the Sylvian fissure (*black arrow*). (Modified with permission from JC Bandres et al: *Clin Infect Dis* 15:799, 1992. © The University of Chicago Press.)

vision, dizziness, ataxia, or confusion, are often evident. Patients with hydrocephalus may develop papilledema or display altered mental status. When cysticerci develop at the base of the brain or in the subarachnoid space, they may cause chronic meningitis or arachnoiditis, communicating hydrocephalus, hemorrhages, or strokes.

Diagnosis The diagnosis of intestinal *T. solium* infection is made by the detection of eggs or proglottids, as described for *T. saginata*. More sensitive methods, including antigen-capture enzyme-linked immunosorbent assay (ELISA), polymerase chain reaction (PCR), and serology for tapeworm stage-specific antigens, are currently available only as research techniques. In cysticercosis, diagnosis can be difficult. A consensus conference has delineated absolute, major, minor, and epidemiologic criteria for diagnosis (Table 260-1). Diagnostic certainty is possible only with definite demonstration of the parasite (absolute criteria). This task can be accomplished by histologic observation of the parasite in excised tissue, by fundoscopic visualization of the parasite in the eye (in the anterior chamber, vitreous, or subretinal spaces), or by neuroimaging studies demonstrating cystic lesions containing a characteristic scolex (Fig. 260-1). With improving resolution of neuroimaging studies, the scolex can now be identified in many cases. In other instances, a clinical diagnosis is based on a combination of clinical presentation, radiographic studies, serologic tests, and exposure history.

Neuroimaging findings suggestive of neurocysticercosis constitute the primary major diagnostic criterion (Fig. 260-1). These findings include cystic lesions with or without enhancement (e.g., ring enhancement), one or more nodular calcifications (which may also have associated enhancement), or focal enhancing lesions. Cysticerci in the brain parenchyma are usually 5–20 mm in diameter and rounded. Cystic lesions in the subarachnoid space or fissures may enlarge up to 6 cm in diameter and may be lobulated. For cysticerci within the subarachnoid space or ventricles, the walls may be very thin and the cyst fluid is often isodense with CSF. Thus, obstructive hydrocephalus or enhancement of the basilar meninges may be the only finding on CT in extraparenchymal neurocysticercosis. Cysticerci in the ventricles or subarachnoid space are usually visible to an experienced neuroradiologist on MRI or on CT with intraventricular contrast injection. CT is more sensitive than MRI in identifying calcified lesions, whereas MRI is better for identifying cystic lesions, scolices, and enhancement.

The second major diagnostic criterion is detection of specific antibodies to cysticerci. Although most tests using unfractionated antigen have high rates of false-positive and false-negative results,

this problem can be overcome by using the more specific immunoblot assay. An immunoblot assay using lentil lectin purified glycoproteins is >99% specific and highly sensitive. However, patients with single intracranial lesions or with calcifications may be seronegative. With this assay, serum samples provide greater diagnostic sensitivity than CSF. All of the diagnostic antigens have been cloned, and assays using recombinant antigens are being developed. Antigen detection assays using monoclonal antibodies to detect parasite antigen in the blood or CSF may also facilitate diagnosis and patient follow-up. These assays are only now becoming available for patient care.

Studies have demonstrated that clinical criteria can aid in diagnosis in selected cases. In patients from endemic areas who had single enhancing lesions presenting with seizures, a normal physical examination, and no evidence of systemic disease (e.g., no fever, adenopathy, or chest radiographic abnormalities), the constellation of rounded CT lesions 5–20 mm in diameter with no midline shift was almost always

caused by neurocysticercosis. Finally, spontaneous resolution or resolution after therapy with albendazole alone is consistent with neurocysticercosis.

Minor diagnostic criteria include neuroimaging findings consistent with but less characteristic of cysticercosis, clinical manifestations suggestive of neurocysticercosis (e.g., seizures, hydrocephalus, or altered mental status), evidence of cysticercosis outside the central nervous system (CNS) (e.g., cigar-shaped soft-tissue calcifications), or detection of antibody in CSF by ELISA. Epidemiologic criteria include

TABLE 260-1 DIAGNOSTIC CRITERIA FOR HUMAN CYSTICERCOSIS^a

1. Absolute criteria
 - a. Demonstration of cysticerci by histologic or microscopic examination of biopsy material
 - b. Visualization of the parasite in the eye by funduscopy
 - c. Neuroradiologic demonstration of cystic lesions containing a characteristic scolex
2. Major criteria
 - a. Neuroradiologic lesions suggestive of neurocysticercosis
 - b. Demonstration of antibodies to cysticerci in serum by enzyme-linked immunoelectrotransfer blot
 - c. Resolution of intracranial cystic lesions spontaneously or after therapy with albendazole or praziquantel alone
3. Minor criteria
 - a. Lesions compatible with neurocysticercosis detected by neuroimaging studies
 - b. Clinical manifestations suggestive of neurocysticercosis
 - c. Demonstration of antibodies to cysticerci or cysticercal antigen in cerebrospinal fluid by enzyme-linked immunosorbent assay
 - d. Evidence of cysticercosis outside the central nervous system (e.g., cigar-shaped soft-tissue calcifications)
4. Epidemiologic criteria
 - a. Residence in a cysticercosis-endemic area
 - b. Frequent travel to a cysticercosis-endemic area
 - c. Household contact with an individual infected with *Taenia solium*

^aDiagnosis is confirmed by either one absolute criterion or a combination of two major criteria, one minor criterion, and one epidemiologic criterion. A probable diagnosis is supported by the fulfillment of (1) one major criterion plus two minor criteria; (2) one major criterion plus one minor criterion and one epidemiologic criterion; or (3) three minor criteria plus one epidemiologic criterion.

Source: Modified from OH Del Brutto et al: *Neurology* 57:177, 2001.

1432 exposure to a tapeworm carrier or household member infected with *T. solium*, current or prior residence in an endemic area, and frequent travel to an endemic area.

The diagnosis is confirmed in patients with either one absolute criterion or a combination of two major criteria, one minor criterion, and one epidemiologic criterion (Table 260-1). A probable diagnosis is supported by the fulfillment of (1) one major criterion plus two minor criteria; (2) one major criterion plus one minor criterion and one epidemiologic criterion; or (3) three minor criteria plus one epidemiologic criterion. Although the CSF is usually abnormal in neurocysticercosis, CSF abnormalities are not pathognomonic. Patients may have CSF pleocytosis with a predominance of lymphocytes, neutrophils, or eosinophils. The protein level in CSF may be elevated; the glucose concentration is usually normal but may be depressed.

TREATMENT TAENIASIS SOLIUM AND CYSTICERCOSIS

Intestinal *T. solium* infection is treated with a single dose of praziquantel (10 mg/kg). However, praziquantel occasionally evokes an inflammatory response in the CNS if concomitant cryptic cysticercosis is present. Niclosamide (2 g) is also effective but is not widely available.

The initial management of neurocysticercosis should focus on symptom-based treatment of seizures or hydrocephalus. Seizures can usually be controlled with antiepileptic treatment. If parenchymal lesions resolve without development of calcifications and patients remain free of seizures, antiepileptic therapy can usually be discontinued after 1–2 years. Placebo-controlled trials are clarifying the clinical advantage of antiparasitic drugs for parenchymal neurocysticercosis. Trends toward faster resolution of neuroradiologic abnormalities have been observed in most studies. The clinical benefits are less dramatic and consist mainly of shortening the period during which recurrent seizures occur and decreasing the number of patients who have many recurrent seizures. For the treatment of patients with brain parenchymal cysticerci, most authorities favor antiparasitic drugs, including albendazole (15 mg/kg per day for 8–28 days) or praziquantel (50–100 mg/kg daily in three divided doses for 15–30 days). A combination of albendazole and praziquantel (50 mg/kg per day) may be more effective in patients with multiple lesions. A longer course or combination therapy is often needed in patients with multiple subarachnoid cysticerci. Both agents may exacerbate the inflammatory response around the dying parasite, thereby exacerbating seizures or hydrocephalus as well. Thus, patients receiving these drugs should be carefully monitored. High-dose glucocorticoids should be used during treatment. Because glucocorticoids induce first-pass metabolism of praziquantel and may decrease its antiparasitic effect, cimetidine should be co-administered to inhibit praziquantel metabolism.

For patients with hydrocephalus, the emergent reduction of intracranial pressure is the mainstay of therapy. In the case of obstructive hydrocephalus, the preferred approach is removal of the cysticercus via endoscopic surgery. However, this intervention is not always possible. An alternative approach is initially to perform a diverting procedure, such as ventriculoperitoneal shunting. Historically, shunts have usually failed, but low failure rates have been attained with administration of antiparasitic drugs and glucocorticoids. Open craniotomy to remove cysticerci is now required only infrequently but is an alternative for fourth-ventricular cysticerci. For patients with subarachnoid cysts or giant cysticerci, anti-inflammatory medications such as glucocorticoids are needed to reduce arachnoiditis and accompanying vasculitis. Most authorities recommend prolonged courses of antiparasitic drugs as well as shunting when hydrocephalus is present. Methotrexate should be used as a steroid-sparing agent in patients requiring prolonged therapy. In patients with diffuse cerebral edema and elevated intracranial pressure due to multiple inflamed lesions, glucocorticoids are the mainstay of therapy, and antiparasitic drugs should be avoided. For ocular and spinal medullary lesions, drug-induced inflammation may cause

irreversible damage. Ocular disease should be managed surgically. Recent data suggest that either medical or surgical therapy can be used for spinal disease.

Prevention Measures for the prevention of intestinal *T. solium* infection consist of the application to pork of precautions similar to those described above for beef with regard to *T. saginata* infection. The prevention of cysticercosis involves minimizing the opportunities for ingestion of fecally derived eggs by means of good personal hygiene, effective fecal disposal, and treatment and prevention of human intestinal infections. Mass chemotherapy has been administered to human and porcine populations in efforts at disease eradication. Finally, vaccines to prevent porcine cysticercosis have shown promise in studies and are under development.

ECHINOCOCCOSIS



Echinococcosis is an infection caused in humans by the larval stage of the *Echinococcus granulosus* complex, *E. multilocularis*, or *E. vogeli*. *E. granulosus* complex parasites produce cystic hydatid disease, with unilocular cystic lesions. These infections are prevalent in most areas where livestock is raised in association with dogs. Molecular evidence suggests that *E. granulosus* strains may actually belong to more than one species; specifically, strains from sheep, cattle, pigs, horses, and camels probably represent separate species. These parasites are found on all continents, with areas of high prevalence in China, central Asia, the Middle East, the Mediterranean region, eastern Africa, and parts of South America. *E. multilocularis*, which causes multilocular alveolar lesions that are locally invasive, is found in Alpine, sub-Arctic, or Arctic regions, including Canada, the United States, and central and northern Europe; China; and central Asia. *E. vogeli* causes polycystic hydatid disease and is found only in Central and South America.

Like other cestodes, echinococcal species have both intermediate and definitive hosts. The definitive hosts are canines that pass eggs in their feces. After the ingestion of eggs, cysts develop in the intermediate hosts—sheep, cattle, humans, goats, camels, and horses for the *E. granulosus* complex and mice and other rodents for *E. multilocularis*. When a dog (*E. granulosus*) or fox (*E. multilocularis*) ingests infected meat containing cysts, the life cycle is completed.

Etiology The small (5-mm-long) adult *E. granulosus* complex worms, which live for 5–20 months in the jejunum of dogs, have only three proglottids: one immature, one mature, and one gravid. The gravid segment splits to release eggs that are morphologically similar to *Taenia* eggs and are extremely hardy. After humans ingest the eggs, embryos escape from the eggs, penetrate the intestinal mucosa, enter the portal circulation, and are carried to various organs, most commonly the liver and lungs. Larvae develop into fluid-filled unilocular hydatid cysts that consist of an external membrane and an inner germinal layer. Daughter cysts develop from the inner aspect of the germinal layer, as do germinating cystic structures called *brood capsules*. New larvae, called *protoscolices*, develop in large numbers within the brood capsule. The cysts expand slowly over a period of years.

The life cycle of *E. multilocularis* is similar except that wild canines, such as foxes, serve as the definitive hosts and small rodents serve as the intermediate hosts. The larval form of *E. multilocularis*, however, is quite different in that it remains in the proliferative phase, the parasite is always multilocular, and vesicles without brood capsule or *protoscolices* progressively invade the host tissue by peripheral extension of processes from the germinal layer.

Clinical Manifestations Slowly enlarging echinococcal cysts generally remain asymptomatic until their expanding size or their space-occupying effect in an involved organ elicits symptoms. The liver and the lungs are the most common sites of these cysts. The liver is involved in about two-thirds of *E. granulosus* infections and in nearly all *E. multilocularis* infections. Because a period of years elapses before cysts enlarge sufficiently to cause symptoms, they may be discovered incidentally on a routine x-ray or ultrasound study.

Patients with hepatic echinococcosis who are symptomatic most often present with abdominal pain or a palpable mass in the right upper quadrant. Compression of a bile duct or leakage of cyst fluid into the biliary tree may mimic recurrent cholelithiasis, and biliary obstruction can result in jaundice. Rupture of or episodic leakage from a hydatid cyst may produce fever, pruritus, urticaria, eosinophilia, or anaphylaxis. Pulmonary hydatid cysts may rupture into the bronchial tree or pleural cavity and produce cough, salty phlegm, dyspnea, chest pain, or hemoptysis. Rupture of hydatid cysts, which can occur spontaneously or at surgery, may lead to multifocal dissemination of protoscolices, which can form additional cysts. Other presentations are due to the involvement of bone (invasion of the medullary cavity with slow bone erosion producing pathologic fractures), the CNS (space-occupying lesions), the heart (conduction defects, pericarditis), and the pelvis (pelvic mass).

The larval forms of *E. multilocularis* characteristically present as a slowly growing hepatic tumor, with progressive destruction of the liver and extension into vital structures. Patients commonly report upper-quadrant and epigastric pain. Liver enlargement and obstructive jaundice may be apparent. The lesions may infiltrate adjoining organs (e.g., diaphragm, kidneys, or lungs) or may metastasize to the spleen, lungs, or brain.

Diagnosis Radiographic and related imaging studies are important in detecting and evaluating echinococcal cysts. Plain x-rays will define pulmonary cysts of *E. granulosus*—usually as rounded masses of uniform density—but may miss cysts in other organs unless there is cyst wall calcification (as occurs in the liver). MRI, CT, and ultrasound reveal well-defined cysts with thick or thin walls. When older cysts contain a layer of hydatid sand that is rich in accumulated protoscolices, these imaging methods may detect this fluid layer of different density. However, the most pathognomonic finding, if demonstrable, is that of daughter cysts within the larger cyst. This finding, like eggshell or mural calcification on CT, is indicative of *E. granulosus* infection and helps to distinguish the cyst from carcinomas, bacterial or amebic liver abscesses, or hemangiomas. In contrast, ultrasound or CT of alveolar hydatid cysts reveals indistinct solid masses with central necrosis and plaque-like calcifications.

A specific diagnosis of *E. granulosus* infection can be made by the examination of aspirated fluids for protoscolices or hooklets, but diagnostic aspiration is not usually recommended because of the risk of

fluid leakage resulting in either dissemination of infection or anaphylactic reactions. Serodiagnostic assays can be useful, although a negative test does not exclude the diagnosis of echinococcosis. Cysts in the liver elicit positive antibody responses in ~90% of cases, whereas up to 50% of individuals with cysts in the lungs are seronegative. Detection of antibody to specific echinococcal antigens by immunoblotting has the highest degree of specificity.

TREATMENT ECHINOCOCCOSIS

Therapy for cystic echinococcosis is based on considerations of the size, location, and manifestations of cysts and the overall health of the patient. Surgery has traditionally been the principal definitive method of treatment. Currently, ultrasound staging is recommended for *E. granulosus* infections (Fig. 260-2). Small CL, CE1, and CE3 lesions may respond to chemotherapy with albendazole. For CE1 lesions and uncomplicated CE3 lesions, PAIR (percutaneous aspiration, infusion of scolicidal agents, and reaspiration) is now recommended instead of surgery. PAIR is contraindicated for superficially located cysts (because of the risk of rupture), for cysts with multiple thick internal septal divisions (honeycombing pattern), and for cysts communicating with the biliary tree. For prophylaxis of secondary peritoneal echinococcosis due to inadvertent spillage of fluid during PAIR, the administration of albendazole (15 mg/kg daily in two divided doses) should be initiated at least 2 days before the procedure and continued for at least 4 weeks afterward. Ultrasound- or CT-guided aspiration allows confirmation of the diagnosis by demonstration of protoscolices in the aspirate. After aspiration, contrast material should be injected to detect occult communications with the biliary tract. Alternatively, the fluid should be checked for bile staining visually and by dipstick. If no bile is found and no communication is visualized, the contrast material is reaspirated, with subsequent infusion of scolicidal agents (usually 95% ethanol; alternatively, hypertonic saline). This approach, when implemented by a skilled practitioner, yields rates of cure and relapse equivalent to those following surgery, with less perioperative morbidity and shorter hospitalization. In experienced hands, some CE2 lesions can be treated by aspiration with a trocar. Daughter cysts within the primary cyst may need to be punctured separately, and catheter drainage may be required.

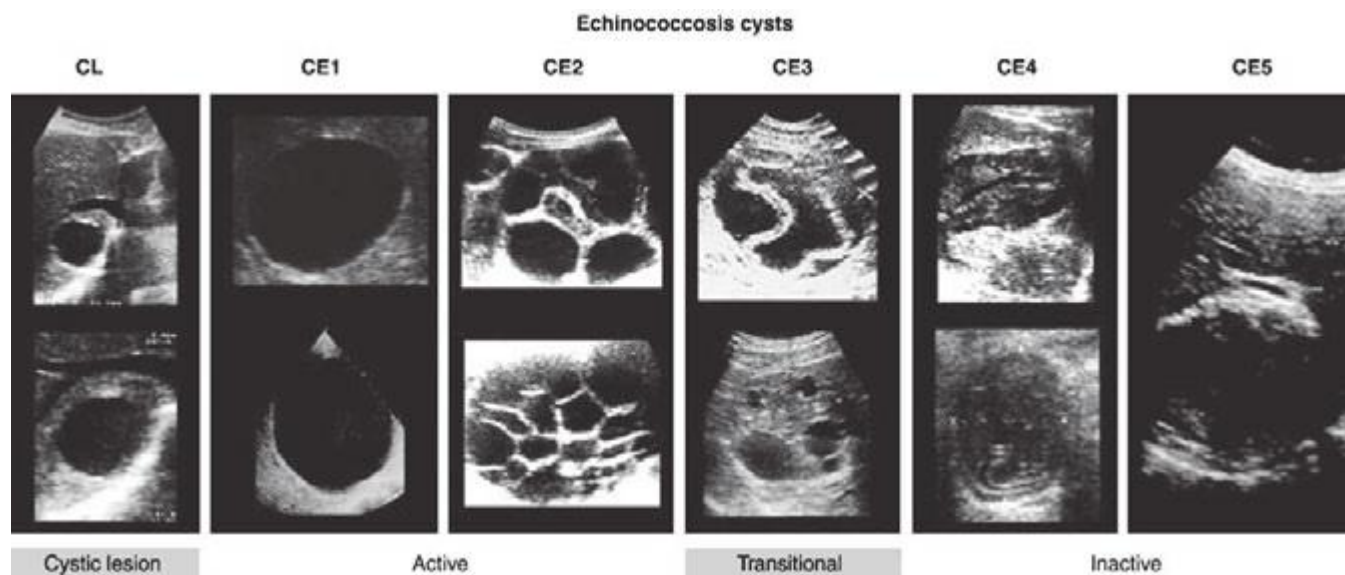




FIGURE 260-2 Management of cystic hydatid disease caused by *Echinococcus granulosus* should be based on viability of the parasite, which can be estimated from radiographic appearance. The ultrasound appearance includes lesions classified as active, transitional, and inactive. *Active* cysts include types CL (with a cystic lesion and no visible cyst wall), CE1 (with a visible cyst wall and internal echoes [snowflake sign]), and CE2 (with a visible cyst wall and internal septation). *Transitional* cysts (CE3) may have detached laminar membranes or may be partially collapsed. *Inactive* cysts include types CE4 (a nonhomogeneous mass) and CE5 (a cyst with a thick calcified wall). (Adapted from RL Guerrant et al [eds]: *Tropical Infectious Diseases: Principles, Pathogens and Practice*, 2nd ed, p 1312. © 2005, with permission from Elsevier Science.)

1434 Surgery remains the treatment of choice for complicated *E. granulosus* cysts (e.g., those communicating with the biliary tract), for most thoracic and intracranial cysts, and for areas where PAIR is not possible. For *E. granulosus* of the liver, the preferred surgical approach is pericystectomy, in which the entire cyst and the surrounding fibrous tissue are removed. The risks posed by leakage of fluid during surgery or PAIR include anaphylaxis and dissemination of infectious protoscolices. The latter complication has been minimized by careful attention to the prevention of spillage of the cyst and by soaking of the drapes with hypertonic saline. Infusion of scolical agents is no longer recommended because of problems with hypernatremia, intoxication, or sclerosing cholangitis. Albendazole, which is active against *Echinococcus*, should be administered adjunctively, beginning several days before resection of the liver and continuing for several weeks for *E. granulosus*. Praziquantel (50 mg/kg daily for 2 weeks) may hasten the death of the protoscolices. Medical therapy with albendazole alone for 12 weeks to 6 months results in cure in ~30% of cases and in improvement in another 50%. In many instances of treatment failure, *E. granulosus* infections are subsequently treated successfully with PAIR or additional courses of medical therapy. Response to treatment is best assessed by serial imaging studies, with attention to cyst size and consistency. Some cysts may not demonstrate complete radiologic resolution even though no viable protoscolices are present. Some of these cysts with partial radiologic resolution (e.g., CE4 or CE5) can be managed with observation only.

Surgical resection remains the treatment of choice for *E. multilocularis* infection. Complete removal of the parasite continues to offer the best chance for cure. Ongoing therapy with albendazole for at least 2 years after presumptively curative surgery is recommended. Positron emission tomography can be used to follow disease activity. Most cases are diagnosed at a stage at which complete resection is not possible; in these cases, albendazole treatment should be continued indefinitely, with careful monitoring. In some cases, liver transplantation has been used because of the size of the necessary liver resection. However, continuous immunosuppression favors the proliferation of *E. multilocularis* larvae and reinfection of the transplant. Thus, indefinite treatment with albendazole is required.

 **Prevention** In endemic areas, echinococcosis can be prevented by administering praziquantel to infected dogs, by denying dogs access to infected animals, or by vaccinating sheep. Limitation of the number of stray dogs is helpful in reducing the prevalence of infection among humans. In Europe, *E. multilocularis* infection has been associated with gardening; gloves should be used when working with soil.

HYMENOLEPIASIS NANA

 Infection with *Hymenolepis nana*, the dwarf tapeworm, is the most common of all the cestode infections. *H. nana* is endemic in both temperate and tropical regions of the world. Infection is spread by fecal/oral contamination and is common among institutionalized children.

Etiology and Pathogenesis *H. nana* is the only cestode of humans that does not require an intermediate host. Both the larval and adult phases of the life cycle take place in the human. The adult—the smallest tapeworm parasitizing humans—is ~2 cm long and dwells in the proximal ileum. Proglottids, which are small and rarely seen in the stool, release spherical eggs 30–44 μm in diameter, each of which contains an oncosphere with six hooklets. The eggs are immediately infective and are unable to survive for >10 days in the external environment. When the egg is ingested by a new host, the oncosphere is freed and penetrates the intestinal villi, becoming a cysticercoid larva. Larvae migrate back into the intestinal lumen, attach to the mucosa, and mature into adult worms over 10–12 days. Eggs may also hatch before passing into the stool, causing internal autoinfection with increasing numbers of intestinal worms. Although the life span of adult *H. nana* worms is only ~4–10 weeks, the autoinfection cycle perpetuates the infection.

Clinical Manifestations *H. nana* infection, even with many intestinal worms, is usually asymptomatic. When infection is intense, anorexia, abdominal pain, and diarrhea develop.

Diagnosis Infection is diagnosed by the finding of eggs in the stool.

TREATMENT HYMENOLEPIASIS NANA


Praziquantel (25 mg/kg once) is the treatment of choice, because it acts against both the adult worms and the cysticercoids in the intestinal villi. Nitazoxanide (500 mg bid for 3 days) may be used as an alternative.

Prevention Good personal hygiene and improved sanitation can eradicate the disease. Epidemics have been controlled by mass chemotherapy coupled with improved hygiene.

HYMENOLEPIASIS DIMINUTA


Hymenolepis diminuta, a cestode of rodents, occasionally infects small children, who ingest the larvae in uncooked cereal foods contaminated by fleas and other insects in which larvae develop. Infection is usually asymptomatic and is diagnosed by the detection of eggs in the stool. Treatment with praziquantel results in cure in most cases.

DIPHYLLOBOTHRIASIS

 *Diphyllobothrium latum* and other *Diphyllobothrium* species are found in the lakes, rivers, and deltas of the Northern Hemisphere, central Africa, and South America.

Etiology and Pathogenesis The adult worm—the longest tapeworm (up to 25 m)—attaches to the ileal and occasionally to the jejunal mucosa by its suckers, which are located on its elongated scolex. The adult worm has 3000–4000 proglottids, which release ~1 million eggs daily into the feces. If an egg reaches water, it hatches and releases a free-swimming embryo that can be eaten by small freshwater crustaceans (*Cyclops* or *Diatomus* species). After an infected crustacean containing a developed proceroid is swallowed by a fish, the larva migrates into the fish's flesh and grows into a plerocercoid, or sparganum larva. Humans acquire the infection by ingesting infected raw or smoked fish. Within 3–5 weeks, the tapeworm matures into an adult in the human intestine.

Clinical Manifestations Most *D. latum* infections are asymptomatic, although manifestations may include transient abdominal discomfort, diarrhea, vomiting, weakness, and weight loss. Occasionally, infection can cause acute abdominal pain and intestinal obstruction; in rare cases, cholangitis or cholecystitis may be produced by migrating proglottids.

 Because the tapeworm absorbs large quantities of vitamin B₁₂ and interferes with ileal B₁₂ absorption, vitamin B₁₂ deficiency can develop, but this effect has been noted only in Scandinavia, where up to 2% of infected patients, especially the elderly, have megaloblastic anemia resembling pernicious anemia and may exhibit neurologic sequelae of B₁₂ deficiency.

Diagnosis The diagnosis is made readily by the detection of the characteristic eggs in the stool. The eggs possess a single shell with an operculum at one end and a knob at the other. Mild to moderate eosinophilia may be detected.

TREATMENT DIPHYLLOBOTHRIASIS

Praziquantel (5–10 mg/kg once) is highly effective. Parenteral vitamin B₁₂ should be given if B₁₂ deficiency is manifest.

Prevention Infection can be prevented by heating fish to 54°C for 5 min or by freezing it at –18°C for 24 h. Placing fish in brine with a high salt concentration for long periods kills the eggs.

DIPYLIDIASIS

Dipylidium caninum, a common tapeworm of dogs and cats, may accidentally infect humans. Dogs, cats, and occasionally humans become infected by ingesting fleas harboring cysticercoids. Children are more likely to become infected than adults. Most infections are asymptomatic, but abdominal pain, diarrhea, anal pruritus, urticaria, eosinophilia, or passage of segments in the stool may occur. The diagnosis is made by the detection of proglottids or ova in the stool. As in *D. latum* infection, therapy consists of praziquantel. Prevention requires anthelmintic treatment and flea control for pet dogs or cats.

SPARGANOSIS

Humans can be infected by the sparganum, or plerocercoid larva, of a diphylobothrid tapeworm of the genus *Spirometra*. Infection can

be acquired by the consumption of water containing infected *Cyclops*; by the ingestion of infected snakes, birds, or mammals; or by the application of infected flesh as poultices. The worm migrates slowly in tissues, and infection commonly presents as a subcutaneous swelling. Periorbital tissues can be involved, and ocular sparganosis may destroy the eye. Surgical excision is used to treat localized sparganosis.

COENUROSIS

This rare infection of humans by the larval stage (coenurus) of the dog tapeworm *Taenia multiceps* or *T. serialis* results in a space-occupying cystic lesion. As in cysticercosis, involvement of the CNS and subcutaneous tissue is most common. Both definitive diagnosis and treatment require surgical excision of the lesion. Chemotherapeutic agents generally are not effective.

This page intentionally left blank

261e Microbial Bioterrorism

H. Clifford Lane, Anthony S. Fauci

Descriptions of the use of microbial pathogens as potential weapons of war or terrorism date from ancient times. Among the most frequently cited of such episodes are the poisoning of water supplies in the sixth century B.C. with the fungus *Claviceps purpurea* (rye ergot) by the Assyrians, the hurling of the dead bodies of plague victims over the walls of the city of Kaffa by the Tartar army in 1346, and the efforts by the British to spread smallpox to the Native American population loyal to the French via contaminated blankets in 1763. The tragic attacks on the World Trade Center and the Pentagon on September 11, 2001, followed closely by the mailing of letters containing anthrax spores to media and congressional offices through the U.S. Postal Service, dramatically changed the mindset of the American public regarding both our vulnerability to microbial bioterrorist attacks and the seriousness and intent of the federal government to protect its citizens against future attacks. Modern science has revealed methods of deliberately spreading or enhancing disease in ways not appreciated by our ancestors. The combination of basic research, good medical practice, and constant vigilance will be needed to defend against such attacks.

Although the potential impact of a bioterrorist attack could be enormous, leading to thousands of deaths and high morbidity rates, acts of bioterrorism would be expected to produce their greatest impact through the fear and terror they generate. In contrast to biowarfare, where the primary goal is destruction of the enemy through mass casualties, an important goal of bioterrorism is to destroy the morale of a society through fear and uncertainty. Although the actual biologic impact of a single act may be small, the degree of disruption created by the realization that such an attack is possible may be enormous. This was readily apparent with the impact on the U.S. Postal Service and the functional interruption of the activities of the legislative branch of the U.S. government following the anthrax attacks noted above. Thus, the key to the defense against these attacks is a highly functioning system of public health surveillance and education so that attacks can be quickly recognized and effectively contained. This is complemented by the availability of appropriate countermeasures in the form of diagnostics, therapeutics, and vaccines, both in response to and in anticipation of bioterrorist attacks.

The Working Group for Civilian Biodefense created a list of key features that characterize the elements of biologic agents that make them particularly effective as weapons (Table 261e-1). Included among these are the ease of spread and transmission of the agent and the presence of an adequate database to allow newcomers to the field to quickly apply the good science of others to bad intentions of their own. Agents of bioterrorism may be used in their naturally occurring forms, or they can be deliberately modified to deliver greater impact. Among the approaches to maximizing the deleterious effects of biologic agents are

TABLE 261e-1 KEY FEATURES OF BIOLOGIC AGENTS USED AS BIOWEAPONS

1. High morbidity and mortality rates
2. Potential for person-to-person spread
3. Low infective dose and highly infectious by aerosol
4. Lack of rapid diagnostic capability
5. Lack of universally available effective vaccine
6. Potential to cause anxiety
7. Availability of pathogen and feasibility of production
8. Environmental stability
9. Database of prior research and development
10. Potential to be “weaponized”

Source: From L Borio et al: JAMA 287:2391, 2002; with permission.

the genetic modification of microbes for the purposes of antimicrobial resistance or evasion by the immune system, creation of fine-particle aerosols, chemical treatment to stabilize and prolong infectivity, and alteration of host range through changes in surface proteins. Certain of these approaches fall under the category of *weaponization*, which is a term generally used to describe the processing of microbes or toxins in a manner that would ensure a devastating effect following release. For example, weaponization of anthrax by the Soviets involved the production of vast numbers of spores of appropriate size to reach the lower respiratory tract easily in a form that maintained aerosolization for prolonged periods of time and that could be delivered in a massive release, such as via widely dispersed bomblets.

The U.S. Centers for Disease Control and Prevention (CDC) classifies potential biologic threats into three categories: A, B, and C (Table 261e-2). Category A agents are the highest-priority pathogens. They pose the greatest risk to national security because they (1) can be easily disseminated or transmitted from person to person, (2) result in high mortality rates and have the potential for major public health impact, (3) might cause public panic and social disruption, and (4) require special action for public health preparedness. Category B agents are the second highest priority pathogens and include those that are moderately easy to disseminate, result in moderate morbidity rates and low mortality rates, and require specifically enhanced diagnostic capacity. Category C agents are the third highest priority. These include certain emerging pathogens to which the general population lacks immunity; that could be engineered for mass dissemination in the future because of availability, ease of production, and ease of dissemination; and that have a major public health impact and the potential for high morbidity and mortality rates. It should be pointed out, however, that these A,

TABLE 261e-2 CDC CATEGORY A, B, AND C AGENTS

Category A
Anthrax (<i>Bacillus anthracis</i>)
Botulism (<i>Clostridium botulinum</i> toxin)
Plague (<i>Yersinia pestis</i>)
Smallpox (<i>Variola major</i>)
Tularemia (<i>Francisella tularensis</i>)
Viral hemorrhagic fevers
Arenaviruses: Lassa, New World (Machupo, Junin, Guanarito, and Sabia)
Bunyaviridae: Crimean-Congo, Rift Valley
Filoviridae: Ebola, Marburg
Category B
Brucellosis (<i>Brucella</i> spp.)
Epsilon toxin of <i>Clostridium perfringens</i>
Food safety threats (e.g., <i>Salmonella</i> spp., <i>Escherichia coli</i> O157:H7, <i>Shigella</i>)
Glanders (<i>Burkholderia mallei</i>)
Melioidosis (<i>Burkholderia pseudomallei</i>)
Psittacosis (<i>Chlamydia psittaci</i>)
Q fever (<i>Coxiella burnetii</i>)
Ricin toxin from <i>Ricinus communis</i> (castor beans)
Staphylococcal enterotoxin B
Typhus fever (<i>Rickettsia prowazekii</i>)
Viral encephalitis (alphaviruses [e.g., Venezuelan, eastern, and western equine encephalitis])
Water safety threats (e.g., <i>Vibrio cholerae</i> , <i>Cryptosporidium parvum</i>)
Category C
Emerging infectious diseases threats such as Nipah, hantavirus, SARS or MERS coronavirus, and pandemic influenza

Abbreviations: MERS, Middle East respiratory syndrome; SARS, severe acute respiratory syndrome.

Source: Centers for Disease Control and Prevention and the National Institute of Allergy and Infectious Diseases.

B, and C designations are empirical and, depending on evolving circumstances such as intelligence-based threat assessments, the priority rating of any given microbe or toxin could change. The CDC classification system also largely reflects the severity of illness produced by a given agent, rather than its accessibility to potential terrorists.

CATEGORY A AGENTS

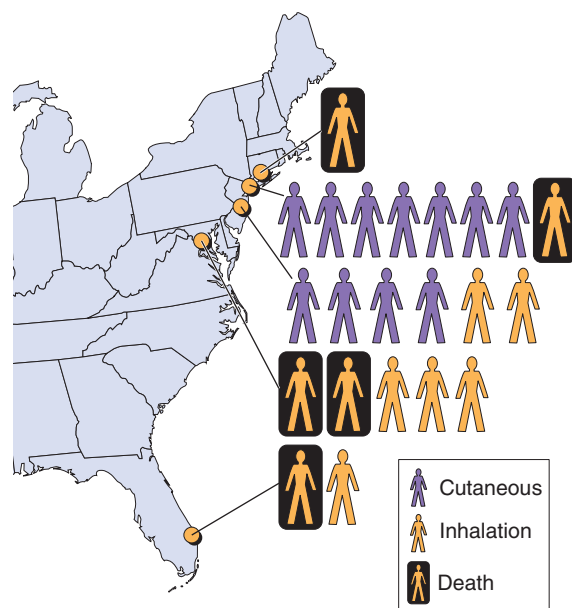
ANTHRAX

See also Chap. 175.

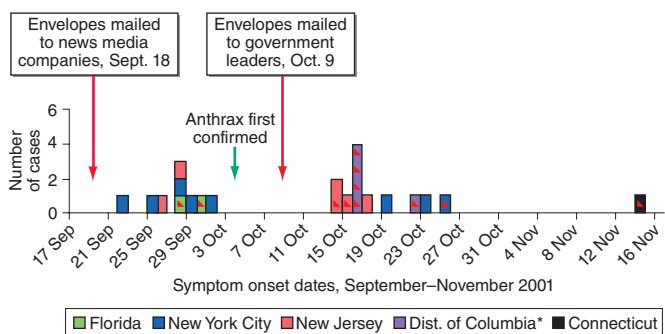
Bacillus anthracis as a Bioweapon Anthrax may be the prototypic disease of bioterrorism. Although rarely, if ever, spread from person to person, the illness embodies the other major features of a disease introduced through terrorism, as outlined in Table 261e-1. U.S. and British government scientists studied anthrax as a potential biologic weapon beginning approximately at the time of World War II (WWII). Offensive bioweapons activity including bioweapons research on microbes and toxins in the United States ceased in 1969 as a result of two executive orders by President Richard M. Nixon. Although the 1972 Biological and Toxin Weapons Convention Treaty outlawed research of this type worldwide, the Soviet Union produced and stored tons of anthrax spores for potential use as a bioweapon until at least the late 1980s. At present, there is suspicion that research on anthrax as an agent of bioterrorism is ongoing by several nations and extremist groups. One example of this was the release of anthrax spores by the Aum Shinrikyo cult in Tokyo in 1993. Fortunately, there were no casualties associated with that episode because of the inadvertent use of a nonpathogenic strain of anthrax by the terrorists.

The potential impact of anthrax spores as a bioweapon was clearly demonstrated in 1979 following the accidental release of spores into the atmosphere from a Soviet Union bioweapons facility in Sverdlovsk, Russia. Although total figures are not known, at least 77 cases of anthrax were diagnosed with certainty, of which 66 were fatal. These victims were exposed in an area within 4 km downwind of the facility, and deaths due to anthrax were also noted in livestock up to 50 km further downwind. Based on recorded wind patterns, the interval between the time of exposure and development of clinical illness ranged from 2 to 43 days. The majority of cases were within the first 2 weeks. Death typically occurred within 1–4 days following the onset of symptoms. It is likely that the widespread use of postexposure penicillin prophylaxis limited the total number of cases. The extended period of time between exposure and disease in some individuals supports the data from nonhuman primate studies, suggesting that anthrax spores can lie dormant in the respiratory tract for at least 4–6 weeks without evoking an immune response. This extended period of microbiologic latency following exposure poses a significant challenge for management of victims in the postexposure period.

In September 2001, the American public was exposed to anthrax spores as a bioweapon delivered through the U.S. Postal Service by an employee of the United States Army Medical Research Institute for Infectious Diseases (USAMRIID) who had access to such materials and who committed suicide prior to being indicted for this crime. The CDC identified 22 confirmed or suspected cases of anthrax as a consequence of this attack. These included 11 patients with inhalational anthrax, of whom 5 died, and 11 patients with cutaneous anthrax (7 confirmed), all of whom survived (Fig. 261e-1). Cases occurred in individuals who opened contaminated letters as well as in postal workers involved in the processing of mail. A minimum of five letters mailed from Trenton, NJ, served as the vehicles for these attacks. One of these letters was reported to contain 2 g of material, equivalent to 100 billion to 1 trillion weapon-grade spores. Based on studies performed in the 1950s using monkeys exposed to aerosolized anthrax demonstrating that ~10,000 spores were required to produce lethal disease in 50% of animals exposed to this dose (the LD_{50}), the contents of one letter had the theoretical potential, under optimal conditions, of causing illness or death in up to 50 million individuals. The strain used in this attack was the Ames strain. Although it was noted to have an inducible β -lactamase and to constitutively express a cephalosporinase, it was susceptible to all antibiotics standard for *B. anthracis*.



A



* = Metropolitan District of Columbia Area, cases were residents of Maryland (3) and Virginia (2).

B

FIGURE 261e-1 Confirmed anthrax cases associated with bioterrorism: United States, 2001. **A.** Geographic location, clinical manifestation, and outcome of the 11 cases of confirmed inhalational and 11 cases of confirmed cutaneous anthrax. **B.** Epidemic curve for 22 cases of anthrax. (From DB Jernigan et al: *Emerg Infect Dis* 8:1019, 2002; with permission.)

Microbiology and Clinical Features Anthrax is caused by *B. anthracis*, a gram-positive, nonmotile, spore-forming rod that is found in soil and predominantly causes disease in herbivores such as cattle, goats, and sheep. Anthrax spores can remain viable for decades. The remarkable stability of these spores makes them an ideal bioweapon, and their destruction in decontamination activities can be a challenge. Naturally occurring human infection is generally the result of contact with anthrax-infected animals or animal products such as goat hair in textile mills or animal skins used in making drums. While an LD_{50} of 10,000 spores is a generally accepted number, it has also been suggested that as few as one to three spores may be adequate to cause disease in some settings. Advanced technology is likely to be necessary to generate a bioweapon containing spores of the optimal size (1–5 μ m) to travel to the alveolar spaces.

The three major clinical forms of anthrax are gastrointestinal, cutaneous, and inhalational. *Gastrointestinal anthrax* typically results from the ingestion of contaminated meat; the condition is rarely seen and is unlikely to be the result of a bioterrorism event. The lesion of *cutaneous anthrax* typically begins as a papule following the introduction of spores through an opening in the skin. This papule then evolves to a painless vesicle followed by the development of a coal-black, necrotic eschar (Fig. 261e-2). It is the Greek word for coal (*anthrax*) that gives the organism and the disease its name. Cutaneous anthrax was ~20% fatal prior to the availability of antibiotics. *Inhalational anthrax* is the

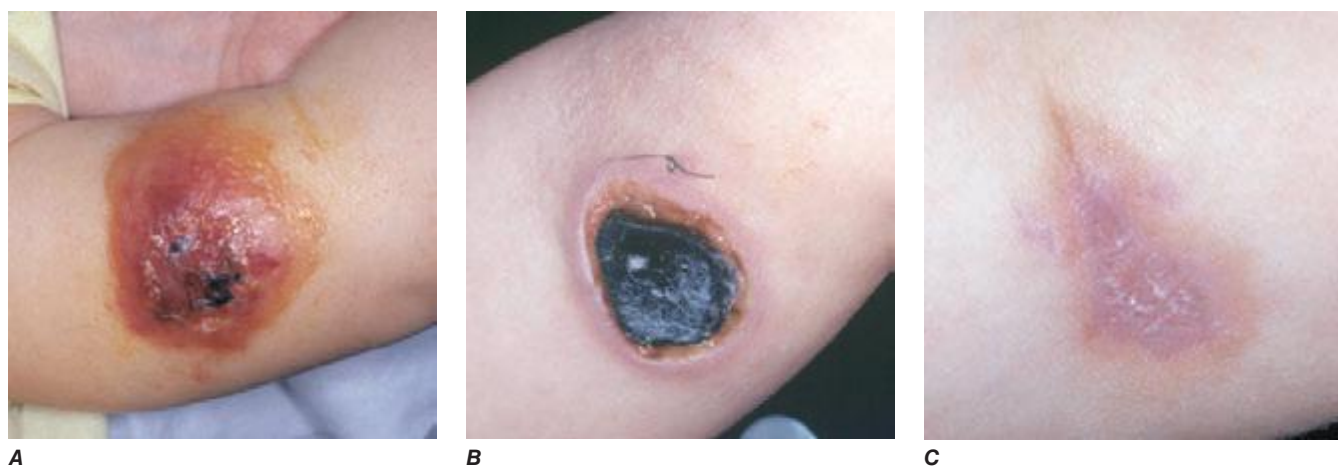


FIGURE 261e-2 Clinical manifestations of a pediatric case of cutaneous anthrax associated with the bioterrorism attack of 2001.

The lesion progresses from vesicular on day 5 (A) to necrotic with the classic black eschar on day 12 (B) to a healed scar 2 months later (C). (Photographs provided by Dr. Mary Wu Chang. Part A reproduced with permission from KJ Roche et al: *N Engl J Med* 345:1611, 2001 and Parts B and C reproduced with permission from A Freedman et al: *JAMA* 287:869, 2002.)

form most likely to be responsible for death in the setting of a bioterrorist attack. It occurs following the inhalation of spores that become deposited in the alveolar spaces. These spores are phagocytized by macrophages and transported to the mediastinal and peribronchial lymph nodes where they germinate, leading to active bacterial growth and elaboration of the bacterial products edema toxin and lethal toxin. Subsequent hematogenous spread of bacteria is accompanied by cardiovascular collapse and death. The earliest symptoms are typically a viral-like prodrome with fever, malaise, and abdominal and/or chest symptoms that progress over the course of a few days to a moribund state. Characteristic findings on chest x-ray include mediastinal widening and pleural effusions (Fig. 261e-3). Although initially thought to be 100% fatal, the experiences at Sverdlovsk in 1979 and in the United States in 2001 (see below) indicate that with prompt initiation of antibiotic therapy, survival is possible. The characteristics of the 11 cases of inhalational anthrax diagnosed in the United States in 2001 following exposure to contaminated letters postmarked September 18 or October 9, 2001, followed the classic pattern established for this illness, with patients presenting with a rapidly progressive course characterized by fever, fatigue or malaise, nausea or vomiting, cough, and shortness of breath. At presentation, the total white blood cell counts were $\sim 10,000$ cells/ μL ; transaminases tended to be elevated, and all 11 had abnormal findings on chest x-ray and computed tomography (CT). Radiologic findings included infiltrates, mediastinal widening, and hemorrhagic pleural effusions. For cases in which the dates of exposure were known, symptoms appeared within 4–6 days. Death occurred within 7 days of diagnosis in the five fatal cases (overall mortality rate 55%). Rapid diagnosis and prompt initiation of antibiotic therapy were key to survival.

TREATMENT ANTHRAX

Anthrax can be successfully treated if the disease is promptly recognized and appropriate therapy with antibiotics and antitoxin is initiated early. Although penicillin, ciprofloxacin, and doxycycline are the currently licensed antibiotics for this indication, clindamycin and rifampin also have *in vitro* activity against the organism and have been used as part of treatment regimens. Until sensitivity results are known, suspected cases are best managed with a combination of broadly active agents (Table 261e-3). The *B. anthracis* toxins lethal factor and edema factor share a component known as protective antigen. Raxibacumab, a monoclonal antibody directed toward protective antigen, was licensed in 2012 under the animal rule (see below) for treatment of adult and pediatric patients with inhalational anthrax in combination with appropriate antibacterial drugs. Patients with inhalational anthrax are not contagious and do not require special isolation procedures.

Vaccination and Prevention The first successful vaccine for anthrax was developed for animals by Louis Pasteur in 1881. At present, the single vaccine licensed for human use is a product produced from the cell-free culture supernatant of an attenuated, nonencapsulated strain of *B. anthracis* (Stern strain), referred to as *anthrax vaccine adsorbed* (AVA). Clinical trials for safety in humans and efficacy in animals are currently under way to evaluate the role of recombinant protective antigen as an alternative to AVA. In a postexposure setting in nonhuman primates, a 2-week course of AVA plus ciprofloxacin was found to be superior to ciprofloxacin alone in preventing the development of clinical disease and death. Although the current recommendation for postexposure prophylaxis is 60 days of antibiotics, it would seem prudent to include immunization with anthrax vaccine if available. Given the potential for *B. anthracis* to be engineered to express penicillin resistance, the empirical regimen of choice in this setting is either ciprofloxacin or doxycycline. In settings where these approaches are not available or appropriate, one can administer the antitoxin monoclonal antibody raxibacumab.

PLAGUE

See also Chap. 196.

***Yersinia pestis* as a Bioweapon** Although it lacks the environmental stability of anthrax, the highly contagious nature and high mortality rate of plague make it a close to ideal agent of bioterrorism, particularly if delivered in a weaponized form. Occupying a unique place in history, plague has been alleged to have been used as a biologic weapon for centuries. The catapulting of plague-infected corpses into besieged fortresses is a practice that was first noted in 1346 during the assault of the Crimean city of Kaffa by the Mongolian Tartars. Although unlikely to have resulted in disease transmission, some believe that this event may have played a role in the start of the Black Death pandemic of the fourteenth and fifteenth centuries in Europe. Given that plague was already moving across Asia toward Europe at this time, it is unclear whether such an allegation is accurate. During WWII, the infamous Unit 731 of the Japanese army was reported to have repeatedly dropped plague-infested fleas over parts of China, including Manchuria. These drops were associated with subsequent outbreaks of plague in the targeted areas. Following WWII, the United States and the Soviet Union conducted programs of research on how to create aerosolized *Y. pestis* that could be used as a bioweapon to cause primary pneumonic plague. As mentioned above, plague was thought to be an excellent bioweapon due to the fact that in addition to causing infection in those inhaling the aerosol, significant numbers of secondary cases of primary pneumonic plague would also likely occur due to the contagious nature of the disease and person-to-person transmission via respiratory aerosol. Secondary reports of research conducted during that time suggest that organisms remain viable for up to 1 h and can be dispersed for

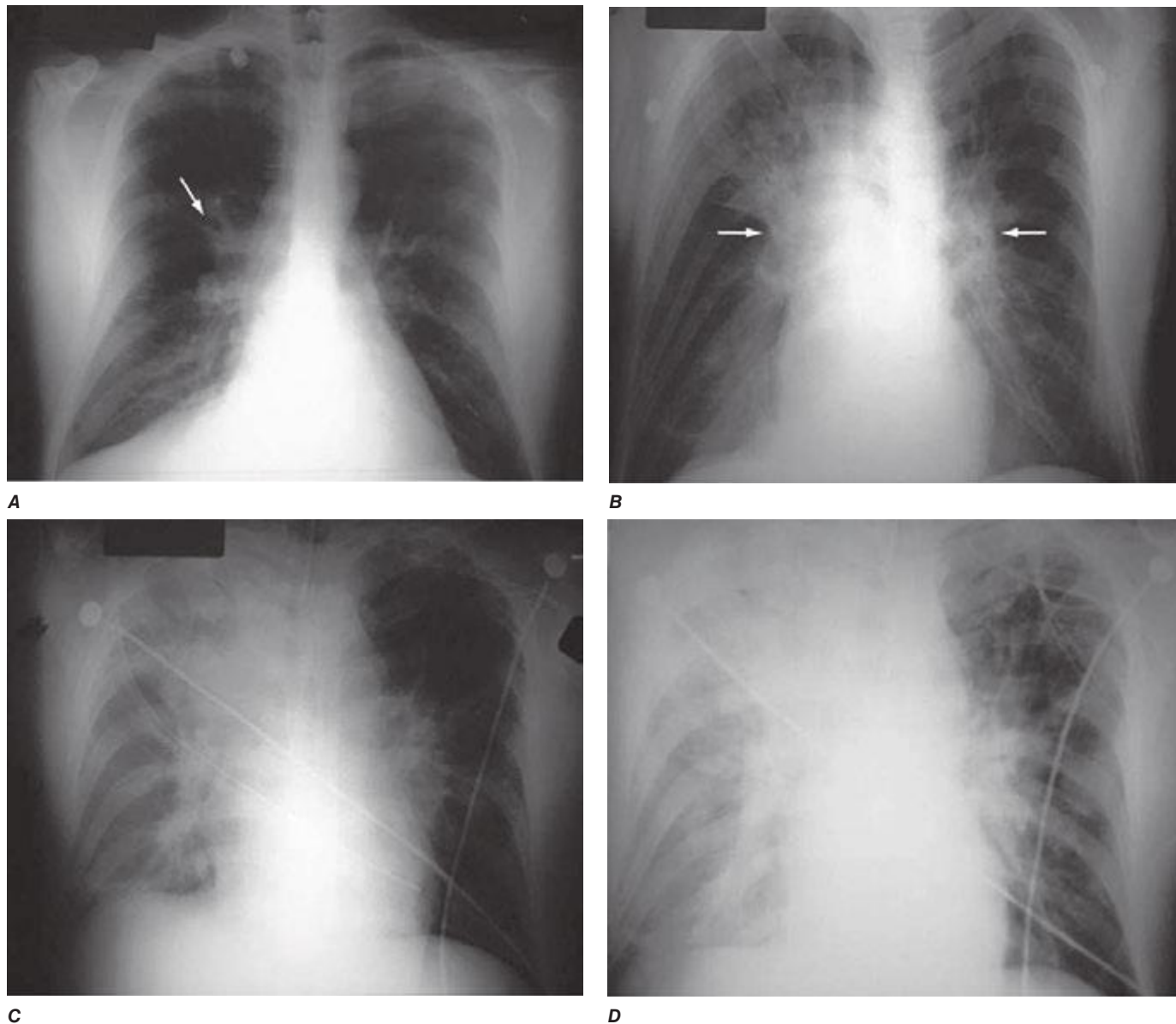


FIGURE 261e-3 Progression of chest x-ray findings in a patient with inhalational anthrax. Findings evolved from subtle hilar prominence and right perihilar infiltrate to a progressively widened mediastinum, marked perihilar infiltrates, peribronchial cuffing, and air bronchograms. (From L Borio et al: JAMA 286:2554, 2001; with permission.)

distances up to 10 km. Although the offensive bioweapons program in the United States was terminated prior to production of sufficient quantities of plague organisms for use as a weapon, it is believed that Soviet scientists did manufacture quantities sufficient for such a purpose. It has also been reported that more than 10 Soviet institutes and >1000 scientists were working with plague as a biologic weapon. Of concern is the fact that in 1995 a microbiologist in Ohio was arrested for having obtained *Y. pestis* in the mail from the American Type Culture Collection, using a credit card and a false letterhead. In the wake of this incident, the U.S. Congress passed a law in 1997 requiring that anyone intending to send or receive any of 42 different agents that could potentially be used as bioweapons first register with the CDC.

Microbiology and Clinical Features Plague is caused by *Y. pestis*, a non-motile, gram-negative bacillus that exhibits bipolar, or “safety pin,” staining with Wright, Giemsa, or Wayson stains. It has had a major impact on the course of history, thus adding to the element of fear evoked by its mention. The earliest reported plague epidemic was in 224 B.C. in China. The most infamous pandemic began in Europe in the fourteenth century, during which time one-third to one-half of the entire population of Europe was killed. During a plague outbreak in India in 1994, even though the number of confirmed cases was

relatively small, it is estimated that 500,000 individuals fled their homes in fear of this disease. In the first decade of the twenty-first century, 21,725 cases of plague were reported to the World Health Organization (WHO). Over 90% of these cases were from Africa, and the overall case fatality rate was 7.4%.

The clinical syndromes of plague generally reflect the mode of infection. *Bubonic plague* is the consequence of an insect bite; primary *pneumonic plague* arises through the inhalation of bacteria. Most of the plague seen in the world today is bubonic plague and is the result of a bite by a plague-infected flea. In part as a consequence of past pandemics, plague infection of rodents exists widely in nature, including in the southwestern United States, and each year thousands of cases of plague occur worldwide through contact with infected animals or fleas. Following inoculation of regurgitated bacteria into the skin by a flea bite, organisms travel through the lymphatics to regional lymph nodes, where they are phagocytized but not destroyed. Inside the cell, they multiply rapidly leading to inflammation, painful lymphadenopathy with necrosis, fever, bacteremia, septicemia, and death. The characteristic enlarged, inflamed lymph nodes, or *buboes*, give this form of plague its name. In some instances, patients may develop bacteremia without lymphadenopathy following infection, a condition referred to as *primary septicemic plague*. Extensive ecchymoses may develop due

TABLE 261e-3 CLINICAL SYNDROMES, PREVENTION, AND TREATMENT STRATEGIES FOR DISEASES CAUSED BY CATEGORY A AGENTS

Agent	Clinical Syndrome	Incubation Period	Diagnosis	Treatment	Prophylaxis
<i>Bacillus anthracis</i> (anthrax)	Cutaneous lesion: Papule to eschar Inhalational disease: Fever, malaise, chest and abdominal discomfort Pleural effusion, widened mediastinum on chest x-ray	1–12 days 1–60 days	Culture, Gram stain, PCR, Wright stain of peripheral smear	Postexposure: Ciprofloxacin, 500 mg, PO bid × 60 d <i>or</i> Doxycycline, 100 mg PO bid × 60 d <i>or</i> Amoxicillin, 500 mg PO q8h × 60 d, likely to be effective if strain is penicillin sensitive <i>Active disease:</i> Ciprofloxacin, 400 mg IV q12h <i>or</i> doxycycline, 100 mg IV q12h <i>plus</i> Clindamycin, 900 mg IV q8h and/or rifampin, 300 mg IV q12h; switch to PO when stable × 60 d total <i>plus</i> <i>Antitoxin</i> Raxibacumab, 40 mg/kg IV over 2.25 h; diphenhydramine to reduce reaction	Anthrax vaccine adsorbed Recombinant protective antigen vaccines are under study Raxibacumab when alternative therapies are not available or appropriate
<i>Yersinia pestis</i> (pneumonic plague)	Fever, cough, dyspnea, hemoptysis Infiltrates and consolidation on chest x-ray	1–6 days	Culture, Gram stain, direct fluorescent antibody, PCR	Gentamicin, 2.0 mg/kg IV loading then 1.7 mg/kg q8h IV <i>or</i> Streptomycin, 1.0 g q12h IM or IV Alternatives include doxycycline, 100 mg bid PO or IV; chloramphenicol, 500 mg qid PO or IV	Doxycycline, 100 mg PO bid <i>or</i> Levofloxacin, 500 mg PO daily Formalin-fixed vaccine (FDA licensed; not available)
<i>Variola major</i> (smallpox)	Fever, malaise, headache, backache, emesis Maculopapular to vesicular to pustular skin lesions	7–17 days	Culture, PCR, electron microscopy	Supportive measures; consideration for cidofovir, tecovirimat, antivaccinia immunoglobulin	Vaccinia immunization
<i>Francisella tularensis</i> (tularemia)	Fever, chills, malaise, myalgia, chest discomfort, dyspnea, headache, skin rash, pharyngitis, conjunctivitis Hilar adenopathy on chest x-ray	1–14 days	Gram stain, culture, immunohistochemistry, PCR	Streptomycin, 1 g IM bid <i>or</i> Gentamicin, 5 mg/kg per day div q8h IV for 14 days <i>or</i> Doxycycline, 100 mg IV bid <i>or</i> Chloramphenicol, 15 mg/kg up to 1 g IV qid <i>or</i> Ciprofloxacin, 400 mg IV bid	Doxycycline, 100 mg PO bid × 14 days <i>or</i> Ciprofloxacin, 500 mg PO bid × 14 days
Viral hemorrhagic fevers	Fever, myalgia, rash, encephalitis, prostration	2–21 days	RT-PCR, serologic testing for antigen or antibody Viral isolation by CDC or U.S. Army Medical Research Institute of Infectious Diseases (USAMRIID)	Supportive measures Ribavirin 30 mg/kg up to 2 g × 1, followed by 16 mg/kg IV up to 1 g q6h for 4 days, followed by 8 mg/kg IV up to 0.5 g q8h × 6 days	No known chemoprophylaxis Consideration for ribavirin or monoclonal antibodies in high-risk situations
Botulinum toxin (<i>Clostridium botulinum</i>)	Dry mouth, blurred vision, ptosis, weakness, dysarthria, dysphagia, dizziness, respiratory failure, progressive paralysis, dilated pupils	12–72 h	Mouse bioassay, toxin immunoassay	Supportive measures including ventilation, HBAT equine antitoxin from the CDC Emergency Operations Center, 770-488-7100	Administration of antitoxin

Abbreviations: CDC, U.S. Centers for Disease Control and Prevention; FDA, U.S. Food and Drug Administration; HBAT, heptavalent botulinum antitoxin; PCR, polymerase chain reaction; RT-PCR, reverse transcriptase polymerase chain reaction.

to disseminated intravascular coagulation, and gangrene of the digits and/or nose may develop in patients with advanced septicemic plague. It is thought that this appearance of some patients gave rise to the term *Black Death* in reference to the plague epidemic of the fourteenth and fifteenth centuries. Some patients may develop pneumonia (secondary pneumonic plague) as a complication of bubonic or septicemic plague. These patients may then transmit the agent to others via the respiratory route, causing cases of primary pneumonic plague. Primary pneumonic plague is the manifestation most likely to occur as the result of a bioterrorist attack, with an aerosol of bacteria spread over a wide area or a particular environment that is densely populated. In this setting, patients would be expected to develop fever, cough with hemoptysis, dyspnea, and gastrointestinal symptoms 1–6 days following exposure. Clinical features of pneumonia would be accompanied by pulmonary

infiltrates and consolidation on chest x-ray. In the absence of antibiotics, the mortality rate of this form of plague is on the order of 85%, and death usually occurs within 2–6 days.

TREATMENT PLAGUE

Streptomycin, tetracycline, doxycycline, and levofloxacin are licensed by the U.S. Food and Drug Administration (FDA) for the treatment of plague. Levofloxacin was approved for this indication in 2012 via the animal rule (see below). Multiple additional antibiotics licensed for other infections are commonly used and are likely effective. Among these are aminoglycosides such as gentamicin, cephalosporins, trimethoprim/sulfamethoxazole, chloramphenicol, and ciprofloxacin (Table 261e-3). A multidrug-resistant strain of

Y. pestis was identified in 1995 from a patient with bubonic plague in Madagascar. Although this organism was resistant to streptomycin, ampicillin, chloramphenicol, sulfonamides, and tetracycline, it retained its susceptibility to other aminoglycosides and cephalosporins. Given the subsequent identification of a similar organism in 1997 coupled with the fact that this resistance is plasmid-mediated, it seems likely that genetically modifying *Y. pestis* to a multidrug-resistant form is possible. Unlike patients with inhalational anthrax (see above), patients with pulmonary plague should be cared for under conditions of strict respiratory isolation comparable to that used for multidrug-resistant tuberculosis.

Vaccination and Prevention A formalin-fixed, whole-organism vaccine was licensed by the FDA for the prevention of plague. That vaccine is no longer being manufactured, but its potential value as a current countermeasure against bioterrorism would likely have been modest at best as it was ineffective against animal models of primary pneumonic plague. Efforts are under way to develop a second generation of vaccines that will protect against aerosol challenge. Among the candidates being tested are recombinant forms of the fraction 1 capsular (F1) antigen and the virulence component of the type III secretion apparatus (V) antigen of *Y. pestis*. It is likely that doxycycline or levofloxacin would provide coverage in a chemoprophylaxis setting. Unlike the case with anthrax, in which one has to be concerned about the persistence of ungerminated spores in the respiratory tract, the duration of prophylaxis against plague need only extend to 7 days following exposure.

SMALLPOX

See also Chap. 220e.

Variola Virus as a Bioweapon Given that most of the world's population was once vaccinated against smallpox, variola virus would not have been considered a good candidate as a bioweapon 30 years ago. However, with the cessation of immunization programs in the United States in 1972 and throughout the world in 1980 due to the successful global eradication of smallpox, close to 50% of the U.S. population is fully susceptible to smallpox today. Given its infectious nature and the 10–30% mortality rate in unimmunized individuals, the deliberate spread of this virus could have a devastating effect on our society and unleash a previously conquered deadly disease. It is estimated that an initial infection of 50–100 persons in a first generation of cases could expand by a factor of 10–20 with each succeeding generation in the absence of any effective containment measures. Although the likely implementation of an effective public health response makes this scenario unlikely, it does illustrate the potential damage and disruption that can result from a smallpox outbreak.

In 1980, the WHO recommended that all immunization programs be terminated; that representative samples of variola virus be transferred to two locations, one at the CDC in Atlanta, GA, in the United States and the other at the Institute of Virus Preparations in the Soviet Union; and that all other stocks of smallpox be destroyed. Several years later, it was recommended that these two authorized collections be destroyed. However, these latter recommendations were placed on hold in the wake of increased concerns on the use of variola virus as a biologic weapon and thus the need to maintain an active program of defensive research. Many of these concerns were based on allegations made by former Soviet officials that extensive programs had been in place in that country for the production and weaponization of large quantities of smallpox virus. The dismantling of these programs with the fall of the Soviet Union and the subsequent weakening of security measures led to fears that stocks of *Variola major* may have made their way to other countries or terrorist organizations. In addition, accounts that efforts had been taken to produce recombinant strains of *Variola* that would be more virulent and more contagious than the wild-type virus have led to an increase in the need to be vigilant for the reemergence of this often fatal infectious disease.

Microbiology and Clinical Features Smallpox is caused by one of two variants of variola virus, *V. major* and *V. minor*. Variola is a double-strand DNA virus and member of the *Orthopoxvirus* genus of the

Poxviridae family. Infections with *V. minor* are generally less severe than those of *V. major*, with milder constitutional symptoms and lower mortality rates; thus *V. major* is the only one considered to be a viable bioweapon. Infection with *V. major* typically occurs following contact with an infected person. Patients are infectious from the time that a maculopapular rash appears on the skin and oropharynx through the resolution and scabbing of the pustular lesions. Infection occurs principally during close contact, through the inhalation of saliva droplets containing virus from the oropharyngeal exanthem. Aerosolized material from contaminated clothing or linen can also spread infection. Several days after exposure, a primary viremia is believed to occur that results in dissemination of virus to lymphoid tissues. A secondary viremia occurs ~4 days later that leads to localization of infection in the dermis. Approximately 12–14 days following the initial exposure, the patient develops high fever, malaise, vomiting, headache, backache, and a maculopapular rash that begins on the face and extremities and spreads to the trunk (centripetal) with lesions in the same developmental stage in any given location. This is in contrast to the rash of varicella (chickenpox) that begins on the trunk and face and spreads to the extremities (centrifugal) with lesions at all stages of development. The lesions are initially maculopapular and evolve to vesicles that eventually become pustules and then scabs. The oral mucosa also develops maculopapular lesions that evolve to ulcers. The lesions appear over a period of 1–2 days and evolve at the same rate. Although virus can be isolated from the scabs on the skin, the conventional thinking is that once the scabs have formed the patient is no longer contagious. Smallpox is associated with 10–30% mortality rates, with patients typically dying of severe systemic illness during the second week of symptoms. Historically, ~5–10% of naturally occurring smallpox cases take either of two highly virulent atypical forms, classified as *hemorrhagic* and *malignant*. These are difficult to diagnose because of their atypical presentations. The hemorrhagic form is uniformly fatal and begins with the relatively abrupt onset of a severely prostrating illness characterized by high fevers and severe headache and back and abdominal pain. This form of the illness resembles a severe systemic inflammatory syndrome, in which patients have a high viremia but die without developing the characteristic rash. Cutaneous erythema develops accompanied by petechiae and hemorrhages into the skin and mucous membranes. Death usually occurs within 5–6 days. The malignant, or “flat,” form of smallpox is frequently fatal and has an onset similar to the hemorrhagic form, but with confluent skin lesions developing more slowly and never progressing to the pustular stage.

TREATMENT SMALLPOX

Given the infectious nature of smallpox and the extreme vulnerability of contemporary society, patients who are suspected cases should be handled with strict isolation procedures. Although laboratory confirmation of a suspected case by culture, polymerase chain reaction (PCR), and electron microscopy is essential, it is equally important that appropriate precautions be used when obtaining samples for culture and laboratory testing. All health care and laboratory workers caring for patients should have been recently immunized with vaccinia, and all samples should be transported in doubly sealed containers. Patients should be cared for in negative-pressure rooms with strict isolation precautions.

There is no licensed specific therapy for smallpox, and historic treatments have focused solely on supportive care. Although several antiviral agents, including cidofovir, that are licensed for other diseases have in vitro activity against *V. major*, they have never been tested in the setting of human disease. For this reason, it is difficult to predict whether or not they would be effective in cases of smallpox and, if effective, whether or not they would be of value in patients with advanced disease. Agents currently being studied as possible antiviral compounds against *V. major* include a viral egress inhibitor (tecovirimat, ST-246, or Arestvyr) and a lipid-conjugated form of cidofovir (brincidofovir, CMX001).

TABLE 261e-4 COMPLICATIONS FROM 438,134 ADMINISTRATIONS OF VACCINIA DURING THE U.S. DEPARTMENT OF DEFENSE (DOD) SMALLPOX IMMUNIZATION CAMPAIGN INITIATED IN DECEMBER 2002

Complication	Number of Cases	DoD Rate per Million Vaccinees (95% Confidence Interval)	Historic Rate Per Million Vaccinees
Mild or temporary:			
Generalized vaccinia, mild	35	67 (52, 85)	45–212 ^a
Inadvertent inoculation, self	62	119 (98, 142)	606 ^a
Vaccinia transfer to contact	28	53 (40, 69)	8–27 ^a
Moderate or serious:			
Encephalitis	1	2.2 (0.6, 7.2)	2.6–8.7 ^a
Acute myopericarditis	69	131 (110, 155)	100 ^b
Eczema vaccinatum	0	0 (0, 3.7)	2–35 ^a
Progressive vaccinia	0	0 (0, 3.7)	1–7 ^a
Death	1 ^c	1.9 (0.2, 5.6)	1–2 ^a

^aBased on adolescent and adult smallpox vaccinations from 1968 studies, both primary and revaccinations. ^bBased on case series in Finnish military recruits given the Finnish strain of smallpox vaccine. ^cPotentially attributable to vaccination; after lupus-like illness.

Source: From JD Grabenstein and W Winkenwerder: <http://www.smallpox.mil/event/SPSafetySum.asp>.

Vaccination and Prevention In 1796, Edward Jenner demonstrated that deliberate infection with cowpox virus could prevent illness on subsequent exposure to smallpox. Today, smallpox is a preventable disease following immunization with vaccinia. The current dilemma facing our society regarding assessment of the risk and benefit of smallpox vaccination is that the degree of risk that someone will deliberately and effectively release smallpox into our society is unknown. Given that there are well-described risks associated with vaccination, the degree of risk/benefit for the general population does not favor immunization. As a prudent first step in preparedness for a smallpox attack, however, members of the U.S. armed services received primary or booster immunizations with vaccinia before 1990 and after 2002. In addition, a number of civilian health care workers who comprise smallpox-response teams at the state and local public health level have been vaccinated.

Initial fears regarding the immunization of a segment of the American population with vaccinia when there are more individuals receiving immunosuppressive drugs and other immunocompromised patients than ever before were dispelled by the data generated from the military and civilian immunization campaigns of 2002–2004. Adverse event rates for the first 450,000 immunizations were similar to and, in certain categories of adverse events, even lower than those from prior historic data, in which most severe sequelae of vaccination occurred in young infants (Table 261e-4). In addition, 11 patients with early-stage HIV infection were inadvertently immunized without problem. One significant concern during that immunization campaign, however, was the description of a syndrome of myopericarditis, which had not been appreciated during prior immunization campaigns with vaccinia. In an effort to provide a safer vaccine to protect against smallpox, ACAM 2000, a cloned virus propagated in tissue culture, was developed and became the first second-generation smallpox vaccine to be licensed. This vaccine is now the only vaccinia product currently licensed in the United States and has been used by the U.S. military since 2008. It is part of the U.S. government stockpile. Research continues on attenuated forms of vaccinia such as modified vaccinia Ankara (MVA). Vaccinia immune globulin is available to treat those who experience a severe reaction to immunization with vaccinia.

TULAREMIA

See also Chap. 195.

Francisella tularensis as a Bioweapon Tularemia has been studied as an agent of bioterrorism since the mid-twentieth century. It has been speculated by some that the outbreak of tularemia among German and Soviet soldiers during fighting on the Eastern Front during WWII was the consequence of a deliberate release. Unit 731 of the Japanese army studied the use of tularemia as a bioweapon during WWII. Large

preparations were made for mass production of *F. tularensis* by the United States, but no stockpiling of any agent took place. Stocks of *F. tularensis* were reportedly generated by the Soviet Union in the mid-1950s. It has also been suggested that the Soviet program extended into the era of molecular biology and that some strains were engineered to be resistant to common antibiotics. *F. tularensis* is an extremely infectious organism, and human infections have occurred from merely examining an uncovered petri dish streaked with colonies. Given these facts, it is reasonable to conclude that this organism might be used as a bioweapon through either an aerosol or contamination of food or drinking water.

Microbiology and Clinical Features Although similar in many ways to anthrax and plague, tularemia, also referred to as rabbit fever or deer fly fever, is neither as lethal nor as fulminant as either of these other two category A bacterial infections. It is, however, extremely infectious, and as few as 10 organisms can lead to estab-

lishment of infection. Despite this fact, it is not spread from person to person. Tularemia is caused by *F. tularensis*, a small, nonmotile, gram-negative coccobacillus. Although it is not a spore-forming organism, it is a hardy bacterium that can survive for weeks in the environment. Infection typically comes from insect bites or contact with organisms in the environment. Infections have occurred in laboratory workers studying the agent. Large waterborne outbreaks have been recorded. It is most likely that the outbreak among German and Russian soldiers and Russian civilians noted above during WWII represented a large waterborne tularemia outbreak in a *Tularensis*-enzootic area devastated by warfare.

Humans can become infected through a variety of environmental sources. Infection is most common in rural areas where a variety of small mammals may serve as reservoirs. Human infections in the summer are often the result of insect bites from ticks, flies, or mosquitoes that have bitten infected animals. In colder months, infections are most likely the result of direct contact with infected mammals and are most common in hunters. In these settings, infection typically presents as a systemic illness with an area of inflammation and necrosis at the site of tissue entry. Drinking of contaminated water may lead to an oropharyngeal form of tularemia characterized by pharyngitis with cervical and/or retropharyngeal lymphadenopathy (Chap. 195). The most likely mode of dissemination of tularemia as a biologic weapon would be as an aerosol, as has occurred in a number of natural outbreaks in rural areas, including Martha's Vineyard in the United States. Approximately 1–14 days following exposure by this route, one would expect to see inflammation of the airways with pharyngitis, pleuritis, and bronchopneumonia. Typical symptoms would include the abrupt onset of fever, fatigue, chills, headache, and malaise (Table 261e-3). Some patients might experience conjunctivitis with ulceration, pharyngitis, and/or cutaneous exanthems. A pulse-temperature dissociation might be present. Approximately 50% of patients would show a pulmonary infiltrate on chest x-ray. Hilar adenopathy might also be present, and a small percentage of patients could have adenopathy without infiltrates. The highly variable presentation makes acute recognition of aerosol-disseminated tularemia very difficult. The diagnosis would likely be made by immunohistochemistry, molecular techniques, or culture of infected tissues or blood. Untreated, mortality rates range from 5 to 15% for cutaneous routes of infection and from 30 to 60% for infection by inhalation. Since the advent of antibiotic therapy, these rates have dropped to <2%.

TREATMENT TULAREMIA

Both streptomycin and doxycycline are licensed for treatment of tularemia. Other agents likely to be effective include gentamicin, chloramphenicol, and ciprofloxacin (Table 261e-3). Given the potential for

genetic modification of this organism to yield antibiotic-resistant strains, broad-spectrum coverage should be the rule until sensitivities have been determined. As mentioned above, special isolation procedures for patients are not required.

Vaccination and Prevention There are no vaccines currently licensed for the prevention of tularemia. Although a live, attenuated strain of the organism has been used in the past with some reported success, there are inadequate data to support its widespread use at this time. Development of a vaccine for this agent is an important part of the current biodefense research agenda. In the absence of an effective vaccine, postexposure chemoprophylaxis with either doxycycline or ciprofloxacin appears to be a reasonable approach (Table 261e-3).

VIRAL HEMORRHAGIC FEVERS

See also Chaps. 233 and 234.

Hemorrhagic Fever Viruses as Bioweapons Several of the hemorrhagic fever viruses have been reported to have been weaponized by the Soviet Union and the United States. Nonhuman primate studies indicate that infection can be established with very few virions and that infectious aerosol preparations can be produced. Under the guise of wanting to aid victims of an Ebola outbreak, members of the Aum Shinrikyo cult in Japan were reported to have traveled to central Africa in 1992 in an attempt to obtain Ebola virus for use in a bioterrorist attack. Thus, although there has been no evidence that these agents have ever been used in a biologic attack, there is clear interest in their potential for this purpose.

Microbiology and Clinical Features The viral hemorrhagic fevers are a group of illnesses caused by any one of a number of similar viruses (Table 261e-2). These viruses are all enveloped, single-strand RNA viruses that are thought to depend on a host reservoir for long-term survival. Although rodents or insects have been identified as the hosts for some of these viruses, for others the hosts are unknown. These viruses tend to be geographically restricted according to the migration patterns of their hosts. Great apes are not a natural reservoir for Ebola virus, but large numbers of these animals in sub-Saharan Africa have died from Ebola infection over the past decade. Humans can become infected with hemorrhagic fever viruses if they come into contact with an infected host or other infected animals. Person-to-person transmission, largely through direct contact with virus-containing body fluids, has been documented for Ebola, Marburg, and Lassa viruses and rarely for the New World arenaviruses. Although there is no clear evidence of respiratory spread among humans, these viruses have been shown in animal models to be highly infectious by the aerosol route. This, coupled with mortality rates as high as 90%, makes them excellent candidate agents of bioterrorism.

The clinical features of the viral hemorrhagic fevers vary depending on the particular agent (Table 261e-3). Initial signs and symptoms typically include fever, myalgia, prostration, and disseminated intravascular coagulation with thrombocytopenia and capillary hemorrhage. These findings are consistent with a cytokine-mediated systemic inflammatory syndrome. A variety of different maculopapular or erythematous rashes may be seen. Leukopenia, temperature-pulse dissociation, renal failure, and seizures may also be part of the clinical presentation.

Outbreaks of most of these diseases are sporadic and unpredictable. As a consequence, most studies of pathogenesis have been performed using laboratory animals. The diagnosis should be suspected in anyone with temperature $>38.3^{\circ}\text{C}$ for <3 weeks who also exhibits at least two of the following: hemorrhagic or purpuric rash, epistaxis, hematemesis, hemoptysis, or hematochezia in the absence of any other identifiable cause. In this setting, samples of blood should be sent after consultation to the CDC or the USAMRIID for serologic testing for antigen and antibody as well as reverse transcriptase polymerase chain reaction (RT-PCR) testing for hemorrhagic fever viruses. All samples should be handled with double-bagging. Given how little is known regarding the human-to-human transmission of these viruses, appropriate isolation measures would include full barrier precautions with negative-pressure rooms and use of powered air-purifying respirators (PAPRs).

Unprotected skin contact with cadavers has been implicated in the transmission of certain hemorrhagic fever viruses such as Ebola, so it is recommended that autopsies of suspected cases be performed using the strictest measures for protection and that burial or cremation be performed promptly without embalming.

TREATMENT VIRAL HEMORRHAGIC FEVERS

There are no approved and effective antiviral therapies for this class of viruses (Table 261e-3). Although there are anecdotal reports of the efficacy of ribavirin, interferon- α , or hyperimmune immunoglobulin, definitive data are lacking. The best data for ribavirin are in arenavirus (Lassa and New World) infections. In some in vitro systems, specific immunoglobulin has been reported to enhance infectivity, and thus these potential treatments must be approached with caution.

Vaccination and Prevention There are no licensed and effective vaccines for these agents. Studies are currently under way examining the potential role of DNA, recombinant viruses, and attenuated viruses as vaccines for several of these infections. Among the most promising at present are vaccines for Argentine, Ebola, Rift Valley, and Kyasanur Forest viruses. A series of monoclonal antibodies directed against the envelope glycoproteins of Ebola have demonstrated protection against infection in a postexposure setting in nonhuman primates and are being further developed for human use.

BOTULISM TOXIN (*CLOSTRIDIUM BOTULINUM*)

See also Chap. 178.

Botulinum Toxin as a Bioweapon In a bioterrorist attack, botulinum toxin would likely be dispersed as an aerosol or as contamination of a food supply. Although contamination of a water supply is possible, it is likely that any toxin would be rapidly inactivated by the chlorine used to purify drinking water. Similarly, toxin can be inactivated by heating any food to $>85^{\circ}\text{C}$ for >5 min. Without external facilitation, the environmental decay rate is estimated at 1% per minute, and thus the time interval between weapon release and ingestion or inhalation needs to be rather short. The Japanese biologic warfare group, Unit 731, is reported to have conducted experiments on botulinum poisoning in prisoners in the 1930s. The United States and the Soviet Union both acknowledged producing botulinum toxin, and there is some evidence that the Soviet Union attempted to create recombinant bacteria containing the gene for botulinum toxin. In records submitted to the United Nations, Iraq admitted to having produced 19,000 L of concentrated toxin—enough toxin to kill the entire population of the world three times over. By many accounts, botulinum toxin was the primary focus of the pre-1991 Iraqi bioweapons program. In addition to these examples of state-supported research into the use of botulinum toxin as a bioweapon, the Aum Shinrikyo cult unsuccessfully attempted on at least three occasions to disperse botulinum toxin into the civilian population of Tokyo.

Microbiology and Clinical Features Unique among the category A agents for not being a live microorganism, botulinum toxin is one of the most potent toxins ever described and is thought by some to be the most poisonous substance in existence. It is estimated that 1 g of botulinum toxin would be sufficient to kill 1 million individuals if adequately dispersed. Botulinum toxin is produced by the gram-positive, spore-forming anaerobe *C. botulinum* (Chap. 178). Its natural habitat is soil. There are seven antigenically distinct forms of botulinum toxin, designated A–G. The majority of naturally occurring human cases are of types A, B, and E. Antitoxin directed toward one of these will have little to no activity against the others. The toxin is a 150-kDa zinc-containing protease that prevents the intracellular fusion of acetylcholine vesicles with the motor neuron membrane, thus preventing the release of acetylcholine. In the absence of acetylcholine-dependent triggering of muscle fibers, a flaccid paralysis develops. Although botulism does

not spread from person to person, the ease of production of the toxin coupled with its high morbidity and 60–100% mortality make it a close to ideal bioweapon.

Botulism can result from the growth of *C. botulinum* infection in a wound or the intestine, the ingestion of contaminated food, or the inhalation of aerosolized toxin. The latter two forms are the most likely modes of transmission for bioterrorism. Once toxin is absorbed into the bloodstream, it binds to the neuronal cell membrane, enters the cell, and cleaves one of the proteins required for the intracellular binding of the synaptic vesicle to the cell membrane, thus preventing release of the neurotransmitter to the membrane of the adjacent muscle cell. Patients initially develop multiple cranial nerve palsies that are followed by a descending flaccid paralysis. The extent of the neuromuscular compromise is dependent on the level of toxemia. The majority of patients experience diplopia, dysphagia, dysarthria, dry mouth, ptosis, dilated pupils, fatigue, and extremity weakness. There are minimal true central nervous system effects, and patients rarely show significant alterations in mental status. Severe cases can involve complete muscular collapse, loss of the gag reflex, and respiratory failure. Recovery requires the regeneration of new motor neuron synapses with the muscle cell, a process that can take weeks to months. In the absence of secondary infections, which may be common during the protracted recovery phase of this illness, patients remain afebrile. The diagnosis is suspected on clinical grounds and confirmed by a mouse bioassay or toxin immunoassay.

TREATMENT BOTULISM

Treatment for botulism is mainly supportive and may require intubation, mechanical ventilation, and parenteral nutrition (Table 261e-3). If diagnosed early enough, administration of equine antitoxin may reduce the extent of nerve injury and decrease the severity of disease. At present, a heptavalent botulinum antitoxin (HBAT) is available through the CDC as an investigational agent for treatment of naturally occurring noninfant botulism. HBAT contains horse serum-derived antibody fragments to all seven known botulinum toxins (A–G). It is composed of <2% intact immunoglobulin and ≥90% Fab and F(ab)₂ immunoglobulin fragments. A single dose of antitoxin is usually adequate to neutralize any circulating toxin. Repeat dosing may be needed in a setting of continued toxin exposure. Given that this product is derived from horse serum, one needs to be vigilant for hypersensitivity reactions, including serum sickness and anaphylaxis following its administration. Once the damage to the nerve axon has been done, however, there is little possible in the way of specific therapy. At this point, vigilance for secondary complications such as infections during the protracted recovery phase is of the utmost importance. Due to their ability to worsen neuromuscular blockade, aminoglycosides and clindamycin should be avoided in the treatment of these infections.

Vaccination and Prevention A botulinum toxoid preparation has been used as a vaccine for laboratory workers at high risk of exposure and in certain military situations; however, it is not currently available in quantities that could be used for the general population. At present, early recognition of the clinical syndrome and use of appropriate equine antitoxin is the mainstay of prevention of full-blown disease in exposed individuals. The development of human monoclonal antibodies as a replacement for equine antitoxin antibodies is an area of active research interest.

CATEGORY B AND C AGENTS

The category B agents include those that are easy or moderately easy to disseminate and result in moderate morbidity and low mortality rates. A listing of the current category B agents is provided in Table 261e-2. As can be seen, it includes a wide array of microorganisms and products of microorganisms. Several of these agents have been used in bioterrorist attacks, although never with the impact of the category A

agents described above. Among the more notorious of these was the contamination of salad bars in Oregon in 1984 with *Salmonella typhimurium* by the religious cult Rajneeshee. In this outbreak, which many consider to be the first bioterrorist attack against U.S. citizens, >750 individuals were poisoned and 40 were hospitalized in an effort to influence a local election. The intentional nature of this outbreak went unrecognized for more than a decade.

Category C agents are the third highest priority agents in the bio-defense agenda. These agents include emerging pathogens to which little or no immunity exists in the general population, such as the severe acute respiratory syndrome (SARS) or Middle East respiratory syndrome (MERS) coronavirus or pandemic-potential strains of influenza that could be obtained from nature and deliberately disseminated. These agents are characterized as being relatively easy to produce and disseminate, having high morbidity and mortality rates, and having a significant public health impact. There is no running list of category C agents at the present time.

PREVENTION AND PREPAREDNESS

As noted above, a large number and diverse array of agents have the potential to be used in a bioterrorist attack. In contrast to the military situation with biowarfare, where the primary objective is to inflict mass casualties on a healthy and prepared militia, the objectives of bioterrorism are to harm civilians as well as to create fear and disruption among the civilian population. Although the military need only to prepare their troops to deal with the limited number of agents that pose a legitimate threat of biowarfare, the public health system needs to prepare the entire civilian population to deal with the multitude of agents and settings that could be used in a bioterrorism attack. This includes anticipating issues specific to the very young and the very old, the pregnant patient, and the immunocompromised individual. The challenges in this regard are enormous and immediate. Whereas military preparedness emphasizes vaccines toward a limited number of agents, civilian preparedness needs to rely on rapid diagnosis and treatment of a wide array of conditions.

The medical profession must maintain a high index of suspicion that unusual clinical presentations or the clustering of cases of a rare disease may not be a chance occurrence but rather the first sign of a bioterrorist event. This is particularly true when such diseases occur in traditionally healthy populations, when surprisingly large numbers of rare conditions occur, and when diseases commonly seen in rural settings appear in urban populations. Given the importance of rapid diagnosis and early treatment for many of these conditions, it is essential that the medical care team report any suspected cases of bioterrorism immediately to local and state health authorities and/or to the CDC (888-246-2675). Enhancements made to the U.S. public health surveillance network after the anthrax attacks of 2001 have greatly facilitated the rapid sharing of information among public health agencies.

A series of efforts are in place to ensure the biomedical security of the civilian population of the United States. The Public Health Service has a more highly trained, fully deployable force. The Strategic National Stockpile (SNS) maintained by the CDC provides rapid access to quantities of pharmaceuticals, antidotes, vaccines, and other medical supplies that may be of value in the event of biologic or chemical terrorism. The SNS has two basic components. The first of these consists of “push packages” that can be deployed anywhere in the United States within 12 h. These push packages are a preassembled set of supplies, pharmaceuticals, and medical equipment ready for immediate delivery to the field. Given that an actual threat may not have been precisely identified at the time of stockpile deployment, they provide treatment for a variety of conditions. The contents of the push packs are constantly updated to ensure that they reflect current needs as determined by national security risk assessments; they include antibiotics for treatment of anthrax, plague, and tularemia as well as a cache of vaccine to deal with a smallpox threat. The second component of the SNS comprises inventories managed by specific vendors and consists of the provision of additional pharmaceuticals, supplies, and/or products tailored to the specific attack.

The number of FDA-approved and licensed drugs and vaccines for category A and B agents is currently limited and not reflective of the pharmacy of today. In an effort to speed the licensure of additional drugs and vaccines for these diseases, the FDA has a rule for the licensure of such countermeasures against agents of bioterrorism when adequate and well-controlled clinical efficacy studies cannot be ethically conducted in humans. This is commonly referred to as the “Animal Rule.” Thus, for indications in which field trials of prophylaxis or therapy for a naturally occurring disease are not feasible, the FDA will rely on evidence solely from laboratory animal studies. For this rule to apply, it must be shown that (1) there are reasonably well-understood pathophysiologic mechanisms for the condition and its treatment; (2) the effect of the intervention is independently substantiated in at least two animal species, including species expected to react with a response predictive for humans; (3) the animal study endpoint is clearly related to the desired benefit in humans; and (4) the data in animals allow selection of an effective dose in humans. As noted above, levofloxacin for treatment of plague and raxibacumab for treatment of inhalational anthrax have been licensed via this mechanism.

Finally, the Biomedical Advanced Research and Development Authority (BARDA) was established in 2006 within the U.S. Department

of Health and Human Services to provide an integrated, systematic approach to the development and purchase of the necessary vaccines, drugs, therapies, and diagnostic tools for public health medical emergencies. As authorized by the All-Hazards Preparedness Reauthorization Act of 2013 in conjunction with the Project Bioshield Act of 2006 and the Pandemic and All Hazards Act of 2006, BARDA manages a series of initiatives designed to facilitate biodefense research within the federal government, create a stable source of funding for the purchase of countermeasures against agents of bioterrorism, and create a category of “emergency use authorization” to allow the FDA to approve the use of unlicensed countermeasures during times of extraordinary unmet needs, as might be present in the context of a bioterrorist attack.

Although the prospect of a deliberate attack on civilians with disease-producing agents may seem to be an act of incomprehensible evil, history shows us that it is something that has been done in the past and will likely be done again in the future. It is the responsibility of health care providers to be aware of this possibility, to be able to recognize early signs of a potential bioterrorist attack and alert the public health system, and to respond quickly to provide care to the individual patient. Among the websites with current information on microbial bioterrorism are www.bt.cdc.gov, www.niaid.nih.gov, and www.cidrap.umn.edu.

262e Chemical Terrorism

Charles G. Hurst, Jonathan Newmark,
James A. Romano, Jr.

The use of chemical warfare agents (CWAs) in modern warfare dates back to World War I (WWI). Sulfur mustard and nerve agents were used by Iraq against the Iranian military and Kurdish civilians. Most recently the nerve agent Sarin, GB, was used by the Syrian military against their civilian population. Since the Japanese sarin attacks in 1994–1995 and the terrorist strikes of September 11, 2001, the all-too-real possibility of chemical or biological terrorism against civilian populations anywhere in the world has attracted increased attention.

Military planners consider the WWI blistering agent sulfur mustard and the organophosphorus nerve agents as the most likely agents to be used on the battlefield. In a civilian or terrorist scenario, the choice widens considerably. For example, many of the CWAs of WWI, including chlorine, phosgene, and cyanide, are used today in large amounts in industry. They are produced in chemical plants, are stockpiled in large tanks, and travel up and down highways and railways in large tanker cars. The rupture of any of these stores by accident or on purpose could cause many injuries and deaths. In three attacks in February 2007, for example, insurgents in Iraq used chlorine gas released from tankers after explosions as a crude form of chemical weaponry; these attacks killed 12 people and intoxicated more than 140 others. Countless hazardous materials (HAZMATs) that are not used on the battlefield can be used as terrorist weapons. Some of them, including insecticides and ammonia, could wreak as much damage and injury as the weaponized chemical agents.

Many mistakenly believe that chemical attacks will always be so severe that little can be done except to bury the dead. History proves the opposite. Even in WWI, when IV fluids, endotracheal tubes, and antibiotics were unavailable, the mortality rate among U.S. forces on the battlefield from CWAs—chiefly sulfur mustard and the pulmonary intoxicants—was only 1.9%. That figure was far lower than the 7% mortality rate from conventional wounds. In the 1995 Tokyo subway sarin incident, among the 5500 patients who sought medical attention at hospitals, 80% were not actually symptomatic and only 12 died. Recent events should prompt not a fatalistic attitude but a realistic wish to understand the pathophysiology of the syndromes these agents cause, with a view to treating expeditiously all patients who present for care and an expectation of saving the vast majority. As we prepare to defend our civilian population from the effects of chemical terrorism, we also must consider the fact that terrorism itself can produce sequelae such as physiologic or neurologic effects that may resemble the effects of nonlethal exposures to CWAs. These effects are due to a general fear of chemicals, fear of decontamination, fear of protective ensembles, or other phobic reactions. The increased difficulty in differentiating between stress reactions and nerve agent-induced organic brain syndromes has been pointed out. Knowledge of the behavioral effects of CWAs and their medical countermeasures is imperative to ensure that military and civilian medical and mental health organizations can deal with possible incidents involving weapons of mass destruction.

For the reader's benefit, the CWAs, their two-letter North Atlantic Treaty Organization (NATO) codes (which were established by a NATO international convention and convey no clinical implications), their unique physical features, and their initial effects are listed in **Table 262e-1**. **Table 262e-2** provides guidelines for immediate treatment. The focus of this chapter is on the blister and nerve CWAs, which have been employed in battle and against civilians and have had a significant public health impact.

VESICANTS: SULFUR MUSTARD

Sulfur mustard has been a military threat since it first appeared on the battlefield in Belgium during WWI. In modern times, it remains a threat on the battlefield as well as a potential chemical terrorism

TABLE 262e-1 RECOGNIZING AND DIAGNOSING HEALTH EFFECTS OF CHEMICAL TERRORISM

Agent Category	Agent Name	Unique Characteristics	Initial Effects
Nerve	Cyclohexyl sarin (GF) ^a	Miosis (pinpoint pupils)	Miosis (pinpoint pupils)
	Sarin (GB)	Copious secretions	Blurred/dim vision
	Soman (GD)	Muscle twitching/fasciculations	Headache
	Tabun (GA)		Nausea, vomiting, diarrhea
	VX		Copious secretions/sweating
	VR		Muscle twitching/fasciculations
Asphyxiant/ blood	Arsine (SA)	Possible cherry-red skin	Confusion
	Cyanogen chloride (CK)	Possible cyanosis	Nausea
	Hydrogen cyanide (AC)	Possible frostbite ^b	Gasping for air in some cases; similar to asphyxiation but more abrupt onset
Choking/ pulmonary- damaging	Chlorine (CL)	Chlorine is a greenish-yellow gas with a pungent odor.	Eye and skin irritation
	Hydrogen chloride		Airway irritation
	Nitrogen oxides		Dyspnea, cough
	Phosgene (CG)	Phosgene gas smells like newly mown hay or grass. Possible frostbite ^b	Sore throat Chest tightness
Blistering/ vesicant	Mustard (HD)/sulfur mustard (H)	Mustard has an odor like burning garlic or horseradish.	Severe irritation
	Mustard gas (H)		Redness and blisters of the skin
	Nitrogen mustard (HN-1, HN-2, HN-3)	Lewisite has an odor like penetrating geranium.	Tearing, conjunctivitis, corneal damage
	Lewisite (L)	Phosgene oxime has a pepperish or pungent odor.	Mild respiratory distress to marked airway damage
	Phosgene oxime (CX)		May cause death
Incapacitating/ behavior- altering	3-Quinuclidinyl benzilate, or Agent 15 (BZ)	May appear as mass drug intoxication with erratic behaviors, shared realistic and distinct hallucinations, disrobing, and confusion	Dry mouth and skin
		Hyperthermia	Initial tachycardia
		Mydriasis (dilated pupils)	Altered consciousness, delusions, denial of illness, belligerence
			Hyperthermia Ataxia (lack of coordination) Hallucinations Mydriasis (dilated pupils)

^aLetters in parentheses indicate NATO codes for designated agents. ^bFrostbite may occur from skin contact with liquid arsine, cyanogen chloride, or phosgene.

Source: State of New York, Department of Health, as modified by the Chemical Casualty Care Division, U.S. Army Medical Research Institute of Chemical Defense.

weapon because of the simplicity of its manufacture and its extreme effectiveness. Sulfur mustard accounted for 70% of the 1.3 million chemical casualties in WWI. Occasional cases of sulfur mustard intoxication continue to occur in the United States among people exposed to WWI- and WWII-era munitions.

Mechanism Sulfur mustard constitutes both a vapor and a liquid threat to all exposed epithelial surfaces. The effects are delayed,

TABLE 262e-2 DECONTAMINATION AND TREATMENT OF CHEMICAL TERRORISM

Agent Category	Decontamination	First Aid	Other Considerations
Nerve	Remove clothing immediately. Gently wash skin with soap and water. Do not abrade skin. For eyes, flush with plenty of water or normal saline.	Atropine before other measures Pralidoxime (2-PAM) chloride	Onset of symptoms from dermal contact with liquid forms may be delayed. Repeated antidote administration may be necessary.
Asphyxiant/blood	Remove clothing immediately if no frostbite. ^a Gently wash skin with soap and water. Do not abrade skin. For eyes, flush with plenty of water or normal saline.	Rapid treatment with oxygen For cyanide, use antidotes (sodium nitrite and then sodium thiosulfate).	Arsine and cyanogen chloride may cause delayed pulmonary edema.
Choking/pulmonary-damaging	Remove clothing immediately if no frostbite. ^a Gently wash skin with soap and water. Do not abrade skin. For eyes, flush with plenty of water or normal saline.	Fresh air, forced rest Semiupright position If signs of respiratory distress are present, oxygen with or without positive airway pressure may be needed. Other supportive therapy, as needed	May cause delayed pulmonary edema, even after a symptom-free period that varies in duration with the amount inhaled
Blistering/vesicant	Immediate decontamination is essential to minimize damage. Remove clothing immediately. Gently wash skin with soap and water. Do not abrade skin. For eyes, flush with plenty of water or normal saline.	Immediately decontaminate skin. Flush eyes with water or normal saline for 10–15 min. If breathing is difficult, give oxygen. Supportive care	Mustard has an asymptomatic latent period. There is no antidote or treatment for mustard. Lewisite causes immediate burning pain, with blisters developing later. Specific antidote—British anti-lewisite—may decrease systemic effects of lewisite. Phosgene oxime causes immediate pain. Possible pulmonary edema
Incapacitating/behavior-altering	Remove clothing immediately. Gently wash skin with water or soap and water. Do not abrade skin.	Remove heavy clothing. Evaluate mental status. Use restraints as needed. Monitor core temperature carefully. Supportive care	Hyperthermia and self-injury are the greatest risks. Hard to detect because it is an odorless and nonirritating substance Possible serious arrhythmias Specific antidote (physostigmine) may be available.

^aFor frostbite areas, DO NOT remove any adhering clothing. Wash area with plenty of warm water to release clothing.

Source: State of New York, Department of Health.

appearing hours after exposure. The organs most commonly affected are the skin (with erythema and vesicles), the eyes (with manifestations ranging from mild conjunctivitis to severe eye damage), and the airways (with effects ranging from mild upper airway irritation to severe bronchiolar damage). After exposure to large quantities of mustard, precursor cells of the bone marrow are damaged, with consequent pancytopenia and secondary infection. The gastrointestinal mucosa may be damaged, and there are sometimes central nervous system (CNS) signs of unknown mechanism. No specific antidotes exist; management is entirely supportive. Immediate decontamination of the liquid is the only way to reduce damage. Complete decontamination in 2 min stops clinical injury; decontamination at 5 min reduces skin injury by ~50%. Table 262e-2 lists approaches to decontamination after exposure to mustard and other CWAs.

Mustard dissolves slowly in aqueous media such as sweat, but, once dissolved, it rapidly forms cyclic ethylene sulfonium ions that are extremely reactive with cell proteins, cell membranes, and especially DNA in rapidly dividing cells. The ability of mustard to react with and alkylate DNA gives rise to the effects by which it has been characterized as “radiomimeti”—i.e., similar to radiation injury. Mustard has many biologic actions, but its actual mechanism of action is largely unknown. Much of the biologic damage from mustard results from DNA alkylation and cross-linking in rapidly dividing cells: corneal epithelium, basal keratinocytes, bronchial mucosal epithelium, gastrointestinal mucosal epithelium, and bone marrow precursor cells. This damage may lead to cellular death and inflammatory reactions. In the skin, proteolytic digestion of anchoring filaments at the epidermal-dermal junction may be the major mechanism of action resulting in

blister formation. Mustard also has mild cholinergic activity, which may be responsible for effects such as early gastrointestinal and CNS symptoms. Mustard reacts with tissue within minutes of entering the body. Its circulating half-life in unaltered form is extremely brief.

Clinical Features Topical effects of mustard occur in the skin, airways, and eyes; the eyes are most sensitive and the airways next most sensitive. Absorbed mustard may produce effects in the bone marrow, gastrointestinal tract, and CNS. Direct injury to the gastrointestinal tract also may occur after ingestion of the compound through contamination of water or food.

Erythema is the mildest and earliest form of mustard skin injury. It resembles sunburn and is associated with pruritus, burning, or stinging pain. Erythema begins to appear within 2 h to 2 days after vapor exposure. Time of onset depends on severity of exposure, ambient temperature and humidity, and type of skin. The most sensitive sites are warm moist locations and areas of thin delicate skin, such as the perineum, external genitalia, axillae, antecubital fossae, and neck.

Within the erythematous areas, small vesicles can develop, which may later coalesce to form bullae (Fig. 262e-1). The typical bulla is large, dome-shaped, flaccid, thin-walled, translucent, and surrounded by erythema. The blister fluid, a transudate, is clear to straw-colored and becomes yellow, tending to coagulate. The fluid does not contain mustard and is not itself a vesicant. Lesions from high-dose liquid exposure may develop a central zone of coagulation necrosis with blister formation at the periphery. These lesions take longer to heal and are more prone to secondary infection than are the uncomplicated lesions seen at lower exposure levels. Severe lesions may require skin grafting.



FIGURE 262e-1 Large bulla formation from mustard burn in a patient. Although the blisters in this case involved only 7% of the body surface area, the patient still required hospitalization in a burn intensive care unit.

The primary airway lesion is necrosis of the mucosa with possible damage to underlying smooth muscle. The damage begins in the upper airways and descends to the lower airways in a dose-dependent manner. Usually the terminal airways and alveoli are affected only as a terminal event. Pulmonary edema is not usually present unless the damage is very severe, and then it becomes hemorrhagic.

The earliest effects of mustard—and perhaps the only effects of a low concentration—involve the nose, sinuses, and pharynx. There may be irritation or burning of the nares, epistaxis, sinus pain, and pharyngeal pain. As the concentration increases, laryngitis, voice changes, and nonproductive cough develop. Damage to the trachea and upper bronchi leads to a productive cough. Lower airway involvement causes dyspnea, severe cough, and increasing quantities of sputum. Terminally, there may be necrosis of the smaller airways with hemorrhagic edema into surrounding alveoli. Hemorrhagic pulmonary edema is rare.

Necrosis of airway mucosa causes “pseudomembrane” formation. These membranes may obstruct the bronchi. During WWI, high-dose mustard exposure caused acute death via this mechanism in a small minority of cases (Fig. 262e-2).

The eyes are the organs most sensitive to mustard vapor injury. The latent period is shorter for eye injury than for skin injury and is also dependent on exposure concentration. After low-dose vapor exposure, irritation evidenced by reddening of the eyes may be the only effect. As the dose increases, the injury includes progressively severe conjunctivitis, photophobia, blepharospasm, pain, and corneal damage



FIGURE 262e-2 Schematic diagram of pseudomembrane formation as is seen in high-dose sulfur mustard vapor inhalation exposure. In World War I, severe inhalation exposure often caused death via obstruction of large airways.



FIGURE 262e-3 World War I photograph of troops exposed to sulfur mustard vapor. The vast majority of these troops survived with no long-term damage to the eyes; however, they were rendered effectively blind for days or weeks.

(Fig. 262e-3). About 90% of eye injuries related to mustard heal in 2 weeks to 2 months without sequelae. Scarring between the iris and the lens may follow severe effects; this scarring may restrict pupillary movements and predispose victims to glaucoma. The most severe damage is caused by liquid mustard. Extensive eye exposure can be followed by severe corneal damage with possible perforation of the cornea and loss of the eye. In some individuals, latent chronic keratitis, sometimes associated with corneal ulcerations, has been described as early as 8 months and as late as 20 years after initial exposure.

The mucosa of the gastrointestinal tract is susceptible to mustard damage from either systemic absorption or ingestion of the agent. Mustard exposure in small amounts will cause nausea and vomiting lasting up to 24 h. The mechanism of the nausea and vomiting is not understood, but mustard does have a cholinergic-like effect. The CNS effects of mustard also remain poorly defined. Exposure to large amounts can cause seizures in animals. Reports from WWI and from the Iran–Iraq war described people exposed to small amounts of mustard acting sluggish, apathetic, and lethargic. These reports suggest that minor psychological problems could linger for ≥ 1 year.

The cause of death in the majority of mustard poisoning cases is sepsis and respiratory failure. Mechanical obstruction via pseudomembrane formation and agent-induced laryngospasm is important in the first 24 h, but only in cases of severe exposure. From the third through the fifth day after exposure, secondary pneumonia due to bacterial invasion of denuded necrotic mucosa can be expected. The third wave of death is caused by agent-induced bone marrow suppression, which peaks 7–21 days after exposure and causes death via sepsis.

TREATMENT SULFUR MUSTARD INTOXICATION

A patient severely ill from mustard poisoning requires the general supportive care provided for any severely ill patient as well as the specific care given to a burn patient. Liberal use of systemic analgesics, maintenance of fluid and electrolyte balance, provision of nutrition, administration of appropriate antibiotics, and other supportive measures are necessary (Table 262e-2).

The management of a patient exposed to mustard may range from simple (as in the provision of symptom-based care for a sunburn-like erythema) to complex (as in total management of a severely ill patient with burns, immunosuppression, and multi-system involvement). Before raw denuded areas of skin develop, especially with less severe exposures, topical cortisone creams or lotions may be of benefit. Some very basic research data point to the early use of anti-inflammatory preparations. Small blisters (<1–2 cm) should be left intact. Because larger bullae eventually will break, they should be unroofed carefully. Denuded areas should be irrigated three or four times daily with saline, other sterile

solutions, or soapy water and then liberally covered with the topical antibiotic of choice, such as silver sulfadiazine, mafenide acetate, or triple antibiotic ointment to a thickness of 1–2 mm. Some physicians advocate sterile needle drainage of large blisters, with collapsing of the blister roof to form a sterile dressing. Mustard blister fluid does not contain sulfur mustard but rather consists only of sterile tissue fluid. Health care staff should not fear possible contamination. If an antibiotic cream is not available, sterile petrolatum will be useful. Modified Dakin's solution (sodium hypochlorite, 0.5%) was used for field-expedient irrigation and antisepsis both in WWI and for Iranian casualties in the Iran–Iraq war (1984–1987). Large areas of vesication require hospitalization, IV therapy, and whirlpool bath irrigation.

Systemic analgesics should be used liberally, particularly before manipulation of the patient. Monitoring of fluids and electrolytes is important in any sick patient, but it must be recognized that fluid loss is not of the magnitude seen with thermal burns. Overly rigorous hydration seems to have precipitated pulmonary edema in a few Iranian casualties sent to European hospitals.

Conjunctival irritation from a low-vapor exposure responds to any of a number of available ophthalmic solutions after the eyes are irrigated thoroughly. A topical antibiotic applied several times a day reduces the incidence and severity of infection. Animal laboratory data reflect remarkable results with early application of commercially available topical antibiotic/glucocorticoid ophthalmologic ointments. An ophthalmologist should be consulted. Topical glucocorticoids are not of proven value, but their use during the first few hours or days may significantly reduce inflammation and subsequent damage. Further use should be relegated to an ophthalmologist.

Petroleum jelly or a similar substance should be applied regularly to the edges of the eyelids to prevent them from sticking together. Topical analgesics, although of limited value, may be useful initially if blepharospasm is too severe to permit an adequate examination.

A productive cough and dyspnea accompanied by fever and leukocytosis occurring within 12–24 h are indicative of chemical pneumonitis. The clinician must resist the urge to use prophylactic antibiotics for this process. Infection often occurs on the third to fifth day and is signaled by increased fever, a pulmonary infiltrate, and increased sputum production with a change in color. Appropriate antibiotic therapy should await confirmation by Gram's stain and, later, by culture and sensitivity assessment.

Intubation may be necessary if laryngeal spasm or edema makes breathing difficult or becomes life-threatening. Intubation permits better ventilation and facilitates suctioning of necrotic and inflammatory debris. Early use of positive end-expiratory pressure (PEEP) or continuous positive airway pressure (CPAP) may be beneficial. Pseudomembrane formation may require fiberoptic bronchoscopy for suctioning of necrotic debris.

Bronchodilators are of benefit for bronchospasm. If additional relief of bronchospasm is needed, glucocorticoids should be used. There is little evidence that the routine use of glucocorticoids is beneficial except for additional relief of bronchospasm.

Leukopenia begins around day 3 with major systemic absorption. Marrow suppression peaks at 7–14 days. In the Iran–Iraq war, a white blood cell count of $\leq 200/\mu\text{L}$ usually resulted in death of the patient. Sterilization of the gut by nonabsorbable antibiotics should be considered to reduce the possibility of sepsis from enteric organisms. Cellular replacement (bone marrow transplants or transfusions) may be successful. Granulocyte colony-stimulating factor produced a 50% reduction in the time required for bone marrow recovery in nonhuman primates exposed to sulfur mustard. Medication for nausea and vomiting may be necessary for gastrointestinal side effects. Lymphopenia precedes general leukopenia by a day or more and may be a useful clinical tip-off to impending leukopenia.

Excellent experimental assessments of the contributions of DNA alkylation, inflammation, activation of proteolytic enzymes, or lipid peroxidation to the mustard injury have been developed in the past 15–20 years. Some examples include (1) the demonstration of a reduction by up to 75% of inflammation and tissue damage in the

mouse ear swelling test by vanilloid compounds and (2) the demonstration of 50–60% protection by *N*-acetylcysteine in the generation of free radicals within guinea pig lung exposed to mustard. In many cases, the demonstration of protection is dependent on the availability of sufficient amounts of drugs with adequate half-lives. Strategies to enhance bioavailability include attachment of polyethylene glycol to the antioxidant drug/enzyme and/or delivery of the drug/enzyme in a liposome.

NERVE AGENTS

The organophosphorus nerve agents are the deadliest of the CWAs. They work by inhibition of tissue synaptic acetylcholinesterase, creating an acute cholinergic crisis. Death ensues because of respiratory depression and can occur within seconds to minutes.

The nerve agents tabun and sarin were first used on the battlefield by Iraq against Iran during the first Persian Gulf War (1984–1987). Estimates of casualties from these agents range from 20,000–100,000. In 1994 and 1995, the Japanese cult Aum Shinrikyo used sarin in two terrorist attacks in Matsumoto and Tokyo. Two U.S. soldiers were exposed to sarin while rendering safe an improvised explosive device in Iraq in 2004.

The “classic” nerve agents include tabun (GA), sarin (GB), soman (GD), cyclosarin (GF), and VX; VR, similar to VX, was manufactured in the former Soviet Union (Table 262e-1). All the nerve agents are organophosphorus compounds, which are liquid at standard temperature and pressure. The “G” agents evaporate at about the rate of water, except for cyclosarin, which is oily and thus probably will have evaporated within 24 h after deposition on the ground. Their high volatility thus makes a spill of any amount a serious vapor hazard. In the Tokyo subway attack in which sarin was used, 100% of the symptomatic patients inhaled sarin vapor that spilled out on the floor of the subway cars. The low vapor pressure of VX, an oily liquid, makes it much less of a vapor hazard but potentially a greater environmental hazard because it persists in the environment far longer.

Mechanism Acetylcholinesterase inhibition accounts for the major life-threatening effects of nerve agent poisoning. The efficacy of antidotal therapy in the reversal of this inhibition proves that this is the primary toxic action of these poisons. At cholinergic synapses, acetylcholinesterase, bound to the postsynaptic membrane, functions as a turn-off switch to regulate cholinergic transmission. Inhibition of acetylcholinesterases causes the released neurotransmitter, acetylcholine, to accumulate abnormally. End-organ overstimulation, which is recognized by clinicians as a cholinergic crisis, ensues (Fig. 262e-4).

Clinical Features Clinical effects of nerve agent exposure are identical for vapor and liquid exposure routes if the dose is sufficiently large. The speed and order of symptom onset will differ (Table 262e-2).

Exposure of a patient to nerve agent vapor, by far the more likely route of exposure in both battlefield and terrorist scenarios, will cause cholinergic symptoms in the order in which the toxin encounters cholinergic synapses. The most exposed synapses on the human integument are in the pupillary muscles. Nerve agent vapor easily crosses the cornea, interacts with these synapses, and produces miosis, described by Tokyo subway victims as “the world going black.” Rarely,

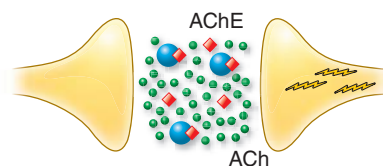


FIGURE 262e-4 Schematic diagram of the pathophysiology of nerve agent exposure. Nerve agent (♦) binds to the active site of acetylcholinesterase (AChE), which is shown as floating free in space but is in reality a postsynaptic membrane-bound enzyme. As a result, acetylcholine (●), which normally is released from presynaptic membrane but then is degraded, accumulates, and this leads (⚡) to organ overstimulation and cholinergic crisis.

this vapor also can cause eye pain and nausea. Exocrine glands in the nose, mouth, and pharynx are next exposed to the vapor, and cholinergic overload here causes increased secretions, rhinorrhea, excess salivation, and drooling. Toxin then interacts with exocrine glands in the upper airway, causing bronchorrhea, and with bronchial smooth muscle, causing bronchospasm. This combination of events can result in hypoxia.

Once the victim has inhaled, vapor can passively cross the alveolar-capillary membrane, enter the bloodstream, and incidentally and asymptotically inhibit circulating cholinesterases, particularly free butyrylcholinesterase and erythrocyte acetylcholinesterase, both of which can be assayed. Unfortunately, the results of this assay may not be easily interpretable without a baseline, since cholinesterase levels vary enormously between individuals and over time in an individual, healthy patient.

Usually the first organ system to become symptomatic from bloodborne nerve agent exposure is the gastrointestinal tract, where cholinergic overload causes abdominal cramping and pain, nausea, vomiting, and diarrhea. After the gastrointestinal tract becomes involved, nerve agents will affect the heart, distant exocrine glands, muscles, and brain. Because there are cholinergic synapses on both the vagal (parasympathetic) and sympathetic sides of the autonomic input to the heart, one cannot predict how heart rate and blood pressure will change once intoxication has occurred. Remote exocrine activity will include oversecretion in the salivary, nasal, respiratory, and sweat glands—the patient will be “wet all over.” Bloodborne nerve agents will overstimulate neuromuscular junctions in skeletal muscles, causing fasciculations followed by frank twitching. If the process goes on long enough, ATP in muscles will eventually be depleted and flaccid paralysis will ensue.

In the brain, since the cholinergic system is so widely distributed, bloodborne nerve agents will, in sufficient doses, cause rapid loss of consciousness, seizures, and central apnea leading to death within minutes. If respiration is supported, status epilepticus that does not respond to usual anticonvulsants may ensue (Chap. 445). If status epilepticus persists, neuronal death and permanent brain dysfunction may occur. Even in mild nerve-agent intoxication, patients may recover but may experience weeks of irritability, sleep disturbance, and nonspecific neurobehavioral manifestations.

The time from exposure to development of the full-blown cholinergic crisis after nerve agent vapor inhalation can be minutes or even seconds, yet there is no depot effect. Since nerve agents have a short

circulating half-life, if the patient is supported and, ideally, treated with antidotes, improvement should be rapid, without subsequent deterioration.

Liquid exposure to nerve agents results in different speeds and orders of symptom onset. A nerve agent on intact skin will partially evaporate and partially begin to travel through the skin, causing localized sweating and then, when it encounters neuromuscular junctions, localized fasciculations. Once in muscle, the agent will cross into the circulation and cause gastrointestinal discomfort, respiratory distress, heart rate changes, generalized fasciculations and twitching, loss of consciousness, seizures, and central apnea. The time course will be much longer than with vapor inhalation; even a large, lethal droplet can take up to 30 min to exert an effect, and a small, sublethal dose could progressively take effect over 18 h. Clinical worsening that occurs hours after treatment has started is far more likely with liquid than with vapor exposure. In addition, miosis, which is practically unavoidable with vapor exposure, is not always present with liquid exposure and may be the last manifestation to develop in this situation; such a delay is due to the relative insulation of the pupillary muscle from the systemic circulation.

Unless the cholinesterase is reactivated by specific therapy (oximes), its binding to the enzyme is essentially irreversible. Erythrocyte acetylcholinesterase activity recovers at ~1% per day. Plasma butyrylcholinesterase recovers more quickly and is a better guide to recovery of tissue enzyme activity.

TREATMENT NERVE AGENT INTOXICATION

Acute nerve agent poisoning is treated by decontamination, respiratory support, and three antidotes: an anticholinergic, an oxime, and an anticonvulsant (Table 262e-3). In acute cases, all these forms of therapy may be given simultaneously.

DECONTAMINATION

Decontamination of a vapor is formally unnecessary; however, in the Tokyo subway attack, sarin vapor trapped in patients' clothing caused miosis in 10% of emergency personnel. Removal of clothing would have prevented most of these instances. Expedient decontamination methods for CWAs are available. For soap and water decontamination, the skin surface and hair are washed in warm or tepid water at least three times, or the exposed individual showers for 2 min, washing with soap and rinsing. The rapid physical removal

TABLE 262e-3 ANTIDOTE RECOMMENDATIONS AFTER EXPOSURE TO NERVE AGENTS

Patient's Age	Antidotes		Other Treatment
	Mild/Moderate Effects ^a	Severe Effects ^b	
Infants (0–2 years)	Atropine (0.05 mg/kg IM or 0.02 mg/kg IV) and 2-PAM Cl (15 mg/kg IM or IV slowly)	Atropine (0.1 mg/kg IM or 0.02 mg/kg IV) and 2-PAM chloride (25 mg/kg IM or 15 mg/kg IV slowly)	Assisted ventilation after antidotes for severe exposure
Child (2–10 years)	Atropine (1 mg IM or 0.02 mg/kg IV) and 2-PAM Cl ^c (15 mg/kg IM or IV slowly)	Atropine (2 mg IM or 0.02 mg/kg IV) and 2-PAM chloride ^c (25 mg/kg IM or 15 mg/kg IV slowly)	Repeat atropine (2 mg IM, or 1 mg IM for infants) at 5- to 10-min intervals until secretions have diminished and breathing is comfortable or airway resistance has returned to nearly normal.
Adolescent (>10 years)	Atropine (2 mg IM or 0.02 mg/kg IV) and 2-PAM Cl ^c (15 mg/kg IM or IV slowly)	Atropine (4 mg IM or 0.02 mg/kg IV) and 2-PAM Cl ^c (25 mg/kg IM or 15 mg/kg IV slowly)	
Adult	Atropine (2–4 mg IM or IV) and 2-PAM Cl (600 mg IM or 15 mg/kg IV slowly)	Atropine (6 mg IM) and 2-PAM Cl (1800 mg IM or 15 mg/kg IV slowly)	Phentolamine for 2-PAM-induced hypertension (5 mg IV for adults; 1 mg IV for children) Diazepam for convulsions (0.2–0.5 mg IV for infants <5 years; 1 mg IV for children >5 years; 5 mg IV for adults).
Elderly, frail	Atropine (1 mg IM) and 2-PAM Cl (10 mg/kg IM or 5–10 mg/kg IV slowly)	Atropine (2–4 mg IM) and 2-PAM Cl (25 mg/kg IM or 5–10 mg/kg IV slowly)	

^aMild/moderate effects include localized sweating, muscle fasciculations, nausea, vomiting, weakness, and dyspnea. ^bSevere effects include unconsciousness, convulsions, apnea, and flaccid paralysis. ^cIf the calculated dose exceeds the adult IM dose, adjust accordingly.

Abbreviation: 2-PAM Cl, 2-pralidoxime (or Protopam[®]) chloride.

Source: State of New York, Department of Health.

of a chemical agent is essential. Scrubbing of exposed skin with a stiff brush or bristles is discouraged, because skin damage may occur and may increase absorption of agent. “Gentle” liquid dish soap and copious amounts of water should be used, with mild to moderate friction applied with a single-use sponge or washcloth in the first and second washes. The third wash should be a rinse with copious amounts of warm or tepid water. Shampoo can be used to wash the hair. If only cold water is available, it should be used; decontamination should not be delayed while warm water is sought. Spot (local) decontamination with reactive skin decontamination lotion (RSDL), followed by a soap and water wash/shower, is the method preferred by the Department of Defense. RSDL is available for purchase by civilians and has been shown to be superior across a broad spectrum of nerve agents as well as sulfur mustard. RSDL is the only product approved by the U.S. Food and Drug Administration (FDA) for initial spot decontamination. An important caveat is that RSDL and 0.5% sodium hypochlorite (dilute bleach military field expedient) should *not* be used concurrently because of a potential exothermic reaction. In any event, decontamination must be accomplished before the patient enters the medical facility to avoid contaminating the facility and its staff. In patients with contaminated wounds, potentially contaminated clothing and other foreign material that may serve as a depot for the liquid agent should be extracted.

RESPIRATORY SUPPORT

Death from nerve agent poisoning is almost always attributable to respiratory causes. Ventilation will be complicated by increased resistance and secretions. Atropine should be given before ventilation or as it begins, since it will make ventilation far easier.

ANTIDOTAL THERAPY

Atropine In theory, any anticholinergic agent could be used to treat nerve agent poisoning. Worldwide, however, the choice is invariably atropine because of its wide temperature stability and rapid effectiveness when administered either IM or IV and because inadvertent administration of this drug usually causes little CNS dysfunction (Table 262e-3). Atropine rapidly reverses cholinergic overload at muscarinic synapses but has little effect at nicotinic synapses. The practical implication is that atropine can quickly treat the life-threatening respiratory effects of nerve agents but probably will not help with neuromuscular (and possibly sympathetic) effects. In the field, military personnel are given a combined autoinjector containing both 2.1 mg of atropine and oxime (2-pralidoxime chloride [2-PAM Cl])—a product licensed by the FDA under the trade name Duodote[®]. Its military designation is the Antidote Treatment Nerve Agent Autoinjector (ATNAA) (Fig. 262e-5). Only full—and not divided—autoinjector doses can be administered. The field loading dose is 2, 4, or 6 mg, with re-treatment every 5–10 min until the patient’s breathing improves and secretions diminish. The Iranian military initially used larger doses during the Iran–Iraq war, in which oximes were in short supply. When the patient reaches a level of medical care at which drugs can be given IV, this is the preferred route. In small children, the IV route may be the initial avenue for atropine therapy; however, pediatric autoinjectors of 0.5 mg and 1 mg are manufactured. There is no upper limit to atropine therapy (whether IM or IV), but the total average dose for a severely afflicted adult is usually 20–30 mg.

In a mildly afflicted patient with miosis and no other systemic symptoms, atropine or homatropine eyedrops may suffice for therapy.

This treatment will result in ~24 h of mydriasis. Frank miosis or imperfect accommodation may persist for weeks or even months after all other signs and symptoms have resolved.

Oximes Oximes are nucleophiles that reactivate the cholinesterase whose active site has been occupied and bound to nerve agent (Table 262e-3). Therapy with oximes therefore restores normal enzyme function. Oxime therapy is limited by a second side reaction, called “aging,” in which a side chain on nerve agents falls off the complex at a characteristic rate. “Aged” complexes are negatively charged, and oximes cannot reactivate negatively charged complexes. The practical effect of this limitation differs from one nerve agent to another since each ages at a characteristic rate. For example, sarin ages in 3–5 h, tabun ages over a longer period (12–13 h), and VX ages much less rapidly (>48 h). All these intervals are so much longer than the patient’s expected life span and expected treatment time after acute nerve-agent toxicity that they are irrelevant. Soman, in contrast, ages in 2 min; thus, only a few minutes after exposure, oximes are useless in treating soman poisoning. The oxime used varies by country; the United States has approved and fielded 2-PAM Cl. MARK 1 kits and Duodotes[®] (Fig. 262e-5A) both contain autoinjectors holding 600 mg of 2-PAM Cl. Initial field loading doses are 600, 1200, and 1800 mg. Since blood pressure may become elevated after administration of 45 mg/kg in adults, field use of 2-PAM Cl is restricted to 1800 mg/h IM. During the time when more oxime cannot be given, atropine alone is recommended. In the hospital setting, 2.5–25 mg/kg of 2-PAM Cl by the IV route has been found to reactivate 50% of inhibited cholinesterase. The usual recommendation is 1000 mg by slow IV drip over 20–30 min, with ≤2500 mg over a period of 1–1.5 h. Active research aims to field a more effective and broader-spectrum oxime than 2-PAM Cl.

Anticonvulsants Nerve agent–induced seizures do not respond to the usual anticonvulsants used for status epilepticus, including phenytoin, phenobarbital, carbamazepine, valproic acid, and lamotrigine (Chap. 445). The only anticonvulsants that have been shown to stop this form of status are the benzodiazepines. Diazepam is the only benzodiazepine approved for seizures in humans, although other FDA-approved benzodiazepines (notably midazolam) work well against nerve agent–induced seizures in animal models. Diazepam



FIGURE 262e-5 Antidotes to nerve agents. **A.** The Antidote Treatment Nerve Agent Autoinjector (ATNAA) replaces the MARK I Kit. It is easier to self-administer and allows prompt distribution of the antidotes atropine and 2-pralidoxime chloride (2-PAM Cl).

B. Diazepam 10-mg autoinjectors are carried by all U.S. military forces in a potential chemical battlefield and are being stockpiled by civilian first responders.

therefore is manufactured in 10-mg injectors for IM use and given to U.S. forces for this purpose (Fig. 262e-5B). Civilian agencies are stockpiling this field product (convulsive antidote for nerve agent, CANA), which generally has not been used in hospital practice. Extrapolation from animal studies indicates that adults will probably require 30–40 mg of diazepam given IM to stop nerve agent–induced status epilepticus. In the hospital or in a small child unable to receive the autoinjector, IV diazepam may be used at similar doses. The clinician may confuse seizures with the neuromuscular signs of nerve agent poisoning. In the hospital, early electroencephalography is advised to distinguish among nonconvulsive status epilepticus, actual seizures, and postictal paralysis. Animal studies have shown that the most effective benzodiazepine in this situation is midazolam, which is not FDA-approved for seizures. At the time of this writing, a new drug application for use of midazolam against seizures has been submitted to the FDA. The superiority of IM midazolam to IV lorazepam in a large community trial of status epilepticus suggests that emergency personnel will soon incorporate autoinjectors into routine clinical practice and that these field products will thus become integrated into clinical medicine.

Peripheral neuropathy and the so-called intermediate syndrome, which are prominent long-term effects of insecticide poisoning, are not described in nerve agent survivors.

Recent research has explored approaches leading to transient “immunity” and eventually to biologic products that will be protective against lethal nerve agents yet be devoid of side effects. A novel approach is to use enzymes to scavenge these highly toxic nerve agents before they attack their intended targets. The accumulated work has shown that if a scavenger is present at the time of nerve agent exposure, toxicant levels are rapidly reduced. This reduction is so rapid and profound that the need to administer a host of pharmacologically active drugs as antidotes is, according to laboratory studies, eliminated.

CYANIDE

Cyanide (CN^-) has become an agent of particular interest in terrorist scenarios because of its applicability to indoor targets. In recent years, for example, attacks with this agent have targeted the water supply of the U.S. Embassy in Italy. The 1993 World Trade Center bombing in New York may have been intended as a cyanide release as well.

Hydrogen cyanide and cyanogen chloride, the major forms of cyanide, are either true gases or liquids very close to their boiling points at standard room temperature. Hydrogen cyanide gas is lighter than air and does not remain concentrated outdoors for long; thus it is a poor military weapon but an effective weapon in an indoor space such as a train station or a sports arena. Cyanide is also water-soluble and poses a threat to the food and water supply from either accidents or malign intent. It is well absorbed from the gastrointestinal tract, through the skin, or via inhalation—the preferred terrorist route. Cyanide smells like bitter almonds, but 50% of persons lack the ability to smell it.

Unique among CWAs, cyanide is a normal constituent of the environment and actually is a required cofactor in many compounds important in metabolism, including vitamin B_{12} . Cyanide is present in many plants, including tobacco; therefore, smokers, for instance, chronically carry cyanide at three times the usual level. Humans have evolved a detoxification mechanism for cyanide. Cyanide poisoning results if a large challenge of CN^- overwhelms this mechanism, while treatment of cyanide poisoning exploits it.

Mechanism Cyanide directly poisons the last step in the mitochondrial electron transport chain, cytochrome a3, which results in a shutdown of cellular energy production. Tissues are poisoned in direct proportion to their metabolic rate, with the carotid baroreceptors and the brain—the most metabolically active tissues in the body—affected fastest and most severely. This poisoning results from cyanide’s high affinity for certain metals, notably Co and Fe^{+++} . Cytochrome a3 contains Fe^{+++} , to which CN^- binds. Cyanide-poisoned tissues cannot extract oxygen from the blood; even though pulmonary oxygen

exchange and cardiac function are preserved, cells die of hypoxia—i.e., of histotoxic rather than cardiopulmonary cause.

Clinical Features Hyperpnea occurs ~15 s after inhalation of a high concentration of cyanide and is followed within 15–30 s by the onset of convulsions and electrical status epilepticus. Respiratory activity stops 2–3 min later, and cardiac activity ceases several minutes after that. Exposure, especially via inhalation, to a large challenge of CN^- can cause death in as little as 8 min. Smaller challenges will cause symptom spread over a longer period; very low doses may produce no effects at all because of the body’s ability to detoxify small amounts. Cyanogen chloride additionally produces mucous membrane irritation. Many but not all patients have a cherry-red appearance because their venous blood remains oxygenated.

Differential Diagnosis In a mass casualty incident caused by a chemical agent, the primary differential diagnosis of cyanide poisoning will be nerve agent poisoning. Cyanide-poisoned patients lack the prominent cholinergic signs seen in nerve agent poisoning, such as miosis and increased secretions. The cherry-red appearance often seen in cyanide poisoning is never seen in nerve agent poisoning. Cyanosis, confusingly, is *not* a prominent early sign in cyanide poisoning.

TREATMENT CYANIDE

Treatment of cyanide poisoning may require simply evacuation of the patient from the source of contamination. Decontamination of a true gas, other than clothing removal to avoid gas trapped in clothing air cells, is probably not a major concern.

Oxygen, supplied via mask, nasal cannula, or endotracheal tube, has been shown to benefit patients, although the benefit is not explained by the known mechanism of action of CN^- .

Cyanide antidotes exploit the body’s innate detoxification mechanism, the hepatic enzyme rhodanese. They also exploit cyanide’s affinity for certain metal ions. Antidote recommendations are summarized in [Table 262e-4](#).

The classic two-step cyanide antidote kit includes two IV solutions: sodium nitrite and sodium thiosulfate. It may also include amyl nitrite perles for inhalation. Nitrites are methemoglobin formers; when administered to a patient, nitrite converts a fraction of the body’s hemoglobin into methemoglobin by converting heme iron from Fe^{++} to Fe^{+++} . CN^- has a greater affinity for methemoglobin Fe^{+++} than for cytochrome a3. As a result, administration of nitrite creates a “sink” of cyanmethemoglobin; formation of methemoglobin pulls CN^- off mitochondrial cytochrome a3, allowing cellular respiration to resume. Recent work suggests that nitrites may also work via a second mechanism involving the neurotransmitter nitrous oxide; if so, this mechanism may explain why cyanide-poisoned patients improve after nitrite administration faster than is explained by the known rate of methemoglobin formation.

Nitrite administration may save the patient acutely but creates an unstable pool of cyanmethemoglobin, the elimination of which requires a sulfur donor: sodium thiosulfate. Sodium thiosulfate donates sulfur to the reaction catalyzed by rhodanese; cyanide is converted to thiocyanate, a compound the body eliminates harmlessly in urine. Sodium thiosulfate alone may be administered to fire victims, whose oxygen-carrying capacity is already reduced and in whom the administration of nitrite may form so much methemoglobin as to render the blood unable to carry oxygen at all.

Hydroxocobalamin, or vitamin B_{12a} , has recently been approved for use as an alternative cyanide antidote. Unlike sodium nitrite and sodium thiosulfate, it must be reconstituted at the scene. Hydroxocobalamin lacks the propensity of nitrites for hypotension, but many cases in which it has been beneficial have also required the use of sodium thiosulfate. Hydroxocobalamin causes an orange discoloration of the skin that is of no functional significance.

All of these antidotes require the placement of an IV line. Amyl nitrite is currently the only non-intravenous cyanide antidote available and has never been formally approved by the FDA. Amyl nitrite

TABLE 262e-4 ANTIDOTE RECOMMENDATIONS AFTER EXPOSURE TO CYANIDE^a

Patient	Mild (Conscious)	Severe (Unconscious)	Other Treatment
Child	If patient is conscious and has no other signs or symptoms, antidotes may not be necessary.	Sodium nitrite ^b : 0.12–0.33 mL/kg, not to exceed 10 mL of 3% solution, ^c by slow IV over no less than 5 min, or more slowly if hypotension develops and Sodium thiosulfate: 1.65 mL/kg of 25% solution IV over 10–20 min	Nasal oxygen supplementation For sodium nitrite–induced orthostatic hypotension, normal saline infusion and a supine position are recommended. If patient is still apneic after antidote administration, consider sodium bicarbonate for severe acidosis.
Adult	If patient is conscious and has no other signs or symptoms, antidotes may not be necessary.	Sodium nitrite ^b : 10–20 mL of 3% solution ^c by slow IV over no less than 5 min, or more slowly if hypotension develops and Sodium thiosulfate: 50 mL of 25% solution IV over 10–20 min Alternative: 5 g of hydroxocobalamin in reconstituted solution IV over 15 min	For amyl nitrite, ^b inhaled ampoules titrated to need or until other forms of IV therapy are initiated

^aVictims whose clothing or skin is contaminated with hydrogen cyanide liquid or solution can secondarily contaminate response personnel by direct contact or through off-gassing vapors. Dermal contact with cyanide-contaminated victims or with the gastric contents of victims who may have ingested cyanide-containing materials should be avoided. Victims exposed only to hydrogen cyanide gas do not pose contamination risks to rescuers. *If the patient is a victim of recent smoke inhalation (and thus may have high carboxyhemoglobin levels), only sodium thiosulfate should be administered.* ^bIf sodium nitrite is unavailable, administer amyl nitrite by inhalation from crushable ampoules. ^cAvailable in cyanide antidote kits, which can be purchased from various manufacturers.

Source: State of New York, Department of Health.

perles in cyanide antidote kits can be crushed and inhaled by a patient who is still breathing. Amyl nitrite can also be given through a respirator.

None of the cyanide antidotes is specifically approved for pediatric use.

Prognosis Cyanide casualties tend to recover much more quickly than casualties exposed to other chemical agents. Many patients with industrial cases have returned to work within the same shift. If a patient receives a large challenge and dies, death usually takes place within minutes of exposure.

263e Radiation Terrorism

Christine E. Hill-Kayser, Eli Glatstein,
Zelig A. Tochner

263e-1

CHAPTER 263e

Radiation Terrorism

The threat of a terror attack employing nuclear or radiation-related devices is unequivocal in the twenty-first century. Such an attack would certainly have the potential to cause unique and devastating medical and psychological effects that would require prompt action by members of the medical community. This chapter outlines the most probable scenarios for an attack involving radiation as well as the medical principles for handling such threats.

Potential terrorist incidents with radiologic consequences may be considered in two major categories. The first is the use of radiologic dispersal devices. Such devices disseminate radioactive material purposefully and without nuclear detonation. An attack with a goal of radiologic dispersal could take place through use of conventional explosives with incorporated radionuclides (“dirty bombs”), one or more fixed nuclear facilities, or nuclear-powered surface vessels or submarines. Other means could include detonation of malfunctioning nuclear weapons with no nuclear yield (nuclear “duds”) and installation of radionuclides in food or water. The second and less probable scenario is the actual use of nuclear weapons. Each scenario poses its own specific medical threats, including “conventional” blast or thermal injury, introduction to a radiation field, and exposure to either external or internal contamination from a radioactive explosion.

TYPES OF RADIOISOTOPIC RADIATION

Atomic isotopes with uneven numbers of protons and/or neutrons are typically unstable; such isotopes discharge particles or energy to matter as they move to stability, a process that is defined as *radiation*. Mass-containing particles, including alpha particles, electrons, and/or neutrons, may be transferred during this process (alpha radiation, beta radiation, and neutron radiation, respectively); alternatively, the transfer may consist only of energy in the form of a gamma ray. *Alpha* (α) *radiation* consists of heavy, positively charged particles, each of which contains two protons and two neutrons. Alpha particles usually are emitted from isotopes that have an atomic number of ≥ 82 , such as uranium and plutonium. Due to their large size, alpha particles have limited penetrating power. Fine obstacles such as cloth and human skin usually can stop these particles from penetrating into the body. Thus they represent a small risk from external exposure. If they are somehow internalized, however, alpha particles can cause significant damage to human cells that are in their immediate proximity.

Beta (β) *radiation* consists of electrons, which are small, light, negatively charged particles (about 1/2000 the mass of a neutron or proton). Electrons can travel only a short, finite distance in tissue, with the precise distance depending on their energy. Exposure to beta particles is common in many radiation accidents. Radioactive iodine released in nuclear plant accidents is the best-known form of beta radiation. Plastic layers and clothing can stop most beta particles, and their penetration is generally measured at a few millimeters. A large quantum of energy delivered via beta particles to the basal stratum of the skin can cause a burn that is similar to a thermal burn and is treated as such.

Gamma (γ) *rays* and *x-rays* (both photons) are similar. Gamma rays are uncharged electromagnetic radiation discharged from a nucleus as a wave of energy. X-rays are the product of abrupt mechanical deceleration of electrons striking a heavy target such as tungsten. Although they are generated by different sources, gamma rays and x-rays have similar properties; that is, they have no charge and no mass, just energy. They travel easily through matter and thus are sometimes referred to as *penetrating radiation*. Gamma rays and x-rays are the principal types of radiation that cause dangerous total-body exposure. Gamma rays and x-rays of the same energy will cause the same biologic effects, and these effects will require the same treatment.

Neutron (n) *particles* are heavy and uncharged and are often emitted during nuclear detonation. They possess a wide energy range; their

ability to penetrate tissues is variable, depending on their energy. They are less likely to be present in most scenarios of radiation bioterrorism than are the other forms of radiation discussed above.

Radiation interactions with atoms can result in ionization and the formation of free radicals that damage tissue by disrupting chemical bonds and molecular structures in the cell, including DNA. Protons, electrons, and gamma rays cause cellular damage through ionization of DNA. Depending on energy and other factors, some fraction of this damage will be caused by a direct strike to the DNA molecule (*direct ionization*). The remainder will be caused by ionization of water molecules to create free radicals that, in turn, damage DNA (*indirect ionization*). Ionization of DNA resulting from neutrons is exclusively indirect. Radiation damage can lead to cell death; the cells that recover may be mutated and at higher risk for subsequent cancer evolution. Cell sensitivity increases as the replication rate increases and cell differentiation decreases.

The commonly used units of radiation are the rad and the gray (Gy). The rad (*radiation absorbed dose*) is energy deposited within living matter and is equal to 100 ergs/g of tissue. The traditional rad has been replaced by the *Système Internationale* (SI) unit of the gray; 100 rad = 1 Gy, while 1 Gy is equal to 1 joule/kg. The sievert (Sv) is the SI unit that refers to the equivalent radiation dose in biologic tissues. While 1 Sv is equal to 1 joule/kg, Sv and Gy are not interchangeable units: Sv refers to the biologic effect of the radiation, while Gy refers to the physical energy being transferred.

TYPES OF EXPOSURE

Whole-body exposure occurs when radiation energy is deposited throughout the entire body. During a whole-body exposure, alpha and beta particles have limited penetration and do not cause significant noncutaneous injury unless emission results from an internalized source. Whole-body exposure from gamma rays, x-rays, or neutrons, which can penetrate through the body (the degree of which depends on their energy), can result in damage to multiple tissues and organs. The damage is proportional to the radiation exposure of the specific organ or tissue.

External contamination is a result of fallout of radioactive particles that land on the body surface, clothing, skin, and hair. This is the dominant element to consider in the mass-casualty situation resulting from a radioactive terrorist strike. The common contaminants primarily emit alpha and beta radiation. Alpha particles do not penetrate beyond the skin and thus have minimal systemic effects. Beta emitters can cause significant cutaneous burns and scarring. Due to their ability to penetrate tissue, gamma emitters can cause not only local damage but also whole-body radiation exposures and injury. Medical treatment primarily entails decontamination of the body, including wounds and burns, to prevent internalization of radioactive contaminants. Removal of contaminated clothing reduces levels of contamination significantly and is a first step in the decontamination process. Generally, patients do not constitute a significant radiation hazard to health care providers, and lifesaving treatment should not be delayed for fear of secondary contamination of the medical team. Although risk is relatively low, any damage to health care personnel will depend directly on the duration of exposure and will be inversely proportional to the square of the distance from any radioactive source. Gowns that can be easily removed offer protection.

Internal contamination occurs when radioactive material is inhaled or ingested or enters the body through open wounds or burns or via skin absorption. In principle, any externally contaminated casualty should be evaluated for internal contamination. Because of their chemical properties, some isotopes may exert toxic effects on specific target organs in addition to causing radiologic injury. The respiratory system is the main portal of entry for internal contamination, and the lung is the organ at greatest risk. Aerosol particles $< 5 \mu\text{m}$ in diameter can reach the alveoli, whereas larger particles will remain in the proximal airways. The tiny particles can be absorbed by the lymphatic system or the bloodstream. Bronchial lavage is often a helpful treatment in this situation. Radioactive material entering the gastrointestinal tract is absorbed according to its chemical structure and solubility.

The insoluble radionuclides may affect the lower gastrointestinal tract. Intact skin is normally an effective barrier to most radionuclides. Penetration through the skin usually takes place when wounds or burns have compromised the skin barrier. Therefore, any skin erosion should be cleaned and decontaminated promptly.

Absorbed radioactive materials travel throughout the body. Liver, kidney, adipose tissue, thyroid, and bone and bone marrow tend to bind and retain radioactive material more than other tissues do. Medical treatment thus includes the prevention of absorption, the reduction of incorporation, and the enhancement of elimination (see below).

Localized exposure refers to close contact between a highly radioactive source and a part of the body, with consequent discrete damage to the skin and deeper tissues that resembles a thermal burn. Later signs include epilation, erythema, moist desquamation, ulceration, blistering, and necrosis in proportion to exposure. Alopecia, transient or permanent, is dose related and starts at cutaneous doses of >3 Gy. Overt tissue damage can take weeks or even months to develop; the healing process can also be very slow, lasting for months. Long-term cutaneous changes, including keratosis, fibrosis, and telangiectasis, may appear years after the exposure. Treatment is based on analgesia and infection prophylaxis. Nevertheless, severe burns often require grafting or even amputation. Long-term radiation effects are characterized by cell loss, cell death, and tissue atrophy.

RADIOLOGIC DISPERSAL EVENTS

Radiologic dispersal incidents are generally of two types, resulting from small, usually localized sources or from wide dispersals over large areas. The methods that could be utilized to formulate an attack using dispersal of radiation are incredibly diverse. Radioactive materials can take the form of solid state, aerosol, gas, or liquid. They can be put into food or water, released from vehicles, or spread by explosion. The principal route of exposure is usually direct contact between the victim's skin and the radioactive particles, although internal contamination can occur if the material is inhaled or ingested. The radiation field is also a potential source of whole-body exposure. The psychosocial effects that accompany such an event are significant and are beyond the scope of this chapter. A list of radioactive materials, including information on their major properties and medical treatment, is given in [Table 263e-1](#).

In a localized event, the amount and spread of the radioactive materials are usually limited and can be treated like a spill of hazardous material. Protective clothing prevents or minimizes the contamination of emergency responders.

The use of explosives coupled with a large amount of radioactive materials can result in wide dispersion of radiation, which is of far greater concern. Other potential sources of radiation are nuclear reactors, spent nuclear fuel, and transport vehicles. Less probable but still possible is the use of a large source of penetrating radiation without explosion. It is expected that most exposures would be low-level and that the principal health and psychosocial effects would be similar to those in the former scenario but on a larger scale. Whenever an explosion is involved, conventional lifesaving treatment should be given first priority. Only then should decontamination and specific treatment be given for the radiation exposure.

Silent exposure represents a scenario in which a powerful radiologic source, often called a *radiologic exposure device*, could be hidden in a crowded place and spread radioactive materials without being recognized or reported. Recognition of the event and the source of exposure might take a long time. A major clue in this situation is the appearance of unusual clinical manifestations in many individuals; such manifestations are often nonspecific and include symptoms of acute radiation sickness (see "Acute Radiation Syndrome," below) such as headache, fatigue, malaise, and opportunistic infections. Gastrointestinal phenomena such as diarrhea, nausea, vomiting, and anorexia may occur. Dermatologic symptoms (e.g., burns, ulceration, and epilation) and hematopoietic manifestations (e.g., bleeding tendency, thrombocytopenia, purpura, lymphopenia, and neutropenia) are also possible and are dose-related. Careful epidemiologic studies may be necessary to identify the source of such exposure.

NUCLEAR WEAPONS

The most likely scenario in a nuclear terror attack is the detonation of a single low-yield device. The estimated yield of such a device is anywhere between 0.01 and 10 kilotons of 2,4,6-trinitrotoluene (TNT). The expected effects of such an explosion are a combination of several components: ground shock, air blast, thermal radiation, initial nuclear radiation, crater formation, and radioactive fallout.

A nuclear detonation, like a conventional explosion, produces a shock wave that can further damage structures and cause many casualties. In addition, the detonation can produce an extremely hot fireball that can ignite materials and cause severe burns. The detonation releases an intense pulse of ionizing radiation consisting mainly of gamma rays and neutrons. The radiation produced in the first minute is termed *initial radiation*, whereas the ongoing radiation due to fallout is termed *residual radiation*. Both types of radiation can cause acute radiation sickness, and winds can carry fallout and contaminate large areas. The $LD_{50/30}$ (i.e., the dose that causes a 50% mortality rate at 30 days) is ~4 Gy for whole-body exposure without medical support; with medical support, the $LD_{50/30}$ ranges between 8 and 10 Gy. On top of its immediate effects, a massive blast forms a crater in the soil and usually produces ground shock that compounds the physical damage and the number of casualties. Inhalation of large amounts of radioactive dust causes pneumonitis that can lead to pulmonary fibrosis. Use of a mask covering the mouth and nose can result in effective prevention. The intense flash of infrared and visible light can cause either temporary or permanent blindness. Cataracts can develop months to years later among survivors.

ACUTE RADIATION SYNDROME

Acute radiation syndrome (ARS) refers to multisystem symptomatology resulting hours to weeks after radiation exposure. As discussed earlier, cell sensitivity to radiation damage increases as the cell replication rate increases and as cell differentiation decreases. Bone marrow and mucosal surfaces of the gastrointestinal tract, which have vast mitotic activity, are significantly more sensitive to radiation than are slowly dividing tissues such as bones and muscles. After exposure of all or most of the human body to ionizing radiation, ARS can develop. The clinical manifestations of ARS reflect the dose and type of radiation as well as the parts of the body exposed.

ARS manifests as three major groups of signs and symptoms: hematopoietic, gastrointestinal, and neurovascular. In addition, ARS exists in four stages: prodrome, latent phase, clinical illness, and recovery or death. The higher the radiation doses, the shorter and more severe each stage. The prodrome appears within minutes to 4 days after exposure, lasts from a few hours to a few days, and can include nausea, vomiting, anorexia, and diarrhea. At the end of the prodrome, ARS progresses to the latent phase. Minimal or no symptoms are present during the latent phase, which commonly lasts up to 2.5 weeks but can last up to 6 weeks. The duration depends on the radiation dose, the prior health of the patient, and coexisting illness or injury. After the latent phase, the exposed person manifests illness that may end in recovery or lead to death.

With exposure to low doses of <1 Gy, ARS is generally mild. At this dose, symptoms can be minimal or nonexistent, even if the entire body is exposed to penetrating radiation. The main feature of the clinical picture is transient depression of bone marrow (lymphopenia) that lasts up to 2–3 weeks and then improves.

ARS is significantly more acute and severe with exposure to very high radiation doses (>30 Gy). At these doses, the prodrome appears in minutes and is followed by 5–6 hours of latency before cardiovascular collapse occurs secondary to irreversible damage to the microcirculation.

Exposure to intermediate radiation doses may result in variable ARS courses. The type and dose of radiation and the part of the body exposed determine not only the timing of the different stages of ARS but also the dominant clinical picture. At low radiation doses of 0.7–4 Gy, hematopoietic depression due to bone marrow suppression is the main constituent of illness. The patient may develop infection and bleeding secondary to low leukocyte and platelet counts, respectively. The bone marrow eventually recovers in almost all patients if

TABLE 263e-1 INTERNAL CONTAMINANT RADIONUCLIDES: PROPERTIES AND TREATMENT

Isotope Name	Symbol	Common Usage	Radiation Type; $t_{1/2}$ Radiologic; $t_{1/2}$ Biologic (days)	Exposure Type	Mode of Contamination	Focal Accumulation in Body	Treatment
Manganese	Mn-56	Reactors, research laboratories	β , γ ; 2.6 h; 5.7	External, internal	N/A	Liver	N/A
Cobalt	Co-60	Medical radiotherapy devices, commercial food irradiators	β , γ ; 5.26 y; 9.5	External, internal	Lungs	Liver	Gastric lavage, purgatives; penicillamine in severe cases
Strontium	Sr-90	Fission product of uranium	β ; 28 y; 18,000	Internal	GI tract	Bones (similar to calcium)	Strontium, calcium, ammonium chloride
Molybdenum	Mo-99	Hospitals: scans	β , γ ; 66.7 h; 3	External, internal	N/A	Kidneys	N/A
Technetium	Tc-99m	Hospitals: scans	β , γ ; 6.049 h; 1	External, internal	IV administration	Kidneys, total body	Potassium perchlorate to reduce thyroid dose
Cesium	Cs-137	Medical radiotherapy devices	β , γ ; 30 y; 70	External, internal	Lungs, GI tract, wounds; follows potassium	Renal excretion	Ion-exchange resins, Prussian blue
Gadolinium	Gd-153	Hospitals	β , γ ; 242 d; 1000	External, internal	N/A	N/A	N/A
Iridium	Ir-192	Commercial radiography	β , γ ; 74 d; 50	External, internal	N/A	Spleen	N/A
Radium	Ra-226	Instrument illumination, industrial applications, old medical equipment, former Soviet Union military equipment	α , β , γ ; 1602 y; 16,400	External, internal	GI tract	Bones	MgSO ₄ lavage, ammonium chloride, calcium alginates
Tritium	H-3	Luminescent gunsights, muzzle-velocity detectors, nuclear weapons	β ; 12.5 y; 12	Internal	Inhalation, GI tract, wounds	Total body	Dilution with controlled water intake, diuretics
Iodine-131	¹³¹ I	Reactors, thyroid ablaters	β , γ ; 8.1 d; 138	Internal	Inhalation, GI tract, wounds	Thyroid	Potassium/sodium iodide, propylthiouracil, methimazole
Uranium	U-235	Depleted uranium, natural uranium, fuel rods, weapons-grade material	α , (α , β , γ); 7.1 \times 10 ⁸ y; 15	Internal	GI tract	Kidneys, bones	NaHCO ₃ , chelation with EDTA
Plutonium	Pu-239	Produced from uranium in reactors, nuclear weapons	α ; 2.2 \times 10 ⁴ y; 73,000	Internal	Limited lung absorption, high retention	Lungs, bones, bone marrow, liver, gonads	Chelation with DTPA or EDTA
Americium	Am-241	Smoke detectors, nuclear weapons (in form of fallout)	α ; 458 y; 73,000	Internal	Inhalation, skin wounds	Lungs, liver, bones, bone marrow	Chelation with DTPA or EDTA
Polonium	Po-210	Calibration source	α ; 138.4 d; 60	Internal	Inhalation, wounds	Spleen, kidneys	Lavage, dimercaprol
Thorium	Th-232	Calibration source	α ; 1.41 \times 10 ¹⁰ y; 73,000	Internal	N/A	N/A	N/A
Phosphorus	P-32	Research laboratories, medical facilities	β ; 14.3 d; 1155	Internal	Inhalation, GI tract, wounds	Bones, bone marrow, rapidly replicating cells	Lavage, aluminum hydroxide, phosphate

Abbreviations: DTPA, diethylenetriamine pentaacetic acid; EDTA, ethylenediamine tetraacetic acid; GI, gastrointestinal; h, hours; N/A, not available; y, years.

they are supported with transfusions and fluids; antibiotics are often needed in addition. Patients with isolated hematopoietic manifestations of ARS can almost always survive with proper supportive care. After exposure to 6–8 Gy, a significantly more complicated clinical picture may ensue. At these doses, the bone marrow does not always recover and death may occur as a result. A gastrointestinal syndrome may accompany the hematopoietic manifestations and further worsen the patient's condition. Gastrointestinal injury due to compromise of the absorptive layer of the gut alters absorption of fluids, electrolytes, and nutrients. Such injury can lead to vomiting, diarrhea, gastrointestinal bleeding, sepsis, and electrolyte and fluid imbalance. Generally, these symptoms are also accompanied by a severe hematopoietic syndrome, with only a slim chance of bone marrow recovery. These factors in constellation often lead to death. Whole-body exposure to >9–10 Gy is almost always fatal. Crucial elements of the bone marrow

simply do not recover. In addition to the gastrointestinal syndrome associated with very high-level exposures, patients may develop a neurovascular syndrome that includes vascular collapse, seizures, and confusion; death occurs within a few days. The neurovascular syndrome dominates after whole-body exposure to >20 Gy. In this variant, the prodrome and latent phase both last only a few hours.

TREATMENT ACUTE RADIATION SICKNESS

The treatment of ARS is focused on maintaining homeostasis, thus giving damaged organs a chance to recover. Aggressive support is provided for every damaged system. Treatment for the hematopoietic system targets mainly neutropenia and infection, with measures that may include transfusion of leukoreduced irradiated

blood as needed and administration of hematopoietic growth factors. The value of bone marrow transplantation in this situation is questionable. None of the bone marrow transplantations that were performed among the victims of the nuclear reactor accident in Chernobyl proved successful. Bone marrow transplantation could be considered for casualties with whole-body exposure to 6–10 Gy when the hematopoietic syndrome is dominant and the bone marrow is less likely to recover with time, although the efficacy of this treatment has not been proved. Another major component of the treatment of ARS is the provision of partial or total parenteral nutrition to bypass the damaged gastrointestinal system. For blast and thermal injuries, standard therapy for trauma is given. Psychological support is essential in many cases. A treatment algorithm is outlined in Fig. 263e-1.

MEDICAL MANAGEMENT OF RADIATION BIOTERRORISM

Victims of radiation bioterrorism can suffer from conventional thermal or blast injuries, exposure to radiation, and contamination by radioactive materials. Many will have combinations of the above, which can cause higher morbidity and mortality rates than each exposure would alone. The number of casualties will be a major factor in determining the response of the medical system to an act of radiation bioterrorism. If only a few persons are affected, no significant changes and adaptations of the system are needed to treat the victims. If a terror attack results in dozens of casualties or more, however, an organized disaster plan at the local and state levels must be invoked to deal with the crisis properly. Useful U.S. planning documents that include many universal planning concepts can be found at http://www.remm.nlm.gov/remm_Preplanning.htm. Ideally, medical personnel will have had a prior assignment and training and be prepared to function in a scenario with which they are familiar. Stockpiles of specific equipment

and medications should be obtained ahead of time and stored safely (see the Centers for Disease Control and Prevention Web site at <http://www.bt.cdc.gov/stockpile/>). One of the goals of terrorist attackers is to overwhelm medical facilities and minimize the salvage of casualties. Initial management consists of *primary triage and transportation* of the wounded to medical facilities for treatment. The rationale behind triage is to sort patients into classes according to the severity of injury for the purpose of expediting clinical care and maximizing the use of the available clinical services and facilities. Triage requires determination of the level of emergency care needed. The higher the number and broader the range of casualties, the more complex and difficult triage becomes. The mildly wounded and victims of contamination only can be sent to evacuation, registration (with disaster response teams), and decontamination/treatment centers. Figure 263e-2 illustrates an evacuation scheme after a radiologic event causing multiple casualties. The goal of such an algorithm is to treat all possible victims of exposure and minor injury outside of the hospital setting. This approach prevents hospitals from being directly overwhelmed and enhances treatment for persons who are severely wounded. Emergency treatment should be administered initially for conventional injuries such as wounds, trauma, and thermal or chemical burns. Individuals with such injuries should be stabilized, if possible, and immediately transported to a medical facility. Removing clothing and wrapping the victim in clean blankets or nylon sheets reduce both the exposure of the patient and the contamination risk to the staff. Less severely injured victims should undergo preliminary decontamination before or during evacuation to a hospital.

One must remember that radionuclide contamination of the skin commonly is not an acute life-threatening situation for the patient or for the personnel who care for the patient. Only powerful gamma emitters are likely to cause real damage from contamination. It is important to emphasize that exposure to a radiation field alone does not necessarily create any contamination. The exposed person, if not contaminated, is not radioactive and does not directly emit any radiation.

To protect the staff, protective gear (gowns, gloves, masks, and caps) should be used. Protective masks with filters and chemically protective overgarments provide excellent protection from contamination. Waterproof shoe covers are also important. Remaining in the contaminated area and dealing with life-saving procedures should take place according to the “ALARA” principle: *as low as reasonably achievable*. It is better to send many people to do the job for short exposure times than to send a few people for longer periods.

Decontamination of victims should take place in the field before their arrival at medical facilities, but radiologic decontamination should never interfere with medical care. Removal of outer clothing and shoes usually reduces a patient’s contamination by 80–90%. Contaminated clothes should be carefully removed by rolling them over themselves, placed in marked plastic bags, and removed to a predefined area for contaminated clothes and equipment. A radiation detector should then be used to check for the presence of any residual radiologic contamination on the patient’s body. To prevent internalization of the radioactive materials, one should cover open wounds before decontamination. Showering or washing of the entire skin and hair is very important and should be done as soon as possible. The skin should then be dried and reassessed for residual contamination until no radiation is found.

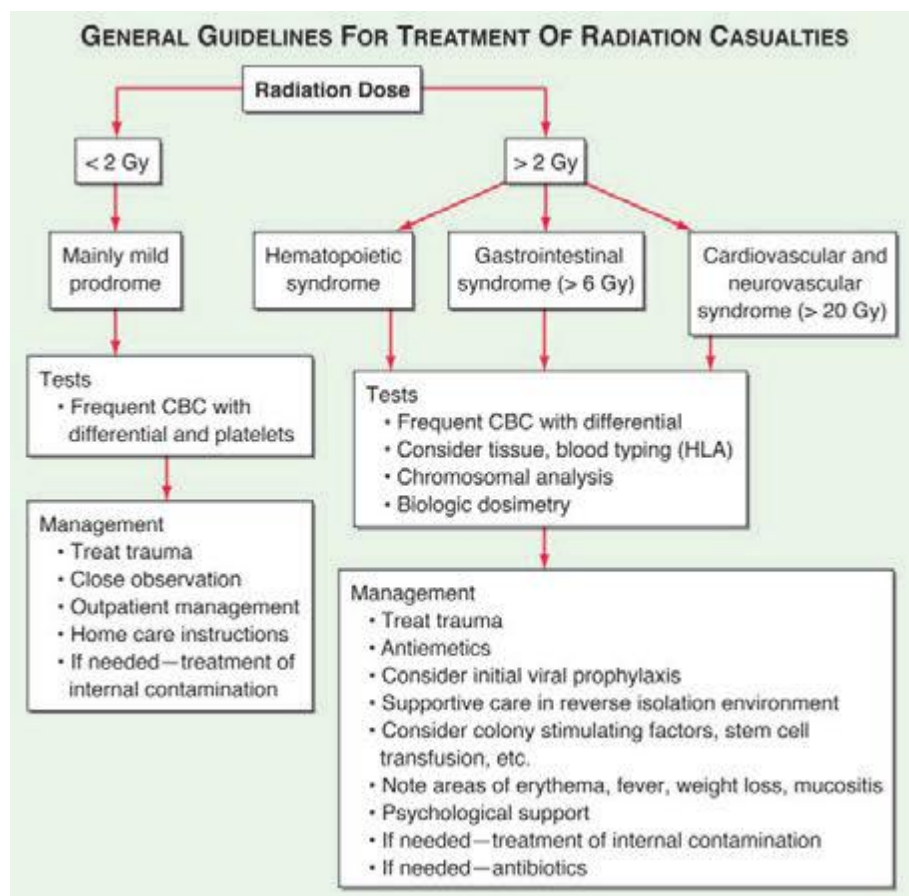


FIGURE 263e-1 General guidelines for treatment of radiation casualties. CBC, complete blood count.

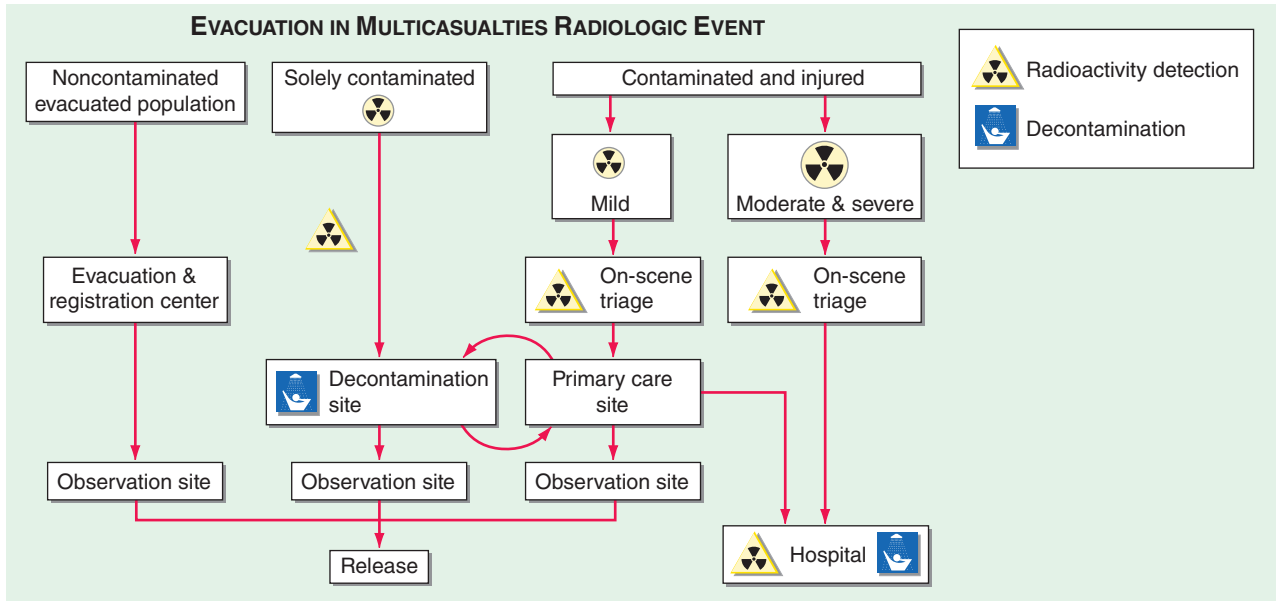


FIGURE 263e-2 Algorithm for evacuation in a multicasualties radiologic event.

Contamination-removing chemical agents are more than sufficient to remove radiologic contamination.

Wound decontamination should be as conservative as possible. The main goal is to prevent both extensive local damage and internal contamination through lacerated skin. The bandages should be removed and the wounds flushed. The wound should then be dried and assessed for radiation. This procedure can be repeated again and again until contamination is undetectable. Excision of contaminated wounds should be attempted only when surgically necessary. Radioactive shrapnel that can penetrate through the skin should be removed.

In the hospital, staff can wear normal hospital barrier clothing, including two pairs of gloves, a gown, shoe covers, a head cover, and a face mask. Eye protection is recommended. Decontamination of medical personnel is obligatory after emergency treatment and decontamination of the patient. After use, all protective clothing should be placed in a designated container for contaminated clothing.

Radiation intensity decays rapidly with the square of the distance from the source; thus increasing the distance from the source and decreasing the time spent near it are basic principles of radiation safety. Shielding with lead can be used as protection from small radioactive gamma sources. Geiger counters can detect gamma and beta radiation. Pocket chambers and dosimeters, film badges, and thermoluminescent dosimeters can measure accumulated exposure to gamma radiation. All these detectors are in common use in medical facilities and should be employed to help define the level of contamination. Alpha radiation is harder to detect due to its poor penetration. An alpha scintillation counter, which is capable of detecting alpha radiation, is not commonly used in medical facilities.

GUIDELINES FOR HOSPITAL MANAGEMENT

Figure 263e-3 shows a model for hospital arrangement for triage. Persons contaminated either externally or internally should be identified, externally decontaminated, and, if need be, treated immediately and specifically for internal contamination as detailed below. In all other cases, the need for treatment of radiation injuries does not constitute a medical emergency. Early actions—e.g., blood sampling both to assess the severity of the exposure and to perform blood typing and cross-matching for possible transfusion—need to be taken promptly if ARS is evident or if whole-body exposure is suspected.

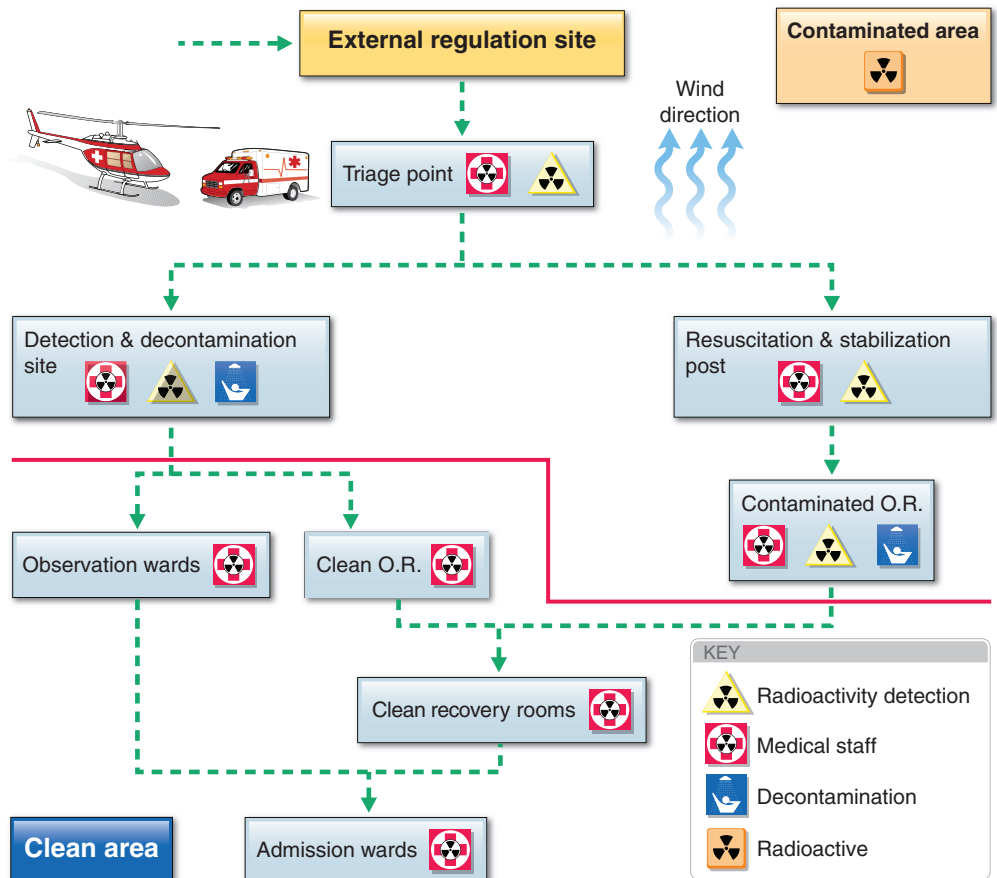


FIGURE 263e-3 Flow chart of hospital triage. O.R., operating room.

In the hospital entrance, a distinct decontamination area should be set up promptly. Separation between clean and contaminated areas is essential. Medical personnel in contaminated areas should wear protective gear, as noted above, and should be rotated in their assignments every 1–2 h to ensure minimal exposure to radiation. If patients are critically wounded and require either surgery or resuscitation, they need to be transported directly to “contaminated” operating rooms or resuscitation sites for lifesaving procedures. Once the condition of such patients is stable, they should be decontaminated. It is important to obtain details concerning the exposure, to look for prodromal signs of radiation sickness, and to conduct a physical examination. One of the simplest ways to estimate exposure clinically is to measure the time of prodromal appearance. The earlier the prodromal signs and symptoms appear, the higher is the dose of radiation exposure. A few laboratory tests need to be done routinely, such as complete blood count and urinalysis. If internal contamination is suspected, specific treatment should be given as outlined below.

TREATMENT RADIONUCLIDE CONTAMINATION

Table 263e-2 summarizes the common treatment regimens for internal radionuclide contamination. Treatment for internal radionuclide contamination, also referred to as *decorporation*, should be started as soon as possible after suspected or known exposure. The approximate upper limit of radionuclide contamination that can reasonably be ignored from a radiation safety point of view

is not well defined. These are judgments that will depend on the circumstances of the event and the resources available. The Clinical Decision Guide within the National Council on Radiation Protection and Measurements (NCRP) Report 161 is a decision tool for determining the need for treatment of a contaminated person. Purchase of these volumes by major triage centers (available at http://www.ncrppublications.org/Reports/161_1) may be a prudent investment that would help health care workers in a critical situation to determine which patients should undergo decorporation.

The goal is to leave the smallest amount of radionuclide possible in the body. Treatment is given to reduce absorption and enhance elimination and excretion. Some decorporation agents are not approved by the U.S. Food and Drug Administration (FDA) for these indications, and few clinical data support the efficacy of their use.

The gastrointestinal tract may be cleared by stomach lavage, with emetics (such as apomorphine, 5–10 mg; or ipecac, 1- to 2-g capsules or 15 mL in syrup), or by use of purgatives, laxatives, ion exchangers, and aluminum antacids. Prussian blue (1 g tid for a minimum of 3 weeks) is an ion exchanger used to treat cesium-137 internal contamination. Aluminum antacids (such as aluminum phosphate gel) may reduce strontium uptake in the gut if given immediately after exposure. Aluminum hydroxide is less effective.

Radionuclide interaction with tissues can be prevented or reversed through use of agents that block absorption; dilute, mobilize, or release radionuclides from tissues; or chelate radionuclides.

TABLE 263e-2 COMMON DRUGS^a FOR TREATMENT OF INTERNAL RADIONUCLIDE CONTAMINATION

Medication	Administered for Radionuclides	Route of Administration	Dosage	Duration	Mechanism of Action
KI	¹³¹ I	PO	130 mg/d for adults >40 with thyroid exposure >500 cGy, for adults 18–40 with thyroid exposure >10 cGy, and for pregnant or lactating women with thyroid exposure >5 cGy; 65 mg/d for children and adolescents 3–18 with thyroid exposure >5 cGy; 32.5 mg/d for infants 1 mo to 3 y with thyroid exposure >5 cGy; 16 mg/d for neonates from birth to 1 mo with thyroid exposure >5 cGy	7–14 d	Blocking agent
Zn-DTPA	Plutonium, <i>trans</i> -plutonium, yttrium, americium, curium	IV Inhalation IM	1 g in 250 mL NS or 5% glucose, given over 1–2 h or as a bolus over 3–4 min 1 g in 1:1 dilution with water or NS over 15–20 min 1 g; not recommended because of pain	Up to 5 d	Chelating agent
Ca-DTPA	Plutonium, <i>trans</i> -plutonium, yttrium, americium, curium	IV Inhalation IM	1 g in 250 mL NS or 5% glucose, given over 1–2 h or as a bolus over 3–4 min 1 g in 1:1 dilution with water or NS over 15–20 min 1 g; not recommended because of pain	Up to 5 d	Chelating agent
Bicarbonate	Uranium	IV PO	2 ampoules NaHCO ₃ (44.3 meq each, 7.5%) in 1000 mL NS, 125 mL/L; or 1 ampoule NaHCO ₃ (44.3 meq, 7.5%) in 500 mL NS, 500 mL/h 2 tablets q4 h until urine pH = 7–8, or 4 g (8 tablets) tid	Usually IV for the first 24 h, PO for additional 2 d; continuation of treatment for >3 d is rare and is based on titration of uranium amounts in the body	Increased excretion via the kidneys
Prussian blue	¹³⁷ Cs	PO	1 g tid with 100–200 mL water, up to 10 g/d	3 wk titrated by urine and fecal bioassay and whole-body counting	Ion exchanger
Water	Tritium (³ H)	PO	>3–4 L/d	3 wk	Excretion of water
Aluminum phosphate gel	Strontium	PO	100 mL immediately after exposure	Once	Decreased gut absorption
Aluminum hydroxide	Strontium	PO	60–100 mL	Once	Decreased gut absorption

^aExcept for KI and Prussian blue, these drugs had not been approved for this purpose by the U.S. Food and Drug Administration at the time of publication.

Abbreviation: NS, normal saline.

Blocking agents prevent the entrance of radioactive materials. The best-recognized effective blocking agent is potassium iodide (KI), which blocks the uptake of radioactive iodine (^{131}I) by the thyroid. KI is most effective if taken within the first hour after exposure and is still effective 6 h after exposure. Its effectiveness subsequently declines until 24 h after exposure; however, it is recommended that KI be taken up to 48 h after exposure. The KI dose is based on age, predicted thyroid exposure, and pregnancy and lactation status. Adults between the ages of 18 and 40 should receive 130 mg/d for 7–14 days if exposed to ≥ 10 cGy of radioactive iodine. Other thyroid-blocking agents include propylthiouracil (100 mg tid for 8 days) and methimazole (10 mg tid for 2 days followed by 5 mg tid for 6 days). These agents are somewhat less effective than KI.

Diluting agents decrease the absorption of the radionuclide; for example, water may be used as a diluting agent in the treatment for tritium (^3H) contamination. The recommended amount is 3–4 L/d for at least 3 weeks.

Mobilizing agents are most effective when given immediately; however, they may be effective for up to 2 weeks after exposure. These agents include antithyroid drugs, parathyroid extract, glucocorticoids, ammonium chloride, diuretics, expectorants, and inhalants. All of the latter agents should induce the release of radionuclides from tissues.

Chelating agents can bind many radioactive materials, after which the complexes are excreted from the body. In this regard, diethylenetriaminepentaacetic acid (DTPA)—as either Ca-DTPA or Zn-DTPA—is superior to ethylenediaminetetraacetic acid (EDTA); DTPA has been approved by the FDA to treat internal contamination with plutonium, americium, and curium, but it also chelates berkelium, californium, and any other material with an atomic number > 92 . Ca-DTPA is more effective than Zn-DTPA during the first 24 h after internal contamination, and the two drugs are equally effective after the initial 24 h. If both drugs are available, Ca-DTPA should be given as the first dose. If additional treatment is needed, treatment should be switched to Zn-DTPA. The dose is 1 g of Ca-DTPA or Zn-DTPA, dissolved in 250 mL of normal saline or 5% glucose and given intravenously over 1 h daily. The duration of chelation treatment depends on the amount of internal contamination and the individual response to treatment. DTPA also can be administered by nebulized inhalation; 1 g is given in a 1:1 dilution with water or saline over 15–20 min. Nebulized Zn-DTPA is recommended if inhalation was the only route of internal contamination. The IV route is recommended and should be used if the route of internal contamination is not known or if multiple routes of internal contamination are likely. DTPA penta-ethyl ester is a prodrug that has a favorable oral-absorption profile and whose therapeutic effects have been demonstrated in initial efficacy studies. Because it can be given orally, this prodrug may ultimately prove more useful in the setting of mass casualties than IV or nebulized forms of the drug. Treating uranium contamination with DTPA is contraindicated due to synergistic damage to the kidneys.

Lung lavage can reduce radiation-induced pneumonitis and is indicated only when a large amount of radionuclide enters the lungs and has the potential to cause acute radiation injury. The procedure requires anesthesia.

MEDICAL ASSAY OF RADIATION IN THE EXPOSED PATIENT

One of the major difficulties in treating victims exposed to radiation is determination of the amount of exposure. Immediately after a terrorist event, when victims are being triaged, information regarding source, dose, and exposure time is unlikely to be available. Clinical assessment of the patient is the best approach and includes history, physical examination, and observation for onset of the ARS prodrome. An early prodrome indicates high-level exposure to radiation. Victims who arrive at the hospital reporting severe weakness, nausea, vomiting, diarrhea, or seizures probably will not survive despite supportive

measures. A very limited number of tests can be performed to estimate radiation exposure and contamination. The Biodosimetry Assessment Tool (BAT) facilitates treatment decisions during radiation exposure incidents. Developed by the U.S. Armed Forces Radiobiology Research Institute (AFRRI), the BAT provides a method of estimating radiation exposure on the basis of a single lymphocyte count, the lymphocyte depletion rate, and the time from exposure to onset of emesis. The BAT algorithms are based on large datasets from human radiation exposure and are available at <http://www.usuhs.mil/afri/outreach/request.htm>. The patient should be observed for clinical symptoms, and the severity and time of onset of nausea, vomiting, headache, anorexia, fever, hypotension, tachycardia, weakness, cognitive changes, skin desquamation, diarrhea, and bloody stools should be recorded. The AFRRI Biodosimetry Worksheet (<http://www.usuhs.mil/afri/outreach/pdf/afriiform331.pdf>) is a useful resource for detailed recording. Baseline tests should include a complete blood count with differential and platelet count, renal evaluation, and determination of electrolytes, serum amylase, and serum C-reactive protein. Urine and stool samples should be obtained if internal contamination is suspected. Nasal swabs taken from each nostril within the first 1–2 h after the exposure may be useful for determination of radionuclide inhalation. After exhalation, each swab is labeled, sealed in a plastic bag, and sent for analysis to appropriate laboratories. Patients exposed to 0.7–4 Gy develop pancytopenia from as early as 10 days to as late as 8 weeks after exposure. Lymphocytes show the most rapid decline, whereas counts of other leukocytes and platelets decline less rapidly. Erythrocytes are the least vulnerable blood elements.

Absolute lymphocyte counts should be repeated every 4–6 h for 5–6 days; they are the most valuable early indicator because they constitute a sensitive marker for radiation damage and correlate with both the exposure and the prognosis. A 50% drop in absolute lymphocyte count within the first 24 h indicates a significant injury. HLA typing is necessary whenever irreversible bone marrow damage is suspected. Lymphocyte chromosomal analysis can detect exposure to as little as 0.03–0.06 Gy, and 15 mL of blood for this purpose should be drawn as early as possible in a heparinized collection tube and kept cool. Radiation-induced chromosomal aberrations visible in peripheral-blood lymphocytes include dicentric chromosomes and ring forms that last for a few weeks. Calibration of a dose-response curve makes it possible to assess the radiation dose on the basis of the presence of these aberrations. Dicentric quantification requires multiple days to perform and is available only in select centers.

Another method for estimating exposure is the *in vitro* cytokinesis-block micronucleus assay. Micronuclei can be the result of small acentric chromosome fragments that arise during exposure to radiation. The technique to score the micronuclei in peripheral-blood lymphocytes has been standardized in the last few years. It can be a useful tool in small-scale exposures but is not feasible in a mass casualty setting.

FOLLOW-UP

It is desirable to continue follow-up over the long term in some circumstances. In general, only persons who are exposed to < 8 – 10 Gy of whole-body irradiation have a chance to survive in the long term, and they are at risk of developing cataracts, sterility, and cancers as well as lung, kidney, and bone marrow problems. In light of their age, their gender, and the amount and type of exposure, they should be followed for many years. A major public health issue is the risk of secondary malignancy in individuals and populations that have been exposed to low doses of radiation. Leukemia and breast, brain, thyroid, and lung cancer develop most commonly, but the exposed population is at increased risk for many other cancers as well. Appropriate follow-up protocols should be based on the type of exposure and the exposed population. In cases of internal contamination, long-term follow-up should be focused on the organ at risk. Substantial psychosocial support will likely be needed for a community in the years after an attack including radiologic agents.

This page intentionally left blank

SECTION 1 INTRODUCTION TO CARDIOVASCULAR DISORDERS

264 Approach to the Patient with Possible Cardiovascular Disease

Joseph Loscalzo

THE MAGNITUDE OF THE PROBLEM

Cardiovascular diseases comprise the most prevalent serious disorders in industrialized nations and are a rapidly growing problem in developing nations (Chap. 266e). Age-adjusted death rates for coronary heart disease have declined by two-thirds in the last four decades in the United States, reflecting the identification and reduction of risk factors as well as improved treatments and interventions for the management of coronary artery disease, arrhythmias, and heart failure. Nonetheless, cardiovascular diseases remain the most common causes of death, responsible for 35% of all deaths, almost 1 million deaths each year. Approximately one-fourth of these deaths are sudden. In addition, cardiovascular diseases are highly prevalent, diagnosed in 80 million adults, or ~35% of the adult population. The growing prevalence of obesity (Chap. 416), type 2 diabetes mellitus (Chap. 417), and metabolic syndrome (Chap. 422), which are important risk factors for atherosclerosis, now threatens to reverse the progress that has been made in the age-adjusted reduction in the mortality rate of coronary heart disease.

For many years cardiovascular disease was considered to be more common in men than in women. In fact, the percentage of all deaths secondary to cardiovascular disease is higher among women (43%) than among men (37%) (Chap. 6e). In addition, although the absolute number of deaths secondary to cardiovascular disease has declined over the past decades in men, this number has actually risen in women. Inflammation, obesity, type 2 diabetes mellitus, and the metabolic syndrome appear to play more prominent roles in the development of coronary atherosclerosis in women than in men. Coronary artery disease (CAD) is more frequently associated with dysfunction of the coronary microcirculation in women than in men. Exercise electrocardiography has a lower diagnostic accuracy in the prediction of epicardial obstruction in women than in men.

NATURAL HISTORY

Cardiovascular disorders often present acutely, as in a previously asymptomatic person who develops an acute myocardial infarction (Chap. 295), or a previously asymptomatic patient with hypertrophic cardiomyopathy (Chap. 287), or with a prolonged QT interval (Chap. 277) whose first clinical manifestation is syncope or even sudden death. However, the alert physician may recognize the patient at risk for these complications long before they occur and often can take measures to prevent their occurrence. For example, a patient with acute myocardial infarction will often have had risk factors for atherosclerosis for many years. Had these risk factors been recognized, their elimination or reduction might have delayed or even prevented the infarction. Similarly, a patient with hypertrophic cardiomyopathy may have had a heart murmur for years and a family history of this disorder. These findings could have led to an echocardiographic examination, recognition of the condition, and appropriate therapy long before the occurrence of a serious acute manifestation.

Patients with valvular heart disease or idiopathic dilated cardiomyopathy, by contrast, may have a prolonged course of gradually increasing dyspnea and other manifestations of chronic heart failure that is punctuated by episodes of acute deterioration only late in the course of the disease. Understanding the natural history of various cardiac

disorders is essential for applying appropriate diagnostic and therapeutic measures to each stage of the condition, as well as for providing the patient and family with the likely prognosis.

CARDIAC SYMPTOMS

The symptoms caused by heart disease result most commonly from myocardial ischemia, disturbance of the contraction and/or relaxation of the myocardium, obstruction to blood flow, or an abnormal cardiac rhythm or rate. Ischemia, which is caused by an imbalance between the heart's oxygen supply and demand, is manifest most frequently as chest discomfort (Chap. 19), whereas reduction of the pumping ability of the heart commonly leads to fatigue and elevated intravascular pressure upstream of the failing ventricle. The latter results in abnormal fluid accumulation, with peripheral edema (Chap. 50) or pulmonary congestion and dyspnea (Chap. 47e). Obstruction to blood flow, as occurs in valvular stenosis, can cause symptoms resembling those of myocardial failure (Chap. 279). Cardiac arrhythmias often develop suddenly, and the resulting symptoms and signs—palpitations (Chap. 52), dyspnea, hypotension, and syncope (Chap. 27)—generally occur abruptly and may disappear as rapidly as they develop.

Although dyspnea, chest discomfort, edema, and syncope are cardinal manifestations of cardiac disease, they occur in other conditions as well. Thus, dyspnea is observed in disorders as diverse as pulmonary disease, marked obesity, and anxiety (Chap. 47e). Similarly, chest discomfort may result from a variety of noncardiac and cardiac causes other than myocardial ischemia (Chap. 19). Edema, an important finding in untreated or inadequately treated heart failure, also may occur with primary renal disease and in hepatic cirrhosis (Chap. 50). Syncope occurs not only with serious cardiac arrhythmias but in a number of neurologic conditions as well (Chap. 27). Whether heart disease is responsible for these symptoms frequently can be determined by carrying out a careful clinical examination (Chap. 267), supplemented by noninvasive testing using electrocardiography at rest and during exercise (Chap. 268), echocardiography, roentgenography, and other forms of myocardial imaging (Chap. 270e).

Myocardial or coronary function that may be adequate at rest may be insufficient during exertion. Thus, dyspnea and/or chest discomfort that appear during activity are characteristic of patients with heart disease, whereas the opposite pattern, i.e., the appearance of these symptoms at rest and their remission during exertion, is rarely observed in such patients. It is important, therefore, to question the patient carefully about the relation of symptoms to exertion.

Many patients with cardiovascular disease may be asymptomatic both at rest and during exertion but may present with an abnormal physical finding such as a heart murmur, elevated arterial pressure, or an abnormality of the electrocardiogram (ECG) or imaging test. It is important to assess the global risk of CAD in asymptomatic individuals, using a combination of clinical assessment and measurement of cholesterol and its fractions, as well as other biomarkers, such as C-reactive protein, in some patients (Chap. 291e). Since the first clinical manifestation of CAD may be catastrophic—sudden cardiac death, acute myocardial infarction, or stroke in previous asymptomatic persons—it is mandatory to identify those at high risk of such events and institute further testing and preventive measures.

DIAGNOSIS

As outlined by the New York Heart Association (NYHA), the elements of a complete cardiac diagnosis include the systematic consideration of the following:

1. *The underlying etiology.* Is the disease congenital, hypertensive, ischemic, or inflammatory in origin?

TABLE 264-1 NEW YORK HEART ASSOCIATION FUNCTIONAL CLASSIFICATION

Class I	Class III
No limitation of physical activity	Marked limitation of physical activity
No symptoms with ordinary exertion	Less than ordinary activity causes symptoms
Class II	Asymptomatic at rest
Slight limitation of physical activity	Class IV
Ordinary activity causes symptoms	Inability to carry out any physical activity without discomfort
	Symptoms at rest

Source: Modified from The Criteria Committee of the New York Heart Association.

2. *The anatomic abnormalities.* Which chambers are involved? Are they hypertrophied, dilated, or both? Which valves are affected? Are they regurgitant and/or stenotic? Is there pericardial involvement? Has there been a myocardial infarction?
3. *The physiologic disturbances.* Is an arrhythmia present? Is there evidence of congestive heart failure or myocardial ischemia?
4. *Functional disability.* How strenuous is the physical activity required to elicit symptoms? The classification provided by the NYHA has been found to be useful in describing functional disability (Table 264-1).

One example may serve to illustrate the importance of establishing a complete diagnosis. In a patient who presents with exertional chest discomfort, the identification of myocardial ischemia as the etiology is of great clinical importance. However, the simple recognition of ischemia is insufficient to formulate a therapeutic strategy or prognosis until the underlying anatomic abnormalities responsible for the myocardial ischemia, e.g., coronary atherosclerosis or aortic stenosis, are identified and a judgment is made about whether other physiologic disturbances that cause an imbalance between myocardial oxygen supply and demand, such as severe anemia, thyrotoxicosis, or supraventricular tachycardia, play contributory roles. Finally, the severity of the disability should govern the extent and tempo of the workup and strongly influence the therapeutic strategy that is selected.

The establishment of a correct and complete cardiac diagnosis usually commences with the history and physical examination (Chap. 267). Indeed, the clinical examination remains the basis for the diagnosis of a wide variety of disorders. The clinical examination may then be supplemented by five types of laboratory tests: (1) ECG (Chap. 268), (2) noninvasive imaging examinations (chest roentgenogram, echocardiogram, radionuclide imaging, computed tomographic imaging, positron emission tomography, and magnetic resonance imaging) (Chap. 270e), (3) blood tests to assess risk (e.g., lipid determinations, C-reactive protein [Chap. 291e]) or cardiac function (e.g., brain natriuretic peptide [BNP] [Chap. 279]), (4) occasionally specialized invasive examinations (i.e., cardiac catheterization and coronary arteriography [Chap. 272]), and (5) genetic tests to identify monogenic cardiac diseases (e.g., hypertrophic cardiomyopathy [Chap. 287], Marfan's syndrome [Chap. 427], and abnormalities of cardiac ion channels that lead to prolongation of the QT interval and an increase in the risk of sudden death [Chap. 276]). These tests are becoming more widely available.

FAMILY HISTORY

In eliciting the history of a patient with known or suspected cardiovascular disease, particular attention should be directed to the family history. Familial clustering is common in many forms of heart disease. Mendelian transmission of single-gene defects may occur, as in hypertrophic cardiomyopathy (Chap. 287), Marfan's syndrome (Chap. 427), and sudden death associated with a prolonged QT syndrome (Chap. 277). Premature coronary disease and essential hypertension, type 2 diabetes mellitus, and hyperlipidemia (the most important risk factors for CAD) are usually polygenic disorders. Although familial transmission may be less obvious than in the monogenic disorders, it is helpful in assessing risk and prognosis in polygenic disorders, as well.

Familial clustering of cardiovascular diseases not only may occur on a genetic basis but also may be related to familial dietary or behavior patterns, such as excessive ingestion of salt or calories and cigarette smoking.

ASSESSMENT OF FUNCTIONAL IMPAIRMENT

When an attempt is made to determine the severity of functional impairment in a patient with heart disease, it is helpful to ascertain the level of activity and the rate at which it is performed before symptoms develop. Thus, it is not sufficient to state that the patient complains of dyspnea. The breathlessness that occurs after running up two long flights of stairs denotes far less functional impairment than do similar symptoms that occur after taking a few steps on level ground. Also, the degree of customary physical activity at work and during recreation should be considered. The development of two-flight dyspnea in a well-conditioned marathon runner may be far more significant than the development of one-flight dyspnea in a previously sedentary person. The history should include a detailed consideration of the patient's therapeutic regimen. For example, the persistence or development of edema, breathlessness, and other manifestations of heart failure in a patient who is receiving optimal doses of diuretics and other therapies for heart failure (Chap. 279) is far graver than are similar manifestations in the absence of treatment. Similarly, the presence of angina pectoris despite treatment with optimal doses of multiple anti-anginal drugs (Chap. 293) is more serious than it is in a patient on no therapy. In an effort to determine the progression of symptoms, and thus the severity of the underlying illness, it may be useful to ascertain what, if any, specific tasks the patient could have carried out 6 months or 1 year earlier that he or she cannot carry out at present.

ELECTROCARDIOGRAM

(See also Chap. 268) Although an ECG usually should be recorded in patients with known or suspected heart disease, with the exception of the identification of arrhythmias, conduction abnormalities, ventricular hypertrophy, and acute myocardial infarction, it generally does not establish a specific diagnosis. The range of normal electrocardiographic findings is wide, and the tracing can be affected significantly by many noncardiac factors, such as age, body habitus, and serum electrolyte concentrations. In general, electrocardiographic changes should be interpreted in the context of other abnormal cardiovascular findings.

ASSESSMENT OF THE PATIENT WITH A HEART MURMUR

(Fig. 264-1) The cause of a heart murmur can often be readily elucidated from a systematic evaluation of its major attributes: timing, duration, intensity, quality, frequency, configuration, location, and radiation when considered in the light of the history, general physical examination, and other features of the cardiac examination, as described in Chap. 267.

The majority of heart murmurs are midsystolic and soft (grades I–II/VI). When such a murmur occurs in an asymptomatic child or young adult *without* other evidence of heart disease on clinical examination, it is usually benign and echocardiography generally is not required. By contrast, two-dimensional and Doppler echocardiography (Chap. 270e) are indicated in patients with loud systolic murmurs (grades \geq III/VI), especially those that are holosystolic or late systolic, and in most patients with diastolic or continuous murmurs.

PITFALLS IN CARDIOVASCULAR MEDICINE

Increasing subspecialization in internal medicine and the perfection of advanced diagnostic techniques in cardiology can lead to several undesirable consequences. Examples include the following:

1. Failure by the *noncardiologist* to recognize important cardiac manifestations of systemic illnesses. For example, the presence of mitral stenosis, patent foramen ovale, and/or transient atrial arrhythmia should be considered in a patient with stroke, or the presence of pulmonary hypertension and cor pulmonale should be considered in a patient with scleroderma or Raynaud's syndrome. A cardiovascular examination should be carried out to identify and estimate

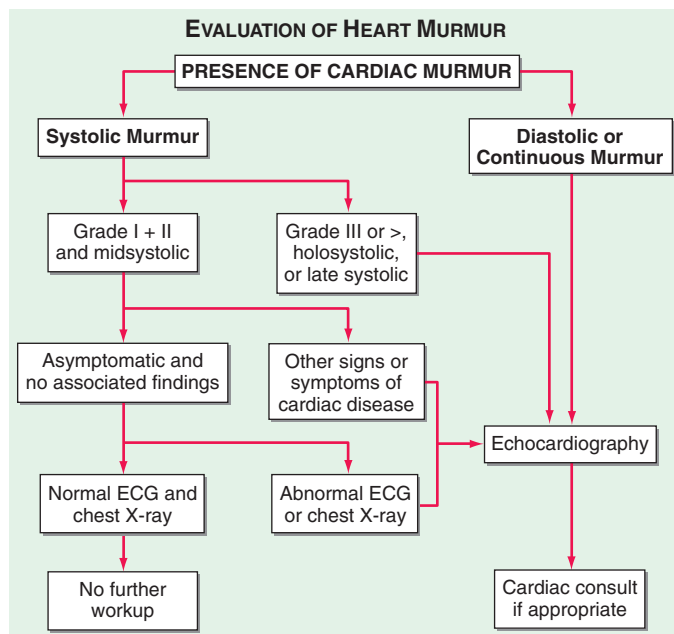


FIGURE 264-1 Approach to the evaluation of a heart murmur.

ECG, electrocardiogram. (From RA O'Rourke, in *Primary Cardiology*, 2nd ed, E Braunwald, L Goldman [eds]. Philadelphia, Saunders, 2003.)

the severity of the cardiovascular involvement that accompanies many noncardiac disorders.

- Failure by the *cardiologist* to recognize underlying systemic disorders in patients with heart disease. For example, hyperthyroidism should be considered in an elderly patient with atrial fibrillation and unexplained heart failure, and Lyme disease should be considered in a patient with unexplained fluctuating atrioventricular block. A cardiovascular abnormality may provide the clue critical to the recognition of some systemic disorders. For example, an unexplained pericardial effusion may provide an early clue to the diagnosis of tuberculosis or a neoplasm.
- Overreliance on and overutilization of laboratory tests, particularly invasive techniques, for the evaluation of the cardiovascular system. Cardiac catheterization and coronary arteriography (Chap. 272) provide precise diagnostic information that may be crucial in developing a therapeutic plan in patients with known or suspected CAD. Although a great deal of attention has been directed to these examinations, it is important to recognize that they serve to *supplement*, not *supplant*, a careful examination carried out with clinical and noninvasive techniques. A coronary arteriogram should not be performed in lieu of a careful history in patients with chest pain suspected of having ischemic heart disease. Although coronary arteriography may establish whether the coronary arteries are obstructed and to what extent, the results of the procedure by themselves often do not provide a definitive answer to the question of whether a patient's complaint of chest discomfort is attributable to coronary atherosclerosis and whether or not revascularization is indicated.

Despite the value of invasive tests in certain circumstances, they entail some small risk to the patient, involve discomfort and substantial cost, and place a strain on medical facilities. Therefore, they should be carried out only if the results can be expected to modify the patient's management.

DISEASE PREVENTION AND MANAGEMENT

The prevention of heart disease, especially of CAD, is one of the most important tasks of primary care health givers as well as cardiologists. Prevention begins with risk assessment, followed by attention to lifestyle, such as achieving optimal weight, physical activity, and smoking cessation, and then aggressive treatment of all abnormal risk factors, such as hypertension, hyperlipidemia, and diabetes mellitus (Chap. 417).

After a complete diagnosis has been established in patients with known heart disease, a number of management options are usually

available. Several examples may be used to demonstrate some of the principles of cardiovascular therapeutics:

- In the absence of evidence of heart disease, the patient should be clearly informed of this assessment and *not* be asked to return at intervals for repeated examinations. If there is no evidence of disease, such continued attention may lead to the patient's developing inappropriate concern about the possibility of heart disease.
- If there is no evidence of cardiovascular disease but the patient has one or more risk factors for the development of ischemic heart disease (Chap. 293), a plan for their reduction should be developed and the patient should be retested at intervals to assess compliance and efficacy in risk reduction.
- Asymptomatic or mildly symptomatic patients with valvular heart disease that is anatomically severe should be evaluated periodically, every 6 to 12 months, by clinical and noninvasive examinations. Early signs of deterioration of ventricular function may signify the need for surgical treatment before the development of disabling symptoms, irreversible myocardial damage, and excessive risk of surgical treatment (Chap. 283).
- In patients with CAD (Chap. 293), available practice guidelines should be considered in the decision on the form of treatment (medical, percutaneous coronary intervention, or surgical revascularization). Mechanical revascularization may be employed too frequently in the United States and too infrequently in Eastern Europe and developing nations. The mere presence of angina pectoris and/or the demonstration of critical coronary arterial narrowing at angiography should not reflexively evoke a decision to treat the patient by revascularization. Instead, these interventions should be limited to patients with CAD whose angina has not responded adequately to medical treatment or in whom revascularization has been shown to improve the natural history (e.g., acute coronary syndrome or multivessel CAD with left ventricular dysfunction).

THE BLOOD VESSEL

VASCULAR ULTRASTRUCTURE

Blood vessels participate in homeostasis on a moment-to-moment basis and contribute to the pathophysiology of diseases of virtually every organ system. Hence, an understanding of the fundamentals of vascular biology furnishes a foundation for understanding the normal function of all organ systems and many diseases. The smallest blood vessels—capillaries—consist of a monolayer of endothelial cells apposed to a basement membrane, adjacent to occasional smooth-muscle-like cells known as *pericytes* (Fig. 265e-1A). Unlike larger vessels, pericytes do not invest the entire microvessel to form a continuous sheath. Arteries typically have a trilaminar structure (Fig. 265e-1B–E). The *intima* consists of a monolayer of endothelial cells continuous with those of the capillaries. The middle layer, or *tunica media*, consists of layers of smooth-muscle cells; in veins, the media can contain just a few layers of smooth-muscle cells (Fig. 265e-1B). The outer layer, the *adventitia*, consists of looser extracellular matrix with occasional fibroblasts, mast cells, and nerve terminals. Larger arteries have their own vasculature, the *vasa vasorum*, which nourishes the outer aspects of the tunica media. The adventitia of many veins surpasses the intima in thickness.

The tone of muscular arterioles regulates blood pressure and flow through various arterial beds. These smaller arteries have a relatively thick tunica media in relation to the adventitia (Fig. 265e-1C). Medium-size muscular arteries similarly contain a prominent tunica media (Fig. 265e-1D); atherosclerosis commonly affects this type of muscular artery. The larger elastic arteries have a much more structured tunica media consisting of concentric bands of smooth-muscle cells, interspersed with strata of elastin-rich extracellular matrix

sandwiched between layers of smooth-muscle cells (Fig. 265e-1E). Larger arteries have a clearly demarcated internal elastic lamina that forms the barrier between the intima and the media. An external elastic lamina demarcates the media of arteries from the surrounding adventitia.

ORIGIN OF VASCULAR CELLS

The intima in human arteries often contains occasional resident smooth-muscle cells beneath the monolayer of vascular endothelial cells. The embryonic origin of smooth-muscle cells in various types of artery differs. Some upper-body arterial smooth-muscle cells derive from the neural crest, whereas lower-body arteries generally recruit smooth-muscle cells from neighboring mesodermal structures during development. Derivatives of the proepicardial organ, which gives rise to the epicardial layer of the heart, contribute to the vascular smooth-muscle cells of the coronary arteries. Bone marrow–derived endothelial progenitors may aid repair of damaged or aging arteries. In addition, multipotent vascular stem cells resident in vessel walls may give rise to the smooth-muscle cells that accumulate in injured or atheromatous arteries (Chaps. 88, 89e, and 90e).

VASCULAR CELL BIOLOGY

Endothelial Cell The key cell of the vascular intima, the endothelial cell, has manifold functions in health and disease. The endothelium forms the interface between tissues and the blood compartment. It therefore must regulate the entry of molecules and cells into tissues in a selective manner. The ability of endothelial cells to serve as a selectively permeable barrier fails in many vascular disorders, including atherosclerosis, hypertension, and renal disease. This dysregulation of permeability also occurs in pulmonary edema and other situations of “capillary leak.”

The endothelium also participates in the local regulation of blood flow and vascular caliber. Endogenous substances produced by endothelial cells such as prostacyclin, endothelium-derived hyperpolarizing factor, nitric oxide (NO), and hydrogen peroxide (H₂O₂) provide tonic vasodilatory stimuli under physiologic conditions in vivo (Table 265e-1). Impaired production or excess catabolism of NO

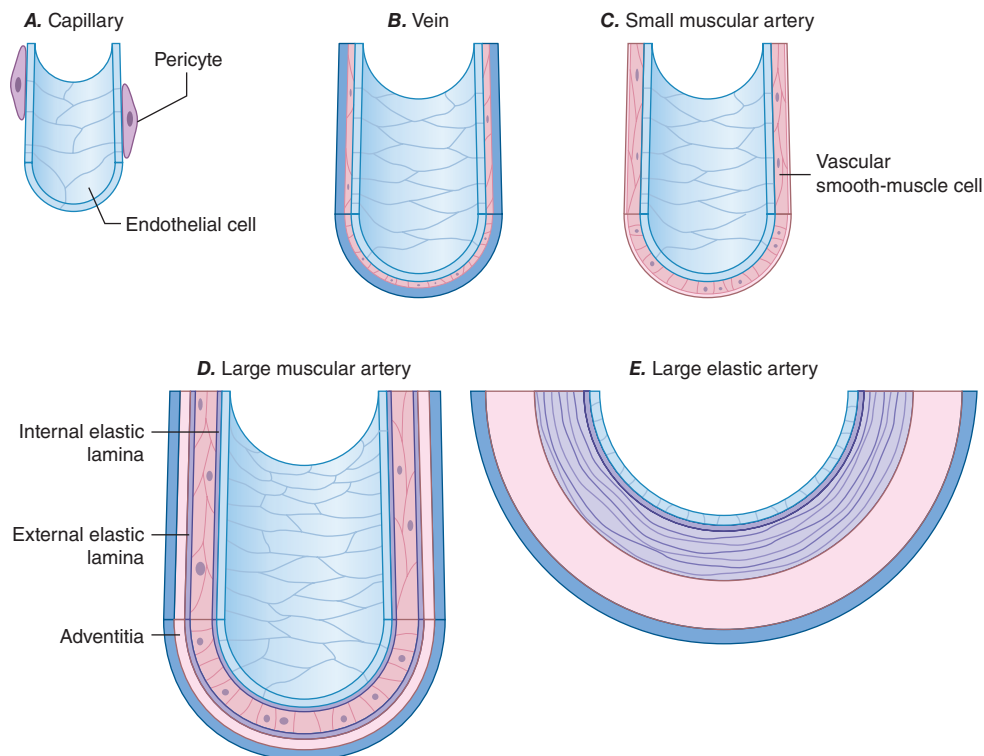


FIGURE 265e-1 Schematics of the structures of various types of blood vessels. **A.** Capillaries consist of an endothelial tube in contact with a discontinuous population of pericytes. **B.** Veins typically have thin medias and thicker adventitias. **C.** A small muscular artery features a prominent tunica media. **D.** Larger muscular arteries have a prominent media with smooth-muscle cells embedded in a complex extracellular matrix. **E.** Larger elastic arteries have cylindrical layers of elastic tissue alternating with concentric rings of smooth-muscle cells.

TABLE 265e-1 ENDOTHELIAL FUNCTIONS IN HEALTH AND DISEASE

Homeostatic Properties	Dysfunctional Properties
Optimize balance between vasodilation and vasoconstriction	Impaired dilation, vasoconstriction
Antithrombotic, profibrinolytic	Prothrombotic, antifibrinolytic
Anti-inflammatory	Proinflammatory
Antiproliferative	Proproliferative
Antioxidant	Prooxidant
Permselectivity	Impaired barrier function

impairs this endothelium-dependent vasodilator function and may contribute to excessive vasoconstriction in various pathologic situations. Measurement of flow-mediated dilatation can assess endothelial vasodilator function in humans (Fig. 265e-2). By contrast, endothelial cells also produce potent vasoconstrictor substances such as endothelin in a regulated fashion. Excessive production of reactive oxygen species, such as superoxide anion (O_2^-), by endothelial or smooth-muscle cells under pathologic conditions (e.g., excessive exposure to angiotensin II), can promote local oxidative stress and inactivate NO.

The endothelial monolayer contributes critically to inflammatory processes involved in normal host defenses and pathologic states. The normal endothelium resists prolonged contact with blood leukocytes; however, when activated by bacterial products such as endotoxin or by proinflammatory cytokines released during infection or injury, endothelial cells express an array of leukocyte adhesion molecules that bind various classes of leukocytes. The endothelial cells appear to recruit selectively different classes of leukocytes in different pathologic conditions. The gamut of adhesion molecules and chemokines generated during acute bacterial infection tends to recruit granulocytes. In chronic inflammatory diseases such as tuberculosis and atherosclerosis, endothelial cells express adhesion molecules that favor the recruitment of mononuclear leukocytes that characteristically accumulate in these conditions.

The endothelium also dynamically regulates thrombosis and hemostasis. NO, in addition to its vasodilatory properties, can limit platelet activation and aggregation. Like NO, prostacyclin produced by endothelial cells under normal conditions not only provides a vasodilatory stimulus but also antagonizes platelet activation and aggregation. Thrombomodulin expressed on the surface of endothelial cells binds thrombin at low concentrations and inhibits coagulation through activation of the protein C pathway, inactivating clotting factors Va and VIIIa and thus combating thrombus formation. The surface of endothelial cells contains heparan sulfate glycosaminoglycans that furnish an endogenous antithrombotic coating to the vasculature. Endothelial cells also participate actively in fibrinolysis and its regulation. They express receptors for plasminogen and plasminogen activators and produce tissue-type plasminogen activator. Through local generation of plasmin, the normal endothelial monolayer can promote the lysis of nascent thrombi.

When activated by inflammatory cytokines, bacterial endotoxin, or angiotensin II, for example, endothelial cells can produce substantial quantities of the major inhibitor of fibrinolysis, plasminogen activator inhibitor 1 (PAI-1). Thus, in pathologic circumstances, the endothelial cell may promote local thrombus accumulation rather than combat it. Inflammatory stimuli also induce the expression of the potent procoagulant tissue factor, a contributor to disseminated intravascular coagulation in sepsis.

Endothelial cells also participate in the pathophysiology of a number of immune-mediated diseases. Lysis of endothelial cells mediated by complement provides an example of immunologically mediated tissue injury. The presentation of foreign histocompatibility complex antigens by endothelial cells in solid-organ allografts can promote allograft arteriopathy. In addition, immune-mediated endothelial injury may contribute in some patients with thrombotic thrombocytopenic purpura and patients with hemolytic-uremic syndrome. Thus, in addition to the involvement of innate immune responses, endothelial cells participate actively in both humoral and cellular limbs of the immune response.

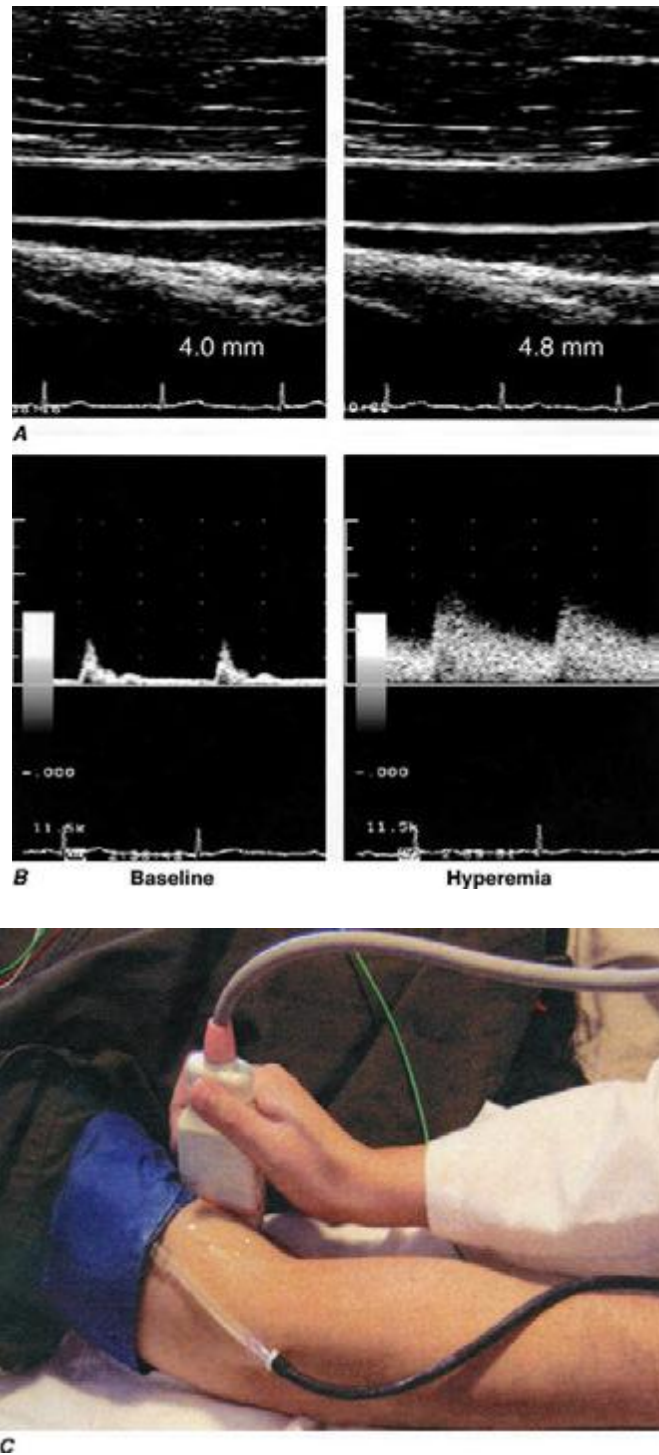


FIGURE 265e-2 Assessment of endothelial function in vivo using blood pressure cuff occlusion and release. Upon deflation of the cuff, an ultrasound probe monitors changes in diameter (A) and blood flow (B) of the brachial artery (C). (Reproduced with permission of J. Vita, MD.)

Endothelial cells regulate growth of subjacent smooth-muscle cells. Heparan sulfate glycosaminoglycans elaborated by endothelial cells can inhibit smooth-muscle proliferation. In contrast, when exposed to various injurious stimuli, endothelial cells can elaborate growth factors and chemoattractants, such as platelet-derived growth factor, that can promote the migration and proliferation of vascular smooth-muscle cells. Dysregulated elaboration of these growth-stimulatory molecules may promote smooth-muscle accumulation in atherosclerotic lesions.

Vascular Smooth-Muscle Cell The vascular smooth-muscle cell, the major cell type of the media layer of blood vessels, also contributes actively to vascular pathobiology. Contraction and relaxation of smooth-muscle cells at the level of the muscular arteries controls blood pressure and, hence, regional blood flow and the afterload experienced by the left ventricle (see below). The vasomotor tone of veins, which is governed by smooth-muscle cell tone, regulates the capacitance of the venous tree and influences the preload experienced by both ventricles. Smooth-muscle cells in the adult vessel seldom replicate in the absence of arterial injury or inflammatory activation. Proliferation and migration of arterial smooth-muscle cells, associated with functional modulation characterized by lower content of contractile proteins and greater production of extracellular matrix macromolecules, can contribute to the development of arterial stenoses in atherosclerosis, arteriolar remodeling that can sustain and propagate hypertension, and the hyperplastic response of arteries injured by percutaneous intervention. In the pulmonary circulation, smooth-muscle migration and proliferation contribute decisively to the pulmonary vascular disease that gradually occurs in response to sustained high-flow states such as left-to-right shunts. Such pulmonary vascular disease provides a major obstacle to the management of many patients with adult congenital heart disease. Among other mediators, microRNAs have emerged as powerful regulators of this transition, offering new targets for intervention.

Smooth-muscle cells secrete the bulk of vascular extracellular matrix. Excessive production of collagen and glycosaminoglycans contributes to the remodeling and altered functions and biomechanics of arteries affected by hypertension or atherosclerosis. In larger elastic arteries, the elastin synthesized by smooth-muscle cells serves

to maintain not only normal arterial structure, but also hemodynamic function. The ability of the larger arteries, such as the aorta, to store the kinetic energy of systole promotes tissue perfusion during diastole. Arterial stiffness associated with aging or disease, as manifested by a widening pulse pressure, increases left ventricular afterload and portends a poor outcome.

Like endothelial cells, vascular smooth-muscle cells do not merely respond to vasomotor or inflammatory stimuli elaborated by other cell types but can themselves serve as a source of such stimuli. For example, when exposed to bacterial endotoxin or other proinflammatory stimuli, smooth-muscle cells can elaborate cytokines and other inflammatory mediators. Like endothelial cells, upon inflammatory activation, arterial smooth-muscle cells can produce prothrombotic mediators such as tissue factor, the antifibrinolytic protein PAI-1, and other molecules that modulate thrombosis and fibrinolysis. Smooth-muscle cells also elaborate autocrine growth factors that can amplify hyperplastic responses to arterial injury.

Vascular Smooth-Muscle Cell Function Vascular smooth-muscle cells govern vessel tone. Those cells contract when stimulated by a rise in intracellular calcium concentration by calcium influx through the plasma membrane and by calcium release from intracellular stores (Fig. 265e-3). In vascular smooth-muscle cells, voltage-dependent L-type calcium channels open with membrane depolarization, which is regulated by energy-dependent ion pumps such as the Na^+, K^+ -ATPase pump and ion channels such as the Ca^{2+} -sensitive K^+ channel. Local changes in intracellular calcium concentration, termed *calcium sparks*, result from the influx of calcium through the voltage-dependent calcium channel and are caused by the coordinated activation of a

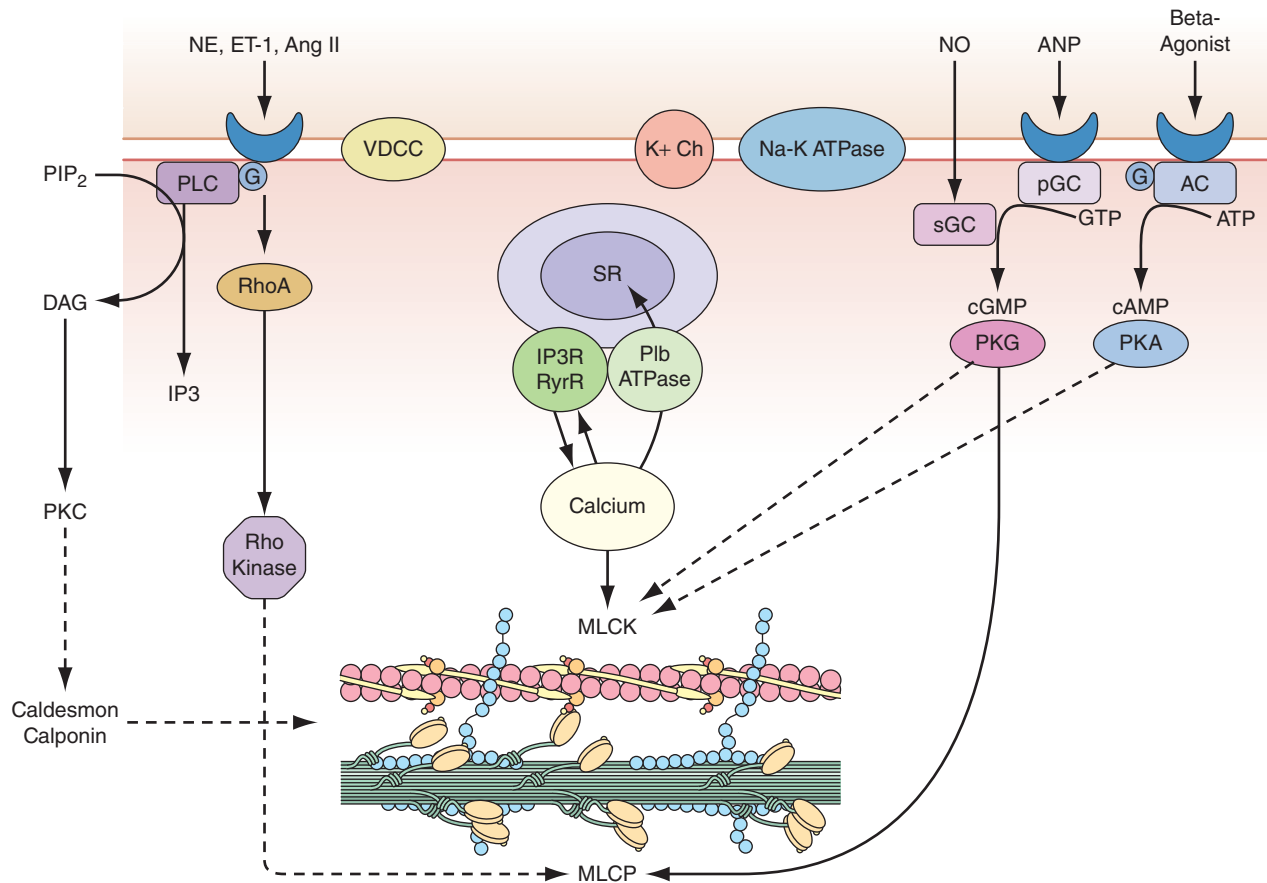


FIGURE 265e-3 Regulation of vascular smooth-muscle cell calcium concentration and actomyosin ATPase-dependent contraction.

AC, adenylyl cyclase; Ang II, angiotensin II; ANP, atrial natriuretic peptide; DAG, diacylglycerol; ET-1, endothelin-1; G, G protein; IP_3 , inositol 1,4,5-trisphosphate; MLCK, myosin light chain kinase; MLCP, myosin light chain phosphatase; NE, norepinephrine; NO, nitric oxide; pGC, particulate guanylyl cyclase; PIP_2 , phosphatidylinositol 4,5-bisphosphate; PKA, protein kinase A; PKC, protein kinase C; PKG, protein kinase G; PLC, phospholipase C; sGC, soluble guanylyl cyclase; SR, sarcoplasmic reticulum; VDCC, voltage-dependent calcium channel. (Modified from B Berk, in *Vascular Medicine*, 3rd ed. Philadelphia, Saunders, Elsevier, 2006, p. 23; with permission.)

cluster of ryanodine-sensitive calcium release channels in the sarcoplasmic reticulum (see below). Calcium sparks directly augment intracellular calcium concentration and indirectly increase intracellular calcium concentration by activating chloride channels. In addition, calcium sparks reduce smooth-muscle contractility by activating large-conductance calcium-sensitive K^+ channels, hyperpolarizing the cell membrane and thereby limiting further voltage-dependent increases in intracellular calcium.

Biochemical agonists also increase intracellular calcium concentration, in this case by receptor-dependent activation of phospholipase C with hydrolysis of phosphatidylinositol 4,5-bisphosphate, resulting in the generation of diacylglycerol (DAG) and inositol 1,4,5-trisphosphate (IP_3). These membrane lipid derivatives in turn activate protein kinase C and increase intracellular calcium concentration. In addition, IP_3 binds to specific receptors on the sarcoplasmic reticulum membrane to increase calcium efflux from this calcium storage pool into the cytoplasm.

Vascular smooth-muscle cell contraction depends principally on the phosphorylation of myosin light chain, which in the steady state, reflects the balance between the actions of myosin light chain kinase and myosin light chain phosphatase. Calcium activates myosin light chain kinase through the formation of a calcium-calmodulin complex. Phosphorylation of myosin light chain by this kinase augments myosin ATPase activity and enhances contraction. Myosin light chain phosphatase dephosphorylates myosin light chain, reducing myosin ATPase activity and contractile force. Phosphorylation of the myosin-binding subunit (thr695) of myosin light chain phosphatase by Rho kinase inhibits phosphatase activity and induces calcium sensitization of the contractile apparatus. Rho kinase is itself activated by the small GTPase RhoA, which is stimulated by guanosine exchange factors and inhibited by GTPase-activating proteins.

Both cyclic AMP and cyclic GMP relax vascular smooth-muscle cells through complex mechanisms. β agonists, acting through their G-protein-coupled receptors activate adenylyl cyclase to convert ATP to cyclic AMP; NO and atrial natriuretic peptide acting directly and via a G-protein-coupled receptor, respectively, activate guanylyl cyclase to convert GTP to cyclic GMP. These agents in turn activate protein kinase A and protein kinase G, respectively, which inactivate myosin light chain kinase and decrease vascular smooth-muscle cell tone. In addition, protein kinase G can interact directly with the myosin-binding substrate subunit of myosin light chain phosphatase, increasing phosphatase activity and decreasing vascular tone. Finally, several mechanisms drive NO-dependent, protein kinase G-mediated reductions in vascular smooth-muscle cell calcium concentration, including phosphorylation-dependent inactivation of RhoA; decreased IP_3 formation; phosphorylation of the IP_3 receptor-associated cyclic GMP kinase substrate, with subsequent inhibition of IP_3 receptor function; phosphorylation of phospholamban, which increases calcium ATPase activity and sequestration of calcium in the sarcoplasmic reticulum; and protein kinase G-dependent stimulation of plasma membrane calcium ATPase activity, perhaps by activation of the Na^+,K^+ -ATPase pump or hyperpolarization of the cell membrane by activation of calcium-dependent K^+ channels.

Control of Vascular Smooth-Muscle Cell Tone The autonomic nervous system and endothelial cells modulate vascular smooth-muscle cells in a tightly regulated manner. Autonomic neurons enter the blood vessel medial layer from the adventitia and modulate vascular smooth-muscle cell tone in response to baroreceptors and chemoreceptors within the aortic arch and carotid bodies and in response to thermoreceptors in the skin. These regulatory components include rapidly acting reflex arcs modulated by central inputs that respond to sensory inputs (olfactory, visual, auditory, and tactile) as well as emotional stimuli. Three classes of nerves mediate autonomic regulation of vascular tone: *sympathetic*, whose principal neurotransmitters are epinephrine and norepinephrine; *parasympathetic*, whose principal neurotransmitter is acetylcholine; and *nonadrenergic/noncholinergic*, which include two subgroups—nitrenergic, whose principal neurotransmitter

is NO, and peptidergic, whose principal neurotransmitters are substance P, vasoactive intestinal peptide, calcitonin gene-related peptide, and ATP.

Each of these neurotransmitters acts through specific receptors on the vascular smooth-muscle cell to modulate intracellular calcium and, consequently, contractile tone. Norepinephrine activates α receptors, and epinephrine activates α and β receptors (adrenergic receptors); in most blood vessels, norepinephrine activates postjunctional α_1 receptors in large arteries and α_2 receptors in small arteries and arterioles, leading to vasoconstriction. Most blood vessels express β_2 -adrenergic receptors on their vascular smooth-muscle cells and respond to β agonists by cyclic AMP-dependent relaxation. Acetylcholine released from parasympathetic neurons binds to muscarinic receptors (of which there are five subtypes, M_{1-5}) on vascular smooth-muscle cells to yield vasorelaxation. In addition, NO stimulates presynaptic neurons to release acetylcholine, which can stimulate the release of NO from the endothelium. Nitrenergic neurons release NO produced by neuronal NO synthase, which causes vascular smooth-muscle cell relaxation via the cyclic GMP-dependent and -independent mechanisms described above. The peptidergic neurotransmitters all potently vasodilate, acting either directly or through endothelium-dependent NO release to decrease vascular smooth-muscle cell tone. **For the detailed molecular physiology of the autonomic nervous system, see Chap. 454.**

The endothelium modulates vascular smooth-muscle tone by the direct release of several effectors, including NO, prostacyclin, hydrogen sulfide, and endothelium-derived hyperpolarizing factor, all of which cause vasorelaxation, and endothelin, which causes vasoconstriction. The release of these endothelial effectors of vascular smooth-muscle cell tone is stimulated by mechanical (shear stress, cyclic strain, etc.) and biochemical mediators (purinergic agonists, muscarinic agonists, peptidergic agonists), with the biochemical mediators acting through endothelial receptors specific to each class. In addition to these local paracrine modulators of vascular smooth-muscle cell tone, circulating mediators can affect tone, including norepinephrine and epinephrine, vasopressin, angiotensin II, bradykinin, and the natriuretic peptides (atrial natriuretic peptide [ANP], brain natriuretic peptide [BNP], C-type natriuretic peptide [CNP], and dendroaspis natriuretic peptide [DNP]), as discussed above.

VASCULAR REGENERATION

Growth of new blood vessels can occur in response to conditions such as chronic hypoxemia and tissue ischemia. Growth factors, including vascular endothelial growth factor (VEGF) and forms of fibroblast growth factor (FGF), activate a signaling cascade that stimulates endothelial proliferation and tube formation, defined as *angiogenesis*. Guidance molecules, including members of the semaphorin family of secreted peptides, direct blood vessel patterning by attracting or repelling nascent endothelial tubes. The development of collateral vascular networks in the ischemic myocardium, an example of angiogenesis, can result from selective activation of endothelial progenitor cells, which may reside in the blood vessel wall or home to the ischemic tissue from the bone marrow. True arteriogenesis, or the development of a new blood vessel that includes all three cell layers, normally does not occur in the cardiovascular system of adult mammals. The molecular mechanisms and progenitor cells that can recapitulate blood vessel development de novo are under rapidly advancing study (**Chaps. 88, 89e, and 90e**).

VASCULAR PHARMACOGENOMICS

The last decade has witnessed considerable progress in efforts to define the genetic differences underlying individual variations in vascular pharmacologic responses. Many investigators have focused on receptors and enzymes associated with neurohumoral modulation of vascular function as well as hepatic enzymes that metabolize drugs that affect vascular tone. The genetic polymorphisms thus far associated with differences in vascular response often (but not invariably) relate to functional differences in the activity or expression of the receptor or enzyme of interest. Some of these

polymorphisms appear to have different allele frequencies in specific ethnic groups.

CELLULAR BASIS OF CARDIAC CONTRACTION

CARDIAC ULTRASTRUCTURE

About three-fourths of the ventricular mass is composed of cardiomyocytes, normally 60–140 μm in length and 17–25 μm in diameter (Fig. 265e-4A). Each cell contains multiple, rodlike cross-banded strands (myofibrils) that run the length of the cell and are composed of serially repeating structures, the sarcomeres. The cytoplasm between the myofibrils contains other cell constituents, including the single

centrally located nucleus, numerous mitochondria, and the intracellular membrane system, the sarcoplasmic reticulum.

The *sarcomere*, the structural and functional unit of contraction, lies between adjacent Z lines, which are dark repeating bands that are apparent on transmission electron microscopy. The distance between Z lines varies with the degree of contraction or stretch of the muscle and ranges between 1.6 and 2.2 μm . Within the confines of the sarcomere are alternating light and dark bands, giving the myocardial fibers their striated appearance under the light microscope. At the center of the sarcomere is a dark band of constant length (1.5 μm), the A band, which is flanked by two lighter bands, the I bands, which are of variable length. The sarcomere of heart muscle, like that of skeletal muscle, consists of two sets of interdigitating myofibrils. Thicker filaments, composed principally of the protein myosin, traverse the A band; they are about 10 nm (100 \AA) in diameter, with tapered ends. Thinner filaments, composed primarily of actin, course from the Z lines through the I band into the A band; they are approximately 5 nm (50 \AA) in diameter and 1.0 μm in length. Thus, thick and thin filaments overlap only within the (dark) A band, whereas the (light) I band contains only thin filaments. On electron-microscopic examination, bridges may be seen to extend between the thick and thin filaments within the A band; these are myosin heads (see below) bound to actin filaments.

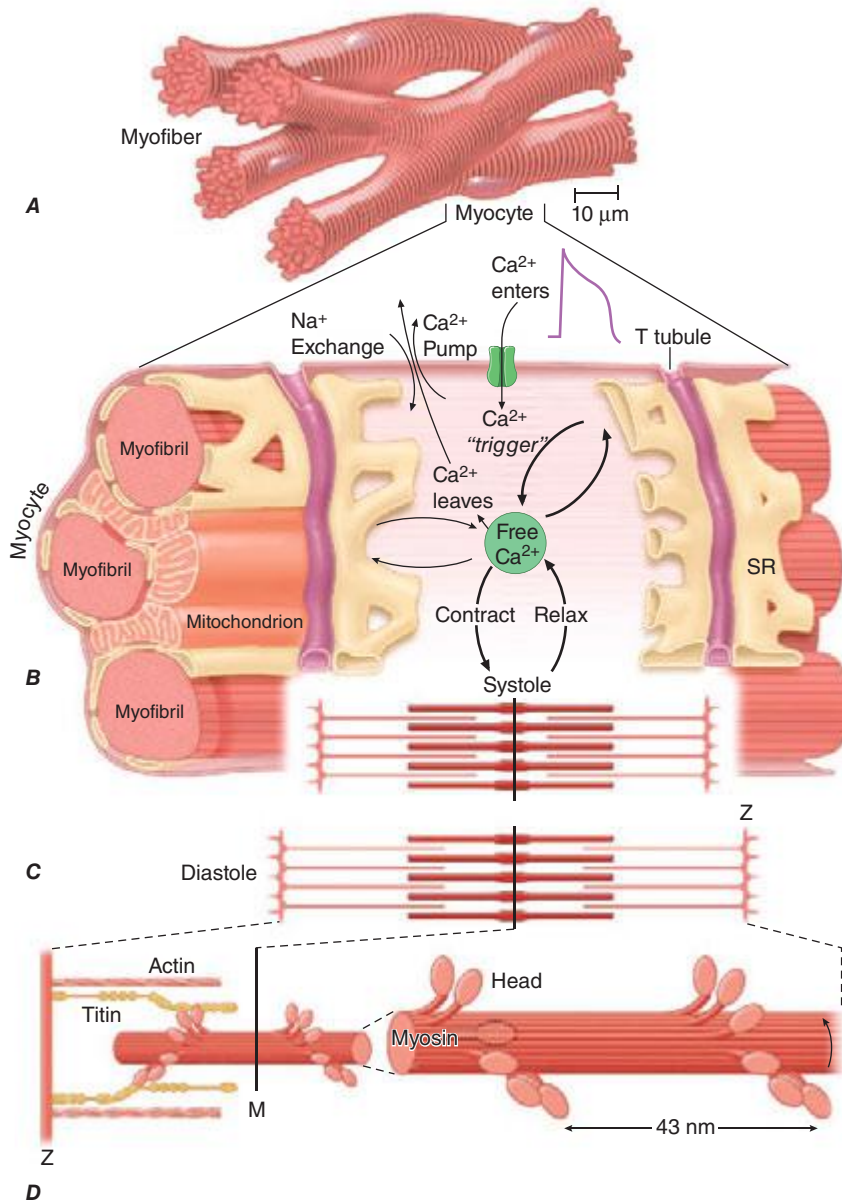


FIGURE 265e-4 A shows the branching myocytes making up the cardiac myofibers. B illustrates the critical role played by the changing $[\text{Ca}^{2+}]$ in the myocardial cytosol. Ca^{2+} ions are schematically shown as entering through the calcium channel that opens in response to the wave of depolarization that travels along the sarcolemma. These Ca^{2+} ions “trigger” the release of more calcium from the sarcoplasmic reticulum (SR) and thereby initiate a contraction-relaxation cycle. Eventually the small quantity of Ca^{2+} that has entered the cell leaves predominantly through an $\text{Na}^+/\text{Ca}^{2+}$ exchanger, with a lesser role for the sarcolemmal Ca^{2+} pump. The varying actin-myosin overlap is shown for (B) systole, when $[\text{Ca}^{2+}]$ is maximal, and (C) diastole, when $[\text{Ca}^{2+}]$ is minimal. D. The myosin heads, attached to the thick filaments, interact with the thin actin filaments. (From LH Opie: *Heart Physiology: From Cell to Circulation*, 4th ed. Philadelphia, Lippincott, Williams & Wilkins, 2004. Reprinted with permission. Copyright LH Opie, 2004.)

THE CONTRACTILE PROCESS

The sliding filament model for muscle contraction rests on the fundamental observation that both the thick and the thin filaments are constant in overall length during both contraction and relaxation. With activation, the actin filaments are propelled farther into the A band. In the process, the A band remains constant in length, whereas the I band shortens and the Z lines move toward one another.

The *myosin* molecule is a complex, asymmetric fibrous protein with a molecular mass of about 500,000 Da; it has a rodlike portion that is about 150 nm (1500 \AA) in length with a globular portion (head) at its end. These globular portions of myosin form the bridges between the myosin and actin molecules and are the site of ATPase activity. In forming the thick myofibril, which is composed of ~300 longitudinally stacked myosin molecules, the rodlike segments of the myosin molecules are laid down in an orderly, polarized manner, leaving the globular portions projecting outward so that they can interact with actin to generate force and shortening (Fig. 265e-4B).

Actin has a molecular mass of about 47,000 Da. The thin filament consists of a double helix of two chains of actin molecules wound about each other on a larger molecule, tropomyosin. A group of regulatory proteins—troponins C, I, and T—are spaced at regular intervals on this filament (Fig. 265e-5). In contrast to myosin, actin lacks intrinsic enzymatic activity but does combine reversibly with myosin in the presence of ATP and Ca^{2+} . The calcium ion activates the myosin ATPase, which in turn breaks down ATP, the energy source for contraction (Fig. 265e-5). The activity of myosin ATPase determines the rate of forming and breaking of the actomyosin cross-bridges and ultimately the velocity of muscle contraction. In relaxed muscle, tropomyosin inhibits this interaction. *Titin* (Fig. 265e-4D) is a large, flexible, myofibrillar protein that connects myosin to the Z line; its stretching contributes to the elasticity of the heart.

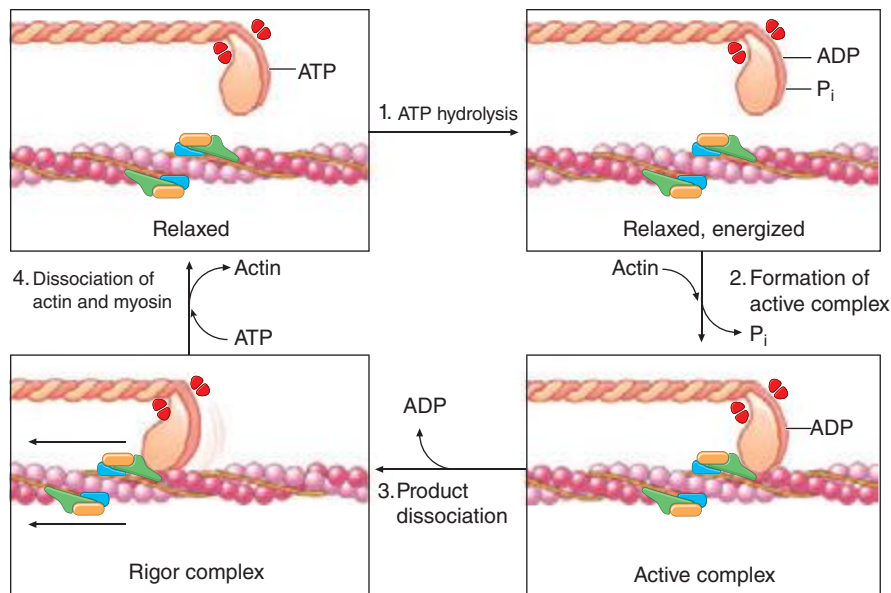


FIGURE 265e-5 Four steps in cardiac muscle contraction and relaxation. In relaxed muscle (*upper left*), ATP bound to the myosin cross-bridge dissociates the thick and thin filaments. **Step 1:** Hydrolysis of myosin-bound ATP by the ATPase site on the myosin head transfers the chemical energy of the nucleotide to the activated cross-bridge (*upper right*). When cytosolic Ca^{2+} concentration is low, as in relaxed muscle, the reaction cannot proceed because tropomyosin and the troponin complex on the thin filament do not allow the active sites on actin to interact with the cross-bridges. Therefore, even though the cross-bridges are energized, they cannot interact with actin. **Step 2:** When Ca^{2+} binding to troponin C has exposed active sites on the thin filament, actin interacts with the myosin cross-bridges to form an active complex (*lower right*) in which the energy derived from ATP is retained in the actin-bound cross-bridge, whose orientation has not yet shifted. **Step 3:** The muscle contracts when ADP dissociates from the cross-bridge. This step leads to the formation of the low-energy rigor complex (*lower left*) in which the chemical energy derived from ATP hydrolysis has been expended to perform mechanical work (the “rowing” motion of the cross-bridge). **Step 4:** The muscle returns to its resting state, and the cycle ends when a new molecule of ATP binds to the rigor complex and dissociates the cross-bridge from the thin filament. This cycle continues until calcium is dissociated from troponin C in the thin filament, which causes the contractile proteins to return to the resting state with the cross-bridge in the energized state. ADP, adenosine diphosphate; ATP, adenosine triphosphate; ATPase, adenosine triphosphatase. (From AM Katz: *Heart failure: Cardiac function and dysfunction*, in *Atlas of Heart Diseases*, 3rd ed, WS Colucci [ed]. Philadelphia, Current Medicine, 2002. Reprinted with permission.)

Dystrophin is a long cytoskeletal protein that has an amino-terminal actin-binding domain and a carboxy-terminal domain that binds to the dystroglycan complex at adherens junctions on the cell membrane, thus tethering the sarcomere to the cell membrane at regions tightly coupled to adjacent contracting myocytes. Mutations in components of the dystrophin complex lead to muscular dystrophy and associated cardiomyopathy.

During activation of the cardiac myocyte, Ca^{2+} becomes attached to one of three components of the heterotrimer troponin C, which results in a conformational change in the regulatory protein tropomyosin; the latter, in turn, exposes the actin cross-bridge interaction sites (Fig. 265e-5). Repetitive interaction between myosin heads and actin filaments is termed *cross-bridge cycling*, which results in sliding of the actin along the myosin filaments, ultimately causing muscle shortening and/or the development of tension. The splitting of ATP then dissociates the myosin cross-bridge from actin. In the presence of ATP (Fig. 265e-5), linkages between actin and myosin filaments are made and broken cyclically as long as sufficient Ca^{2+} is present; these linkages cease when $[Ca^{2+}]$ falls below a critical level, and the troponin-tropomyosin complex once more prevents interactions between the myosin cross-bridges and actin filaments (Fig. 265e-6).

Intracellular Ca^{2+} is a principal determinant of the inotropic state of the heart. Most agents that stimulate myocardial contractility (positive inotropic stimuli), including the digitalis glycosides and

β -adrenergic agonists, increase the $[Ca^{2+}]$ in the vicinity of the myofilaments, which in turn triggers cross-bridge cycling. Increased impulse traffic in the cardiac adrenergic nerves stimulates myocardial contractility as a consequence of the release of norepinephrine from cardiac adrenergic nerve endings. Norepinephrine activates myocardial β receptors and, through the G_s -stimulated guanine nucleotide-binding protein, activates the enzyme adenylyl cyclase, which leads to the formation of the intracellular second messenger cyclic AMP from ATP (Fig. 265e-6). Cyclic AMP in turn activates protein kinase A (PKA), which phosphorylates the Ca^{2+} channel in the myocardial sarcolemma, thereby enhancing the influx of Ca^{2+} into the myocyte. Other functions of PKA are discussed below.

The *sarcoplasmic reticulum* (SR) (Fig. 265e-7), a complex network of anastomosing intracellular channels, invests the myofibrils. Its longitudinally disposed tubules closely invest the surfaces of individual sarcomeres but have no direct continuity with the outside of the cell. However, closely related to the SR, both structurally and functionally, are the transverse tubules, or T system, formed by tubelike invaginations of the sarcolemma that extend into the myocardial fiber along the Z lines, i.e., the ends of the sarcomeres.

CARDIAC ACTIVATION

In the inactive state, the cardiac cell is electrically polarized; i.e., the interior has a negative charge relative to the outside of the cell, with a transmembrane potential of -80 to -100 mV (Chap. 273e). The sarcolemma, which in the resting state is largely impermeable to Na^+ , has a Na^+ - and K^+ -stimulating pump energized by ATP that extrudes Na^+ from the cell; this pump plays a critical role in establishing the resting potential. Thus, intracellular $[K^+]$ is relatively high and $[Na^+]$ is far lower; conversely, extracellular $[Na^+]$ is high and $[K^+]$ is low. At the same time, in the resting state, extracellular $[Ca^{2+}]$ greatly exceeds free intracellular $[Ca^{2+}]$.

The action potential has four phases (see Fig. 273e-1B). During the plateau of the action potential (phase 2), there is a slow inward current through L-type Ca^{2+} channels in the sarcolemma (Fig. 265e-7). The depolarizing current not only extends across the surface of the cell but penetrates deeply into the cell by way of the ramifying T tubular system. The absolute quantity of Ca^{2+} that crosses the sarcolemma and the T system is relatively small and by itself appears to be insufficient to bring about full activation of the contractile apparatus. However, this Ca^{2+} current triggers the release of much larger quantities of Ca^{2+} from the SR, a process termed *Ca^{2+} -induced Ca^{2+} release*. The latter is a major determinant of intracytoplasmic $[Ca^{2+}]$ and therefore of myocardial contractility.

Ca^{2+} is released from the SR through a Ca^{2+} release channel, a cardiac isoform of the ryanodine receptor (RyR2), which controls intracytoplasmic $[Ca^{2+}]$ and, as in vascular smooth-muscle cells, leads to the local changes in intracellular $[Ca^{2+}]$ called calcium sparks. A number of regulatory proteins, including *calstabin 2*, inhibit RyR2 and thereby the release of Ca^{2+} from the SR. PKA dissociates calstabin from the RyR2, enhancing Ca^{2+} release and thereby myocardial contractility. Excessive plasma catecholamine levels and cardiac sympathetic neuronal release of norepinephrine cause hyperphosphorylation of PKA, leading to calstabin 2–depleted RyR2. The latter depletes SR Ca^{2+} stores and thereby impairs cardiac contraction, leading to heart failure, and also triggers ventricular arrhythmias.

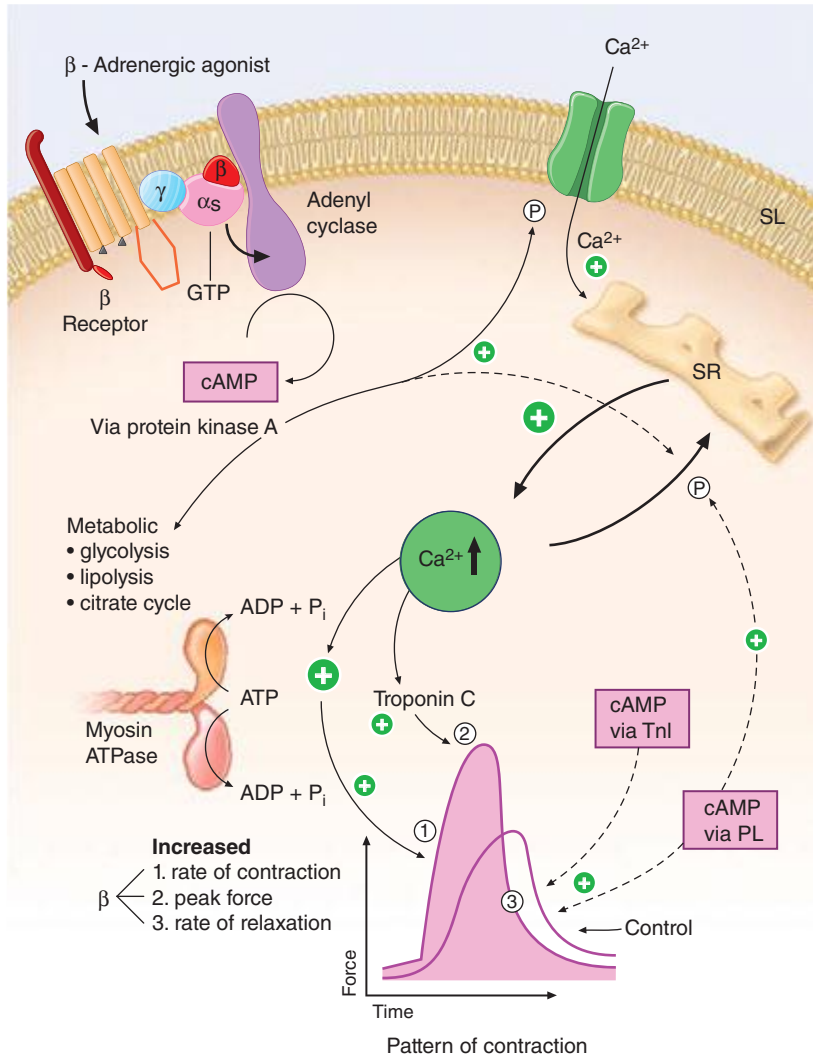


FIGURE 265e-6 Signal systems involved in positive inotropic and lusitropic (enhanced relaxation) effects of β -adrenergic stimulation. When the β -adrenergic agonist interacts with the β receptor, a series of G protein-mediated changes leads to activation of adenylyl cyclase and the formation of cyclic adenosine monophosphate (cAMP). The latter acts via protein kinase A to stimulate metabolism (left) and phosphorylate the Ca^{2+} channel protein (right). The result is an enhanced opening probability of the Ca^{2+} channel, thereby increasing the inward movement of Ca^{2+} ions through the sarcolemma (SL) of the T tubule. These Ca^{2+} ions release more calcium from the sarcoplasmic reticulum (SR) to increase cytosolic Ca^{2+} and activate troponin C. Ca^{2+} ions also increase the rate of breakdown of adenosine triphosphate (ATP) to adenosine diphosphate (ADP) and inorganic phosphate (P_i). Enhanced myosin ATPase activity explains the increased rate of contraction, with increased activation of troponin C explaining increased peak force development. An increased rate of relaxation results from the ability of cAMP to activate as well the protein phospholamban, situated on the membrane of the SR, that controls the rate of uptake of calcium into the SR. The latter effect explains enhanced relaxation (lusitropic effect). P, phosphorylation; PL, phospholamban; Tnl, troponin I. (Modified from LH Opie: *Heart Physiology: From Cell to Circulation*, 4th ed. Philadelphia, Lippincott, Williams & Wilkins, 2004. Reprinted with permission. Copyright LH Opie, 2004.)

The Ca^{2+} released from the SR then diffuses toward the myofibrils, where, as already described, it combines with troponin C (Fig. 265e-6). By repressing this inhibitor of contraction, Ca^{2+} activates the myofilaments to shorten. During repolarization, the activity of the Ca^{2+} pump in the SR, the SR Ca^{2+} ATPase (SERCA_{2A}), reaccumulates Ca^{2+} against a concentration gradient, and the Ca^{2+} is stored in the SR by its attachment to a protein, *calsequestrin*. This reaccumulation of Ca^{2+} is an energy (ATP)-requiring process that lowers the cytoplasmic $[Ca^{2+}]$ to a level that inhibits the actomyosin interaction responsible for contraction, and in this manner leads to myocardial relaxation. Also,

there is an exchange of Ca^{2+} for Na^{+} at the sarcolemma (Fig. 265e-7), reducing the cytoplasmic $[Ca^{2+}]$. Cyclic AMP-dependent PKA phosphorylates the SR protein *phospholamban*; the latter, in turn, permits activation of the Ca^{2+} pump, thereby increasing the uptake of Ca^{2+} by the SR, accelerating the rate of relaxation, and providing larger quantities of Ca^{2+} in the SR for release by subsequent depolarization, thereby stimulating contraction.

Thus, the combination of the cell membrane, transverse tubules, and SR, with their ability to transmit the action potential and release and then reaccumulate Ca^{2+} , plays a fundamental role in the rhythmic contraction and relaxation of heart muscle. Genetic or pharmacologic alterations of any component, whatever its etiology, can disturb these functions.

CONTROL OF CARDIAC PERFORMANCE AND OUTPUT

The extent of shortening of heart muscle and, therefore, the stroke volume of the ventricle in the intact heart depend on three major influences: (1) the length of the muscle at the onset of contraction, i.e., the preload; (2) the tension that the muscle is called on to develop during contraction, i.e., the afterload; and (3) the contractility of the muscle, i.e., the extent and velocity of shortening at any given preload and afterload. The major determinants of preload, afterload, and contractility are shown in Table 265e-2.

THE ROLE OF MUSCLE LENGTH (PRELOAD)

The preload determines the length of the sarcomeres at the onset of contraction. The length of the sarcomeres associated with the most forceful contraction is $\sim 2.2 \mu m$. This length provides the optimum configuration for the interaction between the two sets of myofilaments. The length of the sarcomere also regulates the extent of activation of the contractile system, i.e., its sensitivity to Ca^{2+} . According to this concept, termed *length-dependent activation*, myofilament sensitivity to Ca^{2+} is also maximal at the optimal sarcomere length. The relation between the initial length of the muscle fibers and the developed force has prime importance for the function of heart muscle. This relationship forms the basis of Starling's law of the heart, which states that within limits, the force of ventricular contraction depends on the end-diastolic length of the cardiac muscle; in the intact heart, the latter relates closely to the ventricular end-diastolic volume.

CARDIAC PERFORMANCE

The ventricular end-diastolic or "filling" pressure sometimes is used as a surrogate for the end-diastolic volume. In isolated heart and heart-lung preparations, the stroke volume varies directly with the end-diastolic fiber length (preload) and inversely with the arterial resistance (afterload), and as the heart fails—i.e., as its contractility declines—it delivers a progressively smaller stroke volume from a normal or even elevated end-diastolic volume. The relation between the ventricular end-diastolic pressure and the stroke work of the ventricle (the ventricular function curve) provides a useful definition of the level of contractility of the heart in the intact organism. An increase in contractility is accompanied by a shift of the ventricular function curve upward and to the left (greater stroke work at any level of ventricular end-diastolic pressure, or lower end-diastolic volume at any level of stroke work), whereas a shift downward and to the right characterizes depression of contractility (Fig. 265e-8).

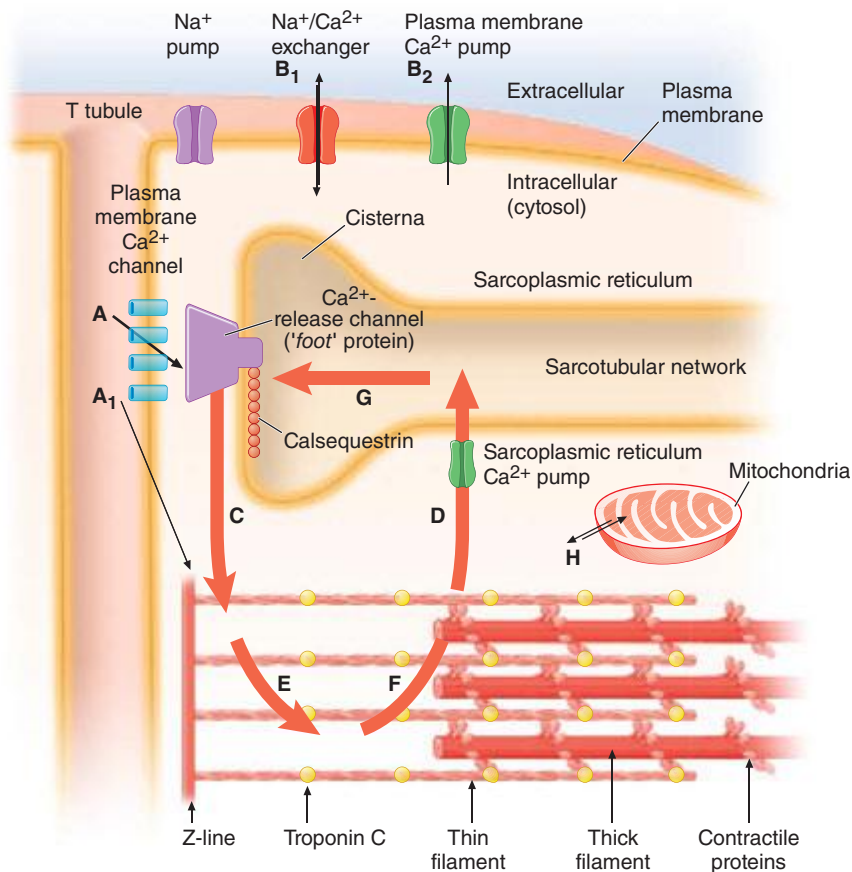


FIGURE 265e-7 The Ca²⁺ fluxes and key structures involved in cardiac excitation-contraction coupling. The arrows denote the direction of Ca²⁺ fluxes. The thickness of each arrow indicates the magnitude of the calcium flux. Two Ca²⁺ cycles regulate excitation-contraction coupling and relaxation. The larger cycle is entirely intracellular and involves Ca²⁺ fluxes into and out of the sarcoplasmic reticulum, as well as Ca²⁺ binding to and release from troponin C. The smaller extracellular Ca²⁺ cycle occurs when this cation moves into and out of the cell. The action potential opens plasma membrane Ca²⁺ channels to allow passive entry of Ca²⁺ into the cell from the extracellular fluid (arrow A). Only a small portion of the Ca²⁺ that enters the cell directly activates the contractile proteins (arrow A₁). The extracellular cycle is completed when Ca²⁺ is actively transported back out to the extracellular fluid by way of two plasma membrane fluxes mediated by the sodium-calcium exchanger (arrow B₁) and the plasma membrane calcium pump (arrow B₂). In the intracellular Ca²⁺ cycle, passive Ca²⁺ release occurs through channels in the cisternae (arrow C) and initiates contraction; active Ca²⁺ uptake by the Ca²⁺ pump of the sarcotubular network (arrow D) relaxes the heart. Diffusion of Ca²⁺ within the sarcoplasmic reticulum (arrow G) returns this activator cation to the cisternae, where it is stored in a complex with calsequestrin and other calcium-binding proteins. Ca²⁺ released from the sarcoplasmic reticulum initiates systole when it binds to troponin C (arrow E). Lowering of cytosolic [Ca²⁺] by the sarcoplasmic reticulum (SR) causes this ion to dissociate from troponin (arrow F) and relaxes the heart. Ca²⁺ also may move between mitochondria and cytoplasm (H). (Adapted from AM Katz: *Physiology of the Heart*, 4th ed. Philadelphia, Lippincott, Williams & Wilkins, 2005, with permission.)

VENTRICULAR AFTERLOAD

In the intact heart, as in isolated cardiac muscle, the extent and velocity of shortening of ventricular muscle fibers at any level of preload and of myocardial contractility relate inversely to the afterload, i.e., the load that opposes shortening. In the intact heart, the afterload may be defined as the tension developed in the ventricular wall during ejection. Afterload is determined by the aortic pressure as well as by the volume and thickness of the ventricular cavity. Laplace's law states that the tension of the myocardial fiber is the product of the intracavitary ventricular pressure and ventricular radius divided by wall thickness. Therefore, at any particular level of aortic pressure, the afterload on a dilated left ventricle exceeds that on a normal-sized

ventricle. Conversely, at the same aortic pressure and ventricular diastolic volume, the afterload on a hypertrophied ventricle is lower than of a normal chamber. The aortic pressure in turn depends on the peripheral vascular resistance, the physical characteristics of the arterial tree, and the volume of blood it contains at the onset of ejection.

Ventricular afterload critically regulates cardiovascular performance (Fig. 265e-9). As already noted, elevations in both preload and contractility increase myocardial fiber shortening, whereas increases in afterload reduce it. The extent of myocardial fiber shortening and left ventricular size determine stroke volume. An increase in arterial pressure induced by vasoconstriction, for example, augments afterload, which opposes myocardial fiber shortening, reducing stroke volume.

When myocardial contractility becomes impaired and the ventricle dilates, afterload rises (Laplace's law) and limits cardiac output. Increased afterload also may result from neural and humoral stimuli that occur in response to a fall in cardiac output. This increased afterload may reduce cardiac output further, thereby increasing ventricular volume and initiating a vicious circle, especially in patients with ischemic heart disease and limited myocardial O₂ supply. Treatment with vasodilators has the opposite effect; when afterload is reduced, cardiac output rises (Chap. 279).

Under normal circumstances, the various influences acting on cardiac performance enumerated above interact in a complex fashion to maintain cardiac output at a level appropriate to the requirements of the metabolizing tissues (Fig. 265e-9); interference with a single mechanism may not influence the cardiac output. For example, a moderate reduction of blood volume or the loss of the atrial contribution to ventricular contraction ordinarily can be sustained without a reduction in the cardiac output at rest. Under these circumstances, other factors, such as increases in the frequency of adrenergic nerve impulses to the heart, heart rate, and venous tone, will serve as compensatory mechanisms and sustain cardiac output in a normal individual.

EXERCISE

The integrated response to exercise illustrates the interactions among the three determinants of stroke volume: preload, afterload, and contractility (Fig. 265e-8). Hyperventilation, the pumping action of the exercising muscles, and venoconstriction during exercise all augment venous return and hence ventricular filling and preload (Table 265e-2). Simultaneously, the increase in the adrenergic nerve impulse traffic to the myocardium, the increased concentration of circulating catecholamines, and the tachycardia that

occur during exercise combine to augment the contractility of the myocardium (Fig. 265e-8, curves 1 and 2) and together elevate stroke volume and stroke work, without a change in or even a reduction of end-diastolic pressure and volume (Fig. 265e-8, points A and B). Vasodilation occurs in the exercising muscles, thus tending to limit the increase in arterial pressure that otherwise would occur as cardiac output rises to levels as high as five times greater than basal levels during maximal exercise. This vasodilation ultimately allows the achievement of a greatly elevated cardiac output during exercise at an arterial pressure only moderately higher than in the resting state.

TABLE 265e-2 DETERMINANTS OF STROKE VOLUME**I. Ventricular Preload**

- A. Blood volume
- B. Distribution of blood volume
 1. Body position
 2. Intrathoracic pressure
 3. Intrapericardial pressure
 4. Venous tone
 5. Pumping action of skeletal muscles
- C. Atrial contraction

II. Ventricular Afterload

- A. Systemic vascular resistance
- B. Elasticity of arterial tree
- C. Arterial blood volume
- D. Ventricular wall tension
 1. Ventricular radius
 2. Ventricular wall thickness

III. Myocardial Contractility^a

- A. Intramyocardial $[Ca^{2+}] \uparrow \downarrow$
- B. Cardiac adrenergic nerve activity $\uparrow \downarrow^b$
- C. Circulating catecholamines $\uparrow \downarrow^b$
- D. Cardiac rate $\uparrow \downarrow^b$
- E. Exogenous inotropic agents \uparrow
- F. Myocardial ischemia \downarrow
- G. Myocardial cell death (necrosis, apoptosis, autophagy) \downarrow
- H. Alterations of sarcomeric and cytoskeletal proteins \downarrow
 1. Genetic
 2. Hemodynamic overload
- I. Myocardial fibrosis \downarrow
- J. Chronic overexpression of neurohormones \downarrow
- K. Ventricular remodeling \downarrow
- L. Chronic and/or excessive myocardial hypertrophy \downarrow

^aArrows indicate directional effects of determinants of contractility. ^bContractility rises initially but later becomes depressed.

ASSESSMENT OF CARDIAC FUNCTION

Several techniques can define impaired cardiac function in clinical practice. The cardiac output and stroke volume may be depressed in the presence of heart failure, but not uncommonly, these variables are within normal limits in this condition. A somewhat more sensitive index of cardiac function is the ejection fraction, i.e., the ratio of stroke volume to end-diastolic volume (normal value = $67 \pm 8\%$), which is frequently depressed in systolic heart failure even when the stroke volume itself is normal. Alternatively, abnormally elevated ventricular end-diastolic volume (normal value = $75 \pm 20 \text{ mL/m}^2$) or end-systolic volume (normal value = $25 \pm 7 \text{ mL/m}^2$) signifies impairment of left ventricular systolic function.

Noninvasive techniques, particularly echocardiography as well as radionuclide scintigraphy and cardiac magnetic resonance imaging (MRI) (Chap. 270e), have great value in the clinical assessment of myocardial function. They provide measurements of end-diastolic and end-systolic volumes, ejection fraction, and systolic shortening rate, and they allow assessment of ventricular filling (see below) as well as regional contraction and relaxation. The latter measurements are particularly important in ischemic heart disease, as myocardial infarction causes regional myocardial damage.

A limitation of measurements of cardiac output, ejection fraction, and ventricular volumes in assessing cardiac function is that ventricular loading conditions strongly influence these variables. Thus, a depressed ejection fraction and lowered cardiac output may occur in patients with normal ventricular function but reduced preload, as occurs in hypovolemia, or with increased afterload, as occurs in acutely elevated arterial pressure.

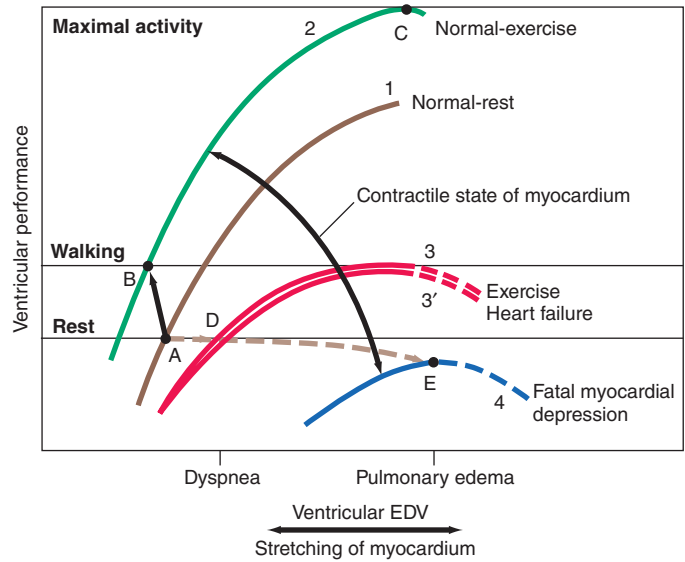


FIGURE 265e-8 The interrelations among influences on ventricular end-diastolic volume (EDV) through stretching of the myocardium and the contractile state of the myocardium. Levels of ventricular EDV associated with filling pressures that result in dyspnea and pulmonary edema are shown on the abscissa. Levels of ventricular performance required when the subject is at rest, while walking, and during maximal activity are designated on the ordinate. The broken lines are the descending limbs of the ventricular-performance curves, which are rarely seen during life but show the level of ventricular performance if end-diastolic volume could be elevated to very high levels. For further explanation, see text. (Modified from WS Colucci and E Braunwald: *Pathophysiology of heart failure, in Braunwald's Heart Disease, 7th ed, DP Zipes et al [eds]. Philadelphia: Elsevier, 2005, pp 509–538.*)

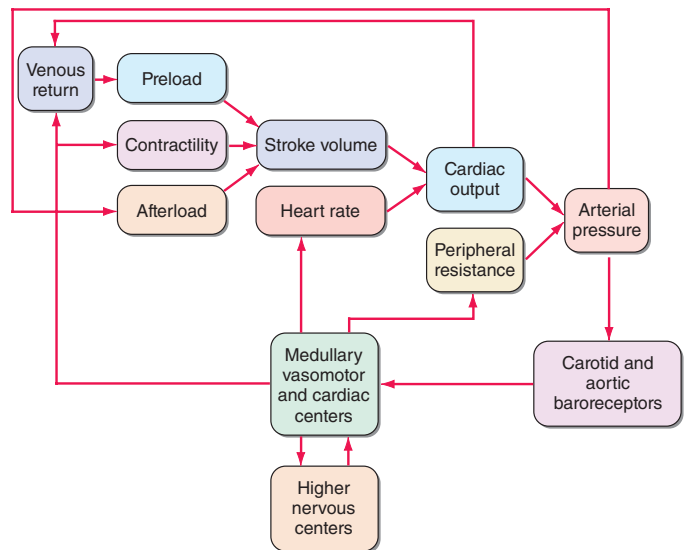


FIGURE 265e-9 Interactions in the intact circulation of preload, contractility, and afterload in producing stroke volume. Stroke volume combined with heart rate determines cardiac output, which, when combined with peripheral vascular resistance, determines arterial pressure for tissue perfusion. The characteristics of the arterial system also contribute to afterload, an increase that reduces stroke volume. The interaction of these components with carotid and aortic arch baroreceptors provides a feedback mechanism to higher medullary and vasomotor cardiac centers and to higher levels in the central nervous system to effect a modulating influence on heart rate, peripheral vascular resistance, venous return, and contractility. (From MR Starling: *Physiology of myocardial contraction, in Atlas of Heart Failure: Cardiac Function and Dysfunction, 3rd ed, WS Colucci and E Braunwald [eds]. Philadelphia: Current Medicine, 2002, pp 19–35.*)

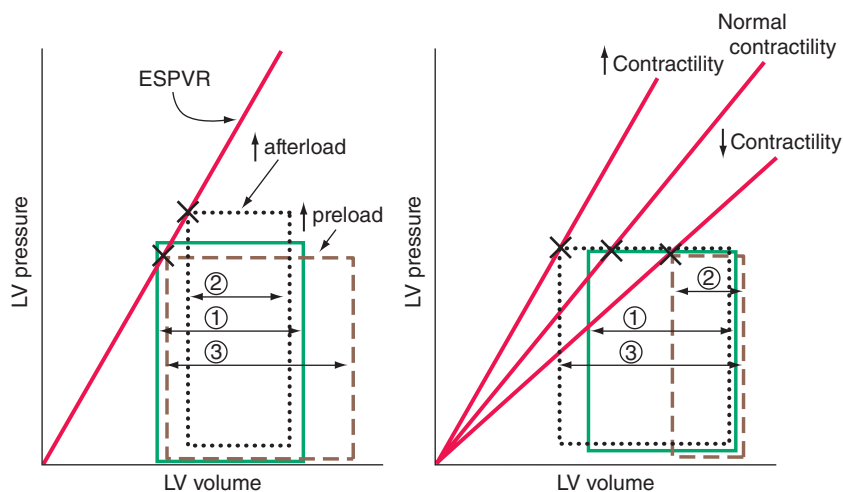


FIGURE 265e-10 The responses of the left ventricle to increased afterload, increased preload, and increased and reduced contractility are shown in the pressure-volume plane. **Left.** Effects of increases in preload and afterload on the pressure-volume loop. Because there has been no change in contractility, the end-systolic pressure-volume relationship (ESPVR) is unchanged. With an increase in afterload, stroke volume falls (1 → 2); with an increase in preload, stroke volume rises (1 → 3). **Right.** With increased myocardial contractility and constant left ventricular end-diastolic volume, the ESPVR moves to the left of the normal line (lower end-systolic volume at any end-systolic pressure) and stroke volume rises (1 → 3). With reduced myocardial contractility, the ESPVR moves to the right; end-systolic volume is increased, and stroke volume falls (1 → 2).

The end-systolic left ventricular pressure-volume relationship is a particularly useful index of ventricular performance because it does not depend on preload and afterload (Fig. 265e-10). At any level of myocardial contractility, left ventricular end-systolic volume varies inversely with end-systolic pressure; as contractility declines, end-systolic volume (at any level of end-systolic pressure) rises.

DIASTOLIC FUNCTION

Ventricular filling is influenced by the extent and speed of myocardial relaxation, which in turn depends on the rate of uptake of Ca^{2+} by the SR; the latter may be enhanced by adrenergic activation and reduced by ischemia, which reduces the ATP available for pumping Ca^{2+} into the SR (see above). The stiffness of the ventricular wall also may impede filling. Ventricular stiffness increases with hypertrophy and conditions that infiltrate the ventricle, such as amyloid, or is caused by an extrinsic constraint (e.g., pericardial compression) (Fig. 265e-11).

Ventricular filling can be assessed by continuously measuring the velocity of flow across the mitral valve using Doppler ultrasound. Normally, the velocity of inflow is more rapid in early diastole than during atrial systole; with mild to moderately impaired relaxation, the rate of early diastolic filling declines, whereas the rate of presystolic filling rises. With further impairment of filling, the pattern is “pseudo-normalized,” and early ventricular filling becomes more rapid as left atrial pressure upstream to the stiff left ventricle rises.

CARDIAC METABOLISM

The heart requires a continuous supply of energy (in the form of ATP) not only to perform its mechanical pumping functions, but also to regulate intracellular and transsarcolemmal ionic movements and concentration gradients. Among its pumping functions, the development of tension, the frequency of contraction, and the level of myocardial contractility are the principal determinants of the heart's substantial energy needs, making its O_2 requirements approximately 15% of that of the entire organism.

Most ATP production depends on the oxidation of substrate (glucose and free fatty acids [FFAs]). Myocardial FFAs are derived from circulating FFAs, which result principally from lipolysis in adipose tissue, whereas the myocyte's glucose derives from plasma as well

as from the cell's breakdown of its glycogen stores (glycogenolysis). These two principal sources of acetyl coenzyme A in cardiac muscle vary reciprocally. Glucose is broken down in the cytoplasm into a three-carbon product, pyruvate, which passes into the mitochondria, where it is metabolized to the two-carbon fragment, acetyl-CoA, and undergoes oxidation. FFAs are converted to acyl-CoA in the cytoplasm and acetyl-CoA in the mitochondria. Acetyl-CoA enters the citric acid (Krebs) cycle to produce ATP by oxidative phosphorylation within the mitochondria; ATP then enters the cytoplasm from the mitochondrial compartment. Intracellular ADP, resulting from the breakdown of ATP, enhances mitochondrial ATP production.

In the fasted, resting state, circulating FFA concentrations and their myocardial uptake are high, and they furnish most of the heart's acetyl-CoA (~70%). In the fed state, with elevations of blood glucose and insulin, glucose oxidation increases and FFA oxidation subsides. Increased cardiac work, the administration of inotropic agents, hypoxia, and mild ischemia all enhance myocardial glucose uptake, glucose production resulting from glycogenolysis, and glucose metabolism to pyruvate (glycolysis). By contrast, β -adrenergic stimulation, as occurs during stress, raises the circulating levels and metabolism of FFAs in favor of glucose. Severe ischemia inhibits the cytoplasmic enzyme pyruvate dehydrogenase, and despite both glycogen and glucose breakdown, glucose is metabolized only to lactic acid (anaerobic glycolysis), which does not enter the citric acid cycle. Anaerobic glycolysis produces much less ATP than does aerobic glucose metabolism, in which glucose is metabolized to pyruvate and subsequently oxidized to CO_2 . High concentrations of circulating FFAs, which can occur when adrenergic stimulation is superimposed on severe ischemia, reduce oxidative phosphorylation and also cause ATP wastage; the myocardial content of ATP declines and impairs myocardial contraction. In addition, products of FFA breakdown can exert toxic effects on cardiac cell membranes and may be arrhythmogenic.

as from the cell's breakdown of its glycogen stores (glycogenolysis). These two principal sources of acetyl coenzyme A in cardiac muscle vary reciprocally. Glucose is broken down in the cytoplasm into a three-carbon product, pyruvate, which passes into the mitochondria, where it is metabolized to the two-carbon fragment, acetyl-CoA, and undergoes oxidation. FFAs are converted to acyl-CoA in the cytoplasm and acetyl-CoA in the mitochondria. Acetyl-CoA enters the citric acid (Krebs) cycle to produce ATP by oxidative phosphorylation within the mitochondria; ATP then enters the cytoplasm from the mitochondrial compartment. Intracellular ADP, resulting from the breakdown of ATP, enhances mitochondrial ATP production.

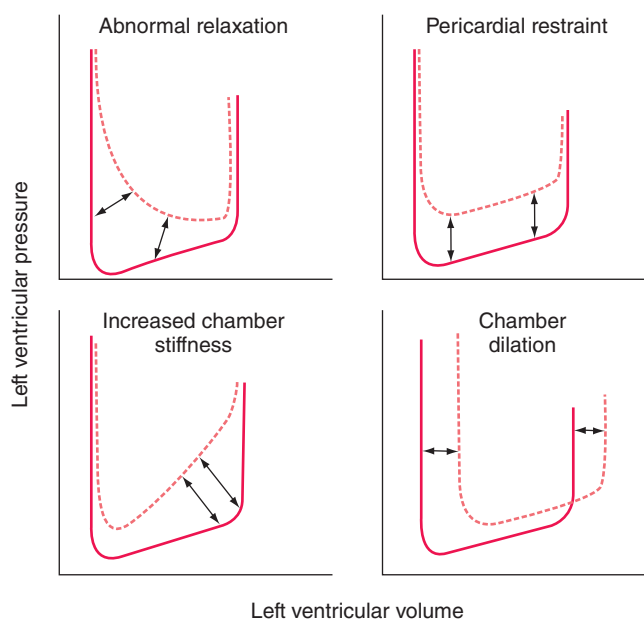


FIGURE 265e-11 Mechanisms that cause diastolic dysfunction reflected in the pressure-volume relation. The bottom half of the pressure-volume loop is depicted. *Solid lines* represent normal subjects; *broken lines* represent patients with diastolic dysfunction. (From JD Carroll et al: *The differential effects of positive inotropic and vasodilator therapy on diastolic properties in patients with congestive cardiomyopathy*. *Circulation* 74:815, 1986; with permission.)

Myocardial energy is stored as creatine phosphate (CP), which is in equilibrium with ATP, the immediate source of energy. In states of reduced energy availability, the CP stores decline first. Cardiac hypertrophy, fibrosis, tachycardia, increased wall tension resulting from ventricular dilation, and increased intracytoplasmic $[Ca^{2+}]$ all contribute to increased myocardial energy needs. When coupled with reduced coronary flow reserve, as occurs with obstruction of coronary arteries or abnormalities of the coronary microcirculation, an imbalance in myocardial ATP production relative to demand may occur, and the resulting ischemia can worsen or cause heart failure.

Developmental Biology of the Cardiovascular System The heart is the first organ to form during embryogenesis (Fig. 265e-12) and must accomplish the simultaneous challenges of circulating blood, nutrients, and oxygen to the other forming organs while continuing to grow and undergo complex morphogenetic changes. Early progenitors of the heart arise within very early crescent-shaped fields of lateral splanchnic mesoderm under the influence of multiple signals, including those derived from neural ectoderm long before neural tube closure. Early cardiac precursors express genes encoding regulatory transcription factors that play reiterated roles in cardiac

development, such as *NKX2-5* and *GATA4*. Mutations in these genes are responsible for some forms of inherited congenital heart disease. Cardiac precursors coalesce to form a midline heart tube composed of a single cell layer of endocardium surrounded by a single layer of myocardial precursors. The caudal, inflow region of the heart tube, which is destined to adopt a more rostral final position, represents the atrial anlagen, whereas the rostral, outflow portion of the tube forms the truncus arteriosus, which divides to produce the aorta and the proximal pulmonary artery. Between these extremes lie the structural precursors of the ventricles.

The linear heart tube undergoes an asymmetric looping process (the first gross evidence of left-right asymmetry in the developing embryo), which positions the portion of the heart tube destined to become the left ventricle to the left of the more rostral precursors of the right ventricle and outflow tract. Looping is coordinated with chamber specification and ballooning of various regions of the heart tube to produce the presumptive atria and ventricles.

Relatively recent work has demonstrated that significant portions of the right ventricle are formed by cells that are added to the developing heart after looping has occurred. These cells, which are derived from what is called the second heart field, migrate to the heart from the ventral pharynx and express markers that allow for their identification, including *Islet-1*. Different embryologic origins of cells within the right and left ventricles may help explain why some forms of congenital and adult heart diseases affect these regions of the heart to varying degrees.

After looping and chamber formation, a series of septation events divide the left and right sides of the heart, separate the atria from the ventricles, and form the aorta and pulmonary artery from the truncus arteriosus. Cardiac valves form between the atria and the ventricles and between the ventricles and the outflow vessels. Early in development, the single layer of myocardial cells secretes an extracellular matrix rich in hyaluronic acid. This extracellular matrix, termed “cardiac jelly,” accumulates within the endocardial cushions, precursors of the cardiac valves. Signals from overlying myocardial cells, including members of the transforming growth factor β family, trigger migration, invasion, and phenotypic changes of underlying endocardial cells, which undergo an epithelial-mesenchymal transformation and invade the cardiac jelly to cellularize the endocardial cushions. Mesenchymal components proliferate and remodel to form the mature valve leaflets.

The great vessels form as a series of bilaterally symmetric aortic arch arteries that undergo asymmetric remodeling events to form the mature vasculature. The immigration of neural crest cells that arise in the dorsal neural tube orchestrates this process. These cells are required for aortic arch remodeling and septation of the truncus arteriosus. They develop into smooth-muscle cells within the tunica media of the aortic arch, the ductus arteriosus, and the carotid arteries. Smooth-muscle cells within the descending aorta arise from a different embryologic source, the lateral plate

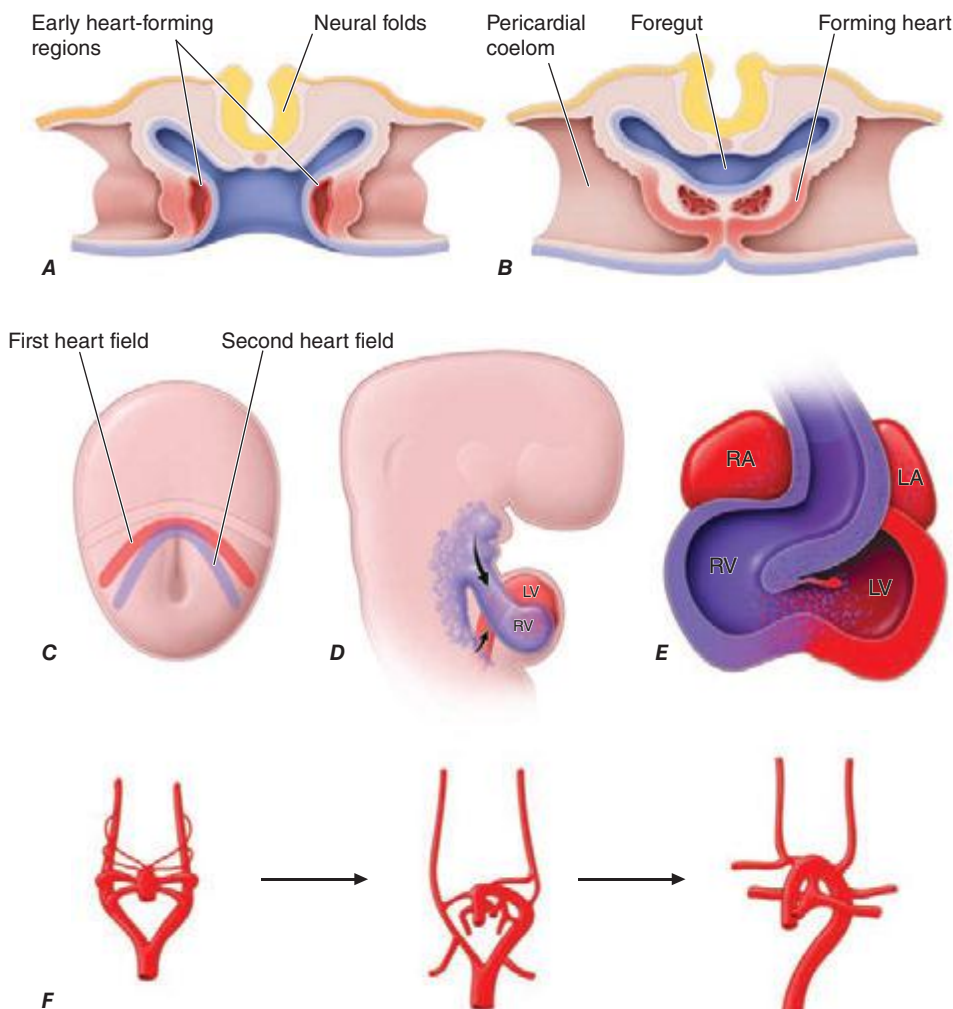


FIGURE 265e-12 **A.** Schematic depiction of a transverse section through an early embryo depicts the bilateral regions where early heart tubes form. **B.** The bilateral heart tubes subsequently migrate to the midline and fuse to form the linear heart tube. **C.** At the early cardiac crescent stage of embryonic development, cardiac precursors include a primary heart field fated to form the linear heart tube and a second heart field fated to add myocardium to the inflow and outflow poles of the heart. **D.** Second heart field cells populate the pharyngeal region before subsequently migrating to the maturing heart. **E.** Large portions of the right ventricle and outflow tract and some cells within the atria derive from the second heart field. **F.** The aortic arch arteries form as symmetric sets of vessels that then remodel under the influence of the neural crest to form the asymmetric mature vasculature. LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle.

265e-12 mesoderm, and smooth muscle of the proximal outflow tract arises from the second heart field. Neural crest cells are sensitive to both vitamin A and folic acid, and congenital heart disease involving abnormal remodeling of the aortic arch arteries has been associated with maternal deficiencies of these vitamins. Congenital heart disease involving the outflow tract can be associated with other defects of neural crest, such as cleft palate or craniofacial abnormalities.

Coronary artery formation requires yet another cell population that initiates extrinsic to the embryonic heart fields. Epicardial cells arise in the proepicardial organ, a derivative of the septum transversum, which also contributes to the fibrous portion of the diaphragm and to the liver. Proepicardial cells contribute to the smooth-muscle cells of the coronary arteries and are required for their proper patterning. Other cell types within the heart, including fibroblasts and potentially some myocardial and endocardial cells, also can arise from the proepicardium.

The cardiac conduction system, which functions both to generate and to propagate electrical impulses, develops primarily from multipotential cardiac precursors, which also give rise to cardiac muscle. The conduction system is composed of slow-conducting (proximal) components, such as the sinoatrial (SA) and atrioventricular (AV) nodes, as well as fast-conducting (distal) components, including the His bundle, bundle branches, and Purkinje fibers. The AV node primarily serves to delay the electrical impulse between atria and ventricles (manifesting decremental conduction), whereas the distal conduction system rapidly delivers the impulse throughout the ventricles. Significant recent attention has been focused on the embryologic origins of various components of the specialized conduction

network. Precursors within the sinus venosus give rise to the SA node, whereas those within the AV canal mature into heterogeneous cell types that compose the AV node. Myocardial cells transdifferentiate into Purkinje fibers to form the distal conduction system. Fast and slow conducting cell types within the nodes and bundles are characterized by expression of distinct gap junction proteins, including connexins, and ion channels that characterize unique cell fates and electrical properties of the tissues. Developmental defects in conduction system morphogenesis and lineage determination can lead to various electrophysiologic disorders, including congenital heart block and preexcitation syndromes such as the Wolff-Parkinson-White syndrome ([Chap. 276](#)).

Studies of cardiac stem and progenitor cells suggest that progressive lineage restriction results in the gradual and stepwise determination of mature cell fates within the heart, with early precursors capable of adopting endothelial, smooth-muscle, or cardiac phenotypes, and subsequent further specialization into atrial, ventricular, and specialized conduction cell types.

REGENERATING CARDIAC TISSUE

Until very recently, adult mammalian myocardial cells were viewed as fully differentiated and without regenerative potential. Evidence currently supports the existence of limited regenerative potential of the mature heart. Considerable current effort is being devoted to evaluating the utility of various putative stem cell populations and regenerative approaches to enhance cardiac repair after injury. The success of such approaches would offer the exciting possibility of reconstructing an infarcted or failing ventricle ([Chaps. 88 and 90e](#)).

266e Epidemiology of Cardiovascular Disease

Thomas A. Gaziano, J. Michael Gaziano

Cardiovascular disease (CVD) is now the most common cause of death worldwide. Before 1900, infectious diseases and malnutrition were the most common causes, and CVD was responsible for less than 10% of all deaths. In 2010, CVD accounted for approximately 16 million deaths worldwide (30%), including nearly 40% of deaths in high-income countries and about 28% in low- and middle-income countries.

THE EPIDEMIOLOGIC TRANSITION



The global rise in CVD is the result of an unprecedented transformation in the causes of morbidity and mortality during the twentieth century. Known as the epidemiologic transition, this shift is driven by industrialization, urbanization, and associated lifestyle changes and is taking place in every part of the world among all races, ethnic groups, and cultures. The transition is divided into four basic stages: pestilence and famine, receding pandemics, degenerative and man-made diseases, and delayed degenerative diseases. A fifth stage, characterized by an epidemic of inactivity and obesity, is emerging in some countries (Table 266e-1).

The *age of pestilence and famine* is marked by malnutrition, infectious diseases, and high infant and child mortality that are offset by high fertility. Tuberculosis, dysentery, cholera, and influenza are often fatal, resulting in a mean life expectancy of about 30 years. CVD, which accounts for less than 10% of deaths, takes the form of rheumatic heart disease and cardiomyopathies due to infection and malnutrition. Approximately 10% of the world's population remains in the age of pestilence and famine.

Per capita income and life expectancy increase during the *age of receding pandemics* as the emergence of public health systems, cleaner water supplies, and improved nutrition combine to drive down deaths from infectious disease and malnutrition. Infant and childhood mortality also decline, but deaths due to CVD increase to between 10 and 35% of all deaths. Rheumatic valvular disease, hypertension, coronary heart disease (CHD), and stroke are the predominant forms of CVD. Almost 40% of the world's population is currently in this stage.

The *age of degenerative and man-made diseases* is distinguished by mortality from noncommunicable diseases—primarily CVD—surpassing mortality from malnutrition and infectious diseases. Caloric intake, particularly from animal fat, increases. CHD and stroke are prevalent, and between 35 and 65% of all deaths can be traced to CVD. Typically, the rate of CHD deaths exceeds that of stroke by a ratio of 2:1 to 3:1. During this period, average life expectancy surpasses the age of 50. Roughly 35% of the world's population falls into this category.

In the *age of delayed degenerative diseases*, CVD and cancer remain the major causes of morbidity and mortality, with CVD accounting for 40% of all deaths. However, age-adjusted CVD mortality declines, aided by preventive strategies (for example, smoking cessation programs and effective blood pressure control), acute hospital management, and technologic advances, such as the availability of bypass surgery. CHD, stroke, and congestive heart failure are the primary forms of CVD. About 15% of the world's population is now in the age of delayed degenerative diseases or is exiting this age and moving into the fifth stage of the epidemiologic transition.

In the industrialized world, physical activity continues to decline while total caloric intake increases. The resulting epidemic of overweight and obesity may signal the start of the *age of inactivity and obesity*. Rates of type 2 diabetes mellitus, hypertension, and lipid abnormalities are on the rise, trends that are particularly evident in children. If these risk factor trends continue, age-adjusted CVD mortality rates could increase in the coming years.

PATTERNS IN THE EPIDEMIOLOGIC TRANSITION

Unique regional features have modified aspects of the transition in various parts of the world. High-income countries experienced declines in CVD death rates by as much as 50–60% over the last 60 years, whereas CVD death rates increased by 15% over the past 20 years in the low- and middle-income range. However, given the large amount of available data, the United States serves as a useful reference point for comparisons. The age of pestilence and famine occurred before 1900, with a largely agrarian economy and population. Infectious diseases accounted for more deaths than any other cause. By the 1930s, the country proceeded through the age of receding pandemics. The establishment of public health infrastructures resulted in dramatic declines in infectious disease mortality rates. Lifestyle changes due to rapid urbanization resulted in a simultaneous increase in CVD mortality rates, reaching approximately 390 per 100,000. Between 1930 and 1965, the country entered the age of degenerative

TABLE 226e-1 FIVE STAGES OF THE EPIDEMIOLOGIC TRANSITION

Stage	Description	Deaths Related to CVD, %	Predominant CVD Type
Pestilence and famine	Predominance of malnutrition and infectious diseases as causes of death; high rates of infant and child mortality; low mean life expectancy	<10	Rheumatic heart disease, cardiomyopathies caused by infection and malnutrition
Receding pandemics	Improvements in nutrition and public health lead to decrease in rates of deaths related to malnutrition and infection; precipitous decline in infant and child mortality rates	10–35	Rheumatic valvular disease, hypertension, CHD, and stroke (predominantly hemorrhagic)
Degenerative and man-made diseases	Increased fat and caloric intake and decrease in physical activity lead to emergence of hypertension and atherosclerosis; with increase in life expectancy, mortality from chronic, noncommunicable diseases exceeds mortality from malnutrition and infectious disease	35–65	CHD and stroke (ischemic and hemorrhagic)
Delayed degenerative diseases	CVD and cancer are the major causes of morbidity and mortality; better treatment and prevention efforts help avoid deaths among those with disease and delay primary events; age-adjusted CVD mortality declines; CVD affecting older and older individuals	40–50	CHD, stroke, and congestive heart failure
Inactivity and obesity	Overweight and obesity increase at alarming rate; diabetes and hypertension increase; decline in smoking rates levels off; a minority of the population meets physical activity recommendations	33	CHD, stroke, and congestive heart failure, peripheral vascular disease

Abbreviations: CHD, coronary heart disease; CVD, cardiovascular disease.

Source: Adapted from AR Omran: The epidemiologic transition: A theory of the epidemiology of population change. *Milbank Mem Fund Q* 49:509, 1971; and SJ Olshansky, AB Ault: The fourth stage of the epidemiologic transition: The age of delayed degenerative diseases. *Milbank Q* 64:355, 1986.

and man-made diseases. Infectious disease mortality rates fell to fewer than 50 per 100,000 per year, whereas CVD mortality rates reached peak levels with increasing urbanization and lifestyle changes in diet, physical activity, and tobacco consumption. The age of delayed degenerative diseases took place between 1965 and 2000. New therapeutic approaches, preventive measures, and exposure to public health campaigns promoting lifestyle modifications led to substantial declines in age-adjusted mortality rates and a steadily rising age at which a first CVD event occurs.



Currently, the United States is entering what appears to be a fifth phase. The decline in the age-adjusted CVD death rate of 3% per year through the 1970s and 1980s has tapered off in the 1990s to 2%. However, CVD death rates have declined by 3–5% per year during the first decade of the new millennium. Competing trends appear to be at play. One the one hand, an increase in the prevalence of diabetes and obesity, a slowing in the rate of decline in smoking, and a leveling off in the rate of detection and treatment for hypertension are in the negative column. On the other hand, cholesterol levels continue to decline in the face of increased statin use.

Many high-income countries (HICs)—which together account for 15% of the population—have proceeded through four stages of the epidemiologic transition in roughly the same pattern as the United States. CHD is the dominant form of CVD in these countries, with rates that tend to be two- to five-fold higher than stroke rates. However, variations exist. Whereas North America, Australia, and central northwestern European HICs experienced significant increases then rapid declines in CVD rates, southern and central European countries experienced a more gradual rise and fall in rates. More specifically, central European countries (i.e., Austria, Belgium, and Germany) declined at slower rates compared to their northern counterparts (i.e., Finland, Sweden, Denmark, and Norway). Countries such as Portugal, Spain, and Japan never reached the high mortality rates that the United States and other countries did, with CHD mortality rates at 200 per 100,000, or less. The countries of Western Europe also exhibit a clear north/south gradient in absolute rates of CVD, with rates highest in northern countries (i.e., Finland, Ireland, and Scotland) and lowest in Mediterranean countries (i.e., France, Spain, and Italy). Japan is unique among the HICs, most likely due to the unique dietary patterns of its population. Although stroke rates increased dramatically, CHD rates did not rise as sharply in Japan. However, Japanese dietary habits are undergoing substantial changes, reflected in an increase in cholesterol levels.

Patterns in low- and middle-income countries (LMICs; gross national income per capita less than U.S. \$12,615) depend, in part, on cultural differences, secular trends, and responses at the country level, with regard to both public health and treatment infrastructure. Although communicable diseases continue to be a major cause of death, CVD has emerged as a significant health concern in LMICs. With 85% of the world's population, LMICs are driving the rates of change in the global burden of CVD (Fig. 266e-1). In most LMICs, an urban/rural gradient has emerged for CHD, stroke, and hypertension, with higher rates in urban centers.

However, although CVD rates are rapidly rising, vast differences exist among the regions and countries, and even within the countries themselves (Fig. 266e-2). The East Asia and Pacific regions appear to be straddling the second and third phases of the epidemiologic transition. CVD is a major cause of death in China, but like Japan, stroke causes more deaths than CHD in a ratio of about three to one. Vietnam and Cambodia, on the other hand, are just emerging from the pestilence and famine transition. The Middle East and North Africa regions also appear to be entering the third phase of the epidemiologic transition, with increasing life expectancy and CVD death rates just below those of HICs. In general, Latin America appears to be in the third phase of the transition, although there is vast regional heterogeneity with some areas in the second phase of the transition and some in the fourth. The Eastern Europe and Central Asia regions, however, are firmly in the peak of the third phase, with the highest death rates due to CVD (~66%) in the world. Importantly, deaths due to CHD are not limited to the elderly in this region and have a significant effect on working-age populations. South Asia—and more specifically, India, which accounts for the greatest proportion of the region's population—is experiencing an alarming increase in heart disease. The transition appears to be in the Western style, with CHD as the dominant form of CVD. However, rheumatic heart disease continues to be a major cause of morbidity and mortality. As in South Asia, rheumatic heart disease is also an important cause of CVD morbidity and mortality in sub-Saharan Africa, which largely remains in the first phase of the epidemiologic transition.

Many factors contribute to this heterogeneity among LMICs. First, the regions are in various stages of the epidemiologic transition. Second, vast differences in lifestyle and behavioral risk factors exist. Third, racial and ethnic differences may lead to altered susceptibilities to various forms of CVD. In addition, it should be noted that for most countries in these regions, accurate country-wide data on cause-specific mortality are not complete.

Global deaths by cause, 2010

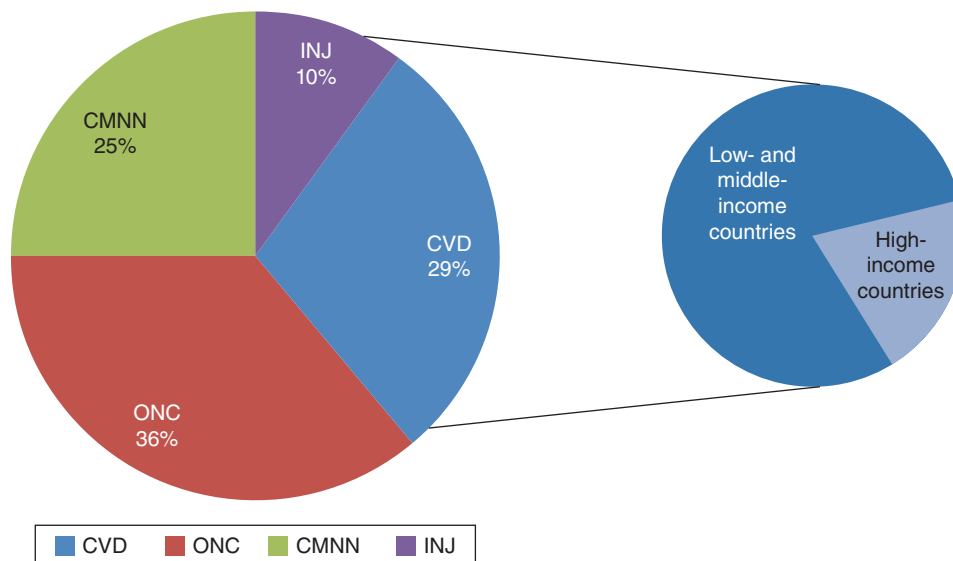


FIGURE 266-1 Global deaths by cause, 2010. CMNN, communicable, maternal, neonatal, and nutritional disorders; CVD, cardiovascular diseases; INJ, injuries; ONC, other noncommunicable diseases. (Based on data from *Global Burden of Disease Study 2010: Global Burden of Disease Study 2010 Mortality Results 1970–2010*. Seattle, Institute for Health Metrics and Evaluation, 2012.)

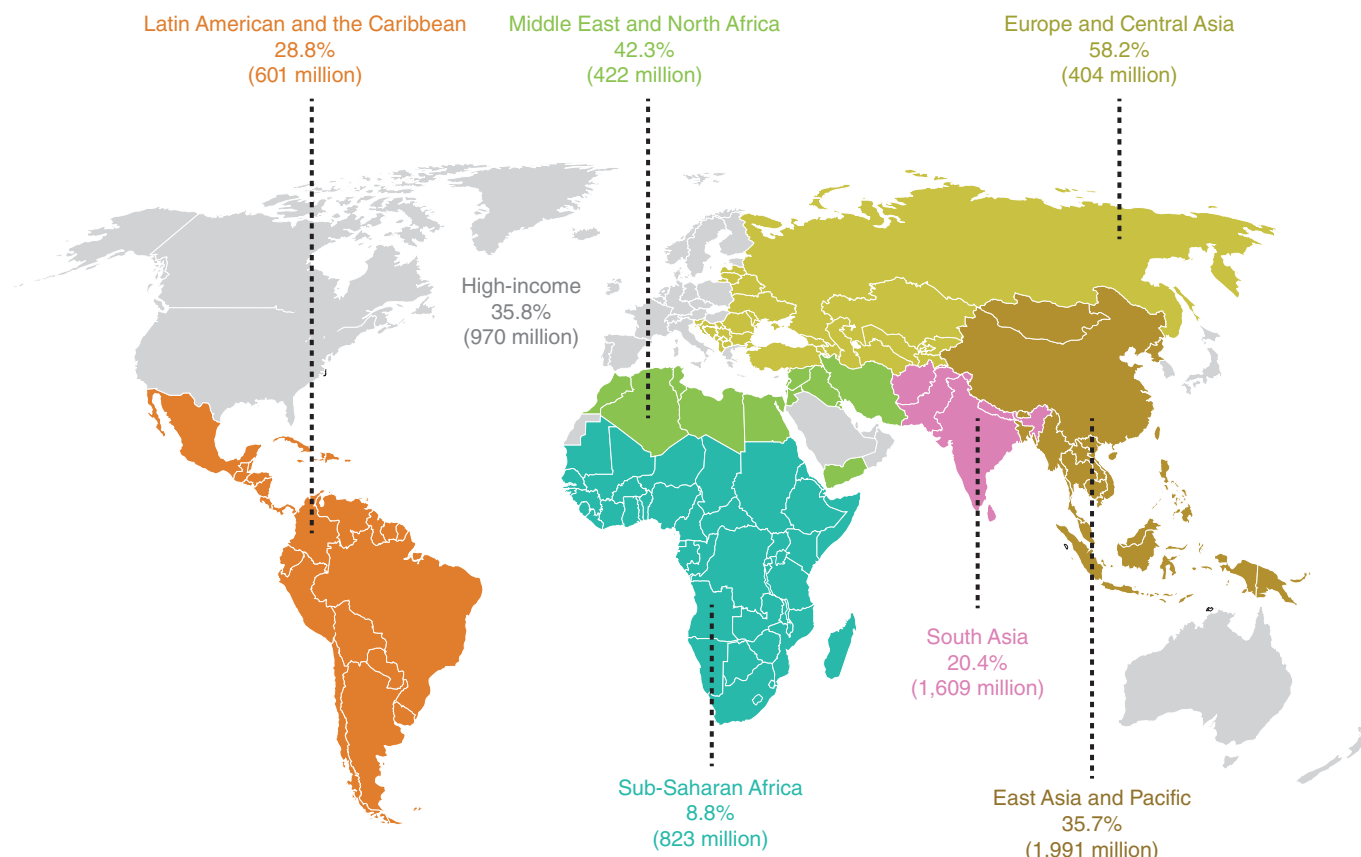



FIGURE 266-2 Cardiovascular disease deaths as a percentage of total deaths and total population in seven economic regions of the world defined by the World Bank. (Based on data from *Global Burden of Disease Study 2010: Global Burden of Disease Study 2010 Mortality Results 1970–2010*. Seattle, Institute for Health Metrics and Evaluation, 2012.)

GLOBAL TRENDS IN CARDIOVASCULAR DISEASE

 CVD accounts for nearly 30% of deaths worldwide, a number that is expected to increase. In 2010, CHD accounted for 13.3% of all deaths globally and the largest portion of global years of life lost (YLLs) and disability-adjusted life-years (DALYs). The second largest cause of death was stroke (11.1% of all deaths), which was also the third largest contributor to global YLLs and DALYs (Table 266e-2). Together, CHD and stroke accounted for nearly a quarter of all deaths worldwide. The burden of stroke is of growing concern among LMICs. The impact of stroke on DALYs and mortality rates is more than three times greater in LMICs as compared to HICs. By 2030, the number of deaths due to stroke is projected to increase by more than 30%, the majority of which will occur in LMICs.

With nearly 85% of the world's population, LMICs largely drive global CVD rates and trends. Ten million CVD deaths occurred in

LMICs in 2010, compared to 5.6 million in HICs. Globally, there is evidence of significant delays in age of occurrence and/or improvements in case fatality rates; between 1990 and 2010, the number of CVD deaths increased by 31%, but age-adjusted death rates decreased by 21.2% in the same period.

Although HIC population growth will be fueled by emigration from LMICs, the populations of HICs will shrink as a proportion of the world's population. The modest decline in CVD death rates that began in the HICs in the latter third of the twentieth century will continue, but the rate of decline appears to be slowing. However, these countries are expected to see an increase in the prevalence of CVD, as well as the absolute number of deaths as the population ages.

Significant portions of the population living in LMICs have entered the third phase of the epidemiologic transition, and some are entering the fourth stage. Changing demographics play a significant role in future predictions for CVD throughout the world. For example, the population growth rate in Eastern Europe and Central Asia was 0.7% in 2012, whereas it was 1.3% in South Asia.

CVD rates will also have an economic impact. Even assuming no increase in CVD risk factors, most countries, but especially India and South Africa, will see a large number of people between 35 and 64 die of CVD over the next 30 years, as well as an increasing level of morbidity among middle-aged people related to heart disease and stroke. In China, it is estimated that there will be 9 million deaths from CVD in 2030—up from 2.4 million in 2002—with half occurring in individuals between 35 and 64 years old.

RISK FACTORS

The global variation in CVD rates is related to temporal and regional variations in known risk behaviors and factors. Ecological analyses of major CVD risk factors and mortality demonstrate high correlations between expected and observed mortality rates for the three main risk factors—smoking, serum cholesterol, and hypertension—and suggest

TABLE 266e-2 MORBIDITY RELATED TO HEART DISEASE: 2010–2030

Deaths	2010	By 2030
CVD deaths: annual number of all deaths	15.6 million	24.2 million
CVD deaths: percentage of all deaths	30.0%	32.5%
CHD deaths: percentage of all male deaths	13.0%	14.9%
CHD deaths: percentage of all female deaths	14.0%	13.1%
Stroke deaths: percentage of all male deaths	9.9%	10.4%
Stroke deaths: percentage of all female deaths	13.0%	11.8%

Abbreviations: CHD, coronary heart disease; CVD, cardiovascular disease.

Source: Adapted from *Global Burden of Disease Study 2010: Global Burden of Disease Study 2010 Mortality Results 1970–2010*. Seattle, Institute for Health Metrics and Evaluation, 2012; J Mackay, G Mensah: *Atlas of Heart Disease and Stroke*. Geneva, World Health Organization, 2004.

Behavioral Risk Factors • TOBACCO Over 1.3 billion people use tobacco worldwide, a number that is projected to increase to 1.6 billion by 2030. Tobacco use currently causes about 5 million deaths annually (9% of all deaths), approximately 1.6 million of which are CVD-related. If current smoking patterns continue, the global burden of disease attributable to tobacco will reach 10 million deaths by 2030. Although tobacco use has been greatest in HICs historically, consumption has shifted dramatically to LMICs in recent decades. Some of the highest tobacco use now occurs in the East Asia and Pacific region. A unique feature of LMICs is easy access to smoking during the early stages of the epidemiologic transition due to the availability of relatively inexpensive tobacco products. In South Asia, the prominence of other locally produced forms of tobacco besides manufactured cigarettes makes control of consumption more challenging. Secondhand smoke is another well-established cause of CHD, responsible for 600,000 deaths of nonsmokers in 2011. Although smoking bans have both immediate and long-term benefits, implementation varies greatly between countries.

DIET Total caloric intake per capita increases as countries develop. With regard to CVD, a key element of dietary change is an increase in intake of saturated animal fats and hydrogenated vegetable fats, which contain atherogenic *trans* fatty acids, along with a decrease in intake of plant-based foods and an increase in simple carbohydrates. Fat contributes less than 20% of calories in rural China and India, less than 30% in Japan, and well above 30% in the United States. Caloric contributions from fat appear to be falling in the HICs. In the United States, between 1971 and 2010, the percentage of calories derived from saturated fat decreased from 13% to 11%.

PHYSICAL INACTIVITY The increased mechanization that accompanies the economic transition leads to a shift from physically demanding, agriculture-based work to largely sedentary industry- and office-based work. In the United States, approximately one-quarter of the population does not participate in any leisure-time physical activity, and only 51.6% of adults report engaging in physical activity three or more times a week. Physical inactivity is similarly high in other regions of the world and is increasing in countries that are rapidly urbanizing as part of their economic transition. In urban China, for example, the proportion of adults who participate in moderate- or high-level activity has decreased significantly, whereas those who participate in low-level activity has increased.

METABOLIC RISK FACTORS

Examination of trends in metabolic risk factors provides insight into changes in the CVD burden globally. Here we describe four metabolic risk factors—lipid levels, hypertension, obesity, and diabetes mellitus—using data from the Global Burden of Disease, Injuries, and Risk Factors Study (GBD 2010). The GBD project identified and compiled mortality and morbidity data from 187 countries from 1980 to 2010.

Lipid Levels Worldwide, high cholesterol levels are estimated to play a role in 56% of ischemic heart disease events and 18% of strokes, amounting to 4.4 million deaths annually. Although mean population plasma cholesterol levels tend to rise as countries move through the epidemiologic transition, mean serum total cholesterol levels have decreased globally between 1980 and 2008 by 0.08 mmol/L per decade in men and 0.07 mmol/L per decade in women. In 2008, age-standardized mean total cholesterol was 4.64 mmol/L (179.4 mg/dL) in men and 4.76 mmol/L (184.2 mg/dL) in women. Large declines occurred in Australasia, North America, and Western Europe (0.19–0.21 mmol/L). Countries in the East Asia and Pacific region experienced increases of greater than 0.08 mmol/L in both men and women. Social and individual changes that accompany urbanization clearly play a role because plasma cholesterol levels tend to be higher among urban residents than among rural residents. This shift is largely driven by greater consumption of dietary fats—primarily from animal products and processed vegetable oils—and decreased physical activ-

ity. In HICs, in general, mean population cholesterol levels are falling, whereas wide variation is seen in the LMICs.

Hypertension Elevated blood pressure is an early indicator of the epidemiologic transition. Worldwide, approximately 62% of strokes and 49% of CHD are attributable to suboptimal (>115 mmHg systolic) blood pressure, which is believed to account for more than 7 million deaths annually. Remarkably, nearly half of this burden occurs among those with systolic blood pressure less than 140 mmHg, even as this level is used at the arbitrary threshold for defining hypertension in many national guidelines. Between 1980 and 2008, the age-standardized prevalence of uncontrolled prevalence has decreased even as the number of people with uncontrolled hypertension has increased. This trend results largely from population growth and aging. Rising mean population blood pressure also occurs as populations industrialize and move from rural to urban settings. For example, the prevalence of hypertension in urban India is 25%, but varies between 10% and 15% in rural communities. One major concern in LMICs is the high rate of undetected, and therefore untreated, hypertension. This may explain, at least in part, the higher stroke rates in these countries in relation to CHD rates during the early stages of the transition. The high rates of hypertension throughout Asia, especially undiagnosed hypertension, likely contribute to the high prevalence of hemorrhagic stroke in the region. Globally, however, mean systolic blood pressure has decreased among both genders (0.8 mmHg per decade among men; 1.0 mmHg per decade among women).

Obesity Although clearly associated with increased risk of CHD, much of the risk posed by obesity may be mediated by other CVD risk factors, including hypertension, diabetes mellitus, and lipid profile imbalances. According to the latest GBD data, nearly 1.46 billion adults were overweight (body mass index ≥ 25 kg/m²) in 2008, and approximately 508 million were obese (BMI ≥ 30 kg/m²). Obesity is increasing throughout the world, particularly in developing countries, where the trajectories are steeper than those experienced by the developed countries. In many of the LMICs, obesity appears to coexist with undernutrition and malnutrition. Adolescents are at particular risk. Currently, 1 in 10 children are estimated to be overweight, a number that is increasing worldwide. Women are also more affected than men, with the number of overweight women generally exceeding underweight women based on data from 36 LMICs.

Diabetes Mellitus As a consequence of, or in addition to, increasing body mass index and decreasing levels of physical activity, worldwide rates of diabetes—predominantly type 2 diabetes—are on the rise. According to the most recent data from the GBD project, mean fasting plasma glucose levels have increased globally between 1980 and 2008. An estimated 346 million people worldwide have diabetes. The International Diabetes Foundation predicts that this number will reach 522 million by 2030, a yearly rate of growth that is higher than that of the world's adult population. Nearly 50% of people with diabetes are undiagnosed, and 80% live in LMICs. The highest regional prevalence for diabetes occurs in the Middle East and North Africa, where an estimated 12.5% of the adult population has diabetes. Future growth will also largely occur in this region, along with other LMICs in South Asia and sub-Saharan Africa. There appear to be clear genetic susceptibilities to diabetes mellitus of various racial and ethnic groups. For example, migration studies suggest that South Asians and Indians tend to be at higher risk than those of European extraction.

SUMMARY

Although CVD rates are declining in the HICs, they are increasing in virtually every other region of the world. The consequences of this preventable epidemic will be substantial on many levels, including individual mortality and morbidity, family suffering, and staggering economic costs.

Three complementary strategies can be used to lessen the impact. First, the overall burden of CVD risk factors can be lowered through population-wide public health measures, such as national campaigns

against cigarette smoking, unhealthy diets, and physical inactivity. Second, it is important to identify higher risk subgroups of the population who stand to benefit the most from specific, low-cost prevention interventions, including screening for and treatment of hypertension and elevated cholesterol. Simple, low-cost interventions, such as the “polypill,” a regimen of aspirin, a statin, and an antihypertensive agent, also need to be explored. Third, resources should be allocated to acute as well as secondary prevention interventions. For countries with limited resources, a critical first step in developing a comprehensive plan

is better assessment of cause-specific mortality and morbidity, as well as the prevalence of the major preventable risk factors.

In the meantime, the HICs must continue to bear the burden of research and development aimed at prevention and treatment, being mindful of the economic limitations of many countries. The concept of the epidemiologic transition provides insight into how to alter the course of the CVD epidemic. The efficient transfer of low-cost preventive and therapeutic strategies could alter the natural course of this epidemic and thereby reduce the excess global burden of preventable CVD.

SECTION 2 DIAGNOSIS OF CARDIOVASCULAR DISORDERS

267 Physical Examination of the Cardiovascular System

Patrick T. O'Gara, Joseph Loscalzo

The approach to a patient with known or suspected cardiovascular disease begins with the time-honored traditions of a directed history and a targeted physical examination. The scope of these activities depends on the clinical context at the time of presentation, ranging from an elective ambulatory follow-up visit to a more focused emergency department encounter. There has been a gradual decline in physical examination skills over the last two decades at every level, from student to faculty specialist, a development of great concern to both clinicians and medical educators. Classic cardiac findings are recognized by only a minority of internal medicine and family practice residents. Despite popular perceptions, clinical performance does not improve predictably as a function of experience; instead, the acquisition of new examination skills may become more difficult for a busy individual practitioner. Less time is now devoted to mentored cardiovascular examinations during the training of students and residents. One widely recognized outcome of these trends is the progressive overutilization of noninvasive imaging studies to establish the presence and severity of cardiovascular disease even when the examination findings imply a low pretest probability of significant pathology. Educational techniques to improve bedside skills include repetition, patient-centered teaching conferences, and visual display feedback of auscultatory events with Doppler echocardiographic imaging.

The evidence base that links the findings from the history and physical examination to the presence, severity, and prognosis of cardiovascular disease has been established most rigorously for coronary artery disease, heart failure, and valvular heart disease. For example, observations regarding heart rate, blood pressure, signs of pulmonary congestion, and the presence of mitral regurgitation (MR) contribute importantly to bedside risk assessment in patients with acute coronary syndromes. Observations from the physical examination in this setting can inform clinical decision making before the results of cardiac biomarkers testing are known. The prognosis of patients with systolic heart failure can be predicted on the basis of the jugular venous pressure (JVP) and the presence or absence of a third heart sound (S_3). Accurate characterization of cardiac murmurs provides important insight into the natural history of many valvular and congenital heart lesions. Finally, the important role played by the physical examination in enhancing the clinician-patient relationship cannot be overestimated.

THE GENERAL PHYSICAL EXAMINATION

Any examination begins with an assessment of the general appearance of the patient, with notation of age, posture, demeanor, and

overall health status. Is the patient in pain or resting quietly, dyspneic or diaphoretic? Does the patient choose to avoid certain body positions to reduce or eliminate pain, as might be the case with suspected acute pericarditis? Are there clues indicating that dyspnea may have a pulmonary cause, such as a barrel chest deformity with an increased anterior-posterior diameter, tachypnea, and pursed-lip breathing? Skin pallor, cyanosis, and jaundice can be appreciated readily and provide additional clues. A chronically ill-appearing emaciated patient may suggest the presence of long-standing heart failure or another systemic disorder, such as a malignancy. Various genetic syndromes, often with cardiovascular involvement, can also be recognized easily, such as trisomy 21, Marfan's syndrome, and Holt-Oram syndrome. Height and weight should be measured routinely, and both body mass index and body surface area should be calculated. Knowledge of the waist circumference and the waist-to-hip ratio can be used to predict long-term cardiovascular risk. Mental status, level of alertness, and mood should be assessed continuously during the interview and examination.

Skin Central cyanosis occurs with significant right-to-left shunting at the level of the heart or lungs, allowing deoxygenated blood to reach the systemic circulation. Peripheral cyanosis or acrocyanosis, in contrast, is usually related to reduced extremity blood flow due to small vessel constriction, as seen in patients with severe heart failure, shock, or peripheral vascular disease; it can be aggravated by the use of β -adrenergic blockers with unopposed α -mediated constriction. Differential cyanosis refers to isolated cyanosis affecting the lower but not the upper extremities in a patient with a large patent ductus arteriosus (PDA) and secondary pulmonary hypertension with right-to-left to shunting at the great vessel level. Hereditary telangiectasias on the lips, tongue, and mucous membranes, as part of the Osler-Weber-Rendu syndrome (hereditary hemorrhagic telangiectasia), resemble spider nevi and can be a source of right-to-left shunting when also present in the lung. Malar telangiectasias also are seen in patients with advanced mitral stenosis and scleroderma. An unusually tan or bronze discoloration of the skin may suggest hemochromatosis as the cause of the associated systolic heart failure. Jaundice, which may be visible first in the sclerae, has a broad differential diagnosis but, in the appropriate setting, can be consistent with advanced right heart failure and congestive hepatomegaly or late-term "cardiac cirrhosis." Cutaneous ecchymoses are seen frequently among patients taking vitamin K antagonists or antiplatelet agents such as aspirin and thienopyridines. Various lipid disorders sometimes are associated with subcutaneous xanthomas, particularly along the tendon sheaths or over the extensor surfaces of the extremities. Severe hypertriglyceridemia can be associated with eruptive xanthomatosis and lipemia retinalis. Palmar crease xanthomas are specific for type III hyperlipoproteinemia. Pseudoxanthoma elasticum, a disease associated with premature atherosclerosis, is manifested by a leathery, cobblestoned appearance of the skin in the axilla and neck creases and by angioid streaks on

fundoscopic examination. Extensive lentiginoses have been described in a variety of development delay–cardiovascular syndromes, including Carney’s syndrome, which includes multiple atrial myxomas. Cutaneous manifestations of sarcoidosis such as lupus pernio and erythema nodosum may suggest this disease as a cause of an associated dilated cardiomyopathy, especially with heart block, intraventricular conduction delay, or ventricular tachycardia.

Head and Neck Dentition and oral hygiene should be assessed in every patient both as a source of potential infection and as an index of general health. A high-arched palate is a feature of Marfan’s syndrome and other connective tissue disease syndromes. Bifid uvula has been described in patients with Loey-Dietz syndrome, and orange tonsils are characteristic of Tangier disease. The ocular manifestations of hyperthyroidism have been well described. Many patients with congenital heart disease have associated hypertelorism, low-set ears, or micrognathia. Blue sclerae are a feature of osteogenesis imperfecta. An arcus senilis pattern lacks specificity as an index of coronary heart disease risk. The fundoscopic examination is an often underused method by which to assess the microvasculature, especially among patients with established atherosclerosis, hypertension, or diabetes mellitus. A mydriatic agent may be necessary for optimal visualization. A fundoscopic examination should be performed routinely in the assessment of patients with suspected endocarditis and those with a history of acute visual change. Branch retinal artery occlusion or visualization of a Hollenhorst plaque can narrow the differential diagnosis rapidly in the appropriate setting. Relapsing polychondritis may manifest as an inflamed pinna or, in its later stages, as a saddle-nose deformity because of destruction of nasal cartilage; granulomatosis with polyangiitis (Wegener’s) can also lead to a saddle-nose deformity.

Chest Midline sternotomy, left posterolateral thoracotomy, or infraclavicular scars at the site of pacemaker/defibrillator generator implantation should not be overlooked and may provide the first clue regarding an underlying cardiovascular disorder in patients unable to provide a relevant history. A prominent venous collateral pattern may suggest subclavian or vena caval obstruction. If the head and neck appear dusky and slightly cyanotic and the venous pressure is grossly elevated without visible pulsations, a diagnosis of superior vena cava syndrome should be entertained. Thoracic cage abnormalities have been well described among patients with connective tissue disease syndromes. They include pectus carinatum (“pigeon chest”) and pectus excavatum (“funnel chest”). Obstructive lung disease is suggested by a barrel chest deformity, especially with tachypnea, pursed-lip breathing, and use of accessory muscles. The characteristically severe kyphosis and compensatory lumbar, pelvic, and knee flexion of ankylosing spondylitis should prompt careful auscultation for a murmur of aortic regurgitation (AR). Straight back syndrome refers to the loss of the normal kyphosis of the thoracic spine and has been described in patients with mitral valve prolapse (MVP) and its variants. In some patients with cyanotic congenital heart disease, the chest wall appears to be asymmetric, with anterior displacement of the left hemithorax. The respiratory rate and pattern should be noted during spontaneous breathing, with additional attention to depth, audible wheezing, and stridor. Lung examination can reveal adventitious sounds indicative of pulmonary edema, pneumonia, or pleuritis.

Abdomen In some patients with advanced obstructive lung disease, the point of maximal cardiac impulse may be in the epigastrium. The liver is frequently enlarged and tender in patients with chronic heart failure. Systolic pulsations over the liver signify severe tricuspid regurgitation (TR). Splenomegaly may be a feature of infective endocarditis, particularly when symptoms have persisted for weeks or months. Ascites is a nonspecific finding but may be present with advanced chronic right heart failure, constrictive pericarditis, hepatic cirrhosis, or an intraperitoneal malignancy. The finding of an elevated JVP implies a cardiovascular etiology. In nonobese patients, the aorta typically is palpated between the epigastrium and the umbilicus. The sensitivity of palpation for the detection of an abdominal aortic aneurysm (pulsatile and expansile mass) decreases as a function of body size. Because palpation alone is not sufficiently accurate to establish

this diagnosis, a screening ultrasound examination is advised. The presence of an arterial bruit over the abdomen suggests high-grade atherosclerotic disease, although precise localization is difficult.

Extremities The temperature and color of the extremities, the presence of clubbing, arachnodactyly, and pertinent nail findings can be surmised quickly during the examination. Clubbing implies the presence of central right-to-left shunting, although it has also been described in patients with endocarditis. Its appearance can range from cyanosis and softening of the root of the nail bed, to the classic loss of the normal angle between the base of the nail and the skin, to the skeletal and periosteal bony changes of hypertrophic osteoarthropathy, which is seen rarely in patients with advanced lung or liver disease. Patients with the Holt-Oram syndrome have an unopposable, “fingerized” thumb, whereas patients with Marfan’s syndrome may have arachnodactyly and a positive “wrist” (overlapping of the thumb and fifth finger around the wrist) or “thumb” (protrusion of the thumb beyond the ulnar aspect of the hand when the fingers are clenched over the thumb in a fist) sign. The Janeway lesions of endocarditis are nontender, slightly raised hemorrhages on the palms and soles, whereas Osler’s nodes are tender, raised nodules on the pads of the fingers or toes. Splinter hemorrhages are classically identified as linear petechiae in the midposition of the nail bed and should be distinguished from the more common traumatic petechiae, which are seen closer to the distal edge.

Lower extremity or presacral edema in the setting of an elevated JVP defines volume overload and may be a feature of chronic heart failure or constrictive pericarditis. Lower extremity edema in the absence of jugular venous hypertension may be due to lymphatic or venous obstruction or, more commonly, to venous insufficiency, as further suggested by the appearance of varicosities, venous ulcers (typically medial in location), and brownish cutaneous discoloration from hemosiderin deposition (eburnation). Pitting edema can also be seen in patients who use dihydropyridine calcium channel blockers. A Homan’s sign (posterior calf pain on active dorsiflexion of the foot against resistance) is neither specific nor sensitive for deep venous thrombosis. Muscular atrophy or the absence of hair along an extremity is consistent with severe arterial insufficiency or a primary neuromuscular disorder.

CARDIOVASCULAR EXAMINATION

Jugular Venous Pressure and Waveform JVP is the single most important bedside measurement from which to estimate the volume status. The internal jugular vein is preferred because the external jugular vein is valved and not directly in line with the superior vena cava and right atrium. Nevertheless, the external jugular vein has been used to discriminate between high and low central venous pressure (CVP) when tested among medical students, residents, and attending physicians. Precise estimation of the central venous or right atrial pressure from bedside assessment of the jugular venous waveform has proved difficult. Venous pressure traditionally has been measured as the vertical distance between the top of the jugular venous pulsation and the sternal inflection point (angle of Louis). A distance >4.5 cm at 30° elevation is considered abnormal. However, the actual distance between the mid-right atrium and the angle of Louis varies considerably as a function of both body size and the patient angle at which the assessment is made (30° , 45° , or 60°). The use of the sternal angle as a reference point leads to systematic underestimation of CVP, and this method should be used less for semiquantification than to distinguish a normal from an abnormally elevated CVP. The use of the clavicle may provide an easier reference for standardization. Venous pulsations above this level in the sitting position are clearly abnormal, as the distance between the clavicle and the right atrium is at least 10 cm. The patient should always be placed in the sitting position, with the legs dangling below the bedside, when an elevated pressure is suspected in the semisupine position. It should also be noted that bedside estimates of CVP are made in centimeters of water but must be converted to millimeters of mercury to provide correlation with accepted hemodynamic norms ($1.36 \text{ cmH}_2\text{O} = 1.0 \text{ mmHg}$).

The venous waveform sometimes can be difficult to distinguish from the carotid pulse, especially during casual inspection. Nevertheless, the venous waveform has several characteristic features, and its individual components can be appreciated in most patients (Fig. 267-1). The arterial pulsation is not easily obliterated with palpation; the venous waveform in patients with sinus rhythm is usually biphasic, while the carotid pulse is monophasic; and the jugular venous pulsation should change with changes in posture or inspiration (unless the venous pressure is quite elevated).

The venous waveform is divided into several distinct peaks. The *a* wave reflects right atrial presystolic contraction and occurs just after the electrocardiographic P wave, preceding the first heart sound (S_1). A prominent *a* wave is seen in patients with reduced right ventricular compliance; a cannon *a* wave occurs with atrioventricular (AV) dissociation and right atrial contraction against a closed tricuspid valve. In a patient with a wide complex tachycardia, the appreciation of cannon *a* waves in the jugular venous waveform identifies the rhythm as ventricular in origin. The *a* wave is not present with atrial fibrillation. The *x* descent defines the fall in right atrial pressure after inscription of the *a* wave. The *c* wave interrupts this *x* descent and is followed by a further descent. The *v* wave represents atrial filling (atrial diastole) and occurs during ventricular systole. The height of the *v* wave is determined by right atrial compliance as well as the volume of blood returning to the right atrium either antegrade from the cavae or retrograde through an incompetent tricuspid valve. In patients with TR, the *v* wave is accentuated and the subsequent fall in pressure (*y* descent) is rapid. With progressive degrees of TR, the *v* wave merges with the *c* wave, and the right atrial and jugular vein waveforms become “ventricularized.” The *y* descent, which follows the peak of the *v* wave, can become prolonged or blunted with obstruction to right ventricular inflow, as may occur with tricuspid stenosis or pericardial tamponade. Normally, the venous pressure should fall by at least 3 mmHg with inspiration. Kussmaul’s sign is defined by either a rise or a lack of fall of the JVP with inspiration and is classically associated with constrictive pericarditis, although it has been reported in patients with restrictive cardiomyopathy, massive pulmonary embolism, right ventricular infarction, and advanced left ventricular systolic heart failure. It is also a common, isolated finding in patients after cardiac surgery without other hemodynamic abnormalities.

Venous hypertension sometimes can be elicited by performance of the abdominojugular reflex or with passive leg elevation. When these signs are positive, a volume-overloaded state with limited compliance of an overly distended or constricted venous system is present. The abdominojugular reflex is elicited with firm and consistent pressure over the upper portion of the abdomen, preferably over the right upper quadrant, for at least 10 s. A positive response is defined by a sustained rise of more than 3 cm in JVP for at least 15 s after release of the hand. Patients must be coached to refrain from breath holding or a Valsalva-like maneuver during the procedure. The abdominojugular reflex is useful in predicting a pulmonary artery wedge pressure in excess of 15 mmHg in patients with heart failure.

Although the JVP estimates right ventricular filling pressure, it has a predictable relationship with the pulmonary artery wedge pressure. In a large study of patients with advanced heart failure, the presence of a right atrial pressure >10 mmHg (as predicted on bedside examination) had a positive value of 88% for the prediction of a pulmonary artery wedge pressure of >22 mmHg. In addition, an elevated JVP has prognostic significance in patients with both symptomatic heart failure and asymptomatic left ventricular systolic dysfunction. The presence of an elevated JVP is associated with a higher risk of subsequent hospitalization for heart failure, death from heart failure, or both.

Assessment of Blood Pressure Measurement of blood pressure usually is delegated to a medical assistant but should be repeated by the clinician. Accurate measurement depends on body position, arm size, time of measurement, place of measurement, device, device size, technique, and examiner. In general, physician-recorded blood pressures are higher than both nurse-recorded pressures and self-recorded pressures at home. Blood pressure is best measured in the seated position with

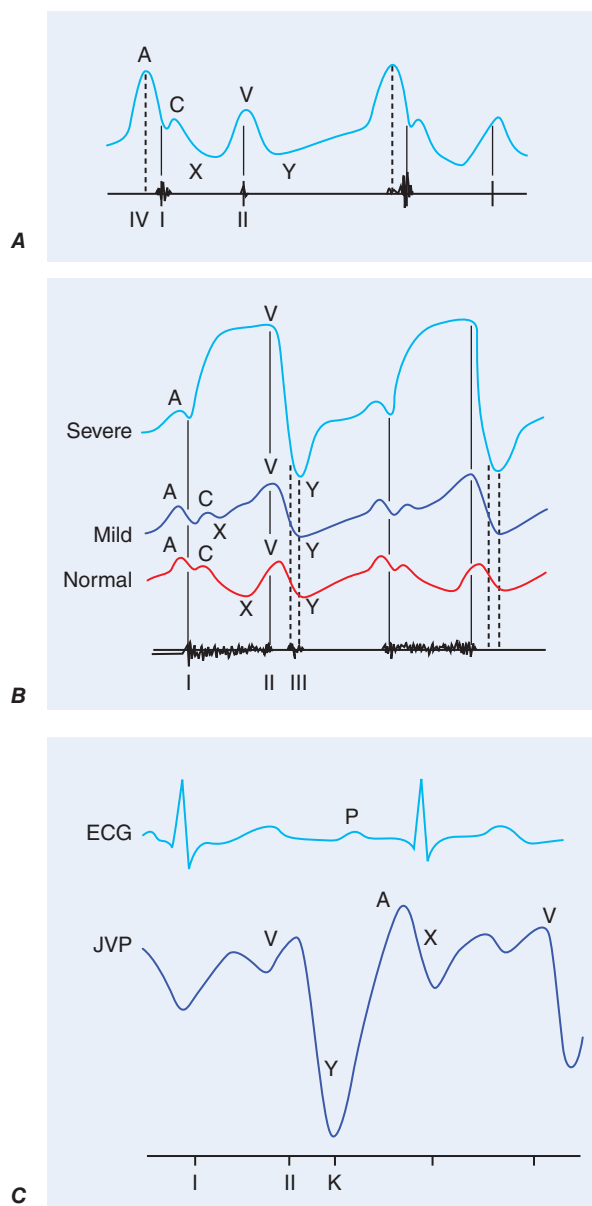


FIGURE 267-1 A. Jugular venous pulse wave tracing (top) with heart sounds (bottom). The A wave represents right atrial presystolic contraction and occurs just after the electrocardiographic P wave and just before the first heart sound (I). In this example, the A wave is accentuated and larger than normal due to decreased right ventricular compliance, as also suggested by the right-sided S_4 (IV). The C wave may reflect the carotid pulsation in the neck and/or an early systolic increase in right atrial pressure as the right ventricle pushes the closed tricuspid valve into the right atrium. The *x* descent follows the A wave just as atrial pressure continues to fall. The V wave represents atrial filling during ventricular systole and peaks at the second heart sound (II). The *y* descent corresponds to the fall in right atrial pressure after tricuspid valve opening. B. Jugular venous wave forms in mild (middle) and severe (top) tricuspid regurgitation, compared with normal, with phonocardiographic representation of the corresponding heart sounds below. With increasing degrees of tricuspid regurgitation, the waveform becomes “ventricularized.” C. Electrocardiogram (ECG) (top), jugular venous waveform (JVP) (middle), and heart sounds (bottom) in pericardial constriction. Note the prominent and rapid *y* descent, corresponding in timing to the pericardial knock (K). (From J Abrams: *Synopsis of Cardiac Physical Diagnosis*, 2nd ed. Boston, Butterworth Heinemann, 2001, pp 25–35.)

the arm at the level of the heart, using an appropriately sized cuff, after 5–10 min of relaxation. When it is measured in the supine position, the arm should be raised to bring it to the level of the mid-right atrium. The length and width of the blood pressure cuff bladder should be 80% and 40% of the arm's circumference, respectively. A common source of error in practice is to use an inappropriately small cuff, resulting in marked overestimation of true blood pressure, or an inappropriately large cuff, resulting in underestimation of true blood pressure. The cuff should be inflated to 30 mmHg above the expected systolic pressure and the pressure released at a rate of 2–3 mmHg/s. Systolic and diastolic pressures are defined by the first and fifth Korotkoff sounds, respectively. Very low (even 0 mmHg) diastolic blood pressures may be recorded in patients with chronic, severe AR or a large arteriovenous fistula because of enhanced diastolic “run-off.” In these instances, both the phase IV and phase V Korotkoff sounds should be recorded. Blood pressure is best assessed at the brachial artery level, though it can be measured at the radial, popliteal, or pedal pulse level. In general, systolic pressure increases and diastolic pressure decreases when measured in more distal arteries. Blood pressure should be measured in both arms, and the difference should be less than 10 mmHg. A blood pressure differential that exceeds this threshold may be associated with atherosclerotic or inflammatory subclavian artery disease, supravalvular aortic stenosis, aortic coarctation, or aortic dissection. Systolic leg pressures are usually as much as 20 mmHg higher than systolic arm pressures. Greater leg–arm pressure differences are seen in patients with chronic severe AR as well as patients with extensive and calcified lower extremity peripheral arterial disease. The ankle-brachial index (lower pressure in the dorsalis pedis or posterior tibial artery divided by the higher of the two brachial artery pressures) is a powerful predictor of long-term cardiovascular mortality.

The blood pressure measured in an office or hospital setting may not accurately reflect the pressure in other venues. “White coat hypertension” is defined by at least three separate clinic-based measurements >140/90 mmHg and at least two non-clinic-based measurements <140/90 mmHg in the absence of any evidence of target organ damage. Individuals with white coat hypertension may not benefit from drug therapy, although they may be more likely to develop sustained hypertension over time. Masked hypertension should be suspected when normal or even low blood pressures are recorded in patients with advanced atherosclerotic disease, especially when evidence of target organ damage is present or bruits are audible.

Orthostatic hypotension is defined by a fall in systolic pressure >20 mmHg or in diastolic pressure >10 mmHg in response to assumption of the upright posture from a supine position within 3 min. There may also be a lack of a compensatory tachycardia, an abnormal response that suggests autonomic insufficiency, as may be seen in patients with diabetes or Parkinson's disease. Orthostatic hypotension is a common cause of postural lightheadedness/syncope and should be assessed routinely in patients for whom this diagnosis might pertain. It can be exacerbated by advanced age, dehydration, certain medications, food, deconditioning, and ambient temperature.

Arterial Pulse The carotid artery pulse occurs just after the ascending aortic pulse. The aortic pulse is best appreciated in the epigastrium, just above the level of the umbilicus. Peripheral arterial pulses that should be assessed routinely include the subclavian, brachial, radial, ulnar, femoral, popliteal, dorsalis pedis, and posterior tibial. In patients in whom the diagnosis of either temporal arteritis or polymyalgia rheumatica is suspected, the temporal arteries also should be examined. Although one of the two pedal pulses may not be palpable in up to 10% of normal subjects, the pair should be symmetric. The integrity of the arcuate system of the hand is assessed by Allen's test, which is performed routinely before instrumentation of the radial artery. The pulses should be examined for their symmetry, volume, timing, contour, amplitude, and duration. If necessary, simultaneous auscultation of the heart can help identify a delay in the arrival of an arterial pulse. Simultaneous palpation of the radial and femoral pulses may reveal a femoral delay in a patient with hypertension and suspected aortic coarctation. The carotid upstrokes should never be examined simultaneously or before listening for a bruit. Light pressure

should always be used to avoid precipitation of carotid hypersensitivity syndrome and syncope in a susceptible elderly individual. The arterial pulse usually becomes more rapid and spiking as a function of its distance from the heart, a phenomenon that reflects the muscular status of the more peripheral arteries and the summation of the incident and reflected waves. In general, the character and contour of the arterial pulse depend on the stroke volume, ejection velocity, vascular compliance, and systemic vascular resistance. The pulse examination can be misleading in patients with reduced cardiac output and in those with stiffened arteries from aging, chronic hypertension, or peripheral arterial disease.

The character of the pulse is best appreciated at the carotid level (Fig. 267-2). A weak and delayed pulse (*pulsus parvus et tardus*) defines severe aortic stenosis (AS). Some patients with AS may also have a slow, notched, or interrupted upstroke (anacrotic pulse) with a thrill or shudder. With chronic severe AR, by contrast, the carotid upstroke has a sharp rise and rapid fall-off (Corrigan's or water-hammer pulse). Some patients with advanced AR may have a bifid or bisferiens pulse, in which two systolic peaks can be appreciated. A bifid pulse is also described in patients with hypertrophic obstructive cardiomyopathy (HOCM), with inscription of percussion and tidal waves. A bifid pulse is easily appreciated in patients on intraaortic balloon counterpulsation (IABP), in whom the second pulse is diastolic in timing.

Pulsus paradoxus refers to a fall in systolic pressure >10 mmHg with inspiration that is seen in patients with pericardial tamponade but also is described in those with massive pulmonary embolism, hemorrhagic shock, severe obstructive lung disease, and tension pneumothorax.

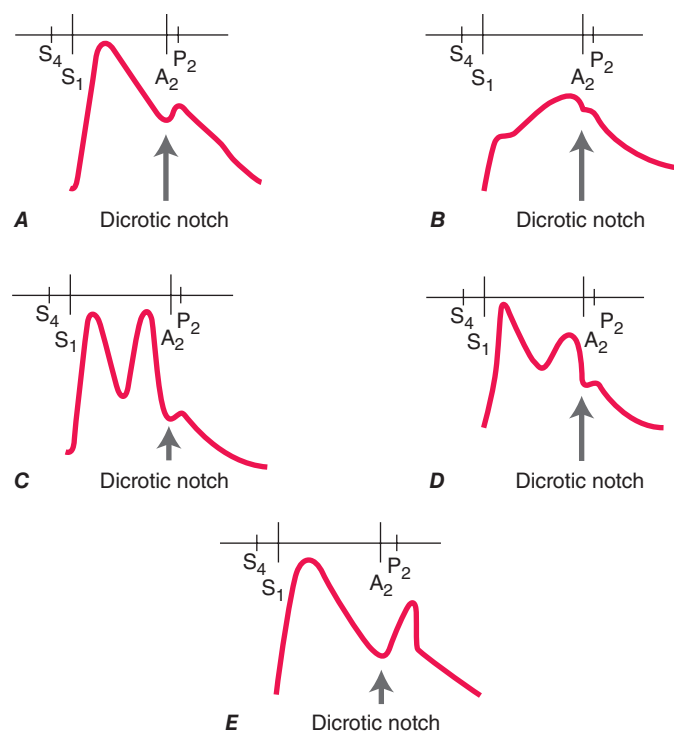


FIGURE 267-2 Schematic diagrams of the configurational changes in carotid pulse and their differential diagnoses. Heart sounds are also illustrated. **A.** Normal. S_4 , fourth heart sound; S_1 , first heart sound; A_2 , aortic component of second heart sound; P_2 , pulmonic component of second heart sound. **B.** Aortic stenosis. Anacrotic pulse with slow upstroke to a reduced peak. **C.** Bisferiens pulse with two peaks in systole. This pulse is rarely appreciated in patients with severe aortic regurgitation. **D.** Bisferiens pulse in hypertrophic obstructive cardiomyopathy. There is a rapid upstroke to the first peak (percussion wave) and a slower rise to the second peak (tidal wave). **E.** Dicotic pulse with peaks in systole and diastole. This waveform may be seen in patients with sepsis or during intraaortic balloon counterpulsation with inflation just after the dicotic notch. (From K Chatterjee, W Parmley [eds]: *Cardiology: An Illustrated Text/Reference*. Philadelphia, Gower Medical Publishers, 1991.)

1446 Pulsus paradoxus is measured by noting the difference between the systolic pressure at which the Korotkoff sounds are first heard (during expiration) and the systolic pressure at which the Korotkoff sounds are heard with each heartbeat, independent of the respiratory phase. Between these two pressures, the Korotkoff sounds are heard only intermittently and during expiration. The cuff pressure must be decreased slowly to appreciate the finding. It can be difficult to measure pulsus paradoxus in patients with tachycardia, atrial fibrillation, or tachypnea. A pulsus paradoxus may be palpable at the brachial artery or femoral artery level when the pressure difference exceeds 15 mmHg. This inspiratory fall in systolic pressure is an exaggerated consequence of interventricular dependence.

Pulsus alternans, in contrast, is defined by beat-to-beat variability of pulse amplitude. It is present only when every other phase I Korotkoff sound is audible as the cuff pressure is lowered slowly, typically in a patient with a regular heart rhythm and independent of the respiratory cycle. Pulsus alternans is seen in patients with severe left ventricular systolic dysfunction and is thought to be due to cyclic changes in intracellular calcium and action potential duration. When pulsus alternans is associated with electrocardiographic T-wave alternans, the risk for an arrhythmic event appears to be increased.

Ascending aortic aneurysms can rarely be appreciated as a pulsatile mass in the right parasternal area. Appreciation of a prominent abdominal aortic pulse should prompt noninvasive imaging for better characterization. Femoral and/or popliteal artery aneurysms should be sought in patients with abdominal aortic aneurysm disease.

The level of a claudication-producing arterial obstruction can often be identified on physical examination (Fig. 267-3). For example, in a

patient with calf claudication, a decrease in pulse amplitude between the common femoral and popliteal arteries will localize the obstruction to the level of the superficial femoral artery, although inflow obstruction above the level of the common femoral artery may coexist. Auscultation for carotid, subclavian, abdominal aortic, and femoral artery bruits should be routine. However, the correlation between the presence of a bruit and the degree of vascular obstruction is poor. A cervical bruit is a weak indicator of the degree of carotid artery stenosis; the absence of a bruit does not exclude the presence of significant luminal obstruction. If a bruit extends into diastole or if a thrill is present, the obstruction is usually severe. Another cause of an arterial bruit is an arteriovenous fistula with enhanced flow.

The likelihood of significant lower extremity peripheral artery disease increases with typical symptoms of claudication, cool skin, abnormalities on pulse examination, or the presence of a vascular bruit. Abnormal pulse oximetry (a >2% difference between finger and toe oxygen saturation) can be used to detect lower extremity peripheral artery disease and is comparable in its performance characteristics to the ankle-brachial index.

Inspection and Palpation of the Heart The left ventricular apex beat may be visible in the midclavicular line at the fifth intercostal space in thin-chested adults. Visible pulsations anywhere other than this expected location are abnormal. The left anterior chest wall may heave in patients with an enlarged or hyperdynamic left or right ventricle. As noted previously, a visible right upper parasternal pulsation may be suggestive of ascending aortic aneurysm disease. In thin, tall patients and patients with advanced obstructive lung disease and flattened

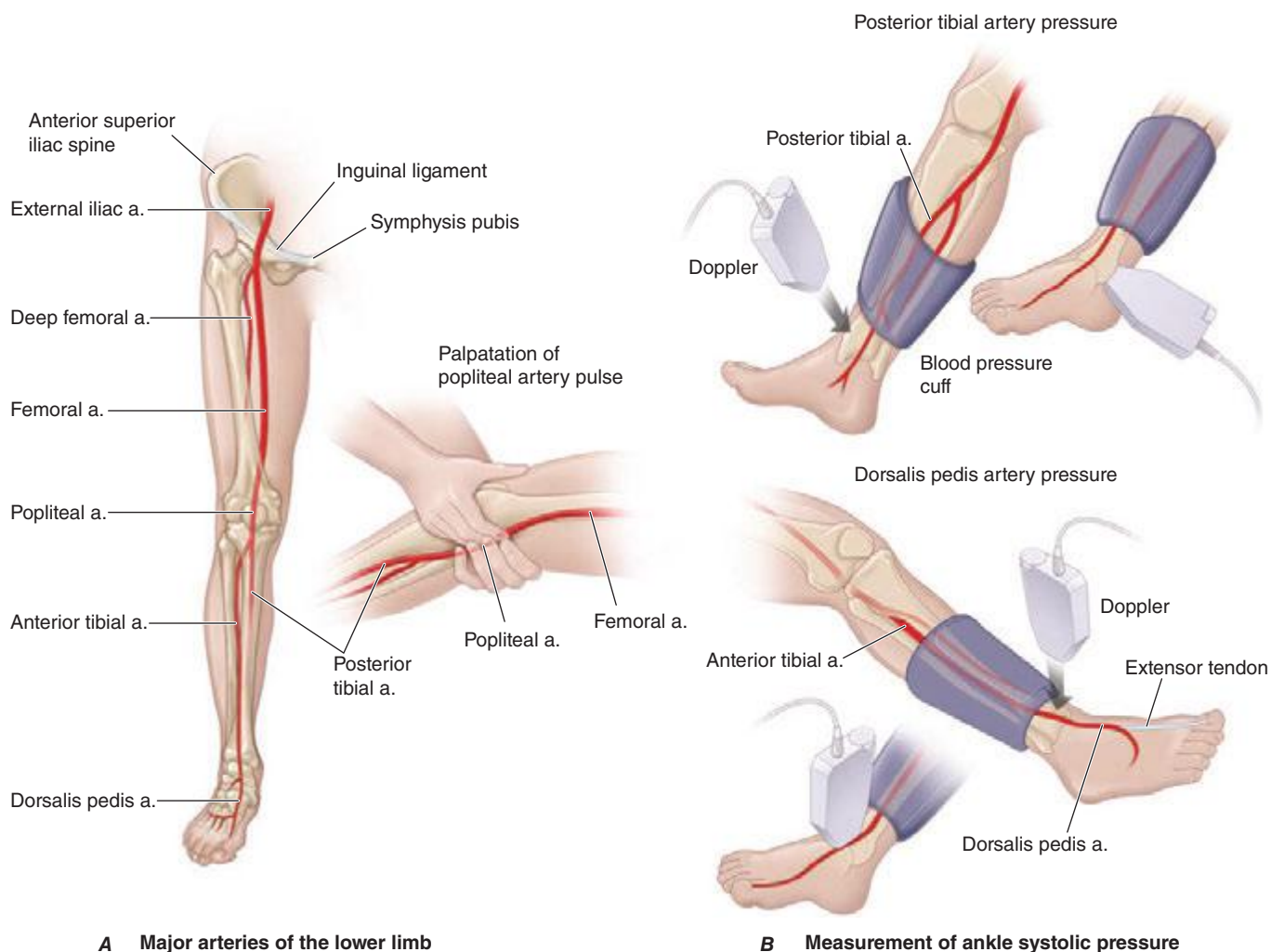


FIGURE 267-3 A. Anatomy of the major arteries of the leg. B. Measurement of the ankle systolic pressure. (From NA Khan et al: JAMA 295:536, 2006.)

diaphragms, the cardiac impulse may be visible in the epigastrium and should be distinguished from a pulsatile liver edge.

Palpation of the heart begins with the patient in the supine position at 30° and can be enhanced by placing the patient in the left lateral decubitus position. The normal left ventricular impulse is less than 2 cm in diameter and moves quickly away from the fingers; it is better appreciated at end expiration, with the heart closer to the anterior chest wall. Characteristics such as size, amplitude, and rate of force development should be noted.

Enlargement of the left ventricular cavity is manifested by a leftward and downward displacement of an enlarged apex beat. A sustained apex beat is a sign of pressure overload, such as that which may be present in patients with AS or chronic hypertension. A palpable presystolic impulse corresponds to the fourth heart sound (S_4) and is indicative of reduced left ventricular compliance and the forceful contribution of atrial contraction to ventricular filling. A palpable third sound (S_3), which is indicative of a rapid early filling wave in patients with heart failure, may be present even when the gallop itself is not audible. A large left ventricular aneurysm may sometimes be palpable as an ectopic impulse, discrete from the apex beat. HOCM may very rarely cause a triple cadence beat at the apex with contributions from a palpable S_4 and the two components of the bisferiens systolic pulse.

Right ventricular pressure or volume overload may create a lateral lift. Signs of either TR (*cv* waves in the jugular venous pulse) and/or pulmonary arterial hypertension (a loud single or palpable P_2) would be confirmatory. The right ventricle can enlarge to the extent that left-sided events cannot be appreciated. A zone of retraction between the right and left ventricular impulses sometimes can be appreciated in patients with right ventricle pressure or volume overload when they are placed in the left lateral decubitus position. Systolic and diastolic thrills signify turbulent and high-velocity blood flow. Their locations help identify the origin of heart murmurs.

CARDIAC AUSCULTATION

Heart Sounds Ventricular systole is defined by the interval between the first (S_1) and second (S_2) heart sounds (Fig. 267-4). The first heart sound (S_1) includes mitral and tricuspid valve closure. Normal splitting can be appreciated in young patients and those with right bundle branch block, in whom tricuspid valve closure is relatively delayed. The intensity of S_1 is determined by the distance over which the anterior leaflet of the mitral valve must travel to return to its annular plane, leaflet mobility, left ventricular contractility, and the PR interval. S_1 is classically loud in the early phases of rheumatic mitral stenosis (MS) and in patients with hyperkinetic circulatory states or short PR intervals. S_1 becomes softer in the later stages of MS when the leaflets are rigid and calcified, after exposure to β -adrenergic receptor blockers, with long PR intervals, and with left ventricular contractile dysfunction. The intensity of heart sounds, however, can be reduced by any process that increases the distance between the stethoscope and the responsible cardiac event, including mechanical ventilation, obstructive lung disease, obesity, pneumothorax, and a pericardial effusion.

Aortic and pulmonic valve closure constitutes the second heart sound (S_2). With normal or physiologic splitting, the A_2 - P_2 interval increases with inspiration and narrows during expiration. This physiologic interval will widen with right bundle branch block because of the further delay in pulmonic valve closure and in patients with severe MR because of the premature closure of the aortic valve. An unusually narrowly split or even a singular S_2 is a feature of pulmonary arterial hypertension. Fixed splitting of S_2 , in which the A_2 - P_2 interval is wide and does not change during the respiratory cycle, occurs in patients with a secundum atrial septal defect. Reversed or paradoxical splitting refers to a pathologic delay in aortic valve closure, such as that which occurs in patients with left bundle branch block, right ventricular pacing, severe AS, HOCM, and acute myocardial ischemia. With reversed or paradoxical splitting, the individual components of S_2 are audible at end expiration, and their interval narrows with inspiration, the opposite of what would be expected under normal physiologic conditions. P_2 is considered loud when its intensity exceeds that of A_2 at the base, when it can be palpated in the area of the proximal main pulmonary

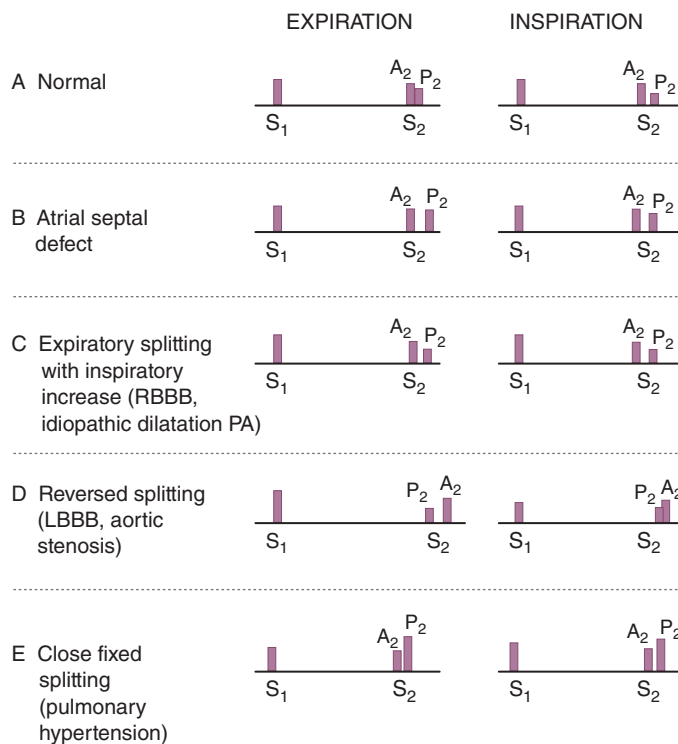


FIGURE 267-4 Heart sounds. **A.** Normal. S_1 , first heart sound; S_2 , second heart sound; A_2 , aortic component of the second heart sound; P_2 , pulmonic component of the second heart sound. **B.** Atrial septal defect with fixed splitting of S_2 . **C.** Physiologic but wide splitting of S_2 with right bundle branch block (RBBB). PA, pulmonary artery. **D.** Reversed or paradoxical splitting of S_2 with left bundle branch block (LBBB). **E.** Narrow splitting of S_2 with pulmonary hypertension. (From NO Fowler: *Diagnosis of Heart Disease*. New York, Springer-Verlag, 1991, p 31.)

artery (second left interspace), or when both components of S_2 can be appreciated at the lower left sternal border or apex. The intensity of A_2 and P_2 decreases with aortic and pulmonic stenosis, respectively. In these conditions, a single S_2 may result.

Systolic Sounds An ejection sound is a high-pitched early systolic sound that corresponds in timing to the upstroke of the carotid pulse. It usually is associated with congenital bicuspid aortic or pulmonic valve disease; however, ejection sounds are also sometimes audible in patients with isolated aortic or pulmonary root dilation and normal semilunar valves. The ejection sound that accompanies bicuspid aortic valve disease becomes softer and then inaudible as the valve calcifies and becomes more rigid. The ejection sound that accompanies pulmonic stenosis (PS) moves closer to the first heart sound as the severity of the stenosis increases. In addition, the pulmonic ejection sound is the only right-sided acoustic event that decreases in intensity with inspiration. Ejection sounds are often heard more easily at the lower left sternal border than they are at the base. Nonejection sounds (clicks), which occur after the onset of the carotid upstroke, are related to MVP and may be single or multiple. The nonejection click may introduce a murmur. This click-murmur complex will move away from the first heart sound with maneuvers that increase ventricular preload, such as squatting. On standing, the click and murmur move closer to S_1 .

Diastolic Sounds The high-pitched opening snap (OS) of MS occurs after a very short interval after the second heart sound. The A_2 -OS interval is inversely proportional to the height of the left atrial-left ventricular diastolic pressure gradient. The intensity of both S_1 and the OS of MS decreases with progressive calcification and rigidity of the anterior mitral leaflets. The pericardial knock (PK) is also high-pitched

and occurs slightly later than the OS, corresponding in timing to the abrupt cessation of ventricular expansion after tricuspid valve opening and to an exaggerated γ descent seen in the jugular venous waveform in patients with constrictive pericarditis. A tumor plop is a lower-pitched sound that rarely can be heard in patients with atrial myxoma. It may be appreciated only in certain positions and arises from the diastolic prolapse of the tumor across the mitral valve.

The third heart sound (S_3) occurs during the rapid filling phase of ventricular diastole. It can be a normal finding in children, adolescents, and young adults; however, in older patients, it signifies heart failure. A left-sided S_3 is a low-pitched sound best heard over the left ventricular (LV) apex. A right-sided S_3 is usually better heard over the lower left sternal border and becomes louder with inspiration. A left-sided S_3 in patients with chronic heart failure is predictive of cardiovascular morbidity and mortality. Interestingly, an S_3 is equally prevalent among heart failure patients with and without LV systolic dysfunction.

The fourth heart sound (S_4) occurs during the atrial filling phase of ventricular diastole and indicates LV presystolic expansion. An S_4 is more common among patients who derive significant benefit from the atrial contribution to ventricular filling, such as those with chronic LV hypertrophy or active myocardial ischemia. An S_4 is not present with atrial fibrillation.

Cardiac Murmurs Heart murmurs result from audible vibrations that are caused by increased turbulence and are defined by their timing within the cardiac cycle. Not all murmurs are indicative of structural heart disease, and the accurate identification of a benign or functional systolic murmur often can obviate the need for additional testing in healthy subjects. The duration, frequency, configuration, and intensity of a heart murmur are dictated by the magnitude, variability, and duration of the responsible pressure difference between two cardiac chambers, the two ventricles, or the ventricles and their respective great arteries. The intensity of a heart murmur is graded on a scale of 1 to 6; a thrill is present with murmurs of grade 4 or greater intensity. Other attributes of the murmur that aid in its accurate identification include its location, radiation, and response to bedside maneuvers. Although clinicians can detect and correctly identify heart murmurs with only fair reliability, a careful and complete bedside examination usually can identify individuals with valvular heart disease for whom transthoracic echocardiography and clinical follow-up are indicated and exclude subjects for whom no further evaluation is necessary.

Systolic murmurs can be early, mid, late, or holosystolic in timing (Fig. 267-5). Acute severe MR results in a decrescendo early systolic murmur, the characteristics of which are related to the progressive attenuation of the left ventricular to left atrial pressure gradient during systole because of the steep and rapid rise in left atrial pressure in this context. Severe MR associated with posterior leaflet prolapse or flail radiates anteriorly and to the base, where it can be confused with the murmur of AS. MR that is due to anterior leaflet involvement radiates posteriorly and to the axilla. With acute TR in patients with normal pulmonary artery pressures, an early systolic murmur that may increase in intensity with inspiration may be heard at the left lower sternal border, with regurgitant *cv* waves visible in the jugular venous pulse.

A midsystolic murmur begins after S_1 and ends before S_2 ; it is typically crescendo-decrescendo in configuration. AS is the most common cause of a midsystolic murmur in an adult. It is often difficult to estimate the severity of the valve lesion on the basis of the physical examination findings, especially in older hypertensive patients with stiffened carotid arteries or patients with low cardiac output in whom the intensity of the systolic heart murmur is misleadingly soft. Examination findings consistent with severe AS would include *parvus et tardus* carotid upstrokes, a late-peaking grade 3 or greater midsystolic murmur, a soft A_2 , a sustained LV apical impulse, and an S_4 . It is sometimes difficult to distinguish aortic sclerosis from more advanced degrees of valve stenosis. The former is defined by focal thickening and calcification of the aortic valve leaflets that is not severe enough to result in obstruction. These valve changes are associated with a Doppler jet velocity across the aortic valve of 2.5 m/s or less. Patients

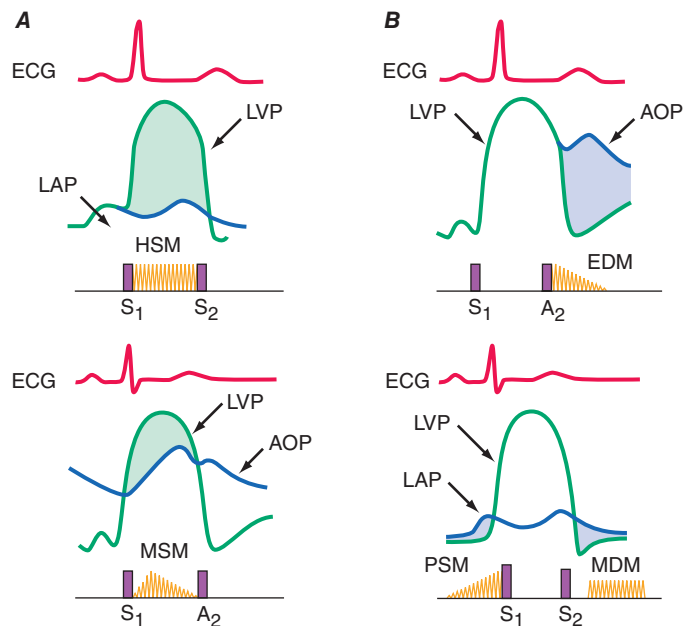


FIGURE 267-5 **A. Top.** Graphic representation of the systolic pressure difference (green shaded area) between left ventricle and left atrium with phonocardiographic recording of a holosystolic murmur (HSM) indicative of mitral regurgitation. ECG, electrocardiogram; LAP, left atrial pressure; LVP, left ventricular pressure; S_1 , first heart sound; S_2 , second heart sound. **Bottom.** Graphic representation of the systolic pressure gradient (green shaded area) between left ventricle and aorta in patient with aortic stenosis. A midsystolic murmur (MSM) with a crescendo-decrescendo configuration is recorded. AOP, aortic pressure. **B. Top.** Graphic representation of the diastolic pressure difference between the aorta and left ventricle (blue shaded area) in a patient with aortic regurgitation, resulting in a decrescendo, early diastolic murmur (EDM) beginning with A_2 . **Bottom.** Graphic representation of the diastolic left atrial–left ventricular gradient (blue areas) in a patient with mitral stenosis with a mid-diastolic murmur (MDM) and late presystolic murmurs (PSM).

with aortic sclerosis can have grade 2 or 3 midsystolic murmurs identical in their acoustic characteristics to the murmurs heard in patients with more advanced degrees of AS. Other causes of a midsystolic heart murmur include pulmonic valve stenosis (with or without an ejection sound), HOCM, increased pulmonary blood flow in patients with a large atrial septal defect and left-to-right shunting, and several states associated with accelerated blood flow in the absence of structural heart disease, such as fever, thyrotoxicosis, pregnancy, anemia, and normal childhood/adolescence.

The murmur of HOCM has features of both obstruction to LV outflow and MR, as would be expected from knowledge of the pathophysiology of this condition. The systolic murmur of HOCM usually can be distinguished from other causes on the basis of its response to bedside maneuvers, including Valsalva, passive leg raising, and standing/squatting. In general, maneuvers that decrease LV preload (or increase LV contractility) will cause the murmur to intensify, whereas maneuvers that increase LV preload or afterload will cause a decrease in the intensity of the murmur. Accordingly, the systolic murmur of HOCM becomes louder during the strain phase of the Valsalva maneuver and after standing quickly from a squatting position. The murmur becomes softer with passive leg raising and when squatting. The murmur of AS is typically loudest in the second right interspace with radiation into the carotids, whereas the murmur of HOCM is best heard between the lower left sternal border and the apex. The murmur of PS is best heard in the second left interspace. The midsystolic murmur associated with enhanced pulmonic blood flow in the setting of a large atrial septal defect (ASD) is usually loudest at the mid-left sternal border.

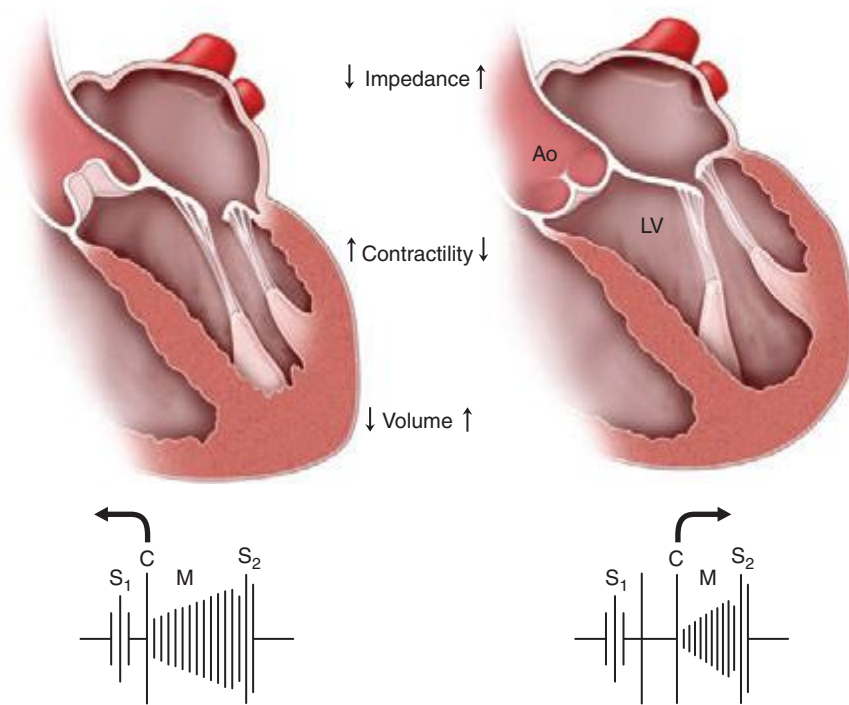


FIGURE 267-6 Behavior of the click (C) and murmur (M) of mitral valve prolapse with changes in loading (volume, impedance) and contractility. S_1 , first heart sound; S_2 , second heart sound. With standing (left side of figure), volume and impedance decrease, as a result of which the click and murmur move closer to S_1 . With squatting (right), the click and murmur move away from S_1 due to the increases in left ventricular volume and impedance (afterload). Ao, aorta; LV, left ventricle. (Adapted from RA O'Rourke, MH Crawford: *Curr Prob Cardiol* 1:9, 1976.)

A late systolic murmur, heard best at the apex, indicates MVP. As previously noted, the murmur may or may not be introduced by a nonejection click. Differential radiation of the murmur, as previously described, may help identify the specific leaflet involved by the myxomatous process. The click-murmur complex behaves in a manner directionally similar to that demonstrated by the murmur of HOCM during the Valsalva and stand/squat maneuvers (Fig. 267-6). The murmur of MVP can be identified by the accompanying nonejection click.

Holosystolic murmurs are plateau in configuration and reflect a continuous and wide pressure gradient between the left ventricle and left atrium with chronic MR, the left ventricle and right ventricle with a ventricular septal defect (VSD), and the right ventricle and right atrium with TR. In contrast to acute MR, in chronic MR the left atrium is enlarged and its compliance is normal or increased to the extent that there is little if any further increase in left atrial pressure from any increase in regurgitant volume. The murmur of MR is best heard over the cardiac apex. The intensity of the murmur increases with maneuvers that increase LV afterload, such as sustained hand grip. The murmur of a VSD (without significant pulmonary hypertension) is holosystolic and loudest at the mid-left sternal border, where a thrill is usually present. The murmur of TR is loudest at the lower left sternal border, increases in intensity with inspiration (Carvallo's sign), and is accompanied by visible *cv* waves in the jugular venous wave form and, on occasion, by pulsatile hepatomegaly.

Diastolic Murmurs In contrast to some systolic murmurs, diastolic heart murmurs always signify structural heart disease (Fig. 267-5). The murmur associated with acute, severe AR is relatively soft and of short duration because of the rapid rise in LV diastolic pressure and the progressive diminution of the aortic-LV diastolic pressure gradient. In contrast, the murmur of chronic severe AR is classically heard as a decrescendo, blowing diastolic murmur along the left sternal border in patients with primary valve pathology and sometimes along the right sternal border in patients with primary aortic root pathology.

With chronic AR, the pulse pressure is wide and the arterial pulses are bounding in character. These signs of significant diastolic run-off are absent in the acute phase. The murmur of pulmonic regurgitation is also heard along the left sternal border. It is most commonly due to pulmonary hypertension and enlargement of the annulus of the pulmonic valve. S_2 is single and loud and may be palpable. There is a right ventricular/parasternal lift that is indicative of chronic right ventricular pressure overload. A less impressive murmur of PR is present after repair of tetralogy of Fallot or pulmonic valve atresia. In this postoperative setting, the murmur is softer and lower-pitched, and the severity of the accompanying pulmonic regurgitation can be underestimated significantly.

MS is the classic cause of a mid- to late diastolic murmur, which is best heard over the apex in the left lateral decubitus position, is low-pitched or rumbling, and is introduced by an OS in the early stages of the rheumatic disease process. Presystolic accentuation refers to an increase in the intensity of the murmur just before the first heart sound and occurs in patients with sinus rhythm. It is absent in patients with atrial fibrillation. The auscultatory findings in patients with rheumatic tricuspid stenosis typically are obscured by left-sided events, although they are similar in nature to those described in patients with MS. "Functional" mitral or tricuspid stenosis refers to the generation of mid-diastolic murmurs that are created by increased and accelerated transvalvular diastolic flow, even in the absence

of valvular obstruction, in the setting of severe MR, severe TR, or a large ASD with left-to-right shunting. The Austin Flint murmur of chronic severe AR is a low-pitched mid- to late apical diastolic murmur that sometimes can be confused with MS. The Austin Flint murmur typically decreases in intensity after exposure to vasodilators, whereas the murmur of MS may be accompanied by an opening snap and also may increase in intensity after vasodilators because of the associated increase in cardiac output. Unusual causes of a mid-diastolic murmur include atrial myxoma, complete heart block, and acute rheumatic mitral valvulitis.

Continuous Murmur A continuous murmur is predicated on a pressure gradient that persists between two cardiac chambers or blood vessels across systole and diastole. The murmurs typically begin in systole, envelop the second heart sound (S_2), and continue through some portion of diastole. They can often be difficult to distinguish from individual systolic and diastolic murmurs in patients with mixed valvular heart disease. The classic example of a continuous murmur is that associated with a PDA, which usually is heard in the second or third interspace at a slight distance from the sternal border. Other causes of a continuous murmur include a ruptured sinus of Valsalva aneurysm with creation of an aortic-right atrial or right ventricular fistula, a coronary or great vessel arteriovenous fistula, and an arteriovenous fistula constructed to provide dialysis access. There are two types of benign continuous murmurs. The cervical venous hum is heard in children or adolescents in the supraclavicular fossa. It can be obliterated with firm pressure applied to the diaphragm of the stethoscope, especially when the subject turns his or her head toward the examiner. The mammary soufflé of pregnancy relates to enhanced arterial blood flow through engorged breasts. The diastolic component of the murmur can be obliterated with firm pressure over the stethoscope.

Dynamic Auscultation Diagnostic accuracy can be enhanced by the performance of simple bedside maneuvers to identify heart murmurs and characterize their significance (Table 267-1). Except for the pulmonic

TABLE 267-1 EFFECTS OF PHYSIOLOGIC AND PHARMACOLOGIC INTERVENTIONS ON THE INTENSITY OF HEART MURMURS AND SOUNDS

Respiration Right-sided murmurs and sounds generally increase with inspiration, except for the PES. Left-sided murmurs and sounds are usually louder during expiration.

Valsalva maneuver Most murmurs decrease in length and intensity. Two exceptions are the systolic murmur of HOCM, which usually becomes much louder, and that of MVP, which becomes longer and often louder. After release of the Valsalva maneuver, right-sided murmurs tend to return to control intensity earlier than do left-sided murmurs.

After VPB or AF Murmurs originating at normal or stenotic semilunar valves increase in the cardiac cycle after a VPB or in the cycle after a long cycle length in AF. By contrast, systolic murmurs due to AV valve regurgitation do not change, diminish (papillary muscle dysfunction), or become shorter (MVP).

Positional changes With *standing*, most murmurs diminish, with two exceptions being the murmur of HOCM, which becomes louder, and that of MVP, which lengthens and often is intensified. With *squatting*, most murmurs become louder, but those of HOCM and MVP usually soften and may disappear. Passive leg raising usually produces the same results.

Exercise Murmurs due to blood flow across normal or obstructed valves (e.g., PS, MS) become louder with both isotonic and submaximal isometric (hand grip) exercise. Murmurs of MR, VSD, and AR also increase with hand grip exercise. However, the murmur of HOCM often decreases with nearly maximum hand grip exercise. Left-sided S_4 and S_3 sounds are often accentuated by exercise, particularly when due to ischemic heart disease.

Abbreviations: AF, atrial fibrillation; AR, aortic regurgitation; HOCM, hypertrophic obstructive cardiomyopathy; MR, mitral regurgitation; MS, mitral stenosis; MVP, mitral valve prolapse; PES, pulmonic ejection sound; PR, pulmonic regurgitation; PS, pulmonic stenosis; TR, tricuspid regurgitation; TS, tricuspid stenosis; VPB, ventricular premature beat; VSD, ventricular septal defect.

ejection sound, right-sided events increase in intensity with inspiration and decrease with expiration; left-sided events behave oppositely (100% sensitivity, 88% specificity). As previously noted, the intensity of the murmurs associated with MR, VSD, and AR will increase in response to maneuvers that increase LV afterload, such as hand grip and vasopressors. The intensity of these murmurs will decrease after exposure to vasodilating agents. Squatting is associated with an abrupt increase in LV preload and afterload, whereas rapid standing results in a sudden decrease in preload. In patients with MVP, the click and murmur move away from the first heart sound with squatting because of the delay in onset of leaflet prolapse at higher ventricular volumes. With rapid standing, however, the click and murmur move closer to the first heart sound as prolapse occurs earlier in systole at a smaller chamber dimension. The murmur of HOCM behaves similarly, becoming softer and shorter with squatting (95% sensitivity, 85% specificity) and longer and louder on rapid standing (95% sensitivity, 84% specificity). A change in the intensity of a systolic murmur in the first beat after a premature beat or in the beat after a long cycle length in patients with atrial fibrillation suggests valvular AS rather than MR, particularly in an older patient in whom the murmur of the AS may be well transmitted to the apex (Gallavardin effect). Of note, however, the systolic murmur of HOCM also increases in intensity in the beat after a premature beat. This increase in intensity of any LV outflow murmur in the beat after a premature beat relates to the combined effects of enhanced LV filling (from the longer diastolic period) and postextrasystolic potentiation of LV contractile function. In either instance, forward flow will accelerate, causing an increase in the gradient across the LV outflow tract (dynamic or fixed) and a louder systolic murmur. In contrast, the intensity of the murmur of MR does not change in a postpremature beat, because there is relatively little change in the nearly constant LV to left atrial pressure gradient or further alteration in mitral valve flow. Bedside exercise can sometimes be performed to increase cardiac output and, secondarily, the intensity of both systolic and diastolic heart murmurs. Most left-sided heart murmurs decrease in intensity and duration during the strain phase of the Valsalva maneuver. The murmurs associated with MVP and HOCM are the two notable exceptions. The Valsalva maneuver also can be used to assess the integrity of the heart and vasculature in the setting of advanced heart failure.

Prosthetic Heart Valves The first clue that prosthetic valve dysfunction may contribute to recurrent symptoms is frequently a change in the quality of the heart sounds or the appearance of a new murmur. The heart sounds with a bioprosthetic valve resemble those generated by native valves. A mitral bioprosthesis usually is associated with a grade 2 or 3 midsystolic murmur along the left sternal border (created by turbulence across the valve struts as they project into the LV outflow tract) as well as by a soft mid-diastolic murmur that occurs with normal LV filling. This diastolic murmur often can be heard only in the left lateral decubitus position and after exercise. A high pitched or holosystolic apical murmur is indicative of pathologic MR due to a paravalvular leak and/or intra-annular bioprosthetic regurgitation from leaflet degeneration, for which additional imaging is usually indicated. Clinical deterioration can occur rapidly after the first expression of mitral bioprosthetic failure. A tissue valve in the aortic position is always associated with a grade 2 to 3 midsystolic murmur at the base or just below the suprasternal notch. A diastolic murmur of AR is abnormal in any circumstance. Mechanical valve dysfunction may first be suggested by a decrease in the intensity of either the opening or the closing sound. A high-pitched apical systolic murmur in patients with a mechanical mitral prosthesis and a diastolic decrescendo murmur in patients with a mechanical aortic prosthesis indicate paravalvular regurgitation. Patients with prosthetic valve thrombosis may present clinically with signs of shock, muffled heart sounds, and soft murmurs.

Pericardial Disease A pericardial friction rub is nearly 100% specific for the diagnosis of acute pericarditis, although the sensitivity of this finding is not nearly as high, because the rub may come and go over the course of an acute illness or be very difficult to elicit. The rub is heard as a leathery or scratchy three-component or two-component sound, although it may be monophasic. Classically, the three components are ventricular systole, rapid early diastolic filling, and late presystolic filling after atrial contraction in patients in sinus rhythm. It is necessary to listen to the heart in several positions. Additional clues may be present from the history and 12-lead electrocardiogram. The rub typically disappears as the volume of any pericardial effusion increases. Pericardial tamponade can be diagnosed with a sensitivity of 98%, a specificity of 83%, and a positive likelihood ratio of 5.9 (95% confidence interval 2.4–14) by a pulsus paradoxus that exceeds 12 mmHg in a patient with a large pericardial effusion.

The findings on physical examination are integrated with the symptoms previously elicited with a careful history to construct an appropriate differential diagnosis and proceed with indicated imaging and laboratory assessment. The physical examination is an irreplaceable component of the diagnostic algorithm and in selected patients can inform prognosis. Educational efforts to improve clinician competence eventually may result in cost saving, particularly if the indications for imaging can be influenced by the examination findings.

268 Electrocardiography

Ary L. Goldberger

An electrocardiogram (ECG or EKG) is a graphic recording of electric potentials generated by the heart. The signals are detected by means of metal electrodes attached to the extremities and chest wall and then are amplified and recorded by the electrocardiograph. ECG *leads* actually display the instantaneous *differences* in potential between the electrodes.

The clinical utility of the ECG derives from its immediate availability as a noninvasive, inexpensive, and highly versatile test. In addition to its use in detecting arrhythmias, conduction disturbances, and myocardial ischemia, electrocardiography may reveal findings related to life-threatening metabolic disturbances (e.g., hyperkalemia) or increased susceptibility to sudden cardiac death (e.g., QT prolongation syndromes).

ELECTROPHYSIOLOGY

(See also Chaps. 274 and 276) Depolarization of the heart is the initiating event for cardiac contraction. The electric currents that spread through the heart are produced by three components: cardiac pacemaker cells, specialized conduction tissue, and the heart muscle itself. The ECG, however, records only the depolarization (stimulation) and repolarization (recovery) potentials generated by the “working” atrial and ventricular myocardium.

The depolarization stimulus for the normal heartbeat originates in the *sinoatrial (SA) node* (Fig. 268-1), or *sinus node*, a collection of *pacemaker cells*. These cells fire spontaneously; that is, they exhibit *automaticity*. The first phase of cardiac electrical activation is the spread of the depolarization wave through the right and left atria, followed by atrial contraction. Next, the impulse stimulates pacemaker and specialized conduction tissues in the atrioventricular (AV) nodal and His-bundle areas; together, these two regions constitute the AV junction. The bundle of His bifurcates into two main branches, the right and left bundles, which rapidly transmit depolarization wavefronts to the right and left ventricular myocardium by way of Purkinje fibers. The main left bundle bifurcates into two primary subdivisions: a left anterior fascicle and a left posterior fascicle. The depolarization wavefronts then spread through the ventricular wall, from endocardium to epicardium, triggering ventricular contraction.

Since the cardiac depolarization and repolarization waves have direction and magnitude, they can be represented by vectors. Vector analysis illustrates a central concept of electrocardiography: The ECG records the complex spatial and temporal summation of electrical potentials from multiple myocardial fibers conducted to the surface of the body. This principle accounts for inherent limitations in both ECG *sensitivity* (activity from certain cardiac regions may be canceled out or may be too weak to be recorded) and *specificity* (the same vectorial sum can result from either a selective gain or a loss of forces in opposite directions).

ECG WAVEFORMS AND INTERVALS

The ECG waveforms are labeled alphabetically, beginning with the P wave, which represents atrial depolarization (Fig. 268-2). The QRS complex represents ventricular depolarization, and the ST-T-U complex (ST segment, T wave, and U wave) represents ventricular repolarization. The J point is the junction between the end of the QRS complex and the beginning of the ST segment. Atrial repolarization (ST_a and T_a) is usually too low in amplitude to be detected, but it may become apparent in conditions such as acute pericarditis and atrial infarction.

The QRS-T waveforms of the surface ECG correspond in a general way with the different phases of simultaneously obtained ventricular *action potentials*, the intracellular recordings from single myocardial

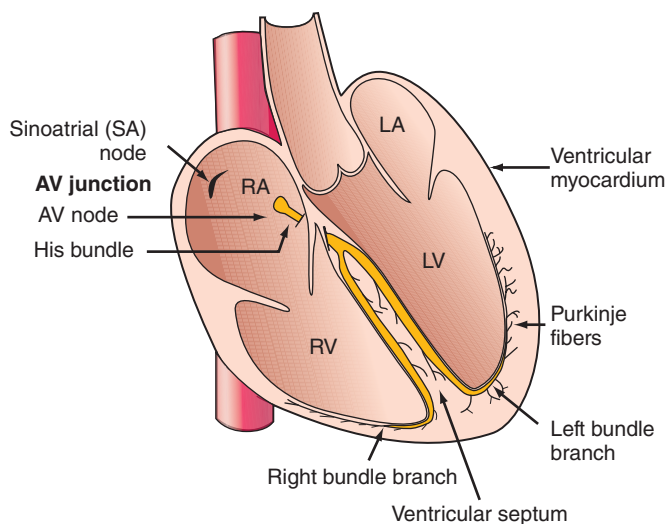


FIGURE 268-1 Schematic of the cardiac conduction system.

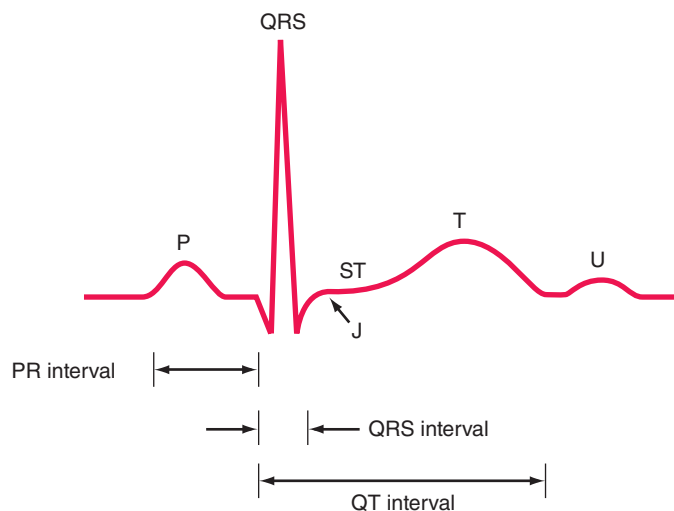


FIGURE 268-2 Basic ECG waveforms and intervals. Not shown is the RR interval, the time between consecutive QRS complexes.

fibers (Chap. 274). The rapid upstroke (phase 0) of the action potential corresponds to the onset of QRS. The plateau (phase 2) corresponds to the isoelectric ST segment, and active repolarization (phase 3) corresponds to the inscription of the T wave. Factors that decrease the slope of phase 0 by impairing the influx of Na^+ (e.g., hyperkalemia and drugs such as flecainide) tend to increase QRS duration. Conditions that prolong phase 2 (amiodarone, hypocalcemia) increase the QT interval. In contrast, shortening of ventricular repolarization (phase 2), such as by digitalis administration or hypercalcemia, abbreviates the ST segment.

The ECG ordinarily is recorded on special graph paper that is divided into 1-mm² gridlike boxes. Since the usual ECG paper speed is 25 mm/s, the smallest (1 mm) horizontal divisions correspond to 0.04 (40 ms), with heavier lines at intervals of 0.20 s (200 ms). Vertically, the ECG graph measures the amplitude of a specific wave or deflection (1 mV = 10 mm with standard calibration; the voltage criteria for hypertrophy mentioned below are given in millimeters). There are four major ECG intervals: RR, PR, QRS, and QT (Fig. 268-2). The heart rate (beats per minute) can be computed readily from the interbeat (RR) interval by dividing the number of large (0.20 s) time units between consecutive R waves into 300 or the number of small (0.04 s) units into 1500. The PR interval measures the time (normally 120–200 ms) between atrial and ventricular depolarization, which includes the physiologic delay imposed by stimulation of cells in the AV junction area. The QRS interval (normally 100–110 ms or less) reflects the duration of ventricular depolarization. The QT interval includes both ventricular depolarization and repolarization times and varies inversely with the heart rate. A rate-related (“corrected”) QT interval, QT_c , can be calculated as QT/\sqrt{RR} and normally is ≤ 0.44 s. (Some references give QT_c upper normal limits as 0.43 s in men and 0.45 s in women. Also, a number of different formulas have been proposed, without consensus, for calculating the QT_c .)

The QRS complex is subdivided into specific deflections or waves. If the initial QRS deflection in a particular lead is negative, it is termed a *Q wave*; the first positive deflection is termed an *R wave*. A negative deflection after an R wave is an *S wave*. Subsequent positive or negative waves are labeled R' and S' , respectively. Lowercase letters (qrs) are used for waves of relatively small amplitude. An entirely negative QRS complex is termed a *QS wave*.

ECG LEADS

The 12 conventional ECG leads record the difference in potential between electrodes placed on the surface of the body. These leads are divided into two groups: six limb (extremity) leads and six chest (precordial) leads. The limb leads record potentials transmitted onto the *frontal plane* (Fig. 268-3A), and the chest leads record potentials transmitted onto the *horizontal plane* (Fig. 268-3B).

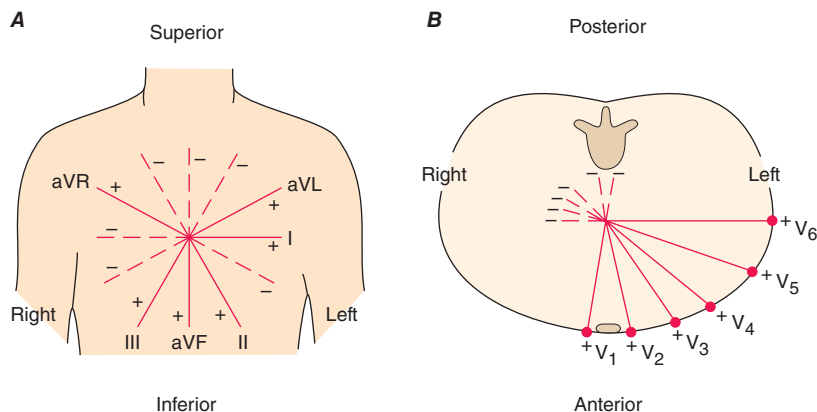


FIGURE 268-3 The six frontal plane (A) and six horizontal plane (B) leads provide a three-dimensional representation of cardiac electrical activity.

The spatial orientation and polarity of the six frontal plane leads is represented on the hexaxial diagram (Fig. 268-4). The six chest leads (Fig. 268-5) are unipolar recordings obtained by electrodes in the following positions: lead V_1 , fourth intercostal space, just to the right of the sternum; lead V_2 , fourth intercostal space, just to the left of the sternum; lead V_3 , midway between V_2 and V_4 ; lead V_4 , midclavicular line, fifth intercostal space; lead V_5 , anterior axillary line, same level as V_4 ; and lead V_6 , midaxillary line, same level as V_4 and V_5 . Additional posterior leads are sometimes placed on the same horizontal plane as V_4 to facilitate detection of acute posterolateral infarction (V_7 , midaxillary line; V_8 , posterior axillary line; and V_9 , posterior scapular line).

Together, the frontal and horizontal plane electrodes provide a three-dimensional representation of cardiac electrical activity. Each lead can be likened to a different video camera angle “looking” at the same events—atrial and ventricular depolarization and repolarization—from different spatial orientations. The conventional 12-lead ECG can be supplemented with additional leads in special circumstances. For example, right precordial leads V_3R , V_4R , etc., are useful in detecting evidence of acute right ventricular ischemia. Bedside monitors and

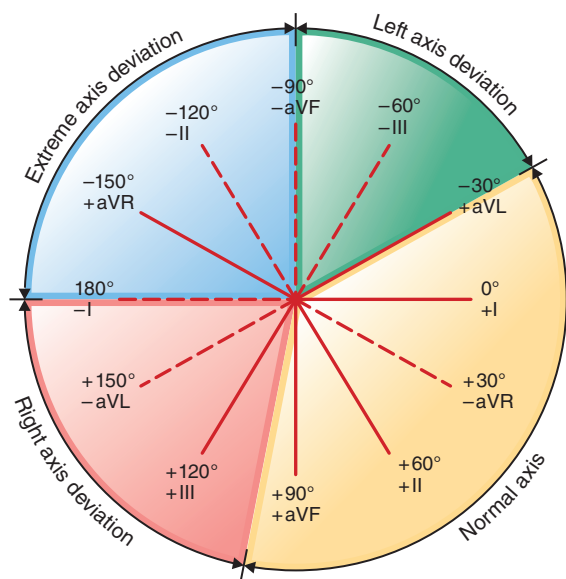


FIGURE 268-4 The frontal plane (limb or extremity) leads are represented on a hexaxial diagram. Each ECG lead has a specific spatial orientation and polarity. The positive pole of each lead axis (solid line) and the negative pole (hatched line) are designated by their angular position relative to the positive pole of lead I (0°). The mean electrical axis of the QRS complex is measured with respect to this display.

ambulatory ECG (Holter) recordings usually employ only one or two modified leads. **Intracardiac electrocardiography and electrophysiologic testing are discussed in Chaps. 274 and 276.**

The ECG leads are configured so that a positive (upright) deflection is recorded in a lead if a wave of depolarization spreads toward the positive pole of that lead, and a negative deflection is recorded if the wave spreads toward the negative pole. If the mean orientation of the depolarization vector is at right angles to a particular lead axis, a biphasic (equally positive and negative) deflection will be recorded.

GENESIS OF THE NORMAL ECG

P WAVE

The normal atrial depolarization vector is oriented downward and toward the subject's left, reflecting the spread of depolarization from the sinus node to the right and then the left atrial myocardium. Since this vector points toward the positive pole of lead II and toward the negative pole of lead aVR, the normal P wave will be positive in lead II and negative in lead aVR. By contrast, activation of the atria from an ectopic pacemaker in the lower part of either atrium or in the AV junction region may produce retrograde P waves (negative in lead II, positive in lead aVR). The normal P wave in lead V_1 may be biphasic with a positive component reflecting right atrial depolarization, followed by a small ($<1 \text{ mm}^2$) negative component reflecting left atrial depolarization.

QRS COMPLEX

Normal ventricular depolarization proceeds as a rapid, continuous spread of activation wave fronts. This complex process can be divided into two major sequential phases, and each phase can be represented by a mean vector (Fig. 268-6). The first phase is depolarization of the interventricular septum from the left to the right and anteriorly (vector 1). The second results from the simultaneous depolarization of the right and left ventricles; it normally is dominated by the more massive left ventricle, so that vector 2 points leftward and posteriorly. Therefore, a right precordial lead (V_1) will record this biphasic depolarization process with a small positive deflection (septal r wave) followed by a larger negative deflection (S wave). A left precordial lead, e.g., V_6 will record the same sequence with a small negative deflection (septal q wave) followed by a relatively tall positive deflection (R wave). Intermediate leads show a relative increase in R-wave amplitude (normal R-wave progression) and a decrease in S-wave amplitude progressing across the chest from right to left. The precordial lead where

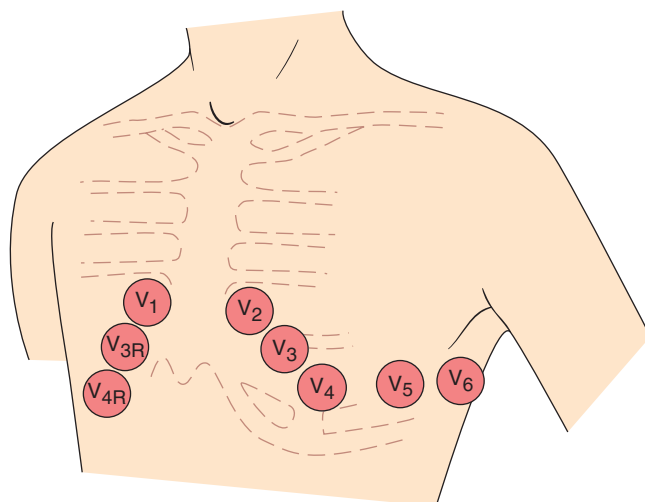


FIGURE 268-5 The horizontal plane (chest or precordial) leads are obtained with electrodes in the locations shown.

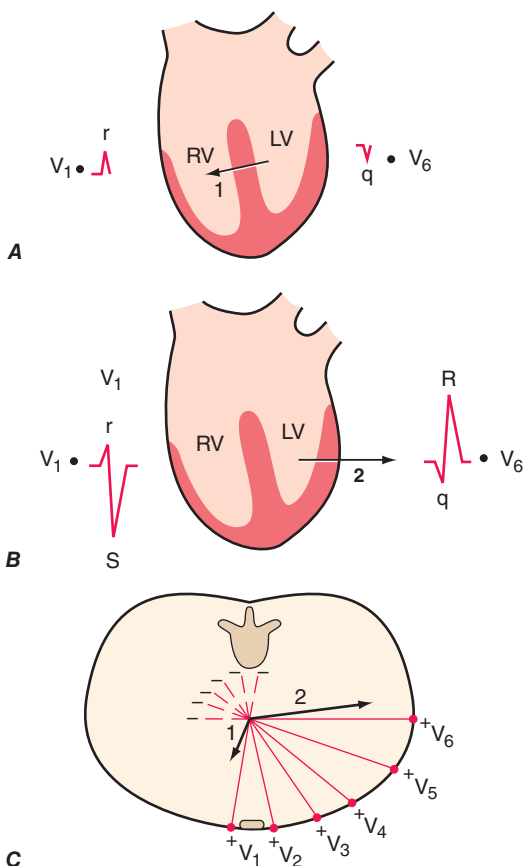


FIGURE 268-6 Ventricular depolarization can be divided into two major phases, each represented by a vector. **A.** The first phase (arrow 1) denotes depolarization of the ventricular septum, beginning on the left side and spreading to the right. This process is represented by a small “septal” r wave in lead V_1 and a small septal q wave in lead V_6 . **B.** Simultaneous depolarization of the left and right ventricles (LV and RV) constitutes the second phase. Vector 2 is oriented to the left and posteriorly, reflecting the electrical predominance of the LV. **C.** Vectors (arrows) representing these two phases are shown in reference to the horizontal plane leads. (After AL Goldberger et al: *Goldberger’s Clinical Electrocardiography: A Simplified Approach*, 8th ed. Philadelphia, Elsevier/Saunders, 2013.)

the R and S waves are of approximately equal amplitude is referred to as the *transition zone* (usually V_3 or V_4) (Fig. 268-7).

The QRS pattern in the extremity leads may vary considerably from one normal subject to another depending on the *electrical axis* of the QRS, which describes the mean orientation of the QRS vector with reference to the six frontal plane leads. Normally, the QRS axis ranges from -30° to $+100^\circ$ (Fig. 268-4). An axis more negative than -30° is referred to as *left axis deviation*, and an axis more positive than $+100^\circ$ is referred to as *right axis deviation*. Left axis deviation may occur as a normal variant but is more commonly associated with left ventricular hypertrophy, a block in the anterior fascicle of the left bundle system (left anterior fascicular block or hemiblock), or inferior myocardial infarction. Right axis deviation

also may occur as a normal variant (particularly in children and young adults), as a spurious finding due to reversal of the left and right arm electrodes, or in conditions such as right ventricular overload (acute or chronic), infarction of the lateral wall of the left ventricle, dextrocardia, left pneumothorax, and left posterior fascicular block.

T WAVE AND U WAVE

Normally, the mean T-wave vector is oriented roughly concordant with the mean QRS vector (within about 45° in the frontal plane). Since depolarization and repolarization are electrically opposite processes, this normal QRS–T-wave vector concordance indicates that repolarization normally must proceed in the reverse direction from depolarization (i.e., from ventricular epicardium to endocardium). The normal T wave is a small, rounded deflection (≤ 1 mm) that follows the T wave and usually has the same polarity as the T wave. An abnormal increase in U-wave amplitude is most commonly due to drugs (e.g., dofetilide, amiodarone, sotalol, quinidine) or hypokalemia. Very prominent U waves are a marker of increased susceptibility to the *torsades de pointes* type of ventricular tachycardia (Chap. 276). Inversion of the U wave in the precordial leads is abnormal and may be a subtle sign of ischemia.

MAJOR ECG ABNORMALITIES

CARDIAC ENLARGEMENT AND HYPERTROPHY

Right atrial overload (acute or chronic) may lead to an increase in P-wave amplitude (≥ 2.5 mm) (Fig. 268-8), sometimes referred to as “P-pulmonale.” Left atrial overload typically produces a biphasic P wave in V_1 with a broad negative component or a broad (≥ 120 ms), often notched P wave in one or more limb leads (Fig. 268-8). This pattern, previously referred to as “P-mitrale,” may also occur with left atrial conduction delays in the absence of actual atrial enlargement, leading to the more general designation of *left atrial abnormality*.

Right ventricular hypertrophy due to a sustained, severe pressure load (e.g., due to tight pulmonic valve stenosis or certain pulmonary artery hypertension syndromes) is characterized by a relatively tall R wave in lead V_1 ($R \geq S$ wave), usually with right axis deviation (Fig. 268-9); alternatively, there may be a qR pattern in V_1 or V_3R . ST depression and T-wave inversion in the right-to-midprecordial leads are also often present. This pattern, formerly called right ventricular “strain,” is attributed to repolarization abnormalities in acutely or chronically overloaded muscle. Prominent S waves may occur in the left lateral precordial leads. Right ventricular hypertrophy due to ostium secundum-type atrial septal defects, with the accompanying

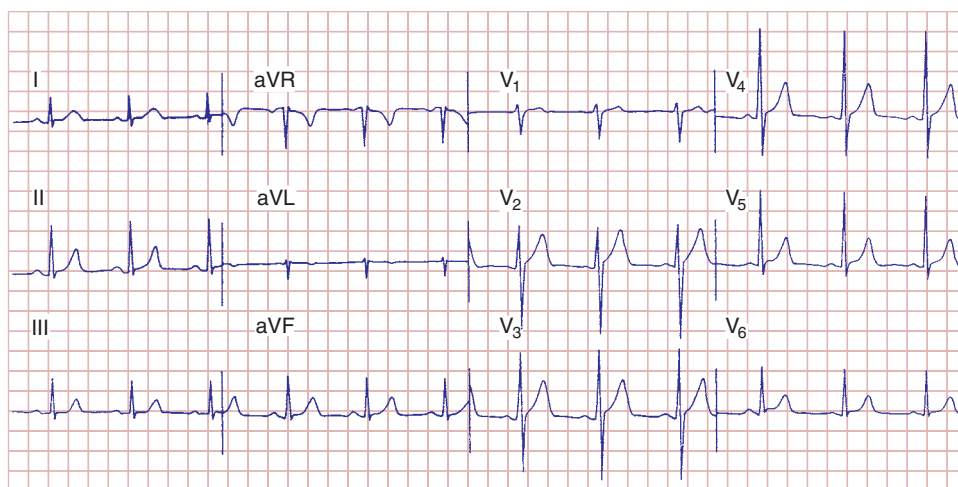


FIGURE 268-7 Normal electrocardiogram from a healthy subject. Sinus rhythm is present with a heart rate of 75 beats per minute. PR interval is 0.16 s; QRS interval (duration) is 0.08 s; QT interval is 0.36 s; QT_c is 0.40 s; the mean QRS axis is about $+70^\circ$. The precordial leads show normal R-wave progression with the transition zone (R wave = S wave) in lead V_3 .

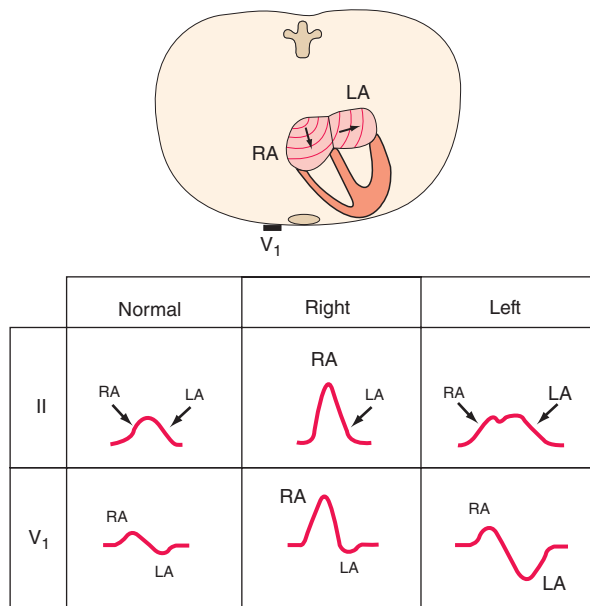


FIGURE 268-8 Right atrial (RA) overload may cause tall, peaked P waves in the limb or precordial leads. Left atrial (LA) abnormality may cause broad, often notched P waves in the limb leads and a biphasic P wave in lead V₁ with a prominent negative component representing delayed depolarization of the LA. (After MK Park, WG Guntheroth: *How to Read Pediatric ECGs*, 4th ed. St. Louis, Mosby/Elsevier, 2006.)

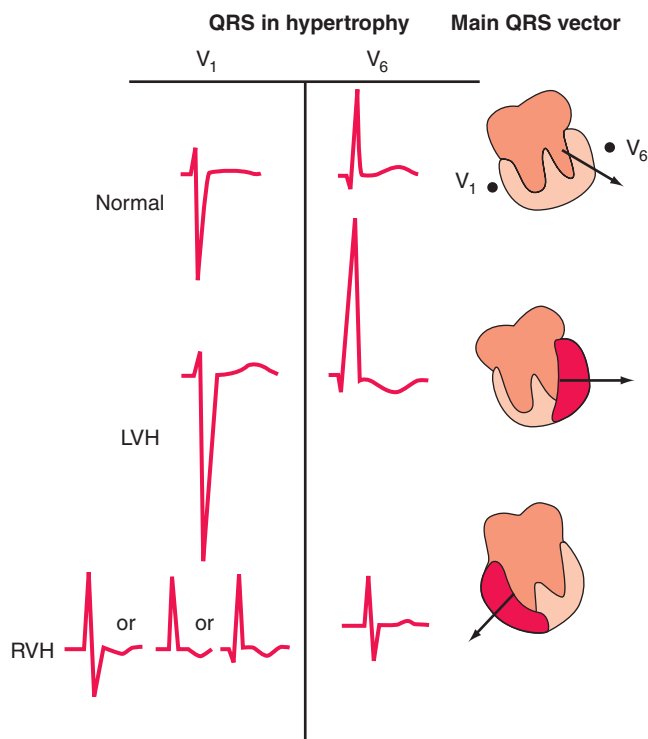


FIGURE 268-9 Left ventricular hypertrophy (LVH) increases the amplitude of electrical forces directed to the left and posteriorly. In addition, repolarization abnormalities may cause ST-segment depression and T-wave inversion in leads with a prominent R wave. Right ventricular hypertrophy (RVH) may shift the QRS vector to the right; this effect usually is associated with an R, RS, or qR complex in lead V₁. T-wave inversions may be present in right precordial leads.

right ventricular volume overload, is commonly associated with an incomplete or complete right bundle branch block pattern with a rightward QRS axis.

Acute cor pulmonale due to pulmonary embolism (Chap. 300), for example, may be associated with a normal ECG or a variety of abnormalities. Sinus tachycardia is the most common arrhythmia, although other tachyarrhythmias, such as atrial fibrillation or flutter, may occur. The QRS axis may shift to the right, sometimes in concert with the so-called S₁Q₃T₃ pattern (prominence of the S wave in lead I and the Q wave in lead III, with T-wave inversion in lead III). Acute right ventricular dilation also may be associated with slow R-wave progression and ST-T abnormalities in V₁ to V₄ simulating acute anterior infarction. A right ventricular conduction disturbance may appear.

Chronic cor pulmonale due to obstructive lung disease (Chap. 279) usually does not produce the classic ECG patterns of right ventricular hypertrophy noted above. Instead of tall right precordial R waves, chronic lung disease more typically is associated with small R waves in right-to-midprecordial leads (slow R-wave progression) due in part to downward displacement of the diaphragm and the heart. Low-voltage complexes are commonly present, owing to hyperaeration of the lungs.

A number of different voltage criteria for *left ventricular hypertrophy* (Fig. 268-9) have been proposed on the basis of the presence of tall left precordial R waves and deep right precordial S waves (e.g., SV₁ + [RV₅ or RV₆] >35 mm). Repolarization abnormalities (ST depression with T-wave inversions, formerly called the left ventricular “strain” pattern) also may appear in leads with prominent R waves. However, prominent precordial voltages may occur as a normal variant, especially in athletic or young individuals. Left ventricular hypertrophy may increase limb lead voltage with or without increased precordial voltage (e.g., RaVL + SV₃ >20 mm in women and >28 mm in men). The presence of left atrial abnormality increases the likelihood of underlying left ventricular hypertrophy in cases with borderline voltage criteria. Left ventricular hypertrophy often progresses to incomplete or complete left bundle branch block. The sensitivity of conventional voltage criteria for left ventricular hypertrophy is decreased in obese persons and smokers. ECG evidence for left ventricular hypertrophy is a major noninvasive marker of increased risk of cardiovascular morbidity and mortality rates, including sudden cardiac death. However, because of false-positive and false-negative diagnoses, the ECG is of limited utility in diagnosing atrial or ventricular enlargement. More definitive information is provided by echocardiography (Chap. 270e).

BUNDLE BRANCH BLOCKS AND RELATED PATTERNS

Intrinsic impairment of conduction in either the right or the left bundle system (intraventricular conduction disturbances) leads to prolongation of the QRS interval. With complete bundle branch blocks, the QRS interval is ≥120 ms in duration; with incomplete blocks, the QRS interval is between 100 and 120 ms. The QRS vector usually is oriented in the direction of the myocardial region where depolarization is delayed (Fig. 268-10). Thus, with right bundle branch block, the terminal QRS vector is oriented to the right and anteriorly (rSR' in V₁ and qRS in V₆, typically). Left bundle branch block alters both early and later phases of ventricular depolarization. The major QRS vector is directed to the left and posteriorly. In addition, the normal early left-to-right pattern of septal activation is disrupted such that septal depolarization proceeds from right to left as well. As a result, left bundle branch block generates wide, predominantly negative (QS) complexes in lead V₁ and entirely positive (R) complexes in lead V₆. A pattern identical to that of left bundle branch block, preceded by a sharp spike, is seen in most cases of electronic right ventricular pacing because of the relative delay in left ventricular activation.

Bundle branch block may occur in a variety of conditions. In subjects without structural heart disease, right bundle branch block is seen more commonly than left bundle branch block. Right bundle branch block also occurs with heart disease, both congenital (e.g., atrial septal defect) and acquired (e.g., valvular, ischemic). Left bundle branch block is often a marker of one of four underlying conditions associated with increased risk of cardiovascular morbidity and mortality rates: coronary heart disease (frequently with impaired left ventricular

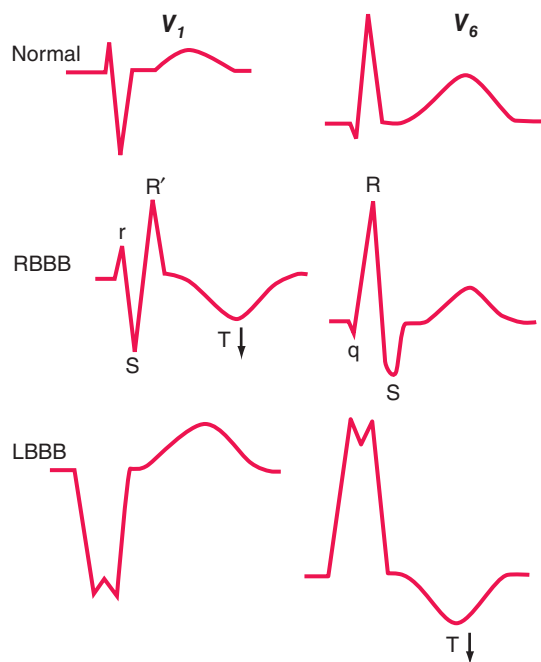


FIGURE 268-10 Comparison of typical QRS-T patterns in right bundle branch block (RBBB) and left bundle branch block (LBBB) with the normal pattern in leads V_1 and V_6 . Note the secondary T-wave inversions (arrows) in leads with an rS complex with RBBB and in leads with a wide R wave with LBBB.

function), hypertensive heart disease, aortic valve disease, and cardiomyopathy. Bundle branch blocks may be chronic or intermittent. A bundle branch block may be rate-related; for example, it often occurs when the heart rate exceeds some critical value.

Bundle branch blocks and depolarization abnormalities secondary to artificial pacemakers not only affect ventricular depolarization (QRS) but also are characteristically associated with *secondary repolarization* (ST-T) abnormalities. With bundle branch blocks, the T wave is typically opposite in polarity to the last deflection of the QRS (Fig. 268-10). This discordance of the QRS–T-wave vectors is caused by the altered sequence of repolarization that occurs secondary to altered depolarization. In contrast, *primary repolarization* abnormalities are independent of QRS changes and are related instead to actual alterations in the electrical properties of the myocardial fibers themselves (e.g., in the resting membrane potential or action potential duration), not just to changes in the sequence of repolarization. Ischemia, electrolyte imbalance, and drugs such as digitalis all cause such primary ST–T-wave changes. Primary and secondary T-wave changes may coexist. For example, T-wave inversions in the right precordial leads with left bundle branch block or in the left precordial leads with right bundle branch block may be important markers of underlying ischemia or other abnormalities. A distinctive abnormality simulating right bundle branch block with ST-segment elevations in the right chest leads is seen with the Brugada pattern (Chap. 276).

Partial blocks (fascicular or “hemiblocks”) in the left bundle system (left anterior or posterior fascicular blocks) generally do not prolong the QRS duration substantially but instead are associated with shifts in the frontal plane QRS axis (leftward or rightward, respectively). Left anterior fascicular block (QRS axis more negative than -45°) is probably the most common cause of marked left axis deviation in adults. In contrast, left posterior fascicular block (QRS axis more

rightward than $+110$ – 120°) is extremely rare as an isolated finding and requires exclusion of other factors causing right axis deviation mentioned earlier.

More complex combinations of fascicular and bundle branch blocks may occur that involve the left and right bundle system. Examples of *bifascicular block* include right bundle branch block and left posterior fascicular block, right bundle branch block with left anterior fascicular block, and complete left bundle branch block. Chronic bifascicular block in an asymptomatic individual is associated with a relatively low risk of progression to high-degree AV heart block. In contrast, new bifascicular block with acute anterior myocardial infarction carries a much greater risk of complete heart block. Alternation of right and left bundle branch block is a sign of *trifascicular disease*. However, the presence of a prolonged PR interval and bifascicular block does not necessarily indicate trifascicular involvement, since this combination may arise with AV node disease and bifascicular block. Intraventricular conduction delays also can be caused by extrinsic (toxic) factors that slow ventricular conduction, particularly hyperkalemia or drugs (e.g., class I antiarrhythmic agents, tricyclic antidepressants, phenothiazines).

Prolongation of QRS duration does not necessarily indicate a conduction delay but may be due to *preexcitation* of the ventricles via a bypass tract, as in Wolff-Parkinson-White (WPW) patterns (Chap. 276) and related variants. The diagnostic triad of WPW consists of a wide QRS complex associated with a relatively short PR interval and slurring of the initial part of the QRS (delta wave), with the latter effect being due to aberrant activation of ventricular myocardium. The presence of a bypass tract predisposes to reentrant supraventricular tachyarrhythmias.

MYOCARDIAL ISCHEMIA AND INFARCTION

(See also Chap. 295) The ECG is a cornerstone in the diagnosis of acute and chronic ischemic heart disease. The findings depend on several key factors: the nature of the process (reversible [i.e., ischemia] versus irreversible [i.e., infarction]), the duration (acute versus chronic), the extent (transmural versus subendocardial), and localization (anterior versus inferoposterior), as well as the presence of other underlying abnormalities (ventricular hypertrophy, conduction defects).

Ischemia exerts complex time-dependent effects on the electrical properties of myocardial cells. Severe, acute ischemia lowers the resting membrane potential and shortens the duration of the action potential. Such changes cause a voltage gradient between normal and ischemic zones. As a consequence, current flows between those regions. These currents of injury are represented on the surface ECG by deviation of the ST segment (Fig. 268-11). When the acute ischemia is *transmural*, the ST vector usually is shifted in the direction of the outer (epicardial) layers, producing ST elevations and sometimes, in the earliest stages of ischemia, tall, positive so-called hyperacute T waves over the ischemic zone. With ischemia confined primarily to the *subendocardium*, the ST vector typically shifts toward the subendocardium and ventricular cavity, so that overlying (e.g., anterior precordial) leads show ST-segment depression (with ST elevation in lead aVR). Multiple factors affect the amplitude of acute ischemic ST deviations. Profound ST elevation or depression in multiple leads usually indicates very severe ischemia.

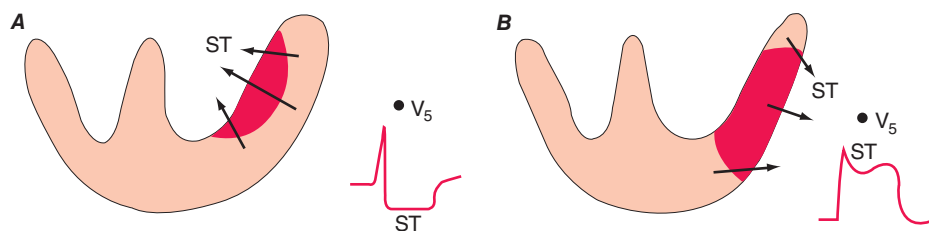


FIGURE 268-11 Acute ischemia causes a current of injury. With predominant subendocardial ischemia (A), the resultant ST vector will be directed toward the inner layer of the affected ventricle and the ventricular cavity. Overlying leads therefore will record ST depression. With ischemia involving the outer ventricular layer (B) (transmural or epicardial injury), the ST vector will be directed outward. Overlying leads will record ST elevation.

1456 From a clinical viewpoint, the division of acute myocardial infarction into ST-segment elevation and non-ST elevation types is useful since the efficacy of acute reperfusion therapy is limited to the former group.

The ECG leads are usually more helpful in localizing regions of ST elevation than non-ST elevation ischemia. For example, acute transmural anterior (including apical and lateral) wall ischemia is reflected by ST elevations or increased T-wave positivity in one or more of the precordial leads (V_1 - V_6) and leads I and aVL.

Inferior wall ischemia produces changes in leads II, III, and aVF. "Posterior" wall ischemia (usually associated with lateral or inferior involvement) may be indirectly recognized by reciprocal ST depressions in leads V_1 to V_3 (thus constituting an ST elevation "equivalent" acute coronary syndrome). Right ventricular ischemia usually produces ST elevations in right-sided chest leads (Fig. 268-5). When ischemic ST elevations occur as the earliest sign of acute infarction, they typically are followed within a period ranging from hours to days by evolving T-wave inversions and often by Q waves occurring in the same lead distribution. Reversible transmural ischemia, for example, due to coronary vasospasm (Prinzmetal's variant angina and possibly the Tako-tsubo "stress" cardiomyopathy syndrome), may cause transient ST-segment elevations without development of Q waves, as may very early reperfusion in acute coronary syndromes. Depending on the severity and duration of ischemia, the ST elevations may resolve completely in minutes or be followed by T-wave inversions that persist for hours or even days. Patients with ischemic chest pain who present with deep T-wave inversions in multiple precordial leads (e.g., V_1 - V_4 , I, and aVL) with or without cardiac enzyme elevations typically have severe obstruction in the left anterior descending coronary artery system (Fig. 268-12). In contrast, patients whose baseline ECG already shows abnormal T-wave inversions may develop T-wave normalization (*pseudonormalization*) during episodes of acute transmural ischemia.

With infarction, depolarization (QRS) changes often accompany repolarization (ST-T) abnormalities. Necrosis of sufficient myocardial

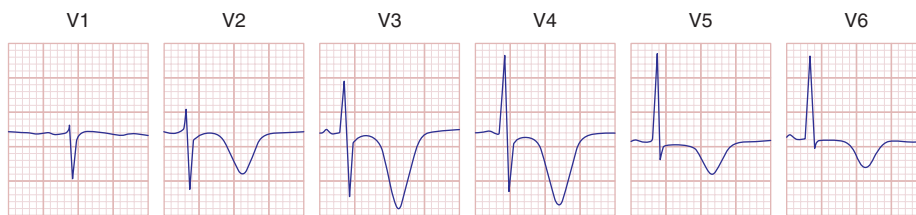


FIGURE 268-12 Severe anterior wall ischemia (with or without infarction) may cause prominent T-wave inversions in the precordial leads. This pattern (sometimes referred to as Wellens T waves) is usually associated with a high-grade stenosis of the left anterior descending coronary artery.

tissue may lead to decreased R-wave amplitude or abnormal Q waves (even in the absence of transmural) in the anterior or inferior leads (Fig. 268-13). Previously, abnormal Q waves were considered markers of transmural myocardial infarction, whereas subendocardial infarcts were thought not to produce Q waves. However, careful ECG-pathology correlative studies have indicated that transmural infarcts may occur without Q waves and that subendocardial (nontransmural) infarcts sometimes may be associated with Q waves. Therefore, infarcts are more appropriately classified as "Q-wave" or "non-Q-wave." The major acute ECG changes in syndromes of ischemic heart disease are summarized schematically in Fig. 268-14. Loss of depolarization forces due to posterior or lateral infarction may cause reciprocal increases in R-wave amplitude in leads V_1 and V_2 without diagnostic Q waves in any of the conventional leads. Atrial infarction may be associated with PR-segment deviations due to an atrial current of injury, changes in P-wave morphology, or atrial arrhythmias. In the weeks and months after infarction, these ECG changes may persist or begin to resolve. Complete normalization of the ECG after Q-wave infarction is uncommon but may occur, particularly with smaller infarcts. In contrast, ST-segment elevations that persist for several weeks or more after a Q-wave infarct usually correlate with a severe underlying wall motion disorder (akinetic or dyskinetic zone), although not necessarily a frank ventricular aneurysm. ECG changes due to ischemia may occur spontaneously or may be provoked by various exercise protocols (stress electrocardiography; Chap. 293).

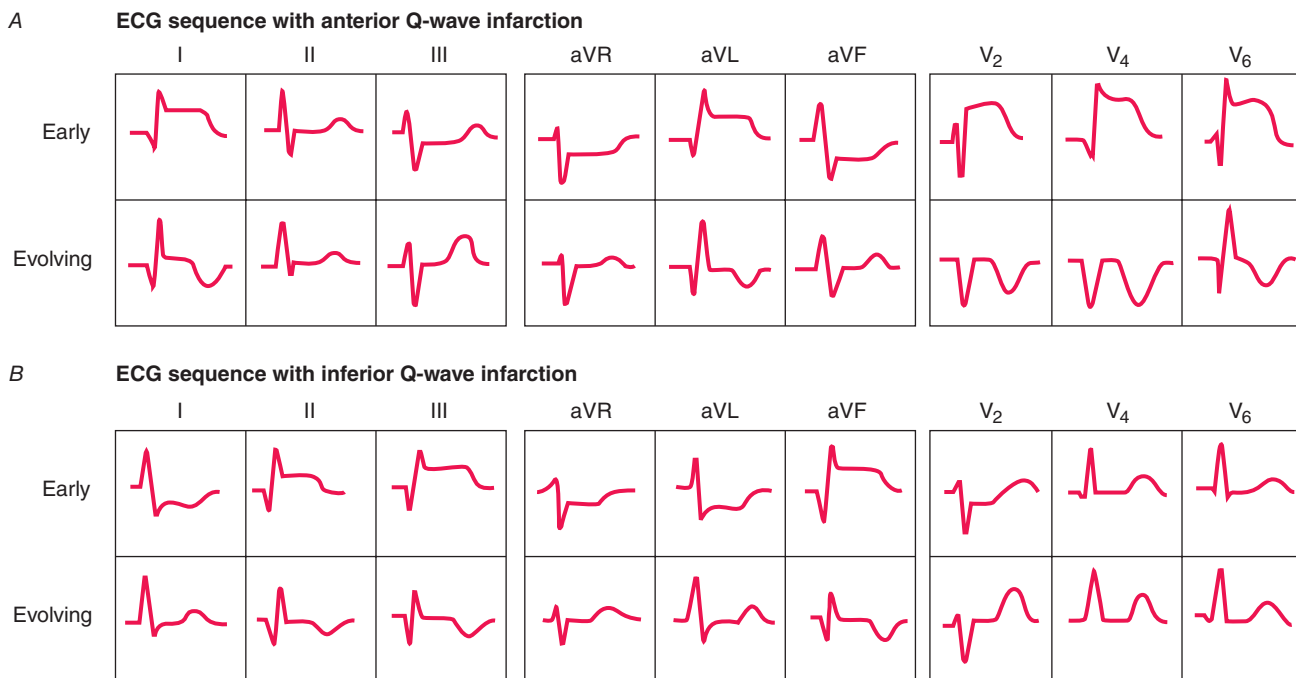


FIGURE 268-13 Sequence of depolarization and repolarization changes with (A) acute anterior and (B) acute inferior wall Q-wave infarctions. With anterior infarcts, ST elevation in leads I and aVL and the precordial leads may be accompanied by reciprocal ST depressions in leads II, III, and aVF. Conversely, acute inferior (or posterolateral) infarcts may be associated with reciprocal ST depressions in leads V_1 to V_3 . (After AL Goldberger et al: *Goldberger's Clinical Electrocardiography: A Simplified Approach*, 8th ed. Philadelphia, Elsevier/Saunders, 2013.)

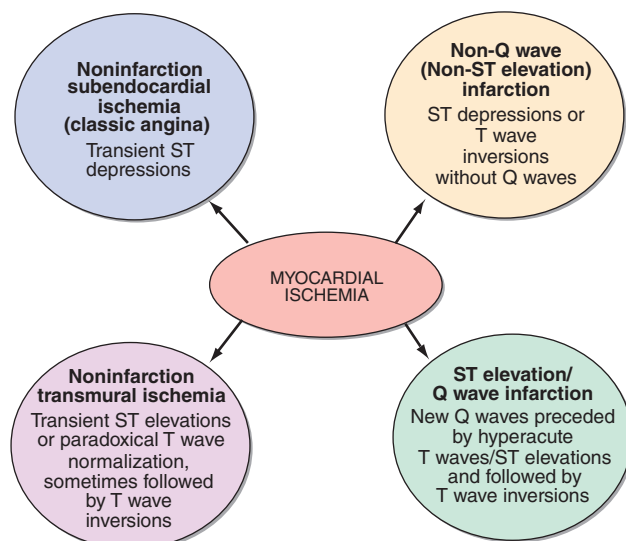


FIGURE 268-14 Variability of ECG patterns with acute myocardial ischemia. The ECG also may be normal or nonspecifically abnormal. Furthermore, these categorizations are not mutually exclusive. (After AL Goldberger et al: *Goldberger's Clinical Electrocardiography: A Simplified Approach*, 8th ed. Philadelphia, Elsevier/Saunders, 2013.)

The ECG has important limitations in both sensitivity and specificity in the diagnosis of ischemic heart disease. Although a single normal ECG does not exclude ischemia or even acute infarction, a normal ECG throughout the course of an acute infarct is distinctly uncommon. Prolonged chest pain without diagnostic ECG changes therefore should always prompt a careful search for other noncoronary causes of chest pain (Chap. 19). Furthermore, the diagnostic changes of acute or evolving ischemia are often masked by the presence of left bundle branch block, electronic ventricular pacemaker patterns, and Wolff-Parkinson-White preexcitation. However, clinicians continue to overdiagnose ischemia or infarction based on the presence of ST-segment elevations or depressions; T-wave inversions; tall, positive T waves; or Q waves *not* related to ischemic heart disease (pseudoinfarct patterns). For example, ST-segment elevations simulating ischemia may occur with acute pericarditis or myocarditis, as a normal variant (including the typical “early repolarization” pattern), or in a variety of other conditions (Table 268-1). Similarly, tall, positive T waves do not invariably represent hyperacute ischemic changes but may also be caused by normal variants, hyperkalemia, cerebrovascular injury, and left ventricular volume overload due to mitral or aortic regurgitation, among other causes.

ST-segment elevations and tall, positive T waves are common findings in leads V_1 and V_2 in left bundle branch block or left ventricular hypertrophy in the absence of ischemia. The differential diagnosis of Q waves includes physiologic or positional variants, ventricular hypertrophy, acute or chronic noncoronary myocardial injury, hypertrophic cardiomyopathy, and ventricular conduction disorders. Digoxin, ventricular hypertrophy, hypokalemia, and a variety of other factors may cause ST-segment depression mimicking subendocardial ischemia. Prominent T-wave inversion may occur with ventricular hypertrophy, cardiomyopathies, myocarditis, and cerebrovascular injury (particularly intracranial bleeds), among many other conditions.

METABOLIC FACTORS AND DRUG EFFECTS

A variety of metabolic and pharmacologic agents alter the ECG and, in particular, cause changes in repolarization (ST-T-U) and sometimes QRS prolongation. Certain life-threatening electrolyte disturbances may be diagnosed initially and monitored from the ECG. *Hyperkalemia* produces a sequence of changes (Fig. 268-15), usually beginning with narrowing and peaking (tenting) of the T waves. Further elevation of extracellular K^+ leads to AV conduction disturbances, diminution in P-wave amplitude, and widening of the QRS interval. Severe hyperkalemia

TABLE 268-1 DIFFERENTIAL DIAGNOSIS OF ST-SEGMENT ELEVATIONS

Ischemia/myocardial infarction
Noninfarction, transmural ischemia (Prinzmetal's angina, and probably Tako-tsubo syndrome, which may also exactly simulate classical acute infarction)
Acute myocardial infarction
Postmyocardial infarction (ventricular aneurysm pattern)
Acute pericarditis
Normal variants (including benign “early repolarization” patterns)
Left ventricular hypertrophy/left bundle branch block ^a
Other (rarer)
Acute pulmonary embolism ^a
Brugada patterns (right bundle branch block–like pattern with ST elevations in right precordial leads) ^a
Class 1C antiarrhythmic drugs ^a
DC cardioversion
Hypercalcemia ^a
Hyperkalemia ^a
Hypothermia (J [Osborn] waves)
Nonischemic myocardial injury
Myocarditis
Tumor invading left ventricle
Trauma to ventricles

^aUsually localized to V_1 – V_2 or V_3 .

Source: Modified from AL Goldberger et al: *Goldberger's Clinical Electrocardiography: A Simplified Approach*, 8th ed. Philadelphia, Elsevier/Saunders, 2013.

eventually causes cardiac arrest with a slow sinusoidal type of mechanism (“sine-wave” pattern) followed by asystole. *Hypokalemia* (Fig. 268-16) prolongs ventricular repolarization, often with prominent U waves. Prolongation of the QT interval is also seen with drugs that increase the duration of the ventricular action potential: class 1A antiarrhythmic agents and related drugs (e.g., quinidine, disopyramide, procainamide, tricyclic antidepressants, phenothiazines) and class III agents (e.g., amiodarone [Fig. 268-16], dofetilide, dronedarone, sotalol, ibutilide). Marked QT prolongation, sometimes with deep, wide T-wave inversions, may occur with intracranial bleeds, particularly subarachnoid hemorrhage (“CVA T-wave” pattern) (Fig. 268-16). Systemic *hypothermia* also prolongs repolarization, usually with a distinctive convex elevation of the J point (Osborn wave). *Hypocalcemia* typically prolongs the QT interval (ST portion), whereas *hypercalcemia* shortens it (Fig. 268-17). Digitalis glycosides also shorten the QT interval, often with a characteristic “scooping” of the ST–T-wave complex (*digitalis effect*).

Many other factors are associated with ECG changes, particularly alterations in ventricular repolarization. T-wave flattening, minimal T-wave inversions, or slight ST-segment depression (“nonspecific ST–T-wave changes”) may occur with a variety of electrolyte and acid-base disturbances, a number of infectious processes, central nervous system disorders, endocrine abnormalities, many drugs, ischemia, hypoxia, and virtually any type of cardiopulmonary abnormality. Although subtle ST–T-wave changes may be markers of ischemia, transient nonspecific repolarization changes may also occur after a meal or with postural (orthostatic) change, hyperventilation, or exercise in healthy individuals.

LOW QRS VOLTAGE

Low QRS voltage is arbitrarily defined as peak-to-trough QRS amplitudes of ≤ 5 mm in the six limb leads and/or ≤ 10 mm in the chest leads. Multiple factors may be responsible. Among the most serious include pericardial (Fig. 268-18) or pleural effusions, chronic obstructive pulmonary disease, infiltrative cardiomyopathies, and anasarca.

ELECTRICAL ALTERNANS

Electrical alternans—a beat-to-beat alternation in one or more components of the ECG signal—is a common type of nonlinear cardiovascular response to a variety of hemodynamic and electrophysiologic

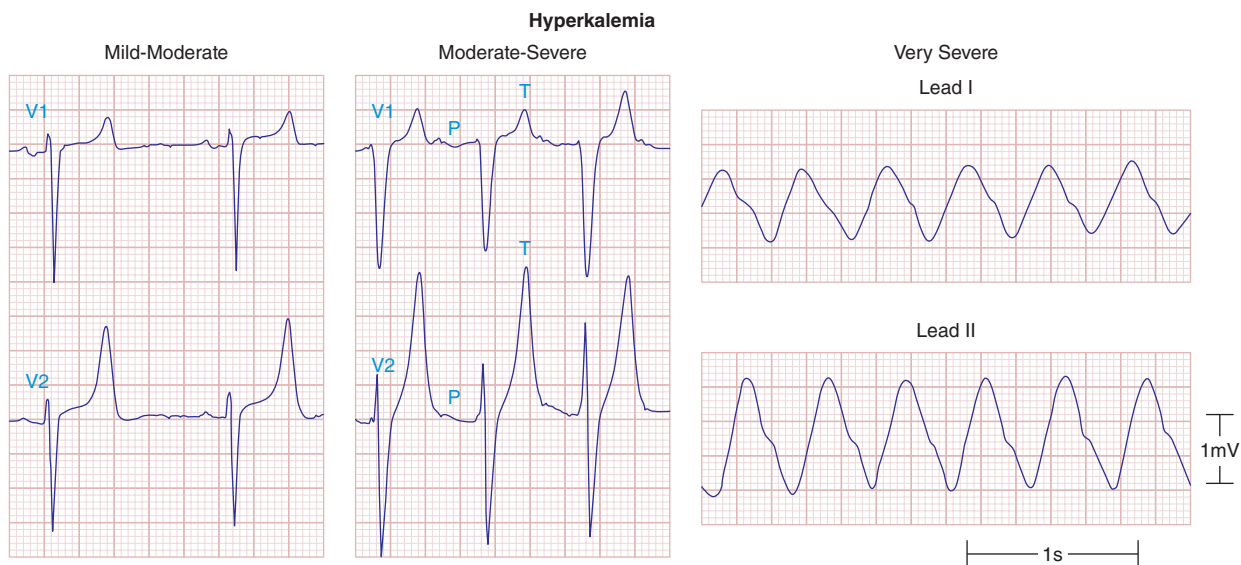


FIGURE 268-15 The earliest ECG change with hyperkalemia is usually peaking (“tenting”) of the T waves. With further increases in the serum potassium concentration, the QRS complexes widen, the P waves decrease in amplitude and may disappear, and finally a sine-wave pattern leads to asystole unless emergency therapy is given. (After AL Goldberger et al: *Goldberger’s Clinical Electrocardiography: A Simplified Approach*, 8th ed. Philadelphia, Elsevier/Saunders, 2013.)

perturbations. Total electrical alternans (P-QRS-T) with sinus tachycardia is a relatively specific sign of pericardial effusion, usually with cardiac tamponade (Fig. 268-18). The mechanism relates to a periodic swinging motion of the heart in the effusion at a frequency exactly one-half the heart rate. In contrast, pure repolarization (ST-T or U wave) alternans is a sign of electrical instability and may precede ventricular tachyarrhythmias.

CLINICAL INTERPRETATION OF THE ECG

Accurate analysis of ECGs requires thoroughness and care. The patient’s age, gender, and clinical status should always be taken into account. Many mistakes in ECG interpretation are errors of omission. Therefore, a systematic approach is essential. The following 14 points should be analyzed carefully in every ECG: (1) standardization (calibration) and technical features (including lead placement and artifacts), (2) rhythm, (3) heart rate, (4) PR interval/AV conduction, (5) QRS interval, (6) QT/QT_c intervals, (7) mean QRS electrical axis, (8) P waves, (9) QRS voltages, (10) precordial R-wave progression,

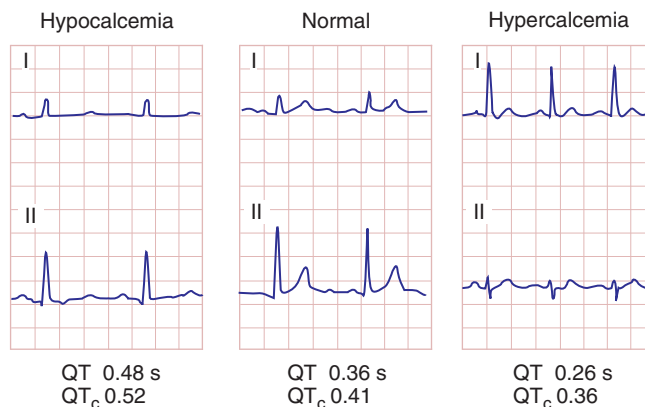


FIGURE 268-17 Prolongation of the Q-T interval (ST-segment portion) is typical of hypocalcemia. Hypercalcemia may cause abbreviation of the ST segment and shortening of the QT interval.

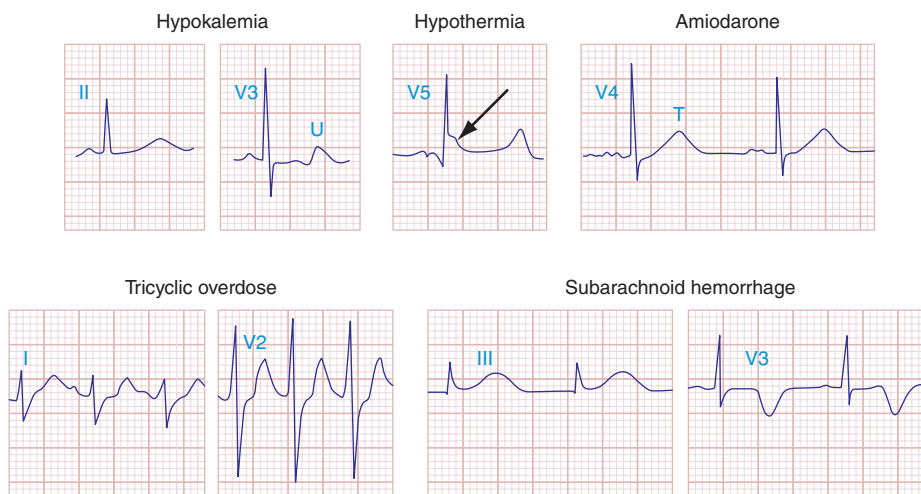


FIGURE 268-16 A variety of metabolic derangements, drug effects, and other factors may prolong ventricular repolarization with QT prolongation or prominent U waves. Prominent repolarization prolongation, particularly if due to hypokalemia, inherited “channelopathies,” or certain pharmacologic agents, indicates increased susceptibility to *torsades des pointes*-type ventricular tachycardia (Chap. 277). Marked systemic hypothermia is associated with a distinctive convex “hump” at the J point (Osborn wave, arrow) due to altered ventricular action potential characteristics. Note QRS and QT prolongation along with sinus tachycardia in the case of tricyclic antidepressant overdose.

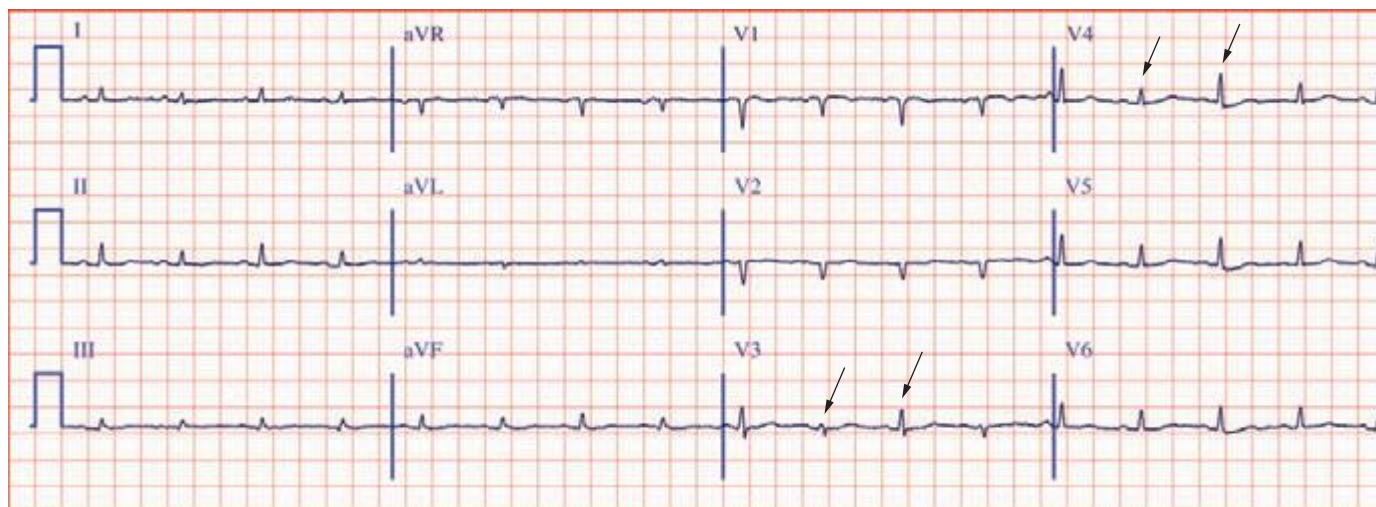


FIGURE 268-18 Classic triad of findings for pericardial effusion with cardiac tamponade: (1) sinus tachycardia; (2) low QRS voltages; and (3) electrical alternans (best seen in leads V_3 and V_4 in this case; arrows). This triad is highly specific for pericardial effusion, usually with tamponade physiology, but of limited sensitivity. (Adapted from LA Nathanson et al: *ECG Wave-Maven*. <http://ecg.bidmc.harvard.edu>.)

(11) abnormal Q waves, (12) ST segments, (13) T waves, and (14) U waves. Comparison with any previous ECGs is invaluable. **The diagnosis and management of specific cardiac arrhythmias and conduction disturbances are discussed in Chaps. 274 and 276.**

COMPUTERIZED ELECTROCARDIOGRAPHY

Computerized ECG systems are widely used for immediate retrieval of thousands of ECG records. Computer interpretation of ECGs still has major limitations. Incomplete or inaccurate readings are most likely with arrhythmias and complex abnormalities. Therefore, computerized interpretation (including measurements of basic ECG intervals) should not be accepted without careful clinician review.

The electrocardiograms (ECGs) in this atlas supplement those illustrated in **Chap. 268**. The interpretations emphasize findings of specific teaching value.

All of the figures are from ECG Wave-Maven, Copyright 2003, Beth Israel Deaconess Medical Center, <http://ecg.bidmc.harvard.edu>.

The abbreviations used in this chapter are as follows:

- AF—atrial fibrillation
- HCM—hypertrophic cardiomyopathy
- LBBB—left bundle branch block
- LVH—left ventricular hypertrophy
- MI—myocardial infarction
- RBBB—right bundle branch block
- RV—right ventricular
- RVH—right ventricular hypertrophy
- SR—sinus rhythm

MYOCARDIAL ISCHEMIA AND INFARCTION

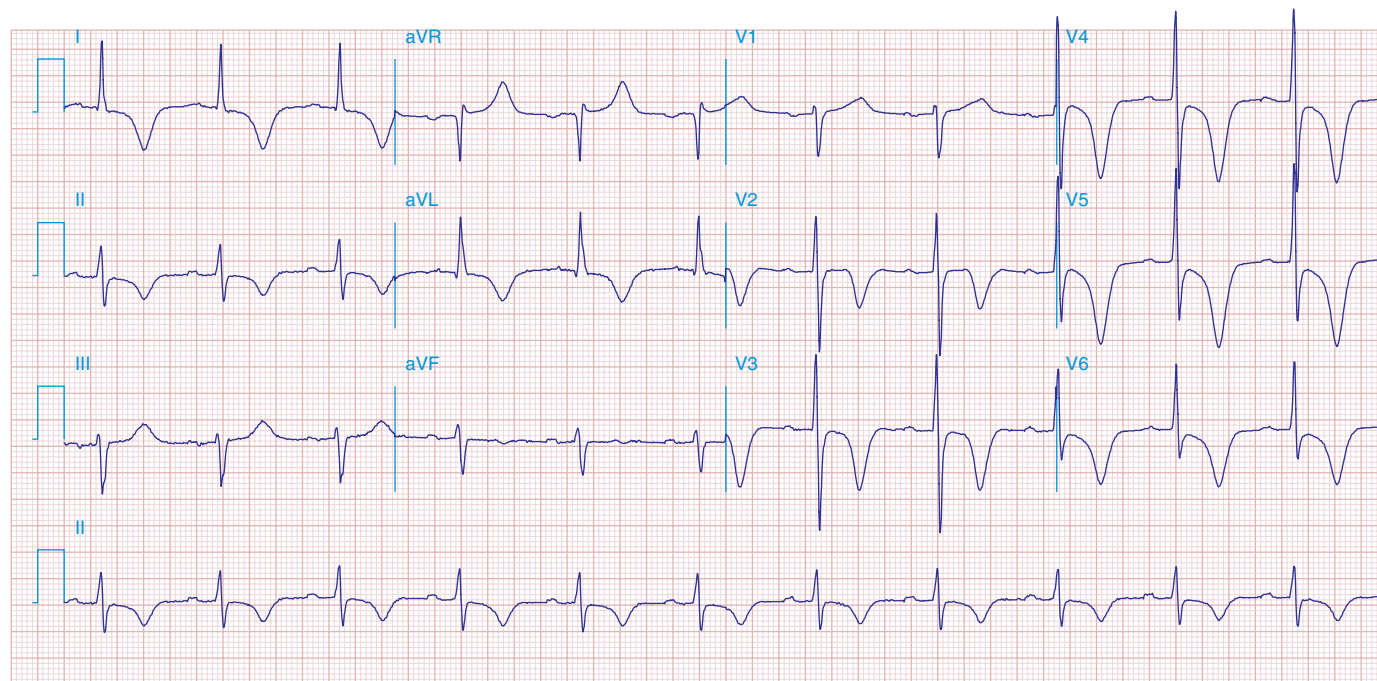


FIGURE 269e-1 Anterior wall ischemia (deep T-wave inversions and ST-segment depressions in I, aVL, V₃–V₆) in a patient with LVH (increased voltage in V₂–V₆).

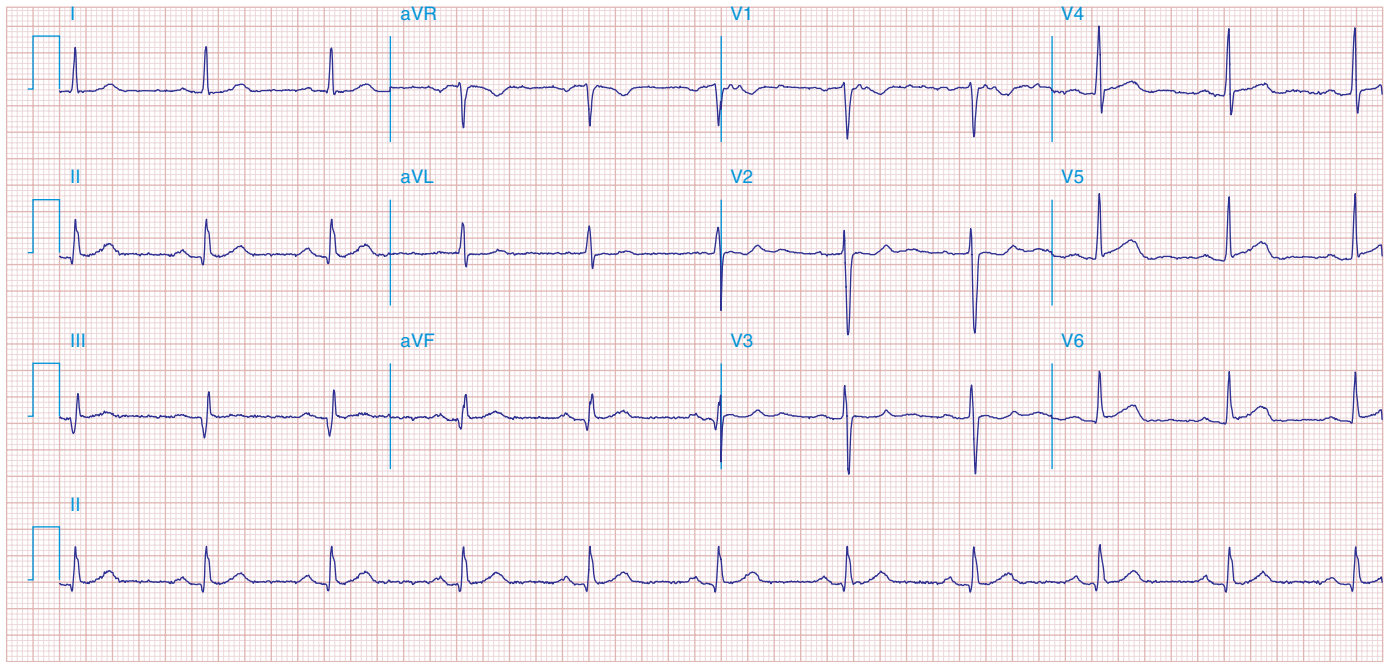


FIGURE 269e-2 Acute anterolateral wall ischemia with ST elevations in V_4 – V_6 . Probable prior inferior MI with Q waves in leads II, III, and aVF.

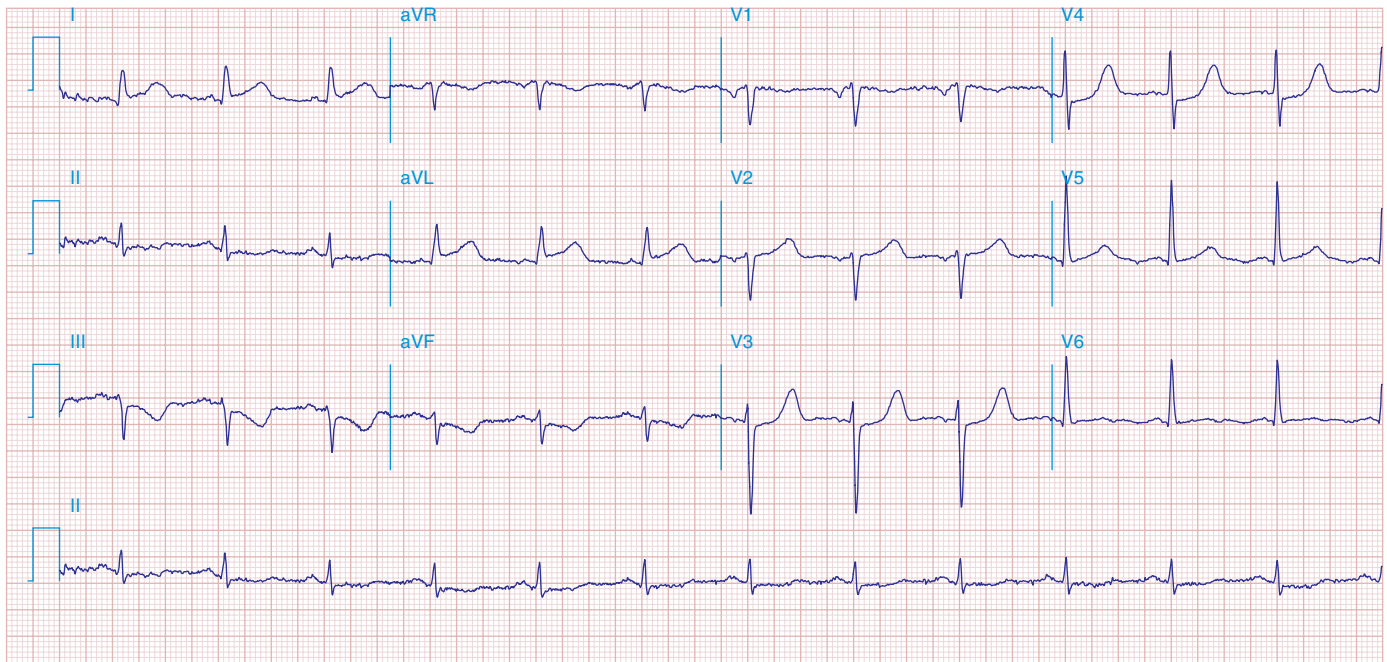


FIGURE 269e-3 Acute lateral ischemia with ST elevations in I and aVL with probable reciprocal ST depressions inferiorly (II, III, and aVF). Ischemic ST depressions also in V_3 and V_4 . Left atrial abnormality.

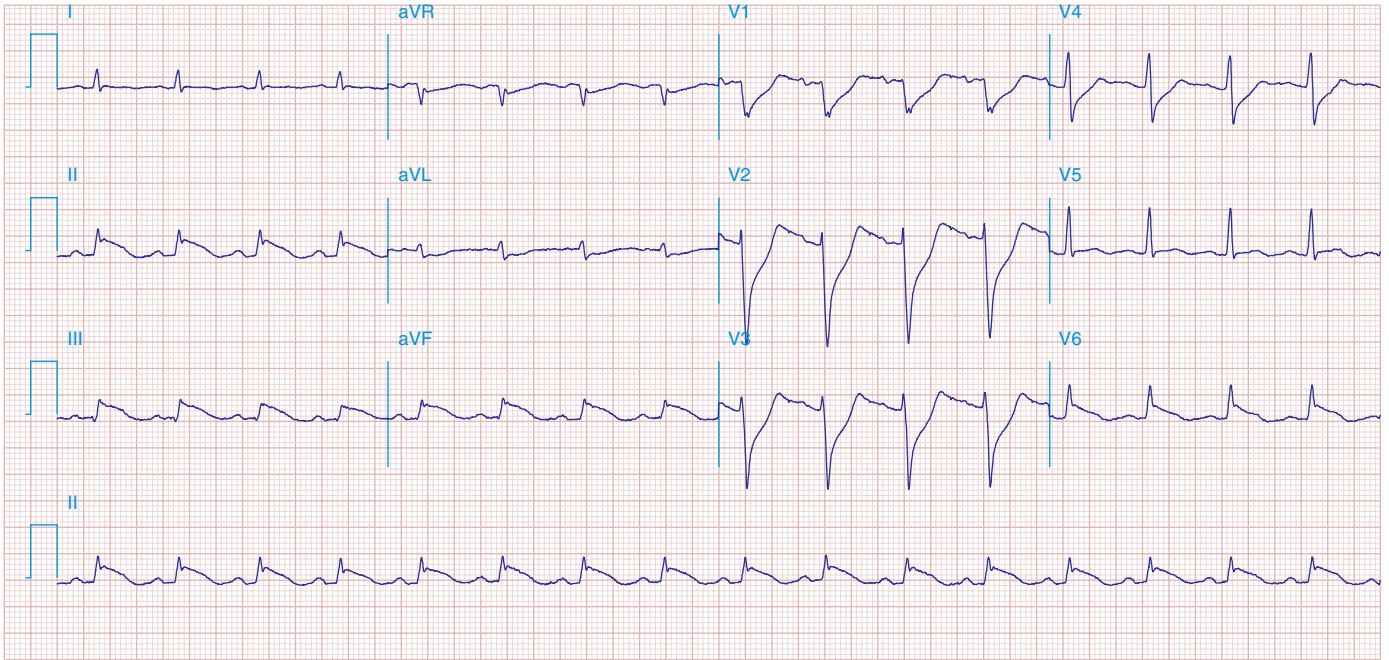


FIGURE 269e-4 Sinus tachycardia. Marked ischemic ST-segment elevations in inferior limb leads (II, III, aVF) and laterally (V_4) suggestive of acute inferolateral MI, and prominent ST-segment depressions with upright T waves in V_1 – V_4 are consistent with associated acute posterior MI.

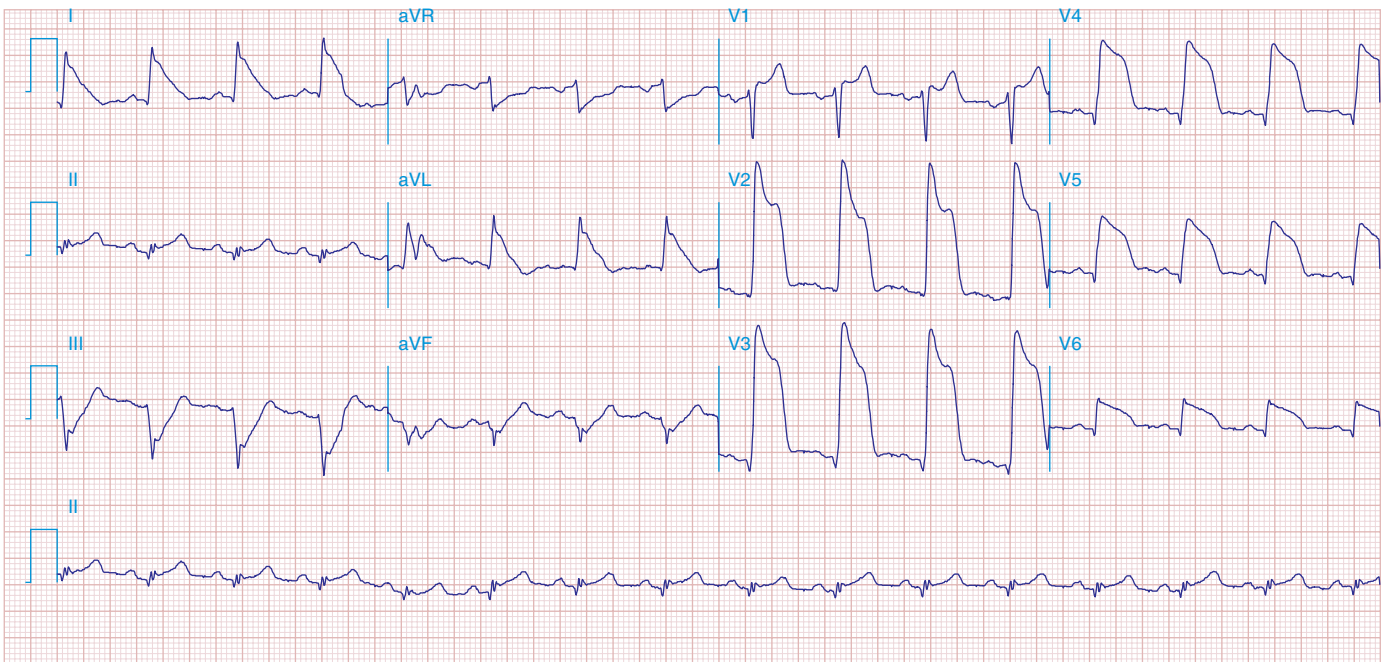


FIGURE 269e-5 Acute, extensive anterior MI with marked ST elevations in I, aVL, V_1 – V_6 and low amplitude pathologic Q waves in V_3 – V_6 . Marked reciprocal ST-segment depressions in III and aVF.

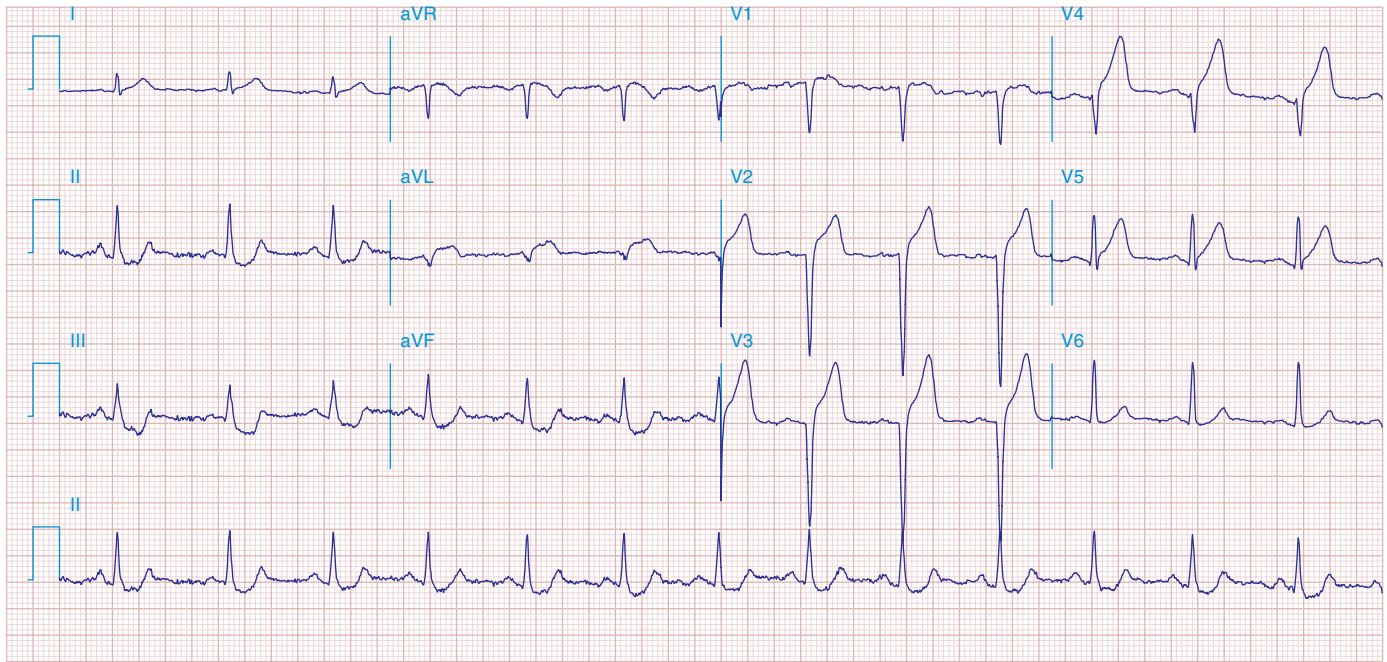


FIGURE 269e-6 Acute anterior wall MI with ST elevations and Q waves in V_1 - V_4 and aVL and reciprocal inferior ST depressions.

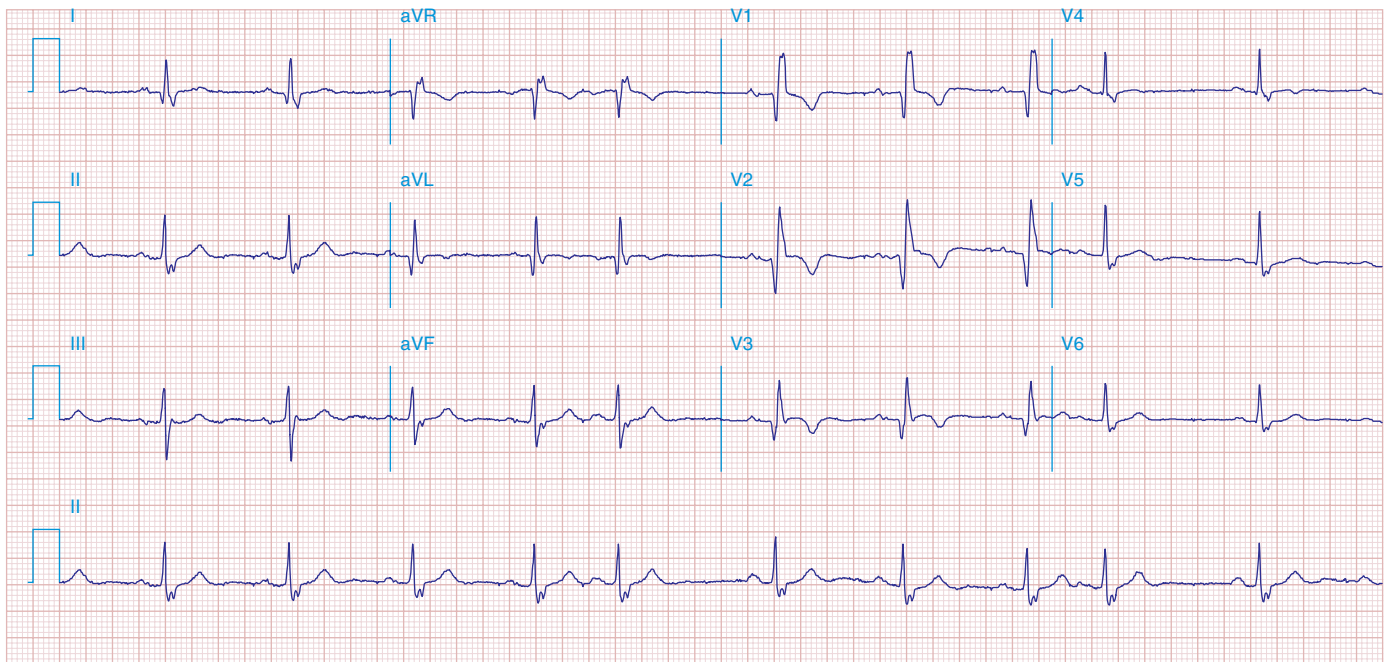


FIGURE 269e-7 SR with premature atrial complexes. RBBB; pathologic Q waves and ST elevation due to acute anterior/septal MI in V_1 - V_3 .

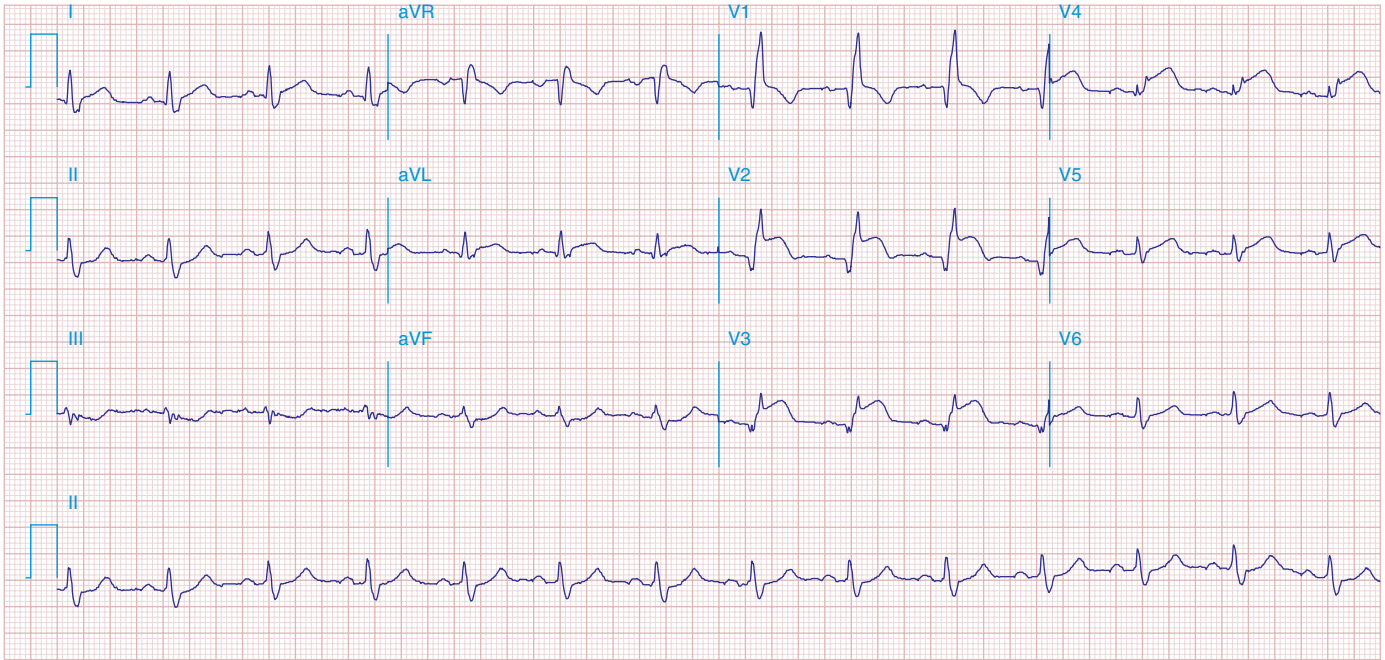


FIGURE 269e-8 Acute anteroseptal MI (Q waves and ST elevations in V_1 - V_4) with RBBB (note terminal R waves in V_1).

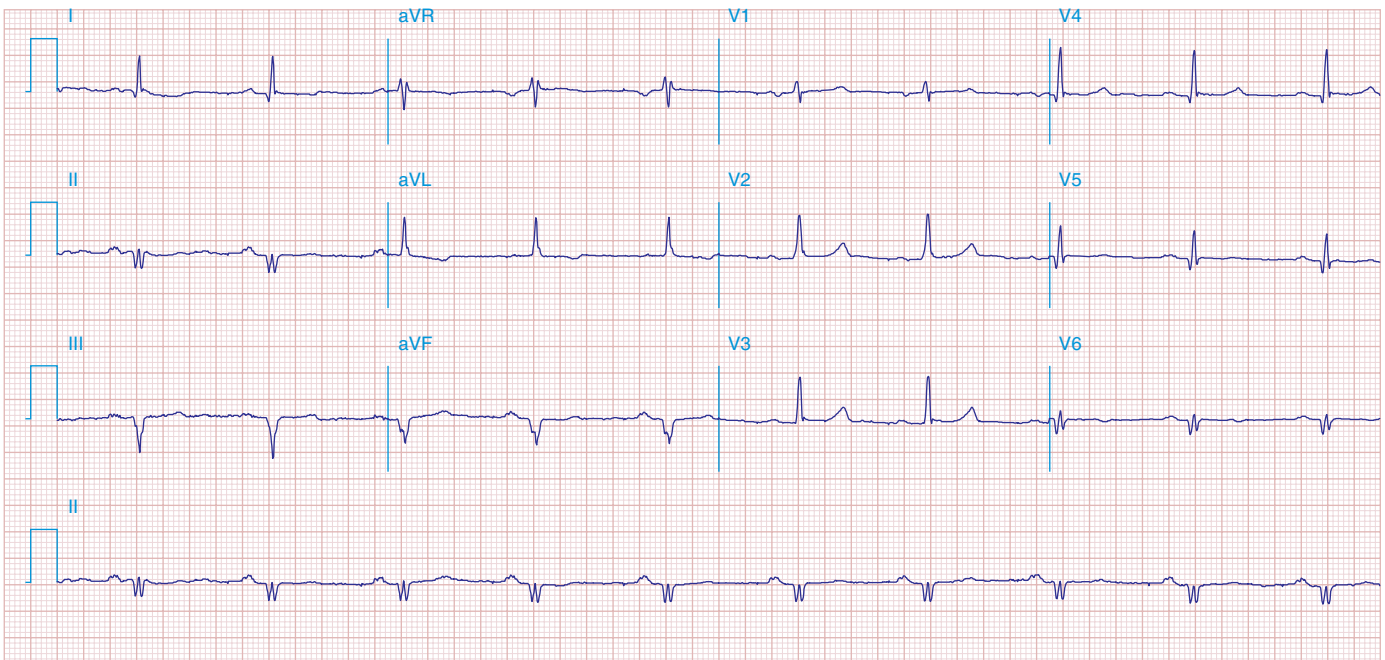


FIGURE 269e-9 Extensive prior MI involving inferior-posterior-lateral wall (Q waves in leads II, III, aVF, tall R waves in V_1 , V_2 , and Q waves in V_5 , V_6). T-wave abnormalities in leads I and aVL, V_5 , and V_6 .

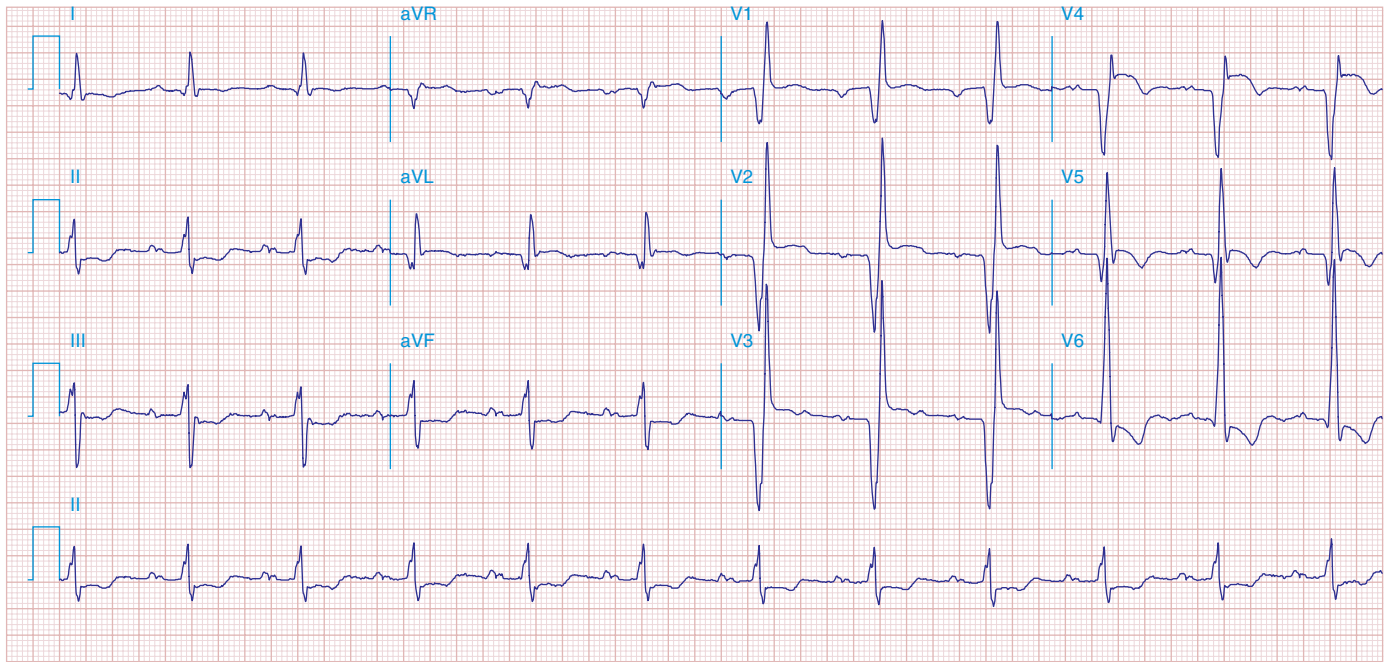


FIGURE 269e-10 SR with PR prolongation (“first-degree AV block”), left atrial abnormality, LVH, and RBBB. Pathologic Q waves in V_1 – V_5 and aVL with ST elevations (a chronic finding in this patient). Findings compatible with prior anterolateral MI and left ventricular aneurysm.

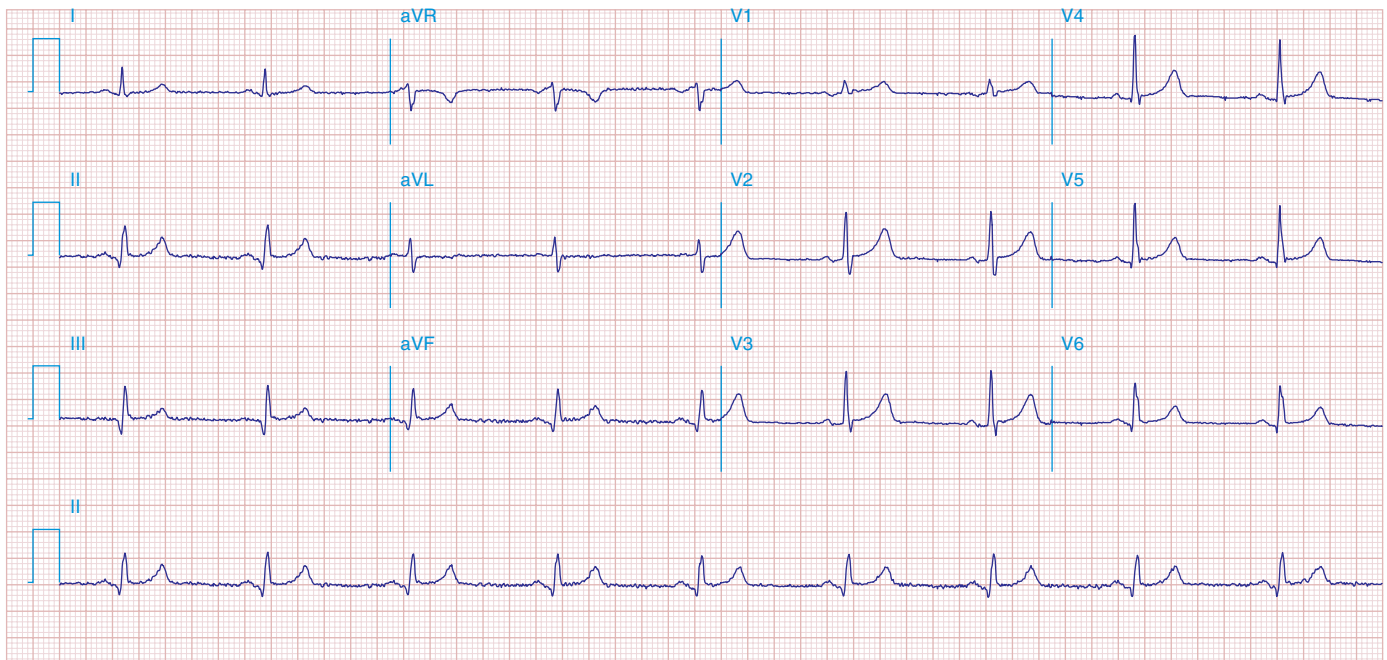


FIGURE 269e-11 Prior inferior-posterior MI. Wide (0.04 s) Q waves in the inferior leads (II, III, aVF); broad R wave in V_1 (a Q wave “equivalent” here). Absence of right-axis deviation and the presence of upright T waves in V_1 – V_2 are also against RVH.

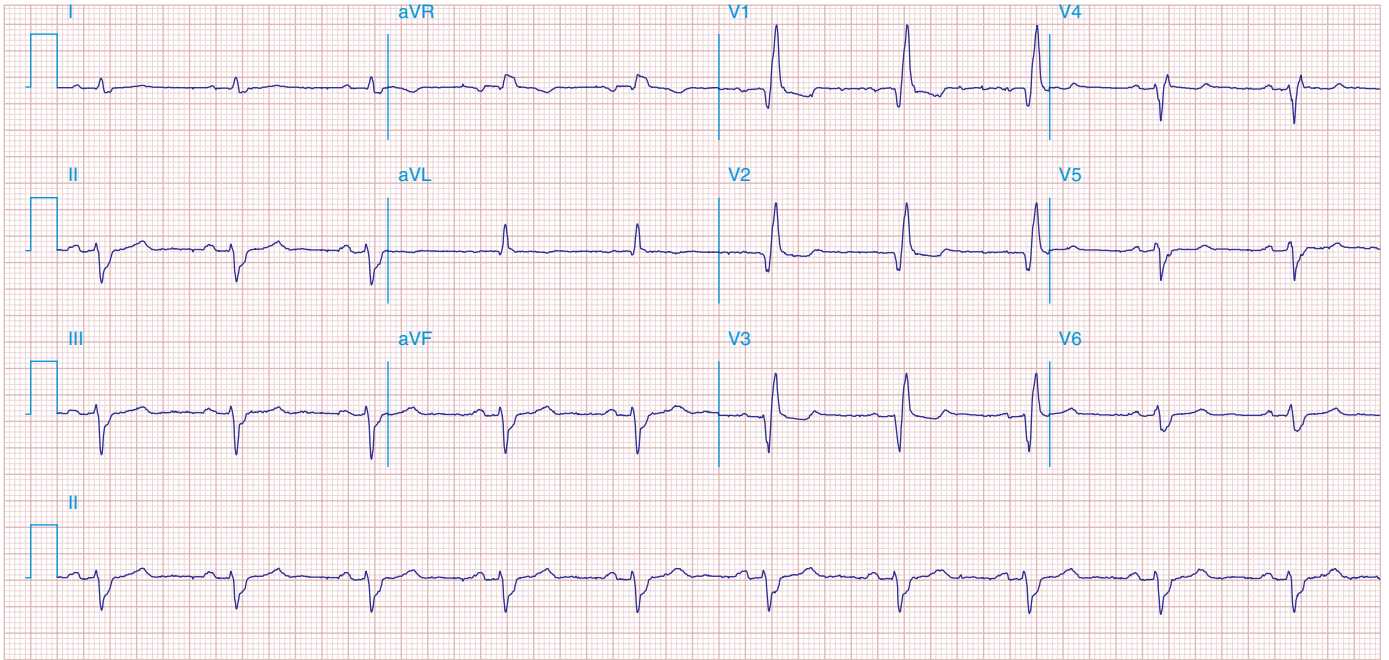


FIGURE 269e-12 SR with RBBB (broad terminal R wave in V₁) and left anterior fascicular block (hemiblock) and pathologic anterior Q waves in V₁–V₃. Patient had severe multivessel coronary artery disease, with echocardiogram showing septal dyskinesis and apical akinesis.

PERICARDITIS

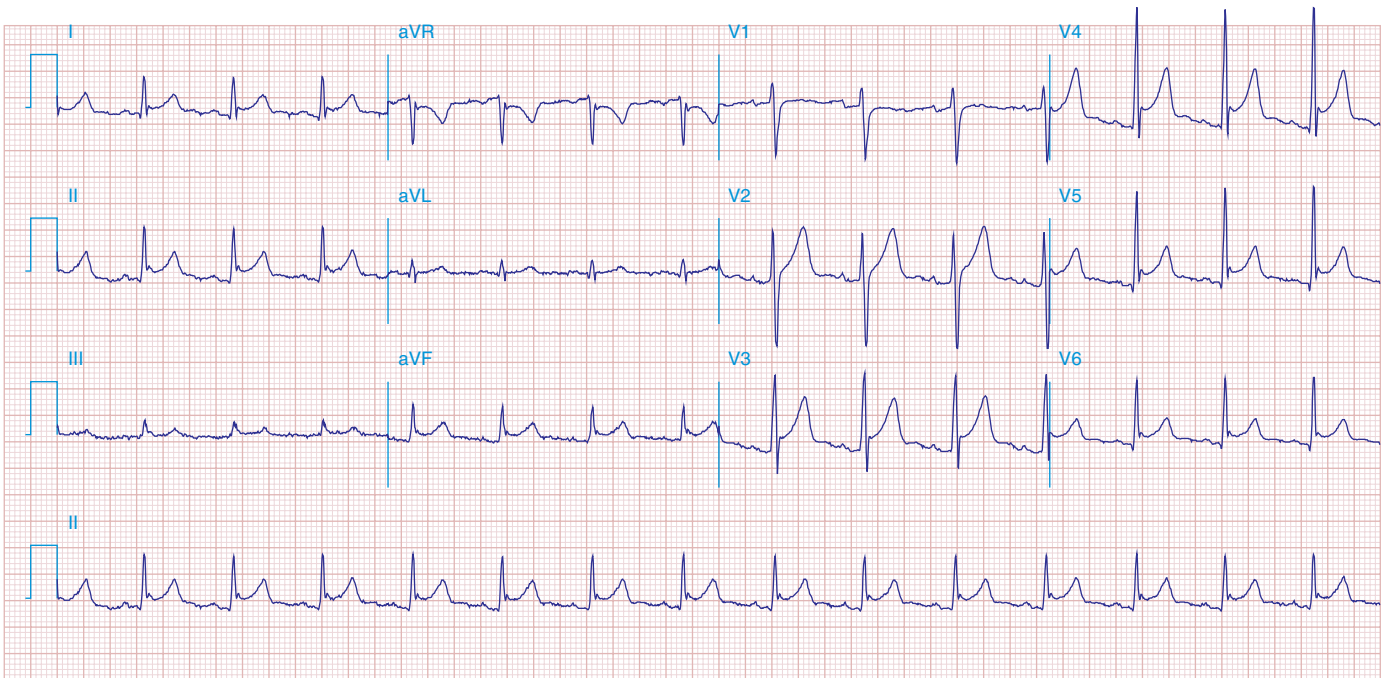


FIGURE 269e-13 Acute pericarditis with diffuse ST elevations in I, II, III, aVF, V₃–V₆, without T-wave inversions. Also note concomitant PR-segment elevation in aVR and PR depression in the inferolateral leads.

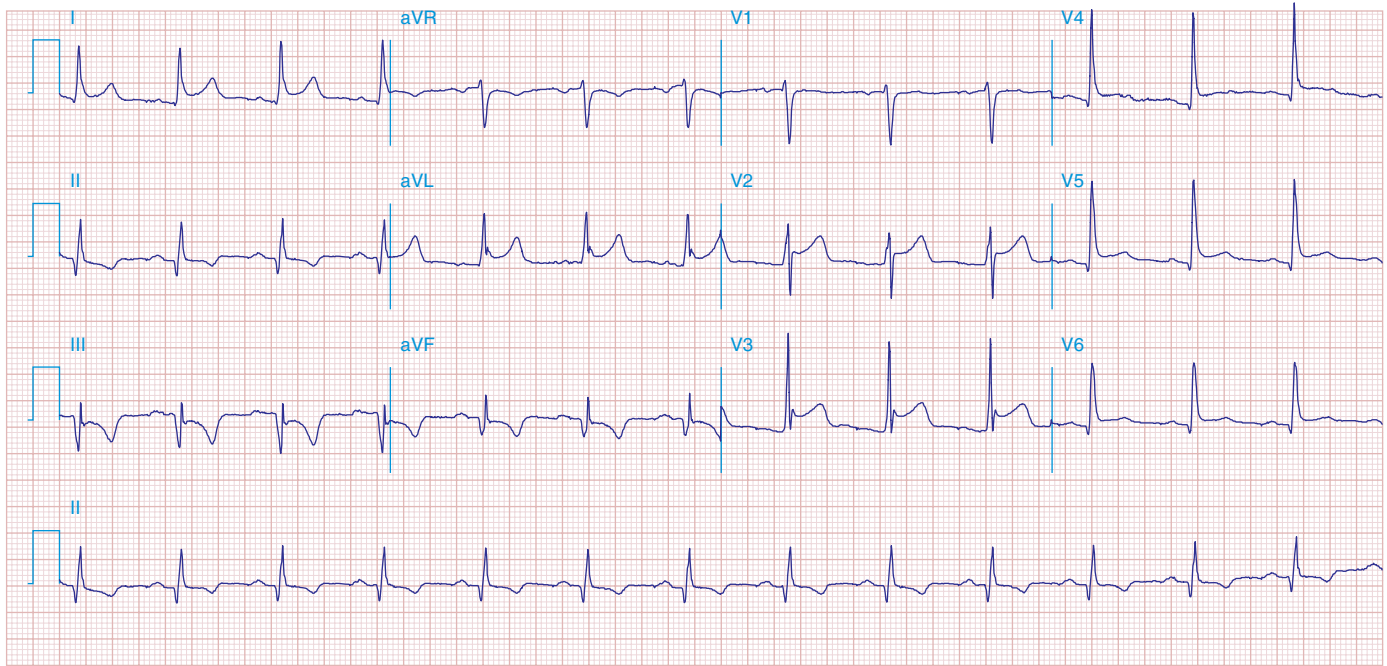


FIGURE 269e-14 SR; diffuse ST elevations (I, II, aVL, aVF, V₂-V₆) with associated PR deviations (elevated PR in aVR; depressed in V₄-V₆); borderline low voltage. Q-wave and T-wave inversions in II, III, and aVF. Diagnosis: acute pericarditis with inferior Q-wave MI.

VALVULAR HEART DISEASE AND HYPERTROPHIC CARDIOMYOPATHY

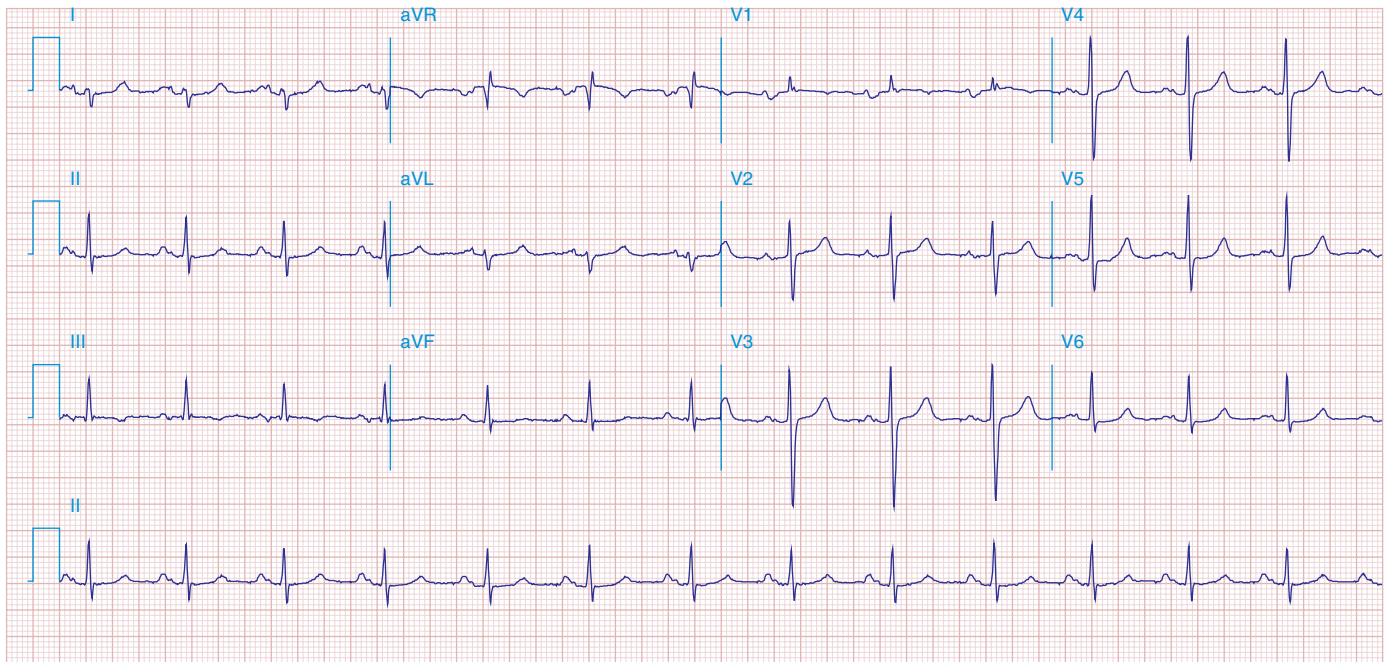


FIGURE 269e-15 SR, prominent left atrial abnormality (see I, II, V₁), right-axis deviation, and RVH (tall, relatively narrow R wave in V₁) in a patient with mitral stenosis.

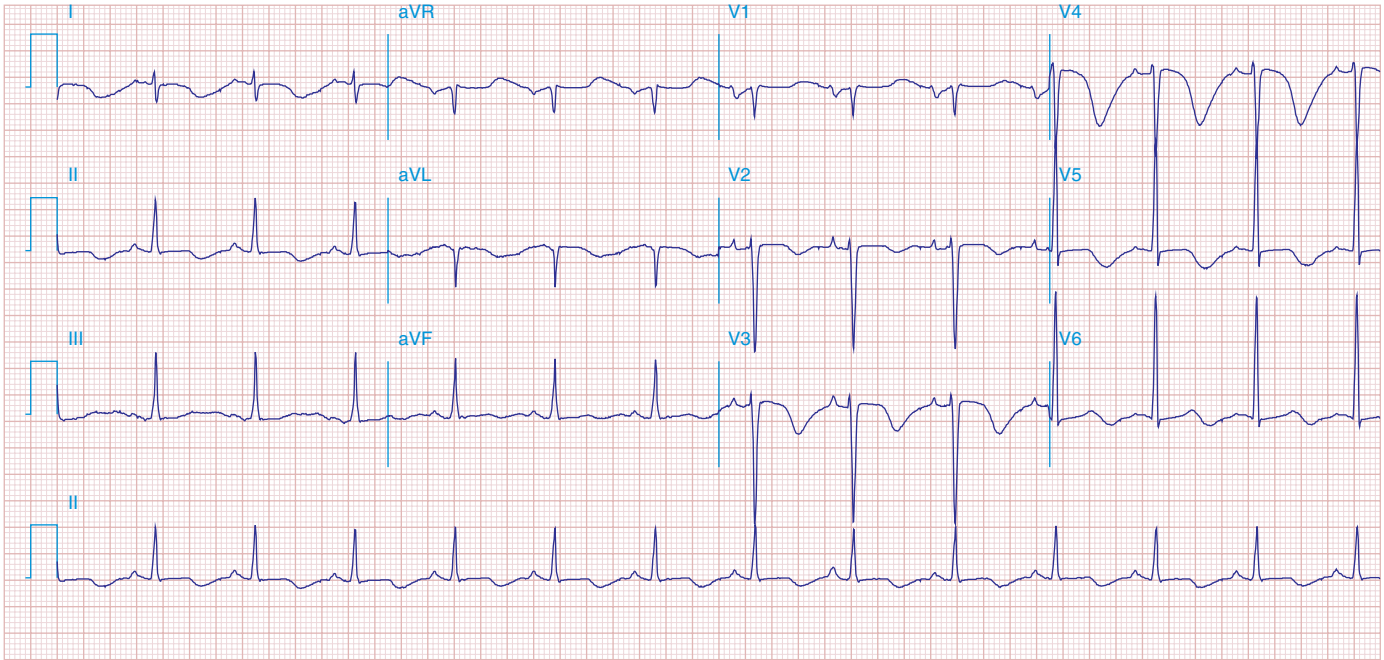


FIGURE 269e-16 SR, left atrial abnormality, and LVH by voltage criteria with borderline right-axis deviation in a patient with mixed mitral stenosis (left atrial abnormality and right-axis deviation) and mitral regurgitation (LVH). Prominent precordial T-wave inversions and QT prolongation also present.

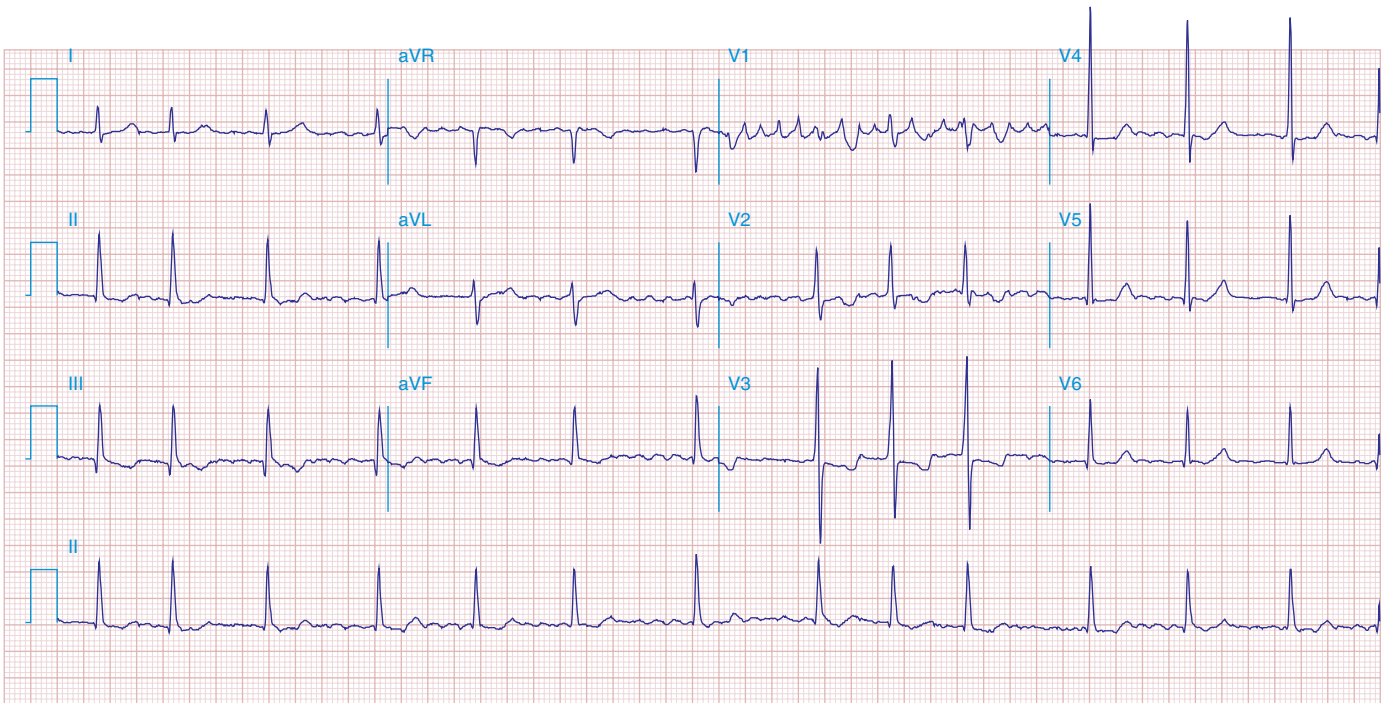


FIGURE 269e-17 Coarse AF, tall R in V_2 with vertical QRS axis (positive R in aVF) indicating RVH. Tall R in V_4 may be due to concomitant LVH. Patient had severe mitral stenosis with moderate mitral regurgitation.

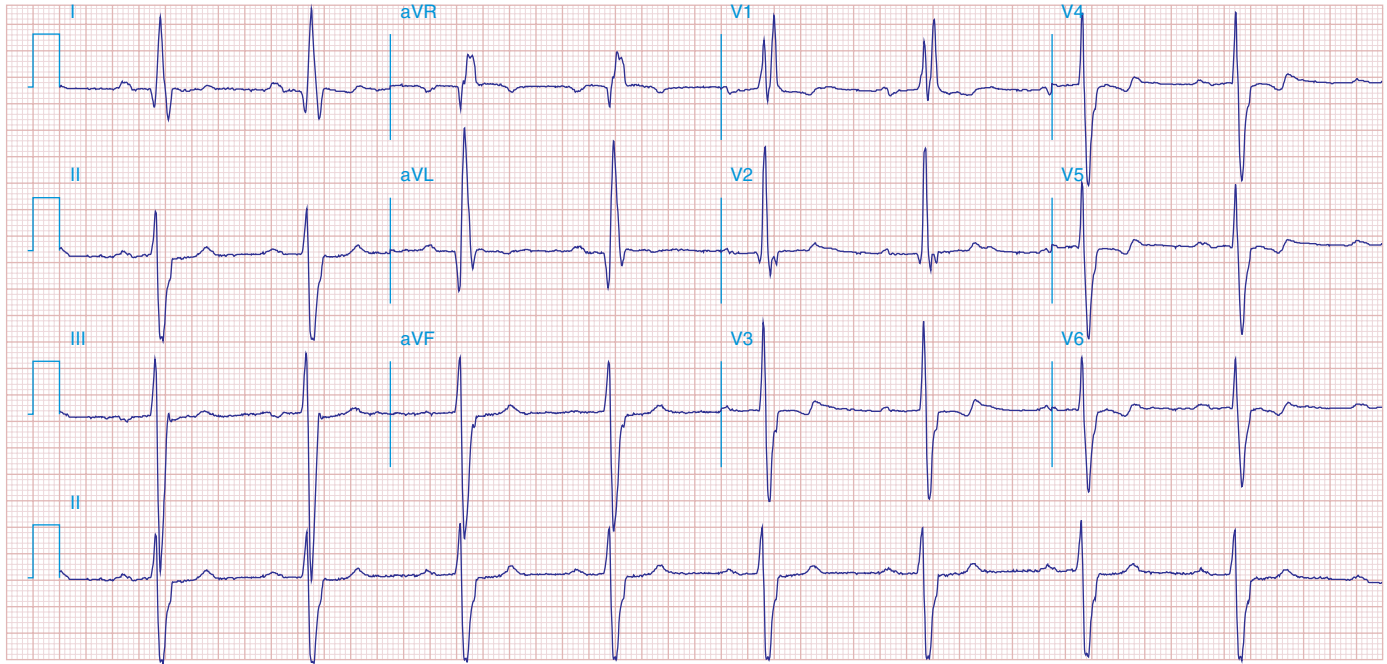


FIGURE 269e-18 SR; first-degree AV “block” (PR prolongation); LVH (tall R in aVL); RBBB (wide multiphasic R wave in V1) and left anterior fascicular block in a patient with HCM. Deep Q waves in I and aVL are consistent with septal hypertrophy.

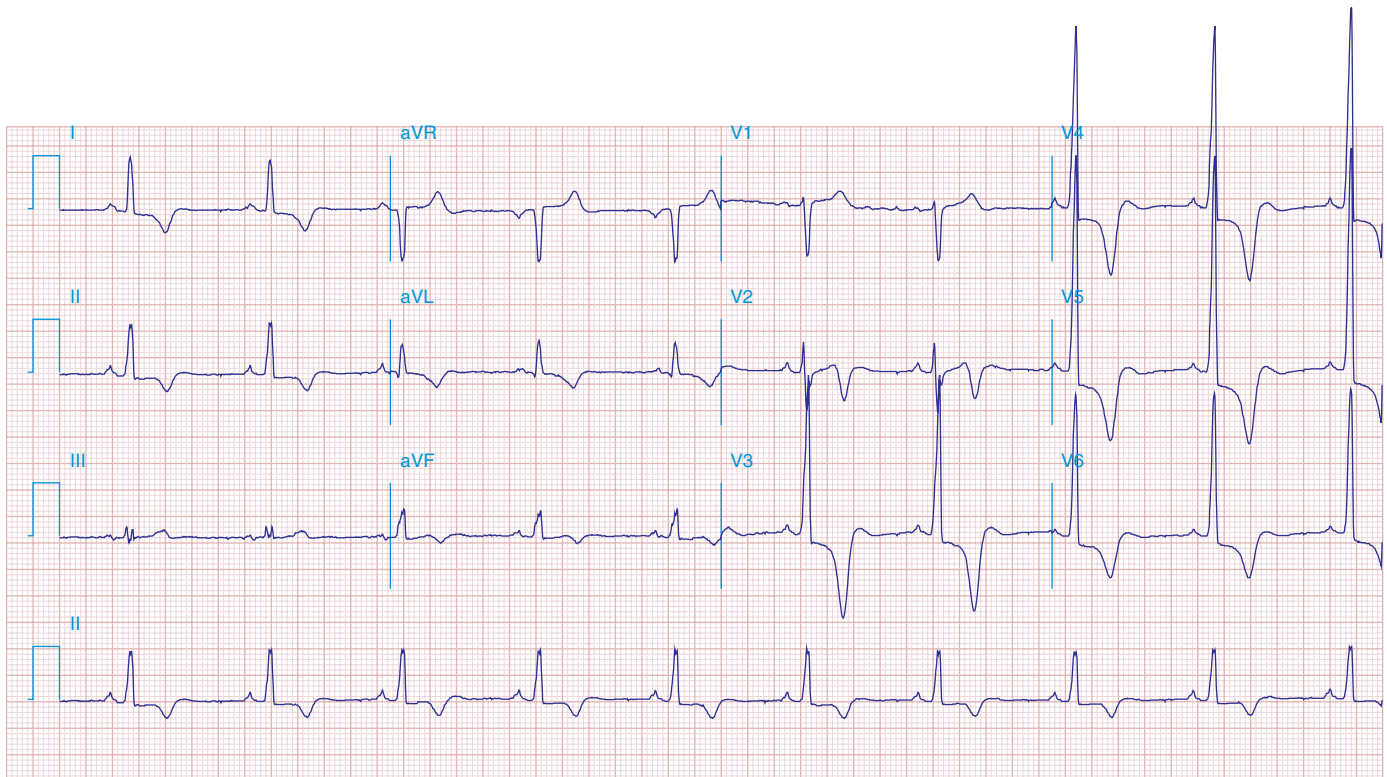


FIGURE 269e-19 LVH with deep T-wave inversions in limb leads and precordial leads. Striking T-wave inversions in mid-precordial leads suggest apical HCM (Yamaguchi’s syndrome).

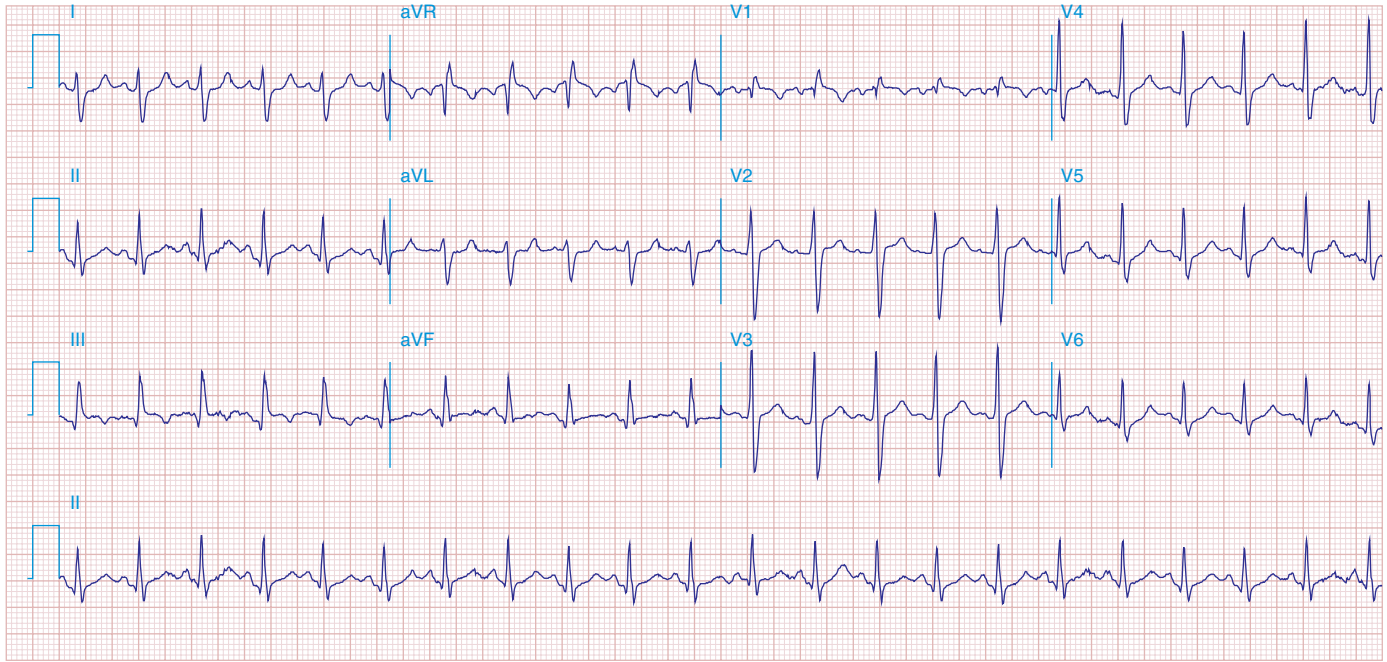


FIGURE 269e-20 Sinus tachycardia with S1Q3T3 pattern (T-wave inversion in III), incomplete RBBB, and right precordial T-wave inversions consistent with acute RV overload in a patient with pulmonary emboli.

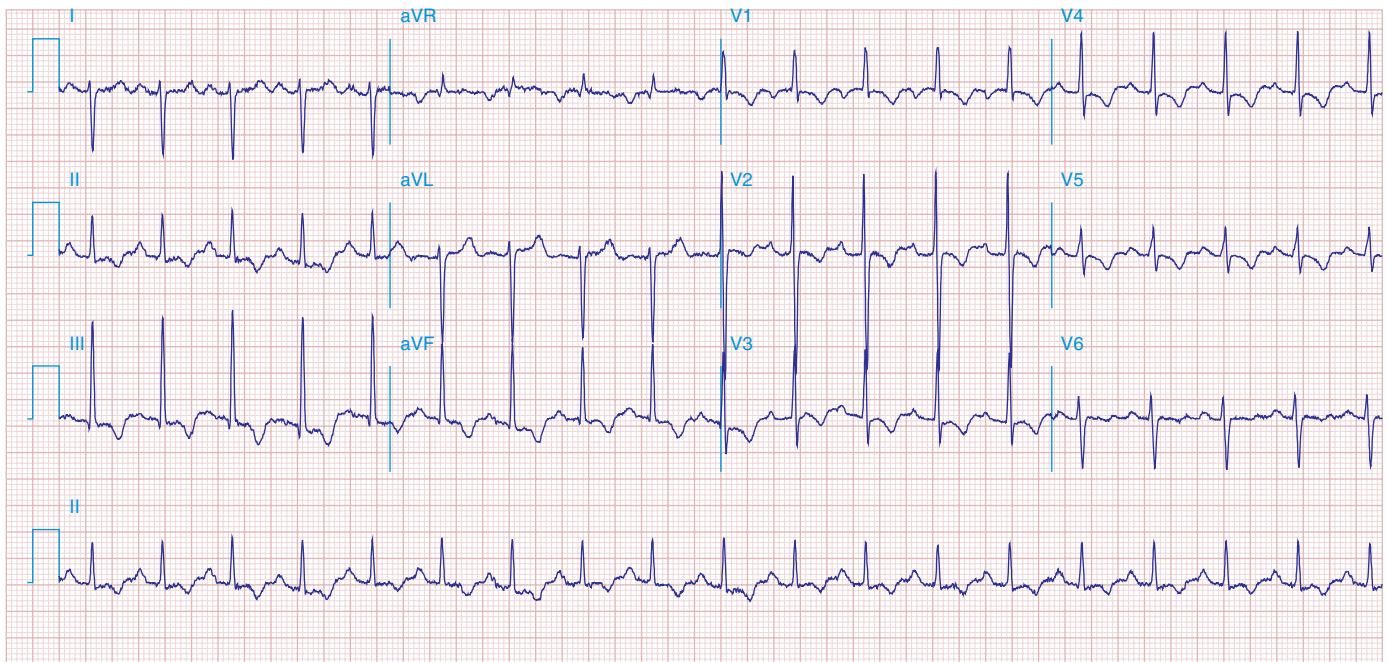


FIGURE 269e-21 Sinus tachycardia, right-axis deviation, RVH with tall R in V_1 and deep S in V_6 , and inverted T waves in II, III, aVF, and V_1 - V_5 in a patient with atrial septal defect and severe pulmonary hypertension.

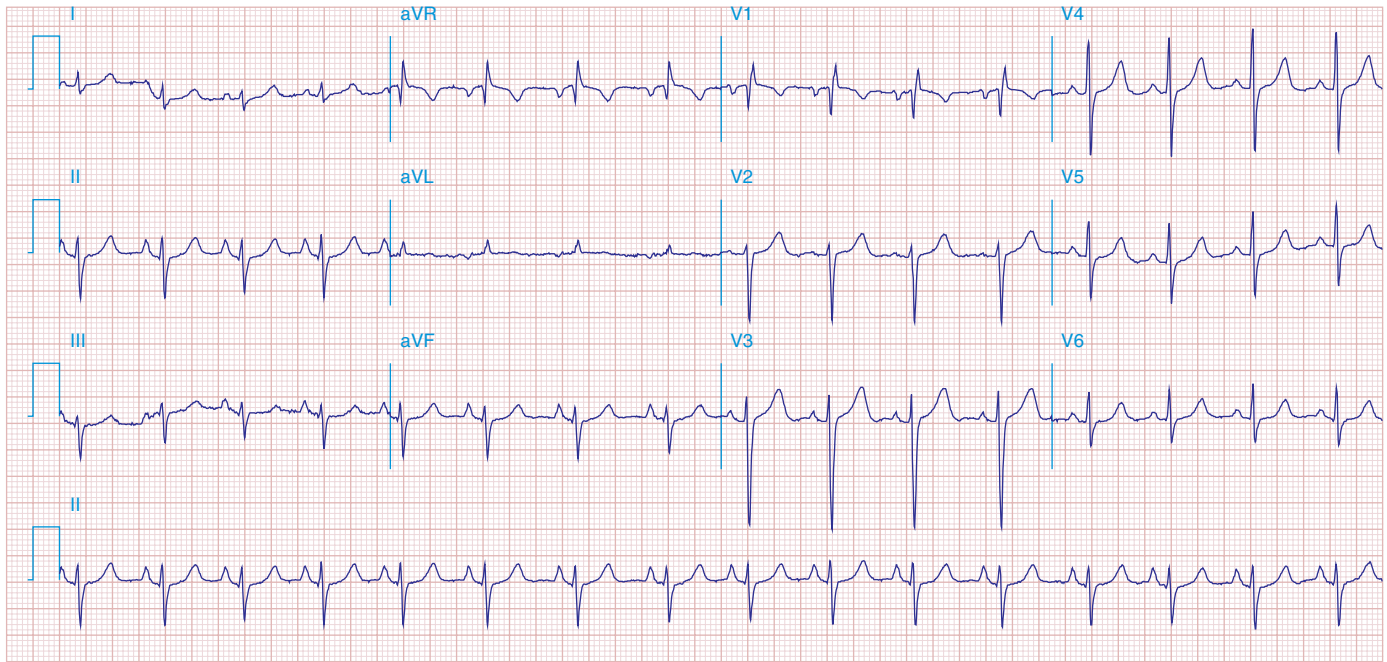


FIGURE 269e-22 Signs of right atrial/RV overload in a patient with chronic obstructive lung disease: (1) peaked P waves in II; (2) QR in V_1 with narrow QRS; (3) delayed precordial transition, with terminal S waves in V_5/V_6 ; (4) superior axis deviation with an S_1 - S_2 - S_3 pattern.

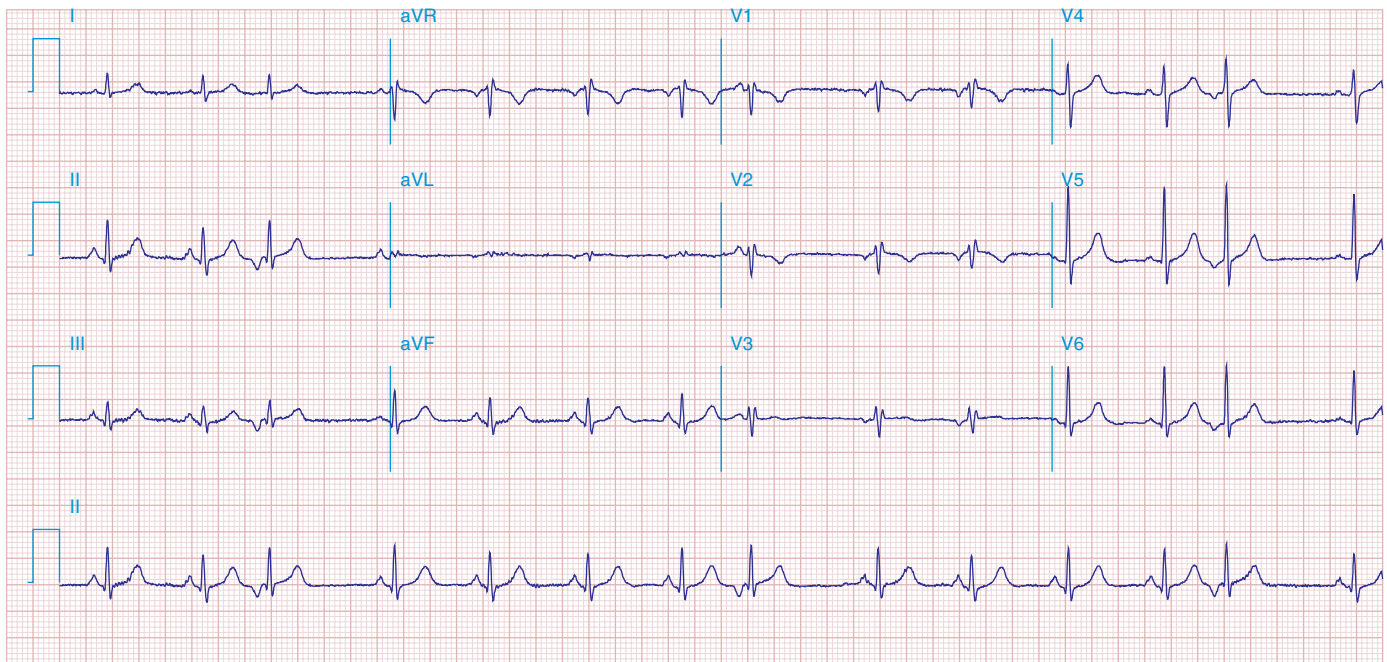


FIGURE 269e-23 (1) Low voltage; (2) incomplete RBBB (rsr' in V_1 - V_3); (3) borderline peaked P waves in lead II with vertical P-wave axis (probable right atrial overload); (4) slow R-wave progression in V_1 - V_3 ; (5) prominent S waves in V_6 ; and (6) atrial premature beats. This combination is seen typically in severe chronic obstructive lung disease.

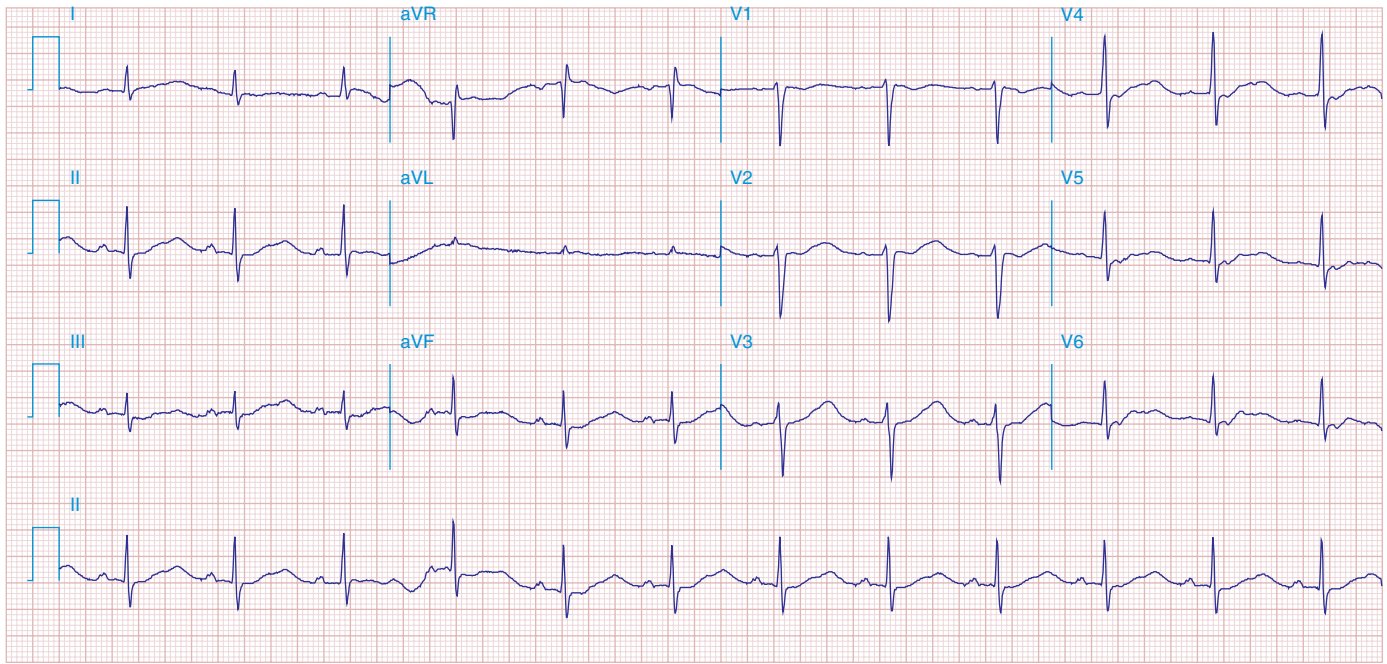


FIGURE 269e-24 Prominent U waves (II, III, and V_4 - V_6) with ventricular repolarization prolongation in a patient with severe hypokalemia.

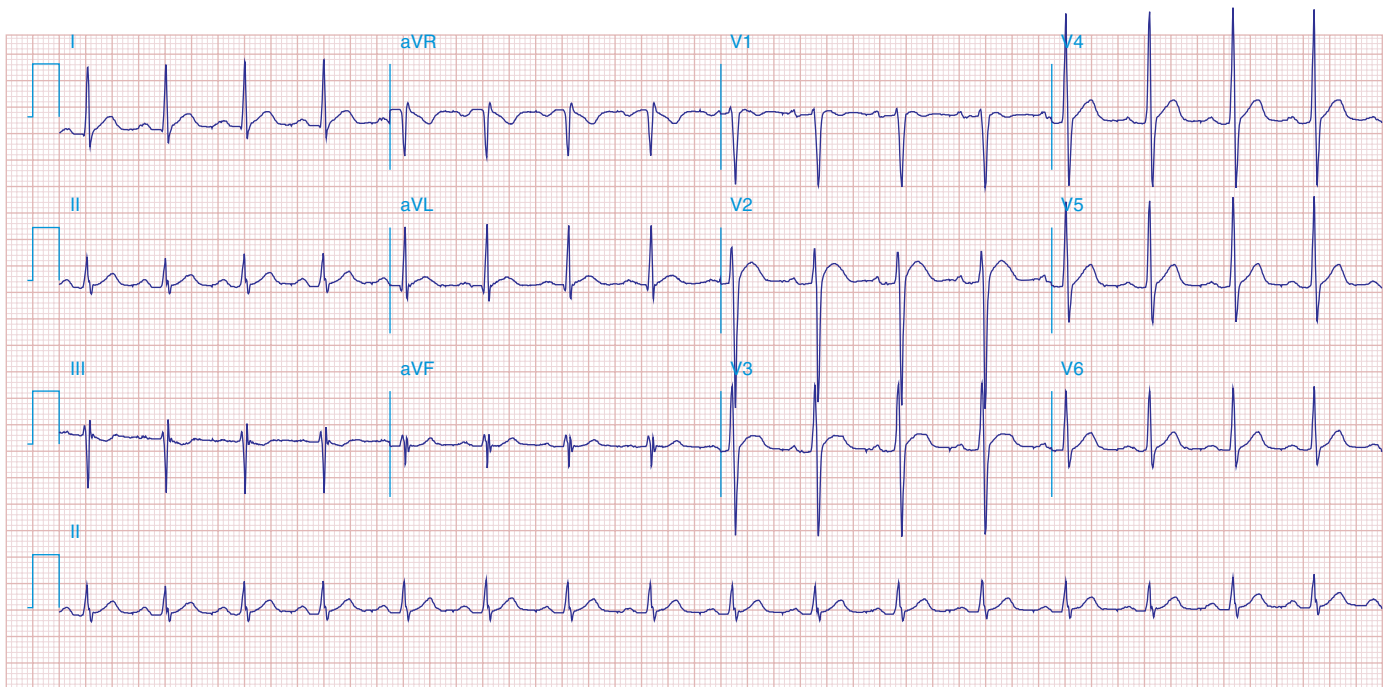


FIGURE 269e-25 Abbreviated ST segment such that the T wave looks like it takes off directly from QRS in some leads (I, V_4 , aVL, and V_2) in a patient with severe hypercalcemia. Note also high takeoff of ST segment in V_2 / V_3 simulating acute ischemia.

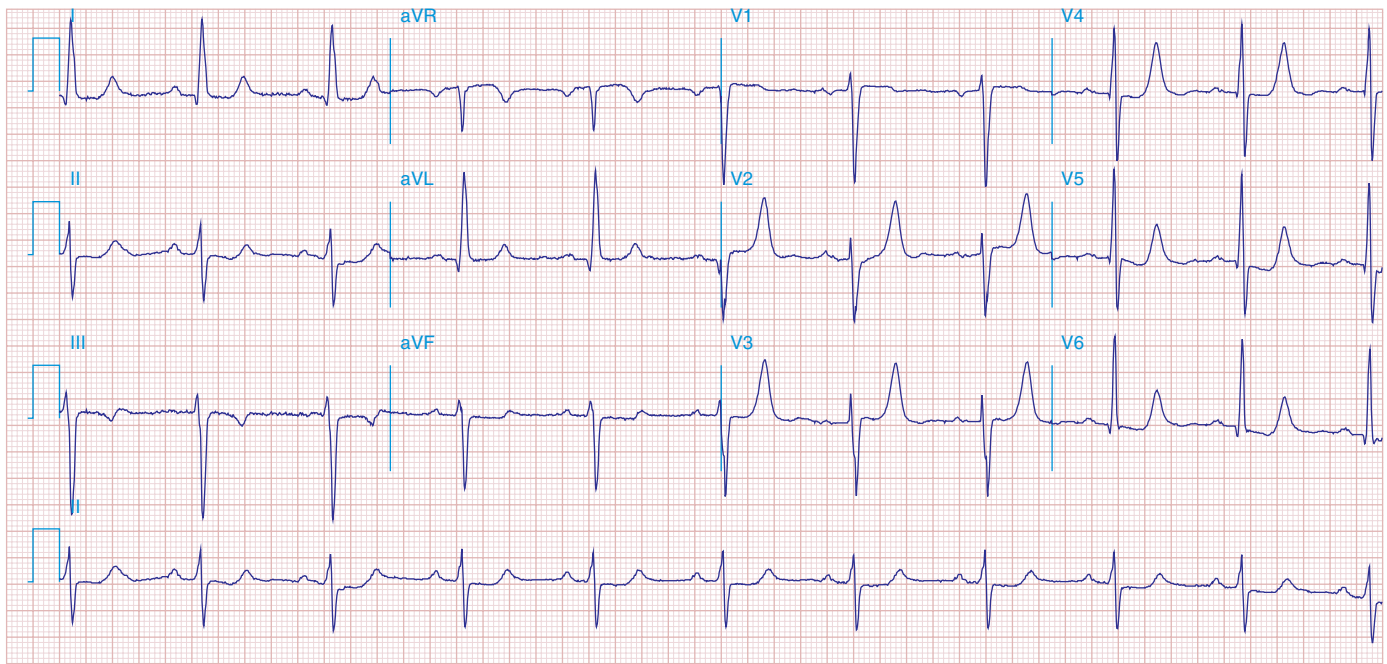


FIGURE 269e-26 SR with LVH, left atrial abnormality, and tall peaked T waves in the precordial leads with inferolateral ST depressions (II, III, aVF, and V₆); left anterior fascicular block and borderline prolonged QT interval in a patient with renal failure, hypertension, and hyperkalemia; prolonged QT is secondary to associated hypocalcemia.

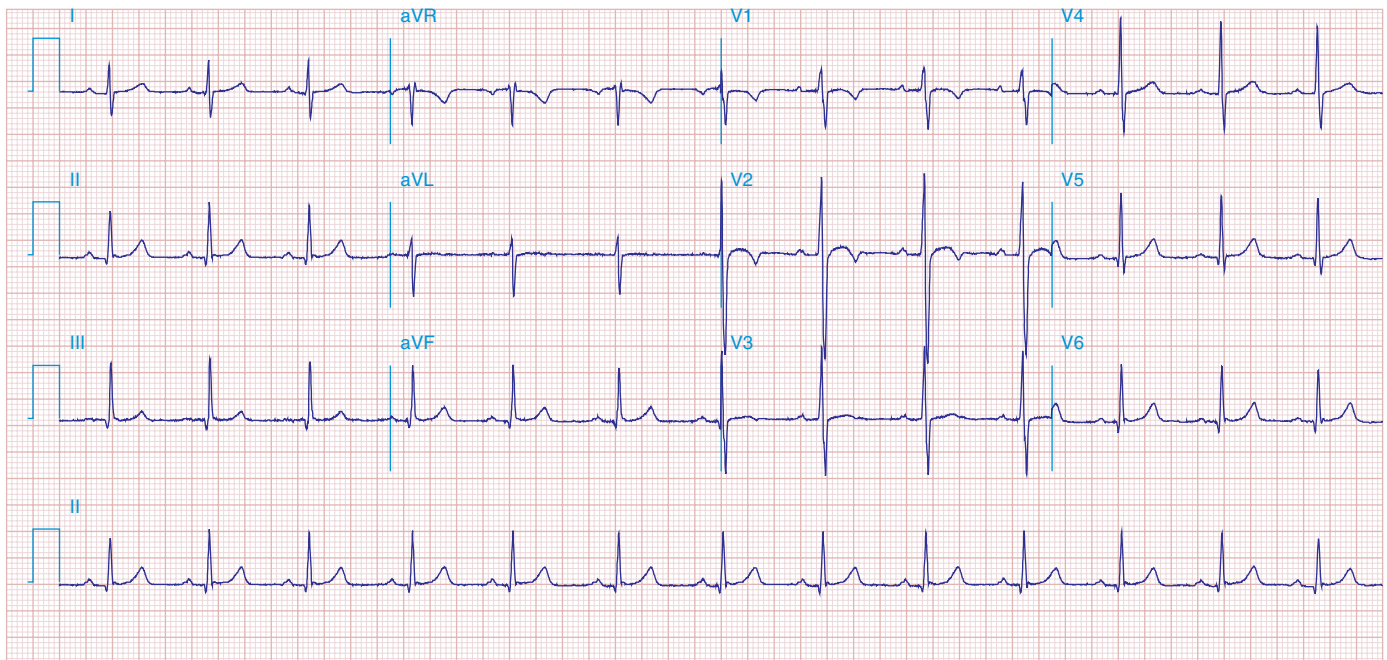


FIGURE 269e-27 Normal ECG in an 11-year-old male. T-wave inversions in V₁-V₃. Vertical QRS axis (+90°) and early precordial transition between V₂ and V₃ are normal findings in children.

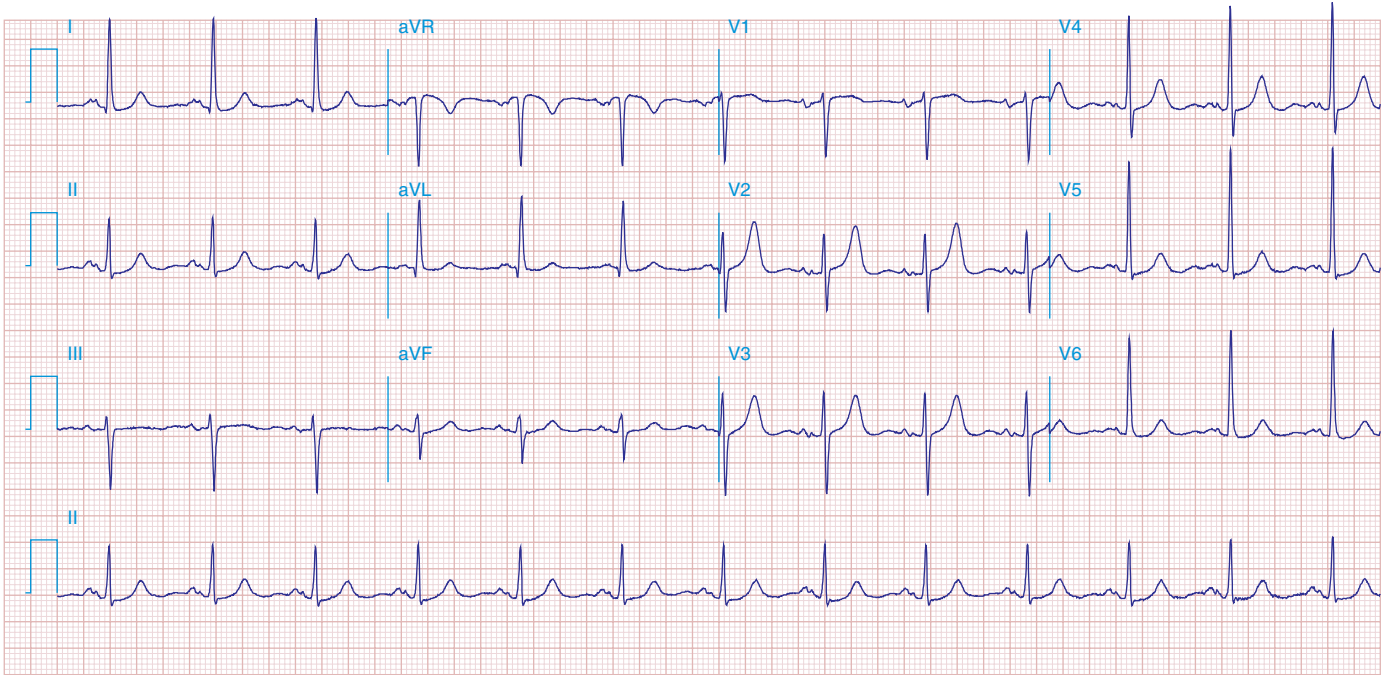


FIGURE 269e-28 Left atrial abnormality and LVH in a patient with long-standing hypertension.

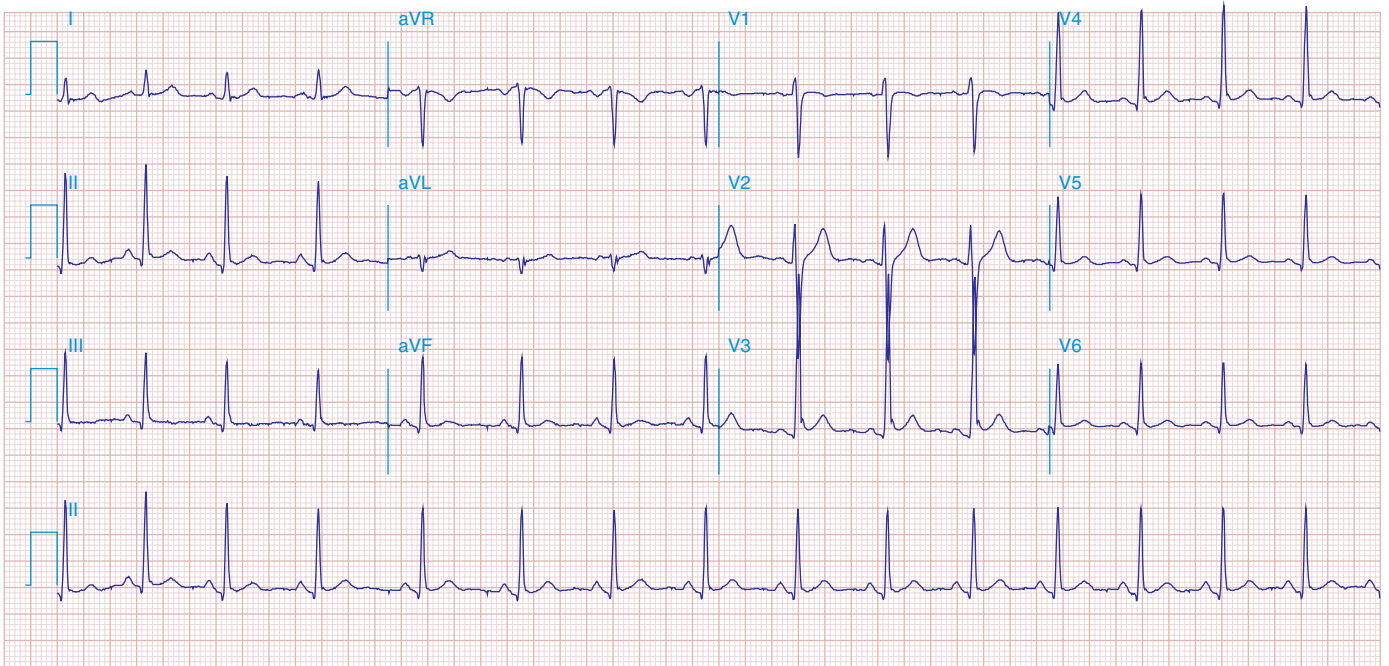


FIGURE 269e-29 Normal variant ST-segment elevations in a healthy 21-year-old male (commonly referred to as *benign early repolarization pattern*). ST elevations exhibit upward concavity and are most apparent in V_3 and V_4 , and less than 1 mm in the limb leads. Precordial QRS voltages are prominent, but within normal limits for a young adult. No evidence of left atrial abnormality or ST depression/T-wave inversions to go along with LVH.

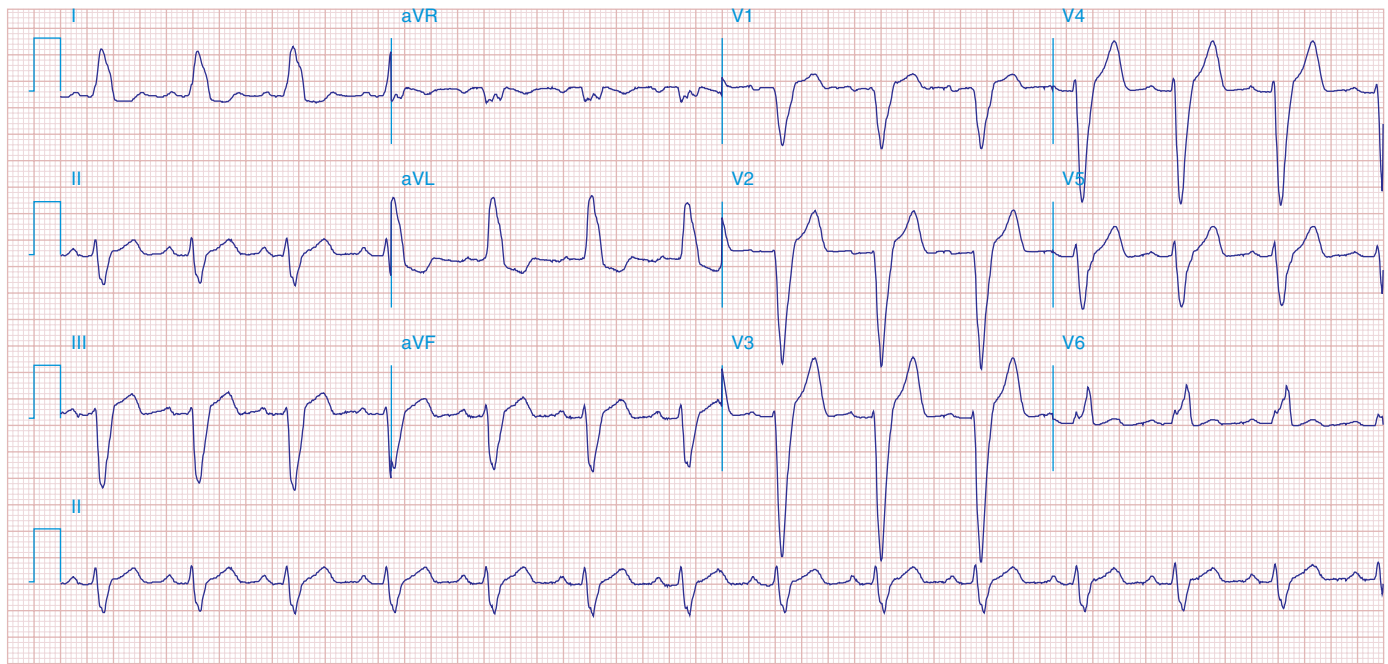


FIGURE 269e-30 SR with first-degree AV “block” (PR interval = 0.24 s) and complete LBBB.

MISCELLANEOUS

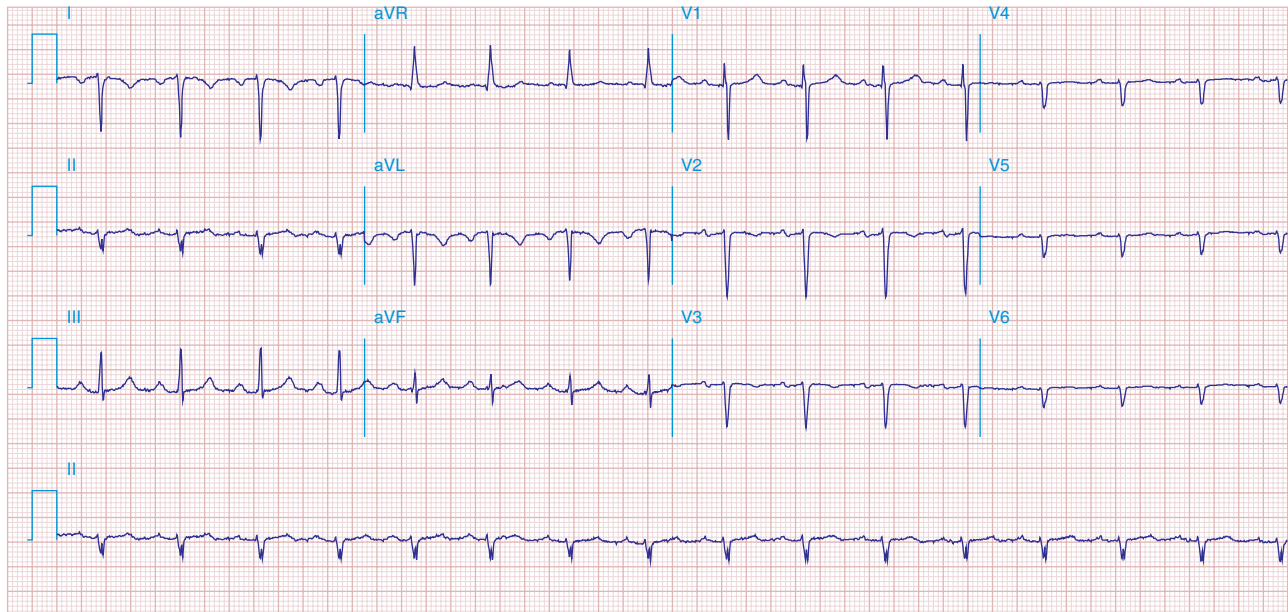


FIGURE 269e-31 Dextrocardia with: (1) inverted P waves in I and aVL; (2) negative QRS complex and T wave in I; and (3) progressively decreasing voltage across the precordium.

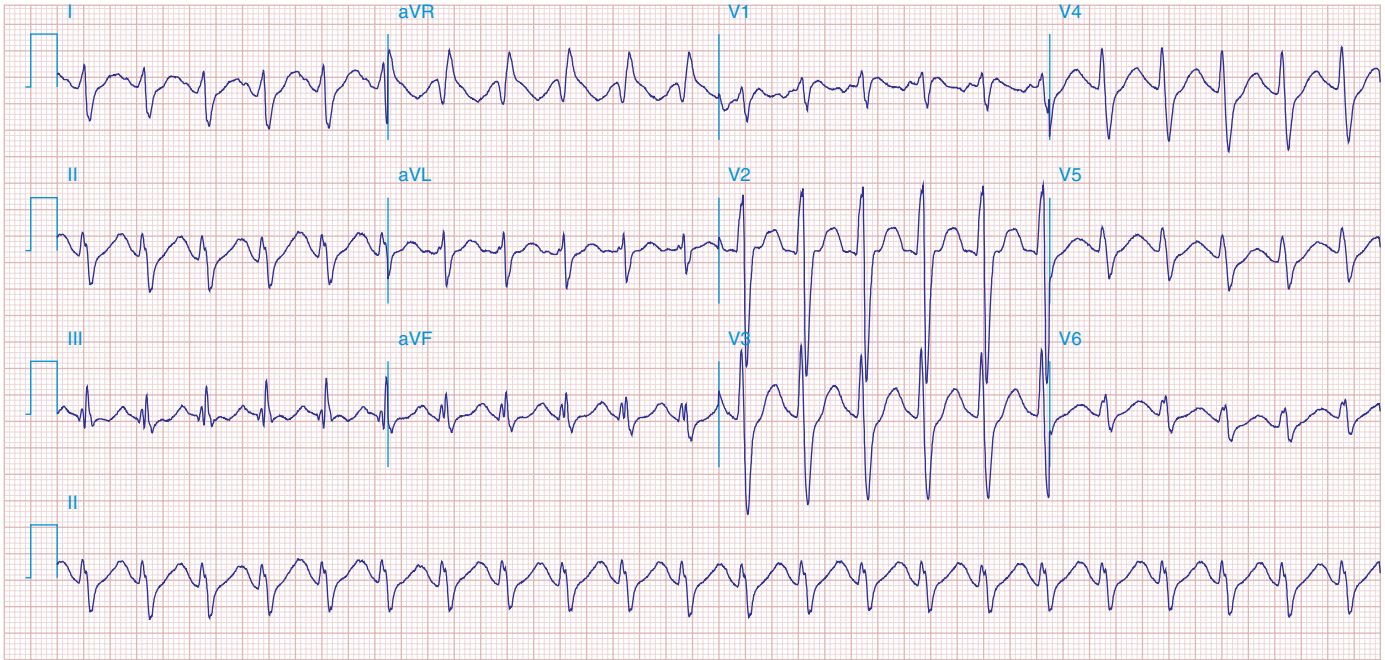


FIGURE 269e-32 Sinus tachycardia; intraventricular conduction delay (IVCD) with a rightward QRS axis. QT interval is prolonged for the rate. The triad of sinus tachycardia, a wide QRS complex, and a long QT in appropriate clinical context suggests tricyclic antidepressant overdose. Terminal S wave (rS) in I and terminal R wave (qR) in aVR are also noted as part of this IVCD variant.

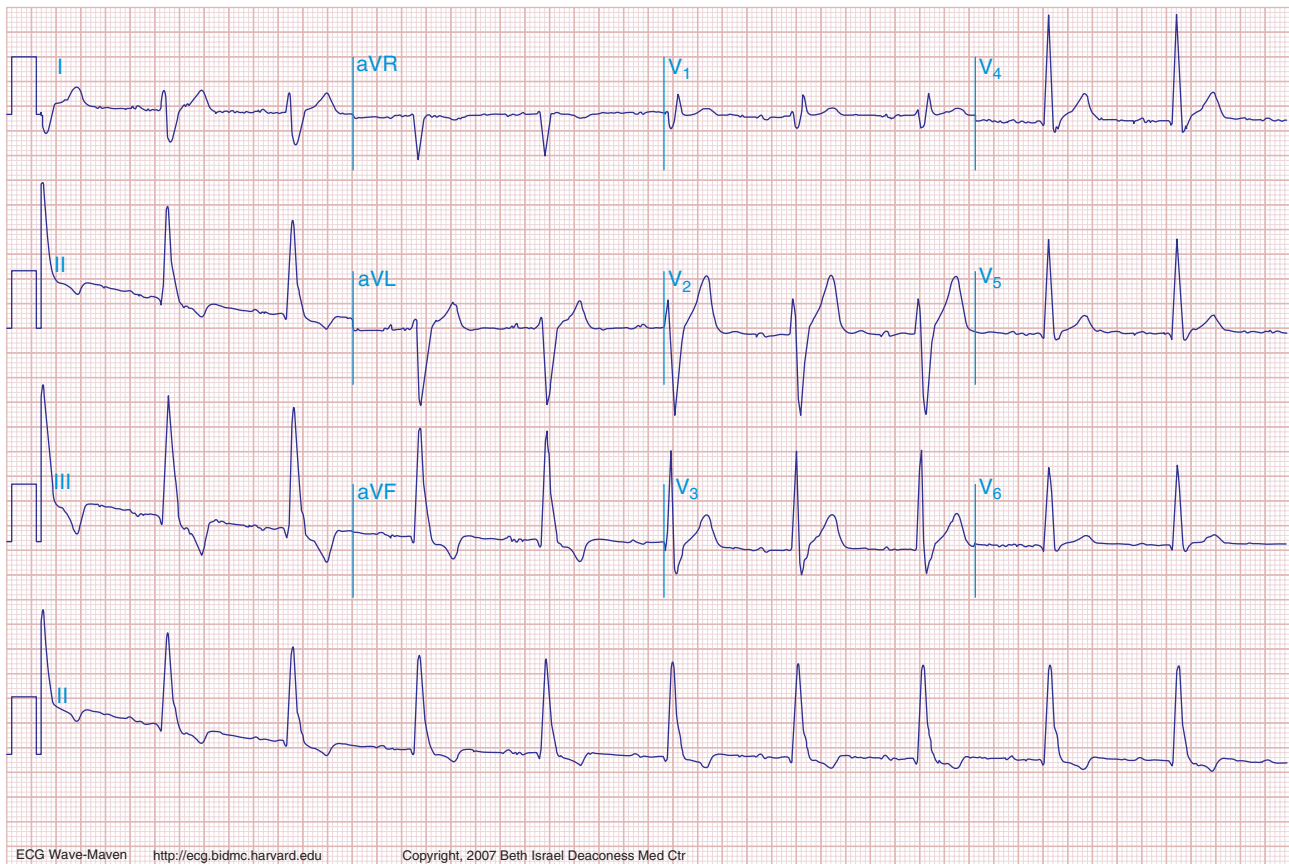


FIGURE 269e-33 Borderline sinus bradycardia (59 beats/min), prolonged PR interval (250 ms), and RBBB are present with marked right-axis deviation (RAD), the latter consistent with left posterior fascicular block (LPFB). LPFB is a diagnosis of exclusion, which requires ruling out lead reversal, normal variant, RV overload syndromes, or lateral MI, in particular, as causes of the RAD. This ECG also shows nondiagnostic Q waves in the inferior leads. In concert with RBBB, the LPFB indicates bifascicular block. (From LA Nathanson et al: ECG Wave-Maven. <http://ecg.bidmc.harvard.edu>.)

Marcelo F. Di Carli, Raymond Y. Kwong,
Scott D. Solomon

The ability to image the heart and blood vessels noninvasively has been one of the greatest advances in cardiovascular medicine since the development of the electrocardiogram. Cardiac imaging complements history taking and physical examination, blood and laboratory testing, and exercise testing in the diagnosis and management of most diseases of the cardiovascular system. Modern cardiovascular imaging consists of echocardiography (cardiac ultrasound), nuclear scintigraphy including positron emission tomography (PET) imaging, magnetic resonance imaging (MRI), and computed tomography (CT). These studies, often used in conjunction with exercise testing, can be used independently or in concert depending on the specific diagnostic needs. In this chapter, we review the principles of each of these modalities and the utility and relative benefits of each for the most common cardiovascular diseases.

PRINCIPLES OF MULTIMODALITY CARDIAC IMAGING

ECHOCARDIOGRAPHY

Echocardiography uses high-frequency sound waves (ultrasound) to penetrate the body, reflect from relevant structures, and generate an image. The basic physical principles of echocardiography are identical to other types of ultrasound imaging, although the hardware and software are optimized for evaluation of cardiac structure and function. Early echocardiography machines displayed “M-mode” echocardiograms in which a single ultrasound beam was displayed over time on a moving sheet of paper (Fig. 270e-1, left panel). Modern

echocardiographic machinery uses phased array transducers that contain up to 512 elements and emit ultrasound in sequence. The reflected ultrasound is then sensed by the receiving elements. A “scan converter” uses information about the timing and magnitude of the reflected ultrasound to generate an image (Fig. 270e-1, right panel). This sequence happens repeatedly in “real time” to generate moving images with frame rates that are typically greater than 30 frames per second, but can exceed 100 frames per second. The gray scale of the image features indicates the intensity of the reflected ultrasound; fluid or blood appears black, and highly reflective structures, such as calcifications on cardiac valves or the pericardium, appear white. Tissues such as myocardium appear more gray, and tissues such as muscle display a unique speckle pattern. Although M-mode echocardiography has largely been supplanted by two-dimensional echocardiography, it is still used because of its high temporal resolution and accuracy for making linear measurements.

The spatial resolution of ultrasound is dependent on the wavelength: the smaller the wavelength and the higher the frequency of the ultrasound beam, the greater are the spatial resolution and ability to discern small structures. Increasing the frequency of ultrasound will increase resolution but at the expense of reduced penetration. Higher frequencies can be used in pediatric imaging or transesophageal echocardiography where the transducer can be much closer to the structures being interrogated, and this is a rationale for using transesophageal echocardiography to obtain higher quality images.

Three-dimensional ultrasound transducers use a waffle-like matrix array transducer and receive a pyramidal data sector. Three-dimensional echocardiography is being increasingly used for assessment of congenital heart disease and valves, although current image quality lags behind two-dimensional ultrasound (Fig. 270e-2).

In addition to the generation of two-dimensional images that provide information about cardiac structure and function, echocardiography can be used to interrogate blood flow within the heart and blood vessels by using the Doppler principle to ascertain the velocity of blood flow. When ultrasound emitted from a transducer reflects off red blood cells that are moving toward the transducer, the reflected ultrasound will return at a slightly higher frequency than emitted; the opposite is true when flow is away from the transducer. That frequency difference, termed the Doppler shift, is directly related to the velocity

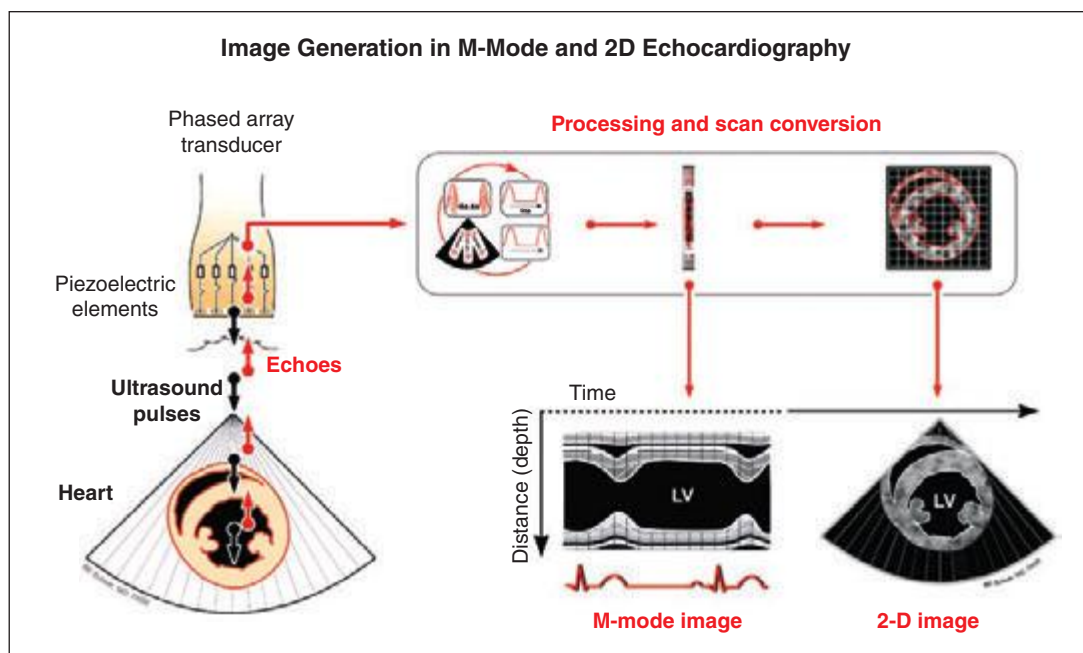


FIGURE 270e-1 Principle of image generation in two-dimensional (2D) echocardiography. An electronically steerable phased-array transducer emits ultrasound from piezoelectric elements, and returning echoes are used to generate a 2D image (right) using a scan converter. Early echocardiography machines used a single ultrasound beam to generate an “M-mode” echocardiogram (see text), although modern equipment generates M-mode echocardiograms digitally from the 2D data. LV, left ventricle.

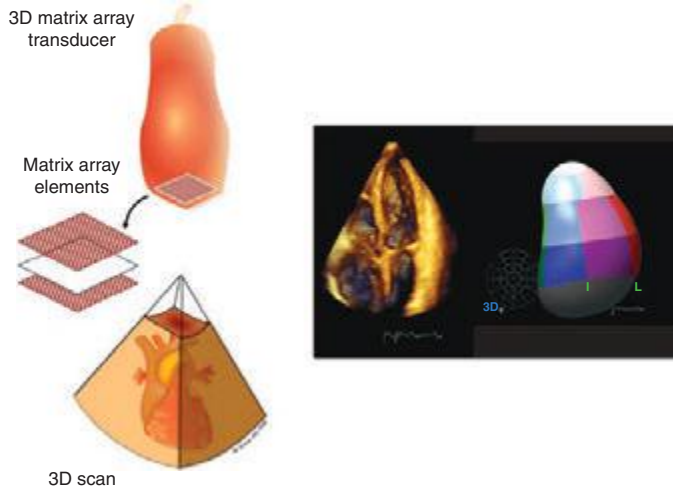


FIGURE 270e-2 Three-dimensional (3D) probe and 3D image.

of the flow of the red blood cells. The velocity of blood flow between two chambers will be directly related to the pressure gradient between those chambers. A modified form of the Bernoulli equation,

$$p = 4v^2$$

where p = the pressure gradient and v = the velocity of blood flow in meters per second, can be used to calculate this pressure gradient in the majority of clinical circumstances. This principle can be used to determine the pressure gradient between chambers and across valves and has become central to the quantitative assessment of valvular heart disease.

There are three types of Doppler ultrasound that are typically used in standard echocardiographic examinations: spectral Doppler, which consists of both pulsed wave Doppler and continuous wave Doppler, and color flow Doppler. Both types of spectral Doppler will display a waveform representing the velocity of blood flow, with time on the horizontal axis and velocity on the vertical axis. Pulsed wave Doppler is used to interrogate relatively low velocity flow and has the ability to determine blood flow velocity at a particular location within the heart. Continuous wave Doppler is used to assess high-velocity flow but can only identify the highest velocity in a particular direction and cannot interrogate the velocity at a specific depth location. Both of these techniques can only accurately assess velocities that are in the direction of the ultrasound scan lines, and velocities that are at an angle to the direction of the ultrasound beam will be underestimated. Color flow Doppler is a form of pulsed wave Doppler in which the velocity of blood flow is color encoded according to a scale and superimposed on a two-dimensional grayscale image in real time, giving the appearance of real-time flow within the heart. The Doppler principle

can also be used to assess the velocity of myocardial motion, which is a sensitive way to assess myocardial function (Fig. 270e-3). A standard full transthoracic echocardiographic examination consists of a series of two-dimensional views made up of different imaging planes from various scanning locations and spectral and color flow Doppler assessment.

Transesophageal echocardiography is a form of echocardiography in which the transducer is located on the tip of an endoscope that can be inserted into the esophagus. This procedure allows closer, less obstructed views of cardiac structures, without having to penetrate through chest wall, muscle, and ribs. Because less penetration is needed, a higher frequency probe can be used, and image quality and spatial resolution are generally higher than with standard transthoracic imaging, particularly for structures that are more posterior. Transesophageal echocardiography has become the test of choice for assessment of small lesions in the heart such as valvular vegetations, especially in the setting of a prosthetic valve disease, and intracardiac thrombi, including assessment of the left atrial appendage, which is difficult to visualize with standard transthoracic imaging, and for assessment of congenital abnormalities. Transesophageal echocardiography requires both topical and systemic anesthesia, generally conscious sedation, and carries additional risks such as potential damage to the esophagus, including the rare possibility of perforation, aspiration, and anesthesia-related complications. Patients generally need to give consent for transesophageal echocardiography and be monitored during and subsequent to the procedure. Transesophageal echocardiography can be carried out in intubated patients and is routinely used for intraoperative monitoring during cardiac surgery.

Stress echocardiography is routinely used to assess cardiac function during exercise and can be used to identify myocardial ischemia or to assess valvular function under exercise conditions. Stress echocardiography is typically performed in conjunction with treadmill or bicycle exercise testing, but can also be performed using pharmacologic stress most typically with an intravenous infusion of dobutamine (see section on stress imaging below).

Whereas typical echocardiographic equipment is large, bulky, and expensive, small hand-held ultrasound equipment developed over the last decade now offers diagnostic quality imaging in a package small enough to be carried on rounds (Fig. 270e-4). These relatively inexpensive point-of-care devices currently lack full diagnostic capabilities but represent an excellent screening tool if used by an experienced operator. As these units become even smaller and less expensive, they are being increasingly used not just by cardiologists, but also by emergency medicine physicians, intensivists, anesthesiologists, and internists.

RADIONUCLIDE IMAGING

Radionuclide imaging techniques are commonly used for the evaluation of patients with known or suspected coronary artery disease (CAD),

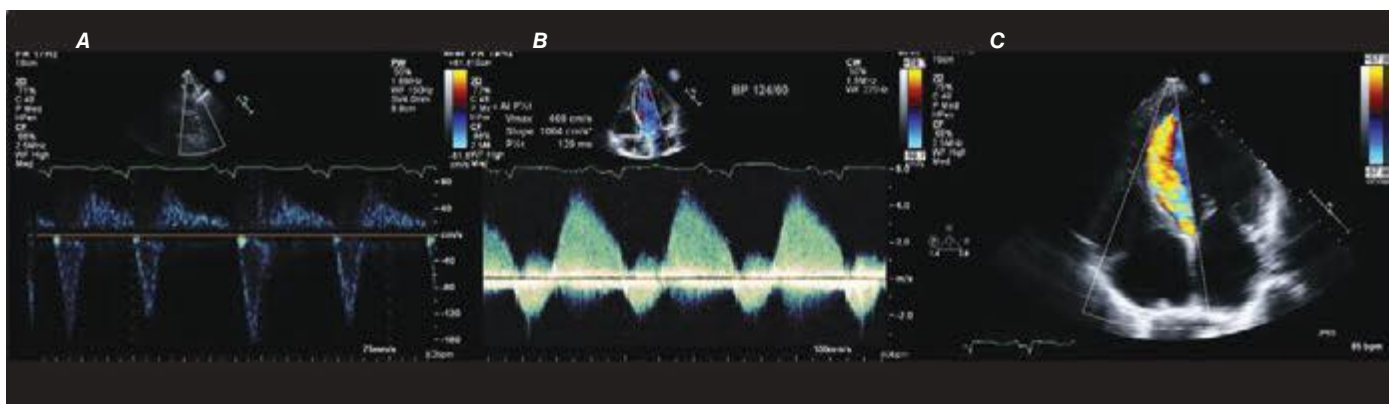


FIGURE 270e-3 Three types of Doppler ultrasound. **A** and **B**. Pulsed and continuous wave Doppler waveforms with time on horizontal axis and velocity of blood flow on vertical axis. **C**. Color flow Doppler, where velocities are encoded by colors according to scale on right side of screen and superimposed on a two-dimensional grayscale image.



FIGURE 270e-4 Two examples of hand-held ultrasound equipment: V-Scan (General Electric, left) and Sonosite (right).

including for initial diagnosis and risk stratification as well as the assessment of myocardial viability. These techniques use small amounts of radiopharmaceuticals (Table 270e-1), which are injected intravenously and trapped in the heart and/or vascular cells. Radioactivity within the heart and vasculature decays by emitting gamma rays. The interaction between these gamma rays and the detectors in specialized scanners (single-photon emission computed tomography [SPECT] and PET) creates a scintillation event or light output, which can be captured by digital recording equipment to form an image of the heart and vasculature. Like CT and MRI, radionuclide images also generate tomographic (three-dimensional) views of the heart and vasculature.

Radiopharmaceuticals Used in Clinical Imaging Table 270e-1 summarizes the most commonly used radiopharmaceuticals in clinical SPECT and PET imaging.

Protocols for Stress Myocardial Perfusion Imaging Both exercise and pharmacologic stress can be used for myocardial perfusion imaging. Exercise stress is generally preferred because it is physiologic and provides additional clinically important information (i.e., clinical and hemodynamic responses, ST-segment changes, exercise duration, and functional status). However, submaximal effort will lower the sensitivity of the test and should be avoided, especially if the test is requested for initial diagnosis of CAD. In patients who are unable to exercise or who exercise submaximally, pharmacologic stress offers an adequate alternative to exercise stress testing. Pharmacologic stress can be accomplished either with coronary vasodilators, such as adenosine, dipyridamole, or regadenoson, or β_1 -receptor agonists, such as dobutamine. For patients unable to exercise, vasodilators are the most commonly used stressors in combination with myocardial perfusion imaging. Dobutamine is a potent β_1 -receptor agonist that increases myocardial oxygen demand by augmenting contractility, heart rate, and blood pressure similar to exercise. It is generally used as an alternative to vasodilator stress in patients with chronic pulmonary disease, in whom vasodilators may be contraindicated. Dobutamine is also

commonly used as a pharmacologic alternative to stress testing in stress echocardiography.

Myocardial Perfusion and Viability Imaging Protocols

Imaging protocols are tailored to the individual patient based on the clinical question, patient's risk, ability to exercise, body mass index, and other factors.

For SPECT imaging, technetium-99m (^{99m}Tc)-labeled tracers are the most commonly used imaging agents because they are associated with the best image quality and the lowest radiation dose to the patient (Fig. 270e-5). Selection of the protocol (stress-only, single-day, or 2-day) depends on the patient and clinical question. After intravenous injection, myocardial uptake of ^{99m}Tc -labeled tracers is rapid (1–2 min). After uptake, these tracers become trapped intracellularly in mitochondria and show minimal change over time. This is why ^{99m}Tc tracers can be helpful in patients with chest pain of unclear etiology occurring at rest, because patients can be injected while having chest pain and imaged some time later after symptoms subside. Because the radiotracer is trapped at the

time of injection, the images provide a snapshot of myocardial perfusion at the time of injection, even if the acquisition is delayed. Indeed, a normal myocardial perfusion study following a rest injection in a patient with active chest pain effectively excludes myocardial ischemia as the cause of chest pain (high negative predictive value). While used commonly in the past for perfusion imaging, thallium-201 protocols are now rarely used because they are typically associated with a higher radiation dose to the patient.

PET myocardial perfusion imaging is an alternative to SPECT and is associated with improved diagnostic accuracy and lower radiation dose to patients due to the fact that radiotracers are typically short lived (Table 270e-1). The ultra-short half-life of some PET radiopharmaceuticals in clinical use (e.g., rubidium-82) is the primary reason why imaging is generally combined with pharmacologic stress, as opposed to exercise, because this allows for faster imaging of these rapidly decaying radiopharmaceuticals. However, exercise is possible for relatively longer lived radiotracers (e.g., ^{13}N -ammonia). PET imaging protocols are typically faster than SPECT, but more expensive. For myocardial perfusion imaging, rubidium-82 does not require an on-site medical cyclotron (it is available from a strontium-82/rubidium-82 generator) and, thus, is the most commonly used radiopharmaceutical. ^{13}N -ammonia has better flow characteristics (higher myocardial extraction) and imaging properties than rubidium-82, but it does require an on-site medical cyclotron. In comparison to SPECT, PET has improved spatial and contrast resolution and provides absolute measures of myocardial perfusion (in mL/min per gram of tissue), thereby providing the patients' regional and global coronary flow reserve. The latter helps improve diagnostic accuracy and risk stratification, especially in obese patients, women, and higher risk individuals (e.g., diabetes mellitus) (Fig. 270e-6). Contemporary PET and SPECT scanners are combined with a CT scanner (so-called *hybrid PET/CT* and *SPECT/CT*). CT is used primarily to guide patient positioning in the field of view and for correcting inhomogeneities in radiotracer distribution due to attenuation by soft tissues (so-called *attenuation*

TABLE 270e-1 RADIOPHARMACEUTICALS FOR CLINICAL NUCLEAR CARDIOLOGY

Radiopharmaceutical	Imaging Technique	Physical Half-Life	Application
Technetium-99m sestamibi	SPECT	6 h	Myocardial perfusion imaging
Technetium-99m tetrofosmin	SPECT	6 h	Myocardial perfusion imaging
Thallium-201	SPECT	72 h	Myocardial perfusion imaging
Iodine-123 metaiodobenzylguanidine (MIBG)	SPECT	13 h	Cardiac sympathetic innervation
Rubidium-82	PET	76 s	Myocardial perfusion imaging
^{13}N -ammonia	PET	10 min	Myocardial perfusion imaging
^{18}F -fluorodeoxyglucose	PET	110 min	Myocardial viability and inflammation imaging

Abbreviations: PET, positron emission tomography; SPECT, single-photon emission computed tomography.

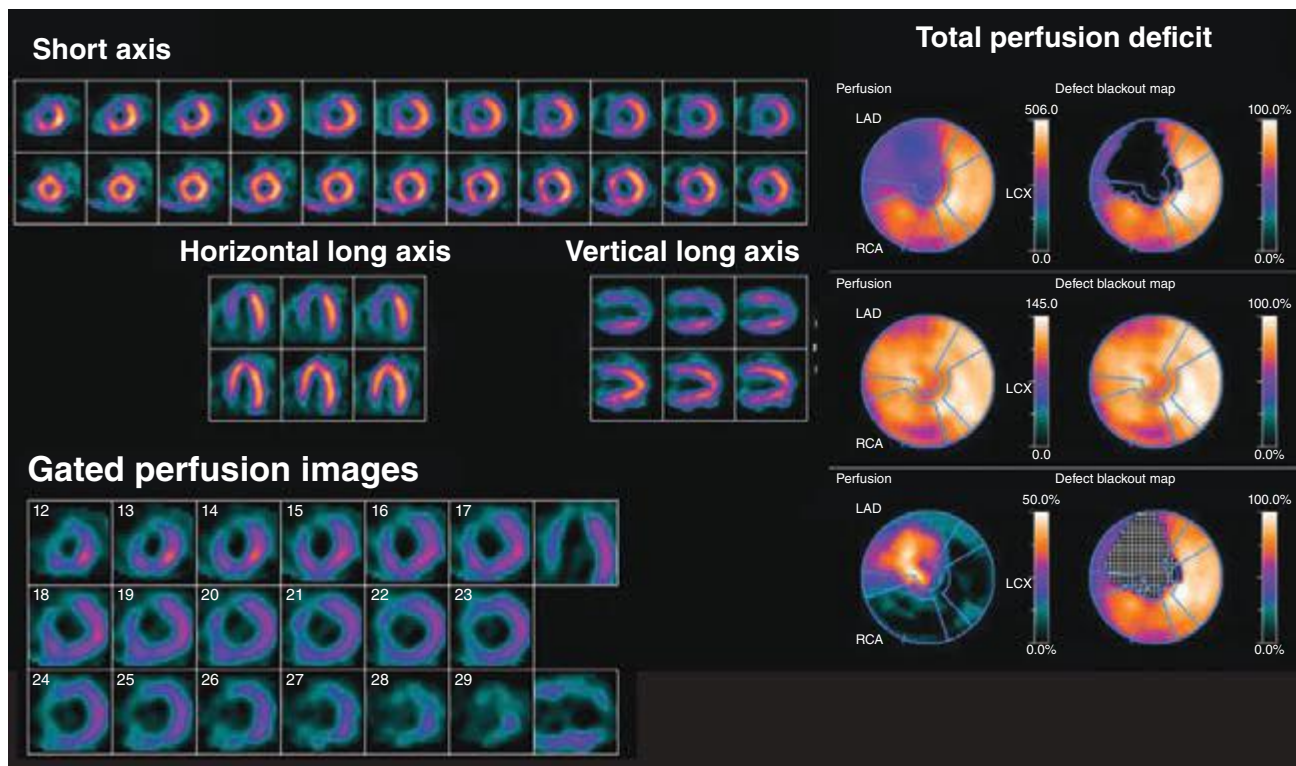


FIGURE 270e-5 Tomographic stress (top of each pair) and rest myocardial perfusion images with technetium-99m sestamibi single-photon emission computed tomography imaging demonstrating a large perfusion defect throughout the anterior and anteroseptal walls. The right panel demonstrates the quantitative extent of the perfusion abnormality at stress (top bull's-eye), at rest (middle bull's-eye), and the extent of defect reversibility (lower bull's-eye). The lower left panel demonstrates electrocardiogram-gated myocardial perfusion images from which one can determine the presence of regional wall motion abnormalities and calculate left ventricular volumes and ejection fraction.

correction). However, it can also be used to obtain diagnostic data including coronary artery calcium score and/or CT coronary angiography (discussed below).

For the evaluation of myocardial viability in patients with ischemic cardiomyopathy, myocardial perfusion imaging (with SPECT or PET) is usually combined with metabolic imaging (i.e., fluorodeoxyglucose [FDG] PET). In hospital settings lacking access to PET scanning, thallium-201 SPECT imaging is an excellent alternative.

CARDIAC COMPUTED TOMOGRAPHY

CT acquires images by passing a thin x-ray beam through the body at many angles to generate cross-sectional images. The x-ray transmission measurements are collected by a detector array and digitized into pixels that form an image. The grayscale information in individual pixels is determined by the attenuation of the x-ray beam along its path by tissues of different densities, referenced to the value for water in units known as Hounsfield units. In the resulting CT images, bone appears bright white, air is black, and blood and muscle show varying shades of gray. However, due to the limited contrast between cardiac chambers and vascular structures, iodinated contrast agents are necessary for most cardiovascular indications. Cardiac CT produces tomographic images of the heart and surrounding structures. With modern CT scanners, a three-dimensional dataset of the heart can be acquired in 5–15 s with submillimeter spatial resolution.

CT Calcium Scoring CT calcium scoring is the simplest application of cardiac CT and does not require administration of iodinated contrast. The presence of coronary artery calcification has been associated with increased burden of atherosclerosis and cardiovascular mortality. Coronary calcium is then quantified (e.g., Agatston score) and categorized as minimal (0–10), mild (10–100), moderate (100–400), or severe (>400) (Fig. 270e-7). Coronary artery calcium (CAC) scores are then normalized by age and gender and reported as percentile scores. Population-based studies in asymptomatic cohorts have reported high cardiac prognostic

value of CT calcium score. With appropriate techniques, the radiation dose associated with CAC scanning is very low (~1–2 mSv).

CT Coronary Angiography Coronary CT angiography (CTA) is emerging as a viable alternative to coronary angiography in selected patients. Imaging of the coronary arteries by CT is challenging because of their small luminal size and because of cardiac and respiratory motion. Respiratory motion can be reduced by breath-holding, and cardiac motion is best reduced by slowing the patient's heart rate, ideally to under 60 beats/min, using intravenous or oral beta blockade or other rate-lowering drugs. When performing a coronary CTA, image quality is further enhanced using sublingual nitroglycerin to enlarge the coronary lumen just prior to contrast injection. Imaging the whole-heart volume is synchronized to the administration of weight-based and appropriately timed intravenous iodinated contrast. Image acquisition is linked to the timing of the cardiac cycle through electrocardiogram (ECG) triggering. Prospective ECG triggering, whereby the x-ray beam is turned on during a specific part of the cardiac cycle (e.g., end systole, combined end-systolic and end-diastolic timing, or mid-diastole), is generally used to limit the radiation exposure to the patient by acquiring data only through that portion of the cardiac cycle with least motion. Dose modulation is another method that should be routinely used to reduce radiation when performing CTA. It delivers a maximal amount of x-ray during the portion of the cardiac cycle of interest, but reduces x-ray delivery throughout the remaining portion of the cardiac cycle. The resulting images are then postprocessed using a three-dimensional workstation, which facilitates interpretation of the coronary anatomy and estimation of the severity of atherosclerosis (Fig. 270e-7).

CARDIAC MAGNETIC RESONANCE

Cardiac magnetic resonance (CMR) imaging is based on imaging of protons in hydrogen. Hydrogen is abundant because 80% of the human body consists of water. When put inside the MRI scanner,

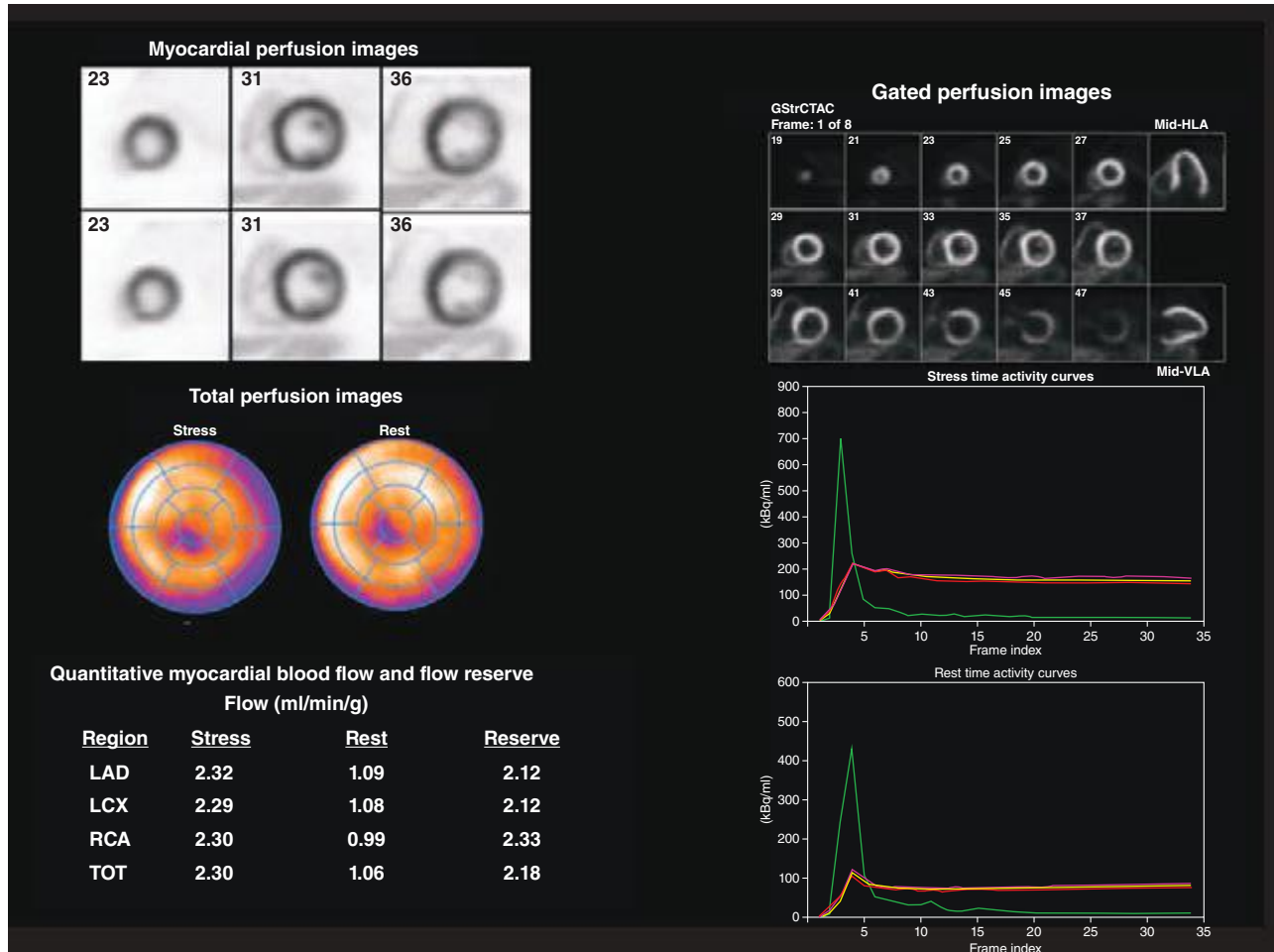


FIGURE 270e-6 Multidimensional cardiac imaging protocol with positron emission tomography. The *left upper panel* demonstrates stress and rest short-axis images of the left and right ventricles demonstrating normal regional myocardial perfusion. The *middle panel* demonstrates the quantitative bull's-eye display to evaluate the extent and severity of perfusion defects. The *lower right panel* illustrates the time-activity curves for quantification of myocardial blood flow. The *right upper panel* demonstrates electrocardiogram-gated myocardial perfusion images from which one can determine the presence of regional wall motion abnormalities and calculate left ventricular volumes and ejection fraction. LAD, left anterior descending artery; LCX, left circumflex artery; RCA, right coronary artery; TOT, total left ventricle.

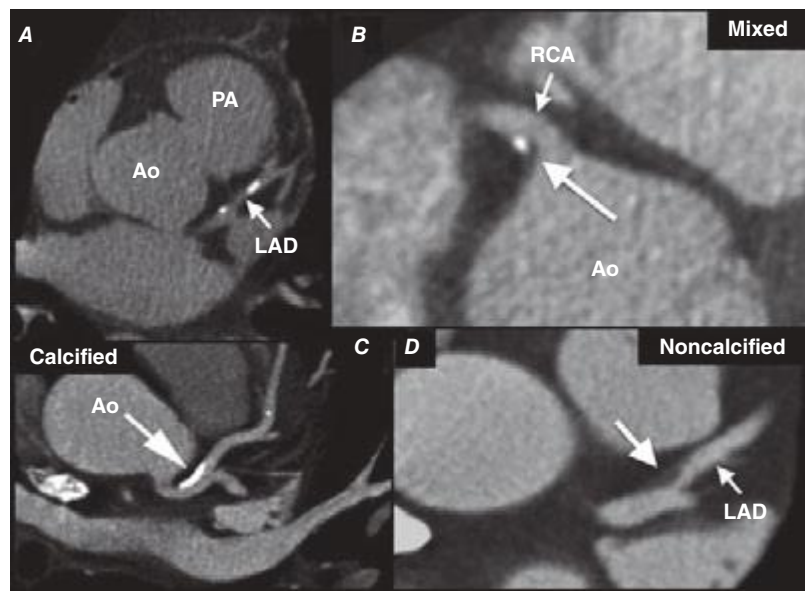


FIGURE 270e-7 Examples of non-contrast- and contrast-enhanced coronary imaging with computed tomography (CT). **A**. Calcified coronary plaques in the distal left main and proximal left anterior descending coronary artery (LAD) in a noncontrast cardiac CT scan. Calcium deposits are dense and present as bright white structures on CT, even without contrast enhancement. **B**, **C**, and **D**. Different types of atherosclerotic plaques on contrast-enhanced CT scans. Importantly, noncalcified plaques are evident only on contrast-enhanced CT scans. AO, aorta; PA, pulmonary artery; RCA, right coronary artery.

TABLE 270e-2 CLINICAL CARDIAC MAGNETIC RESONANCE PULSE SEQUENCES AND THEIR APPLICATION

Pulse Sequence	Key Imaging Interests
Cardiac Morphology	
Still frame imaging (black or bright blood)	Cardiac structures
Cardiac Function	
Cine imaging	Left ventricular volume and function
Cine myocardial tagging	Left ventricular deformation (strain)
Blood Flow Imaging	
Velocity-encoded phase contrast	Cardiac and great vessel flow
Stress Testing	
Myocardial perfusion imaging	Regional myocardial blood flow
Cine imaging	Regional wall motion
Myocardial Tissue Characterization	
Late gadolinium enhancement	Myocardial infarction and infiltrative disease
T2-weighted imaging	Myocardial edema
Iron content imaging	Myocardial iron infiltration
Magnetic Resonance Angiography	
Aorta, peripheral and coronary arteries	Luminal stenosis and vessel wall remodeling

the magnetic field causes the protons (spins) to spin around their axis (a process known as precession) at specific frequencies. Spins within water have a different frequency than spins within more complex macromolecules such as fat or protein. Inside the MRI, a set of gradient coils slightly modifies the magnetic field in each of the three orthogonal directions. As a result, this additional process alters the frequencies of spins, and now the spins can be spatially located inside the MRI bore. This system allows the MRI to selectively deposit radiofrequency energy (in the form of radiofrequency pulse) into specific locations of the body for the purpose of imaging those locations. Once the radiofrequency pulses stop, the energy absorbed by the body will quickly be released back. Using the proper arrangement of surface phased-array coils, this released energy can be read, and important information such as spin locations and frequencies can be digitally recorded in a data matrix known as the K-space, before reconstructed into a magnetic resonance image. Radiofrequency energy deposition into the patient's body can be arranged in many complex ways, known as pulse sequences, that allow extraction of different types of information from the body regions of interests. In CMR, these types of information in general are categorized under T1, T2, or T2* weighting, each of them containing a different combination of diagnostic information regarding cardiac structure, tissue characteristics, blood flow, or other physiologic properties of the heart.

In clinical CMR, most pulse sequences are T1-weighted sequences, which characterize cardiac structure and function, blood flow, and myocardial perfusion, whereas T2-weighted and T2*-weighted pulse sequences characterize myocardial edema and myocardial iron infiltration, respectively. A combination of more than one weighting is possible in some pulse sequences. ECG-triggered cine CMR is the modality that serves as a reference standard for quantifying ventricular volumes and function. Respiratory motion during CMR imaging is suppressed most commonly using repetitive patient breath-holding, but more advanced algorithms such as motion averaging or gating diaphragmatic motion (known as navigator guidance) are also used in clinical CMR. A list of common pulse sequence used in CMR is shown in [Table 270e-2](#).

ASSESSMENT OF CARDIAC STRUCTURE AND FUNCTION

Echocardiography, CMR, and cardiac CT are all capable of assessing cardiac structure and function, although echocardiography is generally considered the primary imaging method for these assessments. Radionuclide imaging can also be used to assess left ventricular

regional and global systolic function. Echocardiography is most often used to assess the size of all four chambers and thickness of ventricular walls, which are affected by both cardiac and systemic diseases.

The structure of the left ventricle is generally assessed by determining its volume and mass. Left ventricular volumes can be easily estimated from two-dimensional echocardiography by using one of several validated methods. The accuracy of these methods by echocardiography is limited by the fact that, as a nontomographic technique, foreshortening of the imaging plane can lead to underestimation of volumes. Moreover, virtually all of these methods require accurate identification of the endocardial border, which is dependent on image quality. In this regard, high-resolution tomographic techniques such as CMR or cardiac CT are considered generally more accurate for volumetric assessment. Three-dimensional echocardiography has several advantages over two-dimensional echocardiography by not requiring any geometric assumptions about the left ventricle for quantification of volumes and ejection fraction. However, acquisition of three-dimensional echocardiographic images requires substantial expertise, and these techniques are not widely used in practice.

Left ventricular dilatation is common to a number of cardiac diseases. For example, regional dysfunction secondary to myocardial infarction can ultimately lead to progressive ventricular dilatation or remodeling. Although dilatation often begins in the region affected by the infarction, subsequent compensatory dilatation can occur in remote myocardial regions as well. The presence of regional wall motion abnormalities associated with ventricular thinning (reflecting scar) in a coronary distribution is strongly suggestive of an ischemic etiology. Direct assessment of infarcted myocardium is possible with both CMR (evident as areas of late gadolinium enhancement [LGE]) and radionuclide imaging (as assessed by regional perfusion or metabolic defects at rest). CMR can be particularly useful in determining etiology of ventricular dilatation and dysfunction, with LGE in coronary distributions being nearly pathognomonic for infarction ([Video 270e-1](#)).

More global ventricular dilatation is seen in cardiomyopathy and dilatation due to valvular heart disease. Idiopathic, nonischemic cardiomyopathies will typically result in global ventricular dilatation and dysfunction, with thinning of the walls. Patients with substantial ventricular dyssynchrony due to conduction abnormalities will have a typical pattern of contraction (e.g., delay of contraction of the lateral wall with left bundle branch block). Although various methods for determining ventricular dyssynchrony have been proposed as ways to identify patients who would benefit from cardiac resynchronization therapy, it is not yet clear that they are superior to ECG assessment of QRS duration and morphology. As discussed later in this chapter, regurgitant lesions of either the mitral or aortic valves can lead to substantial ventricular dilatation, and assessment of ventricular size is integral in the evaluation and timing of surgical correction. Because changes in ventricular size are used clinically to determine which patients should undergo valve surgery, accurate assessment of changes in ventricular size is essential. Although serial echocardiography can provide these data, serial assessment by CMR may be more accurate when appreciation of subtle changes over time is important.

Left ventricular wall thickness and mass are also important measures of cardiac and systemic disease. The left ventricle will hypertrophy under any condition in which its afterload is increased, including conditions that obstruct outflow, such as aortic stenosis, hypertrophic cardiomyopathy, and subaortic membranes; in postcardiac aortic obstruction seen in coarctation; or in systemic conditions characterized by increased afterload, such as hypertension. The pattern of ventricular hypertrophy can change depending on the etiology. Aortic stenosis and hypertension are typically characterized by concentric hypertrophy, in which the ventricular walls thicken "concentrically" and cavity size is usually small. In volume overload conditions such as mitral or aortic regurgitation, there may be minimal increase in ventricular wall thickness, but substantial ventricular dilatation leads to marked increases in left ventricular mass.

Ventricular wall thickness can be measured and ventricular mass can be calculated by either echocardiography or CMR. Although radionuclide imaging and cardiac CT can also provide measures of

left ventricular mass, they are not generally used for this purpose. Although measurement of wall thickness with echocardiography is relatively straightforward and accurate, determining left ventricular mass by echocardiography requires using one of several formulas that takes into account both wall thickness and ventricular cavity dimensions. Assessment of left ventricular mass by CMR has the advantage of not requiring geometric assumptions and is thus more accurate than echocardiography.

ASSESSMENT OF LEFT VENTRICULAR SYSTOLIC FUNCTION

Assessment of ejection fraction, or the percentage of blood ejected with each beat, has been the primary method to assess systolic function and is generally calculated by subtracting end-systolic volume from end-diastolic volume and dividing by end-diastolic volume. All cardiac imaging modalities can provide direct measurements of left ventricular ejection fraction (LVEF). As discussed above, tomographic techniques (e.g., CMR, CT, and radionuclide imaging [SPECT and PET]) are generally more accurate and reproducible than echocardiography because there are no geometric assumptions. A LVEF of 55% or greater is generally considered normal, and an LVEF of 50–55% is considered in the low-normal range.

Newer methods to assess systolic function, such as myocardial strain or deformation imaging using speckle-tracking methods on echocardiography or myocardial tagging on CMR, can provide a more sensitive approach to detection of systolic dysfunction. Additional assessments based on these novel methods include assessment of myocardial twist and torsion. Although these techniques are not used routinely, they may be especially useful in certain conditions such as valvular heart disease and early detection of cardiotoxicity following chemotherapy and/or radiation therapy. In addition to estimation or calculation of ejection fraction, stroke volume can be assessed by virtually all cardiac imaging methods, generally by subtracting the end-systolic volume from the end-diastolic volume, or by Doppler methods (only on echocardiography), and offers another measure of systolic function that provides independent information from ejection fraction.

ASSESSMENT OF LEFT VENTRICULAR DIASTOLIC FUNCTION

Echocardiography remains the primary method for clinical assessment of diastolic function. Recent advances in Doppler tissue imaging allow for accurate assessment of the velocity of myocardial wall motion by assessing the excursion of the mitral annulus in diastole. Mitral annular relaxation velocity, or E' , is inversely related to the time constant of relaxation, tau, and has been shown to have prognostic significance. Dividing the standard mitral inflow maximal velocity, E , by the mitral annular relaxation velocity yields E/E' , which has been shown to correlate with left ventricular filling pressures. The utility of standard E and A wave ratios for assessment of diastolic function has been questioned. Mitral deceleration time can be a useful measure if very short (<150 ms), suggesting restrictive physiology and severe diastolic dysfunction. Several grading methods for diastolic function have been proposed that take into account a number of diastolic parameters, including Doppler tissue-based relaxation velocities, pulmonary venous Doppler, and left atrial size (Fig. 270e-8). Diastolic function worsens with aging, and most diastolic parameters need to be adjusted for age.

ASSESSMENT OF RIGHT VENTRICULAR FUNCTION

Right ventricular size and function have been shown to be prognostically important in a variety of conditions. Right ventricular size and function can be assessed by echocardiography, CMR, CT, or radionuclide imaging methods. CMR is considered the most accurate noninvasive technique to evaluate the structure and right ventricular ejection fraction of the right ventricle (Video 270e-2). Although first-pass imaging by radionuclide angiography can provide accurate and reproducible measurements of right ventricular volumes and ejection fraction, it is not commonly used. Assessment of the right ventricle by echocardiography has generally been qualitative, owing in part to the unusual geometry of the right ventricle. However, several quantitative methods are available for assessment of right ventricular function,

including fractional area change ($FAC = [\text{diastolic area} - \text{systolic area}] / \text{diastolic area}$), which has been shown to correlate with outcomes in heart failure and after myocardial infarction. Excursion of the tricuspid annulus (tricuspid annular plane systolic excursion) is another widely used method to assess right ventricular function, although it is mostly used in research settings.

Abnormalities of right ventricular size and function are generally secondary to either diseases that affect the right ventricle intrinsically or disease in which the right ventricle responds to abnormalities elsewhere in the heart or pulmonary vasculature. Intrinsic diseases that affect the RV include congenital abnormalities, including hypoplastic right ventricle and arrhythmogenic right ventricular dysplasia, and acquired diseases, such as right ventricular infarction and infiltrative diseases that affect the right ventricle. Long-standing pulmonary hypertension or pulmonary outflow tract obstruction leads to right ventricular hypertrophy and ultimately dilatation. Although right ventricular dilatation can occur due to both chronic and acute processes, chronic right ventricular dilatation is usually secondary to long-standing increases in pulmonary pressures and can thus be distinguished from the acute processes that cause right ventricular dilatation. One such acute process that can cause profound right ventricular dilatation and dysfunction is acute pulmonary embolism. In the setting of acute occlusion of a pulmonary artery or branch, an acute rise in pulmonary vascular resistance causes a previously normal right ventricle to dilate and fail due to the increased afterload. In acute pulmonary embolism, right ventricular dilatation and dysfunction are signs of substantial hemodynamic compromise and are associated with a marked increase in likelihood of death. In addition to right ventricular dilatation, acute pulmonary embolism is often associated with a specific pattern of regional right ventricular dysfunction, commonly referred to as the McConnell sign, characterized by preservation of right ventricular wall motion in the basal and apical regions and dyskinesia in the region of the mid right ventricular free wall. This abnormality is highly specific for acute pulmonary embolism and is likely secondary to acute increases in right ventricular load.

Any disease that causes increased pulmonary vascular resistance can lead to right ventricular dilatation and dysfunction. Long-standing chronic obstructive pulmonary disease results in cor pulmonale in which right ventricular pressures become elevated as the right ventricle hypertrophies in response to the increased pulmonary vascular resistance. Acute pneumonia can cause findings that are similar to acute pulmonary embolism. In patients with right ventricular dilatation without obvious pulmonary disease, intracardiac shunts should be considered. The increased flow through the pulmonary vasculature as a result of an atrial septal or ventricular septal defect can, over time, result in elevation in pulmonary resistance with subsequent dilatation and hypertrophy of the right ventricle. Right ventricular dilatation and dysfunction also have prognostic significance in left-sided heart disease and have been shown to be important predictors of outcome in patients with heart failure or acute myocardial infarction.

In addition to assessment of left ventricular structure, assessment of the other cardiac chambers also provides important clues to intracardiac and systemic diseases. Enlargement of the left atrium is common in patients with hypertension and is also suggestive of increased left ventricular filling pressures; indeed, left atrial size is often termed the “hemoglobin A_{1c} ” of diastolic function, because left atrial enlargement reflects long-standing increase in left-sided filling pressures. Right atrial dilatation and dilatation of the inferior vena cava are common in conditions in which central venous pressure is elevated.

PATIENT SAFETY CONSIDERATIONS

RADIATION EXPOSURE

Both cardiac CT and radionuclide imaging expose patients to ionizing radiation. Several recent publications have raised concern regarding the potential harmful effects of ionizing radiation associated with cardiac imaging. The *effective dose* is a measure used to estimate the biologic effects of radiation and is expressed in millisieverts (mSv). However, measuring the radiation effective dose associated with diagnostic imaging is complex and imprecise and often results in varying estimates,

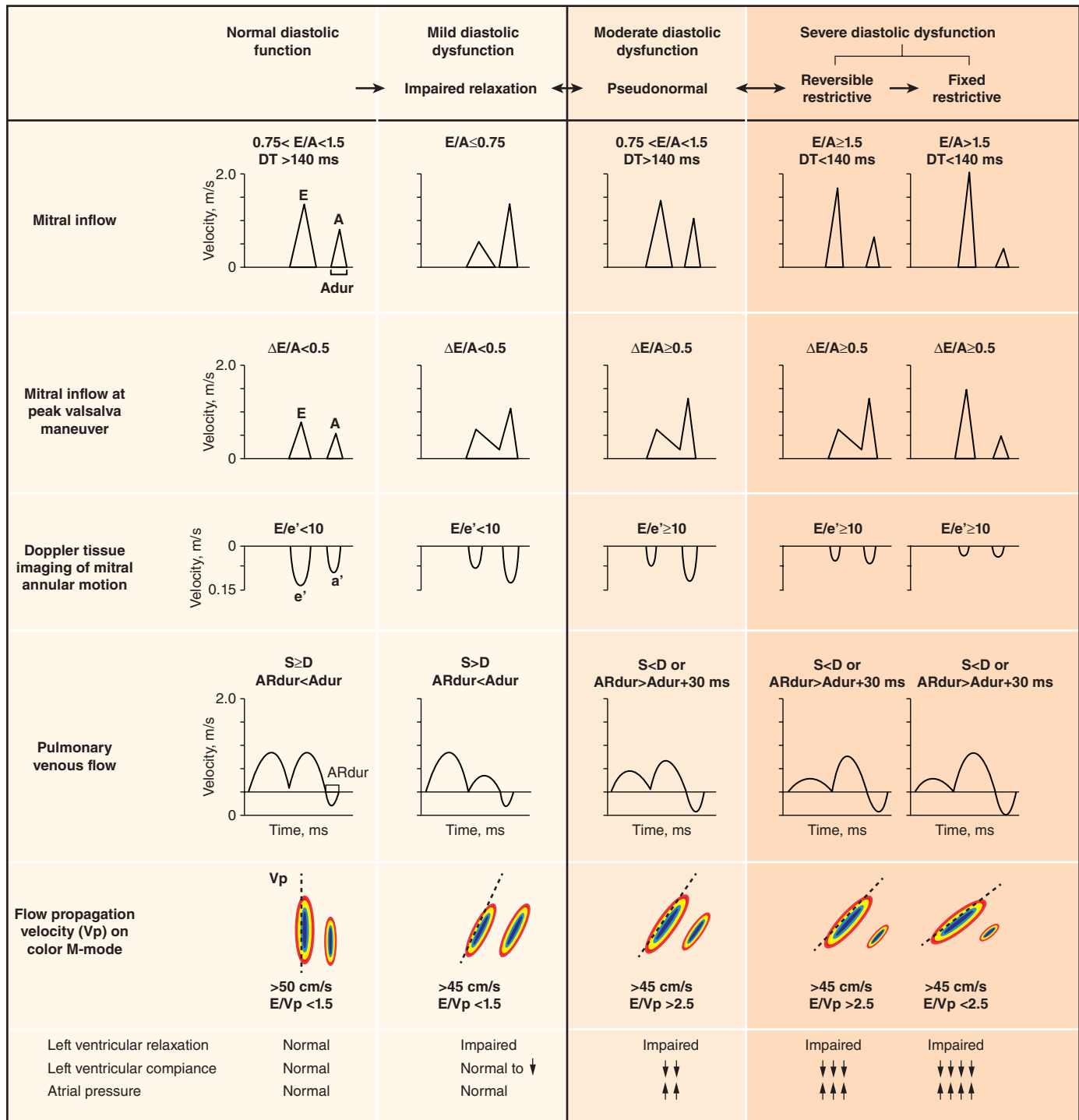


FIGURE 270e-8 Stages of diastolic function based on various parameters, including mitral inflow (with and without Valsalva maneuver), Doppler tissue imaging, pulmonary venous flow, and flow propagation. (Adapted with permission from MM Redfield et al: JAMA 289:194, 2003.)

even among experts. The effective dose from a typical myocardial perfusion SPECT scan ranges between ~4 and 11 mSv, depending on the protocol and type of scanner used. The effective dose from a typical myocardial perfusion PET scan is lower, ~2.5–4 mSv. Radiation exposure associated with cardiac CT is variable and, as with radionuclide imaging, also depends on the imaging protocol and scanner used. Although historic radiation doses with cardiac CT have been quite high, the introduction of newer technologies described above (e.g., x-ray tube modulation, prospective ECG gating) has resulted in a significant dose reduction. The current average radiation dose for a coronary CTA ranges from 5–15 mSv and, in selected cases, can be as low as 1 mSv. Imaging laboratories follow the ALARA (as low as reasonably achievable) principle when balancing the clinical need

and imaging approach. By comparison, the average dose for invasive coronary angiography is ~7 mSv, whereas exposure to radiation from natural sources in the United States amounts to ~3 mSv annually.

The risk of a fatal malignancy from medical imaging–related radiation is difficult to estimate precisely but is likely small and difficult to discern from the background risk of natural malignancies. The small but potential radiation risks from imaging mandate an assessment of the risk-versus-benefit ratio in the individual patient. In this context, one must not fail to take into account the risks of missing important diagnostic information by not performing a test (which could potentially influence near-term management and outcomes) for a theoretical concern of a small long-term risk of malignancy. Before ordering any test, especially one associated with ionizing radiation, we must

ensure the appropriateness of the study and that the potential benefits outweigh the risks. The likelihood that the study being considered will affect clinical management of the patient should be addressed before testing is performed. It is also important that “routine” follow-up scans in asymptomatic individuals be avoided.

CONTRAST AGENTS

Contrast agents are commonly used in cardiac CT, CMR, and echocardiography. Although their use significantly enhances the diagnostic information of each of these tests, there are also potential risks from the administration of contrast agents that should be considered.

The risk of adverse reactions from iodinated contrast agents used in cardiac CT is well established. The precise pathogenesis of contrast reactions following intravascular administration of iodinated contrast media is not known. The overall incidence of contrast reactions is 0.4–3% with nonionic formulations and higher for ionic formulations. Most contrast adverse reactions are mild and self-limiting. The risk of contrast-induced nephropathy (CIN) in patients with relatively normal renal function (glomerular filtration rate [GFR] >60 mL/min) is low. In most patients, CIN is self-limited, and renal function usually returns to baseline within 7–10 days, without progressing to chronic renal failure. However, this risk increases in patients with GFR <60 mL/min, especially older diabetic subjects. In such patients, appropriate screening and pre- and postscan hydration are necessary.

The use of gadolinium-based contrast agents (GBCAs) in CMR imaging enhances the versatility of this technique. Although there are several commercially available GBCAs in the United States, their use in cardiac imaging is considered off-label. Mild reactions from GBCAs occur in ~1% of patients, but severe or anaphylactic reactions are very rare. All GBCAs are chelated to make the compounds nontoxic and to allow renal excretion. Exposure to the nonchelated component of GBCA (Gd³⁺) has been associated with a rare condition known as nephrogenic systemic fibrosis (NSF), which is an interstitial inflammatory reaction that leads to severe skin induration, contracture of the extremities, fibrosis of internal organs, and even death. Risk factors to developing NSF include high-dose (>0.1 mmol/kg) GBCA use in presence of severe renal dysfunction (estimated GFR [eGFR] <30 mL/min per 1.73 m²), need for hemodialysis, an eGFR <15 mL/min per 1.73 m², use of gadodiamine contrast agent, acute renal failure, acute systemic illness, and presence of concurrent pro-inflammatory events. With the use of weight-based dosing and pretest screening, recent data suggest that NSF is extremely rare. Previously, an incidence of 0.02% in 83,121 patients exposed to GBCA over 10 years was noted; however, with current eGFR screening guidelines that have been widely practiced since 2006, a near-zero incidence of NSF has been reported.

Contrast agents can also be used in echocardiography. Injected agitated saline is used routinely to assess cardiac shunts, because these “bubbles” are too large to traverse the pulmonary circulation. After saline injection, the presence of bubbles in the left side of the heart is indicative of shunt, although the location can sometimes be difficult to determine. Dedicated echocardiographic contrast agents have been developed for opacification of left-sided structures and perfusion, although these are only currently U.S. Food and Drug Administration (FDA) approved for

perfusion. These agents are either albumin- or lipid-based microspheres filled with inert gases, typically perfluorocarbons. They are considered extremely safe, although they have, in extremely rare instances, been associated with allergic reactions and neurologic events.

SAFETY CONSIDERATIONS OF CMR IN PATIENTS WITH PACEMAKERS AND DEFIBRILLATORS

The risks of performing CMR in the presence of a pacemaker include generation of electrical current from the metallic hardware (especially if wire loops exist), device movement induced by the magnetic field, inappropriate pacing and sensing, and heating as a result of the “antenna’s effect.” While the presence of a permanent pacemaker remains a contraindication to CMR, highly experienced centers had reported success in performing MRI in these patients in a carefully monitored clinical setting. In general, patients need to be not pacemaker-dependent, the setting of the pacemaker needs to be modified to asynchronous mode, and the pulse sequence needs to be modified to reduce the amount of radiofrequency energy deposition. Pacemakers implanted for less than 6 weeks and the presence of epicardial, abandoned, or nonfixation leads are considered unsafe. Collectively, evidence from combined reports of >250 patients with pacemaker models manufactured after year 2000 suggests that CMR at 1.5 T or less can be performed without significant risk for the patient and with minor nonpermanent alteration of pacemaker settings and function. Similar safety data exists for automatic implantable cardioverter-defibrillators (AICDs), but they are based only on small numbers of patients. In 2011, the first CMR-compatible FDA-approved permanent pacemaker became available commercially. Currently, no AICD has achieved FDA clearance for MRI compatibility.

PATIENT-CENTERED APPLICATIONS OF CARDIAC IMAGING

CORONARY ARTERY DISEASE

The basis for the diagnostic application of imaging tests in patients with known or suspected CAD should be viewed in light of the pretest probability of disease as well as the specific characteristics of imaging tests (i.e., sensitivity and specificity). In symptomatic patients, the prevalence or pretest probability of CAD differs based on the type of symptom (typical angina, atypical angina, noncardiac chest pain), as well as on age, gender, and coronary risk factors. In an individual patient, the results of the initial test informs the posttest likelihood of CAD. In patients undergoing sequential testing (e.g., ECG treadmill testing followed by stress imaging), the posttest probability of disease after the first test becomes the pretest likelihood of disease for the second test. Regardless of the sequence, the expectation is that a test will provide sufficient information to confirm or exclude the diagnosis of CAD and that such information will allow accurate risk stratification to be able to guide management decisions.

Table 270e-3 summarizes the relative diagnostic accuracies of cardiac imaging modalities for the diagnosis of CAD.

It is important to highlight that the vast majority of studies included in meta-analyses of the diagnostic accuracy of cardiac imaging modalities for the diagnosis of CAD were retrospective, small, single-center studies, comprising predominantly male patients with a high prevalence

TABLE 270e-3 COMPARATIVE DIAGNOSTIC ACCURACY OF CARDIAC IMAGING APPROACHES TO CORONARY ARTERY DISEASE

Imaging Modality	Published Data	Sensitivity	Specificity
Exercise echocardiography	15 studies (n = 1849 patients)	84%	82%
Dobutamine echocardiography	28 studies (n = 2246 patients)	80%	84%
SPECT MPI	113 studies (n = 11,212 patients)	88%	76%
Myocardial perfusion PET	9 studies (n = 650 patients)	93%	81%
CMR perfusion	37 studies (n = 2841 patients)	91%	81%
CMR wall motion	14 studies (n = 754 patients)	83%	86%
Coronary CTA	18 studies (n = 1286 patients)	99%	89%

Note: In these studies, the diagnosis of coronary artery disease was based on the presence of a >50% or >70% stenosis on invasive coronary angiography.

Abbreviations: CMR, cardiac magnetic resonance; CTA, computed tomography angiography; MPI, myocardial perfusion imaging; PET, positron emission tomography; SPECT, single-photon emission computed tomography.

270e-10 of CAD (>50–60%). Multicenter studies assessing the performance of individual modalities or comparing different modalities have consistently resulted in more modest diagnostic accuracies, tracking more closely with how these tests perform in practice.

Stress Echocardiography The hallmark of myocardial ischemia during stress echocardiography is the development of new regional wall motion abnormalities and reduced systolic wall thickening (Video 270e-3). Stress echocardiography can be performed in conjunction with exercise or dobutamine stress. Stress echocardiography is best at identifying inducible wall motion abnormalities in previously normally contracting segments. In a patient with wall motion abnormalities at rest, the specificity of stress echocardiography is reduced, and worsening regional function of a previously abnormal segment might reflect worsening contractile function in the setting of increased wall stress rather than new evidence of inducible ischemia.

The advantages of stress echocardiography over other stress imaging techniques include its relatively good diagnostic accuracy, widespread availability, no use of ionizing radiation, and relatively low cost. Limitations of stress echocardiography include (1) the technical challenges associated with image acquisition at peak exercise because of exertional hyperpnea and cardiac excursion, (2) the fact that rapid recovery of wall motion abnormalities can be seen with mild ischemia (especially with one-vessel disease, which limits sensitivity), (3) difficulty detecting residual ischemia within an infarcted territory because of resting wall motion abnormality, (4) high operator dependence for acquisition of echocardiographic data and analysis of images, and (5) the fact that good-quality complete images viewing all myocardial segments occurs in only 85% of patients. Newer techniques including second harmonic imaging and the use of intravenous contrast agents improve image quality, but their effect on diagnostic accuracy has not been well documented. The use of IV contrast agents may also allow assessment of myocardial perfusion, although this is not approved or generally reimbursed, and data concerning the utility of contrast perfusion echocardiography are limited.

As with nuclear perfusion imaging, stress echocardiography is often used for risk stratification in patients with suspected or known CAD. A negative stress echocardiogram is associated with an excellent prognosis, allowing identification of patients at low risk. Conversely, the risk of adverse events increases with the extent and severity of wall motion abnormalities on stress echocardiography.

Stress Radionuclide Imaging SPECT myocardial perfusion imaging is the most common form of stress imaging tests for CAD evaluation. The presence of a reversible myocardial perfusion defect is indicative of ischemia (Fig. 270e-9, left panel), whereas a fixed perfusion defect generally reflects prior myocardial infarction (Fig. 270e-9, right panel). As discussed above, PET has advantages compared to SPECT, but it is not widely available and is more expensive and, thus, considered an emerging technology in clinical practice.

Nuclear perfusion imaging is another robust approach to diagnose obstructive CAD, quantify the magnitude of inducible myocardial ischemia, assess the extent of tissue viability, and guide therapeutic management (i.e., selection of patients for revascularization). One of the most valuable clinical applications of radionuclide perfusion imaging is for risk stratification. It is well established that patients with a normal SPECT or PET study exhibit a median rate of major adverse cardiac events of <1% annually. Importantly, the risks of death and myocardial infarction increase linearly with increasing magnitude of perfusion abnormalities, reflecting the extent and severity of CAD.

Despite the widespread use and clinical acceptance of radionuclide imaging in CAD evaluation, a recognized limitation of this approach is that it often uncovers only coronary territories supplied by the most severe stenoses. Consequently, it is relatively insensitive to accurately

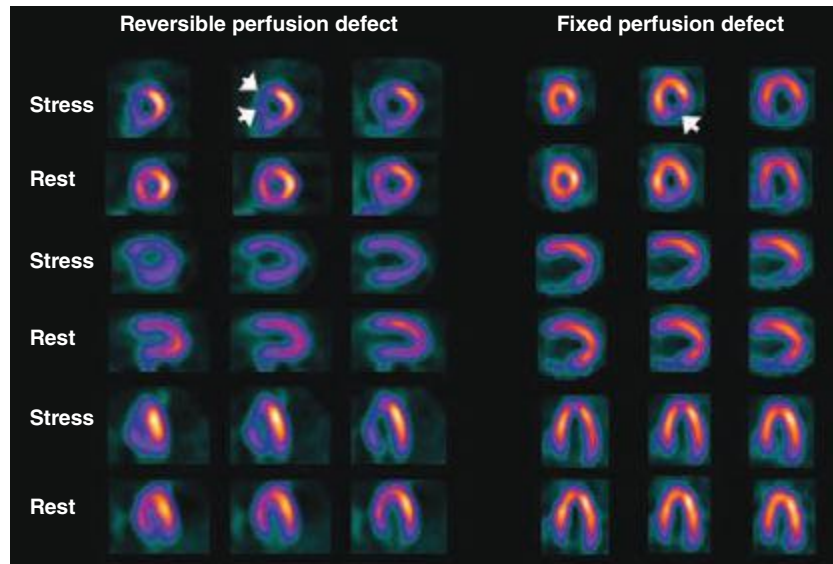


FIGURE 270e-9 Selected technetium-99m sestamibi myocardial perfusion single-photon emission computed tomography images of two different patients demonstrating a reversible perfusion defect involving the anterior and septal left ventricular wall, reflecting ischemia in the left anterior descending coronary territory (arrows in left panel), and a fixed perfusion defect involving the inferior and inferolateral walls consistent with myocardial scar in the right coronary territory (arrow in right panel).

delineate the extent of obstructive angiographic CAD, especially in the setting of multivessel disease. The use of quantitative myocardial blood flow and coronary flow reserve with PET can help mitigate this limitation. In patients with so-called “balanced” ischemia or diffuse CAD, measurements of coronary flow reserve uncover areas of myocardium at risk that would generally be missed by performing only relative assessments of myocardial perfusion (Fig. 270e-10). Conversely, a normal coronary flow reserve is associated with a very high negative predictive value for excluding high-risk angiographic CAD. These measurements of coronary flow reserve also contribute to risk stratification across the spectrum of ischemic changes, including patients with visually normal myocardial perfusion.

HYBRID CT AND NUCLEAR PERFUSION IMAGING Because many of the newer generation nuclear medicine scanners integrate CT and a gamma camera in the same acquisition gantry, it is now possible to acquire and quantify myocardial scar and ischemia and CAC scoring from a single dual-modality study (SPECT/CT or PET/CT) (Fig. 270e-11). The rationale for this integrated approach is predicated on the fact that the perfusion imaging approach is designed to uncover only obstructive atherosclerosis. Conversely, CAC scoring (or CT coronary angiography) provides a quantitative measure of the anatomic extent of atherosclerosis. This provides an opportunity to improve the conventional models for risk assessment using nuclear imaging alone, especially in patients without known CAD.

Cardiac CT Voluminous plaques are more prone to calcification, and stenotic lesions frequently contain large amounts of calcium. Indeed, there is evidence that high CAC scores are generally predictive of a higher likelihood of obstructive CAD, and the available data support the concept of a threshold phenomenon governing this relationship (i.e., Agatston score >400). However, given the fact that CAC scores are not specific markers of obstructive CAD, one should be cautious in using this information as the basis for referral of patients to coronary angiography, especially in symptomatic patients with low-risk stress tests. Conversely, CAC scores <400, especially in symptomatic patients with intermediate-high likelihood of CAD, as in those with typical angina, may be less effective in excluding CAD, especially in young symptomatic men and women who may have primarily noncalcified atherosclerosis (Fig. 270e-12).

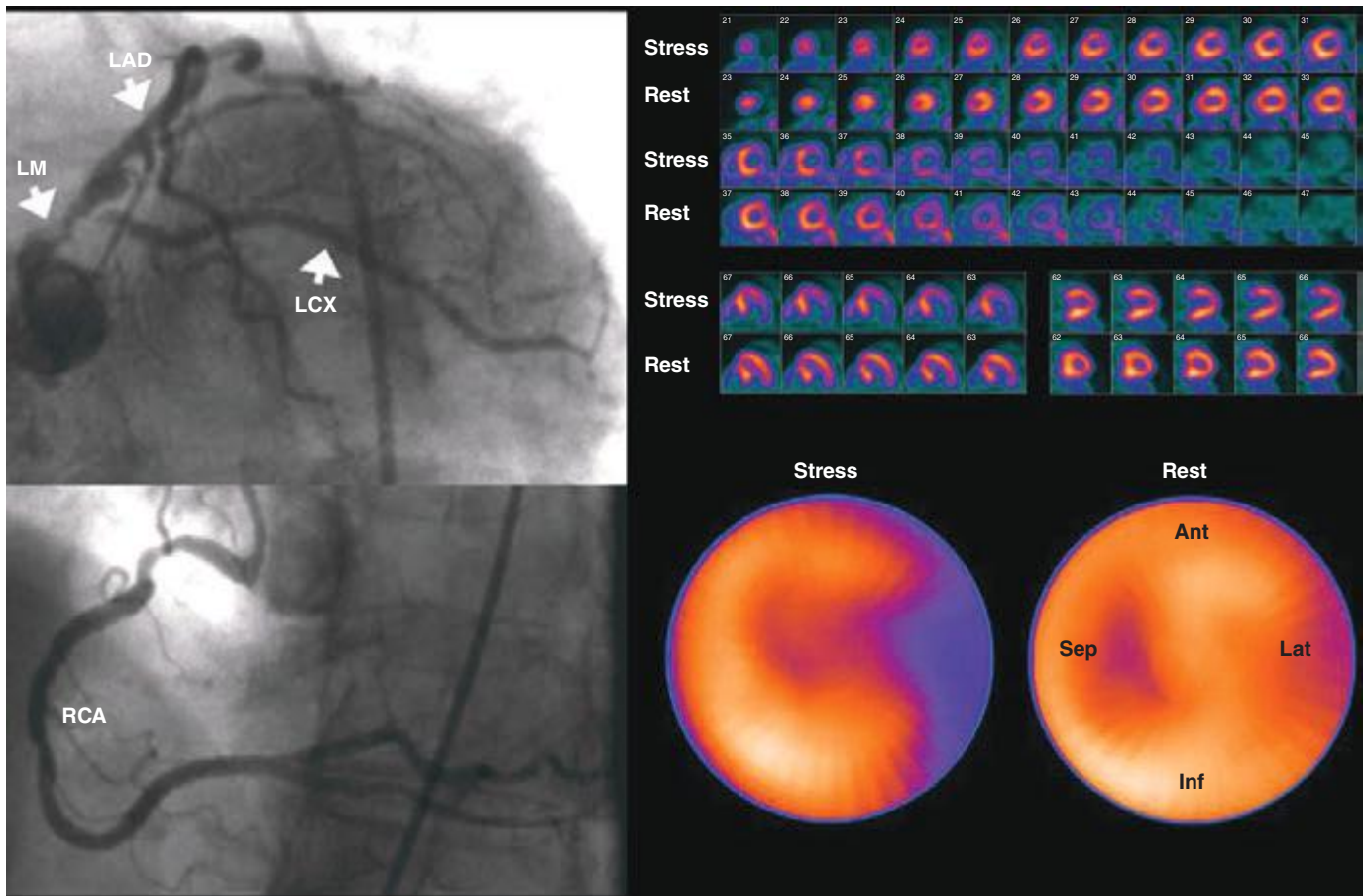


FIGURE 270e-10 Coronary angiographic (*left panel*) and rubidium-82 myocardial perfusion positron emission tomograph images (*right panel*) in an 85-year-old female with diabetes presenting with chest pain. The coronary angiogram demonstrates significant stenoses of the left main and circumflex coronary arteries. However, the perfusion images demonstrate only a reversible lateral wall defect. Quantification of stress and rest myocardial blood flow demonstrated a significant, global reduction on coronary flow reserve (estimated at 1.2, normal value >2.0), reflecting extensive myocardium risk that was underestimated by the semiquantitative estimates of myocardial perfusion. LAD, left anterior descending artery; LCX, left circumflex artery; LM, left main artery; RCA, right coronary artery.

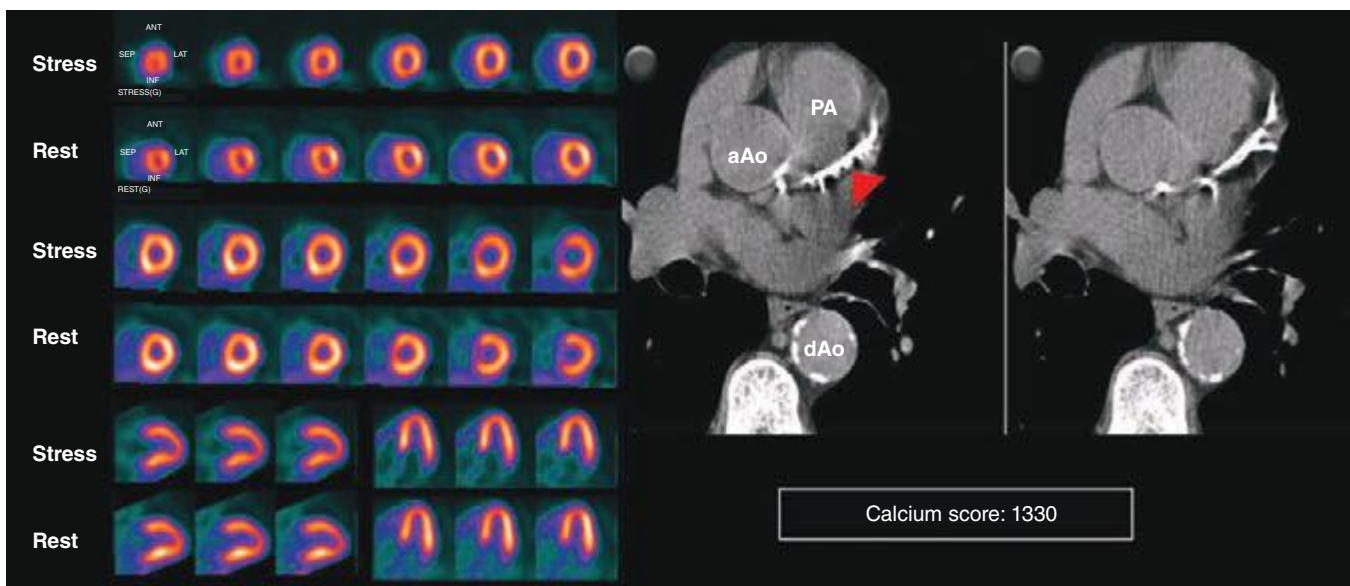


FIGURE 270e-11 Stress and rest rubidium-82 myocardial perfusion positron emission tomography (PET) images (*left*) and noncontrast gated computed tomography (CT) images (*right*) delineating the extent and severity of coronary artery calcifications obtained with integrated PET/CT imaging. The images demonstrate extensive atherosclerosis (Agatston coronary calcium score = 1330) without flow-limiting disease based on the normal perfusion study. aAo, ascending aorta; dAo, descending aorta; PA, pulmonary artery.

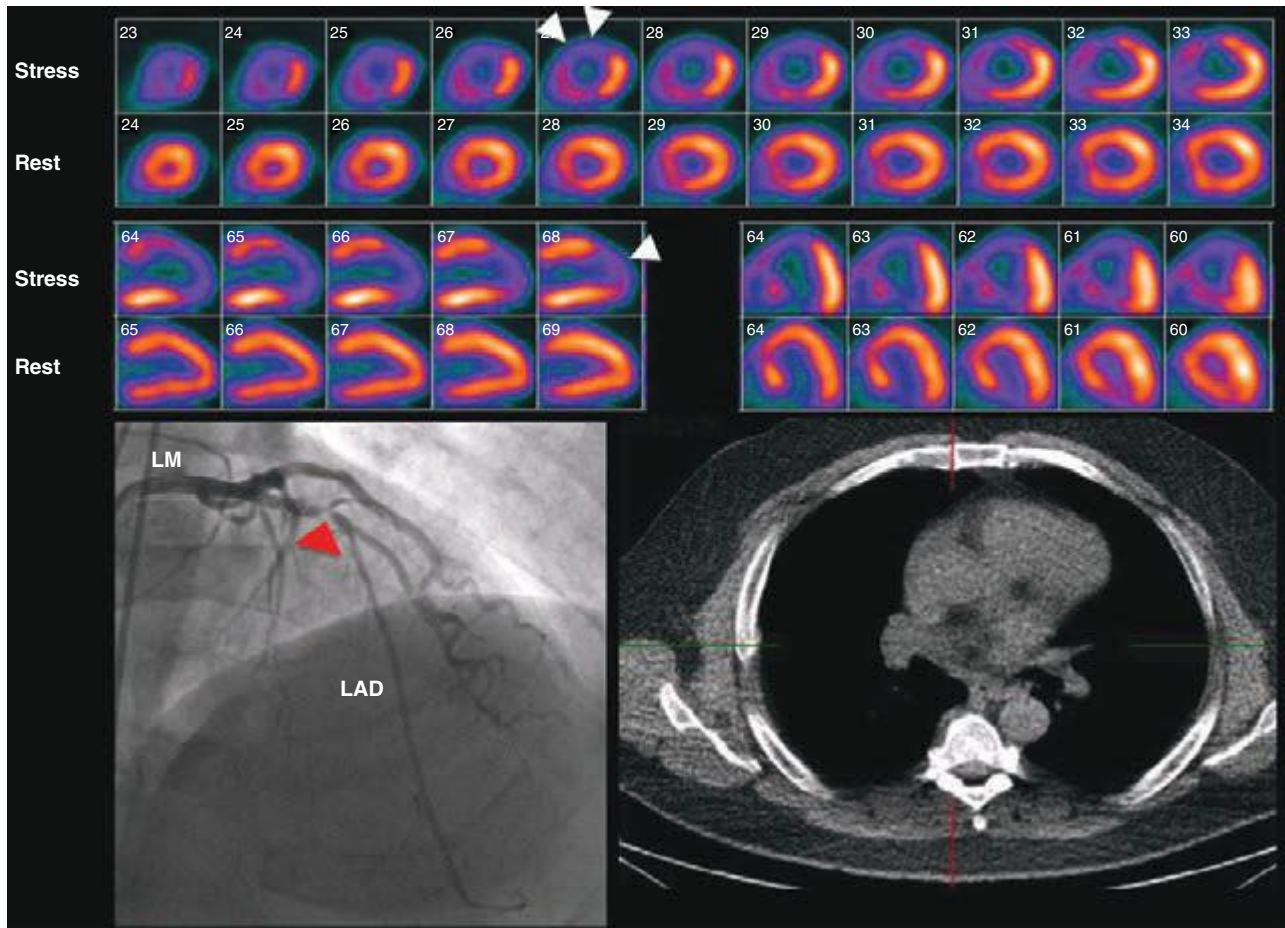


FIGURE 270e-12 Stress and rest rubidium-82 myocardial perfusion positron emission tomography images (*top*), noncontrast gated computed tomography images (*lower right*), and selected coronary angiographic images obtained on a 59-year-old male patient with atypical angina. Despite the absence of significant coronary calcifications (Agatston calcium score = 0), the perfusion images demonstrated a dense and reversible perfusion defect involving the anterior and anteroseptal walls (*arrows*), reflecting significant obstructive disease in the left anterior descending coronary artery (LAD), confirmed on angiography. LM, left main artery.

As discussed above, the improved temporal and spatial resolution of modern multidetector CT scanners offer a unique noninvasive approach to delineate the extent and severity of coronary atherosclerosis. The extremely high sensitivity of this approach offers a very effective means for excluding the presence of CAD (high negative predictive value) (Table 270e-3). In the setting of high coronary calcium scores (e.g., >400), however, specificity is reduced because the blooming artifact of calcium does not allow one to evaluate the vessel lumen accurately. Given the high negative predictive value of CTA, a normal scan result effectively excludes obstructive CAD and abolishes the need for further investigation. As discussed below, this may be quite useful in patients with low-intermediate clinical risk presenting to the emergency room for chest pain. However, the limited capability of this technique to determine the severity of stenosis and to predict which obstructions are flow limiting can make abnormal scan results more difficult to interpret, especially in terms of the possible need of coronary revascularization. There are emerging data suggesting that by adding a stress myocardial perfusion CT evaluation (similar to stress perfusion CMR) (Fig. 270e-13, *top panel*) or an estimated fractional flow reserve (so-called FFR_{CT}) (Fig. 270e-13, *lower panel*), one can define the hemodynamic significance of anatomic stenosis. However, these are not in routine clinical use and remain emerging technologies.

As with invasive coronary angiography, assessments of the extent of CAD by CTA can also provide useful prognostic information. A low 1-year cardiac event rate has been reported for patients without obstructive CAD on CTA. For patients with obstructive CAD, the risk of adverse cardiac events increases proportionally with the extent of angiographically obstructive CAD.

Although CTA can be helpful in assessing patency of bypass grafts, the assessment of stents is somewhat more challenging because the limited spatial resolution of CT and stent diameter (<3 mm being associated with the highest number of partial lumen visualization and nondiagnostic scans) both contribute to limited clinical results.

CMR Imaging The two approaches used with CMR to evaluate known or suspected CAD include the assessment of regional myocardial perfusion or wall motion at rest and during stress, the latter being analogous to dobutamine echocardiography. Although treadmill or bicycle exercise stress CMR is practiced in a small number of specialized centers, the logistics for stress MRI studies currently require the use of pharmacologic stress agents including vasodilators or dobutamine. Myocardial perfusion is evaluated by injecting a bolus of a GBCA followed by continuous data acquisition as the contrast passes through the cardiac chambers and into the myocardium. Relative perfusion deficits are recognized as regions of low signal intensity (black) within the myocardium (Video 270e-4). In addition, LGE imaging allows detection of bright areas of myocardial scar (white), which further enhances the utility of this approach for diagnosis of CAD (Fig. 270e-14).

The major advantage of dobutamine CMR over dobutamine echocardiography is better image quality and sharper definition of endocardial borders from the blood pool. Consequently, dobutamine CMR appears to have better diagnostic accuracy than dobutamine echocardiography for detection of CAD, especially in patients with poor acoustic window (Table 270e-3). A limitation of high-dose dobutamine stress CMR is that it bears the potential risk of severe side effects, such as hypotension and severe ventricular arrhythmias in the inhospitable

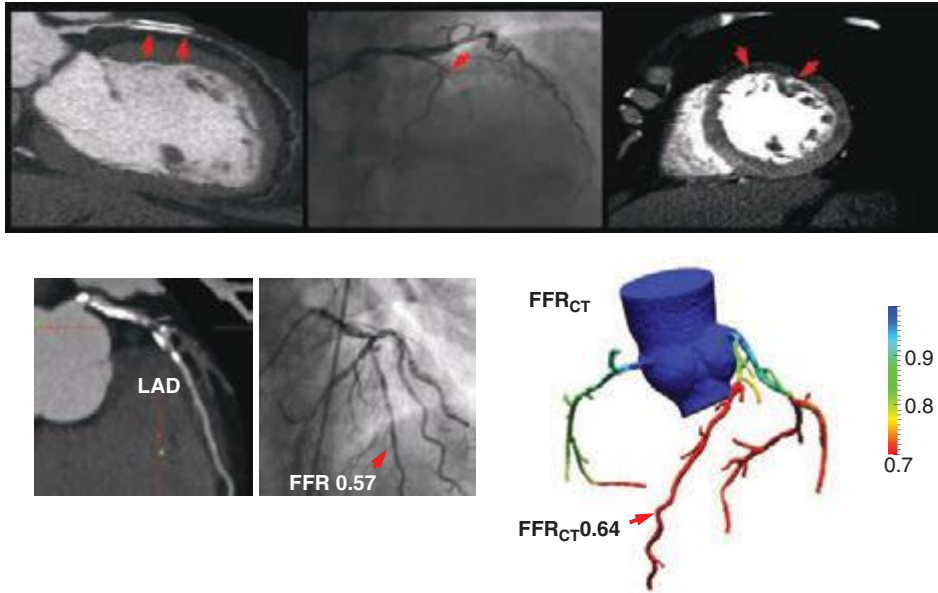


FIGURE 270e-13 Examples of novel approaches to the assessment of flow-limiting coronary artery disease (CAD) with cardiac computed tomography (CT). In the *top panel*, representative views of a coronary CT angiogram (CTA; *left*), coronary angiogram (*middle*), and stress myocardial perfusion CT (*right*) images in a patient with CAD and prior stenting of the left anterior descending coronary artery (LAD) are presented. On the CTA, the stent (*arrows*) is totally occluded as evidenced by the loss of contrast enhancement distal to the stent. The coronary angiogram demonstrates a concordant total occlusion of the LAD. On the perfusion CT images, there is a black rim (*arrows*) involving the anterior and anterolateral walls, indicating the lack of contrast opacification during stress consistent with myocardial ischemia. (*Images courtesy of CORE 320 investigators.*) The *lower panel* illustrates an example of fractional flow reserve (FFR) estimates with coronary CTA (*left*) compared to the reference standard of invasive FFR. The FFR reflects the pressure differential between a coronary segment distal to a stenosis and the aorta. In normal coronary arteries, there is no gradient, and FFR is 1. An FFR <0.80 is consistent with a hemodynamically significant stenosis. (*Images courtesy of Dr. James Min, Cornell University, New York.*)

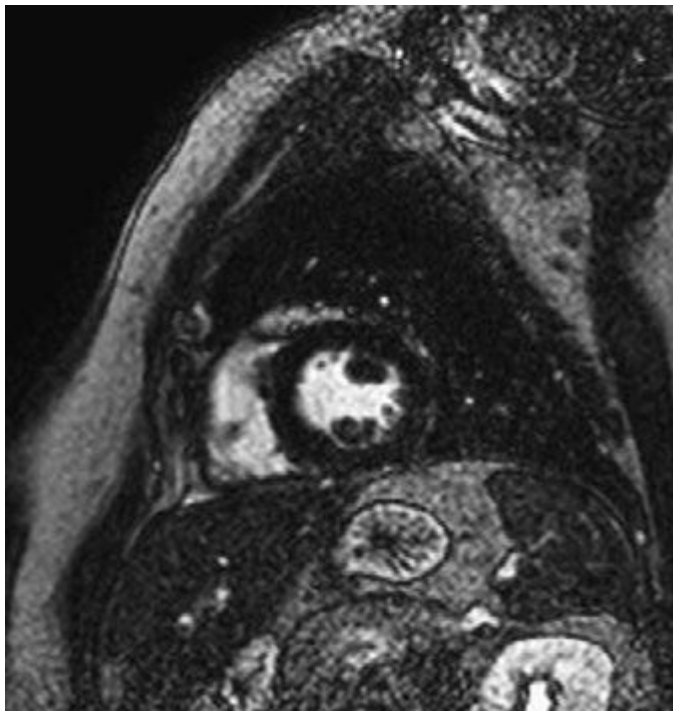


FIGURE 270e-14 The image shows the late gadolinium enhancement image of a mid short-axis view. There is no evidence of infarction in the anterior wall, which would be seen as bright white areas, indicating that the stress perfusion defect primarily represents myocardial ischemia. This patient had a significant stenosis of the left anterior descending coronary artery.

environment of the magnetic resonance scanner. It occurs rarely (~5%), and most cases can be prevented with proper monitoring of vital signs and regional cine function. The advantage of stress perfusion CMR over SPECT is its clearly higher spatial resolution, allowing detection of subendocardial defects that may be missed by SPECT. The addition of the information from LGE imaging allows differentiation of hypoperfused (potentially ischemic) from infarcted myocardium and characterizes the extent of myocardial ischemia.

As with other imaging modalities, there is evidence that ischemia measurements derived from stress CMR studies also have prognostic value. In line with the nuclear and echocardiography literature, a normal CMR study is associated with a good prognosis. Conversely, the presence of new wall motion abnormalities, regional perfusion defects, the combination of wall motion abnormalities and perfusion defects, and the presence of LGE are all predictors of adverse events.

Selecting a Testing Strategy in Patients without Known CAD As discussed above, there are many options for the evaluation of a patient with suspected CAD presenting with chest pain symptoms. The critical questions to be answered by a testing strategy include the following: (1) Does the chest pain reflect obstructive CAD? (2) What are the short- and long-term risks? (3) Does the patient need to be considered for revascularization?

For symptomatic patients without a prior history of CAD and a normal or nearly normal resting ECG who are able to exercise, the American College of Cardiology/American Heart Association guidelines recommend standard exercise treadmill testing (ETT) as the initial testing strategy. The guidelines further suggest that patients who are categorized as low risk by ETT (e.g., those achieving >10 metabolic equivalents [METs] without chest pain or ECG changes) be treated initially with medical therapy, and those with high-risk ETT findings (i.e., typical angina with >2 mm ST-segment depression in multiple leads, ST elevation during exercise, drop in blood pressure, or sustained ventricular arrhythmias) be referred for coronary angiography.

The use of exercise testing in women presents difficulties that are not seen in men, reflecting the differences in the lower prevalence of obstructive CAD in women and the different accuracy of exercise testing in men and women. Compared with men, the lower pretest probability of disease in women means that more test results are false positive. In some of these patients, a positive ETT may reflect true myocardial ischemia caused by microvascular coronary artery dysfunction (so-called *microvascular disease*). In addition, the inability of many women to exercise to maximum aerobic capacity, the greater prevalence of mitral valve prolapse and microvascular disease in women, and possibly other reasons may contribute to the differences with men as well. The difficulties of using exercise testing for diagnosing obstructive CAD in women have led to speculation that stress imaging may be preferred over standard stress testing. However, recent data from the WOMEN study suggests that in symptomatic, low-risk women who are able to exercise, standard ETT is a very effective initial diagnostic strategy as compared to stress radionuclide imaging. Women included in the study were randomized to standard ETT or exercise radionuclide perfusion imaging. The primary endpoint was the 2-year incidence of major adverse cardiac

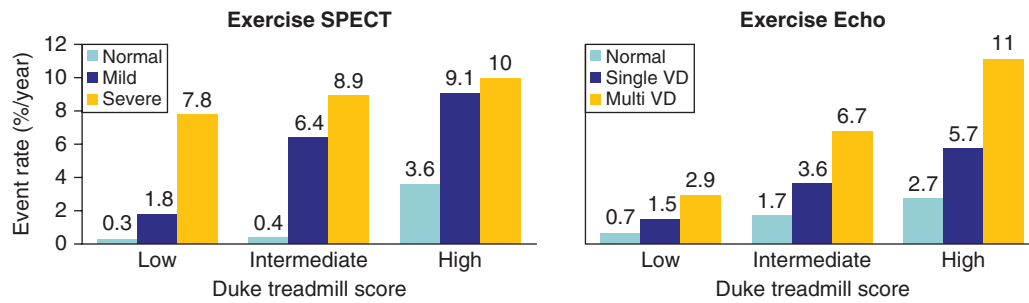


FIGURE 270e-15 Incremental risk stratification of stress imaging over Duke treadmill score in patients with suspected coronary artery disease. Stress imaging is most valuable in the intermediate-risk group. SPECT, single-photon emission computed tomography; VD, vessel disease. (Reproduced with permission from R Hachamovitch et al: *Circulation* 93:905, 1996; and TH Marwick et al: *Circulation* 103:2566, 2001.)

events, defined as CAD death or hospitalization for an acute coronary syndrome or heart failure. At 2 years, there was no difference in major adverse cardiac events. As expected, ETT resulted in 48% lower costs compared to exercise radionuclide imaging.

Patients with intermediate-high risk after ETT (e.g., low exercise duration, chest pain, and/or ST-segment depression without high-risk features) will often require additional testing, either stress imaging or noninvasive CT coronary angiography, to more accurately characterize clinical risk. Most common stress imaging strategies in intermediate-risk patients include stress echocardiography and radionuclide imaging. In such patients, stress imaging with either SPECT or echocardiography has been shown to accurately reclassify patients who are initially classified as intermediate risk by ETT as low or high risk (Fig. 270e-15). Following this staged strategy of applying the low-cost ETT first and reserving more expensive imaging to refine risk stratification to patients initially classified as intermediate risk by ETT is more cost effective than applying stress or anatomic imaging as the initial test routinely.

A stress imaging strategy is the recommended first step for patients who are unable to exercise to an adequate workload and/or those with abnormal resting ECGs (e.g., left ventricular hypertrophy with strain, left bundle branch block). Importantly, the most recent documents regarding appropriate use of radionuclide and echocardiography imaging also considered that an imaging strategy may be an appropriate first step in patients with intermediate-high likelihood of CAD (e.g., diabetics, renal impairment) due to increased overall sensitivity for diagnosis of CAD and improved risk stratification.

In considering the potential clinical application of imaging modalities, the evidence supporting the role of assessment of ischemia versus anatomy must be considered. From the discussion above, a normal CTA is helpful because it effectively excludes the presence of obstructive CAD and the need for further testing, defines a low clinical risk, and makes management decisions regarding referral to coronary angiography straightforward. Because of its limited accuracy to define stenosis severity and predict ischemia, however, abnormal CTA results are more problematic to interpret and to use as the basis for defining the potential need of invasive coronary angiography and revascularization. In such patients, a follow-up stress test is usually required to determine the possible need of revascularization (Fig. 270e-16).

The justification of stress imaging in testing strategies has hinged on the identification of which patients may benefit from a revascularization strategy by means of noninvasive estimates of jeopardized myocardium rather than angiography-derived anatomic stenoses. Indeed, there is evidence that only the presence of moderate-severe ischemia identifies patients with apparent improved survival with revascularization. Patients with mild or no ischemia are better candidates for optimal medical therapy. The advantages of this approach include

avoidance of excess catheterizations with their associated cost and risk and the potential for intervening unnecessarily. The acceptable diagnostic accuracy of stress imaging approaches, along with their robust risk stratification, and the ability of ischemia information to identify patients who would benefit from revascularization suggest a potential role as a first imaging strategy in patients with intermediate-high likelihood of CAD. Although the available data suggest similar diagnostic accuracy for SPECT, PET, echocardiography, and CMR, the choice of strategy depends on availability and local expertise.

Selecting a Testing Strategy in Patients with Known CAD Use and selection of testing strategies in symptomatic patients with established CAD (i.e., prior angiography, prior myocardial infarction, prior revascularization) differ from those in patients without prior CAD. Although standard

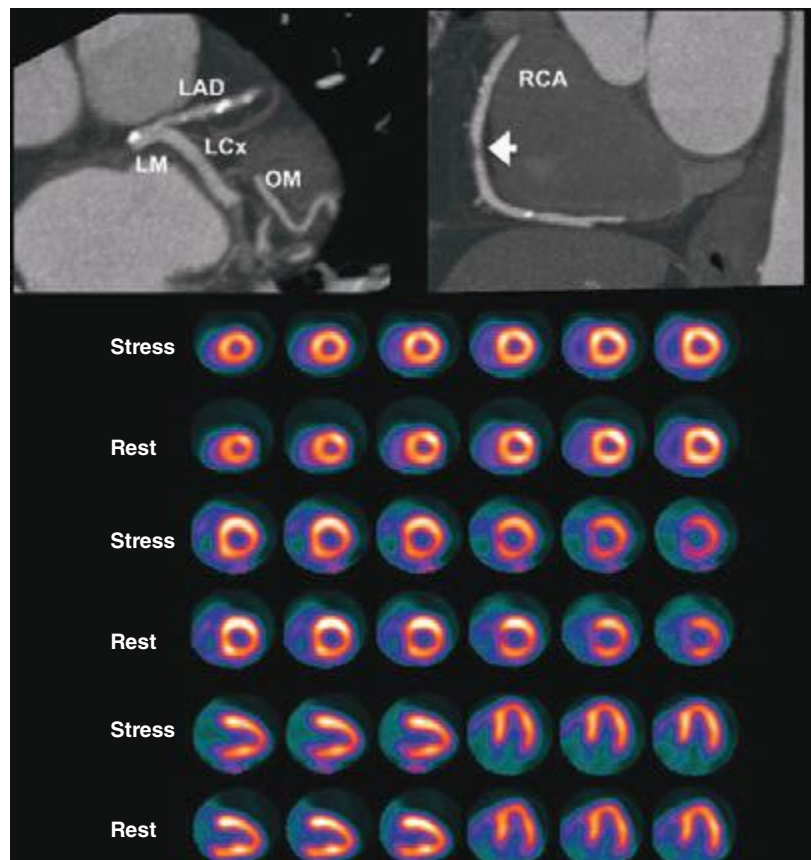


FIGURE 270e-16 Selected views from coronary computed tomography angiographic (CTA) images (top panel) and stress and rest rubidium-82 myocardial perfusion positron emission tomography images (lower panel) obtained on a 64-year-old male patient with atypical angina. The CTA images demonstrate dense focal calcifications in the left main (LM) and left anterior descending (LAD) coronary arteries and a significant noncalcified plaque in the mid right coronary artery (RCA; arrow). The myocardial perfusion images demonstrated no evidence of flow-limiting stenosis. LCx, left circumflex artery; OM, obtuse marginal branch.

ETT may help distinguish cardiac from noncardiac chest pain, exercise ECG has a number of limitations following myocardial infarction and revascularization (especially coronary artery bypass grafting). These patients frequently have rest ECG abnormalities. In addition, there is a clinical need to document both the magnitude and localization of ischemia to be able to direct therapy, especially the potential need for targeted revascularization. Consequently, imaging tests are preferred for evaluating patients with known CAD.

There are also important differences in the effectiveness of imaging tests in these patients. As discussed above, coronary CTA is limited in patients with prior revascularization. Patients with prior coronary artery bypass grafting are a particularly heterogeneous group with respect to the anatomic basis of ischemia and its implications for subsequent morbidity and mortality. In addition to graft attrition, progression of disease in the native coronary arteries is not uncommon in symptomatic patients. While CTA provides excellent visualization of the bypass grafts, the native circulation tends to get heavily calcified and is generally not a good target for imaging with CTA. Likewise, blooming artifacts from metallic stents also limit the application of coronary CTA in patients with prior percutaneous coronary intervention. Although newer stent material may change the potential role of CTA in the future, it is probably not the first line of testing in these patients. If an anatomic strategy is indicated, direct referral to invasive angiography is preferred.

Stress imaging approaches are especially useful and preferred in symptomatic patients with established CAD. As in patients without prior CAD, normal imaging studies in symptomatic patients with established CAD also identify a low-risk cohort. In those with abnormal stress imaging studies, the degree of abnormality relates to posttest risk. In addition, stress imaging approaches can localize and quantify the magnitude of ischemia (especially with perfusion imaging), thereby assisting in planning targeted revascularization procedures. As in patients without prior CAD, the choice of stress imaging strategy depends on availability and local expertise.

Testing Strategy Considerations in Patients Presenting with Chest Pain to the Emergency Department Although acute chest pain is a frequent reason for patient visits to the emergency department (ED), only a small minority of those presentations represent an acute coronary syndrome (ACS). Strategies used in the evaluation of these patients include novel cardiac biomarkers (e.g., serum troponins), conventional stress testing (ETT), and noninvasive cardiac imaging. It is generally accepted that the primary goal of this evaluation is exclusion of ACS and other serious conditions rather than detection of CAD.

The routine evaluation of acute chest pain in most centers in the United States includes admission to a chest pain unit to rule out ACS with the use of serial ECGs and cardiac biomarkers. In selected patients, stress testing with or without imaging may be used for further risk stratification. Stress echocardiography and radionuclide imaging are among the most frequently used imaging approaches in these patients. The relative strengths and weaknesses of these testing options have been discussed above. Both approaches have been shown to be effective for identifying low-risk patients who can be safely discharged from the ED. Multiparametric CMR imaging has also been used successfully in patients with acute chest pain. In addition to the combined assessment of regional and global left ventricular function, myocardial perfusion, and tissue viability, it is also possible to evaluate the presence of myocardial edema to characterize the myocardium at risk secondary to reduced coronary flow (Video 270e-5). Due to its ability to probe multiple aspects of myocardial physiology, cardiac anatomy, and tissue characterization with LGE imaging, CMR is also useful in diagnosing conditions that mimic ACS (e.g., acute myocarditis, takotsubo cardiomyopathy, pericarditis) (Fig. 270e-17). Thus, CMR imaging offers unique information of myocardial pathophysiology in the spectrum of ACS and is, perhaps, the most versatile of all noninvasive imaging techniques. Unfortunately, it is not widely available even at specialized centers and is not a first-line testing strategy. The main disadvantages of the “functional” testing strategy are that it is time consuming and is generally associated with a prolonged length of stay and, thus, is more costly.

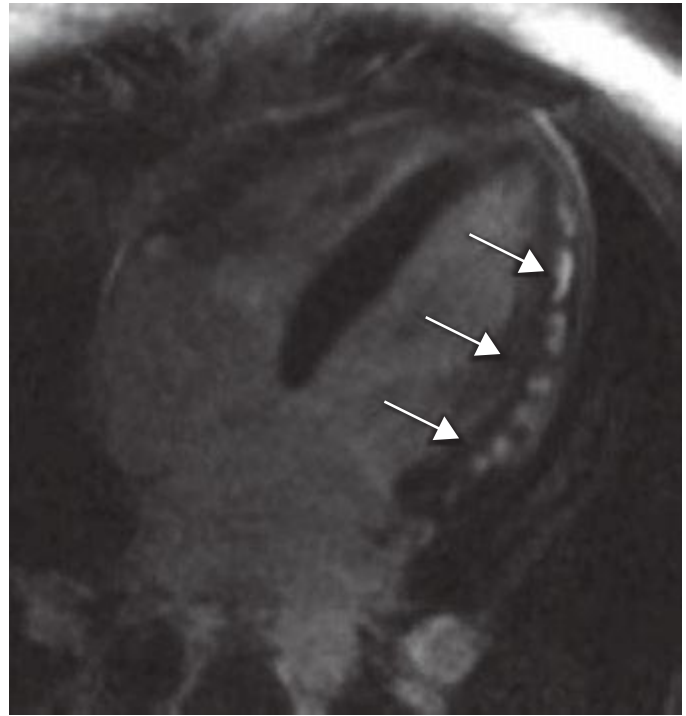


FIGURE 270e-17 A four-chamber long-axis late gadolinium enhancement (LGE) image of a patient with acute myocarditis.

Note that the LGE primarily involved the epicardial aspect of the myocardium (arrows), sparing the endocardium, which is a feature that distinguishes myocarditis from myocardial infarction, which affects the endocardium. Also note the multiple foci of LGE in this case affecting the lateral wall of the left ventricle. Viral myocarditis often presents with this pattern.

As discussed above, coronary CTA is a rapid and accurate imaging technique to exclude the presence of CAD and is well suited for the evaluation of patients with acute chest pain (Fig. 270e-18). Several single-center and, more recently, multicenter studies have demonstrated the feasibility, safety, and accuracy of coronary CTA in the ED. There have been four randomized controlled trials evaluating the efficacy of coronary CTA as the initial testing strategy as compared to usual care (which typically includes stress imaging). Patients in these trials had a very low clinical risk. Overall, there were no deaths and very few myocardial infarctions without differences between the groups. Likewise, there were no differences in postdischarge ED visits or rehospitalizations. These studies showed decreased length of stay with coronary CTA, and most but not all reported cost savings. An observation from a recent meta-analysis was that, compared to usual care, more patients assigned to coronary CTA underwent cardiac catheterization (6.3% vs 8.4%, respectively) and revascularization (2.6% vs 4.6%, respectively). The relative increased frequency in the referral to cardiac catheterization and revascularization after coronary CTA compared to stress imaging testing strategies has also been observed in patients with stable chest pain syndromes.

Taken together, the available data clearly suggest that not all patients presenting with acute chest pain require specialized imaging testing. Patients with very low clinical risk and negative biomarkers (especially high-sensitivity troponin assays) can be safely triaged. The use of imaging tests in patients with low-intermediate risk should be carefully considered, especially given the trade-offs discussed above.

VALVULAR HEART DISEASE

Abnormalities of any of the four valvular structures in the heart can lead to significant cardiac dysfunction, heart failure, or even death. Echocardiography, CMR, and cardiac CT can be used for the evaluation of valvular heart disease, although echocardiography has generally been considered the first imaging test of choice for the assessment of valvular heart disease. In addition, echocardiography is the most cost-effective screening method for valvular heart disease. In some cases,

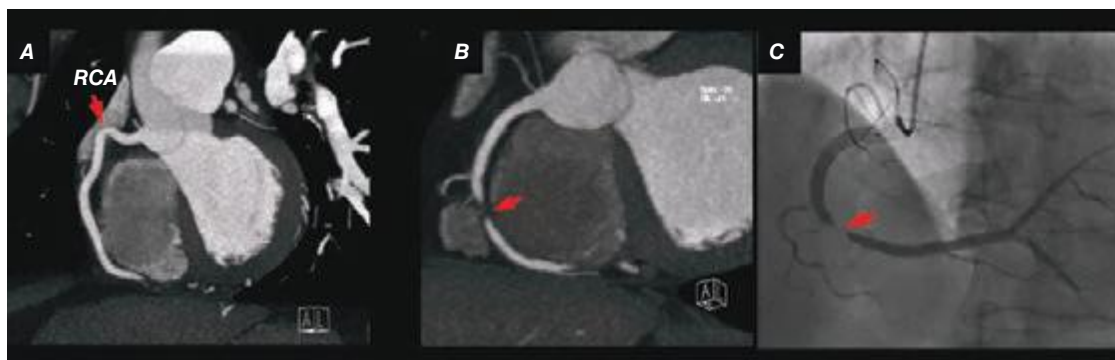


FIGURE 270e-18 Representative coronary computed tomography angiographic (CTA) images of two patients presenting to the emergency department with chest pain and negative biomarkers. The patient in **A** had angiographically normal coronary arteries; the panel shows a representative view of the right coronary artery (RCA). **B** and **C** show a corresponding significant stenosis in the mid portion of the RCA on both the CTA (**B**) and invasive angiographic view (**C**). (Images courtesy of Dr. Quynh Truong, Massachusetts General Hospital, Boston, MA.)

CMR can complement echocardiography when echocardiographic acoustic window is inadequate, quantifying blood flow data more precisely, or providing complimentary assessment of adjacent vascular structures relevant to the valvular condition.

Echocardiography can be used to assess both regurgitant and stenotic lesions of any of the cardiac valves. Typical indications for echocardiography to assess valvular heart disease include cardiac murmurs identified on physical examination, symptoms of breathlessness that may represent valvular heart disease, syncope or presyncope, and preoperative exams in patients undergoing bypass surgery. A standard echocardiographic examination should include qualitative and quantitative assessment of all valves regardless of indication and should serve as an adequate screening test for significant valvular disease.

General Principles of Valvular Assessment • DIRECT VISUALIZATION OF VALVULAR STRUCTURES Direct visualization of valve structures by two-dimensional echocardiography represents the first step in valvular evaluation. The morphology of valvular structures provides useful information regarding the etiology and severity of valvular disease. For example, two-dimensional imaging assessment of the aortic valve can identify the number of leaflets, determine whether the valve is bicuspid or tricuspid, and determine the severity of calcification and degree of leaflet excursion. Similarly, the classic appearance of a rheumatic mitral valve is extremely useful in determining the etiology of mitral stenosis, and mitral valve prolapse can be instantly identified without even the need for Doppler-based quantification.

EVALUATION OF STENOTIC VALVES As described earlier in the chapter, evaluation of stenotic valves generally includes estimation of the pressure gradient across the stenosis and determination of the valve area. Both of these measures have diagnostic and prognostic value. For example, when Doppler echocardiography is used to assess the maximal velocity across a stenotic aortic valve, this calculation will provide an accurate measure of the instantaneous gradient across the valve. This gradient will be higher than the mean gradient, as well as higher than that peak-to-peak gradient obtained at cardiac catheterization. This gradient is dependent on both the degree of stenosis and the contractile function of the left ventricle. Patients with significant left ventricular dysfunction may have severe aortic stenosis but will be unable to generate a high gradient across the valve because generated pressure within the left ventricle will be diminished.

Assessment of stenotic valves generally requires estimation of both the pressure gradient across the valve and the valve area. Pressure gradient is estimated through direct application of the Bernoulli principle, and the formula $p = 4v^2$ is usually sufficient to estimate the gradient across the valve. Several methods can be used to estimate valve areas, including the continuity principle based on the principle of conservation of mass. By this method, flow is estimated in two places. For example, for assessment of the aortic valve area, we measure the flow in the region of the left ventricular outflow tract and the cross-sectional area in this region, the product of which should be equal to the flow

across the stenotic aortic valve and its cross-sectional area. Estimation of the mitral valve area in patients with suspected mitral stenosis can also be performed in a number of ways, including planimetry of the valve directly, estimation with continuity methods, or the most commonly used pressure half-time method, in which the stenosis severity is estimated by the time it takes for the pressure—estimated from velocity by the Bernoulli equation—to reach half of its original value during mitral inflow.

EVALUATION OF REGURGITANT LESIONS Regurgitant lesions are generally assessed by both visual assessment of the valve morphology and a variety of Doppler-based methods to assess the severity of regurgitation. The etiology of regurgitation can often be inferred from visual inspection. For example, prolapse of the mitral valve leaflets—and to a lesser extent, the aortic valve leaflets—can be easily visualized with two-dimensional echocardiography. In general, valvular regurgitation can be caused by abnormalities of the valve leaflets themselves or abnormalities of the annulus and supporting structures, and these can usually be distinguished visually on transthoracic echocardiography (see discussion below).

Quantification of valvular regurgitation is more difficult with echocardiography than quantification of valvular stenoses. Doppler-based methods are best suited to assess blood velocities rather than volumetric flow. The most widely used technique for assessing the severity of valvular regurgitation is color flow Doppler estimation, which is qualitative. More quantitative methods such as the proximal isovelocity surface area (PISA) method (see below) allow for more accurate assessment of regurgitation and provide estimation of the regurgitant fraction and effective regurgitant orifice area but are less widely used. Assessment of regurgitant lesions with CMR also has a number of advantages (see below).

Assessment of Aortic Stenosis Aortic stenosis, one of the most common forms of valvular heart disease, most often occurs because of gradual progression of valvular calcification in both normal and congenitally abnormal valves. Assessment of aortic stenosis is most commonly performed with echocardiography, although techniques for quantitative assessment of aortic stenosis with CMR have been developed and increasingly used over the past decade. Echocardiographic assessment generally begins with visual inspection of the valve, usually in the parasternal long-axis and short-axis views. This allows for assessment of valvular morphology, whether it is tricuspid, bicuspid, or some variant; degree of leaflet calcification; and leaflet excursion.

The normal aortic valve consists of three leaflets or cusps: the right coronary, the left coronary, and the noncoronary cusps. Abnormalities of cusp development are some of the most common congenital heart anomalies, the most common of which is bicuspid aortic valve, with two opening leaflets rather than three (Fig. 270e-19). The aortic valve can be visualized on echocardiography, although sometimes it can be difficult to distinguish true bicuspid aortic valve from variants, including the presence of a vestigial commissure (raphe). Bicuspid aortic



FIGURE 270e-19 Normal aortic valve in the parasternal long-axis view (A) and short-axis view (B), and bicuspid aortic valve showing typical 10 o'clock to 4 o'clock leaflet orientation (C).

valve, one of the most common congenital anomalies, predisposes to both aortic stenosis and aortic insufficiency.

As discussed above, the degree of aortic stenosis is assessed by estimating both the pressure gradient across the valve and the valve area. Patients with moderate aortic stenosis or higher generally have peak instantaneous velocities of 3.0 m/s and higher, and often higher than 4.0 m/s, corresponding to pressure gradients of 36 and 64 mmHg, respectively. Because pressure gradients across the aortic valve can be underestimated in patients with severe left ventricular dysfunction, estimation of valve area by the continuity principle is the most accurate technique for assessing the severity of the stenosis. However, evaluation of the patient with so-called low-flow or low-gradient aortic stenosis can be challenging and can sometimes require provocative testing such as dobutamine echocardiography. In these cases, it is important to distinguish whether the valve is indeed capable of opening further or simply behaving like a stenotic valve because of the low-pressure gradient.

Aortic valve areas less than 1.0 cm² are generally considered severe, and valve areas less than 0.6 cm² are considered critical. Because patients with good left ventricular function can often tolerate severe aortic stenosis for a considerable period of time, valve areas or gradients alone should not be used to determine whether an individual patient should undergo aortic valve surgery, as this remains a clinical decision.

Some patients with apparent aortic stenosis actually have subvalvular or even supra-ventricular obstruction. Hypertrophic cardiomyopathy represents the classic form of subvalvular aortic stenosis, but this is usually easily distinguished from aortic stenosis on echocardiography as the valve leaflets can be seen opening during systole. Subaortic membranes can behave very similarly to leaflet aortic stenosis, and the membranes themselves can be very thin and difficult to visualize, although the presence of a murmur, a gradient across the valve with aortic leaflets that appear to open normally, is highly suggestive of a membrane. Supra-ventricular aortic stenosis, although exceedingly rare, also occurs.

The emergence of transcatheter aortic valve intervention as a therapeutic option for patients with severe aortic stenosis who are not optimal candidates for surgical replacement has resulted in a very important clinical role for multimodality imaging. Imaging plays a critical role in preprocedural planning, intraprocedural implantation optimization, and follow-up of these patients. CT plays an important role in defining the eligibility of the proposed access site (CTA of the aorta and iliac arteries) and in defining the anatomic relationships between the aortic valve and aortic root, left ventricle, and coronary ostia. Cardiac CT and transesophageal echocardiography are also used to define the device size. Transesophageal echocardiography is used during the device implantation to ensure the best prosthesis–patient match, to assess prosthesis position and function after deployment, and to identify immediate complications (e.g.,

aortic insufficiency, paravalvular leak resulting from patient–prosthesis mismatch). Echocardiography is the imaging modality of choice for long-term surveillance.

Assessment of Aortic Regurgitation Assessment of aortic regurgitation requires qualitative assessment of the aortic valve structure. Aortic regurgitation is common with congenital abnormalities of the aortic valve, the most common of which is bicuspid aortic valve. Aortic regurgitation often coexists with aortic stenosis, and it is not uncommon for patients to have both severe aortic stenosis and regurgitation. Congenital abnormalities of the aortic leaflets, such as bicuspid aortic valve, are common causes of aortic insufficiency. Dilatation of the aortic root, as occurs in patients with hypertension and other disorders in which aortic dilatation can occur, can also lead to aortic regurgitation even when the valve leaflets are intrinsically normal due to malcoaptation of the leaflets. Aortic root dilatation is common in patients with aortic regurgitation, both as a cause or coexisting lesion, and the aortic root and ascending aorta should be measured and followed in these patients (Fig. 270e-20).

Because aortic regurgitation can result in dilatation of the left ventricle over time with ultimate reduction in ventricular function, caring for the patient with aortic regurgitation requires serial assessment of ventricular size and function. Patients whose ventricles dilate beyond an end-systolic diameter of 5.5 cm or whose LVEF declines below normal are at significantly higher risk of death or heart failure, and these measures are often used to decide the need for valve surgery. Quantitation of regurgitation itself can be performed using a number of methods. Semiquantitative visual assessment of aortic regurgitant jet width and depth by color flow Doppler remains the most used. The jet diameter as a ratio of the left ventricular outflow tract diameter proximal to the valve represents one of the most reliable indices of severity and correlates well with angiographic assessment. Similarly, the vena contracta, which represents the smallest diameter of the regurgitant flow at the level of the valve, can be used to assess the severity of aortic regurgitation. Other Doppler-based methods include assessing the pressure half-time, or rate of decline of the pressure gradient between the aorta and left ventricle, a measure of acuity of aortic regurgitation, and assessing aortic flow reversal in the descending aorta. The regurgitant volume can be calculated by comparing the

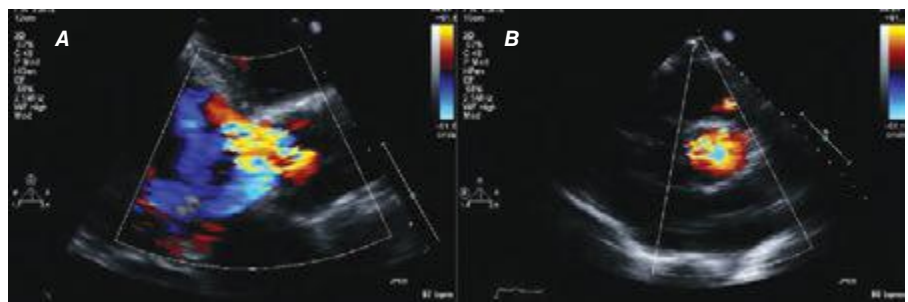


FIGURE 270e-20 Aortic regurgitation visualized by color flow Doppler in the parasternal long-axis view (A) and the parasternal short-axis view (B).

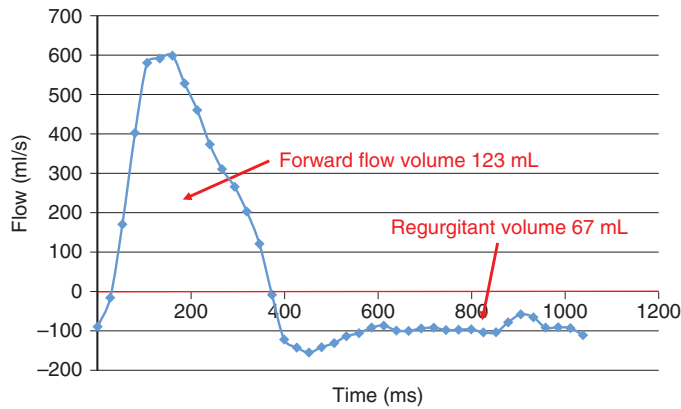


FIGURE 270e-21 The resultant flow curve generated from phase contrast imaging demonstrates a forward flow of 123 mL and a regurgitant volume of 67 mL, yielding a regurgitant fraction of 54% indicating severe aortic regurgitation.

flow across the aortic and pulmonic valves, assuming the pulmonic valve is competent.

CMR offers a number of advantages over echocardiography in the assessment of aortic regurgitation. CMR can be more accurate than echocardiography for assessing small changes in cardiac size or function that can occur over time in patients with aortic insufficiency. In addition, CMR techniques can very accurately quantify regurgitant volume in patients with aortic insufficiency, a known limitation of echocardiography. CMR can also capture three-dimensional imaging

of the aortic size that in some cases can be helpful in determining the etiology of the aortic regurgitation or in monitoring the patient (Fig. 270e-21 and Video 270e-6).

Assessment of Mitral Regurgitation The normal mitral valve consists of an anterior and posterior leaflet in a saddle shape configuration (Fig. 270e-22). The leaflets are attached to the papillary muscles via chordae tendineae that insert on the ventricular side of the leaflets. Mitral regurgitation can occur due to abnormalities of the leaflets, the chordal structures, or the ventricle, or any combination of these (Fig. 270e-23).

Mitral valve prolapse, in which one leaflet moves behind the plane of the other leaflet, can be due to myxomatous degeneration of the valves and leaflet redundancy, disruption of chordal structures secondary to degenerative disease, or papillary muscle rupture or dysfunction following myocardial infarction. Regurgitant jets can be visualized using color flow Doppler. The velocity of regurgitant jets is driven by the pressure gradient between the two chambers. This velocity tends to be quite high for left-sided regurgitant lesions, including mitral regurgitation and aortic regurgitation, resulting in turbulent jets on color flow Doppler (Fig. 270e-23). Visual estimation of color flow Doppler is generally sufficient for qualitative assessment of regurgitation severity but can dramatically under- or overestimate regurgitation severity, particularly when regurgitant jets are quite eccentric. For this reason, quantitative assessment is generally recommended, especially when making clinical decisions about surgical intervention. The PISA method is generally used for quantitative assessment of severity of mitral regurgitation. This method relies on estimation of the velocity of flow acceleration at a specific distance proximal to the valve with the assumption that the flow accelerates in concentric hemispheres.

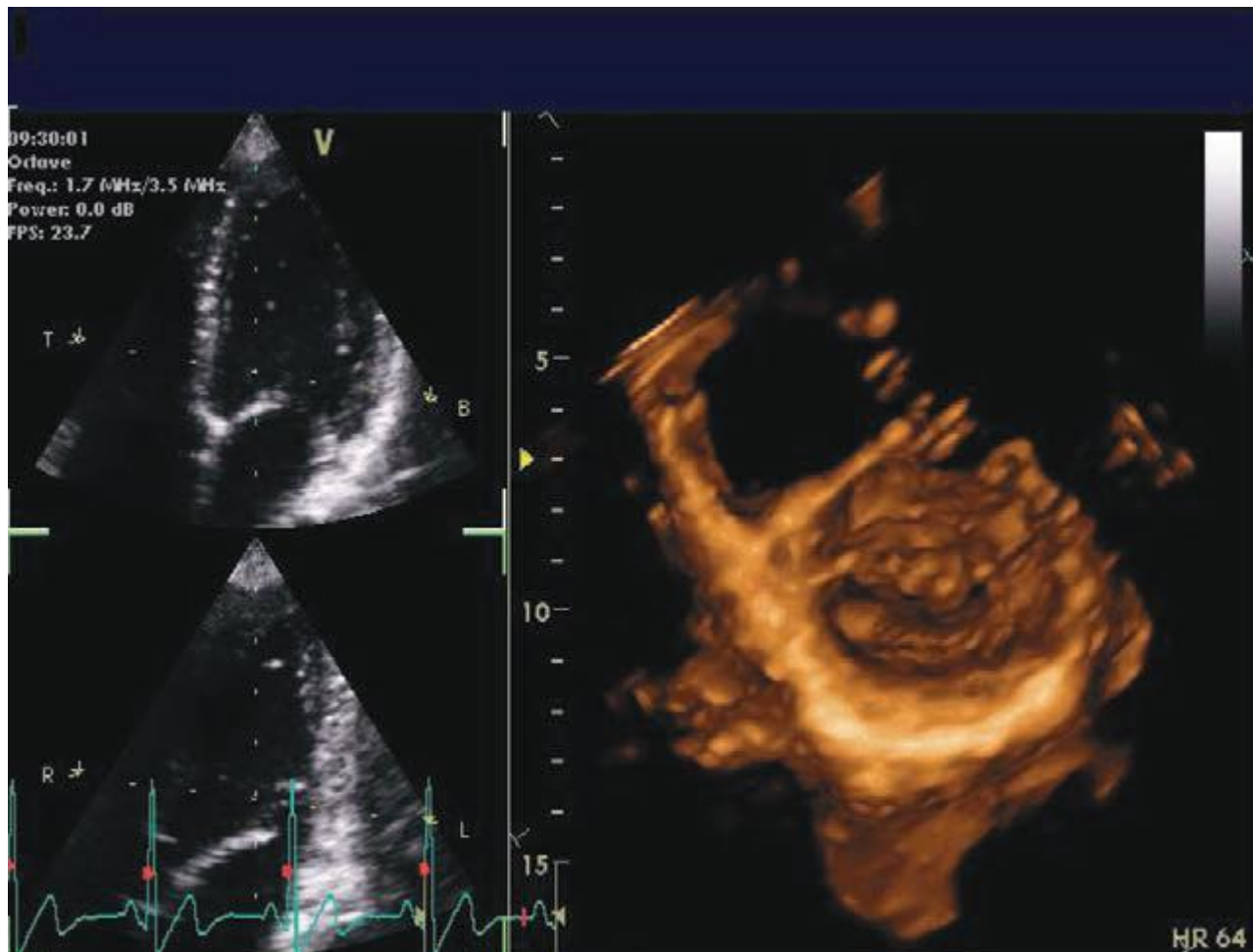


FIGURE 270e-22 Normal mitral valve in two-dimensional views (left) and with three-dimensional imaging (right).

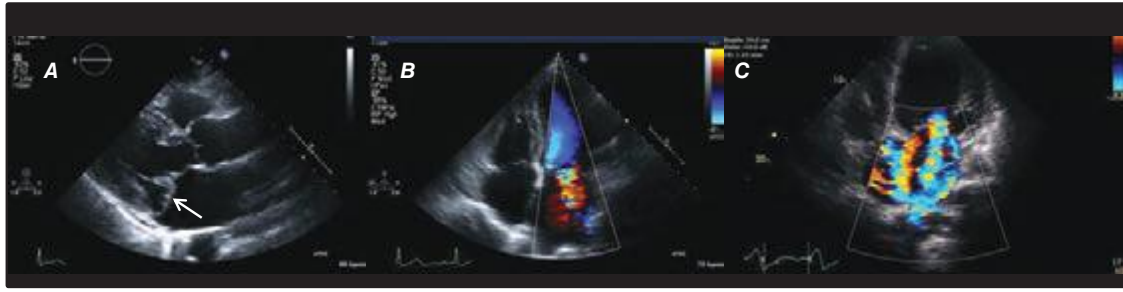


FIGURE 270e-23 **A.** Mitral valve prolapse with posterior leaflet visualized prolapsing behind the plane of the anterior leaflet (*arrow*). **B.** Color flow Doppler showing mitral regurgitation in a patient with mitral valve prolapse. **C.** Severe functional mitral regurgitation in a patient with a dilated left ventricle.

As with aortic insufficiency, assessment of ventricular structure and function is also integral in the evaluation of mitral regurgitation. Although some patients have mitral regurgitation due to intrinsic abnormalities of the valve itself, in others, the valve can be relatively normal but the mitral regurgitation can be secondary to dilatation and remodeling of the left ventricle. So-called functional mitral regurgitation is generally secondary to apical displacement of the papillary muscles in a dilated ventricle, resulting in the leaflets of the mitral valve being pulled toward the apex of the heart, resulting in poor coaptation during systole and resultant relatively central mitral regurgitation. This type of mitral regurgitation can generally be distinguished from intrinsic mitral valve disease, and the surgical or procedural treatment of these conditions can be different. Knowledge of the etiology of mitral regurgitation can be important for a surgeon planning mitral valve surgery. Moreover, new procedural approaches to mitral valve disease may be different depending on the etiology.

Ventricular dilatation is an important predictor of outcome in patients with mitral regurgitation of any cause. It is important to realize that in a patient with significant mitral regurgitation, a large portion of the blood being ejected from the left ventricle with every beat is regurgitant, thus artificially increasing the ejection fraction. Thus, an ejection fraction of 55% in a patient with severe mitral regurgitation may actually represent substantial reduction in myocardial systolic function.

CMR can be helpful in evaluating mitral regurgitation in a subset of patients when echocardiographic assessment is inadequate. CMR can directly quantify regurgitant volume of the mitral regurgitant jet or indirectly quantify regurgitant volume by measuring the difference of left ventricular stroke volume and aortic forward flow.

Assessment of Mitral Stenosis Rheumatic mitral disease remains the most common cause of mitral stenosis, although mitral stenosis can also result from severe calcification of the mitral leaflets. Rheumatic mitral stenosis has a distinct appearance characterized by tethering at the leaflet tips and relative pliability of the leaflets themselves, resulting in a hockey stick-type deformation particularly of the anterior leaflet (**Fig. 270e-24**). Narrowing of the mitral orifice impedes flow from the left atrium to the left ventricle, resulting in increased pressures in the left atrium, which are then transmitted backward into the pulmonary vasculature and the right side of the heart. When mitral stenosis is suspected, echocardiography can be useful for determining etiology (specifically whether it is rheumatic or not), estimating the valve areas and gradients across the valve, assessing the left atrium, and assessing right ventricular size and function. Assessment of left atrial size and right ventricular size and function is particularly useful in helping determine the severity of the mitral stenosis.

MYOCARDIAL INFARCTION AND HEART FAILURE

Role of Imaging after Myocardial Infarction Imaging can be particularly useful in the immediate and long-term follow-up of patients with myocardial infarction. As discussed earlier in the chapter, CMR is the best technique for direct assessment of infarcted myocardium. LGE imaging by CMR provides accurate delineation of infarct size and morphology. In a recent multicenter study, LGE imaging by CMR identified infarct location accurately and detected acute and chronic infarcts with a sensitivity of 99% and 94%, respectively. With an in-plane spatial resolution of 1.5–2 mm and a high contrast-to-noise ratio, LGE by CMR has excellent sensitivity in detecting small areas of myocardial scar. In addition, regions of microvascular obstruction (no-reflow) can be seen as dense hypoenhanced areas within the core of a bright region of infarction. Both the presence of LGE and microvascular obstruction are markers of increased clinical risk.

While echocardiography is often used to assess myocardial function immediately after myocardial infarction, myocardial stunning is common in the early post-myocardial infarction period, especially in patients who undergo reperfusion therapy. In these patients, either partial or complete recovery of ventricular function is common within several days, so that early estimation of ejection fraction may be misleading. In patients with uncomplicated myocardial infarction, imaging can generally be deferred for several days so that a more accurate assessment of cardiac function, including regional wall motion, can be assessed (**Fig. 270e-25**).

Echocardiography is the best method for assessment of patients with suspected mechanical complications after myocardial infarction. These include mitral regurgitation secondary to either papillary muscle dysfunction or rupture of papillary muscle head, ventricular septal defect, or even cardiac rupture. A new severe systolic murmur should raise suspicions for either severe mitral regurgitation or ventricular septal defect. While cardiac rupture is often catastrophic, contained ruptures, also known as pseudoaneurysms, can occur, and early diagnosis and surgical treatment are the best way to maximize survival. The presence of thrombus within the pericardial space following myocardial infarction should immediately raise suspicion of myocardial rupture and represents a surgical emergency.

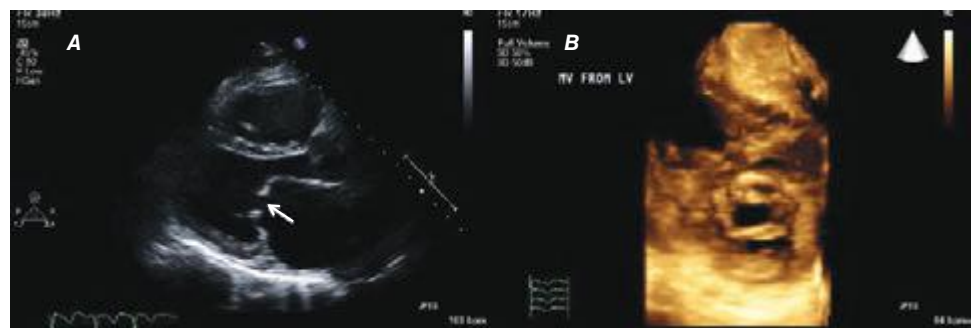


FIGURE 270e-24 **A.** Rheumatic mitral stenosis showing pliable leaflets tethered at the tips (*arrow*). Note the characteristically enlarged left atrium. **B.** Mitral stenosis visualized from a three-dimensional echocardiogram.

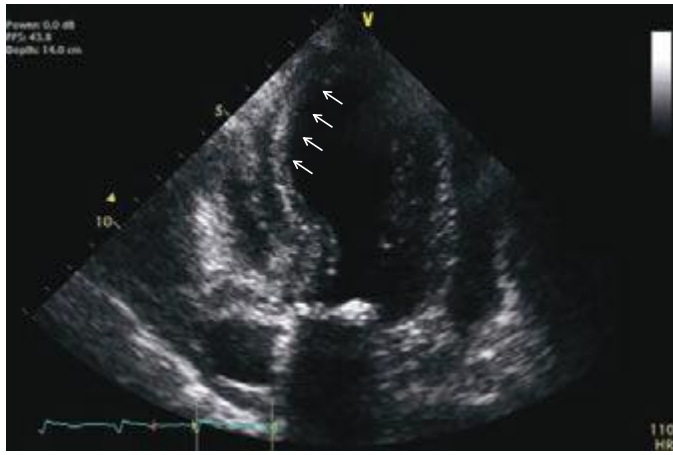


FIGURE 270e-25 Acute left anterior descending artery distribution myocardial infarction at end systole showing akinetic region (arrows).

Some patients demonstrate progressive left ventricular dilatation and dysfunction, known as cardiac remodeling, after myocardial infarction. Assessment of cardiac function and regional wall motion is useful in the follow-up period, generally between 1 and 6 months following infarction. The persistence of left ventricular systolic dysfunction following infarction is used to determine the type of therapy (e.g., angiotensin-converting enzyme inhibitors or angiotensin receptor blockers are typically used in patients with systolic dysfunction following myocardial infarction).

In patients with acute or subacute myocardial infarction, investigation of residual ischemia and/or viability is occasionally an important

clinical question, especially among those with recurrent symptoms after myocardial infarction (Fig. 270e-26). All cardiac imaging techniques can provide information regarding myocardial viability and ischemia. In the absence of definitive trials offering head-to-head comparisons between techniques in large series of patients, uncertainty persists concerning the relative accuracies of each method for predicting functional and prognostic benefit after revascularization. Thus, one should exercise caution in the interpretation of the relative diagnostic accuracy of each imaging technology. Nevertheless, the available data suggest that radionuclide imaging, especially PET, is highly sensitive, with higher negative predictive value than dobutamine echocardiography. In contrast, dobutamine echocardiography tends to be associated with higher specificity and positive predictive accuracy than the radionuclide imaging methods. The experience with CMR suggests that it offers similar predictive accuracies as those seen with dobutamine echocardiography.

Role of Imaging in New-Onset Heart Failure Echocardiography is usually a first-line test in patients presenting with new-onset heart failure. As discussed above, this test provides a direct assessment of ventricular function and can help distinguish patients with reduced ejection fraction from those with preserved ejection fraction. In addition, it provides additional structural information including an assessment of valves, myocardium, and pericardium.

Although coronary angiography is commonly performed in patients with reduced ejection fraction, the determination of heart failure etiology in an individual patient may be difficult even if angiographically obstructive CAD is present. Indeed, patients with heart failure and no angiographic CAD may have typical angina or regional wall motion abnormalities on noninvasive imaging, whereas patients with angiographically obstructive CAD may have no symptoms of angina

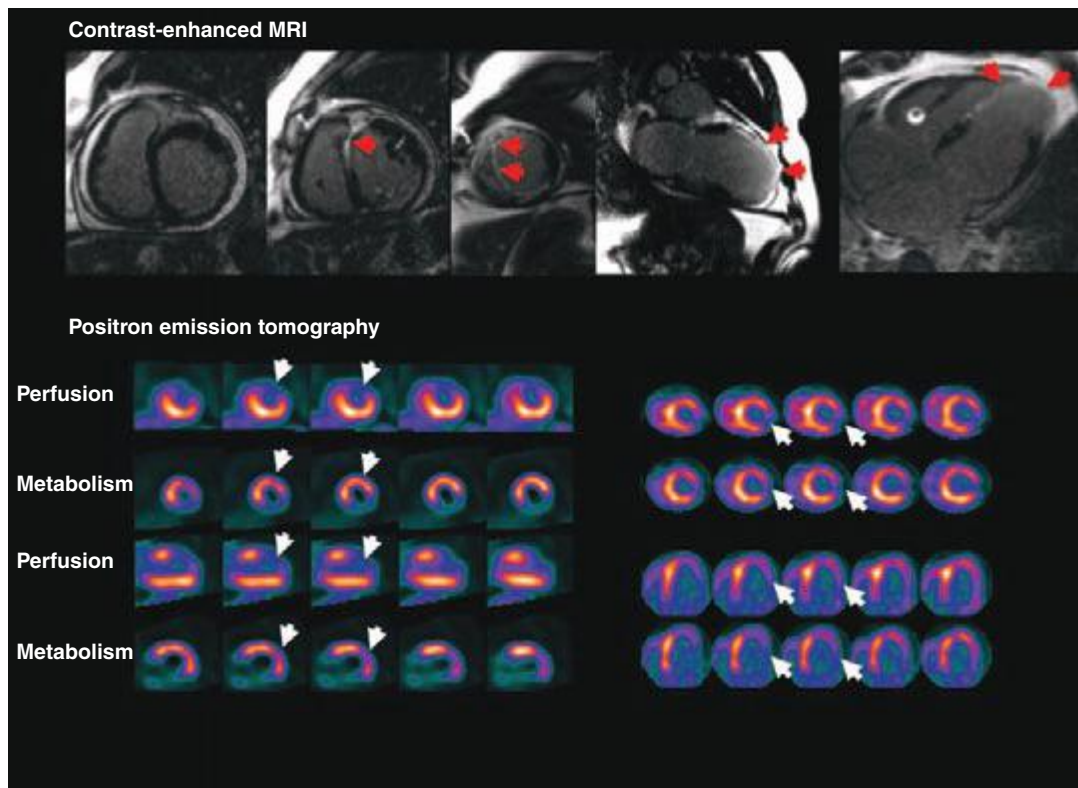


FIGURE 270e-26 Examples of myocardial viability patterns obtained with cardiac magnetic resonance imaging (MRI) and positron emission tomography (PET) in three different patients with coronary artery disease. The top panel demonstrates extensive late gadolinium enhancement (bright white areas) involving the anterior, anteroseptal, and apical left ventricular walls (arrows), consistent with myocardial scar and nonviable myocardium. The lower left panel demonstrates rubidium-82 myocardial perfusion and ^{18}F -fluorodeoxyglucose (FDG) images showing a large and severe perfusion defect in the anterior, anterolateral, and apical walls, indicating preserved glucose metabolism (so-called *perfusion-metabolic mismatch*) consistent with viable myocardium. The right lower panel shows similar PET images demonstrating concordant reduction in perfusion and metabolism (so called *perfusion-metabolic match*) in the lateral wall, consistent with nonviable myocardium.

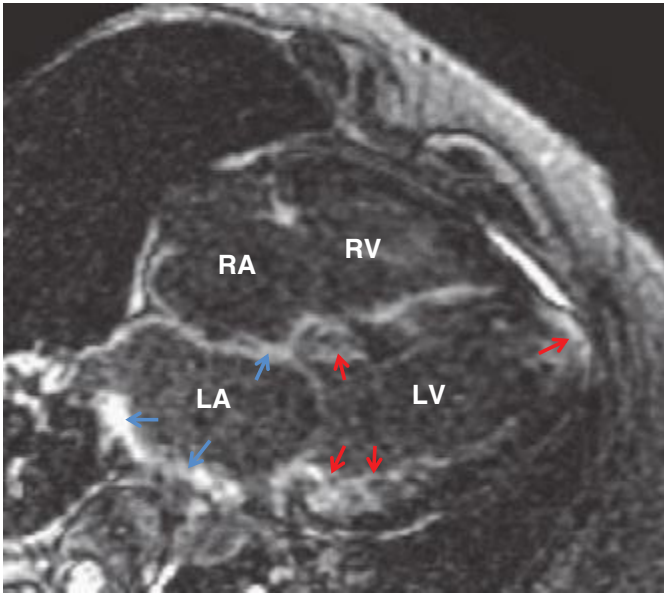


FIGURE 270e-27 A case of cardiac amyloidosis. Note on this late gadolinium enhancement image that there were multiple foci of gadolinium accumulation in the left ventricle (LV) myocardium (*red arrows*), as well as the left atrial (LA) walls (*blue arrows*). The LV walls were markedly increased in thickness, and both atria were dilated, consistent with a restrictive cardiac morphology. The blood pool signal was diminished after contrast injection, which was consistent with high burden of amyloid disease in other organs that causes gadolinium concentration in the blood to rapidly go down. RA, right atrium; RV, right ventricle.

or history of myocardial infarction. Thus, the appropriate classification for any given patient is not always clear, and it often requires the complementary information of coronary angiography and noninvasive imaging. Stress radionuclide imaging and echocardiography can be helpful in delineating the extent and severity of inducible myocardial ischemia and viability. Multiparametric CMR can be quite helpful in the differential diagnosis of heart failure etiologies. Apart from quantifying left and right ventricular volumes and function, CMR can provide information about myocardial ischemia and scar. The pattern of LGE helps differentiate infarction (typically starting in the subendocardium and involving a coronary territory) from other forms of infiltrative or inflammatory cardiomyopathies (typically involving the mid- or subepicardial layers without following a coronary distribution) (*Fig. 270e-27*). In addition, it can assess the presence of myocardial edema (e.g., myocarditis) and quantify myocardial iron deposition that can potentially lead to cardiac toxicity. Infiltrative cardiomyopathy such as amyloidosis typically has a restrictive cardiomyopathy pattern (bilateral atrial enlargement and biventricular increased wall thickness). CMR of patients with cardiac amyloidosis often also demonstrates a characteristic pattern of diffuse endocardial infiltration of the left ventricle and the atria (*Fig. 270e-27*). Hypertrophic

cardiomyopathy has variable degree of increased ventricular thickness, and often is seen to have outflow obstruction and intense LGE in regions with marked hypertrophy (*Fig. 270e-28*). CMR also can quantify myocardial iron content in patients at risk of iron-overload cardiomyopathy (*Video 270e-7*).

PET metabolic imaging has a complementary role in the evaluation of inflammatory cardiomyopathies, especially sarcoidosis. In patients with suspected cardiac sarcoidosis, the presence of focal and/or diffuse glucose uptake can help identify areas of active sarcoidosis. In addition, for patients undergoing immunosuppressive therapy, PET is frequently used to monitor therapeutic response (*Fig. 270e-29*). In patients with ischemic cardiomyopathy, radionuclide imaging in general and PET in particular are frequently used to quantify the presence and extent of myocardial ischemia and viability to assist with clinical decision making related to myocardial revascularization (*Fig. 270e-26*).

ASSESSING CARDIAC FUNCTION IN PATIENTS UNDERGOING CANCER TREATMENT

Therapies used to treat cancer can adversely affect the cardiovascular system. As the efficacy of cancer treatment and survival improve, many patients are presenting with late adverse consequences from chemotherapy and/or radiation therapy on cardiovascular function. Thus, the morbidity and mortality from late cardiovascular complications threaten to offset the early gains in cancer survival, especially among children and young adults. Early recognition and treatment of cardiomyocyte injury are critical for successful application of preventative therapies, but difficult because the adverse effects on cardiac function are a relatively late manifestation after exposure to anticancer therapy.

The accepted standard for clinical diagnosis of cardiotoxicity is defined as a >5% reduction in LVEF to <55% with symptoms of heart failure, or a >10% drop in LVEF to <55% in patients who are asymptomatic. Thus, noninvasive imaging plays a major role in diagnosing and monitoring for cardiac toxicity in patients undergoing cancer treatment. Radionuclide angiography has been the technique of choice

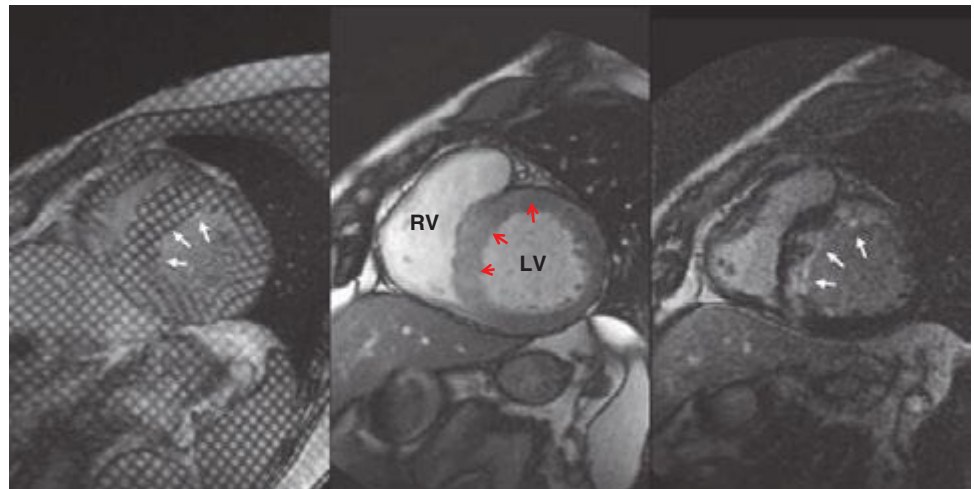


FIGURE 270e-28 This figure demonstrates three pulse sequence techniques by cardiac magnetic resonance that are often used to assess patients with hypertrophic cardiomyopathy, all displayed in the mid short-axis scan plane. The *center panel* demonstrates that the left ventricle (LV) was markedly thickened in its wall thickness especially in the LV septum (*red arrows*). This finding was matched by marked regions of late gadolinium enhancement (LGE), which was consistent with fibrosis in these segments (*right panel, white arrows*). The *left panel* was cine myocardial tagging in the same slice plane. Myocardial tagging is used to assess the normal intramyocardial strain by assessing distortion of the myocardial grids during systole. In this case, despite normal-appearing systolic radial wall thickening, the myocardial strain as assessed by the distortion of grids was markedly reduced (*left panel, white arrows*). This finding is consistent with substantial myofibril disarray in the anterior and anteroseptal segments in this patient. RV, right ventricle.

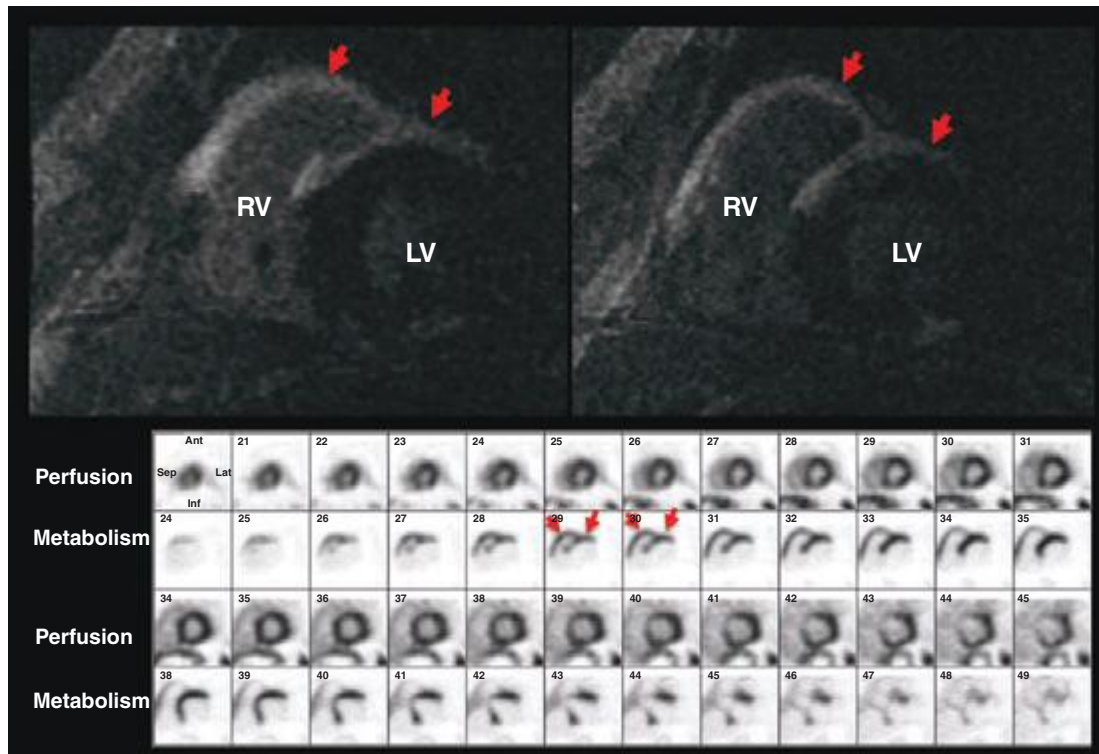


FIGURE 270e-29 Representative cardiac magnetic resonance (CMR; *top panel*) and positron emission tomography (PET; *lower panel*) images from a 45-year-old male presenting with complete heart block. The CMR images demonstrate extensive late gadolinium enhancement in the subepicardial left ventricular (LV) anterior and anteroseptal walls and also in the right ventricular (RV) free wall (*arrows*). The PET images demonstrate extensive fluorodeoxyglucose uptake in the same areas, most consistent with active inflammation due to sarcoidosis.

for quite some time. However, this is rapidly changing, and echocardiography now plays a major role in this application.

Recently, more novel imaging approaches have been advocated, including deformation imaging with echocardiography and fibrosis imaging with CMR. These techniques have shown promising results in experimental animal models and in humans. In addition, there are also proof-of-concept studies in animal models using molecular imaging approaches targeting the mechanisms of cardiac toxicity (e.g., apoptosis and oxidant stress), which can presumably provide the earliest signs of the off-target effects of these therapies. However, all of these techniques are currently considered experimental.

PERICARDIAL DISEASE

The fibroelastic pericardial sac surrounding the heart consists of a visceral, or epicardial, layer and a parietal layer, with a generally small amount of pericardial fluid in between layers. The pericardium is generally quite pliable and moves easily with the heart during contraction and relaxation. Abnormalities of the pericardium can affect cardiac function primarily by impairing the heart's ability to fill. Inflammation of the pericardium can lead to an accumulation of fluid between the two layers, or *pericardial effusion*, which can be visualized by echocardiography, CMR, or CT. Other reasons for accumulation of pericardial fluid include infection, malignancy, and bleeding into the pericardium. The latter can be the result of catastrophic processes such as trauma, cardiac rupture, perforation in the setting of a cardiac procedure, cardiac surgery, or dissection of the aorta with extension in the pericardium.

Echocardiography remains the initial test of choice for assessing pericardial disease, especially effusions (**Fig. 270e-30**). Moreover, echocardiography can be useful in evaluating for pericardial constrictive physiology, in which a thick noncompliant pericardium impairs cardiac filling. The location, size, and physiologic consequences of accumulated pericardial effusion can generally easily be determined by echocardiography. Pericardial tamponade occurs when enough pericardial fluid accumulates so that the intrapericardial pressure exceeds filling pressures of the heart, generally the right ventricle. The balance between intrapericardial pressure and ventricular pressure is

more important than the extent of fluid accumulation. Conditions in which pericardial effusions accumulate over a long period of time, as can be the case in the setting of malignant effusions, can lead to large pericardial fluid accumulations without the classic hemodynamic findings associated with pericardial tamponade. In contrast, rapid accumulations of pericardial fluid, such as those that occur due to cardiac rupture or perforation, can lead to tamponade physiology without very large effusions. In patients with suspected pericardial effusion or tamponade, echocardiography can usually be performed rapidly, at the bedside, and even by operators with limited skill. The distance from the parietal to the visceral pericardial layer can be measured, and when this exceeds approximately 1 cm, an effusion is considered significant. Echocardiographic features suggestive of tamponade include diastolic collapse of the right ventricular free wall, suggestive of pericardial pressures that exceed right ventricular filling pressures, and Doppler evidence of respiratory flow variation, which is the Doppler equivalent



FIGURE 270e-30 Pericardial effusion with tamponade physiology. The right ventricle (*arrow*) is small and collapsing in end diastole due to increased pericardial pressure.

of pulsus paradoxus. Despite the benefits of echocardiography in suspected pericardial tamponade, the diagnosis of tamponade remains a clinical diagnosis, and other important features, such as patient's blood pressure in the presence of pulsus paradoxus, needs to be taken into account when considering therapeutic options.

Chronic inflammation of the pericardium can lead to thickening and potentially calcification of the parietal pericardium, resulting in pericardial constriction in which diastolic filling can be severely impaired. In these cases, filling of the ventricles comes to an abrupt halt when the volume of ventricular filling is impaired by the constricting pericardium. Assessment of pericardial thickness in these patients is important, but it is just as important to note that approximately one in five patients with severe pericardial constriction have no significant pericardial thickening by imaging or at surgery. Thus, a lack of thickened pericardium does not rule out pericardial constriction, and patients' signs and symptomatology and physiologic evidence of constriction should be assessed independently. Pericardial constriction typically demonstrates marked respiratory changes in diastolic flow on Doppler echocardiography, in contrast to restrictive cardiomyopathy, but substantial overlap exists. CT and CMR offer tomographic, whole-heart assessment of pericardial thickening and other anatomy abnormalities in pericardial constriction (enlarged atria, vena cava, pleural and pericardial effusions) (Fig. 270e-31 and Video 270e-8). CMR offers the additional information of pericardial fibrosis and inflammation by LGE imaging, as well as evidence of constrictive physiology (e.g., regional relaxation concordance due to myocardial adhesions, paradoxical septal motion at rest or during Valsalva maneuver).

CARDIAC THROMBUS AND MASS

Echocardiography is usually the modality that first detects the presence of a cardiac mass. Differential diagnoses of an intracardiac mass most often include thrombus, tumor, or vegetation. Given their unrestricted tomographic views and multiplanar three-dimensional imaging, CMR and CT can complement echocardiography by further characterizing the physical features of the cardiac mass. Compared to CT, CMR has the advantage of higher tissue contrast differentiation, more robust cine imaging, and the use of multifaceted techniques within the same imaging session to determine the physiologic characteristics of the mass. Gadolinium contrast enhancement patterns of increased capillary perfusion can help to determine the presence and extent of

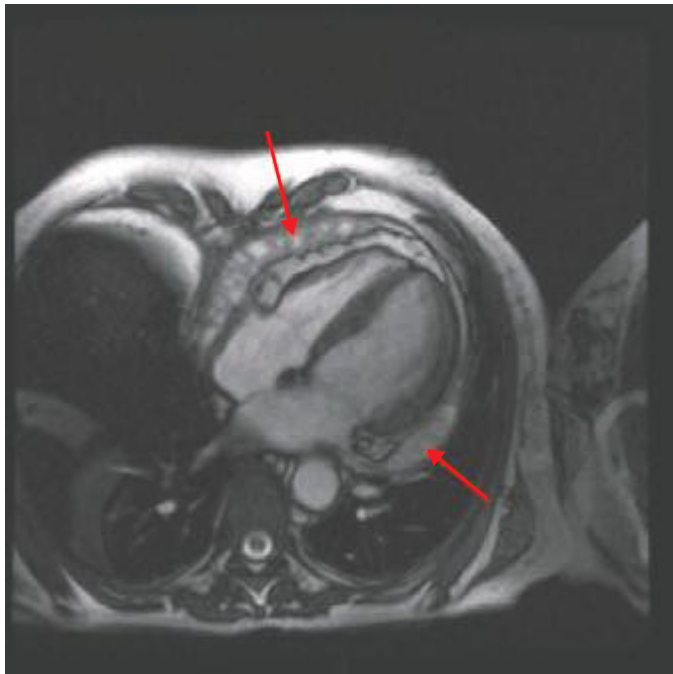


FIGURE 270e-31 A female patient developed pericardial constriction and right heart failure, secondary to radiation therapy for breast cancer. Note the multiple pericardial adhesions (red arrows).



FIGURE 270e-32 Cardiac thrombus (arrow) in an apical aneurysmal region following acute myocardial infarction.

vascularity within the mass, relevant for differentiating tumor from thrombus. Structures that are known to mimic a cardiac mass include (1) anatomic variants, such as the Eustachian valve, Chiari network, crista sagittalis or terminalis, and the right ventricular moderator band, and (2) “pseudotumors,” such as interatrial septal aneurysm, coronary or aortic aneurysm, lipomatous hypertrophy of interatrial septum, hiatal hernia, or a catheter/pacemaker lead. A number of coexisting conditions should raise the likelihood of a cardiac thrombus (Fig. 270e-32), including regional wall motion abnormality from infarction or ventricular aneurysm, atrial fibrillation leading to slow flow in the left atrial appendage, or presence of venous catheters or recent endovascular injury. CMR has the advantage of being able to assess regional wall motion and infarction or ventricular aneurysm in matching scan planes, adjacent to the cardiac thrombus, using cine and LGE imaging, respectively. For ventricular thrombus, gadolinium-enhanced LGE imaging can detect thrombus at a higher sensitivity than echocardiography by depicting high-contrast difference between the dark thrombus and its adjacent structures and by imaging in three dimensions. In addition, mural thrombus does not enhance on first-pass perfusion and often has a characteristic “etched” appearance (black border surrounding a bright center) on LGE imaging, thus providing higher diagnostic specificity than anatomic information alone (Fig. 270e-33). Comparing the signal intensities of a mass before and after contrast injection may confirm the lack of tissue vascularity (i.e., thrombus)

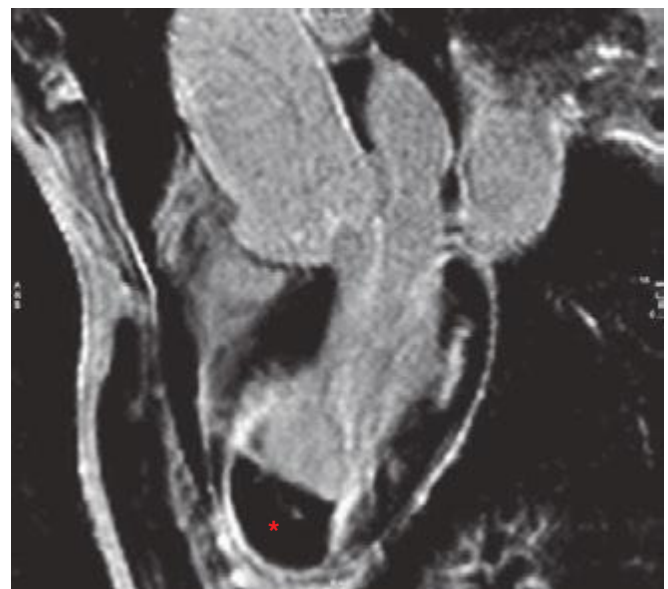


FIGURE 270e-33 Late gadolinium enhancement image of a massive anterior infarction complicated by a dyskinetic left ventricular LV aneurysm and intracavitary thrombus (red asterisk).

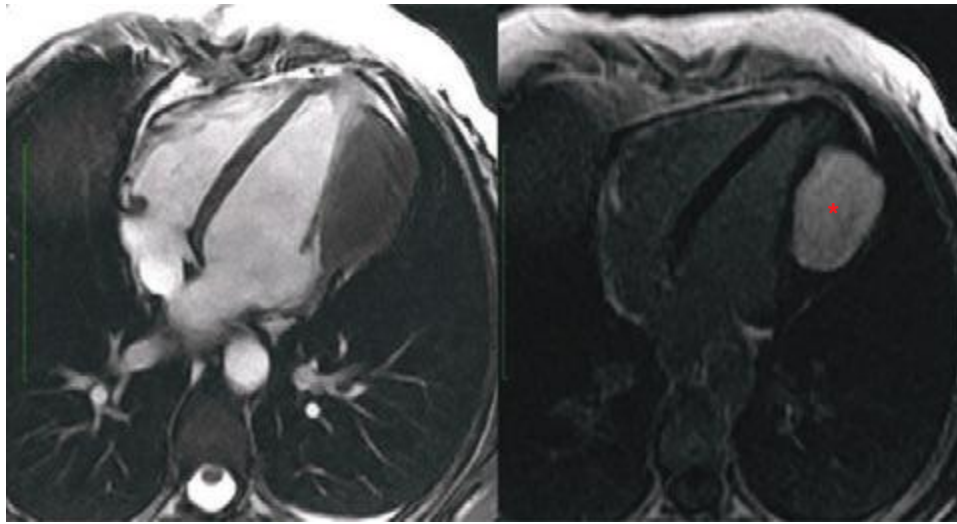


FIGURE 270e-34 A case of a cardiac fibroma. A patient presented with shortness of breath and was found to have a large myocardial mass on echocardiography. Cine cardiac magnetic resonance imaging confirmed the large myocardial mass involving the anterolateral wall. Shortly after gadolinium contrast was injected, the myocardial mass demonstrated intense accumulation of contrast on LGE imaging (*right panel, asterisk*). This is a case of cardiac fibroma. The patient also has gingival hyperplasia and bifid thoracic ribs, a part of the rare Gorlin's syndrome.

by the lack of signal enhancement after contrast administration. Like intracardiac thrombus, regions of microvascular obstruction also appear dark, but microvascular obstruction is confined within the myocardium and surrounded by infarction and thus can be differentiated from intracardiac thrombus. CMR imaging for small thrombus in the left atrial appendage is difficult due to slow flow in the atrium and rhythm irregularity from atrial fibrillation, but it may be helpful in cases where transesophageal echocardiography is suboptimal or not feasible.

The majority of cardiac malignancy is metastatic, and metastatic cardiac malignancy is far more common than primary cardiac malignancies; these metastatic involvements of the heart are the result of direct invasion (e.g., lung and breast), lymphatic spread (e.g., lymphomas and melanomas), or hematogenous spread (e.g., renal cell carcinoma). Primary benign cardiac tumors are seen mostly in children and young adults and include atrial myxoma, rhabdomyoma, fibroma, and endocardial fibroelastoma (*Fig. 270e-34*). Atrial myxomas are often seen as a round or multilobar mass in the left atrium (75%), right atrium (20%), or ventricles or mixed chambers (5%). They typically have inhomogeneous brightness in the center on cine steady-state free precession imaging due to their gelatinous contents and may have a pedunculated attachment to the fossa ovalis. Primary malignant cardiac tumors are extremely rare and include angiosarcoma, fibrosarcoma, rhabdomyosarcoma, and liposarcoma.

ROLE OF IMAGING IN INFECTIOUS AND INFLAMMATORY DISEASE

Patients with suspected endocarditis often undergo echocardiography for the purpose of identifying vegetations or intramyocardial abscesses. Vegetations are generally highly mobile structures that most typically are attached to valves or present in areas of the heart with turbulent flow. The absence of a vegetation on echocardiography does not rule out endocarditis, because small vegetations

below the resolution of the imaging techniques can be present. Echocardiography remains the best technique for assessment of vegetations because its high temporal resolution allows visualization of the typical oscillating motion, although large vegetations can be visualized with other techniques (*Fig. 270e-35*). The size and location of a vegetation do not necessarily provide any specific information about the type of infection. Abscesses, particularly around the aortic and mitral annuli, are particularly concerning in patients with endocarditis and should be suspected in patients with prolongation of cardiac intervals in the setting of endocarditis. Visualization of both vegetations and possible abscesses is best done with transesophageal echocardiography, particularly in patients with prosthetic valves. Indeed, transesophageal echocardiography is the first test of choice in a patient with a mechanical mitral or aortic valve and suspected endocarditis (*Fig. 270e-35*). Vegetations should be measured because their size has prognostic importance and can be used to decide whether a patient should be taken to surgery.

PET metabolic imaging is emerging as a potentially useful imaging technique to identify the source of infection in patients with prosthetic valves, vascular grafts, and implantable pacemakers/defibrillators, especially in patients in whom echocardiography and/or blood cultures are negative. There is an emerging literature documenting the potential value of macrophage-targeted metabolic imaging with ^{18}F -FDG and PET (*Fig. 270e-36*). Likewise, FDG PET is also useful to identify vascular inflammation and monitor the response to immunosuppressive therapy (*Fig. 270e-37*).

EVALUATION OF COMMON CONGENITAL ABNORMALITIES IN THE ADULT

While a discussion of complex congenital heart disease is beyond the scope of this chapter, a number of common congenital abnormalities are present in adults, and cardiac imaging is essential to diagnosing and managing these conditions. Abnormalities of the interatrial



FIGURE 270e-35 Vegetation on native mitral valve (*left panel, arrow*). Left atrium (LA) and left ventricle (LV) are indicated. *Middle panel* shows a vegetation on a mechanical prosthesis (St. Jude) indicated by an *arrow*; *right panel* shows vegetation on prosthesis after excision.

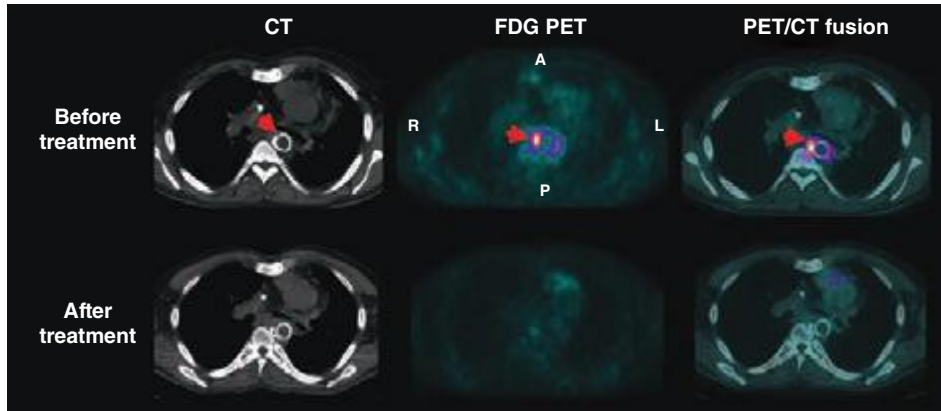


FIGURE 270e-36 Representative cross-sectional computed tomography (CT; *left*), fluorodeoxyglucose (FDG) positron emission tomography (PET; *middle*), and fused CT and PET (*right*) images before and after antibiotic treatment in a patient with fever and suspected infection of the stent placed in the descending portion of the aortic arch (*arrow*) for treatment of aortic coarctation. The FDG images before treatment demonstrate intense glucose uptake within the stent, consistent with inflammation/infection. The *lower panel* demonstrates significant attenuation of the FDG signal after treatment. (*Images courtesy of Dr. Sharmila Dorbala, Brigham and Women's Hospital.*)

septum probably represent the most common adult congenital cardiac abnormalities. Patent foramen ovale (PFO) can be identified in almost 25% of patients. In patients with PFO, a one-way flap in the region of the fossa ovalis is normally kept close by the left atrial pressure, which is generally higher than right atrial pressure for the majority of the cardiac cycle. However, right-to-left flow through a PFO can occur any time the right atrial pressure exceeds the left atrial pressure, including with maneuvers or conditions in which intrathoracic pressure is increased. The presence of a PFO can increase the likelihood of the paradoxical embolus, and thus the presence of a PFO should be determined in patients with stroke or systemic embolus of unknown etiology. Because the one-way flap of the PFO will be closed during much of the cardiac cycle, color flow Doppler will usually not reveal a PFO. Instead, agitated saline (bubble study) is the best way to assess for PFO or atrial septal defect. Saline is agitated and injected peripherally and then enters the right atrium. If no shunt is present, only the right side of the heart will be pacified because the air bubbles will be too small to traverse the lungs. Because PFO is a one-way flap, maneuvers should be used to temporarily increase right atrial pressure. Either a Valsalva maneuver or sniff maneuver can be effective.

Atrial septal defects occur most commonly in the region of the fossa ovalis, referred to as secundum-type defects (**Fig. 270e-38**). Additional atrial septal defects include defects of the sinus venosus and atrium primum. Color flow Doppler echocardiography is usually sufficient for diagnosis of a secundum-type atrial septal defect, but agitated saline is generally needed for the diagnosis of other types of atrial septal defects.

Ventricular septal defects can generally be visualized by color flow Doppler as turbulent high-velocity jets from the left to the right ventricle. In cases where the jet origin is unclear, continuous wave Doppler can estimate the velocities. These would be expected to be

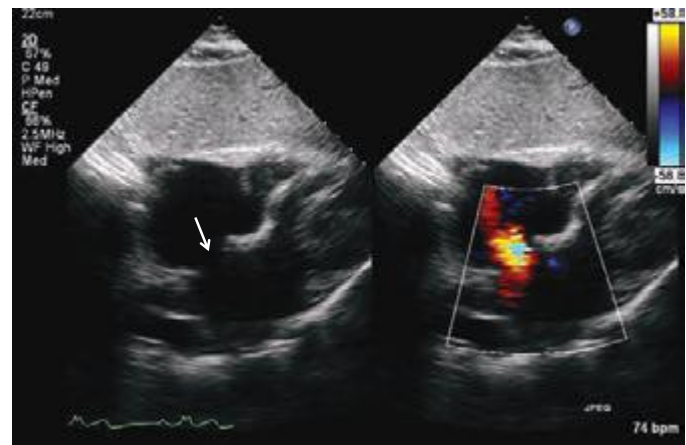


FIGURE 270e-38 Large secundum-type atrial septal defect (*arrow*) noted in the subcostal view with color flow Doppler showing flow through the defect (*right*).

extremely high to reflect the pressure gradient between the left and right ventricles. Defects can occur in both the muscular and membranous portions of the ventricular septum.

In patients with either atrial or ventricular septal defects, estimation of the severity of the left-to-right shunt is essential and can be an important determinant in management decisions. Shunts are generally assessed by echocardiography by assessing the relationship between pulmonary flow and aortic flow, the Qp/Qs ratio. Shunts and cardiac anatomy of most congenital heart diseases can also be accurately evaluated by CMR (**Fig. 270e-39**).

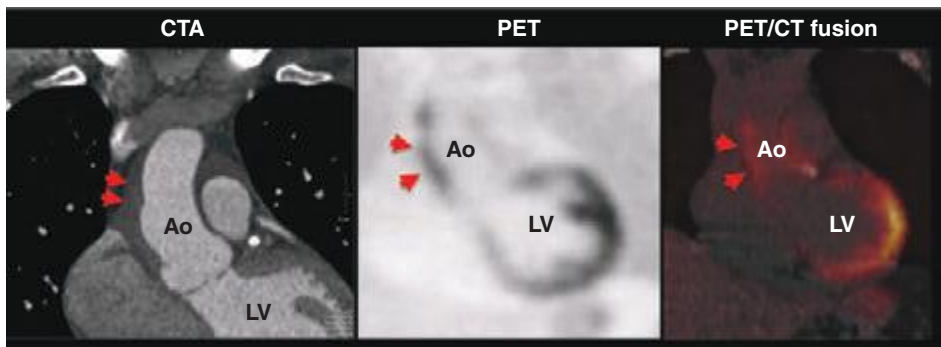


FIGURE 270e-37 Representative coronal computed tomography (CT) angiographic (CTA; *left panel*), fluorodeoxyglucose (FDG) positron emission tomography (PET; *middle panel*), and fused CT and PET (*right panel*) images in a patient with suspected aortitis. The CTA images demonstrate thickening of the ascending aorta (Ao), which correlates with intense, focal FDG uptake consistent with active inflammation. LV, left ventricle.

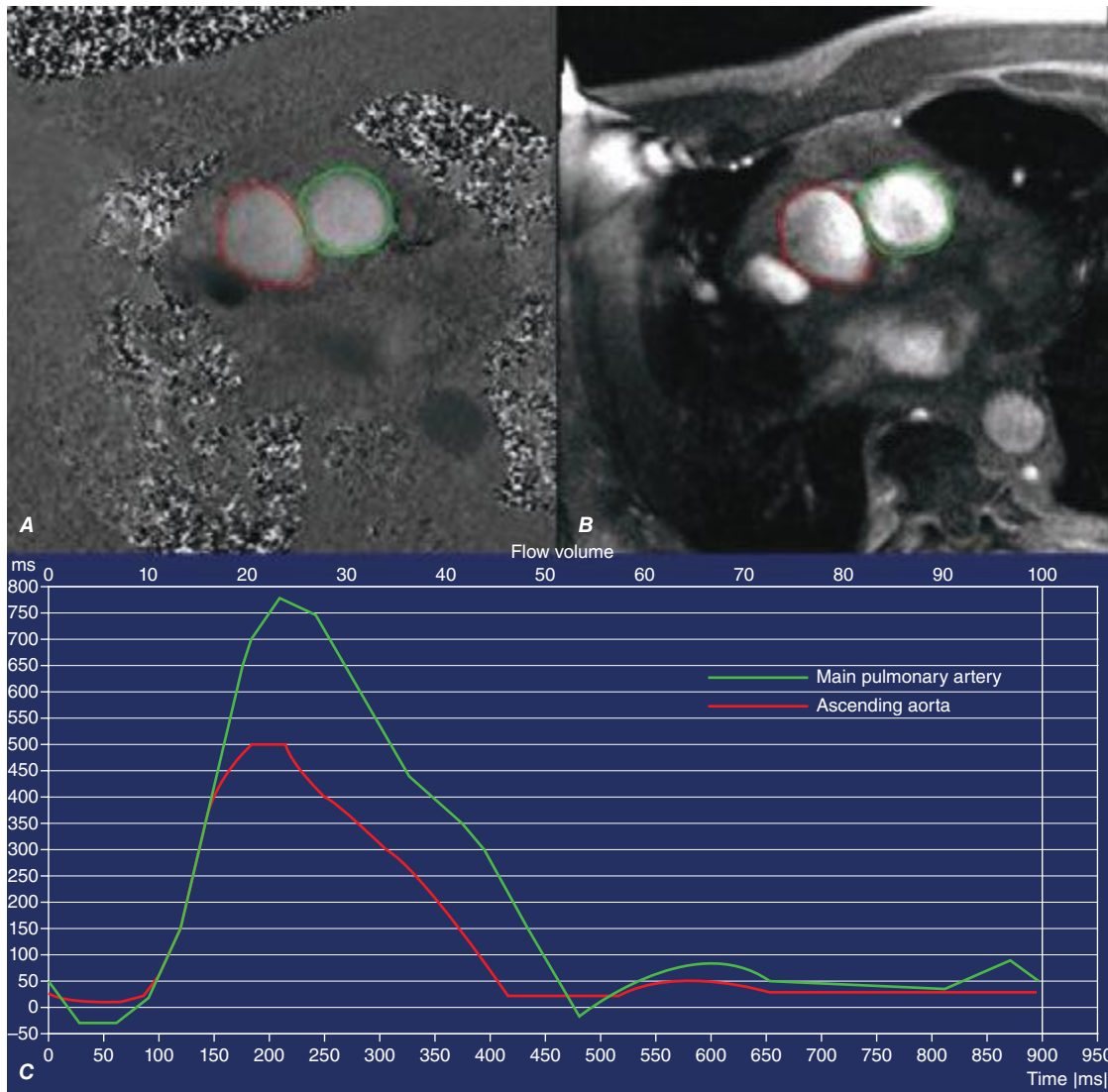


FIGURE 270e-39 **A** and **B** are phase contrast images that display blood flow (phase images on **A**) and anatomy (structural images on **B**) of the aorta (*red*) and pulmonary artery (*green*). **C** demonstrates the flow curves of the aorta (*red*) and the pulmonary artery (*green*). Note that the total flow (area under the curve) was substantially higher in the pulmonary artery than the aorta, indicative of a marked elevated pulmonary-to-systemic shunt ratio, as a result of the partial anomalous pulmonary venous return that drained into the superior vena cava.

VIDEO 270e-1 Cine steady-state free precession (SSFP) imaging (*left*) in short axis in a patient who had a large anterior myocardial infarction. Only one cut of a stack of short axis is shown. This method allows quantification of left ventricular (LV) and right ventricular (RV) volumes in diastole and systole and calculation of the LV ejection fraction, stroke volumes, and cardiac output (a product of LV stroke volume and heart rate). Note that in this case there is anterior and anteroseptal akinesia (lack of systolic wall thickening, as shown by the left cine movie, *red arrows*) matching by a near-transmural myocardial infarction as seen by the matching late gadolinium enhancement (LGE) image (*right picture, white arrows*).

VIDEO 270e-2 This is cine cardiac magnetic resonance (CMR) imaging of a patient in the long-axis four-chamber view. Note that the basal aspect of the right ventricular (RV) free wall is thickened, aneurysmal, and akinetic (*red arrows*). The global RV systolic function is mildly reduced, and the RV is dilated. CMR can image the RV using tomographic views and can quantify the RV volumes and ejection fraction volumetrically. This is a patient who presented with syncopal spells and inducible ventricular tachycardia on subsequent workup. He was diagnosed to have arrhythmogenic right ventricular dysplasia.

VIDEO 270e-3 Exercise echocardiogram showing rest images on left and poststress images on right, with parasternal long-axis, upper panel, and apical four-chamber, lower panel, end-systolic

frames. Following exercise, the distal septal/apical region becomes akinetic. A = upper left (UL); B = upper right (UR); C = lower left (LL); D = lower right (LR).

VIDEO 270e-4 The video shows cardiac magnetic resonance (CMR) myocardial perfusion imaging during vasodilating stress, in three parallel-short-axis views. An bolus of gadolinium contrast was injected intravenously while rapid imaging acquisition occurred. The contrast enhances the right ventricle first, then travels through the pulmonary circulation, enters the left ventricle (LV), and then perfuses the LV myocardium. Myocardial perfusion defects with this technique show as black subendocardial rims, reflecting lack of contrast accumulation due to ischemia and/or scar. In this case, the anterior wall has a severe perfusion defect (*red arrow*). Figure 270-14 shows the late gadolinium enhancement (LGE) image of a mid short-axis view. There is no evidence of infarction in the anterior wall, which would be seen as bright white areas, indicating that the stress perfusion defect primarily represents myocardial ischemia. This patient had a significant stenosis of the left anterior descending coronary artery.

VIDEO 270e-5 A 60-year-old female presented with intermittent chest pain of 3 days in duration but was pain free at the time of assessment in the emergency room. Admission electrocardiogram (ECG) demonstrated T-wave inversion in the anterior precordial lead, but cardiac enzymes were normal. A resting cardiac magnetic

resonance (CMR) study reviewed a large area of anteroseptal hypokinesia (*left picture*, region of hypokinesia shown by the *red arrows*), matching with a large resting perfusion defect (*middle picture*, perfusion defect shown by the *blue arrows*). Late gadolinium enhancement (LGE) imaging (*right picture*), however, did not show any enhancement to indicate any infarction in the anteroseptal wall, suggesting that the hypocontractile and hypoperfused anteroseptal wall was viable. Urgent coronary angiography demonstrated an acute thrombus in the mid left anterior descending coronary artery, which required coronary stenting. This case represents an example of acute coronary syndrome with hibernating but viable myocardium in the anteroseptal wall. The anteroseptal wall recovered contractile function when reassessed 6 months later.

VIDEO 270e-6 A patient with severe aortic regurgitation quantified by cardiac magnetic resonance (CMR). Notice the dark flow jet during diastolic across the aortic valve. For quantitation of the aortic regurgitation severity, a cross-sectional cut was made just below the aortic valve, perpendicular to the aortic regurgitation jet, using phase

contrast flow imaging. Apart from aortic regurgitation fraction and volume, CMR also can volumetrically quantify ventricular sizes and dimensions of the aorta, which are useful in monitoring patients with aortic valve diseases.

VIDEO 270e-7 These are T2* images of the heart (*left panel*) and the liver (*right panel*) of a patient who has hemochromatosis.

Note that iron and the liver are markedly darkened in these movies, indicating high load of iron in the heart muscle and liver. The rate of signal reduction (decay) in the myocardium and liver can be calculated as T2* value expressed in milliseconds. In this case, the T2* was at 10 ms. T2* <20 ms in patients with cardiomyopathy has been shown to indicate iron toxicity as the etiology of the cardiomyopathy, and it carries prognostic value for such patients at risk of cardiac iron toxicity.

VIDEO 270e-8 This video shows the heart in long and short axis.

Note the large atria, thickened pericardium, and extensive pericardial adhesions. Given the extensive pericardial adhesions, there is little shearing motion of the ventricles against the parietal pericardium.

This chapter provides “movie” image clips as they are viewed in clinical practice, as well as additional static images. Noninvasive cardiac imaging is essential to the diagnosis and management of patients with known or suspected cardiovascular disease. This atlas supplements [Chap. 270e](#), which describes the principles and clinical applications of these important techniques.



FIGURE 271e-1 A 48-year-old man with new-onset substernal chest pain. Echocardiography shows evidence of acute anterior myocardial infarction involving the interventricular septum and apex secondary to an occlusion of the left anterior descending coronary artery seen from the parasternal long axis view (*left*) and the apical four-chamber view (*right*). LV, left ventricle; RV, right ventricle. (See [Videos 271e-1](#) and [271e-2](#).)

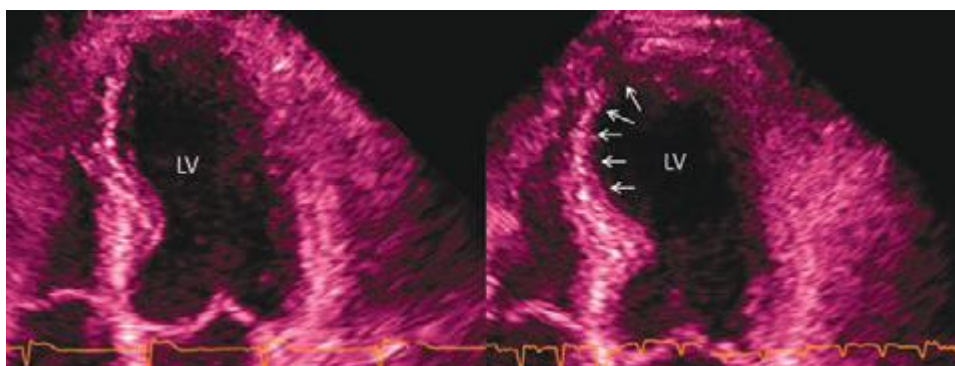


FIGURE 271e-2 A 55-year-old man with exertional chest discomfort and dyspnea. He exercised for 12 min on a standard Bruce protocol, experiencing typical chest pain and ST-segment depression in V_2-V_5 . End-systolic frame of a stress echocardiogram shows apical four-chamber view at rest (*left*) and after exercise (*right*). After exercise, there is a clear regional wall motion abnormality in the distal septum through the apex, consistent with a stenosis in the left anterior descending artery distribution (*arrows*). LV, left ventricle. (See [Videos 271e-3](#) and [271e-4](#).)

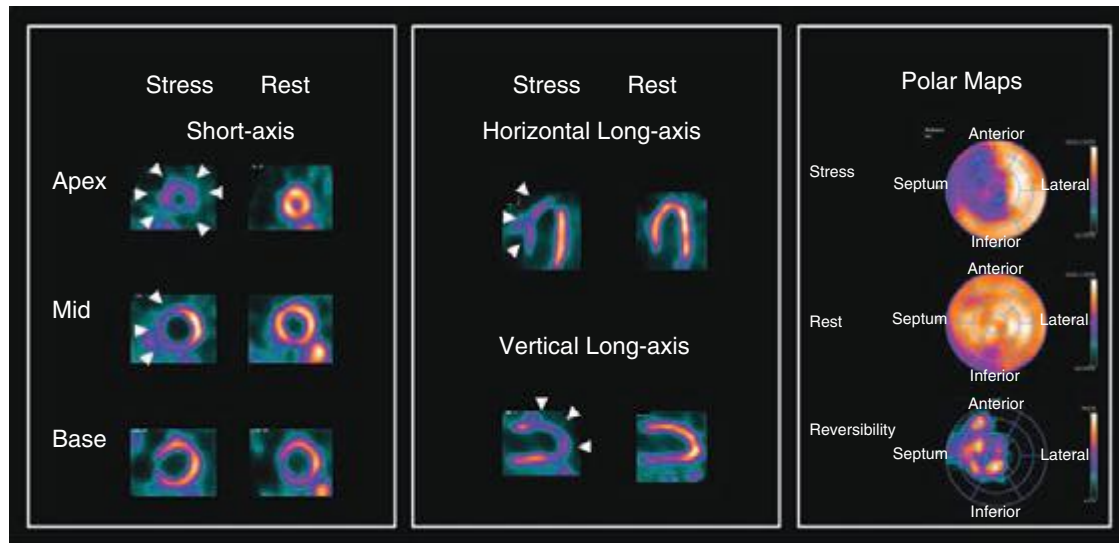


FIGURE 271e-3 Exercise single-photon emission computed tomography (SPECT) myocardial perfusion technetium-99m (^{99m}Tc) sestamibi scan in a 54-year-old male with a history of coronary artery disease and a prior coronary stent. The stress images (left and middle) show a large defect involving the apex, all apical segments, mid-inferior, mid-inferoseptum, and mid-anteroseptum (arrowheads), which is completely reversible at rest (right), reflecting a large area of exercise-induced myocardial ischemia throughout the left anterior descending coronary territory. The bull's eye displays on the right panel depict the semiquantitative extent of ischemia (light yellow and blue areas represent the extent and severity of ischemia).

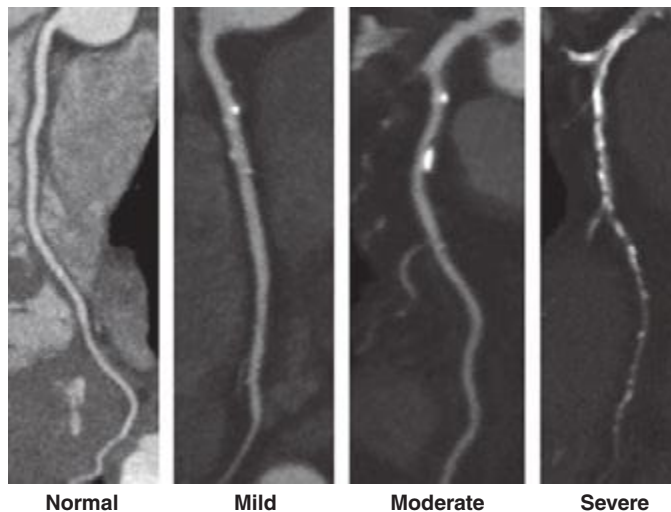
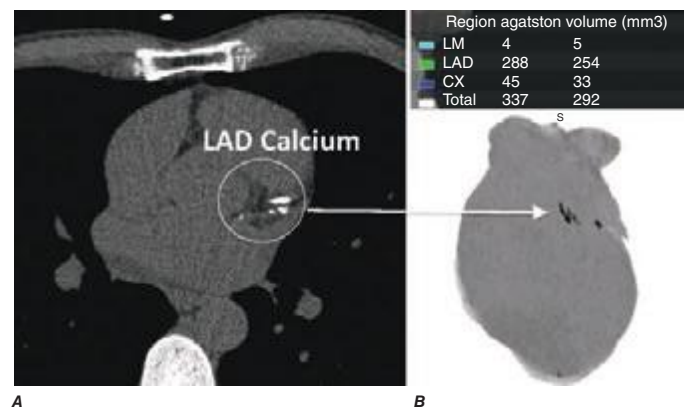


FIGURE 271e-4 Coronary computed tomography angiography (CTA). Curved multiplanar reformations demonstrating coronary artery disease severity, defined as normal (no plaque or stenosis), mild (<40%), moderate (40–69%), and severe (>70%) luminal narrowing. By guidelines for CTA reporting, alternative classification provides for stenosis grading as normal, minimal (1–24%), mild (25–49%), moderate (50–69%), severe (70–99%), and occluded (100%). (From GL Raff et al: SCCT guidelines for the interpretation and reporting of coronary computed tomographic angiography. *J Cardiovasc Comput Tomogr* 3:122, 2009; with permission.)



- CAC score 271e = 96th percentile for age, race and ethnicity¹
 - 10-year hard CHD risk is 6% (observed age) vs 30% (arterial age)^{2#}

FIGURE 271e-5 Coronary artery calcium (CAC) scan on a 51-year-old white male without clinical cardiovascular disease or treated diabetes, referred for CAC for risk stratification to guide preventive therapies. **A.** Gated, noncontrast cardiac computed tomography (CT; 3-mm slice thickness), axial view, demonstrating calcified left anterior descending (LAD) artery atherosclerosis. **B.** Whole-heart three-dimensional image reconstruction, inverted maximum-intensity projection, demonstrating overall burden of CAC with predominant LAD distribution (arrow). **Top right.** CAC scores for each coronary artery with calcified plaque involvement, scored by Agatston method and total volume. #For white male with observed age 51 years, total cholesterol 220 mg/dL, high-density lipoprotein 45 mg/dL, nonsmoker, no hypertension, and systolic blood pressure 120 mmHg. Calculated arterial age is 81 years. CHD, coronary heart disease; CX, left circumflex artery; LM, left main artery. ¹Data from RL McClelland et al: *Circulation* 113:30-37, 2006. ²Data from RL McClelland et al: *Am J Cardiol* 103: 59-63, 2009.

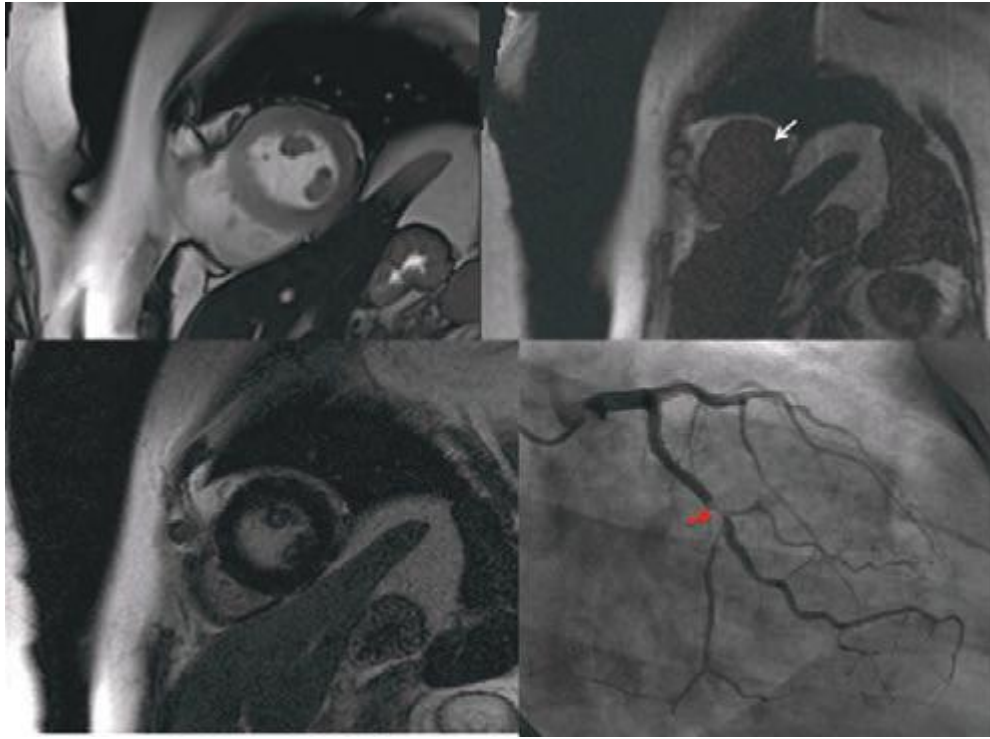


FIGURE 271e-6 Cardiac magnetic resonance (CMR) stress myocardial perfusion images in a 60-year-old patient with atypical chest pain. Cine movie short-axis image (*left upper panel*) shows normal left ventricle size and global and regional function at rest. During vasodilator stress, there is marked reduction of lateral wall perfusion (*white arrow, right upper panel*) as well as a mild defect in the septal wall. This region is confirmed to be viable by matching late gadolinium enhancement imaging (*left lower panel*), which demonstrated no evidence of infarction in the lateral wall. These findings are consistent with a severe coronary stenosis in the left circumflex artery. On angiography performed subsequently, there is a tight lesion in the left circumflex artery (*red arrow, right lower panel*). (See Videos 271e-5 and 271e-6.)

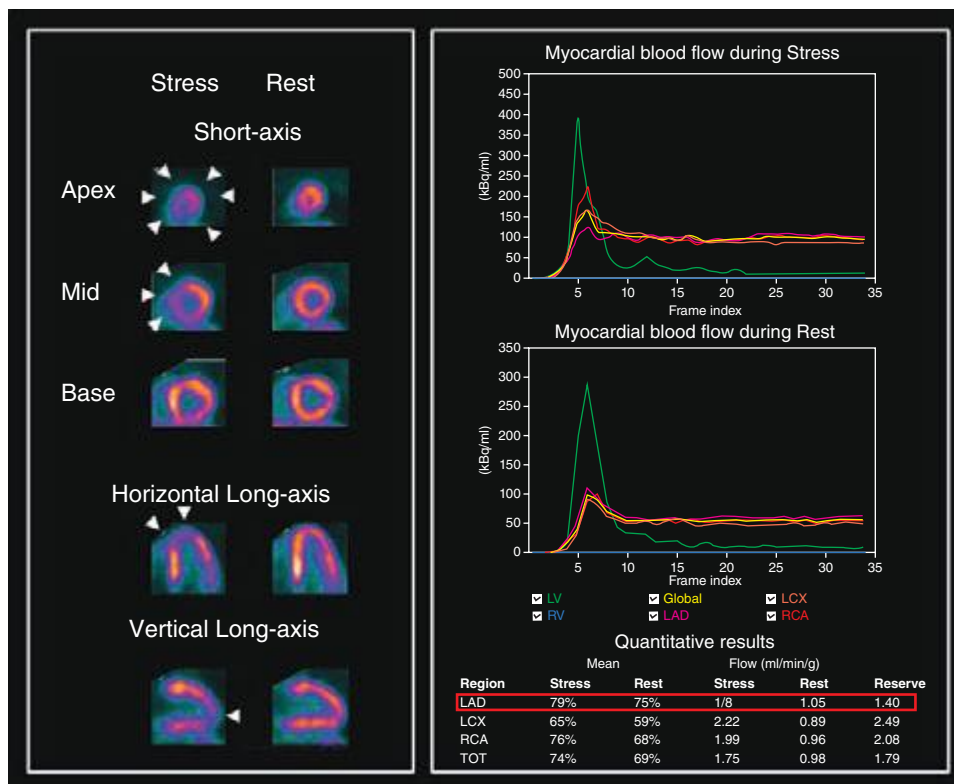


FIGURE 271e-7 Adenosine positron emission tomography (PET) myocardial perfusion ^{13}N -ammonia scan in a 60-year-old female with atypical chest pain. The stress images (*left*) show a large defect involving the apex, all apical segments, mid-inferior, mid-inferoseptum, and mid-anteroseptum (*arrowheads*), which is completely reversible at rest (*right*). This is consistent with a medium-sized area of stress-induced ischemia in the mid portion of the left anterior descending (LAD) coronary artery. The *right panel* illustrates the time-activity curves used for quantification of myocardial blood flow (in mL/min per g of tissue) at peak stress (*upper panel*) and at rest (*lower panel*). Coronary flow reserve is then calculated as the ratio of stress/rest myocardial blood flow. The coronary flow reserve is abnormal in the LAD territory, and normal in the left circumflex (LCX) and right coronary artery (RCA) territories (i.e., >2.0). TOT, total left ventricle.

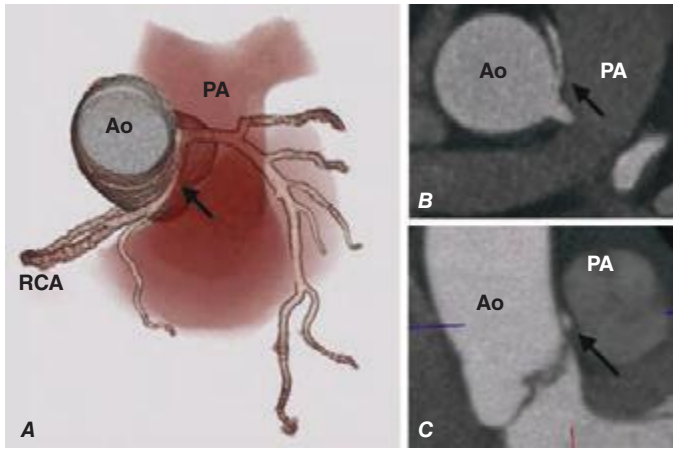


FIGURE 271e-8 Coronary computed tomography angiography (CTA) obtained on a 35-year-old female presenting to an outpatient clinic with a history of unexplained syncope and a 6-month complaint of intermittent, atypical chest pain occurring primarily during rest. Physical examination is normal. An exercise treadmill test is performed demonstrating good exercise capacity with no exertional chest pain or ischemic ECG changes. For persistent, unexplained symptoms, coronary CTA is obtained. **A**, Three-dimensional cardiac CT image reconstruction demonstrating anomalous right coronary artery (RCA) origin from the left coronary cusp with an acute angle takeoff (*arrow*) and an intraarterial course between the aorta (Ao) and main pulmonary artery (PA). **B**, **C**, Contrast-enhanced CTA in two-dimensional axial (**B**) and coronal oblique views (**C**) demonstrating proximal RCA intraarterial course between the Ao and main PA.

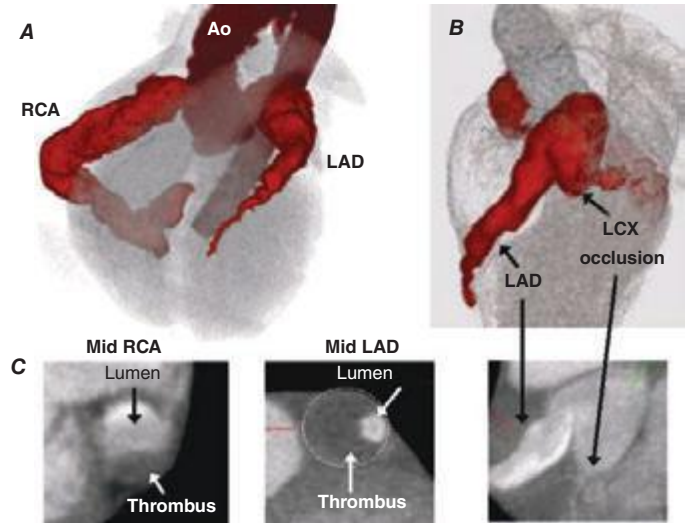


FIGURE 271e-9 Coronary computed tomography angiography (CTA) obtained from a 13-year-old boy with a history of Kawasaki disease who presented with limited exercise capacity and occasional, atypical chest pain. **A**, **B**, Three-dimensional cardiac CT image reconstruction demonstrating large three-vessel coronary artery diffuse aneurysms with proximal, nondominant left circumflex (LCX) artery occlusion. **C**, Two-dimensional contrast-enhanced coronary CTA demonstrating mid-RCA and mid-LAD thrombi that are nonocclusive layered and near circumferential thrombi, respectively, and proximal LCX occlusion. Ao, aorta; CT, computed tomography; LAD, left anterior descending; RCA, right coronary artery.

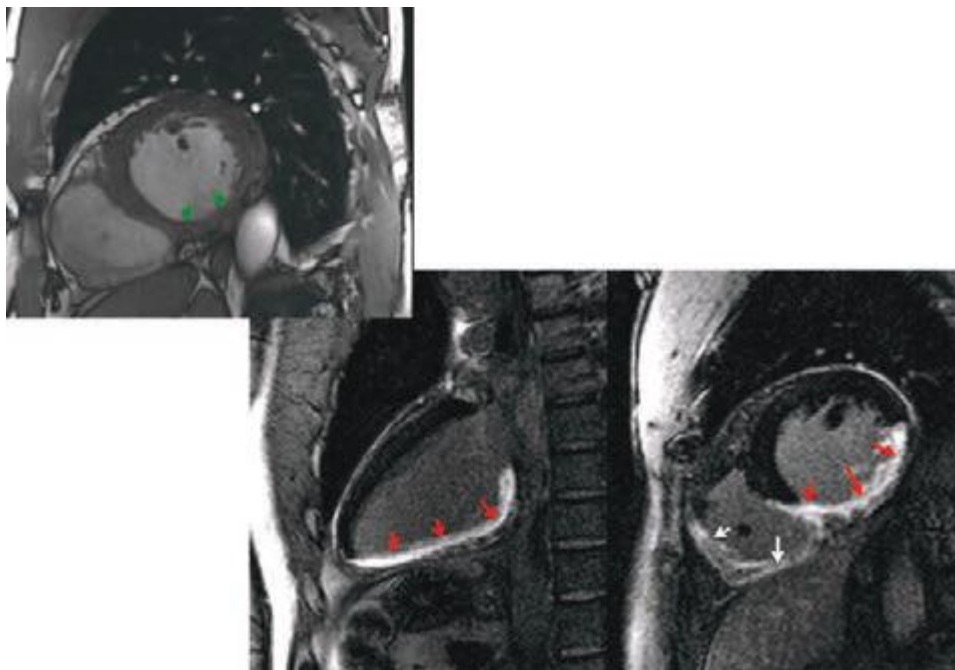


FIGURE 271e-10 A case of viability assessment in a patient with inferior myocardial infarction. The cine movie in the *upper panel* shows an area of inferior akinesis (*green arrows*). Magnetic resonance image demonstrates transmurular contrast enhancement of the inferior wall (*red arrows*) and the right ventricle (*white arrows*), which is consistent with infarction. Imaging the heart 10–15 min after injection of gadolinium allows for the accumulation of gadolinium in infarcted tissue (*red arrows*), which identifies nonviable infarcted myocardium as bright. Viability assessment, as in this case, can provide guidance for any benefits of invasive coronary intervention. In this case, the inferior wall is nonviable. Apart from the inferior wall infarction (*red arrows*), there is extensive right ventricular infarction (*white arrows*). (See [Video 271e-7](#).)

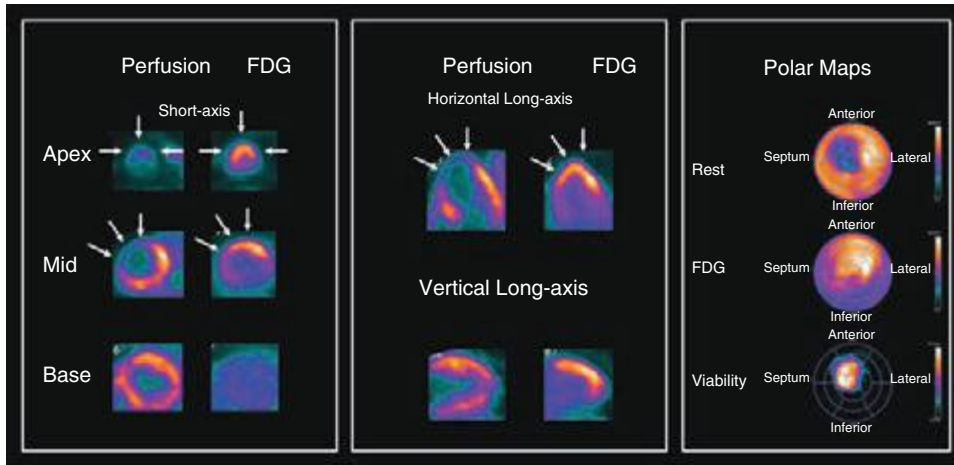


FIGURE 271e-11 Rest myocardial perfusion and metabolism positron emission tomography (PET) scan obtained with ^{13}N -ammonia (perfusion) and ^{18}F -fluorodeoxyglucose (FDG; glucose metabolism) in a 48-year-old male with a prior myocardial infarction. The rest perfusion images show a large defect involving the apex, apical segments, and mid-anteroseptal and anterior segments (*arrowheads*), which has associated increase in glucose uptake (perfusion-metabolic mismatch), reflecting viable but hibernating myocardium throughout the left anterior descending coronary territory.

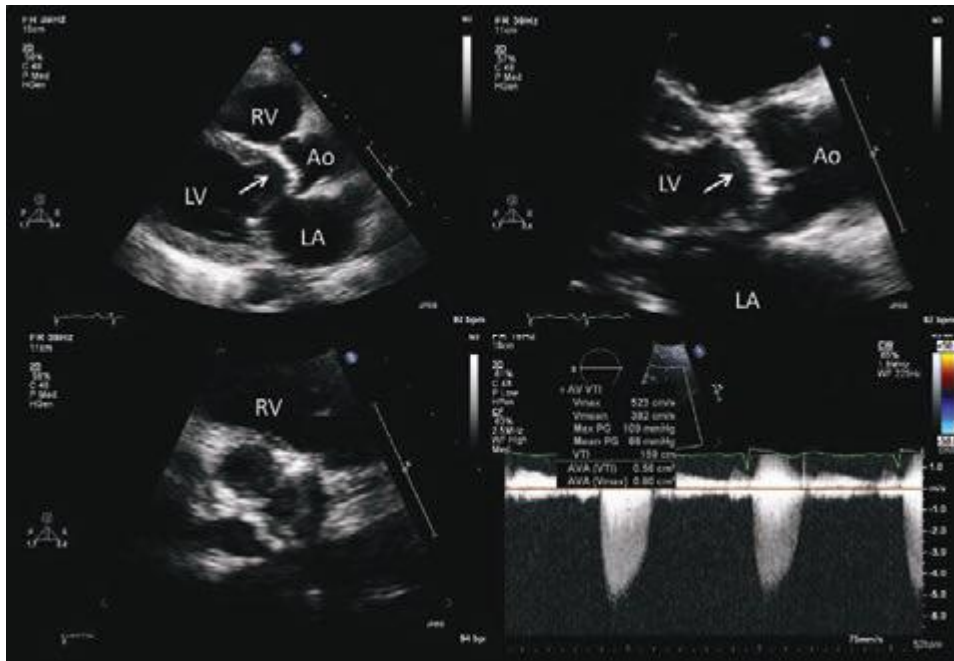


FIGURE 271e-12 A 70-year-old patient with known cardiac murmur and progressive shortness of breath and a recent episode of syncope. Echocardiography shows severe calcific aortic stenosis. A heavily calcified aortic valve (*arrow*) is shown in the parasternal long-axis views (*top panels*) and short-axis view (*bottom left*). Doppler interrogation shows a peak transaortic velocity of 5.2 m/s consistent with a peak instantaneous gradient of 109 mmHg and a mean gradient of 66 mmHg, and a corresponding aortic valve area of $<0.6\text{ cm}^2$ (*lower right*). Ao, aorta; LA, left atrium; LV, left ventricle; RV, right ventricle. (See Videos 271e-8, 271e-9, and 271e-10.)



FIGURE 271e-13 A 66-year-old patient with multiple myeloma and progressive shortness of breath. Echocardiography shows features typical of cardiac amyloidosis, including thickened myocardium with a “sparkly” appearance and left atrial enlargement. Systolic function is mildly reduced, and diastolic function is severely reduced. LA, left atrium; LV, left ventricle; RV, right ventricle. (See Videos 271e-11 and 271e-12.)

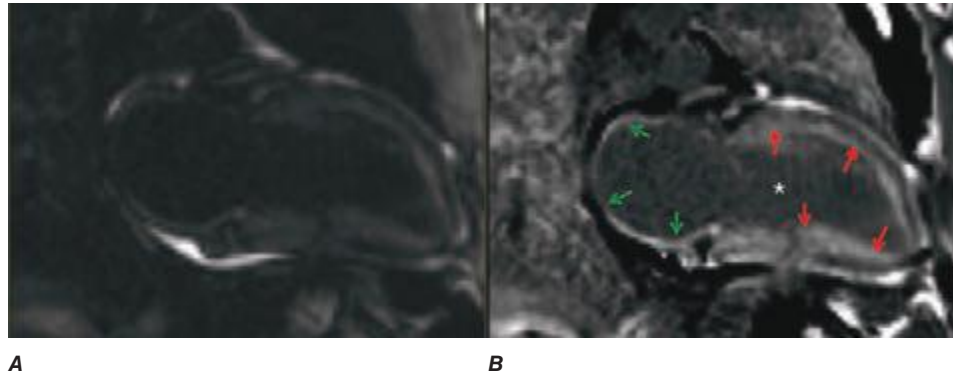


FIGURE 271e-14 Magnetic resonance image with contrast enhancement in magnitude (A), and phase-sensitive reconstructed images (B) 5–10 min after injection of gadolinium in a patient with transthyretin (TTR)-mediated amyloidosis. The phase-sensitive reconstruction (B) enhances the region of abnormal collection of gadolinium, making gadolinium enhancement in the ventricle (red arrows) and the atrium (green arrows) more prominent. Amyloidosis causes accumulation of abnormal interstitial proteins, which results in late gadolinium enhancement in a diffuse subendocardial pattern (red arrows). Blood pool signal is characteristically dark (asterisk) owing to sequestration of gadolinium into other organs.

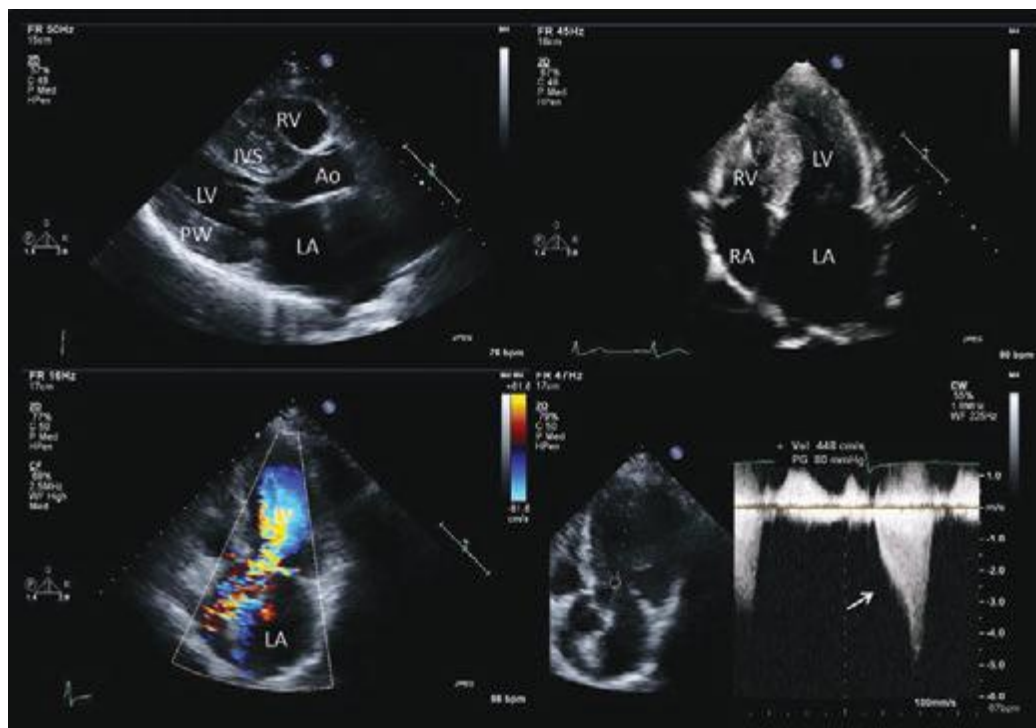


FIGURE 271e-15 A 34-year-old woman with known cardiac murmur and syncope with a family history of sudden cardiac death. Echocardiogram shows classic findings of hypertrophic cardiomyopathy, including marked left ventricular wall thickness, particularly in the interventricular septum, notable in the parasternal long-axis view (upper left) and apical view (upper right). Note reverse septal curvature in the apical view (upper left). There is substantial flow acceleration through the left ventricular outflow tract (lower left) with evidence of a late peaking systolic gradient (arrow, lower right) caused by outflow tract obstruction. Ao, aorta; IVS, interventricular septum; LA, left atrium; LV, left ventricle; PW, posterior wall; RV, right ventricle. (See Videos 271e-13, 271e-14, and 271e-15.)



FIGURE 271e-16 Magnetic resonance image with contrast enhancement in a patient with hypertrophic cardiomyopathy. Note the marked thickened anteroseptal wall (*black arrows, left panel*) consistent with asymmetric septal hypertrophy. After contrast is injected, this region demonstrated heterogeneous foci of contrast enhancement (*right panel, red arrows*) consistent with myocardial fibrosis due to myofibril disarray in this condition. This typical enhancement pattern of hypertrophic cardiomyopathy is found in the areas of maximum wall thickness, typically at the anteroseptum as in this case. (See Video 271e-16.)

Nonischemic cardiomyopathy sarcoid (with both MRI and PET)

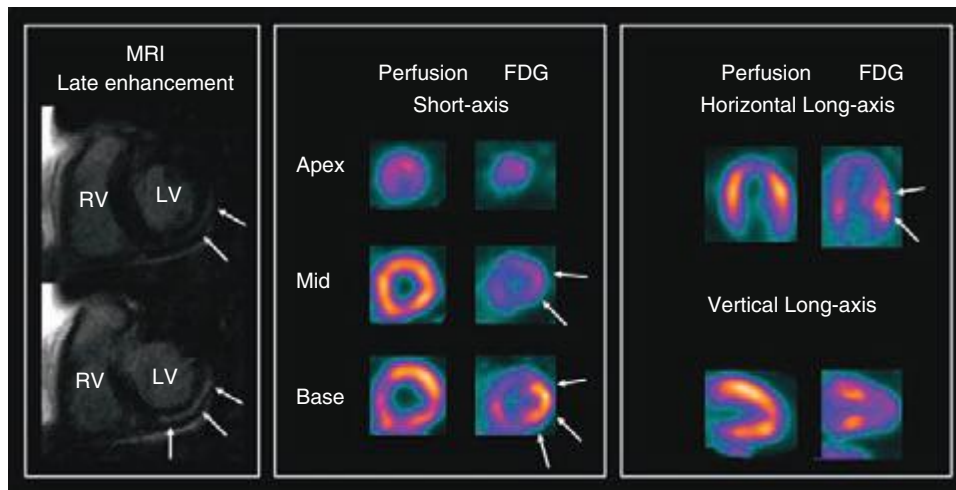


FIGURE 271e-17 Late gadolinium enhancement cardiac magnetic resonance imaging (MRI) (*left panel*) and rest myocardial perfusion and fluorodeoxyglucose (FDG) positron emission tomography (PET)/computed tomography (CT) studies (*middle and right panels*) in a 48-year-old male with a complete heart block. The MRI images demonstrate a linear area of mid-wall late gadolinium enhancement involving the inferior and inferolateral walls (*arrows*). The myocardial perfusion images were normal. However, the FDG images demonstrated a focal area of intense glucose uptake in the inferolateral wall, corresponding to the area of late enhancement on MRI. This is consistent with focal active sarcoidosis. LV, left ventricle; RV, right ventricle.

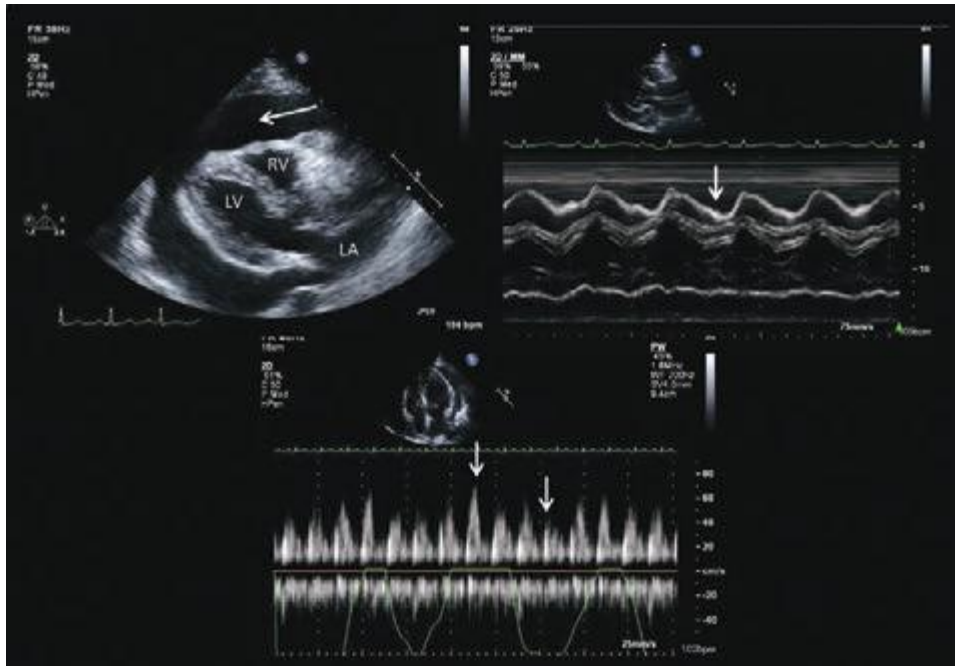


FIGURE 271e-18 A 46-year-old patient with malignant melanoma who presents with acute shortness of breath. Echocardiogram reveals a large pericardial effusion (arrow, upper left) with evidence of cardiac tamponade. M-mode echocardiography (upper right) shows evidence of collapse of the right ventricular free wall during diastole (arrow). Doppler echocardiography (lower panel) shows evidence of respiratory flow variation, consistent with a pulsus paradoxus. LA, left atrium; LV, left ventricle; RV, right ventricle. (See Video 271e-17.)

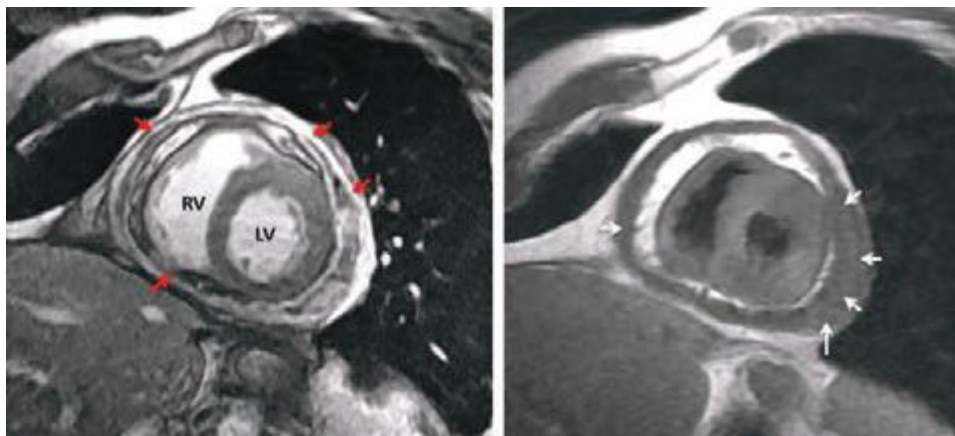


FIGURE 271e-19 Diffuse pericardial thickening (left; red arrows) and circumferential effusion (right; white arrows) associated with effusive-constrictive pericarditis. Effusive-constrictive pericarditis is a progressive condition that has varying degrees of hemodynamic consequences due initially to the collection of pericardial fluid and ultimately to pericardial constriction. It is typically suspected in cases where pericardiocentesis fails to normalize intracardiac pressures. In this example, pericardial fluid analysis resulted in a sterile exudate of leukocytes and erythrocytes. LV, left ventricle; RV, right ventricle.

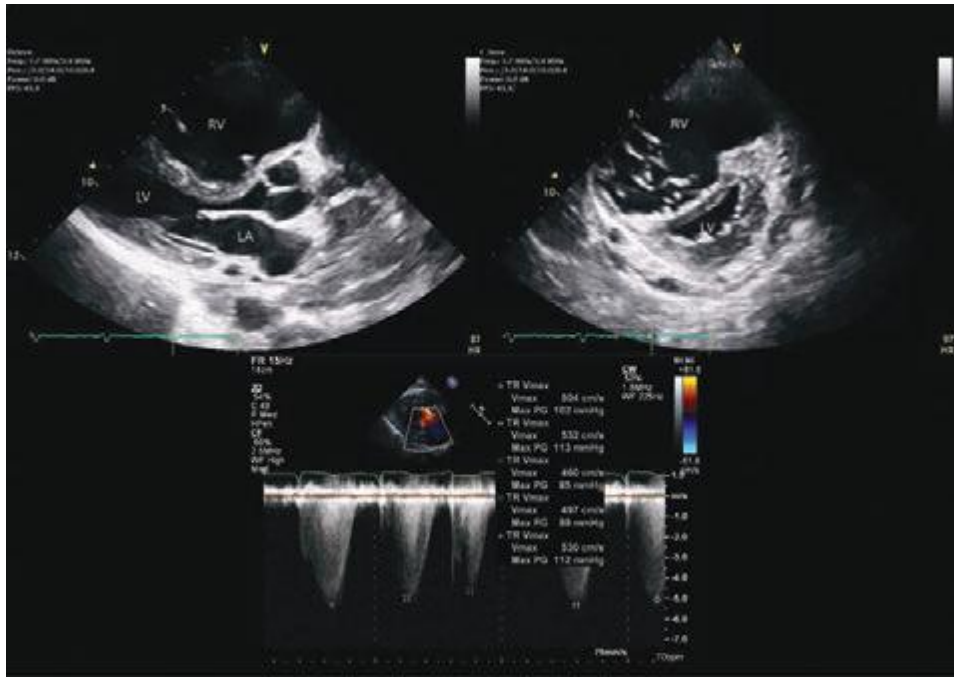


FIGURE 271e-20 A 48-year-old woman with severe idiopathic pulmonary hypertension. Echocardiography reveals evidence of marked right ventricular volume and pressure overload as evidenced by enlarged right ventricle (*upper left and right*), small left ventricle (*upper left and upper right*), and flattening of the interventricular septum (D-shaped septum) in systole and diastole (*upper right*). Tricuspid regurgitation velocity, which reflects the pressure gradient between the right ventricle and the left ventricle, is markedly elevated at 5 m/s, consistent with a right ventricle to right atrial pressure gradient of 100 mmHg, which is consistent with systemic right-sided pressures. LA, left atrium; LV, left ventricle; RV, right ventricle. (See Videos 271e-18 and 271e-19.)

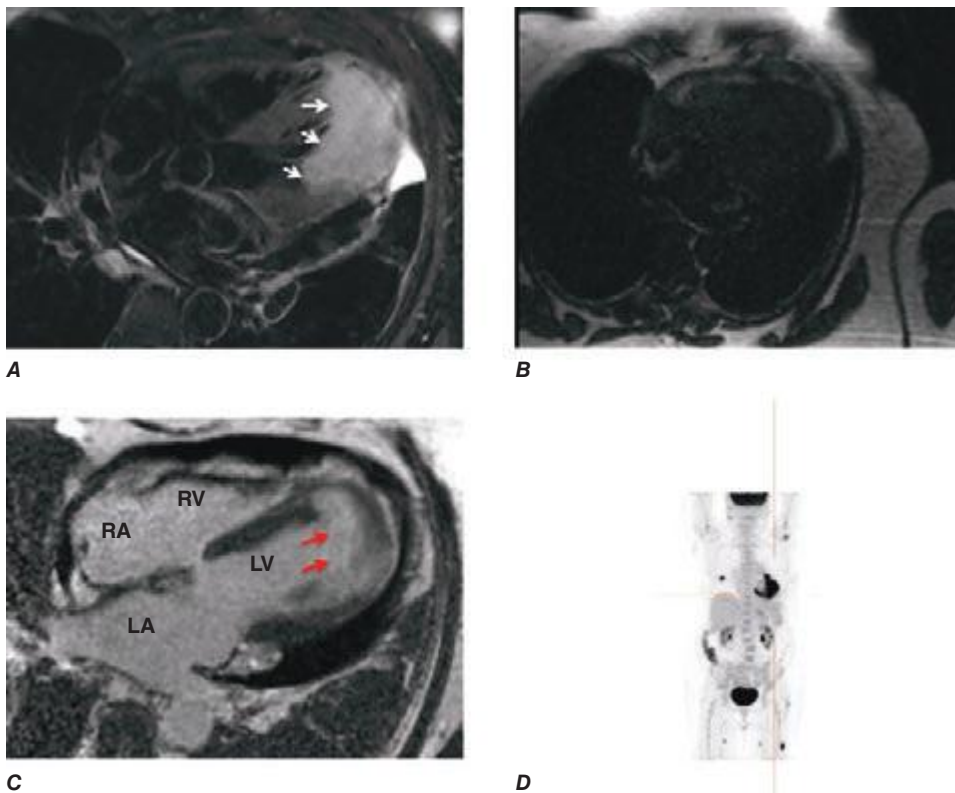


FIGURE 271e-21 Metastatic cardiac tumor diagnosed by cardiac magnetic resonance (MR) in a patient who presented with chest pain and inferior ST elevation. Left heart catheterization was normal. Cardiac MR demonstrates extensive myocardial edema (**A**, white arrows) with marked reduction in first perfusion (**B**) and accumulation of gadolinium within the cardiac mass 10–15 min after injection of gadolinium (**C**, red arrows). **D**. Positron emission tomography scan showed increased fluorodeoxyglucose uptake in a lung mass as well as in the cardiac mass, consistent with cardiac metastasis. Biopsy of the lung mass revealed adenocarcinoma of the lung. LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle. (See Videos 271e-20 and 271e-21.)

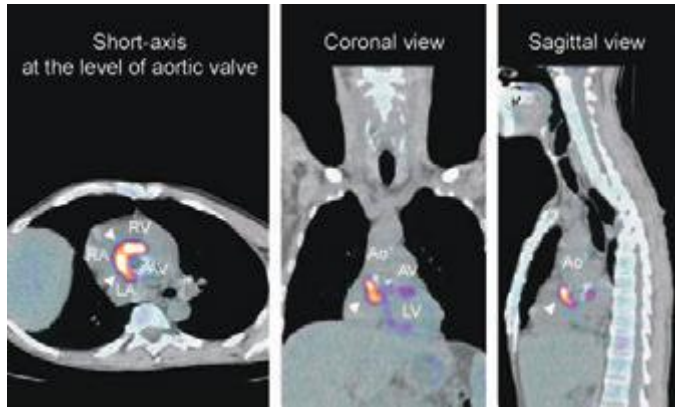


FIGURE 271e-22 Fluorodeoxyglucose (FDG) positron emission tomography (PET)/computed tomography (CT) in a 52-year-old male with a prior aortic valve replacement who presented with fever and was found to have *Haemophilus parainfluenzae* bacteremia. The multiplanar reformatted fused PET/CT images demonstrate intense FDG uptake surrounding the aortic valve prosthesis (arrowheads), compatible with a paravalvular abscess. The patient was found to have purulent fluid around the valve during surgery, and he underwent an aortic valve replacement. Ao, aorta; AV, aortic valve; LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle.

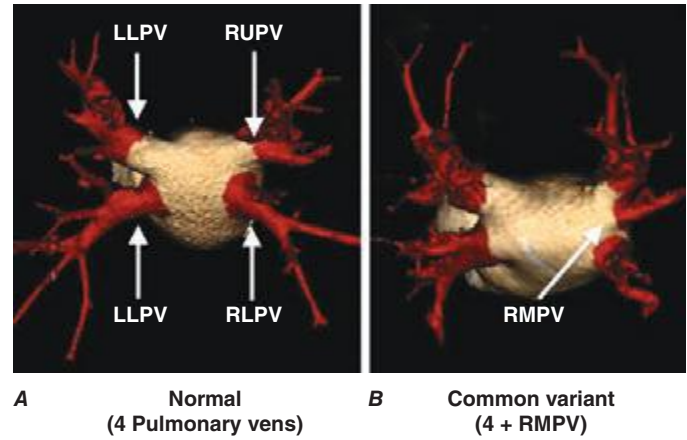


FIGURE 271e-23 Cardiac computed tomography (CT) pulmonary vein mapping in a 62-year-old male with symptomatic, paroxysmal atrial fibrillation referred for cardiac CT for pulmonary vein mapping prior to planned pulmonary vein isolation. Three-dimensional image reconstruction, demonstrating (A) normal pulmonary vein anatomy and (B) common variant with presence of separate right middle pulmonary vein (RMPV) ostium. LLPV, left lower pulmonary vein; LUPV, left upper pulmonary vein; RLPV, right lower pulmonary vein; RUPV, right upper pulmonary vein.

272 Diagnostic Cardiac Catheterization and Coronary Angiography

Jane A. Leopold, David P. Faxon

Diagnostic cardiac catheterization and coronary angiography are considered the gold standard in the assessment of the anatomy and physiology of the heart and its associated vasculature. In 1929, Forssmann demonstrated the feasibility of cardiac catheterization in humans when he passed a urological catheter from a vein in his arm to his right atrium and documented the catheter's position in the heart by x-ray. In the 1940s, Cournand and Richards applied this technique to patients with cardiovascular disease to evaluate cardiac function. These three physicians were awarded the Nobel Prize in 1956. In 1958, Sones inadvertently performed the first selective coronary angiography when a catheter in the left ventricle slipped back across the aortic valve, engaged the right coronary artery, and power-injected 40 mL of contrast down the vessel. The resulting angiogram provided superb anatomic detail of the artery, and the patient suffered no adverse effects. Sones went on to develop selective coronary catheters, which were modified further by Judkins, who developed preformed catheters and allowed coronary artery angiography to gain widespread use as a diagnostic tool. In the United States, cardiac catheterization is the second most common operative procedure, with more than one million procedures performed annually.

CARDIAC CATHETERIZATION

INDICATIONS, RISKS, AND PREPROCEDURE MANAGEMENT

Cardiac catheterization and coronary angiography are indicated to evaluate the extent and severity of cardiac disease in symptomatic patients and to determine if medical, surgical, or catheter-based interventions are warranted (Table 272-1). They are also used to exclude severe disease in symptomatic patients with equivocal findings on noninvasive studies and in patients with chest-pain syndromes of unclear etiology for whom a definitive diagnosis is necessary for management. Cardiac catheterization is not mandatory prior to cardiac surgery in some younger patients who have congenital or valvular heart disease that is well defined by noninvasive imaging and who do not have symptoms or risk factors that suggest concomitant coronary artery disease.

The risks associated with elective cardiac catheterization are relatively low, with a reported risk of 0.05% for myocardial infarction, 0.07% for stroke, and 0.08–0.14% for death. These risks increase substantially if the catheterization is performed emergently, during acute myocardial infarction or in hemodynamically unstable patients.

TABLE 272-1 INDICATIONS FOR CARDIAC CATHETERIZATION AND CORONARY ANGIOGRAPHY

Coronary Artery Disease

Asymptomatic or Symptomatic

High risk for adverse outcome based on noninvasive testing
Sudden cardiac death
Sustained (>30 s) monomorphic ventricular tachycardia
Nonsustained (<30 s) polymorphic ventricular tachycardia

Symptomatic

Canadian Cardiology Society Class II, III, or IV stable angina on medical therapy
Acute coronary syndrome (unstable angina and non-ST-segment elevation myocardial infarction)
Chest-pain syndrome of unclear etiology and equivocal findings on noninvasive tests

ST-Segment Elevation Acute Myocardial Infarction

Reperfusion with primary percutaneous coronary intervention
Persistent or recurrent ischemia
Pulmonary edema and/or reduced ejection fraction
Cardiogenic shock or hemodynamic instability
Risk stratification or positive stress test after acute myocardial infarction
Mechanical complications—mitral regurgitation, ventricular septal defect

Valvular Heart Disease

Suspected severe valve disease in symptomatic patients—dyspnea, angina, heart failure, syncope
Infective endocarditis with need for cardiac surgery
Asymptomatic patients with aortic regurgitation and cardiac enlargement or ↓ ejection fraction
Prior to cardiac surgery in patients with suspected coronary artery disease

Congestive Heart Failure

New onset with angina or suspected undiagnosed coronary artery disease
New-onset cardiomyopathy of uncertain cause or suspected to be due to coronary artery disease

Congenital Heart Disease

Prior to surgical correction, when symptoms or noninvasive testing suggests coronary disease
Suspicion for congenital coronary anomalies

Pericardial Disease

Symptomatic patients with suspected cardiac tamponade or constrictive pericarditis

Cardiac Transplantation

Preoperative and postsurgical evaluation

Other Conditions

Hypertrophic cardiomyopathy with angina
Diseases of the aorta when knowledge of coronary artery involvement is necessary for management

Additional risks of the procedure include tachy- or bradyarrhythmias that require countershock or pharmacologic therapy, acute renal failure leading to transient or permanent dialysis, vascular complications that necessitate surgical repair, and significant access-site bleeding. Of these risks, vascular access-site bleeding is the most common complication, occurring in 1.5–2.0% of patients, with major bleeding events associated with a worse short- and long-term outcome.

In patients who understand and accept the risks associated with cardiac catheterization, there are no absolute contraindications when the procedure is performed in anticipation of a life-saving intervention. Relative contraindications do, however, exist; these include decompensated congestive heart failure; acute renal failure; severe chronic renal insufficiency, unless dialysis is planned; bacteremia; acute stroke; active gastrointestinal bleeding; severe, uncorrected electrolyte abnormalities; a history of an anaphylactic/anaphylactoid reaction to iodinated contrast agents; and a history of allergy/bronchospasm to aspirin in patients for whom progression to a percutaneous coronary intervention is likely and aspirin desensitization has not been performed.

Contrast allergy and contrast-induced acute kidney injury merit further consideration, because these adverse events may occur in otherwise healthy individuals and prophylactic measures exist to reduce risk. Allergic reactions to contrast agents occur in <5% of cases, with severe anaphylactoid (clinically indistinguishable from anaphylaxis, but not mediated by an IgE mechanism) reactions occurring in 0.1%–0.2% of patients. Mild reactions manifest as nausea, vomiting, and urticaria, while severe anaphylactoid reactions lead to hypotensive shock, pulmonary edema, and cardiorespiratory arrest. Patients with a history of significant contrast allergy should be premedicated with corticosteroids and antihistamines (H_1 - and H_2 -blockers) and studies performed with nonionic, low-osmolar contrast agents that have a lower reported rate of allergic reactions.

Contrast-induced acute kidney injury, defined as an increase in creatinine >0.5 mg/dL or 25% above baseline that occurs 48–72 hours after contrast administration, occurs in ~2–7% of patients with rates of 20–30% reported in high-risk patients, including those with diabetes mellitus, congestive heart failure, chronic kidney disease, anemia, and older age. Dialysis is required in 0.3–0.7% of patients and is associated with a fivefold increase in in-hospital mortality. For all patients, adequate intravascular volume expansion with intravenous 0.9% saline (1.0–1.5 mL/kg per hour) for 3–12 hours before and continued 6–24 hours after the procedure limits the risk of contrast-induced acute kidney injury. Pretreatment with *N*-acetylcysteine (Mucomyst) has not reduced the risk of contrast-induced acute kidney injury consistently and, therefore, is no longer recommended routinely. Diabetic patients treated with metformin should stop the drug 48 hours prior to the procedure to limit the associated risk of lactic acidosis. Other strategies to decrease risk include the administration of sodium bicarbonate (3 mL/kg per hour) 1 hour before and 6 hours after the procedure; use of low- or iso-osmolar contrast agents; and limiting the volume of contrast to <100 mL per procedure.

Cardiac catheterization is performed after the patient has fasted for 6 hours and has received intravenous conscious sedation to remain awake but sedated during the procedure. All patients with suspected coronary artery disease are pretreated with 325 mg aspirin. In patients in whom the procedure is likely to progress to a percutaneous coronary intervention, an additional antiplatelet agent should be started: clopidogrel (600-mg loading dose and 75 mg daily) or prasugrel (60-mg loading dose and 10 mg daily), or ticagrelor (180-mg loading and 90 mg twice daily). Prasugrel should not be selected for individuals with prior stroke or transient ischemic attack. Warfarin is held starting 2–3 days prior to the catheterization to allow the international normalized ratio (INR) to fall to <1.7 and limit access-site bleeding complications. Cardiac catheterization is a sterile procedure, so antibiotic prophylaxis is not required.

TECHNIQUE

Cardiac catheterization and coronary angiography provide a detailed hemodynamic and anatomic assessment of the heart and coronary arteries. The selection of procedures is dependent on the patient's symptoms and clinical condition, with some direction provided by noninvasive studies.

Vascular Access Cardiac catheterization procedures are performed using a percutaneous technique to enter the femoral artery and vein as the preferred access sites for left and right heart catheterization, respectively. A flexible sheath is inserted into the vessel over a guidewire, allowing diagnostic catheters to be introduced into the vessel and advanced toward the heart using fluoroscopic guidance. The radial artery (or brachial artery) may also be used as an arterial access site in patients, particularly those with peripheral arterial disease that involves the abdominal aorta, iliac, or femoral vessels; severe iliac artery tortuosity; morbid obesity; or preference for early postprocedure ambulation. Use of radial-artery access is gaining popularity due to a lower rate of access-site bleeding complications. A normal Allen's test confirming dual blood supply to the hand from the radial and ulnar arteries is recommended prior to access at this site. The internal jugular or antecubital veins serve as alternate access sites to the

right heart when the patient has an inferior vena cava filter in place or requires prolonged hemodynamic monitoring.

Right Heart Catheterization This procedure measures pressures in the right heart. Right heart catheterization is no longer a routine part of diagnostic cardiac catheterization, but it is reasonable in patients with unexplained dyspnea, valvular heart disease, pericardial disease, right and/or left ventricular dysfunction, congenital heart disease, and suspected intracardiac shunts. Right heart catheterization uses a balloon-tipped flotation catheter that is advanced sequentially to the right atrium, right ventricle, pulmonary artery, and pulmonary wedge position (as a surrogate for left atrial pressure) using fluoroscopic guidance; in each cardiac chamber, pressure is measured and blood samples are obtained for oxygen saturation analysis to screen for intracardiac shunts.

Left Heart Catheterization This procedure measures pressures in the left heart as a determinant of left ventricular performance. With the aid of fluoroscopy, a catheter is guided to the ascending aorta and across the aortic valve into the left ventricle to provide a direct measure of left ventricular pressure. In patients with a tilting-disc prosthetic aortic valve, crossing the valve with a catheter is contraindicated, and the left heart may be accessed via a transseptal technique from the right atrium using a needle-tipped catheter to puncture the atrial septum at the fossa ovalis. Once the catheter crosses from the right to the left atrium, it can be advanced across the mitral valve to the left ventricle. This technique is also used for mitral valvuloplasty. Heparin is given for prolonged procedures to limit the risk of stroke from embolism of clots that may form on the catheter. For patients with heparin-induced thrombocytopenia, the direct thrombin inhibitors bivalirudin (0.75 mg/kg bolus, 1.75 mg/kg per hour for the duration of the procedure) or argatroban (350 µg/kg bolus, 15 µg/kg per minute for the duration of the procedure) may be used.

HEMODYNAMICS

A comprehensive hemodynamic assessment involves obtaining pressure measurements in the right and left heart and peripheral arterial system and determining the cardiac output ([Table 272-2](#)). The shape

TABLE 272-2 NORMAL VALUES FOR HEMODYNAMIC MEASUREMENTS

Pressures (mmHg)	
Right atrium	
Mean	0–5
<i>a</i> wave	1–7
<i>v</i> wave	1–7
Right ventricle	
Peak systolic/end diastolic	17–32/1–7
Pulmonary artery	
Peak systolic/end diastolic	17–32/1–7
Mean	9–19
Pulmonary capillary wedge (mean)	4–12
Left atrium	
Mean	4–12
<i>a</i> wave	4–15
<i>v</i> wave	4–15
Left ventricle	
Peak systolic/end diastolic	90–130/5–12
Aorta	
Peak systolic/end diastolic	90–130/60–85
Mean	70–100
(Resistances [dyn-s]/cm ⁵)	
Systemic vascular resistance	900–1400
Pulmonary vascular resistance	40–120
Oxygen Consumption Index ([L-min]/m ²)	
Arteriovenous oxygen difference (vol %)	3.5–4.8
Cardiac index ([L-min]/m ²)	2.8–4.2

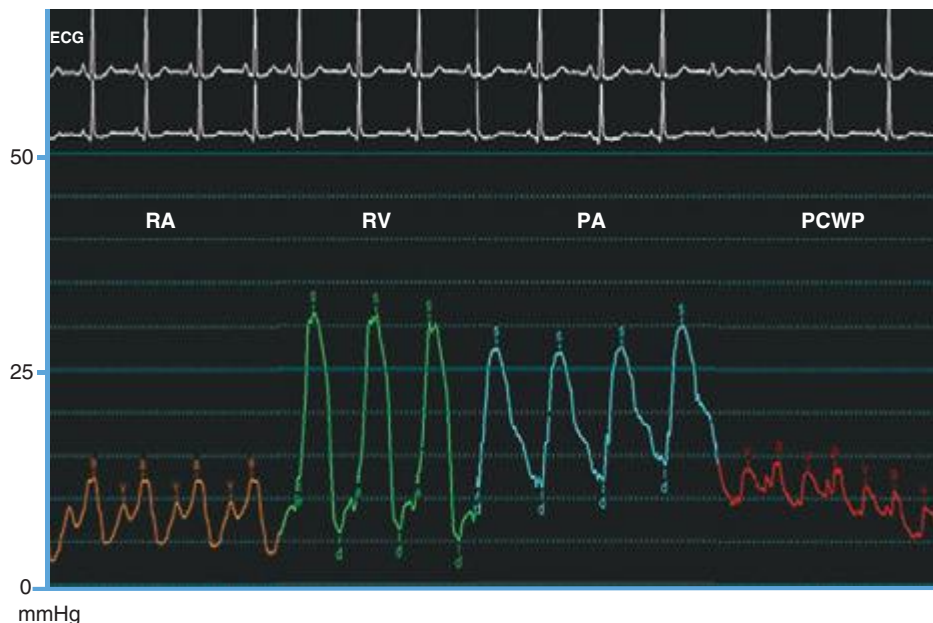


FIGURE 272-1 Normal hemodynamic waveforms recorded during right heart catheterization. Atrial pressure tracings have a characteristic “a” wave that reflects atrial contraction and a “v” wave that reflects pressure changes in the atrium during ventricular systole. Ventricular pressure tracings have a low-pressure diastolic filling period and a sharp rise in pressure that occurs during ventricular systole. d, diastole; PA, pulmonary artery; PCWP, pulmonary capillary wedge pressure; RA, right atrium; RV, right ventricle; s, systole.

and magnitude of the pressure waveforms provide important diagnostic information; an example of normal pressure tracings is shown in Fig. 272-1. In the absence of valvular heart disease, the atria and ventricles are “one chamber” during diastole when the tricuspid and mitral valves are open while in systole, when the pulmonary and aortic valves are open, the ventricles and their respective outflow tracts are considered “one chamber.” These concepts form the basis by which hemodynamic measurements are used to assess valvular stenosis. When aortic stenosis is present, there is a systolic pressure gradient between the left ventricle and the aorta; when mitral stenosis is present, there is a diastolic pressure gradient between the pulmonary capillary wedge (left atrial) pressure and the left ventricle (Fig. 272-2). Hemodynamic measurements also discriminate between aortic stenosis and hypertrophic obstructive cardiomyopathy where the asymmetrically hypertrophied septum creates a dynamic intraventricular pressure gradient during ventricular systole. The magnitude of this obstruction is measured using an end-hole catheter positioned at the left ventricular apex that is pulled back while recording pressure; once the catheter has passed the septal obstruction and is positioned in the apex of the left ventricle, a gradient can be measured between the left ventricular apex and the aorta. Hypertrophic obstructive cardiomyopathy is confirmed by the Brockenbrough-Braunwald sign: following a premature ventricular contraction, there is an increase in the left ventricular–aorta pressure gradient with a simultaneous decrease in the aortic pulse pressure. These findings are absent in aortic stenosis.

Regurgitant valvular lesions increase volume (and pressure) in the “receiving” cardiac chamber. In severe mitral and tricuspid regurgitation, the increase in blood flow to the atria takes place during ventricular systole, leading to an increase in the *v* wave (two times greater than the mean pressure). Severe aortic regurgitation leads to a decrease in aortic diastolic pressure with a concomitant rise in left ventricular end-diastolic pressure, resulting in equalization of pressures between the two chambers at end-diastole.

Hemodynamic measurements are also used to differentiate between cardiac tamponade, constrictive pericarditis, and restrictive cardiomyopathy (Table 272-3). In cardiac tamponade, right atrial pressure is increased with a decreased or absent “y” descent, indicative of impaired right atrial emptying in diastole, and there is diastolic equalization of pressures in all cardiac chambers. In constrictive pericarditis, right atrial pressure is elevated with a prominent “y” descent, indicating rapid filling of the right ventricle during early diastole. A diastolic dip and plateau or “square root sign,” in the ventricular waveforms due to an abrupt halt in ventricular filling during diastole; right ventricular and pulmonary artery pressures are elevated; and discordant pressure changes in the right and left ventricles with inspiration (right ventricular systolic pressure increases while left ventricular systolic pressure decreases) are observed. The latter hemodynamic phenomenon is the most specific for constriction. Restrictive cardiomyopathy may be distinguished from constrictive pericarditis by a marked increase in right ventricular and pulmonary artery systolic pressures (usually >60 mmHg), a separation of the left and right ventricular diastolic pressures by

>5 mmHg (at baseline or with acute volume loading), and concordant changes in left and right ventricular diastolic filling pressures with inspiration (both increase).

Cardiac Output Cardiac output is measured by the Fick method or the thermodilution technique. Typically, the Fick method and thermodilution technique are both performed during cardiac catheterization, although the Fick method is considered more reliable in the presence of tricuspid regurgitation and in low-output states. The Fick method

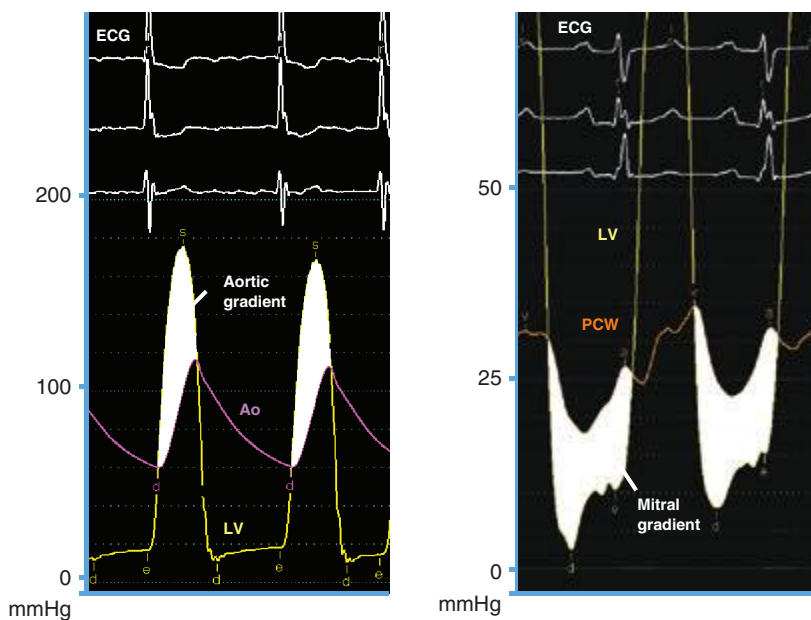


FIGURE 272-2 Severe aortic and mitral stenosis. Simultaneous recording of left ventricular (LV) and aortic (Ao) pressure tracings demonstrates a 62-mmHg mean systolic gradient (shaded area) that corresponds to an aortic valve area of 0.6 cm² (left). Simultaneous recording of LV and pulmonary capillary wedge (PCW) pressure tracings reveals a 14-mmHg mean diastolic gradient (shaded area) that is consistent with critical mitral stenosis (mitral valve area = 0.5 cm²). d, diastole; e, end diastole; s, systole.

TABLE 272-3 HEMODYNAMIC FINDINGS IN TAMPONADE, CONSTRICTIVE PERICARDITIS, AND RESTRICTIVE CARDIOMYOPATHY

	Cardiac Tamponade	Constrictive Pericarditis	Effusive-Constrictive Pericarditis	Restrictive Cardiomyopathy
Pericardial pressure	↑	↑	↑	Normal
Right atrium pressure	↑	↑	↑ (Fails to decrease by 50% or to <10 mmHg after pericardiocentesis)	↑
Right atrium pressure waveform	Prominent “x” descent Diminished or absent “y” descent	Prominent “x” descent Prominent “y” descent	Prominent “x” descent “y” descent less prominent than expected	Prominent “y” descent
Right ventricle systolic pressure	<50 mmHg	<50 mmHg	<50 mmHg	>60 mmHg
Right ventricle end-diastolic pressure	Equals left ventricular end-diastolic pressure within 5 mmHg	>1/3 right ventricular systolic pressure Equals left ventricular end-diastolic pressure within 5 mmHg	>1/3 right ventricular systolic pressure Equals left ventricular end-diastolic pressure within 5 mmHg	<1/3 right ventricular systolic pressure Less than left ventricular end-diastolic pressure by ≥5 mmHg
Right ventricle pressure waveform		Dip and plateau or “square root” sign	Dip and plateau or “square root” sign	Dip and plateau or “square root” sign
Right ventricle–left ventricle systolic pressure relationship with inspiration	Discordant	Discordant	Discordant	Concordant

uses oxygen as the indicator substance and is based on the principle that the amount of a substance taken up or released by an organ (oxygen consumption) is equal to the product of its blood flow (cardiac output) and the difference in the concentration of the substance in the arterial and venous circulation (arterial-venous oxygen difference). Thus, the formula for calculating the Fick cardiac output is:

$$\text{Cardiac output (L/min)} = \frac{\text{oxygen consumption [mL/min]}}{\text{arterial-venous oxygen difference [mL/L]}}$$

Oxygen consumption is estimated as 125 mL oxygen/minute \times body surface area, and the arterial-venous oxygen difference is determined by first calculating the oxygen carrying capacity of blood (hemoglobin [g/100 mL] \times 1.36 [mL oxygen/g hemoglobin] \times 10) and multiplying this product by the fractional oxygen saturation. The thermodilution method measures a substance that is injected into and adequately mixes with blood. In contemporary practice, thermodilution cardiac outputs are measured using temperature as the indicator. Measurements are made with a thermistor-tipped catheter that detects temperature deviations in the pulmonary artery after the injection of 10 mL of room-temperature normal saline into the right atrium.

Vascular Resistance Resistance across the systemic and pulmonary circulations is calculated by extrapolating from Ohm’s law of electrical resistance and is equal to the mean pressure gradient divided by the mean flow (cardiac output). Therefore, systemic vascular resistance is ([mean aortic pressure – mean right atrial pressure]/cardiac output) multiplied by 80 to convert the resistance from Wood units to dyns-cm⁻⁵. Similarly, the pulmonary vascular resistance is ([mean pulmonary artery – mean pulmonary capillary wedge pressure]/cardiac output) \times 80. Pulmonary vascular resistance is lowered by oxygen, nitroprusside, calcium channel blockers, prostacyclin infusions, and inhaled nitric oxide; these therapies may be administered during catheterization to determine if increased pulmonary vascular resistance is fixed or reversible.

Valve Area Hemodynamic data may also be used to calculate the valve area using the Gorlin formula that equates the area to the flow across the valve divided by the pressure gradient between the cardiac chambers surrounding the valve. The formula for the assessment of valve area is: Area = (cardiac output [cm³/min])/[systolic ejection period or diastolic filling period][heart rate])/44.3 C \times square root of the pressure gradient, where C = 1 for aortic valve and 0.85 for the mitral valve. A valve area of <1.0 cm² and a mean gradient of greater than 40 mmHg indicate severe aortic stenosis, while a valve area of <1.5 cm² and a mean gradient >5–10 mmHg is consistent with

moderate-to-severe mitral stenosis; in symptomatic patients with a mitral valve area >1.5 cm², a mean gradient >15 mmHg, pulmonary artery pressure >60 mmHg, or a pulmonary artery wedge pressure >25 mmHg after exercise is also considered significant and may warrant intervention. The modified Hakki formula has also been used to estimate aortic valve area. This formula calculates the valve area as the cardiac output (L/min) divided by the square root of the pressure gradient. Aortic valve area calculations based on the Gorlin formula are flow-dependent and, therefore, for patients with low cardiac outputs, it is imperative to determine if a decreased valve area actually reflects a fixed stenosis or is overestimated by a low cardiac output and stroke volume that is insufficient to open the valve leaflets fully. In these instances, cautious hemodynamic manipulation using dobutamine to increase the cardiac output and recalculation of the aortic valve area may be necessary.

Intracardiac Shunts In patients with congenital heart disease, detection, localization, and quantification of the intracardiac shunt should be evaluated. A shunt should be suspected when there is unexplained arterial desaturation or increased oxygen saturation of venous blood. A “step up” or increase in oxygen content indicates the presence of a left-to-right shunt while a “step down” indicates a right-to-left shunt. The shunt is localized by detecting a difference in oxygen saturation levels of 5–7% between adjacent cardiac chambers. The severity of the shunt is determined by the ratio of pulmonary blood flow (Q_p) to the systemic blood flow (Q_s), or $Q_p/Q_s = ([\text{systemic arterial oxygen content} - \text{mixed venous oxygen content}]/\text{pulmonary vein oxygen content} - \text{pulmonary artery oxygen content})$. For an atrial septal defect, a shunt ratio of 1.5 is considered significant and factored with other clinical variables to determine the need for intervention. When a congenital ventricular septal defect is present, a shunt ratio of ≥ 2.0 with evidence of left ventricular volume overload is a strong indication for surgical correction.

VENTRICULOGRAPHY AND AORTOGRAPHY

Ventriculography to assess left ventricular function may be performed during cardiac catheterization. A pigtail catheter is advanced retrograde across the aortic valve into the left ventricle and 30–45 mL of contrast is power-injected to visualize the left ventricular chamber during the cardiac cycle. The ventriculogram is usually performed in the right anterior oblique projection to examine wall motion and mitral valve function. Normal wall motion is observed as symmetric contraction of all segments; hypokinetic segments have decreased contraction, akinetic segments do not contract, and dyskinetic segments appear to bulge paradoxically during systole (Fig. 272-3). Ventriculography may also reveal a left ventricular aneurysm, pseudoaneurysm, or diverticulum and can be used to assess mitral valve prolapse and the severity of mitral regurgitation. The degree of mitral regurgitation is estimated

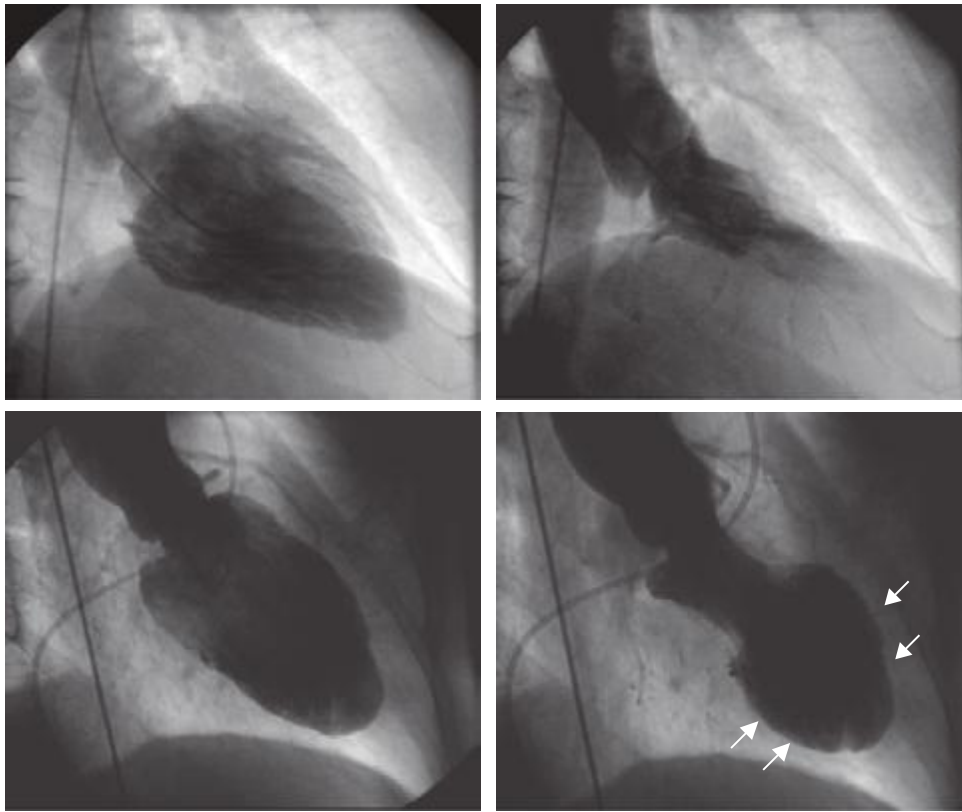


FIGURE 272-3 Left ventriculogram at end diastole (*left*) and end systole (*right*). In patients with normal left ventricular function, the ventriculogram reveals symmetric contraction of all walls (*top*). Patients with coronary artery disease may have wall motion abnormalities on ventriculography as seen in this 60-year-old male following a large anterior myocardial infarction. In systole, the anterior, apical, and inferior walls are akinetic (*white arrows*) (*bottom*).

by comparing the density of contrast opacification of the left atrium with that of the left ventricle. Minimal contrast reflux into the left atrium is considered 1+ mitral regurgitation, while contrast density in the left atrium that is greater than that in the left ventricle with reflux of contrast into the pulmonary veins within three beats defines 4+ mitral regurgitation. Ventriculography performed in the left anterior oblique projection can be used to identify a ventricular septal defect. Calculation of the ventricular volumes in systole and diastole allows calculation of stroke volume and cardiac output.

Aortography in the cardiac catheterization laboratory visualizes abnormalities of the ascending aorta, including aneurysmal dilation and involvement of the great vessels, as well as dissection with compression of the true lumen by an intimal flap that separates the true and false

arteries arising from the left main coronary artery (**Fig. 272-4**). When the right coronary artery is the origin of the atrioventricular nodal branch, the posterior descending artery, and the posterior lateral vessels, the circulation is defined as right dominant; this is found in ~85% of individuals. When these branches arise from the left circumflex artery as occurs in ~5% of individuals, the circulation is defined as left dominant. The remaining ~10% of patients have a codominant circulation with vessels arising from both the right and left coronary circulation. In some patients, a ramus intermedius branch arises directly from the left main coronary artery; this finding is a normal variant. Coronary artery anomalies occur in 1–2% of patients, with separate ostia for the left anterior descending and left circumflex arteries being the most common (0.41%).

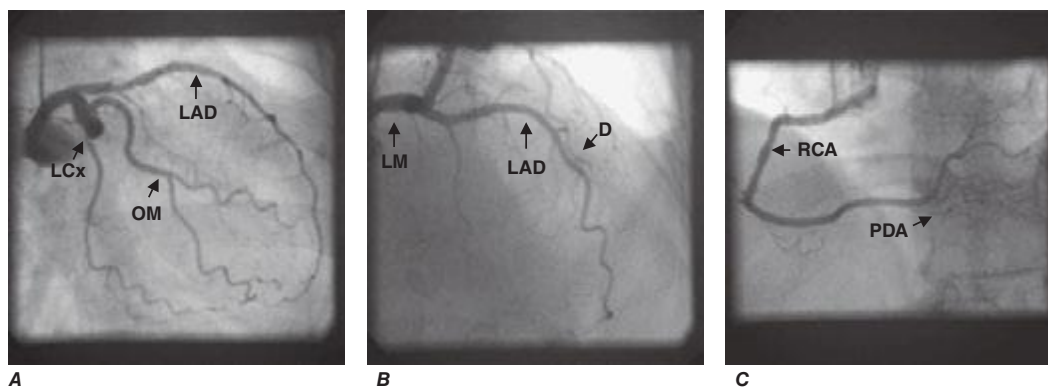


FIGURE 272-4 Normal coronary artery anatomy. **A.** Coronary angiogram showing the left circumflex (LCx) artery and its obtuse marginal (OM) branches. The left anterior descending artery (LAD) is also seen but may be foreshortened in this view. **B.** The LAD and its diagonal (D) branches are best seen in cranial views. In this angiogram, the left main (LM) coronary artery is also seen. **C.** The right coronary artery (RCA) gives off the posterior descending artery (PDA), so this is a right dominant circulation.

lumina. Aortography can also be used to identify patent saphenous vein grafts that elude selective cannulation, identify shunts that involve the aorta such as a patent ductus arteriosus, and provide a qualitative assessment of aortic regurgitation using a 1+–4+ scale similar to that used for mitral regurgitation.

CORONARY ANGIOGRAPHY

Selective coronary angiography is almost always performed during cardiac catheterization and is used to define the coronary anatomy and determine the extent of epicardial coronary artery and coronary artery bypass graft disease. Specially shaped coronary catheters are used to engage the left and right coronary ostia. Hand injection of radiopaque contrast agents creates a coronary “luminogram” that is recorded as radiographic images (cine angiography). Because the coronary arteries are three-dimensional objects that are in motion with the cardiac cycle, angiograms of the vessels using several different orthogonal projections are taken to best visualize the vessels without overlap or foreshortening.

The normal coronary anatomy is highly variable between individuals, but, in general, there are two coronary ostia and three major coronary vessels—the left anterior descending, the left circumflex, and the right coronary arteries with the left anterior descending and left circumflex

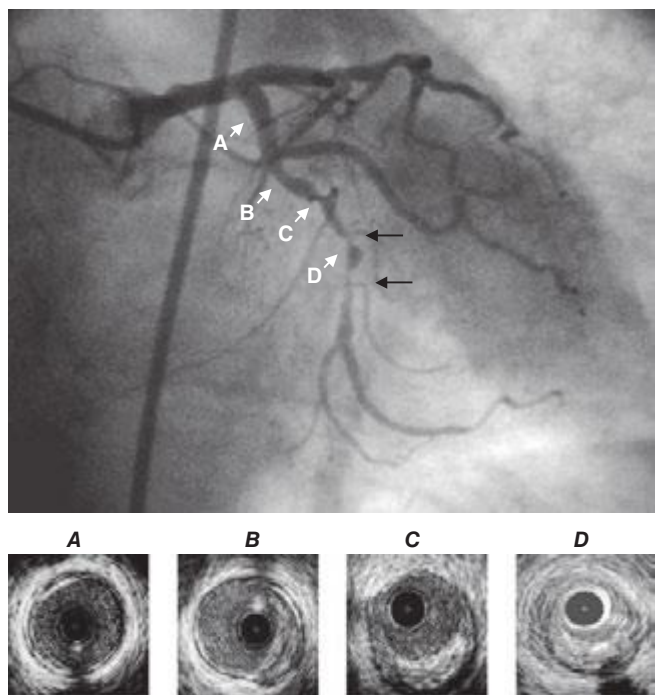


FIGURE 272-5 Coronary stenoses on cine angiogram and intravascular ultrasound. Significant stenoses in the coronary artery are seen as narrowings (*black arrows*) of the vessel. Intravascular ultrasound shows a normal segment of artery (**A**), areas with eccentric plaque (**B, C**), and near total obliteration of the lumen at the site of the significant stenosis (**D**). Note that the intravascular ultrasound catheter is present in the images as a black circle.

Coronary angiography visualizes coronary artery stenoses as luminal narrowings on the cine angiogram. The degree of narrowing is referred to as the percent stenosis and is determined visually by comparing the most severely diseased segment with a proximal or distal “normal segment”; a stenosis $>50\%$ is considered significant (**Fig. 272-5**). Online quantitative coronary angiography can provide a more accurate assessment of the percent stenosis and lessen the tendency to overestimate lesion severity visually. The presence of a myocardial bridge, which most commonly involves the left anterior descending artery, may be mistaken for a significant stenosis; this occurs when a portion of the vessel dips below the epicardial surface into the myocardium and is subject to compressive forces during ventricular systole. The key to differentiating a myocardial bridge from a fixed stenosis is that the “stenosed” part of the vessel returns to normal during diastole. Coronary calcification is also seen during angiography prior to the injection of contrast agents. Collateral blood vessels may be seen traversing from one vessel to the distal vasculature of a severely stenosed or totally occluded vessel. Thrombolysis in myocardial infarction (TIMI) flow grade, a measure of the relative duration of time that it takes for contrast to opacify the coronary artery fully, may provide an additional clue to the degree of lesion severity, and the presence of TIMI grade 1 (minimal filling) or 2 (delayed filling) suggests that a significant coronary artery stenosis is present.

INTRAVASCULAR ULTRASOUND, OPTICAL COHERENCE TOMOGRAPHY, AND FRACTIONAL FLOW RESERVE

During coronary angiography, intermediate stenoses (40–70%), indeterminate findings, or anatomic findings that are incongruous with the patient’s symptoms may require further interrogation. In these cases, intravascular ultrasound provides a more accurate anatomic assessment of the coronary artery and the degree of coronary atherosclerosis

(**Fig. 272-5**). Intravascular ultrasound (IVUS) is performed using a small flexible catheter with a 40-mHz transducer at its tip that is advanced into the coronary artery over a guidewire. Data from intravascular ultrasound studies may be used to image atherosclerotic plaque precisely, determine luminal cross-sectional area, and measure vessel size; it is also used during or following percutaneous coronary intervention to assess the stenosis and determine the adequacy of stent placement. Optical coherence tomography (OCT) is a catheter-based imaging technique that uses near-infrared light to generate images with better spatial resolution than IVUS; however, the depth of field is smaller. The advantage of OCT imaging over IVUS lies in its ability to image characteristics of the atherosclerotic plaque (lipid, fibrous cap) with high definition and to assess coronary stent placement, apposition, and patency (**Fig. 272-6**).

Measurement of the fractional flow reserve provides a functional assessment of the stenosis and is more accurate in predicting long-term clinical outcome than imaging techniques. The fractional flow reserve is the ratio of the pressure in the coronary artery distal to the stenosis divided by the pressure in the artery proximal to the stenosis at maximal vasodilation. Fractional flow reserve is measured using a coronary pressure–sensor guidewire at rest and at maximal hyperemia following the injection of adenosine. A fractional flow reserve of <0.80 indicates a hemodynamically significant stenosis that would benefit from intervention.

POSTPROCEDURE CARE

Once the procedure is completed, vascular access sheaths are removed. If the femoral approach is used, direct manual compression or vascular closure devices that immediately close the arteriotomy site with a staple/clip, collagen plug, or sutures are used to achieve hemostasis. These devices decrease the length of supine bed rest (from 6 hours to 2–4 hours) and improve patient satisfaction but have not been shown definitively to be superior to manual compression with respect to access-site complications. With radial-artery access, bed rest is needed for only 2 hours. When cardiac catheterization is performed as an elective outpatient procedure, the patient completes postprocedure bed rest in a monitored setting and is discharged home with instructions to liberalize fluids because contrast agents promote an osmotic diuresis, to avoid strenuous activity, and to observe the vascular access site for signs of complications. Overnight hospitalization may be required for high-risk patients with significant comorbidities, patients with complications occurring during the catheterization, or patients who have undergone a percutaneous coronary intervention. Hypotension early after the procedure may be due to inadequate fluid replacement or retroperitoneal bleeding from the access site. Patients who received >2 Gy of radiation during the procedure should be examined for signs of erythema. For patients who received higher doses (>5 Gy), clinical follow-up within 1 month to assess for skin injury is recommended.

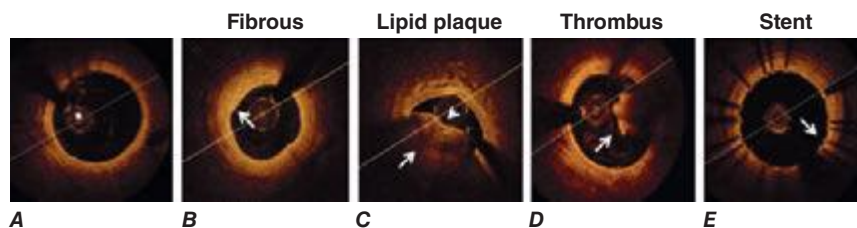


FIGURE 272-6 Optical coherence tomography imaging. **A**. The optical coherence tomography (OCT) catheter (*) in the lumen of a coronary artery with limited neointima formation. The intima is seen with high definition, but unlike intravascular ultrasound imaging, the vessel media and adventitia are not well visualized. **B**. A fibrous plaque (*arrow*) is characterized by a bright signal. **C**. A large, eccentric, lipid-rich plaque obscures part of the vessel lumen. Because lipid in the plaque absorbs light, the lipid-rich plaque appears as a dark area with irregular borders (*arrow*). The plaque is covered by a thin fibrous cap (*arrowhead*) typical of a vulnerable plaque. **D**. A thrombus (*arrow*) adherent to a ruptured plaque that is protruding into the vessel lumen. **E**. A coronary stent is that is well opposed to the vessel wall. The stent struts appear as short bright lines with dropout behind the struts (*arrow*).

SECTION 3 DISORDERS OF RHYTHM

273e Principles of Electrophysiology

David D. Spragg, Gordon F. Tomaselli

HISTORY AND INTRODUCTION

The field of cardiac electrophysiology was ushered in with the development of the electrocardiogram (ECG) by Einthoven at the turn of the twentieth century. Subsequent recording of cellular membrane currents demonstrated that the body surface ECG is the timed sum of the cellular action potentials in the atria and ventricles. In the late 1960s, the development of intracavitary recording, in particular, His bundle electrograms, marked the beginning of contemporary clinical electrophysiology. Adoption of radiofrequency technology to ablate cardiac tissue in the early 1990s heralded the birth of interventional cardiac electrophysiology.

The clinical problem of sudden death caused by ventricular arrhythmias, most commonly in the setting of coronary artery obstruction, was recognized as early as the late nineteenth century. The problem was vexing and led to the development of pharmacologic and non-pharmacologic therapies, including transthoracic defibrillators, cardiac massage, and, most recently, implantable defibrillators. Over time the limitations of antiarrhythmic drug therapy have been highlighted repeatedly in clinical trials, and now ablation and devices are first-line therapy for a number of cardiac arrhythmias.

In the last two decades, the genetic basis of a number of heritable arrhythmias has been elucidated, revealing important insights into the mechanisms not only of these rare arrhythmias but also of similar rhythm disturbances observed in more common forms of heart disease.

DESCRIPTIVE PHYSIOLOGY

The normal cardiac impulse is generated by pacemaker cells in the sinoatrial node situated at the junction of the right atrium and the superior vena cava (see Fig. 268-1). This impulse is transmitted slowly through nodal tissue to the anatomically complex atria, where it is conducted more rapidly to the atrioventricular node (AVN), inscribing the P wave of the ECG (see Fig. 268-2). There is a perceptible delay in conduction through the anatomically and functionally heterogeneous AVN. The time needed for activation of the atria and the AVN delay is represented as the PR interval of the ECG. The AVN is the only electrical connection between the atria and the ventricles in the normal heart. The electrical impulse emerges from the AVN and is transmitted to the His-Purkinje system, specifically the common bundle of His, then the left and right bundle branches, and then to the Purkinje network, facilitating activation of ventricular muscle. In normal circumstances, the ventricles are activated rapidly in a well-defined fashion that is determined by the course of the Purkinje network, and this inscribes the QRS complex (see Fig. 268-2). Recovery of electrical excitability occurs more slowly and is governed by the time of activation and duration of regional action potentials. The relative brevity of epicardial action potentials in the ventricle results in repolarization that occurs first on the epicardial surface and then proceeds to the endocardium, which inscribes a T wave normally of the same polarity as the QRS complex. The duration of ventricular activation and recovery is determined by the action potential duration and is represented on the body surface ECG by the QT interval (see Fig. 268-2).

Cardiac myocytes exhibit a characteristically long action potential (200–400 ms) compared with neurons and skeletal muscle cells (1–5 ms). The action potential profile is sculpted by the orchestrated activity of multiple distinctive time- and voltage-dependent ionic currents (Fig. 273e-1A). The currents are carried by transmembrane

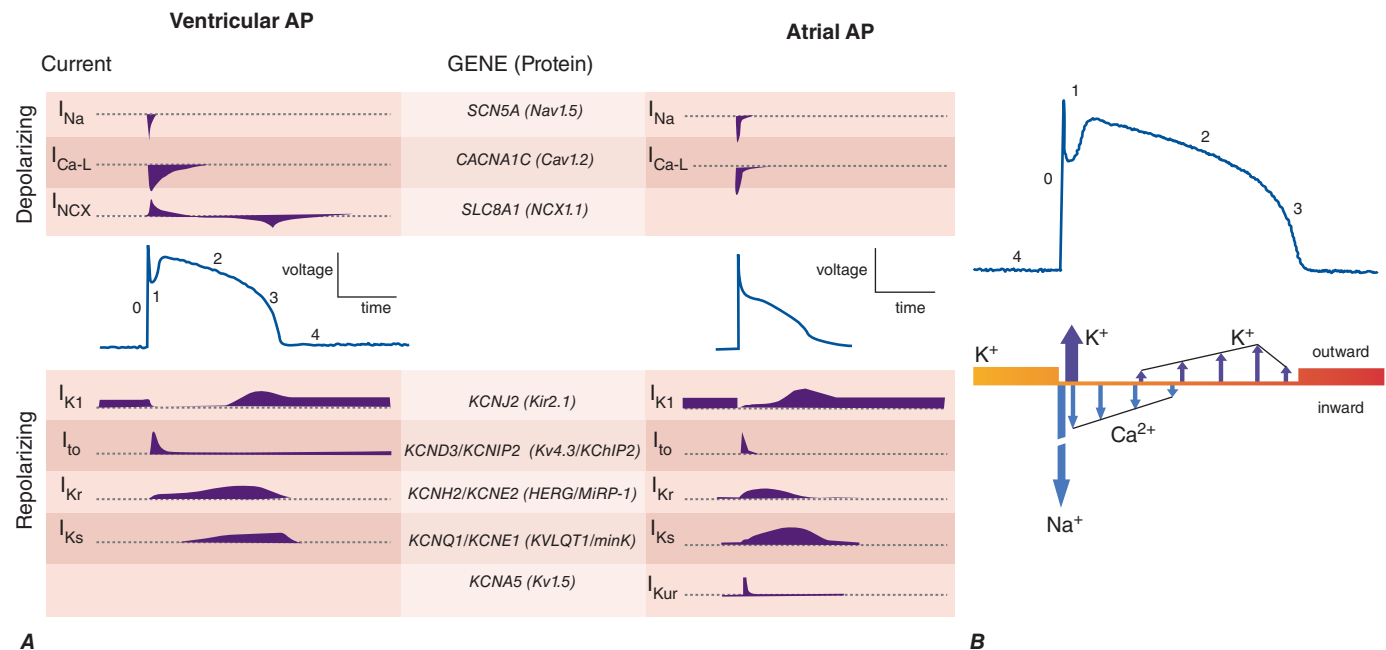


FIGURE 273e-1 **A.** Cellular atrial and ventricular action potentials. Phases 0–4 are the rapid upstroke, early repolarization, plateau, late repolarization, and diastole, respectively. The ionic currents and their respective genes are shown above and below the action potentials. The currents that underlie the action potentials vary in atrial and ventricular myocytes. **B.** A ventricular action potential with a schematic of the ionic currents flowing during the phases of the action potential. Potassium current (I_{K1}) is the principal current during phase 4 and determines the resting membrane potential of the myocyte. Sodium current generates the upstroke of the action potential (phase 0); activation of I_{to} with inactivation of the Na current inscribes early repolarization (phase 1). The plateau (phase 2) is generated by a balance of repolarizing potassium currents and depolarizing calcium current. Inactivation of the calcium current with persistent activation of potassium currents (predominantly I_{Kr} and I_{Ks}) causes phase 3 repolarization.

proteins that passively conduct ions down their electrochemical gradients through selective pores (ion channels), actively transport ions against their electrochemical gradient (pumps, transporters), or electrogenically exchange ionic species (exchangers).

Action potentials in the heart are regionally distinct. The regional variability in cardiac action potentials is a result of differences in the number and types of ion channel proteins expressed by different cell types in the heart. Further, unique sets of ionic currents are active in pacemaking and muscle cells, and the relative contributions of these currents may vary in the same cell type in different regions of the heart (Fig. 273e-1A).

Ion channels are complex, multisubunit transmembrane glycoproteins that open and close in response to a number of biologic stimuli, including a change in membrane voltage, ligand binding (directly to the channel or to a G protein-coupled receptor), and mechanical deformation (Fig. 273e-2). Other ion motive exchangers and transporters contribute importantly to cellular excitability in the heart. Ion pumps establish and maintain the ionic gradients across the cell membrane that serve as the driving force for current flow through ion channels. Transporters or exchangers that do not move ions in an electrically neutral manner (e.g., the sodium-calcium exchanger transports three Na^+ for one Ca^{2+}) are termed *electrogenic* and contribute directly to the action potential profile.

The most abundant superfamily of ion channels expressed in the heart is voltage gated. Several structural themes are common to all voltage-dependent ion channels. First, the architecture is modular, consisting either of four homologous subunits (e.g., K channels) or of four internally homologous domains (e.g., Na and Ca channels). Second, the proteins fold around a central pore lined by amino acids that exhibit exquisite conservation within a given channel family of like selectivity (e.g., all Na channels have very similar P segments). Third, the general strategy for activation gating (opening and closing in response to changes in membrane voltage) is highly conserved: the fourth transmembrane segment (S4), studded with positively charged residues, lies within the membrane field and moves in response to depolarization, opening the channel. Fourth, most ion channel complexes include not only the pore-forming proteins (α subunits) but also auxiliary subunits (e.g., β subunits) that modify channel function (Fig. 273e-2).

Na and Ca channels are the primary carriers of depolarizing current in both the atria and the ventricles; inactivation of these currents and activation of repolarizing K currents hyperpolarize the heart cells, reestablishing the negative resting membrane potential (Fig. 273e-1B). The *plateau phase* is a time when little current is flowing, and relatively minor changes in depolarizing or repolarizing currents can have profound effects on the shape and duration of the action profile. Mutations in subunits of these channel proteins produce arrhythmogenic alterations in the action potentials that cause long and short QT syndrome, Brugada syndrome, idiopathic ventricular fibrillation, familial atrial fibrillation, and some forms of conduction system disease.

MECHANISMS OF CARDIAC ARRHYTHMIAS

Cardiac arrhythmias result from abnormalities of electrical impulse generation, conduction, or both. Bradyarrhythmias typically arise from disturbances in impulse formation at the level of the sinoatrial node or from disturbances in impulse propagation at any level, including exit block from the sinus node, conduction block in the AVN, and impaired conduction in the His-Purkinje system. Tachyarrhythmias can be classified according to mechanism, including enhanced automaticity (spontaneous depolarization of atrial, junctional, or ventricular pacemakers), triggered arrhythmias (initiated by afterdepolarizations occurring during or immediately after cardiac repolarization, during phase 3 or 4 of the action potential), or reentry (circus propagation of a depolarizing wavefront). A variety of mapping and pacing maneuvers typically performed during invasive electrophysiologic testing can often determine the underlying mechanism of a tachyarrhythmia (Table 273e-1).

Alterations in Impulse Initiation: Automaticity Spontaneous (phase 4) diastolic depolarization underlies the property of automaticity characteristic of pacemaking cells in the sinoatrial (SA) and atrioventricular

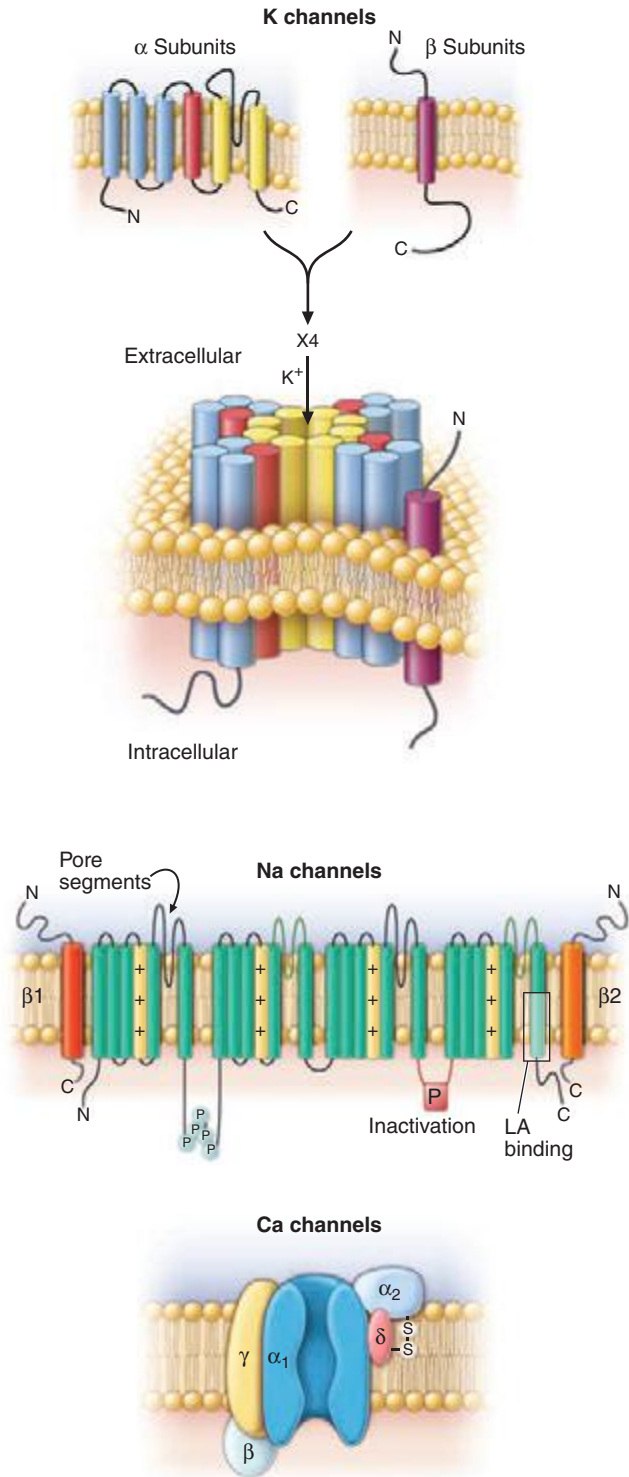


FIGURE 273e-2 Topology and subunit composition of the voltage-dependent ion channels. Potassium channels are formed by the tetramerization of α or pore-forming subunits and one or more β subunits; only single β subunits are shown for clarity. Sodium and calcium channels are composed of α subunits with four homologous domains and one or more ancillary subunits. In all channel types, the loop of protein between the fifth and sixth membrane-spanning repeat in each subunit or domain forms the ion-selective pore. In the case of the sodium channel, the channel is a target for phosphorylation, the linker between the third and fourth homologous domain is critical to inactivation, and the sixth membrane-spanning repeat in the fourth domain is important in local anesthetic antiarrhythmic drug binding. The Ca channel is a multisubunit protein complex with the α_1 subunit containing the pore and major drug binding domain.

TABLE 273e-1 ARRHYTHMIA MECHANISMS

Electrophysiologic Property	Molecular Components	Mechanism	Prototypic Arrhythmias
Cellular			
Impulse Initiation			
Automaticity	I_f , I_{Ca-L} , I_{Ca-T} , I_{K1}	Suppression/acceleration of phase 4	Sinus bradycardia, sinus tachycardia
Triggered automaticity	Calcium overload, I_{T1}	DADs	Digitalis toxicity, reperfusion VT
	I_{Ca-L} , I_{K1} , I_{Na}	EADs	Torsades des pointes, congenital and acquired
Excitation	I_{Na}	Suppression of phase 0	Ischemic VF
	I_{K-ATP}	AP shortening, inexcitability	
	I_{Ca-L}	Suppression	AV block
Repolarization	I_{Na1} , I_{Ca-L} , I_{K1} , I_{K1} , Ca^{2+} homeostasis	AP prolongation, EADs, DADs	Polymorphic VT (HF, LVH)
	I_{Ca-L} , K channels, Ca^{2+} homeostasis	AP shortening	Atrial fibrillation
Multicellular			
Cellular Coupling	Connexins (Cx43), I_{Na1}	Decreased coupling	Ischemic VT/VF
Tissue Structure	I_{K-ATP} , Extracellular matrix, collagen	Excitable gap and functional reentry	Monomorphic VT, atrial fibrillation

Abbreviations: AP, action potential; AV, atrioventricular; DADs, delayed afterdepolarizations; EADs, early afterdepolarizations; HF, heart failure; LVH, left ventricular hypertrophy; VF, ventricular fibrillation; VT, ventricular tachyarrhythmia.

(AV) nodes, His-Purkinje system, coronary sinus, and pulmonary veins. Phase 4 depolarization results from the concerted action of a number of ionic currents, including K^+ currents, Ca^{2+} currents, electrogenic Na, K-ATPase, the Na-Ca exchanger, and the so-called funny, or pacemaker, current (I_f); however, the relative importance of these currents remains controversial.

The rate of phase 4 depolarization and, therefore, the firing rates of pacemaker cells are dynamically regulated. Prominent among the factors that modulate phase 4 is autonomic nervous system tone. The negative chronotropic effect of activation of the parasympathetic nervous system is a result of the release of acetylcholine that binds to muscarinic receptors, releasing G protein $\beta\gamma$ subunits that activate a potassium current (I_{KACH}) in nodal and atrial cells. The resulting increase in K^+ conductance opposes membrane depolarization, slowing the rate of rise of phase 4 of the action potential. Conversely, augmentation of sympathetic nervous system tone increases myocardial catecholamine concentrations, which activate both α - and β -adrenergic receptors. The effect of β_1 -adrenergic stimulation predominates in pacemaking cells, augmenting both L-type Ca current (I_{Ca-L}) and I_f , thus increasing the slope of phase 4. Enhanced sympathetic nervous system activity can dramatically increase the rate of firing of SA nodal cells, producing sinus tachycardia with rates >200 beats/min. By contrast, the increased rate of firing of Purkinje cells is more limited, rarely producing ventricular tachyarrhythmias >120 beats/min.

Normal automaticity may be affected by a number of other factors associated with heart disease. Hypokalemia and ischemia may reduce the activity of Na, K-ATPase, thereby reducing the background repolarizing current and enhancing phase 4 diastolic depolarization. The end result would be an increase in the spontaneous firing rate of pacemaking cells. Modest increases in extracellular potassium may render the maximum diastolic potential more positive, thereby also increasing the firing rate of pacemaking cells. A more significant increase in $[K^+]_o$, however, renders the heart inexcitable by depolarizing the membrane potential.

Normal or enhanced automaticity of subsidiary latent pacemakers produces escape rhythms in the setting of failure of more dominant pacemakers. Suppression of a pacemaker cell by a faster rhythm leads to an increased intracellular Na^+ load ($[Na^+]_i$), and extrusion of Na^+ from the cell by Na, K-ATPase produces an increased background repolarizing current that slows phase 4 diastolic depolarization. At slower rates, $[Na^+]_i$ is decreased, as is the activity of the Na, K-ATPase, resulting in progressively more rapid diastolic depolarization and warm-up of the tachycardia rate. Overdrive suppression and warm-up are characteristic of, but may not be observed in, all automatic tachycardias. Abnormal conduction into tissue with enhanced automaticity (*entrance block*) may blunt or eliminate the phenomena of overdrive suppression and warm-up of automatic tissue.

Abnormal automaticity may produce atrial tachycardia, accelerated idioventricular rhythms, and ventricular tachycardia, particularly associated with ischemia and reperfusion. It has also been suggested that injury currents at the borders of ischemic myocardium may depolarize adjacent nonischemic tissue, predisposing to automatic ventricular tachycardia.

Afterdepolarizations and Triggered Automaticity Triggered automaticity or activity refers to impulse initiation that is dependent on afterdepolarizations (Fig. 273e-3). Afterdepolarizations are membrane voltage oscillations that occur during (early afterdepolarizations, EADs) or after (delayed afterdepolarizations, DADs) an action potential.

The cellular feature common to the induction of DADs is the presence of an increased Ca^{2+} load in the cytosol and sarcoplasmic reticulum. Digitalis glycoside toxicity, catecholamines, and ischemia all can enhance Ca^{2+} loading sufficiently to produce DADs. Accumulation of lysophospholipids in ischemic myocardium with consequent Na^+ and Ca^{2+} overload has been suggested as a mechanism for DADs and triggered automaticity. Cells from damaged areas or cells that survive a myocardial infarction may display spontaneous release of calcium from the sarcoplasmic reticulum, and this may generate “waves” of intracellular calcium elevation and arrhythmias.

EADs occur during the action potential and interrupt the orderly repolarization of the myocyte. Traditionally, EADs have been thought to arise from action potential prolongation and reactivation of depolarizing currents, but more recent experimental evidence suggests a previously unappreciated interrelationship between intracellular calcium loading and EADs. Cytosolic calcium may increase when action potentials are prolonged. This, in turn, appears to enhance L-type Ca current, further prolonging action potential duration as well as providing the inward current driving EADs. Intracellular calcium loading by action potential prolongation may also enhance the likelihood of DADs. The interrelationship among intracellular $[Ca^{2+}]_i$, EADs, and DADs may be one explanation for the susceptibility of hearts that are

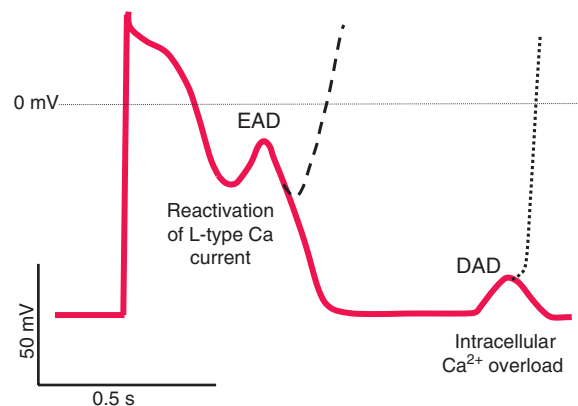


FIGURE 273e-3 Schematic action potentials with early afterdepolarizations (EADs) and delayed afterdepolarizations (DADs).

Afterdepolarizations are spontaneous depolarizations in cardiac myocytes. EADs occur before the end of the action potential (phases 2 and 3), interrupting repolarization. DADs occur during phase 4 of the action potential after completion of repolarization. The cellular mechanisms of EADs and DADs differ (see text).

273e-4 calcium loaded (e.g., in ischemia or congestive heart failure) to develop arrhythmias, particularly on exposure to action potential–prolonging drugs.

EAD-triggered arrhythmias exhibit rate dependence. In general, the amplitude of an EAD is augmented at slow rates when action potentials are longer. Indeed, a fundamental condition that underlies the development of EADs is action potential and QT prolongation. Hypokalemia, hypomagnesemia, bradycardia, and, most commonly, drugs can predispose to the generation of EADs, invariably in the context of prolonging the action potential. Antiarrhythmics with class IA and III action (see below) produce action potential and QT prolongation intended to be therapeutic but frequently causing arrhythmias. Noncardiac drugs such as phenothiazines, nonsedating antihistamines, and some antibiotics can also prolong the action potential duration and predispose to EAD-mediated triggered arrhythmias. Decreased $[K^+]_o$ paradoxically may decrease membrane potassium currents (particularly the delayed rectifier current, I_{Kr}) in the ventricular myocyte, explaining why hypokalemia causes action potential prolongation and EADs. In fact, potassium infusions in patients with the congenital long QT syndrome (LQTS) and in those with drug-induced acquired QT prolongation shorten the QT interval.

EAD-mediated triggered activity probably underlies initiation of the characteristic polymorphic ventricular tachycardia, torsades des pointes, seen in patients with congenital and acquired forms of LQTS. Structural heart disease, such as cardiac hypertrophy and heart failure, may also delay ventricular repolarization (so-called electrical remodeling) and predispose to arrhythmias related to abnormalities of repolarization. The abnormalities of repolarization in hypertrophy and heart failure are often magnified by concomitant drug therapy or electrolyte disturbances.

Abnormal Impulse Conduction: Reentry The most common arrhythmia mechanism is reentry resulting from abnormal electrical impulse conduction and is defined as the circulation of an activation wave around an inexcitable obstacle. The requirements for reentry are two electrophysiologically dissimilar pathways for impulse propagation around an inexcitable region (Fig. 273e-4). Reentry can occur around a fixed anatomic structure (e.g., myocardial scar), with a stable pattern of cardiac depolarization moving in series over the anterograde and retrograde limbs of the circuit. This form of reentry, referred to as *anatomic reentry* or *excitable gap reentry* (see below), is initiated when a depolarizing wavefront encounters an area of unidirectional conduction block in the retrograde limb of the circuit. Conduction across the anterograde limb occurs with a delay that, if of sufficient duration, allows for recovery of conduction in the retrograde limb with reentry of the depolarization wave into the retrograde limb of

the circuit. Sustained reentry requires that the functional dimension of depolarized tissue or the tachycardia wavelength (λ = conduction velocity \times refractory period) fits within the total anatomic length of the circuit, referred to as the path length. When the path length of the circuit exceeds the λ of the tachycardia, the region between the head of the activation wave and the refractory tail is referred to as the excitable gap. Anatomically determined, excitable gap reentry can explain several clinically important tachycardias, such as AV reentry, atrial flutter, bundle branch reentry ventricular tachycardia, and ventricular tachycardia in scarred myocardium.

Reentrant arrhythmias may exist in the heart in the absence of an excitable gap and with a tachycardia wavelength nearly the same size as the path length. In this case, the wavefront propagates through partially refractory tissue without a fixed anatomic obstacle and no fully excitable gap; this is referred to as *leading circle reentry*, a form of functional reentry (reentry that depends on functional properties of the tissue). Unlike excitable gap reentry, there is no fixed anatomic circuit in leading circle reentry, and it may, therefore, not be possible to disrupt the tachycardia with pacing or destruction of a part of the circuit. Furthermore, the circuit in leading circle reentry tends to be less stable than that in excitable gap reentrant arrhythmias, with large variations in cycle length and a predilection to termination. There is strong evidence to suggest that less organized arrhythmias, such as atrial and ventricular fibrillation, are associated with more complex activation of the heart and are due to functional reentry.

Catheter-based and pharmacologic therapies for reentrant arrhythmias are designed to disrupt the anatomic circuit or alter the relationship between the wavelength and path length of the arrhythmia circuit, eliminating pathologic conduction. For example, antiarrhythmic drugs that prolong the action potential (Class III) are effective if they sufficiently prolong the λ such that it can no longer fit within the anatomic circuit. Catheter ablation is often undertaken with the goal of identifying and destroying a critical limb of the reentrant circuit (i.e., ablation of the cavotricuspid isthmus in the treatment of typical, right atrial flutter). Due to the less defined pathways of myocardial activation seen in functional reentry, ablation of these rhythms tends to target initiating triggers (e.g., pulmonary vein potentials in catheter ablation of atrial fibrillation) rather than the anatomic circuit.

Structural heart disease is associated with changes in conduction and refractoriness that increase the risk of reentrant arrhythmias. Chronically ischemic myocardium exhibits a downregulation of the gap junction channel protein (connexin 43) that carries intercellular ionic current. The border zones of infarcted and failing ventricular myocardium exhibit not only functional alterations of ionic currents but also remodeling of tissue and altered distribution of gap junctions. The changes in gap junction channel expression and distribution, in combination with macroscopic tissue alterations, support a role for slowed conduction in reentrant arrhythmias that complicate chronic coronary artery disease (CAD). Aged human atrial myocardium exhibits altered conduction, manifest as highly fractionated atrial electrograms, producing an ideal substrate for the reentry that may underlie the very common development of atrial fibrillation in the elderly.

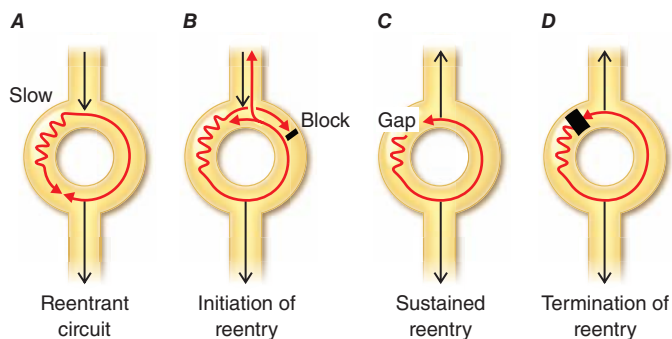


FIGURE 272-4 Schematic diagram of reentry. **A.** The circuit contains two limbs, one with slow conduction. **B.** A premature impulse blocks in the fast pathway and conducts over the slow pathway, allowing the fast pathway to recover so that the activation wave can reenter the fast pathway from the retrograde direction. **C.** During sustained reentry utilizing such a circuit, a gap (excitable gap) exists between the activating head of the wave and the recovering tail. **D.** One mechanism of termination of reentry occurs when the conduction and recovery characteristics of the circuit change and the activating head of the wave collides with the tail, extinguishing the tachycardia.

APPROACH TO THE PATIENT: Cardiac Arrhythmias

The evaluation of patients with suspected cardiac arrhythmias is highly individualized; however, two key features—the history and ECG—are pivotal in directing the diagnostic workup and therapy. Patients with cardiac arrhythmias exhibit a wide spectrum of clinical presentations that range from asymptomatic ECG abnormalities to survival from cardiac arrest. In general, the more severe the presenting symptoms are, the more aggressive the evaluation and treatment are. Loss of consciousness that is believed to be of cardiac origin typically mandates an exhaustive search for the etiology and often requires invasive, device-based therapy. The presence of structural heart disease and prior myocardial infarction dictates a change in the approach to the management of syncope or ventricular

arrhythmias. The presence of a family history of serious ventricular arrhythmias or premature sudden death will influence the evaluation of presumed heritable arrhythmias.

The physical examination is focused on determining whether there is cardiopulmonary disease that is associated with specific cardiac arrhythmias. The absence of significant cardiopulmonary disease often, but not always, suggests benignity of the rhythm disturbance. In contrast, palpitations, syncope, or near syncope in the setting of significant heart or lung disease have more ominous implications. In addition, the physical examination may reveal the presence of a persistent arrhythmia such as atrial fibrillation.

The judicious use of noninvasive diagnostic tests is an important element in the evaluation of patients with arrhythmias, and there is no test more important than the ECG, particularly if recorded at the time of symptoms. Uncommon but diagnostically important signatures of electrophysiologic disturbances may be unearthed on the resting ECG, such as delta waves in Wolff-Parkinson-White (WPW) syndrome, prolongation or shortening of the QT interval, right precordial ST-segment abnormalities in Brugada syndrome, and epsilon waves in arrhythmogenic right ventricular dysplasia. Variants of body surface ECG recording can provide important information about arrhythmia substrates and triggers. Holter monitoring and event recording, either continuous or intermittent, record the body surface ECG over longer periods, enhancing the possibility of observing the cardiac rhythm during symptoms. Holter monitoring is particularly useful in assessing daily symptoms thought to be attributable to arrhythmia or for quantifying a particular arrhythmia phenomenon (e.g., premature ventricular complex burden). Ambulatory event monitors are indicated when symptoms thought to be due to arrhythmia occur less frequently (i.e., several episodes per month), and, because the monitors are typically patient-activated, they are optimal for correlating symptoms with rhythm disturbances. Implantable long-term monitors permit prolonged telemetric monitoring both for diagnosis and to assess the efficacy of therapy. Implantable monitors are typically used for the evaluation of malignant symptoms that occur quite infrequently and that cannot be provoked at diagnostic electrophysiology study.

Exercise electrocardiography is important in determining the presence of myocardial demand ischemia; more recently, analysis of the morphology of the QT interval with exercise has been used to assess the risk of serious ventricular arrhythmias. The exercise ECG may be particularly useful in patients with symptoms that occur during activity. Cardiac imaging plays an important role in the detection and characterization of myocardial structural abnormalities that may render the heart more susceptible to arrhythmia. Ventricular tachyarrhythmias, for instance, occur more frequently in patients with ventricular systolic dysfunction and chamber dilation, in hypertrophic cardiomyopathy, and in the setting of infiltrative diseases such as sarcoidosis. Supraventricular arrhythmias may be associated with particular congenital conditions, including AV reentry in the setting of Ebstein's anomaly. Echocardiography is a frequently employed imaging technique to screen for disorders of cardiac structure and function. Increasingly, magnetic resonance imaging of the myocardium is being used to screen for scar burden, fibrofatty infiltration of the myocardium as seen in arrhythmogenic right ventricular cardiomyopathy, and other structural changes that affect arrhythmia susceptibility.

Head-up tilt (HUT) testing is useful in the evaluation of patients with syncope in whom there is a suspicion that exaggerated vagal tone or vasodepression may play a causal role. The physiologic response to HUT is incompletely understood; however, redistribution of blood volume and increased ventricular contractility occur consistently. Exaggerated activation of a central reflex in response to HUT produces a stereotypic response of an initial increase in heart rate, then a drop in blood pressure followed by a reduction in heart rate characteristic of neurally mediated hypotension. Other responses to HUT may be observed in patients with orthostatic

hypotension and autonomic insufficiency. HUT is used most often in patients with recurrent syncope, although it may be useful in patients with single syncopal episodes with associated injury, particularly in the absence of structural heart disease. In patients with structural heart disease, HUT may be indicated in those with syncope, in whom other causes (e.g., asystole, ventricular tachyarrhythmias) have been excluded. HUT has been suggested as a useful tool in the diagnosis of and therapy for recurrent idiopathic vertigo, chronic fatigue syndrome, recurrent transient ischemic attacks, and repeated falls of unknown etiology in the elderly. Importantly, HUT is relatively contraindicated in the presence of severe CAD with proximal coronary stenoses, known severe cerebrovascular disease, severe mitral stenosis, and obstruction to left ventricular outflow (e.g., aortic stenosis).

Electrophysiologic testing is central to the understanding and treatment of many cardiac arrhythmias. Indeed, most frequently, electrophysiologic testing is interventional, providing both diagnosis and therapy. The indications for electrophysiologic testing fall into several categories: to define the mechanism of an arrhythmia; to deliver catheter-based ablative treatment; and to determine the etiology of symptoms that may be caused by an arrhythmia (e.g., syncope, palpitations). The components of the electrophysiologic test are baseline measurements of conduction under resting and stressed (rate or pharmacologic) conditions and maneuvers, both pacing and pharmacologic, to induce arrhythmias. A number of sophisticated electrical mapping and catheter-guidance techniques have been developed to facilitate catheter-based therapeutics in the electrophysiology laboratory.

TREATMENT CARDIAC ARRHYTHMIAS

ANTIARRHYTHMIC DRUG THERAPY

The interaction of antiarrhythmic drugs with cardiac tissues and the resulting electrophysiologic changes are complex. An incomplete understanding of the effects of these drugs has produced serious missteps that have had adverse effects on patient outcomes and the development of newer pharmacologic agents. Currently, antiarrhythmic drugs have been relegated to an ancillary role in the treatment of most cardiac arrhythmias.

There are several explanations for the complexity of antiarrhythmic drug action: the structural similarity of target ion channels; regional differences in the levels of expression of channels and transporters, which change with disease; time and voltage dependence of drug action; and the effect of these drugs on targets other than ion channels. Because of the limitations of any scheme to classify antiarrhythmic agents, a shorthand that is useful in describing the major mechanisms of action is of some utility. Such a classification scheme was proposed in 1970 by Vaughan-Williams and later modified by Singh and Harrison. The classes of antiarrhythmic action are class I, local anesthetic effect due to blockade of Na⁺ current; class II, interference with the action of catecholamines at the β-adrenergic receptor; class III, delay of repolarization due to inhibition of K⁺ current or activation of depolarizing current; and class IV, interference with calcium conductance (Table 273e-2). Class I antiarrhythmics have been further subdivided based on the kinetics and potency of Na⁺ channel binding; class Ia agents (quinidine, procainamide) are those with moderate potency and intermediate kinetics; class Ib agents (lidocaine, mexiletine) are those with low potency and rapid kinetics; and class Ic drugs (flecainide, propafenone) are those with high potency and the slowest kinetics. The limitations of the Vaughan-Williams classification scheme include multiple actions of most drugs, overwhelming consideration of antagonism as a mechanism of action, and the fact that several agents have none of the four classes of action in the scheme.

CATHETER ABLATION

The use of catheter ablation is based on the principle that there is a critical anatomic region of impulse generation or propagation that

TABLE 273e-2 ANTIARRHYTHMIC DRUG ACTIONS

Drug	Class Actions				Miscellaneous Action
	I	II	III	IV	
Quinidine	++		++		α -Adrenergic blockade
Procainamide	++		++		Ganglionic blockade
Flecainide	+++		+		
Propafenone	++	+			
Sotalol		++	+++		
Dofetilide			+++		
Amiodarone	++	++	+++	+	α -Adrenergic blockade
Ibutilide			+++		Na ⁺ channel activator

is required for the initiation and maintenance of cardiac arrhythmias. Destruction of such a critical region results in the elimination of the arrhythmia. The use of radiofrequency (RF) energy in clinical medicine is nearly a century old. The first catheter ablation using a

DC energy source was performed in the early 1980s by Scheinman and colleagues. By the early 1990s, RF had been adapted for use in catheter-based ablation in the heart (**Fig. 273e-5**).

The RF band (300–30,000 kHz) is used to generate energy for several biomedical applications, including coagulation and cauterization of tissues. Energy of this frequency will not stimulate skeletal muscle or the heart and heats tissue by a resistive mechanism, with the intensity of heating and tissue destruction being proportional to the delivered power. Alternative, less frequently used energy sources for catheter ablation of cardiac arrhythmias include microwaves (915 MHz or 2450 MHz), lasers, ultrasound, and freezing (cryoablation). Of these alternative ablation techniques, cryoablation is being used clinically with the most frequency, especially ablation in the region of the AVN. At temperatures just below 32°C, membrane ion transport is disrupted, producing depolarization of cells, decreased action potential amplitude and duration, and slowed conduction velocity (resulting in local conduction block)—all of which are reversible if the tissue is rewarmed in a timely fashion.

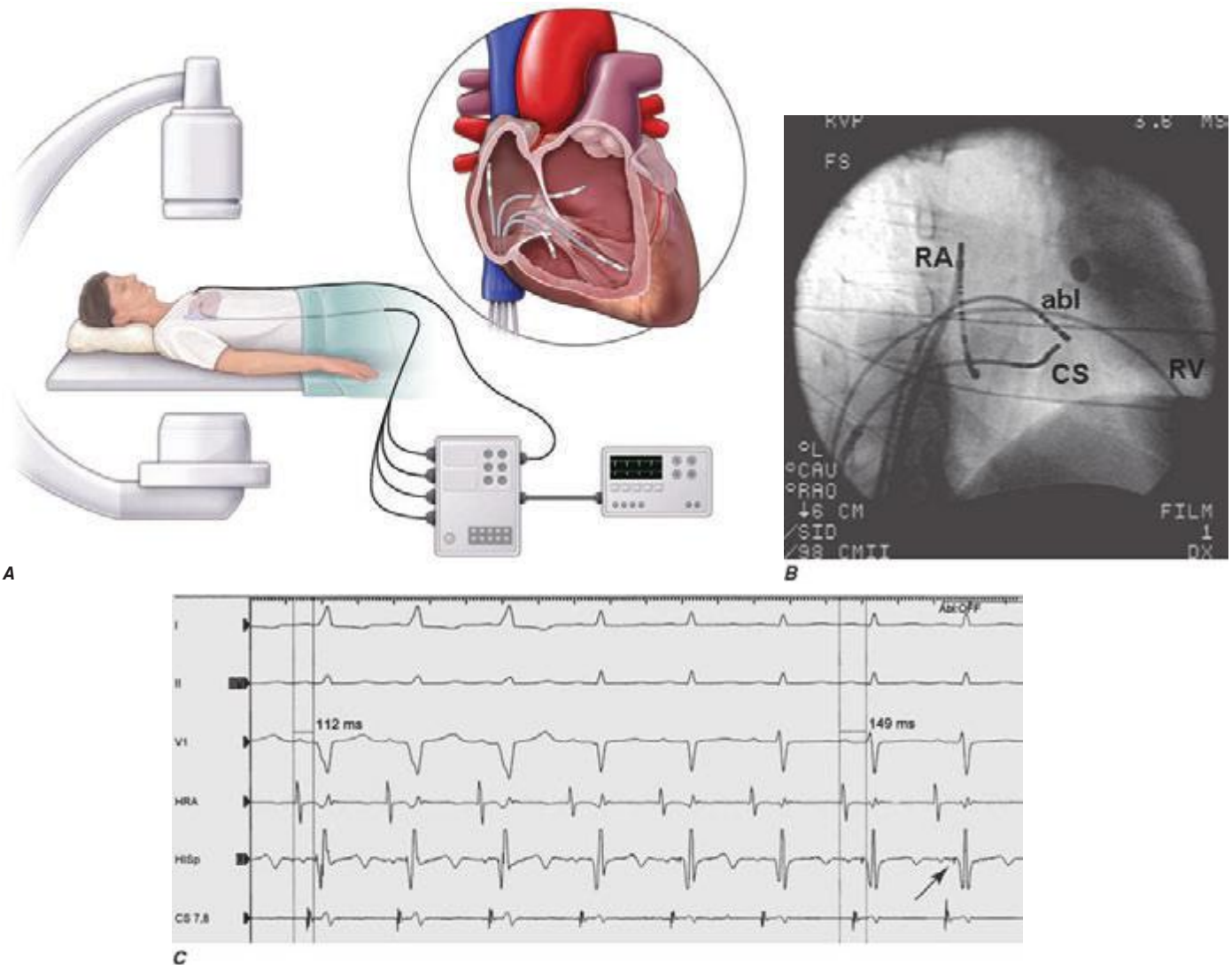


FIGURE 273e-5 Catheter ablation of cardiac arrhythmias. A. A schematic of the catheter system and generator in a patient undergoing radiofrequency catheter ablation (RFCA); the circuit involves the catheter in the heart and a dispersive patch placed on the body surface (usually the back). The inset shows a diagram of the heart with a catheter located at the AV valve ring for ablation of an accessory pathway. **B.** A right anterior oblique fluoroscopic image of the catheter position for ablation of a left-sided accessory pathway. A catheter is placed in the atrial side of the mitral valve ring (abl) via a transseptal puncture. Other catheters are placed in the coronary sinus (CS), in the right atrium (RA), and in the right ventricular (RV) apex to record local electrical activation. **C.** Body surface electrocardiogram recordings (I, II, V₁) and endocardial electrograms (HRA, high right atrium; HISp, proximal His bundle electrogram; CS 7, 8, recordings from poles 7 and 8 of a decapolar catheter placed in the coronary sinus) during RFCA of a left-sided accessory pathway in a patient with Wolff-Parkinson-White syndrome. The QRS narrows at the fourth complex; the arrow shows the His bundle electrogram, which becomes apparent with elimination of ventricular preexcitation over the accessory pathway.

Tissue cooling can be used for mapping and ablation. Cryomapping can be used to confirm the location of a desired ablation target, such as an accessory pathway in WPW syndrome, or can be used to determine the safety of ablation around the AVN by monitoring AV conduction during cooling. Another advantage of cryoablation is that once the catheter tip cools below freezing, it adheres to the tissue, increasing catheter stability independent of the rhythm or pacing.

DEVICE THERAPY

Bradyarrhythmias due either to primary sinus node dysfunction or to atrioventricular conduction defects are readily treated through implantation of a permanent pacemaker. Clinical indications for pacemaker implantation often depend on the presence either of symptomatic bradycardia or of an unreliable endogenous escape rhythm and are more fully reviewed in [Chaps. 274 and 275](#).

Ventricular tachyarrhythmias, particularly those occurring in the context of progressive structural heart diseases such as ischemic cardiomyopathy or arrhythmogenic right ventricular cardiomyopathy, may recur despite therapy with antiarrhythmic drugs or catheter ablation. In appropriate candidates, implantation of an internal cardioverter-defibrillator (ICD) may reduce mortality rates from sudden cardiac death. In a subset of patients with congestive heart failure (CHF) and ventricular mechanical dyssynchrony, ICD or pacemaker platforms can be used to provide cardiac resynchronization therapy, typically through implantation of a left ventricular pacing lead. In patients with dyssynchronous CHF, such therapy has been shown to improve both morbidity and mortality rates.

274 The Bradyarrhythmias: Disorders of the Sinoatrial Node

David D. Spragg, Gordon F. Tomaselli

Electrical activation of the heart normally originates in the sinoatrial (SA) node, the predominant pacemaker. Other subsidiary pacemakers in the atrioventricular (AV) node, specialized conducting system, and muscle may initiate electrical activation if the SA node is dysfunctional or suppressed. Typically, subsidiary pacemakers discharge at a slower rate and, in the absence of an appropriate increase in stroke volume, may result in tissue hypoperfusion.

Spontaneous activation and contraction of the heart are a consequence of the specialized pacemaking tissue in these anatomic locales. As described in [Chap. 273e](#), action potentials in the heart are regionally heterogeneous. The action potentials in cells isolated from nodal tissue are distinct from those recorded from atrial and ventricular myocytes ([Fig. 274-1](#)). The complement of ionic currents present in nodal cells results in a less

negative resting membrane potential compared with atrial or ventricular myocytes. Electrical diastole in nodal cells is characterized by slow diastolic depolarization (phase 4), which generates an action potential as the membrane voltage reaches threshold. The action potential upstrokes (phase 0) are slow compared with atrial or ventricular myocytes, being mediated by calcium rather than sodium current. Cells with properties of SA and AV nodal tissue are electrically connected to the remainder of the myocardium by cells with an electrophysiologic phenotype between that of nodal cells and that of atrial or ventricular myocytes. Cells in the SA node exhibit the most rapid phase 4 depolarization and thus are the dominant pacemakers in a normal heart.

Bradycardia results from a failure of either impulse initiation or impulse conduction. Failure of impulse initiation may be caused by depressed automaticity resulting from a slowing or failure of phase 4 diastolic depolarization ([Fig. 274-2](#)), which may result from disease or exposure to drugs. Prominently, the autonomic nervous system modulates the rate of phase 4 diastolic depolarization and thus the firing rate of both primary (SA node) and subsidiary pacemakers. Failure of conduction of an impulse from nodal tissue to atrial or ventricular myocardium may produce bradycardia as a result of exit block. Conditions that alter the activation and connectivity of cells (e.g., fibrosis) in the heart may result in failure of impulse conduction.

SA node dysfunction and AV conduction block are the most common causes of pathologic bradycardia. SA node dysfunction may be difficult to distinguish from physiologic sinus bradycardia, particularly in the young. SA node dysfunction increases in frequency between the fifth and sixth decades of life and should be considered in patients with fatigue, exercise intolerance, or syncope and sinus bradycardia.

Permanent pacemaking is the only reliable therapy for symptomatic bradycardia in the absence of extrinsic and reversible etiologies such as increased vagal tone, hypoxia, hypothermia, and drugs ([Table 274-1](#)). Approximately 50% of the 150,000 permanent pacemakers implanted in the United States and 20–30% of the 150,000 of those in Europe were implanted for SA node disease.

STRUCTURE AND PHYSIOLOGY OF THE SA NODE

The SA node is composed of a cluster of small fusiform cells in the sulcus terminalis on the epicardial surface of the heart at the right atrial–superior vena caval junction, where they envelop the SA nodal artery. The SA node is structurally heterogeneous, but the central prototypic nodal cells have fewer distinct myofibrils than does the surrounding atrial myocardium, no intercalated disks visible on light microscopy, a poorly developed sarcoplasmic reticulum, and no T-tubules. Cells in the peripheral regions of the SA node are transitional in both structure and function. The SA nodal artery arises from the right coronary artery in 55–60% and the left circumflex artery in 40–45% of persons. The SA node is richly innervated by sympathetic and parasympathetic nerves and ganglia.

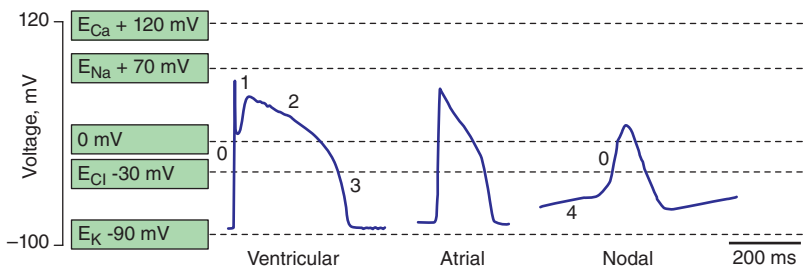


FIGURE 274-1 Action potential profiles recorded in cells isolated from sinoatrial or atrioventricular nodal tissue compared with those of cells from atrial or ventricular myocardium. Nodal cell action potentials exhibit more depolarized resting membrane potentials, slower phase 0 upstrokes, and phase 4 diastolic depolarization.

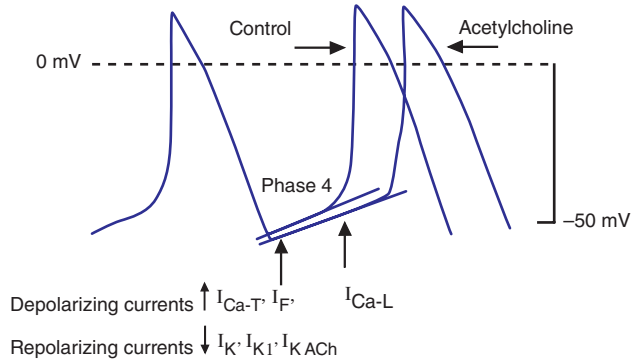


FIGURE 274-2 Schematics of nodal action potentials and the currents that contribute to phase 4 depolarization. Relative increases in depolarizing L- (I_{Ca-L}) and T- (I_{Ca-T}) type calcium and pacemaker currents (I_p) along with a reduction in repolarizing inward rectifier (I_{K1}) and delayed rectifier (I_{Kr}) potassium currents result in depolarization. Activation of ACh-gated (I_{KACH}) potassium current and beta blockade slow the rate of phase 4 and decrease the pacing rate. (Modified from J Jalife et al: *Basic Cardiac Electrophysiology for the Clinician*, Blackwell Publishing, 1999.)

Irregular and slow propagation of impulses from the SA node can be explained by the electrophysiology of nodal cells and the structure of the SA node itself. The action potentials of SA nodal cells are characterized by a relatively depolarized membrane potential (Fig. 274-1) of -40 to -60 mV, slow phase 0 upstroke, and relatively rapid phase 4 diastolic depolarization compared with the action potentials recorded in cardiac muscle cells. The relative absence of inward rectifier

potassium current (I_{K1}) accounts for the depolarized membrane potential; the slow upstroke of phase 0 results from the absence of available fast sodium current (I_{Na}) and is mediated by L-type calcium current (I_{Ca-L}); and phase 4 depolarization is a result of the aggregate activity of a number of ionic currents. Prominently, both L- and T-type (I_{Ca-T}) calcium currents, the pacemaker current (so-called funny current, or I_p) formed by hyperpolarization-activated cyclic nucleotide-gated channels, and the electrogenic sodium-calcium exchanger provide depolarizing current that is antagonized by delayed rectifier (I_{Kr}) and acetylcholine-gated (I_{KACH}) potassium currents. I_{Ca-L} , I_{Ca-T} , and I_p are modulated by β -adrenergic stimulation and I_{KACH} by vagal stimulation, explaining the exquisite sensitivity of diastolic depolarization to autonomic nervous system activity. The slow conduction within the SA node is explained by the absence of I_{Na} and poor electrical coupling of cells in the node, resulting from sizable amounts of interstitial tissue and a low abundance of gap junctions. The poor coupling allows for graded electrophysiologic properties within the node, with the peripheral transitional cells being silenced by electrotonic coupling to atrial myocardium.

ETIOLOGY OF SA NODAL DISEASE

SA nodal dysfunction has been classified as intrinsic or extrinsic. The distinction is important because extrinsic dysfunction is often reversible and generally should be corrected before pacemaker therapy is considered (Table 274-1). The most common causes of extrinsic SA node dysfunction are drugs and autonomic nervous system influences that suppress automaticity and/or compromise conduction. Other extrinsic causes include hypothyroidism, sleep apnea, and conditions likely to occur in critically ill patients such as hypothermia, hypoxia, increased intracranial pressure (Cushing's response), and endotracheal suctioning via activation of the vagus nerve.

Intrinsic sinus node dysfunction is degenerative and often is characterized pathologically by fibrous replacement of the SA node or its connections to the atrium. Acute and chronic coronary artery disease (CAD) may be associated with SA node dysfunction, although in the setting of acute myocardial infarction (MI; typically inferior), the abnormalities are transient. Inflammatory processes may alter SA node function, ultimately producing replacement fibrosis. Pericarditis, myocarditis, and rheumatic heart disease have been associated with SA nodal disease with sinus bradycardia, sinus arrest, and exit block. Carditis associated with systemic lupus erythematosus (SLE), rheumatoid arthritis (RA), and mixed connective tissue disorders (MCTDs) may also affect SA node structure and function. Senile amyloidosis is an infiltrative disorder in patients typically in the ninth decade of life; deposition of amyloid protein in the atrial myocardium can impair SA node function. Some SA node disease is iatrogenic and results from direct injury to the SA node during cardiothoracic surgery.

Rare heritable forms of sinus node disease have been described, and several have been characterized genetically. Autosomal dominant sinus node dysfunction in conjunction with supraventricular tachycardia (i.e., tachycardia-bradycardia variant of sick-sinus syndrome [SSS2]) has been linked to mutations in the pacemaker current (I_p) subunit gene *HCN4* on chromosome 15. An autosomal recessive form of SSS1 with the prominent feature of atrial inexcitability and absence of P waves on the electrocardiogram (ECG) is caused by mutations in the cardiac sodium channel gene, *SCN5A*, on chromosome 3. Variants in myosin heavy chain 6 (*MYH6*) increase the susceptibility to SSS (SSS3). SA node dysfunction associated with myopia has been described but not genetically characterized. There are several neuromuscular diseases, including Kearns-Sayre syndrome (ophthalmoplegia, pigmentary degeneration of the retina, and cardiomyopathy) and myotonic dystrophy, that have a predilection for the conducting system and SA node.

SSS in both the young and the elderly is associated with an increase in fibrous tissue in the SA node. The onset of SSS may be hastened by coexisting disease, such as CAD, diabetes mellitus, hypertension, and valvular diseases and cardiomyopathies.

CLINICAL FEATURES OF SA NODE DISEASE

SA node dysfunction may be completely asymptomatic and manifest as an ECG anomaly such as sinus bradycardia; sinus arrest and

TABLE 274-1 ETIOLOGIES OF SA NODE DYSFUNCTION

Extrinsic	Intrinsic
Autonomic	Sick-sinus syndrome (SSS)
Carotid sinus hypersensitivity	Coronary artery disease (chronic and acute MI)
Vasovagal (cardioinhibitory) stimulation	Inflammatory
Drugs	Pericarditis
Beta blockers	Myocarditis (including viral)
Calcium channel blockers	Rheumatic heart disease
Digoxin	Collagen vascular diseases
Ivabradine	Lyme disease
Antiarrhythmics (class I and III)	Senile amyloidosis
Adenosine	Congenital heart disease
Clonidine (other sympatholytics)	TGA/Mustard and Fontan repairs
Lithium carbonate	Iatrogenic
Cimetidine	Radiation therapy
Amitriptyline	Postsurgical
Phenothiazines	Chest trauma
Narcotics (methadone)	Familial
Pentamidine	SSS2, AD, OMIM #163800 (15q24-25)
Hypothyroidism	SSS1, AR OMIM #608567 (3p21)
Sleep apnea	SSS3, AD, OMIM #614090 (14q11.2)
Hypoxia	SA node disease with myopia, OMIM #182190
Endotracheal suctioning (vagal maneuvers)	Kearns-Sayre syndrome, OMIM #530000
Hypothermia	Myotonic dystrophy
Increased intracranial pressure	Type 1, OMIM #160900 (19q13.2-13.3)
	Type 2, OMIM #602668 (3q13.3-q24)
	Friedreich's ataxia, OMIM #229300 (9q13, 9p23-p11)

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; MI, myocardial infarction; OMIM, Online Mendelian Inheritance in Man (database); TGA, transposition of the great arteries.

exit block; or alternating supraventricular tachycardia, usually atrial fibrillation, and bradycardia. Symptoms associated with SA node dysfunction, in particular tachycardia-bradycardia syndrome, may be related to both slow and fast heart rates. For example, tachycardia may be associated with palpitations, angina pectoris, and heart failure, and bradycardia may be associated with hypotension, syncope, presyncope, fatigue, and weakness. In the setting of SSS, overdrive suppression of the SA node may result in prolonged pauses and syncope upon termination of the tachycardia. In many cases, symptoms associated with SA node dysfunction result from concomitant cardiovascular disease. A significant minority of patients with SSS develop signs and symptoms of heart failure that may be related to slow or fast heart rates.

One-third to one-half of patients with SA node dysfunction develop supraventricular tachycardia, usually atrial fibrillation or atrial flutter. The incidence of persistent atrial fibrillation in patients with SA node dysfunction increases with advanced age, hypertension, diabetes mellitus, left ventricular dilation, valvular heart disease, and ventricular pacing. Remarkably, some symptomatic patients may experience an improvement in symptoms with the development of atrial fibrillation, presumably from an increase in their average heart rate. Patients with the tachycardia-bradycardia variant of SSS, similar to patients with atrial fibrillation, are at risk for thromboembolism, and *those at greatest risk*, including patients ≥ 65 years and patients with a prior history of stroke, valvular heart disease, left ventricular dysfunction, or atrial enlargement, should be treated with anticoagulants. Up to one-quarter of patients with SA node disease will have concurrent AV conduction disease, although only a minority will require specific therapy for high-grade AV block.

The natural history of SA node dysfunction is one of varying intensity of symptoms even in patients who present with syncope. Symptoms related to SA node dysfunction may be significant, but overall mortality usually is not compromised in the absence of other significant comorbid conditions. These features of the natural history need to be taken into account in considering therapy for these patients.

ELECTROCARDIOGRAPHY OF SA NODE DISEASE

The electrocardiographic manifestations of SA node dysfunction include sinus bradycardia, sinus pauses, sinus arrest, sinus exit block, tachycardia (in SSS), and chronotropic incompetence. It is often difficult to distinguish pathologic from physiologic sinus bradycardia. By definition, sinus bradycardia is a rhythm driven by the SA node with a rate of < 60 beats/min; sinus bradycardia is very common and typically benign. Resting heart rates < 60 beats/min are very common in young healthy individuals and physically conditioned subjects. A sinus rate of < 40 beats/min in the awake state in the absence of physical conditioning generally is considered abnormal. Sinus pauses and sinus arrest result from failure of the SA node to discharge, producing a pause without P waves visible on the ECG (Fig. 274-3). Sinus pauses of up to 3 s are common in awake athletes, and pauses of this duration or longer may be observed in asymptomatic elderly subjects. Intermittent failure of conduction from the SA node produces sinus exit block. The severity of sinus exit block may vary in a manner similar to that of AV block (Chap. 275). Prolongation of conduction from the sinus node will not be apparent on the ECG; second-degree SA block will

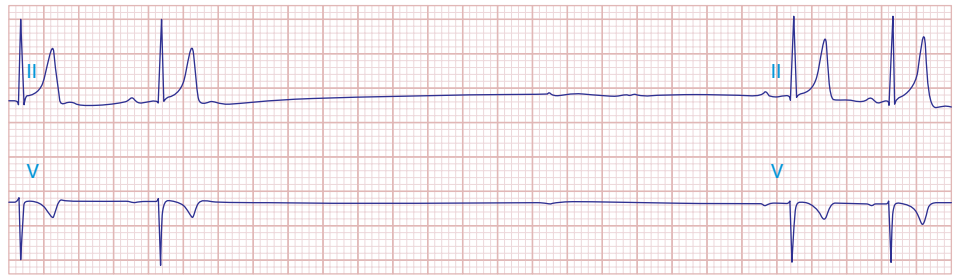


FIGURE 274-3 Sinus slowing and pauses on the electrocardiogram (ECG). The ECG is recorded during sleep in a young patient without heart disease. The heart rate before the pause is slow, and the PR interval is prolonged, consistent with an increase in vagal tone. The P waves have a morphology consistent with sinus rhythm. The recording is from a two-lead telemetry system in which the tracing labeled II mimics frontal lead II and V represents Modified Central Lead 1, which mimics lead V₁ of the standard 12-lead ECG.

produce intermittent conduction from the SA node and a regularly irregular atrial rhythm.

Type I second-degree SA block results from progressive prolongation of SA node conduction with intermittent failure of the impulses originating in the sinus node to conduct to the surrounding atrial tissue. Second-degree SA block appears on the ECG as an intermittent absence of P waves (Fig. 274-4). In type II second-degree SA block, there is no change in SA node conduction before the pause. Complete or third-degree SA block results in no P waves on the ECG. Tachycardia-bradycardia syndrome is manifest as alternating sinus bradycardia and atrial tachyarrhythmias. Although atrial tachycardia, atrial flutter, and atrial fibrillation may be observed, the latter is the most common tachycardia. Chronotropic incompetence is the inability to increase the heart rate in response to exercise or other stress appropriately and is defined in greater detail below.

DIAGNOSTIC TESTING

SA node dysfunction is most commonly a clinical or electrocardiographic diagnosis. Sinus bradycardia or pauses on the resting ECG are rarely sufficient to diagnose SA node disease, and longer-term recording and symptom correlation generally are required. Symptoms in the absence of sinus bradyarrhythmias may be sufficient to exclude a diagnosis of SA node dysfunction.

Electrocardiographic recording plays a central role in the diagnosis and management of SA node dysfunction. Despite the limitations of the resting ECG, longer-term recording employing Holter or event monitors may permit correlation of symptoms with the cardiac rhythm. Many contemporary event monitors may be automatically triggered to record the ECG when certain programmed heart rate criteria are met. Implantable ECG monitors permit long-term recording (12–18 months) in particularly challenging patients.

Failure to increase the heart rate with exercise is referred to as *chronotropic incompetence*. This is alternatively defined as failure to reach 85% of predicted maximal heart rate at peak exercise or failure to achieve a heart rate > 100 beats/min with exercise or a maximal heart rate with exercise less than two standard deviations below that of an age-matched control population. Exercise testing may be useful in dis-

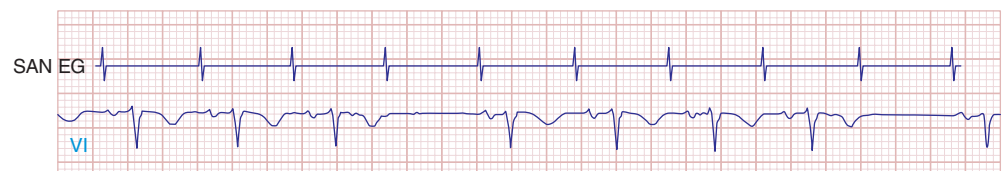


FIGURE 274-4 Mobitz type I SA nodal exit block. A theoretical SA node electrogram (SAN EG) is shown. Note that there is grouped beating producing a regularly irregular heart rhythm. The SA node EG rate is constant with progressive delay in exit from the node and activation of the atria, inscribing the P wave. This produces subtly decreasing P-P intervals before the pause, and the pause is less than twice the cycle length of the last sinus interval.

criminating chronotropic incompetence from resting bradycardia and may aid in the identification of the mechanism of exercise intolerance.

Autonomic nervous system testing is useful in diagnosing carotid sinus hypersensitivity; pauses >3 s are consistent with the diagnosis but may be present in asymptomatic elderly subjects. Determining the intrinsic heart rate (IHR) may distinguish SA node dysfunction from slow heart rates that result from high vagal tone. The normal IHR after administration of 0.2 mg/kg propranolol and 0.04 mg/kg atropine is $117.2 - (0.53 \times \text{age})$ in beats/min; a low IHR is indicative of SA disease.

Electrophysiologic testing may play a role in the assessment of patients with presumed SA node dysfunction and in the evaluation of syncope, particularly in the setting of structural heart disease. In this circumstance, electrophysiologic testing is used to rule out more malignant etiologies of syncope, such as ventricular tachyarrhythmias and AV conduction block. There are several ways to assess SA node function invasively. They include the sinus node recovery time (SNRT), defined as the longest pause after cessation of overdrive pacing of the right atrium near the SA node (normal: <1500 ms or, corrected for sinus cycle length, <550 ms), and the sinoatrial conduction time (SACT), defined as one-half the difference between the intrinsic sinus cycle length and a noncompensatory pause after a premature atrial stimulus (normal <125 ms). The combination of an abnormal SNRT, an abnormal SACT, and a low IHR is a sensitive and specific indicator of intrinsic SA node disease.

TREATMENT SINOATRIAL NODE DYSFUNCTION

Since SA node dysfunction is not associated with increased mortality rates, the aim of therapy is alleviation of symptoms. Exclusion of extrinsic causes of SA node dysfunction and correlation of the cardiac rhythm with symptoms is an essential part of patient management. Pacemaker implantation is the primary therapeutic intervention in patients with symptomatic SA node dysfunction. Pharmacologic considerations are important in the evaluation and management of patients with SA nodal disease. A number of drugs modulate SA node function and are extrinsic causes of dysfunction (Table 274-1). Beta blockers and calcium channel blockers increase SNRT in patients with SA node dysfunction, and antiarrhythmic drugs with class I and III action may promote SA node exit block. In general, such agents should be discontinued before decisions regarding the need for permanent pacing in patients with SA node disease are made. Chronic pharmacologic therapy for sinus bradyarrhythmias is limited. Some pharmacologic agents may improve SA node function; digitalis, for example, has been shown to shorten SNRT in patients with SA node dysfunction. Isoproterenol or atropine administered IV may increase the sinus rate acutely. Theophylline has been used both acutely and chronically to increase heart rate but has liabilities when used in patients with tachycardia-bradycardia syndrome, increasing the frequency of supraventricular tachyarrhythmias, and in patients with structural heart disease, increasing the risk of potentially serious ventricular arrhythmias. Currently, there is only a single randomized study of therapy for SA node dysfunction. In patients with resting heart rates <50 and >30 beats/min on a Holter monitor, patients who received dual-chamber pacemakers experienced significantly fewer syncopal episodes and had symptomatic improvement compared with patients randomized to theophylline or no treatment.

In certain circumstances, sinus bradycardia requires no specific treatment or only temporary rate support. Sinus bradycardia is common in patients with acute inferior or posterior MI and can be exacerbated by vagal activation induced by pain or the use of drugs such as morphine. Ischemia of the SA nodal artery probably occurs in acute coronary syndromes more typically with involvement with the right coronary artery, and even with infarction, the effect on SA node function most often is transient.

Sinus bradycardia is a prominent feature of carotid sinus hypersensitivity and neurally mediated hypotension associated with vasovagal syncope that responds to pacemaker therapy. Carotid

hypersensitivity with recurrent syncope or presyncope associated with a predominant cardioinhibitory component responds to pacemaker implantation. Several randomized trials have investigated the efficacy of permanent pacing in patients with drug-refractory vasovagal syncope, with mixed results. Although initial trials suggested that patients undergoing pacemaker implantation have fewer recurrences and a longer time to recurrence of symptoms, at least one follow-up study did not confirm these results.

PERMANENT PACEMAKERS

Nomenclature and Complications The main therapeutic intervention in SA node dysfunction is permanent pacing. Since the first implementation of permanent pacing in the 1950s, many advances in technology have resulted in miniaturization, increased longevity of pulse generators, improvement in leads, and increased functionality. To better understand pacemaker therapy for bradycardias, it is important to be familiar with the fundamentals of pacemaking. Pacemaker modes and function are named using a five-letter code. The first letter indicates the chamber(s) that is paced (O, none; A, atrium; V, ventricle; D, dual; S, single), the second is the chamber(s) in which sensing occurs (O, none; A, atrium; V, ventricle; D, dual; S, single), the third is the response to a sensed event (O, none; I, inhibition; T, triggered; D, inhibition + triggered), the fourth refers to the programmability or rate response (R, rate responsive), and the fifth refers to the existence of antitachycardia functions if present (O, none; P, antitachycardia pacing; S, shock; D, pace + shock). Almost all modern pacemakers are multiprogrammable and have the capability for rate responsiveness using one of several rate sensors: activity or motion, minute ventilation, or QT interval. The most commonly programmed modes of implanted single- and dual-chamber pacemakers are VVIR and DDDR, respectively, although multiple modes can be programmed in modern pacemakers.

Although pacemakers are highly reliable, they are subject to a number of complications related to implantation and electronic function. In adults, permanent pacemakers are most commonly implanted with access to the heart by way of the subclavian–superior vena cava venous system. Rare, but possible, acute complications of transvenous pacemaker implantation include infection, hematoma, pneumothorax, cardiac perforation, diaphragmatic/phrenic nerve stimulation, and lead dislodgment. Limitations of chronic pacemaker therapy include infection, erosion, lead failure, and abnormalities resulting from inappropriate programming or interaction with the patient's native electrical cardiac function. Rotation of the pacemaker pulse generator in its subcutaneous pocket, either intentionally or inadvertently, often referred to as "twiddler's syndrome," can wrap the leads around the generator and produce dislodgment with failure to sense or pace the heart. The small size and light weight of contemporary pacemakers make this a rare complication.

Complications stemming from chronic cardiac pacing also result from disturbances in atrioventricular synchrony and/or left ventricular mechanical synchrony. Pacing modes that interrupt or fail to restore atrioventricular synchrony may lead to a constellation of signs and symptoms, collectively referred to as pacemaker syndrome, that include neck pulsation, fatigue, palpitations, cough, confusion, exertional dyspnea, dizziness, syncope, elevation in jugular venous pressure, canon A waves, and stigmata of congestive heart failure, including edema, rales, and a third heart sound. Right ventricular apical pacing can induce dyssynchronous activation of the left ventricle, leading to compromised left ventricular systolic function, mitral valve regurgitation, and the previously mentioned stigmata of congestive heart failure. Maintenance of AV synchrony can minimize the sequelae of pacemaker syndrome. Selection of pacing modes that minimize unnecessary ventricular pacing or implantation of a device capable of right and left ventricular pacing (biventricular pacing) can help minimize the deleterious consequences of pacing-induced mechanical dyssynchrony at the ventricular level.

1470 Pacemaker Therapy in SA Node Dysfunction Pacing in SA nodal disease is indicated to alleviate symptoms of bradycardia. Consensus guidelines published by the American Heart Association (AHA)/American College of Cardiology/Heart Rhythm Society (ACC/HRS) outline the indications for the use of pacemakers and categorize them by class based on levels of evidence. Class I conditions are those for which there is evidence or consensus of opinion that therapy is useful and effective. In class II conditions, there is conflicting evidence or a divergence of opinion about the efficacy of a procedure or treatment; in class IIa conditions, the weight of evidence or opinion favors treatment; and in class IIb conditions, efficacy is less well established by the evidence or opinion of experts. In class III conditions, the evidence or weight of opinion indicates that the therapy is not efficacious or useful and may be harmful.

Class I indications for pacing in SA node dysfunction include documented symptomatic bradycardia, sinus node dysfunction–associated long-term drug therapy for which there is no alternative, and symptomatic chronotropic incompetence. Class IIa indications include those outlined previously in which sinus node dysfunction is suspected but not documented and for syncope of unexplained origin in the presence of major abnormalities of SA node dysfunction. Mildly symptomatic individuals with heart rates consistently <40 beats/min constitute a class IIb indication for pacing. Pacing is not indicated in patients with SA node dysfunction who do not have symptoms and in those in whom bradycardia is associated with the use of nonessential drugs (**Table 274-2**).

There is some controversy about the mode of pacing that should be employed in SA node disease. A number of randomized, single-blind trials of pacing mode have been performed. There are no trials that demonstrate an improvement in mortality rate with AV synchronous pacing compared with single-chamber pacing in SA node disease. In some of these studies, the incidence of atrial fibrillation and thromboembolic events was reduced with AV synchronous pacing. In trials of patients with dual-chamber pacemakers designed to compare single-chamber with dual-chamber pacing by crossover design, the need for AV synchronous pacing due to pacemaker syndrome was common. Pacing modes that preserve AV synchrony appear to be associated with a reduction in the incidence of atrial fibrillation and improved quality of life. Because of the low but finite

incidence of AV conduction disease, patients with SA node dysfunction usually undergo dual-chamber pacemaker implantation.

Pacemaker Therapy in Carotid Sinus Hypersensitivity and Vasovagal Syncope Carotid sinus hypersensitivity, if accompanied by a significant cardioinhibitory component, responds well to pacing. In this circumstance, pacing is required only intermittently and single-chamber ventricular pacing is often sufficient. The mechanism of vasovagal syncope is incompletely understood but appears to involve activation of cardiac mechanoreceptors with consequent activation of neural centers that mediate vagal activation and withdrawal of sympathetic nervous system tone. Several randomized clinical trials have been performed in patients with drug-refractory vasovagal syncope, with some studies suggesting reduction in the frequency and the time to recurrent syncope in patients who were paced compared with those who were not. A recent follow-up study to one of those initial trials, however, found less convincing results, casting some doubt on the utility of pacing for vagally mediated syncope.

275 The Bradyarrhythmias: Disorders of the Atrioventricular Node

David D. Spragg, Gordon F. Tomaselli

Impulses generated in the sinoatrial (SA) node or in ectopic atrial loci are conducted to the ventricles through the electrically and anatomically complex atrioventricular (AV) node. As described in **Chap. 274**, the electrophysiologic properties of nodal tissue are distinct from atrial and ventricular myocardium. Cells located in the AV node sit at a relatively higher resting membrane potential than surrounding atrial and ventricular myocytes, exhibit spontaneous depolarization during phase 4 of the action potential, and have slower phase 0 depolarization (mediated by calcium influx in nodal tissue) than that seen in ventricular tissue (mediated by sodium influx).

Bradycardia may occur when conduction across the AV node is compromised, resulting in ineffective ventricular rates, with the possibility of attendant symptoms, including fatigue, syncope, and (if subsidiary pacemaker activity is insufficient) even death. It is important to recognize that in the setting of disturbed AV conduction, SA activation and atrial systole may occur at normal or even accelerated rates, while ventricular activation is either slowed or nonexistent. Transient AV conduction block is common in the young and is most likely the result of high vagal tone found in up to 10% of young adults. Acquired and persistent failure of AV conduction is decidedly rare in healthy adult populations, with an estimated incidence of 200 per million population per year. In the setting of myocardial ischemia, aging and fibrosis, or cardiac infiltrative diseases, however, persistent AV block is much more common.

As with symptomatic bradycardia arising from SA node dysfunction, permanent pacing is the only reliable therapy for symptoms arising from AV conduction block. Approximately 50% of the 150,000 permanent pacemakers implanted in the United States and 70–80% of those in Europe are implanted for disorders of AV conduction.

STRUCTURE AND PHYSIOLOGY OF THE AV NODE

The AV conduction axis is structurally complex, involving the atria and ventricles as well as the AV node. Unlike the SA node, the AV node is a subendocardial structure originating in the transitional zone, which is composed of aggregates of cells in the posterior-inferior right atrium. Superior, medial, and posterior transitional atrionodal bundles converge on the compact AV node. The compact AV node (~1 × 3 × 5 mm) is situated at the apex of the triangle of Koch, which is defined by the coronary sinus ostium posteriorly, the septal tricuspid valve annulus anteriorly, and the tendon of Todaro superiorly. The compact AV node continues as the penetrating AV bundle where it immediately traverses the central fibrous body and is in close proximity to the

TABLE 274-2 SUMMARY OF GUIDELINES FOR PACEMAKER IMPLANTATION IN SA NODE DYSFUNCTION

Class I

1. SA node dysfunction with symptomatic bradycardia or sinus pause
2. Symptomatic SA node dysfunction as a result of essential long-term drug therapy with no acceptable alternatives
3. Symptomatic chronotropic incompetence
4. Atrial fibrillation with bradycardia and pauses >5 s

Class IIa

1. SA node dysfunction with heart rates <40 beats/min without a clear and consistent relationship between bradycardia and symptoms
2. SA node dysfunction with heart rates <40 beats/min on an essential long-term drug therapy with no acceptable alternatives, without a clear and consistent relationship between bradycardia and symptoms
3. Syncope of unknown origin when major abnormalities of SA node dysfunction are discovered or provoked by electrophysiologic testing

Class IIb

1. Mildly symptomatic patients with waking chronic heart rates <40 beats/min

Class III

1. SA node dysfunction in asymptomatic patients, even those with heart rates <40 beats/min
2. SA node dysfunction in which symptoms suggestive of bradycardia are not associated with a slow heart rate
3. SA node dysfunction with symptomatic bradycardia due to nonessential drug therapy

Source: Modified from AE Epstein et al: *J Am Coll Cardiol* 51:e1, 2008 and CM Tracy et al: *J Am Coll Cardiol* 61:e6, 2013.

aortic, mitral, and tricuspid valve annuli; thus, it is subject to injury in the setting of valvular heart disease or its surgical treatment. The penetrating AV bundle continues through the annulus fibrosis and emerges along the ventricular septum adjacent to the membranous septum as the bundle of His. The right bundle branch (RBB) emerges from the distal AV bundle in a band that traverses the right ventricle (moderator band). In contrast, the left bundle branch (LBB) is a broad subendocardial sheet of tissue on the septal left ventricle. The Purkinje fiber network emerges from the RBB and LBB and extensively ramifies on the endocardial surfaces of the right and left ventricles, respectively.

The blood supply to the penetrating AV bundle is from the AV nodal artery and first septal perforator of the left anterior descending coronary artery. The bundle branches also have a dual blood supply from the septal perforators of the left anterior descending coronary artery and branches of the posterior descending coronary artery. The AV node is highly innervated with postganglionic sympathetic and parasympathetic nerves. The bundle of His and distal conducting system are minimally influenced by autonomic tone.

The cells that constitute the AV node complex are heterogeneous with a range of action potential profiles. In the transitional zones, the cells have an electrical phenotype between those of atrial myocytes and cells of the compact node (see Fig. 274-1). Atrionodal transitional connections may exhibit *decremental conduction*, defined as slowing of conduction with increasingly rapid rates of stimulation. Fast and slow AV nodal pathways have been described, but it is controversial whether these two types of pathway are anatomically distinct or represent functional heterogeneities in different regions of the AV nodal complex. Myocytes that constitute the compact node are depolarized (resting membrane potential ~ -60 mV) and exhibit action potentials with low amplitudes, slow upstrokes of phase 0 (<10 V/s), and phase 4 diastolic depolarization; high-input resistance; and relative insensitivity to external $[K^+]$. The action potential phenotype is explained by the complement of ionic currents expressed. AV nodal cells lack a robust inward rectifier potassium current (I_{K1}) and fast sodium current (I_{Na}); L-type calcium current (I_{Ca-L}) is responsible for phase 0; and phase 4 depolarization reflects the composite activity of the depolarizing currents—funny current (I_f), I_{Ca-L} , T-type calcium current (I_{Ca-T}), and sodium calcium exchanger current (I_{NCX})—and the repolarizing currents—delayed rectifier (I_{Kr}) and acetylcholine-gated (I_{KACh}) potassium currents. Electrical coupling between cells in the AV node is tenuous due to the relatively sparse expression of gap junction channels (predominantly connexin-40) and increased extracellular volume.

The His bundle and the bundle branches are insulated from ventricular myocardium. The most rapid conduction in the heart is observed in these tissues. The action potentials exhibit very rapid upstrokes (phase 0), prolonged plateaus (phase 2), and modest automaticity (phase 4 depolarization). Gap junctions, composed largely of connexin-40, are abundant, but bundles are poorly connected transversely to ventricular myocardium.

ETIOLOGY OF AV CONDUCTION DISEASE

Conduction block from the atrium to the ventricle can occur for a variety of reasons in a number of clinical situations, and AV conduction block may be classified in a number of ways. The etiologies may be functional or structural, in part analogous to extrinsic and intrinsic causes of SA nodal dysfunction. The block may be classified by its severity from first to third degree or complete AV block or by the location of block within the AV conduction system. **Table 275-1** summarizes the etiologies of AV conduction block. Those that are functional (autonomic, metabolic/endocrine, and drug-related) tend to be reversible. Most other etiologies produce structural changes, typically fibrosis, in segments of the AV conduction axis that are generally permanent. Heightened vagal tone during sleep or in well-conditioned individuals can be associated with all grades of AV block. Carotid sinus hypersensitivity, vasovagal syncope, and cough and micturition syncope may be associated with SA node slowing and AV conduction block. Transient metabolic and endocrinologic disturbances as well as a number of pharmacologic agents also may produce reversible AV conduction block.

TABLE 275-1 ETIOLOGIES OF ATRIOVENTRICULAR BLOCK

Autonomic	
Carotid sinus hypersensitivity	Vasovagal
Metabolic/Endocrine	
Hyperkalemia	Hypothyroidism
Hypermagnesemia	Adrenal insufficiency
Drug-Related	
Beta blockers	Adenosine
Calcium channel blockers	Antiarrhythmics (class I and III)
Digitalis	Lithium
Infectious	
Endocarditis	Tuberculosis
Lyme disease	Diphtheria
Chagas' disease	Toxoplasmosis
Syphilis	
Heritable/Congenital	
Congenital heart disease	Facioscapulohumeral MD, OMIM #158900 (4q35)
Maternal SLE	
Kearns-Sayre syndrome, OMIM #530000	Emery-Dreifuss MD, OMIM #310300 (Xq28)
Myotonic dystrophy	Progressive familial heart block, type IA OMIM #113900 (3p21)
Type 1, OMIM #160900 (19q13.2-13.3)	Progressive familial heart block, type IB, OMIM #604559 (19q13.32)
Type 2, OMIM #602668 (3q13.3-q24)	Progressive familial heart block, type II, OMIM #140400
Inflammatory	
SLE	MCTD
Rheumatoid arthritis	Scleroderma
Infiltrative	
Amyloidosis	Hemochromatosis
Sarcoidosis	
Neoplastic/Traumatic	
Lymphoma	Radiation
Mesothelioma	Catheter ablation
Melanoma	
Degenerative	
Lev's disease	Lenègre's disease
Coronary Artery Disease	
Acute MI	

Abbreviations: MCTD, mixed connective tissue disease; MI, myocardial infarction; OMIM, Online Mendelian Inheritance in Man (database; designations: #, phenotypic description, molecular basis known; %, phenotypic description); SLE, systemic lupus erythematosus.

Several infectious diseases have a predilection for the conducting system. Lyme disease may involve the heart in up to 50% of cases; 10% of patients with Lyme carditis develop AV conduction block, which is generally reversible but may require temporary pacing support. Chagas' disease, which is common in Latin America, and syphilis may produce more persistent AV conduction disturbances. Some autoimmune and infiltrative diseases may produce AV conduction block, including systemic lupus erythematosus (SLE), rheumatoid arthritis, mixed connective tissue disease, scleroderma, amyloidosis (primary and secondary), sarcoidosis, and hemochromatosis; rare malignancies also may impair AV conduction.

Idiopathic progressive fibrosis of the conduction system is one of the more common and degenerative causes of AV conduction block. Aging is associated with degenerative changes in the summit of the ventricular septum, central fibrous body, and aortic and mitral annuli and has been described as "sclerosis of the left cardiac skeleton." The process typically begins in the fourth decade of life and may be accelerated by atherosclerosis, hypertension, and diabetes mellitus. Accelerated forms

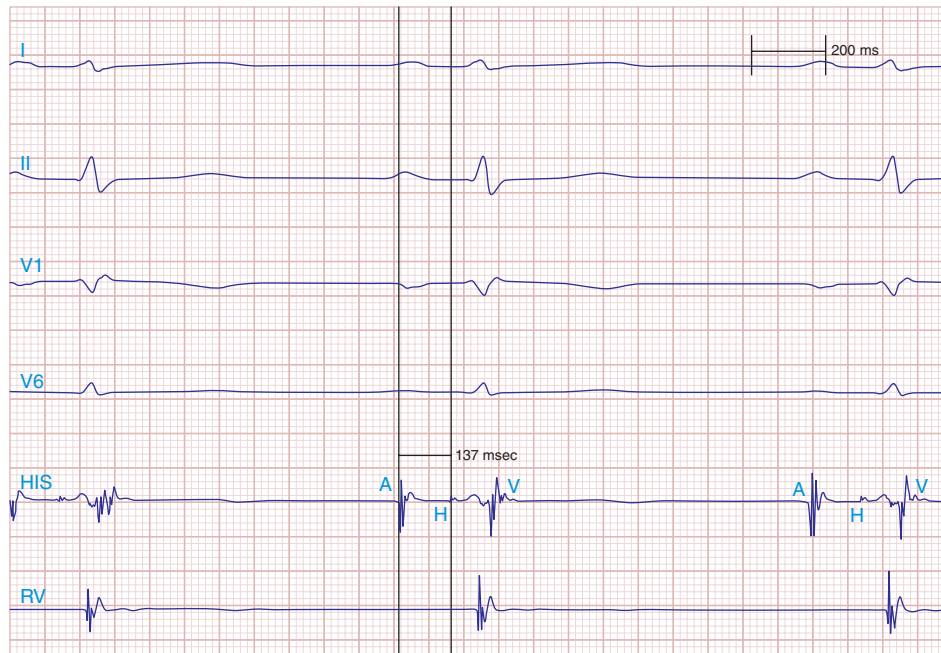


FIGURE 275-1 First-degree AV block with slowing of conduction in the AV node as indicated by the prolonged atrial-to-His bundle electrogram (AH) interval, in this case 157 ms. The His bundle-to-earliest ventricular activation on the surface ECG (HV) interval is normal. The normal HV interval suggests normal conduction below the AV node to the ventricle. I and V₁ are surface ECG leads, and HIS is the recording of the endocavitary electrogram at the His bundle position. A, H, and V are labels for the atrial, His bundle, and right ventricular electrograms, respectively.

of progressive familial heart block have been identified in families with mutations in the cardiac sodium channel gene (*SCN5A*) and other loci that have been mapped to chromosomes 1 and 19.

AV conduction block has been associated with heritable neuromuscular diseases, including the nucleotide repeat disease myotonic dystrophy, the mitochondrial myopathy Kearns-Sayre syndrome (Chap. 462e), and several of the monogenic muscular dystrophies. Congenital AV block may be observed in complex congenital cardiac anomalies (Chap. 282), such as transposition of the great arteries, ostium primum atrial septal defects (ASDs), ventricular septal defects (VSDs), endocardial cushion defects, and some single-ventricle defects. Congenital AV block in the setting of a structurally normal heart has been seen in children born to mothers with SLE. Iatrogenic AV block may occur during mitral or aortic valve surgery, rarely in the setting of thoracic radiation, and as a consequence of catheter ablation. AV block is a decidedly rare complication of the surgical repair of VSDs or ASDs but may complicate repairs of transposition of the great arteries.

Coronary artery disease may produce transient or persistent AV block. In the setting of coronary spasm, ischemia, particularly in the right coronary artery distribution, may produce transient AV block. In acute myocardial infarction (MI), AV block transiently develops in 10–25% of patients; most commonly, this is first- or second-degree AV block, but complete heart block (CHB) may also occur. Second-degree and higher-grade AV block tends to occur more often in inferior than in anterior acute MI; however, the level of block in inferior MI tends to be in the AV node with more stable, narrow escape rhythms. In contrast, acute anterior MI is associated with block in the distal AV

nodal complex, His bundle, or bundle branches and results in wide complex, unstable escape rhythms and a worse prognosis with high mortality rates.

ELECTROCARDIOGRAPHY AND ELECTROPHYSIOLOGY OF AV CONDUCTION BLOCK

AV conduction block typically is diagnosed electrocardiographically, which characterizes the severity of the conduction disturbance and allows one to draw inferences about the location of the block. AV conduction block manifests as slow conduction in its mildest forms and failure to conduct, either intermittent or persistently, in more severe varieties. First-degree AV block (PR interval >200 ms) is a slowing of conduction through the AV junction (Fig. 275-1). The site of delay is typically in the AV node but may be in the atria, bundle of His, or His-Purkinje system. A wide QRS is suggestive of delay in the distal conduction system, whereas a narrow QRS suggests delay in the AV node proper or, less commonly, in the bundle of His. In second-degree AV block there is an intermittent failure of electrical impulse conduction from atrium to ventricle. Second-degree AV block is subclassified as Mobitz type I (Wenckebach) or Mobitz type II. The periodic failure of conduction in Mobitz type I block is characterized by a progressively lengthening PR interval, shortening of the RR interval, and a pause that is less than two times the immediately preceding RR interval on the electrocardiogram (ECG). The ECG complex after the pause exhibits a shorter PR interval than that immediately preceding the pause (Fig. 275-2). This ECG pattern most often arises because of decremental conduction of electrical impulses in the AV node.



FIGURE 275-2 Mobitz type I second-degree AV block. The PR interval prolongs before the pause, as shown in the ladder diagram. The ECG pattern results from slowing of conduction in the AV node.

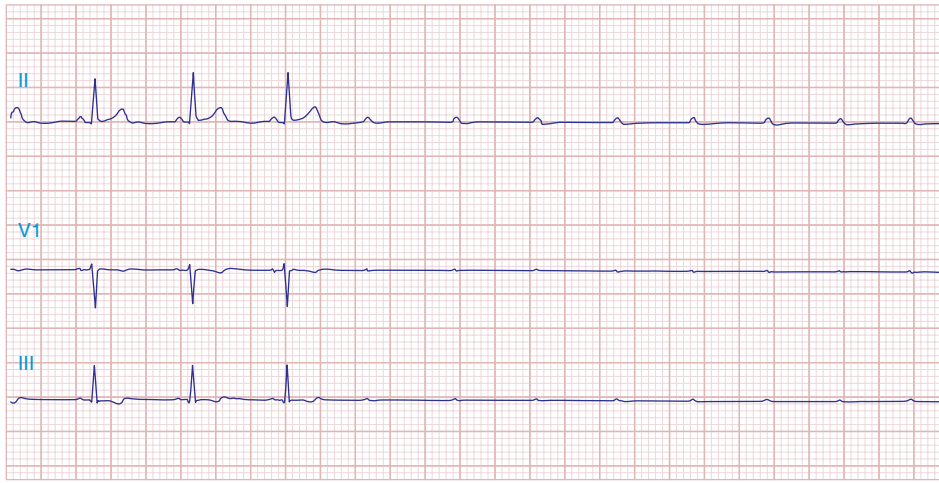


FIGURE 275-3 Paroxysmal AV block. Multiple nonconducted P waves after a period of sinus bradycardia with a normal PR interval. This implies significant conduction system disease, requiring permanent pacemaker implantation.

It is important to distinguish type I from type II second-degree AV nodal block because the latter has more serious prognostic implications. Type II second-degree AV block is characterized by intermittent failure of conduction of the P wave without changes in the preceding PR or RR intervals. When AV block is 2:1, it may be difficult to distinguish type I from type II block. Type II second-degree AV block typically occurs in the distal or infra-His conduction system, is often associated with intraventricular conduction delays (e.g., bundle branch block), and is more likely to proceed to higher grades of AV block than is type I second-degree AV block. Second-degree AV block (particularly type II) may be associated with a series of nonconducted P waves, referred to as *paroxysmal AV block* (Fig. 275-3), and implies significant conduction system disease and is an indication for permanent pacing. Complete failure of conduction from atrium to ventricle is referred to as complete or third-degree AV block. AV block that is intermediate between second degree and third degree is referred to as high-grade AV block and, as with CHB, implies advanced AV conduction system disease. In both cases, the block is most often distal to the AV node, and the duration of the QRS complex can be helpful in determining the level of the block. In the absence of a preexisting bundle branch block, a wide QRS escape rhythm (Fig. 275-4B) implies a block in the distal His or bundle branches; in contrast, a narrow QRS rhythm implies a block in the AV node or proximal His and an escape rhythm originating in the AV junction (Fig. 275-4A). Narrow QRS escape rhythms are typically faster and more stable than wide QRS escape rhythms and originate more proximally in the AV conduction system.

DIAGNOSTIC TESTING

Diagnostic testing in the evaluation of AV block is aimed at determining the level of conduction block, particularly in asymptomatic

patients, since the prognosis and therapy depend on whether the block is in or below the AV node. Vagal maneuvers, carotid sinus massage, exercise, and administration of drugs such as atropine and isoproterenol may be diagnostically informative. Owing to the differences in the innervation of the AV node and infranodal conduction system, vagal stimulation and carotid sinus massage slow conduction in the AV node but have less of an effect on infranodal tissue and may even improve conduction due to a reduced rate of activation of distal tissues. Conversely, atropine, isoproterenol, and exercise improve conduction through the AV node and impair infranodal conduction. In patients with congenital CHB and a narrow QRS complex, exercise typically increases heart rate; by contrast, those with acquired CHB, particularly with wide QRS, do not respond to exercise with an increase in heart rate.

Additional diagnostic evaluation, including electrophysiologic testing, may be indicated in patients with syncope and suspected high-grade AV block. This is particularly relevant if noninvasive testing does not reveal the cause of syncope or if the patient has structural heart disease with ventricular tachyarrhythmias as a cause of symptoms. Electrophysiologic testing provides more precise information regarding the location of AV conduction block and permits studies of AV conduction under conditions of pharmacologic stress and exercise. Recording of the His bundle electrogram by a catheter positioned at the superior margin of the tricuspid valve annulus provides information about conduction at all levels of the AV conduction axis. A properly recorded His bundle electrogram reveals local atrial activity, the His electrogram, and local ventricular activation; when it is monitored simultaneously with recorded body surface electrocardiographic traces, intraatrial, AV nodal, and infranodal conduction times can be assessed (Fig. 275-1). The time from the most rapid deflection of the

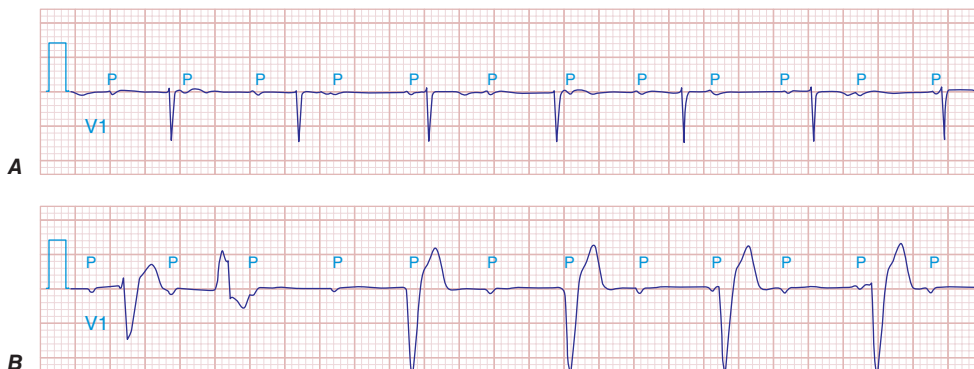


FIGURE 275-4 High-grade AV block. A. Multiple nonconducted P waves with a regular narrow complex QRS escape rhythm probably emanating from the AV junction. **B.** A wide complex QRS escape and a single premature ventricular contraction. In both cases, there is no consistent temporal relationship between the P waves and QRS complexes.

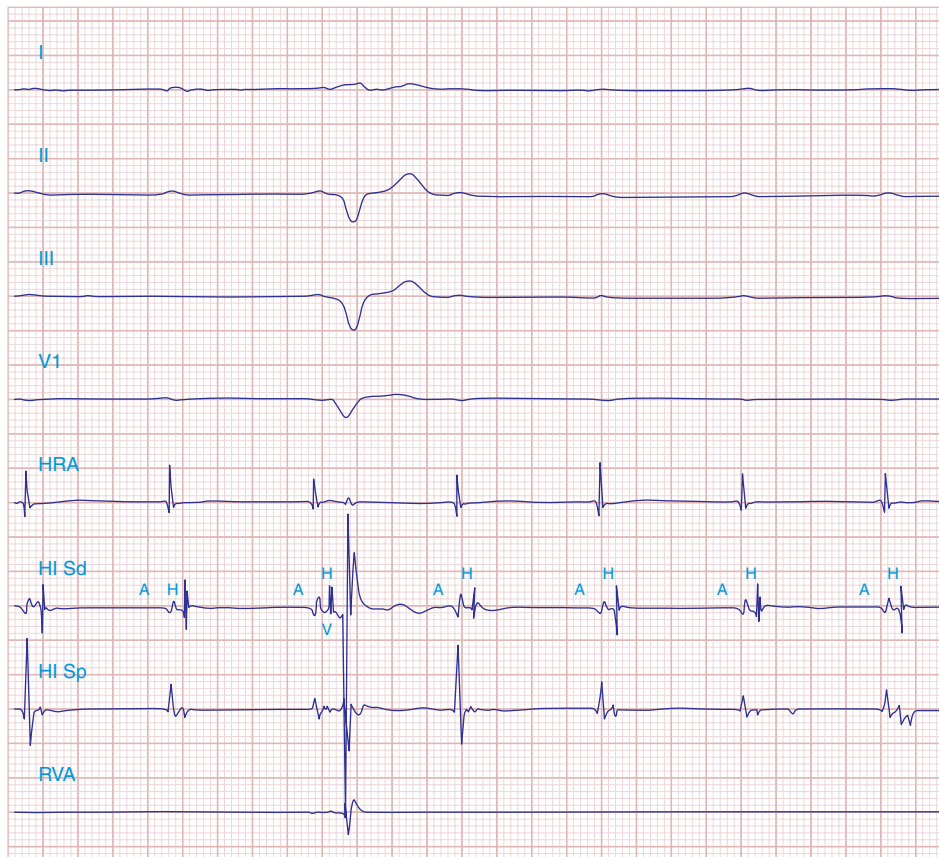


FIGURE 275-5 High-grade AV block below the His. The AH interval is normal and is not changing before the block. Atrial and His bundle electrograms are recorded consistent with block below the distal AV junction. I, II, III, and V₁ are surface ECG leads. HiSp, HiSd, and RVA are the proximal His, distal His, and right ventricular apical electrical recordings, respectively. A, H, and V represent the atrial, His, and ventricular electrograms on the His bundle recording, respectively. (Tracing courtesy of Dr. Joseph Marine; with permission.)

atrial electrogram in the His bundle recording to the His electrogram (*AH interval*) represents conduction through the AV node and is normally <130 ms. The time from the His electrogram to the earliest onset of the QRS on the surface ECG (*HV interval*) represents the conduction time through the His-Purkinje system and is normally ≤55 ms.

Rate stress produced by pacing can unveil abnormal AV conduction. Mobitz I second-degree AV block at short atrial paced cycle lengths is a normal response. However, when it occurs at atrial cycle lengths >500 ms (<120 beats/min) in the absence of high vagal tone, it is abnormal. Typically, type I second-degree AV block is associated with prolongation of the AH interval, representing conduction slowing and block in the AV node. AH prolongation occasionally is due to the effect of drugs (beta blockers, calcium channel blockers, digitalis) or increased vagal tone. Atropine can be used to reverse high vagal tone; however, if AH prolongation and AV block at long pacing cycle lengths persists, intrinsic AV node disease is likely. Type II second-degree block is typically infranodal, often in the His-Purkinje system. Block below the node with prolongation of the HV interval or a His bundle electrogram with no ventricular activation (**Fig. 275-5**) is abnormal unless it is elicited at fast pacing rates or short coupling intervals with extra stimulation. It is often difficult to determine the type of second-degree AV block when 2:1 conduction is present; however, the finding of a His bundle electrogram after every atrial electrogram indicates that block is occurring in the distal conduction system.

Intracardiac recording at electrophysiologic study that reveals prolongation of conduction through the His-Purkinje system (i.e., long HV interval) is associated with an increased risk of progression to higher grades of block and is generally an indication for pacing. In the setting of bundle branch block, the HV interval may reveal the condition of

the unblocked bundle and the prognosis for developing more advanced AV conduction block. Prolongation of the HV interval in patients with asymptomatic bundle branch block is associated with an increased risk of developing higher-grade AV block. The risk increases with greater prolongation of the HV interval such that in patients with an HV interval >100 ms, the annual incidence of complete AV block approaches 10%, indicating a need for pacing. In patients with acquired CHB, even if intermittent, there is little role for electrophysiologic testing, and pacemaker implantation is almost always indicated.

TREATMENT MANAGEMENT OF AV CONDUCTION BLOCK

Temporary or permanent artificial pacing is the most reliable treatment for patients with symptomatic AV conduction system disease. However, exclusion of reversible causes of AV block and the need for temporary heart rate support based on the hemodynamic condition of the patient are essential considerations in each patient. Correction of electrolyte derangements and ischemia, inhibition of excessive vagal tone, and withholding of drugs with AV nodal blocking properties may increase the heart rate. Adjunctive pharmacologic treatment with atropine or isoproterenol may be useful if the block is in the AV node. Since most pharmacologic treatment may take some time to initiate and become effective, temporary pacing may be necessary. The most expeditious technique is the use of transcutaneous pacing, where pacing patches are placed anteriorly over the cardiac apex (cathode) and posteriorly between the spine and the scapula or above the right nipple (anode). Acutely, transcutaneous pacing is highly effective, but its duration is limited by patient discomfort and longer-term failure to capture the ventricle owing to changes in lead impedance. If a patient requires more

TABLE 275-2 GUIDELINE SUMMARY FOR PACEMAKER IMPLANTATION IN ACQUIRED AV BLOCK**Class I**

1. Third-degree or high-grade AV block at any anatomic level associated with:
 - a. Symptomatic bradycardia
 - b. Essential drug therapy that produces symptomatic bradycardia
 - c. Periods of asystole >3 s or any escape rate <40 beats/min while awake, or an escape rhythm originating below the AV node
 - d. Postoperative AV block not expected to resolve
 - e. Catheter ablation of the AV junction
 - f. Neuromuscular diseases such as myotonic dystrophy, Kearns-Sayre syndrome, Erb dystrophy, and peroneal muscular atrophy, regardless of the presence of symptoms
2. Second-degree AV block with symptomatic bradycardia
3. Type II second-degree AV block with a wide QRS complex with or without symptoms
4. Exercise-induced second- or third-degree AV block in the absence of ischemia
5. Atrial fibrillation with bradycardia and pauses >5 s

Class IIa

1. Asymptomatic third-degree AV block regardless of level
2. Asymptomatic type II second-degree AV block with a narrow QRS complex
3. Asymptomatic type II second-degree AV block with block within or below the His at electrophysiologic study
4. First- or second-degree AV block with symptoms similar to pacemaker syndrome

Class IIb

1. AV block in the setting of drug use/toxicity, when the block is expected to recur even with drug discontinuation
2. Neuromuscular diseases such as myotonic dystrophy, Kearns-Sayre syndrome, Erb dystrophy, and peroneal muscular atrophy with any degree of AV block regardless of the presence of symptoms

Class III

1. Asymptomatic first-degree AV block
2. Asymptomatic type I second-degree AV block at the AV node level
3. AV block that is expected to resolve or is unlikely to recur (Lyme disease, drug toxicity)

Source: Modified from AE Epstein et al: *J Am Coll Cardiol* 51:e1, 2008.

than a few minutes of pacemaker support, transvenous temporary pacing should be instituted. Temporary pacing leads can be placed from the jugular or subclavian venous system and advanced to the right ventricle, permitting stable temporary pacing for many days, if necessary. In most circumstances, in the absence of prompt resolution, conduction block distal to the AV node requires permanent pacemaking.

PACEMAKERS IN AV CONDUCTION DISEASE

There are no randomized trials that evaluate the efficacy of pacing in patients with AV block, as there are no reliable therapeutic alternatives for AV block and untreated high-grade AV block is potentially lethal. The consensus guidelines for pacing in acquired AV conduction block in adults provide a general outline for situations in which pacing is indicated (Table 275-2). Pacemaker implantation should be performed in any patient with symptomatic bradycardia and irreversible second- or third-degree AV block, regardless of the cause or level of block in the conducting system. Symptoms may include those directly related to bradycardia and low cardiac output or to worsening heart failure, angina, or intolerance to an essential medication. Pacing in patients with asymptomatic AV block should be individualized; situations in which pacing should be considered are patients with acquired CHB, particularly in the setting of cardiac enlargement; left ventricular dysfunction; and waking heart rates ≤ 40 beats/min. Patients who have asymptomatic second-degree AV block of either type should be considered for pacing if the block is demonstrated to be intra- or infra-His or is associated with a wide QRS complex. Pacing may be indicated in asymptomatic patients in special circumstances, in patients with profound first-degree AV block and left ventricular dysfunction in whom a shorter AV interval produces hemodynamic improvement, and in the setting of milder forms of AV conduction delay (first-degree AV block, intraventricular conduction delay) in patients with neuromuscular diseases that have

a predilection for the conduction system, such as myotonic dystrophy and other muscular dystrophies, and Kearns-Sayre syndrome.

PACEMAKER THERAPY IN MYOCARDIAL INFARCTION

AV block in acute MI is often transient, particularly in inferior infarction. The circumstances in which pacing is indicated in acute MI are persistent second- or third-degree AV block, particularly if symptomatic, and transient second- or third-degree AV block associated with bundle branch block (Table 275-3). Pacing is generally not indicated in the setting of transient AV block in the absence of intraventricular conduction delays or in the presence of fascicular block

TABLE 275-3 GUIDELINE SUMMARY FOR PACEMAKER IMPLANTATION IN AV CONDUCTION BLOCK IN ACUTE MYOCARDIAL INFARCTION (AMI)**Class I**

1. Persistent second-degree AV block in the His-Purkinje system with bilateral bundle branch block or third-degree block within or below the His after AMI
2. Transient advanced (second- or third-degree) infranodal AV block and associated bundle branch block. If the site of block is uncertain, an electrophysiologic study may be necessary
3. Persistent and symptomatic second- or third-degree AV block

Class IIb

1. Persistent second- or third-degree AV block at the AV node level

Class III

1. Transient AV block in the absence of intraventricular conduction defects
2. Transient AV block in the presence of isolated left anterior fascicular block
3. Acquired left anterior fascicular block in the absence of AV block
4. Persistent first-degree AV block in the presence of bundle branch block that is old or age-indeterminate

Source: Modified from AE Epstein et al: *J Am Coll Cardiol* 51:e1, 2008.

TABLE 275-4 INDICATIONS FOR PACEMAKER IMPLANTATION IN CHRONIC BIFASCICULAR AND TRIFASCICULAR BLOCK**Class I**

1. Intermittent third-degree AV block
2. Type II second-degree AV block
3. Alternating bundle branch block

Class IIa

1. Syncope not demonstrated to be due to AV block when other likely causes (e.g., ventricular tachycardia) have been excluded
2. Incidental finding at electrophysiologic study of a markedly prolonged HV interval (>100 ms) in asymptomatic patients
3. Incidental finding at electrophysiologic study of pacing-induced infra-His block that is not physiologic

Class IIb

1. Neuromuscular diseases such as myotonic dystrophy, Kearns-Sayre syndrome, Erb dystrophy, and peroneal muscular atrophy with any degree of fascicular block regardless of the presence of symptoms, because there may be unpredictable progression of AV conduction disease

Class III

1. Fascicular block without AV block or symptoms
2. Fascicular block with first-degree AV block without symptoms

Source: Modified from AE Epstein et al: *J Am Coll Cardiol* 51:e1, 2008.

or first-degree AV block that develops in the setting of preexisting bundle branch block. Fascicular blocks that develop in acute MI in the absence of other forms of AV block also do not require pacing (Table 275-3 and Table 275-4).

PACEMAKER THERAPY IN BIFASCICULAR AND TRIFASCICULAR BLOCK

Distal forms of AV conduction block may require pacemaker implantation in certain clinical settings. Patients with bifascicular or trifascicular block and symptoms, particularly syncope that is not attributable to other causes, should undergo pacemaker implantation. Pacemaking is indicated in asymptomatic patients with bifascicular or trifascicular block who experience intermittent third-degree, type II second-degree AV block or alternating bundle branch block. In patients with fascicular block who are undergoing electrophysiologic study, a markedly prolonged HV interval or block below the His at long cycle lengths also may constitute an indication for permanent pacing. Patients with fascicular block and the neuromuscular diseases previously described should also undergo pacemaker implantation (Table 275-4).

SELECTION OF PACING MODE

In general, a pacing mode that maintains AV synchrony reduces complications of pacing such as pacemaker syndrome and pacemaker-mediated tachycardia. This is particularly true in younger patients; the importance of dual-chamber pacing in the elderly, however, is not well established. Several studies have failed to demonstrate a difference in mortality rate in older patients with AV block treated with a single-(VVI) compared with a dual-(DDD) chamber pacing mode. In some of the studies that randomized pacing mode, the risk of chronic atrial fibrillation and stroke risk decreased with physiologic pacing. In patients with sinus rhythm and AV block, the very modest increase in risk with dual-chamber pacemaker implantation appears to be justified to avoid the possible complications of single-chamber pacing.

276 Supraventricular Tachyarrhythmias

Gregory F. Michaud, William G. Stevenson

Supraventricular tachyarrhythmias originate from or are dependent on conduction through the atrium or atrioventricular (AV) node to the ventricles. Most produce narrow QRS-complex tachycardia (QRS duration <120 ms) characteristic of ventricular activation over the Purkinje system. Conduction block in the left or right bundle branch or activation of the ventricles from an accessory pathway produces a wide QRS complex during supraventricular tachycardia that must be distinguished from ventricular tachycardia (Chap. 277). Supraventricular tachyarrhythmia may be divided into physiologic sinus tachycardia and pathologic tachycardia (Table 276-1). The prognosis and treatment vary considerably depending on the mechanism and underlying heart disease.

Supraventricular tachycardia can be of brief duration, termed *nonsustained*, or can be sustained such that an intervention, such as cardioversion or drug administration, is required for termination. Episodes that occur with sudden onset and termination are referred to as paroxysmal. *Paroxysmal supraventricular tachycardia* (PSVT) refers to a family of tachycardias including AV node reentry, AV reentry using an accessory pathway, and atrial tachycardia.

CLINICAL PRESENTATION

Symptoms of supraventricular arrhythmia vary depending on the rate, duration, associated heart disease, and comorbidities and include palpitations, chest pain, dyspnea, diminished exertional capacity, and occasionally syncope. Rarely, a supraventricular arrhythmia precipitates cardiac arrest in patients with the Wolff-Parkinson-White syndrome or severe heart disease, such as hypertrophic cardiomyopathy.

Diagnosis requires obtaining an electrocardiogram (ECG) at the time of symptoms. For transient arrhythmia, ambulatory ECG recording is warranted (see Table 277-1). Exercise testing is useful for assessing exercise-related symptoms. Occasionally an invasive electrophysiology study is warranted to provoke the arrhythmia with pacing, confirm the mechanism, and often, perform catheter ablation.

Physiologic Sinus Tachycardia The sinus node is comprised of a group of cells dispersed within the superior aspect of the thick ridge of muscle known as the crista terminalis where the posterior smooth atrial wall derived from the sinus venosus meets the trabeculated anterior portion of the right atrium (Fig. 276-1). Sinus p waves are characterized by a frontal plane axis directed inferiorly and leftward, with positive p waves in leads II, III, and aVF; a negative p wave in aVR; and an initially positive biphasic p wave in V1. Normal sinus rhythm has a rate of 60–100 beats/min. Sinus tachycardia (>100 beats/min) typically occurs in response to sympathetic stimulation and vagal withdrawal, whereby the rate of spontaneous depolarization of the sinus node increases and the focus of earliest activation within the node typically shifts more leftward and closer to the superior septal aspect of the crista terminalis, thus producing taller p waves in the inferior limb leads when compared to normal sinus rhythm.

Sinus tachycardia is considered physiologic when it is an appropriate response to exercise, stress, or illness. Sinus tachycardia can be difficult to distinguish from focal atrial tachycardia (see below) that originates from a focus near the sinus node. A causative factor (such as exertion) and a gradual increase and decrease in rate favors sinus tachycardia, whereas an abrupt onset and offset favor atrial tachycardia. The distinction can be difficult and occasionally requires extended ECG monitoring or even invasive electrophysiology study. Treatment for physiologic sinus tachycardia is aimed at the underlying condition (Table 276-2).

Nonphysiologic Sinus Tachycardia *Inappropriate sinus tachycardia* is an uncommon condition in which the sinus rate increases spontaneously at rest or out of proportion to physiologic stress or exertion.

TABLE 276-1 SUPRAVENTRICULAR TACHYCARDIA**I. Physiologic sinus tachycardia**

Defining feature: normal sinus mechanism precipitated by exertion, stress, concurrent illness (Table 276-2)

II. Pathologic supraventricular tachycardia**A. Tachycardia originating from the atrium**

Defining feature: tachycardia may continue despite beats that fail to conduct to the ventricles, indicating that the AV node is not participating in the tachycardia circuit

1. Inappropriate sinus tachycardia

Defining feature: tachycardia from the normal sinus node area that occurs without an identifiable precipitating factor as a result of dysfunctional autonomic regulation

2. Focal atrial tachycardia

Defining feature: Regular atrial tachycardia with defined p wave; may be sustained, nonsustained, paroxysmal, or incessant. Frequent sites of origin occur along the valve annuli of left or right atrium, pulmonary veins, coronary sinus musculature, superior vena cava

3. Atrial flutter – macroreentrant atrial tachycardia

Defining feature: organized reentry creates organized atrial activity, commonly seen as sawtooth flutter waves at rates typically faster than 200 beats/min

a. Common atrial flutter

i. Right atrial reentry parallel to the tricuspid annulus and dependent on conduction through the isthmus between the inferior vena cava and tricuspid annulus

1. Counterclockwise (as viewed from the ventricular aspect)

2. Clockwise

b. Atypical atrial flutter

i. Usually due to reentry in left or right atrium associated with scars usually from prior surgery or catheter ablation for atrial fibrillation, but may be idiopathic

4. Atrial fibrillation

Defining feature: chaotic rapid atrial electrical activity with variable ventricular rate; the most common sustained cardiac arrhythmia in older adults

5. Multifocal atrial tachycardia

Defining feature: multiple discrete p waves often seen in patients with pulmonary disease during acute exacerbations of pulmonary insufficiency

B. AV nodal reentry tachycardia

Defining feature: paroxysmal regular tachycardia with P waves visible at the end of the QRS complex or not visible at all; the most common paroxysmal sustained tachycardia in healthy young adults; more common in women

C. Tachycardias associated with accessory atrioventricular pathways

a. Orthodromic AV reentry tachycardia

Defining feature: paroxysmal sustained tachycardia similar to AV nodal reentry; during sinus rhythm, evidence of ventricular preexcitation may be present (Wolff-Parkinson-White syndrome) or absent (concealed accessory pathway)

b. Preexcited tachycardia

Defining feature: wide QRS tachycardia with QRS morphology similar to VT

1. Antidromic AV reentry – regular paroxysmal tachycardia
2. Atrial fibrillation with preexcitation – irregular wide complex, or intermittently wide complex tachycardia, some with dangerously rapid rates faster than 250/min
3. Atrial tachycardia or flutter with preexcitation

Abbreviations: AV, atrioventricular; VT, ventricular tachycardia.

Affected individuals are often women in the third or fourth decade of life. Fatigue, dizziness, and even syncope may accompany palpitations, which can be disabling. Additional symptoms of chest pain, headaches, and gastrointestinal upset are common. It must be distinguished from appropriate sinus tachycardia and from focal atrial tachycardia, as discussed above. Misdiagnosis of physiologic sinus tachycardia with an anxiety disorder is common. Therapy is often ineffective or poorly tolerated. Careful titration of beta blockers and/

or calcium channel blockers may reduce symptoms. Clonidine and serotonin reuptake inhibitors have also been used. Ivabradine, a drug that blocks the I_f current causing sinus node depolarization, is promising but is not approved for use in the United States. Catheter ablation of the sinus node has been used, but long-term control of symptoms is usually poor, and it often leaves young individuals with a permanent pacemaker.

When symptomatic sinus tachycardia occurs with postural hypotension, the syndrome is called *postural orthostatic tachycardia syndrome* (POTS). Symptoms are often similar to those in patients with inappropriate sinus tachycardia. POTS is sometimes due to autonomic dysfunction following a viral illness and may resolve spontaneously over 3–12 months. Volume expansion with salt supplementation, oral fludrocortisone, compression stockings, and the α -agonist midodrine, often in combination, can be helpful. Exercise training has also been purported to improve symptoms.

Focal Atrial Tachycardia Focal atrial tachycardia (AT) can be due to abnormal automaticity, triggered automaticity, or a small reentry circuit confined to the atrium or atrial tissue extending into a pulmonary vein, the coronary sinus, or vena cava. It can be sustained, nonsustained, paroxysmal, or incessant. Focal AT accounts for approximately 10% of PSVT referred for catheter ablation. Nonsustained AT is commonly observed on 24-h ambulatory ECG recordings, and the prevalence increases with age. Tachycardia can occur in the absence of structural heart disease or can be associated with any form of heart disease that affects the atrium. Sympathetic stimulation is a promoting factor such that AT can be a sign of underlying illness. AT with AV block can occur in digitalis toxicity. Symptoms are similar to other supraventricular tachycardias (SVTs). Incessant AT can cause tachycardia-induced cardiomyopathy.

AT typically presents as an SVT either with 1:1 AV conduction or with AV block that can be Wenckebach type conduction or fixed (e.g., 2:1 or 3:1) block. Because it is not dependent on AV nodal conduction, AT will not terminate with AV block, and the atrial rate will not be affected, which distinguishes AT from most AV nodal-dependent SVTs, such as AV nodal reentry and AV reentry using an accessory pathway (see below). An accelerated warm-up phase after initiation or cool-down phase prior to termination also favors AT rather than AV nodal-dependent SVT. P waves are often discrete, with an intervening isoelectric segment, in contrast to atrial flutter and macroreentrant AT (see below). When 1:1 conduction to the ventricles is present, the arrhythmia can resemble sinus tachycardia typically with a P-R interval shorter than the R-P interval (Fig. 276-2). It can be distinguished from sinus tachycardia by the p-wave morphology, which usually differs from sinus p waves depending on the location of the focus. Focal AT tends to originate in areas of complex atrial anatomy, such as the crista terminalis, valve annuli, atrial septum, and atrial muscle extending along cardiac thoracic veins (superior vena cava, coronary sinus, and pulmonary veins) (Fig. 276-3), and the location can often be estimated by the P-wave morphology. AT from the right atrium has a positive P-wave morphology in lead I and biphasic P-wave morphology in lead V_1 . AT from the atrial septum will frequently have a narrower P-wave duration than sinus rhythm. AT from the left atrium will usually have a monophasic, positive P wave in lead V_1 . AT that originates from superior atrial locations, such as the superior vena cava or superior pulmonary veins, will be positive in the inferior limb leads II, III, and aVF, whereas AT from a more inferior location, such as the ostium of the coronary sinus, will inscribe negative P waves in these same leads. When the focus is in the superior aspect of the crista terminalis, close to the sinus node, however, the p wave will resemble that of sinus tachycardia. Abrupt onset and offset then favor AT rather than sinus tachycardia. Depending on the atrial rate, the P wave may fall on top of the t wave or, during 2:1 conduction, may fall coincident with the QRS. Maneuvers that increase AV block, such as carotid sinus massage, Valsalva maneuver, or administration of AV nodal-blocking agents, such as adenosine, are useful to create AV block that will expose the p wave (Fig. 276-4).

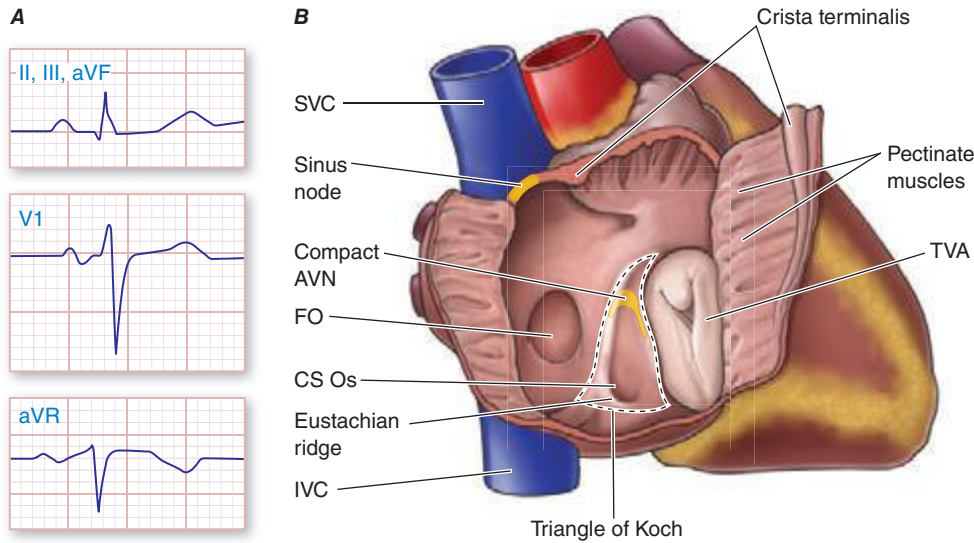


FIGURE 276-1 Right atrial anatomy pertinent to normal sinus rhythm and supraventricular tachycardia. **A.** Typical P-wave morphology during normal sinus rhythm based on standard 12-lead electrocardiogram. There is a positive P wave in leads II, III, and aVF; biphasic, initially positive P wave in V₁; and negative P wave in aVR. **B.** Right atrial anatomy seen from a right lateral perspective with the lateral wall opened to view the septum. AVN, atrioventricular node; CS Os, coronary sinus ostium; FO, fossa ovalis; IVC, inferior vena cava; SVC, superior vena cava; TVA, tricuspid valve annulus.

TABLE 276-2 COMMON CAUSES OF PHYSIOLOGIC SINUS TACHYCARDIA

1. Exercise
2. Acute illness with fever, infection, pain
3. Hypovolemia, anemia
4. Hyperthyroidism
5. Pulmonary insufficiency
6. Drugs that have sympathomimetic, vagolytic, or vasodilator properties, e.g., albuterol, theophylline, tricyclic antidepressants, nifedipine, hydralazine
7. Pheochromocytoma

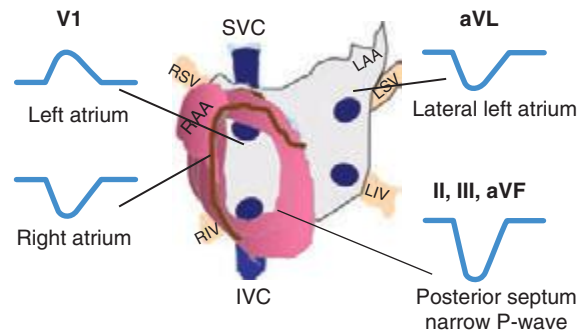


FIGURE 276-3 Location of focal atrial tachycardia focus estimated by P-wave morphology. LAA, left atrial appendage; LIV, left inferior pulmonary vein; LSV, left superior pulmonary vein; RAA, right atrial appendage; RIV, right inferior pulmonary vein; RSV, right superior pulmonary vein; SVC, superior vena cava.

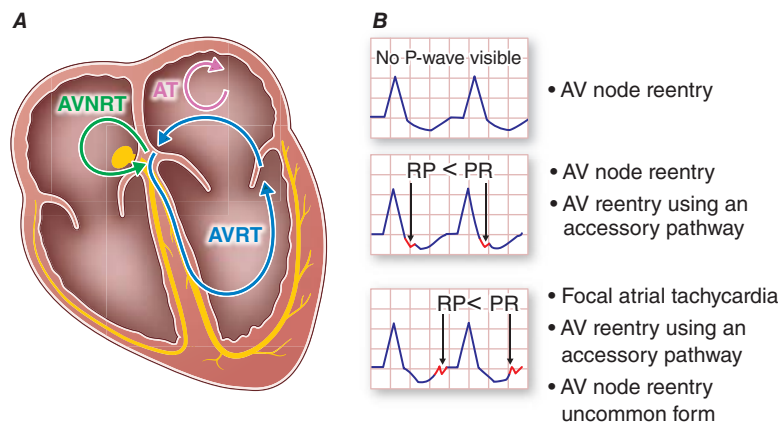
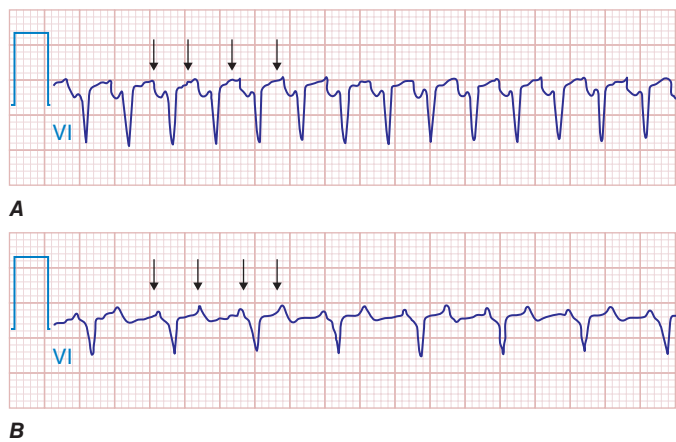


FIGURE 276-2 Common mechanisms underlying paroxysmal supraventricular tachycardia along with typical R-P relationships. **A.** Schematic showing a four-chamber view of the heart with atrioventricular node in green and an accessory pathway between the left atrium and left ventricle in yellow. Atrial tachycardia (AT; red circuit) is confined completely to atrial tissue. Atrioventricular nodal reentry tachycardia (AVNRT; blue circuit) uses atrioventricular (AV) nodal and perinodal atrial tissue. Atrioventricular reentry tachycardia (AVRT; black circuit) uses atrial and ventricular tissue, accessory pathway, AV node, and specialized conduction fibers (His-Purkinje) as part of the reentry circuit. **B.** Typical relation of the p wave to QRS, commonly described as the R-P to P-R relationships for the different tachycardia mechanisms.



B

FIGURE 276-4 Atrial tachycardia (AT) with 1:1 and 2:1 atrioventricular (AV) conduction. Arrows indicate p waves. **A.** AT with 1:1 AV relationship and R-P > P-R. **B.** Same AT with 2:1 AV relationship after AV nodal-blocking agent administered. (Adapted from F Marchlinski: *The tachyarrhythmias*. In Longo DL et al [eds]: *Harrison's Principles of Internal Medicine*, 18th ed. New York, McGraw-Hill, 2012, pp 1878–1900.)

Acute management of sudden-onset, sustained AT is the same as for PSVT (see below), but the response to pharmacologic therapy is variable, likely depending on the mechanism. For AT due to reentry, administration of adenosine or vagal maneuvers may transiently increase AV block without terminating tachycardia. Some ATs terminate with a sufficient dose of adenosine, consistent with triggered activity as the mechanism. Cardioversion can be effective in some, but fails in others, suggesting automaticity as the mechanism. Beta blockers and calcium channel blockers may slow the ventricular rate by increasing AV block, which can improve tolerance of the arrhythmias. Potential precipitating factors and intercurrent illness should be sought and corrected. Underlying heart disease should be considered and excluded.

For patients with recurrent episodes, beta blockers, the calcium channel blockers diltiazem or verapamil, and the antiarrhythmic drugs flecainide, propafenone, disopyramide, sotalol, and amiodarone can be effective, but potential toxicities and adverse effects often warrant avoiding these agents (Tables 276-3, 276-4, and 276-5). Catheter ablation targeting the AT

focus is effective in more than 80% of patients and is recommended for recurrent symptomatic AT when drugs fail or are not desired or for incessant AT causing tachycardia-induced cardiomyopathy.

Atrioventricular Nodal Reentry Tachycardia AV nodal reentry tachycardia (AVNRT) is the most common form of PSVT, representing approximately 60% of cases referred for catheter ablation. It most commonly manifests in the second to fourth decades of life, often in women. It is often well tolerated, but rapid tachycardia, particularly in the elderly, may cause angina, pulmonary edema, hypotension, or syncope. It is not usually associated with structural heart disease.

The mechanism is reentry involving the AV node and likely the perinodal atrium, made possible by the existence of multiple pathways for conduction from the atrium into the AV node (Fig. 276-5). In the most common form, a slowly conducting AV nodal pathway extends from the compact AV node near the bundle of His, inferiorly along the tricuspid annulus, adjacent to the coronary sinus os. The reentry wavefront propagates up this slow pathway to the compact AV node and then exits from the fast pathway at the top of the AV node. The path back to the slow pathway to complete the circuit is not defined. The conduction time from the compact AV node region to the atrium is similar to that from the compact node to the His bundle and ventricles, such that atrial activation occurs at about the same time as ventricular activation. The p wave is therefore inscribed during, slightly before, or slightly after the QRS and can be difficult to discern. Often the P wave is seen at the end of the QRS complex as a pseudo-r' in lead V₁ and pseudo-S waves in leads II, III, and aVF (Fig. 276-5A). The rate can vary with sympathetic tone. Simultaneous atrial and ventricular contraction results in atrial contraction against a closed tricuspid valve that produces cannon waves visible in the jugular venous pulse and that the patient often perceives as a fluttering sensation in the neck. Elevated venous pressures may also lead to release of natriuretic peptides that cause posttachycardia diuresis. Less frequently, the AV nodal reentry circuit revolves in the opposite direction and gives rise to a tachycardia with an R-P interval longer than the P-R interval, similar to AT. The p wave will have the morphology noted above, and in contrast to ATs, maneuvers or medications that produce AV block terminate the arrhythmia.

Acute treatment is the same as for PSVT (discussed below). Whether ongoing therapy is warranted depends on the severity of symptoms and frequency of episodes. Reassurance and instruction as to performance of the Valsalva maneuver to terminate episodes are sufficient for many patients. Administration of an oral beta blocker, verapamil, or diltiazem at the onset of an episode has been used to

TABLE 276-3 COMMONLY USED ANTIARRHYTHMIC AGENTS—INTRAVENOUS DOSE RANGE/PRIMARY INDICATION

Drug	Loading	Maintenance	Primary Indication	Class ^a
Adenosine	6–18 mg (rapid bolus)	N/A	Terminate reentrant SVT involving AV node	—
Amiodarone	15 mg/min for 10 min, 1 mg/min for 6 h	0.5–1 mg/min	AF, AFL, SVT, VT/VF	III
Digoxin	0.25 mg q2h until 1 mg total	0.125–0.25 mg/d	AF/AFL rate control	—
Diltiazem	0.25 mg/kg over 3–5 min (max 20 mg)	5–15 mg/h	SVT, AF/AFL rate control	IV
Esmolol	500 µg/kg over 1 min	50 µg/kg per min	AF/AFL rate control	II
Ibutilide	1 mg over 10 min if over 60 kg	N/A	Terminate AF/AFL	III
Lidocaine	1–3 mg/kg at 20–50 mg/min	1–4 mg/min	VT	IB
Metoprolol	5 mg over 3–5 min × 3 doses	1.25–5 mg q6h	SVT, AF rate control; exercise-induced VT; long QT	II
Procainamide	15 mg/kg over 60 min	1–4 mg/min	Convert/prevent AF/VT	IA
Quinidine	6–10 mg/kg at 0.3–0.5 mg/kg per min	N/A	Convert/prevent AF/VT	IA
Verapamil	5–10 mg over 3–5 min	2.5–10 mg/h	SVT, AF rate control	IV

^aClassification of antiarrhythmic drugs: class I—agents that primarily block inward sodium current; class IA agents also prolong action potential duration; class II—antisympathetic agents; class III—agents that primarily prolong action potential duration; class IV—calcium channel-blocking agents.

Abbreviations: AF, atrial fibrillation; AFL, atrial flutter; AV, atrioventricular; SVT, supraventricular tachycardia; VF, ventricular fibrillation; VT, ventricular tachycardia.

TABLE 276-4 COMMONLY USED ANTIARRHYTHMIC AGENTS: CHRONIC ORAL DOSING/PRIMARY INDICATIONS

Drug	Dosing Oral, mg, Maintenance	Half-Life, h	Primary Route(s) of Metabolism/ Elimination	Most Common Indication	Class ^a
Acebutolol	200–400 q12h	6–7	Renal/hepatic	AF rate control/SVT Long QT/RVOT VT	II
Amiodarone	100–400 qd	40–55 d	Hepatic	AF/VT prevention	III ^b
Atenolol	25–100 per d	6–9	Renal	AF rate control/SVT Long QT/RVOT VT	II
Digoxin	0.125–0.25 qd	38–48	Renal	AF rate control	—
Diltiazem	30–60 q6h	3–4.5	Hepatic	AF rate control/SVT	IV
Disopyramide	100–300 q6–8h	4–10	Renal 50%/hepatic	AF/SVT prevention	Ia
Dofetilide	0.125–0.5 q12h	10	Renal	AF prevention	III
Dronedrone	400 q12h	13–19	Hepatic	AF prevention	IIIb
Flecainide	50–200 q12h	7–22	Hepatic 75%/renal	AF/SVT/VT prevention	Ic
Metoprolol	25–100 q6h	3–8	Hepatic	AF rate control/SVT Long QT/RVOT VT	II
Mexiletine	150–300 q8–12h	10–14	Hepatic	VT prevention	Ib
Nadolol	40–240 per d	10–24	Renal	Same as metoprolol	II
Propafenone	150–300 q8h	2–8	Hepatic	AF/SVT/VT prevention	Ic
Quinidine	300–600 q6h	6–8	Hepatic 75%/renal	AF/SVT/VT prevention	Ia
Sotalol	80–160 q12h	12	Renal	AF/VT prevention	III
Verapamil	80–120 q6–8h	4.5–12	Hepatic/renal	AF rate control/RVOT VT Idiopathic LV VT	IV

^aClassification of antiarrhythmic drugs: class I—agents that primarily block inward sodium current; class II—antisymptomatic agents; class III—agents that primarily prolong action potential duration; class IV—calcium channel-blocking agents. ^bAmiodarone and dronedrone both are grouped in class III, but both also have class I, II, and IV properties.

Abbreviations: AF, atrial fibrillation; LV, left ventricular; RVOT, right ventricular outflow tract; SVT, supraventricular tachycardia; VT, ventricular tachycardia.

facilitate termination. Chronic therapy with these medications or flecainide is an option if prophylactic therapy is needed. Catheter ablation of the slow AV nodal pathway is recommended for patients with recurrent or severe episodes or when drug therapy is ineffective, not tolerated, or not desired by the patient. Catheter ablation is curative in over 95% of patients. The major risk is heart block requiring permanent pacemaker implantation, which occurs in less than 1% of patients.

Junctional Tachycardia Junctional ectopic tachycardia (JET) is due to automaticity within the AV node. It is rare in adults and more

frequently encountered as an incessant tachycardia in children, often in the perioperative period of surgery for congenital heart disease. It presents as a narrow QRS tachycardia, often with ventriculoatrial (VA) block, such that AV dissociation is present. JET can occur as a manifestation of increased adrenergic tone and may be seen after administration of isoproterenol. It may also occur for a short period of time after ablation for AVNRT.

Accelerated junctional rhythm is a junctional automatic rhythm between 50 and 100 beats/min. Initiation may occur with gradual acceleration in rate, suggesting an automatic focus, or after a premature

TABLE 276-5 COMMON AND PROARRHYTHMIC TOXICITIES OF ANTIARRHYTHMIC AGENTS

Drug	Potential Proarrhythmic Toxicities	Common Toxicities
Amiodarone	Sinus bradycardia, AV block, increase in defibrillation threshold. Rare: long QT and torsades des pointes, incessant slow VT in heart disease	Tremor, peripheral neuropathy, pulmonary fibrosis or inflammation, hypo- and hyperthyroidism, hepatitis, photosensitivity
Adenosine	Transient profound pauses, atrial fibrillation	Cough, flushing, chest pain, anxiety
Digoxin	AV block, fascicular tachycardia, accelerated junctional rhythm, atrial tachycardia with AV block	Anorexia, nausea, vomiting, visual changes
Disopyramide	Long QT and torsades des pointes, 1:1 ventricular response to atrial flutter	Anticholinergic effects, acute urinary retention (males), negative inotropy
Dofetilide	Long QT and torsades des pointes	Nausea
Dronedrone	Bradycardias and AV block, long QT and torsades des pointes (rare)	Gastrointestinal intolerance, exacerbation of heart failure
Flecainide	1:1 Ventricular response to atrial flutter; increased risk of ventricular tachycardias in patients with structural heart disease; sinus bradycardia	Dizziness, nausea, headache, decreased myocardial contractility
Ibutilide	Long QT and torsades des pointes	Nausea
Lidocaine	Slow VT in some patients with structural heart disease	Dizziness, confusion, delirium, seizures, coma
Mexiletine	Slow VT in patients with structural heart disease	Ataxia, tremor, gait disturbances, rash, nausea
Procainamide	Long QT and torsades des pointes, accelerated ventricular rate in AF or flutter	Lupus erythematosus-like syndrome (more common in slow acetylators), anorexia, nausea, neutropenia
Propafenone	1:1 Ventricular response to atrial flutter; increased risk ventricular tachycardias in patients with structural heart disease; sinus bradycardia	Taste disturbance, dyspepsia, nausea, vomiting
Quinidine	Long QT and torsades des pointes, accelerated ventricular rate in AF or flutter	Diarrhea, nausea, vomiting, cinchonism, thrombocytopenia
Sotalol	Long QT and torsades des pointes	Hypotension, bronchospasm from β -blocking effect

Abbreviations: AF, atrial fibrillation; AV, atrioventricular; VT, ventricular tachycardia.

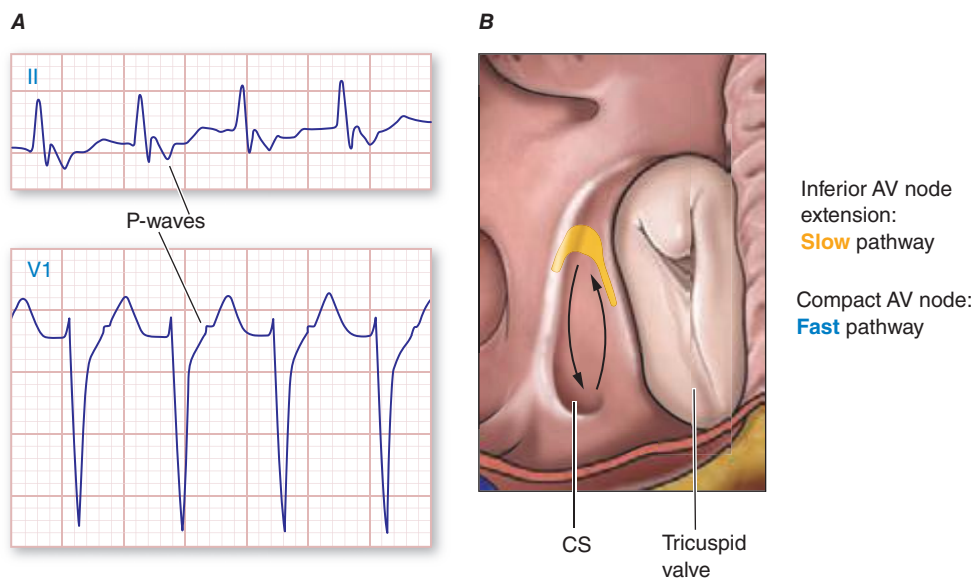


FIGURE 276-5 Atrioventricular (AV) node reentry. **A.** Leads II and V_1 are shown. P waves are visible at the end of the QRS complex and are negative in lead II, and may give the impression of S waves in the inferior limb leads II, III, and aVF and an R' in lead V_1 . **B.** Stylized version of the AV nodal reentry circuit within the triangle of Koch (Fig. 276-1) that involves AV node and its extensions along with perinodal atrial tissue.

ventricular contraction, suggesting a focus of triggered automaticity. VA conduction is usually present, with p-wave morphology and timing such that it resembles slow AVNRT.

ACCESSORY PATHWAYS AND THE WOLFF-PARKINSON-WHITE SYNDROME

Accessory pathways (APs) occur in 1 in 1500–2000 people and are associated with a variety of arrhythmias including narrow-complex PSVT, wide-complex tachycardias, and, rarely, sudden death. Most patients have structurally normal hearts, but APs are associated with Ebstein's anomaly of the tricuspid valve and forms of hypertrophic cardiomyopathy including *PRKAG2* mutations, Danon's disease, and Fabry's disease.

APs are abnormal connections that allow conduction between the atrium and ventricles across the AV ring (Fig. 276-6). They are present from birth and are due to failure of complete partitioning of atrium and ventricle by the fibrous AV rings. They occur across either an AV valve annulus or the septum, most frequently between the left atrium and free wall of the left ventricle, followed by posteroseptal, right free wall, and anteroseptal locations. If the AP conducts from atrium to ventricle (antegrade) with a shorter conduction time than the AV node and His bundle, then the ventricles are preexcited during sinus rhythm, and the ECG shows a short P-R interval (<0.12 s), slurred initial portion of the QRS (delta wave), and prolonged QRS duration produced by slow conduction through direct activation of ventricular myocardium over the AP (Fig. 276-6A). The morphology of the QRS and delta wave is determined by the AP location (Fig. 276-7) and the degree of fusion between the excitation wavefronts from conduction over the AV node and conduction over the AP. Right-sided pathways preexcite the right ventricle, producing a left bundle branch block–like configuration in lead V_1 , and often show marked preexcitation because of relatively close proximity of the AP to the sinus node (Fig. 276-7). Left-sided pathways preexcite the left ventricle and may produce a right bundle branch–like configuration in lead V_1 and a negative delta wave in aVL, indicating initial depolarization of the lateral portion of the left ventricle that can mimic q waves of lateral wall infarction (Fig. 276-7). Preexcitation due to an AP at the diaphragmatic surface of the heart, typically in the paraseptal region, produces delta waves that are negative in leads III and aVF, mimicking the q waves of inferior wall infarction (Fig. 276-7). Preexcitation can be intermittent and disappear during exercise as conduction over the AV node accelerates and takes over ventricular activation completely.

Wolff-Parkinson-White (WPW) syndrome is defined as a preexcited QRS during sinus rhythm and episodes of PSVT. There are a

number of variations of APs, which may not cause preexcitation and/or arrhythmias. Concealed APs allow only retrograde conduction, from ventricle to atrium, so no preexcitation is present during sinus rhythm, but SVT can occur. Fasciculoventricular connections between the His bundle and ventricular septum produce preexcitation but do not cause arrhythmia, nor do fibers such as atrio-Hisian connections, probably because the circuit is too short to promote reentry. Atriofascicular pathways, also known as Mahaim fibers, probably represent a duplicate AV node and His-Purkinje system that connect the right atrium to fascicles of the right bundle branch and conduct slowly only in the anterograde direction.

AV Reentry Tachycardia The most common tachycardia caused by an AP is the PSVT designated *orthodromic AV reentry*. The circulating reentry wavefront propagates from the atrium anterogradely over the AV node and His-Purkinje system to the ventricles and then reenters the atria via retrograde conduction over the AP (Fig. 276-6B). The QRS is narrow or may have typical right or left bundle branch block, but without preexcitation during tachycardia. Because excitation through the normal AV conduction system and AP are necessary, AV or VA block results in tachycardia termination. During sinus rhythm, preexcitation is seen if the pathway also allows anterograde conduction (Fig. 276-6A). Most commonly, during tachycardia the R-P interval is shorter than the P-R interval and can resemble AVNRT (Fig. 276-1). Unlike typical AVNRT, P-wave timing is never simultaneous with a narrow QRS complex because the ventricles must be activated before the reentry wavefront reaches the AP and conducts back to the atrium. The morphology of the P wave is determined by the pathway location, but can be difficult to assess because it is usually inscribed during the ST segment. The p wave in posteroseptal APs is negative in leads II, III, and aVF, similar to that of AV nodal reentry, but P-wave morphology differs from AV nodal reentry for pathways in other locations (Fig. 276-7).

Occasionally, an AP conducts extremely slowly in the retrograde direction, which results in tachycardia with a long R-P interval, similar to most ATs. These pathways are usually located in the septal region and have negative p waves in leads II, III, and aVF. Slow conduction facilitates reentry, often leading to nearly incessant tachycardia, known as *paroxysmal junctional reciprocating tachycardia* (PJRT). Tachycardia-induced cardiomyopathy can occur. Without an invasive electrophysiology study, it may be difficult to distinguish this form of orthodromic AV reentry from atypical AV nodal reentry or AT.

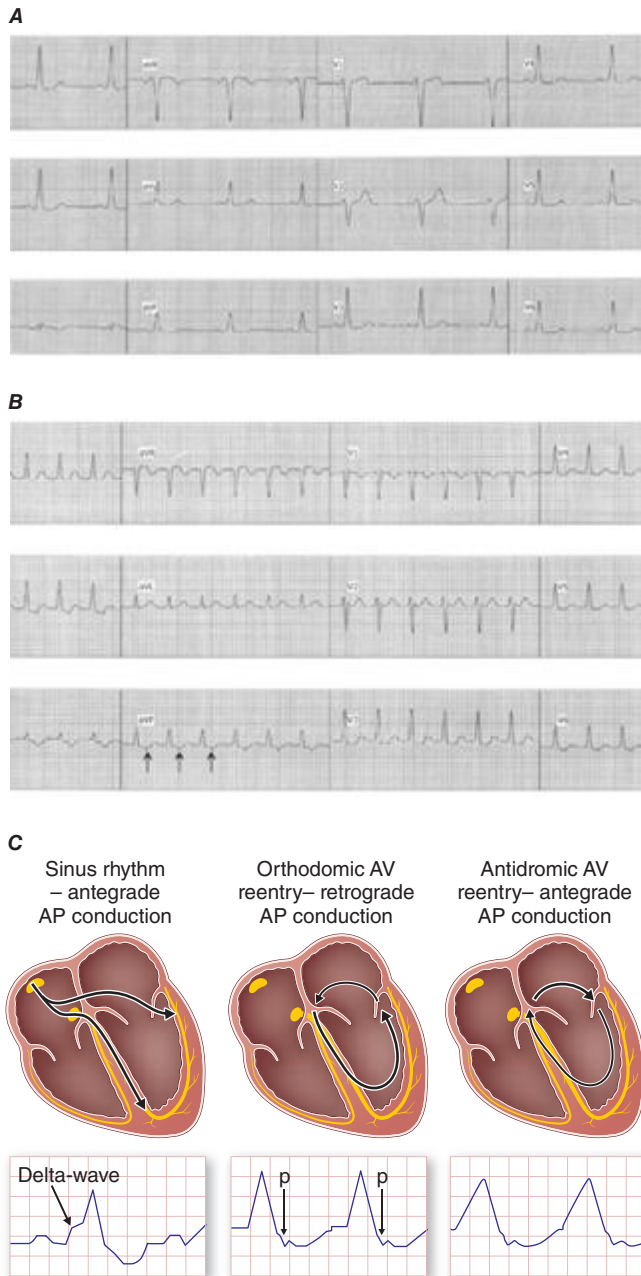


FIGURE 276-6 Wolff-Parkinson-White (WPW) syndrome. **A.** A 12-lead electrocardiogram in sinus rhythm (SR) of a patient with WPW demonstrating short P-R interval, delta waves, and widened QRS complex. This patient had an anteroseptal location of the AP. **B.** Orthodromic AV reentry in a patient with WPW syndrome using a posteroseptal AP. Note the P waves in the ST segment (*arrows*) seen in lead III and normal appearance of QRS complex. **C.** Three most common rhythms associated with WPW syndrome: sinus rhythm demonstrating antegrade conduction over the AP and AV node; orthodromic AVRT using retrograde conduction over the AP and antegrade conduction over the AV node; and antidromic AVRT using retrograde conduction over the AV node and antegrade conduction over the AP. AP, accessory pathway; AV, atrioventricular; AVRT, atrioventricular reentry tachycardia; WPW, Wolff-Parkinson-White.

Preexcited Tachycardias Preexcited tachycardia occurs when the ventricles are activated by antegrade conduction over the AP (Fig. 276-6C). The most common is *antidromic AV reentry* in which activation propagates from atrium to ventricle via the AP and then conducts retrogradely to the atria via the His-Purkinje system and the AV node (or rarely a

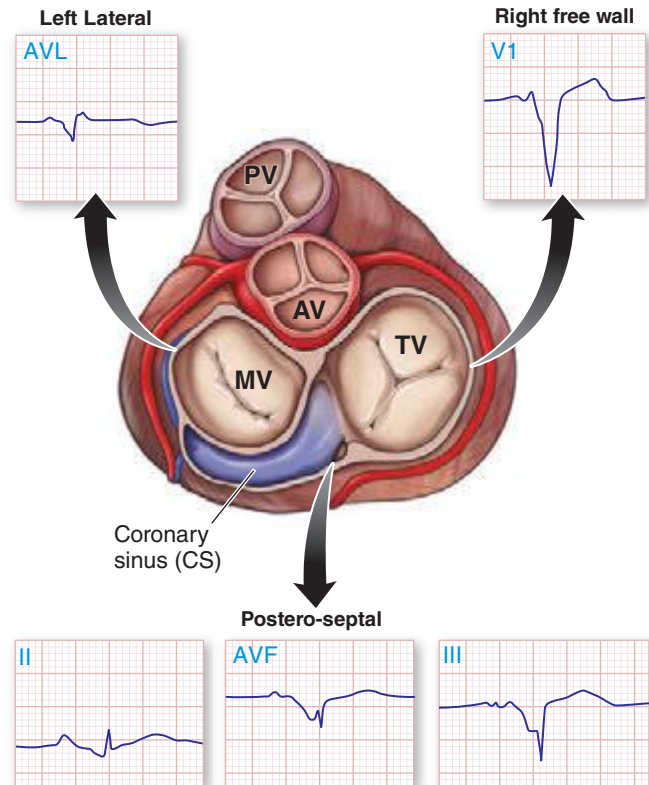


FIGURE 276-7 Potential locations for accessory pathways in patients with Wolff-Parkinson-White Syndrome and typical QRS appearance of delta waves that can mimic underlying structural heart disease such as myocardial infarction of bundle branch block. AV, aortic valve; MV, mitral valve; PV, pulmonary valve; TV, tricuspid valve.

second AP). The wide QRS complex is produced entirely via ventricular excitation over the AP because there is no contribution of ventricular activation over more rapidly conducting specialized His-Purkinje fibers. This tachycardia is often indistinguishable from monomorphic ventricular tachycardia. The presence of preexcitation in sinus rhythm suggests the diagnosis.

Preexcited tachycardia also occurs if an AP allows antegrade conduction to the ventricles during AT, atrial flutter, atrial fibrillation (Fig. 276-8), or AV nodal reentry. Atrial fibrillation and atrial flutter are potentially life threatening if the AP allows very rapid repetitive conduction. Approximately 25% of APs causing preexcitation allow minimum R-to-R intervals of less than 250 ms during atrial fibrillation are therefore associated with a risk of inducing ventricular fibrillation and sudden death. Preexcited atrial fibrillation presents as a wide-complex, very irregular rhythm. During atrial fibrillation, the ventricular rate is determined by the conduction properties of the AP and AV node. The QRS complex can appear quite bizarre and change on a beat-to-beat basis due to the variability in the degree of fusion from activation over the AV node and AP, or all beats may be due to conduction over the AP (Fig. 276-8). Ventricular activation from the Purkinje system may depolarize the ventricular end of the AP and prevent 1:1 atrial wavefront conduction over the AP. Slowing AV nodal conduction can thereby facilitate AP conduction and dangerously accelerate the ventricular rate. Administration of AV nodal-blocking agents including oral or intravenous verapamil, diltiazem, beta blockers, intravenous adenosine, and intravenous amiodarone are contraindicated. Preexcited tachycardias should be treated with electrical cardioversion or intravenous procainamide or ibutilide, which may terminate or slow the ventricular rate.

Management of Patients with Accessory Pathways Acute management of orthodromic AV reentry is discussed below for PSVT. Patients

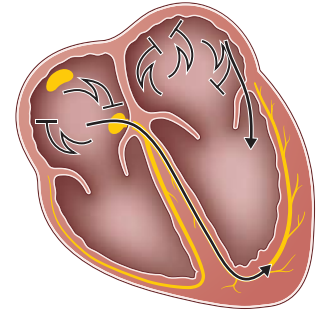
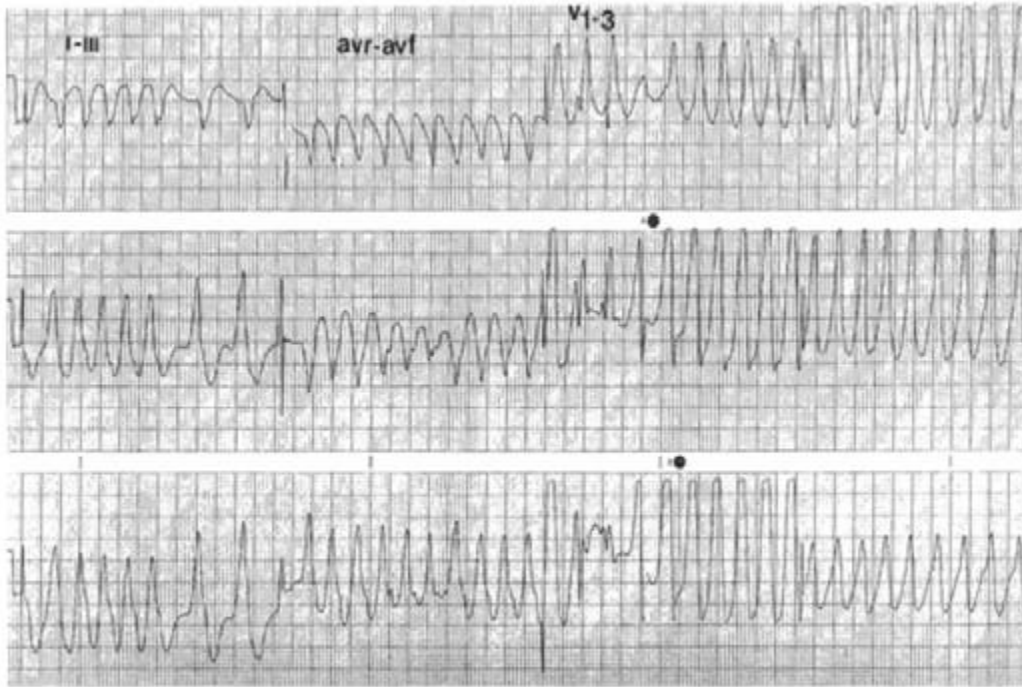


FIGURE 276-8 Preexcited atrial fibrillation (AF) due to conduction over a left free wall accessory pathway (AP). The electrocardiogram shows rapid irregular QRS complexes that represent fusion between conduction over the atrioventricular node and left free wall AP. Shortest R-R intervals between preexcited QRS complexes of less than 250 ms, as in this case, indicate a risk of sudden death with this arrhythmia.

with WPW syndrome may have wide-complex tachycardia due to antidromic AV reentry, orthodromic AV with bundle branch block, or a preexcited tachycardia, and treatment depends on the underlying rhythm.

Initial patient evaluation should include assessment for aggravating factors, including intercurrent illness and factors that increase sympathetic tone. Examination should focus on excluding underlying heart disease. An echocardiogram is reasonable to exclude Ebstein's anomaly and hypertrophic cardiomyopathy.

Patients with preexcitation who have symptoms of arrhythmia are at risk for developing atrial fibrillation and sudden death if they have an AP with high-risk properties. The risk of cardiac arrest is in the range of 2 per 1000 patients in adults but is likely greater in children. An invasive electrophysiology study is warranted to determine if the AP is high enough risk to warrant potentially curative catheter ablation. For patients with concealed APs or known low-risk APs causing orthodromic AV reentry, chronic therapy is guided by symptoms and frequency of events. Vagal maneuvers may terminate episodes, as may a dose of beta blocker, verapamil, or diltiazem taken at the onset of an episode. Chronic therapy with these agents or flecainide can reduce the frequency of episodes in some patients. Catheter ablation is warranted for recurrent arrhythmias when drugs are ineffective, not tolerated, or not desired by the patient or if the AP is considered high risk (Fig. 276-8). Efficacy is in the range of 95% depending on the location of the AP. Serious complications occur in fewer than 3% of patients, but can include AV block, cardiac tamponade, thromboemboli, coronary artery injury, and vascular access complications. Mortality occurs in less than 1 in 1000 patients.

Adults who have preexcitation but no arrhythmia symptoms have a risk of sudden death estimated to be 1 per 1000 patient-years. Electrophysiology study is usually advised for people in occupations for which an arrhythmia occurrence would place them or others at risk, such as police, military, and pilots, or for individuals who desire evaluation for risk. Routine follow-up without therapy is reasonable in others. Children are at greater risk of sudden death, approximately 2 per 1000 patient-years.

TREATMENT PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA

Acute management of narrow QRS PSVT is guided by the clinical presentation. Continuous ECG monitoring should be implemented and a 12-lead ECG should always be obtained when possible. In the presence of hypotension with unconsciousness or respiratory distress, QRS-synchronous direct current cardioversion is warranted, but this is rarely needed, because intravenous adenosine works promptly in most situations (see below). For stable individuals, initial therapy takes advantage of the fact that most PSVTs are dependent on AV nodal conduction (AV nodal reentry or orthodromic AV reentry) and therefore likely to respond to sympatholytic and vagotonic maneuvers and drugs (Fig. 276-9). As these are administered, the ECG should be continuously recorded, because the response can

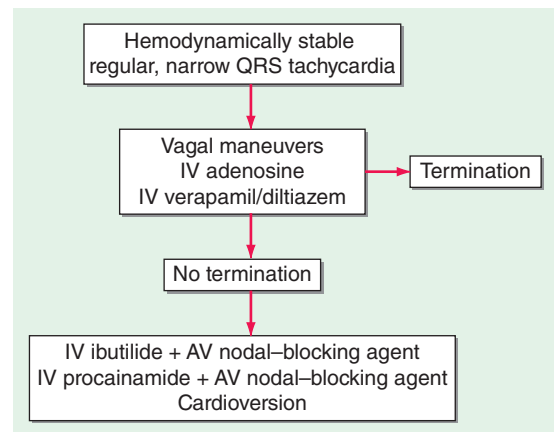


FIGURE 276-9 Treatment algorithm for patients presenting with hemodynamically stable paroxysmal supraventricular tachycardia. AV, atrioventricular.

1484 establish the diagnosis. AV block with only transient slowing of tachycardia may expose ongoing p waves, indicating AT or atrial flutter as the mechanism.

Carotid sinus massage is reasonable provided the risk of carotid vascular disease is low, as indicated by absence of carotid bruits and no prior history of stroke. A Valsalva maneuver should be attempted in cooperative individuals, and if effective, the patient can be taught to perform this maneuver as needed. If vagal maneuvers fail or cannot be performed, intravenous adenosine will terminate the vast majority of PSVT by transiently blocking conduction in the AV node. Adenosine may produce transient chest pain, dyspnea, and anxiety. It is contraindicated in patients with prior cardiac transplantation due to potential hypersensitivity. It can theoretically aggravate bronchospasm. Adenosine precipitates atrial fibrillation, which is usually brief, in up to 15% of patients, so it should be used cautiously in patients with WPW syndrome in whom AF may produce hemodynamic instability. Intravenous beta blockers and calcium channel blockers (verapamil or diltiazem) are also effective but may cause hypotension before and after arrhythmia termination and have a longer duration of action. These agents can also be given orally and can be taken by the patient on an as-needed basis to slow ventricular rate and facilitate termination by Valsalva maneuver.

The differential diagnosis of wide-complex tachycardia includes ventricular tachycardia (Chap. 277), PSVT with bundle branch block aberrancy, and preexcited tachycardia (see above). In general, these should be managed as ventricular tachycardia until proven otherwise. If the tachycardia is regular and the patient is stable, a trial of intravenous adenosine is reasonable. Very irregular wide-complex tachycardia should be managed with cardioversion, intravenous procainamide, or ibutilide, which presumes preexcited atrial fibrillation or flutter (see above). If the diagnosis of PSVT with aberrancy is unequivocal, as may be the case in patients with prior episodes, treatment for PSVT is reasonable. In all cases, continuous ECG monitoring should be implemented, and emergency cardioversion and defibrillation should be available.

COMMON ATRIAL FLUTTER AND MACROREENTRANT ATRIAL TACHYCARDIAS

Macroreentrant atrial tachycardia is due to a large reentry circuit, often associated with areas of scar in the atria. *Common or typical right atrial flutter* is due to a circuit that revolves around the tricuspid valve annulus, bounded anteriorly by the annulus and posteriorly by functional conduction block in the crista terminalis. The wavefront passes through an isthmus between the inferior vena cava and the tricuspid valve annulus, known as the sub-Eustachian or cavotricuspid isthmus, where it is susceptible to interruption by catheter ablation. Thus, common atrial flutter is *cavotricuspid isthmus-dependent atrial flutter*. This circuit most commonly revolves in a counterclockwise direction (as viewed looking toward the tricuspid annulus from the ventricular aspect), which produces the characteristic negative sawtooth flutter waves in leads II, III, and aVF and positive P waves in lead V₁ (Fig. 276-10). When the direction is reversed, clockwise rotation produces the opposite P-wave vector in those leads. The atrial rate is typically 240–300 beats/min but may be slower in the presence of atrial disease or antiarrhythmic drugs. It often conducts to the ventricles with 2:1 AV block, creating a regular tachycardia at 150 beats/min, with p waves that may be difficult to discern. Maneuvers that increase AV nodal block will typically expose flutter waves, allowing diagnosis.

Common right atrial flutter often occurs in association with atrial fibrillation and with atrial scar from senescence or prior cardiac surgery. Some patients with atrial fibrillation that is treated with an antiarrhythmic drug, particularly flecainide, propafenone, or amiodarone, will present with atrial flutter rather than fibrillation.

Macroreentrant ATs that are not dependent on conduction through the cavotricuspid isthmus are referred to as *atypical atrial flutters*. They can occur in either atrium and are usually associated with areas of scar. Left atrial flutter and perimitral left atrial flutter are commonly seen after extensive left atrial ablation for atrial fibrillation or atrial surgery. The clinical presentation is similar to common atrial flutter, but with different P-wave morphologies. They can be difficult to distinguish from focal AT, and in most cases, the mechanism can only be confirmed by an electrophysiology study.

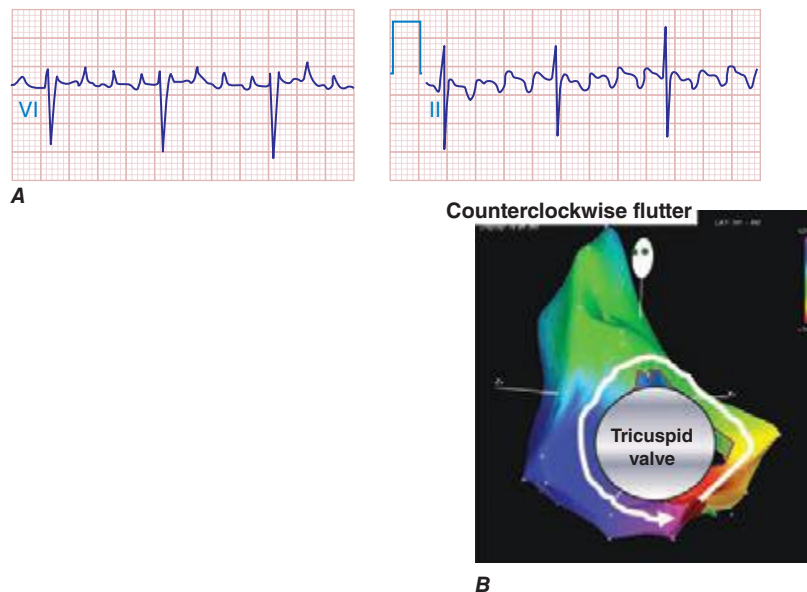


FIGURE 276-10 **A.** Common right atrial flutter, also known as cavotricuspid isthmus flutter, showing positive P waves in lead V₁ and negative “sawtooth” pattern in lead II typical of counterclockwise rotation relative to the tricuspid valve annulus. (Adapted from F Marchlinski: *The tachyarrhythmias*. In Longo DL et al [eds]: *Harrison's Principles of Internal Medicine*, 18th ed. New York, McGraw-Hill, 2012, pp 1878–1900.) **B.** A right atrial map of common counterclockwise flutter is shown. Colors indicate activation time, progressing from red to yellow to green, blue, and purple. The reentry path parallels the tricuspid annulus.

TREATMENT ATRIAL FLUTTER AND MACROREENTRANT ATRIAL TACHYCARDIAS

Initial management of atrial flutter is similar to that for atrial fibrillation, discussed in more detail below. Electrical cardioversion is warranted for hemodynamic instability or severe symptoms. Otherwise, rate control can be achieved with administration of AV nodal-blocking agents, but this is often more difficult than for atrial fibrillation. The risk of thromboembolic events is felt to be similar to that associated with atrial fibrillation. Anticoagulation is warranted prior to conversion for episodes more than 48 h in duration and chronically for patients at increased risk of thromboembolic stroke based on the CHA₂DS₂-VASc scoring system (Table 276-6).

For a first episode of atrial flutter, conversion to sinus rhythm with no antiarrhythmic drug therapy is reasonable. For recurrent episodes, antiarrhythmic drug therapy with sotalol, dofetilide, disopyramide, and amiodarone may be considered, but more than 70% of patients experience recurrences. For recurrent episodes of common atrial flutter, catheter ablation of the cavotricuspid isthmus abolishes the arrhythmia in over 90% of patients with a low risk of complications that are largely related to vascular access and infrequent heart block. Approximately 50% of patients presenting with atrial flutter develop atrial fibrillation within the next 5 years.

MULTIFOCAL ATRIAL TACHYCARDIA

Multifocal AT (MAT) is characterized by at least three distinct P-wave morphologies with rates typically between 100 and 150 beats/min. Unlike atrial fibrillation, there are clear isoelectric intervals between P waves (Fig. 276-11). The mechanism is likely triggered automatically from multiple atrial foci. It is usually encountered in patients with chronic pulmonary disease and acute illness.

TREATMENT MULTIFOCAL ATRIAL TACHYCARDIA

Therapy for MAT is directed at treating the underlying disease and correcting any metabolic abnormalities. Electrical cardioversion has no effect. The calcium channel blockers verapamil or diltiazem may slow the atrial and ventricular rate. Patients with severe pulmonary disease often do not tolerate beta blocker therapy. MAT may respond to amiodarone, but long-term therapy with this agent is usually avoided due to its toxicities, particularly pulmonary fibrosis.

ATRIAL FIBRILLATION

Atrial fibrillation (AF) is characterized by disorganized, rapid, and irregular atrial activation with loss of atrial contraction and with an irregular ventricular rate that is determined by AV nodal conduction (Fig. 276-12). In an untreated patient, the ventricular rate also tends

TABLE 276-6 CHA₂DS₂-VASc RISK ASSESSMENT AND ORAL ANTICOAGULANTS

Risk Factors	Points	CHA ₂ DS ₂ -VASc Score	Estimated Annual Stroke Rate ^a
C – congestive heart failure	1	0	0
H – hypertension	1	1	1.3%
A – age ≥75 y	2	2	2.2%
D – diabetes mellitus	1	3	3.2%
S – stroke or TIA, embolus	2	4	4.0%
V – vascular disease	1	5	6.7%
A – age 65–75 y	1	6–9	>9%
Sex – female	1		

Anticoagulants	Mechanism	Excretion	Dosing Considerations	Risk/Benefit
Warfarin	Vitamin K antagonist	Liver	Adjusted to INR 2–3 Days to therapeutic effect Multiple drug/food interactions (e.g., amiodarone)	Major hemorrhage: 1% per year Intracranial hemorrhage: 0.1–0.6% per year Risk of bleeding increases with INR >3.5 Inexpensive
Dabigatran ^b	Thrombin inhibitor	Kidney CCr >30 mL/min CCr 15–30 mL/min	150 mg bid 75 mg bid P-glycoprotein substrate (inducers – rifampin, reduce concentration) (inhibitors – amiodarone, verapamil, dronedarone, quinidine), Proton pump inhibitors may reduce absorption	Onset of action within hours No reversal agent for bleeding
Rivaroxaban	Xa inhibitor	Kidney CCr ≥50 mL/min CCr 15–50 mL/min	P-glycoprotein substrate 20 mg daily 15 mg daily	No reversal agent for bleeding
Apixaban	Xa inhibitor	Kidney and liver Cr >1.5 mg/dL	P-glycoprotein substrate 2.5 mg bid	No reversal agent for bleeding

^aModified from GY Lip et al: Lancet 379:648, 2012. ^bU.S. Food and Drug Administration recommended dosing; other regimens are available outside the United States.

Abbreviations: CCr, creatinine clearance; Cr, creatinine; INR, international normalized ratio; TIA, transient ischemic attack.



FIGURE 276-11 Multifocal atrial tachycardia. Rhythm strip obtained from a patient with severe pulmonary disease during an acute illness. Arrows note three distinct P-wave morphologies.

to be rapid and variable, between 120 and 160 beats/min, but in some patients, it may exceed 200 beats/min. Patients with high vagal tone or AV nodal conduction disease may have slow rates.

AF is the most common sustained arrhythmia and is a major public health problem. Prevalence increases with age, and more than 95% of AF patients are older than 60 years of age. The prevalence by age 80 is approximately 10%. The lifetime risk of developing AF for individuals 40 years old is approximately 25%. AF is slightly more common in men than women and more common in whites than blacks. Risk factors for developing AF in addition to age include hypertension, diabetes mellitus, cardiac disease, and sleep apnea. AF is a marker for heart disease, the severity of heart disease, and age, and it is therefore difficult to determine the extent to which AF itself contributes to associated increased mortality and morbidity. AF is associated with increased risk of developing heart failure. AF increases the risk of stroke by fivefold and is estimated to be the cause of 25% of strokes. It also increases the risk of dementia.

AF is occasionally associated with an acute precipitating factor such as hyperthyroidism, acute alcohol intoxication, or an acute illness

including myocardial infarction or pulmonary embolism. AF occurs in up to 30% of patients recovering from cardiac surgery, associated with inflammatory pericarditis.

The clinical type of AF suggests the underlying pathophysiology (Fig. 276-12). Paroxysmal AF is defined as episodes that start and stop spontaneously. It is often initiated by small reentrant or rapidly firing foci in sleeves of atrial muscle along the pulmonary veins. Catheter ablation that isolates these foci usually abolishes the AF. Persistent AF has a longer duration, exceeding 7 days, and, in many cases, will continue unless cardioversion is performed. Cardioversion can be followed by prolonged periods of sinus rhythm. Episodes may be initiated by rapidly firing foci, but persistence of the arrhythmia is likely due to single or multiple areas of reentry facilitated by structural and electrophysiologic atrial abnormalities. In patients with long-standing persistent AF (>1 year), significant structural changes are present in the atrium that support reentry and automaticity, making it difficult to restore and maintain sinus rhythm. Some patients progress over years from paroxysmal to persistent AF. Fibrosis that develops with

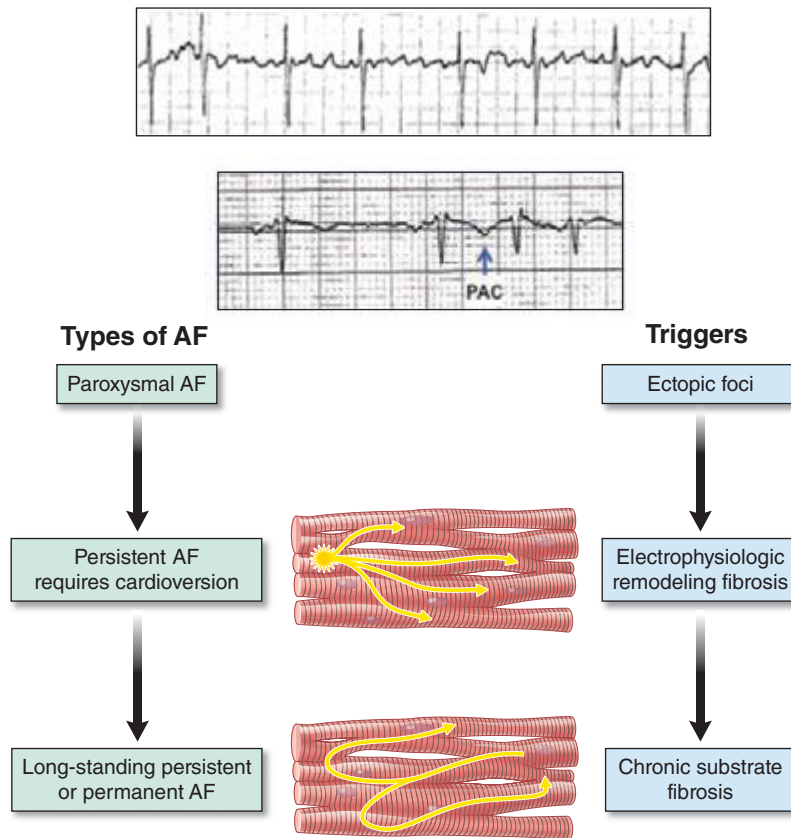


FIGURE 276-12 A rhythm strip of atrial fibrillation (AF) showing no distinct P-wave morphology and irregular ventricular response. Diagram depicts atrial fibrillation types. Paroxysmal AF is initiated by premature beats, as shown in the rhythm strip (arrow) after two sinus beats. Triggering foci are often an important cause of this arrhythmia. Persistent AF is associated with atrial structural and electrophysiologic remodeling, as well as with triggering foci in many patients. Long-standing persistent AF is associated with greater structural remodeling with atrial fibrosis and electrophysiologic remodeling.

aging and atrial hypertrophy in response to hypertension and other cardiac disease may be an important promoting factor, although electrophysiologic changes to conduction and refractoriness occur as well in response to chronic tachycardia in the atrium.

Clinical consequences are related to rapid ventricular rates, loss of atrial contribution to ventricular filling, and predisposition to thrombus formation in the left atrial appendage with potential embolization. Presentations vary with the ventricular rate and underlying heart disease and comorbidities. Many patients are asymptomatic. Rapid rates may cause hemodynamic collapse or heart failure exacerbations particularly in patients with impaired cardiac function, hypertrophic cardiomyopathy, and heart failure with preserved systolic function. Exercise intolerance and easy fatigability are common. Occasionally, dizziness or syncope occurs due to pauses when AF terminates to sinus rhythm (Fig. 276-13).

TREATMENT ATRIAL FIBRILLATION

Treatment for AF is primarily guided by patients' symptoms, the hemodynamic effect of AF, the duration of AF if there are persistent risk factors for stroke, and underlying heart disease. Oral anticoagulation in high-risk patients with AF includes vitamin K antagonists or the newer anticoagulants such as thrombin inhibitors (dabigatran) or factor Xa inhibitors (rivaroxaban, apixaban), but not antiplatelet agents (aspirin and clopidogrel), which have substantially less effect.

New-onset AF that produces severe hypotension, pulmonary edema, or angina should be electrically cardioverted starting with a QRS synchronous shock of 200 J, ideally after sedation or anesthesia is achieved. Greater shock energy and different electrode placements may be tried if the shock fails to terminate AF. If AF terminates and reinitiates, administration of an antiarrhythmic drug, such as ibutilide, and repeat cardioversion may be considered. If the patient is stable, immediate management involves rate control to alleviate or prevent

symptoms, anticoagulation if appropriate, and cardioversion to restore sinus rhythm if AF is persistent. Anticoagulation strategies for new-onset AF are debated. In the absence of contraindications, it is usually appropriate to initiate systemic anticoagulation with heparin immediately, while evaluation and other therapies are implemented.

CARDIOVERSION AND ANTICOAGULATION

Cardioversion *within 48 h of the onset of AF* is common practice in patients who have not been anticoagulated, provided that they are not at high risk for stroke due to a prior history of embolic events, rheumatic mitral stenosis, or hypertrophic cardiomyopathy with marked left atrial enlargement. These patients are usually at risk of recurrence, such that initiation of anticoagulation is considered based on the patient's individual risk for stroke, commonly assessed from the CHA₂DS₂-VASc score.

If the duration of AF exceeds 48 h or is unknown, there is greater concern for thromboembolism with cardioversion, even in patients considered low risk for stroke. There are two approaches to mitigate the risk related to cardioversion. One option is to anticoagulate continuously for 3 weeks before and a minimum of 4 weeks after cardioversion. A second approach is to start anticoagulation and perform a transesophageal echocardiogram to determine if thrombus is present in the left atrial appendage. If thrombus is absent, cardioversion can be performed and anticoagulation continued for a minimum of 4 weeks because recovery of atrial mechanical function after electrical or pharmacologic cardioversion may be delayed and thrombus can form and embolize days after cardioversion. Some patients may merit ongoing anticoagulation after cardioversion, depending on stroke risk profile.

RATE CONTROL

Acute rate control can be achieved with beta blockers and/or the calcium channel blockers verapamil and diltiazem administered either intravenously or orally, as warranted by the urgency of the clinical situation. Digoxin may be added, particularly in heart failure

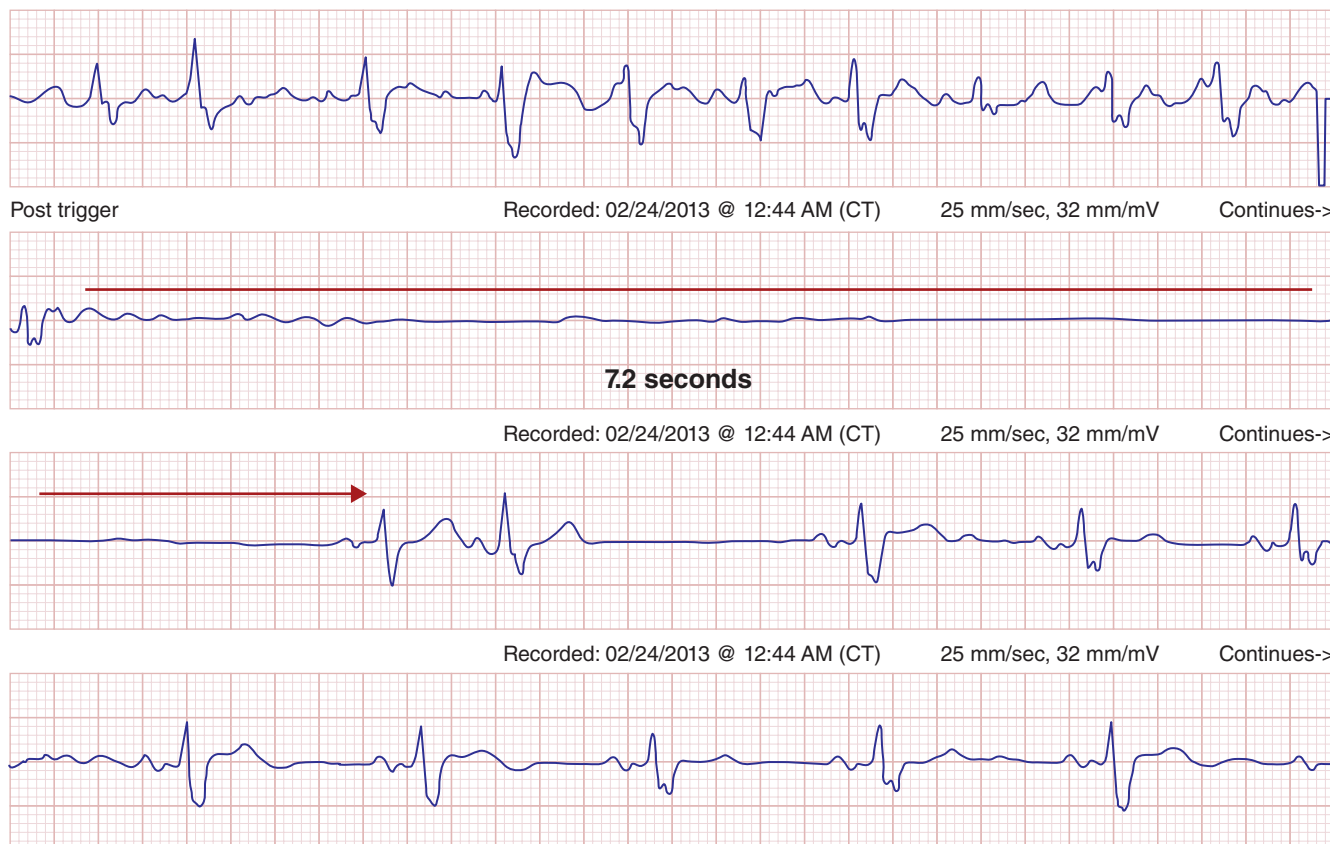


FIGURE 276-13 A continuous rhythm strip is shown. Atrial fibrillation is present at the top and abruptly terminates in the second tracing, with atrial and ventricular standstill for 7.2 s until resumption of sinus rhythm. The patient experienced syncope.

1488 patients, because it does not have negative inotropic effects, particularly if use of AV nodal–blocking agents is limited by poor tolerance or is contraindicated. Its effect is modest but synergistic with the other AV nodal–blocking agents, but it is particularly limited when sympathetic tone is elevated. Typically, the goal of acute rate control is to reduce the ventricular rate to less than 100/min, but the goal must be guided by the clinical situation.

CHRONIC RATE CONTROL

For patients who remain in AF chronically, the goal of rate control is to alleviate and prevent symptoms and prevent deterioration of ventricular function from excessive rates. β -Adrenergic blockers, calcium channel blockers, and digoxin are used, sometimes in combination. Rate should be assessed with exertion and medications adjusted accordingly. Exertion-related symptoms are often an indication of inadequate rate control. The initial goal is a resting heart rate of less than 80 beats/min that increases to less than 100 beats/min with light exertion, such as walking. If it is difficult to slow the ventricular rate to that degree, allowing a resting rate of up to 110 beats/min is acceptable provided it does not cause symptoms and ventricular function remains normal. Periodic assessment of ventricular function is warranted because some patients develop tachycardia-induced cardiomyopathy.

If adequate rate control in AF is difficult to achieve, further consideration should be given to restoring sinus rhythm. Catheter ablation of the AV junction to create heart block and implantation of a permanent pacemaker reliably achieve rate control without the need for AV nodal agents, but implement life-long permanent pacing. Right ventricular apical pacing induces dyssynchronous ventricular activation that can be symptomatic or depress ventricular function in some patients. Biventricular pacing may be used to minimize the degree of ventricular dyssynchrony.

STROKE PREVENTION IN ATRIAL FIBRILLATION

The majority of patients warrant chronic anticoagulation, but selection of therapy should be individualized based on patient profile and risks and benefits of individual agents. Anticoagulation with a vitamin K antagonist is warranted for all patients with AF who have rheumatic mitral stenosis or mechanical heart valves for whom the newer anticoagulants have not been tested. Anticoagulation with a vitamin K antagonist (warfarin) or the newer oral anticoagulants is warranted for patients who have had more than 48 h of AF and are undergoing cardioversion, for patients who have a prior history of stroke, or for patients with a $\text{CHA}_2\text{DS}_2\text{-VASc}$ score of ≥ 2 , but it may be considered in patients with a risk score of 1. The approach to patients with paroxysmal AF is the same as for persistent AF. It is recognized that many patients who appear to have infrequent AF episodes often have asymptomatic episodes that put them at risk. Absence of AF during periodic monitoring is not sufficient to indicate low risk. The role of continuous monitoring with implanted recorders or pacemakers is not yet clear as a guide for anticoagulation in patients with a borderline risk profile. Bleeding is the major risk of anticoagulation. Major bleeding requiring transfusion or in a critical area (e.g., intracranial) occurs in approximately 1% of patients per year. Risk factors for bleeding include age >65 –75 years, heart failure, history of anemia, and excessive alcohol or nonsteroidal anti-inflammatory drug use. Patients with coronary stents who require antiplatelet therapy with aspirin and a thienopyridine are at particularly high risk of bleeding.

Warfarin reduces the annual risk of stroke by 64% compared to placebo and by 37% compared to antiplatelet therapy. The newer anticoagulants, dabigatran, rivaroxaban, and apixaban, have been found to be noninferior to warfarin in individual trials, and analysis of pooled data suggests superiority to warfarin by small absolute margins of 0.4–0.7% in reduction of mortality, stroke, major bleeding, and intracranial hemorrhage. Warfarin is an inconvenient agent that requires several days to achieve a

therapeutic effect (prothrombin time [PT]/international normalized ratio [INR] >2), requires monitoring of PT/INR to adjust dose, and has many drug and food interactions, thus limiting patient compliance. The newer agents are easier to use and achieve reliable anticoagulation promptly without requiring dosage adjustment based on blood tests. Dabigatran, rivaroxaban, and apixaban have renal excretion, cannot be used with severe renal insufficiency, and require dose adjustment for modest renal impairment, which is of particular concern in the elderly, who are at increased bleeding risk. Excretion can also be influenced by P-glycoprotein inducers and inhibitors. Warfarin anticoagulation can be reversed by administration of fresh frozen plasma and vitamin K. Reversing agents for the newer anticoagulants are lacking (but in development), and bleeding must be managed with supportive care, with the expectation that clotting will improve over 12 h as the anticoagulant is excreted.

The antiplatelet agents aspirin and clopidogrel are inferior to warfarin for stroke prevention in AF and do not reduce the risk of bleeding. Clopidogrel combined with aspirin is better than aspirin alone but inferior to warfarin and has greater bleeding risk than aspirin alone.

Chronic anticoagulation is contraindicated in some patients due to bleeding risks. Because most atrial thrombi are felt to originate in the left atrial appendage, surgical removal of the appendage, combined with atrial maze surgery, may be considered for patients undergoing surgery, although removal of the appendage has not been unequivocally shown to reduce the risk of thromboembolism. Percutaneous devices that occlude or ligate the left atrial appendage are being studied for safety and efficacy.

RHYTHM CONTROL

The decision to administer antiarrhythmic drugs or perform catheter ablation to attempt maintenance of sinus rhythm (commonly referred to as the “rhythm control strategy”) is mainly guided by patient symptoms and preferences regarding the benefits and risks of therapies. In general, patients who maintain sinus rhythm have better survival than those who continue to have AF. This is likely because continued AF is a marker of disease severity. In randomized trials, administration of antiarrhythmic medications to maintain sinus rhythm did not improve survival or symptoms compared to a rate control strategy, and the drug therapy group had more hospitalizations. Disappointing efficacy and toxicities of available antiarrhythmic drugs and patient selection bias may be factors that influenced the results of these trials. The impact of catheter ablation on mortality is not known. A rhythm control strategy is usually selected for patients with symptomatic paroxysmal AF, a first episode of symptomatic persistent AF, AF with difficult rate control, and AF that has resulted in depressed ventricular function or that aggravates heart failure. A rhythm control strategy is more likely to be favored in younger patients than in sedentary or elderly patients in whom rate control is usually easily achieved. Even if sinus rhythm is apparently maintained, anticoagulation is recommended according to the $\text{CHA}_2\text{DS}_2\text{-VASc}$ stroke risk profile because asymptomatic episodes of AF are common. Following a first episode of persistent AF, a strategy using AV nodal–blocking agents, cardioversion, and anticoagulation is reasonable, in addition to addressing possible aggravating factors, including hypertension, heart failure, and sleep apnea. If recurrences are infrequent, periodic cardioversion is reasonable.

Pharmacologic Therapy for Maintaining Sinus Rhythm The goal of pharmacologic therapy is to maintain sinus rhythm or reduce episodes of AF. Drug therapy can be instituted once sinus rhythm has been established or in anticipation of cardioversion. β -Adrenergic blockers and calcium channel blockers help control ventricular rate, improve symptoms, and possess a low-risk profile, but have low efficacy for preventing AF episodes. Risks and side effects of antiarrhythmic drugs are a major consideration in selecting

277 Ventricular Arrhythmias

Roy M. John, William G. Stevenson

therapy. Class I sodium channel–blocking agents (e.g., flecainide, propafenone, disopyramide) are options for subjects without significant structural heart disease, but they have negative inotropic and proarrhythmic effects that warrant avoidance in patients with coronary artery disease or heart failure. The class III agents sotalol and dofetilide can be administered to patients with coronary artery disease or structural heart disease but have approximately a 3% risk of inducing excessive QT prolongation and torsades des pointes. Dofetilide should be initiated only in a hospital with ECG monitoring, and many physicians take this approach with sotalol as well. Dronedrone increases mortality in patients with heart failure. All of these agents have modest efficacy in patients with paroxysmal AF, of whom approximately 30–50% will benefit. Amiodarone is more effective, maintaining sinus rhythm in approximately two-thirds of patients. It can be administered to patients with heart failure and coronary artery disease. Over 20% of patients experience toxicities during long-term therapy.

CATHETER AND SURGICAL ABLATION FOR ATRIAL FIBRILLATION

Catheter ablation avoids antiarrhythmic drug toxicities but has procedural risks and requires an experienced center. For patients with previously untreated but recurrent paroxysmal AF, catheter ablation has similar efficacy to antiarrhythmic drug therapy and is superior to antiarrhythmic drugs for patients who have recurrent AF despite drug treatment. The procedure involves cardiac catheterization, transatrial septal puncture, and radiofrequency ablation or cryoablation to electrically isolate the regions around the pulmonary veins, abolishing the effect of triggering foci to interact with the left atrial AF substrate. Extensive areas of ablation are required, and gaps in healed ablation areas necessitate a repeat procedure in 20–50% of patients. Sinus rhythm is maintained for more than 1 year after one procedure in approximately 60% of patients and in 70–80% of patients after multiple procedures. Some patients become more responsive to antiarrhythmic drugs.

There is a 2–7% risk of major complications, including stroke (0.5–1%), cardiac tamponade (1%), phrenic nerve paralysis, bleeding from femoral access sites, and fluid overload with heart failure, that can emerge 1–3 days after the procedure. It is important to recognize the potential for delayed presentation of some complications. Ablation within the pulmonary veins can lead to pulmonary vein stenosis, presenting weeks to months after the procedure with dyspnea or hemoptysis. Esophageal ulcers can form immediately after the procedure and may rarely lead to a fistula between the left atrium and esophagus (estimated incidence of 0.1%) that presents as endocarditis and stroke 10 days to 3 weeks after the procedure.

Catheter ablation is less effective for persistent AF. More extensive ablation is often required, including areas that likely support reentry in regions outside the pulmonary venous antra, but individual strategies are debated. More than one ablation procedure is often required to maintain sinus rhythm.

Surgical ablation of AF is typically performed concomitant with cardiac valve or coronary artery surgery and less commonly as a stand-alone procedure; however, for patients with persistent AF, surgical or hybrid procedures may have higher single-procedure efficacy. Risks include sinus node injury requiring pacemaker implantation. Surgical removal of the left atrial appendage may reduce stroke risk, although thrombus can form in the remnant of the appendage or if the appendage is not completely ligated.

ACKNOWLEDGMENT

Portions of this chapter were retained from the work of the previous author, Francis Marchlinski.

Arrhythmias that originate in the ventricular myocardium or His-Purkinje system include premature ventricular beats, ventricular tachycardias that can be sustained or nonsustained, and ventricular fibrillation. Arrhythmia may emerge from a focus of myocardial or Purkinje cells capable of automaticity, or triggered automaticity, or from reentry through areas of scar or a diseased Purkinje system. Ventricular arrhythmias are often associated with structural heart disease and are an important cause of sudden death (Chap. 327). They also occur in some structurally normal hearts, in which case they are usually benign. Evaluation and management are guided by the risk of arrhythmic death, which is assessed based on symptoms, type of arrhythmia, and associated underlying heart disease.

DEFINITIONS

Ventricular arrhythmias are characterized by their electrocardiographic appearance and duration. Conduction away from the ventricular focus through the ventricular myocardium is slower than activation of the ventricles over the Purkinje system. Hence, the QRS complex during ventricular arrhythmias will be wide, typically >0.12 s.

Premature ventricular beats (also referred to as *premature ventricular contractions* [PVCs]) are single ventricular beats that fall earlier than the next anticipated supraventricular beat (Fig. 277-1). PVCs that originate from the same focus will have the same QRS morphology and are referred to as unifocal (Fig. 277-1A). PVCs that originate from different ventricular sites have different QRS morphologies and are referred to as multifocal (Fig. 277-1B). Two consecutive ventricular beats are *ventricular couplets*.

Ventricular tachycardia (VT) is three or more consecutive beats at a rate faster than 100 beats/min. Three or more consecutive beats at slower rates are designated an *idioventricular rhythm* (Fig. 277-1C). VT that terminates spontaneously within 30 s is designated *nonsustained* (Fig. 277-2), whereas *sustained* VT persists longer than 30 s or is terminated by an active intervention, such as administration of an intravenous medication, external cardioversion, or pacing or a shock from an implanted cardioverter-defibrillator.

Monomorphic VT has the same QRS complex from beat to beat, indicating that the activation sequence is the same from beat to beat and that each beat likely originates from the same source (Fig. 277-3A). The initial site of ventricular activation largely determines the sequence of ventricular activation. Therefore, the QRS morphology of PVCs and monomorphic VT provides an indication of the site of origin within the ventricles (Fig. 277-4). The likely origin often suggests whether an arrhythmia is idiopathic or associated with structural disease. Arrhythmias that originate from the right ventricle or septum result in late activation of much of the left ventricle, thereby producing a prominent S wave in V_1 referred to as a left bundle branch block–like configuration. Arrhythmias that originate from the free wall of the left ventricle have a prominent positive deflection in V_1 , thereby producing a right bundle branch block–like morphology in V_1 . The frontal plane axis of the QRS is also useful. An axis that is directed inferiorly, as indicated by dominant R waves in leads II, III, and AVF, suggests initial activation of the cranial portion of the ventricle, whereas a frontal plane axis that is directed superiorly (dominant S waves in II, III, and AVF) suggests initial activation at the inferior wall.

Very rapid monomorphic VT has a sinusoidal appearance, also called *ventricular flutter*, because it is not possible to distinguish the QRS complex from the T wave (Fig. 277-3B). Relatively slow *sinusoidal* VTs have a wide QRS indicative of slowed ventricular conduction (Fig. 277-3C). Hyperkalemia, toxicity from excessive effects of drugs that blocks sodium channels (e.g., flecainide, propafenone, or

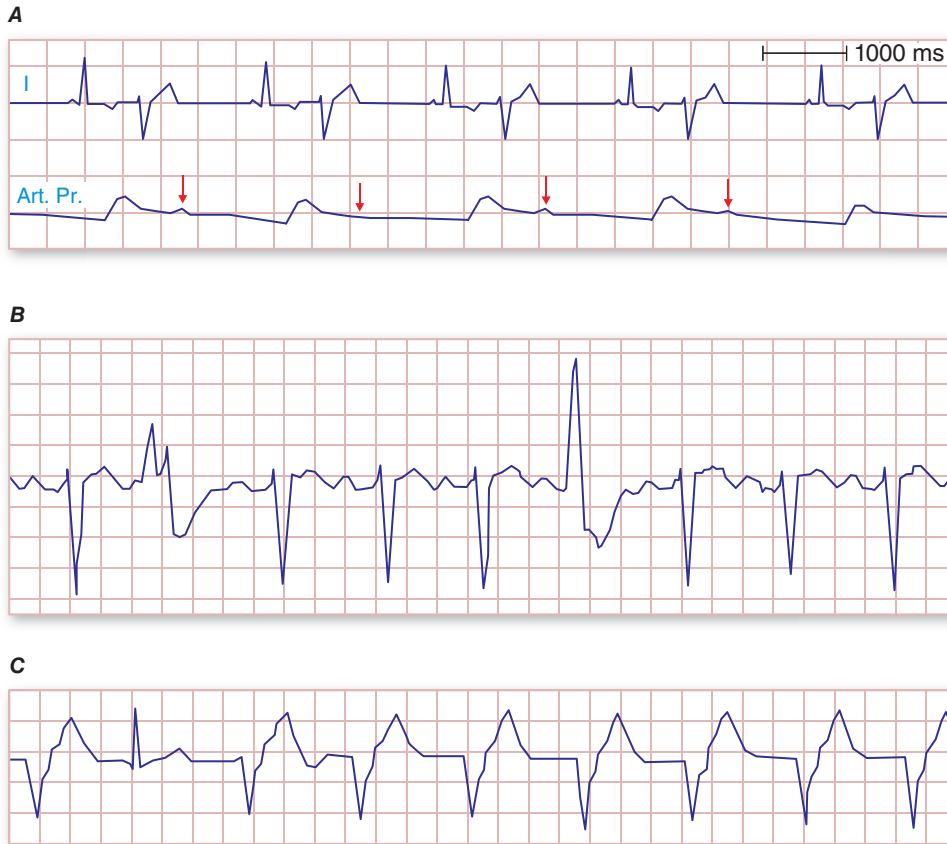


FIGURE 277-1 Examples of types of premature ventricular contractions (PVCs). **A.** Unifocal PVCs follow every sinus beat in a bigeminal frequency. Trace shows electrocardiogram lead 1 and arterial pressure (Art. Pr.). Sinus rhythm beats are followed by normal arterial waveform. The arterial pressure following premature beats is attenuated (*arrows*) and imperceptible to palpation. The pulse in this patient is registered at half the heart rate. **B.** Multifocal PVCs. The two PVCs shown have different morphologies. **C.** Example of accelerated idioventricular rhythm. The second QRS is a normally conducted beat. All other QRS complexes on this rhythm strip are ventricular due to accelerated idioventricular rhythm.

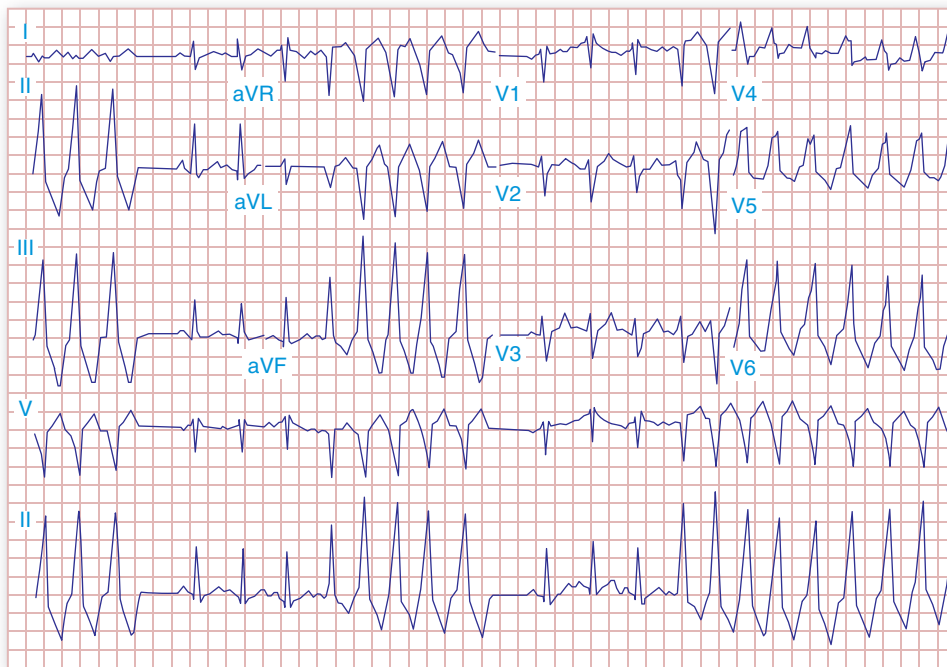


FIGURE 277-2 Repetitive monomorphic nonsustained ventricular tachycardia (VT) of right ventricular outflow tract origin. The VT has a left bundle branch block pattern with inferior axis with tall QRS complexes in the inferior leads.

tricyclic antidepressants), and severe global myocardial ischemia are causes.

Polymorphic VT has a continually changing QRS morphology indicating a changing ventricular activation sequence. Polymorphic VT that occurs in the context of congenital or acquired prolongation of the QT interval often has a waxing and waning QRS amplitude creating a “twisting about the points” appearance referred to as *Torsade de Pointes* (Fig. 277-3D).

Ventricular fibrillation (VF) has continuous irregular activation with no discrete QRS complexes (Fig. 277-3E). Monomorphic or polymorphic VT may transition to VF in susceptible patients.

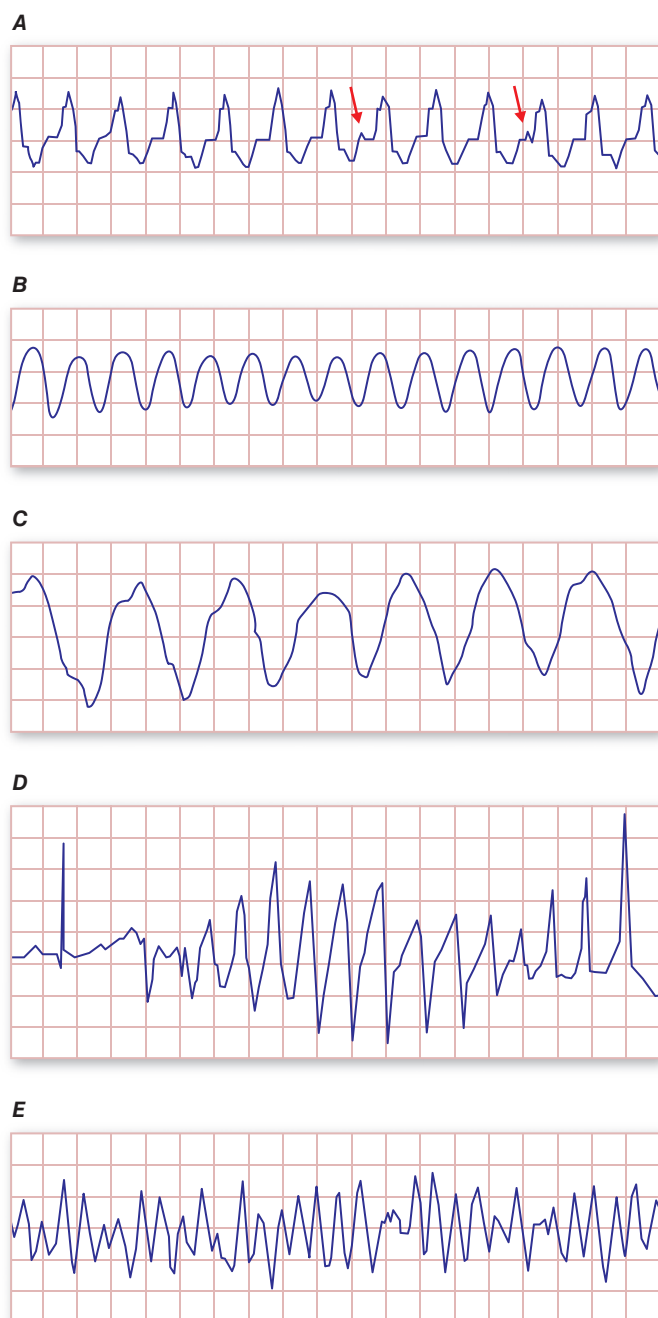


FIGURE 277-3 Examples of types of ventricular tachycardia (VT).

A. Monomorphic VT with dissociated P waves (*short arrows*). **B.** Ventricular flutter. **C.** Sinusoidal VT due to electrolyte disturbance or drug effects. **D.** Polymorphic VT resulting from prolongation of QT interval (*torsade de pointes* VT). **E.** Ventricular fibrillation.

CLINICAL MANIFESTATIONS AND DIAGNOSIS

Common symptoms of ventricular arrhythmias include palpitations, dizziness, exercise intolerance, episodes of lightheadedness, syncope, or sudden death. These arrhythmias can be asymptomatic and encountered unexpectedly as an irregular pulse or heart sounds on examination, or seen on a routine electrocardiogram (ECG), exercise test, or cardiac ECG monitoring.

Syncope is a concerning symptom that can be due to an episode of VT with hypotension. Syncope due to a ventricular arrhythmia often indicates that there is a significant risk for subsequent cardiac arrest and sudden death with arrhythmia recurrence. Although benign causes of syncope, such as reflex-mediated neurocardiogenic (vasovagal) syncope and orthostatic hypotension, are generally more common, it is important to consider the possibility of heart disease or a genetic syndrome causing VT. When these are suspected, hospitalization for further evaluation and monitoring is often appropriate.

Sustained VT may present with cardiac arrest, often with degeneration of the VT to VF. Occasionally a sustained VT will be hemodynamically tolerated and present with diminished exercise capacity or exacerbation of heart failure. Many patients who are at risk for VT have known heart disease and may have an implantable cardioverter-defibrillator (ICD). In patients with an ICD, spontaneous episodes of VT may elicit an episode of transient lightheadedness, palpitations, or syncope that may be followed by a shock from the ICD (see below).

The diagnosis of ventricular arrhythmias is established by recording of the arrhythmia on an ECG or, in some cases, initiation of the arrhythmia during an electrophysiologic study (Table 277-1). A 12-lead ECG of the arrhythmia should be obtained when possible and often provides clues to the potential site of origin and possible presence of underlying heart disease (see above). When the arrhythmia is intermittent with days to weeks between symptoms, prolonged ambulatory monitoring to capture the ECG at the time of symptoms is required to make the diagnosis. Continuous ambulatory monitoring or looping event recording monitors are options. Exercise testing should be considered in patients with exercise-induced symptoms.

APPROACH TO THE PATIENT:

Documented or Suspected Ventricular Arrhythmias

Initial assessment focuses on hemodynamic stability and evaluation for underlying heart disease. A family history of sudden death or cardiomyopathy suggests the possibility of a genetic basis for the arrhythmia and greater risk. The electrocardiogram can provide important clues. Patients with benign idiopathic arrhythmias usually have a completely normal ECG during sinus rhythm.

Cardiac imaging is warranted to assess ventricular function and look for evidence of depressed ventricular function indicative of a cardiomyopathy or ventricular hypertrophy that may indicate hypertrophic cardiomyopathy. Cardiac magnetic resonance imaging (MRI) with late gadolinium enhancement can detect areas of ventricular scar, which are usually present in patients who are at risk for sustained monomorphic VT (Fig. 277-5). Evaluation to exclude atherosclerotic coronary artery disease should be performed in patients at risk, guided by age and other risk factors.

SPECIFIC ARRHYTHMIAS

PVCs and Nonsustained VT Ventricular extrasystoles (Fig. 277-1A) can be due to automaticity or reentry (Chap. 278e). PVCs can be a sign of increased sympathetic tone; myocardial ischemia; hypoxia; electrolyte abnormalities, particularly hypokalemia; or underlying heart disease. During myocardial ischemia or in association with other heart disease, PVCs can be a harbinger of sustained VT or

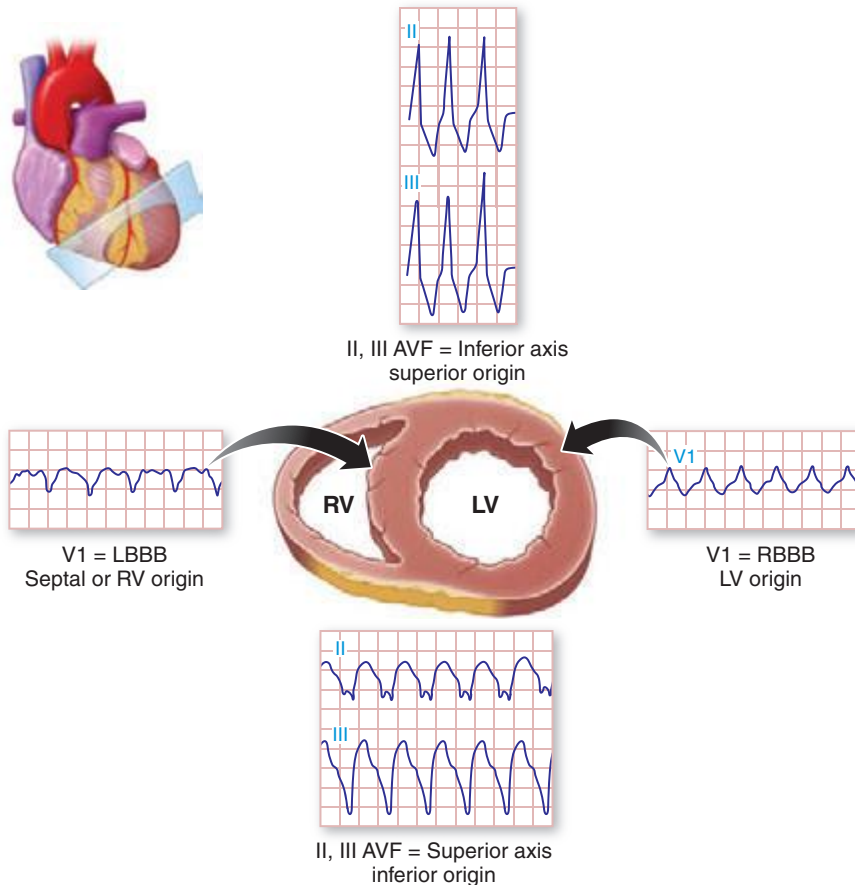


FIGURE 277-4 Site of VT origin based on QRS morphology. LBBB, left bundle branch block; LV, left ventricle; RBBB, right bundle branch block; RV, right ventricle.

TABLE 277-1 DIAGNOSTIC TESTS FOR VENTRICULAR ARRHYTHMIAS

I. 12-Lead ECG
A. Should be obtained for PVCs, nonsustained VT, and monomorphic VT when possible
B. QRS morphology suggests ventricular region of origin
V1 – dominant S = septum or RV
V1 – dominant R = LV
Superior axis = inferior wall origin
Inferior axis = outflow region or anterior wall
II. Ambulatory monitoring
A. 24- to 48-h continuous Holter monitor
Useful for evaluation of daily symptoms to quantitate PVCs
B. Event recorder: can be used for weeks at a time
Useful for evaluation of infrequent symptoms
Some require patient activation and will miss asymptomatic arrhythmias
III. Exercise testing
A. Useful for evaluating exercise-induced arrhythmias and symptoms
B. QT interval response to exercise may be abnormal in long QT syndrome
IV. Invasive electrophysiology study
A. Can establish definitive diagnosis of VT versus supraventricular tachycardia with aberrancy or ventricular preexcitation
B. Can provoke some arrhythmias that are otherwise infrequent
C. Allows potential catheter ablation
D. Procedural risks determined by vascular access, whether ablation is performed, and the location of the arrhythmia substrate

Abbreviation: RV, right ventricle. See text for other abbreviations.

VF. In patients with heart disease, a higher frequency of ectopy and complexity (couplets and nonsustained VT) are associated with more severe disease and, in those with heart failure, with increased mortality. However, suppression of these arrhythmias with antiarrhythmic drugs does not improve survival. In the absence of cardiac disease, PVCs and nonsustained VT generally have a benign prognosis. PVCs that occur at a bigeminal frequency may not generate sufficient cardiac output for a radial pulse and hence may register at rates half that of the heart rate (Fig. 277-1A). Very frequent PVCs can depress ventricular function (see below).

EVALUATION AND MANAGEMENT When encountered during acute illness or as a new finding, evaluation should focus on detection and correction of potential aggravating factors and causes, specifically myocardial ischemia, ventricular dysfunction, and electrolyte abnormalities, most commonly hypokalemia. Underlying heart disease should be defined.

The ECG characteristics of the arrhythmia are often suggestive of whether structural heart disease is present. PVCs with smooth uninterrupted contours and sharp QRS deflections suggest an ectopic focus in relatively normal myocardium, whereas broad notching and slurred QRS deflections suggest a diseased myocardial substrate. The most frequent site of origin for idiopathic ventricular arrhythmias is the right ventricular outflow tract, giving rise to PVCs or VT that have a left bundle branch block configuration, with an inferiorly directed frontal plane axis as discussed below (Fig. 277-2). However, QRS

morphology alone is not reliable as an indicator of disease or subsequent risk. Nonsustained VT is usually monomorphic with rates less than 200 beats/min and typically lasts less than 8 beats (Fig. 277-2). Nonsustained VT that is very rapid, polymorphic, or with a first beat that occurs prior to the peak of the T wave (“short-coupled”) is uncommon and should prompt careful evaluation for underlying disease or genetic syndromes associated with sudden death.

A family history of sudden death should prompt evaluation for genetic syndromes associated with sudden death, including cardiomyopathy, long QT syndrome, and arrhythmogenic right ventricular cardiomyopathy (see below). Any abnormality on the 12-lead ECG warrants further evaluation (Fig. 277-6). Repolarization abnormalities are seen in a number of genetically determined syndromes associated with sudden death, including the long QT syndrome, Brugada syndrome, arrhythmogenic right ventricular cardiomyopathy (ARVC), and hypertrophic cardiomyopathy. An echocardiogram is often necessary to assess ventricular function, wall motion abnormalities, and valvular heart disease. Cardiac magnetic resonance (CMR) imaging is also useful for this purpose and for the detection of ventricular scarring that is the substrate for sustained VT (Fig. 277-5). Exercise stress testing should be performed in patients with effort-related symptoms and in those at risk for coronary artery disease.

IDIOPATHIC PVCs AND NONSUSTAINED VT For PVC and nonsustained VT in the absence of structural heart disease or a genetic sudden death syndrome, no specific therapy is needed unless the patient has significant symptoms or evidence that frequent PVCs are depressing ventricular function (see below). Reassurance that the arrhythmia is benign is often sufficient to allow the patient to cope with the symptoms, which will often wax and wane in frequency over years. Avoiding stimulants, such as caffeine, is helpful in some patients. If symptoms require treatment, β -adrenergic blockers and nondihydropyridine calcium channel blockers (verapamil and diltiazem) are sometimes helpful (see Table 276-3). If these fail, more potent antiarrhythmic drugs or catheter

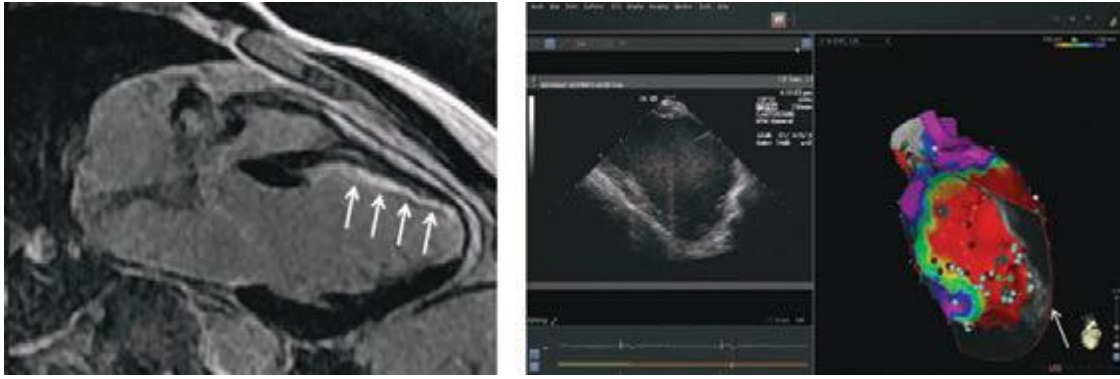


FIGURE 277-5 Imaging studies of the left ventricle (LV) used to assist ablation for ventricle tachycardia (VT). *Left panel* is a magnetic resonance image of a longitudinal section demonstrating thinning of the anterior wall and late gadolinium enhancement in a subendocardial scar (white arrows). The *middle panel* shows a two-dimensional image of the LV in long axis corresponding to the sector through the mid LV (arrow, *right panel*) obtained by an intracardiac echo probe positioned in the right ventricle. An electroanatomic three-dimensional map of the LV in the left anterior oblique projection is displayed in the right panel. The purple color depicts areas of normal voltage (>1.5 mV). Blue, green, and yellow represent progressively lower voltages with the red areas indicating scar (<0.5 mV). Channels of viable myocardium with slow conduction within the scar are identified with the light blue dots. Areas of ablation delivered to regions involved in reentrant VT are indicated by maroon dots.

ablation can be considered. The antiarrhythmic agents flecainide, propafenone, mexiletine, and amiodarone can be effective, but the potential for side effects warrants careful consideration. Catheter ablation can be effective if the arrhythmia occurs with sufficient frequency or is readily provoked such that its origin can be identified for ablation in a similar manner to that for idiopathic monomorphic VT as discussed below. Benefit must be carefully weighed against the procedure-related risks (see below).

PVCs AND NONSUSTAINED VT ASSOCIATED WITH ACUTE CORONARY SYNDROMES
During and early after acute myocardial infarction (MI), PVCs and nonsustained VT are common and can be an early manifestation of ischemia and a harbinger of subsequent VF. Treatment with β -adrenergic blockers and correction of hypokalemia and hypomagnesemia reduce the risk of VF. Routine administration of the

antiarrhythmic drugs such as lidocaine has not been shown to reduce mortality and is not indicated for suppression of PVCs or asymptomatic nonsustained VT.

Following recovery from acute MI, frequent PVCs (typically >10 PVCs per hour), repetitive PVCs with couplets, and nonsustained VT are markers for depressed ventricular function and increased mortality, but routine antiarrhythmic drug therapy to suppress these arrhythmias is not warranted. Treatment with the sodium channel blocker flecainide increased mortality. Amiodarone therapy reduces sudden death, but does not improve total mortality. Therefore, amiodarone is an option for treatment of symptomatic arrhythmias in this population when the potential benefit outweighs its potential toxicities. β -Adrenergic blockers reduce sudden death but have limited effect on spontaneous arrhythmias.

For survivors of an acute MI, an ICD reduces mortality in certain high-risk groups: patients who have survived >40 days after the acute MI and have a left ventricular (LV) ejection fraction of ≤ 0.30 or who have an ejection fraction <0.35 and have symptomatic heart failure (functional class II or III); and patients >5 days after MI who have a reduced LV ejection fraction, nonsustained VT, and inducible sustained VT or VF on electrophysiologic testing. ICDs do not reduce mortality when routinely implanted soon after MI or in patients after recent coronary artery revascularization surgery.

PVCs AND NONSUSTAINED VT ASSOCIATED WITH DEPRESSED VENTRICULAR FUNCTION AND HEART FAILURE

PVCs and nonsustained VT are common in patients with depressed ventricular function and heart failure and are markers for disease severity and increased mortality, but antiarrhythmic drug therapy to suppress these arrhythmias has not been shown to improve survival. Antiarrhythmic drugs whose major action is blockade of the cardiac sodium channel (flecainide, propafenone, mexiletine, quinidine, and disopyramide) are avoided in patients with structural heart disease because of a risk of proarrhythmia, negative inotropic effects, and increased mortality. Therapy with the potassium channel blockers, e.g., dofetilide, does not reduce mortality. Amiodarone suppresses ventricular ectopy and reduces sudden death but does not improve overall survival. ICDs are the major

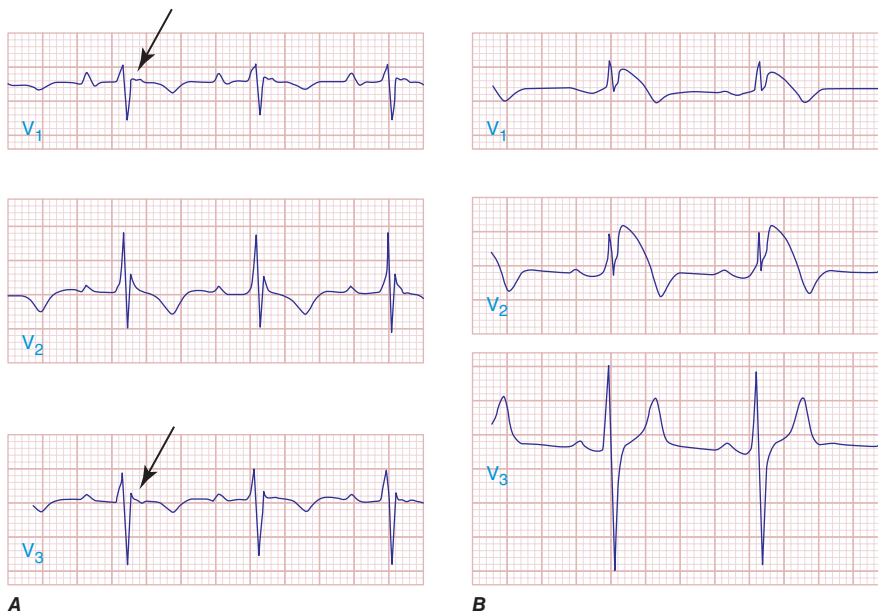


FIGURE 277-6 Precordial chest leads V_1 – V_3 showing typical abnormalities of arrhythmogenic right ventricular cardiomyopathy (ARVC) (A) and Brugada syndrome (B).

In ARVC, there is T inversion and delayed ventricular activation manifest as epsilon waves (arrows). Panel B shows ST elevation in V_1 and V_2 typical of the Brugada syndrome. (Figures reproduced from F Marchlinski: *The tachyarrhythmias*. In Longo DL et al [eds]: *Harrison's Principles of Internal Medicine*, 18th edition. New York, McGraw-Hill, 2012, pp 1878–1900.)

1494 therapy to protect against sudden death in patients at high risk and are recommended for those with LV ejection fraction <0.35 and New York Heart Association class II and III heart failure, in whom they reduce mortality by 20%, from 36% to 29%, over 5 years.

OTHER CARDIAC DISEASES Ventricular ectopy is associated with increased mortality in patients with *hypertrophic cardiomyopathy* (Chap. 287) or with *congenital heart disease* (Chap. 282) associated with right ventricular or LV dysfunction. In these patients, management is similar to that for patients with ventricular dysfunction. Pharmacologic suppression of the arrhythmia has not been shown to improve mortality. ICDs are indicated for patients considered at high risk for sudden cardiac death.

PVC-INDUCED VENTRICULAR DYSFUNCTION Very frequent ventricular ectopy and repetitive nonsustained VT (Fig. 277-2) can depress ventricular function, possibly through an effect similar to chronic tachycardia or by inducing ventricular dyssynchrony. Depression of ventricular function rarely occurs unless PVCs account for more than 10–20% of total beats over a 24-h period. Often the PVCs are idiopathic and unifocal, most commonly originating from the LV papillary muscles or outflow tract regions where they can be targeted for ablation. The distinction between PVC-induced ventricular dysfunction as compared to a cardiomyopathic process causing ventricular dysfunction and arrhythmia is difficult and in some cases can be made only retrospectively by observing an improvement in ventricular function after the arrhythmia is suppressed with an antiarrhythmic drug, such as amiodarone, or by catheter ablation.

Idioventricular Rhythms Three or more ventricular beats at a rate slower than 100 beats/min are termed *idioventricular rhythm* (Fig. 277-1C). Automaticity is the likely mechanism. Idioventricular rhythms are common during acute MI (Chap. 295) and may emerge during sinus bradycardia. Atropine may be administered to increase the sinus rates if the loss of atrioventricular synchrony leads to hemodynamic compromise. This rhythm is also common in patients with cardiomyopathies or sleep apnea. It can also be idiopathic, often emerging when the sinus rate slows during sleep. Therapy should target any underlying cause and correction of bradycardia. Specific therapy for asymptomatic idioventricular rhythm is not necessary.

Sustained Monomorphic VT Sustained monomorphic VT presents as a wide QRS tachycardia that has the same QRS configuration from beat to beat, indicating an identical sequence of ventricular depolarization for each beat (Fig. 277-3A). VT originates from a stable focus or reentry circuit. In structural heart disease, the substrate is often an area of patchy replacement fibrosis due to infarction, inflammation, or prior cardiac surgery that creates anatomical or functional reentry pathways (Fig. 277-5). Less commonly, VT is related to reentry or automaticity in a diseased Purkinje system. In the absence of structural heart disease, idiopathic VT can present as sustained monomorphic VTs that are due to focal automaticity or reentry involving a portion of the Purkinje system.

The clinical presentation can vary depending on the rate of the arrhythmia, underlying cardiac function, and autonomic adaptation in response to the arrhythmia. Whereas patients with normal cardiac function might tolerate rapid VTs, those with severe LV dysfunction often experience symptoms of hypotension, even if VT is not particularly fast. Monomorphic VT may deteriorate to VF, which may be the initial cardiac rhythm recorded at the time of resuscitation.

DIAGNOSIS Sustained monomorphic VT has to be distinguished from other causes of uniform wide QRS tachycardia. These include supraventricular tachycardia with left or right bundle branch block aberrant conduction, supraventricular tachycardias conducted to the ventricles over an accessory pathway (Chap. 276), and rapid cardiac pacing in a patient with a pacemaker or defibrillator. In the presence of known heart disease, VT is the most likely diagnosis of a wide QRS tachycardia. Hemodynamic stability during the arrhythmia does not exclude VT. A number of ECG criteria have been evaluated. The presence of AV dissociation is usually a reliable marker for VT (Fig. 277-7), but P waves can be difficult to define. A P wave following each QRS does not

VT versus Supraventricular Tachycardia with Aberrancy

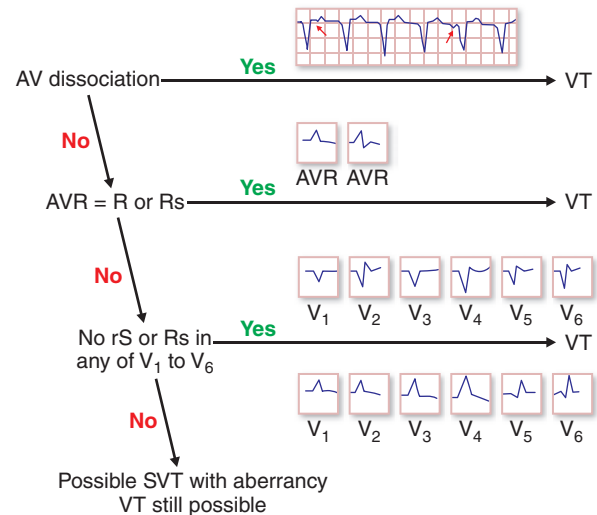


FIGURE 277-7 Algorithm for differentiation of ventricular tachycardia (VT) from supraventricular tachycardia (SVT) with aberrancy. AV, atrioventricular.

exclude VT because 1:1 conduction from ventricle to atrium can occur. A monophasic R wave or Rs complex in AVR or concordance from V₁ to V₆ of monophasic R or S waves is also relatively specific for VT (Fig. 277-7). Other QRS morphology criteria have also been described, but all have limitations and are not very reliable in patients with severe heart disease. In patients with known bundle branch block, the same QRS morphology during tachycardia as during sinus rhythm suggests supraventricular tachycardia rather than VT, but is not absolutely reliable. An electrophysiologic study is sometimes required for definitive diagnosis. Rarely, noise and movement artifacts on telemetry recordings can simulate VT; prompt recognition can avoid unnecessary tests and interventions.

When LV function is depressed or there is evidence of structural myocardial disease, scar-related reentry is the most likely diagnosis. Scars are suggested by pathologic Q waves on the ECG, segmental left or right ventricular wall motion abnormalities on echocardiogram or nuclear imaging, and areas of delayed gadolinium enhancement during MRI (Fig. 277-5).

TREATMENT AND PROGNOSIS Initial management follows Advanced Cardiac Life Support (ACLS) guidelines. If hypotension, impaired consciousness, or pulmonary edema is present, QRS synchronous electrical cardioversion should be performed, ideally after sedation if the patient is conscious. For stable tachycardia, a trial of adenosine is reasonable, as this may clarify a supraventricular tachycardia with aberrancy (Chap. 276). Intravenous amiodarone is the drug of choice if heart disease is present. Following restoration of sinus rhythm, hospitalization and evaluation to define underlying heart disease are required. Assessment of cardiac biomarkers for evidence of MI is appropriate, but acute MI is rarely a cause of sustained monomorphic VT, and elevations in troponin or creatine kinase (CK)-MB are more likely to indicate myocardial damage that is secondary to hypotension and ischemia from the VT. Subsequent management is determined by the underlying heart disease and frequency of VT. If VT recurs frequently or is incessant, administration of antiarrhythmic medications or catheter ablation may be required to restore stability. More commonly, sustained monomorphic VT occurs as an isolated episode, but with a risk of recurrence. ICDs are usually considered for VT associated with structural heart disease.

Sustained Monomorphic VT in Specific Diseases • CORONARY ARTERY DISEASE Patients who present with sustained VT associated with coronary artery disease typically have a history of prior large MI and present years after

TABLE 277-2 VENTRICULAR ARRHYTHMIAS ASSOCIATED WITH DIFFERENT FORMS OF HEART DISEASE

I. Idiopathic VT without structural heart disease
A. Outflow tract origin
1. RV outflow tract: left bundle branch block pattern with inferior axis (tall QRS in inferior leads) and late transition in the precordial leads
2. LV outflow tract: prominent R in V ₁ with inferior axis
B. Left posterior fascicular VT
1. Right bundle branch block pattern with left axis deviation (most common)
II. Ischemic cardiomyopathy
A. Monomorphic VT is common with prior large myocardial infarction
B. Polymorphic VT and VF should prompt ischemia evaluation
III. Nonischemic cardiomyopathy
A. Polymorphic VT and VF more common but fibrotic scars can cause monomorphic VT especially with sarcoidosis and Chagas' disease
IV. Arrhythmogenic right ventricular cardiomyopathy
A. Monomorphic VT usually of right ventricular origin (left bundle branch morphology)
B. Polymorphic VT and VF can occur independently or through degeneration of monomorphic VT
V. Repaired tetralogy of Fallot
A. Monomorphic VT of right ventricular origin (usually left bundle branch morphology)
VI. Hypertrophic cardiomyopathy
A. Polymorphic VT or ventricular fibrillation
B. Less commonly, monomorphic VT associated with myocardial scars
VIII. Genetic arrhythmia syndromes
A. Long QT syndrome: torsade de pointes VT
B. Brugada syndrome: VF
C. Catecholaminergic polymorphic VT: polymorphic VT or bidirectional VT
D. Short QT syndrome: ventricular fibrillation
E. Early repolarization syndrome: polymorphic VT or VF

Abbreviation: RV, right ventricle. See text for other abbreviations.

the acute infarct with a remodeled ventricle and markedly depressed LV function. Even when there is biomarker evidence of acute MI, a preexisting scar from previous MI should be suspected as the cause of the VT. Infarct scars provide a durable substrate for sustained VT, and up to 70% of patients have a recurrence of the arrhythmia within 2 years. Scar-related reentry is not dependent on recurrent acute myocardial ischemia, so coronary revascularization cannot be anticipated to prevent recurrent VT, even when it may be appropriate for other indications. Depressed ventricular function, which is a risk factor for sudden death, is usually present. Implantation of an ICD is warranted for most patients provided that there is a reasonable expectation of survival with acceptable functional status for the next year after recovery from the VT episode. ICDs reduce annual mortality from 12.3% to 8.8% and lower arrhythmic deaths by 50% in patients with hemodynamically significant sustained VT or a history of cardiac arrest compared with pharmacologic therapy. Chronic amiodarone therapy may be considered for patients who are not candidates for or who decline ICD placement.

Following ICD implantation, patients remain at risk for heart failure, recurrent ischemic events, and recurrent VT, with a 5-year mortality that exceeds 30%. Attention to therapies with survival benefit, including β -adrenergic blocking agents, angiotensin-converting enzyme inhibitors, and statins, is important. Patients with frequent symptomatic recurrences of VT require antiarrhythmic drug therapy or catheter ablation.

NONISCHEMIC DILATED CARDIOMYOPATHY Sustained monomorphic VT associated with nonischemic cardiomyopathy is usually due to scar-related reentry. The etiology of scar is often unclear, but progressive replacement fibrosis is the likely cause. On cardiac MRI, scars are detectable as areas of delayed gadolinium enhancement and are more often intramural or subepicardial in location as compared with patients with prior MI. Scars that cause VT are often located adjacent to a valve annulus and can occur in either ventricle. Any cardiomyopathic process can cause scars and VT, but cardiac sarcoidosis (Chap. 390) and Chagas' disease (Chap. 252) are particularly associated with monomorphic VT (Table 277-2). An ICD is usually indicated with additional drugs or ablation for control of recurrent VT.

MONOMORPHIC VT IN ARVC ARVC (Chap. 287) is a rare genetic disorder most commonly due to mutations in genes encoding for cardiac desmosomal proteins. Approximately 50% have a familial transmission with autosomal dominant inheritance. A less common, autosomal recessive form is associated with cardiocutaneous syndromes that include Naxos disease and Carvajal syndrome. Patients typically present between the second and fifth decade with palpitations, syncope, or cardiac arrest owing to sustained monomorphic VT, although polymorphic VT can also occur. Fibrosis and fibro-fatty replacement most commonly involve the right ventricular myocardium and provide the substrate for reentrant VT that usually has a left bundle branch block–like configuration, consistent with the right ventricular origin. The sinus rhythm ECG suggests the disease in more than 85% of patients, most often showing T-wave inversions in V₁–V₃ (Fig. 277-6). Delayed activation of the right ventricle may cause a widened QRS (≥ 110 ms) in the right precordial leads and a prolonged S-wave upstroke in those leads, and occasionally a deflection at the end of the QRS known as an *epsilon wave* (Fig. 277-6). Cardiac imaging may show right ventricular enlargement or areas of abnormal motion or reveal areas of scar on CMR imaging with gadolinium. The monomorphic VT of early ARVC can sometimes be difficult to differentiate from idiopathic right ventricular outflow tract VT.

LV involvement can occur and occasionally precede manifest right ventricular disease. Heart failure is rare except in late stages, and survival to advanced age can be anticipated provided that VT can be controlled. An ICD is recommended. When VT is exercise-induced, it may respond to β -adrenergic blockers and limiting exercise. Sotalol, amiodarone, and catheter ablation have been used to reduce recurrences. Ablation targets are often located in the subepicardium of the RV.

TETRALOGY OF FALLOT VT occurs in 3–14% of patients late after repair of tetralogy of Fallot (Chap. 282) and contributes to a 2% per decade risk of

sudden death. Monomorphic VT is due to reentry around areas of surgically created scar in the RV (Table 277-2). Factors associated with VT risk include age >5 years at the time of repair, high-grade ventricular ectopy, inducible VT on an electrophysiologic study, abnormal right ventricular hemodynamics, and sinus rhythm QRS duration >180 ms. An ICD is usually warranted for patients who have a spontaneous episode of VT, but criteria for a prophylactic ICD in other patients have not been established. Catheter ablation is used to control recurrent episodes.

BUNDLE BRANCH REENTRY VT Reentry through the Purkinje system occurs in approximately 5% of patients with monomorphic VT in the presence of structural heart disease. The reentry circuit typically revolves retrograde via the left bundle and anterograde down the right bundle, thereby producing VT that has a left bundle branch block configuration. Catheter ablation of the right bundle branch abolishes this VT. Bundle branch reentry is usually associated with severe underlying heart disease. Other scar-related VTs are often present and often require additional therapy or ICD implantation.

IDIOPATHIC MONOMORPHIC VT Idiopathic VT in patients without structural heart disease usually presents with palpitations, lightheadedness, and occasionally syncope, often provoked by sympathetic stimulation during exercise or emotional upset. The QRS morphology of the arrhythmia suggests the diagnosis (see below). The sinus rhythm ECG is normal. Cardiac imaging shows normal ventricular function and no evidence of ventricular scar. Occasionally a patient with structural heart disease is found to have concomitant idiopathic VT, unrelated to the structural disease. Sudden death is rare.

Outflow tract VTs originate from a focus, usually with features consistent with triggered automaticity. The arrhythmia may present with

TABLE 277-3 CAUSES OF QT PROLONGATION AND TORSADE DE POINTES VENTRICULAR TACHYCARDIA

1. Congenital long QT syndromes (see text for details)	Antihistamines (histamine 1-receptor antagonists)
Long QT syndrome type 1: Reduced repolarizing current I_{Ks} due to mutation in <i>KCNQ1</i> gene	Terfenadine, astemizole, diphenhydramine, hydroxyzine
Long QT syndrome type 2: Reduced repolarizing current I_{Kr} due to mutation in <i>KCNH2</i> gene	Cholinergic antagonists: Cisapride, organophosphates
Long QT syndrome type 3: Delayed inactivation of the I_{Na} due to mutations in <i>SCN5A</i> gene	Citrate (massive blood transfusions)
Others: Several other types of long QT syndromes have been described; long QT types 1, 2, and 3 account for 80–90% of cases	Cocaine
2. Acquired prolongation of QT interval	Methadone
Electrolyte abnormalities	Fluoxetine (in conjunction with other drugs that prolong QT)
Hypokalemia	Cardiac conditions
Hypomagnesemia	Myocardial ischemia and infarction
Hypocalcemia	Myocarditis
Drugs	Marked bradycardia
Antiarrhythmic drugs	Stress cardiomyopathy
Class IA: Quinidine, disopyramide, procainamide	Endocrine disorders
Class III: Sotalolol, amiodarone (QT prolongation common but torsade ventricular tachycardia is rare), ibutilide, dofetilide, almokalant	Hypothyroidism
Antibiotics	Hyperparathyroidism
Macrolides: Erythromycin, clarithromycin, azithromycin	Pheochromocytoma
Fluoroquinolones: Levofloxacin, moxifloxacin, gatifloxacin	Hyperaldosteronism
Trimethoprim-sulfamethoxazole	Intracranial disorders
Clindamycin	Subarachnoid hemorrhage
Pentamidine	Thalamic hematoma
Chloroquine	Cerebrovascular accident
Antifungals: Ketoconazole, itraconazole	Encephalitis
Antivirals: Amantadine	Head injury
Antipsychotics	Nutritional disorders
Haloperidol, phenothiazines, thioridazine, trifluoperazine, sertindole, zimelidine, ziprasidone	Anorexia nervosa
Tricyclic and tetracyclic antidepressants	Starvation
	Liquid protein diets
	Gastroplasty and ileojejunum bypass
	Celiac disease

sustained VT, nonsustained VT, or PVCs often provoked by exercise or emotional upset. Repeated bursts of nonsustained VTs, which may occur incessantly, are known as repetitive monomorphic VTs and can cause tachycardia-induced cardiomyopathy with depressed ventricular function that recovers after suppression of the arrhythmia (Fig. 277-2). Most originate in the right ventricular outflow tract, which gives rise to VT that has a left bundle branch block configuration in V_1 and an axis that is directed inferiorly, with tall R waves in leads II, III, and AVF (Fig. 277-2). Idiopathic VT can also arise in the LV outflow tract or in sleeves of myocardium that extend along the aortic root. LV origin is suspected when lead V_1 or V_2 has prominent R waves. Although this typical outflow tract QRS morphology favors idiopathic VT, some cardiomyopathies, notably ARVC, can cause PVCs or VT from this region. Excluding these diseases is an initial focus of evaluation.

LV intrafascicular VT presents with sustained VT that has a right bundle branch block–like configuration. It is often exercise-induced and occurs more often in men than women. The mechanism is reentry in or near the septal ramifications of the LV Purkinje system. This VT can be terminated by intravenous administration of verapamil.

MANAGEMENT OF IDIOPATHIC VT Treatment is required for symptoms or when frequent or incessant arrhythmias depress ventricular function. β -Adrenergic blockers are first-line therapy. Nondihydropyridine calcium channel blockers (diltiazem and verapamil) are sometimes effective. Catheter ablation is warranted for severe symptoms or when beta blockers or calcium channel blockers are not effective or not desired. Efficacy and risks of catheter ablation vary with the specific site of origin of the VT, being most favorable for arrhythmias originating in the right ventricular outflow tract.

LV fascicular VT can be terminated by intravenous administration of verapamil, although chronic therapy with oral verapamil is not

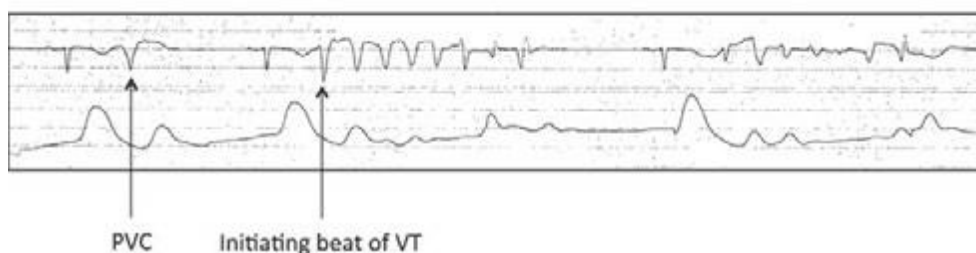
always effective. Catheter ablation is recommended if β -adrenergic blockers or calcium channel blockers are ineffective or not desired.

Polymorphic VT Sustained polymorphic VT can be seen with any form of structural heart disease (Table 277-2). However, unlike sustained monomorphic VT, polymorphic VT does not always indicate a structural abnormality or focus of automaticity. Reentry with continually changing reentrant paths, spiral wave reentry, and multiple automatic foci are potential mechanisms (Chap. 278e). Sustained polymorphic VT usually degenerates into VF. Polymorphic VT is typically seen in association with acute MI or myocardial ischemia, ventricular hypertrophy, and a number of genetic mutations that affect cardiac ion channels (Table 277-3).

POLYMORPHIC VT ASSOCIATED WITH ACUTE MI/MYOCARDIAL ISCHEMIA Acute MI or ischemia is a common cause of polymorphic VT and should be the initial consideration in management. Approximately 10% of patients with acute MI develop VT that degenerates to VF, related to reentry through the infarct border zone. The risk is greatest in the first hour of acute MI. Following resuscitation as per the ACLS guidelines, management is as for acute MI (Chap. 295). β -Adrenergic blockers, correction of electrolyte abnormalities, and prompt myocardial reperfusion are required. Repeated episodes of polymorphic VT suggest ongoing myocardial ischemia and warrant assessment of adequacy of myocardial reperfusion. Polymorphic VT and VF that occur within the first 48 h of acute MI are associated with greater in-hospital mortality, but those who survive past hospital discharge are not at increased risk for arrhythmic sudden death. Long-term therapy for postinfarct ventricular arrhythmia is determined by residual LV function, with an ICD being indicated for persistent severe LV dysfunction (LV ejection fraction <0.35).



A



B

FIGURE 277-8 Electrocardiogram (ECG) of a patient with prolonged QT and episodes of torsade de pointes ventricular tachycardia (VT). **A.** Twelve-lead ECG showing a heart rate of 54, anterior wall T inversion, and QT interval of 600 ms. The corrected QT interval (QTc) is 585 ms. **B.** Telemetry ECG tracing with digital pulse waveform demonstrating bursts of torsade de pointes VT. The initiating sequence of the VT is characteristic, with a PVC inducing a pause followed by a sinus beat that had a longer QT and interruption of the T wave by the PVC that is the first beat of VT. The VT is self-terminating in this case.

REPOLARIZATION ABNORMALITIES AND GENETIC ARRHYTHMIA SYNDROMES • Acquired long QT Abnormal prolongation of the QT interval is associated with the polymorphic VT Torsade de Pointes (Fig. 277-8). The VT often has a characteristic initiation sequence of a premature ventricular beat that induces a pause, followed by a sinus beat that has a longer QT interval and interruption of the T wave by the PVC that is the first beat of the polymorphic VT. This characteristic initiation is termed “pause-dependent” (Fig. 277-8). Causes of QT prolongation include electrolyte abnormalities, bradycardia, and a number of medications that block repolarizing potassium currents, notably the antiarrhythmic drugs sotalol, dofetilide, and ibutilide, but also a number of other medications used for noncardiac diseases, including erythromycin, pentamidine, haloperidol, phenothiazines, and methadone (Table 277-3). Individual susceptibility may be related to genetic polymorphisms or mutations that influence repolarization.

Patients typically present with near-syncope, syncope, or cardiac arrest. Sustained episodes degenerate to VF requiring defibrillation. PVCs and nonsustained VT often precede episodes of sustained VT. Intravenous administration of 1–2 g of magnesium sulphate usually suppresses recurrent episodes. If magnesium alone is ineffective, increasing heart rate with isoproterenol infusion or pacing, to a rate of 100–120 beats/min as required to suppress PVCs, usually suppresses VT recurrences. These maneuvers allow time for correction of associated electrolyte disturbance (hypokalemia and hypocalcemia) and bradycardia and removal of any causative drugs (Table 277-3). Drug interactions that elevate levels of the offending agent are often a precipitating factor. Patients who experience a polymorphic VT induced by QT prolongation should be considered to have a susceptibility to the arrhythmia and should avoid all future exposure to medications known to prolong the QT interval.

Congenital long QT syndrome The congenital long QT syndrome (LQTS) is caused by mutations in genes coding for cardiac ion channels responsible for ventricular repolarization. The corrected QT (QTc) is typically prolonged to greater than 440 ms in men and 460 ms in women. Symptoms are due to Torsade de Pointes VT (Fig. 277-8). Several forms of congenital LQTS have been identified, but three groups of mutations that lead to LQTS type 1 (LQTS-1), LQTS type 2 (LQTS-2), or LQTS type 3 (LQTS-3) account for 90% of cases. The most frequently encountered mutations, *LQTS1* and *LQTS2*, are due to abnormalities of potassium channels, but mutations affecting the sodium channel (*LQTS3*) and calcium channels have also been described (Table 277-3).

Patients often present with syncope or cardiac arrest, usually during childhood. In LQTS-1, episodes tend to occur during exertion, particularly swimming. In LQTS-2, sudden auditory stimuli or emotional upset predispose to events. In LQTS-3, sudden death during sleep is a notable feature. Asymptomatic patients may be discovered in the course of family screening or on a routine ECG. Genotyping can be helpful for family screening and to provide reassurance regarding the diagnosis. Correlations of genotype with risk and response to therapy are beginning to emerge. In most patients with LQTS-1 or LQTS-2, adequate doses of beta blocker therapy (the non-selective agents nadolol or propranolol) are sufficient protection from arrhythmia episodes. Markers of increased risk include QTc interval exceeding 0.5 s, female gender, and a history of syncope or cardiac arrest. Recurrent syncope despite beta blocker therapy or a high-risk profile merits consideration of an ICD. Avoidance of QT-prolonging drugs is critical for all patients with LQTS, including those who are genotype positive but have normal QT intervals.

Short QT syndrome Short QT syndrome is very rare compared to LQTS. The QTc is shorter than 0.36 s, and usually less than 0.3 s. The genetic abnormality causes a gain of function of the potassium channel (I_{Kr}) or reduced inward depolarizing currents. The abnormality is associated with atrial fibrillation, polymorphic VT, and sudden death.

Brugada syndrome Brugada syndrome is a rare syndrome characterized by >0.2 mV of ST-segment elevation with a coved ST segment and negative T wave in more than one anterior precordial lead (V_1 – V_3) (Fig. 277-6) and episodes of syncope or cardiac arrest due to polymorphic VT in the absence of structural heart disease. Cardiac arrest may occur during sleep or be provoked by febrile illness. Males are more commonly affected than females. Mutations involving cardiac sodium channels are identified in approximately 25% of cases. Distinction from patients with similar ST elevation owing to LV hypertrophy, pericarditis, myocardial ischemia or MI hyperkalemia, hypothermia, right bundle branch block, and ARVC is often difficult. Furthermore, the characteristic ST-segment elevation can wax and wane over time and may become pronounced during acute illness and fever. Administration of the sodium channel blocking drug flecainide, ajmaline, or procainamide can augment or unmask ST elevation in affected individuals. An ICD is indicated for individuals who have had unexplained syncope or been resuscitated from cardiac arrest.

1498 Quinidine has been used successfully to suppress frequent episodes of VT.

Early repolarization syndrome Patients resuscitated from VF who have no structural heart disease or other identified abnormality have a higher prevalence of J-point elevation with notching in the terminal QRS. A family history of sudden death is present in some patients, suggesting a potential genetic basis. J-point elevation is also seen in some patients with the Brugada syndrome and associated with a higher risk of arrhythmias. An ICD is recommended for those who have had prior cardiac arrest. It should be noted that J-point elevation is commonly seen as a normal variant, and in the absence of specific symptoms, the clinical relevance is not known.

Catecholaminergic polymorphic VT This rare familial syndrome is due to mutations in the cardiac ryanodine receptor and, less commonly, the sarcoplasmic calcium binding protein, calsequestrin 2. These mutations result in abnormal sarcoplasmic calcium handling and polymorphic ventricular arrhythmias that resemble those seen with digitalis toxicity. The VT is polymorphic or has a characteristic alternating QRS morphology termed bidirectional VT. Patients usually present during childhood with exercise- or emotion-induced palpitations, syncope, or cardiac arrest. β -Adrenergic blockers (e.g., nadolol and propranolol) and an ICD are recommended. Verapamil, flecainide, or surgical left cardiac sympathetic denervation reduces or prevents recurrent VT in some patients.

Hypertrophic cardiomyopathy (HCM) HCM is the most common genetic cardiovascular disorder, occurring in 1 in 500 individuals, and is a prominent cause of sudden death before the age of 35 years (Chap. 287). Sudden death can be due to polymorphic VT/VF. Rarely, sustained monomorphic VT occurs related to areas of ventricular scar. Risk factors include young age, nonsustained VT, failure of blood pressure to increase during exercise, recent (within 6 months) syncope, ventricular wall thickness >3 cm, and possibly the severity of LV outflow obstruction. An ICD is generally indicated for high-risk patients, but the specific risk profile warranting an ICD continues to be debated. Surgical myectomy, performed to relieve outflow obstruction, has been associated with a sudden death rate of less than 1% per year. The reported annual rate of sustained VT or sudden death after transcatheter ethanol septal ablation done to relieve outflow obstruction has been reported to range between 1 and 5%.

Genetic dilated cardiomyopathies Genetic dilated cardiomyopathies account for 30–40% of cases of nonischemic dilated cardiomyopathies. Some are associated with muscular dystrophy. Autosomal dominant, recessive, X-linked, and mitochondrial inheritance patterns are recognized. Mutations in genes coding for structural proteins of the nuclear lamina (lamin A and C) and the *SCN5A* gene are particularly associated with conduction system disease and ventricular arrhythmias. Patients can experience polymorphic VT and cardiac arrest or develop areas of scar causing sustained monomorphic VT. ICDs are recommended for those who have had a sustained VT or are at high risk due to significantly depressed ventricular function (LV ejection fraction of ≤ 0.35 and associated with heart failure) or a malignant family history of sudden death.

Ventricular Fibrillation VF is characterized by disordered electrical ventricular activation without identifiable QRS complexes (Fig. 277-3E). Spiral wave reentry and multiple circulating reentry wavefronts are possible mechanisms. Sustained polymorphic or monomorphic VT that degenerates to VF is a common cause of out-of-hospital cardiac arrest. Treatment follows ACLS guidelines with defibrillation to restore sinus rhythm. If resuscitation is successful, further evaluation is performed to identify and treat underlying heart disease and potential causes of the arrhythmia, including the possibility that monomorphic or polymorphic VT could have initiated VF. If a transient reversible cause such as acute MI is not identified, therapy to reduce the risk of sudden death with an ICD is often warranted. Chronic amiodarone therapy may be considered for individuals who are not ICD candidates.

Incessant VT and Electrical Storm VT is incessant when it continues to recur shortly after electrical, pharmacologic, or spontaneous conversion to sinus rhythm. “VT storm” or “electrical storm” refers to three or more separate episodes of VT within 24 h, most commonly encountered in patients with ICDs. Slow incessant VT is sometimes asymptomatic, but can cause heart failure or tachycardia-induced cardiomyopathy. More commonly, these presentations are life-threatening and require emergent therapy. Measures to reduce sympathetic tone, including β -adrenergic blockade, sedation, and general anesthesia, have been used effectively. Intravenous administration of amiodarone and lidocaine can be effective for suppression. Urgent catheter ablation can be lifesaving.

TREATMENT VENTRICULAR ARRHYTHMIAS

ANTIARRHYTHMIC DRUGS

Use of antiarrhythmic drugs is based on consideration of the risks and potential benefit for the individual patient. The potential to increase the frequency of VT or cause a new VT, an undesirable effect known as “proarrhythmia,” is a potential risk. Many drugs have multiple effects, often blocking more than one channel. **Drug doses, metabolism, and adverse effects are summarized in Chap. 277.**

β -Adrenergic Blockers Many ventricular arrhythmias are sensitive to sympathetic stimulation, and β -adrenergic stimulation also diminishes the electrophysiologic effects of many antiarrhythmic drugs. The safety of β -blocking agents makes them the first choice of therapy for most ventricular arrhythmias. They are particularly useful for exercise-induced arrhythmias and idiopathic arrhythmias, but have limited efficacy for most arrhythmias associated with heart disease. Bradyarrhythmias are the major cardiac toxicity.

Calcium Channel Blockers The nondihydropyridine calcium channel blockers diltiazem and verapamil can be effective for some idiopathic VTs. The risk of proarrhythmia is low, but they have negative inotropic and vasodilatory effects that can aggravate hypotension.

Sodium Channel-Blocking Agents Drugs whose major effect is mediated through sodium channel blockade include mexiletine, quinidine, disopyramide, flecainide, and propafenone, which are available for chronic oral therapy (Table 277-3). Lidocaine, quinidine, and procainamide are available as intravenous formulations. Quinidine, disopyramide, and procainamide also have potassium channel-blocking effects that prolong the QT interval. These agents have potential proarrhythmic effects and, with the possible exception of quinidine, also have negative inotropic effects that may contribute to increased mortality observed in patients with prior MI. Long-term therapy is generally avoided in patients with structural heart disease but may be used to reduce symptomatic arrhythmias in patients with ICDs.

Potassium Channel-Blocking Agents Sotalol and dofetilide block the delayed rectifier potassium channel I_{Kr} , thereby prolonging the QT interval. Sotalol also has nonselective β -adrenergic blocking activity. It has a modest effect on reducing ICD shocks due to ventricular and atrial arrhythmias. Proarrhythmia with Torsade de Pointes due to QT prolongation occurs in 3–5% of patients. Both sotalol and dofetilide are excreted via the kidneys, necessitating dose adjustment or avoidance in renal insufficiency. These drugs must be avoided in patients with other risk factors for Torsade de Pointes, including QT prolongation, hypokalemia, and significant bradycardia.

Amiodarone and Dronedaron Amiodarone, which blocks multiple cardiac ionic currents and has sympatholytic activity, suppresses a variety of ventricular arrhythmias. It is administered intravenously for life-threatening arrhythmias. During chronic oral therapy, electrophysiologic effects develop over several days. It is more effective than sotalol in reducing ICD shocks and is the preferred drug for ventricular arrhythmias in patients with heart

disease who are not candidates for an ICD. Bradyarrhythmias are the major cardiac adverse effect. Ventricular proarrhythmia can occur, but Torsade de Pointes VT is rare. Noncardiac toxicities are a major problem and contribute to drug discontinuation in approximately a third of patients during long-term therapy. Pneumonitis or pulmonary fibrosis occurs in approximately 1% of patients. Photosensitivity is common, and neuropathy and ocular toxicity can occur. Systematic monitoring is recommended during chronic therapy, including assessment for thyroid and liver toxicity every 6 months and lung toxicity with a chest radiograph and/or determination of lung diffusing capacity annually. Dronedarone has structural similarities to amiodarone but without the iodine moiety. Efficacy for ventricular arrhythmias is poor, and it increases mortality in patients with heart failure.

IMPLANTABLE CARディオVERTER-DEFIBRILLATORS

ICDs are highly effective for termination of VT and VF and also provide bradycardia pacing. ICDs decrease mortality in patients at risk for sudden death due to structural heart diseases. In all cases, ICDs are recommended only if there is also expectation for survival of at least a year with acceptable functional capacity. The exception is in cases of patients with end-stage heart disease who are awaiting cardiac transplantation outside the hospital, or who have left bundle branch block QRS prolongation such that they are likely to have improvement in ventricular function with cardiac resynchronization therapy from a biventricular ICD.

ICDs can often terminate monomorphic VT by a burst of rapid pacing faster than the VT, known as antitachycardia pacing (ATP) (Fig. 277-9A). If ATP fails or is not a programmed treatment, as is often the case for rapid VT or VF, a shock is delivered (Fig. 277-9B). Shocks are painful if the patient is conscious. The most common ICD complication is the delivery of unnecessary therapy (either ATP or shocks) in response to a rapid supraventricular tachycardia or

electrical noise as a result of an ICD lead fracture. Interrogation of the ICD, which can be performed remotely and communicated via Internet, is critical after an ICD shock to determine the arrhythmia diagnosis and exclude an unnecessary therapy. Device infection occurs in approximately 1% of patients.

Despite prompt termination of VT or VF by an ICD, the occurrence of these arrhythmias predicts subsequent increased mortality and risk of heart failure. Occurrence of VT or VF should therefore prompt assessment for potential causes including worsening heart failure, electrolyte abnormalities, and ischemia. Repeated shocks, even if appropriate, often induce posttraumatic stress disorder. Antiarrhythmic drugs mostly in the form of amiodarone or catheter ablation are often required for suppression of recurrent arrhythmias. Antiarrhythmic drug therapy can alter the VT rate and the energy required for defibrillation, thereby necessitating programming changes in the ICD's algorithms for detection and therapy.

CATHETER ABLATION FOR VT

Catheter ablation is performed by guiding an electrode catheter to the arrhythmia origin and producing a thermal injury with radiofrequency current. The size and location of the arrhythmia substrate determine the ease and likely effectiveness of the procedure, as well as potential complications. The most common complications, which occur in <5% of patients, are related to vascular access, including bleeding, femoral hematomas, arteriovenous fistulae, and pseudoaneurysms.

Catheter ablation is a reasonable first-line therapy for many patients with symptomatic idiopathic VTs. Success rates for those originating from a focus in the right ventricular outflow tract are in the range of 80–90% but lower for idiopathic VTs arising in less common locations such as from the LV outflow tract or aortic root, along the atrioventricular valve annuli, and from the papillary muscles. Failure of ablation is often due to inability to induce the

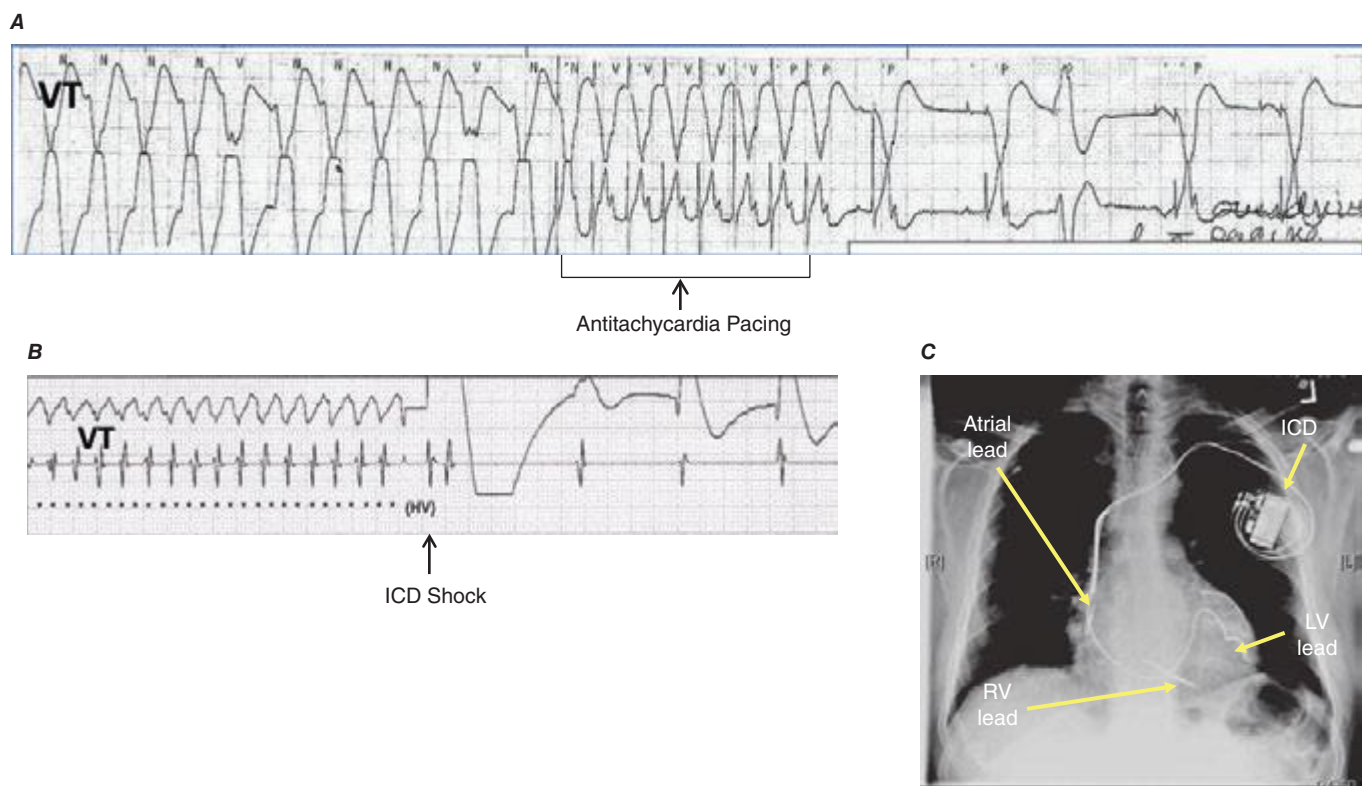


FIGURE 277-9 Implantable cardioverter-defibrillator (ICD) and therapies for ventricular arrhythmias. **A.** A monomorphic ventricular tachycardia (VT) is terminated by a burst of pacing impulses at a rate faster than VT (antitachycardia pacing). **B.** A rapid VT is converted with a high-voltage shock (arrow). The chest x-ray in panel **C** shows the components of an ICD capable of biventricular pacing. ICD generator in the subcutaneous tissue of the left upper chest, pacing leads in the right atrium and the left ventricular (LV) branch of the coronary sinus (LV lead), and a pacing/defibrillating lead in the right ventricle (RV lead) are shown.

1500 arrhythmia for precise localization or because the origin of the VT is from a site that is inaccessible or in close proximity to a coronary artery. Complications are infrequent but can include perforation with cardiac tamponade, atrioventricular block due to injury to the conduction system, and coronary artery injury for foci in proximity to a coronary vessel.

In patients with scar-related VT due to prior infarction or cardiomyopathy, ablation targets abnormal regions in the scar. Because these scars often contain multiple reentry circuits over relatively large regions, extensive areas of ablation are required, and these areas are often identified as regions of low voltage displayed on anatomic reconstructions of the ventricle (Fig. 277-5). If the circuits are not confined to the subendocardial scar, epicardial mapping and ablation can be performed via a subxiphoid pericardial puncture, similar to a pericardiocentesis. Epicardial mapping and ablation are often required for VTs due to nonischemic cardiomyopathy, but also have potentially greater risks of bleeding, coronary injury, and post-procedure pericarditis, which is usually transient. For drug-refractory VT due to prior MI, ablation abolishes VT in approximately half of patients and reduces the frequency of VT in an additional 20%. More than one procedure is necessary in up to 30% of patients. Ablation can be lifesaving for patients with very frequent or incessant VT. Procedure-related mortality is in the range of 3%, with most mortality due to continued uncontrollable VT when the procedure fails. In nonischemic heart disease, the arrhythmia substrate locations are more variable and outcomes are less well defined.

Catheter ablation can also be lifesaving for rare patients with recurrent polymorphic VT and VF that is repeatedly initiated by a uniform PVC. The initiating ectopic beat often originates from the Purkinje system or the right ventricular outflow tract and can be targeted for ablation.

When antiarrhythmic drug therapy and catheter ablation fail or are not options, surgical cryoablation, often combined with aneurysmectomy, can be effective therapy for recurrent VT due to prior MI and has also been used successfully in a few patients with nonischemic heart disease. Few centers now maintain the expertise for this therapy. Injection of absolute ethanol into the coronary arterial blood supply of the arrhythmia substrate has also been used for ablation in a small number of patients who have failed catheter ablation and drugs.

SUMMARY

Patients with ventricular arrhythmias fall into three general groups. The first are those who have associated structural heart disease that must be detected. The risk of life-threatening arrhythmias causing sudden death is indicated by the nature of the arrhythmia—sustained (or causing cardiac arrest) or nonsustained, in which case the risk of life-threatening arrhythmias is assessed from the severity of the heart disease, usually the severity of ventricular dysfunction. ICDs provide the most protection from sudden arrhythmic death. The second group comprises those who do not have recognizable structural heart disease, but have a genetic syndrome associated with increased risk of sudden death. A family history of sudden death and abnormal electrocardiogram most frequently suggest the diagnosis. The third group includes individuals with benign idiopathic arrhythmias who may require therapy to control symptoms, but who are not at significant risk for life-threatening arrhythmias. The appropriate recognition of these patients is facilitated by thoughtful application of ECG and cardiac imaging. High-risk individuals benefit from specialized care for consideration of ICDs, catheter ablation, and antiarrhythmic drug therapy.

The electrocardiograms in this atlas supplement those illustrated in [Chaps. 274 and 276](#). The interpretations emphasize findings of specific teaching value.

All of the figures are adapted from cases in ECG Wave-Maven, Copyright 2003, Beth Israel Deaconess Medical Center, <http://ecg.bidmc.harvard.edu>.

The abbreviations used in this chapter are as follows:

- AF—atrial fibrillation
- AV—atrioventricular
- LBBB—left bundle branch block
- LV—left ventricular
- LVH—left ventricular hypertrophy
- MI—myocardial infarction
- NSR—normal sinus rhythm
- RBBB—right bundle branch block
- VT—ventricular tachycardia
- WPW—Wolff-Parkinson-White

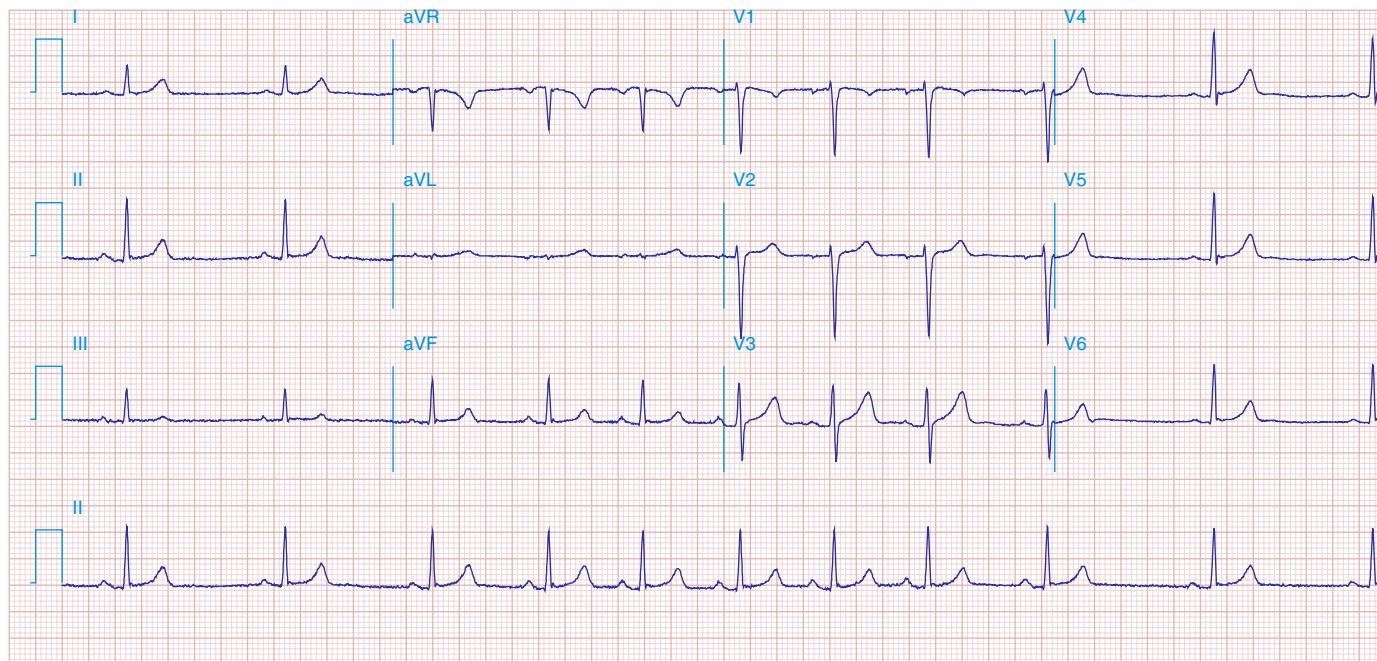


FIGURE 278e-1 Respiratory sinus arrhythmia, a physiologic finding in a healthy young adult. The rate of the sinus pacemaker is relatively slow at the beginning of the strip during expiration, then accelerates during inspiration and slows again with expiration. Changes are due to cardiac vagal tone modulation with breathing.

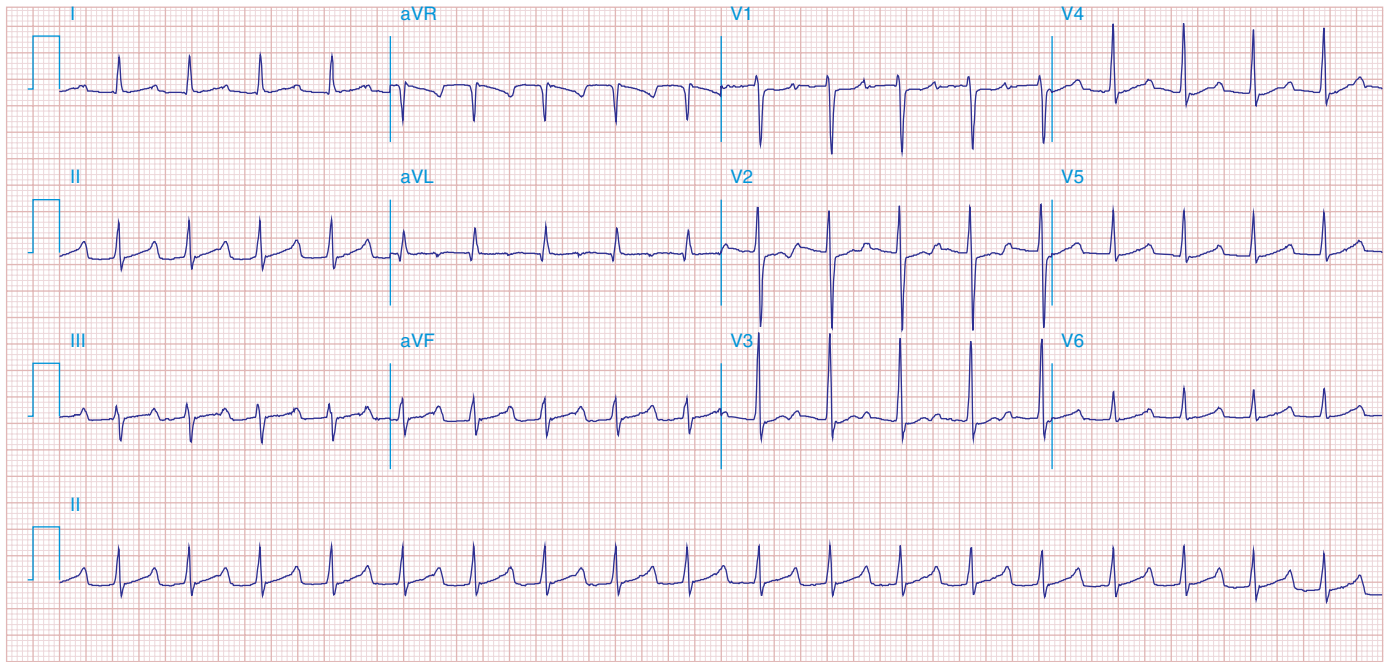


FIGURE 278e-2 Sinus tachycardia (110/min) with first-degree AV “block” (conduction delay) with PR interval = 0.28 s. The P wave is visible after the ST-T wave in V_1 – V_3 and superimposed on the T wave in other leads. Atrial (nonsinus) tachycardias may produce a similar pattern, but the rate is usually faster.

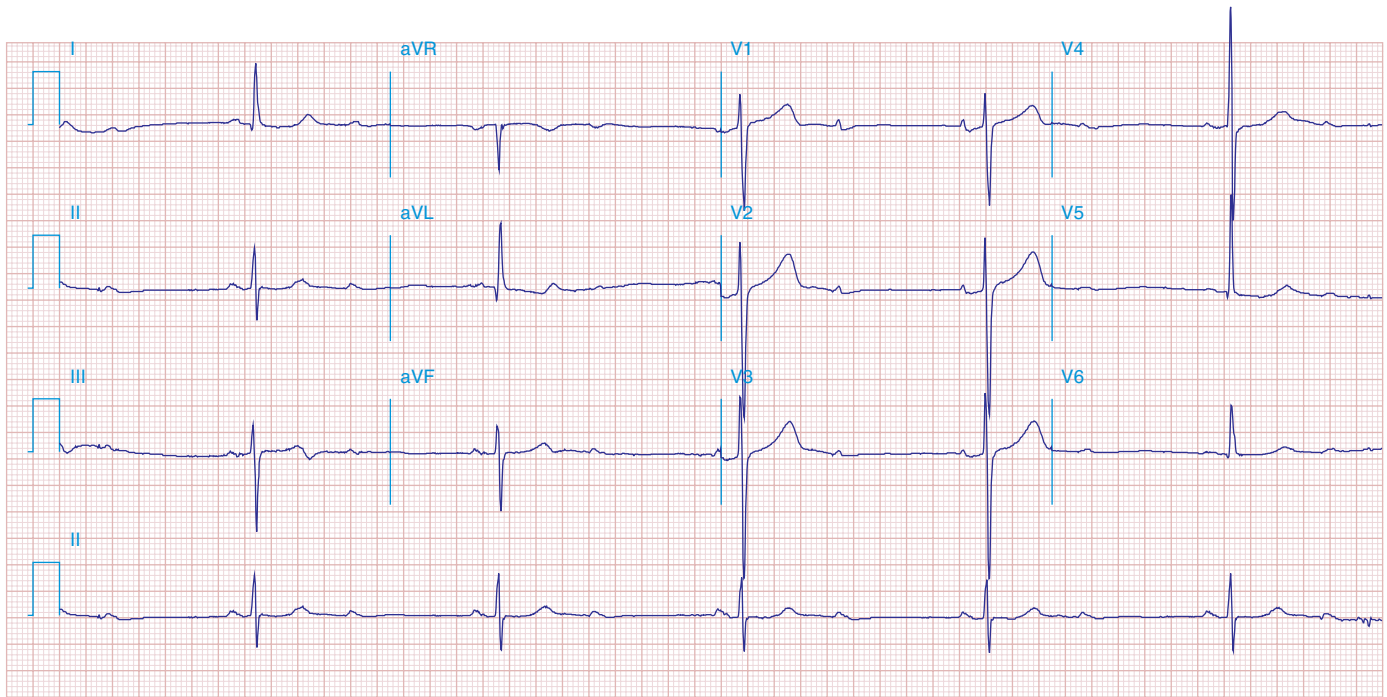


FIGURE 278e-3 Sinus rhythm (P wave rate about 60/min) with 2:1 AV (second-degree) block causing marked bradycardia (ventricular rate of about 30/min). LVH is also present, along with left atrial abnormality.

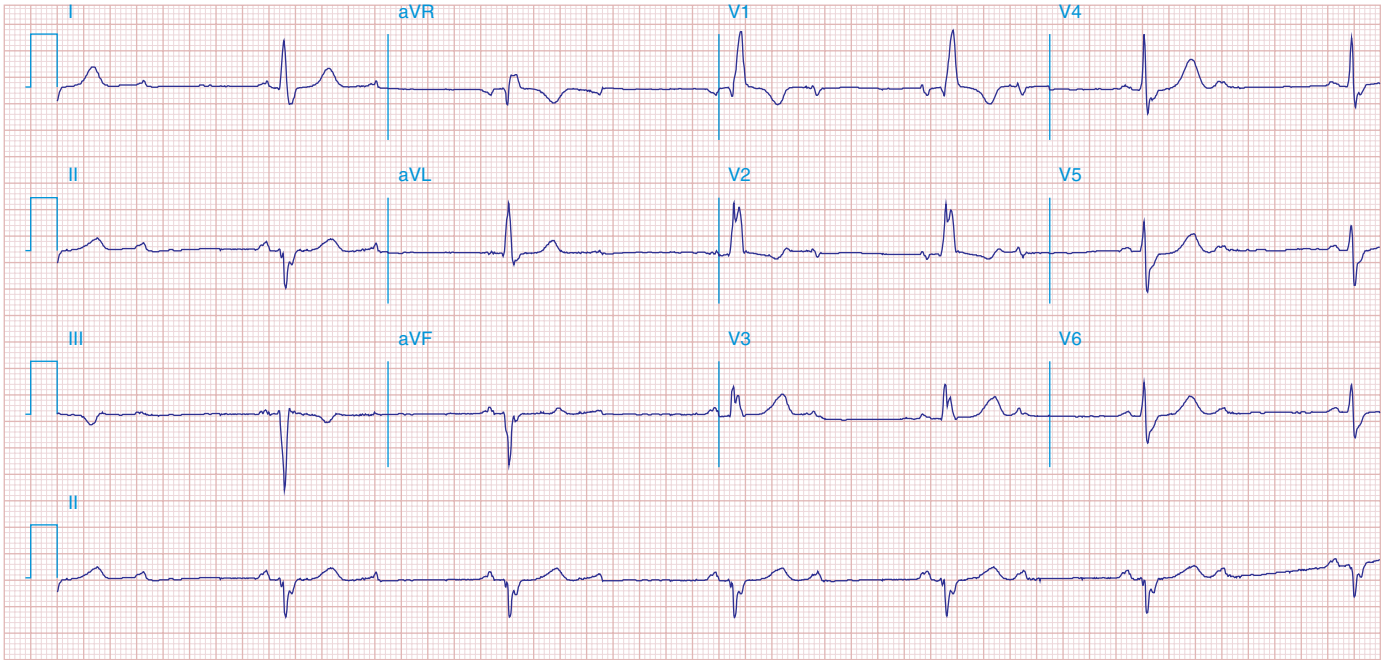


FIGURE 278e-4 Sinus rhythm (P wave rate about 60/min) with 2:1 (second-degree) AV block yielding a ventricular (pulse) rate of about 30/min. Left atrial abnormality. RBBB with left anterior fascicular block. Possible inferior MI.

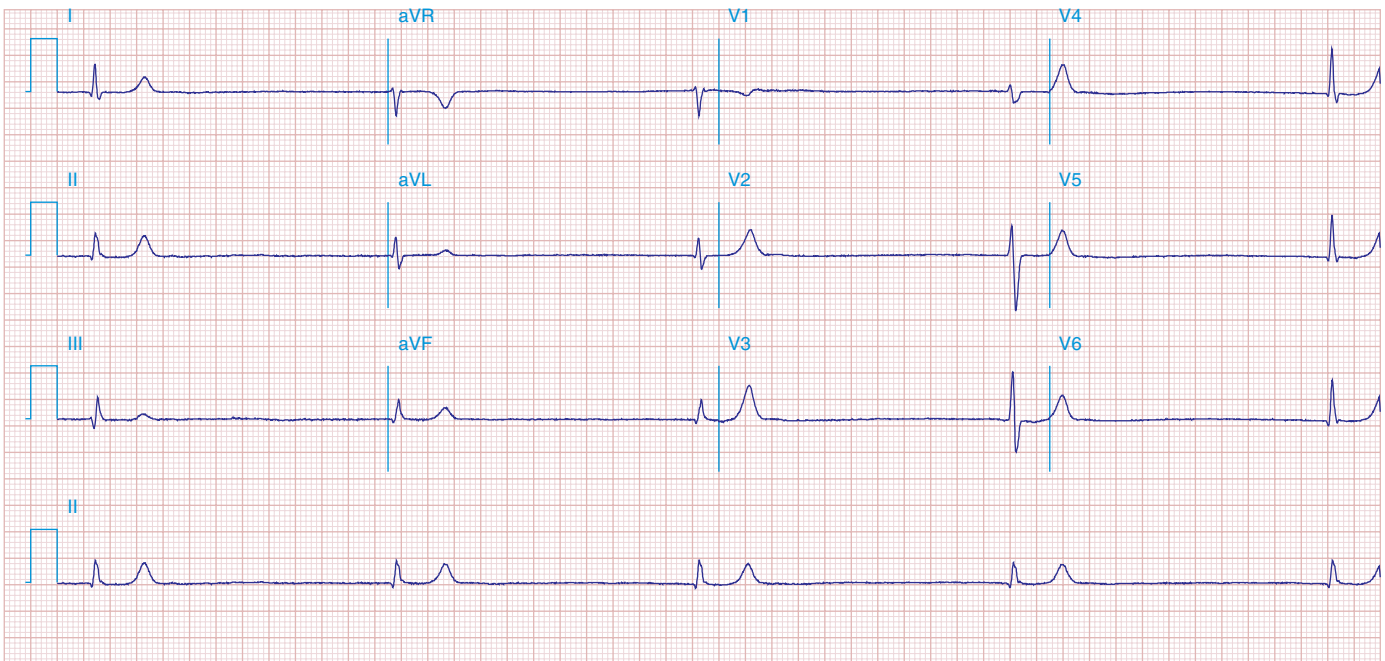


FIGURE 278e-5 Marked junctional bradycardia (25 beats/min). Rate is regular with a flat baseline between narrow QRS complexes, without evident P waves. Patient was on atenolol, with possible underlying sick sinus syndrome. The serum potassium was slightly elevated at 5.5 mEq/L.

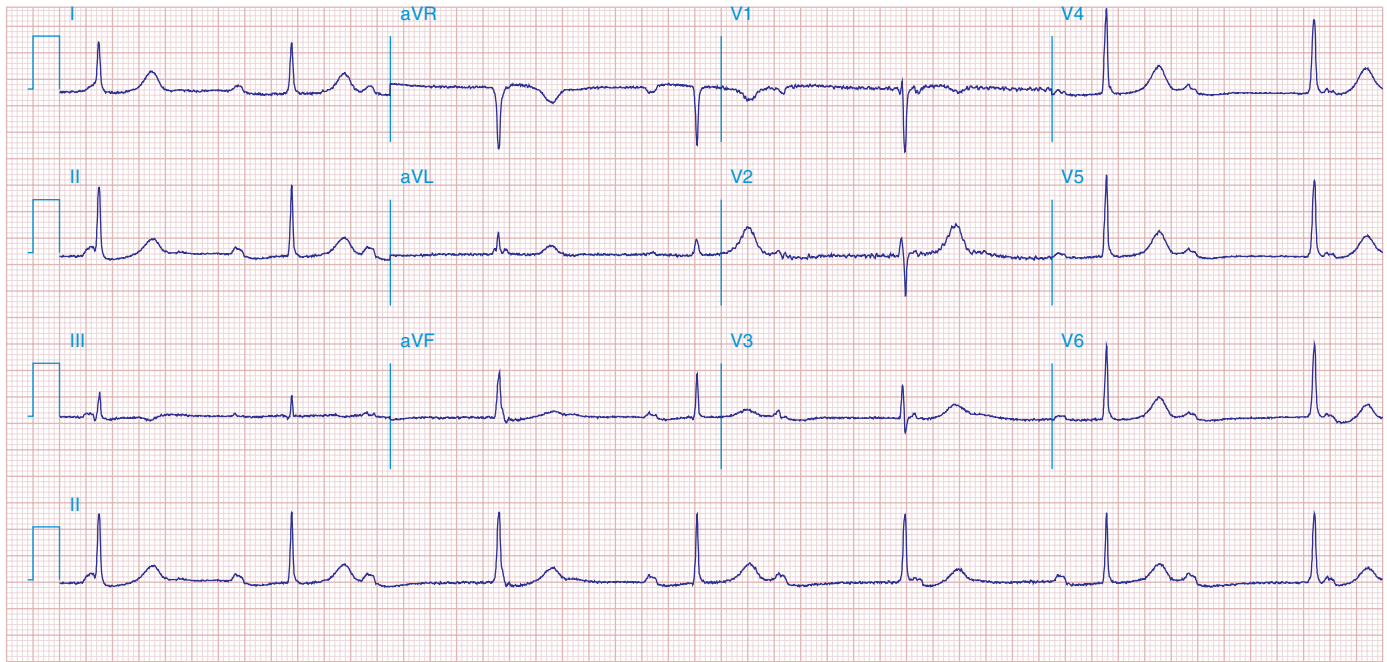


FIGURE 278e-6 Sinus rhythm at a rate of 64/min (P wave rate) with third-degree (complete) AV block yielding an effective heart (pulse) rate of 40/min. The slow, narrow QRS complexes indicate an AV junctional escape pacemaker. Left atrial abnormality.

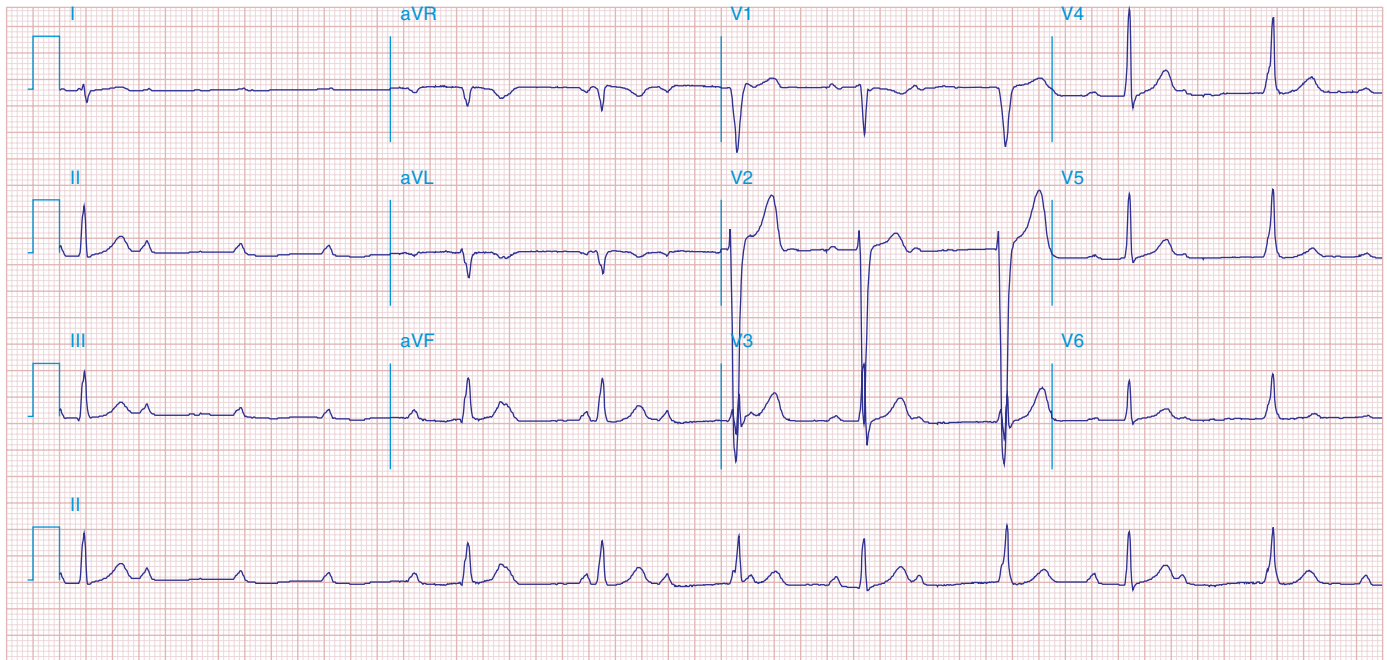


FIGURE 278e-7 Sinus rhythm at a rate of 90/min with advanced second-degree AV block and possible transient complete heart block with Lyme carditis.

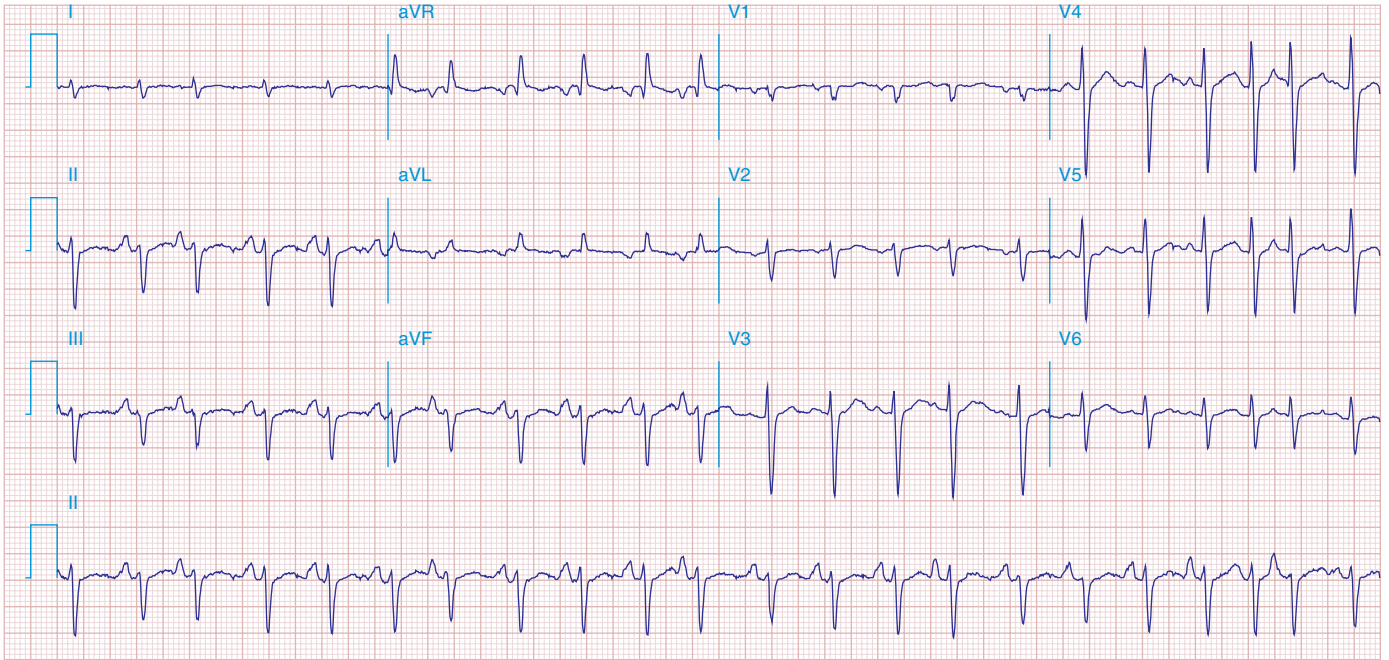


FIGURE 278e-8 Multifocal atrial tachycardia with varying P-wave morphologies and P-P intervals; right atrial overload with peaked P waves in II, III, and aVF (with vertical P-wave axis); superior QRS axis; slow R-wave progression with delayed transition in precordial leads in patient with severe chronic obstructive lung disease.

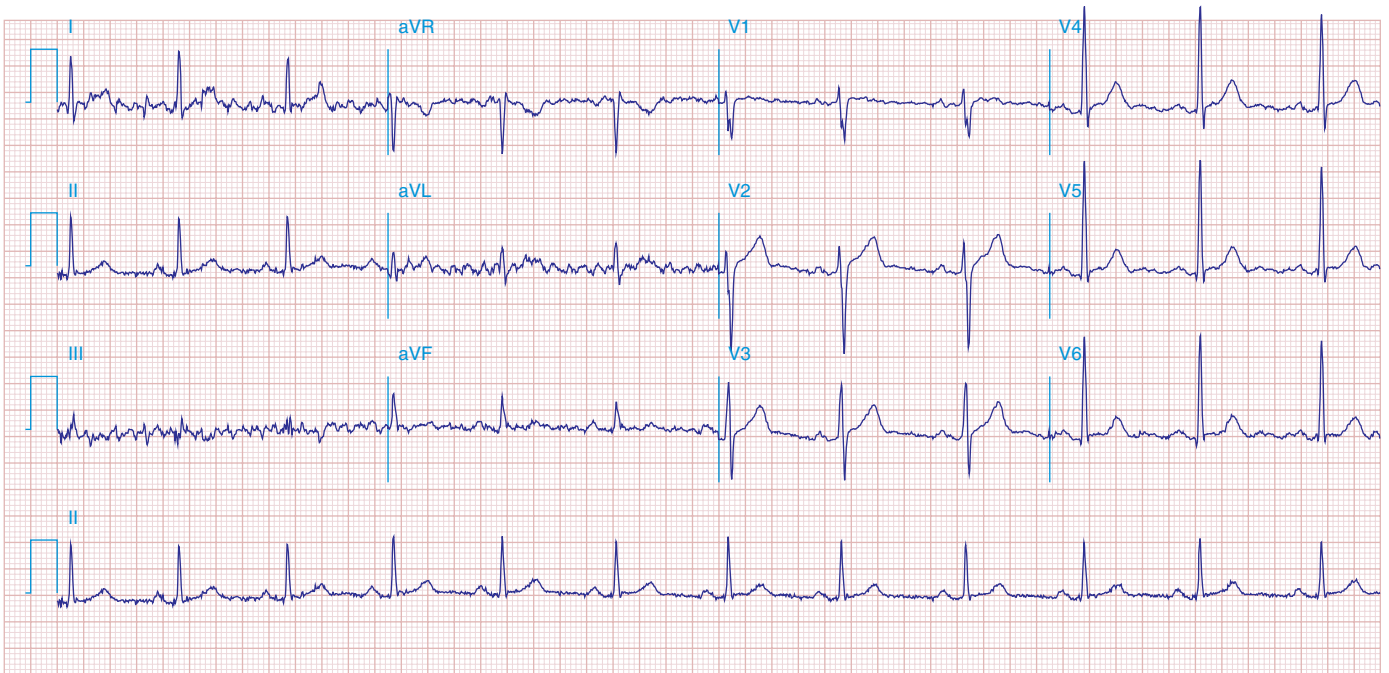


FIGURE 278e-9 NSR in a patient with Parkinson's disease. Tremor artifact, best seen in limb leads. This tremor artifact may sometimes be confused with atrial flutter/fibrillation. Borderline voltage criteria for LVH are present.

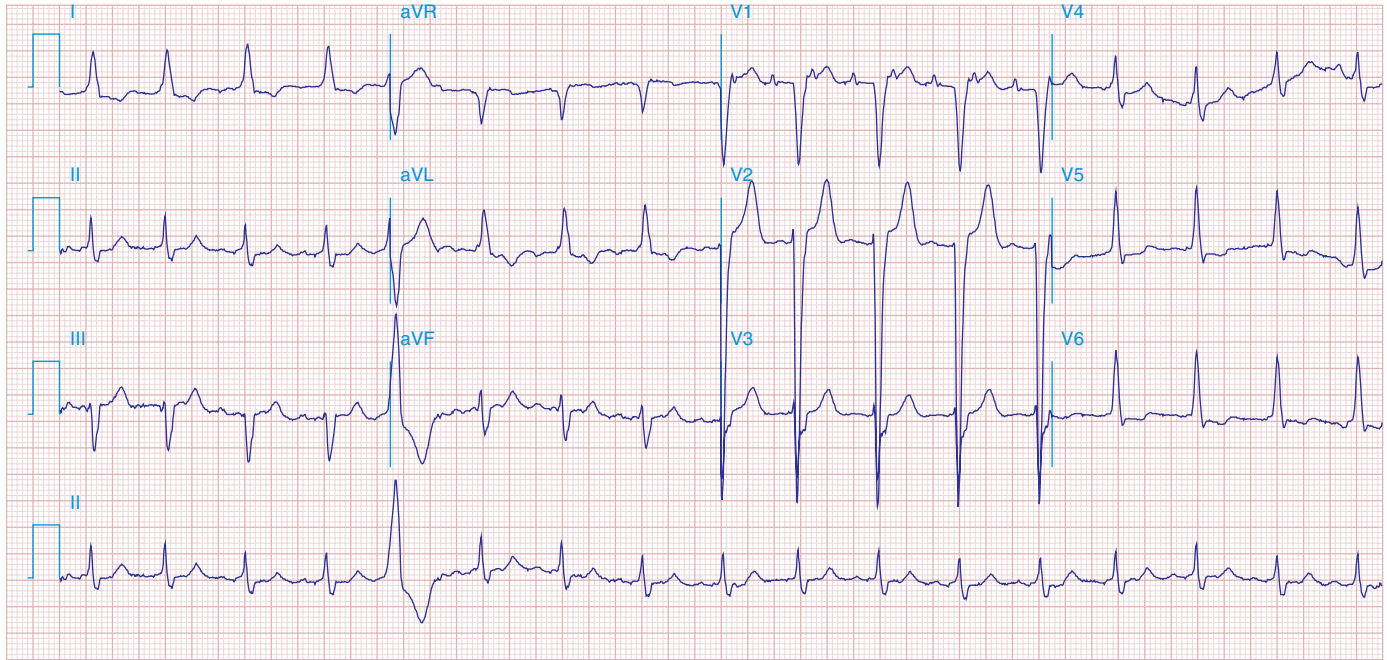


FIGURE 278e-10 Atrial tachycardia with atrial rate of about 200/min (note lead V_1), 2:1 AV block (conduction), and one premature ventricular complex. Also present: LVH with intraventricular conduction delay and slow precordial R-wave progression (cannot rule out prior anterior MI).

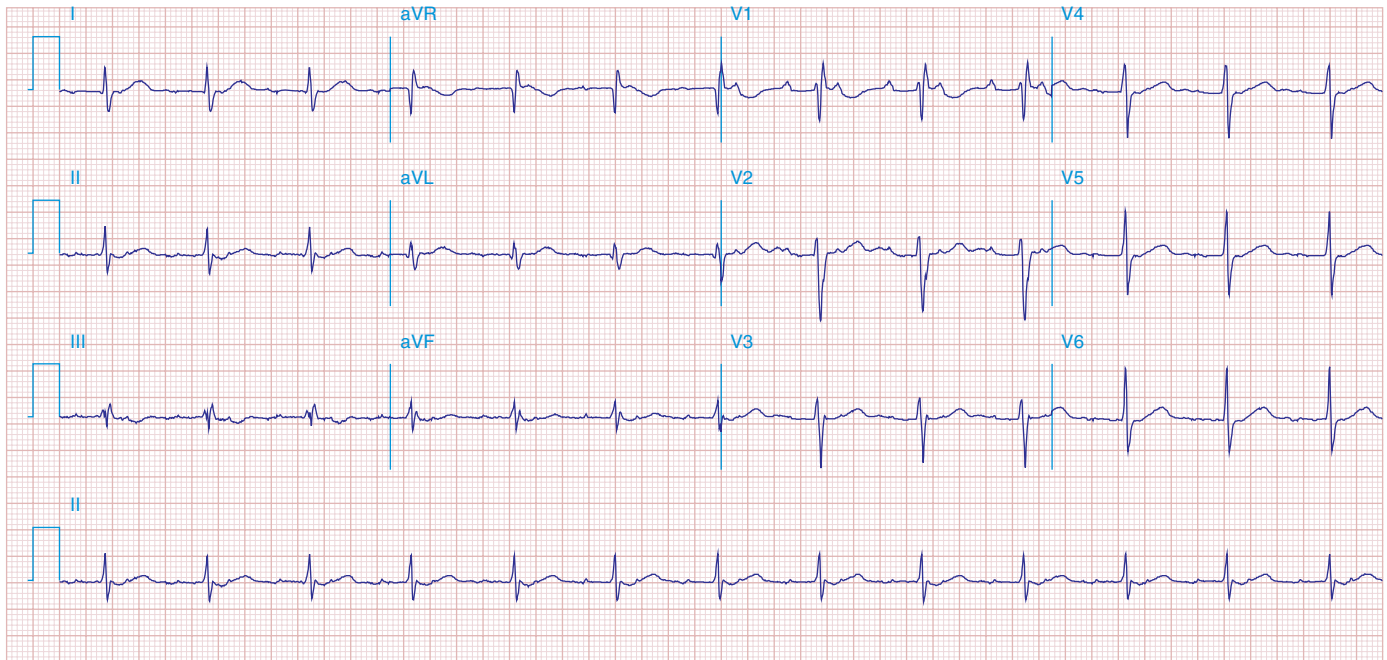


FIGURE 278e-11 Atrial tachycardia with 2:1 block. P-wave rate is about 150/min, with ventricular (QRS) rate of about 75/min. The non-conducted ("extra") P waves just after the QRS complex are best seen in lead V_1 . Also, note incomplete RBBB and borderline QT prolongation.

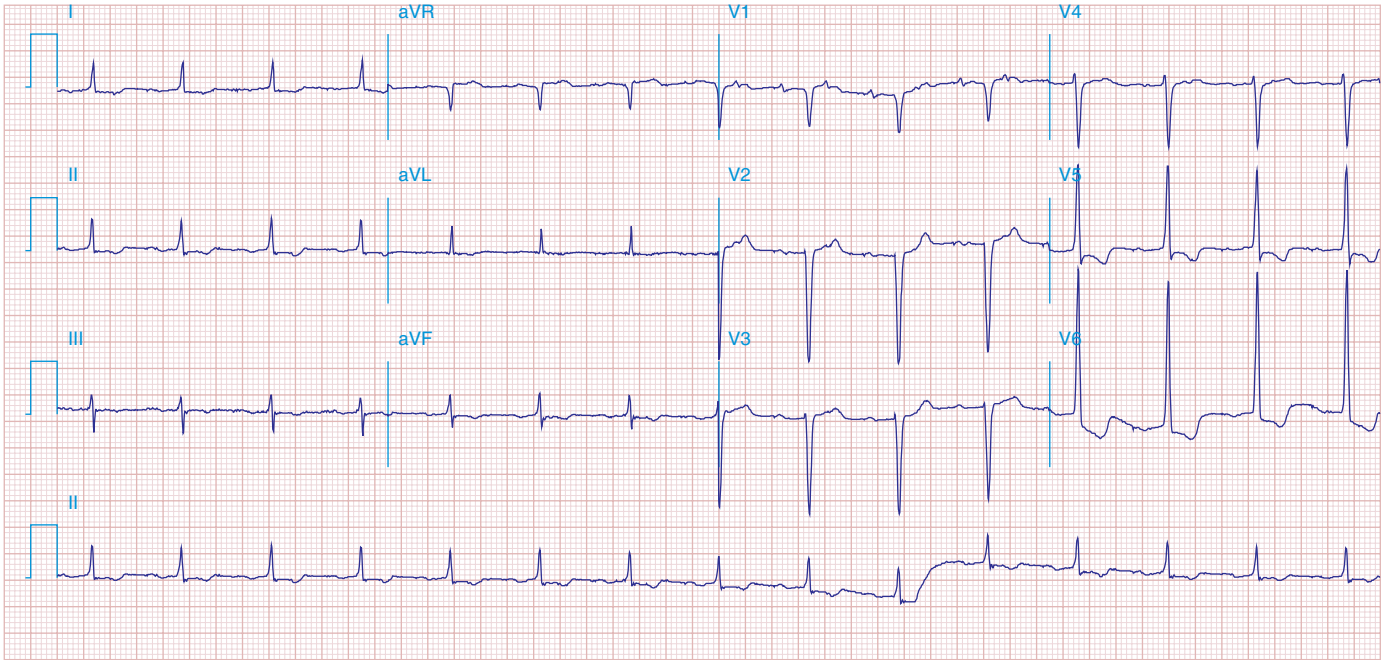


FIGURE 278e-12 Atrial tachycardia (180/min with 2:1 AV block [see lead V₁]). LVH by precordial voltage and nonspecific ST-T changes. Slow R-wave progression (V₁–V₄) raises consideration of prior anterior MI.

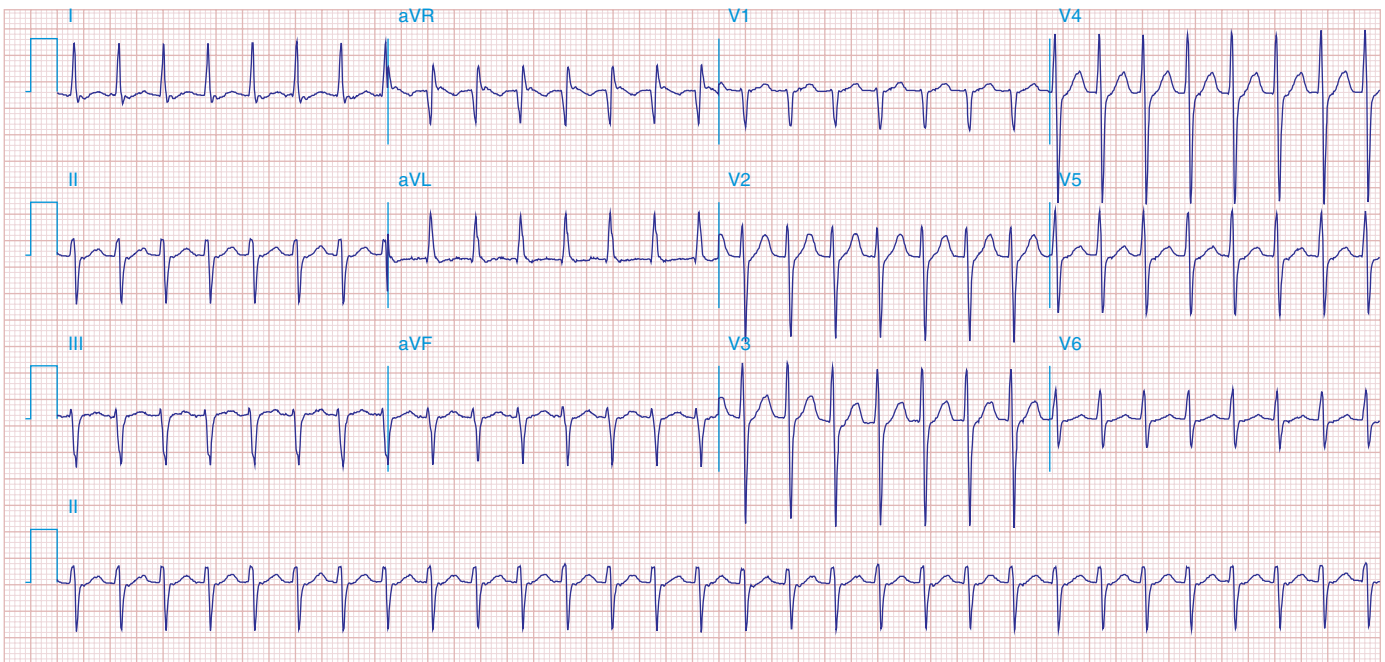


FIGURE 278e-13 AV nodal reentrant tachycardia (AVNRT) at a rate of 150/min. Note subtle “pseudo” R waves in lead aVR due to retrograde atrial activation, which usually occurs nearly simultaneous with ventricles in AVNRT. Left-axis deviation consistent with left anterior fascicular block (hemiblock) is also present.

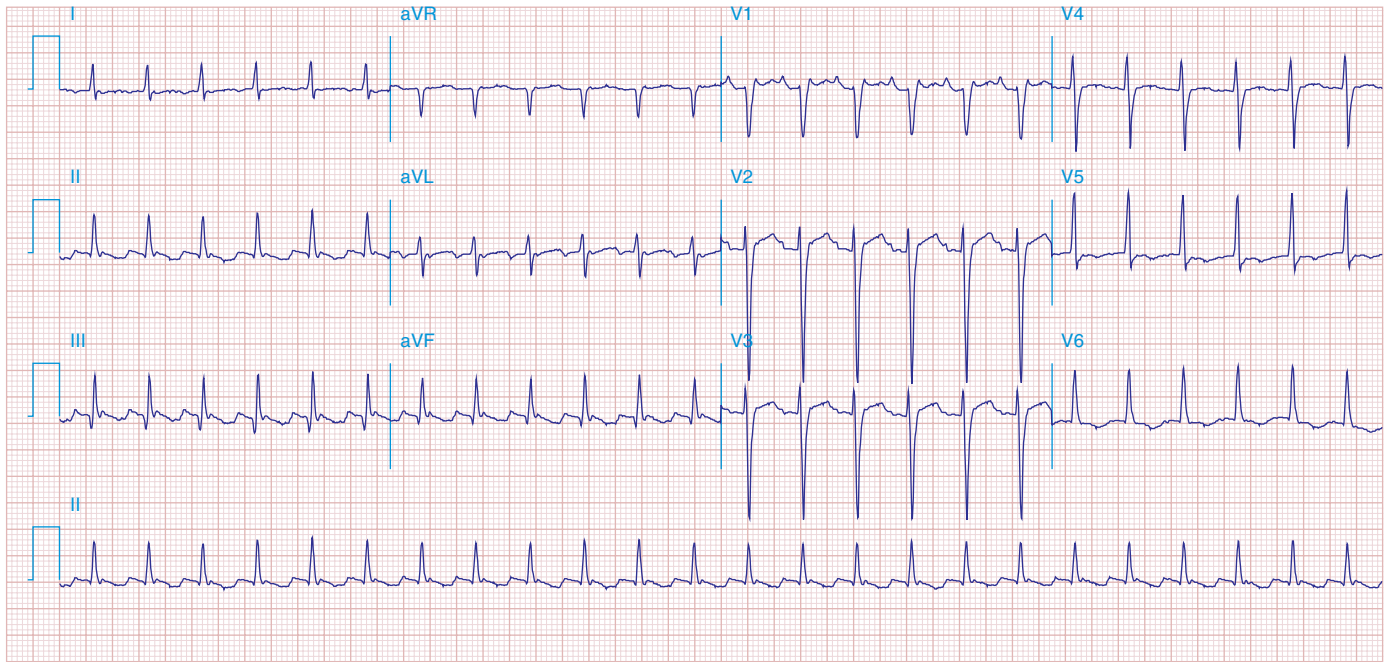


FIGURE 278e-14 Atrial flutter with 2:1 AV conduction. Note typical atrial flutter waves, partly hidden in the early ST segment, seen, for example, in leads II and V_1 .

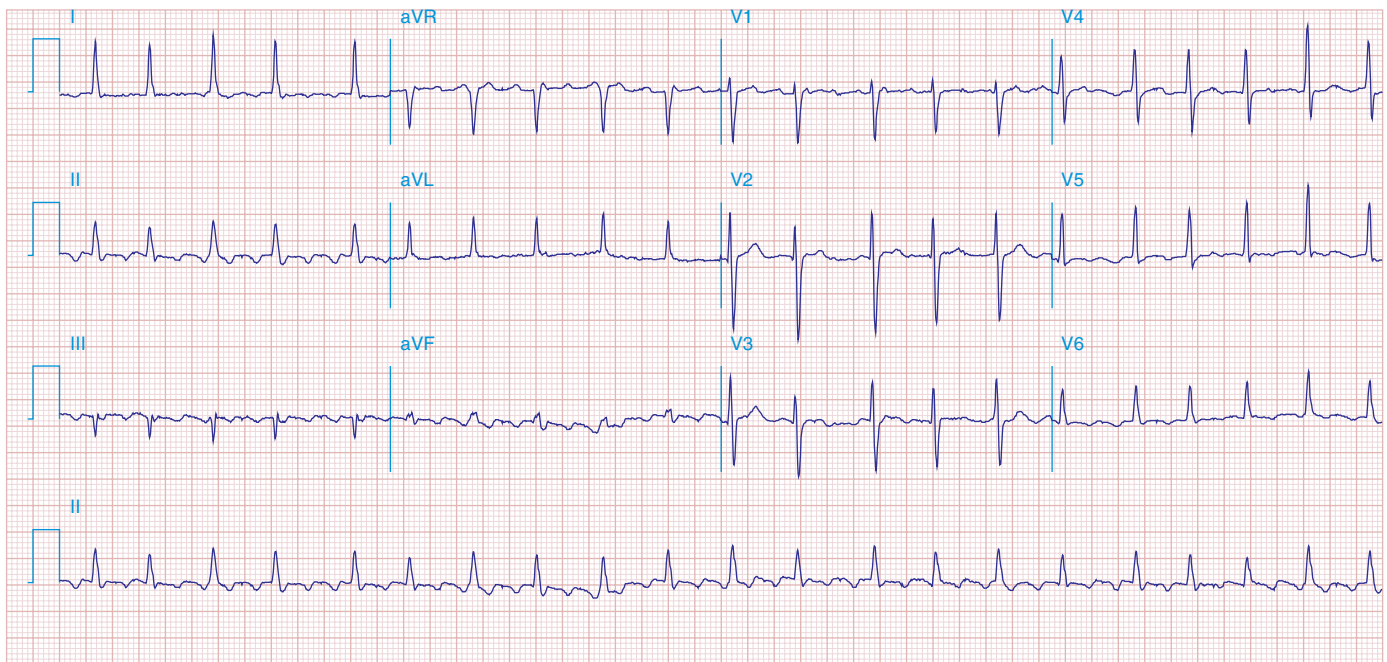


FIGURE 278e-15 Atrial flutter with atrial rate 300/min and variable (predominant 2:1 and 3:1) AV conduction. Typical flutter waves are best seen in lead II.

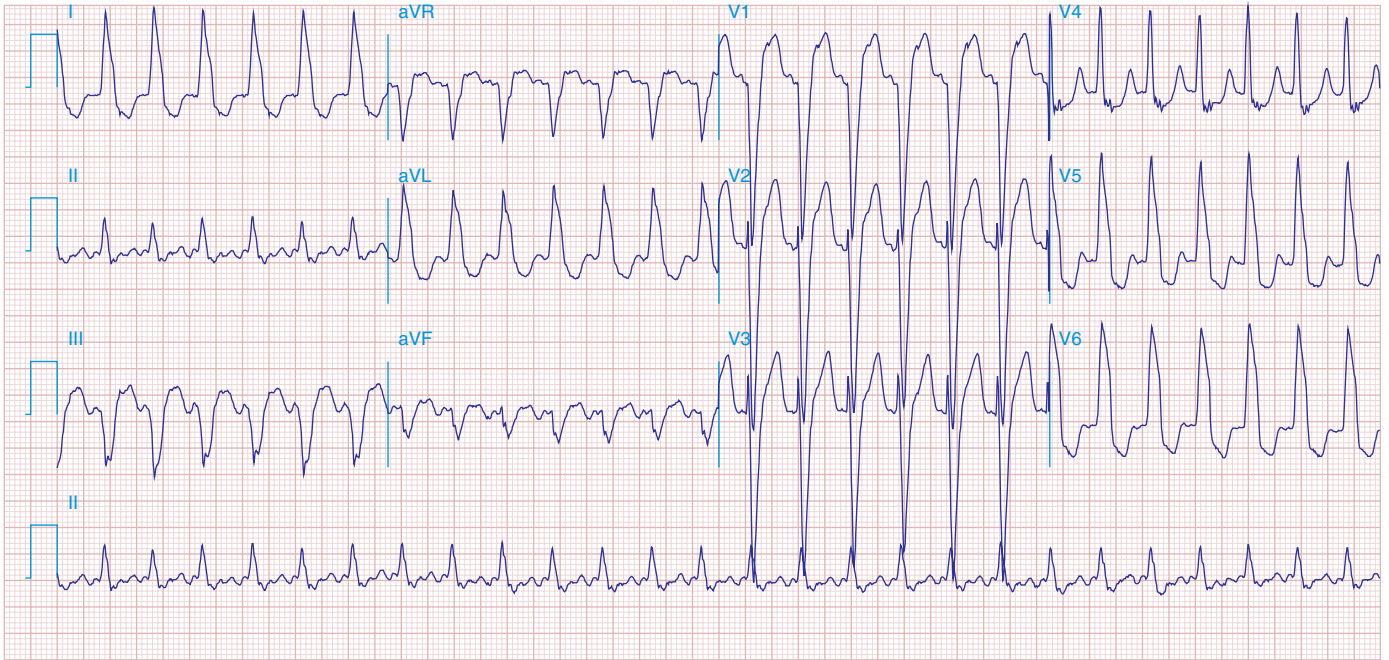


FIGURE 278e-16 Wide complex tachycardia. Atrial flutter with 2:1 AV conduction (block) and LBBB, not to be mistaken for VT. Typical atrial flutter activity is clearly present in lead II, at a cycle rate of about 320/min, yielding an effective ventricular rate of about 160/min.

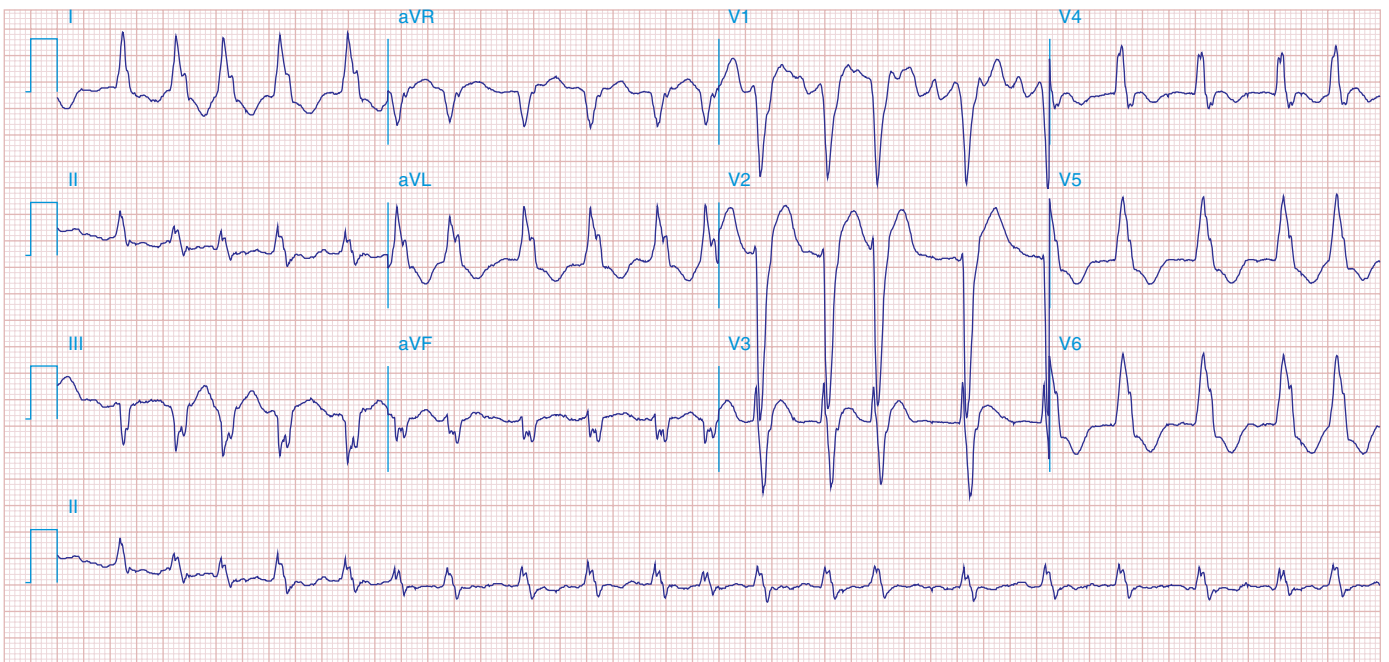


FIGURE 278e-17 AF with LBBB. The ventricular rhythm is erratically irregular. Coarse fibrillatory waves are best seen in lead V₁, with a typical LBBB pattern.

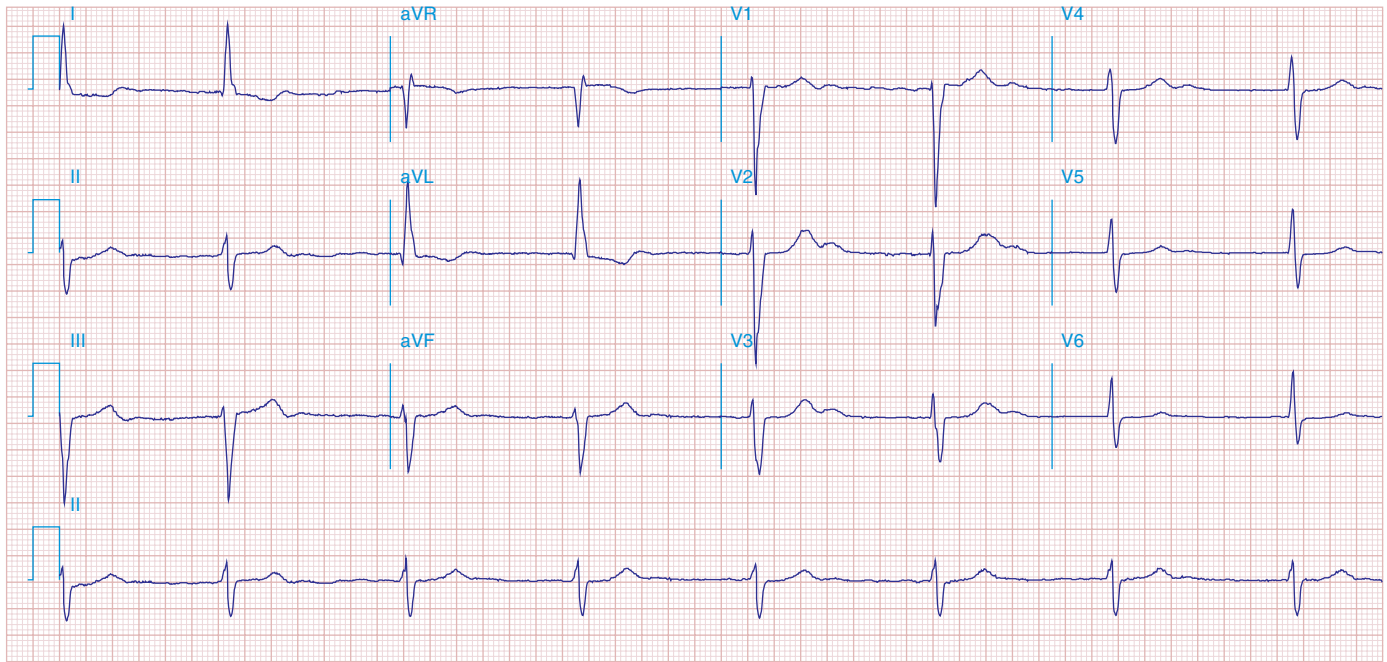


FIGURE 278e-18 AF with complete heart block and a junctional escape mechanism causing a slow regular ventricular response (45/min). The QRS complexes show an intraventricular conduction delay with left-axis deviation and LVH. Q-T (U) prolongation is also present.

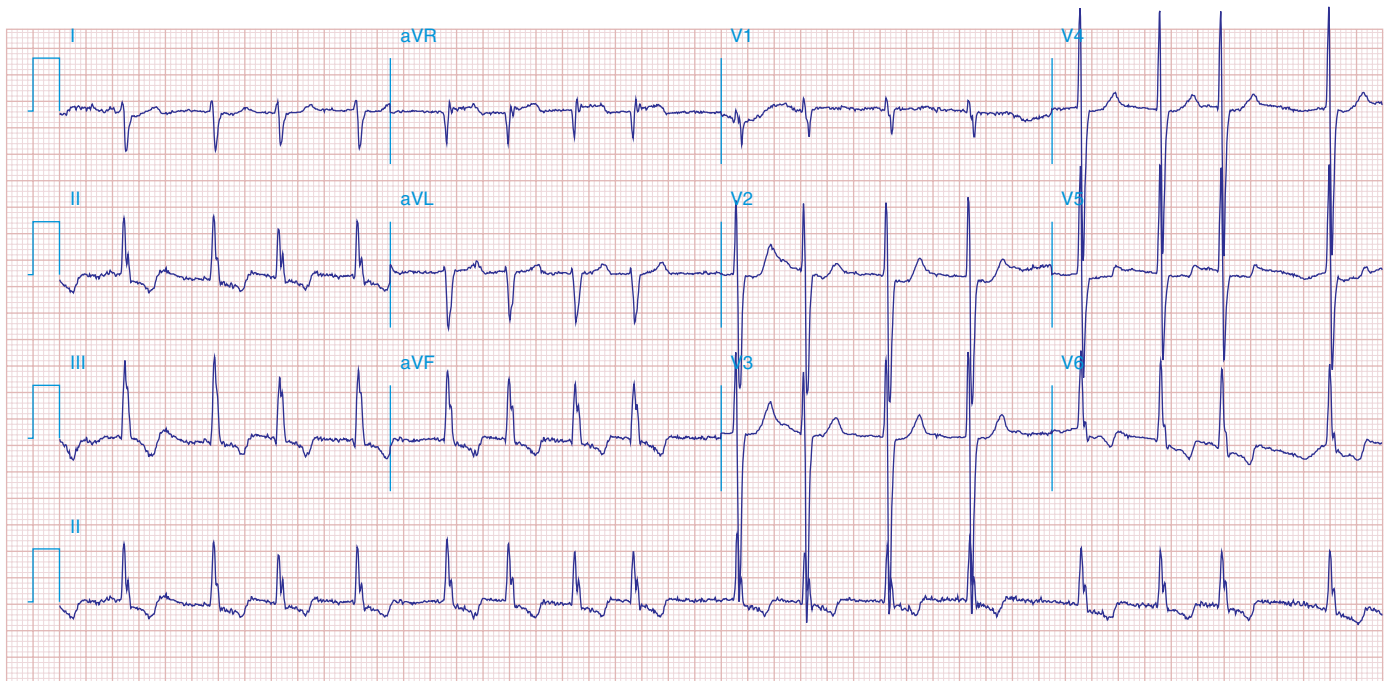


FIGURE 278e-19 AF with right-axis deviation and LVH. Tracing suggests biventricular hypertrophy in a patient with mitral stenosis and aortic valve disease.

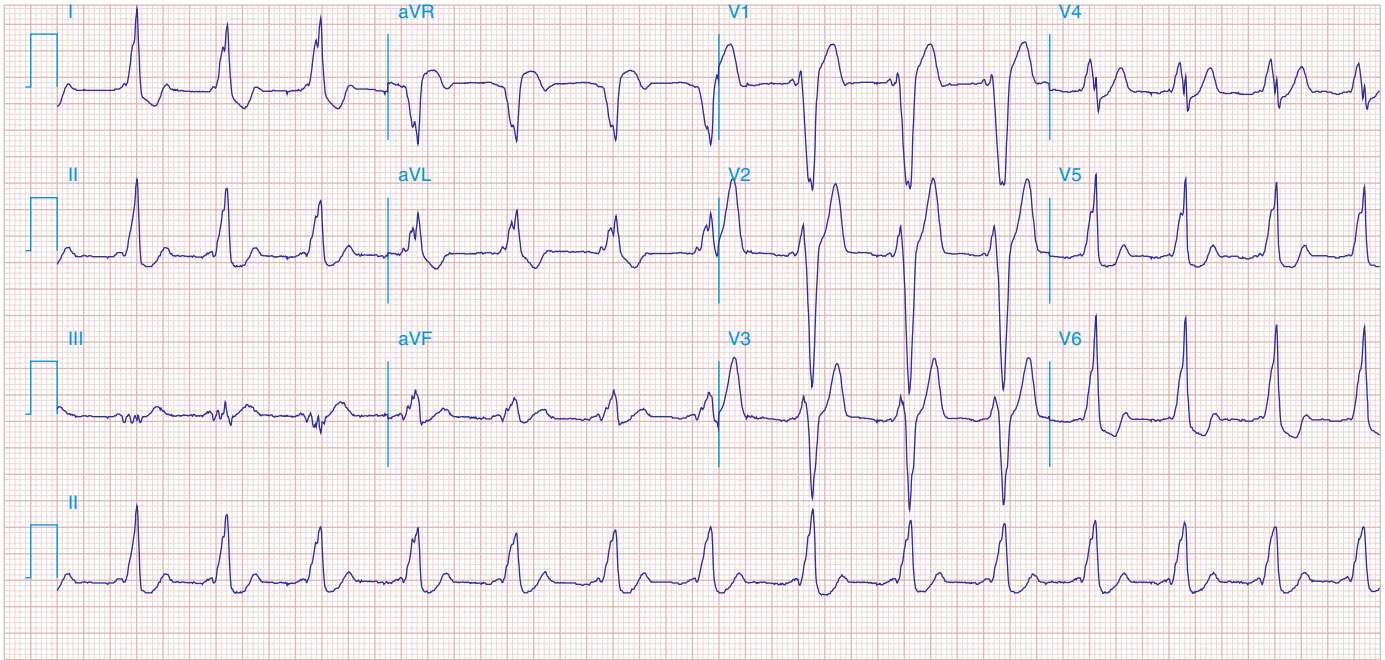


FIGURE 278e-20 WPW preexcitation pattern, with triad of short PR, wide QRS, and delta waves. Polarity of the delta waves (slightly positive in leads V_1 and V_2 and most positive in lead II and lateral chest leads) is consistent with a right-sided bypass tract.

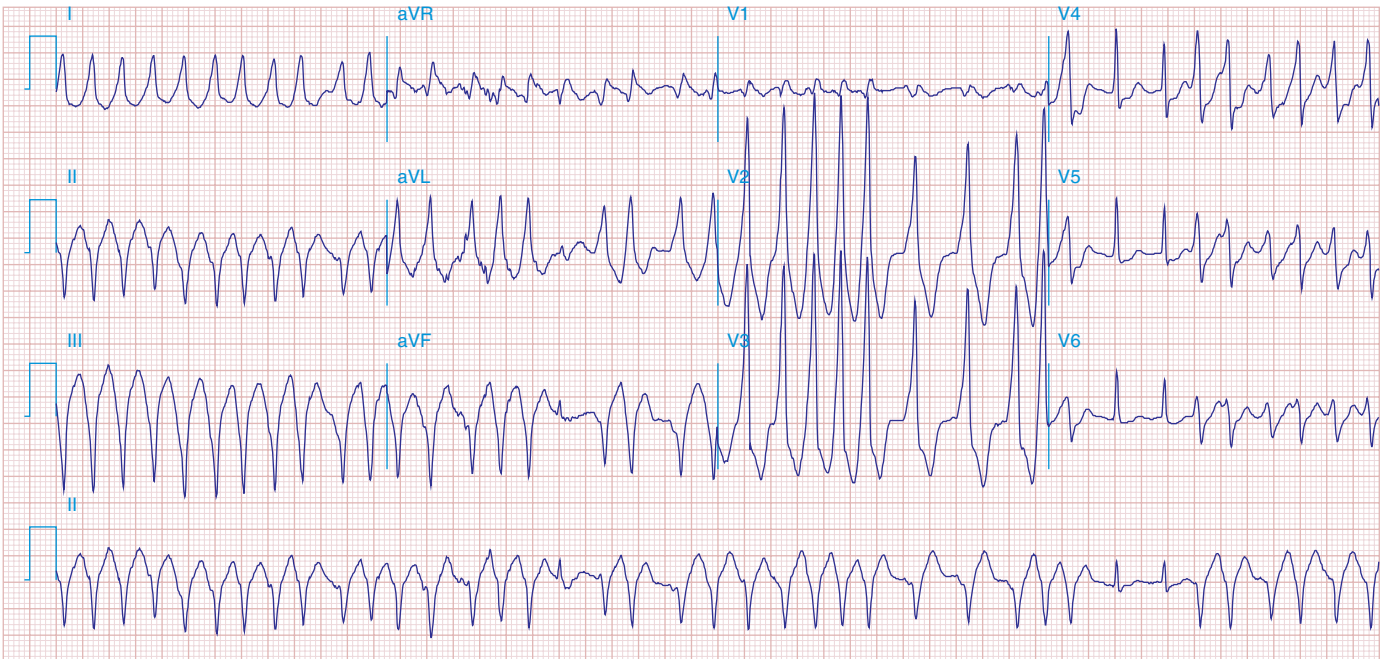


FIGURE 278e-21 AF in patient with the WPW syndrome and antegrade conduction down the bypass tract leading to a wide complex tachycardia. Rhythm is "irregularly irregular," and rate is extremely rapid (about 230/min). Not all beats are preexcited.

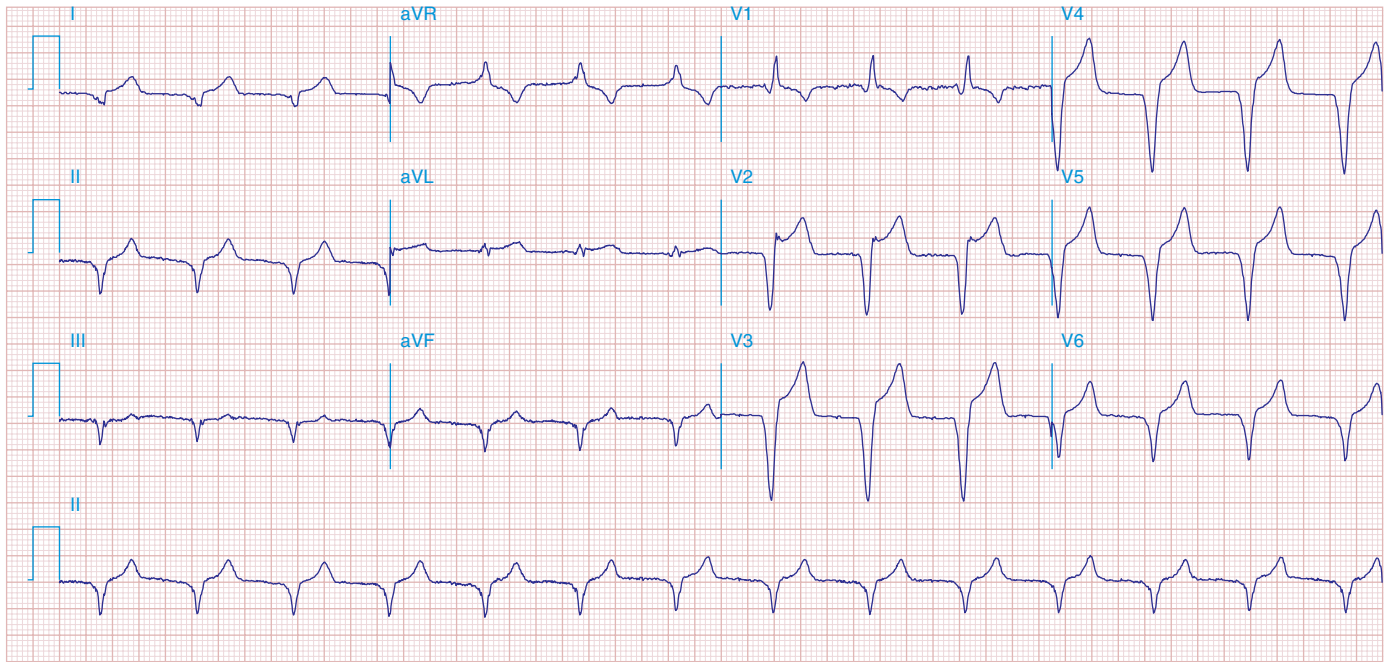


FIGURE 278e-22 Accelerated idioventricular rhythm (AIVR) originating from the LV and accounting for RBBB morphology. ST elevations in the precordial leads from underlying acute MI.

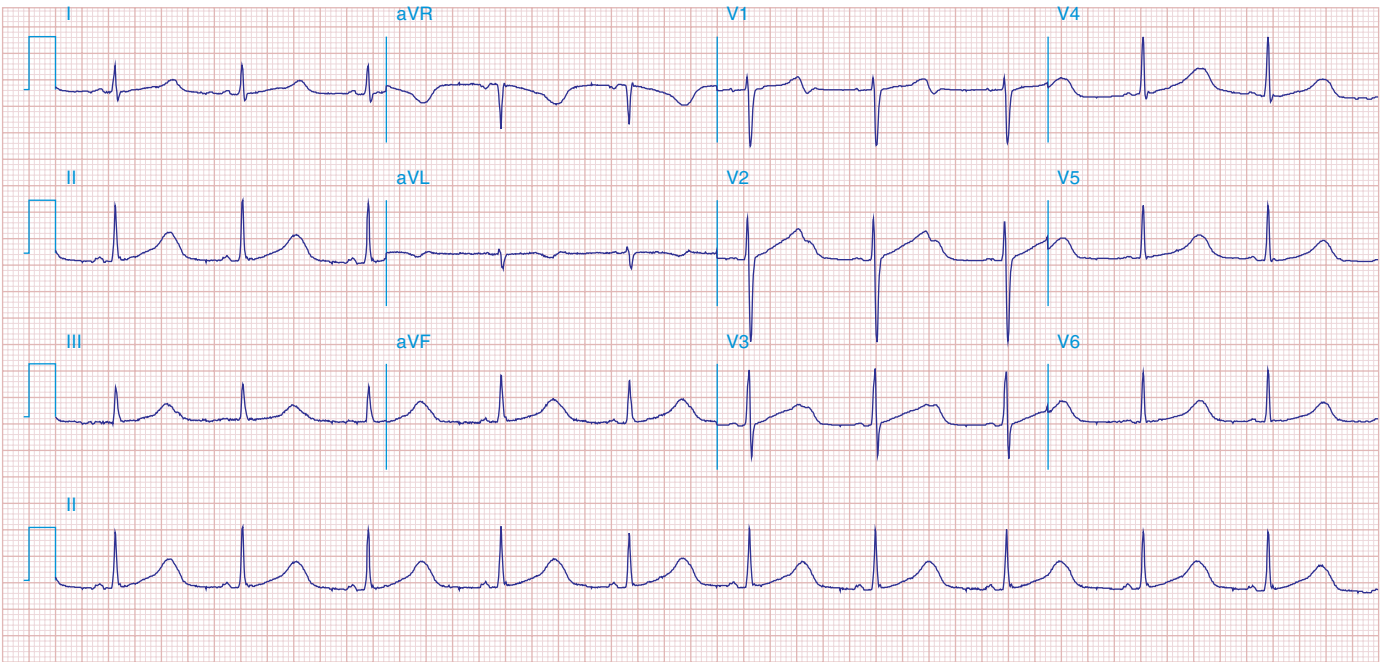


FIGURE 278e-23 Prolonged (0.60 s) QT interval in a patient with a hereditary long QT syndrome.

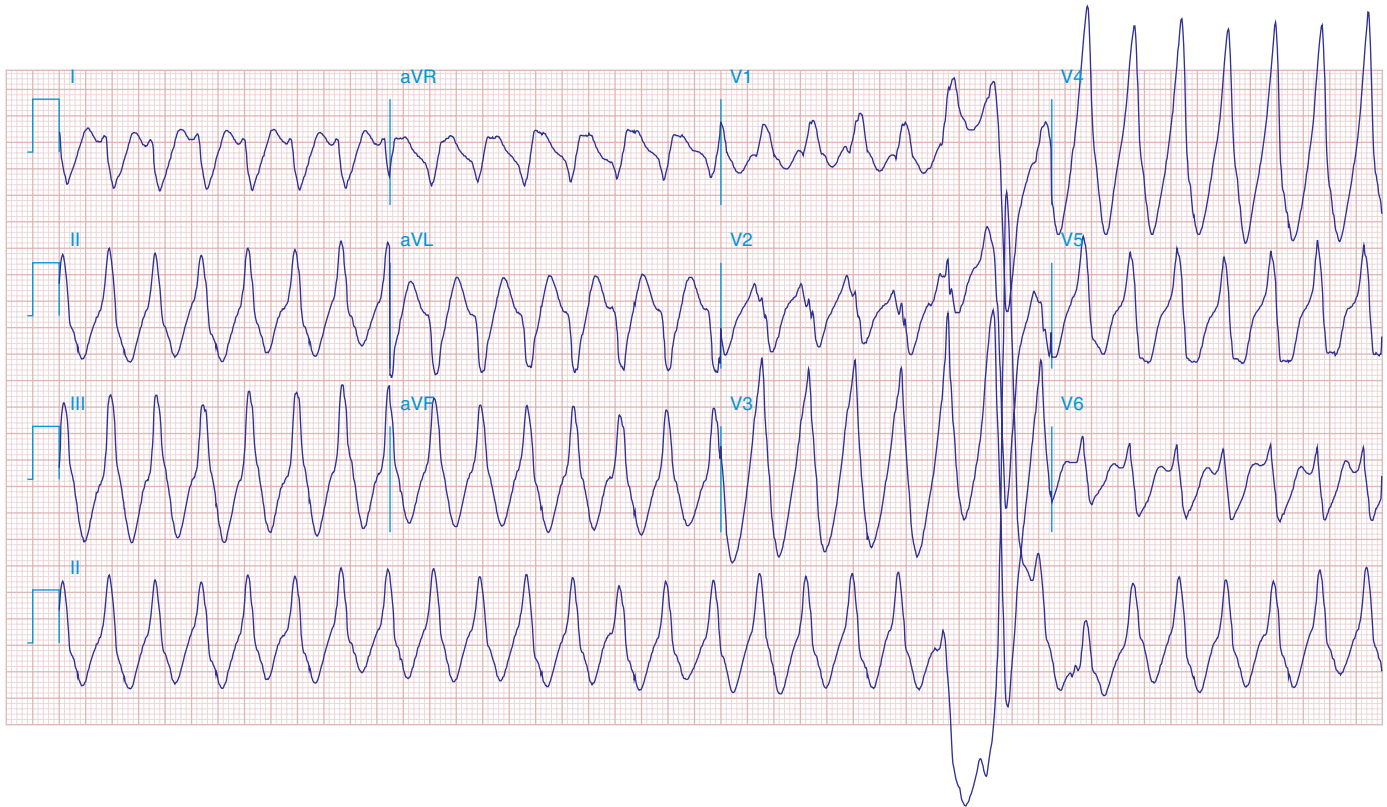


FIGURE 278e-24 Monomorphic VT at rate of 170/min. The RBBB morphology in V_1 and the R:S ratio < 1 in V_6 are both suggestive of VT. The morphology of the VT is suggestive of origin from the left side of the heart, near the base (RBBB with inferior/rightward axis). Baseline artifact is present in leads V_1 – V_3 .

SECTION 4 DISORDERS OF THE HEART

279 Heart Failure: Pathophysiology and Diagnosis

Douglas L. Mann, Murali Chakinala

HEART FAILURE

DEFINITION

Despite repeated attempts to develop a mechanistic definition that encompasses the heterogeneity and complexity of heart failure (HF), no single conceptual paradigm has withstood the test of time. The current American College of Cardiology Foundation (ACCF)/American Heart Association (AHA) guidelines define HF as a complex clinical syndrome that results from structural or functional impairment of ventricular filling or ejection of blood, which in turn leads to the cardinal clinical symptoms of dyspnea and fatigue and signs of HF, namely edema and rales. Because many patients present without signs

or symptoms of volume overload, the term “heart failure” is preferred over the older term “congestive heart failure.”

EPIDEMIOLOGY



HF is a burgeoning problem worldwide, with more than 20 million people affected. The overall prevalence of HF in the adult population in developed countries is 2%. HF prevalence follows an exponential pattern, rising with age, and affects 6–10% of people over age 65. Although the relative incidence of HF is lower in women than in men, women constitute at least one-half the cases of HF because of their longer life expectancy. In North America and Europe, the lifetime risk of developing HF is approximately one in five for a 40-year-old. The overall prevalence of HF is thought to be increasing, in part because current therapies for cardiac disorders, such as myocardial infarction (MI), valvular heart disease, and arrhythmias, are allowing patients to survive longer. Very little is known about the prevalence or risk of developing HF in emerging nations because of the lack of population-based studies in those countries. HF was once thought to arise primarily in the setting of a

depressed left ventricular (LV) ejection fraction (EF); however, epidemiologic studies have shown that approximately one-half of patients who develop HF have a normal or preserved EF (EF \geq 50%). Accordingly, the historical terms “systolic” and “diastolic” HF have been abandoned, and HF patients are now broadly categorized into HF with a reduced EF (HFrEF; formerly *systolic failure*) or HF with a preserved EF (HRpEF; formerly *diastolic failure*).

ETIOLOGY

As shown in **Table 279-1**, any condition that leads to an alteration in LV structure or function can predispose a patient to developing HF. Although the etiology of HF in patients with a preserved EF differs from that of patients with depressed EF, there is considerable overlap between the etiologies of these two conditions. In industrialized countries, coronary artery disease (CAD) has become the predominant cause in men and women and is responsible for 60–75% of cases of HF. Hypertension contributes to the development of HF in 75% of patients, including most patients with CAD. Both CAD and hypertension interact to augment the risk of HF, as does diabetes mellitus.

In 20–30% of the cases of HF with a depressed EF, the exact etiologic basis is not known. These patients are referred to as having nonischemic, dilated, or idiopathic cardiomyopathy if the cause is unknown (**Chap. 287**). Prior viral infection or toxin exposure (e.g., alcoholic or chemotherapeutic) also may lead to a dilated cardiomyopathy. Moreover, it is becoming increasingly clear that a large number of cases of dilated cardiomyopathy are secondary to specific genetic defects, most notably those in the cytoskeleton. Most forms of familial dilated cardiomyopathy are inherited in an autosomal dominant fashion. Mutations of genes that encode cytoskeletal proteins (desmin, cardiac myosin, vinculin) and nuclear membrane proteins (laminin) have been identified thus far. Dilated cardiomyopathy also is associated with Duchenne’s, Becker’s, and limb-girdle muscular dystrophies. Conditions that lead to a high cardiac output (e.g., arteriovenous fistula, anemia) are seldom responsible for the development of HF in a normal heart; however, in the presence of underlying structural heart disease, these conditions can lead to overt HF.

TABLE 279-1 ETIOLOGIES OF HEART FAILURE

Depressed Ejection Fraction (<40%)	
Coronary artery disease	Nonischemic dilated cardiomyopathy
Myocardial infarction ^a	Familial/genetic disorders
Myocardial ischemia ^a	Infiltrative disorders ^a
Chronic pressure overload	Toxic/drug-induced damage
Hypertension ^a	Metabolic disorder ^a
Obstructive valvular disease ^a	Viral
Chronic volume overload	Chagas’ disease
Regurgitant valvular disease	Disorders of rate and rhythm
Intracardiac (left-to-right) shunting	Chronic bradyarrhythmias
Extracardiac shunting	Chronic tachyarrhythmias
Chronic lung disease	
Cor pulmonale	
Pulmonary vascular disorders	
Preserved Ejection Fraction (>40–50%)	
Pathologic hypertrophy	Restrictive cardiomyopathy
Primary (hypertrophic cardiomyopathies)	Infiltrative disorders (amyloidosis, sarcoidosis)
Secondary (hypertension)	Storage diseases (hemosiderinosis)
Aging	Fibrosis
	Endomyocardial disorders
High-Output States	
Metabolic disorders	Excessive blood flow requirements
Thyrotoxicosis	Systemic arteriovenous shunting
Nutritional disorders (beriberi)	Chronic anemia

^aIndicates conditions that can also lead to heart failure with a preserved ejection fraction.

TABLE 279-2 NEW YORK HEART ASSOCIATION CLASSIFICATION

Functional Capacity	Objective Assessment
Class I	Patients with cardiac disease but without resulting limitation of physical activity. Ordinary physical activity does not cause undue fatigue, palpitations, dyspnea, or anginal pain.
Class II	Patients with cardiac disease resulting in slight limitation of physical activity. They are comfortable at rest. Ordinary physical activity results in fatigue, palpitation, dyspnea, or anginal pain.
Class III	Patients with cardiac disease resulting in marked limitation of physical activity. They are comfortable at rest. Less than ordinary activity causes fatigue, palpitation, dyspnea, or anginal pain.
Class IV	Patients with cardiac disease resulting in inability to carry on any physical activity without discomfort. Symptoms of heart failure or the anginal syndrome may be present even at rest. If any physical activity is undertaken, discomfort is increased.

Source: Adapted from New York Heart Association, Inc., *Diseases of the Heart and Blood Vessels: Nomenclature and Criteria for Diagnosis*, 6th ed. Boston, Little Brown, 1964, p. 114.

GLOBAL CONSIDERATIONS



Rheumatic heart disease remains a major cause of HF in Africa and Asia, especially in the young. Hypertension is an important cause of HF in the African and African-American populations. Chagas’ disease is still a major cause of HF in South America. Not surprisingly, anemia is a frequent concomitant factor in HF in many developing nations. As developing nations undergo socioeconomic development, the epidemiology of HF is becoming similar to that of Western Europe and North America, with CAD emerging as the single most common cause of HF. Although the contribution of diabetes mellitus to HF is not well understood, diabetes accelerates atherosclerosis and often is associated with hypertension.

PROGNOSIS

Despite many recent advances in the evaluation and management of HF, the development of symptomatic HF still carries a poor prognosis. Community-based studies indicate that 30–40% of patients die within 1 year of diagnosis and 60–70% die within 5 years, mainly from worsening HF or as a sudden event (probably because of a ventricular arrhythmia). Although it is difficult to predict prognosis in an individual, patients with symptoms at rest (New York Heart Association [NYHA] class IV) have a 30–70% annual mortality rate, whereas patients with symptoms with moderate activity (NYHA class II) have an annual mortality rate of 5–10%. Thus, functional status is an important predictor of patient outcome (**Table 279-2**).

PATHOGENESIS

Figure 279-1 provides a general conceptual framework for considering the development and progression of HFrEF. As shown, HF may be viewed as a progressive disorder that is initiated after an *index event* either damages the heart muscle, with a resultant loss of functioning cardiac myocytes, or, alternatively, disrupts the ability of the myocardium to generate force, thereby preventing the heart from contracting normally. This index event may have an abrupt onset, as in the case of an MI; it may have a gradual or insidious onset, as in the case of hemodynamic pressure or volume overloading; or it may be hereditary, as in the case of many of the genetic cardiomyopathies. Regardless of the nature of the inciting event, the feature that is common to each of these index events is that they all in some manner produce a decline in the pumping capacity of the heart. In most instances, patients remain asymptomatic or minimally symptomatic after the initial decline in pumping capacity of the heart or develop symptoms only after the dysfunction has been present for some time.

Although the precise reasons why patients with LV dysfunction may remain asymptomatic is not certain, one potential explanation is that a number of compensatory mechanisms become activated in the presence of cardiac injury and/or LV dysfunction allowing patients

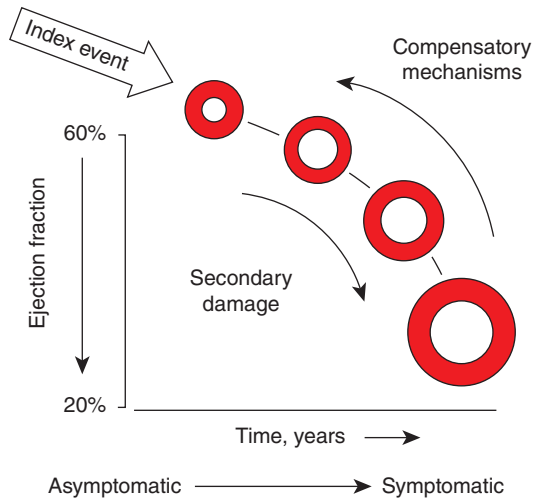


FIGURE 279-1 Pathogenesis of heart failure with a depressed ejection fraction. Heart failure begins after an index event produces an initial decline in the heart's pumping capacity. After this initial decline in pumping capacity, a variety of compensatory mechanisms are activated, including the adrenergic nervous system, the renin-angiotensin-aldosterone system, and the cytokine system. In the short term, these systems are able to restore cardiovascular function to a normal homeostatic range with the result that the patient remains asymptomatic. However, with time, the sustained activation of these systems can lead to secondary end-organ damage within the ventricle, with worsening left ventricular remodeling and subsequent cardiac decompensation. (From D Mann: *Circulation* 100:999, 1999.)

to sustain and modulate LV function for a period of months to years. The compensatory mechanisms that have been described thus far include (1) activation of the renin-angiotensin-aldosterone (RAA) and adrenergic nervous systems, which are responsible, respectively, for maintaining cardiac output through increased retention of salt and water (Fig. 279-2), and (2) increased myocardial contractility. In addition, there is activation of a family of countervailing vasodilatory molecules, including the atrial and brain natriuretic peptides (ANP and BNP), prostaglandins (PGE₂ and PGI₂), and nitric oxide (NO), that offsets the excessive peripheral vascular vasoconstriction. Genetic background, sex, age, or environment may influence these compensatory mechanisms, which are able to modulate LV function within a physiologic/homeostatic range so that the functional capacity of the patient is preserved or is depressed only minimally. Thus, patients may remain asymptomatic or minimally symptomatic for a period of years; however, at some point patients become overtly symptomatic, with a resultant striking increase in morbidity and mortality rates. Although the exact mechanisms that are responsible for this transition are not known, as will be discussed below, the transition to symptomatic HF is accompanied by increasing activation of neurohormonal, adrenergic, and cytokine systems that lead to a series of adaptive changes within the myocardium collectively referred to as *LV remodeling*.

In contrast to our understanding of the pathogenesis of HF with a depressed EF, our understanding of the mechanisms that contribute to the development of HF with a preserved EF is still evolving. That is, although diastolic dysfunction (see below) was thought to be the only mechanism responsible for the development of HF with a preserved EF, community-based studies suggest that additional extracardiac mechanisms may be important, such as increased vascular stiffness and impaired renal function.

BASIC MECHANISMS OF HEART FAILURE

Heart Failure with a Reduced Ejection Fraction LV remodeling develops in response to a series of complex events that occur at the cellular and molecular levels (Table 279-3). These changes include (1) myocyte hypertrophy; (2) alterations in the contractile properties of the myocyte; (3) progressive loss of myocytes through necrosis,

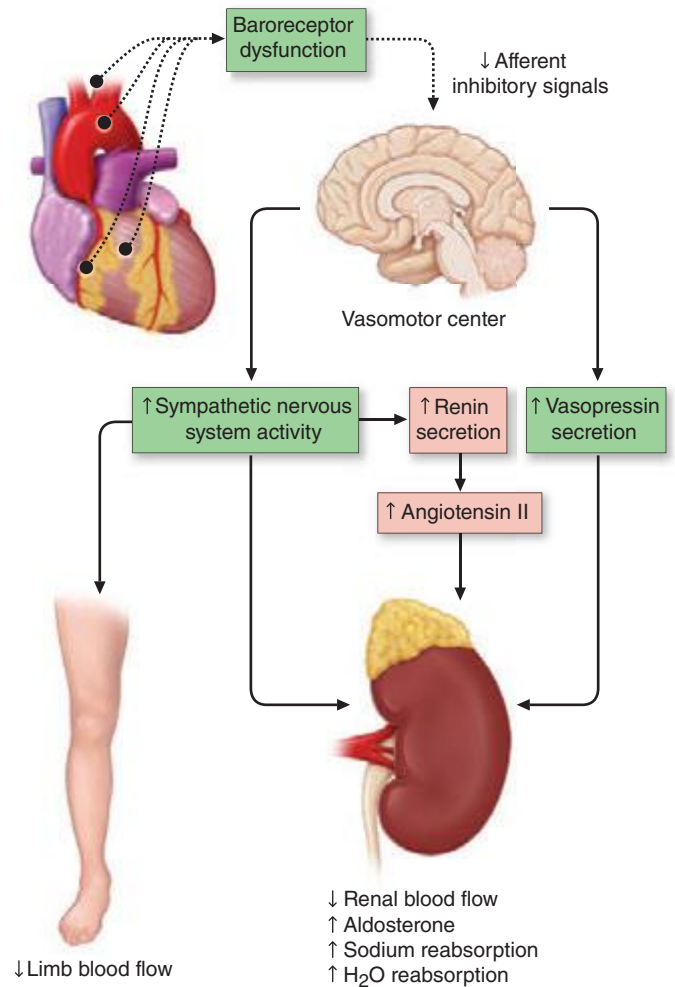


FIGURE 279-2 Activation of neurohormonal systems in heart failure. The decreased cardiac output in heart failure (HF) patients results in an "unloading" of high-pressure baroreceptors (circles) in the left ventricle, carotid sinus, and aortic arch. This unloading of the peripheral baroreceptors leads to a loss of inhibitory parasympathetic tone to the central nervous system (CNS), with a resultant generalized increase in efferent sympathetic tone, and nonosmotic release of arginine vasopressin (AVP) from the pituitary. AVP (or antidiuretic hormone [ADH]) is a powerful vasoconstrictor that increases the permeability of the renal collecting ducts, leading to the reabsorption of free water. These afferent signals to the CNS also activate efferent sympathetic nervous system pathways that innervate the heart, kidney, peripheral vasculature, and skeletal muscles.

Sympathetic stimulation of the kidney leads to the release of renin, with a resultant increase in the circulating levels of angiotensin II and aldosterone. The activation of the renin-angiotensin-aldosterone system promotes salt and water retention and leads to vasoconstriction of the peripheral vasculature, myocyte hypertrophy, myocyte cell death, and myocardial fibrosis. Although these neurohormonal mechanisms facilitate short-term adaptation by maintaining blood pressure, and hence perfusion to vital organs, the same neurohormonal mechanisms are believed to contribute to end-organ changes in the heart and the circulation and to the excessive salt and water retention in advanced HF. (Modified from A Nohria et al: *Neurohormonal, renal and vascular adjustments, in Atlas of Heart Failure: Cardiac Function and Dysfunction*, 4th ed, WS Colucci [ed]. Philadelphia, Current Medicine Group 2002, p. 104.)

apoptosis, and autophagic cell death; (4) β -adrenergic desensitization; (5) abnormal myocardial energetics and metabolism; and (6) reorganization of the extracellular matrix with dissolution of the organized structural collagen weave surrounding myocytes and subsequent replacement by an interstitial collagen matrix that does not

TABLE 279-3 OVERVIEW OF LEFT VENTRICULAR REMODELING

Alterations in Myocyte Biology	
Excitation-contraction coupling	
Myosin heavy chain (fetal) gene expression	
β-Adrenergic desensitization	
Hypertrophy	
Myocytolysis	
Cytoskeletal proteins	
Myocardial Changes	
Myocyte loss	
Necrosis	
Apoptosis	
Autophagy	
Alterations in extracellular matrix	
Matrix degradation	
Myocardial fibrosis	
Alterations in Left Ventricular Chamber Geometry	
Left ventricular (LV) dilation	
Increased LV sphericity	
LV wall thinning	
Mitral valve incompetence	

Source: Adapted from D. Mann: Pathophysiology of heart failure, in *Braunwald's Heart Disease*, 8th ed, PL Libby et al (eds). Philadelphia, Elsevier, 2008, p. 550.

provide structural support to the myocytes. The biologic stimuli for these profound changes include mechanical stretch of the myocyte, circulating neurohormones (e.g., norepinephrine, angiotensin II), inflammatory cytokines (e.g., tumor necrosis factor [TNF]), other peptides and growth factors (e.g., endothelin), and reactive oxygen species (e.g., superoxide). The sustained overexpression of these biologically active molecules is believed to contribute to the progression of HF by virtue of the deleterious effects they exert on the heart and the circulation. Indeed, this insight forms the clinical rationale for using pharmacologic agents that antagonize these systems (e.g., angiotensin-converting enzyme [ACE] inhibitors and beta blockers) in treating patients with HF (**Chap. 280**).

To understand how the changes that occur in the failing cardiac myocyte contribute to depressed LV systolic function in HF, it is instructive first to review the biology of the cardiac muscle cell (**Chap. 265e**). Sustained neurohormonal activation and mechanical overload result in transcriptional and posttranscriptional changes in the genes and proteins that regulate excitation-contraction coupling and cross-bridge interaction (see **Figs. 265e-6 and 265e-7**). The changes that regulate excitation-contraction include decreased function of sarcoplasmic reticulum Ca²⁺ adenosine triphosphatase (SERCA2A), resulting in decreased calcium uptake into the sarcoplasmic reticulum (SR), and hyperphosphorylation of the ryanodine receptor, leading to calcium leakage from the SR. The changes that occur in the cross-bridges include decreased expression of α-myosin heavy chain and increased expression of β-myosin heavy chain, myocytolysis, and disruption of the cytoskeletal links between the sarcomeres and the extracellular matrix. Collectively, these changes impair the ability of the myocyte to contract and therefore contribute to the depressed LV systolic function observed in patients with HF.

Myocardial relaxation is an adenosine triphosphate (ATP)-dependent process that is regulated by uptake of cytoplasmic calcium into the SR by SERCA2A and extrusion of calcium by sarcolemmal pumps (see **Fig. 265e-7**). Accordingly, reductions in ATP concentration, as occurs in ischemia, may interfere with these processes and lead to slowed myocardial relaxation. Alternatively, if LV filling is delayed because LV compliance is reduced (e.g., from hypertrophy or fibrosis), LV filling pressures will similarly remain elevated at end diastole (see **Fig. 265e-11**). An increase in heart rate disproportionately shortens the time for diastolic filling, which may lead to elevated LV filling

pressures, particularly in noncompliant ventricles. Elevated LV end-diastolic filling pressures result in increases in pulmonary capillary pressures, which can contribute to the dyspnea experienced by patients with diastolic dysfunction. In addition to impaired myocardial relaxation, increased myocardial stiffness secondary to cardiac hypertrophy and increased myocardial collagen content may contribute to diastolic failure. Importantly, diastolic dysfunction can occur alone or in combination with systolic dysfunction in patients with HF.

Left Ventricular Remodeling *Ventricular remodeling* refers to the changes in LV mass, volume, and shape and the composition of the heart that occur after cardiac injury and/or abnormal hemodynamic loading conditions. LV remodeling may contribute independently to the progression of HF by virtue of the mechanical burdens that are engendered by the changes in the geometry of the remodeled LV. In addition to the increase in LV end-diastolic volume, LV wall thinning occurs as the left ventricle begins to dilate. The increase in wall thinning, along with the increase in afterload created by LV dilation, leads to a functional *afterload mismatch* that may contribute further to a decrease in stroke volume. Moreover, the high end-diastolic wall stress might be expected to lead to (1) hypoperfusion of the subendocardium, with resultant worsening of LV function; (2) increased oxidative stress, with the resultant activation of families of genes that are sensitive to free radical generation (e.g., TNF and interleukin 1β); and (3) sustained expression of stretch-activated genes (angiotensin II, endothelin, and TNF) and/or stretch activation of hypertrophic signaling pathways. Increasing LV dilation also results in tethering of the papillary muscles with resulting incompetence of the mitral valve apparatus and functional mitral regurgitation, which in turn leads to further hemodynamic overloading of the ventricle. Taken together, the mechanical burdens that are engendered by LV remodeling contribute to the progression of HF. Recent studies have shown that LV remodeling can be reversed following medical and device therapy and that reverse LV remodeling is associated with improved clinical outcomes in patients with HFrEF. Indeed, one of the goals of therapy for HF is to prevent and/or reverse LV remodeling.

CLINICAL MANIFESTATIONS

Symptoms The cardinal symptoms of HF are fatigue and shortness of breath. Although fatigue traditionally has been ascribed to the low cardiac output in HF, it is likely that skeletal-muscle abnormalities and other noncardiac comorbidities (e.g., anemia) also contribute to this symptom. In the early stages of HF, dyspnea is observed only during exertion; however, as the disease progresses, dyspnea occurs with less strenuous activity, and it ultimately may occur even at rest. The origin of dyspnea in HF is probably multifactorial (**Chap. 47e**). The most important mechanism is pulmonary congestion with accumulation of interstitial or intra-alveolar fluid, which activates juxtacapillary J receptors, which in turn stimulate the rapid, shallow breathing characteristic of cardiac dyspnea. Other factors that contribute to dyspnea on exertion include reductions in pulmonary compliance, increased airway resistance, respiratory muscle and/or diaphragm fatigue, and anemia. Dyspnea may become less frequent with the onset of right ventricular (RV) failure and tricuspid regurgitation.

ORTHOPNEA Orthopnea, which is defined as dyspnea occurring in the recumbent position, is usually a later manifestation of HF than is exertional dyspnea. It results from redistribution of fluid from the splanchnic circulation and lower extremities into the central circulation during recumbency, with a resultant increase in pulmonary capillary pressure. Nocturnal cough is a common manifestation of this process and a frequently overlooked symptom of HF. Orthopnea generally is relieved by sitting upright or sleeping with additional pillows. Although orthopnea is a relatively specific symptom of HF, it may occur in patients with abdominal obesity or ascites and patients with pulmonary disease whose lung mechanics favor an upright posture.

PAROXYSMAL NOCTURNAL DYSPNEA (PND) This term refers to acute episodes of severe shortness of breath and coughing that generally occur at night and awaken the patient from sleep, usually 1–3 h after the

1504 patient retires. PND may manifest as coughing or wheezing, possibly because of increased pressure in the bronchial arteries leading to airway compression, along with interstitial pulmonary edema that leads to increased airway resistance. Whereas orthopnea may be relieved by sitting upright at the side of the bed with the legs in a dependent position, patients with PND often have persistent coughing and wheezing even after they have assumed the upright position. *Cardiac asthma* is closely related to PND, is characterized by wheezing secondary to bronchospasm, and must be differentiated from primary asthma and pulmonary causes of wheezing.

CHEYNE-STOKES RESPIRATION Also referred to as periodic respiration or cyclic respiration, Cheyne-Stokes respiration is present in 40% of patients with advanced HF and usually is associated with low cardiac output. Cheyne-Stokes respiration is caused by an increased sensitivity of the respiratory center to arterial PCO_2 . There is an apneic phase, during which arterial PO_2 falls and arterial PCO_2 rises. These changes in the arterial blood gas content stimulate the respiratory center, resulting in hyperventilation and hypocapnia, followed by recurrence of apnea. Cheyne-Stokes respirations may be perceived by the patient or the patient's family as severe dyspnea or as a transient cessation of breathing.

ACUTE PULMONARY EDEMA See Chap. 326

Other Symptoms Patients with HF also may present with gastrointestinal symptoms. Anorexia, nausea, and early satiety associated with abdominal pain and fullness are common complaints and may be related to edema of the bowel wall and/or a congested liver. Congestion of the liver and stretching of its capsule may lead to right upper-quadrant pain. Cerebral symptoms such as confusion, disorientation, and sleep and mood disturbances may be observed in patients with severe HF, particularly elderly patients with cerebral arteriosclerosis and reduced cerebral perfusion. Nocturia is common in HF and may contribute to insomnia.

PHYSICAL EXAMINATION

A careful physical examination is always warranted in the evaluation of patients with HF. The purpose of the examination is to help determine the cause of HF as well as to assess the severity of the syndrome. Obtaining additional information about the hemodynamic profile and the response to therapy and determining the prognosis are important additional goals of the physical examination.

General Appearance and Vital Signs In mild or moderately severe HF, the patient appears to be in no distress at rest except for feeling uncomfortable when lying flat for more than a few minutes. In more severe HF, the patient must sit upright, may have labored breathing, and may not be able to finish a sentence because of shortness of breath. Systolic blood pressure may be normal or high in early HF, but it generally is reduced in advanced HF because of severe LV dysfunction. The pulse pressure may be diminished, reflecting a reduction in stroke volume. Sinus tachycardia is a nonspecific sign caused by increased adrenergic activity. Peripheral vasoconstriction leading to cool peripheral extremities and cyanosis of the lips and nail beds is also caused by excessive adrenergic activity.

Jugular Veins (See also Chap. 267) Examination of the jugular veins provides an estimation of right atrial pressure. The jugular venous pressure is best appreciated with the patient lying recumbent, with the head tilted at 45°. The jugular venous pressure should be quantified in centimeters of water (normal ≤ 8 cm) by estimating the height of the venous column of blood above the sternal angle in centimeters and then adding 5 cm. In the early stages of HF, the venous pressure may be normal at rest but may become abnormally elevated with sustained (~15 seconds) pressure on the abdomen (positive abdominojugular reflux). Giant *v* waves indicate the presence of tricuspid regurgitation.

Pulmonary Examination Pulmonary crackles (rales or crepitations) result from the transudation of fluid from the intravascular space into

the alveoli. In patients with pulmonary edema, rales may be heard widely over both lung fields and may be accompanied by expiratory wheezing (cardiac asthma). When present in patients without concomitant lung disease, rales are specific for HF. Importantly, rales are frequently absent in patients with chronic HF, even when LV filling pressures are elevated, because of increased lymphatic drainage of alveolar fluid. Pleural effusions result from the elevation of pleural capillary pressure and the resulting transudation of fluid into the pleural cavities. Since the pleural veins drain into both the systemic and the pulmonary veins, pleural effusions occur most commonly with biventricular failure. Although pleural effusions are often bilateral in HF, when they are unilateral, they occur more frequently in the right pleural space.

Cardiac Examination Examination of the heart, although essential, frequently does not provide useful information about the severity of HF. If cardiomegaly is present, the point of maximal impulse (PMI) usually is displaced below the fifth intercostal space and/or lateral to the midclavicular line, and the impulse is palpable over two interspaces. Severe LV hypertrophy leads to a sustained PMI. In some patients, a third heart sound (S_3) is audible and palpable at the apex. Patients with enlarged or hypertrophied right ventricles may have a sustained and prolonged left parasternal impulse extending throughout systole. An S_3 (or *protodiastolic gallop*) is most commonly present in patients with volume overload who have tachycardia and tachypnea, and it often signifies severe hemodynamic compromise. A fourth heart sound (S_4) is not a specific indicator of HF but is usually present in patients with diastolic dysfunction. The murmurs of mitral and tricuspid regurgitation are frequently present in patients with advanced HF.

Abdomen and Extremities Hepatomegaly is an important sign in patients with HF. When it is present, the enlarged liver is frequently tender and may pulsate during systole if tricuspid regurgitation is present. Ascites, a late sign, occurs as a consequence of increased pressure in the hepatic veins and the veins draining the peritoneum. Jaundice, also a late finding in HF, results from impairment of hepatic function secondary to hepatic congestion and hepatocellular hypoxemia and is associated with elevations of both direct and indirect bilirubin.

Peripheral edema is a cardinal manifestation of HF, but it is nonspecific and usually is absent in patients who have been treated adequately with diuretics. Peripheral edema is usually symmetric and dependent in HF and occurs predominantly in the ankles and the pretibial region in ambulatory patients. In bedridden patients, edema may be found in the sacral area (*presacral edema*) and the scrotum. Long-standing edema may be associated with indurated and pigmented skin.

Cardiac Cachexia With severe chronic HF, there may be marked weight loss and cachexia. Although the mechanism of cachexia is not entirely understood, it is probably multifactorial and includes elevation of the resting metabolic rate; anorexia, nausea, and vomiting due to congestive hepatomegaly and abdominal fullness; elevation of circulating concentrations of cytokines such as TNF; and impairment of intestinal absorption due to congestion of the intestinal veins. When present, cachexia augurs a poor overall prognosis.

DIAGNOSIS

The diagnosis of HF is relatively straightforward when the patient presents with classic signs and symptoms of HF; however, the signs and symptoms of HF are neither specific nor sensitive. Accordingly, the key to making the diagnosis is to have a high index of suspicion, particularly for high-risk patients. When these patients present with signs or symptoms of HF, additional laboratory testing should be performed.

Routine Laboratory Testing Patients with new-onset HF and those with chronic HF and acute decompensation should have a complete blood count, a panel of electrolytes, blood urea nitrogen, serum creatinine, hepatic enzymes, and a urinalysis. Selected patients should have assessment for diabetes mellitus (fasting serum glucose or oral glucose

tolerance test), dyslipidemia (fasting lipid panel), and thyroid abnormalities (thyroid-stimulating hormone level).

Electrocardiogram (ECG) A routine 12-lead ECG is recommended. The major importance of the ECG is to assess cardiac rhythm and determine the presence of LV hypertrophy or a prior MI (presence or absence of Q waves) as well as to determine QRS width to ascertain whether the patient may benefit from resynchronization therapy (see below). A normal ECG virtually excludes LV systolic dysfunction.

Chest X-Ray A chest x-ray provides useful information about cardiac size and shape, as well as the state of the pulmonary vasculature, and may identify noncardiac causes of the patient's symptoms. Although patients with acute HF have evidence of pulmonary hypertension, interstitial edema, and/or pulmonary edema, the majority of patients with chronic HF do not. The absence of these findings in patients with chronic HF reflects the increased capacity of the lymphatics to remove interstitial and/or pulmonary fluid.

Assessment of LV Function Noninvasive cardiac imaging ([Chap. 270e](#)) is essential for the diagnosis, evaluation, and management of HF. The most useful test is the two-dimensional (2-D) echocardiogram/Doppler, which can provide a semiquantitative assessment of LV size and function as well as the presence or absence of valvular and/or regional wall motion abnormalities (indicative of a prior MI). The presence of left atrial dilation and LV hypertrophy, together with abnormalities of LV diastolic filling provided by pulse-wave and tissue Doppler, is useful for the assessment of HF with a preserved EF. The 2-D echocardiogram/Doppler is also invaluable in assessing RV size and pulmonary pressures, which are critical in the evaluation and management of cor pulmonale (see below). Magnetic resonance imaging (MRI) also provides a comprehensive analysis of cardiac anatomy and function and is now the gold standard for assessing LV mass and volumes. MRI also is emerging as a useful and accurate imaging modality for evaluating patients with HF, both in terms of assessing LV structure and for determining the cause of HF (e.g., amyloidosis, ischemic cardiomyopathy, hemochromatosis).

The most useful index of LV function is the EF (stroke volume divided by end-diastolic volume). Because the EF is easy to measure by noninvasive testing and easy to conceptualize, it has gained wide acceptance among clinicians. Unfortunately, the EF has a number of limitations as a true measure of contractility, since it is influenced by alterations in afterload and/or preload. Nonetheless, with the exceptions indicated above, when the EF is normal ($\geq 50\%$), systolic function is usually adequate, and when the EF is significantly depressed ($<30\text{--}40\%$), contractility is usually depressed.

Biomarkers Circulating levels of natriuretic peptides are useful and important adjunctive tools in the diagnosis of patients with HF. Both B-type natriuretic peptide (BNP) and N-terminal pro-BNP (NT-proBNP), which are released from the failing heart, are relatively sensitive markers for the presence of HF with depressed EF; they also are elevated in HF patients with a preserved EF, albeit to a lesser degree. In ambulatory patients with dyspnea, the measurement of BNP or NT-proBNP is useful to support clinical decision making regarding the diagnosis of HF, especially in the setting of clinical uncertainty. Moreover, the measurement of BNP or NT-proBNP is useful for establishing prognosis or disease severity in chronic HF and can be useful to achieve optimal dosing of medical therapy in select clinically euvolemic patients. However, it is important to recognize that natriuretic peptide levels increase with age and renal impairment, are more elevated in women, and can be elevated in right HF from any cause. Levels can be falsely low in obese patients. Other biomarkers, such as soluble ST-2 and galectin-3, are newer biomarkers that can be used for determining the prognosis of HF patients.

Exercise Testing Treadmill or bicycle exercise testing is not routinely advocated for patients with HF, but either is useful for assessing the need for cardiac transplantation in patients with advanced HF

([Chap. 281](#)). A peak oxygen uptake (vo_2) <14 mL/kg per min is associated with a relatively poor prognosis. Patients with a vo_2 <14 mL/kg per min have been shown, in general, to have better survival when transplanted than when treated medically.

DIFFERENTIAL DIAGNOSIS

HF resembles but should be distinguished from (1) conditions in which there is circulatory congestion secondary to abnormal salt and water retention but in which there is no disturbance of cardiac structure or function (e.g., renal failure), and (2) noncardiac causes of pulmonary edema (e.g., acute respiratory distress syndrome). In most patients who present with classic signs and symptoms of HF, the diagnosis is relatively straightforward. However, even experienced clinicians have difficulty differentiating the dyspnea that arises from cardiac and pulmonary causes ([Chap. 47e](#)). In this regard, noninvasive cardiac imaging, biomarkers, pulmonary function testing, and chest x-ray may be useful. A very low BNP or NT-proBNP may be helpful in excluding a cardiac cause of dyspnea in this setting. Ankle edema may arise secondary to varicose veins, obesity, renal disease, or gravitational effects. When HF develops in patients with a preserved EF, it may be difficult to determine the relative contribution of HF to the dyspnea that occurs in chronic lung disease and/or obesity.

COR PULMONALE

DEFINITION

Cor pulmonale, often referred to as *pulmonary heart disease*, can be defined as altered RV structure and/or function in the context of chronic lung disease and is triggered by the onset of pulmonary hypertension. Although RV dysfunction is also an important sequela of HFpEF and HFrEF, this is not considered as cor pulmonale.

ETIOLOGY AND EPIDEMIOLOGY

Cor pulmonale develops in response to acute or chronic changes in the pulmonary vasculature and/or the lung parenchyma that are sufficient to cause pulmonary hypertension. The true prevalence of cor pulmonale is difficult to ascertain. First, not all patients with chronic lung disease will develop cor pulmonale, which may be subclinical in compensated individuals. Second, our ability to diagnose pulmonary hypertension and cor pulmonale by routine physical examination and laboratory testing is relatively insensitive. However, advances in 2-D echo/Doppler imaging and biomarkers (BNP) can make it easier to identify cor pulmonale.

Once patients with chronic pulmonary or pulmonary vascular disease develop cor pulmonale, the prognosis worsens. Although chronic obstructive pulmonary disease (COPD) and chronic bronchitis are responsible for approximately 50% of the cases of cor pulmonale in North America ([Chap. 314](#)), any disease that affects the pulmonary vasculature ([Chap. 304](#)) or parenchyma can lead to cor pulmonale ([Table 279-4](#)). Primary pulmonary vascular disorders are relatively rare causes of cor pulmonale, but cor pulmonale is extremely common with these conditions, given the magnitude of pulmonary hypertension present.

PATHOPHYSIOLOGY AND BASIC MECHANISMS

Although many conditions can lead to cor pulmonale, the common pathophysiologic mechanism is pulmonary hypertension that is sufficient to alter RV structure (i.e., dilation with or without hypertrophy) and function. Normally, pulmonary artery pressures are only ~ 15 mmHg and do not increase even with multiples of resting cardiac output, because of vasodilation and blood vessel recruitment of the pulmonary circulatory bed. But, in the setting of parenchymal lung diseases, primary pulmonary vascular disorders, or chronic (alveolar) hypoxia, the circulatory bed undergoes varying degrees of vascular remodeling, vasoconstriction, and destruction. As a result, pulmonary artery pressures and RV afterload increase, setting the stage for cor pulmonale ([Table 279-4](#)). The systemic consequences of cor pulmonale relate to alterations in cardiac output as well as salt and

TABLE 279-4 ETIOLOGY OF CHRONIC COR PULMONALE

Diseases of the Lung Parenchyma
Chronic obstructive pulmonary disease
Emphysema
Chronic bronchitis
Cystic fibrosis
Idiopathic interstitial pneumonitis
Idiopathic pulmonary fibrosis
Nonspecific interstitial pneumonitis
Sarcoidosis
Bronchiectasis
Pulmonary Langerhans cell histiocytosis
Lymphangioliomyomatosis
Disorders of Chronic (Alveolar) Hypoxia
Alveolar hypoventilation syndromes
Obesity hypoventilation syndrome
Central hypoventilation syndrome
Neuromuscular respiratory failure
Chest wall disorders
Kyphoscoliosis
Living at high altitude
Diseases of the Pulmonary Vasculature
Pulmonary arterial hypertension (PAH)
Idiopathic PAH
Heritable PAH
Associated PAH
Venoocclusive disease
Chronic thromboembolic pulmonary hypertension
Pulmonary tumor thrombotic microangiopathy

water homeostasis. Anatomically, the RV is a thin-walled, compliant chamber that is better suited to handle volume overload than pressure overload. Thus, the sustained pressure overload imposed by pulmonary hypertension and increased pulmonary vascular resistance eventually causes the RV to fail.

The response of the RV to pulmonary hypertension depends on the acuteness and severity of the pressure overload. Acute cor pulmonale occurs after a sudden and severe stimulus (e.g., massive pulmonary embolus), with RV dilatation and failure but no RV hypertrophy (Chap. 300). Chronic cor pulmonale, however, is associated with a more slowly evolving and progressive pulmonary hypertension that leads to initial modest RV hypertrophy and subsequent RV dilation. Acute decompensation of previously compensated chronic cor pulmonale is a common clinical occurrence. Triggers include worsening hypoxia from any cause (e.g., pneumonia), acidemia (e.g., exacerbation of COPD), acute pulmonary embolus, atrial tachyarrhythmia, hypervolemia, and mechanical ventilation that leads to compressive forces on alveolar blood vessels.

CLINICAL MANIFESTATIONS

Symptoms The symptoms of chronic cor pulmonale generally are related to the underlying pulmonary disorder. Dyspnea, the most common symptom, is usually the result of the increased work of

breathing secondary to changes in elastic recoil of the lung (fibrosing lung diseases), altered respiratory mechanics (e.g., overinflation with COPD), or inefficient ventilation (e.g., primary pulmonary vascular disease). Orthopnea and PND are rarely symptoms of isolated right HF and usually point toward concurrent left heart dysfunction. Rarely, these symptoms reflect increased work of breathing in the supine position resulting from compromised diaphragmatic excursion. Abdominal pain and ascites that occur with cor pulmonale are similar to the right HF that ensues in chronic HF. Lower-extremity edema may occur secondary to neurohormonal activation, elevated RV filling pressures, or increased levels of carbon dioxide and hypoxemia, which can lead to peripheral vasodilation and edema formation.

Signs Many of the signs encountered in cor pulmonale are also present in HF patients with a depressed EF, including tachypnea, elevated jugular venous pressures, hepatomegaly, and lower-extremity edema. Patients may have prominent *v* waves in the jugular venous pulse as a result of tricuspid regurgitation. Other cardiovascular signs include an RV heave palpable along the left sternal border or in the epigastrium. The increase in intensity of the holosystolic murmur of tricuspid regurgitation with inspiration (“Carvallo’s sign”) may be lost eventually as RV failure worsens. Cyanosis is a late finding in cor pulmonale and is secondary to a low cardiac output with systemic vasoconstriction and ventilation-perfusion mismatches in the lung.

DIAGNOSIS

The most common cause of right HF is not pulmonary parenchymal or vascular disease but left HF. Therefore, it is important to evaluate the patient for LV systolic and diastolic dysfunction. The ECG in severe pulmonary hypertension shows P pulmonale, right axis deviation, and RV hypertrophy. Radiographic examination of the chest may show enlargement of the main central pulmonary arteries and hilar vessels. Spirometry and lung volumes can identify obstructive and/or restrictive defects indicative of parenchymal lung diseases; arterial blood gases can demonstrate hypoxemia and/or hypercapnia. Spiral computed tomography (CT) scans of the chest are useful in diagnosing acute thromboembolic disease; however, ventilation-perfusion lung scanning remains best suited for diagnosing *chronic thromboembolic disease* (Chap. 300). A high-resolution CT scan of the chest can identify interstitial lung disease.

Two-dimensional echocardiography is useful for measuring RV thickness and chamber dimensions. Location of the RV behind the sternum and its crescent shape challenge assessment of RV function by echocardiography, especially when parenchymal lung disease is present. Calculated measures of RV function (e.g., tricuspid annular plane systolic excursion [TAPSE] or the Tei Index) supplement more subjective assessments of RV function. The interventricular septum may move paradoxically during systole in the presence of pulmonary hypertension. As noted, Doppler echocardiography can be used to assess pulmonary artery pressures. MRI is also useful for assessing RV structure and function, particularly in patients who are difficult to image with 2-D echocardiography because of severe lung disease. Right-heart catheterization is useful for confirming the diagnosis of pulmonary hypertension and for excluding elevated left-heart pressures (measured as the pulmonary capillary wedge pressure) as a cause for right HF. BNP and N-terminal BNP levels are elevated in patients with cor pulmonale secondary to RV myocardial stretch and may be dramatically elevated in acute pulmonary embolism.

Distinctive phenotypes of presentation with diverse management targets exemplify the vast syndrome of heart failure. These range from chronic heart failure with reduced ejection fraction (HFrEF) or heart failure with preserved ejection fraction (HFpEF), acute decompensated heart failure (ADHF), and advanced heart failure. Early management evolved from symptom control to disease-modifying therapy in HFrEF with the advent of renin-angiotensin-aldosterone system (RAAS)-directed therapy, beta receptor antagonists, mineralocorticoid receptor antagonists, cardiac resynchronization therapy, and implantable cardio-defibrillators. However, similar advances have been elusive in the syndromes of HFpEF and ADHF, which have remained devoid of convincing therapeutic advances to alter their natural history. In advanced heart failure, a stage of disease typically encountered in HFrEF, the patient remains markedly symptomatic with demonstrated refractoriness or inability to tolerate full-dose neurohormonal antagonism, often requires escalating doses of diuretics, and exhibits persistent hyponatremia and renal insufficiency with frequent episodes of heart failure decompensation requiring recurrent hospitalizations. Such individuals are at the highest risk of sudden or progressive pump failure-related deaths (Chap. 281). In contrast, early-stage asymptomatic left ventricular dysfunction is amenable to preventive care, and its natural history is modifiable by neurohormonal antagonism (not further discussed).

HEART FAILURE WITH PRESERVED EJECTION FRACTION

GENERAL PRINCIPLES

Therapeutic targets in HFpEF include control of congestion, stabilization of heart rate and blood pressure, and efforts at improving exercise tolerance. Addressing surrogate targets, such as regression of ventricular hypertrophy in hypertensive heart disease, and use of lusitropic agents, such as calcium channel blockers and beta receptor antagonists, have been disappointing. Experience has demonstrated that lowering blood pressure alleviates symptoms more effectively than targeted therapy with specific agents.

CLINICAL TRIALS IN HFpEF

The Candesartan in Heart Failure—Assessment of Mortality and Morbidity (CHARM) Preserved study showed a statistically significant reduction in hospitalizations but no difference in all-cause mortality in patients with HFpEF who were treated with the angiotensin receptor blocker (ARB), candesartan. Similarly, the Irbesartan in Heart Failure with Preserved Systolic Function (I-PRESERVE) trial demonstrated no differences in meaningful endpoints in such patients treated with irbesartan. An earlier analysis of a subset of the Digitalis Investigation Group (DIG) trial found no role for digoxin in the treatment of HFpEF. In the Study of the Effects of Nebivolol Intervention on Outcomes and Rehospitalization in Seniors with Heart Failure (SENIORS) trial of nebivolol, a vasodilating beta blocker, the subgroup of elderly patients with prior hospitalization and HFpEF did not appear to benefit in terms of all-cause or cardiovascular mortality. Much smaller mechanistic studies in the elderly with the angiotensin-converting enzyme inhibitor (ACEI) enalapril showed no effect on peak exercise oxygen consumption, 6-minute walk distance, aortic distensibility, left ventricular mass, or peripheral neurohormone expression.

NOVEL TARGETS

A small trial demonstrated that the phosphodiesterase-5 inhibitor *sildenafil* improved filling pressures and right ventricular function in a cohort of HFpEF patients with pulmonary venous hypertension. This finding led to the phase II trial, Phosphodiesterase-5 Inhibition to Improve Clinical Status and Exercise Capacity in Diastolic Heart Failure (RELAX), in HFpEF patients (left ventricular ejection fraction [LVEF] >50%) with New York Heart Association (NYHA) functional

class II or III symptoms, who received sildenafil at 20 mg three times daily for 3 months, followed by 60 mg three times daily for another 3 months, compared with a placebo. There was no improvement in functional capacity, quality of life, or other clinical and surrogate parameters. Conceptually targeting myocardial fibrosis in HFpEF, the large-scale Aldosterone Antagonist Therapy in Adults with Preserved Ejection Fraction Congestive Heart Failure (TOPCAT) trial has been completed. This trial demonstrated no improvement in the primary composite end-point, but did show a secondary signal of benefit on HF hospitalizations, counterbalanced, however, by an increase in adverse effects, particularly hyperkalemia. However, pessimism has been generated by the negative outcome of the Aldosterone Receptor Blockade in Diastolic Heart Failure (ALDO-DHF) study wherein *spironolactone* improved echocardiographic indices of diastolic dysfunction but failed to improve exercise capacity, symptoms, or quality-of-life measures. A unique molecule that hybridizes an ARB with an endopeptidase inhibitor, LCZ696, increases the generation of myocardial cyclic guanosine 3',5'-monophosphate, enhances myocardial relaxation, and reduces ventricular hypertrophy. This dual blocker has been shown to reduce circulating natriuretic peptides and reduce left atrial size to a significantly greater extent than valsartan alone in patients with HFpEF.

CLINICAL PEARLS

Even as efforts to control hypertension in HFpEF are critical, evaluation for and correction of underlying ischemia may be beneficial. Appropriate identification and treatment of sleep-disordered breathing should be strongly considered. Excessive decrease in preload with vasodilators may lead to underfilling the ventricle and subsequent hypotension and syncope. Some investigators have suggested that the exercise intolerance in HFpEF is a manifestation of chronotropic insufficiency and that such aberrations could be corrected with use of rate responsive pacemakers, but this remains an inadequately investigated contention (Fig. 280-1).

ACUTE DECOMPENSATED HEART FAILURE

GENERAL PRINCIPLES

ADHF is a heterogeneous clinical syndrome most often resulting in need for hospitalization due to confluence of interrelated abnormalities of decreased cardiac performance, renal dysfunction, and alterations in vascular compliance. Admission with a diagnosis of ADHF is associated with excessive morbidity and mortality, with nearly half of these patients readmitted for management within 6 months, and a high short-term (5–8% in-hospital) and long-term mortality (20% at 1 year). Importantly, long-term aggregate outcomes remain poor, with a combined incidence of cardiovascular deaths, heart failure hospitalizations, myocardial infarction, strokes, or sudden death reaching 50% at 12 months after hospitalization. The management of these patients has remained difficult and principally revolves around volume control and decrease of vascular impedance while maintaining attention to end-organ perfusion (coronary and renal).

The first principle of management of these patients is to identify and tackle known precipitants of decompensation. Identification and management of medication nonadherence and use of prescribed medicines such as nonsteroidal anti-inflammatory drugs, cold and flu preparations with cardiac stimulants, and herbal preparations, including licorice, ginseng, and ma huang (an herbal form of ephedrine now banned in most places), are required. Active infection and overt or covert pulmonary thromboembolism should be sought, identified, and treated when clinical clues suggest such direction. When possible, arrhythmias should be corrected by controlling heart rate or restoring sinus rhythm in patients with poorly tolerated rapid atrial fibrillation and by correcting ongoing ischemia with coronary revascularization or by correcting offenders such as ongoing bleeding in demand-related ischemia. A parallel step in management involves stabilization of hemodynamics in those with instability. The routine use of a pulmonary artery catheter is not recommended and should be restricted to those who respond poorly to diuresis or experience hypotension or signs and symptoms suggestive of a low cardiac output where therapeutic targets are unclear. Analysis of in-hospital registries has identified several

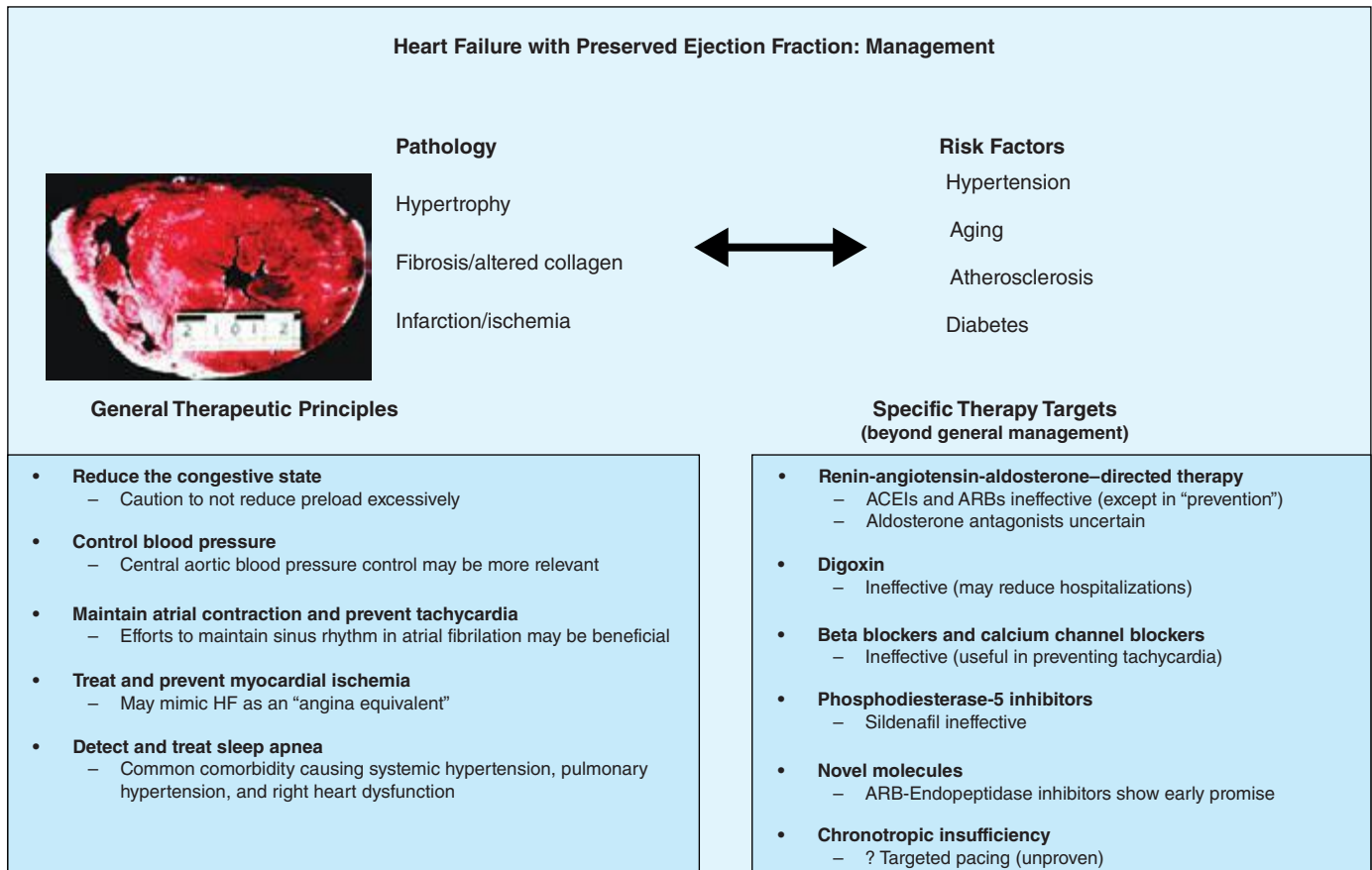


FIGURE 280-1 Pathophysiologic correlations, general therapeutic principles, and results of specific “directed” therapy in heart failure (HF) with preserved ejection fraction. ACEI, angiotensin-converting enzyme inhibitor; ARB, angiotensin receptor blocker.

parameters associated with worse outcomes: a blood urea nitrogen level greater than 43 mg/dL (to convert to mmol/L, multiply by 0.357), systolic blood pressure less than 115 mmHg, a serum creatinine level greater than 2.75 mg/dL (to convert to $\mu\text{mol/L}$, multiply by 88.4), and an elevated troponin I level. A useful clinical schema to identify treatment targets for the various phenotypic presentations and management goals in ADHF is depicted in Fig. 280-2.

VOLUME MANAGEMENT

Intravenous Diuretic Agents Intravenous diuretic agents rapidly and effectively relieve symptoms of congestion and are essential when oral drug absorption is impaired. When high doses of diuretic agents are required or when the effect is suboptimal, a continuous infusion may be needed to reduce toxicity and maintain stable serum drug levels. Randomized clinical trials of high-versus low-dose or bolus versus continuous infusion diuresis have not provided clear justification for the best diuretic strategy in ADHF, and as such, the use of diuretic regimens remains an art rather than science. Addition of a thiazide diuretic agent such as metolazone in combination provides a synergistic effect and is often required in patients receiving long-term therapy with loop diuretic agents. Change in weight is often used as a surrogate for adequate diuresis, but this objective measure of volume status may be surprisingly difficult to interpret, and weight loss during hospitalization does not necessarily correlate closely with outcomes. It is generally advisable to continue diuresis until euvolemia has been achieved. Physical examination findings, specifically the jugular venous pressure coupled with biomarker trends, are useful in timing discharge planning.

The Cardiorenal Syndrome The cardiorenal syndrome is being recognized increasingly as a complication of ADHF. Multiple definitions have been proposed for the cardiorenal syndrome, but at its simplest, it can be thought to reflect the interplay between abnormalities of heart and kidney function, with deteriorating function of one organ

while therapy is administered to preserve the other. Approximately 30% of patients hospitalized with ADHF exhibit abnormal renal function at baseline, and this is associated with longer hospitalizations and increased mortality. However, mechanistic studies have been largely unable to find correlation between deterioration in renal function, cardiac output, left-sided filling pressures, and reduced renal perfusion; most patients with cardiorenal syndrome demonstrate a preserved cardiac output. It is hypothesized that in patients with established heart failure, this syndrome represents a complex interplay of neurohormonal factors, potentially exacerbated by “backward failure” resulting from increased intra-abdominal pressure and impairment in return of renal venous blood flow. Continued use of diuretic therapy may be associated with a reduction in glomerular filtration rate and a worsening of the cardiorenal syndrome when right-sided filling pressures remain elevated. In patients in the late stages of disease characterized by profound low cardiac output state, inotropic therapy or mechanical circulatory support has been shown to preserve or improve renal function in selected individuals in the short term until more definitive therapy such as assisted circulation or cardiac transplantation is implemented.

Ultrafiltration Ultrafiltration (UF) is an invasive fluid removal technique that may supplement the need for diuretic therapy. Proposed benefits of UF include controlled rates of fluid removal, neutral effects on serum electrolytes, and decreased neurohormonal activity. This technique has also been referred to as aquapheresis in recognition of its electrolyte depletion-sparing effects. Current UF systems function with two large-bore, peripherally inserted venous lines. In a pivotal study evaluating UF versus conventional therapy, fluid removal was improved and subsequent heart failure hospitalizations and urgent clinic visits were reduced with UF; however, no improvement in renal function and no subjective differences in dyspnea scores or adverse outcomes were noted. More recently, in the Cardiorenal Rescue Study in Acute Decompensated Heart Failure (CARRESS-HF) trial,

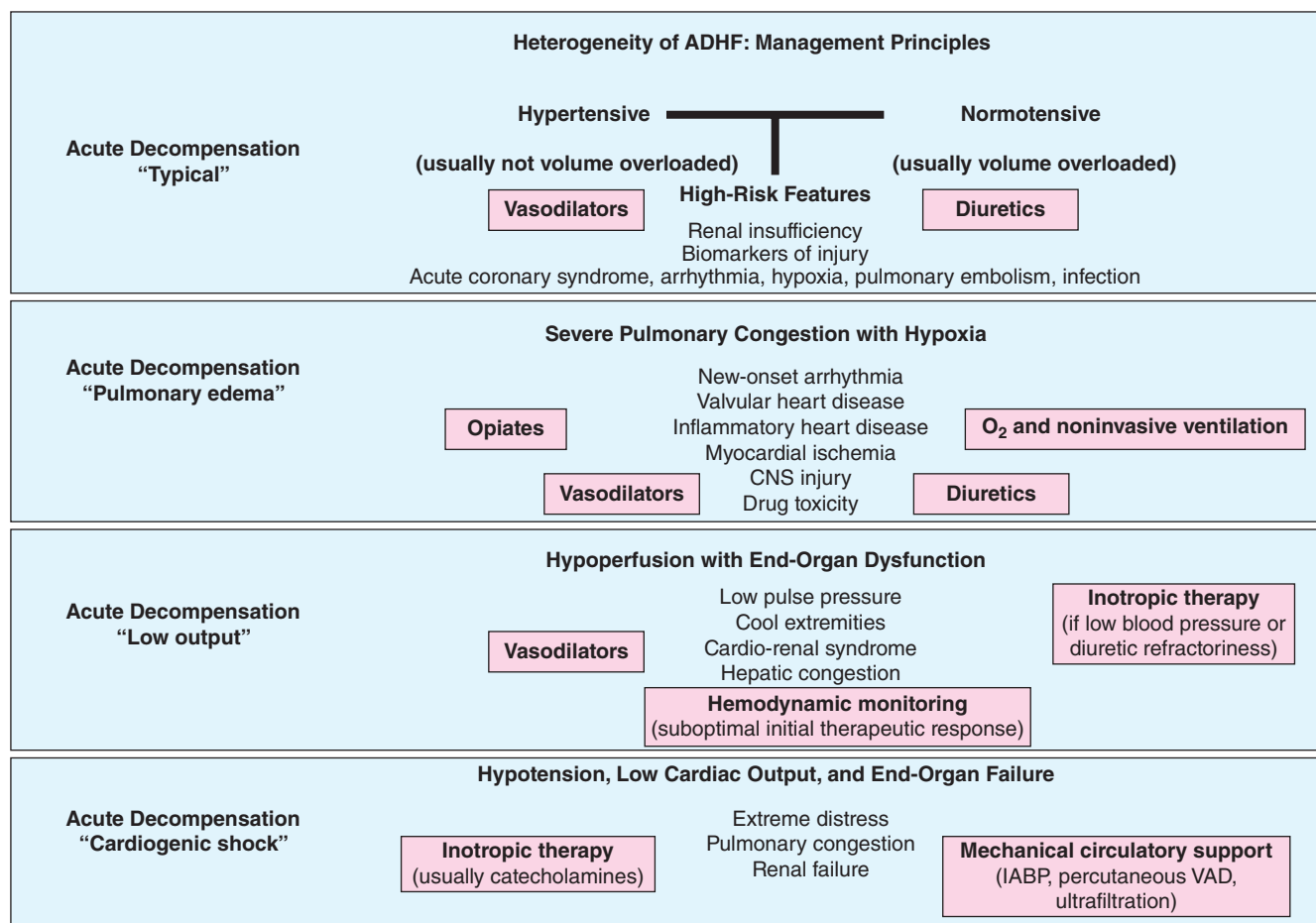


FIGURE 280-2 The distinctive phenotypes of acute decompensated heart failure (ADHF), their presentations, and suggested therapeutic routes. (Unique causes of ADHF, such as isolated right heart failure and pericardial disease, and rare causes, such as aortic and coronary dissection or ruptured valve structures or sinuses of Valsalva, are not delineated and are covered elsewhere.) IABP, intraaortic balloon pump; VAD, ventricular assist device.

188 patients with ADHF and worsening renal failure were randomized to stepped pharmacologic care or UF. The primary endpoint was a change in serum creatinine and change in weight (reflecting fluid removal) at 96 hours. Although similar weight loss occurred in both groups (approximately 5.5 kg), there was worsening in creatinine in the UF group. Deaths and hospitalizations for heart failure were no different between groups, but there were more severe adverse events in the UF group, mainly due to kidney failure, bleeding complications, and intravenous catheter-related complications. This investigation argues against using UF as a primary strategy in patients with ADHF who are nonetheless responsive to diuretics. Whether UF is useful in states of diuretic unresponsiveness remains an open question, and this strategy continues to be employed judiciously in such situations.

VASCULAR THERAPY

Vasodilators including *intravenous nitrates*, *nitroprusside*, and *nesiritide* (a recombinant brain-type natriuretic peptide) have been advocated for upstream therapy in an effort to stabilize ADHF. The latter agent was introduced in a fixed dose for therapy after a comparison with intravenous nitrates suggested more rapid and greater reduction in pulmonary capillary wedge pressure. Enthusiasm for nesiritide waned due to concerns within the pivotal trials for development of renal insufficiency and an increase in mortality. To address these concerns, a large-scale morbidity and mortality trial, the Acute Study of Clinical Effectiveness of Nesiritide in Decompensated Heart Failure (ASCEND-HF) study was completed in 2011 and randomly enrolled 7141 patients with ADHF to nesiritide or placebo for 24 to 168 hours in addition to standard care. Nesiritide was not associated with an increase or a decrease in the rates of death and rehospitalization and had a clinically insignificant benefit on dyspnea. Renal function did

not worsen, but increased rates of hypotension were noted. Although this trial established the safety for this drug, the routine use cannot be advocated due to lack of significant efficacy. *Recombinant human relaxin-2*, or *serelaxin*, is a peptide upregulated in pregnancy and examined in ADHF patients with a normal or elevated blood pressure. In the *Relaxin in Acute Heart Failure (RELAX-AHF)* trial, serelaxin or placebo was added to a regimen of standard therapy in 1161 patients hospitalized with ADHF, evidence of congestion, and systolic pressure >125 mmHg. Serelaxin improved dyspnea, reduced signs and symptoms of congestion, and was associated with less early worsening of HF. Exploratory endpoints of hard outcomes at 6 months suggested positive signals in favor of mortality reduction. This agent is being tested in a large, more confirmatory trial setting.

INOTROPIC THERAPY

Impairment of myocardial contractility often accompanies ADHF, and pharmacologic agents that increase intracellular concentration of cyclic adenosine monophosphate via direct or indirect pathways, such as sympathomimetic amines (dobutamine) and phosphodiesterase-3 inhibitors (milrinone), respectively, serve as positive inotropic agents. Their activity leads to an increase in cytoplasmic calcium. Inotropic therapy in those with a low-output state augments cardiac output, improves perfusion, and relieves congestion acutely. Although milrinone and dobutamine have similar hemodynamic profiles, milrinone is slower acting and is renally excreted and thus requires dose adjustments in the setting of kidney dysfunction. Since milrinone acts downstream from the β_1 -adrenergic receptor, it may provide an advantage in patients receiving beta blockers when admitted to the hospital. Studies are in universal agreement that long-term inotropic therapy increases mortality. However, the short-term use of inotropic agents in ADHF

1510 is also associated with increased arrhythmia, hypotension, and no beneficial effects on hard outcomes. Inotropic agents are currently indicated as bridge therapy (to either left ventricular assist device support or to transplant) or as selectively applied palliation in end-stage heart failure.

Novel inotropic agents that leverage the concept of myofilament calcium sensitization rather than increasing intracellular calcium levels have been introduced. *Levosimendan* is a calcium sensitizer that provides inotropic activity, but also possesses phosphodiesterase-3 inhibition properties that are vasodilators in action. This makes the drug unsuitable in states of low output in the setting of hypotension. Two trials, the second Randomized Multicenter Evaluation of Intravenous Levosimendan Efficacy (REVIVE II) and Survival of Patients with Acute Heart Failure in Need of Intravenous Inotropic Support (SURVIVE), have tested this agent in ADHF. SURVIVE compared levosimendan with dobutamine, and despite an initial reduction in circulating B-type natriuretic peptide levels in the levosimendan group compared with patients in the dobutamine group, this drug did not reduce all-cause mortality at 180 days or affect any secondary clinical outcomes. The second trial compared levosimendan against traditional noninotropic therapy and found a modest improvement in symptoms with worsened short-term mortality and ventricular arrhythmias. Another drug that functions as a selective myosin activator, *omecamtiv mecarbil*, prolongs the ejection period and increases fractional shortening. Distinctively, the force of contraction is not increased, and as such, this agent does not increase myocardial oxygen demand. In a 600-patient trial called ATOMIC-HF

(A Trial of Omecamtiv Mecarbil to Increase Contractility in Acute Heart Failure), this agent showed improvement in dyspnea scores in the highest dose cohort, but not across all enrolled patients. How this agent performs in broader populations remains uncertain. Other inotropic agents that increase myocardial calcium sensitivity through mechanisms that reduce cTnI phosphorylation or inhibit protein kinase A are being developed. (Table 280-1 depicts typical inotropic, vasodilator, and diuretic drugs used in ADHF.)

NEUROHORMONAL ANTAGONISTS

Other trials testing unique agents have yielded disappointing results in the situation of ADHF. The Placebo-Controlled Randomized Study of the Selective A1 Adenosine Receptor Antagonist Rolofylline for Patients Hospitalized with Acute Decompensated Heart Failure and Volume Overload to Assess Treatment Effect on Congestion and Renal Function (PROTECT) trial of selective adenosine antagonism and the Efficacy of Vasopressin Antagonism in Heart Failure Outcome Study with Tolvaptan (EVEREST) trial of an oral selective vasopressin-2 antagonist in ADHF were both negative with respect to hard outcomes.

In patients who fail to respond adequately to medical therapy, mechanical assist devices may be required. This is covered in more detail in Chap. 281.

HEART FAILURE WITH REDUCED EJECTION FRACTION

The past 50 years have witnessed great strides in the management of HFrEF. The treatment of symptomatic heart failure that evolved from a renocentric (diuretics) and hemodynamic therapy model

TABLE 280-1 INTRAVENOUS THERAPY IN ACUTE DECOMPENSATED HEART FAILURE

Drug Class	Generic Drug	Usual Dosing	Special Caution	Comments
Inotropic therapy	Dobutamine	2–20 µg/kg per min	Increased myocardial oxygen demand, arrhythmia	Use in hypotension, end-organ hypoperfusion, or shock states Short acting, an advantage; variable efficacy in presence of beta blockers (requires higher doses); clinical tolerance to prolonged infusions; concerns with hypersensitivity carditis (rare)
	Milrinone	0.375–0.75 µg/kg per min	Hypotension, arrhythmia	Decrease dose in renal insufficiency; avoid initial bolus; effectiveness retained in presence of beta blockers
	Levosimendan	0.1 µg/kg per min, range, 0.05–0.2 µg/kg per min	Hypotension, arrhythmia	Long acting; should not be used in presence of low blood pressure; similar effectiveness as dobutamine but effectiveness retained in presence of beta blockers
	Omecamtiv Mecarbil	N/A	*In trials	Increases contractility without increasing myocardial oxygen demand; in confirmatory trials
Vasodilators				Use in presence of pulmonary congestion for rapid relief of dyspnea, in presence of a preserved blood pressure
	Nitroglycerine	10–20 µg/min, increase up to 200 µg/min	Headache, flushing, tolerance	Most common vasodilator but often underdosed; effective in higher doses
	Nesiritide	Bolus 2 µg/kg and infusion at 0.01 µg/kg per min	Hypotension	Decrease in blood pressure may reduce renal perfusion pressure; bolus may be avoided since it increases hypotension predilection
	Nitroprusside	0.3 µg/kg per min titrated to 5 µg/kg per min	Thiocyanate toxicity in renal insufficiency (>72 hours)	Requires arterial line placement for titration for precise blood pressure management and prevention of hypotension
	Serelaxin	N/A (tested at 30 µg/kg per d)	Baseline blood pressure should be >125 mmHg	Not widely commercially available; undergoing confirmatory trials
Diuretics				First line of therapy in volume overload with congestion; may use bolus or continuous dosing; initial low dose (1× home dose) or high dose (2.5 × home dose) equally effective with higher risk of renal worsening with higher dose
	Furosemide	20–240 mg daily	Monitor for electrolyte loss	In severe congestion, use intravenously and consider continuous infusion (not trial supported)
	Torsemide	10–100 mg daily	Monitor for electrolyte loss	High bioavailability, can be given orally; anecdotally more effective in advanced heart failure states if furosemide less bioavailable (due to gut congestion)
	Bumetanide	0.5–5 mg daily	Monitor for electrolyte loss	Can be used orally; intermediate bioavailability
	Adjuvant diuretics for augmentation	n/a	Metolazone, chlorthalidone, spironolactone, acetazolamide	Acetazolamide is useful in presence of alkalosis; metolazone given in 2.5- to 10-mg doses; causes severe electrolyte imbalance; spironolactone is useful in presence of severe hypokalemia and normal renal function

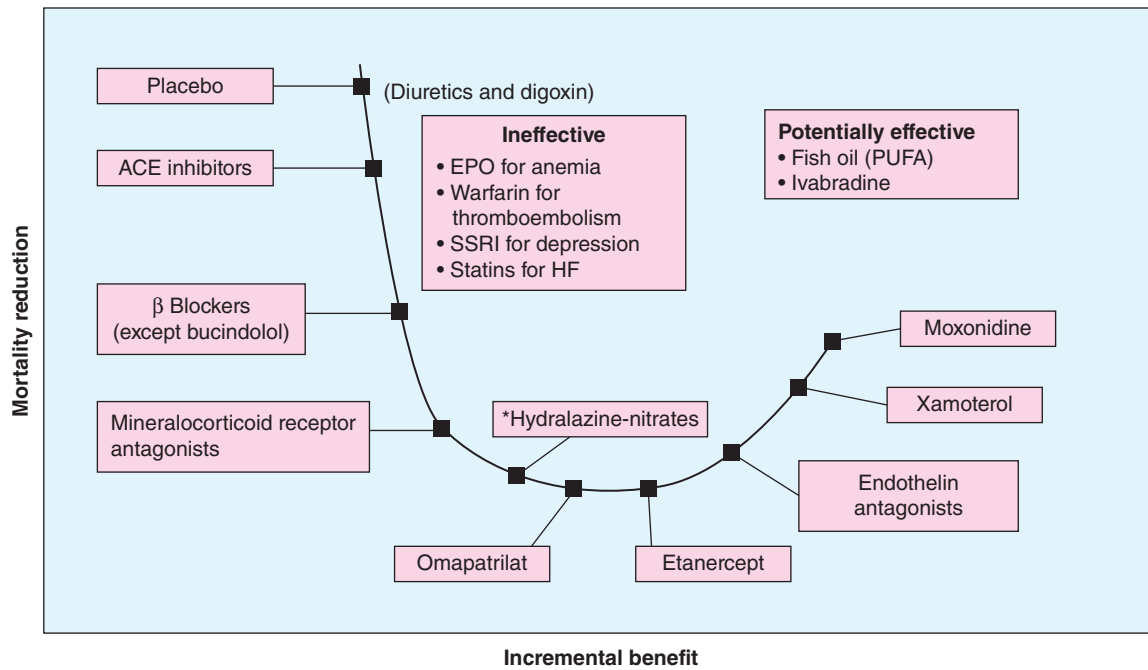


FIGURE 280-3 Progressive decline in mortality with angiotensin-converting enzyme (ACE) inhibitors or angiotensin receptor blockers (ARBs), beta blockers, mineralocorticoid receptor antagonists, and balanced vasodilators (*selected populations such as African Americans); further stack-on neurohormonal therapy is ineffective or results in worse outcome; management of comorbidity is of unclear efficacy. EPO, erythropoietin; HF, heart failure; HFrEF, heart failure with reduced ejection fraction; PUFA, polyunsaturated fatty acid; SSRI, selective serotonin reuptake inhibitor.

(digoxin, inotropic therapy) ushered in the era of disease-modifying therapy with neurohormonal antagonism. In this regard, ACEIs and beta blockers form the cornerstone of pharmacotherapy and lead to attenuation of decline and improvement in cardiac structure and function with consequent reduction in symptoms, improvement in quality of life, decreased burden of hospitalizations, and a decline in mortality from both pump failure and arrhythmic deaths (Fig. 280-3).

NEUROHORMONAL ANTAGONISM

Meta-analyses suggest a 23% reduction in mortality and a 35% reduction in the combination endpoint of mortality and hospitalizations for heart failure in patients treated with ACEIs. Patients treated with beta blockers provide a further 35% reduction in mortality on top of the benefit provided by ACEIs alone. Increased experience with both agents in a broad range of patients with HFrEF has demonstrated the safety of ACEIs in treating patients with mild renal insufficiency and the tolerability of beta blockers in patients with moderately controlled diabetes, asthma, and obstructive lung disease. The benefits of ACEIs and beta blockers extend to advanced symptoms of disease (NYHA class IIIb–IV). However, a substantial number of patients with advanced heart failure may not be able to achieve optimal doses of neurohormonal inhibitors and require cautious reduction in dose exposure to maintain clinical stability. Such individuals with lower exposure to ACEIs and beta blockers represent a high-risk cohort with poor prognosis.

Class Effect and Sequence of Administration ACEIs exert their beneficial effects in HFrEF as a class; however, the beneficial effects of beta blockers are thought to be limited to specific drugs. Beta blockers with intrinsic sympathomimetic activity (xamoterol) and other agents, including bucindolol, have not demonstrated a survival benefit. On the basis of investigations, beta blocker use in HFrEF should be restricted to carvedilol, bisoprolol, and metoprolol succinate—agents tested and proven to improve survival in clinical trials. Whether beta blockers or ACEIs should be started first was answered by the Cardiac Insufficiency Bisoprolol Study (CIBIS) III, in which outcomes did not vary when either agent was initiated first. Thus, it matters little which agent is initiated first; what does matter is that optimally titrated doses of both ACEIs and beta blockers be established in a timely manner.

Dose and Outcome A trial has indicated that higher tolerated doses of ACEIs achieve greater reduction in hospitalizations without materially improving survival. Beta blockers demonstrate a dose-dependent improvement in cardiac function and reductions in mortality and hospitalizations. Clinical experience suggests that, in the absence of symptoms to suggest hypotension (fatigue and dizziness), pharmacotherapy may be up-titrated every 2 weeks in hemodynamically stable and euvolemic ambulatory patients as tolerated.

MINERALOCORTICOID ANTAGONISTS

Aldosterone antagonism is associated with a reduction in mortality in all stages of symptomatic NYHA class II to IV HFrEF. Elevated aldosterone levels in HFrEF promote sodium retention, electrolyte imbalance, and endothelial dysfunction and may directly contribute to myocardial fibrosis. The selective agent eplerenone (tested in NYHA class II and post–myocardial infarction heart failure) and the nonselective antagonist spironolactone (tested in NYHA class III and IV heart failure) reduce mortality and hospitalizations, with significant reductions in sudden cardiac death (SCD). Hyperkalemia and worsening renal function are concerns, especially in patients with underlying chronic kidney disease, and renal function and serum potassium levels must be closely monitored.

RAAS THERAPY AND NEUROHORMONAL “ESCAPE”

Neurohormonal “escape” has been witnessed in patients with HFrEF by the finding that circulating levels of angiotensin II return to pretreatment levels with long-term ACEI therapy. ARBs blunt this phenomenon by binding competitively to the AT₁ receptor. A large meta-analysis of 24 randomized trials showed the superiority of ARBs to placebo in patients with intolerable adverse effects with ACEIs and their noninferiority in all-cause mortality or hospitalizations when compared with ACEIs. The Valsartan Heart Failure Trial (Val-HeFT) suggested that addition of valsartan in patients already receiving treatment with ACEIs and beta blockers was associated with a trend toward worse outcomes. Similarly, adding valsartan to captopril in patients with heart failure after myocardial infarction who were receiving background beta blocker therapy was associated with an increase in adverse events without any added benefit compared with

1512 monotherapy for either group. Thus, the initial clinical strategy should be to use a two-drug combination first (ACEI and beta blocker; if beta blocker intolerant, then ACEI and ARB; if ACEI intolerant, then ARB and beta blocker). In symptomatic patients (NYHA class II–IV), an aldosterone antagonist should be strongly considered, but four-drug therapy should be avoided.

A recent trial called the Aliskiren Trial on Acute Heart Failure Outcomes (ASTRONAUT) tested a direct renin inhibitor, aliskiren, in addition to other heart failure medications, within a week after discharge from a hospitalization for decompensated HFrEF. No significant difference in cardiovascular death or hospitalization at 6 or 12 months was noted. Aliskiren was associated with a reduction in circulating natriuretic peptides, but any disease-modifying effect was overcome by excessive adverse events including hyperkalemia, hypotension, and renal dysfunction.

ARTERIOVENOUS VASODILATION

The combination of hydralazine and nitrates has been demonstrated to improve survival in HFrEF. Hydralazine reduces systemic vascular resistance and induces arterial vasodilatation by affecting intracellular calcium kinetics; nitrates are transformed in smooth muscle cells into nitric oxide, which stimulates cyclic guanosine monophosphate production and consequent arterial-venous vasodilation. This combination improves survival, but not to the magnitude evidenced by ACEIs or ARBs. However, in individuals with HFrEF unable to tolerate renin-angiotensin-aldosterone–based therapy for reasons such as renal insufficiency or hyperkalemia, this combination is preferred as a disease-modifying approach. A trial conducted in self-identified African Americans, the African-American Heart Failure Trial (A-Heft), studied a fixed dose of isosorbide dinitrate with hydralazine in patients with advanced symptoms of HFrEF who were receiving standard background therapy. The study demonstrated benefit in survival and hospitalization recidivism in the treatment group. Adherence to this regimen is limited by the thrice-daily dosing schedule. **Table 280-2** lists the common neurohormonal and vasodilator regimens for HFrEF.

HEART RATE MODIFICATION

Ivabradine, an inhibitor of the I_f current in the sinoatrial node, slows the heart rate without a negative inotropic effect. The Systolic Heart Failure Treatment with Ivabradine Compared with Placebo Trial

(SHIFT) was conducted in patients with class II or III HFrEF, a heart rate >70 beats/min, and history of hospitalization for heart failure during the previous year. Ivabradine reduced hospitalizations and the combined endpoint of cardiovascular-related death and heart failure hospitalization. The study population was not necessarily representative of North American patients with HFrEF since, with a few exceptions, most did not receive internal cardioverter-defibrillation or cardiac resynchronization therapy and 40% did not receive a mineralocorticoid receptor antagonist. Although 90% received beta blockers, only a quarter were on full doses. Whether this agent, now available outside the United States, would have been effective in patients receiving robust, guideline-recommended therapy for heart failure remains enigmatic. In the 2012 European Society of Cardiology guidelines for the treatment of heart failure, ivabradine was suggested as second-line therapy before digoxin in considered in patients who remain symptomatic after guideline-based ACEIs, beta blockers, and mineralocorticoid receptor antagonists and with residual heart rate >70 beats/min. Another group in whom potential benefit may be expected includes those unable to tolerate beta blockers.

DIGOXIN

Digitalis glycosides exert a mild inotropic effect, attenuate carotid sinus baroreceptor activity, and are sympathoinhibitory. These effects decrease serum norepinephrine levels, plasma renin levels, and possibly aldosterone levels. The DIG trial demonstrated a reduction in heart failure hospitalizations in the treatment group but no reduction in mortality or improvement in quality of life. Importantly, treatment with digoxin resulted in a higher mortality rate in women than men. Furthermore, the effects of digoxin in reducing hospitalizations were lower in women than in men. It should be noted that low doses of digoxin are sufficient to achieve any potentially beneficial outcomes, and higher doses breach the therapeutic safety index. Although digoxin levels should be checked to minimize toxicity and although dose reductions are indicated for higher levels, no adjustment is made for low levels. Generally, digoxin is now relegated as therapy for patients who remain profoundly symptomatic despite optimal neurohormonal blockade and adequate volume control.

ORAL DIURETICS

Neurohormonal activation results in avid salt and water retention. Loop diuretic agents are often required because of their increased potency,

TABLE 280-2 PHARMACOLOGIC THERAPY AND TARGET DOSES IN HEART FAILURE WITH REDUCED EJECTION FRACTION

Drug Class	Generic Drug	Mean Daily Dose in Clinical Trials (mg)	Initiation (mg)	Target Dose (mg)
Angiotensin-Converting Enzyme Inhibitors				
	Lisinopril	4.5–33	2.5–5 qd	20–35 qd
	Enalapril	17	2.5 bid	10–20 bid
	Captopril	123	6.25 tid	50 tid
	Trandolapril	N/A	0.5–1 qd	4 qd
Angiotensin Receptor Blockers				
	Losartan	129	50 qd	150 qd
	Valsartan	254	40 bid	160 bid
	Candesartan	24	4–8 qd	32 qd
Aldosterone Antagonists				
	Eplerenone	42.6	25 qd	50 qd
	Spironolactone	26	12.5–25 qd	25–50 qd
Beta Blockers				
	Metoprolol succinate CR/XL	159	12.5–25 qd	200 qd
	Carvedilol	37	3.125 bid	25–50 bid
	Bisoprolol	8.6	1.25 qd	10 qd
Arteriovenous Vasodilators				
	Hydralazine isosorbide dinitrate	270/136	37.5/20 tid	75/40 tid
	Fixed-dose hydralazine/isosorbide dinitrate	143/76	37.5/20 qid	75/40 qid

and frequent dose adjustments may be necessary because of variable oral absorption and fluctuations in renal function. Importantly, clinical trial data confirming efficacy are limited, and no data suggest that these agents improve survival. Thus, diuretic agents should ideally be used in tailored dosing schedules to avoid excessive exposure. Indeed, diuretics are essential at the outset to achieve volume control before neurohormonal therapy is likely to be well tolerated or titrated.

CALCIUM CHANNEL ANTAGONISTS

Amlodipine and felodipine, second-generation calcium channel-blocking agents, safely and effectively reduce blood pressure in HFrEF but do not affect morbidity, mortality, or quality of life. The first-generation agents, including verapamil and diltiazem, may exert negative inotropic effects and destabilize previously asymptomatic patients. Their use should be discouraged.

NOVEL NEUROHORMONAL ANTAGONISM

Despite an abundance of animal and clinical data demonstrating deleterious effects of activated neurohormonal pathways beyond the RAAS and sympathetic nervous system, targeting such pathways with incremental blockade has been largely unsuccessful. As an example, the endothelin antagonist bosentan is associated with worsening heart failure in HFrEF despite demonstrating benefits in right-sided heart failure due to pulmonary arterial hypertension. Similarly, the centrally acting sympatholytic agent moxonidine worsens outcomes in left heart failure. The combined drug omapatrilat hybridizes an ACEI with a neutral endopeptidase inhibitor, and this agent was tested in the Omapatrilat Versus Enalapril Randomized Trial of Utility in Reducing Events (OVERTURE) trial. This drug did not favorably influence the primary outcome measure of the combined risk of death or hospitalization for heart failure requiring intravenous treatment. The risk of angioedema was notably higher with omapatrilat than ACEIs alone. LCZ696 and ARB with an endopeptidase inhibitor have shown benefit in a large trial versus ARB alone.

INFLAMMATION

Targeting inflammatory cytokines such as tumor necrosis factor α (TNF- α) by using anticytokine agents such as infliximab and etanercept has been unsuccessful and associated with worsening heart failure. Nonspecific immunomodulation has been tested in the large Advanced Chronic Heart Failure Clinical Assessment of Immune Modulation Therapy (ACCLAIM-HF) trial of 2426 HFrEF patients with NYHA functional class II to IV symptoms. Ex vivo exposure of a blood sample to controlled oxidative stress initiates apoptosis of leukocytes soon after intramuscular gluteal injection of the treated sample. The physiologic response to apoptotic cells results in a reduction in inflammatory cytokine production and upregulation of anti-inflammatory cytokines. This promising hypothesis was not proven, although certain subgroups (those with no history of previous myocardial infarction and those with mild heart failure) showed signals in favor of immunomodulation. Use of intravenous immunoglobulin therapy in nonischemic etiology of heart failure has not been shown to result in beneficial outcomes.

STATINS

Potent lipid-altering and pleiotropic effects of statins reduce major cardiovascular events and improve survival in non-heart failure populations. Once heart failure is well established, this therapy may not be as beneficial and theoretically could even be detrimental by depleting ubiquinone in the electron transport chain. Two trials, Controlled Rosuvastatin Multinational Trial in Heart Failure (CORONA) and Gruppo Italiano per lo Studio della Sopravvivenza nell'Insufficienza Cardiac (GISSI-HF), have tested low-dose rosuvastatin in patients with HFrEF and demonstrated no improvement in aggregate clinical outcomes. If statins are required to treat progressive coronary artery disease in the background setting of heart failure, then they should be employed. However, no rationale appears to exist for routine statin therapy in nonischemic heart failure.

ANTICOAGULATION AND ANTIPLATELET THERAPY

HFrEF is accompanied by a hypercoagulable state and therefore a high risk of thromboembolic events, including stroke, pulmonary embolism, and peripheral arterial embolism. Although long-term oral anticoagulation is established in certain groups, including patients with atrial fibrillation, the data are insufficient to support the use of warfarin in patients in normal sinus rhythm without a history of thromboembolic events or echocardiographic evidence of left ventricular thrombus. In the large Warfarin versus Aspirin in Reduced Cardiac Ejection Fraction (WARCEF) trial, 2305 patients with HFrEF were randomly allocated to either full-dose aspirin or international normalized ratio-controlled warfarin with follow-up for 6 years. Among patients with reduced LVEF who were in sinus rhythm, there was no significant overall difference in the primary outcome between treatment with warfarin and treatment with aspirin. A reduced risk of ischemic stroke with warfarin was offset by an increased risk of major hemorrhage. Aspirin blunts ACEI-mediated prostaglandin synthesis, but the clinical importance of this finding remains unclear. Current guidelines support the use of aspirin in patients with ischemic cardiomyopathy.

FISH OIL

Treatment with long-chain omega-3 polyunsaturated fatty acids (ω -3 PUFAs) has been shown to be associated with modestly improved clinical outcomes in patients with HFrEF. This observation from the GISSI-HF trial was extended to measurements of ω -3 PUFAs in plasma phospholipids at baseline and after 3 months. Three-month treatment with ω -3 PUFAs enriched circulating eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA). Low EPA levels are inversely related to total mortality in patients with HFrEF.

MICRONUTRIENTS

A growing body of evidence suggests an association between heart failure and micronutrient status. Reversible heart failure has been described as a consequence of severe thiamine and selenium deficiency. Thiamine deficiency has received attention in heart failure due to the fact that malnutrition and diuretics are prime risk factors for thiamine loss. Small exploratory randomized studies have suggested a benefit of supplementation of thiamine in HFrEF with evidence of improved cardiac function. This finding is restricted to chronic heart failure states and does not appear to be beneficial in the ADHF phenotype. Due to the preliminary nature of the evidence, no recommendations for routine supplementation or testing for thiamine deficiency can be made.

ENHANCED EXTERNAL COUNTERPULSATION (EECP)

Peripheral lower extremity therapy using graded external pneumatic compression at high pressure is administered in 1-hour sessions for 35 treatments (7 weeks) and has been proposed to reduce angina symptoms and extend time to exercise-induced ischemia in patients with coronary artery disease. The Prospective Evaluation of Enhanced External Counterpulsation in Congestive Heart Failure (PEECH) study assessed the benefits of enhanced external counterpulsation in the treatment of patients with mild-to-moderate heart failure. This randomized trial improved exercise tolerance, quality of life, and NYHA functional classification but without an accompanying increase in peak oxygen consumption. A placebo effect due to the nature of the intervention simply cannot be excluded.

EXERCISE

The Heart Failure: A Controlled Trial Investigating Outcomes of Exercise Training (HF-ACTION) study investigated short-term (3-month) and long-term (12-month) effects of a supervised exercise training program in patients with moderate HFrEF. Exercise was safe, improved patients' sense of well-being, and correlated with a trend toward mortality reduction. Maximal changes in 6-minute walk distance were evident at 3 months with significant improvements in cardiopulmonary exercise time and peak oxygen consumption persisting at 12 months. Therefore, exercise training is recommended as an adjunctive treatment in patients with heart failure.

Sleep-disordered breathing is common in HF and particularly in HFrEF. A range of presentations exemplified by obstructive sleep apnea, central sleep apnea, and its extreme form of Cheyne-Stokes breathing are noted. Frequent periods of hypoxia and repeated micro- and macro-arousals trigger adrenergic surges, which can worsen hypertension and impair systolic and diastolic function. A high index of suspicion is required, especially in patients with difficult-to-control hypertension or with predominant symptoms of fatigue despite reverse remodeling in response to optimal medical therapy. Worsening of right heart function with improvement of left ventricular function noted on medical therapy should immediately trigger a search for underlying sleep-disordered breathing or pulmonary complications such as occult embolism or pulmonary hypertension. Treatment with nocturnal positive airway pressure improves oxygenation, LVEF, and 6-minute walk distance. However, no conclusive data exist to support this therapy as a disease-modifying approach with reduction in mortality.

Anemia is common in heart failure patients, reduces functional status and quality of life, and is associated with increased proclivity for hospital admissions and mortality. Anemia in heart failure is more common in the elderly, in those with advanced stages of HFrEF, in the presence of renal insufficiency, and in women and African Americans. The mechanisms include iron deficiency, dysregulation of iron metabolism, and occult gastrointestinal bleeding. Intravenous iron using either iron sucrose or carboxymaltose (Ferric Carboxymaltose Assessment in Patients with Iron Deficiency and Chronic Heart Failure [FAIR-HF] trial) has been shown to correct anemia and improve functional capacity. Erythropoiesis-regulating agents such as erythropoietin analogues have been studied with disappointing results. The Reduction of Events by Darbepoetin Alfa in Heart Failure (RED-HF) trial evaluated 2278 mild-to-moderate anemia patients with HFrEF and demonstrated that treatment with darbepoetin alfa did not improve clinical outcomes in patients with systolic heart failure.

Depression is common in HFrEF, with a reported prevalence of one in five patients, and is associated with a poor quality of life, limited functional status, and increased risk of morbidity and mortality in this population. Antidepressants may improve depression, promote vascular health, and decrease systemic inflammation in HFrEF. However, the largest randomized study of depression in HFrEF, the Sertraline Against Depression and Heart Disease in Chronic Heart Failure (SADHART-CHF) trial, showed that sertraline was safe, but did not provide greater reduction in depression or improve cardiovascular status among patients with heart failure and depression compared with nurse-driven multidisciplinary management.

Atrial arrhythmias, especially atrial fibrillation, are common and serve as a harbinger of worse prognosis in patients with heart failure. When rate control is inadequate or symptoms persist, pursuing a rhythm control strategy is reasonable. Rhythm control may be achieved via pharmacotherapy or by percutaneous or surgical techniques, and referral to practitioners or centers experienced in these modalities is recommended. Antiarrhythmic drug therapy should be restricted to amiodarone and dofetilide, both of which have been shown to be safe and effective but do not alter the natural history of the underlying disease. The Antiarrhythmic Trial with Dronedrone in Moderate-to-Severe Congestive Heart Failure Evaluating Morbidity Decrease (ANDROMEDA) studied the effects of the novel antiarrhythmic agent dronedarone and found an increased mortality due to worsening heart failure. Catheter ablation and pulmonary vein isolation appear to be safe and effective in this high-risk cohort and compare favorably with the more established practice of atrioventricular node ablation and biventricular pacing.

CARDIAC RESYNCHRONIZATION THERAPY

Nonsynchronous contraction between the walls of the left ventricle (intraventricular) or between the ventricular chambers (interventricular) impairs systolic function, decreases mechanical efficiency of contraction, and adversely affects ventricular filling. Mechanical

dyssynchrony results in an increase in wall stress and worsens functional mitral regurgitation. The single most important association of extent of dyssynchrony is a widened QRS interval on the surface electrocardiogram, particularly in the presence of a left bundle branch block pattern. With placement of a pacing lead via the coronary sinus to the lateral wall of the ventricle, cardiac resynchronization therapy (CRT) enables a more synchronous ventricular contraction by aligning the timing of activation of the opposing walls. Early studies showed improved exercise capacity, reduction in symptoms, and evidence of reverse remodeling. The Cardiac Resynchronization in Heart Failure Study (CARE-HF) trial was the first study to demonstrate a reduction in all-cause mortality with CRT placement in patients with HFrEF on optimal therapy with continued moderate-to-severe residual symptoms of NYHA class III or IV heart failure. More recent clinical trials have demonstrated disease-modifying properties of CRT in even minimally symptomatic patients with HFrEF, including the Resynchronization–Defibrillation for Ambulatory Heart Failure Trial (RAFT) and Multicenter Automatic Defibrillator Implantation Trial with Cardiac Resynchronization Therapy (MADIT-CRT), both of which sought to use CRT in combination with an implantable defibrillator. Most benefit in mildly symptomatic HFrEF patients accrues from applying this therapy in those with a QRS width of >149 ms and a left bundle branch block pattern. Attempts to further optimize risk stratification and expand indications for CRT using modalities other than electrocardiography have proven disappointing. In particular, echocardiographically derived measures of dyssynchrony vary tremendously, and narrow QRS dyssynchrony has not proven to be a good target for treatment. Uncertainty surrounds the benefits of CRT in those with ADHF, a predominant right bundle branch block pattern, atrial fibrillation, and evidence of scar in the lateral wall, which is the precise location where the CRT lead is positioned.

SUDDEN CARDIAC DEATH PREVENTION IN HEART FAILURE

SCD due to ventricular arrhythmias is the mode of death in approximately half of patients with heart failure and is particularly proportionally prevalent in HFrEF patients with early stages of the disease. Patients who survive an episode of SCD are considered to be at very high risk and qualify for placement of an implantable cardioverter-defibrillator (ICD). Although primary prevention is challenging, the degree of residual left ventricular dysfunction despite optimal medical therapy ($\leq 35\%$) to allow for adequate remodeling and the underlying etiology (post-myocardial infarction or ischemic cardiomyopathy) are the two single most important risk markers for stratification of need and benefit. Currently, patients with NYHA class II or III symptoms of heart failure and an LVEF $< 35\%$, irrespective of etiology of heart failure, are appropriate candidates for ICD prophylactic therapy. In patients with a myocardial infarction and optimal medical therapy with residual LVEF $\leq 30\%$ (even when asymptomatic), placement of an ICD is appropriate. In patients with a terminal illness and a predicted life span of less than 6 months or in those with NYHA class IV symptoms who are refractory to medications and who are not candidates for transplant, the risks of multiple ICD shocks must be carefully weighed against the survival benefits. If a patient meets the QRS criteria for CRT, combined CRT with ICD is often employed ([Table 280-3](#)).

SURGICAL THERAPY IN HEART FAILURE

Coronary artery bypass grafting (CABG) is considered in patients with ischemic cardiomyopathy with multivessel coronary artery disease. The recognition that hibernating myocardium, defined as myocardial tissue with abnormal function but maintained cellular function, could recover after revascularization led to the notion that revascularization with CABG would be useful in those with living myocardium. Revascularization is most robustly supported in individuals with ongoing angina and left ventricular failure. Revascularizing those with left ventricular failure in the absence of angina remains controversial. The Surgical Treatment for Ischemic Heart Failure (STICH) trial enrolled 1212 patients with an ejection fraction of 35% or less and coronary artery disease amenable to CABG and randomly assigned them to

TABLE 280-3 PRINCIPLES OF ICD IMPLANTATION FOR PRIMARY PREVENTION OF SUDDEN DEATH

Principle	Comment
Arrhythmia–sudden death mismatch	Sudden death in heart failure patients is generally due to progressive LVD, not a focal arrhythmia substrate (except in patients with post-MI HF)
Diminishing returns with advanced disease	Intervention at early stages of HF most successful since sudden death diminishes as cause of death with advanced HF
Timing of benefits	LVEF should be evaluated on optimal medical therapy or after revascularization before ICD therapy is employed; no benefit to ICD implant within 40 days of an MI (unless for secondary prevention)
Estimation of benefits and prognosis	Patients and clinicians often overestimate benefits of ICDs; an ICD discharge is not equivalent to an episode of sudden death (some ventricular arrhythmias terminate spontaneously); appropriate ICD discharges are associated with a worse near-term prognosis

Abbreviations: HF, heart failure; ICD, implantable cardioverter-defibrillator; LVD, left ventricular disease; LVEF, left ventricular ejection fraction; MI, myocardial infarction.

medical therapy alone or medical therapy plus CABG. There was no significant difference between groups with respect to the primary endpoint of death from any cause. Patients assigned to CABG had lower rates of death from cardiovascular causes and of death from any cause or hospitalization for cardiovascular causes. An ancillary study of this trial also determined that the detection of hibernation pre-revascularization did not materially influence the efficacy of this approach, nor did it help to define a population unlikely to benefit if hibernation was not detected.

Surgical ventricular restoration (SVR), a technique characterized by infarct exclusion to remodel the left ventricle by reshaping it surgically in patients with ischemic cardiomyopathy and dominant anterior left ventricular dysfunction, has been proposed. However, in a 1000-patient trial in patients with HFrEF who underwent CABG alone or CABG plus SVR, the addition of SVR to CABG had no disease-modifying effect. Cardiac symptoms and exercise tolerance improved from baseline to a similar degree in both study groups. SVR resulted in lower left ventricular volumes at 4 months after operation. However, left ventricular aneurysm surgery is still advocated in those with refractory heart failure, ventricular arrhythmias, or thromboembolism arising from an akinetic aneurysmal segment of the ventricle. Other remodeling procedures, such as use of an external mesh-like net attached around the heart to limit further enlargement, have not been shown to provide hard clinical benefits, although favorable cardiac remodeling was noted.

Mitral regurgitation (MR) occurs with varying degrees in patients with HFrEF and dilated ventricles. Annular dilatation and leaflet non-coaptation in the setting of anatomically normal papillary muscles, chordal structures, and valve leaflets characterize functional MR. In patients who are not candidates for surgical coronary revascularization, mitral valve repair remains controversial. Ischemic MR (or infarct-related MR) is typically associated with leaflet tethering and displacement related to abnormal left ventricular wall motion and geometry. No evidence to support the use of surgical or percutaneous valve correction for functional MR exists as disease-modifying therapy even though MR can be corrected.

CELLULAR AND GENE-BASED THERAPY

The cardiomyocyte is no longer considered a terminally differentiated cell and possesses regenerative capacity. Such renewal is accelerated

under conditions of stress and injury, such as an ischemic event or heart failure. Investigations that use either bone marrow–derived precursor cells or autologous cardiac-derived cells have gained traction. A number of small- and moderate-scale trials of such therapy have focused on post–myocardial infarction patients and have used autologous bone marrow–derived progenitor or stem cells. These trials have had variable results, with most demonstrating modest improvements in parameters of cardiac structure and remodeling. More promising, however, are cardiac-derived stem cells. Two preliminary pilot trials delivering cells via an intracoronary approach have been reported. In one, autologous c-kit–positive cells isolated from the atria obtained from patients undergoing CABG were cultured and reinfused. In another, cardiosphere-derived cells grown from endomyocardial biopsy specimens were used. These small trials demonstrated improvements in left ventricular function but require far more work to usher in a clinical therapeutic success. The appropriate route of administration, the quantity of cells to achieve a minimal therapeutic threshold, the constitution of these cells (single source or mixed), the mechanism by which benefit accrues, and short- and long-term safety remain to be elucidated.

Targeting molecular aberrations using gene transfer therapy, mostly with an adenoviral vector, is emerging in HFrEF. Several methods of gene delivery have been developed, including direct intramyocardial injection, coronary artery or venous infusion, and injection into the pericardial space. Cellular targets under consideration include β_2 -adrenergic receptors and calcium cycling proteins such as inhibitors of phospholamban. SERCA2a is deficient in patients with HFrEF and is primarily responsible for reincorporating calcium into the sarcoplasmic reticulum during diastole. A phase II randomized, double-blind, placebo-controlled trial called CUPID (Efficacy and Safety Study of Genetically Targeted Enzyme Replacement Therapy for Advanced Heart Failure) was completed. This study used coronary arterial infusion of adeno-associated virus type 1 carrying the gene for SERCA2a and demonstrated that natriuretic peptides were decreased, reverse remodeling was noted, and symptomatic improvements were forthcoming. Stromal-derived factor 1 enhances myocardial repair and facilitates “homing” of stem cells to the site of tissue injury. Strategies using intramyocardial injections to deploy this gene at sites of injury are being studied.

More advanced therapies for late-stage heart failure such as left ventricular assist devices and cardiac transplantation are covered in detail in Chap. 281.

DISEASE MANAGEMENT AND SUPPORTIVE CARE

Despite stellar outcomes with medical therapy, admission rates following heart failure hospitalization remain high, with nearly half of all patients readmitted to hospital within 6 months of discharge. Recurrent heart failure and related cardiovascular conditions account for only half of readmissions in patients with heart failure, whereas other comorbidity-related conditions account for the rest. The key to achieving enhanced outcomes must begin with the attention to transitional care at the index hospitalization with facilitated discharge through comprehensive discharge planning, patient and caregiver education, appropriate use of visiting nurses, and planned follow-up. Early postdischarge follow-up, whether by telephone or clinic-based, may be critical to ensuring stability because most heart failure–related readmissions tend to occur within the first 2 weeks after discharge. Although routinely advocated, intensive surveillance of weight and vital signs with use of telemonitoring has not decreased hospitalizations. Intrathoracic impedance measurements have been advocated for the identification of early rise in filling pressure and worsened hemodynamics so that preemptive management may be employed. However, this has not been successful and may worsen outcomes in the short term. Implantable pressure monitoring systems do tend to provide signals for early decompensation, and in patients with moderately advanced symptoms, such systems have been shown to provide information that can allow implementation of therapy to avoid hospitalizations by as much as 39% (in the CardioMEMS Heart Sensor Allows Monitoring of Pressure to Improve Outcomes in NYHA Class III Heart

1516 Failure Patients [CHAMPION] trial). Once heart failure becomes advanced, regularly scheduled review of the disease course and options with the patient and family is recommended including discussions surrounding end-of-life preferences when patients are comfortable in an outpatient setting. As the disease state advances further, integrating care with social workers, pharmacists, and community-based nursing may be critical in improving patient satisfaction with the therapy, enhancing quality of life, and avoiding heart failure hospitalizations. Equally important is attention to seasonal influenza vaccinations and periodic pneumococcal vaccines that may obviate non-heart failure hospitalizations in these ill patients. When nearing end of life, facilitating a shift in priorities to outpatient and hospice palliation is key, as are discussions around advanced therapeutics and continued use of ICD prophylaxis, which may worsen quality of life and prolong death.

options for such patients. Currently, both of the latter approaches are considered to be experimental.

CARDIAC TRANSPLANTATION



Surgical techniques for orthotopic transplantation of the heart were devised in the 1960s and taken into the clinical arena in 1967. The procedures did not gain widespread clinical acceptance until the introduction of “modern” and more effective immunosuppression in the early 1980s. By the 1990s, the demand for transplantable hearts met, and then exceeded, the available donor supply and leveled off at about 4000 heart transplantations annually worldwide, according to data from the Registry of the International Society for Heart and Lung Transplantation (ISHLT). Subsequently, heart transplantation activity in the United States has remained stable at ~2200 per year, but worldwide activity reported to this registry has decreased somewhat. This apparent decline in numbers may be a result of the fact that reporting is legally mandated in the United States but not elsewhere, and several countries have started their own databases.

GLOBAL CONSIDERATIONS

Substantial differences exist in the practice of heart failure therapeutics and outcomes by geographic location. International guidelines produced by the American College of Cardiology/American Heart Association, European Society of Cardiology, and National Institute for Health and Clinical Excellence (United Kingdom) differ in their approach to evaluation of evidence and prioritization of therapy. The penetrance of CRT and ICD is higher in the United States than in Europe. Conversely, therapy unavailable in the United States, such as ivabradine and levosimendan, is designated as useful in Europe. Although ACEIs appear to be similarly effective across populations, variation in the benefits of beta blockers based on world region remains an area of controversy. In oral pharmacologic therapy trials of HF_{rEF}, patients from southwest Europe have a lower incidence of ischemic cardiomyopathy and those in North America tend to have more diabetes and prior coronary revascularization. There is also regional variation in medication use even after accounting for indication. In trials of ADHF, patients in Eastern Europe tend to be younger, with higher ejection fractions and lower natriuretic peptide levels. Patients from South America tend to have the lowest rates of comorbidities, revascularization, and device use. In contrast, patients from North America have the highest comorbidity burden with high revascularization and device use rates. Given geographic differences in baseline characteristics and clinical outcomes, the generalizability of therapeutic outcomes in patients in the United States and Western Europe may require verification.

SURGICAL TECHNIQUE

Donor and recipient hearts are excised in virtually identical operations with incisions made across the atria and atrial septum at the mid-atrial level (with the posterior walls of the atria left in place) and across the great vessels just above the semilunar valves. The donor heart is generally “harvested” by a separate surgical team, transported from the donor hospital in a bag of iced saline solution, and reanastomosed into the waiting recipient in the orthotopic or normal anatomic position. The only change in surgical technique since this method was first described has been a movement in recent years to move the right atrial anastomosis back to the level of the superior and inferior venae cavae to better preserve right atrial geometry and prevent atrial arrhythmias. Both methods of implantation leave the recipient with a surgically denervated heart that does not respond to any direct sympathetic or parasympathetic stimuli but does respond to circulating catecholamines. The physiologic responses of the denervated heart to the demands of exercise are atypical but quite adequate for continuation of normal physical activity.

DONOR ALLOCATION SYSTEM

In the United States, the allocation of donor organs is accomplished under the supervision of the United Network for Organ Sharing, a private organization under contract to the federal government. The United States is divided geographically into eleven regions for donor heart allocation. Allocation of donor hearts within a region is decided according to a system of priority that takes into account (1) the severity of illness, (2) the geographic distance from the donor, and (3) the patient’s time on the waiting list. A physiologic limit of ~3 h of “ischemic” (out-of-body) time for hearts precludes a national sharing of hearts. This allocation system design is reissued annually and is responsive to input from a variety of constituencies, including both donor families and transplantation professionals.

At the current time, the highest priority according to severity of illness is assigned to patients requiring hospitalization at the transplantation center for IV inotropic support, with a pulmonary artery catheter in place for hemodynamic monitoring, or to patients requiring mechanical circulatory support—i.e., use of an intra-aortic balloon pump or a right or left ventricular assist device (RVAD, LVAD), extracorporeal membrane oxygenation, or mechanical ventilation. The second highest priority is given to patients requiring ongoing inotropic support, but without a pulmonary artery catheter in place. All other patients are assigned a priority according to time accrued on the waiting list, and matching generally is based only on compatibility in terms of ABO blood group and gross body size.

While HLA matching of donor and recipient would be ideal, the relatively small numbers of patients as well as the time constraints involved make such matching impractical. However, some patients who are “presensitized” and have preexisting antibodies to human leukocyte antigens (HLAs) undergo prospective cross-matching with the

281 Cardiac Transplantation and Prolonged Assisted Circulation

Sharon A. Hunt, Hari R. Mallidi

Advanced or end-stage heart failure is an increasingly frequent sequela of many types of heart disease, as progressively more effective palliation for the earlier stages of heart disease and prevention of sudden death associated with heart disease become more widely recognized and employed (Chap. 279). When patients with end-stage or refractory heart failure are identified, the physician is faced with the decision of advising compassionate end-of-life care or choosing to recommend extraordinary life-extending measures. For the occasional patient who is relatively young and without serious comorbidities, the latter may represent a reasonable option. Current therapeutic options are limited to cardiac transplantation (with the option of mechanical cardiac assistance as a “bridge” to transplantation) or permanent mechanical assistance of the circulation. In the future, it is possible that genetic modulation of ventricular function or cell-based cardiac repair will be

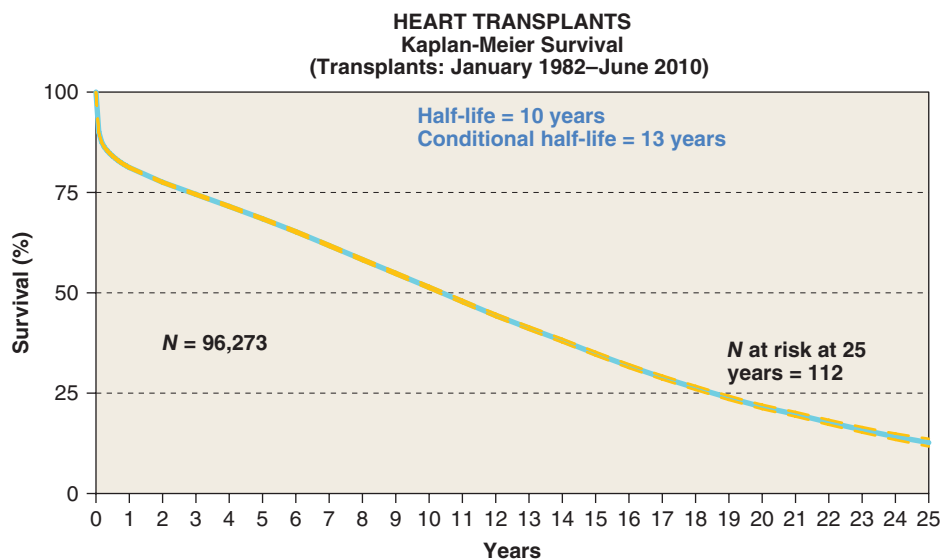


FIGURE 281-1 Global survival rates after heart transplantation since 1982. Rates were calculated by the Kaplan-Meier method, which incorporates information from all transplant recipients for whom any follow-up has been provided. Because many patients are still alive and some patients have been lost to follow-up, the survival rates are estimates rather than exact figures because the time of death is not known for all patients. Therefore, 95% confidence limits are provided. (From J Stehlik et al: *J Heart Lung Transplant* 31:1052, 2012.)

donor; these patients are commonly multiparous women or patients who have received multiple transfusions.

INDICATIONS/CONTRAINDICATIONS

Heart failure is an increasingly common cause of death, particularly in the elderly. Most patients who reach what has recently been categorized as stage D, or refractory end-stage heart failure, are appropriately treated with compassionate end-of-life care. A subset of such patients who are younger and without significant comorbidities can be considered as candidates for heart transplantation. Exact criteria vary in different centers but generally take into consideration the patient's physiologic age and the existence of comorbidities such as peripheral or cerebrovascular disease, obesity, diabetes, cancer, or chronic infection.

RESULTS



A registry organized by the ISHLT has tracked worldwide and U.S. survival rates after heart transplantation since 1982. The most recent update reveals survival rates of 83% and 76% 1 and 3 years after transplantation, respectively, or a posttransplantation “half-life” of 10.00 years (Fig. 281-1). The quality of life of survivors is generally excellent, with well over 90% of patients in the registry returning to normal and unrestricted function after transplantation.

IMMUNOSUPPRESSION

Medical regimens employed to suppress the normal immune response to a solid organ allograft vary from center to center and are in a constant state of evolution, as more effective agents with improved side-effect profiles and less toxicity are introduced. All currently used regimens are nonspecific, providing general hyporeactivity to foreign antigens rather than donor-specific hyporeactivity and also causing the attendant, and unwanted, susceptibility to infections and malignancy. Most cardiac transplantation programs currently use a three-drug regimen that includes a calcineurin inhibitor (cyclosporine or tacrolimus), an inhibitor of T cell proliferation or differentiation (azathioprine, mycophenolate mofetil, or sirolimus), and at least a short initial course of glucocorticoids. Many programs also include an initial “induction” course of polyclonal or monoclonal antibodies to T cells in the perioperative period to decrease the frequency or severity of early posttransplantation rejection. Most recently introduced have been monoclonal antibodies (daclizumab and basiliximab) that block

the interleukin 2 receptor and may prevent allograft rejection without additional global immunosuppression.

Cardiac allograft rejection is usually diagnosed by endomyocardial biopsy conducted either on a surveillance basis or in response to clinical deterioration. Biopsy surveillance is performed on a regular basis in most programs for the first year postoperatively (or the first 5 years in many programs). Therapy consists of augmentation of immunosuppression, the intensity and duration of which are dictated by the severity of rejection.

LATE POSTTRANSPLANTATION MANAGEMENT ISSUES

Increasing numbers of heart transplant recipients are surviving for years following transplantation and constitute a population of patients with a number of long-term management issues.

Allograft Coronary Artery Disease Despite usually having young donor hearts, cardiac allograft recipients are prone to develop coronary artery disease (CAD). This CAD is generally a diffuse, concentric, and longitudinal process that is quite different from “ordinary” atherosclerotic CAD, which is more focal and often eccentric. The underlying etiology most likely is primarily immunologic injury of the vascular endothelium, but a variety of risk factors influence the existence and progression of CAD, including nonimmunologic factors such as dyslipidemia, diabetes mellitus, and cytomegalovirus (CMV) infection. It is hoped that newer and improved immunosuppressive modalities will reduce the incidence and impact of these devastating complications, which currently account for the majority of late posttransplantation deaths. Thus far, the immunosuppressive agents mycophenolate mofetil and the mammalian target of the rapamycin (mTOR) inhibitors sirolimus and everolimus have been shown to be associated with short-term lower incidence and extent of coronary intimal thickening; in anecdotal reports, institution of sirolimus was associated with some reversal of CAD. The use of statins also is associated with a reduced incidence of this vasculopathy, and these drugs are now used almost universally in transplant recipients unless contraindicated. Palliation of CAD with percutaneous interventions is probably safe and effective in the short term, although the disease often advances relentlessly. Because of the denervated status of the organ, patients rarely experience angina pectoris, even in advanced stages of disease.

Retransplantation is the only definitive form of therapy for advanced allograft CAD. However, the scarcity of donor hearts makes the

1518 decision to pursue retransplantation in an individual patient difficult and ethically complex.

Malignancy An increased incidence of malignancy is a well-recognized sequela of any program of chronic immunosuppression, and organ transplantation is no exception. Lymphoproliferative disorders are among the most frequent posttransplantation complications and, in most cases, seem to be driven by Epstein-Barr virus. Effective therapy includes reduction of immunosuppression (a clear “double-edged sword” in the setting of a life-sustaining organ), administration of antiviral agents, and traditional chemotherapy and radiotherapy. Most recently, specific antilymphocyte (CD20) therapy has shown great promise. Cutaneous malignancies (both basal cell and squamous cell carcinomas) also occur with increased frequency among transplant recipients and can follow aggressive courses. The role of decreasing immunosuppression in the treatment of these cancers is far less clear.

Infections The use of currently available nonspecific immunosuppressive modalities to prevent allograft rejection naturally results in increased susceptibility to infectious complications in transplant recipients. Although the incidence has decreased since the introduction of cyclosporine, infections with unusual and opportunistic organisms are still the major cause of death during the first postoperative year and remain a threat to the chronically immunosuppressed patient throughout life. Effective therapy depends on careful surveillance for early signs and symptoms of opportunistic infection, an extremely aggressive approach to obtaining a specific diagnosis, and expertise in recognizing the more common clinical presentations of infections caused by CMV, *Aspergillus*, and other opportunistic agents.

PROLONGED ASSISTED CIRCULATION

The modern era of mechanical circulatory support can be traced back to 1953, when cardiopulmonary bypass was first used in a clinical setting and ushered in the possibility of brief periods of circulatory support to permit open-heart surgery. Subsequently, a variety of extracorporeal pumps to provide circulatory support for brief periods have been developed. The use of a mechanical device to support the circulation for more than a few hours initially progressed slowly, with the implantation of a total artificial heart in 1969 in Texas by Cooley. This patient survived for 60 h until a donor organ became available, at which point he underwent transplantation. Unfortunately, the patient died of pulmonary complications after transplantation. The entire field of mechanical replacement of the heart then took a decade-long hiatus until the 1980s, when total artificial hearts were reintroduced with much publicity; however, they failed to produce the hoped-for treatment of end-stage heart disease. Starting in the 1970s, in parallel with the development of the total artificial heart, intense research had addressed the development of ventricular assist devices, which provide mechanical assistance for (rather than replacing) the failing ventricle.

Although conceived of initially as alternatives to biologic replacement of the heart, LVADs were introduced—and are still employed primarily—as temporary “bridges” to heart transplantation in candidates in whom medical therapy begins to fail before a donor heart becomes available. Several devices are approved by the U.S. Food and Drug Administration (FDA) and are in widespread use (see later). Those that are implantable within the body are compatible with hospital discharge and offer the patient a chance for life at home during a wait for a donor heart. However successful such “bridging” is for the individual patient, it does nothing to alleviate the scarcity of donor hearts; the ultimate goal in the field remains that of providing a reasonable alternative to biologic replacement of the heart—one that is widely and easily available and cost-effective.

CURRENT INDICATIONS AND APPLICATIONS OF VENTRICULAR ASSIST DEVICES

Currently, there are two major indications for ventricular assistance. First, patients at risk of imminent death from cardiogenic shock are eligible for mechanical support. These patients are generally managed with temporary cardiac assist devices. Second, if patients have a left ventricular ejection fraction <25% or a peak VO_2 <14 mL/kg per min

or are dependent on inotropic therapy or support with intra-aortic balloon counterpulsation, they may be eligible for mechanical support. If they are eligible for heart transplantation, the mechanical circulatory assistance is termed the “bridge to transplantation.” By contrast, if the patient has a contraindication to heart transplantation, the use of the device is deemed to constitute “destination” left ventricular assistance therapy.

BASIC CONCEPTS

Pulsatile vs. Nonpulsatile Devices *Pulsatile* devices are ventricular assist devices whose mechanism of action mandates the alternating filling and emptying of a volume chamber within the device that mimics the mechanism of action of the natural heart. *Nonpulsatile* devices have a mechanism of action that results in continuous blood flow through the device, eliminating the need for pulsatility. The pulsatile devices are larger, bulkier, and associated with greater energy requirements and higher rates of complications than the nonpulsatile devices. However, pulsatile devices provide greater degrees of support and may even be capable of replacing the function of the heart entirely in the form of a total artificial heart. Because of the bulkiness of these devices, many patients are too small to be supported with intracorporeal pulsatile pumps. However, paracorporeal versions are available. These devices are versatile and can be used for right, left, or biventricular assistance/replacement.

Continuous-flow (nonpulsatile) devices are further categorized on the basis of impeller design and mechanism. The older designs have tended to be axial-flow pumps, which operate on the Archimedes screw principle. These devices have an impeller that is in line with the direction of blood flow, and the inlet direction of blood is the same as the outlet direction. Continuous-flow devices have been dependent on the presence of blood-washed bearings within the pump housing and may be associated with an increased risk of blood and platelet activation. The newer devices are centrifugal in design; the blood flow takes a 90° turn between the inlet section of the pump and the outlet section. Another major difference in the newer devices is the absence of blood-washed bearings (with most devices having magnetically levitated impellers). This design allows the construction of smaller pumps with less blood-element activation than the axial-flow designs.

Available Devices In the United States, there are currently four FDA-approved devices that are used as bridges to transplantation in adults. Of these four devices, one is also approved for use as destination therapy or as long-term mechanical support of the heart. A number of other devices are approved only for short-term support in post-cardiac surgery shock or in cardiogenic shock secondary to acute myocardial infarction or fulminant myocarditis; these will not be considered here. So far, no long-term device is totally implantable, and, because of the need for transcatheter connections, all share a common problem with infectious complications. Likewise, all share some tendency to thromboembolic complications and are subject to the possibility of mechanical failure common to any machine.

The total artificial heart (TAH) (Syncardia, Tucson, AZ) is a pneumatic, biventricular, orthotopically implanted ventricular assist device with an externalized driveline connecting it to its console. The TAH is currently the only FDA-approved device for use in patients who have severe biventricular failure.

The Thoratec LVAD (Thoratec Corp., Pleasanton, CA) is an extracorporeal pump that takes blood from a large cannula placed in the left ventricular apex and propels it forward through an outflow cannula inserted into the ascending aorta. The extracorporeal nature of this pump allows its use in small adults for whom intracorporeal pumps would be too large. This device provides not only left but also right ventricular assistance and can be utilized for biventricular support within the same patient (BiVentricular Assist Device).

The HeartMate II LVAD (Thoratec) similarly uses a drainage cannula in the left ventricular apex to drain blood into a small chamber, where the blood is driven by an electrically powered motor that spins a rotor, accelerating blood outflow into the ascending aorta (Fig. 281-2). This device is currently the only FDA-approved axial-flow pump that can be used both as a bridge to transplantation and as destination therapy.

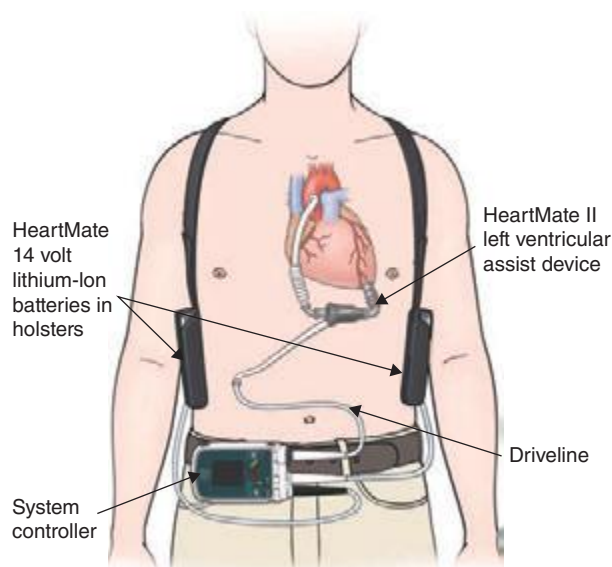


FIGURE 281-2 Diagram of HeartMate II left ventricular assist device (LVAD). (Reprinted with permission from Thoratec Corp., Pleasanton, CA.)

The HeartWare Ventricular Assist System with the HVAD pump (HeartWare Inc., Framingham, MA) is the first third-generation device to be granted FDA approval for use in patients as a bridge to transplantation. The device is a centrifugal pump that is housed completely within the patient's pericardial cavity and provides adequate support for many patients.

RESULTS

The use of these devices in the United States is limited mainly to patients with post-cardiac surgery shock and to those who are bridged to transplantation. The results of bridging to transplantation with the available devices are quite good, with nearly 75% of younger patients receiving a transplant by 1 year and having excellent posttransplantation survival rates.

Publication of the REMATCH (Randomized Evaluation of Mechanical Assistance in the Treatment of Heart Failure) trial in 2001 documented a somewhat improved survival rate in patients who had end-stage heart disease, were not candidates for transplantation, and were randomized to a pulsatile LVAD (albeit with a high rate of complications, especially neurologic issues) as opposed to continued medical therapy. This result led to renewed interest in use of the devices for nonbiologic permanent replacement of heart function as well. Subsequently, this device was supplanted by the HeartMate II axial-flow device, which has dramatically improved the survival of patients with severe end-stage heart disease in whom medical therapy has failed. The patients who had this device implanted had a 2-year survival rate of 58%, whereas the survival rate for patients in the medically treated arm of the original REMATCH trial was only 8%. More recent experience has shown that the mean survival period of patients with a continuous-flow LVAD for destination therapy is approaching 5 years.

Several studies have evaluated the benefit of LVAD therapy as a bridge to transplantation. The most recent data come from a series of 140 patients who underwent implantation of a HeartWare HVAD. Of these patients, 94% achieved the principal outcome (defined as survival to transplantation, recovery of heart function, or ongoing device support) at 180 days. With increased experience and improved outcomes using LVADs as a bridge to transplantation, the ability to maintain end-organ function and limit the progression of pulmonary hypertension—or even to decrease pulmonary vascular resistance—makes mechanical unloading a more attractive option than continued inotropic support. The early bridge-to-transplantation experience demonstrated reduced posttransplantation survival compared with medical management; however, more recent experience has shown equivalent outcomes following transplantation. This result is likely secondary to a trend toward earlier device implantation—i.e., prior to the onset of irreversible end-organ damage.

282 Congenital Heart Disease in the Adult

Jamil A. Aboulhosn, John S. Child

Over a hundred years ago, Sir William Osler, in his classic textbook *The Principles and Practice of Medicine* (New York, Appleton & Co, 1892, pp 659–663), devoted only five pages to “Congenital Affections of the Heart,” with the first sentence declaring that “[t]hese [disorders] have only limited clinical interest, as in a large proportion of cases the anomaly is not compatible with life, and in others nothing can be done to remedy the defect or even to relieve symptoms.” Fortunately, in the intervening century, considerable progress has been made in understanding the basis for these disorders and their effective treatment.

The most common birth defects are cardiovascular in origin. These malformations are due to complex multifactorial genetic and environmental causes. Recognized chromosomal aberrations and mutations of single genes account for <10% of all cardiac malformations. Congenital heart disease (CHD) complicates ~1% of all live births in the general population—about 40,000 births/year—but occurs more frequently in the offspring (about 4–10%, depending on maternal CHD type) of women with CHD. Owing to the remarkable surgical advances over the last 60 years, >90% of afflicted neonates and children now reach adulthood; women with CHD may now frequently successfully bear children after competent repairs. As such, the population with CHD is steadily increasing. Women with CHD are at increased risk for peri- and postpartum complications, but maternal CHD is generally not considered an absolute contraindication to pregnancy unless the mother has certain high-risk features (e.g., cyanosis, pulmonary hypertension, decompensated heart failure, arrhythmias, aortic aneurysm, among others). Consultation with an adult CHD expert is warranted for all females with CHD who desire to become pregnant.

Nearly one and a half million adults with operated or unoperated CHD live in the United States today; there are now more adults than children with CHD in the United States. Because true surgical cures are rare, and all repairs—be they palliative or corrective—may leave residua, sequelae, or complications, most require some degree of lifetime expert surveillance. The anatomic and physiologic changes in the heart and circulation due to any specific CHD lesion are not static but, rather, progress from prenatal life to adulthood. Malformations that are benign or escape detection in childhood may become clinically significant in the adult. Unfortunately, the growing number of adults with CHD has not been paralleled by an adequate increase in the number of specialists and

1520 specialty centers that are trained and equipped to manage this challenging population. Ongoing efforts to increase awareness, resources, and advocacy are essential for the necessary growth of this specialty.

CARDIAC DEVELOPMENT

(See also Chap. 265e) CHD is generally the result of aberrant embryonic development of a normal structure or failure of such a structure to progress beyond an early stage of embryonic or fetal development. This brief section serves to introduce the reader to normal development so that defects may be better understood; by necessity, it is not exhaustive. Cardiogenesis is a finely tuned process with transcriptional control of a complex group of regulatory proteins that activate or inhibit their gene targets in a location- and time-dependent manner. At about 3 weeks of embryonic development, two cardiac cords form and become canalized; at that point, the primordial cardiac tube develops from two sources (cardiac crescent or the first heart field, pharyngeal mesoderm or the second heart field); by 21 days, these fuse into a single cardiac tube beginning at the cranial end. The cardiac tube then elongates and develops discrete constrictions with the following segments from caudal to cranial location: sinus venosus receives the umbilical, vitelline, and common cardinal veins: atrium, ventricle, bulbus cordis, truncus arteriosus, aortic sac, and the aortic arches. The cardiac tube is fixed at the sinus venosus and arterial ends.

Subsequently, in the next few weeks, differential growth of cells causes the tube to elongate and loop as an “S” with the bulboventricular portion moving rightward and the atrium and sinus venosus moving posterior to the ventricle. The primitive atrium and ventricle communicate via the atrioventricular canal from which the endocardial cushion develops into two parts (ventrally and dorsally). The cushions fuse and divide the atrioventricular canal into two atrioventricular inlets and also migrate to help form the ventricular septum. The primitive atrium is divided first by a septum primum membrane, which grows down from the superior wall to the cushions; as this fusion occurs, the mid-portion resorbs in the center forming the ostium secundum. Rightward of the septum primum, a second septum secundum membrane grows down from the ventral-cranial wall toward—but not reaching—the cushions, and covering most, but not all, of the ostium secundum, resulting in a flap of the foramen ovale. The primitive ventricle is partitioned by a finely tuned set of events. The interventricular septum grows up toward the cushions, and the cushions form an upper inlet septum; between the two portions is a hole called the interventricular foramen. The left and right ventricles begin to develop side by side, and the atria and their respective inlet valves align over their ventricles. Finally, these two parts of the septum fuse with the bulboventricular ridges, which, once having septated the truncus arteriosus, extend into the ventricle. The bulboventricular divides into a subaortic portion as the muscular conus resorbs, whereas the subpulmonary section has elongation of its muscular conus. Spiral division of the common truncus arteriosus rotates and aligns the pulmonary artery and aortic portions over their respective outflow tracts, the aortic valve moving posterior over the left ventricle (LV) outflow tract and the pulmonary valve moving anterior over the right ventricle (RV) outflow tract, with a wraparound relationship of the two great arteries.

Early on, the venous systems are bilateral and symmetric and enter two horns of the sinus venosus. Ultimately, except for the coronary sinus, most of the left-sided portions and the left sinus-venosus horn regress, and the systemic venous system empties into the right horn via the inferior and superior vena cavae. The pulmonary venous system, initially connecting to the systemic venous system, develops as buds from the developing lungs, which fuse together in the pulmonary venous confluence, at which point the connection to the systemic system regresses. Simultaneously, a projection from the back wall of the left atrium (the common pulmonary vein) grows posteriorly to merge with the confluence, which then becomes a part of the posterior left atrial wall.

The truncus arteriosus and aortic sac initially develop six paired symmetric arches, which curve posteriorly and become the paired dorsal aortae. The detailed description of the selective regression of some of the arches is not presented in this chapter. In brief summary,

TABLE 282-1 SIMPLE ADULT CONGENITAL HEART DISEASE

Native disease
Uncomplicated congenital aortic valve disease
Mild congenital mitral valve disease (e.g., except parachute valve, cleft leaflet)
Uncomplicated small atrial septal defect
Uncomplicated small ventricular septal defect
Mild pulmonic stenosis
Repaired conditions
Previously ligated or occluded ductus arteriosus
Repaired secundum or sinus venosus atrial septal defect without residua
Repaired ventricular septal defect without residua

this process results in the development of arch 3 as the internal carotid arteries, left arch 4 as the aortic arch and right subclavian artery, and part of arch 6 as the patent ductus arteriosus. The two dorsal thoracic aortae fuse in the abdomen with persistence of the left dorsal aorta.

SPECIFIC CARDIAC DEFECTS

Tables 282-1, 282-2, and 282-3 list CHD malformations as simple, intermediate, or complex. Simple defects generally are single lesions with a shunt or a valvular malformation. Intermediate defects may have two or more simple defects. Complex defects generally have components of an intermediate defect plus more complex cardiac and vascular anatomy, often with cyanosis, and frequently with transposition complexes. The goal of these tables is to suggest when cardiology consultation or advanced CHD specialty care is needed. Patients with complex CHD (which includes most “named” surgeries that usually involve complex CHD) should virtually always be managed in conjunction with an experienced specialty adult CHD center. Patients with intermediate lesions should have an initial consultation and subsequent occasional intermittent follow-up with an adult CHD specialist. Patients with simple lesions often may be managed by a well-informed internist or general cardiologist, although consultation with a specifically trained adult congenital cardiologist is occasionally advisable.

ATRIAL SEPTAL DEFECT

Atrial septal defect (ASD) is a common cardiac anomaly that may be first encountered in the adult and occurs more frequently in females. Sinus venosus ASD occurs high in the atrial septum near the entry of the superior vena cava into the right atrium and is associated frequently with anomalous pulmonary venous connection from the right lung to the superior vena cava or right atrium (Fig. 282-1). Ostium primum ASDs lie adjacent to the atrioventricular valves, either of which may be deformed and regurgitant. Ostium primum ASDs are common in Down’s syndrome, often as part of complex atrioventricular septal defects with a common atrioventricular valve and a posterior defect of the basal portion of the interventricular septum. The most common ostium secundum ASD involves the fossa ovalis and is mid-septal in location; this should not be confused with a patent foramen

TABLE 282-2 INTERMEDIATE COMPLEXITY CONGENITAL HEART DISEASE

Ostium primum or sinus venosus atrial septal defect
Anomalous pulmonary venous drainage, partial or total
Atrioventricular canal defects (partial or complete)
Ventricular septal defect, complicated (e.g., absent or abnormal valves or with associated obstructive lesions, aortic regurgitation)
Coarctation of the aorta
Pulmonic valve stenosis (moderate to severe)
Infundibular right ventricular outflow obstruction of significance
Pulmonary valve regurgitation (moderate to severe)
Patent ductus arteriosus (nonclosed)—moderate to large
Sinus of Valsalva fistula/aneurysm
Subvalvular or supra-valvular aortic stenosis

TABLE 282-3 COMPLEX ADULT CONGENITAL HEART DISEASE

Cyanotic congenital heart diseases (all forms)

Eisenmenger's syndrome

Ebstein's anomaly

Tetralogy of Fallot or pulmonary atresia (all forms)

Transposition of the great arteries

Single ventricle; tricuspid or mitral atresia

Double-outlet ventricle

Truncus arteriosus

Fontan or Rastelli procedures

ovale. Anatomic obliteration of the foramen ovale ordinarily follows its functional closure soon after birth, but residual "probe patency" is a common normal variant; ASD denotes a true deficiency of the atrial septum and implies functional and anatomic patency. The magnitude of the left-to-right shunt depends on the ASD size, ventricular diastolic properties, and the relative impedance in the pulmonary and systemic circulations. The left-to-right shunt causes diastolic overloading of the RV and increased pulmonary blood flow. Patients with ASD are usually asymptomatic in early life, although there may be some physical underdevelopment and an increased tendency for respiratory infections; cardiorespiratory symptoms occur in many older patients. Beyond the fourth decade, a significant number of patients develop atrial arrhythmias, pulmonary arterial hypertension, and right heart failure. Patients exposed to the chronic environmental hypoxemia of high altitude tend to develop pulmonary hypertension at younger ages. In older patients, left-to-right shunting across the ASD increases as progressive systemic hypertension and/or coronary artery disease (CAD) result in reduced compliance of the LV.

Physical Examination Examination usually reveals a prominent RV impulse and palpable pulmonary artery pulsation. The first heart sound is normal or split, with accentuation of the tricuspid valve closure sound. Increased flow across the pulmonic valve is responsible for a midsystolic pulmonary outflow murmur. The second heart sound is widely split and is fixed in relation to respiration. A mid-diastolic rumbling murmur, loudest at the fourth intercostal space and along the left sternal border, reflects increased flow across the tricuspid valve. In ostium primum ASD, an apical holosystolic murmur indicates associated mitral or tricuspid regurgitation or a ventricular septal defect (VSD).

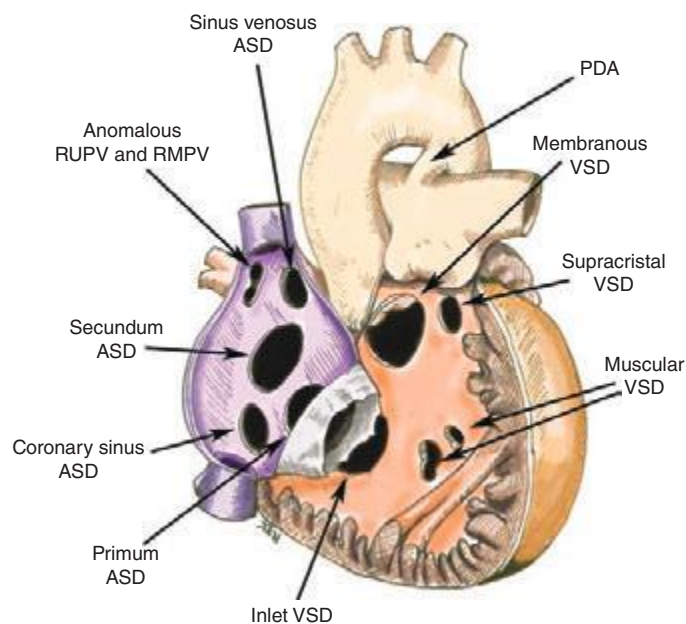


FIGURE 282-1 Types and locations of congenital cardiac defects. ASD, atrial septal defect; PDA, patent ductus arteriosus; RMPV, right middle pulmonary vein; RUPV, right upper pulmonary vein; VSD, ventricular septal defect.

These findings are altered when increased pulmonary vascular resistance causes diminution of the left-to-right shunt. Both the pulmonary outflow and tricuspid inflow murmurs decrease in intensity, the pulmonic component of the second heart sound and a systolic ejection sound are accentuated, the two components of the second heart sound may fuse, and a diastolic murmur of pulmonic regurgitation appears. Cyanosis and clubbing accompany the development of a right-to-left shunt (see "Ventricular Septal Defect" later in this chapter). In adults with an ASD and atrial fibrillation, the physical findings may be confused with mitral stenosis with pulmonary hypertension because the tricuspid diastolic flow murmur and widely split second heart sound may be mistakenly thought to represent the diastolic murmur of mitral stenosis and the mitral "opening snap," respectively.

Electrocardiogram In ostium secundum ASD, electrocardiogram (ECG) usually shows right-axis deviation and an rSr' pattern in the right precordial leads representing enlargement of the RV outflow tract. An ectopic atrial pacemaker or first-degree heart block may occur with the sinus venous ASD. In ostium primum ASD, the RV conduction defect is accompanied by left superior axis deviation and counterclockwise rotation of the frontal plane QRS loop. Varying degrees of RV and right atrial (RA) enlargement or hypertrophy may occur with each type of defect, depending on the presence and degree of pulmonary hypertension. Chest x-ray shows an enlarged RA and RV, and pulmonary artery and its branches; increased pulmonary vascular markings of left-to-right shunt vascularity will diminish if pulmonary vascular disease develops.

Echocardiogram Echocardiography reveals pulmonary arterial and RV and RA dilatation with abnormal (paradoxical) ventricular septal motion in the presence of a significant right heart volume overload. The ASD may be visualized directly by two-dimensional imaging, color-flow imaging, or echocontrast. Echocardiography and Doppler examination have supplanted cardiac catheterization. Transesophageal echocardiography is indicated if the transthoracic echocardiogram is ambiguous, which is often the case with sinus venous defects, or for guiding catheter device closure (Fig. 282-2). Cardiac catheterization is performed if inconsistencies exist in the clinical data, if significant pulmonary hypertension or associated malformations are suspected, if CAD is a possibility, or when attempting transcatheter closure of the ASD.

TREATMENT ATRIAL SEPTAL DEFECT

Operative repair, usually with a patch of pericardium or of prosthetic material or percutaneous transcatheter device closure, if the ASD is of an appropriate size and shape, should be advised for all patients with uncomplicated secundum ASD with significant left-to-right shunting, i.e., pulmonary-to-systemic flow ratios $\geq 1.5:1$. Excellent results may be anticipated, at low risk, even in patients >40 years, in the absence of severe pulmonary hypertension. In ostium primum ASD, cleft mitral valves may require repair in addition to patch closure of the ASD. Closure is not usually carried out in patients with small defects and trivial left-to-right shunts or in those with severe pulmonary vascular disease without a significant left-to-right shunt. However, the use of pulmonary vasodilators with resultant reduction in pulmonary artery pressure and resistance may allow closure of ASD in patients with pulmonary vascular disease.

Patients with sinus venosus or ostium secundum ASDs rarely die before the fifth decade. During the fifth and sixth decades, the incidence of progressive symptoms, often leading to severe disability, increases substantially. Medical management should include prompt treatment of respiratory tract infections; antiarrhythmic medications for atrial fibrillation or supraventricular tachycardia; and the usual measures for hypertension, coronary disease, or heart failure (Chap. 279), if these complications occur. The risk of infective endocarditis is low, and antibiotic prophylaxis is not recommended (Chap. 155).

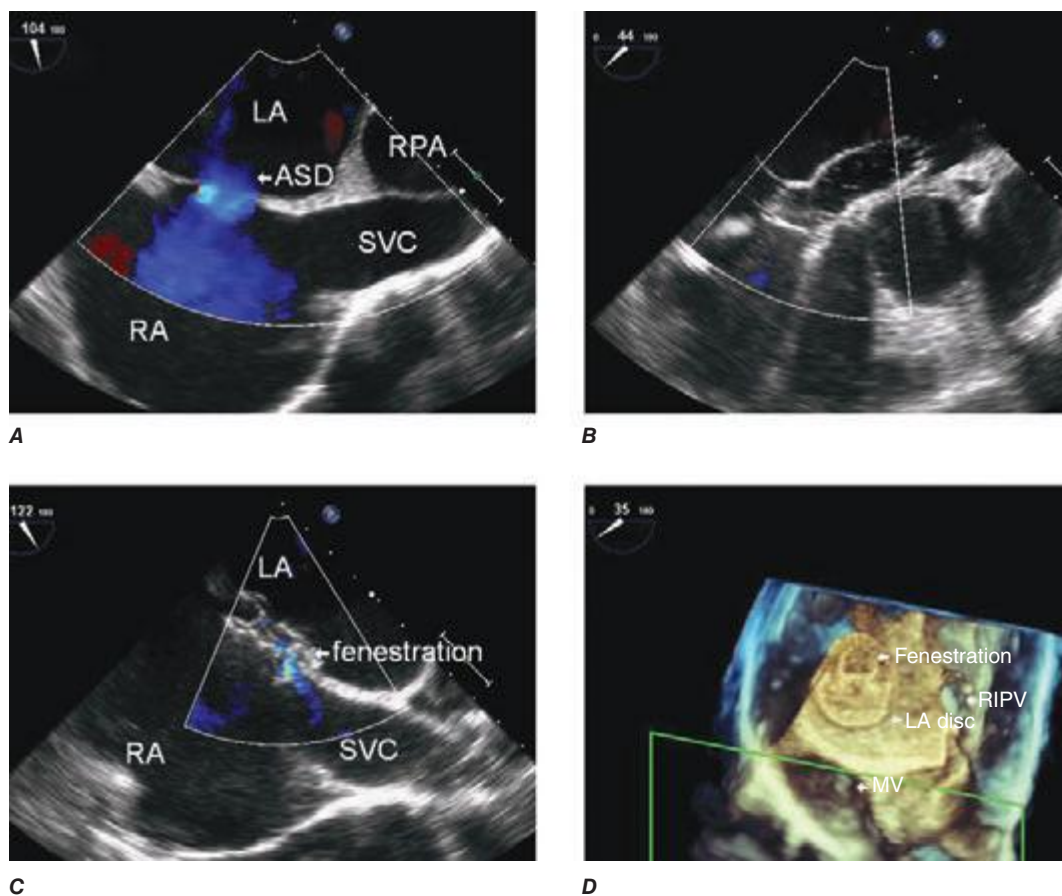


FIGURE 282-2 **A.** Transesophageal echocardiogram demonstrating a secundum-type atrial septal defect (ASD) with shunting from the left atrium (LA) to the right atrium (RA). The right pulmonary artery (RPA) and superior vena cava (SVC) are labeled. **B.** Transcatheter balloon sizing of the ASD. **C.** Atrial septal occluder placement with a small manually created “fenestration” within the device that continues to allow a small amount of flow from the LA to the RA; this is used as a means of preventing left atrial hypertension after ASD closure. Left atrial hypertension may occur in older patients with decreased left ventricular compliance. **D.** Three-dimensional image of the septal occluder en-face; note the fenestration in the LA disc. The mitral valve (MV) and right inferior pulmonary vein (RIPV) are labeled.

VENTRICULAR SEPTAL DEFECT

VSD is one of the most common of all cardiac birth defects, either as an isolated defect or as a component of a combination of anomalies (Fig. 282-1). The VSD is usually single and situated in the membranous or midmuscular portion of the septum. The functional disturbance depends on its size and on the status of the pulmonary vascular bed. Only small- or moderate-size VSDs are seen initially in adulthood, as most patients with an isolated large VSD come to medical or surgical attention early in life.

A wide spectrum exists in the natural history of VSD, ranging from spontaneous closure to congestive cardiac failure and death in infancy. Included within this spectrum is the possible development of pulmonary vascular obstruction, RV outflow tract obstruction, aortic regurgitation, or infective endocarditis. Spontaneous closure is more common in patients born with a small VSD, which occurs in early childhood in most. The pulmonary vascular bed is often a principal determinant of the clinical manifestations and course of a given VSD and feasibility of surgical repair. Increased pulmonary arterial pressure results from increased pulmonary blood flow and/or resistance, the latter usually the result of obstructive, obliterative structural changes within the pulmonary vascular bed. It is important to quantitate and compare pulmonary-to-systemic flows and resistances in patients with severe pulmonary hypertension. The term Eisenmenger’s syndrome is applied to patients with a large communication between the two circulations at the aortopulmonary, ventricular, or atrial levels and bidirectional or predominantly right-to-left shunts because of high resistance and obstructive pulmonary hypertension.

Patients with large VSDs and pulmonary hypertension are at greatest risk for developing pulmonary vascular disease. Large VSDs should

be corrected early in life when pulmonary vascular disease is not severely elevated. In patients with Eisenmenger’s syndrome, symptoms in adult life consist of exertional dyspnea, chest pain, syncope, and hemoptysis. The right-to-left shunt leads to cyanosis, clubbing, and erythrocytosis (see below). The degree to which pulmonary vascular resistance is elevated before operation is a critical factor determining prognosis. If the pulmonary vascular resistance is one-third or less of the systemic value, progression of pulmonary vascular disease after operation is unusual; however, if a moderate to severe increase in pulmonary vascular resistance exists preoperatively, either no change or a progression of pulmonary vascular disease is common postoperatively. Pregnancy is contraindicated in Eisenmenger’s syndrome. The mother’s health is most at risk if she has a cardiovascular lesion associated with pulmonary vascular disease and pulmonary hypertension (e.g., Eisenmenger’s physiology or mitral stenosis) or severe LV outflow tract obstruction (e.g., aortic stenosis), but she is also at risk of death with any malformation that may cause heart failure or a hemodynamically important arrhythmia. The fetus is most at risk with maternal cyanosis, heart failure, or pulmonary hypertension.

RV outflow tract obstruction develops in ~5–10% of patients who present in infancy with a moderate to large left-to-right shunt. With time, as subvalvular RV outflow tract obstruction progresses, the findings in these patients whose VSD remains sizable begin to resemble more closely those of the cyanotic tetralogy of Fallot. In ~5% of patients, aortic valve regurgitation results from insufficient cusp tissue or prolapse of the cusp through the interventricular defect; the aortic regurgitation then complicates and dominates the clinical course. Echocardiography with spectral and color Doppler examination defines the number and location of defects in the ventricular septum

and associated anomalies and the hemodynamic physiology of the defect(s). Hemodynamic and angiographic study may be occasionally required to assess the status of the pulmonary vascular bed and clarify details of the altered anatomy. Cross-sectional imaging modalities such as computed tomography (CT) or magnetic resonance imaging (MRI) are useful in delineating complex anatomy and assessing extra-cardiac structures.

TREATMENT VENTRICULAR SEPTAL DEFECT

Closure is not recommended for patients with normal pulmonary arterial pressures with small shunts (pulmonary-to-systemic flow ratios of <1.5:1). Operative correction or transcatheter closure is indicated when there is a moderate to large left-to-right shunt with a pulmonary-to-systemic flow ratio >1.5:1, in the absence of prohibitively high levels of pulmonary vascular resistance (pulmonary arterial resistance is less than two-thirds of systemic arterial resistance).

In patients with Eisenmenger's VSD, pulmonary arterial vasodilators and both single- or double-lung transplantation with intracardiac defect repair or heart/lung transplantation show promise for improvement in symptoms (Chaps. 281 and 320e). Chronic hypoxemia in cyanotic CHD results in secondary erythrocytosis due to increased erythropoietin production (Chap. 49). The term polycythemia is a misnomer; white cell counts are normal, and platelet counts are normal to decreased. Compensated erythrocytosis with iron-replete equilibrium hematocrits rarely results in symptoms of hyperviscosity at hematocrits <65% and occasionally not even with hematocrits ≥70%. For this reason, therapeutic phlebotomy is rarely required in compensated erythrocytosis. In contrast, patients with decompensated erythrocytosis fail to establish equilibrium with unstable, rising hematocrits and recurrent hyperviscosity symptoms. Therapeutic phlebotomy, a two-edged sword, allows temporary relief of symptoms but limits oxygen delivery, begets instability of the hematocrit, and compounds the problem by iron depletion. Iron-deficiency symptoms are usually indistinguishable from those of hyperviscosity; progressive symptoms after recurrent phlebotomy are usually due to iron depletion with hypochromic microcytosis. Iron depletion results in a larger number of smaller (microcytic) hypochromic red cells that are less capable of carrying oxygen and less deformable in the microcirculation; with more of them relative to plasma volume, viscosity is greater than for an equivalent hematocrit with fewer, larger, iron-replete, deformable cells. As such, iron-depleted erythrocytosis results in increasing symptoms due to decreased oxygen delivery to the tissues.

Hemostasis is abnormal in cyanotic CHD, due, in part, to the increased blood volume and engorged capillaries, abnormalities in platelet function, and sensitivity to aspirin or nonsteroidal anti-inflammatory agents, as well as abnormalities of the extrinsic and intrinsic coagulation system. Oral contraceptives are often contraindicated for cyanotic women because of the enhanced risk of vascular thrombosis. Symptoms of hyperviscosity can be produced in any cyanotic patient with erythrocytosis if dehydration reduces plasma volume. Phlebotomy for symptoms of hyperviscosity not due to dehydration or iron deficiency is a simple outpatient removal of 500 mL of blood over 45 min with isovolumetric replacement with isotonic saline. Acute phlebotomy without volume replacement is contraindicated. Iron repletion in decompensated iron-depleted erythrocytosis reduces iron-deficiency symptoms, but must be done gradually to avoid an excessive rise in hematocrit and resulting hyperviscosity.

PATENT DUCTUS ARTERIOSUS

The ductus arteriosus is a vessel leading from the bifurcation of the pulmonary artery to the aorta just distal to the left subclavian artery (Fig. 282-1). Normally, the vascular channel is open in the fetus but closes immediately after birth. The flow across the ductus is determined by the pressure and resistance relationships between the systemic and pulmonary circulations and by the cross-sectional area and length of the ductus. In most adults with this anomaly, pulmonary pressures are normal, and a gradient and shunt from aorta to pulmonary artery

persist throughout the cardiac cycle, resulting in a characteristic thrill and a continuous “machinery” murmur with late systolic accentuation at the upper left sternal edge. In adults who were born with a large left-to-right shunt through the ductus arteriosus, pulmonary vascular obstruction (Eisenmenger's syndrome) with pulmonary hypertension, right-to-left shunting, and cyanosis have usually developed. Severe pulmonary vascular disease results in reversal of flow through the ductus; unoxygenated blood is shunted to the descending aorta; and the toes—but not the fingers—become cyanotic and clubbed, a finding termed differential cyanosis (Fig. 282-3). The leading causes of death in adults with patent ductus arteriosus are cardiac failure and infective endocarditis; occasionally, severe pulmonary vascular obstruction may cause aneurysmal dilatation, calcification, and rupture of the ductus.

TREATMENT PATENT DUCTUS ARTERIOSUS

In the absence of severe pulmonary vascular disease and predominant left-to-right shunting of blood, the patent ductus should be surgically ligated or divided. Transcatheter closure has become common for appropriately shaped defects. Operation should be deferred for several months in patients treated successfully for infective endocarditis because the ductus may remain somewhat edematous and friable.

AORTIC ROOT-TO-RIGHT-HEART SHUNTS

The three most common causes of aortic root-to-right-heart shunts are congenital aneurysm of an aortic sinus of Valsalva with fistula, coronary arteriovenous fistula, and anomalous origin of the left coronary artery from the pulmonary trunk. Aneurysm of an aortic sinus of Valsalva consists of a separation or lack of fusion between the media of the aorta and the annulus of the aortic valve. Rupture usually occurs in the third or fourth decade of life; most often, the aorticocardiac fistula is between the right coronary cusp and the RV; but occasionally, when the noncoronary cusp is involved, the fistula drains into the RA. Abrupt rupture causes chest pain, bounding pulses, a continuous murmur accentuated in diastole, and volume overload of the heart. Diagnosis is confirmed by two-dimensional and Doppler echocardiographic studies; cardiac catheterization quantitates the left-to-right shunt, and thoracic aortography visualizes the fistula. Medical management is directed at cardiac failure, arrhythmias, or endocarditis. At operation, the aneurysm is closed and amputated, and the aortic wall is reunited with the heart, either by direct suture or with a patch or prosthesis. Transcatheter device closure is a less invasive and effective alternative to surgery.

Coronary arteriovenous fistula, an unusual anomaly, consists of a communication between a coronary artery and another cardiac chamber, usually the coronary sinus, RA, or RV. The shunt is usually of small magnitude, and myocardial blood flow is not usually compromised; if the shunt is large, there may be a coronary “steal” syndrome with myocardial ischemia and possible angina or ventricular arrhythmias. Potential complications include infective endocarditis; thrombus formation with occlusion or distal embolization with myocardial infarction; rupture of an aneurysmal fistula; and, rarely, pulmonary hypertension and congestive failure. A loud, superficial, continuous murmur at the lower or midsternal border usually prompts a further evaluation of asymptomatic patients. Doppler echocardiography demonstrates the site of drainage; if the site of origin is proximal, it may be detectable by two-dimensional echocardiography. Angiography (classic catheterization, CT, or magnetic resonance angiography) permits identification of the size and anatomic features of the fistulous tract, which may be closed by suture or transcatheter obliteration.

The third anomaly causing a shunt from the aortic root to the right heart is anomalous origin of the left coronary artery from the pulmonary artery. In this condition, oxygenated blood from the aortic root flows via a dilated right coronary artery and collaterals to the left coronary artery and retrograde to the lower pressure pulmonary artery circulation via the anomalous left main coronary artery (which emerges from the pulmonary artery). Myocardial infarction and

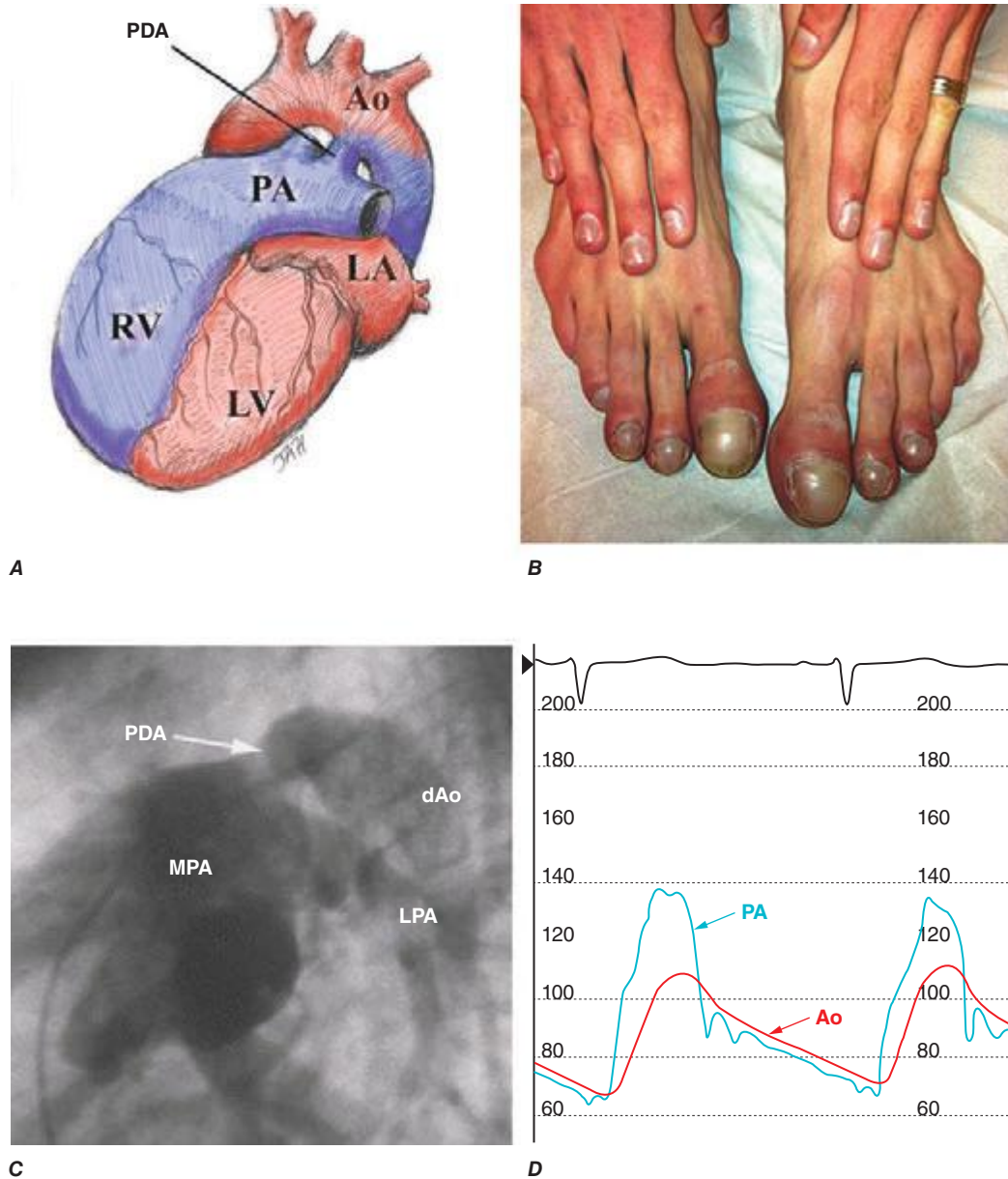


FIGURE 282-3 **A.** Patent ductus arteriosus (PDA) in a patient with severe pulmonary hypertension (Eisenmenger's syndrome). Due to the suprasystemic pulmonary arterial resistance, deoxygenated (cyanotic) blood from the right ventricle (RV) and pulmonary artery (PA) is shunted across the PDA to the aorta (Ao). The left atrium (LA) and left ventricle (LV) are labeled. **B.** Differential clubbing and cyanosis of the toes due to lower extremity perfusion by the deoxygenated blood crossing the PDA. **C.** Angiogram in a dilated main pulmonary artery (MPA) with shunting noted across the PDA to the descending aorta (dAo). The left pulmonary artery (LPA) is labeled. **D.** Direct pressure recordings in the Ao and PA demonstrating suprasystemic PA systolic pressure.

fibrosis commonly lead to death within the first year, although up to 20% of patients survive to adolescence and beyond without surgical correction. The diagnosis is supported by the ECG findings of an anterolateral myocardial infarction and left ventricular hypertrophy (LVH). Operative management of adults consists of coronary artery reimplantation, coronary artery bypass with an internal mammary artery graft, or saphenous vein–coronary artery graft.

CONGENITAL AORTIC STENOSIS

Malformations that cause obstruction to LV outflow include congenital valvular aortic stenosis, discrete subaortic stenosis, or supra-aortic stenosis. Bicuspid aortic valves are more common in males than in females. The congenital bicuspid aortic valve, which may initially be functionally normal, is one of the most common congenital malformations of the heart and may go undetected in early life. Because bicuspid valves may develop stenosis or regurgitation with time or be the site of infective endocarditis, the lesion may be difficult to distinguish in older adults from acquired rheumatic or degenerative calcific aortic

valve disease. The dynamics of blood flow associated with a congenitally deformed, rigid aortic valve commonly lead to thickening of the cusps and, in later life, to calcification. Hemodynamically significant obstruction causes concentric hypertrophy of the LV wall. The ascending aorta is often dilated, misnamed “poststenotic” dilatation; this is due to histologic abnormalities of the aortic media and may result in aortic dissection. Diagnosis is made by echocardiography, which reveals the morphology of the aortic valve and aortic root and quantitates severity of stenosis or regurgitation. **The clinical manifestations and hemodynamic abnormalities are discussed in Chap. 283.**

TREATMENT VALVULAR AORTIC STENOSIS

In patients with diminished cardiac reserve, medical management includes the administration of digoxin and diuretics and sodium restriction while awaiting operation. A dilated aortic root may require beta blockers, angiotensin receptor blockers, or angiotensin-converting enzyme inhibitors. Aortic valve replacement is

indicated in adults with critical obstruction, i.e., with an aortic valve area $<0.45 \text{ cm}^2/\text{m}^2$, with symptoms secondary to LV dysfunction or myocardial ischemia, or with hemodynamic evidence of LV dysfunction. In asymptomatic children or adolescents or young adults with critical aortic stenosis without valvular calcification or these features, aortic balloon valvuloplasty is often useful (Chap. 296e). If surgery is contraindicated in older patients because of a complicating medical problem such as malignancy or renal or hepatic failure, balloon valvuloplasty may provide short-term improvement. This procedure may serve as a bridge to aortic valve replacement in patients with severe heart failure. Transcatheter aortic valve replacement is a potential alternative to surgery.

SUBAORTIC STENOSIS The discrete form of subaortic stenosis consists of a membranous diaphragm or fibromuscular ring encircling the LV outflow tract just beneath the base of the aortic valve. The jet impact from the subaortic stenotic jet on the underside of the aortic valve often begets progressive aortic valve fibrosis and valvular regurgitation. Echocardiography demonstrates the anatomy of the subaortic obstruction; Doppler studies show turbulence proximal to the aortic valve and can quantitate the pressure gradient and severity of aortic regurgitation. Treatment consists of complete excision of the membrane or fibromuscular ring.

SUPRAVALVULAR AORTIC STENOSIS This is a localized or diffuse narrowing of the ascending aorta originating just above the level of the coronary arteries at the superior margin of the sinuses of Valsalva. In contrast to other forms of aortic stenosis, the coronary arteries are subjected to elevated systolic pressures from the LV, are often dilated and tortuous, and are susceptible to premature atherosclerosis. The coronary ostia may also become obstructed by the aortic valve leaflets. In most patients, a genetic defect for the anomaly is located in the same chromosomal region as elastin on chromosome 7. Supravalvular aortic stenosis is the most commonly associated cardiac defect in Williams-Beuren syndrome, typically comprising the following: “elfin” facies, low nasal bridge, cheerful demeanor, mental retardation with retained language skills and love of music, supravalvular aortic stenosis, and transient hypercalcemia.

COARCTATION OF THE AORTA

Narrowing or constriction of the lumen of the aorta may occur anywhere along its length but is most common distal to the origin of the left subclavian artery near the insertion of the ligamentum arteriosum. Coarctation occurs in ~7% of patients with CHD, is more common in males than females, and is particularly frequent in patients with gonadal dysgenesis (e.g., Turner’s syndrome). Clinical manifestations depend on the site and extent of obstruction and the presence of associated cardiac anomalies, most commonly a bicuspid aortic valve. Circle of Willis aneurysms may occur in up to 10%.

Most children and young adults with isolated, discrete coarctation are asymptomatic. Headache, epistaxis, chest pressure, and claudication with exercise may occur, and attention is usually directed to the cardiovascular system when a heart murmur or hypertension in the upper extremities and absence, marked diminution, or delayed pulsations in the femoral arteries are detected on physical examination. Enlarged and pulsatile collateral vessels may be palpated in the intercostal spaces anteriorly, in the axillae, or posteriorly in the interscapular area. The upper extremities and thorax may be more developed than the lower extremities. A midsystolic murmur over the left interscapular space may become continuous if the lumen is narrowed sufficiently to result in a high-velocity jet across the lesion throughout the cardiac cycle. Additional systolic and continuous murmurs over the lateral thoracic wall may reflect increased flow through dilated and tortuous collateral vessels. The ECG usually reveals LV hypertrophy. Chest x-ray may show a dilated left subclavian artery high on the left mediastinal border and a dilated ascending aorta. Indentation of the aorta at the site of coarctation and pre- and poststenotic dilatation (the “3” sign) along the left paramediastinal shadow are essentially pathognomonic. Notching of the third to ninth ribs, an important radiographic sign, is due to inferior rib erosion by dilated collateral vessels.

Two-dimensional echocardiography from suprasternal windows identifies the site of coarctation; Doppler quantitates the pressure gradient. Transesophageal echocardiography and MRI or CT allow visualization of the length and severity of the obstruction and associated collateral arteries. In adults, cardiac catheterization is indicated primarily to evaluate the coronary arteries or to perform catheter-based intervention (angioplasty and stent of the coarctation).

The chief hazards of proximal aortic severe hypertension include cerebral aneurysms and hemorrhage, aortic dissection and rupture, premature coronary arteriosclerosis, aortic valve failure, and LV failure; infective endarteritis may occur on the coarctation site or endocarditis may settle on an associated bicuspid aortic valve, which is estimated to be present in 50% of patients.

TREATMENT COARCTATION OF THE AORTA

Treatment is surgical or involves percutaneous catheter balloon dilatation with stent placement; the details of selection of therapy are beyond this review; however, the use of transcatheter treatment techniques has increased dramatically, and many previously “surgical” cases are treated via percutaneous or hybrid techniques. Late postoperative systemic hypertension in the absence of residual coarctation is related partly to the duration of preoperative hypertension. Follow-up of rest and exercise blood pressures is important; many have systolic hypertension only during exercise, in part due to a diffuse vasculopathy and to noncompliance of the stented or surgically reconstructed region. All operated or stented coarctation patients deserve a high-quality MRI or CT procedure in follow-up.

PULMONARY STENOSIS WITH INTACT VENTRICULAR SEPTUM

Obstruction to RV outflow may be localized to the supravalvular, valvular, or subvalvular levels or occur at a combination of these sites. Multiple sites of narrowing of the peripheral pulmonary arteries are a feature of rubella embryopathy and may occur with both the familial and sporadic forms of supravalvular aortic stenosis. Valvular pulmonic stenosis (PS) is the most common form of isolated RV obstruction.

The severity of the obstructing lesion, rather than the site of narrowing, is the most important determinant of the clinical course. In the presence of a normal cardiac output, a peak systolic pressure gradient $<30 \text{ mmHg}$ indicates mild PS and $>50 \text{ mmHg}$ indicates severe PS; pressures between these limits are considered to indicate moderate stenosis. Patients with mild PS are generally asymptomatic and demonstrate little or no progression in the severity of obstruction with age. In patients with more significant stenosis, the severity may increase with time. Symptoms vary with the degree of obstruction. Fatigue, dyspnea, RV failure, and syncope may limit the activity of older patients, in whom moderate or severe obstruction may prevent an augmentation of cardiac output with exercise. In patients with severe obstruction, the systolic pressure in the RV may exceed that in the LV, because the ventricular septum is intact. RV ejection is prolonged with moderate or severe stenosis, and the sound of pulmonary valve closure is delayed and soft. RV hypertrophy reduces the compliance of that chamber, and a forceful RA contraction is necessary to augment RV filling. A fourth heart sound; prominent a waves in the jugular venous pulse; and, occasionally, presystolic pulsations of the liver reflect vigorous atrial contraction. The clinical diagnosis is supported by a left parasternal lift and harsh systolic crescendo-decrescendo murmur and thrill at the upper left sternal border, typically preceded by a systolic ejection sound if the obstruction is due to a mobile nondysplastic pulmonary valve. The holosystolic murmur of tricuspid regurgitation may accompany severe PS, especially in the presence of congestive heart failure. Cyanosis usually reflects right-to-left shunting through a patent foramen ovale or ASD. In patients with supravalvular or peripheral pulmonary arterial stenosis, the murmur is systolic or continuous and is best heard over the area of narrowing, with radiation to the peripheral lung fields.

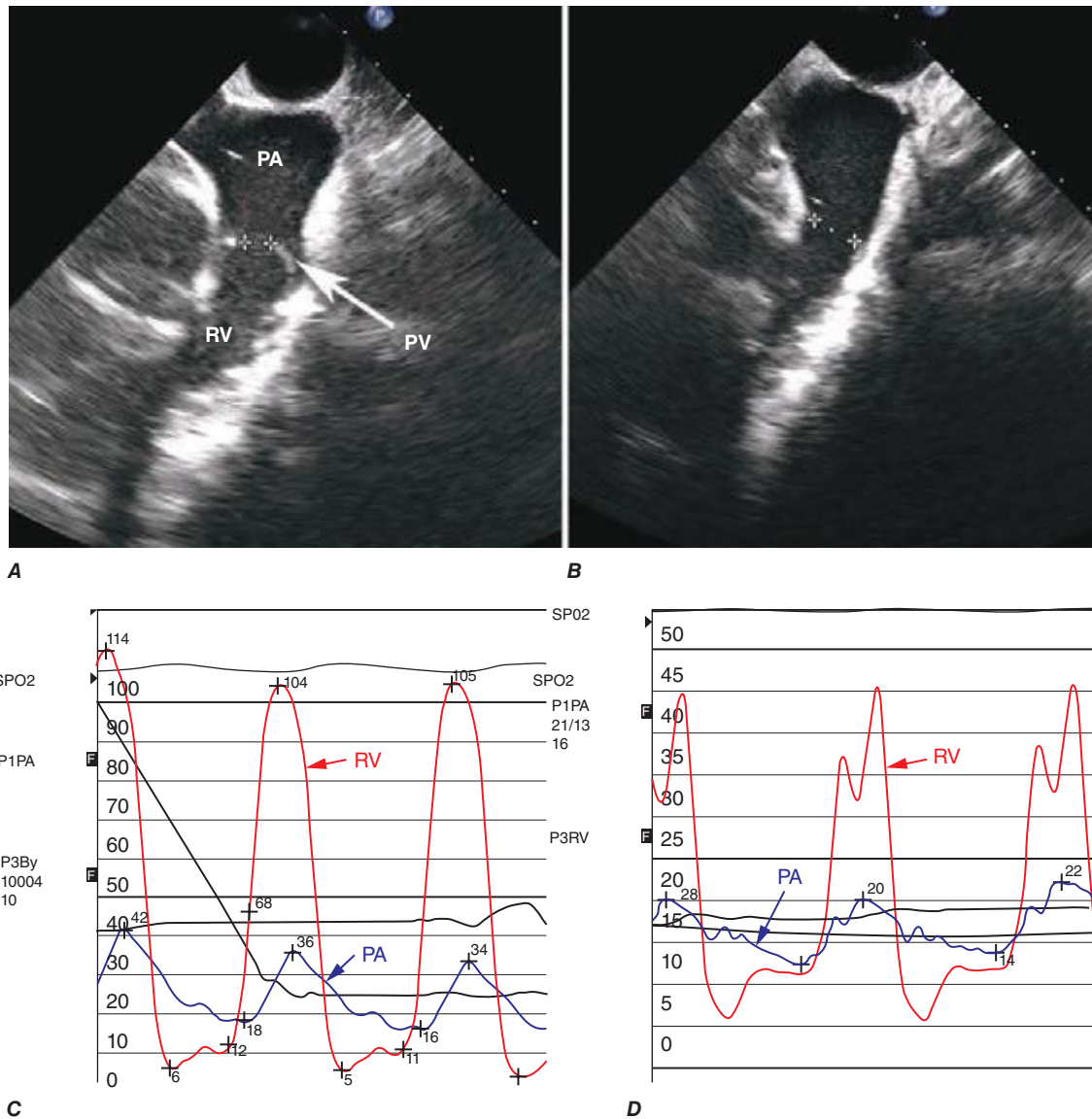


FIGURE 282-4 **A.** Transesophageal echocardiogram of a patient with severe pulmonary stenosis due to a mobile and doming pulmonary valve (PV). The pulmonary artery (PA) and the right ventricle (RV) are labeled. **B.** Following balloon valvuloplasty, the pulmonary valve orifice is larger. **C.** Simultaneous RV and PA pressure tracings before balloon valvuloplasty; the peak-to-peak gradient across the pulmonary valve is ~ 70 mmHg. **D.** After balloon valvuloplasty, the peak-to-peak gradient is reduced to ~ 25 mmHg.

In mild cases, the ECG is normal, whereas moderate and severe stenoses are associated with RV hypertrophy. The chest x-ray with mild or moderate PS shows a heart of normal size with normal lung vascularity. In pulmonary valvular stenosis, dilatation of the main and left pulmonary arteries occurs in part due to the direction of the PS jet and in part due to intrinsic tissue weakness. With severe obstruction, RV hypertrophy is generally evident. The pulmonary vascularity may be reduced with severe stenosis, RV failure, and/or a right-to-left shunt at the atrial level. Two- and three-dimensional echocardiography visualizes pulmonary valve morphology; the outflow tract pressure gradient is quantitated by Doppler echocardiography (Fig. 282-4).

TREATMENT PULMONARY STENOSIS

The cardiac catheter technique of balloon valvuloplasty (Chap. 272) is usually effective, and the surgery is rarely necessary. Multiple stenoses of the peripheral pulmonary arteries are effectively treated with transcatheter angioplasty or stenting.

TETRALOGY OF FALLOT

The four components of the tetralogy of Fallot are malaligned VSD, obstruction to RV outflow, aortic override of the VSD, and RV hypertrophy due to the RV's response to aortic pressure via the large VSD.

The severity of RV outflow obstruction determines the clinical presentation. The severity of hypoplasia of the RV outflow tract varies from mild to complete (pulmonary atresia). Pulmonary valve stenosis and supravalvular and peripheral pulmonary arterial obstruction may coexist; rarely, there is unilateral absence of a pulmonary artery (usually the left). A right-sided aortic arch and descending thoracic aorta occur in $\sim 25\%$.

The relationship between the resistance of blood flow from the ventricles into the aorta and into the pulmonary artery plays a major role in determining the hemodynamic and clinical picture. When the RV outflow obstruction is severe, pulmonary blood flow is reduced markedly, and a large volume of desaturated systemic venous blood shunts right-to-left across the VSD. Severe cyanosis and erythrocytosis occur, and symptoms of systemic hypoxemia are prominent. In many infants and children, the obstruction is mild but progressive.

The ECG shows RV hypertrophy. Chest x-ray shows a normal-sized, boot-shaped heart (*coeur en sabot*) with a prominent RV and a concavity in the region of the pulmonary conus. Pulmonary vascular markings are typically diminished, and the aortic arch and knob may be on the right side. Echocardiography demonstrates the malaligned VSD with the overriding aorta and the site and severity of PS, which may be subpulmonic (fixed or dynamic), at the pulmonary valve or in the main or branch pulmonary arteries. Classic contrast angiography may provide details regarding the RV outflow tract, pulmonary valve and annulus, and caliber of the main branches of the pulmonary artery, as well as about possible associated aortopulmonary collaterals. Coronary arteriography identifies the anatomy and course of the coronary arteries, which may be anomalous. Cardiac MRI and CT complement echocardiography and provide much of the information gathered by angiography as well as additional functional information. MRI is considered the clinical gold standard for quantification of RV volume and function as well as quantification of the pulmonary regurgitation severity.

TREATMENT TETRALOGY OF FALLOT

For a variety of reasons, only a few adults with tetralogy of Fallot have not had some form of previous surgical intervention. Reoperation in adults is most commonly for severe pulmonary regurgitation or pulmonary stenosis. Long-term concerns about ventricular function persist. Ventricular and atrial arrhythmias occur, respectively, in 15% and 25% of adults and may require medical treatment, electrophysiologic study and ablation, defibrillator placement, or transcatheter or surgical intervention, usually including pulmonary valve replacement. Transcatheter pulmonary valve replacement is widely used in patients meeting anatomic criteria. The aortic root has a medial tissue defect; it is commonly enlarged and associated with aortic regurgitation. Endocarditis remains a risk despite surgical repair.

COMPLETE TRANSPOSITION OF THE GREAT ARTERIES

This condition is commonly called *dextro-* or *D-transposition of the great arteries*. The aorta arises rightward anteriorly from the RV, and the pulmonary artery emerges leftward and posteriorly from the LV, which results in two separate parallel circulations; some communication between them must exist after birth to sustain life. Most patients have an interatrial communication, two-thirds have a patent ductus arteriosus, and about one-third have an associated VSD. Transposition is more common in males and accounts for ~10% of cyanotic heart disease. The course is determined by the degree of tissue hypoxemia, the ability of each ventricle to sustain an increased workload in the presence of reduced coronary arterial oxygenation, the nature of the associated cardiovascular anomalies, and the status of the pulmonary vascular bed. Patients who do not undergo surgical palliation generally do not survive to reach adulthood. The long-term outcomes in those that have undergone surgery are in large part determined by the type of surgery performed. By the third decade of life, ~30% of patients with “atrial switch” operations will have developed decreased RV function and progressive tricuspid regurgitation, which may lead to congestive heart failure. Pulmonary vascular obstruction develops by 1–2 years of age in patients with an associated large VSD or large patent ductus arteriosus in the absence of obstruction to LV outflow.

TREATMENT TRANSPOSITION OF THE GREAT ARTERIES

The balloon or blade catheter or surgical creation or enlargement of an interatrial communication in the neonate is the simplest procedure for providing increased intracardiac mixing of systemic and pulmonary venous blood. Systemic pulmonary artery anastomosis may be indicated in the patient with severe obstruction to LV outflow and diminished pulmonary blood flow. Intracardiac repair may be accomplished by rearranging the venous returns

(intraatrial switch, i.e., Mustard or Senning operation) so that the systemic venous blood is directed to the mitral valve and, thence, to the LV and pulmonary artery, while the pulmonary venous blood is diverted through the tricuspid valve and RV to the aorta. The late survival after these repairs is good, but arrhythmias (e.g., atrial flutter) or conduction defects (e.g., sick sinus syndrome) occur in ~50% of such patients by 30 years after the intraatrial switch surgery. Progressive dysfunction of the systemic subaortic RV, tricuspid regurgitation, ventricular arrhythmias, cardiac arrest, and late sudden death are worrisome features. Preferably, this malformation is corrected in infancy by transposing both coronary arteries to the posterior artery and transecting, contraposing, and anastomosing the aorta and pulmonary arteries (arterial-switch operation). For patients with a VSD in whom it is necessary to bypass a severely obstructed LV outflow tract, corrective operation employs an intra-cardiac ventricular baffle and extracardiac prosthetic conduit to replace the pulmonary artery (Rastelli procedure).

SINGLE VENTRICLE

This is a family of complex lesions with both atrioventricular valves or a common atrioventricular valve opening to a single ventricular chamber. Associated anomalies include abnormal great artery positional relationships, pulmonic valvular or subvalvular stenosis, and subaortic stenosis. Survival to adulthood depends on a relatively normal pulmonary blood flow, yet normal pulmonary resistance and good ventricular function. Modifications of the Fontan approach are generally applied to carefully selected patients with creation of a pathway(s) from the systemic veins to the pulmonary arteries.

TRICUSPID ATRESIA

This malformation is characterized by atresia of the tricuspid valve; an interatrial communication; and, frequently, hypoplasia of the RV and pulmonary artery. The clinical picture is usually dominated by severe cyanosis due to obligatory admixture of systemic and pulmonary venous blood in the LV. The ECG characteristically shows RA enlargement, left-axis deviation, and LV hypertrophy.

Atrial septostomy and palliative operations to increase pulmonary blood flow, often by anastomosis of a systemic artery or vein to a pulmonary artery, may allow survival to the second or third decade. A Fontan atriopulmonary or total cavopulmonary connection may then allow functional correction in patients with normal or low pulmonary arterial resistance pressure and good LV function. There are a number of important long-term considerations with the Fontan circulation, including the development of arrhythmias, progressive liver dysfunction, thromboembolic complications, and potential long-term need for heart or heart and liver transplantation.

EBSTEIN'S ANOMALY

Characterized by a downward displacement of the tricuspid valve into the RV, due to anomalous attachment of the tricuspid leaflets, the Ebstein tricuspid valve tissue is dysplastic and results in tricuspid regurgitation. The abnormally situated tricuspid orifice produces an “atrialized” portion of the RV lying between the atrioventricular ring and the origin of the valve, which is continuous with the RA chamber. Often, the RV is hypoplastic. Although the clinical manifestations are variable, some patients come to initial attention because of either (1) progressive cyanosis from right-to-left atrial shunting, (2) symptoms due to tricuspid regurgitation and RV dysfunction, or (3) paroxysmal atrial tachyarrhythmias with or without atrioventricular bypass tracts (Wolff-Parkinson-White [WPW] syndrome). Diagnostic findings by two-dimensional echocardiography include the abnormal positional relation between the tricuspid and mitral valves with abnormally increased apical displacement of the septal tricuspid leaflet. Tricuspid regurgitation is quantitated by Doppler examination. Surgical approaches include prosthetic replacement of the tricuspid valve when the leaflets are tethered or repair of the native valve.

The two fundamental anatomic abnormalities in this malformation are transposition of the ascending aorta and pulmonary trunk and inversion of the ventricles. This arrangement results in desaturated systemic venous blood passing from the RA through the mitral valve to the LV and into the pulmonary trunk, whereas oxygenated pulmonary venous blood flows from the left atrium through the tricuspid valve to the RV and into the aorta. Thus, the circulation is corrected functionally. The clinical presentation, course, and prognosis of patients with congenitally corrected transposition vary depending on the nature and severity of any complicating intracardiac anomalies and of development of dysfunction of the systemic subaortic RV. Progressive RV dysfunction and tricuspid regurgitation may also develop in one-third of patients by age 30; Ebstein-type anomalies of the left-side tricuspid atrioventricular valve are common. VSD or PS due to obstruction to outflow from the right-sided subpulmonary (anatomic left) ventricle may coexist. Complete heart block occurs at a rate of 2–10% per decade. The diagnosis of the malformation and associated lesions can be established by comprehensive two-dimensional echocardiography and Doppler examination.

MALPOSITIONS OF THE HEART

Positional anomalies refer to conditions in which the cardiac apex is in the right side of the chest (*dextrocardia*) or at the midline (*mesocardia*), or in which there is a normal location of the heart in the left side of the chest but abnormal position of the viscera (*isolated levocardia*). Knowledge of the position of the abdominal organs and of the branching pattern of the main stem bronchi is important in categorizing these malpositions. When dextrocardia occurs without situs inversus, when the visceral situs is indeterminate, or if isolated levocardia is present, associated, often complex, multiple cardiac anomalies are usually present. In contrast, mirror-image dextrocardia is usually observed with complete situs inversus, which occurs most frequently in individuals whose hearts are otherwise normal.

SURGICALLY MODIFIED CONGENITAL HEART DISEASE

Owing to the enormous strides in cardiovascular surgical techniques that have occurred in the past 70 years, a large number of long-term survivors of palliative or corrective operations in infancy and childhood have reached adulthood. These patients are often challenging because of the diversity of anatomic, hemodynamic, and electrophysiologic residua and sequelae of cardiac operations.

The proper care of the survivor of an operation for CHD requires that the clinician understand the details of the malformation before operation; pay meticulous attention to the details of the operative procedure; and recognize the postoperative residua (conditions left totally or partially uncorrected), the sequelae (conditions caused by surgery), and the complications that may have resulted from the operation. Except for ligation of an uncomplicated patent ductus arteriosus, almost every other surgical repair leaves behind or causes some abnormality of the heart and circulation that may range from trivial to serious. Thus, even with results that are considered clinically to be good to excellent, continued long-term postoperative follow-up is advisable.

Cardiac operations involving the atria, such as closure of ASD, repair of total or partial anomalous pulmonary venous return, or venous switch corrections of complete transposition of the great arteries (the Mustard or Senning operations), may be followed years later by sinus node or atrioventricular node dysfunction and/or by atrial arrhythmias (especially atrial flutter). Intraventricular surgery may also result in electrophysiologic consequences, including complete heart block necessitating pacemaker insertion to avoid sudden death. Valvular problems may arise late after initial cardiac operation. An example is the progressive stenosis of an initially nonobstructive bicuspid aortic valve in the patient who underwent aortic coarctation repair. Such aortic valves may also be the site of infective endocarditis. After repair of the ostium primum ASD, the cleft mitral valve may become progressively regurgitant. Tricuspid regurgitation may also be progressive in the postoperative patient with tetralogy of Fallot

if RV outflow tract obstruction was not relieved adequately at initial surgery. In many patients with surgically modified CHD, inadequate relief of an obstructive lesion, a residual regurgitant lesion, or a residual shunt will cause or hasten the onset of clinical signs and symptoms of myocardial dysfunction. Despite a good hemodynamic repair, many patients with a subaortic RV develop RV decompensation and signs of left heart failure. In many patients, particularly those who were cyanotic for many years before operation, a preexisting compromise in ventricular performance is due to the original underlying malformation.

A final category of postoperative problems involves the use of prosthetic valves, patches, or conduits in the operative repair. The special risks include infective endocarditis, thrombus formation, and premature degeneration and calcification of the prosthetic materials. There are many patients in whom extracardiac conduits are required to correct the circulation functionally and often to carry blood to the lungs from the RA or RV. These conduits may develop intraluminal obstruction, and if they include a prosthetic valve, it may show progressive calcification and thickening. Many such patients face reintervention (interventional cardiac catheterization or surgical reoperation) one or more times in their lives. Such care should be directed to centers specializing in adults with complex congenital cardiovascular malformations. The effect of pregnancy in postoperative patients depends on the outcome of the repair, including the presence and severity of residua, sequelae, or complications. Contraception is an important topic with such patients. Tubal ligation should be considered in those in whom pregnancy is strictly contraindicated.

Endocarditis Prophylaxis Two major predisposing causes of infective endocarditis are a susceptible cardiovascular substrate and a source of bacteremia. The clinical and bacteriologic profile of infective endocarditis in patients with CHD has changed with the advent of intracardiac surgery and of prosthetic devices. Prophylaxis includes both antimicrobial and hygienic measures. Meticulous dental and skin care are required. Routine antimicrobial prophylaxis is recommended for bacteremic dental procedures or instrumentation through an infected site in most patients with operated CHD, particularly if foreign material, such as a prosthetic valve, conduit, or surgically constructed shunt, is in place. In the case of patches, in the absence of a high-pressure patch leak, or transcatheter devices, prophylaxis is usually recommended for 6 months until there is endothelialization. Individuals with unrepaired cyanotic heart disease are also generally recommended to receive prophylaxis (**Chap. 155**).

283

Aortic Valve Disease

Patrick T. O'Gara, Joseph Loscalzo

GLOBAL BURDEN OF VALVULAR HEART DISEASE



Primary valvular heart disease ranks well below coronary heart disease, stroke, hypertension, obesity, and diabetes as a major threat to the public health. Nevertheless, it is the source of significant morbidity and mortality rates. Rheumatic fever (**Chap. 381**) is the dominant cause of valvular heart disease in developing and low-income countries. Its prevalence has been estimated to range from as low as 1 per 100,000 school-age children in Costa Rica to as high as 150 per 100,000 in China. Rheumatic heart disease accounts for 12–65% of hospital admissions related to cardiovascular disease and 2–10% of hospital discharges in some developing countries. Prevalence and mortality rates vary among communities even within the same country as a function of overcrowding and the availability of medical resources and population-wide programs for detection and treatment of group A streptococcal pharyngitis. In economically deprived areas, tropical and subtropical climates (particularly on the Indian subcontinent), Central

America, and the Middle East, rheumatic valvular disease progresses more rapidly than in more-developed nations and frequently causes serious symptoms in patients younger than 20 years of age. This accelerated natural history may be due to repeated infections with more virulent strains of rheumatogenic streptococci. Approximately 15 million to 20 million people live with rheumatic heart disease worldwide, an estimated prevalence characterized by 300,000 new cases and 233,000 case fatalities per year, with the highest mortality rates reported from Southeast Asia (~7.6 per 100,000).

Although there have been recent reports of isolated outbreaks of streptococcal infection in North America, valve disease in high-income countries is dominated by degenerative or inflammatory processes that lead to valve thickening, calcification, and dysfunction. The prevalence of valvular heart disease increases with age for both men and women. Important left-sided valve disease may affect as many as 12–13% of adults older than the age of 75. In the United States, there were 85,000 hospital discharges with valvular heart disease in 2010, and the vast majority of these were related to surgical procedures for heart valve disease (mostly involving the aortic and mitral valves).

The incidence of infective endocarditis (Chap. 155) has increased with the aging of the population, the more widespread prevalence of vascular grafts and intracardiac devices, the emergence of more virulent multidrug-resistant microorganisms, and the growing epidemic of diabetes. The more restricted use of antibiotic prophylaxis since 2007 has thus far not been associated with an increase in incidence rates. Infective endocarditis has become a relatively more frequent cause of acute valvular regurgitation.

Bicuspid aortic valve disease affects as many as 0.5–1.4% of the general population, with an associated incidence of aortopathy involving root or ascending aortic aneurysm disease or coarctation. An increasing number of childhood survivors of congenital heart disease present later in life with valvular dysfunction. The global burden of valvular heart disease is expected to progress.

As is true for many other chronic health conditions, disparities in access to and quality of care for patients with valvular heart disease have been well documented. Management decisions and outcome differences based on age, gender, race, and geography require educational efforts across all levels of providers.

The role of the physical examination in the evaluation of patients with valvular heart disease is also considered in Chaps. 51e and 267; of electrocardiography (ECG) in Chap. 268; of echocardiography and other noninvasive imaging techniques in Chap. 270e; and of cardiac catheterization and angiography in Chap. 272.

AORTIC STENOSIS

Aortic stenosis (AS) occurs in about one-fourth of all patients with chronic valvular heart disease; approximately 80% of adult patients with symptomatic, valvular AS are male.

ETIOLOGY AND PATHOGENESIS

(Table 283-1) AS in adults is due to degenerative calcification of the aortic cusps and occurs most commonly on a substrate of congenital disease (bicuspid aortic valve), chronic (trileaflet) deterioration, or previous rheumatic inflammation. A pathologic study of specimens removed at the time of aortic valve replacement for AS showed that 53% were bicuspid and 4% unicuspid. The process of aortic valve deterioration and calcification is not a passive one, but rather one that shares many features with vascular atherosclerosis, including endothelial dysfunction, lipid accumulation, inflammatory cell activation, cytokine release, and upregulation of several signaling pathways (Fig. 283-1). Eventually, valvular myofibroblasts differentiate phenotypically into osteoblasts and actively produce bone matrix proteins that allow for the deposition of calcium hydroxyapatite crystals. Genetic polymorphisms involving the vitamin D receptor, the estrogen receptor in postmenopausal women, interleukin 10, and apolipoprotein E4 have been linked to the development of calcific AS, and a strong familial clustering of cases has been reported from western France. Several traditional atherosclerotic risk factors have also been associated with the development and progression of calcific

TABLE 283-1 MAJOR CAUSES OF AORTIC VALVE DISEASE

Valve Lesion	Etiologies
Aortic stenosis	Congenital (bicuspid, unicuspid) Degenerative calcific Rheumatic fever Radiation
Aortic regurgitation	Valvular Congenital (bicuspid) Endocarditis Rheumatic fever Myxomatous (prolapse) Traumatic Syphilis Ankylosing spondylitis Root disease Aortic dissection Cystic medial degeneration Marfan's syndrome Bicuspid aortic valve Nonsyndromic familial aneurysm Aortitis Hypertension

AS, including low-density lipoprotein (LDL) cholesterol, lipoprotein a (Lp[a]), diabetes mellitus, smoking, chronic kidney disease, and the metabolic syndrome. The presence of aortic valve sclerosis (focal thickening and calcification of the leaflets not severe enough to cause obstruction) is associated with an excess risk of cardiovascular death and myocardial infarction (MI) among persons older than age 65. Approximately 30% of persons older than 65 years exhibit aortic valve sclerosis, whereas 2% exhibit frank stenosis.

Rheumatic disease of the aortic leaflets produces commissural fusion, sometimes resulting in a bicuspid-appearing valve. This condition, in turn, makes the leaflets more susceptible to trauma and ultimately leads to fibrosis, calcification, and further narrowing. By the time the obstruction to left ventricular (LV) outflow causes serious clinical disability, the valve is usually a rigid calcified mass, and careful examination may make it difficult or even impossible to determine the etiology of the underlying process. Rheumatic AS is almost always associated with involvement of the mitral valve and with aortic regurgitation. Mediastinal radiation can also result in late scarring, fibrosis, and calcification of the leaflets with AS.

BICUSPID AORTIC VALVE DISEASE

A bicuspid aortic valve (BAV) is the most common congenital heart valve defect and occurs in 0.5–1.4% of the population with a 2–4:1 male-to-female predominance. The inheritance pattern appears to be autosomal dominant with incomplete penetrance, although some have questioned an X-linked component as suggested by the prevalence of BAV disease among patients with Turner's syndrome. The prevalence of BAV disease among first-degree relatives of an affected individual is approximately 10%. A single gene defect to explain the majority of cases has not been identified, although a mutation in the *NOTCH1* gene has been described in some families. Abnormalities in endothelial nitric oxide synthase and NKX2.5 have been implicated as well. Medial degeneration with ascending aortic aneurysm formation occurs commonly among patients with BAV disease; aortic coarctation is less frequently encountered. Patients with BAV disease have larger aortas than patients with comparable tricuspid aortic valve disease. The aortopathy develops independent of the hemodynamic severity of the valve lesion and is a risk factor for aneurysm formation and/or dissection. A BAV can be a component of more complex congenital heart disease with or without other left heart obstructing lesions, as seen in Shone's complex.

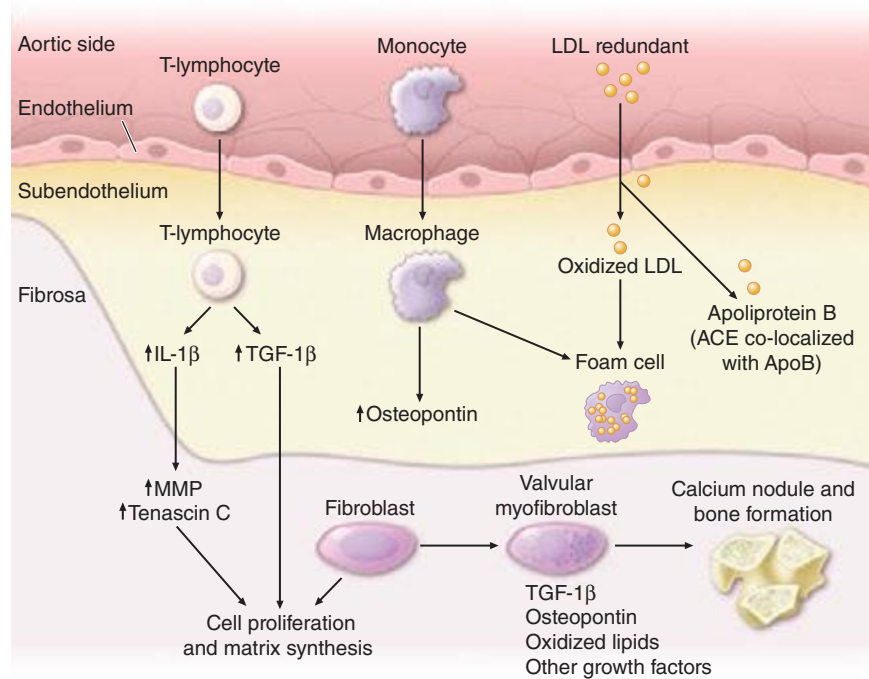


FIGURE 283-1 Pathogenesis of calcific aortic stenosis. Inflammatory cells infiltrate across the endothelial barrier and release cytokines that act on fibroblasts to promote cellular proliferation and matrix remodeling. LDL is oxidatively modified and taken up by macrophage scavengers to become foam cells. Angiotensin-converting enzyme colocalizes with ApoB. A subset of myofibroblasts differentiates into an osteoblast phenotype capable of promoting bone formation. ACE, angiotensin-converting enzyme; ApoB, apolipoprotein B; LDL, low-density lipoprotein; IL, interleukin; MMP, matrix metalloproteinase; TGF, transforming growth factor. (From RV Freeman, CM Otto: *Circulation* 111:3316, 2005; with permission.)

OTHER FORMS OF OBSTRUCTION TO LEFT VENTRICULAR OUTFLOW

In addition to valvular AS, three other lesions may be responsible for obstruction to LV outflow: *hypertrophic obstructive cardiomyopathy* (Chap. 287), *discrete fibromuscular/membranous subaortic stenosis*, and *supravalvular AS* (Chap. 282). The causes of LV outflow obstruction can be differentiated on the basis of the cardiac examination and Doppler echocardiographic findings.

PATHOPHYSIOLOGY

The obstruction to LV outflow produces a systolic pressure gradient between the LV and aorta. When severe obstruction is suddenly produced experimentally, the LV responds by dilation and reduction of stroke volume. However, in some patients, the obstruction may be present at birth and/or increase gradually over the course of many years, and LV contractile performance is maintained by the presence of concentric LV hypertrophy. Initially, this serves as an adaptive mechanism because it reduces toward normal the systolic stress developed by the myocardium, as predicted by the Laplace relation ($S = Pr/h$, where S = systolic wall stress, P = pressure, r = radius, and h = wall thickness). A large transaortic valve pressure gradient may exist for many years without a reduction in cardiac output (CO) or LV dilation; ultimately, however, excessive hypertrophy becomes maladaptive, LV systolic function declines because of afterload mismatch, abnormalities of diastolic function progress, and irreversible myocardial fibrosis develops.

A mean systolic pressure gradient >40 mmHg with a normal CO or an effective aortic orifice area of approximately <1 cm² (or approximately <0.6 cm²/m² body surface area in a normal-sized adult)—i.e., less than approximately one-third of the normal orifice area—is generally considered to represent severe obstruction to LV outflow. The elevated LV end-diastolic pressure observed in many patients with severe AS and preserved ejection fraction (EF) signifies the presence of diminished compliance of the hypertrophied LV. Although the CO at rest is within normal limits in most patients with severe AS, it usually fails to rise normally during exercise. Loss of an appropriately timed, vigorous atrial contraction, as occurs in atrial fibrillation (AF) or atrioventricular dissociation, may cause rapid progression of symptoms. Late in the course, contractile function deteriorates because of

afterload excess, the CO and LV–aortic pressure gradient decline, and the mean left atrial (LA), pulmonary artery (PA), and right ventricular (RV) pressures rise. LV performance can be further compromised by superimposed coronary artery disease (CAD). Stroke volume (and thus CO) can also be reduced in patients with significant hypertrophy and a small LV cavity despite a normal EF. Low-flow, low-gradient AS (with either reduced or normal LV systolic function) is both a diagnostic and therapeutic challenge.

The hypertrophied LV causes an increase in myocardial oxygen requirements. In addition, even in the absence of obstructive CAD, coronary blood flow is impaired to the extent that ischemia can be precipitated under conditions of excess demand. Capillary density is reduced relative to wall thickness, compressive forces are increased, and the elevated LV end-diastolic pressure reduces the coronary driving pressure. The subendocardium is especially vulnerable to ischemia by this mechanism.

SYMPTOMS

AS is rarely of clinical importance until the valve orifice has narrowed to approximately 1 cm². Even severe AS may exist for many years without producing any symptoms because of the ability of the hypertrophied LV to generate the elevated intraventricular pressures required to maintain a normal stroke volume. Once symptoms occur, valve replacement is indicated.

Most patients with pure or predominant AS have gradually increasing obstruction over years but do not become symptomatic until the sixth to eighth decades. Adult patients with BAV disease, however, develop significant valve dysfunction and symptoms one to two decades sooner. Exertional dyspnea, angina pectoris, and syncope are the three cardinal symptoms. Often, there is a history of insidious progression of fatigue and dyspnea associated with gradual curtailment of activities and reduced effort tolerance. *Dyspnea* results primarily from elevation of the pulmonary capillary pressure caused by elevations of LV diastolic pressures secondary to impaired relaxation and reduced LV compliance. *Angina pectoris* usually develops somewhat later and reflects an imbalance between the augmented myocardial oxygen requirements and reduced oxygen availability. CAD may or may not

be present, although its coexistence is common among AS patients older than age 65. *Exertional syncope* may result from a decline in arterial pressure caused by vasodilation in the exercising muscles and inadequate vasoconstriction in nonexercising muscles in the face of a fixed CO, or from a sudden fall in CO produced by an arrhythmia.

Because the CO at rest is usually well maintained until late in the course, marked fatigability, weakness, peripheral cyanosis, cachexia, and other clinical manifestations of a low CO are usually not prominent until this stage is reached. Orthopnea, paroxysmal nocturnal dyspnea, and pulmonary edema, i.e., symptoms of LV failure, also occur only in the advanced stages of the disease. Severe pulmonary hypertension leading to RV failure and systemic venous hypertension, hepatomegaly, AF, and tricuspid regurgitation (TR) are usually late findings in patients with isolated severe AS.

When AS and mitral stenosis (MS) coexist, the reduction in flow (CO) induced by MS lowers the pressure gradient across the aortic valve and, thereby, masks many of the clinical findings produced by AS. The transaortic pressure gradient can be increased in patients with concomitant aortic regurgitation (AR) due to higher aortic valve flow rates.

PHYSICAL FINDINGS

The rhythm is generally regular until late in the course; at other times, AF should suggest the possibility of associated mitral valve disease. The systemic arterial pressure is usually within normal limits. In the late stages, however, when stroke volume declines, the systolic pressure may fall and the pulse pressure narrow. The carotid arterial pulse rises slowly to a delayed peak (*pulsus parvus et tardus*). A thrill or anacrotic “shudder” may be palpable over the carotid arteries, more commonly the left. In the elderly, the stiffening of the arterial wall may mask this important physical sign. In many patients, the *a* wave in the jugular venous pulse is accentuated. This results from the diminished distensibility of the RV cavity caused by the bulging, hypertrophied interventricular septum.

The LV impulse is sometimes displaced laterally in the later stages of the disease. A double apical impulse (with a palpable S_4) may be recognized, particularly with the patient in the left lateral recumbent position. A systolic thrill may be present at the base of the heart to the right of the sternum when leaning forward or in the suprasternal notch.

Auscultation An early systolic ejection sound is frequently audible in children, adolescents, and young adults with congenital BAV disease. This sound usually disappears when the valve becomes calcified and rigid. As AS increases in severity, LV systole may become prolonged so that the aortic valve closure sound no longer precedes the pulmonic valve closure sound, and the two components may become synchronous, or aortic valve closure may even follow pulmonic valve closure, causing paradoxical splitting of S_2 (Chap. 267). The sound of aortic valve closure can be heard most frequently in patients with AS who have pliable valves, and calcification diminishes the intensity of this sound. Frequently, an S_4 is audible at the apex and reflects the presence of LV hypertrophy and an elevated LV end-diastolic pressure; an S_3 generally occurs late in the course, when the LV dilates and its systolic function becomes severely compromised.

The murmur of AS is characteristically an ejection (mid) systolic murmur that commences shortly after the S_1 , increases in intensity to reach a peak toward the middle of ejection, and ends just before aortic valve closure. It is characteristically low-pitched, rough and rasping in character, and loudest at the base of the heart, most commonly in the second right intercostal space. It is transmitted upward along the carotid arteries. Occasionally it is transmitted downward and to the apex, where it may be confused with the systolic murmur of mitral regurgitation (MR) (Gallavardin effect). In almost all patients with severe obstruction and preserved CO, the murmur is at least grade III/VI. In patients with mild degrees of obstruction or in those with severe stenosis with heart failure and low CO in whom the stroke volume and, therefore, the transvalvular flow rate are reduced, the murmur may be relatively soft and brief.

LABORATORY EXAMINATION

ECG In most patients with severe AS, there is LV hypertrophy. In advanced cases, ST-segment depression and T-wave inversion (LV “strain”) in standard leads I and aVL and in the left precordial leads are evident. However, there is no close correlation between the ECG and the hemodynamic severity of obstruction, and the absence of ECG signs of LV hypertrophy does not exclude severe obstruction. Many patients with AS have systemic hypertension, which can also contribute to the development of hypertrophy.

Echocardiogram The key findings on TTE are thickening, calcification, and reduced systolic opening of the valve leaflets and LV hypertrophy. Eccentric closure of the aortic valve cusps is characteristic of congenitally bicuspid valves. TEE imaging can display the obstructed orifice extremely well, but it is not routinely required for accurate characterization of AS. The valve gradient and aortic valve area can be estimated by Doppler measurement of the transaortic velocity. Severe AS is defined by a valve area <1 cm², whereas moderate AS is defined by a valve area of 1–1.5 cm² and mild AS by a valve area of 1.5–2 cm². Aortic valve sclerosis, conversely, is accompanied by a jet velocity of less than 2.5 meters/s (peak gradient <25 mmHg). LV dilation and reduced systolic shortening reflect impairment of LV function. There is increasing experience with the use of longitudinal strain and strain rate to characterize earlier changes in LV systolic function, well before a decline in EF can be appreciated. Doppler indices of impaired diastolic function are frequently seen.

Echocardiography is useful for identifying coexisting valvular abnormalities; for differentiating valvular AS from other forms of LV outflow obstruction; and for measurement of the aortic root and proximal ascending aortic dimensions. These aortic measurements are particularly important for patients with BAV disease. Dobutamine stress echocardiography is useful for the evaluation of patients with AS and severe LV systolic dysfunction (low-flow, low-gradient, severe AS with reduced EF), in whom the severity of the AS can often be difficult to judge. Patients with severe AS (i.e., valve area <1 cm²) with a relatively low mean gradient (<40 mmHg) despite a normal EF (low-flow, low-gradient, severe AS with normal EF) are often hypertensive, and efforts to control their systemic blood pressure should be optimized before Doppler echocardiography is repeated. The use of dobutamine stress echocardiography in this setting is under investigation. When there is continued uncertainty regarding the severity of AS in patients with reduced CO, quantitative analysis of the amount of aortic valve calcium with chest computed tomography (CT) may be helpful.

Chest X-Ray The chest x-ray may show no or little overall cardiac enlargement for many years. Hypertrophy without dilation may produce some rounding of the cardiac apex in the frontal projection and slight backward displacement in the lateral view. A dilated proximal ascending aorta may be seen along the upper right heart border in the frontal view. Aortic valve calcification may be discernible in the lateral view, but is usually readily apparent on fluoroscopic examination or by echocardiography; the absence of valvular calcification on fluoroscopy in an adult suggests that severe valvular AS is *not* present. In later stages of the disease, as the LV dilates, there is increasing roentgenographic evidence of LV enlargement, pulmonary congestion, and enlargement of the LA, PA, and right heart chambers.

Catheterization Right and left heart catheterization for invasive assessment of AS is performed infrequently but can be useful when there is a discrepancy between the clinical and noninvasive findings. Concern has been raised that attempts to cross the aortic valve for measurement of LV pressures are associated with a risk of cerebral embolization. Catheterization is also useful in three distinct categories of patients: (1) *patients with multivalvular disease*, in whom the role played by each valvular deformity should be defined to aid in the planning of operative treatment; (2) *young, asymptomatic patients with noncalcific congenital AS*, to define the severity of obstruction to LV outflow, because operation or percutaneous aortic balloon valvuloplasty (PABV) may be indicated in these patients if severe AS is

1532 present, even in the absence of symptoms; and (3) *patients in whom it is suspected that the obstruction to LV outflow may not be at the level of the aortic valve* but rather at the sub- or supra- valvular level.

Coronary angiography is indicated to screen for CAD in appropriate patients with severe AS who are being considered for surgery. The incidence of significant CAD for which bypass grafting is indicated at the time of aortic valve replacement (AVR) exceeds 50% among adult patients.

NATURAL HISTORY

Death in patients with severe AS occurs most commonly in the seventh and eighth decades. Based on data obtained at postmortem examination in patients before surgical treatment became widely available, the average time to death after the onset of various symptoms was as follows: angina pectoris, 3 years; syncope, 3 years; dyspnea, 2 years; congestive heart failure, 1.5–2 years. Moreover, in >80% of patients who died with AS, symptoms had existed for <4 years. Among adults dying with valvular AS, sudden death, which presumably resulted from an arrhythmia, occurred in 10–20%; however, most sudden deaths occurred in patients who had previously been symptomatic. Sudden death as the first manifestation of severe AS is very uncommon (<1% per year) in asymptomatic adult patients. Calcific AS is a progressive disease, with an annual reduction in valve area averaging 0.1 cm² and annual increases in the peak jet velocity and mean valve gradient averaging 0.3 meters/s and 7 mmHg, respectively (Table 283-2).

TREATMENT AORTIC STENOSIS (FIG. 283-2)

MEDICAL TREATMENT

In patients with severe AS (valve area <1 cm²), strenuous physical activity and competitive sports should be avoided, even in the asymptomatic stage. Care must be taken to avoid dehydration and hypovolemia to protect against a significant reduction in CO. Medications used for the treatment of hypertension or CAD, including beta blockers and angiotensin-converting enzyme (ACE) inhibitors, are generally safe for asymptomatic patients with preserved LV systolic function. Nitroglycerin is helpful in relieving angina pectoris in patients with CAD. Retrospective studies have shown that patients with degenerative calcific AS who receive HMG-CoA reductase inhibitors (“statins”) exhibit slower progression of leaflet calcification and aortic valve area reduction than those who do not. However, randomized prospective studies with either high-dose atorvastatin or combination simvastatin/ezetimibe have failed to show a measurable effect on valve-related outcomes. The use of statin medications should continue to be driven by considerations regarding primary and secondary prevention of atherosclerotic cardiovascular disease (ASCVD) events. ACE inhibitors have not been studied prospectively for AS-related outcomes. The need for endocarditis prophylaxis is restricted to AS patients with a prior history of endocarditis.

SURGICAL TREATMENT

Asymptomatic patients with calcific AS and severe obstruction should be followed carefully for the development of symptoms and by serial echocardiograms for evidence of deteriorating LV function. Operation is indicated in patients with severe AS (valve

area <1 cm² or 0.6 cm²/m² body surface area) who are symptomatic, those who exhibit LV systolic dysfunction (EF <50%), and those with BAV disease and an aneurysmal root or ascending aorta (maximal dimension >5.5 cm). Operation for aneurysm disease is recommended at smaller aortic diameters (4.5–5.0 cm) for patients with a family history of an aortic catastrophe and for patients who exhibit rapid aneurysm growth (>0.5 cm/year). Patients with asymptomatic moderate or severe AS who are referred for coronary artery bypass grafting surgery should also have AVR. In patients without heart failure, the operative risk of AVR (including patients with AS or AR) is approximately 2% (Table 283-2) but increases as a function of age and the need for concomitant aortic surgery or coronary revascularization with bypass grafting. The indications for AVR in the asymptomatic patient have been the subject of intense debate over the past 5 years, as surgical outcomes in selected patients have continued to improve. Relative indications for which surgery can be considered include an abnormal response to treadmill exercise; rapid progression of AS, especially when urgent access to medical care might be compromised; very severe AS, defined by an aortic valve jet velocity >5 meters/s or mean gradient >60 mmHg and low operative risk; and excessive LV hypertrophy in the absence of systemic hypertension. Exercise testing can be safely performed in the asymptomatic patient, as many as one-third of whom will show signs of functional impairment.

Operation should be carried out promptly after symptom onset. In patients with low-flow, low-gradient severe AS with reduced LVEF, the perioperative mortality risk is high (15–20%), and evidence of myocardial disease may persist even when the operation is technically successful. Long-term postoperative survival correlates with preoperative LV function. Nonetheless, in view of the even worse prognosis of such patients when they are treated medically, there is usually little choice but to advise valve replacement, especially in patients in whom contractile reserve can be demonstrated by dobutamine stress echocardiography (defined by a ≥20% increase in stroke volume after dobutamine challenge). Patients in this high surgical risk group may benefit from transcatheter aortic valve replacement (TAVR, see below). The treatment of patients with low-flow, low-gradient severe AS with normal LVEF is also difficult. Outcomes appear to be better with surgery compared with conservative medical care for symptomatic patients with this type of “paradoxical” low-flow AS, but more research is needed to guide therapeutic decision-making. In patients in whom severe AS and CAD coexist, relief of the AS and revascularization may sometimes result in striking clinical and hemodynamic improvement (Table 283-2).

Because many patients with calcific AS are elderly, particular attention must be directed to the adequacy of hepatic, renal, and pulmonary function before AVR is recommended. Age alone is not a contraindication to AVR for AS. The perioperative mortality rate depends to a substantial extent on the patient’s preoperative clinical and hemodynamic state. Treatment decisions for AS patients who are not at low operative risk should be made by a multidisciplinary heart team with representation from general cardiology, interventional cardiology, imaging, cardiac surgery, and other allied specialties as needed, including geriatrics. The 10-year survival rate of older adult patients with AVR is approximately 60%. Approximately 30% of bioprosthetic valves evidence primary valve failure in 10 years, requiring re-replacement, and an approximately equal percentage of patients with mechanical prostheses develop significant hemorrhagic complications as a consequence of treatment with vitamin K antagonists. Homograft AVR is usually reserved for patients with aortic valve endocarditis.

The Ross procedure involves replacement of the diseased aortic valve with the autologous pulmonic valve and implantation of a homograft in the native pulmonic position. Its use has declined considerably in the United States because of the technical complexity of the procedure and the incidence of late postoperative aortic root dilation and autograft failure with AR. There is also a low incidence of pulmonary homograft stenosis.

TABLE 283-2 MORTALITY RATES AFTER AORTIC VALVE SURGERY^a

Operation	Number	Unadjusted Operative Mortality (%)
AVR (isolated)	14,795	2.3
AVR + CAB	9158	4.2
AVR + MVR	876	8.8

^aData are for the first two quarters of calendar year 2013, during which 1004 sites reported a total of 135,666 procedures. Data are available from the Society of Thoracic Surgeons at http://www.sts.org/sites/default/files/documents/2013_3rdHarvestExecutiveSummary.pdf.

Abbreviations: AVR, aortic valve replacement; CAB, coronary artery bypass; MVR, mitral valve replacement.

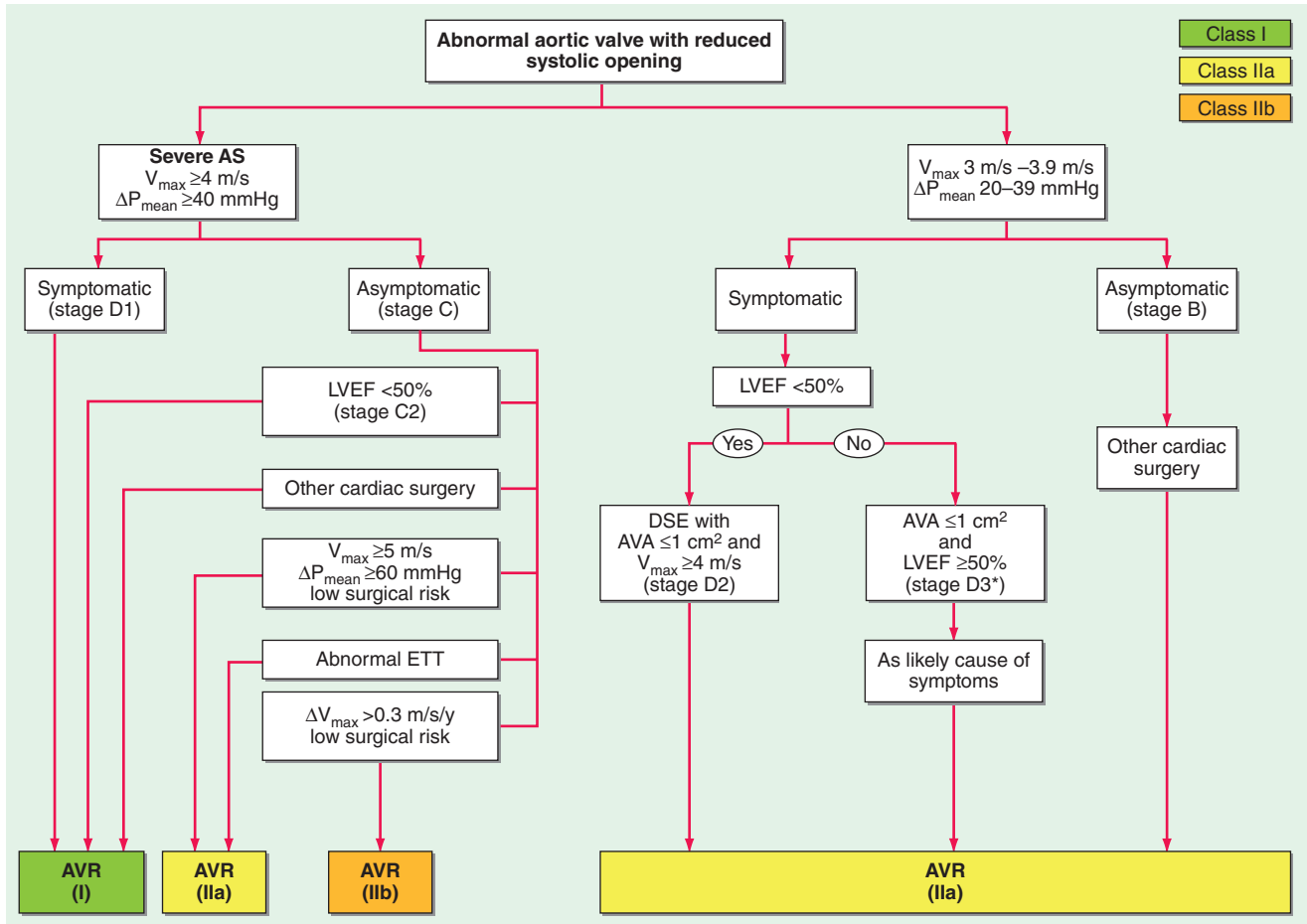


FIGURE 283-2 Management strategy for patients with aortic stenosis. Preoperative coronary angiography should be performed routinely as determined by age, symptoms, and coronary risk factors. Cardiac catheterization and angiography may also be helpful when there is a discrepancy between clinical and noninvasive findings. Patients who do not meet criteria for intervention should be monitored periodically with clinical and echocardiographic follow-up. The class designations refer to the American College of Cardiology/American Heart Association methodology for treatment recommendations. Class I recommendations should be performed or are indicated; Class IIa recommendations are considered reasonable to perform; Class IIb recommendations may be considered. The stages refer to the stages of progression of the disease. At disease stage A, risk factors are present for the development of valve dysfunction; stage B refers to progressive, mild-moderate, asymptomatic valve disease; stage C disease is severe in nature but clinically asymptomatic; stage C1 characterizes asymptomatic patients with severe valve disease but compensated ventricular function; stage C2 refers to asymptomatic, severe disease with ventricular decompensation; stage D refers to severe, symptomatic valve disease. With aortic stenosis, stage D1 refers to symptomatic patients with severe aortic stenosis and a high valve gradient (>40 mmHg mean gradient); stage D2 comprises patients with symptomatic, severe, low-flow, low-gradient aortic stenosis and low left ventricular ejection fraction; and stage D3 characterizes patients with symptomatic, severe, low-flow, low-gradient aortic stenosis and preserved left ventricular ejection fraction (paradoxical, low-flow, low-gradient severe aortic stenosis). AS, aortic stenosis; AVA, aortic valve area; AVR, aortic valve replacement by either surgical or transcatheter approach; BP, blood pressure; DSE, dobutamine stress echocardiography; ETT, exercise treadmill test; LVEF, left ventricular ejection fraction; ΔP_{mean} , mean pressure gradient; and V_{max} , maximum velocity. (Adapted from RA Nishimura et al: 2014 AHA/ACC Guideline for the Management of Patients with Valvular Heart Disease. *J Am Coll Cardiol* doi: 10.1016/j.jacc.2014.02.536, 2014, with permission.)

PERCUTANEOUS AORTIC BALLOON VALVULOPLASTY (PABV)

This procedure is preferable to operation in many children and young adults with congenital, noncalcific AS (Chap. 282). It is not commonly used as definitive therapy in adults with severe calcific AS because of a very high restenosis rate (80% within 1 year) and the risk of procedural complications, but on occasion, it has been used successfully as a “bridge to operation” in patients with severe LV dysfunction and shock who are too ill to tolerate surgery. It is performed routinely as part of the TAVR procedure (see below).

TRANSCATHETER AORTIC VALVE REPLACEMENT

TAVR for treatment of AS has been performed in more than 50,000 prohibitive- or high-surgical-risk adult patients worldwide using one of two available systems, a balloon-expandable valve and a self-expanding valve, both of which incorporate a pericardial prosthesis (Fig. 283-3). More than 250 U.S. centers now offer this procedure.

TAVR is most frequently performed via the transfemoral route, although trans-LV apical, subclavian, carotid, and ascending aortic routes have been used. Aortic balloon valvuloplasty under rapid RV pacing is performed as a first step to create an orifice of sufficient size for the prosthesis. Procedural success rates exceed 90%. Among elderly patients with severe AS who are considered inoperable (i.e., prohibitive surgical risk), 1- and 2-year survival rates are significantly higher with TAVR compared with medical therapy (including PABV) (Fig. 283-4). One- and 2-year survival rates are essentially equal for high-surgical-risk patients treated with TAVR or surgical AVR (SAVR) (Fig. 283-5). TAVR is associated with an early hazard for stroke and a higher incidence of postprocedural, paravalvular AR, a risk factor for mortality over the next 2 years. Postprocedural heart block requiring permanent pacemaker therapy is observed significantly more frequently with the self-expanding valve. Valve performance characteristics are excellent. Overall outcomes with this



FIGURE 283-3 Balloon-expandable (A) and self-expanding (B) valves for transcatheter aortic valve replacement (TAVR). B, inflated balloon; N, nose cone; V, valve. (Part A, courtesy of Edwards Lifesciences, Irvine, CA; with permission. NovaFlex+ is a trademark of Edwards Lifesciences Corporation. Part B, © Medtronic, Inc. 2015. Medtronic CoreValve Transcatheter Aortic Valve. CoreValve is a registered trademark of Medtronic, Inc.)

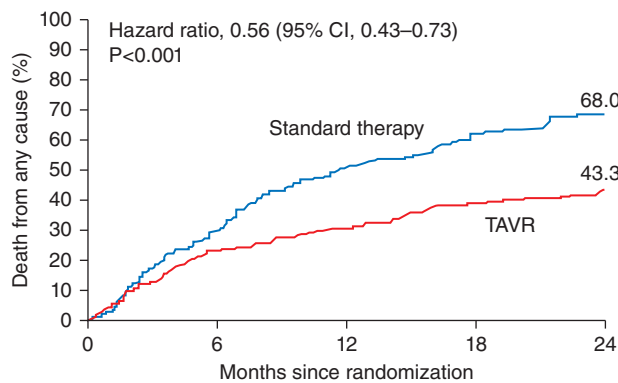
transformative technology have been very favorable and have allowed the extension of AVR to groups of patients previously considered at high or prohibitive risk for conventional surgery. Nevertheless, some patients are not candidates for this procedure because their comorbidity profile, including an assessment of frailty, would make its undertaking inappropriate. The heart team is specifically charged with making challenging decisions of this nature. The use of these devices for the treatment of patients at intermediate operative risk and for those with structural deterioration of bioprosthetic aortic and mitral valves (“valve-in-valve”), as an alternative to reoperative valve replacement, is under active study.

AORTIC REGURGITATION

ETIOLOGY

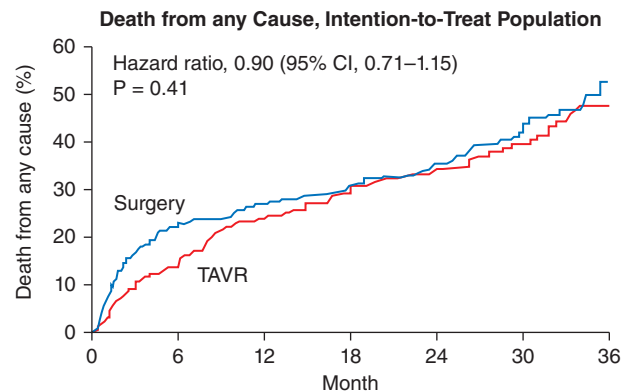
(Table 283-1) AR may be caused by primary valve disease or by primary aortic root disease.

Primary Valve Disease Rheumatic disease results in thickening, deformity, and shortening of the individual aortic valve cusps, changes that prevent their proper opening during systole and closure during diastole. A rheumatic origin is much less common in patients with isolated AR who do not have associated rheumatic mitral valve disease. Patients with congenital BAV disease may develop predominant AR,



No. at Risk					
TAVR	179	138	124	110	83
Standard therapy	179	121	85	62	42

FIGURE 283-4 Twenty-four-month outcomes following transcatheter aortic valve replacement (TAVR) for inoperable patients in the PARTNER I trial (cohort B). CI, confidence interval. (Adapted from RR Makkar et al: *N Engl J Med* 366:1696, 2012; with permission.)



No. at Risk							
TAVR	348	298	260	234	172	70	31
Surgery	351	252	236	217	165	65	32

FIGURE 283-5 Thirty-six-month outcomes following transcatheter aortic valve replacement (TAVR) for high-surgical-risk patients (cohort A) in the PARTNER I trial. CI, confidence interval. (Adapted from SK Kodali et al: *New Engl J Med* 366:1686, 2012.)

and approximately 20% of patients will require aortic valve surgery between 10 and 40 years of age. Congenital fenestrations of the aortic valve occasionally produce mild AR. Membranous subaortic stenosis often leads to thickening and scarring of the aortic valve leaflets with secondary AR. Prolapse of an aortic cusp, resulting in progressive chronic AR, occurs in approximately 15% of patients with ventricular septal defect (Chap. 282) but may also occur as an isolated phenomenon or as a consequence of myxomatous degeneration sometimes associated with mitral and/or tricuspid valve involvement.

AR may result from infective endocarditis, which can develop on a valve previously affected by rheumatic disease, a congenitally deformed valve, or on a normal aortic valve, and may lead to perforation or erosion of one or more leaflets. The aortic valve leaflets may become scarred and retracted during the course of syphilis or ankylosing spondylitis and contribute further to the AR that derives primarily from the associated root disease. Although traumatic rupture or avulsion of an aortic cusp is an uncommon cause of acute AR, it represents the most frequent serious lesion in patients surviving nonpenetrating cardiac injuries. The coexistence of hemodynamically significant AS with AR usually excludes all the rarer forms of AR because it occurs almost exclusively in patients with rheumatic or congenital AR. In patients with AR due to primary valvular disease, dilation of the aortic annulus may occur secondarily and lead to worsening regurgitation.

Primary Aortic Root Disease AR also may be due entirely to marked aortic annular dilation, i.e., aortic root disease, without primary involvement of the valve leaflets; widening of the aortic annulus and separation of the aortic leaflets are responsible for the AR (Chap. 301). Medial degeneration of the ascending aorta, which may or may not be associated with other manifestations of Marfan's syndrome; idiopathic dilation of the aorta; annuloaortic ectasia; osteogenesis imperfecta; and severe, chronic hypertension may all widen the aortic annulus and lead to progressive AR. Occasionally AR is caused by retrograde dissection of the aorta involving the aortic annulus. Syphilis and ankylosing spondylitis, both of which may affect the aortic leaflets, may also be associated with cellular infiltration and scarring of the media of the thoracic aorta, leading to aortic dilation, aneurysm formation, and severe regurgitation. In syphilis of the aorta (Chap. 206), now a very rare condition, the involvement of the intima may narrow the coronary ostia, which in turn may be responsible for myocardial ischemia.

PATHOPHYSIOLOGY

The total stroke volume ejected by the LV (i.e., the sum of the effective forward stroke volume and the volume of blood that regurgitates back into the LV) is increased in patients with AR. In patients with severe AR, the volume of regurgitant flow may equal the effective forward stroke volume. In contrast to MR, in which a portion of the LV stroke volume is delivered into the low-pressure LA, in AR the entire LV stroke volume is ejected into a high-pressure zone, the aorta. An increase in the LV end-diastolic volume (increased preload) constitutes the major hemodynamic compensation for AR. The dilation and eccentric hypertrophy of the LV allow this chamber to eject a larger stroke volume without requiring any increase in the relative shortening of each myofibril. Therefore, severe AR may occur with a normal effective forward stroke volume and a normal LVEF (total [forward plus regurgitant] stroke volume/end-diastolic volume), together with an elevated LV end-diastolic pressure and volume. However, through the operation of Laplace's law, LV dilation increases the LV systolic tension required to develop any given level of systolic pressure. Chronic AR is, thus, a state in which LV preload and afterload are both increased. Ultimately, these adaptive measures fail. As LV function deteriorates, the end-diastolic volume rises further and the forward stroke volume and EF decline. Deterioration of LV function often precedes the development of symptoms. Considerable thickening of the LV wall also occurs with chronic AR, and at autopsy, the hearts of these patients may be among the largest encountered, sometimes weighing >1000 g.

The reverse pressure gradient from aorta to LV, which drives the AR flow, falls progressively during diastole, accounting for the decrescendo

nature of the diastolic murmur. Equilibration between aortic and LV pressures may occur toward the end of diastole in patients with chronic severe AR, particularly when the heart rate is slow. In patients with acute severe AR, the LV is unprepared for the regurgitant volume load. LV compliance is normal or reduced, and LV diastolic pressures rise rapidly, occasionally to levels >40 mmHg. The LV pressure may exceed the LA pressure toward the end of diastole, and this reversed pressure gradient closes the mitral valve prematurely.

In patients with chronic severe AR, the effective forward CO usually is normal or only slightly reduced at rest, but often it fails to rise normally during exertion. An early sign of LV dysfunction is a reduction in the EF. In advanced stages, there may be considerable elevation of the LA, PA wedge, PA, and RV pressures and lowering of the forward CO at rest.

Myocardial ischemia may occur in patients with AR because myocardial oxygen requirements are elevated by LV dilation, hypertrophy, and elevated LV systolic tension, and coronary blood flow may be compromised. A large fraction of coronary blood flow occurs during diastole, when arterial pressure is low, thereby reducing coronary perfusion or driving pressure. This combination of increased oxygen demand and reduced supply may cause myocardial ischemia, particularly of the subendocardium, even in the absence of epicardial CAD.

HISTORY

Approximately three-fourths of patients with pure or predominant valvular AR are men; women predominate among patients with primary valvular AR who have associated rheumatic mitral valve disease. A history compatible with infective endocarditis may sometimes be elicited from patients with rheumatic or congenital involvement of the aortic valve, and the infection often precipitates or seriously aggravates preexisting symptoms.

In patients with *acute severe AR*, as may occur in infective endocarditis, aortic dissection, or trauma, the LV cannot dilate sufficiently to maintain stroke volume, and LV diastolic pressure rises rapidly with associated marked elevations of LA and PA wedge pressures. Pulmonary edema and/or cardiogenic shock may develop rapidly.

Chronic severe AR may have a long latent period, and patients may remain relatively asymptomatic for as long as 10–15 years. However, uncomfortable awareness of the heartbeat, especially on lying down, may be an early complaint. Sinus tachycardia, during exertion or with emotion, or premature ventricular contractions may produce particularly uncomfortable palpitations as well as head pounding. These complaints may persist for many years before the development of exertional dyspnea, usually the first symptom of diminished cardiac reserve. The dyspnea is followed by orthopnea, paroxysmal nocturnal dyspnea, and excessive diaphoresis. Anginal chest pain even in the absence of CAD may occur in patients with severe AR, even in younger patients. Anginal pain may develop at rest as well as during exertion. Nocturnal angina may be a particularly troublesome symptom, and it may be accompanied by marked diaphoresis. The anginal episodes can be prolonged and often do not respond satisfactorily to sublingual nitroglycerin. Systemic fluid accumulation, including congestive hepatomegaly and ankle edema, may develop late in the course of the disease.

PHYSICAL FINDINGS

In chronic severe AR, the jarring of the entire body and the bobbing motion of the head with each systole can be appreciated, and the abrupt distention and collapse of the larger arteries are easily visible. The examination should be directed toward the detection of conditions predisposing to AR, such as bicuspid valve, endocarditis, Marfan's syndrome, and ankylosing spondylitis.

Arterial Pulse A rapidly rising “water-hammer” pulse, which collapses suddenly as arterial pressure falls rapidly during late systole and diastole (Corrigan's pulse), and capillary pulsations, an alternate flushing and paling of the skin at the root of the nail while pressure is applied to the tip of the nail (Quincke's pulse), are characteristic

1536 of chronic severe AR. A booming “pistol-shot” sound can be heard over the femoral arteries (Traube’s sign), and a to-and-fro murmur (Duroziez’s sign) is audible if the femoral artery is lightly compressed with a stethoscope.

The arterial pulse pressure is widened as a result of both systolic hypertension and a lowering of the diastolic pressure. The measurement of arterial diastolic pressure with a sphygmomanometer may be complicated by the fact that systolic sounds are frequently heard with the cuff completely deflated. However, the level of cuff pressure at the time of muffling of the Korotkoff sounds (phase IV) generally corresponds fairly closely to the true intraarterial diastolic pressure. As the disease progresses and the LV end-diastolic pressure rises, the arterial diastolic pressure may actually rise as well, because the aortic diastolic pressure cannot fall below the LV end-diastolic pressure. For the same reason, acute severe AR may also be accompanied by only a slight widening of the pulse pressure. Such patients are invariably tachycardic as the heart rate increases in an attempt to preserve the CO.

Palpation In patients with chronic severe AR, the LV impulse is heaving and displaced laterally and inferiorly. The systolic expansion and diastolic retraction of the apex are prominent. A diastolic thrill may be palpable along the left sternal border in thin-chested individuals, and a prominent systolic thrill may be palpable in the suprasternal notch and transmitted upward along the carotid arteries. This systolic thrill and the accompanying murmur do not necessarily signify the coexistence of AS. In some patients with AR or with combined AS and AR, the carotid arterial pulse may be bisferiens, i.e., with two systolic waves separated by a trough (see Fig. 267-2D).

Auscultation In patients with severe AR, the aortic valve closure sound (A_2) is usually absent. A systolic ejection sound is audible in patients with BAV disease, and occasionally an S_4 also may be heard. The murmur of chronic AR is typically a high-pitched, blowing, decrescendo diastolic murmur, heard best in the third intercostal space along the left sternal border (see Fig. 267-5B). In patients with mild AR, this murmur is brief, but as the severity increases, it generally becomes louder and longer, indeed holodiastolic. When the murmur is soft, it can be heard best with the diaphragm of the stethoscope and with the patient sitting up, leaning forward, and with the breath held in forced expiration. In patients in whom the AR is caused by primary valvular disease, the diastolic murmur is usually louder along the left than the right sternal border. However, when the murmur is heard best along the right sternal border, it suggests that the AR is caused by aneurysmal dilation of the aortic root. “Cooing” or musical diastolic murmurs suggest eversion of an aortic cusp vibrating in the regurgitant stream.

A mid-systolic ejection murmur is frequently audible in isolated AR. It is generally heard best at the base of the heart and is transmitted along the carotid arteries. This murmur may be quite loud without signifying aortic obstruction. A third murmur sometimes heard in patients with severe AR is the *Austin Flint murmur*, a soft, low-pitched, rumbling mid-to-late diastolic murmur. It is probably produced by the diastolic displacement of the anterior leaflet of the mitral valve by the AR stream and is not associated with hemodynamically significant mitral obstruction. The auscultatory features of AR are intensified by strenuous and sustained handgrip, which augments systemic vascular resistance.

In acute severe AR, the elevation of LV end-diastolic pressure may lead to early closure of the mitral valve, a soft S_1 , a pulse pressure that is not particularly wide, and a soft, short, early diastolic murmur of AR.

LABORATORY EXAMINATION

ECG In patients with chronic severe AR, the ECG signs of LV hypertrophy become manifest (Chap. 268). In addition, these patients frequently exhibit ST-segment depression and T-wave inversion in leads I, aVL, V_5 , and V_6 (“LV strain”). Left-axis deviation and/or QRS prolongation denote diffuse myocardial disease, generally associated with patchy fibrosis, and usually signify a poor prognosis.

Echocardiogram LV size is increased in chronic AR and systolic function is normal or even supernormal until myocardial contractility declines, as signaled by a decrease in EF or increase in the end-systolic dimension. A rapid, high-frequency diastolic fluttering of the anterior mitral leaflet produced by the impact of the regurgitant jet is a characteristic finding. The echocardiogram is also useful in determining the cause of AR, by detecting dilation of the aortic annulus and root, aortic dissection (see Fig. 270e-5), or primary leaflet pathology. With severe AR, the central jet width assessed by color flow Doppler imaging exceeds 65% of the LV outflow tract, the regurgitant volume is ≥ 60 mL/beat, the regurgitant fraction is $\geq 50\%$, and there is diastolic flow reversal in the proximal descending thoracic aorta. The continuous-wave Doppler profile of the AR jet shows a rapid deceleration time in patients with acute severe AR, due to the rapid increase in LV diastolic pressure. Surveillance transthoracic echocardiography forms the cornerstone of longitudinal follow-up and allows for the early detection of changes in LV size and/or function. For patients in whom transthoracic echocardiography (TTE) is limited by poor acoustical windows or inadequate semiquantitative assessment of LV function or the severity of the regurgitation, cardiac magnetic resonance imaging (MRI) can be performed. This modality also allows for accurate assessment of aortic size and contour. Transesophageal echocardiography (TEE) can also provide detailed anatomic assessment of the valve, root, and portions of the aorta.

Chest X-Ray In chronic severe AR, the apex is displaced downward and to the left in the frontal projection. In the left anterior oblique and lateral projections, the LV is displaced posteriorly and encroaches on the spine. When AR is caused by primary disease of the aortic root, aneurysmal dilation of the aorta may be noted, and the aorta may fill the retrosternal space in the lateral view. Echocardiography, cardiac MRI, and chest CT angiography are more sensitive than the chest x-ray for the detection of root and ascending aortic enlargement.

Cardiac Catheterization and Angiography When needed, right and left heart catheterization with contrast aortography can provide confirmation of the magnitude of regurgitation and the status of LV function. Coronary angiography is performed routinely in appropriate patients prior to surgery.

TREATMENT AORTIC REGURGITATION

ACUTE AORTIC REGURGITATION (FIG. 283-6)

Patients with acute severe AR may respond to intravenous diuretics and vasodilators (such as sodium nitroprusside), but stabilization is usually short-lived and operation is indicated urgently. Intraaortic balloon counterpulsation is contraindicated. Beta blockers are also best avoided so as not to reduce the CO further or slow the heart rate, thus allowing more time for diastolic filling of the LV. Surgery is the treatment of choice and is usually necessary within 24 h of diagnosis.

CHRONIC AORTIC REGURGITATION

Early symptoms of dyspnea and effort intolerance respond to treatment with diuretics; vasodilators (ACE inhibitors, dihydropyridine calcium channel blockers, or hydralazine) may be useful as well. Surgery can then be performed in a more controlled setting. The use of vasodilators to extend the compensated phase of chronic severe AR before the onset of symptoms or the development of LV dysfunction is more controversial and less well established. Systolic blood pressure should be controlled (goal < 140 mmHg) in patients with chronic AR, and vasodilators are an excellent first choice as antihypertensive agents. It is often difficult to achieve adequate control because of the increased stroke volume that accompanies severe AR. Cardiac arrhythmias and systemic infections are poorly tolerated in patients with severe AR and must be treated promptly and vigorously. Although nitroglycerin and long-acting nitrates are not as helpful in relieving anginal pain as they are in patients with ischemic heart disease, they are worth a trial. Patients with

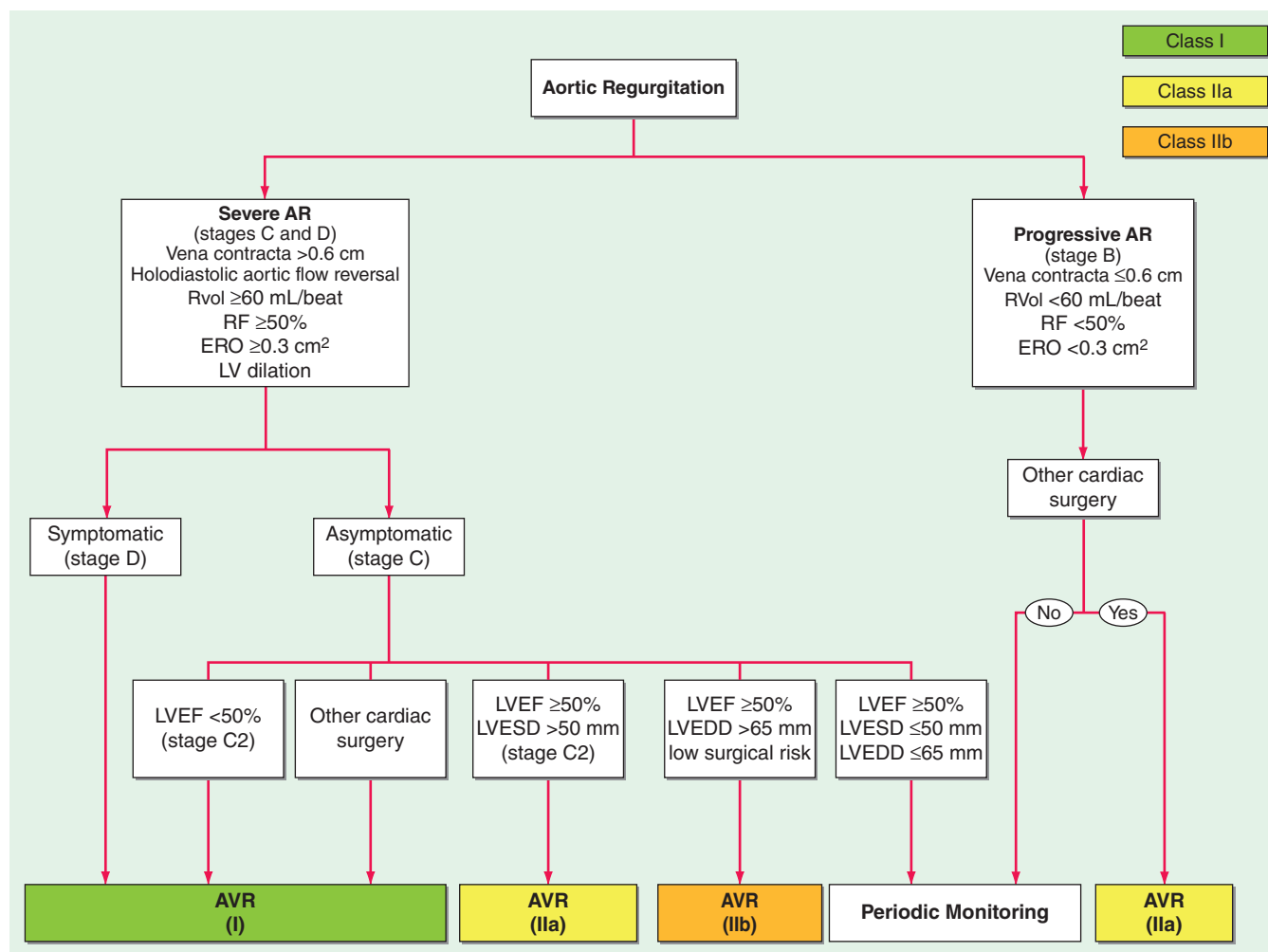


FIGURE 283-6 Management of patients with aortic regurgitation. See legend for Fig. 283-2 for explanation of treatment recommendations (Class I, IIa, and IIb) and disease stages (B, C1, C2, D). Preoperative coronary angiography should be performed routinely as determined by age, symptoms, and coronary risk factors. Cardiac catheterization and angiography may also be helpful when there is a discrepancy between clinical and noninvasive findings. Patients who do not meet criteria for intervention should be monitored periodically with clinical and echocardiographic follow-up. AR, aortic regurgitation; AVR, aortic valve replacement (valve repair may be appropriate in selected patients); ERO, effective regurgitant orifice; LV, left ventricular; LVEDD, left ventricular end-diastolic dimension; LVEF, left ventricular ejection fraction; LVESD, left ventricular end-systolic dimension; RF, regurgitant fraction; RVol, regurgitant volume. (Adapted from RA Nishimura et al: 2014 AHA/ACC Guideline for the Management of Patients with Valvular Heart Disease. *J Am Coll Cardiol* doi: 10.1016/j.jacc.2014.02.536, 2014, with permission.)

siphilitic aortitis should receive a full course of penicillin therapy (Chap. 206). Beta blockers and the angiotensin receptor blocker losartan may be useful to retard the rate of aortic root enlargement in young patients with Marfan's syndrome and aortic root dilation. Early reports of the efficacy of losartan in patients with Marfan's syndrome have led to its use in other populations of patients including those with BAV disease and aortopathy. The use of beta blockers in patients with valvular AR was previously felt to be relatively contraindicated due to concerns that the resulting slowing of the heart rate would allow more time for diastolic regurgitation. More recent observational reports, however, suggest that beta blockers may provide functional benefit in patients with chronic AR. Beta blockers can sometimes provide incremental blood pressure lowering in patients with chronic AR and hypertension. Patients with severe AR, particularly those with an associated aortopathy, should avoid isometric exercises.

SURGICAL TREATMENT

In deciding on the advisability and proper timing of surgical treatment, two points should be kept in mind: (1) patients with chronic severe AR usually do not become symptomatic until

after the development of myocardial dysfunction; and (2) when delayed too long (defined as >1 year from onset of symptoms or LV dysfunction), surgical treatment often does not restore normal LV function. Therefore, in patients with chronic severe AR, careful clinical follow-up and noninvasive testing with echocardiography at approximately 6- to 12-month intervals are necessary if operation is to be undertaken at the optimal time, i.e., after the onset of LV dysfunction but prior to the development of severe symptoms. Exercise testing may be helpful to assess effort tolerance more objectively. Operation can be deferred as long as the patient both remains asymptomatic and retains normal LV function without severe chamber dilation.

AVR is indicated for the treatment of severe AR in symptomatic patients irrespective of LV function. In general, the operation should be carried out in asymptomatic patients with severe AR and progressive LV dysfunction defined by an LVEF <50%, an LV end-systolic dimension >50 mm, or an LV diastolic dimension >65 mm. Smaller dimensions may be appropriate thresholds in individuals of smaller stature. Patients with severe AR without indications for operation should be followed by clinical and echocardiographic examination every 6–12 months.

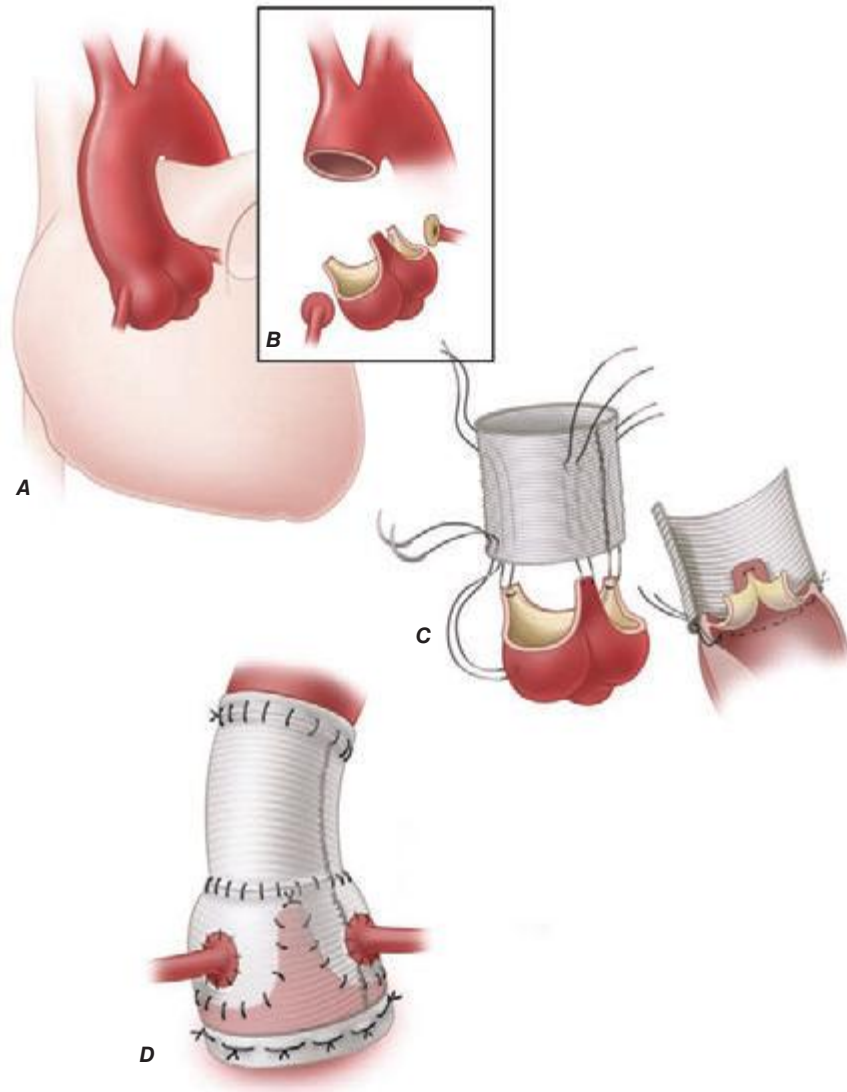


FIGURE 283-7 Valve-sparing aortic root reconstruction (David procedure). (From P Steltzer et al [eds]: *Valvular Heart Disease: A Companion to Braunwald's Heart Disease*, 3rd ed, Fig 12-27, p. 200.)

Surgical options for management of aortic valve and root disease have expanded considerably over the past decade. AVR with a suitable mechanical or tissue prosthesis is generally necessary in patients with rheumatic AR and in many patients with other forms of regurgitation. Rarely, when a leaflet has been perforated during infective endocarditis or torn from its attachments to the aortic annulus by thoracic trauma, primary surgical repair may be possible. When AR is due to aneurysmal dilation of the root or proximal ascending aorta rather than to primary valve involvement, it may be possible to reduce or eliminate the regurgitation by narrowing the annulus or by excising a portion of the aortic root without replacing the valve. Elective, valve-sparing aortic root reconstruction generally involves reimplantation of the valve in a contoured graft with reattachment of the coronary artery buttons into the side of the graft and is best undertaken in specialized surgical centers (Fig. 283-7). Resuspension of the native aortic valve leaflets is possible in approximately 50% of patients with acute AR in the setting of type A aortic dissection. In other conditions, however, regurgitation can be effectively eliminated only by replacing the aortic valve, the dilated or aneurysmal

ascending aorta responsible for the regurgitation, and implanting a composite valve-graft conduit. This formidable procedure entails a higher risk than isolated AVR.

As in patients with other valvular abnormalities, both the operative risk and the late mortality rate are largely dependent on the stage of the disease and myocardial function at the time of operation. The overall operative mortality rate for isolated AVR (performed for either or both AS or AR) is approximately 2% (Table 283-2). However, patients with AR, marked cardiac enlargement, and prolonged LV dysfunction experience an operative mortality rate of approximately 10% and a late mortality rate of approximately 5% per year due to LV failure despite a technically satisfactory operation. Nonetheless, because of the very poor prognosis with medical management, even patients with LV systolic failure should be considered for operation.

Patients with acute severe AR require prompt surgical treatment, which may be lifesaving.

The role of the physical examination in the evaluation of patients with valvular heart disease is also considered in **Chaps. 51e and 267**; of electrocardiography (ECG) in **Chap. 268**; of echocardiography and other noninvasive imaging techniques in **Chap. 270e**; and of cardiac catheterization and angiography in **Chap. 272**.

MITRAL STENOSIS

ETIOLOGY AND PATHOLOGY

Rheumatic fever is the leading cause of mitral stenosis (MS) (**Table 284-1**). Other less common etiologies of obstruction to left ventricular inflow include congenital mitral valve stenosis, cor triatriatum, mitral annular calcification with extension onto the leaflets, systemic lupus erythematosus, rheumatoid arthritis, left atrial myxoma, and infective endocarditis with large vegetations. Pure or predominant MS occurs in approximately 40% of all patients with rheumatic heart disease and a history of rheumatic fever (**Chap. 381**). In other patients with rheumatic heart disease, lesser degrees of MS may accompany mitral regurgitation (MR) and aortic valve disease. With reductions in the incidence of acute rheumatic fever, particularly in temperate climates and developed countries, the incidence of MS has declined considerably over the past several decades. However, it remains a major problem in developing nations, especially in tropical and semi-tropical climates.

In rheumatic MS, chronic inflammation leads to diffuse thickening of the valve leaflets with formation of fibrous tissue and/or calcific deposits. The mitral commissures fuse, the chordae tendineae fuse and shorten, the valvular cusps become rigid, and these changes, in turn, lead to narrowing at the apex of the funnel-shaped (“fish-mouth”) valve. Although the initial insult to the mitral valve is rheumatic, later changes may be exacerbated by a nonspecific process resulting from trauma to the valve due to altered flow patterns. Calcification of the

stenotic mitral valve immobilizes the leaflets and narrows the orifice further. Thrombus formation and arterial embolization may arise from the calcific valve itself, but in patients with atrial fibrillation (AF), thrombi arise more frequently from the dilated left atrium (LA), particularly from within the LA appendage.

PATHOPHYSIOLOGY

In normal adults, the area of the mitral valve orifice is 4–6 cm². In the presence of significant obstruction, i.e., when the orifice area is reduced to < ~2 cm², blood can flow from the LA to the left ventricle (LV) only if propelled by an abnormally elevated left atrioventricular pressure gradient, the hemodynamic hallmark of MS. When the mitral valve opening is reduced to <1.5 cm², referred to as “severe” MS, an LA pressure of ~25 mmHg is required to maintain a normal cardiac output (CO). The elevated pulmonary venous and pulmonary arterial (PA) wedge pressures reduce pulmonary compliance, contributing to exertional dyspnea. The first bouts of dyspnea are usually precipitated by clinical events that increase the rate of blood flow across the mitral orifice, resulting in further elevation of the LA pressure (see below).

To assess the severity of obstruction hemodynamically, both the transvalvular pressure gradient and the flow rate must be measured (**Chap. 272**). The latter depends not only on the CO but on the heart rate, as well. An increase in heart rate shortens diastole proportionately more than systole and diminishes the time available for flow across the mitral valve. Therefore, at any given level of CO, tachycardia, including that associated with rapid AF, augments the transvalvular pressure gradient and elevates further the LA pressure. Similar considerations apply to the pathophysiology of tricuspid stenosis.

The LV diastolic pressure and ejection fraction (EF) are normal in isolated MS. In MS and sinus rhythm, the elevated LA and PA wedge pressures exhibit a prominent atrial contraction pattern (*a* wave) and a gradual pressure decline after the *v* wave and mitral valve opening (*y* descent). In severe MS and whenever pulmonary vascular resistance is significantly increased, the PA pressure (PAP) is elevated at rest and rises further during exercise, often causing secondary elevations of right ventricular (RV) end-diastolic pressure and volume.

Cardiac Output In patients with severe MS (mitral valve orifice 1–1.5 cm²), the CO is normal or almost so at rest, but rises subnormally during exertion. In patients with very severe MS (valve area <1 cm²), particularly those in whom pulmonary vascular resistance is markedly elevated, the CO is subnormal at rest and may fail to rise or may even decline during activity.

Pulmonary Hypertension The clinical and hemodynamic features of MS are influenced importantly by the level of the PAP. Pulmonary hypertension results from: (1) passive backward transmission of the elevated LA pressure; (2) pulmonary arteriolar constriction (the so-called “second stenosis”), which presumably is triggered by LA and pulmonary venous hypertension (reactive pulmonary hypertension); (3) interstitial edema in the walls of the small pulmonary vessels; and (4) at end stage, organic obliterative changes in the pulmonary vascular bed. Severe pulmonary hypertension results in RV enlargement, secondary tricuspid regurgitation (TR), and pulmonic regurgitation (PR), as well as right-sided heart failure.

SYMPTOMS

In temperate climates, the latent period between the initial attack of rheumatic carditis (in the increasingly rare circumstances in which a history of one can be elicited) and the development of symptoms due to MS is generally about two decades; most patients begin to experience disability in the fourth decade of life. Studies carried out before the development of mitral valvotomy revealed that once a patient with MS became seriously symptomatic, the disease progressed inexorably to death within 2–5 years.

In patients whose mitral orifices are large enough to accommodate a normal blood flow with only mild elevations of LA pressure, marked elevations of this pressure leading to dyspnea and cough may be precipitated by sudden changes in the heart rate, volume status, or CO, as, for example, with severe exertion, excitement, fever, severe anemia, paroxysmal AF

TABLE 284-1 MAJOR CAUSES OF MITRAL VALVE DISEASE

Valve Lesion	Etiologies
Mitral stenosis	Rheumatic fever
	Congenital
	Severe mitral annular calcification SLE, RA
Mitral regurgitation	Acute
	Endocarditis
	Papillary muscle rupture (post-MI)
	Trauma
	Chordal rupture/leaflet flail (MVP, IE)
	Chronic
	Myxomatous (MVP)
	Rheumatic fever
	Endocarditis (healed)
	Mitral annular calcification
	Congenital (cleft, AV canal)
	HOCM with SAM
	Ischemic (LV remodeling)
Dilated cardiomyopathy	
Radiation	

Abbreviations: AV, atrioventricular; IE, infective endocarditis; HOCM, hypertrophic obstructive cardiomyopathy; LV, left ventricular; MI, myocardial infarction; MVP, mitral valve prolapse; RA, rheumatoid arthritis; SAM, systolic anterior motion; SLE, systemic lupus erythematosus.

and other tachycardias, sexual intercourse, pregnancy, and thyrotoxicosis. As MS progresses, lesser degrees of stress precipitate dyspnea, the patient becomes limited in daily activities, and orthopnea and paroxysmal nocturnal dyspnea develop. The development of persistent AF often marks a turning point in the patient's course and is generally associated with acceleration of the rate at which symptoms progress. *Hemoptysis* (Chap. 48) results from rupture of pulmonary-bronchial venous connections secondary to pulmonary venous hypertension. It occurs most frequently in patients who have elevated LA pressures without markedly elevated pulmonary vascular resistances and is rarely fatal. *Recurrent pulmonary emboli* (Chap. 300), sometimes with infarction, are an important cause of morbidity and mortality late in the course of MS. *Pulmonary infections*, i.e., bronchitis, bronchopneumonia, and lobar pneumonia, commonly complicate untreated MS, especially during the winter months.

Pulmonary Changes In addition to the aforementioned changes in the pulmonary vascular bed, fibrous thickening of the walls of the alveoli and pulmonary capillaries occurs commonly in MS. The vital capacity, total lung capacity, maximal breathing capacity, and oxygen uptake per unit of ventilation are reduced (Chap. 306e). Pulmonary compliance falls further as pulmonary capillary pressure rises during exercise.

Thrombi and Emboli *Thrombi* may form in the left atria, particularly within the enlarged atrial appendages of patients with MS. Systemic embolization, the incidence of which is 10–20%, occurs more frequently in patients with AF, in patients >65 years of age, and in those with a reduced CO. However, systemic embolization may be the presenting feature in otherwise asymptomatic patients with only mild MS.

PHYSICAL FINDINGS

(See also Chaps. 51e and 267)

Inspection and Palpation In patients with severe MS, there may be a malar flush with pinched and blue facies. In patients with sinus rhythm and severe pulmonary hypertension or associated tricuspid stenosis (TS), the jugular venous pulse reveals prominent *a* waves due to vigorous right atrial systole. The systemic arterial pressure is usually normal or slightly low. An RV tap along the left sternal border signifies an enlarged RV. A diastolic thrill may rarely be present at the cardiac apex, with the patient in the left lateral recumbent position.

Auscultation The first heart sound (S_1) is usually accentuated in the early stages of the disease and slightly delayed. The pulmonic component of the second heart sound (P_2) also is often accentuated with elevated PA pressures, and the two components of the second heart sound (S_2) are closely split. The opening snap (OS) of the mitral valve is most readily audible in expiration at, or just medial to, the cardiac apex. This sound generally follows the sound of aortic valve closure (A_2) by 0.05–0.12 s. The time interval between A_2 and OS varies inversely with the severity of the MS. The OS is followed by a low-pitched, rumbling, diastolic murmur, heard best at the apex with the patient in the left lateral recumbent position (see Fig. 267-5); it is accentuated by mild exercise (e.g., a few rapid sit-ups) carried out just before auscultation. In general, the duration of this murmur correlates with the severity of the stenosis in patients with preserved CO. In patients with sinus rhythm, the murmur often reappears or becomes louder during atrial systole (presystolic accentuation). Soft, grade I or II/VI systolic murmurs are commonly heard at the apex or along the left sternal border in patients with pure MS and do not necessarily signify the presence of MR. Hepatomegaly, ankle edema, ascites, and pleural effusion, particularly in the right pleural cavity, may occur in patients with MS and RV failure.

Associated Lesions With severe pulmonary hypertension, a pansystolic murmur produced by functional TR may be audible along the left sternal border. This murmur is usually louder during inspiration and diminishes during forced expiration (Carvallo's sign). When the CO is markedly reduced in MS, the typical auscultatory findings, including the diastolic rumbling murmur, may not be detectable (silent MS), but they may reappear as compensation is restored. The *Graham Steell murmur* of PR, a high-pitched, diastolic, decrescendo blowing murmur along the left sternal border, results from dilation of the pulmonary

valve ring and occurs in patients with mitral valve disease and severe pulmonary hypertension. This murmur may be indistinguishable from the more common murmur produced by aortic regurgitation (AR), although it may increase in intensity with inspiration and is accompanied by a loud and often palpable P_2 .

LABORATORY EXAMINATION

ECG In MS and sinus rhythm, the P wave usually suggests LA enlargement (see Fig. 268-8). It may become tall and peaked in lead II and upright in lead V_1 when severe pulmonary hypertension or TS complicates MS and right atrial (RA) enlargement occurs. The QRS complex is usually normal. However, with severe pulmonary hypertension, right axis deviation and RV hypertrophy are often present.

Echocardiogram (See also Chap. 270e) Transthoracic echocardiography (TTE) with color flow and spectral Doppler imaging provides critical information, including measurements of mitral inflow velocity during early (E wave) and late (A wave in patients in sinus rhythm) diastolic filling, estimates of the transvalvular peak and mean gradients and of the mitral orifice area, the presence and severity of any associated MR, the extent of leaflet calcification and restriction, the degree of distortion of the subvalvular apparatus, and the anatomic suitability for percutaneous mitral balloon valvotomy (percutaneous mitral balloon valvuloplasty [PMBV]; see below). In addition, TTE provides an assessment of LV and RV function, chamber sizes, an estimation of the PAP based on the tricuspid regurgitant jet velocity, and an indication of the presence and severity of any associated valvular lesions, such as aortic stenosis and/or regurgitation. Transesophageal echocardiography (TEE) provides superior images and should be used when TTE is inadequate for guiding management decisions. TEE is especially indicated to exclude the presence of LA thrombus prior to PMBV. The performance of TTE with exercise to evaluate the mean mitral diastolic gradient and PA pressures can be very helpful in the evaluation of patients with MS when there is a discrepancy between the clinical findings and the resting hemodynamics.

Chest X-Ray The earliest changes are straightening of the upper left border of the cardiac silhouette, prominence of the main PAs, dilation of the upper lobe pulmonary veins, and posterior displacement of the esophagus by an enlarged LA. Kerley B lines are fine, dense, opaque, horizontal lines that are most prominent in the lower and mid-lung fields and that result from distention of interlobular septae and lymphatics with edema when the resting mean LA pressure exceeds approximately 20 mmHg.

DIFFERENTIAL DIAGNOSIS

Like MS, significant MR may also be associated with a prominent diastolic murmur at the apex due to increased antegrade transmitral flow, but in patients with isolated MR, this diastolic murmur commences slightly later than in patients with MS, and there is often clear-cut evidence of LV enlargement. An OS and increased P_2 are absent, and S_1 is soft or absent. An apical pansystolic murmur of at least grade III/VI intensity as well as an S_3 suggest significant MR. Similarly, the apical mid-diastolic murmur associated with severe AR (*Austin Flint murmur*) may be mistaken for MS but can be differentiated from it because it is not intensified in presystole and becomes softer with administration of amyl nitrite or other arterial vasodilators. TS, which occurs rarely in the absence of MS, may mask many of the clinical features of MS or be clinically silent; when present, the diastolic murmur of TS increases with inspiration and the γ descent in the jugular venous pulse is delayed.

Atrial septal defect (Chap. 282) may be mistaken for MS; in both conditions, there is often clinical, ECG, and chest x-ray evidence of RV enlargement and accentuation of pulmonary vascularity. However, the absence of LA enlargement and of Kerley B lines and the demonstration of fixed splitting of S_2 with a grade II or III mid-systolic murmur at the mid to upper left sternal border all favor atrial septal defect over MS. Atrial septal defects with large left-to-right shunts may result in functional TS because of the enhanced diastolic flow.

Left atrial myxoma (Chap. 289e) may obstruct LA emptying, causing dyspnea, a diastolic murmur, and hemodynamic changes resembling those of MS. However, patients with an LA myxoma often have features suggestive of a systemic disease, such as weight loss, fever, anemia, systemic emboli, and elevated serum IgG and interleukin 6 (IL-6) concentrations. The auscultatory findings may change markedly with body position. The diagnosis can be established by the demonstration of a characteristic echo-producing mass in the LA with TTE.

CARDIAC CATHETERIZATION

Left and right heart catheterization can be useful when there is a discrepancy between the clinical and noninvasive findings, including those from TEE and exercise echocardiographic testing as appropriate. Catheterization is helpful in assessing associated lesions, such as aortic stenosis (AS) and AR. Catheterization and coronary angiography are not usually necessary to aid in decision-making about surgery in patients younger than 65 years of age with typical findings of severe mitral obstruction on physical examination and TTE. In men older than 40 years of age, women older than 45 years of age, and younger patients with coronary risk factors, especially those with positive noninvasive stress tests for myocardial ischemia, coronary angiography is advisable preoperatively to identify patients with critical coronary obstructions that should be bypassed at the time of operation. Computed tomographic coronary angiography (CTCA) (Chap. 270e) is now often used to screen preoperatively for the

presence of coronary artery disease (CAD) in patients with valvular heart disease and low pretest likelihood of CAD. Catheterization and left ventriculography may be useful in patients who have undergone PMBV or previous mitral valve surgery for MS, and who have redeveloped limiting symptoms, especially if questions regarding the severity of the valve lesion(s) remain after noninvasive study.

TREATMENT MITRAL STENOSIS

(Fig. 284-1) Penicillin prophylaxis of group A β -hemolytic streptococcal infections (Chap. 381) for secondary prevention of rheumatic fever is important for at-risk patients with rheumatic MS. Recommendations for infective endocarditis prophylaxis are similar to those for other valve lesions and are restricted to patients at high risk for complications from infection, including patients with a history of endocarditis. In symptomatic patients, some improvement usually occurs with restriction of sodium intake and small doses of oral diuretics. Beta blockers, nondihydropyridine calcium channel blockers (e.g., verapamil or diltiazem), and digitalis glycosides are useful in slowing the ventricular rate of patients with AF. Warfarin therapy targeted to an international normalized ratio (INR) of 2–3 should be administered indefinitely to patients with MS who have AF or a history of thromboembolism. The routine use of warfarin in patients in sinus rhythm with LA enlargement (maximal dimension

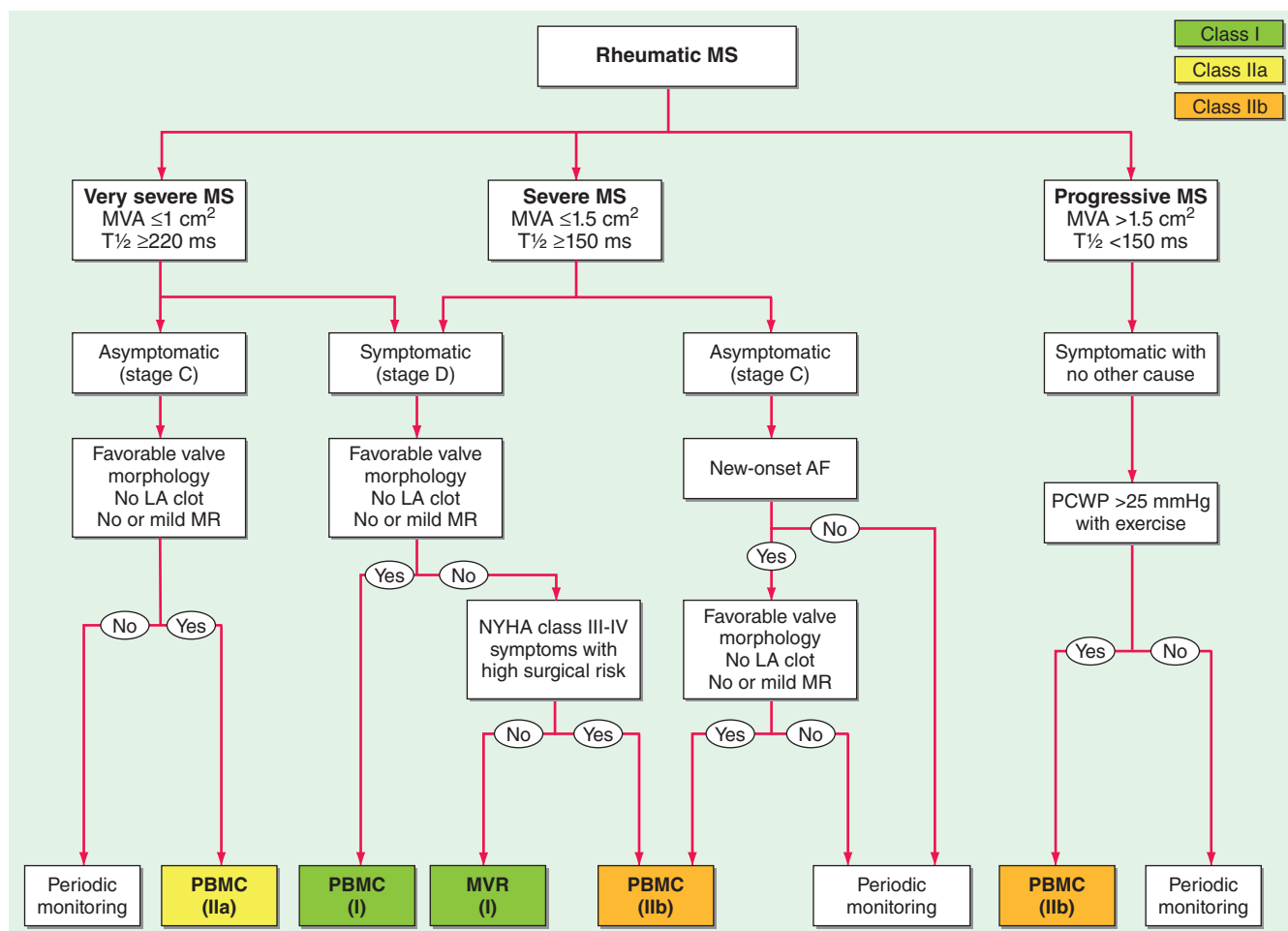


FIGURE 284-1 Management of rheumatic mitral stenosis. See legend for Fig. 283-2 for explanation of treatment recommendations (class I, IIa, IIb) and disease stages (C, D). Preoperative coronary angiography should be performed routinely as determined by age, symptoms, and coronary risk factors. Cardiac catheterization and angiography may also be helpful when there is a discrepancy between clinical and noninvasive findings. AF, atrial fibrillation; LA, left atrial; MR, mitral regurgitation; MS, mitral stenosis; MVA, mitral valve area; MVR, mitral valve surgery (repair or replacement); NYHA, New York Heart Association; PCWP, pulmonary capillary wedge pressure; PMBC, percutaneous mitral balloon commissurotomy; and T_{1/2}, pressure half-time. (Adapted from RA Nishimura et al: 2014 AHA/ACC Guideline for the Management of Patients with Valvular Heart Disease. *J Am Coll Cardiol* doi: 10.1016/j.jacc.2014.02.536, 2014, with permission.)

1542 >5.5 cm) with or without spontaneous echo contrast is more controversial. The novel oral anticoagulants are not approved for use in patients with significant valvular heart disease.

If AF is of relatively recent onset in a patient whose MS is not severe enough to warrant PMBV or surgical commissurotomy, reversion to sinus rhythm pharmacologically or by means of electrical countershock is indicated. Usually, cardioversion should be undertaken after the patient has had at least 3 consecutive weeks of anticoagulant treatment to a therapeutic INR. If cardioversion is indicated more urgently, then intravenous heparin should be provided and TEE performed to exclude the presence of LA thrombus before the procedure. Conversion to sinus rhythm is rarely successful or sustained in patients with severe MS, particularly those in whom the LA is especially enlarged or in whom AF has been present for more than 1 year.

MITRAL VALVOTOMY

Unless there is a contraindication, mitral valvotomy is indicated in symptomatic (New York Heart Association [NYHA] Functional Class II–IV) patients with isolated severe MS, whose effective orifice (valve area) is $< 1 \text{ cm}^2/\text{m}^2$ body surface area, or $< 1.5 \text{ cm}^2$ in normal-sized adults. Mitral valvotomy can be carried out by two techniques: PMBV and surgical valvotomy. In PMBV (Figs. 284-2 and 284-3), a catheter is directed into the LA after transseptal puncture, and a single balloon is directed across the valve and inflated in the valvular orifice. Ideal patients have relatively pliable leaflets with little or no commissural calcium. In addition, the subvalvular structures should not be significantly scarred or thickened, and there should be no LA thrombus. The short- and long-term results of this procedure in appropriate patients are similar to those of surgical valvotomy, but with less morbidity and a lower periprocedural mortality rate. Event-free

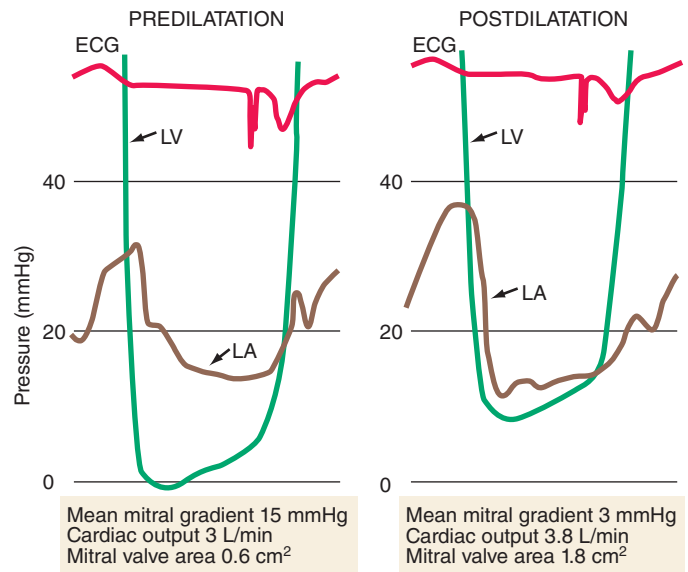


FIGURE 284-3 Simultaneous left atrial (LA) and left ventricular (LV) pressure before and after percutaneous mitral balloon valvuloplasty (PMBV) in a patient with severe mitral stenosis. ECG, electrocardiogram. (Courtesy of Raymond G. McKay, MD; with permission.)

survival in younger (<45 years) patients with pliable valves is excellent, with rates as high as 80–90% over 3–7 years. Therefore, PMBV has become the procedure of choice for such patients when it can be performed by a skilled operator in a high-volume center.

TTE is helpful in identifying patients for the percutaneous procedure, and TEE is performed routinely to exclude LA thrombus and to assess the degree of MR at the time of the scheduled procedure. An “echo score” has been developed to help guide decision-making. The score accounts for the degree of leaflet thickening, calcification, and mobility, and for the extent of subvalvular thickening. A lower score predicts a higher likelihood of successful PMBV.

In patients in whom PMBV is not possible or unsuccessful, or in many patients with restenosis after previous surgery, an “open” valvotomy using cardiopulmonary bypass is necessary. In addition to opening the valve commissures, it is important to loosen any subvalvular fusion of papillary muscles and chordae tendineae; to remove large deposits of calcium, thereby improving valvular function; and to remove atrial thrombi. The perioperative mortality rate is ~2%.

Successful valvotomy is defined by a 50% reduction in the mean mitral valve gradient and a doubling of the mitral valve area. Successful valvotomy, whether balloon or surgical, usually results in striking symptomatic and hemodynamic improvement and prolongs survival. However, there is no evidence that the procedure improves the prognosis of patients with slight or no functional impairment. Therefore, unless recurrent systemic embolization or severe pulmonary hypertension has occurred (PA systolic pressures $> 50 \text{ mmHg}$ at rest or $> 60 \text{ mmHg}$ with exercise), valvotomy is *not* recommended for patients who are entirely asymptomatic and/or who have mild or moderate stenosis (mitral valve area $> 1.5 \text{ cm}^2$). When there is little symptomatic improvement after valvotomy, it is likely that the procedure was ineffective, that it induced MR, or that associated valvular or myocardial disease was present. About half of all patients undergoing surgical mitral valvotomy require reoperation by 10 years. In the pregnant patient with MS, valvotomy should be carried out if pulmonary congestion occurs despite intensive medical treatment. PMBV is the preferred strategy in this setting and is performed with TEE and no or minimal x-ray exposure.

Mitral valve replacement (MVR) is necessary in patients with MS and significant associated MR, those in whom the valve has been severely distorted by previous transcatheter or operative

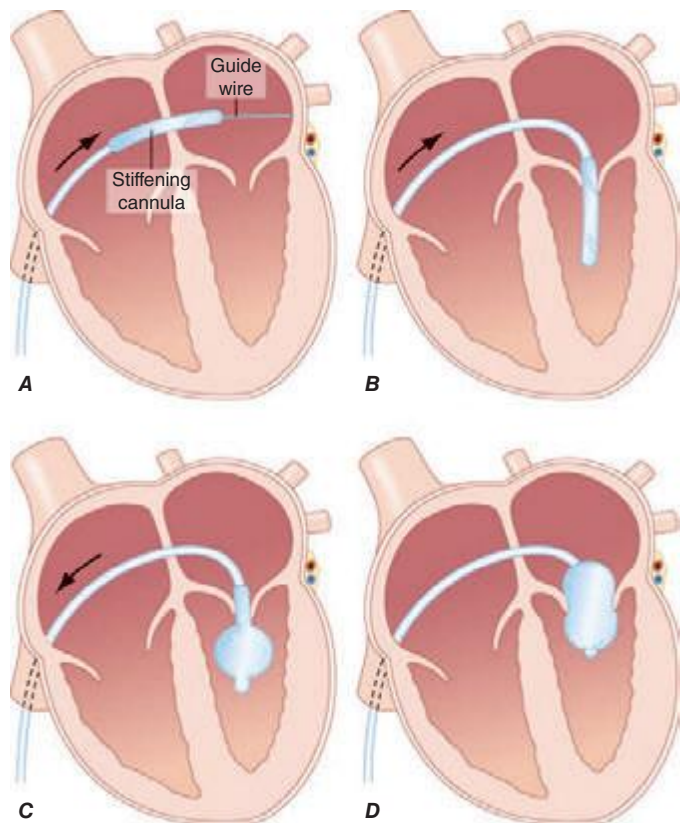


FIGURE 284-2 Inoue balloon technique for percutaneous mitral balloon valvotomy. **A.** After transseptal puncture, the deflated balloon catheter is advanced across the interatrial septum, then across the mitral valve and into the left ventricle. **B–D.** The balloon is inflated stepwise within the mitral orifice.

TABLE 284-2 MORTALITY RATES AFTER MITRAL VALVE SURGERY^a

Operation	Number	Unadjusted Operative Mortality (%)
MVR (isolated)	3154	5.2
MVR + CAB	1184	10.0
MVRp	4215	1.0
MVRp + CAB	2330	4.8

^aData are for the first two quarters of calendar year 2013, during which 1004 sites reported a total of 135,666 procedures. Data are available from the Society of Thoracic Surgeons at http://www.sts.org/sites/default/files/documents/2013_3rdHarvestExecutiveSummary.pdf.

Abbreviations: CAB, coronary artery bypass; MVR, mitral valve replacement; MVRp, mitral valve repair.

manipulation, or those in whom the surgeon does not find it possible to improve valve function significantly with valvotomy. MVR is now routinely performed with preservation of the chordal attachments to optimize LV functional recovery. Perioperative mortality rates with MVR vary with age, LV function, the presence of CAD, and associated comorbidities. They average 5% overall but are lower in young patients and may be twice as high in patients >65 years of age with significant comorbidities (Table 284-2). Because there are also long-term complications of valve replacement, patients in whom preoperative evaluation suggests the possibility that MVR may be required should be operated on only if they have severe MS—i.e., an orifice area ≤ 1.5 cm²—and are in NYHA Class III, i.e., symptomatic with ordinary activity despite optimal medical therapy. The overall 10-year survival of surgical survivors is ~70%. Long-term prognosis is worse in patients >65 years of age and those with marked disability and marked depression of the CO preoperatively. Pulmonary hypertension and RV dysfunction are additional risk factors for poor outcome.

MITRAL REGURGITATION

ETIOLOGY

MR may result from an abnormality or disease process that affects any one or more of the five functional components of the mitral valve apparatus (leaflets, annulus, chordae tendineae, papillary muscles, and subjacent myocardium) (Table 284-1). Acute MR can occur in the setting of acute myocardial infarction (MI) with papillary muscle rupture (Chap. 295), following blunt chest wall trauma, or during the course of infective endocarditis. With acute MI, the posteromedial papillary muscle is involved much more frequently than the anterolateral papillary muscle because of its singular blood supply. Transient, acute MR can occur during periods of active ischemia and bouts of angina pectoris. Rupture of chordae tendineae can result in “acute-on-chronic MR” in patients with myxomatous degeneration of the valve apparatus.

Chronic MR can result from rheumatic disease, mitral valve prolapse (MVP), extensive mitral annular calcification, congenital valve defects, hypertrophic obstructive cardiomyopathy (HOCM), and dilated cardiomyopathy (Chap. 287). Distinction also should be drawn between primary (degenerative, organic) MR, in which the leaflets and/or chordae tendineae are primarily responsible for abnormal valve function, and functional (secondary) MR, in which the leaflets and chordae tendineae are structurally normal but the regurgitation is caused by annular enlargement, papillary muscle displacement, leaflet tethering, or their combination. The rheumatic process produces rigidity, deformity, and retraction of the valve cusps and commissural fusion, as well as shortening, contraction, and fusion of the chordae tendineae. The MR associated with both MVP and HOCM is usually dynamic in nature. MR in HOCM occurs as a consequence of anterior papillary muscle displacement and systolic anterior motion of the anterior mitral valve leaflet into the narrowed LV outflow tract. Annular calcification is especially prevalent among patients with advanced renal disease and is commonly observed in women >65 years of age with hypertension and diabetes. MR may occur as a congenital

anomaly (Chap. 282), most commonly as a defect of the endocardial cushions (atrioventricular cushion defects). A cleft anterior mitral valve leaflet accompanies primum atrial septal defect. Chronic MR is frequently secondary to ischemia and may occur as a consequence of ventricular remodeling, papillary muscle displacement, and leaflet tethering, or with fibrosis of a papillary muscle, in patients with healed MI(s) and ischemic cardiomyopathy. Similar mechanisms of annular dilation and ventricular remodeling contribute to the MR that occurs among patients with nonischemic forms of dilated cardiomyopathy once the LV end-diastolic dimension reaches 6 cm.

Irrespective of cause, chronic severe MR is often progressive, because enlargement of the LA places tension on the posterior mitral leaflet, pulling it away from the mitral orifice and thereby aggravating the valvular dysfunction. Similarly, LV dilation increases the regurgitation, which, in turn, enlarges the LA and LV further, resulting in a vicious circle; hence the aphorism, “mitral regurgitation begets mitral regurgitation.”

PATHOPHYSIOLOGY

The resistance to LV emptying (LV afterload) is reduced in patients with MR. As a consequence, the LV is decompressed into the LA during ejection, and with the reduction in LV size during systole, there is a rapid decline in LV tension. The initial compensation to MR is more complete LV emptying. However, LV volume increases progressively with time as the severity of the regurgitation increases and as LV contractile function deteriorates. This increase in LV volume is often accompanied by a reduced forward CO. LV compliance is often increased, and thus, LV diastolic pressure does not increase until late in the course. The regurgitant volume varies directly with the LV systolic pressure and the size of the regurgitant orifice; the latter, in turn, is influenced by the extent of LV and mitral annular dilation. Because EF rises in severe MR in the presence of normal LV function, even a modest reduction in this parameter (<60%) reflects significant dysfunction.

During early diastole, as the distended LA empties, there is a particularly rapid *y* descent in the absence of accompanying MS. A brief, early diastolic LA-LV pressure gradient (often generating a rapid filling sound [S₃] and mid-diastolic murmur masquerading as MS) may occur in patients with pure, severe MR as a result of the very rapid flow of blood across a normal-sized mitral orifice.

Semiquantitative estimates of LV ejection fraction (LVEF), CO, PA systolic pressure, regurgitant volume, regurgitant fraction (RF), and the effective regurgitant orifice area can be obtained during a careful Doppler echocardiographic examination. These measurements can also be obtained accurately with cardiac magnetic resonance (CMR) imaging, although this technology is not widely available. Left and right heart catheterization with contrast ventriculography is used less frequently. Severe, nonischemic MR is defined by a regurgitant volume ≥ 60 mL/beat, RF $\geq 50\%$, and effective regurgitant orifice area ≥ 0.40 cm². Severe ischemic MR, however, is usually associated with an effective regurgitant orifice area of >0.2 cm². In the latter instance, lesser degrees of MR carry relatively greater prognostic weight.

LA Compliance In acute severe MR, the regurgitant volume is delivered into a normal-sized LA having normal or reduced compliance. As a result, LA pressures rise markedly for any increase in LA volume. The *v* wave in the LA pressure pulse is usually prominent, LA and pulmonary venous pressures are markedly elevated, and pulmonary edema is common. Because of the rapid rise in LA pressures during ventricular systole, the murmur of acute MR is early in timing and decrescendo in configuration ending well before S₂, as a reflection of the progressive diminution in the LV-LA pressure gradient. LV systolic function in acute MR may be normal, hyperdynamic, or reduced, depending on the clinical context.

Patients with chronic severe MR, on the other hand, develop marked LA enlargement and *increased* LA compliance with little if any increase in LA and pulmonary venous pressures for any increase in LA volume. The LA *v* wave is relatively less prominent. The murmur of chronic MR is classically holosystolic in timing and plateau in configuration, as a reflection of the near-constant LV-LA pressure gradient. These patients usually complain of severe fatigue and exhaustion secondary

1544 to a low forward CO, whereas symptoms resulting from pulmonary congestion are less prominent initially; AF is almost invariably present once the LA dilates significantly.

SYMPTOMS

Patients with chronic mild-to-moderate, isolated MR are usually asymptomatic. This form of LV volume overload is well tolerated. Fatigue, exertional dyspnea, and orthopnea are the most prominent complaints in patients with chronic severe MR. Palpitations are common and may signify the onset of AF. Right-sided heart failure, with painful hepatic congestion, ankle edema, distended neck veins, ascites, and secondary TR, occurs in patients with MR who have associated pulmonary vascular disease and pulmonary hypertension. Acute pulmonary edema is common in patients with acute severe MR.

PHYSICAL FINDINGS

In patients with chronic severe MR, the arterial pressure is usually normal, although the carotid arterial pulse may show a sharp, low-volume upstroke owing to the reduced forward CO. A systolic thrill is often palpable at the cardiac apex, the LV is hyperdynamic with a brisk systolic impulse and a palpable rapid-filling wave (S_3), and the apex beat is often displaced laterally.

In patients with acute severe MR, the arterial pressure may be reduced with a narrow pulse pressure, the jugular venous pressure and wave forms may be normal or increased and exaggerated, the apical impulse is not displaced, and signs of pulmonary congestion are prominent.

Auscultation S_1 is generally absent, soft, or buried in the holosystolic murmur of chronic, severe MR. In patients with severe MR, the aortic valve may close prematurely, resulting in wide but physiologic splitting of S_2 . A low-pitched S_3 occurring 0.12–0.17 s after the aortic valve closure sound, i.e., at the completion of the rapid-filling phase of the LV, is believed to be caused by the sudden tensing of the papillary muscles, chordae tendineae, and valve leaflets. It may be followed by a short, rumbling, mid-diastolic murmur, even in the absence of structural MS. A fourth heart sound is often audible in patients with acute severe MR who are in sinus rhythm. A presystolic murmur is not ordinarily heard with isolated MR.

A systolic murmur of at least grade III/VI intensity is the most characteristic auscultatory finding in chronic severe MR. It is usually holosystolic (see Fig. 267-5A), but as previously noted, it is decrescendo and ceases in mid to late systole in patients with acute severe MR. The systolic murmur of chronic MR is usually most prominent at the apex and radiates to the axilla. However, in patients with ruptured chordae tendineae or primary involvement of the posterior mitral leaflet with prolapse or flail, the regurgitant jet is eccentric, directed anteriorly, and strikes the LA wall adjacent to the aortic root. In this situation, the systolic murmur is transmitted to the base of the heart and, therefore, may be confused with the murmur of AS. In patients with ruptured chordae tendineae, the systolic murmur may have a cooing or “seagull” quality, whereas a flail leaflet may produce a murmur with a musical quality. The systolic murmur of chronic MR not due to MVP is intensified by isometric exercise (handgrip) but is reduced during the strain phase of the Valsalva maneuver because of the associated decrease in LV preload.

LABORATORY EXAMINATION

ECG In patients with sinus rhythm, there is evidence of LA enlargement, but RA enlargement also may be present when pulmonary hypertension is significant and affects RV function. Chronic severe MR is frequently associated with AF. In many patients, there is no clear-cut ECG evidence of enlargement of either ventricle. In others, the signs of eccentric LV hypertrophy are present.

Echocardiogram TTE is indicated to assess the mechanism of the MR and its hemodynamic severity. LV function can be assessed from LV end-diastolic and end-systolic volumes and EF. Observations can be made regarding leaflet structure and function, chordal integrity, LA and LV size, annular calcification, and regional and global LV systolic function. Doppler imaging should demonstrate the width or

area of the color flow MR jet within the LA, the duration and intensity of the continuous wave Doppler signal, the pulmonary venous flow contour, the early peak mitral inflow velocity, and quantitative measures of regurgitant volume, RF, and effective regurgitant orifice area. In addition, the PAPs can be estimated from the TR jet velocity. TTE is also indicated to follow the course of patients with chronic MR and to provide rapid assessment for any clinical change. The echocardiogram in patients with MVP is described in the next section. TEE provides greater anatomic detail than TTE (see Fig. 270e-5). Exercise testing with TTE can be useful to assess exercise capacity as well as any dynamic change in MR severity, PA systolic pressures, and biventricular function, for patients in whom there is a discrepancy between clinical findings and the results of functional testing performed at rest.

Chest X-Ray The LA and LV are the dominant chambers in chronic MR. Late in the course of the disease, the LA may be massively enlarged and forms the right border of the cardiac silhouette. Pulmonary venous congestion, interstitial edema, and Kerley B lines are sometimes noted. Marked calcification of the mitral leaflets occurs commonly in patients with long-standing, combined rheumatic MR and MS. Calcification of the mitral annulus may be visualized, particularly on the lateral view of the chest. Patients with acute severe MR may have asymmetric pulmonary edema if the regurgitant jet is directed predominantly to the orifice of an upper lobe pulmonary vein.

TREATMENT MITRAL REGURGITATION

MEDICAL TREATMENT (FIG. 284-4)

The management of chronic severe MR depends to some degree on its cause. Warfarin should be provided once AF intervenes with a target INR of 2–3. Novel oral anticoagulants are not approved for this indication. Cardioversion should be considered depending on the clinical context and LA size. In contrast to the acute setting, there are no large, long-term prospective studies to substantiate the use of vasodilators for the treatment of chronic, isolated severe MR with preserved LV systolic function in the absence of systemic hypertension. The severity of MR in the setting of an ischemic or nonischemic dilated cardiomyopathy may diminish with aggressive guideline-directed treatment of heart failure including the use of diuretics, beta blockers, angiotensin-converting enzyme (ACE) inhibitors, digitalis, and biventricular pacing (cardiac resynchronization therapy [CRT]) when otherwise indicated. Asymptomatic patients with severe MR in sinus rhythm with normal LV size and systolic function should avoid isometric forms of exercise.

Patients with acute severe MR require urgent stabilization and preparation for surgery. Diuretics, intravenous vasodilators (particularly sodium nitroprusside), and even intraaortic balloon counterpulsation may be needed for patients with post-MI papillary muscle rupture or other forms of acute severe MR.

SURGICAL TREATMENT

In the selection of patients with chronic, nonischemic, primary or organic, severe MR for surgical treatment, the often slowly progressive nature of the condition must be balanced against the immediate and long-term risks associated with operation. These risks are significantly lower for primary valve repair than for valve replacement (Table 287-2). Repair usually consists of valve reconstruction using a variety of valvuloplasty techniques and insertion of an annuloplasty ring. Repair spares the patient the long-term adverse consequences of valve replacement, including thromboembolic and hemorrhagic complications in the case of mechanical prostheses and late valve failure necessitating repeat valve replacement in the case of bioprostheses. In addition, by preserving the integrity of the papillary muscles, subvalvular apparatus, and chordae tendineae, mitral repair and valvuloplasty maintain LV function to a relatively greater degree.

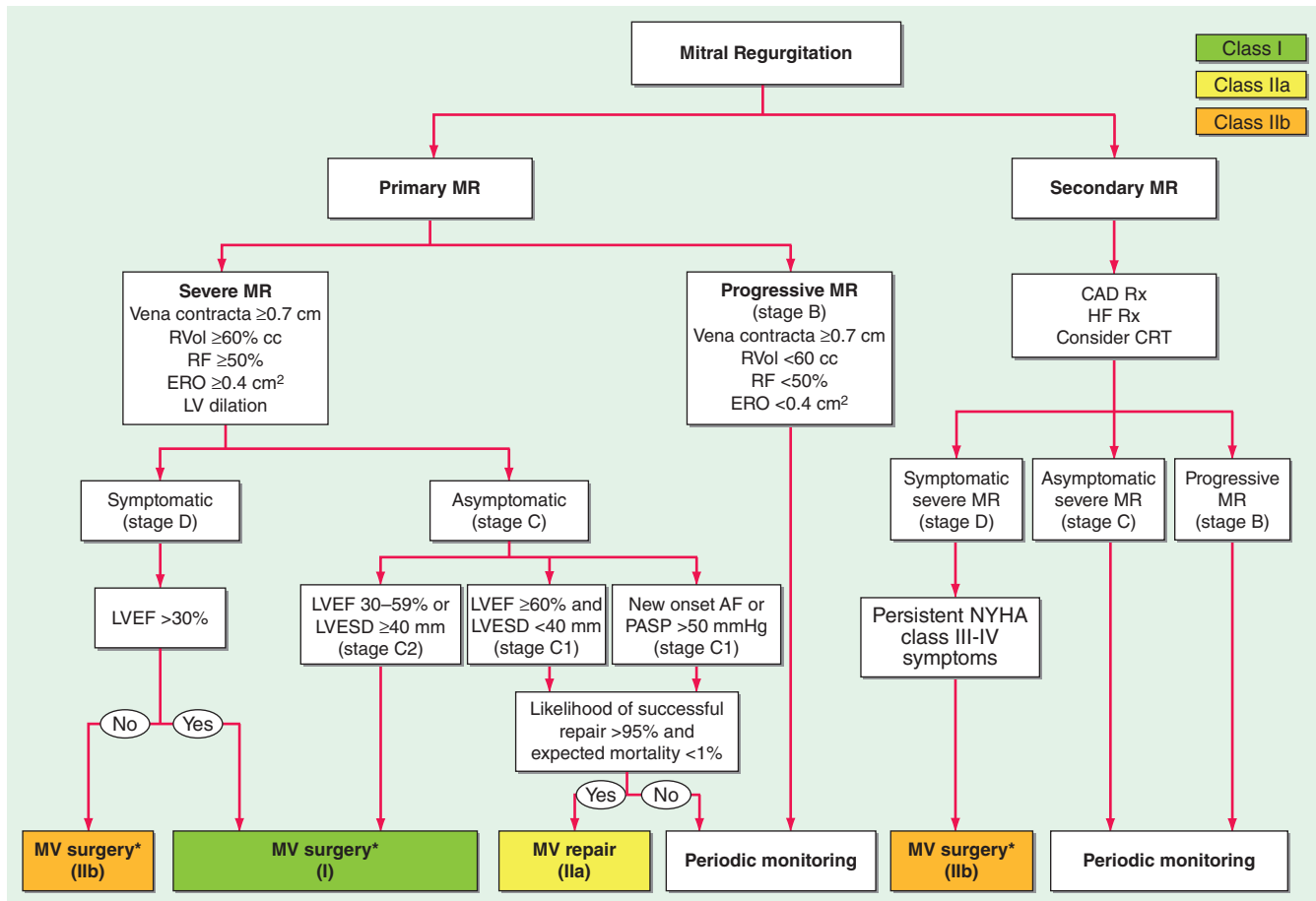


FIGURE 284-4 Management of mitral regurgitation. See legend for Fig. 283-2 for explanation of treatment recommendations (class I, IIa, IIb) and disease stages (B, C1, C2, D). Preoperative coronary angiography should be performed routinely as determined by age, symptoms, and coronary risk factors. Cardiac catheterization and angiography may also be helpful when there is a discrepancy between clinical and noninvasive findings. AF, atrial fibrillation; CAD, coronary artery disease; CRT, cardiac resynchronization therapy; ERO, effective regurgitant orifice; HF, heart failure; LV, left ventricular; LVEF, left ventricular ejection fraction; LVESD, left ventricular end-systolic dimension; MR, mitral regurgitation, MV, mitral valve; MVR, mitral valve replacement; NYHA, New York Heart Association; PASP, pulmonary artery systolic pressure; RF, regurgitant fraction; RVol, regurgitant volume; and Rx, therapy. *Mitral valve repair preferred over MVR when possible. (Adapted from RA Nishimura et al: 2014 AHA/ACC Guideline for the Management of Patients with Valvular Heart Disease. *J Am Coll Cardiol* doi: 10.1016/j.jacc.2014.02.536, 2014, with permission.)

Surgery for chronic nonischemic severe MR is indicated once symptoms occur, especially if valve repair is feasible (Fig. 284-4). Other indications for early consideration of mitral valve repair include recent-onset AF and pulmonary hypertension defined as a systolic PA pressure ≥ 50 mmHg at rest or ≥ 60 mmHg with exercise. Surgical treatment of chronic nonischemic severe MR is indicated for asymptomatic patients when LV dysfunction is progressive with the LVEF falling below 60% and/or end-systolic dimension increasing beyond 40 mm. These aggressive recommendations for surgery are predicated on the outstanding results achieved with mitral valve repair particularly when applied to patients with myxomatous disease such as that associated with prolapse or flail leaflet. Indeed primary valvuloplasty repair of patients younger than 75 years with normal LV systolic function and no CAD can now be performed by experienced surgeons with $< 1\%$ perioperative mortality risk. The risk of stroke, however, is also approximately 1%. Repair is feasible in up to 95% of patients with myxomatous disease operated on by a high-volume surgeon in a referral center of excellence. Long-term durability is excellent; the incidence of reoperative surgery for failed primary repair is $\sim 1\%$ per year for the first 10 years after surgery. For patients with AF, left or biatrial maze surgery, or radiofrequency, isolation of the pulmonary veins is often performed to reduce the risk of recurrent postoperative AF.

The surgical management of patients with functional, ischemic MR is more complicated and most often involves simultaneous

coronary artery revascularization. Current surgical practice includes annuloplasty repair with an undersized, rigid ring or chord-sparing valve replacement for patients with moderate or greater degrees of MR. Valve repair for ischemic MR is associated with lower perioperative mortality rates but higher rates of recurrent MR over time. In patients with ischemic MR and significantly impaired LV systolic function (EF $< 30\%$), the risk of surgery is higher, recovery of LV performance is incomplete, and long-term survival is reduced. Referral for surgery must be individualized and made only after aggressive attempts with guideline-directed medical therapy and CRT, when indicated. The routine performance of valve repair in patients with significant MR in the setting of severe, functional, nonischemic dilated cardiomyopathy has not been shown to improve long-term survival compared with optimal medical therapy. Patients with acute severe MR can often be stabilized temporarily with appropriate medical therapy, but surgical correction will be necessary emergently in the case of papillary muscle rupture and within days to weeks in most other settings.

When surgical treatment is contemplated, left and right heart catheterization and left ventriculography may be helpful in confirming the presence of severe MR in patients in whom there is a discrepancy between the clinical and TTE findings that cannot be resolved with TEE or CMR. Coronary angiography identifies patients who require concomitant coronary revascularization.



FIGURE 284-5 Clip used to grasp the free edges of the anterior and posterior leaflets in their midsections during transcatheter repair of selected patients with mitral regurgitation. (Courtesy of Abbott Vascular. © 2014 Abbott Laboratories. All rights reserved.)

TRANSCATHETER MITRAL VALVE REPAIR

A transcatheter approach to the treatment of either organic or functional MR may be feasible in selected patients with appropriate anatomy. The proper role of currently available techniques remains under active investigation. One approach involves the deployment of a clip delivered via transseptal puncture that grasps the leading edges of the mitral leaflets in their mid-portion (anterior scallop to posterior scallop or A2-P2; Fig. 284-5). The length and width of the gap between these leading edges dictate patient eligibility. The device is commercially available for the treatment of prohibitive surgical risk patients with severe, degenerative (organic) MR and is undergoing study in the United States for treatment of patients with symptomatic heart failure, reduced LVEF, and severe, functional MR despite guideline-directed medical therapy. A second approach involves the deployment of a device within the coronary sinus that can be adjusted to reduce its circumference, thus secondarily decreasing the circumference of the mitral annulus and the effective orifice area of the valve much like a surgically implanted ring. Variations in the anatomic relationship of the coronary sinus to the mitral annulus and circumflex coronary artery have limited the applicability of this technique. Attempts to reduce the septal-lateral dimension of a dilated annulus using adjustable cords placed across the LV in a subvalvular location have also been investigated.

MITRAL VALVE PROLAPSE

MVP, also variously termed the *systolic click-murmur syndrome*, *Barlow's syndrome*, *floppy-valve syndrome*, and *billowing mitral leaflet syndrome*, is a relatively common but highly variable clinical syndrome resulting from diverse pathologic mechanisms of the mitral valve apparatus. Among these are excessive or redundant mitral leaflet tissue, which is commonly associated with myxomatous degeneration and greatly increased concentrations of certain glycosaminoglycans.

In most patients with MVP, the cause is unknown, but in some, it appears to be genetically determined. A reduction in the production of type III collagen has been incriminated, and electron microscopy has revealed fragmentation of collagen fibrils.

MVP is a frequent finding in patients with heritable disorders of connective tissue, including Marfan's syndrome (Chap. 427), osteogenesis imperfecta, and Ehlers-Danlos syndrome. MVP may be associated with thoracic skeletal deformities similar to but not as severe as those in Marfan's syndrome, such as a high-arched palate and alterations of the chest and thoracic spine, including the so-called straight back syndrome.

In most patients with MVP, myxomatous degeneration is confined to the mitral valve, although the tricuspid and aortic valves may also be affected. The posterior mitral leaflet is usually more affected than the anterior, and the mitral valve annulus is often dilated. In many patients, elongated, redundant, or ruptured chordae tendineae cause or contribute to the regurgitation.

MVP also may occur rarely as a sequel to acute rheumatic fever, in ischemic heart disease, and in various cardiomyopathies, as well as in 20% of patients with ostium secundum atrial septal defect.

MVP may lead to excessive stress on the papillary muscles, which, in turn, leads to dysfunction and ischemia of the papillary muscles and the subjacent ventricular myocardium. Rupture of chordae tendineae and progressive annular dilation and calcification contribute to valvular regurgitation, which then places more stress on the diseased mitral valve apparatus, thereby creating a vicious circle. ECG changes (see below) and ventricular arrhythmias described in some patients with MVP appear to result from regional ventricular dysfunction related to the increased stress placed on the papillary muscles.

CLINICAL FEATURES

MVP is more common in women and occurs most frequently between the ages of 15 and 30 years; the clinical course is most often benign. MVP may also be observed in older (>50 years) patients, often men, in whom MR is often more severe and requires surgical treatment. There is an increased familial incidence for some patients, suggesting an autosomal dominant form of inheritance with incomplete penetrance. MVP varies in its clinical expression, ranging from only a systolic click and murmur with mild prolapse of the posterior leaflet to severe MR due to chordal rupture and leaflet flail. The degree of myxomatous change of the leaflets can also vary widely. In many patients, the condition progresses over years or decades; in others, it worsens rapidly as a result of chordal rupture or endocarditis.

Most patients are asymptomatic and remain so for their entire lives. However, in North America, MVP is now the most common cause of isolated severe MR requiring surgical treatment. Arrhythmias, most commonly ventricular premature contractions and paroxysmal supraventricular and ventricular tachycardia, as well as AF, have been reported and may cause palpitations, light-headedness, and syncope. Sudden death is a very rare complication and occurs most often in patients with severe MR and depressed LV systolic function. There may be an excess risk of sudden death among patients with a flail leaflet. Many patients have chest pain that is difficult to evaluate; it is often substernal, prolonged, and not related to exertion, but may rarely resemble angina pectoris. Transient cerebral ischemic attacks secondary to emboli from the mitral valve due to endothelial disruption have been reported. Infective endocarditis may occur in patients with MR and/or leaflet thickening.

Auscultation A frequent finding is the mid or late (nonejection) systolic click, which occurs 0.14 s or more after S_1 and is thought to be generated by the sudden tensing of slack, elongated chordae tendineae or by the prolapsing mitral leaflet when it reaches its maximal excursion. Systolic clicks may be multiple and may be followed by a high-pitched, mid-late systolic crescendo-decrescendo murmur, which occasionally is "whooping" or "honking" and is heard best at the apex. The click and murmur occur earlier with standing, during the strain phase of the Valsalva maneuver, and with any intervention that decreases LV volume, exaggerating the propensity of mitral leaflet prolapse. Conversely, squatting and isometric exercises, which increase LV volume, diminish MVP; the click-murmur complex is delayed, moves away from S_1 , and may even disappear. Some patients have a mid-systolic click without a murmur; others have a murmur without a click. Still others have both sounds at different times.

LABORATORY EXAMINATION

The ECG most commonly is normal but may show biphasic or inverted T waves in leads II, III, and aVF, and occasionally supraventricular or ventricular premature beats. TTE is particularly effective in identifying the abnormal position and prolapse of the mitral valve leaflets. A useful echocardiographic definition of MVP is systolic displacement (in the

parasternal long axis view) of the mitral valve leaflets by at least 2 mm into the LA superior to the plane of the mitral annulus. Color flow and continuous wave Doppler imaging is helpful to evaluate the associated MR and provide semiquantitative estimates of severity. The jet lesion of MR due to MVP is most often eccentric, and assessment of RF and effective regurgitant orifice area can be difficult. TEE is indicated when more accurate information is required and is performed routinely for intraoperative guidance for valve repair. Invasive left ventriculography is rarely necessary but can also show prolapse of the posterior and sometimes of both mitral valve leaflets.

TREATMENT MITRAL VALVE PROLAPSE

Infective endocarditis prophylaxis is indicated only for patients with a prior history of endocarditis. Beta blockers sometimes relieve chest pain and control palpitations. If the patient is symptomatic from severe MR, mitral valve repair (or rarely, chord-sparing replacement) is indicated (Fig. 284-4). Antiplatelet agents, such as aspirin, should be given to patients with transient ischemic attacks, and if these are not effective, warfarin should be considered. Warfarin is also indicated once AF intervenes.

285 Tricuspid and Pulmonic Valve Disease

Patrick T. O'Gara, Joseph Loscalzo

TRICUSPID STENOSIS

Tricuspid stenosis (TS), which is much less prevalent than mitral stenosis (MS) in North America and Western Europe, is generally rheumatic in origin, and is more common in women than men (Table 285-1). It does not occur as an isolated lesion and is usually associated with MS. Hemodynamically significant TS occurs in 5–10% of patients with severe MS; rheumatic TS is commonly associated with some degree of tricuspid regurgitation (TR). Nonrheumatic causes of TS are rare.

PATHOPHYSIOLOGY

A diastolic pressure gradient between the right atrium (RA) and right ventricle (RV) defines TS. It is augmented when the transvalvular blood flow increases during inspiration and declines during expiration. A mean diastolic pressure gradient of 4 mmHg is usually sufficient to elevate the mean RA pressure to levels that result in systemic venous congestion. Unless sodium intake has been restricted and diuretics administered, this venous congestion is associated with hepatomegaly, ascites, and edema, sometimes severe. In patients with sinus rhythm, the RA *a* wave may be extremely tall and may even approach the level of the RV systolic pressure. The *y* descent is prolonged. The cardiac output (CO) at rest is usually depressed, and it fails to rise during exercise. The low CO is responsible for the normal or only slightly elevated left atrial (LA), pulmonary artery (PA), and RV systolic pressures despite the presence of MS. Thus, the presence of TS can mask the hemodynamic and clinical features of any associated MS.

SYMPTOMS

Because the development of MS generally precedes that of TS, many patients initially have symptoms of pulmonary congestion and fatigue. Characteristically, patients with severe TS complain of relatively little dyspnea for the degree of hepatomegaly, ascites, and edema that they have. However, fatigue secondary to a low CO and discomfort due to refractory edema, ascites, and marked hepatomegaly are common in patients with advanced TS and/or TR. In some patients, TS may be

TABLE 285-1 CAUSES OF TRICUSPID AND PULMONIC VALVE DISEASES

Valve Lesion	Etiologies
Tricuspid stenosis	Rheumatic Congenital
Tricuspid regurgitation	Primary (organic) Rheumatic Endocarditis Myxomatous (TVP) Carcinoid Radiation Congenital (Ebstein's) Trauma Papillary muscle injury (post-MI) Secondary (functional) RV and tricuspid annular dilatation due to multiple causes of RV enlargement (e.g., long-standing pulmonary HTN, remodeling post-RV MI) Chronic RV apical pacing
Pulmonic stenosis	Congenital Carcinoid Tumor Endocarditis
Pulmonic regurgitation	Primary valve disease Congenital Postvalvotomy Endocarditis Carcinoid Annular enlargement Pulmonary hypertension Idiopathic dilation Marfan's syndrome

Abbreviations: HTN, hypertension; MI, myocardial infarction; RV, right ventricular; TVP, tricuspid valve prolapse.

suspected for the first time when symptoms of right-sided failure persist after an adequate mitral valvotomy.

PHYSICAL FINDINGS

Because TS usually occurs in the presence of other obvious valvular disease, the diagnosis may be missed unless it is considered. Severe TS is associated with marked hepatic congestion, often resulting in cirrhosis, jaundice, serious malnutrition, anasarca, and ascites. Congestive hepatomegaly and, in cases of severe tricuspid valve disease, splenomegaly are present. The jugular veins are distended, and in patients with sinus rhythm, there may be giant *a* waves. The *v* waves are less conspicuous, and because tricuspid obstruction impedes RA emptying during diastole, there is a slow *y* descent. In patients with sinus rhythm, there may be prominent presystolic pulsations of the enlarged liver as well.

On auscultation, an opening snap (OS) of the tricuspid valve may rarely be heard approximately 0.06 s after pulmonic valve closure. The diastolic murmur of TS has many of the qualities of the diastolic murmur of MS, and because TS almost always occurs in the presence of MS, it may be missed. However, the tricuspid murmur is generally heard best along the left lower sternal border and over the xiphoid process, and is most prominent during presystole in patients with sinus rhythm. The murmur of TS is augmented during inspiration, and it is reduced during expiration and particularly during the strain phase of the Valsalva maneuver, when tricuspid transvalvular flow is reduced.

LABORATORY EXAMINATION

The electrocardiogram (ECG) features of RA enlargement (see Fig. 268-8) include tall, peaked P waves in lead II, as well as prominent, upright P waves in lead V₁. The absence of ECG evidence of RV

1548 hypertrophy (RVH) in a patient with right-sided heart failure who is believed to have MS should suggest associated tricuspid valve disease. The chest x-ray in patients with combined TS and MS shows particular prominence of the RA and superior vena cava without much enlargement of the PA and with less evidence of pulmonary vascular congestion than occurs in patients with isolated MS. On echocardiographic examination, the tricuspid valve is usually thickened and domes in diastole; the transvalvular gradient can be estimated by continuous wave Doppler echocardiography. Severe TS is characterized by a valve area ≤ 1 cm² or pressure half-time of ≥ 190 ms. The RA and inferior vena cava (IVC) are enlarged. Transthoracic echocardiography (TTE) provides additional information regarding the severity of any associated TR, mitral valve structure and function, left ventricle (LV) and RV size and function, and PA pressure. Cardiac catheterization is not routinely necessary for assessment of TS.

can lead to “ventricularization” of the RA wave form (see Fig. 267-1B). Severe TR is also characterized by RV dilation (RV volume overload) and eventual systolic dysfunction, the rate of which can be accelerated by a concomitant pressure load from PA hypertension or by myocardial fibrosis from previous injury.

SYMPTOMS

Mild or moderate degrees of TR are usually well tolerated in the absence of other hemodynamic disturbances. Because TR most often coexists with left-sided valve lesions, LV dysfunction, and/or PA hypertension, symptoms related to these lesions may dominate the clinical picture. Fatigue and exertional dyspnea owing to reduced forward CO are early symptoms of isolated, severe TR. As the disease progresses and RV function declines, patients may report cervical pulsations, abdominal fullness/bloating, diminished appetite, and muscle wasting, although with progressive weight gain and painful swelling of the lower extremities.

PHYSICAL FINDINGS

The neck veins in patients with severe TR are distended with prominent *c-v* waves and rapid *y* descents (in the absence of TS). TR is more often diagnosed by examination of the neck veins than by auscultation of the heart sounds. Other findings may include marked hepatomegaly with systolic pulsations, ascites, pleural effusions, edema, and a positive hepatojugular reflex. A prominent RV pulsation along the left parasternal region and a blowing holosystolic murmur along the lower left sternal margin, which may be intensified during inspiration (Carvalho’s sign) and reduced during expiration or the strain phase of the Valsalva maneuver, are characteristic findings. The murmur of TR may sometimes be confused with that of MR unless attention is paid to its variation during the respiratory cycle and the extent of RV enlargement is appreciated. Atrial fibrillation (AF) is usually present in the chronic phase of the disease.

LABORATORY EXAMINATION

The ECG may show changes characteristic of the lesion responsible for the TR, e.g., an inferior Q-wave MI suggestive of a prior RV MI, RVH, or a bizarre right bundle branch block type pattern with preexcitation in patients with Ebstein’s anomaly. ECG signs of RA enlargement may be present in patients with sinus rhythm; AF is frequently noted. The chest x-ray may show RA and RV enlargement, depending on the chronicity and severity of TR. TTE is usually definitive with demonstration of RA dilation and RV volume overload and prolapsing, flail, scarred, or displaced/tethered tricuspid leaflets; the diagnosis and assessment of TR can be made by color flow Doppler imaging (see Fig. 270e-8). Severe TR is accompanied by hepatic vein systolic flow reversal. Continuous wave Doppler of the TR velocity profile is useful in estimating PA systolic pressure. Accurate assessment of TR severity, PA pressures, and RV size and systolic function with TTE can be quite challenging in many patients. Real-time three-dimensional echocardiography and cardiac magnetic resonance (CMR) imaging provide alternative imaging modalities, although they are not widely available. In patients with severe TR, the CO is usually markedly reduced, and the RA pressure pulse may exhibit no *x* descent during early systole but a prominent *c-v* wave with a rapid *y* descent. The mean RA and RV end-diastolic pressures are often elevated. Exercise testing can be used to assess functional capacity in patients with asymptomatic severe TR. The prognostic significance of exercise-induced changes in TR severity and RV function has not been well studied.

TREATMENT TRICUSPID REGURGITATION (FIG. 285-1)

Diuretics can be useful for patients with severe TR and signs of right heart failure. An aldosterone antagonist may be particularly helpful because many patients have secondary hyperaldosteronism from marked hepatic congestion. Therapies to reduce elevated PA pressures and/or pulmonary vascular resistance, including those targeted at left-sided heart disease, can also be considered for patients with PA hypertension and severe functional TR. Tricuspid valve surgery is recommended for patients with severe TR who are undergoing

TREATMENT TRICUSPID STENOSIS

Patients with TS generally exhibit marked systemic venous congestion; salt restriction, bed rest, and diuretic therapy are required during the preoperative period. Such a preparatory period may diminish hepatic congestion and thereby improve hepatic function sufficiently so that the risks of operation, particularly bleeding, are diminished. Surgical relief of the TS should be carried out, preferably at the time of surgical mitral valvotomy or mitral valve replacement (MVR) for mitral valve disease, in patients with moderate or severe TS who have mean diastolic pressure gradients exceeding ~ 4 mmHg and tricuspid orifice areas < 1.5 – 2 cm². TS is almost always accompanied by significant TR. Operative repair may permit substantial improvement of tricuspid valve function. If repair cannot be accomplished, the tricuspid valve may have to be replaced. Meta-analysis has shown no difference in overall survival between mechanical and tissue valve replacement. Mechanical valves in the tricuspid position are more prone to thromboembolic complications than in other positions. Percutaneous tricuspid balloon valvuloplasty for isolated severe TS without significant TR is very rarely performed.

TRICUSPID REGURGITATION

In at least 80% of cases, TR is secondary to marked dilation of the tricuspid annulus from RV enlargement due to PA hypertension (Table 285-1). Functional TR may complicate RV enlargement of any cause, however, including an inferior myocardial infarction (MI) that involves the RV. It is commonly seen in the late stages of heart failure due to rheumatic or congenital heart disease with severe PA hypertension (PA systolic pressure > 55 mmHg), as well as in ischemic and idiopathic dilated cardiomyopathies. It is reversible in part if PA hypertension can be relieved. Functional TR can also develop from chronic RV apical pacing. Rheumatic fever may produce primary (organic) TR, often associated with TS. Infarction of RV papillary muscles, tricuspid valve prolapse, carcinoid heart disease, endomyocardial fibrosis, radiation, infective endocarditis, and leaflet trauma all may produce TR. Less commonly, TR results from congenitally deformed tricuspid valves, and it occurs with defects of the atrioventricular canal, as well as with Ebstein’s malformation of the tricuspid valve (Chap. 282).

PATHOPHYSIOLOGY

The incompetent tricuspid valve allows blood to flow backward from the RV into the RA, the volume of which is dependent on the driving pressure (i.e., RV systolic pressure) and the size of the regurgitant orifice. The severity and physical signs of TR can vary as a function of PA systolic pressure (in the absence of RV outflow tract stenosis), the dimension of the tricuspid valve annulus, the respiratory cycle-dependent changes in RV preload, and RA compliance. RV filling is increased during inspiration. Forward CO is reduced and does not augment with exercise. Significant degrees of TR will lead to RA enlargement and elevation of the RA and jugular venous pressures with prominent *c-v* waves in the pulse tracings. Progressively severe TR

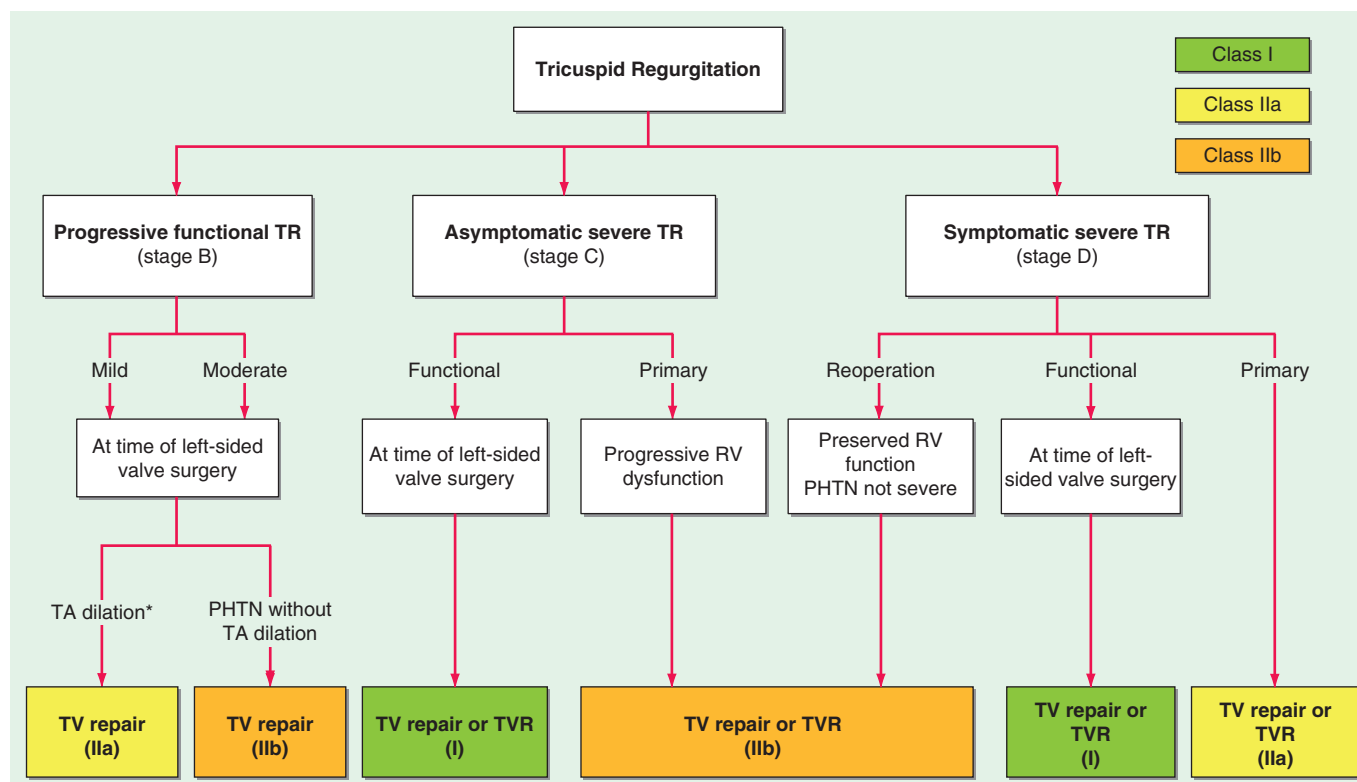


FIGURE 285-1 Management of tricuspid regurgitation. See legend for Fig. 283-2 for explanation of treatment recommendations (Class I, IIa, IIb) and disease stages (B, C, D). Preoperative coronary angiography should be performed routinely as determined by age, symptoms, and coronary risk factors. Cardiac catheterization and angiography may also be helpful when there is a discrepancy between clinical and noninvasive findings. PHTN, pulmonary hypertension; RV, right ventricular; TA, tricuspid annular; TTE, transthoracic echocardiogram; TR, tricuspid regurgitation; TV, tricuspid valve; TVR, tricuspid valve replacement. *TA dilation is defined by >40 mm on TTE (>21 mm/m²) or >70 mm on direct intraoperative measurement. (Adapted from RA Nishimura et al: 2014 AHA/ACC Guideline for the Management of Patients with Valvular Heart Disease. *J Am Coll Cardiol* doi: 10.1016/j.jacc.2014.02.536, 2014, with permission.)

left-sided valve surgery and is also undertaken frequently for treatment of even moderate TR in patients undergoing left-sided valve surgery who have tricuspid annular dilation (>40 mm), a history of right heart failure, or PA hypertension. Operation most often comprises repair rather than replacement in these settings and has become routine in most major surgical centers. Surgery may also infrequently be required for treatment of severe, primary TR with right heart failure not responsive to standard medical therapy or because of progressively declining RV systolic function. Reported perioperative mortality rates for isolated tricuspid valve surgery (repair and replacement) are high (~8-9%) and likely are influenced by the hazards encountered during reoperation on patients who have undergone previous left-sided valve surgery and have reduced RV function. Indwelling pacemaker or defibrillator leads can also pose technical challenges.

PULMONIC STENOSIS

Pulmonic valve stenosis (PS) is essentially a congenital disorder (Table 285-1). With isolated PS, the valve is typically domed. Dysplastic pulmonic valves are seen as part of the Noonan's syndrome (Chap. 302), which maps to chromosome 12. Much less common etiologies include carcinoid and obstructing tumors or bulky vegetations. The pulmonic valve is only very rarely affected by the rheumatic process.

PATHOPHYSIOLOGY

PS is defined hemodynamically by a systolic pressure gradient between the RV and main PA. RV hypertrophy develops as a consequence of sustained obstruction to RV outflow, and systolic ejection is prolonged. Compared with the ability of the LV to compensate for the pressure overload imposed by aortic stenosis (AS), RV dysfunction from afterload mismatch occurs earlier in the course of PS and at lower peak systolic pressures, because the RV adapts less well to this type of

hemodynamic burden. With normal systolic function and CO, severe PS is defined by a peak systolic gradient across the pulmonic valve of >50 mmHg; moderate PS correlates with a peak gradient of 30–50 mmHg. PS rarely progresses in patients with peak gradients less than 30 mmHg, but may worsen in those with moderate disease due to valve thickening and calcification with age. The RA *a* wave elevates in relation to the higher pressures needed to fill a noncompliant, hypertrophied RV. A prominent RA *v* wave signifies functional TR from RV and annular dilation. The CO is maintained until late in the course of the disease.

SYMPTOMS

Patients with mild or even moderate PS are usually asymptomatic and first come to medical attention because of a heart murmur that leads to echocardiography. With severe PS, patients may report exertional dyspnea or early-onset fatigue. Anginal chest pain from RV oxygen supply-demand mismatch and syncope may occur with very severe forms of obstruction, particularly in the presence of a destabilizing trigger such as atrial fibrillation, fever, infection, or anemia.

PHYSICAL FINDINGS

The murmur of mild or moderate PS is mid-systolic in timing, crescendo-decrescendo in configuration, heard best in the left second interspace, and usually introduced by an ejection sound (click) in younger adults whose valves are still pliable. The ejection sound is the only right-sided acoustic event that decreases in intensity with inspiration. This phenomenon reflects premature opening of the pulmonic valve by the elevated RV end-diastolic (postatrial *a* wave) pressure. The systolic murmur increases in intensity during inspiration. With progressively severe PS, the ejection sound moves closer to the first heart sound and eventually becomes inaudible. A right-sided fourth heart sound may emerge. The systolic murmur peaks later and may persist through the aortic component of the second heart sound (*A*₂). Pulmonic valve

1550 closure is delayed, and the pulmonic component of the second heart sound (P_2) is reduced or absent. A prominent *a* wave, indicative of the higher atrial pressure necessary to fill the noncompliant RV, may be seen in the jugular venous pulse. A parasternal or RV lift can be felt with significant pressure overload. Signs of right heart failure, such as hepatomegaly, ascites, and edema, are uncommon but may appear very late in the disease.

LABORATORY EXAMINATION

The ECG will show right axis deviation, RVH, and RA enlargement in adult patients with severe PS. Chest x-ray findings include poststenotic dilation of the main PA in the frontal plane projection and filling of the retrosternal airspace due to RV enlargement on the lateral film. In some patients with RVH, the cardiac apex appears to be lifted off the left hemidiaphragm. The RA may also be enlarged. TTE allows definitive diagnosis and characterization in most cases, with depiction of the valve and assessment of the gradient, RV function, PA pressures (which should be low), and any associated cardiac lesions. TEE may be useful in some patients for improved delineation of the RV outflow tract (RVOT) and assessment of infundibular hypertrophy. Cardiac catheterization is not usually necessary, but if performed, pressures should be obtained from just below and above the pulmonic valve with attention to the possibility that a dynamic component to the gradient may exist. The correlation between Doppler assessment of peak instantaneous gradient and catheterization-measured peak-to-peak gradient is weak. The latter may correlate better with the Doppler mean gradient.

TREATMENT PULMONIC STENOSIS

Diuretics can be used to treat symptoms and signs of right heart failure. Provided there is less than moderate pulmonic regurgitation, pulmonic balloon valvotomy is recommended for symptomatic patients with a domed valve and a peak gradient >50 mmHg (or mean gradient >30 mmHg) and for asymptomatic patients with a peak gradient >60 mmHg (or mean gradient >40 mmHg). Surgery may be required when the valve is dysplastic (as seen in patients with Noonan's syndrome and other disorders). A multidisciplinary heart team is best positioned to make treatment decisions of this nature.

PULMONIC REGURGITATION

Pulmonic regurgitation (PR) may develop as a consequence of primary valve pathology, annular enlargement, or their combination; after surgical treatment of RVOT obstruction in children with such disorders as tetralogy of Fallot; or after pulmonic balloon valvotomy (Table 285-1). Carcinoid usually causes mixed pulmonic valve disease with PR and PS. Long-standing severe PA hypertension from any cause can result in dilation of the pulmonic valve ring and PR.

PATHOPHYSIOLOGY

Severe PR results in RV chamber enlargement and eccentric hypertrophy. As is the case for aortic regurgitation (AR), PR is a state of increased preload and afterload. The reverse pressure gradient from the PA to the RV, which drives the PR, progressively decreases throughout diastole and accounts for the decrescendo nature of the diastolic murmur. As RV diastolic pressure increases, the murmur becomes shorter in duration. The forward CO is preserved during the early stages of the disease, but may not increase normally with exercise and declines over time. A reduction in RV ejection fraction may be an early indicator of hemodynamic compromise. In advanced stages, there is significant enlargement of the RV and RA with marked elevation of the jugular venous pressure.

SYMPTOMS

Mild or moderate degrees of PR do not, by themselves, result in symptoms. Other problems, such as PA hypertension, may dominate the clinical picture. With progressively severe PR and RV dysfunction,

fatigue, exertional dyspnea, abdominal fullness/bloating, and lower extremity swelling may be reported.

PHYSICAL FINDINGS

The physical examination hallmark of PR is a high-pitched, decrescendo diastolic murmur (Graham Steell murmur) heard along the left sternal border that can be difficult to distinguish from the more frequently appreciated murmur of aortic regurgitation. The Graham Steell murmur may become louder with inspiration and is usually associated with a loud and sometimes palpable P_2 and an RV lift, as would be expected in patients with significant PA hypertension of any cause. Survivors of childhood surgery for tetralogy of Fallot or PS/pulmonary atresia may have an RV-PA conduit that is freely regurgitant because it does not contain a valve. PA pressures in these individuals are not elevated and the diastolic murmur can be misleadingly low pitched and of short duration despite significant degrees of PR and RV volume overload.

LABORATORY EXAMINATION

Depending on both the etiology and severity of PR, the ECG may show findings of RVH and RA enlargement. On chest x-ray, the RV and RA may be enlarged. Pulmonic valve morphology and function can be assessed with transthoracic Doppler echocardiography. PA pressures can be estimated from the tricuspid valve systolic jet velocity. CMR provides greater anatomic detail, particularly in patients with repaired congenital heart disease, and more precise assessment of RV volumes. Cardiac catheterization is not routinely necessary but would be performed as part of a planned transcatheter procedure.

TREATMENT PULMONIC REGURGITATION

In patients with functional PR due to PA hypertension and annular dilation, efforts to reduce PA vascular resistance and pressure should be optimized. Such efforts may include pharmacologic/vasodilator and/or surgical/interventional strategies, depending on the cause of the PA hypertension. Diuretics can be used to treat the manifestations of right heart failure. Surgical valve replacement for primary, severe, pulmonic valve disease, such as carcinoid or endocarditis, is rarely undertaken. Transcatheter pulmonic valve replacement has been successfully performed in many patients with severe PR after childhood repair of tetralogy of Fallot or pulmonic valve stenosis or atresia. This procedure was introduced clinically prior to transcatheter aortic valve replacement.

286 Multiple and Mixed Valvular Heart Disease

Patrick T. O'Gara, Joseph Loscalzo

Many acquired and congenital cardiac lesions may result in stenosis and/or regurgitation of one or more heart valves. For example, rheumatic heart disease can involve the mitral (mitral stenosis [MS], mitral regurgitation [MR], or MS and MR), aortic (aortic stenosis [AS], aortic regurgitation [AR], or AS and AR), and/or tricuspid (tricuspid stenosis [TS], tricuspid regurgitation [TR], or TS and TR) valve, alone or in combination. The common association of functional TR with significant mitral valve disease is discussed in [Chap. 285](#). Severe mitral annular calcification can result in regurgitation (due to decreased annular shortening during systole) and mild stenosis (caused by extension of the calcification onto the leaflets resulting in restricted valve opening). Patients with severe AS may develop functional MR that may not improve after isolated aortic valve replacement (AVR). Chordal rupture

has been described infrequently in patients with severe AS. Aortic valve infective endocarditis may secondarily involve the mitral apparatus either by abscess formation and contiguous spread via the intervalvular fibrosa or by “drop metastases” from the aortic leaflets onto the anterior leaflet of the mitral valve. Mediastinal radiation may result in aortic, mitral, and even tricuspid valve disease, most often with mixed stenosis and regurgitation. Carcinoid heart disease may cause mixed lesions of either or both the tricuspid and pulmonic valves. Ergotamines, and the previously used combination of fenfluramine and phentermine, can rarely result in mixed lesions of the aortic and/or mitral valve. Patients with Marfan’s syndrome may have both AR from aortic root dilation and MR due to mitral valve prolapse (MVP). Myxomatous degeneration causing prolapse of multiple valves (mitral, aortic, tricuspid) can also occur in the absence of an identifiable connective tissue disorder. Bicuspid aortic or pulmonic valve disease can result in mixed stenosis and regurgitation.

PATHOPHYSIOLOGY

In patients with multivalvular heart disease, the pathophysiologic derangements associated with the more proximal valve disease can mask the full expression of the attributes of the more distal valve lesion. For example, in patients with rheumatic mitral and aortic valve disease, the reduction in cardiac output (CO) imposed by the mitral valve disease will decrease the magnitude of the hemodynamic derangements related to the severity of the aortic valve lesion (stenotic, regurgitant, or both). Alternatively, the development of atrial fibrillation (AF) during the course of MS can lead to sudden worsening in a patient whose aortic valve disease was not previously felt to be significant. The development of reactive pulmonary vascular disease, sometimes referred to as a “secondary obstructive lesion in series,” can impose an additional challenge in these settings. As CO falls with progressive tricuspid valve disease, the severity of any associated mitral or aortic disease can be underestimated.

One of the most common examples of multivalve disease is that of functional TR in the setting of significant mitral valve disease. Functional TR occurs as a consequence of right ventricular and annular dilation; pulmonary artery (PA) hypertension is often present. The tricuspid leaflets are morphologically normal. Progressive degrees of TR lead to right ventricular volume overload and continued chamber and annular dilation. The TR is usually central in origin; reflux into the right atrium (RA) is expressed as large, systolic *c-v* waves in the RA pressure pulse. The height of the *c-v* wave is dependent on RA compliance and the volume of regurgitant flow. The RA wave form may become “ventricularized” in advanced stages of chronic, severe TR with PA hypertension. CO falls and the severity of the associated mitral valve disease may become more difficult to appreciate. Primary rheumatic tricuspid valve disease may occur with rheumatic mitral disease and cause hemodynamic changes reflective of TR, TS, or their combination. With TS, the γ descent in the RA pressure pulse is prolonged.

Another example of rheumatic, multivalve disease involves the combination of mitral and aortic valve pathology, frequently characterized by MS and AR. In isolated MS, left ventricular (LV) preload and diastolic pressure are reduced as a function of the severity of inflow obstruction. With concomitant AR, however, LV filling is enhanced and diastolic pressure may rise depending on the compliance characteristics of the chamber. Because the CO falls with progressive degrees of MS, transaortic valve flows will decline, masking the potential severity of the aortic valve lesion (AR, AS, or its combination). As noted above, onset of AF in such patients can be especially deleterious.

Functional MR may complicate the course of some patients with severe AS. The mitral valve leaflets and chordae tendineae are usually normal. Incompetence is related to changes in LV geometry (remodeling) and abnormal systolic tethering of the leaflets in the context of markedly elevated LV systolic pressures. Relief of the excess afterload with surgical or transcatheter AVR often, but not always, results in reduction or elimination of the MR. Persistence of significant MR following AVR is associated with impaired functional outcomes and reduced survival. Identification of patients who would benefit from concomitant treatment of their functional MR at time of AVR is quite

challenging. Most surgeons advocate for repair of moderate-to-severe or severe functional MR at time of surgical AVR.

In patients with mixed AS and AR, assessment of valve stenosis can be influenced by the magnitude of the regurgitant valve flow. Because transvalvular systolic flow velocities are augmented in patients with AR and preserved LV systolic function, the LV-aortic Doppler-derived pressure gradient and the intensity of the systolic murmur will be elevated to values higher than expected for the true systolic valve orifice size as delineated by planimetry. Uncorrected, the Gorlin formula, which relies on forward CO (systolic transvalvular flow) and the mean pressure gradient for calculation of valve area, is not accurate in the setting of mixed aortic valve disease. Similar considerations apply to patients with mixed mitral valve disease. The peak mitral valve Doppler E wave velocity (v_p) is increased in the setting of severe MR because of enhanced early diastolic flow and may not accurately reflect the contribution to left atrial (LA) hypertension from any associated MS. When either AR or MR is the dominant lesion in patients with mixed aortic or mitral valve disease, respectively, the LV is dilated. When AS or MS predominates, LV chamber size will be normal or small. It can sometimes be difficult to ascertain whether stenosis or regurgitation is the dominant lesion in patients with mixed valve disease, although an integrated clinical and noninvasive assessment can usually provide clarification for purposes of patient management and follow-up.

Patients with significant AS, a nondilated LV chamber, and concentric hypertrophy will poorly tolerate the abrupt development of aortic regurgitation, as may occur, for example, with infective endocarditis or after surgical or transcatheter AVR complicated by paravalvular leakage. The noncompliant LV is not prepared to accommodate the sudden volume load, and as a result, LV diastolic pressure rises rapidly and severe heart failure develops. Indeed, paravalvular regurgitation is a significant risk factor for short- to intermediate-term death following transcatheter AVR. Conditions in which the LV may not be able to dilate in response to chronic AR (or MR) include radiation heart disease and, in some patients, the cardiomyopathy associated with obesity and diabetes. Noncompliant ventricles of small chamber size predispose to earlier onset diastolic dysfunction and heart failure in response to any further perturbation in valve function.

SYMPTOMS

Compared with patients with isolated, single-lesion valve disease, patients with multiple or mixed valve disease may develop symptoms at a relatively earlier stage in the natural history of their disease. Symptoms such as exertional dyspnea and fatigue are usually related to elevated filling pressures, reduced CO, or their combination. Palpitations may signify AF and identify mitral valve disease as an important component of the clinical presentation, even when not previously suspected. Chest pain compatible with angina could reflect left or right ventricular oxygen supply/demand mismatch on a substrate of hypertrophy and pressure/volume overload with or without superimposed coronary artery disease. Symptoms related to right heart failure (abdominal fullness/bloating, edema) are late manifestations of advanced disease.

PHYSICAL FINDINGS

Mixed disease of a single valve is most often manifested by systolic and diastolic murmurs, each with the attributes expected for the valve in question. Thus, patients with AS and AR will have characteristic mid-systolic, crescendo-decrescendo and blowing, decrescendo diastolic murmurs at the base of the heart in the second right interspace and along the left sternal edge, respectively. Many patients with significant AR have mid-systolic outflow murmurs even in the absence of valve sclerosis/stenosis, and other findings of AS must be sought. The separate murmurs of AS and AR can occasionally be difficult to distinguish from the continuous murmurs associated with either a patent ductus arteriosus (PDA) or ruptured sinus of Valsalva aneurysm. With mixed aortic valve disease, the systolic murmur should end before, and not envelope or extend through, the second heart sound (S_2). The murmur associated with a PDA is heard best to the left of the upper sternum. The continuous murmur heard with a ruptured sinus of Valsalva

1552 aneurysm is often first appreciated after an episode of acute chest pain. An early ejection click, which usually defines bicuspid aortic valve disease in young adults, is often not present in patients with congenital, mixed AS and AR. As noted above, both the intensity and duration of these separate murmurs can be influenced by a reduction in CO and transvalvular flow due to coexistent mitral valve disease. In patients with isolated MS and MR, expected findings would include a blowing, holosystolic murmur and a mid-diastolic rumble (with or without an opening snap) best heard at the cardiac apex. An irregularly irregular heart rhythm in such patients would likely signify AF. Findings with TS and TR would mimic those of left-sided MS and MR, save for the expected changes in the murmurs with respiration. The murmurs of pulmonic stenosis and regurgitation behave in a fashion directionally similar to AS and AR; dynamic changes during respiration should be noted. **Specific attributes of these cardiac murmurs are reviewed in Chap. 285.**

LABORATORY EXAMINATION

The electrocardiogram (ECG) may show evidence of ventricular hypertrophy and/or atrial enlargement. ECG signs indicative of right-sided cardiac abnormalities in patients with left-sided valve lesions should prompt additional assessment for PA hypertension and/or right-sided valve disease. The presence of AF in patients with aortic valve disease may be a clue to the presence of previously unsuspected mitral valve disease in the appropriate context. The chest x-ray can be reviewed for evidence of cardiac chamber enlargement, valve and/or annular calcification, and any abnormalities in the appearance of the pulmonary vasculature. The latter could include enlargement of the main and proximal pulmonary arteries with PA hypertension and pulmonary venous redistribution/engorgement or Kerley B lines with increasing degrees of LA hypertension. An enlarged azygos vein in the frontal projection indicates RA hypertension. Roentgenographic findings not expected based on a single or mixed valve lesion may reflect other valve disease.

Transthoracic echocardiography (TTE) is the most commonly used imaging modality for the diagnosis and characterization of multiple and/or mixed valvular heart disease and may often demonstrate findings not clinically suspected. Transesophageal echocardiography (TEE) may sometimes be required for more accurate assessment of valve anatomy (specifically, the mitral valve) and when infective endocarditis (IE) is considered responsible for the clinical presentation. TTE findings of particular interest include those related to valve morphology and function, calcification, chamber size, ventricular wall thickness, estimated PA systolic pressure, and the dimensions of the great vessels, including the root and ascending aorta, PA, and inferior vena cava. Exercise testing (with or without echocardiography) can be useful when the degree of functional limitation reported by the patient is not adequately explained by the findings on TTE performed at rest. An integrated assessment of the clinical and TTE findings is needed to help determine the dominant valve lesion(s) and establish an appropriate plan for treatment and follow-up. Natural history is usually influenced to a relatively greater degree by the dominant lesion. Exercise testing (with or without echocardiography) can

Cardiac magnetic resonance (CMR) can be used to provide additional anatomic and physiologic information when echocardiography proves suboptimal, but is less well suited to the evaluation of valve morphology. Cardiac computed tomography (CT) has been used to assess intracardiac structures in patients with complicated IE. Coronary CT angiography provides a noninvasive alternative for the assessment of coronary artery anatomy prior to surgery.

Invasive hemodynamic evaluation with right and left heart catheterization may be required to characterize more completely the individual contributions of each lesion in patients with either multiple or mixed valvular heart disease. Measurement of PA pressures and calculation of pulmonary vascular resistance (PVR) can help inform clinical decision-making in certain patient subsets, such as those with advanced mitral and tricuspid valve disease. Attention to the accurate assessment of CO is essential. Coronary angiography (if indicated) can be performed as part of the procedure. Contrast ventriculography and great vessel angiography are performed infrequently.

TREATMENT MULTIPLE AND MIXED VALVE DISEASE

Management of patients with multiple or mixed valve disease can be challenging. As noted above, it is helpful to determine the dominant valve lesion and proceed according to the treatment and follow-up recommendations for it (**Chaps. 283 to 285**), being mindful of deviations from the expected course because of problems related to another valve disorder. For example, AF that emerges in the course of moderate mitral valve disease may precipitate heart failure in patients with concomitant, severe aortic valve disease.

Medical therapies are limited and include diuretics when indicated for relief of congestion and vitamin K antagonists for anticoagulation to prevent stroke and thromboembolism in patients with AF. The novel oral anticoagulants are not approved for use in the setting of significant valvular heart disease. Blood pressure-lowering medications may be needed to treat systemic hypertension, which may aggravate left-sided regurgitant valve lesions, but should be initiated and titrated carefully. Pulmonary vasodilators to lower PVR are not generally effective in this context.

There is a paucity of evidence to inform practice guidelines for surgical and/or transcatheter valve intervention in patients with multiple or mixed valve disease. When there is a clear, dominant lesion, as for example in a patient with severe AS and mild AR, indications for intervention are straightforward and follow those recommended for patients with AS (**Chap. 283**). In other patients, however, there is less clarity, and decisions regarding intervention should be based on several considerations, including those related to lesion severity, ventricular remodeling, functional capacity, and PA pressures. In this regard, it is important to realize that patients with multiple and/or mixed valve disease may develop limiting symptoms or signs of physiologic impairment even with moderate valve lesions.

Concomitant aortic and mitral valve replacement surgery is associated with a significantly higher perioperative mortality risk than replacement of either valve alone (**see Tables 283-2 and 284-2**), and operation should be carefully considered. Double valve replacement surgery is usually performed for treatment of severe (unrepairable) valve disease at both locations and for the combination of severe disease at one location with moderate disease at the other, so as to avoid the hazards of reoperation in the intermediate to late term for progressive disease of the unoperated valve. In addition, the presence of a prosthesis in the aortic position significantly restricts surgical exposure of the native mitral valve. The need for double valve replacement may also impact the decision regarding the type of prosthesis (i.e., mechanical vs tissue).

Tricuspid valve repair for moderate or severe functional TR at the time of left-sided valve surgery is now commonplace, particularly if there is dilation of the tricuspid annulus (>40 mm). The addition of tricuspid valve repair, consisting usually of insertion of an annuloplasty ring, adds little time or complexity to the procedure and is well tolerated. Reoperation for repair (or replacement) of progressive TR years after initial surgery for left-sided valve disease, on the other hand, is associated with a relatively high perioperative mortality risk. Repair of moderate or severe functional MR at time of AVR for AS can usually be undertaken with acceptable risk for perioperative death or major complication.

The presence of moderate or severe MR in patients with rheumatic MS is a contraindication to percutaneous mitral balloon valvotomy (PMBV). Likewise, the presence of significant AR in patients with AS disqualifies them from percutaneous aortic balloon valvotomy (PABV). The presence of severe, coexistent AR was an exclusion criterion for enrollment in the initial PARTNER trials of transcatheter AVR (TAVR) in prohibitive- and high-surgical-risk patients with severe, calcific AS. Transcatheter management of both severe AS (with TAVR) and functional MR (with deployment of a MitralClip) has been reported. Further advances in transcatheter treatments for multiple and mixed valve disease are anticipated.

DEFINITION AND CLASSIFICATION

Cardiomyopathy is disease of the heart muscle. It is estimated that cardiomyopathy accounts for 5–10% of the heart failure in the 5–6 million patients carrying that diagnosis in the United States. This term is intended to exclude cardiac dysfunction that results from other structural heart disease, such as coronary artery disease, primary valve disease, or severe hypertension; however, in general usage, the phrase *ischemic cardiomyopathy* is sometimes applied to describe diffuse dysfunction attributed to multivessel coronary artery disease, and *nonischemic cardiomyopathy* to describe cardiomyopathy from other causes. As of 2006, cardiomyopathies are defined as “a heterogeneous group of diseases of the myocardium associated with mechanical and/or electrical dysfunction that usually (but not invariably) exhibit inappropriate ventricular hypertrophy or dilatation and are due to a variety of causes that frequently are genetic.”¹

The traditional classification of cardiomyopathies into a triad of dilated, restrictive, and hypertrophic was based initially on autopsy specimens and later on echocardiographic findings. Dilated and hypertrophic cardiomyopathies can be distinguished on the basis of left ventricular wall thickness and cavity dimension; however, restrictive cardiomyopathy can have variably increased wall thickness and chamber dimensions that range from reduced to slightly increased, with prominent atrial enlargement. Restrictive cardiomyopathy is now defined more on the basis of abnormal diastolic function, which is also present but initially less prominent in dilated and hypertrophic cardiomyopathy. Restrictive cardiomyopathy can overlap in presentation, gross morphology, and etiology with both hypertrophic and dilated cardiomyopathies (Table 287-1).

Expanding information renders this classification triad based on phenotype increasingly inadequate to define disease or therapy. Identification of more genetic determinants of cardiomyopathy has suggested a four-way classification scheme of etiology as primary (affecting primarily the heart) and secondary to other systemic disease. The primary causes are then divided into genetic, mixed genetic and acquired, and acquired; however, genetic information is often unavailable at the time of initial presentation, the phenotypic expression of a given mutation varies widely, and genetic predisposition influences the clinical phenotype of acquired cardiomyopathies, as well. Although the proposed genetic classification does not yet guide many current clinical strategies, it will likely become increasingly relevant as classification of disease moves beyond individual organ pathology to more integrated systems approaches.

GENERAL PRESENTATION

For all cardiomyopathies, the early symptoms often relate to exertional intolerance with breathlessness or fatigue, usually from inadequate cardiac reserve during exercise. These symptoms may initially go unnoticed or be attributed to other causes, commonly lung disease or age-dependent exercise limitation. As fluid retention leads to elevation of resting filling pressures, shortness of breath may occur during routine daily activity such as dressing and may manifest as dyspnea or cough when lying down at night. Although often considered the hallmark of congestion, peripheral edema may be absent despite severe fluid retention, particularly in younger patients in whom ascites and abdominal discomfort may dominate. The non-specific term *congestive heart failure* describes only the resulting syndrome of fluid retention, which is common to all three types of cardiomyopathy and also to cardiac structural diseases associated

with elevated filling pressures. All three types of cardiomyopathy can be associated with atrioventricular valve regurgitation, typical and atypical chest pain, atrial and ventricular tachyarrhythmias, and embolic events (Table 287-1). Initial evaluation begins with a detailed clinical history and examination, looking for clues to cardiac, extracardiac, and familial disease (Table 287-2).

GENETIC ETIOLOGIES OF CARDIOMYOPATHY



Estimates for the prevalence of genetic etiology for cardiomyopathy continue to rise, with increasing attention paid to the family history and the availability of genetic testing. Well-recognized in hypertrophic cardiomyopathy, heritability is also present in at least 30% of dilated cardiomyopathy without other clear etiology. Careful family history should elicit not only known cardiomyopathy and heart failure, but also family members who have had sudden death, often incorrectly attributed to “a massive heart attack,” who have had atrial fibrillation or pacemaker implantation by middle age, or who have muscular dystrophy.

Most familial cardiomyopathies are inherited in an autosomal dominant pattern, with occasional autosomal recessive and X-linked inheritance (Table 287-3). Missense mutations with amino acid substitutions are the most common in cardiomyopathy. Expressed mutant proteins may interfere with function of the normal allele through a dominant negative mechanism. Mutations introducing a premature stop codon (nonsense) or shift in the reading frame (frameshift) may create a truncated or unstable protein the lack of which causes cardiomyopathy (haploinsufficiency). Deletions or duplications of an entire exon or gene are uncommon causes of cardiomyopathy, except for the dystrophinopathies.

Many different genes have been implicated in human cardiomyopathy (locus heterogeneity), and many mutations within those genes have been associated with disease (allelic heterogeneity). Although most identified mutations are “private” to individual families, several specific mutations are found repeatedly, either due to a founder effect or recurrent mutations at a common residue.

Genetic cardiomyopathy is characterized by age dependence and incomplete penetrance. The defining phenotype of cardiomyopathy is rarely present at birth and, in some individuals, may never manifest. Related individuals who carry the *same* mutation may differ in the severity of cardiomyopathy and associated consequences of rhythm disorders and need for transplantation, indicating the important role of other genetic, epigenetic, and environmental modifiers in disease expression. Sex appears to play a role, as penetrance and clinical severity may be greater in men for most cardiomyopathies. Clinical disease expression is generally more severe in the 3–5% of individuals who harbor two or more mutations linked to cardiomyopathy. However, the clinical course of a patient usually cannot be predicted based on which mutation is present; thus, current therapy is based on the phenotype rather than the genetic defect. Currently, the greatest utility of genetic testing for cardiomyopathy is to inform family evaluations. However, genetic testing occasionally enables the detection of a disease for which specific therapy is indicated, such as the replacements for defective metabolic enzymes in Fabry’s disease and Gaucher disease.

GENES AND PATHWAYS IN CARDIOMYOPATHY

Mutations in sarcomeric genes, encoding the thick and thin myofibrillar proteins, are the best characterized. While the majority are associated with hypertrophic cardiomyopathy, an increasing number of sarcomeric mutations have now been implicated in dilated cardiomyopathy, and some in left ventricular noncompaction. Few mutations have been identified in excitation-contraction coupling proteins, perhaps because they are too crucial for survival to allow variation. The most commonly recognized genetic causes of dilated cardiomyopathy are structural mutations of the giant protein titin, encoded *TTN*, which maintains sarcomere structure and acts as a key signaling molecule.

As cytoskeletal proteins play crucial roles in the structure, connection, and stability of the myocyte, multiple defects in these proteins can lead to cardiomyopathy, usually with a dilated phenotype (Fig. 287-1). For example, desmin forms intermediate filaments that connect the nuclear and plasma membranes, Z-lines,

¹From BJ Maron et al: Circulation 113:1807, 2006.

TABLE 287-1 PRESENTATION WITH SYMPTOMATIC CARDIOMYOPATHY

	Dilated	Restrictive	Hypertrophic
Ejection fraction (normal >55%)	Usually <30% when symptoms severe	25–50%	>60%
Left ventricular diastolic dimension (normal <55 mm)	≥60 mm	<60 mm (may be decreased)	Often decreased
Left ventricular wall thickness	Normal or decreased	Normal or increased	Markedly increased
Atrial size	Increased, may also be primarily affected	Increased; may be massive	Increased; related to elevated filling pressures
Valvular regurgitation	Related to annular dilation; mitral appears earlier during decompensation; tricuspid regurgitation with right ventricular dysfunction	Related to endocardial involvement; frequent mitral and tricuspid regurgitation, rarely severe	Related to valve-septum interaction; mitral regurgitation
Common first symptoms	Exertional intolerance	Exertional intolerance, fluid retention early, may have dominant right-sided symptoms	Exertional intolerance; may have chest pain
Congestive symptoms ^a	Left before right, except right prominent in young adults	Right often dominates	Left-sided congestion at rest may develop late
Arrhythmias	Ventricular tachyarrhythmia; conduction block in Chagas' disease, and some families. Atrial fibrillation.	Ventricular uncommon except in sarcoidosis, conduction block in sarcoidosis and amyloidosis. Atrial fibrillation.	Ventricular tachyarrhythmias; atrial fibrillation

^aLeft-sided symptoms of pulmonary congestion: dyspnea on exertion, orthopnea, paroxysmal nocturnal dyspnea. Right-sided symptoms of systemic venous congestion: hepatic and abdominal distention, discomfort on bending, peripheral edema.

TABLE 287-2 INITIAL EVALUATION OF CARDIOMYOPATHY

Clinical Evaluation
Thorough history and physical examination to identify cardiac and noncardiac disorders ^a
Detailed family history of heart failure, cardiomyopathy, skeletal myopathy, conduction disorders, tachyarrhythmias, and sudden death
History of alcohol, illicit drugs, chemotherapy or radiation therapy ^a
Assessment of ability to perform routine and desired activities ^a
Assessment of volume status, orthostatic blood pressure, body mass index ^a
Laboratory Evaluation
Electrocardiogram ^a
Chest radiograph ^a
Two-dimensional and Doppler echocardiogram ^a
Magnetic resonance imaging for evidence of myocardial inflammation and fibrosis
Chemistry:
Serum sodium, ^a potassium, ^a calcium, ^a magnesium ^a
Fasting glucose (glycohemoglobin in diabetes mellitus)
Creatinine, ^a blood urea nitrogen ^a
Albumin, ^a total protein, ^a liver function tests ^a
Lipid profile
Thyroid-stimulating hormone ^a
Serum iron, transferrin saturation
Urinalysis
Creatine kinase isoforms
Cardiac troponin levels
Hematology:
Hemoglobin/hematocrit ^a
White blood cell count with differential, ^a including eosinophils
Erythrocyte sedimentation rate
Initial Evaluation When Specific Diagnoses Are Suspected
Titers for infection in the setting of clinical suspicion:
Acute viral (coxsackie, echovirus, influenza)
Human immunodeficiency virus
Chagas' (<i>Trypanosoma cruzi</i>), Lyme (<i>Borrelia burgdorferi</i>), toxoplasmosis
Catheterization with coronary angiography in patients with angina who are candidates for intervention ^a
Serologies for active rheumatologic disease
Endomyocardial biopsy including sample for electron microscopy when suspecting specific diagnosis with therapeutic implications
Screening for sleep-disordered breathing

^aLevel I recommendations from ACC/AHA Practice Guidelines for Chronic Heart Failure in the Adult.

and the intercalated disks between muscle cells. Desmin mutations impair the transmission of force and signaling for both cardiac and skeletal muscle and may cause combined cardiac and skeletal myopathy.

Sarcolemmal membrane protein defects are associated with dilated cardiomyopathy. The best known is dystrophin, encoded by the X chromosome gene *DMD*, abnormalities of which cause Duchenne's and Becker's muscle dystrophy. (Interestingly, abnormal dystrophin can be acquired when the coxsackie virus cleaves dystrophin during viral myocarditis.) This protein provides a network that supports the sarcolemma and also connects to the sarcomere. The progressive functional defect in both cardiac and skeletal muscle reflects vulnerability to mechanical stress. Dystrophin is associated at the membrane with a complex of other proteins, such as metavinculin, abnormalities of which also cause dilated cardiomyopathy. Defects in the sarcolemmal channel proteins (*channelopathies*) are generally associated with primary arrhythmias, but mutations in *SCN5A*, distinct from those that cause the Brugada or long-QT syndromes, have been implicated in dilated cardiomyopathy with conduction disease.

Nuclear membrane protein defects in cardiac and skeletal muscle occur in either autosomal (lamin A/C) or X-linked (emerin) patterns. These defects are associated with a high prevalence of atrial arrhythmias and conduction system disease, which can occur in some family members without or before detectable cardiomyopathy.

Intercalated disks contribute to intracellular connections, allowing mechanical and electrical coupling between cells and also connections to desmin filaments within the cell. Mutations in proteins of the desmosomal complex compromise attachment of the myocytes, which can become disconnected and die, to be replaced by fat and fibrous tissue. These areas are highly arrhythmogenic and may dilate to form aneurysms. Although more often noted in the right ventricle (arrhythmogenic right ventricular dysplasia), this condition can affect both ventricles and has also been termed "arrhythmogenic cardiomyopathy."

Owing to the conservation of signaling pathways in multiple systems, we may expect to discover more extracardiac manifestations of genetic abnormalities initially considered to manifest exclusively in the heart. In contrast, the monogenic disorders of metabolism that affect the heart are already clearly recognized to affect multiple organ systems. Currently, it is most important to diagnose defective enzymes for which specific enzyme replacement therapy can now ameliorate the course of disease, such as with alpha-galactosidase A deficiency (Fabry's disease). Abnormalities of mitochondrial DNA (maternally transmitted) impair energy production with multiple clinical manifestations, including impaired cognitive function and skeletal myopathy. The phenotypic expression is highly variable depending on the

TABLE 287-3 SELECTED GENETIC DEFECTS ASSOCIATED WITH CARDIOMYOPATHY

	Gene Product	Inheritance	Cardiac Phenotype	Isolated Cardiac Phenotype	Extracardiac Manifestations
Sarcomere	<i>MYH7</i> (β myosin heavy chain)	AD	HCM, DCM, LVNC	Yes	Skeletal myopathy
	<i>MYBPC3</i> (myosin binding protein C)	AD	HCM	Yes	
	<i>TNNT2</i> (cardiac troponin T)	AD	HCM, DCM, LVNC	Yes	
	<i>TNNI3</i> (cardiac troponin I)	AD, AR	HCM, DCM, RCM	Yes	
	<i>TTN</i> (Titin)	AD	DCM	Yes	
	<i>TPM1</i> (α -tropomyosin)	AD	HCM, DCM	Yes	
	<i>TNNC1</i> (cardiac troponin C)	AD	DCM	Yes	
	<i>ACTC</i> (α -actin)	AD	HCM, DCM, (LVNC)	Yes	
	<i>MYL2</i> (myosin regulatory light chain)	AD	HCM	Yes	Skeletal myopathy
	<i>MYL3</i> (myosin essential light chain)	AD	HCM	Yes	
Z-disk and Cytoskeleton	<i>DES</i> (Desmin)	AD	DCM, RCM	Yes	Skeletal myopathy
	<i>LDB3</i> (Cypher-ZASP)	AD	DCM, LVNC	Yes	Skeletal myopathy
	<i>MYOZ2</i> (Myozenin)	AD	HCM	Yes	
	<i>TCAP</i> (Telethonin)	AD	DCM, HCM	Yes	
	<i>ANKRD1</i> (CARP)	AD	HCM, (DCM)	Yes	
	<i>CSRP3</i> (MLP)	AD	DCM, (HCM)	Yes	
	<i>ACTN2</i> (α -actinin-2)	AD	DCM	Yes	
	<i>CRYAB</i> (α B-crystallin)	AD	DCM	Yes	
Nuclear Membrane	<i>LMNA</i> (Lamin A/C)	AD, AR	CDDC	Yes	Skeletal myopathy
	<i>EMD</i> (Emerin)	X-linked	CDDC	No	Skeletal myopathy, contractures
Excitation-Contraction Coupling	<i>PLN</i> (Phospholamban)	AD	DCM	Yes	
	<i>SCN5A</i> (NAV 1.5)	AD	CDDC	Yes	Note other mutations associated with Brugada syndrome
	<i>RYR2</i> (cardiac ryanodine receptor)	AD	ARVC	Yes	
	<i>CASQ2</i> (calsequestrin 2)	AR	ARVC	Yes	
Cellular Metabolism	<i>PRKAG2</i> (γ -subunit of AMP kinase)	AD	HCM+	Yes	
	<i>LAMP2</i> (lysosomal associated membrane protein)	X-linked	HCM+	No ^a	Danon's disease: skeletal myopathy, cognitive impairment
	<i>TAZ</i> (Tafazzin)	X-linked	DCM, LVNC	No	Barth's syndrome: skeletal myopathy, cognitive impairment, neutropenia
	<i>FXN</i> (Frataxin)	AR	HCM	No	Friedreich's ataxia: ataxia, diabetes mellitus type 2
	<i>TMEM43</i> (transmembrane protein 43)	AD	ARVC	Yes	
	<i>GLA</i> (α -galactosidase-A)	X-linked	HCM+	Yes	Fabry's disease: renal failure, angiokeratomas and painful neuropathy
Mitochondria	Mitochondrial DNA	Maternal transmission	DCM, HCM	No	MELAS, MERRF, Kearns-Sayre syndrome, ocular myopathy
Sarcolemmal Membrane	<i>DMD</i> (Dystrophin)	X-linked	DCM	No ^a	Duchenne's and Becker's muscular dystrophy
	<i>DMPK</i> (dystrophica myotonica protein kinase)	AD	DCM	No	Myotonic dystrophy type 1
	<i>SGCD</i> (δ -sarcoglycan)	AD	DCM	Yes	
Desmosome	<i>DSP</i> (Desmoplakin) <i>JUP</i> (Plakoglobin)	AD, AR	ARVC	Yes	Carvajal syndrome (AR), Naxos syndrome (AR), "woolly hair" and hyperkeratosis of palms and soles
	<i>DSG2</i> (Desmoglein 2) <i>DSC2</i> (Desmocollin 2) <i>PKP2</i> (Plakophilin 2)	AD	ARVC	Yes	
	<i>PKP2</i> (Plakophilin 2)				
Other Examples	<i>RBM20</i> (RNA binding motif 20)	AD	DCM	Yes	
	<i>PSEN1</i> (Presenilin-1,2)	AD	DCM	Yes	Dementia
	<i>BAG3</i> (BCL2-associated athanogene 3)	AD	DCM	Yes	

^aIndicates that isolated cardiac phenotype can occur in women with the X-linked defects.

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; ARVC, arrhythmogenic right ventricular cardiomyopathy; CDDC, conduction disease with dilated cardiomyopathy; DCM, dilated cardiomyopathy; HCM, hypertrophic cardiomyopathy; HCM+, HCM with preexcitation; HCMc, HCM with conduction disease; LVNC, left ventricular noncompaction; MELAS, (mitochondrial) myopathy, encephalopathy, lactic acidosis, and stroke-like episodes syndrome; MERRF, myoclonic epilepsy with ragged red fibers; RCM, restrictive cardiomyopathy.

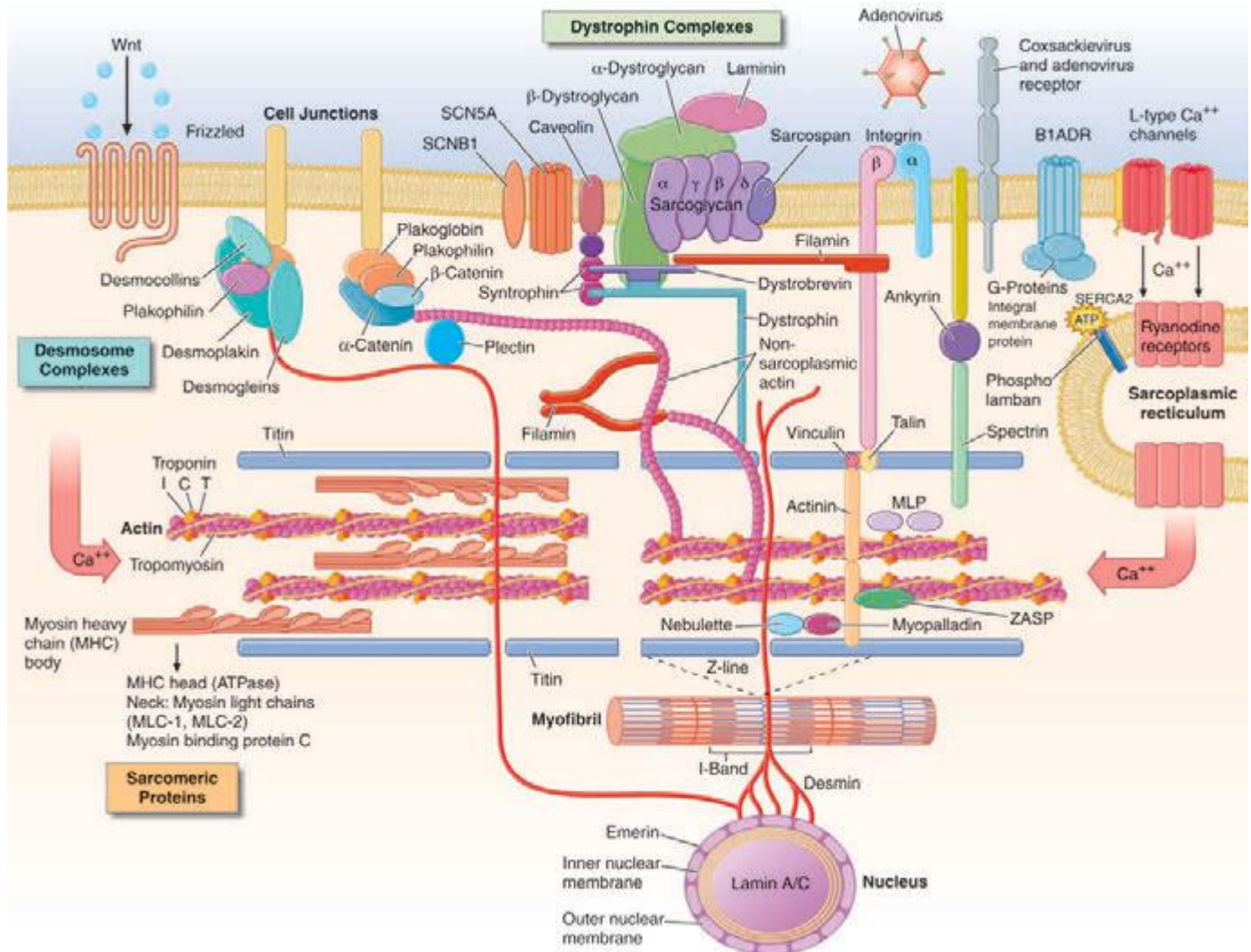


FIGURE 287-1 Drawing of myocyte indicating multiple sites of abnormal gene products associated with cardiomyopathy. Major functional groups include the sarcomeric proteins (actin, myosin, tropomyosin, and the associated regulatory proteins), the dystrophin complex stabilizing and connecting the cell membrane to intracellular structures, the desmosome complexes associated with cell-cell connections and stability, and multiple cytoskeletal proteins that integrate and stabilize the myocyte. ATP, adenosine triphosphate. (Figure adapted from Jeffrey A. Towbin, MD, University of Cincinnati, with permission.)

distribution of the maternal mitochondria during embryonic development. Heritable systemic diseases, such as familial amyloidosis and hemochromatosis, can affect the heart without mutation of genes expressed in the heart.

For any patient with suspected or proven genetic disease, family members should be considered and evaluated in a longitudinal fashion. Screening includes an echocardiogram and electrocardiogram (ECG). The indications and implications for confirmatory specific genetic testing vary depending on the specific mutation. The profound questions raised by families about diseases shared and passed down merit serious and sensitive discussion, ideally provided by a trained genetic counselor.

DILATED CARDIOMYOPATHY

An enlarged left ventricle with decreased systolic function as measured by left ventricular ejection fraction characterizes dilated cardiomyopathy (Figs. 287-2, 287-3, and 287-4). *Systolic failure* is more marked than diastolic dysfunction. Although the syndrome of dilated cardiomyopathy has multiple etiologies (Table 287-4), there appear to be common pathways of secondary response and disease progression. When myocardial injury is acquired, some myocytes may die initially, whereas others survive only to have later programmed cell death (apoptosis), and remaining myocytes hypertrophy in response to

increased wall stress. Local and circulating factors stimulate deleterious secondary responses that contribute to progression of disease. Dynamic remodeling of the interstitial scaffolding affects diastolic function and the amount of ventricular dilation. Mitral regurgitation commonly develops as the valvular apparatus is distorted and is usually substantial by the time heart failure is severe. Many cases that present “acutely” have progressed silently through these stages over months to years. Dilation and decreased function of the right ventricle may result from the initial injury and occasionally dominate, but more commonly appear later in relation to mechanical interactions with the failing left ventricle and the elevated afterload presented by secondary pulmonary hypertension.

Regardless of the nature and degree of direct cell injury, the resulting functional impairment often includes some contribution from secondary responses that may be modifiable or reversible. Almost half of all patients with new-onset cardiomyopathy demonstrate substantial spontaneous recovery. Even with long-standing disease, some patients have dramatic improvement to near-normal ejection fractions during pharmacologic therapy, particularly notable with the β -adrenergic antagonists coupled with renin-angiotensin system inhibition. For patients in whom left bundle branch block precedes clinical heart failure by many years, cardiac resynchronization pacing may be particularly likely to improve ejection fraction and decrease ventricular size. Interest in the potential for recovery of cardiomyopathy has been



FIGURE 287-2 Dilated cardiomyopathy. This gross specimen of a heart removed at the time of transplantation shows massive left ventricular dilation and moderate right ventricular dilation. Although the left ventricular wall in particular appears thinned, there is significant hypertrophy of this heart, which weighs more than 800 g (upper limit of normal = 360 g). A defibrillator lead is seen traversing the tricuspid valve into the right ventricular apex. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

further stimulated by occasional “recovery” of left ventricular function after prolonged mechanical circulatory support. The diagnosis and therapy for dilated cardiomyopathy are generally dictated by the stage of heart failure (Chap. 279), with specific aspects discussed for relevant etiologies below.

MYOCARDITIS

Myocarditis (inflammation of the heart) can result from multiple causes but is most commonly attributed to infective agents that can injure the myocardium through direct invasion, production of cardiotoxic substances, or chronic inflammation with or without persistent infection. Myocarditis cannot be assumed from a presentation of decreased systolic function in the setting of an acute infection, as any severe infection causing systemic cytokine release can depress cardiac function transiently. Infectious myocarditis has been reported with almost all types of infective agents but is most commonly associated with viruses and the protozoan *Trypanosoma cruzi*.

INFECTIVE MYOCARDITIS

The pathogenesis of viral myocarditis has been extensively studied in murine models. After viruses gain entry through the respiratory or gastrointestinal tract, they can infect organs possessing specific receptors, such as the coxsackie-adenovirus receptor on the heart. Viral infection and replication can cause myocardial injury and lysis. For example, the enteroviral protease 2A facilitates viral replication and infection through degradation of the myocyte protein dystrophin, which is crucial for myocyte stability. Activation of viral receptor proteins can also activate host tyrosine kinases, which modify the cytoskeleton to facilitate further viral entry.

The first host response to infection is the nonspecific innate immune response, heavily dependent on Toll-like receptors that recognize common antigenic patterns. Cytokine release is rapid, followed by triggered activation and expansion of specific T- and B-cell populations. This initial response appears to be crucial, as early immunosuppression

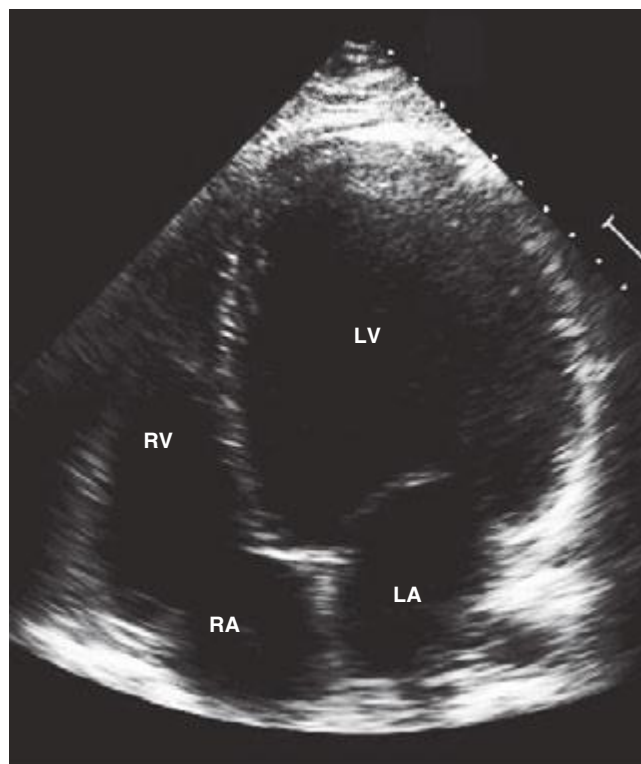


FIGURE 287-3 Dilated cardiomyopathy. This echocardiogram of a young man with dilated cardiomyopathy shows massive global dilation and thinning of the walls of the left ventricle (LV). The left atrium (LA) is also enlarged compared to normal. Note that the echocardiographic and pathologic images are vertically opposite, such that the LV is by convention on the top right in the echocardiographic image and bottom right in the pathologic images. RA, right atrium; RV, right ventricle. (Image courtesy of Justina Wu, MD, Brigham and Women's Hospital, Boston.)

in animal models can increase viral replication and worsen cardiac injury. However, successful recovery from viral infection depends not only on the efficacy of the immune response to limit viral infection, but also on timely downregulation to prevent overreaction and autoimmune injury to the host.

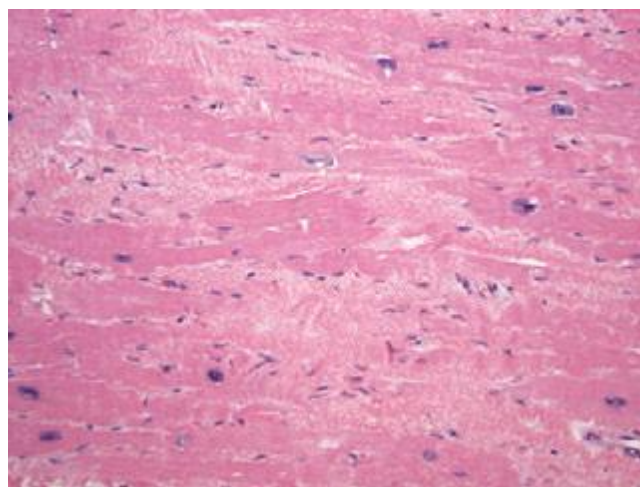


FIGURE 287-4 Dilated cardiomyopathy. Microscopic specimen of a dilated cardiomyopathy showing the nonspecific changes of interstitial fibrosis and myocyte hypertrophy characterized by increased myocyte size and enlarged, irregular nuclei. Hematoxylin and eosin–stained section, 100× original magnification. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

TABLE 287-4 MAJOR CAUSES OF DILATED CARDIOMYOPATHY (WITH COMMON EXAMPLES)

Inflammatory Myocarditis	
Infective	
Viral (coxsackie, ^a adenovirus, ^a HIV, hepatitis C)	
Parasitic (<i>T. cruzi</i> —Chagas' disease, trypanosomiasis, toxoplasmosis)	
Bacterial (diphtheria)	
Spirochetal (<i>Borrelia burgdorferi</i> —Lyme disease)	
Rickettsial (Q fever)	
Fungal (with systemic infection)	
Noninfective	
Granulomatous inflammatory disease	
Sarcoidosis	
Giant cell myocarditis	
Eosinophilic myocarditis	
Polymyositis, dermatomyositis	
Collagen vascular disease	
Peripartum cardiomyopathy	
Transplant rejection	
Toxic	
Alcohol	
Catecholamines: amphetamines, cocaine	
Chemotherapeutic agents (anthracyclines, trastuzumab)	
Interferon	
Other therapeutic agents (hydroxychloroquine, chloroquine)	
Drugs of misuse (emetine, anabolic steroids)	
Heavy metals: lead, mercury	
Occupational exposure: hydrocarbons, arsenicals	
Metabolic^a	
Nutritional deficiencies: thiamine, selenium, carnitine	
Electrolyte deficiencies: calcium, phosphate, magnesium	
Endocrinopathy	
Thyroid disease	
Pheochromocytoma	
Diabetes	
Obesity	
Hemochromatosis	
Inherited Metabolic Pathway Defects^a	
Familial^a (See Table 287-3)	
Skeletal and cardiac myopathy	
Dystrophin-related dystrophy (Duchenne's, Becker's)	
Mitochondrial myopathies (e.g., Kearns-Sayre syndrome)	
Arrhythmogenic ventricular dysplasia	
Hemochromatosis	
Associated with other systemic diseases	
Susceptibility to immune-mediated myocarditis	
Overlap with Nondilated Cardiomyopathy	
"Minimally dilated cardiomyopathy"	
Hemochromatosis ^a	
Amyloidosis ^a	
Hypertrophic cardiomyopathy ^a ("burned-out")	
"Idiopathic"^a	
Miscellaneous (Shared Elements of Above Etiologies)	
Arrhythmogenic right ventricular dysplasia (may also affect left ventricle) ^a	
Left ventricular noncompaction ^a	
Peripartum cardiomyopathy	
Tachycardia-related cardiomyopathy	
Supraventricular arrhythmias with uncontrolled rate	
Very frequent nonsustained ventricular tachycardia or high premature ventricular complex burden	
Left bundle branch block (LBBB) has been implicated as a cause of dilated cardiomyopathy appearing late after idiopathic LBBB and responding with near-normal left ventricle size and function after cardiac resynchronization therapy.	

^aSome specific cases can be linked now to specific genetic mutation in a familial cardiomyopathy; others with similar phenotypes that appear to be acquired or idiopathic may represent genetic factors not yet identified.

The secondary acquired immune response is more specifically addressed against the viral proteins and can include both T-cell infiltration and antibodies to viral proteins. If unchecked, the acquired immune response can perpetuate secondary cardiac damage. Ongoing cytokine release activates matrix metalloproteinases that can disrupt the collagen and elastin scaffolding of the heart, potentiating ventricular dilation. Stimulation of profibrotic factors leads to pathologic interstitial fibrosis. Some of the antibodies triggered through co-stimulation or molecular mimicry also recognize targets within the host myocyte, such as the β -adrenergic receptor, troponin, and Na^+/K^+ ATPase, but it remains unclear whether these antibodies contribute actively to cardiac dysfunction in humans or merely serve as markers of cardiac injury.

It is not known how long the viruses persist in the human heart, whether late persistence of the viral genome continues to be deleterious, or how often a dormant virus can again become pathogenic. Genomes of common viruses have frequently been detected in patients with clinical diagnoses of myocarditis or dilated cardiomyopathy, but there is little information on how often these are present in patients without cardiac disease (see below). Further information is needed to understand the relative timing and contribution of infection, immune responses, and secondary adaptations in the progression of heart failure after viral myocarditis (Fig 287-5).

Clinical Presentation of Viral Myocarditis *Acute viral myocarditis* often presents with symptoms and signs of heart failure. Some patients present with chest pain suggestive of pericarditis or acute myocardial infarction. Occasionally, the presentation is dominated by atrial or ventricular tachyarrhythmias, or by pulmonary or systemic emboli from intracardiac thrombi. Electrocardiographic or echocardiographic abnormalities may also be detected incidentally during evaluation for other diagnoses. The typical patient with presumed viral myocarditis is a young to middle-aged adult who develops progressive dyspnea and weakness within a few days to weeks after a viral syndrome that was accompanied by fever and myalgias.

A small number of patients present with fulminant myocarditis, with rapid progression from a severe febrile respiratory syndrome to cardiogenic shock that may involve multiple organ systems, leading to renal failure, hepatic failure, and coagulopathy. These patients are typically young adults who have recently been dismissed from urgent care settings with antibiotics for bronchitis or oseltamivir for viral syndromes, only to return within a few days in rapidly progressive cardiogenic shock. Prompt triage is vital to provide aggressive support with high-dose intravenous catecholamine therapy and sometimes with temporary mechanical circulatory support. Recognition of patients with this fulminant presentation is potentially life-saving as more than half can survive, with marked improvement demonstrable within the first few weeks. The ejection fraction function of these patients often recovers to near-normal, although residual diastolic dysfunction may limit vigorous exercise for some survivors.

Chronic viral myocarditis is often invoked, but rarely proven, as a diagnosis when no other cause of dilated cardiomyopathy can be identified. However, some cases of otherwise unexplained cardiomyopathy will later be recognized to have a genetic basis, or ultimately found to have resulted from excess alcohol consumption or illicit drugs. There are likely many other causes that cannot yet be identified. The prevalence of previous or persistent viral infection as the cause for chronic dilated cardiomyopathy remains highly controversial.

Laboratory evaluation for myocarditis The initial evaluation for suspected myocarditis includes an ECG, an echocardiogram, and serum levels of troponin and creatine phosphokinase fractions. Magnetic resonance imaging is increasingly used for the diagnosis of myocarditis, which is supported by evidence of increased tissue edema and gadolinium enhancement (Fig. 287-6), particularly in the mid-wall (as distinct from usual coronary artery territories).

Endomyocardial biopsy is not often indicated for the initial evaluation of suspected viral myocarditis unless ventricular tachyarrhythmias suggest possible etiologies of sarcoidosis or giant cell myocarditis. The indications and benefit of endomyocardial biopsy for evaluation of myocarditis or new-onset cardiomyopathy remain controversial.

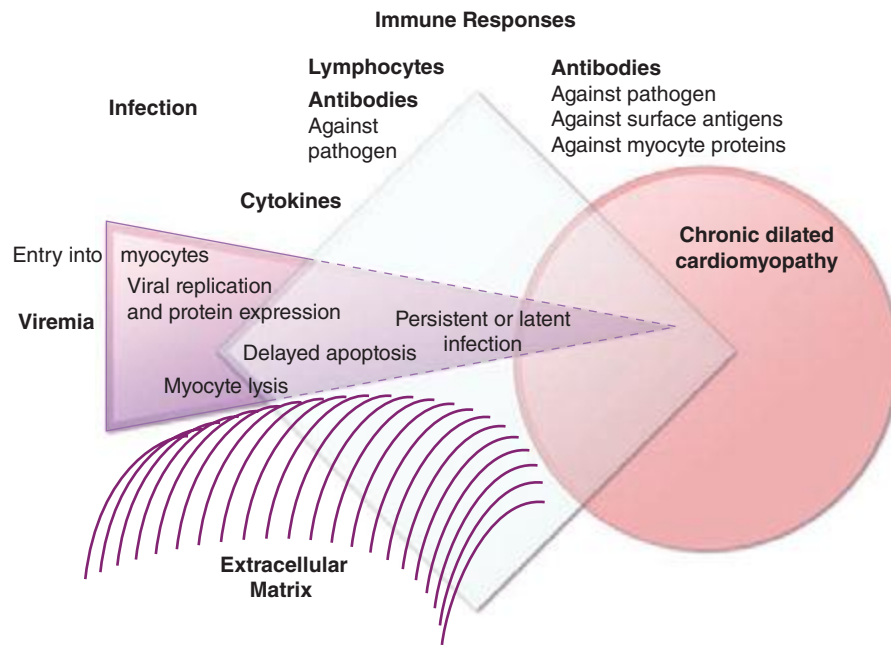


FIGURE 287-5 Schematic diagram demonstrating the possible progression from infection through direct, secondary, and autoimmune responses to dilated cardiomyopathy. Most of the supporting evidence for this sequence is derived from animal models. It is not known to what degree persistent infection and/or ongoing immune responses contribute to ongoing myocardial injury in the chronic phase.

The Dallas Criteria for myocarditis on endomyocardial biopsy include lymphocytic infiltrate with evidence of myocyte necrosis (Fig. 287-7) and are negative in 80–90% of patients with clinical myocarditis. Negative Dallas Criteria can reflect sampling error or early resolution of lymphocytic infiltrates, but also the insensitivity of the test when inflammation results from cytokines and antibody-mediated injury. Routine histologic examination of endomyocardial biopsy rarely reveals a specific infective etiology, such as toxoplasmosis or *Cytomegalovirus*. Immunohistochemistry of myocardial biopsy samples is commonly used to identify active lymphocyte subtypes and may also detect upregulation of HLA antigens and the presence of complement components attributed to inflammation, but the specificity and significance of these findings are uncertain.

An increase in circulating viral titers between acute and convalescent blood samples supports a diagnosis of acute viral myocarditis with potential spontaneous improvement. There is no established role for

measuring circulating anti-heart antibodies, which may be the result, rather than a cause, of myocardial injury and have been found also in patients with coronary artery disease and genetic cardiomyopathy.

Patients with recent or ongoing viral syndromes can be classified into three levels of diagnosis:

1. Possible subclinical acute myocarditis is diagnosed when a patient has a typical viral syndrome but *no* cardiac symptoms, with one or more of the following:
 - Elevated biomarkers of cardiac injury (troponin or CK-MB)
 - ECG findings suggestive of acute injury
 - Reduced left ventricular ejection fraction or regional wall motion
 - Abnormality on cardiac imaging, usually echocardiography

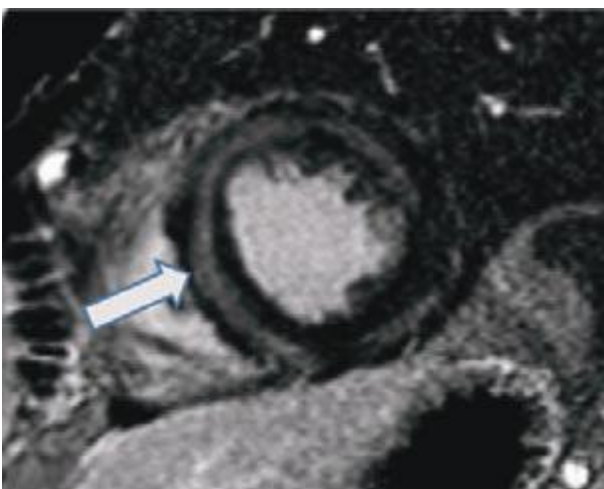


FIGURE 287-6 Magnetic resonance image of myocarditis showing the typical mid-wall location (arrow) for late gadolinium enhancement from cardiac inflammation and scarring. (Image courtesy of Ron Blankstein, MD, and Marcelo Di Carli, MD, Division of Nuclear Medicine, Brigham and Women's Hospital, Boston.)

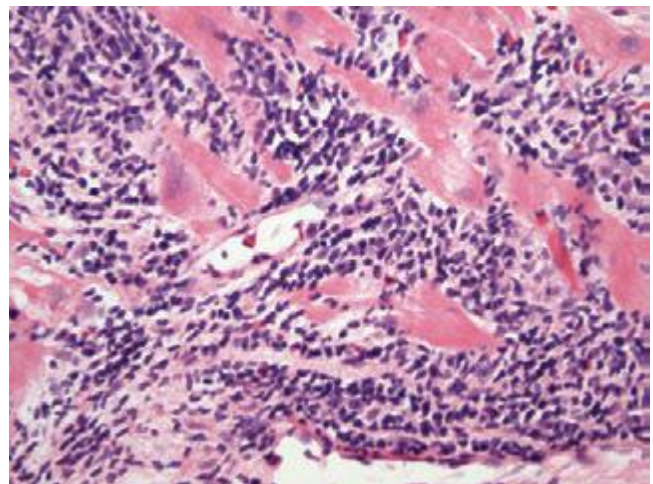


FIGURE 287-7 Acute myocarditis. Microscopic image of an endomyocardial biopsy showing massive infiltration with mononuclear cells and occasional eosinophils associated with clear myocyte damage. The myocyte nuclei are enlarged and reactive. Such extensive involvement of the myocardium would lead to extensive replacement fibrosis even if the inflammatory response could be suppressed. Hematoxylin and eosin–stained section, 200× original magnification. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

2. Probable acute myocarditis is diagnosed when the above criteria are met and accompanied also by cardiac symptoms, such as shortness of breath or chest pain, which can result from pericarditis or myocarditis. When clinical findings of pericarditis (pleuritic chest pain, ECG abnormalities, pericardial rub or effusion) are accompanied by elevated troponin or CK-MB or abnormal cardiac wall motion, the terms perimyocarditis or myopericarditis are sometimes used.
3. Definite myocarditis is diagnosed when there is histologic or immunohistologic evidence of inflammation on endomyocardial biopsy (see below) and does *not* require any other laboratory or clinical criteria.

SPECIFIC VIRUSES IMPLICATED IN MYOCARDITIS

In humans, viruses are often suspected but rarely proven to be the direct cause of clinical myocarditis. First implicated was the picornavirus family of RNA viruses, principally the enteroviruses, coxsackievirus, echovirus, and poliovirus. Influenza, another RNA virus, is implicated with varying frequency every winter and spring as epitopes change. Of the DNA viruses, adenovirus, vaccinia (smallpox vaccine), and the herpesviruses (varicella zoster, cytomegalovirus, Epstein-Barr virus, and human herpesvirus 6 [HHV6]) are well-recognized to cause myocarditis but also occur commonly in the healthy population. Polymerase chain reaction (PCR) detects viral genomes in the majority of patients with dilated cardiomyopathy, but also in normal “control” hearts. Most often detected are parvovirus B19 and HHV6, which may affect the cardiovascular system, in part, through infection of vascular endothelial cells. However, their contribution to chronic cardiomyopathy is uncertain, as serologic evidence of exposure is present in many children and most adults.

Human immunodeficiency virus (HIV) was associated with an incidence of dilated cardiomyopathy of 1–2%; however, with the advent of highly active antiretroviral therapy (HAART), HIV has been associated with a significantly lower incidence of cardiac disease. Cardiomyopathy in HIV may result from cardiac involvement with other associated viruses, such as cytomegalovirus and hepatitis C, as well as by HIV directly. Antiviral drugs to treat chronic HIV can cause cardiomyopathy, both directly and through drug hypersensitivity. The clinical picture may be complicated by pericardial effusions and pulmonary hypertension. There is a high frequency of lymphocytic myocarditis found at autopsy, and viral particles have been demonstrated in the myocardium in some cases, consistent with direct causation.

Hepatitis C has been repeatedly implicated in cardiomyopathy, particularly in Germany and Asia. Cardiac dysfunction may improve after interferon therapy. As this cytokine itself often depresses cardiac function transiently, careful coordination of administration and ongoing clinical evaluation are critical. Involvement of the heart with hepatitis B is uncommon, but can be seen when associated with systemic vasculitis (polyarteritis nodosa).

Additional viruses implicated specifically in myocarditis include *mumps*, *respiratory syncytial virus*, the *arboviruses* (*dengue fever* and *yellow fever*), and *arenaviruses* (*Lassa fever*). However, for any serious infection, the systemic inflammatory response can cause nonspecific depression of cardiac function, which is generally reversible if the patient survives.

THERAPY

There is currently no specific therapy recommended during any stage of viral myocarditis. During acute infection, therapy with anti-inflammatory or immunosuppressive medications is avoided, as their use has been shown to increase viral replication and myocardial injury in animal models. Therapy with specific antiviral agents (such as oseltamivir) has not been studied in relation to cardiac involvement. There is ongoing investigation into the impact of antiviral therapy to treat chronic viral persistence identified from endomyocardial biopsy. Large trials of immunosuppressive therapy for Dallas Criteria–positive myocarditis have been negative. There are some initial encouraging results and ongoing investigations with immunosuppressive therapy for immune-mediated myocarditis defined by immunohistologic criteria on biopsy or circulating anti-heart antibodies in the absence

of myocardial viral genomes. However, neither antiviral nor anti-inflammatory therapies are currently recommended. Until we have a better understanding of the different phases of viral myocarditis and its sequelae and the effects of timed or targeted therapies, treatment will continue to be directed to the clinical cardiovascular stage of the disease, for dilated cardiomyopathy in general.

Parasitic Myocarditis *Chagas’ disease* is the third most common parasitic infection in the world and the most common infective cause of cardiomyopathy. The protozoan *T. cruzi* is transmitted by the bite of the reduviid bug, endemic in the rural areas of South and Central America. Transmission can also occur through blood transfusion, organ donation, from mother to fetus, and occasionally orally. While programs to eradicate the insect vector have decreased the prevalence from about 16 million to less than 10 million in South America, cases are increasingly recognized in Western developed countries. Approximately 100,000 affected individuals are currently living in the United States, most of whom contracted the disease in endemic areas.

Multiple pathogenic mechanisms are implicated. The parasite itself can cause myocyte lysis and primary neuronal damage, and specific immune responses may recognize the parasites or related antigens and lead to chronic immune activation in the absence of detectable parasites. Molecular techniques have revealed persistent parasite DNA fragments in infected individuals. Further evidence for persistent infection is the eruption of parasitic skin lesions during immunosuppression after cardiac transplantation. As with viral myocarditis, the relative roles of persistent infection and of secondary autoimmune injury have not been resolved (Fig. 287-5). An additional factor in the progression of Chagas’ disease is the autonomic dysfunction and microvascular damage that may contribute to cardiac and gastrointestinal disease.

The acute phase of Chagas’ disease with parasitemia is usually unrecognized, but in fewer than 5% of cases, it presents clinically within a few weeks of infection, with nonspecific symptoms or occasionally with acute myocarditis and meningoencephalitis. In the absence of antiparasitic therapy, the silent stage progresses slowly over 10–30 years in almost half of patients to manifest in the cardiac and gastrointestinal systems in the chronic stages. Features typical of Chagas’ disease are conduction system abnormalities, particularly sinus node and atrioventricular (AV) node dysfunction and right bundle branch block. Atrial fibrillation and ventricular tachyarrhythmias also occur. Small ventricular aneurysms are common, particularly at the ventricular apex. These dilated ventricles are particularly thrombogenic, giving rise to pulmonary and systemic emboli. Xenodiagnosis, detection of the parasite itself, is rarely performed. The serologic tests for specific IgG antibodies against the trypanosome lack sufficient specificity and sensitivity, thereby requiring two separate positive tests required to make a diagnosis.

Treatment of the advanced stages focuses on clinical manifestations of the disease and includes heart failure medications, pacemaker-defibrillators, and anticoagulation. Increasing attention is directed to antiparasitic therapy even in chronic disease without obvious active infection. The most common effective antiparasitic therapies are benznidazole and nifurtimox, both associated with multiple severe reactions, including dermatitis, gastrointestinal distress, and neuropathy. Survival is less than 30% at 5 years after the onset of overt clinical heart failure. Patients without major extracardiac disease have occasionally undergone transplantation, after which they may require lifelong therapy to suppress reactivation of infection.

African trypanosomiasis infection results from the tsetse fly bite and can occur in travelers exposed during trips to Africa. The West African form is caused by *Trypanosoma brucei gambiense* and progresses silently over years. The East African form caused by *T. brucei rhodesiense* can progress rapidly through perivascular infiltration to myocarditis and heart failure, with frequent arrhythmias. The diagnosis is made by identification of trypanosomes in blood, lymph nodes, or other affected sites. Antiparasitic therapy has limited efficacy and is determined by the specific type and the stage of infection (hemolymphatic or neurologic).

Toxoplasmosis is contracted through undercooked infected beef or pork, transmission from feline feces, organ transplantation, transfusion, or maternal-fetal transmission. Immunocompromised hosts are most likely to experience reactivation of latent infection from cysts. The cysts have been found in up to 40% of autopsies of patients dying from HIV infection. Toxoplasmosis may present with encephalitis or chorioretinitis and, in the heart, can cause myocarditis, pericardial effusion, constrictive pericarditis, and heart failure. The diagnosis in an immunocompetent patient is made when the IgM is positive and the IgG becomes positive later. Active toxoplasmosis may be suspected in an immunocompromised patient with myocarditis and a positive IgG titer for toxoplasmosis, particularly when avidity testing identifies high specificity of the antibody. Fortuitous sampling occasionally reveals the cysts in the myocardium. Combination therapy can include pyrimethamine and sulfadiazine or clindamycin.

Trichinellosis is caused by *Trichinella spiralis* larva ingested with undercooked meat. Larvae migrating into skeletal muscles cause myalgias, weakness, and fever. Periorbital and facial edema and conjunctival and retinal hemorrhage may also be seen. Although the larva may occasionally invade the myocardium, clinical heart failure is rare and, when observed, attributed to the eosinophilic inflammatory response. The diagnosis is made from the specific serum antibody and is further supported by the presence of eosinophilia. Treatment includes antihelminthic drugs (albendazole, mebendazole) and glucocorticoids if inflammation is severe.

Cardiac involvement with *Echinococcus* is rare, but cysts can form and rupture in the myocardium and pericardium.

Bacterial Infections Most bacterial infections can involve the heart occasionally through direct invasion and abscess formation, but do so rarely. More commonly, systemic inflammatory responses depress contractility in severe infection and sepsis. *Diphtheria* specifically affects the heart in almost one-half of cases, and cardiac involvement is the most common cause of death in patients with this infection. The prevalence of vaccines has shifted the incidence of diphtheria from children worldwide to countries without routine immunization and to older populations who have lost their immunity. The bacillus releases a toxin that impairs protein synthesis and may particularly affect the conduction system. The specific antitoxin should be administered as soon as possible, with higher priority than antibiotic therapy. Other systemic bacterial infections that can involve the heart include *brucellosis*, *chlamydia*, *legionella*, *meningococcus*, *mycoplasma*, *psittacosis*, and *salmonellosis*, for which specific treatment is directed at the systemic infection.

Clostridial infections cause myocardial damage from the released toxin. Gas bubbles can be detected in the myocardium, and occasionally abscesses can form in the myocardium and pericardium. *Streptococcal infection* with β -hemolytic streptococci is most commonly associated with acute rheumatic fever and is characterized by inflammation and fibrosis of cardiac valves and systemic connective tissue, but it can also lead to a myocarditis with focal or diffuse infiltrates of mononuclear cells.

Tuberculosis can involve the myocardium directly as well as through tuberculous pericarditis, but rarely does so when the disease is treated with antibiotics. *Whipple's disease* is caused by *Tropheryma whippeli*. The usual manifestations are in the gastrointestinal tract, but pericarditis, coronary arteritis, valvular lesions, and occasionally clinical heart failure may also occur. Multidrug antituberculous regimens are effective, but the disease tends to relapse even with appropriate treatment.

Other Infections *Spirochetal myocarditis* has been diagnosed from myocardial biopsies containing *Borrelia burgdorferi* that causes *Lyme disease*. Lyme carditis most often presents with arthritis and conduction system disease that resolves within 1–2 weeks of antibiotic treatment, only rarely implicated in chronic heart failure.

Fungal myocarditis can occur due to hematogenous or direct spread of infection from other sites, as has been described for aspergillosis, actinomycosis, blastomycosis, candidiasis, coccidioidomycosis, cryptococcosis, histoplasmosis, and mucormycosis. However, cardiac involvement is rarely the dominant clinical feature of these infections.

The **rickettsial infections**, *Q fever*, *Rocky Mountain spotted fever*, and *scrub typhus* are frequently accompanied by ECG changes, but most clinical manifestations relate to systemic vascular involvement.

NONINFECTIVE MYOCARDITIS

Myocardial inflammation can occur without apparent preceding infection. The paradigm of noninfective inflammatory myocarditis is cardiac transplant rejection, from which we have learned that myocardial depression can develop and reverse quickly, that noncellular mediators such as antibodies and cytokines play a major role in addition to lymphocytes, and that myocardial antigens are exposed by prior physical injury and viral infection.

The most commonly diagnosed noninfective inflammation is granulomatous myocarditis, including both sarcoidosis and giant cell myocarditis. Sarcoidosis, as discussed in **Chap. 390**, is a multisystem disease most commonly affecting the lungs. Although classically presenting with higher prevalence in young African-American men, the epidemiology appears to be changing, with increasing recognition of sarcoidosis in Caucasian patients in nonurban areas. Patients with pulmonary sarcoid are at high risk for cardiac involvement, but cardiac sarcoidosis also occurs without clinical lung disease. Regional clustering of the disease supports the suspicion that the granulomatous reaction is triggered by an infectious or environmental allergen not yet identified.

The sites and density of cardiac granulomata, the time course, and the degree of extracardiac involvement are remarkably variable. Patients may present with rapid-onset heart failure and ventricular tachyarrhythmias, conduction block, chest pain syndromes, or minor cardiac findings in the setting of ocular involvement, an infiltrative skin rash, or a nonspecific febrile illness. They may also present less acutely after months to years of fluctuating cardiac symptoms. When ventricular tachycardia or conduction block dominates the initial presentation of heart failure without coronary artery disease, suspicion should be high for these granulomatous myocarditides.

Depending on the time course, the ventricles may appear restrictive or dilated. There is often right ventricular predominance of both dilation and ventricular arrhythmias, sometimes initially attributed to arrhythmogenic right ventricular dysplasia. Small ventricular aneurysms are common. Computed tomography of the chest often reveals pulmonary lymphadenopathy even in the absence of clinical lung disease. Metabolic imaging (positron emission tomography (PET)) of the whole chest can highlight active sarcoid lesions that are avid for glucose. Magnetic resonance imaging (MRI) of the heart can identify areas likely to be inflammatory. To rule out chronic infections, such as tuberculosis or histoplasmosis as the cause of adenopathy, the diagnosis usually requires pathologic confirmation. Biopsy of enlarged mediastinal nodes may provide the highest yield. The scattered granulomata of sarcoidosis can easily be missed on cardiac biopsy (**Fig. 287-8**).

Immunosuppressive treatment for sarcoidosis is initiated with high-dose glucocorticoids, which are often more effective for arrhythmias than for the heart failure. Patients with sarcoid lesions that persist or recur during tapering of corticosteroids are considered candidates for other immunosuppressive therapies, frequently with agents also used for cardiac transplantation. Pacemakers and implantable defibrillators are generally indicated to prevent life-threatening heart block or ventricular tachycardia, respectively. Because the inflammation often resolves into extensive fibrosis that impairs cardiac function and provides pathways for reentrant arrhythmias, the prognosis for improvement is best when the granulomata are not extensive and the ejection fraction is not severely reduced.

Giant cell myocarditis is less common than sarcoidosis, but accounts for 10–20% of biopsy-positive cases of myocarditis. Giant cell myocarditis typically presents with rapidly progressive heart failure and tachyarrhythmias. Diffuse granulomatous lesions are surrounded by extensive inflammatory infiltrate unlikely to be missed on endomyocardial biopsy, often with extensive eosinophilic infiltration. Associated conditions are thymomas, thyroiditis, pernicious anemia, other autoimmune diseases, and occasionally recent infections. Glucocorticoid therapy is less effective than for sarcoidosis and is

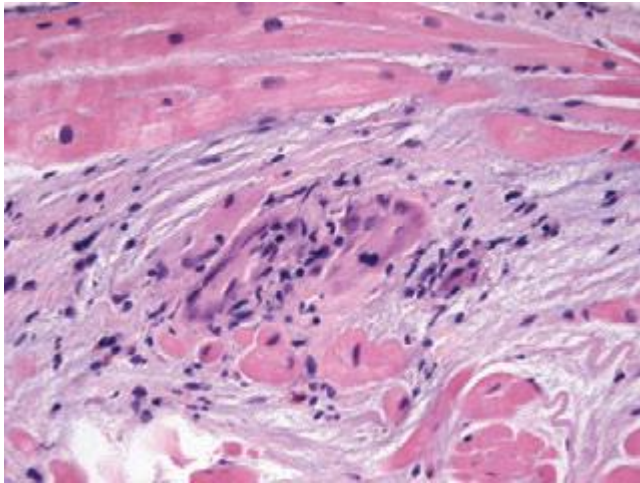


FIGURE 287-8 Sarcoidosis. Microscopic image of an endomyocardial biopsy showing a noncaseating granuloma and associated interstitial fibrosis typical of sarcoidosis. No microorganisms were present on special stains, and no foreign material was identified. Hematoxylin and eosin–stained section, 200× original magnification. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women’s Hospital, Boston.)

sometimes combined with other immunosuppressive agents. The course is generally of rapid deterioration requiring urgent transplantation. Although the severity of presentation and myocardial histology are more fulminant than with sarcoidosis, the occasional finding of giant cell myocarditis after sarcoidosis suggests that they may in some cases represent different stages of the same disease spectrum.

Eosinophilic myocarditis can be an important manifestation of the hypereosinophilic syndrome, which in Western countries is often idiopathic, although in Mediterranean and African countries, it is likely a consequence of antecedent infection. It may also be seen with systemic eosinophilic syndromes such as Churg–Strauss syndrome or malignancies. *Hypersensitivity myocarditis* is often an unexpected diagnosis, made when the biopsy reveals infiltration with lymphocytes and mononuclear cells with a high proportion of eosinophils. Most commonly, the reaction is attributed to antibiotics, particularly those taken chronically, but thiazides, anticonvulsants, indomethacin, and methyl dopa have also been implicated. Occasional associations with the smallpox vaccine have been reported. Although the circulating eosinophil count may be slightly elevated in hypersensitivity myocarditis, it does not reach the high levels of the hypereosinophilic syndrome. High-dose glucocorticoids and discontinuation of the trigger agent can be curative for hypersensitivity myocarditis.

Myocarditis is often associated with systemic inflammatory diseases, such as polymyositis and *dermatomyositis*, which affect skeletal and cardiac muscle. Although noninfective inflammatory myocarditis is sometimes included in the differential diagnosis of cardiac findings in patients with connective tissue disease such as systemic lupus erythematosus, pericarditis, vasculitis, pulmonary hypertension, and accelerated coronary artery disease are more common cardiac manifestations of connective tissue disease.

Peripartum cardiomyopathy (PPCM) develops during the last trimester or within the first 6 months after pregnancy, with a frequency between 1:2000 and 1:15,000 deliveries. Risk factors are increased maternal age, increased parity, twin pregnancy, malnutrition, use of tocolytic therapy for premature labor, and preeclampsia or toxemia of pregnancy. Heart failure early after delivery was previously common in Nigeria, when the custom for new mothers included salt ingestion while reclining on a warm bed, which likely impaired mobilization of the excess circulating volume after delivery. In the Western world, lymphocytic myocarditis has sometimes been found on myocardial biopsy. This inflammation has been hypothesized to reflect increased susceptibility to viral myocarditis or an autoimmune myocarditis due

to cross-reactivity of anti-uterine antibodies against cardiac muscle. Another proposed mechanism invokes an abnormal prolactin cleavage fragment, which is induced by oxidative stress and may trigger myocardial apoptosis; this observation has led to preliminary investigation of bromocriptine as possible therapy.

Very recently, PPCM has been found to be associated with increased antiangiogenic signaling, a process that is exacerbated by preeclampsia. In animal models of this disease, proangiogenic therapies have proven curative.

As the increased circulatory demand of pregnancy can aggravate other cardiac disease that was clinically unrecognized, it is crucial to the diagnosis of PPCM that there be no evidence for a preexisting cardiac disorder. By contrast, heart failure presenting earlier in pregnancy has been termed pregnancy-associated cardiomyopathy (PACM). Both PPCM and PACM have been found in some families with other presentations of dilated cardiomyopathy, in some cases with known sarcomeric protein mutations. Pregnancy may, thus, represent an environmental trigger for accelerated phenotypic expression of genetic cardiomyopathy.

TOXIC CARDIOMYOPATHY

Cardiotoxicity has been reported with multiple environmental and pharmacologic agents. Often these associations are seen only with very high levels of exposure or acute overdoses, in which acute electrocardiographic and hemodynamic abnormalities may reflect both direct drug effect and systemic toxicity.

Alcohol is the most common toxin implicated in chronic dilated cardiomyopathy. Excess consumption may contribute to more than 10% of cases of heart failure, including exacerbation of cases with other primary etiologies such as valvular disease or previous infarction. Toxicity is attributed both to alcohol and to its primary metabolite, acetaldehyde. Polymorphisms of the genes encoding alcohol dehydrogenase and the angiotensin-converting enzyme increase the likelihood of alcoholic cardiomyopathy in an individual with excess consumption. Superimposed vitamin deficiencies and toxic alcohol additives are rarely implicated. The alcohol consumption necessary to produce cardiomyopathy in an otherwise normal heart has been estimated to be five to six drinks (about 4 ounces of pure ethanol) daily for 5–10 years, but frequent binge drinking may also be sufficient. Many patients with alcoholic cardiomyopathy are fully functional in their daily lives without apparent stigmata of alcoholism. The cardiac impairment in severe alcoholic cardiomyopathy is the sum of both permanent damage and a substantial component that is reversible after cessation of alcohol consumption. Atrial fibrillation occurs commonly both early in the disease (“holiday heart”) and in advanced stages. Medical therapy includes neurohormonal antagonists and diuretics as needed for fluid management. Withdrawal should be supervised to avoid exacerbations of heart failure or arrhythmias, and ongoing support arranged. Even with severe disease, marked improvement can occur within 3–6 months of abstinence. Implantable defibrillators are generally deferred until an adequate period of abstinence, after which they may not be necessary if the ejection fraction has improved. With continued consumption, the prognosis is grim.

Cocaine, *amphetamines*, and related catecholaminergic stimulants can produce chronic cardiomyopathy as well as acute ischemia and tachyarrhythmias. Pathology reveals microinfarcts consistent with small vessel ischemia, similar to those seen with pheochromocytoma.

Chemotherapy agents are the most common drugs implicated in toxic cardiomyopathy. Judicious use of these drugs requires balancing the risks of the malignancy and the risks of cardiotoxicity, as many cancers have a chronic course with better prognosis than heart failure.

Anthracyclines cause characteristic histologic changes of vacuolar degeneration and myofibrillar loss. Generation of reactive oxygen species involving heme compounds is currently the favored explanation for myocyte injury and fibrosis. Disruption of the large titin protein may contribute to loss of sarcomere organization. Risk for cardiotoxicity increases with higher doses, preexisting cardiac disease, and concomitant chest irradiation. There are three different presentations of anthracycline-induced cardiomyopathy. (1) Heart failure can develop acutely during administration of a single dose, but may clinically resolve

in a few weeks. (2) Early-onset doxorubicin cardiotoxicity develops in about 3% of patients during or shortly after a chronic course, relating closely to total dose. It may be rapidly progressive, but may also resolve to good, but not normal, ventricular function. (3) The chronic presentation differs according to whether therapy was given before or after puberty. Patients who received doxorubicin while still growing may have impaired development of the heart, which leads to clinical heart failure by the time the patient reaches the early twenties. Late after adult exposure, patients may develop the gradual onset of symptoms or an acute onset precipitated by a reversible second insult, such as influenza or atrial fibrillation. Doxorubicin cardiotoxicity leads to a relatively nondilated ventricle, perhaps due to the accompanying fibrosis. Thus, the stroke volume may be severely reduced with an ejection fraction of 30–40%, which would be well tolerated in a patient with a larger ventricle typical of other cardiomyopathies with systolic dysfunction. Therapy includes angiotensin-converting enzyme inhibitors and β -adrenergic blocking agents used for other causes of heart failure, with careful suppression of “inappropriate” sinus tachycardia, and attention to postural hypotension that can occur in these patients. Once thought to have an inexorable downward course, some patients with doxorubicin cardiotoxicity improve under careful management to near-normal clinical function for many years.

Trastuzumab (Herceptin) is a monoclonal antibody that interferes with cell surface receptors crucial for some tumor growth and for cardiac adaptation. The incidence of cardiotoxicity is lower than for anthracyclines but enhanced by coadministration with them. Although considered to be more often reversible, trastuzumab cardiotoxicity does not always resolve, and some patients progress to clinical heart failure and death. As with anthracycline cardiotoxicity, therapy is as usual for heart failure, but it is not clear whether the spontaneous rate of improvement is enhanced by neurohormonal antagonists.

Cardiotoxicity with *cyclophosphamide* and *ifosfamide* generally occurs acutely and with very high doses. 5-Fluorouracil, cisplatin, and some other alkylating agents can cause recurrent coronary spasm that occasionally leads to depressed contractility. Acute administration of *interferon- α* can cause hypotension and arrhythmias. Clinical heart failure occurring during repeated chronic administration usually resolves after discontinuation.

Many small-molecule *tyrosine kinase inhibitors* are under development for different malignancies. Although these agents are “targeted” at specific tumor receptors or pathways, the biologic conservation of signaling pathways can cause these inhibitors to have “off-target” effects that include the heart and vasculature. Recognition of cardiotoxicity during therapy with these agents is complicated because they occasionally cause peripheral fluid accumulation (ankle edema, periorbital swelling, pleural effusions) due to local factors rather than elevated central venous pressures. Therapeutic approaches include withdrawal of the tyrosine kinase inhibitor (when possible) and substitution with a congener (when available), as well as conventional treatment for heart failure. Prophylactic treatment with beta blockers and angiotensin-converting enzyme inhibitors prior to and during chemotherapy is a topic of ongoing investigation.

Other therapeutic drugs that can cause cardiotoxicity during chronic use include hydroxychloroquine, chloroquine, emetine, and antiretroviral therapies.

Toxic exposures can cause arrhythmias or respiratory injury acutely during accidents. Chronic exposures implicated in cardiotoxicity include hydrocarbons, fluorocarbons, arsenicals, lead, and mercury.

METABOLIC CAUSES OF CARDIOMYOPATHY

Endocrine disorders affect multiple organ systems, including the heart. *Hyperthyroidism* and *hypothyroidism* do not often cause clinical heart failure in an otherwise normal heart, but commonly exacerbate heart failure. Clinical signs of thyroid disease may be masked, so tests of thyroid function are part of the routine evaluation of cardiomyopathy. Hyperthyroidism should always be considered with new-onset atrial fibrillation or ventricular tachycardia or atrial fibrillation in which the rapid ventricular response is difficult to control. The most common current reason for thyroid abnormalities in the cardiac

population is the treatment of tachyarrhythmias with amiodarone, a drug with substantial iodine content. Hypothyroidism should be treated with very slow escalation of thyroid supplements to avoid exacerbating tachyarrhythmias and heart failure. Hyperthyroidism and heart failure are a dangerous combination that merits very close supervision, often hospitalization, during titration of antithyroid medications, during which decompensation of heart failure may occur precipitously and fatally.

Pheochromocytoma is rare, but should be considered when a patient has heart failure and very labile blood pressure and heart rate, sometimes with episodic palpitations (Chap. 407). Patients with pheochromocytoma often have postural hypotension. In addition to α -adrenergic receptor antagonists, definitive therapy requires surgical extirpation. Very high renin states, such as those caused by renal artery stenosis, can lead to modest depression in ejection fraction with little or no ventricular dilation and markedly labile symptoms with flash pulmonary edema, related to sudden shifts in vascular tone and intravascular volume.

Controversies remain regarding whether *diabetes* and *obesity* are sufficient to cause cardiomyopathy. Most heart failure in diabetes results from epicardial coronary disease, with further increase in coronary artery risk due to accompanying hypertension and renal dysfunction. Cardiomyopathy may result in part from insulin resistance and increased advanced-glycosylation end products, which impair both systolic and diastolic function. However, much of the dysfunction can be attributed to scattered focal ischemia resulting from distal coronary artery tapering and limited microvascular perfusion even without proximal focal stenoses. Diabetes is a typical factor in heart failure with “preserved” ejection fraction, along with hypertension, advanced age, and female gender.

The existence of a cardiomyopathy due to *obesity* is generally accepted. In addition to cardiac involvement from associated diabetes, hypertension, and vascular inflammation of the metabolic syndrome, obesity alone is associated with impaired excretion of excess volume load, which, over time, can lead to increased wall stress and secondary adaptive neurohumoral responses. Fluid retention may be aggravated by large fluid intake and the rapid clearance of natriuretic peptides by adipose tissue. In the absence of another obvious cause of cardiomyopathy in an obese patient with systolic dysfunction without marked ventricular dilation, effective weight reduction is often associated with major improvement in ejection fraction and clinical function. Improvement in cardiac function has been described after successful bariatric surgery, although all major surgical therapy poses increased risk for patients with heart failure. Postoperative malabsorption and nutritional deficiencies, such as calcium and phosphate deficiencies, may be particularly deleterious for patients with cardiomyopathy.

Nutritional deficiencies can occasionally cause dilated cardiomyopathy but are not commonly implicated in developed Western countries. *Beri-beri heart disease* due to thiamine deficiency can result from poor nutrition in undernourished populations and in patients deriving most of their calories from alcohol, and has been reported in teenagers subsisting only on highly processed foods. This disease is initially a vasodilated state with very high output heart failure that can later progress to a low output state; thiamine repletion can lead to prompt recovery of cardiovascular function. Abnormalities in *carnitine* metabolism can cause dilated or restrictive cardiomyopathies, usually in children. Deficiency of trace elements such as *selenium* can cause cardiomyopathy (Keshan’s disease).

Calcium is essential for excitation-contraction coupling. Chronic deficiencies of calcium, such as can occur with hypoparathyroidism (particularly postsurgical) or intestinal dysfunction (from diarrheal syndromes and following extensive resection), can cause severe chronic heart failure that responds over days or weeks to vigorous calcium repletion. *Phosphate* is a component of high-energy compounds needed for efficient energy transfer and multiple signaling pathways. *Hypophosphatemia* can develop during starvation and early refeeding following a prolonged fast, and occasionally during hyperalimentation. *Magnesium* is a cofactor for thiamine-dependent reactions and for the sodium-potassium adenosine triphosphatase (ATPase), but

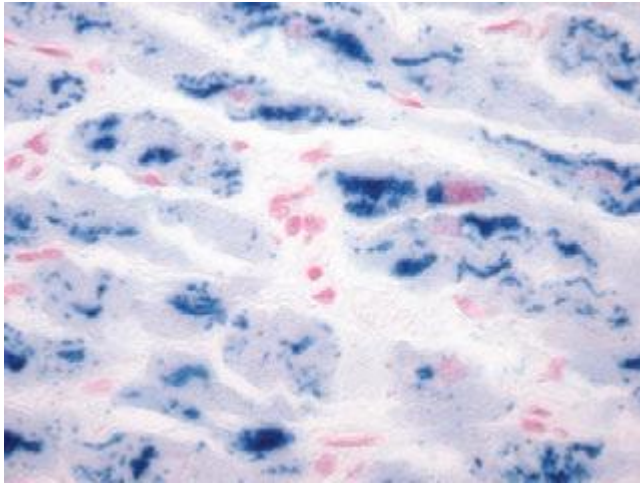


FIGURE 287-9 Hemochromatosis. Microscopic image of an endomyocardial biopsy showing extensive iron deposition within the cardiac myocytes with the Prussian blue stain (400 \times original magnification). (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

hypomagnesemia rarely becomes sufficiently profound to cause clinical cardiomyopathy.

Hemochromatosis is variably classified as a metabolic or storage disease (Chap. 428). It is included among the causes of restrictive cardiomyopathy, but the clinical presentation is often that of a dilated cardiomyopathy. The autosomal recessive form is related to the *HFE* gene. With up to 10% of the population heterozygous for one mutation, the clinical prevalence might be as high as 1 in 500. The lower observed rates highlight the limited penetrance of the disease, suggesting the role of additional genetic and environmental factors for clinical expression. Hemochromatosis can also be acquired from iron overload due to hemolytic anemia and transfusions. Excess iron is deposited in the perinuclear compartment of cardiomyocytes, with resulting disruption of intracellular architecture and mitochondrial function. Diagnosis is easily made from measurement of serum iron and transferrin saturation, with a threshold of >60% for men and >45–50% for women. MRI can help to quantitate iron stores in the liver and heart, and endomyocardial biopsy tissue can be stained for iron (Fig. 287-9), which is particularly important if the patient has another cause for cardiomyopathy. If diagnosed early, hemochromatosis can often be managed by repeated phlebotomy to remove iron. For more severe iron overload, iron chelation therapy with desferrioxamine (deferrioxamine) or deferasirox can help to improve cardiac function if myocyte loss and replacement fibrosis are not too severe.

Inborn disorders of metabolism occasionally present with dilated cardiomyopathy, although they are most often associated with restrictive cardiomyopathy (Table 287-4).

FAMILIAL DILATED CARDIOMYOPATHY

The genetic basis for cardiomyopathy is discussed in the section, “Genetic Etiologies of Cardiomyopathy.” The recognized frequency of familial involvement in dilated cardiomyopathy has increased to over 30%. Mutations in *TTN*, encoding the giant sarcomeric protein titin, are the most common cause of dilated cardiomyopathy, accounting for up to 25% of familial disease. On average, men with *TTN* mutations develop cardiomyopathy a decade before women, without distinctive clinical features. Mutations in thick and thin filament genes account for ~8% of dilated cardiomyopathy and may manifest in early childhood.

The most recognizable familial cardiomyopathy syndromes with extracardiac manifestations are the *muscular dystrophies*. Both Duchenne's and the milder Becker's dystrophy result from abnormalities in the X-linked dystrophin gene of the sarcolemmal membrane. Skeletal myopathy is present in multiple other genetic cardiomyopathies (Table 287-3), some of which are associated with creatine kinase elevations.

Families with a history of atrial arrhythmias, conduction system disease, and cardiomyopathy may have abnormalities of the nuclear membrane lamin proteins. While all dilated cardiomyopathies carry a risk of sudden death, a family history of cardiomyopathy with sudden death raises suspicion for a particularly arrhythmogenic mutation; affected family members may be considered for implantable defibrillators even before meeting the reduced ejection fraction threshold for primary prevention of sudden death.

A prominent family history of sudden death or ventricular tachycardia before clinical cardiomyopathy suggests genetic defects in the desmosomal proteins (Fig. 287-10). Originally described as affecting the right ventricle (arrhythmogenic right ventricular dysplasia [ARVD]), this disorder (arrhythmogenic ventricular dysplasia) can affect either or both ventricles. Patients often present first with ventricular tachycardia. Genetic defects in proteins of the desmosomal complex disrupt myocyte junctions and adhesions, leading to replacement of myocardium by deposits of fat. Thin ventricular walls may be recognized on echocardiography but are better visualized on MRI. Because desmosomes are also important for elasticity of hair and skin, some of the defective desmosomal proteins are associated with striking “woolly hair” and thickened skin on the palms and soles. Implantable defibrillators are usually indicated to prevent sudden death. There is variable progression to right, left, or biventricular failure.

Left ventricular noncompaction is a condition of unknown prevalence that is increasingly revealed with the refinement of imaging techniques. The diagnostic criteria include the presence of multiple trabeculations in the left ventricle distal to the papillary muscles, creating a “spongy” appearance of the apex. Noncompaction has been associated with multiple genetic variants in the sarcomeric and other genes, such as *TAZ* (encoding tafazzin). The diagnosis may be made incidentally or in patients previously diagnosed with cardiomyopathy, in whom the criteria for noncompaction may appear and resolve with changing left ventricular size and function. The three cardinal clinical features are ventricular arrhythmias, embolic events, and heart failure. Treatment generally includes anticoagulation and early consideration for an implantable defibrillator, in addition to neurohormonal antagonists as indicated by stage of disease.

Some families inherit a susceptibility to viral-induced myocarditis. This propensity may relate to abnormalities in cell surface receptors, such as the coxsackie-adenovirus receptor, that bind viral proteins.

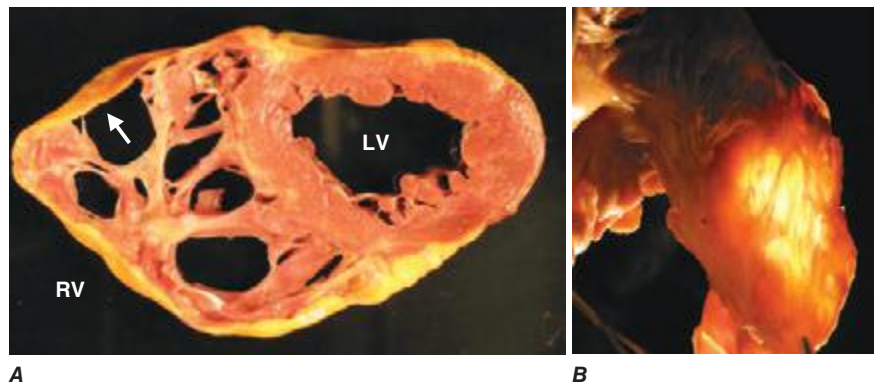


FIGURE 287-10 Arrhythmogenic right ventricular dysplasia. **A.** Cross-sectional slice of a pathology specimen removed at transplantation, showing severe dysplasia of the right ventricle (RV) with extensive fatty replacement of right ventricular myocardium. **B.** The remarkably thin right ventricular free wall is revealed by transillumination. LV, left ventricle. (Images courtesy of Gayle Winters, MD, and Richard Mitchell, MD, PhD, Division of Pathology, Brigham and Women's Hospital, Boston.)

Some may have partial homology with viral proteins such that an autoimmune response is triggered against the myocardium.

Prognosis and therapy of familial dilated cardiomyopathy are dictated primarily by the stage of clinical disease and the risk for sudden death. In some cases, the familial etiology facilitates prognostic decisions, particularly regarding the likelihood of recovery after a new diagnosis, which is unlikely for familial disease. The rate of progression of disease, once manifest, is, to some extent, heritable, although marked variation can be seen. However, there have been cases of remarkable clinical remission after acute presentation, likely after a reversible additional insult, such as prolonged tachycardia or infective myocarditis.

TAKOTSUBO CARDIOMYOPATHY

The apical ballooning syndrome, or stress-induced cardiomyopathy, occurs typically in older women after sudden intense emotional or physical stress. The ventricle shows global ventricular dilation with basal contraction, forming the shape of the narrow-necked jar (*takotsubo*) used in Japan to trap octopi. Originally described in Japan, it is increasingly recognized elsewhere during emergency cardiac catheterization and intensive care unit admissions for noncardiac conditions. Presentations include pulmonary edema, hypotension, and chest pain with ECG changes mimicking an acute infarction. The left ventricular dysfunction extends beyond a specific coronary artery distribution and generally resolves within days to weeks. Animal models and ventricular biopsies suggest that this acute cardiomyopathy may result from intense sympathetic activation with heterogeneity of myocardial autonomic innervation, diffuse microvascular spasm, and/or direct catecholamine toxicity. Coronary angiography may be required to rule out acute coronary occlusion. No therapies have been proven beneficial, but reasonable strategies include nitrates for pulmonary edema, intraaortic balloon pump if needed for low output, combined alpha and beta blockers rather than selective beta blockade if hemodynamically stable, and magnesium for arrhythmias related to QT prolongation. Anticoagulation is generally withheld due to the occasional occurrence of ventricular rupture. While the prognosis is generally good, recurrences have been described in up to 10% of patients.

IDIOPATHIC DILATED CARDIOMYOPATHY

Idiopathic dilated cardiomyopathy is a diagnosis of exclusion, when all other known factors have been excluded. Approximately two-thirds of dilated cardiomyopathies are still labeled as idiopathic; however, a substantial proportion of these may reflect unrecognized genetic disease. Continued reconsideration of etiology during chronic heart failure management often reveals specific causes later in a patient's course.

OVERLAPPING TYPES OF CARDIOMYOPATHY

The limitations of our phenotypic classification are revealed through the multiple overlaps between the etiologies and presentations of the three types. Cardiomyopathy with reduced systolic function but without severe dilation can represent early dilated cardiomyopathy, "minimally dilated cardiomyopathy," or restrictive diseases without marked increases in ventricular wall thickness. For example, sarcoidosis and hemochromatosis can present as dilated or restrictive disease. Early stages of amyloidosis are often mistaken for hypertrophic cardiomyopathy. Progression of hypertrophic cardiomyopathy into a "burned-out" phase occurs occasionally, with decreased contractility and modest ventricular dilation. Overlaps are particularly common with the inherited metabolic disorders, which can present as any of the three major phenotypes (Fig. 287-4).

DISORDERS OF METABOLIC PATHWAYS

Multiple genetic disorders of metabolic pathways can cause myocardial disease, due to infiltration of abnormal products or cells containing them between the myocytes, and storage disease, due to their accumulation within cells (see HPIM 18e, Table 238-4, and 287-5). The restrictive phenotype is most common, but mildly dilated cardiomyopathy may occur. Hypertrophic cardiomyopathy may be mimicked by the myocardium thickened with these abnormal products causing "pseudohypertrophy." Most of these diseases are diagnosed during childhood.

TABLE 287-5 CAUSES OF RESTRICTIVE CARDIOMYOPATHIES

Infiltrative (Between Myocytes)
Amyloidosis
Primary (light chain amyloid)
Familial (abnormal transthyretin) ^a
Senile (normal transthyretin or atrial peptides)
Inherited metabolic defects ^a
Storage (Within Myocytes)
Hemochromatosis (iron) ^a
Inherited metabolic defects ^a
Fabry's disease
Glycogen storage disease (II, III)
Fibrotic
Radiation
Scleroderma
Endomyocardial
Possibly related fibrotic diseases
Tropical endomyocardial fibrosis
Hypereosinophilic syndrome (Löffler's endocarditis)
Carcinoid syndrome
Radiation
Drugs: e.g., serotonin, ergotamine
Overlap with Other Cardiomyopathies
Hypertrophic cardiomyopathy/"pseudohypertrophic" ^a
"Minimally dilated" cardiomyopathy
Early-stage dilated cardiomyopathy
Partial recovery from dilated cardiomyopathy
Sarcoidosis
Idiopathic ^a

^aCan be familial.

Fabry's disease results from a deficiency of the lysosomal enzyme alpha-galactosidase A caused by one of more than 160 mutations. This disorder of glycosphingolipid metabolism is an X-linked recessive disorder that may also cause clinical disease in female carriers. Glycolipid accumulation may be limited to the cardiac tissues or may also involve the skin and kidney. Electron microscopy of endomyocardial biopsy tissue shows diagnostic vesicles containing concentric lamellar figures (Fig. 287-11). Diagnosis is crucial because enzyme replacement can reduce abnormal deposits and improve cardiac and clinical function. The magnitude of clinical impact has not been well-established for this therapy, which requires frequent infusions of the enzyme at a cost of over \$100,000 a year. Enzyme replacement can also improve the course of Gaucher's disease, in which cerebroside-rich cells accumulate in multiple organs due to a deficiency of beta-glucosidase. Cerebroside-rich cells infiltrate the heart, which can also lead to a hemorrhagic pericardial effusion and valvular disease.

Glycogen storage diseases lead to accumulation of lysosomal storage products and intracellular glycogen accumulation, particularly with *glycogen storage disease type III*, due to a defective debranching enzyme. There are more than 10 types of *mucopolysaccharidoses*, in which autosomal dominant or X-linked deficiencies of lysosomal enzymes lead to the accumulation of glycosaminoglycans in the skeleton, nervous system, and occasionally the heart. With characteristic facies, short stature, and frequent cognitive impairment, most individuals are diagnosed early in childhood and die before adulthood.

Carnitine is an essential cofactor in long-chain fatty acid metabolism. Multiple defects have been described that lead to carnitine deficiency, causing intracellular lipid inclusions and restrictive or dilated cardiomyopathy, often presenting in children. Fatty acid oxidation requires many metabolic steps with specific enzymes that can be deficient, with complex interactions with carnitine. Depending on the defect, cardiac and skeletal myopathy can be ameliorated with replacement of fatty acid intermediates and carnitine.

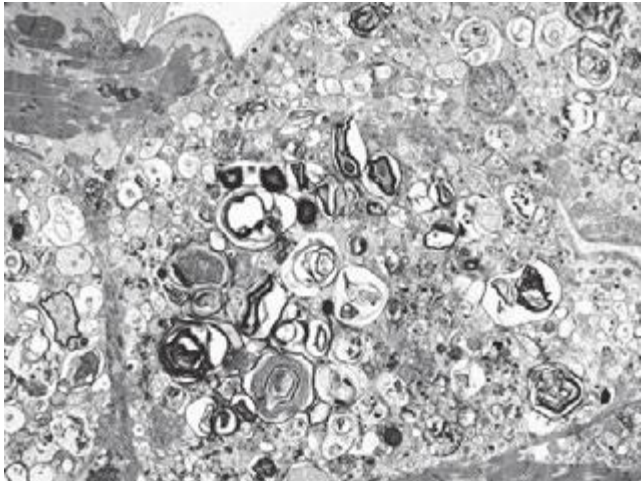


FIGURE 287-11 Fabry's disease. Transmission electron micrograph of a right ventricular endomyocardial biopsy specimen at high magnification showing the characteristic concentric lamellar inclusions of glycosphingolipids accumulating as a result of deficiency of the lysosomal enzyme alpha-galactosidase A. Image taken at 15,000 \times original magnification. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

Two monogenic metabolic cardiomyopathies have recently been described as causes of increased ventricular wall thickness without an increase of muscle subunits or an increase in contractility. Mutations in the gamma-2 regulatory subunit of the adenosine monophosphate (AMP)-activated protein kinase important for glucose metabolism (*PRKAG2*) have been associated with a high prevalence of conduction abnormalities, such as AV block and ventricular preexcitation (Wolff-Parkinson-White syndrome). Several defects have been reported in an X-linked lysosome-associated membrane protein (*LAMP2*). This defect can be maternally transmitted or sporadic and has occasionally been isolated to the heart, although it often leads to a syndrome of skeletal myopathy, mental retardation, and hepatic dysfunction referred to as *Danon's disease*. Extreme left ventricular hypertrophy appears early, often in childhood, and can progress rapidly to end-stage heart failure with low ejection fraction. Electron microscopy of these metabolic disorders shows that the myocytes are enlarged by multiple intracellular vacuoles of metabolic by-products.

RESTRICTIVE CARDIOMYOPATHY

The least common of the physiologic triad of cardiomyopathies is restrictive cardiomyopathy, which is dominated by abnormal diastolic function, often with mildly decreased contractility and ejection fraction (usually >30–50%). Both atria are enlarged, sometimes massively. Modest left ventricular dilation can be present, usually with an end-diastolic dimension <6 cm. End-diastolic pressures are elevated in both ventricles, with preservation of cardiac output until late in the disease. Subtle exercise intolerance is usually the first symptom but is often not recognized until after clinical presentation with congestive symptoms. The restrictive diseases often present with relatively more right-sided symptoms, such as edema, abdominal discomfort, and ascites, although filling pressures are elevated in both ventricles. The cardiac impulse is less displaced than in dilated cardiomyopathy and less dynamic than in hypertrophic cardiomyopathy. A fourth heart sound is more common than a third heart sound in sinus rhythm, but atrial fibrillation is common. Jugular venous pressures often show rapid Y descents and may increase during inspiration (positive Kussmaul's sign). Most restrictive cardiomyopathies are due to infiltration of abnormal substances between myocytes, storage of abnormal metabolic products within myocytes, or fibrotic injury (Table 287-5). The differential diagnosis should include constrictive pericardial disease, which may also be dominated by right-sided heart failure.

INFILTRATIVE DISEASE

Amyloidosis is the major cause of restrictive cardiomyopathy (Figs. 287-12, 287-13, and 287-14). Several proteins can self-assemble to form the beta-sheets of amyloid proteins, which deposit with different consequences depending on the type of protein. The systemic amyloidoses are discussed in Chap. 137. In addition to cardiac infiltration, neurologic involvement occurs commonly with primary amyloidosis (immunoglobulin light chains) and with familial amyloidosis (genetic abnormalities of transthyretin). There are over 100 identified mutations in transthyretin on chromosome 13, among which the V122I transthyretin mutation has been identified in about 4% of African Americans and in 10% of African Americans with heart failure and may contribute importantly to heart failure in general in the elderly African-American population. Organ dysfunction was previously attributed solely to physical disruption from the infiltrating amyloid fibrils, but newer information suggests additional direct toxicity from the immunoglobulin light chain and abnormal transthyretin protein aggregates themselves. In senile amyloidosis, there is abnormal accumulation of normal transthyretin or natriuretic peptide folding, detected in 10% of people over 80 years and half of those over 90 years but often without apparent clinical disease. Men show a greater burden of amyloid deposition and 20-fold greater likelihood of clinical disease with senile amyloidosis. The aging of the population will soon render senile amyloidosis the most common of the amyloidoses.

Cardiac amyloid is classically suspected from thickened ventricular walls with an ECG that shows low voltage. However, low voltage is not always present and is less common in familial or senile amyloidosis than in primary AL amyloidosis. A characteristic refractile brightness in the septum on echocardiography is suggestive of the diagnosis, but neither sensitive nor specific. Both atria are dilated, often dramatically, and diastolic dysfunction may be more obvious than in left ventricular hypertrophy from other causes. Amyloid infiltration can also be detected with gadolinium enhancement in MRI.



FIGURE 287-12 Restrictive cardiomyopathy—amyloidosis. Gross specimen of a heart with amyloidosis. The heart is firm and rubbery with a waxy cut surface. The atria are markedly dilated, and the left atrial endocardium, normally smooth, has yellow-brown amyloid deposits that give texture to the surface. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

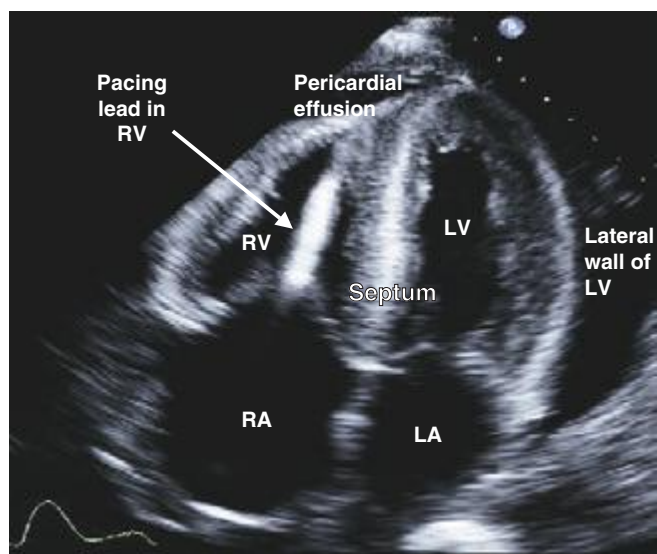


FIGURE 287-13 Restrictive cardiomyopathy—amyloidosis.

Echocardiogram showing thickened walls of both ventricles without major chamber dilation. The atria are markedly dilated, consistent with chronically elevated ventricular filling pressures. In this example, there is a characteristic hyperrefractile “glittering” of the myocardium typical of amyloid infiltration, which is often absent (especially with more recent echocardiographic systems of better resolution). The mitral and tricuspid valves are thickened. A pacing lead is visible in the right ventricle (RV), and a pericardial effusion is evident. Note that the echocardiographic and pathologic images are vertically opposite, such that the left ventricle (LV) is by convention on the top right in the echocardiographic image and bottom right in the pathologic images. LA, left atrium; RA, right atrium. (Image courtesy of Justina Wu, MD, Brigham and Women’s Hospital, Boston.)

The diagnosis of primary or familial amyloidosis can sometimes be made from biopsies of an abdominal fat pad or the rectum, but cardiac amyloidosis is most reliably identified from a biopsy of the heart, in which amyloid fibrils infiltrate the myocardium diffusely, particularly around the conduction system and coronary vessels (Fig. 287-14). Diagnosis of the type of amyloid protein requires immunohistochemistry of biopsied tissue rather than serum or urine electrophoresis, which can lead to incorrect classification.

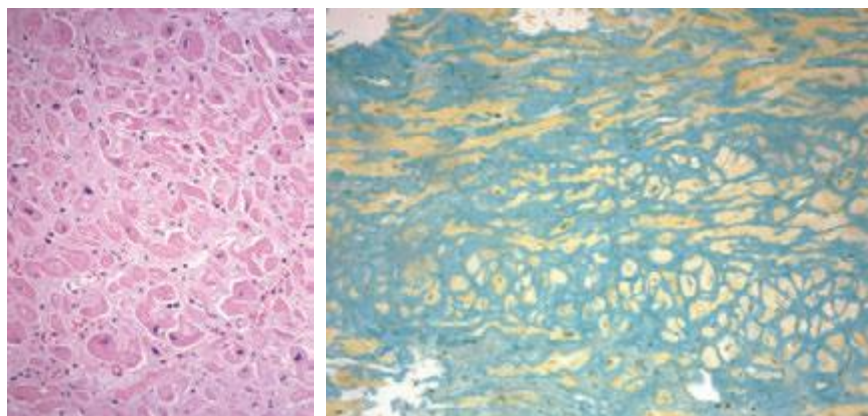


FIGURE 287-14 Amyloidosis—microscopic images of amyloid involving the myocardium. The left panel (hematoxylin and eosin stain) shows glassy, grey-pink amorphous material infiltrating between cardiomyocytes, which stain a darker pink. The right panel shows a sulfated blue stain that highlights the amyloid green and stains the cardiac myocytes yellow. (The Congo red stain can also be used to highlight amyloid; under polarized light, amyloid will have an apple-green birefringence when stained with Congo red.) Images at 100× original magnification. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women’s Hospital, Boston.)

Therapy for all types of amyloid is predominantly for symptoms of fluid retention, which often requires high doses of loop diuretics. Digoxin bound to the amyloid fibrils can reach toxic levels, and should therefore be used only in very low doses, if at all. There is no evidence regarding use of neurohormonal antagonists in amyloid heart disease, where the possible theoretical benefit has to be balanced against the possibility of aggravating postural hypotension and diminishing the crucial heart rate reserve. The risk of intracardiac thrombi may warrant chronic anticoagulation.

The prognosis is worst for primary amyloid, with a median survival of 6–12 months after symptoms of heart failure. If present, multiple myeloma is treated with chemotherapy, the extent of which is often limited by the potential of worsening cardiac dysfunction. Immunoglobulin-associated amyloid has occasionally been treated with sequential heart transplantation and delayed bone marrow transplant, with frequent recurrence of amyloid in the transplanted heart. Abnormal transthyretin-associated cardiac amyloid has a somewhat better prognosis and can be treated in selected patients with heart and liver transplantation. Senile cardiac amyloid has the slowest progression and best overall prognosis.

FIBROTIC RESTRICTIVE CARDIOMYOPATHY

Progressive fibrosis can cause restrictive myocardial disease without ventricular dilation. Thoracic radiation, common for breast and lung cancer or mediastinal lymphoma, can produce early or late restrictive cardiomyopathy. Patients with *radiation cardiomyopathy* may present with a possible diagnosis of constrictive pericarditis, as the two conditions often coexist. Careful hemodynamic evaluation and, often, endomyocardial biopsy should be performed if considering pericardial stripping surgery, which is unlikely to be successful in the presence of underlying restrictive cardiomyopathy.

Scleroderma causes small vessel spasm and ischemia that can lead to a small, stiff heart with reduced ejection fraction without dilation. The pulmonary hypertension associated with scleroderma may lead to more clinical right heart failure because of concomitant fibrotic disease of the right ventricle. Doxorubicin causes direct myocyte injury usually leading to dilated cardiomyopathy, but the limited degree of dilation may result from increased fibrosis, which restricts remodeling.

ENDOMYOCARDIAL DISEASE

The physiologic picture of elevated filling pressures with atrial enlargement and preserved ventricular contractility with normal or reduced ventricular volumes can result from extensive fibrosis of the endocardium, without transmural myocardial disease. For patients who have not lived in the equatorial regions, this picture is rare, and when seen is often associated with a history of chronic hypereosinophilic syndrome (*Löffler’s endocarditis*), which is more common in men than women. In this disease, persistent hypereosinophilia of >1500 eos/mm³ for at least 6 months can cause an acute phase of eosinophilic injury in the endocardium (see earlier discussion of eosinophilic myocarditis), with systemic illness and injury to other organs. There is usually no obvious cause, but the hypereosinophilia can occasionally be explained by allergic, parasitic, or malignant disease. It is postulated to be followed by a period in which cardiac inflammation is replaced by evidence of fibrosis with superimposed thrombosis. In severe disease, the dense fibrotic layer can obliterate the ventricular apices and extend to thicken and tether the AV valve leaflets. The clinical disease may present with heart failure, embolic events, and atrial arrhythmias. While plausible, the sequence of transition from eosinophilic myocarditis or *Löffler’s endocarditis* to endomyocardial fibrosis has not been clearly demonstrated.

In tropical countries, up to one-quarter of heart failure may be due to *endomyocardial fibrosis*,

1568 affecting either or both ventricles. This condition shares with the previous condition the partial obliteration of the ventricular apex with fibrosis extending into the valvular inflow tract and leaflets; however, it is not clear that the etiologies are the same for all cases. Pericardial effusions frequently accompany endomyocardial fibrosis but are not common in Löffler's endocarditis. For endomyocardial fibrosis, there is no gender difference, but a higher prevalence in African-American populations. While tropical endomyocardial fibrosis could represent the end-stage of previous hypereosinophilic disease triggered by endemic parasites, neither prior parasitic infection nor hypereosinophilia is usually documented. Geographic nutritional deficiencies have also been proposed as an etiology.

Medical treatment focuses on glucocorticoids and chemotherapy to suppress hypereosinophilia when present. Fluid retention may become increasingly resistant to diuretic therapy. Anticoagulation is recommended. Atrial fibrillation is associated with worse symptoms and prognosis, but may be difficult to suppress. Surgical resection of the apices and replacement of the fibrotic valves can improve symptoms, but surgical morbidity and mortality and later recurrence rates are high.

The serotonin secreted by *carcinoid* tumors can produce fibrous plaques in the endocardium and right-sided cardiac valves, occasionally affecting left-sided valves, as well. Valvular lesions may be stenotic or regurgitant. Systemic symptoms include flushing and diarrhea. Liver disease from hepatic metastases may play a role by limiting hepatic function and thereby allowing more serotonin to reach the venous circulation.

HYPERTROPHIC CARDIOMYOPATHY

Hypertrophic cardiomyopathy is defined as left ventricular hypertrophy that develops in the absence of causative hemodynamic factors, such as hypertension, aortic valve disease, or systemic infiltrative or storage diseases (Figs. 287-15 and 287-16). It has previously been termed *hypertrophic obstructive cardiomyopathy* (HOCM), *asymmetric septal hypertrophy* (ASH), and *idiopathic hypertrophic subaortic stenosis* (IHSS). However, the accepted terminology is now hypertrophic

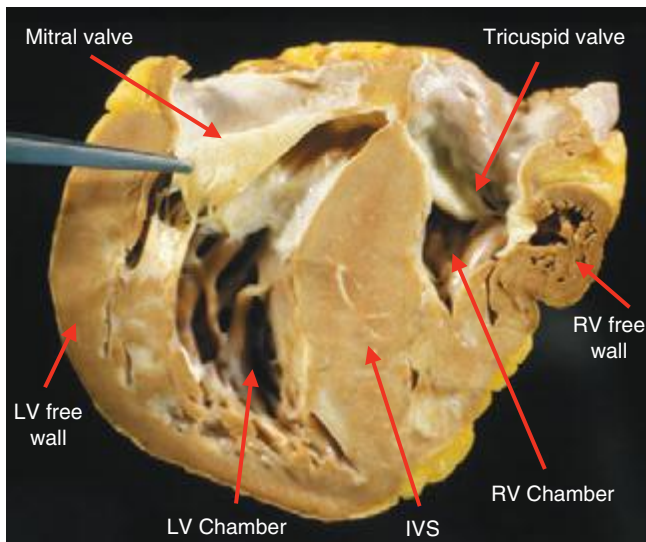


FIGURE 287-15 Hypertrophic cardiomyopathy. Gross specimen of a heart with hypertrophic cardiomyopathy removed at the time of transplantation, showing asymmetric septal hypertrophy (septum much thicker than left ventricular free wall) with the septum bulging into the left ventricular outflow tract causing obstruction. The forceps are retracting the anterior leaflet of the mitral valve, demonstrating the characteristic plaque of systolic anterior motion, manifest as endocardial fibrosis on the interventricular septum in a mirror-image pattern to the valve leaflet. There is patchy replacement fibrosis, and small thick-walled arterioles can be appreciated grossly, especially in the interventricular septum. IVS, interventricular septum; LV, left ventricle; RV, right ventricle. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

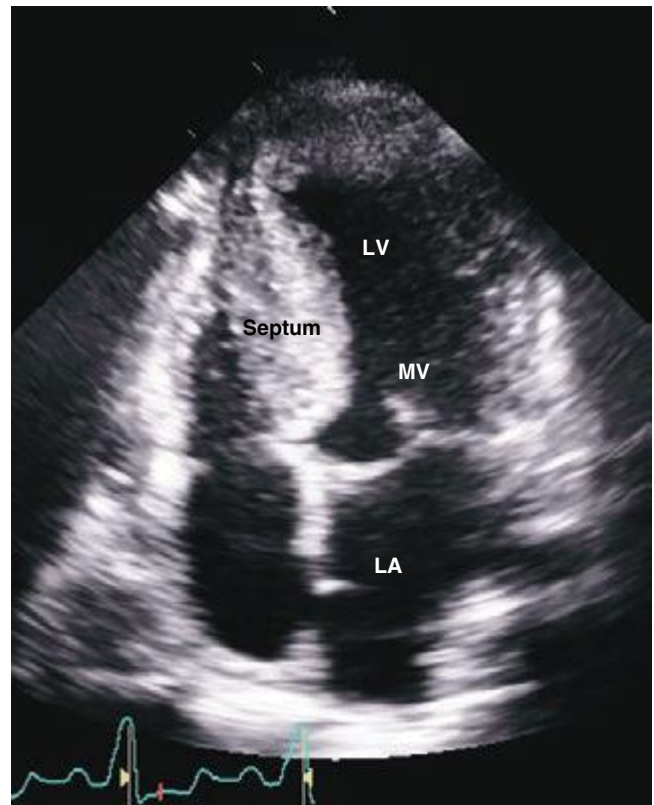


FIGURE 287-16 Hypertrophic cardiomyopathy. This echocardiogram of hypertrophic cardiomyopathy shows asymmetric hypertrophy of the septum compared to the lateral wall of the left ventricle (LV). The mitral valve (MV) is moving anteriorly toward the hypertrophied septum in systole. The left atrium (LA) is enlarged. Note that the echocardiographic and pathologic images are vertically opposite, such that the LV is by convention on the top right in the echocardiographic image and bottom right in the pathologic images. (Image courtesy of Justina Wu, MD, Brigham and Women's Hospital, Boston.)

cardiomyopathy with or without obstruction. Prevalence in North America, Japan, and China is about 1:500. It is the leading cause of sudden death in the young and is an important cause of heart failure. Although pediatric presentation is associated with increased early morbidity and mortality, the prognosis for patients diagnosed as adults is generally favorable.

The clustering of hypertrophic cardiomyopathy within families has been appreciated since recognition of the disease approximately 55 years ago. Echocardiographic screening of families revealed an autosomal dominant pattern of inheritance. Initial genetic studies using linkage analysis in large families identified disease-causing mutations in sarcomeric genes. A sarcomere mutation is present in ~60% of patients with hypertrophic cardiomyopathy and is more common in those with familial disease and characteristic asymmetric septal hypertrophy. More than nine different sarcomere genes with over 1400 mutations have been implicated, although ~80% of patients have a mutation in either *MYH7* or *MYBPC3* (Table 287-3), most of which are unique to individual families ("private" mutations).

Hypertrophic cardiomyopathy is characterized by age-dependent and incomplete penetrance. The defining phenotype of left ventricular hypertrophy is rarely present at birth and usually develops later in life. Accordingly, screening of family members should begin in adolescence and extend through adulthood. In *MYBPC3* mutation carriers, the average age of disease development is 40 years, while 30% remain free from hypertrophy after 70 years. Related individuals who carry the *same* mutation may have a different extent and pattern of hypertrophy (e.g., asymmetric versus concentric), occurrence of outflow tract obstruction, and associated clinical outcomes (e.g., sudden death, atrial fibrillation).

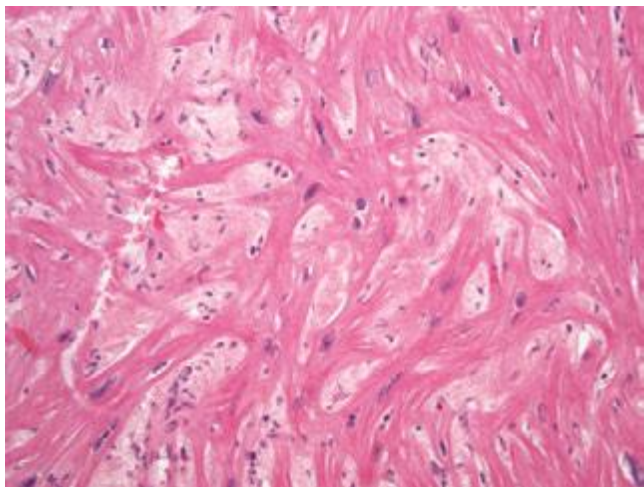


FIGURE 287-17 Hypertrophic cardiomyopathy. Microscopic image of hypertrophic cardiomyopathy showing the characteristic disordered myocyte architecture with swirling and branching rather than the usual parallel arrangement of myocyte fibers. Myocyte nuclei vary markedly in size and interstitial fibrosis is present. (Image courtesy of Robert Padera, MD, PhD, Department of Pathology, Brigham and Women's Hospital, Boston.)

At the level of the sarcomere, hypertrophic cardiomyopathy mutations lead to enhanced calcium sensitivity, maximal force generation, and ATPase activity. Calcium handling is affected through modification of regulatory proteins. Sarcomere mutations lead to abnormal energetics and impaired relaxation, both directly and as a result of hypertrophy. Hypertrophic cardiomyopathy is characterized by misalignment and disarray of the enlarged myofibrils and myocytes (Fig. 287-17), which can also occur to a lesser extent in other cardiac diseases. Although hypertrophy is the defining feature of hypertrophic cardiomyopathy, fibrosis and microvascular disease are also present. Interstitial fibrosis is detectable before overt hypertrophy develops and likely results from early activation of profibrotic pathways. In the majority of patients with overt cardiomyopathy, focal areas of replacement fibrosis can be readily detected with MRI. These areas of “scar” may represent substrate for the development of ventricular arrhythmias. Increased thickness and decreased luminal area of the intramural vessels in hypertrophied myocardium contribute to microvascular ischemia and angina. Microinfarction of hypertrophied myocardium is a hypothesized mechanism for replacement scar formation.

Macroscopically, hypertrophy is typically manifest as nonuniform ventricular thickening (Fig. 287-15). The interventricular septum is the typical location of maximal hypertrophy, although other patterns of hypertrophic remodeling include concentric and midventricular. Hypertrophy confined to the ventricular apex (apical hypertrophic cardiomyopathy) is less often familial and has a different genetic substrate, with sarcomere mutations present in only ~15%. Left ventricular outflow tract obstruction represents the most common focus of diagnosis and intervention, although diastolic dysfunction, myocardial fibrosis, and microvascular ischemia also contribute to contractile dysfunction and elevated intracardiac pressures. Obstruction is present in ~30% of patients at rest and can be provoked by exercise in another ~30%. Systolic obstruction is initiated by drag forces, which push an anteriorly displaced and enlarged anterior mitral leaflet into contact with the hypertrophied ventricular septum. Mitral leaflet coaptation may ensue, leading to posteriorly directed mitral regurgitation. In order to maintain stroke volume across outflow tract obstruction, the ventricle generates higher pressures, leading to higher wall stress and myocardial oxygen demand. Smaller chamber size and increased contractility exacerbate the severity of obstruction. Conditions of low preload, such as dehydration, and low afterload, such as arterial vasodilation, may lead to transient hypotension and near-syncope. The systolic ejection murmur of left ventricular outflow tract obstruction is harsh and late peaking and can be enhanced by bedside maneuvers that diminish

ventricular volume and transiently worsen obstruction, such as standing from a squatting position or the Valsalva maneuver.

DIAGNOSIS

The substantial variability of hypertrophic cardiomyopathy pathology is reflected in the diversity of clinical presentations. Patients may be diagnosed after undergoing evaluations triggered by the abnormal physical findings (murmur) or symptoms of exertional dyspnea, angina, or syncope. Alternatively, diagnosis may follow evaluations prompted by the detection of disease in family members. Cardiac imaging (Fig. 287-16) is central to diagnosis due to the insensitivity of examination and ECG and the need to exclude other causes for hypertrophy. The identification of a disease-causing mutation in a proband can focus family evaluations on mutation carriers, but this strategy requires a high degree of certainty that the mutation is truly pathogenic and not a benign DNA variant. Biopsy is not needed to diagnose hypertrophic cardiomyopathy but can be used to exclude infiltrative and metabolic diseases. Rigorous athletic training (athlete's heart) may cause intermediate degrees of physiologic hypertrophy difficult to differentiate from mild hypertrophic cardiomyopathy. Unlike hypertrophic cardiomyopathy, hypertrophy in the athlete's heart regresses with cessation of training, and is accompanied by supernormal exercise capacity ($VO_{2max} > 50$ mL/kg/min), mild ventricular dilation, and normal diastolic function.

TREATMENT

HYPERTROPHIC CARDIOMYOPATHY

Management focuses on treatment of symptoms and prevention of sudden death and stroke (Fig. 287-18). Left ventricular outflow tract obstruction can be controlled medically in the majority of patients. β -Adrenergic blocking agents and L-type calcium channel blockers (e.g., verapamil) are first-line agents that reduce the severity of obstruction by slowing heart rate, enhancing diastolic filling, and decreasing contractility. Persistent symptoms of exertional dyspnea or chest pain can sometimes be controlled with the addition of disopyramide, an antiarrhythmic agent with potent negative inotropic properties. Patients with or without obstruction may develop heart failure symptoms due to fluid retention and require diuretic therapies for venous congestion. Severe medically refractory symptoms develop in ~5% of patients, for whom surgical myectomy or alcohol septal ablation may be effective. Developed over 50 years ago, surgical myectomy effectively relieves outflow tract obstruction by excising part of the septal myocardium involved in the dynamic obstruction. In selected patients, perioperative mortality is extremely low with excellent long-term survival free from recurrent obstruction and symptoms. Mitral valve repair or replacement is usually unnecessary as associated eccentric mitral regurgitation resolves with myectomy alone. Alcohol septal ablation in patients with suitable coronary anatomy can relieve outflow tract obstruction via a controlled infarction of the proximal septum, which produces similar periprocedural outcomes and gradient reduction as surgical myectomy. Until long-term outcomes are demonstrated for this procedure, it is relegated primarily to patients who wish to avoid surgery or who have limiting comorbidities. Neither procedure has been shown to improve outcomes other than symptoms. With both procedures, the most common complication is the development of complete heart block necessitating permanent pacing. However, ventricular pacing as a primary therapy for outflow tract obstruction is ineffective and not generally advised.

Patients with hypertrophic cardiomyopathy have an increased risk of sudden cardiac death from ventricular tachyarrhythmias. Vigorous physical activity and competitive sport are prohibited. Factors that increase the risk of sudden death from a baseline of 0.5% per year are presented in Table 287-6. As sudden death has not been reduced by medical or procedural interventions, an implantable cardioverter-defibrillator is advised for patients with two or more risk factors and is advised on a selected basis for patient with one risk factor. Nevertheless, the positive predictive value of most risk factors is low, and many patients receiving a defibrillator

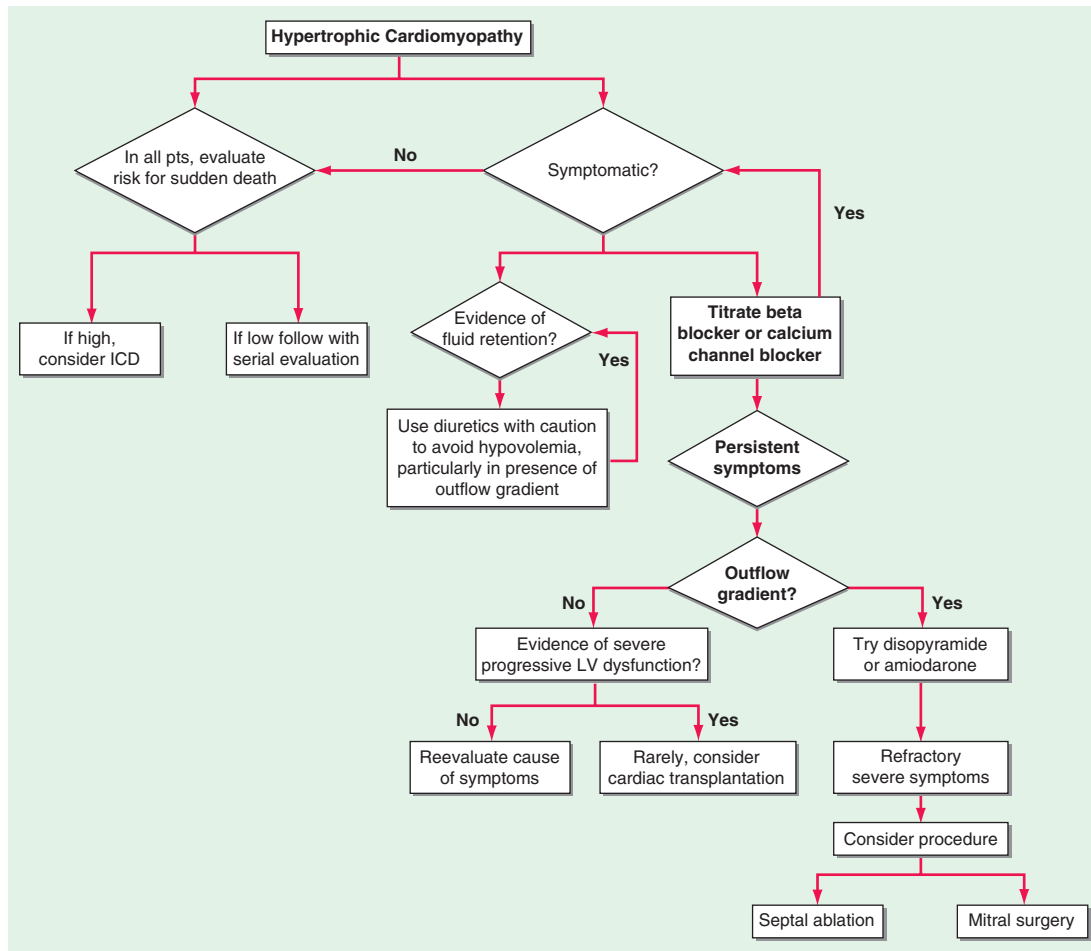


FIGURE 287-18 Treatment algorithm for hypertrophic cardiomyopathy depending on the presence and severity of symptoms and the presence of an intraventricular gradient with obstruction to outflow. Note that all patients with hypertrophic cardiomyopathy should be evaluated for atrial fibrillation and risk of sudden death, whether or not they require treatment for symptoms. ICD, implantable cardioverter-defibrillator; LV, left ventricular.

never receive an appropriate therapy. Long-term use of a defibrillator may be associated with serious device-related complications, particularly in young active patients. Refinement of sudden death risk through the application of contemporary technologies such as cardiac MRI is ongoing.

TABLE 287-6 RISK FACTORS FOR SUDDEN DEATH IN HYPERTROPHIC CARDIOMYOPATHY		
Major Risk Factor		Screening Technique
History of cardiac arrest or spontaneous sustained ventricular tachycardia ^a		History
Syncope	Nonvagal, often with or after exertion	History
Family history of sudden cardiac death		Family history
Spontaneous nonsustained ventricular tachycardia ^b	>3 beats at rate >120	Exercise or 24- to 48-hour ambulatory recording
LV thickness >30 mm	Present in <10% of patients	Echocardiography
Abnormal blood pressure response to exercise ^b	Systolic blood pressure fall or failure to increase at peak exercise	Maximal upright exercise testing

^aImplantable cardioverter-defibrillator advised for patients with prior arrest or sustained ventricular tachycardia regardless of other risk factors. ^bPrognostic value most applicable to patients less than 40 years old.

Abbreviation: LV, left ventricle.

Atrial fibrillation is common in patients with hypertrophic cardiomyopathy and may lead to hemodynamic deterioration and embolic stroke. Rapid ventricular response is poorly tolerated and may worsen outflow tract obstruction. β -Adrenergic blocking agents and L-type calcium channel blockers slow AV nodal conduction and improve symptoms; cardiac glycosides should be avoided, as they may increase contractility and worsen obstruction. Symptoms exacerbated by atrial fibrillation may persist despite adequate rate control due to loss of AV synchrony and may require restoration of sinus rhythm. Disopyramide and amiodarone are the preferred antiarrhythmic agents, with radiofrequency ablation considered for medically refractory cases. Anticoagulation to prevent embolic stroke in atrial fibrillation is recommended.

PROGNOSIS The general prognosis for hypertrophic cardiomyopathy is good, better than in early studies of referral populations. For patients diagnosed as adults, survival is comparable to an age-matched population without cardiomyopathy. The sudden death risk is less than 1% per year; however, up to 1 in 20 patients will progress to overt systolic dysfunction with a reduced ejection fraction with or without dilated remodeling (“burned out” or end-stage hypertrophic cardiomyopathy). These patients suffer from low cardiac output and have a high risk of death from progressive heart failure and sudden death unless they undergo cardiac transplantation.

NORMAL FUNCTIONS OF THE PERICARDIUM

The normal pericardium is a double-layered sac; the visceral pericardium is a serous membrane that is separated by a small quantity (15–50 mL) of fluid, an ultrafiltrate of plasma, from the fibrous parietal pericardium. The normal pericardium, by exerting a restraining force, prevents sudden dilation of the cardiac chambers, especially the right atrium and ventricle, during exercise and with hypervolemia. It also restricts the anatomic position of the heart, and probably retards the spread of infections from the lungs and pleural cavities to the heart. Nevertheless, *total* absence of the pericardium, either congenital or after surgery, does not produce obvious clinical disease. In *partial* left pericardial defects, the main pulmonary artery and left atrium may bulge through the defect; very rarely, herniation and subsequent strangulation of the left atrium may cause sudden death.

ACUTE PERICARDITIS

Acute pericarditis, by far the most common pathologic process involving the pericardium (Table 288-1), has four principal diagnostic features:

1. *Chest pain* is usually present in acute infectious pericarditis and in many of the forms presumed to be related to hypersensitivity or autoimmunity. The pain of acute pericarditis is often severe, retrosternal, and left precordial, and referred to the neck, arms, or left shoulder. Frequently the pain is pleuritic, consequent to accompanying pleural inflammation (i.e., sharp and aggravated by inspiration and coughing), but sometimes it is steady, constricting, radiates into either arm or both arms, and resembles that of myocardial ischemia; therefore, confusion with acute myocardial infarction (AMI) is common. Characteristically, however, pericardial pain may be relieved by sitting up and leaning forward and is intensified by lying supine (Chap. 19). Pain is often absent in slowly developing tuberculous, postirradiation, and neoplastic, uremic, and constrictive pericarditis.

The differentiation of AMI from acute pericarditis may become perplexing when, with acute pericarditis, serum biomarkers of myocardial damage such as troponin and creatine kinase-MB rise, presumably because of concomitant involvement of the epicardium in the inflammatory process (an epi-myocarditis) with resulting myocyte necrosis. However, these elevations, if they occur, are quite modest given the extensive electrocardiographic ST-segment elevation in pericarditis. This dissociation is useful in differentiating between these conditions.
2. A *pericardial friction rub* is audible at some point in about 85% of patients with acute pericarditis, may have up to three components per cardiac cycle, is high-pitched, and is described as rasping, scratching, or grating (Chap. 267). It is heard most frequently at end expiration with the patient upright and leaning forward.
3. The *electrocardiogram* (ECG) in acute pericarditis without massive effusion usually displays changes secondary to acute subepicardial inflammation (Fig. 288-1). It typically evolves through four stages. In stage 1, there is widespread elevation of the ST segments, often with upward concavity, involving two or three standard limb leads and V_2 to V_6 , with reciprocal depressions only in aVR and sometimes V_1 . Also, there is depression of the PR segment below the TP segment, reflecting atrial involvement. Usually there are no significant changes in QRS complexes. After several days, the ST segments return to normal (stage 2), and only then, or even later, do the T waves become inverted (stage 3). Weeks or months after the onset of acute pericarditis, the ECG returns to normal (stage 4). In contrast, in AMI, ST elevations are convex, and reciprocal depression is usually more prominent; these changes may return to normal within a

TABLE 288-1 CLASSIFICATION OF PERICARDITIS

Clinical Classification	
I.	Acute pericarditis (<6 weeks) <ol style="list-style-type: none"> A. Fibrinous B. Effusive (serous or sanguineous)
II.	Subacute pericarditis (6 weeks to 6 months) <ol style="list-style-type: none"> A. Effusive-constrictive B. Constrictive
III.	Chronic pericarditis (>6 months) <ol style="list-style-type: none"> A. Constrictive B. Effusive C. Adhesive (nonconstrictive)
Etiologic Classification	
I.	Infectious pericarditis <ol style="list-style-type: none"> A. Viral (coxsackievirus A and B, echovirus, mumps, adenovirus, hepatitis, HIV) B. Pyogenic (pneumococcus, <i>Streptococcus</i>, <i>Staphylococcus</i>, <i>Neisseria</i>, <i>Legionella</i>) C. Tuberculous D. Fungal (histoplasmosis, coccidioidomycosis, <i>Candida</i>, blastomycosis) E. Other infections (syphilitic, protozoal, parasitic)
II.	Noninfectious pericarditis <ol style="list-style-type: none"> A. Acute myocardial infarction B. Uremia C. Neoplasia <ol style="list-style-type: none"> 1. Primary tumors (benign or malignant, mesothelioma) 2. Tumors metastatic to pericardium (lung and breast cancer, lymphoma, leukemia) D. Myxedema E. Cholesterol F. Chylopericardium G. Trauma <ol style="list-style-type: none"> 1. Penetrating chest wall 2. Nonpenetrating H. Aortic dissection (with leakage into pericardial sac) I. Postirradiation J. Familial Mediterranean fever K. Familial pericarditis <ol style="list-style-type: none"> 1. Mulibrey nanism^a L. Acute idiopathic M. Whipple's disease N. Sarcoidosis
III.	Pericarditis presumably related to hypersensitivity or autoimmunity <ol style="list-style-type: none"> A. Rheumatic fever B. Collagen vascular disease (systemic lupus erythematosus, rheumatoid arthritis, ankylosing spondylitis, scleroderma, acute rheumatic fever, granulomatosis with polyangiitis [Wegener's]) C. Drug-induced (e.g., procainamide, hydralazine, phenytoin, isoniazid, minoxidil, anticoagulants, methysergide) D. Postcardiac injury <ol style="list-style-type: none"> 1. Postmyocardial infarction (Dressler's syndrome) 2. Postpericardiotomy 3. Posttraumatic

^aAn autosomal recessive syndrome characterized by growth failure, muscle hypotonia, hepatomegaly, ocular changes, enlarged cerebral ventricles, mental retardation, ventricular hypertrophy, and chronic constrictive pericarditis

day or two. Q waves may develop, with loss of R-wave amplitude, and T-wave inversions are usually seen within hours *before* the ST segments have become isoelectric (Chaps. 294 and 295).

4. *Pericardial effusion* is usually associated with pain and/or the ECG changes mentioned above, as well as electrical alternans.

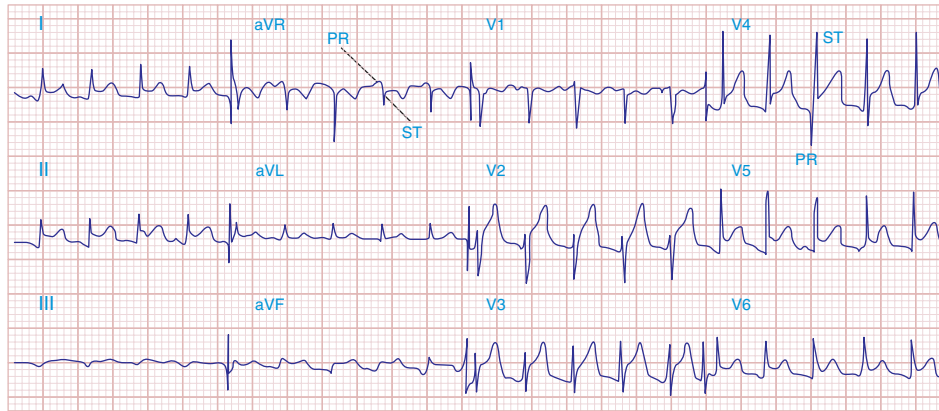


FIGURE 288-1 Acute pericarditis. There are diffuse ST-segment elevations (in this case in leads I, II, aVF, and V_2 to V_6) due to a ventricular current of injury. There is PR-segment deviation (opposite in polarity to the ST segment) due to a concomitant atrial injury current.

Pericardial effusion is especially important clinically when it develops within a relatively short time because it may lead to cardiac tamponade (see below). Differentiation from cardiac enlargement may be difficult on physical examination, but heart sounds may be fainter with pericardial effusion. The friction rub and the apex impulse may disappear. The base of the left lung may be compressed by pericardial fluid, producing *Ewart's sign*, a patch of dullness and increased fremitus (and egophony) beneath the angle of the left scapula. The chest roentgenogram may show enlargement of the cardiac silhouette, with a “water bottle” configuration, but may be normal.

Diagnosis Echocardiography (Chap. 270e) is the most widely used imaging technique. It is sensitive, specific, simple, noninvasive, may be performed at the bedside, and can identify accompanying cardiac tamponade (see below) (Fig. 288-2). The presence of pericardial fluid is recorded by two-dimensional transthoracic echocardiography as a relatively echo-free space between the posterior pericardium and left ventricular epicardium in patients with small effusions and as a space between the anterior right ventricle and the parietal pericardium just beneath the anterior chest wall. In patients with large effusions, the heart may swing freely within the pericardial sac. When severe, the extent of this motion alternates and may be associated with electrical alternans (Fig. 288-3). Echocardiography allows localization and identification of the quantity of pericardial fluid.

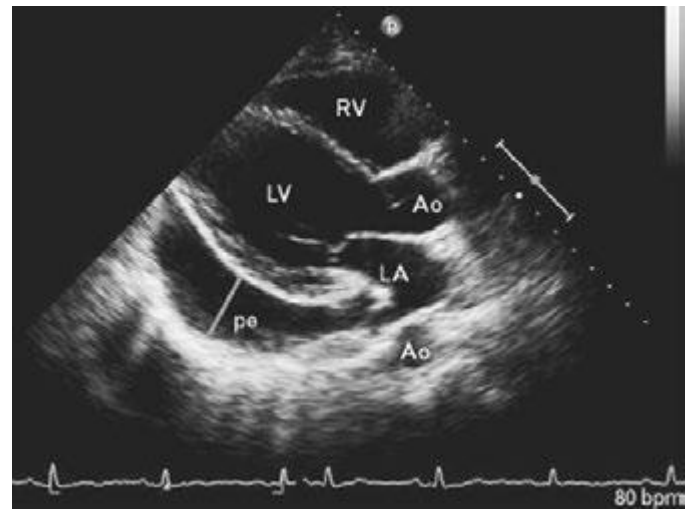


FIGURE 288-2 Two-dimensional echocardiogram in lateral view in a patient with a large pericardial effusion. Ao, aorta; LV, left ventricle; pe, pericardial effusion; RV, right ventricle. (From M Imazio: *Curr Opin Cardiol* 27:308, 2012.)

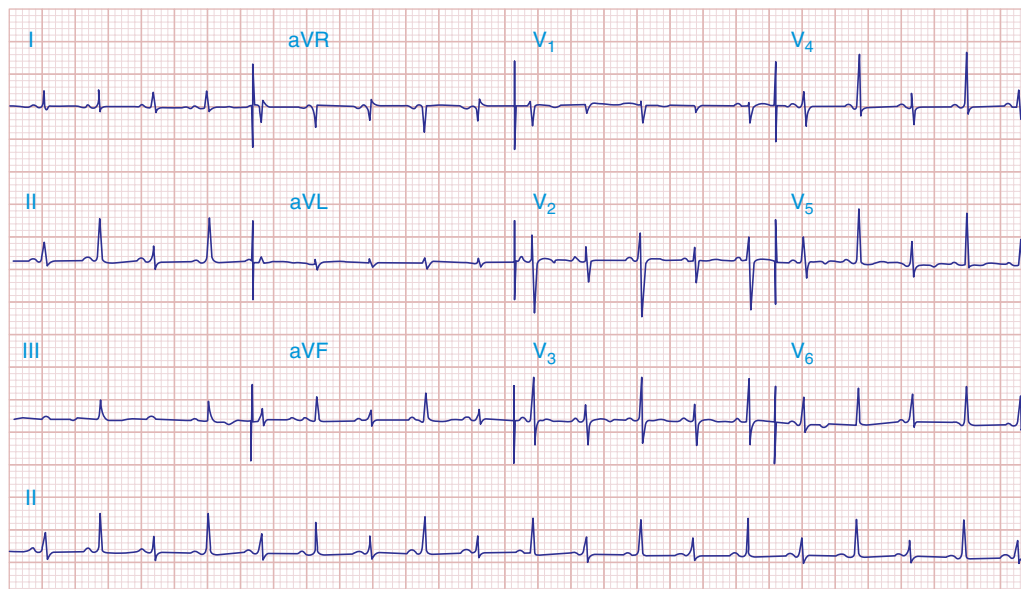


FIGURE 288-3 Electrical alternans. This tracing was obtained from a patient with a large pericardial effusion with tamponade. (Reproduced from DM Mirvis, AL Goldberger: *Electrocardiography*, in RO Bonow et al [eds]: *Braunwald's Heart Disease*, 9th ed. Philadelphia: Elsevier, 2012.)

The diagnosis of pericardial fluid or thickening may be confirmed by computed tomography (CT) or magnetic resonance imaging (MRI). These techniques may be superior to echocardiography in detecting loculated pericardial effusions, pericardial thickening, and the identification of pericardial masses.

TREATMENT ACUTE PERICARDITIS

There is no specific therapy for acute idiopathic pericarditis, but bed rest and anti-inflammatory treatment with aspirin (2–4 g/d), with gastric protection (e.g., omeprazole 20 mg/d), may be given. If this is ineffective, one of the nonsteroidal anti-inflammatory drugs (NSAIDs), such as ibuprofen (400–600 mg tid) or indomethacin (25–50 mg tid), should be tried. In responsive patients, these doses should be continued for 1–2 weeks and then tapered over several weeks. In patients who are unresponsive, colchicine (0.5 mg bid, given for 4–8 weeks) has been found to be effective, not only in acute pericarditis, but also in reducing the risk of recurrent pericarditis. Colchicine is concentrated in and interferes with the migration of neutrophils, is contraindicated in patients with hepatic or renal dysfunction, and may cause diarrhea and other gastrointestinal side effects. Glucocorticoids (e.g., prednisone 1 mg/kg per day) usually suppress the clinical manifestations of acute pericarditis in patients who have failed therapy with the anti-inflammatory therapies described above, but appear to increase the risk of subsequent recurrence. Therefore, full-dose corticosteroids should be given for only 2–4 days and then tapered. Anticoagulants should be avoided because their use could cause bleeding into the pericardial cavity and tamponade.

In patients with recurrences that are multiple, frequent, disabling, continue for more than 2 years, and are not prevented by colchicine and other NSAIDs and are not controlled by glucocorticoids,

pericardial stripping may be necessary to terminate the illness, and usually does so.

CARDIAC TAMPONADE

The accumulation of fluid in the pericardial space in a quantity sufficient to cause serious obstruction of the inflow of blood into the ventricles results in cardiac tamponade. This complication may be fatal if it is not recognized and treated promptly. The most common causes of tamponade are idiopathic pericarditis and pericarditis secondary to neoplastic disease. Tamponade may also result from bleeding into the pericardial space after leakage from an aortic dissection, cardiac operations, trauma, and treatment of patients with acute pericarditis with anticoagulants.

The three principal features of tamponade (*Beck's triad*) are hypotension, soft or absent heart sounds, and jugular venous distention with a prominent *x* descent but an absent *y* descent. The limitations of ventricular filling are responsible for a reduction of cardiac output. The quantity of fluid necessary to produce cardiac tamponade may be as small as 200 mL when the fluid develops rapidly to as much as >2000 mL in slowly developing effusions when the pericardium has had the opportunity to stretch and adapt to an increasing volume. Tamponade may also develop more slowly, and in these circumstances, the clinical manifestations can resemble those of heart failure, including dyspnea, orthopnea, and hepatic engorgement.

A high index of suspicion for cardiac tamponade is required because in many instances no obvious cause for pericardial disease is apparent, and this diagnosis should be considered in any patient with otherwise unexplained enlargement of the cardiac silhouette, hypotension, and elevation of jugular venous pressure. There may be reduction in amplitude of the QRS complexes, and *electrical alternans* of the P, QRS, or T waves should raise the suspicion of cardiac tamponade (Fig. 288-3).

Table 288-2 lists the features that distinguish acute cardiac tamponade from constrictive pericarditis.

TABLE 288-2 FEATURES THAT DISTINGUISH CARDIAC TAMPONADE FROM CONSTRICTIVE PERICARDITIS AND SIMILAR CLINICAL DISORDERS

Characteristic	Tamponade	Constrictive Pericarditis	Restrictive Cardiomyopathy	RVMI	Effusive Constrictive Pericarditis
Clinical					
Pulsus paradoxus	+++	+	+	+	+++
Jugular veins					
Prominent <i>y</i> descent	–	++	+	+	–
Prominent <i>x</i> descent	+++	++	+++	+	+++
Kussmaul's sign	–	+++	+	+++	++
Third heart sound	–	–	+	+	+
Pericardial knock	–	++	–	–	–
Electrocardiogram					
Low ECG voltage	++	++	++	–	++
Electrical alternans	++	–	–	–	+
Echocardiogram					
Thickened pericardium	–	+++	–	–	++
Pericardial calcification	–	++	–	–	–
Pericardial effusion	+++	–	–	++	–
RV size	Usually small	Usually normal	Usually normal	Enlarged	–
RA and RV	+++	–	–	–	–
Exaggerated respiratory variation in flow velocity	+++	+++	–	+++	–
CT/MRI					
Thickened pericardium	–	+++	–	++	–
Cardiac catheterization					
Equalization of diastolic pressures	+++	+++	–	++	–

Abbreviations: +++, always present; ++, usually present; +, rare; –, absent; DC, diastolic collapse; ECG, electrocardiograph; RA, right atrium; RV, right ventricle; RVMI, right ventricular myocardial infarction.

Source: Adapted from GM Brockington et al: *Cardiol Clin* 8:645, 1990, with permission.

1574 Paradoxical Pulse This important clue to the presence of cardiac tamponade consists of a greater than normal (10 mmHg) inspiratory decline in systolic arterial pressure. When severe, it may be detected by palpating weakness or disappearance of the arterial pulse during inspiration, but usually sphygmomanometric measurement of systolic pressure during slow respiration is required.

Because both ventricles share a tight incompressible covering, i.e., the pericardial sac, the inspiratory enlargement of the right ventricle in cardiac tamponade compresses and reduces left ventricular volume; leftward bulging of the interventricular septum reduces further the left ventricular cavity as the right ventricle enlarges during inspiration. Thus, in cardiac tamponade, the normal inspiratory augmentation of right ventricular volume causes an exaggerated reduction of left ventricular volume, stroke volume, and systolic pressure. Paradoxical pulse also occurs in approximately one-third of patients with constrictive pericarditis (see below), and in some cases of hypovolemic shock, acute and chronic obstructive airway disease, and pulmonary embolus. Right ventricular infarction (**Chap. 295**) may resemble cardiac tamponade with hypotension, elevated jugular venous pressure, an absent y descent in the jugular venous pulse, and, occasionally, a paradoxical pulse (Table 288-2).

Low-pressure tamponade refers to mild tamponade in which the intrapericardial pressure is increased from its slightly subatmospheric levels to +5 to +10 mmHg; in some instances, hypovolemia coexists. As a consequence, the central venous pressure is normal or only slightly elevated, whereas arterial pressure is unaffected and there is no paradoxical pulse. These patients are asymptomatic or complain of mild weakness and dyspnea. The diagnosis is aided by echocardiography, and both hemodynamic and clinical manifestations improve after pericardiocentesis.

Diagnosis Because immediate treatment of cardiac tamponade may be lifesaving, prompt measures to establish the diagnosis by echocardiography should be undertaken. When pericardial effusion causes tamponade, Doppler ultrasound shows that tricuspid and pulmonic valve flow velocities increase markedly during inspiration, whereas pulmonic vein, mitral, and aortic flow velocities diminish (as in constrictive pericarditis, see below) (**Fig. 288-4**). In tamponade, there is late diastolic inward motion (collapse) of the right ventricular free wall and the right atrium. Transesophageal echocardiography, CT, or cardiac MRI may be necessary to diagnose a loculated effusion responsible for cardiac tamponade.

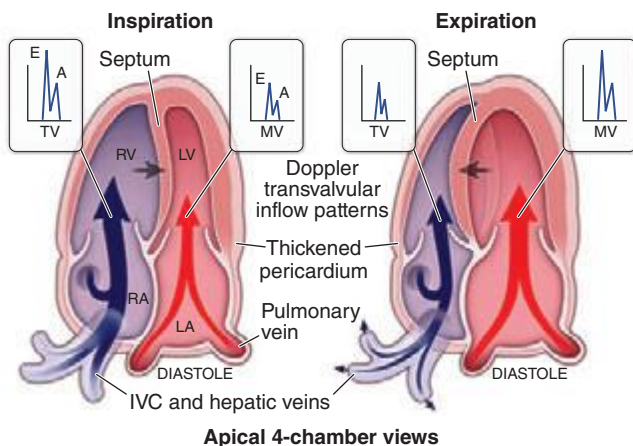


FIGURE 288-4 Constrictive pericarditis. Doppler schema of respirophasic changes in mitral and tricuspid inflow. Reciprocal patterns of ventricular filling are assessed on pulsed Doppler examination of mitral valve (MV) and tricuspid valve (TV) inflow. IVC, inferior vena cava; LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle. (Courtesy of Bernard E. Bulwer, MD; with permission.)

TREATMENT CARDIAC TAMPONADE

Patients with acute pericarditis should be observed frequently for the development of an effusion; if a large effusion is present, pericardiocentesis should be carried out or the patient watched closely for signs of tamponade. Arterial and venous pressures should be monitored and serial echocardiograms obtained.

PERICARDIOCENTESIS

If manifestations of tamponade appear, echocardiographically guided pericardiocentesis using an apical, parasternal, or, most commonly, subxiphoid approach must be carried out at once because reduction of the elevated intrapericardial pressure may be lifesaving. Intravenous saline may be administered as the patient is being readied for the procedure, but the pericardiocentesis must not be delayed. If possible, intrapericardial pressure should be measured before fluid is withdrawn, and the pericardial cavity should be drained as completely as possible. A small, multiholed catheter advanced over the needle inserted into the pericardial cavity may be left in place to allow draining of the pericardial space if fluid reaccumulates. Surgical drainage through a limited (subxiphoid) thoracotomy may be required in recurrent tamponade, when it is necessary to remove loculated effusions, and/or when it is necessary to obtain tissue for diagnosis.

Pericardial fluid obtained from an effusion often has the physical characteristics of an exudate. Bloody fluid is most commonly due to neoplasm, renal failure, or dialysis in the United States and tuberculosis in developing nations but may also be found in the effusion of acute rheumatic fever, after cardiac injury, and after myocardial infarction. Transudative pericardial effusions may occur in heart failure.

The pericardial fluid should be analyzed for red and white blood cells and cytologic studies, and cultures should be obtained. The presence of DNA of *Mycobacterium tuberculosis* determined by the polymerase chain reaction strongly supports the diagnosis of tuberculous pericarditis (**Chap. 202**).

VIRAL OR IDIOPATHIC ACUTE PERICARDITIS

In many instances, acute pericarditis occurs in association with illnesses of known or presumed viral origin and probably is caused by the same agent. Commonly, there is an antecedent infection of the respiratory tract, and viral isolation and serologic studies are negative. In some cases, coxsackievirus A or B or the virus of influenza, echovirus, mumps, herpes simplex, chickenpox, adenovirus, or cytomegalovirus has been isolated from pericardial fluid and/or appropriate elevations in viral antibody titers have been noted. Pericardial effusion is a common cardiac manifestation of HIV; it is usually secondary to infection (often mycobacterial) or neoplasm, most often lymphoma. Frequently, a viral cause cannot be established, and the term *idiopathic acute pericarditis* is then appropriate.

Viral or idiopathic acute pericarditis occurs at all ages but is more common in young adults and is often associated with pleural effusions and pneumonitis. The almost simultaneous development of fever and precordial pain, often 10–12 days after a presumed viral illness, constitutes an important feature in the differentiation of acute pericarditis from AMI, in which chest pain precedes fever. The constitutional symptoms are usually mild to moderate, and a pericardial friction rub is often audible. The disease ordinarily runs its course in a few days to 4 weeks. The ST-segment alterations in the ECG usually disappear after 1 or more weeks, but the abnormal T waves may persist for several years and be a source of confusion in persons without a clear history of pericarditis. Pleuritis and pneumonitis frequently accompany viral or idiopathic acute pericarditis. Accumulation of some pericardial fluid is common, and both tamponade and constrictive pericarditis are possible, but infrequent, complications.

The most frequent complication is recurrent (relapsing) pericarditis, which occurs in about one-fourth of patients with acute idiopathic

pericarditis. In a smaller number, there are multiple recurrences. For treatment, see earlier section on treatment of acute pericarditis.

Postcardiac Injury Syndrome Acute pericarditis may appear in a variety of circumstances that have one common feature—previous injury to the myocardium with blood in the pericardial cavity. The syndrome may develop after a cardiac operation (postpericardiotomy syndrome), after blunt or penetrating cardiac trauma (**Chap. 289e**), or after perforation of the heart with a catheter. Rarely, it follows AMI.

The clinical picture mimics acute viral or idiopathic pericarditis. The principal symptom is the pain of acute pericarditis, which usually develops 1–4 weeks after the cardiac injury but earlier (1–3 days) after AMI. Recurrences are common and may occur up to 2 years or more following the injury. Fever, pleuritis, and pneumonitis are the outstanding features, and the bout of illness usually subsides in 1 or 2 weeks. The pericarditis may be of the fibrinous variety, or it may be a pericardial effusion, which is often serosanguineous but rarely causes tamponade. ECG changes typical of acute pericarditis may also occur. This syndrome is probably the result of a hypersensitivity reaction to antigen(s) that originate from injured myocardial tissue and/or pericardium.

Often no treatment is necessary aside from aspirin and analgesics. When the illness is severe or followed by a series of disabling recurrences, therapy with an NSAID, colchicine, or a glucocorticoid, such as described for treatment of acute pericarditis, is usually effective.

DIFFERENTIAL DIAGNOSIS

Because there is no specific test for *acute idiopathic pericarditis*, the diagnosis is one of exclusion. Consequently, all other disorders that may be associated with acute fibrinous pericarditis must be considered. A common diagnostic error is mistaking acute viral or idiopathic pericarditis for AMI and vice versa. When acute fibrinous pericarditis is associated with AMI (**Chap. 295**), it is characterized by fever, pain, and a friction rub in the first 4 days after the development of the infarct. ECG abnormalities (such as the appearance of Q waves, brief ST-segment elevations with reciprocal changes, and earlier T-wave changes in AMI) and the extent of the elevations of markers of myocardial necrosis (higher in AMI) are helpful in differentiating pericarditis from AMI.

Pericarditis secondary to postcardiac injury is differentiated from acute idiopathic pericarditis chiefly by timing. If it occurs within a few days or weeks of an AMI, a chest blow, a cardiac perforation, or a cardiac operation, it may be justified to conclude that the two are probably related.

It is important to distinguish *pericarditis due to collagen vascular disease* from acute idiopathic pericarditis. Most important in the differential diagnosis is the pericarditis due to systemic lupus erythematosus (SLE; **Chap. 378**) or drug-induced (procainamide or hydralazine) lupus. When pericarditis occurs in the absence of any obvious underlying disorder, the diagnosis of SLE may be suggested by a rise in the titer of antinuclear antibodies. Acute pericarditis is an occasional complication of *rheumatoid arthritis*, *scleroderma*, and *polyarteritis nodosa*, and other evidence of these diseases is usually obvious.

Pyogenic (purulent) pericarditis is usually secondary to cardiothoracic operations, by extension of infection from the lungs or pleural cavities, from rupture of the esophagus into the pericardial sac, or from rupture of a ring abscess in a patient with infective endocarditis. It may also complicate the viral, pyogenic, mycobacterial, and fungal infections that occur with HIV infection. It is generally accompanied by fever, chills, septicemia, and evidence of infection elsewhere and generally has a poor prognosis. The diagnosis is made by examination of the pericardial fluid. It requires drainage as well as vigorous antibiotic treatment.

Pericarditis of renal failure occurs in up to one-third of patients with chronic uremia (*uremic pericarditis*), and is also seen in patients undergoing chronic dialysis who have normal levels of blood urea and creatinine (*dialysis-associated pericarditis*). These two forms of pericarditis may be fibrinous and are generally associated with serosanguineous

effusions. A pericardial friction rub is common, but pain is usually absent or mild. Treatment with an NSAID and intensification of dialysis are usually adequate. Occasionally, tamponade occurs and pericardiocentesis is required. When the pericarditis of renal failure is recurrent or persistent, a pericardial window should be created or pericardiectomy may be necessary.

Pericarditis due to *neoplastic diseases* results from extension or invasion of metastatic tumors (most commonly carcinoma of the lung and breast, malignant melanoma, lymphoma, and leukemia) to the pericardium; pain, atrial arrhythmias, and tamponade are complications that occur occasionally. Diagnosis is made by pericardial fluid cytology or pericardial biopsy. *Mediastinal irradiation* for neoplasm may cause acute pericarditis and/or chronic constrictive pericarditis. Unusual causes of acute pericarditis include syphilis, fungal infection (histoplasmosis, blastomycosis, aspergillosis, and candidiasis), and parasitic infestation (amebiasis, toxoplasmosis, echinococcosis, and trichinosis).

CHRONIC PERICARDIAL EFFUSIONS

Chronic pericardial effusions are sometimes encountered in patients without an antecedent history of acute pericarditis. They may cause few symptoms per se, and their presence may be detected by finding an enlarged cardiac silhouette on a chest roentgenogram. Tuberculosis is a common cause. *Myxedema* may be responsible for chronic pericardial effusion that is sometimes massive but rarely, if ever, causes cardiac tamponade. The cardiac silhouette may be markedly enlarged, and an echocardiogram distinguishes cardiomegaly from pericardial effusion. The diagnosis of myxedema can be confirmed by tests of thyroid function (**Chap. 405**). Myxedematous pericardial effusion responds to thyroid hormone replacement. Neoplasms, SLE, rheumatoid arthritis, mycotic infections, radiation therapy to the chest, pyogenic infections, and chylopericardium may also cause chronic pericardial effusion and should be considered and specifically sought in such patients.

Aspiration and analysis of the pericardial fluid are often helpful in diagnosis. Pericardial fluid should be analyzed as described in pericardiocentesis. Grossly sanguineous pericardial fluid results most commonly from a neoplasm, tuberculosis, renal failure, or slow leakage from an aortic dissection. Pericardiocentesis may resolve large effusions, but pericardiectomy may be required in patients with recurrence. Intrapericardial instillation of sclerosing agents may be used to prevent reaccumulation of fluid.

CHRONIC CONSTRICTIVE PERICARDITIS

This disorder results when the healing of an acute fibrinous or serofibrinous pericarditis or the resorption of a chronic pericardial effusion is followed by obliteration of the pericardial cavity with the formation of granulation tissue. The latter gradually contracts and forms a firm scar encasing the heart, which may be calcified. In developing nations where the condition is prevalent, a high percentage of cases are of tuberculous origin, but this is now an uncommon cause in North America. Chronic constrictive pericarditis may follow acute or relapsing viral or idiopathic pericarditis, trauma with organized blood clot, or cardiac surgery of any type or result from mediastinal irradiation, purulent infection, histoplasmosis, neoplastic disease (especially breast cancer, lung cancer, and lymphoma), rheumatoid arthritis, SLE, or chronic renal failure treated by chronic dialysis. In many patients, the cause of the pericardial disease is undetermined, and in these patients, an asymptomatic or forgotten bout of viral pericarditis, acute or idiopathic, may have been the inciting event.

The basic physiologic abnormality in patients with chronic constrictive pericarditis is the inability of the ventricles to fill because of the limitations imposed by the rigid, thickened pericardium. Ventricular filling is unimpeded during early diastole but is reduced abruptly when the elastic limit of the pericardium is reached, whereas in cardiac tamponade, ventricular filling is impeded throughout diastole. In both conditions, ventricular end-diastolic and stroke volumes are reduced and the end-diastolic pressures in both ventricles and

1576 the mean pressures in the atria, pulmonary veins, and systemic veins are all elevated to similar levels (i.e., within 5 mmHg of one another). Despite these hemodynamic changes, systolic function may be normal or only slightly impaired. However, in advanced cases, the fibrotic process may extend into the myocardium and cause myocardial scarring and atrophy, and venous congestion may then be due to the combined effects of the pericardial and myocardial lesions.

In constrictive pericarditis, the right and left atrial pressure pulses display an M-shaped contour, with prominent *x* and *y* descents. The *y* descent, which is absent or diminished in cardiac tamponade, is the most prominent deflection in constrictive pericarditis; it reflects rapid early filling of the ventricles. The *y* descent is interrupted by a rapid rise in atrial pressure during early diastole, when ventricular filling is impeded by the constricting pericardium. These characteristic changes are transmitted to the jugular veins, where they may be recognized by inspection. In constrictive pericarditis, the ventricular pressure pulses in both ventricles exhibit characteristic “square root” signs during diastole. These hemodynamic changes, although characteristic, are not pathognomonic of constrictive pericarditis and may also be observed in restrictive cardiomyopathies (Chap. 287, Table 287-2).

CLINICAL AND LABORATORY FINDINGS

Weakness, fatigue, weight gain, increased abdominal girth, abdominal discomfort, and edema are common. The patient often appears chronically ill, and in advanced cases, anasarca, skeletal muscle wasting, and cachexia may be present. Exertional dyspnea is common, and orthopnea may occur, although it is usually not severe. Acute left ventricular failure (acute pulmonary edema) is very uncommon. The cervical veins are distended and may remain so even after intensive diuretic treatment, and venous pressure may fail to decline during inspiration (*Kussmaul's sign*). The latter is common in chronic pericarditis but may also occur in tricuspid stenosis, right ventricular infarction, and restrictive cardiomyopathy.

The pulse pressure is normal or reduced. A paradoxical pulse can be detected in about one-third of cases. Congestive hepatomegaly is pronounced and may impair hepatic function and cause jaundice; ascites is common and is usually more prominent than dependent edema. The apical pulse is reduced and may retract in systole (*Broadbent's sign*). The heart sounds may be distant; an early third heart sound (i.e., a pericardial knock, occurring at the cardiac apex 0.09–0.12 s after aortic valve closure) with the abrupt cessation of ventricular filling is often conspicuous.

The ECG frequently displays low voltage of the QRS complexes and diffuse flattening or inversion of the T waves. Atrial fibrillation is present in about one-third of patients. The chest roentgenogram shows a normal or slightly enlarged heart. Pericardial calcification is most common in tuberculous pericarditis. Pericardial calcification may, however, occur in the absence of constriction, and constriction may occur without calcification.

Inasmuch as the usual physical signs of cardiac disease (murmurs, cardiac enlargement) may be inconspicuous or absent in chronic constrictive pericarditis, hepatic enlargement and dysfunction associated with jaundice and intractable ascites may lead to a mistaken diagnosis of hepatic cirrhosis. This error can be avoided if the neck veins are inspected and found to be distended.

The transthoracic echocardiogram typically shows pericardial thickening, dilation of the inferior vena cava and hepatic veins, and a sharp halt in ventricular filling in early diastole, with normal ventricular systolic function and flattening of the left ventricular posterior wall. There is a distinctive pattern of transvalvular flow velocity on Doppler echocardiography. During inspiration, there is an exaggerated reduction in blood flow velocity in the pulmonary veins and across the mitral valve and a leftward shift of the ventricular septum; the opposite occurs during expiration. Diastolic flow velocity in the inferior vena cava into the right atrium and across the tricuspid valve increases in an exaggerated manner during inspiration and declines during expiration (Fig. 288-4). However, echocardiography cannot definitively exclude the diagnosis of constrictive pericarditis. CT and MRI scanning (Fig. 288-5) are

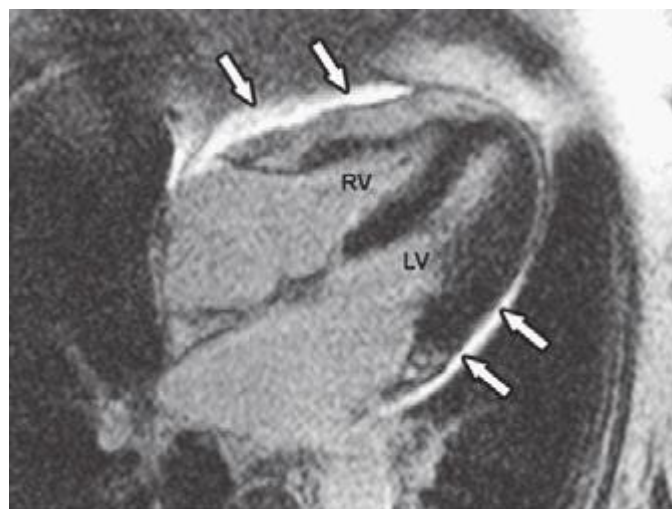


FIGURE 288-5 Magnetic resonance imaging in chronic constrictive pericarditis. The arrows point to a thickened pericardium, which shows late enhancement after gadolinium, characteristic of intense inflammation. LV, left ventricle; RV, right ventricle. (From RY Kwong: *Cardiovascular magnetic resonance imaging*, in RO Bonow et al [eds]: *Braunwald's Heart Disease*, 9th ed. Philadelphia: Elsevier, 2012.)

more accurate than echocardiography in establishing or excluding the presence of a thickened pericardium.

DIFFERENTIAL DIAGNOSIS

Like chronic constrictive pericarditis, cor pulmonale (Chap. 279) may be associated with severe systemic venous hypertension but little pulmonary congestion; the heart is usually not enlarged, and a paradoxical pulse may be present. However, in cor pulmonale, advanced parenchymal pulmonary disease is usually apparent and venous pressure falls during inspiration (i.e., Kussmaul's sign is negative). *Tricuspid stenosis* (Chap. 283) may also simulate chronic constrictive pericarditis; congestive hepatomegaly, splenomegaly, ascites, and venous distention may be equally prominent. However, in tricuspid stenosis, a characteristic murmur and the murmur of accompanying mitral stenosis are usually present.

Because constrictive pericarditis can be corrected surgically, it is important to distinguish chronic constrictive pericarditis from restrictive cardiomyopathy (Chap. 287), which has a similar physiologic abnormality (i.e., restriction of ventricular filling). The differentiating features are summarized in Table 288-2. When a patient has progressive, disabling, and unresponsive congestive heart failure and displays any of the features of constrictive heart disease, Doppler echocardiography to record respiratory effects on transvalvular flow and an MRI or CT scan should be obtained to detect or exclude constrictive pericarditis, because the latter is usually correctable.

TREATMENT CONSTRUCTIVE PERICARDITIS

Pericardial resection is the only definitive treatment of constrictive pericarditis and should be as complete as possible. Dietary sodium restriction and diuretics are useful during preoperative preparation. Coronary arteriography should be carried out preoperatively in patients older than 50 years to exclude unsuspected accompanying coronary artery disease. The benefits derived from cardiac decortication are usually progressive over a period of months. The risk of this operation depends on the extent of penetration of the myocardium by the fibrotic and calcific process, the severity of myocardial atrophy, the extent of secondary impairment of hepatic and/or renal function, and the patient's general condition. Operative mortality is in the range of 5 to 10% even in experienced centers; the patients

with the most severe disease are at highest risk. Therefore, surgical treatment should, if possible, be carried out as early as possible in the course.

Subacute Effusive-Constrictive Pericarditis This form of pericardial disease is characterized by the combination of a tense effusion in the pericardial space and constriction of the heart by thickened pericardium. It shares a number of features with both chronic pericardial effusion producing cardiac compression and with pericardial constriction. It may be caused by tuberculosis (see below), multiple attacks of acute idiopathic pericarditis, radiation, traumatic pericarditis, renal failure, scleroderma, and neoplasms. The heart is generally enlarged, and a paradoxical pulse and a prominent *x* descent (without a prominent *y* descent) are present in the atrial and jugular venous pressure pulses. After pericardiocentesis, the physiologic findings may change from those of cardiac tamponade to those of pericardial constriction. Furthermore, the intrapericardial pressure and the central venous pressure may decline, but not to normal. The diagnosis can be established by pericardiocentesis followed by pericardial biopsy. Wide excision of both the visceral and parietal pericardium is usually effective therapy.

Tuberculous Pericardial Disease This chronic infection is a common cause of chronic pericardial effusion, although less so in North America than in the developing world where active tuberculosis is endemic. The clinical picture is that of a chronic, systemic illness in a patient with pericardial effusion. It is important to consider this diagnosis in a patient with known tuberculosis, with HIV, and with fever, chest pain, weight loss, and enlargement of the cardiac silhouette of undetermined origin. If the etiology of chronic pericardial effusion remains obscure despite detailed analysis of the pericardial fluid (see above), a pericardial biopsy, preferably by a limited thoracotomy, should be performed. If definitive evidence is still lacking but the specimen shows granulomas with caseation, antituberculous chemotherapy (Chap. 202) is indicated.

If the biopsy specimen shows a thickened pericardium after 2–4 weeks of antituberculin therapy, pericardiectomy should be carried out to prevent the development of constriction. Tubercular cardiac constriction should be treated surgically while the patient is receiving antituberculous chemotherapy.

OTHER DISORDERS OF THE PERICARDIUM

Pericardial cysts appear as rounded or lobulated deformities of the cardiac silhouette, most commonly at the right cardiophrenic angle. They usually do not cause symptoms, and their major clinical significance lies in the possibility of confusion with a tumor, ventricular aneurysm, or massive cardiomegaly. *Tumors* involving the pericardium are most commonly secondary to malignant neoplasms originating in or invading the mediastinum, including carcinoma of the bronchus and breast, lymphoma, and melanoma. Mesothelioma is the most common *primary* malignant tumor. The usual clinical picture of malignant pericardial tumor is an insidiously developing, often bloody pericardial effusion. Surgical exploration is required to establish a definitive diagnosis and to carry out definitive or, more commonly, palliative treatment.

289e Tumors and Trauma of the Heart

Eric H. Awtry, Wilson S. Colucci

TUMORS OF THE HEART

PRIMARY TUMORS

Primary tumors of the heart are rare. Approximately three-quarters are histologically benign, and the majority of these tumors are myxomas. Malignant tumors, almost all of which are sarcomas, account for 25% of primary cardiac tumors. All cardiac tumors, regardless of pathologic type, have the potential to cause life-threatening complications. Many tumors are now surgically curable; thus, early diagnosis is imperative.

Clinical Presentation Cardiac tumors may present with a wide array of cardiac and noncardiac manifestations. These manifestations depend in large part on the location and size of the tumor and are often nonspecific features of more common forms of heart disease, such as chest pain, syncope, heart failure, murmurs, arrhythmias, conduction disturbances, and pericardial effusion with or without tamponade. Additionally, embolic phenomena and constitutional symptoms may occur.

Myxoma Myxomas are the most common type of primary cardiac tumor in adults, accounting for one-third to one-half of all cases at postmortem examination, and about three-quarters of the tumors treated surgically. They occur at all ages, most commonly in the third through sixth decades, with a female predilection. Approximately 90% of myxomas are sporadic; the remainder are familial with autosomal dominant transmission. The familial variety often occurs as part of a syndrome complex (Carney complex) that includes (1) myxomas (cardiac, skin, and/or breast), (2) lentiginos and/or pigmented nevi, and (3) endocrine overactivity (primary nodular adrenal cortical disease with or without Cushing's syndrome, testicular tumors, and/or pituitary adenomas with gigantism or acromegaly). Certain constellations of findings have been referred to as the *NAME* syndrome (nevi, atrial myxoma, myxoid neurofibroma, and ephelides) or the *LAMB* syndrome (lentiginos, atrial myxoma, and blue nevi), although these syndromes probably represent subsets of the Carney complex. The genetic basis of this complex has not been elucidated completely; however, patients frequently have inactivating mutations in the tumor-suppressor gene *PRKARIA*, which encodes the protein kinase A type I- α regulatory subunit.

Pathologically, myxomas are gelatinous structures that consist of myxoma cells embedded in a stroma rich in glycosaminoglycans. Most are solitary, arise from the interatrial septum in the vicinity of the fossa ovalis (particularly the left atrium), and are often pedunculated on a fibrovascular stalk. In contrast to sporadic tumors, familial or syndromic tumors tend to occur in younger individuals, are often multiple, may be ventricular in location, and are more likely to recur after initial resection.

Myxomas commonly present with obstructive signs and symptoms. The most common clinical presentation mimics that of mitral valve disease: either stenosis owing to tumor prolapse into the mitral orifice or regurgitation resulting from tumor-induced valvular trauma. Ventricular myxomas may cause outflow obstruction similar to that caused by subaortic or subpulmonic stenosis. The symptoms and signs of myxoma may be sudden in onset or positional in nature, owing to the effects of gravity on tumor position. A characteristic low-pitched sound, a "tumor plop," may be appreciated on auscultation during early or mid-diastole and is thought to result from the impact of the tumor against the mitral valve or ventricular wall. Myxomas also may present with peripheral or pulmonary emboli or with constitutional signs and symptoms, including fever, weight loss, cachexia, malaise, arthralgias, rash, digital clubbing, Raynaud's phenomenon, hypergammaglobulinemia, anemia, polycythemia, leukocytosis, elevated erythrocyte sedimentation rate, thrombocytopenia, and thrombocytosis. These features account for the frequent misdiagnosis of patients with myxomas as having endocarditis, collagen vascular disease, or a paraneoplastic syndrome.

Two-dimensional transthoracic or omniplane transesophageal echocardiography is useful in the diagnosis of cardiac myxoma and allows assessment of tumor size and determination of the site of tumor attachment, both of which are important considerations in the planning of surgical excision (Fig. 289e-1). Computed tomography (CT) and magnetic resonance imaging (MRI) may provide important information regarding size, shape, composition, and surface characteristics of the tumor (Fig. 289e-2).

Although cardiac catheterization and angiography were previously performed routinely before tumor resection, they no longer are considered mandatory when adequate noninvasive information is available and other cardiac disorders (e.g., coronary artery disease) are not considered likely. Additionally, catheterization of the chamber from which the tumor arises carries the risk of tumor embolization. Because myxomas may be familial, echocardiographic screening of first-degree relatives is appropriate, particularly if the patient is young and has multiple tumors or evidence of myxoma syndrome.

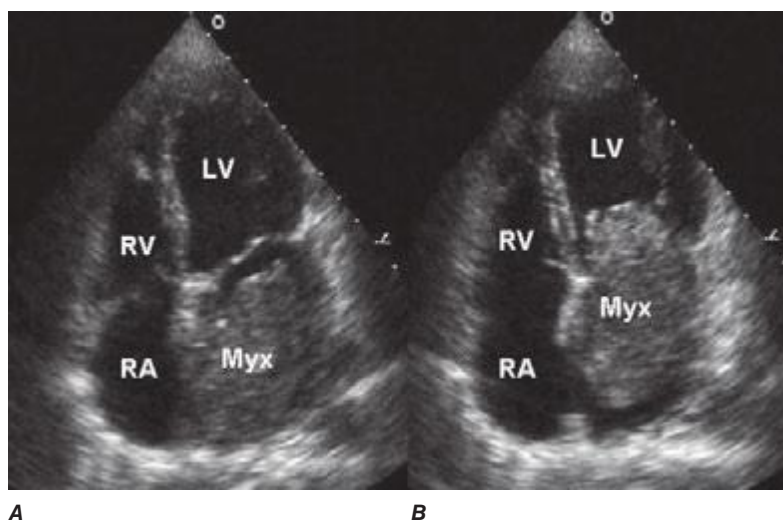


FIGURE 289e-1 Transthoracic echocardiogram demonstrating a large atrial myxoma. The myxoma (Myx) fills the entire left atrium in systole (A) and prolapses across the mitral valve and into the left ventricle (LV) during diastole (B). RA, right atrium; RV, right ventricle. (Courtesy of Dr. Michael Tsang; with permission.)

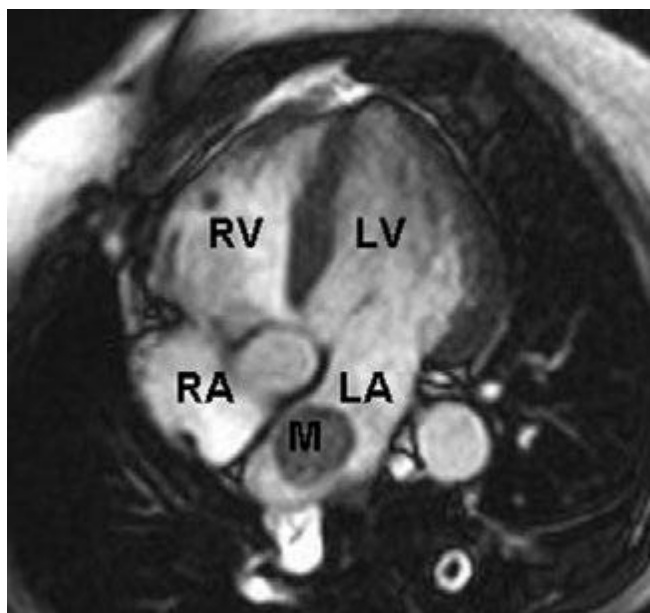


FIGURE 289e-2 Cardiac magnetic resonance imaging demonstrating a rounded mass (M) within the left atrium (LA). Pathologic evaluation at the time of surgery revealed it to be an atrial myxoma. LV, left ventricle; RA, right atrium; RV, right ventricle.

TREATMENT MYXOMA

Surgical excision using cardiopulmonary bypass is indicated regardless of tumor size and is generally curative. Myxomas recur in 12–22% of familial cases but in only 1–2% of sporadic cases. Tumor recurrence most likely is due to multifocal lesions in the former and inadequate resection in the latter.

Other Benign Tumors Cardiac *lipomas*, although relatively common, are usually incidental findings at postmortem examination; however, they may grow as large as 15 cm and may present with symptoms owing to mechanical interference with cardiac function, arrhythmias, or conduction disturbances or as an abnormality of the cardiac silhouette on chest x-ray. *Papillary fibroelastomas* are the most common tumors of the cardiac valves. Although usually clinically silent, they can cause valve dysfunction and may embolize distally, resulting in transient ischemic attacks, stroke, or myocardial infarction. In general, these tumors should be resected even when asymptomatic, although a more conservative approach may be considered for small, right-sided lesions. *Rhabdomyomas* and *fibromas* are the most common cardiac tumors in infants and children and usually occur in the ventricles, where they may produce mechanical obstruction to blood flow, thereby mimicking valvular stenosis, congestive heart failure (CHF), restrictive or hypertrophic cardiomyopathy, or pericardial constriction. *Rhabdomyomas* are probably hamartomatous growths, are multiple in 90% of cases, and are strongly associated with tuberous sclerosis. These tumors have a tendency to regress completely or partially; only tumors that cause obstruction require surgical resection. *Fibromas* are usually single, are often calcified, tend to grow and cause obstructive symptoms, and should be resected. *Hemangiomas* and *mesotheliomas* are generally small tumors, most often intramyocardial in location, and may cause atrioventricular (AV) conduction disturbances and even sudden death as a result of their propensity to develop in the region of the AV node. Other benign tumors arising from the heart include *teratoma*, *chemodectoma*, *neurilemoma*, *granular cell myoblastoma*, and *bronchogenic cysts*.

Sarcoma Almost all malignant primary cardiac tumors are sarcomas, which may be of several histologic types. In general, these tumors are characterized by rapid progression that culminates in the patient's death within weeks to months from the time of presentation as a result

of hemodynamic compromise, local invasion, or distant metastases. Sarcomas commonly involve the right side of the heart, are characterized by rapid growth, frequently invade the pericardial space, and may obstruct the cardiac chambers or venae cavae. Sarcomas also may occur on the left side of the heart and may be mistaken for myxomas.

TREATMENT SARCOMA

At the time of presentation, these tumors have often spread too extensively to allow for surgical excision. Although there are scattered reports of palliation with surgery, radiotherapy, and/or chemotherapy, the response of cardiac sarcomas to these therapies is generally poor. The one exception appears to be cardiac lymphosarcomas, which may respond to a combination of chemo- and radiotherapy.

TUMORS METASTATIC TO THE HEART

Tumors metastatic to the heart are much more common than primary tumors, and their incidence is likely to increase as the life expectancy of patients with various forms of malignant neoplasms is extended by more effective therapy. Although cardiac metastases may occur with any tumor type, the relative incidence is especially high in malignant melanoma and, to a somewhat lesser extent, leukemia and lymphoma. In absolute terms, the most common primary originating sites of cardiac metastases are carcinoma of the breast and lung, reflecting the high incidence of those cancers. Cardiac metastases almost always occur in the setting of widespread primary disease, and most often there is either primary or metastatic disease elsewhere in the thoracic cavity. Nevertheless, cardiac metastasis occasionally may be the initial presentation of an extrathoracic tumor.

Cardiac metastases may occur via hematogenous or lymphangitic spread or by direct tumor invasion. They generally manifest as small, firm nodules; diffuse infiltration also may occur, especially with sarcomas or hematologic neoplasms. The pericardium is most often involved, followed by myocardial involvement of any chamber and, rarely, by involvement of the endocardium or cardiac valves.

Cardiac metastases are clinically apparent only ~10% of the time, are usually not the cause of the patient's presentation, and rarely are the cause of death. The vast majority occur in the setting of a previously recognized malignant neoplasm. As with primary cardiac tumors, the clinical presentation reflects more the location and size of the tumor than its histologic type. When symptomatic, cardiac metastases may result in a variety of clinical features, including dyspnea, acute pericarditis, cardiac tamponade, ectopic tachyarrhythmias, heart block, and CHF. Importantly, many of these signs and symptoms may also result from myocarditis, pericarditis, or cardiomyopathy induced by radiotherapy or chemotherapy.

Electrocardiographic (ECG) findings are nonspecific. On chest x-ray, the cardiac silhouette is most often normal but may be enlarged or exhibit a bizarre contour. Echocardiography is useful for identifying pericardial effusions and visualizing larger metastases, although CT and radionuclide imaging with gallium or thallium may define the tumor burden more clearly. Cardiac MRI offers superb image quality and plays a central role in the diagnostic evaluation of cardiac metastases and cardiac tumors in general. Pericardiocentesis may allow for a specific cytologic diagnosis in patients with malignant pericardial effusions. Angiography is rarely necessary but may delineate discrete lesions.

TREATMENT TUMORS METASTATIC TO THE HEART

Most patients with cardiac metastases have advanced malignant disease; thus, therapy is generally palliative and consists of treatment of the primary tumor. Symptomatic malignant pericardial effusions should be drained by pericardiocentesis. Concomitant instillation of a sclerosing agent (e.g., tetracycline or bleomycin) may delay or prevent reaccumulation of the effusion, and creation of a pericardial window allows drainage of the effusion to the pleural or peritoneal space.

TRAUMATIC CARDIAC INJURY

Traumatic cardiac injury may be caused by either penetrating or nonpenetrating trauma. *Penetrating injuries* most often result from gunshot or knife wounds, and the site of entry is usually obvious. *Nonpenetrating injuries* most often occur during motor vehicle accidents, either from rapid deceleration or from impact of the chest against the steering wheel, and may be associated with significant cardiac injury even in the absence of external signs of thoracic trauma.

NONPENETRATING CARDIAC INJURY

Myocardial contusions are the most common form of nonpenetrating cardiac injury and may initially be overlooked in trauma patients as the clinical focus is directed toward other, more obvious injuries. Myocardial necrosis may occur as a direct result of the blunt injury or as a result of traumatic coronary laceration or thrombosis. The contused myocardium is pathologically similar to infarcted myocardium and may be associated with atrial or ventricular arrhythmias; conduction disturbances, including bundle branch block; or ECG abnormalities resembling those of infarction or pericarditis. Thus, it is important to consider contusion as a cause of otherwise unexplained ECG changes in a trauma patient. Serum creatine kinase, myocardial band (CK-MB) isoenzyme levels are increased in ~20% of patients who experience blunt chest trauma but may be falsely elevated in the presence of massive skeletal muscle injury. Cardiac troponin levels are more specific for identifying cardiac injury in this setting. Echocardiography is useful in detecting structural and functional sequelae of contusion, including wall motion abnormalities (most commonly involving the right ventricle, interventricular septum, or left ventricular apex), pericardial effusion, valvular dysfunction, and ventricular rupture.

Rupture of the cardiac valves or their supporting structures, most commonly of the tricuspid or mitral valve, leads to acute valvular incompetence. This complication is usually heralded by the development of a loud murmur, may be associated with rapidly progressive heart failure, and can be diagnosed by either transthoracic or transesophageal echocardiography.

The most serious consequence of nonpenetrating cardiac injury is myocardial rupture, which may result in hemopericardium and tamponade (free wall rupture) or intracardiac shunting (ventricular septal rupture). Although it generally is fatal, up to 40% of patients with cardiac rupture have been reported to survive long enough to reach a specialized trauma center. Hemopericardium also may result from traumatic rupture of a pericardial vessel or a coronary artery. Additionally, a pericardial effusion may develop weeks or even months after blunt chest trauma as a manifestation of the post-cardiac injury syndrome, which resembles the postpericardiotomy syndrome ([Chap. 288](#)).

Blunt, nonpenetrating, often innocent-appearing injuries to the chest may trigger ventricular fibrillation even in absence of overt signs of injury. This syndrome, referred to as *commotio cordis*, occurs most often in adolescents during sporting events (e.g., baseball, hockey, football, and lacrosse) and probably results from an impact to the chest wall overlying the heart during the susceptible phase of repolarization just before the peak of the T wave. Survival depends on prompt defibrillation. Sudden emotional or physical trauma, even in the absence of direct cardiac trauma, may precipitate a transient catecholamine-mediated

cardiomyopathy referred to as *tako-tsubo syndrome* or the *apical ballooning syndrome* ([Chap. 287](#)).

Rupture or transection of the aorta, usually just above the aortic valve or at the site of the ligamentum arteriosum, is a common consequence of nonpenetrating chest trauma and is the most common vascular deceleration injury. The clinical presentation is similar to that of aortic dissection ([Chap. 301](#)); the arterial pressure and pulse amplitude may be increased in the upper extremities and decreased in the lower extremities, and chest x-ray may reveal mediastinal widening. Occasionally, aortic rupture is contained by the aortic adventitia, resulting in a false, or *pseudo*-, aneurysm that may be discovered months or years after the initial injury.

PENETRATING CARDIAC INJURY

Penetrating injuries of the heart produced by knife or bullet wounds usually result in rapid clinical deterioration and frequently in death as a result of hemopericardium/pericardial tamponade or massive hemorrhage. Nonetheless, up to half of such patients may survive long enough to reach a specialized trauma center if immediate resuscitation is performed. Prognosis in these patients relates to the mechanism of injury, their clinical condition at presentation, and the specific cardiac chamber(s) involved. Iatrogenic cardiac or coronary arterial perforation may complicate placement of central venous or intracardiac catheters, pacemaker leads, or intracoronary stents and is associated with a better prognosis than are other forms of penetrating cardiac trauma.

Traumatic rupture of a great vessel from penetrating injury is usually associated with hemothorax and, less often, hemopericardium. Local hematoma formation may compress major vessels and produce ischemic symptoms, and AV fistulas may develop, occasionally resulting in high-output CHF.

Occasionally, patients who survive penetrating cardiac injuries may subsequently present with a new cardiac murmur or CHF as a result of mitral regurgitation or an intracardiac shunt (i.e., ventricular or atrial septal defect, aortopulmonary fistula, or coronary AV fistula) that was undetected at the time of the initial injury or developed subsequently. Therefore, trauma patients should be examined carefully several weeks after the injury. If a mechanical complication is suspected, it can be confirmed by echocardiography or cardiac catheterization.

TREATMENT TRAUMATIC CARDIAC INJURY

The treatment of an uncomplicated myocardial contusion is similar to the medical therapy for a myocardial infarction, except that anticoagulation is contraindicated, and should include monitoring for the development of arrhythmias and mechanical complications such as cardiac rupture ([Chap. 295](#)). Acute myocardial failure resulting from traumatic valve rupture usually requires urgent operative correction. Immediate thoracotomy should be carried out for most cases of penetrating injury, or if there is evidence of cardiac tamponade and/or shock regardless of the type of trauma. Pericardiocentesis may be lifesaving in patients with tamponade but is usually only a temporizing measure while awaiting definitive surgical therapy. Pericardial hemorrhage often leads to constriction ([Chap. 288](#)), which must be treated by surgical decortication.

290e Cardiac Manifestations of Systemic Disease

Eric H. Awtry, Wilson S. Colucci

The common systemic disorders that have associated cardiac manifestations are summarized in [Table 290e-1](#).

DIABETES MELLITUS

(See also [Chap. 417](#)) Diabetes mellitus, both insulin- and non-insulin-dependent, is an independent risk factor for coronary artery disease (CAD; [Chap. 291e](#)) and accounts for 14–50% of new cases of cardiovascular disease. Furthermore, CAD is the most common cause of death in adults with diabetes mellitus. In the diabetic population, the incidence of CAD relates to the duration of diabetes and the level of glycemic control, and its pathogenesis involves endothelial dysfunction, increased lipoprotein peroxidation, increased inflammation, a prothrombotic state, and associated metabolic abnormalities.

Compared to their nondiabetic counterparts, diabetic patients are more likely to have a myocardial infarction, have a greater burden of CAD, have larger infarct size, and have more postinfarct complications, including heart failure, shock, and death. Importantly, diabetic patients are more likely to have atypical ischemic symptoms; nausea, dyspnea, pulmonary edema, arrhythmias, heart block, or syncope may be their anginal equivalent. Additionally, “silent ischemia,” resulting from autonomic nervous system dysfunction, is more common in diabetic patients, accounting for up to 90% of their ischemic episodes. Thus, one must have a low threshold for suspecting CAD in diabetic patients. The treatment of diabetic patients with CAD must include aggressive risk factor management ([Chap. 418](#)). Considerations regarding pharmacologic therapy and revascularization strategies are similar in diabetic and nondiabetic patients except that diabetic patients have higher morbidity and mortality rates associated with revascularization, have an increased risk of restenosis after percutaneous coronary intervention (PCI), and have improved survival when treated with surgical bypass compared with PCI for multivessel CAD.

Patients with diabetes mellitus also may have abnormal left ventricular systolic and diastolic function, reflecting concomitant epicardial CAD and/or hypertension, coronary microvascular disease, endothelial dysfunction, ventricular hypertrophy, and autonomic dysfunction. Furthermore, the increase in intramyocardial lipid deposition (predominantly nonesterified fatty acids) that is characteristic of diabetic states may contribute to both systolic and diastolic dysfunction by impairing insulin signaling, reducing trans-sarcolemma calcium flux, and inducing myocyte apoptosis. A restrictive cardiomyopathy may be present with abnormal myocardial relaxation and elevated ventricular filling pressures. Histologically, interstitial fibrosis is seen, and intramural arteries may demonstrate intimal thickening, hyaline deposition, and inflammatory changes. Diabetic patients have an increased risk of developing clinical heart failure, which probably contributes to their excessive cardiovascular morbidity and mortality rates. There is some evidence that insulin therapy may ameliorate diabetes-related myocardial dysfunction.

MALNUTRITION AND VITAMIN DEFICIENCY

Malnutrition (See also [Chap. 97](#)) In patients whose intake of protein, calories, or both is severely deficient, the heart may become thin, pale, and hypokinetic with myofibrillar atrophy and interstitial edema. The systolic pressure and cardiac output fall, and the pulse pressure narrows. Generalized edema is common and relates to a variety of factors, including reduced serum oncotic pressure and myocardial dysfunction. Such profound states of protein and calorie malnutrition, termed *kwashiorkor* and *marasmus*, respectively, are most common in underdeveloped countries. However, significant nutritional heart disease also may occur in developed nations, particularly in patients with chronic diseases such as AIDS, patients with anorexia nervosa, and patients with severe cardiac failure in whom gastrointestinal hypoperfusion and venous congestion may lead to anorexia and malabsorption. Open-heart surgery poses increased risk in malnourished patients; such patients may benefit from preoperative hyperalimentation.

Thiamine Deficiency (Beriberi) (See also [Chap. 96e](#)) Generalized malnutrition often is accompanied by thiamine deficiency; however, this hypovitaminosis also may occur in the presence of an adequate protein and caloric intake, particularly in East Asia, where polished rice

TABLE 290e-1 COMMON SYSTEMIC DISORDERS AND THEIR ASSOCIATED CARDIAC MANIFESTATIONS

Systemic Disorder	Common Cardiac Manifestations	Chapter
Diabetes mellitus	CAD, atypical angina, CMP, systolic or diastolic CHF	417
Protein-calorie malnutrition	Dilated CMP, CHF	97
Thiamine deficiency	High-output failure, dilated CMP	96e
Hyperhomocysteinemia	Premature atherosclerosis	96e
Obesity	CMP, systolic or diastolic CHF	415e
Hyperthyroidism	Palpitations, SVT, atrial fibrillation, hypertension	405
Hypothyroidism	Hypotension, bradycardia, dilated CMP, CHF, pericardial effusion	405
Malignant carcinoid	Tricuspid and pulmonary valve disease, right heart failure	113
Pheochromocytoma	Hypertension, palpitations, CHF	407
Acromegaly	Systolic or diastolic heart failure	401e
Rheumatoid arthritis	Pericarditis, pericardial effusions, coronary arteritis, myocarditis, valvulitis	380
Seronegative arthropathies	Aortitis, aortic and mitral insufficiency, conduction abnormalities	384
Systemic lupus erythematosus	Pericarditis, Libman-Sacks endocarditis, myocarditis, arterial and venous thrombosis	378
HIV	Myocarditis, dilated CMP, pericardial effusion	226
Amyloidosis	CHF, restrictive CMP, valvular regurgitation, pericardial effusion	137
Sarcoidosis	CHF, dilated or restrictive CMP, ventricular arrhythmias, heart block	390
Hemochromatosis	CHF, arrhythmias, heart block	428
Marfan syndrome	Aortic aneurysm and dissection, aortic insufficiency, mitral valve prolapse	427
Ehlers-Danlos syndrome	Aortic and coronary aneurysms, mitral and tricuspid valve prolapse	427
Systemic sclerosis	Pericardial effusion, CHF (systolic and diastolic), myocarditis, coronary microvascular vasospasm, tachyarrhythmias	382

Abbreviations: CAD, coronary artery disease; CHF, congestive heart failure; CMP, cardiomyopathy; SVT, supraventricular tachycardia.

deficient in thiamine may be a major dietary component. In Western nations where the use of thiamine-enriched flour is widespread, clinical thiamine deficiency is limited primarily to alcoholics, food faddists, and patients receiving chemotherapy. Nonetheless, when thiamine stores are measured using the thiamine-pyrophosphate effect (TPPE), thiamine deficiency has been found in 20–90% of patients with chronic heart failure. This deficiency appears to result from both reduced dietary intake and a diuretic-induced increase in the urinary excretion of thiamine. The acute administration of thiamine to these patients increases the left ventricular ejection fraction and the excretion of salt and water.

Clinically, patients with thiamine deficiency usually have evidence of generalized malnutrition, peripheral neuropathy, glossitis, and anemia. The classic associated cardiovascular syndrome is characterized by high-output heart failure, tachycardia, and often elevated biventricular filling pressures. The major cause of the high-output state is vasomotor depression leading to reduced systemic vascular resistance, the precise mechanism of which is not understood. The cardiac examination may reveal a wide pulse pressure, tachycardia, a third heart sound, and an apical systolic murmur. The electrocardiogram (ECG) may reveal decreased voltage, a prolonged QT interval, and T-wave abnormalities. The chest x-ray generally reveals cardiomegaly and signs of congestive heart failure (CHF). The response to thiamine is often dramatic, with an increase in systemic vascular resistance, a decrease in cardiac output, clearing of pulmonary congestion, and a reduction in heart size often occurring in 12–48 h. Although the response to inotropes and diuretics may be poor before thiamine therapy, these agents may be important *after* thiamine repletion, since the left ventricle may not be able to handle the increased work load presented by the return of vascular tone.

Vitamin B₆, B₁₂, and Folate Deficiency (See also Chap. 96e) Vitamin B₆, vitamin B₁₂, and folate are cofactors in the metabolism of homocysteine. Their deficiency probably contributes to the majority of cases of hyperhomocysteinemia, a disorder associated with increased atherosclerotic risk. Supplementation of these vitamins has reduced the incidence of hyperhomocysteinemia in the United States; however, the clinical cardiovascular benefit of normalizing elevated homocysteine levels has not been proved.

OBESITY

(See also Chap. 415e) Obesity is associated with an increased prevalence of hypertension, glucose intolerance, atherosclerotic CAD, atrial fibrillation, obstructive sleep apnea, and pulmonary hypertension, and is associated with increased cardiovascular morbidity and mortality rates. In addition, obese patients have a distinct hemodynamic profile characterized by increased total and central blood volumes, increased cardiac output, and elevated left ventricular filling pressure. The elevated cardiac output appears to be required to support the metabolic demands of the excess adipose tissue. Left ventricular filling pressure is often at the upper limits of normal at rest and rises excessively with exercise, contributing to exertional dyspnea. In part as a result of chronic volume overload, eccentric cardiac hypertrophy with cardiac dilation and ventricular diastolic and/or systolic dysfunction may develop. In addition, altered levels of adipokines secreted by adipose tissue may contribute to adverse myocardial remodeling via direct effects on cardiac myocytes and other cells. Pathologically, there is left and, in some cases, right ventricular hypertrophy and generalized cardiac dilation. Pulmonary congestion, peripheral edema, and exercise intolerance may all ensue; however, the recognition of these findings may be difficult in massively obese patients.

Treatment with angiotensin-converting enzyme inhibitors, sodium restriction, and diuretics may be useful to control heart failure symptoms. Weight reduction, however, is the most effective therapy and results in reduction in blood volume and the return of cardiac output toward normal. However, rapid weight reduction may be dangerous, as cardiac arrhythmias and sudden death owing to electrolyte imbalance have been described.

THYROID DISEASE

(See also Chap. 405) Thyroid hormone exerts a major influence on the cardiovascular system by a number of direct and indirect mechanisms, and not surprisingly, cardiovascular effects are prominent in both hypo- and hyperthyroidism. Thyroid hormone causes increases in total-body metabolism and oxygen consumption that indirectly increase the cardiac workload. In addition, thyroid hormone exerts direct inotropic, chronotropic, and dromotropic effects that are similar to those seen with adrenergic stimulation (e.g., tachycardia, increased cardiac output); they are mediated at least partly by both transcriptional and nontranscriptional effects of thyroid hormone on myosin, calcium-activated ATPase, Na⁺-K⁺-ATPase, and myocardial β-adrenergic receptors.

Hyperthyroidism Common cardiovascular manifestations of hyperthyroidism include palpitations, systolic hypertension, and fatigue. Sinus tachycardia is present in ~40% of hyperthyroid patients, and atrial fibrillation is present in ~15%. Physical examination may reveal a hyperdynamic precordium, a widened pulse pressure, increases in the intensity of the first heart sound and the pulmonic component of the second heart sound, and a third heart sound. An increased incidence of mitral valve prolapse has been described in hyperthyroid patients, in which case a midsystolic murmur may be heard at the left sternal border with or without a midsystolic click. A systolic pleuropericardial friction rub (*Means-Lerman scratch*) may be heard at the left second intercostal space during expiration and is thought to result from the hyperdynamic cardiac motion.

Elderly patients with hyperthyroidism may present with only cardiovascular manifestations of thyrotoxicosis such as sinus tachycardia, atrial fibrillation, and hypertension, all of which may be resistant to therapy until the hyperthyroidism is controlled. Angina pectoris and CHF are unusual with hyperthyroidism unless there is coexistent heart disease; in such cases, symptoms often resolve with treatment of the hyperthyroidism.

Hypothyroidism Cardiac manifestations of hypothyroidism include a reduction in cardiac output, stroke volume, heart rate, systolic blood pressure, and pulse pressure. Pericardial effusions are present in about one-third of patients, rarely progress to tamponade, and probably result from increased capillary permeability. Other clinical signs include cardiomegaly, bradycardia, weak arterial pulses, distant heart sounds, and pleural effusions. Although the signs and symptoms of myxedema may mimic those of CHF, in the absence of other cardiac disease, myocardial failure is uncommon. The ECG generally reveals sinus bradycardia and low voltage and may show prolongation of the QT interval, decreased P-wave voltage, prolonged AV conduction time, intraventricular conduction disturbances, and nonspecific ST-T-wave abnormalities. Chest x-ray may show cardiomegaly, often with a “water bottle” configuration; pleural effusions; and, in some cases, evidence of CHF. Pathologically, the heart is pale and dilated and often demonstrates myofibrillar swelling, loss of striations, and interstitial fibrosis.

Patients with hypothyroidism frequently have elevations of cholesterol and triglycerides, resulting in premature atherosclerotic CAD. Before treatment with thyroid hormone, patients with hypothyroidism frequently do not have angina pectoris, presumably because of the low metabolic demands caused by their condition. However, angina and myocardial infarction may be precipitated during initiation of thyroid hormone replacement, especially in elderly patients with underlying heart disease. Therefore, replacement should be done with care, starting with low doses that are increased gradually.

MALIGNANT CARCINOID

(See also Chap. 113) Carcinoid tumors most often originate in the small bowel and elaborate a variety of vasoactive amines (e.g., serotonin), kinins, indoles, and prostaglandins that are believed to be responsible for the diarrhea, flushing, and labile blood pressure that characterize the carcinoid syndrome. Some 50% of patients with carcinoid syndrome have cardiac involvement, usually manifesting as abnormalities of the tricuspid or pulmonic valves. These patients

invariably have hepatic metastases that allow vasoactive substances to circumvent hepatic metabolism. Left-sided cardiac involvement is rare and indicates either pulmonary carcinoid or an intracardiac shunt. Pathologically, carcinoid lesions are fibrous plaques that consist of smooth-muscle cells embedded in a stroma of glycosaminoglycans and collagen. They occur on the cardiac valves, where they cause valvular dysfunction, as well as on the endothelium of the cardiac chambers and great vessels.

Carcinoid heart disease most often presents as tricuspid regurgitation, pulmonic stenosis, or both. In some cases, a high cardiac output state may occur, presumably as a result of a decrease in systemic vascular resistance resulting from vasoactive substances released by the tumor. Treatment with somatostatin analogues (e.g., octreotide) or interferon α improves symptoms and survival in patients with carcinoid heart disease but does not appear to improve valvular abnormalities. Treatment with diuretics usually mitigates the symptoms of right heart failure; in some severely symptomatic patients, valve replacement is indicated. Coronary artery spasm, presumably due to a circulating vasoactive substance, may occur in patients with carcinoid syndrome.

PHEOCHROMOCYTOMA

(See also Chap. 407) In addition to causing labile or sustained hypertension, the high circulating levels of catecholamines resulting from a pheochromocytoma may cause direct myocardial injury. Focal myocardial necrosis and inflammatory cell infiltration are present in ~50% of patients who die with pheochromocytoma and may contribute to clinically significant left ventricular failure and pulmonary edema. In addition, associated hypertension results in left ventricular hypertrophy. Left ventricular dysfunction and CHF may resolve after removal of the tumor.

ACROMEGALY

(See also Chap. 401e) Exposure of the heart to excessive growth hormone may cause CHF as a result of high cardiac output, diastolic dysfunction owing to ventricular hypertrophy (with increased left ventricular chamber size or wall thickness), or global systolic dysfunction. Hypertension occurs in up to one-third of patients with acromegaly and is characterized by suppression of the renin-angiotensin-aldosterone axis and increases in total-body sodium and plasma volume. Some form of cardiac disease occurs in about one-third of patients with acromegaly and is associated with a doubling of the risk of cardiac death.

RHEUMATOID ARTHRITIS AND THE COLLAGEN VASCULAR DISEASES

Rheumatoid Arthritis (See also Chap. 380) Rheumatoid arthritis may be associated with inflammatory changes in any or all cardiac structures, although pericarditis is the most common clinical entity. Pericardial effusions are found on echocardiography in 10–50% of patients with rheumatoid arthritis, particularly those with subcutaneous nodules. Nonetheless, only a small fraction of these patients have symptomatic pericarditis, and when present, it usually follows a benign course, only occasionally progressing to cardiac tamponade or constrictive pericarditis. The pericardial fluid is generally exudative,

with decreased concentrations of complement and glucose and elevated cholesterol. Coronary arteritis with intimal inflammation and edema is present in ~20% of cases but only rarely results in angina pectoris or myocardial infarction. Inflammation and granuloma formation may affect the cardiac valves, most often the mitral and aortic valves, and may cause clinically significant regurgitation owing to valve deformity. Myocarditis is uncommon and rarely results in cardiac dysfunction.

Treatment is directed at the underlying rheumatoid arthritis and may include glucocorticoids. Urgent pericardiocentesis should be performed in patients with tamponade, but pericardiectomy usually is required in cases of pericardial constriction.

Seronegative Arthropathies (See also Chap. 384) The seronegative arthropathies, including ankylosing spondylitis, reactive arthritis, psoriatic arthritis, and the arthritides associated with ulcerative colitis and regional enteritis, are all strongly associated with the HLA-B27 histocompatibility antigen and may be accompanied by a pancarditis and proximal aortitis. The aortic inflammation usually is limited to the aortic root but may extend to involve the aortic valve, mitral valve, and ventricular myocardium, resulting in aortic and mitral regurgitation, conduction abnormalities, and ventricular dysfunction. One-tenth of these patients have significant aortic insufficiency, and one-third have conduction disturbances; both are more common in patients with peripheral joint involvement and long-standing disease. Treatment with aortic valve replacement and permanent pacemaker implantation may be required. Occasionally, aortic regurgitation precedes the onset of arthritis, and therefore, the diagnosis of a seronegative arthritis should be considered in young males with isolated aortic regurgitation.

Systemic Lupus Erythematosus (SLE) (See also Chap. 378) A significant percentage of patients with SLE have cardiac involvement. Pericarditis is common, occurring in about two-thirds of patients, and generally follows a benign course, although rarely tamponade or constriction may result. The characteristic endocardial lesions of SLE are verrucous valvular abnormalities known as *Libman-Sacks endocarditis*. They most often are located on the left-sided cardiac valves, particularly on the ventricular surface of the posterior mitral leaflet, and are made up almost entirely of fibrin. These lesions may embolize or become infected but rarely cause hemodynamically important valvular regurgitation. Myocarditis generally parallels the activity of the disease and, although common histologically, seldom results in clinical heart failure unless associated with hypertension. Although arteritis of epicardial coronary arteries may occur, it rarely results in myocardial ischemia. There is, however, an increased incidence of coronary atherosclerosis that probably is related more to associated risk factors and glucocorticoid use than to SLE itself. Patients with the antiphospholipid antibody syndrome may have a higher incidence of cardiovascular abnormalities, including valvular regurgitation, venous and arterial thrombosis, premature stroke, myocardial infarction, pulmonary hypertension, and cardiomyopathy.

291e

The Pathogenesis, Prevention, and Treatment of Atherosclerosis

Peter Libby

PATHOGENESIS

Atherosclerosis remains the major cause of death and premature disability in developed societies. Moreover, current predictions estimate that by the year 2020 cardiovascular diseases, notably atherosclerosis, will become the leading global cause of total disease burden. Although many generalized or systemic risk factors predispose to its development, atherosclerosis affects various regions of the circulation preferentially and has distinct clinical manifestations that depend on the particular circulatory bed affected. Atherosclerosis of the coronary arteries commonly causes myocardial infarction (MI) (Chap. 295) and angina pectoris (Chap. 293). Atherosclerosis of the arteries supplying the central nervous system frequently provokes strokes and transient cerebral ischemia (Chap. 446). In the peripheral circulation, atherosclerosis causes intermittent claudication and gangrene and can jeopardize limb viability. Involvement of the splanchnic circulation can cause mesenteric ischemia. Atherosclerosis can affect the kidneys either directly (e.g., renal artery stenosis) or as a common site of atheroembolic disease (Chap. 301).

Even within a particular arterial bed, stenoses due to atherosclerosis tend to occur focally, typically in certain predisposed regions. In the coronary circulation, for example, the proximal left anterior descending coronary artery exhibits a particular predilection for developing atherosclerotic disease. Similarly, atherosclerosis preferentially affects the proximal portions of the renal arteries and, in the extracranial circulation to the brain, the carotid bifurcation. Indeed, atherosclerotic lesions often form at branch points of arteries, regions characterized by disturbed hydrodynamics. Not all manifestations of atherosclerosis result from stenotic, occlusive disease. Ectasia and the development of aneurysmal disease, for example, frequently occur in the aorta (Chap. 301). In addition to focal, flow-limiting stenoses, nonocclusive intimal atherosclerosis also occurs diffusely in affected arteries, as shown by intravascular imaging and postmortem studies.

Atherogenesis in humans typically occurs over a period of many years, usually many decades. Growth of atherosclerotic plaques probably does not occur in a smooth, linear fashion but discontinuously, with periods of relative quiescence punctuated by periods of rapid evolution. After a generally prolonged “silent” period, atherosclerosis may become clinically manifest. The clinical expressions of atherosclerosis may be *chronic*, as in the development of stable, effort-induced angina pectoris or predictable and reproducible intermittent claudication. Alternatively, a dramatic *acute* clinical event such as MI, stroke, or sudden cardiac death may first herald the presence of atherosclerosis. Other individuals may never experience clinical manifestations of arterial disease despite the presence of widespread atherosclerosis demonstrated postmortem.

INITIATION OF ATHEROSCLEROSIS

An integrated view of experimental results in animals and studies of human atherosclerosis suggests that the “fatty streak” represents the initial lesion of atherosclerosis. These early lesions most often seem to arise from focal increases in the content of lipoproteins within regions of the intima. In particular, the fraction of lipoproteins related to low-density lipoprotein (LDL) that bear apolipoprotein B appear causally related to atherosclerosis. This accumulation of lipoprotein particles may not result simply from increased permeability, or

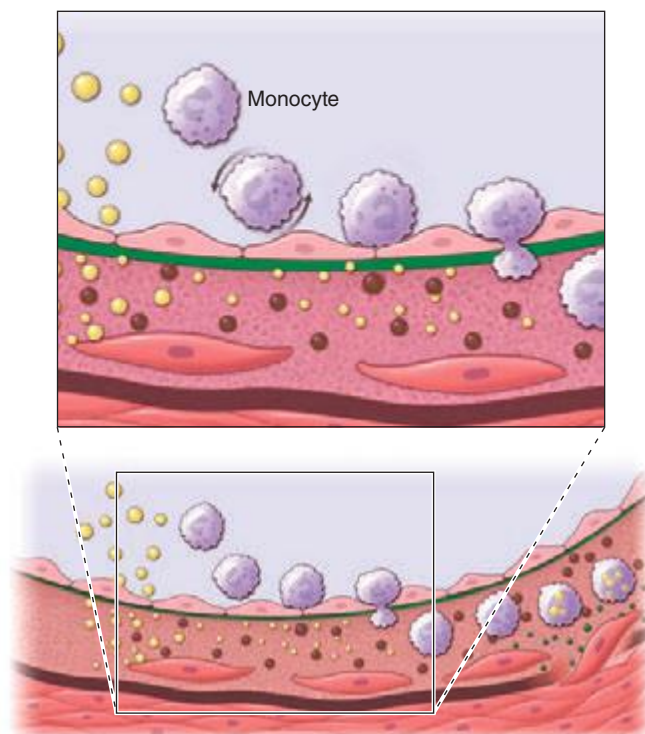


FIGURE 291e-1 Cross-sectional view of an artery depicting steps in development of an atheroma, from left to right.

The upper panel shows a detail of the boxed area below. The endothelial monolayer overlying the intima contacts blood. Hypercholesterolemia promotes accumulation of low-density lipoprotein (LDL) particles (yellow spheres) in the intima. The lipoprotein particles often associate with constituents of the extracellular matrix, notably proteoglycans. Sequestration within the intima separates lipoproteins from some plasma antioxidants and favors oxidative modification. Such modified lipoprotein particles (darker spheres) may trigger a local inflammatory response that signals subsequent steps in lesion formation. The augmented expression of various adhesion molecules for leukocytes recruits monocytes to the site of a nascent arterial lesion.

Once adherent, some white blood cells migrate into the intima. The directed migration of leukocytes probably depends on chemoattractant factors, including modified lipoprotein particles themselves and chemoattractant cytokines (depicted by the smaller green spheres), such as the chemokine macrophage chemoattractant protein-1 produced by vascular wall cells in response to modified lipoproteins. Leukocytes in the evolving fatty streak can divide and exhibit augmented expression of receptors for modified lipoproteins (scavenger receptors). These mononuclear phagocytes ingest lipids and become foam cells, represented by a cytoplasm filled with lipid droplets. As the fatty streak evolves into a more complicated atherosclerotic lesion, smooth-muscle cells migrate from the media (bottom of lower panel hairline) through the internal elastic membrane (solid wavy line) and accumulate within the expanding intima, where they lay down extracellular matrix that forms the bulk of the advanced lesion (bottom panel, right side).

“leakiness,” of the overlying endothelium (Fig. 291e-1). Rather, the lipoproteins may collect in the intima of arteries because they bind to constituents of the extracellular matrix, increasing the residence time of the lipid-rich particles within the arterial wall. Lipoproteins that accumulate in the extracellular space of the intima of arteries often associate with proteoglycans of the arterial extracellular matrix, an interaction that may slow the egress of these lipid-rich particles from

the intima. Lipoprotein particles in the extracellular space of the intima, particularly those retained by binding to matrix macromolecules, may undergo oxidative modifications. Considerable evidence supports a pathogenic role for products of oxidized lipoproteins in atherogenesis. Lipoproteins sequestered from (plasma) antioxidants in the extracellular space of the intima become particularly susceptible to oxidative modification, giving rise to hydroperoxides, lysophospholipids, oxysterols, and aldehydic breakdown products of fatty acids and phospholipids. Modifications of the apoprotein moieties may include breaks in the peptide backbone as well as derivatization of certain amino acid residues. Local production of hypochlorous acid by myeloperoxidase associated with inflammatory cells within the plaque yields chlorinated species such as chlorotyrosyl moieties. Considerable evidence supports the presence of such oxidation products in atherosclerotic lesions.

Leukocyte Recruitment Accumulation of leukocytes characterizes the formation of early atherosclerotic lesions (Fig. 291e-1). Thus, from its very inception, atherogenesis involves elements of inflammation, a process that now provides a unifying theme in the pathogenesis of this disease. The inflammatory cell types typically found in the evolving atheroma include monocyte-derived macrophages and dendritic cells, T and B lymphocytes, and mast cells. Hypercholesterolemia augments the portion of particularly proinflammatory monocytes in blood that preferentially enter the nascent atheroma in mice. A number of adhesion molecules or receptors for leukocytes expressed on the surface of the arterial endothelial cell probably participate in the recruitment of leukocytes to the nascent atheroma. Proinflammatory cytokines can augment the expression of leukocyte adhesion molecules.

Laminar shear forces such as those encountered in most regions of normal arteries also can suppress the expression of leukocyte adhesion molecules. Sites of predilection for atherosclerotic lesions (e.g., distal to flow dividers) often have low shear stress and/or disturbed flow. Ordered, pulsatile laminar shear of normal blood flow augments the production of nitric oxide by endothelial cells. This molecule, in addition to its vasodilator properties, can act at the low levels constitutively produced by arterial endothelium as a local anti-inflammatory autacoid, e.g., limiting local adhesion molecule expression. Exposure of endothelial cells to laminar shear stress increases the transcription of Krüppel-like factor 2 (KLF2), which augments the activity of numerous salutary endothelial functions including nitric oxide synthase. Laminar shear stress also stimulates endothelial cells to produce superoxide dismutase, an antioxidant enzyme. These examples indicate how hemodynamic forces may influence the cellular events that underlie atherosclerotic lesion initiation and potentially explain the favored localization of atherosclerotic lesions at sites that experience disturbed flow or low shear stress.

Once captured on the surface of the arterial endothelial cell by adhesion receptors, the leukocytes penetrate the endothelial layer and take up residence in the intima. In addition to products of modified lipoproteins, cytokines (protein mediators of inflammation) can regulate the expression of adhesion molecules involved in leukocyte recruitment. For example, interleukin 1 (IL-1) and tumor necrosis factor (TNF) induce or augment the expression of leukocyte adhesion molecules on endothelial cells. Because products of lipoprotein oxidation can induce cytokine release from vascular wall cells, this pathway may provide an additional link between arterial accumulation of lipoproteins and leukocyte recruitment. Chemoattractant cytokines appear to direct the migration of leukocytes into the arterial wall.

Foam-Cell Formation Once resident within the intima, the mononuclear phagocytes mature into macrophages and become lipid-laden foam cells, a conversion that requires the uptake of lipoprotein particles by receptor-mediated endocytosis. One might suppose that the “classic” LDL receptor mediates this lipid uptake; however, humans or animals lacking effective LDL receptors due to genetic alterations (e.g., familial hypercholesterolemia) have abundant arterial lesions and extraarterial xanthomata rich in macrophage-derived foam cells. In addition, the exogenous cholesterol suppresses expression of the LDL receptor; thus, the level of this cell-surface receptor for LDL decreases under

conditions of cholesterol excess. Candidates for alternative receptors that can mediate lipid loading of foam cells include a number of macrophage “scavenger” receptors, which preferentially endocytose modified lipoproteins, and other receptors for oxidized LDL or very low-density lipoprotein (VLDL). Monocyte attachment to the endothelium, migration into the intima, and maturation to form lipid-laden macrophages thus represent key steps in the formation of the fatty streak, the precursor of fully formed atherosclerotic plaques.

ATHEROMA EVOLUTION AND COMPLICATIONS

Although the fatty streak commonly precedes the development of a more advanced atherosclerotic plaque, not all fatty streaks progress to form complex atheromata. By ingesting lipids from the extracellular space, the mononuclear phagocytes bearing such scavenger receptors may remove lipoproteins from the developing lesion. Some lipid-laden macrophages may leave the artery wall, exporting lipid in the process. Lipid accumulation, and hence the propensity to form an atheroma, ensues if the amount of lipid entering the artery wall exceeds that removed by mononuclear phagocytes or other pathways. Macrophages also proliferate in plaques in response to hematopoietic growth factors overexpressed in lesions, another aspect of the dynamic regulation and flux of cells during atherogenesis.

Export by phagocytes may constitute one response to local lipid overload in the evolving lesion. Another mechanism, reverse cholesterol transport mediated by high-density lipoproteins (HDLs), probably provides an independent pathway for lipid removal from atheroma. This transfer of cholesterol from the cell to the HDL particle involves specialized cell-surface molecules such as the ATP binding cassette (ABC) transporters. *ABCA1*, the gene mutated in Tangier disease, a condition characterized by very low HDL levels, transfers cholesterol from cells to nascent HDL particles and *ABCG1* to mature HDL particles. “Reverse cholesterol transport” mediated by these ABC transporters allows HDL loaded with cholesterol to deliver it to hepatocytes by binding to scavenger receptor B1 or other receptors. The liver cell can metabolize the sterol to bile acids that can be excreted. Thus, macrophages may play a vital role in the dynamic economy of lipid accumulation in the arterial wall during atherogenesis.

Some lipid-laden foam cells within the expanding intimal lesion perish. Some foam cells may die as a result of programmed cell death, or *apoptosis*. This death of mononuclear phagocytes results in the formation of the lipid-rich center, often called the *necrotic core*, in established atherosclerotic plaques. Impaired clearance of dead foam cells (efferocytosis) in plaques may hasten lipid core formation. Macrophages loaded with modified lipoproteins may elaborate microparticles or exosomes (which may contain regulatory microRNAs), cytokines, and growth factors that can further signal some of the cellular events in lesion complication. Whereas accumulation of lipid-laden macrophages characterizes the fatty streak, buildup of fibrous tissue formed by extracellular matrix typifies the more advanced atherosclerotic lesion. The smooth-muscle cell synthesizes the bulk of the extracellular matrix of the complex atherosclerotic lesion. A number of growth factors or cytokines elaborated by mononuclear phagocytes can stimulate smooth-muscle cell proliferation and production of extracellular matrix. Cytokines found in the plaque, including IL-1 and TNF, can induce local production of growth factors, including forms of platelet-derived growth factor (PDGF), fibroblast growth factors, and others, which may contribute to plaque evolution and complication. Other cytokines, notably interferon γ (IFN- γ) derived from activated T cells within lesions, can limit the synthesis of interstitial forms of collagen by smooth-muscle cells. These examples illustrate how atherogenesis involves a complex mix of mediators that in the balance determines the characteristics of particular lesions.

The accumulation of smooth-muscle cells and their elaboration of extracellular matrix probably provide a critical transition, yielding a fibrofatty lesion in place of a simple accumulation of macrophage-derived foam cells. For example, PDGF elaborated by activated platelets, macrophages, and endothelial cells can stimulate the migration of smooth-muscle cells normally resident in the tunica media into the intima. Such growth factors and cytokines produced locally can

stimulate the proliferation of resident smooth-muscle cells or resident stem cells in the intima as well as those that may migrate in from the media. Transforming growth factor β (TGF- β), among other mediators, potently stimulates interstitial collagen production by smooth-muscle cells. These mediators may arise not only from neighboring vascular cells or leukocytes (a “paracrine” pathway), but also, in some instances, from the same cell that responds to the factor (an “autocrine” pathway). Together, these alterations in smooth-muscle cells, signaled by these mediators acting at short distances, can hasten transformation of the fatty streak into a more fibrous smooth-muscle cell and extracellular matrix—rich lesion.

In addition to locally produced mediators, products of blood coagulation and thrombosis likely contribute to atheroma evolution and complication. This involvement justifies the use of the term *atherothrombosis* to convey the inextricable links between atherosclerosis and thrombosis. Fatty streak formation begins beneath a morphologically intact endothelium. In advanced fatty streaks, however, microscopic breaches in endothelial integrity may occur. Microthrombi rich in platelets can form at such sites of limited endothelial denudation, owing to exposure of the thrombogenic extracellular matrix of the underlying basement membrane. Activated platelets release numerous factors that can promote the fibrotic response, including PDGF and TGF- β . Thrombin not only generates fibrin during coagulation, but also stimulates protease-activated receptors that can signal smooth-muscle migration, proliferation, and extracellular matrix production. Many arterial mural microthrombi resolve without clinical manifestation by a process of local fibrinolysis, resorption, and endothelial repair, yet can lead to lesion progression by stimulating these profibrotic functions of smooth-muscle cells (Fig. 291e-2D).

Microvessels As atherosclerotic lesions advance, abundant plexi of microvessels develop in connection with the artery’s vasa vasorum. Newly developing microvascular networks may contribute to lesion complications in several ways. These blood vessels provide an abundant surface area for leukocyte trafficking and may serve as the portal for entry and exit of white blood cells from the established atheroma. Microvessels in the plaques may also furnish foci for intraplaque hemorrhage. Like the neovessels in the diabetic retina, microvessels in the atheroma may be friable and prone to rupture and can produce focal hemorrhage. Such a vascular leak can provoke thrombosis in situ, yielding local thrombin generation, which in turn can activate smooth-muscle and endothelial cells through ligation of protease-activated receptors. Atherosclerotic plaques often contain fibrin and hemosiderin, an indication that episodes of intraplaque hemorrhage contribute to plaque complications.

CALCIFICATION As they advance, atherosclerotic plaques also accumulate calcium. Microvesicles derived from lesional cells can stimulate calcification, and this process co-localizes with regions of heightened inflammation. Mineralization of the atherosclerotic plaque recapitulates many aspects of bone formation, including the regulatory participation of transcription factors such as Runx2.

Plaque Evolution Smooth-muscle cells and macrophages die in the atherosclerotic plaque. Indeed, complex atheromata often have a mostly fibrous character and lack the cellularity of less advanced lesions. This relative paucity of smooth-muscle cells in advanced atheromata may result from the predominance of cytostatic mediators such as TGF- β and IFN- γ (which can inhibit smooth-muscle cell proliferation) and also from smooth-muscle cell apoptosis. Thus, during the evolution of the atherosclerotic plaque, a complex and highly regulated balance between entry and egress of lipoproteins and leukocytes, cell proliferation and cell death, extracellular matrix production, and remodeling, as well as calcification and neovascularization, contribute to lesion formation. Many mediators related to atherogenic risk factors, including those derived from lipoproteins, cigarette smoking, and angiotensin II, provoke the production of proinflammatory cytokines and alter the behavior of the intrinsic vascular wall cells and infiltrating leukocytes that underlie the complex pathogenesis of these lesions. Thus, advances in vascular biology have led to increased

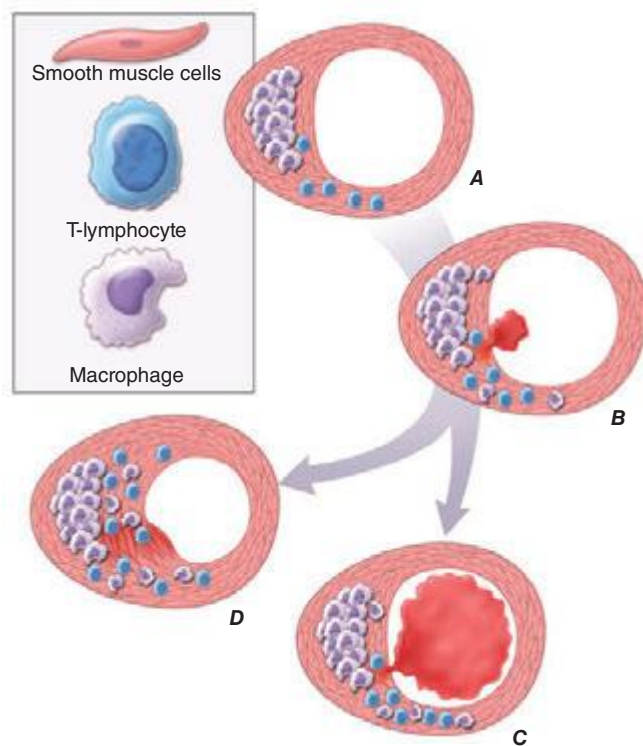


FIGURE 291e-2 Plaque rupture, thrombosis, and healing. **A.** Arterial remodeling during atherogenesis. During the initial part of the life history of an atheroma, growth is often outward, preserving the caliber of the lumen. This phenomenon of “compensatory enlargement” accounts in part for the tendency of coronary arteriography to underestimate the degree of atherosclerosis. **B.** Rupture of the plaque’s fibrous cap causes thrombosis. Physical disruption of the atherosclerotic plaque commonly causes arterial thrombosis by allowing blood coagulant factors to contact thrombogenic collagen found in the arterial extracellular matrix and tissue factor produced by macrophage-derived foam cells in the lipid core of lesions. In this manner, sites of plaque rupture form the nidus for thrombi. The normal artery wall has several fibrinolytic or antithrombotic mechanisms that tend to resist thrombosis and lyse clots that begin to form in situ. Such antithrombotic or thrombolytic molecules include thrombomodulin, tissue- and urokinase-type plasminogen activators, heparan sulfate proteoglycans, prostacyclin, and nitric oxide. **C.** When the clot overwhelms the endogenous fibrinolytic mechanisms, it may propagate and lead to arterial occlusion. The consequences of this occlusion depend on the degree of existing collateral vessels. In a patient with chronic multivessel occlusive coronary artery disease (CAD), collateral channels have often formed. In such circumstances, even a total arterial occlusion may not lead to myocardial infarction (MI), or it may produce an unexpectedly modest or a non-ST-segment elevation infarct because of collateral flow. In a patient with less advanced disease and without substantial stenotic lesions to provide a stimulus for collateral vessel formation, sudden plaque rupture and arterial occlusion commonly produces an ST-segment elevation infarction. These are the types of patients who may present with MI or sudden death as a first manifestation of coronary atherosclerosis. In some cases, the thrombus may lyse or organize into a mural thrombus without occluding the vessel. Such instances may be clinically silent. **D.** The subsequent thrombin-induced fibrosis and healing causes a fibroproliferative response that can lead to a more fibrous lesion that can produce an eccentric plaque that causes a hemodynamically significant stenosis. In this way, a nonocclusive mural thrombus, even if clinically silent or causing unstable angina rather than infarction, can provoke a healing response that can promote lesion fibrosis and luminal encroachment. Such a sequence of events may convert a “vulnerable” atheroma with a thin fibrous cap that is prone to rupture into a more “stable” fibrous plaque with a reinforced cap. Angioplasty of unstable coronary lesions may “stabilize” the lesions by a similar mechanism, producing a wound followed by healing.

291e-4 understanding of the mechanisms that link risk factors to the pathogenesis of atherosclerosis and its complications.

PATHOPHYSIOLOGIC CONSEQUENCES OF ATHEROSCLEROSIS

Atherosclerotic lesions occur ubiquitously in Western societies, and the prevalence of this disease is on the rise globally. Most atheromata produce no symptoms, and many never cause clinical manifestations. Numerous patients with diffuse atherosclerosis may succumb to unrelated illnesses without ever having experienced a clinically significant manifestation of atherosclerosis. Arterial remodeling during atheroma formation accounts for some of this variability in the clinical expression of atherosclerotic disease (Fig. 291e-2A). During the initial phases of atheroma development, the plaque usually grows outward, in an abluminal direction. Vessels affected by atherogenesis tend to increase in diameter, a phenomenon known as *compensatory enlargement*, a type of vascular remodeling. The growing atheroma does not encroach on the arterial lumen until the burden of atherosclerotic plaque exceeds ~40% of the area encompassed by the internal elastic lamina. Thus, during much of its life history, an atheroma will not cause stenosis that can limit tissue perfusion.

Flow-limiting stenoses commonly form later in the history of the plaque. Many such plaques cause stable syndromes such as demand-induced angina pectoris or intermittent claudication in the extremities. In the coronary circulation and other circulations, even total vascular occlusion by an atheroma does not invariably lead to infarction. The hypoxic stimulus of repeated bouts of ischemia characteristically induces formation of collateral vessels in the myocardium, mitigating the consequences of an acute occlusion of an epicardial coronary artery. By contrast, many lesions that cause acute or unstable atherosclerotic syndromes, particularly in the coronary circulation, may arise from atherosclerotic plaques that do not produce a flow-limiting stenosis. Such lesions may produce only minimal luminal irregularities on traditional angiograms and often do not meet the traditional criteria for “significance” by arteriography. Thrombi arising from such nonocclusive stenoses may explain the frequency of MI as an initial manifestation of coronary artery disease (CAD) (in at least one-third of cases) in patients who report no prior history of angina pectoris, a syndrome usually caused by flow-limiting stenoses.

Plaque Instability and Rupture Postmortem studies afford considerable insight into the microanatomic substrate underlying the “instability” of plaques that do not cause critical stenoses. A superficial erosion of the endothelium or a frank plaque rupture or fissure usually produces the thrombus that causes episodes of unstable angina pectoris or the occlusive and relatively persistent thrombus that causes acute MI (Fig. 291e-2B). Rupture of the plaque’s fibrous cap (Fig. 291e-2C) permits contact between coagulation factors in the blood and highly thrombogenic tissue factor expressed by macrophage foam cells in the plaque’s lipid-rich core. If the ensuing thrombus is nonocclusive or transient, the episode of plaque disruption may not cause symptoms or may result in episodic ischemic symptoms such as rest angina. Occlusive thrombi that endure often cause acute MI, particularly in the absence of a well-developed collateral circulation that supplies the affected territory. Repetitive episodes of plaque disruption and healing provide one likely mechanism of transition of the fatty streak to a more complex fibrous lesion (Fig. 291e-2D). The healing process in arteries, as in skin wounds, involves the laying down of new extracellular matrix and fibrosis.

Not all atheromata exhibit the same propensity to rupture. Pathologic studies of culprit lesions that have caused acute MI reveal several characteristic features. Plaques that have caused thromboses tend to have thin fibrous caps, relatively large lipid cores, a high content of macrophages, outward remodeling, and spotty (rather than dense) calcification. Morphometric studies of such culprit lesions show that at sites of plaque rupture, macrophages and T lymphocytes predominate and contain relatively few smooth-muscle cells. The cells that concentrate at sites of plaque rupture bear markers of inflammatory activation. In addition, patients with active atherosclerosis and

acute coronary syndromes display signs of disseminated inflammation. Inflammatory mediators regulate processes that govern the integrity of the plaque’s fibrous cap and, hence, its propensity to rupture. For example, the T cell–derived cytokine IFN- γ , which is found in atherosclerotic plaques, can inhibit growth and collagen synthesis of smooth-muscle cells, as noted above. Cytokines derived from activated macrophages and lesional T cells can boost production of proteolytic enzymes that can degrade the extracellular matrix of the plaque’s fibrous cap. Thus, inflammatory mediators can impair the collagen synthesis required for maintenance and repair of the fibrous cap and trigger degradation of extracellular matrix macromolecules, processes that weaken the plaque’s fibrous cap and enhance its susceptibility to rupture (so-called vulnerable plaques, Fig. 291e-3). In contrast to plaques with these features of vulnerability, those with a dense extracellular matrix and relatively thick fibrous cap without substantial tissue factor–rich lipid cores seem generally resistant to rupture and unlikely to provoke thrombosis.

Functional features of the atheromatous plaque, in addition to its degree of luminal encroachment, influence the clinical manifestations of this disease. This enhanced understanding of plaque biology provides insight into the diverse ways in which atherosclerosis can present clinically and the reasons why the disease may remain silent or stable for prolonged periods, punctuated by acute complications at certain times. Increased understanding of atherogenesis provides new insight into the mechanisms linking it to the risk factors discussed below, indicates the ways in which current therapies may improve outcomes, and suggests new targets for future intervention.

PREVENTION AND TREATMENT

THE CONCEPT OF ATHEROSCLEROTIC RISK FACTORS

The systematic study of risk factors for atherosclerosis emerged from a coalescence of experimental results, as well as from cross-sectional and ultimately longitudinal studies in humans. The prospective, community-based Framingham Heart Study provided rigorous support for the concept that hypercholesterolemia, hypertension, and other factors correlate with cardiovascular risk. Similar observational studies performed worldwide bolstered the concept of “risk factors” for cardiovascular disease.

From a practical viewpoint, the cardiovascular risk factors that have emerged from such studies fall into two categories: those modifiable by lifestyle and/or pharmacotherapy, and those that are immutable, such as age and sex. The weight of evidence supporting various risk factors differs. For example, hypercholesterolemia and hypertension certainly predict coronary risk, but the magnitude of the contributions of other so-called nontraditional risk factors, such as levels of homocysteine, levels of lipoprotein (a) [Lp(a)], and infection, remains controversial. Moreover, some biomarkers that predict cardiovascular risk may not participate in the causal pathway for the disease or its complications. Genetic studies using genome-wide association (GWAS) approaches and Mendelian randomization approaches have helped to distinguish between risk markers and factors that contribute causally to the disease. For example, recent genetic studies suggest that C-reactive protein (CRP) does not itself mediate atherogenesis, despite its ability to predict risk, whereas Lp(a) and apolipoprotein C3 have emerged as a causal risk factor. Table 291e-1 lists a number of risk factors implicated in atherosclerosis. The sections below will consider some of these factors and approaches to their modification.

Lipid Disorders

Abnormalities in plasma lipoproteins and derangements in lipid metabolism rank among the most firmly established and best understood risk factors for atherosclerosis. Chapter 421 describes the lipoprotein classes and provides a detailed discussion of lipoprotein metabolism.

The American College of Cardiology and American Heart Association (ACC/AHA) promulgated new guidelines on risk assessment, lifestyle measures, and cholesterol management in 2013. The panels that produced these guidelines followed an evidence-based approach.

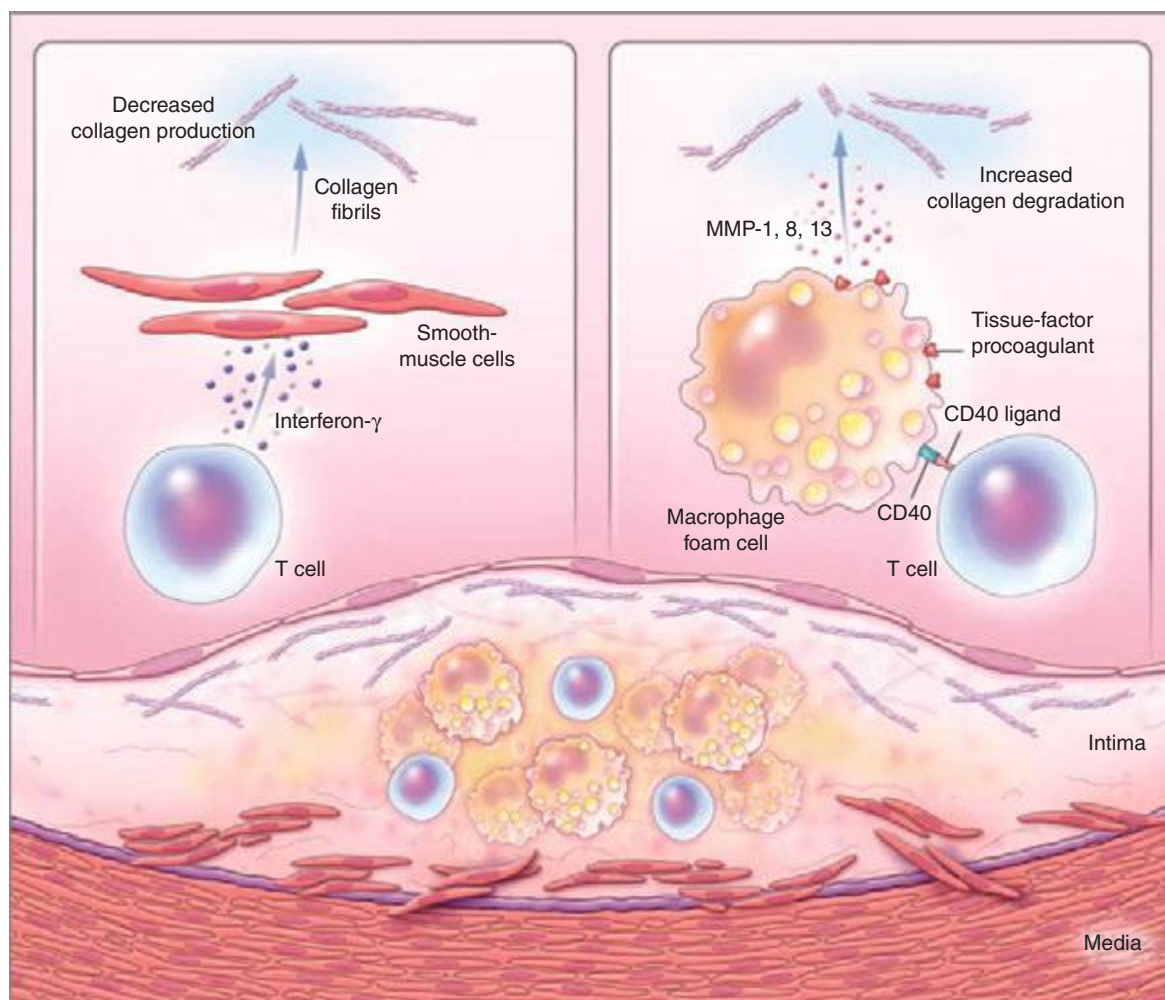


FIGURE 291e-3 Inflammatory pathways that predispose atherosclerotic plaques to rupture and provoke thrombosis. A cross-section of an atheromatous plaque at the bottom of the figure shows the central lipid core that contains macrophage foam cells (yellow) and T cells (blue). The intima and media also contain arterial smooth-muscle cells (red), which are the source of arterial collagen (depicted as triple helical coiled structures). Activated T cells (of the type 1 helper T cell subtype) secrete cytokine interferon γ , which inhibits the production of the new, interstitial collagen that is required to repair and maintain the plaque's protective fibrous cap (upper left). The T cells can also activate the macrophages in the intimal lesion by expressing the inflammatory mediator CD40 ligand (CD154), which engages its cognate receptor (CD40) on the phagocyte. This inflammatory signalling causes overproduction of interstitial collagenases (matrix metalloproteinases [MMPs] 1, 8, and 13) that catalyze the initial rate-limiting step in collagen breakdown (top right). CD40 ligation also causes macrophages to overproduce tissue-factor procoagulant. Thus, inflammatory signalling puts the collagen in the plaque's fibrous cap in double jeopardy—decreasing synthesis and increasing breakdown—rendering the cap susceptible to rupture. Inflammatory activation also boosts tissue-factor production, which triggers thrombus formation in the disrupted plaque. These mechanisms link inflammation in the plaque to the thrombotic complications of atherosclerosis, including the acute coronary syndromes. (Adapted from P Libby: *N Engl J Med* 368:2004, 2013.)

The 2013 cholesterol guideline focused on 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase inhibitors (statins) rather than other classes of lipid-modulating drugs, including fibric acid derivatives, cholesterol absorption inhibitors such as ezetimibe, and niacin products. The guideline cites the lack of contemporary randomized clinical trial evidence that supports the efficacy of these nonstatin lipid-modifying agents in cardiovascular event reduction. The cholesterol guideline defined four statin benefit groups (Table 291e-2): (1) all individuals who have clinical atherosclerotic cardiovascular disease (ASCVD), therefore considered “secondary prevention”; (2) those with LDL cholesterol ≥ 190 mg/dL without a secondary cause such as a high intake of saturated or *trans* fats, various drugs, or certain diseases; (3) individuals with diabetes without established cardiovascular disease who are 40–75 years old and have LDL cholesterol of 70–189 mg/dL; and (4) those without established ASCVD without diabetes who are 40–75 years old and who have LDL cholesterol of 70–189 mg/dL and a calculated ASCVD risk $\geq 7.5\%$. An online risk calculator based on pooled cohorts was provided to aid clinicians and patients in calculating their risk (<http://my.americanheart.org/professional/StatementsGuidelines/PreventionGuidelines/Prevention->

[Guidelines_UCM_457698_SubHomePage.jsp](http://my.americanheart.org/professional/StatementsGuidelines/PreventionGuidelines/Prevention-Guidelines_UCM_457698_SubHomePage.jsp)). Other validated risk calculators that incorporate family history of CAD and a marker of inflammation (high-sensitivity CRP [hsCRP]) that apply to U.S. women and men exist (<http://www.reynoldsriskscore.org>). Downloadable applications for risk calculation on handheld devices are readily available.

The 2013 guideline emphasized a patient-centered approach and recommended that clinicians and patients engage in a risk-benefit conversation before starting statin therapy and not rely solely on calculated risks or arbitrary category assignment. It further emphasizes that medications do not supplant a healthy lifestyle. The guideline also provides some practical suggestions regarding management of muscle symptoms attributed to statins, an issue of considerable concern to many patients and practitioners alike.

In a major departure from prior guidelines, the 2013 guideline eliminates LDL targets as goals of therapy. The panel did so because major clinical trials did not titrate therapy to a goal, but rather used fixed doses of statins. Instead, the new guideline suggests different intensities of statin therapy based on risk category (Fig. 291e-4).

The 2013 guideline focus on statins reflects an extensive body of rigorous evidence that supports the effectiveness of this class of drugs

TABLE 291e-1 MAJOR RISK FACTORS FOR ATHEROSCLEROSIS

High LDL cholesterol
Cigarette smoking
Hypertension (BP \geq 140/90 mmHg or on antihypertensive medication)
Low HDL cholesterol ^a (<1.0 mmol/L [$<$ 40 mg/dL])
Diabetes mellitus
Family history of premature CHD
Age (men \geq 45 years; women \geq 55 years)
Lifestyle risk factors
Obesity (BMI \geq 30 kg/m ²)
Physical inactivity
Atherogenic diet
Emerging risk factors
Lipoprotein (a)
Prothrombotic factors
Proinflammatory factors
Impaired fasting glucose
Subclinical atherosclerosis

^aHDL cholesterol \geq 1.6 mmol/L (\geq 60 mg/dL) has been viewed as a “negative” risk factor.

Abbreviations: BMI, body mass index; BP, blood pressure; CHD, coronary heart disease; HDL, high-density lipoprotein; LDL, low-density lipoprotein.

in cardiovascular event reduction and an acceptable risk-benefit relationship (Fig. 291e-5). Moreover, because almost all statins are now available as generic statins medications, cost has become much less of an impediment to their use.

The clinical use of effective pharmacologic strategies for lowering LDL has reduced cardiovascular events markedly, but a considerable burden of residual risk remains even in patients treated with high-intensity statins. Hence, current studies are evaluating other avenues to address the residual burden of cardiovascular disease that persists despite statin treatment. Inhibitors of genetic studies identified proprotein convertase subtilisin kexin-like 9 (PCSK9) as a regulator of LDL levels associated with cardiovascular outcomes. Interaction of the LDL receptor with PCSK9 hastens the receptor’s degradation, and hence yields higher circulating LDL concentrations. Genetic variants that lower PCSK9 activity appear to protect against cardiovascular events. Monoclonal antibodies that neutralize PCSK9 lower LDL levels even in statin-treated patients and are currently under investigation as novel therapeutics to lower cardiovascular risk.

LDL-lowering therapies do not appear to exert their beneficial effect on cardiovascular events by causing a marked “regression” of stenoses. Studies of lipid lowering monitored by angiography or by intravascular imaging modalities have shown at best a modest reduction in coronary artery stenoses over the duration of study, despite abundant evidence of event reduction. These results suggest that the beneficial mechanism of lipid lowering by statins does not require a substantial reduction in the fixed stenoses. Rather, the benefit may derive from “stabilization” of atherosclerotic lesions without substantially decreased stenosis.

TABLE 291e-2 SUMMARY OF THE FOUR STATIN BENEFIT GROUPS DESCRIBED IN THE 2013 ACC/AHA GUIDELINE ON THE TREATMENT OF BLOOD CHOLESTEROL TO REDUCE ATHEROSCLEROTIC CARDIOVASCULAR RISK IN ADULTS

- Clinical ASCVD “secondary prevention”
- LDL-C \geq 190 mg/dL without secondary cause (e.g., saturated/trans fats, drugs, certain diseases)
- Primary prevention *with* diabetes mellitus: age 40–75 years, LDL-C 70–189 mg/dL
- Primary prevention *without* diabetes mellitus: age 40–75 years, LDL-C 70–189 mg/dL, estimated ASCVD risk \geq 7.5%

Abbreviations: ACC/AHA, American College of Cardiology and American Heart Association; ASCVD, atherosclerotic cardiovascular disease; LDL-C, low-density lipoprotein cholesterol.

Source: Adapted from NJ Stone et al: 2013 ACC/AHA Guideline on the Treatment of Blood Cholesterol to Reduce Atherosclerotic Cardiovascular Risk in Adults. *J Am Coll Cardiol* 2013, doi: 10.1016/j.jacc.2013.11.002.

Such stabilization of atherosclerotic lesions and the attendant decrease in coronary events may result from the egress of lipids or from favorably influencing aspects of the biology of atherogenesis discussed above. In addition, as sizable lesions may protrude abuminally rather than into the lumen due to complementary enlargement, shrinkage of such plaques may not be apparent on angiograms. The consistent benefit of statins may depend not only on their salutary effects on the lipid profile, but also on direct modulation of plaque biology independent of lipid lowering.

As the prevalence of metabolic syndrome and diabetes increases, many patients present with low concentrations of HDL (HDL cholesterol $<$ 1.0 mmol/L [$<$ 40 mg/dL]). A baseline measurement of HDL cholesterol indubitably correlates with future cardiovascular risk. Yet, the utility of therapies that raise HDL cholesterol levels in blood as effective interventions to reduce cardiovascular vascular events has come into question. Blood HDL levels vary inversely with those of triglycerides, and the independent role of HDL versus triglycerides as a cardiovascular risk factor remains unsettled. The 2013 guideline does not advocate any specific therapy for raising HDL. Indeed, multiple recent trials failed to show that raising HDL cholesterol levels improves cardiovascular outcomes, and recent genetic studies cast doubt on low HDL as a causal risk factor for atherosclerotic events. Weight loss and physical activity can raise HDL, and these lifestyle measures merit universal adoption (Table 291e-3). Nicotinic acid, particularly in combination with statins, can robustly raise HDL, but clinical trial data do not support the effectiveness of nicotinic acid in cardiovascular risk reduction. Agonists of nuclear receptors provide another potential avenue for raising HDL levels. Yet patients treated with peroxisome proliferator-activated receptors alpha and gamma (PPAR- α and - γ) agonists have not consistently shown improved cardiovascular outcomes, and at least some PPAR agonists have been associated with worsened cardiovascular outcomes. Other agents in clinical development raise HDL levels by inhibiting cholesteryl ester transfer protein (CETP). Two such agents have undergone large-scale clinical evaluation and have not shown efficacy in improving cardiovascular outcomes. Clinical studies currently under way will assess the effectiveness of two other CETP inhibitors that lack some of the adverse off-target actions encountered with the first agent tested.

The mechanism by which elevated LDL levels promote atherogenesis may involve oxidative modification. Yet, rigorous and well-controlled clinical trials have failed to demonstrate that antioxidant vitamin therapy improves coronary heart disease (CHD) outcomes. In regard to nontraditional risk factors including homocysteine and infection, large-scale clinical trials using vitamins to lower homocysteine or using antibiotics have not reduced cardiovascular events. Therefore, the current evidence base does *not* support the use of vitamins or antibiotics to lower cardiovascular risk.

Hypertension (See also Chap. 298) A wealth of epidemiologic data support a relationship between hypertension and atherosclerotic risk, and extensive clinical trial evidence has established that pharmacologic treatment of hypertension can reduce the risk of stroke, heart failure, and CHD events.

Diabetes Mellitus, Insulin Resistance, and the Metabolic Syndrome (See also Chap. 417) Most patients with diabetes mellitus die of atherosclerosis and its complications. Aging and rampant obesity underlie a current epidemic of type 2 diabetes mellitus. The abnormal lipoprotein profile associated with insulin resistance, known as *diabetic dyslipidemia*, accounts for part of the elevated cardiovascular risk in patients with type 2 diabetes. Although diabetic individuals often have LDL cholesterol levels near the average, the LDL particles tend to be smaller and denser and, therefore, more atherogenic. Other features of diabetic dyslipidemia include low HDL and elevated triglyceride levels. Hypertension also frequently accompanies obesity, insulin resistance, and dyslipidemia. This commonly encountered clinical cluster of risk factors has become known as the *metabolic syndrome* (Chap. 422). Despite legitimate concerns about whether clustered components confer more risk than the individual components, the metabolic syndrome concept may offer clinical utility.

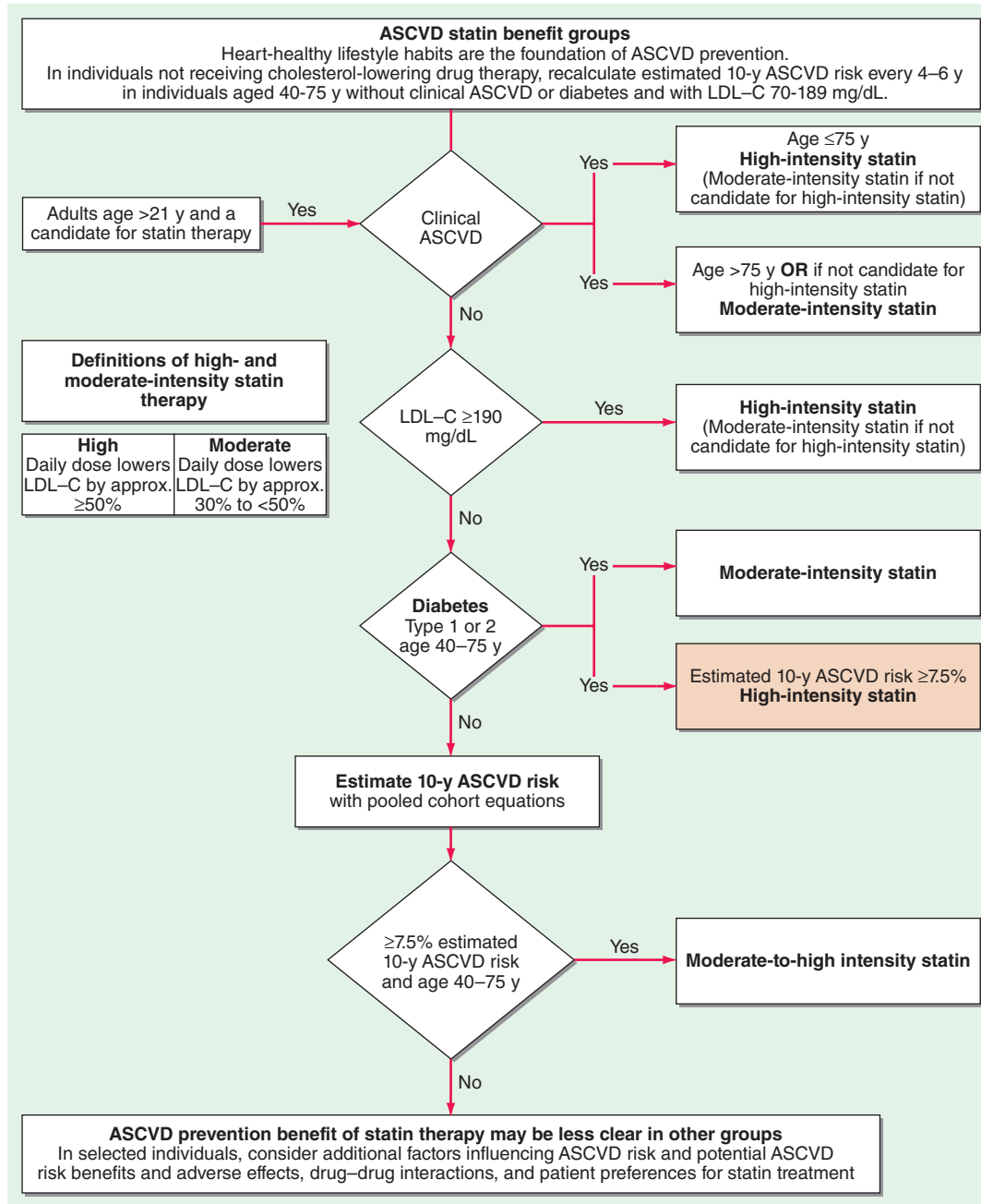


FIGURE 291e-4 Major recommendations for statin therapy for atherosclerotic cardiovascular disease (ASCVD) prevention. LDL-C, low-density lipoprotein cholesterol. (From NJ Stone et al: *J Am Coll Cardiol*, 2013, doi: 10.1016/j.jacc.2013.11.002.)

Therapeutic objectives for intervention in these patients include addressing the underlying causes, including obesity and low physical activity, by initiating lifestyle measures (see below). Establishing that strict glycemic control reduces the risk of macrovascular complications of diabetes has proved much more elusive than the beneficial effects on microvascular complications such as retinopathy and renal disease. Indeed, “tight” glycemic control may increase adverse events in patients with type 2 diabetes, lending even greater importance to aggressive control of other aspects of risk in this patient population. In this regard, multiple clinical trials have demonstrated unequivocal benefit of statin therapy in diabetic patients over all ranges of LDL cholesterol levels (but not those with end-stage renal disease or advanced heart failure). Among the oral hypoglycemic agents, metformin possesses the best evidence base for cardiovascular event reduction. The novel oral hypoglycemic agents tested in sufficiently powered trials, the dipeptidyl peptidase-4 (DPP-4) inhibitors saxagliptin and alogliptin, did not show cardiovascular benefit. Indeed, saxagliptin was associated

with a slight increase in heart failure. Diabetic populations appear to derive particular benefit from antihypertensive strategies that block the action of angiotensin II. Thus, the antihypertensive regimen for patients with the metabolic syndrome should include angiotensin-converting enzyme inhibitors or angiotensin receptor blockers when possible. Many of these individuals will require more than one antihypertensive agent to reach the 2013 goals for individuals 18 years of age or older with diabetes to achieve a systolic blood pressure of less than 140 mmHg and a diastolic blood pressure of less than 90 mmHg.

Male Sex/Postmenopausal State Decades of observational studies have verified excess coronary risk in men compared with premenopausal women. After menopause, however, coronary risk accelerates in women. Although observational and experimental studies have suggested that estrogen therapy reduces coronary risk, large-scale randomized clinical trials have not demonstrated a net benefit of estrogen with or without progestins on CHD outcomes. In the Heart

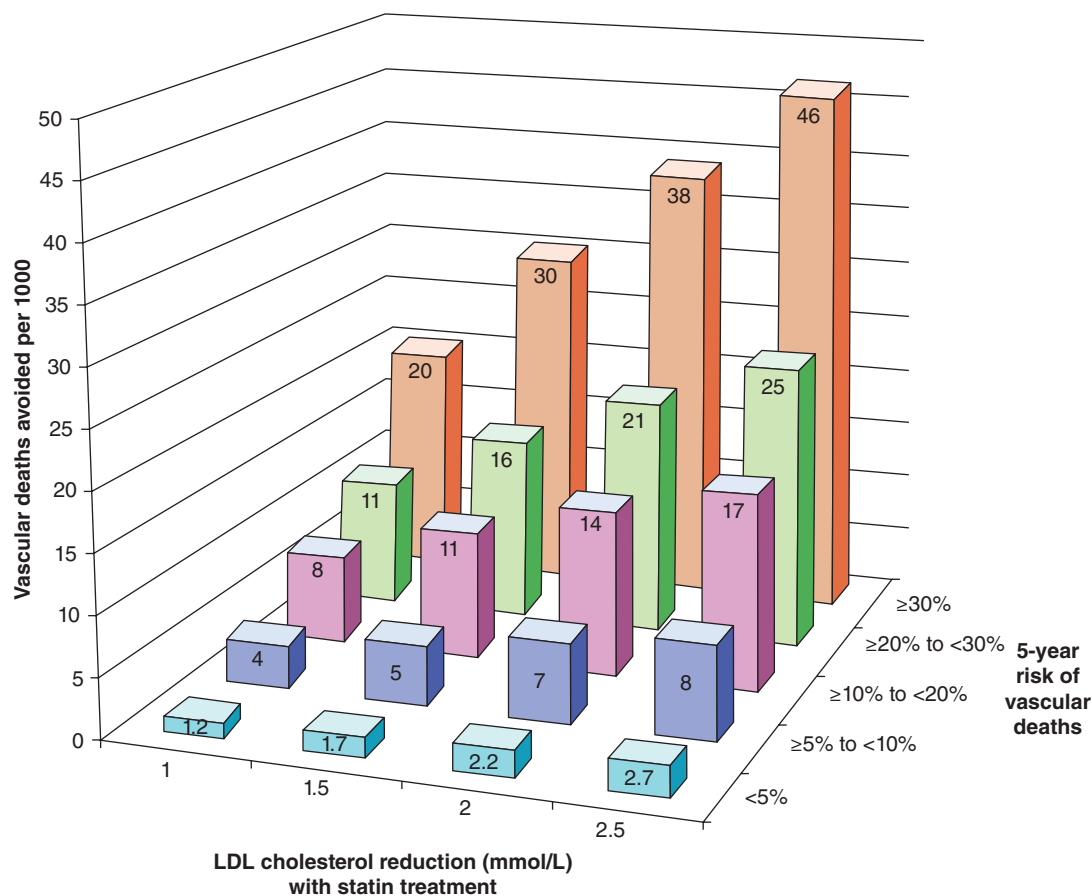


FIGURE 291e-5 The Cholesterol Treatment Trialists Collaboration meta-analyzed 27 randomized clinical trials evaluating statin therapy. They found profound decreases in both major vascular events and vascular death (not shown) proportional to the magnitude of low-density lipoprotein (LDL) cholesterol reduction achieved with statin treatment. This diagram shows the results of this meta-analysis for vascular death. (From *Lancet* 380:581, 2012.)

and Estrogen/Progestin Replacement Study (HERS), postmenopausal female survivors of acute MI were randomized to an estrogen/progestin combination or to placebo. This study showed no overall reduction in recurrent coronary events in the active treatment arm. Indeed, early in the 5-year course of this trial, a trend occurred toward an increase

in vascular events in the treated women. Extended follow-up of this cohort did not disclose an accrual of benefit in the treatment group. The Women's Health Initiative (WHI) study arm, using a similar estrogen plus progesterone regimen, was halted due to a small but significant hazard of cardiovascular events, stroke, and breast cancer. The estrogen without progestin arm of WHI (conducted in women without a uterus) was stopped early due to an increase in strokes, and failed to afford protection from MI or CHD death during observation over 7 years. The excess cardiovascular events in these trials may result from an increase in thromboembolism (Chap. 413). Physicians should work with women to provide information and help weigh the small but evident CHD risk of estrogen ± progestin versus the benefits for postmenopausal symptoms and osteoporosis, taking personal preferences into account. Post hoc analyses of observational studies suggest that estrogen therapy in women younger than or closer to menopause than the women enrolled in WHI might confer cardiovascular benefit. Thus, the timing in relation to menopause or the age at which estrogen therapy begins may influence its risk/benefit balance.

The lack of efficacy of estrogen therapy in cardiovascular risk reduction highlights the need for redoubled attention to known modifiable risk factors in women. Meta-analysis supports the efficacy of statins to reduce cardiovascular events in women in primary prevention, as well as in those who have already experienced a cardiovascular event.

TABLE 291e-3 HEART HEALTHY NUTRITION AND PHYSICAL ACTIVITY BEHAVIORS RECOMMENDED IN THE 2013 ACC/AHA GUIDELINE ON LIFESTYLE MANAGEMENT TO REDUCE CARDIOVASCULAR RISK

The adult population should be encouraged to practice heart healthy lifestyle behaviors, including:

- Consume a dietary pattern that emphasizes intake of vegetables, fruits, and whole grains; include low-fat dairy products, poultry, fish, legumes, nontropical vegetable oils, and nuts; and *limit* intake of sodium, sweets, sugar-sweetened beverages, and red meats.
- Adapt this dietary pattern to appropriate calorie requirements, personal and cultural food preferences, and nutrition therapy for other medical conditions (including diabetes mellitus).
- Achieve this pattern by following plans such as the DASH dietary pattern, the USDA Food Pattern, or the AHA Diet.
- Engage in 2 h and 30 min a week of moderate-intensity or 1 h and 15 min (75 min) a week of vigorous-intensity aerobic physical activity, or an equivalent combination of moderate- and vigorous-intensity aerobic physical activity. Aerobic activity should be performed in episodes of at least 10 min, preferably spread throughout the week.
- Achieve and maintain a healthy weight. Refer to the 2013 Obesity Expert Panel Report for recommendations on weight loss and maintenance.

Abbreviations: ACC/AHA, American College of Cardiology and American Heart Association; DASH, Dietary Approaches to Stop Hypertension; USDA, U.S. Department of Agriculture.

Source: Adapted from RH Eckel et al: 2013 AHA/ACC Guideline on Lifestyle Management to Reduce Cardiovascular Risk. *J Am Coll Cardiol* 2013, doi: 10.1016/j.jacc.2013.11.003.

Dysregulated Coagulation or Fibrinolysis Thrombosis ultimately causes the gravest complications of atherosclerosis. The propensity to form thrombi and/or lyse clots once they form influences the manifestations of atherosclerosis. Thrombosis provoked by atheroma rupture and subsequent healing may promote plaque growth. Certain individual characteristics can influence thrombosis or fibrinolysis and have received attention as potential coronary risk factors. For example,

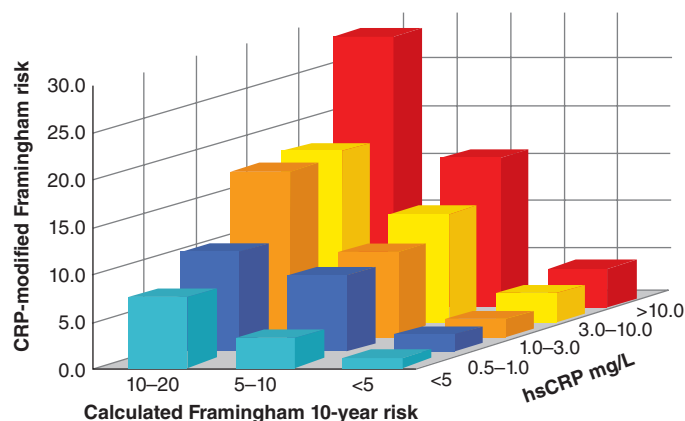


FIGURE 291e-6 C-reactive protein (CRP) level adds to the predictive value of the Framingham score. hsCRP, high-sensitivity measurement of CRP. (Adapted from PM Ridker et al: *Circulation* 109:2818, 2004.)

fibrinogen levels correlate with coronary risk and provide information about coronary risk independent of the lipoprotein profile.

The stability of an arterial thrombus depends on the balance between fibrinolytic factors, such as plasmin, and inhibitors of the fibrinolytic system, such as plasminogen activator inhibitor 1 (PAI-1). Individuals with diabetes mellitus or the metabolic syndrome have elevated levels of PAI-1 in plasma, and this probably contributes to the increased risk of thrombotic events. Lp(a) (Chap. 421) may modulate fibrinolysis, and individuals with elevated Lp(a) levels have increased CHD risk.

Aspirin reduces CHD events in several contexts. Chapter 293 discusses aspirin therapy in stable ischemic heart disease, Chap. 294 reviews recommendations for aspirin treatment in acute coronary syndromes, and Chap. 446 describes aspirin's role in preventing recurrent ischemic stroke. In primary prevention, pooled trial data show that low-dose aspirin treatment (81 mg/d to 325 mg on alternate days) can reduce the risk of a first MI in men. Although the Women's Health Study (WHS) showed that aspirin (100 mg on alternate days) reduced strokes by 17%, it did not prevent MI in women. Current AHA guidelines recommend the use of low-dose aspirin (75–160 mg/d) for women with high cardiovascular risk ($\geq 20\%$ 10-year risk), for men with a $\geq 10\%$ 10-year risk of CHD, and for all aspirin-tolerant patients with established cardiovascular disease who lack contraindications.

Inflammation An accumulation of clinical evidence shows that markers of inflammation correlate with coronary risk. For example, plasma levels of CRP, as measured by a high-sensitivity assay (hsCRP), prospectively predict the risk of MI. CRP levels also correlate with the outcome in patients with acute coronary syndromes. In contrast to several other novel risk factors, CRP adds predictive information to that derived from established risk factors, such as those included in the Framingham score (Fig. 291e-6). Mendelian randomization studies do not support a causal role for CRP in cardiovascular disease. Thus, CRP serves as a validated biomarker of risk, but probably not as a direct contributor to pathogenesis.

Elevations in acute-phase reactants such as fibrinogen and CRP reflect the overall inflammatory burden, not just vascular foci of inflammation. Visceral adipose tissue releases proinflammatory cytokines that drive CRP production and may represent a major extravascular

stimulus to the elevation of inflammatory markers in obese and overweight individuals. Indeed, CRP levels rise with body mass index (BMI) or visceral adipose depot as assessed by imaging, and weight reduction lowers CRP levels. Infectious agents might also furnish inflammatory stimuli related to cardiovascular risk.

Statin therapy likely reduces cardiovascular events in part by muting the inflammatory aspects of the pathogenesis of atherosclerosis. For example, in statin trials conducted in both primary (JUPITER) and secondary (PROVE-IT/TIMI-22) prevention populations, prespecified analyses showed that those who achieved lower levels of both LDL and CRP had better clinical outcomes than did those who only reached the lower level of either the inflammatory marker or the atherogenic lipoprotein (Fig. 291e-7). The anti-inflammatory effect of statins appears independent of LDL lowering, because these two variables correlated very poorly in individual subjects in multiple clinical trials.

Lifestyle Modification The prevention of atherosclerosis presents a long-term challenge to all health care professionals and for public health policy. Both individual practitioners and organizations providing health care should strive to help patients optimize their risk factor profiles long before atherosclerotic disease becomes manifest. The current accumulation of cardiovascular risk in youth and in certain minority populations presents a particularly vexing concern from a public health perspective.

The ACC/AHA 2013 Guideline on Lifestyle Management to Reduce Cardiovascular Risk relied on rigorous evidentiary reviews. Few lifestyle interventions have undergone rigorous evaluation in randomized clinical trials. Therefore, these guidelines reflected judicious analysis of carefully selected observational studies and of intervention studies that relied primarily on biomarkers or surrogate endpoints rather than “hard” cardiovascular outcomes. Table 291e-3 summarizes the ACC/AHA lifestyle recommendations.

The care plan for all patients seen by internists should include measures to assess and minimize cardiovascular risk. Physicians must counsel patients about the health risks of tobacco use and provide guidance and resources regarding smoking cessation. Similarly, physicians should advise all patients about prudent dietary and physical activity habits for maintaining ideal body weight. Both National Institutes of Health (NIH) and AHA statements recommend at least 30 min of moderate-intensity physical activity per day. Obesity, particularly the male pattern of centripetal or visceral fat accumulation, can contribute to the elements of the “metabolic syndrome” cluster. Physicians should encourage their patients to take personal responsibility for behavior related to modifiable risk factors for the development of premature atherosclerotic disease. Conscientious counseling and patient education may forestall the need for pharmacologic measures intended to reduce coronary risk.

Issues in Risk Assessment A growing panel of markers of coronary risk presents a perplexing array to the practitioner. Markers measured in peripheral blood include size fractions of LDL particles and concentrations of homocysteine, Lp(a), fibrinogen, CRP, PAI-1, myeloperoxidase, lipoprotein-associated phospholipase A₂, and imaging assessment of subclinical atherosclerosis, among many others. In general, such specialized tests add little to the information available from a careful history and physical examination combined with measurement of a plasma lipoprotein profile and fasting blood glucose. The hsCRP measurement may well prove an exception in view of its robustness in risk

Group	N	Rate
Placebo	7832	1.11
LDL ≥ 70 mg/dL, hsCRP ≥ 2 mg/L	1384	1.11
LDL < 70 mg/dL, hsCRP ≥ 2 mg/L	2921	0.62
LDL ≥ 70 mg/dL, hsCRP < 2 mg/L	726	0.54
LDL < 70 mg/dL, hsCRP < 2 mg/L	2685	0.38

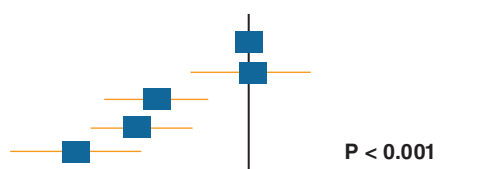


FIGURE 291e-7 Evidence from the JUPITER study that both low-density lipoprotein (LDL)-lowering and anti-inflammatory actions contribute to the benefit of statin therapy in primary prevention. See text for explanation. hsCRP, high-sensitivity measurement of C-reactive protein (CRP). (Adapted from PM Ridker et al: *Lancet* 373:1175, 2009.)

291e-10 prediction, ease of reproducible and standardized measurement, relative stability in individuals over time, ability to add to the risk information disclosed by standard measurements such as the components of the Framingham risk score, and most importantly, the demonstration in a large-scale trial (JUPITER) that allocating therapy can reduce cardiovascular events in those deemed ineligible by traditional risk assessment criteria. The addition of information regarding a family history of premature atherosclerosis (a simply obtained indicator of genetic susceptibility), together with the inflammation marker hsCRP, permits correct reclassification of risk in individuals, especially those whose Framingham scores place them at intermediate risk.

Available data do not support the routine use of imaging studies to screen for subclinical disease (e.g., measurement of carotid intima-media thickness, coronary artery calcification, and use of computed tomographic coronary angiograms [CTA]). Inappropriate use of such imaging modalities may promote excessive alarm in asymptomatic individuals and prompt invasive diagnostic and therapeutic procedures of unproven value for both asymptomatic atherosclerosis and incidental findings. Widespread application of such modalities for screening should await proof that targeting therapies based on their application provides clinical benefit.

The 2013 ACC/AHA Guideline on the Assessment of Cardiovascular Risk recommends the use of newer risk markers if uncertainty persists after assessing quantitative risk using the pooled cohort calculator. The guideline states that family history, hsCRP, coronary artery calcium (CAC) score, or ankle-brachial index (ABI) may then be considered to inform treatment decision making. It discourages carotid intima-media thickness (CIMT) for routine measurement in clinical practice for risk assessment for a first ASCVD event. The guideline panel deemed the contribution to risk assessment for a first ASCVD event using apolipoprotein B (ApoB), chronic kidney disease, albuminuria, or cardiorespiratory fitness as uncertain at present.

Progress in human genetics holds considerable promise for risk prediction and for individualization of cardiovascular therapy. Many early reports identified single-nucleotide polymorphisms (SNPs) in candidate genes as predictors of cardiovascular risk. The validation of such genetic markers of risk and drug responsiveness in multiple populations often proved disappointing. The era of GWAS has led to discovery of sites of genetic variation that reproducibly indicate heightened cardiovascular risk (e.g., chromosome 9p21). The advent of technology that permits relatively rapid and inexpensive exome or whole-genome sequencing promises to identify new therapeutic targets, sharpen risk prediction, and deploy preventive or therapeutic measures in a more personalized manner. Despite this considerable promise, genetic scores for risk prediction have not yet demonstrated consistent improvement over algorithms that use traditional tools.

THE CHALLENGE OF IMPLEMENTATION: CHANGING PHYSICIAN AND PATIENT BEHAVIOR

Despite declining age-adjusted rates of coronary death, cardiovascular mortality worldwide is rising due to aging of the population and due to subsiding of communicable diseases and increased prevalence of risk factors in developing countries. Enormous challenges remain regarding translation of the current evidence base into practice. Physicians must learn how to help individuals adopt a healthy lifestyle in a culturally appropriate manner and to deploy their increasingly powerful pharmacologic tools most economically and effectively. The obstacles to implementation of current evidence-based prevention and treatment of atherosclerosis involve economics, education, physician awareness, and patient adherence to recommended regimens. Future goals in the treatment of atherosclerosis should include more widespread implementation of the current evidence-based guidelines regarding risk factor management and, when appropriate, drug therapy.

Knowledge about the biology of human atherosclerosis and the risk factors for the disease has expanded considerably. The application of vascular biology to human atherosclerosis has revealed many new insights into the mechanisms that promote clinical events. The series of animated video presentations presented here illustrates some of the evolving information about risk factors for atherosclerosis and the pathophysiology of clinical events.

The importance of blood pressure as a risk factor for atherosclerosis and cardiovascular events has long been recognized. More recent clinical information has highlighted the importance of pulse pressure—the difference between the systolic pressure and minimum diastolic arterial pressure—as a prognostic indicator of cardiovascular risk. The video clip on pulse pressure explains the pathophysiology of this readily measured clinical variable.

Physicians possess a great deal of knowledge about the role of cholesterol in the prediction of atherosclerosis and its complications, but knowledge about the mechanism that links hypercholesterolemia to cardiovascular events has lagged the epidemiologic and observational findings. Low-density lipoprotein (LDL) provides an example of a well-understood cardiovascular risk factor. Several of the animations included in this series highlight the role of modified LDL as a trigger for inflammation and other aspects of the pathobiology of arterial plaques that lead to their aggravation and clinical events. Physicians have useful tools for modulating LDL, but other aspects of dyslipidemia are on the rise and provide a growing challenge to the practitioner. In particular, low levels of high-density lipoprotein (HDL) and elevated levels of triglycerides characterize the constellation of findings denoted by some as the “metabolic syndrome.” In the wake of increasing obesity worldwide, these features of the lipoprotein profile require renewed focus. Several of the animations in this collection discuss the concept of the metabolic syndrome and the role of lipid profile components other than LDL in atherogenesis.

The traditional approach to atherosclerosis focused on arterial stenoses as a cause of ischemia and cardiovascular events. Physicians now have effective revascularization modalities for addressing flow-limiting stenoses, but atherosclerotic plaques that do not cause stenoses nonetheless may precipitate clinical events, such as unstable angina and acute myocardial infarction. Thus, it is necessary to add to the traditional focus on stenosis an enlarged appreciation of the pathobiology of atherosclerosis that underlies many acute coronary syndromes. The animation on the development and complication of atherosclerotic plaque explains some of these emerging concepts in plaque

activation as they apply to the precipitation of thrombotic complications of atherosclerosis.

ACKNOWLEDGMENTS

From Peter Libby, MD: Changes and Challenges in Cardiovascular Protection: A Special CME Activity for Physicians. Created under an unrestricted educational grant from Merck & Co., Inc. Copyright © 2002, Cardinal Health; used with permission.

VIDEO 292e-1 Pulse pressure. Considerable evidence suggests that pulse pressure serves as an important risk factor for future cardiovascular events. This video clip explains the derivation of pulse pressure and some of the pathophysiology that determines this parameter. *(With permission from the Academy for Health Care Education.)*

VIDEO 292e-2 Plaque instability. Most coronary thromboses result from a physical disruption of the atherosclerotic plaque. This animation explains some of the current concepts of the pathophysiology of atherosclerotic plaque disruption and how it triggers arterial thrombosis.

VIDEO 292e-3 Lipoprotein menagerie. The lipid profile confers important information regarding cardiovascular risk and the effects of therapies; understanding lipoprotein metabolism provides insight into the pathophysiology of arterial disease. This animation presents the rudiments of lipoprotein metabolism that are important in clinical medicine.

VIDEO 292e-4 Formation and complication of atherosclerotic plaques. Physicians now understand the generation of atherosclerotic plaques as a dynamic process involving an interchange between cells of the artery wall, inflammatory cells recruited from blood, and risk factors such as lipoproteins. This animation reviews current thinking about how risk factors alter the biology of the artery wall and can incite initiation and progression of atherosclerosis. It also discusses the importance of inflammation in these processes and portrays the role of inflammation in plaque disruption and thrombosis. Finally, this animation depicts the concept of stabilization of atherosclerotic plaques by interventions such as lipid lowering.

VIDEO 292e-5 Atherogenesis. This video clip highlights some of the current thinking about mechanisms of atherogenesis.

VIDEO 292e-6 Metabolic syndrome. A number of important cardiovascular risk factors tend to cluster in a pattern that has been described by some as the metabolic syndrome. Although controversy persists regarding whether cardiovascular risk due to these factors is additive or synergistic, their clinical importance is growing. This animation discusses some of the metabolic derangements that underlie the metabolic syndrome.

293 Ischemic Heart Disease

Elliott M. Antman, Joseph Loscalzo

Ischemic heart disease (IHD) is a condition in which there is an inadequate supply of blood and oxygen to a portion of the myocardium; it typically occurs when there is an imbalance between myocardial oxygen supply and demand. The most common cause of myocardial ischemia is atherosclerotic disease of an epicardial coronary artery (or arteries) sufficient to cause a regional reduction in myocardial blood flow and inadequate perfusion of the myocardium supplied by the involved coronary artery. **Chapter 291e** deals with the development and treatment of atherosclerosis. This chapter focuses on the chronic manifestations and treatment of IHD. The subsequent chapters address the acute phases of IHD.

EPIDEMIOLOGY AND GLOBAL TRENDS



IHD causes more deaths and disability and incurs greater economic costs than any other illness in the developed world. IHD is the most common, serious, chronic, life-threatening illness in the United States, where 13 million persons have IHD, >6 million have angina pectoris, and >7 million have sustained a myocardial infarction. Genetic factors, a high-fat and energy-rich diet, smoking, and a sedentary lifestyle are associated with the emergence of IHD (**Chap. 291e**). In the United States and Western Europe, IHD is growing among low-income groups, but primary prevention has delayed the disease to later in life in all socioeconomic groups. Despite these sobering statistics, it is worth noting that epidemiologic data show a decline in the rate of deaths due to IHD, about half of which is attributable to treatments and half to prevention by risk factor modification.

Obesity, insulin resistance, and type 2 diabetes mellitus are increasing and are powerful risk factors for IHD. These trends are occurring in the general context of population growth and as a result of the increase in the average age of the world's population. With urbanization in countries with emerging economies and a growing middle class, elements of the energy-rich Western diet are being adopted. As a result, the prevalence of risk factors for IHD and the prevalence of IHD itself are both increasing rapidly, so that in analyses of the global burden of disease, there is a shift from communicable to noncommunicable diseases. Population subgroups that appear to be particularly affected are men in South Asian countries, especially India and the Middle East. In light of the projection of large increases in IHD throughout the world, IHD is likely to become the most common cause of death worldwide by 2020.

PATHOPHYSIOLOGY

Central to an understanding of the pathophysiology of myocardial ischemia is the concept of myocardial supply and demand. In normal conditions, for any given level of a demand for oxygen, the myocardium will control the supply of oxygen-rich blood to prevent underperfusion of myocytes and the subsequent development of ischemia and infarction. The major determinants of myocardial oxygen demand (MVO_2) are heart rate, myocardial contractility, and myocardial wall tension (stress). An adequate supply of oxygen to the myocardium requires a satisfactory level of oxygen-carrying capacity of the blood (determined by the inspired level of oxygen, pulmonary function, and hemoglobin concentration and function) and an adequate level of coronary blood flow. Blood flows through the coronary arteries in a phasic fashion, with the majority occurring during diastole. About 75% of the total coronary resistance to flow occurs across three sets of arteries: (1) large epicardial arteries (R_1), (2) prearteriolar vessels (R_2), and (3) arteriolar and intramyocardial capillary vessels (R_3). In the absence of significant flow-limiting atherosclerotic

obstructions, R_1 is trivial; the major determinant of coronary resistance is found in R_2 and R_3 .

The normal coronary circulation is dominated and controlled by the heart's requirements for oxygen. This need is met by the ability of the coronary vascular bed to vary its resistance (and, therefore, blood flow) considerably while the myocardium extracts a high and relatively fixed percentage of oxygen. Normally, intramyocardial resistance vessels demonstrate a great capacity for dilation (R_2 and R_3 decrease). For example, the changing oxygen needs of the heart with exercise and emotional stress affect coronary vascular resistance and in this manner regulate the supply of oxygen and substrate to the myocardium (*metabolic regulation*). The coronary resistance vessels also adapt to physiologic alterations in blood pressure to maintain coronary blood flow at levels appropriate to myocardial needs (*autoregulation*).

By reducing the lumen of the coronary arteries, atherosclerosis limits appropriate increases in perfusion when the demand for flow is augmented, as occurs during exertion or excitement. When the luminal reduction is severe, myocardial perfusion in the basal state is reduced. Coronary blood flow also can be limited by spasm (see Prinzmetal's angina in [Chap. 294](#)), arterial thrombi, and, rarely, coronary emboli as well as by ostial narrowing due to aortitis. Congenital abnormalities such as the origin of the left anterior descending coronary artery from the pulmonary artery may cause myocardial ischemia and infarction in infancy, but this cause is very rare in adults.

Myocardial ischemia also can occur if myocardial oxygen demands are markedly increased and particularly when coronary blood flow may be limited, as occurs in severe left ventricular hypertrophy due to aortic stenosis. The latter can present with angina that is indistinguishable from that caused by coronary atherosclerosis largely owing to subendocardial ischemia ([Chap. 283](#)). A reduction in the oxygen-carrying capacity of the blood, as in extremely severe anemia or in the presence of carboxyhemoglobin, rarely causes myocardial ischemia by itself but may lower the threshold for ischemia in patients with moderate coronary obstruction.

Not infrequently, two or more causes of ischemia coexist in a patient, such as an increase in oxygen demand due to left ventricular hypertrophy secondary to hypertension and a reduction in oxygen supply secondary to coronary atherosclerosis and anemia. Abnormal constriction or failure of normal dilation of the coronary resistance vessels also can cause ischemia. When it causes angina, this condition is referred to as *microvascular angina*.

CORONARY ATHEROSCLEROSIS

Epicardial coronary arteries are the major site of atherosclerotic disease. The major risk factors for atherosclerosis (high levels of plasma low-density lipoprotein [LDL], low plasma high-density lipoprotein [HDL], cigarette smoking, hypertension, and diabetes mellitus [[Chap. 291e](#)]) disturb the normal functions of the vascular endothelium. These functions include local control of vascular tone, maintenance of an antithrombotic surface, and control of inflammatory cell adhesion and diapedesis. The loss of these defenses leads to inappropriate constriction, luminal thrombus formation, and abnormal interactions between blood cells, especially monocytes and platelets, and the activated vascular endothelium. Functional changes in the vascular milieu ultimately result in the subintimal collections of fat, smooth muscle cells, fibroblasts, and intercellular matrix that define the atherosclerotic plaque. Rather than viewing atherosclerosis strictly as a vascular problem, it is useful to consider it in the context of alterations in the nature of the circulating blood (hyperglycemia; increased concentrations of LDL cholesterol, tissue factor, fibrinogen, von Willebrand factor, coagulation factor VII, and platelet microparticles). The combination of a "vulnerable vessel" in a patient with "vulnerable blood" promotes a state of hypercoagulability and hypofibrinolysis. This is especially true in patients with diabetes mellitus.

Atherosclerosis develops at irregular rates in different segments of the epicardial coronary tree and leads eventually to segmental reductions in cross-sectional area, i.e., plaque formation. There is also a

predilection for atherosclerotic plaques to develop at sites of increased turbulence in coronary flow, such as at branch points in the epicardial arteries. When a stenosis reduces the diameter of an epicardial artery by 50%, there is a limitation of the ability to increase flow to meet increased myocardial demand. When the diameter is reduced by ~80%, blood flow at rest may be reduced, and further minor decreases in the stenotic orifice area can reduce coronary flow dramatically to cause myocardial ischemia at rest or with minimal stress.

Segmental atherosclerotic narrowing of epicardial coronary arteries is caused most commonly by the formation of a plaque, which is subject to rupture or erosion of the cap separating the plaque from the bloodstream. Upon exposure of the plaque contents to blood, two important and interrelated processes are set in motion: (1) platelets are activated and aggregate, and (2) the coagulation cascade is activated, leading to deposition of fibrin strands. A thrombus composed of platelet aggregates and fibrin strands traps red blood cells and can reduce coronary blood flow, leading to the clinical manifestations of myocardial ischemia.

The location of the obstruction influences the quantity of myocardium rendered ischemic and determines the severity of the clinical manifestations. Thus, critical obstructions in vessels, such as the left main coronary artery and the proximal left anterior descending coronary artery, are particularly hazardous. Chronic severe coronary narrowing and myocardial ischemia frequently are accompanied by the development of collateral vessels, especially when the narrowing develops gradually. When well developed, such vessels can by themselves provide sufficient blood flow to sustain the viability of the myocardium at rest but not during conditions of increased demand.

With progressive worsening of a stenosis in a proximal epicardial artery, the distal resistance vessels (when they function normally) dilate to reduce vascular resistance and maintain coronary blood flow. A pressure gradient develops across the proximal stenosis, and poststenotic pressure falls. When the resistance vessels are maximally dilated, myocardial blood flow becomes dependent on the pressure in the coronary artery distal to the obstruction. In these circumstances, ischemia, manifest clinically by angina or electrocardiographically by ST-segment deviation, can be precipitated by increases in myocardial oxygen demand caused by physical activity, emotional stress, and/or tachycardia. Changes in the caliber of the stenosed coronary artery due to physiologic vasomotion, loss of endothelial control of dilation (as occurs in atherosclerosis), pathologic spasm (Prinzmetal's angina), or small platelet-rich plugs also can upset the critical balance between oxygen supply and demand and thereby precipitate myocardial ischemia.

EFFECTS OF ISCHEMIA

During episodes of inadequate perfusion caused by coronary atherosclerosis, myocardial tissue oxygen tension falls and may cause transient disturbances of the mechanical, biochemical, and electrical functions of the myocardium ([Fig. 293-1](#)). Coronary atherosclerosis is a focal process that usually causes nonuniform ischemia. During ischemia, regional disturbances of ventricular contractility cause segmental hypokinesia, akinesia, or, in severe cases, bulging (dyskinesia), which can reduce myocardial pump function.

The abrupt development of severe ischemia, as occurs with total or subtotal coronary occlusion, is associated with almost instantaneous failure of normal muscle relaxation and then contraction. The relatively poor perfusion of the subendocardium causes more intense ischemia of this portion of the wall (compared with the subepicardial region). Ischemia of large portions of the ventricle causes transient left ventricular failure, and if the papillary muscle apparatus is involved, mitral regurgitation can occur. When ischemia is transient, it may be associated with angina pectoris; when it is prolonged, it can lead to myocardial necrosis and scarring with or without the clinical picture of acute myocardial infarction ([Chap. 295](#)).

A wide range of abnormalities in cell metabolism, function, and structure underlie these mechanical disturbances during ischemia. The normal myocardium metabolizes fatty acids and glucose to carbon

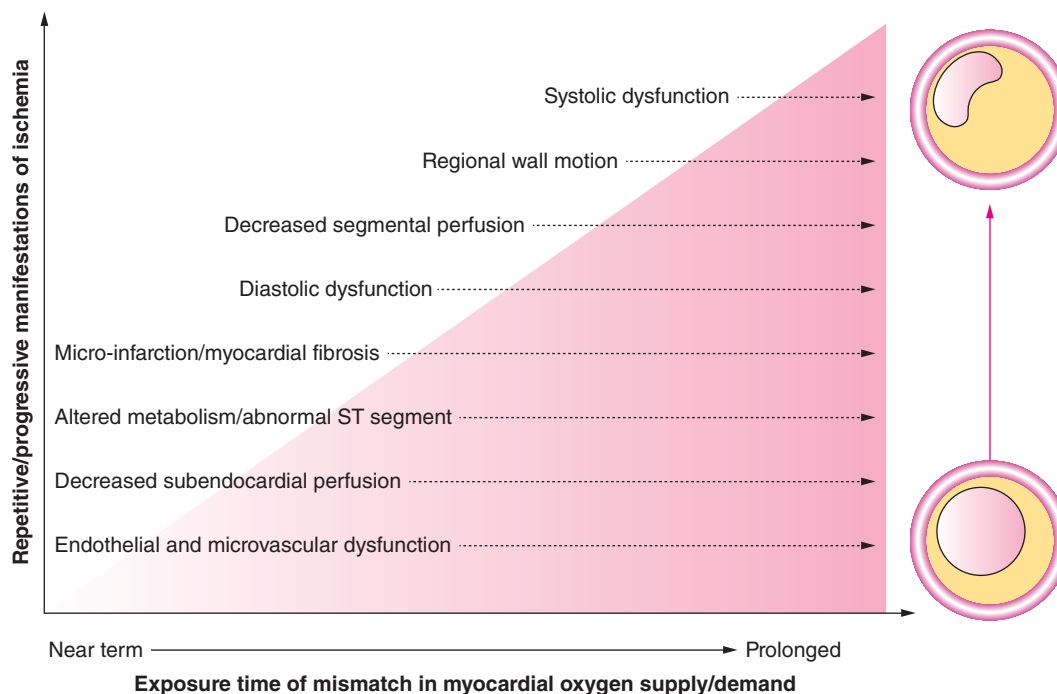


FIGURE 293-1 Cascade of mechanisms and manifestations of ischemia. (Modified from LJ Shaw et al: *J Am Coll Cardiol* 54:1561, 2009. Original figure illustration by Rob Flewell.)

dioxide and water. With severe oxygen deprivation, fatty acids cannot be oxidized, and glucose is converted to lactate; intracellular pH is reduced, as are the myocardial stores of high-energy phosphates, i.e., ATP and creatine phosphate. Impaired cell membrane function leads to the leakage of potassium and the uptake of sodium by myocytes as well as an increase in cytosolic calcium. The severity and duration of the imbalance between myocardial oxygen supply and demand determine whether the damage is reversible (≤ 20 min for total occlusion in the absence of collaterals) or permanent, with subsequent myocardial necrosis (>20 min).

Ischemia also causes characteristic changes in the electrocardiogram (ECG) such as repolarization abnormalities, as evidenced by inversion of T waves and, when more severe, displacement of ST segments (Chap. 268). Transient T-wave inversion probably reflects non-transmural, intramyocardial ischemia; transient ST-segment depression often reflects patchy subendocardial ischemia; and ST-segment elevation is thought to be caused by more severe transmural ischemia. Another important consequence of myocardial ischemia is electrical instability, which may lead to isolated ventricular premature beats or even ventricular tachycardia or ventricular fibrillation (Chap. 277). Most patients who die suddenly from IHD do so as a result of ischemia-induced ventricular tachyarrhythmias (Chap. 327).

ASYMPTOMATIC VERSUS SYMPTOMATIC IHD

Although the prevalence is decreasing, postmortem studies of accident victims and military casualties in Western countries still show that coronary atherosclerosis can begin before age 20 and is present even among adults who were asymptomatic during life. Exercise stress tests in asymptomatic persons may show evidence of silent myocardial ischemia, i.e., exercise-induced ECG changes not accompanied by angina pectoris; coronary angiographic studies of such persons may reveal coronary artery plaques and previously unrecognized obstructions (Chap. 272). Postmortem examination of patients with such obstructions without a history of clinical manifestations of myocardial ischemia often shows macroscopic scars secondary to myocardial infarction in regions supplied by diseased coronary arteries, with or without collateral circulation. According to population studies, $\sim 25\%$ of patients who survive acute myocardial infarction may not come to medical attention, and these patients have the same adverse prognosis as do those who present with the classic clinical picture of acute

myocardial infarction (Chap. 295). Sudden death may be unheralded and is a common presenting manifestation of IHD (Chap. 327).

Patients with IHD also can present with cardiomegaly and heart failure secondary to ischemic damage of the left ventricular myocardium that may have caused no symptoms before the development of heart failure; this condition is referred to as *ischemic cardiomyopathy*. In contrast to the asymptomatic phase of IHD, the symptomatic phase is characterized by chest discomfort due to either angina pectoris or acute myocardial infarction (Chap. 295). Having entered the symptomatic phase, the patient may exhibit a stable or progressive course, revert to the asymptomatic stage, or die suddenly.

STABLE ANGINA PECTORIS

This episodic clinical syndrome is due to transient myocardial ischemia. Various diseases that cause myocardial ischemia and the numerous forms of discomfort with which it may be confused are discussed in Chap. 19. Males constitute $\sim 70\%$ of all patients with angina pectoris and an even greater proportion of those less than 50 years of age. It is, however, important to note that angina pectoris in women is often atypical in presentation (see below).

HISTORY

The typical patient with angina is a man >50 years or a woman >60 years of age who complains of episodes of chest discomfort, usually described as heaviness, pressure, squeezing, smothering, or choking and only rarely as frank pain. When the patient is asked to localize the sensation, he or she typically places a hand over the sternum, sometimes with a clenched fist, to indicate a squeezing, central, substernal discomfort (Levine's sign). Angina is usually crescendo-decrescendo in nature, typically lasts 2 to 5 min, and can radiate to either shoulder and to both arms (especially the ulnar surfaces of the forearm and hand). It also can arise in or radiate to the back, interscapular region, root of the neck, jaw, teeth, and epigastrium. Angina is rarely localized below the umbilicus or above the mandible. A useful finding in assessing a patient with chest discomfort is the fact that myocardial ischemic discomfort does not radiate to the trapezius muscles; that radiation pattern is more typical of pericarditis.

Although episodes of angina typically are caused by exertion (e.g., exercise, hurrying, or sexual activity) or emotion (e.g., stress, anger,

fright, or frustration) and are relieved by rest, they also may occur at rest (Chap. 294) and while the patient is recumbent (angina decubitus). The patient may be awakened at night by typical chest discomfort and dyspnea. Nocturnal angina may be due to episodic tachycardia, diminished oxygenation as the respiratory pattern changes during sleep, or expansion of the intrathoracic blood volume that occurs with recumbency; the latter causes an increase in cardiac size (end-diastolic volume), wall tension, and myocardial oxygen demand that can lead to ischemia and transient left ventricular failure.

The threshold for the development of angina pectoris may vary by time of day and emotional state. Many patients report a fixed threshold for angina, which occurs predictably at a certain level of activity, such as climbing two flights of stairs at a normal pace. In these patients, coronary stenosis and myocardial oxygen supply are fixed, and ischemia is precipitated by an increase in myocardial oxygen demand; they are said to have stable exertional angina. In other patients, the threshold for angina may vary considerably within any particular day and from day to day. In such patients, variations in myocardial oxygen supply, most likely due to changes in coronary vasomotor tone, may play an important role in defining the pattern of angina. A patient may report symptoms upon minor exertion in the morning (a short walk or shaving) yet by midday be capable of much greater effort without symptoms. Angina may also be precipitated by unfamiliar tasks, a heavy meal, exposure to cold, or a combination of these factors.

Exertional angina typically is relieved in 1–5 min by slowing or ceasing activities and even more rapidly by rest and sublingual nitroglycerin (see below). Indeed, the diagnosis of angina should be suspect if it does not respond to the combination of these measures. The severity of angina can be conveniently summarized by the Canadian Cardiac Society functional classification (Table 293-1). Its impact on the patient's functional capacity can be described by using the New York Heart Association functional classification (Table 293-1).

TABLE 293-1 CARDIOVASCULAR DISEASE CLASSIFICATION CHART

Class	New York Heart Association Functional Classification	Canadian Cardiovascular Society Functional Classification
I	Patients have cardiac disease but <i>without</i> the resulting limitations of physical activity. Ordinary physical activity does not cause undue fatigue, palpitation, dyspnea, or anginal pain.	Ordinary physical activity, such as walking and climbing stairs, <i>does not cause angina</i> . Angina present with strenuous or rapid or prolonged exertion at work or recreation.
II	Patients have cardiac disease resulting in <i>slight limitation</i> of physical activity. They are comfortable at rest. Ordinary physical activity results in fatigue, palpitation, dyspnea, or anginal pain.	<i>Slight limitation</i> of ordinary activity. Walking or climbing stairs rapidly, walking uphill, walking or stair climbing after meals, in cold, or when under emotional stress or only during the few hours after awakening. Walking more than two blocks on the level and climbing more than one flight of stairs at a normal pace and in normal conditions.
III	Patients have cardiac disease resulting in <i>marked limitation</i> of physical activity. They are comfortable at rest. Less than ordinary physical activity causes fatigue, palpitation, dyspnea, or anginal pain.	<i>Marked limitation</i> of ordinary physical activity. Walking one to two blocks on the level and climbing more than one flight of stairs in normal conditions.
IV	Patients have cardiac disease resulting in <i>inability</i> to carry on any physical activity without discomfort. Symptoms of cardiac insufficiency or of the anginal syndrome may be present even at rest. If any physical activity is undertaken, discomfort is increased.	<i>Inability</i> to carry on any physical activity without discomfort—anginal syndrome <i>may</i> be present at rest.

Source: Modified from L Goldman et al: Circulation 64:1227, 1981.

Sharp, fleeting chest pain or a prolonged, dull ache localized to the left submammary area is rarely due to myocardial ischemia. However, especially in women and diabetic patients, angina pectoris may be atypical in location and not strictly related to provoking factors. In addition, this symptom may exacerbate and remit over days, weeks, or months. Its occurrence can be seasonal, occurring more frequently in the winter in temperate climates. Anginal “equivalents” are symptoms of myocardial ischemia other than angina. They include dyspnea, nausea, fatigue, and faintness and are more common in the elderly and in diabetic patients.

Systematic questioning of a patient with suspected IHD is important to uncover the features of an unstable syndrome associated with increased risk, such as angina occurring with less exertion than in the past, occurring at rest, or awakening the patient from sleep. Since coronary atherosclerosis often is accompanied by similar lesions in other arteries, a patient with angina should be questioned and examined for peripheral arterial disease (intermittent claudication [Chap. 302]), stroke, or transient ischemic attacks (Chap. 446). It is also important to uncover a family history of premature IHD (<55 years in first-degree male relatives and <65 in female relatives) and the presence of diabetes mellitus, hyperlipidemia, hypertension, cigarette smoking, and other risk factors for coronary atherosclerosis (Chap. 291e).

The history of typical angina pectoris establishes the diagnosis of IHD until proven otherwise. The coexistence of advanced age, male sex, the postmenopausal state, and risk factors for atherosclerosis increase the likelihood of hemodynamically significant coronary disease. A particularly challenging problem is the evaluation and management of patients with persistent ischemic-type chest discomfort but no flow-limiting obstructions in their epicardial coronary arteries. This situation arises more often in women than in men. Potential etiologies include microvascular coronary disease (detectable on coronary reactivity testing in response to vasoactive agents such as intracoronary adenosine, acetylcholine, and nitroglycerin) and abnormal cardiac nociception. Treatment of microvascular coronary disease should focus on efforts to improve endothelial function, including nitrates, beta blockers, calcium antagonists, statins, and angiotensin-converting enzyme (ACE) inhibitors. Abnormal cardiac nociception is more difficult to manage and may be ameliorated in some cases by imipramine.

PHYSICAL EXAMINATION

The physical examination is often normal in patients with stable angina when they are asymptomatic. However, because of the increased likelihood of IHD in patients with diabetes and/or peripheral arterial disease, clinicians should search for evidence of atherosclerotic disease at other sites, such as an abdominal aortic aneurysm, carotid arterial bruits, and diminished arterial pulses in the lower extremities. The physical examination also should include a search for evidence of risk factors for atherosclerosis such as xanthelasmas and xanthomas (Chap. 291e). Evidence for peripheral arterial disease should be sought by evaluating the pulse contour at multiple locations and comparing the blood pressure between the arms and between the arms and the legs (ankle-brachial index). Examination of the fundi may reveal an increased light reflex and arteriovenous nicking as evidence of hypertension. There also may be signs of anemia, thyroid disease, and nicotine stains on the fingertips from cigarette smoking.

Palpation may reveal cardiac enlargement and abnormal contraction of the cardiac impulse (left ventricular dyskinesia). Auscultation can uncover arterial bruits, a third and/or fourth heart sound, and, if acute ischemia or previous infarction has impaired papillary muscle function, an apical systolic murmur due to mitral regurgitation. These auscultatory signs are best appreciated with the patient in the left lateral decubitus position. Aortic stenosis, aortic regurgitation (Chap. 283), pulmonary hypertension (Chap. 304), and hypertrophic cardiomyopathy (Chap. 287) must be excluded, since these disorders may cause angina in the absence of coronary atherosclerosis. Examination during an anginal attack is useful, since ischemia can cause transient left ventricular failure with the appearance of a third

and/or fourth heart sound, a dyskinetic cardiac apex, mitral regurgitation, and even pulmonary edema. Tenderness of the chest wall, localization of the discomfort with a single fingertip on the chest, or reproduction of the pain with palpation of the chest makes it unlikely that the pain is caused by myocardial ischemia. A protuberant abdomen may indicate that the patient has the metabolic syndrome and is at increased risk for atherosclerosis.

LABORATORY EXAMINATION

Although the diagnosis of IHD can be made with a high degree of confidence from the history and physical examination, a number of simple laboratory tests can be helpful. The urine should be examined for evidence of diabetes mellitus and renal disease (including microalbuminuria) since these conditions accelerate atherosclerosis. Similarly, examination of the blood should include measurements of lipids (cholesterol—total, LDL, HDL—and triglycerides), glucose (hemoglobin A_{1c}), creatinine, hematocrit, and, if indicated based on the physical examination, thyroid function. A chest x-ray is important as it may show the consequences of IHD, i.e., cardiac enlargement, ventricular aneurysm, or signs of heart failure. These signs can support the diagnosis of IHD and are important in assessing the degree of cardiac damage. Evidence exists that an elevated level of high-sensitivity C-reactive protein (CRP) (specifically, between 0 and 3 mg/dL) is an independent risk factor for IHD and may be useful in therapeutic decision making about the initiation of hypolipidemic treatment. The major benefit of high-sensitivity CRP is in reclassifying the risk of IHD in patients in the “intermediate” risk category on the basis of traditional risk factors.

ELECTROCARDIOGRAM

A 12-lead ECG recorded at rest may be normal in patients with typical angina pectoris, but there may also be signs of an old myocardial infarction (Chap. 268). Although repolarization abnormalities, i.e., ST-segment and T-wave changes, as well as left ventricular hypertrophy and disturbances of cardiac rhythm or intraventricular conduction are suggestive of IHD, they are nonspecific, since they also can occur in pericardial, myocardial, and valvular heart disease or, in the case of the former, transiently with anxiety, changes in posture, drugs, or esophageal disease. The presence of left ventricular hypertrophy (LVH) is a significant indication of increased risk of adverse outcomes from IHD. Of note, even though LVH and cardiac rhythm disturbances are nonspecific indicators of the development of IHD, they may be contributing factors to episodes of angina in patients in whom IHD has developed as a consequence of conventional risk factors. Dynamic ST-segment and T-wave changes that accompany episodes of angina pectoris and disappear thereafter are more specific.

STRESS TESTING

Electrocardiographic The most widely used test for both the diagnosis of IHD and the estimation of risk and prognosis involves recording the 12-lead ECG before, during, and after exercise, usually on a treadmill (Fig. 293-2). The test consists of a standardized incremental increase in external workload (Table 293-2) while symptoms, the ECG, and arm blood pressure are monitored. Exercise duration is usually symptom-limited, and the test is discontinued upon evidence of chest discomfort, severe shortness of breath, dizziness, severe fatigue, ST-segment depression >0.2 mV (2 mm), a fall in systolic blood pressure >10 mmHg, or the development of a ventricular tachyarrhythmia. This test is used to discover any limitation in exercise performance, detect typical ECG signs of myocardial ischemia, and establish their relationship to chest discomfort. The ischemic ST-segment response generally is defined as flat or downsloping depression of the ST segment >0.1 mV below baseline (i.e., the PR segment) and lasting longer than 0.08 s (Fig. 293-1). Upsloping or junctional ST-segment changes are not considered characteristic of ischemia and do not constitute a positive test. Although T-wave abnormalities, conduction disturbances, and ventricular arrhythmias that develop during exercise should be noted, they are also not diagnostic. Negative exercise tests in which the target heart rate (85% of maximal predicted heart rate for age and sex) is not achieved are considered nondiagnostic.

In interpreting ECG stress tests, the probability that coronary artery disease (CAD) exists in the patient or population under study (i.e., pre-test probability) should be considered. Overall, false-positive or false-negative results occur in one-third of cases. However, a positive result on exercise indicates that the likelihood of CAD is 98% in males who are >50 years with a history of typical angina pectoris and who develop chest discomfort during the test. The likelihood decreases if the patient has atypical or no chest pain by history and/or during the test.

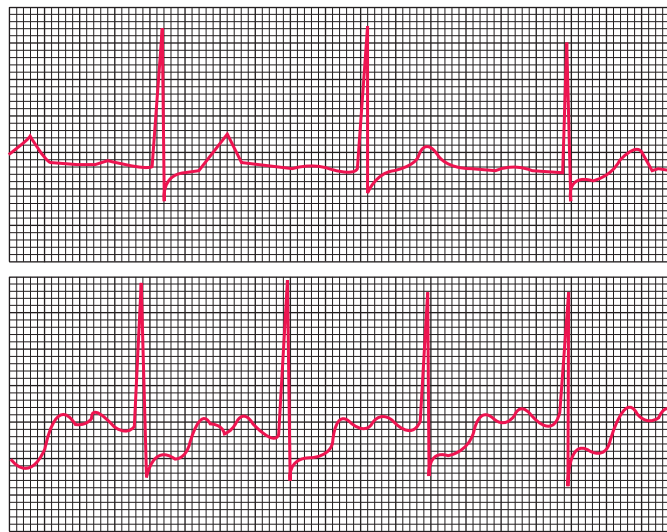
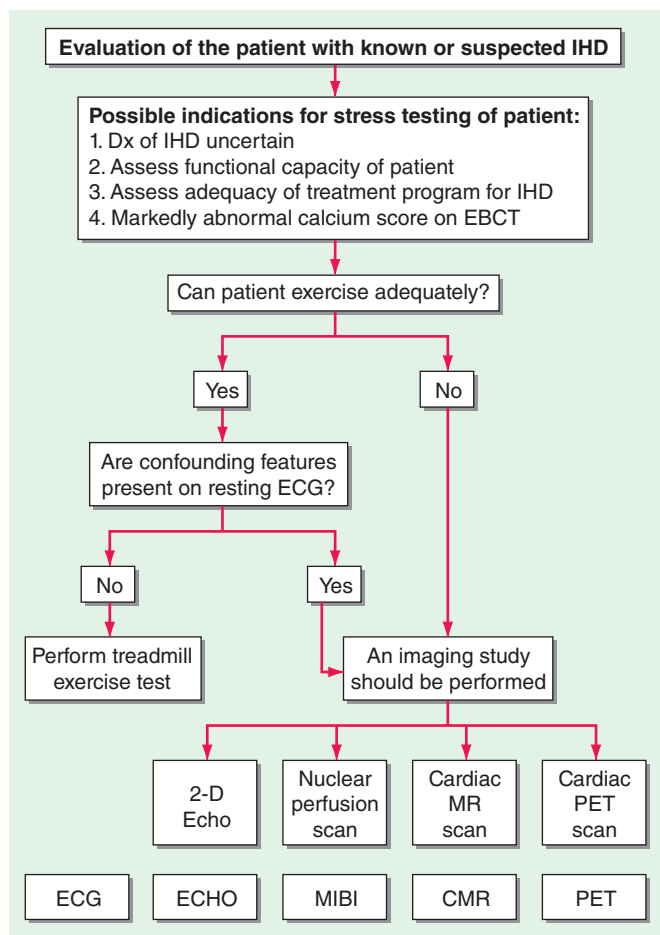
The incidence of false-positive tests is significantly increased in patients with low probabilities of IHD, such as asymptomatic men age <40 or premenopausal women with no risk factors for premature atherosclerosis. It is also increased in patients taking cardioactive drugs, such as digitalis and antiarrhythmic agents, and in those with intraventricular conduction disturbances, resting ST-segment and T-wave abnormalities, ventricular hypertrophy, or abnormal serum potassium levels. Obstructive disease limited to the circumflex coronary artery may result in a false-negative stress test since the lateral portion of the heart that this vessel supplies is not well represented on the surface 12-lead ECG. Since the overall sensitivity of exercise stress electrocardiography is only $\sim 75\%$, a negative result does not exclude CAD, although it makes the likelihood of three-vessel or left main CAD extremely unlikely.

A medical professional should be present throughout the exercise test. It is important to measure total duration of exercise, the times to the onset of ischemic ST-segment change and chest discomfort, the external work performed (generally expressed as the stage of exercise), and the internal cardiac work performed, i.e., by the heart rate–blood pressure product. The depth of the ST-segment depression and the time needed for recovery of these ECG changes are also important. Because the risks of exercise testing are small but real—estimated at one fatality and two nonfatal complications per 10,000 tests—equipment for resuscitation should be available. Modified (heart rate–limited rather than symptom–limited) exercise tests can be performed safely in patients as early as 6 days after uncomplicated myocardial infarction (Table 293-2). Contraindications to exercise stress testing include rest angina within 48 h, unstable rhythm, severe aortic stenosis, acute myocarditis, uncontrolled heart failure, severe pulmonary hypertension, and active infective endocarditis.

The normal response to graded exercise includes progressive increases in heart rate and blood pressure. Failure of the blood pressure to increase or an actual decrease with signs of ischemia during the test is an important adverse prognostic sign, since it may reflect ischemia-induced global left ventricular dysfunction. The development of angina and/or severe (>0.2 mV) ST-segment depression at a low workload, i.e., before completion of stage II of the Bruce protocol, and/or ST-segment depression that persists >5 min after the termination of exercise increases the specificity of the test and suggests severe IHD and a high risk of future adverse events.

Cardiac Imaging (See also Chap. 270e) When the resting ECG is abnormal (e.g., preexcitation syndrome, >1 mm of resting ST-segment depression, left bundle branch block, paced ventricular rhythm), information gained from an exercise test can be enhanced by stress myocardial radionuclide perfusion imaging after the intravenous administration of thallium-201 or 99m-technetium sestamibi during exercise (or with pharmacologic stress). Contemporary data also suggest positron emission tomography (PET) imaging (with exercise or pharmacologic stress) using N-13 ammonia or rubidium-82 nuclide as another technique for assessing perfusion. Images obtained immediately after cessation of exercise to detect regional ischemia are compared with those obtained at rest to confirm reversible ischemia and regions of persistently absent uptake that signify infarction.

A sizable fraction of patients who need noninvasive stress testing to identify myocardial ischemia and increased risk of coronary events cannot exercise because of peripheral vascular or musculoskeletal disease, exertional dyspnea, or deconditioning. In these circumstances, an intravenous pharmacologic challenge is used in place of exercise. For example, dipyridamole or adenosine can be given to create a coronary “steal” by temporarily increasing flow in nondiseased



A

FIGURE 293-2 Evaluation of the patient with known or suspected ischemic heart disease. On the left of the figure is an algorithm for identifying patients who should be referred for stress testing and the decision pathway for determining whether a standard treadmill exercise with electrocardiogram (ECG) monitoring alone is adequate. A specialized imaging study is necessary if the patient cannot exercise adequately (pharmacologic challenge is given) or if there are confounding features on the resting ECG (symptom-limited treadmill exercise may be used to stress the coronary circulation). Panels B–E, continued on the next page, are examples of the data obtained with ECG monitoring and specialized imaging procedures. CMR, cardiac magnetic resonance; EBCT, electron beam computed tomography; ECHO, echocardiography; IHD, ischemic heart disease; MIBI, methoxyisobutyl isonitrite; MR, magnetic resonance; PET, positron emission tomography. **A.** Lead V_4 at rest (*top panel*) and after 4.5 min of exercise (*bottom panel*). There is 3 mm (0.3 mV) of horizontal ST-segment depression, indicating a positive test for ischemia. (Modified from BR Chaitman, in E Braunwald et al [eds]: *Heart Disease*, 8th ed, Philadelphia, Saunders, 2008.) **B.** A 45-year-old avid jogger who began experiencing classic substernal chest pressure underwent an exercise echo study. With exercise the patient's heart rate increased from 52 to 153 beats/min. The left ventricular chamber dilated with exercise, and the septal and apical portions became akinetic to dyskinetic (*red arrow*). These findings are strongly suggestive of a significant flow-limiting stenosis in the proximal left anterior descending artery, which was confirmed at coronary angiography. (Modified from SD Solomon, in E. Braunwald et al [eds]: *Primary Cardiology*, 2nd ed, Philadelphia, Saunders, 2003.) **C.** Stress and rest myocardial perfusion single-photon emission computed tomography images obtained with 99m -technetium sestamibi in a patient with chest pain and dyspnea on exertion. The images demonstrate a medium-size and severe stress perfusion defect involving the inferolateral and basal inferior walls, showing nearly complete reversibility, consistent with moderate ischemia in the right coronary artery territory (*red arrows*). (Images provided by Dr. Marcello Di Carli, Nuclear Medicine Division, Brigham and Women's Hospital, Boston, MA.) **D.** A patient with a prior myocardial infarction presented with recurrent chest discomfort. On cardiac magnetic resonance (CMR) cine imaging, a large area of anterior akinesia was noted (marked by the *arrows* in the top left and right images, systolic frame only). This area of akinesia was matched by a larger extent of late gadolinium-DTPA enhancements consistent with a large transmural myocardial infarction (marked by *arrows* in the middle left and right images). Resting (*bottom left*) and adenosine vasodilating stress (*bottom right*) first-pass perfusion images revealed reversible perfusion abnormality that extended to the inferior septum. This patient was found to have an occluded proximal left anterior descending coronary artery with extensive collateral formation. This case illustrates the utility of different modalities in a CMR examination in characterizing ischemic and infarcted myocardium. DTPA, diethylenetriamine penta-acetic acid. (Images provided by Dr. Raymond Kwong, Cardiovascular Division, Brigham and Women's Hospital, Boston, MA.) **E.** Stress and rest myocardial perfusion PET images obtained with rubidium-82 in a patient with chest pain on exertion. The images demonstrate a large and severe stress perfusion defect involving the mid and apical anterior, anterolateral, and anteroseptal walls and the left ventricular apex, showing complete reversibility, consistent with extensive and severe ischemia in the mid-left anterior descending coronary artery territory (*red arrows*). (Images provided by Dr. Marcello Di Carli, Nuclear Medicine Division, Brigham and Women's Hospital, Boston, MA.)

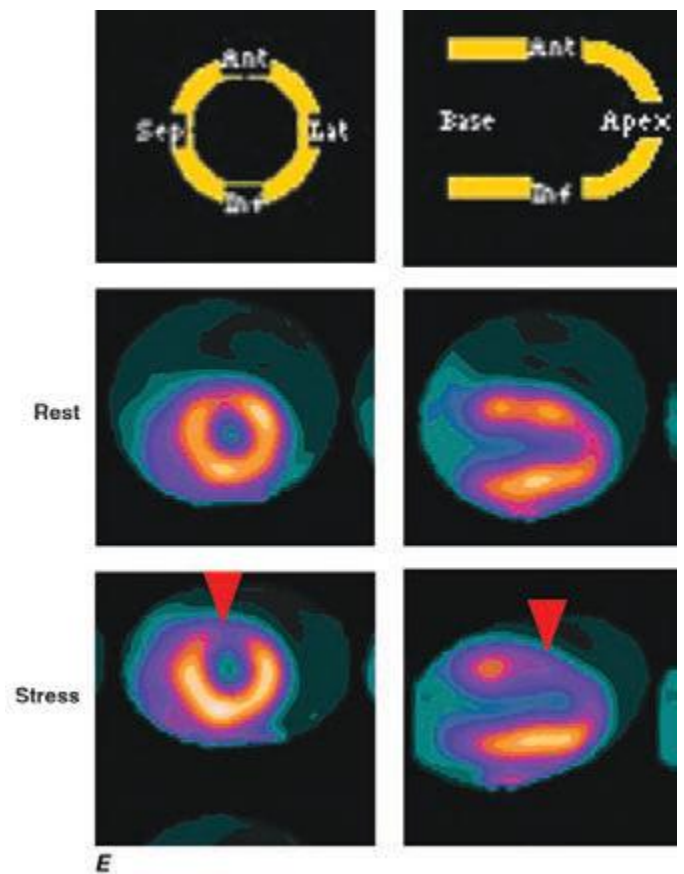
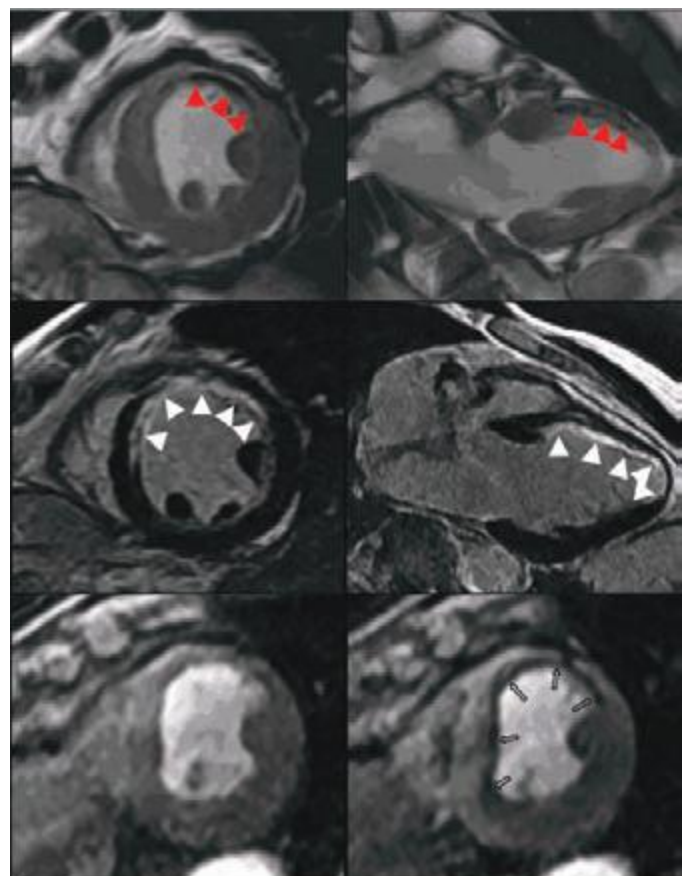
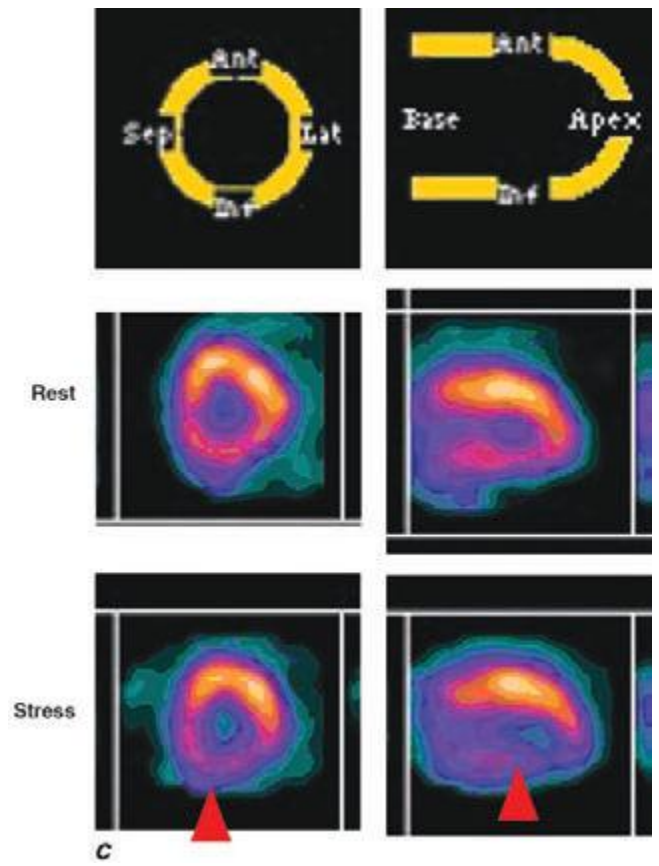
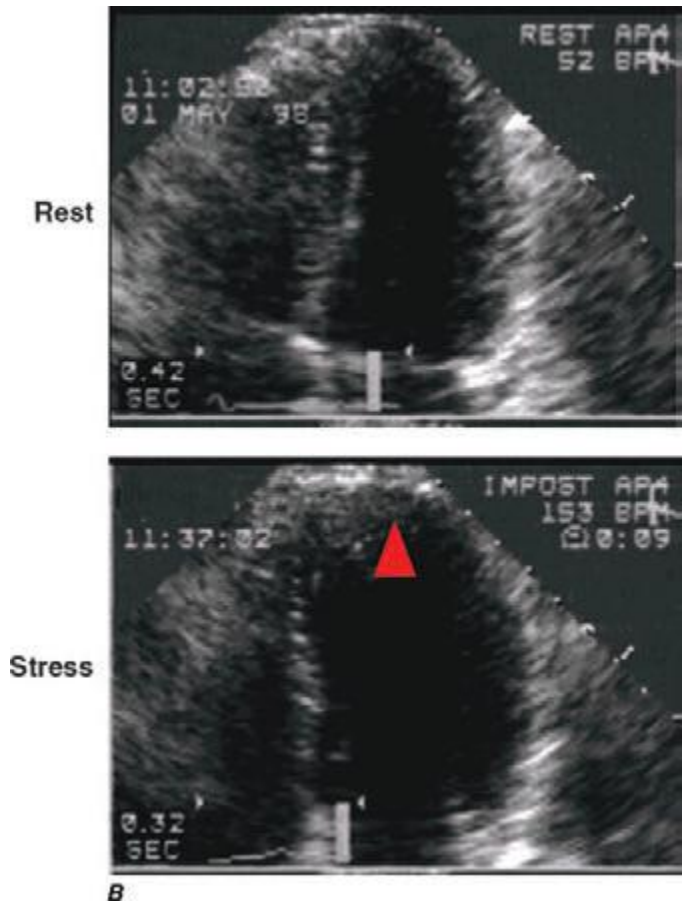


FIGURE 293-2 (Continued)

TABLE 293-2 RELATION OF METABOLIC EQUIVALENT TASKS (METs) TO STAGES IN VARIOUS TESTING PROTOCOLS

Functional Class	Clinical Status	O ₂ Cost mL/kg/min	METs	Treadmill Protocols					
				BRUCE Modified 3 min Stages		BRUCE 3 min Stages			
				MPH	%GR	MPH	%GR		
NORMAL AND I	HEALTHY, DEPENDENT ON AGE, ACTIVITY			6.0	22	6.0	22		
				5.5	20	5.2	20		
				5.0	18	5.0	18		
				56.0	16				
				52.5	15				
				49.0	14				
				45.5	13	4.2	16	4.2	16
				42.0	12				
				38.5	11	3.4	14	3.4	14
				35.0	10				
				31.5	9				
				28.0	8				
		24.5	7	2.5	12	2.5	12		
II	SEMI-HEALTHY LIMITED SYMPTOMATIC								
				21.0	6				
					17.5	5	1.7	10	
					14.0	4			
III									
				10.5	3	1.7	5		
				7.0	2	1.7	0		
IV				3.5	1				

Abbreviations: GR, grade; MPH, miles per hour.

Source: Modified from GF Fletcher et al: Circulation 104:1694, 2001.

segments of the coronary vasculature at the expense of diseased segments. Alternatively, a graded incremental infusion of dobutamine may be administered to increase MVO_2 . A variety of imaging options are available to accompany these pharmacologic stressors (Fig. 293-2). The development of a transient perfusion defect with a tracer such as thallium-201 or 99m-technetium sestamibi is used to detect myocardial ischemia.

Echocardiography is used to assess left ventricular function in patients with chronic stable angina and patients with a history of a prior myocardial infarction, pathologic Q waves, or clinical evidence of heart failure. Two-dimensional echocardiography can assess both global and regional wall motion abnormalities of the left ventricle that are transient when due to ischemia. Stress (exercise or dobutamine) echocardiography may cause the emergence of regions of akinesis or dyskinesis that are not present at rest. Stress echocardiography, like stress myocardial perfusion imaging, is more sensitive than exercise electrocardiography in the diagnosis of IHD. Cardiac magnetic resonance (CMR) stress testing is also evolving as an alternative to radio-nuclide, PET, or echocardiographic stress imaging. CMR stress testing performed with dobutamine infusion can be used to assess wall motion abnormalities accompanying ischemia, as well as myocardial perfusion. CMR can be used to provide more complete ventricular evaluation using multislice magnetic resonance imaging (MRI) studies.

Atherosclerotic plaques become progressively calcified over time, and coronary calcification in general increases with age. For this reason, methods for detecting coronary calcium have been developed as a measure of the presence of coronary atherosclerosis. These methods involve computed tomography (CT) applications that achieve rapid acquisition of images (electron beam [EBCT] and multidetector [MDCT] detection). Coronary calcium detected by these imaging techniques most commonly is quantified by using the Agatston score, which is based on the area and density of calcification. Although the diagnostic accuracy of this imaging method is high (sensitivity, 90–94%; specificity, 95–97%; negative predictive value, 93–99%), its prognostic utility has not been defined. Thus, its role in CT, EBCT, and MDCT scans for the detection and management of patients with IHD has not been clarified.

CORONARY ARTERIOGRAPHY

(See also Chap. 272) This diagnostic method outlines the lumina of the coronary arteries and can be used to detect or exclude serious coronary obstruction. However, coronary arteriography provides no information about the arterial wall, and severe atherosclerosis that does not encroach on the lumen may go undetected. Of note, atherosclerotic plaques characteristically are scattered throughout the coronary tree, tend to occur more frequently at branch points, and grow progressively in the intima and media of an epicardial coronary artery at first without encroaching on the lumen, causing an outward bulging of the artery—a process referred to as remodeling (Chap. 291e). Later in the course of the disease, further growth causes luminal narrowing.

Indications Coronary arteriography is indicated in (1) patients with chronic stable angina pectoris who are severely symptomatic despite medical therapy and are being considered for revascularization, i.e., a percutaneous coronary intervention (PCI) or coronary artery bypass grafting (CABG); (2) patients with troublesome symptoms that present diagnostic difficulties in whom there is a need to confirm or rule out the diagnosis of IHD; (3) patients with known or possible angina pectoris who have survived cardiac arrest; (4) patients with angina or evidence of ischemia on noninvasive testing with clinical or laboratory evidence of ventricular dysfunction; and (5) patients judged to be at high risk of sustaining coronary events based on signs of severe ischemia on noninvasive testing, regardless of the presence or severity of symptoms (see below).

Examples of other indications for coronary arteriography include the following:

1. Patients with chest discomfort suggestive of angina pectoris but a negative or nondiagnostic stress test who require a definitive diagnosis for guiding medical management, alleviating psychological stress, career or family planning, or insurance purposes.
2. Patients who have been admitted repeatedly to the hospital for a suspected acute coronary syndrome (Chaps. 294 and 295), but in whom this diagnosis has not been established and in whom the presence or absence of CAD should be determined.

3. Patients with careers that involve the safety of others (e.g., pilots, firefighters, police) who have questionable symptoms or suspicious or positive noninvasive tests and in whom there are reasonable doubts about the state of the coronary arteries.
4. Patients with aortic stenosis or hypertrophic cardiomyopathy and angina in whom the chest pain could be due to IHD.
5. Male patients >45 years and females >55 years who are to undergo a cardiac operation such as valve replacement or repair and who may or may not have clinical evidence of myocardial ischemia.
6. Patients after myocardial infarction, especially those who are at high risk after myocardial infarction because of the recurrence of angina or the presence of heart failure, frequent ventricular premature contractions, or signs of ischemia on the stress test.
7. Patients with angina pectoris, regardless of severity, in whom noninvasive testing indicates a high risk of coronary events (poor exercise performance or severe ischemia).
8. Patients in whom coronary spasm or another nonatherosclerotic cause of myocardial ischemia (e.g., coronary artery anomaly, Kawasaki disease) is suspected.

Noninvasive alternatives to diagnostic coronary arteriography include CT angiography and CMR angiography (Chap. 270e). Although these new imaging techniques can provide information about obstructive lesions in the epicardial coronary arteries, their exact role in clinical practice has not been rigorously defined. Important aspects of their use that should be noted include the substantially higher radiation exposure with CT angiography compared to conventional diagnostic arteriography and the limitations on CMR imposed by cardiac movement during the cardiac cycle, especially at high heart rates.

PROGNOSIS

The principal prognostic indicators in patients known to have IHD are age, the functional state of the left ventricle, the location(s) and severity of coronary artery narrowing, and the severity or activity of myocardial ischemia. Angina pectoris of recent onset, unstable angina (Chap. 294), early postmyocardial infarction angina, angina that is unresponsive or poorly responsive to medical therapy, and angina accompanied by symptoms of congestive heart failure all indicate an increased risk for adverse coronary events. The same is true for the physical signs of heart failure, episodes of pulmonary edema, transient third heart sounds, and mitral regurgitation and for echocardiographic or radioisotopic (or roentgenographic) evidence of cardiac enlargement and reduced (<0.40) ejection fraction.

Most important, any of the following signs during noninvasive testing indicates a high risk for coronary events: inability to exercise for 6 min, i.e., stage II (Bruce protocol) of the exercise test; a strongly positive exercise test showing onset of myocardial ischemia at low workloads (≥ 0.1 mV ST-segment depression before completion of stage II, ≥ 0.2 mV ST-segment depression at any stage, ST-segment depression for >5 min after the cessation of exercise, a decline in systolic pressure >10 mmHg during exercise, or the development of ventricular tachyarrhythmias during exercise); the development of large or multiple perfusion defects or increased lung uptake during stress radioisotope perfusion imaging; and a decrease in left ventricular ejection fraction during exercise on radionuclide ventriculography or during stress echocardiography. Conversely, patients who can complete stage III of the Bruce exercise protocol and have a normal stress perfusion scan or negative stress echocardiographic evaluation are at very low risk for future coronary events. The finding of frequent episodes of ST-segment deviation on ambulatory ECG monitoring (even in the absence of symptoms) is also an adverse prognostic finding.

On cardiac catheterization, elevations of left ventricular end-diastolic pressure and ventricular volume and reduced ejection fraction are the most important signs of left ventricular dysfunction and are associated with a poor prognosis. Patients with chest discomfort but normal left ventricular function and normal coronary arteries have an excellent prognosis. Obstructive lesions of the left main (>50% luminal diameter) or left anterior descending coronary artery proximal to the origin of the first septal artery are associated with

a greater risk than are lesions of the right or left circumflex coronary artery because of the greater quantity of myocardium at risk. Atherosclerotic plaques in epicardial arteries with fissuring or filling defects indicate increased risk. These lesions go through phases of inflammatory cellular activity, degeneration, endothelial dysfunction, abnormal vasomotion, platelet aggregation, and fissuring or hemorrhage. These factors can temporarily worsen the stenosis and cause thrombosis and/or abnormal reactivity of the vessel wall, thus exacerbating the manifestations of ischemia. The recent onset of symptoms, the development of severe ischemia during stress testing (see above), and unstable angina pectoris (Chap. 294) all reflect episodes of rapid progression in coronary lesions.

With any degree of obstructive CAD, mortality is greatly increased when left ventricular function is impaired; conversely, at any level of left ventricular function, the prognosis is influenced importantly by the quantity of myocardium perfused by critically obstructed vessels. Therefore, it is essential to collect all the evidence substantiating past myocardial damage (evidence of myocardial infarction on ECG, echocardiography, radioisotope imaging, or left ventriculography), residual left ventricular function (ejection fraction and wall motion), and risk of future damage from coronary events (extent of coronary disease and severity of ischemia defined by noninvasive stress testing). The larger the quantity of established myocardial necrosis is, the less the heart is able to withstand additional damage and the poorer the prognosis is. Risk estimation must include age, presenting symptoms, all risk factors, signs of arterial disease, existing cardiac damage, and signs of impending damage (i.e., ischemia).

The greater the number and severity of risk factors for coronary atherosclerosis (advanced age [>75 years], hypertension, dyslipidemia, diabetes, morbid obesity, accompanying peripheral and/or cerebrovascular disease, previous myocardial infarction), the worse the prognosis of an angina patient. Evidence exists that elevated levels of CRP in the plasma, extensive coronary calcification on electron beam CT (see above), and increased carotid intimal thickening on ultrasound examination also indicate an increased risk of coronary events.

TREATMENT STABLE ANGINA PECTORIS

Once the diagnosis of IHD has been made, each patient must be evaluated individually with respect to his or her level of understanding, expectations and goals, control of symptoms, and prevention of adverse clinical outcomes such as myocardial infarction and premature death. The degree of disability and the physical and emotional stress that precipitates angina must be recorded carefully to set treatment goals. The management plan should include the following components: (1) explanation of the problem and reassurance about the ability to formulate a treatment plan, (2) identification and treatment of aggravating conditions, (3) recommendations for adaptation of activity as needed, (4) treatment of risk factors that will decrease the occurrence of adverse coronary outcomes, (5) drug therapy for angina, and (6) consideration of revascularization.

EXPLANATION AND REASSURANCE

Patients with IHD need to understand their condition and realize that a long and productive life is possible even though they have angina pectoris or have experienced and recovered from an acute myocardial infarction. Offering results of clinical trials showing improved outcomes can be of great value in encouraging patients to resume or maintain activity and return to work. A planned program of rehabilitation can encourage patients to lose weight, improve exercise tolerance, and control risk factors with more confidence.

IDENTIFICATION AND TREATMENT OF AGGRAVATING CONDITIONS

A number of conditions may increase oxygen demand or decrease oxygen supply to the myocardium and may precipitate or exacerbate angina in patients with IHD. Left ventricular hypertrophy,

aortic valve disease, and hypertrophic cardiomyopathy may cause or contribute to angina and should be excluded or treated. Obesity, hypertension, and hyperthyroidism should be treated aggressively to reduce the frequency and severity of anginal episodes. Decreased myocardial oxygen supply may be due to reduced oxygenation of the arterial blood (e.g., in pulmonary disease or, when carboxyhemoglobin is present, due to cigarette or cigar smoking) or decreased oxygen-carrying capacity (e.g., in anemia). Correction of these abnormalities, if present, may reduce or even eliminate angina pectoris.

ADAPTATION OF ACTIVITY

Myocardial ischemia is caused by a discrepancy between the demand of the heart muscle for oxygen and the ability of the coronary circulation to meet that demand. Most patients can be helped to understand this concept and utilize it in the rational programming of activity. Many tasks that ordinarily evoke angina may be accomplished without symptoms simply by reducing the speed at which they are performed. Patients must appreciate the diurnal variation in their tolerance of certain activities and should reduce their energy requirements in the morning, immediately after meals, and in cold or inclement weather. On occasion, it may be necessary to recommend a change in employment or residence to avoid physical stress.

Physical conditioning usually improves the exercise tolerance of patients with angina and has substantial psychological benefits. A regular program of isotonic exercise that is within the limits of the individual patient's threshold for the development of angina pectoris and that does not exceed 80% of the heart rate associated with ischemia on exercise testing should be strongly encouraged. Based on the results of an exercise test, the number of metabolic equivalent tasks (METs) performed at the onset of ischemia can be estimated (Table 293-2) and a practical exercise prescription can be formulated to permit daily activities that will fall below the ischemic threshold (Table 293-3).

TREATMENT OF RISK FACTORS

A *family history* of premature IHD is an important indicator of increased risk and should trigger a search for treatable risk factors such as hyperlipidemia, hypertension, and diabetes mellitus. *Obesity* impairs the treatment of other risk factors and increases the risk of adverse coronary events. In addition, obesity often is accompanied by three other risk factors: diabetes mellitus, hypertension, and hyperlipidemia. The treatment of obesity and these accompanying risk factors is an important component of any management plan. A diet low in saturated and *trans*-unsaturated fatty acids and a reduced caloric intake to achieve optimal body weight are a cornerstone in the management of chronic IHD. It is especially important to emphasize weight loss and regular exercise in patients with the metabolic syndrome or overt diabetes mellitus.

Cigarette smoking accelerates coronary atherosclerosis in both sexes and at all ages and increases the risk of thrombosis, plaque instability, myocardial infarction, and death (Chap. 291e). In addition, by increasing myocardial oxygen needs and reducing oxygen supply, it aggravates angina. Smoking cessation studies have demonstrated important benefits with a significant decline in the occurrence of these adverse outcomes. The physician's message must be clear and strong and supported by programs that achieve and monitor abstinence (Chap. 470). *Hypertension* (Chap. 298) is associated with an increased risk of adverse clinical events from coronary atherosclerosis as well as stroke. In addition, the left ventricular hypertrophy that results from sustained hypertension aggravates ischemia. There is evidence that long-term effective treatment of hypertension can decrease the occurrence of adverse coronary events.

Diabetes mellitus (Chap. 417) accelerates coronary and peripheral atherosclerosis and is frequently associated with dyslipidemias and increases in the risk of angina, myocardial infarction, and sudden coronary death. Aggressive control of the dyslipidemia (target LDL cholesterol <70 mg/dL) and hypertension (target blood pressure 120/80 mmHg) that are frequently found in diabetic patients is highly effective and therefore essential, as described below.

TABLE 293-3 ENERGY REQUIREMENTS FOR SOME COMMON ACTIVITIES

Less Than 3 METs	3–5 METs	5–7 METs	7–9 METs	More Than 9 METs
Self-Care				
Washing/shaving	Cleaning windows	Easy digging in garden	Heavy shoveling	Carrying loads up stairs (objects more than 90 lb)
Dressing	Raking	Level hand lawn mowing	Carrying objects (60–90 lb)	Climbing stairs (quickly)
Light housekeeping	Power lawn mowing	Carrying objects (30–60 lb)		Shoveling heavy snow
Desk work	Bed making/stripping			
Driving auto	Carrying objects (15–30 lb)			
Occupational				
Sitting (clerical/assembly)	Stocking shelves (light objects)	Carpentry (exterior)	Digging ditches (pick and shovel)	Heavy labor
Desk work	Light welding/carpentry	Shoveling dirt		
Standing (store clerk)		Sawing wood		
Recreational				
Golf (cart)	Dancing (social)	Tennis (singles)	Canoeing	Squash
Knitting	Golf (walking)	Snow skiing (downhill)	Mountain climbing	Ski touring
	Sailing	Light backpacking		Vigorous basketball
	Tennis (doubles)	Basketball		
		Stream fishing		
Physical Conditioning				
Walking (2 mph)	Level walking (3–4 mph)	Level walking (4.5–5.0 mph)	Level jogging (5 mph)	Running more than 6 mph
Stationary bike	Level biking (6–8 mph)	Bicycling (9–10 mph)	Swimming (crawl stroke)	Bicycling (more than 13 mph)
Very light calisthenics	Light calisthenics	Swimming, breast stroke	Rowing machine	Rope jumping
			Heavy calisthenics	Walking uphill (5 mph)
			Bicycling (12 mph)	

Abbreviation: METs, metabolic equivalent tasks.

Source: Modified from WL Haskell: Rehabilitation of the coronary patient, in NK Wenger, HK Hellerstein (eds): *Design and Implementation of Cardiac Conditioning Program*. New York, Churchill Livingstone, 1978.

The treatment of dyslipidemia is central in aiming for long-term relief from angina, reduced need for revascularization, and reduction in myocardial infarction and death. The control of lipids can be achieved by the combination of a diet low in saturated and *trans*-unsaturated fatty acids, exercise, and weight loss. Nearly always, HMG-CoA reductase inhibitors (statins) are required and can lower LDL cholesterol (25–50%), raise HDL cholesterol (5–9%), and lower triglycerides (5–30%). A powerful treatment effect of statins on atherosclerosis, IHD, and outcomes is seen regardless of the pretreatment LDL cholesterol level. Fibrates or niacin can be used to raise HDL cholesterol and lower triglycerides (**Chaps. 291e and 421**). Controlled trials with lipid-regulating regimens have shown equal proportional benefit for men, women, the elderly, diabetic patients, and smokers.

Compliance with the health-promoting behaviors listed above is generally very poor, and a conscientious physician must not underestimate the major effort required to meet this challenge. Many patients who are discharged from the hospital with proven coronary disease do not receive adequate treatment for dyslipidemia. In light of the proof that treating dyslipidemia brings major benefits, physicians need to establish treatment pathways, monitor compliance, and follow up regularly.

RISK REDUCTION IN WOMEN WITH IHD

The incidence of clinical IHD in premenopausal women is very low; however, after menopause, the atherogenic risk factors increase (e.g., increased LDL, reduced HDL) and the rate of clinical coronary events accelerates to the levels observed in men. Women have not given up cigarette smoking as effectively as have men. Diabetes mellitus, which is more common in women, greatly increases the occurrence of clinical IHD and amplifies the deleterious effects of hypertension, hyperlipidemia, and smoking. Cardiac catheterization and coronary revascularization are underused in women and are performed at a later and more severe stage of the disease than in men. When cholesterol lowering, beta blockers after myocardial infarction, and coronary artery bypass grafting are applied in the appropriate patient groups, women benefit to the same degree as men.

DRUG THERAPY

The commonly used drugs for the treatment of angina pectoris are summarized in **Tables 293-4 through 293-6**. Pharmacotherapy for IHD is designed to reduce the frequency of anginal episodes, myocardial infarction, and coronary death. There is a wealth of trial

data to emphasize how important this medical management is when added to the health-promoting behaviors discussed above. To achieve maximum benefit from medical therapy for IHD, it is frequently necessary to combine agents from different classes and titrate the doses as guided by the individual profile of risk factors, symptoms, hemodynamic responses, and side effects.

NITRATES

The organic nitrates are a valuable class of drugs in the management of angina pectoris (Table 293-4). Their major mechanisms of action include systemic venodilation with concomitant reduction in left ventricular end-diastolic volume and pressure, thereby reducing myocardial wall tension and oxygen requirements; dilation of epicardial coronary vessels; and increased blood flow in collateral vessels. When metabolized, organic nitrates release nitric oxide (NO) that binds to guanylyl cyclase in vascular smooth muscle cells, leading to an increase in cyclic guanosine monophosphate, which causes relaxation of vascular smooth muscle. Nitrates also exert antithrombotic activity by NO-dependent activation of platelet guanylyl cyclase, impairment of intraplatelet calcium flux, and platelet activation.

The absorption of these agents is most rapid and complete through the mucous membranes. For this reason, nitroglycerin is most commonly administered sublingually in tablets of 0.4 or 0.6 mg. Patients with angina should be instructed to take the medication both to relieve angina and also approximately 5 min before stress that is likely to induce an episode. The value of this prophylactic use of the drug cannot be overemphasized.

Nitrates improve exercise tolerance in patients with chronic angina and relieve ischemia in patients with unstable angina as well as patients with Prinzmetal's variant angina (**Chap. 294**). A diary of angina and nitroglycerin use may be valuable for detecting changes in the frequency, severity, or threshold for discomfort that may signify the development of unstable angina pectoris and/or herald an impending myocardial infarction.

Long-Acting Nitrates None of the long-acting nitrates are as effective as sublingual nitroglycerin for the acute relief of angina. These organic nitrate preparations can be swallowed, chewed, or administered as a patch or paste by the transdermal route (Table 293-4). They can provide effective plasma levels for up to 24 h, but the therapeutic response is highly variable. Different preparations and/or administration during the daytime should be tried only to prevent discomfort while avoiding side effects such as headache and dizziness. Individual dose titration is important to prevent side effects. To minimize the effects of tolerance, the minimum effective dose should be used and a minimum of 8 h each day kept free of the drug to restore any useful response(s).

β -Adrenergic Blockers These drugs represent an important component of the pharmacologic treatment of angina pectoris (Table 293-5). They reduce myocardial oxygen demand by inhibiting the increases in heart rate, arterial pressure, and myocardial contractility caused by adrenergic activation. Beta blockade reduces these variables most strikingly during exercise but causes only small reductions at rest. Long-acting beta-blocking drugs or sustained-release formulations offer the advantage of once-daily dosing (Table 293-5). The therapeutic aims include relief of angina and ischemia. These drugs also can reduce mortality and reinfarction rates in patients after myocardial infarction and are moderately effective antihypertensive agents.

Relative contraindications include asthma and reversible airway obstruction in patients with chronic lung disease, atrioventricular conduction disturbances, severe bradycardia, Raynaud's phenomenon, and a history of mental depression. Side effects include fatigue, reduced exercise tolerance, nightmares, impotence, cold extremities, intermittent claudication, bradycardia (sometimes severe), impaired atrioventricular conduction, left ventricular failure, bronchial asthma, worsening claudication, and intensification of the hypoglycemia produced by oral hypoglycemic agents and insulin.

TABLE 293-4 NITRATE THERAPY IN PATIENTS WITH ISCHEMIC HEART DISEASE

Preparation of Agent	Dose	Schedule
Nitroglycerin ^a		
Ointment	0.5–2 inches	Two or three times daily
Transdermal patch	0.2–0.8 mg/h	Every 24 h; remove at bedtime for 12–14 h
Sublingual tablet	0.3–0.6 mg	As needed, up to three doses 5 min apart
Spray	One or two sprays	As needed, up to three doses 5 min apart
Isosorbide dinitrate ^a		
Oral	10–40 mg	Two or three times daily
Oral sustained release	80–120 mg	Once or twice daily (eccentric schedules)
Isosorbide 5-mononitrate		
Oral	20 mg	Twice daily (given 7–8 h apart)
Oral sustained release	30–240 mg	Once daily

^aA 10- to 12-h nitrate-free interval is recommended.

Source: Modified from DA Morrow, WE Boden: Stable ischemic heart disease. In RO Bonow et al (eds): *Braunwald's Heart Disease: A Textbook of Cardiovascular Medicine*. 9th edition. Philadelphia, Saunders, 2012, p. 1224.

TABLE 293-5 PROPERTIES OF BETA BLOCKERS IN CLINICAL USE FOR ISCHEMIC HEART DISEASE

Drugs	Selectivity	Partial Agonist Activity	Usual Dose for Angina
Acebutolol	β_1	Yes	200–600 mg twice daily
Atenolol	β_1	No	50–200 mg/d
Betaxolol	β_1	No	10–20 mg/d
Bisoprolol	β_1	No	10 mg/d
Esmolol (intravenous) ^a	β_1	No	50–300 μ g/kg/min
Labetalol ^b	None	Yes	200–600 mg twice daily
Metoprolol	β_1	No	50–200 mg twice daily
Nadolol	None	No	40–80 mg/d
Nebivolol	β_1 (at low doses)	No	5–40 mg/d
Pindolol	None	Yes	2.5–7.5 mg 3 times daily
Propranolol	None	No	80–120 mg twice daily
Timolol	None	No	10 mg twice daily

^aEsmolol is an ultra-short-acting beta blocker that is administered as a continuous intravenous infusion. Its rapid offset of action makes esmolol an attractive agent to use in patients with relative contraindications to beta blockade. ^bLabetalol is a combined alpha and beta blocker.

Note: This list of beta blockers that may be used to treat patients with angina pectoris is arranged alphabetically. The agents for which there is the greatest clinical experience include atenolol, metoprolol, and propranolol. It is preferable to use a sustained-release formulation that may be taken once daily to improve the patient's compliance with the regimen.

Source: Modified from RJ Gibbons et al: *J Am Coll Cardiol* 41:159, 2003.

Reducing the dose or even discontinuation may be necessary if these side effects develop and persist. Since sudden discontinuation can intensify ischemia, the doses should be tapered over 2 weeks. Beta blockers with relative β_1 -receptor specificity such as metoprolol and atenolol may be preferable in patients with mild bronchial obstruction and insulin-requiring diabetes mellitus.

Calcium Channel Blockers Calcium channel blockers (Table 293-6) are coronary vasodilators that produce variable and dose-dependent reductions in myocardial oxygen demand, contractility, and arterial pressure. These combined pharmacologic effects are advantageous and make these agents as effective as beta blockers in the treatment of angina pectoris. They are indicated when beta blockers are contraindicated, poorly tolerated, or ineffective. Because of differences in the dose-response relationship on cardiac electrical activity between the dihydropyridine and nondihydropyridine calcium channel blockers, verapamil and diltiazem may produce symptomatic disturbances in cardiac conduction and bradyarrhythmias. They also exert negative inotropic actions and are more

likely to aggravate left ventricular failure, particularly when used in patients with left ventricular dysfunction, especially if the patients are also receiving beta blockers. Although useful effects usually are achieved when calcium channel blockers are combined with beta blockers and nitrates, individual titration of the doses is essential with these combinations. Variant (Prinzmetal's) angina responds particularly well to calcium channel blockers (especially members of the dihydropyridine class), supplemented when necessary by nitrates (**Chap. 294**).

Verapamil ordinarily should not be combined with beta blockers because of the combined adverse effects on heart rate and contractility. Diltiazem can be combined with beta blockers in patients with normal ventricular function and no conduction disturbances. Amlodipine and beta blockers have complementary actions on coronary blood supply and myocardial oxygen demands. Whereas the former decreases blood pressure and dilates coronary arteries, the latter slows heart rate and decreases contractility. Amlodipine and the other second-generation dihydropyridine calcium antagonists (nicardipine, isradipine, long-acting nifedipine, and felodipine) are

TABLE 293-6 CALCIUM CHANNEL BLOCKERS IN CLINICAL USE FOR ISCHEMIC HEART DISEASE

Drugs	Usual Dose	Duration of Action	Side Effects
Dihydropyridines			
Amlodipine	5–10 mg qd	Long	Headache, edema
Felodipine	5–10 mg qd	Long	Headache, edema
Isradipine	2.5–10 mg bid	Medium	Headache, fatigue
Nicardipine	20–40 mg tid	Short	Headache, dizziness, flushing, edema
Nifedipine	Immediate release: ^a 30–90 mg daily orally Slow release: 30–180 mg orally	Short	Hypotension, dizziness, flushing, nausea, constipation, edema
Nisoldipine	20–40 mg qd	Short	Similar to nifedipine
Nondihydropyridines			
Diltiazem	Immediate release: 30–80 mg 4 times daily Slow release: 120–320 mg qd	Short Long	Hypotension, dizziness, flushing, bradycardia, edema
Verapamil	Immediate release: 80–160 mg tid Slow release: 120–480 mg qd	Short Long	Hypotension, myocardial depression, heart failure, edema, bradycardia

^aMay be associated with increased risk of mortality if administered during acute myocardial infarction.

Note: This list of calcium channel blockers that may be used to treat patients with angina pectoris is divided into two broad classes, dihydropyridines and nondihydropyridines, and arranged alphabetically within each class. Among the dihydropyridines, the greatest clinical experience has been obtained with amlodipine and nifedipine. After the initial period of dose titration with a short-acting formulation, it is preferable to switch to a sustained-release formulation that may be taken once daily to improve patient compliance with the regimen.

Source: Modified from RJ Gibbons et al: *J Am Coll Cardiol* 41:159, 2003.

1590 potent vasodilators and are useful in the simultaneous treatment of angina and hypertension. Short-acting dihydropyridines should be avoided because of the risk of precipitating infarction, particularly in the absence of concomitant beta blocker therapy.

Choice Between Beta Blockers and Calcium Channel Blockers for Initial Therapy Since beta blockers have been shown to improve life expectancy after acute myocardial infarction (**Chaps. 294 and 295**) and calcium channel blockers have not, the former may also be preferable in patients with angina and a damaged left ventricle. However, calcium channel blockers are indicated in patients with the following: (1) inadequate responsiveness to the combination of beta blockers and nitrates; many of these patients do well with a combination of a beta blocker and a dihydropyridine calcium channel blocker; (2) adverse reactions to beta blockers such as depression, sexual disturbances, and fatigue; (3) angina and a history of asthma or chronic obstructive pulmonary disease; (4) sick-sinus syndrome or significant atrioventricular conduction disturbances; (5) Prinzmetal's angina; or (6) symptomatic peripheral arterial disease.

Antiplatelet Drugs Aspirin is an irreversible inhibitor of platelet cyclooxygenase and thereby interferes with platelet activation. Chronic administration of 75–325 mg orally per day has been shown to reduce coronary events in asymptomatic adult men over age 50, patients with chronic stable angina, and patients who have or have survived unstable angina and myocardial infarction. There is a dose-dependent increase in bleeding when aspirin is used chronically. It is preferable to use an enteric-coated formulation in the range of 81–162 mg/d. Administration of this drug should be considered in all patients with IHD in the absence of gastrointestinal bleeding, allergy, or dyspepsia. Clopidogrel (300–600 mg loading and 75 mg/d) is an oral agent that blocks P2Y₁₂ ADP receptor-mediated platelet aggregation. It provides benefits similar to those of aspirin in patients with stable chronic IHD and may be substituted for aspirin if aspirin causes the side effects listed above. Clopidogrel combined with aspirin reduces death and coronary ischemic events in patients with an acute coronary syndrome (**Chap. 294**) and also reduces the risk of thrombus formation in patients undergoing implantation of a stent in a coronary artery (**Chap. 296e**). Alternative antiplatelet agents that block the P2Y₁₂ platelet receptor such as prasugrel and ticagrelor have been shown to be more effective than clopidogrel for prevention of ischemic events after placement of a stent for an acute coronary syndrome but are associated with an increased risk of bleeding. Although combined treatment with clopidogrel and aspirin for at least a year is recommended in patients with an acute coronary syndrome treated with implantation of a drug-eluting stent, studies have not shown any benefit from the routine addition of clopidogrel to aspirin in patients with chronic stable IHD.

OTHER THERAPIES

The angiotensin-converting enzyme (ACE) inhibitors are widely used in the treatment of survivors of myocardial infarction, patients with hypertension or chronic IHD including angina pectoris, and those at high risk of vascular diseases such as diabetes. The benefits of ACE inhibitors are most evident in IHD patients at increased risk, especially if diabetes mellitus or left ventricle dysfunction is present, and those who have not achieved adequate control of blood pressure and LDL cholesterol on beta blockers and statins. However, the routine administration of ACE inhibitors to IHD patients who have normal left ventricular function and have achieved blood pressure and LDL goals on other therapies does not reduce the incidence of events and therefore is not cost-effective.

Despite treatment with nitrates, beta blockers, or calcium channel blockers, some patients with IHD continue to experience angina, and additional medical therapy is now available to alleviate their symptoms. Ranolazine, a piperazine derivative, may be useful for patients with chronic angina despite standard medical therapy. Its antianginal action is believed to occur via inhibition of the late inward sodium current (I_{Na}). The benefits of I_{Na} inhibition include

limitation of the Na overload of ischemic myocytes and prevention of Ca^{2+} overload via the Na^+-Ca^{2+} exchanger. A dose of 500–1000 mg orally twice daily is usually well tolerated. Ranolazine is contraindicated in patients with hepatic impairment or with conditions or drugs associated with QT_c prolongation and when drugs that inhibit the CYP3A metabolic system (e.g., ketoconazole, diltiazem, verapamil, macrolide antibiotics, HIV protease inhibitors, and large quantities of grapefruit juice) are being used.

Nonsteroidal anti-inflammatory drug (NSAID) use in patients with IHD may be associated with a small but finite increased risk of myocardial infarction and mortality. For this reason, they generally should be avoided in IHD patients. If they are required for symptom relief, it is advisable to coadminister aspirin and strive to use an NSAID associated with the lowest risk of cardiovascular events, in the lowest dose required, and for the shortest period of time.

Another class of agents opens ATP-sensitive potassium channels in myocytes, leading to a reduction of free intracellular calcium ions. The major drug in this class is nicorandil, which typically is administered orally in a dose of 20 mg twice daily for prevention of angina. (Nicorandil is not available for use in the United States but is used in several other countries.)

Angina and Heart Failure Transient left ventricular failure with angina can be controlled by the use of nitrates. For patients with established congestive heart failure, the increased left ventricular wall tension raises myocardial oxygen demand. Treatment of congestive heart failure with an ACE inhibitor, a diuretic, and digoxin (**Chap. 279**) reduces heart size, wall tension, and myocardial oxygen demand, which helps control angina and ischemia. If the symptoms and signs of heart failure are controlled, an effort should be made to use beta blockers not only for angina but because trials in heart failure have shown significant improvement in survival. A trial of the intravenous ultra-short-acting beta blocker esmolol may be useful to establish the safety of beta blockade in selected patients. Nocturnal angina often can be relieved by the treatment of heart failure.

The combination of congestive heart failure and angina in patients with IHD usually indicates a poor prognosis and warrants serious consideration of cardiac catheterization and coronary revascularization.

CORONARY REVASCULARIZATION

Clinical trials have confirmed that with the initial diagnosis of stable IHD, it is first appropriate to initiate a thorough medical regimen as described above. Revascularization should be considered in the presence of unstable phases of the disease, intractable symptoms, severe ischemia or high-risk coronary anatomy, diabetes, and impaired left ventricular (LV) function. *Revascularization should be employed in conjunction with but not replace the continuing need to modify risk factors and assess medical therapy.* An algorithm for integrating medical therapy and revascularization options in patients with IHD is shown in **Fig. 293-3**.

PERCUTANEOUS CORONARY INTERVENTION

(**See also Chap. 296e**) Percutaneous coronary intervention (PCI) involving balloon dilatation usually accompanied by coronary stenting is widely used to achieve revascularization of the myocardium in patients with symptomatic IHD and suitable stenoses of epicardial coronary arteries. Whereas patients with stenosis of the left main coronary artery and those with three-vessel IHD (especially with diabetes and/or impaired LV function) who require revascularization are best treated with CABG, PCI is widely employed in patients with symptoms and evidence of ischemia due to stenoses of one or two vessels and even in selected patients with three-vessel disease (and, perhaps, in some patients with left main disease) and may offer many advantages over surgery.

Indications and Patient Selection The most common clinical indication for PCI is symptom-limiting angina pectoris, despite medical therapy, accompanied by evidence of ischemia during a stress test. PCI is more effective than medical therapy for the relief of angina. PCI improves

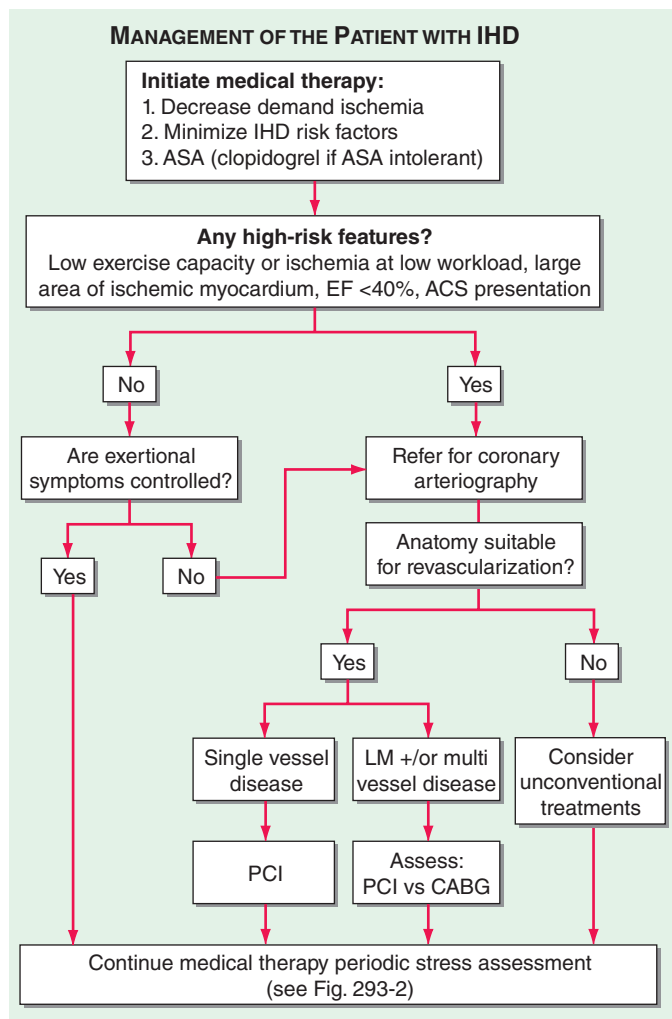


FIGURE 293-3 Algorithm for management of a patient with ischemic heart disease. All patients should receive the core elements of medical therapy as shown at the top of the algorithm. If high-risk features are present, as established by the clinical history, exercise test data, and imaging studies, the patient should be referred for coronary arteriography. Based on the number and location of the diseased vessels and their suitability for revascularization, the patient is treated with a percutaneous coronary intervention (PCI) or coronary artery bypass graft (CABG) surgery or should be considered for unconventional treatments. See text for further discussion. ACS, acute coronary syndrome; ASA, aspirin; EF, ejection fraction; IHD, ischemic heart disease; LM, left main.

outcomes in patients with unstable angina or when used early in the course of myocardial infarction with and without cardiogenic shock. However, in patients with stable exertional angina, clinical trials have confirmed that PCI does not reduce the occurrence of death or myocardial infarction compared to optimum medical therapy. PCI can be used to treat stenoses in native coronary arteries as well as in bypass grafts in patients who have recurrent angina after CABG.

Risks When coronary stenoses are discrete and symmetric, two and even three vessels can be treated in sequence. However, case selection is essential to avoid a prohibitive risk of complications, which are usually due to dissection or thrombosis with vessel occlusion, uncontrolled ischemia, and ventricular failure (Chap. 296e). Oral aspirin, a P2Y12 antagonist, and an antithrombin agent are given to reduce coronary thrombus formation. Left main coronary artery stenosis generally is regarded as a contraindication to PCI; such patients should be treated with CABG. In selected cases such as patients with prohibitive surgical risks, PCI of an unprotected left main can be considered, but such a procedure should be performed only by a highly skilled

operator; importantly, there are regional differences in the use of this approach internationally.

Efficacy Primary success, i.e., adequate dilation (an increase in luminal diameter >20% to a residual diameter obstruction <50%) with relief of angina, is achieved in >95% of cases. Recurrent stenosis of the dilated vessels occurs in ~20% of cases within 6 months of PCI with bare metal stents, and angina will recur within 6 months in 10% of cases. Restenosis is more common in patients with diabetes mellitus, arteries with small caliber, incomplete dilation of the stenosis, long stents, occluded vessels, obstructed vein grafts, dilation of the left anterior descending coronary artery, and stenoses containing thrombi. In diseased vein grafts, procedural success has been improved by the use of capture devices or filters that prevent embolization, ischemia, and infarction.

It is usual clinical practice to administer aspirin indefinitely and a P2Y12 antagonist for 1–3 months after the implantation of a bare metal stent. Although aspirin in combination with a thienopyridine may help prevent coronary thrombosis during and shortly after PCI with stenting, there is no evidence that these medications reduce the incidence of restenosis.

The use of drug-eluting stents that locally deliver antiproliferative drugs can reduce restenosis to less than 10%. Advances in PCI, especially the availability of drug-eluting stents, have vastly extended the use of this revascularization option in patients with IHD. Of note, however, the delayed endothelial healing in the region of a drug-eluting stent also extends the period during which the patient is at risk for subacute stent thrombosis. Current recommendations are to administer aspirin indefinitely and a P2Y12 antagonist daily for at least 1 year after implantation of a drug-eluting stent. When a situation arises in which temporary discontinuation of antiplatelet therapy is necessary, the clinical circumstances should be reviewed with the operator who performed the PCI and a coordinated plan should be established for minimizing the risk of late stent thrombus; central to this plan is the discontinuation of antiplatelet therapy for the shortest acceptable period. The risk of stent thrombosis is dependent on stent size and length, complexity of the lesions, age, diabetes, and technique. However, compliance with dual antiplatelet therapy and individual responsiveness to platelet inhibition are very important factors as well.

Successful PCI produces effective relief of angina in >95% of cases. The majority of patients with symptomatic IHD who require revascularization can be treated initially by PCI. Successful PCI is less invasive and expensive than CABG and permits savings in the *initial* cost of care. Successful PCI avoids the risk of stroke associated with CABG surgery, allows earlier return to work, and allows the resumption of an active life. However, the early health-related and economic benefit of PCI is reduced over time because of the greater need for follow-up and the increased need for repeat procedures. When directly compared in patients with diabetes or three-vessel or left main CAD, CABG was superior to PCI in preventing major adverse cardiac or cerebrovascular events over a 12-month follow-up.

CORONARY ARTERY BYPASS GRAFTING

Anastomosis of one or both of the internal mammary arteries or a radial artery to the coronary artery distal to the obstructive lesion is the preferred procedure. For additional obstructions that cannot be bypassed by an artery, a section of a vein (usually the saphenous) is used to form a connection between the aorta and the coronary artery distal to the obstructive lesion.

Although some indications for CABG are controversial, certain areas of agreement exist:

1. The operation is relatively safe, with mortality rates <1% in patients without serious comorbid disease and normal LV function and when the procedure is performed by an experienced surgical team.
2. Intraoperative and postoperative mortality rates increase with the severity of ventricular dysfunction, comorbidities, age >80 years, and lack of surgical experience. The effectiveness and risk of CABG vary widely depending on case selection and the skill and experience of the surgical team.

3. Occlusion of *venous* grafts is observed in 10–20% of patients during the first postoperative year and in approximately 2% per year during 5- to 7-year follow-up and 4% per year thereafter. Long-term patency rates are considerably higher for internal mammary and radial artery implantations than for saphenous vein grafts. In patients with left anterior descending coronary artery obstruction, survival is better when coronary bypass involves the internal mammary artery rather than a saphenous vein. Graft patency and outcomes are improved by meticulous treatment of risk factors, particularly dyslipidemia.
4. Angina is abolished or greatly reduced in ~90% of patients after complete revascularization. Although this usually is associated with graft patency and restoration of blood flow, the pain may also have been alleviated as a result of infarction of the ischemic segment or a placebo effect. Within 3 years, angina recurs in about one-fourth of patients but is rarely severe.
5. Survival may be improved by operation in patients with stenosis of the left main coronary artery as well as in patients with three- or two-vessel disease with significant obstruction of the proximal left anterior descending coronary artery. The survival benefit is greater in patients with abnormal LV function (ejection fraction <50%). Survival *may* also be improved in the following patients: (a) patients with obstructive CAD who have survived sudden cardiac death or sustained ventricular tachycardia; (b) patients who have undergone previous CABG and have multiple saphenous vein graft stenoses, especially of a graft supplying the left anterior descending coronary artery; and (c) patients with recurrent stenosis after PCI and high-risk criteria on noninvasive testing.
6. Minimally invasive CABG through a small thoracotomy and/or off-pump surgery can reduce morbidity and shorten convalescence in suitable patients but does not appear to reduce significantly the risk of neurocognitive dysfunction postoperatively.
7. Among patients with type 2 diabetes mellitus and multivessel coronary disease, CABG surgery plus optimal medical therapy is superior to optimal medical therapy alone in preventing major cardiovascular events, a benefit mediated largely by a significant reduction in nonfatal myocardial infarction. The benefits of CABG are especially evident in diabetic patients treated with an insulin-sensitizing strategy as opposed to an insulin-providing strategy. CABG has also been shown to be superior to PCI (including the use of drug-eluting stents) in preventing death, myocardial infarction, and repeat revascularization in patients with diabetes mellitus and multivessel IHD.

Indications for CABG usually are based on the severity of symptoms, coronary anatomy, and ventricular function. The ideal candidate is male, <80 years of age, has no other complicating disease, and has troublesome or disabling angina that is not adequately controlled by medical therapy or does not tolerate medical therapy. The patient wishes to lead a more active life and has severe stenoses of two or three epicardial coronary arteries with objective evidence of myocardial ischemia as a cause of the chest discomfort. Great symptomatic benefit can be anticipated in such patients. Congestive heart failure and/or LV dysfunction, advanced age (>80 years), reoperation, urgent need for surgery, and the presence of diabetes mellitus are all associated with a higher perioperative mortality rate.

LV dysfunction can be due to noncontractile or hypocontractile segments that are viable but are chronically ischemic (hibernating myocardium). As a consequence of chronic reduction in myocardial blood flow, these segments downregulate their contractile function. They can be detected by using radionuclide scans of myocardial perfusion and metabolism, PET, cardiac MRI, or delayed scanning with thallium-201 or by improvement of regional functional impairment provoked by low-dose dobutamine. In such patients, revascularization improves myocardial blood flow, can return function, and can improve survival.

The Choice Between PCI and CABG All the clinical characteristics of each individual patient must be used to decide on the method of revascularization (e.g., LV function, diabetes, lesion complexity). A number of randomized clinical trials have compared PCI and CABG in patients

with multivessel CAD who were suitable technically for both procedures. The redevelopment of angina requiring repeat coronary angiography and repeat revascularization is higher with PCI. This is a result of restenosis in the stented segment (a problem largely solved with drug-eluting stents) and the development of new stenoses in unstented portions of the coronary vasculature. It has been argued that PCI with stenting focuses on culprit lesions, whereas a bypass graft to the target vessel also provides a conduit around future culprit lesions proximal to the anastomosis of the graft to the native vessel (Fig. 293-4). By contrast, stroke rates are lower with PCI.

Based on available evidence, it is now recommended that patients with an unacceptable level of angina despite optimal medical management be considered for coronary revascularization. Patients with single- or two-vessel disease with normal LV function and anatomically suitable lesions ordinarily are advised to undergo PCI (Chap. 296e). Patients with three-vessel disease (or two-vessel disease that includes the proximal left descending coronary artery) and impaired global LV function (LV ejection fraction <50%) or diabetes mellitus and those with left main CAD or other lesions unsuitable for catheter-based procedures should be considered for CABG as the initial method of revascularization. In light of the complexity of the decision making, it is desirable to have a multidisciplinary team, including a cardiologist and a cardiac surgeon in conjunction with the patient's primary care physician, provide input along with ascertaining the patient's preferences before committing to a particular revascularization option.

UNCONVENTIONAL TREATMENTS FOR IHD

On occasion clinicians will encounter a patient who has persistent disabling angina despite maximally tolerated medical therapy and for whom revascularization is not an option (e.g., small diffusely diseased vessels not amenable to stent implantation or acceptable targets for bypass grafting). In such situations, unconventional treatments should be considered.

Enhanced external counterpulsation utilizes pneumatic cuffs on the lower extremities to provide diastolic augmentation and systolic unloading of blood pressure to decrease cardiac work and oxygen consumption while enhancing coronary blood flow. Clinical trials have shown that regular application improves angina, exercise capacity, and regional myocardial perfusion. Experimental approaches such as gene and stem cell therapies are also under active study.

ASYMPTOMATIC (SILENT) ISCHEMIA

Obstructive CAD, acute myocardial infarction, and transient myocardial ischemia are frequently asymptomatic. During continuous ambulatory ECG monitoring, the majority of ambulatory patients with typical chronic stable angina are found to have objective evidence of myocardial ischemia (ST-segment depression) during episodes of chest discomfort while they are active outside the hospital. In addition, many of these patients also have more frequent episodes of asymptomatic ischemia. Frequent episodes of ischemia (symptomatic and asymptomatic) during daily life appear to be associated with an increased likelihood of adverse coronary events (death and myocardial infarction). In addition, patients with asymptomatic ischemia after a myocardial infarction are at greater risk for a second coronary event. The widespread use of exercise ECG during routine examinations has also identified some of these previously unrecognized patients with asymptomatic CAD. Longitudinal studies have demonstrated an increased incidence of coronary events in asymptomatic patients with positive exercise tests.

TREATMENT ASYMPTOMATIC ISCHEMIA

The management of patients with asymptomatic ischemia must be individualized. When coronary disease has been confirmed, the aggressive treatment of hypertension and dyslipidemia is essential and will decrease the risk of infarction and death. In addition, the

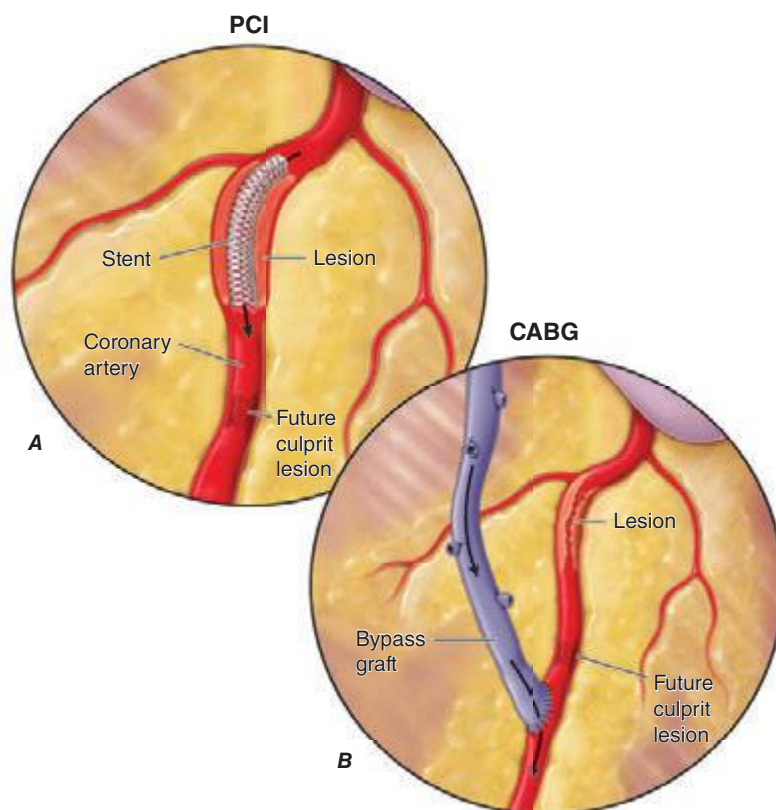


FIGURE 293-4 Difference in the approach to the lesion with percutaneous coronary intervention (PCI) and coronary artery bypass grafting (CABG). PCI is targeted at the “culprit” lesion or lesions, whereas CABG is directed at the epicardial vessel, including the culprit lesion or lesions and future culprits, proximal to the insertion of the vein graft, a difference that may account for the superiority of CABG, at least in the intermediate term, in patients with multivessel disease. (Reproduced from BJ Gersh, RL Frye: *N Engl J Med* 352:2235, 2005.)

physician should consider the following: (1) the degree of positivity of the stress test, particularly the stage of exercise at which ECG signs of ischemia appear; the magnitude and number of the ischemic zones of myocardium on imaging; and the change in LV ejection fraction that occurs on radionuclide ventriculography or echocardiography during ischemia and/or during exercise; (2) the ECG leads showing a positive response, with changes in the anterior precordial leads indicating a less favorable prognosis than changes in the inferior leads; and (3) the patient’s age, occupation, and general medical condition.

Most would agree that an asymptomatic 45-year-old commercial airline pilot with significant (0.4-mV) ST-segment depression in leads V_1 to V_4 during mild exercise should undergo coronary arteriography, whereas an asymptomatic, sedentary 85-year-old retiree with 0.1-mV ST-segment depression in leads II and III during maximal activity need not. However, there is no consensus about the most appropriate approach in the large majority of patients for whom the situation is less extreme. Asymptomatic patients with silent ischemia, three-vessel CAD, and impaired LV function may be considered appropriate candidates for CABG.

The treatment of risk factors, particularly lipid lowering and blood pressure control as described above, and the use of aspirin, statins, and beta blockers after infarction have been shown to reduce events and improve outcomes in asymptomatic as well as symptomatic patients with ischemia and proven CAD. Although the incidence of asymptomatic ischemia can be reduced by treatment with beta blockers, calcium channel blockers, and long-acting nitrates, it is not clear whether this is necessary or desirable in patients who have not had a myocardial infarction.

294 Non-ST-Segment Elevation Acute Coronary Syndrome (Non-ST-Segment Elevation Myocardial Infarction and Unstable Angina)

Christopher P. Cannon, Eugene Braunwald

Patients with ischemic heart disease fall into two large groups: patients with chronic coronary artery disease (CAD) who most commonly present with stable angina (Chap. 293) and patients with acute coronary syndromes (ACSs). These include patients with acute myocardial infarction with ST-segment elevation (STEMI) on their presenting electrocardiogram (Chap. 295) and those with non-ST-segment elevation acute coronary syndrome (NSTEMI). The latter include patients with non-ST-segment elevation myocardial infarction (NSTEMI), who, by definition, have evidence of myocyte necrosis, and those with unstable angina (UA), who do not. The relative incidence of NSTEMI compared to STEMI appears to be increasing (Fig. 294-1). Every year in the United States, approximately 1.1 million patients are admitted to hospitals with NSTEMI-ACS as compared with ~300,000 patients with acute STEMI. Women comprise more than one-third of patients with NSTEMI-ACS, but less than one-fourth of patients with STEMI.

PATHOPHYSIOLOGY

NSTEMI-ACS is most commonly caused by an imbalance between oxygen supply and oxygen demand resulting from a partially occluding thrombus forming on a disrupted atherothrombotic coronary plaque

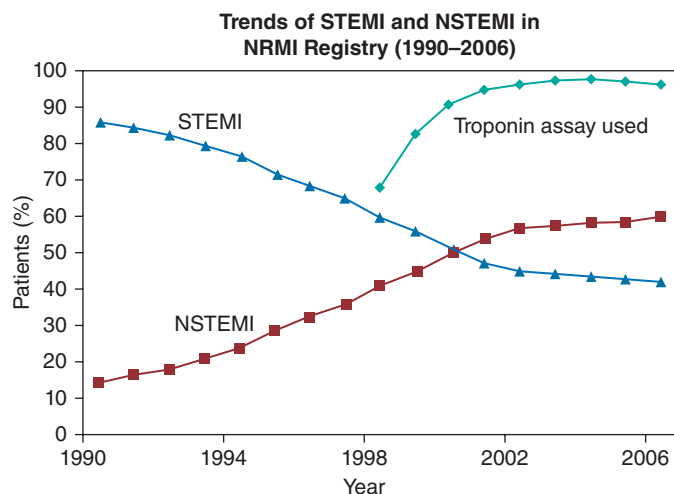


FIGURE 294-1 Trends of incidence of ST-segment elevation myocardial infarction (STEMI) and non-ST-segment elevation myocardial infarction (NSTEMI) and of frequency of use of troponin assay to diagnose acute myocardial infarction. NRM, National Registry of Myocardial Infarction. (From N Arora, RG Brindis, CP Cannon: *Acute coronary syndrome in North America*, in Theroux P [ed]: *Acute Coronary Syndromes*, 2nd ed. Philadelphia: Elsevier, 2011.)

or on eroded coronary artery endothelium. Severe ischemia or myocardial necrosis may occur consequent to the reduction of coronary blood flow caused by the thrombus and by downstream embolization of platelet aggregates and/or atherosclerotic debris. Other causes of NSTEMI-ACS include: (1) dynamic obstruction (e.g., coronary spasm, as in Prinzmetal's variant angina [see "Prinzmetal's Variant Angina" later]); (2) severe mechanical obstruction due to progressive coronary atherosclerosis; and (3) increased myocardial oxygen demand produced by conditions such as fever, tachycardia, and thyrotoxicosis in the presence of fixed epicardial coronary obstruction. More than one of these processes may be involved.

Among patients with NSTEMI-ACS studied at angiography, approximately 10% have stenosis of the left main coronary artery, 35% have three-vessel CAD, 20% have two-vessel disease, 20% have single-vessel disease, and 15% have no apparent critical epicardial coronary artery stenosis; some of the latter may have obstruction of the coronary microcirculation and/or spasm. The "culprit lesion" responsible for ischemia may show an eccentric stenosis with scalloped or overhanging edges and a narrow neck on coronary angiography. Optical coherence tomography (an invasive technique) and contrast-enhanced coronary computed tomographic angiography (CCTA), a noninvasive technique (Fig. 294-2), have shown that culprit lesions are composed of a lipid-rich core with a thin fibrous cap. Patients with NSTEMI-ACS frequently have multiple such plaques that are at risk of disruption (vulnerable plaques).

CLINICAL PRESENTATION

Diagnosis The diagnosis of NSTEMI-ACS is based largely on the clinical presentation. Typically, chest discomfort is severe and has at least one of three features: (1) it occurs at rest (or with minimal exertion), lasting >10 minutes; (2) it is of relatively recent onset (i.e., within the prior 2 weeks); and/or (3) it occurs with a crescendo pattern (i.e., distinctly more severe, prolonged, or frequent than previous episodes). The diagnosis of NSTEMI is established if a patient with these clinical features develops evidence of myocardial necrosis, as reflected in abnormally elevated levels of biomarkers of cardiac necrosis (see below).

History and Physical Examination The chest discomfort, often severe enough to be described as frank pain, is typically located in the substernal region or sometimes in the epigastrium, and radiates to the left



FIGURE 294-2 Coronary computed tomographic angiogram showing an obstructive plaque in the right coronary artery. (From PJ de Feyter, K Nieman. *Multislice computed tomography in acute coronary syndromes*, in Theroux P [ed]: *Acute Coronary Syndromes*, 2nd ed. Philadelphia: Elsevier, 2011.)

arm, left shoulder, and/or neck. Anginal "equivalents" such as dyspnea, epigastric discomfort, nausea, or weakness may occur instead of chest pain and appear to be more frequent in women, the elderly, and patients with diabetes mellitus. The physical examination resembles that in patients with stable angina (Chap. 293) and may be unremarkable. If the patient has a large area of myocardial ischemia or a large NSTEMI, the physical findings can include diaphoresis; pale, cool skin; sinus tachycardia; a third and/or fourth heart sound; basilar rales; and, sometimes, hypotension.

Electrocardiogram ST-segment depression occurs in 20 to 25% of patients; it may be transient in patients without biomarker evidence of myocardial necrosis, but may be persistent for several days in NSTEMI. T-wave changes are common but are less specific signs of ischemia, unless they are new and deep T-wave inversions (≥ 0.3 mV).

Cardiac Biomarkers Patients with NSTEMI have elevated biomarkers of necrosis, such as cardiac troponin I or T, which are specific, sensitive, and the preferred markers of myocardial necrosis. The MB isoform of creatine kinase (CK-MB) is a less sensitive alternative. Elevated levels of these markers distinguish patients with NSTEMI from those with UA. There is a characteristic temporal rise and fall of the plasma concentration of these markers and a direct relationship between the degree of elevation and mortality (see Fig. 294-4B). However, in patients without a clear clinical history of myocardial ischemia, minor cardiac troponin (cTn) elevations have been reported and can be caused by congestive heart failure, myocarditis, or pulmonary embolism, or using high-sensitivity assays, they may occur in ostensibly normal subjects. Thus, in patients with an unclear history, small elevations of cTn, especially if they are persistent, may not be diagnostic of an ACS.

With more widespread measurement of troponin, especially using high-sensitivity assays, an increasing fraction of patients with NSTEMI-ACS are found to have NSTEMI, whereas the fraction of patients with UA is dwindling.

DIAGNOSTIC EVALUATION

In addition to the clinical examination, three major noninvasive tools are used in the evaluation of NSTEMI-ACS: the electrocardiogram (ECG), cardiac biomarkers, and stress testing. CCTA is an additional emerging option (Fig. 294-2). The goals are to: (1) recognize or exclude myocardial infarction (MI) using cardiac biomarkers, preferably cTn;

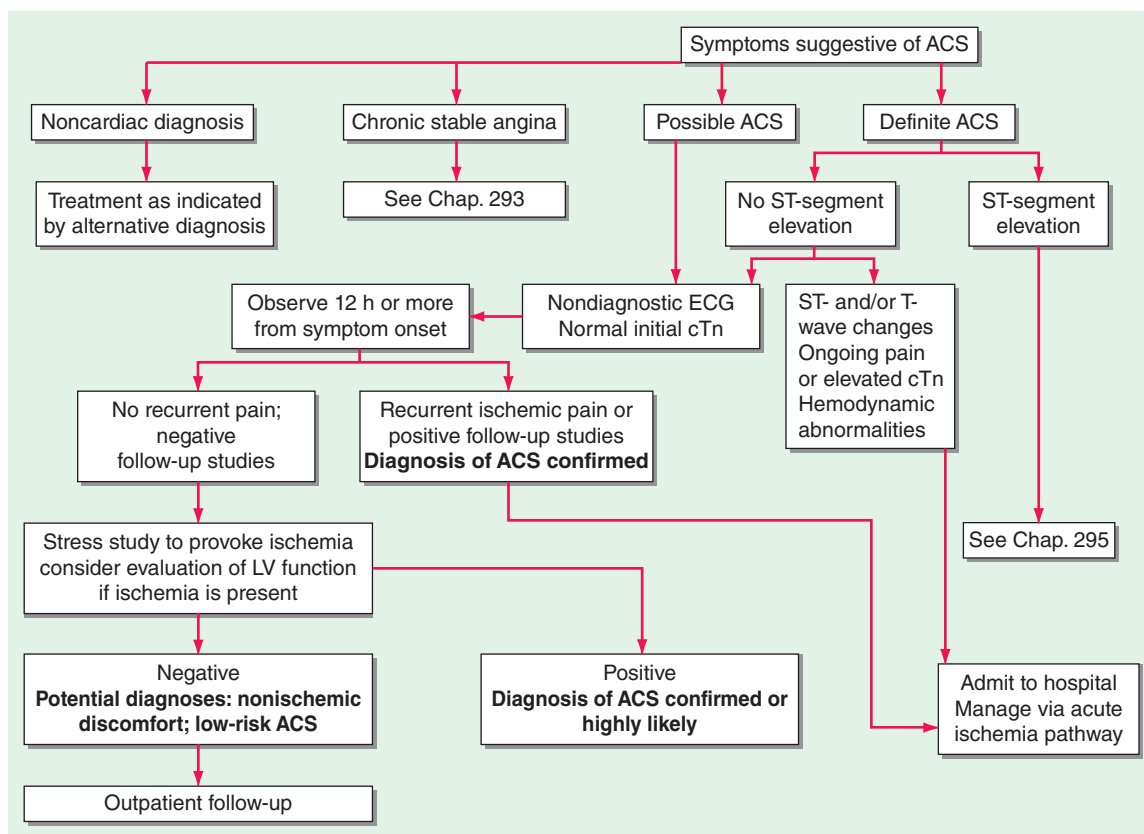


FIGURE 294-3 Algorithm for evaluation and management of patients with suspected acute coronary syndrome (ACS). Follow-up studies refer to ST deviations and elevation of troponin levels. cTn, cardiac troponin; ECG, electrocardiogram; LV, left ventricular. (Modified from JL Anderson et al: *J Am Coll Cardiol* 61:e179, 2013.)

(2) detect rest ischemia (using serial or continuous ECGs); and (3) detect significant coronary obstruction at rest with CCTA and myocardial ischemia using stress testing (Chap 270e).

Patients with a low likelihood of ischemia are usually managed with an emergency department–based critical pathway (which, in some institutions, is carried out in a “chest pain unit”) (Fig. 294-3). Evaluation of such patients includes clinical monitoring for recurrent ischemic discomfort and continuous monitoring of ECGs and cardiac markers, typically obtained at baseline and at 4–6 h and 12 h after presentation. If new elevations in cardiac markers or ST-T-wave changes on the ECG are noted, the patient should be admitted to the hospital. Patients who remain pain free with negative markers may proceed to stress testing to determine the presence of ischemia or CCTA to detect coronary luminal obstruction (Fig. 294-2).

RISK STRATIFICATION

Patients with documented NSTEMI-ACS exhibit a wide spectrum of early (30 days) risk of death, ranging from 1 to 10%, and a recurrent ACS rate of 5 to 15% during the first year. Assessment of risk can be accomplished by clinical risk scoring systems such as that developed from the Thrombolysis in Myocardial Infarction (TIMI) Trials, which includes seven independent risk factors (Fig. 294-4A). The presence of an abnormally elevated cTn is especially important, as is its peak level, which correlates with the extent of myocardial damage (Fig. 294-4B). Other risk factors include diabetes mellitus, left ventricular dysfunction, renal dysfunction, and elevated levels of B-type natriuretic peptides and C-reactive protein. Multimarker strategies are now gaining favor, both to define more fully the pathophysiologic mechanisms underlying a given patient’s presentation and to stratify the patient’s risk further. Patients with ACS without elevated levels of cTn (infrequently encountered with the new sensitive troponin assays) are considered to have UA and have a more favorable prognosis than those with cTn elevations (NSTEMI).

Early risk assessment is useful both in predicting the risk of recurrent cardiac events and in identifying patients who would derive the greatest benefit from an early invasive strategy. For example, in the TACTICS-TIMI 18 Trial, an early invasive strategy conferred a 40% reduction in recurrent cardiac events in patients with an elevated cTn level, whereas no benefit was observed in those without detectable troponin.

TREATMENT NON-ST-SEGMENT ELEVATION ACUTE CORONARY SYNDROME (NON-ST-SEGMENT ELEVATION MYOCARDIAL INFARCTION AND UNSTABLE ANGINA)

MEDICAL TREATMENT

Patients should be placed at bed rest with continuous ECG monitoring for ST-segment deviation and cardiac arrhythmias. Ambulation is permitted if the patient shows no recurrence of ischemia (symptoms or ECG changes) and does not develop an elevation of a biomarker of necrosis for 12–24 h. Medical therapy involves simultaneous anti-ischemic and antithrombotic treatments and consideration of coronary revascularization.

ANTI-ISCHEMIC TREATMENT (TABLE 294-1)

To provide relief and prevention of recurrence of chest pain, initial treatment should include bed rest, nitrates, beta adrenergic blockers, and inhaled oxygen in the presence of hypoxemia.

Nitrates These should first be given sublingually or by buccal spray (0.3–0.6 mg) if the patient is experiencing ischemic pain. If pain persists after three doses given 5 min apart, intravenous nitroglycerin (5–10 µg/min using nonabsorbing tubing) is recommended. The rate of the infusion may be increased by 10 µg/min every 3–5 min until symptoms are relieved, systolic arterial pressure falls to

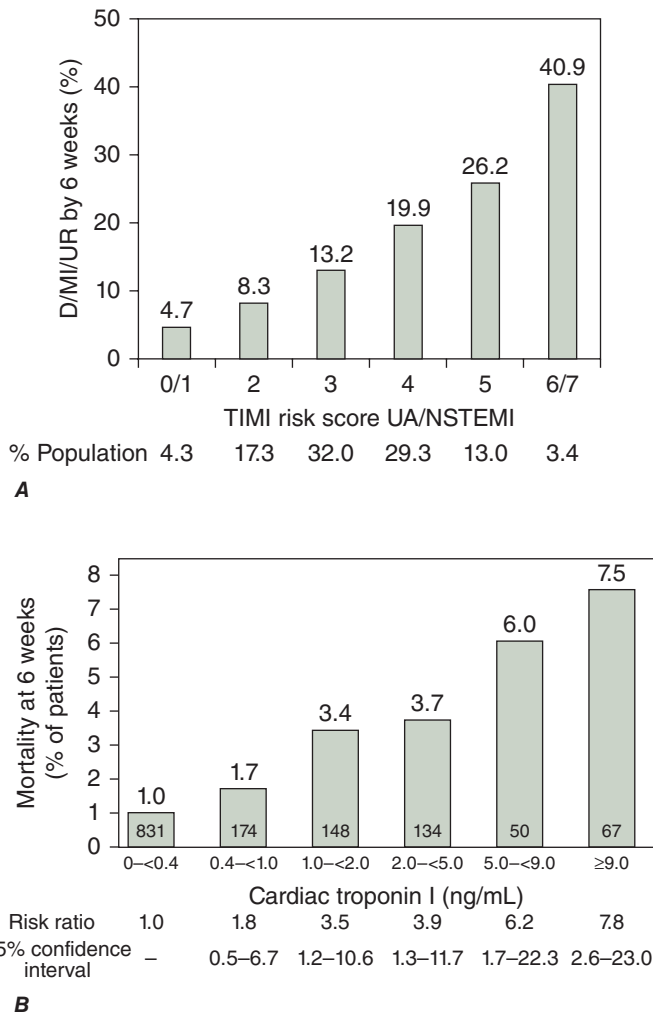


FIGURE 294-4 A. Death (D), myocardial infarction (MI), or need for urgent revascularization (UR) through 6 weeks by Thrombolysis in Myocardial Infarction (TIMI) Risk Score in the unfractionated heparin arm of the TIMI 11B trial. (From EM Antman et al: *JAMA* 284:835, 2000.) B. Mortality rate at 42 days by baseline cardiac troponin I levels in the TIMI 3B trial. (From EM Antman et al: *N Engl J Med* 335:1342, 1996.)

<100 mmHg, or the dose reaches 200 μ g/min. Topical or oral nitrates (Chap. 293) can be used when the pain has resolved, or they may replace intravenous nitroglycerin when the patient has been pain-free for 12–24 h. The only absolute contraindications to the use of nitrates are hypotension or the use of sildenafil or other phosphodiesterase-5 inhibitors within the previous 24–48 h.

Beta Adrenergic Blockers and Other Agents Beta blockers are the other mainstay of anti-ischemic treatment. They may be started by the intravenous route in patients with severe ischemia, but this is contraindicated in the presence of heart failure. Ordinarily, oral beta blockade targeted to a heart rate of 50–60 beats/min is recommended. Heart rate–slowing calcium channel blockers, e.g., verapamil or diltiazem, are recommended for patients who have persistent symptoms or ECG signs of ischemia after treatment with full-dose nitrates and beta blockers and in patients with contraindications to either class of these agents. Additional medical therapy includes angiotensin-converting enzyme (ACE) inhibitors or, if these are not tolerated, angiotensin receptor blockers. Early administration of intensive HMG-CoA reductase inhibitors (statins), such as atorvastatin 80 mg/d, prior to percutaneous coronary intervention (PCI), and continued thereafter, has been shown to reduce complications of the procedure and recurrences of ACS.

ANTITHROMBOTIC THERAPY (TABLE 294-2)

This is the second major cornerstone of treatment. There are two components of antithrombotic therapy: antiplatelet drugs and anticoagulants.

Antiplatelet Drugs (See Chap. 143) Initial treatment should begin with the platelet cyclooxygenase inhibitor aspirin. The typical initial dose is 325 mg/d, with lower doses (75–100 mg/d) recommended thereafter. Contraindications are active bleeding or aspirin intolerance. “Aspirin resistance” has been noted in 2–8% of patients but frequently has been related to noncompliance.

In the absence of a high risk for bleeding, patients with NSTEMI-ACS, irrespective of whether an invasive or conservative strategy (see below) is selected, should receive a platelet P2Y₁₂ receptor blocker to inhibit platelet activation. The thienopyridine clopidogrel is an inactive prodrug that is converted into an active metabolite that causes irreversible blockade of the platelet P2Y₁₂ receptor. When added to aspirin, so-called dual antiplatelet therapy, it has been shown to confer a 20% relative reduction in cardiovascular death, MI, or stroke, compared to aspirin alone, but to be associated with a moderate (absolute 1%) increase in major bleeding.

Continued benefit of treatment with the combination of aspirin and clopidogrel has been observed both in patients treated conservatively and in those who underwent PCI. This regimen should continue for at least 1 year in patients with NSTEMI-ACS, especially those with a drug-eluting stent, to prevent stent thrombosis. Up to one-third of patients have an inadequate response to clopidogrel, and a substantial proportion of these cases are related to a genetic variant of the cytochrome P450 system. A variant of the 2C19 gene leads to reduced conversion of clopidogrel into its active metabolite, which, in turn, reduces platelet inhibition and is associated with increases in the incidence of adverse cardiovascular events. Alternate P2Y₁₂ blockers, such as prasugrel or ticagrelor (see below) used with aspirin, should be considered in patients with NSTEMI-ACS who develop a coronary event while receiving clopidogrel and aspirin or who are hyporesponsive to clopidogrel as identified by platelet and/or genetic testing, although such testing is not yet widespread.

A second P2Y₁₂ blocker, prasugrel, also a thienopyridine, achieves a more rapid onset and higher level of platelet inhibition than clopidogrel. It has been approved for ACS patients following angiography in whom PCI is planned. It should be administered at a loading dose of 60 mg followed by 10 mg/d for up to 15 months. The TRITON-TIMI 38 trial showed that relative to clopidogrel, prasugrel reduced the risk of cardiovascular death, MI, or stroke significantly, albeit with an increase in major bleeding. Stent thrombosis was reduced by half. This agent is contraindicated in patients with prior stroke or transient ischemic attack or at high risk for bleeding. It has not been found to be effective in patients treated by a conservative strategy (see below).

Ticagrelor is a novel, potent, *reversible* platelet P2Y₁₂ inhibitor. It has been shown in the PLATO trial to reduce the risk of cardiovascular death, MI, or stroke compared with clopidogrel in ACS patients who are treated by either an invasive or a conservative strategy. This agent reduced mortality but increased the risk of bleeding not associated with coronary artery bypass grafting. After a loading dose of 180 mg, 90 mg bid is administered as maintenance.

Prior to the development of the oral P2Y₁₂ receptor blockers, many trials had shown the benefit of intravenous glycoprotein IIb/IIIa inhibitors. Their benefit, however, has been small (i.e., only a 10% reduction in death or MI, with a significant increase in major bleeding). Two recent studies failed to show a benefit of routine early initiation of a drug in this class compared with their use only in patients who undergo PCI. The addition of these agents to aspirin and a P2Y₁₂ inhibitor (i.e., triple antiplatelet therapy) should be reserved for unstable patients with recurrent rest pain, elevated cTn, and ECG changes, as well as those who have a coronary thrombus evident on angiography when they undergo PCI.

TABLE 294-1 DRUGS COMMONLY USED IN INTENSIVE MEDICAL MANAGEMENT OF PATIENTS WITH UNSTABLE ANGINA AND NON-ST-SEGMENT ELEVATION MYOCARDIAL INFARCTION

Drug Category	Clinical Condition	When to Avoid ^a	Dosage
Nitrates	Administer sublingually, and, if symptoms persist, intravenously	Hypotension Patient receiving sildenafil or other PDE-5 inhibitor	Topical, oral, or buccal nitrates are acceptable alternatives for patients without ongoing or refractory symptoms 5–10 µg/min by continuous infusion titrated up to 75–100 µg/min until relief of symptoms or limiting side effects (headache or hypotension with a systolic blood pressure <90 mmHg or more than 30% below starting mean arterial pressure levels if significant hypertension is present)
Beta blockers ^b	Unstable angina	PR interval (ECG) <0.24 s 2° or 3° atrioventricular block Heart rate <60 beats/min Systolic pressure <90 mmHg Shock Left ventricular failure Severe reactive airway disease	Metoprolol 25–50 mg by mouth every 6 h If needed, and no heart failure, 5-mg increments by slow (over 1–2 min) IV administration
Calcium channel blockers	Patients whose symptoms are not relieved by adequate doses of nitrates and beta blockers, or in patients unable to tolerate adequate doses of one or both of these agents, or in patients with variant angina	Pulmonary edema Evidence of left ventricular dysfunction (for diltiazem or verapamil)	Dependent on specific agent
Morphine sulfate	Patients whose symptoms are not relieved after three serial sublingual nitroglycerin tablets or whose symptoms recur with adequate anti-ischemic therapy	Hypotension Respiratory depression Confusion Obtundation	2–5 mg IV dose May be repeated every 5–30 min as needed to relieve symptoms and maintain patient comfort

^aAllergy or prior intolerance is a contraindication for all categories of drugs listed in this chart. ^bChoice of the specific agent is not as important as ensuring that appropriate candidates receive this therapy. If there are concerns about patient intolerance due to existing pulmonary disease, especially asthma, left ventricular dysfunction, risk of hypotension, or severe bradycardia, initial selection should favor a short-acting agent, such as propranolol or metoprolol or the ultra-short-acting agent esmolol. Mild wheezing or a history of chronic obstructive pulmonary disease should prompt a trial of a short-acting agent at a reduced dose (e.g., 2.5 mg IV metoprolol, 12.5 mg oral metoprolol, or 25 µg/kg per min esmolol as initial doses) rather than complete avoidance of beta blocker therapy.

Note: Some of the recommendations in this guide suggest the use of agents for purposes or in doses other than those specified by the U.S. Food and Drug Administration. Such recommendations are made after consideration of concerns regarding nonapproved indications. Where made, such recommendations are based on more recent clinical trials or expert consensus. 2°, second-degree; 3°, third-degree; ECG, electrocardiogram; IV, intravenous.

Source: Modified from J Anderson et al: *J Am Coll Cardiol* 61:e179, 2013.

Anticoagulants (See Chap. 143) Four options are available for anticoagulant therapy to be added to antiplatelet agents: (1) unfractionated heparin (UFH), long the mainstay of therapy; (2) the low-molecular-weight heparin (LMWH), enoxaparin, which has been shown to be superior to UFH in reducing recurrent cardiac events, especially in patients managed by a conservative strategy but with some increase in bleeding; (3) bivalirudin, a direct thrombin inhibitor that is similar in efficacy to either UFH or LMWH but causes less bleeding and is used just prior to and/or during PCI; and (4) the indirect factor Xa inhibitor, fondaparinux, which is equivalent in efficacy to enoxaparin but appears to have a lower risk of major bleeding.

Excessive bleeding is the most important adverse effect of all antithrombotic agents, including both antiplatelet agents and anticoagulants. Therefore, attention must be directed to the doses of antithrombotic agents, accounting for body weight, creatinine clearance, and a previous history of excessive bleeding, as a means of reducing the risk of bleeding. Patients who have experienced a stroke are at higher risk of intracranial bleeding with potent antiplatelet agents and combinations of antithrombotic drugs.

INVASIVE VERSUS CONSERVATIVE STRATEGY

Multiple clinical trials have demonstrated the benefit of an early invasive strategy in high-risk patients (i.e., patients with multiple

clinical risk factors, ST-segment deviation, and/or positive biomarkers) (Table 294-3). In this strategy, following treatment with anti-ischemic and antithrombotic agents, coronary arteriography is carried out within ~48 h of presentation, followed by coronary revascularization (PCI or coronary artery bypass grafting), depending on the coronary anatomy. In low-risk patients, the outcomes from an invasive strategy are similar to those obtained from a conservative strategy. The latter consists of anti-ischemic and antithrombotic therapy followed by “watchful waiting,” in which the patient is closely observed and coronary arteriography is carried out only if rest pain or ST-segment changes recur, a biomarker of necrosis becomes positive, or there is evidence of severe ischemia on a stress test.

LONG-TERM MANAGEMENT

The time of hospital discharge is a “teachable moment” for the patient with NSTEMI-ACS, when the physician can review and optimize the medical regimen. Risk-factor modification is key, and the caregiver should discuss with the patient the importance of smoking cessation, achieving optimal weight, daily exercise, blood-pressure control, following an appropriate diet, control of hyperglycemia (in diabetic patients), and lipid management as recommended for patients with chronic stable angina (Chap. 293).

TABLE 294-2 CLINICAL USE OF ANTITHROMBOTIC THERAPY**Oral Antiplatelet Therapy**

Aspirin	Initial dose of 325 mg nonenteric formulation followed by 75–100 mg/d of an enteric or a nonenteric formulation
Clopidogrel	Loading dose of 300–600 mg followed by 75 mg/d
Prasugrel	Pre-PCI: Loading dose 60 mg followed by 10 mg/d
Ticagrelor	Loading dose of 180 mg followed by 90 mg twice daily

Intravenous Antiplatelet Therapy

Abciximab	0.25 mg/kg bolus followed by infusion of 0.125 µg/kg per min (maximum 10 µg/min) for 12–24 h
Eptifibatid	180 µg/kg bolus followed 10 min later by second bolus of 180 µg with infusion of 2.0 µg/kg per min for 72–96 h following first bolus
Tirofiban	25 µg/kg per min followed by infusion of 0.15 µg/kg per min for 48–96 h

Heparins^a

Unfractionated heparin (UFH)	^b Bolus 70–100 U/kg (maximum 5000 U) IV followed by infusion of 12–15 U/kg per h (initial maximum 1000 U/h) titrated to ACT 250–300 s
Enoxaparin	1 mg/kg SC every 12 h; the first dose may be preceded by a 30-mg IV bolus; renal adjustment to 1 mg/kg once daily if creatine clearance <30 cc/min
Fondaparinux	2.5 mg SC qd
Bivalirudin	Initial IV bolus of 0.75 mg/kg and an infusion of 1.75 mg/kg per h.

^aOther low-molecular-weight heparins exist beyond those listed. ^bIf no glycoprotein IIb/IIIa inhibitor planned.

Abbreviations: ACT, activated clotting time for HemoTec; IV, intravenous; SC, subcutaneously.

Source: Modified from J Anderson et al: J Am Coll Cardiol 61:e179, 2013.

There is evidence of benefit with long-term therapy with five classes of drugs that are directed at different components of the atherothrombotic process. Beta blockers, statins (at a high dose, e.g., atorvastatin 80 mg/d), and ACE inhibitors or angiotensin receptor blockers are recommended for long-term plaque stabilization. Antiplatelet therapy,

now recommended to be the combination of low-dose (75–100 mg/d) aspirin and a P2Y₁₂ inhibitor (clopidogrel, prasugrel, or ticagrelor) for 1 year, with aspirin continued thereafter, prevents or reduces the severity of any thrombosis that would occur if a plaque were to rupture.

Registries have shown that women and racial minorities, as well as patients with NSTEMI-ACS at high risk, including the elderly and patients with diabetes or chronic kidney disease, are less likely to receive evidence-based pharmacologic and interventional therapies with resultant poorer clinical outcomes and quality of life. Special attention should be directed to these groups.

PRINZMETAL'S VARIANT ANGINA

In 1959 Prinzmetal et al. described a syndrome of severe ischemic pain that usually occurs at rest and is associated with transient ST-segment elevation. Prinzmetal's variant angina (PVA) is caused by focal spasm of an epicardial coronary artery, leading to severe transient myocardial ischemia and occasionally infarction. The cause of the spasm is not well defined, but it may be related to hypercontractility of vascular smooth muscle due to adrenergic vasoconstrictors, leukotrienes, or serotonin. For reasons that are not clear, the prevalence of PVA has decreased substantially during the past few decades.

Clinical and Angiographic Manifestations Patients with PVA are generally younger and have fewer coronary risk factors (with the exception of cigarette smoking) than do patients with NSTEMI-ACS. Cardiac examination is usually unremarkable in the absence of ischemia. The clinical diagnosis of PVA is made by the detection of transient ST-segment elevation with rest pain. Many patients also exhibit multiple episodes of asymptomatic ST-segment elevation (*silent ischemia*). Small elevations of troponin may occur in patients with prolonged attacks.

Coronary angiography demonstrates transient coronary spasm as the diagnostic hallmark of PVA. Atherosclerotic plaques in at least one proximal coronary artery occur in about half of patients, and in these patients, spasm usually occurs within 1 cm of the plaque. Focal spasm is most common in the right coronary artery, and it may occur at one or more sites in one artery or in multiple arteries simultaneously. Hyperventilation or intracoronary acetylcholine has been used to provoke focal coronary stenosis on angiography or to provoke rest angina with ST-segment elevation to establish the diagnosis.

TABLE 294-3 CLASS I RECOMMENDATIONS FOR USE OF AN EARLY INVASIVE STRATEGY IN PATIENTS WITH NON-ST-SEGMENT ELEVATION ACUTE CORONARY SYNDROME^a**Class I (Level of Evidence: A) Indications**

Recurrent angina at rest/low-level activity despite treatment
Elevated TnT or TnI
New ST-segment depression
CHF symptoms, rales, MR
EF <0.40
Sustained VT
PCI <6 months, prior CABG
High-risk findings from noninvasive testing
Hemodynamic instability
Mild-to-moderate renal dysfunction
Diabetes mellitus
High TIMI Risk Score (>3) ^b

^aAny one of the high-risk indicators. ^bSee Antman (JAMA 284:835, 2000).

Abbreviations: CABG, coronary artery bypass grafting; CHF, congestive heart failure; EF, ejection fraction; MR, mitral regurgitation; PCI, percutaneous coronary intervention; TIMI, Thrombolysis in Myocardial Infarction; TnI, troponin I; TnT, troponin T; VT, ventricular tachycardia.

Source: Modified from J Anderson et al: J Am Coll Cardiol 61:e179, 2013.

TREATMENT PRINZMETAL'S VARIANT ANGINA

Nitrates and calcium channel blockers are the main therapeutic agents. Aspirin may actually increase the severity of ischemic episodes, possibly as a result of the sensitivity of coronary tone to modest changes in the synthesis of prostacyclin. The response to beta blockers is variable. Coronary revascularization may be helpful in patients who also have discrete, flow-limiting, proximal fixed obstructive lesions.

Prognosis Many patients with PVA pass through an acute, active phase, with frequent episodes of angina and cardiac events during the first 6 months after presentation. Survival at 5 years is excellent (~90–95%). Patients with no or mild fixed coronary obstruction tend to experience a more benign course than do patients with associated severe obstructive lesions. Nonfatal MI occurs in up to 20% of patients by 5 years. Patients with PVA who develop serious arrhythmias during spontaneous episodes of pain are at a higher risk for sudden cardiac death. In most patients who survive an infarction or the initial 3- to 6-month period of frequent episodes, there is a tendency for symptoms and cardiac events to diminish over time.

295 ST-Segment Elevation Myocardial Infarction

Elliott M. Antman, Joseph Loscalzo

Acute myocardial infarction (AMI) is one of the most common diagnoses in hospitalized patients in industrialized countries. In the United States, approximately 525,000 patients experience a new AMI, and 190,000 experience a recurrent AMI each year. More than half of AMI-related deaths occur before the stricken individual reaches the hospital. The in-hospital mortality rate after admission for AMI has declined from 10% to about 6% over the past decade. The 1-year mortality rate after AMI is about 15%. Mortality is approximately fourfold higher in elderly patients (over age 75) as compared with younger patients.

When patients with prolonged ischemic discomfort at rest are first seen, the working clinical diagnosis is that they are suffering from an acute coronary syndrome (Fig. 295-1). The 12-lead electrocardiogram (ECG) is a pivotal diagnostic and triage tool because it is at the center of the decision pathway for management; it permits distinction of those patients presenting with ST-segment elevation from those presenting without ST-segment elevation. Serum cardiac biomarkers are obtained to distinguish unstable angina (UA) from non-ST-segment elevation myocardial infarction (NSTEMI) and to assess the magnitude of an ST-segment elevation myocardial infarction (STEMI). This chapter focuses on the evaluation and management of patients with STEMI, while Chap. 294 discusses UA/NSTEMI.

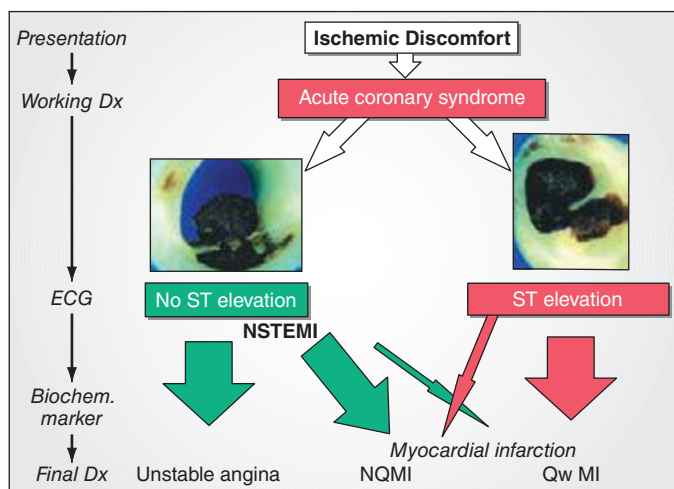


FIGURE 295-1 Acute coronary syndromes. Following disruption of a vulnerable plaque, patients experience ischemic discomfort resulting from a reduction of flow through the affected epicardial coronary artery. The flow reduction may be caused by a completely occlusive thrombus (right) or subtotally occlusive thrombus (left). Patients with ischemic discomfort may present with or without ST-segment elevation. Of patients with ST-segment elevation, the majority (wide red arrow) ultimately develop a Q wave on the ECG (Qw MI), while a minority (thin red arrow) do not develop Q wave and, in older literature, were said to have sustained a non-Q-wave MI (NQMI). Patients who present without ST-segment elevation are suffering from either unstable angina or a non-ST-segment elevation MI (NSTEMI) (wide green arrows), a distinction that is ultimately made based on the presence or absence of a serum cardiac marker such as CK-MB or a cardiac troponin detected in the blood. The majority of patients presenting with NSTEMI do not develop a Q wave on the ECG; a minority develop a Qw MI (thin green arrow). Dx, diagnosis; ECG, electrocardiogram; MI, myocardial infarction. (Adapted from CW Hamm et al: *Lancet* 358:1533, 2001, and MJ Davies: *Heart* 83:361, 2000; with permission from the BMJ Publishing Group.)

STEMI usually occurs when coronary blood flow decreases abruptly after a thrombotic occlusion of a coronary artery previously affected by atherosclerosis. Slowly developing, high-grade coronary artery stenoses do not typically precipitate STEMI because of the development of a rich collateral network over time. Instead, STEMI occurs when a coronary artery thrombus develops rapidly at a site of vascular injury. This injury is produced or facilitated by factors such as cigarette smoking, hypertension, and lipid accumulation. In most cases, STEMI occurs when the surface of an atherosclerotic plaque becomes disrupted (exposing its contents to the blood) and conditions (local or systemic) favor thrombogenesis. A mural thrombus forms at the site of plaque disruption, and the involved coronary artery becomes occluded. Histologic studies indicate that the coronary plaques prone to disruption are those with a rich lipid core and a thin fibrous cap (Chap. 291e). After an initial platelet monolayer forms at the site of the disrupted plaque, various agonists (collagen, ADP, epinephrine, serotonin) promote platelet activation. After agonist stimulation of platelets, thromboxane A₂ (a potent local vasoconstrictor) is released, further platelet activation occurs, and potential resistance to fibrinolysis develops.

In addition to the generation of thromboxane A₂, activation of platelets by agonists promotes a conformational change in the glycoprotein IIb/IIIa receptor (Chap. 140). Once converted to its functional state, this receptor develops a high affinity for soluble adhesive proteins (i.e., integrins) such as fibrinogen. Since fibrinogen is a multivalent molecule, it can bind to two different platelets simultaneously, resulting in platelet cross-linking and aggregation.

The coagulation cascade is activated on exposure of tissue factor in damaged endothelial cells at the site of the disrupted plaque. Factors VII and X are activated, ultimately leading to the conversion of prothrombin to thrombin, which then converts fibrinogen to fibrin (Chap. 141). Fluid-phase and clot-bound thrombin participates in an autoamplification reaction leading to further activation of the coagulation cascade. The culprit coronary artery eventually becomes occluded by a thrombus containing platelet aggregates and fibrin strands.

In rare cases, STEMI may be due to coronary artery occlusion caused by coronary emboli, congenital abnormalities, coronary spasm, and a wide variety of systemic—particularly inflammatory—diseases. The amount of myocardial damage caused by coronary occlusion depends on (1) the territory supplied by the affected vessel, (2) whether or not the vessel becomes totally occluded, (3) the duration of coronary occlusion, (4) the quantity of blood supplied by collateral vessels to the affected tissue, (5) the demand for oxygen of the myocardium whose blood supply has been suddenly limited, (6) endogenous factors that can produce early spontaneous lysis of the occlusive thrombus, and (7) the adequacy of myocardial perfusion in the infarct zone when flow is restored in the occluded epicardial coronary artery.

Patients at increased risk for developing STEMI include those with multiple coronary risk factors (Chap. 291e) and those with UA (Chap. 294). Less common underlying medical conditions predisposing patients to STEMI include hypercoagulability, collagen vascular disease, cocaine abuse, and intracardiac thrombi or masses that can produce coronary emboli.

There have been major advances in the management of STEMI with recognition that the “chain of survival” involves a highly integrated system starting with prehospital care and extending to early hospital management so as to provide expeditious implementation of a reperfusion strategy.

CLINICAL PRESENTATION

In up to one-half of cases, a precipitating factor appears to be present before STEMI, such as vigorous physical exercise, emotional stress, or a medical or surgical illness. Although STEMI may commence at any time of the day or night, circadian variations have been reported such that clusters are seen in the morning within a few hours of awakening.

Pain is the most common presenting complaint in patients with STEMI. The pain is deep and visceral; adjectives commonly used to

describe it are *heavy, squeezing, and crushing*, although, occasionally, it is described as stabbing or burning (**Chap. 19**). It is similar in character to the discomfort of angina pectoris (**Chap. 293**) but commonly occurs at rest, is usually more severe, and lasts longer. Typically, the pain involves the central portion of the chest and/or the epigastrium, and, on occasion, it radiates to the arms. Less common sites of radiation include the abdomen, back, lower jaw, and neck. The frequent location of the pain beneath the xiphoid and epigastrium and the patients' denial that they may be suffering a heart attack are chiefly responsible for the common mistaken impression of indigestion. The pain of STEMI may radiate as high as the occipital area but not below the umbilicus. It is often accompanied by weakness, sweating, nausea, vomiting, anxiety, and a sense of impending doom. The pain may commence when the patient is at rest, but when it begins during a period of exertion, it does not usually subside with cessation of activity, in contrast to angina pectoris.

The pain of STEMI can simulate pain from acute pericarditis (**Chap. 288**), pulmonary embolism (**Chap. 300**), acute aortic dissection (**Chap. 301**), costochondritis, and gastrointestinal disorders. These conditions should therefore be considered in the differential diagnosis. Radiation of discomfort to the trapezius is not seen in patients with STEMI and may be a useful distinguishing feature that suggests pericarditis is the correct diagnosis. However, *pain is not uniformly present in patients with STEMI*. The proportion of painless STEMIs is greater in patients with diabetes mellitus, and it increases with age. In the elderly, STEMI may present as sudden-onset breathlessness, which may progress to pulmonary edema. Other less common presentations, with or without pain, include sudden loss of consciousness, a confusional state, a sensation of profound weakness, the appearance of an arrhythmia, evidence of peripheral embolism, or merely an unexplained drop in arterial pressure.

PHYSICAL FINDINGS

Most patients are anxious and restless, attempting unsuccessfully to relieve the pain by moving about in bed, altering their position, and stretching. Pallor associated with perspiration and coolness of the extremities occurs commonly. The combination of substernal chest pain persisting for >30 min and diaphoresis strongly suggests STEMI. Although many patients have a normal pulse rate and blood pressure within the first hour of STEMI, about one-fourth of patients with anterior infarction have manifestations of sympathetic nervous system hyperactivity (tachycardia and/or hypertension), and up to one-half with inferior infarction show evidence of parasympathetic hyperactivity (bradycardia and/or hypotension).

The precordium is usually quiet, and the apical impulse may be difficult to palpate. In patients with anterior wall infarction, an abnormal systolic pulsation caused by dyskinetic bulging of infarcted myocardium may develop in the periapical area within the first days of the illness and then may resolve. Other physical signs of ventricular dysfunction include fourth and third heart sounds, decreased intensity of the first heart sound, and paradoxical splitting of the second heart sound (**Chap. 267**). A transient midsystolic or late systolic apical systolic murmur due to dysfunction of the mitral valve apparatus may be present. A pericardial friction rub may be heard in patients with transmural STEMI at some time in the course of the disease, if they are examined frequently. The carotid pulse is often decreased in volume, reflecting reduced stroke volume. Temperature elevations up to 38°C may be observed during the first week after STEMI. The arterial pressure is variable; in most patients with transmural infarction, systolic pressure declines by approximately 10–15 mmHg from the preinfarction state.

LABORATORY FINDINGS

STEMI progresses through the following temporal stages: (1) acute (first few hours–7 days), (2) healing (7–28 days), and (3) healed (≥ 29 days). When evaluating the results of diagnostic tests for STEMI, the temporal phase of the infarction must be considered. The laboratory tests of value in confirming the diagnosis may be divided into four groups: (1) ECG, (2) serum cardiac biomarkers, (3) cardiac imaging, and (4) nonspecific indices of tissue necrosis and inflammation.

ELECTROCARDIOGRAM

The electrocardiographic manifestations of STEMI are described in **Chap. 268**. During the initial stage, total occlusion of an epicardial coronary artery produces ST-segment elevation. Most patients initially presenting with ST-segment elevation ultimately evolve Q waves on the ECG. However, Q waves in the leads overlying the infarct zone may vary in magnitude and even appear only transiently, depending on the reperfusion status of the ischemic myocardium and restoration of transmembrane potentials over time. A small proportion of patients initially presenting with ST-segment elevation will not develop Q waves when the obstructing thrombus is not totally occlusive, obstruction is transient, or if a rich collateral network is present. Among patients presenting with ischemic discomfort but *without* ST-segment elevation, if a serum cardiac biomarker of necrosis (see below) is detected, the diagnosis of NSTEMI is ultimately made (**Fig. 295-1**). A minority of patients who present initially without ST-segment elevation may develop a Q-wave MI. Previously, it was believed that transmural myocardial infarction (MI) is present if the ECG demonstrates Q waves or loss of R waves, and nontransmural MI may be present if the ECG shows only transient ST-segment and T-wave changes. However, electrocardiographic-pathologic correlations are far from perfect and terms such as *Q-wave MI, non-Q-wave MI, transmural MI, and nontransmural MI*, have been replaced by STEMI and NSTEMI (**Fig. 295-1**). Contemporary studies using magnetic resonance imaging (MRI) suggest that the development of a Q wave on the ECG is more dependent on the volume of infarcted tissue rather than the transmural extent of infarction.

SERUM CARDIAC BIOMARKERS

Certain proteins, called serum cardiac biomarkers, are released from necrotic heart muscle after STEMI. The rate of liberation of specific proteins differs depending on their intracellular location, their molecular weight, and the local blood and lymphatic flow. Cardiac biomarkers become detectable in the peripheral blood once the capacity of the cardiac lymphatics to clear the interstitium of the infarct zone is exceeded and spillover into the venous circulation occurs. The temporal pattern of protein release is of diagnostic importance. The criteria for AMI require a rise and/or fall in cardiac biomarker values with at least one value above the 99th percentile of the upper reference limit for normal individuals.

Cardiac-specific troponin T (cTnT) and *cardiac-specific troponin I (cTnI)* have amino-acid sequences different from those of the skeletal muscle forms of these proteins. These differences permitted the development of quantitative assays for cTnT and cTnI with highly specific monoclonal antibodies. Since cTnT and cTnI are not normally detectable in the blood of healthy individuals but may increase after STEMI to levels many times higher than the upper reference limit (the highest value seen in 99% of a reference population not suffering from MI), the measurement of cTnT or cTnI is of considerable diagnostic usefulness, and they are now the preferred biochemical markers for MI (**Fig. 295-2**). With improvements in the assays for the cardiac-specific troponins, it is now possible to detect concentrations <1 ng/L in patients without ischemic-type chest discomfort. The cardiac troponins are particularly valuable when there is clinical suspicion of either skeletal muscle injury or a small MI that may be below the detection limit for creatine phosphokinase (CK) and its MB isoenzyme (CK-MB) measurements, and they are, therefore, of particular value in distinguishing UA from NSTEMI. In practical terms, the high-sensitivity troponin assays are of less immediate value in patients with STEMI. Contemporary urgent reperfusion strategies necessitate making a decision (based largely on a combination of clinical and ECG findings) before the results of blood tests have returned from the laboratory. Levels of cTnI and cTnT may remain elevated for 7–10 days after STEMI.

CK rises within 4–8 h and generally returns to normal by 48–72 h (**Fig. 295-2**). An important drawback of total CK measurement is its lack of specificity for STEMI, as CK may be elevated with skeletal muscle disease or trauma, including intramuscular injection. The MB isoenzyme of CK has the advantage over total CK that it is not present in significant concentrations in extracardiac tissue and, therefore,

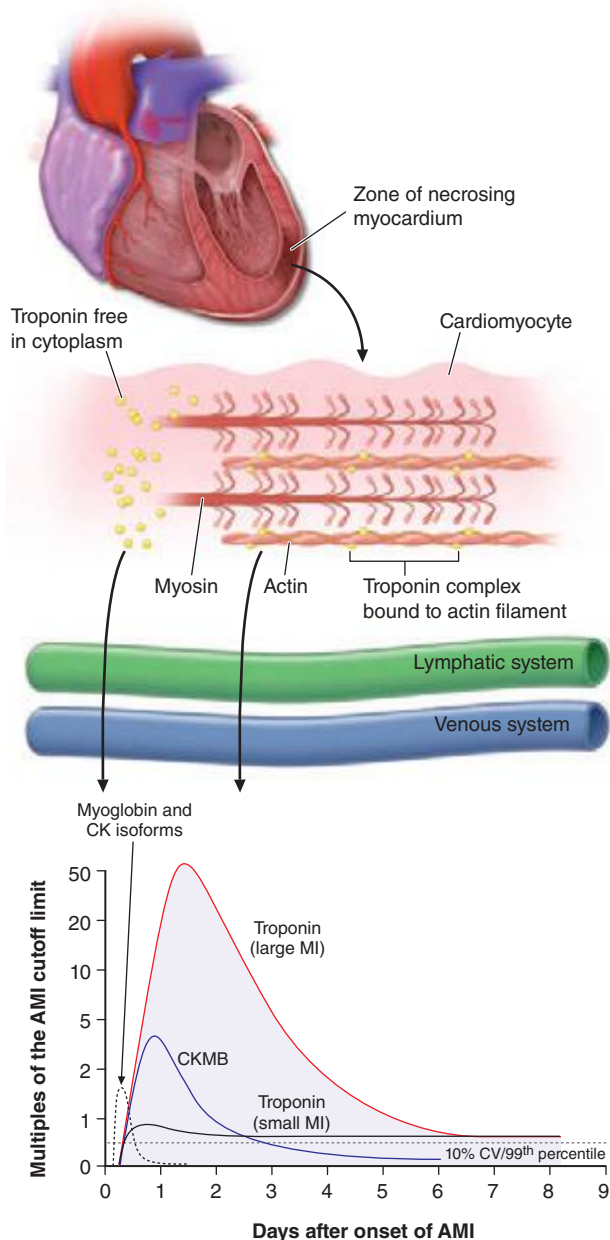


FIGURE 295-2 The zone of necrosing myocardium is shown at the top of the figure, followed in the middle portion of the figure by a diagram of a cardiomyocyte that is in the process of releasing biomarkers. The biomarkers that are released into the interstitium are first cleared by lymphatics followed subsequently by spillover into the venous system. After disruption of the sarcolemmal membrane of the cardiomyocyte, the cytoplasmic pool of biomarkers is released first (left-most arrow in bottom portion of figure). Markers such as myoglobin and CK isoforms are rapidly released, and blood levels rise quickly above the cutoff limit; this is then followed by a more protracted release of biomarkers from the disintegrating myofilaments that may continue for several days. Cardiac troponin levels rise to about 20 to 50 times the upper reference limit (the 99th percentile of values in a reference control group) in patients who have a “classic” acute myocardial infarction (MI) and sustain sufficient myocardial necrosis to result in abnormally elevated levels of the MB fraction of creatine kinase (CK-MB). Clinicians can now diagnose episodes of microinfarction by sensitive assays that detect cardiac troponin elevations above the upper reference limit, even though CK-MB levels may still be in the normal reference range (not shown). CV, coefficient of variation. (Modified from EM Antman: *Decision making with cardiac troponin tests*. *N Engl J Med* 346:2079, 2002 and AS Jaffe, L Babuin, FS Apple: *Biomarkers in acute cardiac disease: The present and the future*. *J Am Coll Cardiol* 48:1, 2006.)

is considerably more specific. However, cardiac surgery, myocarditis, and electrical cardioversion often result in elevated serum levels of the MB isoenzyme. A ratio (relative index) of CK-MB mass to CK activity ≥ 2.5 suggests but is not diagnostic of a myocardial rather than a skeletal muscle source for the CK-MB elevation.

Many hospitals are using cTnT or cTnI rather than CK-MB as the routine serum cardiac marker for diagnosis of STEMI, although any of these analytes remain clinically acceptable. It is *not* cost-effective to measure both a cardiac-specific troponin and CK-MB at all time points in every patient.

While it has long been recognized that the total quantity of protein released correlates with the size of the infarct, the peak protein concentration correlates only weakly with infarct size. Recanalization of a coronary artery occlusion (either spontaneously or by mechanical or pharmacologic means) in the early hours of STEMI causes earlier peaking of biomarker measurements (Fig. 295-2) because of a rapid washout from the interstitium of the infarct zone, quickly overwhelming lymphatic clearance of the proteins.

The *nonspecific reaction* to myocardial injury is associated with polymorphonuclear leukocytosis, which appears within a few hours after the onset of pain and persists for 3–7 days; the white blood cell count often reaches levels of 12,000–15,000/ μ L. The erythrocyte sedimentation rate rises more slowly than the white blood cell count, peaking during the first week and sometimes remaining elevated for 1 or 2 weeks.

CARDIAC IMAGING

Abnormalities of wall motion on *two-dimensional echocardiography* (Chap. 270e) are almost universally present. Although acute STEMI cannot be distinguished from an old myocardial scar or from acute severe ischemia by echocardiography, the ease and safety of the procedure make its use appealing as a screening tool in the Emergency Department setting. When the ECG is not diagnostic of STEMI, early detection of the presence or absence of wall motion abnormalities by echocardiography can aid in management decisions, such as whether the patient should receive reperfusion therapy (e.g., fibrinolysis or a percutaneous coronary intervention [PCI]). Echocardiographic estimation of left ventricular (LV) function is useful prognostically; detection of reduced function serves as an indication for therapy with an inhibitor of the renin-angiotensin-aldosterone system. Echocardiography may also identify the presence of right ventricular (RV) infarction, ventricular aneurysm, pericardial effusion, and LV thrombus. In addition, Doppler echocardiography is useful in the detection and quantitation of a ventricular septal defect and mitral regurgitation, two serious complications of STEMI.

Several *radionuclide imaging techniques* (Chap. 270e) are available for evaluating patients with suspected STEMI. However, these imaging modalities are used less often than echocardiography because they are more cumbersome and lack sensitivity and specificity in many clinical circumstances. Myocardial perfusion imaging with [^{201}Tl] or [$^{99\text{m}}\text{Tc}$]-sestamibi, which are distributed in proportion to myocardial blood flow and concentrated by viable myocardium (Chap. 293), reveals a defect (“cold spot”) in most patients during the first few hours after development of a transmural infarct. Although perfusion scanning is extremely sensitive, it cannot distinguish acute infarcts from chronic scars and, thus, is not specific for the diagnosis of *acute* MI. Radionuclide ventriculography, carried out with [$^{99\text{m}}\text{Tc}$]-labeled red blood cells, frequently demonstrates wall motion disorders and reduction in the ventricular ejection fraction in patients with STEMI. While of value in assessing the hemodynamic consequences of infarction and in aiding in the diagnosis of RV infarction when the RV ejection fraction is depressed, this technique is nonspecific, as many cardiac abnormalities other than MI alter the radionuclide ventriculogram.

MI can be detected accurately with high-resolution cardiac MRI (Chap. 270e) using a technique referred to as late enhancement. A standard imaging agent (gadolinium) is administered and images are obtained after a 10-min delay. Since little gadolinium enters normal myocardium, where there are tightly packed myocytes, but does percolate into the expanded intercellular region of the infarct zone, there

Criteria for Acute Myocardial Infarction

The term acute myocardial infarction (MI) should be used when there is evidence of myocardial necrosis in a clinical setting consistent with acute myocardial ischemia. Under these conditions, any one of the following criteria meets the diagnosis for MI:

- Detection of a rise and/or fall of cardiac biomarker values (preferably cardiac troponin [cTn]) with at least one value above the 99th percentile upper reference limit (URL) and with at least one of the following:
 - Symptoms of ischemia
 - New or presumed new significant ST-segment T-wave (ST-T) changes or new left bundle branch block (LBBB)
 - Development of pathologic Q waves in the electrocardiogram (ECG)
 - Imaging evidence of new loss of viable myocardium or new regional wall motion abnormality
 - Identification of an intracoronary thrombus by angiography or autopsy
- Cardiac death with symptoms suggestive of myocardial ischemia and presumed new ischemic ECG changes of new LBBB, but death occurred before cardiac biomarkers were obtained or before cardiac biomarker values would be increased.
- Percutaneous coronary intervention (PCI)-related MI is arbitrarily defined by elevation of cTn values ($>5 \times 99$ th percentile URL) in patients with normal baseline values (≤ 99 th percentile URL) or a rise of cTn values $>20\%$ if the baseline values are elevated and are stable or falling. In addition, either (i) symptoms suggestive of myocardial ischemia, or (ii) new ischemic ECG changes, or (iii) angiographic findings consistent with a procedural complication, or (iv) imaging demonstration of new loss of viable myocardium or new regional wall motion abnormality are required.
- Stent thrombosis associated with MI when detected by coronary angiography or autopsy in the setting of myocardial ischemia and with a rise and/or fall of cardiac biomarker values with at least one value above the 99th percentile URL.
- Coronary artery bypass grafting (CABG)-related MI is arbitrarily defined by elevation of cardiac biomarker values ($>10 \times 99$ th percentile URL) in patients with normal baseline cTn values (≤ 99 th percentile URL). In addition, either (i) new pathologic Q waves or new LBBB, or (ii) angiographic documented new graft or new native coronary artery occlusion, or (iii) imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.

Criteria for Prior Myocardial Infarction

Any one of the following criteria meets the diagnosis for prior MI:

- Pathologic Q waves with or without symptoms in the absence of nonischemic causes.
- Imaging evidence of a region of loss of viable myocardium that is thinned and fails to contract, in the absence of a nonischemic cause.
- Pathologic findings of a prior MI.

Source: K Thygesen: Eur Heart J 33:2551, 2012.

is a bright signal in areas of infarction that appears in stark contrast to the dark areas of normal myocardium.

An Expert Consensus Task Force for the Universal Definition of Myocardial Infarction has provided a comprehensive set of criteria for the definition of MI that integrates the clinical and laboratory findings discussed earlier (Table 295-1) as well as a classification of MI into five types that reflect the clinical circumstances in which it may occur (Table 295-2).

INITIAL MANAGEMENT**PREHOSPITAL CARE**

The prognosis in STEMI is largely related to the occurrence of two general classes of complications: (1) electrical complications (arrhythmias) and (2) mechanical complications (“pump failure”). Most out-of-hospital deaths from STEMI are due to the sudden development of ventricular fibrillation. The vast majority of deaths due to ventricular fibrillation occur within the first 24 h of the onset of symptoms, and of these, over half occur in the first hour. Therefore, the major elements of prehospital care of patients with suspected STEMI include (1) recognition of symptoms by the patient and prompt seeking of medical attention; (2) rapid deployment of an emergency medical team capable

Type I: Spontaneous Myocardial Infarction

Spontaneous myocardial infarction related to atherosclerotic plaque rupture, ulceration, fissuring, erosion, or dissection with resulting intraluminal thrombus in one or more of the coronary arteries leading to decreased myocardial blood flow or distal platelet emboli with ensuing myocyte necrosis. The patient may have underlying severe coronary artery disease (CAD) but on occasion nonobstructive or no CAD.

Type 2: Myocardial Infarction Secondary to an Ischemic Imbalance

In instances of myocardial injury with necrosis where a condition other than CAD contributes to an imbalance between myocardial oxygen supply and/or demand, e.g., coronary endothelial dysfunction, coronary artery spasm, coronary embolism, tachy-brady-arrhythmias, anemia, respiratory failure, hypotension, and hypertension with or without left ventricular hypertrophy.

Type 3: Myocardial Infarction Resulting in Death When Biomarker Values Are Unavailable

Cardiac death with symptoms suggestive of myocardial ischemia and presumed new ischemic electrocardiogram (ECG) changes or new left bundle branch block (LBBB), but death occurring before blood samples could be obtained or before cardiac biomarker could rise, or in rare cases, cardiac biomarkers were not collected.

Type 4a: Myocardial Infarction Related to Percutaneous Coronary Intervention (PCI)

Myocardial infarction associated with PCI is arbitrarily defined by elevation of cardiac troponin (cTn) values $>5 \times 99$ th percentile upper reference limit (URL) in patients with normal baseline values (≤ 99 th percentile URL) or a rise of cTn values $>20\%$ if the baseline values are elevated and are stable or falling. In addition, either (i) symptoms suggestive of myocardial ischemia, or (ii) new ischemic ECG changes or new LBBB, or (iii) angiographic loss of patency of a major coronary artery or a side branch or persistent slow or no flow or embolization, or (iv) imaging demonstration of new loss of viable myocardium or new regional wall motion abnormality is required.

Type 4b: Myocardial Infarction Related to Stent Thrombosis

Myocardial infarction associated with stent thrombosis is detected by coronary angiography or autopsy in the setting of myocardial ischemia and with a rise and/or fall of cardiac biomarker values with at least one value above the 99th percentile URL.

Type 5: Myocardial Infarction Related to Coronary Artery Bypass Grafting (CABG)

Myocardial infarction associated with CABG is arbitrarily defined by elevation of cardiac biomarker values $>10 \times 99$ th percentile URL in patients with normal baseline cTn values (≤ 99 th percentile URL). In addition, either (i) new pathologic Q waves or new LBBB, or (ii) angiographic documented new graft or new native coronary artery occlusion, or (iii) imaging evidence of new loss of viable myocardium or new regional wall motion abnormality.

Source: K Thygesen: Eur Heart J 33:2551, 2012.

of performing resuscitative maneuvers, including defibrillation; (3) expeditious transportation of the patient to a hospital facility that is continuously staffed by physicians and nurses skilled in managing arrhythmias and providing advanced cardiac life support; and (4) expeditious implementation of reperfusion therapy (Fig. 295-3). The greatest delay usually occurs not during transportation to the hospital but, rather, between the onset of pain and the patient’s decision to call for help. This delay can best be reduced by health care professionals educating the public concerning the significance of chest discomfort and the importance of seeking early medical attention. Regular office visits with patients having a history of or who are at risk for ischemic heart disease are important “teachable moments” for clinicians to review the symptoms of STEMI and the appropriate action plan.

Increasingly, monitoring and treatment are carried out by trained personnel in the ambulance, further shortening the time between the onset of the infarction and appropriate treatment. General guidelines for initiation of fibrinolysis in the prehospital setting include the ability to transmit 12-lead ECGs to confirm the diagnosis, the presence of paramedics in the ambulance, training of paramedics in the interpretation of ECGs and management of STEMI, and online medical command and control that can authorize the initiation of treatment in the field.

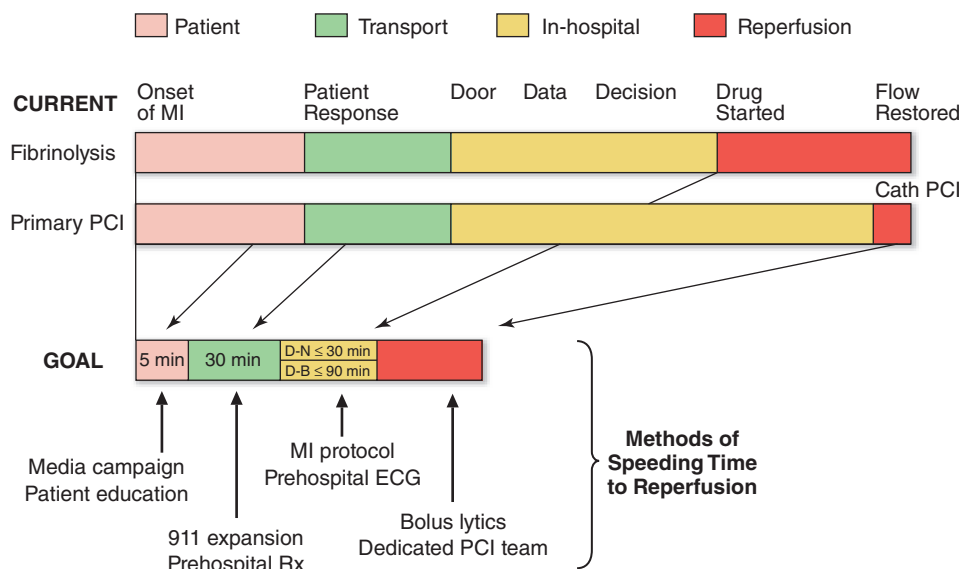


FIGURE 295-3 Major components of time delay between onset of symptoms from ST-segment elevation myocardial infarction and restoration of flow in the infarct-related artery. Plotted sequentially from left to right are the times for patients to recognize symptoms and seek medical attention, transportation to the hospital, in-hospital decision making, implementation of reperfusion strategy, and restoration of flow once the reperfusion strategy has been initiated. The time to initiate fibrinolytic therapy is the “door-to-needle” (D-N) time; this is followed by the period of time required for pharmacologic restoration of flow. More time is required to move the patient to the catheterization laboratory for a percutaneous coronary interventional (PCI) procedure, referred to as the “door-to-balloon” (D-B) time, but restoration of flow in the epicardial infarct-related artery occurs promptly after PCI. At the bottom is a variety of methods for speeding the time to reperfusion along with the goals for the time intervals for the various components of the time delay. (Adapted from CP Cannon et al: *J Thromb Thrombol* 1:27, 1994.)

MANAGEMENT IN THE EMERGENCY DEPARTMENT

In the Emergency Department, the goals for the management of patients with suspected STEMI include control of cardiac discomfort, rapid identification of patients who are candidates for urgent reperfusion therapy, triage of lower-risk patients to the appropriate location in the hospital, and avoidance of inappropriate discharge of patients with STEMI. Many aspects of the treatment of STEMI are initiated in the Emergency Department and then continued during the in-hospital phase of management (Fig. 295-4). The overarching goal is to minimize the time from first medical contact to initiation of reperfusion therapy. This may involve transfer from a non-PCI hospital to one that is PCI capable, with a goal of initiating PCI within 120 min of first medical contact (Fig. 295-4).

Aspirin is essential in the management of patients with suspected STEMI and is effective across the entire spectrum of acute coronary syndromes (Fig. 295-1). Rapid inhibition of cyclooxygenase-1 in platelets followed by a reduction of thromboxane A_2 levels is achieved by buccal absorption of a chewed 160–325-mg tablet in the Emergency Department. This measure should be followed by daily oral administration of aspirin in a dose of 75–162 mg.

In patients whose arterial O_2 saturation is normal, supplemental O_2 is of limited if any clinical benefit and therefore is not cost-effective. However, when hypoxemia is present, O_2 should be administered by nasal prongs or face mask (2–4 L/min) for the first 6–12 h after infarction; the patient should then be reassessed to determine if there is a continued need for such treatment.

CONTROL OF DISCOMFORT

Sublingual *nitroglycerin* can be given safely to most patients with STEMI. Up to three doses of 0.4 mg should be administered at about 5-min intervals. In addition to diminishing or abolishing chest discomfort, nitroglycerin may be capable of both decreasing myocardial oxygen demand (by lowering preload) and increasing myocardial oxygen supply (by dilating infarct-related coronary vessels or collateral vessels). In patients whose initially favorable response to sublingual nitroglycerin is followed by the return of chest discomfort, particularly

if accompanied by other evidence of ongoing ischemia such as further ST-segment or T-wave shifts, the use of intravenous nitroglycerin should be considered. Therapy with nitrates should be avoided in patients who present with low systolic arterial pressure (<90 mmHg) or in whom there is clinical suspicion of RV infarction (inferior infarction on ECG, elevated jugular venous pressure, clear lungs, and hypotension). Nitrates should not be administered to patients who have taken a phosphodiesterase-5 inhibitor for erectile dysfunction within the preceding 24 h, because it may potentiate the hypotensive effects of nitrates. An idiosyncratic reaction to nitrates, consisting of sudden marked hypotension, sometimes occurs but can usually be reversed promptly by the rapid administration of intravenous atropine.

Morphine is a very effective analgesic for the pain associated with STEMI. However, it may reduce sympathetically mediated arteriolar and venous constriction, and the resulting venous pooling may reduce cardiac output and arterial pressure. These hemodynamic disturbances usually respond promptly to elevation of the legs, but in some patients, volume expansion with intravenous saline is required. The patient may experience diaphoresis and nausea, but these events usually pass and are replaced by a feeling of well-being associated with the relief of pain. Morphine also has a vagotonic effect and may cause bradycardia or advanced degrees of heart block, particularly in patients with inferior infarction. These side effects usually respond to atropine (0.5 mg intravenously). Morphine is routinely administered by repetitive (every 5 min) intravenous injection of small doses (2–4 mg), rather than by the subcutaneous administration of a larger quantity, because absorption may be unpredictable by the latter route.

Intravenous *beta blockers* are also useful in the control of the pain of STEMI. These drugs control pain effectively in some patients, presumably by diminishing myocardial O_2 demand and hence ischemia. More important, there is evidence that intravenous beta blockers reduce the risks of reinfarction and ventricular fibrillation (see “Beta-Adrenoceptor Blockers” below). However, patient selection is important when considering beta blockers for STEMI. Oral beta blocker therapy should be initiated in the first 24 h for patients who do not have any of the following: (1) signs of heart failure, (2) evidence of a low-output state, (3) increased risk for cardiogenic shock, or (4) other

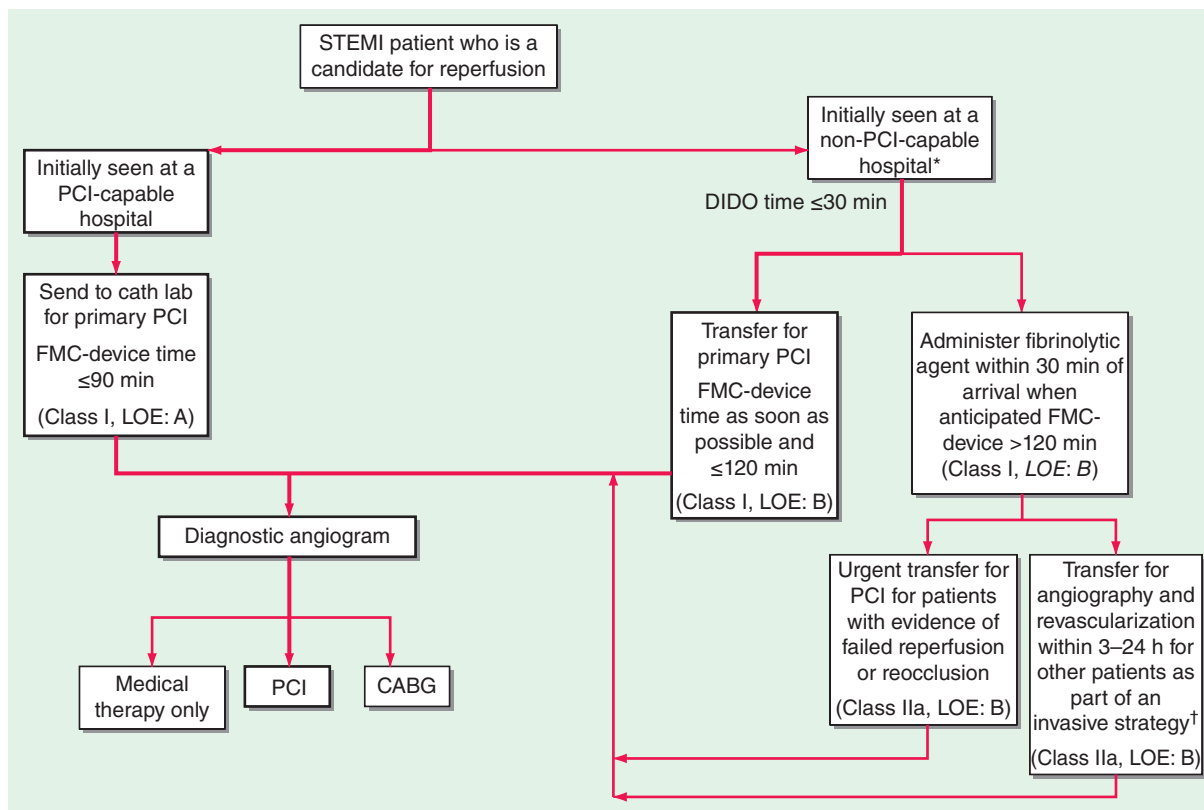


FIGURE 295-4 Reperfusion therapy for patients with ST-segment elevation myocardial infarction (STEMI). The **bold arrows** and **boxes** are the preferred strategies. Performance of percutaneous coronary intervention (PCI) is dictated by an anatomically appropriate culprit stenosis. *Patients with cardiogenic shock or severe heart failure initially seen at a non-PCI-capable hospital should be transferred for cardiac catheterization and revascularization as soon as possible, irrespective of time delay from myocardial infarction (MI) onset (Class I, LOE: B). †Angiography and revascularization should not be performed within the first 2 to 3 hours after administration of fibrinolytic therapy. CABG, coronary artery bypass graft; DIDO, door-in-door-out; FMC, first medical contact; LOE, level of evidence; STEMI, ST-elevation myocardial infarction. (Adapted with permission from P O'Gara et al: *Circulation* 127:e362, 2013.)

relative contraindications to beta blockade (PR interval greater than 0.24 seconds, second- or third-degree heart block, active asthma, or reactive airway disease). A commonly employed regimen is metoprolol, 5 mg every 2–5 min for a total of three doses, provided the patient has a heart rate >60 beats/min, systolic pressure >100 mmHg, a PR interval <0.24 s, and rales that are no higher than 10 cm up from the diaphragm. Fifteen minutes after the last intravenous dose, an oral regimen is initiated of 50 mg every 6 h for 48 h, followed by 100 mg every 12 h.

Unlike beta blockers, calcium antagonists are of little value in the acute setting, and there is evidence that short-acting dihydropyridines may be associated with an increased mortality risk.

MANAGEMENT STRATEGIES

The primary tool for screening patients and making triage decisions is the initial 12-lead ECG. When ST-segment elevation of at least 2 mm in two contiguous precordial leads and 1 mm in two adjacent limb leads is present, a patient should be considered a candidate for reperfusion therapy (Fig. 295-4). The process of selecting patients for fibrinolysis versus primary PCI (angioplasty or stenting; Chap. 296e) is discussed below. In the absence of ST-segment elevation, fibrinolysis is not helpful, and evidence exists suggesting that it may be harmful.

LIMITATION OF INFARCT SIZE

The quantity of myocardium that becomes necrotic as a consequence of a coronary artery occlusion is determined by factors other than just the site of occlusion. While the central zone of the infarct contains necrotic tissue that is irretrievably lost, the fate of the surrounding ischemic myocardium (ischemic penumbra) may be improved by timely restoration of coronary perfusion, reduction of myocardial

O₂ demands, prevention of the accumulation of noxious metabolites, and blunting of the impact of mediators of reperfusion injury (e.g., calcium overload and oxygen-derived free radicals). Up to one-third of patients with STEMI may achieve *spontaneous* reperfusion of the infarct-related coronary artery within 24 h and experience improved healing of infarcted tissue. Reperfusion, either pharmacologically (by fibrinolysis) or by PCI, accelerates the opening of infarct-related arteries in those patients in whom spontaneous fibrinolysis ultimately would have occurred and also greatly increases the number of patients in whom restoration of flow in the infarct-related artery is accomplished. Timely restoration of flow in the epicardial infarct-related artery combined with improved perfusion of the downstream zone of infarcted myocardium results in a limitation of infarct size. Protection of the ischemic myocardium by the maintenance of an optimal balance between myocardial O₂ supply and demand through pain control, treatment of congestive heart failure (CHF), and minimization of tachycardia and hypertension extends the “window” of time for the salvage of myocardium by reperfusion strategies.

Glucocorticoids and nonsteroidal anti-inflammatory agents, with the exception of aspirin, should be avoided in patients with STEMI. They can impair infarct healing and increase the risk of myocardial rupture, and their use may result in a larger infarct scar. In addition, they can increase coronary vascular resistance, thereby potentially reducing flow to ischemic myocardium.

PRIMARY PERCUTANEOUS CORONARY INTERVENTION

(See also Chap. 296e) PCI, usually angioplasty and/or stenting without preceding fibrinolysis, referred to as *primary PCI*, is effective in restoring perfusion in STEMI when carried out on an emergency basis in the first few hours of MI. It has the advantage of being applicable to patients who have contraindications to fibrinolytic therapy (see below)

but otherwise are considered appropriate candidates for reperfusion. It appears to be more effective than fibrinolysis in opening occluded coronary arteries and, *when performed by experienced operators in dedicated medical centers*, is associated with better short-term and long-term clinical outcomes. Compared with fibrinolysis, primary PCI is generally preferred when the diagnosis is in doubt, cardiogenic shock is present, bleeding risk is increased, or symptoms have been present for at least 2–3 h when the clot is more mature and less easily lysed by fibrinolytic drugs. However, PCI is expensive in terms of personnel and facilities, and its applicability is limited by its availability, around the clock, in only a minority of hospitals (Fig. 295-4).

FIBRINOLYSIS

If no contraindications are present (see below), fibrinolytic therapy should ideally be initiated within 30 min of presentation (i.e., door-to-needle time \leq 30 min). The principal goal of fibrinolysis is prompt restoration of full coronary arterial patency. The fibrinolytic agents tissue plasminogen activator (tPA), streptokinase, tenecteplase (TNK), and reteplase (rPA) have been approved by the U.S. Food and Drug Administration for intravenous use in patients with STEMI. These drugs all act by promoting the conversion of plasminogen to plasmin, which subsequently lyses fibrin thrombi. Although considerable emphasis was first placed on a distinction between more fibrin-specific agents, such as tPA, and non-fibrin-specific agents, such as streptokinase, it is now recognized that these differences are only relative, as some degree of systemic fibrinolysis occurs with the former agents. TNK and rPA are referred to as *bolus fibrinolytics* since their administration does not require a prolonged intravenous infusion.

When assessed angiographically, flow in the culprit coronary artery is described by a simple qualitative scale called the *Thrombolysis in Myocardial Infarction (TIMI) grading system*: grade 0 indicates complete occlusion of the infarct-related artery; grade 1 indicates some penetration of the contrast material beyond the point of obstruction but without perfusion of the distal coronary bed; grade 2 indicates perfusion of the entire infarct vessel into the distal bed, but with flow that is delayed compared with that of a normal artery; and grade 3 indicates full perfusion of the infarct vessel with normal flow. The latter is the goal of reperfusion therapy, because full perfusion of the infarct-related coronary artery yields far better results in terms of limiting infarct size, maintenance of LV function, and reduction of both short- and long-term mortality rates. Additional methods of angiographic assessment of the efficacy of fibrinolysis include counting the number of frames on the cine film required for dye to flow from the origin of the infarct-related artery to a landmark in the distal vascular bed (*TIMI frame count*) and determining the rate of entry and exit of contrast dye from the microvasculature in the myocardial infarct zone (*TIMI myocardial perfusion grade*). These methods have an even tighter correlation with outcomes after STEMI than the more commonly employed TIMI flow grade.

tPA and the other relatively fibrin-specific plasminogen activators, rPA and TNK, are more effective than streptokinase at restoring full perfusion—i.e., TIMI grade 3 coronary flow—and have a small edge in improving survival as well. The current recommended regimen of tPA consists of a 15-mg bolus followed by 50 mg intravenously over the first 30 min, followed by 35 mg over the next 60 min. Streptokinase is administered as 1.5 million units (MU) intravenously over 1 h. rPA is administered in a double-bolus regimen consisting of a 10-MU bolus given over 2–3 min, followed by a second 10-MU bolus 30 min later. TNK is given as a single weight-based intravenous bolus of 0.53 mg/kg over 10 s. In addition to the fibrinolytic agents discussed earlier, pharmacologic reperfusion typically involves adjunctive antiplatelet and antithrombotic drugs, as discussed subsequently.

Clear contraindications to the use of fibrinolytic agents include a history of cerebrovascular hemorrhage at any time, a nonhemorrhagic stroke or other cerebrovascular event within the past year, marked hypertension (a reliably determined systolic arterial pressure >180 mmHg and/or a diastolic pressure >110 mmHg) at any time during the acute presentation, suspicion of aortic dissection, and active internal bleeding (excluding menses). While advanced age is associated with

an increase in hemorrhagic complications, the benefit of fibrinolytic therapy in the elderly appears to justify its use if no other contraindications are present and the amount of myocardium in jeopardy appears to be substantial.

Relative contraindications to fibrinolytic therapy, which require assessment of the risk-to-benefit ratio, include current use of anti-coagulants (international normalized ratio ≥ 2), a recent (<2 weeks) invasive or surgical procedure or prolonged (>10 min) cardiopulmonary resuscitation, known bleeding diathesis, pregnancy, a hemorrhagic ophthalmic condition (e.g., hemorrhagic diabetic retinopathy), active peptic ulcer disease, and a history of severe hypertension that is currently adequately controlled. Because of the risk of an allergic reaction, patients should not receive streptokinase if that agent had been received within the preceding 5 days to 2 years.

Allergic reactions to streptokinase occur in $\sim 2\%$ of patients who receive it. While a minor degree of hypotension occurs in 4–10% of patients given this agent, marked hypotension occurs, although rarely, in association with severe allergic reactions.

Hemorrhage is the most frequent and potentially the most serious complication. Because bleeding episodes that require transfusion are more common when patients require invasive procedures, unnecessary venous or arterial interventions should be avoided in patients receiving fibrinolytic agents. Hemorrhagic stroke is the most serious complication and occurs in $\sim 0.5\text{--}0.9\%$ of patients being treated with these agents. This rate increases with advancing age, with patients >70 years experiencing roughly twice the rate of intracranial hemorrhage as those <65 years. Large-scale trials have suggested that the rate of intracranial hemorrhage with tPA or rPA is slightly higher than with streptokinase.

INTEGRATED REPERFUSION STRATEGY

Evidence has emerged that suggests PCI plays an increasingly important role in the management of STEMI. Prior approaches that segregated the pharmacologic and catheter-based approaches to reperfusion have now been replaced with an integrated approach to triage and transfer of STEMI patients to receive PCI (Fig. 295-4). To achieve the degree of integration required to care for a patient with STEMI, all communities should create and maintain a regional system of STEMI care that includes assessment and continuous quality improvement of emergency medical services and hospital-based activities.

Cardiac catheterization and coronary angiography should be carried out after fibrinolytic therapy if there is evidence of either (1) failure of reperfusion (persistent chest pain and ST-segment elevation >90 min), in which case a *rescue PCI* should be considered; or (2) coronary artery reocclusion (re-elevation of ST segments and/or recurrent chest pain) or the development of recurrent ischemia (such as recurrent angina in the early hospital course or a positive exercise stress test before discharge), in which case an *urgent PCI* should be considered. Routine angiography and *elective PCI* even in asymptomatic patients following administration of fibrinolytic therapy are used with less frequency, given the numerous technologic advances that have occurred in the catheterization laboratory and the increasing number of skilled interventionalists. Coronary artery bypass surgery should be reserved for patients whose coronary anatomy is unsuited to PCI but in whom revascularization appears to be advisable because of extensive jeopardized myocardium or recurrent ischemia.

HOSPITAL PHASE MANAGEMENT

CORONARY CARE UNITS

These units are routinely equipped with a system that permits continuous monitoring of the cardiac rhythm of each patient and hemodynamic monitoring in selected patients. Defibrillators, respirators, noninvasive transthoracic pacemakers, and facilities for introducing pacing catheters and flow-directed balloon-tipped catheters are also usually available. Equally important is the organization of a highly trained team of nurses who can recognize arrhythmias; adjust the dosage of antiarrhythmic, vasoactive, and anticoagulant drugs; and perform cardiac resuscitation, including electroshock, when necessary.

Patients should be admitted to a coronary care unit early in their illness when it is expected that they will derive benefit from the sophisticated and expensive care provided. The availability of electrocardiographic monitoring and trained personnel outside the coronary care unit has made it possible to admit lower-risk patients (e.g., those not hemodynamically compromised and without active arrhythmias) to “intermediate care units.”

The duration of stay in the coronary care unit is dictated by the ongoing need for intensive care. If symptoms are controlled with oral therapy, patients may be transferred out of the coronary care unit. Also, patients who have a confirmed STEMI but who are considered to be at low risk (no prior infarction and no persistent chest discomfort, CHF, hypotension, or cardiac arrhythmias) may be safely transferred out of the coronary care unit within 24 h.

Activity Factors that increase the work of the heart during the initial hours of infarction may increase the size of the infarct. Therefore, patients with STEMI should be kept at bed rest for the first 6–12 h. However, in the absence of complications, patients should be encouraged, under supervision, to resume an upright posture by dangling their feet over the side of the bed and sitting in a chair within the first 24 h. This practice is psychologically beneficial and usually results in a reduction in the pulmonary capillary wedge pressure. In the absence of hypotension and other complications, by the second or third day, patients typically are ambulating in their room with increasing duration and frequency, and they may shower or stand at the sink to bathe. By day 3 after infarction, patients should be increasing their ambulation progressively to a goal of 185 m (600 ft) at least three times a day.

Diet Because of the risk of emesis and aspiration soon after STEMI, patients should receive either nothing or only clear liquids by mouth for the first 4–12 h. The typical coronary care unit diet should provide ≤30% of total calories as fat and have a cholesterol content of ≤300 mg/d. Complex carbohydrates should make up 50–55% of total calories. Portions should not be unusually large, and the menu should be enriched with foods that are high in potassium, magnesium, and fiber, but low in sodium. Diabetes mellitus and hypertriglyceridemia are managed by restriction of concentrated sweets in the diet.

Bowel Management Bed rest and the effect of the narcotics used for the relief of pain often lead to constipation. A bedside commode rather than a bedpan, a diet rich in bulk, and the routine use of a stool softener such as dioctyl sodium sulfosuccinate (200 mg/d) are recommended. If the patient remains constipated despite these measures, a laxative can be prescribed. Contrary to prior belief, it is safe to perform a gentle rectal examination on patients with STEMI.

Sedation Many patients require sedation during hospitalization to withstand the period of enforced inactivity with tranquility. Diazepam (5 mg), oxazepam (15–30 mg), or lorazepam (0.5–2 mg), given three to four times daily, is usually effective. An additional dose of any of the above medications may be given at night to ensure adequate sleep. Attention to this problem is especially important during the first few days in the coronary care unit, where the atmosphere of 24-h vigilance may interfere with the patient’s sleep. However, sedation is no substitute for reassuring, quiet surroundings. Many drugs used in the coronary care unit, such as atropine, H₂ blockers, and narcotics, can produce delirium, particularly in the elderly. This effect should not be confused with agitation, and it is wise to conduct a thorough review of the patient’s medications before arbitrarily prescribing additional doses of anxiolytics.

PHARMACOTHERAPY

ANTITHROMBOTIC AGENTS

The use of antiplatelet and anticoagulant therapy during the initial phase of STEMI is based on extensive laboratory and clinical evidence that thrombosis plays an important role in the pathogenesis of this condition. The primary goal of treatment with antiplatelet and anticoagulant agents is to maintain patency of the infarct-related artery, in conjunction with reperfusion strategies. A secondary goal is to reduce the patient’s tendency to thrombosis and, thus, the likelihood of mural

thrombus formation or deep venous thrombosis, either of which could result in pulmonary embolization. The degree to which antiplatelet and anticoagulant therapy achieves these goals partly determines how effectively it reduces the risk of mortality from STEMI.

As noted previously (see “Management in the Emergency Department” earlier), aspirin is the standard antiplatelet agent for patients with STEMI. The most compelling evidence for the benefits of antiplatelet therapy (mainly with aspirin) in STEMI is found in the comprehensive overview by the Antiplatelet Trialists’ Collaboration. Data from nearly 20,000 patients with MI enrolled in 15 randomized trials were pooled and revealed a relative reduction of 27% in the mortality rate, from 14.2% in control patients to 10.4% in patients receiving antiplatelet agents.

Inhibitors of the P2Y₁₂ ADP receptor prevent activation and aggregation of platelets. The addition of the P2Y₁₂ inhibitor clopidogrel to background treatment with aspirin to STEMI patients reduces the risk of clinical events (death, reinfarction, stroke) and, in patients receiving fibrinolytic therapy, has been shown to prevent reocclusion of a successfully reperfused infarct artery. New P2Y₁₂ ADP receptor antagonists, such as prasugrel and ticagrelor, are more effective than clopidogrel in preventing ischemic complications in STEMI patients undergoing PCI, but are associated with an increased risk of bleeding. Glycoprotein IIb/IIIa receptor inhibitors appear useful for preventing thrombotic complications in patients with STEMI undergoing PCI.

The standard anticoagulant agent used in clinical practice is unfractionated heparin (UFH). The available data suggest that when UFH is added to a regimen of aspirin and a non-fibrin-specific thrombolytic agent such as streptokinase, additional mortality benefit occurs (about 5 lives saved per 1000 patients treated). It appears that the immediate administration of intravenous UFH, in addition to a regimen of aspirin and relatively fibrin-specific fibrinolytic agents (tPA, rPA, or TNK), helps to maintain patency of the infarct-related artery. This effect is achieved at the cost of a small increased risk of bleeding. The recommended dose of UFH is an initial bolus of 60 U/kg (maximum 4000 U) followed by an initial infusion of 12 U/kg per hour (maximum 1000 U/h). The activated partial thromboplastin time during maintenance therapy should be 1.5–2 times the control value.

Alternatives to UFH for anticoagulation of patients with STEMI are the low-molecular-weight heparin (LMWH) preparations, a synthetic version of the critical pentasaccharide sequence (fondaparinux), and the direct antithrombin bivalirudin. Advantages of LMWHs include high bioavailability permitting administration subcutaneously, reliable anticoagulation without monitoring, and greater antiXa:IIa activity. Enoxaparin has been shown to reduce significantly the composite endpoints of death/nonfatal reinfarction and death/nonfatal reinfarction/urgent revascularization compared with UFH in STEMI patients who receive fibrinolysis. Treatment with enoxaparin is associated with higher rates of serious bleeding, but net clinical benefit—a composite endpoint that combines efficacy and safety—still favors enoxaparin over UFH. Interpretation of the data on fondaparinux is difficult because of the complex nature of the pivotal clinical trial evaluating it in STEMI (OASIS-6). Fondaparinux appears superior to placebo in STEMI patients not receiving reperfusion therapy, but its relative efficacy and safety compared with UFH is less certain. Due to the risk of catheter thrombosis, fondaparinux should not be used alone at the time of coronary angiography and PCI but should be combined with another anticoagulant with antithrombin activity such as UFH or bivalirudin. Contemporary trials of bivalirudin used an open-label design to evaluate its efficacy and safety compared with UFH plus a glycoprotein IIb/IIIa inhibitor. Bivalirudin was associated with a lower rate of bleeding, largely driven by reductions in vascular access site hematomas ≥5 cm or the administration of blood transfusions.

Patients with an anterior location of the infarction, severe LV dysfunction, heart failure, a history of embolism, two-dimensional echocardiographic evidence of mural thrombus, or atrial fibrillation are at increased risk of systemic or pulmonary thromboembolism. Such individuals should receive full therapeutic levels of anticoagulant therapy (LMWH or UFH) while hospitalized, followed by at least 3 months of warfarin therapy.

BETA-ADRENOCEPTOR BLOCKERS

The benefits of beta blockers in patients with STEMI can be divided into those that occur immediately when the drug is given acutely and those that accrue over the long term when the drug is given for secondary prevention after an infarction. Acute intravenous beta blockade improves the myocardial O₂ supply-demand relationship, decreases pain, reduces infarct size, and decreases the incidence of serious ventricular arrhythmias. In patients who undergo fibrinolysis soon after the onset of chest pain, no incremental reduction in mortality rate is seen with beta blockers, but recurrent ischemia and reinfarction are reduced.

Thus, beta-blocker therapy after STEMI is useful for most patients (including those treated with an angiotensin-converting enzyme [ACE] inhibitor) except those in whom it is specifically contraindicated (patients with heart failure or severely compromised LV function, heart block, orthostatic hypotension, or a history of asthma) and perhaps those whose excellent long-term prognosis (defined as an expected mortality rate of <1% per year, patients <55 years, no previous MI, with normal ventricular function, no complex ventricular ectopy, and no angina) markedly diminishes any potential benefit.

INHIBITION OF THE RENIN-ANGIOTENSIN-ALDOSTERONE SYSTEM

ACE inhibitors reduce the mortality rate after STEMI, and the mortality benefits are additive to those achieved with aspirin and beta blockers. The maximum benefit is seen in high-risk patients (those who are elderly or who have an anterior infarction, a prior infarction, and/or globally depressed LV function), but evidence suggests that a short-term benefit occurs when ACE inhibitors are prescribed unselectively to all hemodynamically stable patients with STEMI (i.e., those with a systolic pressure >100 mmHg). The mechanism involves a reduction in ventricular remodeling after infarction (see “Ventricular Dysfunction” later) with a subsequent reduction in the risk of CHF. The rate of recurrent infarction may also be lower in patients treated chronically with ACE inhibitors after infarction.

Before hospital discharge, LV function should be assessed with an imaging study. ACE inhibitors should be continued indefinitely in patients who have clinically evident CHF, in patients in whom an imaging study shows a reduction in global LV function or a large regional wall motion abnormality, or in those who are hypertensive.

Angiotensin receptor blockers (ARBs) should be administered to STEMI patients who are intolerant of ACE inhibitors and who have either clinical or radiologic signs of heart failure. Long-term aldosterone blockade should be prescribed for STEMI patients without significant renal dysfunction (creatinine ≥ 2.5 mg/dL in men and ≥ 2.0 mg/dL in women) or hyperkalemia (potassium ≥ 5.0 mEq/L) who are already receiving therapeutic doses of an ACE inhibitor, have an LV ejection fraction $\leq 40\%$, and have either symptomatic heart failure or diabetes mellitus. A multidrug regimen for inhibiting the renin-angiotensin-aldosterone system has been shown to reduce both heart failure–related and sudden cardiac death–related cardiovascular mortality after STEMI, but has not been as thoroughly explored as ACE inhibitors in STEMI patients.

OTHER AGENTS

Favorable effects on the ischemic process and ventricular remodeling (see below) previously led many physicians to routinely use *intravenous nitroglycerin* (5–10 $\mu\text{g}/\text{min}$ initial dose and up to 200 $\mu\text{g}/\text{min}$ as long as hemodynamic stability is maintained) for the first 24–48 h after the onset of infarction. However, the benefits of routine use of intravenous nitroglycerin are less in the contemporary era where beta-adrenoceptor blockers and ACE inhibitors are routinely prescribed for patients with STEMI.

Results of multiple trials of different calcium antagonists have failed to establish a role for these agents in the treatment of most patients with STEMI. Therefore, the routine use of calcium antagonists cannot be recommended. Strict control of blood glucose in diabetic patients with STEMI has been shown to reduce the mortality rate. Serum magnesium should be measured in all patients on admission, and any demonstrated deficits should be corrected to minimize the risk of arrhythmias.

COMPLICATIONS AND THEIR MANAGEMENT

VENTRICULAR DYSFUNCTION

After STEMI, the left ventricle undergoes a series of changes in shape, size, and thickness in both the infarcted and noninfarcted segments. This process is referred to as *ventricular remodeling* and generally precedes the development of clinically evident CHF in the months to years after infarction. Soon after STEMI, the left ventricle begins to dilate. Acutely, this results from expansion of the infarct, i.e., slippage of muscle bundles, disruption of normal myocardial cells, and tissue loss within the necrotic zone, resulting in disproportionate thinning and elongation of the infarct zone. Later, lengthening of the noninfarcted segments occurs as well. The overall chamber enlargement that occurs is related to the size and location of the infarct, with greater dilation following infarction of the anterior wall and apex of the left ventricle and causing more marked hemodynamic impairment, more frequent heart failure, and a poorer prognosis. Progressive dilation and its clinical consequences may be ameliorated by therapy with ACE inhibitors and other vasodilators (e.g., nitrates). In patients with an ejection fraction <40%, regardless of whether or not heart failure is present, ACE inhibitors or ARBs should be prescribed (see “Inhibition of the Renin-Angiotensin-Aldosterone System” earlier).

HEMODYNAMIC ASSESSMENT

Pump failure is now the primary cause of in-hospital death from STEMI. The extent of infarction correlates well with the degree of pump failure and with mortality, both early (within 10 days of infarction) and later. The most common clinical signs are pulmonary rales and S₃ and S₄ gallop sounds. Pulmonary congestion is also frequently seen on the chest roentgenogram. Elevated LV filling pressure and elevated pulmonary artery pressure are the characteristic hemodynamic findings, but these findings may result from a reduction of ventricular compliance (diastolic failure) and/or a reduction of stroke volume with secondary cardiac dilation (systolic failure) ([Chap. 279](#)).

A classification originally proposed by Killip divides patients into four groups: class I, no signs of pulmonary or venous congestion; class II, moderate heart failure as evidenced by rales at the lung bases, S₃ gallop, tachypnea, or signs of failure of the right side of the heart, including venous and hepatic congestion; class III, severe heart failure, pulmonary edema; and class IV, shock with systolic pressure <90 mmHg and evidence of peripheral vasoconstriction, peripheral cyanosis, mental confusion, and oliguria. When this classification was established in 1967, the expected hospital mortality rate of patients in these classes was as follows: class I, 0–5%; class II, 10–20%; class III, 35–45%; and class IV, 85–95%. With advances in management, the mortality rate in each class has fallen, perhaps by as much as one-third to one-half.

Hemodynamic evidence of abnormal global LV function appears when contraction is seriously impaired in 20–25% of the left ventricle. Infarction of $\geq 40\%$ of the left ventricle usually results in cardiogenic shock ([Chap. 326](#)). Positioning of a balloon flotation (Swan-Ganz) catheter in the pulmonary artery permits monitoring of LV filling pressure; this technique is useful in patients who exhibit hypotension and/or clinical evidence of CHF. Cardiac output can also be determined with a pulmonary artery catheter. With the addition of intra-arterial pressure monitoring, systemic vascular resistance can be calculated as a guide to adjusting vasopressor and vasodilator therapy. Some patients with STEMI have markedly elevated LV filling pressures (>22 mmHg) and normal cardiac indices (2.6–3.6 L/[min·m²]), while others have relatively low LV filling pressures (<15 mmHg) and reduced cardiac indices. The former patients usually benefit from diuresis, while the latter may respond to volume expansion.

HYPVOLEMIA

This is an easily corrected condition that may contribute to the hypotension and vascular collapse associated with STEMI in some patients. It may be secondary to previous diuretic use, to reduced fluid intake during the early stages of the illness, and/or to vomiting associated with pain or medications. Consequently, hypovolemia should be identified and corrected in patients with STEMI and hypotension before

more vigorous forms of therapy are begun. Central venous pressure reflects RV rather than LV filling pressure and is an inadequate guide for adjustment of blood volume, because LV function is almost always affected much more adversely than RV function in patients with STEMI. The optimal LV filling or pulmonary artery wedge pressure may vary considerably among patients. Each patient's ideal level (generally ~20 mmHg) is reached by cautious fluid administration during careful monitoring of oxygenation and cardiac output. Eventually, the cardiac output level plateaus, and further increases in LV filling pressure only increase congestive symptoms and decrease systemic oxygenation without raising arterial pressure.

TREATMENT CONGESTIVE HEART FAILURE

The management of CHF in association with STEMI is similar to that of acute heart failure secondary to other forms of heart disease (avoidance of hypoxemia, diuresis, afterload reduction, inotropic support) (**Chap. 279**), except that the benefits of digitalis administration to patients with STEMI are unimpressive. By contrast, diuretic agents are extremely effective, as they diminish pulmonary congestion in the presence of systolic and/or diastolic heart failure. LV filling pressure falls and orthopnea and dyspnea improve after the intravenous administration of furosemide or other loop diuretics. These drugs should be used with caution, however, as they can result in a massive diuresis with associated decreases in plasma volume, cardiac output, systemic blood pressure, and, hence, coronary perfusion. Nitrates in various forms may be used to decrease preload and congestive symptoms. Oral isosorbide dinitrate, topical nitroglycerin ointment, and intravenous nitroglycerin all have the advantage over a diuretic of lowering preload through venodilation without decreasing the total plasma volume. In addition, nitrates may improve ventricular compliance if ischemia is present, as ischemia causes an elevation of LV filling pressure. Vasodilators must be used with caution to prevent serious hypotension. As noted earlier, ACE inhibitors are an ideal class of drugs for management of ventricular dysfunction after STEMI, especially for the long term. (See "Inhibition of the Renin-Angiotensin-Aldosterone System" earlier.)

CARDIOGENIC SHOCK

Prompt reperfusion, efforts to reduce infarct size and treatment of ongoing ischemia and other complications of MI appear to have reduced the incidence of cardiogenic shock from 20% to about 7%. Only 10% of patients with this condition present with it on admission, while 90% develop it during hospitalization. Typically, patients who develop cardiogenic shock have severe multivessel coronary artery disease with evidence of "piecemeal" necrosis extending outward from the original infarct zone. **The evaluation and management of cardiogenic shock and severe power failure after STEMI are discussed in detail in Chap. 326.**

RIGHT VENTRICULAR INFARCTION

Approximately one-third of patients with inferior infarction demonstrate at least a minor degree of RV necrosis. An occasional patient with inferoposterior LV infarction also has extensive RV infarction, and rare patients present with infarction limited primarily to the RV. Clinically significant RV infarction causes signs of severe RV failure (jugular venous distention, Kussmaul's sign, hepatomegaly [**Chap. 267**]) with or without hypotension. ST-segment elevations of right-sided precordial ECG leads, particularly lead V₄R, are frequently present in the first 24 h in patients with RV infarction. Two-dimensional echocardiography is helpful in determining the degree of RV dysfunction. Catheterization of the right side of the heart often reveals a distinctive hemodynamic pattern resembling constrictive pericarditis (steep right atrial "y" descent and an early diastolic dip and plateau in RV waveforms) (**Chap. 288**). Therapy consists of volume expansion to maintain adequate RV preload and efforts to improve LV performance

with attendant reduction in pulmonary capillary wedge and pulmonary arterial pressures.

ARRHYTHMIAS

(See also **Chaps. 274 and 276**) The incidence of arrhythmias after STEMI is higher in patients seen early after the onset of symptoms. The mechanisms responsible for infarction-related arrhythmias include autonomic nervous system imbalance, electrolyte disturbances, ischemia, and slowed conduction in zones of ischemic myocardium. An arrhythmia can usually be managed successfully if trained personnel and appropriate equipment are available when it develops. Since most deaths from arrhythmia occur during the first few hours after infarction, the effectiveness of treatment relates directly to the speed with which patients come under medical observation. The prompt management of arrhythmias constitutes a significant advance in the treatment of STEMI.

Ventricular Premature Beats Infrequent, sporadic ventricular premature depolarizations occur in almost all patients with STEMI and do not require therapy. Whereas in the past, frequent, multifocal, or early diastolic ventricular extrasystoles (so-called warning arrhythmias) were routinely treated with antiarrhythmic drugs to reduce the risk of development of ventricular tachycardia and ventricular fibrillation, pharmacologic therapy is now reserved for patients with sustained ventricular arrhythmias. Prophylactic antiarrhythmic therapy (either intravenous lidocaine early or oral agents later) is contraindicated for ventricular premature beats in the absence of clinically important ventricular tachyarrhythmias, because such therapy may actually increase the mortality rate. Beta-adrenoceptor blocking agents are effective in abolishing ventricular ectopic activity in patients with STEMI and in the prevention of ventricular fibrillation. As described earlier (see "Beta-Adrenoceptor Blockers"), they should be used routinely in patients without contraindications. In addition, hypokalemia and hypomagnesemia are risk factors for ventricular fibrillation in patients with STEMI; to reduce the risk, the serum potassium concentration should be adjusted to approximately 4.5 mmol/L and magnesium to about 2.0 mmol/L.

Ventricular Tachycardia and Fibrillation Within the first 24 h of STEMI, ventricular tachycardia and fibrillation can occur without prior warning arrhythmias. The occurrence of ventricular fibrillation can be reduced by prophylactic administration of intravenous lidocaine. However, prophylactic use of lidocaine has not been shown to reduce overall mortality from STEMI. In fact, in addition to causing possible noncardiac complications, lidocaine may predispose to an excess risk of bradycardia and asystole. For these reasons, and with earlier treatment of active ischemia, more frequent use of beta-blocking agents, and the nearly universal success of electrical cardioversion or defibrillation, routine prophylactic antiarrhythmic drug therapy is *no longer recommended*.

Sustained ventricular tachycardia that is well tolerated hemodynamically should be treated with an intravenous regimen of amiodarone (bolus of 150 mg over 10 min, followed by infusion of 1.0 mg/min for 6 h and then 0.5 mg/min) or procainamide (bolus of 15 mg/kg over 20–30 min; infusion of 1–4 mg/min); if it does not stop promptly, electroversion should be used (**Chap. 276**). An unsynchronized discharge of 200–300 J (monophasic waveform; approximately 50% of these energies with biphasic waveforms) is used immediately in patients with ventricular fibrillation or when ventricular tachycardia causes hemodynamic deterioration. Ventricular tachycardia or fibrillation that is refractory to electroshock may be more responsive after the patient is treated with epinephrine (1 mg intravenously or 10 mL of a 1:10,000 solution via the intracardiac route) or amiodarone (a 75–150-mg bolus).

Ventricular arrhythmias, including the unusual form of ventricular tachycardia known as torsades des pointes (**Chaps. 276 and 277**), may occur in patients with STEMI as a consequence of other concurrent problems (such as hypoxia, hypokalemia, or other electrolyte disturbances) or of the toxic effects of an agent being administered to the patient (such as digoxin or quinidine). A search for such secondary causes should always be undertaken.

Although the in-hospital mortality rate is increased, the long-term survival is excellent in patients who survive to hospital discharge after *primary* ventricular fibrillation; i.e., ventricular fibrillation that is a primary response to acute ischemia that occurs during the first 48 h and is not associated with predisposing factors such as CHF, shock, bundle branch block, or ventricular aneurysm. This result is in sharp contrast to the poor prognosis for patients who develop ventricular fibrillation *secondary* to severe pump failure. For patients who develop ventricular tachycardia or ventricular fibrillation late in their hospital course (i.e., after the first 48 h), the mortality rate is increased both in-hospital and during long-term follow-up. Such patients should be considered for electrophysiologic study and implantation of a cardioverter-defibrillator (ICD) (Chap. 276). A more challenging issue is the prevention of sudden cardiac death from ventricular fibrillation late after STEMI in patients who have not exhibited sustained ventricular tachyarrhythmias during their index hospitalization. An algorithm for selection of patients who warrant prophylactic implantation of an ICD is shown in Fig. 295-5.

Accelerated Idioventricular Rhythm Accelerated idioventricular rhythm (AIVR, “slow ventricular tachycardia”), a ventricular rhythm with a rate of 60–100 beats/min, often occurs transiently during fibrinolytic therapy at the time of reperfusion. For the most part, AIVR, whether it occurs in association with fibrinolytic therapy or spontaneously, is benign and does not presage the development of classic ventricular tachycardia. Most episodes of AIVR do not require treatment if the patient is monitored carefully, as degeneration into a more serious arrhythmia is rare.

Supraventricular Arrhythmias Sinus tachycardia is the most common supraventricular arrhythmia. If it occurs secondary to another cause

(such as anemia, fever, heart failure, or a metabolic derangement), the primary problem should be treated first. However, if it appears to be due to sympathetic overstimulation (e.g., as part of a hyperdynamic state), then treatment with a beta blocker is indicated. Other common arrhythmias in this group are atrial flutter and atrial fibrillation, which are often secondary to LV failure. Digoxin is usually the treatment of choice for supraventricular arrhythmias if heart failure is present. If heart failure is absent, beta blockers, verapamil, or diltiazem are suitable alternatives for controlling the ventricular rate, as they may also help to control ischemia. If the abnormal rhythm persists for >2 h with a ventricular rate >120 beats/min, or if tachycardia induces heart failure, shock, or ischemia (as manifested by recurrent pain or ECG changes), a synchronized electroshock (100–200 J monophasic waveform) should be used.

Accelerated junctional rhythms have diverse causes but may occur in patients with inferoposterior infarction. Digitalis excess must be ruled out. In some patients with severely compromised LV function, the loss of appropriately timed atrial systole results in a marked reduction of cardiac output. Right atrial or coronary sinus pacing is indicated in such instances.

Sinus Bradycardia Treatment of sinus bradycardia is indicated if hemodynamic compromise results from the slow heart rate. Atropine is the most useful drug for increasing heart rate and should be given intravenously in doses of 0.5 mg initially. If the rate remains <50–60 beats/min, additional doses of 0.2 mg, up to a total of 2.0 mg, may be given. Persistent bradycardia (<40 beats/min) despite atropine may be treated with electrical pacing. Isoproterenol should be avoided.

Atrioventricular and Intraventricular Conduction Disturbances (See also Chap. 274) Both the in-hospital mortality rate and the postdischarge mortality rate of patients who have complete atrioventricular (AV) block in association with anterior infarction are markedly higher than those of patients who develop AV block with inferior infarction. This difference is related to the fact that heart block in inferior infarction is commonly a result of increased vagal tone and/or the release of adenosine and therefore is transient. In anterior wall infarction, however, heart block is usually related to ischemic malfunction of the conduction system, which is commonly associated with extensive myocardial necrosis.

Temporary electrical pacing provides an effective means of increasing the heart rate of patients with bradycardia due to AV block. However, acceleration of the heart rate may have only a limited impact on prognosis in patients with anterior wall infarction and complete heart block in whom the large size of the infarct is the major factor determining outcome. It should be carried out if it improves hemodynamics. Pacing does appear to be beneficial in patients with inferoposterior infarction who have complete heart block associated with heart failure, hypotension, marked bradycardia, or significant ventricular ectopic activity. A subgroup of these patients, those with RV infarction, often respond poorly to ventricular pacing because of the loss of the atrial contribution to ventricular filling. In such patients, dual-chamber AV sequential pacing may be required.

External noninvasive pacing electrodes should be positioned in a “demand” mode for patients with sinus bradycardia (rate <50 beats/min) that is unresponsive to drug therapy, Mobitz II second-degree AV block, third-degree heart block, or bilateral bundle branch block (e.g., right bundle branch block plus left anterior fascicular block). Retrospective studies suggest that permanent pacing may reduce the long-term risk of sudden death due to bradyarrhythmias in the rare patient who develops combined persistent bifascicular and transient third-degree heart block during the acute phase of MI.

OTHER COMPLICATIONS

Recurrent Chest Discomfort Because recurrent or persistent ischemia often heralds extension of the original infarct or reinfarction in a new myocardial zone and is associated with a near tripling of mortality after STEMI, patients with these symptoms should be referred for prompt coronary arteriography and mechanical revascularization.

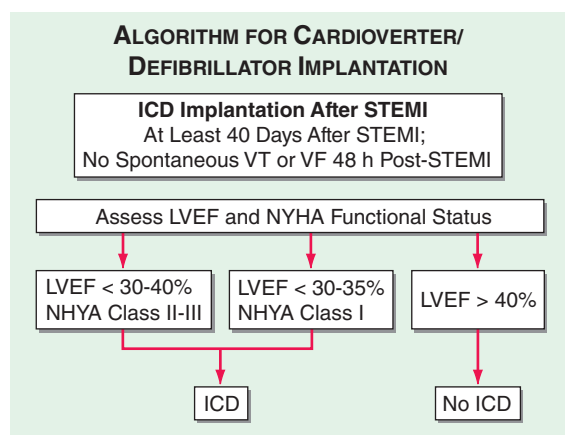


FIGURE 295-5 Algorithm for assessment of need for implantation of a cardioverter-defibrillator. The appropriate management is selected based on measurement of left ventricular ejection fraction and assessment of the New York Heart Association (NYHA) functional class. Patients with depressed left ventricular function at least 40 days after ST-segment elevation myocardial infarction (STEMI) are referred for insertion of an implantable cardioverter-defibrillator (ICD) if the left ventricular ejection fraction (LVEF) is <30–40% and they are in NYHA class II–III or if the LVEF is <30–35% and they are in NYHA class I functional status. Patients with preserved left ventricular function (LVEF >40%) do not receive an ICD regardless of NYHA functional class. All patients are treated with medical therapy after STEMI. VF, ventricular fibrillation; VT, ventricular tachycardia. (Adapted from data contained in DP Zipes et al: ACC/AHA/ESC 2006 guidelines for management of patients with ventricular arrhythmias and the prevention of sudden cardiac death; a report of the American College of Cardiology/American Heart Association Task Force and the European Society of Cardiology Committee for Practice Guidelines [Writing Committee to Develop Guidelines for Management of Patients with Ventricular Arrhythmias and the Prevention of Sudden Cardiac Death]. *J Am Coll Cardiol* 48:1064, 2006.)

1610 Administration of a fibrinolytic agent is an alternative to early mechanical revascularization.

Pericarditis (See also Chap. 288) Pericardial friction rubs and/or pericardial pain are frequently encountered in patients with STEMI involving the epicardium. This complication can usually be managed with aspirin (650 mg four times daily). It is important to diagnose the chest pain of pericarditis accurately, because failure to recognize it may lead to the erroneous diagnosis of recurrent ischemic pain and/or infarct extension, with resulting inappropriate use of anticoagulants, nitrates, beta blockers, or coronary arteriography. When it occurs, complaints of pain radiating to either trapezius muscle is helpful, because such a pattern of discomfort is typical of pericarditis but rarely occurs with ischemic discomfort. Anticoagulants potentially could cause tamponade in the presence of acute pericarditis (as manifested by either pain or persistent rub) and therefore should not be used unless there is a compelling indication.

Thromboembolism Clinically apparent thromboembolism complicates STEMI in ~10% of cases, but embolic lesions are found in 20% of patients in necropsy series, suggesting that thromboembolism is often clinically silent. Thromboembolism is considered to be an important contributing cause of death in 25% of patients with STEMI who die after admission to the hospital. Arterial emboli originate from LV mural thrombi, while most pulmonary emboli arise in the leg veins.

Thromboembolism typically occurs in association with large infarcts (especially anterior), CHF, and an LV thrombus detected by echocardiography. The incidence of arterial embolism from a clot originating in the ventricle at the site of an infarction is small but real. Two-dimensional echocardiography reveals LV thrombi in about one-third of patients with anterior wall infarction but in few patients with inferior or posterior infarction. Arterial embolism often presents as a major complication, such as hemiparesis when the cerebral circulation is involved or hypertension if the renal circulation is compromised. When a thrombus has been clearly demonstrated by echocardiographic or other techniques or when a large area of regional wall motion abnormality is seen even in the absence of a detectable mural thrombus, systemic anticoagulation should be undertaken (in the absence of contraindications), as the incidence of embolic complications appears to be markedly lowered by such therapy. The appropriate duration of therapy is unknown, but 3–6 months is probably prudent.

Left Ventricular Aneurysm The term *ventricular aneurysm* is usually used to describe *dyskinesis* or local expansile paradoxical wall motion. Normally functioning myocardial fibers must shorten more if stroke volume and cardiac output are to be maintained in patients with ventricular aneurysm; if they cannot, overall ventricular function is impaired. True aneurysms are composed of scar tissue and neither predispose to nor are associated with cardiac rupture.

The complications of LV aneurysm do not usually occur for weeks to months after STEMI; they include CHF, arterial embolism, and ventricular arrhythmias. Apical aneurysms are the most common and the most easily detected by clinical examination. The physical finding of greatest value is a double, diffuse, or displaced apical impulse. Ventricular aneurysms are readily detected by two-dimensional echocardiography, which may also reveal a mural thrombus in an aneurysm.

Rarely, myocardial rupture may be contained by a local area of pericardium, along with organizing thrombus and hematoma. Over time, this *pseudoaneurysm* enlarges, maintaining communication with the LV cavity through a narrow neck. Because a pseudoaneurysm often ruptures spontaneously, it should be surgically repaired if recognized.

POSTINFARCTION RISK STRATIFICATION AND MANAGEMENT

Many clinical and laboratory factors have been identified that are associated with an increase in cardiovascular risk after initial recovery from STEMI. Some of the most important factors include persistent ischemia (spontaneous or provoked), depressed LV ejection fraction (<40%), rales above the lung bases on physical examination or congestion on

chest radiograph, and symptomatic ventricular arrhythmias. Other features associated with increased risk include a history of previous MI, age >75, diabetes mellitus, prolonged sinus tachycardia, hypotension, ST-segment changes at rest without angina (“silent ischemia”), an abnormal signal-averaged ECG, nonpatency of the infarct-related coronary artery (if angiography is undertaken), and persistent advanced heart block or a new intraventricular conduction abnormality on the ECG. Therapy must be individualized on the basis of the relative importance of the risk(s) present.

The goal of preventing reinfarction and death after recovery from STEMI has led to strategies to evaluate risk after infarction. In stable patients, submaximal exercise stress testing may be carried out before hospital discharge to detect residual ischemia and ventricular ectopy and to provide the patient with a guideline for exercise in the early recovery period. Alternatively, or in addition, a maximal (symptom-limited) exercise stress test may be carried out 4–6 weeks after infarction. Evaluation of LV function is usually warranted as well. Recognition of a depressed LV ejection fraction by echocardiography or radionuclide ventriculography identifies patients who should receive medications to inhibit the renin-angiotensin-aldosterone system. Patients in whom angina is induced at relatively low workloads, those who have a large reversible defect on perfusion imaging or a depressed ejection fraction, those with demonstrable ischemia, and those in whom exercise provokes symptomatic ventricular arrhythmias should be considered at high risk for recurrent MI or death from arrhythmia (Fig. 295-5). Cardiac catheterization with coronary angiography and/or invasive electrophysiologic evaluation is advised.

Exercise tests also aid in formulating an individualized exercise prescription, which can be much more vigorous in patients who tolerate exercise without any of the previously mentioned adverse signs. In addition, pre-discharge stress testing may provide an important psychological benefit, building the patient’s confidence by demonstrating a reasonable exercise tolerance.

In many hospitals, a cardiac rehabilitation program with progressive exercise is initiated in the hospital and continued after discharge. Ideally, such programs should include an educational component that informs patients about their disease and its risk factors.

The usual duration of hospitalization for an uncomplicated STEMI is about 5 days. The remainder of the convalescent phase may be accomplished at home. During the first 1–2 weeks, the patient should be encouraged to increase activity by walking about the house and outdoors in good weather. Normal sexual activity may be resumed during this period. After 2 weeks, the physician must regulate the patient’s activity on the basis of exercise tolerance. Most patients will be able to return to work within 2–4 weeks.

SECONDARY PREVENTION

Various secondary preventive measures are at least partly responsible for the improvement in the long-term mortality and morbidity rates after STEMI. Long-term treatment with an antiplatelet agent (usually aspirin) after STEMI is associated with a 25% reduction in the risk of recurrent infarction, stroke, or cardiovascular mortality (36 fewer events for every 1000 patients treated). An alternative antiplatelet agent that may be used for secondary prevention in patients intolerant of aspirin is clopidogrel (75 mg orally daily). ACE inhibitors or ARBs and, in appropriate patients, aldosterone antagonists should be used indefinitely by patients with clinically evident heart failure, a moderate decrease in global ejection fraction, or a large regional wall motion abnormality to prevent late ventricular remodeling and recurrent ischemic events.

The chronic routine use of oral beta-adrenoceptor blockers for at least 2 years after STEMI is supported by well-conducted, placebo-controlled trials.

Evidence suggests that warfarin lowers the risk of late mortality and the incidence of reinfarction after STEMI. Most physicians prescribe aspirin routinely for all patients without contraindications and add warfarin for patients at increased risk of embolism (see “Thromboembolism” earlier). Several studies suggest that in patients

<75 years old a low dose of aspirin (75–81 mg/d) in combination with warfarin administered to achieve an international normalized ratio >2.0 is more effective than aspirin alone for preventing recurrent MI and embolic cerebrovascular accident. However, there is an increased risk of bleeding and a high rate of discontinuation of warfarin that has limited clinical acceptance of combination antithrombotic therapy. There is increased risk of bleeding when warfarin is added to dual antiplatelet therapy (aspirin and clopidogrel). However, patients who have had a stent implanted and have an indication for anticoagulation should receive dual antiplatelet therapies in combination with warfarin. Such patients should also receive a proton pump inhibitor to minimize the risk of gastrointestinal bleeding and should have regular monitoring of their hemoglobin levels and stool hematest while on combination antithrombotic therapy.

Finally, risk factors for *atherosclerosis* ([Chap. 265e](#)) should be discussed with the patient and, when possible, favorably modified.

David P. Faxon, Deepak L. Bhatt

Percutaneous transluminal coronary angioplasty (PTCA) was first introduced by Andreas Gruentzig in 1977 as an alternative to coronary bypass surgery. The concept was initially demonstrated by Charles Dotter in 1964 in peripheral vessels. The development of a small inelastic balloon catheter by Gruentzig allowed expansion of the technique into smaller peripheral and coronary vessels. Initial coronary experience was limited to single-vessel coronary disease and discrete proximal lesions due to the technical limitations of the equipment. Advances in technology and greater operator experience allowed the procedure to grow rapidly with expanded use in patients with more complex lesions and multivessel disease. The introduction of coronary stents in 1994 was one of the major advances in the field. These devices reduced acute complications and reduced by half the significant problem of restenosis (or recurrence of the stenosis). Further reductions in restenosis were achieved by the introduction of drug-eluting stents in 2003. These stents slowly release antiproliferative drugs directly into the plaque over a few months. Percutaneous coronary intervention (PCI) is the most common revascularization procedure in the United States and is performed more than twice as often as coronary artery bypass surgery: nearly 600,000 patients a year.

Interventional cardiology is a separate discipline in cardiology that requires a dedicated 1-year interventional cardiology fellowship following a 3-year general cardiology fellowship in order to obtain a separate board certification. The discipline has also expanded to include interventions for structural heart disease including treatment of congenital heart disease and valvular heart disease; it also includes interventions to treat peripheral vascular disease, including atherosclerotic and nonatherosclerotic lesions in the carotid, renal, aortic, and peripheral circulations.

TECHNIQUE

The initial procedure is performed in a similar manner as a diagnostic cardiac catheterization (Chap. 272). Arterial access is obtained via the femoral or radial artery. To prevent thrombotic complications during the procedure, patients who are anticipated to need an angioplasty

are given aspirin (325 mg) and may be given clopidogrel (loading dose of 300–600 mg), prasugrel (loading dose of 60 mg), or ticagrelor (loading dose of 180 mg) before the procedure. During the procedure, anticoagulation is achieved by administration of unfractionated heparin, enoxaparin (a low-molecular-weight heparin), or bivalirudin (a direct thrombin inhibitor). The latter has gained popularity due to the significant reduction in bleeding complications. In patients with ST-elevation myocardial infarction, high-risk acute coronary syndrome, or a large thrombus in the coronary artery, an intravenous glycoprotein IIb/IIIa inhibitor (abciximab, tirofiban, or eptifibatid) may also be given.

Following placement of an introducing sheath, preformed guiding catheters are used to cannulate selectively the origins of the coronary arteries. Through the guiding catheter, a flexible, steerable guidewire is negotiated down the coronary artery lumen using fluoroscopic guidance; it is then advanced through the stenosis and into the vessel beyond. This guidewire then serves as a “rail” over which angioplasty balloons, stents, or other therapeutic devices can be advanced to enlarge the narrowed segment of coronary artery. The artery is usually dilated with a balloon catheter followed by placement of a stent. The catheters and introducing sheath are removed and the artery manually held or closed using one of several femoral arterial closure devices to achieve hemostasis. Because PCI is performed under local anesthesia and mild sedation, it requires only a short (1-day) hospitalization or less.

Angioplasty works by stretching the artery and displacing the plaque away from the lumen, enlarging the entire vessel (Figs. 296e-1 and 296e-2). The procedure rarely results in embolization of atherosclerotic material. Owing to inelastic elements in the plaque, the stretching of the vessel by the balloon results in small localized dissections that can protrude into the lumen and be a nidus for acute thrombus formation. If the dissections are severe, then they can obstruct the lumen or induce a thrombotic occlusion of the artery (acute closure). Stents have largely prevented this complication by holding the dissection flaps up against the vessel wall (Fig. 296e-1).

Stents are currently used in more than 90% of coronary angioplasty procedures. Stents are wire meshes (usually made of stainless steel) that are compressed over a deflated angioplasty balloon. When the balloon is inflated, the stent is enlarged to approximate the “normal” vessel lumen. The balloon is then deflated and removed, leaving the stent behind to provide a permanent scaffold in the artery. Owing to the design of the struts, these devices are flexible, allowing their passage through diseased and tortuous coronary vessels. Stents are rigid enough to prevent elastic recoil of the vessel and have dramatically improved the success and safety of the procedure as a result.

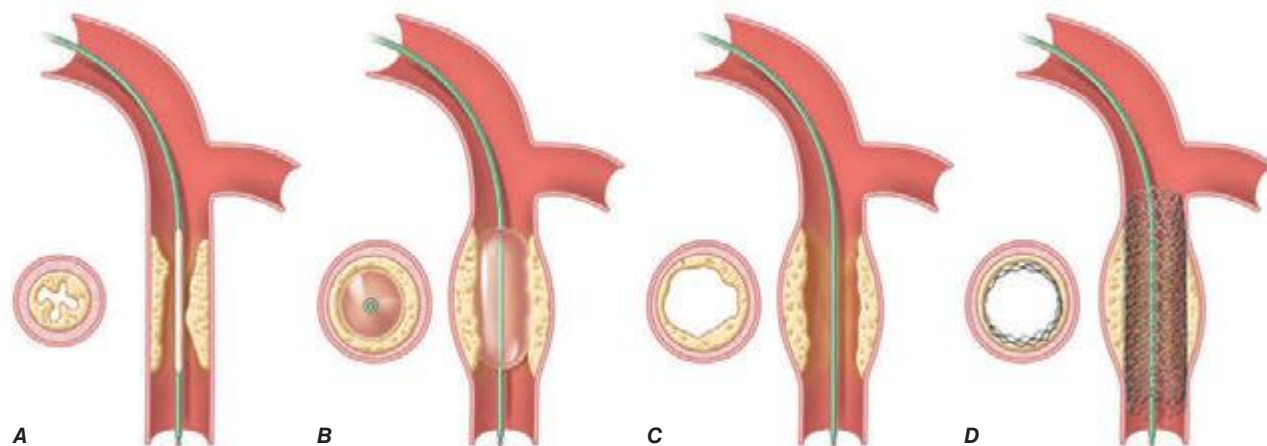


FIGURE 296e-1 Schematic diagram of the primary mechanisms of balloon angioplasty and stenting. **A.** A balloon angioplasty catheter is positioned into the stenosis over a guidewire under fluoroscopic guidance. **B.** The balloon is inflated temporarily occluding the vessel. **C.** The lumen is enlarged primarily by stretching the vessel, often resulting in small dissections in the neointima. **D.** A stent mounted on a deflated balloon is placed into the lesion and pressed against the vessel wall with balloon inflation (not shown). The balloon is deflated and removed, leaving the stent permanently against the wall acting as a scaffold to hold the dissections against the wall and prevent vessel recoil. (Adapted from EJ Topol: *Textbook of Cardiovascular Medicine*, 2nd ed. Philadelphia, Lippincott Williams & Wilkins, 2002.)

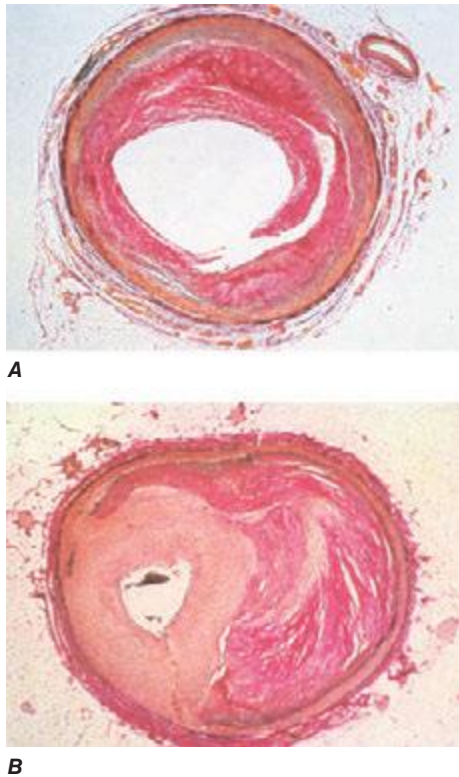


FIGURE 296e-2 Pathology of acute effects of balloon angioplasty with intimal dissection and vessel stretching (A) and an example of neointimal hyperplasia and restenosis showing renarrowing of the vessel (B). (Panel A from M Ueda et al: *Eur Heart J* 12:937, 1991; with permission. Panel B from CE Essed et al: *Br Heart J* 49:393, 1983; with permission.)

Drug-eluting stents further enhanced the efficacy of PCI. An antiproliferative agent is attached to the metal stent by use of a thin polymer coating. The antiproliferative drug elutes from the stent over a 1- to 3-month period after implantation. Drug-eluting stents have been shown to reduce clinical restenosis by 50%, so that in uncomplicated lesions symptomatic restenosis occurs in 5–10% of patients. Not surprisingly, this led to the rapid acceptance of these devices; currently 80–90% of all stents implanted are drug-eluting. The first-generation devices were coated with either sirolimus or paclitaxel. Second-generation drug-eluting stents use newer agents such as everolimus, biolimus, and zotarolimus. These second-generation drug-eluting stents appear to be more effective with fewer complications, such as early or late stent thrombosis, than the first-generation devices and, therefore, have replaced the first-generation stents. Biodegradable polymers that are used to attach the drugs to the stents may be superior to permanent polymers in preventing late stent thrombosis and are under investigation. In addition, the everolimus-eluting biodegradable vascular scaffold (BVS) stent has been shown to be safe and effective with gradual degradation over several years with return of normal vessel function. It is currently approved in Europe. Additional stents are under investigation. Other interventional devices include atherectomy devices and thrombectomy catheters. These devices are designed to remove atherosclerotic plaque or thrombus and are used in conjunction with balloon dilatation and stent placement. Rotational atherectomy is the most commonly used adjunctive device and is modeled after a dentist's drill, with small round burrs of 1.25–2.5 mm at the tip of a flexible wire shaft. They are passed over the guidewire up to the stenosis and drill away atherosclerotic material. Because the atherosclerotic particles are $\leq 25 \mu\text{m}$, they pass through the coronary microcirculation and rarely cause problems. The device is particularly useful in heavily calcified plaques that are resistant to balloon dilatation. Given the current advances in stents, rotational atherectomy is infrequently used. Directional atherectomy catheters are not used in the coronaries any longer but are used in peripheral arterial disease.

In acute ST-elevation myocardial infarction, specialized catheters without a balloon are used to aspirate thrombus in order to prevent embolization down the coronary vessel and to improve blood flow before angioplasty and stent placement. Some data suggest that manual catheter thrombus aspiration may reduce mortality in addition to improving blood flow in primary PCI.

PCI of degenerated saphenous vein graft lesions has been associated with a significant incidence of distal embolization of atherosclerotic material, unlike PCI of native vessel disease. A number of distal protection devices have been shown to significantly reduce embolization and myocardial infarction in this setting. Most devices work by using a collapsible wire filter at the end of a guidewire that is expanded in the distal vessel before PCI. If atherosclerotic debris is dislodged, the basket captures the material, and at the end of the PCI, the basket is pulled into a delivery catheter and the debris safely removed from the patient.

SUCCESS AND COMPLICATIONS

A successful procedure (angiographic success), defined as a reduction of the stenosis to less than a 20% diameter narrowing, occurs in 95–99% of patients. Lower success rates are seen in patients with tortuous, small, or calcified vessels or chronic total occlusions. Chronic total occlusions have the lowest success rates (60–70%), and their recanalization is usually not attempted unless the occlusion is recent (within 3 months) or there are favorable anatomic features. Improvements in equipment and technique have increased the success rates of recanalization of chronic total occlusions.

Serious complications are rare but include a mortality rate of 0.1–0.3% for elective cases, a large myocardial infarction in less than 3%, and stroke in less than 0.1%. Patients who are elderly (>65 years), undergoing an emergent or urgent procedure, have chronic kidney disease, present with an ST-segment elevation myocardial infarction (STEMI), or are in shock have significantly higher risk. Scoring systems can help to estimate the risk of the procedure. Myocardial infarction during PCI can occur for multiple reasons including an acute occluding thrombus, severe coronary dissection, embolization of thrombus or atherosclerotic material, or closure of a side branch vessel at the site of angioplasty. Most myocardial infarctions are small and only detected by a rise in the creatine phosphokinase (CPK) or troponin level after the procedure. Only those with significant enzyme elevations (more than three to five times the upper limit of normal) are associated with a less favorable long-term outcome. Coronary stents have largely prevented coronary dissections due to the scaffolding effect of the stent.

Metallic stents are also prone to stent thrombosis (1–3%), either acute (<24 h) or subacute (1–30 days), which can be ameliorated by greater attention to full initial stent deployment and the use of dual antiplatelet therapy (DAPT) (aspirin, plus a platelet P2Y₁₂ receptor blocker [clopidogrel, prasugrel, or ticagrelor]). Late (30 days–1 year) and very late stent thromboses (>1 year) occur very infrequently with stents but are slightly more common with first-generation drug-eluting stents, necessitating DAPT for up to 1 year or longer. Use of the second-generation stents is associated with lower rates of late and very late stent thromboses, and shorter durations of DAPT may be possible. Premature discontinuation of DAPT, particularly in the first month after implantation, is associated with a significantly increased risk for stent thrombosis (three- to ninefold greater). Stent thrombosis results in death in 10–20% and myocardial infarction in 30–70% of patients. Elective surgery that requires discontinuation of antiplatelet therapy after drug-eluting stent implantation should be postponed until after 6 months and preferably after 1 year, if at all possible.

Restenosis, or renarrowing of the dilated coronary stenosis, is the most common complication of angioplasty and occurs in 20–50% of patients with balloon angioplasty alone, 10–30% of patients with bare metal stents, and 5–15% of patients with drug-eluting stents within the first year. The fact that stent placement provides a larger acute luminal area than balloon angioplasty alone reduces the incidence of subsequent restenosis. Drug-eluting stents further reduce restenosis through a reduction in excessive neointimal growth over the stent. If restenosis does not occur, the long-term outcome is excellent

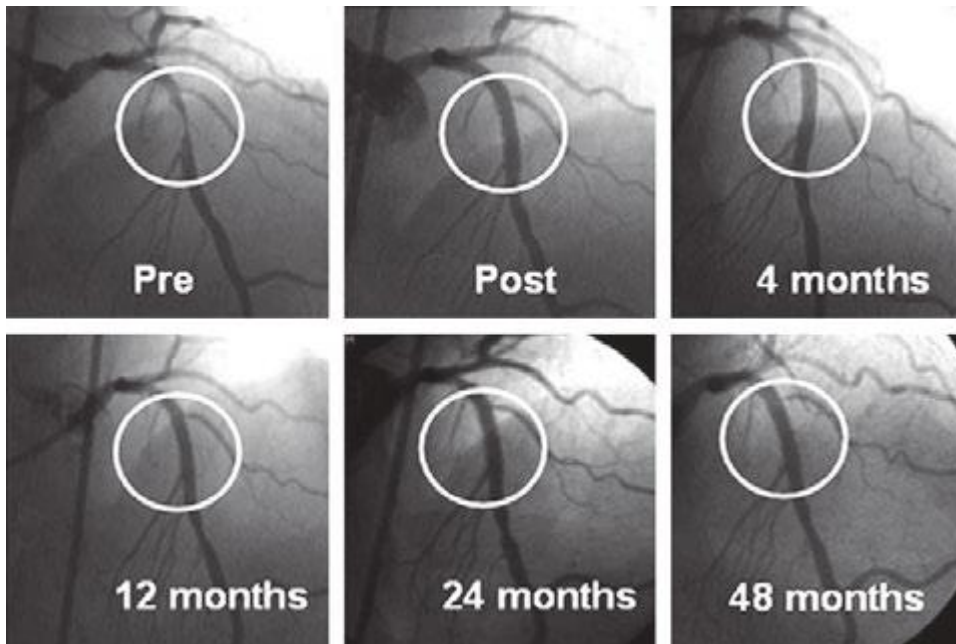


FIGURE 296e-3 Long-term results from one of the first patients to receive a sirolimus-eluting stent from early Sao Paulo experience. (From: GW Stone, in D Baim [ed]: *Cardiac Catheterization, Angiography and Intervention*, 7th ed. Philadelphia, Lippincott Williams & Wilkins, 2006; with permission.)

(Fig. 296e-3). Clinical restenosis is recognized by recurrence of angina or symptoms within 9 months of the procedure. Less frequently, patients with restenosis can present with non-ST-segment elevation myocardial infarction (NSTEMI) (10%) or STEMI (2%) as well. Clinical restenosis requires confirmation of a significant stenosis at the site of the prior PCI. *Target lesion revascularization (TLR)* or *target vessel revascularization (TVR)* is defined as angiographic restenosis with repeat PCI or coronary artery bypass grafting (CABG). By angiography, the incidence of restenosis is significantly higher than clinical restenosis (TLR or TVR) because many patients have mild restenosis that does not result in a recurrence of symptoms. The management of clinical restenosis is usually to repeat the PCI with balloon dilatation and placement of a drug-eluting stent. Once a patient has had restenosis, the risk of a second restenosis is further increased. The risk factors for restenosis are diabetes, myocardial infarction, long lesions, small-diameter vessels, and suboptimal initial PCI result.

INDICATIONS

The American College of Cardiology (ACC)/American Heart Association (AHA) guidelines extensively review the indications for PCI in patients with stable angina, unstable angina, NSTEMI, and STEMI and should be referred to for a comprehensive discussion of the indications. Briefly, the two principal indications for coronary revascularization in patients with *chronic stable angina* (Chap. 293) are (1) to improve anginal symptoms in patients who remain symptomatic despite adequate medical therapy and (2) to reduce mortality rates in patients with severe coronary disease. In patients with stable angina who are well controlled on medical therapy, studies such as the Clinical Outcomes Utilizing Revascularization and Aggressive Drug Evaluation (COURAGE) and Bypass Angioplasty Revascularization Investigation 2 Diabetes (BARI 2D) trials have shown that initial revascularization does not lead to better outcomes and can be safely delayed until symptoms worsen or evidence of severe ischemia on noninvasive testing occurs. When revascularization is indicated, the choice of PCI or CABG depends on a number of clinical and anatomic factors (Fig. 296e-4). The Synergy between Percutaneous Coronary Intervention with Taxus and Cardiac Surgery (SYNTAX) trial compared PCI with the paclitaxel drug-eluting stent to CABG in 1800 patients with three-vessel coronary disease or left main disease. The study found no difference in death or myocardial infarction at 1 year,

but repeat revascularization was significantly higher in the stent-treated group (13.5 vs. 5.9%), while stroke was significantly higher in the surgical group (2.2 vs. 0.6%). The primary endpoint of death, myocardial infarction, stroke, or revascularization was significantly better with CABG, particularly in those with the most extensive coronary artery disease. The 3-year results confirm these findings. The Future Revascularization Evaluation in Patients With Diabetes Mellitus: Optimal Management of Multivessel Disease (FREEDOM) trial randomized 1900 patients with diabetes and multivessel disease and showed a significantly lower primary endpoint of death, myocardial infarction, or stroke with CABG than PCI. These studies support CABG for those with the most severe left main and three-vessel disease or those with diabetes. Lesser degrees of multivessel disease in patients with or without diabetes have an equal outcome with PCI.

The choice of PCI versus CABG is also related to the anticipated procedural success and complications of PCI and the risks of CABG. For PCI, the

characteristics of the coronary anatomy are critically important. The location of the lesion in the vessel (proximal or distal), the degree of tortuosity, and the size of the vessel are considered. In addition, the

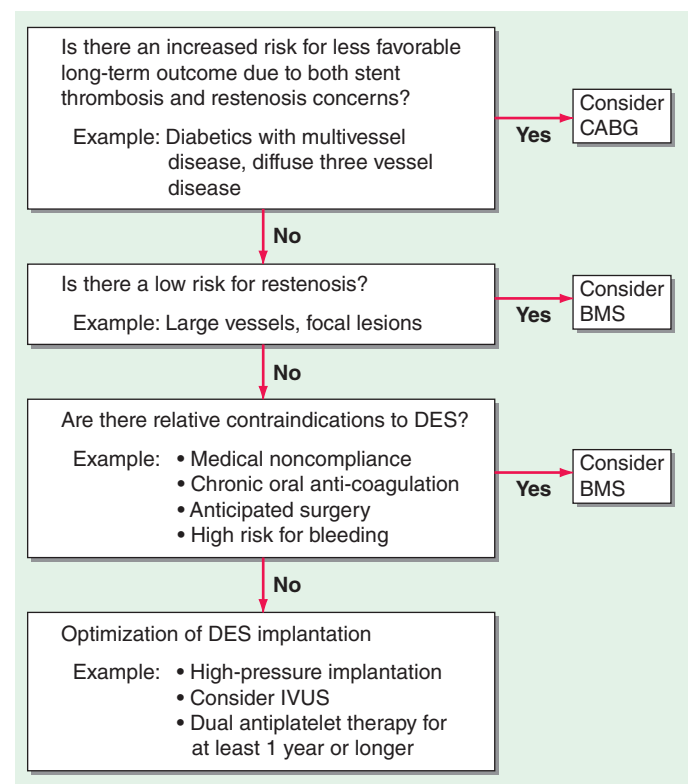


FIGURE 296e-4 In patients requiring revascularization, several factors need to be considered in choosing between bare metal stents, drug-eluting stents, or coronary artery bypass surgery.

ACS, acute coronary syndrome; BMS, bare metal stent; CABG, coronary artery bypass grafting; DES, drug-eluting stent; IVUS, intravascular ultrasound; STEMI, ST-segment elevation myocardial infarction. (From AA Bavy, DL Bhatt: *Circulation* 116:696, 2007; with permission.)

lesion characteristics, including the degree of the stenosis, the presence of calcium, lesion length, and presence of thrombus, are assessed. The most common reason to decide not to do PCI is that the lesion felt to be responsible for the patient's symptoms is not treatable. This is most commonly due to the presence of a chronic total occlusion (>3 months in duration). In this setting, the historical success rate has been low (30–70%) and complications are more common. A lesion classification to characterize the likelihood of success or failure of PCI has been developed by the ACC/AHA. Lesions with the highest success are called type A lesions (such as proximal noncalcified subtotal lesions), and those with the lowest success or highest complication rate are type C lesions (such as chronic total occlusions). Intermediate lesions are classified as type B1 or B2 depending on the number of unfavorable characteristics. Approximately 25–30% of patients will not be candidates for PCI due to unfavorable anatomy, whereas only 5% of CABG patients will not be candidates for surgery due to coronary anatomy. The primary reason for being considered inoperable with CABG is the presence of severe comorbidities such as advanced age, frailty, severe chronic obstructive pulmonary disease (COPD), or poor left ventricular function.

Another consideration in choosing a revascularization strategy is the degree of revascularization. In patients with multivessel disease, bypass grafts can usually be placed to all vessels with significant stenosis, whereas PCI may be able to treat only some of the lesions due to the presence of unfavorable anatomy. Assessment of the significance of intermediate lesions using fractional flow reserve (FFR) ([Chap. 272](#)) can assist in determining which lesions should be revascularized. The Fractional Flow Reserve versus Angiography for Multivessel Evaluation (FAME) trial showed a 30% reduction in adverse events when revascularization by PCI was restricted to those lesions that were hemodynamically significant (FFR ≤ 0.80) rather than when guided by angiography alone. Thus, complete revascularization of all functionally significant lesions should be favored and considered when choosing the optimal revascularization strategy. Given the multiple factors that need to be considered in choosing the best revascularization for an individual patient with multivessel disease, it is optimal to have a discussion among the cardiac surgeon, interventional cardiologist, and the physicians caring for the patient (so-called Heart Team) to properly weigh the choices.

Patients with acute coronary syndrome are at excess risk of short- and long-term mortality. Randomized clinical trials have shown that PCI is superior to intensive medical therapy in reducing mortality and myocardial infarction, with the benefit largely confined to those patients who are high risk. High-risk patients are defined as those with any one of the following: refractory ischemia, recurrent angina, positive cardiac-specific enzymes, new ST-segment depression, low ejection fraction, severe arrhythmias, or a recent PCI or CABG. PCI is preferred over surgical therapy in most high-risk patients with acute coronary syndromes unless they have severe multivessel disease or the culprit lesion responsible for the unstable presentation cannot be adequately treated. In STEMI, thrombolysis or PCI (primary PCI) are effective methods to restore coronary blood flow and salvage myocardium within the first 12 h after onset of chest pain. Because PCI is more effective in restoring flow than thrombolysis, it is preferred if readily available. PCI is also performed following thrombolysis to facilitate adequate reperfusion or as a rescue procedure in those who do not achieve reperfusion from thrombolysis or in those who develop cardiogenic shock.

OTHER INTERVENTIONAL TECHNIQUES

STRUCTURAL HEART DISEASE

Interventional treatment for structural heart disease (adult congenital heart disease and valvular heart disease) is a significant and growing component of the field of interventional cardiology.

The most common adult congenital lesion to be treated with percutaneous techniques is closure of atrial septal defects ([Chap. 282](#)). The procedure is done as in a diagnostic right heart catheterization with the passage of a catheter up the femoral vein into the right atrium. With echo and fluoroscopic guidance, the size and location of the defect can

be accurately defined, and closure is accomplished using one of several approved devices. All devices use a left atrial and right atrial wire mesh or covered disk that are pulled together to capture the atrial septum around the defect and seal it off. The Amplatzer Septal Occluder device (AGA Medical, Minneapolis, Minnesota) is the most commonly used in the United States. The success rate in selected patients is 85–95%, and the device complications are rare and include device embolization, infection, or erosion. Closure of patent foramen ovale (PFO) is done in a similar way. PFO closure may be considered in patients who have had recurrent paradoxical stroke or transient ischemic attack (TIA) despite adequate medical therapy including anticoagulation or antiplatelet therapy. The benefit, however, has not been proven. The CLOSURE I trial randomized 909 patients with cryptogenic stroke or TIA who had a PFO. Closure did not reduce the primary endpoint of death within 30 days or death following a neurologic cause during 2 years of follow-up or stroke/TIA within 2 years. Other trials have confirmed these findings. The use in the treatment of migraine is under clinical investigation and is not an approved indication.

Similar devices can also be used to close patent ductus arteriosus and ventricular septal defects. Other congenital diseases that can be treated percutaneously include coarctation of the aorta, pulmonic stenosis, peripheral pulmonary stenosis, and other abnormal communications between the cardiac chambers or vessels.

The treatment of valvular heart disease is the most rapidly growing area in interventional cardiology. Until recently, the only available techniques were balloon valvuloplasty for the treatment of aortic, mitral, or pulmonic stenosis ([Chap. 283](#)). Mitral valvuloplasty is the preferred treatment for symptomatic patients with rheumatic mitral stenosis who have favorable anatomy. The outcome in these patients is equal to that of surgical commissurotomy. The success is highly related to the echocardiographic appearance of the valve. The most favorable setting is commissural fusion without calcification or subchordal fusion and the absence of significant mitral regurgitation. Access is obtained from the femoral vein using a transseptal technique in which a long metal catheter with a needle tip is advanced from the femoral vein through the right atrium and atrial septum at the level of the foramen ovale into the left atrium. A guidewire is advanced into the left ventricle, and a balloon-dilatation catheter is negotiated across the mitral valve and inflated to a predetermined size to enlarge the valve. The most commonly used dilatation catheter is the Inoue balloon. The technique splits the commissural fusion and commonly results in a doubling of the mitral valve area. The success of the procedure in favorable anatomy is 95% and severe complications are rare (1–2%). The most common complications are tamponade due to puncture into the pericardium or the creation of severe mitral regurgitation.

For regurgitant valvular lesions, only severe mitral regurgitation can be effectively treated percutaneously using the MitraClip (Abbott, Abbott Park, Ill) device. The procedure involves the passage of a catheter into the left atrium using the transseptal technique. A special catheter with a metallic clip on the end is passed through the mitral valve and retracted to catch and clip together the mid portion of the anterior and posterior mitral valve leaflet. The clip creates a double opening in the mitral valve and thereby reduces mitral regurgitation similar to the surgical Alfieri repair. In the Endovascular Valve Edge-to-Edge Repair Study (EVEREST II) trial, the device was less effective than surgical repair or replacement but was shown to be safe. It is currently used for patients who are not good candidates for surgical repair, particularly when the regurgitation is due to functional causes.

Severe aortic stenosis can be treated with balloon valvuloplasty as well. In this setting, the valvuloplasty balloon catheter is placed retrograde across the aortic valve from the femoral artery and briefly inflated to stretch open the valve. The success is much less favorable, with only 50% achieving an aortic valve area of >1 cm² and a restenosis rate of 25–50% after 6–12 months. This poor success rate has limited its use to patients who are not surgical candidates or as a bridge to surgery or transcatheter aortic valve replacement (TAVR). In this setting, the intermediate-term mortality rate of the procedure is high (10%). Repeat aortic valvuloplasty as a treatment for aortic valve restenosis has been reported.

Percutaneous aortic valve replacement (TAVR) has been shown to be an effective treatment for high-risk and inoperable patients with aortic stenosis. Currently, two valve models, the Edwards SAPIEN valve (Edwards Lifescience, Irvine, California) and the CoreValve ReValving system (Medtronic, Minneapolis, Minnesota) are available. In more than 10,000 cases worldwide, follow-up shows no evidence of restenosis or severe prosthetic valve dysfunction in the midterm. The CoreValve is self-expanding, while the Edwards valve is balloon expanded. The cannulas are large (14–22 French), and retrograde access via the femoral artery is most commonly chosen, if possible. In patients with peripheral artery disease, access via the subclavian artery or transapically through a surgical incision can be used. Following balloon valvuloplasty, the valve is positioned across the valve and deployed with postdeployment balloon inflation to ensure full contact with the aortic annulus. The success rate is 80–90%, and the 30-day mortality rate is 10–15%, not unexpectedly as only high-risk patients are undergoing the procedure currently. The Placement of Aortic Transcatheter Valve (PARTNER) randomized trial of the Edwards valve showed a 55% reduction in 1-year mortality and major adverse events in the extreme-risk group randomized to TAVR compared to medical therapy. In a separate randomized trial, high-risk patients had a similar outcome to surgical valve replacement at 1 year. As a result, this valve is approved for both high-risk and extreme-risk patients with severe aortic stenosis.

Pulmonic stenosis can also be effectively treated with balloon valvuloplasty and percutaneously replaced with the Melody stent (Medtronic). Tricuspid valve interventions remain experimental.

PERIPHERAL ARTERY INTERVENTIONS

The use of percutaneous interventions to treat symptomatic patients with arterial obstruction in the carotid, renal, aortic, and peripheral vessels is also part of the field of interventional cardiology. Randomized clinical trial data already support the use of carotid stenting in patients at high risk of complications from carotid endarterectomy (Fig. 296e-5). Recent trials suggest similar outcomes with carotid stenting and carotid endarterectomy in patients at average risk, although depending on the patient's risk for periprocedural stroke or myocardial infarction, one procedure may be preferred over the other. The success rate of peripheral artery interventional procedures has been improving, including for long segments of occlusive disease historically treated by peripheral bypass surgery (Fig. 296e-6). Peripheral intervention is increasingly part of the training of an interventional cardiologist, and most programs now require an additional year of training after the interventional cardiology training year. The techniques and outcomes are described in detail in the chapter on peripheral vascular disease (Chap. 302).

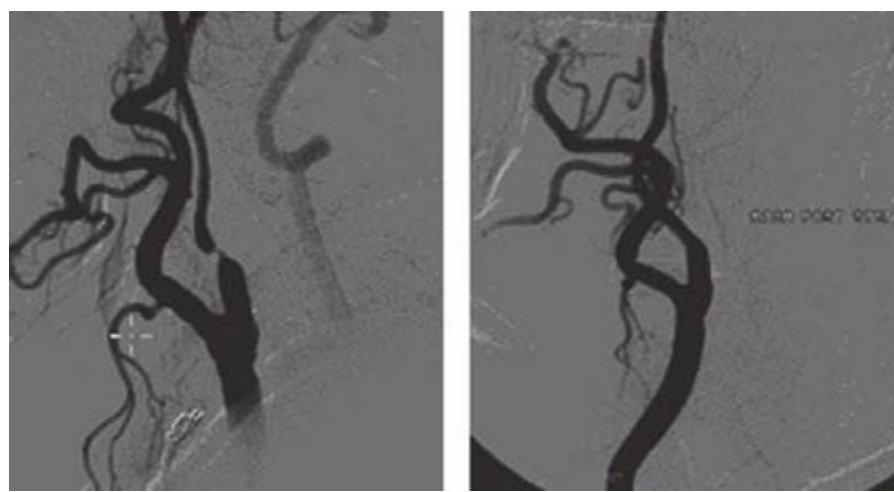


FIGURE 296e-5 An example of a high-risk patient who requires carotid revascularization, but who is not a candidate for carotid endarterectomy. Carotid artery stenting resulted in an excellent angiographic result. (From M Belkin, DL Bhatt: *Circulation* 119:2302, 2009; with permission.)

CIRCULATORY SUPPORT TECHNIQUES

The use of circulatory support techniques is occasionally needed in order to safely perform PCI on hemodynamically unstable patients. It also can be useful in helping to stabilize patients before surgical interventions. The most commonly used device is the percutaneous intraaortic balloon pump developed in the early 1960s. A 7- to 10-French, 25- to 50-mL balloon catheter is placed retrograde from the femoral artery into the descending aorta between the aortic arch and the abdominal aortic bifurcation. It is connected to a helium gas inflation system that synchronizes the inflation to coincide with early diastole with deflation by mid-diastole. As a result, it increases early diastolic pressure, lowers systolic pressure, and lowers late diastolic pressure through displacement of blood from the descending aorta (counterpulsation). This results in an increase in coronary blood flow and a decrease in afterload. It is contraindicated in patients with aortic regurgitation, aortic dissection, or severe peripheral artery disease. The major complications are vascular and thrombotic. Intravenous heparin is given in order to reduce thrombotic complications.

Another potentially useful tool is the Impella device (Abiomed, Danvers, Massachusetts). The catheter is placed percutaneously from the femoral artery into the left ventricle. The catheter has a small microaxial pump at its tip that can pump up to 2.5–5 L/min from the left ventricle to the aorta. Other support devices include TandemHeart (CardiacAssist, Pittsburgh, Pennsylvania), which involves placement of a large 21-French catheter from the femoral vein through the right atrium into the left atrium using the transeptal technique and a catheter in the femoral artery. A centrifugal pump can deliver 5 L of blood per minute. It may be useful in patients in shock or with STEMI or very-high-risk PCI. Patients can also be placed on peripheral extracorporeal membrane oxygenation using large cannulas placed in the femoral artery and vein.

INTERVENTIONS FOR PULMONARY EMBOLISM

The treatment of deep vein thrombosis is intravenous anticoagulation, with placement of an inferior vena cava filter if recurrent pulmonary emboli occur. Postphlebotic syndrome is a serious condition due to chronic venous obstruction that can lead to chronic leg edema and venous ulcers. Preliminary studies suggest that mechanical treatments may have a role in treatment, and a large trial is ongoing.

Pulmonary emboli (PE) should be treated with fibrinolytic agents if massive and in some cases if submassive. Surgical pulmonary embolectomy is an option for the treatment of massive PE with hemodynamic instability in patients who have contraindications for systemic fibrinolysis or those in whom it has failed. Catheter-based therapies for submassive and massive PEs are still evolving, but studies have shown promise. The techniques employed include the use of aspiration of the clot with a large catheter (10 French), intraclot infusion of a thrombolytic agent followed by aspiration, ultrasound-assisted catheter-directed thrombolysis, and use of rheolytic thrombectomy. Success for these techniques has been reported to be 80–90%, with major complications occurring in 2–4% of patients.

INTERVENTIONS FOR REFRACTORY HYPERTENSION

The recent recognition of the importance of the renal sympathetic nerves in modulating blood pressure has led to a technique to selectively denervate renal sympathetic nerves in patients with refractory hypertension. The procedure involves applying low-power radiofrequency treatment via a catheter along the length of both renal arteries. In the randomized Symplicity HTN-2 trial, renal denervation significantly reduced blood pressure compared with medical therapy. The Symplicity device (Medtronic) is approved in Europe, though the randomized and blinded U.S. Symplicity HTN-3 trial showed no effect.

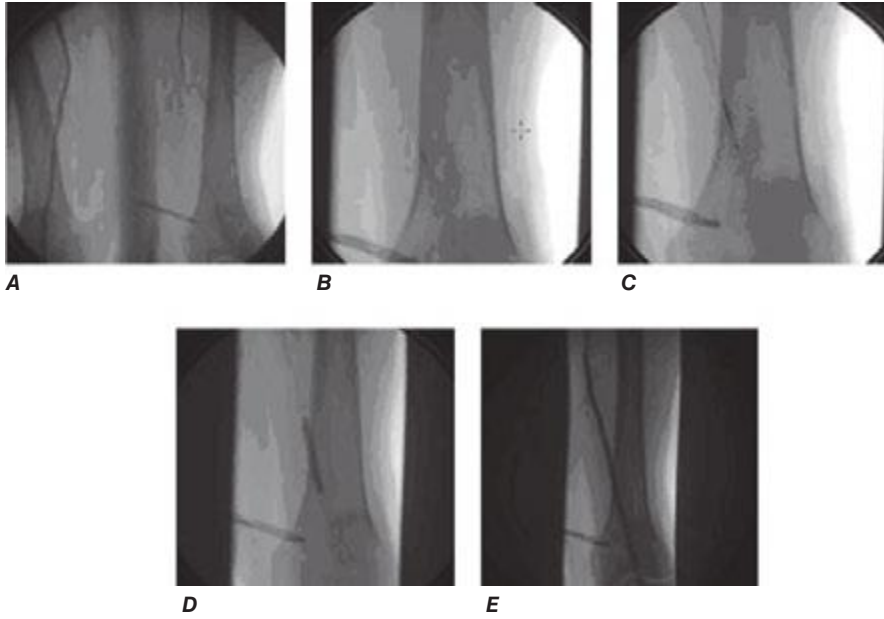


FIGURE 296e-6 Peripheral interventional procedures have become highly effective at treating anatomic lesions previously amenable only to bypass surgery. **A.** Complete occlusion of the left superficial femoral artery. **B.** Wire and catheter advanced into subintimal space. **C.** Intravascular ultrasound positioned in the subintimal space to guide retrograde wire placement through the occluded vessel. **D.** Balloon dilation of the occlusion. **E.** Stent placement with excellent angiographic result. (From A Al Mahameed, DL Bhatt: *Cleve Clin J Med* 73:S45, 2006; with permission.)

CONCLUSION

Interventional cardiology continues to expand its borders. Treatment for coronary artery disease, including complex anatomic subsets, continues to advance, encroaching on what has traditionally been treated by CABG. Technological advances such as drug-eluting stents, now already in their second generation, and manual thrombus aspiration devices are improving the results of PCI. In particular, PCI has been shown to prevent future ischemic events in acute coronary syndromes. For patients with stable coronary disease, PCI has an important role in symptom alleviation. Treatment of peripheral and cerebrovascular disease has also benefited from the application of percutaneous techniques. Structural heart disease is increasingly being treated with percutaneous options, with a high likelihood that interventional approaches will compete with open-heart surgery in a significant proportion of cases in years to come.

297e Atlas of Percutaneous Revascularization

Jane A. Leopold, Deepak L. Bhatt, David P. Faxon

Percutaneous coronary intervention (PCI) is the most widely employed coronary revascularization procedure worldwide (Chap. 296e). It is now applied to patients with stable angina; patients with acute coronary syndromes, including unstable angina and non-ST-segment elevation myocardial infarction (NSTEMI); and as a primary treatment strategy in patients with ST-segment elevation myocardial infarction (STEMI). PCI is also applicable to patients with either single-vessel or multivessel disease.

In this chapter, the use of PCI will be illustrated in a variety of commonly encountered clinical and anatomic situations, such as chronic total occlusion of a coronary artery, bifurcation disease, acute STEMI, saphenous vein graft disease, left main coronary artery disease, multivessel disease, and stent thrombosis. In addition, the use of interventional techniques to treat structural heart disease will be shown, including closure of an atrial septal defect (ASD) and transcatheter aortic valve replacement (TAVR).

CASE 1: CHRONIC TOTAL OCCLUSION

(Videos 297e-1 to 297e-7)

- An 81-year-old male with angina, New York Heart Association (NYHA) class IV congestive heart failure and inferoapicoposterior ischemia on an exercise technetium-99m scan.
- Diagnostic cardiac catheterization revealed a left dominant system with a totally occluded left circumflex (LCx) artery. The distal LCx filled via collaterals from the left anterior descending (LAD) artery, indicating chronicity of the total occlusion.

VIDEO 297e-1 Baseline left coronary angiogram shows an occluded LCx with left-to-left collaterals originating from LAD septal vessels.

VIDEO 297e-2 Attempts to cross the total occlusion in the LCx using a hydrophilic wire and an antegrade approach were not successful with the wire tracking to the right of the trajectory.

VIDEO 297e-3 The LAD septal collateral is accessed with a guidewire that is directed toward the distal LCx to cross the total occlusion retrograde.

VIDEO 297e-4 The total occlusion is crossed retrograde. The wire is snared in the guide, exteriorized, and used to provide antegrade access to the LCx.

VIDEO 297e-5 Antegrade flow in the LCx is restored after balloon inflation.

VIDEO 297e-6 Following stenting of the total occlusion, blood flow in the distal vessel is improved and a second significant stenosis is seen.

VIDEO 297e-7 Final result after LCx stenting.

SUMMARY

- Approximately 15–30% of all patients referred for cardiac catheterization will have a chronic total occlusion (CTO) of a coronary artery.
- CTO often leads to a surgical referral for complete revascularization.
- Incomplete revascularization due to an untreated CTO is associated with an increased mortality rate (hazard ratio = 1.36; 95% confidence interval [CI], 1.12–1.66, $p < .05$).
- Successful PCI of a CTO leads to a 3.8–8.4% absolute reduction in mortality, symptom relief, and improved left ventricular function.
- Newer techniques, such as the retrograde approach to crossing total occlusions, are useful when the antegrade approach fails or is not feasible and there are well-developed collateral vessels.

(Case contributed with permission by Dr. Frederick G.P. Welt.)

CASE 2: BIFURCATION STENTING

(Fig. 297e-1; Videos 297e-8 to 297e-16)

- A 52-year-old male with an acute coronary syndrome and a troponin I = 0.18 (upper limit of normal, <0.04).
- Diagnostic cardiac catheterization showed single-vessel coronary artery disease with a significant stenosis in the mid-LAD and a bifurcation lesion involving a large diagonal branch.

VIDEO 297e-8 Baseline angiogram of the left coronary circulation shows the significant stenosis in the mid-LAD and the bifurcation lesion involving a large diagonal branch.

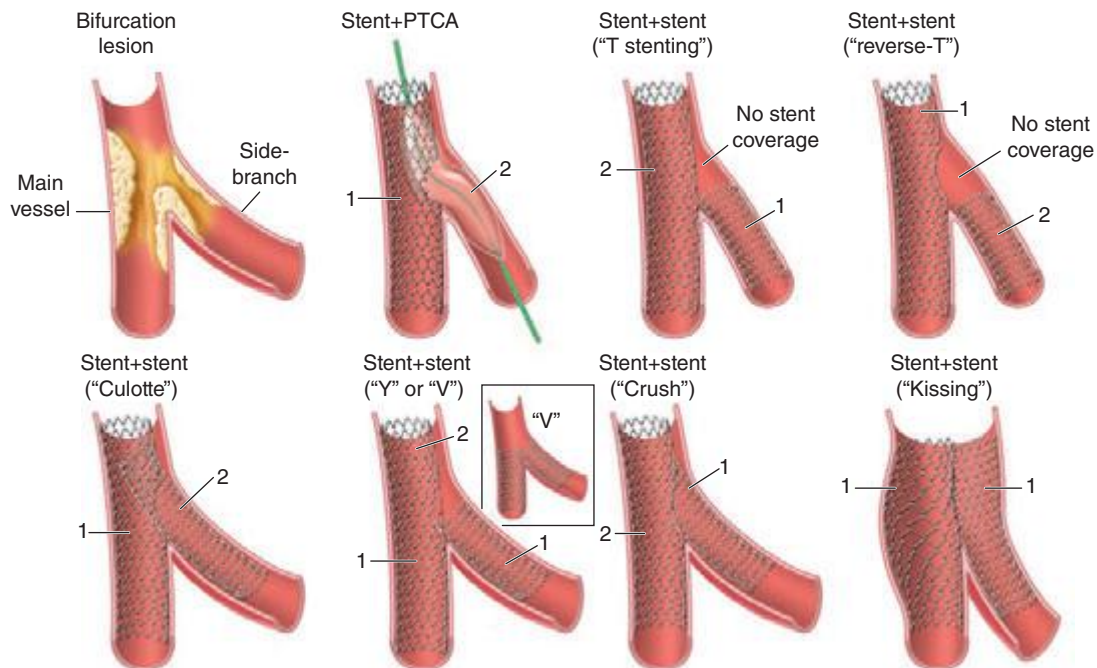


FIGURE 297e-1 Schematic representation of one-stent and two-stent techniques to treat bifurcation lesions. PTCA, percutaneous transluminal coronary angioplasty

297e-2 **VIDEO 297e-9** Both vessels are accessed with guidewires and pre-treated with balloon angioplasty.

VIDEO 297e-10 Result after balloon angioplasty.

VIDEO 297e-11 Stent being positioned in the LAD.

VIDEO 297e-12 LAD poststent result.

VIDEO 297e-13 Stent deployed in the diagonal branch through the stent struts in the LAD using the “culotte” technique.

VIDEO 297e-14 Diagonal branch poststent result.

VIDEO 297e-15 Simultaneous inflation of two 2.5-mm “kissing” balloons.

VIDEO 297e-16 Final postbifurcation stenting result.

SUMMARY

- Approximately 15–20% of PCIs will involve the treatment of bifurcation lesions.
- Bifurcation lesions require consideration of PCI strategies that protect side-branch patency.
- There are both one-stent and two-stent techniques to treat bifurcation lesions; the selection of technique depends on anatomic considerations, including plaque burden, angle of side-branch take-off, plaque shift during angioplasty, and side-branch distribution.
- Rates of target lesion revascularization and stent thrombosis are similar between one-stent and two-stent procedures.

CASE 3: INFERIOR MYOCARDIAL INFARCTION—THROMBUS AND MANUAL THROMBECTOMY

(Figs. 297e-2 to 297e-4; Videos 297e-17 to 297e-22)

- A 59-year-old male presented to the emergency room with 2 h of severe midsternal chest pressure.
- His systolic blood pressure was 100 mmHg, and he was tachycardic in sinus rhythm with a heart rate of 90–100 beats/min.
- His initial electrocardiogram (ECG) showed inferior ST-segment elevations with lateral ST-segment depressions.
- He was referred emergently to the cardiac catheterization laboratory for primary PCI.

VIDEO 297e-17 The right coronary artery (RCA) is totally occluded with filling defects in the vessel after contrast injection, indicating thrombus is present in the vessel.

VIDEO 297e-18 An angioplasty wire is threaded through the thrombotic lesion, but this does not restore blood flow to the distal vessel.

VIDEO 297e-19 Result after manual thrombectomy and thrombus extraction. The “culprit” ruptured plaque and residual thrombus are now apparent in the vessel.

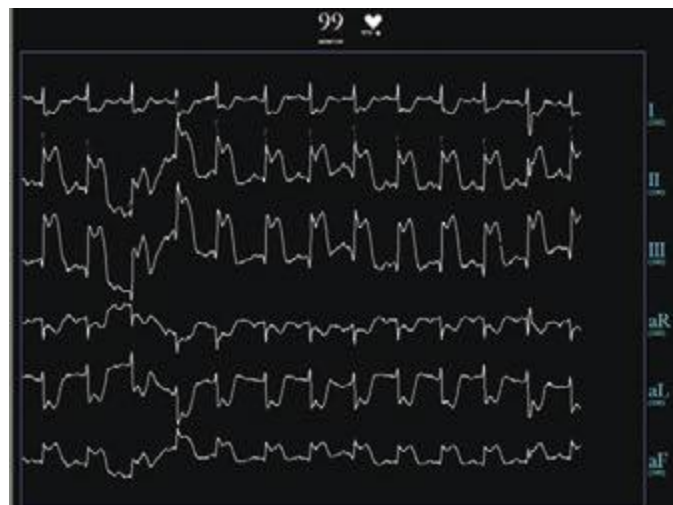


FIGURE 297e-2 Preprocedure ECG showing inferior ST-segment elevations and lateral ST-segment depressions.

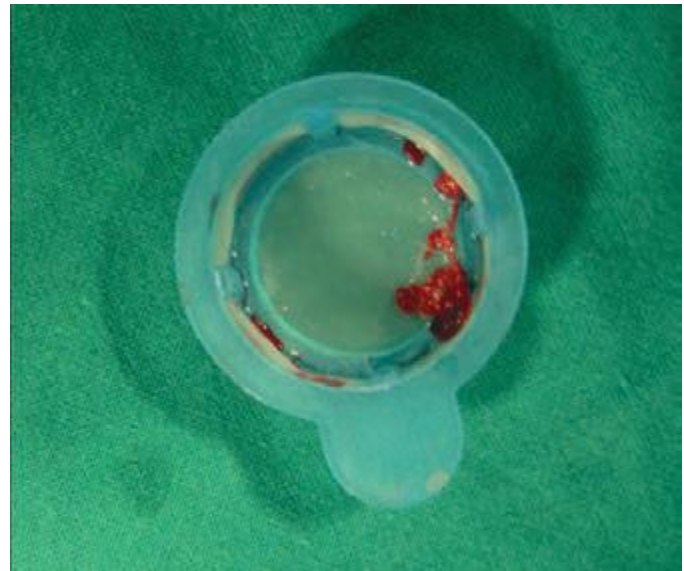


FIGURE 297e-3 Example of an organized red thrombus retrieved by manual thrombectomy.

VIDEO 297e-20 After balloon angioplasty and stenting, thrombus is still present.

VIDEO 297e-21 After repeat manual thrombectomy and expansion of the stent, the thrombus is no longer present.

VIDEO 297e-22 Final result.

SUMMARY

- An acute STEMI occurs following plaque rupture that promotes thrombotic occlusion of a coronary artery.
- Despite successful revascularization of the epicardial coronary artery, microemboli liberated during balloon angioplasty and stenting may lead to persistent microvascular dysfunction. When present, microvascular dysfunction is associated with a larger infarct size, heart failure, malignant ventricular arrhythmias, and death.
- Manual thrombectomy is used to aspirate or remove thrombus in the vessel and limit distal embolization during angioplasty and stenting.
- Manual thrombectomy in primary PCI is associated with improved myocardial perfusion and a reduction in mortality.
- Adjunctive antiplatelet and antithrombin agents are important to aid in the resolution of intracoronary thrombus.

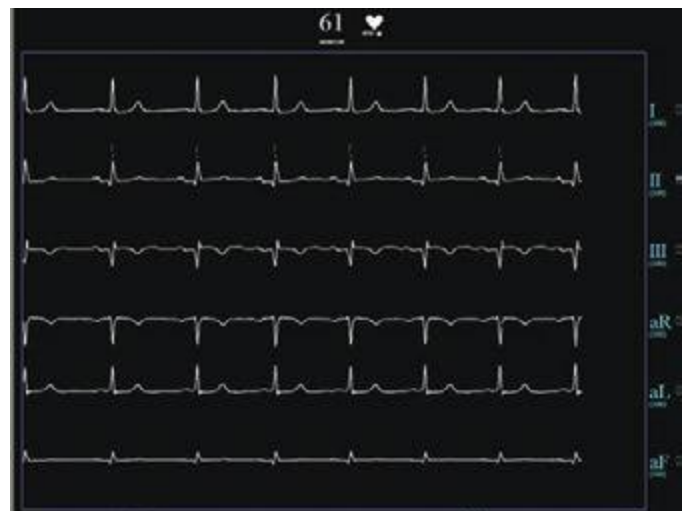


FIGURE 297e-4 Postprocedure ECG showing resolution of ST-segment elevations.

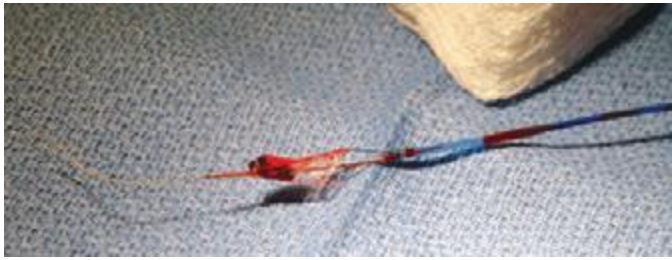


FIGURE 297e-5 Distal protection device showing captured atherosclerotic debris liberated by initial balloon dilation.

CASE 4: SAPHENOUS VEIN GRAFT INTERVENTION WITH DISTAL PROTECTION

(Fig. 297e-5; Videos 297e-23 to 297e-26)

- A 62-year-old male with a history of chronic stable angina.
- A four-vessel coronary artery bypass grafting (CABG) surgery was performed 17 years earlier with a left internal mammary artery graft to the LAD, a right internal mammary artery graft to the right coronary artery (RCA), a saphenous vein graft to the first obtuse marginal branch, and a saphenous vein graft to the first diagonal branch.
- The patient had a recent increase in angina with exertion and was found to have lateral ischemia on an exercise technetium-99m scan.
- Diagnostic cardiac catheterization revealed a significant stenosis in the body of the saphenous vein graft to the first obtuse marginal branch.

VIDEO 297e-23 Saphenous vein graft to a first obtuse marginal branch with an 80% eccentric stenosis in the midgraft.

VIDEO 297e-24 A distal protection device is deployed past the lesion.

VIDEO 297e-25 Angioplasty balloon inflation with the distal protection device in place.

VIDEO 297e-26 Final result after stent placement.

SUMMARY

- Saphenous vein grafts have a failure rate of up to 20% after 1 year and as high as 50% by 5 years.
- Graft failure (after >1 month) results from intimal hyperplasia and atherosclerosis.

- Saphenous vein graft PCI is associated with distal embolization of atherosclerotic debris and microthrombi leading to microvascular occlusion, reduced antegrade blood flow (the “no-reflow” phenomenon), and myocardial infarction.
- Embolic distal protection devices decrease the risk of distal embolization, as well as the incidence of no-reflow and myocardial infarction associated with saphenous vein graft interventions.

CASE 5: UNPROTECTED LEFT MAIN PCI IN A HIGH-RISK PATIENT

(Figs. 297e-6 and 297e-7; Videos 297e-27 to 297e-34)

- An 89-year-old woman presented with a NSTEMI associated with 5-mm ST-segment depression in the apical leads occurring 2 weeks after hospitalization for a NSTEMI that was treated conservatively.
- Chronic obstructive lung disease, elderly age, and the patient’s refusal to consider cardiac surgery restricted the choice of therapeutic options to medical and/or percutaneous interventions.
- Diagnostic catheterization revealed a left dominant circulation with a heavily calcified 80% distal left main coronary artery stenosis extending into the LAD and into the proximal LCx coronary arteries. A 70% proximal LAD lesion was also present.
- After consultation with the patient, family, and a cardiac surgeon, PCI was performed with intraaortic balloon pump support and a temporary pacemaker in the right ventricle.

VIDEO 297e-27 Baseline left coronary artery injection in right anterior oblique (RAO) cranial projection shows a high-grade calcified stenosis in the left main coronary artery and a significant stenosis in the proximal LAD.

VIDEO 297e-28 In the left anterior oblique (LAO) caudal view, the left main coronary artery lesion can be seen to extend into the ostia of both the LCx and the LAD.

VIDEO 297e-29 Guidewires were placed into both the LCx and LAD. After the left main coronary artery and LCx are dilated with balloon angioplasty, the proximal LAD is dilated, and a long drug-eluting stent is placed to cover a lesion dissection that occurred with wiring of the vessel.

VIDEO 297e-30 The bifurcation lesion in the left main coronary artery extending into the LCx and LAD ostia is treated using a “culotte” technique. First, a drug-eluting stent is placed in the left main coronary artery and into the proximal LCx.

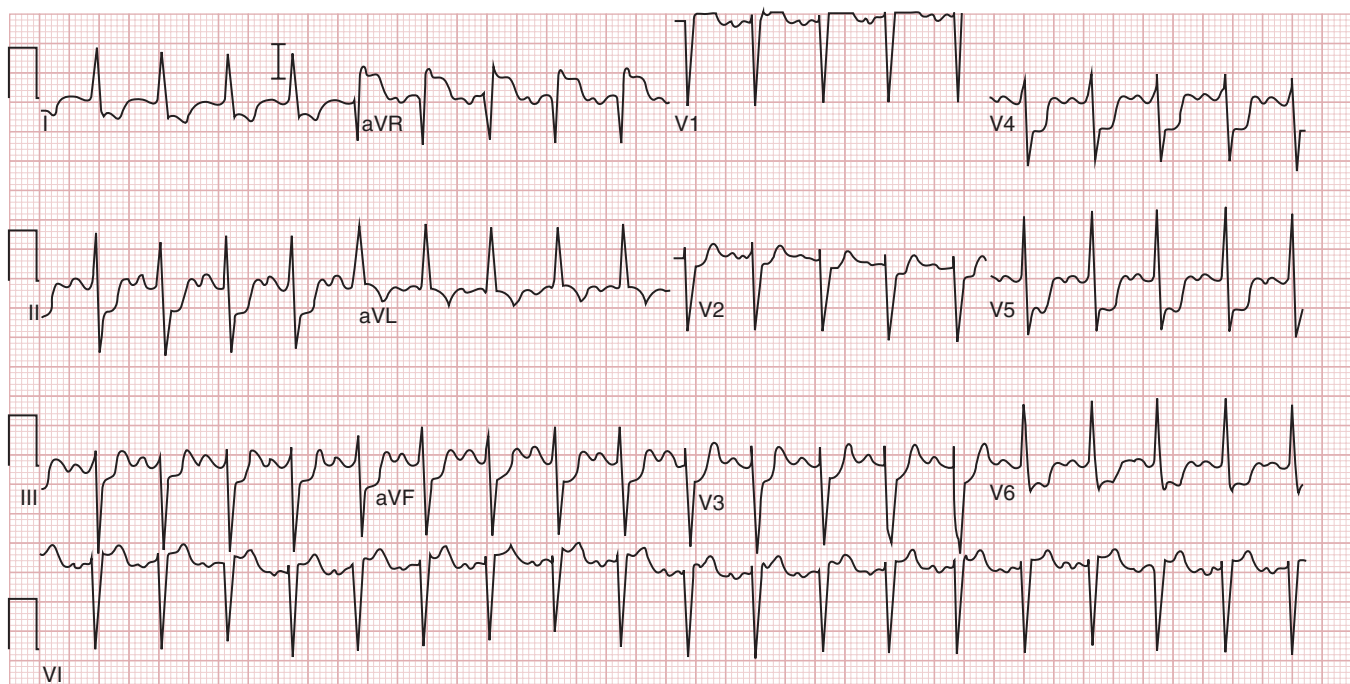


FIGURE 297e-6 During chest pain, the ECG showed diffuse ST-segment depression of up to 5 mm in the inferior and lateral leads.

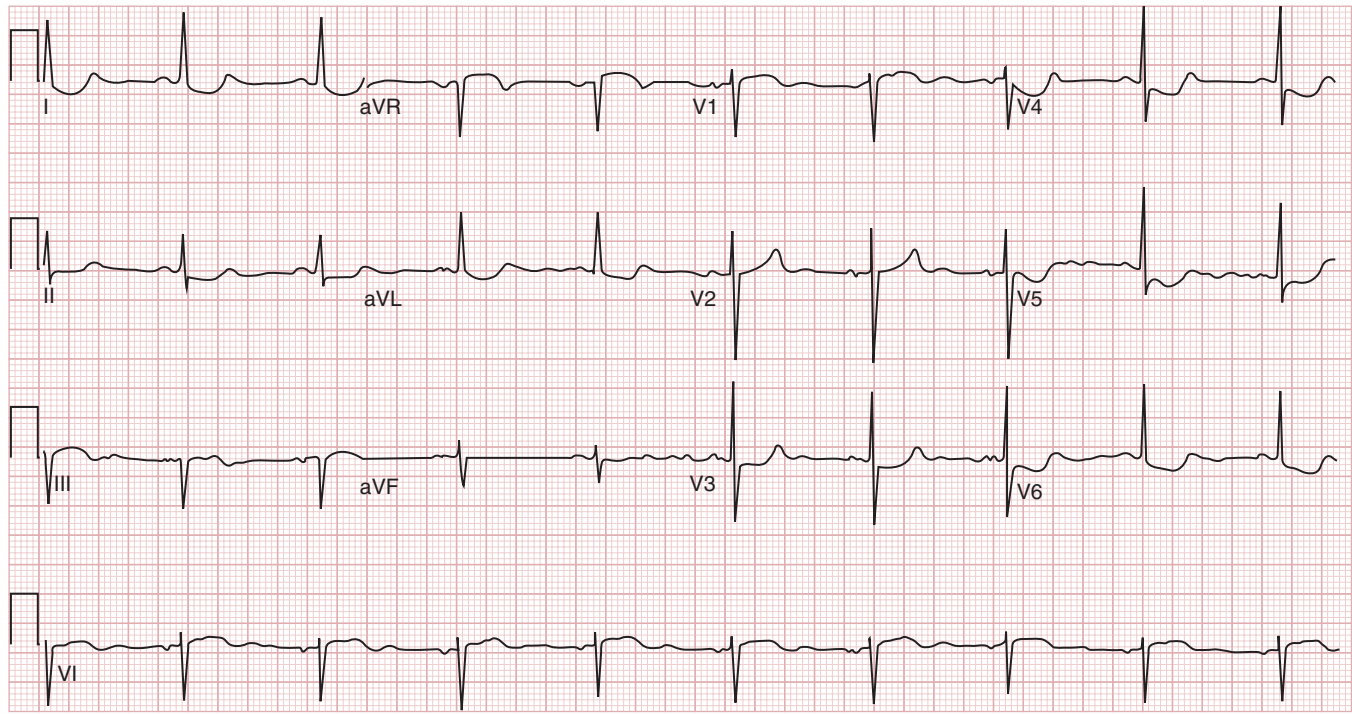


FIGURE 297e-7 Following resolution of the chest pain, the ST-segment depression is less marked.

VIDEO 297e-31 Next, the LAD wire is removed and passed through the stent into the distal LAD. A second drug-eluting stent is deployed through the struts of the left main coronary artery/LCx stent.

VIDEO 297e-32 Following rewiring of the LCx, both stents are re-dilated simultaneously ("kissing" balloons).

VIDEO 297e-33 The final result in the LAO caudal view.

VIDEO 297e-34 The final result in the RAO cranial view showing patent left main, LCx, and LAD coronary arteries.

SUMMARY

- Left main coronary artery disease occurs in 5–10% of patients with symptomatic coronary artery disease.
- In patients with left main coronary artery disease, revascularization with CABG has been shown to decrease mortality significantly over 5–10 years of follow-up.
- PCI with drug-eluting stents in selected cases has been shown to have equal in-hospital and 1-year death and myocardial infarction rates compared with CABG in the Synergy between PCI with Taxus and Cardiac Surgery (SYNTAX) trial. Long-term outcome differences between the two treatment strategies are not known.
- Indications for PCI of left main coronary artery lesions are high-risk surgical patients and patients with protected left main coronary artery disease (i.e., prior CABG with patent bypass grafts). Patients who are good candidates for bypass surgery may also undergo a stenting procedure, but discussion with the patient, the interventional cardiologist, and the cardiac surgeon should be undertaken to determine the best treatment option in an individual case.
- Outcomes are better for patients with an isolated lesion in the ostium and body of the left main coronary artery where a single stent can be placed compared to bifurcation lesions that involve the ostium of the LAD and LCx.

CASE 6: MULTIVESSEL PCI IN A DIABETIC PATIENT

(Videos 297e-35 to 297e-42)

- A 58-year-old man presented with a NSTEMI.
- The patient has hyperlipidemia and type 2 diabetes mellitus treated with oral hypoglycemic agents.
- Diagnostic catheterization revealed two-vessel disease with a total occlusion of the second obtuse marginal branch that was felt to be

responsible for the patient's symptoms (culprit lesion). In addition, there was a high-grade stenosis in a large ramus intermedius branch, and the RCA had a significant stenosis in the midsegment of the vessel.

VIDEO 297e-35 Baseline angiogram of the left coronary circulation in the RAO view shows the total occlusion of the second obtuse marginal branch with delayed retrograde filling via collateral vessels and a high-grade stenosis in the ramus intermedius.

VIDEO 297e-36 A guidewire is passed through the total occlusion, and the lesion is pretreated with balloon angioplasty.

VIDEO 297e-37 Following placement of a drug-eluting stent in the lesion, the vessel is widely patent. A third obtuse marginal vessel, not previously seen, now fills faintly (Thrombolysis in Myocardial Infarction [TIMI] grade 1 flow) with contrast but was not treated.

VIDEO 297e-38 The ramus intermedius lesion was crossed with a guidewire and pretreated with balloon angioplasty.

VIDEO 297e-39 A drug-eluting stent is placed across the ramus lesion and deployed. The final result shows no residual stenosis in either the ramus or second obtuse marginal vessels.

VIDEO 297e-40 Baseline angiogram of the RCA shows a high-grade lesion in the midsegment of the vessel.

VIDEO 297e-41 The lesion was pretreated with balloon dilation followed by stent deployment.

VIDEO 297e-42 The final result shows no residual stenosis in the mid-RCA.

SUMMARY

- Multivessel PCI is performed commonly and may be done in one setting or staged with two or more procedures.
- Acute and long-term studies of multivessel PCI have shown comparable rates of death and myocardial infarction when compared to CABG, but a higher incidence of repeat revascularization as a result of restenosis is associated with PCI.
- In the randomized Bypass Angioplasty Revascularization Investigation (BARI) trial, diabetic patients treated with PCI had a worse long-term mortality than diabetic patients treated with CABG. However, the BARI registry found that in selected diabetic patients with favorable anatomy, PCI can result in outcomes equal to those observed with CABG.

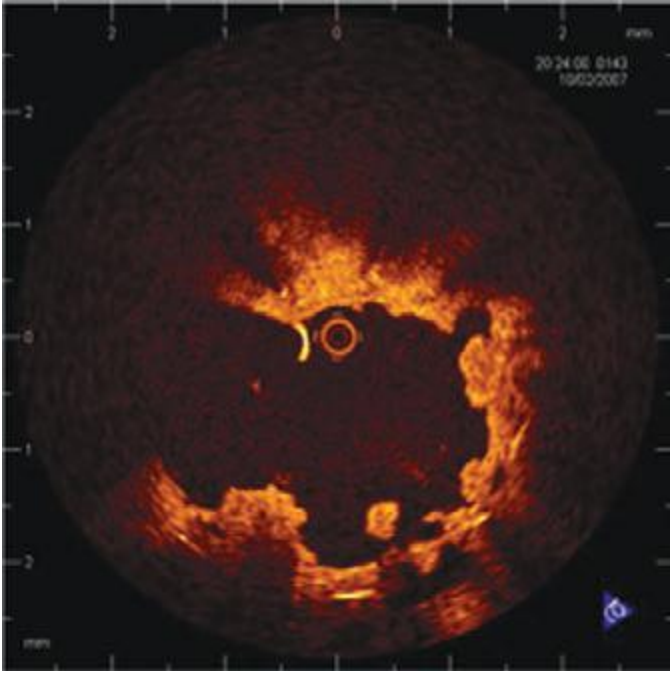


FIGURE 297e-8 Optical coherence tomography image following initial balloon dilation. Residual thrombus that is adherent to the stent struts is seen.

CASE 7: VERY LATE STENT THROMBOSIS OF A PROXIMAL LAD DRUG-ELUTING STENT

(Figs. 297e-8 and 297e-9; Videos 297e-43 to 297e-46)

- A 62-year-old male had a drug-eluting stent placed in a proximal LAD lesion to treat severe angina. He received dual antiplatelet therapy with aspirin and clopidogrel for 1 year and then discontinued clopidogrel per protocol.
- He remained asymptomatic until 15 months after the initial stent placement when he presented with severe chest pain due to an acute anterior STEMI.
- He was taken to the catheterization laboratory within 70 min of presentation, and his initial angiogram showed a total occlusion of the proximal LAD stent.

VIDEO 297e-43 Baseline angiogram showing a total occlusion of the proximal LAD within the drug-eluting stent and a significant stenosis at the origin of the LCx.

VIDEO 297e-44 The LAO view shows the LCx stenosis with a filling defect indicating that thrombus is present in the vessel lumen.

VIDEO 297e-45 The LAD lesion was crossed with a guidewire, which resulted in slow filling of the mid-LAD (TIMI 2 flow) and revealed thrombus filling the stent.

VIDEO 297e-46 The final result after LAD and LCx stenting. The LAD lesion was pretreated with balloon angioplasty, and a bare metal stent was deployed to cover the proximal lesion. The LCx ostial lesion was dilated with balloon angioplasty, and a bare metal stent was placed using a “V stenting” technique.

SUMMARY

- Stent thrombosis is an infrequent (1–2%) but serious complication of stent placement. It occurs most commonly within the first month, but rarely can occur as late as 1 year (0.2–0.6%) with bare metal stents. Very late stent thrombosis (VLST), which occurs after 1 year, is very rare with bare metal stents but can occur with drug-eluting stents.
- Premature discontinuation of dual antiplatelet therapy is the most common cause of early and late stent thrombosis; however, the etiology of VLST is not clear.
- The majority of patients with stent thrombosis present with acute coronary syndromes or STEMI; this presentation is associated with a high mortality rate (10%).
- Treatment is immediate PCI with balloon angioplasty or re-stenting.

CASE 8: TRANSCATHETER AORTIC VALVE REPLACEMENT

(Figs. 297e-10 to 297e-14; Videos 297e-47 to 297e-50)

- A 75-year-old female with symptomatic aortic stenosis and a valve area of 0.58 cm² by transthoracic echocardiogram.
- Chronic obstructive pulmonary disease (forced expiratory volume in 1 s [FEV₁] = 0.54) and other comorbidities contributed to an unacceptably high cardiac surgical risk (calculated logistic Euroscore = 29.57%) for aortic valve replacement.
- She was referred for transcatheter aortic valve replacement (TAVR) as part of a clinical trial.

VIDEO 297e-47 Aortogram shows patent coronary arteries and minimal aortic insufficiency.

VIDEO 297e-48 Balloon valvuloplasty is performed with rapid ventricular pacing at 180 beats/min.

VIDEO 297e-49 A 26-mm Edwards SAPIEN valve is positioned using fluoroscopic and transesophageal echo guidance and deployed.

VIDEO 297e-50 Aortogram after valve deployment shows a functional valve with mild aortic insufficiency and without impingement of the coronary ostia.

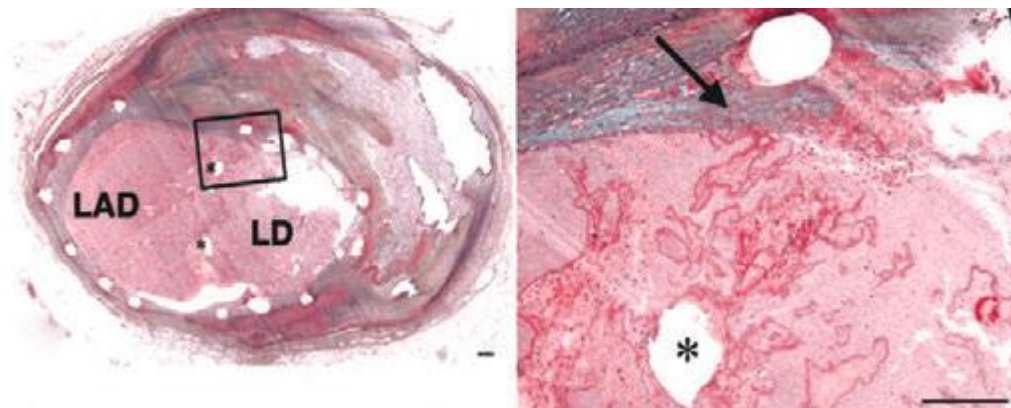


FIGURE 297e-9 Pathologic specimen of late stent thrombosis obtained at autopsy. Thrombus is seen filling the LAD vessel lumen and extending into a diagonal branch (LD). Stent struts occupied the space denoted by the asterisk (*) (left). A magnified view of the vessel reveals thrombus around the stent strut and neointima formation (arrow) (right).



FIGURE 297e-10 Transesophageal echocardiogram shows a calcified trileaflet aortic valve (left) with reduced leaflet excursion and a narrowed orifice in peak systole (right).



FIGURE 297e-11 Hemodynamically significant aortic (AO) stenosis. Simultaneous recording of AO and left ventricle (LV) pressures shows an 82 mmHg peak-to-peak gradient and a 63.3 mmHg mean gradient between the LV (154/9 mmHg) and AO (72/29 mmHg) pressures. This is consistent with an aortic valve area of 0.58 cm².

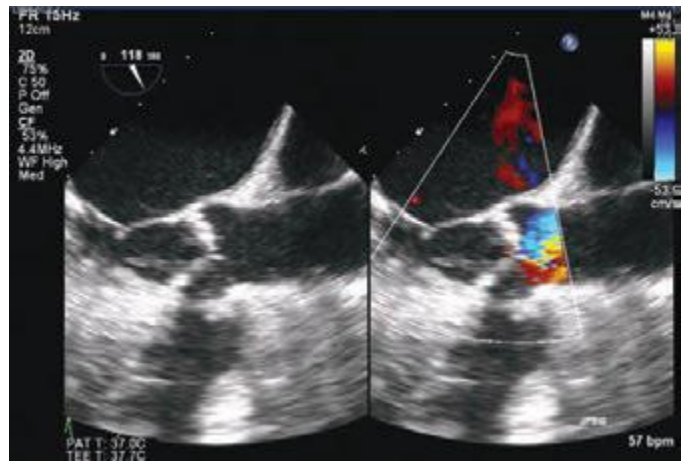


FIGURE 297e-13 The Edwards SAPIEN transcatheter heart valve. (Reprinted with permission from A Zajarias, AG Cribier: *J Am Coll Cardiol* 53:1829, 2009.)

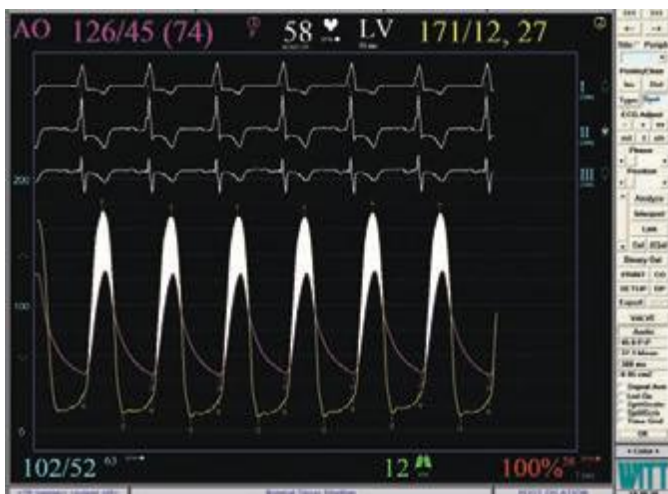


FIGURE 297e-12 After balloon valvuloplasty, the LV-AO mean pressure gradient decreased to 37.3 mmHg, indicating that the aortic valve area increased to 0.95 cm².

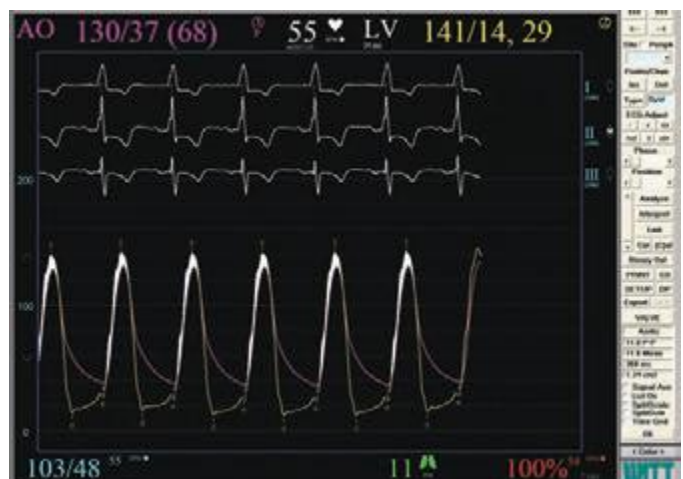


FIGURE 297e-14 Once the valve was deployed, the pressure gradient between the LV and AO decreased to 11.6 mmHg, and the functional valve area is 1.34 cm².

SUMMARY

- The prevalence of calcific aortic stenosis is 2–3% in individuals age ≥ 75 years.
- Symptomatic aortic stenosis is associated with an average survival of 2–3 years and an increased risk of sudden death; aortic valve replacement improves both symptoms and survival.
- In high-risk patients with severe aortic stenosis who are not surgical candidates, 1- and 5-year survival rates are ~62% and 38%, respectively.
- TAVR is approved in Europe and was recently approved in the United States as an alternative to surgical aortic valve replacement in high-risk patients.

(Case contributed with permission by Dr. Andrew C. Eisenhauer.)

CASE 9: ATRIAL SEPTAL DEFECT CLOSURE

(Figs. 297e-15 to 297e-21; Videos 297e-51 to 297e-53)

- A 48-year-old female with increased shortness of breath, exercise intolerance, and an 18-mm secundum ASD.
- Echocardiogram showed a dilated right atrium (RA) and right ventricle (RV) with evidence of right ventricular volume overload.

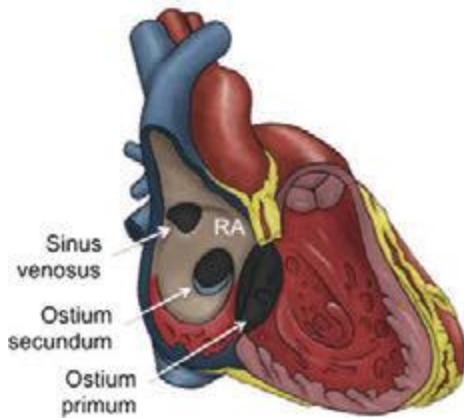


FIGURE 297e-15 Anatomic location of common atrial septal defects (ASDs). The most common ASDs are the sinus venosus, ostium secundum, and ostium primum ASDs. The sinus venosus ASD is located at the junction of the superior vena cava (SVC) and the right atrium (RA). This ASD is often associated with anomalous drainage of the right-side pulmonary veins into the RA instead of the left atrium. The secundum ASD is located at the foramen ovale and allows blood to flow between the RA and the left atrium. The primum ASD, also known as an atrioventricular septal defect, connects the RA and right ventricle with the left atrium and ventricle. (Illustration by Justin E. Tribuna.)

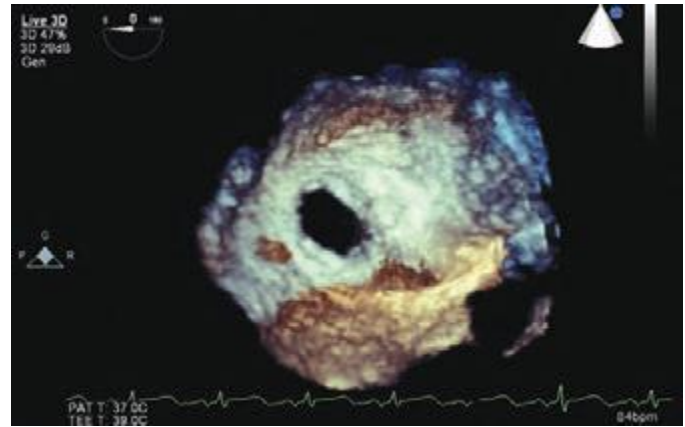


FIGURE 297e-17 Three-dimensional echocardiographic reconstruction of the secundum ASD. The ASD is round and has an acceptable margin of tissue to seat a septal occluder device.

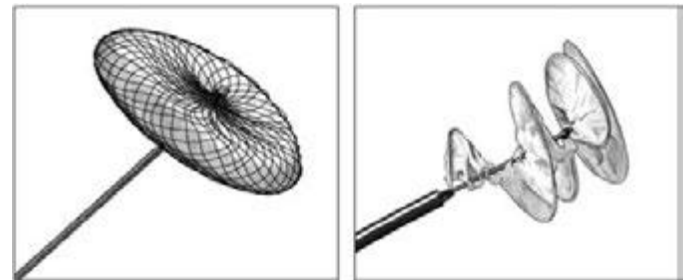


FIGURE 297e-18 ASD percutaneous closure devices. The Amplatzer® septal occluder (left) and the HELEX® septal occluder (right) are among the devices used for percutaneous closure of ASDs. The Amplatzer® septal occluder is a plug between two disks that are positioned on each side of the ASD to obstruct blood flow. The HELEX® Septal Occluder is positioned on each side of the ASD to allow circular rings covered with rapidly stretching polytetrafluoroethylene device to limit blood flow. The delivery catheters are detached and the devices are left in place. (Illustration by Justin E. Tribuna.)

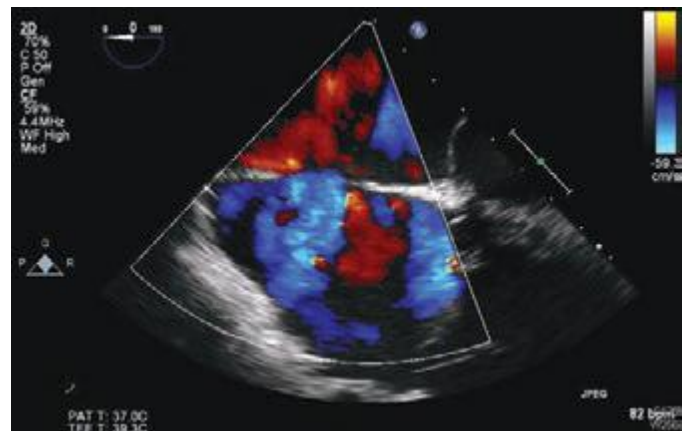
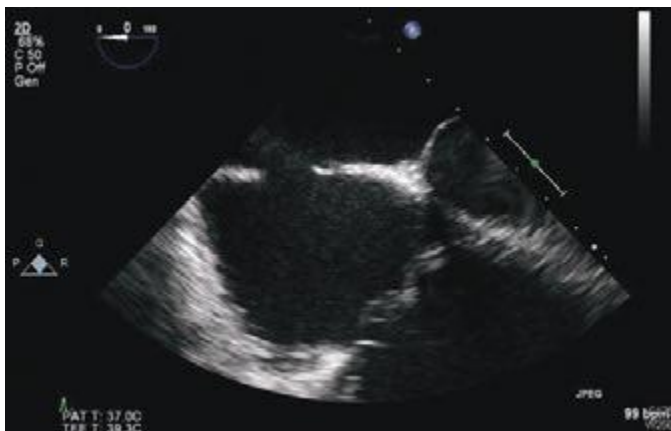


FIGURE 297e-16 Transesophageal echocardiogram of a secundum ASD. The ASD is seen as “dropout” in the interatrial septum between the left atrium (LA) and RA (left). Doppler color flow imaging shows blue in the RA consistent with left-to-right flow (right).



FIGURE 297e-19 Transesophageal echocardiogram showing sizing balloon (*left*) and no flow (*right*) across the atrial septal defect.

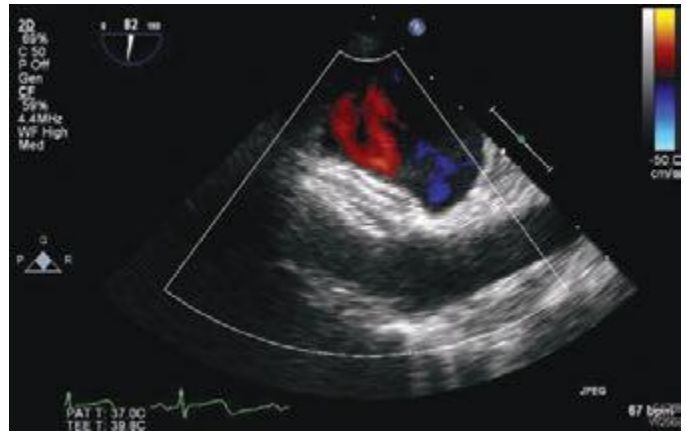
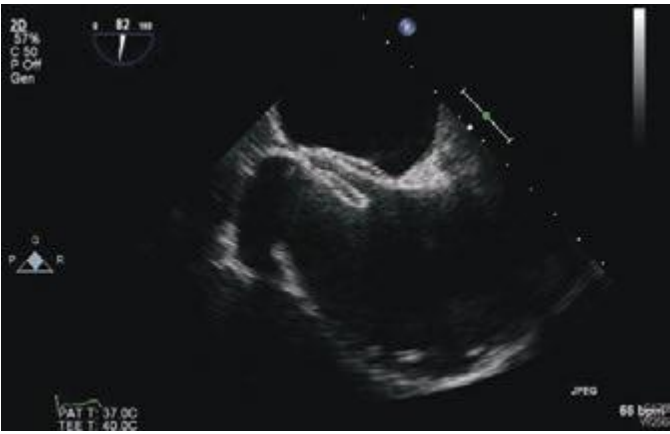


FIGURE 297e-20 Amplatzer® septal occluder in place (*left*). There is no blood flow across the device (*right*).

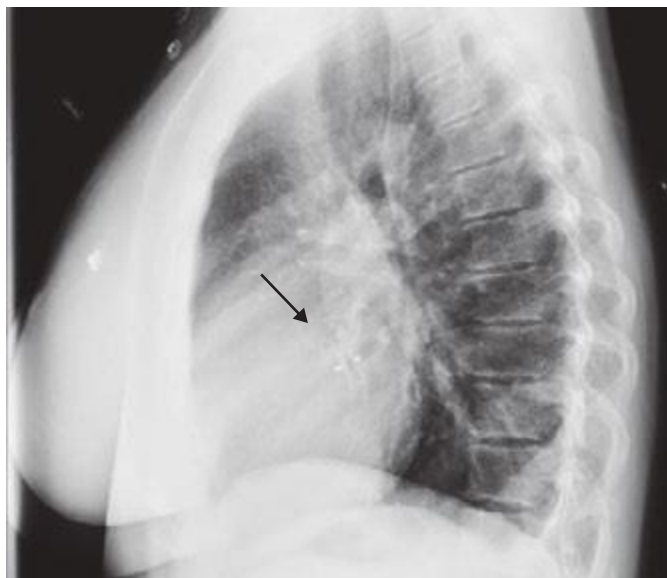


FIGURE 297e-21 Postprocedure lateral chest x-ray showing the Amplatzer® septal occluder in place.

- A shunt ratio (Qp/Qs) of 2.3:1 was determined at cardiac catheterization.
- Based on her symptoms, evidence of right-side chamber dilation, and a moderately sized ASD, the patient was referred for percutaneous closure of the ASD.
- Percutaneous closure of an ASD may be recommended for individuals with a secundum ASD and evidence of RA and RV enlargement with or without symptoms.
- Percutaneous closure is contraindicated in patients with irreversible pulmonary arterial hypertension and no left-to-right shunt. It is not recommended for closure of sinus venosus, coronary sinus, or primum ASDs.

VIDEO 297e-51 A sizing balloon is placed across the ASD.

VIDEO 297e-52 An Amplatzer® septal occluder is being positioned across the ASD.

VIDEO 297e-53 The two disks of the device in place across the ASD.

SUMMARY

- Unrepaired ASDs lead to signs and symptoms of increased pulmonary blood flow and right heart failure, dyspnea, exercise intolerance, fatigue, palpitations and atrial arrhythmias, and pulmonary infections.

- After the atrial septal occluder device is placed, patients are treated with antiplatelet agents and use antibiotic prophylaxis for certain procedures for 6 months. Follow-up echocardiograms to assess for device migration or erosion, residual shunting, thrombus, or pericardial effusion are recommended at 1 day, 1 month, 6 months, 1 year, and periodically thereafter.

(Case contributed with permission by Dr. Andrew C. Eisenhauer.)

298 Hypertensive Vascular Disease

Theodore A. Kotchen

Hypertension is one of the leading causes of the global burden of disease. Approximately 7.6 million deaths (13–15% of the total) and 92 million disability-adjusted life years worldwide were attributable to high blood pressure in 2001. Hypertension doubles the risk of cardiovascular diseases, including coronary heart disease (CHD), congestive heart failure (CHF), ischemic and hemorrhagic stroke, renal failure, and peripheral arterial disease. It often is associated with additional cardiovascular disease risk factors, and the risk of cardiovascular disease increases with the total burden of risk factors. Although anti-hypertensive therapy reduces the risks of cardiovascular and renal disease, large segments of the hypertensive population are either untreated or inadequately treated.

EPIDEMIOLOGY

Blood pressure levels, the rate of age-related increases in blood pressure, and the prevalence of hypertension vary among countries and among subpopulations within a country. Hypertension is present in all populations except for a small number of individuals living in developing countries. In industrialized societies, blood pressure increases steadily during the first two decades of life. In children and adolescents, blood pressure is associated with growth and maturation. Blood pressure “tracks” over time in children and between adolescence and young adulthood. In the United States, average systolic blood pressure is higher for men than for women during early adulthood, although among older individuals the age-related rate of rise is steeper for women. Consequently, among individuals age 60 and older, systolic blood pressures of women are higher than those of men. Among adults, diastolic blood pressure also increases progressively with age until ~55 years, after which it tends to decrease. The consequence is a widening of pulse pressure (the difference between systolic and diastolic blood pressure) beyond age 60.

In the United States, based on results of the National Health and Nutrition Examination Survey (NHANES), approximately 30% (age-adjusted prevalence) of adults, or at least 65 million individuals, have hypertension (defined as any one of the following: systolic blood pressure ≥ 140 mmHg, diastolic blood pressure ≥ 90 mmHg, taking antihypertensive medications). Hypertension prevalence is 33.5% in non-Hispanic blacks, 28.9% in non-Hispanic whites, and 20.7% in Mexican Americans. The likelihood of hypertension increases with age, and among individuals age ≥ 60 , the prevalence is 65.4%. Recent evidence suggests that the prevalence of hypertension in the United States may be increasing, possibly as a consequence of increasing obesity. The prevalence of hypertension and stroke mortality rates are higher in the southeastern United States than in other regions. In African Americans, hypertension appears earlier, is generally more severe, and results in higher rates of morbidity and mortality from stroke, left ventricular hypertrophy, CHF, and end-stage renal disease (ESRD) than in white Americans.

Both environmental and genetic factors may contribute to regional and racial variations in hypertension prevalence. Studies of societies undergoing “acculturation” and studies of migrants from a less to a more urbanized setting indicate a profound environmental contribution to blood pressure. Obesity and weight gain are strong, independent risk factors for hypertension. It has been estimated that 60% of hypertensives are $>20\%$ overweight. Among populations, hypertension prevalence is related to dietary NaCl intake, and the age-related increase in blood pressure may be augmented by a high NaCl intake. Low dietary intakes of calcium and potassium also may contribute to the risk of hypertension. The urine sodium-to-potassium ratio (an index of both sodium and potassium intakes) is a stronger correlate of blood pressure than is either sodium or potassium alone. Alcohol consumption, psychosocial stress, and low levels of physical activity also may contribute to hypertension.

Adoption, twin, and family studies document a significant heritable component to blood pressure levels and hypertension. Family studies controlling for a common environment indicate that blood pressure heritabilities are in the range 15–35%. In twin studies, heritability estimates of blood pressure are $\sim 60\%$ for males and 30–40% for females. High blood pressure before age 55 occurs 3.8 times more frequently among persons with a positive family history of hypertension. However, to date, only a fraction of high heritability estimates are accounted for by specific genetic determinants.

GENETIC CONSIDERATIONS



Although specific genetic variants have been identified in rare Mendelian forms of hypertension (Table 298–5), these variants are not applicable to the vast majority ($>98\%$) of patients with hypertension. For most individuals, it is likely that hypertension represents a polygenic disorder in which a combination of genes acts in concert with environmental exposures to make only a modest contribution to blood pressure. Further, different subsets of genes may lead to different phenotypes associated with hypertension, e.g., obesity, dyslipidemia, insulin resistance.

Several strategies are being used in the search for specific hypertension-related genes. Animal models (including selectively bred rats and congenic rat strains) provide a powerful approach for evaluating genetic loci and genes associated with hypertension. Comparative mapping strategies allow for the identification of syntenic genomic regions between the rat and human genomes that may be involved in blood pressure regulation. In association studies, different alleles (or combinations of alleles at different loci) of specific candidate genes or chromosomal regions are compared in hypertensive patients and normotensive control subjects. Current evidence suggests that genes that encode components of the renin-angiotensin-aldosterone system, along with angiotensinogen and angiotensin-converting enzyme (ACE) polymorphisms, may be related to hypertension and to blood pressure sensitivity to dietary NaCl. The alpha-adducin gene is thought to be associated with increased renal tubular absorption of sodium, and variants of this gene may be associated with hypertension and salt sensitivity of blood pressure. Other genes possibly related to hypertension include

genes encoding the AT_1 receptor, aldosterone synthase, atrial natriuretic peptide, and the β_2 adrenoreceptor. Genomewide association studies involve rapidly scanning markers across the entire genome to identify loci (not specific genes) associated with an observable trait (e.g., blood pressure) or a particular disease. This strategy has been facilitated by the availability of dense genotyping chips and the International HapMap. Results of candidate gene studies often have not been replicated, and in contrast to several other polygenic disorders, genomewide association studies have had limited success in identifying genetic determinants of hypertension.

Preliminary evidence suggests that there may also be genetic determinants of target organ damage attributed to hypertension. Family studies indicate significant heritability of left ventricular mass, and there is considerable individual variation in the responses of the heart to hypertension. Family studies and variations in candidate genes associated with renal damage suggest that genetic factors also may contribute to hypertensive nephropathy. Specific genetic variants have been linked to CHD and stroke.

In the future, it is possible that DNA analysis will predict individual risk for hypertension and target organ damage and will identify responders to specific classes of antihypertensive agents. However, with the exception of the rare, monogenic hypertensive diseases, the genetic variants associated with hypertension remain to be confirmed, and the intermediate steps by which these variants affect blood pressure remain to be determined.

MECHANISMS OF HYPERTENSION

To provide a framework for understanding the pathogenesis of and treatment options for hypertensive disorders, it is useful to understand factors involved in the regulation of both normal and elevated arterial pressure. Cardiac output and peripheral resistance are the two determinants of arterial pressure (Fig. 298-1). Cardiac output is determined by stroke volume and heart rate; stroke volume is related to myocardial contractility and to the size of the vascular compartment. Peripheral resistance is determined by functional and anatomic changes in small arteries (lumen diameter 100–400 μm) and arterioles.

INTRAVASCULAR VOLUME

Sodium is predominantly an extracellular ion and is a primary determinant of the extracellular fluid volume. When NaCl intake exceeds the capacity of the kidney to excrete sodium, vascular volume may initially expand and cardiac output may increase. However, many vascular beds have the capacity to autoregulate blood flow, and if constant blood flow is to be maintained in the face of increased arterial pressure, resistance within that bed must increase, since

$$\text{Blood flow} = \frac{\text{pressure across the vascular bed}}{\text{vascular resistance}}$$

The initial elevation of blood pressure in response to vascular volume expansion may be related to an increase of cardiac output; however, over time, peripheral resistance increases and cardiac output reverts toward normal. Whether this hypothesized sequence of events occurs in the pathogenesis of hypertension is not clear. What is clear is that salt can activate a number of neural, endocrine/paracrine, and vascular mechanisms, all of which have the potential to increase arterial pressure. The effect of sodium on blood pressure is related to the provision of sodium with chloride; nonchloride salts of sodium have

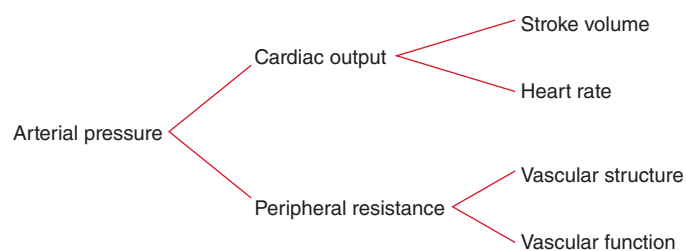


FIGURE 298-1 Determinants of arterial pressure.

little or no effect on blood pressure. As arterial pressure increases in response to a high NaCl intake, urinary sodium excretion increases and sodium balance is maintained at the expense of an increase in arterial pressure. The mechanism for this “pressure-natriuresis” phenomenon may involve a subtle increase in the glomerular filtration rate, decreased absorbing capacity of the renal tubules, and possibly hormonal factors such as atrial natriuretic factor. In individuals with an impaired capacity to excrete sodium, greater increases in arterial pressure are required to achieve natriuresis and sodium balance.

NaCl-dependent hypertension may be a consequence of a decreased capacity of the kidney to excrete sodium, due either to intrinsic renal disease or to increased production of a salt-retaining hormone (mineralocorticoid) resulting in increased renal tubular reabsorption of sodium. Renal tubular sodium reabsorption also may be augmented by increased neural activity to the kidney. In each of these situations, a higher arterial pressure may be required to achieve sodium balance. Conversely, salt-wasting disorders are associated with low blood pressure levels. ESRD is an extreme example of volume-dependent hypertension. In ~80% of these patients, vascular volume and hypertension can be controlled with adequate dialysis; in the other 20%, the mechanism of hypertension is related to increased activity of the renin-angiotensin system and is likely to be responsive to pharmacologic blockade of renin-angiotensin.

AUTONOMIC NERVOUS SYSTEM

Adrenergic reflexes modulate blood pressure over the short term, and adrenergic function, in concert with hormonal and volume-related factors, contributes to the long-term regulation of arterial pressure. Norepinephrine, epinephrine, and dopamine all play important roles in tonic and phasic cardiovascular regulation.

The activities of the adrenergic receptors are mediated by guanosine nucleotide-binding regulatory proteins (G proteins) and by intracellular concentrations of downstream second messengers. In addition to receptor affinity and density, physiologic responsiveness to catecholamines may be altered by the efficiency of receptor-effector coupling at a site “distal” to receptor binding. The receptor sites are relatively specific both for the transmitter substance and for the response that occupancy of the receptor site elicits. Based on their physiology and pharmacology, adrenergic receptors have been divided into two principal types: α and β . These types have been differentiated further into α_1 , α_2 , β_1 , and β_2 receptors. Recent molecular cloning studies have identified several additional subtypes. α Receptors are occupied and activated more avidly by norepinephrine than by epinephrine, and the reverse is true for β receptors. α_1 Receptors are located on postsynaptic cells in smooth muscle and elicit vasoconstriction. α_2 Receptors are localized on presynaptic membranes of postganglionic nerve terminals that synthesize norepinephrine. When activated by catecholamines, α_2 receptors act as negative feedback controllers, inhibiting further norepinephrine release. In the kidney, activation of α_1 -adrenergic receptors increases renal tubular reabsorption of sodium. Different classes of antihypertensive agents either inhibit α_1 receptors or act as agonists of α_2 receptors and reduce systemic sympathetic outflow. Activation of myocardial β_1 receptors stimulates the rate and strength of cardiac contraction and consequently increases cardiac output. β_1 Receptor activation also stimulates renin release from the kidney. Another class of antihypertensive agents acts by inhibiting β_1 receptors. Activation of β_2 receptors by epinephrine relaxes vascular smooth muscle and results in vasodilation.

Circulating catecholamine concentrations may affect the number of adrenoceptors in various tissues. Downregulation of receptors may be a consequence of sustained high levels of catecholamines and provides an explanation for decreasing responsiveness, or tachyphylaxis, to catecholamines. For example, orthostatic hypotension frequently is observed in patients with pheochromocytoma, possibly due to the lack of norepinephrine-induced vasoconstriction with assumption of the upright posture. Conversely, with chronic reduction of neurotransmitter substances, adrenoceptors may increase in number or be upregulated, resulting in increased responsiveness to the neurotransmitter. Chronic administration of agents that block adrenergic receptors may

result in upregulation, and abrupt withdrawal of those agents may produce a condition of temporary hypersensitivity to sympathetic stimuli. For example, clonidine is an antihypertensive agent that is a centrally acting α_2 agonist that inhibits sympathetic outflow. Rebound hypertension may occur with the abrupt cessation of clonidine therapy, probably as a consequence of upregulation of α_1 receptors.

Several reflexes modulate blood pressure on a minute-to-minute basis. One arterial baroreflex is mediated by stretch-sensitive sensory nerve endings in the carotid sinuses and the aortic arch. The rate of firing of these baroreceptors increases with arterial pressure, and the net effect is a decrease in sympathetic outflow, resulting in decreases in arterial pressure and heart rate. This is a primary mechanism for rapid buffering of acute fluctuations of arterial pressure that may occur during postural changes, behavioral or physiologic stress, and changes in blood volume. However, the activity of the baroreflex declines or adapts to sustained increases in arterial pressure such that the baroreceptors are reset to higher pressures. Patients with autonomic neuropathy and impaired baroreflex function may have extremely labile blood pressures with difficult-to-control episodic blood pressure spikes associated with tachycardia.

In both normal-weight and obese individuals, hypertension often is associated with increased sympathetic outflow. Based on recordings of postganglionic muscle nerve activity (detected by a microelectrode inserted in a peroneal nerve in the leg), sympathetic outflow tends to be higher in hypertensive than in normotensive individuals. Sympathetic outflow is increased in obesity-related hypertension and in hypertension associated with obstructive sleep apnea. Baroreceptor activation via electrical stimulation of carotid sinus afferent nerves lowers blood pressure in patients with “resistant” hypertension. Drugs that block the sympathetic nervous system are potent antihypertensive agents, indicating that the sympathetic nervous system plays a permissive, although not necessarily a causative, role in the maintenance of increased arterial pressure.

Pheochromocytoma is the most blatant example of hypertension related to increased catecholamine production, in this instance by a tumor. Blood pressure can be reduced by surgical excision of the tumor or by pharmacologic treatment with an α_1 receptor antagonist or with an inhibitor of tyrosine hydroxylase, the rate-limiting step in catecholamine biosynthesis.

RENIN-ANGIOTENSIN-ALDOSTERONE

The renin-angiotensin-aldosterone system contributes to the regulation of arterial pressure primarily via the vasoconstrictor properties of angiotensin II and the sodium-retaining properties of aldosterone. Renin is an aspartyl protease that is synthesized as an enzymatically inactive precursor, prorenin. Most renin in the circulation is synthesized in the renal afferent renal arteriole. Prorenin may be secreted directly into the circulation or may be activated within secretory cells and released as active renin. Although human plasma contains two to five times more prorenin than renin, there is no evidence that prorenin contributes to the physiologic activity of this system. There are three primary stimuli for renin secretion: (1) decreased NaCl transport in the distal portion of the thick ascending limb of the loop of Henle that abuts the corresponding afferent arteriole (macula densa), (2) decreased pressure or stretch within the renal afferent arteriole (baroreceptor mechanism), and (3) sympathetic nervous system stimulation of renin-secreting cells via β_1 adrenoceptors. Conversely, renin secretion is inhibited by increased NaCl transport in the thick ascending limb of the loop of Henle, by increased stretch within the renal afferent arteriole, and by β_1 receptor blockade. In addition, angiotensin II directly inhibits renin secretion due to angiotensin II type 1 receptors on juxtaglomerular cells, and renin secretion increases in response to pharmacologic blockade of either ACE or angiotensin II receptors.

Once released into the circulation, active renin cleaves a substrate, angiotensinogen, to form an inactive decapeptide, angiotensin I (Fig. 298-2). A converting enzyme, located primarily but not exclusively in the pulmonary circulation, converts angiotensin I to the active octapeptide, angiotensin II, by releasing the C-terminal histidyl-leucine dipeptide. The same converting enzyme cleaves a number

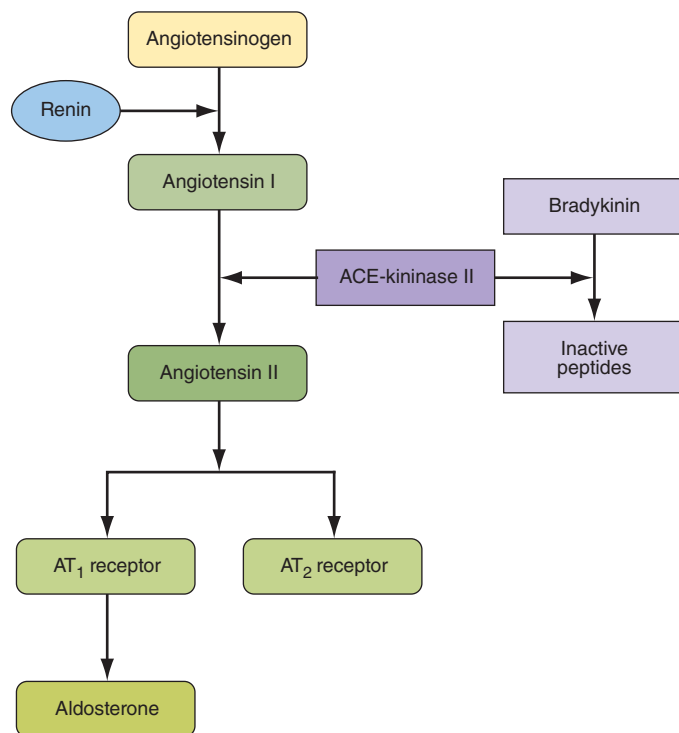


FIGURE 298-2 Renin-angiotensin-aldosterone axis. ACE, angiotensin-converting enzyme.

of other peptides, including and thereby inactivating the vasodilator bradykinin. Acting primarily through angiotensin II type 1 (AT_1) receptors on cell membranes, angiotensin II is a potent pressor substance, the primary tropic factor for the secretion of aldosterone by the adrenal zona glomerulosa, and a potent mitogen that stimulates vascular smooth muscle cell and myocyte growth. Independent of its hemodynamic effects, angiotensin II may play a role in the pathogenesis of atherosclerosis through a direct cellular action on the vessel wall. The angiotensin II type 2 (AT_2) receptor has the opposite functional effects of the AT_1 receptor. The AT_2 receptor induces vasodilation, sodium excretion, and inhibition of cell growth and matrix formation. Experimental evidence suggests that the AT_2 receptor improves vascular remodeling by stimulating smooth muscle cell apoptosis and contributes to the regulation of glomerular filtration rate. AT_1 receptor blockade induces an increase in AT_2 receptor activity.

Renin-secreting tumors are clear examples of renin-dependent hypertension. In the kidney, these tumors include benign hemangiopericytomas of the juxtaglomerular apparatus and, infrequently, renal carcinomas, including Wilms' tumors. Renin-producing carcinomas also have been described in lung, liver, pancreas, colon, and adrenals. In these instances, in addition to excision and/or ablation of the tumor, treatment of hypertension includes pharmacologic therapies targeted to inhibit angiotensin II production or action. Renovascular hypertension is another renin-mediated form of hypertension. Obstruction of the renal artery leads to decreased renal perfusion pressure, thereby stimulating renin secretion. Over time, possibly as a consequence of secondary renal damage, this form of hypertension may become less renin dependent.

Angiotensinogen, renin, and angiotensin II are also synthesized locally in many tissues, including the brain, pituitary, aorta, arteries, heart, adrenal glands, kidneys, adipocytes, leukocytes, ovaries, testes, uterus, spleen, and skin. Angiotensin II in tissues may be formed by the enzymatic activity of renin or by other proteases, e.g., tonin, chymase, and cathepsins. In addition to regulating local blood flow, tissue angiotensin II is a mitogen that stimulates growth and contributes to modeling and repair. Excess tissue angiotensin II may contribute to atherosclerosis, cardiac hypertrophy, and renal failure and consequently may be a target for pharmacologic therapy to prevent target organ damage.

Angiotensin II is the primary tropic factor regulating the synthesis and secretion of aldosterone by the zona glomerulosa of the adrenal

cortex. Aldosterone synthesis is also dependent on potassium, and aldosterone secretion may be decreased in potassium-depleted individuals. Although acute elevations of adrenocorticotropic hormone (ACTH) levels also increase aldosterone secretion, ACTH is not an important tropic factor for the chronic regulation of aldosterone.

Aldosterone is a potent mineralocorticoid that increases sodium reabsorption by amiloride-sensitive epithelial sodium channels (ENaC) on the apical surface of the principal cells of the renal cortical collecting duct (Chap. 332e). Electric neutrality is maintained by exchanging sodium for potassium and hydrogen ions. Consequently, increased aldosterone secretion may result in hypokalemia and alkalosis.

Cortisol also binds to the mineralocorticoid receptor but normally functions as a less potent mineralocorticoid than aldosterone because cortisol is converted to cortisone by the enzyme 11 β -hydroxysteroid dehydrogenase type 2. Cortisone has no affinity for the mineralocorticoid receptor. Primary aldosteronism is a compelling example of mineralocorticoid-mediated hypertension. In this disorder, adrenal aldosterone synthesis and release are independent of renin-angiotensin, and renin release is suppressed by the resulting volume expansion.

Mineralocorticoid receptors are expressed in a number of tissues in addition to the kidney, and mineralocorticoid receptor activation induces structural and functional alterations in the heart, kidney, and blood vessels, leading to myocardial fibrosis, nephrosclerosis, and vascular inflammation and remodeling, perhaps as a consequence of oxidative stress. These effects are amplified by a high salt intake. In animal models, high circulating aldosterone levels stimulate cardiac fibrosis and left ventricular hypertrophy, and spironolactone (an aldosterone antagonist) prevents aldosterone-induced myocardial fibrosis. Pathologic patterns of left ventricular geometry also have been associated with elevations of plasma aldosterone concentration in hypertensive patients. In patients with CHF, low-dose spironolactone reduces the risk of progressive heart failure and sudden death from cardiac causes by 30%. Due to a renal hemodynamic effect, in patients with primary aldosteronism, high circulating levels of aldosterone also may cause glomerular hyperfiltration and albuminuria. These renal effects are reversible after removal of the effects of excess aldosterone by adrenalectomy or spironolactone.

Increased activity of the renin-angiotensin-aldosterone axis is not invariably associated with hypertension. In response to a low-NaCl diet or to volume contraction, arterial pressure and volume homeostasis may be maintained by increased activity of the renin-angiotensin-aldosterone axis. Secondary aldosteronism (i.e., increased aldosterone secondary to increased renin-angiotensin), but not hypertension, also is observed in edematous states such as CHF and liver disease.

VASCULAR MECHANISMS

Vascular radius and compliance of resistance arteries are important determinants of arterial pressure. Resistance to flow varies inversely with the fourth power of the radius, and consequently, small decreases in lumen size significantly increase resistance. In hypertensive patients, structural, mechanical, or functional changes may reduce the lumen diameter of small arteries and arterioles. Remodeling refers to geometric alterations in the vessel wall without a change in vessel volume. Hypertrophic (increased cell size, and increased deposition of intercellular matrix) or eutrophic vascular remodeling results in decreased lumen size and, hence, increased peripheral resistance. Apoptosis, low-grade inflammation, and vascular fibrosis also contribute to remodeling. Lumen diameter also is related to elasticity of the vessel. Vessels with a high degree of elasticity can accommodate an increase of volume with relatively little change in pressure, whereas in a semirigid vascular system, a small increment in volume induces a relatively large increment of pressure.

Hypertensive patients may have stiffer arteries due to arteriosclerosis, and high systolic blood pressures and wide pulse pressures are a consequence of decreased vascular compliance. Due to arterial stiffness, central blood pressures (aortic, carotid) may not correspond to brachial artery pressures. Ejection of blood into the aorta elicits a pressure wave that is propagated at a given velocity. The forward travelling wave generates a reflected wave that travels backward toward

the ascending aorta. Although mean arterial pressure is determined by cardiac output and peripheral resistance, pulse pressure is related to the functional properties of large arteries and the amplitude and timing of the incident and reflected waves. Increased arterial stiffness results in increased pulse wave velocity of both incident and reflected waves. Due to the timing of these waves, the consequence is augmentation of aortic systolic pressure and a reduction of aortic diastolic pressure, i.e., an increase in pulse pressure. The aortic augmentation index, an index of arterial stiffening, is calculated as the ratio of central arterial pressure-to-pulse pressure. Central blood pressure may be measured directly by placing a sensor in the aorta or noninvasively by radial tonometry using commercially available devices. Central blood pressure and the aortic augmentation index are strong, independent predictors of cardiovascular disease and all-cause mortality.

Ion transport by vascular smooth muscle cells may contribute to hypertension-associated abnormalities of vascular tone and vascular growth, both of which are modulated by intracellular pH (pH_i). Three ion transport mechanisms participate in the regulation of pH_i : (1) Na^+ - H^+ exchange, (2) Na^+ -dependent HCO_3^- - Cl^- exchange, and (3) cation-independent HCO_3^- - Cl^- exchange. Based on measurements in cell types that are more accessible than vascular smooth muscle (e.g., leukocytes, erythrocytes, platelets, skeletal muscle), activity of the Na^+ - H^+ exchanger is increased in hypertension, and this may result in increased vascular tone by two mechanisms. First, increased sodium entry may lead to increased vascular tone by activating Na^+ - Ca^{2+} exchange and thereby increasing intracellular calcium. Second, increased pH_i enhances calcium sensitivity of the contractile apparatus, leading to an increase in contractility for a given intracellular calcium concentration. Additionally, increased Na^+ - H^+ exchange may stimulate growth of vascular smooth muscle cells by enhancing sensitivity to mitogens.

Vascular endothelial function also modulates vascular tone. The vascular endothelium synthesizes and releases several vasoactive substances, including nitric oxide, a potent vasodilator. Endothelium-dependent vasodilation is impaired in hypertensive patients. This impairment often is assessed with high-resolution ultrasonography before and after the hyperemic phase of reperfusion that follows 5 minutes of forearm ischemia. Alternatively, endothelium-dependent vasodilation may be assessed in response to an intra-arterially infused endothelium-dependent vasodilator, e.g., acetylcholine. Endothelin is a vasoconstrictor peptide produced by the endothelium, and orally active endothelin antagonists may lower blood pressure in patients with resistant hypertension.

Currently, it is not known if the hypertension-related vascular abnormalities of ion transport and endothelial function are primary alterations or secondary consequences of elevated arterial pressure. Limited evidence suggests that vascular compliance and endothelium-dependent vasodilation may be improved by aerobic exercise, weight loss, and antihypertensive agents. It remains to be determined whether these interventions affect arterial structure and stiffness via a blood pressure-independent mechanism and whether different classes of antihypertensive agents preferentially affect vascular structure and function.

PATHOLOGIC CONSEQUENCES OF HYPERTENSION

Hypertension is an independent predisposing factor for heart failure, coronary artery disease, stroke, renal disease, and peripheral arterial disease (PAD).

HEART

Heart disease is the most common cause of death in hypertensive patients. Hypertensive heart disease is the result of structural and functional adaptations leading to left ventricular hypertrophy, CHF, abnormalities of blood flow due to atherosclerotic coronary artery disease and microvascular disease, and cardiac arrhythmias.

Individuals with left ventricular hypertrophy are at increased risk for CHD, stroke, CHF, and sudden death. Aggressive control of hypertension can regress or reverse left ventricular hypertrophy and reduce

the risk of cardiovascular disease. It is not clear whether different classes of antihypertensive agents have an added impact on reducing left ventricular mass, independent of their blood pressure-lowering effect.

CHF may be related to systolic dysfunction, diastolic dysfunction, or a combination of the two. Abnormalities of diastolic function that range from asymptomatic heart disease to overt heart failure are common in hypertensive patients. Approximately one-third of patients with CHF have normal systolic function but abnormal diastolic function. Diastolic dysfunction is an early consequence of hypertension-related heart disease and is exacerbated by left ventricular hypertrophy and ischemia. Cardiac catheterization provides the most accurate assessment of diastolic function. Alternatively, diastolic function can be evaluated by several noninvasive methods, including echocardiography and radionuclide angiography.

BRAIN

Stroke is the second most frequent cause of death in the world; it accounts for 5 million deaths each year, with an additional 15 million persons having nonfatal strokes. Elevated blood pressure is the strongest risk factor for stroke. Approximately 85% of strokes are due to infarction, and the remainder are due to either intracerebral or subarachnoid hemorrhage. The incidence of stroke rises progressively with increasing blood pressure levels, particularly systolic blood pressure in individuals >65 years. Treatment of hypertension decreases the incidence of both ischemic and hemorrhagic strokes.

Hypertension also is associated with impaired cognition in an aging population, and longitudinal studies support an association between midlife hypertension and late-life cognitive decline. Hypertension-related cognitive impairment and dementia may be a consequence of a single infarct due to occlusion of a "strategic" larger vessel or multiple lacunar infarcts due to occlusive small vessel disease resulting in subcortical white matter ischemia. Several clinical trials suggest that antihypertensive therapy has a beneficial effect on cognitive function, although this remains an active area of investigation.

Cerebral blood flow remains unchanged over a wide range of arterial pressures (mean arterial pressure of 50–150 mmHg) through a process termed *autoregulation* of blood flow. In patients with the clinical syndrome of malignant hypertension, encephalopathy is related to failure of autoregulation of cerebral blood flow at the upper pressure limit, resulting in vasodilation and hyperperfusion. Signs and symptoms of hypertensive encephalopathy may include severe headache, nausea and vomiting (often of a projectile nature), focal neurologic signs, and alterations in mental status. Untreated, hypertensive encephalopathy may progress to stupor, coma, seizures, and death within hours. It is important to distinguish hypertensive encephalopathy from other neurologic syndromes that may be associated with hypertension, e.g., cerebral ischemia, hemorrhagic or thrombotic stroke, seizure disorder, mass lesions, pseudotumor cerebri, delirium tremens, meningitis, acute intermittent porphyria, traumatic or chemical injury to the brain, and uremic encephalopathy.

KIDNEY

The kidney is both a target and a cause of hypertension. Primary renal disease is the most common etiology of secondary hypertension. Mechanisms of kidney-related hypertension include a diminished capacity to excrete sodium, excessive renin secretion in relation to volume status, and sympathetic nervous system overactivity. Conversely, hypertension is a risk factor for renal injury and ESRD. The increased risk associated with high blood pressure is graded, continuous, and present throughout the distribution of blood pressure above optimal pressure. Renal risk appears to be more closely related to systolic than to diastolic blood pressure, and black men are at greater risk than white men for developing ESRD at every level of blood pressure.

Atherosclerotic, hypertension-related vascular lesions in the kidney primarily affect preglomerular arterioles, resulting in ischemic changes in the glomeruli and postglomerular structures. Glomerular injury also may be a consequence of direct damage to the glomerular capillaries due to glomerular hyperperfusion. Studies of hypertension-related

1616 renal damage, primarily in experimental animals, suggest that loss of autoregulation of renal blood flow at the afferent arteriole results in transmission of elevated pressures to an unprotected glomerulus with ensuing hyperfiltration, hypertrophy, and eventual focal segmental glomerular sclerosis. With progressive renal injury there is a loss of autoregulation of renal blood flow and glomerular filtration rate, resulting in a lower blood pressure threshold for renal damage and a steeper slope between blood pressure and renal damage. The result may be a vicious cycle of renal damage and nephron loss leading to more severe hypertension, glomerular hyperfiltration, and further renal damage. Glomerular pathology progresses to glomerulosclerosis, and eventually the renal tubules may also become ischemic and gradually atrophic. The renal lesion associated with malignant hypertension consists of fibrinoid necrosis of the afferent arterioles, sometimes extending into the glomerulus, and may result in focal necrosis of the glomerular tuft.

Clinically, macroalbuminuria (a random urine albumin/creatinine ratio >300 mg/g) or microalbuminuria (a random urine albumin/creatinine ratio 30–300 mg/g) are early markers of renal injury. These are also risk factors for renal disease progression and cardiovascular disease.

PERIPHERAL ARTERIES

In addition to contributing to the pathogenesis of hypertension, blood vessels are a target organ for atherosclerotic disease secondary to long-standing elevated blood pressure. In hypertensive patients, vascular disease is a major contributor to stroke, heart disease, and renal failure. Further, hypertensive patients with arterial disease of the lower extremities are at increased risk for future cardiovascular disease. Although patients with stenotic lesions of the lower extremities may be asymptomatic, intermittent claudication is the classic symptom of PAD. The ankle-brachial index is a useful approach for evaluating PAD and is defined as the ratio of noninvasively assessed ankle to brachial (arm) systolic blood pressure. An ankle-brachial index <0.90 is considered diagnostic of PAD and is associated with >50% stenosis in at least one major lower limb vessel. An ankle-brachial index <0.80 is associated with elevated blood pressure, particularly systolic blood pressure.

DEFINING HYPERTENSION

From an epidemiologic perspective, there is no obvious level of blood pressure that defines hypertension. In adults, there is a continuous, incremental risk of cardiovascular disease, stroke, and renal disease across levels of both systolic and diastolic blood pressure. The Multiple Risk Factor Intervention Trial (MRFIT), which included >350,000 male participants, demonstrated a continuous and graded influence of both systolic and diastolic blood pressure on CHD mortality, extending down to systolic blood pressures of 120 mmHg. Similarly, results of a meta-analysis involving almost 1 million participants indicate that ischemic heart disease mortality, stroke mortality, and mortality from other vascular causes are directly related to the height of the blood pressure, beginning at 115/75 mmHg, without evidence of a threshold. Cardiovascular disease risk doubles for every 20-mmHg increase in systolic and 10-mmHg increase in diastolic pressure. Among older individuals, systolic blood pressure and pulse pressure are more powerful predictors of cardiovascular disease than is diastolic blood pressure.

Clinically, hypertension may be defined as that level of blood pressure at which the institution of therapy reduces blood pressure–related morbidity and mortality. Current clinical criteria for defining hypertension generally are based on the average of two or more seated blood pressure readings during each of two or more outpatient visits. A recent classification recommends blood pressure criteria for defining normal blood pressure, prehypertension, hypertension (stages I and II), and isolated systolic hypertension, which is frequent among the elderly (Table 298-1). In children and adolescents, hypertension generally is defined as systolic and/or diastolic blood pressure consistently >95th percentile for age, sex, and height. Blood pressures between the 90th

TABLE 298-1 BLOOD PRESSURE CLASSIFICATION

Blood Pressure Classification	Systolic, mmHg	Diastolic, mmHg
Normal	<120	and <80
Prehypertension	120–139	or 80–89
Stage 1 hypertension	140–159	or 90–99
Stage 2 hypertension	≥160	or ≥100
Isolated systolic hypertension	≥140	and <90

Source: Adapted from AV Chobanian et al: JAMA 289:2560, 2003.

and 95th percentiles are considered prehypertensive and are an indication for lifestyle interventions.

Home blood pressure and average 24-h ambulatory blood pressure measurements are generally lower than clinic blood pressures. Because ambulatory blood pressure recordings yield multiple readings throughout the day and night, they provide a more comprehensive assessment of the vascular burden of hypertension than do a limited number of office readings. Increasing evidence suggests that home blood pressures, including 24-h blood pressure recordings, more reliably predict target organ damage than do office blood pressures. Blood pressure tends to be higher in the early morning hours, soon after waking, than at other times of day. Myocardial infarction and stroke are more common in the early morning hours. Nighttime blood pressures are generally 10–20% lower than daytime blood pressures, and an attenuated nighttime blood pressure “dip” may be associated with increased cardiovascular disease risk. Recommended criteria for a diagnosis of hypertension, based on 24-h blood pressure monitoring, are average awake blood pressure ≥135/85 mmHg and asleep blood pressure ≥120/75 mmHg. These levels approximate a clinic blood pressure of 140/90 mmHg.

Approximately 15–20% of patients with stage 1 hypertension (as defined in Table 298-1) based on office blood pressures have average ambulatory readings <135/85 mmHg. This phenomenon, so-called white coat hypertension, also may be associated with an increased risk of target organ damage, although to a lesser extent than in individuals with elevated office and ambulatory readings. Individuals with white coat hypertension are also at increased risk for developing sustained hypertension.

CLINICAL DISORDERS OF HYPERTENSION

Depending on methods of patient ascertainment, ~80–95% of hypertensive patients are diagnosed as having primary, or “essential,” hypertension. In the remaining 5–20% of hypertensive patients, a specific underlying disorder causing the elevation of blood pressure can be identified (Tables 298-2 and 298-3). In individuals with “secondary” hypertension, a specific mechanism for the blood pressure elevation is often more apparent.

PRIMARY HYPERTENSION

Primary hypertension tends to be familial and is likely to be the consequence of an interaction between environmental and genetic factors. The prevalence of primary hypertension increases with age, and individuals with relatively high blood pressures at younger ages are at increased risk for the subsequent development of hypertension. It is

TABLE 298-2 SYSTOLIC HYPERTENSION WITH WIDE PULSE PRESSURE

1. Decreased vascular compliance (arteriosclerosis)
2. Increased cardiac output
 - a. Aortic regurgitation
 - b. Thyrotoxicosis
 - c. Hyperkinetic heart syndrome
 - d. Fever
 - e. Arteriovenous fistula
 - f. Patent ductus arteriosus

TABLE 298-3 SECONDARY CAUSES OF SYSTOLIC AND DIASTOLIC HYPERTENSION

Renal	Parenchymal diseases, renal cysts (including polycystic kidney disease), renal tumors (including renin-secreting tumors), obstructive uropathy
Renovascular	Arteriosclerotic, fibromuscular dysplasia
Adrenal	Primary aldosteronism, Cushing's syndrome, 17 α -hydroxylase deficiency, 11 β -hydroxylase deficiency, 11-hydroxysteroid dehydrogenase deficiency (licorice), pheochromocytoma
Aortic coarctation	
Obstructive sleep apnea	
Preeclampsia/eclampsia	
Neurogenic	Psychogenic, diencephalic syndrome, familial dysautonomia, polyneuritis (acute porphyria, lead poisoning), acute increased intracranial pressure, acute spinal cord section
Miscellaneous endocrine	Hypothyroidism, hyperthyroidism, hypercalcemia, acromegaly
Medications	High-dose estrogens, adrenal steroids, decongestants, appetite suppressants, cyclosporine, tricyclic antidepressants, monoamine oxidase inhibitors, erythropoietin, nonsteroidal anti-inflammatory agents, cocaine
Mendelian forms of hypertension	See Table 298-4

likely that primary hypertension represents a spectrum of disorders with different underlying pathophysiologies. In the majority of patients with established hypertension, peripheral resistance is increased and cardiac output is normal or decreased; however, in younger patients with mild or labile hypertension, cardiac output may be increased and peripheral resistance may be normal.

When plasma renin activity (PRA) is plotted against 24-h sodium excretion, ~10–15% of hypertensive patients have high PRA and 25% have low PRA. High-renin patients may have a vasoconstrictor form of hypertension, whereas low-renin patients may have volume-dependent hypertension. Inconsistent associations between plasma aldosterone and blood pressure have been described in patients with primary hypertension. The association between aldosterone and blood pressure is more striking in African Americans, and PRA tends to be low in hypertensive African Americans. This raises the possibility that subtle increases in aldosterone may contribute to hypertension in at least some groups of patients who do not have overt primary aldosteronism. Furthermore, spironolactone, an aldosterone antagonist, may be a particularly effective antihypertensive agent for some patients with primary hypertension, including some patients with “drug-resistant” hypertension.

OBESITY AND THE METABOLIC SYNDROME

(See also Chap. 422) There is a well-documented association between obesity (body mass index >30 kg/m²) and hypertension. Further, cross-sectional studies indicate a direct linear correlation between body weight (or body mass index) and blood pressure. Centrally located body fat is a more important determinant of blood pressure elevation than is peripheral body fat. In longitudinal studies, a direct correlation exists between change in weight and change in blood pressure over time. Sixty percent of hypertensive adults are more than 20% overweight. It has been established that 60–70% of hypertension in adults may be directly attributable to adiposity.

Hypertension and dyslipidemia frequently occur together and in association with resistance to insulin-stimulated glucose uptake. This clustering of risk factors is often, but not invariably, associated with obesity, particularly abdominal obesity. Insulin resistance also is associated with an unfavorable imbalance in the endothelial production of mediators that regulate platelet aggregation, coagulation, fibrinolysis, and vessel tone. When these risk factors cluster, the risks for CHD, stroke, diabetes, and cardiovascular disease mortality are increased further.

Depending on the populations studied and the methodologies for defining insulin resistance, ~25–50% of nonobese, nondiabetic hypertensive persons are insulin resistant. The constellation of insulin resistance, abdominal obesity, hypertension, and dyslipidemia has been designated as the *metabolic syndrome*. As a group, first-degree relatives of patients with primary hypertension are also insulin resistant, and hyperinsulinemia (a surrogate marker of insulin resistance) may predict the eventual development of hypertension and cardiovascular disease. Although the metabolic syndrome may in part be heritable as a polygenic condition, the expression of the syndrome is modified by environmental factors, such as degree of physical activity and diet. Insulin sensitivity increases and blood pressure decreases in response to weight loss. The recognition that cardiovascular disease risk factors tend to cluster within individuals has important implications for the evaluation and treatment of hypertension. Evaluation of both hypertensive patients and individuals at risk for developing hypertension should include assessment of overall cardiovascular disease risk. Similarly, introduction of lifestyle modification strategies and drug therapies should address overall risk and not focus exclusively on hypertension.

RENAL PARENCHYMAL DISEASES

Virtually all disorders of the kidney may cause hypertension (Table 298-3), and renal disease is the most common cause of secondary hypertension. Hypertension is present in >80% of patients with chronic renal failure. In general, hypertension is more severe in glomerular diseases than in interstitial diseases such as chronic pyelonephritis. Conversely, hypertension may cause nephrosclerosis, and in some instances it may be difficult to determine whether hypertension or renal disease was the initial disorder. Proteinuria >1000 mg/d and an active urine sediment are indicative of primary renal disease. In either instance, the goals are to control blood pressure and retard the rate of progression of renal dysfunction.

RENOVASCULAR HYPERTENSION

Hypertension due to an occlusive lesion of a renal artery, renovascular hypertension, is a potentially curable form of hypertension. In the initial stages, the mechanism of hypertension generally is related to activation of the renin-angiotensin system. However, renin activity and other components of the renin-angiotensin system may be elevated only transiently; over time, recruitment of other pressure mechanisms may contribute to elevated arterial pressure. Two groups of patients are at risk for this disorder: older arteriosclerotic patients who have a plaque obstructing the renal artery, frequently at its origin, and patients with fibromuscular dysplasia. Atherosclerosis accounts for the large majority of patients with renovascular hypertension. Although fibromuscular dysplasia may occur at any age, it has a strong predilection for young white women. The prevalence in females is eightfold that in males. There are several histologic variants of fibromuscular dysplasia, including medial fibroplasia, perimedial fibroplasia, medial hyperplasia, and intimal fibroplasia. Medial fibroplasia is the most common variant and accounts for approximately two-thirds of patients. The lesions of fibromuscular dysplasia are frequently bilateral and, in contrast to atherosclerotic renovascular disease, tend to affect more distal portions of the renal artery.

Several clues from the history and physical examination may suggest renovascular hypertension. The diagnosis should be considered in patients with other evidence of atherosclerotic vascular disease. Although response to antihypertensive therapy does not exclude the diagnosis, severe or refractory hypertension, recent loss of hypertension control or recent onset of moderately severe hypertension, and unexplained deterioration of renal function or deterioration of renal function associated with an ACE inhibitor should raise the possibility of renovascular hypertension. Approximately 50% of patients with renovascular hypertension have an abdominal or flank bruit, and the bruit is more likely to be hemodynamically significant if it lateralizes or extends throughout systole into diastole.

If blood pressure is adequately controlled with a simple antihypertensive regimen and renal function remains stable, there may be little

1618 impetus to pursue an evaluation for renal artery stenosis, particularly in an older patient with atherosclerotic disease and comorbid conditions. Patients with long-standing hypertension, advanced renal insufficiency, or diabetes mellitus are less likely to benefit from renal vascular repair. The most effective medical therapies include an ACE inhibitor or an angiotensin II receptor blocker; however, these agents decrease glomerular filtration rate in a stenotic kidney owing to efferent renal arteriolar dilation. In the presence of bilateral renal artery stenosis or renal artery stenosis to a solitary kidney, progressive renal insufficiency may result from the use of these agents. Importantly, the renal insufficiency is generally reversible after discontinuation of the offending drug.

If renal artery stenosis is suspected and if the clinical condition warrants an intervention such as percutaneous transluminal renal angioplasty (PTRA), placement of a vascular endoprosthesis (stent), or surgical renal revascularization, imaging studies should be the next step in the evaluation. As a screening test, renal blood flow may be evaluated with a radionuclide [¹³¹I]-orthoiodohippurate (OIH) scan, or glomerular filtration rate may be evaluated with a [^{99m}Tc]-diethylenetriamine pentaacetic acid (DTPA) scan before and after a single dose of captopril (or another ACE inhibitor). The following are consistent with a positive study: (1) decreased relative uptake by the involved kidney, which contributes <40% of total renal function, (2) delayed uptake on the affected side, and (3) delayed washout on the affected side. In patients with normal, or nearly normal, renal function, a normal captopril renogram essentially excludes functionally significant renal artery stenosis; however, its usefulness is limited in patients with renal insufficiency (creatinine clearance <20 mL/min) or bilateral renal artery stenosis. Additional imaging studies are indicated if the scan is positive. Doppler ultrasound of the renal arteries produces reliable estimates of renal blood flow velocity and offers the opportunity to track a lesion over time. Positive studies usually are confirmed at angiography, whereas false-negative results occur frequently, particularly in obese patients. Gadolinium-contrast magnetic resonance angiography offers clear images of the proximal renal artery but may miss distal lesions. An advantage is the opportunity to image the renal arteries with an agent that is not nephrotoxic. Contrast arteriography remains the “gold standard” for evaluation and identification of renal artery lesions. Potential risks include nephrotoxicity, particularly in patients with diabetes mellitus or preexisting renal insufficiency.

Some degree of renal artery obstruction may be observed in almost 50% of patients with atherosclerotic disease, and there are several approaches for evaluating the functional significance of such a lesion to predict the effect of vascular repair on blood pressure control and renal function. Each approach has varying degrees of sensitivity and specificity, and no single test is sufficiently reliable to determine a causal relationship between a renal artery lesion and hypertension. Functionally significant lesions generally occlude more than 70% of the lumen of the affected renal artery. On angiography, the presence of collateral vessels to the ischemic kidney suggests a functionally significant lesion. A lateralizing renal vein renin ratio (ratio >1.5 of affected side/contralateral side) has a 90% predictive value for a lesion that would respond to vascular repair; however, the false-negative rate for blood pressure control is 50–60%. Measurement of the pressure gradient across a renal artery lesion does not reliably predict the response to vascular repair.

In the final analysis, a decision concerning vascular repair vs. medical therapy and the type of repair procedure should be individualized. Patients with fibromuscular disease have more favorable outcomes than do patients with atherosclerotic lesions, presumably owing to their younger age, shorter duration of hypertension, and less systemic disease. Because of its low risk-versus-benefit ratio and high success rate (improvement or cure of hypertension in 90% of patients and restenosis rate of 10%), PTRA is the initial treatment of choice for these patients. Surgical revascularization may be undertaken if PTRA is unsuccessful or if a branch lesion is present. In atherosclerotic patients, vascular repair should be considered if blood pressure cannot be controlled adequately despite optimal medical therapy or if renal function deteriorates. Surgery may be the preferred initial approach

for younger atherosclerotic patients without comorbid conditions; however, for most atherosclerotic patients, depending on the location of the lesion, the initial approach may be PTRA and/or stenting. Surgical revascularization may be indicated if these approaches are unsuccessful, the vascular lesion is not amenable to PTRA or stenting, or concomitant aortic surgery is required, e.g., to repair an aneurysm. A National Institutes of Health–sponsored prospective, randomized clinical trial is in progress comparing medical therapy alone with medical therapy plus renal artery stenting regarding Cardiovascular Outcomes for Renal Atherosclerotic Lesions (CORAL).

PRIMARY ALDOSTERONISM

Excess aldosterone production due to primary aldosteronism is a potentially curable form of hypertension. In patients with primary aldosteronism, increased aldosterone production is independent of the renin-angiotensin system, and the consequences are sodium retention, hypertension, hypokalemia, and low PRA. The reported prevalence of this disorder varies from <2% to ~15% of hypertensive individuals. In part, this variation is related to the intensity of screening and the criteria for establishing the diagnosis.

History and physical examination provide little information about the diagnosis. The age at the time of diagnosis is generally the third through fifth decade. Hypertension is usually mild to moderate but occasionally may be severe; primary aldosteronism should be considered in all patients with refractory hypertension. Hypertension in these patients may be associated with glucose intolerance. Most patients are asymptomatic; however, infrequently, polyuria, polydipsia, paresthesias, or muscle weakness may be present as a consequence of hypokalemic alkalosis. Although aldosterone is a salt-retaining hormone, patients with primary aldosteronism rarely have edema. Renal dysfunction and cardiovascular disease are strikingly increased in patients with primary aldosteronism compared to those with primary hypertension.

In a hypertensive patient with unprovoked hypokalemia (i.e., unrelated to diuretics, vomiting, or diarrhea), the prevalence of primary aldosteronism approaches 40–50%. In patients on diuretics, serum potassium <3.1 mmol/L (<3.1 meq/L) also raises the possibility of primary aldosteronism; however, serum potassium is an insensitive and nonspecific screening test. Serum potassium is normal in ~25% of patients subsequently found to have an aldosterone-producing adenoma, and higher percentages of patients with other etiologies of primary aldosteronism are not hypokalemic. Additionally, hypokalemic hypertension may be a consequence of secondary aldosteronism, other mineralocorticoid- and glucocorticoid-induced hypertensive disorders, and pheochromocytoma.

The ratio of plasma aldosterone to plasma renin activity (PA/PRA) is a useful screening test. These measurements preferably are obtained in ambulatory patients in the morning. A ratio >30:1 in conjunction with a plasma aldosterone concentration >555 pmol/L (>20 ng/dL) reportedly has a sensitivity of 90% and a specificity of 91% for an aldosterone-producing adenoma. In a Mayo Clinic series, an aldosterone-producing adenoma subsequently was confirmed surgically in >90% of hypertensive patients with a PA/PRA ratio ≥20 and a plasma aldosterone concentration ≥415 pmol/L (≥15 ng/dL). There are, however, several caveats to interpreting the ratio. The cutoff for a “high” ratio is laboratory- and assay-dependent. Some antihypertensive agents may affect the ratio (e.g., aldosterone antagonists, angiotensin receptor antagonists, and ACE inhibitors may increase renin; aldosterone antagonists may increase aldosterone). Current recommendations are to withdraw aldosterone antagonists for at least 4–6 weeks before obtaining these measurements. Because aldosterone biosynthesis is potassium-dependent, hypokalemia should be corrected with oral potassium supplements prior to screening. With these caveats, the ratio has been reported to be useful as a screening test in measurements obtained with patients taking their usual antihypertensive medications. A high ratio in the absence of an elevated plasma aldosterone level is considerably less specific for primary aldosteronism since many patients with primary hypertension have low renin levels in this setting, particularly African Americans and elderly patients.

In patients with renal insufficiency, the ratio may also be elevated because of decreased aldosterone clearance. In patients with an elevated PA/PRA ratio, the diagnosis of primary aldosteronism can be confirmed by demonstrating failure to suppress plasma aldosterone to <277 pmol/L (<10 ng/dL) after IV infusion of 2 L of isotonic saline over 4 h; post-saline infusion plasma aldosterone values between 138 and 277 pmol/L (5–10 ng/dL) are not determinant. Alternative confirmatory tests include failure to suppress aldosterone (based on test specific criteria) in response to an oral NaCl load, fludrocortisone, or captopril.

Several sporadic and familial adrenal abnormalities may culminate in the syndrome of primary aldosteronism, and appropriate therapy depends on the specific etiology. The two most common causes of sporadic primary aldosteronism are an aldosterone-producing adenoma and bilateral adrenal hyperplasia. Together, they account for >90% of all patients with primary aldosteronism. The tumor is almost always unilateral, and most often measures <3 cm in diameter. Most of the remainder of these patients have bilateral adrenocortical hyperplasia (idiopathic hyperaldosteronism). Rarely, primary aldosteronism may be caused by an adrenal carcinoma or an ectopic malignancy, e.g., ovarian arrhenoblastoma. Most aldosterone-producing carcinomas, in contrast to adrenal adenomas and hyperplasia, produce excessive amounts of other adrenal steroids in addition to aldosterone. Functional differences in hormone secretion may assist in the diagnosis of adenoma vs. hyperplasia. Aldosterone biosynthesis is more responsive to ACTH in patients with adenoma and more responsive to angiotensin in patients with hyperplasia. Consequently, patients with adenoma tend to have higher plasma aldosterone in the early morning that decreases during the day, reflecting the diurnal rhythm of ACTH, whereas plasma aldosterone tends to increase with upright posture in patients with hyperplasia, reflecting the normal postural response of the renin-angiotensin-aldosterone axis. However, there is overlap in the ability of these measurements to discriminate between adenoma and hyperplasia. Rare familial forms of primary aldosteronism include glucocorticoid-remediable primary aldosteronism and familial aldosteronism types II and III. Genetic testing may assist in the diagnosis of these familial disorders.

Adrenal computed tomography (CT) should be carried out in all patients diagnosed with primary aldosteronism. High-resolution CT may identify tumors as small as 0.3 cm and is positive for an adrenal tumor 90% of the time. If the CT is not diagnostic, an adenoma may be detected by adrenal scintigraphy with 6 β -[I¹³¹] iodomethyl-19-norcholesterol after dexamethasone suppression (0.5 mg every 6 h for 7 days); however, this technique has decreased sensitivity for adenomas <1.5 cm.

When carried out by an experienced radiologist, bilateral adrenal venous sampling for measurement of plasma aldosterone is the most accurate means of differentiating unilateral from bilateral forms of primary aldosteronism. The sensitivity and specificity of adrenal venous sampling (95% and 100%, respectively) for detecting unilateral aldosterone hypersecretion are superior to those of adrenal CT; success rates are 90–96%, and complication rates are <2.5%. One frequently used protocol involves sampling for aldosterone and cortisol levels in response to ACTH stimulation. An ipsilateral/contralateral aldosterone ratio >4, with symmetric ACTH-stimulated cortisol levels, is indicative of unilateral aldosterone production.

Hypertension generally is responsive to surgery in patients with adenoma but not in patients with bilateral adrenal hyperplasia. Unilateral adrenalectomy, often done via a laparoscopic approach, is curative in 40–70% of patients with an adenoma. Transient hypoadosteronism may occur up to 3 months postoperatively, resulting in hyperkalemia. Potassium should be monitored during this time, and hyperkalemia should be treated with potassium-wasting diuretics and with fludrocortisone, if needed. Patients with bilateral hyperplasia should be treated medically. The drug regimen for these patients, as well as for patients with an adenoma who are poor surgical candidates, should include an aldosterone antagonist and, if necessary, other potassium-sparing diuretics.

Glucocorticoid-remediable hyperaldosteronism is a rare, monogenic autosomal dominant disorder characterized by moderate to

severe hypertension, often occurring at an early age. These patients may have a family history of hemorrhagic stroke at a young age. Hypokalemia is usually mild or absent. Normally, angiotensin II stimulates aldosterone production by the adrenal zona glomerulosa, whereas ACTH stimulates cortisol production in the zona fasciculata. Owing to a chimeric gene on chromosome 8, ACTH also regulates aldosterone secretion by the zona fasciculata in patients with glucocorticoid-remediable hyperaldosteronism. The consequence is overproduction in the zona fasciculata of both aldosterone and hybrid steroids (18-hydroxycortisol and 18-oxocortisol) due to oxidation of cortisol. The diagnosis may be established by urine excretion rates of these hybrid steroids that are 20 to 30 times normal or by direct genetic testing. Therapeutically, suppression of ACTH with low-dose glucocorticoids corrects the hyperaldosteronism, hypertension, and hypokalemia. Aldosterone antagonists are also therapeutic options. Patients with familial aldosteronism types II and III are treated with aldosterone antagonists or adrenalectomy.

CUSHING'S SYNDROME

(See also Chap. 406) Cushing's syndrome is related to excess cortisol production due either to excess ACTH secretion (from a pituitary tumor or an ectopic tumor) or to ACTH-independent adrenal production of cortisol. Hypertension occurs in 75–80% of patients with Cushing's syndrome. The mechanism of hypertension may be related to stimulation of mineralocorticoid receptors by cortisol and increased secretion of other adrenal steroids. If clinically suspected based on phenotypic characteristics, in patients not taking exogenous glucocorticoids, laboratory screening may be carried out with measurement of 24-h excretion rates of urine free cortisol or an overnight dexamethasone-suppression test. Late night salivary cortisol is also a sensitive and convenient screening test. Further evaluation is required to confirm the diagnosis and identify the specific etiology of Cushing's syndrome. Appropriate therapy depends on the etiology.

PHEOCHROMOCYTOMA

(See also Chap. 407) Catecholamine-secreting tumors are located in the adrenal medulla (pheochromocytoma) or in extra-adrenal paraganglion tissue (paraganglioma) and account for hypertension in ~0.05% of patients. If unrecognized, pheochromocytoma may result in lethal cardiovascular consequences. Clinical manifestations, including hypertension, are primarily related to increased circulating catecholamines, although some of these tumors may secrete a number of other vasoactive substances. In a small percentage of patients, epinephrine is the predominant catecholamine secreted by the tumor, and these patients may present with hypotension rather than hypertension. The initial suspicion of the diagnosis is based on symptoms and/or the association of pheochromocytoma with other disorders (Table 298-4). Approximately 20% of pheochromocytomas are familial with autosomal dominant inheritance. Inherited pheochromocytomas may be associated with multiple endocrine neoplasia (MEN) type 2A and type 2B, von Hippel-Lindau disease, and neurofibromatosis (Table 298-4). Each of these syndromes is related to specific, identifiable germ-line mutations. Additionally, mutations of succinate dehydrogenase genes are associated with paraganglioma syndromes, generally characterized by head and neck paragangliomas. Laboratory testing consists of measuring catecholamines in either urine or plasma, e.g., 24-h urine metanephrine excretion or fractionated plasma free metanephrines. The urine measurement is less sensitive but more specific. Genetic screening is available for evaluating patients and relatives suspected of harboring a pheochromocytoma associated with a familial syndrome. Surgical excision is the definitive treatment of pheochromocytoma and results in cure in ~90% of patients.

MISCELLANEOUS CAUSES OF HYPERTENSION

Independent of obesity, hypertension occurs in >50% of individuals with *obstructive sleep apnea*. The severity of hypertension correlates with the severity of sleep apnea. Approximately 70% of patients with obstructive sleep apnea are obese. Hypertension related to obstructive sleep apnea also should be considered in patients with drug-resistant

TABLE 298-4 RARE MENDELIAN FORMS OF HYPERTENSION

Disease	Phenotype	Genetic Cause
Glucocorticoid-remediable hyperaldosteronism	Autosomal dominant Absent or mild hypokalemia	Chimeric 11 β -hydroxylase/aldosterone gene on chromosome 8
17 α -hydroxylase deficiency	Autosomal recessive Males: pseudohermaphroditism Females: primary amenorrhea, absent secondary sexual characteristics	Random mutations of the <i>CYP17</i> gene on chromosome 10
11 β -hydroxylase deficiency	Autosomal recessive Masculinization	Mutations of the <i>CYP11B1</i> gene on chromosome 8q21-q22
11 β -hydroxysteroid dehydrogenase deficiency (apparent mineralocorticoid excess syndrome)	Autosomal recessive Hypokalemia, low renin, low aldosterone	Mutations in the 11 β -hydroxysteroid dehydrogenase gene
Liddle's syndrome	Autosomal dominant Hypokalemia, low renin, low aldosterone	Mutation subunits of the epithelial sodium channel <i>SCNN1B</i> and <i>SCNN1C</i> genes
Pseudohypoaldosteronism type II (Gordon's syndrome)	Autosomal dominant Hyperkalemia, normal glomerular filtration rate	Linkage to chromosomes 1q31-q42 and 17p11-q21
Hypertension exacerbated in pregnancy	Autosomal dominant Severe hypertension in early pregnancy	Missense mutation with substitution of leucine for serine at codon 810 (<i>MR_{1B10}</i>)
Polycystic kidney disease	Autosomal dominant Large cystic kidneys, renal failure, liver cysts, cerebral aneurysms, valvular heart disease	Mutations in the <i>PKD1</i> gene on chromosome 16 and <i>PKD2</i> gene on chromosome 4
Pheochromocytoma	Autosomal dominant (a) Multiple endocrine neoplasia, type 2A Medullary thyroid carcinoma, hyperparathyroidism (b) Multiple endocrine neoplasia, type 2B Medullary thyroid carcinoma, mucosal neuromas, thickened corneal nerves, alimentary ganglioneuromatoses, marfanoid habitus (c) von Hippel-Lindau disease Retinal angiomas, hemangioblastomas of the cerebellum and spinal cord, renal cell carcinoma (d) Neurofibromatosis type 1 Multiple neurofibromas, café-au-lait spots	(a) Mutations in the <i>RET</i> protooncogene (b) Mutations in the <i>RET</i> protooncogene (c) Mutations in the <i>VHL</i> tumor-suppressor gene (d) Mutations in the <i>NF1</i> tumor-suppressor gene

hypertension and patients with a history of snoring. The diagnosis can be confirmed by polysomnography. In obese patients, weight loss may alleviate or cure sleep apnea and related hypertension. Continuous positive airway pressure (CPAP) or bilevel positive airway pressure (BiPAP) administered during sleep is an effective therapy for obstructive sleep apnea. With CPAP or BiPAP, patients with apparently drug-resistant hypertension may be more responsive to antihypertensive agents.

Coarctation of the aorta is the most common congenital cardiovascular cause of hypertension (Chap. 282). The incidence is 1–8 per 1000 live births. It is usually sporadic but occurs in 35% of children with Turner's syndrome. Even when the anatomic lesion is surgically corrected in infancy, up to 30% of patients develop subsequent hypertension and are at risk of accelerated coronary artery disease and cerebrovascular events. Patients with less severe lesions may not be diagnosed until young adulthood. Physical findings include diminished and delayed femoral pulses and a systolic pressure gradient between the right arm and the legs and, depending on the location of the coarctation, between the right and left arms. A blowing systolic murmur may be heard in the posterior left interscapular areas. The diagnosis may be confirmed by chest x-ray and transesophageal echocardiography. Therapeutic options include surgical repair and balloon angioplasty, with or without placement of an intravascular stent. Subsequently, many patients do not have a normal life expectancy but may have persistent hypertension, with death due to ischemic heart disease, cerebral hemorrhage, or aortic aneurysm.

Several additional endocrine disorders, including *thyroid diseases* and *acromegaly*, cause hypertension. Mild diastolic hypertension may be a consequence of hypothyroidism, whereas hyperthyroidism may result in systolic hypertension. *Hypercalcemia* of any etiology, the most

common being primary hyperparathyroidism, may result in hypertension. Hypertension also may be related to a number of prescribed or over-the-counter *medications*.

MONOGENIC HYPERTENSION

In addition to glucocorticoid-remediable primary aldosteronism, a number of rare forms of monogenic hypertension have been identified (Table 298-4). These disorders may be recognized by their characteristic phenotypes, and in many instances the diagnosis may be confirmed by genetic analysis. Several inherited defects in adrenal steroid biosynthesis and metabolism result in mineralocorticoid-induced hypertension and hypokalemia. In patients with a 17 α -hydroxylase deficiency, synthesis of sex hormones and cortisol is decreased (Fig. 298-3). Consequently, these individuals do not mature sexually; males may present with pseudohermaphroditism and females with primary amenorrhea and absent secondary sexual characteristics. Because cortisol-induced negative feedback on pituitary ACTH production is diminished, ACTH-stimulated adrenal steroid synthesis proximal to the enzymatic block is increased. Hypertension and hypokalemia are consequences of increased synthesis of mineralocorticoids proximal to the enzymatic block, particularly desoxycorticosterone. Increased steroid production and, hence, hypertension may be treated with low-dose glucocorticoids. An 11 β -hydroxylase deficiency results in a salt-retaining adrenogenital syndrome that occurs in 1 in 100,000 live births. This enzymatic defect results in decreased cortisol synthesis, increased synthesis of mineralocorticoids (e.g., desoxycorticosterone), and shunting of steroid biosynthesis into the androgen pathway. In the severe form, the syndrome may present early in life, including the

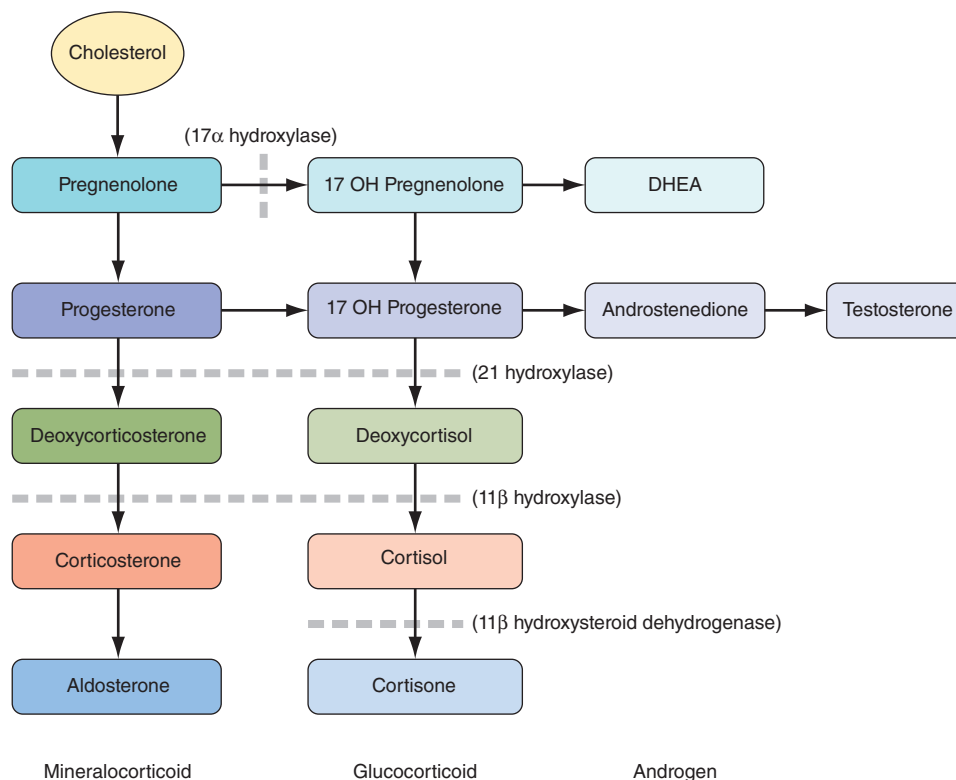


FIGURE 298-3 Adrenal enzymatic defects. DHEA, dehydroepiandrosterone.

newborn period, with virilization and ambiguous genitalia in females and penile enlargement in males, or in older children as precocious puberty and short stature. Acne, hirsutism, and menstrual irregularities may be the presenting features when the disorder is first recognized in adolescence or early adulthood. Hypertension is less common in the late-onset forms. Patients with an 11 β -hydroxysteroid dehydrogenase deficiency have an impaired capacity to metabolize cortisol to its inactive metabolite, cortisone, and hypertension is related to activation of mineralocorticoid receptors by cortisol. This defect may be inherited or acquired, due to licorice-containing glycyrrhizin acid. The same substance is present in the paste of several brands of chewing tobacco. The defect in Liddle's syndrome (Chaps. 63 and 406) results from constitutive activation of amiloride-sensitive epithelial sodium channels on the distal renal tubule, resulting in excess sodium reabsorption; the syndrome is ameliorated by amiloride. Hypertension exacerbated in pregnancy (Chap. 8) may be due to activation of the mineralocorticoid receptor by progesterone.

APPROACH TO THE PATIENT: Hypertension

HISTORY

The initial assessment of a hypertensive patient should include a complete history and physical examination to confirm a diagnosis of hypertension, screen for other cardiovascular disease risk factors, screen for secondary causes of hypertension, identify cardiovascular consequences of hypertension and other comorbidities, assess blood pressure-related lifestyles, and determine the potential for intervention.

Most patients with hypertension have no specific symptoms referable to their blood pressure elevation. Although popularly considered a symptom of elevated arterial pressure, headache generally occurs only in patients with severe hypertension. Characteristically, a "hypertensive headache" occurs in the morning and is localized to the occipital region. Other nonspecific symptoms that may be related to elevated blood pressure include

dizziness, palpitations, easy fatigability, and impotence. When symptoms are present, they are generally related to hypertensive cardiovascular disease or to manifestations of secondary hypertension. Table 298-5 lists salient features that should be addressed in obtaining a history from a hypertensive patient.

MEASUREMENT OF BLOOD PRESSURE

Reliable measurements of blood pressure depend on attention to the details of the technique and conditions of the measurement. Proper training of observers, positioning of the patient, and selection of cuff size are essential. Owing to recent regulations preventing the use of mercury because of concerns about its potential toxicity, most office measurements are made with aneroid sphygmomanometers or with oscillometric devices. These instruments should be calibrated periodically, and their accuracy confirmed. Before the blood pressure measurement is taken, the individual should be seated quietly in a chair (not the exam table) with feet on the floor for 5 min in a private, quiet setting with a comfortable room temperature. At least two measurements should be made. The center of the cuff should be at heart level, and the width of the bladder cuff

TABLE 298-5 PATIENT'S RELEVANT HISTORY

Duration of hypertension
Previous therapies: responses and side effects
Family history of hypertension and cardiovascular disease
Dietary and psychosocial history
Other risk factors: weight change, dyslipidemia, smoking, diabetes, physical inactivity
Evidence of secondary hypertension: history of renal disease; change in appearance; muscle weakness; spells of sweating, palpitations, tremor; erratic sleep, snoring, daytime somnolence; symptoms of hypo- or hyperthyroidism; use of agents that may increase blood pressure
Evidence of target organ damage: history of TIA, stroke, transient blindness; angina, myocardial infarction, congestive heart failure; sexual function
Other comorbidities

Abbreviation: TIA, transient ischemic attack.

should equal at least 40% of the arm circumference; the length of the cuff bladder should encircle at least 80% of the arm circumference. It is important to pay attention to cuff placement, stethoscope placement, and the rate of deflation of the cuff (2 mmHg/s). Systolic blood pressure is the first of at least two regular “tapping” Korotkoff sounds, and diastolic blood pressure is the point at which the last regular Korotkoff sound is heard. In current practice, a diagnosis of hypertension generally is based on seated, office measurements.

Currently available ambulatory monitors are fully automated, use the oscillometric technique, and typically are programmed to take readings every 15–30 min. Twenty-four-hour ambulatory blood pressure monitoring more reliably predicts cardiovascular disease risk than do office measurements. However, ambulatory monitoring is not used routinely in clinical practice and generally is reserved for patients in whom white coat hypertension is suspected. The Seventh Report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC 7) has also recommended ambulatory monitoring for treatment resistance, symptomatic hypotension, autonomic failure, and episodic hypertension.

PHYSICAL EXAMINATION

Body habitus, including weight and height, should be noted. At the initial examination, blood pressure should be measured in both arms and preferably in the supine, sitting, and standing positions to evaluate for postural hypotension. Even if the femoral pulse is normal to palpation, arterial pressure should be measured at least once in the lower extremity in patients in whom hypertension is discovered before age 30. Heart rate also should be recorded. Hypertensive individuals have an increased prevalence of atrial fibrillation. The neck should be palpated for an enlarged thyroid gland, and patients should be assessed for signs of hypo- and hyperthyroidism. Examination of blood vessels may provide clues about underlying vascular disease and should include fundoscopic examination, auscultation for bruits over the carotid and femoral arteries, and palpation of femoral and pedal pulses. The retina is the only tissue in which arteries and arterioles can be examined directly. With increasing severity of hypertension and atherosclerotic disease, progressive fundoscopic changes include increased arteriolar light reflex, arteriovenous crossing defects, hemorrhages and exudates, and, in patients with malignant hypertension, papilledema. Examination of the heart may reveal a loud second heart sound due to closure of the aortic valve and an S_4 gallop attributed to atrial contraction against a noncompliant left ventricle. Left ventricular hypertrophy may be detected by an enlarged, sustained, and laterally displaced apical impulse. An abdominal bruit, particularly a bruit that lateralizes and extends throughout systole into diastole, raises the possibility of renovascular hypertension. Kidneys of patients with polycystic kidney disease may be palpable in the abdomen. The physical examination also should include evaluation for signs of CHF and a neurologic examination.

LABORATORY TESTING

Table 298-6 lists recommended laboratory tests in the initial evaluation of hypertensive patients. Repeat measurements of renal function, serum electrolytes, fasting glucose, and lipids may be obtained

TABLE 298-6 BASIC LABORATORY TESTS FOR INITIAL EVALUATION

System	Test
Renal	Microscopic urinalysis, albumin excretion, serum BUN and/or creatinine
Endocrine	Serum sodium, potassium, calcium, ?TSH
Metabolic	Fasting blood glucose, total cholesterol, HDL and LDL (often computed) cholesterol, triglycerides
Other	Hematocrit, electrocardiogram

Abbreviations: BUN, blood urea nitrogen; HDL, high-density lipoprotein; LDL, low-density lipoprotein; TSH, thyroid-stimulating hormone.

after the introduction of a new antihypertensive agent and then annually or more frequently if clinically indicated. More extensive laboratory testing is appropriate for patients with apparent drug-resistant hypertension or when the clinical evaluation suggests a secondary form of hypertension.

TREATMENT HYPERTENSION

LIFESTYLE INTERVENTIONS

Implementation of lifestyles that favorably affect blood pressure has implications for both the prevention and the treatment of hypertension. Health-promoting lifestyle modifications are recommended for individuals with prehypertension and as an adjunct to drug therapy in hypertensive individuals. These interventions should address overall cardiovascular disease risk. Although the impact of lifestyle interventions on blood pressure is more pronounced in persons with hypertension, in short-term trials, weight loss and reduction of dietary NaCl have been shown to prevent the development of hypertension. In hypertensive individuals, even if these interventions do not produce a sufficient reduction in blood pressure to avoid drug therapy, the number of medications or doses required for blood pressure control may be reduced. Dietary modifications that effectively lower blood pressure are weight loss, reduced NaCl intake, increased potassium intake, moderation of alcohol consumption, and an overall healthy dietary pattern (Table 298-7).

Prevention and treatment of obesity are important for reducing blood pressure and cardiovascular disease risk. In short-term trials, even modest weight loss can lead to a reduction of blood pressure and an increase in insulin sensitivity. Average blood pressure reductions of 6.3/3.1 mmHg have been observed with a reduction in mean body weight of 9.2 kg. Regular physical activity facilitates weight loss, decreases blood pressure, and reduces the overall risk of cardiovascular disease. Blood pressure may be lowered by 30 min of moderately intense physical activity, such as brisk walking, 6–7 days a week, or by more intense, less frequent workouts.

There is individual variability in the sensitivity of blood pressure to NaCl, and this variability may have a genetic basis. Based on results of meta-analyses, lowering of blood pressure by limiting daily NaCl intake to 4.4–7.4 g (75–125 meq) results in blood pressure reductions of 3.7–4.9/0.9–2.9 mmHg in hypertensive individuals and lesser reductions in normotensive individuals. Several long-term, prospective, randomized clinical trials have reported that a reduced salt intake results in a decreased incidence of cardiovascular events. Although reduced salt intakes are generally recommended for both the prevention and treatment of hypertension, overly rigorous salt restriction may have adverse cardiovascular outcomes in diabetic patients and in patients with CHF aggressively treated with diuretics. Potassium and calcium supplementation have inconsistent, modest antihypertensive effects, and, independent of blood pressure, potassium supplementation may be associated with reduced stroke mortality. Consuming three or more alcoholic drinks per day (a standard drink contains ~14 g ethanol) is associated with higher blood pressures, and a reduction of alcohol consumption is associated with a

TABLE 298-7 LIFESTYLE MODIFICATIONS TO MANAGE HYPERTENSION

Weight reduction	Attain and maintain BMI <25 kg/m ²
Dietary salt reduction	<6 g NaCl/d
Adapt DASH-type dietary plan	Diet rich in fruits, vegetables, and low-fat dairy products with reduced content of saturated and total fat
Moderation of alcohol consumption	For those who drink alcohol, consume ≤2 drinks/day in men and ≤1 drink/day in women
Physical activity	Regular aerobic activity, e.g., brisk walking for 30 min/d

Abbreviations: BMI, body mass index; DASH, Dietary Approaches to Stop Hypertension (trial).

reduction of blood pressure. In patients with advanced renal disease, dietary protein restriction may have a modest effect in mitigating renal damage by reducing the intrarenal transmission of systemic arterial pressure.

The DASH (Dietary Approaches to Stop Hypertension) trial convincingly demonstrated that over an 8-week period a diet high in fruits, vegetables, and low-fat dairy products lowers blood pressure in individuals with high-normal blood pressures or mild hypertension. Reduction of daily NaCl intake to <6 g (100 meq) augmented the effect of this diet on blood pressure. Fruits and vegetables are enriched sources of potassium, magnesium, and fiber, and dairy products are an important source of calcium.

PHARMACOLOGIC THERAPY

Drug therapy is recommended for individuals with blood pressures $\geq 140/90$ mmHg. The degree of benefit derived from antihypertensive agents is related to the magnitude of the blood pressure reduction. Lowering systolic blood pressure by 10–12 mmHg and diastolic blood pressure by 5–6 mmHg confers relative risk reductions of 35–40% for stroke and 12–16% for CHD within 5 years of the initiation of treatment. Risk of heart failure is reduced by >50%. Hypertension control is the single most effective intervention for slowing the rate of progression of hypertension-related kidney disease.

There is considerable variation in individual responses to different classes of antihypertensive agents, and the magnitude of response to any single agent may be limited by activation of counter-regulatory mechanisms. Most available agents reduce systolic blood pressure by 7–13 mmHg and diastolic blood pressure by 4–8 mmHg when corrected for placebo effect. More often than not, combinations of agents, with complementary antihypertensive mechanisms, are required to achieve goal blood pressure reductions. Selection of antihypertensive agents and combinations of agents should be individualized, taking into account age, severity of hypertension, other cardiovascular disease risk factors, comorbid conditions, and practical considerations related to cost, side effects, and frequency of dosing (**Table 298-8**).

Diuretics Low-dose thiazide diuretics may be used alone or in combination with other antihypertensive drugs. Thiazides inhibit the Na^+/Cl^- pump in the distal convoluted tubule and hence increase sodium excretion. In the long term, they also may act as vasodilators. Thiazides are safe, efficacious, inexpensive, and reduce clinical events. They provide additive blood pressure-lowering effects when combined with beta blockers, angiotensin-converting enzyme inhibitors (ACEIs), or angiotensin receptor blockers (ARBs). In contrast, addition of a diuretic to a calcium channel blocker is less effective. Usual doses of hydrochlorothiazide range from 6.25–50 mg/d. Owing to an increased incidence of metabolic side effects (hypokalemia, insulin resistance, increased cholesterol), higher doses generally are not recommended. Chlorthalidone is a diuretic structurally similar to hydrochlorothiazide, and like hydrochlorothiazide, it blocks sodium-chloride cotransport in the early distal tubule. However, chlorthalidone has a longer half-life (40–60 h vs. 9–15 h) and an antihypertensive potency ~1.5–2.0 times that of hydrochlorothiazide. Potassium loss is also greater with chlorthalidone. Two potassium-sparing diuretics, amiloride and triamterene, act by inhibiting epithelial sodium channels in the distal nephron. These agents are weak antihypertensive agents but may be used in combination with a thiazide to protect against hypokalemia. The main pharmacologic target for loop diuretics is the $\text{Na}^+-\text{K}^+-2\text{Cl}^-$ cotransporter in the thick ascending limb of the loop of Henle. Loop diuretics generally are reserved for hypertensive patients with reduced glomerular filtration rates (reflected in serum creatinine >220 $\mu\text{mol/L}$ [>2.5 mg/dL]), CHF, or sodium retention and edema for some other reason, such as treatment with a potent vasodilator, e.g., minoxidil.

Blockers of the Renin-Angiotensin System ACEIs decrease the production of angiotensin II, increase bradykinin levels, and reduce sympathetic nervous system activity. ARBs provide selective blockade of AT_1

receptors, and the effect of angiotensin II on unblocked AT_2 receptors may augment their hypotensive effect. Both classes of agents are effective antihypertensive agents that may be used as monotherapy or in combination with diuretics, calcium antagonists, and alpha blocking agents. ACEIs and ARBs improve insulin action and ameliorate the adverse effects of diuretics on glucose metabolism. Although the overall impact on the incidence of diabetes is modest, compared with amlodipine (a calcium antagonist), valsartan (an ARB) has been shown to reduce the risk of developing diabetes in high-risk hypertensive patients. ACEI/ARB combinations are less effective in lowering blood pressure than is the case when either class of these agents is used in combination with other classes of agents. In patients with vascular disease or a high risk of diabetes, combination ACEI/ARB therapy has been associated with more adverse events (e.g., cardiovascular death, myocardial infarction, stroke, and hospitalization for heart failure) without increases in benefit.

Side effects of ACEIs and ARBs include functional renal insufficiency due to efferent renal arteriolar dilation in a kidney with a stenotic lesion of the renal artery. Additional predisposing conditions to renal insufficiency induced by these agents include dehydration, CHF, and use of nonsteroidal anti-inflammatory drugs. Dry cough occurs in ~15% of patients, and angioedema occurs in <1% of patients taking ACEIs. Angioedema occurs most commonly in individuals of Asian origin and more commonly in African Americans than in whites. Hyperkalemia due to hypoaldosteronism is an occasional side effect of both ACEIs and ARBs.

An alternative approach to blocking the renin-angiotensin system has recently been introduced into clinical practice for the treatment of hypertension: direct renin inhibitors. Blockade of the renin-angiotensin system is more complete with renin inhibitors than with ACEIs or ARBs. Aliskiren is the first of a class of oral, nonpeptide competitive inhibitors of the enzymatic activity of renin. Monotherapy with aliskiren seems to be as effective as an ACEI or ARB for lowering blood pressure, but not more effective. Further blood reductions may be achieved when aliskiren is used in combination with a thiazide diuretic or a calcium antagonist. Currently, aliskiren is not considered a first-line antihypertensive agent.

Aldosterone Antagonists Spironolactone is a nonselective aldosterone antagonist that may be used alone or in combination with a thiazide diuretic. It may be a particularly effective agent in patients with low-renin primary hypertension, resistant hypertension, and primary aldosteronism. In patients with CHF, low-dose spironolactone reduces mortality and hospitalizations for heart failure when given in addition to conventional therapy with ACEIs, digoxin, and loop diuretics. Because spironolactone binds to progesterone and androgen receptors, side effects may include gynecomastia, impotence, and menstrual abnormalities. These side effects are circumvented by a newer agent, eplerenone, which is a selective aldosterone antagonist.

Beta Blockers β -Adrenergic receptor blockers lower blood pressure by decreasing cardiac output, due to a reduction of heart rate and contractility. Other proposed mechanisms by which beta blockers lower blood pressure include a central nervous system effect and inhibition of renin release. Beta blockers are particularly effective in hypertensive patients with tachycardia, and their hypotensive potency is enhanced by coadministration with a diuretic. In lower doses, some beta blockers selectively inhibit cardiac β_1 receptors and have less influence on β_2 receptors on bronchial and vascular smooth muscle cells; however, there seems to be no difference in the antihypertensive potencies of cardioselective and nonselective beta blockers. Some beta blockers have intrinsic sympathomimetic activity, although it is uncertain whether this constitutes an overall advantage or disadvantage in cardiac therapy. Beta blockers without intrinsic sympathomimetic activity decrease the rate of sudden death, overall mortality, and recurrent myocardial infarction. In patients with CHF, beta blockers have been shown to reduce the risks of hospitalization and mortality. Overall, beta blockers

TABLE 298-8 EXAMPLES OF ORAL DRUGS USED IN TREATMENT OF HYPERTENSION

Drug Class	Examples	Usual Total Daily Dose ^a (Dosing Frequency/Day)	Other Indications	Contraindications/Cautions
Diuretics				
Thiazides	Hydrochlorothiazide	6.25–50 mg (1–2)		Diabetes, dyslipidemia, hyperuricemia, gout, hypokalemia
	Chlorthalidone	25–50 mg (1)		
Loop diuretics	Furosemide	40–80 mg (2–3)	CHF due to systolic dysfunction, renal failure	Diabetes, dyslipidemia, hyperuricemia, gout, hypokalemia
	Ethacrynic acid	50–100 mg (2–3)		
Aldosterone antagonists	Spirolactone	25–100 mg (1–2)	CHF due to systolic dysfunction, primary aldosteronism	Renal failure, hyperkalemia
	Eplerenone	50–100 mg (1–2)		
K ⁺ retaining	Amiloride	5–10 mg (1–2)		Renal failure, hyperkalemia
	Triamterene	50–100 mg (1–2)		
Beta blockers				
Cardioselective	Atenolol	25–100 mg (1)	Angina, CHF due to systolic dysfunction, post-MI, sinus tachycardia, ventricular tachyarrhythmias	Asthma, COPD, 2nd- or 3rd-degree heart block, sick-sinus syndrome
Nonselective	Metoprolol	25–100 mg (1–2)		
	Propranolol	40–160 mg (2)		
Combined alpha/beta	Propranolol LA	60–180 (1)		
	Labetalol	200–800 mg (2)	?Post-MI, CHF	
Carvedilol	12.5–50 mg (2)			
Alpha antagonists				
Selective	Prazosin	2–20 mg (2–3)	Prostatism	
	Doxazosin	1–16 mg (1)		
	Terazosin	1–10 mg (1–2)		
Nonselective	Phenoxybenzamine	20–120 mg (2–3)	Pheochromocytoma	
Sympatholytics				
Central	Clonidine	0.1–0.6 mg (2)		
	Clonidine patch	0.1–0.3 mg (1/week)		
	Methyldopa	250–1000 mg (2)		
	Reserpine	0.05–0.25 mg (1)		
	Guanfacine	0.5–2 mg (1)		
ACE inhibitors				
	Captopril	25–200 mg (2)	Post-MI, coronary syndromes, CHF with low ejection fraction, nephropathy	Acute renal failure, bilateral renal artery stenosis, pregnancy, hyperkalemia
	Lisinopril	10–40 mg (1)		
	Ramipril	2.5–20 mg (1–2)		
Angiotensin II antagonists				
	Losartan	25–100 mg (1–2)	CHF with low ejection fraction, nephropathy, ACE inhibitor cough	Renal failure, bilateral renal artery stenosis, pregnancy, hyperkalemia
	Valsartan	80–320 mg (1)		
	Candesartan	2–32 mg (1–2)		
Renin inhibitors	Aliskiren	150–300 mg (1)	Diabetic nephropathy	Pregnancy
Calcium antagonists				
Dihydropyridines	Nifedipine (long-acting)	30–60 mg (1)		
Nondihydropyridines	Verapamil (long-acting)	120–360 mg (1–2)	Post-MI, supraventricular tachycardias, angina	2nd- or 3rd-degree heart block
	Diltiazem (long-acting)	180–420 mg (1)		
Direct vasodilators				
	Hydralazine	25–100 mg (2)		Severe coronary artery disease
	Minoxidil	2.5–80 mg (1–2)		

^aAt the initiation of therapy, lower doses may be preferable for elderly patients and for select combinations of antihypertensive agents.

Abbreviations: ACE, angiotensin-converting enzyme; CHF, congestive heart failure; COPD, chronic obstructive pulmonary disease; MI, myocardial infarction.

may be less protective against cardiovascular and cerebrovascular endpoints, and some beta blockers may have less effect on central aortic pressure than other classes of antihypertensive agents. However, beta blockers remain appropriate therapy for hypertensive patients with concomitant heart disease and related comorbidities. Carvedilol and labetalol block both β receptors and peripheral α -adrenergic receptors. The potential advantages of combined β - and α -adrenergic blockade in treating hypertension remain to be determined. Nebivolol represents another class of cardioselective beta blockers that has additional vasodilator actions related to enhancement of nitric oxide activity. Whether this confers greater clinical effectiveness remains to be determined.

α -Adrenergic Blockers Postsynaptic, selective α -adrenoreceptor antagonists lower blood pressure by decreasing peripheral vascular resistance. They are effective antihypertensive agents used either as monotherapy or in combination with other agents. However, in clinical trials of hypertensive patients, alpha blockade has not been shown to reduce cardiovascular morbidity and mortality or to provide as much protection against CHF as other classes of antihypertensive agents. These agents are also effective in treating lower urinary tract symptoms in men with prostatic hypertrophy. Nonselective α -adrenoreceptor antagonists bind to postsynaptic and presynaptic receptors and are used primarily for the management of patients with pheochromocytoma.

Sympatholytic Agents Centrally acting α_2 sympathetic agonists decrease peripheral resistance by inhibiting sympathetic outflow. They may be particularly useful in patients with autonomic neuropathy who have wide variations in blood pressure due to baroreceptor denervation. Drawbacks include somnolence, dry mouth, and rebound hypertension on withdrawal. Peripheral sympatholytics decrease peripheral resistance and venous constriction by depleting nerve terminal norepinephrine. Although they are potentially effective antihypertensive agents, their usefulness is limited by orthostatic hypotension, sexual dysfunction, and numerous drug-drug interactions. Rebound hypertension is another concern with abrupt cessation of drugs with a short half-life.

Calcium Channel Blockers Calcium antagonists reduce vascular resistance through L-channel blockade, which reduces intracellular calcium and blunts vasoconstriction. This is a heterogeneous group of agents that includes drugs in the following three classes: phenylalkylamines (verapamil), benzothiazepines (diltiazem), and 1,4-dihydropyridines (nifedipine-like). Used alone and in combination with other agents (ACEIs, beta blockers, α_1 -adrenergic blockers), calcium antagonists effectively lower blood pressure; however, it is unclear if adding a diuretic to a calcium blocker results in a further lowering of blood pressure. Side effects of flushing, headache, and edema with dihydropyridine use are related to their potencies as arteriolar dilators; edema is due to an increase in transcapillary pressure gradients, not to net salt and water retention.

Direct Vasodilators Direct vasodilators decrease peripheral resistance and concomitantly activate mechanisms that defend arterial pressure, notably the sympathetic nervous system, the renin-angiotensin-aldosterone system, and sodium retention. Usually, they are not considered first-line agents but are most effective when added to a combination that includes a diuretic and a beta blocker. Hydralazine is a potent direct vasodilator that has antioxidant and nitric oxide-enhancing actions, and minoxidil is a particularly potent agent and is used most frequently in patients with renal insufficiency who are refractory to all other drugs. Hydralazine may induce a lupus-like syndrome, and side effects of minoxidil include hypertrichosis and pericardial effusion. Intravenous nitroprusside can be used to treat malignant hypertension and life-threatening left ventricular heart failure associated with elevated arterial pressure.

COMPARISONS OF ANTIHYPERTENSIVES

Based on pooling results from clinical trials, meta-analyses of the efficacy of different classes of antihypertensive agents suggest essentially equivalent blood pressure-lowering effects of the following six major classes of antihypertensive agents when used as monotherapy: thiazide diuretics, beta blockers, ACEIs, ARBs, calcium antagonists, and α_1 blockers. On average, standard doses of most antihypertensive agents reduce blood pressure by 8–10/4–7 mmHg; however, there may be subgroup differences in responsiveness. Younger patients may be more responsive to beta blockers and ACEIs, whereas patients over age 50 may be more responsive to diuretics and calcium antagonists. There is a limited relationship between plasma renin and blood pressure response. Patients with high-renin hypertension may be more responsive to ACEIs and ARBs than to other classes of agents, whereas patients with low-renin hypertension are more responsive to diuretics and calcium antagonists. Hypertensive African Americans tend to have low renin and may require higher doses of ACEIs and ARBs than whites for optimal blood pressure control, although this difference is abolished when these agents are combined with a diuretic. Beta blockers also appear to be less effective than thiazide diuretics in African Americans than in non-African Americans. Early pharmacogenetic studies, utilizing either a candidate gene approach or genome-wide scans, have shown associations of gene polymorphisms with blood pressure responsiveness to specific antihypertensive drugs. However, the reported effects have generally been too small to affect clinical decisions, and associated polymorphisms remain to be confirmed.

Currently, in practical terms, the presence of comorbidities often influences the selection of antihypertensive agents.

A meta-analysis of more than 30 randomized trials of blood pressure-lowering therapy indicates that for a given reduction in blood pressure, the major drug classes seem to produce similar overall net effects on total cardiovascular events. In both nondiabetic and diabetic hypertensive patients, most trials have failed to show significant differences in cardiovascular outcomes with different drug regimens as long as equivalent decreases in blood pressure were achieved. For example, the Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT) demonstrated that the occurrence of CHD and nonfatal myocardial infarction, as well as overall mortality, was virtually identical in hypertensive patients treated with either an ACEI (lisinopril), a diuretic (chlorthalidone), or a calcium antagonist (amlodipine).

However, in specific patient groups, ACEIs may have particular advantages, beyond that of blood pressure control, in reducing cardiovascular and renal outcomes. ACEIs and ARBs decrease intraglomerular pressure and proteinuria and may retard the rate of progression of renal insufficiency, not totally accounted for by their hypotensive effects, in both diabetic and nondiabetic renal diseases. In patients with type 2 diabetes, treatment with an ACEI, an ARB, or aliskiren decreases proteinuria and delays the progression of renal disease. In experimental models of hypertension and diabetes, renal protection with aliskiren is comparable to that with ACEIs and ARBs. However, in patients with type 2 diabetes, addition of aliskiren to an ACEI provides no additional protection against cardiovascular or renal disease and may be associated with more adverse outcomes. Among African Americans with hypertension-related renal disease, ACEIs appear to be more effective than beta blockers or dihydropyridine calcium channel blockers in slowing, although not preventing, the decline of glomerular filtration rate. The renoprotective effect of these renin-angiotensin blockers, compared with other antihypertensive drugs, is less obvious at lower blood pressures. In most patients with hypertension and heart failure due to systolic and/or diastolic dysfunction, the use of diuretics, ACEIs or ARBs, and beta blockers is recommended to improve survival. Independent of blood pressure, in both hypertensive and normotensive individuals, ACEIs attenuate the development of left ventricular hypertrophy, improve symptomatology and risk of death from CHF, and reduce morbidity and mortality rates in post-myocardial infarction patients. Similar benefits in cardiovascular morbidity and mortality rates in patients with CHF have been observed with the use of ARBs. ACEIs provide better coronary protection than do calcium channel blockers, whereas calcium channel blockers provide more stroke protection than do either ACEIs or beta blockers. Results of a large, double-blind, prospective clinical trial (Avoiding Cardiovascular Events through Combination Therapy in Patients Living with Systolic Hypertension [ACCOMPLISH Trial]) indicated that combination treatment with an ACEI (benazepril) plus a calcium antagonist (amlodipine) was superior to treatment with the ACEI plus a diuretic (hydrochlorothiazide) in reducing the risk of cardiovascular events and death among high-risk patients with hypertension. However, the combination of an ACEI and a diuretic has recently been shown to produce major reductions in morbidity and mortality in the very elderly.

After a stroke, combination therapy with an ACEI and a diuretic, but not with an ARB, has been reported to reduce the rate of recurrent stroke. Some of these apparent differences may reflect differences in trial design and/or patient groups.

There is a recent resurgence of interest in two nonpharmacologic, antihypertensive therapies that interrupt sympathetic outflow: (1) device-based carotid baroreflex activation by electrical stimulation of the carotid sinus; and (2) endovascular radiofrequency ablation of the renal sympathetic nerves. Whereas renal denervation is a minimally invasive procedure, carotid baroreceptor stimulation is a surgical procedure, usually performed under general anesthesia, that currently involves implanting electrodes on both the right and left carotid arteries. Both interventions inhibit sympathetic drive and

1626 decrease blood pressure by increasing the capacity of the kidney to excrete sodium and by decreasing renin release. Sustained activation of the baroreflex most likely lowers blood pressure by other mechanisms as well. Clinical experience with these interventions is limited. In the short term, blood pressure is lowered in 75–80% of patients, and the magnitude of the blood pressure reduction is similar for both procedures. To date, the most impressive results have been observed in patients with “resistant” hypertension and patients with obesity-related hypertension. Awaiting the results of long-term, multicenter clinical trials to evaluate their efficacy and safety, it remains to be seen whether these interventions will be adopted into clinical practice.

BLOOD PRESSURE GOALS OF ANTIHYPERTENSIVE THERAPY

Based on clinical trial data, the maximum protection against combined cardiovascular endpoints is achieved with pressures <135–140 mmHg for systolic blood pressure and <80–85 mmHg for diastolic blood pressure; however, treatment has not reduced cardiovascular disease risk to the level in nonhypertensive individuals. In diabetic patients, effective blood pressure control reduces the risk of cardiovascular events and death as well as the risk for microvascular disease (nephropathy, retinopathy). Although guidelines for hypertension control have recommended more aggressive blood pressure targets (e.g., office or clinic blood pressure <130/80 mmHg) for patients with diabetes, CHD, chronic kidney disease, or additional cardiovascular disease risk factors, recent evidence suggests that overly aggressive targets for blood pressure control may not be advantageous, particularly in high-risk patients. For example, among hypertensive patients with diabetes and coronary heart disease, “tight control” of systolic blood pressure (<130 mmHg) is not associated with improved cardiovascular outcomes. The concept of a “J-curve” suggests that the risk of cardiovascular events increases at blood pressures that are either too high or too low. Theoretically blood pressures that are too low may exceed the autoregulatory capacity of cerebral, coronary, and renal blood flows. There is some suggestive evidence from recent randomized clinical trials for a J-shaped relationship between blood pressure and cardiovascular outcomes (including all-cause mortality) in high-risk patients. Consequently, caution should be exercised in lowering blood pressure <130/80 mmHg in patients with diabetes, CHD, and other high-risk patients. In patients with chronic renal insufficiency, a small, nonprogressive increase in the serum creatinine concentration may occur. This generally reflects a hemodynamic response, not structural renal injury, indicating that intraglomerular pressure has been reduced. Blood pressure control should not be allowed to deteriorate in order to prevent the modest creatinine rise. Among older patients with isolated systolic hypertension, further lowering of diastolic blood pressure does not result in harm. However, relatively little information is available concerning the risk-versus-benefit ratio of antihypertensive therapy in individuals >80 years of age, and in this population, gradual blood pressure reduction to a less aggressive target level of control may be appropriate.

To achieve recommended blood pressure goals, the majority of individuals with hypertension will require treatment with more than one drug. Three or more drugs frequently are needed in patients with diabetes and renal insufficiency. For most agents, reduction of blood pressure at half-standard doses is only ~20% less than at standard doses. Appropriate combinations of agents at these lower doses may have additive or almost additive effects on blood pressure with a lower incidence of side effects.

The term *resistant hypertension* refers to patients with blood pressures persistently >140/90 mmHg despite taking three or more antihypertensive agents, including a diuretic. Resistant or difficult-to-control hypertension is more common in patients >60 years than in younger patients. Resistant hypertension may be related to “pseudoresistance” (high office blood pressures and lower home blood pressures), nonadherence to therapy, identifiable causes of hypertension (including obesity and excessive alcohol intake), and the use of any of a number of nonprescription and prescription

drugs (Table 298-3). Rarely, in older patients, pseudohypertension may be related to the inability to measure blood pressure accurately in severely sclerotic arteries. This condition is suggested if the radial pulse remains palpable despite occlusion of the brachial artery by the cuff (Osler maneuver). The actual blood pressure can be determined by direct intra-arterial measurement. Evaluation of patients with resistant hypertension might include home blood pressure monitoring to determine if office blood pressures are representative of the usual blood pressure. A more extensive evaluation for a secondary form of hypertension should be undertaken if no other explanation for hypertension resistance becomes apparent.

HYPERTENSIVE EMERGENCIES

Probably due to the widespread availability of antihypertensive therapy, in the United States there has been a decline in the numbers of patients presenting with “crisis levels” of blood pressure. Most patients who present with severe hypertension are chronically hypertensive, and in the absence of acute end organ damage, precipitous lowering of blood pressure may result in significant morbidity and should be avoided. The key to successful management of severe hypertension is to differentiate hypertensive crises from hypertensive urgencies. The degree of target organ damage, rather than the level of blood pressure alone, determines the rapidity with which blood pressure should be lowered. **Tables 298-9 and 298-10** list a number of hypertension-related emergencies and recommended therapies.

Malignant hypertension is a syndrome associated with an abrupt increase of blood pressure in a patient with underlying hypertension or related to the sudden onset of hypertension in a previously normotensive individual. The absolute level of blood pressure is not as important as its rate of rise. Pathologically, the syndrome is associated with diffuse necrotizing vasculitis, arteriolar thrombi, and fibrin deposition in arteriolar walls. Fibrinoid necrosis has been observed in arterioles of kidney, brain, retina, and other organs. Clinically, the syndrome is recognized by progressive retinopathy (arteriolar spasm, hemorrhages, exudates, and papilledema), deteriorating renal function with proteinuria, microangiopathic hemolytic anemia, and encephalopathy. Historic inquiry should include questions about the use of monoamine oxidase inhibitors and recreational drugs (e.g., cocaine, amphetamines).

Although blood pressure should be lowered rapidly in patients with hypertensive encephalopathy, there are inherent risks of overly aggressive therapy. In hypertensive individuals, the upper and lower limits of autoregulation of cerebral blood flow are shifted to higher levels of arterial pressure, and rapid lowering of blood pressure to below the lower limit of autoregulation may precipitate cerebral ischemia or infarction as a consequence of decreased cerebral blood flow. Renal and coronary blood flows also may decrease with overly aggressive acute therapy. The initial goal of therapy is to reduce

TABLE 298-9 PREFERRED PARENTERAL DRUGS FOR SELECTED HYPERTENSIVE EMERGENCIES

Hypertensive encephalopathy	Nitroprusside, nicardipine, labetalol
Malignant hypertension (when IV therapy is indicated)	Labetalol, nicardipine, nitroprusside, enalaprilat
Stroke	Nicardipine, labetalol, nitroprusside
Myocardial infarction/unstable angina	Nitroglycerin, nicardipine, labetalol, esmolol
Acute left ventricular failure	Nitroglycerin, enalaprilat, loop diuretics
Aortic dissection	Nitroprusside, esmolol, labetalol
Adrenergic crisis	Phentolamine, nitroprusside
Postoperative hypertension	Nitroglycerin, nitroprusside, labetalol, nicardipine
Preeclampsia/eclampsia of pregnancy	Hydralazine, labetalol, nicardipine

Source: Adapted from DG Vidt, in S Oparil, MA Weber (eds): *Hypertension*, 2nd ed. Philadelphia, Elsevier Saunders, 2005.

TABLE 298-10 USUAL INTRAVENOUS DOSES OF ANTIHYPERTENSIVE AGENTS USED IN HYPERTENSIVE EMERGENCIES^a

Antihypertensive Agent	Intravenous Dose
Nitroprusside	Initial 0.3 (μg/kg)/min; usual 2–4 (μg/kg)/min; maximum 10 (μg/kg)/min for 10 min
Nicardipine	Initial 5 mg/h; titrate by 2.5 mg/h at 5–15 min intervals; max 15 mg/h
Labetalol	2 mg/min up to 300 mg or 20 mg over 2 min, then 40–80 mg at 10-min intervals up to 300 mg total
Enalaprilat	Usual 0.625–1.25 mg over 5 min every 6–8 h; maximum 5 mg/dose
Esmolol	Initial 80–500 μg/kg over 1 min, then 50–300 (μg/kg)/min
Phentolamine	5–15 mg bolus
Nitroglycerin	Initial 5 μg/min, then titrate by 5 μg/min at 3–5-min intervals; if no response is seen at 20 μg/min, incremental increases of 10–20 μg/min may be used
Hydralazine	10–50 mg at 30-min intervals

^aConstant blood pressure monitoring is required. Start with the lowest dose. Subsequent doses and intervals of administration should be adjusted according to the blood pressure response and duration of action of the specific agent.

mean arterial blood pressure by no more than 25% within minutes to 2 h or to a blood pressure in the range of 160/100–110 mmHg. This may be accomplished with IV nitroprusside, a short-acting vasodilator with a rapid onset of action that allows for minute-to-minute control of blood pressure. Parenteral labetalol and nicardipine are also effective agents for the treatment of hypertensive encephalopathy.

In patients with malignant hypertension without encephalopathy or another catastrophic event, it is preferable to reduce blood pressure over hours or longer rather than minutes. This goal may effectively be achieved initially with frequent dosing of short-acting oral agents such as captopril, clonidine, and labetalol.

Acute, transient blood pressure elevations that last days to weeks frequently occur after thrombotic and hemorrhagic strokes. Autoregulation of cerebral blood flow is impaired in ischemic cerebral tissue, and higher arterial pressures may be required to maintain cerebral blood flow. Although specific blood pressure targets have not been defined for patients with acute cerebrovascular events, aggressive reductions of blood pressure are to be avoided. With the increasing availability of improved methods for measuring cerebral blood flow (using CT technology), studies are in progress to evaluate the effects of different classes of antihypertensive agents on both blood pressure and cerebral blood flow after an acute stroke. Currently, in the absence of other indications for acute therapy, for patients with cerebral infarction who are not candidates for thrombolytic therapy, one recommended guideline is to institute antihypertensive therapy only for patients with a systolic blood pressure >220 mmHg or a diastolic blood pressure >130 mmHg. If thrombolytic therapy is to be used, the recommended goal blood pressure is <185 mmHg systolic pressure and <110 mmHg diastolic pressure. In patients with hemorrhagic stroke, suggested guidelines for initiating antihypertensive therapy are systolic >180 mmHg or diastolic pressure >130 mmHg. The management of hypertension after subarachnoid hemorrhage is controversial. Cautious reduction of blood pressure is indicated if mean arterial pressure is >130 mmHg.

In addition to pheochromocytoma, an adrenergic crisis due to catecholamine excess may be related to cocaine or amphetamine overdose, clonidine withdrawal, acute spinal cord injuries, and an interaction of tyramine-containing compounds with monoamine oxidase inhibitors. These patients may be treated with phentolamine or nitroprusside.

Treatment of hypertension in patients with acute aortic dissection is discussed in Chap. 301, and treatment of hypertension in pregnancy is discussed in Chap. 8.

299 Renovascular Disease

Stephen C. Textor

The renal vasculature is unusually complex with rich arteriolar flow to the cortex in excess of metabolic requirements, consistent with its primary function as a filtering organ. After delivering blood to cortical glomeruli, the postglomerular circulation supplies deeper medullary segments that support energy-dependent solute transport at multiple levels of the renal tubule. These postglomerular vessels carry less blood, and high oxygen consumption leaves the deeper medullary regions at the margin of hypoxemia. Vascular disorders that commonly threaten the blood supply of the kidney include large-vessel atherosclerosis, fibromuscular diseases, and embolic disorders. **Microvascular injury, including inflammatory and primary hematologic disorders, is described in Chap. 341.**

The glomerular capillary endothelium shares susceptibility to oxidative stress, pressure injury, and inflammation with other vascular territories. Rates of urinary albumin excretion (UAE) are predictive of systemic atherosclerotic disease events. Increased UAE may develop years before cardiovascular events. UAE and the risk of cardiovascular events are both reduced with pharmacologic therapy such as statins. Experimental studies demonstrate functional changes and rarefaction of renal microvessels under conditions of accelerated atherosclerosis and/or compromise of proximal perfusion pressures with large-vessel disease (Fig. 299-1).

MACROVASCULAR DISEASE

Large-vessel renal artery occlusive disease can result from extrinsic compression of the vessel, fibromuscular dysplasia, or, most commonly, atherosclerotic disease. Any disorder that reduces perfusion pressure to the kidney can activate mechanisms that tend to restore renal pressures at the expense of developing systemic hypertension. Because restoration of perfusion pressures can reverse these pathways, renal artery stenosis is considered a specifically treatable “secondary” cause of hypertension.

Renal artery stenosis is common and often has only minor hemodynamic effects. Fibromuscular dysplasia (FMD) is reported in 3–5% of normal subjects presenting as potential kidney donors without hypertension. It may present clinically with hypertension in younger individuals (between age 15 and 50), most often women. FMD does not often threaten kidney function, but sometimes produces total occlusion and can be associated with renal artery aneurysms. Atherosclerotic renal artery stenosis (ARAS) is common in the general population (6.8% of a community-based sample above age 65), and the prevalence increases with age and for patients with other vascular conditions such as coronary artery disease (18–23%) and/or peripheral aortic or lower extremity disease (>30%). If untreated, ARAS progresses in nearly 50% of cases over a 5-year period, sometimes to total occlusion. Intensive treatment of arterial blood pressure and statin therapy appear to slow these rates and improve clinical outcomes.

Critical levels of stenosis lead to a reduction in perfusion pressure that activates the renin-angiotensin system, reduces sodium excretion, and activates sympathetic adrenergic pathways. These events lead to systemic hypertension characterized by angiotensin dependence in the early stages, widely varying pressures, loss of circadian blood pressure (BP) rhythms, and accelerated target organ injury, including left ventricular hypertrophy and renal fibrosis. Renovascular hypertension can be treated with agents that block the renin-angiotensin system and other drugs that modify these pressor pathways. It can also be treated with restoration of renal blood flow by either endovascular or surgical revascularization. Most patients require continued antihypertensive drug therapy because revascularization alone rarely lowers BP to normal.

ARAS and systemic hypertension tend to affect both the post-stenotic and contralateral kidneys, reducing overall glomerular filtration rate (GFR) in ARAS. When kidney function is threatened by

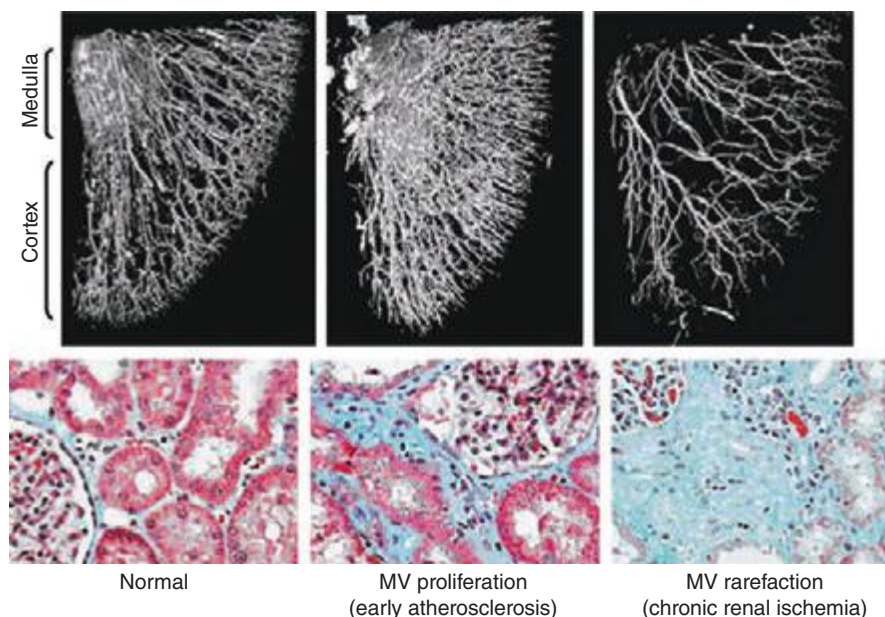


FIGURE 299-1 Examples of micro-CT images from vessels defined by radiopaque casts injected into the renal vasculature. These illustrate the complex, dense cortical capillary network supplying the kidney cortex that can either proliferate or succumb to rarefaction under the influence of atherosclerosis and/or occlusive disease. Changes in blood supply are followed by tubulointerstitial fibrosis and loss of kidney function. MV, microvascular. (From LO Lerman, AR Chade: *Curr Opin Nephrol Hyper* 18:160, 2009, with permission.)

large-vessel disease primarily, it has been labeled ischemic nephropathy. Moderately reduced blood flow that develops gradually is associated with reduced GFR and limited oxygen consumption with preserved tissue oxygenation. Hence, kidney function can remain stable during medical therapy, sometimes for years. With more advanced disease, reductions in cortical perfusion and frank tissue hypoxia develop. Unlike FMD, ARAS develops in patients with other risk factors for atherosclerosis and is commonly superimposed upon preexisting small-vessel disease in the kidney resulting from hypertension, aging, and diabetes. Nearly 85% of patients considered for renal revascularization have stage 3–5 chronic kidney disease (CKD) with GFR below 60 mL/min per 1.73 m². The presence of ARAS is a strong predictor of morbidity- and mortality-related cardiovascular events, independent of whether renal revascularization is undertaken.

Diagnostic approaches to renal artery stenosis depend partly on the specific issues to be addressed. Noninvasive characterization of the renal vasculature may be achieved by several techniques, summarized in [Table 299-1](#). Although activation of the renin-angiotensin system is a key step in developing renovascular hypertension, it is transient. Levels of renin activity are therefore subject to timing, the effects of

drugs, and sodium intake, and do not reliably predict the response to vascular therapy. Renal artery velocities by Doppler ultrasound above 200 cm/s generally predict hemodynamically important lesions (above 60% vessel lumen occlusion), although treatment trials require velocity above 300 cm/s to avoid false positives. The renal resistive index has predictive value regarding the viability of the kidney. It remains operator- and institution-dependent, however. Captopril-enhanced renography has a strong negative predictive value when entirely normal. Magnetic resonance angiography (MRA) is now less often used, as gadolinium contrast has been associated with nephrogenic systemic fibrosis. Contrast-enhanced computed tomography (CT) with vascular reconstruction provides excellent vascular images and functional assessment, but carries a small risk of contrast toxicity.

TREATMENT RENAL ARTERY STENOSIS

While restoring renal blood flow and perfusion seems intuitively beneficial for high-grade occlusive lesions, revascularization procedures also pose hazards and expense. Patients with FMD are commonly younger females with otherwise normal vessels and a

TABLE 299-1 SUMMARY OF IMAGING MODALITIES FOR EVALUATING THE KIDNEY VASCULATURE

Perfusion Studies to Assess Differential Renal Blood Flow			
Captopril renography with technetium ^{99m} Tc mertiatide (^{99m} Tc MAG3)	Captopril-mediated fall in filtration pressure amplifies differences in renal perfusion	Normal study excludes renovascular hypertension	Multiple limitations in patients with advanced atherosclerosis or creatinine >2.0 mg/dL (177 μmol/L)
Vascular Studies to Evaluate the Renal Arteries			
Duplex ultrasonography	Shows the renal arteries and measures flow velocity as a means of assessing the severity of stenosis	Inexpensive; widely available	Heavily dependent on operator's experience; less useful than invasive angiography for the diagnosis of fibromuscular dysplasia and abnormalities in accessory renal arteries
Magnetic resonance angiography	Shows the renal arteries and perirenal aorta	Not nephrotoxic, but concerns for gadolinium toxicity exclude use in GFR <30 mL/min/1.73 m ² ; provides excellent images	Expensive; gadolinium excluded in renal failure, unable to visualize stented vessels
Computed tomographic angiography	Shows the renal arteries and perirenal aorta	Provides excellent images; stents do not cause artifacts	Expensive, moderate volume of contrast required, potentially nephrotoxic
Intraarterial angiography	Shows location and severity of vascular lesion	Considered "gold standard" for diagnosis of large-vessel disease, usually performed simultaneous with planned intervention	Expensive, associated hazard of atheroemboli, contrast toxicity, procedure-related complications, e.g., dissection

Abbreviation: GFR, glomerular filtration rate.

long life expectancy. These patients often respond well to percutaneous renal artery angioplasty. If BP can be controlled to goal levels and kidney function remains stable in patients with ARAS, it may be argued that medical therapy with follow-up for disease progression is equally effective. Prospective trials up to now have failed to identify compelling benefits for interventional procedures regarding short-term results of BP and renal function, and long-term studies regarding cardiovascular outcomes, such as stroke, congestive heart failure, myocardial infarction, and end-stage renal failure, are not yet complete. Medical therapy should include blockade of the renin-angiotensin system, attainment of goal BPs, cessation of tobacco, statins, and aspirin. Renal revascularization is now often reserved for patients failing medical therapy or developing additional complications.

Techniques of renal revascularization are improving. With experienced operators, major complications occur in about 9% of cases, including renal artery dissection, capsular perforation, hemorrhage, and occasional atheroembolic disease. Although not common, atheroembolic disease can be catastrophic and accelerate both hypertension and kidney failure, precisely the events that revascularization is intended to prevent. Although renal blood flow usually can be restored by endovascular stenting, recovery of renal function is limited to about 25% of cases, with no change in 50% and some deterioration evident in others. Patients with rapid loss of kidney function, sometimes associated with antihypertensive drug therapy, or with vascular disease affecting the entire functioning kidney mass are more likely to recover function after restoring blood flow. When hypertension is refractory to effective therapy, revascularization offers real benefits. **Table 299-2** summarizes currently accepted guidelines for considering renal revascularization.

ATHEROEMBOLIC RENAL DISEASE

Emboli to the kidneys arise most frequently as a result of cholesterol crystals breaking free of atherosclerotic vascular plaque and lodging in downstream microvessels. Most clinical atheroembolic events follow angiographic procedures, often of the coronary vessels. It has been argued that nearly all vascular interventional procedures lead to plaque fracture and release of microemboli, but clinical manifestations develop only in a fraction of these. The incidence of clinical atheroemboli has been increasing with more vascular procedures and

longer life spans. Atheroembolic renal disease is suspected in more than 3% of elderly subjects with end-stage renal disease (ESRD) and is likely underdiagnosed. It is more frequent in males with a history of diabetes, hypertension, and ischemic cardiac disease. Atheroemboli in the kidney are strongly associated with aortic aneurysmal disease and renal artery stenosis. Most clinical cases can be linked to precipitating events, such as angiography, vascular surgery, anticoagulation with heparin, thrombolytic therapy, or trauma. Clinical manifestations of this syndrome commonly develop between 1 and 14 days after an inciting event and may continue to develop for weeks thereafter. Systemic embolic disease manifestations, such as fever, abdominal pain, and weight loss, are present in less than half of patients, although cutaneous manifestations including livedo reticularis and localized toe gangrene may be more common. Worsening hypertension and deteriorating kidney function are common, sometimes reaching a malignant phase. Progressive renal failure can occur and require dialytic support. These cases often develop after a stuttering onset over many weeks and have an ominous prognosis. Mortality rate after 1 year reaches 38%, and although some may eventually recover sufficiently to no longer require dialysis, many do not.

Beyond the clinical manifestations above, laboratory findings include rising creatinine, transient eosinophilia (60–80%), elevated sedimentation rate, and hypocomplementemia (15%). Establishing this diagnosis can be difficult and is often by exclusion. Definitive diagnosis depends on kidney biopsy demonstrating microvessel occlusion with cholesterol crystals that leave a “cleft” in the vessel. Biopsies obtained from patients undergoing surgical revascularization of the kidney indicate that silent cholesterol emboli are frequently present before any further manipulation is performed.

No effective therapy is available for atheroembolic disease once it has developed. Withdrawal of anticoagulation is recommended. Late recovery of kidney function after supportive measures sometimes occurs, and statin therapy may improve outcome. The role of embolic protection devices in the renal circulation is unclear, but a few prospective trials have failed to demonstrate major benefits. These devices are limited to distal protection during the endovascular procedure and offer no protection from embolic debris after removal.

THROMBOEMBOLIC RENAL DISEASE

Thrombotic occlusion of renal vessels or branch arteries can lead to declining renal function and hypertension. It is difficult to diagnose and is often overlooked, especially in elderly patients. Thrombosis can develop as a result of local vessel abnormalities, such as local dissection, trauma, or inflammatory vasculitis. Local microdissections sometimes lead to patchy, transient areas of infarctions labeled “segmental arteriolar mediolysis.” Although hypercoagulability conditions sometimes present as renal artery thrombosis, this is rare. It can also derive from distant embolic events, e.g., the left atrium in patients with atrial fibrillation or from fat emboli originating from traumatized tissue, most commonly large bone fractures. Cardiac sources include vegetations from subacute bacterial endocarditis. Systemic emboli to the kidneys may also arise from the venous circulation if right-to-left shunting occurs, e.g., through a patent foramen ovale.

Clinical manifestations vary depending on the rapidity of onset and extent of occlusion. Acute arterial thrombosis may produce flank pain, fever, leukocytosis, nausea, and vomiting. If kidney infarction results, enzymes such as lactate dehydrogenase (LDH) rise to extreme levels. If both kidneys are affected, renal function will decline precipitously with a drop in urine output. If a single kidney is involved, renal functional changes may be minor. Hypertension related to sudden release of renin from ischemic tissue can develop rapidly, as long as some viable tissue in the “peri-infarct” border zone remains. If the infarct zone demarcates precisely, the rise in BP and renin activity may resolve. Diagnosis of renal infarction may be established by vascular imaging with MRI, CT angiography, or arteriography (**Fig. 299-2**).

MANAGEMENT OF ARTERIAL THROMBOSIS OF THE KIDNEY

Options for interventions of newly detected arterial occlusion include surgical reconstruction, anticoagulation, thrombolytic therapy,

TABLE 299-2 CLINICAL FACTORS FAVORING MEDICAL THERAPY AND REVASCUARIZATION OR SURVEILLANCE FOR RENAL ARTERY STENOSIS

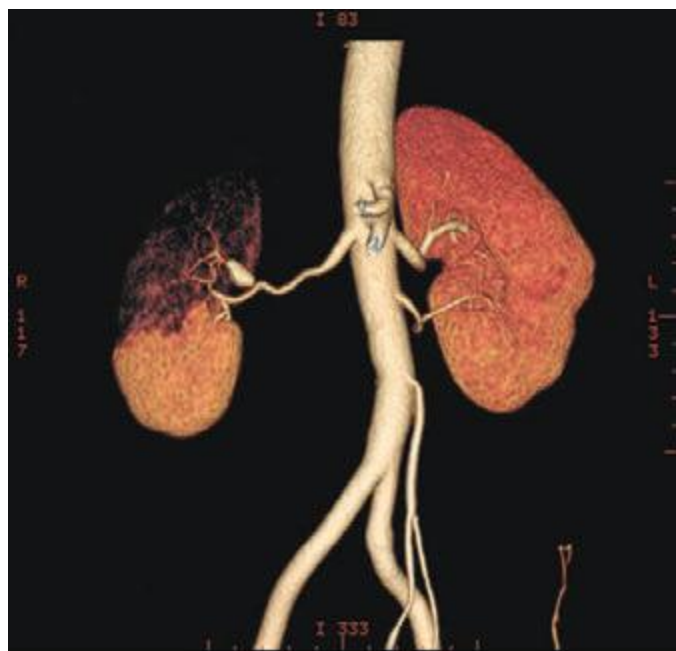
Factors Favoring Medical Therapy and Revascularization for Renal Artery Stenosis

- Progressive decline in GFR during treatment of systemic hypertension
- Failure to achieve adequate blood pressure control with optimal medical therapy (medical failure)
- Rapid or recurrent decline in the GFR in association with a reduction in systemic pressure
- Decline in the GFR during therapy with ACE inhibitors or ARBs
- Recurrent congestive heart failure in a patient in whom the adequacy of left ventricular function does not explain a cause

Factors Favoring Medical Therapy and Surveillance of Renal Artery Disease

- Controlled blood pressure with stable renal function (e.g., stable renal insufficiency)
- Stable renal artery stenosis without progression on surveillance studies (e.g., serial duplex ultrasound)
- Very advanced age and/or limited life expectancy
- Extensive comorbidity that make revascularization too risky
- High risk for or previous experience with atheroembolic disease
- Other concomitant renal parenchymal diseases that cause progressive renal dysfunction (e.g., interstitial nephritis, diabetic nephropathy)

Abbreviations: ACE, angiotensin-converting enzyme; ARBs, angiotensin receptor blockers; GFR, glomerular filtration rate.



A



B

FIGURE 299-2 **A.** CT angiogram illustrating loss of circulation to the upper pole of the right kidney in a patient with fibromuscular disease and a renal artery aneurysm. Activation of the renin-angiotensin system produced rapidly developing hypertension. **B.** Angiogram illustrating high-grade renal artery stenosis affecting the left kidney. This lesion is often part of widespread atherosclerosis and sometimes is an extension of aortic plaque. This lesion develops in older individuals with preexisting atherosclerotic risk factors.

endovascular procedures, and supportive care, particularly antihypertensive drug therapy. Application of these methods depends on the patient's overall condition, the precipitating factors (e.g., local trauma or systemic illness), the magnitude of renal tissue and function at risk, and the likelihood of recurrent events in the future. For unilateral disease, e.g., arterial dissection with thrombosis, supportive care with anticoagulation may suffice. Acute, bilateral occlusion is potentially catastrophic, producing anuric renal failure. Depending on the precipitating event, surgical or thrombolytic therapies can sometimes restore kidney viability.

MICROVASCULAR INJURY ASSOCIATED WITH HYPERTENSION

ARTERIOLONEPHROSCLEROSIS

“Malignant” Hypertension Although BP rises with age, it has long been recognized that some individuals develop rapidly progressive BP elevations with target organ injury including retinal hemorrhages, encephalopathy, and declining kidney function. Placebo arms during the controlled trials of hypertension therapy identified progression to severe levels in 20% of subjects over 5 years. If untreated, patients with target organ injury including papilledema and declining kidney function suffered mortality rates in excess of 50% over 6–12 months, hence the designation “malignant.” Postmortem studies of such patients identified vascular lesions, designated “fibrinoid necrosis,” with breakdown of the vessel wall, deposition of eosinophilic material including fibrin, and a perivascular cellular infiltrate. A separate lesion was identified in the larger interlobular arteries in many patients with hyperplastic proliferation of the vascular wall cellular elements, deposition of collagen, and separation of layers, designated the “onionskin” lesion. For many of these patients, fibrinoid necrosis led to obliteration of glomeruli and loss of tubular structures. Progressive kidney failure ensued and, without dialysis support, led to early mortality in untreated malignant-phase hypertension. These vascular changes could develop with pressure-related injury from a variety of hypertensive pathways, including but not limited to activation of the renin-angiotensin system and severe vasospasm associated with catecholamine release. Occasionally, endothelial injury is sufficient to induce microangiopathic hemolysis, as discussed below.

Antihypertensive therapy is the mainstay of therapy for malignant hypertension. With effective BP reduction, manifestations of vascular

injury including microangiopathic hemolysis and renal dysfunction can improve over time. Whereas series reported before the era of drug therapy suggested that 1-year mortality rates exceeded 90%, current survival over 5 years exceeds 50%.

Malignant hypertension is less common in Western countries, although it persists in parts of the world where medical care and antihypertensive drug therapy are less available. It most commonly develops in patients with treated hypertension who neglect to take medications or who may use vasospastic drugs, such as cocaine. Renal abnormalities typically include rising serum creatinine and occasionally hematuria and proteinuria. Biochemical findings may include evidence of hemolysis (anemia, schistocytes, and reticulocytosis) and changes associated with kidney failure. African-American males are more likely to develop rapidly progressive hypertension and kidney failure than are whites in the United States. Genetic polymorphisms (first identified as *MYH9*, but now thought to be *APOL1*) that are common in the African-American population predispose to subtle focal sclerosing glomerular disease, with severe hypertension developing at younger ages secondary to renal disease in this instance.

“Hypertensive Nephrosclerosis” Based on experience with malignant hypertension and epidemiologic evidence linking BP with long-term risks of kidney failure, it has long been assumed that lesser degrees of hypertension induce less severe, but prevalent, changes in kidney vessels and loss of kidney function. As a result, a large portion of patients reaching ESRD without a specific etiologic diagnosis are assigned the designation “hypertensive nephrosclerosis.” Pathologic examination commonly identifies afferent arteriolar thickening with deposition of homogeneous eosinophilic material (hyaline arteriosclerosis) associated with narrowing of vascular lumina. Clinical manifestations include retinal vessel changes associated with hypertension (arteriolar narrowing, crossing changes), left ventricular hypertrophy, and elevated BP. The role of these vascular changes in kidney function is unclear. Postmortem and biopsy samples from normotensive kidney donors demonstrate similar vessel changes associated with aging, dyslipidemia, and glucose intolerance. Although BP reduction does slow progression of proteinuric kidney diseases and is warranted to reduce the excessive cardiovascular risks associated with CKD, antihypertensive therapy does not alter the course of kidney dysfunction identified specifically as hypertensive nephrosclerosis.

EPIDEMIOLOGY

Venous thromboembolism (VTE) encompasses deep venous thrombosis (DVT) and pulmonary embolism (PE) and causes cardiovascular death and disability. In the United States, the Surgeon General estimates there are 100,000 to 180,000 deaths annually from PE and has declared that PE is the most common preventable cause of death among hospitalized patients. Survivors may succumb to the disabilities of chronic thromboembolic pulmonary hypertension or postthrombotic syndrome. Chronic thromboembolic pulmonary hypertension causes breathlessness, especially with exertion. Postthrombotic syndrome (also known as *chronic venous insufficiency*) damages the venous valves of the leg and causes ankle or calf swelling and leg aching, especially after prolonged standing. In its most severe form, postthrombotic syndrome causes skin ulceration (Fig. 300-1).

PATHOPHYSIOLOGY

Inflammation and Platelet Activation Virchow's triad of inflammation, hypercoagulability, and endothelial injury leads to recruitment of activated platelets, which release microparticles. These microparticles contain proinflammatory mediators that bind neutrophils, stimulating them to release their nuclear material and form web-like extracellular networks called neutrophil extracellular traps. These prothrombotic networks contain histones that stimulate platelet aggregation and promote platelet-dependent thrombin generation. Venous thrombi form and flourish in an environment of stasis, low oxygen tension, and upregulation of proinflammatory genes.

Prothrombotic States The two most common autosomal dominant genetic mutations are factor V Leiden, which causes resistance to the



FIGURE 300-1 Skin ulceration in the lateral malleolus from post-thrombotic syndrome of the leg.



FIGURE 300-2 Deep venous thrombosis at autopsy.

endogenous anticoagulant, activated protein C (which inactivates clotting factors V and VIII), and the prothrombin gene mutation, which increases the plasma prothrombin concentration (Chaps. 78 and 142). Antithrombin, protein C, and protein S are naturally occurring coagulation inhibitors. Deficiencies of these inhibitors are associated with VTE but are rare. Antiphospholipid antibody syndrome is the most common acquired cause of thrombophilia and is associated with venous or arterial thrombosis. Other common predisposing factors include cancer, obesity, cigarette smoking, systemic arterial hypertension, chronic obstructive pulmonary disease, chronic kidney disease, blood transfusion, long-haul air travel, air pollution, oral contraceptives, pregnancy, postmenopausal hormone replacement, surgery, and trauma.

Embolization When deep venous thrombi (Fig. 300-2) detach from their site of formation, they embolize to the vena cava, right atrium, and right ventricle, and lodge in the pulmonary arterial circulation, thereby causing acute PE. Paradoxically, these thrombi occasionally embolize to the arterial circulation through a patent foramen ovale or atrial septal defect. Many patients with PE have no evidence of DVT because the clot has already embolized to the lungs.

Physiology The most common gas exchange abnormalities are arterial hypoxemia and an increased alveolar-arterial O_2 tension gradient, which represents the inefficiency of O_2 transfer across the lungs. Anatomic dead space increases because breathed gas does not enter gas exchange units of the lung. Physiologic dead space increases because ventilation to gas exchange units exceeds venous blood flow through the pulmonary capillaries.

Other pathophysiologic abnormalities include:

1. *Increased pulmonary vascular resistance* due to vascular obstruction or platelet secretion of vasoconstricting neurohumoral agents such as serotonin. Release of vasoactive mediators can produce ventilation-perfusion mismatching at sites remote from the embolus, thereby accounting for discordance between a small PE and a large alveolar-arterial O_2 gradient.
2. *Impaired gas exchange* due to increased alveolar dead space from vascular obstruction, hypoxemia from alveolar hypoventilation relative to perfusion in the nonobstructed lung, right-to-left shunting, or impaired carbon monoxide transfer due to loss of gas exchange surface.
3. *Alveolar hyperventilation* due to reflex stimulation of irritant receptors.
4. *Increased airway resistance* due to constriction of airways distal to the bronchi.
5. *Decreased pulmonary compliance* due to lung edema, lung hemorrhage, or loss of surfactant.

Pulmonary Hypertension, Right Ventricular (RV) Dysfunction, and RV Microinfarction Pulmonary artery obstruction causes a rise in pulmonary

1632 artery pressure and in pulmonary vascular resistance. When RV wall tension rises, RV dilation and dysfunction ensue, with release of the cardiac biomarker, brain natriuretic peptide. The interventricular septum bulges into and compresses an intrinsically normal left ventricle (LV). Diastolic LV dysfunction reduces LV distensibility and impairs LV filling. Increased RV wall tension also compresses the right coronary artery, limits myocardial oxygen supply, and precipitates right coronary artery ischemia and RV microinfarction, with release of cardiac biomarkers such as troponin. Underfilling of the LV may lead to a fall in LV cardiac output and systemic arterial pressure, with consequent circulatory collapse and death.

CLASSIFICATION OF PULMONARY EMBOLISM AND DEEP VEIN THROMBOSIS

Pulmonary Embolism Massive PE accounts for 5–10% of cases, and is characterized by extensive thrombosis affecting at least half of the pulmonary vasculature. Dyspnea, syncope, hypotension, and cyanosis are hallmarks of massive PE. Patients with massive PE may present in cardiogenic shock and can die from multisystem organ failure. **Submassive PE** accounts for 20–25% of patients, and is characterized by RV dysfunction despite normal systemic arterial pressure. The combination of right heart failure and release of cardiac biomarkers indicates an increased likelihood of clinical deterioration. **Low-risk PE** constitutes about 70–75% of cases. These patients have an excellent prognosis.

Deep Vein Thrombosis Lower extremity DVT usually begins in the calf and propagates proximally to the popliteal vein, femoral vein, and iliac veins. Leg DVT is about 10 times more common than upper extremity DVT, which is often precipitated by placement of pacemakers, internal cardiac defibrillators, or indwelling central venous catheters. The likelihood of upper extremity DVT increases as the catheter diameter and number of lumens increase. **Superficial venous thrombosis** usually presents with erythema, tenderness, and a “palpable cord.” Patients are at risk for extension of the thrombosis to the deep venous system.

DIAGNOSIS

Clinical Evaluation PE is known as “the Great Masquerader.” Diagnosis is difficult because symptoms and signs are nonspecific. The most common symptom is unexplained breathlessness. When occult PE occurs concomitantly with overt congestive heart failure or pneumonia, clinical improvement often fails to occur despite standard medical treatment of the concomitant illness. This scenario presents a clinical clue to the possible coexistence of PE.

With DVT, the most common symptom is a cramp or “charley horse” in the lower calf that persists and intensifies over several days. Point score criteria help estimate the clinical likelihood of DVT and PE (Table 300-1). Patients with a low-to-moderate likelihood of DVT or PE should undergo initial diagnostic evaluation with D-dimer testing alone (see “Blood Tests”) without obligatory imaging tests (Fig. 300-3). However, patients with a high clinical likelihood of VTE should skip D-dimer testing and undergo imaging as the next step in the diagnostic algorithm.

Clinical Pearls Not all leg pain is due to DVT, and not all dyspnea is due to PE (Table 300-2). Sudden, severe calf discomfort suggests a ruptured Baker’s cyst. Fever and chills usually herald cellulitis rather than DVT. Physical findings, if present, may consist only of mild palpation discomfort in the lower calf. However, massive DVT often presents with marked thigh swelling, tenderness, and erythema. If the leg is diffusely edematous, DVT is unlikely. More probable is an acute exacerbation of venous insufficiency due to postthrombotic syndrome. Upper extremity venous thrombosis may present with asymmetry in the supraclavicular fossa or in the circumference of the upper arms.

Pulmonary infarction usually indicates a small PE. This condition is exquisitely painful because the thrombus lodges peripherally, near the innervation of pleural nerves. *Nonthrombotic PE* etiologies include fat embolism after pelvic or long bone fracture, tumor embolism, bone marrow, and air embolism. Cement embolism and bony fragment embolism

TABLE 300-1 CLINICAL DECISION RULES

Low Clinical Likelihood of DVT if Point Score Is Zero or Less; Moderate Likelihood if Score Is 1 to 2; High Likelihood if Score Is 3 or Greater	
Clinical Variable	DVT Score
Active cancer	1
Paralysis, paresis, or recent cast	1
Bedridden for >3 days; major surgery <12 weeks	1
Tenderness along distribution of deep veins	1
Entire leg swelling	1
Unilateral calf swelling >3 cm	1
Pitting edema	1
Collateral superficial nonvaricose veins	1
Alternative diagnosis at least as likely as DVT	-2
High Clinical Likelihood of PE if Point Score Exceeds 4	
Clinical Variable	PE Score
Signs and symptoms of DVT	3.0
Alternative diagnosis less likely than PE	3.0
Heart rate >100/min	1.5
Immobilization >3 days; surgery within 4 weeks	1.5
Prior PE or DVT	1.5
Hemoptysis	1.0
Cancer	1.0

can occur after total hip or knee replacement. Intravenous drug users may inject themselves with a wide array of substances that can embolize such as hair, talc, and cotton. *Amniotic fluid embolism* occurs when fetal membranes leak or tear at the placental margin.

Nonimaging Diagnostic Modalities • BLOOD TESTS The quantitative *plasma D-dimer enzyme-linked immunosorbent assay (ELISA)* rises in the presence of DVT or PE because of the breakdown of fibrin by plasmin. Elevation of D-dimer indicates endogenous although often clinically ineffective thrombolysis. The sensitivity of the D-dimer is >80% for DVT (including isolated calf DVT) and >95% for PE. The D-dimer is less sensitive for DVT than for PE because the DVT thrombus size is smaller. A normal D-dimer is a useful “rule out” test. However, the D-dimer assay is not specific. Levels increase in patients with myocardial infarction, pneumonia, sepsis, cancer, and the postoperative state and those in the second or third trimester of pregnancy. Therefore, D-dimer rarely has a useful role among hospitalized patients, because levels are frequently elevated due to systemic illness.

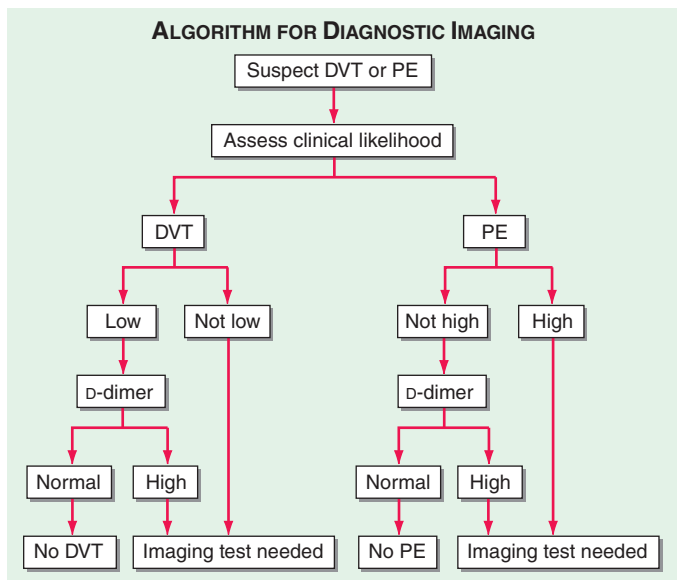


FIGURE 300-3 How to decide whether diagnostic imaging is needed. For assessment of clinical likelihood, see Table 300-1.

TABLE 300-2 DIFFERENTIAL DIAGNOSIS

DVT
Ruptured Baker's cyst
Cellulitis
Postphlebitic syndrome/venous insufficiency
PE
Pneumonia, asthma, chronic obstructive pulmonary disease
Congestive heart failure
Pericarditis
Pleurisy: "viral syndrome," costochondritis, musculoskeletal discomfort
Rib fracture, pneumothorax
Acute coronary syndrome
Anxiety

ELEVATED CARDIAC BIOMARKERS Serum troponin and plasma heart-type fatty acid-binding protein levels increase because of RV microinfarction. Myocardial stretch causes release of brain natriuretic peptide or NT-pro-brain natriuretic peptide.

ELECTROCARDIOGRAM The most frequently cited abnormality, in addition to sinus tachycardia, is the S1Q3T3 sign: an S wave in lead I, a Q wave in lead III, and an inverted T wave in lead III (Chap. 268). This finding is relatively specific but insensitive. RV strain and ischemia cause the most common abnormality, T-wave inversion in leads V₁ to V₄.

Noninvasive Imaging Modalities • VENOUS ULTRASONOGRAPHY Ultrasonography of the deep venous system relies on loss of vein compressibility as the primary criterion for DVT. When a normal vein is imaged in cross-section, it readily collapses with gentle manual pressure from the ultrasound transducer. This creates the illusion of a "wink." With acute DVT, the vein loses its compressibility because of passive distention by acute thrombus. The diagnosis of acute DVT is even more secure when thrombus is directly visualized. It appears homogeneous and has low echogenicity (Fig. 300-4). The vein itself often appears mildly dilated, and collateral channels may be absent.

Venous flow dynamics can be examined with Doppler imaging. Normally, manual calf compression causes augmentation of the Doppler flow pattern. Loss of normal respiratory variation is caused by an obstructing DVT or by any obstructive process within the pelvis. For patients with a technically poor or nondiagnostic venous ultrasound, one should consider alternative imaging modalities for DVT, such as computed tomography (CT) and magnetic resonance imaging.

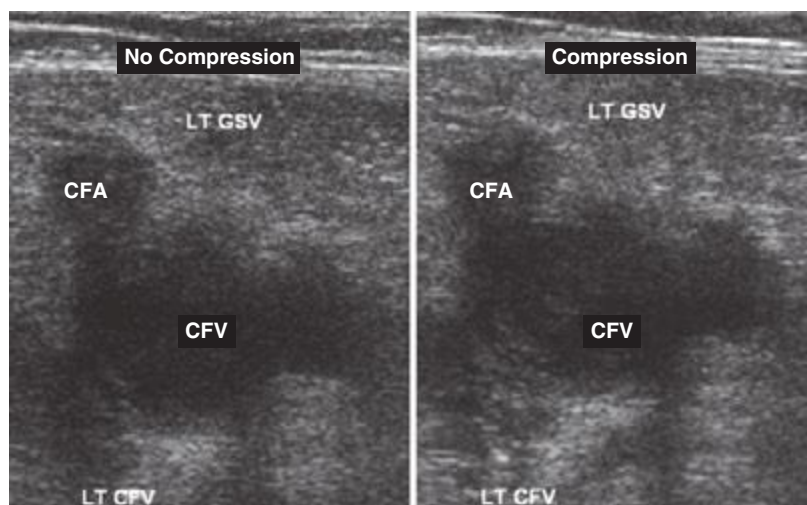


FIGURE 300-4 Venous ultrasound, with and without compression of the leg veins. CFA, common femoral artery; CFV, common femoral vein; GSV, great saphenous vein; LT, left.

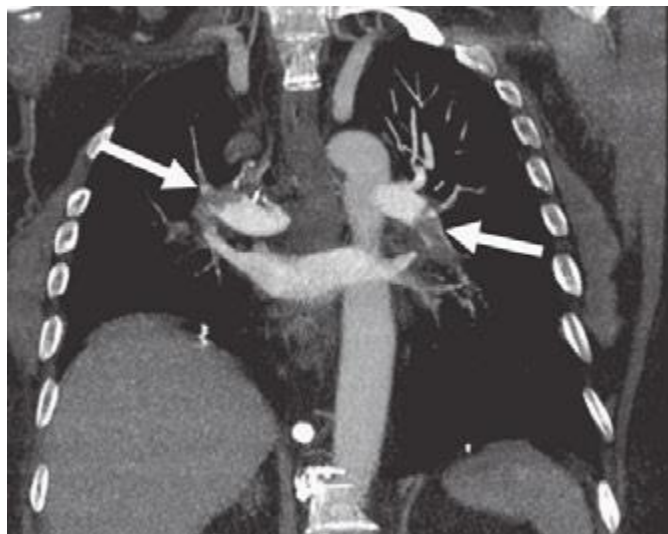


FIGURE 300-5 Large bilateral proximal PE on a coronal chest CT image in a 54-year-old man with lung cancer and brain metastases. He had developed sudden onset of chest heaviness and shortness of breath while at home. There are filling defects in the main and segmental pulmonary arteries bilaterally (white arrows). Only the left upper lobe segmental artery is free of thrombus.

CHEST ROENTGENOGRAPHY A normal or nearly normal chest x-ray often occurs in PE. Well-established abnormalities include focal oligemia (Westermark's sign), a peripheral wedged-shaped density above the diaphragm (Hampton's hump), and an enlarged right descending pulmonary artery (Palla's sign).

CHEST CT CT of the chest with intravenous contrast is the principal imaging test for the diagnosis of PE (Fig. 300-5). Multidetector-row spiral CT acquires all chest images with ≤ 1 mm of resolution during a short breath hold. Sixth-order branches can be visualized with resolution superior to that of conventional invasive contrast pulmonary angiography. The CT scan also provides an excellent four-chamber view of the heart. RV enlargement on chest CT indicates an increased likelihood of death within the next 30 days compared with PE patients who have normal RV size. When imaging is continued below the chest to the knee, pelvic and proximal leg DVT also can be diagnosed by CT scanning. In patients without PE, the lung parenchymal images may establish alternative diagnoses not apparent on chest x-ray that explain the presenting symptoms and signs such as pneumonia, emphysema, pulmonary fibrosis, pulmonary mass, and aortic pathology. Sometimes asymptomatic early-stage lung cancer is diagnosed incidentally.

LUNG SCANNING Lung scanning has become a second-line diagnostic test for PE, used mostly for patients who cannot tolerate intravenous contrast. Small particulate aggregates of albumin labeled with a gamma-emitting radionuclide are injected intravenously and are trapped in the pulmonary capillary bed. The perfusion scan defect indicates absent or decreased blood flow, possibly due to PE. Ventilation scans, obtained with a radiolabeled inhaled gas such as xenon or krypton, improve the specificity of the perfusion scan. Abnormal ventilation scans indicate abnormal nonventilated lung, thereby providing possible explanations for perfusion defects other than acute PE, such as asthma and chronic obstructive pulmonary disease. A high-probability scan for PE is defined as two or more segmental perfusion defects in the presence of normal ventilation.

The diagnosis of PE is very unlikely in patients with normal and nearly normal scans and is about 90% certain in patients with high-probability scans. Unfortunately, most

1634 patients have nondiagnostic scans, and fewer than one-half of patients with angiographically confirmed PE have a high probability scan. As many as 40% of patients with high clinical suspicion for PE but “low-probability” scans do, in fact, have PE at angiography.

MAGNETIC RESONANCE (MR) (CONTRAST-ENHANCED) IMAGING When ultrasound is equivocal, MR venography with gadolinium contrast is an excellent imaging modality to diagnose DVT. MR pulmonary angiography may detect large proximal PE but is not reliable for smaller segmental and subsegmental PE.

ECHOCARDIOGRAPHY Echocardiography is *not* a reliable diagnostic imaging tool for acute PE because most patients with PE have normal echocardiograms. However, echocardiography is a very useful diagnostic tool for detecting conditions that may mimic PE, such as acute myocardial infarction, pericardial tamponade, and aortic dissection. Transthoracic echocardiography rarely images thrombus directly. The best-known indirect sign of PE on transthoracic echocardiography is McConnell’s sign: hypokinesis of the RV free wall with normal or hyperkinetic motion of the RV apex. One should consider transesophageal echocardiography when CT scanning facilities are not available or when a patient has renal failure or severe contrast allergy that precludes administration of contrast despite premedication with high-dose steroids. This imaging modality can identify saddle, right main, or left main PE.

Invasive Diagnostic Modalities • PULMONARY ANGIOGRAPHY Chest CT with contrast (see above) has virtually replaced invasive pulmonary angiography as a diagnostic test. Invasive catheter-based diagnostic testing is reserved for patients with technically unsatisfactory chest CTs and for those in whom an interventional procedure such as catheter-directed thrombolysis is planned. A definitive diagnosis of PE depends on visualization of an intraluminal filling defect in more than one projection. Secondary signs of PE include abrupt occlusion (“cut-off”) of vessels, segmental oligemia or avascularity, a prolonged arterial phase with slow filling, and tortuous, tapering peripheral vessels.

CONTRAST PHLEBOGRAPHY Venous ultrasonography has virtually replaced contrast phlebography as the diagnostic test for suspected DVT.

Integrated Diagnostic Approach An integrated diagnostic approach (Fig. 300-3) streamlines the workup of suspected DVT and PE (Fig. 300-6).

TREATMENT DEEP VEIN THROMBOSIS

PRIMARY THERAPY

Primary therapy consists of clot dissolution with pharmacomechanical therapy that usually includes low-dose catheter-directed thrombolysis. This approach is reserved for patients with extensive femoral, iliofemoral, or upper extremity DVT. The open vein hypothesis postulates that patients who receive primary therapy will sustain less long-term damage to venous valves, with consequent lower rates of postthrombotic syndrome. A National Heart, Lung, and Blood Institute–sponsored randomized controlled trial called ATTRACT (NCT00790335) is testing this hypothesis.

SECONDARY PREVENTION

Anticoagulation or placement of an inferior vena caval filter constitutes secondary prevention of VTE. To lessen the severity of postthrombotic syndrome of the legs, below-knee graduated compression stockings may be prescribed, 30–40 mmHg, for 2 years after the DVT episode. They should be replaced every 3 months because they lose their elasticity.

TREATMENT PULMONARY EMBOLISM

RISK STRATIFICATION

Hemodynamic instability, RV dysfunction on echocardiography, RV enlargement on chest CT, or elevation of the troponin level

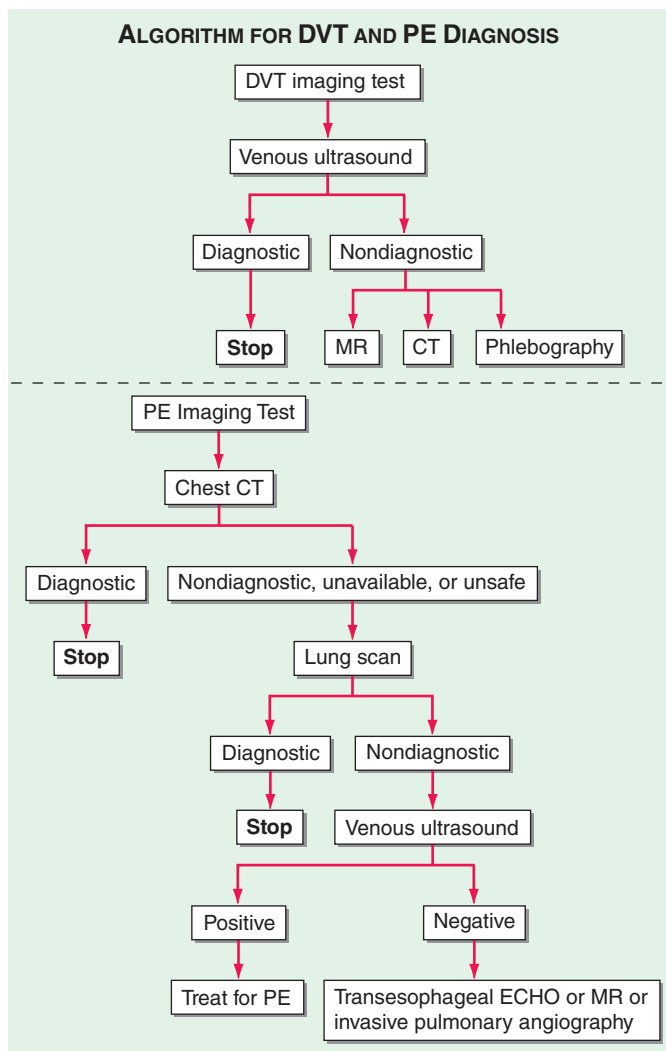


FIGURE 300-6 Imaging tests to diagnose DVT and PE. ECHO, echocardiography.

due to RV microinfarction portend a high risk of an adverse clinical outcome. When RV function remains normal in a hemodynamically stable patient, a good clinical outcome is highly likely with anticoagulation alone (Fig. 300-7).

ANTICOAGULATION

Effective anticoagulation is the foundation for successful treatment of DVT and PE. There are three options: (1) the conventional strategy

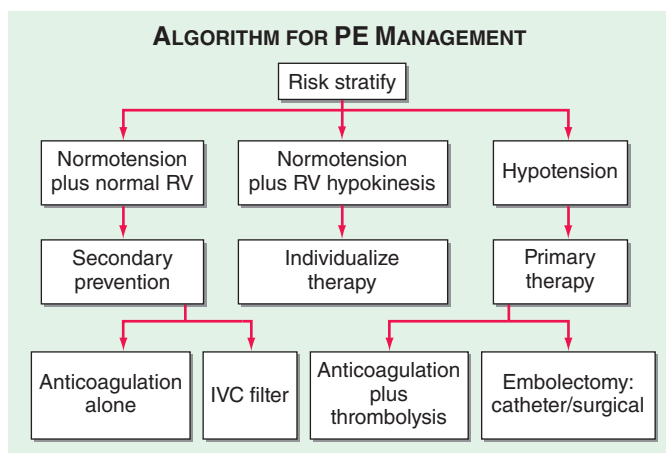


FIGURE 300-7 Acute management of pulmonary thromboembolism. RV, right ventricular; IVC, inferior vena cava.

TABLE 300-3 ANTICOAGULATION OF VTE**Immediate Anticoagulation**

Unfractionated heparin, bolus and continuous infusion, to achieve aPTT 2–3 times the upper limit of the laboratory normal, or
 Enoxaparin 1 mg/kg twice daily with normal renal function, or
 Dalteparin 200 U/kg once daily or 100 U/kg twice daily, with normal renal function, or
 Tinzaparin 175 U/kg once daily with normal renal function, or
 Fondaparinux weight-based once daily; adjust for impaired renal function
 Direct thrombin inhibitors: argatroban or bivalirudin
 Rivaroxaban 15 mg twice daily for 3 weeks, followed by 20 mg once daily with the dinner meal thereafter
 Apixaban (not yet licensed)

Warfarin Anticoagulation

Requires 5–10 days of administration to achieve effectiveness as monotherapy (Unfractionated heparin, low-molecular-weight heparin, and fondaparinux are the usual immediately effective “bridging agents” used when initiating warfarin)
 Usual start dose is 5 mg
 Titrate to INR, target 2.0–3.0
 Continue parenteral anticoagulation for a minimum of 5 days and until two sequential INR values, at least 1 day apart, achieve the target INR range

Novel Oral Anticoagulants for Extended-Duration Anticoagulation following Initial Parenteral Anticoagulation

Edoxaban (not yet licensed)
 Dabigatran (not yet licensed)

of parenteral therapy “bridged” to warfarin, (2) parenteral therapy “bridged” to a novel oral anticoagulant such as dabigatran (a direct thrombin inhibitor) or edoxaban (an anti-Xa agent), or (3) oral anticoagulation with rivaroxaban or apixaban (both are anti-Xa agents) with a loading dose followed by a maintenance dose as monotherapy without parenteral anticoagulation.

The three heparin-based parenteral anticoagulants are (1) unfractionated heparin (UFH), (2) low-molecular-weight heparin (LMWH), and (3) fondaparinux. For patients with suspected or proven heparin-induced thrombocytopenia, there are two parenteral direct thrombin inhibitors: argatroban and bivalirudin (Table 300-3).

Unfractionated Heparin UFH anticoagulates by binding to and accelerating the activity of antithrombin, thus preventing additional thrombus formation. UFH is dosed to achieve a target activated partial thromboplastin time (aPTT) of 60–80 s. The most popular nomogram uses an initial bolus of 80 U/kg, followed by an initial infusion rate of 18 U/kg per h.

The major advantage of UFH is its short half-life, which is especially useful in patients in whom hour-to-hour control of the intensity of anticoagulation is desired.

Low-Molecular-Weight Heparins These fragments of UFH exhibit less binding to plasma proteins and endothelial cells and consequently have greater bioavailability, a more predictable dose response, and a longer half-life than does UFH. No monitoring or dose adjustment is needed unless the patient is markedly obese or has chronic kidney disease.

Fondaparinux Fondaparinux, an anti-Xa pentasaccharide, is administered as a weight-based once-daily subcutaneous injection in a pre-filled syringe. No laboratory monitoring is required. Fondaparinux is synthesized in a laboratory and, unlike LMWH or UFH, is not derived from animal products. It does not cause heparin-induced thrombocytopenia. The dose must be adjusted downward for patients with renal dysfunction.

Warfarin This vitamin K antagonist prevents carboxylation activation of coagulation factors II, VII, IX, and X. The full effect of warfarin requires at least 5 days, even if the prothrombin time,

used for monitoring, becomes elevated more rapidly. If warfarin is initiated as monotherapy during an acute thrombotic illness, a paradoxical exacerbation of hypercoagulability increases the likelihood of thrombosis. Overlapping UFH, LMWH, fondaparinux, or parenteral direct thrombin inhibitors with warfarin for at least 5 days will nullify the early procoagulant effect of warfarin.

WARFARIN DOSING In an average-size adult, warfarin is often initiated in a dose of 5 mg. The prothrombin time is standardized by calculating the international normalized ratio (INR), which assesses the anticoagulant effect of warfarin (Chap. 78). The target INR is usually 2.5, with a range of 2.0–3.0.

The warfarin dose is usually titrated empirically to achieve the target INR. Proper dosing is difficult because hundreds of drug-drug and drug-food interactions affect warfarin metabolism. Increasing age and systemic illness reduce the required warfarin dose. Pharmacogenomics may provide more precise initial dosing of warfarin. *CYP2C9* variant alleles impair the hydroxylation of S-warfarin, thereby lowering the dose requirement. Variants in the gene encoding the vitamin K epoxide reductase complex 1 (*VKORC1*) can predict whether patients require low, moderate, or high warfarin doses.

Centralized anticoagulation clinics have improved the efficacy and safety of warfarin dosing. Patients can self-monitor their INR with a home point-of-care fingerstick machine and can occasionally be taught to self-dose their warfarin.

Novel Oral Anticoagulants Novel oral anticoagulants are administered in a fixed dose, establish effective anticoagulation within hours of ingestion, require no laboratory coagulation monitoring, and have few of the drug-drug or drug-food interactions that make warfarin so difficult to dose. Rivaroxaban, a factor Xa inhibitor, is approved for treatment of acute DVT and acute PE as monotherapy, without a parenteral “bridging” anticoagulant. Apixaban is likely to receive similar approval for oral monotherapy. Dabigatran, a direct thrombin inhibitor, and edoxaban, a factor Xa inhibitor, are likely to be approved for treatment of VTE after an initial course of parenteral anticoagulation.

Complications of Anticoagulants The most serious adverse effect of anticoagulation is hemorrhage. For life-threatening or intracranial hemorrhage due to heparin or LMWH, protamine sulfate can be administered. Heparin-induced thrombocytopenia is less common with LMWH than with UFH. There is no specific reversal agent for bleeding caused by fondaparinux, direct thrombin inhibitors, or factor Xa inhibitors.

Major bleeding from warfarin is best managed with prothrombin complex concentrate. With serious but non-life-threatening bleeding, fresh-frozen plasma or intravenous vitamin K can be used. Recombinant human coagulation factor VIIa (rFVIIa) is an off-label option to manage catastrophic bleeding from warfarin, but prothrombin complex concentrate is a better choice. Oral vitamin K is effective for managing minor bleeding or an excessively high INR in the absence of bleeding.

Duration of Anticoagulation For DVT isolated to an upper extremity or calf that has been provoked by surgery, trauma, estrogen, or an indwelling central venous catheter or pacemaker, 3 months of anticoagulation usually suffice. For an initial episode of provoked proximal leg DVT or PE, 3 to 6 months of anticoagulation are considered sufficient. For patients with cancer and VTE, prescribe LMWH as monotherapy without warfarin and continue anticoagulation indefinitely unless the patient is rendered cancer-free.

Among patients with idiopathic, unprovoked VTE, the recurrence rate is high after cessation of anticoagulation. VTE that occurs during long-haul air travel is considered unprovoked. Unprovoked VTE may be caused by an exacerbation of an underlying inflammatory state and can be conceptualized as a chronic illness, with latent periods between flares of recurrent episodes. American College of Chest Physicians (ACCP) guidelines recommend considering

1636 anticoagulation for an indefinite duration with a target INR between 2 and 3 for patients with idiopathic VTE. An alternative approach after the first 6 months of anticoagulation is to reduce the intensity of anticoagulation and to lower the target INR range to between 1.5 and 2.

Counterintuitively, the presence of genetic mutations such as heterozygous factor V Leiden and prothrombin gene mutation does not appear to increase the risk of recurrent VTE. However, patients with antiphospholipid antibody syndrome may warrant indefinite-duration anticoagulation, even if the initial VTE was provoked by trauma or surgery.

INFERIOR VENA CAVAL (IVC) FILTERS

The two principal indications for insertion of an IVC filter are (1) active bleeding that precludes anticoagulation and (2) recurrent venous thrombosis despite intensive anticoagulation. Prevention of recurrent PE in patients with right heart failure who are not candidates for fibrinolysis and prophylaxis of extremely high-risk patients are “softer” indications for filter placement. The filter itself may fail by permitting the passage of small-to medium-size clots. Large thrombi may embolize to the pulmonary arteries via collateral veins that develop. A more common complication is caval thrombosis with marked bilateral leg swelling.

Paradoxically, by providing a nidus for clot formation, filters increase the DVT rate, even though they usually prevent PE (over the short term). Retrievable filters can now be placed for patients with an anticipated temporary bleeding disorder or for patients at temporary high risk of PE, such as individuals undergoing bariatric surgery who have a prior history of perioperative PE. The filters can be retrieved up to several months after insertion unless thrombus forms and is trapped within the filter. The retrievable filter becomes permanent if it remains in place or if, for technical reasons such as rapid endothelialization, it cannot be removed.

MANAGEMENT OF MASSIVE PE

For patients with massive PE and hypotension, replete volume with 500 mL of normal saline. Additional fluid should be infused with extreme caution because excessive fluid administration exacerbates RV wall stress, causes more profound RV ischemia, and worsens LV compliance and filling by causing further interventricular septal shift toward the LV. Dopamine and dobutamine are first-line inotropic agents for treatment of PE-related shock. Maintain a low threshold for initiating these pressors. Often, a “trial-and-error” approach works best; other agents that may be effective include norepinephrine, vasopressin, or phenylephrine.

FIBRINOLYSIS

Successful fibrinolytic therapy rapidly reverses right heart failure and may result in a lower rate of death and recurrent PE by (1) dissolving much of the anatomically obstructing pulmonary arterial thrombus, (2) preventing the continued release of serotonin and other neurohumoral factors that exacerbate pulmonary hypertension, and (3) lysing much of the source of the thrombus in the pelvic or deep leg veins, thereby decreasing the likelihood of recurrent PE.

The preferred fibrinolytic regimen is 100 mg of recombinant tissue plasminogen activator (tPA) administered as a continuous peripheral intravenous infusion over 2 h. The sooner thrombolysis is administered, the more effective it is. However, this approach can be used for at least 14 days after the PE has occurred.

Contraindications to fibrinolysis include intracranial disease, recent surgery, and trauma. The overall major bleeding rate is about 10%, including a 1–3% risk of intracranial hemorrhage. Careful screening of patients for contraindications to fibrinolytic therapy (**Chap. 295**) is the best way to minimize bleeding risk.

The only Food and Drug Administration–approved indication for PE fibrinolysis is massive PE. For patients with submassive PE, who have preserved systolic blood pressure but moderate or severe RV dysfunction, use of fibrinolysis remains controversial. Results of a 1006-patient European multicentered randomized trial of submassive

PE, using the thrombolytic agent tenecteplase, were published in 2014. Death or hemodynamic collapse within 7 days of randomization was reduced by 56% in the tenecteplase group. However, hemorrhagic stroke occurred in 2% of tenecteplase patients versus 0.2% in patients who only received heparin.

PHARMACOMECHANICAL CATHETER-DIRECTED THERAPY

Many patients have relative contraindications to full-dose thrombolysis. Pharmacomechanical catheter-directed therapy usually combines physical fragmentation or pulverization of thrombus with catheter-directed low-dose thrombolysis. Mechanical techniques include catheter maceration and intentional embolization of clot more distally, suction thrombectomy, rheolytic hydrolysis, and low-energy ultrasound-facilitated thrombolysis. The dose of alteplase can be markedly reduced, usually to a range of 20 to 25 mg instead of the peripheral intravenous systemic dose of 100 mg.

PULMONARY EMBOLCTOMY

The risk of major hemorrhage with systemically administered fibrinolysis has prompted a renaissance of interest in surgical embolectomy, an operation that had almost become extinct. More rapid referral before the onset of irreversible multisystem organ failure and improved surgical technique have resulted in a high survival rate.

PULMONARY THROMBOENDARTERECTOMY

Chronic thromboembolic pulmonary hypertension develops in 2–4% of acute PE patients. Therefore, PE patients who have initial pulmonary hypertension (usually diagnosed with Doppler echocardiography) should be followed up at about 6 weeks with a repeat echocardiogram to determine whether pulmonary arterial pressure has normalized. Patients impaired by dyspnea due to chronic thromboembolic pulmonary hypertension should be considered for pulmonary thromboendarterectomy, which, if successful, can markedly reduce, and sometimes even cure, pulmonary hypertension (**Chap. 304**). The operation requires median sternotomy, cardiopulmonary bypass, deep hypothermia, and periods of hypothermic circulatory arrest. The mortality rate at experienced centers is approximately 5%. Inoperable patients should be managed with pulmonary vasodilator therapy.

EMOTIONAL SUPPORT

Patients with VTE may feel overwhelmed when they learn that they are suffering from PE or DVT. Some have never previously encountered serious cardiovascular illness. They wonder whether they will be able to adapt to the new limitations imposed by anticoagulation. They worry about the health of their families and the genetic implications of their illness. Those who are advised to discontinue anticoagulation may feel especially vulnerable about the potential for suffering recurrent VTE. At Brigham and Women’s Hospital, a physician-nurse–facilitated PE support group was initiated to address these concerns and has met monthly for more than 20 years.

PREVENTION OF VTE

Prevention of DVT and PE (**Table 300-4**) is of paramount importance because VTE is difficult to detect and poses a profound medical and economic burden. Low-dose UFH or LMWH is the most common form of in-hospital prophylaxis. Computerized reminder systems can increase the use of preventive measures and, at Brigham and Women’s Hospital, have reduced the symptomatic VTE rate by more than 40%. Audits of hospitals to ensure that prophylaxis protocols are being used will also increase utilization of preventive measures. Duration of prophylaxis is an important consideration. Extended-duration prophylaxis has not been shown to be both effective and safe in medically ill patients after hospital discharge in separate large trials that have tested enoxaparin, apixaban, and rivaroxaban. There is an ongoing trial of a novel oral anticoagulant, betrixaban, for extended-duration VTE prophylaxis in medically ill patients.

TABLE 300-4 PREVENTION OF VENOUS THROMBOEMBOLISM AMONG HOSPITALIZED PATIENTS

Condition	Prophylaxis Strategy
High-risk nonorthopedic surgery	Unfractionated heparin 5000 units SC bid or tid Enoxaparin 40 mg daily Dalteparin 2500 or 5000 units daily
Cancer surgery, including gynecologic cancer surgery	Enoxaparin 40 mg daily, consider 1 month of prophylaxis
Major orthopedic surgery	Warfarin (target INR 2.0–3.0) Enoxaparin 40 mg daily Enoxaparin 30 mg bid Dalteparin 2500 or 5000 units daily Fondaparinux 2.5 mg daily Rivaroxaban 10 mg daily Aspirin 81–325 mg daily Dabigatran 220 mg daily (not in the United States) Apixaban 2.5 mg bid (not in the United States) Intermittent pneumatic compression (with or without pharmacologic prophylaxis)
Medically ill patients, especially if immobilized, with a history of prior VTE, with an indwelling central venous catheter, or with cancer (but without active gastroduodenal ulcer, major bleeding within 3 months, or platelet count <50,000)	Unfractionated heparin 5000 units bid or tid Enoxaparin 40 mg daily Dalteparin 2500 or 5000 units daily Fondaparinux 2.5 mg daily
Anticoagulation contraindicated	Intermittent pneumatic compression devices (but whether graduated compression stockings are effective in medical patients is controversial)

Patients who have undergone total hip or knee replacement or cancer surgery will benefit from extended pharmacologic VTE prophylaxis after hospital discharge. For hip replacement or extensive cancer surgery, the duration of prophylaxis is usually at least 1 month.

double aortic arch, origin of the right subclavian artery distal to the left subclavian artery, and right-sided aortic arch with an aberrant left subclavian artery. A Kommerell's diverticulum is an anatomic remnant of a right aortic arch. Most congenital anomalies of the aorta do not cause symptoms and are detected during catheter-based procedures. The diagnosis of suspected congenital anomalies of the aorta typically is confirmed by computed tomographic (CT) or magnetic resonance (MR) angiography. Surgery is used to treat symptomatic anomalies.

301 Diseases of the Aorta

Mark A. Creager, Joseph Loscalzo

The aorta is the conduit through which blood ejected from the left ventricle is delivered to the systemic arterial bed. In adults, its diameter is approximately 3 cm at the origin and in the ascending portion, 2.5 cm in the descending portion in the thorax, and 1.8–2 cm in the abdomen. The aortic wall consists of a thin intima composed of endothelium, subendothelial connective tissue, and an internal elastic lamina; a thick tunica media composed of smooth muscle cells and extracellular matrix; and an adventitia composed primarily of connective tissue enclosing the vasa vasorum and nervi vascularis. In addition to the conduit function of the aorta, its viscoelastic and compliant properties serve a buffering function. The aorta is distended during systole to allow a portion of the stroke volume and elastic energy to be stored, and it recoils during diastole so that blood continues to flow to the periphery. Owing to its continuous exposure to high pulsatile pressure and shear stress, the aorta is particularly prone to injury and disease resulting from mechanical trauma. The aorta is also more prone to rupture than is any other vessel, especially with the development of aneurysmal dilation, since its wall tension, as governed by Laplace's law (i.e., proportional to the product of pressure and radius), will be increased.

CONGENITAL ANOMALIES OF THE AORTA

Congenital anomalies of the aorta usually involve the aortic arch and its branches. Symptoms such as dysphagia, stridor, and cough may occur if an anomaly causes a ring around or otherwise compresses the esophagus or trachea. Anomalies associated with symptoms include

AORTIC ANEURYSM

An *aneurysm* is defined as a pathologic dilation of a segment of a blood vessel. A *true aneurysm* involves all three layers of the vessel wall and is distinguished from a *pseudoaneurysm*, in which the intimal and medial layers are disrupted and the dilated segment of the aorta is lined by adventitia only and, at times, by perivascular clot. Aneurysms also may be classified according to their gross appearance. A *fusiform aneurysm* affects the entire circumference of a segment of the vessel, resulting in a diffusely dilated artery. In contrast, a *saccular aneurysm* involves only a portion of the circumference, resulting in an outpouching of the vessel wall. Aortic aneurysms also are classified according to location, i.e., abdominal versus thoracic. Aneurysms of the descending thoracic aorta are usually contiguous with infradiaphragmatic aneurysms and are referred to as *thoracoabdominal aortic aneurysms*.

ETIOLOGY

Aortic aneurysms result from conditions that cause degradation or abnormal production of the structural components of the aortic wall: elastin and collagen. The causes of aortic aneurysms may be broadly categorized as degenerative disorders, genetic or developmental diseases, vasculitis, infections, and trauma (Table 301-1). Inflammation, oxidative stress, proteolysis, and biomechanical wall stress contribute to the degenerative processes that characterize most aneurysms of the abdominal and descending thoracic aorta. These are mediated by B cell and T cell lymphocytes, macrophages, inflammatory cytokines, and matrix metalloproteinases that degrade elastin and collagen and alter the tensile strength and ability of the aorta to accommodate pulsatile stretch. The associated histopathology demonstrates destruction of elastin and collagen, decreased vascular smooth muscle, in-growth of new blood vessels, and inflammation. Factors associated with degenerative aortic aneurysms include aging, cigarette smoking, hypercholesterolemia, hypertension, and male sex.

Aortic aneurysm
Degenerative
Aging
Cigarette smoking
Hypercholesterolemia
Hypertension
Atherosclerosis
Genetic or developmental
Marfan's syndrome
Loeys-Dietz syndrome
Ehlers-Danlos syndrome type IV
Turner's syndrome
Familial
Bicuspid aortic valve
Chronic aortic dissection
Aortitis (see below)
Infective (see below)
Trauma
Acute aortic syndromes (aortic dissection, acute intramural hematoma, penetrating atherosclerotic ulcer)
Degenerative disorders (see above)
Genetic/developmental disorders (see above)
Hypertension
Aortitis (see below)
Pregnancy
Trauma
Aortic occlusion
Atherosclerosis
Thromboembolism
Aortitis
Vasculitis
Takayasu's arteritis
Giant cell arteritis
Rheumatic
HLA-B27-associated spondyloarthropathies
Behçet's syndrome
Cogan's syndrome
Idiopathic aortitis
Infective
Syphilis
Tuberculosis
Mycotic (<i>Salmonella</i> , staphylococcal, streptococcal, fungal)

The most common pathologic condition associated with degenerative aortic aneurysms is *atherosclerosis*. Many patients with aortic aneurysms have coexisting risk factors for atherosclerosis (Chap. 291e), as well as atherosclerosis in other blood vessels.

Medial degeneration, previously designated *cystic medial necrosis*, is the histopathologic term used to describe the degeneration of collagen and elastic fibers in the tunica media of the aorta as well as the loss of medial cells that are replaced by multiple clefts of mucoid material, such as proteoglycans. Medial degeneration characteristically affects the proximal aorta, results in circumferential weakness and dilation, and leads to the development of fusiform aneurysms involving the ascending aorta and the sinuses of Valsalva. This condition is particularly prevalent in patients with Marfan's syndrome, Loeys-Dietz syndrome, Ehlers-Danlos syndrome type IV (Chap. 427), hypertension, congenital bicuspid aortic valves, and familial thoracic aortic aneurysm syndromes; sometimes it appears as an isolated condition in patients without any other apparent disease.

Familial clusterings of aortic aneurysms occur in 20% of patients, suggesting a hereditary basis for the disease. Mutations of the gene that encodes fibrillin-1 are present in patients with Marfan's syndrome. Fibrillin-1 is an important component of extracellular microfibrils, which support the architecture of elastic fibers and other connective tissue. Deficiency of fibrillin-1 in the extracellular matrix leads to excessive signaling by transforming growth factor β (TGF- β). Loeys-Dietz syndrome is caused by mutations in the genes that encode TGF- β receptors 1 (*TGFBR1*) and 2 (*TGFBR2*). Increased signaling by TGF- β and mutations of *TGFBR1* and *TGFBR2* may cause thoracic aortic aneurysms. Mutations of type III procollagen have been implicated in Ehlers-Danlos type IV syndrome. Mutations of *SMAD3*, which encodes a downstream signaling protein involved with TGF binding to its receptors, have been described in a syndrome of thoracic aortic aneurysm; craniofacial, skeletal, and cutaneous anomalies; and osteoarthritis. Mutations of the genes encoding the smooth muscle-specific alpha-actin (*ACTA2*), smooth muscle cell-specific myosin heavy chain 11 (*MHC11*), and myosin light chain kinase (*MYLK*) and mutations of *TGFBR2* and *SMAD3* have been reported in some patients with non-syndromic familial thoracic aortic aneurysms.

The infectious causes of aortic aneurysms include syphilis, tuberculosis, and other bacterial infections. *Syphilis* (Chap. 206) is a relatively uncommon cause of aortic aneurysm. Syphilitic periaortitis and meso-aortitis damage elastic fibers, resulting in thickening and weakening of the aortic wall. Approximately 90% of syphilitic aneurysms are located in the ascending aorta or aortic arch. *Tuberculous aneurysms* (Chap. 202) typically affect the thoracic aorta and result from direct extension of infection from hilar lymph nodes or contiguous abscesses as well as from bacterial seeding. Loss of aortic wall elasticity results from granulomatous destruction of the medial layer. A *mycotic aneurysm* is a rare condition that develops as a result of staphylococcal, streptococcal, *Salmonella*, or other bacterial or fungal infections of the aorta, usually at an atherosclerotic plaque. These aneurysms are usually saccular. Blood cultures are often positive and reveal the nature of the infective agent.

Vasculitides associated with aortic aneurysm include Takayasu's arteritis and giant cell arteritis, which may cause aneurysms of the aortic arch and descending thoracic aorta. Spondyloarthropathies such as ankylosing spondylitis, rheumatoid arthritis, psoriatic arthritis, relapsing polychondritis, and reactive arthritis (formerly known as Reiter's syndrome) are associated with dilation of the ascending aorta. Aortic aneurysms occur in patients with Behçet's syndrome (Chap. 387), Cogan's syndrome, and IgG4-related systemic disease. Aortic aneurysms also result from idiopathic aortitis. *Traumatic aneurysms* may occur after penetrating or nonpenetrating chest trauma and most commonly affect the descending thoracic aorta just beyond the site of insertion of the ligamentum arteriosum. Chronic aortic dissections are associated with weakening of the aortic wall that may lead to the development of aneurysmal dilatation.

THORACIC AORTIC ANEURYSMS

The clinical manifestations and natural history of thoracic aortic aneurysms depend on their location. Medial degeneration is the most common pathology associated with ascending aortic aneurysms, whereas atherosclerosis is the condition most frequently associated with aneurysms of the descending thoracic aorta. The average growth rate of thoracic aneurysms is 0.1–0.2 cm per year. Thoracic aortic aneurysms associated with Marfan's syndrome or aortic dissection may expand at a greater rate. The risk of rupture is related to the size of the aneurysm and the presence of symptoms, ranging approximately from 2–3% per year for thoracic aortic aneurysms <4.0 cm in diameter to 7% per year for those >6 cm in diameter. Most thoracic aortic aneurysms are asymptomatic; however, compression or erosion of adjacent tissue by aneurysms may cause symptoms such as chest pain, shortness of breath, cough, hoarseness, and dysphagia. Aneurysmal dilation of the ascending aorta may cause congestive heart failure as a consequence of aortic regurgitation, and compression of the superior vena cava may produce congestion of the head, neck, and upper extremities.

A chest x-ray may be the first test that suggests the diagnosis of a thoracic aortic aneurysm (Fig. 301-1). Findings include widening



FIGURE 301-1 A chest x-ray of a patient with a thoracic aortic aneurysm.

of the mediastinal shadow and displacement or compression of the trachea or left main stem bronchus. Echocardiography, particularly transesophageal echocardiography, can be used to assess the proximal ascending aorta and descending thoracic aorta. Contrast-enhanced CT, magnetic resonance imaging (MRI), and conventional invasive aortography are sensitive and specific tests for assessment of aneurysms of the thoracic aorta and involvement of branch vessels (Fig. 301-2). In asymptomatic patients whose aneurysms are too small to justify surgery, noninvasive testing with either contrast-enhanced CT or MRI should be performed at least every 6–12 months to monitor expansion.

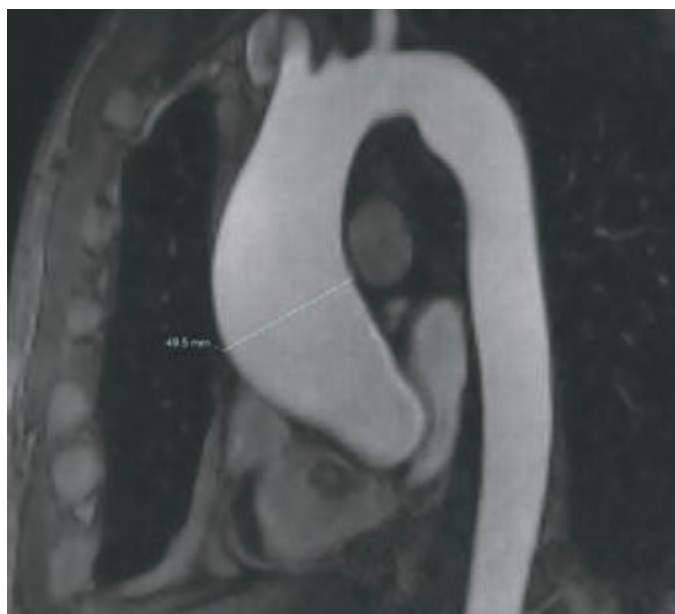


FIGURE 301-2 A magnetic resonance angiogram demonstrating a fusiform aneurysm of the ascending thoracic aorta. (Courtesy of Dr. Michael Steigner, Brigham and Women's Hospital, Boston, MA, with permission.)

β -Adrenergic blockers currently are recommended for patients with thoracic aortic aneurysms, particularly those with Marfan's syndrome, who have evidence of aortic root dilatation to reduce the rate of further expansion. Additional medical therapy should be given as necessary to control hypertension. Recent studies indicate that angiotensin receptor antagonists and angiotensin-converting enzyme inhibitors reduce the rate of aortic dilation in patients with Marfan's syndrome by blocking TGF- β signaling; clinical outcome trials of this treatment approach are in progress. Operative repair with placement of a prosthetic graft is indicated in patients with symptomatic ascending thoracic aortic aneurysms and for most asymptomatic aneurysms when the ascending aortic diameter is >5.5 cm. In patients with Marfan's syndrome or bicuspid aortic valve, ascending thoracic aortic aneurysms of 4–5 cm should be considered for surgery. Operative repair is indicated for patients with descending thoracic aortic aneurysms when the diameter is >6 cm, and endovascular repair should be considered if feasible when the diameter is >5.5 cm. Repair is also recommended when the diameter of an aneurysm has increased >1 cm per year.

ABDOMINAL AORTIC ANEURYSMS

Abdominal aortic aneurysms occur more frequently in males than in females, and the incidence increases with age. Abdominal aortic aneurysms ≥ 4.0 cm may affect 1–2% of men older than 50 years. At least 90% of all abdominal aortic aneurysms >4.0 cm are related to atherosclerotic disease, and most of these aneurysms are below the level of the renal arteries. Prognosis is related to both the size of the aneurysm and the severity of coexisting coronary artery and cerebrovascular disease. The risk of rupture increases with the size of the aneurysm: the 5-year risk for aneurysms <5 cm is 1–2%, whereas it is 20–40% for aneurysms >5 cm in diameter. The formation of mural thrombi within aneurysms may predispose to peripheral embolization.

An abdominal aortic aneurysm commonly produces no symptoms. It usually is detected on routine examination as a palpable, pulsatile, expansile, and nontender mass, or it is an incidental finding observed on an abdominal imaging study performed for other reasons. As abdominal aortic aneurysms expand, however, they may become painful. Some patients complain of strong pulsations in the abdomen; others experience pain in the chest, lower back, or scrotum. Aneurysmal pain is usually a harbinger of rupture and represents a medical emergency. More often, acute rupture occurs without any prior warning, and this complication is always life-threatening. Rarely, there is leakage of the aneurysm with severe pain and tenderness. Acute pain and hypotension occur with rupture of the aneurysm, which requires an emergency operation.

Abdominal radiography may demonstrate the calcified outline of the aneurysm; however, about 25% of aneurysms are not calcified and cannot be visualized by x-ray imaging. An abdominal ultrasound can delineate the transverse and longitudinal dimensions of an abdominal aortic aneurysm and may detect mural thrombus. Abdominal ultrasound is useful for serial documentation of aneurysm size and can be used to screen patients at risk for developing an aortic aneurysm. In one large study, ultrasound screening of men age 65–74 years was associated with a risk reduction in aneurysm-related death of 42%. For this reason, screening by ultrasonography is recommended for men age 65–75 years who have ever smoked. In addition, siblings or offspring of persons with abdominal aortic aneurysms, as well as individuals with thoracic aortic or peripheral arterial aneurysms, should be considered for screening for abdominal aortic aneurysms. CT with contrast and MRI are accurate noninvasive tests to determine the location and size of abdominal aortic aneurysms and to plan endovascular or open surgical repair (Fig. 301-3). Contrast aortography may be used for the evaluation of patients with aneurysms, but the procedure carries a small risk of complications such as bleeding, allergic reactions, and atheroembolism. Since the presence of mural thrombi may reduce the luminal size, aortography may underestimate the diameter of an aneurysm.



FIGURE 301-3 A computed tomographic angiogram depicting a fusiform abdominal aortic aneurysm before (left) and after (right) treatment with a bifurcated stent graft. (Courtesy of Drs. Elizabeth George and Frank Rybicki, Brigham and Women's Hospital, Boston, MA, with permission.)

TREATMENT ABDOMINAL AORTIC ANEURYSMS

Operative repair of the aneurysm with insertion of a prosthetic graft or endovascular placement of an aortic stent graft (Fig. 301-3) is indicated for abdominal aortic aneurysms of any size that are expanding rapidly or are associated with symptoms. For asymptomatic aneurysms, abdominal aortic aneurysm repair is indicated if the diameter is >5.5 cm. In randomized trials of patients with abdominal aortic aneurysms <5.5 cm, there was no difference in the long-term (5- to 8-year) mortality rate between those followed with ultrasound surveillance and those undergoing elective surgical repair. Thus, serial noninvasive follow-up of smaller aneurysms (<5 cm) is an alternative to immediate repair. The decision to perform an open surgical operation or endovascular repair is based in part on the vascular anatomy and comorbid conditions. Endovascular repair of abdominal aortic aneurysms has a lower short-term morbidity rate but a comparable long-term mortality rate with open surgical reconstruction. Long-term surveillance with CT or MR aortography is indicated after endovascular repair to detect leaks and possible aneurysm expansion.

In surgical candidates, careful preoperative cardiac and general medical evaluations (followed by appropriate therapy for complicating conditions) are essential. Preexisting coronary artery disease, congestive heart failure, pulmonary disease, diabetes mellitus, and advanced age add to the risk of surgery. β -Adrenergic blockers decrease perioperative cardiovascular morbidity and mortality. With careful preoperative cardiac evaluation and postoperative care, the operative mortality rate approximates 1–2%. After acute rupture, the mortality rate of emergent operation is 45–50%. Endovascular repair with stent placement is an alternative approach to treat ruptured aneurysms and may be associated with a lower mortality rate.

ACUTE AORTIC SYNDROMES

The four major acute aortic syndromes are aortic rupture (discussed earlier), aortic dissection, intramural hematoma, and penetrating atherosclerotic ulcer. Aortic dissection is caused by a circumferential or, less frequently, transverse tear of the intima. It often occurs along the right lateral wall of the ascending aorta where the hydraulic shear

stress is high. Another common site is the descending thoracic aorta just below the ligamentum arteriosum. The initiating event is either a primary intimal tear with secondary dissection into the media or a medial hemorrhage that dissects into and disrupts the intima. The pulsatile aortic flow then dissects along the elastic lamellar plates of the aorta and creates a false lumen. The dissection usually propagates distally down the descending aorta and into its major branches, but it may propagate proximally. Distal propagation may be limited by atherosclerotic plaque. In some cases, a secondary distal intimal disruption occurs, resulting in the reentry of blood from the false to the true lumen.

There are at least two important pathologic and radiologic variants of aortic dissection: intramural hematoma without an intimal flap and penetrating atherosclerotic ulcer. Acute intramural hematoma is thought to result from rupture of the vasa vasorum with hemorrhage into the wall of the aorta. Most of these hematomas occur in the descending thoracic aorta. Acute intramural hematomas may progress to dissection and rupture. Penetrating atherosclerotic ulcers are caused by erosion of a plaque into the aortic media, are usually localized, and are not associated with extensive propagation. They are found primarily in the middle and distal portions of the descending thoracic aorta and are associated with extensive atherosclerotic disease. The ulcer can erode beyond the internal elastic lamina, leading to medial hematoma, and may progress to false aneurysm formation or rupture.

Several classification schemes have been developed for thoracic aortic dissections. DeBakey and colleagues initially classified aortic dissections as type I, in which an intimal tear occurs in the ascending aorta but involves the descending aorta as well; type II, in which the dissection is limited to the ascending aorta; and type III, in which the intimal tear is located in the descending aorta with distal propagation of the dissection (Fig. 301-4). Another classification (Stanford) is that of type A, in which the dissection involves the ascending aorta (proximal dissection), and type B, in which it is limited to the arch and/or descending aorta (distal dissection). From a management standpoint, classification of aortic dissections and intramural hematomas into type A or B is more practical and useful, since DeBakey types I and II are managed in a similar manner.

The factors that predispose to aortic dissection include those associated with medial degeneration and others that increase aortic wall stress (Table 301-1). Systemic hypertension is a coexisting condition in

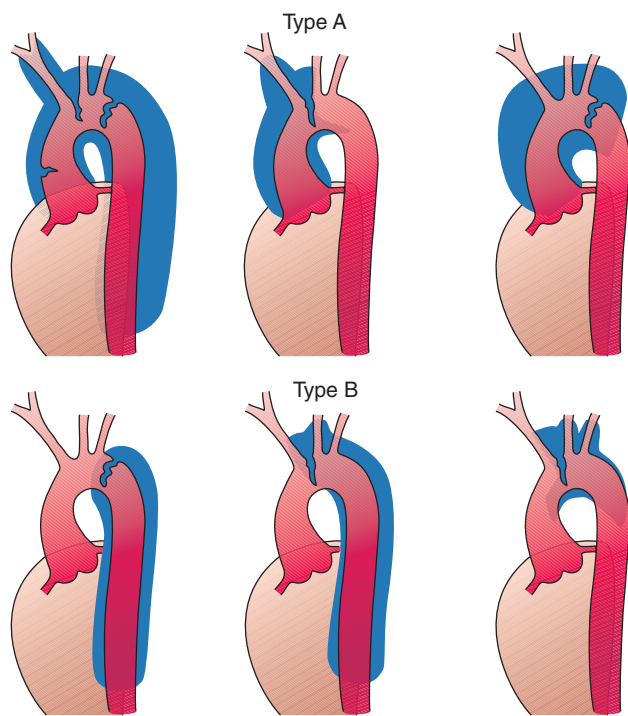


FIGURE 301-4 Classification of aortic dissections. Stanford classification: Type A dissections (*top*) involve the ascending aorta independent of site of tear and distal extension; type B dissections (*bottom*) involve transverse and/or descending aorta without involvement of the ascending aorta. DeBakey classification: Type I dissection involves ascending to descending aorta (*top left*); type II dissection is limited to ascending or transverse aorta, without descending aorta (*top center + top right*); type III dissection involves descending aorta only (*bottom left*). (From DC Miller, in RM Doroghazi, EE Slater [eds]: *Aortic Dissection*. New York, McGraw-Hill, 1983, with permission.)

70% of patients. Aortic dissection is the major cause of morbidity and mortality in patients with Marfan's syndrome ([Chap. 427](#)) or Loeys-Dietz syndrome, and similarly may affect patients with Ehlers-Danlos syndrome. The incidence also is increased in patients with inflammatory aortitis (i.e., Takayasu's arteritis, giant cell arteritis), congenital aortic valve anomalies (e.g., bicuspid valve), coarctation of the aorta, and a history of aortic trauma. In addition, the risk of dissection is increased in otherwise normal women during the third trimester of pregnancy. Aortic dissection also may occur as a consequence of weight lifting, cocaine use, or deceleration injury.

CLINICAL MANIFESTATIONS

The peak incidence of aortic dissection is in the sixth and seventh decades. Men are more affected than women by a ratio of 2:1. The presentations of aortic dissection and its variants are the consequences of intimal tear, dissecting hematoma, occlusion of involved arteries, and compression of adjacent tissues. Acute aortic dissection presents with the sudden onset of pain ([Chap. 19](#)), which often is described as very severe and tearing and is associated with diaphoresis. The pain may be localized to the front or back of the chest, often the interscapular region, and typically migrates with propagation of the dissection. Other symptoms include syncope, dyspnea, and weakness. Physical findings may include hypertension or hypotension, loss of pulses, aortic regurgitation, pulmonary edema, and neurologic findings due to carotid artery obstruction (hemiplegia, hemianesthesia) or spinal cord ischemia (paraplegia). Bowel ischemia, hematuria, and myocardial ischemia have all been observed. These clinical manifestations reflect complications resulting from the dissection occluding the major arteries. Furthermore, clinical manifestations may result from the compression of adjacent structures (e.g., superior cervical ganglia, superior vena cava, bronchus, esophagus) by the expanding dissection, causing aneurysmal dilation, and include Horner's syndrome, superior

vena cava syndrome, hoarseness, dysphagia, and airway compromise. Hemopericardium and cardiac tamponade may complicate a type A lesion with retrograde dissection. Acute aortic regurgitation is an important and common (>50%) complication of proximal dissection. It is the outcome of either a circumferential tear that widens the aortic root or a disruption of the annulus by a dissecting hematoma that tears a leaflet(s) or displaces it, inferior to the line of closure. Signs of aortic regurgitation include bounding pulses, a wide pulse pressure, a diastolic murmur often radiating along the right sternal border, and evidence of congestive heart failure. The clinical manifestations depend on the severity of the regurgitation.

In dissections involving the ascending aorta, the chest x-ray often reveals a widened superior mediastinum. A pleural effusion (usually left-sided) also may be present. This effusion is typically serosanguineous and not indicative of rupture unless accompanied by hypotension and falling hematocrit. In dissections of the descending thoracic aorta, a widened mediastinum may be observed on chest x-ray. In addition, the descending aorta may appear to be wider than the ascending portion. An electrocardiogram that shows no evidence of myocardial ischemia is helpful in distinguishing aortic dissection from myocardial infarction. Rarely, the dissection involves the right or, less commonly, left coronary ostium and causes acute myocardial infarction.

The diagnosis of aortic dissection can be established by noninvasive techniques such as echocardiography, CT, and MRI. Aortography is used less commonly because of the accuracy of these noninvasive techniques. Transthoracic echocardiography can be performed simply and rapidly and has an overall sensitivity of 60–85% for aortic dissection. For diagnosing proximal ascending aortic dissections, its sensitivity exceeds 80%; it is less useful for detecting dissection of the arch and descending thoracic aorta. Transesophageal echocardiography requires greater skill and patient cooperation but is very accurate in identifying dissections of the ascending and descending thoracic aorta but not the arch, achieving 98% sensitivity and approximately 90% specificity. Echocardiography also provides important information regarding the presence and severity of aortic regurgitation and pericardial effusion. CT and MRI are both highly accurate in identifying the intimal flap and the extent of the dissection and involvement of major arteries; each has a sensitivity and specificity >90%. They are useful in recognizing intramural hemorrhage and penetrating ulcers. The relative utility of transesophageal echocardiography, CT, and MRI depends on the availability and expertise in individual institutions as well as on the hemodynamic stability of the patient, with CT and MRI obviously less suitable for unstable patients.

TREATMENT AORTIC DISSECTION

Medical therapy should be initiated as soon as the diagnosis is considered. The patient should be admitted to an intensive care unit for hemodynamic monitoring. Unless hypotension is present, therapy should be aimed at reducing cardiac contractility and systemic arterial pressure, and thus shear stress. For acute dissection, unless contraindicated, β -adrenergic blockers should be administered parenterally, using intravenous propranolol, metoprolol, or the short-acting esmolol to achieve a heart rate of approximately 60 beats/min. This should be accompanied by sodium nitroprusside infusion to lower systolic blood pressure to ≤ 120 mmHg. Labetalol ([Chap. 298](#)), a drug with both β - and α -adrenergic blocking properties, also may be used as a parenteral agent in acute therapy for dissection.

The calcium channel antagonists verapamil and diltiazem may be used intravenously if nitroprusside or β -adrenergic blockers cannot be employed. The addition of a parenteral angiotensin-converting enzyme (ACE) inhibitor such as enalaprilat to a β -adrenergic blocker also may be considered. Isolated use of a direct vasodilator such as hydralazine is contraindicated because these agents can increase hydraulic shear and may propagate the dissection.

Emergent or urgent surgical correction is the preferred treatment for acute ascending aortic dissections and intramural hematomas (type A) and for complicated type B dissections, including

1642 those characterized by propagation, compromise of major aortic branches, impending rupture, or continued pain. Surgery involves excision of the intimal flap, obliteration of the false lumen, and placement of an interposition graft. A composite valve-graft conduit is used if the aortic valve is disrupted. The overall in-hospital mortality rate after surgical treatment of patients with aortic dissection is reported to be 15–25%. The major causes of perioperative mortality and morbidity include myocardial infarction, paraplegia, renal failure, tamponade, hemorrhage, and sepsis. Endoluminal stent grafts may be considered in selected patients. Other transcatheter techniques, such as fenestration of the intimal flaps and stenting of narrowed branch vessels to increase flow to compromised organs, are used in selected patients. For uncomplicated and stable distal dissections and intramural hematomas (type B), medical therapy is the preferred treatment. The in-hospital mortality rate of medically treated patients with type B dissection is 10–20%. Long-term therapy for patients with aortic dissection and intramural hematomas (with or without surgery) consists of control of hypertension and reduction of cardiac contractility with the use of beta blockers plus other antihypertensive agents, such as ACE inhibitors or calcium antagonists. Patients with chronic type B dissection and intramural hematomas should be followed on an outpatient basis every 6–12 months with contrast-enhanced CT or MRI to detect propagation or expansion. Patients with Marfan's syndrome are at high risk for postdissection complications. The long-term prognosis for patients with treated dissections is generally good with careful follow-up; the 10-year survival rate is approximately 60%.

CHRONIC ATHEROSCLEROTIC OCCLUSIVE DISEASE

Atherosclerosis may affect the thoracic and abdominal aorta. Occlusive aortic disease caused by atherosclerosis usually is confined to the distal abdominal aorta below the renal arteries. Frequently the disease extends to the iliac arteries (Chap. 302). Claudication characteristically involves the buttocks, thighs, and calves and may be associated with impotence in males (Leriche's syndrome). The severity of the symptoms depends on the adequacy of collaterals. With sufficient collateral blood flow, a complete occlusion of the abdominal aorta may occur without the development of ischemic symptoms. The physical findings include the absence of femoral and other distal pulses bilaterally and the detection of an audible bruit over the abdomen (usually at or below the umbilicus) and the common femoral arteries. Atrophic skin, loss of hair, and coolness of the lower extremities usually are observed. In advanced ischemia, rubor on dependency and pallor on elevation can be seen.

The diagnosis usually is established by physical examination and noninvasive testing, including leg pressure measurements, Doppler velocity analysis, pulse volume recordings, and duplex ultrasonography. The anatomy may be defined by MRI, CT, or conventional aortography, typically performed when one is considering revascularization. Catheter-based endovascular or operative treatment is indicated in patients with lifestyle-limiting or debilitating symptoms of claudication and patients with critical limb ischemia.

ACUTE AORTIC OCCLUSION

Acute occlusion in the distal abdominal aorta constitutes a medical emergency because it threatens the viability of the lower extremities; it usually results from an occlusive (saddle) embolus that almost always originates from the heart. Rarely, acute occlusion may occur as the result of in situ thrombosis in a preexisting severely narrowed segment of the aorta.

The clinical picture is one of acute ischemia of the lower extremities. Severe rest pain, coolness, and pallor of the lower extremities and the absence of distal pulses bilaterally are the usual manifestations. Diagnosis should be established rapidly by MRI, CT, or aortography. Emergency thrombectomy or revascularization is indicated.

AORTITIS

Aortitis, a term referring to inflammatory disease of the aorta, may be caused by large vessel vasculitides such as Takayasu's arteritis and

giant cell arteritis, rheumatic and HLA-B27-associated spondyloarthropathies, Behçet's syndrome, antineutrophil cytoplasmic antibody (ANCA)-associated vasculitides, Cogan's syndrome, IgG4-related systemic disease, and infections such as syphilis, tuberculosis, and *Salmonella*, or may be associated with retroperitoneal fibrosis. Aortitis may result in aneurysmal dilation and aortic regurgitation, occlusion of the aorta and its branch vessels, or acute aortic syndromes.

TAKAYASU'S ARTERITIS

This inflammatory disease often affects the ascending aorta and aortic arch, causing obstruction of the aorta and its major arteries. Takayasu's arteritis is also termed *pulseless disease* because of the frequent occlusion of the large arteries originating from the aorta. It also may involve the descending thoracic and abdominal aorta and occlude large branches such as the renal arteries. Aortic aneurysms also may occur. The pathology is a panarteritis characterized by mononuclear cells and occasionally giant cells, with marked intimal hyperplasia, medial and adventitial thickening, and, in the chronic form, fibrotic occlusion. The disease is most prevalent in young females of Asian descent but does occur in women of other geographic and ethnic origins and also in young men. During the acute stage, fever, malaise, weight loss, and other systemic symptoms may be evident. Elevations of the erythrocyte sedimentation rate and C-reactive protein are common. The chronic stages of the disease, which is intermittently active, present with symptoms related to large artery occlusion, such as upper extremity claudication, cerebral ischemia, and syncope. The process is progressive, and there is no definitive therapy. Glucocorticoids and immunosuppressive agents are effective in some patients during the acute phase. Surgical bypass or endovascular intervention of a critically stenotic artery may be necessary.

GIANT CELL ARTERITIS

(See also Chap. 385) This vasculitis occurs in older individuals and affects women more often than men. Primarily large and medium-size arteries are affected. The pathology is that of focal granulomatous lesions involving the entire arterial wall; it may be associated with polymyalgia rheumatica. Obstruction of medium-size arteries (e.g., temporal and ophthalmic arteries) and major branches of the aorta and the development of aortitis and aortic regurgitation are important complications of the disease. High-dose glucocorticoid therapy may be effective when given early.

RHEUMATIC AORTITIS

Rheumatoid arthritis (Chap. 380), ankylosing spondylitis (Chap. 384), psoriatic arthritis (Chap. 384), reactive arthritis (formerly known as Reiter's syndrome) (Chap. 384), relapsing polychondritis, and inflammatory bowel disorders may all be associated with aortitis involving the ascending aorta. The inflammatory lesions usually involve the ascending aorta and may extend to the sinuses of Valsalva, the mitral valve leaflets, and adjacent myocardium. The clinical manifestations are aneurysm, aortic regurgitation, and involvement of the cardiac conduction system.

IDIOPATHIC AORTITIS

Idiopathic abdominal aortitis is characterized by adventitial and periaortic inflammation with thickening of the aortic wall. It is associated with abdominal aortic aneurysms and idiopathic retroperitoneal fibrosis. Affected individuals may present with vague constitutional symptoms, fever, and abdominal pain. Retroperitoneal fibrosis can cause ureteral obstruction and hydronephrosis. Glucocorticoids and immunosuppressive agents may reduce the inflammation.

INFECTIVE AORTITIS

Infective aortitis may result from direct invasion of the aortic wall by bacterial pathogens such as *Staphylococcus*, *Streptococcus*, and *Salmonella* or by fungi. These bacteria cause aortitis by infecting the aorta at sites of atherosclerotic plaque. Bacterial proteases lead to degradation of collagen, and the ensuing destruction of the aortic wall leads to the formation of a saccular aneurysm referred to as a mycotic

aneurysm. Mycotic aneurysms have a predilection for the suprarenal abdominal aorta. The pathologic characteristics of the aortic wall include acute and chronic inflammation, abscesses, hemorrhage, and necrosis. Mycotic aneurysms typically affect the elderly and occur in men three times more frequently than in women. Patients may present with fever, sepsis, and chest, back, or abdominal pain; there may have been a preceding diarrheal illness. Blood cultures are positive in the majority of patients. Both CT and MRI are useful to diagnose mycotic aneurysms. Treatment includes antibiotic therapy and surgical removal of the affected part of the aorta and revascularization of the lower extremities with grafts placed in uninfected tissue.

Syphilitic aortitis is a late manifestation of luetic infection (**Chap. 206**) that usually affects the proximal ascending aorta, particularly the aortic root, resulting in aortic dilation and aneurysm formation. Syphilitic aortitis occasionally may involve the aortic arch or the descending aorta. The aneurysms may be saccular or fusiform and are usually asymptomatic, but compression of and erosion into adjacent structures may result in symptoms; rupture also may occur.

The initial lesion is an obliterative endarteritis of the vasa vasorum, especially in the adventitia. This is an inflammatory response to the invasion of the adventitia by the spirochetes. Destruction of the aortic media occurs as the spirochetes spread into this layer, usually via the lymphatics accompanying the vasa vasorum. Destruction of collagen and elastic tissues leads to dilation of the aorta, scar formation, and calcification. These changes account for the characteristic radiographic appearance of linear calcification of the ascending aorta.

The disease typically presents as an incidental chest radiographic finding 15–30 years after initial infection. Symptoms may result from aortic regurgitation, narrowing of coronary ostia due to syphilitic aortitis, compression of adjacent structures (e.g., esophagus), or rupture. Diagnosis is established by a positive serologic test, i.e., rapid plasmin reagin (RPR) or fluorescent treponemal antibody. Treatment includes penicillin and surgical excision and repair.

302 Arterial Diseases of the Extremities

Mark A. Creager, Joseph Loscalzo

PERIPHERAL ARTERY DISEASE

Peripheral artery disease (PAD) is defined as a clinical disorder in which there is a stenosis or occlusion in the aorta or the arteries of the limbs. Atherosclerosis is the leading cause of PAD in patients >40 years old. Other causes include thrombosis, embolism, vasculitis, fibromuscular dysplasia, entrapment, cystic adventitial disease, and trauma. The highest prevalence of atherosclerotic PAD occurs in the sixth and seventh decades of life. As in patients with atherosclerosis of the coronary and cerebral vasculature, there is an increased risk of developing PAD in cigarette smokers and in persons with diabetes mellitus, hypercholesterolemia, hypertension, or renal insufficiency.

Pathology (See also **Chap. 291e**) Segmental lesions that cause stenosis or occlusion are usually localized to large and medium-size vessels. The pathology of the lesions includes atherosclerotic plaques with calcium deposition, thinning of the media, patchy destruction of muscle and elastic fibers, fragmentation of the internal elastic lamina, and thrombi composed of platelets and fibrin. The primary sites of involvement are the abdominal aorta and iliac arteries (30% of symptomatic patients), the femoral and popliteal arteries (80–90% of patients), and the more distal vessels, including the tibial and peroneal arteries (40–50% of patients). Atherosclerotic lesions occur preferentially at arterial branch points, which are sites of increased turbulence, altered shear stress, and intimal injury. Involvement of the distal vasculature is most common in elderly individuals and patients with diabetes mellitus.

Clinical Evaluation Fewer than 50% of patients with PAD are symptomatic, although many have a slow or impaired gait. The most common *symptom* is intermittent claudication, which is defined as a pain, ache, cramp, numbness, or a sense of fatigue in the muscles; it occurs during exercise and is relieved by rest. The site of claudication is distal to the location of the occlusive lesion. For example, buttock, hip, thigh, and calf discomfort occurs in patients with aortoiliac disease, whereas calf claudication develops in patients with femoral-popliteal disease. Symptoms are far more common in the lower than in the upper extremities because of the higher incidence of obstructive lesions in the former region. In patients with severe arterial occlusive disease in whom resting blood flow cannot accommodate basal nutritional needs of the tissues, critical limb ischemia may develop. Patients complain of rest pain or a feeling of cold or numbness in the foot and toes. Frequently, these symptoms occur at night when the legs are horizontal and improve when the legs are in a dependent position. With severe ischemia, rest pain may be persistent.

Important *physical findings* of PAD include decreased or absent pulses distal to the obstruction, the presence of bruits over the narrowed artery, and muscle atrophy. With more severe disease, hair loss, thickened nails, smooth and shiny skin, reduced skin temperature, and pallor or cyanosis are common physical signs. In patients with critical limb ischemia, ulcers or gangrene may occur. Elevation of the legs and repeated flexing of the calf muscles produce pallor of the soles of the feet, whereas rubor, secondary to reactive hyperemia, may develop when the legs are dependent. The time required for rubor to develop or for the veins in the foot to fill when the patient's legs are transferred from an elevated to a dependent position is related to the severity of the ischemia and the presence of collateral vessels. Patients with severe ischemia may develop peripheral edema because they keep their legs in a dependent position much of the time. Ischemic neuropathy can result in numbness and hyporeflexia.

Noninvasive Testing The history and physical examination are often sufficient to establish the diagnosis of PAD. An objective assessment of the presence and severity of disease is obtained by noninvasive techniques. Arterial pressure can be recorded noninvasively in the legs by placement of sphygmomanometric cuffs at the ankles and the use of a Doppler device to auscultate or record blood flow from the dorsalis pedis and posterior tibial arteries. Normally, systolic blood pressure in the legs and arms is similar. Indeed, ankle pressure may be slightly higher than arm pressure due to pulse-wave amplification. In the presence of hemodynamically significant stenoses, the systolic blood pressure in the leg is decreased. Thus, the ratio of the ankle and brachial artery pressures (termed the *ankle:brachial index*, or ABI) is 1.00–1.40 in normal individuals. ABI values of 0.91–0.99 are considered “borderline,” and those <0.90 are abnormal and diagnostic of PAD. ABIs >1.40 indicate noncompressible arteries secondary to vascular calcification.

Other noninvasive tests include segmental pressure measurements, segmental pulse volume recordings, duplex ultrasonography (which combines B-mode imaging and Doppler flow velocity waveform analysis examination), transcutaneous oximetry, and stress testing (usually using a treadmill). Placement of pneumatic cuffs enables assessment of systolic pressure along the legs. The presence of pressure gradients between sequential cuffs provides evidence of the presence and location of hemodynamically significant stenoses. In addition, the amplitude of the pulse volume contour becomes blunted in the presence of significant PAD. Duplex ultrasonography is used to image and detect stenotic lesions in native arteries and bypass grafts.

Treadmill testing allows the physician to assess functional limitations objectively. Decline of the ABI immediately after exercise provides further support for the diagnosis of PAD in patients with equivocal symptoms and findings on examination.

Magnetic resonance angiography (MRA), computed tomographic angiography (CTA), and conventional catheter-based angiography should not be used for routine diagnostic testing but are performed before potential revascularization (**Fig. 302-1**). Each test is useful in defining the anatomy to assist planning for endovascular and surgical revascularization procedures.

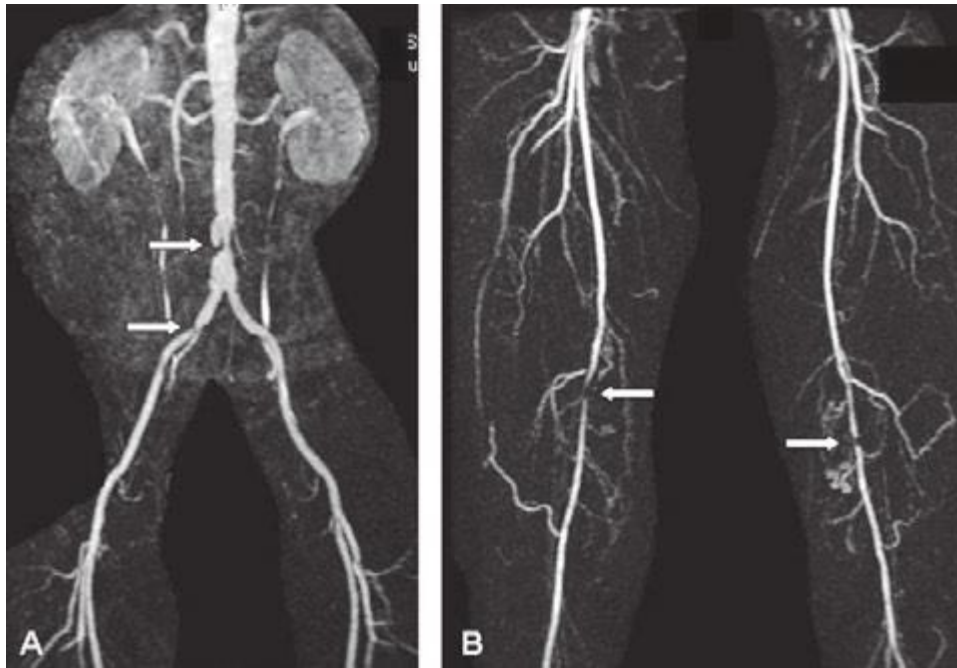


FIGURE 302-1 Magnetic resonance angiography of a patient with intermittent claudication, showing stenoses of the distal abdominal aorta and right iliac common iliac artery (A) and stenoses of the right and left superficial femoral arteries (B). (Courtesy of Dr. Edwin Gravereaux, with permission.)

Prognosis The natural history of patients with PAD is influenced primarily by the extent of coexisting coronary artery and cerebrovascular disease. Approximately one-third to one-half of patients with symptomatic PAD have evidence of coronary artery disease (CAD) based on clinical presentation and electrocardiogram, and over one-half have significant CAD by coronary angiography. Patients with PAD have a 15–30% 5-year mortality rate and a two- to sixfold increased risk of death from coronary heart disease. Mortality rates are highest in those with the most severe PAD. Measurement of ABI is useful for detecting PAD and identifying persons at risk for future atherothrombotic events. The likelihood of symptomatic progression of PAD is lower than the chance of succumbing to CAD. Approximately 75–80% of nondiabetic patients who present with mild to moderate claudication remain symptomatically stable. Deterioration is likely to occur in the remainder, with approximately 1–2% of the group ultimately developing critical limb ischemia each year. Approximately 25–30% of patients with critical limb ischemia undergo amputation within 1 year. The prognosis is worse in patients who continue to smoke cigarettes or have diabetes mellitus.

TREATMENT PERIPHERAL ARTERY DISEASE

Patients with PAD should receive therapies to reduce the risk of associated cardiovascular events, such as myocardial infarction and death, and to improve limb symptoms, prevent progression to critical limb ischemia, and preserve limb viability. Risk factor modification and antiplatelet therapy should be initiated to improve cardiovascular outcomes. The importance of discontinuing cigarette smoking cannot be overemphasized. The physician must assume a major role in this lifestyle modification. Counseling and adjunctive drug therapy with the nicotine patch, bupropion, or varenicline increase smoking cessation rates and reduce recidivism. It is important to control blood pressure in hypertensive patients. Angiotensin-converting enzyme inhibitors may reduce the risk of cardiovascular events in patients with symptomatic PAD. β -Adrenergic blockers do not worsen claudication and may be used to treat hypertension, especially in patients with coexistent CAD. Treatment of hypercholesterolemia with statins is advocated to reduce the risk of myocardial infarction, stroke, and death. The 2013 ACC/AHA Guideline on the Treatment

of Blood Cholesterol to Reduce Atherosclerotic Cardiovascular Risk in Adults recommends high intensity statin treatment in patients with atherosclerotic disorders, including peripheral artery disease. Platelet inhibitors, including aspirin and clopidogrel, reduce the risk of adverse cardiovascular events in patients with atherosclerosis and are recommended for patients with symptomatic PAD, including those with intermittent claudication or critical limb ischemia or prior lower extremity revascularization. Dual antiplatelet therapy with both aspirin and clopidogrel is not more effective than aspirin alone in reducing cardiovascular morbidity and mortality rates in patients with PAD. The anticoagulant warfarin is as effective as antiplatelet therapy in preventing adverse cardiovascular events but causes more major bleeding; therefore, it is not indicated to improve outcomes in patients with chronic PAD.

Therapies for intermittent claudication and critical limb ischemia include supportive measures, medications, nonoperative interventions, and surgery. Supportive measures include meticulous care of the feet, which should be kept clean and protected against excessive drying with moisturizing creams. Well-fitting and protective shoes are advised to reduce trauma. Elastic support hose should be avoided, as it reduces blood flow to the skin. In patients with critical limb ischemia, shock blocks under the head of the bed together with a canopy over the feet may improve perfusion pressure and ameliorate some of the rest pain.

Patients with claudication should be encouraged to exercise regularly and at progressively more strenuous levels. Supervised exercise training programs for 30- to 45-min sessions, three to five times per week for at least 12 weeks, prolong walking distance. Patients also should be advised to walk until nearly maximum claudication discomfort occurs and then rest until the symptoms resolve before resuming ambulation. The beneficial effect of supervised exercise training on walking performance in patients with claudication often is similar to or greater than that realized after a revascularization procedure. Pharmacologic treatment of PAD has not been as successful as the medical treatment of CAD (Chap. 293). In particular, vasodilators as a class have not proved to be beneficial. During exercise, peripheral vasodilation occurs distal to sites of significant arterial stenoses. As a result, perfusion pressure falls, often to levels lower than that generated in the interstitial tissue by the exercising muscle. Drugs such as α -adrenergic blocking agents, calcium

channel antagonists, and other vasodilators have not been shown to be effective in patients with PAD.

Cilostazol, a phosphodiesterase inhibitor with vasodilator and antiplatelet properties, increases claudication distance by 40–60% and improves measures of quality of life. The mechanism of action accounting for its beneficial effects is not known. Pentoxifylline, a substituted xanthine derivative, increases blood flow to the microcirculation and enhances tissue oxygenation. Although several placebo-controlled studies have found that pentoxifylline increases the duration of exercise in patients with claudication, its efficacy has not been confirmed in all clinical trials. Statins and angiotensin-converting enzyme inhibitors appear promising for treatment of intermittent claudication in initial clinical trials, but more studies are needed to confirm the efficacy of each class of drugs. There is no definitive medical therapy for critical limb ischemia, although several studies have suggested that long-term parenteral administration of vasodilator prostaglandins decreases pain and facilitates healing of ulcers. Enthusiasm for therapy with angiogenic growth factors abated when clinical trials of intramuscular gene transfer of DNA encoding vascular endothelial growth factor, fibroblast growth factor, hepatocyte growth factor, or hypoxia-inducible factor 1 α failed to demonstrate improvement in symptoms or outcomes in patients with intermittent claudication or critical limb ischemia. Clinical trials assessing the ability of bone marrow–derived vascular progenitor cells to promote angiogenesis and preserve limb viability in patients with critical limb ischemia are ongoing.

REVASCULARIZATION

Revascularization procedures, including catheter-based and surgical interventions, are usually indicated for patients with disabling, progressive, or severe symptoms of intermittent claudication despite medical therapy and for those with critical limb ischemia. MRA, CTA, or conventional angiography should be performed to assess vascular anatomy in patients who are being considered for revascularization. Nonoperative interventions include percutaneous transluminal angiography (PTA) and stent placement (**Chap. 296e**). PTA and stenting of the iliac artery are associated with higher success rates than are PTA and stenting of the femoral and popliteal arteries. Approximately 90–95% of iliac PTAs are initially successful, and the 3-year patency rate is >75%. Patency rates may be higher if a stent is placed in the iliac artery. The initial success rates for femoral-popliteal PTA and stenting are approximately 80%, with 60% 3-year patency rates. Patency rates are influenced by the severity of pretreatment stenoses; the prognosis of occlusive lesions is worse than that of nonocclusive stenotic lesions. The role of drug-eluting stents and drug-coated balloons in PAD is under investigation.

Several operative procedures are available for treating patients with aortoiliac and femoral-popliteal artery disease. The preferred operative procedure depends on the location and extent of the obstruction(s) and the general medical condition of the patient. Operative procedures for aortoiliac disease include aortobifemoral bypass, axillofemoral bypass, femoro-femoral bypass, and aortoiliac endarterectomy. The most frequently used procedure is the aortobifemoral bypass using knitted Dacron grafts. Immediate graft patency approaches 99%, and 5- and 10-year graft patency rates in survivors are >90% and 80%, respectively. Operative complications include myocardial infarction and stroke, infection of the graft, peripheral embolization, and sexual dysfunction from interruption of autonomic nerves in the pelvis. The operative mortality rate ranges from 1–3%, mostly due to ischemic heart disease.

Operative therapy for femoral-popliteal artery disease includes in situ and reverse autogenous saphenous vein bypass grafts, placement of polytetrafluoroethylene (PTFE) or other synthetic grafts, and thromboendarterectomy. The operative mortality rate ranges from 1–3%. The long-term patency rate depends on the type of graft used, the location of the distal anastomosis, and the patency of runoff vessels beyond the anastomosis. Patency rates of femoral-popliteal saphenous vein bypass grafts approach 90% at 1 year and 70–80% at 5 years. Five-year patency rates of infrapopliteal saphenous vein

bypass grafts are 60–70%. In contrast, 5-year patency rates of infrapopliteal PTFE grafts are <30%.

Preoperative cardiac risk assessment may identify individuals who are especially likely to experience an adverse cardiac event during the perioperative period. Patients with angina, prior myocardial infarction, ventricular ectopy, heart failure, or diabetes are among those at increased risk. Stress testing with treadmill exercise (if feasible), radionuclide myocardial perfusion imaging, or echocardiography permits further stratification of risk in these patients (**Chap. 296e**). Patients with abnormal test results require close supervision and adjunctive management with anti-ischemic medications. β -Adrenergic blockers and statins reduce the risk of postoperative cardiovascular complications. Coronary angiography and coronary artery revascularization compared with optimal medical therapy do not improve outcomes in most patients undergoing peripheral vascular surgery, but cardiac catheterization should be considered in patients with unstable angina and angina refractory to medical therapy as well as those suspected of having left main or three-vessel CAD.

FIBROMUSCULAR DYSPLASIA

Fibromuscular dysplasia is a hyperplastic disorder that affects medium-size and small arteries. It occurs predominantly in females and usually involves the renal and carotid arteries but can affect extremity vessels such as the iliac and subclavian arteries. The histologic classification includes intimal fibroplasia (also classified as focal), medial dysplasia (multifocal), and adventitial hyperplasia. Medial dysplasia is subdivided into medial fibroplasia, perimedial fibroplasia, and medial hyperplasia. Medial fibroplasia is the most common type and is characterized by alternating areas of thinned media and fibromuscular ridges. The internal elastic lamina usually is preserved. The iliac arteries are the limb arteries most likely to be affected by fibromuscular dysplasia. It is identified angiographically by a “string of beads” appearance caused by thickened fibromuscular ridges contiguous with thin, less-involved portions of the arterial wall, which is typical of medial fibroplasia. When limb vessels are involved, clinical manifestations are similar to those for atherosclerosis, including claudication and rest pain. PTA and surgical reconstruction have been beneficial in patients with debilitating symptoms or threatened limbs.

THROMBOANGIITIS OBLITERANS

Thromboangiitis obliterans (Buerger’s disease) is an inflammatory occlusive vascular disorder involving small and medium-size arteries and veins in the distal upper and lower extremities. Cerebral, visceral, and coronary vessels may be affected rarely. This disorder develops most frequently in men <40 years of age. The prevalence is higher in Asians and individuals of Eastern European descent. Although the cause of thromboangiitis obliterans is not known, there is a definite relationship to cigarette smoking in patients with this disorder.

In the initial stages of thromboangiitis obliterans, polymorphonuclear leukocytes infiltrate the walls of the small and medium-size arteries and veins. The internal elastic lamina is preserved, and a cellular, inflammatory thrombus develops in the vascular lumen. As the disease progresses, mononuclear cells, fibroblasts, and giant cells replace the neutrophils. Later stages are characterized by perivascular fibrosis, organized thrombus, and recanalization.

The clinical features of thromboangiitis obliterans often include a triad of claudication of the affected extremity, Raynaud’s phenomenon, and migratory superficial vein thrombophlebitis. Claudication usually is confined to the calves and feet or the forearms and hands because this disorder primarily affects distal vessels. In the presence of severe digital ischemia, trophic nail changes, painful ulcerations, and gangrene may develop at the tips of the fingers or toes. The physical examination shows normal brachial and popliteal pulses but reduced or absent radial, ulnar, and/or tibial pulses. MRA, CTA, and conventional arteriography are helpful in making the diagnosis. Smooth, tapering segmental lesions in the distal vessels are characteristic, as are

1646 collateral vessels at sites of vascular occlusion. Proximal atherosclerotic disease is usually absent. The diagnosis can be confirmed by excisional biopsy and pathologic examination of an involved vessel.

There is no specific treatment except abstention from tobacco. The prognosis is worse in individuals who continue to smoke, but results are discouraging even in those who stop smoking. Arterial bypass of the larger vessels may be used in selected instances, as well as local debridement, depending on the symptoms and severity of ischemia. Antibiotics may be useful; anticoagulants and glucocorticoids are not helpful. If these measures fail, amputation may be required.

VASCULITIS

Other vasculitides may affect the arteries that supply the upper and lower extremities. **Takayasu's arteritis and giant cell (temporal) arteritis are discussed in Chap. 385.**

ACUTE LIMB ISCHEMIA

Acute limb ischemia occurs when arterial occlusion results in the sudden cessation of blood flow to an extremity. The severity of ischemia and the viability of the extremity depend on the location and extent of the occlusion and the presence and subsequent development of collateral blood vessels. Principal causes of acute arterial occlusion include embolism, thrombus in situ, arterial dissection, and trauma.

The most common sources of arterial emboli are the heart, aorta, and large arteries. Cardiac disorders that cause thromboembolism include atrial fibrillation, both chronic and paroxysmal; acute myocardial infarction; ventricular aneurysm; cardiomyopathy; infectious and marantic endocarditis; thrombi associated with prosthetic heart valves; and atrial myxoma. Emboli to the distal vessels may also originate from proximal sites of atherosclerosis and aneurysms of the aorta and large vessels. Less frequently, an arterial occlusion results paradoxically from a venous thrombus that has entered the systemic circulation via a patent foramen ovale or another septal defect. Arterial emboli tend to lodge at vessel bifurcations because the vessel caliber decreases at those sites; in the lower extremities, emboli lodge most frequently in the femoral artery, followed by the iliac artery, aorta, and popliteal and tibioperoneal arteries.

Acute arterial thrombosis in situ occurs most frequently in atherosclerotic vessels at the site of an atherosclerotic plaque or aneurysm and in arterial bypass grafts. Trauma to an artery may disrupt continuity of blood flow and cause acute limb ischemia via formation of an acute arterial thrombus or by disruption of an artery's integrity and extravasation of blood. Arterial occlusion may complicate arterial punctures and placement of catheters; it also may result from arterial dissection if the intimal flap obstructs the artery. Less common causes include thoracic outlet compression syndrome, which causes subclavian artery occlusion, and entrapment of the popliteal artery by abnormal placement of the medial head of the gastrocnemius muscle. Polycythemia and hypercoagulable disorders (**Chaps. 131 and 141**) are also associated with acute arterial thrombosis.

CLINICAL FEATURES

The symptoms of an acute arterial occlusion depend on the location, duration, and severity of the obstruction. Often, severe pain, paresthesia, numbness, and coldness develop in the involved extremity within 1 hour. Paralysis may occur with severe and persistent ischemia. Physical findings include loss of pulses distal to the occlusion, cyanosis or pallor, mottling, decreased skin temperature, muscle stiffening, loss of sensation, weakness, and/or absent deep tendon reflexes. If acute arterial occlusion occurs in the presence of an adequate collateral circulation, as is often the case in acute graft occlusion, the symptoms and findings may be less impressive. In this situation, the patient complains about an abrupt decrease in the distance walked before claudication occurs or of modest pain and paresthesia. Pallor and coolness are evident, but sensory and motor functions generally are preserved. The diagnosis of acute limb ischemia is usually apparent from the clinical presentation. In most circumstances, MRA, CTA, or catheter-based arteriography is used to confirm the diagnosis and demonstrate the location and extent of arterial occlusion.

TREATMENT ACUTE LIMB ISCHEMIA

Once the diagnosis is made, the patient should be anticoagulated with intravenous heparin to prevent propagation of the clot. In cases of severe ischemia of recent onset, particularly when limb viability is jeopardized, immediate intervention to ensure reperfusion is indicated. Catheter-directed thrombolysis/thrombectomy, surgical thromboembolectomy, and arterial bypass procedures are used to restore blood flow to the ischemic extremity promptly, particularly when a large proximal vessel is occluded.

Intraarterial thrombolytic therapy with recombinant tissue plasminogen activator, reteplase, or tenecteplase is most effective when acute arterial occlusion is recent (<2 weeks) and caused by a thrombus in an atherosclerotic vessel, arterial bypass graft, or occluded stent. Thrombolytic therapy is also indicated when the patient's overall condition contraindicates surgical intervention or when smaller distal vessels are occluded, thus preventing surgical access. Meticulous observation for hemorrhagic complications is required during intraarterial thrombolytic therapy. Another endovascular approach to thrombus removal is percutaneous mechanical thrombectomy using devices that employ hydrodynamic forces or rotating baskets to fragment and remove the clot. These treatments may be used alone but usually are used in conjunction with pharmacologic thrombolysis. Surgical revascularization is preferred when restoration of blood flow must occur within 24 h to prevent limb loss or when symptoms of occlusion have been present for more than 2 weeks. Amputation is performed when the limb is not viable, as characterized by loss of sensation, paralysis, and the absence of Doppler-detected blood flow in both arteries and veins.

If the limb is not in jeopardy, a more conservative approach that includes observation and administration of anticoagulants may be taken. Anticoagulation prevents recurrent embolism and reduces the likelihood of thrombus propagation; it can be initiated with intravenous heparin and followed by oral warfarin. Recommended doses are the same as those used for deep vein thrombosis (**Chap. 300**). Emboli resulting from infective endocarditis, the presence of prosthetic heart valves, or atrial myxoma often require surgical intervention to remove the cause.

ATHEROEMBOLISM

Atheroembolism is another cause of limb ischemia. In this condition, multiple small deposits of fibrin, platelets, and cholesterol debris embolize from proximal atherosclerotic lesions or aneurysmal sites. Large protruding aortic atheromas are a source of emboli that may lead to limb ischemia, as well as stroke and renal insufficiency. Atheroembolism may occur after intraarterial procedures. Since atheroemboli to limbs tend to lodge in the small vessels of the muscle and skin and may not occlude the large vessels, distal pulses usually remain palpable. Patients complain of acute pain and tenderness at the site of embolization. Digital vascular occlusion may result in ischemia and the "blue toe" syndrome; digital necrosis and gangrene may develop (**Fig. 302-2**). Localized areas of tenderness, pallor, and livedo reticularis (see below) occur at sites of emboli. Skin or muscle biopsy may demonstrate cholesterol crystals.

Ischemia resulting from atheroemboli is notoriously difficult to treat. Usually neither surgical revascularization procedures nor thrombolytic therapy is helpful because of the multiplicity, composition, and distal location of the emboli. There is limited evidence that antithrombotic therapy with platelet inhibitors or anticoagulants prevents atheroembolism. Statins may stabilize plaque and potentially reduce the risk of atheroembolism. Surgical intervention to remove or bypass the atherosclerotic vessel or aneurysm that causes the recurrent atheroemboli may be necessary.

THORACIC OUTLET COMPRESSION SYNDROME

This is a symptom complex resulting from compression of the neurovascular bundle (artery, vein, or nerves) at the thoracic outlet as it courses through the neck and shoulder. Cervical ribs, abnormalities of



FIGURE 302-2 Atheroembolism causing cyanotic discoloration and impending necrosis of the toes (“blue toe” syndrome).

the scalenus anticus muscle, proximity of the clavicle to the first rib, or abnormal insertion of the pectoralis minor muscle may compress the subclavian artery, subclavian vein, and brachial plexus as these structures pass from the thorax to the arm. Depending on the structures affected, thoracic outlet compression syndrome is divided into arterial, venous, and neurogenic forms. Patients with neurogenic thoracic outlet compression may develop shoulder and arm pain, weakness, and paresthesias. Patients with arterial compression may experience claudication, Raynaud’s phenomenon, and even ischemic tissue loss and gangrene. Venous compression may cause thrombosis of the subclavian and axillary veins; this is often associated with effort and is referred to as *Paget-Schroetter syndrome*.

APPROACH TO THE PATIENT

Thoracic Outlet Compression Syndrome

Examination of a patient with arterial thoracic outlet compression syndrome is often normal unless provocative maneuvers are performed. Occasionally, distal pulses are decreased or absent and digital cyanosis and ischemia may be evident.

Several maneuvers that support the diagnosis of arterial thoracic outlet compression syndrome may be used to precipitate symptoms, cause a subclavian artery bruit, and diminish arm pulses. These maneuvers include the abduction and external rotation test, in which the affected arm is abducted by 90° and the shoulder is externally rotated; the scalene maneuver (extension of the neck and rotation of the head to the side of the symptoms); the costoclavicular maneuver (posterior rotation of shoulders); and the hyperabduction maneuver (raising the arm 180°). A chest x-ray will indicate the presence of cervical ribs. Duplex ultrasonography, MRA, and contrast angiography can be performed during provocative maneuvers to demonstrate thoracic outlet compression of the subclavian artery. Neurophysiologic tests such as the electromyogram, nerve conduction studies, and somatosensory evoked potentials may be abnormal if the brachial plexus is involved, but the diagnosis of neurogenic thoracic outlet syndrome is not necessarily excluded if these tests are normal owing to their low sensitivity.

Most patients can be managed conservatively. They should be advised to avoid the positions that cause symptoms. Many patients benefit from shoulder girdle exercises. Surgical procedures such as removal of the first rib and resection of the scalenus anticus muscle are necessary occasionally for relief of symptoms or treatment of ischemia.

POPLITEAL ARTERY ENTRAPMENT

Popliteal artery entrapment typically affects young athletic men and women when the gastrocnemius or popliteus muscle compresses the

popliteal artery and causes intermittent claudication. Thrombosis, embolism, or popliteal artery aneurysm may occur. The pulse examination may be normal unless provocative maneuvers such as ankle dorsiflexion and plantar flexion are performed. The diagnosis is confirmed by duplex ultrasound, CTA, MRA, or conventional angiography. Treatment involves surgical release of the popliteal artery or vascular reconstruction.

POPLITEAL ARTERY ANEURYSM

Popliteal artery aneurysms are the most common peripheral artery aneurysms. Approximately 50% are bilateral. Patients with popliteal artery aneurysms often have aneurysms of other arteries, especially the aorta. The most common clinical presentation is limb ischemia secondary to thrombosis or embolism. Rupture occurs less frequently. Other complications include compression of the adjacent popliteal vein or peroneal nerve. Popliteal artery aneurysm can be detected by palpation and confirmed by duplex ultrasonography. Repair is indicated for symptomatic aneurysms or when the diameter exceeds 2–3 cm, owing to the risk of thrombosis, embolism, or rupture.

ARTERIOVENOUS FISTULA

Abnormal communications between an artery and a vein, bypassing the capillary bed, may be congenital or acquired. Congenital arteriovenous fistulas are a result of persistent embryonic vessels that fail to differentiate into arteries and veins; they may be associated with birthmarks, can be located in almost any organ of the body, and frequently occur in the extremities. Acquired arteriovenous fistulas either are created to provide vascular access for hemodialysis or occur as a result of a penetrating injury such as a gunshot or knife wound or as complications of arterial catheterization or surgical dissection. An uncommon cause of arteriovenous fistula is rupture of an arterial aneurysm into a vein.

The clinical features depend on the location and size of the fistula. Frequently, a pulsatile mass is palpable, and a thrill and a bruit lasting throughout systole and diastole are present over the fistula. With long-standing fistulas, clinical manifestations of chronic venous insufficiency, including peripheral edema; large, tortuous varicose veins; and stasis pigmentation become apparent because of the high venous pressure. Evidence of ischemia may occur in the distal portion of the extremity. Skin temperature is higher over the arteriovenous fistula. Large arteriovenous fistulas may result in an increased cardiac output with consequent cardiomegaly and high-output heart failure ([Chap. 279](#)).

The diagnosis is often evident from the physical examination. Compression of a large arteriovenous fistula may cause reflex slowing of the heart rate (Nicoladoni-Branham sign). Duplex ultrasonography may detect an arteriovenous fistula, especially one that affects the femoral artery and vein at the site of catheter access. CTA and conventional angiography can confirm the diagnosis and are useful in demonstrating the site and size of the arteriovenous fistula.

Management of arteriovenous fistulas may involve surgery, radiotherapy, or embolization. Congenital arteriovenous fistulas are often difficult to treat because the communications may be numerous and extensive, and new communications frequently develop after ligation of the most obvious ones. Many of these lesions are best treated conservatively using elastic support hose to reduce the consequences of venous hypertension. Occasionally, embolization with autologous material, such as fat or muscle, or with hemostatic agents, such as gelatin sponges or silicon spheres, is used to obliterate the fistula. Acquired arteriovenous fistulas are usually amenable to surgical treatment that involves division or excision of the fistula. Occasionally, autogenous or synthetic grafting is necessary to reestablish continuity of the artery and vein.

RAYNAUD’S PHENOMENON

Raynaud’s phenomenon is characterized by episodic digital ischemia, manifested clinically by the sequential development of digital blanching, cyanosis, and rubor of the fingers or toes after cold exposure



FIGURE 302-3 Vascular diseases associated with temperature: (A) Raynaud's phenomenon; (B) acrocyanosis; (C) livedo reticularis; (D) pernio; (E) erythromelalgia; and (F) frostbite.

and subsequent rewarming. Emotional stress may also precipitate Raynaud's phenomenon. The color changes are usually well demarcated and are confined to the fingers or toes. Typically, one or more digits will appear white when the patient is exposed to a cold environment or touches a cold object (Fig. 302-3A). The blanching, or pallor, represents the ischemic phase of the phenomenon and results from vasospasm of digital arteries. During the ischemic phase, capillaries and venules dilate, and cyanosis results from the deoxygenated blood that is present in these vessels. A sensation of cold or numbness or paresthesia of the digits often accompanies the phases of pallor and cyanosis.

With rewarming, the digital vasospasm resolves, and blood flow into the dilated arterioles and capillaries increases dramatically. This "reactive hyperemia" imparts a bright red color to the digits. In addition to rubor and warmth, patients often experience a throbbing, painful sensation during the hyperemic phase. Although the triphasic color response is typical of Raynaud's phenomenon, some patients may develop only pallor and cyanosis; others may experience only cyanosis.

Raynaud's phenomenon is broadly separated into two categories: idiopathic, termed primary Raynaud's phenomenon, and secondary Raynaud's phenomenon, which is associated with other disease states or known causes of vasospasm (Table 302-1).

Primary Raynaud's Phenomenon This appellation is applied when the secondary causes of Raynaud's phenomenon have been excluded. Over 50% of patients with Raynaud's phenomenon have the primary form. Women are affected about five times more often than men, and the age of presentation is usually between 20 and 40 years. The fingers are involved more frequently than the toes. Initial episodes may involve only one or two fingertips, but subsequent attacks may involve

the entire finger and may include all the fingers. The toes are affected in 40% of patients. Although vasospasm of the toes usually occurs in patients with symptoms in the fingers, it may happen alone. Rarely, the earlobes, the tip of the nose, and the penis are involved. Raynaud's phenomenon occurs frequently in patients who also have migraine headaches or variant angina. These associations suggest that there may be a common predisposing cause for the vasospasm.

Results of physical examination are often entirely normal; the radial, ulnar, and pedal pulses are normal. The fingers and toes may

TABLE 302-1 CLASSIFICATION OF RAYNAUD'S PHENOMENON

Primary or idiopathic Raynaud's phenomenon
Secondary Raynaud's phenomenon
Collagen vascular diseases: scleroderma, systemic lupus erythematosus, rheumatoid arthritis, dermatomyositis, polymyositis, mixed connective tissue disease, Sjögren's syndrome
Arterial occlusive diseases: atherosclerosis of the extremities, thromboangiitis obliterans, acute arterial occlusion, thoracic outlet syndrome
Pulmonary hypertension
Neurologic disorders: intervertebral disk disease, syringomyelia, spinal cord tumors, stroke, poliomyelitis, carpal tunnel syndrome, complex regional pain syndrome
Blood dyscrasias: cold agglutinins, cryoglobulinemia, cryofibrinogenemia, myeloproliferative disorders, lymphoplasmacytic lymphoma
Trauma: vibration injury, hammer hand syndrome, electric shock, cold injury, typing, piano playing
Drugs and toxins: ergot derivatives, methysergide, β -adrenergic receptor blockers, bleomycin, vinblastine, cisplatin, gemcitabine, vinyl chloride

be cool between attacks and may perspire excessively. Thickening and tightening of the digital subcutaneous tissue (*sclerodactyly*) develop in 10% of patients. Angiography of the digits for diagnostic purposes is not indicated.

In general, patients with primary Raynaud's disease have milder clinical manifestations. Fewer than 1% of these patients lose a part of a digit. After the diagnosis is made, the disease improves spontaneously in approximately 15% of patients and progresses in about 30%.

Secondary Causes of Raynaud's Phenomenon Raynaud's phenomenon occurs in 80–90% of patients with systemic sclerosis (scleroderma) and is the presenting symptom in 30% (**Chap. 382**). It may be the only symptom of scleroderma for many years. Abnormalities of the digital vessels may contribute to the development of Raynaud's phenomenon in this disorder. Ischemic fingertip ulcers may develop and progress to gangrene and autoamputation. About 20% of patients with systemic lupus erythematosus (SLE) have Raynaud's phenomenon (**Chap. 378**). Occasionally, persistent digital ischemia develops and may result in ulcers or gangrene. In most severe cases, the small vessels are occluded by a proliferative endarteritis. Raynaud's phenomenon occurs in about 30% of patients with dermatomyositis or polymyositis (**Chap. 388**). It frequently develops in patients with rheumatoid arthritis and may be related to the intimal proliferation that occurs in the digital arteries.

Atherosclerosis of the extremities is a common cause of Raynaud's phenomenon in men >50 years. Thromboangiitis obliterans is an uncommon cause of Raynaud's phenomenon but should be considered in young men, particularly those who are cigarette smokers. The development of cold-induced pallor in these disorders may be confined to one or two digits of the involved extremity. Occasionally, Raynaud's phenomenon may follow acute occlusion of large and medium-size arteries by a thrombus or embolus. Embolization of atheroembolic debris may cause digital ischemia. The latter situation often involves one or two digits and should not be confused with Raynaud's phenomenon. In patients with thoracic outlet compression syndrome, Raynaud's phenomenon may result from diminished intravascular pressure, stimulation of sympathetic fibers in the brachial plexus, or a combination of both. Raynaud's phenomenon occurs in patients with primary pulmonary hypertension (**Chap. 304**); this is more than coincidental and may reflect a neurohumoral abnormality that affects both the pulmonary and digital circulations.

A variety of blood dyscrasias may be associated with Raynaud's phenomenon. Cold-induced precipitation of plasma proteins, hyperviscosity, and aggregation of red cells and platelets may occur in patients with cold agglutinins, cryoglobulinemia, or cryofibrinogenemia. Hyperviscosity syndromes that accompany myeloproliferative disorders and lymphoplasmacytic lymphoma (Waldenström's macroglobulinemia) should also be considered in the initial evaluation of patients with Raynaud's phenomenon.

Raynaud's phenomenon occurs often in patients whose vocations require the use of vibrating hand tools, such as chain saws or jackhammers. The frequency of Raynaud's phenomenon also seems to be increased in pianists and keyboard operators. Electric shock injury to the hands or frostbite may lead to the later development of Raynaud's phenomenon.

Several drugs have been causally implicated in Raynaud's phenomenon. They include ergot preparations, methysergide, β -adrenergic receptor antagonists, and the chemotherapeutic agents bleomycin, vinblastine, cisplatin, and gemcitabine.

TREATMENT RAYNAUD'S PHENOMENON

Most patients with Raynaud's phenomenon experience only mild and infrequent episodes. These patients need reassurance and should be instructed to dress warmly and avoid unnecessary cold exposure. In addition to gloves and mittens, patients should protect the trunk, head, and feet with warm clothing to prevent cold-induced reflex vasoconstriction. Tobacco use is contraindicated.

Drug treatment should be reserved for severe cases. Dihydropyridine calcium channel antagonists such as nifedipine, isradipine, felodipine, and amlodipine decrease the frequency and severity of Raynaud's phenomenon. Diltiazem may be considered but is less effective. The postsynaptic α_1 -adrenergic antagonist prazosin has been used with favorable responses; doxazosin and terazosin may also be effective. Phosphodiesterase type 5 inhibitors such as sildenafil and tadalafil may improve symptoms in patients with secondary Raynaud's phenomenon, as occurs with systemic sclerosis. Digital sympathectomy is helpful in some patients who are unresponsive to medical therapy.

ACROCYNANOSIS

In this condition, there is arterial vasoconstriction and secondary dilation of the capillaries and venules with resulting persistent cyanosis of the hands and, less frequently, the feet. Cyanosis may be intensified by exposure to a cold environment. Acrocyanosis may be categorized as primary or secondary to an underlying condition. In primary acrocyanosis, women are affected much more frequently than men, and the age of onset is usually <30 years. Generally, patients are asymptomatic but seek medical attention because of the discoloration. The prognosis is favorable, and pain, ulcers, and gangrene do not occur. Examination reveals normal pulses, peripheral cyanosis, and moist palms (**Fig. 302-3B**). Trophic skin changes and ulcerations do not occur. The disorder can be distinguished from Raynaud's phenomenon because it is persistent and not episodic, the discoloration extends proximally from the digits, and blanching does not occur. Ischemia secondary to arterial occlusive disease can usually be excluded by the presence of normal pulses. Central cyanosis and decreased arterial oxygen saturation are not present. Patients should be reassured and advised to dress warmly and avoid cold exposure. Pharmacologic intervention is not indicated.

Secondary acrocyanosis may result from hypoxemia, vasopressor medications, connective tissue diseases, atheroembolism, antiphospholipid antibodies, cold agglutinins, or cryoglobulins and is associated with anorexia nervosa and postural orthostatic tachycardia syndrome. Treatment should be directed at the underlying disorder.

LIVEDO RETICULARIS

In this condition, localized areas of the extremities develop a mottled or rete (netlike) appearance of reddish to blue discoloration (**Fig. 302-3C**). The mottled appearance may be more prominent after cold exposure. There are primary and secondary forms of livedo reticularis. The primary, or idiopathic, form of this disorder may be benign or associated with ulcerations. The benign form occurs more frequently in women than in men, and the most common age of onset is the third decade. Patients with the benign form are usually asymptomatic and seek attention for cosmetic reasons. These patients should be reassured and advised to avoid cold environments. No drug treatment is indicated. Primary livedo reticularis with ulceration is also called *atrophia blanche en plaque*. The ulcers are painful and may take months to heal. Secondary livedo reticularis can occur with atheroembolism (see above), SLE and other vasculitides, anticardiolipin antibodies, hyperviscosity, cryoglobulinemia, and Sneddon's syndrome (ischemic stroke and livedo reticularis). Rarely, skin ulcerations develop.

PERNIO (CHILBLAINS)

Pernio is a vasculitic disorder associated with exposure to cold; acute forms have been described. Raised erythematous lesions develop on the lower part of the legs and feet in cold weather (**Fig. 302-3D**). They are associated with pruritus and a burning sensation, and they may blister and ulcerate. Pathologic examination demonstrates angitis characterized by intimal proliferation and perivascular infiltration of mononuclear and polymorphonuclear leukocytes. Giant cells may be present in the subcutaneous tissue. Patients should avoid exposure to cold, and ulcers should be kept clean and protected with sterile dressings. Sympatholytic drugs and dihydropyridine calcium channel antagonists may be effective in some patients.

This disorder is characterized by burning pain and erythema of the extremities (Fig. 302-3E). The feet are involved more frequently than the hands, and males are affected more frequently than females. Erythromelalgia may occur at any age but is most common in middle age. It may be primary (also termed erythermalgia) or secondary. Mutations in the *SCN9A* gene, which encodes the Nav1.7 voltage-gated sodium channel expressed in sensory and sympathetic nerves, has been described in inherited forms of erythromelalgia. The most common causes of secondary erythromelalgia are myeloproliferative disorders such as polycythemia vera and essential thrombocytosis. Less common causes include drugs, such as calcium channel blockers, bromocriptine, and pergolide; neuropathies; connective tissue diseases such as SLE; and paraneoplastic syndromes. Patients complain of burning in the extremities that is precipitated by exposure to a warm environment and aggravated by a dependent position. The symptoms are relieved by exposing the affected area to cool air or water or by elevation. Erythromelalgia can be distinguished from ischemia secondary to peripheral arterial disorders because the peripheral pulses are present. There is no specific treatment; aspirin may produce relief in patients with erythromelalgia secondary to myeloproliferative disease. Treatment of associated disorders in secondary erythromelalgia may be helpful.

FROSTBITE

In this condition, tissue damage results from severe environmental cold exposure or from direct contact with a very cold object. Tissue injury results from both freezing and vasoconstriction. Frostbite usually affects the distal aspects of the extremities or exposed parts of the face, such as the ears, nose, chin, and cheeks. Superficial frostbite involves the skin and subcutaneous tissue. Patients experience pain or paresthesia, and the skin appears white and waxy. After rewarming, there is cyanosis and erythema, wheal-and-flare formation, edema, and superficial blisters. Deep frostbite involves muscle, nerves, and deeper blood vessels. It may result in edema of the hand or foot, vesicles and bullae, tissue necrosis, and gangrene (Fig. 302-3F).

Initial treatment is rewarming, performed in an environment where reexposure to freezing conditions will not occur. Rewarming is accomplished by immersion of the affected part in a water bath at temperatures of 40°–44°C (104°–111°F). Massage, application of ice water, and extreme heat are contraindicated. The injured area should be cleansed with soap or antiseptic, and sterile dressings should be applied. Analgesics are often required during rewarming. Antibiotics are used if there is evidence of infection. The efficacy of sympathetic blocking drugs is not established. After recovery, the affected extremity may exhibit increased sensitivity to cold.

VENOUS ANATOMY

Veins in the extremities can be broadly classified as either superficial or deep. The superficial veins are located between the skin and deep fascia. In the legs, these include the great and small saphenous veins and their tributaries. The great saphenous vein is the longest vein in the body. It originates on the medial side of the foot and ascends anterior to the medial malleolus and then along the medial side of the calf and thigh, and drains into the common femoral vein. The small saphenous vein originates on the dorsolateral aspect of the foot, ascends posterior to the lateral malleolus and along the posterolateral aspect of the calf, and drains into the popliteal vein. The deep veins of the leg accompany the major arteries. There are usually paired peroneal, anterior tibial, and posterior tibial veins in the calf, which converge to form the popliteal vein. Soleal tributary veins drain into the posterior tibial or peroneal veins, and gastrocnemius tributary veins drain into the popliteal vein. The popliteal vein ascends in the thigh as the femoral vein. The confluence of the femoral vein and deep femoral vein form the common femoral vein, which ascends in the pelvis as the external iliac and then common iliac vein, which converges with the contralateral common iliac vein at the inferior vena cava. Perforating veins connect the superficial and deep systems in the legs at multiple locations, normally allowing blood to flow from the superficial to deep veins. In the arms, the superficial veins include the basilic, cephalic, and median cubital veins and their tributaries. The basilic and cephalic veins course along the medial and lateral aspects of the arm, respectively, and these are connected via the median cubital vein in the antecubital fossa. The deep veins of the arms accompany the major arteries and include the radial, ulnar, brachial, axillary, and subclavian veins. The subclavian vein converges with the internal jugular vein to form the brachiocephalic vein, which joins the contralateral brachiocephalic vein to form the superior vena cava. Bicuspid valves are present throughout the venous system to direct the flow of venous blood centrally.

Pathophysiology of Chronic Venous Disease *Varicose veins* are dilated, bulging, tortuous superficial veins, measuring at least 3 mm in diameter. The smaller and less tortuous reticular veins are dilated intradermal veins, which appear blue-green, measure 1 to 3 mm in diameter, and do not protrude from the skin surface. Telangiectasias, or spider veins, are small, dilated veins, less than 1 mm in diameter, located near the skin surface, and form blue, purple, or red linear, branching, or spider-web patterns.

Varicose veins can be categorized as primary or secondary. Primary varicose veins originate in the superficial system and result from defective structure and function of the valves of the saphenous veins, intrinsic weakness of the vein wall, and high intraluminal pressure. Approximately one-half of these patients have a family history of varicose veins. Other factors associated with primary varicose veins include aging, pregnancy, hormonal therapy, obesity, and prolonged standing. Secondary varicose veins result from venous hypertension, associated with deep venous insufficiency or deep venous obstruction, and incompetent perforating veins that cause enlargement of superficial veins. Arteriovenous fistulas also cause varicose veins in the affected limb.

Chronic venous insufficiency is a consequence of incompetent veins in which there is venous hypertension and extravasation of fluid and blood elements into the tissue of the limb. It may occur in patients with varicose veins but usually is caused by disease in the deep veins. It also is categorized as primary or secondary. Primary deep venous insufficiency is a consequence of an intrinsic structural or functional abnormality in the vein wall or venous valves leading to valvular reflux. Secondary deep venous insufficiency is caused by obstruction and/or valvular incompetence from previous deep vein thrombosis (Chap. 300). Deep venous insufficiency occurs following deep vein thrombosis, as the delicate valve leaflets become thickened and contracted and can no longer prevent retrograde flow of blood and the vein itself becomes rigid and thick walled. Although most veins recanalize after an episode of thrombosis, the large proximal veins may remain occluded. Secondary incompetence develops in distal valves because high pressures distend the vein and separate the leaflets. Other

303 Chronic Venous Disease and Lymphedema

Mark A. Creager, Joseph Loscalzo

CHRONIC VENOUS DISEASE

Chronic venous diseases range from telangiectasias and reticular veins, to varicose veins, to chronic venous insufficiency with edema, skin changes, and ulceration. This section of the chapter will focus on identification and treatment of varicose veins and chronic venous insufficiency, since these problems are encountered frequently by the internist. The estimated prevalence of varicose veins in the United States is approximately 15% in men and 30% in women. Chronic venous insufficiency with edema affects approximately 7.5% of men and 5% of women, and the prevalence increases with age ranging from 2% among those less than 50 years of age to 10% of those 70 years of age. Approximately 20% of patients with chronic venous insufficiency develop venous ulcers.

causes of secondary deep venous insufficiency include May-Thurner syndrome, where the left iliac vein is occluded or stenosed by extrinsic compression from the overlapping right common iliac artery; arteriovenous fistulas resulting in increased venous pressure; congenital deep vein agenesis or hypoplasia; and venous malformations as may occur in Klippel-Trénaunay-Weber and Parkes-Weber syndromes.

Clinical Presentation Patients with venous varicosities are often asymptomatic but still concerned about the cosmetic appearance of their legs. Superficial venous thrombosis may be a recurring problem, and, rarely, a varicosity ruptures and bleeds. Symptoms in patients with varicose veins or venous insufficiency, when they occur, include a dull ache, throbbing or heaviness, or pressure sensation in the legs typically after prolonged standing; these symptoms usually are relieved with leg elevation. Additional symptoms may include cramping, burning, pruritus, leg swelling, and skin ulceration.

The legs are examined in both the supine and standing positions. Visual inspection and palpation of the legs in the standing position confirm the presence of varicose veins. The location and extent of the varicose veins should be noted. Edema, stasis dermatitis, and skin ulceration near the ankle may be present if there is superficial venous insufficiency and venous hypertension. Findings of deep venous insufficiency include increased leg circumference, venous varicosities, edema, and skin changes. The edema, which is usually pitting, may be confined to the ankles, extend above the ankles to the knees, or involve the thighs in severe cases. Over time, the edema may become less pitting and more indurated. Dermatologic findings associated with venous stasis include hyperpigmentation, erythema, eczema, lipodermatosclerosis, *atrophie blanche*, and a phlebectasia corona. Lipodermatosclerosis is the combination of induration, hemosiderin deposition, and inflammation, and typically occurs in the lower part of the leg just above the ankle. *Atrophie blanche* is a white patch of scar tissue, often with focal telangiectasias and a hyperpigmented border; it usually develops near the medial malleolus. A phlebectasia corona is a fan-shaped pattern of intradermal veins near the ankle or on the foot. Skin ulceration may occur near the medial and lateral malleoli. A venous ulcer is often shallow and characterized by an irregular border, a base of granulation tissue, and the presence of exudate (Fig. 303-1).

Bedside maneuvers can be used to distinguish primary varicose veins from secondary varicose veins caused by deep venous insufficiency. With the contemporary use of venous ultrasound (see below), however, these maneuvers are employed infrequently. The Brodie-Trendelenburg



FIGURE 303-1 Venous insufficiency with active venous ulcer near the medial malleolus. (Courtesy of Dr. Steven Dean, with permission.)

test is used to determine whether varicose veins are secondary to deep venous insufficiency. As the patient is lying supine, the leg is elevated and the veins allowed to empty. Then, a tourniquet is placed on the proximal part of the thigh and the patient is asked to stand. Filling of the varicose veins within 30 s indicates that the varicose veins are caused by deep venous insufficiency and incompetent perforating veins. Primary varicose veins with superficial venous insufficiency are the likely diagnosis if venous refilling occurs promptly after tourniquet removal. The Perthes test assesses the possibility of deep venous obstruction. A tourniquet is placed on the midthigh after the patient has stood, and the varicose veins are filled. The patient is then instructed to walk for 5 min. A patent deep venous system and competent perforating veins enable the superficial veins below the tourniquet to collapse. Deep venous obstruction is likely to be present if the superficial veins distend further with walking.

Differential Diagnosis The duration of leg edema helps to distinguish chronic venous insufficiency from acute deep vein thrombosis. Lymphedema, as discussed later in this chapter, is often confused with chronic venous insufficiency, and both may occur together. Other disorders that cause leg swelling should be considered and excluded when evaluating a patient with presumed venous insufficiency. Bilateral leg swelling occurs in patients with congestive heart failure, hypoalbuminemia secondary to nephrotic syndrome or severe hepatic disease, myxedema caused by hypothyroidism or pretibial myxedema associated with Graves' disease, and with drugs such as dihydropyridine calcium channel blockers and thiazolidinediones. Unilateral causes of leg swelling also include ruptured leg muscles, hematomas secondary to trauma, and popliteal cysts. Cellulitis may cause erythema and swelling of the affected limb. Leg ulcers may be caused by severe peripheral artery disease and critical limb ischemia; neuropathies, particularly those associated with diabetes; and less commonly, skin cancer, vasculitis, or rarely as a complication of hydroxyurea. The location and characteristics of venous ulcers help to differentiate these from other causes.

Classification of Chronic Venous Disease The CEAP (clinical, etiologic, anatomic, pathophysiologic) classification schema incorporates the range of symptoms and signs of chronic venous disease to characterize its severity. It also broadly categorizes the etiology as congenital, primary, or secondary; identifies the affected veins as superficial, deep, or perforating; and characterizes the pathophysiology as reflux, obstruction, both, or neither (Table 303-1).

Diagnostic Testing The principal diagnostic test to evaluate patients with chronic venous disease is venous duplex ultrasonography. A venous duplex ultrasound examination uses a combination of B-mode imaging and spectral Doppler to detect the presence of venous obstruction and venous reflux in superficial and deep veins. Color-assisted Doppler ultrasound is useful to visualize venous flow patterns. Obstruction may be diagnosed by absence of flow, the presence of an echogenic thrombus within the vein, or failure of the vein to collapse when a compression maneuver is applied by the sonographer, the last implicating the presence of an intraluminal thrombus. Venous reflux is detected by prolonged reversal of venous flow direction during a Valsalva maneuver, particularly for the common femoral vein or saphenofemoral junction, or after compression and release of a cuff placed on the limb distal to the area being interrogated.

Some vascular laboratories use air or strain gauge plethysmography to assess the severity of venous reflux and complement findings from the venous ultrasound examination. Venous volume and venous refilling time are measured when the legs are placed in a dependent position and after calf exercise to quantify the severity of venous reflux and the efficiency of the calf muscle pump to affect venous return.

Magnetic resonance, computed tomographic, and conventional venography are rarely required to determine the cause and plan treatment for chronic venous insufficiency unless there is suspicion for pathology that might warrant intervention. These modalities are used to identify obstruction or stenosis of the inferior vena cava and iliofemoral veins, as may occur in patients with previous proximal

TABLE 303-1 CEAP (CLINICAL, ETIOLOGIC, ANATOMIC, PATHOPHYSIOLOGIC) CLASSIFICATION

Clinical Classification	
C0	No visible or palpable signs of venous disease
C1	Telangiectasias, reticular veins
C2	Varicose veins
C3	Edema without skin changes
C4	Skin changes, including pigmentation, eczema, lipodermatosclerosis, and atrophie blanche
C5	Healed venous ulcer
C6	Active venous ulcer
Etiologic Classification	
Ec	Congenital
Ep	Primary
Es	Secondary (postthrombotic)
En	No venous etiology identified
Anatomic Classification	
As	Superficial veins
Ap	Perforator veins
Ad	Deep veins
An	No venous location identified
Pathophysiologic Classification	
Pr	Reflux
Po	Obstruction
Pr,o	Reflux and obstruction
Pn	No venous pathophysiology identifiable

Source: B Eklöf et al: *J Vasc Surg* 40:1248, 2004.

deep vein thrombosis; occlusion of inferior vena cava filters; extrinsic compression from tumors; and May-Thurner syndrome.

TREATMENT CHRONIC VENOUS DISEASE

SUPPORTIVE MEASURES

Varicose veins usually are treated with conservative measures. Symptoms often decrease when the legs are elevated periodically, prolonged standing is avoided, and elastic support hose are worn. External compression with elastic stockings or stretch bandages provides a counterbalance to the hydrostatic pressure in the veins. Although compression garments may improve symptoms, they do not prevent progression of varicose veins. Graduated compression stockings with pressures of 20–30 mmHg are suitable for most patients with simple varicose veins, although pressures of 30–40 mmHg may be required for patients with manifestations of venous insufficiency such as edema and ulcers.

Patients with chronic venous insufficiency also should be advised to avoid prolonged standing or sitting; frequent leg elevation is helpful. Graded compression therapy consisting of stockings or multilayered compression bandages is the standard of care for advanced chronic venous insufficiency characterized by edema, skin changes, or venous ulcers defined as CEAP clinical class C3–C6. Graduated compression stockings of 30–40 mmHg are more effective than lesser grades for healing venous ulcers. The length of stocking depends on the distribution of edema. Calf-length stockings are tolerated better by most patients, particularly elderly patients; for patients with varicose veins or edema extending to the thigh, thigh-length stockings or panty hose should be considered. Overweight and obese patients should be advised to lose weight via caloric restriction and exercise.

In addition to a compression bandage or stocking, patients with venous ulcers also may be treated with low adherent absorbent dressings that take up exudates while maintaining a moist environment. Other types of dressings include hydrocolloid (an adhesive dressing comprised of polymers such as carboxymethylcellulose that absorbs exudates by forming a gel), hydrogel (a nonabsorbent

dressing comprising over 80% water or glycerin that moisturizes wounds), foam (an absorbent dressing made with polymers such as polyurethane), and alginate (an absorbent, biodegradable dressing that is derived from seaweed), but there is little evidence that these are more effective than low adherent absorbent dressings. The choice of specific dressing depends on the amount of drainage, presence of infection, and integrity of the skin surrounding the ulcer. Antibiotics are not indicated unless the ulcer is infected. The multilayered compression bandage or graduated compression garment is then put over the dressing.

MEDICAL THERAPIES

There are no drugs approved by the U.S. Food and Drug Administration for the treatment of chronic venous insufficiency. Diuretics may reduce edema, but at the risk of volume depletion and compromise in renal function. Topical steroids may be used for a short period of time to treat inflammation associated with stasis dermatitis. Several herbal supplements, such as horse chestnut seed extract (aescin); flavonoids including diosmin, hesperidin, or the two combined as micronized purified flavonoid fraction; and French maritime pine bark extract, are touted to have venoconstrictive and anti-inflammatory properties. Although meta-analyses have suggested that aescin reduces edema, pruritus, and pain and that micronized purified flavonoid fraction in conjunction with compression therapy facilitates venous ulcer healing, there is insufficient evidence to recommend the general use of these substances in patients with chronic venous insufficiency.

INTERVENTIONAL AND SURGICAL THERAPIES

Ablative procedures, including endovenous thermal ablation, sclerotherapy, and surgery, are used to treat varicose veins in selected patients who have persistent symptoms, great saphenous vein incompetency, and complications of venous insufficiency including dermatitis, edema, and ulcers. Ablative therapy may also be indicated for cosmetic reasons.

Endovenous thermal ablation procedures of the saphenous veins include endovenous laser therapy and radiofrequency ablation. To ablate the great saphenous vein, a catheter is placed percutaneously and advanced from the level of the knee to just below the saphenofemoral junction via ultrasound guidance. Thermal energy is then delivered as the catheter is pulled back. The heat injures the endothelium and media and promotes thrombosis and fibrosis, resulting in venous occlusion. Average 1- and 5-year occlusion rates exceed 90% following endovenous laser therapy and are slightly less after radiofrequency ablation. Deep vein thrombosis of the common femoral vein adjacent to the saphenofemoral junction is an uncommon but potential complication of endovenous thermal ablation. Other adverse effects of thermal ablation procedures include pain, paresthesias, bruising, hematoma, and hyperpigmentation.

Sclerotherapy involves the injection of a chemical into a vein to cause fibrosis and obstruction. Sclerosing agents approved by the U.S. Food and Drug Administration include sodium tetradecyl sulfate, polidocanol, sodium morrhuate, and glycerin. The sclerosing agent is administered as a liquid or mixed with air or CO₂/O₂ to create a foam. It first is injected into the great saphenous vein or its affected tributaries, often with ultrasound guidance. Thereafter, smaller more distal veins and incompetent perforating veins are injected. Following completion of the procedure, elastic bandages are applied, or 30–40 mmHg compression stockings are worn for 1–2 weeks. Average 1- and 5-year occlusion rates are 81% and 74%, respectively, following sclerotherapy. Complications are uncommon and include deep vein thrombosis, hematomas, damage to adjacent saphenous or sural nerves, and infection. Anaphylaxis is a very rare but severe complication.

Surgical therapy usually involves ligation and stripping of the great and small saphenous veins. The procedure is performed under general anesthesia. Incisions are made at the groin and the upper calf. The great saphenous vein is ligated below the saphenofemoral junction, and a wire is inserted into the great saphenous vein and

advanced distally. The proximal part of the great saphenous vein is secured to the wire and retrieved, i.e., stripped, via the calf incision. Stripping of the great saphenous vein below the knee and stripping of the small saphenous vein usually are not performed because of the respective risks of saphenous and sural nerve injury. Complications of great saphenous vein ligation and stripping include deep vein thrombosis, bleeding, hematoma, infection, and nerve injury. Recurrent varicose veins occur in up to 50% patients by 5 years, due to technical failures, deep venous insufficiency, and incompetent perforating veins.

Stab phlebectomy is another surgical treatment for of varicose veins. A small incision is made alongside the varicose vein, and it is avulsed by means of a forceps or hook. This procedure may be performed in conjunction with saphenous vein ligation and stripping or thermal ablation. Subfascial endoscopic perforator surgery (SEPS) uses endoscopy to identify and occlude incompetent perforating veins. It also may be performed along with other ablative procedures.

Endovascular interventions, surgical bypass, and reconstruction of the valves of the deep veins are performed when feasible to treat patients with advanced chronic venous insufficiency who have not responded to other therapies. Catheter-based interventions, usually involving placement of endovenous stents, may be considered to treat some patients with chronic occlusions of the iliac veins. Technical success rates exceed 85% in most series, and long-term patency is achieved in approximately 75% of these patients. Iliocaval bypass, femoroiliac venous bypass, and femorofemoral crossover venous bypass are procedures used occasionally to treat iliofemoral vein occlusion; saphenopopliteal vein bypass can be used to treat chronic femoropopliteal vein obstruction. Long-term patency rates for venous bypass procedures generally exceed 60% and are associated with improvement in symptoms. Surgical reconstruction of the valves of the deep veins and valve transfer procedures are used to treat valvular incompetence. Valvuloplasty involves tightening the valve by commissural apposition. With valve transfer procedures, a segment of vein with a competent valve, such as a brachial or axillary vein, or adjacent saphenous or deep femoral vein, is inserted as an interposition graft in the incompetent vein. Both valvuloplasty and vein transfer operations result in ulcer healing in the majority of patients, although success rates are somewhat better with valvuloplasty.

Lymphedema Lymphedema is a chronic condition caused by impaired transport of lymph and characterized by swelling of one or more limbs and occasionally the trunk and genitalia. Fluid accumulates in interstitial tissues when there is an imbalance between lymph production and lymph absorption, a process governed in large part by Starling forces. Deficiency, reflux, or obstruction of lymph vessels perturbs the ability of the lymphatic system to reabsorb proteins that had been filtered by blood vessels, and the tissue osmotic load promotes interstitial accumulation of water. Persistent lymphedema leads to inflammatory and immune responses characterized by infiltration of mononuclear cells, fibroblasts, and adipocytes, leading to adipose and collagen deposition in the skin and subcutaneous tissues.

Lymphatic Anatomy Lymphatic capillaries are blind-ended tubes formed by a single layer of endothelial cells. The absent or widely fenestrated basement membrane of lymphatic capillaries allows access to interstitial proteins and particles. Lymphatic capillaries merge to form microlymphatic precollector vessels, which contain few smooth muscle cells. The precollector vessels drain into collecting lymphatic vessels, which comprise endothelial cells, a basement membrane, smooth muscle, and bileaflet valves. The collecting lymphatic vessels in term merge to form larger lymphatic conduits. Analogous to venous anatomy, there are superficial and deep lymphatic vessels in the legs, which communicate at the popliteal and inguinal lymph nodes. Pelvic lymphatic vessels drain into the thoracic duct, which ascends from the abdomen to the thorax and connects with the left brachiocephalic

vein. Lymph is propelled centrally by the phasic contractile activity of lymphatic smooth muscle and facilitated by the contractions of contiguous skeletal muscle. The presence of lymphatic valves ensures unidirectional flow.

Etiology Lymphedema may be categorized as primary or secondary (Table 303-2). The prevalence of primary lymphedema is approximately 1.15 per 100,000 persons less than 20 years of age. Females are affected more frequently than males. Primary lymphedema may be caused by agenesis, hypoplasia, hyperplasia, or obstruction of the lymphatic vessels. There are three clinical subtypes: congenital lymphedema, which appears shortly after birth; lymphedema praecox, which has its onset at the time of puberty; and lymphedema tarda, which usually begins after age 35. Familial forms of congenital lymphedema (Milroy's disease) and lymphedema praecox (Meige's disease) may be inherited in an autosomal dominant manner with variable penetrance; autosomal or sex-linked recessive forms are less common. Mutations in genes expressing vascular endothelial growth factor receptor 3 (VEGFR3), which is a determinant of lymphangiogenesis, have been described in patients with Milroy's disease. A mutation on chromosome 15q is associated with the cholestasis-lymphedema syndrome. A mutation in the *FOXC2* gene, which encodes a transcription factor

TABLE 303-2 CAUSES OF LYMPHEDEMA

Primary
Sporadic (no identified cause)
Genetic disorders
Milroy's disease
Meige's disease
Lymphedema-distichiasis syndrome
Cholestasis-lymphedema
Hypotrichosis-lymphedema-telangiectasia
Turner's syndrome
Klinefelter's syndrome
Trisomy 13, 18, or 21
Noonan's syndrome
Klippel-Trénaunay syndrome
Parkes-Weber syndrome
Hennekam's syndrome
Yellow nail syndrome
Intestinal lymphangiectasia syndrome
Lymphangiomyomatosis
Neurofibromatosis type 1
Secondary
Infection
Bacterial lymphangitis (<i>Streptococcus pyogenes</i> , <i>Staphylococcus aureus</i>)
Lymphogranuloma venereum (<i>Chlamydia trachomatis</i>)
Filariasis (<i>Wucheria bancrofti</i> , <i>Brugia malayi</i> , <i>B. timori</i>)
Tuberculosis
Neoplastic infiltration of lymph nodes
Lymphoma
Prostate
Others
Surgery or irradiation of axillary or inguinal lymph nodes for treatment of cancer
Iatrogenic
Lymphatic division (during peripheral bypass surgery, varicose vein surgery, or harvesting of saphenous veins)
Miscellaneous
Contact dermatitis
Rheumatoid arthritis
Pregnancy
Factitious

that interacts with a signaling pathway involved in the development of lymphatic vessels, has been reported in patients with the lymphedema-distichiasis syndrome, in which lymphedema praecox occurs in patients who also have a double row of eyelashes. A mutation of *SOX18*, a transcription factor upstream of lymphatic endothelial cell differentiation, has been described in patients with lymphedema, alopecia, and telangiectasias (hypotrichosis, lymphedema, telangiectasia syndrome). Patients with a chromosomal aneuploidy, such as Turner's syndrome, Klinefelter's syndrome, or trisomy 18, 13, or 21, may develop lymphedema. Syndromic vascular anomalies associated with lymphedema include Klippel-Trénaunay syndrome, Parkes-Weber syndrome, and Hennekam's syndrome. Other disorders associated with lymphedema include Noonan's syndrome, yellow nail syndrome, intestinal lymphangiectasia syndrome, lymphangiomyomatosis, and neurofibromatosis type 1.

Secondary lymphedema is an acquired condition that results from damage to or obstruction of previously normal lymphatic channels. Recurrent episodes of bacterial lymphangitis, usually caused by streptococci, are a very common cause of lymphedema. The most common cause of secondary lymphedema worldwide is lymphatic filariasis, affecting approximately 129 million children and adults worldwide and causing lymphedema and elephantiasis in 14 million of these affected individuals (Chap. 258). Recurrent bacterial lymphangitis by *Streptococcus* may result in chronic lymphedema. Other infectious causes include lymphogranuloma venereum and tuberculosis. In developed countries, the most common secondary cause of lymphedema is surgical excision or irradiation of axillary and inguinal lymph nodes for treatment of cancers, such as breast, cervical, endometrial, and prostate cancer, sarcomas, and malignant melanoma. Lymphedema of the arm occurs in 13% of breast cancer patients after axillary node dissection and in 22% after both surgery and radiotherapy. Lymphedema of the leg affects approximately 15% of patients with cancer after inguinal lymph node dissection. Tumors, such as prostate cancer and lymphoma, also can infiltrate and obstruct lymphatic vessels. Less common causes include contact dermatitis, rheumatoid arthritis, pregnancy, and self-induced or factitious lymphedema after application of tourniquets.

Clinical Presentation Lymphedema is generally a painless condition, but patients may experience a chronic dull, heavy sensation in the leg, and most often they are concerned about the appearance of the leg. Lymphedema of the lower extremity initially involves the foot and gradually progresses up the leg so that the entire limb becomes edematous (Fig. 303-2). In the early stages, the edema is soft and pits easily with pressure. Over time, subcutaneous adipose tissue accumulates, the limb enlarges further and loses its normal contour, and the toes appear square. Thickening of the skin is detected by Stemmer's sign, which is the inability to tent the skin at the base of the toes. *Peau d'orange* is a term used to describe dimpling of the skin, resembling that of an orange peel, caused by lymphedema. In the chronic stages, the edema no longer pits and the limb acquires a woody texture as the tissues become indurated and fibrotic. The International Society of Lymphology describes four clinical stages of lymphedema (Table 303-3).

Differential Diagnosis Lymphedema should be distinguished from other disorders that cause unilateral leg swelling, such as deep vein thrombosis and chronic venous insufficiency. In the latter condition, the edema is softer, and there is often evidence of a stasis dermatitis, hyperpigmentation, and superficial venous varicosities, as described earlier. Other causes of leg swelling that resemble lymphedema are myxedema and lipedema. Lipedema usually occurs in women and is caused by accumulation of adipose tissue in the leg from the thigh to the ankle with sparing of the feet.

Diagnostic Testing The evaluation of patients with lymphedema should include diagnostic studies to clarify the cause. Abdominal and pelvic ultrasound and computed tomography (CT) can be used to detect obstructing lesions such as neoplasms. Magnetic resonance imaging (MRI) of the affected limb may reveal a honeycomb pattern



A Lymphedema characterized by swelling of the leg, nonpitting edema, and squaring of the toes. (Courtesy of Dr. Marie Gerhard-Herman, with permission.) **B** Advanced chronic stage of lymphedema illustrating the woody appearance of the leg with acanthosis and verrucous overgrowths. (Courtesy of Dr. Jeffrey Olin, with permission.)

characteristic of lymphedema in the epifascial compartment and identify enlarged lymphatic channels and lymph nodes. MRI also is useful to distinguish lymphedema from lipedema. Lymphoscintigraphy and lymphangiography are rarely indicated, but either can be used to confirm the diagnosis or differentiate primary from secondary lymphedema. Lymphoscintigraphy involves the injection of radioactively labeled technetium-containing colloid into the distal subcutaneous tissue of the affected extremity, which is imaged with a scintigraphic camera to visualize lymphatic vessels and lymph nodes. Findings indicative of primary lymphedema include absent or delayed filling of the lymphatic vessels or dermal back flow caused by lymphatic reflux. Findings of secondary lymphedema include dilated lymphatic vessels distal to an area of obstruction. In lymphangiography, iodinated radiocontrast material is injected into a distal lymphatic vessel that has been isolated and cannulated. In primary lymphedema, lymphatic channels are absent, hypoplastic, or ectatic. In secondary lymphedema, lymphatic channels often appear dilated beneath the level of obstruction.

TABLE 303-3 STAGES OF LYMPHEDEMA

Stage 0 (or Ia)	A latent or subclinical condition where swelling is not evident despite impaired lymph transport. It may exist for months or years before overt edema occurs.
Stage I	Early accumulation of fluid relatively high in protein content that subsides with limb elevation. Pitting may occur. An increase in proliferating cells may also be seen.
Stage II	Limb elevation alone rarely reduces tissue swelling, and pitting is manifest. Late in stage II, the limb may or may not pit as excess fat and fibrosis supervene.
Stage III	Lymphostatic elephantiasis where pitting can be absent and trophic skin changes such as acanthosis, further deposition of fat and fibrosis, and warty overgrowths have developed.

Source: Adapted from The 2013 Consensus Document of the International Society of Lymphology: *Lymphology* 46:1, 2013.

The complexities of lymphatic cannulation and the risk of lymphangitis associated with the contrast agent limit the utility of lymphangiography. A novel technique of optical imaging with a near-infrared fluorescence dye may enable quantitative imaging of lymph flow.

TREATMENT LYMPHEDEMA

Patients with lymphedema of the lower extremities must be instructed to take meticulous care of their feet to prevent recurrent lymphangitis. Skin hygiene is important, and emollients can be used to prevent drying. Prophylactic antibiotics are often helpful, and fungal infection should be treated aggressively. Patients should be encouraged to participate in physical activity; frequent leg elevation can reduce the amount of edema. Psychosocial support is indicated to assist patients cope with anxiety or depression related to body image, self-esteem, functional disability, and fear of limb loss.

Physical therapy, including massage to facilitate lymphatic drainage, may be helpful. The type of massage used in decongestive physiotherapy for lymphedema involves mild compression of the skin of the affected extremity to dilate the lymphatic channels and enhance lymphatic motility. Multilayered, compressive bandages are applied after each massage session to reduce recurrent edema. After optimal reduction in limb volume by decongestive physiotherapy, patients can be fitted with graduated compression hose. Occasionally, intermittent pneumatic compression devices can be applied at home to facilitate reduction of the edema. Diuretics are contraindicated and may cause depletion of intravascular volume and metabolic abnormalities.

Liposuction in conjunction with decongestive physiotherapy may be considered to treat lymphedema, particularly postmastectomy lymphedema. Other surgical interventions are rarely used and often not successful in ameliorating lymphedema. Microsurgical lymphaticovenous anastomotic procedures have been performed to rechannel lymph flow from obstructed lymphatic vessels into the venous system. Limb reduction procedures to resect subcutaneous tissue and excessive skin are performed occasionally in severe cases of lymphedema to improve mobility.

Therapeutic lymphangiogenesis has been studied in rodent models of lymphedema, but not as yet in humans. Overexpression of vascular endothelial growth factor (VEGF) C generates new lymphatic vessels and improves lymphedema in a murine model of primary lymphedema, and administration of recombinant VEGF-C or VEGF-D stimulated lymphatic growth in preclinical models of post-surgical lymphedema. Clinical trials in patients with lymphedema are required to determine efficacy of gene transfer (cell-based) therapies for lymphedema.

304 Pulmonary Hypertension

Aaron B. Waxman, Joseph Loscalzo

Pulmonary hypertension (PH) is a spectrum of diseases involving the pulmonary vasculature, and is defined as an elevation in pulmonary arterial pressures (mean pulmonary artery pressure >22 mmHg). Pulmonary arterial hypertension (PAH) is a relatively rare form of PH and is characterized by symptoms of dyspnea, chest pain, and syncope. If left untreated, the disease carries a high mortality rate, with the most common cause of death being decompensated right heart failure. There have been significant advances in this field in regard to understanding the pathogenesis, diagnosis, and classification of PAH. Despite these significant advances, there is still a substantial delay in diagnosis of up to 2 years. In many cases, patients whose primary complaint is dyspnea on exertion are frequently misdiagnosed with more

common diseases such as asthma or chronic obstructive pulmonary disease. The availability of newer drugs has resulted in a radical change in the management of this disease with significant improvement in both quality of life and mortality. A delay in diagnosis results in an obvious delay in the initiation of appropriate treatment. Clinicians should be able to recognize the signs and symptoms of PH and to complete a systematic workup in patients suspected of having it. In this way, early diagnosis, prompt treatment, and improved outcomes for patients become achievable.

PATHOBIOLOGY

Vasoconstriction, vascular proliferation, thrombosis, and inflammation appear to underlie the development of PAH (Fig. 304-1). In long-standing PH, intimal proliferation and fibrosis, medial hypertrophy, and in situ thrombosis characterize the pathologic findings in the pulmonary vasculature. Vascular remodeling at earlier stages may be confined to the small pulmonary arteries. As the disease advances, intimal proliferation and pathologic remodeling progress, resulting in decreased compliance and increased elastance of the pulmonary vasculature. The outcome is a progressive increase in the right ventricular afterload or total pulmonary vascular resistance (PVR) and, thus, right ventricular work. In subjects with moderate to severe pulmonary vascular disease with significantly increased PVR, as the resting PVR increases, there will be a corresponding increase in mean pulmonary artery pressure (PAP) until the cardiac output (CO) is compromised and starts to fall. With a decline in CO, the PAP will fall. As CO declines as a result of increased afterload and decreased contractility, tachycardia is a compensatory response. Tachycardia decreases filling time and, thus, preload, and results in a reduced fraction of stroke volume available to distend the pulmonary vascular tree.

Abnormalities in multiple molecular pathways and genes that regulate the pulmonary vascular endothelial and smooth muscle cells have been identified (Table 304-1). These abnormalities include decreased expression of the voltage-regulated potassium channel, mutations in the bone morphogenetic protein receptor-2, increased tissue factor expression, overactivation of the serotonin transporter, hypoxia-induced activation of hypoxia-inducible factor-1 α , and activation of nuclear factor of activated T cells. As a result, there is a decrease in apoptosis of the smooth muscle cells and the emergence of apoptosis-resistant endothelial cells that promote their accumulation and can obliterate the vascular lumen. In addition, thrombin deposition in the pulmonary vasculature from the prothrombotic state that develops as an independent abnormality or as a result of endothelial dysfunction may amplify vascular cell proliferation and the obliterative arteriopathy.

DIAGNOSIS AND CLASSIFICATION

The diagnosis of PH can be missed without a reasonable index of suspicion. Dyspnea is the most common presenting symptom, but this complaint is far from specific for the diagnosis of PH. PH symptoms are insidious and overlap considerably with many common conditions, including asthma and other lung disease and cardiac disease. The symptoms of PH are often nonspecific and variable. Most patients will present with dyspnea and/or fatigue, whereas edema, chest pain, presyncope, and frank syncope are less common and associated with more advanced disease. On examination, there may be evidence of right ventricular failure with elevated jugular venous pressure, lower extremity edema, and ascites. Additionally, the cardiovascular examination may reveal an accentuated P₂ component of the second heart sound, a right-sided S₃ or S₄, and a holosystolic tricuspid regurgitant murmur. It is also important to seek signs of the diseases that are often concurrent with PH: clubbing may be seen in some chronic lung diseases, sclerodactyly and telangiectasia may signify scleroderma, and crackles and systemic hypertension may be clues to left-sided systolic or diastolic heart failure.

Once clinical suspicion is raised, a systematic approach to diagnosis and assessment is essential. An echocardiogram with (if indicated) a *bubble study* is the most important screening test. Echocardiography is important for the diagnosis of PH and often essential for determining the cause. All forms of PH may demonstrate a hypertrophied and

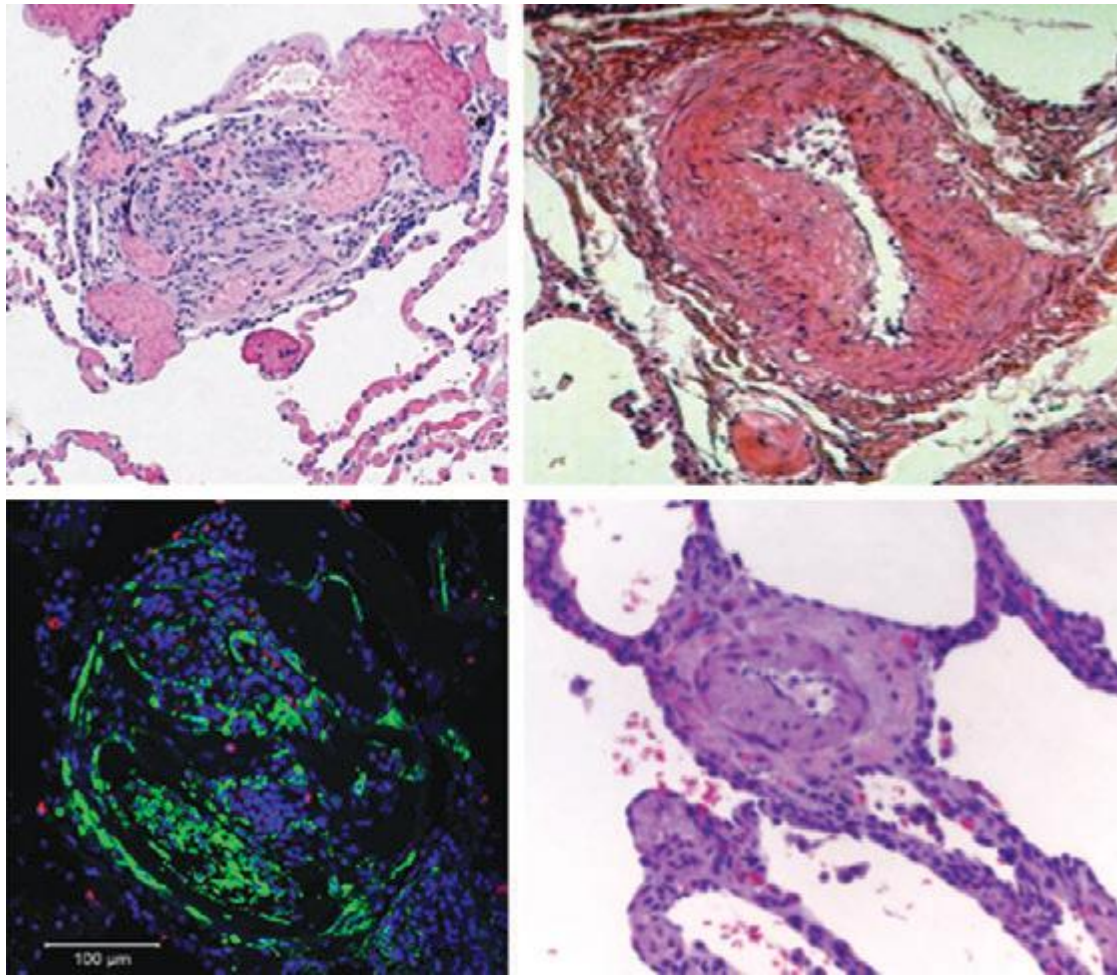


FIGURE 304-1 The left panels show examples of plexogenic pulmonary arteriopathy. These are obstructive and proliferative lesions of the small muscular pulmonary arteries, composed primarily of endothelial cells with intermixed inflammatory cells, myofibroblasts, and connective tissue components. The lower left panel demonstrates proliferating cells (red PCNA stained cells). Panels on the right demonstrate medial hypertrophy of muscular pulmonary arteries. (Photographs on the left are courtesy of Dr. Stephen Archer, Queen's University School of Medicine, Kingston, Ontario, Canada.)

dilated right ventricle (Fig. 304-2) with elevated estimated pulmonary artery systolic pressure. Important additional information can be gleaned about specific etiologies of PH such as valvular disease, left ventricular systolic and diastolic function, intracardiac shunts, and other cardiac diseases.

Although the accuracy of Doppler echocardiography is often debated, a high-quality echocardiogram that is absolutely normal may obviate the need for further evaluation for PH. An echocardiogram is a screening test, whereas invasive hemodynamic monitoring is the gold standard for diagnosis and assessment of disease severity. With a normal echocardiogram, there may still be some concern for PH; this is particularly true if there is unexplained dyspnea or hypoxemia. In this setting, it is reasonable to proceed to right heart catheterization for definitive diagnosis. Alternatively, if the patient has a reasonable functional capacity, a cardiopulmonary exercise test may help to identify a true physiologic limitation as well as differentiate between cardiac and pulmonary causes of dyspnea. If this test is normal, there is no indication for a right heart catheterization. If a cardiovascular limitation to exercise is found, a right heart catheterization should be pursued.

If the echocardiogram or cardiopulmonary exercise test (CPET) suggests PH and the diagnosis is confirmed by catheterization, a reasonable effort must be made to establish the etiology because this will largely determine the therapeutic approach. A stepwise approach to evaluation is outlined below.

Chest imaging and lung function tests are essential because lung disease is an important cause of PH. A sign of PH that may be evident on chest x-ray include enlargement of the central pulmonary arteries

associated with “vascular pruning,” a relative paucity of peripheral vessels (Fig. 304-3). Cardiomegaly, with specific evidence of right atrial and ventricular enlargement, can often be observed. The chest x-ray may also demonstrate significant interstitial lung disease or suggest hyperinflation from obstructive lung disease, which may be the underlying cause or contributor to the development of PH. High-resolution computed tomography (CT) may provide additional useful information. Classic findings of PH on CT include those found on chest x-ray: enlarged pulmonary arteries (Fig. 304-4), peripheral pruning of the small vessels, and enlarged right ventricle and atrium. However, high-resolution CT may also reveal signs of venous congestion including centrilobular ground-glass infiltrate and thickened septal lines. In the absence of left heart disease, these findings suggest pulmonary veno-occlusive disease, a rare cause of PAH that can be quite challenging to diagnose.

CT angiograms are commonly used to evaluate acute thromboembolic disease and have demonstrated excellent sensitivity and specificity for that purpose. Ventilation-perfusion (\dot{V}/\dot{Q}) scanning has traditionally been used for screening because of its high sensitivity and its role in qualifying patients for surgical intervention. The role of CT angiograms in the diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH) remains controversial, even with the advent of spiral CT. Although a negative \dot{V}/\dot{Q} virtually rules out CTEPH, some cases may be missed through the use of CT angiograms.

Pulmonary function tests are an important component of the evaluation. Although an isolated reduction in DL_{CO} is the classic finding in PAH, results of pulmonary function tests may also suggest restrictive or obstructive lung diseases as the cause of dyspnea or PH. The 6-minute walk test

TABLE 304-1 COMPONENTS OF THE PATHOGENESIS OF PULMONARY ARTERIAL HYPERTENSION

Alterations in regulators of proliferation

- Growth factors
 - Platelet-derived growth factor
 - Fibroblast growth factor
 - Vascular endothelial growth factor
 - Epidermal growth factor
- Transforming growth factor β (TGF- β)
- Bone morphogenetic protein
- Transcription factors
- Matrix metalloproteinases
- Cytokines
- Chemokines
- Mitochondria

Alterations in inflammatory mediators

- Altered T cell subsets
- Monocytes and macrophages
- Interleukin (IL) 1 β
- IL-6
- MCP-1
- RANTES
- Fractalkine

Alterations in vascular tone

- Endothelin
- Nitric oxide
- Serotonin
- Prostaglandin
- K⁺ channels
- Ca²⁺ channels

Hypoxia-induced remodeling

- HIF-1 α
- ROS
- Mitochondria

TGF- β signaling

- BMPR2
- ALK1
- Endoglin
- Smad9
- TGF- β 1

Abbreviations: PDGF, platelet-derived growth factor; EGF, epidermal-derived growth factor; FGF, fetal-derived growth factor; VEGF, vascular endothelial-derived growth factor; MCP-1, monocyte chemoattractant protein-1; IL, interleukin.

is also important to evaluate the degree of exertional hypoxemia and limitation and to monitor progression and response to therapy.

Sleep-disordered breathing is another important cause of PH, but a sleep study is generally necessary only when indicated by the patient's history. Nocturnal desaturation is a common finding in PH, even in the absence of sleep-disordered breathing. Thus, all patients should undergo nocturnal oximetry screening, regardless of whether classic symptoms of obstructive sleep apnea or obesity-hypoventilation syndrome are observed. Laboratory tests that are important for screening include an HIV test when clinically indicated. In addition, all patients should have antinuclear antibodies, rheumatoid factor, and scl-70 antibodies assessed to screen for the most common rheumatologic diseases associated with PH if clinically indicated. Liver function and hepatitis serology tests are important to screen for underlying liver disease. Finally, there is an increasing role for brain natriuretic peptide (BNP) testing in the diagnosis and management of PH. BNP and the N-terminus of its propeptide (NT-proBNP) correlate with right ventricular function, hemodynamic severity, and functional status in PAH.

Right heart catheterization with pulmonary vasodilator testing remains the gold standard both to establish the diagnosis of PH and to enable selection of appropriate medical therapy. The definition of precapillary PH or PAH requires (1) an increased mean pulmonary artery pressure (mPAP ≥ 25 mmHg); (2) a pulmonary capillary wedge pressure (PCWP), left atrial pressure, or left ventricular end-diastolic pressure ≤ 15 mmHg; and (3) PVR > 3 Wood units. Postcapillary PH is differentiated from precapillary PH by a PCWP of ≥ 15 mmHg; this is further differentiated into passive, based on a transpulmonary gradient < 12 mmHg, or reactive, based on a transpulmonary gradient > 12 mmHg and an increased PVR. In either case, the CO may be normal or reduced.

Vasodilators with a short duration of action, such as inhaled nitric oxide, inhaled epoprostenol, or intravenous adenosine, are preferred for vasodilator testing. A decrease in mPAP by ≥ 10 mmHg to an absolute level of ≤ 40 mmHg without a decrease in CO is defined as a positive pulmonary vasodilator response, and responders are considered for long-term treatment with calcium channel blockers (CCBs). Less than 12% of patients are deemed vasoreactive during testing, and even fewer exhibit long-term responsiveness to CCBs. Acute vasodilator-induced reductions in PVR and mPAP predict better long-term survival even among patients not treated with CCBs. The need for invasive hemodynamic measurements to diagnose PH accurately poses an additional problem when evaluating older patients. Physicians are often reluctant to refer older patients for invasive procedures. However, the diagnosis of PH is increasing in the older population, at least in part because of increased awareness of this disease in the elderly and increased use of screening echocardiograms. Furthermore, the increased availability of oral and less complicated therapeutic options has encouraged the referral of older patients for evaluation and treatment.

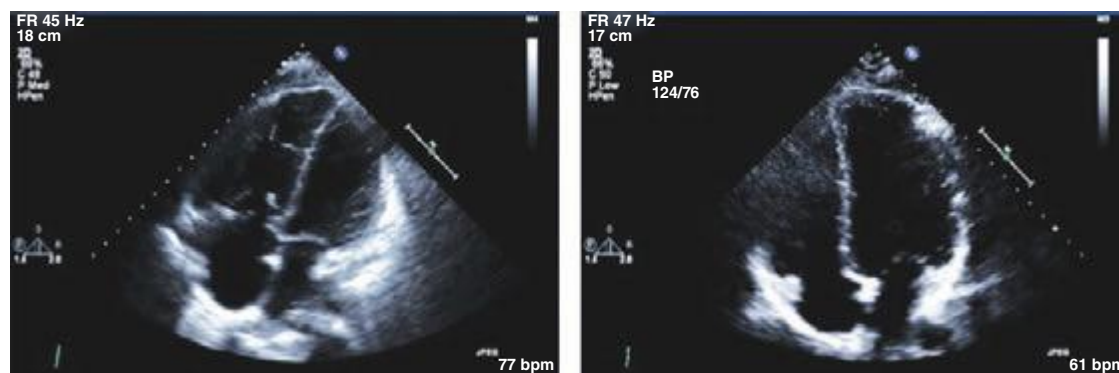


FIGURE 304-2 **A.** Representative echocardiogram showing the apical four-chamber view from a patient with pulmonary hypertension demonstrating an enlarged right atrium and ventricle with some compression of the left side of the heart. **B.** Same echocardiographic view showing a normal echocardiogram.

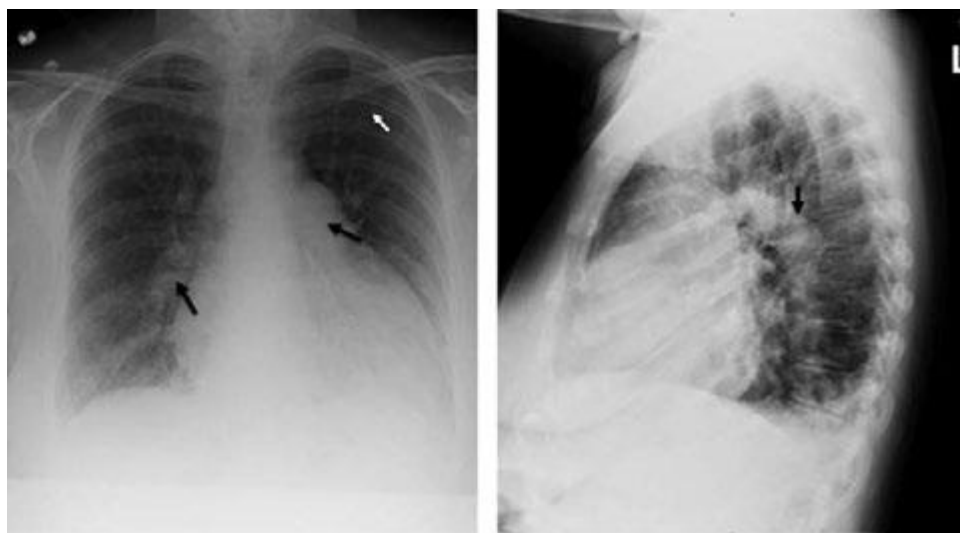


FIGURE 304-3 Posteroanterior (*left*) and lateral (*right*) chest radiograph showing enlarged pulmonary arteries (*black arrows*) and pruning of the distal pulmonary vasculature (*white arrow*) commonly seen with advanced pulmonary arterial hypertension.

PULMONARY HYPERTENSION AS A COMORBID DISEASE

PAH is just one of a number of disease classifications that affect the pulmonary vascular bed. PH was previously classified as primary or secondary, but as understanding of the various contributing diseases has increased, classification systems have attempted to group these diseases by clinical features to aid in diagnosis. The World Health Organization (WHO) formulated a clinical classification of the various manifestations of PH, of which PAH is a subgroup, according to similarities in pathophysiologic mechanisms and clinical presentation. PH is a diverse mix of pathologies in which the only unifying theme is elevated PAP relative to left atrial pressure. The categorization of PH was designed by convenience for the purpose of facilitating novel treatments to be tested across different presentations and is not based on a molecular understanding of the pathology and is not a guide for management decisions.



FIGURE 304-4 Representative computed tomography scan of the chest demonstrating enlarged main pulmonary arteries. There is also a mosaic pattern evident in both lungs.

The current classification system, last revised in 2013 during the Fifth World Symposium on Pulmonary Hypertension, recognizes five categories of PH, including PAH, PH due to left heart disease, PH due to chronic lung disease, PH associated with chronic thromboemboli, and a group of miscellaneous diseases that only rarely cause PH.

Pulmonary Arterial Hypertension WHO Group I PH, pulmonary arterial hypertension (PAH), is a relatively rare cause of PH. PAH includes a group of diseases that result in pulmonary arterial precapillary remodeling marked by intimal fibrosis, increased medial thickness, pulmonary arteriolar occlusion, and classic plexiform lesions. PAH is defined as a sustained elevation in resting mPAP ≥ 25 mmHg, PVR > 240 dyne-s/cm⁵, and PCWP or left ventricle end-diastolic pressure of ≤ 15 mmHg based on a right

heart catheterization. With a normal PCWP and an elevated mPAP, these diseases demonstrate an increased transpulmonary gradient (mPAP – PCWP); in addition, the PVR is elevated.

Idiopathic pulmonary arterial hypertension (IPAH) is a progressive disease that leads to right heart failure and death. It is typically seen in young women. The National Institutes of Health registry, the first large registry of patients with PAH, reported that the average age at diagnosis was 36 years, with only 9% of patients with IPAH over the age of 60 at diagnosis. However, the more current clinical data suggest that the patient demographics are changing. The Pulmonary Hypertension Connection registry found that the average age of diagnosis for IPAH was 45 years, with 8.5% of patients older than 70 years at diagnosis. This finding is supported by data from the Registry to Evaluate Early and Long-Term PAH Disease Management (REVEAL), the largest cohort of PAH to date, which reported that the average age at diagnosis of IPAH was 44.9 ± 0.6 years.

Other forms of PAH that deserve specific consideration in patients are those associated with HIV, connective tissue disease, and portal hypertension. Although HIV is a rare cause of PAH, this form of PAH is indistinguishable from IPAH and is an important cause of mortality in the HIV-infected population. Importantly, there is no correlation between the stage of HIV infection and the development of PAH.

Among connective tissue diseases, the prevalence of PAH has been established only for systemic sclerosis, especially in those with limited cutaneous scleroderma. Although the average age of scleroderma onset is 30 to 50 years old, patients who eventually develop scleroderma-associated PAH tend to be older at the time of scleroderma diagnosis. Outcomes of scleroderma are closely linked to the development of PAH and are associated with a poor prognosis, although modern therapies have improved outcomes.

Portopulmonary hypertension occurs in 2–10% of patients with established portal hypertension. Its occurrence appears to be independent of the cause of liver disease and is observed in patients with nonhepatic causes of portal hypertension. A hyperdynamic circulatory state is common, as in most patients with advanced liver disease; however, the same pulmonary vascular remodeling observed in other forms of PAH is seen in the pulmonary vascular bed in portopulmonary hypertension. It is important to distinguish this process from hepatopulmonary syndrome, which can also manifest with dyspnea and hypoxemia but is pathophysiologically distinct from portopulmonary hypertension in that abnormal vasodilation of the pulmonary vasculature leads to intrapulmonary shunting.

Pulmonary Hypertension Associated with Left Heart Disease WHO Group II PH includes patients with left heart systolic failure, aortic and

mitral valve disease, and heart failure with preserved ejection fraction (HFpEF). PH can develop as a result of all of these conditions. The hallmark of Group II PH (i.e., PH due to left heart disease) is elevated left atrial pressure with resulting pulmonary venous hypertension. In general, the transpulmonary gradient and PVR remain normal. Although this phenomenon is well described in both left-sided valvular disease and left-sided systolic heart failure, studies suggest that HFpEF may carry a higher overall risk of PH.

Whatever the cause of elevated left atrial pressure (i.e., systolic or diastolic heart failure or valvular disease), the increased pulmonary venous pressure indirectly leads to a rise in pulmonary arterial pressure. The presence of PH portends a poor prognosis in all forms of heart failure. In particular, chronic pulmonary venous hypertension may lead to a reactive pulmonary arterial vasculopathy, seen as an elevated transpulmonary gradient (>12 mmHg) and elevated PVR (>3 Wood units). Pathologically, this process is marked by pulmonary arteriolar remodeling with intimal fibrosis and medial hyperplasia akin to that seen in PAH.

Pulmonary Hypertension Associated with Lung Disease Intrinsic lung disease is the second most common cause of PH, although its actual prevalence is difficult to ascertain. PH has been observed in both chronic obstructive lung disease and interstitial lung disease. It can also be seen in diseases with mixed obstructive/restrictive physiology: bronchiectasis, cystic fibrosis, mixed obstructive restrictive disease marked by fibrosis in the lower lung zones, and emphysema predominantly in the upper lung zones. As in patients with left heart disease, PH associated with chronic lung disease is usually modest; however, some of these patients appear to have PH “out of proportion” to their parenchymal lung disease, suggesting intrinsic pulmonary arterial disease. These patients typically have more severe PH, with results of pulmonary function tests demonstrating a very low DL_{CO} .

Although PH is described in most forms of interstitial lung disease, it has been most extensively studied in idiopathic pulmonary fibrosis; however, the individual studies have been small. Early echocardiographic data suggested that the prevalence of PH in interstitial lung diseases was high, but invasive hemodynamic monitoring suggests that the incidence is considerably lower than originally believed. The diagnosis of PH portends poor outcome in pulmonary fibrosis.

Also included in Group III PH is sleep-disordered breathing. Sleep apnea has long been associated with PH. However, PH associated with sleep-disordered breathing is generally mild.

Pulmonary Hypertension Associated with Chronic Thromboembolic Disease The development of PH after chronic thromboembolic obstruction of the pulmonary arteries is well described, but its incidence is not known. The incidence of PH after a single pulmonary embolic event is thought to be quite low and likely increases following recurrent embolism. The risk factors for developing CTEPH are unclear. Many patients have no history of clinical venous thromboembolism. The pathogenesis of CTEPH is poorly understood. Obstruction of the proximal pulmonary vasculature is important and often the dominant factor; however, additional pulmonary vascular remodeling occurs. Approximately 10–15% of patients will develop a disease very similar clinically and pathologically to PAH after resection of the proximal thrombus.

OTHER DISORDERS AFFECTING THE PULMONARY VASCULATURE

Sarcoidosis Patients with sarcoidosis can develop PH as a result of lung involvement. Consequently, patients with sarcoidosis who present with progressive dyspnea and PH require a thorough evaluation. Although the majority of sarcoidosis patients with PH generally do not respond to therapy for PAH, a subset of patients with sarcoidosis and severe PH do have a beneficial response to therapy.

Sickle Cell Disease Cardiovascular system abnormalities are prominent in the clinical spectrum of sickle cell disease, including PH. The etiology is multifactorial, including hemolysis, hypoxemia, thromboembolism, chronic high CO, and chronic liver disease. The presence of PH in patients with sickle cell disease is rare.

Schistosomiasis Globally, schistosomiasis is one of the most common causes of PH. The development of PH occurs in the setting of hepatosplenic disease and portal hypertension. Studies suggest that inflammation from the infection triggers the pulmonary vascular changes that occur. The diagnosis is confirmed by finding the parasite ova in the urine or stool of patients with symptoms, which can be difficult. The efficacy of therapies directed toward PH in these patients is unknown.

PHARMACOLOGIC TREATMENT OF PAH

PH was a consistently fatal condition with no effective medical treatment options before 1996; however, since that time, there has been an upsurge in the development of novel therapeutic agents for PAH. There are several approved agents for PAH, including prostacyclin and prostacyclin analogues, phosphodiesterase-5 inhibitors, a soluble guanylyl cyclase stimulator, and endothelin receptor antagonists, that have improved the outlook dramatically. Although there is no cure for PAH, current pharmacologic therapies improve morbidity and, in some cases, mortality.

PROSTANOIDS

In PAH, endothelial dysfunction and platelet activation cause an imbalance of arachidonic acid metabolites with reduced prostacyclin levels and increased thromboxane A_2 production. Prostacyclin (PGI_2) activates cyclic adenosine monophosphate (cAMP)-dependent pathways that mediate vasodilation. PGI_2 also has antiproliferative effects on vascular smooth muscle and inhibits platelet aggregation. Protein levels of prostacyclin synthase are decreased in pulmonary arteries of patients with PAH. This imbalance of mediators is addressed by the exogenous administration of prostanoids as therapy in advanced PAH.

Epoprostenol was the first prostanoid available for the management of PAH. Epoprostenol delivered as a continuous intravenous infusion improves functional capacity and survival in PAH. The efficacy of epoprostenol in WHO functional class 3 and 4 PAH patients was demonstrated in a clinical trial that showed improved quality of life, mPAP, PVR, 6-minute walk distance (6MWD), and mortality. Treprostinil has a longer half-life than epoprostenol (~4 h vs ~6 min), which allows for continuous subcutaneous and intravenous administration. Treprostinil has been shown to improve pulmonary hemodynamics, symptoms, exercise capacity, and survival in PAH.

Inhaled prostacyclins provide the beneficial effects of infused prostacyclin therapy without the inconvenience and side effects (risk of infection and infusion site reactions) of infusion catheters. Both inhaled iloprost and treprostinil have been approved for patients with WHO class 3 and 4 PAH. The main advantage of treprostinil is less frequent administration. Inhaled formulations can be efficacious in moderately symptomatic patients with PAH and may be appropriate when used in combination with an oral medication. Phosphodiesterase-5 (PDE5) inhibitors (e.g., sildenafil) increase cyclic guanosine monophosphate (cGMP) levels and activate cGMP-dependent signaling pathways that also mediate vasodilation and platelet inhibition. Thus, the addition of a PDE5 inhibitor augments the pulmonary hemodynamic and functional capacity benefits of prostanoids in PAH.

Endothelin Receptor Antagonists Endothelin receptor antagonists (ERAs) target endothelin-1 (ET-1), a potent endogenous vasoconstrictor and vascular smooth muscle mitogen that is elevated in PAH patients. Endothelin levels are increased coincident with increased PVR and mPAP and decreased CO and 6MWD.

ERAs block the binding of ET-1 to either endothelin receptor A (ET-A) and/or B (ET-B). ET-A receptors found on pulmonary artery smooth muscle cells mediate vasoconstriction. In the normal pulmonary vasculature, ET-B receptors are found on endothelial cells and mediate vasodilation via production of prostacyclin and nitric oxide as well as ET-1 clearance. Three ERAs approved for use in the United States are bosentan and macitentan both, nonselective receptor antagonists, and ambrisentan, a selective ET-A receptor antagonist.

Studies have shown that both bosentan and macitentan improve hemodynamics and exercise capacity and delay clinical worsening.

TABLE 304-2 FDA-APPROVED THERAPIES FOR THE TREATMENT OF PAH

Generic Name	Route of Administration	Drug Class	Indication
Epoprostenol	IV	Prostacyclin derivative	Treatment of PAH to improve exercise capacity
Iloprost	Inhaled	Prostacyclin derivative	Treatment of PAH to improve a composite endpoint consisting of exercise tolerance, symptoms (NYHA Class), and lack of deterioration
Treprostinil	IV or SC	Prostacyclin derivative	Treatment of PAH to diminish symptoms associated with exercise
Treprostinil	Inhaled	Prostacyclin derivative	Treatment of PAH to improve exercise ability
Treprostinil	Oral	Prostacyclin derivative	Treatment of PAH to improve exercise ability
Bosentan	Oral	Non-selective endothelin receptor antagonist	Treatment of PAH to improve exercise capacity and to decrease clinical worsening
Ambrisentan	Oral	Endothelin receptor antagonist	Treatment of PAH to improve exercise capacity and delay clinical worsening
Macitentan	Oral	Non-selective endothelin antagonist	Treatment of PAH to improve exercise capacity and delay clinical worsening
Sildenafil	Oral	PDE5 inhibitor	Treatment of PAH to improve exercise capacity and delay clinical worsening
Tadalafil	Oral	PDE5 inhibitor	Treatment of PAH to improve exercise ability
Riociguat	Oral	Soluble guanylyl cyclase stimulator	Treatment of PAH to improve exercise capacity and delay clinical worsening

Abbreviations: FDA, U.S. Food and Drug Administration; NYHA, New York Heart Association; PAH, pulmonary arterial hypertension; PDES, phosphodiesterase-5.

The randomized, placebo-controlled, phase III Bosentan Randomized Trial of Endothelin Antagonist Therapy (BREATHE)-1 comparing bosentan with placebo demonstrated improved symptoms, 6MWD, and WHO functional class. The Endothelin Antagonist Trial in Mildly Symptomatic Pulmonary Arterial Hypertension Patients (EARLY) comparing bosentan with placebo demonstrated improved PVR and 6MWD.

Several studies, including the phase III, placebo-controlled Ambrisentan in Pulmonary Arterial Hypertension-1 (ARIES-1) trial, suggest that ambrisentan improves exercise tolerance, WHO functional class, hemodynamics, and quality of life in patients with PAH. There are no trial data to evaluate whether the selective ET-A receptor antagonism of ambrisentan has any advantage over the nonselective ET receptor antagonism of bosentan.

Phosphodiesterase Type-5 Inhibitors Nitric oxide derived from endothelial cells activates guanylyl cyclase, which, in turn, generates cGMP in vascular smooth muscle cells and platelets. cGMP is a second messenger that induces vasodilation through relaxation of the arterial smooth muscle cells and inhibits platelet activation. PDE5 enzymes metabolize cGMP. Therefore, cGMP PDE5 inhibitors prolong the vasodilatory effect of nitric oxide, especially within the pulmonary arterial bed where high concentrations of cGMP are found. There

are currently two PDE5 inhibitors used for the treatment of PAH, sildenafil and tadalafil. Both agents have been shown to improve hemodynamics and 6MWD. Recently, the oral soluble guanylyl cyclase stimulator, riociguat, was approved for the treatment of both PAH and CTEPH.

Unmet and Future Research Needs in Pulmonary Hypertension Presently there are only three classes of therapy for patients with PAH, and even with therapy, the median survival for a person with PAH is only 5–6 years (Table 304-2). Although there are five subtypes of PH, current approved therapies only address one subtype. Not only do we need to expand the treatment options for patients with PAH, we also need to develop effective therapies for all patients with PH. Limited survival is, in part, a result of delay in diagnosis. Improved awareness among clinicians and patients could lead to more timely diagnosis that will affect the response to therapy and survival. PH needs to be diagnosed in a timely manner so that therapy can be initiated as soon as possible. Patients should also have the option of referral to a specialty center that focuses on treatment of patients with pulmonary vascular disease, which will ensure their access to state-of-the-art care and a multidisciplinary approach to care. Finally, there need to be continued efforts at developing new therapies that target the increasingly complex and overlapping pathways involved in the various forms of PH.

SECTION 1 DIAGNOSIS OF RESPIRATORY DISORDERS

305 Approach to the Patient with Disease of the Respiratory System

Patricia A. Kritek, Augustine M. K. Choi

The majority of diseases of the respiratory system fall into one of three major categories: (1) obstructive lung diseases; (2) restrictive disorders; and (3) abnormalities of the vasculature. Obstructive lung diseases are most common and primarily include disorders of the airways, such as asthma, chronic obstructive pulmonary disease (COPD), bronchiectasis, and bronchiolitis. Diseases resulting in restrictive pathophysiology include parenchymal lung diseases, abnormalities of the chest wall and pleura, and neuromuscular disease. Disorders of the pulmonary vasculature include pulmonary embolism, pulmonary hypertension, and pulmonary veno-occlusive disease. Although many specific diseases fall into these major categories, both infective and neoplastic processes can affect the respiratory system and result in myriad pathologic findings, including those listed in the three categories above (Table 305-1).

Disorders can also be grouped according to gas exchange abnormalities, including hypoxemic, hypercarbic, or combined impairment. However, many diseases of the lung do not manifest as gas exchange abnormalities.

As with the evaluation of most patients, the approach to a patient with disease of the respiratory system begins with a thorough history and a focused physical examination. Many patients will subsequently undergo pulmonary function testing, chest imaging, blood and sputum analysis, a variety of serologic or microbiologic studies, and diagnostic procedures, such as bronchoscopy. This stepwise approach is discussed in detail below.

TABLE 305-1 CATEGORIES OF RESPIRATORY DISEASE

Category	Examples
Obstructive lung disease	Asthma
	Chronic obstructive pulmonary disease (COPD)
	Bronchiectasis
	Bronchiolitis
Restrictive pathophysiology—parenchymal disease	Idiopathic pulmonary fibrosis (IPF)
	Asbestosis
	Desquamative interstitial pneumonitis (DIP)
Restrictive pathophysiology—neuromuscular weakness	Sarcoidosis
Restrictive pathophysiology—chest wall/pleural disease	Amyotrophic lateral sclerosis (ALS)
	Guillain-Barré syndrome
Restrictive pathophysiology—pulmonary vascular disease	Kyphoscoliosis
	Ankylosing spondylitis
	Chronic pleural effusions
Malignancy	Pulmonary embolism
	Pulmonary arterial hypertension (PAH)
Infectious diseases	Bronchogenic carcinoma (non-small-cell and small-cell)
	Metastatic disease
	Pneumonia
Infectious diseases	Bronchitis
	Tracheitis

HISTORY

Dyspnea and Cough The cardinal symptoms of respiratory disease are dyspnea and cough (Chaps. 47e and 48). Dyspnea has many causes, some of which are not predominantly due to lung pathology. The words a patient uses to describe shortness of breath can suggest certain etiologies for dyspnea. Patients with obstructive lung disease often complain of “chest tightness” or “inability to get a deep breath,” whereas patients with congestive heart failure more commonly report “air hunger” or a sense of suffocation.

The tempo of onset and the duration of a patient’s dyspnea are likewise helpful in determining the etiology. Acute shortness of breath is usually associated with sudden physiologic changes, such as laryngeal edema, bronchospasm, myocardial infarction, pulmonary embolism, or pneumothorax. Patients with COPD and idiopathic pulmonary fibrosis (IPF) experience a gradual progression of dyspnea on exertion, punctuated by acute exacerbations of shortness of breath. In contrast, most asthmatics have normal breathing the majority of the time with recurrent episodes of dyspnea that are usually associated with specific triggers, such as an upper respiratory tract infection or exposure to allergens.

Specific questioning should focus on factors that incite dyspnea as well as on any intervention that helps resolve the patient’s shortness of breath. Asthma is commonly exacerbated by specific triggers, although this can also be true of COPD. Many patients with lung disease report dyspnea on exertion. Determining the degree of activity that results in shortness of breath gives the clinician a gauge of the patient’s degree of disability. Many patients adapt their level of activity to accommodate progressive limitation. For this reason, it is important, particularly in older patients, to delineate the activities in which they engage and how these activities have changed over time. Dyspnea on exertion is often an early symptom of underlying lung or heart disease and warrants a thorough evaluation.

Cough generally indicates disease of the respiratory system. The clinician should inquire about the duration of the cough, whether or not it is associated with sputum production, and any specific triggers that induce it. Acute cough productive of phlegm is often a symptom of infection of the respiratory system, including processes affecting the upper airway (e.g., sinusitis, tracheitis), the lower airways (e.g., bronchitis, bronchiectasis), and the lung parenchyma (e.g., pneumonia). Both the quantity and quality of the sputum, including whether it is blood-streaked or frankly bloody, should be determined. Hemoptysis warrants an evaluation as delineated in Chap. 48.

Chronic cough (defined as that persisting for >8 weeks) is commonly associated with obstructive lung diseases, particularly asthma and chronic bronchitis, as well as “nonrespiratory” diseases, such as gastroesophageal reflux and postnasal drip. Diffuse parenchymal lung diseases, including IPF, frequently present as a persistent, nonproductive cough. As with dyspnea, all causes of cough are not respiratory in origin, and assessment should encompass a broad differential, including cardiac and gastrointestinal diseases as well as psychogenic causes.

Additional Symptoms Patients with respiratory disease may report wheezing, which is suggestive of airways disease, particularly asthma. Hemoptysis can be a symptom of a variety of lung diseases, including infections of the respiratory tract, bronchogenic carcinoma, and pulmonary embolism. In addition, chest pain or discomfort is often thought to be respiratory in origin. As the lung parenchyma is not innervated with pain fibers, pain in the chest from respiratory disorders usually results from either diseases of the parietal pleura (e.g., pneumothorax) or pulmonary vascular diseases (e.g., pulmonary hypertension). As many diseases of the lung can result in strain on

1662 the right side of the heart, patients may also present with symptoms of cor pulmonale, including abdominal bloating or distention and pedal edema (Chap. 279).

Additional History A thorough social history is an essential component of the evaluation of patients with respiratory disease. All patients should be asked about current or previous cigarette smoking, as this exposure is associated with many diseases of the respiratory system, most notably COPD and bronchogenic lung cancer but also a variety of diffuse parenchymal lung diseases (e.g., desquamative interstitial pneumonitis and pulmonary Langerhans cell histiocytosis). For most disorders, longer duration and greater intensity of exposure to cigarette smoke increases the risk of disease. There is growing evidence that “second-hand smoke” is also a risk factor for respiratory tract pathology; for this reason, patients should be asked about parents, spouses, or housemates who smoke. Possible inhalational exposures should be explored, including those at the work place (e.g., asbestos, wood smoke) and those associated with leisure (e.g., excrement from pet birds) (Chap. 311). Travel predisposes to certain infections of the respiratory tract, most notably the risk of tuberculosis. Potential exposure to fungi found in specific geographic regions or climates (e.g., *Histoplasma capsulatum*) should be explored.

Associated symptoms of fever and chills should raise the suspicion of infective etiologies, both pulmonary and systemic. A comprehensive review of systems may suggest rheumatologic or autoimmune disease presenting with respiratory tract manifestations. Questions should focus on joint pain or swelling, rashes, dry eyes, dry mouth, or constitutional symptoms. In addition, carcinomas from a variety of primary sources commonly metastasize to the lung and cause respiratory symptoms. Finally, therapy for other conditions, including both irradiation and medications, can result in diseases of the chest.

Physical Examination The clinician’s suspicion of respiratory disease often begins with a patient’s vital signs. The respiratory rate is often informative, whether elevated (tachypnea) or depressed (hypopnea). In addition, pulse oximetry should be measured, as many patients with respiratory disease have hypoxemia, either at rest or with exertion. The classic structure of the respiratory examination proceeds through inspection, percussion, palpation, and auscultation as described below. Often, however, auscultatory findings will lead the clinician to perform further percussion or palpation in order to clarify these findings.

The first step of the physical examination is inspection. Patients with respiratory disease may be in distress, often using accessory muscles of respiration to breathe. Severe kyphoscoliosis can result in restrictive pathophysiology. Inability to complete a sentence in conversation is generally a sign of severe impairment and should result in an expedited evaluation of the patient.

Percussion of the chest is used to establish diaphragm excursion and lung size. In the setting of decreased breath sounds, percussion is used to distinguish between pleural effusions (dull to percussion) and pneumothorax (hyper-resonant note).

The role of palpation is limited in the respiratory examination. Palpation can demonstrate subcutaneous air in the setting of barotrauma. It can also be used as an adjunctive assessment to determine whether an area of decreased breath sounds is due to consolidation (increased tactile fremitus) or a pleural effusion (decreased tactile fremitus).

The majority of the manifestations of respiratory disease present as abnormalities of auscultation. Wheezes are a manifestation of airway obstruction. While most commonly a sign of asthma, peribronchial edema in the setting of congestive heart failure can also result in diffuse wheezes, as can any other process that causes narrowing of small airways. For this reason, clinicians must take care not to attribute all wheezing to asthma.

Rhonchi are a manifestation of obstruction of medium-sized airways, most often with secretions. In the acute setting, this manifestation may be a sign of viral or bacterial bronchitis. Chronic rhonchi suggest bronchiectasis or COPD. Stridor, a high-pitched, focal inspiratory wheeze, usually heard over the neck, is a manifestation of upper airway obstruction and should prompt expedited evaluation of the patient,

as it can precede complete upper airway obstruction and respiratory failure.

Crackles, or rales, are commonly a sign of alveolar disease. A variety of processes that fill the alveoli with fluid may result in crackles. Pneumonia can cause focal crackles. Pulmonary edema is associated with crackles, generally more prominent at the bases. Interestingly, diseases that result in fibrosis of the interstitium (e.g., IPF) also result in crackles often sounding like Velcro being ripped apart. Although some clinicians make a distinction between “wet” and “dry” crackles, this distinction has not been shown to be a reliable way to differentiate among etiologies of respiratory disease.

One way to help distinguish between crackles associated with alveolar fluid and those associated with interstitial fibrosis is to assess for egophony. *Egophony* is the auscultation of the sound “AH” instead of “EEE” when a patient phonates “EEE.” This change in note is due to abnormal sound transmission through consolidated parenchyma and is present in pneumonia but not in IPF. Similarly, areas of alveolar filling have increased whispered *pectoriloquy* as well as transmission of larger-airway sounds (i.e., bronchial breath sounds in a lung zone where vesicular breath sounds are expected).

The lack or diminution of breath sounds can also help determine the etiology of respiratory disease. Patients with emphysema often have a quiet chest with diffusely decreased breath sounds. A pneumothorax or pleural effusion may present with an area of absent breath sounds.

Other Systems Pedal edema, if symmetric, may suggest cor pulmonale; if asymmetric, it may be due to deep venous thrombosis and associated pulmonary embolism. Jugular venous distention may also be a sign of volume overload associated with right heart failure. *Pulsus paradoxus* is an ominous sign in a patient with obstructive lung disease, as it is associated with significant negative intrathoracic (pleural) pressures required for ventilation and impending respiratory failure.

As stated earlier, rheumatologic disease may manifest primarily as lung disease. Owing to this association, particular attention should be paid to joint and skin examination. Clubbing can be found in many lung diseases, including cystic fibrosis, IPF, and lung cancer. Cyanosis is seen in hypoxemic respiratory disorders that result in >5 g of deoxygenated hemoglobin/dL.

DIAGNOSTIC EVALUATION

The sequence of studies is dictated by the clinician’s differential diagnosis, as determined by the history and physical examination. Acute respiratory symptoms are often evaluated with multiple tests performed at the same time in order to diagnose any life-threatening diseases rapidly (e.g., pulmonary embolism or multilobar pneumonia). In contrast, chronic dyspnea and cough can be evaluated in a more protracted, stepwise fashion.

Pulmonary Function Testing (See also Chap. 307) The initial pulmonary function test obtained is spirometry. This study is an effort-dependent test used to assess for obstructive pathophysiology as seen in asthma, COPD, and bronchiectasis. A diminished-forced expiratory volume in 1 sec (FEV_1)/forced vital capacity (FVC) (often defined as <70% of the predicted value) is diagnostic of obstruction. In addition to measuring FEV_1 and FVC, the clinician should examine the flow-volume loop (which is effort-independent). A plateau of the inspiratory and expiratory curves suggests large-airway obstruction in extrathoracic and intrathoracic locations, respectively.

Spirometry with symmetric decreases in FEV_1 and FVC warrants further testing, including measurement of lung volumes and the diffusion capacity of the lung for carbon monoxide (D_LCO). A total lung capacity <80% of the predicted value for a patient’s age, race, sex, and height defines restrictive pathophysiology. Restriction can result from parenchymal disease, neuromuscular weakness, or chest wall or pleural diseases. Restriction with impaired gas exchange, as indicated by a decreased D_LCO , suggests parenchymal lung disease. Additional testing, such as measurements of maximal expiratory pressure and maximal inspiratory pressure, can help diagnose neuromuscular weakness. Normal spirometry, normal lung volumes, and a low D_LCO should prompt further evaluation for pulmonary vascular disease.

Arterial blood gas testing is often helpful in assessing respiratory disease. Hypoxemia, while usually apparent with pulse oximetry, can be further evaluated with the measurement of arterial PO_2 and the calculation of an alveolar gas and arterial blood oxygen tension difference ($[A-a]DO_2$). Patients with diseases that cause ventilation-perfusion mismatch or shunt physiology have an increased $(A-a)DO_2$ at rest. Arterial blood gas testing also allows the measurement of arterial PCO_2 . Hypercarbia can accompany severe airway obstruction (e.g., COPD) or progressive restrictive physiology, as in patients with neuromuscular weakness.

Chest Imaging (See Chap. 308e) Most patients with disease of the respiratory system undergo imaging of the chest as part of the initial evaluation. Clinicians should generally begin with a plain chest radiograph, preferably posterior-anterior and lateral films. Several findings, including opacities of the parenchyma, blunting of the costophrenic angles, mass lesions, and volume loss, can be very helpful in determining an etiology. However, many diseases of the respiratory system, particularly those of the airways and pulmonary vasculature, are associated with a normal chest radiograph.

CT of the chest is often performed subsequently and allows better delineation of parenchymal processes, pleural disease, masses or nodules, and large airways. If the test includes administration of contrast, the pulmonary vasculature can be assessed with particular utility for determination of pulmonary emboli. Intravenous contrast also allows lymph nodes to be delineated in greater detail.

FURTHER STUDIES

Depending on the clinician's suspicion, a variety of other studies may be done. Concern about large-airway lesions may warrant bronchoscopy. This procedure may also be used to sample the alveolar space with bronchoalveolar lavage or to obtain nonsurgical lung biopsies. Blood testing may include assessment for hypercoagulable states in the setting of pulmonary vascular disease, serologic testing for infectious or rheumatologic disease, or assessment of inflammatory markers or leukocyte counts (e.g., eosinophils). Sputum evaluation for malignant cells or microorganisms may be appropriate. An echocardiogram to assess right- and left-sided heart function is often obtained. Finally, at times, a surgical lung biopsy is needed to diagnose certain diseases of the respiratory system. All of these studies will be guided by the preceding history, physical examination, pulmonary function testing, and chest imaging.

The primary functions of the respiratory system—to oxygenate blood and eliminate carbon dioxide—require virtual contact between blood and fresh air, which facilitates diffusion of respiratory gases between blood and gas. This process occurs in the lung alveoli, where blood flowing through alveolar wall capillaries is separated from alveolar gas by an extremely thin membrane of flattened endothelial and epithelial cells, across which respiratory gases diffuse and equilibrate. Blood flow through the lung is unidirectional via a continuous vascular path, along which venous blood absorbs oxygen from and loses CO_2 to inspired gas. The path for airflow, in contrast, reaches a dead end at the alveolar walls; thus the alveolar space must be ventilated tidally, with inflow of fresh gas and outflow of alveolar gas alternating periodically at the respiratory rate (RR). To provide an enormous alveolar surface area (typically 70 m^2) for blood-gas diffusion within the modest volume of a thoracic cavity (typically 7 L), nature has distributed both blood flow and ventilation among millions of tiny alveoli through multigenerational branching of both pulmonary arteries and bronchial airways. As a consequence of variations in tube lengths and calibers along these pathways as well as the effects of gravity, tidal pressure fluctuations, and anatomic constraints from the chest wall, the alveoli vary in their relative ventilations and perfusions. Not surprisingly, for the lung to be most efficient in exchanging gas, the fresh gas ventilation of a given alveolus must be matched to its perfusion.

For the respiratory system to succeed in oxygenating blood and eliminating CO_2 , it must be able to ventilate the lung tidally and thus to freshen alveolar gas; it must provide for perfusion of the individual alveolus in a manner proportional to its ventilation; and it must allow adequate diffusion of respiratory gases between alveolar gas and capillary blood. Furthermore, it must accommodate severalfold increases in the demand for oxygen uptake or CO_2 elimination imposed by metabolic needs or acid-base derangement. Given these multiple requirements for normal operation, it is not surprising that many diseases disturb respiratory function. This chapter considers in some detail the physiologic determinants of lung ventilation and perfusion, elucidates how the matching distributions of these processes and rapid gas diffusion allow normal gas exchange, and discusses how common diseases derange these normal functions, thereby impairing gas exchange—or at least increasing the work required by the respiratory muscles or heart to maintain adequate respiratory function.

VENTILATION

It is useful to think about the respiratory system as three independently functioning components: the lung, including its airways; the neuromuscular system; and the chest wall, which includes everything that is not lung or active neuromuscular system. Accordingly, the mass of the respiratory muscles is part of the chest wall, while the force these muscles generate is part of the neuromuscular system; the abdomen (especially an obese abdomen) and the heart (especially an enlarged heart) are, for these purposes, part of the chest wall. Each of these three components has mechanical properties that

relate to its enclosed volume (or—in the case of the neuromuscular system—the respiratory system volume at which it is operating) and to the rate of change of its volume (i.e., flow).

Volume-Related Mechanical Properties—Statics Figure 306e-1 shows the volume-related properties of each component of the respiratory system. Due both to surface tension at the air-liquid interface between alveolar wall lining fluid and alveolar gas and to elastic recoil of the lung tissue itself, the lung requires a positive transmural pressure difference between alveolar gas and its pleural surface to stay inflated; this difference is called the *elastic recoil pressure* of the lung, and it increases with lung volume. The lung becomes rather stiff at high volumes, so that relatively small volume changes are accompanied by large changes in transpulmonary pressure; in contrast, the lung is compliant at lower volumes, including those at which tidal breathing normally occurs. At zero inflation pressure, even normal lungs retain some air in the alveoli because the small peripheral airways are tethered open by radially outward pull from inflated lung parenchyma attached to adventitia; as the lung deflates during exhalation, those small airways are pulled open progressively less, and eventually they close, trapping some gas in the alveoli. This effect can be exaggerated with age and especially with obstructive airway diseases, resulting in gas trapping at quite large lung volumes.

The elastic behavior of the passive chest wall (i.e., in the absence of neuromuscular activation) differs markedly from that of the lung. Whereas the lung tends toward full deflation with no distending (transmural) pressure, the chest wall encloses a large volume when pleural pressure equals body surface (atmospheric) pressure. Furthermore, the chest wall is compliant at high enclosed volumes, readily expanding even further in response to increases in transmural pressure. The chest wall also remains compliant at small negative transmural pressures (i.e., when pleural pressure falls slightly below atmospheric pressure), but as the volume enclosed by the chest wall becomes quite small in response to large negative transmural pressures, the passive chest wall becomes stiff due to squeezing together of ribs and intercostal muscles, diaphragm stretch, displacement of abdominal contents, and straining of ligaments and bony articulations. Under normal circumstances, the lung and the passive chest wall enclose essentially the same volume, the only difference being the volumes of the pleural fluid and of the lung parenchyma (both quite small). For this reason and because the lung and chest wall function in mechanical series, the pressure required to displace the passive respiratory system (lungs plus chest wall) at any volume is simply the sum of the elastic recoil pressure of the lungs and the transmural pressure across the chest wall. When plotted against respiratory system volume, this relationship assumes a sigmoid shape, exhibiting stiffness at high lung volumes (imparted by the lung), stiffness at low lung volumes (imparted by the chest wall or sometimes by airway closure), and compliance in the middle range of lung volumes.

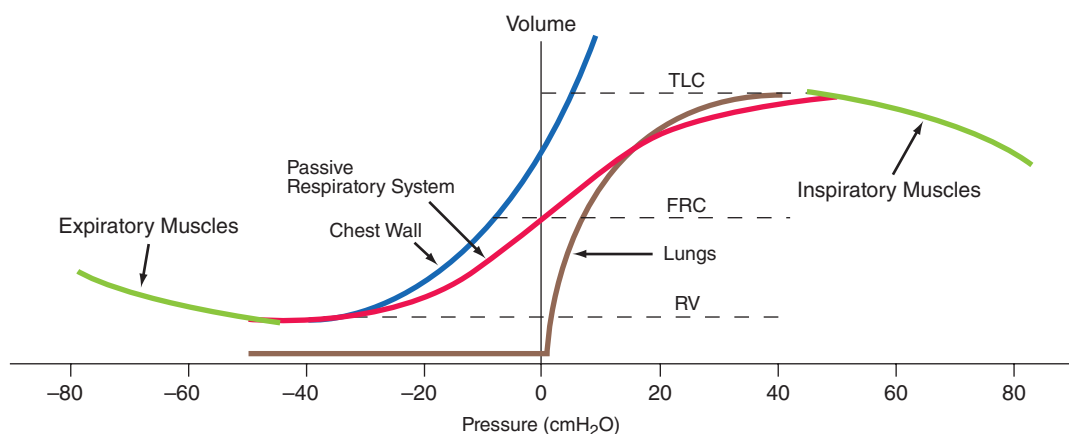


FIGURE 306e-1 Pressure-volume curves of the isolated lung, isolated chest wall, combined respiratory system, inspiratory muscles, and expiratory muscles. FRC, functional residual capacity; RV, residual volume; TLC, total lung capacity.

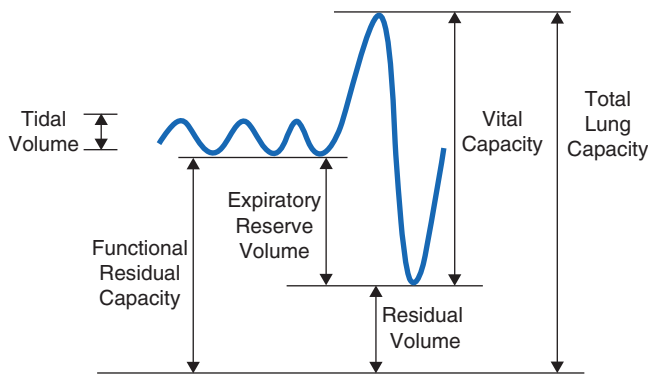


FIGURE 306e-2 Spirogram demonstrating a slow vital capacity maneuver and various lung volumes.

In addition, a passive resting point of the respiratory system is attained when alveolar gas pressure equals body surface pressure (i.e., when the transrespiratory system pressure is zero). At this volume (called the *functional residual capacity* [FRC]), the outward recoil of the chest wall is balanced exactly by the inward recoil of the lung. As these recoils are transmitted through the pleural fluid, the lung is pulled both outward and inward simultaneously at FRC, and thus its pressure falls below atmospheric pressure (typically, $-5 \text{ cmH}_2\text{O}$).

The normal passive respiratory system would equilibrate at the FRC and remain there were it not for the actions of respiratory muscles. The inspiratory muscles act on the chest wall to generate the equivalent of positive pressure across the lungs and passive chest wall, while the expiratory muscles generate the equivalent of negative transrespiratory pressure. The maximal pressures these sets of muscles can generate vary with the lung volume at which they operate. This variation is due to length-tension relationships in striated muscle sarcomeres and to changes in mechanical advantage as the angles of insertion change with lung volume (Fig. 306e-1). Nonetheless, under normal conditions, the respiratory muscles are substantially “overpowered” for their roles and generate more than adequate force to drive the respiratory system to its stiffness extremes, as determined by the lung (total lung capacity [TLC]) or by chest wall or airway closure (residual volume [RV]); the airway closure always prevents the adult lung from emptying completely under normal circumstances. The excursion between full and minimal lung inflation is called *vital capacity* (VC; Fig. 306e-2) and is readily seen to be the difference between volumes at two unrelated stiffness extremes—one determined by the lung (TLC) and the other by the chest wall or airways (RV). Thus, although VC is easy to measure (see below), it provides little information about the intrinsic properties of the respiratory system. As will become clear, it is much more useful for the clinician to consider TLC and RV individually.

Flow-Related Mechanical Properties—Dynamics The passive chest wall and active neuromuscular system do exhibit mechanical behaviors related to the rate of change of volume, but these behaviors become quantitatively important only at markedly supraphysiologic breathing frequencies (e.g., during high-frequency mechanical ventilation) and thus will not be addressed here. In contrast, the dynamic airflow properties of the lung substantially affect its ability to ventilate and contribute importantly to the work of breathing, and these properties are often deranged by disease. Understanding dynamic airflow properties is therefore worthwhile.

As with the flow of any fluid (gas or liquid) in any tube, maintenance of airflow within the pulmonary airways requires a pressure gradient that falls along the direction of flow, the magnitude of which is determined by the flow rate and the frictional resistance to flow. During quiet tidal breathing, the pressure gradients driving inspiratory or expiratory flow are small owing to the very low frictional resistance of normal pulmonary airways (R_{aw} , normally $<2 \text{ cmH}_2\text{O/L}$ per second). However, during rapid exhalation, another phenomenon reduces flow below that which would have been expected if frictional resistance were the only impediment to flow. This phenomenon is called *dynamic*

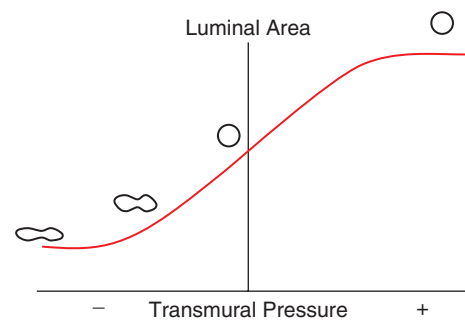


FIGURE 306e-3 Luminal area versus transmural pressure relationship. Transmural pressure represents the pressure difference across the airway wall from inside to outside.

airflow limitation, and it occurs because the bronchial airways through which air is exhaled are collapsible rather than rigid (Fig. 306e-3). An important anatomic feature of the pulmonary airways is its treelike branching structure. While the individual airways in each successive generation, from most proximal (trachea) to most distal (respiratory bronchioles), are smaller than those of the parent generation, their number increases exponentially such that the summed cross-sectional area of the airways becomes very large toward the lung periphery. Because flow (volume/time) is constant along the airway tree, the velocity of airflow (flow/summed cross-sectional area) is much greater in the central airways than in the peripheral airways. During exhalation, gas leaving the alveoli must therefore gain velocity as it proceeds toward the mouth. The energy required for this “convective” acceleration is drawn from the component of gas energy manifested as its local pressure, which reduces intraluminal gas pressure, airway transmural pressure, airway size (Fig. 306e-3), and flow. This is the Bernoulli effect, the same effect that keeps an airplane airborne, generating a lifting force by decreasing pressure above the curved upper surface of the wing due to acceleration of air flowing over the wing. If an individual tries to exhale more forcefully, the local velocity increases further and reduces airway size further, resulting in no net increase in flow. Under these circumstances, flow has reached its maximum possible value, or its *flow limit*. Lungs normally exhibit such dynamic airflow limitation. This limitation can be assessed by spirometry, in which an individual inhales fully to TLC and then forcibly exhales to RV. One useful spirometric measure is the volume of air exhaled during the first second of expiration (FEV_1), as discussed later. Maximal expiratory flow at any lung volume is determined by gas density, airway cross-section and distensibility, elastic recoil pressure of the lung, and frictional pressure loss to the flow-limiting airway site. Under normal conditions, maximal expiratory flow falls with lung volume (Fig. 306e-4), primarily because of the dependence of lung recoil pressure on lung volume (Fig. 306e-1). In pulmonary fibrosis, lung recoil pressure is increased at any lung volume, and thus the maximal expiratory flow is elevated when considered in relation to lung volume. Conversely, in emphysema, lung recoil pressure is reduced; this reduction is a principal mechanism by which maximal expiratory flows fall. Diseases that narrow the airway lumen at any transmural pressure (e.g., asthma or chronic bronchitis) or that cause excessive airway collapsibility (e.g., tracheomalacia) also reduce maximal expiratory flow.

The Bernoulli effect also applies during inspiration, but the more negative pleural pressures during inspiration lower the pressure outside of airways, thereby increasing transmural pressure and promoting airway expansion. Thus inspiratory airflow limitation seldom occurs due to diffuse pulmonary airway disease. Conversely, extrathoracic airway narrowing (e.g., due to a tracheal adenoma or post-tracheostomy stricture) can lead to inspiratory airflow limitation (Fig. 306e-4).

The Work of Breathing In health, the elastic (volume change-related) and dynamic (flow-related) loads that must be overcome to ventilate the lungs at rest are small, and the work required of the respiratory muscles is minimal. However, the work of breathing can

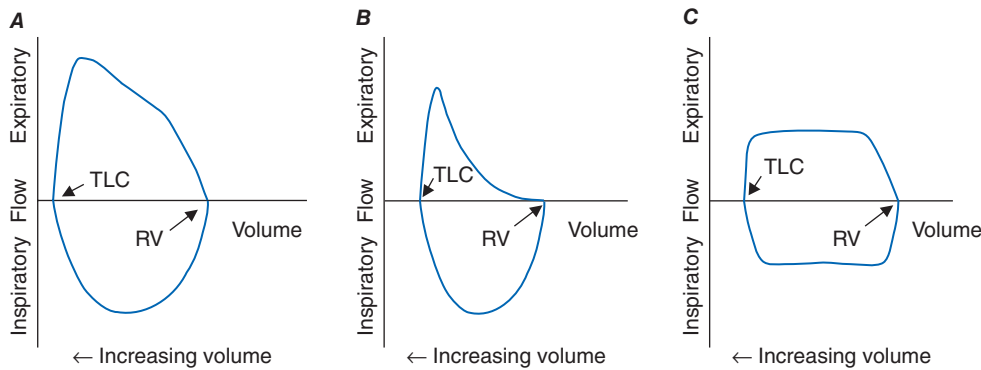


FIGURE 306e-4 Flow-volume loops. **A.** Normal. **B.** Airflow obstruction. **C.** Fixed central airway obstruction. RV, residual volume; TLC, total lung capacity.

increase considerably due to a metabolic requirement for substantially increased ventilation, an abnormally increased mechanical load, or both. As discussed below, the rate of ventilation is primarily set by the need to eliminate carbon dioxide, and thus ventilation increases during exercise (sometimes by more than twentyfold) and during metabolic acidosis as a compensatory response. Naturally, the work rate required to overcome the elasticity of the respiratory system increases with both the depth and the frequency of tidal breaths, while the work required to overcome the dynamic load increases with total ventilation. A modest increase of ventilation is most efficiently achieved by increasing tidal volume but not respiratory rate, which is the normal ventilatory response to lower-level exercise. At high levels of exercise, deep breathing persists, but respiratory rate also increases. The pattern chosen by the respiratory controller minimizes the work of breathing.

The work of breathing also increases when disease reduces the compliance of the respiratory system or increases the resistance to airflow. The former occurs commonly in diseases of the lung parenchyma (interstitial processes or fibrosis, alveolar filling diseases such as pulmonary edema or pneumonia, or substantial lung resection), and the latter occurs in obstructive airway diseases such as asthma, chronic bronchitis, emphysema, and cystic fibrosis. Furthermore, severe airflow obstruction can functionally reduce the compliance of the respiratory system by leading to dynamic hyperinflation. In this scenario, expiratory flows slowed by the obstructive airways disease may be insufficient to allow complete exhalation during the expiratory phase of tidal breathing; as a result, the “functional residual capacity” from which the next breath is inhaled is greater than the static FRC. With repetition of incomplete exhalations of each tidal breath, the operating FRC becomes dynamically elevated, sometimes to a level that approaches TLC. At these high lung volumes, the respiratory system is much less compliant than at normal breathing volumes, and thus the elastic work of each tidal breath is also increased. The dynamic pulmonary hyperinflation that accompanies severe airflow obstruction causes patients to sense difficulty in inhaling—even though the root cause of this pathophysiologic abnormality is expiratory airflow obstruction.

Adequacy of Ventilation As noted above, the respiratory control system that sets the rate of ventilation responds to chemical signals, including arterial CO_2 and oxygen tensions and blood pH, and to volitional needs, such as the need to inhale deeply before playing a long phrase on the trumpet. Disturbances in ventilation are discussed in [Chap. 318](#). The focus of this chapter is on the relationship between ventilation of the lung and CO_2 elimination.

At the end of each tidal exhalation, the conducting airways are filled with alveolar gas that had not reached the mouth when expiratory flow stopped. During the ensuing inhalation, fresh gas immediately enters the airway tree at the mouth, but the gas first entering the alveoli at the start of inhalation is that same alveolar gas in the conducting airways that had just left the alveoli. Accordingly, fresh gas does not enter the alveoli until the volume of the conducting airways has been inspired.

This volume is called the *anatomic dead space* (V_D). Quiet breathing with tidal volumes smaller than the anatomic dead space introduces no fresh gas into the alveoli at all; only that part of the inspired tidal volume (V_T) that is greater than the V_D introduces fresh gas into the alveoli. The dead space can be further increased functionally if some of the inspired tidal volume is delivered to a part of the lung that receives no pulmonary blood flow and thus cannot contribute to gas exchange (e.g., the portion of the lung distal to a large pulmonary embolus). In this situation, exhaled minute ventilation ($\dot{V}_E = V_T \times \text{RR}$) includes a component of dead space ventilation ($\dot{V}_D = V_D \times \text{RR}$) and

a component of fresh gas alveolar ventilation ($\dot{V}_A = [V_T - V_D] \times \text{RR}$). CO_2 elimination from the alveoli is equal to \dot{V}_A times the difference in CO_2 fraction between inspired air (essentially zero) and alveolar gas (typically ~5.6% after correction for humidification of inspired air, corresponding to 40 mmHg). In the steady state, the alveolar fraction of CO_2 is equal to metabolic CO_2 production divided by alveolar ventilation. Because, as discussed below, alveolar and arterial CO_2 tensions are equal, and because the respiratory controller normally strives to maintain arterial P_{CO_2} (P_{aCO_2}) at ~40 mmHg, the adequacy of alveolar ventilation is reflected in P_{aCO_2} . If the P_{aCO_2} falls much below 40 mmHg, alveolar hyperventilation is present; if the P_{aCO_2} exceeds 40 mmHg, then alveolar hypoventilation is present. Ventilatory failure is characterized by extreme alveolar hypoventilation.

As a consequence of oxygen uptake of alveolar gas into capillary blood, alveolar oxygen tension falls below that of inspired gas. The rate of oxygen uptake (determined by the body’s metabolic oxygen consumption) is related to the average rate of metabolic CO_2 production, and their ratio—the “respiratory quotient” ($R = \dot{V}_{\text{CO}_2} / \dot{V}_{\text{O}_2}$)—depends largely on the fuel being metabolized. For a typical American diet, R is usually around 0.85, and more oxygen is absorbed than CO_2 is excreted. Together, these phenomena allow the estimation of alveolar oxygen tension, according to the following relationship, known as the *alveolar gas equation*:

$$P_{\text{A}_{\text{O}_2}} = F_{\text{I}_{\text{O}_2}} \times (P_{\text{bar}} - P_{\text{H}_2\text{O}}) - P_{\text{A}_{\text{CO}_2}} / R$$

The alveolar gas equation also highlights the influences of inspired oxygen fraction ($F_{\text{I}_{\text{O}_2}}$), barometric pressure (P_{bar}), and vapor pressure of water ($P_{\text{H}_2\text{O}} = 47$ mmHg at 37°C) in addition to alveolar ventilation (which sets $P_{\text{A}_{\text{CO}_2}}$) in determining $P_{\text{A}_{\text{O}_2}}$. An implication of the alveolar gas equation is that severe arterial hypoxemia rarely occurs as a pure consequence of alveolar hypoventilation at sea level while an individual is breathing air. The potential for alveolar hypoventilation to induce severe hypoxemia with otherwise normal lungs increases as P_{bar} falls with increasing altitude.

GAS EXCHANGE

Diffusion For oxygen to be delivered to the peripheral tissues, it must pass from alveolar gas into alveolar capillary blood by diffusing through alveolar membrane. The aggregate alveolar membrane is highly optimized for this process, with a very large surface area and minimal thickness. Diffusion through the alveolar membrane is so efficient in the human lung that in most circumstances a red blood cell’s hemoglobin becomes fully oxygen saturated by the time the cell has traveled just one-third the length of the alveolar capillary. Thus the uptake of alveolar oxygen is ordinarily limited by the amount of blood transiting the alveolar capillaries rather than by the rapidity with which oxygen can diffuse across the membrane; consequently, oxygen uptake from the lung is said to be “perfusion limited.” CO_2 also equilibrates rapidly across the alveolar membrane. Therefore, the oxygen and CO_2 tensions in capillary blood leaving a normal alveolus are essentially equal to those in alveolar gas. Only in rare circumstances (e.g., at high

306e-4 altitude or in high-performance athletes exerting maximal effort) is oxygen uptake from normal lungs diffusion limited. Diffusion limitation can also occur in interstitial lung disease if substantially thickened alveolar walls remain perfused.

Ventilation/Perfusion Heterogeneity As noted above, for gas exchange to be most efficient, ventilation to each individual alveolus (among the millions of alveoli) should match perfusion to its accompanying capillaries. Because of the differential effects of gravity on lung mechanics and blood flow throughout the lung and because of differences in airway and vascular architecture among various respiratory paths, there is minor ventilation/perfusion heterogeneity even in the normal lung; however, \dot{V}/\dot{Q} heterogeneity can be particularly marked in disease. Two extreme examples are (1) ventilation of unperfused lung distal to a pulmonary embolus, in which ventilation of the physiologic dead space is “wasted” in the sense that it does not contribute to gas exchange; and (2) perfusion of nonventilated lung (a “shunt”), which allows venous blood to pass through the lung unaltered. When mixed with fully oxygenated blood leaving other well-ventilated lung units, shunted venous blood disproportionately lowers the mixed arterial $P_{a_{O_2}}$ as a result of the nonlinear oxygen content versus P_{O_2} relationship of hemoglobin (Fig. 306e-5). Furthermore, the resulting arterial hypoxemia is refractory to supplemental inspired oxygen. The reason is that (1) raising the inspired F_{iO_2} has no effect on alveolar gas tensions in nonventilated alveoli and (2) while raising inspired F_{iO_2} does increase $P_{A_{CO_2}}$ in ventilated alveoli, the oxygen content of blood

exiting ventilated units increases only slightly, as hemoglobin will already have been nearly fully saturated and the solubility of oxygen in plasma is quite small.

A more common occurrence than the two extreme examples given above is a widening of the distribution of ventilation/perfusion ratios; such \dot{V}/\dot{Q} heterogeneity is a common consequence of lung disease. In this circumstance, perfusion of relatively underventilated alveoli results in the incomplete oxygenation of exiting blood. When mixed with well-oxygenated blood leaving higher \dot{V}/\dot{Q} regions, this partially reoxygenated blood disproportionately lowers arterial $P_{a_{O_2}}$, although to a lesser extent than does a similar perfusion fraction of blood leaving regions of pure shunt. In addition, in contrast to shunt regions, inhalation of supplemental oxygen does raise the $P_{A_{O_2}}$, even in relatively underventilated low \dot{V}/\dot{Q} regions, and so the arterial hypoxemia induced by \dot{V}/\dot{Q} heterogeneity is typically responsive to oxygen therapy (Fig. 306e-5).

In sum, arterial hypoxemia can be caused by substantial reduction of inspired oxygen tension; by severe alveolar hypoventilation; by perfusion of relatively underventilated (low \dot{V}/\dot{Q}) or completely unventilated (shunt) lung regions; and, in unusual circumstances, by limitation of gas diffusion.

PATHOPHYSIOLOGY

Although many diseases injure the respiratory system, this system responds to injury in relatively few ways. For this reason, the pattern of

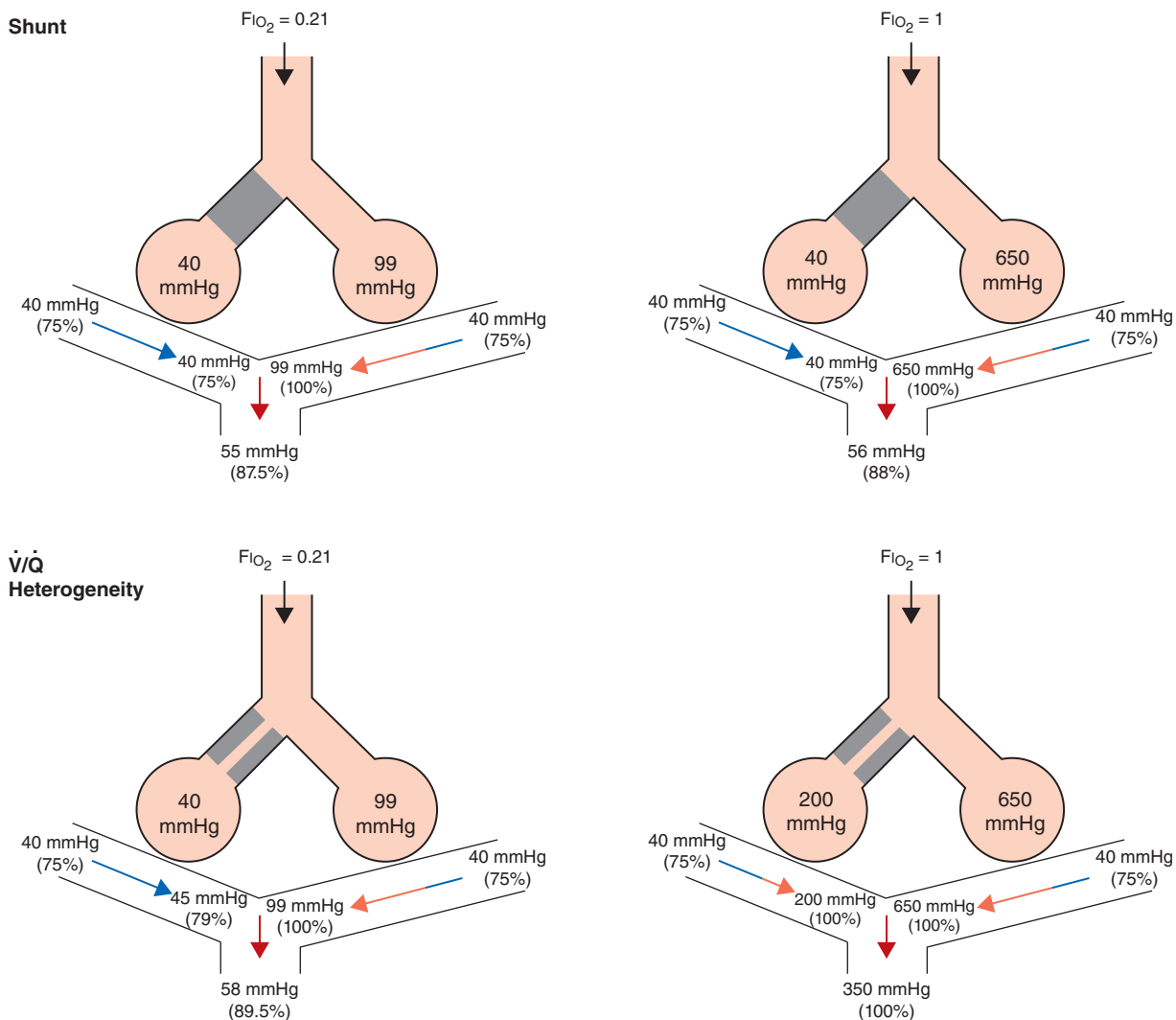


FIGURE 306e-5 Influence of air versus oxygen breathing on mixed arterial oxygenation in shunt and ventilation/perfusion heterogeneity. Partial pressure of oxygen (mmHg) and oxygen saturations are shown for mixed venous blood, for end capillary blood (normal versus affected alveoli), and for mixed arterial blood. F_{iO_2} , fraction of inspired oxygen; \dot{V}/\dot{Q} , ventilation/perfusion.

	Restriction due to increased lung elastic recoil (pulmonary fibrosis)	Restriction due to chest wall abnormality (moderate obesity)	Restriction due to respiratory muscle weakness (myasthenia gravis)	Obstruction due to airway narrowing (acute asthma)	Obstruction due to decreased elastic recoil (severe emphysema)
TLC	60%	95%	75%	100%	130%
FRC	60%	65%	100%	104%	220%
RV	60%	100%	120%	120%	310%
FVC	60%	92%	60%	90%	60%
FEV ₁	75%	92%	60%	35% pre-b.d. 75% post-b.d.	35% pre-b.d. 38% post-b.d.
R _{aw}	1.0	1.0	1.0	2.5	1.5
DL _{CO}	60%	95%	80%	120%	40%

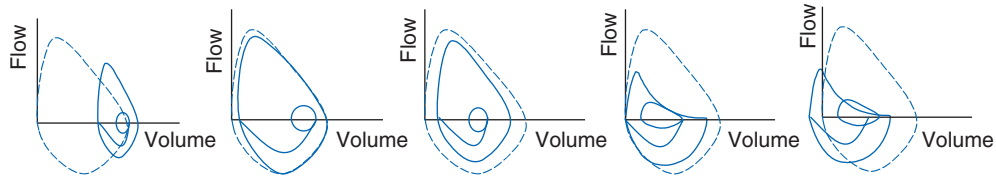


FIGURE 306e-6 Common abnormalities of pulmonary function (see text). Pulmonary function values are expressed as a percentage of normal predicted values, except for R_{aw} , which is expressed as $\text{cmH}_2\text{O/L}$ per sec (normal, $<2 \text{ cmH}_2\text{O/L}$ per second). The figures at the bottom of each column show the typical configuration of flow-volume loops in each condition, including the flow-volume relationship during tidal breathing. b.d., bronchodilator; DL_{CO} , diffusion capacity of lung for carbon monoxide; FEV₁, forced expiratory volume in 1 sec; FRC, functional residual capacity; FVC, forced vital capacity; R_{aw} , airways resistance; RV, residual volume; TLC, total lung capacity.

physiologic abnormalities may or may not provide sufficient information by which to discriminate among conditions.

Figure 306e-6 lists abnormalities in pulmonary function testing that are typically found in a number of common respiratory disorders and highlights the simultaneous occurrence of multiple physiologic abnormalities. The coexistence of some of these respiratory disorders results in more complex superposition of these abnormalities. Methods to measure respiratory system function clinically are described later in this chapter.

Ventilatory Restriction Due to Increased Elastic Recoil—Example: Idiopathic Pulmonary Fibrosis Idiopathic pulmonary fibrosis raises lung recoil at all lung volumes, thereby lowering TLC, FRC, and RV as well as forced vital capacity (FVC). Maximal expiratory flows are also reduced from normal values but are elevated when considered in relation to lung volumes. Increased flow occurs both because the increased lung recoil drives greater maximal flow at any lung volume and because airway diameters are relatively increased due to greater radially outward traction exerted on bronchi by the stiff lung parenchyma. For the same reason, airway resistance is also normal. Destruction of the pulmonary capillaries by the fibrotic process results in a marked reduction in diffusing capacity (see below). Oxygenation is often severely reduced by persistent perfusion of alveolar units that are relatively underventilated due to fibrosis of nearby (and mechanically linked) lung. The flow-volume loop (see below) looks like a miniature version of a normal loop but is shifted toward lower absolute lung volumes and displays maximal expiratory flows that are increased for any given volume over the normal tracing.

Ventilatory Restriction Due to Chest Wall Abnormality—Example: Moderate Obesity As the size of the average American continues to increase, this pattern may become the most common of pulmonary function abnormalities. In moderate obesity, the outward recoil of the chest wall is blunted by the weight of chest wall fat and the space occupied by intraabdominal fat. In this situation, preserved inward recoil of the lung overbalances the reduced outward recoil of the chest wall, and FRC falls. Because respiratory muscle strength and lung recoil remain normal, TLC is typically unchanged (although it may fall in massive obesity) and RV is normal (but may be reduced in massive obesity).

Mild hypoxemia may be present due to perfusion of alveolar units that are poorly ventilated because of airway closure in dependent portions of the lung during breathing near the reduced FRC. Flows remain normal, as does the diffusion capacity of the lung for carbon monoxide (DL_{CO}), unless obstructive sleep apnea (which often accompanies obesity) and associated chronic intermittent hypoxemia have induced pulmonary arterial hypertension, in which case DL_{CO} may be low.

Ventilatory Restriction Due to Reduced Muscle Strength—Example: Myasthenia Gravis In this circumstance, FRC remains normal, as both lung recoil and passive chest wall recoil are normal. However, TLC is low and RV is elevated because respiratory muscle strength is insufficient to push the passive respiratory system fully toward either volume extreme. Caught between the low TLC and the elevated RV, FVC and FEV₁ are reduced as “innocent bystanders.” As airway size and lung vasculature are unaffected, both R_{aw} and DL_{CO} are normal. Oxygenation is normal unless weakness becomes so severe that the patient has insufficient strength to reopen collapsed alveoli during sighs, with resulting atelectasis.

Airflow Obstruction Due to Decreased Airway Diameter—Example: Acute Asthma During an episode of acute asthma, luminal narrowing due to smooth muscle constriction as well as inflammation and thickening within the small- and medium-sized bronchi raise frictional resistance and reduce airflow. “Scooping” of the flow-volume loop is caused by reduction of airflow, especially at lower lung volumes. Often, airflow obstruction can be reversed by inhalation of β_2 -adrenergic agonists acutely or by treatment with inhaled steroids chronically. TLC usually remains normal (although elevated TLC is sometimes seen in long-standing asthma), but FRC may be dynamically elevated. RV is often increased due to exaggerated airway closure at low lung volumes, and this elevation of RV reduces FVC. Because central airways are narrowed, R_{aw} is usually elevated. Mild arterial hypoxemia is often present due to perfusion of relatively underventilated alveoli distal to obstructed airways (and is responsive to oxygen supplementation), but DL_{CO} is normal or mildly elevated.

Airflow Obstruction Due to Decreased Elastic Recoil—Example: Severe Emphysema Loss of lung elastic recoil in severe emphysema results in

pulmonary hyperinflation, of which elevated TLC is the hallmark. FRC is more severely elevated due both to loss of lung elastic recoil and to dynamic hyperinflation—the same phenomenon as autoPEEP, which is the positive end-expiratory alveolar pressure that occurs when a new breath is initiated before the lung volume is allowed to return to FRC. Residual volume is very severely elevated because of airway closure and because exhalation toward RV may take so long that RV cannot be reached before the patient must inhale again. Both FVC and FEV_1 are markedly decreased, the former because of the severe elevation of RV and the latter because loss of lung elastic recoil reduces the pressure driving maximal expiratory flow and also reduces tethering open of small intrapulmonary airways. The flow-volume loop demonstrates marked scooping, with an initial transient spike of flow attributable largely to expulsion of air from collapsing central airways at the onset of forced exhalation. Otherwise, the central airways remain relatively unaffected, so R_{aw} is normal in “pure” emphysema. Loss of alveolar surface and capillaries in the alveolar walls reduces DL_{CO} ; however, because poorly ventilated emphysematous acini are also poorly perfused (due to loss of their capillaries), arterial hypoxemia usually is not seen at rest until emphysema becomes very severe. However, during exercise, Pa_{O_2} may fall precipitously if extensive destruction of the pulmonary vasculature prevents a sufficient increase in cardiac output and mixed venous oxygen content falls substantially. Under these circumstances, any venous admixture through low \dot{V}/\dot{Q} units has a particularly marked effect in lowering mixed arterial oxygen tension.

FUNCTIONAL MEASUREMENTS

Measurement of Ventilatory Function • LUNG VOLUMES Figure 306e-2 demonstrates a spirometry tracing in which the volume of air entering or exiting the lung is plotted over time. In a slow vital capacity maneuver, the subject inhales from FRC, fully inflating the lungs to TLC, and then exhales slowly to RV; VC, the difference between TLC and RV, represents the maximal excursion of the respiratory system. Spirometry discloses relative volume changes during these maneuvers but cannot reveal the absolute volumes at which they occur. To determine absolute lung volumes, two approaches are commonly used: inert gas dilution and body plethysmography. In the former, a known amount of a nonabsorbable inert gas (usually helium or neon) is inhaled in a single large breath or is rebreathed from a closed circuit; the inert gas is diluted by the gas resident in the lung at the time of inhalation, and its final concentration reveals the volume of resident gas contributing to the dilution. A drawback of this method is that regions of the lung that ventilate poorly (e.g., due to airflow obstruction) may not receive much inspired inert gas and so do not contribute to its dilution. Therefore, inert gas dilution (especially in the single-breath method) often underestimates true lung volumes.

In the second approach, FRC is determined by measuring the compressibility of gas within the chest, which is proportional to the volume of gas being compressed. The patient sits in a body plethysmograph (a chamber usually made of transparent plastic to minimize claustrophobia) and, at the end of a normal tidal breath (i.e., when lung volume is at FRC), is instructed to pant against a closed shutter, thus periodically compressing air within the lung slightly. Pressure fluctuations at the mouth and volume fluctuations within the body box (equal but opposite to those in the chest) are determined, and from these measurements the thoracic gas volume is calculated by means of Boyle’s law. Once FRC is obtained, TLC and RV are calculated by adding the value for inspiratory capacity and subtracting the value for expiratory reserve volume, respectively (both values having been obtained during spirometry) (Fig. 306e-2). The most important determinants of healthy individuals’ lung volumes are height, age, and sex, but there is considerable additional normal variation beyond that accounted for by these parameters. In addition, race influences lung volumes; on average, TLC values are ~12% lower in African Americans and 6% lower in Asian Americans than in Caucasian Americans. In practice, a mean “normal” value is predicted by multivariate regression equations using height, age, and sex, and the patient’s value is divided by the predicted value (often with “race correction” applied) to determine “percent predicted.” For most measures of lung function, 85–115% of the predicted value

can be normal; however, in health, the various lung volumes tend to scale together. For example, if one is “normal big” with a TLC 110% of the predicted value, then all other lung volumes and spirometry values will also approximate 110% of their respective predicted values. This pattern is particularly helpful in evaluating airflow, as discussed below.

AIR FLOW As noted above, spirometry plays a key role in lung volume determination. Even more often, spirometry is used to measure airflow, which reflects the dynamic properties of the lung. During an FVC maneuver, the patient inhales to TLC and then exhales rapidly and forcefully to RV; this method ensures that flow limitation has been achieved, so that the precise effort made has little influence on actual flow. The total amount of air exhaled is the FVC, and the amount of air exhaled in the first second is the FEV_1 ; the FEV_1 is a flow rate, revealing volume change per time. Like lung volumes, an individual’s maximal expiratory flows should be compared with predicted values based on height, age, and sex. While the FEV_1/FVC ratio is typically reduced in airflow obstruction, this condition can also reduce FVC by raising RV, sometimes rendering the FEV_1/FVC ratio “artificially normal” with the erroneous implication that airflow obstruction is absent. To circumvent this problem, it is useful to compare FEV_1 as a fraction of its predicted value with TLC as a fraction of its predicted value. In health, the results are usually similar. In contrast, even an FEV_1 value that is 95% of its predicted value may actually be relatively low if TLC is 110% of its respective predicted value. In this case, airflow obstruction may be present, despite the “normal” value for FEV_1 .

The relationships among volume, flow, and time during spirometry are best displayed in two plots—the spirogram (volume vs. time) and the flow-volume loop (flow vs. volume) (Fig. 306e-4). In conditions that cause airflow obstruction, the site of obstruction is sometimes correlated with the shape of the flow-volume loop. In diseases that cause lower airway obstruction, such as asthma and emphysema, flows decrease more rapidly with declining lung volumes, leading to a characteristic scooping of the flow-volume loop. In contrast, fixed upper-airway obstruction typically leads to inspiratory and/or expiratory flow plateaus (Fig. 306e-4).

AIRWAYS RESISTANCE The total resistance of the pulmonary and upper airways is measured in the same body plethysmograph used to measure FRC. The patient is asked once again to pant, but this time against a closed and then opened shutter. Panting against the closed shutter reveals the thoracic gas volume as described above. When the shutter is opened, flow is directed to and from the body box, so that volume fluctuations in the box reveal the extent of thoracic gas compression, which in turn reveals the pressure fluctuations driving flow. Simultaneous measurement of flow allows the calculation of lung resistance (as flow divided by pressure). In health, R_{aw} is very low (<2 cmH₂O/L per second), and half of the detected resistance resides within the upper airway. In the lung, most resistance originates in the central airways. For this reason, airways resistance measurement tends to be insensitive to peripheral airflow obstruction.

RESPIRATORY MUSCLE STRENGTH To measure respiratory muscle strength, the patient is instructed to exhale or inhale with maximal effort against a closed shutter while pressure is monitored at the mouth. Pressures greater than ± 60 cmH₂O at FRC are considered adequate and make it unlikely that respiratory muscle weakness accounts for any other resting ventilatory dysfunction that is identified.

Measurement of Gas Exchange • DIFFUSING CAPACITY (DL_{CO}) This test uses a small (and safe) amount of carbon monoxide (CO) to measure gas exchange across the alveolar membrane during a 10-sec breath hold. CO in exhaled breath is analyzed to determine the quantity of CO crossing the alveolar membrane and combining with hemoglobin in red blood cells. This “single-breath diffusing capacity” (DL_{CO}) value increases with the surface area available for diffusion and the amount of hemoglobin within the capillaries, and it varies inversely with alveolar membrane thickness. Thus, DL_{CO} decreases in diseases that thicken or destroy alveolar membranes (e.g., pulmonary fibrosis, emphysema), curtail the pulmonary vasculature (e.g., pulmonary hypertension), or reduce alveolar capillary hemoglobin (e.g., anemia). Single-breath

diffusing capacity may be elevated in acute congestive heart failure, asthma, polycythemia, and pulmonary hemorrhage.

ARTERIAL BLOOD GASES The effectiveness of gas exchange can be assessed by measuring the partial pressures of oxygen and CO₂ in a sample of blood obtained by arterial puncture. The oxygen content of blood (Ca_{O₂}) depends upon arterial saturation (%O₂Sat), which is set by Pa_{O₂}, pH, and Pa_{CO₂} according to the oxyhemoglobin dissociation curve. Ca_{O₂} can also be measured by oximetry (see below):

$$\text{Ca}_{\text{O}_2} \text{ (mL/dL)} = 1.39 \text{ (mL/dL)} \times [\text{hemoglobin}] \text{ (g)} \times \% \text{ O}_2 \text{ Sat} \\ + 0.003 \text{ (mL/dL/mmHg)} \times \text{Pa}_{\text{O}_2} \text{ (mmHg)}$$

If hemoglobin saturation alone needs to be determined, this task can be accomplished noninvasively with pulse oxymetry.

ACKNOWLEDGMENT

The authors wish to acknowledge the contributions of Drs. Steven E. Weinberger and Irene M. Rosen to this chapter in previous editions as well as the helpful contributions of Drs. Mary Streck and Jeffrey Jacobson.

307 Diagnostic Procedures in Respiratory Disease

Anne L. Fuhlbrigge, Augustine M. K. Choi

The diagnostic modalities available for assessing the patient with suspected or known respiratory system disease include imaging studies and techniques for acquiring biologic specimens, some of which involve direct visualization of part of the respiratory system. Methods to characterize the functional changes developing as a result of disease, including pulmonary function tests and measurements of gas exchange, are discussed in [Chap. 306e](#).

IMAGING STUDIES

ROUTINE RADIOGRAPHY

Routine chest radiography, including both posteroanterior (PA) and lateral views, is an integral part of the diagnostic evaluation of diseases involving the pulmonary parenchyma, the pleura, and, to a lesser extent, the airways and the mediastinum ([see Chaps. 305 and 308e](#)). Lateral decubitus views are useful for determining whether pleural abnormalities represent freely flowing fluid, whereas apical lordotic views can visualize disease at the lung apices better than the standard PA view. Portable equipment is often used for acutely ill patients who cannot be transported to a radiology suite but are more difficult to interpret owing to several limitations: (1) the single anteroposterior (AP) projection obtained; (2) variability in over- and underexposure of film; (3) a shorter focal spot-film distance leading to lack of edge sharpness and loss of fine detail; and (4) magnification of the cardiac silhouette and other anterior structures by the AP projection. Common radiographic patterns and their clinical correlates are reviewed in [Chap. 308e](#).

Advances in computer technology have allowed the development of digital or computed radiography, which has several benefits: (1) immediate availability of the images; (2) significant postprocessing analysis of images to improve diagnostic information; and (3) ability to store images electronically and to transfer them within or between health care systems.

ULTRASOUND

Diagnostic ultrasound (US) produces images using echoes or reflection of the US beam from interfaces between tissues with differing acoustic properties. US is nonionizing and safe to perform on pregnant patients and children. It can detect and localize pleural abnormalities and is a quick and effective way of guiding percutaneous needle biopsy of peripheral lung, pleural, or chest wall lesions. US is also helpful in identifying septations within loculated collections and can facilitate placement of a needle for sampling of pleural liquid (i.e., for thoracentesis), improving the yield and safety of the procedure. Bedside availability makes it valuable in the intensive care setting. Real-time imaging can be used to assess the movement of the diaphragm. Because US energy is rapidly dissipated in air, it is not useful for evaluation of

1664 the pulmonary parenchyma and cannot be used if there is any aerated lung between the US probe and the abnormality of interest.

Endobronchial US, in which the US probe is passed through a bronchoscope, is a valuable adjunct to bronchoscopy, allowing identification and localization of pathology adjacent to airway walls or within the mediastinum.

NUCLEAR MEDICINE TECHNIQUES

Nuclear imaging depends on the selective uptake of various compounds by organs of the body. In thoracic imaging, these compounds are concentrated by one of three mechanisms: blood pool or compartmentalization (e.g., within the heart), physiologic incorporation (e.g., bone or thyroid) and capillary blockage (e.g., lung scan). Radioactive isotopes can be administered by either the IV or inhaled routes or both. When injected intravenously, albumin macroaggregates labeled with technetium-99m (^{99m}Tc) become lodged in pulmonary capillaries; the distribution of the trapped radioisotope follows the distribution of blood flow. When inhaled, radiolabeled xenon gas can be used to demonstrate the distribution of ventilation. Using these techniques, ventilation-perfusion lung scanning was a commonly used technique for the evaluation of pulmonary embolism. Pulmonary thromboembolism produces one or more regions of ventilation-perfusion mismatch (i.e., regions in which there is a defect in perfusion that follows the distribution of a vessel and that is not accompanied by a corresponding defect in ventilation [Chap. 300]). However, with advances in computed tomography (CT) scanning, scintigraphic imaging has been largely replaced by CT angiography in patients with suspected pulmonary embolism.

Another common use of ventilation-perfusion scans is in patients with impaired lung function, who are being considered for lung resection. Many patients with bronchogenic carcinoma have coexisting chronic obstructive pulmonary disease (COPD), and the question arises as to whether or not a patient can tolerate lung resection. The distribution of the isotope(s) can be used to assess the regional distribution of blood flow and ventilation, allowing the physician to estimate the level of postoperative lung function.

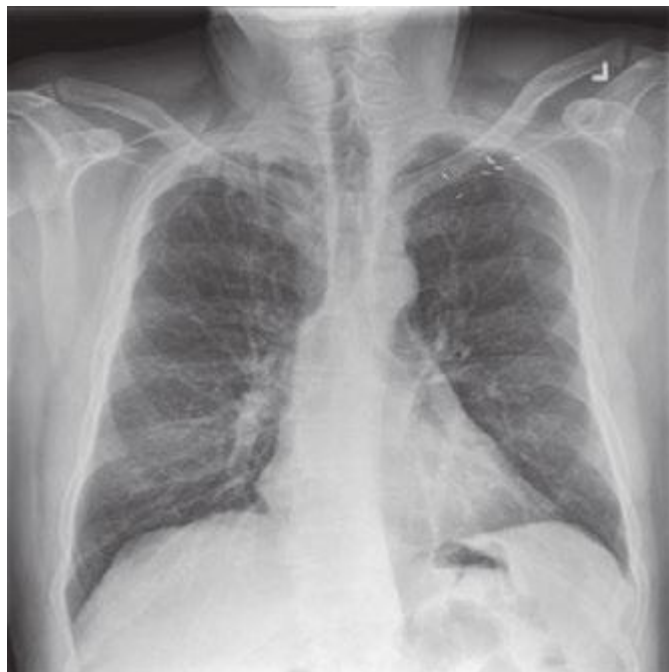
COMPUTED TOMOGRAPHY

CT offers several advantages over routine chest radiography (Figs. 307-1A, B and 307-2A, B; see also Figs. 315-3, 315-4, and 322-4). First, the use of cross-sectional images allows distinction between densities that would be superimposed on plain radiographs. Second, CT is far better than routine radiographic studies at characterizing tissue density and providing accurate size assessment of lesions.

CT is particularly valuable in assessing hilar and mediastinal disease (often poorly characterized by plain radiography), in identifying and characterizing disease adjacent to the chest wall or spine (including pleural disease), and in identifying areas of fat density or calcification in pulmonary nodules (Fig. 307-2). Its utility in the assessment of mediastinal disease has made CT an important tool in the staging of lung cancer (Chap. 107). With the additional use of contrast material, CT also makes it possible to distinguish vascular from nonvascular structures, which is particularly important in distinguishing lymph nodes and masses from vascular structures primarily in the mediastinum, and vascular disorders such as pulmonary embolism.

In high-resolution CT (HRCT), the thickness of individual cross-sectional images is ~1–2 mm, rather than the usual 7–10 mm in conventional CT. The visible detail on HRCT scans allows better recognition of subtle parenchymal and airway disease, thickened interlobular septa, ground-glass opacification, small nodules, and the abnormally thickened or dilated airways seen in bronchiectasis. Using HRCT, characteristic patterns are recognized for many interstitial lung diseases such as lymphangitic carcinoma, idiopathic pulmonary fibrosis, sarcoidosis, and eosinophilic granuloma. However, there is debate about the settings in which the presence of a characteristic pattern on HRCT eliminates the need for obtaining lung tissue to make a diagnosis.

Helical CT and Multidetector CT Helical scanning is currently the standard method for thoracic CT. Helical CT technology results in faster scans with improved contrast enhancement and thinner collimation.



A



B

FIGURE 307-1 Chest x-ray (A) and computed tomography (CT) scan (B) from a patient with emphysema. The extent and distribution of emphysema are not well appreciated on plain film but clearly evident on the CT scan obtained.

Images are obtained during a single breath-holding maneuver that allows less motion artifact and collection of continuous data over a larger volume of lung than is possible with conventional CT. Data from the imaging procedure can be reconstructed in coronal or sagittal planes (Fig. 307-3A), as well as the traditional cross-sectional (axial) view.

Further refinements in detector technology have allowed production of scanners with additional detectors along the scanning axis (z-axis). These *multidetector CT* (MDCT) scanners can obtain multiple slices in a single rotation that are thinner and can be acquired in a shorter period of time. This results in enhanced resolution and



A



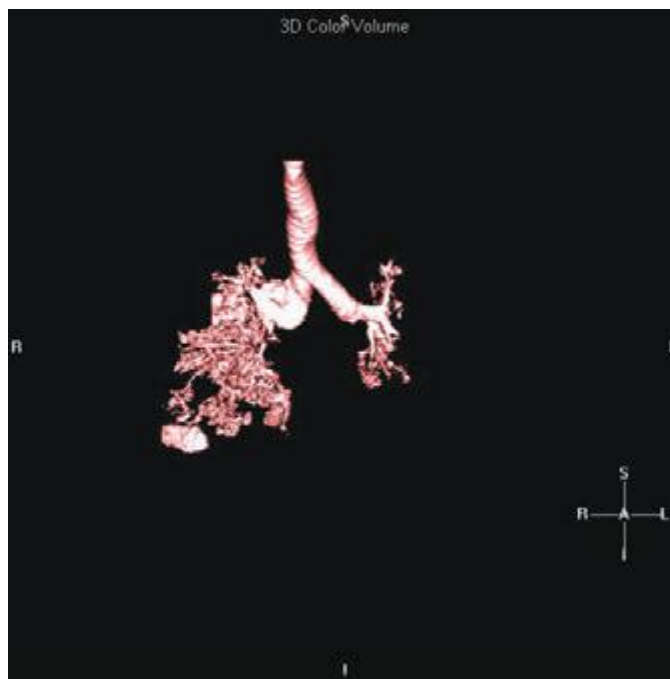
B

FIGURE 307-2 Chest x-ray (A) and computed tomography (CT) scan (B) demonstrating a right lower-lobe mass. The mass is not well appreciated on the plain film because of the hilar structures and known calcified adenopathy. CT is superior to plain radiography for the detection of abnormal mediastinal densities and the distinction of masses from adjacent vascular structures.

increased image reconstruction ability. As the technology has progressed, higher numbers (currently up to 64) of detectors are used to produce clearer final images. MDCT allows for even shorter breath holds, which are beneficial for all patients but especially children, the elderly, and the critically ill. However, it should be noted that despite the advantages of MDCT, there is an increase in radiation dose compared to single-detector CT to consider.



A



B

FIGURE 307-3 Spiral computed tomography (CT) with reconstruction of images in planes other than axial view. Spiral CT in a lung transplant patient with a dehiscence and subsequent aneurysm of the anastomosis. CT images were reconstructed in the sagittal view (A) and using digital subtraction to view images of the airways only (B), which demonstrate the exact location and extent of the abnormality.

In MDCT, the additional detectors along the z-axis result in improved use of the contrast bolus. This and the faster scanning times and increased resolution have all led to improved imaging of the pulmonary vasculature and the ability to detect segmental and subsegmental emboli. CT pulmonary angiography (CTPA) also allows simultaneous detection of parenchymal abnormalities that may be contributing to a patient's clinical presentation. Secondary to these

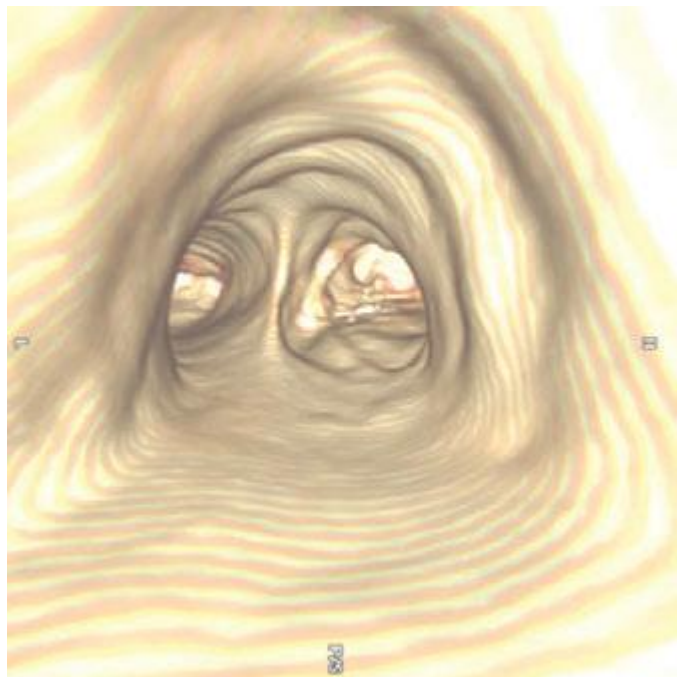


FIGURE 307-4 Virtual bronchoscopic image of the trachea. The view projected is one that would be obtained from the trachea looking down to the carina. The left and right main stem airways are seen bifurcating from the carina.

advantages and increasing availability, CTPA has rapidly become the test of choice for many clinicians in the evaluation of pulmonary embolism; compared with pulmonary angiography, it is considered equal in terms of accuracy and with less associated risks.

VIRTUAL BRONCHOSCOPY

The three-dimensional (3D) image of the thorax obtained by MDCT can be digitally stored, reanalyzed, and displayed as 3D reconstructions of the airways down to the sixth to seventh generation. Using these reconstructions, a “virtual” bronchoscopy can be performed (Fig. 307-4). Virtual bronchoscopy has been proposed as an adjunct to conventional bronchoscopy in several clinical situations: It can allow accurate assessment of the extent and length of an airway stenosis, including the airway distal to the narrowing; it can provide useful information about the relationship of the airway abnormality to adjacent mediastinal structures; and it allows preprocedure planning for therapeutic bronchoscopy to help ensure the appropriate equipment is available for the procedure.

Virtual bronchoscopy can be used to help target the area of peripheral lung for endobronchial lung volume reduction surgery that is being used in the management of pulmonary emphysema. The extent of emphysema in each segmental region together with other anatomic details may help in choosing the most appropriate subsegments. However, software packages for the generation of virtual bronchoscopic images are relatively early in development, and their utilization and potential impact on patient care are still unknown. Electromagnetic navigational bronchoscopy systems (EMN or ENB) using virtual bronchoscopy have been developed to allow accurate navigation to peripheral pulmonary target lesions, using technology similar to a car global positioning system (GPS) unit.

POSITRON EMISSION TOMOGRAPHIC SCANNING

Positron emission tomographic (PET) scanning is commonly used to identify malignant lesions in the lung, based on their increased uptake and metabolism of glucose. The technique involves injection of a radiolabeled glucose analogue, [¹⁸F]-fluoro-2-deoxyglucose (FDG), which is taken up by metabolically active malignant cells. However, FDG is trapped within the cells following phosphorylation, and the

unstable [¹⁸F] decays by emission of positrons, which can be detected by a specialized PET camera or by a gamma camera that has been adapted for imaging of positron-emitting nuclides. This technique has been used in the evaluation of solitary pulmonary nodules and in staging lung cancer. Detection or exclusion of mediastinal lymph node involvement and identification of extrathoracic disease can be achieved. The limited anatomical definition of radionuclide imaging has been improved by the development of hybrid imaging that allows the superimposition of PET and CT images, a technique known as functional-anatomical mapping. Hybrid PET/CT scans provide images that help pinpoint the abnormal metabolic activity to anatomical structures seen on CT and provide more accurate diagnoses than the two scans performed separately. FDG-PET can differentiate benign from malignant lesions as small as 1 cm. However, false-negative findings can occur in lesions with low metabolic activity such as carcinoid tumors and bronchioloalveolar cell carcinomas, or in lesions <1 cm in which the required threshold of metabolically active malignant cells is not present for PET diagnosis. False-positive results can be seen due to FDG uptake in inflammatory conditions such as pneumonia and granulomatous diseases.

MAGNETIC RESONANCE IMAGING

The role of magnetic resonance imaging (MRI) in the evaluation of respiratory system disease is less well-defined than that of CT. Magnetic resonance (MR) provides poorer spatial resolution and less detail of the pulmonary parenchyma and, for these reasons, is currently not considered a substitute for CT in imaging the thorax. However, the use of hyperpolarized gas in conjunction with MR has led to the investigational use of MR for imaging the lungs, particularly in obstructive lung disease. In addition, imaging performed during an inhalation and exhalation can provide dynamic information on lung function. Of note, MR examinations are difficult to obtain among several subgroups of patients. Patients who cannot lie still or who cannot lie on their backs may have MRIs that are of poor quality; some tests require patients to hold their breaths for 15–25 seconds at a time in order to get good MRIs. MRI is generally avoided in unstable and/or ventilated patients and those with severe trauma because of the hazards of the MR environment and the difficulties in monitoring patients within the MR room. The presence of metallic foreign bodies, pacemakers, and intracranial aneurysm clips also preclude use of MRI.

An advantage of MR is the use of nonionizing electromagnetic radiation. Additionally, MR is well suited to distinguish vascular from nonvascular structures without the need for contrast. Blood vessels appear as hollow tubular structures because flowing blood does not produce a signal on MRI. Therefore, MR can be useful in demonstrating pulmonary emboli, defining aortic lesions such as aneurysms or dissection, or other vascular abnormalities (Fig. 307-5) if radiation and IV contrast medium cannot be used. Gadolinium can be used as an intravascular contrast agent for MR angiography (MRA); however, synchronization of data acquisition with the peak arterial bolus is one of the major challenges of MRA. The flow of contrast medium from the peripheral injection site to the vessel of interest is affected by a number of factors including heart rate, stroke volume, and the presence of proximal stenotic lesions.

PULMONARY ANGIOGRAPHY

The pulmonary arterial system can be visualized by pulmonary angiography, in which radiopaque contrast medium is injected through a catheter placed in the pulmonary artery. When performed in cases of pulmonary embolism, pulmonary angiography demonstrates the consequences of an intravascular thrombus—either a defect in the lumen of a vessel (a filling defect) or an abrupt termination (cutoff) of the vessel. Other, less common indications for pulmonary angiography include visualization of a suspected pulmonary arteriovenous malformation and assessment of pulmonary arterial invasion by a neoplasm. The risks associated with modern arteriography are small, generally of greatest concern in patients with severe pulmonary hypertension or chronic kidney disease. With advances in CT scanning, MDCT



FIGURE 307-5 Magnetic resonance angiography image of the vasculature of a patient after lung transplant. The image demonstrates the detailed view of the vasculature that can be obtained using digital subtraction techniques. Images from a patient after lung transplant show the venous and arterial anastomosis on the right; a slight narrowing is seen at the site of the anastomosis, which is considered within normal limits and not suggestive of obstruction.

angiography (MDCTA) is replacing conventional angiography for the diagnosis of pulmonary embolism.

MEDICAL TECHNIQUES FOR OBTAINING BIOLOGIC SPECIMENS

COLLECTION OF SPUTUM

Sputum can be collected either by spontaneous expectoration or induced (after inhalation of an irritating aerosol such as hypertonic saline). *Sputum induction* is used either because sputum is not spontaneously being produced or because of an expected higher yield of certain types of findings. Because sputum consists mainly of secretions from the tracheobronchial tree rather than the upper airway, the finding of alveolar macrophages and other inflammatory cells is consistent with a lower respiratory tract origin of the sample, whereas the presence of squamous epithelial cells in a “sputum” sample indicates contamination by secretions from the upper airways.

In addition to processing for routine bacterial pathogens by Gram’s method and culture, sputum can be processed for a variety of other pathogens, including staining and culture for mycobacteria or fungi, culture for viruses, and staining for *Pneumocystis jiroveci*. In the specific case of sputum obtained for evaluation of *P. jiroveci* pneumonia, for example, sputum should be collected by induction rather than spontaneous expectoration, and an immunofluorescent stain should be used to detect the organisms. Traditional stains and cultures are now also being supplemented in some cases by immunologic techniques and by molecular biologic methods, including the use of polymerase chain reaction amplification and DNA probes. Cytologic staining of sputum for malignant cells, using the traditional Papanicolaou method, allows noninvasive evaluation for suspected lung cancer.

PERCUTANEOUS NEEDLE ASPIRATION (TRANSTHORACIC)

A needle can be inserted through the chest wall into a pulmonary lesion to obtain an aspirate or tissue core for cytologic/histologic or microbiologic analysis. Aspiration can be performed to obtain a diagnosis or to decompress and/or drain a fluid collection. The procedure is usually carried out under CT or ultrasound guidance to assist positioning of

the needle and assure localization in the lesion. The low potential risk of this procedure (intrapulmonary bleeding or creation of a pneumothorax with collapse of the underlying lung) in experienced hands is usually acceptable compared with the information obtained. However, a limitation of the technique is sampling error due to the small size of the tissue sample. Thus, findings other than a specific cytologic or microbiologic diagnosis are of limited clinical value.

THORACENTESIS

Sampling of pleural liquid by thoracentesis is commonly performed for diagnostic purposes or, in the case of a large effusion, for palliation of dyspnea. Diagnostic sampling, either by blind needle aspiration or after localization by US, allows the collection of liquid for microbiologic and cytologic studies. Analysis of the fluid obtained for its cellular composition and chemical constituents allows classification of the effusion and can help with diagnosis and treatment (Chap. 316).

BRONCHOSCOPY

Bronchoscopy is the process of direct visualization of the tracheobronchial tree. Although bronchoscopy is now performed almost exclusively with flexible fiberoptic instruments, rigid bronchoscopy, generally performed in an operating room on a patient under general anesthesia, still has a role in selected circumstances, primarily because of a larger suction channel and the fact that the patient can be ventilated through the bronchoscope channel. These situations include the retrieval of a foreign body and the suctioning of a massive hemorrhage, for which the small suction channel of the bronchoscope may be insufficient.

FLEXIBLE FIBEROPTIC BRONCHOSCOPY

This outpatient procedure is usually performed in an awake but sedated patient (conscious sedation). The bronchoscope is passed through either the mouth or the nose, between the vocal cords, and into the trachea. The ability to flex the scope makes it possible to visualize virtually all airways to the level of subsegmental bronchi. The bronchoscopist is able to identify endobronchial pathology, including tumors, granulomas, bronchitis, foreign bodies, and sites of bleeding. Samples from airway lesions can be taken by several methods, including washing, brushing, and biopsy. Washing involves instillation of sterile saline through a channel of the bronchoscope and onto the surface of a lesion. A portion of the liquid is collected by suctioning through the bronchoscope, and the recovered material can be analyzed for cells (cytology) or organisms (by standard stains and cultures). Brushing or biopsy of the surface of the lesion, using a small brush or biopsy forceps at the end of a long cable inserted through a channel of the bronchoscope, allows recovery of cellular material or tissue for analysis by standard cytologic and histopathologic methods.

The bronchoscope can be used to sample material not only from the regions that can be directly visualized (i.e., the airways) but also from the more distal pulmonary parenchyma. With the bronchoscope wedged into a subsegmental airway, aliquots of sterile saline can be instilled through the scope, allowing sampling of cells and organisms from alveolar spaces. This procedure, called *bronchoalveolar lavage*, has been particularly useful for the recovery of organisms such as *P. jiroveci*.

Brushing and biopsy of the distal lung parenchyma can also be performed with the same instruments that are used for endobronchial sampling. These instruments can be passed through the scope into small airways. When biopsies are performed, the forceps penetrate the airway wall, allowing biopsy of peribronchial alveolar tissue. This procedure, called *transbronchial biopsy*, is used when there is either relatively diffuse disease or a localized lesion of adequate size. With the aid of fluoroscopic imaging, the bronchoscopist is able to determine not only whether and when the instrument is in the area of abnormality, but also the proximity of the instrument to the pleural surface. If the forceps are too close to the pleural surface, there is a risk of violating the visceral pleura and creating a pneumothorax; the other potential complication of transbronchial biopsy is pulmonary hemorrhage. The incidence of these complications is less than several percent.

Another procedure involves use of a hollow-bore needle passed through the bronchoscope for sampling of tissue adjacent to the trachea or a large bronchus. The needle is passed through the airway wall (transbronchial), and cellular material can be aspirated from mass lesions or enlarged lymph nodes, generally in a search for malignant cells. Mediastinoscopy has been considered the gold standard for mediastinal staging; however, transbronchial needle aspiration (TBNA) allows sampling from the lungs and surrounding lymph nodes without the need for surgery or general anesthesia.

ENDOBROCHIAL ULTRASOUND (EBUS)–TRANSBRONCHIAL NEEDLE ASPIRATION (TBNA)

Further advances in needle aspiration techniques have been accomplished with the development of endobronchial ultrasound (EBUS). The technology uses an ultrasonic bronchoscope fitted with a probe that allows for needle aspiration of mediastinal and hilar lymph nodes guided by real-time US images. EBUS allows sampling of mediastinal lymph nodes and masses under direct vision to better identify and localize peribronchial and mediastinal pathology and offers access to more difficult-to-reach areas and smaller lymph nodes in the staging of malignancies. EBUS-TBNA has the potential to access the same paratracheal and subcarinal lymph node stations as mediastinoscopy, but also extends out to the hilar lymph nodes (levels 10 and 11). The usefulness of EBUS for clinical indications other than lung cancer is improving and has been recommended in the evaluation of mediastinal masses of unknown origin early in the diagnostic process.

EMERGING BRONCHOSCOPIC TECHNIQUES

Emerging techniques that can be performed using bronchoscopy include video/autofluorescence bronchoscopy (AFB), narrow band imaging (NBI), optical coherence tomography (OCT), and endomicroscopy using confocal fluorescent laser microscopy (CFM). AFB uses bronchoscopy with an additional light source to screen high-risk individuals and identify premalignant lesions (airway dysplasia) and carcinoma in situ. NBI capitalizes on the increased absorption of blue and green wavelengths of light by hemoglobin to enhance the visibility of vessels of the mucosa and differentiate between inflammatory versus malignant mucosal lesions. CFM uses a blue laser to induce fluorescence, and its high degree of resolution provides a real-time view of living tissue at an almost histologic resolution. OCT uses near-infrared light source and has spatial resolution advantages over CT and MRI. It can penetrate the airway wall up to three times deeper than CFM and is less susceptible to motion artifacts from cardiac pulsation and respiratory movements. However, careful assessment is required before these methods find a place in the evaluation strategy of early lung cancer and other lung diseases.

THERAPEUTIC BRONCHOSCOPY

The bronchoscope may provide the opportunity for treatment as well as diagnosis. A central role of the interventional pulmonology (IP) physician is the performance of therapeutic bronchoscopy. For example, an aspirated foreign body may be retrieved with an instrument passed through the bronchoscope (either flexible or rigid), and bleeding may be controlled with a balloon catheter similarly introduced. Newer interventional techniques performed through a bronchoscope include methods for achieving and maintaining patency of airways that are partially or completely occluded, especially by tumors. These techniques include laser therapy, cryotherapy, argon plasma coagulation, electrocautery, balloon bronchoplasty and dilation, and stent placement. Many IP physicians are also trained in performing percutaneous tracheotomy.

MEDICAL THORACOSCOPY

Medical thoracoscopy (or pleuroscopy) focuses on the diagnosis of pleural-based problems. The procedure is performed with a conventional rigid or a semi-rigid pleuroscope (similar in design to a bronchoscope and enabling the operator to inspect the pleural surface,

sample and/or drain pleural fluid, or perform targeted biopsies of the parietal pleura). Medical thoracoscopy can be performed in the endoscopy suite or operating room with the patient under conscious sedation and local anesthesia. In contrast, video-assisted thoracoscopic surgery (VATS) requires general anesthesia and is only performed in the operating room. A common diagnostic indication for medical thoracoscopy is the evaluation of a pleural effusion or biopsy of presumed parietal pleural carcinomatosis. It can also be used to place a chest tube under visual guidance, or perform chemical or talc pleurodesis, a therapeutic intervention to prevent a recurrent pleural effusion (usually malignant) or recurrent pneumothorax.

The increasing availability of advanced bronchoscopic and pleuroscopic techniques has motivated the development of IP programs. IP can be defined as “the art and science of medicine as related to the performance of diagnostic and invasive therapeutic procedures, that which require additional training and expertise beyond that which required in a standard pulmonary medicine training program.” IP physicians provide alternatives to surgery for patients with a wide variety of thoracic disorders and problems.

SURGICAL TECHNIQUES FOR OBTAINING BIOLOGIC SPECIMENS

Evaluation and diagnosis of disorders of the chest commonly involve collaboration between pulmonologists and thoracic surgeons. Although procedures such as mediastinoscopy, VATS, and thoracotomy are performed by thoracic surgeons, there is overlap in many minimally invasive techniques that can be performed by a pulmonologist, an interventional pulmonologist, or a thoracic surgeon.

MEDIASTINOSCOPY AND MEDIASTINOTOMY

Proper staging of lung cancer is of paramount concern when determining a treatment regimen. Although CT and PET scanning are useful for determining the size and nature of mediastinal lymph nodes as part of the staging of lung cancer, tissue biopsy and histopathologic examination are often critical for the diagnosis of mediastinal masses or enlarged mediastinal lymph nodes. The two major surgical procedures used to obtain specimens from masses or nodes in the mediastinum are mediastinoscopy (via a suprasternal approach) and mediastinotomy (via a parasternal approach). Both procedures are performed under general anesthesia by a qualified surgeon. In the case of suprasternal mediastinoscopy, a rigid mediastinoscope is inserted at the suprasternal notch and passed into the mediastinum along a pathway just anterior to the trachea. Tissue can be obtained with biopsy forceps passed through the scope, sampling masses or nodes that are in a paratracheal or pretracheal position (levels 2R, 2L, 3, 4R, 4L). Aortopulmonary lymph nodes (levels 5, 6) are not accessible by this route and thus are commonly sampled by parasternal mediastinotomy (the Chamberlain procedure). This approach involves a parasternal incision and dissection directly down to a mass or node that requires biopsy.

As an alternative to surgery, a bronchoscope can be used to perform TBNA to obtain tissue from the mediastinum, and, when combined with EBUS, can allow access to the same lymph node stations associated with mediastinoscopy, but also extend access out to the hilar lymph nodes (levels 10, 11). Finally, endoscopic ultrasound (EUS)–fine-needle aspiration (FNA) is a second procedure that complements EBUS-FNA in the staging of lung cancer. EUS-FNA is performed via the esophagus and is ideally suited for sampling lymph nodes in the posterior mediastinum (levels 7, 8, 9). Because US imaging cannot penetrate air-filled spaces, the area directly anterior to the trachea cannot accurately be assessed and is a “blind spot” for EUS-FNA. However, EBUS-FNA can visualize the anterior lymph nodes and can complement EUS-FNA. The combination of EUS-FNA and EBUS-FNA is a technique that is becoming an alternative to surgery for staging the mediastinum in thoracic malignancies.

VIDEO-ASSISTED THORACOSCOPIC SURGERY

VATS has become a standard technique for the diagnosis and management of pleural as well as parenchymal lung disease. This procedure is performed in the operating room using single-lung ventilation

with double-lumen endotracheal intubation and involves the passage of a rigid scope with a distal lens through a trocar inserted into the pleura. A high-quality image is shown on a monitor screen, allowing the operator to manipulate instruments passed into the pleural space through separate small intercostal incisions. With these instruments the operator can biopsy lesions of the pleura under direct visualization. In addition, this procedure is now used commonly to biopsy peripheral lung tissue or to remove peripheral nodules for both diagnostic and therapeutic purposes. This much less invasive procedure has largely supplanted the traditional “open lung biopsy” performed via thoracotomy. The decision to use a VATS technique versus performing an open thoracotomy is made by the thoracic surgeon and is based on whether a patient can tolerate the single-lung ventilation that is required to allow adequate visualization of the lung. With further advances in instrumentation and experience, VATS can be used to perform procedures previously requiring thoracotomy, including stapled lung biopsy, resection of pulmonary nodules, lobectomy, pneumonectomy, pericardial window, or other standard thoracic surgical procedures, but allows them to be performed in a minimally invasive manner.

THORACOTOMY

Although frequently replaced by VATS, thoracotomy remains an option for the diagnostic sampling of lung tissue. It provides the largest

amount of material, and it can be used to biopsy and/or excise lesions that are too deep or too close to vital structures for removal by VATS. The choice between VATS and thoracotomy needs to be made on a case-by-case basis.

308e Atlas of Chest Imaging

Patricia A. Kritek, John J. Reilly, Jr.

This atlas of chest imaging is a collection of interesting chest radiographs and computed tomograms (CTs) of the chest. The readings of the films are meant to be illustrative of specific, major findings. The associated text is not intended as a comprehensive assessment of the images.

EXAMPLES OF NORMAL IMAGING

R

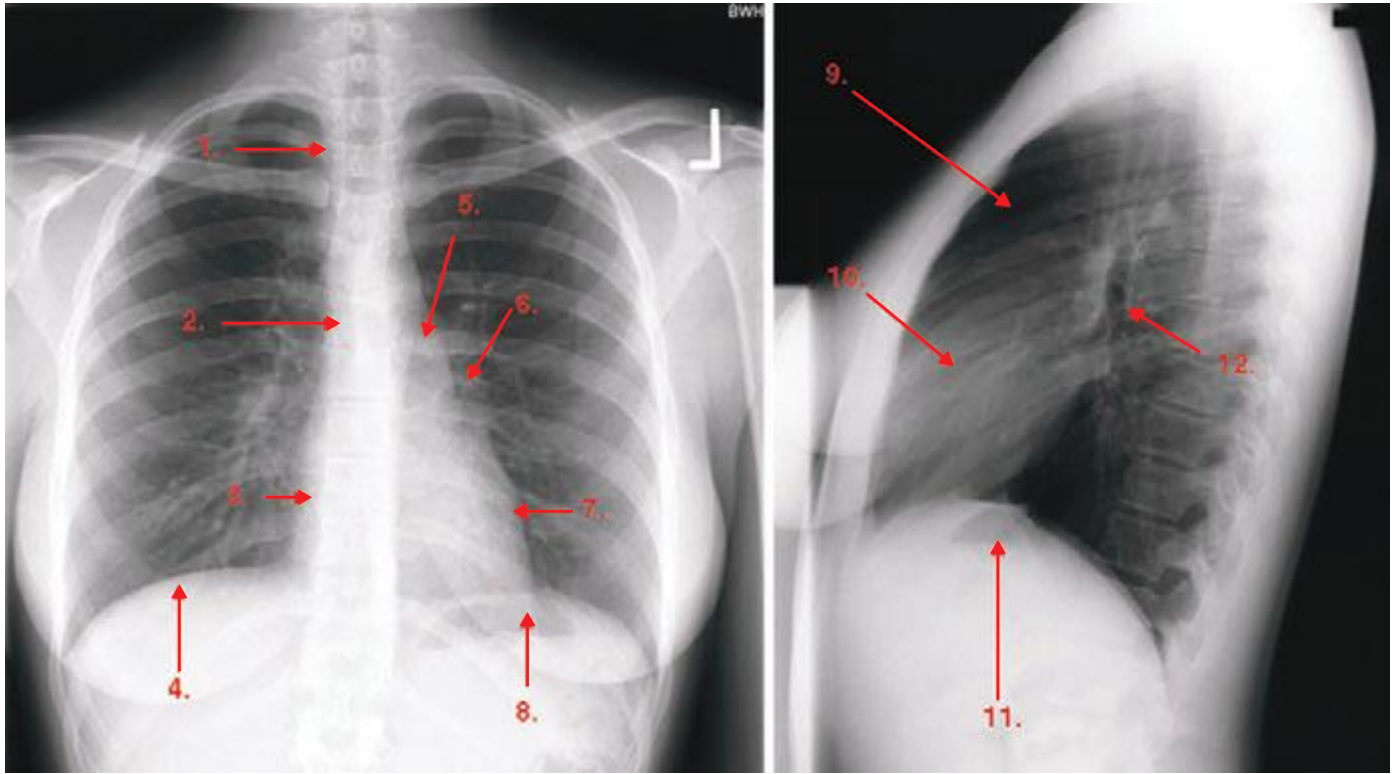


FIGURE 308e-1 Normal chest radiograph—review of anatomy. 1. Trachea. 2. Carina. 3. Right atrium. 4. Right hemidiaphragm. 5. Aortic knob. 6. Left hilum. 7. Left ventricle. 8. Left hemidiaphragm (with stomach bubble). 9. Retrosternal clear space. 10. Right ventricle. 11. Left hemidiaphragm (with stomach bubble). 12. Left upper lobe bronchus.

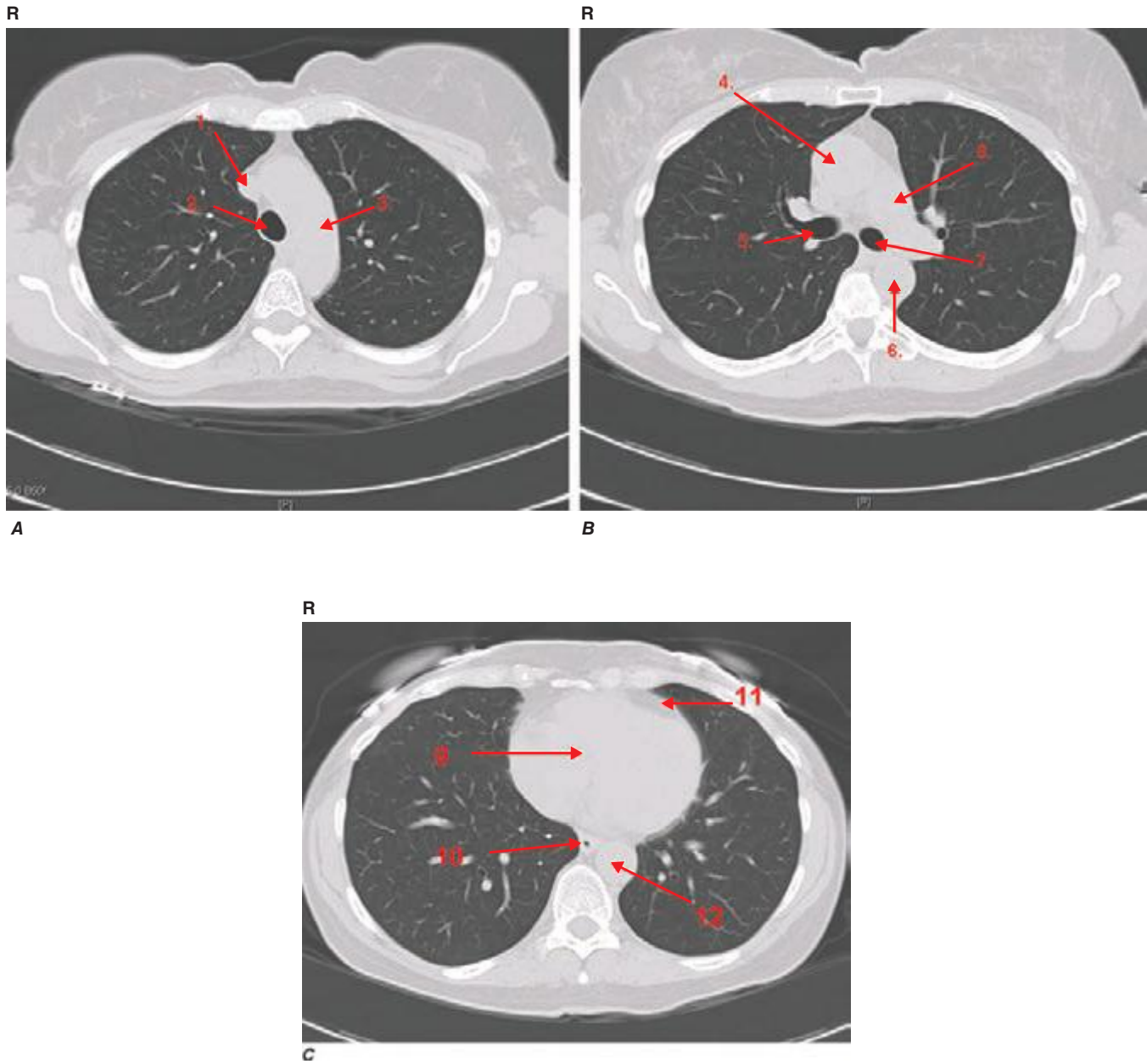


FIGURE 308e-2 Normal chest tomogram—note anatomy. 1. Superior vena cava. 2. Trachea. 3. Aortic arch. 4. Ascending aorta. 5. Right mainstem bronchus. 6. Descending aorta. 7. Left mainstem bronchus. 8. Main pulmonary artery. 9. Heart. 10. Esophagus. 11. Pericardium. 12. Descending aorta.

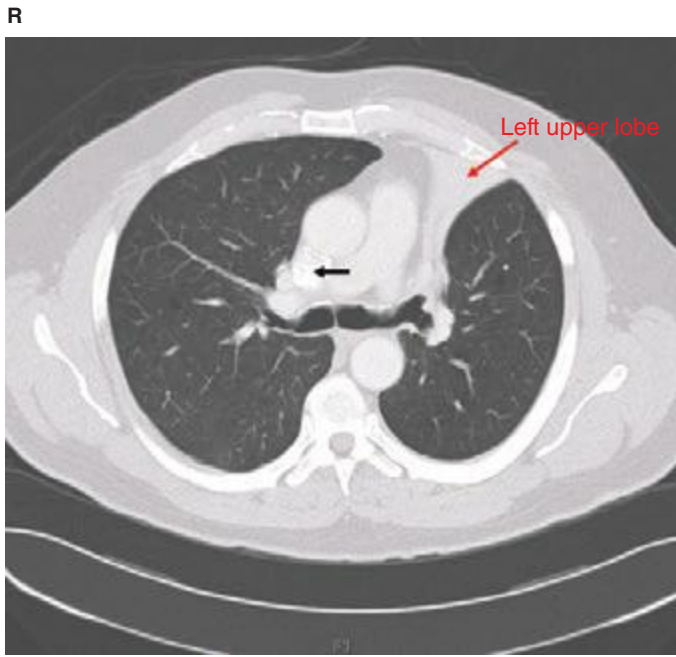


FIGURE 308e-3 CT scan demonstrating left upper lobe collapse. The patient was found to have an endobronchial lesion (not visible on the CT scan) resulting in this finding. The superior vena cava (*black arrow*) is partially opacified by intravenous contrast.

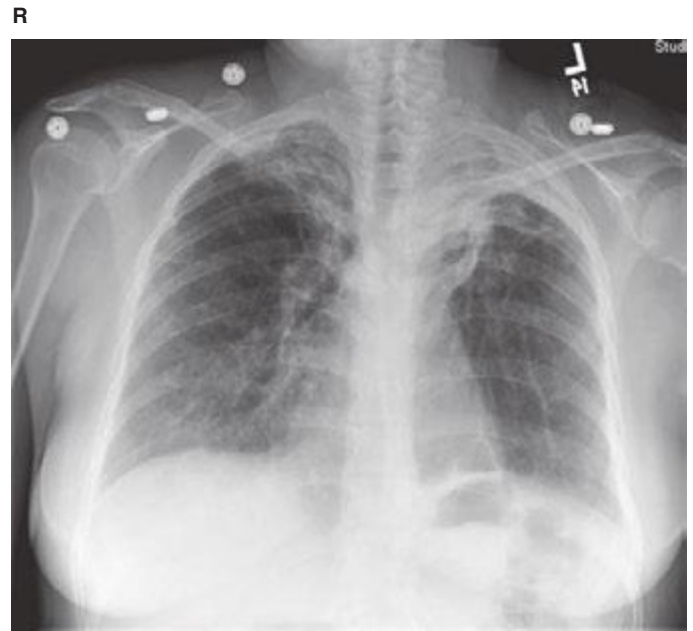


FIGURE 308e-5 Left upper lobe scarring with hilar retraction with less prominent scarring in right upper lobe as well. Findings consistent with previous tuberculosis infection in an immigrant from Ecuador.

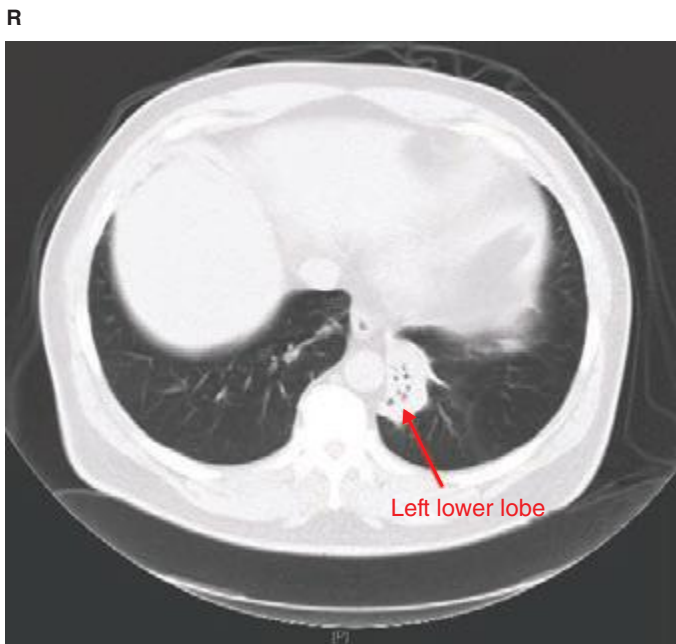


FIGURE 308e-4 CT scan revealing chronic left lower lobe collapse. Note dramatic volume loss with minimal aeration. There is subtle mediastinal shift to the left.



FIGURE 308e-6 Apical scarring, traction bronchiectasis (*red arrow*), and decreased lung volume consistent with previous tuberculosis infection. Findings most significant in left lung.

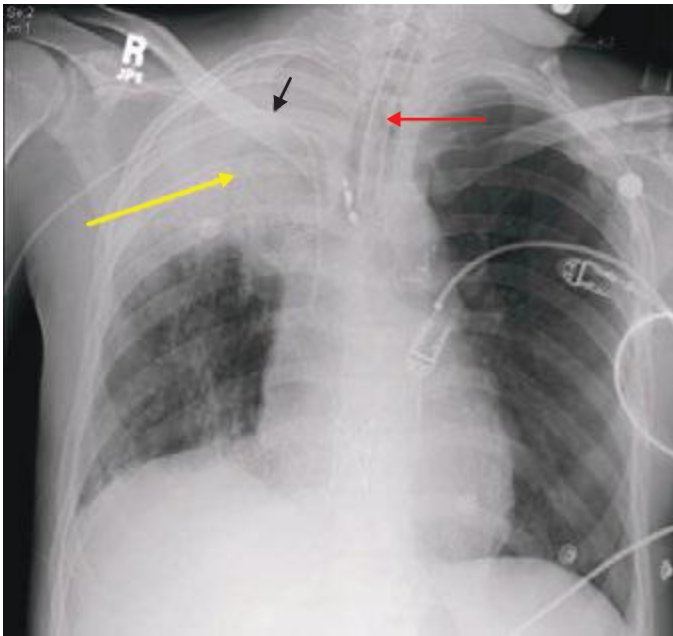


FIGURE 308e-7 Chest radiograph demonstrating right upper lobe collapse (yellow arrow). Note the volume loss as demonstrated by the elevated right hemidiaphragm as well as mediastinal shift to the right. Also apparent on the film are an endotracheal tube (red arrow) and a central venous catheter (black arrow).

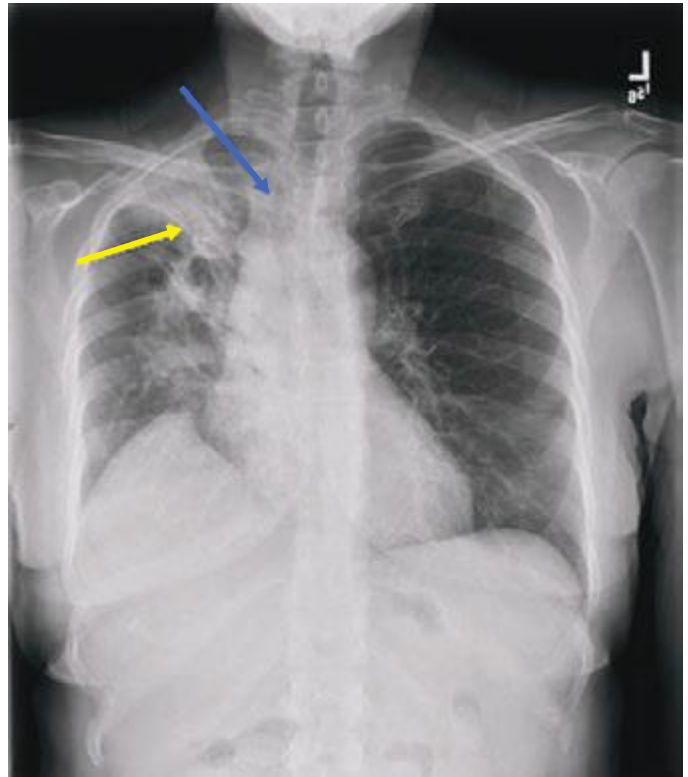


FIGURE 308e-8 Opacity in the right upper lobe. Note the volume loss as indicated by the elevation of the right hemidiaphragm, elevation of minor fissure (yellow arrow), and deviation of the trachea to the right (blue arrow).

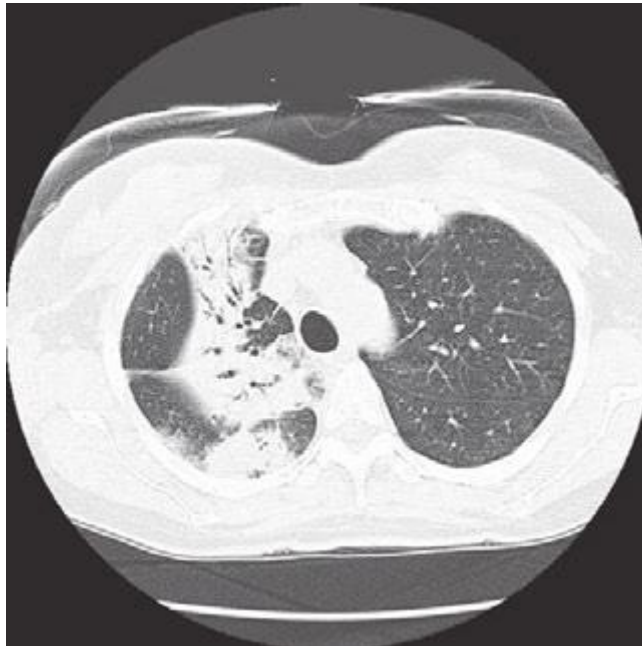


FIGURE 308e-9 CT scan of the same right upper lobe opacity. Note the air bronchograms and areas of consolidation.

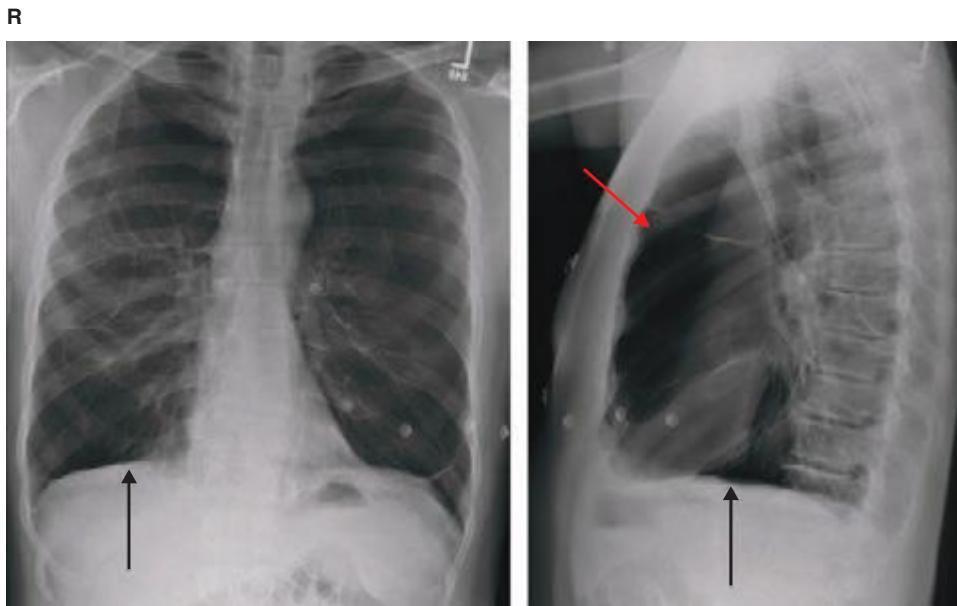


FIGURE 308e-10 Emphysema with increased lucency, flattened diaphragms (*black arrows*), increased anteroposterior (AP) diameter, and increased retrosternal clear space (*red arrow*).



FIGURE 308e-11 CT scan of diffuse, bilateral emphysema.



FIGURE 308e-12 CT scan of bullous emphysema.



FIGURE 308e-13 Lymphangioleiomyomatosis—note multiple thin-walled parenchymal cysts.

R

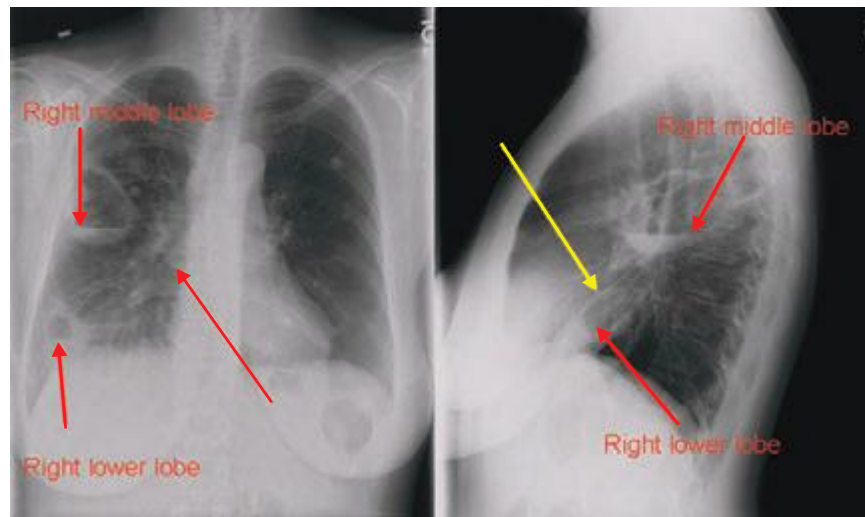


FIGURE 308e-14 Two cavities on posteroanterior (PA) and lateral chest radiograph. Cavities and air-fluid levels identified by red arrows. The smaller cavity is in the right lower lobe (located below the major fissure, identified with the yellow arrow), and the larger cavity is located in the right middle lobe, which is located between the minor (red arrow) and major fissures. There is an associated opacity surrounding the cavity in the right lower lobe.

R



FIGURE 308e-15 CT scan of parenchymal cavity.

R

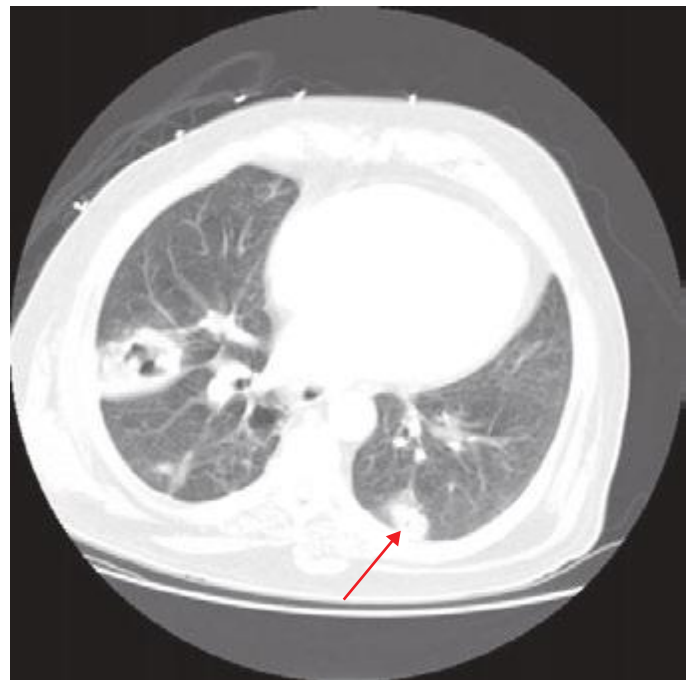


FIGURE 308e-16 Thick-walled cavitary lung lesions. The mass in the right lung has thick walls and advanced cavitation, whereas the smaller nodule on the left has early cavitary changes (arrow). This patient was diagnosed with *Nocardia* infection.

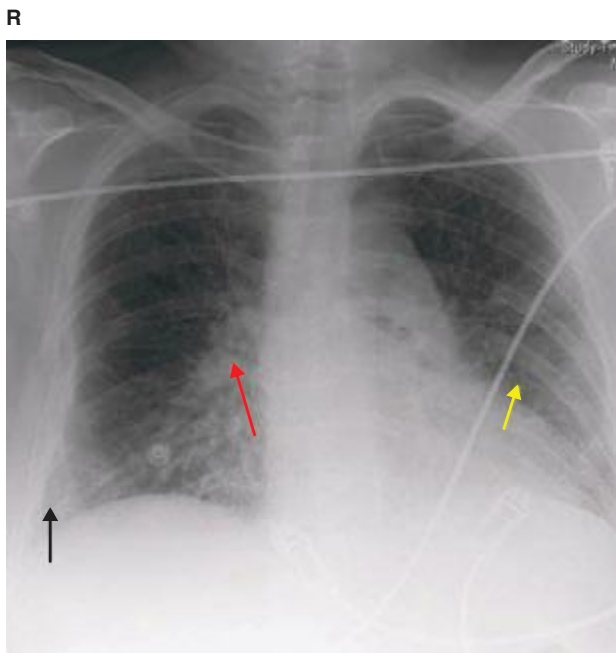


FIGURE 308e-17 Mild congestive heart failure. Note the Kerley B lines (*black arrow*) and perivascular cuffing (*yellow arrow*) as well as the pulmonary vascular congestion (*red arrow*).



FIGURE 308e-18 Pulmonary edema. Note indistinct vasculature, perihilar opacities, and peripheral interstitial reticular opacities. Although this is an anteroposterior film making cardiac size more difficult to assess, the cardiac silhouette still appears enlarged.



FIGURE 308e-19 Chest radiograph demonstrates reticular nodular opacities bilaterally with small lung volumes consistent with usual interstitial pneumonitis (UIP) on pathology. Clinically, UIP is used interchangeably with idiopathic pulmonary fibrosis (IPF).

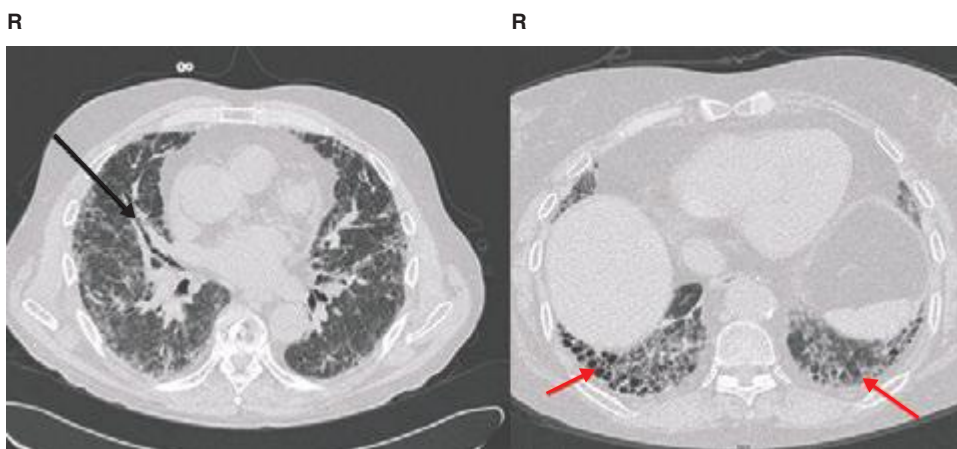
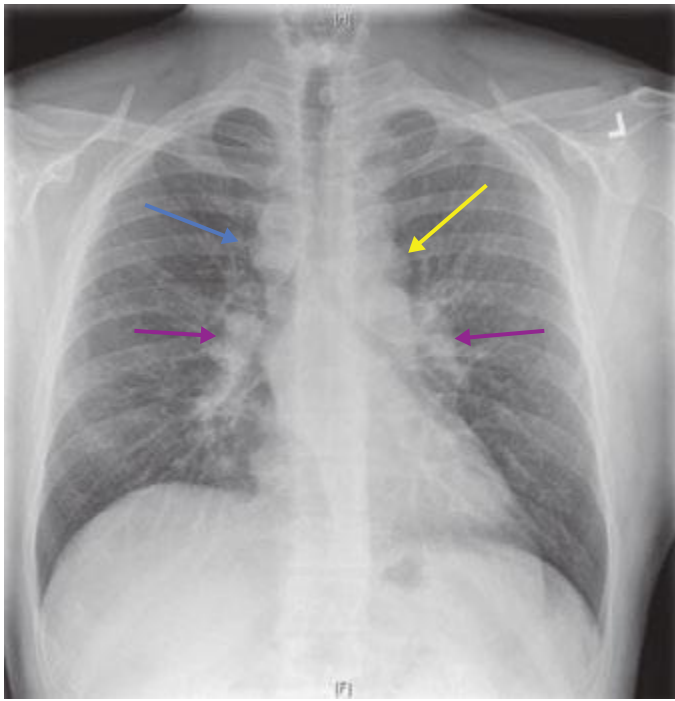
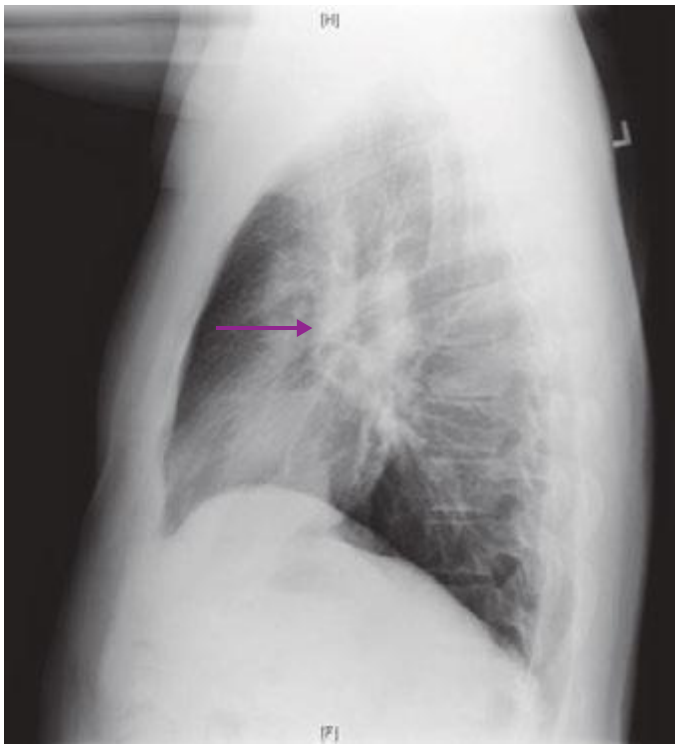


FIGURE 308e-20 CT scan of usual interstitial pneumonitis (UIP), also known as idiopathic pulmonary fibrosis (IPF). Classic findings include traction bronchiectasis (*black arrow*) and honeycombing (*red arrows*). Note subpleural, basilar predominance of the honeycombing.



A



B

FIGURE 308e-21 A. PA chest film—note presence of paratracheal (blue arrow), aortopulmonary window (yellow arrow), and hilar lymphadenopathy (purple arrows). B. Lateral film—note hilar lymphadenopathy (purple arrow).

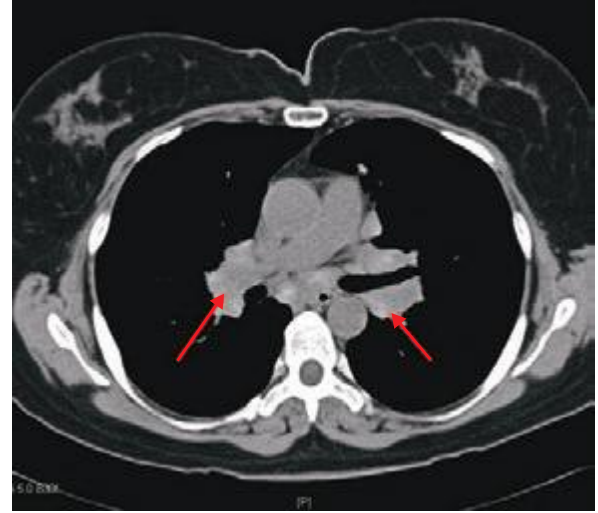
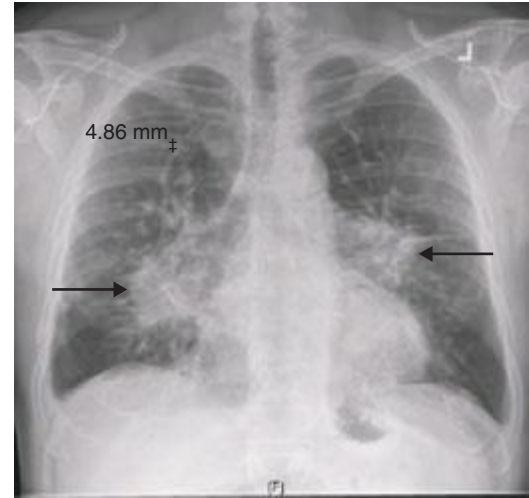
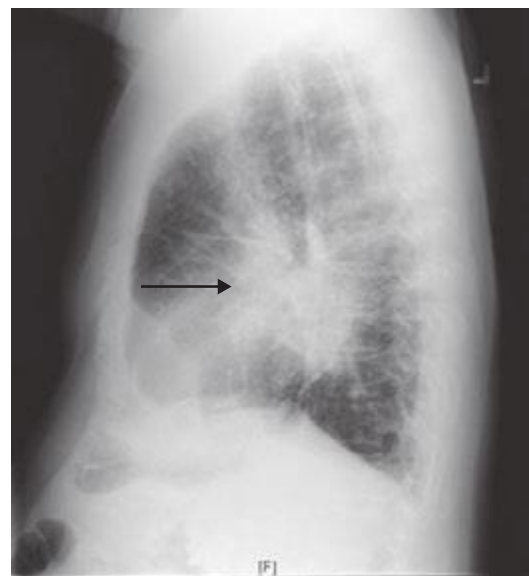


FIGURE 308e-22 Sarcoid—CT scan of stage I demonstrating bulky hilar and mediastinal lymphadenopathy (red arrows).

R



A



B

FIGURE 308e-23 Sarcoid—chest radiograph of stage II. A. PA film with hilar lymphadenopathy (black arrows) and parenchymal changes. B. Lateral film with hilar adenopathy (black arrow) and parenchymal changes.



FIGURE 308e-24 Sarcoid—CT scan of stage II (calcified lymphadenopathy, parenchymal infiltrates).

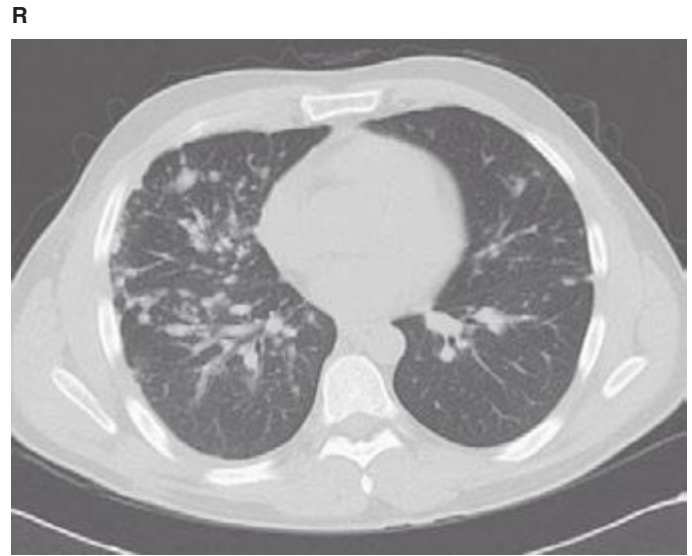


FIGURE 308e-25 Sarcoid—CT scan of stage II (nodular opacities tracking along bronchovascular bundles).

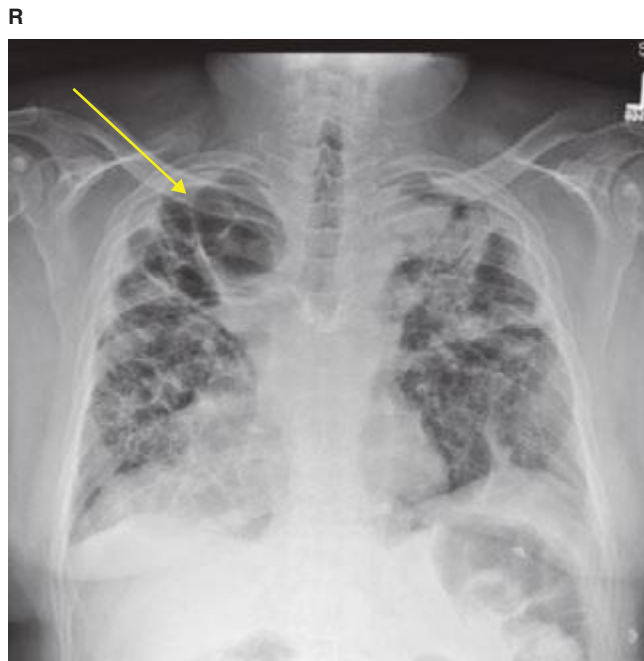


FIGURE 308e-26 Sarcoid—stage IV with fibrotic lung disease and cavitary areas (yellow arrow).

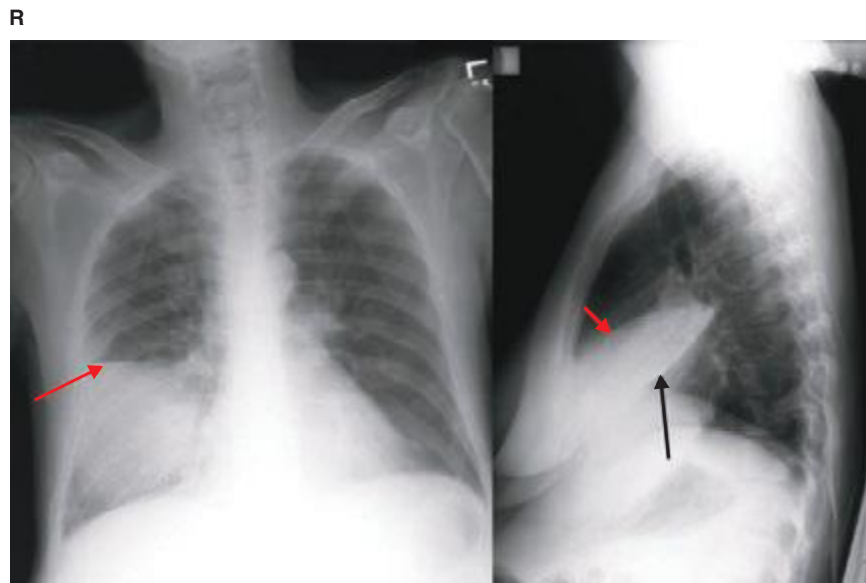


FIGURE 308e-27 Right middle lobe opacity illustrates major (*black arrow*) and minor fissures (*red arrows*) as well as the “silhouette sign” on the right heart border. The silhouette sign is the loss of clear demarcation between normal lung and soft tissue (e.g., heart, diaphragm). This occurs when the lung parenchyma is no longer filled with air and the contrast between air and soft tissue is lost.

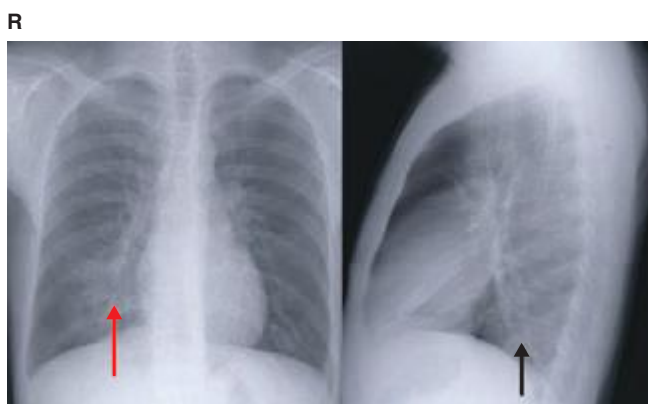


FIGURE 308e-28 Right lower lobe pneumonia—subtle opacity on PA film (*red arrow*), while the lateral film illustrates the “spine sign” (*black arrow*) where the lower spine does not become more lucent.

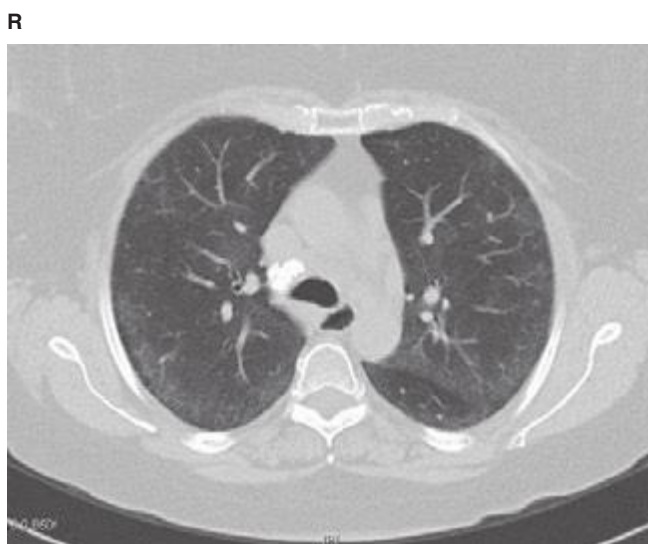


FIGURE 308e-29 CT scan of diffuse, bilateral “ground-glass” opacities. This finding is consistent with fluid density in the alveolar space.

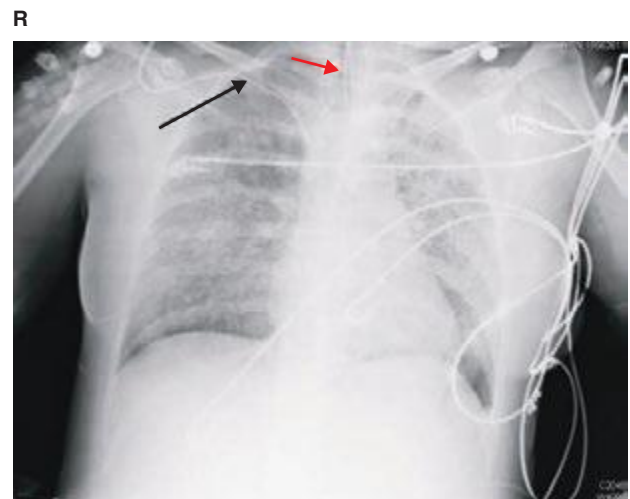


FIGURE 308e-30 Chest radiograph reveals diffuse, bilateral alveolar opacities without pleural effusions, consistent with acute respiratory distress syndrome (ARDS). Note that the patient has an endotracheal tube (*red arrow*) and a central venous catheter (*black arrow*).



FIGURE 308e-31 CT scan of ARDS demonstrates “ground-glass” opacities with more consolidated areas in the dependent lung zones.

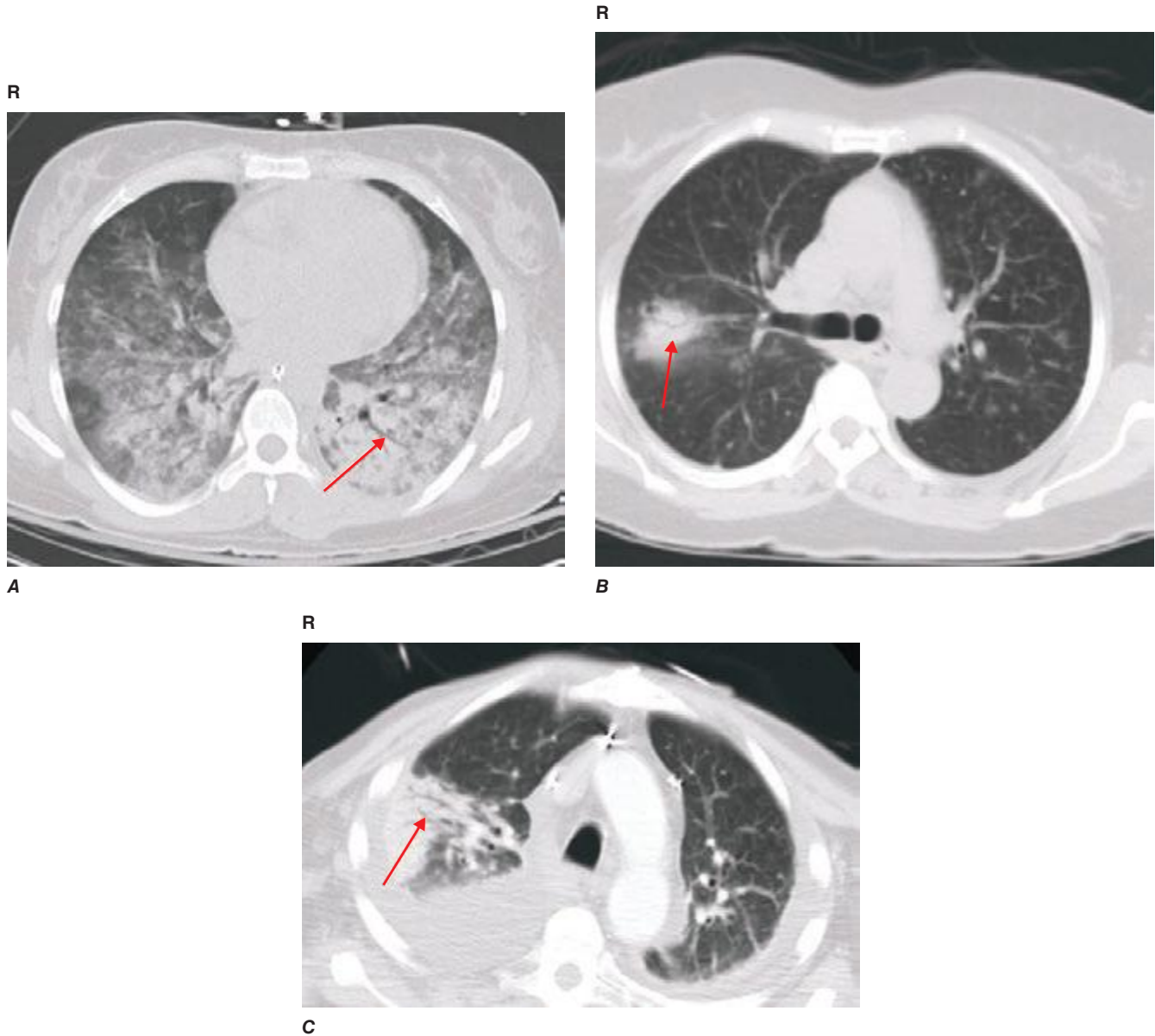


FIGURE 308e-32 Three examples of air bronchograms (red arrows) on chest CT.

BRONCHIECTASIS AND AIRWAY ABNORMALITIES

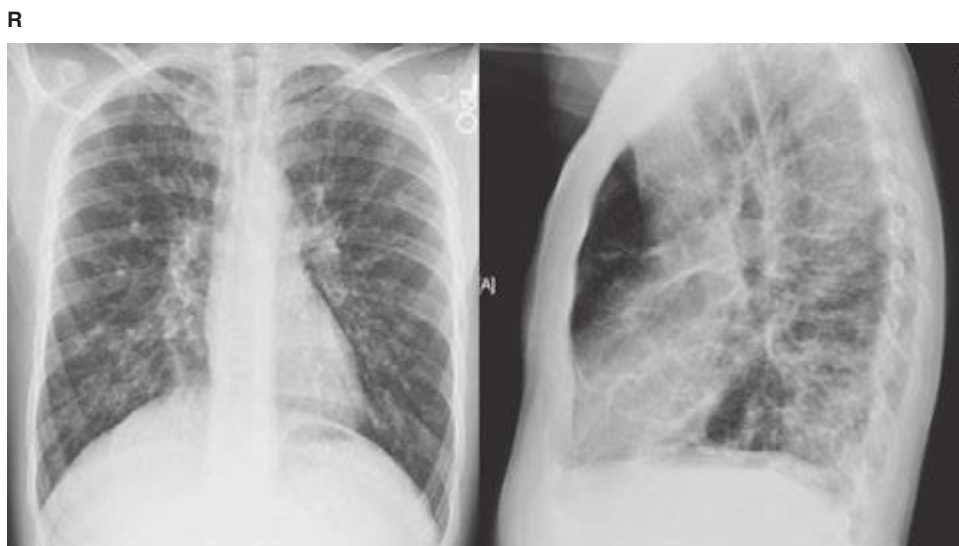


FIGURE 308e-33 Cystic fibrosis with bronchiectasis, apical disease.

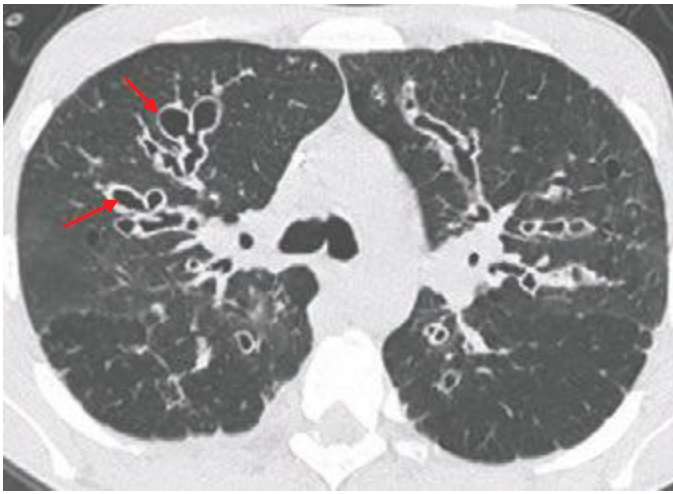


FIGURE 308e-34 CT scan of diffuse, cystic bronchiectasis (*red arrows*) in a patient with cystic fibrosis.

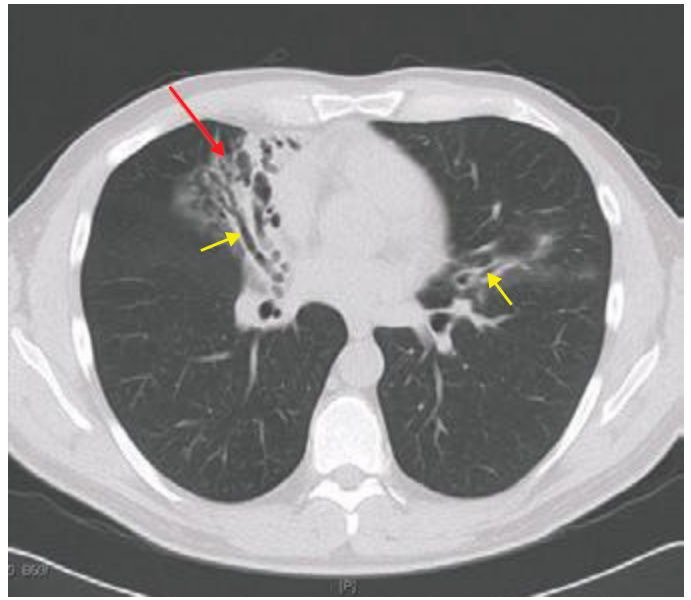


FIGURE 308e-35 CT scan of focal right middle lobe and lingular bronchiectasis (*yellow arrows*). Note that there is near total collapse of the right middle lobe (*red arrow*).

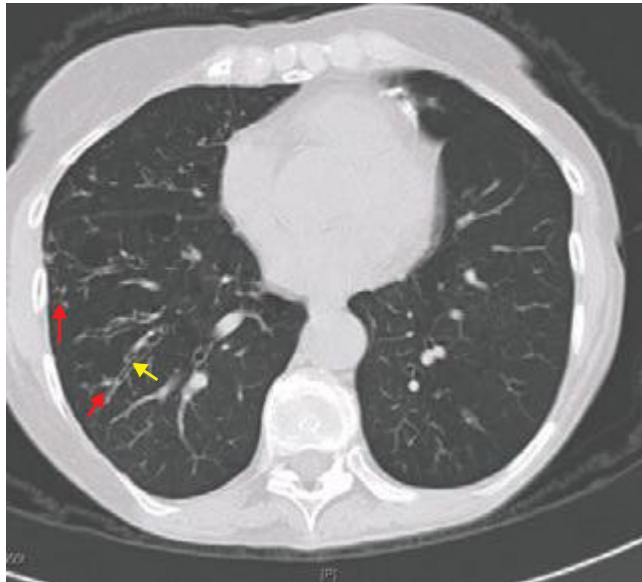


FIGURE 308e-36 "Tree in bud" opacities (*red arrows*) and bronchiectasis (*yellow arrow*) consistent with atypical mycobacterial infection. "Tree in bud" refers to small nodules clustered around the centrilobular arteries as well as increased prominence of the centrilobular branching. These findings are consistent with bronchiolitis.

R

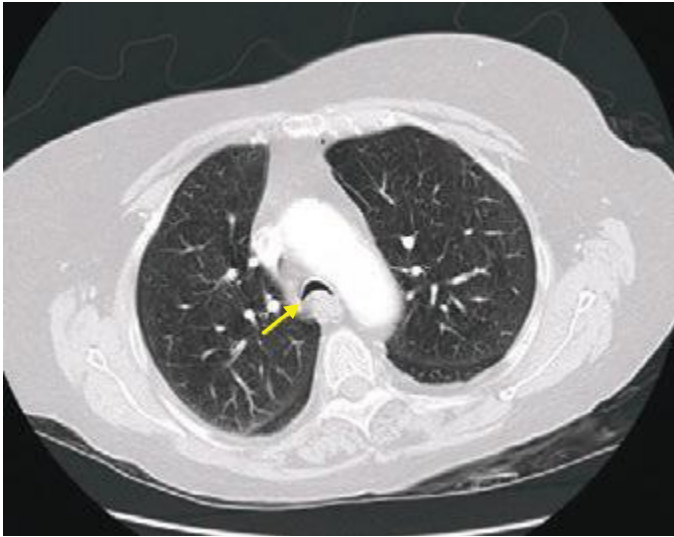


FIGURE 308e-37 CT scan demonstrating tracheomalacia (*yellow arrow*). Tracheomalacia is dynamic collapse of the trachea, most prominent during exhalation, due to loss of cartilaginous support.

R

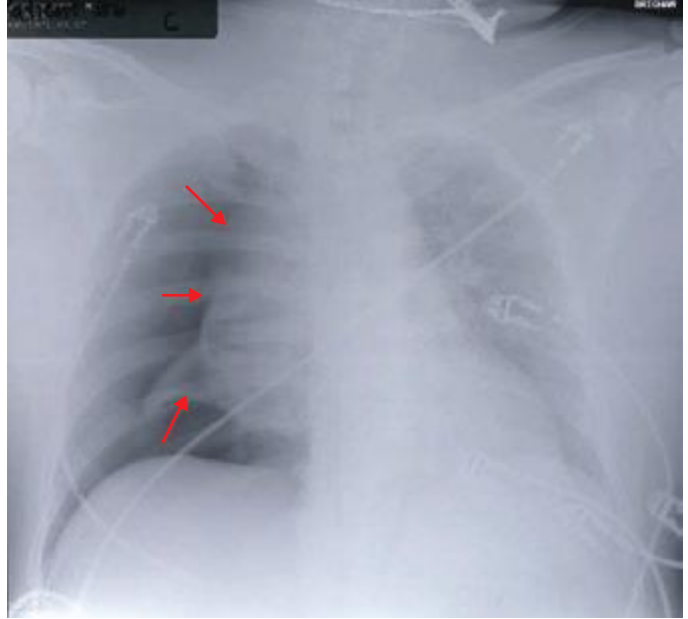


FIGURE 308e-38 Large right pneumothorax with near complete collapse of right lung. Pleural reflection highlighted with *red arrows*.

R

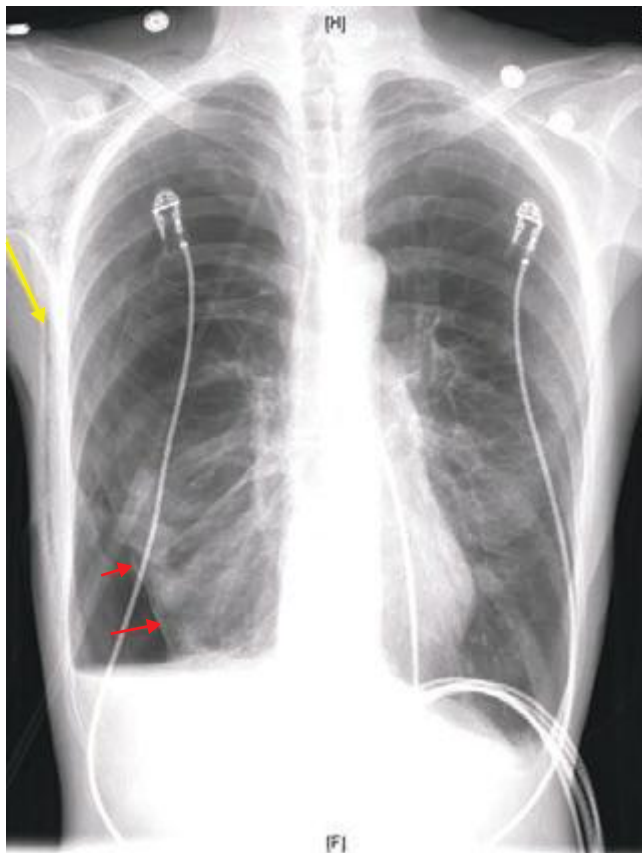


FIGURE 308e-39 Basilar pneumothorax with visible pleural reflection (*red arrows*). Also note, patient has subcutaneous emphysema (*yellow arrow*).

R

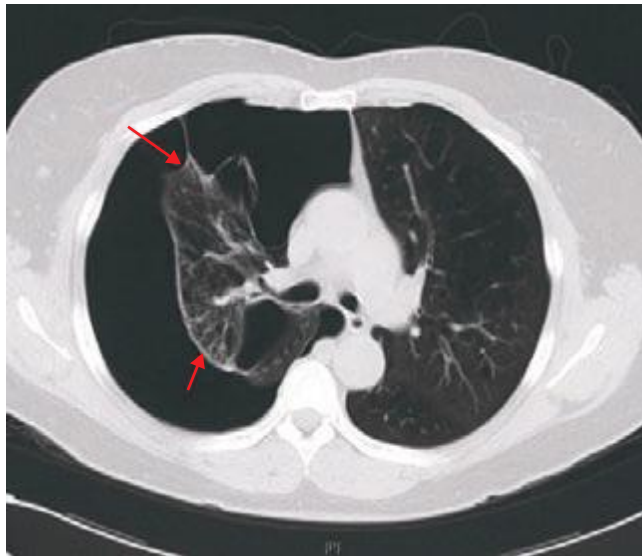


FIGURE 308e-40 CT scan of large right-sided pneumothorax. Note significant collapse of right lung with adhesion to anterior chest wall. Pleural reflection highlighted with *red arrows*. The patient has severe underlying emphysema.

R



FIGURE 308e-41 Small right pleural effusion (*red arrows* highlight blunted right costophrenic angles) with associated pleural thickening. Note fluid in the major fissure (*black arrow*) visible on the lateral film as well as the meniscus of the right pleural effusion.

R

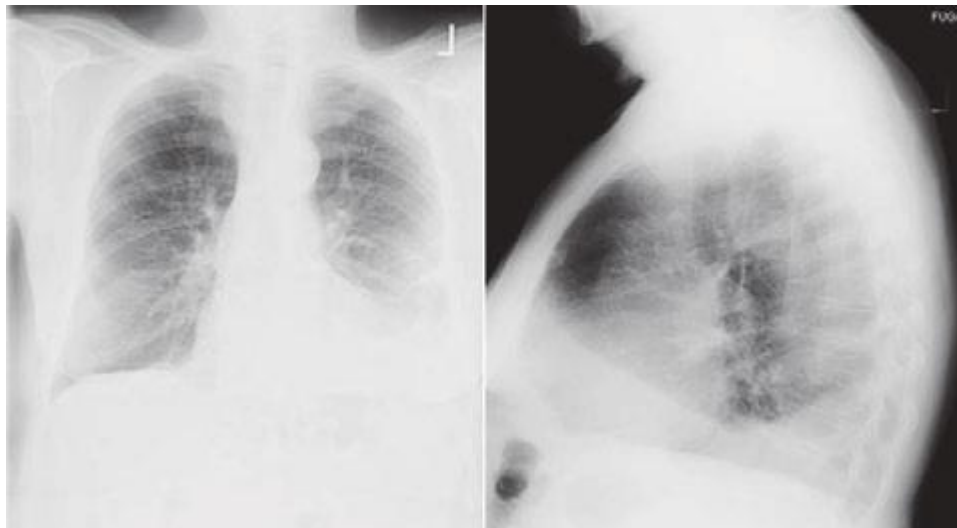


FIGURE 308e-42 Left pleural effusion with clear meniscus seen on both PA and lateral chest radiographs.

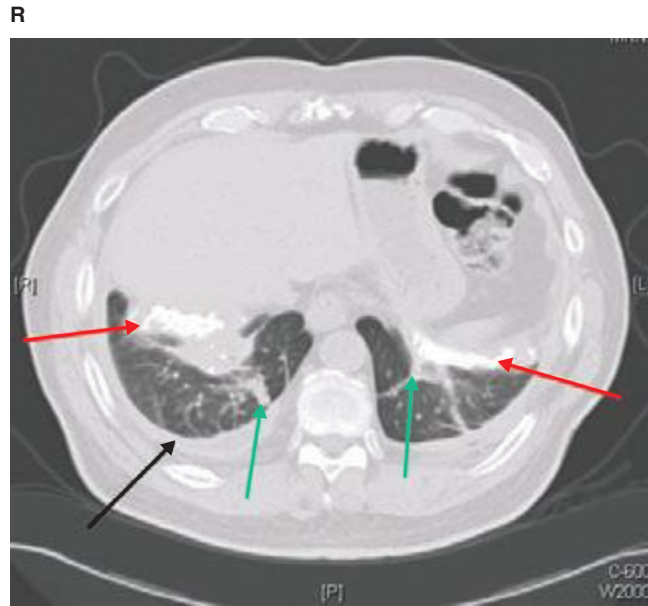


FIGURE 308e-43 Asbestosis. Note calcified pleural plaques (*red arrows*), pleural thickening (*black arrow*), and subpleural atelectasis (*green arrows*).

NODULES AND MASSES

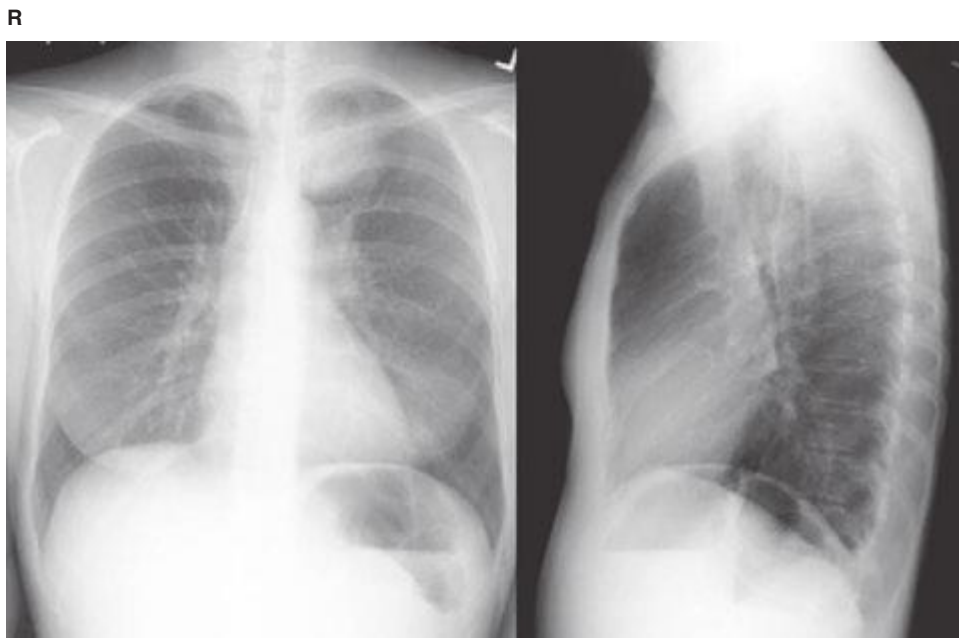


FIGURE 308e-44 Left upper lobe mass, which biopsy revealed to be squamous cell carcinoma.

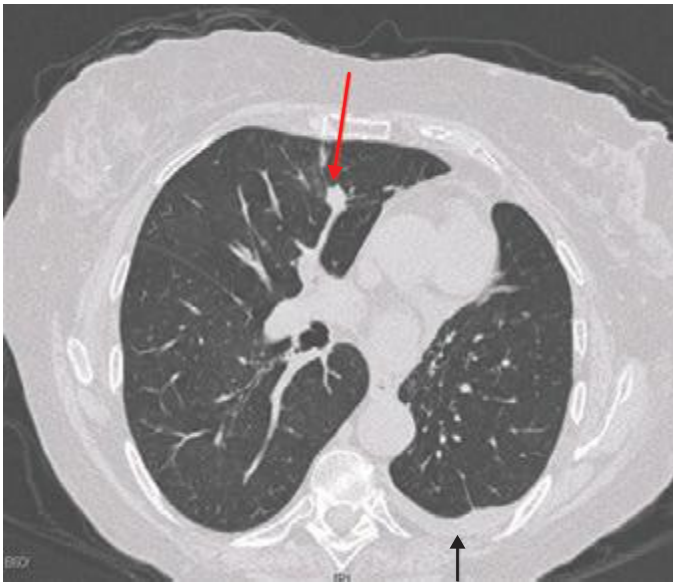


FIGURE 308e-45 Solitary pulmonary nodule on the right (*red arrow*) with a spiculated pattern concerning for lung cancer. Note also that the patient is status post left upper lobectomy with resultant volume loss and associated effusion (*black arrow*).

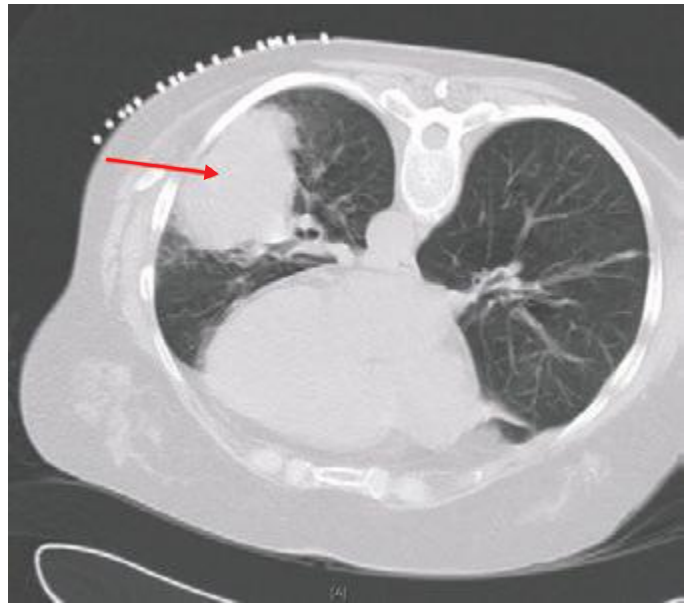


FIGURE 308e-47 Left lower lobe lung mass (*red arrow*) abutting pleura. Biopsy demonstrated small-cell lung cancer.

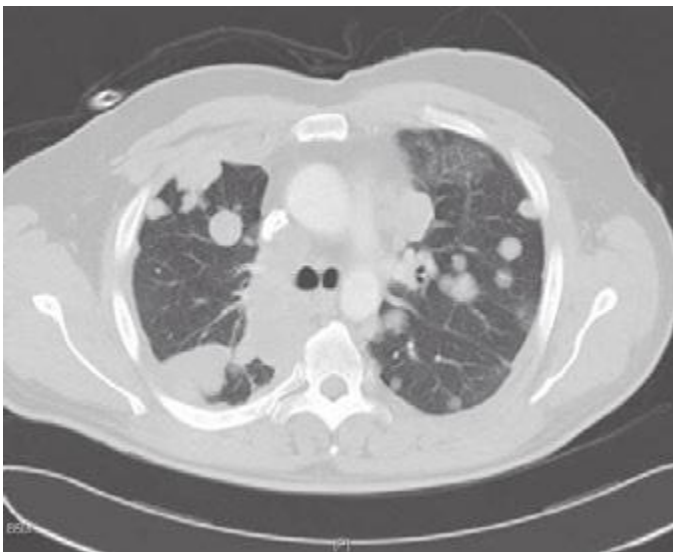


FIGURE 308e-46 Metastatic sarcoma. Note the multiple, well-circumscribed nodules of different size.



FIGURE 308e-48 CT scan of soft tissue mass encircling the trachea (*red arrow*) and invading tracheal lumen. Biopsy demonstrated adenoid cystic carcinoma (cylindroma).

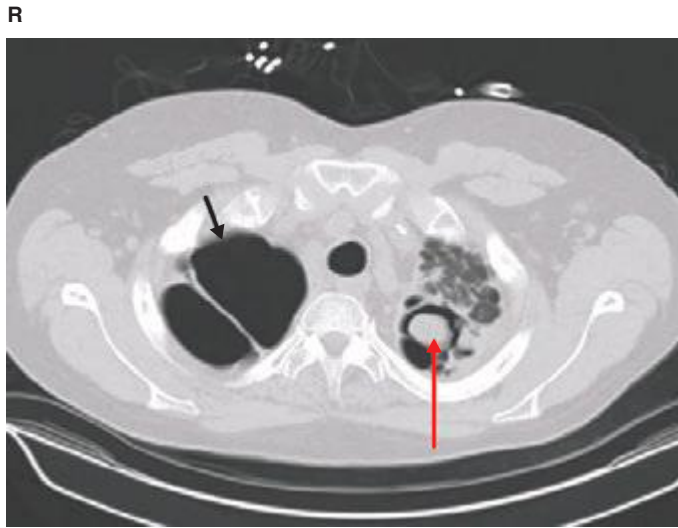


FIGURE 308e-49 Mycetoma. Fungal ball (*red arrow*) growing in preexisting cavity on the left. Right upper lobe has a large bulla (*black arrow*).

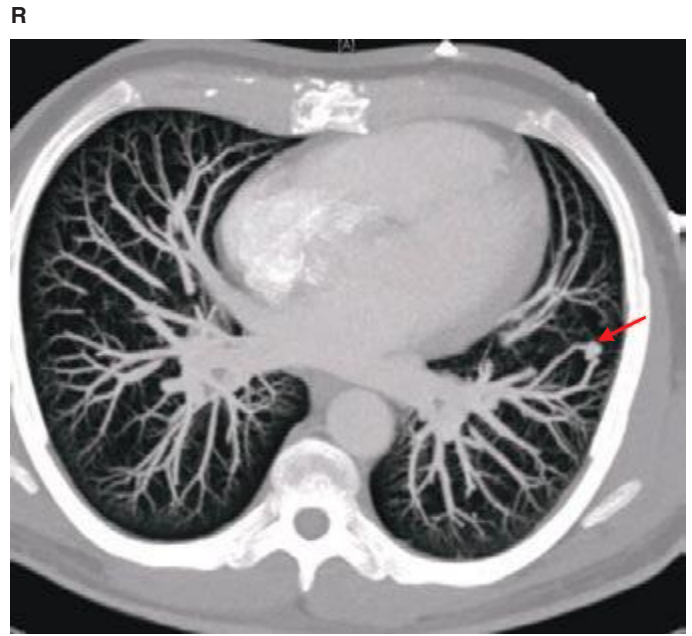


FIGURE 308e-50 Pulmonary arteriovenous malformation (AVM) demonstrated on reformatted CT angiogram (*red arrow*).

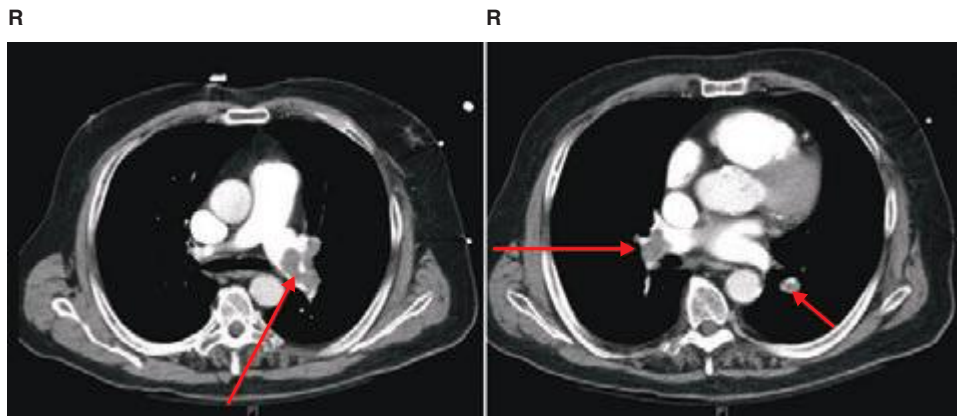


FIGURE 308e-51 Large bilateral pulmonary emboli (intravascular filling defects in contrast scan identified by *red arrows*).

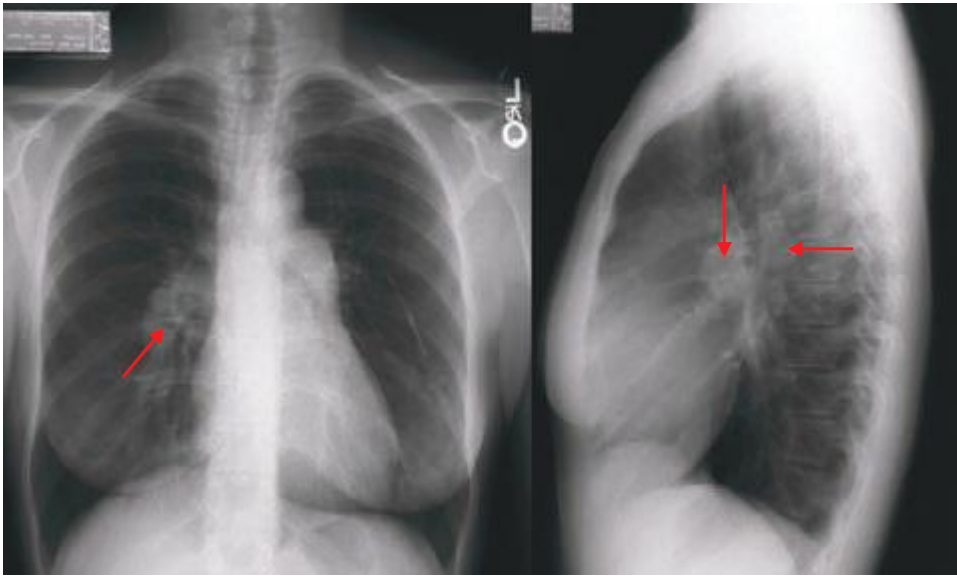


FIGURE 308e-52 Chest radiograph of a patient with severe pulmonary hypertension. Note the enlarged pulmonary arteries (*red arrows*) visible on both PA and lateral films.



FIGURE 308e-53 CT scan of the same patient as in Fig. 308e-52. Note the markedly enlarged pulmonary arteries (*red arrow*).

SECTION 2 DISEASES OF THE RESPIRATORY SYSTEM

309 Asthma

Peter J. Barnes

Asthma is a syndrome characterized by airflow obstruction that varies markedly, both spontaneously and with treatment. Asthmatics harbor a special type of inflammation in the airways that makes them more responsive than nonasthmatics to a wide range of triggers, leading to excessive narrowing with consequent reduced airflow and symptomatic wheezing and dyspnea. Narrowing of the airways is usually reversible, but in some patients with chronic asthma there may be an element of irreversible airflow obstruction. The increasing global prevalence of asthma, the large burden it now imposes on patients, and the high health care costs have led to extensive research into its mechanisms and treatment.

PREVALENCE

Asthma is one of the most common chronic diseases globally and currently affects approximately 300 million people worldwide. The prevalence of asthma has risen in affluent countries over the last 30 years but now appears to have stabilized, with approximately 10–12% of adults and 15% of children affected by the disease. In developing countries where the prevalence of asthma had been much lower, there is a rising prevalence, which is associated with increased urbanization. The prevalence of atopy and other allergic diseases has also increased over the same time, suggesting that the reasons for the increase are likely to be systemic rather than confined to the lungs. Most patients with asthma in affluent countries are atopic, with allergic sensitization to the house dust mite *Dermatophagoides pteronyssinus* and other environmental allergens, such as animal fur and pollens.

Asthma can present at any age, with a peak age of 3 years. In childhood, twice as many males as females are asthmatic, but by adulthood the sex ratio has equalized. Long-term studies that have followed children until they reach the age of 40 years suggest that many with asthma

become asymptomatic during adolescence but that asthma returns in some during adult life, particularly in those with persistent symptoms and severe asthma. Adults with asthma, including those with onset during adulthood, rarely become permanently asymptomatic. The severity of asthma does not vary significantly within a given patient; those with mild asthma rarely progress to more severe disease, whereas those with severe asthma usually have severe disease at the onset.

Deaths from asthma are uncommon, and in many affluent countries have been steadily declining over the last decade. A rise in asthma mortality seen in several countries during the 1960s was associated with increased use of short-acting inhaled β_2 -adrenergic agonists (as rescue therapy), but there is now compelling evidence that the more widespread use of inhaled corticosteroids (ICS) in patients with persistent asthma is responsible for the decrease in mortality in recent years. Major risk factors for asthma deaths are poorly controlled disease with frequent use of bronchodilator inhalers, lack of or poor compliance with ICS therapy, and previous admissions to hospital with near-fatal asthma.

It has proved difficult to agree on a definition of asthma, but there is good agreement on the description of the clinical syndrome and disease pathology. Until the etiologic mechanisms of the disease are better understood, it will be difficult to provide an accurate definition.

RISK FACTORS AND TRIGGERS

Asthma is a heterogeneous disease with interplay between genetic and environmental factors. Several risk factors that predispose to asthma have been identified (Table 309-1). These should be distinguished from triggers, which are environmental factors that worsen asthma in a patient with established disease.

Atopy Atopy is the major risk factor for asthma, and nonatopic individuals have a very low risk of developing asthma. Patients with asthma commonly suffer from other atopic diseases, particularly allergic rhinitis, which may be found in over 80% of asthmatic patients, and atopic dermatitis (eczema). Atopy may be found in 40–50% of the population in affluent countries, with only a proportion of atopic

TABLE 309-1 RISK FACTORS AND TRIGGERS INVOLVED IN ASTHMA

Endogenous Factors	Environmental Factors
Genetic predisposition	Indoor allergens
Atopy	Outdoor allergens
Airway hyperresponsiveness	Occupational sensitizers
Gender	Passive smoking
Ethnicity	Respiratory infections
Obesity	Diet
Early viral infections	Acetaminophen (paracetamol)
Triggers	
Allergens	
Upper respiratory tract viral infections	
Exercise and hyperventilation	
Cold air	
Sulfur dioxide and irritant gases	
Drugs (β blockers, aspirin)	
Stress	
Irritants (household sprays, paint fumes)	

individuals becoming asthmatic. This observation suggests that some other environmental or genetic factor(s) predispose to the development of asthma in atopic individuals. The allergens that lead to sensitization are usually proteins that have protease activity, and the most common allergens are derived from house dust mites, cat and dog fur, cockroaches (in inner cities), grass and tree pollens, and rodents (in laboratory workers). Atopy is due to the genetically determined production of specific IgE antibody, with many patients showing a family history of allergic diseases.



Genetic Predisposition The familial association of asthma and a high degree of concordance for asthma in identical twins indicate a genetic predisposition to the disease; however, whether or not the genes predisposing to asthma are similar or in addition to those predisposing to atopy is not yet clear. It now seems likely that different genes may also contribute to asthma specifically, and there is increasing evidence that the severity of asthma is also genetically determined. Genetic screens with classical linkage analysis and single-nucleotide polymorphisms of various candidate genes indicate that asthma is polygenic, with each gene identified having a small effect that is often not replicated in different populations. This observation suggests that the interaction of many genes is important, and these may differ in different populations. The most consistent findings have been associations with polymorphisms of genes on chromosome 5q, including the T helper 2 (T_H2) cells interleukin (IL)-4, IL-5, IL-9, and IL-13, which are associated with atopy. There is increasing evidence for a complex interaction between genetic polymorphisms and environmental factors that will require very large population studies to unravel. Novel genes that have been associated with asthma, including *ADAM-33*, and *DPP-10*, have also been identified by positional cloning, but their function in disease pathogenesis is not yet clear. Recent genome-wide association studies have identified further novel genes, such as *ORMDL3*, although their functional role is not yet clear. Genetic polymorphisms may also be important in determining the response to asthma therapy. For example, the Arg-Gly-16 variant in the β_2 -receptor has been associated with reduced response to β_2 -agonists, and repeats of an Sp1 recognition sequence in the promoter region of 5-lipoxygenase may affect the response to antileukotrienes. However, these effects are small and inconsistent and do not yet have any implications for asthma therapy.

It is likely that environmental factors in early life determine which atopic individuals become asthmatic. The increasing prevalence of asthma, particularly in developing countries, over the last few decades also indicates the importance of environmental mechanisms interacting with a genetic predisposition.

Infections Although viral infections (especially rhinovirus) are common triggers of asthma exacerbations, it is uncertain whether they play a role in etiology. There is some association between respiratory syncytial virus infection in infancy and the development of asthma, but the specific pathogenesis is difficult to elucidate because this infection is very common in children. Atypical bacteria, such as *Mycoplasma* and *Chlamydia*, have been implicated in the mechanism of severe asthma, but thus far, the evidence is not very convincing of a true association.

The observation that allergic sensitization and asthma were less common in children with older siblings first suggested that lower levels of infection may be a factor in affluent societies that increase the risks of asthma. This “hygiene hypothesis” proposes that lack of infections in early childhood preserves the T_H2 cell bias at birth, whereas exposure to infections and endotoxin results in a shift toward a predominant protective T_H1 immune response. Children brought up on farms who are exposed to a high level of endotoxin are less likely to develop allergic sensitization than children raised on dairy farms. Intestinal parasite infection, such as hookworm, may also be associated with a reduced risk of asthma. Although there is considerable epidemiologic support for the hygiene hypothesis, it cannot account for the parallel increase in T_H1 -driven diseases such as diabetes mellitus over the same period.

Diet The role of dietary factors is controversial. Observational studies have shown that diets low in antioxidants such as vitamin C and vitamin A, magnesium, selenium, and omega-3 polyunsaturated fats (fish oil) or high in sodium and omega-6 polyunsaturated fats are associated with an increased risk of asthma. Vitamin D deficiency may also predispose to the development of asthma. However, interventional studies with supplementary diets have not supported an important role for these dietary factors. Obesity is also an independent risk factor for asthma, particularly in women, but the mechanisms are thus far unknown.

Air Pollution Air pollutants, such as sulfur dioxide, ozone, and diesel particulates, may trigger asthma symptoms, but the role of different air pollutants in the etiology of the disease is much less certain. Most evidence argues against an important role for air pollution because asthma is no more prevalent in cities with a high ambient level of traffic pollution than in rural areas with low levels of pollution. Asthma had a much lower prevalence in East Germany compared to West Germany despite a much higher level of air pollution, but since reunification, these differences have decreased as eastern Germany has become more affluent. Indoor air pollution may be more important with exposure to nitrogen oxides from cooking stoves and exposure to passive cigarette smoke. There is some evidence that maternal smoking is a risk factor for asthma, but it is difficult to dissociate this association from an increased risk of respiratory infections.

Allergens Inhaled allergens are common triggers of asthma symptoms and have also been implicated in allergic sensitization. Exposure to house dust mites in early childhood is a risk factor for allergic sensitization and asthma, but rigorous allergen avoidance has not shown any evidence for a reduced risk of developing asthma. The increase in house dust mites in centrally heated poorly ventilated homes with fitted carpets has been implicated in the increasing prevalence of asthma in affluent countries. Domestic pets, particularly cats, have also been associated with allergic sensitization, but early exposure to cats in the home may be protective through the induction of tolerance.

Occupational Exposure Occupational asthma is relatively common and may affect up to 10% of young adults. Over 300 sensitizing agents have been identified. Chemicals such as toluene diisocyanate and trimellitic anhydride, may lead to sensitization independent of atopy. Individuals may also be exposed to allergens in the workplace such as small animal allergens in laboratory workers and fungal amylase in wheat flour in bakers. Occupational asthma may be suspected when symptoms improve during weekends and holidays.

Obesity Asthma occurs more frequently in obese people (body mass index >30 kg/m²) and is often more difficult to control. Although

mechanical factors may contribute, it may also be linked to the pro-inflammatory adipokines and reduced anti-inflammatory adipokines that are released from fat stores.

Other Factors Several other factors have been implicated in the etiology of asthma, including lower maternal age, duration of breast-feeding, prematurity and low birthweight, and inactivity, but are unlikely to contribute to the recent global increase in asthma prevalence. There is also an association with acetaminophen (paracetamol) consumption in childhood, which may be linked to increased oxidative stress.

Intrinsic Asthma A minority of asthmatic patients (approximately 10%) have negative skin tests to common inhalant allergens and normal serum concentrations of IgE. These patients, with nonatopic or intrinsic asthma, usually show later onset of disease (adult-onset asthma), commonly have concomitant nasal polyps, and may be aspirin-sensitive. They usually have more severe, persistent asthma. Little is understood about mechanism, but the immunopathology in bronchial biopsies and sputum appears to be identical to that found in atopic asthma. There is recent evidence for increased local production of IgE in the airways, suggesting that there may be common IgE-mediated mechanisms; staphylococcal enterotoxins, which serve as “superantigens,” have been implicated.

Asthma Triggers Several stimuli trigger airway narrowing, wheezing, and dyspnea in asthmatic patients. Although a previous view held that these stimuli should be avoided, the triggering of asthma by these stimuli is now seen as evidence for poor control and an indicator of the need to increase controller (preventive) therapy.

ALLERGENS Inhaled allergens activate mast cells with bound IgE directly leading to the immediate release of bronchoconstrictor mediators, resulting in the early response that is reversed by bronchodilators. Often, experimental allergen challenge is followed by a late response when there is airway edema and an acute inflammatory response with increased eosinophils and neutrophils that are not very reversible with bronchodilators. The most common allergens to trigger asthma are *Dermatophagoides* species, and environmental exposure leads to low-grade chronic symptoms that are perennial. Other perennial allergens are derived from cats and other domestic pets, as well as cockroaches. Other allergens, including grass pollen, ragweed, tree pollen, and fungal spores, are seasonal. Pollens usually cause allergic rhinitis rather than asthma, but in thunderstorms, the pollen grains are disrupted and the particles that may be released can trigger severe asthma exacerbations (thunderstorm asthma).

VIRUS INFECTIONS Upper respiratory tract virus infections such as rhinovirus, respiratory syncytial virus, and coronavirus are the most common triggers of acute severe exacerbations and may invade epithelial cells of the lower as well as the upper airways. The mechanism whereby these viruses cause exacerbations is poorly understood, but there is an increase in airway inflammation with increased numbers of eosinophils and neutrophils. There is evidence for reduced production of type I interferons by epithelial cells from asthmatic patients, resulting in increased susceptibility to these viral infections and a greater inflammatory response.

PHARMACOLOGIC AGENTS Several drugs may trigger asthma. Beta-adrenergic blockers commonly acutely worsen asthma, and their use may be fatal. The mechanisms are not clear, but are likely mediated through increased cholinergic bronchoconstriction. All β blockers need to be avoided, and even selective β_2 blockers or topical application (e.g., timolol eye drops) may be dangerous. Angiotensin-converting enzyme inhibitors are theoretically detrimental as they inhibit breakdown of kinins, which are bronchoconstrictors; however, they rarely worsen asthma, and the characteristic cough is no more frequent in asthmatics than in nonasthmatics. Aspirin may worsen asthma in some patients (aspirin-sensitive asthma is discussed below under “Special Considerations”).

EXERCISE Exercise is a common trigger of asthma, particularly in children. The mechanism is linked to hyperventilation, which results

in increased osmolality in airway lining fluid and triggers mast cell mediator release, resulting in bronchoconstriction. Exercise-induced asthma (EIA) typically begins after exercise has ended and resolves spontaneously within about 30 min. EIA is worse in cold, dry climates than in hot, humid conditions. It is, therefore, more common in sports activities such as cross-country running in cold weather, overland skiing, and ice hockey than in swimming. It may be prevented by prior administration of β_2 -agonists and antileukotrienes, but is best prevented by regular treatment with ICSs, which reduce the population of surface mast cells required for this response.

PHYSICAL FACTORS Cold air and hyperventilation may trigger asthma through the same mechanisms as exercise. Laughter may also be a trigger. Many patients report worsening of asthma in hot weather and when the weather changes. Some asthmatics become worse when exposed to strong smells or perfumes, but the mechanism of this response is uncertain.

FOOD AND DIET There is little evidence that allergic reactions to food lead to increased asthma symptoms, despite the belief of many patients that their symptoms are triggered by particular food constituents. Exclusion diets are usually unsuccessful at reducing the frequency of episodes. Some foods such as shellfish and nuts may induce anaphylactic reactions that may include wheezing. Patients with aspirin-induced asthma may benefit from a salicylate-free diet, but these are difficult to maintain. Certain food additives may trigger asthma. Metabisulfite, which is used as a food preservative, may trigger asthma through the release of sulfur dioxide gas in the stomach. Tartrazine, a yellow food-coloring agent, was believed to be a trigger for asthma, but there is little convincing evidence for this.

AIR POLLUTION Increased ambient levels of sulfur dioxide, ozone, and nitrogen oxides are associated with increased asthma symptoms.

OCCUPATIONAL FACTORS Several substances found in the workplace may act as sensitizing agents, as discussed above, but may also act as triggers of asthma symptoms. Occupational asthma is characteristically associated with symptoms at work with relief on weekends and holidays. If removed from exposure within the first 6 months of symptoms, there is usually complete recovery. More persistent symptoms lead to irreversible airway changes, and thus, early detection and avoidance are important.

HORMONES Some women show premenstrual worsening of asthma, which can occasionally be very severe. The mechanisms are not completely understood, but are related to a fall in progesterone and in severe cases may be improved by treatment with high doses of progesterone or gonadotropin-releasing factors. Thyrotoxicosis and hypothyroidism can both worsen asthma, although the mechanisms are uncertain.

GASTROESOPHAGEAL REFLUX Gastroesophageal reflux is common in asthmatic patients because it is increased by bronchodilators. Although acid reflux might trigger reflex bronchoconstriction, it rarely causes asthma symptoms, and antireflux therapy usually fails to reduce asthma symptoms in most patients.

STRESS Many asthmatics report worsening of symptoms with stress. Psychological factors can induce bronchoconstriction through cholinergic reflex pathways. Paradoxically, very severe stress such as bereavement usually does not worsen, and may even improve, asthma symptoms.

PATHOPHYSIOLOGY

Asthma is associated with a specific chronic inflammation of the mucosa of the lower airways. One of the main aims of treatment is to reduce this inflammation.

Pathology The pathology of asthma has been revealed through examining the lungs of patients who have died of asthma and from bronchial biopsies. The airway mucosa is infiltrated with activated eosinophils and T lymphocytes, and there is activation of mucosal mast cells. The degree of inflammation is poorly related to disease

1672 severity and may even be found in atopic patients without asthma symptoms. This inflammation is usually reduced by treatment with ICS. There are also structural changes in the airways (described as remodeling). A characteristic finding is thickening of the basement membrane due to subepithelial collagen deposition. This feature is also found in patients with eosinophilic bronchitis presenting as cough who do not have asthma and is, therefore, likely to be a marker of eosinophilic inflammation in the airway as eosinophils release fibrogenic mediators. The epithelium is often shed or friable, with reduced attachments to the airway wall and increased numbers of epithelial cells in the lumen. The airway wall itself may be thickened and edematous, particularly in fatal asthma. Another common finding in fatal asthma is occlusion of the airway lumen by a mucous plug, which is comprised of mucous glycoproteins secreted from goblet cells and plasma proteins from leaky bronchial vessels (Fig. 309-1). There is also vasodilation and increased numbers of blood vessels (angiogenesis). Direct observation by bronchoscopy indicates that the airways may be narrowed, erythematous, and edematous. The pathology of asthma is remarkably uniform in different phenotypes of asthma, including atopic (extrinsic), nonatopic (intrinsic), occupational, aspirin-sensitive, and pediatric asthma. These pathologic changes are found in all airways, but do not extend to the lung parenchyma; peripheral airway inflammation is found particularly in patients with severe asthma. The involvement of airways may be patchy, and this is consistent with bronchographic findings of uneven narrowing of the airways.

Airway Inflammation There is inflammation in the respiratory mucosa from the trachea to terminal bronchioles, but with a predominance in the bronchi (cartilaginous airways); however, it is still uncertain as to how inflammatory cells interact and how inflammation translates into the symptoms of asthma (Fig. 309-2). There is good evidence that the specific pattern of airway inflammation in asthma is associated with airway hyperresponsiveness (AHR), the physiologic abnormality of asthma, which is correlated with variable airflow obstruction. The pattern of inflammation in asthma is characteristic of allergic diseases, with similar inflammatory cells seen in the nasal mucosa in rhinitis. However, an indistinguishable pattern of inflammation is found in intrinsic asthma, and this may reflect local rather than systemic IgE production. Although most attention has focused on the acute inflammatory changes seen in asthma, this is a chronic condition, with inflammation persisting over many years in most patients. The mechanisms involved in persistence of inflammation in asthma are still poorly understood. Superimposed on this chronic inflammatory state are acute inflammatory episodes, which correspond to exacerbations of asthma. Although the common pattern of inflammation in asthma is characterized by eosinophil infiltration, some patients with severe asthma show a neutrophilic pattern of inflammation that is less sensitive to corticosteroids. However, many inflammatory cells are involved in asthma with no key cell that is predominant (Fig. 309-3).

MAST CELLS Mast cells are important in initiating the acute bronchoconstrictor responses to allergens and several other indirectly acting stimuli, such as exercise and hyperventilation (via osmolality changes), as well as fog. Activated mucosal mast cells are found at the airway surface in asthma patients and also in the airway smooth-muscle layer, whereas this is not seen in normal subjects or patients with eosinophilic bronchitis. Mast cells are activated by allergens through an IgE-dependent mechanism, and binding of specific IgE to mast cells renders them more sensitive to activation by physical stimuli such as osmolality. The importance of IgE in the pathophysiology of asthma has been highlighted by clinical studies with humanized anti-IgE antibodies, which inhibit IgE-mediated effects, reduce asthma symptoms, and reduce exacerbations. There are, however, uncertainties about the

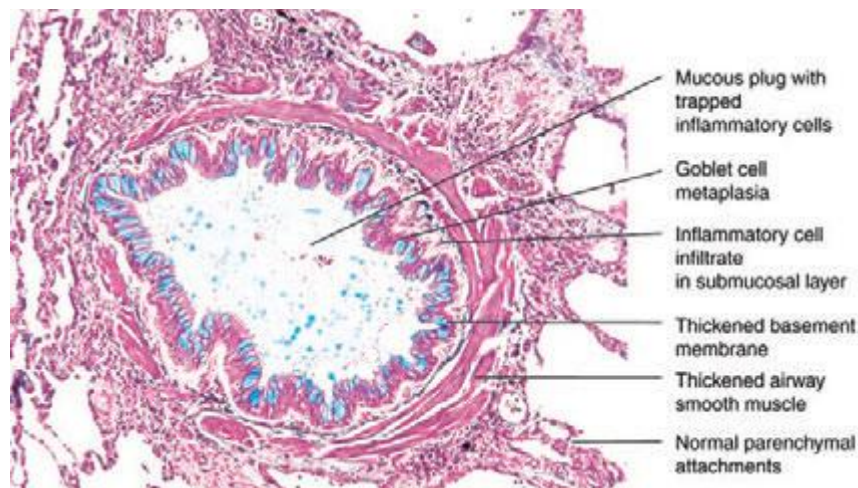


FIGURE 309-1 Histopathology of a small airway in fatal asthma. The lumen is occluded with a mucous plug, there is goblet cell metaplasia, and the airway wall is thickened, with an increase in basement membrane thickness and airway smooth muscle. (Courtesy of Dr. J. Hogg, University of British Columbia.)

role of mast cells in more chronic allergic inflammatory events. Mast cells release several bronchoconstrictor mediators, including histamine, prostaglandin D_2 , and cysteinyl-leukotrienes, but also several cytokines, chemokines, growth factors, and neurotrophins.

MACROPHAGES AND DENDRITIC CELLS Macrophages, which are derived from blood monocytes, may traffic into the airways in asthma and may be activated by allergens via low-affinity IgE receptors ($Fc\epsilon RII$). Macrophages have the capacity to initiate a type of inflammatory response via the release of a certain pattern of cytokines, but these cells also release anti-inflammatory mediators (e.g., IL-10), and thus, their roles in asthma are uncertain. Dendritic cells are specialized macrophage-like cells in the airway epithelium, which are the major antigen-presenting cells. Dendritic cells take up allergens, process them to peptides, and migrate to local lymph nodes where they present the allergenic peptides to uncommitted T lymphocytes to program the production of allergen-specific T cells. Immature dendritic cells in the respiratory tract promote T_H2 cell differentiation and require cytokines, such as IL-12 and tumor necrosis factor α (TNF- α), to promote the normally preponderant T_H1 response. The cytokine thymic stromal lymphopoietin (TSLP) released from epithelial cells in asthmatic patients instructs dendritic cells to release chemokines that attract T_H2 cells into the airways.

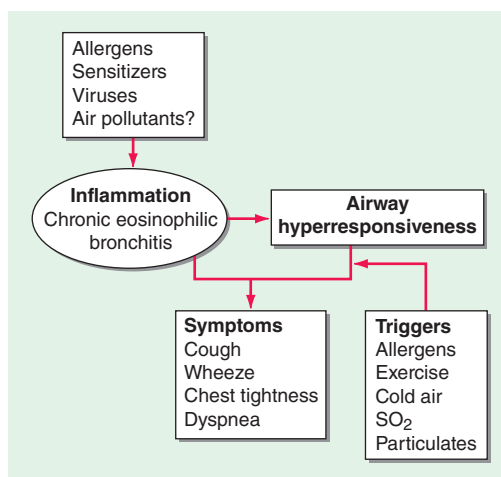


FIGURE 309-2 Inflammation in the airways of asthmatic patients leads to airway hyperresponsiveness and symptoms. SO_2 , sulfur dioxide.

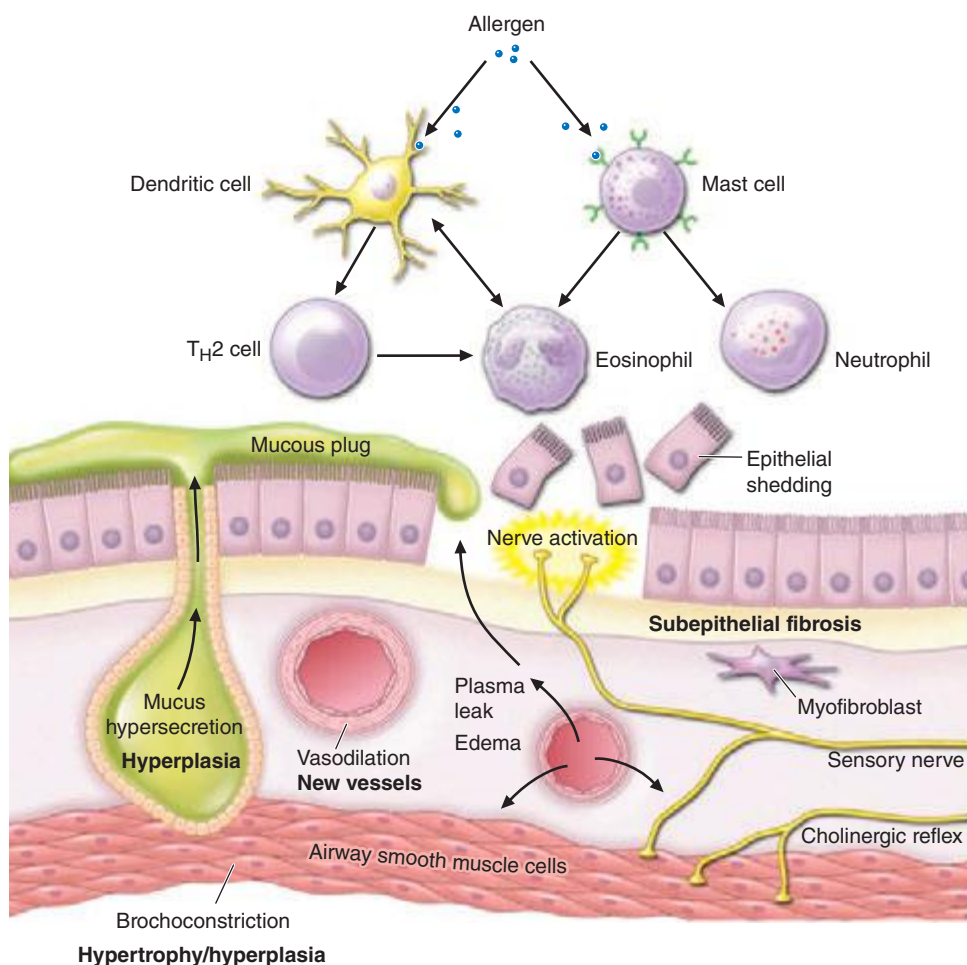


FIGURE 309-3 The pathophysiology of asthma is complex with participation of several interacting inflammatory cells, which result in acute and chronic inflammatory effects on the airway.

EOSINOPHILS Eosinophil infiltration is a characteristic feature of asthmatic airways. Allergen inhalation results in a marked increase in activated eosinophils in the airways at the time of the late reaction. Eosinophils are linked to the development of AHR through the release of basic proteins and oxygen-derived free radicals. Eosinophil recruitment involves adhesion of eosinophils to vascular endothelial cells in the airway circulation due to interaction between adhesion molecules, migration into the submucosa under the direction of chemokines, and their subsequent activation and prolonged survival. Blocking antibodies to IL-5 causes a profound and prolonged reduction in circulating and sputum eosinophils, but is not associated with reduced AHR or asthma symptoms, although in selected patients with steroid-resistant airway eosinophils, there is a reduction in exacerbations. Eosinophils may be important in release of growth factors involved in airway remodeling and in exacerbations but probably not in AHR.

NEUTROPHILS Increased numbers of activated neutrophils are found in sputum and airways of some patients with severe asthma and during exacerbations, although there is a proportion of patients even with mild or moderate asthma who have a predominance of neutrophils. The roles of neutrophils in asthma that are resistant to the anti-inflammatory effects of corticosteroids are currently unknown.

T LYMPHOCYTES T lymphocytes play a very important role in coordinating the inflammatory response in asthma through the release of specific patterns of cytokines, resulting in the recruitment and survival of eosinophils and in the maintenance of a mast cell population in the airways. The naïve immune system and the immune system of asthmatics are skewed to express the T_H2 phenotype, whereas in normal airways, T_H1 cells predominate. T_H2 cells, through the release of IL-5, are associated with eosinophilic inflammation and, through the release of IL-4 and IL-13, are associated with increased IgE formation. Recently,

bronchial biopsies have demonstrated a preponderance of natural killer CD4+ T lymphocytes that express high levels of IL-4. Regulatory T cells play an important role in determining the expression of other T cells, and there is evidence for a reduction in a certain subset of regulatory T cells (CD4+CD25+) in asthma that is associated with increased T_H2 cells. Recently, innate T cells (ILC2) without T cell receptors have been identified that release T_H2 cytokines and are regulated by epithelial cytokines, such as IL-25 and IL-33.

STRUCTURAL CELLS Structural cells of the airways, including epithelial cells, fibroblasts, and airway smooth-muscle cells, are also important sources of inflammatory mediators, such as cytokines and lipid mediators, in asthma. Indeed, because structural cells far outnumber inflammatory cells, they may become the major sources of mediators driving chronic inflammation in asthmatic airways. In addition, epithelial cells may have key roles in translating inhaled environmental signals into an airway inflammatory response and are probably major target cells for ICS.

Inflammatory Mediators Multiple inflammatory mediators have been implicated in asthma, and they may have a variety of effects on the airways that account for the pathologic features of asthma (Fig. 309-4). Mediators such as histamine, prostaglandin D_2 , and cysteinyl-leukotrienes contract airway smooth muscle, increase

microvascular leakage, increase airway mucus secretion, and attract other inflammatory cells. Because each mediator has many effects, the role of individual mediators in the pathophysiology of asthma is not yet clear. Although the multiplicity of mediators makes it unlikely that preventing the synthesis or action of a single mediator will have a major impact in clinical asthma, recent clinical studies with antileukotrienes suggest that cysteinyl-leukotrienes have clinically important effects.

CYTOKINES Multiple cytokines regulate the chronic inflammation of asthma. The T_H2 cytokines IL-4, IL-5, and IL-13 mediate allergic inflammation, whereas proinflammatory cytokines, such as TNF- α and IL-1 β , amplify the inflammatory response and play a role in more

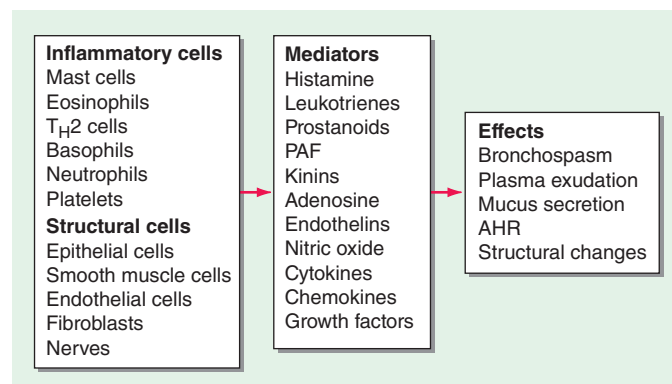


FIGURE 309-4 Many cells and mediators are involved in asthma and lead to several effects on the airways. AHR, airway hyperresponsiveness; PAF, platelet-activating factor.

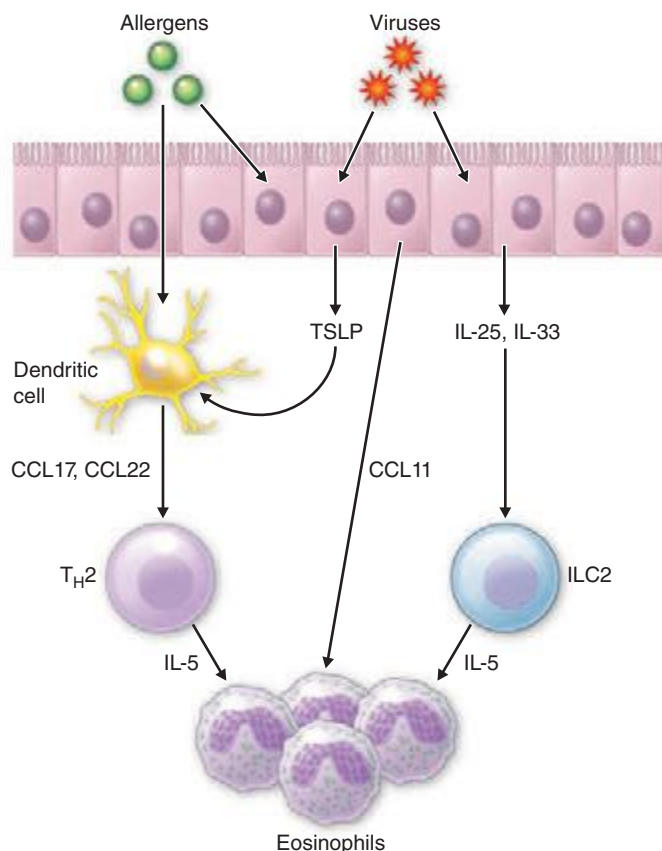


FIGURE 309-5 T lymphocytes in asthma. Allergen interacts with dendritic cells and releases thymus stimulated lymphopoeitin (TSLP), which stimulates activated dendritic cells to release the chemokines CCL17 and CCL22, which attract T helper 2 (T_H2) lymphocytes. Allergens and viral infection may release interleukin (IL)-25 and -33, which recruit and activate type 2 innate lymphoid cells (ILC2). Both T_H2 and ILC2 cells release IL-5 and epithelial cells release CCL11 (eotaxin), which together lead to recruitment of eosinophils into the airways.

severe disease. TSLP is an upstream cytokine released from epithelial cells of asthmatics that orchestrates the release of chemokines that selectively attract T_H2 cells. Some cytokines such as IL-10 and IL-12 are anti-inflammatory and may be deficient in asthma.

CHEMOKINES Chemokines are involved in attracting inflammatory cells from the bronchial circulation into the airways. Eotaxin (CCL11) is selectively attractant to eosinophils via CCR3 and is expressed by epithelial cells of asthmatics, whereas CCL17 (TARC) and CCL22 (MDC) from epithelial cells attract T_H2 cells via CCR4 (Fig. 309-5).

OXIDATIVE STRESS Activated inflammatory cells such as macrophages and eosinophils produce reactive oxygen species. Evidence for increased oxidative stress in asthma is provided by the increased concentrations of 8-isoprostane (a product of oxidized arachidonic acid) in exhaled breath condensates and increased ethane (a product of lipid peroxidation) in the expired air of asthmatic patients. Increased oxidative stress is related to disease severity, may amplify the inflammatory response, and may reduce responsiveness to corticosteroids.

NITRIC OXIDE Nitric oxide (NO) is produced by NO synthases in several cells in the airway, particularly airway epithelial cells and macrophages. The level of NO in the expired air of patients with asthma is higher than normal and is related to the eosinophilic inflammation. Increased NO may contribute to the bronchial vasodilation observed in asthma. Fractional exhaled NO (F_{eNO}) is increasingly used in the diagnosis and monitoring of asthmatic inflammation, although it is not yet used routinely in clinical practice.

TRANSCRIPTION FACTORS Proinflammatory transcription factors, such as nuclear factor- κ B (NF- κ B) and activator protein-1, are activated in asthmatic airways and orchestrate the expression of multiple inflammatory genes. More specific transcription factors that are involved include nuclear factor of activated T cells and GATA-3, which regulate the expression of T_H2 cytokines in T cells.

Effects of Inflammation The chronic inflammatory response has several effects on the target cells of the airways, resulting in the characteristic pathophysiologic and remodeling changes associated with asthma. Asthma may be regarded as a disease with continuous inflammation and repair proceeding simultaneously, although the relationship between chronic inflammatory processes and asthma symptoms is often obscure.

AIRWAY EPITHELIUM Airway epithelial shedding may be important in contributing to AHR and may explain how several mechanisms, such as ozone exposure, virus infections, chemical sensitizers, and allergens (usually proteases), can lead to its development, as all of these stimuli may lead to epithelial disruption. Epithelial damage may contribute to AHR in a number of ways, including loss of its barrier function to allow penetration of allergens; loss of enzymes (such as neutral endopeptidase) that degrade certain peptide inflammatory mediators; loss of a relaxant factor (so called epithelial-derived relaxant factor); and exposure of sensory nerves, which may lead to reflex neural effects on the airway.

FIBROSIS In all asthmatic patients, the basement membrane is apparently thickened due to subepithelial fibrosis with deposition of types III and V collagen below the true basement membrane and is associated with eosinophil infiltration, presumably through the release of profibrotic mediators such as transforming growth factor- β . Mechanical manipulations can alter the phenotype of airway epithelial cells in a profibrotic fashion. In more severe patients, there is also fibrosis within the airway wall, which may contribute to irreversible narrowing of the airways.

AIRWAY SMOOTH MUSCLE In vitro airway smooth muscle from asthmatic patients usually shows no increased responsiveness to constrictors. Reduced responsiveness to β -agonists has also been reported in post-mortem or surgically removed bronchi from asthmatics, although the number of β -receptors is not reduced, suggesting that β -receptors have been uncoupled. These abnormalities of airway smooth muscle may be secondary to the chronic inflammatory process. Inflammatory mediators may modulate the ion channels that serve to regulate the resting membrane potential of airway smooth-muscle cells, thus altering the level of excitability of these cells. In asthmatic airways there is also a characteristic hypertrophy and hyperplasia of airway smooth muscle, which is presumably the result of stimulation of airway smooth-muscle cells by various growth factors such as platelet-derived growth factor (PDGF) or endothelin-1 released from inflammatory or epithelial cells.

VASCULAR RESPONSES There is increased airway mucosal blood flow in asthma, which may contribute to airway narrowing. There is an increase in the number of blood vessels in asthmatic airways as a result of angiogenesis in response to growth factors, particularly vascular endothelial growth factor. Microvascular leakage from postcapillary venules in response to inflammatory mediators is observed in asthma, resulting in airway edema and plasma exudation into the airway lumen.

MUCUS HYPERSECRETION Increased mucus secretion contributes to the viscid mucous plugs that occlude asthmatic airways, particularly in fatal asthma. There is hyperplasia of submucosal glands that are confined to large airways and of increased numbers of epithelial goblet cells. IL-13 induces mucus hypersecretion in experimental models of asthma.

NEURAL REGULATION Various defects in autonomic neural control may contribute to AHR in asthma, but these are likely to be secondary to the disease, rather than primary defects. Cholinergic pathways, through the release of acetylcholine acting on muscarinic receptors, cause bronchoconstriction and may be activated reflexly in asthma.

Inflammatory mediators may activate sensory nerves, resulting in reflex cholinergic bronchoconstriction or release of inflammatory neuropeptides. Inflammatory products may also sensitize sensory nerve endings in the airway epithelium such that the nerves become hyperalgesic. Neurotrophins, which may be released from various cell types in airways, including epithelial cells and mast cells, may cause proliferation and sensitization of airway sensory nerves. Airway nerves may also release neurotransmitters, such as substance P, which have inflammatory effects.

Airway Remodeling Several changes in the structure of the airway are characteristically found in asthma, and these may lead to irreversible narrowing of the airways. Population studies have shown a greater decline in lung function over time than in normal subjects; however, most patients with asthma preserve normal or near-normal lung function throughout life if appropriately treated.

The accelerated decline in lung function occurs in a smaller proportion of asthmatics, and these are usually patients with more severe disease. There is some evidence that the early use of ICS may reduce the decline in lung function. The characteristic structural changes are increased airway smooth muscle, fibrosis, angiogenesis, and mucus hyperplasia.

Physiology Limitation of airflow is due mainly to bronchoconstriction, but airway edema, vascular congestion, and luminal occlusion with exudate may contribute. This results in a reduction in forced expiratory volume in 1 second (FEV_1), FEV_1 /forced vital capacity (FVC) ratio, and peak expiratory flow (PEF), as well as an increase in airway resistance. Early closure of peripheral airway results in lung hyperinflation (air trapping) and increased residual volume, particularly during acute exacerbations and in severe persistent asthma. In more severe asthma, reduced ventilation and increased pulmonary blood flow result in mismatching of ventilation and perfusion and in bronchial hyperemia. Ventilatory failure is very uncommon, even in patients with severe asthma, and arterial PCO_2 tends to be low due to increased ventilation.

Airway Hyperresponsiveness AHR is the characteristic physiologic abnormality of asthma and describes the excessive bronchoconstrictor response to multiple inhaled triggers that would have no effect on normal airways. The increase in AHR is linked to the frequency of asthma symptoms, and, thus, an important aim of therapy is to reduce AHR. Increased bronchoconstrictor responsiveness is seen with *direct* bronchoconstrictors such as histamine and methacholine, which contract airway smooth muscle, but is characteristically also seen with many *indirect* stimuli, which release bronchoconstrictors from mast cells or activate sensory nerves. Most of the triggers for asthma symptoms appear to act indirectly, including allergens, exercise, hyperventilation, fog (via mast cell activation), irritant dusts, and sulfur dioxide (via a cholinergic reflex).

CLINICAL FEATURES AND DIAGNOSIS

The characteristic symptoms of asthma are wheezing, dyspnea, and coughing, which are variable, both spontaneously and with therapy. Symptoms may be worse at night, and patients typically awake in the early morning hours. Patients may report difficulty in filling their lungs with air. There is increased mucus production in some patients, with typically tenacious mucus that is difficult to expectorate. There may be increased ventilation and use of accessory muscles of ventilation. Prodromal symptoms may precede an attack, with itching under the chin, discomfort between the scapulae, or inexplicable fear (impending doom).

Typical physical signs are inspiratory, and to a greater extent expiratory, rhonchi throughout the chest, and there may be hyperinflation. Some patients, particularly children, may present with a

predominant nonproductive cough (cough-variant asthma). There may be no abnormal physical findings when asthma is under control.

DIAGNOSIS

The diagnosis of asthma is usually apparent from the symptoms of variable and intermittent airways obstruction, but must be confirmed by objective measurements of lung function.

Lung Function Tests Simple spirometry confirms airflow limitation with a reduced FEV_1 , FEV_1 /FVC ratio, and PEF (Fig. 309-6). Reversibility is demonstrated by a $>12\%$ and 200-mL increase in FEV_1 15 min after an inhaled short-acting β_2 -agonist or in some patients by a 2- to 4-week trial of oral corticosteroids (OCS) (prednisone or prednisolone 30–40 mg daily). Measurements of PEF twice daily may confirm the diurnal variations in airflow obstruction. Flow-volume loops show reduced peak flow and reduced maximum expiratory flow. Further lung function tests are rarely necessary, but whole-body plethysmography shows increased airway resistance and may show increased total lung capacity and residual volume. Gas diffusion is usually normal, but there may be a small increase in gas transfer in some patients.

Airway Responsiveness The increased AHR is normally measured by methacholine or histamine challenge with calculation of the provocative concentration that reduces FEV_1 by 20% (PC_{20}). This is rarely useful in clinical practice, but can be used in the differential diagnosis of chronic cough and when the diagnosis is in doubt in the setting of normal pulmonary function tests. Occasionally exercise testing is done to demonstrate the postexercise bronchoconstriction if there is a predominant history of EIA. Allergen challenge is rarely necessary and should only be undertaken by a specialist if specific occupational agents are to be identified.

Hematologic Tests Blood tests are not usually helpful. Total serum IgE and specific IgE to inhaled allergens (radioallergosorbent test [RAST]) may be measured in some patients.

Imaging Chest roentgenography is usually normal but in more severe patients may show hyperinflated lungs. In exacerbations, there may be evidence of a pneumothorax. Lung shadowing usually indicates pneumonia or eosinophilic infiltrates in patients with bronchopulmonary aspergillosis. High-resolution computed tomography (CT) may show areas of bronchiectasis in patients with severe asthma, and there may be thickening of the bronchial walls, but these changes are not diagnostic of asthma.

Skin Tests Skin prick tests to common inhalant allergens (house dust mite, cat fur, grass pollen) are positive in allergic asthma and negative

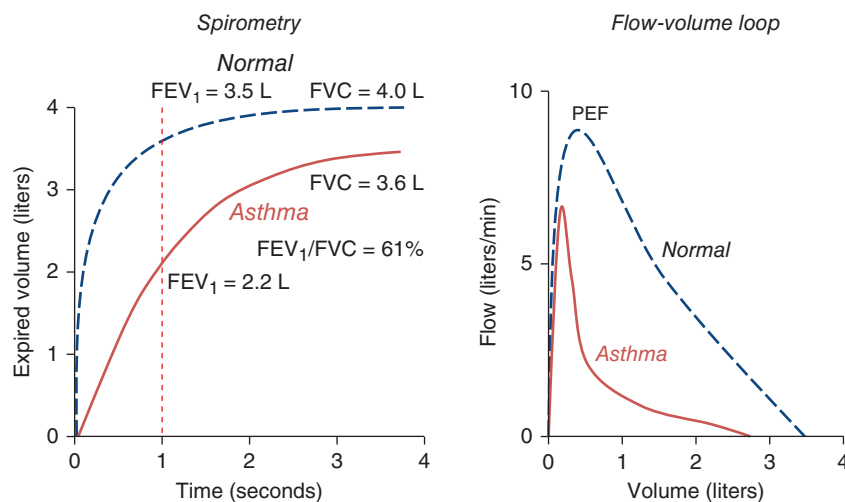


FIGURE 309-6 Spirometry and flow-volume loop in asthmatic compared to normal subject. There is a reduction in forced expiratory volume in 1 second (FEV_1) but less reduction in forced vital capacity (FVC), giving a reduced FEV_1 /FVC ratio ($<70\%$). The flow-volume loop shows reduced peak expiratory flow and a typical scalloped appearance indicating widespread airflow obstruction.

1676 in intrinsic asthma, but are not helpful in diagnosis. Positive skin responses may be useful in persuading patients to undertake allergen avoidance measures.

Exhaled Nitric Oxide $F_E NO$ is now being used as a noninvasive test to measure airway inflammation. The typically elevated levels in asthma are reduced by ICS, so this may be a test of compliance with therapy. It may also be useful in demonstrating insufficient anti-inflammatory therapy and may be useful in down-titrating ICS. However, studies in unselected patients have not convincingly demonstrated improved clinical outcomes, and it may be necessary to select patients who are poorly controlled.

Differential Diagnosis It is usually not difficult to differentiate asthma from other conditions that cause wheezing and dyspnea. Upper airway obstruction by a tumor or laryngeal edema can mimic severe asthma, but patients typically present with stridor localized to large airways. The diagnosis is confirmed by a flow-volume loop that shows a reduction in inspiratory as well as expiratory flow, and bronchoscopy to demonstrate the site of upper airway narrowing. Persistent wheezing in a specific area of the chest may indicate endobronchial obstruction with a foreign body. Left ventricular failure may mimic the wheezing of asthma, but basilar crackles are present in contrast to asthma. Vocal chord dysfunction may mimic asthma and is thought to be an hysterical conversion syndrome.

Eosinophilic pneumonias and systemic vasculitis, including Churg-Strauss syndrome and polyarteritis nodosa, may be associated with wheezing. Chronic obstructive pulmonary disease (COPD) is usually easy to differentiate from asthma as symptoms show less variability, never completely remit, and show much less (or no) reversibility to bronchodilators. Approximately 10% of COPD patients have features of asthma, with increased sputum eosinophils and a response to OCSs; these patients probably have both diseases concomitantly.

TREATMENT ASTHMA

The treatment of asthma is straightforward, and the majority of patients are now managed by internists and family doctors with effective and safe therapies. There are several aims of therapy (Table 309-2). Most emphasis has been placed on drug therapy, but several nonpharmacologic approaches have also been used. The main drugs for asthma can be divided into bronchodilators, which give rapid relief of symptoms mainly through relaxation of airway smooth muscle, and controllers, which inhibit the underlying inflammatory process.

BRONCHODILATOR THERAPIES

Bronchodilators act primarily on airway smooth muscle to reverse the bronchoconstriction of asthma. This gives rapid relief of symptoms but has little or no effect on the underlying inflammatory process. Thus, bronchodilators are not sufficient to control asthma in patients with persistent symptoms. There are three classes of bronchodilators in current use: β_2 -adrenergic agonists, anticholinergics, and theophylline; of these, β_2 -agonists are by far the most effective.

β_2 -Agonists β_2 -Agonists activate β_2 -adrenergic receptors, which are widely expressed in the airways. β_2 -Receptors are coupled through a stimulatory G protein to adenylyl cyclase, resulting in increased

intracellular cyclic adenosine monophosphate (AMP), which relaxes smooth-muscle cells and inhibits certain inflammatory cells, particularly mast cells.

MODE OF ACTION The primary action of β_2 -agonists is to relax airway smooth-muscle cells of all airways, where they act as functional antagonists, reversing and preventing contraction of airway smooth-muscle cells by all known bronchoconstrictors. This generalized action is likely to account for their great efficacy as bronchodilators in asthma. There are also additional nonbronchodilator effects that may be clinically useful, including inhibition of mast cell mediator release, reduction in plasma exudation, and inhibition of sensory nerve activation. Inflammatory cells express small numbers of β_2 -receptors, but these are rapidly downregulated with β_2 -agonist activation so that, in contrast to corticosteroids, there are no effects on inflammatory cells in the airways and there is no reduction in AHR.

CLINICAL USE β_2 -Agonists are usually given by inhalation to reduce side effects. Short-acting β_2 -agonists (SABAs) such as albuterol and terbutaline have a duration of action of 3–6 h. They have a rapid onset of bronchodilatation and are, therefore, used as needed for symptom relief. Increased use of SABA indicates that asthma is not controlled. They are also useful in preventing EIA if taken prior to exercise. SABAs are used in high doses by nebulizer or via a metered-dose inhaler with a spacer. Long-acting β_2 -agonists (LABAs) include salmeterol and formoterol, both of which have a duration of action over 12 h and are given twice daily by inhalation; indacaterol is given once daily. LABAs have replaced the regular use of SABAs, but LABAs should not be given in the absence of ICS therapy because they do not control the underlying inflammation. They do, however, improve asthma control and reduce exacerbations when added to ICS, which allows asthma to be controlled at lower doses of corticosteroids. This observation has led to the widespread use of fixed-combination inhalers that contain a corticosteroid and a LABA, which have proved to be highly effective in the control of asthma.

SIDE EFFECTS Adverse effects are not usually a problem with β_2 -agonists when given by inhalation. The most common side effects are muscle tremor and palpitations, which are seen more commonly in elderly patients. There is a small fall in plasma potassium due to increased uptake by skeletal muscle cells, but this effect does not usually cause any clinical problem.

TOLERANCE Tolerance is a potential problem with any agonist given chronically, but although there is downregulation of β_2 -receptors, this does not reduce the bronchodilator response because there is a large receptor reserve in airway smooth-muscle cells. By contrast, mast cells become rapidly tolerant, but their tolerance may be prevented by concomitant administration of ICS.

SAFETY The safety of β_2 -agonists has been an important issue. There is an association between asthma mortality and the amount of SABA used, but careful analysis demonstrates that the increased use of rescue SABA reflects poor asthma control, which is a risk factor for asthma death. The slight excess in mortality that has been associated with the use of LABA is related to the lack of use of concomitant ICS, as the LABA therapy fails to suppress the underlying inflammation. This highlights the importance of always using an ICS when LABAs are given, which is most conveniently achieved by using a combination inhaler.

Anticholinergics Muscarinic receptor antagonists such as ipratropium bromide prevent cholinergic nerve-induced bronchoconstriction and mucus secretion. They are less effective than β_2 -agonists in asthma therapy because they inhibit only the cholinergic reflex component of bronchoconstriction, whereas β_2 -agonists prevent all bronchoconstrictor mechanisms. Anticholinergics, including once-daily tiotropium bromide, may be used as an additional bronchodilator in patients with asthma that is not controlled by ICS and LABA combinations. High doses may be given by nebulizer in treating acute severe asthma but should only be given following β_2 -agonists, because they have a slower onset of bronchodilatation.

TABLE 309-2 AIMS OF ASTHMA THERAPY

- Minimal (ideally no) chronic symptoms, including nocturnal
- Minimal (infrequent) exacerbations
- No emergency visits
- Minimal (ideally no) use of a required β_2 -agonist
- No limitations on activities, including exercise
- Peak expiratory flow circadian variation <20%
- (Near) normal peak expiratory flow
- Minimal (or no) adverse effects from medicine

Side effects are not usually a problem because there is little or no systemic absorption. The most common side effect is dry mouth; in elderly patients, urinary retention and glaucoma may also be observed.

Theophylline Theophylline was widely prescribed as an oral bronchodilator several years ago, especially because it was inexpensive. It has now fallen out of favor because side effects are common and inhaled β_2 -agonists are much more effective as bronchodilators. The bronchodilator effect is due to inhibition of phosphodiesterases in airway smooth-muscle cells, which increases cyclic AMP, but doses required for bronchodilatation commonly cause side effects that are mediated mainly by phosphodiesterase inhibition. There is increasing evidence that theophylline at lower doses has anti-inflammatory effects, and these are likely to be mediated through different molecular mechanisms. Theophylline activates the key nuclear enzyme histone deacetylase-2 (HDAC2), which is a critical mechanism for switching off activated inflammatory genes and may, therefore, reduce corticosteroid insensitivity in severe asthma.

CLINICAL USE Oral theophylline is usually given as a slow-release preparation once or twice daily because this gives more stable plasma concentrations than normal theophylline tablets. It may be used as an additional bronchodilator in patients with severe asthma when plasma concentrations of 10–20 mg/L are required, although these concentrations are often associated with side effects. Low doses of theophylline, giving plasma concentrations of 5–10 mg/L, have additive effects to ICS and are particularly useful in patients with severe asthma. Indeed, withdrawal of theophylline from these patients may result in marked deterioration in asthma control. At low doses, the drug is well tolerated. IV aminophylline (a soluble salt of theophylline) was used for the treatment of severe asthma but has now been largely replaced by high doses of inhaled SABA, which are more effective and have fewer side effects. Aminophylline is occasionally used (via slow IV infusion) in patients with severe exacerbations that are refractory to SABA.

SIDE EFFECTS Oral theophylline is well absorbed and is largely inactivated in the liver. Side effects are related to plasma concentrations; measurement of plasma theophylline may be useful in determining the correct dose. The most common side effects are nausea, vomiting, and headaches and are due to phosphodiesterase inhibition. Diuresis and palpitations may also occur, and at high concentrations, cardiac arrhythmias, epileptic seizures, and death may occur due to adenosine A_1 -receptor antagonism. Theophylline side effects are related to plasma concentration and are rarely observed at plasma concentrations below 10 mg/L. Theophylline is metabolized by CYP450 in the liver, and thus, plasma concentrations may be elevated by drugs that block CYP450 such as erythromycin and allopurinol. Other drugs may also reduce clearance by other mechanisms leading to increased plasma concentrations (Table 309-3).

TABLE 309-3 FACTORS AFFECTING CLEARANCE OF THEOPHYLLINE

Increased Clearance

- Enzyme induction (rifampicin, phenobarbitone, ethanol)
- Smoking (tobacco, marijuana)
- High-protein, low-carbohydrate diet
- Barbecued meat
- Childhood

Decreased Clearance

- Enzyme inhibition (cimetidine, erythromycin, ciprofloxacin, allopurinol, zileuton, zafirlukast)
- Congestive heart failure
- Liver disease
- Pneumonia
- Viral infection and vaccination
- High-carbohydrate diet
- Old age

CONTROLLER THERAPIES

Inhaled Corticosteroids ICSs are by far the most effective controllers for asthma, and their early use has revolutionized asthma therapy.

MODE OF ACTION ICSs are the most effective anti-inflammatory agents used in asthma therapy, reducing inflammatory cell numbers and their activation in the airways. ICSs reduce eosinophils in the airways and sputum and the numbers of activated T lymphocytes and surface mast cells in the airway mucosa. These effects may account for the reduction in AHR that is seen with chronic ICS therapy.

The molecular mechanism of action of corticosteroids involves several effects on the inflammatory process. The major effect of corticosteroids is to switch off the transcription of multiple activated genes that encode inflammatory proteins such as cytokines, chemokines, adhesion molecules, and inflammatory enzymes. This effect involves several mechanisms, including inhibition of the transcription factor NF- κ B, but an important mechanism is recruitment of HDAC2 to the inflammatory gene complex, which reverses the histone acetylation associated with increased gene transcription. Corticosteroids also activate anti-inflammatory genes, such as mitogen-activated protein (MAP) kinase phosphatase-1, and increase the expression of β_2 -receptors. Most of the metabolic and endocrine side effects of corticosteroids are also mediated through transcriptional activation.

CLINICAL USE ICSs are by far the most effective controllers in the management of asthma and are beneficial in treating asthma of any severity and age. ICSs are usually given twice daily, but some may be effective once daily in mildly symptomatic patients. ICSs rapidly improve the symptoms of asthma, and lung function improves over several days. They are effective in preventing asthma symptoms, such as EIA and nocturnal exacerbations, but also prevent severe exacerbations. ICSs reduce AHR, but maximal improvement may take several months of therapy. Early treatment with ICS appears to prevent irreversible changes in airway function that occur with chronic asthma. Withdrawal of ICS results in slow deterioration of asthma control, indicating that they suppress inflammation and symptoms, but do not cure the underlying condition. ICSs are now given as first-line therapy for patients with persistent asthma, but if they do not control symptoms at low doses, it is usual to add a LABA as the next step.

SIDE EFFECTS Local side effects include hoarseness (dysphonia) and oral candidiasis, which may be reduced with the use of a large-volume spacer device. There has been concern about systemic side effects from lung absorption, but many studies have demonstrated that ICS have minimal systemic effects (Fig. 309-7). At the highest recommended doses, there may be some suppression of plasma and urinary cortisol concentrations, but there is no convincing evidence that long-term treatment leads to impaired growth in children or to osteoporosis in adults. Indeed effective control of asthma with ICS reduces the number of courses of OCS that are needed and, thus, reduces systemic exposure to ICS.

Systemic Corticosteroids Corticosteroids are used intravenously (hydrocortisone or methylprednisolone) for the treatment of acute severe asthma, although several studies now show that OCSs are as effective and easier to administer. A course of OCS (usually prednisone or prednisolone 30–45 mg once daily for 5–10 days) is used to treat acute exacerbations of asthma; no tapering of the dose is needed. Approximately 1% of asthma patients may require maintenance treatment with OCS; the lowest dose necessary to maintain control needs to be determined. Systemic side effects, including truncal obesity, bruising, osteoporosis, diabetes, hypertension, gastric ulceration, proximal myopathy, depression, and cataracts, may be a major problem, and steroid-sparing therapies may be considered if side effects are a significant problem. If patients require maintenance treatment with OCS, it is important to monitor bone density so that preventive treatment with bisphosphonates or estrogen in postmenopausal women may be initiated if bone density is low.

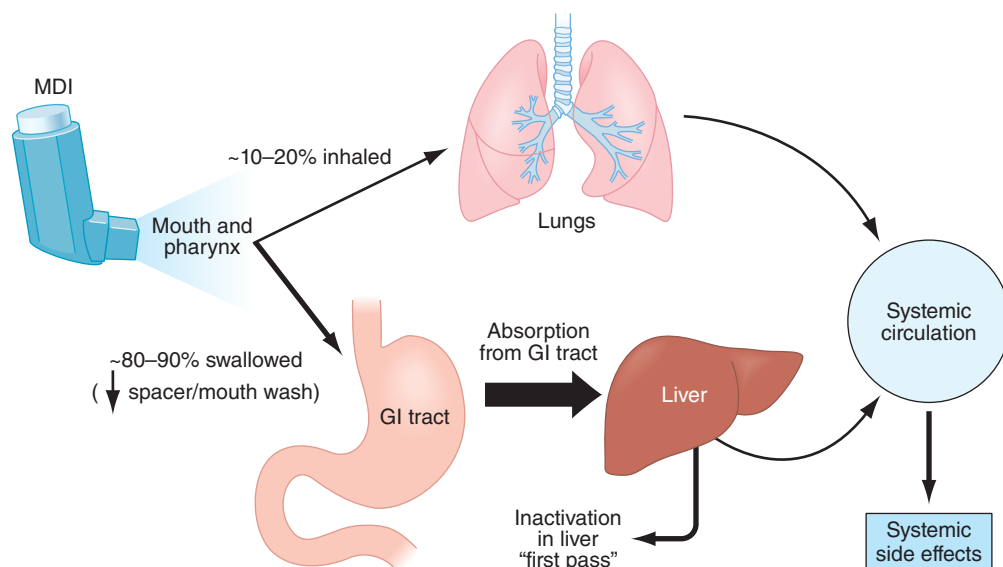


FIGURE 309-7 Pharmacokinetics of inhaled corticosteroids. GI, gastrointestinal; MDI, metered-dose inhaler.

Intramuscular triamcinolone acetonide is a depot preparation that is occasionally used in noncompliant patients, but proximal myopathy is a major problem with this therapy.

Antileukotrienes Cysteinyl-leukotrienes are potent bronchoconstrictors, cause microvascular leakage, and increase eosinophilic inflammation through the activation of cys-LT_1 -receptors. These inflammatory mediators are produced predominantly by mast cells and, to a lesser extent, eosinophils in asthma. Antileukotrienes, such as montelukast, block cys-LT_1 -receptors and provide modest clinical benefit in asthma. They are less effective than ICS in controlling asthma and have less effect on airway inflammation, but are useful as an add-on therapy in some patients not controlled with low doses of ICS, although less effective than LABA. They are given orally once or twice daily and are well tolerated. Some patients show a better response than others to antileukotrienes, but this has not been convincingly linked to any genomic differences in the leukotriene pathway.

Cromones Cromolyn sodium and nedocromil sodium are asthma controller drugs that appear to inhibit mast cell and sensory nerve activation and are, therefore, effective in blocking trigger-induced asthma such as EIA and allergen- and sulfur dioxide-induced symptoms. Cromones have relatively little benefit in the long-term control of asthma due to their short duration of action (at least four times daily by inhalation). They are very safe and were popular in the treatment of childhood asthma, although now low doses of ICS are preferred because they are more effective and have a proven safety profile.

Steroid-Sparing Therapies Various immunomodulatory treatments have been used to reduce the requirement for OCS in patients with severe asthma who have serious side effects with this therapy. Methotrexate, cyclosporin A, azathioprine, gold, and IV gamma globulin have all been used as steroid-sparing therapies, but none of these treatments has any long-term benefit, and each is associated with a relatively high risk of side effects.

Anti-IgE Omalizumab is a blocking antibody that neutralizes circulating IgE without binding to cell-bound IgE and, thus, inhibits IgE-mediated reactions. This treatment has been shown to reduce the number of exacerbations in patients with severe asthma and may improve asthma control. However, the treatment is very expensive and is only suitable for highly selected patients who are not controlled on maximal doses of inhaler therapy and have a circulating IgE within a specified range. Patients should be given a 3- to 4-month trial of therapy to show objective benefit. Omalizumab is usually given as a subcutaneous injection every 2–4 weeks and

appears not to have significant side effects, although anaphylaxis is very occasionally seen.

Immunotherapy Specific immunotherapy using injected extracts of pollens or house dust mites has not been very effective in controlling asthma and may cause anaphylaxis. Side effects may be reduced by sublingual dosing. It is not recommended in most asthma treatment guidelines because of lack of evidence of clinical efficacy.

Alternative Therapies Nonpharmacologic treatments, including hypnosis, acupuncture, chiropraxis, breathing control, yoga, and speleotherapy, may be popular with some patients. However, placebo-controlled studies have shown that each of these treatments lacks efficacy and cannot be recommended. However, they are not detrimental and may be used as long as conventional pharmacologic therapy is continued.

Future Therapies It has proved very difficult to discover novel pharmaceutical therapies, particularly because current therapy with corticosteroids and β_2 -agonists is so effective in the majority of patients. There is, however, a need for the development of new therapies for patients with refractory asthma who have side effects with systemic corticosteroids. Antagonists of specific mediators have little or no benefit in asthma, apart from antileukotrienes, which have rather weak effects, presumably reflecting the fact that multiple mediators are involved. Blocking antibodies against IL-5 may reduce exacerbations in highly selected patients who have sputum eosinophils despite high doses of corticosteroids, whereas anti-TNF- α antibodies are not effective in severe asthma. Novel anti-inflammatory treatments that are in clinical development include inhibitors of phosphodiesterase-4, NF- κ B, and p38 MAP kinase. However, these drugs, which act on signal transduction pathways common to many cells, are likely to have troublesome side effects, necessitating their delivery by inhalation. Safer and more effective immunotherapy using T cell peptide fragments of allergens or DNA vaccination is also being investigated. Bacterial products, such as CpG oligonucleotides that stimulate T_H1 immunity or regulatory T cells, are also currently under evaluation.

MANAGEMENT OF CHRONIC ASTHMA

There are several aims of chronic therapy in asthma (Table 309-2). It is important to establish the diagnosis objectively using spirometry or PEF measurements at home. Triggers that worsen asthma control, such as allergens or occupational agents, should be avoided, whereas triggers, such as exercise and fog, which result in transient symptoms, provide an indication that more controller therapy is needed. It is important to assess asthma control, determined by

TABLE 309-4 ASTHMA CONTROL

Characteristic	Controlled (all of the following)	Partly Controlled	Uncontrolled
Daytime symptoms	None (≤ 2 /week)	> 2 /week	Three or more features of partly controlled
Limitation of activities	None	Any	
Nocturnal symptoms/awakening	None	Any	
Need for reliever/rescue treatment	None (≤ 2 /week)	> 2 /week	
Lung function (PEF or FEV ₁)	Normal	$< 80\%$ predicted or personal best (if known)	

Abbreviations: FEV₁, forced expiratory volume in 1 s; PEF, peak expiratory flow.

symptoms, night awakening, need for reliever inhalers, limitation of activity, and lung function (Table 309-4). Avoidance of side effects and expense of medications are also important. There are several validated questionnaires for quantifying asthma control, such as the Asthma Quality of Life Questionnaire (AQLQ) and Asthma Control Test (ACT).

Stepwise Therapy For patients with mild, intermittent asthma, a short-acting β_2 -agonist is all that is required (Fig. 309-8). However, use of a reliever medication more than twice a week indicates the need for regular controller therapy. The treatment of choice for all patients is an ICS given twice daily. It is usual to start with an intermediate dose (e.g., 200 μg bid of beclomethasone dipropionate [BDP]) or equivalent and to decrease the dose if symptoms are controlled after 3 months. If symptoms are not controlled, a LABA should be added, which is most conveniently given by switching to a combination inhaler. The dose of controller should be adjusted accordingly, as judged by the need for a rescue inhaler. Low doses of theophylline or an antileukotriene may also be considered as an add-on therapy, but these are less effective than LABA. In patients with severe asthma, low-dose oral theophylline is also helpful, and when there is irreversible airway narrowing, the long-acting anticholinergic tiotropium bromide may be tried. If asthma is not controlled despite the maximal recommended dose of inhaled therapy, it is important to check compliance and inhaler technique. In these patients, maintenance treatment with an OCS may be needed, and the lowest dose that maintains control should be used. Occasionally omalizumab may be tried in steroid-dependent asthmatics who are not well controlled. Once asthma is controlled, it is important to slowly decrease therapy in order to find the optimal dose to control symptoms.

Education Patients with asthma need to understand how to use their medications and the difference between reliever and controller

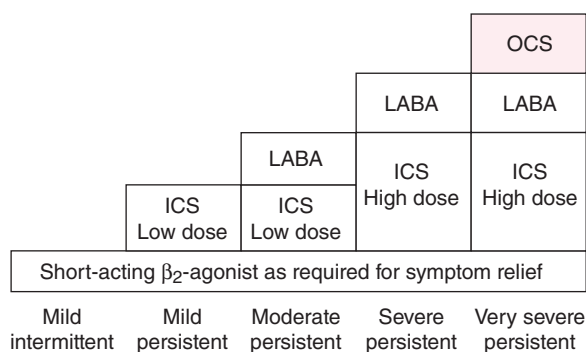


FIGURE 309-8 Stepwise approach to asthma therapy according to the severity of asthma and ability to control symptoms. ICS, inhaled corticosteroids; LABA, long-acting β_2 -agonist; OCS, oral corticosteroid.

therapies. Education may improve compliance, particularly with ICS. All patients should be taught how to use their inhalers correctly. In particular, they need to understand how to recognize worsening of asthma and how to step up therapy. Written action plans have been shown to reduce hospital admissions and morbidity rates in adults and children, and are recommended particularly in patients with unstable disease who have frequent exacerbations.

ACUTE SEVERE ASTHMA

Exacerbations of asthma are feared by patients and may be life threatening. One of the main aims of controller therapy is to prevent exacerbations; in this respect, ICS and combination inhalers are very effective.

Clinical Features Patients are aware of increasing chest tightness, wheezing, and dyspnea that are often not or poorly relieved by their usual reliever inhaler. In severe exacerbations, patients may be so breathless that they are unable to complete sentences and may become cyanotic. Examination usually shows increased ventilation, hyperinflation, and tachycardia. Pulsus paradoxus may be present, but this is rarely a useful clinical sign. There is a marked fall in spirometric values and PEF. Arterial blood gases on air show hypoxemia, and PCO_2 is usually low due to hyperventilation. A normal or rising PCO_2 is an indication of impending respiratory failure and requires immediate monitoring and therapy. A chest roentgenogram is not usually informative but may show pneumonia or pneumothorax.

TREATMENT ACUTE SEVERE ASTHMA

A high concentration of oxygen should be given by face mask to achieve oxygen saturation of $> 90\%$. The mainstay of treatment are high doses of SABA given either by nebulizer or via a metered-dose inhaler with a spacer. In severely ill patients with impending respiratory failure, IV β_2 -agonists may be given. A nebulized anticholinergic may be added if there is not a satisfactory response to β_2 -agonists alone, as there are additive effects. In patients who are refractory to inhaled therapies, a slow infusion of aminophylline may be effective, but it is important to monitor blood levels, especially if patients have already been treated with oral theophylline. Magnesium sulfate given intravenously or by nebulizer is effective when added to inhaled β_2 -agonists, and is relatively well tolerated but is not routinely recommended. Prophylactic intubation may be indicated for impending respiratory failure, when the PCO_2 is normal or rises. For patients with respiratory failure, it is necessary to intubate and institute ventilation. These patients may benefit from an anesthetic such as halothane if they have not responded to conventional bronchodilators. Sedatives should never be given because they may depress ventilation. Antibiotics should not be used routinely unless there are signs of pneumonia.

SPECIAL CONSIDERATIONS

Refractory Asthma Although most patients with asthma are easily controlled with appropriate medication, a small proportion of patients (approximately 5–10% of asthmatics) are difficult to control despite maximal inhaled therapy. Some of these patients will require maintenance treatment with OCS. In managing these patients, it is important to investigate and correct any mechanisms that may be aggravating asthma. There are two major patterns of difficult asthma: some patients have persistent symptoms and poor lung function, despite appropriate therapy, whereas others may have normal or near-normal lung function but intermittent, severe (sometimes life-threatening) exacerbations.

MECHANISMS The most common reason for poor control of asthma is noncompliance with medication, particularly ICS. Compliance with ICS may be low because patients do not feel any immediate clinical benefit or may be concerned about side effects. Compliance with ICS is difficult to monitor because there are no useful plasma measurements that can be made, but measuring the fractional excretion of

1680 induced NO ($F_E NO$) may identify the problem. Compliance may be improved by giving the ICS as a combination with a LABA that gives symptom relief. Compliance with OCS may be measured by suppression of plasma cortisol and the expected concentration of prednisone/prednisolone in the plasma. There are several factors that may make asthma more difficult to control, including exposure to high, ambient levels of allergens or unidentified occupational agents. Severe rhinosinusitis may make asthma more difficult to control; upper airway disease should be vigorously treated. Drugs such as beta-adrenergic blockers, aspirin, and other cyclooxygenase (COX) inhibitors may worsen asthma. Some women develop severe premenstrual worsening of asthma, which is unresponsive to corticosteroids and requires treatment with progesterone or gonadotropin-releasing factors. Few systemic diseases make asthma more difficult to control, but hyper- and hypothyroidism may increase asthma symptoms and should be investigated if suspected.

Bronchial biopsy studies in refractory asthma may show the typical eosinophilic pattern of inflammation, whereas others have a predominantly neutrophilic pattern. There may be an increase in T_H1 cells, T_H17 cells, and CD8 lymphocytes compared to mild asthma and increased expression of TNF- α . Structural changes in the airway, including fibrosis, angiogenesis, and airway smooth-muscle thickening, are more commonly seen in these patients.

Corticosteroid-Resistant Asthma A few patients with asthma show a poor response to corticosteroid therapy and may have various molecular abnormalities that impair the anti-inflammatory action of corticosteroids. Complete resistance to corticosteroids is extremely uncommon and affects less than 1 in 1000 patients. It is defined by a failure to respond to a high dose of oral prednisone/prednisolone (40 mg once daily over 2 weeks), ideally with a 2-week run-in with matched placebo. More common is reduced responsiveness to corticosteroids where control of asthma requires OCS (corticosteroid-dependent asthma). In patients with poor responsiveness to corticosteroids, there is a reduction in the response of circulating monocytes and lymphocytes to the anti-inflammatory effects of corticosteroids *in vitro* and reduced skin blanching in response to topical corticosteroids. There are several mechanisms that have been described, including an increase in the alternatively spliced form of the glucocorticoid receptor (GR)- β , an abnormal pattern of histone acetylation in response to corticosteroids, a defect in IL-10 production, and a reduction in HDAC2 activity (as in COPD). These observations suggest that there are likely to be heterogeneous mechanisms for corticosteroid resistance; whether these mechanisms are genetically determined has yet to be decided.

Brittle Asthma Some patients show chaotic variations in lung function despite taking appropriate therapy. Some show a persistent pattern of variability and may require oral corticosteroids or, at times, continuous infusion of β_2 -agonists (type 1 brittle asthma), whereas others have generally normal or near-normal lung function but precipitous, unpredictable falls in lung function that may result in death (type 2 brittle asthma). These latter patients are difficult to manage because they do not respond well to corticosteroids, and the worsening of asthma does not reverse well with inhaled bronchodilators. The most effective therapy is subcutaneous epinephrine, which suggests that the worsening is likely to be a localized airway anaphylactic reaction with edema. In some of these patients, there may be allergy to specific foods. These patients should be taught to self-administer epinephrine and should carry a medical warning accordingly.

TREATMENT REFRACTORY ASTHMA

Refractory asthma is difficult to control, by definition. It is important to check compliance and the correct use of inhalers and to identify and eliminate any underlying triggers. Low doses of theophylline may be helpful in some patients, and theophylline withdrawal has been found to worsen in many patients. Most of these patients will require maintenance treatment with oral corticosteroids, and the minimal dose that achieves satisfactory control should be determined by careful dose titration. Steroid-sparing therapies are rarely

effective. In some patients with allergic asthma, omalizumab is effective, particularly when there are frequent exacerbations. Anti-TNF therapy is not effective in severe asthma and should not be used. A few patients may benefit from infusions of β_2 -agonists. New therapies are needed for these patients, who currently consume a disproportionate amount of health care spending.

Aspirin-Sensitive Asthma A small proportion (1–5%) of asthmatics become worse with aspirin and other COX inhibitors, although this is much more commonly seen in severe cases and in patients with frequent hospital admission. Aspirin-sensitive asthma is a well-defined phenotype of asthma that is usually preceded by perennial rhinitis and nasal polyps in nonatopic patients with a late onset of the disease. Aspirin, even in small doses, characteristically provokes rhinorrhea, conjunctival injection, facial flushing, and wheezing. There is a genetic predisposition to increased production of cysteinyl-leukotrienes with functional polymorphism of *cys*-leukotriene C_4 synthase. Asthma is triggered by COX inhibitors but is persistent even in their absence. All nonselective COX inhibitors should be avoided, but selective COX2 inhibitors are safe to use when an anti-inflammatory analgesic is needed. Aspirin-sensitive asthma responds to usual therapy with ICS. Although antileukotrienes should be effective in these patients, they are no more effective than in allergic asthma. Occasionally, aspirin desensitization is necessary, but this should only be undertaken in specialized centers.

Asthma in the Elderly Asthma may start at any age, including in elderly patients. The principles of management are the same as in other asthmatics, but side effects of therapy may be a problem, including muscle tremor with β_2 -agonists and more systemic side effects with ICS. Comorbidities are more frequent in this age group, and interactions with drugs such as β_2 -blockers, COX inhibitors, and agents that may affect theophylline metabolism need to be considered. COPD is more likely in elderly patients and may coexist with asthma. A trial of OCS may be very useful in documenting the steroid responsiveness of asthma.

Pregnancy Approximately one-third of asthmatic patients who are pregnant improve during the course of a pregnancy, one-third deteriorate, and one-third are unchanged. It is important to maintain good control of asthma because poor control may have adverse effects on fetal development. Compliance may be a problem because there is often concern about the effects of antiasthma medications on fetal development. The drugs that have been used for many years in asthma therapy have now been shown to be safe and without teratogenic potential. These drugs include SABA, ICS, and theophylline; there is less safety information about newer classes of drugs such as LABA, antileukotrienes, and anti-IgE. If an OCS is needed, it is better to use prednisone rather than prednisolone because it cannot be converted to the active prednisolone by the fetal liver, thus protecting the fetus from systemic effects of the corticosteroid. There is no contraindication to breast-feeding when patients are using these drugs.

Cigarette Smoking Approximately 20% of asthmatics smoke, which may adversely affect asthma in several ways. Smoking asthmatics have more severe disease, more frequent hospital admissions, a faster decline in lung function, and a higher risk of death from asthma than nonsmoking asthmatics. There is evidence that smoking interferes with the anti-inflammatory actions of corticosteroids by reducing HDAC2, necessitating higher doses for asthma control. Smoking cessation improves lung function and reduces the steroid resistance, and thus, vigorous smoking cessation strategies should be used. Some patients report a temporary worsening of asthma when they first stop smoking, possibly due to the loss of the bronchodilating effect of NO in cigarette smoke.

Surgery If asthma is well controlled, there is no contraindication to general anesthesia and intubation. Patients who are treated with OCS will have adrenal suppression and should be treated with an increased dose of OCS immediately prior to surgery. Patients with $FEV_1 < 80\%$

of their normal levels should also be given a boost of OCS prior to surgery. High-maintenance doses of corticosteroids may be a contraindication to surgery because of increased risks of infection and delayed wound healing.

Bronchopulmonary Aspergillosis Bronchopulmonary aspergillosis (BPA) is uncommon and results from an allergic pulmonary reaction to inhaled spores of *Aspergillus fumigatus* and, occasionally, other *Aspergillus* species. A skin prick test to *A. fumigatus* is always positive, whereas serum *Aspergillus* precipitins are low or undetectable. Characteristically, there are fleeting eosinophilic infiltrates in the lungs, particularly in the upper lobes. Airways become blocked with mucoid plugs rich in eosinophils, and patients may cough up brown plugs and have hemoptysis. BPA may result in bronchiectasis, particularly affecting central airways, if not suppressed by corticosteroids. Asthma is controlled in the usual way by ICS, but it is necessary to give a course of OCS if any sign of worsening or pulmonary shadowing is found. Treatment with the oral antifungal itraconazole is beneficial in preventing exacerbations.

310 Hypersensitivity Pneumonitis and Pulmonary Infiltrates with Eosinophilia

Praveen Akuthota, Michael E. Wechsler

HYPERSENSITIVITY PNEUMONITIS

INTRODUCTION AND DEFINITION

Hypersensitivity pneumonitis (HP), also referred to as extrinsic allergic alveolitis, is a pulmonary disease that occurs due to inhalational exposure to a variety of antigens leading to an inflammatory response of the alveoli and small airways. Systemic manifestations such as fever and fatigue can accompany respiratory symptoms. Although sensitization to an inhaled antigen as manifested by specific circulating IgG antibodies is necessary for the development of HP, sensitization alone is not sufficient as a defining characteristic, because many sensitized individuals do not develop HP. The incidence and prevalence of HP are variable, depending on geography, occupation, avocation, and environment of the cohort being studied. As yet unexplained is the decreased risk of developing HP in smokers.

OFFENDING ANTIGENS

HP can be caused by any of a large list of potential offending inhaled antigens (Table 310-1). The various antigens and environmental conditions described to be associated with HP give rise to an expansive list of monikers given to specific forms of HP. Antigens derived from fungal, bacterial, mycobacterial, bird-derived, and chemical sources have all been implicated in causing HP.

Categories of individuals at particular risk in the United States include farmers, bird owners, industrial workers, and hot tub users. Farmer's lung occurs as a result of exposure to one of several possible sources of bacterial or fungal antigens such as grain, moldy hay, or silage. Potential offending antigens include thermophilic actinomycetes or *Aspergillus* species. Bird fancier's lung (also referred to by names corresponding to specific birds) must be considered in patients who give a history of keeping birds in their home and is precipitated by exposure to antigens derived from feathers, droppings, and serum proteins. Occupational exposure to birds may also cause HP, as is seen in poultry worker's lung. Chemical worker's lung is provoked by exposure to occupational chemical antigens such as diphenylmethane diisocyanate and toluene diisocyanate. Mycobacteria may cause HP rather than frank infection, a phenomenon observed in hot tub lung and in HP due to metalworking fluid.

TABLE 310-1 EXAMPLES OF HYPERSENSITIVITY PNEUMONITIS

Disease	Antigen	Source
Farming/Food Processing		
Farmer's lung	Thermophilic actinomycetes (e.g., <i>Saccharopolyspora rectivirgula</i>); fungus	Grain, moldy hay, silage
Bagassosis	Thermophilic actinomycetes	Sugarcane
Cheese washer's lung	<i>Penicillium casei</i> ; <i>Aspergillus clavatus</i>	Cheese
Coffee worker's lung	Coffee bean dust	Coffee beans
Malt worker's lung	<i>Aspergillus</i> species	Barley
Miller's lung	<i>Sitophilus granarius</i> (wheat weevil)	Wheat flour
Mushroom worker's lung	Thermophilic actinomycetes; mushroom spores	Mushrooms
Potato riddler's lung	Thermophilic actinomycetes; <i>Aspergillus</i> species	Moldy hay around potatoes
Tobacco grower's lung	<i>Aspergillus</i> species	Tobacco
Wine maker's lung	<i>Botrytis cinerea</i>	Grapes
Birds and Other Animals		
Bird fancier's lung (also named by specific bird exposures)	Proteins derived by parakeets, pigeons, budgerigars	Bird feathers, droppings, serum proteins
Duck fever	Duck feathers, serum proteins	Ducks
Fish meal worker's lung	Fish meal dust	Fish meal
Furrier's lung	Dust from animal furs	Animal furs
Laboratory worker's lung	Rat urine, serum, fur	Laboratory rats
Pituitary snuff taker's lung	Animal proteins	Pituitary snuff from bovine and porcine sources
Poultry worker's lung	Chicken serum proteins	Chickens
Turkey handling disease	Turkey serum proteins	Turkeys
Other Occupational and Environmental Exposures		
Chemical worker's lung	Isocyanates	Polyurethane foam, varnish, lacquer
Detergent worker's lung	<i>Bacillus subtilis</i> enzymes	Detergent
Hot tub lung	<i>Cladosporium</i> species; <i>Mycobacterium avium</i> complex	Contaminated water, mold on ceiling
Humidifier fever (and air conditioner lung)	Several microorganisms including: <i>Aureobasidium pullulans</i> ; <i>Candida albicans</i> ; thermophilic actinomycetes; <i>Mycobacterium</i> species; <i>Klebsiella oxytoca</i> ; <i>Naegleria gruberi</i>	Humidifiers and air conditioners (contaminated water)
Machine operator's lung	<i>Pseudomonas</i> species; <i>Mycobacteria</i> species	Metal working fluid
Sauna taker's lung	<i>Aureobasidium</i> species; other antigens	Sauna water
Suberosis	<i>Penicillium glabrum</i> ; <i>Chrysonilia sitophila</i>	Cork dust
Summer-type pneumonitis	<i>Trichosporon cutaneum</i>	House dust mites, bird droppings
Woodworker's lung	<i>Alternaria</i> species; <i>Bacillus subtilis</i>	Oak, cedar, pine, mahogany dusts

The pathophysiology of HP has not been characterized in depth on an immunologic level, although it has been established that HP is an immune-mediated condition that occurs in response to inhaled antigens that are small enough to deposit in distal airways and alveoli. From a lymphocyte perspective, HP has been categorized as a condition with a T_H1 inflammatory pattern. However, emerging evidence suggests that T_H17 lymphocyte subsets may be involved in the pathogenesis of the disease as well. Although the presence of precipitating IgG antibodies against specific antigens in HP suggests a prominent role for adaptive immunity in the pathophysiology of HP, innate immune mechanisms may also make an important contribution. This is highlighted by the observation that Toll-like receptors and downstream signaling proteins such as MyD88 are activated in HP. Although no clear genetic basis for HP has been established, in specific cohorts, polymorphisms in genes involved in antigen processing and presentation, including TAP1 and major histocompatibility complex type II, have been observed.

CLINICAL PRESENTATION

Given the heterogeneity among patients, variability in offending antigens, and differences in the intensity and duration of exposure to antigen, the presentation of HP is accordingly variable. Although these categories are not fully satisfactory in capturing this variability, HP has been traditionally categorized as having *acute*, *subacute*, and *chronic* forms. Acute HP usually manifests itself 4–8 h following exposure to the inciting antigen, often intense in nature. Systemic symptoms, including fevers, chills, and malaise, are prominent and are accompanied by dyspnea. Symptoms resolve within hours to days if no further exposure to the offending antigen occurs. In subacute HP resulting from ongoing antigen exposure, the onset of respiratory and systemic symptoms is typically more gradual over the course of weeks. A similar presentation may occur as a culmination of intermittent episodes of acute HP. Although respiratory impairment may be quite severe, antigen avoidance generally results in resolution of the symptoms, although with a slower time course, on the order of weeks to months, than that seen with acute HP. Chronic HP can present with an even more gradual onset of symptoms than subacute HP, with progressive dyspnea, cough, fatigue, weight loss, and clubbing of the digits. The insidious onset of symptoms and frequent lack of an antecedent episode of acute HP make diagnosing chronic HP a challenge. Unlike with the other forms of HP, there can be an irreversible component to the respiratory impairment that is not responsive to removal of the responsible antigen from the patient's environment. The disease progression to hypoxemic respiratory failure can mirror that seen in idiopathic pulmonary fibrosis (IPF). Fibrotic lung disease is a potential feature of chronic HP due to exposure to bird antigens, whereas an emphysematous phenotype may be seen in farmer's lung.

The categories of acute, subacute, and chronic HP are not completely sufficient in classifying HP. The HP Study Group found on cluster analysis that a cohort of HP patients is best described in bipartite fashion, with one group featuring recurrent systemic signs and symptoms and the other featuring more severe respiratory findings.

Concordant with the variability in the presentation of HP is the observed variability in outcome. HP that has not progressed to chronic lung disease has a more favorable outcome with likely resolution if antigen avoidance can be achieved. However, chronic HP resulting in lung fibrosis has a poorer prognosis, with patients with chronic pigeon breeder's lung having demonstrated a similar mortality as seen in IPF.

DIAGNOSIS

Although there is no set of universally accepted criteria for arriving at a diagnosis of HP, diagnosis depends foremost on establishing a history of exposure to an offending antigen that correlates with respiratory and systemic symptoms. A careful occupational and home exposure history should be taken and may be supplemented if necessary by a clinician visit to the work or home environment. Specific inquiries will be influenced by geography and the occupation of the patient. When HP is suspected by history, the additional workup is aimed at establishing an

immunologic and physiologic response to inhalational antigen exposure with chest imaging, pulmonary function testing, serologic studies, bronchoscopy, and, on occasion, lung biopsy.

Chest Imaging Chest x-ray findings in HP are nonspecific and can even lack any discernible abnormalities. In cases of acute and subacute HP, findings may be transient and can include ill-defined micronodular opacities or hazy ground-glass airspace opacities. Findings on chest x-ray will often resolve with removal from the offending antigen, although the time course of resolution may vary. With chronic HP, the abnormalities seen on the chest radiograph are frequently more fibrotic in nature and may be difficult to distinguish from IPF.

With the wide availability of high-resolution computed tomography (HRCT), this modality has become a common component in the diagnostic workup for HP. Although the HRCT may be normal in acute forms of HP, this may be due to lack of temporal correlation between exposure to the offending antigen and obtaining the imaging. Additionally, because of the transient nature of acute HP, HRCT is not always performed. In subacute forms of the disease, ground-glass airspace opacities are characteristic, as is the presence of centrilobular nodules. Expiratory images may show areas of air trapping that are likely caused by involvement of the small airways (Fig. 310-1). Reticular changes and traction bronchiectasis can be observed in chronic HP. Subpleural honeycombing similar to that seen in IPF may be present in advanced cases, although unlike in IPF, the lung bases are frequently spared.

Pulmonary Function Testing (PFT) Either restrictive or obstructive PFTs can be present in HP, so the pattern of PFT change is not useful in establishing the diagnosis of HP. However, obtaining PFTs is of use in characterizing the physiologic impairment of an individual patient and in gauging the response to antigen avoidance and/or corticosteroid therapy. Diffusion capacity for carbon monoxide may be significantly impaired, particularly in cases of chronic HP with fibrotic pulmonary parenchymal changes.

Serum Precipitins Assaying for precipitating IgG antibodies against specific antigens can be a useful adjunct in the diagnosis of HP. However, the presence of an immunologic response alone is not sufficient for establishing the diagnosis, because many asymptomatic individuals with high levels of exposure to antigen may display serum precipitins, as has been observed in farmers and in pigeon breeders. It should also be noted that panels that test for several specific serum

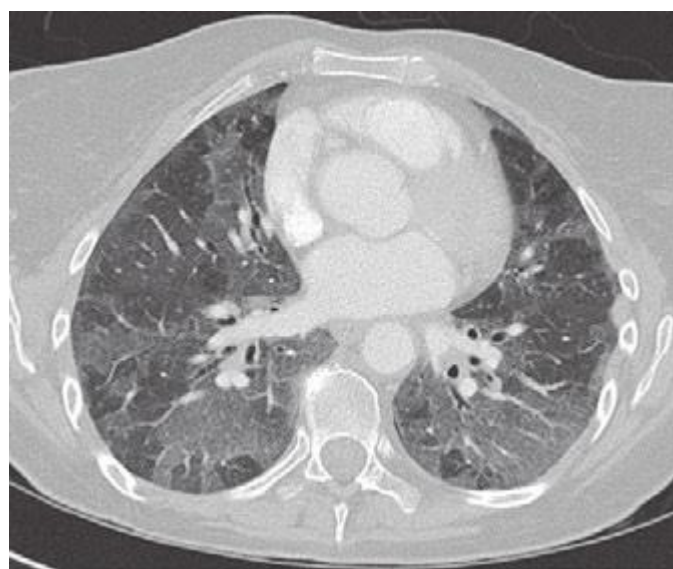


FIGURE 310-1 Chest computed tomography scan of a patient with subacute hypersensitivity pneumonitis in which scattered regions of ground-glass infiltrates in a mosaic pattern consistent with air trapping are seen bilaterally. This patient had bird fancier's lung. (Courtesy of TJ Gross; with permission.)

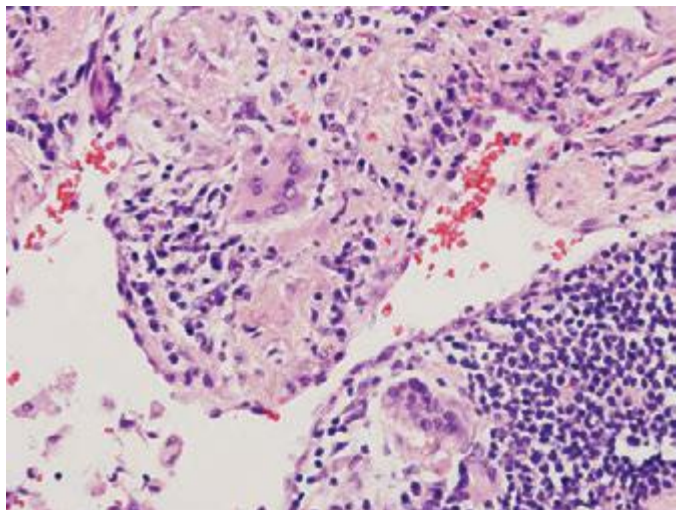


FIGURE 310-2 Open-lung biopsy from a patient with subacute hypersensitivity pneumonitis demonstrating a loose, nonnecrotizing granuloma made up of histiocytes and multinucleated giant cells. Peribronchial inflammatory infiltrate made up of lymphocytes and plasma cells is also seen. (Courtesy of TJ Gross; with permission.)

precipitins often provide false-negative results, because they represent an extremely limited proportion of the universe of potential offending environmental antigens.

Bronchoscopy Bronchoscopy with bronchoalveolar lavage (BAL) may be used in the evaluation of HP. Although not a specific finding, BAL lymphocytosis is characteristic of HP. However, in active smokers, a lower threshold should be used to establish BAL lymphocytosis, because smoking will result in lower lymphocyte percentages. Most cases of HP have a CD4+/CD8+ lymphocyte ratio of less than 1, but again, this is not a specific finding and has limited utility in the diagnosis of HP.

Lung Biopsy Tissue samples may be obtained by a bronchoscopic approach using transbronchial biopsy, or more architecturally preserved specimens may be obtained by a surgical approach (video-assisted thoracoscopy or open approach). As is the case with BAL, histologic specimens are not absolutely necessary to establish the diagnosis of HP, but they can be useful in the correct clinical context. A common histologic feature in HP is the presence of noncaseating granulomas in the vicinity of small airways (Fig. 310-2). As opposed to pulmonary sarcoidosis, in which noncaseating granulomas are well defined, the granulomas seen in HP are loose and poorly defined in nature. Within the alveolar spaces and in the interstitium, a mixed cellular infiltrate with a lymphocytic predominance is observed that is frequently patchy in distribution. Bronchiolitis with the presence of organizing exudate is also often observed. Fibrosis may be present as well, particularly as the disease progresses to its chronic form. Fibrotic changes may be focal but can be diffuse and severe with honeycombing in advanced cases, similar to findings in IPF.

Clinical Prediction Rule Although not meant as a set of validated diagnostic criteria, a clinical prediction rule for predicting the presence of HP has been published by the HP Study Group. They identified six statistically significant predictors for HP, the strongest of which was exposure to an antigen known to cause HP. Other predictive criteria were the presence of serum precipitins, recurrent symptoms, symptoms occurring 4–8 h after antigen exposure, crackles on inspiration, and weight loss.

DIFFERENTIAL DIAGNOSIS

Differentiating HP from other conditions that cause a similar constellation of respiratory and systemic symptoms requires an increased index of suspicion based on obtaining a history of possible exposure to an offending antigen. Presentations of acute or subacute HP can be mistaken for respiratory infection. In cases of chronic disease, HP

must be differentiated from interstitial lung disease, such as IPF or nonspecific interstitial pneumonitis (NSIP); this can be a difficult task even with lung biopsy. Given the presence of pulmonary infiltrates and noncaseating granulomas on biopsy, sarcoidosis is also a consideration in the differential diagnosis of HP. Unlike in HP, however, hilar adenopathy may be prominent on chest x-ray, organs other than the lung may be involved, and noncaseating granulomas in pathologic specimens tend to be well formed. Other inhalational syndromes, such as organic toxic dust syndrome (OTDS), can be misdiagnosed as HP. OTDS occurs with exposure to organic dusts, including those produced by grains or mold silage, but neither requires prior antigen sensitization nor is characterized by positive serum precipitins.

TREATMENT HYPERSENSITIVITY PNEUMONITIS

The mainstay of treatment for HP is antigen avoidance. A careful exposure history must be obtained to attempt to identify the potential offending antigen and to identify the location where a patient is exposed. Once a potential antigen and location are identified, efforts should be made to modify the environment to minimize patient exposure. This may be accomplished with measures such as removal of birds, removal of molds, and improved ventilation. Personal protective equipment including respirators and ventilated helmets can be used but may not provide adequate protection for sensitized individuals. In some cases, fully avoiding specific environments may be necessary, although such a recommendation must be balanced against the effects to an individual's lifestyle or occupation. It is not uncommon for patients with HP due to exposure to household birds to be unwilling to remove them from the home.

Because acute HP is generally a self-limited disease after a discrete exposure to an offending antigen, pharmacologic therapy is generally not necessary. However, in so-called subacute and chronic forms of the disease, there is a role for glucocorticoid therapy. In patients with particularly severe symptoms as a result of subacute HP, antigen avoidance may be insufficient after establishing the diagnosis. Although glucocorticoids do not change the long-term outcome in these patients, they can accelerate the resolution of symptoms. While there is significant variability in the approach to glucocorticoid therapy by individual clinicians, prednisone therapy can be initiated at 0.5–1 mg/kg of ideal body weight per day (not to exceed 60 mg/d or alternative glucocorticoid equivalent) over a duration of 1–2 weeks, followed by a taper over the next 2–6 weeks. In chronic HP, a similar trial of corticosteroids may be used, although a variable component of fibrotic disease may be irreversible.

GLOBAL CONSIDERATIONS



As the ever-expanding list of antigens and exposures associated with the development of HP suggests, populations at risk for HP will vary globally based on specifics of local occupational, avocational, and environmental factors. Specific examples of geographically limited HP include summer-type pneumonitis seen in Japan and suberosis seen in cork workers in Portugal and Spain.

PULMONARY INFILTRATES WITH EOSINOPHILIA

Although eosinophils are normal constituents of the lungs, there are several pulmonary eosinophilic syndromes that are characterized by pulmonary infiltrates on imaging along with an increased number of eosinophils in lung tissue, in sputum, and/or in BAL fluid, with resultant increased respiratory symptoms and the potential for systemic manifestations. Because the eosinophil plays such an important role in each of these syndromes, it is often difficult to distinguish between them, but there are important clinical and pathologic differences as well as differences in prognosis and treatment paradigms.

CLASSIFYING PULMONARY INFILTRATES WITH EOSINOPHILIA AND GENERAL APPROACH

Because there are so many different diagnoses associated with pulmonary infiltrates with eosinophilia, the first step in classifying pulmonary

TABLE 310-2 PULMONARY INFILTRATES WITH EOSINOPHILIA

Primary Pulmonary Eosinophilic Disorders
Acute eosinophilic pneumonia
Chronic eosinophilic pneumonia
Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
Hypereosinophilic syndrome
Pulmonary Disorders of Known Cause Associated with Eosinophilia
Asthma and eosinophilic bronchitis
Allergic bronchopulmonary aspergillosis
Bronchocentric granulomatosis
Drug/toxin reaction
Infection (Table 310-4)
Parasitic/helminthic disease
Nonparasitic infection
Lung Diseases Associated with Eosinophilia
Cryptogenic organizing pneumonia
Hypersensitivity pneumonitis
Idiopathic pulmonary fibrosis
Pulmonary Langerhans cell granulomatosis
Malignant Neoplasms Associated with Eosinophilia
Leukemia
Lymphoma
Lung cancer
Adenocarcinoma of various organs
Squamous cell carcinoma of various organs
Systemic Disease Associated with Eosinophilia
Postradiation pneumonitis
Rheumatoid arthritis
Sarcoidosis
Sjögren's syndrome

eosinophilic syndromes is distinguishing between primary pulmonary eosinophilic lung disorders and those with eosinophilia that are secondary to a specific cause such as a drug reaction, an infection, a malignancy, or another pulmonary condition such as asthma. **Table 310-2** lists primary and secondary pulmonary eosinophilic disorders.

For each patient, a detailed history is of utmost importance and can help elucidate what the underlying disease is. Details regarding onset, timing, and precipitants of specific symptoms can help discern one diagnosis from another. History regarding pharmacologic, occupational, and environmental exposures is instructive, and family and travel history are crucial. In addition to details about the sinuses and lungs, it is important to inquire about systemic manifestations and assess for physical findings of cardiac, gastrointestinal (GI), neurologic, dermatologic, and genitourinary involvement, all of which may give clues to specific diagnoses. Once the details from history and physical are teased out, laboratory testing (including measurements of blood eosinophils, cultures, and markers of inflammation), spirometry and radiographic imaging can help distinguish between different diseases. Often, however, BAL, transbronchial, or open lung biopsies are required. In many cases, biopsies or noninvasive diagnostic studies of other organs (e.g., echocardiogram, electromyogram, or bone marrow biopsy) can be helpful.

PATHOPHYSIOLOGY

Pathologically, the pulmonary eosinophilic syndromes are characterized by tissue infiltration by eosinophils (Fig. 310-2). In eosinophilic granulomatosis with polyangiitis (EGPA), extravascular granulomas and necrotizing vasculitis may occur in the lungs, as well as in the heart, skin, muscle, liver, spleen, and kidneys, and may be associated with fibrinoid necrosis and thrombosis.

The exact etiology of the various pulmonary eosinophilic syndromes is unknown; however, it is felt that these syndromes result from dysregulated eosinophilopoiesis or an autoimmune process because of

TABLE 310-3 DIAGNOSTIC CRITERIA OF ACUTE EOSINOPHILIC PNEUMONIA

Acute febrile illness with respiratory manifestations of <1 month in duration
Hypoxemic respiratory failure
Diffuse pulmonary infiltrates on chest x-ray
Bronchoalveolar lavage eosinophilia >25%
Absence of parasitic, fungal, or other infection
Absence of drugs known to cause pulmonary eosinophilia
Quick clinical response to corticosteroids
Failure to relapse after discontinuation of corticosteroids

the prominence of allergic features and the presence of immune complexes, heightened T cell immunity, and altered humoral immunity as evidenced by elevated IgE and rheumatoid factor. Because of its integral involvement in eosinophilopoiesis, interleukin 5 (IL-5) has been hypothesized to play an etiologic role, and efforts to block this cytokine are being investigated. Antineutrophil cytoplasmic antibodies (ANCA) are present in about half of patients with EGPA; binding of ANCA to vascular walls likely contributes to vascular inflammation and injury as well as chemotaxis of inflammatory cells.

ACUTE EOSINOPHILIC PNEUMONIA

Acute eosinophilic pneumonia is a syndrome characterized by fevers, acute respiratory failure that often requires mechanical ventilation, diffuse pulmonary infiltrates, and pulmonary eosinophilia in a previously healthy individual (**Table 310-3**).

Clinical Features and Etiology At presentation, acute eosinophilic pneumonia is often mistaken for acute lung injury or acute respiratory distress syndrome (ARDS), until a BAL is performed and reveals >25% eosinophils. Although the predominant symptoms of acute eosinophilic pneumonia are cough, dyspnea, malaise, myalgias, night sweats, and pleuritic chest pain, physical exam findings include high fevers, basilar rales, and rhonchi on forced expiration. Acute eosinophilic pneumonia most often affects males between age 20 and 40 with no history of asthma. Although no clear etiology has been identified, several case reports have linked acute eosinophilic pneumonia to recent initiation of tobacco smoking or exposure to other environmental stimuli including dust from indoor renovations.

In addition to a suggestive history, the key to establishing a diagnosis of acute eosinophilic pneumonia is the presence of >25% eosinophilia on BAL fluid. While lung biopsies show eosinophilic infiltration with acute and organizing diffuse alveolar damage, it is generally not necessary to proceed to biopsy to establish a diagnosis. Although patients present with an elevated white blood cell count, in contrast to other pulmonary eosinophilic syndromes, acute eosinophilic pneumonia is often not associated with peripheral eosinophilia upon presentation. However, between 7 and 30 days of disease onset, peripheral eosinophilia often occurs with mean eosinophil counts of 1700. Erythrocyte sedimentation rate (ESR), C-reactive protein, and IgE levels are high but nonspecific, whereas HRCT is always abnormal with bilateral random patchy ground-glass or reticular opacities, and small pleural effusions in as many as two-thirds of patients. Pleural fluid is characterized by a high pH with marked eosinophilia.

Clinical Course and Response to Therapy Although some patients improve spontaneously, most patients require admission to an intensive care unit and respiratory support with either invasive (intubation) or noninvasive mechanical ventilation. However, what distinguishes acute eosinophilic pneumonia from both other cases of acute lung injury as well as some of the other pulmonary eosinophilic syndromes is the absence of organ dysfunction or multisystem organ failure other than respiratory failure. One of the characteristic features of acute eosinophilic pneumonia is the high degree of corticosteroid responsiveness and the excellent prognosis. Another distinguishing feature of acute eosinophilic pneumonia is that complete clinical and radiographic recovery without recurrence or residual sequelae occurs in almost all patients within several weeks of initiation of therapy.

CHRONIC EOSINOPHILIC PNEUMONIA

In contrast to acute eosinophilic pneumonia, chronic eosinophilic pneumonia is a more indolent syndrome that is characterized by pulmonary infiltrates and eosinophilia in both the tissue and blood. Most patients are female nonsmokers with a mean age of 45, and patients do not usually develop the acute respiratory failure and significant hypoxemia appreciated in acute eosinophilic pneumonia. Similar to EGPA, a majority have asthma, with many having a history of allergies.

Patients present with a subacute illness over weeks to months, with cough, low-grade fevers, progressive dyspnea, weight loss, wheezing, malaise, and night sweats, and a chest x-ray with migratory bilateral peripheral or pleural-based opacities. Although this “photographic negative pulmonary edema” appearance on chest x-ray and chest CT is pathognomonic of chronic eosinophilic pneumonia, less than 25% of patients present with this finding. Other radiographic findings include atelectasis, pleural effusions, lymphadenopathy, and septal line thickening.

Almost 90% of patients have peripheral eosinophilia, with mean eosinophil counts of over 30% of total white blood cell count. BAL eosinophilia is also an important distinguishing feature with mean BAL eosinophil counts of close to 60%. Both peripheral and BAL eosinophilia are very responsive to treatment with corticosteroids. Other laboratory features of chronic eosinophilic pneumonia include increased ESR, C-reactive protein, platelets, and IgE. Lung biopsy is also often not required to establish a diagnosis, but may show accumulation of eosinophils and histiocytes in the lung parenchyma and interstitium, as well as cryptogenic organizing pneumonia, but with minimal fibrosis. Nonrespiratory manifestations are uncommon, but arthralgias, neuropathy, and skin and GI symptoms have all been reported; their presence may suggest EGPA or hypereosinophilic syndrome. Another similarity is the rapid response to corticosteroids with quick resolution of peripheral and BAL eosinophilia and improvement in symptoms. In contrast to acute eosinophilic pneumonia, though, over 50% of patients relapse, and many require prolonged courses of corticosteroids for months to years.

EOSINOPHILIC GRANULOMATOSIS WITH POLYANGIITIS (EGPA)

Previously known as allergic angitis granulomatosis or Churg-Strauss syndrome, this complex syndrome is characterized by eosinophilic vasculitis that may involve multiple organ systems including the lungs, heart, skin, GI tract, and nervous system. Although EGPA is characterized by peripheral and pulmonary eosinophilia with infiltrates on chest x-ray, the primary features that distinguish EGPA from other pulmonary eosinophilic syndromes are the presence of eosinophilic vasculitis in the setting of asthma and involvement of multiple end organs (a feature it shares with hypereosinophilic syndrome). Although perceived to be quite rare, in the last few years, there has appeared to be an increased incidence of this disease, particularly in association with various asthma therapies.

The primary features of EGPA include asthma, peripheral eosinophilia, neuropathy, pulmonary infiltrates, paranasal sinus abnormality, and presence of eosinophilic vasculitis. It typically occurs in several phases. The prodromal phase is characterized by asthma and allergic rhinitis, and usually begins when the individual is in his or her twenties or thirties, typically persisting for many years. The eosinophilic infiltrative phase is characterized by peripheral eosinophilia and eosinophilic tissue infiltration of various organs including the lungs and GI tract. The third phase is the vasculitic phase and may be associated with constitutional signs and symptoms including fever, weight loss, malaise, and fatigue. The mean age at diagnosis is 48 years, with a range of 14 to 74 years; the average length of time between diagnosis of asthma and vasculitis is 9 years.

Similar to other pulmonary eosinophilic syndromes, constitutional symptoms are very common in EGPA and include weight loss of 10–20 lb, fevers, and diffuse myalgias and migratory polyarthralgias. Myositis may be present with evidence of vasculitis on muscle biopsies. In contrast to the eosinophilic pneumonias, EGPA involves many organ systems including the lungs, skin, nerves, heart, GI tract, and kidneys.

Symptoms and Clinical Manifestations • RESPIRATORY Most EGPA patients have asthma that arises later in life and in individuals who have no family history of atopy. The asthma can often be severe, and oral corticosteroids are often required to control symptoms but may lead to suppression of vasculitic symptoms. In addition to the more common symptoms of cough, dyspnea, sinusitis, and allergic rhinitis, alveolar hemorrhage and hemoptysis may also occur.

NEUROLOGIC Over three-fourths of EGPA patients have neurologic manifestations. Mononeuritis multiplex most commonly involves the peroneal nerve, but also involves the ulnar, radial, internal popliteal, and occasionally, cranial nerves. Cerebral hemorrhage and infarction may also occur and are important causes of death. Despite treatment, neurologic sequelae often do not completely resolve.

DERMATOLOGIC Approximately half of EGPA patients develop dermatologic manifestations. These include palpable purpura, skin nodules, urticarial rashes, and livedo.

CARDIOVASCULAR Granulomas, vasculitis, and widespread myocardial damage may be found on biopsy or at autopsy, and cardiomyopathy and heart failure may be seen in up to half of all patients but are often at least partially reversible. Acute pericarditis, constrictive pericarditis, myocardial infarction, and other electrocardiographic changes all may occur. The heart is a primary target organ in EGPA, and cardiac involvement often portends a worse prognosis.

GI GI symptoms are common in EGPA and likely represent an eosinophilic gastroenteritis characterized by abdominal pain, diarrhea, GI bleeding, and colitis. Ischemic bowel, pancreatitis, and cholecystitis have also been reported in association with EGPA and usually portend a worse prognosis.

RENAL Renal involvement is more common than once thought, and approximately 25% of patients have some degree of renal involvement. This may include proteinuria, glomerulonephritis, renal insufficiency, and rarely, renal infarct.

Lab Abnormalities Systemic eosinophilia is the hallmark laboratory finding in patients with EGPA and reflects the likely pathogenic role that the eosinophil plays in this disease. Eosinophilia greater than 10% is one of the defining features of this illness and may be as high as 75% of the peripheral white blood cell count. It is present at the time of diagnosis in over 80% of patients but may respond quickly (often within 24 h) to initiation of systemic corticosteroid therapy. Even in the absence of systemic eosinophilia, tissue eosinophilia may be present.

Although not specific to EGPA, ANCA are present in up to two-thirds of patients, mostly with a perinuclear staining pattern. Nonspecific lab abnormalities that may be present in patients with EGPA include a marked elevation in ESR, a normochromic normocytic anemia, an elevated IgE, hypergammaglobulinemia, and positive rheumatoid factor and antinuclear antibodies (ANA). Although BAL often reveals significant eosinophilia, this may be seen in other eosinophilic lung diseases. Similarly, PFT often reveals an obstructive defect similar to asthma.

Radiographic Features Chest x-ray abnormalities are extremely common in EGPA and consist of bilateral, nonsegmental, patchy infiltrates that often migrate and may be interstitial or alveolar in appearance. Reticulonodular and nodular disease without cavitation can be seen, as can pleural effusions and hilar adenopathy. The most common CT findings include bilateral ground-glass opacity and airspace consolidation that is predominantly subpleural. Other CT findings include bronchial wall thickening, hyperinflation, interlobular septal thickening, lymph node enlargement, and pericardial and pleural effusions. Angiography may be used diagnostically and may show signs of vasculitis in the coronary, central nervous system, and peripheral vasculature.

Treatment and Prognosis of EGPA Most patients diagnosed with EGPA have previously been diagnosed with asthma, rhinitis, and sinusitis, and have received treatment with inhaled or systemic corticosteroids.

1686 Because these agents are also the initial treatment of choice for EGPA patients, institution of these therapies in patients with EGPA who are perceived to have severe asthma may delay the diagnosis of EGPA because signs of vasculitis may be masked. Corticosteroids dramatically alter the course of EGPA: up to 50% of those who are untreated die within 3 months of diagnosis, whereas treated patients have a 6-year survival of over 70%. Common causes of death include heart failure, cerebral hemorrhage, renal failure, and GI bleeding. Recent data suggest that clinical remission may be obtained in over 90% of patients treated; approximately 25% of those patients may relapse, often due to corticosteroid tapering, with a rising eosinophil count heralding the relapse. Myocardial, GI, and renal involvement most often portend a poor prognosis. In such cases, treatment with higher doses of corticosteroids or the addition of cytotoxic agents such as cyclophosphamide is often warranted. Although survival does not differ between those treated or untreated with cyclophosphamide, cyclophosphamide is associated with a reduced incidence of relapse and an improved clinical response to treatment. Other therapies that have been used successfully in the management of EGPA include azathioprine, methotrexate, intravenous gamma globulin, and interferon α . Plasma exchange has not been shown to provide any additional benefit. Recent studies examining the efficacy of anti-IL-5 therapy have shown promise.

HYPEREOSINOPHILIC SYNDROMES

Hyper eosinophilic syndromes (HES) constitute a heterogeneous group of disease entities manifest by persistent eosinophilia >1500 eosinophils/ μL in association with end organ damage or dysfunction, in the absence of secondary causes of eosinophilia. In addition to familial, undefined, and overlap syndromes with incomplete criteria, the predominant HES subtypes are the myeloproliferative and lymphocytic variants. The myeloproliferative variant may be divided into three subgroups: (1) chronic eosinophilic leukemia with demonstrable cytogenetic abnormalities and/or blasts on peripheral smear; (2) the platelet-derived growth factor receptor α (PDGFR α)-associated HES, attributed to a constitutively activated tyrosine kinase fusion protein (Fip1L1-PDGFR α) due to a chromosomal deletion on 4q12; this variant is often responsive to imatinib; and (3) the FIP1-negative variant associated with clonal eosinophilia and at least four of the following: dysplastic peripheral eosinophils, increased serum vitamin B₁₂, increased tryptase, anemia, thrombocytopenia, splenomegaly, bone marrow cellularity $>80\%$, spindle-shaped mast cells, and myelofibrosis.

Extrapulmonary Manifestations of HES More common in men than in women, HES occurs between the ages of 20 and 50 and is characterized by significant extrapulmonary involvement, including infiltration of the heart, GI tract, kidney, liver, joints, and skin. Cardiac involvement includes myocarditis and/or endomyocardial fibrosis, as well as a restrictive cardiomyopathy.

Pulmonary Manifestations of HES Similar to the other pulmonary eosinophilic syndromes, these HES are manifest by high levels of blood, BAL, and tissue eosinophilia. Lung involvement occurs in 40% of these patients and is characterized by cough and dyspnea, as well as pulmonary infiltrates. Although it is often difficult to discern the pulmonary infiltrates and effusions seen on chest x-ray from pulmonary edema resulting from cardiac involvement, CT scan findings include interstitial infiltrates, ground-glass opacities, and small nodules. HES are typically not associated with ANCA or elevated IgE.

Course and Response to Therapy Unlike the other pulmonary eosinophilic syndromes, less than half of patients with these HES respond to corticosteroids as first-line therapy. Although other treatment options include hydroxyurea, cyclosporine, and interferon, the tyrosine kinase inhibitor imatinib has emerged as an important therapeutic option for patients with the myeloproliferative variant. Anti-IL-5 therapy with mepolizumab also holds promise for these patients and is currently being investigated.

ALLERGIC BRONCHOPULMONARY ASPERGILLOSIS

Allergic bronchopulmonary aspergillosis (ABPA) is an eosinophilic pulmonary disorder that occurs in response to allergic sensitization to antigens from *Aspergillus* species fungi. The predominant clinical presentation of ABPA is an asthmatic phenotype, often accompanied by cough with production of brownish plugs of mucus. ABPA has also been well described as a complication of cystic fibrosis. A workup for ABPA may be beneficial in patients who carry a diagnosis of asthma but have proven refractory to usual therapy. ABPA is a distinct diagnosis from simple asthma, characterized by prominent peripheral eosinophilia and elevated circulating levels of IgE (>417 IU/mL). Establishing a diagnosis of ABPA also requires establishing sensitivity to *Aspergillus* antigens by skin test reactivity, positive serum precipitins for *Aspergillus*, and/or direct measurement of circulating specific IgG and IgE to *Aspergillus*. Central bronchiectasis is described as a classic finding on chest imaging in ABPA but is not necessary for making a diagnosis. Other possible findings on chest imaging include patchy infiltrates and evidence of mucus impaction.

Systemic glucocorticoids may be used in the treatment of ABPA that is persistently symptomatic despite the use of inhaled therapies for asthma. Courses of glucocorticoids should be tapered over 3–6 months, and their use must be balanced against the risks of prolonged steroid therapy. Antifungal agents such as fluconazole and voriconazole given over a 4-month course reduce the antigenic stimulus in ABPA and may therefore modulate disease activity in selected patients. The use of monoclonal antibody against IgE (omalizumab) has been described in treating severe ABPA, particularly in individuals with ABPA as a complication of cystic fibrosis.

ABPA-like syndromes have been reported as a result to sensitization to several non-*Aspergillus* species fungi. However, these conditions are substantially rarer than ABPA, which may be present in a significant proportion of patients with refractory asthma.

INFECTIOUS PROCESSES

Infectious etiologies of pulmonary eosinophilia are largely due to helminths and are of particular importance in the evaluation of pulmonary eosinophilia in tropical environments and in the developing world (Table 310-4). These infectious conditions may also be considered in recent travelers to endemic regions. Löffler syndrome refers to transient pulmonary infiltrates with eosinophilia that occurs in response to passage of helminthic larvae through the lungs, most

TABLE 310-4 INFECTIOUS CAUSES OF PULMONARY EOSINOPHILIA

Löffler Syndrome
<i>Ascaris</i>
Hookworm
Schistosomiasis
Heavy Parasite Burden
Strongyloidiasis
Direct Pulmonary Penetration
Paragonimiasis
Visceral larval migrans
Immunologic Response to Organisms in Lungs
Filariasis
Dirofilariasis
Cystic Disease
<i>Echinococcus</i>
Cysticercosis
Other Nonparasitic
Coccidioidomycosis
Basidiobolomycosis
Paracoccidioidomycosis
Tuberculosis

Source: Adapted from P Akuthota, PF Weller: Clin Microbiol Rev 25:649, 2012.

commonly larvae of *Ascaris* species (roundworm). Symptoms are generally self-limited and may include dyspnea, cough, wheeze, and hemoptysis. Löffler syndrome may also occur in response to hookworm infection with *Ancylostoma duodenale* or *Necator americanus*. Chronic *Strongyloides stercoralis* infection can lead to recurrent respiratory symptoms with peripheral eosinophilia between flares. In immunocompromised hosts, including patients on glucocorticoids, a severe, potentially fatal, hyperinfection syndrome can result from *Strongyloides* infection. Paragonimiasis, filariasis, and visceral larval migrans can all cause pulmonary eosinophilia as well.

DRUGS AND TOXINS

A host of medications are associated with the development of pulmonary infiltrates with peripheral eosinophilia. Therefore, drug reaction must always be included in the differential diagnosis of pulmonary eosinophilia. Although the list of medications associated with pulmonary eosinophilia is ever expanding, common culprits include nonsteroidal anti-inflammatory medications and systemic antibiotics, most specifically nitrofurantoin. Additionally, various and diverse environmental exposures such as particulate metals, scorpion stings, and inhalational drugs of abuse may also cause pulmonary eosinophilia. Radiation therapy for breast cancer has been linked with eosinophilic pulmonary infiltration as well. The mainstay of treatment is removal of the offending exposure, although glucocorticoids may be necessary if respiratory symptoms are severe.

GLOBAL CONSIDERATIONS



In the United States, drug-induced eosinophilic pneumonias are the most common cause of eosinophilic pulmonary infiltrates. A travel history or evidence of recent immigration should prompt the consideration of parasite-associated disorders. Tropical eosinophilia is usually caused by filarial infection; however, eosinophilic pneumonias also occur with other parasites such as *Ascaris* spp., *Ancylostoma* spp., *Toxocara* spp., and *Strongyloides stercoralis*. Tropical eosinophilia due to *Wuchereria bancrofti* or *Wuchereria malayi* occurs most commonly in southern Asia, Africa, and South America and is treated successfully with diethylcarbamazine. In the United States, *Strongyloides* is endemic to the southeastern and Appalachian regions.

ACKNOWLEDGMENTS

We acknowledge the contributions of Dr. Alicia K. Gerke and Dr. Gary W. Hunninghake to the previous edition of this chapter.

311 Occupational and Environmental Lung Disease

John R. Balmes, Frank E. Speizer

Occupational and environmental lung diseases are difficult to distinguish from those of nonenvironmental origin. Virtually all major categories of pulmonary disease can be caused by environmental agents, and environmentally related disease usually presents clinically in a manner indistinguishable from that of disease not caused by such agents. In addition, the etiology of many diseases may be multifactorial; occupational and environmental factors may interact with other factors (such as smoking and genetic risk). It is often only after a careful exposure history is taken that the underlying workplace or general environmental exposure is uncovered.

Why is knowledge of occupational or environmental etiology so important? Patient management and prognosis are affected significantly by such knowledge. For example, patients with occupational

asthma or hypersensitivity pneumonitis often cannot be managed adequately without cessation of exposure to the offending agent. Establishment of cause may have significant legal and financial implications for a patient who no longer can work in his or her usual job. Other exposed people may be identified as having the disease or prevented from getting it. In addition, new associations between exposure and disease may be identified (e.g., nylon flock worker's lung disease and diacetyl-induced bronchiolitis obliterans).

Although the exact proportion of lung disease due to occupational and environmental factors is unknown, a large number of individuals are at risk. For example, 15–20% of the burden of adult asthma and chronic obstructive pulmonary disease (COPD) has been estimated to be due to occupational factors.

HISTORY AND EXPOSURE ASSESSMENT

The patient's history is of paramount importance in assessing any potential occupational or environmental exposure. Inquiry into specific work practices should include questions about the specific contaminants involved, the presence of visible dusts, chemical odors, the size and ventilation of workspaces, the use of respiratory protective equipment, and whether co-workers have similar complaints. The temporal association of exposure at work and symptoms may provide clues to occupation-related disease. In addition, the patient must be questioned about alternative sources of exposure to potentially toxic agents, including hobbies, home characteristics, exposure to secondhand smoke, and proximity to traffic or industrial facilities. Short-term and long-term exposures to potential toxic agents in the distant past also must be considered.

Workers in the United States have the right to know about potential hazards in their workplaces under federal Occupational Safety and Health Administration (OSHA) regulations. Employers must provide specific information about potential hazardous agents in products being used through Material Safety Data Sheets as well as training in personal protective equipment and environmental control procedures. However, the introduction of new processes and/or new chemical compounds may change exposure significantly, and often only the employee on the production line is aware of the change. For the physician caring for a patient with a suspected work-related illness, a visit to the work site can be very instructive. Alternatively, an affected worker can request an inspection by OSHA. If reliable environmental sampling data are available, that information should be used in assessing a patient's exposure. Because many of the chronic diseases result from exposure over many years, current environmental measurements should be combined with work histories to arrive at estimates of past exposure.

PULMONARY FUNCTION TESTS AND CHEST IMAGING

Exposures to inorganic and organic dusts can cause interstitial lung disease that presents with a restrictive pattern and a decreased diffusing capacity (Chap. 306e). Similarly, exposures to a number of organic dusts or chemical agents may result in occupational asthma or COPD that is characterized by airway obstruction. Measurement of change in forced expiratory volume (FEV₁) before and after a working shift can be used to detect an acute bronchoconstrictive response.

The chest radiograph is useful in detecting and monitoring the pulmonary response to mineral dusts, certain metals, and organic dusts capable of inducing hypersensitivity pneumonitis. The International Labour Organisation (ILO) International Classification of Radiographs of Pneumoconioses classifies chest radiographs by the nature and size of opacities seen and the extent of involvement of the parenchyma. In general, small rounded opacities are seen in silicosis or coal worker's pneumoconiosis, and small linear opacities are seen in asbestosis. Although useful for epidemiologic studies and screening large numbers of workers, the ILO system can be problematic when applied to an individual worker's chest radiograph. With dusts causing rounded opacities, the degree of involvement on the chest radiograph may be extensive, whereas pulmonary function may be only minimally impaired. In contrast, in pneumoconiosis causing linear, irregular opacities like those seen in asbestosis, the radiograph may lead to

1688 underestimation of the severity of the impairment until relatively late in the disease. For patients with a history of asbestos exposure, conventional computed tomography (CT) is more sensitive for the detection of pleural thickening, and high-resolution CT (HRCT) improves the detection of asbestosis.

Other procedures that may be of use in identifying the role of environmental exposures in causing lung disease include skin prick testing or specific IgE antibody titers for evidence of immediate hypersensitivity to agents capable of inducing occupational asthma (flour antigens in bakers), specific IgG precipitating antibody titers for agents capable of causing hypersensitivity pneumonitis (pigeon antigen in bird handlers), and assays for specific cell-mediated immune responses (beryllium lymphocyte proliferation testing in nuclear workers or tuberculin skin testing in health care workers). Sometimes a bronchoscopy to obtain transbronchial biopsies of lung tissue may be required for histologic diagnosis (chronic beryllium disease). Rarely, video-assisted thoracoscopic surgery to obtain a larger sample of lung tissue may be required to determine the specific diagnosis of environmentally induced lung disease (hypersensitivity pneumonitis or giant cell interstitial pneumonitis due to cobalt exposure).

DETERMINANTS OF INHALATIONAL EXPOSURE

The chemical and physical characteristics of inhaled agents affect both the dose and the site of deposition in the respiratory tract. Water-soluble gases such as ammonia and sulfur dioxide are absorbed in the lining fluid of the upper and proximal airways and thus tend to produce irritative and bronchoconstrictive responses. In contrast, nitrogen dioxide and phosgene, which are less soluble, may penetrate to the bronchioles and alveoli in sufficient quantities to produce acute chemical pneumonitis.

Particle size of air contaminants must also be considered. Because of their settling velocities in air, particles >10–15 μm in diameter do not penetrate beyond the nose and throat. Particles <10 μm in size are deposited below the larynx. These particles are divided into three size fractions on the basis of their size characteristics and sources. Particles

~2.5–10 μm (coarse-mode fraction) contain crustal elements such as silica, aluminum, and iron. These particles mostly deposit relatively high in the tracheobronchial tree. Although the total mass of an ambient sample is dominated by these larger respirable particles, the number of particles, and therefore the surface area on which potential toxic agents can deposit and be carried to the lower airways, is dominated by particles <2.5 μm (fine-mode fraction). These fine particles are created primarily by the burning of fossil fuels or high-temperature industrial processes resulting in condensation products from gases, fumes, or vapors. The smallest particles, those <0.1 μm in size, represent the ultrafine fraction and make up the largest number of particles; they tend to remain in the airstream and deposit in the lung only on a random basis as they come into contact with the alveolar walls. If they do deposit, however, particles of this size range may penetrate into the circulation and be carried to extrapulmonary sites. New technologies create particles of this size (“nanoparticles”) for use in many commercial applications. Besides the size characteristics of particles and the solubility of gases, the actual chemical composition, mechanical properties, and immunogenicity or infectivity of inhaled material determine in large part the nature of the diseases found among exposed persons.

OCCUPATIONAL EXPOSURES AND PULMONARY DISEASE

Table 311-1 provides broad categories of exposure in the workplace and diseases associated with chronic exposure in those industries.

ASBESTOS-RELATED DISEASES

Asbestos is a generic term for several different mineral silicates, including chrysolite, amosite, anthophyllite, and crocidolite. In addition to workers involved in the production of asbestos products (mining, milling, and manufacturing), many workers in the shipbuilding and construction trades, including pipe fitters and boilermakers, were occupationally exposed because asbestos was widely used during the twentieth century for its thermal and electrical insulation properties. Asbestos also was used in the manufacture of fire-resistant textiles, in cement and floor tiles, and in friction materials such as brake and clutch linings.

TABLE 311-1 CATEGORIES OF OCCUPATIONAL EXPOSURE AND ASSOCIATED RESPIRATORY CONDITIONS

Occupational Exposures	Nature of Respiratory Responses	Comment
Inorganic Dusts		
Asbestos: mining, processing, construction, ship repair	Fibrosis (asbestosis), pleural disease, cancer, mesothelioma	Virtually all new mining and construction with asbestos done in developing countries
Silica: mining, stone cutting, sandblasting, quarrying	Fibrosis (silicosis), progressive massive fibrosis (PMF), cancer, tuberculosis, chronic obstructive pulmonary disease (COPD)	Improved protection in United States; persistent risk in developing countries
Coal dust: mining	Fibrosis (coal worker’s pneumoconiosis), PMF, COPD	Risk persists in certain areas of United States, increasing in countries where new mines open
Beryllium: processing alloys for high-tech industries	Acute pneumonitis (rare), chronic granulomatous disease, lung cancer (highly suspect)	Risk in high-tech industries persists
Other metals: aluminum, chromium, cobalt, nickel, titanium, tungsten carbide, or “hard metal” (contains cobalt)	Wide variety of conditions from acute pneumonitis to lung cancer and asthma	New diseases appear with new process development
Organic Dusts		
Cotton dust: milling, processing	Byssinosis (an asthma-like syndrome), chronic bronchitis, COPD	Increasing risk in developing countries with drop in United States as jobs shift overseas
Grain dust: elevator agents, dock workers, milling, bakers	Asthma, chronic bronchitis, COPD	Risk shifting more to migrant labor pool
Other agricultural dusts: fungal spores, vegetable products, insect fragments, animal dander, bird and rodent feces, endotoxins, microorganisms, pollens	Hypersensitivity pneumonitis (farmer’s lung), asthma, chronic bronchitis	Important in migrant labor pool but also resulting from in-home exposures
Toxic chemicals: wide variety of industries; see Table 311-2	Asthma, chronic bronchitis, COPD, hypersensitivity pneumonitis, pneumoconiosis, and cancer	Reduced risk with recognized hazards; increasing risk for developing countries where controlled labor practices are less stringent
Other respiratory environmental agents: uranium and radon daughters, secondhand tobacco smoke, polycyclic aromatic hydrocarbons (PAHs), biomass smoke, diesel exhaust, welding fumes, wood finishing	Occupational exposures estimated to contribute to up to 10% of all lung cancers; chronic bronchitis, COPD, and fibrosis	In-home exposures important; in developing countries, biomass smoke is a major risk factor for COPD among women

Exposure to asbestos is not limited to persons who directly handle the material. Cases of asbestos-related diseases have been encountered in individuals with only bystander exposure, such as painters and electricians who worked alongside insulation workers in a shipyard. Community exposure resulted from the use of asbestos-containing mine and mill tailings as landfill, road surface, and playground material (e.g., Libby, MT, the site of a vermiculite mine in which the ore was contaminated with asbestos). Finally, exposure can occur from the disturbance of naturally occurring asbestos (e.g., from increasing residential development in the foothills of the Sierra Mountains in California).

Asbestos has largely been replaced in the developed world with synthetic mineral fibers such as fiberglass and refractory ceramic fibers, but it continues to be used in the developing world. The major health effects from exposure to asbestos are pleural and pulmonary fibrosis, cancers of the respiratory tract, and pleural and peritoneal mesothelioma.

Asbestosis is a diffuse interstitial fibrosing disease of the lung that is directly related to the intensity and duration of exposure. The disease resembles other forms of diffuse interstitial fibrosis (Chap. 315). Usually, exposure has taken place for at least 10 years before the disease becomes manifest. The mechanisms by which asbestos fibers induce lung fibrosis are not completely understood but are known to involve oxidative injury due to the generation of reactive oxygen species by the transition metals on the surface of the fibers as well as from cells engaged in phagocytosis.

Past exposure to asbestos is specifically indicated by pleural plaques on chest radiographs, which are characterized by either thickening or calcification along the parietal pleura, particularly along the lower lung fields, the diaphragm, and the cardiac border. Without additional manifestations, pleural plaques imply only exposure, not pulmonary impairment. Benign pleural effusions also may occur. The fluid is typically a serous or bloody exudate. The effusion may be slowly progressive or may resolve spontaneously.

Irregular or linear opacities that usually are first noted in the lower lung fields are the chest radiographic hallmark of asbestosis. An indistinct heart border or a “ground-glass” appearance in the lung fields may be seen. HRCT may show distinct changes of subpleural curvilinear lines 5–10 mm in length that appear to be parallel to the pleural surface (Fig. 311-1).

Pulmonary function testing in asbestosis reveals a restrictive pattern with a decrease in both lung volumes and diffusing capacity. There may also be evidence of mild airflow obstruction (due to peribronchiolar fibrosis).

Because no specific therapy is available for asbestosis, supportive care is the same as that given to any patient with diffuse interstitial fibrosis of any cause. In general, newly diagnosed cases will have resulted from exposures that occurred many years before.

Lung cancer (Chap. 107) is the most common cancer associated with asbestos exposure. The excess frequency of lung cancer (all histologic types) in asbestos workers is associated with a minimum latency of 15–19 years between first exposure and development of the disease. Persons with more exposure are at greater risk of disease. In addition, there is a significant interactive effect of smoking and asbestos exposure that results in greater risk than what would be expected from the additive effect of each factor.

Mesotheliomas (Chap. 316), both pleural and peritoneal, are also associated with asbestos exposure. In contrast to lung cancers, these tumors do not appear to be associated with smoking. Relatively short-term asbestos exposures of ≤ 1 –2 years, occurring up to 40 years in the past, have been associated with the development of mesotheliomas (an observation that emphasizes the importance of obtaining a complete environmental exposure history). Although the risk of mesothelioma is much less than that of lung cancer among asbestos-exposed workers, over 2000 cases were reported in the United States per year at the start of the twenty-first century.

Because epidemiologic studies have shown that >80% of mesotheliomas may be associated with asbestos exposure, documented mesothelioma in a patient with occupational or environmental exposure to asbestos may be compensable.

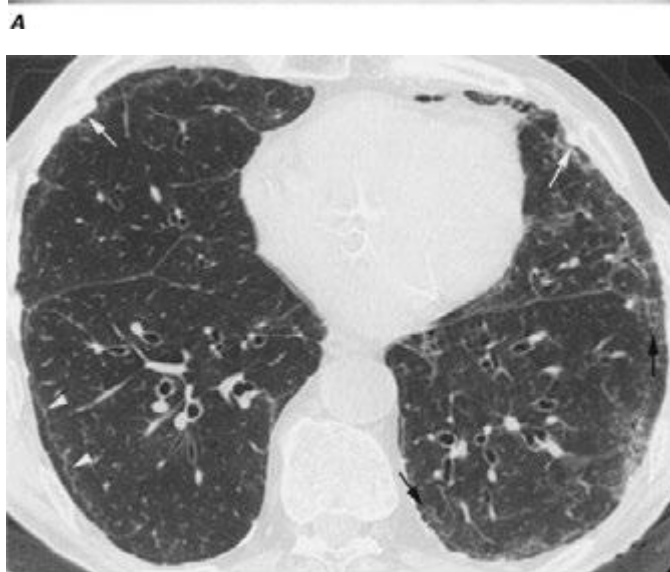
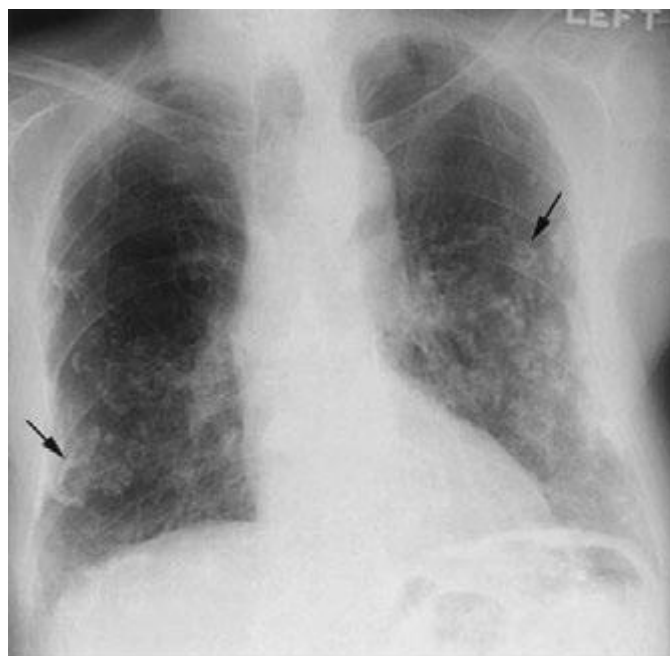


FIGURE 311-1 Asbestosis. **A.** Frontal chest radiograph shows bilateral calcified pleural plaques consistent with asbestos-related pleural disease. Poorly defined linear and reticular abnormalities are seen in the lower lobes bilaterally. **B.** Axial high-resolution computed tomography of the thorax obtained through the lung bases shows bilateral, subpleural reticulation (black arrows), representing fibrotic lung disease due to asbestosis. Subpleural lines are also present (arrowheads), characteristic of, though not specific for, asbestosis. Calcified pleural plaques representing asbestos-related pleural disease (white arrows) are also evident.

SILICOSIS

Despite being one of the oldest known occupational pulmonary hazards, *free silica* (SiO_2), or crystalline quartz, is still a major cause of disease. The major occupational exposures include mining; stonecutting; sand blasting; glass and cement manufacturing; foundry work; packing of silica flour; and quarrying, particularly of granite. Most often, pulmonary fibrosis due to silica exposure (silicosis) occurs in a dose-response fashion after many years of exposure.

Workers heavily exposed through sandblasting in confined spaces, tunneling through rock with a high quartz content (15–25%), or the manufacture of abrasive soaps may develop acute silicosis with as little as 10 months of exposure. The clinical and pathologic features of acute silicosis are similar to those of pulmonary alveolar proteinosis



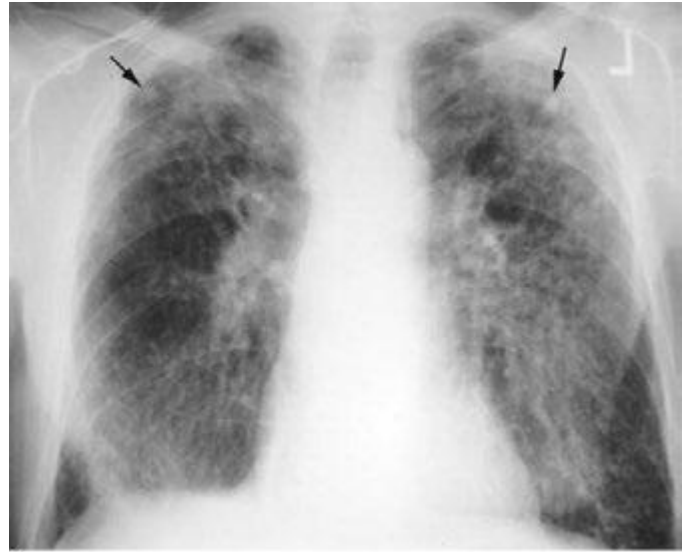
FIGURE 311-2 Acute silicosis. This high-resolution computed tomography scan shows multiple small nodules consistent with silicosis but also diffuse ground-glass densities with thickened intralobular and interlobular septa producing polygonal shapes. This has been referred to as “crazy paving.”

(Chap. 315). The chest radiograph may show profuse miliary infiltration or consolidation, and there is a characteristic HRCT pattern known as “crazy paving” (Fig. 311-2). The disease may be quite severe and progressive despite the discontinuation of exposure. Whole-lung lavage may provide symptomatic relief and slow the progression.

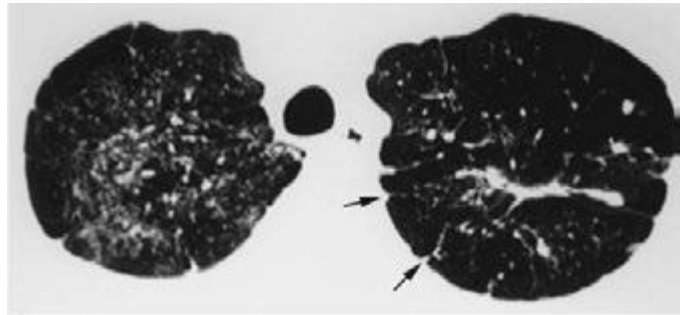
With long-term, less intense exposure, small rounded opacities in the upper lobes may appear on the chest radiograph after 15–20 years of exposure, usually without associated impairment of lung function (*simple silicosis*). Calcification of hilar nodes may occur in as many as 20% of cases and produces a characteristic “eggshell” pattern. Silicotic nodules may be identified more readily by HRCT (Fig. 311-3). The nodular fibrosis may be progressive in the absence of further exposure, with coalescence and formation of nonsegmental conglomerates of irregular masses >1 cm in diameter (*complicated silicosis*). These masses can become quite large, and when this occurs, the term *progressive massive fibrosis* (PMF) is applied. Significant functional impairment with both restrictive and obstructive components may be associated with PMF.

Because silica is cytotoxic to alveolar macrophages, patients with silicosis are at greater risk of acquiring lung infections that involve these cells as a primary defense (*Mycobacterium tuberculosis*, atypical mycobacteria and fungi). Because of the increased risk of active tuberculosis, the recommended treatment of latent tuberculosis in these patients is longer. Another potential clinical complication of silicosis is autoimmune connective tissue disorders such as rheumatoid arthritis and scleroderma. In addition, there are sufficient epidemiologic data that the International Agency for Research on Cancer lists silica as a probable lung carcinogen.

Other, less hazardous silicates include fuller’s earth, kaolin, mica, diatomaceous earths, silica gel, soapstone, carbonate dusts, and cement dusts. The production of fibrosis in workers exposed to these agents is believed to be related either to the free silica content of these dusts or, for substances that contain no free silica, to the potentially large dust loads to which these workers may be exposed. Some silicates, including *talc* and *vermiculite*, may be contaminated with asbestos. Fibrosis of lung or pleura, lung cancer, and mesothelioma have been associated with chronic exposure to talc and vermiculite dusts.



A



B

FIGURE 311-3 Chronic silicosis. **A.** Frontal chest radiograph in a patient with silicosis shows variably sized, poorly defined nodules (arrows) predominating in the upper lobes. **B.** Axial thoracic computed tomography image through the lung apices shows numerous small nodules, more pronounced in the right upper lobe. A number of the nodules are subpleural in location (arrows).

COAL WORKER’S PNEUMOCONIOSIS (CWP)

Occupational exposure to *coal dust* can lead to CWP, which has enormous social, economic, and medical significance in every nation in which coal mining is an important industry. Simple radiographically identified CWP is seen in ~10% of all coal miners and in as many as 50% of anthracite miners with more than 20 years of work on the coal face. The prevalence of disease is lower in workers in bituminous coal mines.

With prolonged exposure to coal dust (i.e., 15–20 years), small, rounded opacities similar to those of silicosis may develop. As in silicosis, the presence of these nodules (*simple CWP*) usually is not associated with pulmonary impairment. In addition to CWP, coal dust can cause chronic bronchitis and COPD (Chap. 314). The effects of coal dust are additive to those of cigarette smoking.

Complicated CWP is manifested by the appearance on the chest radiograph of nodules ≥ 1 cm in diameter generally confined to the upper half of the lungs. As in silicosis, this condition can progress to PMF that is accompanied by severe lung function deficits and associated with premature mortality. Despite improvements in technology to protect coal miners, cases of PMF still occur in the United States at a disturbing rate.

Caplan syndrome (Chap. 380), first described in coal miners but subsequently in patients with silicosis, is the combination of pneumoconiotic nodules and seropositive rheumatoid arthritis. Silica has immunoadjuvant properties and is often present in anthracitic coal dust.

CHRONIC BERYLLIUM DISEASE

Beryllium is a lightweight metal with tensile strength, good electrical conductivity, and value in the control of nuclear reactions through its ability to quench neutrons. Although beryllium may produce an acute pneumonitis, it is far more commonly associated with a chronic granulomatous inflammatory disease that is similar to sarcoidosis (**Chap. 390**). Unless one inquires specifically about occupational exposures to beryllium in the manufacture of alloys, ceramics, or high-technology electronics in a patient with sarcoidosis, one may miss entirely the etiologic relationship to the occupational exposure. What distinguishes chronic beryllium disease (CBD) from sarcoidosis is evidence of a specific cell-mediated immune response (i.e., delayed hypersensitivity) to beryllium.

The test that usually provides this evidence is the beryllium lymphocyte proliferation test (BeLPT). The BeLPT compares the *in vitro* proliferation of lymphocytes from blood or bronchoalveolar lavage in the presence of beryllium salts with that of unstimulated cells. Proliferation is usually measured by lymphocyte uptake of radiolabeled thymidine.

Chest imaging findings are similar to those of sarcoidosis (nodules along septal lines) except that hilar adenopathy is somewhat less common. As with sarcoidosis, pulmonary function test results may show restrictive and/or obstructive ventilatory deficits and decreased diffusing capacity. With early disease, both chest imaging studies and pulmonary function tests may be normal. Fiberoptic bronchoscopy with transbronchial lung biopsy usually is required to make the diagnosis of CBD. In a beryllium-sensitized individual, the presence of noncaseating granulomas or monocytic infiltration in lung tissue establishes the diagnosis. Accumulation of beryllium-specific CD4+ T cells occurs in the granulomatous inflammation seen on lung biopsy. Susceptibility to CBD is highly associated with human leukocyte antigen DP (HLA-DP) alleles that have a glutamic acid in position 69 of the β chain.

OTHER METALS

Aluminum and titanium dioxide have been rarely associated with a sarcoid-like reaction in lung tissue. Exposure to dust containing tungsten carbide, also known as “hard metal,” may produce giant cell interstitial pneumonitis. Cobalt is a constituent of tungsten carbide and is the likely etiologic agent of both the interstitial pneumonitis and the occupational asthma that may occur. The most common exposures to tungsten carbide occur in tool and dye, saw blade, and drill bit manufacture. Diamond polishing may also involve exposure to cobalt dust. In patients with interstitial lung disease, one should always inquire about exposure to metal fumes and/or dusts. Especially when sarcoidosis appears to be the diagnosis, one should always consider possible CBD.

OTHER INORGANIC DUSTS

Most of the inorganic dusts discussed thus far are associated with the production of either dust macules or interstitial fibrotic changes in the lung. Other inorganic and organic dusts (see categories in Table 311-1), along with some of the dusts previously discussed, are associated with chronic mucus hypersecretion (chronic bronchitis), with or without reduction of expiratory flow rates. Cigarette smoking is the major cause of these conditions, and any effort to attribute some component of the disease to occupational and environmental exposures must take cigarette smoking into account. Most studies suggest an additive effect of dust exposure and smoking. The pattern of the irritant dust effect is similar to that of cigarette smoking, suggesting that small airway inflammation may be the initial site of pathologic response in those cases and continued exposure may lead to chronic bronchitis and COPD.

ORGANIC DUSTS

Some of the specific diseases associated with organic dusts are discussed in detail in the chapters on asthma (**Chap. 309**) and hypersensitivity pneumonitis (**Chap. 310**). Many of these diseases are named for the specific setting in which they are found, e.g., farmer’s lung, malt worker’s disease, and mushroom worker’s disease. Often the temporal relation of symptoms to exposure furnishes the best evidence for the

diagnosis. Three occupational exposures are singled out for discussion here because they affect the largest proportions of workers.

Cotton Dust (Byssinosis) Workers occupationally exposed to cotton dust (but also to flax, hemp, or jute dust) in the production of yarns for textiles and rope making are at risk for an asthma-like syndrome known as byssinosis. Exposure occurs throughout the manufacturing process but is most pronounced in the portions of the factory involved with the treatment of the cotton before spinning, i.e., blowing, mixing, and carding (straightening of fibers). The risk of byssinosis is associated with both cotton dust and endotoxin levels in the workplace environment.

Byssinosis is characterized clinically as occasional (early-stage) and then regular (late-stage) chest tightness toward the end of the first day of the workweek (“Monday chest tightness”). Exposed workers may show a significant drop in FEV₁ over the course of a Monday workshift. Initially the symptoms do not recur on subsequent days of the week. However, in 10–25% of workers, the disease may be progressive, with chest tightness recurring or persisting throughout the workweek. After >10 years of exposure, workers with recurrent symptoms are more likely to have an obstructive pattern on pulmonary function testing. The highest grades of impairment generally are seen in smokers.

Dust exposure can be reduced by the use of exhaust hoods, general increases in ventilation, and wetting procedures, but respiratory protective equipment may be required during certain operations. Regular surveillance of pulmonary function in cotton dust–exposed workers using spirometry before and after the workshift is required by OSHA. All workers with persistent symptoms or significantly reduced levels of pulmonary function should be moved to areas of lower risk of exposure.

Grain Dust Worldwide, many farmers and workers in grain storage facilities are exposed to grain dust. The presentation of obstructive airway disease in grain dust–exposed workers is virtually identical to the characteristic findings in cigarette smokers, i.e., persistent cough, mucus hypersecretion, wheeze and dyspnea on exertion, and reduced FEV₁ and FEV₁/FVC (forced vital capacity) ratio (**Chap. 306e**).

Dust concentrations in grain elevators vary greatly but can be >10,000 $\mu\text{g}/\text{m}^3$ with many particles in the respirable size range. The effect of grain dust exposure is additive to that of cigarette smoking, with ~50% of workers who smoke having symptoms. Smoking grain dust–exposed workers are more likely to have obstructive ventilatory deficits on pulmonary function testing. As in byssinosis, endotoxin may play a role in grain dust–induced chronic bronchitis and COPD.

Farmer’s Lung This condition results from exposure to moldy hay containing spores of thermophilic actinomycetes that produce a hypersensitivity pneumonitis (**Chap. 310**). A patient with acute farmer’s lung presents 4–8 h after exposure with fever, chills, malaise, cough, and dyspnea without wheezing. The history of exposure is obviously essential to distinguish this disease from influenza or pneumonia with similar symptoms. In the chronic form of the disease, the history of repeated attacks after similar exposure is important in differentiating this syndrome from other causes of patchy fibrosis (e.g., sarcoidosis).

A wide variety of other organic dusts are associated with the occurrence of hypersensitivity pneumonitis (**Chap. 310**). For patients who present with hypersensitivity pneumonitis, specific and careful inquiry about occupations, hobbies, and other home environmental exposures is necessary to uncover the source of the etiologic agent.

TOXIC CHEMICALS

Exposure to toxic chemicals affecting the lung generally involves gases and vapors. A common accident is one in which the victim is trapped in a confined space where the chemicals have accumulated to harmful levels. In addition to the specific toxic effects of the chemical, the victim often sustains considerable anoxia, which can play a dominant role in determining whether the individual survives.

Table 311-2 lists a variety of toxic agents that can produce acute and sometimes life-threatening reactions in the lung. All these agents in sufficient concentrations have been demonstrated, at least in animal studies, to affect the lower airways and disrupt alveolar architecture, either acutely or as a result of chronic exposure. Some of these agents may be generated acutely in the environment (see below).

TABLE 311-2 SELECTED COMMON TOXIC CHEMICAL AGENTS THAT AFFECT THE LUNG

Agent(s)	Selected Exposures	Acute Effects from High or Accidental Exposure	Chronic Effects from Relatively Low Exposure
Acid anhydrides	Manufacture of resin esters, polyester resins, thermoactivated adhesives	Nasal irritation, cough	Asthma, chronic bronchitis, hypersensitivity pneumonitis
Acid fumes: H ₂ SO ₄ , HNO ₃	Manufacture of fertilizers, chlorinated organic compounds, dyes, explosives, rubber products, metal etching, plastics	Mucous membrane irritation, followed by chemical pneumonitis 2–3 days later	Bronchitis and suggestion of mildly reduced pulmonary function in children with lifelong residential exposure to high levels
Acrolein and other aldehydes	By-product of burning plastics, woods, tobacco smoke	Mucous membrane irritant, decrease in lung function	Upper respiratory tract irritation
Ammonia	Refrigeration; petroleum refining; manufacture of fertilizers, explosives, plastics, and other chemicals	Same as for acid fumes, but bronchiectasis also has been reported	Upper respiratory tract irritation, chronic bronchitis
Cadmium fumes	Smelting, soldering, battery production	Mucous membrane irritant, acute respiratory distress syndrome (ARDS)	Chronic obstructive pulmonary disease (COPD)
Formaldehyde	Manufacture of resins, leathers, rubber, metals, and woods; laboratory workers, embalmers; emission from urethane foam insulation	Same as for acid fumes	Nasopharyngeal cancer
Halides and acid salts (Cl, Br, F)	Bleaching in pulp, paper, textile industry; manufacture of chemical compounds; synthetic rubber, plastics, disinfectant, rocket fuel, gasoline	Mucous membrane irritation, pulmonary edema; possible reduced forced vital capacity (FVC) 1–2 years after exposure	Upper respiratory tract irritation, epistaxis, tracheobronchitis
Hydrogen sulfide	By-product of many industrial processes, oil, other petroleum processes and storage	Increase in respiratory rate followed by respiratory arrest, lactic acidosis, pulmonary edema, death	Conjunctival irritation, chronic bronchitis, recurrent pneumonitis
Isocyanates (TDI, HDI, MDI)	Production of polyurethane foams, plastics, adhesives, surface coatings	Mucous membrane irritation, dyspnea, cough, wheeze, pulmonary edema	Upper respiratory tract irritation, cough, asthma, hypersensitivity pneumonitis, reduced lung function
Nitrogen dioxide	Silage, metal etching, explosives, rocket fuels, welding, by-product of burning fossil fuels	Cough, dyspnea, pulmonary edema may be delayed 4–12 h; possible result from acute exposure: bronchiolitis obliterans in 2–6 weeks	Emphysema in animals, ? chronic bronchitis, associated with reduced lung function in children with lifelong residential exposure
Ozone	Arc welding, flour bleaching, deodorizing, emissions from copying equipment, photochemical air pollutant	Mucous membrane irritant, pulmonary hemorrhage and edema, reduced pulmonary function transiently in children and adults, and increased hospitalization with exposure to summer haze	Excess cardiopulmonary mortality rates
Phosgene	Organic compound, metallurgy, volatilization of chlorine-containing compounds	Delayed onset of bronchiolitis and pulmonary edema	Chronic bronchitis
Sulfur dioxide	Manufacture of sulfuric acid, bleaches, coating of nonferrous metals, food processing, refrigerant, burning of fossil fuels, wood pulp industry	Mucous membrane irritant, epistaxis, bronchospasm (especially in people with asthma)	Chronic bronchitis

Abbreviations: HDI, hexamethylene diisocyanate; MDI, methylene diphenyl diisocyanate; TDI, toluene diisocyanate.

Firefighters and fire victims are at risk of *smoke inhalation*, an important cause of acute cardiorespiratory failure. Smoke inhalation kills more fire victims than does thermal injury. Carbon monoxide poisoning with resulting significant hypoxemia can be life-threatening (Chap. 473e). Synthetic materials (plastic, polyurethanes), when burned, may release a variety of other toxic agents (such as cyanide and hydrochloric acid), and this must be considered in evaluating smoke inhalation victims. Exposed victims may have some degree of lower respiratory tract inflammation and/or pulmonary edema.

Exposure to certain highly reactive, low-molecular-weight agents used in the manufacture of synthetic polymers, paints, and coatings (*diisocyanates* in polyurethanes, *aromatic amines* and *acid anhydrides* in epoxies) is associated with a high risk of occupational asthma. Although this occupational asthma manifests clinically as if sensitization has occurred, an IgE antibody-mediated mechanism is not necessarily involved. Hypersensitivity pneumonitis-like reactions also have been described in diisocyanate and acid anhydride-exposed workers.

Fluoropolymers such as Teflon, which at normal temperatures produce no reaction, become volatilized upon heating. The inhaled agents cause a characteristic syndrome of fever, chills, malaise, and occasionally mild wheezing, leading to the diagnosis of *polymer fume fever*. A similar self-limited, influenza-like syndrome—*metal fume fever*—results from acute exposure to fumes containing zinc oxide, typically from welding of galvanized steel. These inhalational fever syndromes may begin several hours after work and resolve within 24 h, only to return on repeated exposure.

Two other agents have been associated with potentially severe lung disease. Occupational exposure to nylon flock has been shown to induce a lymphocytic bronchiolitis, and workers exposed to diacetyl, which is used to provide “butter” flavor in the manufacture of microwave popcorn and other foods, have developed bronchiolitis obliterans (Chap. 315).

World Trade Center Disaster A consequence of the attack on the World Trade Center (WTC) on September 11, 2001, was relatively heavy exposure of a large number of firefighters and other rescue workers to the dust generated by the collapse of the buildings. Environmental monitoring and chemical characterization of WTC dust has revealed a wide variety of potentially toxic constituents, although much of the dust was pulverized cement. Possibly because of the high alkalinity of WTC dust, significant cough, wheeze, and phlegm production occurred among firefighters and cleanup crews. New cough and wheeze syndromes also occurred among local residents. Heavier exposure to WTC dust among New York City firefighters was associated with accelerated decline of lung function over the first year after the disaster. More recently, concerns have been raised about risk of interstitial lung disease, especially of a granulomatous nature.

OCCUPATIONAL RESPIRATORY CARCINOGENS

Exposures at work have been estimated to contribute to 10% of all lung cancer cases. In addition to asbestos, other agents either proven or suspected to be respiratory carcinogens include acrylonitrile, arsenic

compounds, beryllium, bis(chloromethyl) ether, chromium (hexavalent), formaldehyde (nasal), isopropanol (nasal sinuses), mustard gas, nickel carbonyl (nickel smelting), polycyclic aromatic hydrocarbons (coke oven emissions and diesel exhaust), secondhand tobacco smoke, silica (both mining and processing), talc (possible asbestos contamination in both mining and milling), vinyl chloride (sarcomas), wood (nasal cancer only), and uranium. Workers at risk of radiation-related lung cancer include not only those involved in mining or processing uranium but also those exposed in underground mining operations of other ores where radon daughters may be emitted from rock formations.

ASSESSMENT OF DISABILITY

Disability is the term used to describe the decreased ability to work due to the effects of a medical condition. Physicians are generally able to assess physiologic dysfunction, or *impairment*, but the rating of disability for compensation of loss of income also involves nonmedical factors such as the education and employability of the individual. The disability rating scheme differs with the compensation-granting agency. For example, the U.S. Social Security Administration requires that an individual be unable to do any work (i.e., *total* disability) before he or she will receive income replacement payments. Many state workers' compensation systems allow for payments for *partial* disability. In the Social Security scheme, no determination of cause is done, whereas work-relatedness must be established in workers' compensation systems.

For respiratory impairment rating, resting pulmonary function tests (spirometry and diffusing capacity) are used as the initial assessment tool, with cardiopulmonary exercise testing (to assess maximal oxygen consumption) used if the results of the resting tests do not correlate with the patient's symptoms. Methacholine challenge (to assess airway reactivity) can also be useful in patients with asthma who have normal spirometry when evaluated. Some compensation agencies (e.g., Social Security) have proscribed disability classification schemes based on pulmonary function test results. When no specific scheme is proscribed, the *Guidelines of the American Medical Association* should be used.

GENERAL ENVIRONMENTAL EXPOSURES

OUTDOOR AIR POLLUTION

In 1971, the U.S. government established national air quality standards for several pollutants believed to be responsible for excess cardiorespiratory diseases. Primary standards regulated by the U.S. Environmental Protection Agency (EPA) designed to protect the public health with an adequate margin of safety exist for sulfur dioxide, particulates matter, nitrogen dioxide, ozone, lead, and carbon monoxide. Standards for each of these pollutants are updated regularly through an extensive review process conducted by the EPA. (For details on current standards, go to <http://www.epa.gov/air/criteria.html>.)

Pollutants are generated from both stationary sources (power plants and industrial complexes) and mobile sources (motor vehicles), and none of the regulated pollutants occurs in isolation. Furthermore, pollutants may be changed by chemical reactions after being emitted. For example, sulfur dioxide and particulate matter emissions from a coal-fired power plant may react in air to produce acid sulfates and aerosols, which can be transported long distances in the atmosphere. Oxides of nitrogen and volatile organic compounds from automobile exhaust react with sunlight to produce ozone. Although originally thought to be confined to Los Angeles, photochemically derived pollution ("smog") is now known to be a problem throughout the United States and in many other countries. Both acute and chronic effects of these exposures have been documented in large population studies.

The symptoms and diseases associated with air pollution are the same as conditions commonly associated with cigarette smoking. In addition, decreased growth of lung function and asthma have been associated with chronic exposure to only modestly elevated levels of traffic-related gases and respirable particles. Multiple population-based time-series studies within cities have demonstrated excess health care utilization for asthma and other cardiopulmonary conditions as well as increased mortality rates. Cohort studies comparing cities that

have relatively high levels of particulate exposures with less polluted communities suggest excess morbidity and mortality rates from cardiopulmonary conditions in long-term residents of the former. The strong epidemiologic evidence that fine particulate matter is a risk factor for cardiovascular morbidity and mortality has prompted toxicologic investigations into the underlying mechanisms. The inhalation of fine particles from combustion sources probably generates oxidative stress followed by local injury and inflammation in the lungs that in turn lead to autonomic and systemic inflammatory responses that can induce endothelial dysfunction and/or injury. Recent research findings on the health effects of air pollutants has led to stricter U.S. ambient air quality standards for ozone, oxides of nitrogen, and particulate matter as well as greater emphasis on publicizing pollution alerts to encourage individuals with significant cardiopulmonary impairment to stay indoors during high-pollution episodes.

INDOOR EXPOSURES

Secondhand tobacco smoke (Chap. 470), radon gas, wood smoke, and other biologic agents generated indoors must be considered. Several studies have shown that the respirable particulate load in any household is directly proportional to the number of cigarette smokers living in that home. Increases in prevalence of respiratory illnesses, especially asthma, and reduced levels of pulmonary function measured with simple spirometry have been found in the children of smoking parents in a number of studies. Recent meta-analyses for lung cancer and cardiopulmonary diseases, combining data from multiple secondhand tobacco smoke epidemiologic studies, suggest an ~25% increase in relative risk for each condition, even after adjustment for major confounders.

Exposure to *radon gas* in homes is a risk factor for lung cancer. The main radon product (radon-222) is a gas that results from the decay series of uranium-238, with the immediate precursor being radium-226. The amount of radium in earth materials determines how much radon gas will be emitted. Levels associated with excess lung cancer risk may be present in as many as 10% of the houses in the United States. When smokers reside in the home, the problem is potentially greater, because the molecular size of radon particles allows them to attach readily to smoke particles that are inhaled. Fortunately, technology is available for assessing and reducing the level of exposure.

Other indoor exposures of concern are bioaerosols that contain antigenic material (fungi, cockroaches, dust mites, and pet danders) associated with an increased risk of atopy and asthma. Indoor chemical agents include strong cleaning agents (bleach, ammonia), formaldehyde, perfumes, pesticides, and oxides of nitrogen from gas appliances. Nonspecific responses associated with "tight-building syndrome," perhaps better termed "building-associated illness," in which no particular agent has been implicated, have included a wide variety of complaints, among them respiratory symptoms that are relieved only by avoiding exposure in the building in question. The degree to which "smells" and other sensory stimuli are involved in the triggering of potentially incapacitating psychological or physical responses has yet to be determined, and the long-term consequences of such environmental exposures are unknown.

GLOBAL CONSIDERATIONS



Indoor exposure to *biomass smoke* (wood, dung, crop residues, charcoal) is estimated to be responsible for >4% of worldwide disability-adjusted life-years (DALYs) lost, due to acute lower respiratory infections in children, COPD and lung cancer in women, and cardiovascular disease among men. This burden of disease places indoor exposure to biomass smoke as the leading environmental hazard for poor health and the third most important risk factor overall.

Almost one-half of the world's population uses biomass fuel for cooking, heating, or baking. This occurs predominantly in the rural areas of developing countries. Because many families burn biomass fuels in open stoves, which are highly inefficient, and inside homes with poor ventilation, women and young children are exposed on a daily basis to high levels of smoke. In these homes, 24-h mean levels

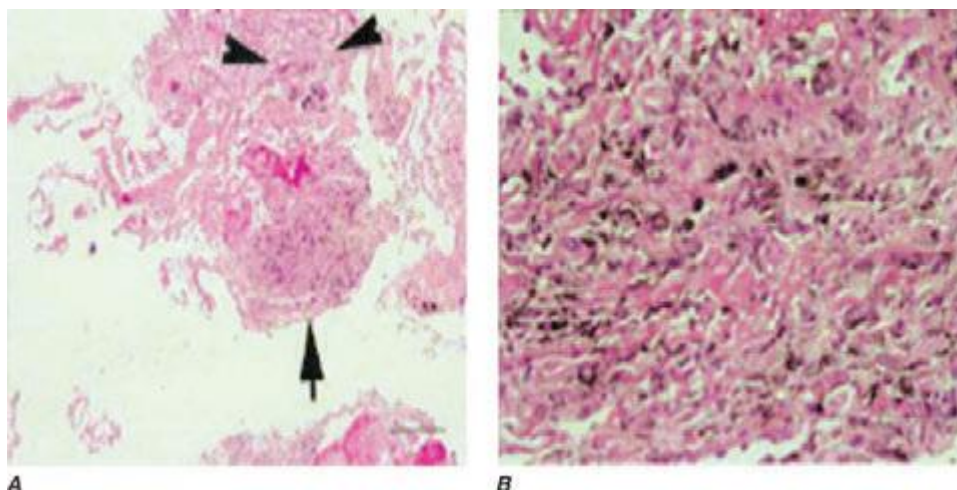


FIGURE 311-4 Histopathologic features of biomass smoke-induced interstitial lung disease. **A.** Anthracitic pigment is seen accumulating along alveolar septae (arrowheads) and within a pigmented dust macule (single arrow). **B.** A high-power photomicrograph contains a mixture of fibroblasts and carbon-laden macrophages.

of fine particulate matter, a component of biomass smoke, have been reported to be 2–30 times higher than the National Ambient Air Quality Standards set by the U.S. EPA.

Epidemiologic studies have consistently shown associations between exposure to biomass smoke and both chronic bronchitis and COPD, with odds ratios ranging between 3 and 10 and increasing with longer exposures. In addition to the common occupational exposure to biomass smoke of women in developing countries, men from such countries may be occupationally exposed. Because of increased migration to the United States from developing countries, clinicians need to be aware of the chronic respiratory effects of exposure to biomass smoke, which can include interstitial lung disease (Fig. 311-4). Evidence is beginning to emerge that improved stoves with chimneys can reduce biomass smoke-induced respiratory illness in both children and women.

312 Bronchiectasis

Rebecca M. Baron, Miriam Baron Barshak

Bronchiectasis refers to an irreversible airway dilation that involves the lung in either a focal or a diffuse manner and that classically has been categorized as cylindrical or tubular (the most common form), varicose, or cystic.

ETIOLOGY

Bronchiectasis can arise from infectious or noninfectious causes (Table 312-1). Clues to the underlying etiology are often provided by the pattern of lung involvement. *Focal bronchiectasis* refers to bronchiectatic changes in a localized area of the lung and can be a consequence of obstruction of the airway—either extrinsic (e.g., due to compression by adjacent lymphadenopathy or parenchymal tumor mass) or intrinsic (e.g., due to an airway tumor or aspirated foreign body, a scarred/stenotic airway, or bronchial atresia from congenital underdevelopment of the airway). *Diffuse bronchiectasis* is characterized by widespread bronchiectatic changes throughout the lung and often arises from an underlying systemic or infectious disease process.

More pronounced involvement of the upper lung fields is most common in cystic fibrosis (CF) and is also observed in postradiation fibrosis, corresponding to the lung region encompassed by the radiation port. Bronchiectasis with predominant involvement of the lower lung fields usually has its source in chronic recurrent aspiration (e.g., due to esophageal motility disorders like those in scleroderma), end-stage fibrotic lung disease (e.g., traction bronchiectasis from idiopathic

TABLE 312-1 MAJOR ETIOLOGIES OF BRONCHIECTASIS AND PROPOSED WORKUP

Pattern of Lung Involvement	Etiology by Category (Examples)	Workup
Focal	Obstruction (aspirated foreign body, tumor mass)	Chest imaging (chest x-ray and/or chest computed tomography); bronchoscopy
Diffuse	Infection (bacterial, nontuberculous mycobacterial)	Sputum Gram's stain/culture; stains/cultures for acid-fast bacilli and fungi. If no pathogen is identified, consider bronchoscopy with bronchoalveolar lavage.
	Immunodeficiency (hypogammaglobulinemia, HIV infection, bronchiolitis obliterans after lung transplantation)	Complete blood count with differential; immunoglobulin measurement; HIV testing
	Genetic causes (cystic fibrosis, Kartagener's syndrome, α_1 antitrypsin deficiency)	Measurement of chloride levels in sweat (for cystic fibrosis), α_1 antitrypsin levels; nasal or respiratory tract brush/biopsy (for dyskinetic/immotile cilia syndrome); genetic testing
	Autoimmune or rheumatologic causes (rheumatoid arthritis, Sjögren's syndrome, inflammatory bowel disease); immune-mediated disease (allergic bronchopulmonary aspergillosis)	Clinical examination with careful joint exam, serologic testing (e.g., for rheumatoid factor). Consider workup for allergic bronchopulmonary aspergillosis, especially in patients with refractory asthma. ^a
	Recurrent aspiration	Test of swallowing function and general neuromuscular strength
	Miscellaneous (yellow nail syndrome, traction bronchiectasis from postradiation fibrosis or idiopathic pulmonary fibrosis)	Guided by clinical condition
	Idiopathic	Exclusion of other causes

^aSkin testing for *Aspergillus* reactivity; measurement of serum precipitins for *Aspergillus*, serum IgE levels, serum eosinophils, etc.

pulmonary fibrosis), or recurrent immunodeficiency-associated infections (e.g., hypogammaglobulinemia). Bronchiectasis resulting from infection by nontuberculous mycobacteria (NTM), most commonly the *Mycobacterium avium-intracellulare* complex (MAC), often preferentially affects the midlung fields. Congenital causes of bronchiectasis

with predominant midlung field involvement include the dyskinetic/immotile cilia syndrome. Finally, predominant involvement of the central airways is reported in association with allergic bronchopulmonary aspergillosis (ABPA), in which an immune-mediated reaction to *Aspergillus* damages the bronchial wall. Congenital causes of central airway–predominant bronchiectasis resulting from cartilage deficiency include tracheobronchomegaly (Mounier-Kuhn syndrome) and Williams–Campbell syndrome.

In many cases, the etiology of bronchiectasis is not determined. In case series, as many as 25–50% of patients referred for bronchiectasis have idiopathic disease.

EPIDEMIOLOGY

The overall reported prevalence of bronchiectasis in the United States has recently increased, but the epidemiology of bronchiectasis varies greatly with the underlying etiology. For example, patients born with CF often develop significant clinical bronchiectasis in late adolescence or early adulthood, although atypical presentations of CF in adults in their thirties and forties are also possible. In contrast, bronchiectasis resulting from MAC infection classically affects nonsmoking women >50 years of age. In general, the incidence of bronchiectasis increases with age. Bronchiectasis is more common among women than among men.



In areas where tuberculosis is prevalent, bronchiectasis more frequently occurs as a sequela of granulomatous infection.

Focal bronchiectasis can arise from extrinsic compression of the airway by enlarged granulomatous lymph nodes and/or from development of intrinsic obstruction as a result of erosion of a calcified lymph node through the airway wall (e.g., broncholithiasis). Especially in reactivated tuberculosis, parenchymal destruction from infection can result in areas of more diffuse bronchiectasis. Apart from cases associated with tuberculosis, an increased incidence of non-CF bronchiectasis with an unclear underlying mechanism has been reported as a significant problem in developing nations. It has been suggested that the high incidence of malnutrition in certain areas may predispose to immune dysfunction and development of bronchiectasis.

PATHOGENESIS AND PATHOLOGY

The most widely cited mechanism of infectious bronchiectasis is the “vicious cycle hypothesis,” in which susceptibility to infection and poor mucociliary clearance result in microbial colonization of the bronchial tree. Some organisms, such as *Pseudomonas aeruginosa*, exhibit a particular propensity for colonizing damaged airways and evading host defense mechanisms. Impaired mucociliary clearance can result from inherited conditions such as CF or dyskinetic cilia syndrome, and it has been proposed that a single severe infection (e.g., pneumonia caused by *Bordetella pertussis* or *Mycoplasma pneumoniae*) can result in significant airway damage and poor secretion clearance. The presence of the microbes incites continued chronic inflammation, with consequent damage to the airway wall, continued impairment of secretion and microbial clearance, and ongoing propagation of the infectious/inflammatory cycle. Moreover, it has been proposed that mediators released directly from bacteria can interfere with mucociliary clearance.

Classic studies of the pathology of bronchiectasis from the 1950s demonstrated significant small-airway wall inflammation and larger-airway wall destruction as well as dilation, with loss of elastin, smooth muscle, and cartilage. It has been proposed that inflammatory cells in the small airways release proteases and other mediators, such as reactive oxygen species and proinflammatory cytokines, that damage the larger-airway walls. Furthermore, the ongoing inflammatory process in the smaller airways results in airflow obstruction. It is thought that antiproteases, such as α_1 antitrypsin, play an important role in neutralizing the damaging effects of neutrophil elastase and in enhancing bacterial killing. Bronchiectasis and emphysema have been observed in patients with α_1 antitrypsin deficiency.

Proposed mechanisms for noninfectious bronchiectasis include immune-mediated reactions that damage the bronchial wall (e.g., those associated with systemic autoimmune conditions such as Sjögren’s

syndrome and rheumatoid arthritis). *Traction bronchiectasis* refers to dilated airways arising from parenchymal distortion as a result of lung fibrosis (e.g., postradiation fibrosis or idiopathic pulmonary fibrosis).

CLINICAL MANIFESTATIONS

The most common clinical presentation is a persistent productive cough with ongoing production of thick, tenacious sputum. Physical findings often include crackles and wheezing on lung auscultation, and some patients with bronchiectasis exhibit clubbing of the digits. Mild to moderate airflow obstruction is often detected on pulmonary function tests, overlapping with that seen at presentation with other conditions, such as chronic obstructive pulmonary disease (COPD). Acute exacerbations of bronchiectasis are usually characterized by changes in the nature of sputum production, with increased volume and purulence. However, typical signs and symptoms of lung infection, such as fever and new infiltrates, may not be present.

DIAGNOSIS

The diagnosis is usually based on presentation with a persistent chronic cough and sputum production accompanied by consistent radiographic features. Although chest radiographs lack sensitivity, the presence of “tram tracks” indicating dilated airways is consistent with bronchiectasis. Chest computed tomography (CT) is more specific for bronchiectasis and is the imaging modality of choice for confirming the diagnosis. CT findings include airway dilation (detected as parallel “tram tracks” or as the “signet-ring sign”—a cross-sectional area of the airway with a diameter at least 1.5 times that of the adjacent vessel), lack of bronchial tapering (including the presence of tubular structures within 1 cm from the pleural surface), bronchial wall thickening in dilated airways, inspissated secretions (e.g., the “tree-in-bud” pattern), or cysts emanating from the bronchial wall (especially pronounced in cystic bronchiectasis; Fig. 312-1).

APPROACH TO THE PATIENT: Bronchiectasis

The evaluation of a patient with bronchiectasis entails elicitation of a clinical history, chest imaging, and a workup to determine the underlying etiology. Evaluation of focal bronchiectasis almost always requires bronchoscopy to exclude airway obstruction by an underlying mass or foreign body. A workup for diffuse bronchiectasis includes analysis for the major etiologies (Table 312-1), with an initial focus on excluding CF. Pulmonary function testing is an important component of a functional assessment of the patient.



FIGURE 312-1 Representative chest computed tomography (CT) image of severe bronchiectasis. This patient’s CT demonstrates many severely dilated airways, seen both longitudinally (arrowhead) and in cross-section (arrow).

Treatment of infectious bronchiectasis is directed at the control of active infection and improvements in secretion clearance and bronchial hygiene so as to decrease the microbial load within the airways and minimize the risk of repeated infections.

ANTIBIOTIC TREATMENT

Antibiotics targeting the causative or presumptive pathogen (with *Haemophilus influenzae* and *P. aeruginosa* isolated commonly) should be administered in acute exacerbations, usually for a minimum of 7–10 days and perhaps for as long as 14 days. Decisions about treatment of NTM infection can be difficult, given that these organisms can be colonizers as well as pathogens and the prolonged treatment course often is not well tolerated. Consensus guidelines have advised that diagnostic criteria for true clinical infection with NTM should be considered in patients with symptoms and radiographic findings of lung disease who have at least two sputum samples positive on culture; at least one bronchoalveolar lavage (BAL) fluid sample positive on culture; a biopsy sample displaying histopathologic features of NTM infection (e.g., granuloma or a positive stain for acid-fast bacilli) along with one positive sputum culture; or a pleural fluid sample (or a sample from another sterile extrapulmonary site) positive on culture. MAC strains are the most common NTM pathogens, and the recommended regimen for HIV-negative patients includes a macrolide combined with rifampin and ethambutol. Consensus guidelines also recommend macrolide susceptibility testing for clinically significant MAC isolates.

BRONCHIAL HYGIENE

The numerous approaches used to enhance secretion clearance in bronchiectasis include hydration and mucolytic administration, aerosolization of bronchodilators and hyperosmolar agents (e.g., hypertonic saline), and chest physiotherapy (e.g., postural drainage, traditional mechanical chest percussion via hand clapping to the chest, or use of devices such as an oscillatory positive expiratory pressure flutter valve or a high-frequency chest wall oscillation vest). Pulmonary rehabilitation and a regular exercise program may assist with secretion clearance as well as with other aspects of bronchiectasis, including improved exercise capacity and quality of life. The mucolytic dornase (DNase) is recommended routinely in CF-related bronchiectasis but not in non-CF bronchiectasis, given concerns about lack of efficacy and potential harm in the non-CF population.

ANTI-INFLAMMATORY THERAPY

It has been proposed that control of the inflammatory response may be of benefit in bronchiectasis, and relatively small-scale trials have yielded evidence of alleviated dyspnea, decreased need for inhaled β -agonists, and reduced sputum production with inhaled glucocorticoids. However, no significant differences in lung function or bronchiectasis exacerbation rates have been observed. Risks of immunosuppression and adrenal suppression must be carefully considered with use of anti-inflammatory therapy in infectious bronchiectasis. Nevertheless, administration of oral/systemic glucocorticoids may be important in treatment of bronchiectasis due to certain etiologies, such as ABPA, or of noninfectious bronchiectasis due to underlying conditions, especially that in which an autoimmune condition is believed to be active (e.g., rheumatoid arthritis or Sjögren's syndrome). Patients with ABPA may also benefit from a prolonged course of treatment with the oral antifungal agent itraconazole.

REFRACTORY CASES

In select cases, surgery can be considered, with resection of a focal area of suppuration. In advanced cases, lung transplantation can be considered.

COMPLICATIONS

In more severe cases of infectious bronchiectasis, recurrent infections and repeated courses of antibiotics can lead to microbial resistance to antibiotics. In certain cases, combinations of antibiotics that have their own independent toxicity profiles may be necessary to treat resistant organisms.

Recurrent infections can result in injury to superficial mucosal vessels, with bleeding and, in severe cases, life-threatening hemoptysis. Management of massive hemoptysis usually requires intubation to stabilize the patient, identification of the source of bleeding, and protection of the nonbleeding lung. Control of bleeding often necessitates bronchial artery embolization and, in severe cases, surgery.

PROGNOSIS

Outcomes of bronchiectasis can vary widely with the underlying etiology and may also be influenced by the frequency of exacerbations and (in infectious cases) the specific pathogens involved. In one study, the decline of lung function in patients with non-CF bronchiectasis was similar to that in patients with COPD, with the forced expiratory volume in 1 s (FEV₁) declining by 50–55 mL per year as opposed to 20–30 mL per year for healthy controls.

PREVENTION

Reversal of an underlying immunodeficient state (e.g., by administration of gamma globulin for immunoglobulin-deficient patients) and vaccination of patients with chronic respiratory conditions (e.g., influenza and pneumococcal vaccines) can decrease the risk of recurrent infections. Patients who smoke should be counseled about smoking cessation.

After resolution of an acute infection in patients with recurrences (e.g., ≥ 3 episodes per year), the use of suppressive antibiotics to minimize the microbial load and reduce the frequency of exacerbations has been proposed, although there is less consensus with regard to this approach in non-CF-associated bronchiectasis than in patients with CF-related bronchiectasis. Possible suppressive treatments include (1) administration of an oral antibiotic (e.g., ciprofloxacin) daily for 1–2 weeks per month; (2) use of a rotating schedule of oral antibiotics (to minimize the risk of development of drug resistance); (3) administration of a macrolide antibiotic (see below) daily or three times per week (with mechanisms of possible benefit related to non-antimicrobial properties, such as anti-inflammatory effects and reduction of gram-negative bacillary biofilms); (4) inhalation of aerosolized antibiotics (e.g., tobramycin inhalation solution) by select patients on a rotating schedule (e.g., 30 days on, 30 days off), with the goal of decreasing the microbial load without eliciting the side effects of systemic drug administration; and (5) intermittent administration of IV antibiotics (e.g., “clean-outs”) for patients with more severe bronchiectasis and/or resistant pathogens. In relation to macrolide therapy (point 3 above), a number of double-blind, placebo-controlled, randomized trials have recently been published in non-CF bronchiectasis and support a benefit of long-term macrolides (6–12 months of azithromycin or erythromycin) in decreasing rates of bronchiectasis exacerbation, mucus production, and decline in lung function. However, two of these studies also reported increased macrolide resistance in commensal pathogens, dampening enthusiasm for universal use of macrolides in this setting and raising the question of whether there might be select non-CF bronchiectasis patients with higher morbidity for whom benefits of long-term macrolides might outweigh the risks of emergence of antibiotic resistance. In particular, development of macrolide-resistant NTM is a significant concern, making treatment of that pathogen much more difficult. Therefore, it is advised to rule out NTM infection before chronic macrolide therapy is considered.

In addition, ongoing consistent attention to bronchial hygiene can promote secretion clearance and decrease the microbial load in the airways.

CLINICAL FEATURES

Cystic fibrosis (CF) is an autosomal recessive exocrinopathy affecting multiple epithelial tissues. The gene product responsible for CF (the cystic fibrosis transmembrane conductance regulator [CFTR]) serves as an anion channel in the apical (luminal) plasma membranes of epithelial cells and regulates volume and composition of exocrine secretion. An increasingly sophisticated understanding of CFTR molecular genetics and membrane protein biochemistry has facilitated CF drug discovery, with a number of new agents advancing through the clinical testing phase.

Respiratory Manifestations The major morbidity and mortality associated with CF is attributable to respiratory compromise, characterized by copious hyperviscous and adherent pulmonary secretions that obstruct small and medium-sized airways. CF airway secretions are exceedingly difficult to clear, and a complex bacterial flora that includes *Staphylococcus aureus*, *Haemophilus influenzae*, and *Pseudomonas aeruginosa* (among other pathogens) is routinely cultured from CF sputum. Robust pulmonary inflammation in the setting of inspissated mucus and chronic bacterial infection leads to collateral tissue injury and further aggravates respiratory decline. Organisms such as *P. aeruginosa* exhibit a stereotypic mode of pathogenesis; a sentinel and early colonization event often engenders lifelong pulmonary infection by the same genetic strain. Over the course of many years, *P. aeruginosa* evolves in CF lungs to adopt a mucoid phenotype (attributable to release of alginate exoproduct) that confers selective advantage for the pathogen and poor prognosis for the host. Infection with other bacterial organisms such as *Burkholderia cepacia* also indicates a less favorable pulmonary outlook. Strategies to eradicate organisms such as *P. aeruginosa* early in the pathogenesis cascade have been successful and are thought to improve prognosis significantly if sustained.

Pancreatic Findings The complete name of the disease, *cystic fibrosis of the pancreas*, refers to profound tissue destruction of the exocrine pancreas, with fibrotic scarring and/or fatty replacement, cyst proliferation, loss of acinar tissue, and ablation of normal pancreatic architecture. As in the lung, tenacious exocrine secretions (sometimes termed *concretions*) obstruct pancreatic ducts and impair production and flow of digestive enzymes to the duodenum. The sequelae of exocrine pancreatic insufficiency include chronic malabsorption, poor growth, fat-soluble vitamin insufficiency, high levels of serum immunoreactive trypsinogen (a diagnostic test used in newborn screening), and loss of pancreatic islet cell mass. CF-related diabetes mellitus is a manifestation in over 30% of adults with the disease and is likely multifactorial in nature (attributable to progressive destruction of the endocrine pancreas, insulin resistance due to stress hormones, and other factors).

Other Organ System Damage As in CF lung and pancreas, thick and tenacious secretions compromise numerous other exocrine tissues. Obstruction of intrahepatic bile ducts and parenchymal fibrosis are commonly observed in pathologic specimens, with multilobular cirrhosis in 4–15% of patients with CF and significant hepatic insufficiency as a resulting manifestation among adults. Contents of the intestinal lumen are often difficult to excrete, leading to meconium ileus (a presentation in approximately 10–20% of newborns with CF) or distal intestinal obstructive syndrome in older individuals. Men typically exhibit complete involution of the vas deferens and infertility (despite functioning spermatogenesis), and approximately 99% of males with CF are infertile. The etiology of this dramatic anatomic defect in the male genitourinary system is not understood but may represent a developmental abnormality secondary to secretory obstruction of the vas. Abnormalities of female reproductive tract secretions are likely contributors to an increased incidence of infertility among women with CF. Radiographic evidence of sinusitis occurs in most CF patients and is associated with pathogens similar to those recovered

from lower airways, suggesting that the sinus may serve as a reservoir for bacterial seeding.

PATHOGENESIS

Cystic Fibrosis Transmembrane Conductance Regulator (CFTR) CFTR is an integral membrane protein that functions as an epithelial anion channel. The ~1480-amino-acid molecule encodes a passive conduit for chloride and bicarbonate transport across plasma membranes of epithelial tissues, with direction of ion flow dependent on the electrochemical driving force. Gating of CFTR involves conformational cycling between an open and closed configuration and is augmented by hydrolysis of adenosine triphosphate (ATP). Anion flux mediated by CFTR does not involve active transport against a concentration gradient but utilizes the energy provided from ATP hydrolysis as a central feature of ion channel mechanochemistry and gating.

CFTR is situated in the apical plasma membranes of acinar and other epithelial cells where it regulates the amount and composition of secretion by exocrine glands. In numerous epithelia, chloride and bicarbonate release is followed passively by the flow of water, allowing for mobilization and clearance of exocrine products. Along respiratory mucosa, CFTR is necessary to provide sufficient depth of the periciliary fluid layer (PCL), allowing normal ciliary extension and mucociliary transport. CFTR-deficient airway cells exhibit depleted PCL, causing ciliary collapse and failure to clear overlying mucus (**Video 313-1**). In airway submucosal glands, CFTR is highly expressed in acini and may participate both in the formation of mucus and extrusion of glandular secretion onto the airway surface (**Fig. 313-1**). In other exocrine glands characterized by abrogated mucus transport (e.g., pancreatic acini and ducts, bile canaliculi, intestinal lumen), similar pathogenic mechanisms have been implicated. In these tissues, a driving force for apical chloride and/or bicarbonate secretion is believed to promote CFTR-mediated fluid and electrolyte release into the lumen, which confers proper rheology of mucins and other exocrine products. Failure of this mechanism disrupts normal hydration and transport of glandular secretion and is widely viewed as a proximate cause of ductular obstruction, with concomitant tissue injury.

Pulmonary Inflammation and Remodeling The CF airway is characterized by an aggressive, unrelenting, neutrophilic inflammatory response with release of proteases and oxidants leading to airway remodeling and bronchiectasis. Intense pulmonary inflammation is largely driven by chronic respiratory infection. Macrophages resident in CF lungs augment elaboration of proinflammatory cytokines, which contribute to innate and adaptive immune reactivity. CFTR-dependent abnormalities of airway surface fluid composition (e.g., pH) have been reported as contributors to impaired bacterial killing in CF lungs. The role of CFTR as a direct mediator of inflammatory responsiveness and/or pulmonary remodeling represents an important and topical area of investigation.

MOLECULAR GENETICS

DNA sequencing of *CFTR* from patients (and others) worldwide has revealed almost 2000 allelic variants; however, only about 10% of these have been well-characterized as disease-causing mutations. Distinguishing the single nucleotide transversions or other polymorphisms with causal relevance often presents a significant challenge. The CFTR2 resource (www.cftr2.org/) delineates gene variants with a clear etiologic role.

CFTR defects known to elicit disease are often categorized based on molecular mechanism. For example, the common F508del mutation (nomenclature denotes omission of a single phenylalanine residue [F] at CFTR position 508) leads to a folding abnormality recognized by cellular quality control pathways. CFTR encoding F508del retains partial ion channel function, but protein maturation is arrested in the endoplasmic reticulum, and CFTR fails to arrive at the plasma membrane. Instead, F508del CFTR is misrouted and undergoes endoplasmic reticulum-associated degradation via the proteasome. CFTR mutations that disrupt protein maturation are termed class II defects and are by far the most common genetic abnormalities. F508del alone accounts for ~70% of defective *CFTR* alleles in the United States, where

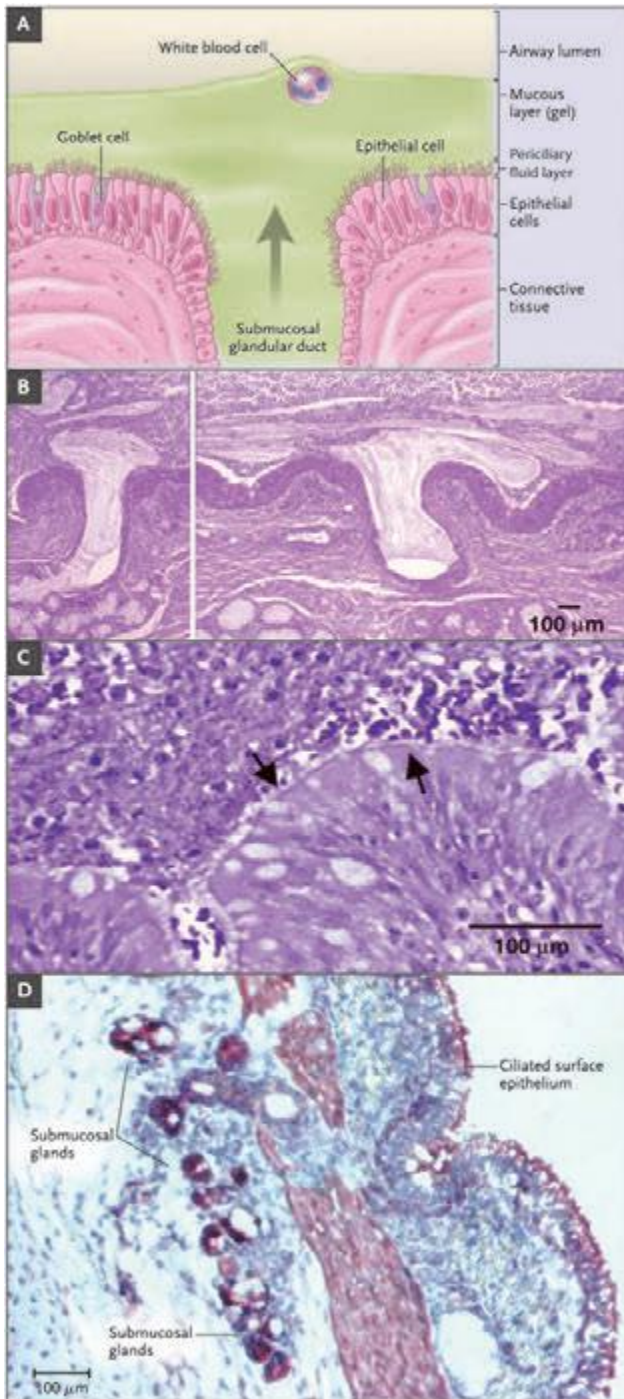


FIGURE 313-1 Extrusion of mucus secretion onto the epithelial surface of airways in cystic fibrosis. **A.** Schematic of the surface epithelium and supporting glandular structure of the human airway. **B.** The submucosal glands of a patient with cystic fibrosis are filled with mucus, and mucopurulent debris overlies the airway surfaces, essentially burying the epithelium. **C.** A higher magnification view of a mucus plug tightly adhering to the airway surface, with arrows indicating the interface between infected and inflamed secretions and the underlying epithelium to which the secretions adhere. (Both **B** and **C** were stained with hematoxylin and eosin, with the colors modified to highlight structures.) Infected secretions obstruct airways and, over time, dramatically disrupt the normal architecture of the lung. **D.** CFTR is expressed in surface epithelium and serous cells at the base of submucosal glands in a porcine lung sample, as shown by the dark staining, signifying binding by CFTR antibodies to epithelial structures (aminoethylcarbazole detection of horseradish peroxidase with hematoxylin counterstain). (From SM Rowe, S Miller, EJ Sorscher: *N Engl J Med* 352:1992, 2005.)

approximately 90% of individuals with CF carry at least one F508del mutation.

Other gene defects include CFTR ion channels properly trafficked to the apical cell surface but unable to open and/or gate. Such channel proteins include G551D (a glycine to aspartic acid replacement at CFTR position 551), which leads to an inability to transport Cl^- or HCO_3^- in the presence of ATP (a class III abnormality). Individuals with at least one G551D allele represent 4–5% of CF patients in North America. CFTR nonsense alleles such as G542X, R553X, and W1282X (premature termination codon replaces glycine, arginine, or tryptophan at positions 542, 553, or 1282, respectively) are among the common class I defects, in addition to large deletions or other major disruptions of the gene. The W1282X mutation, for example, is prevalent among individuals of Ashkenazi descent and is a predominant CF genotype in Israel. Additional categories of CFTR mutation include defects in the ion channel pore (class IV), RNA splicing (class V), and increased plasma membrane turnover (class VI) (Fig. 313-2).

DIAGNOSIS

The diagnosis of CF is based in part on clinical symptoms, family history, or positive newborn screening. CFTR mutation analysis together with sweat electrolyte measurements represent cardinal diagnostic tests. DNA-based evaluation typically surveys numerous disease-associated mutations; panels that identify 20–80 gene defects are

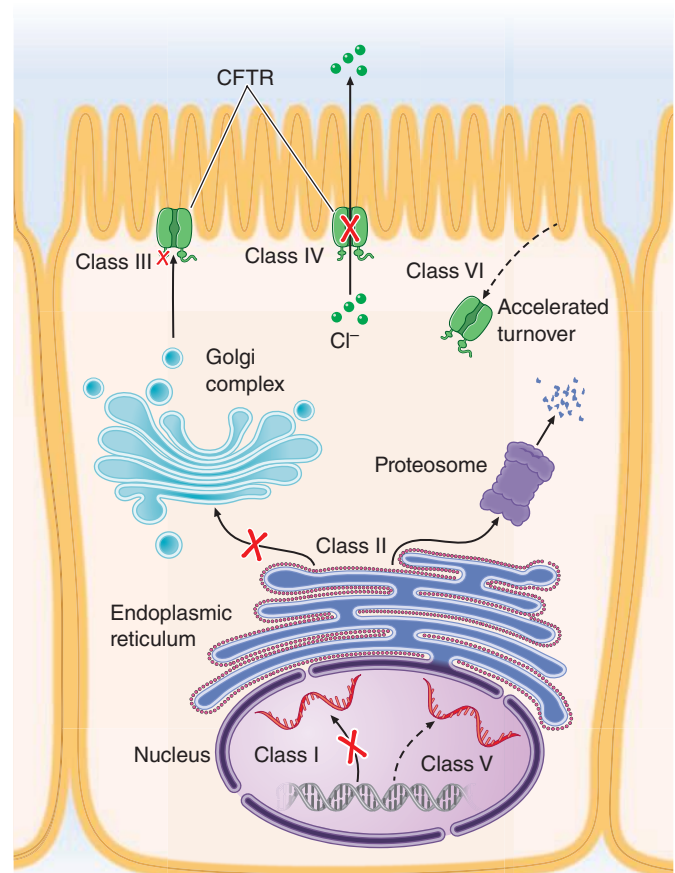


FIGURE 313-2 Categories of CFTR mutations. Classes of defects in the CFTR gene include the absence of synthesis (class I); defective protein maturation and premature degradation (class II); disordered gating/regulation, such as diminished adenosine triphosphate (ATP) binding and hydrolysis (class III); defective conductance through the ion channel pore (class IV); a reduced number of CFTR transcripts due to a promoter or splicing abnormality (class V); and accelerated turnover from the cell surface (class VI). (From SM Rowe, S Miller, EJ Sorscher: *N Engl J Med* 352:1992, 2005.)

available through commercial sources. For difficult cases, complete *CFTR* exonic sequencing together with analysis of splice junctions and key regulatory elements can be obtained. Sweat electrolytes following pilocarpine iontophoresis comprise an invaluable diagnostic measurement, with levels of chloride markedly elevated in CF compared to non-CF individuals. The sweat test result is highly specific and served as the mainstay of diagnosis for many decades prior to availability of *CFTR* genotyping. Notably, hyperviscosity of eccrine sweat is not a clinical feature of the disease. Sweat ducts function to reabsorb chloride from a primary sweat secretion produced by the glandular coil. Malfunction of *CFTR* leads to diminished chloride uptake from the ductular lumen, and sweat emerges on the skin with markedly elevated levels of chloride. For the unusual situation in which both *CFTR* genotype and sweat electrolytes are inconclusive, *in vivo* measurement of ion transport across the nasal airways can serve as a specific test for CF and is used by a number of referral centers. For example, elevated (sodium-dependent) transepithelial charge separation across airway epithelial tissue and failure of isoproterenol-dependent chloride secretion (via *CFTR*) represent bioelectric findings highly specific for the disease. Measurements of *CFTR* activity in excised rectal mucosal biopsies can also be obtained.

COMPLEXITY OF A CF PHENOTYPE

CF classically presents in childhood with chronic productive cough, malabsorption including steatorrhea, and failure to thrive. The disease is most common among whites (~1 in 3300 live births) and much less frequent among African-American (~1 in 15,000) or Asian populations (~1 in 33,000). Several “severe” defects that impair *CFTR* activity (including F508del, G551D, and truncation alleles) are predictive of pancreatic insufficiency, which is clinically evident in 80–90% of individuals with CF. These few specific genotype-phenotype correlations notwithstanding, genotype is, in general, a poor predictor of overall respiratory prognosis.

A spectrum of *CFTR*-related diseases with features resembling classic CF has been well described. In addition to multiorgan involvement, *forme frustes*, such as isolated congenital bilateral absence of the vas deferens or pancreatitis (without other organ system findings), are strongly associated with *CFTR* mutations in at least one allele. Although CF is a classic monogenic disease, the importance of non-*CFTR* gene modifiers and proteins that regulate ion flux, inflammatory pathways, and airway remodeling has been increasingly appreciated as influencing clinical course. For example, the magnitude of transepithelial sodium reabsorption in CF airways, which helps control periciliary fluid depth and composition, is strongly influenced by *CFTR* and represents a molecular target for disease intervention.

THERAPEUTICS DIRECTED TOWARD CF SEQUELAE

Standard care for outpatients with CF is intensive, with regimens that include exogenous pancreatic enzymes taken with meals, nutritional supplementation, anti-inflammatory medication, bronchodilators, and chronic or periodic administration of oral or aerosolized antibiotics (e.g., as maintenance therapy for patients with *P. aeruginosa*). Recombinant DNase aerosols (degraded DNA strands that contribute to mucus viscosity) and nebulized hypertonic saline (serves to augment PCL depth, activate mucociliary clearance, and mobilize inspissated airway secretions) are administered routinely. Chest physiotherapy several times each day is a standard means to promote clearance of airway mucus. Among older individuals with CF, malabsorption, chronic inflammation, and endocrine abnormalities can lead to poor bone mineralization, requiring treatment with vitamin D, calcium, and other measures. The time, complexity, and expense of home care are considerable and take a significant toll on patients and their families.

Severe respiratory exacerbation is commonly managed by hospital admission for frequent chest physiotherapy and parenteral antibiotics directed against serious (and often multiply resistant) bacterial pathogens. Aggressive intervention in this setting can restore a large component of lung function, but ongoing and cumulative loss of pulmonary reserve reflects the natural history of the disease. Poor prognostic indicators such as sputum culture containing *B. cepacia*, mucoid

P. aeruginosa, or atypical mycobacteria are rigorously monitored in the CF patient population. An increasing incidence of methicillin-resistant *S. aureus* has also been observed, although the clinical significance of this finding has not been fully elucidated. Typical inpatient antibiotic coverage includes combination drug therapy with an aminoglycoside and β -lactam for up to 14 days. Maximal improvement in lung function is often achieved by 8–10 days in this setting. Many families elect parenteral antibiotic treatment at home, and additional studies are needed to evaluate specific drug combinations, duration of therapy, and home versus inpatient management. Other CF respiratory sequelae that may require hospitalization include hemoptysis and pneumothorax. Hypersensitivity to *Aspergillus* (allergic bronchopulmonary aspergillosis) occurs in approximately 5% of individuals with the disease and should be suspected in the absence of a response to conventional treatment.

Lung transplantation remains a viable therapeutic option in the setting of end-stage CF pulmonary failure, with 5-year postoperative survival rates on the order of 50–60%. Determining the optimal timing for surgery presents a substantial challenge, particularly because overall prognosis for individuals with severe lung disease is sometimes difficult to predict, and mortality associated with transplantation is significant (1-year survival rates of approximately 80%). Forced expiratory volume in 1 s (FEV_1) measurements less than 30% predicted, together with an assortment of other clinical features, are often used as thresholds for entry onto transplantation lists, although waiting periods for healthy donor lungs can be quite protracted. Based on clinical outcome and limited access to healthy donor lungs, many CF patients and their families do not pursue this option.

CFTR MODULATION

Potiation of Mutant *CFTR* Gating A massive effort directed toward high-throughput drug analysis of large compound libraries (containing millions of individual agents) has identified novel and promising approaches to CF therapy. The approved compound ivacaftor, for example, robustly potentiates *CFTR* channel opening and stimulates ion transport. Ivacaftor overcomes the G551D *CFTR* gating defect, and individuals carrying this mutation exhibit dramatic improvement in lung function, weight gain, and other clinical parameters after only a few weeks of oral therapy. Remarkably, sweat chloride values are significantly improved with this treatment in patients with G551D *CFTR*. No clinical intervention of any sort has previously been shown to normalize the CF sweat chloride abnormality. Long-term studies of the drug in patients with G551D *CFTR* are ongoing. Ivacaftor has been viewed as the harbinger of a new era for CF therapeutics directed at treating the most fundamental causes of the disease.

Correction of the F508del Processing Abnormality Advancement of new drugs that address specific *CFTR* defects in protein folding and maturation has been bolstered by clinical studies of F508del rescue in combination with ivacaftor. So-called “corrector” molecules (as distinct from *CFTR* gating “potentiators” such as ivacaftor) discovered through compound library screening are suitable for promoting cell surface localization of the F508del protein. Significant improvement in pulmonary function of F508del homozygous individuals has been achieved with potentiator/corrector combination therapy in early clinical trials, and several candidate molecules are under evaluation.

Personalized Molecular Therapies The advent of modulators with robust clinical impact has engendered new optimism regarding care of patients with CF. It is clear that future interventions will be tailored to specific genotypic abnormalities. Drug screening campaigns and other research programs have identified agents capable of suppressing *CFTR* nonsense alleles, augmenting potentiator activity, and promoting F508del correction. Efforts to apply these compounds in a fashion that will benefit CF subjects carrying a single copy of F508del (i.e., with a distinct or unusual *CFTR* mutation on the second allele) comprise an essential priority for the future. Progress in CF drug discovery is emblematic of what might be accomplished in other refractory genetic diseases using an approach grounded in molecular mechanism and unbiased compound library screening.

As a direct result of advances in basic research, new therapies have transformed CF from a disease typically leading to death in early childhood to a condition with frequent survival well into the fourth decade of life. It has also become increasingly clear that specified approaches to patient management can have an impact on overall prognosis. For example, standardization of clinical intervention throughout the United States has led to remarkable benefit among the CF population. Well-defined measures for outpatient care are now established, including thresholds for hospital admission, antibiotic regimens, nutritional guidelines, periodicity of diagnostic tests, and other clinical parameters. These therapeutic recommendations have become standardized throughout approximately 110 specialized CF care centers and 55 affiliated programs. The initiative has improved endpoints such as weight gain, body mass index, and pulmonary function. Information regarding standardized protocols for CF therapy can be accessed at (www.cff.org/treatments/cfcareguidelines/) or through a number of excellent reviews.

Newborn screening for CF is now universal throughout the United States, most of the Canadian provinces, Australia, New Zealand, and much of Europe, and will facilitate early CF intervention. Based on data indicating that early nutritional and other therapies can be beneficial, newborn diagnosis is expected to significantly promote health in the CF population. Export of quality control measures and novel therapeutics worldwide has become an increasing imperative. For example, median survival of individuals with CF is less than 20 years in much of Central and South America (compared to ~40 years in the United States and Canada), and efforts to apply state-of-the-art management to underdiagnosed and underserved CF patient populations are expected to improve outcome and mitigate CF health disparities in the future.

VIDEO 313-1 Initial video sequences describe establishment of the normal periciliary fluid layer bathing the surface airway epithelium, with spheres representing chloride and bicarbonate ions secreted through CFTR and across the apical (mucosal) respiratory surface. Later video sequences depict failure of CFTR anion transport and resulting depletion of the periciliary layer, “plastering” of cilia against the mucosal surface, and accumulation of mucus in the airway with resulting bacterial infection. (Video courtesy of the Cystic Fibrosis Foundation.)

314 Chronic Obstructive Pulmonary Disease

John J. Reilly, Jr., Edwin K. Silverman, Steven D. Shapiro

Chronic obstructive pulmonary disease (COPD) is defined as a disease state characterized by airflow limitation that is not fully reversible (<http://www.goldcopd.com/>). COPD includes *emphysema*, an anatomically defined condition characterized by destruction and enlargement of the lung alveoli; *chronic bronchitis*, a clinically defined condition with chronic cough and phlegm; and *small airways disease*, a condition in which small bronchioles are narrowed. COPD is present only if chronic airflow obstruction occurs; chronic bronchitis *without* chronic airflow obstruction is *not* included within COPD.

COPD is the third leading cause of death and affects >10 million persons in the United States. COPD is also a disease of increasing public health importance around the world. Estimates suggest that COPD will rise from the sixth to the third most common cause of death worldwide by 2020.

PATHOGENESIS

Airflow limitation, the major physiologic change in COPD, can result from both small airway obstruction and emphysema. As described below, small airways may become narrowed by cells (hyperplasia and accumulation), mucus, and fibrosis. Of note, activation of transforming growth factor β (TGF- β) contributes to airway fibrosis, while lack of TGF- β may contribute to parenchymal inflammation and emphysema. Largely due to greater similarity of animal air spaces than airways to humans, we know more about mechanisms involved in emphysema than small airway obstruction.

The dominant paradigm of the pathogenesis of emphysema comprises four interrelated events (Fig. 314-1): (1) Chronic exposure to cigarette smoke leads to inflammatory and immune cell recruitment within the terminal air spaces of the lung. (2) These inflammatory cells release elastolytic and other proteinases that damage the extracellular matrix of the lung. (3) Structural cell death (endothelial and epithelial cells) occurs directly through oxidant-induced cigarette smoke damage and senescence as well as indirectly via proteolytic loss of matrix attachment. (4) Ineffective repair of elastin and other extracellular matrix components result in air space enlargement that defines pulmonary emphysema.

THE ELASTASE:ANTI-ELASTASE HYPOTHESIS

Elastin, the principal component of elastic fibers, is a highly stable component of the extracellular matrix that is critical to the integrity of the lung. The elastase:antielastase hypothesis proposed in the mid-1960s states that the balance of elastin-degrading enzymes and their inhibitors determines the susceptibility of the lung to destruction resulting in air space enlargement. This hypothesis was based on the clinical observation that patients with genetic deficiency in α_1 antitrypsin (α_1 AT), the inhibitor of the serine proteinase neutrophil elastase, were at increased risk of emphysema, and that instillation of elastases, including neutrophil elastase, into experimental animals results in emphysema. The elastase:antielastase hypothesis remains a prevailing mechanism for the development of emphysema. However, a complex

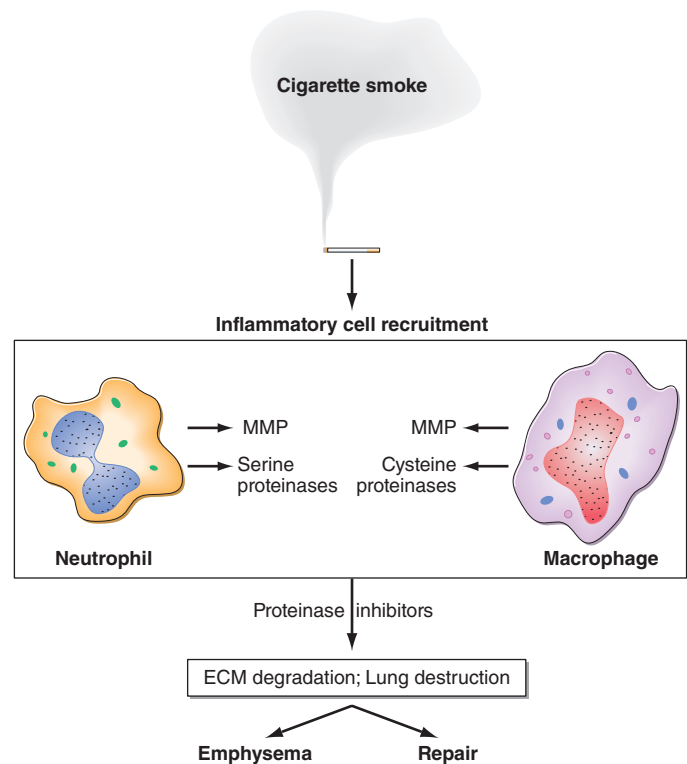


FIGURE 314-1 Pathogenesis of emphysema. Upon long-term exposure to cigarette smoke, inflammatory cells are recruited to the lung; they release proteinases in excess of inhibitors, and if repair is abnormal, this leads to air space destruction and enlargement or emphysema. ECM, extracellular matrix; MMP, matrix metalloproteinase.

network of immune and inflammatory cells and additional proteinases that contribute to emphysema have subsequently been identified.

INFLAMMATION AND EXTRACELLULAR MATRIX PROTEOLYSIS

Upon exposure to oxidants from cigarette smoke, macrophages and epithelial cells become activated, producing proteinases and chemokines that attract other inflammatory and immune cells. One mechanism of macrophage activation occurs via oxidant-induced inactivation of histone deacetylase-2, shifting the balance toward acetylated or loose chromatin, exposing nuclear factor- κ B sites, and resulting in transcription of matrix metalloproteinases, proinflammatory cytokines such as interleukin 8 (IL-8), and tumor necrosis factor α (TNF- α); this leads to neutrophil recruitment. CD8+ T cells are also recruited in response to cigarette smoke and release interferon-inducible protein-10 (IP-10, CXCL-7), which in turn leads to macrophage production of macrophage elastase (matrix metalloproteinase-12 [MMP-12]). Matrix metalloproteinases and serine proteinases, most notably neutrophil elastase, work together by degrading the inhibitor of the other, leading to lung destruction. Proteolytic cleavage products of elastin also serve as a macrophage chemokine, fueling this destructive positive feedback loop.

Autoimmune mechanisms may promote the progression of disease. Increased B cells and lymphoid follicles are present in patients, particularly those with advanced disease. Antibodies have been found against elastin fragments as well; IgG autoantibodies with avidity for pulmonary epithelium and the potential to mediate cytotoxicity have been detected.

Concomitant cigarette smoke-induced loss of cilia in the airway epithelium and impaired macrophage phagocytosis predispose to bacterial infection with neutrophilia. In end-stage lung disease, long after smoking cessation, there remains an exuberant inflammatory response, suggesting that mechanisms of cigarette smoke-induced inflammation that initiate the disease differ from mechanisms that sustain inflammation after smoking cessation.

Cell Death Cigarette smoke oxidant-mediated structural cell death occurs via a variety of mechanisms including mTOR inhibition of mammalian target of rapamycin (mTOR), leading to cell death as well as inflammation and proteolysis. Involvement of mTOR and other senescence markers has led to the recent concept that emphysema resembles premature aging of the lung. Uptake of apoptotic cells by macrophages results in production of growth factors and dampens inflammation, promoting lung repair. Cigarette smoke impairs macrophage uptake of apoptotic cells, limiting repair.

Ineffective Repair The ability of the adult lung to repair damaged alveoli appears limited. It is unlikely that the process of septation that is responsible for alveologenesis during lung development can be reinitiated. The capacity of stem cells to repopulate the lung is under active investigation. It appears difficult for an adult human to completely restore an appropriate extracellular matrix, particularly functional elastic fibers.

PATHOLOGY

Cigarette smoke exposure may affect the large airways, small airways (≤ 2 mm diameter), and alveoli. Changes in large airways cause cough and sputum, while changes in small airways and alveoli are responsible for physiologic alterations. Emphysema and small airway pathology are both present in most persons with COPD; however, they do not appear to be mechanistically related to each other, and their relative contributions to obstruction vary from one person to another.

LARGE AIRWAY

Cigarette smoking often results in mucus gland enlargement and goblet cell hyperplasia, leading to cough and mucus production that define chronic bronchitis, but these abnormalities are not related to airflow limitation. Goblet cells not only increase in number but in extent through the bronchial tree. Bronchi also undergo squamous metaplasia, predisposing to carcinogenesis and disrupting mucociliary clearance. Although not as prominent as in asthma, patients may have smooth-muscle hypertrophy and bronchial hyperreactivity

leading to airflow limitation. Neutrophil influx has been associated with purulent sputum of upper respiratory tract infections. Independent of its proteolytic activity, neutrophil elastase is among the most potent secretagogues identified.

SMALL AIRWAYS

The major site of increased resistance in most individuals with COPD is in airways ≤ 2 mm diameter. Characteristic cellular changes include goblet cell metaplasia, with these mucus-secreting cells replacing surfactant-secreting Clara cells. Smooth-muscle hypertrophy may also be present. These abnormalities may cause luminal narrowing by fibrosis, excess mucus, edema, and cellular infiltration. Reduced surfactant may increase surface tension at the air-tissue interface, predisposing to airway narrowing or collapse. Respiratory bronchiolitis with mononuclear inflammatory cells collecting in distal airway tissues may cause proteolytic destruction of elastic fibers in the respiratory bronchioles and alveolar ducts where the fibers are concentrated as rings around alveolar entrances. Narrowing and drop-out of small airways precede the onset of emphysematous destruction.

LUNG PARENCHYMA

Emphysema is characterized by destruction of gas-exchanging air spaces, i.e., the respiratory bronchioles, alveolar ducts, and alveoli. Their walls become perforated and later obliterated with coalescence of small distinct air spaces into abnormal and much larger air spaces. Macrophages accumulate in respiratory bronchioles of essentially all young smokers. Bronchoalveolar lavage fluid from such individuals contains roughly five times as many macrophages as lavage from nonsmokers. In smokers' lavage fluid, macrophages comprise $>95\%$ of the total cell count, and neutrophils, nearly absent in nonsmokers' lavage, account for 1–2% of the cells. T lymphocytes, particularly CD8+ cells, are also increased in the alveolar space of smokers.

Emphysema is classified into distinct pathologic types, the most important being centriacinar and panacinar. *Centriacinar emphysema*, the type most frequently associated with cigarette smoking, is characterized by enlarged air spaces found (initially) in association with respiratory bronchioles. Centriacinar emphysema is usually most prominent in the upper lobes and superior segments of lower lobes and is often quite focal. *Panacinar emphysema* refers to abnormally large air spaces evenly distributed within and across acinar units. Panacinar emphysema is usually observed in patients with α_1 AT deficiency, which has a predilection for the lower lobes.

PATHOPHYSIOLOGY

Persistent reduction in forced expiratory flow rates is the most typical finding in COPD. Increases in the residual volume and the residual volume/total lung capacity ratio, nonuniform distribution of ventilation, and ventilation-perfusion mismatching also occur.

AIRFLOW OBSTRUCTION

Airflow limitation, also known as airflow obstruction, is typically determined by spirometry, which involves forced expiratory maneuvers after the subject has inhaled to total lung capacity. Key parameters obtained from spirometry include the volume of air exhaled within the first second of the forced expiratory maneuver (FEV_1) and the total volume of air exhaled during the entire spirometric maneuver (forced vital capacity [FVC]). Patients with airflow obstruction related to COPD have a chronically reduced ratio of FEV_1/FVC . In contrast to asthma, the reduced FEV_1 in COPD seldom shows large responses to inhaled bronchodilators, although improvements up to 15% are common. Asthma patients can also develop chronic (not fully reversible) airflow obstruction.

Airflow during forced exhalation is the result of the balance between the elastic recoil of the lungs promoting flow and the resistance of the airways limiting flow. In normal lungs, as well as in lungs affected by COPD, maximal expiratory flow diminishes as the lungs empty because the lung parenchyma provides progressively less elastic recoil and because the cross-sectional area of the airways falls, raising the resistance to airflow. The decrease in flow coincident with decreased lung

1702 volume is readily apparent on the expiratory limb of a flow-volume curve. In the early stages of COPD, the abnormality in airflow is only evident at lung volumes at or below the functional residual capacity (closer to residual volume), appearing as a scooped-out lower part of the descending limb of the flow-volume curve. In more advanced disease, the entire curve has decreased expiratory flow compared to normal.

HYPERINFLATION

Lung volumes are also routinely assessed in pulmonary function testing. In COPD there is often “air trapping” (increased residual volume and increased ratio of residual volume to total lung capacity) and progressive hyperinflation (increased total lung capacity) late in the disease. Hyperinflation of the thorax during tidal breathing preserves maximum expiratory airflow, because as lung volume increases, elastic recoil pressure increases, and airways enlarge so that airway resistance decreases.

Despite compensating for airway obstruction, hyperinflation can push the diaphragm into a flattened position with a number of adverse effects. First, by decreasing the zone of apposition between the diaphragm and the abdominal wall, positive abdominal pressure during inspiration is not applied as effectively to the chest wall, hindering rib cage movement and impairing inspiration. Second, because the muscle fibers of the flattened diaphragm are shorter than those of a more normally curved diaphragm, they are less capable of generating inspiratory pressures than normal. Third, the flattened diaphragm (with increased radius of curvature, r) must generate greater tension (t) to develop the transpulmonary pressure (p) required to produce tidal breathing. This follows from Laplace’s law, $p = 2t/r$. Also, because the thoracic cage is distended beyond its normal resting volume, during tidal breathing the inspiratory muscles must do work to overcome the resistance of the thoracic cage to further inflation instead of gaining the normal assistance from the chest wall recoiling outward toward its resting volume.

GAS EXCHANGE

Although there is considerable variability in the relationships between the FEV₁ and other physiologic abnormalities in COPD, certain generalizations may be made. The partial pressure of oxygen in arterial blood Pao₂ usually remains near normal until the FEV₁ is decreased to ~50% of predicted, and even much lower FEV₁ values can be associated with a normal Pao₂, at least at rest. An elevation of arterial level of carbon dioxide (Paco₂) is not expected until the FEV₁ is <25% of predicted and even then may not occur. Pulmonary hypertension severe enough to cause cor pulmonale and right ventricular failure due to COPD typically occurs in individuals who have marked decreases in FEV₁ (<25% of predicted) and chronic hypoxemia (Pao₂ <55 mmHg); however, recent evidence suggests that some patients will develop significant pulmonary hypertension independent of COPD severity (**Chap. 304**).

Nonuniform ventilation and ventilation-perfusion mismatching are characteristic of COPD, reflecting the heterogeneous nature of the disease process within the airways and lung parenchyma. Physiologic studies are consistent with multiple parenchymal compartments having different rates of ventilation due to regional differences in compliance and airway resistance. Ventilation-perfusion mismatching accounts for essentially all of the reduction in Pao₂ that occurs in COPD; shunting is minimal. This finding explains the effectiveness of modest elevations of inspired oxygen in treating hypoxemia due to COPD and therefore the need to consider problems other than COPD when hypoxemia is difficult to correct with modest levels of supplemental oxygen.

RISK FACTORS

CIGARETTE SMOKING

By 1964, the Advisory Committee to the Surgeon General of the United States had concluded that cigarette smoking was a major risk factor for mortality from chronic bronchitis and emphysema. Subsequent longitudinal studies have shown accelerated decline in FEV₁ in a dose-response relationship to the intensity of cigarette smoking, which is typically expressed as pack-years (average number of packs of cigarettes smoked per day multiplied by the total number

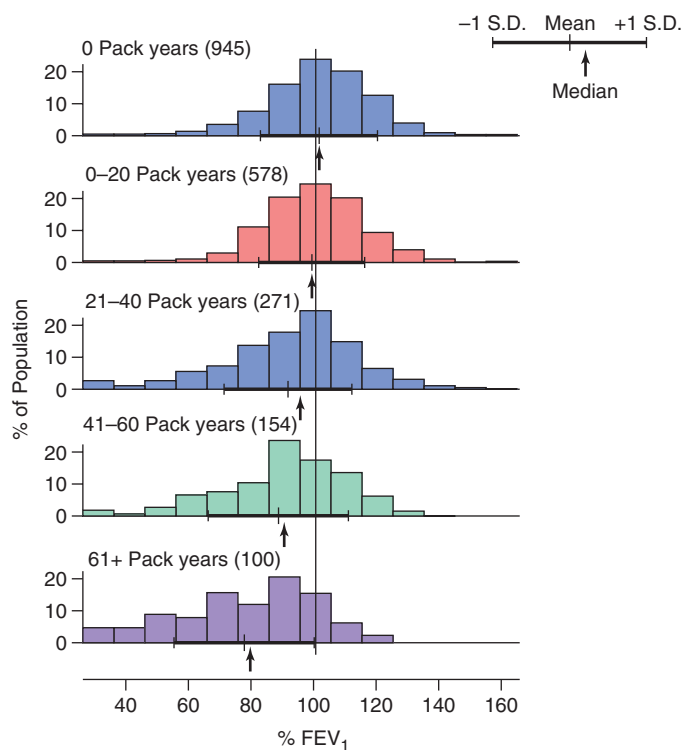


FIGURE 314-2 Distributions of forced expiratory volume in 1 s (FEV₁) values in a general population sample, stratified by pack-years of smoking. Means, medians, and ± 1 standard deviation of percent predicted FEV₁ are shown for each smoking group. Although a dose-response relationship between smoking intensity and FEV₁ was found, marked variability in pulmonary function was observed among subjects with similar smoking histories. (From B Burrows et al: *Am Rev Respir Dis* 115:95, 1977; with permission.)

of years of smoking). This dose-response relationship between reduced pulmonary function and cigarette smoking intensity accounts for the higher prevalence rates of COPD with increasing age. The historically higher rate of smoking among males is the likely explanation for the higher prevalence of COPD among males; however, the prevalence of COPD among females is increasing as the gender gap in smoking rates has diminished in the past 50 years.

Although the causal relationship between cigarette smoking and the development of COPD has been absolutely proved, there is considerable variability in the response to smoking. Although pack-years of cigarette smoking is the most highly significant predictor of FEV₁ (**Fig. 314-2**), only 15% of the variability in FEV₁ is explained by pack-years. This finding suggests that additional environmental and/or genetic factors contribute to the impact of smoking on the development of airflow obstruction.

Although cigar and pipe smoking may also be associated with the development of COPD, the evidence supporting such associations is less compelling, likely related to the lower dose of inhaled tobacco by-products during cigar and pipe smoking.

AIRWAY RESPONSIVENESS AND COPD

A tendency for increased bronchoconstriction in response to a variety of exogenous stimuli, including methacholine and histamine, is one of the defining features of asthma (**Chap. 309**). However, many patients with COPD also share this feature of airway hyperresponsiveness. The considerable overlap between persons with asthma and those with COPD in airway responsiveness, airflow obstruction, and pulmonary symptoms led to the formulation of the Dutch hypothesis. This suggests that asthma, chronic bronchitis, and emphysema are variations of the same basic disease, which is modulated by environmental and genetic factors to produce these pathologically distinct entities. The alternative British hypothesis contends that asthma and COPD

are fundamentally different diseases: Asthma is viewed as largely an allergic phenomenon, whereas COPD results from smoking-related inflammation and damage. Determination of the validity of the Dutch hypothesis versus the British hypothesis awaits identification of all of the genetic predisposing factors for asthma and/or COPD, as well as the interactions between these postulated genetic factors and environmental risk factors.

Longitudinal studies that compared airway responsiveness at the beginning of the study to subsequent decline in pulmonary function have demonstrated that increased airway responsiveness is clearly a significant predictor of subsequent decline in pulmonary function. Thus, airway hyperresponsiveness is a risk factor for COPD.

RESPIRATORY INFECTIONS

The impact of adult respiratory infections on decline in pulmonary function is controversial, but significant long-term reductions in pulmonary function are not typically seen following an episode of bronchitis or pneumonia. The impact of the effects of childhood respiratory illnesses on the subsequent development of COPD has been difficult to assess due to a lack of adequate longitudinal data. Thus, although respiratory infections are important causes of exacerbations of COPD, the association of both adult and childhood respiratory infections with the development and progression of COPD remains to be proven.

OCCUPATIONAL EXPOSURES

Increased respiratory symptoms and airflow obstruction have been suggested to result from exposure to dust and fumes at work. Several specific occupational exposures, including coal mining, gold mining, and cotton textile dust, have been suggested as risk factors for chronic airflow obstruction. Although nonsmokers in these occupations can develop some reductions in FEV₁, the importance of dust exposure as a risk factor for COPD, independent of cigarette smoking, is not certain for most of these exposures. However, among coal miners, coal mine dust exposure was a significant risk factor for emphysema in both smokers and nonsmokers. In most cases, the magnitude of these occupational exposures on COPD risk is likely substantially less important than the effect of cigarette smoking.

AMBIENT AIR POLLUTION

Some investigators have reported increased respiratory symptoms in those living in urban compared to rural areas, which may relate to increased pollution in the urban settings. However, the relationship of air pollution to chronic airflow obstruction remains unproved. Prolonged exposure to smoke produced by biomass combustion—a common mode of cooking in some countries—also appears to be a significant risk factor for COPD among women in those countries. However, in most populations, ambient air pollution is a much less important risk factor for COPD than cigarette smoking.

PASSIVE, OR SECOND-HAND, SMOKING EXPOSURE

Exposure of children to maternal smoking results in significantly reduced lung growth. In utero, tobacco smoke exposure also contributes to significant reductions in postnatal pulmonary function. Although passive smoke exposure has been associated with reductions in pulmonary function, the importance of this risk factor in the development of the severe pulmonary function reductions in COPD remains uncertain.

GENETIC CONSIDERATIONS



Although cigarette smoking is the major environmental risk factor for the development of COPD, the development of airflow obstruction in smokers is highly variable. Severe α_1 AT deficiency is a proven genetic risk factor for COPD; there is increasing evidence that other genetic determinants also exist.

α_1 Antitrypsin Deficiency Many variants of the protease inhibitor (PI or SERPINA1) locus that encodes α_1 AT have been described. The common M allele is associated with normal α_1 AT levels. The S allele, associated with slightly reduced α_1 AT levels, and the Z allele, associated with

markedly reduced α_1 AT levels, also occur with frequencies of >1% in most white populations. Rare individuals inherit null alleles, which lead to the absence of any α_1 AT production through a heterogeneous collection of mutations. Individuals with two Z alleles or one Z and one null allele are referred to as Pi^Z, which is the most common form of severe α_1 AT deficiency.

Although only approximately 1% of COPD patients are found to have severe α_1 AT deficiency as a contributing cause of COPD, these patients demonstrate that genetic factors can have a profound influence on the susceptibility for developing COPD. Pi^Z individuals often develop early-onset COPD, but the ascertainment bias in the published series of Pi^Z individuals—which have usually included many Pi^Z subjects who were tested for α_1 AT deficiency because they had COPD—means that the fraction of Pi^Z individuals who will develop COPD and the age-of-onset distribution for the development of COPD in Pi^Z subjects remain unknown. Approximately 1 in 3000 individuals in the United States inherits severe α_1 AT deficiency, but only a small minority of these individuals has been identified. The clinical laboratory test used most frequently to screen for α_1 AT deficiency is measurement of the immunologic level of α_1 AT in serum (see “Laboratory Findings”).

A significant percentage of the variability in pulmonary function among Pi^Z individuals is explained by cigarette smoking; cigarette smokers with severe α_1 AT deficiency are more likely to develop COPD at early ages. However, the development of COPD in Pi^Z subjects, even among current or ex-smokers, is not absolute. Among Pi^Z nonsmokers, impressive variability has been noted in the development of airflow obstruction. Asthma and male gender also appear to increase the risk of COPD in Pi^Z subjects. Other genetic and/or environmental factors likely contribute to this variability.

Specific treatment in the form of α_1 AT augmentation therapy is available for severe α_1 AT deficiency as a weekly IV infusion (see “Treatment,” below).

The risk of lung disease in heterozygous Pi^{MZ} individuals, who have intermediate serum levels of α_1 AT (~60% of Pi^{MM} levels), is controversial. Several recent large studies have suggested that Pi^{MZ} subjects are at slightly increased risk for the development of airflow obstruction, but it remains unclear if all Pi^{MZ} subjects are at slightly increased risk for COPD or if a subset of Pi^{MZ} subjects are at substantially increased risk for COPD due to other genetic or environmental factors.

Other Genetic Risk Factors Studies of pulmonary function measurements performed in general population samples have suggested that genetic factors other than PI type influence variation in pulmonary function. Familial aggregation of airflow obstruction within families of COPD patients has also been demonstrated.

Association studies have compared the distribution of variants in candidate genes hypothesized to be involved in the development of COPD in COPD patients and control subjects. However, the results have been quite inconsistent, often due to underpowered studies. However, a well-powered association study comprising 8300 patients and 7 separate cohorts found that a minor allele single nucleotide polymorphism (SNP) of *MMP12* (rs2276109) associated with decreased MMP12 expression has a positive effect on lung function in children with asthma and in adult smokers. Recent genome-wide association studies have identified several COPD susceptibility loci, including a region near the hedgehog interacting protein (*HHIP*) gene on chromosome 4, a cluster of genes on chromosome 15 (including components of the nicotinic acetylcholine receptor), and a region within a gene of unknown function (*FAM13A*). A regulatory SNP upstream from the *HHIP* gene has been identified as one potential functional variant; the specific genetic determinants in the other genomic regions have yet to be definitively identified.

NATURAL HISTORY

The effects of cigarette smoking on pulmonary function appear to depend on the intensity of smoking exposure, the timing of smoking exposure during growth, and the baseline lung function of the individual; other environmental factors may have similar effects. Most individuals follow a steady trajectory of increasing pulmonary function with

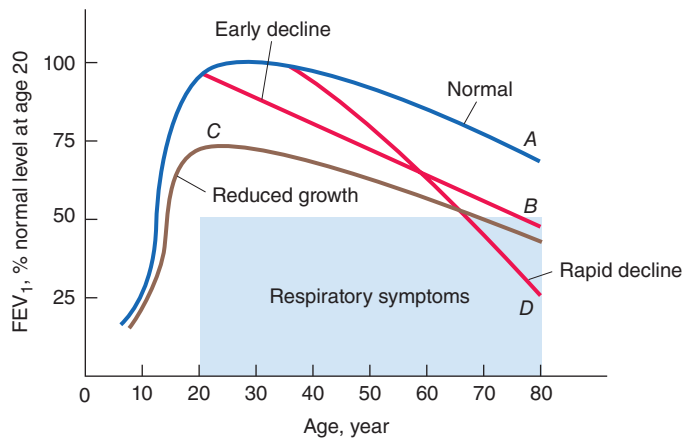


FIGURE 314-3 Hypothetical tracking curves of forced expiratory volume in 1 s (FEV_1) for individuals throughout their life spans.

The normal pattern of growth and decline with age is shown by curve A. Significantly reduced FEV_1 (<65% of predicted value at age 20) can develop from a normal rate of decline after a reduced pulmonary function growth phase (curve C), early initiation of pulmonary function decline after normal growth (curve B), or accelerated decline after normal growth (curve D). (From B Rijcken: Doctoral dissertation, p 133, University of Groningen, 1991; with permission.)

growth during childhood and adolescence, followed by a gradual decline with aging. Individuals appear to track in their quantile of pulmonary function based on environmental and genetic factors that put them on different tracks. The risk of eventual mortality from COPD is closely associated with reduced levels of FEV_1 . A graphic depiction of the natural history of COPD is shown as a function of the influences on tracking curves of FEV_1 in Fig. 314-3. Death or disability from COPD can result from a normal rate of decline after a reduced growth phase (curve C), an early initiation of pulmonary function decline after normal growth (curve B), or an accelerated decline after normal growth (curve D). The rate of decline in pulmonary function can be modified by changing environmental exposures (i.e., quitting smoking), with smoking cessation at an earlier age providing a more beneficial effect than smoking cessation after marked reductions in pulmonary function have already developed. Genetic factors likely contribute to the level of pulmonary function achieved during growth and to the rate of decline in response to smoking and potentially to other environmental factors as well.

CLINICAL PRESENTATION

HISTORY

The three most common symptoms in COPD are cough, sputum production, and exertional dyspnea. Many patients have such symptoms for months or years before seeking medical attention. Although the development of airflow obstruction is a gradual process, many patients date the onset of their disease to an acute illness or exacerbation. A careful history, however, usually reveals the presence of symptoms prior to the acute exacerbation. The development of exertional dyspnea, often described as increased effort to breathe, heaviness, air hunger, or gasping, can be insidious. It is best elicited by a careful history focused on typical physical activities and how the patient's ability to perform them has changed. Activities involving significant arm work, particularly at or above shoulder level, are particularly difficult for patients with COPD. Conversely, activities that allow the patient to brace the arms and use accessory muscles of respiration are better tolerated. Examples of such activities include pushing a shopping cart or walking on a treadmill. As COPD advances, the principal feature is worsening dyspnea on exertion with increasing intrusion on the ability to perform vocational or avocational activities. In the most advanced stages, patients are breathless doing simple activities of daily living.

Accompanying worsening airflow obstruction is an increased frequency of exacerbations (described below). Patients may also develop resting hypoxemia and require institution of supplemental oxygen.

PHYSICAL FINDINGS

In the early stages of COPD, patients usually have an entirely normal physical examination. Current smokers may have signs of active smoking, including an odor of smoke or nicotine staining of fingernails. In patients with more severe disease, the physical examination is notable for a prolonged expiratory phase and may include expiratory wheezing. In addition, signs of hyperinflation include a barrel chest and enlarged lung volumes with poor diaphragmatic excursion as assessed by percussion. Patients with severe airflow obstruction may also exhibit use of accessory muscles of respiration, sitting in the characteristic "tripod" position to facilitate the actions of the sternocleidomastoid, scalene, and intercostal muscles. Patients may develop cyanosis, visible in the lips and nail beds.

Although traditional teaching is that patients with predominant emphysema, termed "pink puffers," are thin and noncyanotic at rest and have prominent use of accessory muscles, and patients with chronic bronchitis are more likely to be heavy and cyanotic ("blue bloaters"), current evidence demonstrates that most patients have elements of both bronchitis and emphysema and that the physical examination does not reliably differentiate the two entities.

Advanced disease may be accompanied by cachexia, with significant weight loss, bitemporal wasting, and diffuse loss of subcutaneous adipose tissue. This syndrome has been associated with both inadequate oral intake and elevated levels of inflammatory cytokines (TNF- α). Such wasting is an independent poor prognostic factor in COPD. Some patients with advanced disease have paradoxical inward movement of the rib cage with inspiration (Hoover's sign), the result of alteration of the vector of diaphragmatic contraction on the rib cage as a result of chronic hyperinflation.

Signs of overt right heart failure, termed *cor pulmonale*, are relatively infrequent since the advent of supplemental oxygen therapy.

Clubbing of the digits is not a sign of COPD, and its presence should alert the clinician to initiate an investigation for causes of clubbing. In this population, the development of lung cancer is the most likely explanation for newly developed clubbing.

LABORATORY FINDINGS

The hallmark of COPD is airflow obstruction (discussed above). Pulmonary function testing shows airflow obstruction with a reduction in FEV_1 and FEV_1/FVC (Chap. 306e). With worsening disease severity, lung volumes may increase, resulting in an increase in total lung capacity, functional residual capacity, and residual volume. In patients with emphysema, the diffusing capacity may be reduced, reflecting the lung parenchymal destruction characteristic of the disease. The degree of airflow obstruction is an important prognostic factor in COPD and is the basis for the Global Initiative for Lung Disease (GOLD) severity classification (Table 314-1). More recently it has been shown that a multifactorial index incorporating airflow obstruction, exercise performance, dyspnea, and body mass index is a better predictor of mortality rate than pulmonary function alone. In 2011, the GOLD added an additional classification system incorporating symptoms and exacerbation history; the utility of this system remains to be defined.

Arterial blood gases and oximetry may demonstrate resting or exertional hypoxemia. Arterial blood gases provide additional information

TABLE 314-1 GOLD CRITERIA FOR SEVERITY OF AIRFLOW OBSTRUCTION IN COPD

GOLD Stage	Severity	Spirometry
I	Mild	$FEV_1/FVC < 0.7$ and $FEV_1 \geq 80\%$ predicted
II	Moderate	$FEV_1/FVC < 0.7$ and $FEV_1 \geq 50\%$ but $< 80\%$ predicted
III	Severe	$FEV_1/FVC < 0.7$ and $FEV_1 \geq 30\%$ but $< 50\%$ predicted
IV	Very severe	$FEV_1/FVC < 0.7$ and $FEV_1 < 30\%$ predicted

Abbreviations: COPD, chronic obstructive pulmonary disease; GOLD, Global Initiative for Lung Disease.

Source: From the Global Strategy for Diagnosis, Management and Prevention of COPD 2014, © Global Initiative for Chronic Obstructive Lung Disease (GOLD), all rights reserved. Available from <http://www.goldcopd.org>.



FIGURE 314-4 Chest computed tomography scan of a patient with chronic obstructive pulmonary disease who underwent a left single-lung transplant. Note the reduced parenchymal markings in the right lung (left side of figure) as compared to the left lung, representing emphysematous destruction of the lung, and mediastinal shift to the left, indicative of hyperinflation.

about alveolar ventilation and acid-base status by measuring arterial P_{CO_2} and pH. The change in pH with P_{CO_2} is 0.08 units/10 mmHg acutely and 0.03 units/10 mmHg in the chronic state. Knowledge of the arterial pH therefore allows the classification of ventilatory failure, defined as $P_{CO_2} > 45$ mmHg, into acute or chronic conditions. The arterial blood gas is an important component of the evaluation of patients presenting with symptoms of an exacerbation. An elevated hematocrit suggests the presence of chronic hypoxemia, as does the presence of signs of right ventricular hypertrophy.

Radiographic studies may assist in the classification of the type of COPD. Obvious bullae, paucity of parenchymal markings, or hyperlucency suggests the presence of emphysema. Increased lung volumes and flattening of the diaphragm suggest hyperinflation but do not provide information about chronicity of the changes. Computed tomography (CT) scan is the current definitive test for establishing the presence or absence of emphysema in living subjects (Fig. 314-4). From a practical perspective, the CT scan currently does little to influence therapy of COPD except in individuals considering surgical therapy for their disease (described below) and as screening for lung cancer.

Recent guidelines have suggested testing for α_1 AT deficiency in all subjects with COPD or asthma with chronic airflow obstruction. Measurement of the serum α_1 AT level is a reasonable initial test. For subjects with low α_1 AT levels, the definitive diagnosis of α_1 AT deficiency requires protease inhibitor (PI) type determination. This is typically performed by isoelectric focusing of serum, which reflects the genotype at the PI locus for the common alleles and many of the rare PI alleles as well. Molecular genotyping of DNA can be performed for the common PI alleles (M, S, and Z).

TREATMENT CHRONIC OBSTRUCTIVE PULMONARY DISEASE

STABLE PHASE COPD

Only three interventions—smoking cessation, oxygen therapy in chronically hypoxemic patients, and lung volume reduction surgery in selected patients with emphysema—have been demonstrated

to influence the natural history of patients with COPD. There is currently suggestive, but not definitive, evidence that the use of inhaled glucocorticoids may alter mortality rate (but not lung function). All other current therapies are directed at improving symptoms and decreasing the frequency and severity of exacerbations. The institution of these therapies should involve an assessment of symptoms, potential risks, costs, and benefits of therapy. This should be followed by an assessment of response to therapy, and a decision should be made whether or not to continue treatment.

PHARMACOTHERAPY

Smoking Cessation (See also Chap. 470) It has been shown that middle-aged smokers who were able to successfully stop smoking experienced a significant improvement in the rate of decline in pulmonary function, returning to annual changes similar to that of nonsmoking patients. Thus, all patients with COPD should be strongly urged to quit smoking and educated about the benefits of quitting. An emerging body of evidence demonstrates that combining pharmacotherapy with traditional supportive approaches considerably enhances the chances of successful smoking cessation. There are three principal pharmacologic approaches to the problem: bupropion; nicotine replacement therapy available as gum, transdermal patch, lozenge, inhaler, and nasal spray; and varenicline, a nicotinic acid receptor agonist/antagonist. Current recommendations from the U.S. Surgeon General are that all adult, nonpregnant smokers considering quitting be offered pharmacotherapy, in the absence of any contraindication to treatment.

Bronchodilators In general, bronchodilators are used for symptomatic benefit in patients with COPD. The inhaled route is preferred for medication delivery because the incidence of side effects is lower than that seen with the use of parenteral medication delivery.

Anticholinergic Agents Ipratropium bromide improves symptoms and produces acute improvement in FEV_1 . Tiotropium, a long-acting anticholinergic, has been shown to improve symptoms and reduce exacerbations. Studies of both ipratropium and tiotropium have failed to demonstrate that either influences the rate of decline in FEV_1 . In a large randomized clinical trial, there was a trend toward reduced mortality rate in the tiotropium-treated patients that approached, but did not reach, statistical significance. Side effects are minor, and a trial of inhaled anticholinergics is recommended in symptomatic patients with COPD. Recent retrospective analyses raised the possibility that anticholinergic use is associated with increased cardiovascular events in the COPD population. This was not demonstrated in a large, prospective randomized trial of tiotropium.

Beta Agonists These provide symptomatic benefit. The main side effects are tremor and tachycardia. Long-acting inhaled β agonists, such as salmeterol or formoterol, have benefits comparable to ipratropium bromide. Their use is more convenient than short-acting agents. The addition of a β agonist to inhaled anticholinergic therapy has been demonstrated to provide incremental benefit. A recent report in asthma suggests that those patients, particularly African Americans, using a long-acting β agonist without concomitant inhaled corticosteroids have an increased risk of deaths from respiratory causes. The applicability of these data to patients with COPD is unclear.

Inhaled Glucocorticoids Although a recent trial demonstrated an apparent benefit from the regular use of inhaled glucocorticoids on the rate of decline of lung function, a number of other well-designed randomized trials have not. Patients studied included those with mild to severe airflow obstruction and current and ex-smokers. Patients with significant acute response to inhaled β agonists were excluded from many of these trials, which may impact the generalizability of the findings. Their use has been associated with increased rates of oropharyngeal candidiasis and an increased rate of loss of bone density. Available data suggest that inhaled glucocorticoids reduce exacerbation frequency by ~25%. The impact of

1706 inhaled corticosteroids on mortality rates in COPD is controversial. A meta-analysis and several retrospective studies suggest a mortality benefit, but in a recently published randomized trial, differences in mortality rate approached, but did not reach, conventional criteria for statistical significance. A trial of inhaled glucocorticoids should be considered in patients with frequent exacerbations, defined as two or more per year, and in patients who demonstrate a significant amount of acute reversibility in response to inhaled bronchodilators.

Oral Glucocorticoids The chronic use of oral glucocorticoids for treatment of COPD is not recommended because of an unfavorable benefit/risk ratio. The chronic use of oral glucocorticoids is associated with significant side effects, including osteoporosis, weight gain, cataracts, glucose intolerance, and increased risk of infection. A recent study demonstrated that patients tapered off chronic low-dose prednisone (~10 mg/d) did not experience any adverse effect on the frequency of exacerbations, health-related quality of life, or lung function. On average, patients lost ~4.5 kg (~10 lb) when steroids were withdrawn.

Theophylline Theophylline produces modest improvements in expiratory flow rates and vital capacity and a slight improvement in arterial oxygen and carbon dioxide levels in patients with moderate to severe COPD. Nausea is a common side effect; tachycardia and tremor have also been reported. Monitoring of blood theophylline levels is typically required to minimize toxicity. The selective phosphodiesterase 4 (PDE4) inhibitor roflumilast has been demonstrated to reduce exacerbation frequency in COPD patients with chronic bronchitis and a prior history of exacerbations; its effects on airflow obstruction and symptoms are modest.

Antibiotics As outlined below, there are strong data implicating bacterial infection as a precipitant of a substantial portion of exacerbations. Early trials of prophylactic or suppressive antibiotics, given either seasonally or year round, failed to show a positive impact on exacerbation occurrence. More recently, a randomized clinical trial of azithromycin, chosen for both its anti-inflammatory and antimicrobial properties, administered daily to subjects with a history of exacerbation in the past 6 months demonstrated a reduced exacerbation frequency and longer time to first exacerbation in the macrolide-treated cohort (hazard ratio, 0.73).

Oxygen Supplemental O₂ is the only pharmacologic therapy demonstrated to unequivocally decrease mortality rates in patients with COPD. For patients with resting hypoxemia (resting O₂ saturation ≤88% or <90% with signs of pulmonary hypertension or right heart failure), the use of O₂ has been demonstrated to have a significant impact on mortality rate. Patients meeting these criteria should be on continual oxygen supplementation because the mortality benefit is proportional to the number of hours per day oxygen is used. Various delivery systems are available, including portable systems that patients may carry to allow mobility outside the home.

Supplemental O₂ is commonly prescribed for patients with exertional hypoxemia or nocturnal hypoxemia. Although the rationale for supplemental O₂ in these settings is physiologically sound, the benefits of such therapy are not well substantiated.

Other Agents N-acetyl cysteine has been used in patients with COPD for both its mucolytic and antioxidant properties. A prospective trial failed to find any benefit with respect to decline in lung function or prevention of exacerbations. Specific treatment in the form of IV α₁AT augmentation therapy is available for individuals with severe α₁AT deficiency. Despite sterilization procedures for these blood-derived products and the absence of reported cases of viral infection from therapy, some physicians recommend hepatitis B vaccination prior to starting augmentation therapy. Although biochemical efficacy of α₁AT augmentation therapy has been shown, a randomized controlled trial of α₁AT augmentation therapy has not definitively established the efficacy of augmentation therapy in reducing decline of pulmonary function. Eligibility for α₁AT augmentation therapy requires a serum α₁AT level <11 μM (approximately

50 mg/dL). Typically, Pi² individuals will qualify, although other rare types associated with severe deficiency (e.g., null-null) are also eligible. Because only a fraction of individuals with severe α₁AT deficiency will develop COPD, α₁AT augmentation therapy is not recommended for severely α₁AT-deficient persons with normal pulmonary function and a normal chest CT scan.

NONPHARMACOLOGIC THERAPIES

General Medical Care Patients with COPD should receive the influenza vaccine annually. Polyvalent pneumococcal vaccine is also recommended, although proof of efficacy in this patient population is not definitive. Similar recommendations and limitations of evidence also exist for vaccination for *Bordetella pertussis*.

Pulmonary Rehabilitation This refers to a treatment program that incorporates education and cardiovascular conditioning. In COPD, pulmonary rehabilitation has been demonstrated to improve health-related quality of life, dyspnea, and exercise capacity. It has also been shown to reduce rates of hospitalization over a 6- to 12-month period.

Lung Volume Reduction Surgery (LVRS) Surgery to reduce the volume of lung in patients with emphysema was first introduced with minimal success in the 1950s and was reintroduced in the 1990s. Patients are excluded if they have significant pleural disease, a pulmonary artery systolic pressure >45 mmHg, extreme deconditioning, congestive heart failure, or other severe comorbid conditions. Patients with an FEV₁ <20% of predicted and either diffusely distributed emphysema on CT scan or diffusing capacity of lung for carbon monoxide (DL_{CO}) <20% of predicted have an increased mortality rate after the procedure and thus are not candidates for LVRS.

The National Emphysema Treatment trial demonstrated that LVRS offers both a mortality benefit and a symptomatic benefit in certain patients with emphysema. The anatomic distribution of emphysema and post-rehabilitation exercise capacity are important prognostic characteristics. Patients with upper lobe-predominant emphysema and a low post-rehabilitation exercise capacity are most likely to benefit from LVRS.

Lung Transplantation (See also Chap. 320e) COPD is currently the second leading indication for lung transplantation (Fig. 314-4). Current recommendations are that candidates for lung transplantation should have severe disability despite maximal medical therapy and be free of comorbid conditions such as liver, renal, or cardiac disease. In contrast to LVRS, the anatomic distribution of emphysema and the presence of pulmonary hypertension are not contraindications to lung transplantation.

EXACERBATIONS OF COPD

Exacerbations are a prominent feature of the natural history of COPD. Exacerbations are episodes of increased dyspnea and cough and change in the amount and character of sputum. They may or may not be accompanied by other signs of illness, including fever, myalgias, and sore throat. Self-reported health-related quality of life correlates with frequency of exacerbations more closely than it does with the degree of airflow obstruction. Economic analyses have shown that >70% of COPD-related health care expenditures go to emergency department visits and hospital care; this translates to >\$10 billion annually in the United States. The frequency of exacerbations increases as airflow obstruction increases; patients with moderate to severe airflow obstruction (GOLD stage III or IV; Table 314-1) on average have one to three episodes per year. However, some individuals with very severe airflow obstruction do not have frequent exacerbations; the history of prior exacerbations is a strong predictor of future exacerbations. Recently, an elevated ratio of the diameter of the pulmonary artery to aorta on chest CT has been associated with increased risk of COPD exacerbations.

The approach to the patient experiencing an exacerbation includes an assessment of the severity of the patient's illness, both acute and chronic components; an attempt to identify the precipitant of the exacerbation; and the institution of therapy.

Precipitating Causes and Strategies to Reduce Frequency of Exacerbations

A variety of stimuli may result in the final common pathway of airway inflammation and increased symptoms that are characteristic of COPD exacerbations. Studies suggest that acquiring a new strain of bacteria is associated with increased near-term risk of exacerbation and that bacterial infection/superinfection is involved in over 50% of exacerbations. Viral respiratory infections are present in approximately one-third of COPD exacerbations. In a significant minority of instances (20–35%), no specific precipitant can be identified.

The role of pharmacotherapy in reducing exacerbation frequency is less well studied. Chronic oral glucocorticoids are not recommended for this purpose. Inhaled glucocorticoids reduce the frequency of exacerbations by 25–30% in most analyses. The use of inhaled glucocorticoids should be considered in patients with frequent exacerbations or those who have an asthmatic component, i.e., significant reversibility on pulmonary function testing or marked symptomatic improvement after inhaled bronchodilators. Similar magnitudes of reduction have been reported for anticholinergic and long-acting β -agonist therapy. The influenza vaccine has been shown to reduce exacerbation rates in patients with COPD. As outlined above, daily azithromycin administered to subjects with COPD and an exacerbation history reduces exacerbation frequency.

Patient Assessment An attempt should be made to establish the severity of the exacerbation as well as the severity of preexisting COPD. The more severe either of these two components, the more likely that the patient will require hospital admission. The history should include quantification of the degree of dyspnea by asking about breathlessness during activities of daily living and typical activities for the patient. The patient should be asked about fever; change in character of sputum; any ill contacts; and associated symptoms such as nausea, vomiting, diarrhea, myalgias, and chills. Inquiring about the frequency and severity of prior exacerbations can provide important information.

The physical examination should incorporate an assessment of the degree of distress of the patient. Specific attention should be focused on tachycardia, tachypnea, use of accessory muscles, signs of perioral or peripheral cyanosis, the ability to speak in complete sentences, and the patient's mental status. The chest examination should establish the presence or absence of focal findings, degree of air movement, presence or absence of wheezing, asymmetry in the chest examination (suggesting large airway obstruction or pneumothorax mimicking an exacerbation), and the presence or absence of paradoxical motion of the abdominal wall.

Patients with severe underlying COPD, who are in moderate or severe distress, or those with focal findings should have a chest x-ray. Approximately 25% of x-rays in this clinical situation will be abnormal, with the most frequent findings being pneumonia and congestive heart failure. Patients with advanced COPD, those with a history of hypercarbia, those with mental status changes (confusion, sleepiness), or those in significant distress should have an arterial blood-gas measurement. The presence of hypercarbia, defined as a $P_{CO_2} > 45$ mmHg, has important implications for treatment (discussed below). In contrast to its utility in the management of exacerbations of asthma, measurement of pulmonary function has not been demonstrated to be helpful in the diagnosis or management of exacerbations of COPD.

There are no definitive guidelines concerning the need for inpatient treatment of exacerbations. Patients with respiratory acidosis and hypercarbia, significant hypoxemia, or severe underlying disease or those whose living situation is not conducive to careful observation and the delivery of prescribed treatment should be admitted to the hospital.

ACUTE EXACERBATIONS

Bronchodilators Typically, patients are treated with an inhaled β agonist, often with the addition of an anticholinergic agent. These may be administered separately or together, and the frequency of administration depends on the severity of the exacerbation.

Patients are often treated initially with nebulized therapy, as such treatment is often easier to administer in older patients or to those in respiratory distress. It has been shown, however, that conversion to metered-dose inhalers is effective when accompanied by education and training of patients and staff. This approach has significant economic benefits and also allows an easier transition to outpatient care. The addition of methylxanthines (such as theophylline) to this regimen can be considered, although convincing proof of its efficacy is lacking. If added, serum levels should be monitored in an attempt to minimize toxicity.

Antibiotics Patients with COPD are frequently colonized with potential respiratory pathogens, and it is often difficult to identify conclusively a specific species of bacteria responsible for a particular clinical event. Bacteria frequently implicated in COPD exacerbations include *Streptococcus pneumoniae*, *Haemophilus influenzae*, and *Moraxella catarrhalis*. In addition, *Mycoplasma pneumoniae* or *Chlamydia pneumoniae* are found in 5–10% of exacerbations. The choice of antibiotic should be based on local patterns of antibiotic susceptibility of the above pathogens as well as the patient's clinical condition. Most practitioners treat patients with moderate or severe exacerbations with antibiotics, even in the absence of data implicating a specific pathogen.

Glucocorticoids Among patients admitted to the hospital, the use of glucocorticoids has been demonstrated to reduce the length of stay, hasten recovery, and reduce the chance of subsequent exacerbation or relapse for a period of up to 6 months. One study demonstrated that 2 weeks of glucocorticoid therapy produced benefit indistinguishable from 8 weeks of therapy. The GOLD guidelines recommend 30–40 mg of oral prednisolone or its equivalent for a period of 10–14 days. Hyperglycemia, particularly in patients with preexisting diagnosis of diabetes, is the most frequently reported acute complication of glucocorticoid treatment.

Oxygen Supplemental O_2 should be supplied to keep arterial saturations $\geq 90\%$. Hypoxemic respiratory drive plays a small role in patients with COPD. Studies have demonstrated that in patients with both acute and chronic hypercarbia, the administration of supplemental O_2 does not reduce minute ventilation. It does, in some patients, result in modest increases in arterial P_{CO_2} , chiefly by altering ventilation-perfusion relationships within the lung. This should not deter practitioners from providing the oxygen needed to correct hypoxemia.

Mechanical Ventilatory Support The initiation of noninvasive positive-pressure ventilation (NIPPV) in patients with respiratory failure, defined as $P_{aCO_2} > 45$ mmHg, results in a significant reduction in mortality rate, need for intubation, complications of therapy, and hospital length of stay. Contraindications to NIPPV include cardiovascular instability, impaired mental status or inability to cooperate, copious secretions or the inability to clear secretions, craniofacial abnormalities or trauma precluding effective fitting of mask, extreme obesity, or significant burns.

Invasive (conventional) mechanical ventilation via an endotracheal tube is indicated for patients with severe respiratory distress despite initial therapy, life-threatening hypoxemia, severe hypercarbia and/or acidosis, markedly impaired mental status, respiratory arrest, hemodynamic instability, or other complications. The goal of mechanical ventilation is to correct the aforementioned conditions. Factors to consider during mechanical ventilatory support include the need to provide sufficient expiratory time in patients with severe airflow obstruction and the presence of auto-PEEP (positive end-expiratory pressure), which can result in patients having to generate significant respiratory effort to trigger a breath during a demand mode of ventilation. The mortality rate of patients requiring mechanical ventilatory support is 17–30% for that particular hospitalization. For patients age > 65 admitted to the intensive care unit for treatment, the mortality rate doubles over the next year to 60%, regardless of whether mechanical ventilation was required.

315 Interstitial Lung Diseases

Talmadge E. King, Jr.

Patients with interstitial lung diseases (ILDs) come to medical attention mainly because of the onset of progressive exertional dyspnea or a persistent nonproductive cough. Hemoptysis, wheezing, and chest pain may be present. Often, the identification of interstitial opacities on chest x-ray focuses the diagnostic approach on one of the ILDs.

ILDs represent a large number of conditions that involve the parenchyma of the lung—the alveoli, the alveolar epithelium, the capillary endothelium, and the spaces between those structures—as well as the perivascular and lymphatic tissues. The disorders in this heterogeneous group are classified together because of similar clinical, roentgenographic, physiologic, or pathologic manifestations. These disorders often are associated with considerable rates of morbidity and mortality, and there is little consensus regarding the best management of most of them.

ILDs have been difficult to classify because >200 known individual diseases are characterized by diffuse parenchymal lung involvement, either as the primary condition or as a significant part of a multiorgan process, as may occur in the connective tissue diseases (CTDs). One useful approach to classification is to separate the ILDs into two groups based on the major underlying histopathology: (1) those associated with predominant inflammation and fibrosis and (2) those with a predominantly granulomatous reaction in interstitial or vascular areas (Table 315-1). Each of these groups can be subdivided further according to whether the cause is known or unknown. For each ILD there may be an acute phase, and there is usually a chronic one as well. Rarely, some are recurrent, with intervals of subclinical disease.

Sarcoidosis (Chap. 390), idiopathic pulmonary fibrosis (IPF), and pulmonary fibrosis associated with CTDs (Chaps. 378, 382, 388, and 427) are the most common ILDs of unknown etiology. Among the ILDs of known cause, the largest group includes occupational and environmental exposures, especially the inhalation of inorganic dusts, organic dusts, and various fumes or gases (Chap. 311). A multidisciplinary approach—requiring close communication between clinician, radiologist, and when appropriate, pathologist—is often required to make the diagnosis. High-resolution computed tomography (HRCT) scanning improves the diagnostic accuracy and may eliminate the need for tissue examination in many cases, especially in IPF. For other forms, tissue examination, usually obtained by thoracoscopic lung biopsy, is critical to confirmation of the diagnosis.

PATHOGENESIS

The ILDs are nonmalignant disorders and are not caused by identified infectious agents. The precise pathway(s) leading from injury to fibrosis is not known. Although there are multiple initiating agent(s) of injury, the immunopathogenic responses of lung tissue are limited, and the mechanisms of repair have common features (Fig. 315-1).

As mentioned above, the two major histopathologic patterns are a granulomatous pattern and a pattern in which inflammation and fibrosis predominate.

Granulomatous Lung Disease This process is characterized by an accumulation of T lymphocytes, macrophages, and epithelioid cells organized into discrete structures (granulomas) in the lung parenchyma. The granulomatous lesions can progress to fibrosis. Many patients with granulomatous lung disease remain free of severe impairment of lung function or, when symptomatic, improve after treatment. The main differential diagnosis is between sarcoidosis (Chap. 390) and hypersensitivity pneumonitis (Chap. 310).

Inflammation and Fibrosis The initial insult is an injury to the epithelial surface that causes inflammation in the air spaces and alveolar walls. If the disease becomes chronic, inflammation spreads to adjacent portions of the interstitium and vasculature and eventually causes interstitial fibrosis. Important histopathologic patterns found in the ILDs include

TABLE 315-1 MAJOR CATEGORIES OF ALVEOLAR AND INTERSTITIAL INFLAMMATORY LUNG DISEASE

Lung Response: Alveolitis, Interstitial Inflammation, and Fibrosis

Known Cause

Asbestos	Residual of acute respiratory distress syndrome
Fumes, gases	Smoking-related
Drugs (antibiotics, amiodarone, gold) and chemotherapy drugs	Desquamative interstitial pneumonia
Radiation	Respiratory bronchiolitis-associated interstitial lung disease
Aspiration pneumonia	Pulmonary Langerhans cell granulomatosis

Unknown Cause

Idiopathic interstitial pneumonias	Pulmonary alveolar proteinosis
Idiopathic pulmonary fibrosis (usual interstitial pneumonia)	Lymphocytic infiltrative disorders (lymphocytic interstitial pneumonitis associated with connective tissue disease)
Acute interstitial pneumonia (diffuse alveolar damage)	Eosinophilic pneumonias
Cryptogenic organizing pneumonia	Lymphangioleiomyomatosis
Nonspecific interstitial pneumonia	Inherited diseases
Idiopathic lymphocytic interstitial pneumonia	Tuberous sclerosis, neurofibromatosis, Niemann-Pick disease, Gaucher disease, Hermansky-Pudlak syndrome
Rare and ill-defined entities	
Idiopathic pleuroparenchymal fibroelastosis	
Acute fibrinous and organizing pneumonia	
Bronchiolocentric patterns of interstitial pneumonia	
Connective tissue diseases	Gastrointestinal or liver diseases (Crohn disease, primary biliary cirrhosis, chronic active hepatitis, ulcerative colitis)
Systemic lupus erythematosus, rheumatoid arthritis, ankylosing spondylitis, systemic sclerosis, Sjögren syndrome, polymyositis-dermatomyositis	
Pulmonary hemorrhage syndromes	Graft-versus-host disease (bone marrow transplantation; solid organ transplantation)
Goodpasture syndrome, idiopathic pulmonary hemosiderosis, isolated pulmonary capillaritis	
Amyloidosis	

Lung Response: Granulomatous

Known Cause

Hypersensitivity pneumonitis (organic dusts)	Inorganic dusts: beryllium, silica
--	------------------------------------

Unknown Cause

Sarcoidosis	Bronchocentric granulomatosis
Granulomatous vasculitides	Lymphomatoid granulomatosis
Granulomatosis with polyangiitis (Wegener)	
Eosinophilic granulomatosis with polyangiitis (Churg-Strauss)	

usual interstitial pneumonia (UIP), nonspecific interstitial pneumonia, respiratory bronchiolitis/desquamative interstitial pneumonia, organizing pneumonia, diffuse alveolar damage (acute or organizing), and lymphocytic interstitial pneumonia. The development of irreversible scarring (fibrosis) of alveolar walls, airways, or vasculature is the most feared outcome in all of these conditions because it is often progressive and leads to significant derangement of ventilatory function and gas exchange.

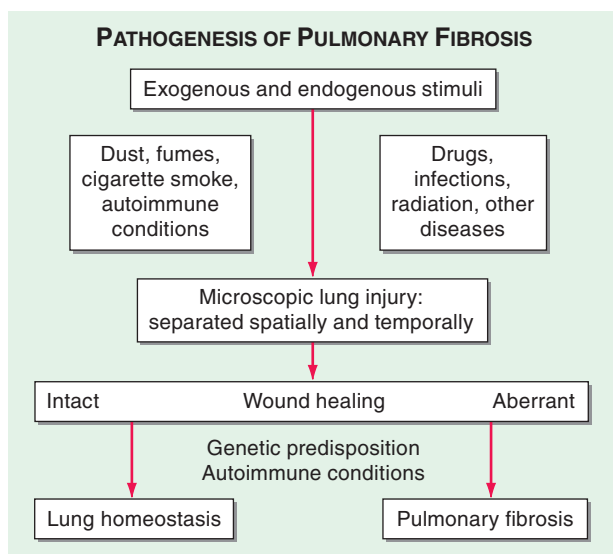


FIGURE 315-1 Proposed mechanism for the pathogenesis of pulmonary fibrosis. The lung is naturally exposed to repetitive injury from a variety of exogenous and endogenous stimuli. Several local and systemic factors (e.g., fibroblasts, circulating fibrocytes, chemokines, growth factors, and clotting factors) contribute to tissue healing and functional recovery. Dysregulation of this intricate network through genetic predisposition, autoimmune conditions, or superimposed diseases can lead to aberrant wound healing, with the result of pulmonary fibrosis. Alternatively, excessive injury to the lung may overwhelm even intact reparative mechanisms and lead to pulmonary fibrosis. (From S Garantziotis et al: *J Clin Invest* 114:319, 2004.)

HISTORY

Duration of Illness *Acute presentation* (days to weeks), although unusual, occurs with allergy (drugs, fungi, helminths), acute interstitial pneumonia (AIP), eosinophilic pneumonia, and hypersensitivity pneumonitis. These conditions may be confused with atypical pneumonias because of diffuse alveolar opacities on chest x-ray. *Subacute presentation* (weeks to months) may occur in all ILDs but is seen especially in sarcoidosis, drug-induced ILDs, the alveolar hemorrhage syndromes, cryptogenic organizing pneumonia (COP), and the acute immunologic pneumonia that complicates systemic lupus erythematosus (SLE) or polymyositis. In most ILDs, the symptoms and signs form a *chronic presentation* (months to years). Examples include IPF, sarcoidosis, pulmonary Langerhans cell histiocytosis (PLCH), pneumoconioses, and CTDs. *Episodic presentations* are unusual and include eosinophilic pneumonia, hypersensitivity pneumonitis, COP, vasculitides, pulmonary hemorrhage, and Churg-Strauss syndrome.

Age Most patients with sarcoidosis, ILD associated with CTD, lymphangioleiomyomatosis (LAM), PLCH, and inherited forms of ILD (familial IPF, Gaucher disease, Hermansky-Pudlak syndrome) present between the ages of 20 and 40 years. Most patients with IPF are older than 60 years.

Gender LAM and pulmonary involvement in tuberous sclerosis occur exclusively in premenopausal women. In addition, ILD in Hermansky-Pudlak syndrome and in the CTDs is more common in women; an exception is ILD in rheumatoid arthritis (RA), which is more common in men. IPF is more common in men. Because of occupational exposures, pneumoconioses also occur more frequently in men.

Family History Familial lung fibrosis has been associated with mutations in the surfactant protein C gene, the surfactant protein A2 gene, telomerase reverse transcriptase (*TERT*), telomerase RNA component (*TERC*), and the promoter of a mucin gene (*MUC5B*). Familial lung fibrosis is characterized by several patterns of interstitial pneumonia, including nonspecific interstitial pneumonia, desquamative interstitial pneumonia, and UIP. Older age, male sex, and a history of cigarette

smoking have been identified as risk factors for familial lung fibrosis. Family associations (with an autosomal dominant pattern) have been identified in tuberous sclerosis and neurofibromatosis. Familial clustering has been identified increasingly in sarcoidosis. The genes responsible for several rare ILDs have been identified, i.e., alveolar microlithiasis, Gaucher disease, Hermansky-Pudlak syndrome, and Niemann-Pick disease, along with the genes for surfactant homeostasis in pulmonary alveolar proteinosis and for control of cell growth and differentiation in LAM.

Smoking History Two-thirds to 75% of patients with IPF and familial lung fibrosis have a history of smoking. Patients with PLCH, respiratory bronchiolitis/desquamative interstitial pneumonia (DIP), Goodpasture syndrome, respiratory bronchiolitis, and pulmonary alveolar proteinosis are usually current or former smokers.

Occupational and Environmental History A strict chronologic listing of the patient's lifelong employment must be sought, including specific duties and known exposures. In hypersensitivity pneumonitis (see Fig. 310-1), respiratory symptoms, fever, chills, and an abnormal chest roentgenogram are often temporally related to a hobby (pigeon breeder's disease) or to the workplace (farmer's lung) (Chap. 310). Symptoms may diminish or disappear after the patient leaves the site of exposure for several days; similarly, symptoms may reappear when the patient returns to the exposure site.

Other Important Past History Parasitic infections may cause pulmonary eosinophilia, and therefore a travel history should be taken in patients with known or suspected ILD. History of risk factors for HIV infection should be elicited because several processes may occur at the time of initial presentation or during the clinical course, e.g., HIV infection, organizing pneumonia, AIP, lymphocytic interstitial pneumonitis, and diffuse alveolar hemorrhage.

Respiratory Symptoms and Signs Dyspnea is a common and prominent complaint in patients with ILD, especially the idiopathic interstitial pneumonias, hypersensitivity pneumonitis, COP, sarcoidosis, eosinophilic pneumonias, and PLCH. Some patients, especially those with sarcoidosis, silicosis, PLCH, hypersensitivity pneumonitis, lipid pneumonia, or lymphangitis carcinomatosa, may have extensive parenchymal lung disease on chest imaging studies without significant dyspnea, especially early in the course of the illness. Wheezing is an uncommon manifestation of ILD but has been described in patients with chronic eosinophilic pneumonia, Churg-Strauss syndrome, respiratory bronchiolitis, and sarcoidosis. Clinically significant chest pain is uncommon in most ILDs. However, substernal discomfort is common in sarcoidosis. Sudden worsening of dyspnea, especially if associated with acute chest pain, may indicate a spontaneous pneumothorax, which occurs in PLCH, tuberous sclerosis, LAM, and neurofibromatosis. Frank hemoptysis and blood-streaked sputum are rarely presenting manifestations of ILD but can be seen in the diffuse alveolar hemorrhage (DAH) syndromes, LAM, tuberous sclerosis, and the granulomatous vasculitides. Fatigue and weight loss are common in all ILDs.

PHYSICAL EXAMINATION

The findings are usually not specific. Most commonly, physical examination reveals tachypnea and bibasilar end-inspiratory dry crackles, which are common in most forms of ILD associated with inflammation but are less likely to be heard in the granulomatous lung diseases. Crackles may be present in the absence of radiographic abnormalities on the chest radiograph. Scattered late inspiratory high-pitched rhonchi—so-called inspiratory squeaks—are heard in patients with bronchiolitis. The cardiac examination is usually normal except in the middle or late stages of the disease, when findings of pulmonary hypertension and cor pulmonale may become evident (Chap. 304). Cyanosis and clubbing of the digits occur in some patients with advanced disease.

LABORATORY

Antinuclear antibodies and anti-immunoglobulin antibodies (rheumatoid factors) are identified in some patients, even in the absence

1710 of a defined CTD. A raised lactate dehydrogenase (LDH) level is a nonspecific finding common to ILDs. Elevation of the serum level of angiotensin-converting enzyme is common in ILDs, especially sarcoidosis. Serum precipitins confirm exposure when hypersensitivity pneumonitis is suspected, although they are not diagnostic of the process. Antineutrophil cytoplasmic or anti-basement membrane antibodies are useful if vasculitis is suspected. The electrocardiogram is usually normal unless pulmonary hypertension is present; then it demonstrates right-axis deviation, right ventricular hypertrophy, or right atrial enlargement or hypertrophy. Echocardiography also reveals right ventricular dilation and/or hypertrophy in the presence of pulmonary hypertension.

CHEST IMAGING STUDIES

Chest X-Ray ILD may be first suspected on the basis of an abnormal chest radiograph, which most commonly reveals a bibasilar reticular pattern. A nodular or mixed pattern of alveolar filling and increased reticular markings also may be present. Subgroups of ILDs exhibit nodular opacities with a predilection for the upper lung zones (sarcoidosis, PLCH, chronic hypersensitivity pneumonitis, silicosis, berylliosis, RA [necrobiotic nodular form], ankylosing spondylitis). The chest x-ray correlates poorly with the clinical or histopathologic stage of the disease. The radiographic finding of honeycombing correlates with pathologic findings of small cystic spaces and progressive fibrosis; when present, it portends a poor prognosis. In most cases, the chest radiograph is nonspecific and usually does not allow a specific diagnosis.

Computed Tomography HRCT is superior to the plain chest x-ray for early detection and confirmation of suspected ILD (Fig. 315-2). In addition, HRCT allows better assessment of the extent and distribution of disease, and it is especially useful in the investigation of patients with a normal chest radiograph. Coexisting disease is often best recognized on HRCT scanning, e.g., mediastinal adenopathy, carcinoma, or emphysema. In the appropriate clinical setting, HRCT may be sufficiently characteristic to preclude the need for lung biopsy in IPF, sarcoidosis, hypersensitivity pneumonitis, asbestosis, lymphangitic carcinoma, and PLCH. When a lung biopsy is required, HRCT scanning is useful for determining the most appropriate area from which biopsy samples should be taken.

PULMONARY FUNCTION TESTING

Spirometry and Lung Volumes Measurement of lung function is important in assessing the extent of pulmonary involvement in patients with ILD. Most forms of ILD produce a restrictive defect with reduced total lung capacity (TLC), functional residual capacity, and residual volume

(Chap. 306e). Forced expiratory volume in 1 second (FEV_1) and forced vital capacity (FVC) are reduced, but these changes are related to the decreased TLC. The FEV_1/FVC ratio is usually normal or increased. Lung volumes decrease as lung stiffness worsens with disease progression. A few disorders produce interstitial opacities on chest x-ray and obstructive airflow limitation on lung function testing (uncommon in sarcoidosis and hypersensitivity pneumonitis but common in tuberous sclerosis and LAM). Pulmonary function studies have been proved to have prognostic value in patients with idiopathic interstitial pneumonias, particularly IPF and nonspecific interstitial pneumonia (NSIP).

Diffusing Capacity A reduction in the diffusing capacity of the lung for carbon monoxide (DL_{CO}) is a common but nonspecific finding in most ILDs. This decrease is due in part to effacement of the alveolar capillary units but, more important, to mismatching of ventilation and perfusion (\dot{V}/\dot{Q}). Lung regions with reduced compliance due to either fibrosis or cellular infiltration may be poorly ventilated but may still maintain adequate blood flow, and the ventilation-perfusion mismatch in these regions acts like true venous admixture. The severity of the reduction in DL_{CO} does not correlate with disease stage.

Arterial Blood Gas The resting arterial blood gas may be normal or reveal hypoxemia (secondary to a mismatching of ventilation to perfusion) and respiratory alkalosis. A normal arterial O_2 tension (or saturation by oximetry) at rest does not rule out significant hypoxemia during exercise or sleep. Carbon dioxide (CO_2) retention is rare and is usually a manifestation of end-stage disease.

CARDIOPULMONARY EXERCISE TESTING

Because hypoxemia at rest is not always present and because severe exercise-induced hypoxemia may go undetected, it is useful to perform exercise testing with measurement of arterial blood gases to detect abnormalities of gas exchange. Arterial oxygen desaturation, a failure to decrease dead space appropriately with exercise (i.e., a high V_D/V_T [dead space/tidal volume] ratio [Chap. 306e]), and an excessive increase in respiratory rate with a lower than expected recruitment of tidal volume provide useful information about physiologic abnormalities and extent of disease. Serial assessment of resting and exercise gas exchange is an excellent method for following disease activity and responsiveness to treatment, especially in patients with IPF. Increasingly, the 6-min walk test is used to obtain a global evaluation of submaximal exercise capacity in patients with ILD. The walk distance and level of oxygen desaturation tend to correlate with the patient's baseline lung function and mirror the patient's clinical course.

FIBEROPTIC BRONCHOSCOPY AND BRONCHOALVEOLAR LAVAGE (BAL)

In selected diseases (e.g., sarcoidosis, hypersensitivity pneumonitis, DAH syndrome, cancer, pulmonary alveolar proteinosis), cellular analysis of BAL fluid may be useful in narrowing the differential diagnostic possibilities among various types of ILD (Table 315-2). The role of BAL in defining the stage of disease and assessment of disease progression or response to therapy remains poorly understood, and the usefulness of BAL in the clinical assessment and management remains to be established.

TISSUE AND CELLULAR EXAMINATION

Lung biopsy is the most effective method for confirming the diagnosis and assessing disease activity. The findings may identify a more treatable process than originally suspected, particularly chronic hypersensitivity pneumonitis, COP, respiratory bronchiolitis-associated ILD, or sarcoidosis. Biopsy should be obtained before the initiation of treatment. A definitive diagnosis avoids confusion and anxiety later in the clinical course if the patient does not respond to therapy or experiences serious side effects from it.

Fiberoptic bronchoscopy with multiple transbronchial lung biopsies (four to eight biopsy samples) is often the initial procedure of choice, especially when sarcoidosis, lymphangitic carcinomatosis, eosinophilic pneumonia, Goodpasture syndrome, or infection is suspected. If a specific diagnosis is not made by transbronchial biopsy, surgical lung biopsy by video-assisted thoracic surgery or open thoracotomy is



FIGURE 315-2 Idiopathic pulmonary fibrosis. High-resolution computed tomography image shows bibasilar, peripheral predominant reticular abnormality with traction bronchiectasis and honeycombing. The lung biopsy showed the typical features of usual interstitial pneumonia.

TABLE 315-2 DIAGNOSTIC VALUE OF BRONCHOALVEOLAR LAVAGE IN INTERSTITIAL LUNG DISEASE

Condition	Bronchoalveolar Lavage Finding
Sarcoidosis	Lymphocytosis; CD4:CD8 ratio >3.5 most specific of diagnosis
Hypersensitivity pneumonitis	Marked lymphocytosis (>50%)
Organizing pneumonia	Foamy macrophages; mixed pattern of increased cells characteristic; decreased CD4:CD8 ratio
Eosinophilic lung disease	Eosinophils >25%
Diffuse alveolar bleeding	Hemosiderin-laden macrophages, red blood cells
Diffuse alveolar damage, drug toxicity	Atypical hyperplastic type II pneumocytes
Opportunistic infections	<i>Pneumocystis carinii</i> , fungi, cytomegalovirus-transformed cells
Lymphangitic carcinomatosis, alveolar cell carcinoma, pulmonary lymphoma	Malignant cells
Alveolar proteinosis	Milky effluent, foamy macrophages and lipoproteinaceous intraalveolar material (periodic acid–Schiff stain–positive)
Lipoid pneumonia	Fat globules in macrophages
Pulmonary Langerhans cell histiocytosis	Increased CD1+ Langerhans cells, electron microscopy demonstrating Birbeck granule in lavaged macrophage (expensive and difficult to perform)
Asbestos-related pulmonary disease	Dust particles, ferruginous bodies
Berylliosis	Positive lymphocyte transformation test to beryllium
Silicosis	Dust particles by polarized light microscopy
Lipoidosis	Accumulation of specific lipopigment in alveolar macrophages

indicated. Adequate-sized biopsies from multiple sites, usually from two lobes, should be obtained. Relative contraindications to lung biopsy include serious cardiovascular disease, honeycombing and other roentgenographic evidence of diffuse end-stage disease, severe pulmonary dysfunction, and other major operative risks, especially in the elderly.

TREATMENT INTERSTITIAL LUNG DISEASE

Although the course of ILD is variable, progression is common and often insidious. All treatable possibilities should be carefully considered. Because therapy does not reverse fibrosis, the major goals of treatment are permanent removal of the offending agent, when known, and early identification and aggressive suppression of the acute and chronic inflammatory process, thereby reducing further lung damage. Hypoxemia ($\text{PaO}_2 < 55$ mmHg) at rest and/or with exercise should be managed with supplemental oxygen. Management of cor pulmonale may be required as the disease progresses (**Chaps. 280 and 304**). Pulmonary rehabilitation has been shown to improve the quality of life in patients with ILD.

DRUG THERAPY

Glucocorticoids are the mainstay of therapy for suppression of the inflammation present in ILD, but the success rate is low. There have been no placebo-controlled trials of glucocorticoids in ILD, and so there is no direct evidence that steroids improve survival in many of the diseases for which they are commonly used. Glucocorticoid therapy is recommended for symptomatic ILD patients with eosinophilic pneumonias, COP, CTD, sarcoidosis, hypersensitivity pneumonitis, acute inorganic dust exposures, acute radiation pneumonitis, DAH, and drug-induced ILD. In organic dust disease, glucocorticoids are recommended for both the acute and chronic stages.

The optimal dose and proper length of therapy with glucocorticoids in the treatment of most ILDs are not known. A common

starting dose is prednisone, 0.5–1 mg/kg in a once-daily oral dose (based on the patient's lean body weight). This dose is continued for 4–12 weeks, at which time the patient is reevaluated. If the patient is stable or improved, the dose is tapered to 0.25–0.5 mg/kg and is maintained at this level for an additional 4–12 weeks, depending on the course. Rapid tapering or a shortened course of glucocorticoid treatment can result in recurrence. If the patient's condition continues to decline on glucocorticoids, a second agent (see below) often is added and the prednisone dose is lowered to or maintained at 0.25 mg/kg per day.

Cyclophosphamide, azathioprine (1–2 mg/kg lean body weight per day), and mycophenolate mofetil, with or without glucocorticoids, have been tried with variable success in IPF, vasculitis, progressive systemic sclerosis, and other ILDs. An objective response usually requires at least 8–12 weeks to occur. In situations in which these drugs have failed or could not be tolerated, other agents, including methotrexate and cyclosporine, have been tried. However, their role in the treatment of ILDs remains to be determined.

Many cases of ILD are chronic and irreversible despite the therapy discussed above, and lung transplantation may then be considered (**Chap. 320e**).

INDIVIDUAL FORMS OF INTERSTITIAL LUNG DISEASE

IDIOPATHIC PULMONARY FIBROSIS

IPF is the most common form of idiopathic interstitial pneumonia. Separating IPF from other forms of lung fibrosis is an important step in the evaluation of all patients presenting with ILD. IPF has a distinctly poor response to therapy and a bad prognosis.

Clinical Manifestations Exertional dyspnea, a nonproductive cough, and inspiratory crackles with or without digital clubbing may be present on physical examination. HRCT lung scans typically show patchy, predominantly basilar, subpleural reticular opacities, often associated with traction bronchiectasis and honeycombing (Fig. 315-2). A definite UIP pattern on HRCT is highly accurate for the presence of a UIP pattern on surgical lung biopsy. Atypical findings that should suggest an alternative diagnosis include extensive ground-glass abnormality, nodular opacities, upper or midzone predominance, and prominent hilar or mediastinal lymphadenopathy. Pulmonary function tests often reveal a restrictive pattern, a reduced DL_{CO} , and arterial hypoxemia that is exaggerated or elicited by exercise.

Histologic Findings Confirmation of the presence of the UIP pattern on histologic examination is essential to confirm this diagnosis. Transbronchial biopsies are not helpful in making the diagnosis of UIP, and surgical biopsy usually is required. The histologic hallmark and chief diagnostic criterion of UIP is a heterogeneous appearance at low magnification with alternating areas of normal lung, interstitial inflammation, foci of proliferating fibroblasts, dense collagen fibrosis, and honeycomb changes. These histologic changes affect the peripheral, subpleural parenchyma most severely. The interstitial inflammation is usually patchy and consists of a lymphoplasmacytic infiltrate in the alveolar septa, associated with hyperplasia of type 2 pneumocytes. The fibrotic zones are composed mainly of dense collagen, although scattered foci of proliferating fibroblasts are a consistent finding. The extent of fibroblastic proliferation is predictive of disease progression. Areas of honeycomb change are composed of cystic fibrotic air spaces that frequently are lined by bronchiolar epithelium and filled with mucin. Smooth-muscle hyperplasia is commonly seen in areas of fibrosis and honeycomb change. A fibrotic pattern with some features similar to UIP may be found in the chronic stage of several specific disorders, such as pneumoconioses (e.g., asbestosis), radiation injury, certain drug-induced lung diseases (e.g., nitrofurantoin), chronic aspiration, sarcoidosis, chronic hypersensitivity pneumonitis, organized chronic eosinophilic pneumonia, and PLCH. Commonly, other histopathologic features are present in these situations, thus allowing separation of these lesions from the UIP-like pattern. Consequently, the term *usual interstitial pneumonia* is used for patients in whom the lesion is idiopathic and not associated with another condition.

Untreated patients with IPF show continued progression of their disease and have a high mortality rate. There is no effective therapy for IPF. Thalidomide appears to improve cough in patients with IPF. Chronic microaspiration secondary to gastroesophageal reflux may play a role in the pathogenesis and natural history of IPF. Gastroesophageal reflux (GER) therapy may be of benefit in IPF. In patients with IPF, treatment with the three-drug regimen of prednisone, azathioprine, and *N*-acetylcysteine (NAC) or warfarin (in IPF patients who lacked other indications for anticoagulation) has been shown to increase the risks of hospitalization and death.

Patients with IPF and coexisting emphysema (combined pulmonary fibrosis and emphysema [CPFE]) are more likely to require long-term oxygen therapy and develop pulmonary hypertension and may have a more dismal outcome than those without emphysema.

Patients with IPF may have acute deterioration secondary to infections, pulmonary embolism, or pneumothorax. Heart failure and ischemic heart disease are common problems in patients with IPF, accounting for nearly one-third of deaths. These patients also commonly experience an accelerated phase of rapid clinical decline that is associated with a poor prognosis (so-called acute exacerbations of IPF). These acute exacerbations are defined by worsening of dyspnea within a few days to 4 weeks; newly developing diffuse ground-glass abnormality and/or consolidation superimposed on a background reticular or honeycomb pattern consistent with the UIP pattern; worsening hypoxemia; and absence of infectious pneumonia, heart failure, and sepsis. The rate of these acute exacerbations ranges from 10–57%, apparently depending on the length of follow-up. During these episodes, the histopathologic pattern of diffuse alveolar damage is often found on the background of UIP. No therapy has been found to be effective in the management of acute exacerbations of IPF. Often mechanical ventilation is required, but it is usually not successful, with a hospital mortality rate of up to three-fourths of patients. In those who survive, a recurrence of acute exacerbation is common and usually results in death at those times.

Patients should be referred early for lung transplant because of the unpredictability of disease progression (e.g., acute exacerbations) (**Chap. 320e**).

NONSPECIFIC INTERSTITIAL PNEUMONIA

This condition defines a subgroup of the idiopathic interstitial pneumonias that can be distinguished clinically and pathologically from UIP, DIP, AIP, and COP. Importantly, many cases with this histopathologic pattern occur in the context of an underlying disorder, such as a CTD, drug-induced ILD, or chronic hypersensitivity pneumonitis.

Clinical Manifestations Patients with idiopathic NSIP have clinical, serologic, radiographic, and pathologic characteristics highly suggestive of autoimmune disease and meet the criteria for undifferentiated CTD. Idiopathic NSIP is a subacute restrictive process with a presentation similar to that of IPF but usually at a younger age, most commonly in women who have never smoked. It is often associated with a febrile illness. HRCT shows bilateral, subpleural ground-glass opacities, often associated with lower lobe volume loss (**Fig. 315-3**). Patchy areas of airspace consolidation and reticular abnormalities may be present, but honeycombing is unusual.

Histologic Findings The key histopathologic feature of NSIP is the uniformity of interstitial involvement across the biopsy section, and this may be predominantly cellular or fibrosing. There is less temporal and spatial heterogeneity than in UIP, and little or no honeycombing is found. The cellular variant is rare.

Treatment The majority of patients with NSIP have a good prognosis (5-year mortality rate estimated at <15%), with most showing improvement after treatment with glucocorticoids, often used in combination with azathioprine or mycophenolate mofetil.



FIGURE 315-3 Nonspecific interstitial pneumonia. High-resolution computed tomography through the lower lung shows volume loss with extensive ground-glass abnormality, reticular abnormality, and traction bronchiectasis. There is sparing on the lung immediately adjacent to the pleura. Histology showed a combination of inflammation and mild fibrosis.

ACUTE INTERSTITIAL PNEUMONIA (HAMMAN-RICH SYNDROME)

Clinical Manifestations AIP is a rare, fulminant form of lung injury characterized histologically by diffuse alveolar damage on lung biopsy. Most patients are older than 40 years. AIP is similar in presentation to the acute respiratory distress syndrome (ARDS) (**Chap. 322**) and probably corresponds to the subset of cases of idiopathic ARDS. The onset is usually abrupt in a previously healthy individual. A prodromal illness, usually lasting 7–14 days before presentation, is common. Fever, cough, and dyspnea are common manifestations at presentation. Diffuse, bilateral, air-space opacification is present on the chest radiograph. HRCT scans show bilateral, patchy, symmetric areas of ground-glass attenuation. Bilateral areas of air-space consolidation also may be present. A predominantly subpleural distribution may be seen.

Histologic Findings The diagnosis of AIP requires the presence of a clinical syndrome of idiopathic ARDS and pathologic confirmation of organizing diffuse alveolar damage. Therefore, lung biopsy is required to confirm the diagnosis.

Treatment Most patients have moderate to severe hypoxemia and develop respiratory failure. Mechanical ventilation is often required. The mortality rate is high (>60%), with most patients dying within 6 months of presentation. Recurrences have been reported. However, those who recover often have substantial improvement in lung function. The main treatment is supportive. It is not clear that glucocorticoid therapy is effective.

CRYPTOGENIC ORGANIZING PNEUMONIA

Clinical Manifestations COP is a clinicopathologic syndrome of unknown etiology. The onset is usually in the fifth and sixth decades. The presentation may be of a fluke illness with cough, fever, malaise, fatigue, and weight loss. Inspiratory crackles are frequently present on examination. Pulmonary function is usually impaired, with a restrictive defect and arterial hypoxemia being most common. The roentgenographic manifestations are distinctive, revealing bilateral, patchy, or diffuse alveolar opacities in the presence of normal lung volume. Recurrent and migratory pulmonary opacities are common. HRCT shows areas of air-space consolidation, ground-glass opacities, small nodular opacities, and bronchial wall thickening and dilation. These changes occur more frequently in the periphery of the lung and in the lower lung zone.

Histologic Findings Lung biopsy shows granulation tissue within small airways, alveolar ducts, and airspaces, with chronic inflammation in the surrounding alveoli. Foci of organizing pneumonia are a nonspecific reaction to lung injury found adjacent to other pathologic processes or as a component of other primary pulmonary disorders (e.g., cryptococcosis, granulomatosis with polyangiitis [Wegener], lymphoma, hypersensitivity pneumonitis, and eosinophilic pneumonia). Consequently, the clinician must carefully reevaluate any patient found to have this histopathologic lesion to rule out these possibilities.

Treatment Glucocorticoid therapy induces clinical recovery in two-thirds of patients. A few patients have rapidly progressive courses with fatal outcomes despite glucocorticoids.

ILD ASSOCIATED WITH CIGARETTE SMOKING

Desquamative Interstitial Pneumonia • CLINICAL MANIFESTATIONS DIP is a rare but distinct clinical and pathologic entity found almost exclusively in cigarette smokers. The histologic hallmark is the extensive accumulation of macrophages in intraalveolar spaces with minimal interstitial fibrosis. The peak incidence is in the fourth and fifth decades. Most patients present with dyspnea and cough. Lung function testing shows a restrictive pattern with reduced DL_{CO} and arterial hypoxemia. The chest x-ray and HRCT scans usually show diffuse hazy opacities.

HISTOLOGIC FINDINGS A diffuse and uniform accumulation of macrophages in the alveolar spaces is the hallmark of DIP. The macrophages contain golden, brown, or black pigment of tobacco smoke. There may be mild thickening of the alveolar walls by fibrosis and scanty inflammatory cell infiltration.

TREATMENT Clinical recognition of DIP is important because the process is associated with a better prognosis (10-year survival rate is ~70%) in response to smoking cessation. There are no clear data showing that systemic glucocorticoids are effective in DIP.

Respiratory Bronchiolitis–Associated ILD • CLINICAL MANIFESTATIONS Respiratory bronchiolitis–associated ILD (RB-ILD) is considered to be a subset of DIP and is characterized by the accumulation of macrophages in peribronchial alveoli. The clinical presentation is similar to that of DIP. Crackles are often heard on chest examination and occur throughout inspiration; sometimes they continue into expiration. The process is best seen on HRCT lung scanning, which shows bronchial wall thickening, centrilobular nodules, ground-glass opacity, and emphysema with air trapping. There is a spectrum of CT features in asymptomatic smokers (and elderly asymptomatic individuals) that may not necessarily represent clinically relevant disease.

HISTOLOGIC FINDINGS The histologic findings in RB-ILD include alveolar macrophage accumulation in respiratory bronchioles, with a variable chronic inflammatory cell infiltrate in bronchiolar and surrounding alveolar walls and occasional peribronchial alveolar septal fibrosis. The pulmonary parenchyma may show presence of smoking-related emphysema.

TREATMENT RB-ILD appears to resolve in most patients after smoking cessation alone.

Pulmonary Langerhans Cell Histiocytosis • CLINICAL MANIFESTATIONS This is a rare, smoking-related, diffuse lung disease that primarily affects men between the ages of 20 and 40 years. The clinical presentation varies from an asymptomatic state to a rapidly progressive condition. The most common clinical manifestations at presentation are cough, dyspnea, chest pain, weight loss, and fever. Pneumothorax occurs in ~25% of patients. Hemoptysis and diabetes insipidus are rare manifestations. The radiographic features vary with the stage of the disease. The combination of ill-defined or stellate nodules (2–10 mm in diameter), reticular or nodular opacities, bizarre-shaped upper zone cysts, preservation of lung volume, and sparing of the costophrenic angles are characteristics of PLCH. HRCT that reveals a combination of nodules and thin-walled cysts is virtually diagnostic of PLCH. The most common pulmonary function abnormality is a markedly reduced

DL_{CO} , although varying degrees of restrictive disease, airflow limitation, and diminished exercise capacity may occur.

HISTOLOGIC FINDINGS The characteristic histopathologic finding in PLCH is the presence of nodular sclerosing lesions that contain Langerhans cells accompanied by mixed cellular infiltrates. The nodular lesions are poorly defined and are distributed in a bronchiolocentric fashion with intervening normal lung parenchyma. As the disease advances, fibrosis progresses to involve adjacent lung tissue, leading to pericatricial air space enlargement, which accounts for the concomitant cystic changes.

TREATMENT Discontinuance of smoking is the key treatment, resulting in clinical improvement in one-third of patients. Most patients with PLCH experience persistent or progressive disease. Death due to respiratory failure occurs in ~10% of patients.

ILD ASSOCIATED WITH CONNECTIVE TISSUE DISORDERS

Clinical findings suggestive of a CTD (musculoskeletal pain, weakness, fatigue, fever, joint pain or swelling, photosensitivity, Raynaud's phenomenon, pleuritis, dry eyes, dry mouth) should be sought in any patient with ILD. The CTDs may be difficult to rule out since the pulmonary manifestations occasionally precede the more typical systemic manifestations by months or years. The most common form of pulmonary involvement is the nonspecific interstitial pneumonia histopathologic pattern. However, determining the precise nature of lung involvement in most of the CTDs is difficult due to the high incidence of lung involvement caused by disease-associated complications of esophageal dysfunction (predisposing to aspiration and secondary infections), respiratory muscle weakness (atelectasis and secondary infections), complications of therapy (opportunistic infections), and associated malignancies. For the majority of CTDs, with the exception of progressive system sclerosis, recommended initial treatment for ILD includes oral glucocorticoids often in association with an immunosuppressive agent (usually oral or intravenous cyclophosphamide or oral azathioprine) or mycophenolate mofetil.

Progressive Systemic Sclerosis (PSS) • CLINICAL MANIFESTATIONS (See also Chap. 382) Clinical evidence of ILD is present in about one-half of patients with PSS, and pathologic evidence in three-quarters. Pulmonary function tests show a restrictive pattern and impaired diffusing capacity, often before any clinical or radiographic evidence of lung disease appears. The HRCT features of lung disease in PSS range from predominant ground-glass attenuation to a predominant reticular pattern and are mostly similar to idiopathic NSIP.

HISTOLOGIC FINDINGS NSIP is the histopathologic pattern in most patients (~75%); the UIP pattern is rare (<10%).

TREATMENT Therapy is similar to that in idiopathic NSIP. UIP in PSS has a better outcome than IPF. The most widely used initial treatment regimen is low-dose glucocorticoid therapy and an immunosuppressive agent, usually oral or pulse cyclophosphamide. There are no convincing data showing this regime to be efficacious, and there is concern that the risk of renal crisis rises substantially with corticosteroids. Pulmonary vascular disease alone or in association with pulmonary fibrosis, pleuritis, or recurrent aspiration pneumonitis is strikingly resistant to current modes of therapy.

Rheumatoid Arthritis • CLINICAL MANIFESTATIONS (See also Chap. 380) ILD associated with RA is more common in men. Pulmonary manifestations of RA include pleurisy with or without effusion, ILD in up to 20% of cases, necrobiotic nodules (nonpneumoconiotic intrapulmonary rheumatoid nodules) with or without cavities, Caplan syndrome (rheumatoid pneumoconiosis), pulmonary hypertension secondary to rheumatoid pulmonary vasculitis, organized pneumonia, and upper airway obstruction due to cricoarytenoid arthritis.

HISTOLOGIC FINDINGS There are two primary histopathologic patterns of ILD that are observed in patients with ILD associated with RA: NSIP pattern and UIP pattern.

1714 TREATMENT Little data exist to support the management of ILD in RA. Initial treatment of rheumatoid ILD, if required, is typically with oral glucocorticoids, which should be tried for 1–3 months. The potential benefit of anti-tumor necrosis factor α (TNF- α) therapy has been clouded by concerns about the development of a rapid and occasionally fatal lung disease in patients with RA-associated ILD treated with anti-TNF- α therapy.

Systemic Lupus Erythematosus • CLINICAL MANIFESTATIONS (See also Chap. 378) Lung disease is a common complication in SLE. Pleuritis with or without effusion is the most common pulmonary manifestation. Other lung manifestations include the following: atelectasis, diaphragmatic dysfunction with loss of lung volumes, pulmonary vascular disease, pulmonary hemorrhage, uremic pulmonary edema, infectious pneumonia, and organized pneumonia. Acute lupus pneumonitis characterized by pulmonary capillaritis leading to alveolar hemorrhage is uncommon. Chronic, progressive ILD is uncommon (<10%). It is important to exclude pulmonary infection. Although pleuropulmonary involvement may not be evident clinically, pulmonary function testing, particularly DL_{CO}, reveals abnormalities in many patients with SLE.

HISTOLOGIC FINDINGS The most common pathologic patterns seen include NSIP, UIP, LIP, and, on occasion, organizing pneumonia and amyloidosis.

TREATMENT There have been no controlled trials of treatment for ILD in SLE. Treatment involves the use of a glucocorticoid, either alone or, more often, in combination with an additional immunomodulating agent.

Polymyositis and Dermatomyositis (PM/DM) • CLINICAL MANIFESTATIONS (See also Chap. 388) ILD occurs in ~10% of patients with PM/DM. Diffuse reticular or nodular opacities with or without an alveolar component occur radiographically, with a predilection for the lung bases (NSIP pattern). ILD occurs more commonly in the subgroup of patients with an anti-Jo-1 antibody that is directed to histidyl tRNA synthetase. Weakness of respiratory muscles contributing to aspiration pneumonia may be present. A rapidly progressive illness characterized by diffuse alveolar damage may cause respiratory failure.

HISTOLOGIC FINDINGS NSIP predominates over UIP, organizing pneumonia, or other patterns of interstitial pneumonia.

TREATMENT The optimal treatment is unknown. The most widely used initial treatment is oral glucocorticoids. Fulminant disease may require high-dose intravenous methylprednisolone (1.0 g/d) for 3–5 days.

Sjögren Syndrome • CLINICAL MANIFESTATIONS (See also Chap. 383) General dryness and lack of airway secretion cause the major problems of hoarseness, cough, and bronchitis.

HISTOLOGIC FINDINGS Lung biopsy is frequently required to establish a precise pulmonary diagnosis. Fibrotic NSIP is most common. Lymphocytic interstitial pneumonitis, lymphoma, pseudolymphoma, bronchiolitis, and bronchiolitis obliterans are associated with this condition.

TREATMENT Glucocorticoids have been used in the management of ILD associated with Sjögren syndrome with some degree of clinical success.

DRUG-INDUCED ILD

Clinical Manifestations Many classes of drugs have the potential to induce diffuse ILD, which is manifest most commonly as exertional dyspnea and nonproductive cough. A detailed history of the medications taken by the patient is needed to identify drug-induced disease, including over-the-counter medications, oily nose drops, and petroleum products (mineral oil). In most cases, the pathogenesis is unknown, although a combination of direct toxic effects of the drug (or its metabolite) and indirect inflammatory and immunologic events

are likely. The onset of the illness may be abrupt and fulminant, or it may be insidious, extending over weeks to months. The drug may have been taken for several years before a reaction develops (e.g., amiodarone), or the lung disease may occur weeks to years after the drug has been discontinued (e.g., carmustine). The extent and severity of disease are usually dose-related.

Histologic Findings The patterns of lung injury vary widely and depend on the agent.

Treatment Treatment consists of discontinuation of any possible offending drug and supportive care.

EOSINOPHILIC PNEUMONIA

(See Chap. 310)

PULMONARY ALVEOLAR PROTEINOSIS (PAP)

Clinical Manifestations Although not strictly an ILD, PAP resembles and is therefore considered with these conditions. It has been proposed that a defect in macrophage function, more specifically an impaired ability to process surfactant, may play a role in the pathogenesis of PAP. PAP is an autoimmune disease with a neutralizing antibody of immunoglobulin G isotype against granulocyte-macrophage colony-stimulating factor (GM-CSF). These findings suggest that neutralization of GM-CSF bioactivity by the antibody causes dysfunction of alveolar macrophages, which results in reduced surfactant clearance. There are three distinct classes of PAP: acquired (>90% of all cases), congenital, and secondary. *Congenital PAP* is transmitted in an autosomal recessive manner and is caused by homozygosity for a frameshift mutation (121ins2) in the *SP-B* gene, which leads to an unstable *SP-B* mRNA, reduced protein levels, and secondary disturbances of *SP-C* processing. *Secondary PAP* is rare among adults and is caused by lysinuric protein intolerance, acute silicosis and other inhalational syndromes, immunodeficiency disorders, and malignancies (almost exclusively of hematopoietic origin) and hematopoietic disorders.

The typical age of presentation is 30–50 years, and males predominate. The clinical presentation is usually insidious and is manifested by progressive exertional dyspnea, fatigue, weight loss, and low-grade fever. A nonproductive cough is common, but occasionally expectoration of “chunky” gelatinous material may occur. Polycythemia, hypergammaglobulinemia, and increased LDH levels are common. Markedly elevated serum levels of lung surfactant proteins A and D have been found in PAP. In the absence of any known secondary cause of PAP, an elevated serum anti-GM-CSF titer is highly sensitive and specific for the diagnosis of acquired PAP. BAL fluid levels of anti-GM-CSF antibodies correlate better with the severity of PAP than do serum titers. Radiographically, bilateral symmetric alveolar opacities located centrally in middle and lower lung zones result in a “bat-wing” distribution. HRCT shows a ground-glass opacification and thickened intralobular structures and interlobular septa.

Histologic Findings This diffuse disease is characterized by the accumulation of an amorphous, periodic acid–Schiff–positive lipoproteinaceous material in the distal air spaces. There is little or no lung inflammation, and the underlying lung architecture is preserved.

Treatment Whole-lung lavage(s) through a double-lumen endotracheal tube provides relief to many patients with dyspnea or progressive hypoxemia and also may provide long-term benefit.

PULMONARY LYMPHANGIOLEIOMYOMATOSIS

Clinical Manifestations Pulmonary LAM is a rare condition that afflicts premenopausal women and should be suspected in young women with “emphysema,” recurrent pneumothorax, or chylous pleural effusion. It is often misdiagnosed as asthma or chronic obstructive pulmonary disease. Whites are affected much more commonly than are members of other racial groups. The disease accelerates during pregnancy and abates after oophorectomy. Common complaints at presentation are dyspnea, cough, and chest pain. Hemoptysis may

be life threatening. Spontaneous pneumothorax occurs in 50% of patients; it may be bilateral and necessitate pleurodesis. Meningioma and renal angiomyolipomas (hamartomas), characteristic findings in the genetic disorder tuberous sclerosis, are also common in patients with LAM. Chylothorax, chyloperitoneum (chylous ascites), chyluria, and chylopericardium are other complications. Pulmonary function testing usually reveals an obstructive or mixed obstructive-restrictive pattern, and gas exchange is often abnormal. HRCT shows thin-walled cysts surrounded by normal lung without zonal predominance.

Histologic Findings Pathologically, LAM is characterized by the proliferation of atypical pulmonary interstitial smooth muscle and cyst formation. The immature-appearing smooth-muscle cells react with monoclonal antibody HMB45, which recognizes a 100-kDa glycoprotein (gp100) originally found in human melanoma cells.

Treatment Progression is common, with a median survival of 8–10 years from diagnosis. No therapy is of proven benefit in LAM. Sirolimus, an inhibitor of the mammalian target of rapamycin (mTOR), appears to be an active agent for LAM. After 12 months, it stabilized lung function (FVC, FEV₁, and functional residual capacity) and was associated with a reduction in symptoms and improvement in quality of life. Adverse effects (e.g., mucositis, diarrhea, nausea, hypercholesterolemia, acneiform rash, peripheral edema) were more common in the sirolimus group, but serious adverse effects were not increased. Subjects were followed off sirolimus for an additional 12 months, during which time pulmonary function declined at the same rate as in the placebo group. Progesterone and luteinizing hormone–releasing hormone analogues have been used. Oophorectomy is no longer recommended, and estrogen-containing drugs should be discontinued. Lung transplantation offers the only hope for cure despite reports of recurrent disease in the transplanted lung.

SYNDROMES OF ILD WITH DIFFUSE ALVEOLAR HEMORRHAGE

Clinical Manifestations The clinical onset is often abrupt, with cough, fever, and dyspnea. Severe respiratory distress requiring ventilatory support may be evident at initial presentation. Although hemoptysis is expected, it can be absent at the time of presentation in one-third of the cases. For patients without hemoptysis, new alveolar opacities, a falling hemoglobin level, and hemorrhagic BAL fluid point to the diagnosis. The chest radiograph is nonspecific and most commonly shows new patchy or diffuse alveolar opacities. Recurrent episodes of DAH may lead to pulmonary fibrosis, resulting in interstitial opacities on the chest radiograph. An elevated white blood cell count and falling hematocrit are common. Evidence for impaired renal function caused by focal segmental necrotizing glomerulonephritis, usually with crescent formation, also may be present. Varying degrees of hypoxemia may occur and are often severe enough to require ventilatory support. Dr_{CO} may be increased, resulting from the increased hemoglobin within the alveoli compartment.

Histologic Findings Injury to arterioles, venules, and the alveolar septal (alveolar wall or interstitial) capillaries can result in hemoptysis secondary to disruption of the alveolar-capillary basement membrane. This results in bleeding into the alveolar spaces, which characterizes DAH. Pulmonary capillaritis, characterized by a neutrophilic infiltration of the alveolar septae, may lead to necrosis of these structures, loss of capillary structural integrity, and the pouring of red blood cells into the alveolar space. Fibrinoid necrosis of the interstitium and red blood cells within the interstitial space are sometimes seen. Bland pulmonary hemorrhage (i.e., DAH without inflammation of the alveolar structures) also may occur.

Evaluation of either lung or renal tissue by immunofluorescent techniques indicates an absence of immune complexes (pauci-immune) in granulomatosis with polyangiitis (Wegener), microscopic polyangiitis, pauci-immune glomerulonephritis, and isolated pulmonary capillaritis. A granular pattern is found in the CTDs, particularly SLE, and a characteristic linear deposition is found in Goodpasture syndrome. Granular deposition of IgA-containing immune complexes is present in Henoch-Schönlein purpura.

Treatment The mainstay of therapy for the DAH associated with systemic vasculitis, CTD, Goodpasture syndrome, and isolated pulmonary capillaritis is IV methylprednisolone, 0.5–2 g daily in divided doses for up to 5 days, followed by a gradual tapering, and then maintenance on an oral preparation. Prompt initiation of therapy is important, particularly in the face of renal insufficiency, because early initiation of therapy has the best chance of preserving renal function. The decision to start other immunosuppressive therapy (cyclophosphamide or azathioprine) acutely depends on the severity of illness.

Goodpasture Syndrome • CLINICAL MANIFESTATIONS Pulmonary hemorrhage and glomerulonephritis are features in most patients with this disease. Autoantibodies to renal glomerular and lung alveolar basement membranes are present. This syndrome can present and recur as DAH without an associated glomerulonephritis. In such cases, circulating anti-basement membrane antibody is often absent, and the only way to establish the diagnosis is by demonstrating linear immunofluorescence in lung tissue.

HISTOLOGIC FINDINGS The underlying histology may be bland hemorrhage or DAH associated with capillaritis.

TREATMENT Plasmapheresis has been recommended as adjunctive treatment.

INHERITED DISORDERS ASSOCIATED WITH ILD

Pulmonary opacities and respiratory symptoms typical of ILD can develop in related family members and in several inherited diseases. These diseases include the phakomatoses, tuberous sclerosis and neurofibromatosis (Chap. 118), and the lysosomal storage diseases, Niemann-Pick disease and Gaucher disease (Chap. 432e). The Hermansky-Pudlak syndrome is an autosomal recessive disorder in which granulomatous colitis and ILD may occur. It is characterized by oculocutaneous albinism, bleeding diathesis secondary to platelet dysfunction, and the accumulation of a chromolipid, lipofuscin material in cells of the reticuloendothelial system. A fibrotic pattern is found on lung biopsy, but the alveolar macrophages may contain cytoplasmic ceroid-like inclusions.

ILD WITH A GRANULOMATOUS RESPONSE IN LUNG TISSUE OR VASCULAR STRUCTURES

Inhalation of organic dusts, which cause hypersensitivity pneumonitis, or of inorganic dust, such as silica, which elicits a granulomatous inflammatory reaction leading to ILD, produces diseases of known etiology (Table 315-1) that are discussed in Chaps. 310 and 311. Sarcoidosis (Chap. 390) is prominent among granulomatous diseases of unknown cause in which ILD is an important feature.

Granulomatous Vasculitides (See also Chap. 385) The granulomatous vasculitides are characterized by pulmonary angiitis (i.e., inflammation and necrosis of blood vessels) with associated granuloma formation (i.e., infiltrates of lymphocytes, plasma cells, epithelioid cells, or histiocytes, with or without the presence of multinucleated giant cells, sometimes with tissue necrosis). The lungs are almost always involved, although any organ system may be affected. Granulomatosis with polyangiitis (Wegener) and Eosinophilic granulomatosis with polyangiitis (Churg-Strauss) primarily affect the lung but are associated with a systemic vasculitis as well. The granulomatous vasculitides generally limited to the lung include necrotizing sarcoid granulomatosis and benign lymphocytic angiitis and granulomatosis. Granulomatous infection and pulmonary angiitis due to irritating embolic material (e.g., talc) are important known causes of pulmonary vasculitis.

LYMPHOCYTIC INFILTRATIVE DISORDERS

This group of disorders features lymphocyte and plasma cell infiltration of the lung parenchyma. The disorders either are benign or can behave as low-grade lymphomas. Included is angioimmunoblastic lymphadenopathy with dysproteinemia, a rare lymphoproliferative disorder characterized by diffuse lymphadenopathy, fever, hepatosplenomegaly, and hemolytic anemia, with ILD in some cases.

1716 Lymphocytic Interstitial Pneumonitis This rare form of ILD occurs in adults, some of whom have an autoimmune disease or dysproteinemia. It has been reported in patients with Sjögren syndrome and HIV infection.

Lymphomatoid Granulomatosis • CLINICAL MANIFESTATIONS Pulmonary lymphomatoid granulomatosis generally presents predominantly in men between the ages of 30 and 50, although patients can be affected at any age. The effects of race and geography on disease incidence are not known, although a higher diagnosis rate is reported in Western countries. Although it may affect virtually any organ, it is most frequently characterized by pulmonary (>90%), skin, and central nervous system involvement. The most common presenting symptoms and signs include cough, fever, rash/nodules, malaise, weight loss, neurologic abnormalities, dyspnea, and chest pain.

HISTOLOGIC FINDINGS This multisystem disorder of unknown etiology is an angiocentric malignant (T cell) lymphoma characterized by a polymorphic lymphoid infiltrate, an angiitis, and granulomatosis.

TREATMENT The clinical course of lymphomatoid granulomatosis ranges from remission without treatment to death from malignant lymphoma within 2 years. The choice of a treatment strategy should be based upon the presence of symptoms, history of using an inciting medication, extent of extrapulmonary involvement, and careful assessment of the histopathologic grade of the lesion. Referral to a hematologic oncology specialist for consultation is recommended.

BRONCHOCENTRIC GRANULOMATOSIS

Clinical Manifestations Rather than a specific clinical entity, bronchocentric granulomatosis (BG) is a descriptive histologic term that is applied to an uncommon and nonspecific pathologic response to a variety of airway injuries. There is evidence that BG is caused by a hypersensitivity reaction to *Aspergillus* or other fungi in patients with asthma. About one-half of the patients described have had chronic asthma with severe wheezing and peripheral blood eosinophilia. In patients with asthma, BG probably represents one pathologic manifestation of allergic bronchopulmonary aspergillosis or another allergic mycosis. In patients without asthma, BG has been associated with RA and a variety of infections, including tuberculosis, echinococcosis, histoplasmosis, coccidioidomycosis, and nocardiosis. The chest roentgenogram reveals irregularly shaped nodular or mass lesions with ill-defined margins, which are usually unilateral and solitary, with upper lobe predominance.

Histologic Findings Bronchocentric granulomatosis is characterized by peribronchial and peribronchiolar necrotizing granulomatous inflammation. Destruction of airway walls and adjacent parenchyma leads to granulomatous replacement of mucosa and submucosa by palisading, epithelioid, and multinucleated histiocytes. Bronchocentric granulomatosis does not typically involve the pulmonary arteries.

Treatment Glucocorticoids are the treatment of choice, often with an excellent outcome, although recurrences may occur as therapy is tapered or stopped.

GLOBAL CONSIDERATIONS



Limited epidemiologic data exist describing the prevalence or incidence of ILD in the general population. With a few exceptions, e.g., sarcoidosis and certain occupational and environmental exposures, there appear to be no significant differences in the prevalence or incidence of ILD among various populations. For sarcoidosis, there are important environmental, racial, and genetic differences (Chap. 390).

316 Disorders of the Pleura

Richard W. Light

PLEURAL EFFUSION

The pleural space lies between the lung and the chest wall and normally contains a very thin layer of fluid, which serves as a coupling system. A pleural effusion is present when there is an excess quantity of fluid in the pleural space.

Etiology Pleural fluid accumulates when pleural fluid formation exceeds pleural fluid absorption. Normally, fluid enters the pleural space from the capillaries in the parietal pleura and is removed via the lymphatics in the parietal pleura. Fluid also can enter the pleural space from the interstitial spaces of the lung via the visceral pleura or from the peritoneal cavity via small holes in the diaphragm. The lymphatics have the capacity to absorb 20 times more fluid than is formed normally. Accordingly, a pleural effusion may develop when there is excess pleural fluid formation (from the interstitial spaces of the lung, the parietal pleura, or the peritoneal cavity) or when there is decreased fluid removal by the lymphatics.

Diagnostic Approach Patients suspected of having a pleural effusion should undergo chest imaging to diagnose its extent. Chest ultrasound has replaced the lateral decubitus x-ray in the evaluation of suspected pleural effusions and as a guide to thoracentesis. When a patient is found to have a pleural effusion, an effort should be made to determine the cause (Fig. 316-1). The first step is to determine whether the effusion is a transudate or an exudate. A *transudative pleural effusion* occurs when *systemic factors* that influence the formation and absorption of pleural fluid are altered. The leading causes of transudative pleural effusions in the United States are left ventricular failure and cirrhosis. An *exudative pleural effusion* occurs when *local factors* that influence the formation and absorption of pleural fluid are altered. The leading causes of exudative pleural effusions are bacterial pneumonia, malignancy, viral infection, and pulmonary embolism. The primary reason for making this differentiation is that additional diagnostic procedures are indicated with exudative effusions to define the cause of the local disease.

Transudative and exudative pleural effusions are distinguished by measuring the lactate dehydrogenase (LDH) and protein levels in the pleural fluid. Exudative pleural effusions meet at least one of the following criteria, whereas transudative pleural effusions meet none:

1. Pleural fluid protein/serum protein >0.5
2. Pleural fluid LDH/serum LDH >0.6
3. Pleural fluid LDH more than two-thirds the normal upper limit for serum

These criteria misidentify ~25% of transudates as exudates. If one or more of the exudative criteria are met and the patient is clinically thought to have a condition producing a transudative effusion, the difference between the protein levels in the serum and the pleural fluid should be measured. If this gradient is >31 g/L (3.1 g/dL), the exudative categorization by these criteria can be ignored because almost all such patients have a transudative pleural effusion.

If a patient has an exudative pleural effusion, the following tests on the pleural fluid should be obtained: description of the appearance of the fluid, glucose level, differential cell count, microbiologic studies, and cytology.

Effusion Due to Heart Failure The most common cause of pleural effusion is left ventricular failure. The effusion occurs because the increased amounts of fluid in the lung interstitial spaces exit in part across the visceral pleura; this overwhelms the capacity of the lymphatics in the parietal pleura to remove fluid. In patients with heart failure,

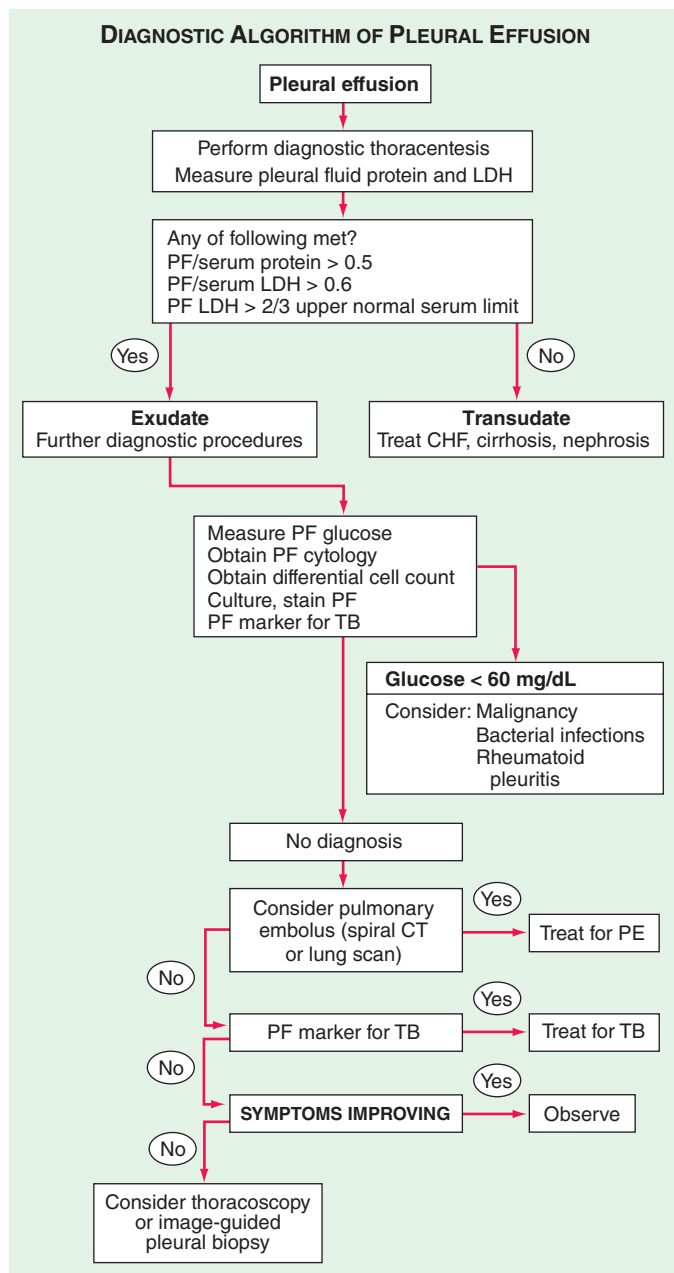


FIGURE 316-1 Approach to the diagnosis of pleural effusions.

CHF, congestive heart failure; CT, computed tomography; LDH, lactate dehydrogenase; PE, pulmonary embolism; PF, pleural fluid; TB, tuberculosis.

a diagnostic thoracentesis should be performed if the effusions are not bilateral and comparable in size, if the patient is febrile, or if the patient has pleuritic chest pain to verify that the patient has a transudative effusion. Otherwise the patient's heart failure is treated. If the effusion persists despite therapy, a diagnostic thoracentesis should be performed. A pleural fluid N-terminal pro-brain natriuretic peptide (NT-proBNP) >1500 pg/mL is virtually diagnostic that the effusion is secondary to congestive heart failure.

Hepatic Hydrothorax Pleural effusions occur in ~5% of patients with cirrhosis and ascites. The predominant mechanism is the direct movement of peritoneal fluid through small openings in the diaphragm into the pleural space. The effusion is usually right-sided and frequently is large enough to produce severe dyspnea.

Parapneumonic Effusion Parapneumonic effusions are associated with bacterial pneumonia, lung abscess, or bronchiectasis and are probably the most common cause of exudative pleural effusion in the United States. *Empyema* refers to a grossly purulent effusion.

Patients with aerobic bacterial pneumonia and pleural effusion present with an acute febrile illness consisting of chest pain, sputum production, and leukocytosis. Patients with anaerobic infections present with a subacute illness with weight loss, a brisk leukocytosis, mild anemia, and a history of some factor that predisposes them to aspiration.

The possibility of a parapneumonic effusion should be considered whenever a patient with bacterial pneumonia is initially evaluated. The presence of free pleural fluid can be demonstrated with a lateral decubitus radiograph, computed tomography (CT) of the chest, or ultrasound. If the free fluid separates the lung from the chest wall by >10 mm, a therapeutic thoracentesis should be performed. Factors indicating the likely need for a procedure more invasive than a thoracentesis (in increasing order of importance) include the following:

1. Loculated pleural fluid
2. Pleural fluid pH <7.20
3. Pleural fluid glucose <3.3 mmol/L (<60 mg/dL)
4. Positive Gram stain or culture of the pleural fluid
5. Presence of gross pus in the pleural space

If the fluid recurs after the initial therapeutic thoracentesis and if any of these characteristics are present, a repeat thoracentesis should be performed. If the fluid cannot be completely removed with the therapeutic thoracentesis, consideration should be given to inserting a chest tube and instilling the combination of a fibrinolytic agent (e.g., tissue plasminogen activator, 10 mg) and deoxyribonuclease (5 mg) or performing a thoracoscopy with the breakdown of adhesions. Decortication should be considered when these measures are ineffective.

Effusion Secondary to Malignancy Malignant pleural effusions secondary to metastatic disease are the second most common type of exudative pleural effusion. The three tumors that cause ~75% of all malignant pleural effusions are lung carcinoma, breast carcinoma, and lymphoma. Most patients complain of dyspnea, which is frequently out of proportion to the size of the effusion. The pleural fluid is an exudate, and its glucose level may be reduced if the tumor burden in the pleural space is high.

The diagnosis usually is made via cytology of the pleural fluid. If the initial cytologic examination is negative, thoracoscopy is the best next procedure if malignancy is strongly suspected. At the time of thoracoscopy, a procedure such as pleural abrasion should be performed to effect a pleurodesis. An alternative to thoracoscopy is CT- or ultrasound-guided needle biopsy of pleural thickening or nodules. Patients with a malignant pleural effusion are treated symptomatically for the most part, since the presence of the effusion indicates disseminated disease and most malignancies associated with pleural effusion are not curable with chemotherapy. The only symptom that can be attributed to the effusion itself is dyspnea. If the patient's lifestyle is compromised by dyspnea and if the dyspnea is relieved with a therapeutic thoracentesis, one of the following procedures should be considered: (1) insertion of a small indwelling catheter or (2) tube thoracostomy with the instillation of a sclerosing agent such as doxycycline (500 mg).

Mesothelioma Malignant mesotheliomas are primary tumors that arise from the mesothelial cells that line the pleural cavities; most are related to asbestos exposure. Patients with mesothelioma present with chest pain and shortness of breath. The chest radiograph reveals a pleural effusion, generalized pleural thickening, and a shrunken hemithorax. The diagnosis is usually established with image-guided needle biopsy or thoracoscopy.

Effusion Secondary to Pulmonary Embolization The diagnosis most commonly overlooked in the differential diagnosis of a patient with an undiagnosed pleural effusion is pulmonary embolism. Dyspnea is the most common symptom. The pleural fluid is almost always an exudate. The diagnosis is established by spiral CT scan or pulmonary arteriography (Chap. 300). Treatment of a patient with a pleural effusion secondary to pulmonary embolism is the same as it is for any patient with pulmonary emboli. If the pleural effusion increases in size after anticoagulation, the patient probably has recurrent emboli or another complication, such as a hemothorax or a pleural infection.

1718 Tuberculous Pleuritis (See also Chap. 202) In many parts of the world, the most common cause of an exudative pleural effusion is tuberculosis (TB), but tuberculous effusions are relatively uncommon in the United States. Tuberculous pleural effusions usually are associated with primary TB and are thought to be due primarily to a hypersensitivity reaction to tuberculous protein in the pleural space. Patients with tuberculous pleuritis present with fever, weight loss, dyspnea, and/or pleuritic chest pain. The pleural fluid is an exudate with predominantly small lymphocytes. The diagnosis is established by demonstrating high levels of TB markers in the pleural fluid (adenosine deaminase >40 IU/L or interferon γ >140 pg/mL). Alternatively, the diagnosis can be established by culture of the pleural fluid, needle biopsy of the pleura, or thoracoscopy. The recommended treatments of pleural and pulmonary TB are identical (Chap. 202).

Effusion Secondary to Viral Infection Viral infections are probably responsible for a sizable percentage of undiagnosed exudative pleural effusions. In many series, no diagnosis is established for ~20% of exudative effusions, and these effusions resolve spontaneously with no long-term residua. The importance of these effusions is that one should not be too aggressive in trying to establish a diagnosis for the undiagnosed effusion, particularly if the patient is improving clinically.

Chylothorax A chylothorax occurs when the thoracic duct is disrupted and chyle accumulates in the pleural space. The most common cause of chylothorax is trauma (most frequently thoracic surgery), but it also may result from tumors in the mediastinum. Patients with chylothorax present with dyspnea, and a large pleural effusion is present on the chest radiograph. Thoracentesis reveals milky fluid, and biochemical analysis reveals a triglyceride level that exceeds 1.2 mmol/L (110 mg/dL). Patients with chylothorax and no obvious trauma should have a lymphangiogram and a mediastinal CT scan to assess the mediastinum for lymph nodes. The treatment of choice for most chylothoraxes is insertion of a chest tube plus the administration of octreotide. If these modalities fail, a pleuroperitoneal shunt should be placed unless the patient has chylous ascites. Alternative treatments are ligation of the thoracic duct and percutaneous transabdominal thoracic duct blockage. Patients with chylothoraxes should not undergo prolonged tube thoracostomy with chest tube drainage because this will lead to malnutrition and immunologic incompetence.

Hemothorax When a diagnostic thoracentesis reveals bloody pleural fluid, a hematocrit should be obtained on the pleural fluid. If the hematocrit is more than one-half of that in the peripheral blood, the patient is considered to have a hemothorax. Most hemothoraxes are the result of trauma; other causes include rupture of a blood vessel or tumor. Most patients with hemothorax should be treated with tube thoracostomy, which allows continuous quantification of bleeding. If the bleeding emanates from a laceration of the pleura, apposition of the two pleural surfaces is likely to stop the bleeding. If the pleural hemorrhage exceeds 200 mL/h, consideration should be given to thoracoscopy or thoracotomy.

Miscellaneous Causes of Pleural Effusion There are many other causes of pleural effusion (Table 316-1). Key features of some of these conditions are as follows: If the pleural fluid amylase level is elevated, the diagnosis of esophageal rupture or pancreatic disease is likely. If the patient is febrile, has predominantly polymorphonuclear cells in the pleural fluid, and has no pulmonary parenchymal abnormalities, an intraabdominal abscess should be considered.

The diagnosis of an asbestos pleural effusion is one of exclusion. Benign ovarian tumors can produce ascites and a pleural effusion (Meigs' syndrome), as can the ovarian hyperstimulation syndrome. Several drugs can cause pleural effusion; the associated fluid is usually eosinophilic. Pleural effusions commonly occur after coronary artery bypass surgery. Effusions occurring within the first weeks are typically left-sided and bloody, with large numbers of eosinophils, and respond to one or two therapeutic thoracenteses. Effusions occurring after the first few weeks are typically left-sided and clear yellow, with predominantly small lymphocytes, and tend to recur. Other medical manipulations that induce pleural effusions include abdominal

TABLE 316-1 DIFFERENTIAL DIAGNOSES OF PLEURAL EFFUSIONS

Transudative Pleural Effusions

1. Congestive heart failure
2. Cirrhosis
3. Nephrotic syndrome
4. Peritoneal dialysis
5. Superior vena cava obstruction
6. Myxedema
7. Urinothorax

Exudative Pleural Effusions

1. Neoplastic diseases
 - a. Metastatic disease
 - b. Mesothelioma
2. Infectious diseases
 - a. Bacterial infections
 - b. Tuberculosis
 - c. Fungal infections
 - d. Viral infections
 - e. Parasitic infections
3. Pulmonary embolization
4. Gastrointestinal disease
 - a. Esophageal perforation
 - b. Pancreatic disease
 - c. Intraabdominal abscesses
 - d. Diaphragmatic hernia
 - e. After abdominal surgery
 - f. Endoscopic variceal sclerotherapy
 - g. After liver transplant
5. Collagen vascular diseases
 - a. Rheumatoid pleuritis
 - b. Systemic lupus erythematosus
 - c. Drug-induced lupus
 - d. Immunoblastic lymphadenopathy
 - e. Sjögren syndrome
 - f. Granulomatosis with polyangiitis (Wegener)
 - g. Churg-Strauss syndrome
6. Post-coronary artery bypass surgery
7. Asbestos exposure
8. Sarcoidosis
9. Uremia
10. Meigs' syndrome
11. Yellow nail syndrome
12. Drug-induced pleural disease
 - a. Nitrofurantoin
 - b. Dantrolene
 - c. Methysergide
 - d. Bromocriptine
 - e. Procarbazine
 - f. Amiodarone
 - g. Dasatinib
13. Trapped lung
14. Radiation therapy
15. Post-cardiac injury syndrome
16. Hemothorax
17. Iatrogenic injury
18. Ovarian hyperstimulation syndrome
19. Pericardial disease
20. Chylothorax

surgery; radiation therapy; liver, lung, or heart transplantation; and the intravascular insertion of central lines.

PNEUMOTHORAX

Pneumothorax is the presence of gas in the pleural space. A *spontaneous pneumothorax* is one that occurs without antecedent trauma to the thorax. A *primary spontaneous pneumothorax* occurs in the absence of underlying lung disease, whereas a *secondary pneumothorax* occurs in its presence. A *traumatic pneumothorax* results from penetrating or nonpenetrating chest injuries. A *tension pneumothorax* is a pneumothorax in which the pressure in the pleural space is positive throughout the respiratory cycle.

Primary Spontaneous Pneumothorax Primary spontaneous pneumothoraxes are usually due to rupture of apical pleural blebs, small cystic spaces that lie within or immediately under the visceral pleura. Primary spontaneous pneumothoraxes occur almost exclusively in smokers; this suggests that these patients have subclinical lung disease. Approximately one-half of patients with an initial primary spontaneous pneumothorax will have a recurrence. The initial recommended treatment for primary spontaneous pneumothorax is simple aspiration. If the lung does not expand with aspiration or if the patient has a recurrent pneumothorax, thoracoscopy with stapling of blebs and pleural abrasion is indicated. Thoracoscopy or thoracotomy with pleural abrasion is almost 100% successful in preventing recurrences.

Secondary Pneumothorax Most secondary pneumothoraxes are due to chronic obstructive pulmonary disease, but pneumothoraxes have been reported with virtually every lung disease. Pneumothorax in patients with lung disease is more life-threatening than it is in normal individuals because of the lack of pulmonary reserve in these patients. Nearly all patients with secondary pneumothorax should be treated with tube thoracostomy. Most should also be treated with thoracoscopy or thoracotomy with the stapling of blebs and pleural abrasion. If the patient is not a good operative candidate or refuses surgery, pleurodesis should be attempted by the intrapleural injection of a sclerosing agent such as doxycycline.

Traumatic Pneumothorax Traumatic pneumothoraxes can result from both penetrating and nonpenetrating chest trauma. Traumatic pneumothoraxes should be treated with tube thoracostomy unless they are very small. If a hemopneumothorax is present, one chest tube should be placed in the superior part of the hemithorax to evacuate the air and another should be placed in the inferior part of the hemithorax to remove the blood. Iatrogenic pneumothorax is a type of traumatic pneumothorax that is becoming more common. The leading causes are transthoracic needle aspiration, thoracentesis, and the insertion of central intravenous catheters. Most can be managed with supplemental oxygen or aspiration, but if these measures are unsuccessful, a tube thoracostomy should be performed.

Tension Pneumothorax This condition usually occurs during mechanical ventilation or resuscitative efforts. The positive pleural pressure is life-threatening both because ventilation is severely compromised and because the positive pressure is transmitted to the mediastinum, resulting in decreased venous return to the heart and reduced cardiac output.

Difficulty in ventilation during resuscitation or high peak inspiratory pressures during mechanical ventilation strongly suggest the diagnosis. The diagnosis is made by physical examination showing an enlarged hemithorax with no breath sounds, hyperresonance to percussion, and shift of the mediastinum to the contralateral side. Tension pneumothorax must be treated as a medical emergency. If the tension in the pleural space is not relieved, the patient is likely to die from inadequate cardiac output or marked hypoxemia. A large-bore needle should be inserted into the pleural space through the second anterior intercostal space. If large amounts of gas escape from the needle after insertion, the diagnosis is confirmed. The needle should be left in place until a thoracostomy tube can be inserted.

317 Disorders of the Mediastinum

Richard W. Light

The mediastinum is the region between the pleural sacs. It is separated into three compartments (Table 317-1). The *anterior mediastinum* extends from the sternum anteriorly to the pericardium and brachiocephalic vessels posteriorly. It contains the thymus gland, the anterior mediastinal lymph nodes, and the internal mammary arteries and veins. The *middle mediastinum* lies between the anterior and posterior mediastina and contains the heart; the ascending and transverse arches of the aorta; the venae cavae; the brachiocephalic arteries and veins; the phrenic nerves; the trachea, the main bronchi, and their contiguous lymph nodes; and the pulmonary arteries and veins. The *posterior mediastinum* is bounded by the pericardium and trachea anteriorly and the vertebral column posteriorly. It contains the descending thoracic aorta, the esophagus, the thoracic duct, the azygos and hemiazygos veins, and the posterior group of mediastinal lymph nodes.

MEDIASTINAL MASSES

The first step in evaluating a mediastinal mass is to place it in one of the three mediastinal compartments, since each has different characteristic lesions (Table 317-1). The most common lesions in the anterior mediastinum are thymomas, lymphomas, teratomatous neoplasms, and thyroid masses. The most common masses in the middle mediastinum are vascular masses, lymph node enlargement from metastases or granulomatous disease, and pleuropericardial and bronchogenic cysts. In the posterior mediastinum, neurogenic tumors, meningoceles, meningomyeloceles, gastroenteric cysts, and esophageal diverticula are commonly found.

Computed tomography (CT) scanning is the most valuable imaging technique for evaluating mediastinal masses and is the only imaging technique that should be done in most instances. Barium studies of the gastrointestinal tract are indicated in many patients with posterior mediastinal lesions, because hernias, diverticula, and achalasia are readily diagnosed in this manner. An iodine-131 scan can efficiently establish the diagnosis of intrathoracic goiter.

A definite diagnosis can be obtained with mediastinoscopy or anterior mediastinotomy in many patients with masses in the anterior or middle mediastinal compartments. A diagnosis can be established without thoracotomy via percutaneous fine-needle aspiration biopsy or endoscopic transesophageal or endobronchial ultrasound-guided biopsy of mediastinal masses in most cases. An alternative way to establish the diagnosis is video-assisted thoracoscopy. In many cases, the diagnosis can be established and the mediastinal mass removed with video-assisted thoracoscopy.

ACUTE MEDIASTITIS

Most cases of acute mediastinitis either are due to esophageal perforation or occur after median sternotomy for cardiac surgery. Patients with esophageal rupture are acutely ill with chest pain and dyspnea due to the mediastinal infection. The esophageal rupture can occur spontaneously or as a complication of esophagoscopy or the insertion of a Blakemore tube. Appropriate treatment consists of exploration of the mediastinum with primary repair of the esophageal tear and drainage of the pleural space and the mediastinum.

The incidence of mediastinitis after median sternotomy is 0.4–5.0%. Patients most commonly present with wound drainage. Other presentations include sepsis and a widened mediastinum. The diagnosis usually is established with mediastinal needle aspiration. Treatment includes immediate drainage, debridement, and parenteral antibiotic therapy, but the mortality rate still exceeds 20%.

CHRONIC MEDIASTITIS

The spectrum of chronic mediastinitis ranges from granulomatous inflammation of the lymph nodes in the mediastinum to fibrosing

TABLE 317-1 THE THREE COMPARTMENTS OF THE MEDIASTINUM

	Anterior Compartment	Middle Compartment	Posterior Compartment
Anatomical boundaries	Manubrium and sternum anteriorly, pericardium, aorta, and brachiocephalic vessels posteriorly	Anterior mediastinum anteriorly, posterior mediastinum posteriorly	Pericardium and trachea anteriorly; vertebral column posteriorly
Contents	Thymus gland, anterior mediastinal lymph nodes, internal mammary arteries and veins	Pericardium, heart, ascending and transverse arch of aorta, superior and inferior vena cavae, brachiocephalic arteries and veins, phrenic nerves, trachea, and main bronchi and their contiguous lymph nodes, pulmonary arteries, and veins	Descending thoracic aorta, esophagus, thoracic duct, azygos and hemiazygos veins, sympathetic chains, and the posterior group of mediastinal lymph nodes
Common abnormalities	Thymoma, lymphomas, teratomatous neoplasms, thyroid masses, parathyroid masses, mesenchymal tumors, giant lymph node hyperplasia, hernia through foramen of Morgagni	Metastatic lymph node enlargement, granulomatous lymph node enlargement, pleuropericardial cysts, bronchogenic cysts, masses of vascular origin	Neurogenic tumors, meningocele, meningomyelocele, gastroenteric cysts, esophageal diverticula, hernia through foramen of Bochdalek, extramedullary hematopoiesis

mediastinitis. Most cases are due to histoplasmosis or tuberculosis, but sarcoidosis, silicosis, and other fungal diseases are at times causative. Patients with granulomatous mediastinitis are usually asymptomatic. Those with fibrosing mediastinitis usually have signs of compression of a mediastinal structure such as the superior vena cava or large airways, phrenic or recurrent laryngeal nerve paralysis, or obstruction of the pulmonary artery or proximal pulmonary veins. Other than antituberculous therapy for tuberculous mediastinitis, no medical or surgical therapy has been demonstrated to be effective for mediastinal fibrosis.

PNEUMOMEDIASTINUM

In this condition, there is gas in the interstices of the mediastinum. The three main causes are (1) alveolar rupture with dissection of air into the mediastinum; (2) perforation or rupture of the esophagus, trachea, or main bronchi; and (3) dissection of air from the neck or the abdomen into the mediastinum. Typically, there is severe substernal chest pain with or without radiation into the neck and arms. The physical examination usually reveals subcutaneous emphysema in the suprasternal notch and *Hamman's sign*, which is a crunching or clicking noise synchronous with the heartbeat and is best heard in the left lateral decubitus position. The diagnosis is confirmed with the chest radiograph. Usually no treatment is required, but the mediastinal air will be absorbed faster if the patient inspires high concentrations of oxygen. If mediastinal structures are compressed, the compression can be relieved with needle aspiration.

Diseases that alter \dot{V}_{CO_2} are often acute (e.g., sepsis, burns, or pyrexia), and their contribution to ventilatory abnormalities and/or respiratory failure is reviewed elsewhere. Chronic ventilatory disorders typically involve inappropriate levels of minute ventilation or increased dead space fraction. Characterization of these disorders requires a review of the normal respiratory cycle.

The spontaneous cycle of inspiration and expiration is automatically generated in the brainstem. Two groups of neurons located within the medulla are particularly important: the dorsal respiratory group (DRG) and the ventral respiratory column (VRC). These neurons have widespread projections including the descending projections into the contralateral spinal cord where they perform many functions. They initiate activity in the phrenic nerve/diaphragm, project to the upper airway muscle groups and spinal respiratory neurons, and innervate the intercostal and abdominal muscles that participate in normal respiration. The DRG acts as the initial integration site for many of the afferent nerves relaying information about Pa_{O_2} , Pa_{CO_2} , pH, and blood pressure from the carotid and aortic chemoreceptors and baroreceptors to the central nervous system (CNS). In addition, the vagus nerve relays information from stretch receptors and juxtapulmonary-capillary receptors in the lung parenchyma and chest wall to the DRG. The respiratory rhythm is generated within the VRC as well as the more rostrally located parafacial respiratory group (pFRG), which is particularly important for the generation of active expiration. One particularly important area within the VRC is the so-called pre-Bötzinger complex. This area is responsible for the generation of various forms of inspiratory activity, and lesioning of the pre-Bötzinger complex leads to the complete cessation of breathing. The neural output of these medullary respiratory networks can be voluntarily suppressed or augmented by input from higher brain centers and the autonomic nervous system. During normal sleep, there is an attenuated response to hypercapnia and hypoxemia, resulting in mild nocturnal hypoventilation that corrects upon awakening.

Once neural input has been delivered to the respiratory pump muscles, normal gas exchange requires an adequate amount of respiratory muscle strength to overcome the elastic and resistive loads of the respiratory system (Fig. 318-1A) (Chap. 306e). In health, the strength of the respiratory muscles readily accomplishes this, and normal respiration continues indefinitely. Reduction in respiratory drive or neuromuscular competence or substantial increase in respiratory load can diminish minute ventilation, resulting in hypercapnia (Fig. 318-1B). Alternatively, if normal respiratory muscle strength is coupled with excessive respiratory drive, then alveolar hyperventilation ensues and leads to hypocapnia (Fig. 318-1C).

HYPOVENTILATION

CLINICAL FEATURES

Diseases that reduce minute ventilation or increase dead space fall into four major categories: parenchymal lung and chest wall disease, sleep-disordered breathing, neuromuscular disease, and respiratory drive disorders (Fig. 318-1B). The clinical manifestations of hypoventilation

318 Disorders of Ventilation

John F. McConville, Babak Mokhlesi, Julian Solway

DEFINITION AND PHYSIOLOGY

In health the arterial level of carbon dioxide (Pa_{CO_2}) is maintained between 37 and 43 mmHg at sea level. All disorders of ventilation result in abnormal measurements of Pa_{CO_2} . This chapter reviews chronic ventilatory disorders.

The continuous production of CO_2 by cellular metabolism necessitates its efficient elimination by the respiratory system. The relationship between CO_2 production and Pa_{CO_2} is described by the equation: $Pa_{CO_2} = (k) (\dot{V}_{CO_2})/\dot{V}_A$, where \dot{V}_{CO_2} represents the carbon dioxide production, k is a constant, and \dot{V}_A is fresh gas alveolar ventilation (Chap. 306e). \dot{V}_A can be calculated as minute ventilation $\times (1 - V_d/V_t)$, where the dead space fraction V_d/V_t represents the portion of a tidal breath that remains within the conducting airways at the conclusion of inspiration and so does not contribute to alveolar ventilation. As such, all disturbances of Pa_{CO_2} must reflect altered CO_2 production, minute ventilation, or dead space fraction.

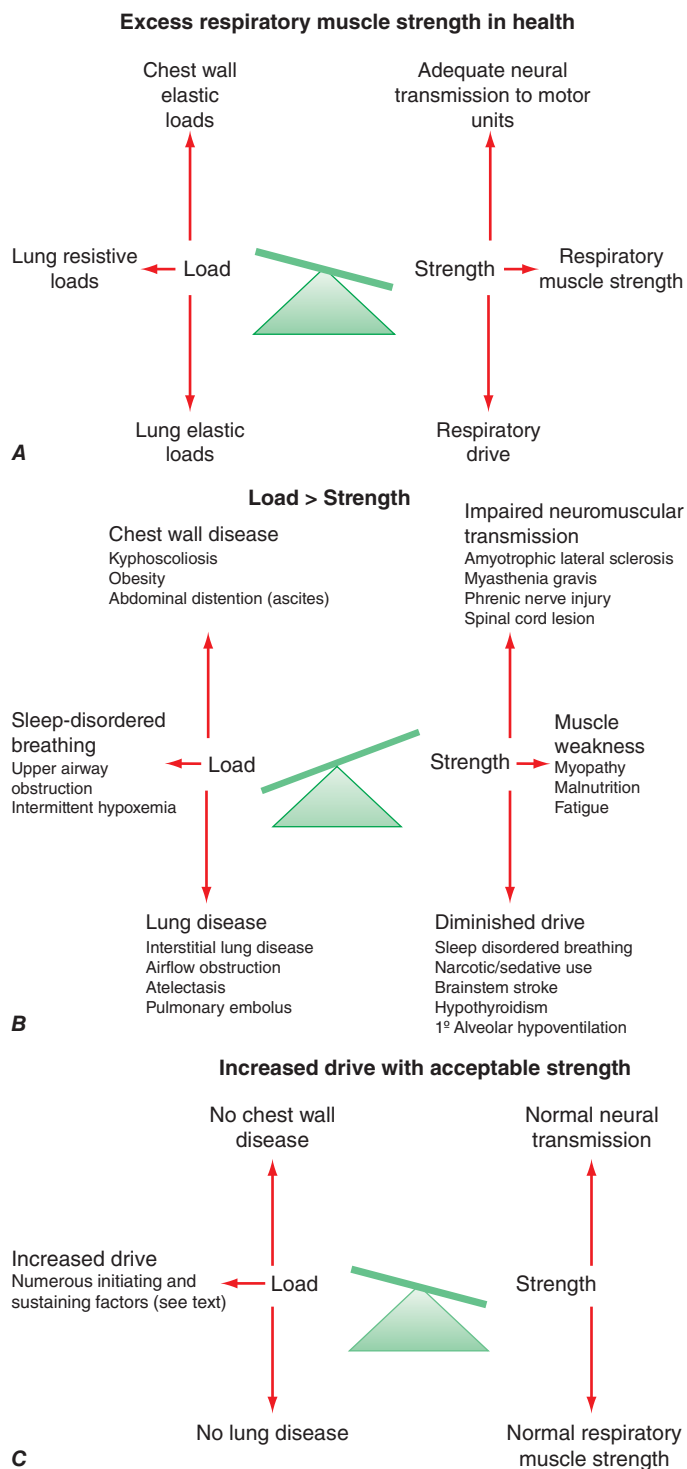


FIGURE 318-1 Examples of balance between respiratory system strength and load. **A.** Excess respiratory muscle strength in health. **B.** Load greater than strength. **C.** Increased drive with acceptable strength.

syndromes are nonspecific (Table 318-1) and vary depending on the severity of hypoventilation, the rate at which hypercapnia develops, the degree of compensation for respiratory acidosis, and the underlying disorder. Patients with parenchymal lung or chest wall disease typically present with shortness of breath and diminished exercise tolerance. Episodes of increased dyspnea and sputum production are hallmarks of obstructive lung diseases such as chronic obstructive pulmonary disease, whereas progressive dyspnea and cough are common in interstitial lung diseases. Excessive daytime somnolence, poor-quality sleep, and snoring are common among patients with sleep-disordered breathing. Sleep disturbance and orthopnea are also described in

TABLE 318-1 SIGNS AND SYMPTOMS OF HYPOVENTILATION

Dyspnea during activities of daily living
Orthopnea in diseases affecting diaphragm function
Poor-quality sleep
Daytime hypersomnolence
Early morning headaches
Anxiety
Impaired cough in neuromuscular diseases

neuromuscular disorders. As neuromuscular weakness progresses, the respiratory muscles, including the diaphragm, are placed at a mechanical disadvantage in the supine position due to the upward movement of the abdominal contents. New-onset orthopnea is frequently a sign of reduced respiratory muscle force generation. More commonly, however, extremity weakness or bulbar symptoms develop prior to sleep disturbance in neuromuscular diseases such as amyotrophic lateral sclerosis (ALS) or muscular dystrophy. Patients with respiratory drive disorders do not have symptoms distinguishable from other causes of chronic hypoventilation.

The clinical course of patients with chronic hypoventilation from neuromuscular or chest wall disease follows a characteristic sequence: an asymptomatic stage where daytime P_{aO_2} and P_{aCO_2} are normal followed by nocturnal hypoventilation, initially during rapid eye movement (REM) sleep and later in non-REM sleep. Finally, if vital capacity drops further, daytime hypercapnia develops. Symptoms can develop at any point along this time course and often depend on the pace of respiratory muscle functional decline. Regardless of cause, the hallmark of all alveolar hypoventilation syndromes is an increase in alveolar P_{CO_2} (P_{ACO_2}) and therefore in P_{aCO_2} . The resulting respiratory acidosis eventually leads to a compensatory increase in plasma bicarbonate concentration. The increase in P_{ACO_2} results in an obligatory decrease in P_{AO_2} , often resulting in hypoxemia. If severe, the hypoxemia manifests clinically as cyanosis and can stimulate erythropoiesis and thus induce secondary erythrocytosis. The combination of chronic hypoxemia and hypercapnia may also induce pulmonary vasoconstriction, leading eventually to pulmonary hypertension, right ventricular hypertrophy, and right heart failure.

DIAGNOSIS

Elevated plasma bicarbonate in the absence of volume depletion is suggestive of hypoventilation. An arterial blood gas demonstrating elevated P_{aCO_2} with a normal pH confirms chronic alveolar hypoventilation. The subsequent evaluation to identify an etiology should initially focus on whether the patient has lung disease or chest wall abnormalities. Physical examination, imaging studies (chest x-ray and/or computed tomography [CT] scan), and pulmonary function tests are sufficient to identify most lung/chest wall disorders leading to hypercapnia. If these evaluations are unrevealing, then the clinician should screen for obesity hypoventilation syndrome (OHS), the most frequent sleep disorder leading to chronic hypoventilation, which is typically accompanied by obstructive sleep apnea (OSA). Several screening tools have been developed to identify patients at risk for OSA. The Berlin Questionnaire has been validated in a primary care setting and identifies patients likely to have OSA. The Epworth Sleepiness Scale (ESS) and the STOP-Bang questionnaires have not been validated in outpatient primary care settings but are quick and easy to use. The ESS measures daytime sleepiness, with a score of ≥ 10 identifying individuals who warrant additional investigation. The STOP-Bang survey has been used in preoperative clinics to identify patients at risk of having OSA. In this population, it has 93% sensitivity and 90% negative predictive value.

If the ventilatory apparatus (lungs, airways, chest wall) is not responsible for chronic hypercapnia, then the focus should shift to respiratory drive and neuromuscular disorders. There is an attenuated increase in minute ventilation in response to elevated C_{O_2} and/or low O_2 in respiratory drive disorders. These diseases are difficult to diagnose and should be suspected when patients with hypercapnia are

1722 found to have normal respiratory muscle strength, normal pulmonary function, and normal alveolar-arterial P_{O_2} difference. Hypoventilation is more marked during sleep in patients with respiratory drive defects, and polysomnography often reveals central apneas, hypopneas, or hypoventilation. Brain imaging (CT scan or magnetic resonance imaging [MRI]) can sometimes identify structural abnormalities in the pons or medulla that result in hypoventilation. Chronic narcotic use or significant hypothyroidism can depress the central respiratory drive and lead to chronic hypercapnia as well.

Respiratory muscle weakness has to be profound before lung volumes are compromised and hypercapnia develops. Typically physical examination reveals decreased strength in major muscle groups prior to the development of hypercapnia. Measurement of maximum inspiratory and expiratory pressures or forced vital capacity (FVC) can be used to monitor for respiratory muscle involvement in diseases with progressive muscle weakness. These patients also have increased risk for sleep-disordered breathing, including hypopneas, central and obstructive apneas, and hypoxemia. Nighttime oximetry and capnometry during polysomnography are helpful in better characterizing sleep disturbances in this patient population.

TREATMENT HYPOVENTILATION

Nocturnal noninvasive positive-pressure ventilation (NIPPV) has been used successfully in the treatment of hypoventilation and apneas, both central and obstructive, in patients with neuromuscular and chest wall disorders. Nocturnal NIPPV has been shown to improve daytime hypercapnia, prolong survival, and improve health-related quality of life when daytime hypercapnia is documented. ALS guidelines recommend consideration of nocturnal NIPPV if symptoms of hypoventilation exist and one of the following criteria is present: $P_{aCO_2} \geq 45$ mmHg; nocturnal oximetry demonstrates oxygen saturation $\leq 88\%$ for 5 consecutive min; maximal inspiratory pressure < 60 cmH₂O; FVC $< 50\%$ predicted; or sniff nasal pressure < 40 cmH₂O. However, at present, there is inconclusive evidence to support preemptive nocturnal NIPPV use in all patients with neuromuscular and chest wall disorders who demonstrate nocturnal but not daytime hypercapnia. Nevertheless, at some point, the institution of full-time ventilatory support with either pressure or volume-preset modes is required in progressive neuromuscular disorders. There is less evidence to direct the timing of this decision, but ventilatory failure requiring mechanical ventilation and chest infections related to ineffective cough are frequent triggers for the institution of full-time ventilatory support.

Treatment of chronic hypoventilation from lung or neuromuscular diseases should be directed at the underlying disorder. Pharmacologic agents that stimulate respiration, such as medroxyprogesterone and acetazolamide, have been poorly studied in chronic hypoventilation and should not replace treatment of the underlying disease process. Regardless of the cause, excessive metabolic alkalosis should be corrected, because plasma bicarbonate levels elevated out of proportion for the degree of chronic respiratory acidosis can result in additional hypoventilation. When indicated, administration of supplemental oxygen is effective in attenuating hypoxemia, polycythemia, and pulmonary hypertension. However, in some patients, supplemental oxygen can worsen hypercapnia.

Phrenic nerve or diaphragm pacing is a potential therapy for patients with hypoventilation from high cervical spinal cord lesions or respiratory drive disorders. Prior to surgical implantation, patients should have nerve conduction studies to ensure normal bilateral phrenic nerve function. Small case series suggest that effective diaphragmatic pacing can improve quality of life in these patients.

HYPOVENTILATION SYNDROMES

OBESITY HYPOVENTILATION SYNDROME

The diagnosis of OHS requires body mass index (BMI) ≥ 30 kg/m² and chronic daytime alveolar hypoventilation, defined as P_{aCO_2}

≥ 45 mmHg at sea level in the absence of other known causes of hypercapnia. In almost 90% of cases, the sleep-disordered breathing is in the form of OSA. Several international studies in different populations confirm that the overall prevalence of OSA syndrome, defined by an apnea-hypopnea index (AHI) ≥ 5 and daytime sleepiness, is approximately 3–4% in middle-aged men and 2% in middle-aged women. Thus, the population at risk for the development of OHS continues to rise as the worldwide obesity epidemic persists. Although no population-based prevalence studies of OHS have been performed, some estimates suggest there may be as many as 500,000 individuals with OHS in the United States.

Some, but not all, studies suggest that severe obesity (BMI > 40 kg/m²) and severe OSA (AHI > 30 events per hour) are risk factors for the development of OHS. The pathogenesis of hypoventilation in these patients is the result of multiple physiologic variables and conditions including OSA, increased work of breathing, respiratory muscle impairment, ventilation-perfusion mismatching, and depressed central ventilatory responsiveness to hypoxemia and hypercapnia. These defects in central respiratory drive often improve with treatment, which suggests that decreased ventilatory responsiveness is a consequence rather than a primary cause of OHS. The treatment of OHS is similar to that for OSA: weight reduction and nocturnal NIPPV. There is evidence that weight loss alone lowers P_{aCO_2} in patients with OHS. However, treatment with NIPPV should never be delayed while the patient attempts to lose weight. Continuous positive airway pressure (CPAP) improves daytime hypercapnia and hypoxemia in more than half of patients with OHS and concomitant OSA. Bilevel positive airway pressure should be reserved for patients not able to tolerate high levels of CPAP support or patients who remain hypoxemic despite resolution of obstructive respiratory events. NIPPV with bilevel positive airway pressure should be strongly considered if hypercapnia persists after several weeks of CPAP therapy with objectively proven adherence. Patients with OHS and no evidence of OSA are typically started on bilevel positive airway pressure, as are patients presenting with acute decompensated OHS. Finally, comorbid conditions that impair ventilation, such as chronic obstructive pulmonary disease, should be aggressively treated in conjunction with coexisting OHS.

CENTRAL HYPOVENTILATION SYNDROME

This syndrome can present later in life or in the neonatal period where it is often called Ondine's curse or congenital central hypoventilation syndrome. Abnormalities in the gene encoding PHOX2b, a transcription factor with a role in neuronal development, have been implicated in the pathogenesis of congenital central hypoventilation syndrome. Regardless of the age of onset, these patients have absent respiratory response to hypoxia or hypercapnia, mildly elevated P_{aCO_2} while awake, and markedly elevated P_{aCO_2} during non-REM sleep. Interestingly these patients are able to augment their ventilation and "normalize" P_{aCO_2} during exercise and during REM sleep. These patients typically require NIPPV or mechanical ventilation as therapy and should be considered for phrenic nerve or diaphragmatic pacing at centers with experience performing these procedures.

HYPERVENTILATION

CLINICAL FEATURES

Hyperventilation is defined as ventilation in excess of metabolic requirements (CO_2 production) leading to a reduction in P_{aCO_2} . The physiology of patients with chronic hyperventilation is poorly understood, and there is no typical clinical presentation. Symptoms can include dyspnea, paresthesias, tetany, headache, dizziness, visual disturbances, and atypical chest pain. Because symptoms can be so diverse, patients with chronic hyperventilation present to a variety of health care providers, including internists, neurologists, psychologists, psychiatrists, and pulmonologists.

It is helpful to think of hyperventilation as having initiating and sustaining factors. Some investigators believe that an initial event leads to increased alveolar ventilation and a drop in P_{aCO_2}

to ~20 mmHg. The ensuing onset of chest pain, breathlessness, paresthesia, or altered consciousness can be alarming. The resultant increase in minute volume to relieve these acute symptoms only serves to exacerbate symptoms that are often misattributed by the patient and health care workers to cardiopulmonary disorders. An unrevealing evaluation for causes of these symptoms often results in patients being anxious and fearful of additional attacks. It is important to note that **anxiety disorders and panic attacks are not synonymous with hyperventilation**. Anxiety disorders can be both an initiating and sustaining factor in the pathogenesis of chronic hyperventilation, but these are not necessary for the development of chronic hypocapnia.

DIAGNOSIS

Respiratory symptoms associated with acute hyperventilation can be the initial manifestation of systemic illnesses such as diabetic ketoacidosis. Causes of acute hyperventilation need to be excluded before a diagnosis of chronic hyperventilation is considered. Arterial blood gas sampling that demonstrates a compensated respiratory alkalosis with a near normal pH, low P_{aCO_2} , and low calculated bicarbonate is necessary to confirm chronic hyperventilation. Other causes of respiratory alkalosis, such as mild asthma, need to be diagnosed and treated before chronic hyperventilation can be considered. A high index of suspicion is required because increased minute ventilation can be difficult to detect on physical examination. Once chronic hyperventilation is established, a sustained 10% increase in alveolar ventilation is enough to perpetuate hypocapnia. This increase can be accomplished with subtle changes in the respiratory pattern, such as occasional sigh breaths or yawning two to three times per minute.

TREATMENT HYPERVENTILATION

There are few well-controlled treatment studies of chronic hyperventilation due to its diverse features and the lack of a universally accepted diagnostic process. Clinicians often spend considerable time identifying initiating factors, excluding alternative diagnoses, and discussing the patient's concerns and fears. In some patients, reassurance and frank discussion about hyperventilation can be liberating. Identifying and eliminating habits that perpetuate hypocapnia, such as frequent yawning or sigh breathing, can be helpful. Some evidence suggests that breathing exercises and diaphragmatic retraining may be beneficial for some patients. The evidence for using medications to treat hyperventilation is scant. Beta blockers may be helpful in patients with sympathetically mediated symptoms such as palpitations and tremors.

ACKNOWLEDGMENT

We would like to acknowledge Eliot A. Phillipson for earlier versions of this chapter and Jan-Marino Ramirez for his careful critique and helpful suggestions.

319 Sleep Apnea

Andrew Wellman, Susan Redline

Obstructive sleep apnea/hypopnea syndrome (OSAHS) and central sleep apnea (CSA) are both classified as sleep-related breathing disorders. OSAHS and CSA share some risk factors and physiological bases but also have unique features. Each disorder is associated with impaired ventilation during sleep and disruption of sleep, and each diagnosis requires careful elicitation of the patient's history, physical examination, and physiological testing. OSAHS, the more common disorder, causes daytime sleepiness, impairs daily function, and is a

major contributor to cardiovascular disease in adults and to behavioral problems in children. CSA is less common and may occur in combination with obstructive sleep apnea, as a primary condition, or secondary to a medical condition or medication. CSA impairs overnight gas exchange and may result in symptoms of either insomnia or excessive sleepiness.

OBSTRUCTIVE SLEEP APNEA/HYPOPNEA SYNDROME (OSAHS)

Definition OSAHS is defined on the basis of nocturnal and daytime symptoms as well as sleep study findings. Diagnosis requires the patient to have (1) either symptoms of nocturnal breathing disturbances (snoring, snorting, gasping, or breathing pauses during sleep) or daytime sleepiness or fatigue that occurs despite sufficient opportunities to sleep and is unexplained by other medical problems; and (2) five or more episodes of obstructive apnea or hypopnea per hour of sleep (the apnea-hypopnea index [AHI], calculated as the number of episodes divided by the number of hours of sleep) documented during a sleep study. OSAHS also may be diagnosed in the absence of symptoms if the AHI is above 15. Each episode of apnea or hypopnea represents a reduction in breathing for at least 10 sec. OSAHS is often identified when associated with a $\geq 3\%$ drop in oxygen saturation and/or a brain cortical arousal. OSAHS severity is based on the frequency of breathing disturbances (AHI), the amount of oxygen desaturation with respiratory events, the duration of apneas and hypopneas, the degree of sleep fragmentation, and the level of daytime sleepiness.

Pathophysiology During inspiration, intraluminal pharyngeal pressure becomes increasingly negative, creating a "suctioning" force. Because the pharyngeal airway has no bone or cartilage, airway patency is dependent on the stabilizing influence of the pharyngeal dilator muscles. Although these muscles are continuously activated during wakefulness, neuromuscular output declines with sleep onset. In patients with a collapsible airway, the reduction in neuromuscular output results in transient episodes of pharyngeal collapse (manifesting as an "apnea") or near collapse (manifesting as a "hypopnea"). The episodes of collapse are terminated when ventilatory reflexes are activated and cause arousal, thus stimulating an increase in neuromuscular activity and opening of the airway. The airway may collapse at various levels: the soft palate (most common), tongue base, lateral pharyngeal walls, and/or epiglottis (Fig. 319-1). OSAHS may be most

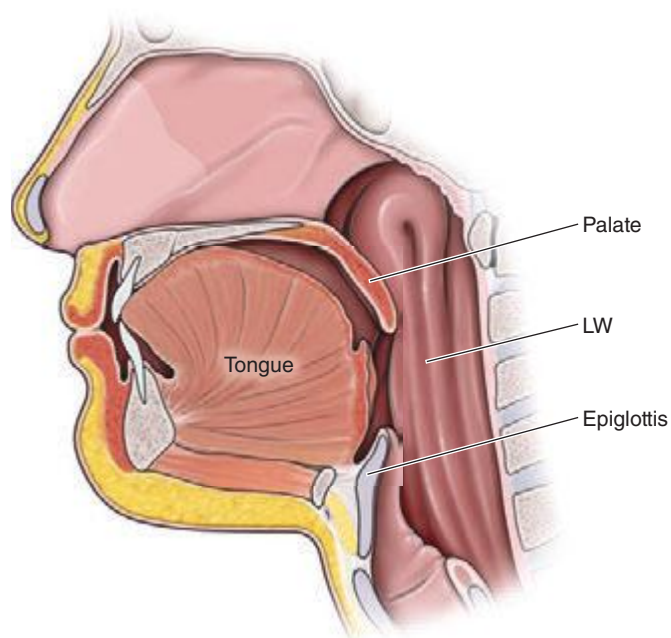


FIGURE 319-1 Common sites of airway collapse. For example, the palate, tongue, and/or epiglottis (Ep) can be posteriorly displaced, and the lateral pharyngeal walls (LW) can collapse.

1724 severe during REM (rapid eye movement) sleep, when neuromuscular output to the skeletal muscles is particularly low, and in the supine position due to gravitational forces.

Individuals with a small pharyngeal lumen require relatively high levels of neuromuscular innervation to maintain patency during wakefulness and thus are predisposed to excessive airway collapsibility during sleep. The airway lumen may be narrowed with enlargement of soft tissue structures (tongue, palate, and uvula) due to fat deposition, increased lymphoid tissue, or genetic variation. Craniofacial factors such as mandibular retroposition or micrognathia, reflecting genetic variation or developmental influences, also can reduce lumen dimensions. In addition, lung volumes influence the caudal traction on the pharynx and consequently the stiffness of the pharyngeal wall. Accordingly, low lung volume in the recumbent position, which is particularly pronounced in the obese, contributes to collapse. A high degree of nasal resistance (e.g., due to nasal septal deviation or polyps) can contribute to airway collapse by increasing the negative intraluminal suction pressure. High-level nasal resistance also may trigger mouth opening during sleep, which breaks the seal between the tongue and the teeth and allows the tongue to fall posteriorly and occlude the airway.

Pharyngeal muscle activation is integrally linked to ventilatory drive. Thus, factors related to ventilatory control, particularly ventilatory sensitivity, arousal threshold, and neuromuscular responses to CO₂, contribute to the pathogenesis of OSAHS. A buildup in CO₂ during sleep activates both the diaphragm and the pharyngeal muscles, which stiffen the upper airway and can counteract inspiratory suction pressures and maintain airway patency to an extent that depends on the anatomic predisposition to collapse. However, pharyngeal collapse can occur when the ventilatory control system is overly sensitive to CO₂, with resultant wide fluctuations in ventilation and ventilatory drive and in upper airway instability. Moreover, increasing levels of CO₂ during sleep result in central nervous system arousal, causing the individual to move from a deeper to a lighter level of sleep or to awaken. A low arousal threshold (i.e., awaken to a low level of CO₂ or ventilatory drive) can preempt the CO₂-mediated process of pharyngeal muscle compensation and prevent airway stabilization. A high arousal threshold, conversely, may prevent appropriate termination of apneas, prolonging apnea duration and oxyhemoglobin desaturation severity. Finally, any impairment in the ability of the muscles to compensate during sleep can contribute to collapse of the pharynx. The relative contributions of risk factors vary among individuals. Approaches to the measurement of these factors in clinical settings, with consequent enhancement of “personalized” therapeutic interventions, are being actively investigated.

Risk Factors and Prevalence The major risk factors for OSAHS are obesity and male sex. Additional risk factors include mandibular retrognathia and micrognathia, a positive family history of OSAHS, genetic syndromes that reduce upper airway patency (e.g., Down syndrome, Treacher-Collins syndrome), adenotonsillar hypertrophy (especially in children), menopause (in women), and various endocrine syndromes (e.g., acromegaly, hypothyroidism).

Approximately 40–60% of cases of OSAHS are attributable to excess weight. Obesity predisposes to OSAHS through the narrowing effects of upper airway fat on the pharyngeal lumen. Obesity also reduces chest wall compliance and decreases lung volumes, resulting in a loss of caudal traction on upper airway structures. Obese individuals are at a fourfold or greater risk for OSAHS than their normal-weight counterparts. A 10% weight gain is associated with a >30% increase in AHI. Even modest weight loss or weight gain can influence the risk and severity of OSAHS. However, the absence of obesity does not exclude this diagnosis.

The prevalence of OSAHS is two- to fourfold higher among men than among women. Factors that predispose men to OSAHS include android patterns of obesity (resulting in upper-airway fat deposition) and relatively great pharyngeal length, which exacerbates collapsibility. Premenopausal women are relatively protected from OSAHS by the influence of sex hormones on ventilatory drive. The decline in sex

differences in older age is associated with an increased OSAHS prevalence in women after menopause.

Variations in craniofacial morphology that reduce the size of the posterior airway space increase OSAHS risk. The contribution of hard-tissue structural features to OSAHS is most evident in nonobese patients. Identification of features such as retrognathia can influence therapeutic decision-making.

OSAHS has a strong genetic basis, as evidenced by its significant familial aggregation and heritability. For a first-degree relative of a patient with OSAHS, the odds ratio of having OSAHS is approximately twofold higher than that for someone without an affected relative.

OSAHS prevalence varies with age, from 2–15% among middle-aged adults to >20% among elderly individuals. There is a peak due to lymphoid hypertrophy among children between the ages of 3 and 8 years; with airway growth and lymphoid tissue regression during later childhood, prevalence declines. Then, as obesity prevalence increases in middle life and women enter menopause, OSAHS again increases.

The prevalence of OSAHS may be especially high among patients with diabetes or hypertension. Individuals of Asian ancestry appear to be at increased risk of OSAHS at relatively low levels of body mass index, possibly because of the influence of craniofacial risk factors that narrow the nasopharynx. In the United States, African Americans, especially children and young adults, are at higher risk for OSAHS than their Caucasian counterparts. In a majority of adults with OSAHS, the disorder is undiagnosed.

Course of the Disorder The precise onset of OSAHS is usually hard to identify. A person may snore for many years, often beginning in childhood, before OSAHS is identified. Weight gain may precipitate an increase in symptoms, which in turn may lead the patient to pursue an evaluation. OSAHS may become less severe with weight loss, particularly after bariatric surgery. Marked increases and decreases in the AHI are uncommon unless accompanied by weight change.

APPROACH TO THE PATIENT:

Obstructive Sleep Apnea/Hypopnea Syndrome (OSAHS)

An evaluation for OSAHS should be considered in patients with symptoms of OSAHS and one or more risk factors. Screening also should be considered in patients who report symptoms consistent with OSAHS and who are at high risk for OSAHS-related morbidities, such as hypertension, diabetes mellitus, and cardiac and cerebrovascular diseases.

SYMPTOMS AND HISTORY

When possible, a sleep history should be obtained in the presence of a bed partner. Snoring is the most common complaint; however, its absence does not exclude the diagnosis, as pharyngeal collapse may occur without tissue vibration. Gasping or snorting during sleep may also be reported, reflecting termination of individual apneas with abrupt airway opening. Dyspnea is unusual, and its absence generally distinguishes OSAHS from paroxysmal nocturnal dyspnea, nocturnal asthma, and acid reflux with laryngospasm. Patients also may describe frequent awakening or sleep disruption, which is more common among women and older adults. The most common daytime symptom is sleepiness. This symptom can be difficult to elicit and may be hard to distinguish from exercise-related fatigue, deconditioning, and malaise. In contrast to true sleepiness, the latter symptoms generally improve with rest. Other symptoms include a dry mouth, nocturnal heartburn, diaphoresis of the chest and neck, nocturia, morning headaches, trouble concentrating, irritability, and mood disturbances. Several questionnaires that evaluate snoring frequency, self-reported apneas, and daytime sleepiness can facilitate OSAHS screening. The predictive ability of a questionnaire can be enhanced by a consideration of whether the patient is male or has risk factors such as obesity or hypertension.

PHYSICAL FINDINGS

Physical findings often reflect the etiologic factors for the disorder as well as comorbid conditions, particularly vascular disease. On examination, patients may exhibit hypertension and regional (central) obesity, as indicated by a large waist and neck circumference. The oropharynx may reveal a small orifice with crowding due to an enlarged tongue, a low-lying soft palate with a bulky uvula, large tonsils, a high arched palate, and/or micro/retrognathia. Since high-level nasal resistance can increase pharyngeal collapsibility, the nasal cavity should be inspected for polyps, septal deviation, and other signs of obstruction. Because patients with heart failure are at increased risk for both OSAHS and CSA, a careful cardiac examination should be conducted to detect possible left- or right-sided cardiac dysfunction. Evidence of cor pulmonale suggests severe OSAHS or a comorbid cardiopulmonary condition. A neurologic evaluation is needed to evaluate for conditions such as neuromuscular and cerebrovascular diseases, which increase OSAHS risk.

LABORATORY FINDINGS

Diagnostic Findings Since symptoms and signs do not accurately predict the severity of sleep-related breathing disturbances, specific diagnosis and categorization of OSAHS severity require objective measurement of breathing during the period of sleep. The gold standard for diagnosis of OSAHS is an overnight polysomnogram (PSG). A negative in-laboratory PSG rules out OSAHS except in unusual circumstances—e.g., with insufficient REM sleep or supine sleep. Home sleep tests that record only a few respiratory and cardiac channels commonly are used as a cost-effective means for diagnosing patients without significant comorbidity who have a high pretest probability of OSAHS. However, a home study may yield a false-negative result if sleep time is not accurately estimated, and further evaluation may therefore be required.

The key physiological information collected during a sleep study for OSAHS assessment includes measurement of breathing (changes in airflow, respiratory excursion), oxygenation (hemoglobin oxygen saturation), body position, and cardiac rhythm. In addition, PSGs and some home sleep studies measure sleep continuity and sleep stages (by electroencephalography, chin electromyography, and electro-oculography), limb movements (by leg sensors), and snoring intensity. This information is used to quantify the frequency and subtypes of abnormal respiratory events during sleep as well as associated changes in oxygen saturation, arousals, and sleep stage distributions. **Tables 319-1** and **319-2** define the respiratory events scored and the severity guidelines employed during a sleep study. **Figure 319-2** shows examples of sleep-related respiratory events. A typical sleep study report provides quantitative data such as the AHI and the profile of oxygen saturation over the night (mean, nadir, time at low levels). Reports may also include the respiratory disturbance index, which includes the number of respiratory effort-related arousals in addition to the number of apneas plus hypopneas. In-laboratory PSG also quantifies sleep latency (time from “lights off” to first sleep onset), sleep efficiency (percentage of

TABLE 319-1 RESPIRATORY EVENT DEFINITIONS

- **Apnea:** Cessation of airflow for ≥ 10 sec during sleep, accompanied by:
 - Persistent respiratory effort (obstructive apneas, Fig. 319-2A), or
 - Absence of respiratory effort (central apneas, Fig. 319-2B)
- **Hypopnea:** A $\geq 30\%$ reduction in airflow for at least 10 sec during sleep that is accompanied by either a $\geq 3\%$ desaturation or an arousal (Fig. 319-2C)
- **Respiratory effort–related arousal (RERA):** A partially obstructed breath that does not meet the criteria for hypopnea but provides evidence of increasing inspiratory effort (usually through pleural pressure monitoring) punctuated by an arousal (Fig. 319-2D)
- **Flow-limited breath:** A partially obstructed breath, typically within a hypopnea or RERA, identified by a flattened or “scooped-out” inspiratory flow shape (Fig. 319-3)

TABLE 319-2 OBSTRUCTIVE SLEEP APNEA/HYPOPNEA SYNDROME (OSAHS): QUANTIFICATION AND SEVERITY SCALE

- **Apnea-hypopnea index (AHI):**^a Number of apneas plus hypopneas per hour of sleep
- **Respiratory disturbance index (RDI):** Number of apneas plus hypopneas plus RERAs per hour of sleep
- **Mild OSAHS:** AHI of 5–14 events/h
- **Moderate OSAHS:** AHI of 15–29 events/h
- **Severe OSAHS:** AHI of ≥ 30 events/h

^aEach level of AHI can be further quantified by level of sleepiness and associated hypoxemia.

time asleep relative to time in bed), arousal index (number of cortical arousals per hour of sleep), time in each sleep stage, and periodic limb movement index. OSAHS severity can be further characterized according to the degree of sleep fragmentation associated with respiratory disturbances. Relevant metrics include the frequency of cortical micro-arousals or awakenings per sleep hour, reduction in sleep continuity (low sleep efficiency), reduction of time in deeper stages of sleep (stage N3 and REM sleep) and increases in light sleep (stage N1). The detection of autonomic arousals, such as surges in blood pressure, changes in heart rate, and abnormalities in cardiac rhythm, also provides relevant information on OSAHS severity.

Other Laboratory Findings Various imaging studies, including cephalometric radiography, MRI, CT, and fiberoptic endoscopy, can be used to identify anatomic risk factors for OSAHS. Cardiac testing may yield evidence of impaired systolic or diastolic ventricular function or abnormal cardiac structure. Overnight blood pressure monitoring often displays a “non-dipping” pattern (absence of the typical 10-mmHg fall during sleep from blood pressure while awake). Arterial blood gas measurements made during wakefulness are usually normal. Waking hypoxemia or hypercarbia suggests coexisting lung disease or hypoventilation syndrome. Patients with severe nocturnal hypoxemia may have elevated hemoglobin values. A multiple sleep latency test or a maintenance of wakefulness test can be useful in quantifying sleepiness and helping to distinguish OSAHS from narcolepsy.

Health Consequences and Comorbidities OSAHS is a major contributor to cardiac, cerebrovascular, and metabolic disorders as well as to premature death. It is the most common medical cause of daytime sleepiness and negatively influences quality of life. This broad range of health effects is attributable to the impact of sleep fragmentation, cortical arousal, and intermittent hypoxemia on vascular, cardiac, metabolic, and neurologic functions. OSAHS-related respiratory events stimulate sympathetic overactivity, leading to acute blood pressure surges during sleep, endothelial damage, and nocturnal as well as daytime hypertension. OSAHS-related hypoxemia also stimulates release of acute-phase proteins and reactive oxygen species that exacerbate insulin resistance and lipolysis and cause an augmented prothrombotic and proinflammatory state. Inspiratory effort against an occluded airway causes large intrathoracic negative pressure swings, altering cardiac preload and afterload and resulting in cardiac remodeling and reduced cardiac function. Hypoxemia and sympathetic-parasympathetic imbalance also may cause electrical remodeling of the heart and myocyte injury.

HYPERTENSION OSAHS can raise blood pressure to prehypertensive and hypertensive ranges, increase the prevalence of a non-dipping overnight blood pressure pattern, and increase the risk of resistant hypertension. Elevations in blood pressure are due to augmented sympathetic nervous system activation as well as alterations in the rennin-angiotensin-aldosterone system and fluid balance. Treatment of OSAHS with nocturnal continuous positive airway pressure (CPAP) has been shown to reduce 24-h ambulatory blood pressure. Although the overall impact of CPAP on blood pressure levels is relatively modest (averaging 2–4 mmHg), larger improvements are observed among patients with high AHIs and sleepiness.

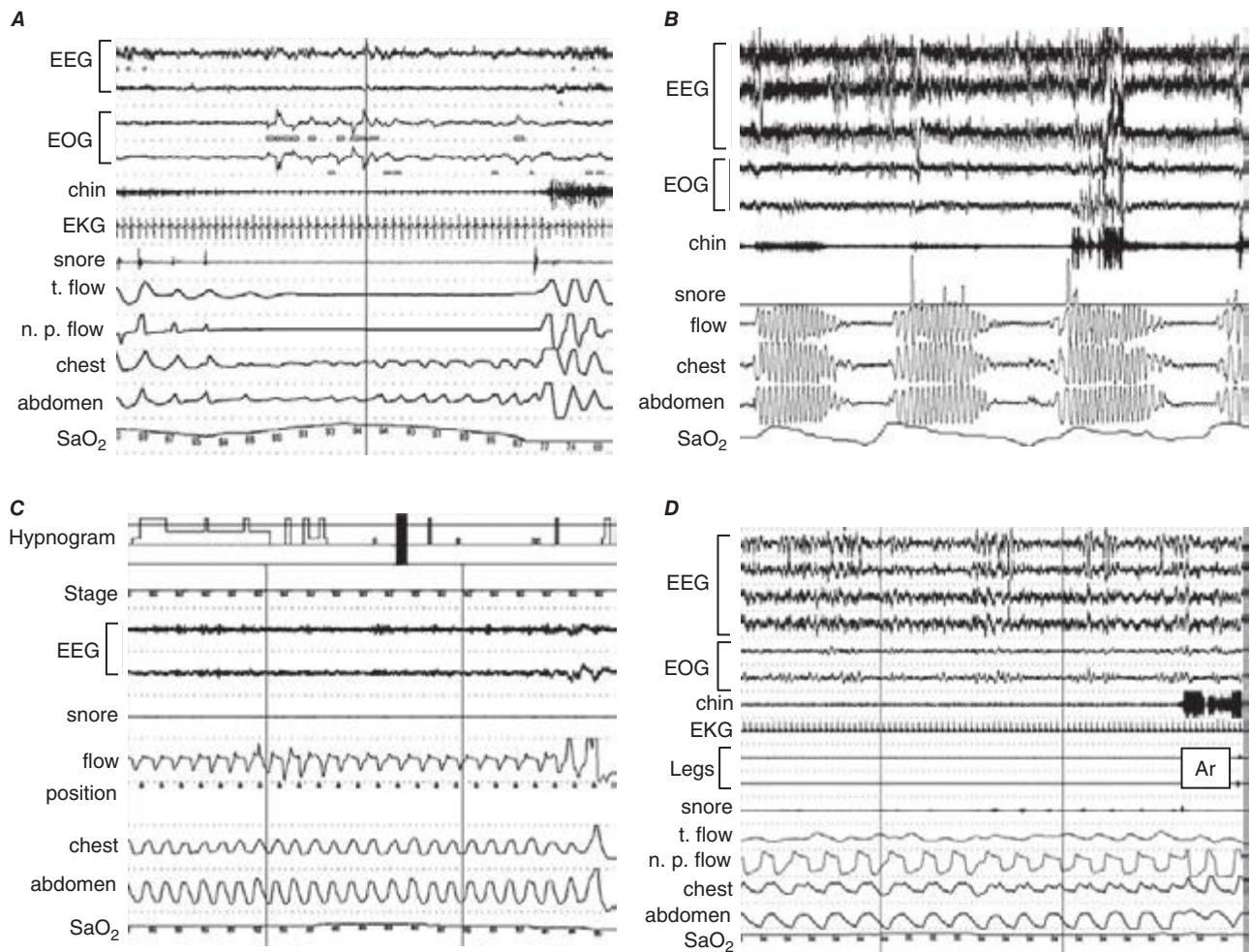


FIGURE 319-2 **A.** Obstructive apnea. There are 30 sec of no airflow, as shown in the nasal pressure (n. p. flow) and thermistor-measured flow (t. flow). Note the presence of chest-abdomen motion, indicating respiratory effort against an occluded airway. **B.** Central apnea in a patient with Cheyne-Stokes respiration due to congestive heart failure. The flat chest-abdomen tracings indicate the absence of inspiratory effort during the central apneas. **C.** Hypopnea. Partial obstruction of the pharyngeal airway can limit ventilation, leading to desaturation (a mild decrease in this patient, from 93% to 90%) and arousal. **D.** Respiratory effort–related arousal (RERA). Minimal flow reduction terminated by an arousal (Ar) without desaturation constitutes a RERA. EEG, electroencephalogram; EOG, electro-oculogram; EKG, electrocardiogram.

CARDIOVASCULAR, CEREBROVASCULAR, AND METABOLIC DISEASES Among the most serious health consequences of OSAHS is its impact on cardiac and metabolic functions. Strong epidemiologic evidence indicates that OSAHS significantly increases the risk of coronary artery disease, heart failure with and without reduced ejection fraction, atrial and ventricular arrhythmias, atherosclerosis and coronary artery disease, stroke, and diabetes. Treatment of OSAHS has been shown to reduce several markers of cardiovascular risk, improve insulin resistance, decrease the recurrence rate of atrial fibrillation, and improve various outcomes in patients with active cardiovascular disease. Large-scale trials are under way to evaluate the role of OSAHS treatment in reducing cardiac event rates and in prolonging the survival of patients with cardiac disease.

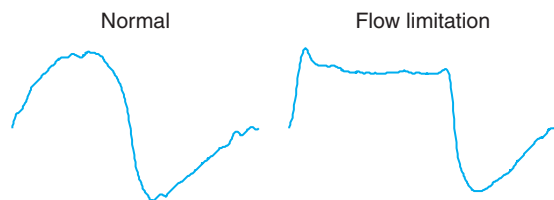


FIGURE 319-3 **Example of flow limitation.** The inspiratory flow pattern in a patent airway is rounded and peaks in the middle. In contrast, a partially obstructed airway exhibits an early peak followed by mid-inspiratory flattening, yielding a scooped-out appearance.

SLEEPINESS More than 50% of patients with moderate to severe OSAHS report daytime sleepiness. Patients with OSAHS symptoms have a twofold increased risk of occupational accidents. Individuals with elevated AHIs are involved in motor vehicle crashes as much as seven times more often than persons with normal AHIs. Randomized controlled trials have shown that treatment of OSAHS with nasal CPAP therapy alleviates sleepiness as measured by either questionnaire or objective testing. However, the degree of improvement varies widely. Residual sleepiness may be due to several factors, including suboptimal treatment adherence, insufficient sleep time, other sleep disorders, or prior hypoxic-mediated damage in brain areas involved in alertness. Visceral adipose tissue, whose amounts are increased in patients with OSAHS, releases somnogenic cytokines that may contribute to sleepiness. Thus, even after treatment, it is important to assess and monitor patients for residual sleepiness and to evaluate the necessity of optimizing treatment adherence, improving sleep patterns, and identifying other disorders contributing to sleepiness.

QUALITY OF LIFE AND MOOD Reductions in health-related quality of life are common in patients with OSAHS, with the largest decrements on the physical and vitality subscales. Treatment with CPAP often results in improvement in these patient-reported outcomes. Depression, in particular symptoms of somatic depression (irritability, fatigue, lack of energy) is commonly reported in OSAHS.

TREATMENT OBSTRUCTIVE SLEEP APNEA/HYPOPNEA SYNDROME -OSAHS-

A comprehensive approach to the management of OSAHS is needed to reduce risk factors and comorbidities. The clinician should seek to identify and address lifestyle and behavioral factors as well as comorbidities that may be exacerbating OSAHS. As appropriate, treatment should aim to reduce weight; optimize sleep duration (7–9 hours); regulate sleep schedules (with similar bedtimes and wake times across the week); encourage the patient to avoid sleeping in the supine position; treat nasal allergies; increase physical activity; eliminate alcohol ingestion within 3 h of bedtime; and minimize use of sedating medications. Patients should be counseled to avoid drowsy driving.

CPAP is the standard medical therapy with the highest level of evidence for efficacy. Delivered through a nasal or nasal-oral mask, CPAP works as a mechanical splint to hold the airway open, thus maintaining airway patency during sleep. An overnight CPAP titration study, performed either in a laboratory or with a home “autotitrating” device, is required to determine the optimal pressure setting that reduces the number of apneas/hypopneas during sleep, improves gas exchange, and reduces arousals. Rates of adherence to CPAP treatment are highly variable (average, 50–80%) and may be improved with support by a skilled health care team who can address side effects (Table 319-3). Despite the limitations of CPAP, controlled studies have demonstrated its beneficial effect on blood pressure, alertness, mood, and insulin sensitivity. Uncontrolled studies also indicate a favorable effect on cardiovascular outcomes, cardiac ejection fraction, atrial fibrillation recurrence, and mortality risk.

Oral appliances for OSAHS work by advancing the mandible, thus opening the airway by repositioning the lower jaw and pulling the tongue forward. These devices generally work better when customized for patient use; maximal adaptation can take several weeks. Efficacy studies show that these devices can reduce the AHI by 50% in two-thirds of individuals, although these data are based largely on patients with mild OSAHS. Side effects of oral appliances include temporomandibular joint pain and tooth movement. Oral appliances are most often used for treating patients with mild OSAHS or patients who do not tolerate CPAP. However, since adherence to the use of oral appliances sometimes exceeds CPAP adherence, these devices are under investigation for treatment of more severe disease.

Upper airway surgery for OSAHS is less effective than CPAP and is mostly reserved for the treatment of patients who snore, have mild OSAHS, and cannot tolerate CPAP. Uvulopalatopharyngoplasty (removal of the uvula and the margin of the soft palate) is the most common surgery and, although results vary greatly, has a success

rate similar to or slightly lower than treatment with oral appliances. Upper airway surgery is less effective in severe OSAHS and in obese patients. Success rates may be higher for multilevel surgery (involving more than one site/structure) performed by an experienced surgeon, but the selection of patients is an important factor and relies on careful targeting of culprit areas for surgical resection. Bariatric surgery is an option for obese patients with OSAHS and can improve not only OSAHS but also other obesity-associated health conditions. Other procedures that can decrease snoring but have minimal effects on OSAHS include injection of the soft palate (resulting in stiffening), radiofrequency ablation, laser-assisted uvulopalatoplasty, and palatal implants.

Supplemental oxygen can improve oxygen saturation, but there is little evidence that it improves OSAHS symptoms or the AHI.

CENTRAL SLEEP APNEA

CSA, which is less common than OSAHS, may occur in isolation or, more often, in combination with obstructive events in the form of “mixed” apneas. CSA is often caused by an increased sensitivity to pCO₂, which leads to an unstable breathing pattern that manifests as hyperventilation alternating with apnea. A prolonged circulation delay between the pulmonary capillaries and carotid chemoreceptors is also a contributing cause; thus individuals with congestive heart failure are at risk for CSA. With prolonged circulation delay, there is a crescendo-decrescendo breathing pattern known as *Cheyne-Stokes respiration* (Fig. 319-2B). Other risk factors for CSA include opioid medications (which appear to have a dose-dependent effect on CSA) and hypoxia (e.g., breathing at high altitude). In some individuals, CPAP—particularly at high pressures—seems to induce central apnea; this condition is referred to as *complex sleep apnea*. Rarely, CSA may be caused by blunted chemosensitivity due to congenital disorders (congenital central hypoventilation syndrome) or acquired factors. Treatment of CSA is difficult and depends on the underlying cause. Limited data suggest that supplemental oxygen can reduce the frequency of central apneas, particularly in patients with hypoxemia. Cheyne-Stokes respiration is treated by optimizing therapy for heart failure and, in some cases, using CPAP with or without supplemental oxygen. *Adaptive servoventilation*, a form of ventilatory support that dynamically changes inspiratory support levels across periods of apnea and hypopnea, can minimize large fluctuations in PCO₂ that produce central apnea and can be effective for the treatment of CSA.

TABLE 319-3 SIDE EFFECTS OF CONTINUOUS POSITIVE AIRWAY PRESSURE CPAP AND THEIR TREATMENTS

Side Effect	Treatment
Nasal congestion	Provide heated humidification, administer saline/steroid nasal sprays
Claustrophobia	Change mask interface (e.g., to nasal prongs), promote habituation (i.e., practice breathing on CPAP while awake)
Difficulty exhaling	Temporarily reduce pressure, provide bilevel positive airway pressure
Bruised nasal ridge	Change mask interface, provide protective padding
Aerophagia	Administer antacids

320e Lung Transplantation

Elbert P. Trulock

Lung transplantation is a therapeutic consideration for many patients with nonmalignant end-stage lung disease, and it prolongs survival and improves quality of life in appropriately selected recipients. Since 1985 almost 40,000 procedures have been recorded worldwide, and since 2009 more than 3000 transplants have been reported annually.

INDICATIONS

The indications span the gamut of lung diseases, but in some respects the distribution of indications differs among countries. According to aggregate international data, the most common indications in the last few years have been chronic obstructive pulmonary disease (COPD), ~29%; idiopathic pulmonary fibrosis (IPF), ~28%; cystic fibrosis (CF), ~16%; α_1 -antitrypsin deficiency emphysema, ~3.5%; and idiopathic pulmonary arterial hypertension (IPAH), ~3%. Other diseases have made up the balance of primary indications, and retransplantation has accounted for ~3% of procedures.

RECIPIENT SELECTION

Transplantation should be considered when other therapeutic options have been exhausted and when the patient's prognosis is expected to improve as a result of the procedure. Survival rates after transplantation can be compared with predictive indices for the patient's disease, but each patient's clinical course must be integrated into the assessment as well. Moreover, quality of life is a primary motive for transplantation for many patients, and the prospect of improved quality-adjusted survival is often attractive even if the survival advantage itself may be marginal.

Disease-specific consensus guidelines for referring patients for evaluation and for proceeding with transplantation are summarized in [Table 320e-1](#) and are linked to clinical, physiologic, radiographic, and pathologic features that influence the prognosis of the respective diseases. Candidates for lung transplantation are also thoroughly screened for comorbidities that might affect the outcome adversely. Conditions such as systemic hypertension, diabetes mellitus, gastroesophageal reflux, and osteoporosis are not unusual; however, if uncomplicated and adequately managed, they do not disqualify patients from transplantation. The upper age limit is ~70 years at most centers, but the median age of recipients has been increasing steadily over the last decade. In the United States in 2009, 22% of recipients were ≥ 65 years old.

Standard exclusions include HIV infection, chronic active hepatitis B or C infection, uncontrolled or untreatable pulmonary or extrapulmonary infection, uncured malignancy, active cigarette smoking, drug or alcohol dependency, irreversible physical deconditioning, chronic nonadherence with medical care, significant disease of another vital organ (e.g., heart, liver, or kidney), and psychiatric or psychosocial situations that could substantially interfere with post-transplantation management. Other problems that may compromise outcome constitute relative contraindications. Some typical issues are ventilator-dependent respiratory failure, previous thoracic surgical procedures, obesity, and coronary artery disease. Chronic infection with antibiotic-resistant *Pseudomonas* species, *Burkholderia* species, *Aspergillus* species, or nontuberculous mycobacteria is a unique concern in some patients with CF. The potential impact of these and other factors must be judged in the clinical context to determine an individual candidate's suitability for transplantation.

WAITING LIST AND ORGAN ALLOCATION

Organ allocation policies are influenced by medical, ethical, geographic, and political factors, with systems varying from country to country. Regardless of the system, potential recipients are placed on a waiting list and must be matched for blood group compatibility and, with some latitude, for lung size with an acceptable donor. Most lungs are procured from deceased donors after total brain failure ("brain

TABLE 320e-1 DISEASE-SPECIFIC GUIDELINES FOR REFERRAL AND TRANSPLANTATION

Chronic Obstructive Pulmonary Disease

Referral

BODE index >5

Transplantation

BODE index 7–10

or

Any of the following criteria:

Hospitalization for exacerbation, with $Pa_{CO_2} >50$ mmHg

Pulmonary hypertension or cor pulmonale, despite oxygen therapy

$FEV_1 <20\%$ with either $DL_{CO} <20\%$ or diffuse emphysema

Cystic Fibrosis / Bronchiectasis

Referral

$FEV_1 <30\%$ or rapidly declining FEV_1

Hospitalization in ICU for exacerbation

Increasing frequency of exacerbations

Refractory or recurrent pneumothorax

Recurrent hemoptysis not controlled by bronchial artery embolization

Transplantation

Oxygen-dependent respiratory failure

Hypercapnia

Pulmonary hypertension

Idiopathic Pulmonary Fibrosis

Referral

Pathologic or radiographic evidence of UIP, regardless of vital capacity

Transplantation

Pathologic or radiographic evidence of UIP

and

Any of the following criteria:

$DL_{CO} <39\%$

Decrement in FVC $\geq 10\%$ during 6 months of follow-up

Decrease in Sp_{O_2} to $<88\%$ during a 6-min walk test

Honeycombing on HRCT (fibrosis score >2)

Idiopathic Pulmonary Arterial Hypertension

Referral

NYHA functional class III or IV, regardless of therapy

Rapidly progressive disease

Transplantation

Failure of therapy with IV epoprostenol (or equivalent drug)

Persistent NYHA functional class III or IV during maximal medical therapy

Low (<350 m) or declining 6-min walk test

Cardiac index <2 L/min per m^2

Right atrial pressure >15 mmHg

Abbreviations: BODE, body mass index (B), airflow obstruction (O), dyspnea (D), exercise capacity (E); DL_{CO} , diffusing capacity for carbon monoxide; FEV_1 , forced expiratory volume in 1 s; FVC, forced vital capacity; HRCT, high-resolution computed tomography; ICU, intensive care unit; NYHA, New York Heart Association; Pa_{CO_2} , partial pressure of carbon dioxide in arterial blood; Sp_{O_2} , arterial oxygen saturation by pulse oximetry; UIP, usual interstitial pneumonitis.

Source: Summarized from JB Orens et al: J Heart Lung Transplant 25:745, 2006. For BODE index, BR Celli et al: N Engl J Med 350:1005, 2004.

death"), but only ~15–20% of brain-death organ donors yield either one or two lungs suitable for transplantation. Lungs from donors after cardiac death have been utilized to a limited extent (~2% of lung donors in the United States in 2009). Recently, ex vivo lung perfusion has been used by some centers to assess donor lungs that are marginal or high-risk for implantation by standard criteria; if the results of ex vivo testing are satisfactory, these lungs have been transplanted successfully.

In the United States, a lung allocation scoring system is used to prioritize patients on the waiting list. The lung allocation score (LAS) for a patient is based on the patient's risk of death during 1 year on

320e-2 the waiting list and the patient's likelihood of survival for 1 year after transplantation. The LAS can range from 0 to 100, and precedence for transplantation is ranked from highest to lowest scores. Both the lung disease and its severity affect a patient's LAS; parameters in the LAS must be updated biannually but can be submitted for recalculation whenever the patient's condition changes. The median LAS for all candidates on the waiting list is usually ~35, but the LAS tends to be higher among patients with IPF and CF than among patients with COPD and IPAH.

Under this system in the United States, the median waiting time for transplantation has been ~135 days. The overall death rate on the waiting list has been ~6.5%, but death rates vary substantially with the diagnosis (e.g., COPD, ~3%; IPF, ~7%) and with the LAS (e.g., 40–49, ~7%; 50–59, ~15%; ≥60, ~25%). The indications for transplantation depend not only on the prevalence and natural history of the various lung diseases but also on the LAS typically associated with these diseases. While patients with IPF constitute ~20% of the waiting list, they make up ~34% of recipients because their allocation scores are typically higher than those of patients with other diseases.

TRANSPLANT PROCEDURE

Bilateral transplantation is mandatory for CF and other forms of bronchiectasis because the risk of spillover infection from a remaining native lung precludes single-lung transplantation. Heart-lung transplantation is obligatory for Eisenmenger syndrome with complex anomalies that cannot readily be repaired in conjunction with lung transplantation and for concomitant end-stage lung and heart disease. However, cardiac replacement is not necessary for cor pulmonale because right ventricular function will recover when pulmonary vascular afterload is normalized by lung transplantation.

Either bilateral or single-lung transplantation is an option for other diseases unless there is a special consideration, but bilateral transplantation has been utilized increasingly for most indications. Recently, ~65% of procedures in the United States have been bilateral, and ~70% of transplants for COPD, ~55% of those for IPF, and ~95% of those for IPAH in the international registry have been bilateral.

Living-donor lobar transplantation has had a limited role in adult lung transplantation but is now rarely performed. It has been used predominantly for teenagers or young adults with CF and has usually been reserved for patients who were unlikely to survive the wait for a deceased-donor organ.

POSTTRANSPLANTATION MANAGEMENT

Induction therapy with an antilymphocyte globulin or an interleukin 2 receptor antagonist is utilized by ~55% of centers, and a three-drug maintenance immunosuppressive regimen that includes a calcineurin

inhibitor (cyclosporine or tacrolimus), a purine synthesis antagonist (azathioprine or a mycophenolic acid precursor), and prednisone is traditional. Subsequently, other drugs (e.g., sirolimus) may be substituted into the regimen for various reasons. Prophylaxis against *Pneumocystis jirovecii* pneumonia is standard, and prophylaxis against cytomegalovirus (CMV) infection and fungal infection is part of many protocols. The dose of cyclosporine, tacrolimus, or sirolimus is adjusted by blood-level monitoring. All of these agents are metabolized by the hepatic cytochrome P450 system, and interactions with medications that affect this pathway can significantly alter their clearance and blood level.

Routine management focuses on monitoring of the allograft, regulation of immunosuppressive therapy, and expeditious detection of problems or complications. Regular contact with a nurse coordinator, physician follow-up, chest radiography, blood tests, and spirometry are customary, and periodic surveillance bronchoscopies are employed in some programs. If recovery is uncomplicated, lung function rapidly improves and then stabilizes by 3–6 months after transplantation. Subsequently, the variation in spirometric measurements is small, and a sustained decline of ≥10–15% signals a potentially significant problem.

OUTCOMES

Survival Major registries publish survival rates (Table 320e-2) and other outcomes annually (www.ishlt.org; www.srtr.org). In the international registry, the survival half-life for recipients with IPF is 4.4 years; IPAH, 5 years; COPD, 5.3 years; and CF, 7.5 years. However, age and transplantation procedure have a significant impact on outcome. For recipients 18–59 years of age, the survival half-life is 5–6 years, but this figure decreases to 4.4 years among patients 60–65 years old and to 3.6 years for those >65 years old. Survival rates at >15 years have been significantly higher after bilateral transplantation than after unilateral transplantation for COPD, α₁-antitrypsin deficiency emphysema, IPF, and IPAH.

The main sources of perioperative mortality include technical complications of the operation, primary graft dysfunction, and infections. Acute rejection and CMV infection are common problems in the first year, but neither is usually fatal. Beyond the first year, chronic rejection and non-CMV infections cause the majority of deaths.

Risk factors for mortality have been analyzed in the international and U. S. registries. In these analyses, factors associated with an increased risk of death, especially in the first year after transplantation, have included the following: recipients hospitalized at the time of transplantation; recipients supported by mechanical ventilation, extracorporeal membrane oxygenation, inotropic drugs, or dialysis at the time of transplantation; and recipients undergoing retransplantation.

TABLE 320e-2 RECIPIENT SURVIVAL, BY PRETRANSPLANTATION DIAGNOSIS (1990–2010)

Diagnosis, Transplant Type	n	Survival Rate, %					
		3 Months	1 Year	3 Years	5 Years	10 Years	15 Years
Chronic obstructive pulmonary disease							
Bilateral	5147	92	83	68	57	34	20
Single	6797	90	81	63	49	21	7
α ₁ -antitrypsin deficiency emphysema							
Bilateral	1403	88	80	68	59	39	22
Single	1086	87	78	62	52	29	14
Cystic fibrosis	5608	90	82	69	59	42	32
Idiopathic pulmonary fibrosis							
Bilateral	3057	84	75	62	52	33	19
Single	4481	86	74	57	43	20	8
Idiopathic pulmonary arterial hypertension							
Bilateral	1037	78	72	61	53	38	24
Single	271	71	63	51	41	25	18
Sarcoidosis	849	84	73	58	51	30	

Source: Data from www.ishlt.org/registries/slides.asp?slides=heartLungRegistry.

However, other factors have contributed as well. The mortality risk has been higher at centers with <20–30 transplantations per year.

Function Regardless of the disease, successful transplantation impressively restores cardiopulmonary function. After bilateral transplantation, pulmonary function tests are typically normal; after unilateral transplantation, a mild abnormality characteristic of the remaining diseased lung is still apparent. Formal exercise testing usually demonstrates some impairment in maximal work rate and maximal oxygen uptake, but few recipients report any limitation to activities of daily living.

Quality of Life Both overall and health-related quality-of-life scores are enhanced. With multidimensional profiles, improvements extend across most domains and are sustained longitudinally unless chronic rejection or some other complication develops. Other problems that detract from quality of life include renal dysfunction and drug side effects.

Cost The cost of transplantation depends on the health care system, other health care policies, and economic factors that vary from country to country. In the United States in 2011, the average billed charge for the period from 30 days before bilateral lung transplantation until 180 days after discharge from the transplantation admission was \$797,300. The total cost included the following charges: all care during 30 days before transplantation, \$21,400; organ procurement, \$90,300; hospital transplantation admission, \$458,500; physician fees during transplantation admission, \$56,300; all inpatient and outpatient care for 180 days after discharge, \$142,600; and all outpatient drugs (including immunosuppressants) for 180 days after discharge, \$28,200.

COMPLICATIONS

Lung transplantation can be complicated by a variety of problems (Table 320e-3). Aside from predicaments that are unique to transplantation,

TABLE 320e-3 MAJOR POTENTIAL COMPLICATIONS OF LUNG TRANSPLANTATION AND IMMUNOSUPPRESSION

Category	Complication
Allograft	Primary graft dysfunction; anastomotic dehiscence or stenosis; ischemic airway injury with bronchostenosis or bronchomalacia; rejection; infection; recurrence of primary disease (sarcoidosis, lymphangioleiomyomatosis, giant cell interstitial pneumonitis, diffuse panbronchiolitis, pulmonary alveolar proteinosis, Langerhans cell histiocytosis)
Thoracic	Phrenic nerve injury/diaphragmatic dysfunction; recurrent laryngeal nerve injury/vocal cord dysfunction; cervical ganglia injury/Horner's syndrome; pneumothorax; pleural effusion; chylothorax; empyema
Cardiovascular	Intraoperative or perioperative air embolism; postoperative pericarditis; perioperative myocardial injury/infarction; venous thromboembolism; supraventricular dysrhythmias; systemic hypertension
Gastrointestinal	Esophagitis (especially <i>Candida</i> , herpesvirus, or cytomegalovirus [CMV]); gastroparesis; gastroesophageal reflux; diarrhea (<i>Clostridium difficile</i> ; medications, especially mycophenolate mofetil and sirolimus); colitis (<i>C. difficile</i> ; CMV)
Hepatobiliary	Hepatitis (especially CMV or medications); acalculous cholecystitis
Renal	Calcineurin inhibitor nephropathy; hemolytic-uremic syndrome (thrombotic microangiopathy)
Neurologic	Perioperative stroke; tremors; seizures; reversible posterior leukoencephalopathy; headaches
Musculoskeletal	Steroid myopathy; rhabdomyolysis (cyclosporine + HMG-coA reductase inhibitor treatment); osteoporosis; avascular necrosis
Metabolic	Obesity; diabetes mellitus; hyperlipidemia; idiopathic hyperammonemia
Hematologic	Anemia; leukopenia; thrombocytopenia; thrombotic microangiopathy
Oncologic	Lymphoproliferative disease and lymphoma; skin cancers; other malignancies

side effects and toxicities of immunosuppressive medications can cause new medical problems or aggravate preexisting conditions.

Graft Dysfunction Primary graft dysfunction (PGD), an acute lung injury, is a manifestation of multiple potential insults to the donor organ that are inherent in the transplantation process. The principal clinical features are diffuse pulmonary infiltrates and hypoxemia within 72 h of transplantation; however, the presentation can be mimicked by pulmonary venous obstruction, hyperacute rejection, pulmonary edema, and pneumonia.

The severity is variable, and a standardized grading system has been established. Up to 50% of lung transplant recipients may have some degree of PGD, and ~10–20% have severe PGD. The treatment follows the conventional supportive paradigm for acute lung injury. Inhalation of nitric oxide and extracorporeal membrane oxygenation have been used in severe cases; retransplantation has also been performed, but when undertaken in the first 30 days this procedure is associated with a poor survival rate (~30% at 1 year). Most recipients with mild PGD recover, but the mortality rate for severe PGD has been ~40–60%. PGD is also associated with longer postoperative ventilator support, longer intensive care unit and hospital stays, higher costs, and excess morbidity, and severe PGD is a risk factor for the later development of chronic rejection (see below).

Airway Complications The bronchial blood supply to the donor lung is disrupted during procurement. Bronchial revascularization during transplantation is technically feasible in some cases, but it is not widely practiced. Consequently, after implantation, the donor bronchus is dependent on retrograde bronchial blood flow from the pulmonary circulation and is vulnerable to ischemia.

The spectrum of airway problems includes anastomotic necrosis and dehiscence, occlusive granulation tissue, anastomotic or bronchial stenosis, and bronchomalacia. The incidence has been in the range of 7–18%, but the associated mortality rate has been low. These problems usually can be managed bronchoscopically with techniques such as simple endoscopic debridement, laser photoresection, balloon dilation, and bronchial stenting.

Rejection Rejection is the main deterrent to higher medium- and long-term survival rates. In this immunologic response to alloantigen recognition, both cell-mediated and antibody-mediated (humoral) cascades can play a role. Cellular rejection is effected by T lymphocyte interactions with donor alloantigens, mainly in the major histocompatibility complex (MHC), whereas humoral rejection is driven by antibodies to donor MHC alloantigens or possibly to non-MHC antigens on epithelial or endothelial cells.

Rejection is often categorized as acute or chronic without reference to the underlying mechanism. Acute rejection is cell-mediated, and its incidence is highest in the first 6–12 months after transplantation. In contrast, chronic rejection generally emerges later, and both alloimmune and non-alloimmune fibroproliferative reactions may contribute to its pathogenesis.

Acute Cellular Rejection With current immunosuppressive regimens, ~30–40% of recipients experience acute rejection in the first year. Acute cellular rejection (ACR) can be clinically silent or can be manifested by nonspecific symptoms or signs that may include cough, low-grade fever, dyspnea, hypoxemia, inspiratory crackles, interstitial infiltrates, and declining lung function; however, clinical impressions are not reliable. The diagnosis is confirmed by transbronchial biopsies showing the characteristic lymphocytic infiltrates around arterioles or bronchioles, and a standardized pathologic scheme is used to grade the biopsies.

Minimal ACR on a surveillance biopsy in a clinically stable recipient is often left untreated, but higher grades generally are treated regardless of the clinical situation. Treatment usually includes a short course of high-dose glucocorticoids and adjustment of the maintenance immunosuppressive regimen. Most episodes respond to this approach; however, more intensive therapy is sometimes necessary for persistent or recurrent episodes.

320e-4 Chronic Rejection This complication is the main impediment to long-term survival and is the source of substantial morbidity because of its impact on lung function and quality of life. Clinically, chronic rejection is characterized physiologically by airflow limitation and pathologically by bronchiolitis obliterans; the process is designated *bronchiolitis obliterans syndrome* (BOS). Transbronchial biopsies are relatively insensitive for detecting bronchiolitis obliterans, and pathologic confirmation is not required for diagnosis. Thus, after other causes of graft dysfunction have been excluded, the diagnosis of BOS is based primarily on a sustained decrement ($\geq 20\%$) in forced expiratory volume in 1 s (FEV_1), although smaller declines in FEV_1 ($\geq 10\%$) or in midexpiratory flow rate ($FEF_{25-75\%}$) may presage BOS. Spirometric criteria for diagnosis and staging of BOS have been standardized.

The prevalence of BOS approaches 50% by 5 years after transplantation. Antecedent ACR is the main risk factor, but PGD, CMV pneumonitis, other community-acquired respiratory viral infections, and gastroesophageal reflux have been implicated as well. BOS can present acutely and imitate infectious bronchitis, or it can manifest as an insidious decline in lung function. The chest radiograph is typically unchanged; CT may reveal mosaic perfusion, air trapping, ground-glass opacities, or bronchiolectasis. Bronchoscopy is indicated to rule out other processes, but transbronchial biopsies identify bronchiolitis obliterans in a minority of cases.

BOS usually is treated with augmented immunosuppression, but there is no consensus about therapy. Strategies include changes in the maintenance drug regimen, including the addition of azithromycin, antilymphocyte globulin, photopheresis, and total lymphoid irradiation. Although therapy may stabilize lung function, the overall results of treatment have been disappointing; the median survival period after onset has been ~3–4 years. Retransplantation is a consideration if clinical circumstances and other comorbidities are not prohibitive, but survival rates have been inferior to those with primary transplantation.

Humoral Rejection Consensus on the role of antibody-mediated rejection is still evolving. Hyperacute rejection is caused by preformed HLA antibodies in the recipient, but it is minimized by pretransplantation antibody screening coupled with virtual or direct cross-matching with any potential donor. Donor-specific HLA antibodies develop after transplantation in up to 50% of recipients, and their presence

has been associated with an increased risk of both ACR and BOS and with poorer overall survival. However, the mechanisms by which these antibodies could contribute to ACR or BOS or could otherwise be detrimental have not been unraveled. Formal criteria for antibody-mediated rejection have been defined for renal transplantation, but few cases in lung transplantation fulfill these criteria. Nonetheless, episodes of acute lung allograft dysfunction occasionally have been attributed directly to antibody-mediated injury. If treatment is indicated, potential therapies include plasmapheresis and administration of IV immune globulin, rituximab, bortezomib, or eculizumab.

Infection The lung allograft is especially susceptible to infection, which has been one of the leading causes of death in recipients. In addition to a blunted immune response from immunosuppressive drugs, other normal defenses are compromised: the cough reflex is diminished, and mucociliary clearance is impaired in the transplanted lung. The spectrum of infections includes both opportunistic and non-opportunistic pathogens.

Bacterial bronchitis or pneumonia can occur at any time but is very common in the perioperative period. Later, bronchitis occurs frequently in recipients with BOS, and *Pseudomonas aeruginosa* or methicillin-resistant *Staphylococcus aureus* is often the culprit.

CMV is the most common cause of viral infection. Although gastroenteritis, colitis, and hepatitis can occur, CMV viremia and CMV pneumonia are the main illnesses. Most episodes occur in the first 6 months, and treatment with ganciclovir is effective unless resistance develops. Other community-acquired viruses, such as influenza, parainfluenza, and respiratory syncytial viruses, also contribute to respiratory complications. The most problematic fungal infections are caused by *Aspergillus* species. The spectrum encompasses simple pulmonary colonization, tracheobronchitis, invasive pulmonary aspergillosis, and disseminated aspergillosis, and the clinical scenario dictates treatment.

Other Complications Other potential complications are listed in Table 320e-3. Many of them are related to side effects or toxicities of immunosuppressive drugs. Management of these general medical problems is guided by standard practices, but the complex milieu of transplantation requires close collaboration and good communication among health care providers.

This page intentionally left blank

SECTION 1 RESPIRATORY CRITICAL CARE

321 Approach to the Patient with Critical Illness

John P. Kress, Jesse B. Hall

The care of critically ill patients requires a thorough understanding of pathophysiology and centers initially on the resuscitation of patients at the extremes of physiologic deterioration. This resuscitation is often fast-paced and occurs early, without a detailed awareness of the patient's chronic medical problems. While physiologic stabilization is taking place, intensivists attempt to gather important background medical information to supplement the real-time assessment of the patient's current physiologic conditions. Numerous tools are available to assist intensivists in the accurate assessment of pathophysiology and management of incipient organ failure, offering a window of opportunity for diagnosing and treating underlying disease(s) in a stabilized patient. Indeed, the use of invasive interventions such as mechanical ventilation and renal replacement therapy is commonplace in the intensive care unit (ICU). An appreciation of the risks and benefits of such aggressive and often invasive interventions is vital to ensure an optimal outcome. Nonetheless, intensivists must recognize when a patient's chances for recovery are remote or nonexistent and must counsel and comfort dying patients and their significant others. Critical care physicians often must redirect the goals of care from resuscitation and cure to comfort when the resolution of an underlying illness is not possible.

ASSESSMENT OF ILLNESS SEVERITY

In the ICU, illnesses are frequently categorized by degree of severity. Numerous severity-of-illness (SOI) scoring systems have been developed and validated over the past three decades. Although these scoring systems have been validated as tools to assess populations of critically ill patients, their utility in predicting individual patient outcomes is not clear. SOI scoring systems are important for defining populations of critically ill patients. Such systematic scoring allows effective comparison of groups of patients enrolled in clinical trials. In verifying a purported benefit of therapy, investigators must be confident that different groups involved in a clinical trial have similar illness severities. SOI scores are also useful in guiding hospital administrative policies, directing the allocation of resources such as nursing and ancillary care and assisting in assessments of quality of ICU care over time. Scoring system validations are based on the premise that age, chronic medical illnesses, and derangements from normal physiology are associated with increased mortality rates. All existing SOI scoring systems are derived from patients who have already been admitted to the ICU.

SOI scoring systems cannot be used to predict survival in individual patients. No established scoring systems that purport to direct clinicians' decision-making regarding criteria for admission to an ICU are available, although such models are being developed. Thus the use of SOI scoring systems to direct therapy and clinical decision-making cannot be recommended at present. Instead, these tools should be used as a source of important data to complement clinical bedside decision-making.

The most commonly utilized scoring systems are the APACHE (Acute Physiology and Chronic Health Evaluation) and the SAPS (Simplified Acute Physiology Score) systems.

THE APACHE II SCORING SYSTEM

The APACHE II system is the most commonly used SOI scoring system in North America. Age, type of ICU admission (after elective

surgery vs. nonsurgical or after emergency surgery), chronic health problems, and 12 physiologic variables (the worst values for each in the first 24 h after ICU admission) are used to derive a score. The predicted hospital mortality rate is derived from a formula that takes into account the APACHE II score, the need for emergency surgery, and a weighted, disease-specific diagnostic category (Table 321-1). The relationship between APACHE II score and mortality risk is illustrated in Fig. 321-1. Updated versions of the APACHE scoring system (APACHE III and APACHE IV) have been published.

THE SAPS SCORING SYSTEM

The SAPS II score, used more frequently in Europe than in the United States, was derived in a manner similar to the APACHE score. This score is not disease specific but rather incorporates three underlying disease variables: AIDS, metastatic cancer, and hematologic malignancy. SAPS 3, which utilizes a 1-h rather than a 24-h window for measuring physiologic derangement scores, was developed in 2005.

SHOCK

See also Chap. 324.

INITIAL EVALUATION

Shock, a common condition necessitating ICU admission or occurring in the course of critical care, is defined by the presence of multisystem end-organ hypoperfusion. Clinical indicators include reduced mean arterial pressure (MAP), tachycardia, tachypnea, cool skin and extremities, acute altered mental status, and oliguria. Hypotension is usually, though not always, present. The end result of multiorgan hypoperfusion is tissue hypoxia, often with accompanying lactic acidosis. Since the MAP is the product of cardiac output and systemic vascular resistance (SVR), reductions in blood pressure can be caused by decreases in cardiac output and/or SVR. Accordingly, once shock is contemplated, the initial evaluation of a hypotensive patient should include an early bedside assessment of the adequacy of cardiac output (Fig. 321-2). Clinical evidence of *diminished* cardiac output includes a narrow pulse pressure—a marker that correlates with stroke volume—and cool extremities with delayed capillary refill. Signs of *increased* cardiac output include a widened pulse pressure (particularly with a reduced diastolic pressure), warm extremities with bounding pulses, and rapid capillary refill. If a hypotensive patient has clinical signs of increased cardiac output, it can be inferred that the reduced blood pressure is from decreased SVR.

In hypotensive patients with signs of reduced cardiac output, an assessment of intravascular volume status is appropriate. A hypotensive patient with decreased intravascular volume status may have a history suggesting hemorrhage or other volume losses (e.g., vomiting, diarrhea, polyuria). Although evidence of a reduced jugular venous pressure (JVP) is often sought, static measures of right atrial pressure do not predict fluid responsiveness reliably; the *change* in right atrial pressure as a function of spontaneous respiration is a better predictor of fluid responsiveness (Fig. 321-3). Patients with fluid-responsive (i.e., hypovolemic) shock also may manifest large changes in pulse pressure as a function of respiration during mechanical ventilation (Fig. 321-4). A hypotensive patient with increased intravascular volume and cardiac dysfunction may have S_3 and/or S_4 gallops on examination, increased JVP, extremity edema, and crackles on lung auscultation. The chest x-ray may show cardiomegaly, widening of the vascular pedicle, Kerley B lines, and pulmonary edema. Chest pain and electrocardiographic changes consistent with ischemia may be noted (Chap. 326).

In hypotensive patients with clinical evidence of increased cardiac output, a search for causes of decreased SVR is appropriate. The

TABLE 321-1 CALCULATION OF ACUTE PHYSIOLOGY AND CHRONIC HEALTH EVALUATION II (APACHE II) SCORE^a

Acute Physiology Score									
Score	4	3	2	1	0	1	2	3	4
Rectal temperature (°C)	≥41	39.0–40.9		38.5–38.9	36.0–38.4	34.0–35.9	32.0–33.9	30.0–31.9	≤29.9
Mean blood pressure (mmHg)	≥160	130–159	110–129		70–109		50–69		≤49
Heart rate (beats/min)	≥180	140–179	110–139		70–109		55–69	40–54	≤39
Respiratory rate (breaths/min)	≥50	35–49		25–34	12–24	10–11	6–9		≤5
Arterial pH	≥7.70	7.60–7.69		7.50–7.59	7.33–7.49		7.25–7.32	7.15–7.24	<7.15
Oxygenation									
If $FI_{O_2} > 0.5$, use $(A - a) D_{O_2}$	≥500	350–499	200–349		<200				
If $FI_{O_2} \leq 0.5$, use Pa_{O_2}					>70	61–70		55–60	<55
Serum sodium (meq/L)	≥180	160–179	155–159	150–154	130–149		120–129	111–119	≤110
Serum potassium (meq/L)	≥7.0	6.0–6.9		5.5–5.9	3.5–5.4	3.0–3.4	2.5–2.9		<2.5
Serum creatinine (mg/dL)	≥3.5	2.0–3.4	1.5–1.9		0.6–1.4		<0.6		
Hematocrit (%)	≥60		50–59.9	46–49.9	30–45.9		20–29.9		<20
WBC count (10^3 /mL)	≥40		20–39.9	15–19.9	3–14.9		1–2.9		<1
Glasgow Coma Score ^{b,c}									
<i>Eye Opening</i>	<i>Verbal (Nonintubated)</i>			<i>Verbal (Intubated)</i>			<i>Motor Activity</i>		
4—Spontaneous	5—Oriented and talks			5—Seems able to talk			6—Verbal command		
3—Verbal stimuli	4—Disoriented and talks			3—Questionable ability to talk			5—Localizes to pain		
2—Painful stimuli	3—Inappropriate words			1—Generally unresponsive			4—Withdraws from pain		
1—No response	2—Incomprehensible sounds						3—Decorticate		
	1—No response						2—Decerebrate		
							1—No response		
Points Assigned to Age and Chronic Disease									
<i>Age, Years</i>	<i>Score</i>								
<45	0								
45–54	2								
55–64	3								
65–74	5								
≥75	6								
<i>Chronic Health (History of Chronic Conditions)^d</i>	<i>Score</i>								
None	0								
If patient is admitted after elective surgery	2								
If patient is admitted after emergency surgery or for reason other than after elective surgery	5								

^aThe APACHE II score is the sum of the acute physiology score (vital signs, oxygenation, laboratory values), the Glasgow coma score, age, and chronic health points. The worst values during the first 24 h in the ICU should be used. ^bGlasgow coma score (GCS) = eye-opening score + verbal (intubated or nonintubated) score + motor score. ^cFor GCS component of acute physiology score, subtract GCS from 15 to obtain points assigned. ^dHepatic: cirrhosis with portal hypertension or encephalopathy; cardiovascular: class IV angina (at rest or with minimal self-care activities); pulmonary: chronic hypoxemia or hypercapnia, polycythemia, ventilator dependence; renal: chronic peritoneal or hemodialysis; immune: immunocompromised host.

Abbreviations: (A – a) D_{O_2} , alveolar-arterial oxygen difference; FI_{O_2} , fraction of inspired oxygen; Pa_{O_2} , partial pressure of oxygen; WBC, white blood cell count.

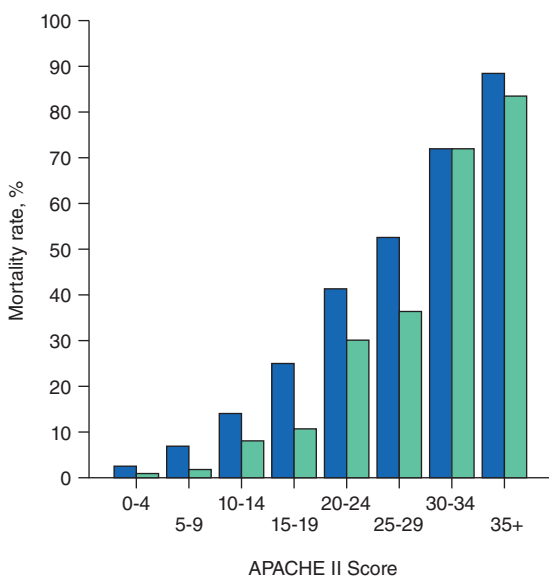


FIGURE 321-1 APACHE II survival curve. Blue, nonoperative; green, postoperative.

most common cause of high-cardiac-output hypotension is sepsis (Chap. 325). Other causes include liver failure, severe pancreatitis, burns and other trauma that elicit the systemic inflammatory response syndrome (SIRS), anaphylaxis, thyrotoxicosis, and peripheral arteriovenous shunts.

In summary, the most common categories of shock are hypovolemic, cardiogenic, and high-cardiac-output with decreased SVR (high-output hypotension). Certainly more than one category can occur simultaneously (e.g., hypovolemic and septic shock).

The initial assessment of a patient in shock should take only a few minutes. It is important that aggressive resuscitation is instituted on the basis of the initial assessment, particularly since early resuscitation from septic and cardiogenic shock may improve survival (see below). If the initial bedside assessment yields equivocal or confounding data, more objective assessments such as echocardiography and/or invasive vascular monitoring may be useful. The goal of early resuscitation is to reestablish adequate tissue perfusion and thus to prevent or minimize end-organ injury.

MECHANICAL VENTILATORY SUPPORT

(See also Chap. 323) During the initial resuscitation of patients in shock, principles of advanced cardiac life support should be followed.

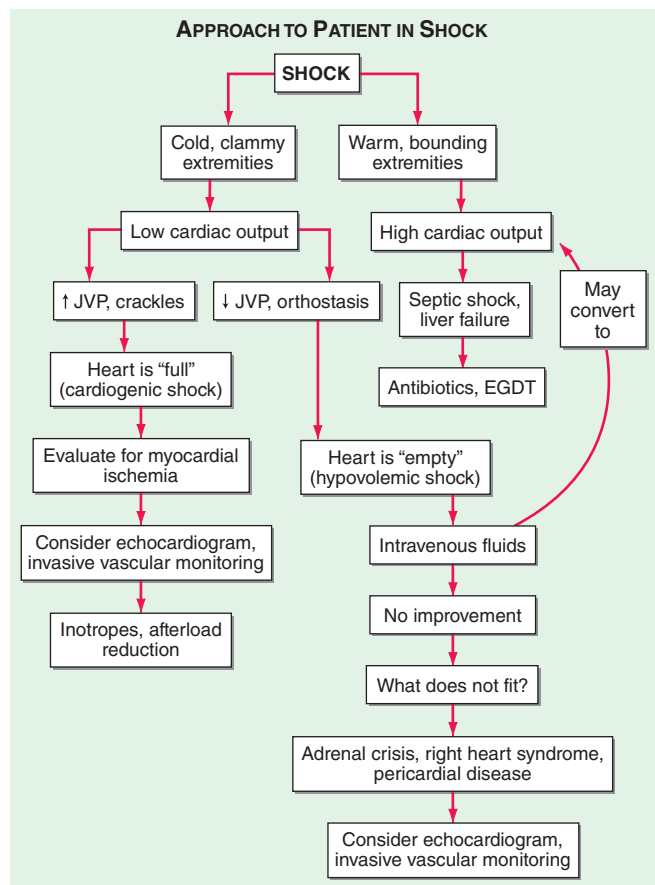


FIGURE 321-2 Approach to the patient in shock. EGDT, early goal-directed therapy; JVP, jugular venous pulse.

As such patients may be obtunded and unable to protect the airway, an early assessment of the airway is mandatory. Early intubation and mechanical ventilation often are required. Reasons for the institution of endotracheal intubation and mechanical ventilation include acute hypoxemic respiratory failure and ventilatory failure, which frequently accompany shock. Acute hypoxemic respiratory failure may occur in patients with cardiogenic shock and pulmonary edema (Chap. 326) as well as in those who are in septic shock with pneumonia or acute respiratory distress syndrome (ARDS) (Chaps. 322 and 325). Ventilatory failure often occurs as a consequence of an increased load on the respiratory system in the form of acute metabolic (often lactic) acidosis or decreased lung compliance due to pulmonary edema. Inadequate perfusion to respiratory muscles in the setting of shock may be another reason for early intubation and mechanical ventilation. Normally, the respiratory muscles receive a very small percentage of the cardiac output. However, in patients who are in shock with respiratory distress, the percentage of cardiac output dedicated to respiratory muscles may increase by tenfold or more. Lactic acid production from inefficient respiratory muscle activity presents an additional ventilatory load.

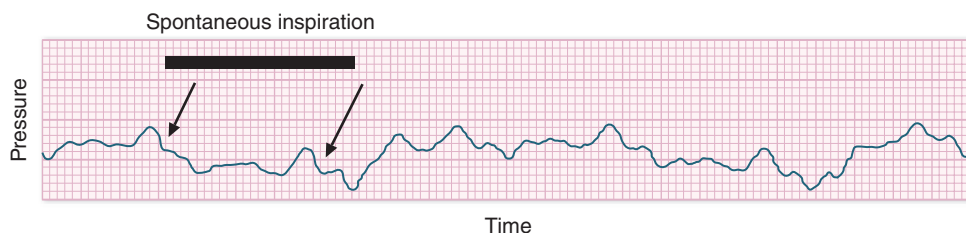


FIGURE 321-3 Right atrial pressure change during spontaneous respiration in a patient with shock whose cardiac output will increase in response to intravenous fluid administration. The right atrial pressure decreases from 7 mmHg to 4 mmHg. The horizontal bar marks the time of spontaneous inspiration.

Mechanical ventilation may relieve the work of breathing and allow redistribution of a limited cardiac output to other vital organs. Patients demonstrate respiratory distress by an inability to speak full sentences, accessory use of respiratory muscles, paradoxical abdominal muscle activity, extreme tachypnea (>40 breaths/min), and decreasing respiratory rate despite an increasing drive to breathe. When patients with shock are treated with mechanical ventilation, a major goal is for the ventilator to assume all or the majority of the work of breathing, facilitating a state of minimal respiratory muscle work. With the institution of mechanical ventilation for shock, further declines in MAP are frequently seen. The reasons include impeded venous return from positive-pressure ventilation, reduced endogenous catecholamine secretion once the stress associated with respiratory failure abates, and the actions of drugs used to facilitate endotracheal intubation (e.g., propofol, opiates). Accordingly, hypotension should be anticipated during endotracheal intubation. Because many of these patients may be fluid responsive, IV volume administration should be considered. Figure 321-2 summarizes the diagnosis and treatment of different types of shock. For further discussion of individual forms of shock, see Chaps. 324, 325, and 326.

RESPIRATORY FAILURE

Respiratory failure is one of the most common reasons for ICU admission. In some ICUs, $\geq 75\%$ of patients require mechanical ventilation during their stay. Respiratory failure can be categorized mechanistically on the basis of pathophysiologic derangements in respiratory function.

TYPE I: ACUTE HYPOXEMIC RESPIRATORY FAILURE

This type of respiratory failure occurs with alveolar flooding and subsequent intrapulmonary shunt physiology. Alveolar flooding may be a consequence of pulmonary edema, pneumonia, or alveolar hemorrhage. Pulmonary edema can be further categorized as occurring due to elevated pulmonary microvascular pressures, as seen in heart failure and intravascular volume overload or ARDS (“low-pressure pulmonary edema,” Chap. 322). This syndrome is defined by acute onset (≤ 1 week) of bilateral opacities on chest imaging that are not fully explained by cardiac failure or fluid overload and of shunt physiology requiring positive end-expiratory pressure (PEEP). Type I respiratory failure occurs in clinical settings such as sepsis, gastric aspiration, pneumonia, near-drowning, multiple blood transfusions, and pancreatitis. The mortality rate among patients with ARDS was traditionally very high (50–70%), although changes in patient care have led to mortality rates closer to 30% (see below).

For many years, physicians have suspected that mechanical ventilation of patients with ARDS may propagate lung injury. Cyclical collapse and reopening of alveoli may be partly responsible for this adverse effect. As seen in Fig. 321-5, the pressure-volume relationship of the lung in ARDS is not linear. Alveoli may collapse at very low lung volumes. Animal studies have suggested that stretching and overdistention of injured alveoli during mechanical ventilation can further injure the lung. Concern over this alveolar overdistention, termed *ventilator-induced “volutrauma,”* led to a multicenter, randomized, prospective trial comparing traditional ventilator strategies for ARDS

(large tidal volume: 12 mL/kg of ideal body weight) with a low tidal volume (6 mL/kg of ideal body weight). This study showed a dramatic reduction in mortality rate in the low-tidal-volume group from that in the high-tidal-volume group (31% versus 39.8%). In addition, a “fluid-conservative” management strategy (maintaining a low central venous pressure [CVP] or pulmonary capillary wedge pressure [PCWP]) is associated with fewer days of mechanical ventilation than a “fluid-liberal” strategy (maintaining a relatively high CVP or PCWP) in ARDS.

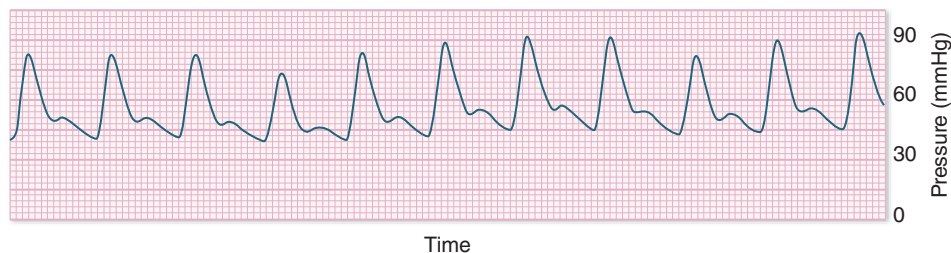


FIGURE 321-4 Pulse pressure change during mechanical ventilation in a patient with shock whose cardiac output will increase in response to intravenous fluid administration. The pulse pressure (systolic minus diastolic blood pressure) changes during mechanical ventilation in a patient with septic shock.

TYPE II RESPIRATORY FAILURE

This type of respiratory failure is a consequence of alveolar hypoventilation and results from the inability to eliminate carbon dioxide effectively. Mechanisms are categorized by impaired central nervous system (CNS) drive to breathe, impaired strength with failure of neuromuscular function in the respiratory system, and increased load(s) on the respiratory system. Reasons for diminished CNS drive to breathe include drug overdose, brainstem injury, sleep-disordered breathing, and severe hypothyroidism. Reduced strength can be due to impaired neuromuscular transmission (e.g., myasthenia gravis, Guillain-Barré syndrome, amyotrophic lateral sclerosis) or respiratory muscle weakness (e.g., myopathy, electrolyte derangements, fatigue).

The overall load on the respiratory system can be subclassified into resistive loads (e.g., bronchospasm), loads due to reduced lung compliance (e.g., alveolar edema, atelectasis, intrinsic positive end-expiratory pressure [auto-PEEP]—see below), loads due to reduced chest wall compliance (e.g., pneumothorax, pleural effusion, abdominal distention), and loads due to increased minute ventilation requirements (e.g., pulmonary embolus with increased dead-space fraction, sepsis).

The mainstays of therapy for type II respiratory failure are directed at reversing the underlying cause(s) of ventilatory failure. Noninvasive positive-pressure ventilation with a tight-fitting facial or nasal mask, with avoidance of endotracheal intubation, often stabilizes these patients. This approach has been shown to be beneficial in treating patients with exacerbations of chronic obstructive pulmonary disease; it has been tested less extensively in other kinds of respiratory failure but may be attempted nonetheless in the absence of contraindications (hemodynamic instability, inability to protect the airway, respiratory arrest).

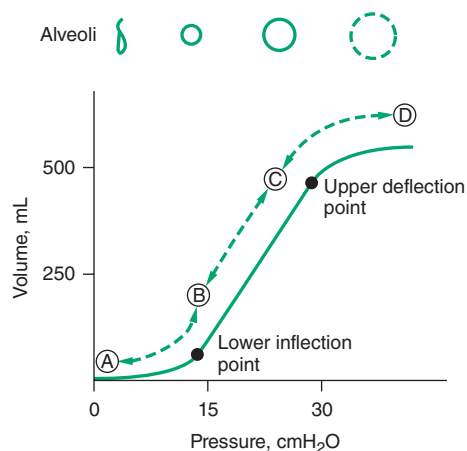


FIGURE 321-5 Pressure-volume relationship in the lungs of a patient with acute respiratory distress syndrome (ARDS). At the lower inflection point, collapsed alveoli begin to open and lung compliance changes. At the upper deflection point, alveoli become overdistended. The shape and size of alveoli are illustrated at the top of the figure.

TYPE III RESPIRATORY FAILURE

This form of respiratory failure results from lung atelectasis. Because atelectasis occurs so commonly in the perioperative period, this form is also called *perioperative respiratory failure*. After general anesthesia, decreases in functional residual capacity lead to collapse of dependent lung units. Such atelectasis can be treated by frequent changes in position, chest physiotherapy, upright positioning, and control of incisional and/or abdominal pain. Noninvasive positive-pressure ventilation may also be used to reverse regional atelectasis.

TYPE IV RESPIRATORY FAILURE

This form results from hypoperfusion of respiratory muscles in patients in shock. Normally, respiratory muscles consume <5% of total cardiac output and oxygen delivery. Patients in shock often experience respiratory distress due to pulmonary edema (e.g., in cardiogenic shock), lactic acidosis, and anemia. In this setting, up to 40% of cardiac output may be distributed to the respiratory muscles. Intubation and mechanical ventilation can allow redistribution of the cardiac output away from the respiratory muscles and back to vital organs while the shock is treated.

CARE OF THE MECHANICALLY VENTILATED PATIENT

(See also Chap. 323) Whereas a thorough understanding of the pathophysiology of respiratory failure is essential for optimal patient care, recognition of a patient's readiness to be liberated from mechanical ventilation is likewise important. Several studies have shown that daily spontaneous breathing trials can identify patients who are ready for extubation. Accordingly, all intubated, mechanically ventilated patients should undergo daily screening of respiratory function. If oxygenation is stable (i.e., Pa_{O_2}/FI_{O_2} [partial pressure of oxygen/fraction of inspired oxygen] >200 and $P\dot{E}EP \leq 5$ cmH₂O), cough and airway reflexes are intact, and no vasopressor agents or sedatives are being administered, the patient has passed the screening test and should undergo a spontaneous breathing trial. This trial consists of a period of breathing through the endotracheal tube without ventilator support (both continuous positive airway pressure [CPAP] of 5 cmH₂O and an open T-piece breathing system can be used) for 30–120 min. The spontaneous breathing trial is declared a failure and stopped if *any* of the following occur: (1) respiratory rate >35/min for >5 min, (2) O₂ saturation <90%, (3) heart rate >140/min or a 20% increase or decrease from baseline, (4) systolic blood pressure <90 mmHg or >180 mmHg, or (5) increased anxiety or diaphoresis. If, at the end of the spontaneous breathing trial, none of the above events has occurred and the ratio of the respiratory rate and tidal volume in liters (f/V_T) is <105, the patient can be extubated. Such protocol-driven approaches to patient care can have an important impact on the duration of mechanical ventilation and ICU stay. In spite of such a careful approach to liberation from mechanical ventilation, up to 10% of patients develop respiratory distress after extubation and may require resumption of mechanical ventilation. Many of these patients will require reintubation. The use of noninvasive ventilation in patients in whom extubation fails may be associated with worse outcomes than are obtained with immediate reintubation.

Mechanically ventilated patients frequently require sedatives and analgesics. Opiates are the mainstay of therapy for pain control in mechanically ventilated patients. After adequate pain control has been ensured, additional indications for sedation include anxiolysis; treatment of subjective dyspnea; psychosis; facilitation of nursing care; reduction of autonomic hyperactivity, which may precipitate myocardial ischemia; and reduction of total O₂ consumption (V_{O_2}).

Neuromuscular blocking agents are occasionally needed to facilitate mechanical ventilation in patients with profound ventilator dyssynchrony despite optimal sedation, particularly in the setting of severe

ARDS. Use of these agents may result in prolonged weakness—a myopathy known as the *postparalytic syndrome*. For this reason, neuromuscular blocking agents typically are used as a last resort when aggressive sedation fails to achieve patient-ventilator synchrony. Because neuromuscular blocking agents result in pharmacologic paralysis without altering mental status, sedative-induced amnesia is mandatory when these agents are administered.

Amnesia can be achieved reliably with benzodiazepines such as lorazepam and midazolam as well as the IV anesthetic agent propofol. Outside the setting of pharmacologic paralysis, few data support the idea that amnesia is mandatory in all patients who require intubation and mechanical ventilation. Since many of these critically ill patients have impaired hepatic and renal function, sedatives and opiates may accumulate when given for prolonged periods. A nursing protocol-driven approach to sedation of mechanically ventilated patients or daily interruption of sedative infusions paired with daily spontaneous breathing trials has been shown to prevent excessive drug accumulation and shorten the duration of both mechanical ventilation and ICU stay.

MULTIORGAN SYSTEM FAILURE

Multiorgan system failure, which is commonly associated with critical illness, is defined by the simultaneous presence of physiologic dysfunction and/or failure of two or more organs. Typically, this syndrome occurs in the setting of severe sepsis, shock of any kind, severe inflammatory conditions such as pancreatitis, and trauma. The fact that multiorgan system failure occurs commonly in the ICU is a testament to our current ability to stabilize and support single-organ failure. The ability to support single-organ failure aggressively (e.g., by mechanical ventilation or by renal replacement therapy) has reduced rates of early mortality in critical illness. As a result, it is uncommon for critically ill patients to die in the initial stages of resuscitation. Instead, many patients succumb to critical illness later in the ICU stay, after the initial presenting problem has been stabilized.

Although there is debate regarding specific definitions of organ failure, several general principles governing the syndrome of multiorgan system failure apply. First, organ failure, no matter how it is defined, must persist beyond 24 h. Second, mortality risk increases with the accrual of failing organs. Third, the prognosis worsens with increased duration of organ failure. These observations remain true across various critical care settings (e.g., medical versus surgical). SIRS is a common basis for multiorgan system failure. Although infection is a common cause of SIRS, “sterile” triggers such as pancreatitis, trauma, and burns often are invoked to explain multiorgan system failure.

MONITORING IN THE ICU

Because respiratory failure and circulatory failure are common in critically ill patients, monitoring of the respiratory and cardiovascular systems is undertaken frequently. Evaluation of respiratory gas exchange is routine in critical illness. The “gold standard” remains arterial blood-gas analysis, in which pH, PaO₂, partial pressure of carbon dioxide (PCO₂), and O₂ saturation are measured directly. With arterial blood-gas analysis, the two main functions of the lung—oxygenation of arterial blood and elimination of CO₂—can be assessed directly. In fact, the blood pH, which has a profound effect on the drive to breathe, can be assessed only by such sampling. Although sampling of arterial blood is generally safe, it may be painful and cannot provide continuous information. In light of these limitations, noninvasive monitoring of respiratory function is often employed.

PULSE OXIMETRY

The most commonly utilized noninvasive technique for monitoring respiratory function, pulse oximetry takes advantage of differences in the absorptive properties of oxygenated and deoxygenated hemoglobin. At wavelengths of 660 nm, oxyhemoglobin reflects light more effectively than does deoxyhemoglobin, whereas the reverse is true in the infrared spectrum (940 nm). A pulse oximeter passes both wavelengths of light through a perfused digit such as a finger, and the relative intensity of light transmission at these two wavelengths

is recorded. From this information, the relative percentage of oxyhemoglobin is derived. Since arterial pulsations produce phasic changes in the intensity of transmitted light, the pulse oximeter is designed to detect only light of alternating intensity. This feature allows distinction of arterial and venous blood O₂ saturations.

RESPIRATORY SYSTEM MECHANICS

Respiratory system mechanics can be measured in patients during mechanical ventilation (Chap. 323). When volume-controlled modes of mechanical ventilation are used, accompanying airway pressures can easily be measured as long as the patient is passive. The peak airway pressure is determined by two variables: airway resistance and respiratory system compliance. At the end of inspiration, inspiratory flow can be stopped transiently. This end-inspiratory pause (*plateau pressure*) is a static measurement, affected only by respiratory system compliance and not by airway resistance. Therefore, during volume-controlled ventilation, the difference between the peak (airway resistance + respiratory system compliance) and plateau (respiratory system compliance only) airway pressures provides a quantitative assessment of airway resistance. Accordingly, during volume-controlled ventilation, patients with increases in airway resistance typically have increased peak airway pressures as well as abnormally high gradients between peak and plateau airway pressures (typically >15 cmH₂O) at an inspiratory flow rate of 1 L/sec. The compliance of the respiratory system is defined by the change in pressure of the respiratory system per unit change in volume.

The respiratory system can be divided into two components: the lungs and the chest wall. Normally, respiratory system compliance is ~100 mL/cmH₂O. Pathophysiologic processes such as pleural effusions, pneumothorax, and increased abdominal girth all reduce chest wall compliance. Lung compliance may be reduced by pneumonia, pulmonary edema, interstitial lung disease, or auto-PEEP. Accordingly, patients with abnormalities in compliance of the respiratory system (lungs and/or chest wall) typically have elevated peak and plateau airway pressures but a normal gradient between these two pressures. Auto-PEEP occurs when there is insufficient time for emptying of alveoli before the next inspiratory cycle. Since the alveoli have not decompressed completely, alveolar pressure remains positive at the end of exhalation (*functional residual capacity*). This phenomenon results most commonly from critical narrowing of distal airways in disease processes such as asthma and COPD. Auto-PEEP with resulting alveolar overdistention may result in diminished lung compliance, reflected by abnormally increased plateau airway pressures. Modern mechanical ventilators allow breath-to-breath display of pressure and flow, permitting detection of problems such as patient-ventilator dyssynchrony, airflow obstruction, and auto-PEEP (Fig. 321–6).

CIRCULATORY STATUS

Oxygen delivery (Q_{O₂}) is a function of cardiac output and the content of O₂ in the arterial blood (Ca_{O₂}). The Ca_{O₂} is determined by the hemoglobin concentration, the arterial hemoglobin saturation, and dissolved O₂ not bound to hemoglobin. For normal adults:

$$\begin{aligned} Q_{O_2} &= 50 \text{ dL/min} \times (1.39 \times 15 \text{ g/dL [hemoglobin concentration]} \\ &\quad \times 1.0 \text{ [hemoglobin \% saturation]} + 0.0031 \times 100 \text{ [Pa}_{O_2}\text{]}) \\ &= 50 \text{ dL/min (cardiac output)} \times 21.6 \text{ mL O}_2 \text{ per dL blood (Ca}_{O_2}\text{)} \\ &= 1058 \text{ mL O}_2 \text{ per min} \end{aligned}$$

It is apparent that nearly all of the O₂ delivered to tissues is bound to hemoglobin and that the dissolved O₂ (Pa_{O₂}) contributes very little to O₂ content in arterial blood or to O₂ delivery. Normally, the content of O₂ in mixed venous blood (C \bar{v} _{O₂}) is 15.76 mL/dL since the mixed venous blood is 75% saturated. Therefore, the normal tissue extraction ratio for O₂ is Ca_{O₂} – C \bar{v} _{O₂}/Ca_{O₂} ((21.16–15.76)/21.16) or ~25%. A pulmonary artery catheter allows measurements of O₂ delivery and the O₂ extraction ratio.

Information on the mixed venous O₂ saturation allows assessment of global tissue perfusion. A reduced mixed venous O₂ saturation may be caused by inadequate cardiac output, reduced hemoglobin concentration, and/or reduced arterial O₂ saturation. An abnormally high V_{O₂} may also lead to a reduced mixed venous O₂ saturation if O₂ delivery is

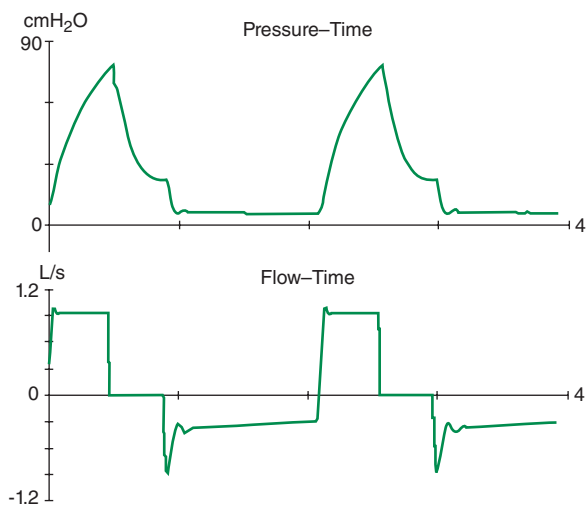


FIGURE 321-6 Increased airway resistance with auto-PEEP. The top waveform (airway pressure vs. time) shows a large difference between the peak airway pressure (80 cmH₂O) and the plateau airway pressure (20 cmH₂O). The bottom waveform (flow vs. time) demonstrates airflow throughout expiration (reflected by the flow tracing on the negative portion of the abscissa) that persists up to the next inspiratory effort.

not concomitantly increased. Abnormally increased V_{O_2} in peripheral tissues may be caused by problems such as fever, agitation, shivering, and thyrotoxicosis.

The pulmonary artery catheter originally was designed as a tool to guide therapy for acute myocardial infarction but has been used in the ICU for evaluation and treatment of a variety of other conditions, such as ARDS, septic shock, congestive heart failure, and acute renal failure. This device has never been validated as a tool associated with reduction in morbidity and mortality rates. Indeed, despite numerous prospective studies, mortality or morbidity rate benefits associated with use of the pulmonary artery catheter have never been reported in any setting. Accordingly, it appears that routine pulmonary artery catheterization is not indicated as a means of monitoring and characterizing circulatory status in most critically ill patients.

Static measurements of circulatory parameters (e.g., CVP, PCWP) do not provide reliable information on the circulatory status of critically ill patients. In contrast, dynamic assessments measuring the impact of breathing on the circulation are more reliable predictors of responsiveness to IV fluid administration. A decrease in CVP of >1 mmHg during inspiration in a spontaneously breathing patient may predict an increase in cardiac output after IV fluid administration. Similarly, a changing pulse pressure during mechanical ventilation has been shown to predict an increase in cardiac output after IV fluid administration in patients with septic shock.

PREVENTION OF COMPLICATIONS OF CRITICAL ILLNESS

SEPSIS IN THE CRITICAL CARE UNIT

(See also Chap. 325) Sepsis, defined as the presence of SIRS in the setting of known or suspected infection, is a significant problem in the care of critically ill patients, who often progress to severe sepsis with the failure of one or more organs. Sepsis is the leading cause of death in noncoronary ICUs in the United States, with case rates expected to increase as the population ages and a higher percentage of people are vulnerable to infection.

NOSOCOMIAL INFECTIONS IN THE ICU

Many therapeutic interventions in the ICU are invasive and predispose patients to infectious complications. These interventions include endotracheal intubation, indwelling vascular catheters, transurethral bladder catheters, and other catheters placed into sterile body cavities (e.g., tube thoracostomy, percutaneous intraabdominal drainage

catheterization). The longer such devices remain in place, the more prone to these infections patients become. For example, ventilator-associated pneumonia correlates strongly with the duration of intubation and mechanical ventilation. Therefore, an important aspect of preventive care is the timely removal of invasive devices as soon as they are no longer needed. Moreover, multidrug-resistant organisms are commonplace in the ICU.

Infection control is critical in the ICU. Care bundles, which include measures such as frequent hand washing, are effective but underutilized strategies. Other components of care bundles, such as protective isolation of patients colonized or infected by drug-resistant organisms, are also commonly used. Silver-coated endotracheal tubes reportedly reduce the incidence of ventilator-associated pneumonia. Studies evaluating multifaceted, evidence-based strategies to decrease catheter-related bloodstream infections have shown improved outcomes with strict adherence to measures such as hand washing, full-barrier precautions during catheter insertion, chlorhexidine skin preparation, avoidance of the femoral site, and timely catheter removal.

DEEP VEIN THROMBOSIS (DVT)

(See also Chap. 300) All ICU patients are at high risk for this complication because of their predilection for immobility. Therefore, all should receive some form of prophylaxis against DVT. The most commonly employed forms of prophylaxis are subcutaneous low-dose heparin injections and sequential compression devices for the lower extremities. Observational studies report an alarming incidence of DVTs despite the use of these standard prophylactic regimens. Furthermore, heparin prophylaxis may result in heparin-induced thrombocytopenia, another nosocomial complication in critically ill patients.

Low-molecular-weight heparins such as enoxaparin are more effective than unfractionated heparin for DVT prophylaxis in high-risk patients (e.g., those undergoing orthopedic surgery) and are associated with a lower incidence of heparin-induced thrombocytopenia. Fondaparinux, a selective factor Xa inhibitor, is even more effective than enoxaparin in high-risk orthopedic patients.

STRESS ULCERS

Prophylaxis against stress ulcers is frequently administered in most ICUs; typically, histamine-2 antagonists or proton pump inhibitors are given. Available data suggest that high-risk patients, such as those with coagulopathy, shock, or respiratory failure requiring mechanical ventilation, benefit from such prophylactic treatment.

NUTRITION AND GLYCEMIC CONTROL

These are important issues that may be associated with respiratory failure, impaired wound healing, and dysfunctional immune response in critically ill patients. Early enteral feeding is reasonable, though no data are available to suggest that this treatment improves patient outcome per se. Certainly, enteral feeding, if possible, is preferred over parenteral nutrition, which is associated with numerous complications, including hyperglycemia, fatty liver, cholestasis, and sepsis. When parenteral feeding is necessary to supplement enteral nutrition, delaying this intervention until day 8 in the ICU results in better recovery and fewer ICU-related complications. Tight glucose control is an area of controversy in critical care. Although one study showed a significant mortality benefit when glucose levels were aggressively normalized in a large group of surgical ICU patients, more recent data for a large population of both medical and surgical ICU patients suggested that tight glucose control resulted in increased rates of mortality.

ICU-ACQUIRED WEAKNESS

ICU-acquired weakness occurs frequently in patients who survive critical illness, particularly those with SIRS and/or sepsis. Both neuropathies and myopathies have been described, most commonly after ~1 week in the ICU. The mechanisms behind ICU-acquired weakness syndromes are poorly understood. Intensive insulin therapy may reduce polyneuropathy in critical illness. Very early physical and occupational therapy in mechanically ventilated patients reportedly results in significant improvements in functional independence at hospital

discharge as well as in reduced durations of mechanical ventilation and delirium.

ANEMIA

Studies have shown that most ICU patients are anemic as a result of chronic inflammation. Phlebotomy also contributes to ICU anemia. A large multicenter study involving patients in many different ICU settings challenged the conventional notion that a hemoglobin level of 100 g/L (10 g/dL) is needed in critically ill patients, with similar outcomes noted in those whose transfusion trigger was 7 g/dL. Red blood cell transfusion is associated with impairment of immune function and increased risk of infections as well as of ARDS and volume overload, all of which may explain the findings in this study. Recently, a conservative transfusion strategy enhanced survival among patients with active upper gastrointestinal hemorrhage.

ACUTE RENAL FAILURE

(See also Chap. 334) Acute renal failure occurs in a significant percentage of critically ill patients. The most common underlying etiology is acute tubular necrosis, usually precipitated by hypoperfusion and/or nephrotoxic agents. Currently, no pharmacologic agents are available for prevention of renal injury in critical illness. Studies have shown convincingly that low-dose dopamine is *not* effective in protecting the kidneys from acute injury.

NEUROLOGIC DYSFUNCTION IN CRITICALLY ILL PATIENTS

DELIRIUM

(See also Chaps. 34 and 328) This state is defined by (1) an acute onset of changes or fluctuations in mental status, (2) inattention, (3) disorganized thinking, and (4) an altered level of consciousness (i.e., a state other than alertness). Delirium is reported to occur in a wide range of mechanically ventilated ICU patients and can be detected by the Confusion Assessment Method (CAM)-ICU or the Intensive Care Delirium Screening Checklist. These tools are used to ask patients to answer simple questions and perform simple tasks and can be used readily at the bedside. The differential diagnosis of delirium in ICU patients is broad and includes infectious etiologies (including sepsis), medications (particularly sedatives and analgesics), drug withdrawal, metabolic/electrolyte derangements, intracranial pathology (e.g., stroke, intracranial hemorrhage), seizures, hypoxia, hypertensive crisis, shock, and vitamin deficiencies (particularly thiamine). Patients with ICU delirium have increases in length of hospital stay, time on mechanical ventilation, cognitive impairment at hospital discharge, and 6-month mortality rate. Interventions to reduce ICU delirium are limited. The sedative dexmedetomidine has been less strongly associated with ICU delirium than midazolam. In addition, as mentioned above, very early physical and occupational therapy in mechanically ventilated patients has been demonstrated to reduce delirium.

ANOXIC CEREBRAL INJURY

(See also Chap. 330) This condition is common after cardiac arrest and often results in severe and permanent brain injury in survivors. Active cooling of patients after cardiac arrest has been shown to improve neurologic outcomes. Therefore, patients who present to the ICU after circulatory arrest from ventricular fibrillation or pulseless ventricular tachycardia should be actively cooled to achieve a core body temperature of 32–34°C.

STROKE

(See also Chap. 446) Stroke is a common cause of neurologic critical illness. Hypertension must be managed carefully, since abrupt reductions in blood pressure may be associated with further brain ischemia and injury. Acute ischemic stroke treated with tissue plasminogen activator (tPA) has an improved neurologic outcome when treatment is given within 3 h of onset of symptoms. The mortality rate is not reduced when tPA is compared with placebo, despite the improved neurologic outcome. The risk of cerebral hemorrhage is significantly higher in patients given tPA. No benefit is seen when tPA therapy is given beyond 3 h after symptom onset. Heparin has not been convincingly shown to improve

outcomes in patients with acute ischemic stroke. Decompressive craniectomy is a surgical procedure that relieves increased intracranial pressure in the setting of space-occupying brain lesions or brain swelling from stroke; available evidence suggests that this procedure may improve survival among select patients (≤ 55 years of age), albeit at a cost of increased disability for some.

SUBARACHNOID HEMORRHAGE

(See also Chap. 446) Subarachnoid hemorrhage may occur secondary to aneurysm rupture and is often complicated by cerebral vasospasm, re-bleeding, and hydrocephalus. Vasospasm can be detected by either transcranial Doppler assessment or cerebral angiography; it is typically treated with the calcium channel blocker nimodipine, aggressive IV fluid administration, and therapy aimed at increasing blood pressure, typically with vasoactive drugs such as phenylephrine. The IV fluids and vasoactive drugs (hypertensive hypervolemic therapy) are used to overcome the cerebral vasospasm. Early surgical clipping or endovascular coiling of aneurysms is advocated to prevent complications related to re-bleeding. Hydrocephalus, typically heralded by a decreased level of consciousness, may require ventriculostomy drainage.

STATUS EPILEPTICUS

(See also Chap. 445) Recurrent or relentless seizure activity is a medical emergency. Cessation of seizure activity is required to prevent irreversible neurologic injury. Lorazepam is the most effective benzodiazepine for treating status epilepticus and is the treatment of choice for controlling seizures acutely. Phenytoin or fosphenytoin should be given concomitantly since lorazepam has a short half-life. Other drugs, such as gabapentin, carbamazepine, and phenobarbital, should be reserved for patients with contraindications to phenytoin (e.g., allergy or pregnancy) or ongoing seizures despite phenytoin.

BRAIN DEATH

(See also Chap. 330) Although deaths of critically ill patients usually are attributable to irreversible cessation of circulatory and respiratory function, a diagnosis of death also may be established by irreversible cessation of all functions of the entire brain, including the brainstem, even if circulatory and respiratory functions remain intact on artificial life support. Such a diagnosis requires demonstration of the absence of cerebral function (no response to any external stimulus) and brainstem functions (e.g., unreactive pupils, lack of ocular movement in response to head turning or ice-water irrigation of ear canals, positive apnea test [no drive to breathe]). Absence of brain function must have an established cause and be permanent without possibility of recovery; a sedative effect, hypothermia, hypoxemia, neuromuscular paralysis, and severe hypotension must be ruled out. If there is uncertainty about the cause of coma, studies of cerebral blood flow and electroencephalography should be performed.

WITHHOLDING OR WITHDRAWING CARE

(See also Chap. 10) Withholding or withdrawal of care occurs commonly in the ICU setting. The Task Force on Ethics of the Society of Critical Care Medicine reported that it is ethically sound to withhold or withdraw care if a patient or the patient's surrogate makes such a request or if the physician judges that the goals of therapy are not achievable. Since all medical treatments are justified by their expected benefits, the loss of such an expectation justifies the act of withdrawing or withholding such treatment; these two actions are judged to be fundamentally similar. An underlying stipulation derived from this report is that an informed patient should have his or her wishes respected with regard to life-sustaining therapy. Implicit in this stipulation is the need to ensure that patients are thoroughly and accurately informed regarding the plausibility and expected results of various therapies.

The act of informing patients and/or surrogate decision-makers is the responsibility of the physician and other health care providers. If a patient or surrogate desires therapy deemed futile by the treating physician, the physician is not obligated ethically to provide such treatment. Rather, arrangements may be made to transfer the patient's care to another care provider. Whether the decision to withdraw

1736 life support should be initiated by the physician or left to surrogate decision-makers alone is not clear. One study reported that slightly more than half of surrogate decision-makers preferred to receive such a recommendation, whereas the rest did not. Critical care providers should meet regularly with patients and/or surrogates to discuss prognosis when the withholding or withdrawal of care is being considered. After a consensus among caregivers has been reached, this information should be relayed to the patient and/or surrogate decision-maker. If a decision to withhold or withdraw life-sustaining care for a patient has been made, aggressive attention to analgesia and anxiolysis is needed.

322 Acute Respiratory Distress Syndrome

Bruce D. Levy, Augustine M. K. Choi

Acute respiratory distress syndrome (ARDS) is a clinical syndrome of severe dyspnea of rapid onset, hypoxemia, and diffuse pulmonary infiltrates leading to respiratory failure. ARDS is caused by diffuse lung injury from many underlying medical and surgical disorders. The lung injury may be direct, as occurs in toxic inhalation, or indirect, as occurs in sepsis (Table 322-1). The clinical features of ARDS are listed in Table 322-2. By expert consensus, ARDS is defined by three categories based on the degrees of hypoxemia (Table 322-2). These stages of mild, moderate, and severe ARDS are associated with mortality risk and with the duration of mechanical ventilation in survivors.

The annual incidence of ARDS is estimated to be as high as 60 cases/100,000 population. Approximately 10% of all intensive care unit (ICU) admissions involve patients with acute respiratory failure; ~20% of these patients meet the criteria for ARDS.

ETIOLOGY

While many medical and surgical illnesses have been associated with the development of ARDS, most cases (>80%) are caused by a relatively small number of clinical disorders: severe sepsis syndrome and/or bacterial pneumonia (~40–50%), trauma, multiple transfusions, aspiration of gastric contents, and drug overdose. Among patients with trauma, the most frequently reported surgical conditions in ARDS are pulmonary contusion, multiple bone fractures, and chest wall trauma/flail chest, whereas head trauma, near-drowning, toxic inhalation, and burns are rare causes. The risks of developing ARDS are increased in patients with more than one predisposing medical or surgical condition.

Several other clinical variables have been associated with the development of ARDS. These include older age, chronic alcohol abuse, metabolic acidosis, and severity of critical illness. Trauma patients with an Acute Physiology and Chronic Health Evaluation (APACHE) II score ≥ 16 (Chap. 321) have a 2.5-fold increased risk of developing ARDS, and those with a score >20 have an incidence of ARDS that

TABLE 322-1 CLINICAL DISORDERS COMMONLY ASSOCIATED WITH ARDS

Direct Lung Injury	Indirect Lung Injury
Pneumonia	Sepsis
Aspiration of gastric contents	Severe trauma
Pulmonary contusion	Multiple bone fractures
Near-drowning	Flail chest
Toxic inhalation injury	Head trauma
	Burns
	Multiple transfusions
	Drug overdose
	Pancreatitis
	Postcardiopulmonary bypass

TABLE 322-2 DIAGNOSTIC CRITERIA FOR ARDS

Severity: Oxygenation	Onset	Chest Radiograph	Absence of Left Atrial Hypertension
Mild: 200 mmHg < $P_{aO_2}/F_{iO_2} \leq 300$ mmHg	Acute	Bilateral alveolar or interstitial infiltrates	PCWP ≤ 18 mmHg or no clinical evidence of increased left atrial pressure
Moderate: 100 mmHg < $P_{aO_2}/F_{iO_2} \leq 200$ mmHg			
Severe: $P_{aO_2}/F_{iO_2} \leq 100$ mmHg			

Abbreviations: ARDS, acute respiratory distress syndrome; F_{iO_2} , inspired O_2 percentage; P_{aO_2} , arterial partial pressure of O_2 ; PCWP, pulmonary capillary wedge pressure.

is more than threefold greater than the incidence among those with APACHE II scores ≤ 9 .

CLINICAL COURSE AND PATHOPHYSIOLOGY

The natural history of ARDS is marked by three phases—exudative, proliferative, and fibrotic—that each have characteristic clinical and pathologic features (Fig. 322-1).

Exudative Phase In this phase (Fig. 322-2), alveolar capillary endothelial cells and type I pneumocytes (alveolar epithelial cells) are injured, with consequent loss of the normally tight alveolar barrier to fluid and macromolecules. Edema fluid that is rich in protein accumulates in the interstitial and alveolar spaces. Significant concentrations of cytokines (e.g., interleukin 1, interleukin 8, and tumor necrosis factor α) and lipid mediators (e.g., leukotriene B_4) are present in the lung in this acute phase. In response to proinflammatory mediators, leukocytes (especially neutrophils) traffic into the pulmonary interstitium and alveoli. In addition, condensed plasma proteins aggregate in the air spaces with cellular debris and dysfunctional pulmonary surfactant to form hyaline membrane whorls. Pulmonary vascular injury also occurs early in ARDS, with vascular obliteration by microthrombi and fibrocellular proliferation (Fig. 322-3).

Alveolar edema predominantly involves *dependent* portions of the lung, with diminished aeration and atelectasis. Collapse of large sections of dependent lung markedly decreases lung compliance. Consequently, intrapulmonary shunting and hypoxemia develop and the work of breathing increases, leading to dyspnea. The pathophysiologic alterations in alveolar spaces are exacerbated by microvascular occlusion that results in reductions in pulmonary arterial blood flow to ventilated portions of the lung (and thus in increased dead space) and in pulmonary hypertension. Thus, in addition to severe hypoxemia, hypercapnia secondary to an increase in pulmonary dead space is prominent in early ARDS.

The exudative phase encompasses the first 7 days of illness after exposure to a precipitating ARDS risk factor, with the patient experiencing the onset of respiratory symptoms. Although usually presenting within 12–36 h after the initial insult, symptoms can be delayed by 5–7 days. Dyspnea develops, with a sensation of rapid shallow breathing

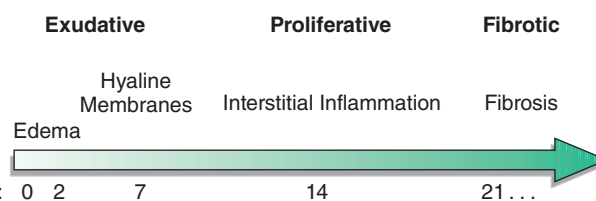


FIGURE 322-1 Diagram illustrating the time course for the development and resolution of ARDS. The exudative phase is notable for early alveolar edema and neutrophil-rich leukocytic infiltration of the lungs, with subsequent formation of hyaline membranes from diffuse alveolar damage. Within 7 days, a proliferative phase ensues with prominent interstitial inflammation and early fibrotic changes. Approximately 3 weeks after the initial pulmonary injury, most patients recover. However, some patients enter the fibrotic phase, with substantial fibrosis and bullae formation.



FIGURE 322-2 A representative anteroposterior chest x-ray in the exudative phase of ARDS shows diffuse interstitial and alveolar infiltrates that can be difficult to distinguish from left ventricular failure.

and an inability to get enough air. Tachypnea and increased work of breathing result frequently in respiratory fatigue and ultimately in respiratory failure. Laboratory values are generally nonspecific and are primarily indicative of underlying clinical disorders. The chest radiograph usually reveals alveolar and interstitial opacities involving at least three-quarters of the lung fields (Fig. 322-2). While characteristic for ARDS, these radiographic findings are not specific and can be indistinguishable from cardiogenic pulmonary edema (Chap. 326). Unlike the latter, however, the chest x-ray in ARDS rarely shows cardiomegaly, pleural effusions, or pulmonary vascular redistribution. Chest CT in ARDS reveals extensive heterogeneity of lung involvement (Fig. 322-4).

Because the early features of ARDS are nonspecific, alternative diagnoses must be considered. In the differential diagnosis of ARDS, the most common disorders are cardiogenic pulmonary edema, diffuse pneumonia, and alveolar hemorrhage. Less common diagnoses to consider include acute interstitial lung diseases (e.g., acute interstitial pneumonitis; Chap. 315), acute immunologic injury (e.g., hypersensitivity pneumonitis; Chap. 310), toxin injury (e.g., radiation pneumonitis; Chap. 263), and neurogenic pulmonary edema (Chap. 47e).

Proliferative Phase This phase of ARDS usually lasts from day 7 to day 21. Most patients recover rapidly and are liberated from mechanical

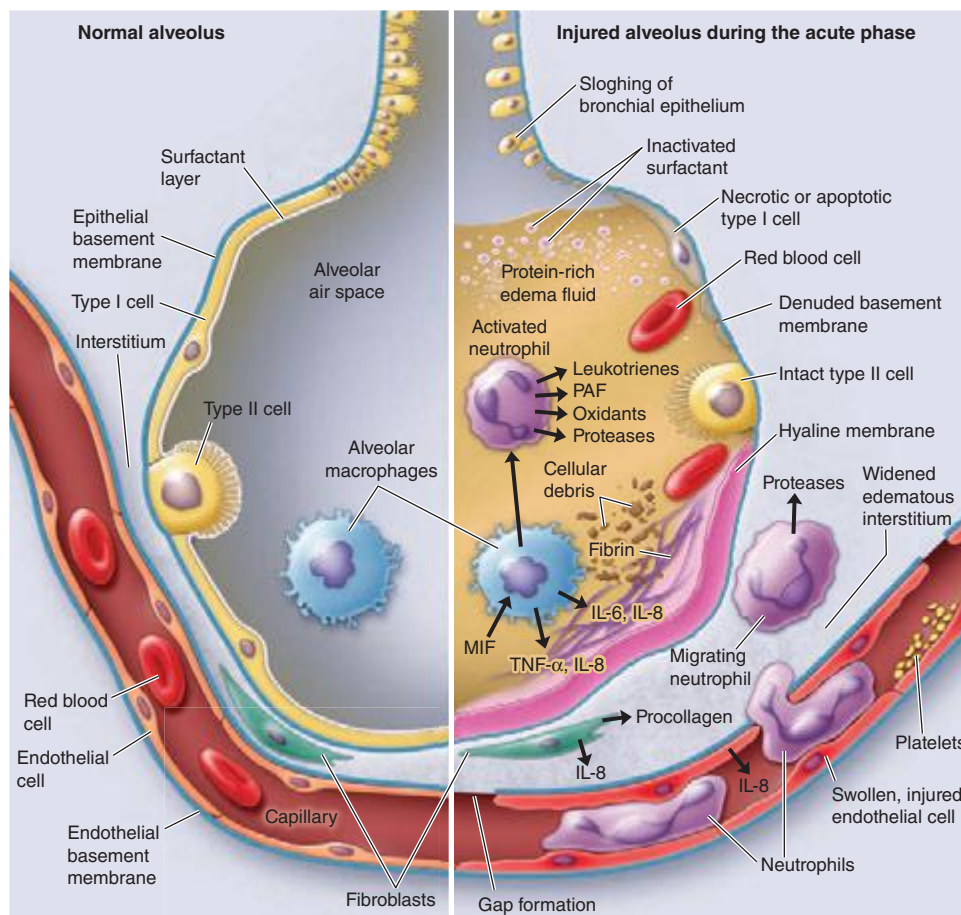


FIGURE 322-3 The normal alveolus (left) and the injured alveolus in the acute phase of acute lung injury and the acute respiratory distress syndrome (right). In the acute phase of the syndrome (right), there is sloughing of both the bronchial and alveolar epithelial cells, with the formation of protein-rich hyaline membranes on the denuded basement membrane. Neutrophils are shown adhering to the injured capillary endothelium and transmigrating through the interstitium into the air space, which is filled with protein-rich edema fluid. In the air space, an alveolar macrophage is secreting cytokines—i.e., interleukins 1, 6, 8, and 10 (IL-1, -6, -8, and -10) and tumor necrosis factor α (TNF- α)—that act locally to stimulate chemotaxis and activate neutrophils. Macrophages also secrete other cytokines, including IL-1, -6, and -10. IL-1 can also stimulate the production of extracellular matrix by fibroblasts. Neutrophils can release oxidants, proteases, leukotrienes, and other proinflammatory molecules, such as platelet-activating factor (PAF). A number of antiinflammatory mediators are also present in the alveolar milieu, including the IL-1-receptor antagonist, soluble TNF- α receptor, autoantibodies to IL-8, and cytokines such as IL-10 and IL-11 (not shown). The influx of protein-rich edema fluid into the alveolus has led to the inactivation of surfactant. MIF, macrophage inhibitory factor. (From LB Ware, MA Matthay: *N Engl J Med* 342:1334, 2000, with permission.)



FIGURE 322-4 A representative CT scan of the chest during the exudative phase of ARDS, in which *dependent* alveolar edema and atelectasis predominate.

ventilation during this phase. Despite this improvement, many patients still experience dyspnea, tachypnea, and hypoxemia. Some patients develop progressive lung injury and early changes of pulmonary fibrosis during the proliferative phase. Histologically, the first signs of resolution are often evident in this phase, with the initiation of lung repair, the organization of alveolar exudates, and a shift from a neutrophil- to a lymphocyte-predominant pulmonary infiltrate. As part of the reparative process, type II pneumocytes proliferate along alveolar basement membranes. These specialized epithelial cells synthesize new pulmonary surfactant and differentiate into type I pneumocytes.

Fibrotic Phase While many patients with ARDS recover lung function 3–4 weeks after the initial pulmonary injury, some enter a fibrotic phase that may require long-term support on mechanical ventilators and/or supplemental oxygen. Histologically, the alveolar edema and inflammatory exudates of earlier phases are now converted to extensive alveolar-duct and interstitial fibrosis. Marked disruption of acinar architecture leads to emphysema-like changes, with large bullae. Intimal fibroproliferation in the pulmonary microcirculation causes progressive vascular occlusion and pulmonary hypertension. The physiologic consequences include an increased risk of pneumothorax, reductions in lung compliance, and increased pulmonary dead space. Patients in this late phase experience a substantial burden of excess morbidity. Lung biopsy evidence for pulmonary fibrosis in any phase of ARDS is associated with increased mortality risk.

TREATMENT ACUTE RESPIRATORY DISTRESS SYNDROME

GENERAL PRINCIPLES

Recent reductions in ARDS mortality rates are largely the result of general advances in the care of critically ill patients (Chap. 321). Thus, caring for these patients requires close attention to (1) the recognition and treatment of underlying medical and surgical disorders (e.g., sepsis, aspiration, trauma); (2) the minimization of procedures and their complications; (3) prophylaxis against venous thromboembolism, gastrointestinal bleeding, aspiration, excessive sedation, and central venous catheter infections; (4) prompt recognition of nosocomial infections; and (5) provision of adequate nutrition.

MANAGEMENT OF MECHANICAL VENTILATION

(See also Chap. 323) Patients meeting clinical criteria for ARDS frequently become fatigued from increased work of breathing and progressive hypoxemia, requiring mechanical ventilation for support.

Ventilator-Induced Lung Injury Despite its life-saving potential, mechanical ventilation can aggravate lung injury. Experimental models have demonstrated that ventilator-induced lung injury appears to require two processes: repeated alveolar overdistention

and recurrent alveolar collapse. As is clearly evident from chest CT (Fig. 322-4), ARDS is a heterogeneous disorder, principally involving dependent portions of the lung with relative sparing of other regions. Because compliance differs in affected versus more “normal” areas of the lung, attempts to fully inflate the consolidated lung may lead to overdistention of and injury to the more normal areas. Ventilator-induced injury can be demonstrated in experimental models of acute lung injury, with high-tidal-volume (V_T) ventilation resulting in additional, synergistic alveolar damage.

A large-scale, randomized controlled trial sponsored by the National Institutes of Health and conducted by the ARDS Network compared low V_T ventilation (6 mL/kg of predicted body weight) to conventional V_T ventilation (12 mL/kg predicted body weight). The mortality rate was significantly lower in the low V_T patients (31%) than in the conventional V_T patients (40%). This improvement in survival represents the most substantial ARDS-mortality benefit that has been demonstrated for *any* therapeutic intervention to date.

Prevention of Alveolar Collapse In ARDS, the presence of alveolar and interstitial fluid and the loss of surfactant can lead to a marked reduction of lung compliance. Without an increase in end-expiratory pressure, significant alveolar collapse can occur at end-expiration, with consequent impairment of oxygenation. In most clinical settings, positive end-expiratory pressure (PEEP) is empirically set to minimize F_{iO_2} (inspired O_2 percentage) and maximize P_{aO_2} (arterial partial pressure of O_2). On most modern mechanical ventilators, it is possible to construct a static pressure–volume curve for the respiratory system. The lower inflection point on the curve represents alveolar opening (or “recruitment”). The pressure at this point, usually 12–15 mmHg in ARDS, is a theoretical “optimal PEEP” for alveolar recruitment. Titration of the PEEP to the lower inflection point on the static pressure–volume curve has been hypothesized to keep the lung open, improving oxygenation and protecting against lung injury. Three large randomized trials have investigated the utility of PEEP-based strategies to keep the lung open. In all three trials, improvement in lung function was evident but overall mortality rates were not altered significantly. Until more data become available on the clinical utility of high PEEP, it is advisable to set PEEP to minimize F_{iO_2} and optimize P_{aO_2} (Chap. 323). Measurement of esophageal pressures to estimate transpulmonary pressure may help identify an optimal PEEP in some cases.

Oxygenation can also be improved by increasing mean airway pressure with *inverse-ratio ventilation*. In this technique, the inspiratory time (I) is lengthened so that it is longer than the expiratory time (E)—that is, $I:E > 1:1$. With diminished time to exhale, dynamic hyperinflation leads to increased end-expiratory pressure, similar to ventilator-prescribed PEEP. This mode of ventilation has the advantage of improving oxygenation with lower peak pressures than are required for conventional ventilation. Although inverse-ratio ventilation can improve oxygenation and can help reduce F_{iO_2} to ≤ 0.60 , thus avoiding possible oxygen toxicity, no benefit in ARDS mortality risk has been demonstrated. Recruitment maneuvers that transiently increase PEEP to “recruit” atelectatic lung can also increase oxygenation, but a mortality benefit has not been established.

In several randomized trials, mechanical ventilation in the prone position improved arterial oxygenation, but its effect on survival and other important clinical outcomes remains uncertain. Moreover, unless the critical-care team is experienced in “proning,” repositioning critically ill patients can be hazardous, leading to accidental endotracheal extubation, loss of central venous catheters, and orthopedic injury.

OTHER STRATEGIES IN MECHANICAL VENTILATION

Several additional mechanical-ventilation strategies that use specialized equipment have been tested in ARDS patients; most of these approaches have had mixed or disappointing results in adults. *High-frequency ventilation* (HFV) entails ventilating at extremely high respiratory rates (5–20 cycles per second) and low V_T s (1–2 mL/kg). Use of *partial liquid ventilation* (PLV) with perfluorocarbon—an inert, high-density liquid that easily solubilizes oxygen and carbon

dioxide—has yielded promising preliminary results, enhancing pulmonary function in patients with ARDS, but also has provided no survival benefit. *Lung-replacement therapy with extracorporeal membrane oxygenation* (ECMO), which provides a clear survival benefit in neonatal respiratory distress syndrome, may also have utility in selected adult patients with ARDS.

Data supporting the efficacy of “adjunctive” ventilator therapies (e.g., high PEEP, inverse ratio ventilation, recruitment maneuvers, prone positioning, HFV, ECMO, and PLV) remain incomplete. Accordingly, these modalities are reserved for use as rescue rather than primary therapies.

FLUID MANAGEMENT

(See also Chap. 321) Increased pulmonary vascular permeability leading to interstitial and alveolar edema fluid rich in protein is a central feature of ARDS. In addition, impaired vascular integrity augments the normal increase in extravascular lung water that occurs with increasing left atrial pressure. Maintaining a low left atrial filling pressure minimizes pulmonary edema and prevents further decrements in arterial oxygenation and lung compliance; improves pulmonary mechanics; shortens ICU stay and the duration of mechanical ventilation; and is associated with a lower mortality rate in both medical and surgical ICU patients. Thus, aggressive attempts to reduce left atrial filling pressures with fluid restriction and diuretics should be an important aspect of ARDS management, limited only by hypotension and hypoperfusion of critical organs such as the kidneys.

NEUROMUSCULAR BLOCKADE

In severe ARDS, sedation alone can be inadequate for the patient-ventilator synchrony required for lung-protective ventilation. This clinical problem was recently addressed in a multicenter, randomized, placebo-controlled trial of early neuromuscular blockade (with cisatracurium besylate) for 48 h. In severe ARDS, early neuromuscular blockade increased the rate of survival and ventilator-free days without increasing ICU-acquired paresis. These promising findings support the early administration of neuromuscular blockade if needed to facilitate mechanical ventilation in severe ARDS; however, these results must be replicated prior to their widespread application in clinical practice.

GLUCOCORTICOIDS

Many attempts have been made to treat both early and late ARDS with glucocorticoids, with the goal of reducing potentially deleterious pulmonary inflammation. Few studies have shown any benefit. Current evidence does *not* support the use of high-dose glucocorticoids in the care of ARDS patients.

OTHER THERAPIES

Clinical trials of surfactant replacement and multiple other medical therapies have proved disappointing. Inhaled nitric oxide and inhaled epoprostenol sodium can transiently improve oxygenation but do not improve survival or decrease time on mechanical ventilation.

RECOMMENDATIONS

Many clinical trials have been undertaken to improve the outcome of patients with ARDS; most have been unsuccessful in modifying the natural history. While results of large clinical trials must be judiciously applied to *individual* patients, evidence-based recommendations are summarized in Table 322-3, and an algorithm for the initial therapeutic goals and limits in ARDS management is provided in Fig. 322-5.

PROGNOSIS

Mortality Recent mortality estimates for ARDS range from 26% to 44%. There is substantial variability, but a trend toward improved ARDS outcomes appears evident. Of interest, mortality in ARDS is largely attributable to nonpulmonary causes, with sepsis and nonpulmonary organ failure accounting for >80% of deaths. Thus, improvement in survival is

TABLE 322-3 EVIDENCE-BASED RECOMMENDATIONS FOR ARDS THERAPIES

Treatment	Recommendation ^a
Mechanical ventilation	
Low tidal volume	A
Minimized left atrial filling pressures	B
High-PEEP or “open lung”	C
Prone position	C
Recruitment maneuvers	C
High-frequency ventilation	D
ECMO	C
Early neuromuscular blockade	A
Glucocorticoid treatment	D
Surfactant replacement, inhaled NO, inhaled epoprostenol, and other anti-inflammatory therapy (e.g., ketoconazole, PGE1, NSAIDs)	D

^aKey: A, recommended therapy based on strong clinical evidence from randomized clinical trials; B, recommended therapy based on supportive but limited clinical data; C, recommended only as alternative therapy on the basis of indeterminate evidence; D, not recommended on the basis of clinical evidence against efficacy of therapy.

Abbreviations: ARDS, acute respiratory distress syndrome; ECMO, extracorporeal membrane oxygenation; NO, nitric oxide; NSAIDs, nonsteroidal anti-inflammatory drugs; PEEP, positive end-expiratory pressure; PGE1, prostaglandin E₁.

likely secondary to advances in the care of septic/infected patients and those with multiple organ failure (Chap. 321).

The major risk factors for ARDS mortality are nonpulmonary. Advanced age is an important risk factor. Patients >75 years of age have a substantially higher mortality risk (~60%) than those <45 (~20%). Moreover, patients >60 years of age with ARDS and sepsis have a threefold higher mortality risk than those <60. Other risk factors include preexisting organ dysfunction from chronic medical illness—in particular, chronic liver disease, cirrhosis, chronic alcohol abuse, chronic immunosuppression, sepsis, chronic renal disease, failure of any nonpulmonary organ, and increased APACHE III scores (Chap. 321). Patients with ARDS arising from direct lung injury (including pneumonia, pulmonary contusion, and aspiration;

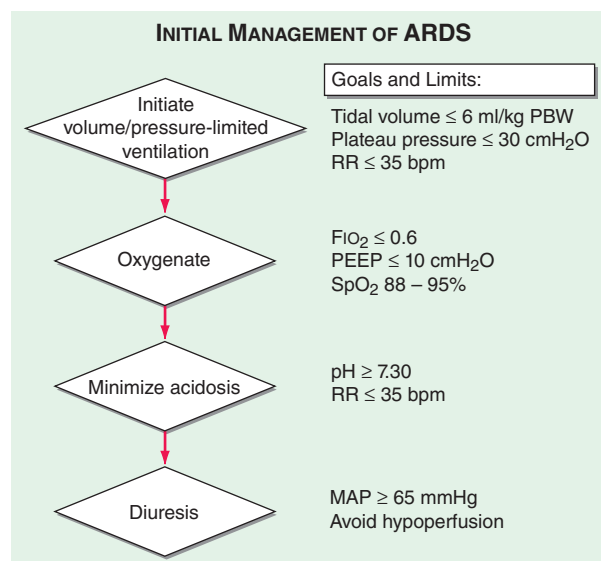


FIGURE 322-5 Algorithm for the initial management of ARDS.

Clinical trials have provided evidence-based therapeutic goals for a stepwise approach to the early mechanical ventilation, oxygenation, and correction of acidosis and diuresis of critically ill patients with ARDS. FiO₂, inspired O₂ percentage; MAP, mean arterial pressure; PBW, predicted body weight; PEEP, positive end expiratory pressure; RR, respiratory rate; SpO₂, arterial oxyhemoglobin saturation measured by pulse oximetry.

1740 Table 322-1) are nearly twice as likely to die as those with indirect causes of lung injury, while surgical and trauma patients with ARDS—especially those without direct lung injury—have a higher survival rate than other ARDS patients.

An early (within 24 h of presentation) elevation in pulmonary dead space (>0.60) and severe arterial hypoxemia (P_{aO_2}/F_{iO_2} , <100 mmHg) predict increased mortality risk from ARDS; however, there is surprisingly little additional value in predicting ARDS mortality from other measures of the severity of lung injury, including the level of PEEP (≥ 10 cm H_2O), respiratory system compliance (≤ 40 mL/cm H_2O), the extent of alveolar infiltrates on chest radiography, and the corrected expired volume per minute (≥ 10 L/min).

Functional Recovery in ARDS Survivors While it is common for patients with ARDS to experience prolonged respiratory failure and remain dependent on mechanical ventilation for survival, it is a testament to the resolving powers of the lung that the majority of patients recover nearly normal lung function. Patients usually recover maximal lung function within 6 months. One year after endotracheal extubation, more than one-third of ARDS survivors have normal spirometry values and diffusion capacity. Most of the remaining patients have only mild abnormalities in pulmonary function. Unlike mortality risk, recovery of lung function is strongly associated with the extent of lung injury in early ARDS. Low static respiratory compliance, high levels of required PEEP, longer durations of mechanical ventilation, and high lung injury scores are all associated with less recovery of pulmonary function. Of note, when physical function is assessed 5 years after ARDS, exercise limitation and decreased physical quality of life are often documented despite normal or nearly normal pulmonary function. When caring for ARDS survivors, it is important to be aware of the potential for a substantial burden of psychological problems in patients and family caregivers, including significant rates of depression and posttraumatic stress disorder.

WEBSITES

ARDS Support Center for patient-oriented education: www.ards.org
NHLBI ARDS Clinical Trials information: www.ardsnet.org
ARDS Foundation: www.ardsusa.org

ACKNOWLEDGMENT

The authors acknowledge the contribution to this chapter by the previous author, Dr. Steven D. Shapiro.

INDICATIONS

The most common reasons for instituting mechanical ventilation are acute respiratory failure with hypoxemia (acute respiratory distress syndrome, heart failure with pulmonary edema, pneumonia, sepsis, complications of surgery and trauma), which accounts for ~65% of all ventilated cases, and hypercarbic ventilatory failure—e.g., due to coma (15%), exacerbations of chronic obstructive pulmonary disease (COPD; 13%), and neuromuscular diseases (5%). The primary objectives of mechanical ventilation are to decrease the work of breathing, thus avoiding respiratory muscle fatigue, and to reverse life-threatening hypoxemia and progressive respiratory acidosis.

In some cases, mechanical ventilation is used as an adjunct to other forms of therapy. For example, it is used to reduce cerebral blood flow in patients with increased intracranial pressure. Mechanical ventilation also is used frequently in conjunction with endotracheal intubation for airway protection to prevent aspiration of gastric contents in otherwise unstable patients during gastric lavage for suspected drug overdose or during gastrointestinal endoscopy. In critically ill patients, intubation and mechanical ventilation may be indicated before the performance of essential diagnostic or therapeutic studies if it appears that respiratory failure may occur during those maneuvers.

TYPES OF MECHANICAL VENTILATION

There are two basic methods of mechanical ventilation: noninvasive ventilation (NIV) and invasive (or conventional mechanical) ventilation (MV).

Noninvasive Ventilation NIV has gained acceptance because it is effective in certain conditions, such as acute or chronic respiratory failure, and is associated with fewer complications—namely, pneumonia and tracheolaryngeal trauma. NIV usually is provided with a tight-fitting face mask or nasal mask similar to the masks traditionally used for treatment of sleep apnea. NIV has proved highly effective in patients with respiratory failure arising from acute exacerbations of chronic obstructive pulmonary disease. It is most frequently implemented as bilevel positive airway pressure ventilation or pressure-support ventilation. Both modes, which apply a preset positive pressure during inspiration and a lower pressure during expiration at the mask, are well tolerated by a conscious patient and optimize patient-ventilator synchrony. The major limitation to the widespread application of NIV has been patient intolerance: the tight-fitting mask required for NIV can cause both physical and psychological discomfort. In addition, NIV has had limited success in patients with acute hypoxemic respiratory failure, for whom endotracheal intubation and conventional MV remain the ventilatory method of choice.

The most important group of patients who benefit from a trial of NIV are those with exacerbations of COPD and respiratory acidosis ($pH < 7.35$). Experience from several randomized trials has shown that, in patients with ventilatory failure characterized by blood pH levels between 7.25 and 7.35, NIV is associated with low failure rates (15–20%) and good outcomes (as judged by intubation rate, length of stay in intensive care, and—in some series—mortality rates). In more severely ill patients with a blood pH < 7.25 , the rate of NIV failure is inversely related to the severity of respiratory acidosis, with higher failure rates as the pH decreases. In patients with milder acidosis ($pH > 7.35$), NIV is not better than conventional treatment that includes controlled oxygen delivery and pharmacotherapy for exacerbations of COPD (systemic glucocorticoids, bronchodilators, and, if needed, antibiotics).

Despite its benign outcomes, NIV is not useful in the majority of cases of respiratory failure and is contraindicated in patients with the conditions listed in [Table 323-1](#). NIV can delay lifesaving ventilatory support in those cases and, in fact, can actually result in aspiration or hypoventilation. Once NIV is initiated, patients should be monitored; a reduction in respiratory frequency and a decrease in the use of accessory muscles (scalene, sternomastoid, and intercostals) are good clinical indicators of adequate therapeutic benefit. Arterial blood gases should be determined at least within hours of the initiation of therapy to ensure that NIV is having the desired effect. Lack of benefit within

323 Mechanical Ventilatory Support

Bartolome R. Celli

MECHANICAL VENTILATORY SUPPORT

Mechanical ventilation is used to assist or replace spontaneous breathing. It is implemented with special devices that can support ventilatory function and improve oxygenation through the application of high-oxygen-content gas and positive pressure. The primary indication for initiation of mechanical ventilation is respiratory failure, of which there are two basic types: (1) *hypoxemic*, which is present when arterial O_2 saturation (SaO_2) $< 90\%$ occurs despite an increased inspired O_2 fraction and usually results from ventilation-perfusion mismatch or shunt; and (2) *hypercarbic*, which is characterized by elevated arterial carbon dioxide partial pressure (PCO_2) values (usually > 50 mmHg) resulting from conditions that decrease minute ventilation or increase physiologic dead space such that alveolar ventilation is inadequate to meet metabolic demands. When respiratory failure is chronic, neither of the two types is obligatorily treated with mechanical ventilation, but when it is acute, mechanical ventilation may be lifesaving.

TABLE 323-1 CONTRAINDICATIONS FOR NONINVASIVE VENTILATION

Cardiac or respiratory arrest
Severe encephalopathy
Severe gastrointestinal bleed
Hemodynamic instability
Unstable angina and myocardial infarction
Facial surgery or trauma
Upper airway obstruction
High-risk aspiration and/or inability to protect airways
Inability to clear secretions

that time frame should alert the physician to the possible need for conventional MV.

Conventional Mechanical Ventilation Conventional MV is implemented once a cuffed tube is inserted into the trachea to allow conditioned gas (warmed, oxygenated, and humidified) to be delivered to the airways and lungs at pressures above atmospheric pressure. Care should be taken during intubation to avoid brain-damaging hypoxia. In most cases, the administration of mild sedation may facilitate the procedure. Opiates and benzodiazepines are good choices but can have a deleterious effect on hemodynamics in patients with depressed cardiac function or low systemic vascular resistance. Morphine can promote histamine release from tissue mast cells and may worsen bronchospasm in patients with asthma; fentanyl, sufentanil, and alfentanil are acceptable alternatives. Ketamine may increase systemic arterial pressure and has been associated with hallucinatory responses. The shorter-acting agents etomidate and propofol have been used for both induction and maintenance of anesthesia in ventilated patients because they have fewer adverse hemodynamic effects, but both are significantly more expensive than older agents. Great care must be taken to avoid the use of neuromuscular paralysis during intubation of patients with renal failure, tumor lysis syndrome, crush injuries, medical conditions associated with elevated serum potassium levels, and muscular dystrophy syndromes; in particular, the use of agents whose mechanism of action includes depolarization at the neuromuscular junction, such as succinylcholine chloride, must be avoided.

PRINCIPLES OF MECHANICAL VENTILATION

Once the patient has been intubated, the basic goals of MV are to optimize oxygenation while avoiding ventilator-induced lung injury due to overstretch and collapse/re-recruitment. This concept, known as the “protective ventilatory strategy” (see below and Fig. 323-1) is supported by evidence linking high airway pressures and volumes and overstretching of the lung as well as collapse/re-recruitment to poor clinical outcomes (barotrauma and volume trauma). Although normalization of pH through elimination of CO₂ is desirable, the risk of lung damage associated with the large volume and high pressures needed to achieve this goal has led to the acceptance of permissive hypercapnia. This condition is well tolerated when care is taken to avoid excess acidosis by pH buffering.

MODES OF VENTILATION

Mode refers to the manner in which ventilator breaths are triggered, cycled, and limited. The *trigger*, either an inspiratory effort or a time-based signal, defines what the ventilator senses to initiate an assisted breath. *Cycle* refers to the factors that determine the end of inspiration. For example, in volume-cycled ventilation, inspiration ends when a specific tidal volume is delivered. Other types of cycling include pressure cycling and time cycling. The *limiting factors* are operator-specified values, such as airway pressure, that are monitored by transducers internal to the ventilator circuit throughout the respiratory cycle; if the specified values are exceeded, inspiratory flow is terminated, and the ventilator circuit is vented to atmospheric pressure or the specified pressure at the end of expiration (positive end-expiratory pressure, or PEEP). Most patients are ventilated with assist-control ventilation,

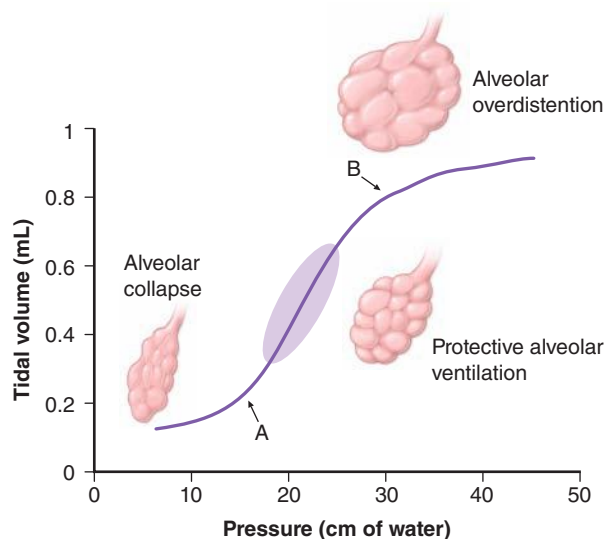


FIGURE 323-1 Hypothetical pressure-volume curve of the lung in a patient undergoing mechanical ventilation. Alveoli tend to close if the distending pressure falls below the lower inflection point (A), whereas they overstretch if the pressure within them is higher than that of the upper inflection point (B). Collapse and opening of ventilated alveoli are associated with poor outcomes in patients with acute respiratory failure. Protective ventilation (purple shaded area), using a lower tidal volume (6 mL/kg of ideal body weight) and maintaining positive end-expiratory pressure to prevent overstretching and collapse/opening of alveoli, has resulted in improved survival rates among patients receiving mechanical ventilatory support.

intermittent mandatory ventilation, or pressure-support ventilation, with the latter two modes often used simultaneously (Table 323-2).

Assist-Control Ventilation (ACMV) ACMV is the most widely used mode of ventilation. In this mode, an inspiratory cycle is initiated either by the patient’s inspiratory effort or, if none is detected within a specified time window, by a timer signal within the ventilator. Every breath delivered, whether patient- or timer-triggered, consists of the operator-specified tidal volume. Ventilatory rate is determined either by the patient or by the operator-specified backup rate, whichever is of higher frequency. ACMV is commonly used for initiation of mechanical ventilation because it ensures a backup minute ventilation in the absence of an intact respiratory drive and allows for synchronization of the ventilator cycle with the patient’s inspiratory effort.

Problems can arise when ACMV is used in patients with tachypnea due to nonrespiratory or nonmetabolic factors, such as anxiety, pain, and airway irritation. Respiratory alkalemia may develop and trigger myoclonus or seizures. Dynamic hyperinflation leading to increased intrathoracic pressures (so-called auto-PEEP) may occur if the patient’s respiratory mechanics are such that inadequate time is available for complete exhalation between inspiratory cycles. Auto-PEEP can limit venous return, decrease cardiac output, and increase airway pressures, predisposing to barotrauma.

Intermittent Mandatory Ventilation (IMV) With this mode, the operator sets the number of mandatory breaths of fixed volume to be delivered by the ventilator; between those breaths, the patient can breathe spontaneously. In the most frequently used synchronized mode (SIMV), mandatory breaths are delivered in synchrony with the patient’s inspiratory efforts at a frequency determined by the operator. If the patient fails to initiate a breath, the ventilator delivers a fixed-tidal-volume breath and resets the internal timer for the next inspiratory cycle. SIMV differs from ACMV in that only a preset number of breaths are ventilator-assisted.

SIMV allows patients with an intact respiratory drive to exercise inspiratory muscles between assisted breaths; thus it is useful for both supporting and weaning intubated patients. SIMV may be difficult to

TABLE 323-2 CHARACTERISTICS OF THE MOST COMMONLY USED FORMS OF MECHANICAL VENTILATION

Ventilatory Mode	Variables Set by User (Independent)	Variables Monitored by User (Dependent)	Trigger Cycle Limit	Advantages	Disadvantages
ACMV (assist-control ventilation)	Tidal volume Ventilator rate FiO ₂ PEEP level Pressure limit	Peak, mean, and plateau airway pressures VE ABG I/E ratio	Patient effort Timer Pressure limit	Patient control Guaranteed ventilation	Potential hyperventilation Barotrauma and volume trauma Every effective breath generates a ventilator volume
IMV (intermittent mandatory ventilation)	Tidal volume Mandatory ventilator rate FiO ₂ PEEP level Pressure limit Spontaneous breaths between assisted breaths	Peak, mean, and plateau airway pressures VE ABG I/E ratio	Patient effort Timer Pressure limit	Patient control Comfort from spontaneous breaths Guaranteed ventilation	Potential dysynchrony Potential hypoventilation
PSV (pressure-support ventilation)	Inspiratory pressure level FiO ₂ PEEP Pressure limit	Tidal volume Respiratory rate VE ABG	Pressure limit Inspiratory flow	Patient control Comfort Assures synchrony	No timer backup Potential hypoventilation
NIV (noninvasive ventilation)	Inspiratory and expiratory pressure level FiO ₂	Tidal volume Respiratory rate VE ABG	Pressure limit Inspiratory flow	Patient control	Mask interface may cause discomfort and facial bruising Leaks are common Hypoventilation

Abbreviations: ABG, arterial blood gases; FiO₂, fraction of inspired oxygen; PEEP, positive end-expiratory pressure; I/E, inspiratory to expiratory time ratio; VE, minute ventilation.

use in patients with tachypnea because they may attempt to exhale during the ventilator-programmed inspiratory cycle. Consequently, the airway pressure may exceed the inspiratory pressure limit, the ventilator-assisted breath will be aborted, and minute volume may drop below that programmed by the operator. In this setting, if the tachypnea represents a response to respiratory or metabolic acidosis, a change in ACMV will increase minute ventilation and help normalize the pH while the underlying process is further evaluated and treated.

Pressure-Support Ventilation (PSV) This form of ventilation is patient-triggered, flow-cycled, and pressure-limited. It provides graded assistance and differs from the other two modes in that the operator sets the pressure level (rather than the volume) to augment every spontaneous respiratory effort. The level of pressure is adjusted by observing the patient's respiratory frequency. During PSV, the inspiration is terminated when inspiratory airflow falls below a certain level; in most ventilators, this flow rate cannot be adjusted by the operator. With PSV, patients receive ventilator assistance only when the ventilator detects an inspiratory effort. PSV is often used in combination with SIMV to ensure volume-cycled backup for patients whose respiratory drive is depressed. PSV is well tolerated by most patients who are being weaned from MV; PSV parameters can be set to provide full ventilatory support and can be withdrawn to load the respiratory muscles gradually.

Other Modes of Ventilation There are other modes of ventilation, each with its own acronym and each with specific modifications of the manner and duration in which pressure is applied to the airway and lungs and of the interaction between the mechanical assistance provided by the ventilator and the patient's respiratory effort. Although their use in acute respiratory failure is limited, the following modes have been used with varying levels of enthusiasm and adoption.

PRESSURE-CONTROL VENTILATION (PCV) This form of ventilation is time-triggered, time-cycled, and pressure-limited. A specified pressure is imposed at the airway opening throughout inspiration. Since the inspiratory pressure is specified by the operator, tidal volume and

inspiratory flow rate are *dependent*, rather than *independent*, variables and are not operator-specified. PCV is the preferred mode of ventilation for patients in whom it is desirable to regulate peak airway pressures, such as those with preexisting barotrauma, and for post-thoracic surgery patients, in whom the shear forces across a fresh suture line should be limited. When PCV is used, minute ventilation is altered through changes in rate or in the pressure-control value, with consequent changes in tidal volume.

INVERSE-RATIO VENTILATION (IRV) This mode is a variant of PCV that incorporates the use of a prolonged inspiratory time with the appropriate shortening of the expiratory time. IRV has been used in patients with severe hypoxemic respiratory failure. This approach increases mean distending pressures without increasing peak airway pressures. It is thought to work in conjunction with PEEP to open collapsed alveoli and improve oxygenation. However, no clinical-trial data have shown that IRV improves outcomes.

CONTINUOUS POSITIVE AIRWAY PRESSURE (CPAP) CPAP is not a true support mode of ventilation because all ventilation occurs through the patient's spontaneous efforts. The ventilator provides fresh gas to the breathing circuit with each inspiration and sets the circuit to a constant, operator-specified pressure. CPAP is used to assess extubation potential in patients who have been effectively weaned and who require little ventilatory support and in patients with intact respiratory system function who require an endotracheal tube for airway protection.

Nonconventional Ventilatory Strategies Several nonconventional strategies have been evaluated for their ability to improve oxygenation and reduce mortality rates in patients with advanced hypoxemic respiratory failure. These strategies include high-frequency oscillatory ventilation (HFOV), airway pressure release ventilation (APRV), extracorporeal membrane oxygenation (ECMO), and partial liquid ventilation (PLV) using perfluorocarbons. Although case reports and small uncontrolled cohort studies have shown benefit, randomized controlled trials have failed to demonstrate consistent improvements in outcome with most

of these strategies. A recent randomized trial of ECMO documented positive outcomes, but the technique remains controversial because older studies failed to document positive results. Currently, these approaches should be thought of as “salvage” techniques and considered for patients with hypoxemia refractory to conventional therapy. Prone positioning of patients with refractory hypoxemia has also been explored because, in theory, lying prone should improve ventilation-perfusion matching. Several randomized trials in patients with acute lung injury did not demonstrate a survival advantage with prone positioning despite demonstration of a transient physiologic benefit. The administration of nitric oxide gas, which has bronchodilator and pulmonary vasodilator effects when delivered through the airways and improves arterial oxygenation in many patients with advanced hypoxemic respiratory failure, also failed to improve outcomes in these patients with acute lung injury.

The design of new ventilator modes reflect attempts to improve patient-ventilator synchrony—a major practical issue during MV—by allowing patients to trigger the ventilator with their own effort while also incorporating flow algorithms that terminate the cycles once certain preset criteria are reached; this approach has greatly improved patient comfort. New modes of ventilation that synchronize not only the timing but also the levels of assistance to match the patient’s effort have been developed. Proportional assist ventilation (PAV) and neurally adjusted ventilatory-assist ventilation (NAV) are two modes that are designed to deliver assisted breaths through algorithms incorporating not only pressure, volume, and time but also overall respiratory resistance as well as compliance (in the case of PAV) and neural activation of the diaphragm (in the case of NAV). Although these modes enhance patient-ventilator synchrony, their practical use in the everyday management of patients undergoing MV needs further study.

PROTECTIVE VENTILATORY STRATEGY

Whichever mode of MV is used in acute respiratory failure, the evidence from several important controlled trials indicates that a protective ventilation approach guided by the following principles (and summarized in Fig. 323-1) is safe and offers the best chance of a good outcome: (1) Set a target tidal volume close to 6 mL/kg of ideal body weight. (2) Prevent plateau pressure (static pressure in the airway at the end of inspiration) exceeding 30 cm H₂O. (3) Use the lowest possible fraction of inspired oxygen (F_{IO₂}) to keep the SaO₂ at ≥90%. (4) Adjust the PEEP to maintain alveolar patency while preventing overdistention and closure/reopening. With the application of these techniques, the mortality rate among patients with acute hypoxemic respiratory failure has decreased to ~30% from close to 50% a decade ago.

PATIENT MANAGEMENT

Once the patient has been stabilized with respect to gas exchange, definitive therapy for the underlying process responsible for respiratory failure is initiated. Subsequent modifications in ventilator therapy must be provided in parallel with changes in the patient’s clinical status. As improvement in respiratory function is noted, the first priority is to reduce the level of mechanical ventilatory support. Patients on full ventilatory support should be monitored frequently, with the goal of switching to a mode that allows for weaning as soon as possible. Protocols and guidelines that can be applied by paramedical personnel when physicians are not readily available have proved to be of value in shortening ventilator and intensive care unit (ICU) time, with very good outcomes. Patients whose condition continues to deteriorate after ventilatory support is initiated may require increased O₂, PEEP, or one of the alternative modes of ventilation.

GENERAL SUPPORT DURING VENTILATION

Patients for whom mechanical ventilation has been initiated usually require sedation and analgesia to maintain an acceptable level of comfort. Often, this treatment consists of a combination of a benzodiazepine and an opiate administered intravenously. Medications commonly used for this purpose include lorazepam, midazolam, diazepam, morphine, and fentanyl. Oversedation must be avoided in the ICU because most (but not all) studies show that daily interruption of sedation in

patients with improved ventilatory status results in a shorter time on the ventilator and a shorter ICU stay.

Immobilized patients receiving mechanical ventilatory support are at risk for deep venous thrombosis and decubitus ulcers. Venous thrombosis should be prevented with the use of subcutaneous heparin and/or pneumatic compression boots. Fractionated low-molecular-weight heparin appears to be equally effective for this purpose. To help prevent decubitus ulcers, frequent changes in body position and the use of soft mattress overlays and air mattresses are employed. Prophylaxis against diffuse gastrointestinal mucosal injury is indicated for patients undergoing MV. Histamine-receptor (H₂-receptor) antagonists, antacids, and cytoprotective agents such as sucralfate have all been used for this purpose and appear to be effective. Nutritional support by enteral feeding through either a nasogastric or an orogastric tube should be initiated and maintained whenever possible. Delayed gastric emptying is common in critically ill patients taking sedative medications but often responds to promotility agents such as metoclopramide. Parenteral nutrition is an alternative to enteral nutrition in patients with severe gastrointestinal pathology who need prolonged MV.

COMPLICATIONS OF MECHANICAL VENTILATION

Endotracheal intubation and mechanical ventilation have direct and indirect effects on the lung and upper airways, the cardiovascular system, and the gastrointestinal system. Pulmonary complications include barotrauma, nosocomial pneumonia, oxygen toxicity, tracheal stenosis, and deconditioning of respiratory muscles. Barotrauma and volutrauma overdistend and disrupt lung tissue; may be clinically manifest by interstitial emphysema, pneumomediastinum, subcutaneous emphysema, or pneumothorax; and can result in the liberation of cytokines from overdistended tissues, further promoting tissue injury. Clinically significant pneumothorax requires tube thoracostomy. Intubated patients are at high risk for ventilator-associated pneumonia as a result of aspiration from the upper airways through small leaks around the endotracheal tube cuff; the most common organisms responsible for this condition are *Pseudomonas aeruginosa*, enteric gram-negative rods, and *Staphylococcus aureus*. Given the high associated mortality rates, early initiation of empirical antibiotics directed against likely pathogens is recommended. Hypotension resulting from elevated intrathoracic pressures with decreased venous return is almost always responsive to intravascular volume repletion. In patients who are judged to have respiratory failure on the basis of alveolar edema but in whom the cardiac or pulmonary origin of the edema is unclear, hemodynamic monitoring with a pulmonary arterial catheter may be of value in helping to clarify the cause of the edema. Gastrointestinal effects of positive-pressure ventilation include stress ulceration and mild to moderate cholestasis.

WEANING FROM MECHANICAL VENTILATION

The Decision to Wean It is important to consider discontinuation of mechanical ventilation once the underlying respiratory disease begins to reverse. Although the predictive capacities of multiple clinical and physiologic variables have been explored, the consensus from a ventilatory weaning task force cites the following conditions as indicating amenability to weaning: (1) Lung injury is stable or resolving. (2) Gas exchange is adequate, with low PEEP/F_{IO₂} (<8 cmH₂O) and F_{IO₂} (<0.5). (3) Hemodynamic variables are stable, and the patient is no longer receiving vasopressors. (4) The patient is capable of initiating spontaneous breaths. A “wean screen” based on these variables should be done at least daily. If the patient is deemed capable of beginning to wean, the recommendation is to perform a spontaneous breathing trial (SBT), whose value is supported by several randomized trials (Fig. 323-2). The SBT involves an integrated patient assessment during spontaneous breathing with little or no ventilatory support. The SBT is usually implemented with a T-piece using 1–5 cmH₂O CPAP with 5–7 cmH₂O or PSV from the ventilator to offset resistance from the endotracheal tube. Once it is determined that the patient can breathe spontaneously, a decision must be made about the removal of the artificial airway, which should be undertaken only when it is concluded that the patient has the ability to protect the airway, is able to cough and clear

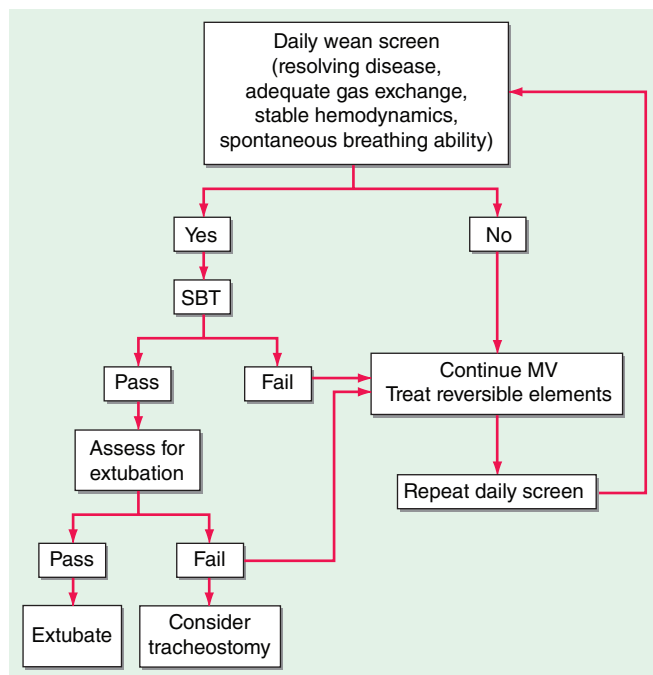


FIGURE 323-2 Flow chart to guide the daily approach to management of patients being considered for weaning off mechanical ventilation (MV). If attempts at extubation fail, a tracheostomy should be considered. SBT, spontaneous breathing trial.

secretions, and is alert enough to follow commands. In addition, other factors must be taken into account, such as the possible difficulty of replacing the tube if that maneuver is required. If upper airway difficulty is suspected, an evaluation using a “cuff-leak” test (assessing the

presence of air movement around a deflated endotracheal tube cuff) is supported by some internists. Despite all precautions, ~10–15% of extubated patients require reintubation. Several studies suggest that NIV can be used to obviate reintubation, particularly in patients with ventilatory failure secondary to COPD exacerbation; in this setting, earlier extubation with the use of prophylactic NIV has yielded good results. The use of NIV to facilitate weaning in respiratory failure of other etiologies is not currently indicated.

Prolonged Mechanical Ventilation and Tracheostomy From 5% to 13% of patients undergoing MV will go on to require prolonged MV (>21 days). In these instances, critical care personnel must decide whether and when to perform a tracheostomy. This decision is individualized and is based on the risk and benefits of tracheostomy and prolonged intubation as well as the patient’s preferences and expected outcomes. A tracheostomy is thought to be more comfortable, to require less sedation, and to provide a more secure airway and may also reduce weaning time. However, tracheostomy carries the risk of complications, which occur in 5–40% of these procedures and include bleeding, cardiopulmonary arrest, hypoxia, structural damage, pneumothorax, pneumomediastinum, and wound infection. In patients with long-term tracheostomy, complex complications include tracheal stenosis, granulation, and erosion of the innominate artery. In general, if a patient needs MV for more than 10–14 days, a tracheostomy, planned under optimal conditions, is indicated. Whether it is completed at the bedside or as an operative procedure depends on local resources and experience. Some 5–10% of patients are deemed unable to wean in the ICU. These patients may benefit from transfer to special units where a multidisciplinary approach, including nutrition optimization, physical therapy with rehabilitation, and slower weaning methods (including SIMV with PSV), results in successful weaning rates of up to 30%. Unfortunately, close to 2% of ventilated patients may ultimately become dependent on ventilatory support to maintain life. Most of these patients remain in chronic care institutions, although some with strong social, economic, and family support may live a relatively fulfilling life with at-home ventilation.

SECTION 2 SHOCK AND CARDIAC ARREST

324 Approach to the Patient with Shock

Ronald V. Maier

Shock is the clinical syndrome that results from inadequate tissue perfusion. Irrespective of cause, the hypoperfusion-induced imbalance between the delivery of and requirements for oxygen and substrate leads to cellular dysfunction. The cellular injury created by the inadequate delivery of oxygen and substrates also induces the production and release of damage-associated molecular patterns (DAMPs or “danger signals”) and inflammatory mediators that further compromise perfusion through functional and structural changes within the microvasculature. This leads to a vicious cycle in which impaired perfusion is responsible for cellular injury that causes maldistribution of blood flow, further compromising cellular perfusion; the latter ultimately causes multiple organ failure (MOF) and, if the process is not interrupted, leads to death. The clinical manifestations of shock are also the result, in part, of autonomic neuroendocrine responses to hypoperfusion as well as the breakdown in organ function induced by severe cellular dysfunction (Fig. 324-1).

When very severe and/or persistent, inadequate oxygen delivery leads to irreversible cell injury, only rapid restoration of oxygen

delivery can reverse the progression of the shock state. The fundamental approach to management, therefore, is to recognize overt and impending shock in a timely fashion and to intervene emergently to restore perfusion. Doing so often requires the expansion or reexpansion of intravascular blood volume. Control of any inciting pathologic process (e.g., continued hemorrhage, impairment of cardiac function, or infection) must occur simultaneously.

Clinical shock is usually accompanied by hypotension (i.e., a mean arterial pressure [MAP] <60 mmHg in previously normotensive persons). Multiple classification schemes have been developed in an attempt to synthesize the seemingly dissimilar processes leading to shock. Strict adherence to a classification scheme may be difficult from a clinical standpoint because of the frequent combination of two or more causes of shock in any individual patient, but the classification shown in Table 324-1 provides a useful reference point from which to discuss and further delineate the underlying processes.

PATHOGENESIS AND ORGAN RESPONSE

MICROCIRCULATION

Normally when cardiac output falls, systemic vascular resistance rises to maintain a level of systemic pressure that is adequate for perfusion of the heart and brain at the expense of other tissues such as muscle, skin, and especially the gastrointestinal (GI) tract. Systemic vascular resistance is determined primarily by the luminal diameter of arterioles. The metabolic rates of the heart and brain are high, and their stores of energy substrate are low. These organs are critically dependent on a

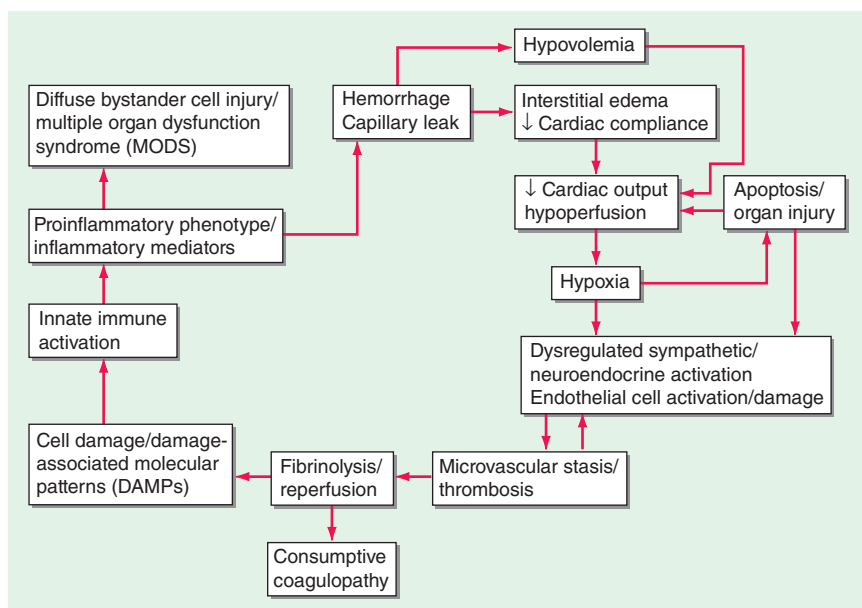


FIGURE 324-1 Shock-induced vicious cycle.

continuous supply of oxygen and nutrients, and neither tolerates severe ischemia for more than brief periods (minutes). Autoregulation (i.e., the maintenance of blood flow over a wide range of perfusion pressures) is critical in sustaining cerebral and coronary perfusion despite significant hypotension. However, when MAP drops to ≤ 60 mmHg, blood flow to these organs falls, and their function deteriorates.

Arteriolar vascular smooth muscle has both α - and β -adrenergic receptors. The α_1 receptors mediate vasoconstriction, while the β_2 receptors mediate vasodilation. Efferent sympathetic fibers release norepinephrine, which acts primarily on α_1 receptors as one of the most fundamental compensatory responses to reduced perfusion pressure. Other constrictor substances that are increased in most forms of shock include angiotensin II, vasopressin, endothelin 1, and thromboxane A_2 . Both norepinephrine and epinephrine are released by the adrenal medulla, and the concentrations of these catecholamines in the bloodstream rise. Circulating vasodilators in shock include prostacyclin (prostaglandin [PG] I_2), nitric oxide (NO), and, importantly, products of local metabolism such as adenosine that match flow to the tissue's metabolic needs. The balance between these various vasoconstrictors and vasodilators influences the microcirculation and determines local perfusion.

Transport to cells depends on microcirculatory flow; capillary permeability; the diffusion of oxygen, carbon dioxide, nutrients, and products of metabolism through the interstitium; and the exchange of these products across cell membranes. Impairment of the microcirculation that is central to the pathophysiologic responses in the late stages of all forms of shock results in the derangement of cellular metabolism that is ultimately responsible for organ failure.

The endogenous response to mild or moderate hypovolemia is an attempt at restitution of intravascular volume through alterations in hydrostatic pressure and osmolarity. Constriction of arterioles leads to reductions in both the capillary hydrostatic pressure and the number of capillary beds perfused, thereby limiting the capillary surface area across which filtration occurs. When filtration is reduced while intravascular oncotic pressure remains constant or rises, there is net

reabsorption of fluid into the vascular bed, in accord with Starling's law of capillary interstitial liquid exchange. Metabolic changes (including hyperglycemia and elevations in the products of glycolysis, lipolysis, and proteolysis) raise extracellular osmolarity, leading to an osmotic gradient that increases interstitial and intravascular volume at the expense of intracellular volume.

CELLULAR RESPONSES

Interstitial transport of nutrients is impaired in shock, leading to a decline in intracellular high-energy phosphate stores. Mitochondrial dysfunction and uncoupling of oxidative phosphorylation are the most likely causes for decreased amounts of adenosine triphosphate (ATP). As a consequence, there is an accumulation of hydrogen ions, lactate, reactive oxygen species, and other products of anaerobic metabolism. As shock progresses, these vasodilator metabolites override vasomotor tone, causing further hypotension and hypoperfusion. Dysfunction of cell membranes is thought to represent a common end-stage pathophysiologic pathway in the various forms of shock. Normal cellular transmembrane potential falls, and there is an associated increase in

intracellular sodium and water, leading to cell swelling that interferes further with microvascular perfusion. In a preterminal event, homeostasis of calcium via membrane channels is lost with flooding of calcium into the cytosol and concomitant extracellular hypocalcemia. There is also evidence for a widespread but selective apoptotic (programmed cell death) loss of cells, contributing to organ and immune failure.

NEUROENDOCRINE RESPONSE

Hypovolemia, hypotension, and hypoxia are sensed by baroreceptors and chemoreceptors that contribute to an autonomic response that attempts to restore blood volume, maintain central perfusion, and mobilize metabolic substrates. Hypotension disinhibits the vasomotor center, resulting in increased adrenergic output and reduced vagal activity. Release of norepinephrine from adrenergic neurons induces significant peripheral and splanchnic vasoconstriction, a major contributor to the maintenance of central organ perfusion, while reduced vagal activity increases the heart rate and cardiac output. Loss of vagal activity is also recognized to upregulate the innate immune inflammatory response. The effects of circulating epinephrine released by the adrenal medulla in shock are largely metabolic, causing increased glycogenolysis and gluconeogenesis and reduced pancreatic insulin release. However, epinephrine also inhibits production and release of inflammatory mediators through stimulation of β -adrenergic receptors on innate immune cells.

Severe pain or other stresses cause the hypothalamic release of adrenocorticotrophic hormone (ACTH). This stimulates cortisol secretion that contributes to decreased peripheral uptake of glucose and amino acids, enhances lipolysis, and increases gluconeogenesis. Increased pancreatic secretion of glucagon during stress accelerates hepatic gluconeogenesis and further elevates blood glucose concentration. These hormonal actions act synergistically to increase blood glucose for both selective tissue metabolism and the maintenance of blood volume. Many critically ill patients have recently been shown to exhibit low plasma cortisol levels and an impaired response to ACTH stimulation, which is linked to a decrease in survival. The importance of the cortisol response to stress is illustrated by the profound circulatory collapse that occurs in patients with adrenocortical insufficiency (**Chap. 406**).

Renin release is increased in response to adrenergic discharge and reduced perfusion of the juxtaglomerular apparatus in the kidney. Renin induces the formation of angiotensin I that is then converted to angiotensin II by the angiotensin converting enzyme; angiotensin II is an extremely potent vasoconstrictor and stimulator of aldosterone release by the adrenal cortex and of vasopressin by the posterior pituitary. Aldosterone contributes to the maintenance of intravascular

TABLE 324-1 CLASSIFICATION OF SHOCK

Hypovolemic	Septic
Traumatic	Hyperdynamic (early)
Cardiogenic	Hypodynamic (late)
Intrinsic	Neurogenic
Compressive	Hypoadrenal

1746 volume by enhancing renal tubular reabsorption of sodium, resulting in the excretion of a low-volume, concentrated, sodium-free urine. Vasopressin has a direct action on vascular smooth muscle, contributing to vasoconstriction, and acts on the distal renal tubules to enhance water reabsorption.

CARDIOVASCULAR RESPONSE

Three variables—ventricular filling (preload), the resistance to ventricular ejection (afterload), and myocardial contractility—are paramount in controlling stroke volume (**Chap. 265e**). Cardiac output, the major determinant of tissue perfusion, is the product of stroke volume and heart rate. Hypovolemia leads to decreased ventricular preload that, in turn, reduces the stroke volume. An increase in heart rate is a useful but limited compensatory mechanism to maintain cardiac output. A shock-induced reduction in myocardial compliance is frequent, reducing ventricular end-diastolic volume and, hence, stroke volume at any given ventricular filling pressure. Restoration of intravascular volume may return stroke volume to normal but only at elevated filling pressures. Increased filling pressures stimulate release of brain natriuretic peptide (BNP) to secrete sodium and volume to relieve the pressure on the heart. Levels of BNP correlate with outcome following severe stress. In addition, sepsis, ischemia, myocardial infarction (MI), severe tissue trauma, hypothermia, general anesthesia, prolonged hypotension, and acidemia may all also impair myocardial contractility and reduce the stroke volume at any given ventricular end-diastolic volume. The resistance to ventricular ejection is significantly influenced by the systemic vascular resistance, which is elevated in most forms of shock. However, resistance is decreased in the early hyperdynamic stage of septic shock or neurogenic shock (**Chap. 325**), thereby initially allowing the cardiac output to be maintained or elevated.

The venous system contains nearly two-thirds of the total circulating blood volume, most in the small veins, and serves as a dynamic reservoir for autoinfusion of blood. Active venoconstriction as a consequence of α -adrenergic activity is an important compensatory mechanism for the maintenance of venous return and, therefore, of ventricular filling during shock. By contrast, venous dilation, as occurs in neurogenic shock, reduces ventricular filling and hence stroke volume and potentially cardiac output.

PULMONARY RESPONSE

The response of the pulmonary vascular bed to shock parallels that of the systemic vascular bed, and the relative increase in pulmonary vascular resistance, particularly in septic shock, may exceed that of the systemic vascular resistance, leading to right heart failure. Shock-induced tachypnea reduces tidal volume and increases both dead space and minute ventilation. Relative hypoxia and the subsequent tachypnea induce a respiratory alkalosis. Recumbency and involuntary restriction of ventilation secondary to pain reduce functional residual capacity and may lead to atelectasis. Shock and, in particular, resuscitation-induced reactive oxygen species (oxidant radical) generation are recognized as major causes of acute lung injury and subsequent acute respiratory distress syndrome (ARDS; **Chap. 322**). These disorders are characterized by noncardiogenic pulmonary edema secondary to diffuse pulmonary capillary endothelial and alveolar epithelial injury, hypoxemia, and bilateral diffuse pulmonary infiltrates. Hypoxemia results from perfusion of underventilated and nonventilated alveoli. Loss of surfactant and lung volume in combination with increased interstitial and alveolar edema reduces lung compliance. The work of breathing and the oxygen requirements of respiratory muscles increase.

RENAL RESPONSE

Acute kidney injury (**Chap. 334**), a serious complication of shock and hypoperfusion, occurs less frequently than heretofore because of early aggressive volume repletion. Acute tubular necrosis is now more frequently seen as a result of the interactions of shock, sepsis, the administration of nephrotoxic agents (such as aminoglycosides and angiographic contrast media), and rhabdomyolysis; the latter may be particularly severe in skeletal muscle trauma. The physiologic response

of the kidney to hypoperfusion is to conserve salt and water. In addition to decreased renal blood flow, increased afferent arteriolar resistance accounts for diminished glomerular filtration rate (GFR) that, together with increased aldosterone and vasopressin, is responsible for reduced urine formation. Toxic injury causes necrosis of tubular epithelium and tubular obstruction by cellular debris with back leak of filtrate. The depletion of renal ATP stores that occurs with prolonged renal hypoperfusion contributes to subsequent impairment of renal function.

METABOLIC DERANGEMENTS

During shock, there is disruption of the normal cycles of carbohydrate, lipid, and protein metabolism. Through the citric acid cycle, alanine in conjunction with lactate, which is converted from pyruvate in the periphery in the presence of oxygen deprivation, enhances the hepatic production of glucose. With reduced availability of oxygen, the breakdown of glucose to pyruvate, and ultimately lactate, represents an inefficient cycling of substrate with minimal net energy production. An elevated plasma lactate/pyruvate ratio is preferable to lactate alone as a measure of anaerobic metabolism and reflects inadequate tissue perfusion. Decreased clearance of exogenous triglycerides coupled with increased hepatic lipogenesis causes a significant rise in serum triglyceride concentrations. There is increased protein catabolism as energy substrate, a negative nitrogen balance, and, if the process is prolonged, severe muscle wasting.

INFLAMMATORY RESPONSES

Activation of an extensive network of proinflammatory mediator pathways by the innate immune system plays a significant role in the progression of shock and contributes importantly to the development of multiple organ injury, multiple organ dysfunction (MOD), and MOF (**Fig. 324-2**). In those surviving the acute insult, there is a prolonged endogenous counterregulatory response to “turn off” or balance the excessive proinflammatory response. If balance is restored, the patient does well. If the response is excessive, adaptive immunity is suppressed and the patient is highly susceptible to secondary nosocomial infections, which may then drive the inflammatory response and lead to delayed MOF.

Multiple humoral mediators are activated during shock and tissue injury. The complement cascade, activated through both the classic and alternate pathways, generates the anaphylatoxins C3a and C5a (**Chap. 372e**). Direct complement fixation to injured tissues can progress to the C5-C9 attack complex, causing further cell damage. Activation of the coagulation cascade (**Chap. 141**) causes microvascular thrombosis, with subsequent fibrinolysis leading to repeated episodes of ischemia and reperfusion. Components of the coagulation system (e.g., thrombin) are potent proinflammatory mediators that cause expression of adhesion molecules on endothelial cells and activation of neutrophils, leading to microvascular injury. Coagulation also activates the kallikrein-kininogen cascade, contributing to hypotension.

Eicosanoids are vasoactive and immunomodulatory products of arachidonic acid metabolism that include cyclooxygenase-derived prostaglandins (PGs) and thromboxane A_2 , as well as lipoxygenase-derived leukotrienes and lipoxins. Thromboxane A_2 is a potent vasoconstrictor that contributes to the pulmonary hypertension and acute tubular necrosis of shock. PGI₂ and PGE₂ are potent vasodilators that enhance capillary permeability and edema formation. The cysteinyl leukotrienes LTC₄ and LTD₄ are pivotal mediators of the vascular sequelae of anaphylaxis, as well as of shock states resulting from sepsis or tissue injury. LTB₄ is a potent neutrophil chemoattractant and secretagogue that stimulates the formation of reactive oxygen species. Platelet-activating factor, an ether-linked, arachidonyl-containing phospholipid mediator, causes pulmonary vasoconstriction, bronchoconstriction, systemic vasodilation, increased capillary permeability, and the priming of macrophages and neutrophils to produce enhanced levels of inflammatory mediators.

Tumor necrosis factor α (TNF- α), produced by activated macrophages, reproduces many components of the shock state, including

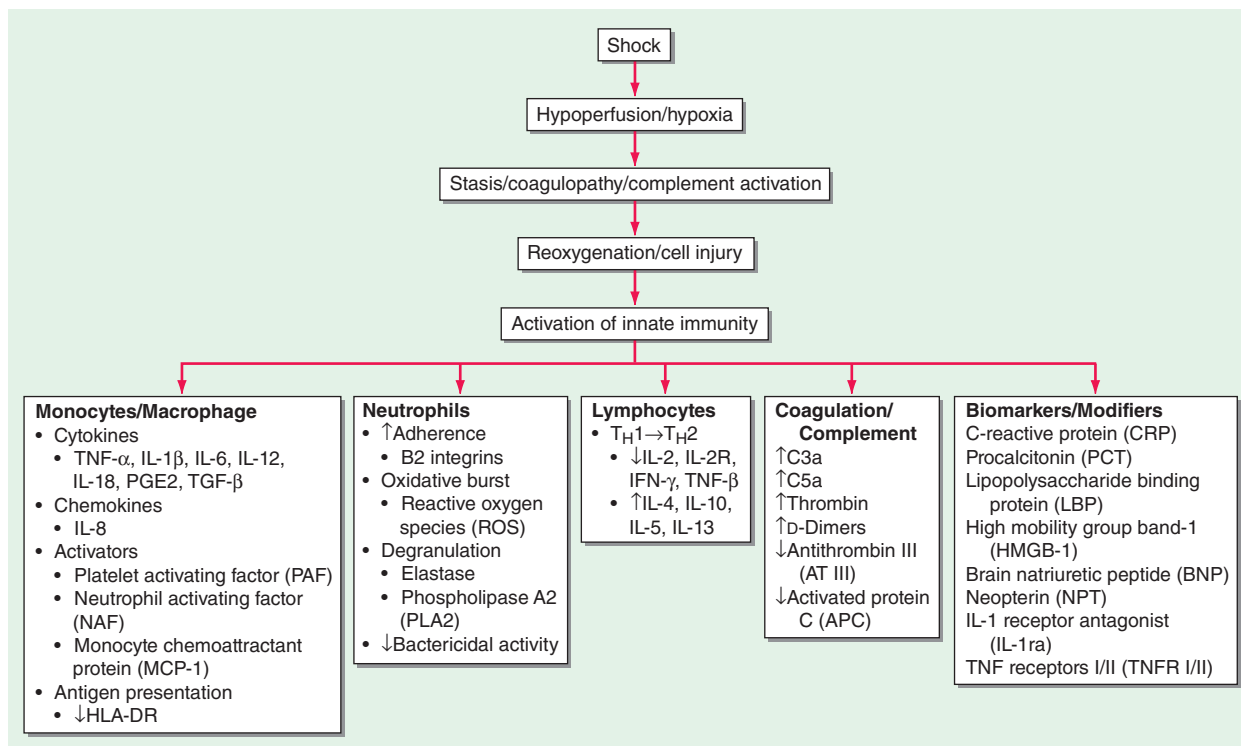


FIGURE 324-2 A schematic of the host immunoinflammatory response to shock. IFN, interferon; IL, interleukin; PG, prostaglandin; TGF, tumor growth factor; TNF, tumor necrosis factor.

hypotension, lactic acidosis, and respiratory failure. Interleukin 1 β (IL-1 β), originally defined as “endogenous pyrogen” and produced by tissue-fixed macrophages, is critical to the inflammatory response. Both are significantly elevated immediately following trauma and shock. IL-6, also produced predominantly by the macrophage, has a slightly delayed peak response but is the best single predictor of prolonged recovery and development of MOF following shock. Chemokines such as IL-8 are potent neutrophil chemoattractants and activators that upregulate adhesion molecules on the neutrophil to enhance aggregation, adherence, and damage to the vascular endothelium. While the endothelium normally produces low levels of NO, the inflammatory response stimulates the inducible isoform of NO synthase (iNOS), which is overexpressed and produces toxic nitroxyl- and oxygen-derived free radicals that contribute to the hyperdynamic cardiovascular response and tissue injury in sepsis.

Multiple inflammatory cells, including neutrophils, macrophages, and platelets, are major contributors to inflammation-induced injury. Margination of activated neutrophils in the microcirculation is a common pathologic finding in shock, causing secondary injury due to the release of toxic oxygen radicals, lipases (primarily PLA₂), and proteases. Release of high levels of reactive oxygen intermediates/species (ROI/ROS) rapidly consumes endogenous essential antioxidants and generates diffuse oxygen radical damage. Newer efforts to control ischemia/reperfusion injury include treatment with carbon monoxide, hydrogen sulfide, or other agents to reduce oxidant stress. Tissue-fixed macrophages produce virtually all major mediators of the inflammatory response and orchestrate the progression and duration of the inflammatory response. A major source of activation of the monocyte/macrophage is through the highly conserved membrane toll-like receptors (TLRs) that recognize DAMPs, such as HMGB-1, and pathogen-associated molecular patterns (PAMPs), such as endotoxins released following tissue injury, and by pathogenic microbial organisms, respectively. TLRs also appear important in the chronic inflammation seen in Crohn’s disease, ulcerative colitis, and transplant rejection. The variability in individual responses is a genetic predisposition that, in part, is due to variants in genetic sequences affecting the function and production of various inflammatory mediators.

TREATMENT SHOCK

MONITORING

Patients in shock require care in an intensive care unit (ICU). Careful and continuous assessment of the physiologic status is necessary. Arterial pressure through an indwelling line, pulse, and respiratory rate should be monitored continuously; a Foley catheter should be inserted to follow urine flow; and mental status should be assessed frequently. Sedated patients should be allowed to awaken (“drug holiday”) daily to assess their neurologic status and to shorten duration of ventilator support.

There is ongoing debate as to the indications for using the flow-directed pulmonary artery catheter (PAC; Swan-Ganz catheter) in the ICU. A recent Cochrane analysis showed that the use of a PAC did not alter mortality, length of stay, or cost for adult ICU patients. Most patients in the ICU can be safely managed without the use of a PAC. However, in shock with significant ongoing blood loss, fluid shifts, and underlying cardiac dysfunction, a PAC may be useful. The PAC is placed percutaneously via the subclavian or jugular vein through the central venous circulation and right heart into the pulmonary artery. There are ports both proximal in the right atrium and distal in the pulmonary artery to provide access for infusions and for cardiac output measurements. Right atrial and pulmonary artery pressures (PAPs) are measured, and the pulmonary capillary wedge pressure (PCWP) serves as an approximation of the left atrial pressure. Normal hemodynamic parameters and their derivation are summarized in [Table 272-2](#) and [Table 324-2](#).

Cardiac output is determined by the thermodilution technique, and high-resolution thermistors can also be used to determine right ventricular end-diastolic volume to monitor further the response of the right heart to fluid resuscitation. A PAC with an oximeter port offers the additional advantage of online monitoring of the mixed venous oxygen saturation, an important index of overall tissue perfusion. Systemic and pulmonary vascular resistances are calculated as the ratio of the pressure drop across these vascular beds to the cardiac output ([Chap. 272](#)). Determinations of oxygen content in arterial and venous blood, together with cardiac output and hemoglobin concentration, allow calculation of oxygen delivery,

TABLE 324-2 NORMAL HEMODYNAMIC PARAMETERS

Parameter	Calculation	Normal Values
Cardiac output (CO)	SV × HR	4–8 L/min
Cardiac index (CI)	CO/BSA	2.6–4.2 (L/min)/m ²
Stroke volume (SV)	CO/HR	50–100 mL/beat
Systemic vascular resistance (SVR)	((MAP – RAP)/CO) × 80	700–1600 dynes·s/cm ⁵
Pulmonary vascular resistance (PVR)	((PAP _m – PCWP)/CO) × 80	20–130 dynes·s/cm ⁵
Left ventricular stroke work (LVS _W)	SV(MAP – PCWP) × 0.0136	60–80 g·m/beat
Right ventricular stroke work (RVS _W)	SV(PAP _m – RAP)	10–15 g·m/beat

Abbreviations: BSA, body surface area; HR, heart rate; MAP, mean arterial pressure; PAP_m, pulmonary artery pressure—mean; PCWP, pulmonary capillary wedge pressure; RAP, right atrial pressure.

oxygen consumption, and oxygen-extraction ratio (Table 324-3). The hemodynamic patterns associated with the various forms of shock are shown in Table 324-4.

In resuscitation from shock, it is critical to restore tissue perfusion and optimize oxygen delivery, hemodynamics, and cardiac function rapidly. A reasonable goal of therapy is to achieve a normal mixed venous oxygen-saturation and arteriovenous oxygen-extraction ratio. To enhance oxygen delivery, red cell mass, arterial oxygen saturation, and cardiac output may be augmented singly or simultaneously. An increase in oxygen delivery not accompanied by an increase in oxygen consumption implies that oxygen availability is adequate and that oxygen consumption is not flow dependent. Conversely, an elevation of oxygen consumption with increased delivery implies that the oxygen supply was inadequate. However, cautious interpretation is required due to the link among increased oxygen delivery, cardiac work, and oxygen consumption. A reduction in systemic vascular resistance accompanying an increase in cardiac output indicates that compensatory vasoconstriction is reversing due to improved tissue perfusion. The determination of stepwise expansion of blood volume on cardiac performance allows identification of the optimum preload (Starling's law). An algorithm for the resuscitation of the patient in shock is shown in Fig. 324-3.

TABLE 324-3 OXYGEN TRANSPORT CALCULATIONS

Parameter	Calculation	Normal Values
Oxygen-carrying capacity of hemoglobin		1.39 mL/g
Plasma O ₂ concentration		Po ₂ × 0.0031
Arterial O ₂ concentration (Cao ₂)	1.39 Sao ₂ + 0.0031 Pao ₂	20 vol%
Venous O ₂ concentration (Cvo ₂)	1.39 Svo ₂ + 0.0031 Pvo ₂	15.5 vol%
Arteriovenous O ₂ difference (Cao ₂ – Cvo ₂)	1.39 (Sao ₂ – Svo ₂) + 0.0031 (Pao ₂ – Pvo ₂)	3.5 vol%
Oxygen delivery (Do ₂)	Cao ₂ × CO (L/min) × 10 (dL/L) 1.39 Sao ₂ × CO × 10	800–1600 mL/min
Oxygen uptake (Vo ₂)	(Cao ₂ – Cvo ₂) × CO × 10 1.39 (Sao ₂ – Svo ₂) × CO × 10	150–400 mL/min
Oxygen delivery index (Do ₂ I)	Do ₂ /BSA	520–720 (mL/min)/m ²
Oxygen uptake index (Vo ₂ I)	Vo ₂ /BSA	115–165 (mL/min)/m ²
Oxygen extraction ratio (O ₂ ER)	[1 – (Vo ₂ /Do ₂)] × 100	22–32%

Abbreviations: BSA, body surface area; CO, cardiac output; Po₂, partial pressure of oxygen; Pao₂, partial pressure of oxygen in arterial blood; Pvo₂, partial pressure of oxygen in venous blood; Sao₂, saturation of hemoglobin with oxygen in arterial blood; Svo₂, saturation of hemoglobin with oxygen in venous blood.

TABLE 324-4 PHYSIOLOGIC CHARACTERISTICS OF THE VARIOUS FORMS OF SHOCK

Type of Shock	CVP and PCWP	Cardiac Output	Systemic Vascular Resistance	Venous O ₂ Saturation
Hypovolemic	↓	↓	↑	↓
Cardiogenic	↑	↓	↑	↓
Septic				
Hyperdynamic	↓↑	↑	↓	↑
Hypodynamic	↓↑	↓	↑	↓↑
Traumatic	↓	↓↑	↑↓	↓
Neurogenic	↓	↓	↓	↓
Hypoadrenal	↓	↓	=↓	↓

Abbreviations: CVP, central venous pressure; PCWP, pulmonary capillary wedge pressure.

SPECIFIC FORMS OF SHOCK

HYPOVOLEMIC SHOCK

This most common form of shock results either from the loss of red blood cell mass and plasma from hemorrhage or from the loss of plasma volume alone due to extravascular fluid sequestration or GI, urinary, and insensible losses. The signs and symptoms of nonhemorrhagic hypovolemic shock are the same as those of hemorrhagic shock, although they may have a more insidious onset. The normal physiologic response to hypovolemia is to maintain perfusion of the brain and heart while attempting to restore an effective circulating blood volume. There is an increase in sympathetic activity, hyperventilation, collapse of venous capacitance vessels, release of stress hormones, and an attempt to replace the loss of intravascular volume through the recruitment of interstitial and intracellular fluid and by reduction of urine output.

Mild hypovolemia (≤20% of the blood volume) generates mild tachycardia but relatively few external signs, especially in a supine young patient (Table 324-5). With moderate hypovolemia (~20–40% of the blood volume), the patient becomes increasingly anxious and tachycardic; although normal blood pressure may be maintained in the supine position, there may be significant postural hypotension and tachycardia. If hypovolemia is severe (≥40% of the blood volume), the classic signs of shock appear; the blood pressure declines and becomes unstable even in the supine position, and the patient develops marked tachycardia, oliguria, and agitation or confusion. Perfusion of the central nervous system is well maintained until shock becomes severe. Hence, mental obtundation is an ominous clinical sign. The transition from mild to severe hypovolemic shock can be insidious or extremely rapid. If severe shock is not reversed rapidly, especially in elderly patients and those with comorbid illnesses, death is imminent. A very narrow time frame separates the derangements found in severe shock that can be reversed with aggressive resuscitation from those of progressive decompensation and irreversible cell injury.

Diagnosis Hypovolemic shock is readily diagnosed when there are signs of hemodynamic instability and the source of volume loss is obvious. The diagnosis is more difficult when the source of blood loss is occult, as into the GI tract, or when plasma volume alone is depleted. Even after acute hemorrhage, hemoglobin and hematocrit values do not change until compensatory fluid shifts have occurred or exogenous fluid is administered. Thus, an initial normal hematocrit does not disprove the presence of significant blood loss. Plasma losses cause hemoconcentration, and free water loss leads to hypernatremia. These findings should suggest the presence of hypovolemia.

It is essential to distinguish between hypovolemic and cardiogenic shock (Chap. 326) because, although both may respond to volume initially, definitive therapy differs significantly. Both forms are associated with a reduced cardiac output and a compensatory sympathetic mediated response characterized by tachycardia and elevated systemic vascular resistance. However, the findings in cardiogenic shock of jugular venous distention, rales, and an S₃ gallop distinguish it from hypovolemic shock and signify that ongoing volume expansion is undesirable and may cause further organ dysfunction.

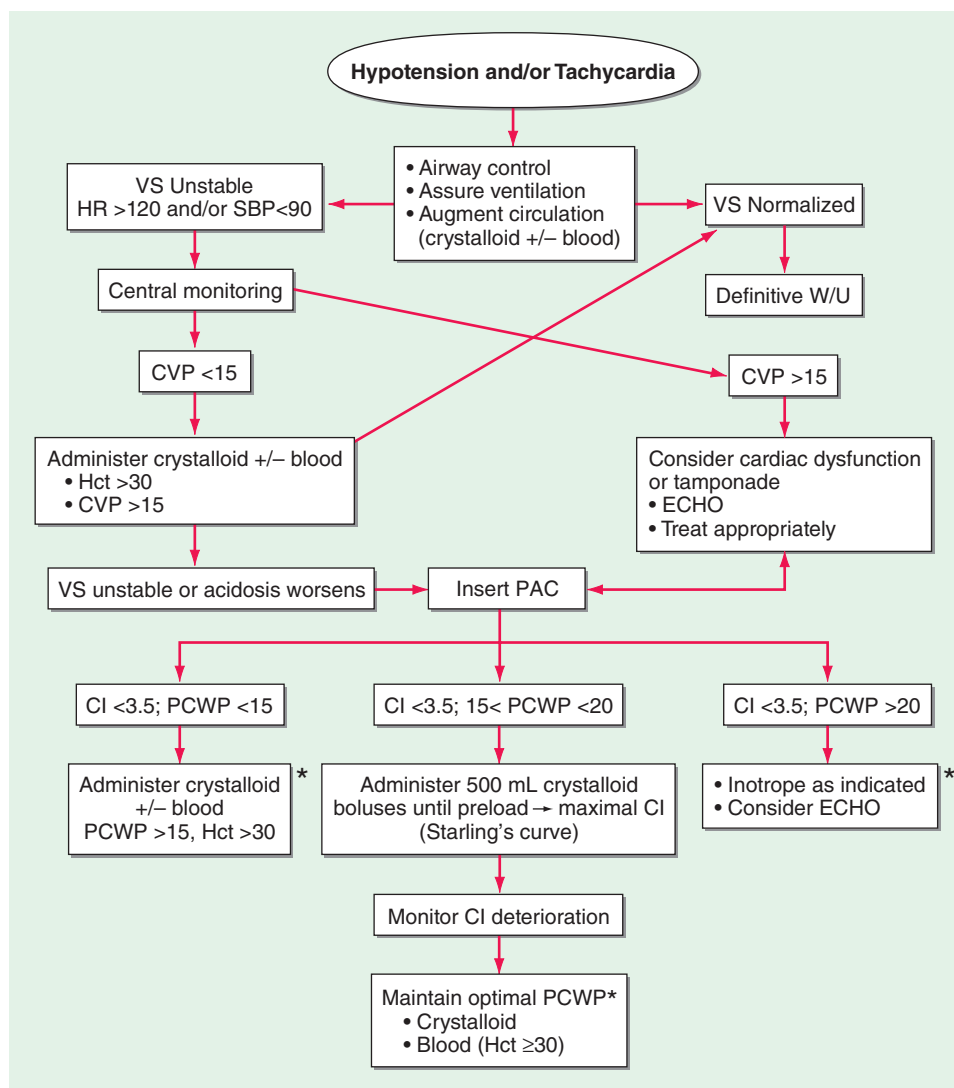


FIGURE 324-3 An algorithm for the resuscitation of the patient in shock. *Monitor Sv_{O_2} , SVRI, and RVEDVI as additional markers of correction for perfusion and hypovolemia. Consider age-adjusted CI. CI, cardiac index in (L/min) per m^2 ; CVP, central venous pressure; ECHO, echocardiogram; Hct, hematocrit; HR, heart rate; PAC, pulmonary artery catheter; PCWP, pulmonary capillary wedge pressure in mmHg; RVEDVI, right ventricular end-diastolic volume index; SBP, systolic blood pressure; Sv_{O_2} , saturation of hemoglobin with O_2 in venous blood; SVRI, systemic vascular resistance index; VS, vital signs; W/U, workup.

TREATMENT HYPOVOLEMIC SHOCK

Initial resuscitation requires rapid reexpansion of the circulating intravascular blood volume along with interventions to control ongoing losses. In accordance with Starling's law (Chap. 265e), stroke volume and cardiac output rise with the increase in preload. After resuscitation, the compliance of the ventricles may remain reduced due to increased interstitial fluid in the myocardium.

TABLE 324-5 HYPOVOLEMIC SHOCK

Mild (<20% Blood Volume)	Moderate (20–40% Blood Volume)	Severe (>40% Blood Volume)
Cool extremities	Same, plus:	Same, plus:
Increased capillary refill time	Tachycardia	Hemodynamic instability
Diaphoresis	Tachypnea	Marked tachycardia
Collapsed veins	Oliguria	Hypotension
Anxiety	Postural changes	Mental status deterioration (coma)

Therefore, elevated filling pressures are frequently required to maintain adequate ventricular performance.

Volume resuscitation is initiated with the rapid infusion of either isotonic saline (although care must be taken to avoid hyperchloremic acidosis from loss of bicarbonate buffering capacity and replacement with excess chloride) or a balanced salt solution such as Ringer's lactate (being cognizant of the presence of potassium and potential renal dysfunction) through large-bore intravenous lines. Data, particularly on severe traumatic brain injury (TBI), regarding benefits of small volumes of hypertonic saline that more rapidly restore blood pressure are variable but tend to show improved survival thought to be linked to immunomodulation. No distinct benefit from the use of colloid has been demonstrated, and in trauma patients, it is associated with a higher mortality particularly in patients with TBI. The infusion of 2–3 L of salt solution over 20–30 min should restore normal hemodynamic parameters. Continued hemodynamic instability implies that shock has not been reversed and/or there are significant ongoing blood or other volume losses. Continuing acute blood loss with hemoglobin concentrations declining to ≤ 100 g/L (10 g/dL) should initiate blood transfusion preferably as fully cross-matched, recently banked (<14 days old) blood. Resuscitated patients are often coagulopathic due

1750 to deficient clotting factors in crystalloids and banked packed red blood cells (PRBCs). Early administration of component therapy during massive transfusion (fresh-frozen plasma [FFP] and platelets) approaching a 1:1 ratio of PRBC/FFP appears to improve survival. In extreme emergencies, type-specific or O-negative packed red cells may be transfused. Following severe and/or prolonged hypovolemia, inotropic support with norepinephrine, vasopressin, or dopamine may be required to maintain adequate ventricular performance *but only after* blood volume has been restored. Increases in peripheral vasoconstriction with inadequate resuscitation lead to tissue loss and organ failure. Once hemorrhage is controlled and the patient has stabilized, blood transfusions should not be continued unless the hemoglobin is <7 g/dL. Studies have demonstrated an increased survival in patients treated with this restrictive blood transfusion protocol.

Successful resuscitation also requires support of respiratory function. Supplemental oxygen should always be provided, and endotracheal intubation may be necessary to maintain arterial oxygenation. Following resuscitation from isolated hemorrhagic shock, end-organ damage is frequently less than following septic or traumatic shock. This may be due to the absence of massive activation of the inflammatory innate immune response and consequent nonspecific organ injury and failure.

TRAUMATIC SHOCK

Shock following trauma is, in large measure, due to hemorrhage. However, even when hemorrhage has been controlled, patients can continue to suffer loss of plasma volume into the interstitium of injured tissues. These fluid losses are compounded by injury-induced inflammatory responses, which contribute to the secondary microcirculatory injury. Proinflammatory mediators are induced by DAMPs released from injured tissue and are recognized by the highly conserved membrane receptors of the TLR family (see “Inflammatory Responses” above). These receptors on cells of the innate immune system, particularly the circulating monocyte, tissue-fixed macrophage, and dendritic cell, are potent activators of an excessive proinflammatory phenotype in response to cellular injury. This causes secondary tissue injury and maldistribution of blood flow, intensifying tissue ischemia and leading to multiple organ system failure. In addition, direct structural injury to the heart, chest, or head can also contribute to shock. For example, pericardial tamponade or tension pneumothorax impairs ventricular filling, whereas myocardial contusion depresses myocardial contractility.

TREATMENT TRAUMATIC SHOCK

Inability of the patient to maintain a systolic blood pressure ≥ 90 mmHg after trauma-induced hypovolemia is associated with a mortality rate up to $\sim 50\%$. To prevent this decompensation of homeostatic mechanisms, therapy must be promptly administered.

The initial management of the seriously injured patient requires attention to the “ABCs” of resuscitation: assurance of an *airway* (A), adequate ventilation (*breathing*, B), and establishment of an adequate blood volume to support the *circulation* (C). Control of ongoing hemorrhage requires immediate attention. Early stabilization of fractures, debridement of devitalized or contaminated tissues, and evacuation of hematoma all reduce the subsequent inflammatory response to the initial insult and minimize damaged tissue release of DAMPs and subsequent diffuse organ injury. Supplementation of depleted endogenous antioxidants also reduces subsequent organ failure and mortality.

CARDIOGENIC SHOCK
See Chap. 326.

COMPRESSIVE CARDIOGENIC SHOCK

With extrinsic compression, the heart and surrounding structures are less compliant, and therefore, normal filling pressures generate inadequate diastolic filling and stroke volume. Blood or fluid within the poorly distensible pericardial sac may cause tamponade (Chap. 288). Any cause of increased intrathoracic pressure, such as tension pneumothorax, herniation of abdominal viscera through a diaphragmatic hernia, or excessive positive-pressure ventilation to support pulmonary function, can also initiate compressive cardiogenic shock while simultaneously impeding venous return and preload. Although initially responsive to increased filling pressures produced by volume expansion, as compression increases, cardiogenic shock recurs. The window of opportunity gained by volume loading may be very brief until irreversible shock recurs. Diagnosis and intervention must occur urgently.

The diagnosis of compressive cardiogenic shock is most frequently based on clinical findings, the chest radiograph, and an echocardiogram. The diagnosis of compressive cardiac shock may be more difficult to establish in the setting of trauma when hypovolemia and cardiac compression are present simultaneously. The classic findings of pericardial tamponade include the triad of hypotension, neck vein distention, and muffled heart sounds (Chap. 288). Pulsus paradoxus (i.e., an inspiratory reduction in systolic pressure >10 mmHg) may also be noted. The diagnosis is confirmed by echocardiography, and treatment consists of immediate pericardiocentesis or the creation of an open subxiphoid pericardial window. A tension pneumothorax produces ipsilateral decreased breath sounds, tracheal deviation away from the affected thorax, and jugular venous distention. Radiographic findings include increased intrathoracic volume, depression of the diaphragm of the affected hemithorax, and shifting of the mediastinum to the contralateral side. Chest decompression must be carried out immediately and, ideally, should occur based on clinical findings rather than awaiting a chest radiograph. Release of air and restoration of normal cardiovascular dynamics are both diagnostic and therapeutic.

SEPTIC SHOCK

See Chap. 325.

NEUROGENIC SHOCK

Interruption of sympathetic vasomotor input after a high cervical spinal cord injury, inadvertent cephalad migration of spinal anesthesia, or devastating head injury may result in neurogenic shock. In addition to arteriolar dilation, venodilation causes pooling in the venous system, which decreases venous return and cardiac output. The extremities are often warm, in contrast to the usual sympathetic vasoconstriction-induced coolness in hypovolemic or cardiogenic shock. Treatment involves a simultaneous approach to the relative hypovolemia and to the loss of vasomotor tone. Excessive volumes of fluid may be required to restore normal hemodynamics if given alone. Once hemorrhage has been ruled out, norepinephrine or a pure α -adrenergic agent (phenylephrine) may be necessary to augment vascular resistance and maintain an adequate MAP.

HYPOADRENAL SHOCK

(See also Chap. 406) The normal host response to the stress of illness, operation, or trauma requires that the adrenal glands hypersecrete cortisol in excess of that normally required. Hypoadrenal shock occurs in settings in which unrecognized adrenal insufficiency complicates the host response to the stress induced by acute illness or major surgery. Adrenocortical insufficiency may occur as a consequence of the chronic administration of high doses of exogenous glucocorticoids. In addition, recent studies have shown that critical illness, including trauma and sepsis, may also induce a relative hypoadrenal state. Other, less common causes include adrenal insufficiency secondary to idiopathic atrophy, use of etomidate for intubation, tuberculosis, metastatic disease, bilateral hemorrhage, and amyloidosis. The shock produced by adrenal insufficiency is characterized by loss of homeostasis with reductions in systemic vascular resistance, hypovolemia, and reduced cardiac output. The diagnosis of adrenal insufficiency may be established by means of an ACTH stimulation test.

In the persistently hemodynamically unstable patient, dexamethasone sodium phosphate, 4 mg, should be given intravenously. This agent is preferred if empiric therapy is required because, unlike hydrocortisone, it does not interfere with the ACTH stimulation test. If the diagnosis of absolute or relative adrenal insufficiency is established as shown by nonresponse to corticotropin stimulation (cortisol ≤ 9 $\mu\text{g/dL}$ change after stimulation), the patient has a reduced risk of death if treated with hydrocortisone, 100 mg every 6–8 h, and tapered as the patient achieves hemodynamic stability. Simultaneous volume resuscitation and pressor support are required. The need for simultaneous mineralocoid is unclear.

ADJUNCTIVE THERAPIES

The sympathomimetic amines dobutamine, dopamine, and norepinephrine are widely used in the treatment of all forms of shock. Dobutamine is inotropic with simultaneous afterload reduction, thus minimizing cardiac-oxygen consumption increases as cardiac output increases. Dopamine is an inotropic and chronotropic agent that also supports vascular resistance in those whose blood pressure will not tolerate peripheral vascular dilation. Norepinephrine primarily supports blood pressure through vasoconstriction and increases myocardial oxygen consumption while placing marginally perfused tissues, such as extremities and splanchnic organs, at risk for ischemia or necrosis, but it is also inotropic without significant chronotropy. Arginine-vasopressin (antidiuretic hormone) is being used increasingly to increase afterload and may better protect vital organ blood flow and prevent pathologic vasodilation.

REWARMING

Hypothermia is a frequent adverse consequence of massive volume resuscitation (Chap. 478e). The infusion of large volumes of refrigerated blood products and room temperature crystalloid solutions can rapidly drop core temperatures if fluid is not run through warming devices. Hypothermia may depress cardiac contractility and thereby further impair cardiac output and oxygen delivery/utilization. Hypothermia, particularly temperatures $<35^{\circ}\text{C}$ ($<95^{\circ}\text{F}$), directly impairs the coagulation pathway, sometimes causing a significant coagulopathy. Rapid rewarming to $>35^{\circ}\text{C}$ ($>95^{\circ}\text{F}$) significantly decreases the requirement for blood products and produces an improvement in cardiac function. The most effective method for rewarming is endovascular countercurrent warmers through femoral vein cannulation. This process does not require a pump and can rewarm a patient from 30° to 35°C (86° to 95°F) in 30–60 min.

325 Severe Sepsis and Septic Shock

Robert S. Munford

DEFINITIONS

(Table 325-1) Animals mount both local and systemic responses to microbes that traverse their epithelial barriers and enter underlying tissues. Fever or hypothermia, leukocytosis or leukopenia, tachypnea, and tachycardia are cardinal signs of the systemic response. To date, attempts to devise precise definitions for the harmful systemic reaction to infection (“sepsis”) have not resulted in a clinically useful level of specificity, in part because the systemic responses to infection, trauma, and other major stresses can be so similar. In general, when an infectious etiology is proven or strongly suspected and the response results in hypofunction of uninfected organs, the term *sepsis* (or *severe sepsis*) should be used. *Septic shock* refers to sepsis accompanied by hypotension that cannot be corrected by the infusion of fluids.

TABLE 325-1 DEFINITIONS USED TO DESCRIBE THE CONDITION OF SEPTIC PATIENTS

Bacteremia	Presence of bacteria in blood, as evidenced by positive blood cultures
Signs of possibly harmful systemic response	Two or more of the following conditions: (1) fever (oral temperature $>38^{\circ}\text{C}$ [$>100.4^{\circ}\text{F}$]) or hypothermia ($<36^{\circ}\text{C}$ [$<96.8^{\circ}\text{F}$]); (2) tachypnea (>24 breaths/min); (3) tachycardia (heart rate >90 beats/min); (4) leukocytosis ($>12,000/\mu\text{L}$), leukopenia ($<4000/\mu\text{L}$), or $>10\%$ bands
Sepsis (or severe sepsis)	The harmful host response to infection; systemic response to proven or suspected infection plus some degree of organ hypofunction, i.e.: <ol style="list-style-type: none"> Cardiovascular: Arterial systolic blood pressure ≤ 90 mmHg or mean arterial pressure ≤ 70 mmHg that responds to administration of IV fluid Renal: Urine output <0.5 mL/kg per hour for 1 h despite adequate fluid resuscitation Respiratory: $\text{Pao}_2/\text{Fio}_2 \leq 250$ or, if the lung is the only dysfunctional organ, ≤ 200 Hematologic: Platelet count $<80,000/\mu\text{L}$ or 50% decrease in platelet count from highest value recorded over previous 3 days Unexplained metabolic acidosis: A pH ≤ 7.30 or a base deficit ≥ 5.0 mEq/L and a plasma lactate level >1.5 times upper limit of normal for reporting lab
Septic shock	Sepsis with hypotension (arterial blood pressure <90 mmHg systolic, or 40 mmHg less than patient’s normal blood pressure) for at least 1 h despite adequate fluid resuscitation ^a or Need for vasopressors to maintain systolic blood pressure ≥ 90 mmHg or mean arterial pressure ≥ 70 mmHg
Refractory septic shock	Septic shock that lasts for >1 h and does not respond to fluid or pressor administration

^aFluid resuscitation is considered adequate when the pulmonary artery wedge pressure is ≥ 12 mmHg or the central venous pressure is ≥ 8 mmHg.

ETIOLOGY


 The systemic response to any class of microorganism can be harmful. Microbial invasion of the bloodstream is not essential because local inflammation can also elicit distant organ dysfunction and hypotension. In fact, blood cultures yield bacteria or fungi in only ~20–40% of cases of severe sepsis and 40–70% of cases of septic shock. In a prevalence study of 14,414 patients in intensive care units (ICUs) from 75 countries in 2007, 51% of patients were considered infected. Respiratory infection was most common (64%). Microbiologic results were positive in 70% of individuals considered infected; of the isolates, 62% were gram-negative bacteria (*Pseudomonas* species and *Escherichia coli* were most common), 47% were gram-positive bacteria (*Staphylococcus aureus* was most common), and 19% were fungi (*Candida* species). This distribution is similar to that reported a decade earlier from eight academic centers in the United States (Table 325-2). In patients whose blood cultures are negative, the etiologic agent is often established by culture or microscopic examination of infected material from a local site; specific identification of microbial DNA or RNA in blood or tissue samples is also used. In some case series, a majority of patients with a clinical picture of severe sepsis or septic shock have had negative microbiologic data.

TABLE 325-2 MICROORGANISMS INVOLVED IN EPISODES OF SEVERE SEPSIS AT EIGHT ACADEMIC MEDICAL CENTERS

Microorganisms	Episodes with Bloodstream Infection, % (n = 436)	Episodes with Documented Infection but No Bloodstream Infection, % (n = 430)	Total Episodes, % (n = 866)
Gram-negative bacteria ^a	35	44	40
Gram-positive bacteria ^b	40	24	31
Fungi	7	5	6
Polymicrobial	11	21	16
Classic pathogens ^c	<5	<5	<5

^aEnterobacteriaceae, pseudomonads, *Haemophilus* spp., other gram-negative bacteria. ^b*Staphylococcus aureus*, coagulase-negative staphylococci, enterococci, *Streptococcus pneumoniae*, other streptococci, other gram-positive bacteria. ^cSuch as *Neisseria meningitidis*, *S. pneumoniae*, *Haemophilus influenzae*, and *Streptococcus pyogenes*.

Source: Adapted from KE Sands et al: JAMA 278:234, 1997.

EPIDEMIOLOGY

Severe sepsis is a contributing factor in >200,000 deaths per year in the United States. The incidence of severe sepsis and septic shock has increased over the past 30 years, and the annual number of cases is now >750,000 (~3 per 1000 population). Approximately two-thirds of the cases occur in patients with significant underlying illness. Sepsis-related incidence and mortality rates increase with age and preexisting comorbidity. The rising incidence of severe sepsis in the United States has been attributable to the aging of the population, the increasing longevity of patients with chronic diseases, and the relatively high frequency with which sepsis has occurred in patients with AIDS. The widespread use of immunosuppressive drugs, indwelling catheters, and mechanical devices has also played a role. In the aforementioned international ICU prevalence study, the case-fatality rate among infected patients (33%) greatly exceeded that among uninfected patients (15%).



Invasive bacterial infections are prominent causes of death around the world, particularly among young children. In sub-Saharan Africa, for example, careful screening for positive blood cultures found that community-acquired bacteremia accounted for at least one-fourth of deaths of children >1 year of age. Nontyphoidal *Salmonella* species, *Streptococcus pneumoniae*, *Haemophilus influenzae*, and *E. coli* were the most commonly isolated bacteria. Bacteremic children often had HIV infection or were severely malnourished.

PATHOPHYSIOLOGY

Sepsis is triggered most often by bacteria or fungi that do not ordinarily cause systemic disease in immunocompetent hosts (Table 325-2). To survive within the human body, these microbes often exploit acquired deficiencies in host defenses, indwelling catheters or other foreign matter, or obstructed fluid drainage conduits. Microbial pathogens, in contrast, can circumvent innate defenses because they (1) lack molecules that can be recognized by host receptors (see below) or (2) elaborate toxins or other virulence factors. In both cases, the body can mount a vigorous inflammatory reaction that results in sepsis or septic shock yet fails to kill the invaders. The septic response may also be induced by microbial exotoxins that act as superantigens (e.g., toxic shock syndrome toxin 1; Chap. 172) as well as by many pathogenic viruses.

Host Mechanisms for Sensing Microbes Animals have exquisitely sensitive mechanisms for recognizing and responding to certain highly conserved microbial molecules. Recognition of the lipid A moiety of lipopolysaccharide (LPS, also called *endotoxin*; Chap. 145e) is the best-studied example. A host protein (LPS-binding protein) binds lipid A and transfers the LPS to CD14 on the surfaces of monocytes, macrophages, and neutrophils. LPS then is passed to MD-2, a small receptor protein that is bound to Toll-like receptor (TLR) 4 to form a molecular complex that transduces the LPS recognition signal to the interior of the cell. This signal rapidly triggers the production and release of mediators, such as tumor necrosis factor (TNF; see below), that amplify the LPS signal and transmit it to other cells and tissues. Bacterial peptidoglycan and lipopeptides elicit responses in animals that are generally similar to those induced by LPS, although they interact with different TLRs. Having numerous TLR-based receptor

complexes (10 different TLRs have been identified in humans) allows animals to recognize many conserved microbial molecules; others include lipopeptides (TLR2/1, TLR2/6), flagellin (TLR5), undermethylated DNA CpG sequences (TLR9), single-stranded RNA (TLR7, 8), and double-stranded RNA (TLR3). The ability of some TLRs to serve as receptors for host ligands (e.g., hyaluronans, heparan sulfate, saturated fatty acids, high-mobility group box 1) raises the possibility that they also play a role in producing noninfectious sepsis-like states. Other host pattern-recognition proteins that are important for sensing microbes include the intracellular NOD1 and NOD2 proteins, which recognize discrete fragments of bacterial peptidoglycan; the inflammasome, which senses some pathogens and produces interleukin (IL) 1 β and IL-18; early complement components (principally in the alternative pathway); mannose-binding lectin and C-reactive protein, which activate the classic complement pathway; and Dectin-1 and complement receptor 3, which sense fungal β -glucan.

A host's ability to recognize certain microbial molecules may influence both the potency of its own defenses and the pathogenesis of severe sepsis. For example, MD-2-TLR4 best senses LPS that has a bisphosphorylated, hexaacyl lipid A moiety (i.e., one with two phosphates and six fatty acyl chains). Most of the commensal aerobic and facultatively anaerobic gram-negative bacteria that trigger severe sepsis and shock (including *E. coli*, *Klebsiella*, and *Enterobacter*) make this lipid A structure. When they invade human hosts, often through breaks in an epithelial barrier, they are typically confined to the subepithelial tissue by a localized inflammatory response. Bacteremia, if it occurs, is intermittent and low grade because these bacteria are efficiently cleared from the bloodstream by TLR4-expressing Kupffer cells and splenic macrophages. These mucosal commensals seem to induce severe sepsis most often by triggering severe local tissue inflammation rather than by circulating within the bloodstream. One exception is *Neisseria meningitidis*. Its hexaacyl LPS seems to be shielded from host recognition by its polysaccharide capsule. This protection may allow meningococci to transit undetected from the nasopharyngeal mucosa into the bloodstream, where they can infect vascular endothelial cells and release large amounts of endotoxin and DNA. Host recognition of lipid A may nonetheless influence pathogenesis, as meningococci that produce pentaacyl LPS were isolated from the blood of patients with less severe coagulopathy than was found in patients whose isolates produced hexaacyl lipid A; underacylated *N. meningitidis* LPS has also been found in many isolates from patients with chronic meningococcemia. In contrast, gram-negative bacteria that make lipid A with fewer than six acyl chains (*Yersinia pestis*, *Francisella tularensis*, *Vibrio vulnificus*, *Pseudomonas aeruginosa*, and *Burkholderia pseudomallei*, among others) are poorly recognized by MD-2-TLR4. When these bacteria enter the body, they may initially induce relatively little inflammation. When they do trigger severe sepsis, it is often after they have multiplied to high density in tissues and blood. The importance of LPS recognition in disease pathogenesis was demonstrated by engineering of a virulent strain of *Y. pestis* that makes tetraacyl LPS at 37°C to produce hexaacyl LPS; unlike its virulent parent, the mutant strain stimulated local inflammation and was rapidly cleared from tissues. These findings were subsequently replicated in *F. tularensis*. For at least one large class of microbes—gram-negative aerobic bacteria—the

pathogenesis of sepsis thus depends, at least in part, on whether the bacterium's major signal molecule, LPS, can be sensed by the host.

Local and Systemic Host Responses to Invading Microbes Recognition of microbial molecules by tissue phagocytes triggers the production and/or release of numerous host molecules (cytokines, chemokines, prostanoids, leukotrienes, and others) that increase blood flow to the infected tissue (*rubor*), enhance the permeability of local blood vessels (*tumor*), recruit neutrophils and other cells to the site of infection (*calor*), and elicit pain (*dolor*). These reactions are familiar elements of local inflammation, the body's frontline innate immune mechanism for eliminating microbial invaders. Systemic responses are activated by neural and/or humoral communication with the hypothalamus and brainstem; these responses enhance local defenses by increasing blood flow to the infected area, augmenting the number of circulating neutrophils, and elevating blood levels of numerous molecules (such as the microbial recognition proteins discussed above) that have anti-infective functions.

CYTOKINES AND OTHER MEDIATORS Cytokines can exert endocrine, paracrine, and autocrine effects (Chap. 372e). TNF- α stimulates leukocytes and vascular endothelial cells to release other cytokines (as well as additional TNF- α), to express cell-surface molecules that enhance neutrophil endothelial adhesion at sites of infection, and to increase prostaglandin and leukotriene production. Whereas blood levels of TNF- α are not elevated in individuals with localized infections, they increase in most patients with severe sepsis or septic shock. Moreover, IV infusion of TNF- α can elicit fever, tachycardia, hypotension, and other responses. In animals, larger doses of TNF- α induce shock and death.

Although TNF- α is a central mediator, it is only one of many proinflammatory molecules that contribute to innate host defense. Chemokines, most prominently IL-8 and IL-17, attract circulating neutrophils to the infection site. IL-1 β exhibits many of the same activities as TNF- α . TNF- α , IL-1 β , interferon γ , IL-12, IL-17, and other proinflammatory cytokines probably interact synergistically with one another and with additional mediators. The nonlinearity and multiplicity of these interactions have made it difficult to interpret the roles played by individual mediators in both tissues and blood.

COAGULATION FACTORS Intravascular thrombosis, a hallmark of the local inflammatory response, may help wall off invading microbes and prevent infection and inflammation from spreading to other tissues. IL-6 and other mediators promote intravascular coagulation initially by inducing blood monocytes and vascular endothelial cells to express tissue factor (Chap. 78). When tissue factor is expressed on cell surfaces, it binds to factor VIIa to form an active complex that can convert factors X and IX to their enzymatically active forms. The result is activation of both extrinsic and intrinsic clotting pathways, culminating in the generation of fibrin. Clotting is also favored by impaired function of the protein C–protein S inhibitory pathway and depletion of antithrombin and proteins C and S, whereas fibrinolysis is reduced by increases in plasma levels of plasminogen activator inhibitor 1. Thus, there may be a striking propensity toward intravascular fibrin deposition, thrombosis, and bleeding; this propensity has been most apparent in patients with intravascular endothelial infections such as meningococcemia (Chap. 180). Evidence points to tissue factor–expressing microparticles derived from leukocytes as a potential trigger for intravascular coagulation. The contact system is activated during sepsis but contributes more to the development of hypotension than to that of disseminated intravascular coagulation (DIC).

Neutrophil extracellular traps (NETs) are produced when neutrophils, stimulated by microbial agonists or IL-8, release granule proteins and chromatin to form an extracellular fibrillar matrix. NETs kill bacteria and fungi with antimicrobial granule proteins (e.g., elastase) and histones. It has been reported that NETs can form within hepatic sinusoids in animals injected with large amounts of LPS, and platelets can induce NET formation without killing neutrophils. A role played by NETs in organ hypofunction during sepsis has been proposed but not established.

CONTROL MECHANISMS Elaborate control mechanisms operate within both local sites of inflammation and the systemic compartment.

Local control mechanisms Host recognition of invading microbes within subepithelial tissues typically ignites immune responses that rapidly kill the invaders and then subside to allow tissue recovery. The forces that put out the fire and clean up the battleground include molecules that neutralize or inactivate microbial signals. Among these molecules are intracellular factors (e.g., suppressor of cytokine signaling 3 and IL-1 receptor–associated kinase 3) that diminish the production of proinflammatory mediators by neutrophils and macrophages; anti-inflammatory cytokines (IL-10, IL-4); and molecules derived from essential polyunsaturated fatty acids (lipoxins, resolvins, and protectins) that promote tissue restoration. Enzymatic inactivation of microbial signal molecules (e.g., LPS) may be required to restore homeostasis; a leukocyte enzyme, acyloxycyl hydrolase, has been shown to prevent prolonged inflammation in mice by inactivating LPS.

Systemic control mechanisms The signaling apparatus that links microbial recognition to cellular responses in tissues is less active in the blood. For example, whereas LPS-binding protein plays a role in recognizing LPS, in plasma it also prevents LPS signaling by transferring LPS molecules into plasma lipoprotein particles that sequester the lipid A moiety so that it cannot interact with cells. At the high concentrations found in blood, LPS-binding protein also inhibits monocyte responses to LPS, and the soluble (circulating) form of CD14 strips off LPS that has bound to monocyte surfaces.

Systemic responses to infection also diminish cellular responses to microbial molecules. Circulating levels of cortisol and anti-inflammatory cytokines (e.g., IL-6 and IL-10) increase even in patients with minor infections. Glucocorticoids inhibit cytokine synthesis by monocytes in vitro; the increase in blood cortisol levels that occurs early in the systemic response presumably plays a similarly inhibitory role. Epinephrine inhibits the TNF- α response to endotoxin infusion in humans while augmenting and accelerating the release of IL-10; prostaglandin E₂ has a similar “reprogramming” effect on the responses of circulating monocytes to LPS and other bacterial agonists. Cortisol, epinephrine, IL-10, and C-reactive protein reduce the ability of neutrophils to attach to vascular endothelium, favoring their demargination and thus contributing to leukocytosis while preventing neutrophil-endothelial adhesion in uninflamed organs. Studies in rodents have found that macrophage cytokine synthesis is inhibited by acetylcholine that is produced by choline acetyltransferase–secreting CD4+ T cells in response to stimulation by norepinephrine, whereas acetylcholine-producing B cells reduce neutrophil infiltration into tissues. Several lines of evidence thus suggest that the body's neuroendocrine responses to injury and infection normally prevent inflammation within organs distant from a site of infection. There is also evidence that these responses may be immunosuppressive.

IL-6 plays important roles in the systemic compartment. Released by many different cell types, IL-6 is an important stimulus to the hypothalamic-pituitary-adrenal axis, is the major procoagulant cytokine, and is a principal inducer of the acute-phase response, which increases the blood concentrations of numerous molecules that have anti-infective, procoagulant, or anti-inflammatory actions. Blood levels of IL-1 receptor antagonist often greatly exceed those of circulating IL-1 β , for example, and this excess may inhibit the binding of IL-1 β to its receptors. High levels of soluble TNF receptors neutralize TNF- α that enters the circulation. Other acute-phase proteins are protease inhibitors or antioxidants; these may neutralize potentially harmful molecules released from neutrophils and other inflammatory cells. Increased hepatic production of hepcidin (stimulated largely by IL-6) promotes the sequestration of iron in hepatocytes, intestinal epithelial cells, and erythrocytes; this effect reduces iron acquisition by invading microbes while contributing to the normocytic, normochromic anemia associated with inflammation.

It may thus be said that both local and systemic responses to infectious agents benefit the host in important ways. Most of these responses and the molecules responsible for them have been highly conserved during animal evolution and therefore may be adaptive.

1754 Elucidating how they become maladaptive and contribute to lethality remains a major challenge for sepsis research.

Organ Dysfunction and Shock As the body's responses to infection intensify, the mixture of circulating cytokines and other molecules becomes very complex: elevated blood levels of more than 60 molecules have been found in patients with septic shock. Although high concentrations of both pro- and anti-inflammatory molecules are found, the net mediator balance in the plasma of these extremely sick patients seems to be anti-inflammatory. For example, blood leukocytes from patients with severe sepsis are often hyporesponsive to agonists such as LPS. In patients with severe sepsis, persistence of leukocyte hyporesponsiveness has been associated with an increased risk of dying; at this time, the most predictive biomarker is a decrease in the expression of HLA-DR (class II) molecules on the surfaces of circulating monocytes, a response that seems to be induced by cortisol and/or IL-10. Apoptotic death of B cells, follicular dendritic cells, and CD4+ T lymphocytes also may contribute significantly to the immunosuppressive state.

ENDOTHELIAL INJURY Given the vascular endothelium's important roles in regulating vascular tone, vascular permeability, and coagulation, many investigators have favored widespread vascular endothelial injury as the major mechanism for multiorgan dysfunction. In keeping with this idea, one study found high numbers of vascular endothelial cells in the peripheral blood of septic patients. Leukocyte-derived mediators and platelet-leukocyte-fibrin thrombi may contribute to vascular injury, but the vascular endothelium also seems to play an active role. Stimuli such as TNF- α induce vascular endothelial cells to produce and release cytokines, procoagulant molecules, platelet-activating factor, nitric oxide, and other mediators. In addition, regulated cell-adhesion molecules promote the adherence of neutrophils to endothelial cells. Although these responses can attract phagocytes to infected sites and activate their antimicrobial arsenals, endothelial cell activation can also promote increased vascular permeability, microvascular thrombosis, DIC, and hypotension.

Tissue oxygenation may decrease as the number of functional capillaries is reduced by luminal obstruction due to swollen endothelial cells, decreased deformability of circulating erythrocytes, leukocyte-platelet-fibrin thrombi, or compression by edema fluid. On the other hand, studies using orthogonal polarization spectral imaging of the microcirculation in the tongue found that sepsis-associated derangements in capillary flow could be reversed by applying acetylcholine to the surface of the tongue or by giving nitroprusside intravenously; these observations suggest a neuroendocrine basis for the loss of capillary filling. Oxygen utilization by tissues may also be impaired by changes (possibly induced by nitric oxide) that decrease oxidative phosphorylation and ATP production while increasing glycolysis. The local accumulation of lactic acid, a consequence of increased glycolysis, may decrease extracellular pH and contribute to the slowdown in cellular metabolism that occurs within affected tissues.

Remarkably, poorly functioning "septic" organs usually appear normal at autopsy. There is typically very little necrosis or thrombosis, and apoptosis is largely confined to lymphoid organs and the gastrointestinal tract. Moreover, organ function usually returns to normal if patients recover. These points suggest that organ dysfunction during severe sepsis has a basis that is principally biochemical, not structural.

SEPTIC SHOCK The hallmark of septic shock is a decrease in peripheral vascular resistance that occurs despite increased levels of vasopressor catecholamines. Before this vasodilatory phase, many patients experience a period during which oxygen delivery to tissues is compromised by myocardial depression, hypovolemia, and other factors. During this "hypodynamic" period, the blood lactate concentration is elevated and central venous oxygen saturation is low. Fluid administration is usually followed by the hyperdynamic vasodilatory phase, during which cardiac output is normal (or even high) and oxygen consumption declines despite adequate oxygen delivery. The blood lactate level may be normal or increased, and normalization of central venous oxygen saturation may reflect improved oxygen delivery, decreased oxygen uptake by tissues, or left-to-right shunting.

Prominent hypotensive molecules include nitric oxide, β -endorphin, bradykinin, platelet-activating factor, and prostacyclin. Agents that inhibit the synthesis or action of each of these mediators can prevent or reverse endotoxic shock in animals. However, in clinical trials, neither a platelet-activating factor receptor antagonist nor a bradykinin antagonist improved survival rates among patients with septic shock, and a nitric oxide synthase inhibitor, L-N^G-methylarginine HCl, actually increased the mortality rate.

Severe Sepsis: A Single Pathogenesis? In some cases, circulating bacteria and their products almost certainly elicit multiorgan dysfunction and hypotension by directly stimulating inflammatory responses within the vasculature. In patients with fulminant meningococcemia, for example, mortality rates have correlated directly with blood levels of endotoxin and bacterial DNA and with the occurrence of DIC (**Chap. 180**). In most patients infected with other gram-negative bacteria, in contrast, circulating bacteria or bacterial molecules may reflect uncontrolled infection at a local tissue site and have little or no direct impact on distant organs; in these patients, inflammatory mediators or neural signals arising from the local site seem to be the key triggers for severe sepsis and septic shock. In a large series of patients with positive blood cultures, the risk of developing severe sepsis was strongly related to the site of primary infection: bacteremia arising from a pulmonary or abdominal source was eightfold more likely to be associated with severe sepsis than was bacteremic urinary tract infection, even after the investigators controlled for age, the kind of bacteria isolated from the blood, and other factors. A third pathogenesis may be represented by severe sepsis due to superantigen-producing *S. aureus* or *Streptococcus pyogenes*; the T cell activation induced by these toxins produces a cytokine profile that differs substantially from that elicited by gram-negative bacterial infection. Further evidence for different pathogenic pathways has come from observations that the pattern of mRNA expression in peripheral-blood leukocytes from children with sepsis is different for gram-positive, gram-negative, and viral pathogens.

The pathogenesis of severe sepsis thus may differ according to the infecting microbe, the ability of the host's innate defense mechanisms to sense and respond to it, the site of the primary infection, the presence or absence of immune defects, and the prior physiologic status of the host. Genetic factors are probably important as well, yet despite much study very few allelic polymorphisms have been associated with sepsis severity in more than one or two analyses. Further studies in this area are needed.

CLINICAL MANIFESTATIONS

The manifestations of the septic response are superimposed on the symptoms and signs of the patient's underlying illness and primary infection. The rate at which severe sepsis develops may differ from patient to patient, and there are striking individual variations in presentation. For example, some patients with sepsis are normo- or hypothermic; the absence of fever is most common in neonates, in elderly patients, and in persons with uremia or alcoholism.

Hyperventilation, producing respiratory alkalosis, is often an early sign of the septic response. Disorientation, confusion, and other manifestations of encephalopathy may also develop early on, particularly in the elderly and in individuals with preexisting neurologic impairment. Focal neurologic signs are uncommon, although preexisting focal deficits may become more prominent.

Hypotension and DIC predispose to acrocyanosis and ischemic necrosis of peripheral tissues, most commonly the digits. Cellulitis, pustules, bullae, or hemorrhagic lesions may develop when hematogenous bacteria or fungi seed the skin or underlying soft tissue. Bacterial toxins may also be distributed hematogenously and elicit diffuse cutaneous reactions. On occasion, skin lesions may suggest specific pathogens. When sepsis is accompanied by cutaneous petechiae or purpura, infection with *N. meningitidis* (or, less commonly, *H. influenzae*) should be suspected (**see Fig. 25e-42**); in a patient who has been bitten by a tick while in an endemic area, petechial lesions also suggest Rocky Mountain spotted fever (**see Fig. 211-1**). A cutaneous lesion seen almost exclusively in neutropenic patients is ecthyma gangrenosum, often caused by *P. aeruginosa*. This bullous lesion surrounded by edema undergoes central hemorrhage and necrosis (**see Fig. 189-1**).

Histopathologic examination shows bacteria in and around the wall of a small vessel, with little or no neutrophilic response. Hemorrhagic or bullous lesions in a septic patient who has recently eaten raw oysters suggest *V. vulnificus* bacteremia, whereas such lesions in a patient who has recently sustained a dog bite may indicate bloodstream infection due to *Capnocytophaga canimorsus* or *Capnocytophaga cynodegmi*. Generalized erythroderma in a septic patient suggests the toxic shock syndrome due to *S. aureus* or *S. pyogenes*.

Gastrointestinal manifestations such as nausea, vomiting, diarrhea, and ileus may suggest acute gastroenteritis. Stress ulceration can lead to upper gastrointestinal bleeding. Cholestatic jaundice, with elevated levels of serum bilirubin (mostly conjugated) and alkaline phosphatase, may precede other signs of sepsis. Hepatocellular or canalicular dysfunction appears to underlie most cases, and the results of hepatic function tests return to normal with resolution of the infection. Prolonged or severe hypotension may induce acute hepatic injury or ischemic bowel necrosis.

Many tissues may be unable to extract oxygen normally from the blood, so that anaerobic metabolism occurs despite near-normal mixed venous oxygen saturation. Blood lactate levels rise early because of increased glycolysis as well as impaired clearance of the resulting lactate and pyruvate by the liver and kidneys. The blood glucose concentration often increases, particularly in patients with diabetes, although impaired gluconeogenesis and excessive insulin release on occasion produce hypoglycemia. The cytokine-driven acute-phase response inhibits the synthesis of transthyretin while enhancing the production of C-reactive protein, fibrinogen, and complement components. Protein catabolism is often markedly accelerated. Serum albumin levels decline as a result of decreased hepatic synthesis and the movement of albumin into interstitial spaces.

MAJOR COMPLICATIONS

Cardiopulmonary Complications Ventilation-perfusion mismatching produces a fall in arterial Po_2 early in the course. Increasing alveolar epithelial injury and capillary permeability result in increased pulmonary water content, which decreases pulmonary compliance and interferes with oxygen exchange. In the absence of pneumonia or heart failure, progressive diffuse pulmonary infiltrates and arterial hypoxemia occurring within 1 week of a known insult indicate the development of mild acute respiratory distress syndrome (ARDS) ($200 \text{ mmHg} < \text{PaO}_2/\text{FiO}_2 \leq 300 \text{ mmHg}$), moderate ARDS ($100 \text{ mmHg} < \text{PaO}_2/\text{FiO}_2 \leq 200 \text{ mmHg}$), or severe ARDS ($\text{PaO}_2/\text{FiO}_2 \leq 100 \text{ mmHg}$). Acute lung injury or ARDS develops in ~50% of patients with severe sepsis or septic shock. Respiratory muscle fatigue can exacerbate hypoxemia and hypercapnia. An elevated pulmonary capillary wedge pressure ($>18 \text{ mmHg}$) suggests fluid volume overload or cardiac failure rather than ARDS. Pneumonia caused by viruses or by *Pneumocystis* may be clinically indistinguishable from ARDS.

Sepsis-induced hypotension (see “Septic Shock,” above) usually results initially from a generalized maldistribution of blood flow and blood volume and from hypovolemia that is due, at least in part, to diffuse capillary leakage of intravascular fluid. Other factors that may decrease effective intravascular volume include dehydration from antecedent disease or insensible fluid losses, vomiting or diarrhea, and polyuria. During early septic shock, systemic vascular resistance is usually elevated and cardiac output may be low. After fluid repletion, in contrast, cardiac output typically increases and systemic vascular resistance falls. Indeed, normal or increased cardiac output and decreased systemic vascular resistance distinguish septic shock from cardiogenic, extracardiac obstructive, and hypovolemic shock; other processes that can produce this combination include anaphylaxis, beriberi, cirrhosis, and overdoses of nitroprusside or narcotics.

Depression of myocardial function, manifested as increased end-diastolic and systolic ventricular volumes with a decreased ejection fraction, develops within 24 h in most patients with severe sepsis. Cardiac output is maintained despite the low ejection fraction because ventricular dilation permits a normal stroke volume. In survivors, myocardial function returns to normal over several days. Although myocardial dysfunction may contribute to hypotension, refractory

hypotension is usually due to low systemic vascular resistance, and death most often results from refractory shock or the failure of multiple organs rather than from cardiac dysfunction per se.

Adrenal Insufficiency The diagnosis of adrenal insufficiency may be very difficult in critically ill patients. Whereas a plasma cortisol level of $\leq 15 \text{ }\mu\text{g/mL}$ ($\leq 10 \text{ }\mu\text{g/mL}$ if the serum albumin concentration is $<2.5 \text{ mg/dL}$) indicates adrenal insufficiency (inadequate production of cortisol), many experts now feel that the adrenocorticotropic hormone (CoSyntropin[®]) stimulation test is not useful for detecting less profound degrees of corticosteroid deficiency in patients who are critically ill. The concept of critical illness–related corticosteroid insufficiency (CIRCI) was proposed to encompass the different mechanisms that may produce corticosteroid activity that is inadequate for the severity of a patient’s illness. Although CIRCI may result from structural damage to the adrenal gland, it is more commonly due to reversible dysfunction of the hypothalamic-pituitary axis or to tissue corticosteroid resistance resulting from abnormalities of the glucocorticoid receptor or increased conversion of cortisol to cortisone. The major clinical manifestation of CIRCI is hypotension that is refractory to fluid replacement and requires pressor therapy. Some classic features of adrenal insufficiency, such as hyponatremia and hyperkalemia, are usually absent; others, such as eosinophilia and modest hypoglycemia, may sometimes be found. Specific etiologies include fulminant *N. meningitidis* bacteremia, disseminated tuberculosis, AIDS (with cytomegalovirus, *Mycobacterium avium-intracellulare*, or *Histoplasma capsulatum* disease), or the prior use of drugs that diminish glucocorticoid production, such as glucocorticoids, megestrol, etomidate, or ketoconazole.

Renal Complications Oliguria, azotemia, proteinuria, and nonspecific urinary casts are frequently found. Many patients are inappropriately polyuric; hyperglycemia may exacerbate this tendency. Most renal failure is due to acute tubular necrosis induced by hypovolemia, arterial hypotension, or toxic drugs, although some patients also have glomerulonephritis, renal cortical necrosis, or interstitial nephritis. Drug-induced renal damage may greatly complicate therapy, particularly when hypotensive patients are given aminoglycoside antibiotics. Nosocomial sepsis following acute renal injury is associated with a high mortality rate.

Coagulopathy Although thrombocytopenia occurs in 10–30% of patients, the underlying mechanisms are not understood. Platelet counts are usually very low ($<50,000/\mu\text{L}$) in patients with DIC; these low counts may reflect diffuse endothelial injury or microvascular thrombosis, yet thrombi have only infrequently been found on biopsy of septic organs.

Neurologic Complications Delirium (acute encephalopathy) is often an early manifestation of sepsis. Depending on the diagnostic criteria used, it occurs in 10–70% of septic patients at some point during the hospital course. When the septic illness lasts for weeks or months, “critical illness” polyneuropathy may prevent weaning from ventilatory support and produce distal motor weakness. Electrophysiologic studies are diagnostic. Guillain-Barré syndrome, metabolic disturbances, and toxin activity must be ruled out. Recent studies have documented long-term cognitive loss in survivors of severe sepsis.

Immunosuppression Patients with severe sepsis often become profoundly immunosuppressed. Manifestations include loss of delayed-type hypersensitivity reactions to common antigens, failure to control the primary infection, and increased risk for secondary infections (e.g., by opportunists such as *Stenotrophomonas maltophilia*, *Acinetobacter calcoaceticus-baumannii*, and *Candida albicans*). Approximately one-third of patients experience reactivation of herpes simplex virus, varicella-zoster virus, or cytomegalovirus infections; the latter are thought to contribute to adverse outcomes in some instances.

LABORATORY FINDINGS

Abnormalities that occur early in the septic response may include leukocytosis with a left shift, thrombocytopenia, hyperbilirubinemia, and

1756 proteinuria. Leukopenia may develop. The neutrophils may contain toxic granulations, Döhle bodies, or cytoplasmic vacuoles. As the septic response becomes more severe, thrombocytopenia worsens (often with prolonged thrombin time, decreased fibrinogen, and the presence of D-dimers, suggesting DIC), azotemia and hyperbilirubinemia become more prominent, and levels of aminotransferases rise. Active hemolysis suggests clostridial bacteremia, malaria, a drug reaction, or DIC; in the case of DIC, microangiopathic changes may be seen on a blood smear.

During early sepsis, hyperventilation induces respiratory alkalosis. With respiratory muscle fatigue and the accumulation of lactate, metabolic acidosis (with increased anion gap) typically supervenes. Evaluation of arterial blood gases reveals hypoxemia that is initially correctable with supplemental oxygen but whose later refractoriness to 100% oxygen inhalation indicates right-to-left shunting. The chest radiograph may be normal or may show evidence of underlying pneumonia, volume overload, or the diffuse infiltrates of ARDS. The electrocardiogram may show only sinus tachycardia or nonspecific ST-T wave abnormalities.

Most diabetic patients with sepsis develop hyperglycemia. Severe infection may precipitate diabetic ketoacidosis that may exacerbate hypotension (Chap. 417). Hypoglycemia occurs rarely and may indicate adrenal insufficiency. The serum albumin level declines as sepsis continues. Hypocalcemia is rare.

DIAGNOSIS

There is no specific diagnostic test for sepsis. Diagnostically sensitive findings in a patient with suspected or proven infection include fever or hypothermia, tachypnea, tachycardia, and leukocytosis or leukopenia (Table 325-1); acutely altered mental status, thrombocytopenia, an elevated blood lactate level, respiratory alkalosis, or hypotension also should suggest the diagnosis. The systemic response can be quite variable, however. In one study, 36% of patients with severe sepsis had a normal temperature, 40% had a normal respiratory rate, 10% had a normal pulse rate, and 33% had normal white blood cell counts. Moreover, the systemic responses of uninfected patients with other conditions may be similar to those characteristic of sepsis. Examples include pancreatitis, burns, trauma, adrenal insufficiency, pulmonary embolism, dissecting or ruptured aortic aneurysm, myocardial infarction, occult hemorrhage, cardiac tamponade, postcardiopulmonary bypass syndrome, anaphylaxis, tumor-associated lactic acidosis, and drug overdose.

Definitive etiologic diagnosis requires identification of the causative microorganism from blood or a local site of infection. At least two blood samples should be obtained (from two different venipuncture sites) for culture; in a patient with an indwelling catheter, one sample should be collected from each lumen of the catheter and another via venipuncture. In many cases, blood cultures are negative; this result can reflect prior antibiotic administration, the presence of slow-growing or fastidious organisms, or the absence of microbial invasion of the bloodstream. In these cases, Gram's staining and culture of material from the primary site of infection or from infected cutaneous lesions may help establish the microbial etiology. Identification of microbial DNA in peripheral blood or tissue samples by polymerase chain reaction may also be definitive. The skin and mucosae should be examined carefully and repeatedly for lesions that might yield diagnostic information. With overwhelming bacteremia (e.g., pneumococcal sepsis in splenectomized individuals; fulminant meningococcemia; or infection with *V. vulnificus*, *B. pseudomallei*, or *Y. pestis*), microorganisms are sometimes visible on buffy coat smears of peripheral blood.

TREATMENT SEVERE SEPSIS AND SEPTIC SHOCK

Patients in whom sepsis is suspected must be managed expeditiously. This task is best accomplished by personnel who are experienced in the care of the critically ill. Successful management requires urgent measures to treat the infection, to provide hemodynamic and respiratory support, and to remove or drain infected tissues. These measures should be initiated within 1 h of

the patient's presentation with severe sepsis or septic shock. Rapid assessment and diagnosis are therefore essential.

ANTIMICROBIAL AGENTS

Antimicrobial chemotherapy should be started as soon as samples of blood and other relevant sites have been obtained for culture. A large retrospective review of patients who developed septic shock found that the interval between the onset of hypotension and the administration of appropriate antimicrobial chemotherapy was the major determinant of outcome; a delay of as little as 1 h was associated with lower survival rates. Use of "inappropriate" antibiotics, defined on the basis of local microbial susceptibilities and published guidelines for empirical therapy (see below), was associated with fivefold lower survival rates, even among patients with negative cultures.

It is therefore very important to promptly initiate empirical antimicrobial therapy that is effective against both gram-positive and gram-negative bacteria (Table 325-3). Maximal recommended doses of antimicrobial drugs should be given intravenously, with adjustment for impaired renal function when necessary. Available information about patterns of antimicrobial susceptibility among bacterial isolates from the community, the hospital, and the patient should be taken into account. When culture results become available, the regimen can often be simplified because a single antimicrobial agent is usually adequate for the treatment of a known pathogen. Meta-analyses have concluded that, with one exception, combination antimicrobial therapy is not superior to monotherapy for treating gram-negative bacteremia; the exception is that aminoglycoside monotherapy for *P. aeruginosa* bacteremia is less effective than the combination of an aminoglycoside with an antipseudomonal β -lactam agent. Empirical antifungal therapy should be strongly considered if the septic patient is already receiving broad-spectrum antibiotics or parenteral nutrition, has been neutropenic for ≥ 5 days, has had a long-term central venous catheter in place, or has been hospitalized in an ICU for a prolonged period. The chosen antimicrobial regimen should be reconsidered daily in order to provide maximal efficacy with minimal resistance, toxicity, and cost.

Most patients require antimicrobial therapy for at least 1 week. The duration of treatment is typically influenced by factors such as the site of tissue infection, the adequacy of surgical drainage, the patient's underlying disease, and the antimicrobial susceptibility of the microbial isolate(s). The absence of an identified microbial pathogen is not necessarily an indication for discontinuing antimicrobial therapy because "appropriate" antimicrobial regimens seem to be beneficial in both culture-negative and culture-positive cases.

REMOVAL OF THE SOURCE OF INFECTION

Removal or drainage of a focal source of infection is essential. In one series, a focus of ongoing infection was found in ~80% of surgical ICU patients who died of severe sepsis or septic shock. Sites of occult infection should be sought carefully, particularly in the lungs, abdomen, and urinary tract. Indwelling IV or arterial catheters should be removed and the tip rolled over a blood agar plate for quantitative culture; after antibiotic therapy has been initiated, a new catheter should be inserted at a different site. Foley and drainage catheters should be replaced. The possibility of paranasal sinusitis (often caused by gram-negative bacteria) should be considered if the patient has undergone nasal intubation or has an indwelling nasogastric or feeding tube. Even in patients without abnormalities on chest radiographs, computed tomography (CT) of the chest may identify unsuspected parenchymal, mediastinal, or pleural disease. In the neutropenic patient, cutaneous sites of tenderness and erythema, particularly in the perianal region, must be carefully sought. In patients with sacral or ischial decubitus ulcers, it is important to exclude pelvic or other soft tissue pus collections with CT or magnetic resonance imaging (MRI). In patients with severe sepsis arising from the urinary tract, sonography or CT should be used to rule out ureteral obstruction, perinephric abscess, and renal abscess. Sonographic or CT imaging of the upper abdomen may disclose

TABLE 325-3 INITIAL ANTIMICROBIAL THERAPY FOR SEVERE SEPSIS WITH NO OBVIOUS SOURCE IN ADULTS WITH NORMAL RENAL FUNCTION

Clinical Condition	Antimicrobial Regimens (Intravenous Therapy)
Immunocompetent adult	The many acceptable regimens include (1) piperacillin-tazobactam (3.375 g q4–6h); (2) imipenem-cilastatin (0.5 g q6h), ertapenem (1 g q24h), or meropenem (1 g q8h); or (3) cefepime (2 g q12h). If the patient is allergic to β -lactam agents, use ciprofloxacin (400 mg q12h) or levofloxacin (500–750 mg q12h) plus clindamycin (600 mg q8h). Vancomycin (15 mg/kg q12h) should be added to each of the above regimens.
Neutropenia (<500 neutrophils/ μ L)	Regimens include (1) imipenem-cilastatin (0.5 g q6h) or meropenem (1 g q8h) or cefepime (2 g q8h) or (2) piperacillin-tazobactam (3.375 g q4h) plus tobramycin (5–7 mg/kg q24h). Vancomycin (15 mg/kg q12h) should be added if the patient has an indwelling vascular catheter, has received quinolone prophylaxis, or has received intensive chemotherapy that produces mucosal damage; if staphylococci are suspected; if the institution has a high incidence of MRSA infections; or if there is a high prevalence of MRSA isolates in the community. Empirical antifungal therapy with an echinocandin (for caspofungin: a 70-mg loading dose, then 50 mg daily), voriconazole (6 mg/kg q12h for 2 doses, then 3 mg/kg q12h), or a lipid formulation of amphotericin B should be added if the patient is hypotensive, has been receiving broad-spectrum antibacterial drugs, or remains febrile 5 days after initiation of empirical antibacterial therapy.
Splenectomy	Cefotaxime (2 g q6–8h) or ceftriaxone (2 g q12h) should be used. If the local prevalence of cephalosporin-resistant pneumococci is high, add vancomycin. If the patient is allergic to β -lactam drugs, vancomycin (15 mg/kg q12h) plus either moxifloxacin (400 mg q24h) or levofloxacin (750 mg q24h) should be used.
IV drug user	Vancomycin (15 mg/kg q12h) is essential.
AIDS	Cefepime alone (2 g q8h) or piperacillin-tazobactam (3.375 g q4h) plus tobramycin (5–7 mg/kg q24h) should be used. If the patient is allergic to β -lactam drugs, ciprofloxacin (400 mg q12h) or levofloxacin (750 mg q12h) plus vancomycin (15 mg/kg q12h) plus tobramycin should be used.

Abbreviation: MRSA, methicillin-resistant *Staphylococcus aureus*.

Source: Adapted in part from DN Gilbert et al: *The Sanford Guide to Antimicrobial Therapy*, 43rd ed, 2013.

evidence of cholecystitis, bile duct dilation, and pus collections in the liver, subphrenic space, or spleen.

HEMODYNAMIC, RESPIRATORY, AND METABOLIC SUPPORT

The primary goals are to restore adequate oxygen and substrate delivery to the tissues as quickly as possible and to improve tissue oxygen utilization and cellular metabolism. Adequate organ perfusion

is thus essential. Circulatory adequacy is assessed by measurement of arterial blood pressure and monitoring of parameters such as mentation, urine output, and skin perfusion. Indirect indices of oxygen delivery and consumption, such as central venous oxygen saturation, may also be useful. Initial management of hypotension should include the administration of IV fluids, typically beginning with 1–2 L of normal saline over 1–2 h. To avoid pulmonary edema, the central venous pressure should be maintained at 8–12 cmH₂O. The urine output rate should be kept at >0.5 mL/kg per hour by continuing fluid administration; a diuretic such as furosemide may be used if needed. In about one-third of patients, hypotension and organ hypoperfusion respond to fluid resuscitation; a reasonable goal is to maintain a mean arterial blood pressure of >65 mmHg (systolic pressure >90 mmHg). If these guidelines cannot be met by volume infusion, vasopressor therapy is indicated (**Chap. 326**). Titrated doses of norepinephrine should be administered through a central catheter. If myocardial dysfunction produces elevated cardiac filling pressures and low cardiac output, inotropic therapy with dobutamine is recommended. Dopamine is rarely used.

In patients with septic shock, plasma vasopressin levels increase transiently but then decrease dramatically. Early studies found that vasopressin infusion can reverse septic shock in some patients, reducing or eliminating the need for catecholamine pressors. Although vasopressin may benefit patients who require less norepinephrine, its role in the treatment of septic shock seems to be a minor one overall.

CIRCI (see “Adrenal Insufficiency,” above) should be strongly considered in patients who develop hypotension that does not respond to fluid replacement therapy. Hydrocortisone (50 mg IV every 6 h) should be given; if clinical improvement occurs over 24–48 h, most experts would continue hydrocortisone therapy for 5–7 days before slowly tapering and discontinuing it. Meta-analyses of recent clinical trials have concluded that hydrocortisone therapy hastens recovery from sepsis-induced hypotension without increasing long-term survival.

Ventilator therapy is indicated for progressive hypoxemia, hypercapnia, neurologic deterioration, or respiratory muscle failure. Sustained tachypnea (respiratory rate, >30 breaths/min) is frequently a harbinger of impending respiratory collapse; mechanical ventilation is often initiated to ensure adequate oxygenation, to divert blood from the muscles of respiration, to prevent aspiration of oropharyngeal contents, and to reduce the cardiac afterload. The results of recent studies favor the use of low tidal volumes (6 mL/kg of ideal body weight, or as low as 4 mL/kg if the plateau pressure exceeds 30 cmH₂O). Patients undergoing mechanical ventilation require careful sedation, with daily interruptions; elevation of the head of the bed helps to prevent nosocomial pneumonia. Stress-ulcer prophylaxis with a histamine H₂-receptor antagonist may decrease the risk of gastrointestinal hemorrhage in ventilated patients.

Erythrocyte transfusion is generally recommended when the blood hemoglobin level decreases to \leq 7 g/dL, with a target level of 9 g/dL in adults. Erythropoietin is not used to treat sepsis-related anemia. Bicarbonate is sometimes administered for severe metabolic acidosis (arterial pH <7.2), but there is little evidence that it improves either hemodynamics or the response to vasopressor hormones. DIC, if complicated by major bleeding, should be treated with transfusion of fresh-frozen plasma and platelets. Successful treatment of the underlying infection is essential to reverse both acidosis and DIC. Patients who are hypercatabolic and have acute renal failure may benefit greatly from intermittent hemodialysis or continuous veno-venous hemofiltration.

GENERAL SUPPORT

In patients with prolonged severe sepsis (i.e., that lasting more than 2 or 3 days), nutritional supplementation may reduce the impact of protein hypercatabolism; the available evidence favors the enteral delivery route. Prophylactic heparinization to prevent deep venous thrombosis is indicated for patients who do not have active bleeding or coagulopathy; when heparin is contraindicated, compression

1758 stockings or an intermittent compression device should be used. Recovery is also assisted by prevention of skin breakdown, nosocomial infections, and stress ulcers.

The role of tight control of the blood glucose concentration in recovery from critical illness has been addressed in numerous controlled trials. Meta-analyses of these trials have concluded that use of insulin to lower blood glucose levels to 100–120 mg/dL is potentially harmful and does not improve survival rates. Most experts now recommend using insulin only if it is needed to maintain the blood glucose concentration below ~180 mg/dL. Patients receiving intravenous insulin must be monitored frequently (every 1–2 h) for hypoglycemia.

OTHER MEASURES

Despite aggressive management, many patients with severe sepsis or septic shock die. Numerous interventions have been tested for their ability to improve survival rates among patients with severe sepsis. The list includes endotoxin-neutralizing proteins, inhibitors of cyclooxygenase or nitric oxide synthase, anticoagulants, polyclonal immunoglobulins, glucocorticoids, a phospholipid emulsion, and antagonists to TNF- α , IL-1, platelet-activating factor, and bradykinin. Unfortunately, none of these agents has improved rates of survival among patients with severe sepsis/septic shock in more than one large-scale, randomized, placebo-controlled clinical trial. Many factors have contributed to this lack of reproducibility, including (1) heterogeneity of the patient populations studied, the primary infection sites, the preexisting illnesses, and the inciting microbes; and (2) the nature of the “standard” therapy also used. A dramatic example of this problem was seen in a trial of tissue factor pathway inhibitor. Whereas the drug appeared to improve survival rates after 722 patients had been studied ($p = .006$), it did not do so in the next 1032 patients, and the overall result was negative. This inconsistency argues that the results of a clinical trial may not apply to individual patients, even within a carefully selected patient population. It also suggests that, at a minimum, a sepsis intervention should show a significant survival benefit in more than one placebo-controlled, randomized clinical trial before it is accepted as routine clinical practice. In one attempt to reduce patient heterogeneity in clinical trials, experts have called for changes that would restrict these trials to patients who have similar underlying diseases (e.g., major trauma) and inciting infections (e.g., pneumonia). Other investigators have proposed using specific biomarkers, such as IL-6 levels in blood or the expression of HLA-DR on peripheral-blood monocytes, to identify the patients most likely to benefit from certain interventions.

Recombinant activated protein C (aPC) was the first immunomodulatory drug to be approved by the U.S. Food and Drug Administration (FDA) for the treatment of patients with severe sepsis or septic shock. Approval was based on the results of a single randomized controlled trial in which the drug was given within 24 h of the patient’s first sepsis-related organ dysfunction; the 28-day survival rate was significantly higher among aPC recipients who were very sick (APACHE II score, ≥ 25) before infusion of the protein than among placebo-treated controls. Subsequent trials failed to show a benefit of aPC treatment in patients who were less sick (APACHE II score, < 25) or in children, and, a decade after its licensure by the FDA, the drug was withdrawn from the market when a European trial failed to confirm its efficacy in adults with sepsis. Agents in ongoing or planned clinical trials include intravenous immunoglobulin, a polymyxin B hemofiltration column, and granulocyte-macrophage colony-stimulating factor, which has been reported to restore monocyte immunocompetence in patients with sepsis-associated immunosuppression.

A careful retrospective analysis found that the apparent efficacy of all sepsis therapeutics studied to date has been greatest among the patients at greatest risk of dying before treatment; conversely, use of many of these drugs has been associated with increased mortality rates among patients who are less ill. It is possible that neutralizing one of many different mediators may help patients who are very sick, whereas disrupting the mediator balance may

be harmful to patients whose adaptive defense mechanisms are working well. This analysis suggests that if more aggressive early resuscitation improves survival rates among sicker patients, it will become more difficult to obtain additional benefit from other therapies; that is, if an intervention improves patients’ risk status, moving them into a “less severe illness” category, it will be harder to show that adding another agent to the therapeutic regimen is beneficial.

THE SURVIVING SEPSIS CAMPAIGN

An international consortium has advocated “bundling” of multiple therapeutic maneuvers into a unified algorithmic approach that will become the standard of care for severe sepsis. In theory, such a strategy would improve care by mandating measures that seem to bring maximal benefit, such as the rapid administration of appropriate antimicrobial therapy, fluids, and blood pressure support. Caution may be engendered by the fact that three of the key elements of the initial algorithm were eventually withdrawn for lack of evidence; moreover, the benefit of the current sepsis bundles has not been established in randomized controlled clinical trials.

PROGNOSIS

Approximately 20–35% of patients with severe sepsis and 40–60% of patients with septic shock die within 30 days. Others die within the ensuing 6 months. Late deaths often result from poorly controlled infection, immunosuppression, complications of intensive care, failure of multiple organs, or the patient’s underlying disease. Case-fatality rates are similar for culture-positive and culture-negative severe sepsis. Prognostic stratification systems such as APACHE II indicate that factoring in the patient’s age, underlying condition, and various physiologic variables can yield useful estimates of the risk of dying of severe sepsis. Age and prior health status are probably the most important risk factors (Fig. 325-1). In patients with no known preexisting morbidity, the case-fatality rate remains $< 10\%$ until the fourth decade of life, after which it gradually increases to $> 35\%$ in the very elderly. Death is significantly more likely in severely septic patients with preexisting illness. Septic shock is also a strong predictor of

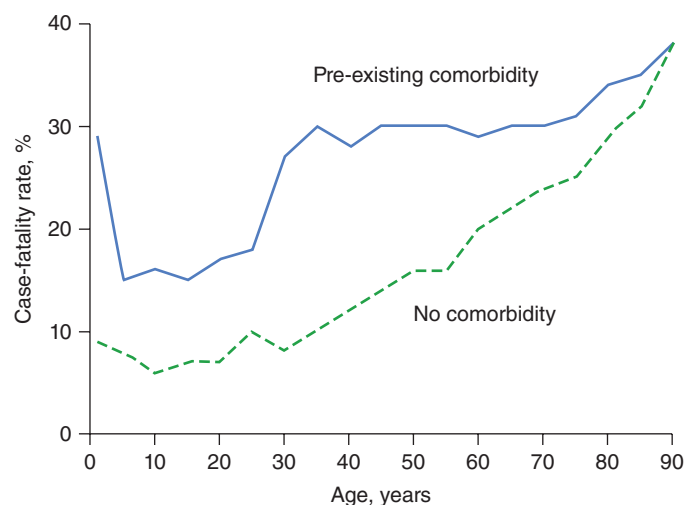


FIGURE 325-1 Influence of age and prior health status on outcome of severe sepsis. With modern therapy, fewer than 10% of previously healthy young individuals (below 35 years of age) die with severe sepsis; the case-fatality rate then increases slowly through middle and old age. The most commonly identified etiologic agents in patients who die are *Staphylococcus aureus*, *Streptococcus pyogenes*, *Streptococcus pneumoniae*, and *Neisseria meningitidis*. Individuals with preexisting comorbidities are at greater risk of dying of severe sepsis at any age. The etiologic agents in these cases are likely to be *S. aureus*, *Pseudomonas aeruginosa*, various Enterobacteriaceae, enterococci, or fungi. (Adapted from DC Angus et al: *Crit Care Med* 29:1303, 2001.)

both short- and long-term mortality. Cognitive impairment may be significant in survivors, particularly those who are elderly.

PREVENTION

Prevention offers the best opportunity to reduce morbidity and mortality from severe sepsis. In developed countries, most episodes of severe sepsis and septic shock are complications of nosocomial infections. These cases might be prevented by reducing the number of invasive procedures undertaken, by limiting the use (and duration of use) of indwelling vascular and bladder catheters, by reducing the incidence and duration of profound neutropenia (<500 neutrophils/ μL), and by more aggressively treating localized nosocomial infections. Indiscriminate use of antimicrobial agents and glucocorticoids should be avoided, and optimal infection-control measures (Chap. 168) should be used. Studies indicate that 50–70% of patients who develop nosocomial severe sepsis or septic shock have experienced a less severe stage of the septic response on at least one previous day in the hospital. Research is needed to identify patients at increased risk and to develop adjunctive agents that can modulate the septic response before organ dysfunction or hypotension occurs.

326 Cardiogenic Shock and Pulmonary Edema

Judith S. Hochman, David H. Ingbar

Cardiogenic shock and pulmonary edema are life-threatening conditions that should be treated as medical emergencies. The most common joint etiology is severe left ventricular (LV) dysfunction that leads to pulmonary congestion and/or systemic hypoperfusion (Fig. 326-1). The pathophysiology of pulmonary edema and shock is discussed in Chaps. 47e and 324, respectively.

CARDIOGENIC SHOCK

Cardiogenic shock (CS) is characterized by systemic hypoperfusion due to severe depression of the cardiac index (<2.2 [L/min]/ m^2) and sustained systolic arterial hypotension (<90 mmHg) despite an elevated filling pressure (pulmonary capillary wedge pressure [PCWP] >18 mmHg). It is associated with in-hospital mortality rates $>50\%$. The major causes of CS are listed in Table 326-1. Circulatory failure based on cardiac dysfunction may be caused by primary myocardial failure, most commonly secondary to acute myocardial infarction (MI) (Chap. 295), and less frequently by cardiomyopathy or myocarditis (Chap. 287), cardiac tamponade (Chap. 288), or critical valvular heart disease (Chap. 283).

Incidence The rate of CS complicating acute MI was 20% in the 1960s, stayed at $\sim 8\%$ for >20 years, but decreased to 5–7% in the first decade of this millennium largely due to increasing use of early reperfusion therapy for acute MI. Shock is more common with ST elevation MI (STEMI) than with non-ST elevation MI (Chap. 295).

LV failure accounts for $\sim 80\%$ of cases of CS complicating acute MI. Acute severe mitral regurgitation (MR), ventricular septal rupture (VSR), predominant right ventricular (RV) failure, and free wall rupture or tamponade account for the remainder.

Pathophysiology CS is characterized by a vicious circle in which depression of myocardial contractility, usually due to ischemia, results in reduced cardiac output and arterial blood pressure (BP), which result in hypoperfusion of the myocardium and further ischemia and depression of cardiac output (Fig. 326-1). Systolic myocardial dysfunction reduces stroke volume and, together with diastolic dysfunction, leads to elevated LV end-diastolic pressure and PCWP as well as to pulmonary congestion. Reduced coronary perfusion leads to worsening ischemia and progressive myocardial dysfunction and a rapid

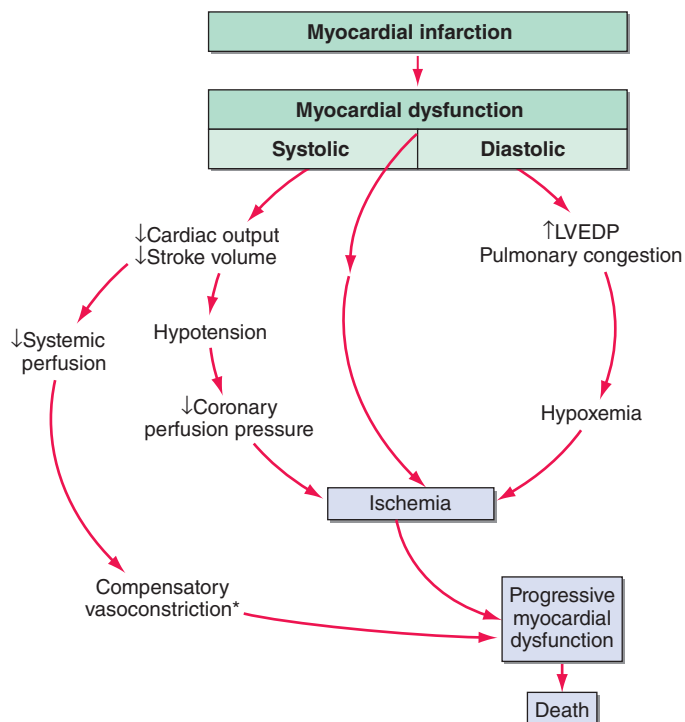


FIGURE 326-1 Pathophysiology of cardiogenic shock. Systolic and diastolic myocardial dysfunction results in a reduction in cardiac output and often pulmonary congestion. Systemic and coronary hypoperfusion occur, resulting in progressive ischemia. Although a number of compensatory mechanisms are activated in an attempt to support the circulation, these compensatory mechanisms may become maladaptive and produce a worsening of hemodynamics. *Release of inflammatory cytokines after myocardial infarction may lead to inducible nitric oxide expression, excess nitric oxide, and inappropriate vasodilation. This causes further reduction in systemic and coronary perfusion. A vicious spiral of progressive myocardial dysfunction occurs that ultimately results in death if it is not interrupted. LVEDP, left ventricular end-diastolic pressure. (From SM Hollenberg et al: *Ann Intern Med* 131:47, 1999.)

downward spiral, which, if uninterrupted, is often fatal. A systemic inflammatory response syndrome may accompany large infarctions and shock. Inflammatory cytokines, inducible nitric oxide synthase, and excess nitric oxide and peroxynitrite may contribute to the genesis of CS as they do to that of other forms of shock (Chap. 324). Lactic acidosis and hypoxemia from CS contribute to the vicious circle by worsening myocardial ischemia and hypotension. Severe acidosis reduces the efficacy of endogenous and exogenously administered catecholamines. Refractory sustained ventricular or atrial tachyarrhythmias can cause or exacerbate CS.

Patient Profile Older age, female sex, prior MI, diabetes, anterior MI location, and extensive coronary artery stenoses are associated with an increased risk of CS complicating MI. Shock associated with a first inferior MI should prompt a search for a mechanical cause. CS may rarely occur in the absence of significant stenosis, as seen in LV apical ballooning/Takotsubo's cardiomyopathy.

Timing Shock is present on admission in only one-quarter of patients who develop CS complicating MI; one-quarter develop it rapidly thereafter, within 6 h of MI onset. Another quarter develop shock later on the first day. Subsequent onset of CS may be due to reinfarction, marked infarct expansion, or a mechanical complication.

Diagnosis Due to the unstable condition of these patients, supportive therapy must be initiated simultaneously with diagnostic evaluation (Fig. 326-2). A focused history and physical examination should be performed, blood specimens sent to the laboratory, and an electrocardiogram (ECG) and chest x-ray obtained.

TABLE 326-1 ETIOLOGIES OF CARDIOGENIC SHOCK (CS)^a AND CARDIOGENIC PULMONARY EDEMA

Etiologies of Cardiogenic Shock or Pulmonary Edema
Acute myocardial infarction/ischemia
LV failure
Ventricular septal rupture
Papillary muscle/chordal rupture–severe MR
Ventricular free wall rupture with subacute tamponade
Other conditions complicating large MIs
Hemorrhage
Infection
Excess negative inotropic or vasodilator medications
Prior valvular heart disease
Hyperglycemia/ketoacidosis
Post-cardiac arrest
Post-cardiotomy
Refractory sustained tachyarrhythmias
Acute fulminant myocarditis
End-stage cardiomyopathy
LV apical ballooning
Takotsubo's cardiomyopathy
Hypertrophic cardiomyopathy with severe outflow obstruction
Aortic dissection with aortic insufficiency or tamponade
Severe valvular heart disease
Critical aortic or mitral stenosis
Acute severe aortic regurgitation or mitral regurgitation
Toxic/metabolic
β blocker or calcium channel antagonist overdose
Other Etiologies of Cardiogenic Shock^b
RV failure due to:
Acute myocardial infarction
Acute cor pulmonale
Refractory sustained bradyarrhythmias
Pericardial tamponade
Toxic/metabolic
Severe acidosis, severe hypoxemia

^aThe etiologies of CS are listed. Most of these can cause pulmonary edema instead of shock or pulmonary edema with CS. ^bThese cause CS but not pulmonary edema.

Abbreviations: LV, left ventricular; MI, myocardial infarction; MR, mitral regurgitation; RV, right ventricular; VSR, ventricular septal rupture.

Echocardiography is an invaluable diagnostic tool in patients with suspected CS.

CLINICAL FINDINGS Most patients have dyspnea and appear pale, apprehensive, and diaphoretic, and mental status may be altered. The pulse is typically weak and rapid, often in the range of 90–110 beats/min, or severe bradycardia due to high-grade heart block may be present. Systolic BP is reduced (<90 mmHg or ≥30 mmHg below baseline) with a narrow pulse pressure (<30 mmHg), but occasionally BP may be maintained by very high systemic vascular resistance. Tachypnea, Cheyne-Stokes respirations, and jugular venous distention may be present. There is typically a weak apical pulse and soft S₁, and an S₃ gallop may be audible. Acute, severe MR and VSR usually are associated with characteristic systolic murmurs (**Chap. 295**). Rales are audible in most patients with LV failure. Oliguria is common.

LABORATORY FINDINGS The white blood cell count is typically elevated with a left shift. Renal function is initially unchanged, but blood urea nitrogen and creatinine rise progressively. Hepatic transaminases may be markedly elevated due to liver hypoperfusion. The lactic acid level is elevated. Arterial blood gases usually demonstrate hypoxemia and anion gap metabolic acidosis, which may be compensated by respiratory alkalosis. Cardiac markers, creatine phosphokinase and its MB fraction, and troponins I and T are typically markedly elevated.

ELECTROCARDIOGRAM In CS due to acute MI with LV failure, Q waves and/or >2-mm ST elevation in multiple leads or left bundle branch block are usually present. More than one-half of all infarcts associated with shock are anterior. Global ischemia due to severe left main stenosis usually is accompanied by severe (e.g., >3 mm) ST depressions in multiple leads.

CHEST ROENTGENOGRAM The chest x-ray typically shows pulmonary vascular congestion and often pulmonary edema, but these findings may be absent in up to a third of patients. The heart size is usually normal when CS results from a first MI but is enlarged when it occurs in a patient with a previous MI.

ECHOCARDIOGRAM A two-dimensional echocardiogram with color-flow Doppler (**Chap. 270e**) should be obtained promptly in patients with suspected CS to help define its etiology. Doppler mapping demonstrates a left-to-right shunt in patients with VSR and the severity of MR when the latter is present. Proximal aortic dissection with aortic regurgitation or tamponade may be visualized, or evidence for pulmonary embolism may be obtained (**Chap. 300**).

PULMONARY ARTERY CATHETERIZATION The use of pulmonary artery (Swan-Ganz) catheters in patients with established or suspected CS is controversial (**Chaps. 272 and 321**). Their use is generally recommended for measurement of filling pressures and cardiac output to confirm the diagnosis and to optimize the use of IV fluids, inotropic agents, and vasopressors in persistent shock (**Table 326-2**). O₂ saturation measurement from right atrial, RV, and pulmonary arterial blood samples can rule out a left-to-right shunt. In CS, low mixed venous O₂ saturations and elevated arteriovenous (AV) O₂ differences reflect low cardiac index and high fractional O₂ extraction. However, when sepsis accompanies CS, AV O₂ differences may not be elevated (**Chap. 324**). The PCWP is elevated. Use of sympathomimetic amines may return these measurements and the systemic BP to normal. Systemic vascular resistance may be low, normal, or elevated in CS. Equalization of right- and left-sided filling pressures (right atrial and PCWP) suggests cardiac tamponade as the cause of CS (**Chap. 288**).

LEFT HEART CATHETERIZATION AND CORONARY ANGIOGRAPHY Measurement of LV pressure and definition of the coronary anatomy provide useful information and are indicated in most patients with CS complicating MI. Cardiac catheterization should be performed when there is a plan and capability for immediate coronary intervention (see below) or when a definitive diagnosis has not been made by other tests.

TREATMENT ACUTE MYOCARDIAL INFARCTION

GENERAL MEASURES

(Fig. 326-2) In addition to the usual treatment of acute MI (**Chap. 295**), initial therapy is aimed at maintaining adequate systemic and coronary perfusion by raising systemic BP with vasopressors and adjusting volume status to a level that ensures optimum LV filling pressure. There is interpatient variability, but the values that generally are associated with adequate perfusion are systolic BP ~90 mmHg or mean BP >60 mmHg and PCWP >20 mmHg. Hypoxemia and acidosis must be corrected; most patients require ventilatory support (see “Pulmonary Edema,” below). Negative inotropic agents should be discontinued and the doses of renally cleared medications adjusted. Hyperglycemia should be controlled with insulin. Bradyarrhythmias may require transvenous pacing. Recurrent ventricular tachycardia or rapid atrial fibrillation may require immediate treatment (**Chap. 276**).

VASOPRESSORS

Various IV drugs may be used to augment BP and cardiac output in patients with CS. All have important disadvantages, and none has been shown to change the outcome in patients with established shock. *Norepinephrine* is a potent vasoconstrictor and inotropic stimulant that is useful for patients with CS. As first line of therapy norepinephrine was associated with fewer adverse events, including arrhythmias, compared to a dopamine randomized trial of patients

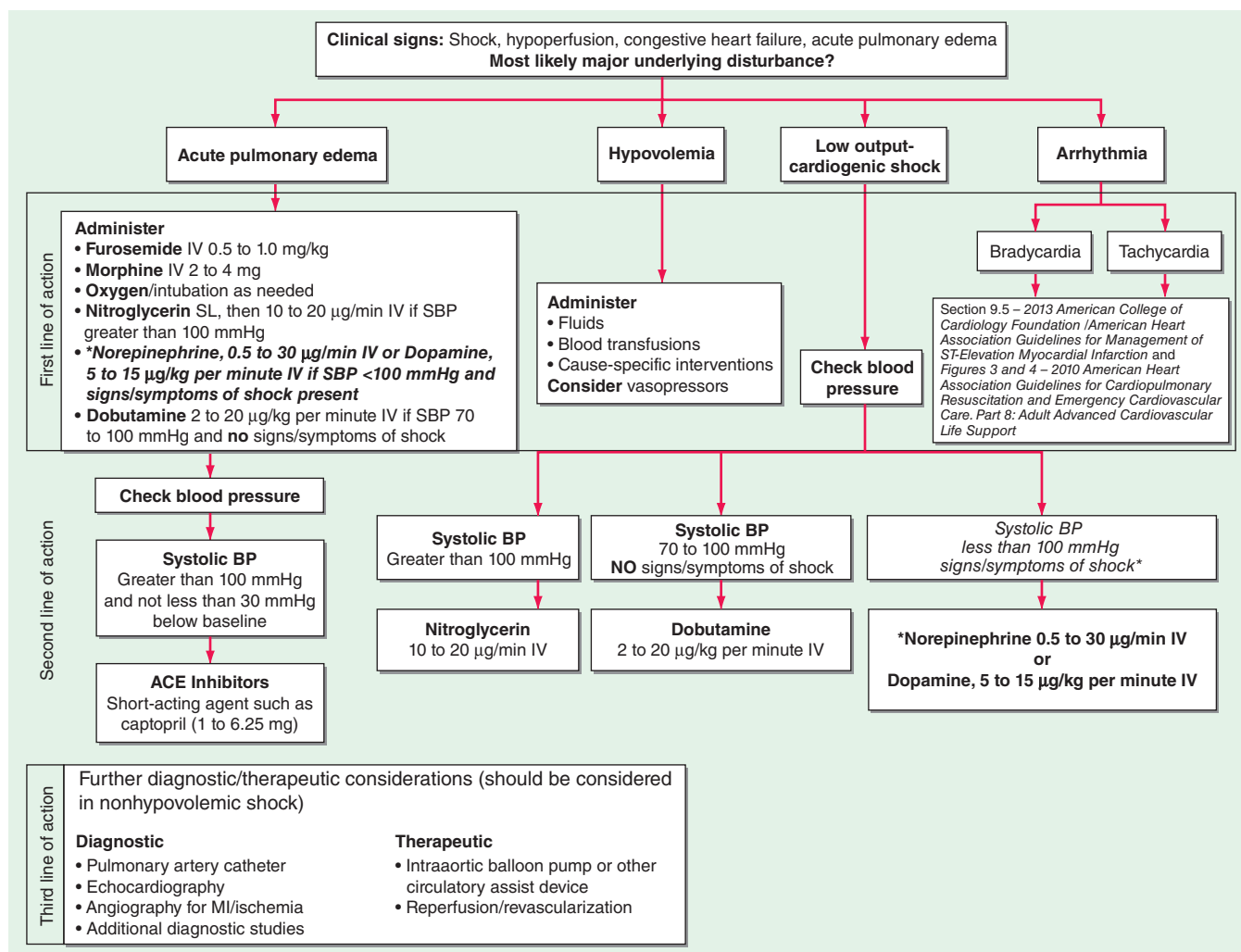


FIGURE 326-2 The emergency management of patients with cardiogenic shock, acute pulmonary edema, or both is outlined.

*Furosemide: <0.5 mg/kg for new-onset acute pulmonary edema without hypervolemia; 1 mg/kg for acute on chronic volume overload, renal insufficiency. †For management of bradycardia and tachycardia, see [Chaps. 274 and 276](#). Additional information can also be found in Section 9.5 of the 2013 American College of Cardiology Foundation/American Heart Association Guidelines for Management of ST-Elevation Myocardial Infarction and Figures 3 and 4 of the 2010 American Heart Association Guidelines for Cardiopulmonary Resuscitation and Emergency Cardiovascular Care. Part 8: Adult Advanced Cardiovascular Life Support. *Indicates modification from published guidelines. ACE, angiotensin-converting enzyme; BP, blood pressure; MI, myocardial infarction. (Modified from *Guidelines 2000 for Cardiopulmonary Resuscitation and Emergency Cardiovascular Care. Part 7: The era of reperfusion: Section 1: Acute coronary syndromes [acute myocardial infarction]*. The American Heart Association in collaboration with the International Liaison Committee on Resuscitation. *Circulation* 102:1172, 2000.)

with several etiologies of circulatory shock. Although it did not significantly improve survival compared to dopamine, its relative safety suggests that norepinephrine is reasonable as initial vasopressor therapy. Norepinephrine should be started at a dose of 2 to 4 $\mu\text{g}/\text{min}$ and titrated upward as necessary. If systemic perfusion or systolic pressure cannot be maintained at >90 mmHg with a dose of 15 $\mu\text{g}/\text{min}$, it is unlikely that a further increase will be beneficial.

Dopamine has varying hemodynamic effects based on the dose; at low doses ($\leq 2 \mu\text{g}/\text{kg}$ per min), it dilates the renal vascular bed, although its outcome benefits at this low dose have not been demonstrated conclusively; at moderate doses (2–10 $\mu\text{g}/\text{kg}$ per min), it has positive chronotropic and inotropic effects as a consequence of β -adrenergic receptor stimulation. At higher doses, a vasoconstrictor effect results from α -receptor stimulation. It is started at an infusion rate of 2–5 $\mu\text{g}/\text{kg}$ per min, and the dose is increased every 2–5 min to a maximum of 20–50 $\mu\text{g}/\text{kg}$ per min. *Dobutamine* is a synthetic sympathomimetic amine with positive inotropic action and minimal positive chronotropic activity at low doses (2.5 $\mu\text{g}/\text{kg}$ per min) but moderate chronotropic activity at higher doses. Although the usual dose is up to 10 $\mu\text{g}/\text{kg}$ per min, its vasodilating activity precludes its use when a vasoconstrictor effect is required.

MECHANICAL CIRCULATORY SUPPORT

Circulatory assist devices can be placed percutaneously or surgically and can be used to support the left, right, or both ventricles. Venoarterial extracorporeal membrane oxygenation (VA ECMO, a pump in combination with an oxygenator) may be used when respiratory failure accompanies biventricular failure. Temporary percutaneous devices can be used as a bridge to surgically implanted devices in community hospital settings or when neurologic status is uncertain. The most commonly used device is an intraaortic balloon pump (IABP), which is inserted into the aorta via the femoral artery and provides temporary hemodynamic support. However, routine IABP use in conjunction with early revascularization (predominantly with percutaneous coronary intervention [PCI]) did not reduce 30-day mortality in the IABP-SHOCK II trial. Although other percutaneous devices, including VA ECMO, result in better hemodynamic support compared to IABP, the effects on clinical outcomes are unknown. Surgically implanted devices can support the circulation as bridging therapy for cardiac transplant candidates or as destination therapy ([Chap. 281](#)). Assist devices should be used selectively in suitable patients in consultation with advanced heart failure specialists.

TABLE 326-2 HEMODYNAMIC PATTERNS^a

	RA, mmHg	RVS, mmHg	RVD, mmHg	PAS, mmHg	PAD, mmHg	PCW, mmHg	CI, (L/min)/m ²	SVR, (dyn · s)/cm ⁵
Normal values	<6	<25	0–12	<25	0–12	<6–12	≥2.5	(800–1600)
MI without pulmonary edema ^b	–	–	–	–	–	~13 (5–18)	~2.7 (2.2–4.3)	–
Pulmonary edema	↔↑	↔↑	↔↑	↑	↑	↑	↔↓	↑
Cardiogenic shock								
LV failure	↔↑	↔↑	↔↑	↔↑	↑	↑	↓	↔↑
RV failure ^c	↑	↓↔↑ ^d	↑	↓↔↑ ^d	↔↓↑ ^d	↓↔↑ ^d	↓	↑
Cardiac tamponade	↑	↔↑	↑	↔↑	↔↑	↔↑	↓	↑
Acute mitral regurgitation	↔↑	↑	↔↑	↑	↑	↑	↔↓	↔↑
Ventricular septal rupture	↑	↔↑	↑	↔↑	↔↑	↔↑	↑PBF ↓SBF	↔↑
Hypovolemic shock	↓	↔↓	↔↓	↓	↓	↓	↓	↑
Septic shock	↓	↔↓	↔↓	↓	↓	↓	↑	↓

^aThere is significant patient-to-patient variation. Pressure may be normalized if cardiac output is low. ^bForrester et al classified nonreperfused MI patients into four hemodynamic subsets. (From JS Forrester et al: N Engl J Med 295:1356, 1976.) PCW pressure and CI in clinically stable subset 1 patients are shown. Values in parentheses represent range. ^c“Isolated” or predominant RV failure. ^dPCW and pulmonary artery pressures may rise in RV failure after volume loading due to RV dilation and right-to-left shift of the interventricular septum, resulting in impaired LV filling. When biventricular failure is present, the patterns are similar to those shown for LV failure.

Abbreviations: CI, cardiac index; MI, myocardial infarction; P/SBF, pulmonary/systemic blood flow; PAS/D, pulmonary artery systolic/diastolic; PCW, pulmonary capillary wedge; RA, right atrium; RVS/D, right ventricular systolic/diastolic; SVR, systemic vascular resistance.

Source: Table prepared with the assistance of Krishnan Ramanathan, MD.

REPERFUSION-REVASCULARIZATION

The rapid establishment of blood flow in the infarct-related artery is essential in the management of CS and forms the centerpiece of management. The randomized SHOCK Trial demonstrated that 132 lives were saved per 1000 patients treated with early revascularization with PCI or coronary artery bypass graft (CABG) compared with initial medical therapy including IABP with fibrinolytics followed by delayed revascularization. The benefit is seen across the risk strata and is sustained up to 11 years after an MI. Early revascularization with PCI or CABG is recommended in candidates suitable for aggressive care.

Prognosis Within this high-risk condition, there is a wide range of expected death rates based on age, severity of hemodynamic abnormalities, severity of the clinical manifestations of hypoperfusion, and the performance of early revascularization.

SHOCK SECONDARY TO RIGHT VENTRICULAR INFARCTION

Although transient hypotension is common in patients with RV infarction and inferior MI (Chap. 295), persistent CS due to RV failure accounts for only 3% of CS complicating MI. The salient features of RV shock are absence of pulmonary congestion, high right atrial pressure (which may be seen only after volume loading), RV dilation and dysfunction, only mildly or moderately depressed LV function, and predominance of single-vessel proximal right coronary artery occlusion. Management includes IV fluid administration to optimize right atrial pressure (10–15 mmHg); avoidance of excess fluids, which cause a shift of the interventricular septum into the LV; sympathomimetic amines; the early reestablishment of infarct-artery flow; and assist devices.

MITRAL REGURGITATION

(See also Chap. 295) Acute severe MR due to papillary muscle dysfunction and/or rupture may complicate MI and result in CS and/or pulmonary edema. This complication most often occurs on the first day, with a second peak several days later. The diagnosis is confirmed by echo-Doppler. Rapid stabilization with IABP is recommended, with administration of dobutamine as needed to raise cardiac output.

Reducing the load against which the LV pumps (afterload) reduces the volume of regurgitant flow of blood into the left atrium. Mitral valve surgery is the definitive therapy and should be performed early in the course in suitable candidates.

VENTRICULAR SEPTAL RUPTURE

(See also Chap. 295) Echo-Doppler demonstrates shunting of blood from the left to the right ventricle and may visualize the opening in the interventricular septum. Timing and management are similar to those for MR with IABP support and surgical correction for suitable candidates.

FREE WALL RUPTURE

Myocardial rupture is a dramatic complication of STEMI that is most likely to occur during the first week after the onset of symptoms; its frequency increases with the age of the patient. The clinical presentation typically is a sudden loss of pulse, blood pressure, and consciousness but sinus rhythm on ECG (pulseless electrical activity) due to cardiac tamponade (Chap. 288). Free wall rupture may also result in CS due to subacute tamponade when the pericardium temporarily seals the rupture sites. Definitive surgical repair is required.

ACUTE FULMINANT MYOCARDITIS

(See also Chap. 287) Myocarditis can mimic acute MI with ST deviation or bundle branch block on the ECG and marked elevation of cardiac markers. Acute myocarditis causes CS in a small proportion of cases. These patients are typically younger than those with CS due to acute MI and often do not have typical ischemic chest pain. Echocardiography usually shows global LV dysfunction. Initial management is the same as for CS complicating acute MI (Fig. 326-2) but does not involve coronary revascularization. Endomyocardial biopsy is recommended to determine the diagnosis and need for immunosuppressives for entities such as giant cell myocarditis. Refractory CS can be managed with assist devices with or without ECMO.

PULMONARY EDEMA

The etiologies and pathophysiology of pulmonary edema are discussed in Chap. 47e.

Diagnosis Acute pulmonary edema usually presents with the rapid onset of dyspnea at rest, tachypnea, tachycardia, and severe hypoxemia. Crackles and wheezing due to alveolar flooding and airway compression from peribronchial cuffing may be audible. Release of endogenous catecholamines often causes hypertension.

It is often difficult to distinguish between cardiogenic and noncardiogenic causes of acute pulmonary edema. *Echocardiography* may identify systolic and diastolic ventricular dysfunction and valvular lesions. Electrocardiographic ST elevation and evolving Q waves are usually diagnostic of acute MI and should prompt immediate institution of MI protocols and coronary artery reperfusion therapy (**Chap. 295**). Brain natriuretic peptide levels, when substantially elevated, support heart failure as the etiology of acute dyspnea with pulmonary edema (**Chap. 279**).

The use of a *Swan-Ganz catheter* permits measurement of PCWP and helps differentiate high-pressure (cardiogenic) from normal-pressure (noncardiogenic) causes of pulmonary edema. Pulmonary artery catheterization is indicated when the etiology of the pulmonary edema is uncertain, when edema is refractory to therapy, or when it is accompanied by hypotension. Data derived from use of a catheter often alter the treatment plan, but no impact on mortality rates has been demonstrated.

TREATMENT PULMONARY EDEMA

The treatment of pulmonary edema depends on the specific etiology. As an acute, life-threatening condition, a number of measures must be applied immediately to support the circulation, gas exchange, and lung mechanics. Simultaneously, conditions that frequently complicate pulmonary edema, such as infection, acidemia, anemia, and acute kidney dysfunction, must be corrected.

SUPPORT OF OXYGENATION AND VENTILATION

Patients with acute cardiogenic pulmonary edema generally have an identifiable cause of acute LV failure—such as arrhythmia, ischemia/infarction, or myocardial decompensation (**Chap. 279**)—that may be rapidly treated, with improvement in gas exchange. In contrast, noncardiogenic edema usually resolves much less quickly, and most patients require mechanical ventilation.

Oxygen Therapy Support of oxygenation is essential to ensure adequate O_2 delivery to peripheral tissues, including the heart.

Positive-Pressure Ventilation Pulmonary edema increases the work of breathing and the O_2 requirements of this work, imposing a significant physiologic stress on the heart. When oxygenation or ventilation is not adequate in spite of supplemental O_2 , positive-pressure ventilation by face or nasal mask or by endotracheal intubation should be initiated. Noninvasive ventilation (**Chap. 323**) can rest the respiratory muscles, improve oxygenation and cardiac function, and reduce the need for intubation. In refractory cases, mechanical ventilation can relieve the work of breathing more completely than can noninvasive ventilation. Mechanical ventilation with positive end-expiratory pressure can have multiple beneficial effects on pulmonary edema: (1) decreases both preload and afterload, thereby improving cardiac function; (2) redistributes lung water from the intraalveolar to the extraalveolar space, where the fluid interferes less with gas exchange; and (3) increases lung volume to avoid atelectasis.

REDUCTION OF PRELOAD

In most forms of pulmonary edema, the quantity of extravascular lung water is determined by both the PCWP and the intravascular volume status.

Diuretics The “loop diuretics” furosemide, bumetanide, and torsemide are effective in most forms of pulmonary edema, even in the presence of hypoalbuminemia, hyponatremia, or hypochloremia. Furosemide is also a venodilator that rapidly reduces preload before any diuresis, and is the diuretic of choice. The initial dose of furosemide should be ≤ 0.5 mg/kg, but a higher dose (1 mg/kg) is required

in patients with renal insufficiency, chronic diuretic use, or hypervolemia or after failure of a lower dose.

Nitrates Nitroglycerin and isosorbide dinitrate act predominantly as venodilators but have coronary vasodilating properties as well. They are rapid in onset and effective when administered by a variety of routes. Sublingual nitroglycerin (0.4 mg \times 3 every 5 min) is first-line therapy for acute cardiogenic pulmonary edema. If pulmonary edema persists in the absence of hypotension, sublingual may be followed by IV nitroglycerin, commencing at 5–10 μ g/min. IV nitroprusside (0.1–5 μ g/kg per min) is a potent venous and arterial vasodilator. It is useful for patients with pulmonary edema and hypertension but is not recommended in states of reduced coronary artery perfusion. It requires close monitoring and titration using an arterial catheter for continuous BP measurement.

Morphine Given in 2- to 4-mg IV boluses, morphine is a transient venodilator that reduces preload while relieving dyspnea and anxiety. These effects can diminish stress, catecholamine levels, tachycardia, and ventricular afterload in patients with pulmonary edema and systemic hypertension.

Angiotensin-Converting Enzyme (ACE) Inhibitors ACE inhibitors reduce both afterload and preload and are recommended for hypertensive patients. A low dose of a short-acting agent may be initiated and followed by increasing oral doses. In acute MI with heart failure, ACE inhibitors reduce short- and long-term mortality rates.

Other Preload-Reducing Agents IV recombinant brain natriuretic peptide (nesiritide) is a potent vasodilator with diuretic properties and is effective in the treatment of cardiogenic pulmonary edema. It should be reserved for refractory patients and is not recommended in the setting of ischemia or MI.

Physical Methods In nonhypotensive patients, venous return can be reduced by use of the sitting position with the legs dangling along the side of the bed.

Inotropic and Inodilator Drugs The sympathomimetic amines dopamine and dobutamine (see above) are potent inotropic agents. The bipyridine phosphodiesterase-3 inhibitors (inodilators), such as milrinone (50 μ g/kg followed by 0.25–0.75 μ g/kg per min), stimulate myocardial contractility while promoting peripheral and pulmonary vasodilation. Such agents are indicated in patients with cardiogenic pulmonary edema and severe LV dysfunction.

Digitalis Glycosides Once a mainstay of treatment because of their positive inotropic action (**Chap. 279**), digitalis glycosides are rarely used at present. However, they may be useful for control of ventricular rate in patients with rapid atrial fibrillation or flutter and LV dysfunction, because they do not have the negative inotropic effects of other drugs that inhibit atrioventricular nodal conduction.

Intraaortic Balloon Counterpulsation IABP or other LV-assist devices (**Chap. 281**) may help relieve cardiogenic pulmonary edema and are indicated when refractory pulmonary edema results from the etiologies discussed in the CS section, especially in preparation for surgical repair.

Treatment of Tachyarrhythmias and Atrial-Ventricular Resynchronization (See also **Chap. 277**) Sinus tachycardia or atrial fibrillation can result from elevated left atrial pressure and sympathetic stimulation. Tachycardia itself can limit LV filling time and raise left atrial pressure further. Although relief of pulmonary congestion will slow the sinus rate or ventricular response in atrial fibrillation, a primary tachyarrhythmia may require cardioversion. In patients with reduced LV function and without atrial contraction or with lack of synchronized atrioventricular contraction, placement of an atrioventricular sequential pacemaker should be considered (**Chap. 274**).

Stimulation of Alveolar Fluid Clearance A variety of drugs can stimulate alveolar epithelial ion transport and upregulate the clearance of alveolar solute and water, but this strategy has not been proven beneficial in clinical trials thus far.

Risk of Iatrogenic Cardiogenic Shock In the treatment of pulmonary edema, vasodilators lower BP, and their use, particularly in combination, may lead to hypotension, coronary artery hypoperfusion, and shock (Fig. 326-1). In general, patients with a *hypertensive* response to pulmonary edema tolerate and benefit from these medications. In normotensive patients, low doses of single agents should be instituted sequentially, as needed.

Acute Coronary Syndromes (See also Chap. 295) Acute STEMI complicated by pulmonary edema is associated with in-hospital mortality rates of 20–40%. After immediate stabilization, coronary artery blood flow must be reestablished rapidly. When available, primary PCI is preferable; alternatively, a fibrinolytic agent should be administered. Early coronary angiography and revascularization by PCI or CABG also are indicated for patients with non-ST elevation acute coronary syndrome. Assist devices may be used selectively as noted for refractory pulmonary edema.

Extracorporeal Membrane Oxygenation For patients with acute, severe noncardiogenic edema with a potential rapidly reversible cause, ECMO may be considered as a temporizing supportive measure to achieve adequate gas exchange. Usually venovenous ECMO is used in this setting.

Unusual Types of Edema Specific etiologies of pulmonary edema may require particular therapy. Reexpansion pulmonary edema can develop after removal of longstanding pleural space air or fluid. These patients may develop hypotension or oliguria resulting from rapid fluid shifts into the lung. Diuretics and preload reduction are contraindicated, and intravascular volume repletion often is needed while supporting oxygenation and gas exchange.

High-altitude pulmonary edema often can be prevented by use of dexamethasone, calcium channel–blocking drugs, or long-acting inhaled β_2 -adrenergic agonists. Treatment includes descent from altitude, bed rest, oxygen, and, if feasible, inhaled nitric oxide; nifedipine may also be effective.

For pulmonary edema resulting from upper airway obstruction, recognition of the obstructing cause is key, because treatment then is to relieve or bypass the obstruction.

Biological death may be delayed by interventions, but the relevant pathophysiologic event remains the sudden and unexpected cardiac arrest. Accordingly, for statistical purposes, deaths that occur during hospitalization or within 30 days after resuscitated cardiac arrest are counted as sudden deaths.

The majority of natural deaths are caused by cardiac disorders. However, it is common for underlying heart diseases—often far advanced—to go unrecognized before the fatal event. As a result, up to two-thirds of all SCDs occur as the first clinical expression of previously undiagnosed disease or in patients with known heart disease, the extent of which suggests low individual risk. The magnitude of sudden cardiac death as a public health problem is highlighted by the estimate that ~50% of all cardiac deaths are sudden and unexpected, accounting for a total SCD burden estimated to range from <200,000 to >450,000 deaths each year in the United States. SCD is a direct consequence of cardiac arrest, which may be reversible if addressed promptly. Because resuscitation techniques and emergency rescue systems are available to respond to victims of out-of-hospital cardiac arrest, which was uniformly fatal in the past, understanding the SCD problem has practical clinical importance.

CLINICAL DEFINITION OF FORMS OF CARDIOVASCULAR COLLAPSE

Cardiovascular collapse is a general term connoting loss of sufficient cerebral blood flow to maintain consciousness due to acute dysfunction of the heart and/or peripheral vasculature. It may be caused by vasodepressor syncope (vasovagal syncope, postural hypotension with syncope, neurocardiogenic syncope; Chap. 27), a transient severe bradycardia, or cardiac arrest. The latter is distinguished from the transient forms of cardiovascular collapse in that it usually requires an active intervention to restore spontaneous blood flow. In contrast, vasodepressor syncope and other primary bradyarrhythmic syncopal events are transient and non-life-threatening, with spontaneous return of consciousness.

In the past, the most common electrical mechanism for cardiac arrest was ventricular fibrillation (VF) or pulseless sustained ventricular tachycardia (PVT). These were the initial rhythms recorded in 60–80% of cardiac arrests, with VF being the far more common of the two. Severe persistent bradyarrhythmias, asystole, and pulseless electrical activity (PEA; organized electrical activity, unusually slow, without mechanical response, formerly called electromechanical dissociation [EMD]) caused another 20–30%. Currently, asystole has emerged as the most common mechanism recorded at initial contact (45–50% of cases). PEA accounts for 20–25%, and VF is now present on initial contact in 25–35%. Undoubtedly, a significant proportion of the asystole cases began as VF and deteriorated to asystole because of long response times, but there are data suggesting an absolute reduction in VF as well. Acute low cardiac output states, having a precipitous onset, also may present clinically as a cardiac arrest. These hemodynamic causes include massive acute pulmonary emboli, internal blood loss from a ruptured aortic aneurysm, intense anaphylaxis, and cardiac rupture with tamponade after myocardial infarction (MI).

ETIOLOGY, INITIATING EVENTS, AND CLINICAL EPIDEMIOLOGY

Clinical, epidemiologic, and pathologic studies have provided information on the underlying *structural substrates* in victims of SCD and identified subgroups at high risk for SCD. In addition, studies of clinical physiology have begun to identify *transient functional factors* that may convert a long-standing underlying structural abnormality from a stable to an unstable state, leading to the onset of cardiac arrest (Table 327-2).

Cardiac disorders constitute the most common causes of sudden natural death. After an initial peak incidence of sudden death between birth and 6 months of age (sudden infant death syndrome [SIDS]), the incidence of sudden death declines sharply and remains low through childhood and adolescence. Among adolescents and young adults, the incidence of SCD is approximately 1 per 100,000 population per year. The incidence begins to increase in adults over age 30 years, reaching a second peak in the age range of 45–75 years, when it approximates 1–2 per 1000 per year among the unselected adult population. Increasing

327 Cardiovascular Collapse, Cardiac Arrest, and Sudden Cardiac Death

Robert J. Myerburg, Agustin Castellanos

OVERVIEW AND DEFINITIONS

Sudden cardiac death (SCD) is defined as *natural death due to cardiac causes* in a person who may or may not have previously recognized heart disease but in whom the time and mode of death are *unexpected*. The term “sudden,” in the context of SCD, is defined for most clinical and epidemiologic purposes as *1 h or less* between a change in clinical status heralding the onset of the terminal clinical event and the cardiac arrest itself. One exception is unwitnessed deaths, in which pathologists may expand the temporal definition to 24 h after the victim was last seen to be alive and stable.

Another exception is the variable interval between cardiac arrest and biological death that results from community-based interventions, following which victims may remain biologically alive for days or even weeks after a cardiac arrest that has resulted in irreversible central nervous system damage. Confusion in terms can be avoided by adhering strictly to definitions of cardiovascular collapse, cardiac arrest, and death (Table 327-1). Although cardiac arrest is often potentially reversible by appropriate and timely interventions, death is biologically, legally, and literally an absolute and irreversible event.

TABLE 327-1 DISTINCTION BETWEEN CARDIOVASCULAR COLLAPSE, CARDIAC ARREST, AND DEATH

Term	Definition	Qualifiers	Mechanisms
Cardiovascular collapse	Sudden loss of effective blood flow due to cardiac and/or peripheral vascular factors that may reverse spontaneously (e.g., neurocardiogenic syncope, vasovagal syncope) or require interventions (e.g., cardiac arrest)	Nonspecific term; includes cardiac arrest and its consequences and transient events that characteristically revert spontaneously	Same as "Cardiac Arrest," plus vasodepressor syncope or other causes of transient loss of blood flow
Cardiac arrest	Abrupt cessation of cardiac mechanical function, which may be reversible by a prompt intervention but will lead to death in its absence	Rare spontaneous reversions; likelihood of successful intervention relates to mechanism of arrest, clinical setting, and prompt return of circulation	Ventricular fibrillation, ventricular tachycardia, asystole, bradycardia, pulseless electrical activity, noncardiac mechanical factors (e.g., pulmonary embolism)
Sudden cardiac death	Sudden, irreversible cessation of all biological functions	None	

Source: Modified from RJ Myerburg, A Castellanos: Cardiac arrest and sudden cardiac death, in P Libby et al (eds): *Braunwald's Heart Disease*, 8th ed. Philadelphia, Saunders, 2008.

TABLE 327-2 CARDIAC ARREST AND SUDDEN CARDIAC DEATH**Structural Substrates and Causes**

- I. Coronary heart disease
 - A. Coronary artery abnormalities
 1. Chronic atherosclerotic lesions
 2. Active lesions (plaque fissuring, platelet aggregation, acute thrombosis)
 3. Anomalous coronary artery anatomy
 - B. Myocardial infarction
 1. Healed
 2. Acute
- II. Myocardial hypertrophy
 - A. Secondary
 - B. Hypertrophic cardiomyopathy
 1. Obstructive
 2. Nonobstructive
- III. Dilated cardiomyopathy—primary muscle disease
- IV. Inflammatory and infiltrative disorders
 - A. Myocarditis
 - B. Noninfectious inflammatory diseases
 - C. Infiltrative diseases
- V. Valvular heart disease
- VI. Electrophysiologic abnormalities, structural
 - A. Anomalous pathways in Wolff-Parkinson-White syndrome
 - B. Conducting system disease
- VII. Inherited disorders associated with electrophysiologic abnormalities (congenital long QT syndromes, right ventricular dysplasia, Brugada syndrome, catecholaminergic polymorphic ventricular tachycardia, etc.)

Triggers for Expression of Cardiac Arrest

- I. Alterations of coronary blood flow
 - A. Transient ischemia
 - B. Reperfusion after ischemia
- II. Low cardiac output states
 - A. Heart failure
 1. Chronic
 2. Acute decompensation
 - B. Shock
- III. Systemic metabolic abnormalities
 - A. Electrolyte imbalance (e.g., hypokalemia)
 - B. Hypoxemia, acidosis
- IV. Neurologic disturbances
 - A. Autonomic fluctuations: central, neural, humoral
 - B. Receptor function
- V. Toxic responses
 - A. Proarrhythmic drug effects
 - B. Cardiac toxins (e.g., cocaine, digitalis intoxication)
 - C. Drug interactions

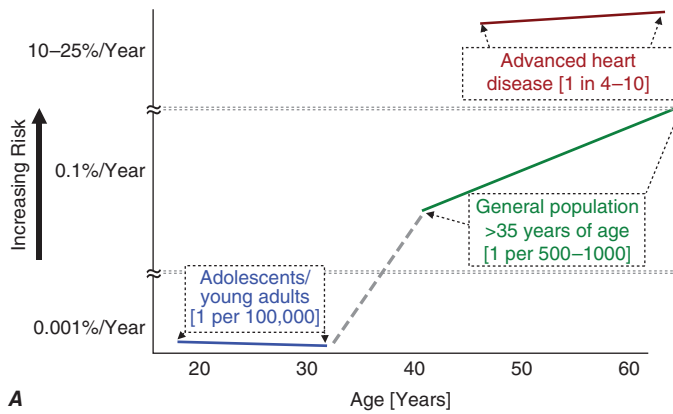
age within this range is associated with increasing risk for sudden cardiac death (Fig. 327-1A). From 1 to 13 years of age, only one of five sudden natural deaths is due to cardiac causes. Between 14 and 21 years of age, the proportion increases to 30%, and it rises to 88% in the middle-aged and elderly.

Young and middle-aged men and women have different susceptibilities to SCD, but the sex differences decrease and ultimately disappear with advancing age. The difference in risk for SCD parallels the differences in age-related risks for other manifestations of coronary heart disease (CHD) between men and women. As the gender gap for manifestations of CHD closes in the sixth to eighth decades of life, the excess risk of SCD in males progressively narrows. Despite the lower incidence among younger women, coronary risk factors such as cigarette smoking, diabetes, hyperlipidemia, and hypertension are highly influential, and SCD remains an important clinical and epidemiologic problem. The incidence of SCD among the African-American population appears to be higher than it is among the white population; the reasons remain uncertain.

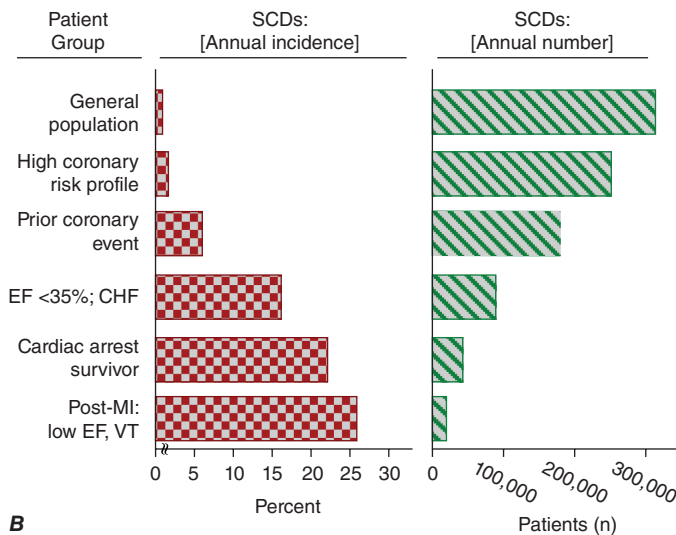
Genetic factors contribute to the risk of acquiring CHD, and a genetic basis for its expression as SCD is being explored. A genetic hypothesis for at least part of the SCD risk is supported by data suggesting a familial predisposition to SCD as a specific form of expression of CHD. A parental history of SCD as a first cardiac event increases the probability that an acute coronary event in the offspring will express similarly. In a number of less common syndromes, such as hypertrophic cardiomyopathy, congenital long QT interval syndromes, right ventricular dysplasia, and the syndrome of right bundle branch block and nonischemic ST-segment elevations (Brugada syndrome), and other more rare syndromes, there is a specific inherited risk of ventricular arrhythmias and SCD (Chap. 277).

The etiologic structural substrates and functional factors contributing to expression of the SCD syndrome are listed in Table 327-2. Worldwide, and especially in Western cultures, coronary atherosclerotic heart disease is the most common structural abnormality associated with SCD in middle-aged and older adults. Up to 80% of all SCDs in the United States are due to the consequences of coronary atherosclerosis. The nonischemic cardiomyopathies (dilated and hypertrophic, collectively; Chap. 273e) account for another 10–15% of SCDs, and all the remaining diverse etiologies cause only 5–10% of all SCDs. The inherited arrhythmia syndromes (see above and Table 327-2) are proportionally more common causes in adolescents and young adults. For some of these syndromes, such as hypertrophic cardiomyopathy (Chap. 287), the risk of SCD increases significantly after the onset of puberty.

Transient ischemia in a previously scarred or hypertrophied heart, hemodynamic and fluid and electrolyte disturbances, fluctuations in autonomic nervous system activity, and transient electrophysiologic changes caused by drugs or other chemicals (e.g., proarrhythmia) have all been implicated as mechanisms responsible for the transition from electrophysiologic stability to instability. In addition, reperfusion of ischemic myocardium may cause transient electrophysiologic instability and arrhythmias.



A



B

FIGURE 327-1 **Panel A** demonstrates age-related risk for sudden cardiac death (SCD). For the general population age 35 years and older, SCD risk is 0.1–0.2% per year (1 per 500–1000 population). Among the general population of adolescents and adults younger than age 30 years, the overall risk of SCD is 1 per 100,000 population, or 0.001% per year. The risk of SCD increases dramatically beyond age 35 years. The greatest rate of increase is between 40 and 65 years (vertical axis is discontinuous). Among patients older than 30 years of age, with advanced structural heart disease and markers of high risk for cardiac arrest, the event rate may exceed 25% per year, and age-related risk attenuates. (Modified from RJ Myerburg, A Castellanos: *Cardiac arrest and sudden cardiac death*, in P Libby et al [eds]: *Braunwald's Heart Disease*, 8th ed. Philadelphia, Saunders, 2008.) **Panel B** demonstrates the incidence of SCD in population subgroups and the relation of total number of events per year to incidence figures. Approximations of subgroup incidence figures and the related population pool from which they are derived are presented. Approximately 50% of all cardiac deaths are sudden and unexpected. The incidence bars on the left (percent/year) indicate the approximate percentage of sudden and nonsudden deaths in each of the population subgroups indicated, ranging from the lowest percentage in unselected adult populations (0.1–2% per year) to the highest percentage in patients with severe left ventricular dysfunction and heart failure (approximately 25% per year). The bars on the right indicate the total number of events per year in each of these groups with the population impact size of each of the subgroups. The highest risk categories identify the smallest number of total annual events, and the lowest incidence category accounts for the largest number of events per year. CHF, congestive heart failure; EF, ejection fraction; MI, myocardial infarction; VT, ventricular tachycardia. (After RJ Myerburg et al: *Circulation* 85:2, 1992.)

PATHOLOGY

Data from postmortem examinations of SCD victims parallel the clinical observations on the prevalence of CHD as the major structural etiologic factor. More than 80% of SCD victims have pathologic findings of CHD. The pathologic description often includes a combination of long-standing, extensive atherosclerosis of the epicardial coronary arteries and unstable coronary artery lesions, which include various permutations of eroded, fissured, or ruptured plaques; platelet aggregates; hemorrhage; and/or thrombosis. As many as 70–75% of males who die suddenly have preexisting healed MIs, whereas only 20–30% have recent acute MIs, despite the prevalence of unstable plaques and thrombi. The latter suggests transient ischemia as the mechanism of onset. Regional or global left ventricular (LV) hypertrophy often coexists with prior MIs.

PREDICTION AND PREVENTION OF CARDIAC ARREST AND SUDDEN CARDIAC DEATH

SCD accounts for approximately one-half the total number of cardiovascular deaths. As shown in Fig. 327-1B, the very-high-risk subgroups consist of more focused populations at higher risk of cardiac arrest or SCD, with better individual prediction, but the representation of such subgroups within the overall population burden of SCD is small. This is indicated by the absolute number of events (“events per year”), in contrast to the percentage per year in the subgroup. To achieve a major population impact, effective prevention of underlying diseases and the development of new epidemiologic and clinical probes that will allow better individual risk prediction by identifying specific high-risk subgroups within the large general populations are needed.

Strategies for predicting and preventing SCD are classified as primary and secondary. *Primary prevention* refers to the attempt to identify individual patients at specific risk for SCD and institute preventive strategies. *Secondary prevention* refers to measures taken to prevent recurrent cardiac arrest or death in individuals who have survived a prior cardiac arrest.

The effectiveness of the prevention strategies currently used depends on the magnitude of risk among the various population subgroups. Because the annual incidence of SCD among the unselected adult population is limited to approximately 1 per 1000 population per year (Fig. 327-1) and ~50% of all SCDs due to coronary artery disease occur as the first clinical manifestation of the disease (Fig. 327-2A), the only currently practical strategies are profiling for risk of developing CHD and risk factor control (Fig. 327-2B). The most powerful long-term risk factors include age, cigarette smoking, elevated serum cholesterol, diabetes mellitus, elevated blood pressure, LV hypertrophy, and non-specific electrocardiographic abnormalities. Markers of inflammation (e.g., levels of C-reactive protein) that may predict plaque destabilization have been added to risk classifications. The presence of multiple risk factors progressively increases incidence, but not sufficiently or specifically enough to warrant therapies targeted to potentially fatal arrhythmias (Fig. 327-1A). However, recent studies suggesting familial clustering of SCD associated with a first acute coronary syndrome offer hope that genetic markers for specific risk may be forthcoming.

After coronary artery disease has been identified in a patient, additional strategies for risk profiling become available (Fig. 327-2B), but the majority of SCDs occur among the large unselected groups rather than in the specific high-risk subgroups that become evident among populations with established disease (compare events per year with percentage per year in Fig. 327-1B). After a major cardiovascular event, such as acute MI, recent onset of heart failure, or survival after out-of-hospital cardiac arrest, the highest risk of death occurs during the initial 6–18 months after the event and then plateaus toward the baseline risk associated with the extent of underlying disease. However, many of the early deaths are nonsudden, diluting the potential benefit of strategies targeted specifically to SCD. Thus, although post-MI beta blocker therapy has an identifiable benefit for both early SCD and nonsudden mortality risk, a total mortality benefit for implantable cardioverter-defibrillator (ICD) therapy early after MI has not been observed.

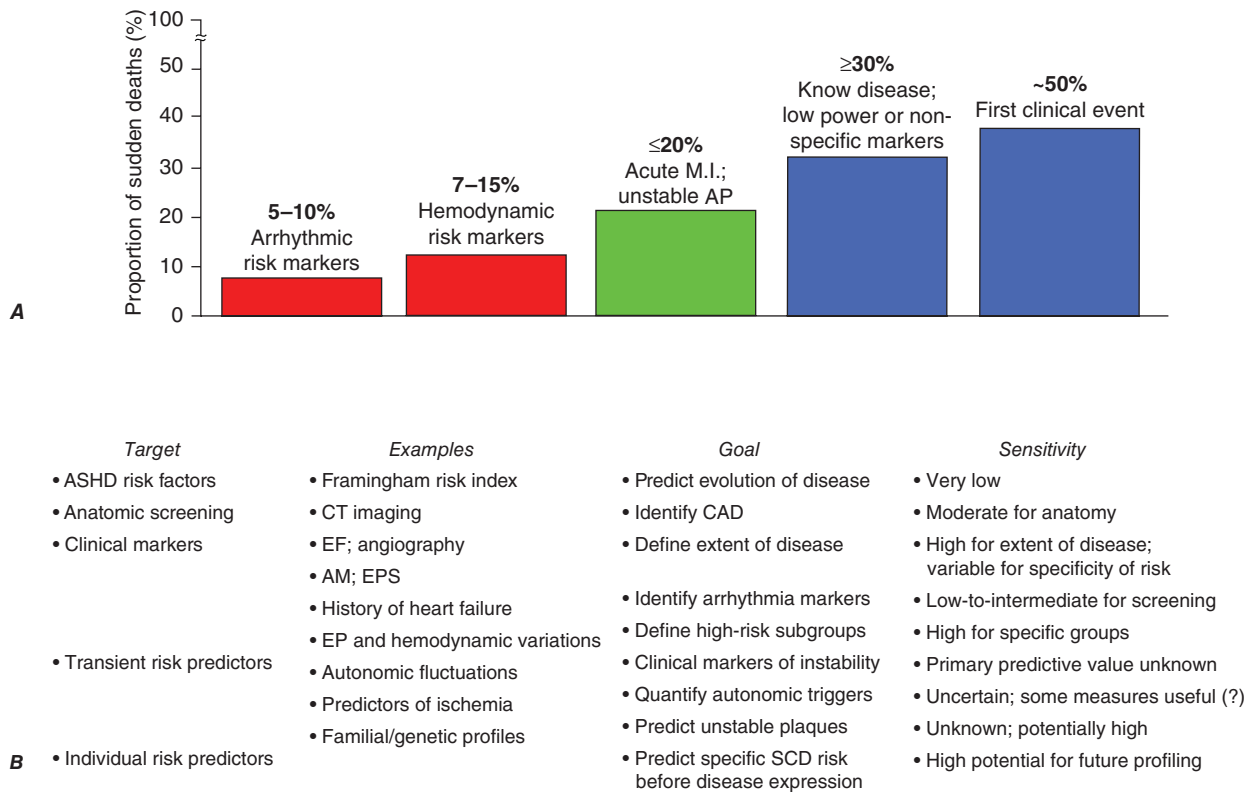


FIGURE 327-2 Population subsets, risk predictors, and distribution of sudden cardiac deaths (SCDs) according to clinical circumstances.

A. The population subset with high-risk arrhythmia markers in conjunction with low ejection fraction is a group at high risk of SCD but accounts for <10% of the total SCD burden attributable to coronary artery disease. In contrast, 50% of all SCD victims present with SCD as the first and only manifestation of underlying disease, and up to 30% have known disease but are considered relatively low risk because of the absence of high-risk markers. **B.** Profiling for individual prediction and prevention of SCD is difficult. The highest absolute numbers of events occur among the general population who may have risk factors for coronary heart disease or expressions of disease that do not predict high risk. This results in a low sensitivity for predicting and preventing SCD. New approaches that include epidemiologic modeling of transient risk factors and genetic predictors of individual patient risk offer hope for greater sensitivity in the future. AM, ambulatory monitoring; AP, angina pectoris; ASHD, arteriosclerotic heart disease; CAD, coronary artery disease; CT, computed tomography; EF, ejection fraction; EP, electrophysiologic; EPS, electrophysiologic study; MI, myocardial infarction. (Modified from RJ Myerburg: *J Cardiovasc Electrophysiol* 12:369–381, 2001.)

Among patients in the acute, convalescent, and chronic phases of MI (Chap. 295), subgroups at high absolute risk of SCD can be identified. During the acute phase, the potential risk of cardiac arrest from onset through the first 48 h used to be as high as 15%, but is now reported in the range of 2.3–4.4% because of early patient awareness of the significance of symptoms and the availability of emergency revascularization strategies. Those who survive acute-phase VF are not at continuing risk for recurrent cardiac arrest indexed to that event. During the convalescent phase after MI (3 days to ~6 weeks), an episode of sustained ventricular tachycardia (VT) or VF, which is usually associated with a large infarct, predicts a natural history mortality risk of >25% at 12 months. At least one-half of the deaths are sudden. Aggressive intervention techniques may reduce this incidence.

During the chronic phase after MI, the longer-term risk for total mortality and SCD mortality is predicted by a number of factors (Fig. 327-2B). The most important for both SCD and nonsudden death is the extent of myocardial damage sustained as a result of the acute MI. This is measured by the magnitude of reduction of the ejection fraction (EF) and/or the occurrence of heart failure. Various studies have demonstrated that ventricular arrhythmias identified by ambulatory monitoring contribute significantly to this risk, especially in patients with an EF <40%. In addition, inducibility of VT or VF during electrophysiologic testing of patients who have ambient ventricular arrhythmias (premature ventricular contractions [PVCs] and nonsustained VT) and an EF <35% is a strong predictor of SCD risk. Patients in this subgroup are now considered candidates for ICDs (see below). Risk falls off sharply with EFs >35% and the absence of

ambient arrhythmias after MI, and conversely is high with EFs <30% even without the ambient arrhythmia markers.

The cardiomyopathies (dilated and hypertrophic, Chap. 287) are the second most common category of diseases associated with risk of SCD (Table 327-2). Some risk factors have been identified, largely related to extent of disease, presence of heart failure, documented ventricular arrhythmias, and syncope thought to be due to arrhythmias. The less common causes of SCD include valvular heart disease (primarily aortic) and inflammatory and infiltrative disorders of the myocardium. The latter include viral myocarditis, sarcoidosis, and amyloidosis.

Among adolescents and young adults, rare inherited disorders such as hypertrophic cardiomyopathy, the long QT interval syndromes, right ventricular dysplasia, and the Brugada syndrome have received attention as important causes of SCD, as has acute myocarditis and other less common acquired diseases. Among the subgroup of young competitive athletes, the incidence of SCD may be higher than it is for the general adolescent and young adult population, perhaps up to 1 in 75,000–100,000. Hypertrophic cardiomyopathy (Chap. 287) is the most common cause in the United States.

Secondary prevention strategies should be applied to survivors of cardiac arrest that was not associated with an acute MI or other controllable transient risk factors, such as certain drug exposures and correctable electrolyte imbalances. Multivessel coronary artery disease and dilated cardiomyopathy, especially with markedly reduced left ventricular EF, predict a high risk of recurrence of cardiac arrest or SCD and are indications for specific interventions, such as ICDs (see

1768 below). The occurrence of otherwise unexplained syncope or documented life-threatening arrhythmias in patients with long QT syndromes or right ventricular dysplasia are also associated with increased risk of SCD.

CLINICAL CHARACTERISTICS OF CARDIAC ARREST

PRODROME, ONSET, ARREST, DEATH

SCD may be presaged by days to months of increasing angina, dyspnea, palpitations, easy fatigability, and other nonspecific complaints. However, these *prodromal symptoms* are generally predictive of any major cardiac event; they are not specific for predicting SCD.

The *onset of the clinical transition*, leading to cardiac arrest, is defined as an acute change in cardiovascular status preceding cardiac arrest by up to 1 h. When the onset is instantaneous or abrupt, the probability that the arrest is cardiac in origin is >95%. Continuous electrocardiogram (ECG) recordings fortuitously obtained at the onset of a cardiac arrest commonly demonstrate a tendency for the heart rate to increase and for advanced grades of PVCs to evolve during the minutes or hours before the event.

The probability of achieving successful resuscitation from cardiac arrest is related to the interval from onset of loss of circulation to return of spontaneous circulation (ROSC), the setting in which the event occurs, the mechanism (VF, VT, PEA, asystole), and the clinical status of the patient before the cardiac arrest. ROSC and survival rates as a result of defibrillation decrease almost linearly from the first minute to 10 min. After 4–5 min, survival rates are no better than 25–30% in out-of-hospital settings without bystander cardiopulmonary resuscitation (CPR). Those settings in which it is possible to institute prompt CPR followed by prompt defibrillation provide a better chance of a successful outcome. The outcome in intensive care units and other in-hospital environments is heavily influenced by the patient's preceding clinical status. The immediate outcome is good for cardiac arrest occurring in the intensive care unit in the presence of an acute cardiac event or transient metabolic disturbance, but survival among patients with far-advanced chronic cardiac disease or advanced noncardiac diseases (e.g., renal failure, pneumonia, sepsis, diabetes, cancer) is low and not much better in the in-hospital setting. Survival rates after unexpected cardiac arrest in unmonitored areas in a hospital do not differ from witnessed out-of-hospital arrests. Since implementation of community response systems, survival from out-of-hospital cardiac arrest has improved, although it still remains low, under most circumstances. Survival probabilities in public sites exceed those in the home environment, where the majority of cardiac arrests occur.

The success rate for initial resuscitation and survival to hospital discharge after an out-of-hospital cardiac arrest depends heavily on the mechanism of the event. When the mechanism is pulseless VT, the outcome is best; VF is the next most successful; and asystole and PEA, now the most common mechanisms, generate dismal outcome statistics. Advanced age also adversely influences the chances of successful resuscitation.

The probability of *progression to biologic death* is a function of the mechanism of cardiac arrest and the length of the delay before interventions. VF without CPR within the first 4–6 min has a poor outcome even if defibrillation is successful because of secondary brain damage; the prompt interposition of bystander CPR (basic life support; see below) improves outcome at any point along the time scale, especially when followed by early successful defibrillation. However, there are few survivors among patients who had no life support activities for the first 8 min after onset. Evaluations of deployment of automatic external defibrillators (AEDs) in communities (e.g., police vehicles, large buildings, airports, and stadiums) are beginning to generate encouraging data, but the data for home deployment has been less impressive.

Death during the hospitalization after a successfully resuscitated cardiac arrest relates closely to the severity of central nervous system injury. Anoxic encephalopathy and infections subsequent to prolonged respirator dependence account for 60% of the deaths. Another 30% occur as a consequence of low cardiac output states that fail to

respond to interventions. Recurrent arrhythmias are the least common cause of death, accounting for only 10% of in-hospital deaths.

In the setting of acute MI (Chap. 295), it is important to distinguish between primary and secondary cardiac arrests. *Primary cardiac arrests* are those that occur in the absence of hemodynamic instability, and *secondary cardiac arrests* are those that occur in patients in whom abnormal hemodynamics dominate the clinical picture before cardiac arrest. The success rate for immediate resuscitation in primary cardiac arrest during acute MI in a monitored setting should exceed 90%. In contrast, as many as 70% of patients with secondary cardiac arrest succumb immediately or during the same hospitalization.

TREATMENT CARDIAC ARREST

An individual who collapses suddenly is managed in five stages: (1) initial evaluation and basic life support if cardiac arrest is confirmed, (2) public access defibrillation (when available), (3) advanced life support, (4) postresuscitation care, and (5) long-term management. The initial response, including confirmation of loss of circulation, followed by basic life support and public access defibrillation, can be carried out by physicians, nurses, paramedical personnel, and trained laypersons.

INITIAL EVALUATION AND BASIC LIFE SUPPORT

Confirmation that a sudden collapse with loss of consciousness (LOC) is due to a cardiac arrest includes prompt observations of the state of consciousness, respiratory movements, skin color, and the presence or absence of pulses in the carotid or femoral arteries. For lay responders, the pulse check is no longer recommended because it is unreliable. As soon as a cardiac arrest is suspected, confirmed, or even considered to be impending, calling an emergency rescue system (e.g., 911) is the immediate priority. With the development of AEDs that are easily used by nonconventional emergency responders, an additional layer for response has evolved (see below).

Careful attention to the respiratory status after abrupt LOC is important. Although normal breathing or tachypnea after LOC makes cardiac arrest less likely, gasping respiratory movements may persist during a true cardiac arrest, and their presence should not deter appropriate responses. In fact, continued gasping is considered a good prognostic sign for successful outcome. It is also important to observe for severe stridor with a persistent pulse as a clue to aspiration of a foreign body or food. If this is suspected, a Heimlich maneuver (see below) may dislodge the obstructing body. A precordial blow, or "thump," delivered firmly with a clenched fist to the junction of the middle and lower thirds of the sternum may occasionally revert VT or VF, but there is concern about converting VT to VF. Therefore, it is recommended to use precordial thumps as a life support technique only when monitoring and defibrillation are available. This conservative application of the technique remains controversial.

The third action during the initial response is to clear the airway. The head is tilted back and the chin lifted so that the oropharynx can be explored to clear the airway. Dentures or foreign bodies are removed, and the Heimlich maneuver is performed if there is reason to suspect that a foreign body is lodged in the oropharynx. If respiratory arrest precipitating cardiac arrest is suspected, a second precordial thump is delivered after the airway is cleared.

Basic life support, more popularly known as CPR, is intended to maintain organ perfusion until definitive interventions can be instituted. The initial and primary element of CPR is maintenance of perfusion until spontaneous circulation can be restored. Closed chest cardiac compression maintains a pump function by sequential filling and emptying of the chambers, with competent valves maintaining forward direction of flow. The palm of one hand is placed over the lower sternum, with the heel of the other resting on the dorsum of the lower hand. The sternum is depressed, with the arms remaining straight, at a rate of 100 per minute. Sufficient force is applied to depress the sternum 4–5 cm, and relaxation is abrupt.

Until recently, providing ventilation of the lungs by mouth-to-mouth respiration was used if no specific rescue equipment was immediately available (e.g., plastic oropharyngeal airways, esophageal obturators, masked Ambu bag). However, ventilatory support during CPR has yielded to evidence that continuous chest compressions (“hands only” CPR) results in better outcomes. Compressions are interrupted only for single shocks from an AED when available, with 2 min of CPR between each single shock.

AUTOMATED EXTERNAL DEFIBRILLATION (AED)

AEDs that are easily used by nonconventional responders, such as nonparamedic firefighters, police officers, ambulance drivers, trained security guards, and minimally trained or untrained laypersons, have been developed. This advance has inserted another level of response into the cardiac arrest paradigm. A number of studies have demonstrated that AED use by nonconventional responders in strategic response systems and public access lay responders can improve cardiac arrest survival rates. The rapidity with which defibrillation/cardioversion is achieved is an important element for successful resuscitation, both for ROSC and for protection of the central nervous system. Chest compressions should be carried out while the defibrillator is being charged. As soon as a diagnosis of VF or VT is established, a biphasic waveform shock of 150–200 J (360 J if a monophasic waveform device is used) should be delivered. If 5 min has elapsed between collapse and first contact with the victim, there is some evidence that 60–90 s of CPR before the first shock may improve probability of survival without neurologic damage. If the initial shock does not successfully revert VT or VF, chest compression at a rate of 100 per minute is resumed for 2 min, and then

a second shock is delivered. Multiple shocks given in sequence are no longer recommended, in order to minimize interruptions of chest compressions. This sequence is continued until personnel capable of, and equipped for, advanced life support are available, although not much data support the notion that shocks and chest compressions alone will revert VF after three shocks have failed.

ADVANCED CARDIAC LIFE SUPPORT (ACLS)

ACLS is intended to achieve and maintain organ perfusion and adequate ventilation, control cardiac arrhythmias, and stabilize blood pressure and cardiac output. The activities carried out to achieve these goals include (1) defibrillation/cardioversion and/or pacing, (2) intubation with an endotracheal tube, and (3) insertion of an intravenous line.

As in basic life support, the major emphasis during ACLS is minimizing interruptions of chest compressions until ROSC is achieved. After two or three unsuccessful defibrillation attempts, epinephrine, 1 mg IV, is given and attempts to defibrillate are repeated. The dose of epinephrine may be repeated after intervals of 3–5 min (Fig. 327-3A). Vasopressin (a single 40-unit dose given IV) has been suggested as an alternative to epinephrine.

If the patient is less than fully conscious upon reversion or if two or three attempts fail, prompt intubation, ventilation, and arterial blood gas analysis should be carried out. Ventilation with O₂ (room air if O₂ is not immediately available) may promptly reverse hypoxemia and acidosis. Quantitative waveform capnography is now recommended for confirmation and monitoring of endotracheal tube placement. A patient who is persistently acidotic after successful defibrillation and intubation or had acidosis prior to arrest, may be

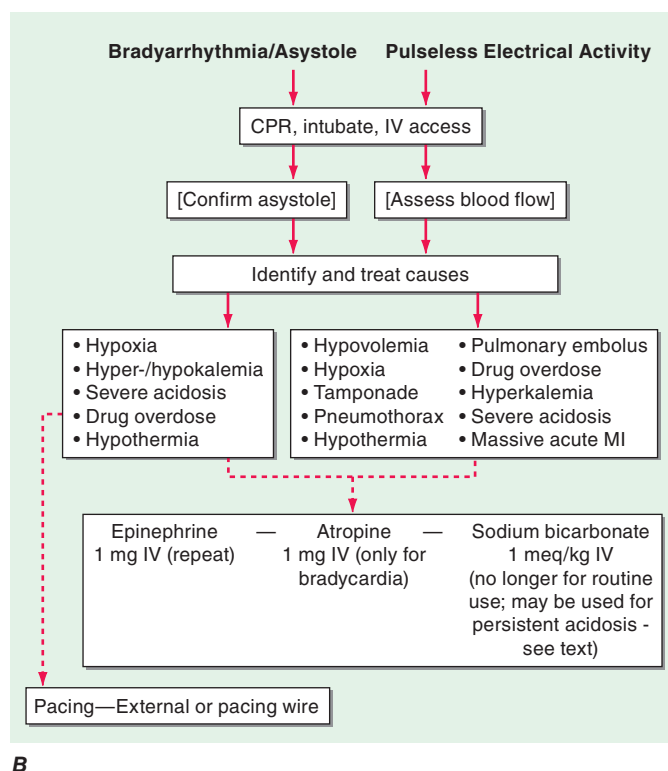
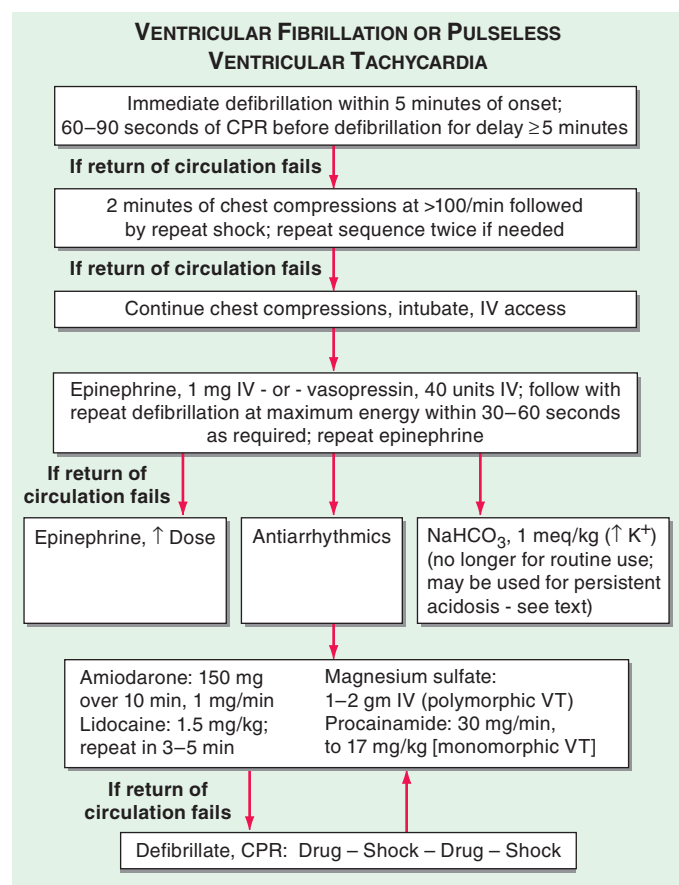


FIGURE 327-3 **A.** The algorithm of ventricular fibrillation or pulseless ventricular tachycardia begins with and initial defibrillate on attempt. If a single shock fails to restore a pulse, it is followed by 2 min of cardiopulmonary resuscitation (CPR; chest compressions), followed by another single shock. After three such sequences, epinephrine and then antiarrhythmic drugs are added to the protocol. See text for details. **B.** The algorithms for bradyarrhythmia/asystole (left) or pulseless electrical activity (right) are dominated first by continued life support and a search for reversible causes. Subsequent therapy is nonspecific and is accompanied by a low success rate. See text for details. MI, myocardial infarction; VT, ventricular tachycardia.

1770 given 1 meq/kg NaHCO₃ initially and an additional 50% of the dose repeated every 10–15 min. However, it should not be used routinely.

After initial unsuccessful defibrillation attempts or with persistent/recurrent electrical instability, antiarrhythmic therapy should be instituted. Intravenous amiodarone has emerged as the initial treatment of choice (150 mg over 10 min, followed by 1 mg/min for up to 6 h and 0.5 mg/min thereafter) (Fig. 327-3A). For cardiac arrest due to VF in the early phase of an acute coronary syndrome, a bolus of 1 mg/kg of lidocaine may be given intravenously as an alternative, and the dose may be repeated in 2 min. It also may be tried in patients in whom amiodarone is unsuccessful. Intravenous procainamide (loading infusion of 100 mg/5 min to a total dose of 500–800 mg, followed by continuous infusion at 2–5 mg/min) is now rarely used in this setting but may be tried for persisting, hemodynamically stable arrhythmias. Intravenous calcium gluconate is no longer considered safe or necessary for routine administration. It is used only in patients in whom acute hyperkalemia is known to be the triggering event for resistant VF, in the presence of known hypocalcemia, or in patients who have received toxic doses of calcium channel antagonists.

Cardiac arrest due to bradyarrhythmias or asystole (B/A cardiac arrest) is managed differently (Fig. 327-3B). The patient is promptly intubated, CPR is continued, and an attempt is made to control hypoxemia and acidosis and identify other reversible causes. Epinephrine may be given intravenously or by an intraosseous route. Atropine is no longer considered effective for asystole or PEA, but can be used for bradyarrhythmias. External pacing devices are used to attempt to establish a regular rhythm when atropine fails for a bradyarrhythmia, but chronotropic agents given intravenously are now recognized as an equally effective alternative.

The success rate may be good when B/A arrest is due to acute inferior wall MI or to correctable airway obstruction or drug-induced respiratory depression or with prompt resuscitation efforts. For acute airway obstruction, prompt removal of foreign bodies by the Heimlich maneuver or, in hospitalized patients, by intubation and suctioning of obstructing secretions in the airway is often successful. The prognosis is generally very poor in other causes of this form of cardiac arrest, such as end-stage cardiac or noncardiac diseases. Treatment of PEA is similar to that for bradyarrhythmias, but its outcome is also dismal.

POST-CARDIAC ARREST SYNDROME AND POSTRESUSCITATION CARE

After return of spontaneous or stable assisted circulation, attention shifts to the diagnostic and therapeutic elements of the post-cardiac arrest syndrome. This recently developed clinical classification emerged from the organization of the elements of injury following cardiac arrest into a multidisciplinary continuum. The four components of post-cardiac arrest syndrome include brain injury, myocardial dysfunction, systemic ischemia/reperfusion responses, and control of persistent precipitating factors. The therapeutic goal is to maintain a stable electrical, hemodynamic, and central nervous system status.

Postresuscitation care is determined by the specific clinical circumstances. The most pressing is the presence of anoxic encephalopathy, which is a strong predictor of in-hospital death and postarrest disability. Mild therapeutic hypothermia is indicated for resuscitated cardiac arrest victims who are hemodynamically stable, but remain comatose. Core body temperature is decreased to 32–34°C, by several available techniques (external and/or internal [core]), as soon as practical after resuscitation and maintained for a minimum of 12–24 h. By reducing metabolic demands and cerebral edema, this intervention improves probability of survival with better neurologic outcome.

Primary VF in acute MI (not accompanied by low-output states) (Chap. 295) is generally very responsive to life support techniques and easily controlled after the initial event. In the in-hospital setting, respirator support is usually not necessary or is needed for only a short time, and hemodynamics stabilize promptly after defibrillation or cardioversion. In secondary VF in acute MI (those events in

which hemodynamic abnormalities predispose to the potentially fatal arrhythmia), resuscitative efforts are less often successful, and in patients who are successfully resuscitated, the recurrence rate is high. The clinical picture and outcome are dominated by hemodynamic instability and the ability to control hemodynamic dysfunction. Bradyarrhythmias, asystole, and PEA are commonly secondary events in hemodynamically unstable patients.

The outcome after in-hospital cardiac arrest associated with noncardiac diseases is poor, and in the few successfully resuscitated patients, the postresuscitation course is dominated by the nature of the underlying disease. Patients with end-stage cancer, renal failure, acute central nervous system disease, and uncontrolled infections, as a group, have a survival rate of <10% after in-hospital cardiac arrest. Some major exceptions are patients with transient airway obstruction, electrolyte disturbances, proarrhythmic effects of drugs, and severe metabolic abnormalities, most of whom may have a good chance of survival if they can be resuscitated promptly and stabilized while the transient abnormalities are being corrected.

LONG-TERM MANAGEMENT AFTER SURVIVAL OF OUT-OF-HOSPITAL CARDIAC ARREST

Patients who survive cardiac arrest without irreversible damage to the central nervous system and who achieve hemodynamic stability should have diagnostic testing to define appropriate therapeutic interventions for their long-term management. This approach is driven by the fact that survival after out-of-hospital cardiac arrest is followed by a 10–25% mortality rate during the first 2 years after the event, and there are data suggesting that significant survival benefits can be achieved by prescription of an ICD.

Among patients in whom an acute ST elevation MI or transient and reversible myocardial ischemia is identified as the specific mechanism triggering an out-of-hospital cardiac arrest, the management is dictated in part by the transient nature of life-threatening arrhythmia risk during the acute coronary syndrome (ACS) and in part by the extent of permanent myocardial damage that results. Cardiac arrest during the acute ischemic phase is not an ICD indication, but survivors of cardiac arrest not associated with an ACS do benefit. In addition, patients who survive MI with an EF less than 30–35% appear to benefit from ICDs.

For patients with cardiac arrest determined to be due to a treatable transient ischemic mechanism, particularly with higher EFs, catheter interventional, surgical, and/or pharmacologic anti-ischemic therapy is generally accepted for long-term management.

Survivors of cardiac arrest due to other categories of disease, such as the hypertrophic or dilated cardiomyopathies and the various rare inherited disorders (e.g., right ventricular dysplasia, long QT syndrome, Brugada syndrome, catecholaminergic polymorphic VT, and so-called idiopathic VF), are all considered ICD candidates.

PREVENTION OF SCD IN HIGH-RISK INDIVIDUALS WITHOUT PRIOR CARDIAC ARREST

Post-MI patients with EFs <35% and other markers of risk such as ambient ventricular arrhythmias, inducible ventricular tachyarrhythmias in the electrophysiology laboratory, and a history of heart failure are considered candidates for ICDs 40 days or more after the MI. Total mortality benefits in the range of a 20–35% reduction over 2–5 years have been observed in a series of clinical trials. One study suggested that an EF <30% was a sufficient marker of risk to indicate ICD benefit, and another demonstrated benefit for patients with Functional Class 2 or 3 heart failure and EFs ≤35%, regardless of etiology (ischemic or nonischemic) or the presence of ambient or induced arrhythmias (Chaps. 277 and 279). For patients with newly diagnosed heart failure and an EF <35%, the required delay between diagnosis and institution of medical therapy, and subsequent implantation of an ICD, is 90 days. In general, there appears to be a gradient of increasing ICD benefit with EFs ranging lower than the threshold indications. However, patients with very low EFs (e.g., <20%) may receive less benefit.

Decision making for primary prevention in disorders other than coronary artery disease and dilated cardiomyopathy is generally driven by observational data and judgment based on clinical observations. Controlled clinical trials providing evidence-based indicators for ICDs are lacking for these smaller population subgroups. In general, for

the rare disorders listed above, indicators of arrhythmic risk such as syncope, documented ventricular tachyarrhythmias, aborted cardiac arrest, or a family history of premature SCD in some conditions, and a number of other clinical or ECG markers, may be used as indicators for ICDs.

SECTION 3

NEUROLOGIC CRITICAL CARE

328 Coma

Allan H. Ropper

Coma is among the most common and striking problems in general medicine. It accounts for a substantial portion of admissions to emergency wards and occurs on all hospital services. It demands immediate attention and requires an organized approach.

There is a continuum of states of reduced alertness, the most severe form being coma, defined as a deep sleeplike state from which the patient cannot be aroused. Stupor refers to a higher degree of arousability in which the patient can be transiently awakened by vigorous stimuli, accompanied by motor behavior that leads to avoidance of uncomfortable or aggravating stimuli. Drowsiness, which is familiar to all persons, simulates light sleep and is characterized by easy arousal and the persistence of alertness for brief periods. Drowsiness and stupor are usually accompanied by some degree of confusion (Chap. 34). A precise narrative description of the level of arousal and of the type of responses evoked by various stimuli as observed at the bedside is preferable to ambiguous terms such as lethargy, semicomatose, or obtundation.

Several conditions that render patients unresponsive and simulate coma are considered separately because of their special significance. The vegetative state signifies an awake-appearing but nonresponsive state in a patient who has emerged from coma. In the vegetative state, the eyelids may open, giving the appearance of wakefulness. Respiratory and autonomic functions are retained. Yawning, coughing, swallowing, and limb and head movements persist, and the patient may follow visually presented objects, but there are few, if any, meaningful responses to the external and internal environment—in essence, an “awake coma.” The term *vegetative* is unfortunate because it is subject to misinterpretation. There are always accompanying signs that indicate extensive damage in both cerebral hemispheres, e.g., decerebrate or decorticate limb posturing and absent responses to visual stimuli (see below). In the closely related but less severe minimally conscious state, the patient displays rudimentary vocal or motor behaviors, often spontaneous, but some in response to touch, visual stimuli, or command. Cardiac arrest with cerebral hypoperfusion and head injuries are the most common causes of the vegetative and minimally conscious states (Chaps. 327 and 330). The prognosis for regaining mental faculties once the vegetative state has supervened for several months is very poor, and after a year, almost nil; hence the term *persistent vegetative state*. Most reports of dramatic recovery, when investigated carefully, are found to yield to the usual rules for prognosis, but there have been rare instances in which recovery has occurred to a severely disabled condition and, in rare childhood cases, to an even better state. The possibility of incorrectly attributing meaningful behavior to patients in the vegetative and minimally conscious states creates inordinate problems and anguish. On the other hand, the question of whether these patients lack any capability for cognition has been reopened by functional imaging studies that have demonstrated, in a small proportion of posttraumatic cases, meaningful cerebral activation in response to verbal and other stimuli.

Apart from the above conditions, several syndromes that affect alertness are prone to be misinterpreted as stupor or coma. Akinetic mutism refers to a partially or fully awake state in which the patient is able to form impressions and think, as demonstrated by later recounting of events, but remains virtually immobile and mute. The condition results from damage in the regions of the medial thalamic nuclei or the frontal lobes (particularly lesions situated deeply or on the orbitofrontal surfaces) or from extreme hydrocephalus. The term *abulia* describes a milder form of akinetic mutism characterized by mental and physical slowness and diminished ability to initiate activity. It is also usually the result of damage to the frontal lobes and its connections (Chap. 36).

Catatonia is a curious hypomobile and mute syndrome that occurs as part of a major psychosis, usually schizophrenia or major depression. Catatonic patients make few voluntary or responsive movements, although they blink, swallow, and may not appear distressed. There are nonetheless signs that the patient is responsive, although it may take ingenuity on the part of the examiner to demonstrate them. For example, eyelid elevation is actively resisted, blinking occurs in response to a visual threat, and the eyes move concomitantly with head rotation, all of which are inconsistent with the presence of a brain lesion causing unresponsiveness. It is characteristic but not invariable in catatonia for the limbs to retain the postures in which they have been placed by the examiner (“waxy flexibility,” or *cataplexy*). With recovery, patients often have some memory of events that occurred during their catatonic stupor. Catatonia is superficially similar to akinetic mutism, but clinical evidence of cerebral damage such as Babinski signs and hypertonicity of the limbs is lacking. The special problem of coma in brain death is discussed below.

The locked-in state describes yet another type of pseudocomatose state in which an awake patient has no means of producing speech or volitional movement but retains voluntary vertical eye movements and lid elevation, thus allowing the patient to signal with a clear mind. The pupils are normally reactive. Such individuals have written entire treatises using Morse code. The usual cause is an infarction or hemorrhage of the ventral pons that transects all descending motor (corticospinal and corticobulbar) pathways. A similar awake but de-efferented state occurs as a result of total paralysis of the musculature in severe cases of Guillain-Barré syndrome (Chap. 460), critical illness neuropathy (Chap. 330), and pharmacologic neuromuscular blockade.

THE ANATOMY AND PHYSIOLOGY OF COMA

Almost all instances of diminished alertness can be traced to widespread abnormalities of the cerebral hemispheres or to reduced activity of a special thalamocortical alerting system termed the reticular activating system (RAS). The proper functioning of this system, its ascending projections to the cortex, and the cortex itself are required to maintain alertness and coherence of thought. It follows that the principal causes of coma are (1) lesions that damage the RAS in the upper midbrain or its projections; (2) destruction of large portions of both cerebral hemispheres; or (3) suppression of reticulocerebral function by drugs, toxins, or metabolic derangements such as hypoglycemia, anoxia, uremia, and hepatic failure.

The proximity of the RAS to midbrain structures that control pupillary function and eye movements permits clinical localization of the cause of coma in many cases. Pupillary enlargement with loss of light

1772 reaction and loss of vertical and adduction movements of the eyes suggests that the lesion is in the upper brainstem where the nuclei subserving these functions reside. Conversely, preservation of pupillary light reactivity and of eye movements absolves the upper brainstem and indicates that widespread structural lesions or metabolic suppression of the cerebral hemispheres is responsible for coma.

Coma Due to Cerebral Mass Lesions and Herniations In addition to the fixed restriction of the skull, the cranial cavity is separated into compartments by infoldings of the dura. The two cerebral hemispheres are separated by the falx, and the anterior and posterior fossae by the tentorium. Herniation refers to displacement of brain tissue by an overlying or adjacent mass into a contiguous compartment that it normally does not occupy. Coma and many of its associated signs can be attributed to these tissue shifts, and certain clinical features are characteristic of specific configurations of herniation (Fig. 328-1). They are in essence “false localizing” signs because they derive from compression of brain structures at a distance from the mass.

In the most common form of herniation, brain tissue is displaced from the supratentorial to the infratentorial compartment through the tentorial opening; this is referred to as transtentorial herniation. Uncal transtentorial herniation refers to impaction of the anterior medial temporal gyrus (the uncus) into the tentorial opening just anterior to and adjacent to the midbrain (Fig. 328-1A). The uncus compresses the third nerve as the nerve traverses the subarachnoid space, causing enlargement of the ipsilateral pupil (the fibers subserving parasympathetic pupillary function are located peripherally in the nerve). The coma that follows is due to compression of the midbrain against the opposite tentorial edge by the displaced parahippocampal gyrus (Fig. 328-2). Lateral displacement of the midbrain may compress the opposite cerebral peduncle against the tentorial edge, producing a Babinski sign and hemiparesis contralateral to the hemiparesis that resulted from the mass (the Kernohan-Woltman sign). Herniation may also compress the anterior and posterior cerebral arteries as they pass over the tentorial reflections, with resultant brain infarction. The distortions may also entrap portions of the ventricular system, resulting in hydrocephalus.

Central transtentorial herniation denotes a symmetric downward movement of the thalamic structures through the tentorial opening with compression of the upper midbrain (Fig. 328-1B). Miotic pupils and drowsiness are the heralding signs, in contrast to a unilaterally

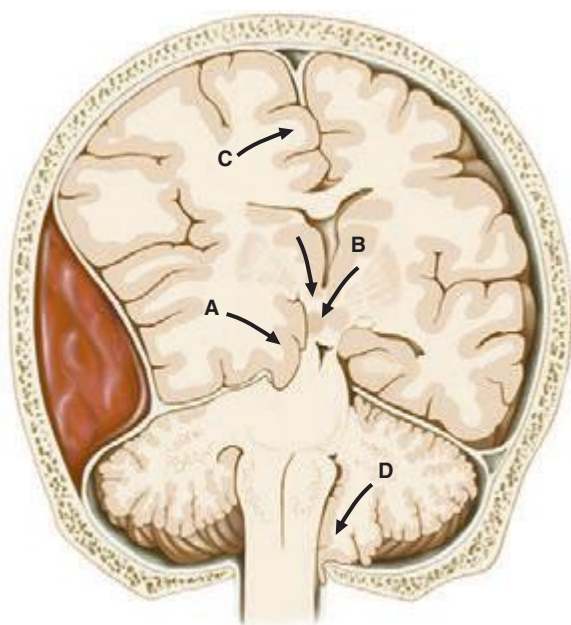


FIGURE 328-1 Types of cerebral herniation: (A) uncal; (B) central; (C) transfalcial; and (D) foraminal.

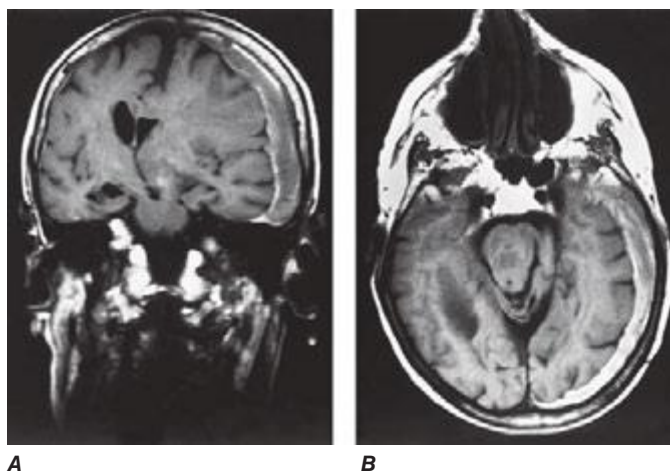


FIGURE 328-2 Coronal (A) and axial (B) magnetic resonance images from a stuporous patient with a left third nerve palsy as a result of a large left-sided subdural hematoma (seen as a gray-white rim). The upper midbrain and lower thalamic regions are compressed and displaced horizontally away from the mass, and there is transtentorial herniation of the medial temporal lobe structures, including the uncus anteriorly. The lateral ventricle opposite to the hematoma has become enlarged as a result of compression of the third ventricle.

enlarged pupil of the uncal syndrome. Both uncal and central transtentorial herniations cause progressive compression of the brainstem, with initial damage to the midbrain, then the pons, and finally the medulla. The result is an approximate sequence of neurologic signs that corresponds to each affected level. Other forms of herniation are transfalcial herniation (displacement of the cingulate gyrus under the falx and across the midline, Fig. 328-1C) and foraminal herniation (downward forcing of the cerebellar tonsils into the foramen magnum, Fig. 328-1D), which causes compression of the medulla, respiratory arrest, and death.

A direct relationship between the various configurations of transtentorial herniation and coma is not always found. Drowsiness and stupor can occur with moderate horizontal displacement of the diencephalon (thalamus), before transtentorial herniation is evident. This lateral shift may be quantified on axial images of computed tomography (CT) and magnetic resonance imaging (MRI) scans (Fig. 328-2). In cases of acutely enlarging masses, horizontal displacement of the pineal calcification of 3–5 mm is generally associated with drowsiness, 6–8 mm with stupor, and >9 mm with coma. Intrusion of the medial temporal lobe into the tentorial opening is also apparent on MRI and CT scans as obliteration of the cisterna that surrounds the upper brainstem.

Coma due to Metabolic Disorders Many systemic metabolic abnormalities cause coma by interrupting the delivery of energy substrates (e.g., oxygen, glucose) or by altering neuronal excitability (drugs and alcohol, anesthesia, and epilepsy). The metabolic abnormalities that produce coma may, in milder forms, induce an acute confusional state. Thus, in metabolic encephalopathies, clouded consciousness and coma are in a continuum.

Cerebral neurons are fully dependent on cerebral blood flow (CBF) and the delivery of oxygen and glucose. CBF is ~75 mL per 100 g/min in gray matter and 30 mL per 100 g/min in white matter (mean ~55 mL per 100 g/min); oxygen consumption is 3.5 mL per 100 g/min, and glucose utilization is 5 mg per 100 g/min. Brain stores of glucose are able to provide energy for ~2 min after blood flow is interrupted, and oxygen stores last 8–10 s after the cessation of blood flow. Simultaneous hypoxia and ischemia exhaust glucose more rapidly. The electroencephalogram (EEG) rhythm in these circumstances becomes diffusely slowed, typical of metabolic encephalopathies, and as substrate delivery worsens, eventually brain electrical activity ceases.

Unlike hypoxia-ischemia, which causes neuronal destruction, most metabolic disorders such as hypoglycemia, hyponatremia, hyperosmolarity, hypercapnia, hypercalcemia, and hepatic and renal failure cause only minor neuropathologic changes. The reversible effects of these conditions on the brain are not understood but may result from impaired energy supplies, changes in ion fluxes across neuronal membranes, and neurotransmitter abnormalities. For example, the high ammonia concentration of hepatic coma interferes with cerebral energy metabolism and with the Na^+ , K^+ -ATPase pump, increases the number and size of astrocytes, and causes increased concentrations of potentially toxic products of ammonia metabolism; it may also affect neurotransmitters, including the production of putative “false” neurotransmitters that are active at receptor sites. Apart from hyperammonemia, which of these mechanisms is of critical importance is not clear. The mechanism of the encephalopathy of renal failure is also not known. Unlike ammonia, urea does not produce central nervous system (CNS) toxicity, and a multifactorial causation has been proposed for the encephalopathy, including increased permeability of the blood-brain barrier to toxic substances such as organic acids and an increase in brain calcium and cerebrospinal fluid (CSF) phosphate content.

Coma and seizures are common accompaniments of large shifts in sodium and water balance in the brain. These changes in osmolarity arise from systemic medical disorders, including diabetic ketoacidosis, the nonketotic hyperosmolar state, and hyponatremia from any cause (e.g., water intoxication, excessive secretion of antidiuretic hormone, or atrial natriuretic peptides). Sodium levels <125 mmol/L induce confusion, and levels <115 mmol/L are typically associated with coma and convulsions. In hyperosmolar coma, the serum osmolarity is generally >350 mosmol/L. Hypercapnia depresses the level of consciousness in proportion to the rise in carbon dioxide (CO_2) tension in the blood. In all of these metabolic encephalopathies, the degree of neurologic change depends to a large extent on the rapidity with which the serum changes occur. The pathophysiology of other metabolic encephalopathies such as those due to hypercalcemia, hypothyroidism, vitamin B_{12} deficiency, and hypothermia are incompletely understood but must reflect derangements of CNS biochemistry, membrane function, or neurotransmitters.

Epileptic Coma Generalized electrical seizures are associated with coma, even in the absence of motor convulsions (nonconvulsive status epilepticus). The self-limited coma that follows a seizure, the postictal state, may be due to exhaustion of energy reserves or effects of locally toxic molecules that are the by-product of seizures. The postictal state produces continuous, generalized slowing of the background EEG activity similar to that of metabolic encephalopathies.

Toxic (Including Drug-Induced) Coma This common class of encephalopathy is in large measure reversible and leaves no residual damage provided there has not been cardiorespiratory failure. Many drugs and toxins are capable of depressing nervous system function. Some produce coma by affecting both the brainstem nuclei, including the RAS, and the cerebral cortex. The combination of cortical and brainstem signs, which occurs in certain drug overdoses, may lead to an incorrect diagnosis of structural brainstem disease. Overdose of medications that have atropinic actions produces signs such as dilated pupils, tachycardia, and dry skin; opiate overdose produces pinpoint pupils <1 mm in diameter.

Coma due to Widespread Damage to the Cerebral Hemispheres This category, comprising a number of unrelated disorders, results from widespread structural cerebral damage that simulates a metabolic disorder of the cortex. Hypoxia-ischemia is perhaps the best characterized and one in which it is not possible initially to distinguish the acute reversible effects of oxygen deprivation of the brain from the subsequent effects of anoxic neuronal damage. Similar widespread cerebral damage may be produced by disorders that occlude small blood vessels throughout the brain; examples include cerebral malaria, thrombotic thrombocytopenic purpura, and hyperviscosity. Diffuse white matter damage from cranial trauma or inflammatory demyelinating diseases can cause a similar coma syndrome.

APPROACH TO THE PATIENT:

Coma

A video examination of the comatose patient is shown in Chap. 329e.

Acute respiratory and cardiovascular problems should be attended to prior to neurologic assessment. In most instances, a complete medical evaluation, except for vital signs, funduscopy, and examination for nuchal rigidity, may be deferred until the neurologic evaluation has established the severity and nature of coma. **The approach to the patient with coma from cranial trauma is discussed in Chap. 457e.**

HISTORY

The cause of coma may be immediately evident as in cases of trauma, cardiac arrest, or observed drug ingestion. In the remainder, certain points are useful: (1) the circumstances and rapidity with which neurologic symptoms developed; (2) the antecedent symptoms (confusion, weakness, headache, fever, seizures, dizziness, double vision, or vomiting); (3) the use of medications, drugs, or alcohol; and (4) chronic liver, kidney, lung, heart, or other medical disease. Direct interrogation of family, observers, and ambulance technicians on the scene, in person or by telephone, is an important part of the evaluation when possible.

GENERAL PHYSICAL EXAMINATION

Fever suggests a systemic infection, bacterial meningitis, encephalitis, heat stroke, neuroleptic malignant syndrome, malignant hyperthermia due to anesthetics, or anticholinergic drug intoxication. Only rarely is fever attributable to a lesion that has disturbed hypothalamic temperature-regulating centers (“central fever”). A slight elevation in temperature may follow vigorous convulsions. Hypothermia is observed with exposure that attends alcohol, barbiturate, sedative, or phenothiazine intoxication; hypoglycemia; peripheral circulatory failure; or extreme hypothyroidism. Hypothermia itself causes coma when the temperature is $<31^\circ\text{C}$ (87.8°F). Tachypnea may indicate systemic acidosis or pneumonia or, rarely, infiltration of the brain with lymphoma. Aberrant respiratory patterns that reflect brainstem disorders are discussed below. Marked hypertension suggests hypertensive encephalopathy or cerebral hemorrhage or head injury. Hypotension is characteristic of coma from alcohol or barbiturate intoxication, internal hemorrhage, myocardial infarction, sepsis, profound hypothyroidism, or Addisonian crisis. The funduscopic examination can detect subarachnoid hemorrhage (subhyaloid hemorrhages), hypertensive encephalopathy (exudates, hemorrhages, vessel-crossing changes, papilledema), and increased intracranial pressure (ICP) (papilledema). Cutaneous petechiae suggest thrombotic thrombocytopenic purpura, meningococcemia, or a bleeding diathesis associated with an intracerebral hemorrhage. Cyanosis and reddish or anemic skin coloration are other indications of an underlying systemic disease or carbon monoxide as responsible for the coma.

NEUROLOGIC EXAMINATION

The patient should be observed without intervention by the examiner. Tossing about in the bed, reaching up toward the face, crossing legs, yawning, swallowing, coughing, or moaning reflect a drowsy state that is close to normal awakesness. Lack of restless movements on one side or an outturned leg suggests a hemiplegia. Intermittent twitching movements of a foot, finger, or facial muscle may be the only sign of seizures. Multifocal myoclonus almost always indicates a metabolic disorder, particularly uremia, anoxia, drug intoxication (especially with lithium or haloperidol), or a prion disease (Chap. 453e). In a drowsy and confused patient, bilateral asterixis is a certain sign of metabolic encephalopathy or drug intoxication.

Decorticate rigidity and decerebrate rigidity, or “posturing,” describe stereotyped arm and leg movements occurring spontaneously

or elicited by sensory stimulation. Flexion of the elbows and wrists and supination of the arm (decorticate posturing) suggests bilateral damage rostral to the midbrain, whereas extension of the elbows and wrists with pronation (decerebrate posturing) indicates damage to motor tracts in the midbrain or caudal diencephalon. The less frequent combination of arm extension with leg flexion or flaccid legs is associated with lesions in the pons. These concepts have been adapted from animal work and cannot be applied with precision to coma in humans. In fact, acute and widespread disorders of any type, regardless of location, frequently cause limb extension, and almost all extensor posturing becomes predominantly flexor as time passes.

LEVEL OF AROUSAL

A sequence of increasingly intense stimuli is used to determine the threshold for arousal and the motor response of each side of the body. The results of testing may vary from minute to minute, and serial examinations are useful. Tickling the nostrils with a cotton wisp is a moderate stimulus to arousal—all but deeply stuporous and comatose patients will move the head away and arouse to some degree. An even greater degree of responsiveness is present if the patient uses his hand to remove an offending stimulus. Pressure on the knuckles or bony prominences and pinprick stimulation are humane forms of noxious stimuli; pinching the skin causes unsightly ecchymoses and is generally not necessary but may be useful in eliciting abduction withdrawal movements of the limbs. Posturing in response to noxious stimuli indicates severe damage to the corticospinal system, whereas abduction-avoidance movement of a limb is usually purposeful and denotes an intact corticospinal system. Posturing may also be unilateral and coexist with purposeful limb movements, reflecting incomplete damage to the motor system.

BRAINSTEM REFLEXES

Assessment of brainstem function is essential to localization of the lesion in coma (Fig. 328-3). The brainstem reflexes that are examined are pupillary size and reaction to light, spontaneous and elicited eye movements, corneal responses, and the respiratory pattern. As a rule, coma due to bilateral hemispherical disease preserves these brainstem activities, particularly the pupillary reactions and eye movements. However, the presence of abnormal brainstem signs does not always indicate that the primary lesion is in the brainstem because hemispherical masses can cause secondary brainstem damage by the earlier described transtentorial herniations.

Pupillary Signs Pupillary reactions are examined with a bright, diffuse light (preferably not an ophthalmoscope, which illuminates only a limited part of the retina). Reactive and round pupils of midsize (2.5–5 mm) essentially exclude midbrain damage, either primary or secondary to compression. A response to light may be difficult to appreciate in pupils <2 mm in diameter, and bright room lighting mutes pupillary reactivity. One enlarged and poorly reactive pupil (>6 mm) signifies compression or stretching of the third nerve from the effects of a cerebral mass above. Enlargement of the pupil contralateral to a hemispherical mass may occur but is infrequent. An oval and slightly eccentric pupil is a transitional sign that accompanies early midbrain–third nerve compression. The most extreme pupillary sign, bilaterally dilated and unreactive pupils, indicates severe midbrain damage, usually from compression by a supratentorial mass. Ingestion of drugs with anticholinergic activity, the use of mydriatic eye drops, and direct ocular trauma are among the causes of misleading pupillary enlargement.

Unilateral miosis in coma has been attributed to dysfunction of sympathetic efferents originating in the posterior hypothalamus and descending in the tegmentum of the brainstem to the cervical cord. It is therefore of limited localizing value but is an occasional finding in patients with a large cerebral hemorrhage that affects the thalamus. Reactive and bilaterally small (1–2.5 mm) but not pinpoint pupils are seen in metabolic encephalopathies or in deep bilateral hemispherical lesions such as hydrocephalus or thalamic

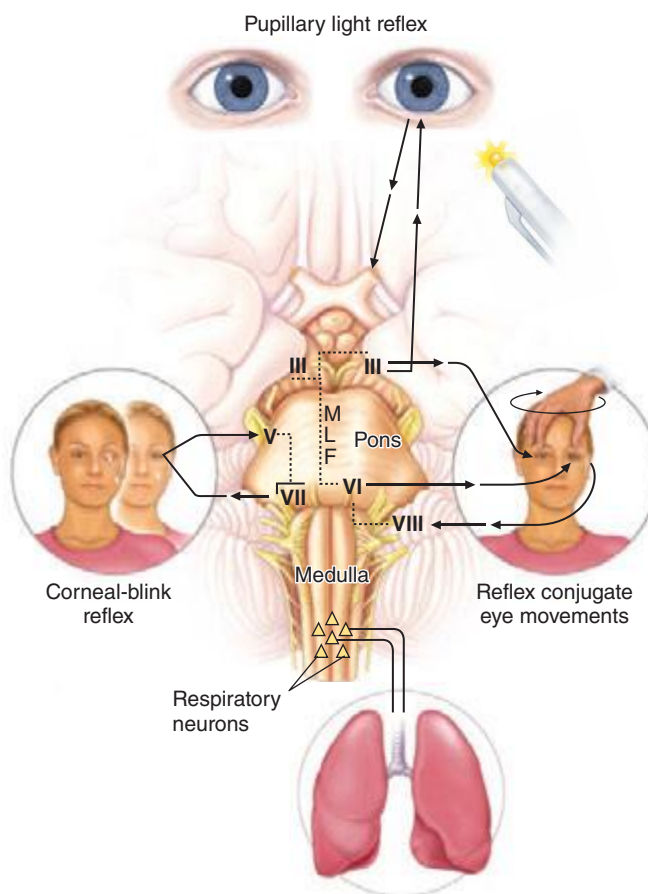


FIGURE 328-3 Examination of brainstem reflexes in coma. Midbrain and third nerve function are tested by pupillary reaction to light, pontine function by spontaneous and reflex eye movements and corneal responses, and medullary function by respiratory and pharyngeal responses. Reflex conjugate, horizontal eye movements are dependent on the medial longitudinal fasciculus (MLF) interconnecting the sixth and contralateral third nerve nuclei. Head rotation (oculocephalic reflex) or caloric stimulation of the labyrinths (oculovestibular reflex) elicits contraversive eye movements (for details see text).

hemorrhage. Even smaller reactive pupils (<1 mm) characterize narcotic or barbiturate overdoses but also occur with extensive pontine hemorrhage. The response to naloxone and the presence of reflex eye movements (see below) assist in distinguishing between these.

Ocular Movements The eyes are first observed by elevating the lids and observing the resting position and spontaneous movements of the globes. Lid tone, tested by lifting the eyelids and noting their resistance to opening and the speed of closure, is progressively reduced as unresponsiveness progresses. Horizontal divergence of the eyes at rest is normal in drowsiness. As coma deepens, the ocular axes may become parallel again.

Spontaneous eye movements in coma often take the form of conjugate horizontal roving. This finding alone exonerates damage in the midbrain and pons and has the same significance as normal reflex eye movements (see below). Conjugate horizontal ocular deviation to one side indicates damage to the pons on the opposite side or, alternatively, to the frontal lobe on the same side. This phenomenon is summarized by the following maxim: *The eyes look toward a hemispherical lesion and away from a brainstem lesion.* Seizures also drive the eyes to one side but usually with superimposed clonic movements of the globes. The eyes may occasionally turn paradoxically away from the side of a deep hemispherical lesion (“wrong-way eyes”). The eyes turn down and inward with thalamic and upper midbrain lesions, typically thalamic hemorrhage.

“Ocular bobbing” describes brisk downward and slow upward movements of the eyes associated with loss of horizontal eye movements and is diagnostic of bilateral pontine damage, usually from thrombosis of the basilar artery. “Ocular dipping” is a slower, arrhythmic downward movement followed by a faster upward movement in patients with normal reflex horizontal gaze; it usually indicates diffuse cortical anoxic damage.

The oculocephalic reflexes, elicited by moving the head from side to side or vertically and observing eye movements in the direction opposite to the head movement, depend on the integrity of the ocular motor nuclei and their interconnecting tracts that extend from the midbrain to the pons and medulla (Fig. 328-3). The movements, called somewhat inappropriately “doll’s eyes” (which refers more accurately to the reflex elevation of the eyelids with flexion of the neck), are normally suppressed in the awake patient. The ability to elicit them therefore reflects both reduced cortical influence on the brainstem and intact brainstem pathways, indicating that coma is caused by a lesion or dysfunction in the cerebral hemispheres. The opposite, an absence of reflex eye movements, usually signifies damage within the brainstem but can result from overdoses of certain drugs. In this circumstance, normal pupillary size and light reaction distinguishes most drug-induced comas from structural brainstem damage.

Thermal, or “caloric,” stimulation of the vestibular apparatus (oculovestibular response) provides a more intense stimulus for the oculocephalic reflex but provides essentially the same information. The test is performed by irrigating the external auditory canal with cool water in order to induce convection currents in the labyrinths. After a brief latency, the result is tonic deviation of both eyes to the side of cool-water irrigation and nystagmus in the opposite direction. (The acronym “COWS” has been used to remind generations of medical students of the direction of nystagmus—“cold water opposite, warm water same.”) The loss of induced conjugate ocular movements indicates brainstem damage. The presence of corrective nystagmus indicates that the frontal lobes are functioning and connected to the brainstem; thus catatonia or hysterical coma is likely.

By touching the cornea with a wisp of cotton, a response consisting of brief bilateral lid closure is normally observed. The corneal reflex depends on the integrity of pontine pathways between the fifth (afferent) and both seventh (efferent) cranial nerves; in conjunction with reflex eye movements, it is a useful test of pontine function. CNS-depressant drugs diminish or eliminate the corneal responses soon after reflex eye movements are paralyzed but before the pupils become unreactive to light. The corneal (and pharyngeal) response may be lost for a time on the side of an acute hemiplegia.

Respiratory Patterns These are of less localizing value in comparison to other brainstem signs. Shallow, slow, but regular breathing suggests metabolic or drug depression. Cheyne-Stokes respiration in its typical cyclic form, ending with a brief apneic period, signifies bihemispherical damage or metabolic suppression and commonly accompanies light coma. Rapid, deep (Kussmaul) breathing usually implies metabolic acidosis but may also occur with pontomesencephalic lesions. Agonal gasps are the result of lower brainstem (medullary) damage and are recognized as the terminal respiratory pattern of severe brain damage. A number of other cyclic breathing variations have been described but are of lesser significance.

LABORATORY STUDIES AND IMAGING

The studies that are most useful in the diagnosis of coma are chemical-toxicologic analysis of blood and urine, cranial CT or MRI, EEG, and CSF examination. Arterial blood gas analysis is helpful in patients with lung disease and acid-base disorders. The metabolic aberrations commonly encountered in clinical practice are usually exposed by measurement of electrolytes, glucose, calcium, osmolarity, and renal (blood urea nitrogen) and hepatic (NH_3) function. Toxicologic analysis may be necessary in any case of acute coma where the diagnosis is not immediately clear. However, the presence of exogenous drugs or toxins, especially alcohol, does not exclude the possibility that

other factors, particularly head trauma, are also contributing to the clinical state. An ethanol level of 43 mmol/L (0.2 g/dL) in nonhabituated patients generally causes impaired mental activity; a level of >65 mmol/L (0.3 g/dL) is associated with stupor. The development of tolerance may allow the chronic alcoholic to remain awake at levels >87 mmol/L (0.4 g/dL).

The availability of CT and MRI has focused attention on causes of coma that are detectable by imaging (e.g., hemorrhage, tumor, or hydrocephalus). Resorting primarily to this approach, although at times expedient, is imprudent because most cases of coma (and confusion) are metabolic or toxic in origin. Furthermore, the notion that a normal CT scan excludes an anatomic lesion as the cause of coma is erroneous. Bilateral hemisphere infarction, acute brainstem infarction, encephalitis, meningitis, mechanical shearing of axons as a result of closed head trauma, sagittal sinus thrombosis, and subdural hematoma isodense to adjacent brain are some of the disorders that may not be detected. Nevertheless, if the source of coma remains unknown, a scan should be obtained.

The EEG (**Chap. 442e**) is useful in metabolic or drug-induced states but is rarely diagnostic. However, it is the essential test to reveal coma that is due to clinically unrecognized, nonconvulsive seizures, and shows fairly characteristic patterns in herpesvirus encephalitis and prion (Creutzfeldt-Jakob) disease. The EEG may be further helpful in disclosing generalized slowing of the background activity, a reflection of the severity of an encephalopathy. Predominant high-voltage slowing (δ or triphasic waves) in the frontal regions is typical of metabolic coma, as from hepatic failure, and widespread fast (β) activity implicates sedative drugs (e.g., benzodiazepines). A special pattern of “alpha coma,” defined by widespread, variable 8- to 12-Hz activity, superficially resembles the normal α rhythm of waking but, unlike normal α activity, is not altered by environmental stimuli. Alpha coma results from pontine or diffuse cortical damage and is associated with a poor prognosis. Normal α activity on the EEG, which is suppressed by stimulating the patient, also alerts the clinician to the locked-in syndrome or to hysteria or catatonia. Still, the most important use of EEG recordings in coma is to reveal clinically inapparent epileptic discharges.

Lumbar puncture is performed less frequently than in the past for coma diagnosis because neuroimaging effectively excludes intracerebral and extensive subarachnoid hemorrhage. However, examination of the CSF remains indispensable in the diagnosis of meningitis and encephalitis. For patients with an altered level of consciousness, it is generally recommended that an imaging study be performed prior to lumbar puncture to exclude a large intracranial mass lesion. Blood culture and antibiotic administration usually precede the imaging study if meningitis is suspected (**Chap. 164**).

DIFFERENTIAL DIAGNOSIS OF COMA

(**Table 328-1**) The causes of coma can be divided into three broad categories: those cases without focal neurologic signs (e.g., metabolic and toxic encephalopathies); meningitis syndromes, characterized by fever or stiff neck and an excess of cells in the spinal fluid (e.g., bacterial meningitis, subarachnoid hemorrhage, encephalitis); and diseases associated with prominent focal signs (e.g., stroke, cerebral hemorrhage). Conditions that cause sudden coma include drug ingestion, cerebral hemorrhage, trauma, cardiac arrest, epilepsy, and basilar artery occlusion from an embolism. Coma that appears subacutely is usually related to a preexisting medical or neurologic problem or, less often, to secondary brain swelling surrounding a mass such as tumor or cerebral infarction.

The diagnosis of coma due to cerebrovascular disease can be difficult (**Chap. 446**). The most common diseases are (1) basal ganglia and thalamic hemorrhage (acute but not instantaneous onset, vomiting, headache, hemiplegia, and characteristic eye signs); (2) pontine hemorrhage (sudden onset, pinpoint pupils, loss of reflex eye movements and corneal responses, ocular bobbing, posturing, and hyperventilation); (3) cerebellar hemorrhage (occipital headache, vomiting, gaze paresis, and inability to stand and walk); (4) basilar artery thrombosis (neurologic prodrome or warning spells, diplopia, dysarthria, vomiting, eye movement and corneal response abnormalities, and asymmetric limb paresis); and (5) subarachnoid hemorrhage (precipitous

TABLE 328-1 DIFFERENTIAL DIAGNOSIS OF COMA

1. Diseases that cause no focal or lateralizing neurologic signs, usually with normal brainstem functions; CT scan and cellular content of the CSF are normal
 - a. Intoxications: alcohol, sedative drugs, opiates, etc.
 - b. Metabolic disturbances: anoxia, hyponatremia, hypernatremia, hypercalcemia, diabetic acidosis, nonketotic hyperosmolar hyperglycemia, hypoglycemia, uremia, hepatic coma, hypercarbia, Addisonian crisis, hypo- and hyperthyroid states, profound nutritional deficiency
 - c. Severe systemic infections: pneumonia, septicemia, typhoid fever, malaria, Waterhouse-Friderichsen syndrome
 - d. Shock from any cause
 - e. Postseizure states, status epilepticus, nonconvulsive status epilepticus
 - f. Hypertensive encephalopathy, eclampsia
 - g. Severe hyperthermia, hypothermia
 - h. Concussion
 - i. Acute hydrocephalus
2. Diseases that cause meningeal irritation with or without fever, and with an excess of WBCs or RBCs in the CSF, usually without focal or lateralizing cerebral or brainstem signs; CT or MRI shows no mass lesion
 - a. Subarachnoid hemorrhage from ruptured aneurysm, arteriovenous malformation, trauma
 - b. Acute bacterial meningitis
 - c. Viral encephalitis
 - d. Miscellaneous: fat embolism, cholesterol embolism, carcinomatous and lymphomatous meningitis, etc.
3. Diseases that cause focal brainstem or lateralizing cerebral signs, with or without changes in the CSF; CT and MRI are abnormal
 - a. Hemispherical hemorrhage (basal ganglionic, thalamic) or infarction (large middle cerebral artery territory) with secondary brainstem compression
 - b. Brainstem infarction due to basilar artery thrombosis or embolism
 - c. Brain abscess, subdural empyema
 - d. Epidural and subdural hemorrhage, brain contusion
 - e. Brain tumor with surrounding edema
 - f. Cerebellar and pontine hemorrhage and infarction
 - g. Widespread traumatic brain injury
 - h. Metabolic coma (see above) with preexisting focal damage
 - i. Miscellaneous: Cortical vein thrombosis, herpes simplex encephalitis, multiple cerebral emboli due to bacterial endocarditis, acute hemorrhagic leukoencephalitis, acute disseminated (postinfectious) encephalomyelitis, thrombotic thrombocytopenic purpura, cerebral vasculitis, gliomatosis cerebri, pituitary apoplexy, intravascular lymphoma, etc.

Abbreviations: CSF, cerebrospinal fluid; CT, computed tomography; MRI, magnetic resonance imaging; RBCs, red blood cells; WBCs, white blood cells.

coma after sudden severe headache and vomiting). The most common stroke, infarction in the territory of the middle cerebral artery, does not cause coma, but edema surrounding large infarctions may expand over several days and cause coma from mass effect.

The syndrome of acute hydrocephalus accompanies many intracranial diseases, particularly subarachnoid hemorrhage. It is characterized by headache and sometimes vomiting that may progress quickly to coma with extensor posturing of the limbs, bilateral Babinski signs, small unreactive pupils, and impaired oculocephalic movements in the vertical direction.

The majority of medical causes of coma can be established without a neuroimaging study but if the history and examination do not indicate the cause of coma, CT or MRI is needed. Sometimes imaging results can be misleading such as when small subdural hematomas or old strokes are found, but the patient's coma is due to intoxication.

BRAIN DEATH

This is a state of irreversible cessation of all cerebral function with preservation of cardiac activity and maintenance of respiratory and somatic function by artificial means. It is the only type of brain damage recognized as equivalent to death. Criteria have been advanced for

the diagnosis of brain death, and it is essential to adhere to standards endorsed by the local medical community. Ideal criteria are simple, can be assessed at the bedside, and allow no chance of diagnostic error. They contain three essential elements: (1) widespread cortical destruction that is reflected by deep coma and unresponsiveness to all forms of stimulation; (2) global brainstem damage demonstrated by absent pupillary light reaction and by the loss of oculovestibular and corneal reflexes; and (3) destruction of the medulla, manifested by complete and irreversible apnea. The heart rate is invariant and does not accelerate to atropine. Diabetes insipidus is usually present but may only develop hours or days after the other clinical signs of brain death. The pupils are usually midsized but may be enlarged; they should not, however, be small. Loss of deep tendon reflexes is not required because the spinal cord remains functional. Babinski signs are generally absent and the toe response is instead, often flexor.

Demonstration that apnea is due to structural medullary damage requires that the P_{CO_2} be high enough to stimulate respiration during a test of spontaneous breathing. Apnea testing can be done safely by the use of diffusion oxygenation prior to removing the ventilator. This is accomplished by preoxygenation with 100% oxygen, which is then sustained during the test by oxygen administered through a tracheal cannula. CO_2 tension increases ~ 0.3 – 0.4 kPa/min (2–3 mmHg/min) during apnea. At the end of a period of observation, typically several minutes, arterial P_{CO_2} should be at least >6.6 – 8.0 kPa (50–60 mmHg) for the test to be valid. Apnea is confirmed if no respiratory effort has been observed in the presence of a sufficiently elevated P_{CO_2} . Other techniques, including the administration of CO_2 to accelerate the test, are used in special circumstances. The apnea test is usually stopped if there is serious cardiovascular instability.

An isoelectric EEG may be used as a confirmatory test for total cerebral damage. Radionuclide brain scanning, cerebral angiography, or transcranial Doppler measurements may also be included to demonstrate the absence of CBF, but they have not been as extensively correlated with pathologic changes.

The possibility of profound drug-induced or hypothermic depression of the nervous system must be excluded, and some period of observation, usually 6–24 h, is desirable, during which the clinical signs of brain death are sustained. It is advisable to delay clinical testing for at least 24 h if a cardiac arrest has caused brain death or if the inciting disease is not known.

Although it is largely accepted in Western society that the respirator can be disconnected from a brain-dead patient and that organ donation is subsequently possible, problems frequently arise because of poor communication and inadequate preparation of the family by the physician. Reasonable medical practice, ideally with the agreement of the family, also allows the removal of support or transfer out of an intensive care unit of patients who are not brain dead but whose neurologic conditions are nonetheless hopeless.

TREATMENT COMA

The immediate goal in a comatose patient is prevention of further nervous system damage. Hypotension, hypoglycemia, hypercalcemia, hypoxia, hypercapnia, and hyperthermia should be corrected rapidly. An oropharyngeal airway is adequate to keep the pharynx open in a drowsy patient who is breathing normally. Tracheal intubation is indicated if there is apnea, upper airway obstruction, hypoventilation, or emesis, or if the patient is liable to aspirate because of coma. Mechanical ventilation is required if there is hypoventilation or a need to induce hypocapnia in order to lower ICP. IV access is established, and naloxone and dextrose are administered if narcotic overdose or hypoglycemia is a possibility; thiamine is given along with glucose to avoid provoking Wernicke's disease in malnourished patients. In cases of suspected basilar thrombosis with brainstem ischemia, IV heparin or a thrombolytic agent is often used, after cerebral hemorrhage has been excluded by a neuroimaging study. Physostigmine may awaken patients with anticholinergic-type drug overdose but should be used only with careful monitoring; many physicians believe that it should only be used to treat

anticholinergic overdose–associated cardiac arrhythmias. The use of benzodiazepine antagonists offers some prospect of improvement after overdose of soporific drugs and has transient benefit in hepatic encephalopathy. Certain other toxic and drug-induced comas have specific treatments such as fomepizole for ethylene glycol ingestion.

Administration of hypotonic intravenous solutions should be monitored carefully in any serious acute brain illness because of the potential for exacerbating brain swelling. Cervical spine injuries must not be overlooked, particularly before attempting intubation or evaluation of oculocephalic responses. Fever and meningismus indicate an urgent need for examination of the CSF to diagnose meningitis. If the lumbar puncture in a case of suspected meningitis is delayed, an antibiotic such as a third-generation cephalosporin may be administered, preferably after obtaining blood cultures. **The management of raised ICP is discussed in Chap. 330.**

PROGNOSIS

One hopes to avoid the difficult outcome of a patient who is left severely disabled or vegetative. Children and young adults may have ominous early clinical findings such as abnormal brainstem reflexes and yet recover; temporization in offering a prognosis in this group of patients is wise. Metabolic comas have a far better prognosis than traumatic ones. All systems for estimating prognosis in adults should be taken as approximations, and medical judgments must be tempered by factors such as age, underlying systemic disease, and general medical condition. In an attempt to collect prognostic information from large numbers of patients with head injury, the Glasgow Coma Scale was devised; empirically, it has predictive value in cases of brain trauma (see Table 457e-2). For anoxic and metabolic coma, clinical signs such as the pupillary and motor responses after 1 day, 3 days, and 1 week have been shown to have predictive value. Other studies suggest that the absence of corneal responses may have the most discriminative value. The absence of the cortical waves of the somatosensory evoked potentials has also proved a strong indicator of poor outcome in coma from any cause.

The uniformly poor outcome of the prolonged vegetative state has already been mentioned, but recent reports that a small number of such patients display consistent cortical activation on functional MRI in response to salient stimuli have begun to alter the perception of the possible internal mental milieu of such individuals. These findings do not change the poor prognosis. For example, in one series, about 10% of vegetative patients after traumatic brain injury could activate their frontal or temporal lobes in response to requests by an examiner to imagine certain visuospatial tasks. In one case, a rudimentary form of communication could be established. There are also reports in exceptional patients of improvement in cognitive function with the implantation of thalamic-stimulating electrodes. It is prudent to avoid generalizations from these findings.

329e Examination of the Comatose Patient

S. Andrew Josephson

This chapter features a video illustrating the examination of a comatose patient. Proper techniques are demonstrated and supplemented with a discussion of interpretation of findings and implications for management. Also included is an overview of coma and its anatomic basis.

330 Neurologic Critical Care, Including Hypoxic-Ischemic Encephalopathy, and Subarachnoid Hemorrhage

J. Claude Hemphill, III, Wade S. Smith, Daryl R. Gress

Life-threatening neurologic illness may be caused by a primary disorder affecting any region of the neuraxis or may occur as a consequence of a systemic disorder such as hepatic failure, multisystem organ failure, or cardiac arrest (**Table 330-1**). Neurologic critical care focuses on preservation of neurologic tissue and prevention of secondary brain injury caused by ischemia, hemorrhage, edema, herniation, and elevated intracranial pressure (ICP). Management of other organ systems proceeds concurrently and may need to be modified in order to maintain the overall focus on neurologic issues.

PATHOPHYSIOLOGY

Brain Edema Swelling, or edema, of brain tissue occurs with many types of brain injury. The two principal types of edema are vasogenic and cytotoxic. *Vasogenic edema* refers to the influx of fluid and solutes into the brain through an incompetent blood-brain barrier (BBB). In the normal cerebral vasculature, endothelial tight junctions associated with astrocytes create an impermeable barrier (the BBB), through which access into the brain interstitium is dependent upon specific transport mechanisms. The BBB may be compromised in ischemia, trauma, infection, and metabolic derangements. *Vasogenic edema* results from abnormal permeability of the BBB, and typically develops rapidly following injury. *Cytotoxic edema* results from cellular swelling, membrane breakdown, and ultimately cell death. Clinically significant brain edema usually represents a combination of vasogenic and cytotoxic components. Edema can lead to increased ICP as well as tissue shifts and brain displacement or herniation from focal processes (**Chap. 328**). These tissue shifts can cause injury by mechanical distention and compression in addition to the ischemia of impaired perfusion consequent to the elevated ICP.

Ischemic Cascade and Cellular Injury When delivery of substrates, principally oxygen and glucose, is inadequate to sustain cellular function, a series of interrelated biochemical reactions known as the *ischemic cascade* is initiated (**see Fig. 446-2**). The release of excitatory amino acids, especially glutamate, leads to influx of calcium and sodium ions, which disrupt cellular homeostasis. An increased intracellular calcium concentration may activate proteases and lipases, which then lead to lipid peroxidation and free radical-mediated cell membrane injury. Cytotoxic edema ensues, and ultimately necrotic cell death and tissue infarction occur. This pathway to irreversible cell death is common to ischemic stroke, global cerebral ischemia, and traumatic brain injury.

Penumbra refers to areas of ischemic brain tissue that have not yet undergone irreversible infarction, implying that these regions are potentially salvageable if ischemia can be reversed. Factors that may exacerbate ischemic brain injury include systemic hypotension and hypoxia, which further reduce substrate delivery to vulnerable brain tissue, and fever, seizures, and hyperglycemia, which can increase cellular metabolism, outstripping compensatory processes. Clinically, these events are known as *secondary brain insults* because they lead to exacerbation of the primary brain injury. Prevention, identification, and treatment of secondary brain insults are fundamental goals of management.

An alternative pathway of cellular injury is *apoptosis*. This process implies programmed cell death, which may occur in the setting of ischemic stroke, global cerebral ischemia, traumatic brain injury, and possibly intracerebral hemorrhage. Apoptotic cell death can be distinguished histologically from the necrotic cell death of ischemia and is mediated through a different set of biochemical pathways; apoptotic

TABLE 330-1 NEUROLOGIC DISORDERS IN CRITICAL ILLNESS

Localization Along Neuroaxis	Syndrome	
Central Nervous System		
Brain: Cerebral hemispheres	Global encephalopathy	
	Delirium	
	Sepsis	
	Organ failure—hepatic, renal	
	Medication related—sedatives, hypnotics, analgesics, H ₂ blockers, antihypertensives	
	Drug overdose	
	Electrolyte disturbance—hyponatremia, hypoglycemia	
	Hypotension/hypoperfusion	
	Hypoxia	
	Meningitis	
	Subarachnoid hemorrhage	
	Wernicke's disease	
	Seizure—postictal or nonconvulsive status	
	Hypertensive encephalopathy	
	Hypothyroidism—myxedema	
	Focal deficits	
	Ischemic stroke	
	Tumor	
	Abscess, subdural empyema	
	Intraparenchymal hemorrhage	
	Subdural/epidural hematoma	
	Brainstem/cerebellum	Mass effect and compression
		Basilar artery thrombosis
Intraparenchymal hemorrhage		
Central pontine myelinolysis		
Spinal cord	Mass effect and compression	
	Disk herniation	
	Epidural hematoma	
	Ischemia—hypotension/embolic	
	Epidural abscess	
	Trauma	
Myelitis		
Peripheral Nervous System		
Peripheral nerve		
	Axonal	Critical illness polyneuropathy Neuromuscular blocking agent complications Metabolic disturbances, uremia, hyperglycemia Medication effects—chemotherapeutic, antiretroviral
Demyelinating		
		Guillain-Barré syndrome Chronic inflammatory demyelinating polyneuropathy
Neuromuscular junction	Prolonged effect of neuromuscular blockade	
Muscle		Medication effects—aminoglycosides Myasthenia gravis, Lambert-Eaton syndrome, botulism
		Critical illness myopathy Cachectic myopathy Acute necrotizing myopathy Thick-filament myopathy
		Electrolyte disturbances—hypokalemia/hyperkalemia, hypophosphatemia Rhabdomyolysis

cell death occurs without cerebral edema and therefore is often not seen on brain imaging. At present, interventions for prevention and treatment of apoptotic cell death remain less well defined than those for ischemia. **Excitotoxicity and mechanisms of cell death are discussed in more detail in Chap. 444e.**

Cerebral Perfusion and Autoregulation Brain tissue requires constant perfusion in order to ensure adequate delivery of substrate. The hemodynamic response of the brain has the capacity to preserve perfusion across a wide range of systemic blood pressures. Cerebral perfusion pressure (CPP), defined as the mean systemic arterial pressure (MAP) minus the ICP, provides the driving force for circulation across the capillary beds of the brain. *Autoregulation* refers to the physiologic response whereby cerebral blood flow (CBF) is regulated via alterations in cerebrovascular resistance in order to maintain perfusion over wide physiologic changes such as neuronal activation or changes in hemodynamic function. If systemic blood pressure drops, cerebral perfusion is preserved through vasodilation of arterioles in the brain; likewise, arteriolar vasoconstriction occurs at high systemic pressures to prevent hyperperfusion, resulting in fairly constant perfusion across a wide range of systemic blood pressures (Fig. 330-1). At the extreme limits of MAP or CPP (high or low), flow becomes directly related to perfusion pressure. These autoregulatory changes occur in the microcirculation and are mediated by vessels below the resolution of those seen on angiography. CBF is also strongly influenced by pH and PaCO₂. CBF increases with hypercapnia and acidosis and decreases with hypocapnia and alkalosis because of pH related changes in cerebral vascular resistance. This forms the basis for the use of hyperventilation to lower ICP, and this effect on ICP is mediated through a decrease in both CBF and intracranial blood volume. Cerebral autoregulation is a complex process critical to the normal homeostatic functioning of the brain, and this process may be disordered focally and unpredictably in disease states such as traumatic brain injury and severe focal cerebral ischemia.

Cerebrospinal Fluid and Intracranial Pressure The cranial contents consist essentially of brain, cerebrospinal fluid (CSF), and blood. CSF is produced principally in the choroid plexus of each lateral ventricle, exits the brain via the foramina of Luschka and Magendie, and flows over the cortex to be absorbed into the venous system along the superior sagittal sinus. In adults, approximately 150 mL of CSF are contained within the ventricles and surrounding the brain and spinal cord; the cerebral blood volume is also ~150 mL. The bony skull offers excellent protection for the brain but allows little tolerance for additional volume. Significant

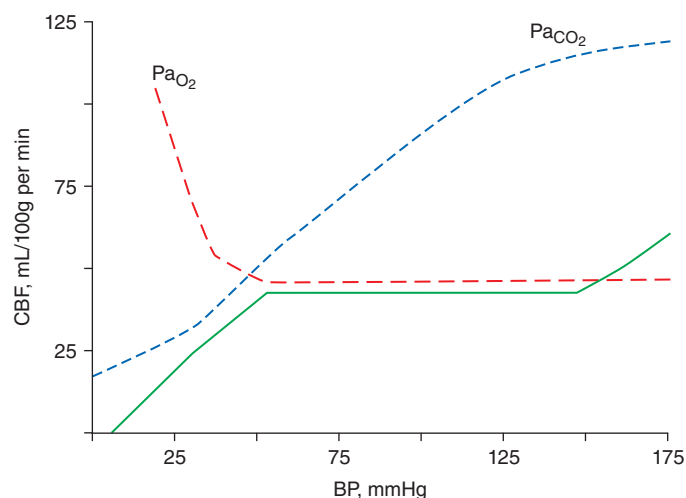


FIGURE 330-1 Autoregulation of cerebral blood flow (solid line).

Cerebral perfusion is constant over a wide range of systemic blood pressure. Perfusion is increased in the setting of hypoxia or hypercarbia. BP, blood pressure; CBF, cerebral blood flow. (Reprinted with permission from HM Shapiro: *Anesthesiology* 43:447, 1975. Copyright 1975, Lippincott Company.)

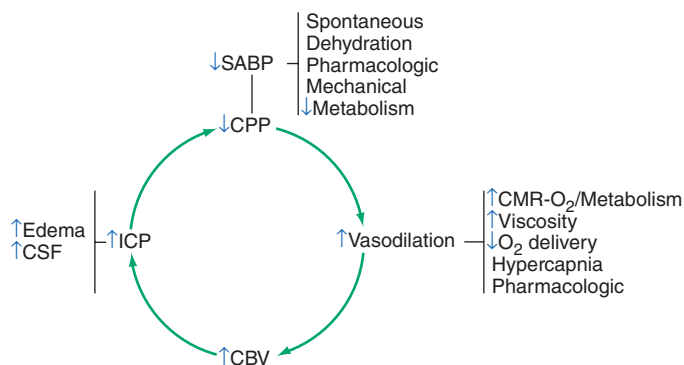


FIGURE 330-2 Ischemia and vasodilatation. Reduced cerebral perfusion pressure (CPP) leads to increased ischemia, vasodilation, increased intracranial pressure (ICP), and further reductions in CPP, a cycle leading to further neurologic injury. CBV, cerebral blood volume; CMR, cerebral metabolic rate; CSF, cerebrospinal fluid; SABP, systolic arterial blood pressure. (Adapted from MJ Rosner et al: *J Neurosurg* 83:949, 1995; with permission.)

increases in volume eventually result in increased ICP. Obstruction of CSF outflow, edema of cerebral tissue, or increases in volume from tumor or hematoma may increase ICP. Elevated ICP diminishes cerebral perfusion and can lead to tissue ischemia. Ischemia in turn may lead to vasodilation via autoregulatory mechanisms designed to restore cerebral perfusion. However, vasodilation also increases cerebral blood volume, which in turn then increases ICP, lowers CPP, and provokes further ischemia (Fig. 330-2). This vicious cycle is commonly seen in traumatic brain injury, massive intracerebral hemorrhage, and large hemispheric infarcts with significant tissue shifts.

APPROACH TO THE PATIENT: Severe Central Nervous System Dysfunction

Critically ill patients with severe central nervous system (CNS) dysfunction require rapid evaluation and intervention in order to limit primary and secondary brain injury. Initial neurologic evaluation should be performed concurrent with stabilization of basic respiratory, cardiac, and hemodynamic parameters. Significant barriers may exist to neurologic assessment in the critical care unit, including endotracheal intubation and the use of sedative or paralytic agents to facilitate procedures.

An impaired level of consciousness is common in critically ill patients. The essential first task in assessment is to determine whether the cause of dysfunction is related to a diffuse, usually metabolic, process or whether a focal, usually structural, process is implicated. Examples of diffuse processes include metabolic encephalopathies related to organ failure, drug overdose, or hypoxia-ischemia. Focal processes include ischemic and hemorrhagic stroke and traumatic brain injury, especially with intracranial hematomas. Because these two categories of disorders have fundamentally different causes, treatments, and prognoses, the initial focus is on making this distinction rapidly and accurately.

The approach to the comatose patient is discussed in Chap. 328; etiologies are listed in Table 328-1.

Minor focal deficits may be present on the neurologic examination in patients with metabolic encephalopathies. However, the finding of prominent focal signs such as pupillary asymmetry, hemiparesis, gaze palsy, or paraplegia should suggest the possibility of a structural lesion. All patients with a decreased level of consciousness associated with focal findings should undergo an urgent neuroimaging procedure, as should all patients with coma of unknown etiology. Computed tomography (CT) scanning is usually the most appropriate initial study because it can be performed

quickly in critically ill patients and demonstrates hemorrhage, hydrocephalus, and intracranial tissue shifts well. Magnetic resonance imaging (MRI) may provide more specific information in some situations, such as acute ischemic stroke (diffusion-weighted imaging [DWI]) and cerebral venous sinus thrombosis (magnetic resonance venography [MRV]). Any suggestion of trauma from the history or examination should alert the examiner to the possibility of cervical spine injury and prompt an imaging evaluation using plain x-rays, CT, or MRI.

Acute brainstem ischemia due to basilar artery thrombosis may cause brief episodes of spontaneous extensor posturing superficially resembling generalized seizures. Coma of sudden onset, accompanied by these movements and cranial nerve abnormalities, necessitates emergency imaging. A noncontrast CT scan of the brain may reveal a hyperdense basilar artery indicating thrombus in the vessel, and subsequent CT or MR angiography can assess basilar artery patency.

Acute brainstem ischemia due to basilar artery thrombosis may cause brief episodes of spontaneous extensor posturing superficially resembling generalized seizures. Coma of sudden onset, accompanied by these movements and cranial nerve abnormalities, necessitates emergency imaging. A noncontrast CT scan of the brain may reveal a hyperdense basilar artery indicating thrombus in the vessel, and subsequent CT or MR angiography can assess basilar artery patency.

Other diagnostic studies are best used in specific circumstances, usually when neuroimaging studies fail to reveal a structural lesion and the etiology of the altered mental state remains uncertain. Electroencephalography (EEG) can be important in the evaluation of critically ill patients with severe brain dysfunction. The EEG of metabolic encephalopathy typically reveals generalized slowing. One of the most important uses of EEG is to help exclude inapparent seizures, especially nonconvulsive status epilepticus. Untreated continuous or frequently recurrent seizures may cause neuronal injury, making the diagnosis and treatment of seizures crucial in this patient group. Lumbar puncture (LP) may be necessary to exclude infectious or inflammatory processes, and an elevated opening pressure may be an important clue to cerebral venous sinus thrombosis. In patients with coma or profound encephalopathy, it is preferable to perform a neuroimaging study prior to LP. If bacterial meningitis is suspected, an LP may be performed first or antibiotics may be empirically administered before the diagnostic studies are completed. Standard laboratory evaluation of critically ill patients should include assessment of serum electrolytes (especially sodium and calcium), glucose, renal and hepatic function, complete blood count, and coagulation. Serum or urine toxicology screens should be performed in patients with encephalopathy of unknown cause. EEG, LP, and other specific laboratory tests are most useful when the mechanism of the altered level of consciousness is uncertain; they are not routinely performed in clear-cut cases of stroke or traumatic brain injury.

Monitoring of ICP can be an important tool in selected patients. In general, patients who should be considered for ICP monitoring are those with primary neurologic disorders, such as stroke or traumatic brain injury, who are at significant risk for secondary brain injury due to elevated ICP and decreased CPP. Included are patients with the following: severe traumatic brain injury (Glasgow Coma Scale [GCS] score ≤ 8 [see Table 457e-2]); large tissue shifts from supratentorial ischemic or hemorrhagic stroke; or hydrocephalus from subarachnoid hemorrhage (SAH), intraventricular hemorrhage, or posterior fossa stroke. An additional disorder in which ICP monitoring can add important information is fulminant hepatic failure, in which elevated ICP may be treated with barbiturates or, eventually, liver transplantation. In general, ventriculostomy is preferable to ICP monitoring devices that are placed in the brain parenchyma, because ventriculostomy allows CSF drainage as a method of treating elevated ICP. However, parenchymal ICP monitoring is most appropriate for patients with diffuse edema and small ventricles (which may make ventriculostomy placement more difficult) or any degree of coagulopathy (in which ventriculostomy carries a higher risk of hemorrhagic complications) (Fig 330-3).

Treatment of Elevated ICP Elevated ICP may occur in a wide range of disorders, including head trauma, intracerebral hemorrhage, SAH with hydrocephalus, and fulminant hepatic failure. Because CSF and blood volume can be redistributed initially, by the time

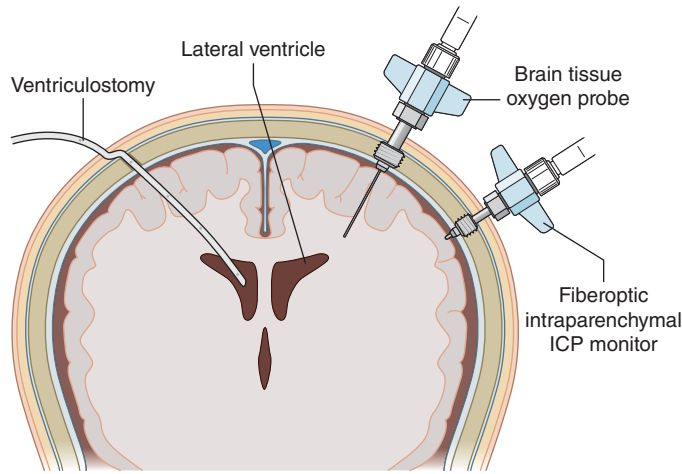


FIGURE 330-3 Intracranial pressure and brain tissue oxygen monitoring. A ventriculostomy allows for drainage of cerebrospinal fluid to treat elevated intracranial pressure (ICP). Fiberoptic ICP and brain tissue oxygen monitors are usually secured using a screwlike skull bolt. Cerebral blood flow and microdialysis probes (not shown) may be placed in a manner similar to the brain tissue oxygen probe.

elevated ICP occurs, intracranial compliance is severely impaired. At this point, any small increase in the volume of CSF, intravascular blood, edema, or a mass lesion may result in a significant increase in ICP and a decrease in cerebral perfusion. This is a fundamental mechanism of secondary ischemic brain injury and constitutes an emergency that requires immediate attention. In general, ICP should be maintained at <20 mmHg and CPP should be maintained at ≥ 60 mmHg.

Interventions to lower ICP are ideally based on the underlying mechanism responsible for the elevated ICP (Table 330-2). For

TABLE 330-2 STEPWISE APPROACH TO TREATMENT OF ELEVATED INTRACRANIAL PRESSURE (ICP)^a

Insert ICP monitor—ventriculostomy versus parenchymal device

General goals: maintain ICP <20 mmHg and CPP ≥ 60 mmHg. For ICP >20–25 mmHg for >5 min:

1. Elevate head of the bed; midline head position
2. Drain CSF via ventriculostomy (if in place)
3. Osmotherapy—mannitol 25–100 g q4h as needed (maintain serum osmolality <320 mosmol) or hypertonic saline (30 mL, 23.4% NaCl bolus)
4. Glucocorticoids—dexamethasone 4 mg q6h for vasogenic edema from tumor, abscess (avoid glucocorticoids in head trauma, ischemic and hemorrhagic stroke)
5. Sedation (e.g., morphine, propofol, or midazolam); add neuromuscular paralysis if necessary (patient will require endotracheal intubation and mechanical ventilation at this point, if not before)
6. Hyperventilation—to PaCO₂ 30–35 mmHg (short-term use or skip this step)
7. Pressor therapy—phenylephrine, dopamine, or norepinephrine to maintain adequate MAP to ensure CPP ≥ 60 mmHg (maintain euvolemia to minimize deleterious systemic effects of pressors). May adjust target CPP in individual patients based on autoregulation status.
8. Consider second-tier therapies for refractory elevated ICP
 - a. Decompressive craniectomy
 - b. High-dose barbiturate therapy (“pentobarb coma”)
 - c. Hypothermia to 33°C

^aThroughout ICP treatment algorithm, consider repeat head computed tomography to identify mass lesions amenable to surgical evacuation. May alter order of steps based on directed treatment to specific cause of elevated ICP.

Abbreviations: CPP, cerebral perfusion pressure; CSF, cerebrospinal fluid; MAP, mean arterial pressure; PaCO₂, arterial partial pressure of carbon dioxide.

example, in hydrocephalus from SAH, the principal cause of elevated ICP is impairment of CSF drainage. In this setting, ventricular drainage of CSF is likely to be sufficient and most appropriate. In head trauma and stroke, cytotoxic edema may be most responsible, and the use of osmotic agents such as mannitol or hypertonic saline becomes an appropriate early step. As described above, elevated ICP may cause tissue ischemia, and, if cerebral autoregulation is intact, the resulting vasodilation can lead to a cycle of worsening ischemia. Paradoxically, administration of vasopressor agents to increase mean arterial pressure may actually lower ICP by improving perfusion, thereby allowing autoregulatory vasoconstriction as ischemia is relieved and ultimately decreasing intracranial blood volume.

Early signs of elevated ICP include drowsiness and a diminished level of consciousness. Neuroimaging studies may reveal evidence of edema and mass effect. Hypotonic IV fluids should be avoided, and elevation of the head of the bed is recommended. Patients must be carefully observed for risk of aspiration and compromise of the airway as the level of alertness declines. Coma and unilateral pupillary changes are late signs and require immediate intervention. Emergent treatment of elevated ICP is most quickly achieved by intubation and hyperventilation, which causes vasoconstriction and reduces cerebral blood volume. To avoid provoking or worsening cerebral ischemia, hyperventilation, if used at all, is best administered only for short periods of time until a more definitive treatment can be instituted. Furthermore, the effects of hyperventilation on ICP are short-lived, often lasting only for several hours because of the buffering capacity of the cerebral interstitium, and rebound elevations of ICP may accompany abrupt discontinuation of hyperventilation. As the level of consciousness declines to coma, the ability to follow the neurologic status of the patient by examination lessens and measurement of ICP assumes greater importance. If a ventriculostomy device is in place, direct drainage of CSF to reduce ICP is possible. Finally, high-dose barbiturates, decompressive hemicraniectomy, and hypothermia are sometimes used for refractory elevations of ICP, although these have significant side effects and have not been proven to improve outcome.

Secondary Brain Insults Patients with primary brain injuries, whether due to trauma or stroke, are at risk for ongoing secondary ischemic brain injury. Because secondary brain injury can be a major determinant of a poor outcome, strategies for minimizing secondary brain insults are an integral part of the critical care of all patients. Although elevated ICP may lead to secondary ischemia, most secondary brain injury is mediated through other clinical events that exacerbate the ischemic cascade already initiated by the primary brain injury. Episodes of secondary brain insults are usually not associated with apparent neurologic worsening. Rather, they lead to cumulative injury limiting eventual recovery, which manifests as a higher mortality rate or worsened long-term functional outcome. Thus, close monitoring of vital signs is important, as is early intervention to prevent secondary ischemia. Avoiding hypotension and hypoxia is critical, as significant hypotensive events (systolic blood pressure <90 mmHg) as short as 10 min in duration have been shown to adversely influence outcome after traumatic brain injury. Even in patients with stroke or head trauma who do not require ICP monitoring, close attention to adequate cerebral perfusion is warranted. Hypoxia (pulse oximetry saturation <90%), particularly in combination with hypotension, also leads to secondary brain injury. Likewise, fever and hyperglycemia both worsen experimental ischemia and have been associated with worsened clinical outcome after stroke and head trauma. Aggressive control of fever with a goal of normothermia is warranted but may be difficult to achieve with antipyretic medications and cooling blankets. The value of newer surface or intravascular temperature control devices for the management of refractory fever is under investigation. The use of IV insulin infusion is encouraged for control of hyperglycemia because this allows better regulation of serum glucose levels than SC insulin. A reasonable goal is to

maintain the serum glucose level at <10.0 mmol/L (<180 mg/dL), although episodes of hypoglycemia appear equally detrimental and the optimal targets remain uncertain. New cerebral monitoring tools that allow continuous evaluation of brain tissue oxygen tension, CBF, and metabolism (via microdialysis) may further improve the management of secondary brain injury.

CRITICAL CARE DISORDERS OF THE CENTRAL NERVOUS SYSTEM

HYPOXIC-ISCHEMIC ENCEPHALOPATHY

This occurs from lack of delivery of oxygen to the brain because of extreme hypotension (hypoxia-ischemia) or hypoxia due to respiratory failure. Causes include myocardial infarction, cardiac arrest, shock, asphyxiation, paralysis of respiration, and carbon monoxide or cyanide poisoning. In some circumstances, hypoxia may predominate. Carbon monoxide and cyanide poisoning are sometimes termed *histotoxic hypoxia* because they cause a direct impairment of the respiratory chain.

Clinical Manifestations Mild degrees of pure hypoxia, such as occur at high altitudes, cause impaired judgment, inattentiveness, motor incoordination, and, at times, euphoria. However, with hypoxia-ischemia, such as occurs with circulatory arrest, consciousness is lost within seconds. If circulation is restored within 3–5 min, full recovery may occur, but if hypoxia-ischemia lasts beyond 3–5 min, some degree of permanent cerebral damage usually results. Except in extreme cases, it may be difficult to judge the precise degree of hypoxia-ischemia, and some patients make a relatively full recovery after even 8–10 min of global cerebral ischemia. The brain is more tolerant to pure hypoxia than it is to hypoxia-ischemia. For example, a PaO_2 as low as 20 mmHg (2.7 kPa) can be well tolerated if it develops gradually and normal blood pressure is maintained, whereas short durations of very low or absent cerebral circulation usually result in permanent impairment.

Clinical examination at different time points after a hypoxic-ischemic insult (especially cardiac arrest) is useful in assessing prognosis for long-term neurologic outcome. The prognosis is better for patients with intact brainstem function, as indicated by normal pupillary light responses and intact oculoccephalic (doll's eyes), oculovestibular (caloric), and corneal reflexes. Absence of these reflexes and the presence of persistently dilated pupils that do not react to light are grave prognostic signs. A low likelihood of a favorable outcome from hypoxic-ischemic coma is strongly suggested by an absent pupillary light reflex or extensor or absent motor response to pain on day 3 following the injury, excluding patients with metabolic disturbances and those treated with high-dose barbiturates or hypothermia, which confound interpretation of these signs. Electrophysiologically, the bilateral absence of the N20 component of the somatosensory evoked potential (SSEP) in the first several days also conveys a poor prognosis. A very elevated serum level (>33 $\mu\text{g/L}$) of the biochemical marker neuron-specific enolase (NSE) is indicative of brain damage after resuscitation from cardiac arrest and predicts a poor outcome. However, at present, SSEPs and NSE levels may be difficult to obtain in a timely fashion, with SSEP testing requiring substantial expertise in interpretation and NSE measurements not yet standardized. Recent studies suggest that the administration of mild hypothermia after cardiac arrest (see "Treatment") may affect the time points when these clinical and electrophysiologic predictors become reliable in identifying patients with a very low likelihood of clinically meaningful recovery. For example, the false-positive rate for incorrect prediction of poor neurologic outcome may be as high as 21% (95% confidence interval [CI] 8–43%) for patients treated with mild hypothermia who exhibit 3-day motor function no better than extensor posturing. Long-term consequences of hypoxic-ischemic encephalopathy include persistent coma or a vegetative state (Chap. 328), dementia, visual agnosia (Chap. 36), parkinsonism, choreoathetosis, cerebellar ataxia, myoclonus, seizures, and an amnesic state, which may be a consequence of selective damage to the hippocampus.

Pathology Principal histologic findings are extensive multifocal or diffuse laminar cortical necrosis (Fig. 330-4), with frequent involvement

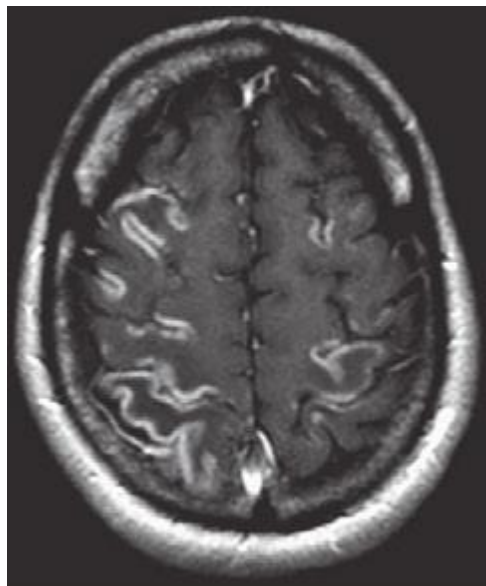


FIGURE 330-4 Cortical laminar necrosis in hypoxic-ischemic encephalopathy. T1-weighted postcontrast magnetic resonance imaging shows cortical enhancement in a watershed distribution consistent with laminar necrosis.

of the hippocampus. The hippocampal CA1 neurons are vulnerable to even brief episodes of hypoxia-ischemia, perhaps explaining why selective persistent memory deficits may occur after brief cardiac arrest. Scattered small areas of infarction or neuronal loss may be present in the basal ganglia, hypothalamus, or brainstem. In some cases, extensive bilateral thalamic scarring may affect pathways that mediate arousal, and this pathology may be responsible for the persistent vegetative state. A specific form of hypoxic-ischemic encephalopathy, so-called watershed infarcts, occurs at the distal territories between the major cerebral arteries and can cause cognitive deficits, including visual agnosia, and weakness that is greater in proximal than in distal muscle groups.

Diagnosis Diagnosis is based on the history of a hypoxic-ischemic event such as cardiac arrest. Blood pressure <70 mmHg systolic or PaO_2 <40 mmHg is usually necessary, although both absolute levels and duration of exposure are important determinants of cellular injury. Carbon monoxide intoxication can be confirmed by measurement of carboxyhemoglobin and is suggested by a cherry red color of the venous blood and skin, although the latter is an inconsistent clinical finding.

TREATMENT HYPOXIC-ISCHEMIC ENCEPHALOPATHY

Treatment should be directed at restoration of normal cardiorespiratory function. This includes securing a clear airway, ensuring adequate oxygenation and ventilation, and restoring cerebral perfusion, whether by cardiopulmonary resuscitation, fluid, pressors, or cardiac pacing. Hypothermia may target the neuronal cell injury cascade and has substantial neuroprotective properties in experimental models of brain injury. In two trials, mild hypothermia (33°C) improved functional outcome in patients who remained comatose after resuscitation from a cardiac arrest. Treatment was initiated within minutes of cardiac resuscitation and continued for 12 h in one study and 24 h in the other. Potential complications of hypothermia include coagulopathy and an increased risk of infection. Based on these studies, the International Liaison Committee on Resuscitation issued the following advisory statement: "Unconscious adult patients with spontaneous circulation after out-of-hospital cardiac arrest should be cooled to 32°–34°C for 12–24 h when the initial rhythm was ventricular fibrillation. Such cooling may also be beneficial for other rhythms or in-hospital cardiac arrest."

1782 Severe carbon monoxide intoxication may be treated with hyperbaric oxygen. Anticonvulsants may be needed to control seizures, although these are not usually given prophylactically. Posthypoxic myoclonus may respond to oral administration of clonazepam at doses of 1.5–10 mg daily or valproate at doses of 300–1200 mg daily in divided doses. Myoclonic status epilepticus within 24 h after a primary circulatory arrest generally portends a very poor prognosis, even if seizures are controlled.

Carbon monoxide and cyanide intoxication can also cause a delayed encephalopathy. Little clinical impairment is evident when the patient first regains consciousness, but a parkinsonian syndrome characterized by akinesia and rigidity without tremor may develop. Symptoms can worsen over months, accompanied by increasing evidence of damage in the basal ganglia as seen on both CT and MRI.

METABOLIC ENCEPHALOPATHIES

Altered mental states, variously described as confusion, delirium, disorientation, and encephalopathy, are present in many patients with severe illness in an intensive care unit (ICU). Older patients are particularly vulnerable to delirium, a confusional state characterized by disordered perception, frequent hallucinations, delusions, and sleep disturbance. This is often attributed to medication effects, sleep deprivation, pain, and anxiety. The presence of delirium is associated with worsened outcome in critically ill patients, even in those without an identifiable CNS pathology such as stroke or brain trauma. In these patients, the cause of delirium is often multifactorial, resulting from organ dysfunction, sepsis, and especially the use of medications given to treat pain, agitation, or anxiety. Critically ill patients are often treated with a variety of sedative and analgesic medications, including opiates, benzodiazepines, neuroleptics, and sedative-anesthetic medications, such as propofol. In critically ill patients requiring sedation, use of the centrally acting α_2 agonist dexmedetomidine may reduce delirium and shorten the duration of mechanical ventilation compared to the use of benzodiazepines such as lorazepam or midazolam. The presence of family members in the ICU may also help to calm and orient agitated patients, and in severe cases, low doses of neuroleptics (e.g., haloperidol 0.5–1 mg) can be useful. Current strategies focus on limiting the use of sedative medications when this can be done safely.

In the ICU setting, several metabolic causes of an altered level of consciousness predominate. Hypercarbic encephalopathy can present with headache, confusion, stupor, or coma. Hypoventilation syndrome occurs most frequently in patients with a history of chronic CO_2 retention who are receiving oxygen therapy for emphysema or chronic pulmonary disease (Chap. 318). The elevated PaCO_2 leading to CO_2 narcosis may have a direct anesthetic effect, and cerebral vasodilation from increased PaCO_2 can lead to increased ICP. Hepatic encephalopathy is suggested by asterixis and can occur in chronic liver failure or acute fulminant hepatic failure. Both hyperglycemia and hypoglycemia can cause encephalopathy, as can hyponatremia and hypernatremia. Confusion, impairment of eye movements, and gait ataxia are the hallmarks of acute Wernicke's disease (see below).

SEPSIS-ASSOCIATED ENCEPHALOPATHY

Pathogenesis In patients with sepsis, the systemic response to infectious agents leads to the release of circulating inflammatory mediators that appear to contribute to encephalopathy. Critical illness, in association with the systemic inflammatory response syndrome (SIRS), can lead to multisystem organ failure. This syndrome can occur in the setting of apparent sepsis, severe burns, or trauma, even without clear identification of an infectious agent. Many patients with critical illness, sepsis, or SIRS develop encephalopathy without obvious explanation. This condition is broadly termed *sepsis-associated encephalopathy*. Although the specific mediators leading to neurologic dysfunction remain uncertain, it is clear that the encephalopathy is not simply the result of metabolic derangements of multiorgan failure. The cytokines tumor necrosis factor, interleukin (IL)-1, IL-2, and IL-6 are thought to play a role in this syndrome.

Diagnosis Sepsis-associated encephalopathy presents clinically as a diffuse dysfunction of the brain without prominent focal findings. Confusion, disorientation, agitation, and fluctuations in level of alertness are typical. In more profound cases, especially with hemodynamic compromise, the decrease in level of alertness can be more prominent, at times resulting in coma. Hyperreflexia and frontal release signs such as a grasp or snout reflex (Chap. 36) can be seen. Abnormal movements such as myoclonus, tremor, or asterixis can occur. Sepsis-associated encephalopathy is quite common, occurring in the majority of patients with sepsis and multisystem organ failure. Diagnosis is often difficult because of the multiple potential causes of neurologic dysfunction in critically ill patients and requires exclusion of structural, metabolic, toxic, and infectious (e.g., meningitis or encephalitis) causes. The mortality rate of patients with sepsis-associated encephalopathy severe enough to produce coma approaches 50%, although this principally reflects the severity of the underlying critical illness and is not a direct result of the encephalopathy. Patients dying from severe sepsis or septic shock may have elevated levels of the serum brain injury biomarker S-100 β and neuropathologic findings of neuronal apoptosis and cerebral ischemic injury. Successful treatment of the underlying critical illness almost always results in substantial improvement of the encephalopathy. However, although severe disability to the level of chronic vegetative or minimally conscious states is uncommon, long-term cognitive dysfunction clinically similar to dementia is being increasingly recognized in some survivors.

CENTRAL PONTINE MYELINOLYSIS

This disorder typically presents in a devastating fashion as quadriplegia and pseudobulbar palsy. Predisposing factors include severe underlying medical illness or nutritional deficiency; most cases are associated with rapid correction of hyponatremia or with hyperosmolar states. The pathology consists of demyelination without inflammation in the base of the pons, with relative sparing of axons and nerve cells. MRI is useful in establishing the diagnosis (Fig. 330-5) and may also identify partial forms that present as confusion, dysarthria, and/or disturbances of conjugate gaze without quadriplegia. Occasional cases present with lesions outside of the brainstem. Therapeutic guidelines for the restoration of severe hyponatremia should aim for gradual correction, i.e., by ≤ 10 mmol/L (10 meq/L) within 24 h and 20 mmol/L (20 meq/L) within 48 h.

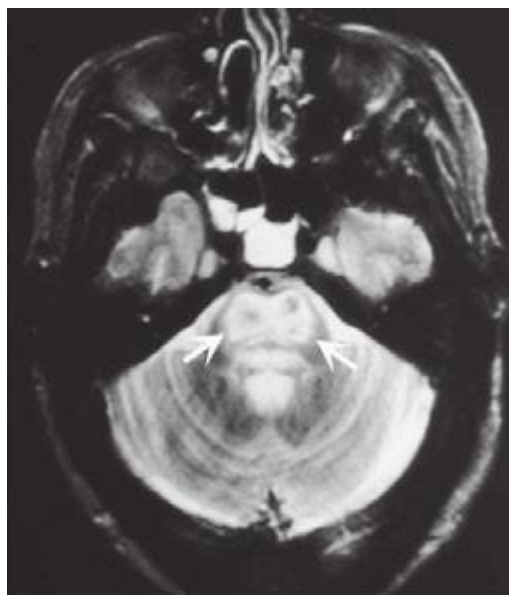


FIGURE 330-5 Central pontine myelinolysis. Axial T2-weighted magnetic resonance scan through the pons reveals a symmetric area of abnormal high signal intensity within the basis pontis (arrows).

WERNICKE'S DISEASE

Wernicke's disease is a common and preventable disorder due to a deficiency of thiamine (**Chap. 96e**). In the United States, alcoholics account for most cases, but patients with malnutrition due to hyperemesis, starvation, renal dialysis, cancer, AIDS, or rarely gastric surgery are also at risk. The characteristic clinical triad is that of ophthalmoplegia, ataxia, and global confusion. However, only one-third of patients with acute Wernicke's disease present with the classic clinical triad. Most patients are profoundly disoriented, indifferent, and inattentive, although rarely they have an agitated delirium related to ethanol withdrawal. If the disease is not treated, stupor, coma, and death may ensue. Ocular motor abnormalities include horizontal nystagmus on lateral gaze, lateral rectus palsy (usually bilateral), conjugate gaze palsies, and rarely ptosis. Gait ataxia probably results from a combination of polyneuropathy, cerebellar involvement, and vestibular paresis. The pupils are usually spared, but they may become miotic with advanced disease.

Wernicke's disease is usually associated with other manifestations of nutritional disease, such as polyneuropathy. Rarely, amblyopia or myelopathy occurs. Tachycardia and postural hypotension may be related to impaired function of the autonomic nervous system or to the coexistence of cardiovascular beriberi. Patients who recover show improvement in ocular palsies within hours after the administration of thiamine, but horizontal nystagmus may persist. Ataxia improves more slowly than the ocular motor abnormalities. Approximately half recover incompletely and are left with a slow, shuffling, wide-based gait and an inability to tandem walk. Apathy, drowsiness, and confusion improve more gradually. As these symptoms recede, an amnesic state with impairment in recent memory and learning may become more apparent (*Korsakoff's psychosis*). Korsakoff's psychosis is frequently persistent; the residual mental state is characterized by gaps in memory, confabulation, and disordered temporal sequencing.

Pathology Periventricular lesions surround the third ventricle, aqueduct, and fourth ventricle, with petechial hemorrhages in occasional acute cases and atrophy of the mammillary bodies in most chronic cases. There is frequently endothelial proliferation, demyelination, and some neuronal loss. These changes may be detected by MRI scanning (**Fig. 330-6**). The amnesic defect is related to lesions in the dorsal medial nuclei of the thalamus.

Pathogenesis Thiamine is a cofactor of several enzymes, including transketolase, pyruvate dehydrogenase, and α -ketoglutarate

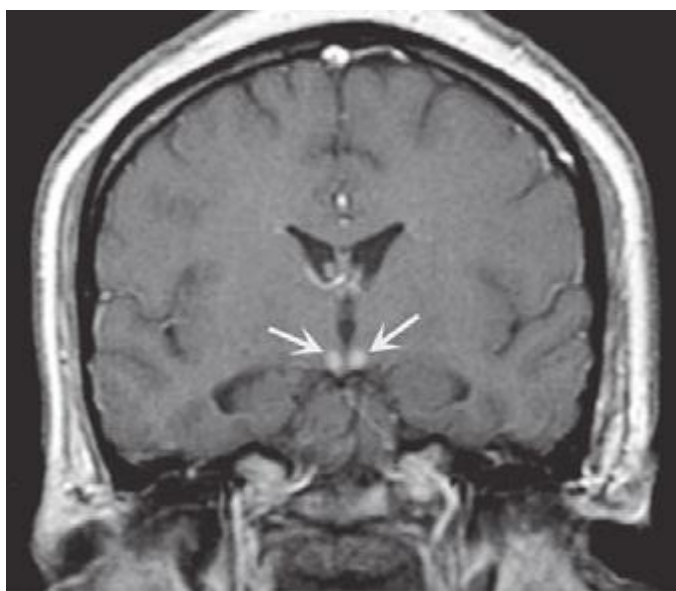


FIGURE 330-6 Wernicke's disease. Coronal T1-weighted postcontrast magnetic resonance imaging reveals abnormal enhancement of the mammillary bodies (arrows), typical of acute Wernicke's encephalopathy.

dehydrogenase. Thiamine deficiency produces a diffuse decrease in cerebral glucose utilization and results in mitochondrial damage. Glutamate accumulates due to impairment of α -ketoglutarate dehydrogenase activity and, in combination with the energy deficiency, may result in excitotoxic cell damage.

TREATMENT WERNICKE'S DISEASE

Wernicke's disease is a medical emergency and requires immediate administration of thiamine, in a dose of 100 mg either IV or IM. The dose should be given daily until the patient resumes a normal diet and should be begun prior to treatment with IV glucose solutions. Larger doses, 100 mg four times a day or more, have been advocated by some. Glucose infusions may precipitate Wernicke's disease in a previously unaffected patient or cause a rapid worsening of an early form of the disease. For this reason, thiamine should be administered to all alcoholic patients requiring parenteral glucose.

CRITICAL CARE DISORDERS OF THE PERIPHERAL NERVOUS SYSTEM

Critical illness with disorders of the peripheral nervous system (PNS) arises in two contexts: (1) primary neurologic diseases that require critical care interventions such as intubation and mechanical ventilation, and (2) secondary PNS manifestations of systemic critical illness, often involving multisystem organ failure. The former include acute polyneuropathies such as Guillain-Barré syndrome (**Chap. 460**), neuromuscular junction disorders including myasthenia gravis (**Chap. 461**) and botulism (**Chap. 178**), and primary muscle disorders such as polymyositis (**Chap. 462e**). The latter result either from the systemic disease itself or as a consequence of interventions.

General principles of respiratory evaluation in patients with PNS involvement, regardless of cause, include assessment of pulmonary mechanics, such as maximal inspiratory force (MIF) and vital capacity (VC), and evaluation of strength of bulbar muscles. Regardless of the cause of weakness, endotracheal intubation should be considered when the MIF falls to <-25 cmH₂O or the VC is <1 L. Also, patients with severe palatal weakness may require endotracheal intubation in order to prevent acute upper airway obstruction or recurrent aspiration. Arterial blood gases and oxygen saturation from pulse oximetry are used to follow patients with potential respiratory compromise from PNS dysfunction. However, intubation and mechanical ventilation should be undertaken based on clinical assessment rather than waiting until oxygen saturation drops or CO₂ retention develops from hypoventilation. Noninvasive mechanical ventilation may be considered initially in lieu of endotracheal intubation but is generally insufficient in patients with severe bulbar weakness or ventilatory failure with hypercarbia. **Principles of mechanical ventilation are discussed in Chap. 323.**

NEUROPATHY

Although encephalopathy may be the most obvious neurologic dysfunction in critically ill patients, dysfunction of the PNS is also quite common. It is typically present in patients with prolonged critical illnesses lasting several weeks and involving sepsis; clinical suspicion is aroused when there is failure to wean from mechanical ventilation despite improvement of the underlying sepsis and critical illness. *Critical illness polyneuropathy* refers to the most common PNS complication related to critical illness; it is seen in the setting of prolonged critical illness, sepsis, and multisystem organ failure. Neurologic findings include diffuse weakness, decreased reflexes, and distal sensory loss. Electrophysiologic studies demonstrate a diffuse, symmetric, distal axonal sensorimotor neuropathy, and pathologic studies have confirmed axonal degeneration. The precise mechanism of critical illness polyneuropathy remains unclear, but circulating factors such as cytokines, which are associated with sepsis and SIRS, are thought to play a role. It has been reported that up to 70% of patients with the sepsis syndrome have some degree of neuropathy, although far fewer

1784 have a clinical syndrome profound enough to cause severe respiratory muscle weakness requiring prolonged mechanical ventilation or resulting in failure to wean. Aggressive glycemic control with insulin infusions appears to decrease the risk of critical illness polyneuropathy. Treatment is otherwise supportive, with specific intervention directed at treating the underlying illness. Although spontaneous recovery is usually seen, the time course may extend over weeks to months and necessitate long-term ventilatory support and care even after the underlying critical illness has resolved.

DISORDERS OF NEUROMUSCULAR TRANSMISSION

A defect in neuromuscular transmission may be a source of weakness in critically ill patients. Botulism (**Chap. 178**) may be acquired by ingesting botulinum toxin from improperly stored food or may arise from an anaerobic abscess from *Clostridium botulinum* (wound botulism). Infants can present with generalized weakness from gut-derived *Clostridium* infection, especially if they are fed honey. Diplopia and dysphagia are early signs of foodborne botulism. Treatment is mostly supportive, although use of antitoxin early in the course may limit the duration of the neuromuscular blockade. General ICU care is similar to patients with Guillain-Barré syndrome or myasthenia gravis with focused care to avoid ulcer formation at pressure points, deep venous thromboprophylaxis, and infection prevention. Public health officers should be rapidly informed when the diagnosis is made to prevent further exposure to others from the tainted food or source of wound botulism (such as injection drug use).

Undiagnosed myasthenia gravis (**Chap. 461**) may be a consideration in weak ICU patients; however, persistent weakness secondary to impaired neuromuscular junction transmission is almost always due to administration of drugs. A number of medications impair neuromuscular transmission; these include antibiotics, especially aminoglycosides, and beta-blocking agents. In the ICU, the nondepolarizing neuromuscular blocking agents (nd-NMBAs), also known as muscle relaxants, are most commonly responsible. Included in this group of drugs are such agents as pancuronium, vecuronium, rocuronium, and cisatracurium. They are often used to facilitate mechanical ventilation or other critical care procedures, but with prolonged use persistent neuromuscular blockade may result in weakness even after discontinuation of these agents hours or days earlier. Risk factors for this prolonged action of neuromuscular blocking agents include female sex, metabolic acidosis, and renal failure.

Prolonged neuromuscular blockade does not appear to produce permanent damage to the PNS. Once the offending medications are discontinued, full strength is restored, although this may take days. In general, the lowest dose of neuromuscular blocking agent should be used to achieve the desired result and, when these agents are used in the ICU, a peripheral nerve stimulator should be used to monitor neuromuscular junction function.

MYOPATHY

Critically ill patients, especially those with sepsis, frequently develop muscle weakness and wasting, often in the face of seemingly adequate nutritional support. *Critical illness myopathy* is an overall term that describes several different discrete muscle disorders that may occur in critically ill patients. The assumption has been that a catabolic myopathy may develop as a result of multiple factors, including elevated cortisol and catecholamine release and other circulating factors induced by the SIRS. In this syndrome, known as *cachectic myopathy*, serum creatine kinase levels and electromyography (EMG) are normal. Muscle biopsy shows type II fiber atrophy. Panfascicular muscle fiber necrosis may also occur in the setting of profound sepsis. This less common *acute necrotizing intensive care myopathy* is characterized clinically by weakness progressing to a profound level over just a few days. There may be associated elevations in serum creatine kinase and urine myoglobin. Both EMG and muscle biopsy may be normal initially but eventually show abnormal spontaneous activity and panfascicular necrosis with an accompanying inflammatory reaction. Acute rhabdomyolysis can occur from alcohol ingestion or from compartment syndromes.

A *thick-filament myopathy* may occur in the setting of glucocorticoid and nd-NMBA use. The most frequent scenario in which this is encountered is the asthmatic patient who requires high-dose glucocorticoids and nd-NMBA to facilitate mechanical ventilation. This muscle disorder is not due to prolonged action of nd-NMBAs at the neuromuscular junction but, rather, is an actual myopathy with muscle damage; it has occasionally been described with high-dose glucocorticoid use or sepsis alone. Clinically this syndrome is most often recognized when a patient fails to wean from mechanical ventilation despite resolution of the primary pulmonary process. Pathologically, there may be loss of thick (myosin) filaments. Thick-filament critical illness myopathy has a good prognosis. If patients survive their underlying critical illness, the myopathy invariably improves and most patients return to normal. However, because this syndrome is a result of true muscle damage, not just prolonged blockade at the neuromuscular junction, this process may take weeks or months, and tracheotomy with prolonged ventilatory support may be necessary. Some patients do have residual long-term weakness, with atrophy and fatigue limiting ambulation. At present, it is unclear how to prevent this myopathic complication, except by avoiding use of nd-NMBAs, a strategy not always possible. Monitoring with a peripheral nerve stimulator can help to avoid the overuse of these agents. However, this is more likely to prevent the complication of prolonged neuromuscular junction blockade than it is to prevent this myopathy.

SUBARACHNOID HEMORRHAGE

Subarachnoid hemorrhage (SAH) renders the brain critically ill from both primary and secondary brain insults. Excluding head trauma, the most common cause of SAH is rupture of a saccular aneurysm. Other causes include bleeding from a vascular malformation (arteriovenous malformation or dural arteriovenous fistula) and extension into the subarachnoid space from a primary intracerebral hemorrhage. Some idiopathic SAHs are localized to the perimesencephalic cisterns and are benign; they probably have a venous or capillary source, and angiography is unrevealing.

SACCULAR ("BERRY") ANEURYSM

Autopsy and angiography studies have found that about 2% of adults harbor intracranial aneurysms, for a prevalence of 4 million persons in the United States; the aneurysm will rupture, producing SAH, in 25,000–30,000 cases per year. For patients who arrive alive at hospital, the mortality rate over the next month is about 45%. Of those who survive, more than half are left with major neurologic deficits as a result of the initial hemorrhage, cerebral vasospasm with infarction, or hydrocephalus. If the patient survives but the aneurysm is not obliterated, the rate of rebleeding is about 20% in the first 2 weeks, 30% in the first month, and about 3% per year afterward. Given these alarming figures, the major therapeutic emphasis is on preventing the predictable early complications of the SAH.

Unruptured, asymptomatic aneurysms are much less dangerous than a recently ruptured aneurysm. The annual risk of rupture for aneurysms <10 mm in size is ~0.1%, and for aneurysms ≥10 mm in size is ~0.5–1%; the surgical morbidity rate far exceeds these percentages. Because of the longer length of exposure to risk of rupture, younger patients with aneurysms >10 mm in size may benefit from prophylactic treatment. As with the treatment of asymptomatic carotid stenosis, this risk-benefit ratio strongly depends on the complication rate of treatment.

Giant aneurysms, those >2.5 cm in diameter, occur at the same sites (see below) as small aneurysms and account for 5% of cases. The three most common locations are the terminal internal carotid artery, middle cerebral artery (MCA) bifurcation, and top of the basilar artery. Their risk of rupture is ~6% in the first year after identification and may remain high indefinitely. They often cause symptoms by compressing the adjacent brain or cranial nerves.

Mycotic aneurysms are usually located distal to the first bifurcation of major arteries of the circle of Willis. Most result from infected emboli due to bacterial endocarditis causing septic degeneration of arteries and subsequent dilation and rupture. Whether these lesions

should be sought and repaired prior to rupture or left to heal spontaneously with antibiotic treatment is controversial.

Pathophysiology Saccular aneurysms occur at the bifurcations of the large- to medium-sized intracranial arteries; rupture is into the subarachnoid space in the basal cisterns and often into the parenchyma of the adjacent brain. Approximately 85% of aneurysms occur in the anterior circulation, mostly on the circle of Willis. About 20% of patients have multiple aneurysms, many at mirror sites bilaterally. As an aneurysm develops, it typically forms a neck with a dome. The length of the neck and the size of the dome vary greatly and are important factors in planning neurosurgical obliteration or endovascular embolization. The arterial internal elastic lamina disappears at the base of the neck. The media thins, and connective tissue replaces smooth-muscle cells. At the site of rupture (most often the dome), the wall thins, and the tear that allows bleeding is often ≤ 0.5 mm long. Aneurysm size and site are important in predicting risk of rupture. Those >7 mm in diameter and those at the top of the basilar artery and at the origin of the posterior communicating artery are at greater risk of rupture.

Clinical Manifestations Most unruptured intracranial aneurysms are completely asymptomatic. Symptoms are usually due to rupture and resultant SAH, although some unruptured aneurysms present with mass effect on cranial nerves or brain parenchyma. At the moment of aneurysmal rupture with major SAH, the ICP suddenly rises. This may account for the sudden transient loss of consciousness that occurs in nearly half of patients. Sudden loss of consciousness may be preceded by a brief moment of excruciating headache, but most patients first complain of headache upon regaining consciousness. In 10% of cases, aneurysmal bleeding is severe enough to cause loss of consciousness for several days. In $\sim 45\%$ of cases, severe headache associated with exertion is the presenting complaint. The patient often calls the headache “the worst headache of my life”; however, the most important characteristic is sudden onset. Occasionally, these ruptures may present as headache of only moderate intensity or as a change in the patient’s usual headache pattern. The headache is usually generalized, often with neck stiffness, and vomiting is common.

Although sudden headache in the absence of focal neurologic symptoms is the hallmark of aneurysmal rupture, focal neurologic deficits may occur. Anterior communicating artery or MCA bifurcation aneurysms may rupture into the adjacent brain or subdural space and form a hematoma large enough to produce mass effect. The deficits that result can include hemiparesis, aphasia, and abulia.

Occasionally, prodromal symptoms suggest the location of a progressively enlarging unruptured aneurysm. A third cranial nerve palsy, particularly when associated with pupillary dilation, loss of ipsilateral (but retained contralateral) light reflex, and focal pain above or behind the eye, may occur with an expanding aneurysm at the junction of the posterior communicating artery and the internal carotid artery. A sixth nerve palsy may indicate an aneurysm in the cavernous sinus, and visual field defects can occur with an expanding supraclinoid carotid or anterior cerebral artery aneurysm. Occipital and posterior cervical pain may signal a posterior inferior cerebellar artery or anterior inferior cerebellar artery aneurysm (**Chap. 446**). Pain in or behind the eye and in the low temple can occur with an expanding MCA aneurysm. Thunderclap headache is a variant of migraine that simulates an SAH. Before concluding that a patient with sudden, severe headache has thunderclap migraine, a definitive workup for aneurysm or other intracranial pathology is required.

Aneurysms can undergo small ruptures and leaks of blood into the subarachnoid space, so-called *sentinel bleeds*. Sudden unexplained headache at any location should raise suspicion of SAH and be investigated, because a major hemorrhage may be imminent.

The initial clinical manifestations of SAH can be graded using the Hunt-Hess or World Federation of Neurosurgical Societies classification schemes (**Table 330-3**). For ruptured aneurysms, prognosis for good outcomes falls as the grade increases. For example, it is unusual for a Hunt-Hess grade 1 patient to die if the aneurysm is treated, but the mortality rate for grade 4 and 5 patients may be as high as 80%.

TABLE 330-3 GRADING SCALES FOR SUBARACHNOID HEMORRHAGE

Grade	Hunt-Hess Scale	World Federation of Neurosurgical Societies (WFNS) Scale
1	Mild headache, normal mental status, no cranial nerve or motor findings	GCS ^a score 15, no motor deficits
2	Severe headache, normal mental status, may have cranial nerve deficit	GCS score 13–14, no motor deficits
3	Somnolent, confused, may have cranial nerve or mild motor deficit	GCS score 13–14, with motor deficits
4	Stupor, moderate to severe motor deficit, may have intermittent reflex posturing	GCS score 7–12, with or without motor deficits
5	Coma, reflex posturing or flaccid	GCS score 3–6, with or without motor deficits

^aGlasgow Coma Scale; see **Table 457e-1**.

Delayed Neurologic Deficits There are four major causes of delayed neurologic deficits: rerupture, hydrocephalus, vasospasm, and hyponatremia.

1. **Rerupture.** The incidence of rerupture of an untreated aneurysm in the first month following SAH is $\sim 30\%$, with the peak in the first 7 days. Rerupture is associated with a 60% mortality rate and poor outcome. Early treatment eliminates this risk.
2. **Hydrocephalus.** Acute hydrocephalus can cause stupor and coma and can be mitigated by placement of an external ventricular drain. More often, subacute hydrocephalus may develop over a few days or weeks and causes progressive drowsiness or slowed mentation (abulia) with incontinence. Hydrocephalus is differentiated from cerebral vasospasm with a CT scan, CT angiogram, transcranial Doppler (TCD) ultrasound, or conventional x-ray angiography. Hydrocephalus may clear spontaneously or require temporary ventricular drainage. Chronic hydrocephalus may develop weeks to months after SAH and manifest as gait difficulty, incontinence, or impaired mentation. Subtle signs may be a lack of initiative in conversation or a failure to recover independence.
3. **Vasospasm.** Narrowing of the arteries at the base of the brain following SAH causes symptomatic ischemia and infarction in $\sim 30\%$ of patients and is the major cause of delayed morbidity and death. Signs of ischemia appear 4–14 days after the hemorrhage, most often at 7 days. The severity and distribution of vasospasm determine whether infarction will occur.
4. Delayed vasospasm is believed to result from direct effects of clotted blood and its breakdown products on the arteries within the subarachnoid space. In general, the more blood that surrounds the arteries, the greater the chance of symptomatic vasospasm. Spasm of major arteries produces symptoms referable to the appropriate vascular territory (**Chap. 446**). All of these focal symptoms may present abruptly, fluctuate, or develop over a few days. In most cases, focal spasm is preceded by a decline in mental status.
5. Vasospasm can be detected reliably with conventional x-ray angiography, but this invasive procedure is expensive and carries the risk of stroke and other complications. TCD ultrasound is based on the principle that the velocity of blood flow within an artery will rise as the lumen diameter is narrowed. By directing the probe along the MCA and proximal anterior cerebral artery (ACA), carotid terminus, and vertebral and basilar arteries on a daily or every-other-day basis, vasospasm can be reliably detected and treatments initiated to prevent cerebral ischemia (see below). CT angiography is another method that can detect vasospasm.
6. Severe cerebral edema in patients with infarction from vasospasm may increase the ICP enough to reduce cerebral perfusion pressure. Treatment may include mannitol, hyperventilation, and hemicraniectomy; moderate hypothermia may have a role as well.
7. **Hyponatremia.** Hyponatremia may be profound and can develop quickly in the first 2 weeks following SAH. There is both natriuresis and volume depletion with SAH, so that patients become both hyponatremic and hypovolemic. Both atrial natriuretic peptide and brain natriuretic peptide have a role in producing this “cerebral

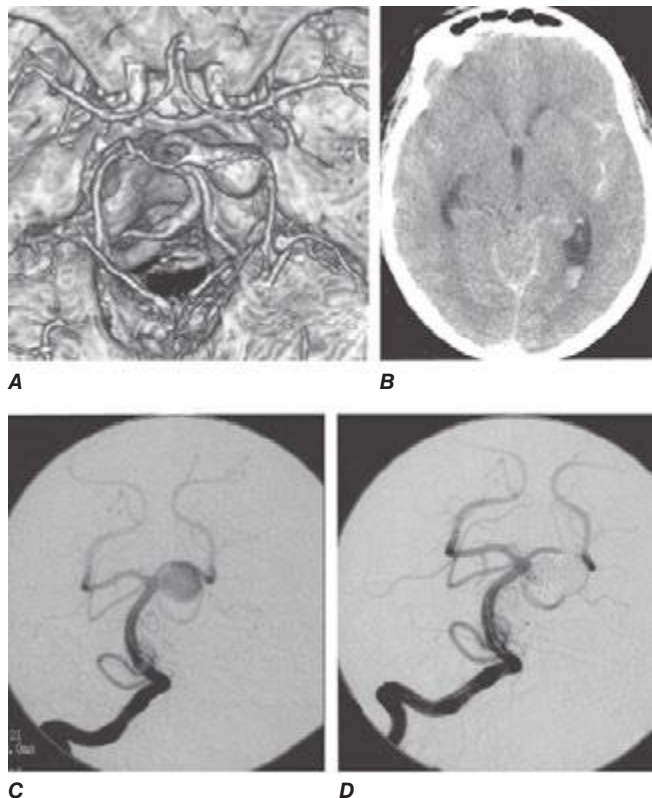


FIGURE 330-7 Subarachnoid hemorrhage. **A.** Computed tomography (CT) angiography revealing an aneurysm of the left superior cerebellar artery. **B.** Noncontrast CT scan at the level of the third ventricle revealing subarachnoid blood (*bright*) in the left sylvian fissure and within the left lateral ventricle. **C.** Conventional anteroposterior x-ray angiogram of the right vertebral and basilar artery showing the large aneurysm. **D.** Conventional angiogram following coil embolization of the aneurysm, whereby the aneurysm body is filled with platinum coils delivered through a microcatheter navigated from the femoral artery into the aneurysm neck.

salt-wasting syndrome.” Typically, it clears over the course of 1–2 weeks and, in the setting of SAH, should not be treated with free-water restriction as this may increase the risk of stroke (see below).

Laboratory Evaluation and Imaging (Fig. 330-7) The hallmark of aneurysmal rupture is blood in the CSF. More than 95% of cases have enough blood to be visualized on a high-quality noncontrast CT scan obtained within 72 h. If the scan fails to establish the diagnosis of SAH and no mass lesion or obstructive hydrocephalus is found, a lumbar puncture should be performed to establish the presence of subarachnoid blood. Lysis of the red blood cells and subsequent conversion of hemoglobin to bilirubin stains the spinal fluid yellow within 6–12 h. This xanthochromic spinal fluid peaks in intensity at 48 h and lasts for 1–4 weeks, depending on the amount of subarachnoid blood.

The extent and location of subarachnoid blood on a noncontrast CT scan help locate the underlying aneurysm, identify the cause of any neurologic deficit, and predict delayed vasospasm. A high incidence of symptomatic vasospasm in the MCA and ACA has been found when early CT scans show subarachnoid clots $>5 \times 3$ mm in the basal cisterns, or layers of blood >1 mm thick in the cerebral fissures. CT scans less reliably predict vasospasm in the vertebral, basilar, or posterior cerebral arteries.

Lumbar puncture prior to an imaging procedure is indicated only if a CT scan is not available at the time of the suspected SAH. Once the diagnosis of hemorrhage from a ruptured saccular aneurysm is suspected, four-vessel conventional x-ray angiography (both carotids and both vertebrals) is generally performed to localize and define the anatomic details of the aneurysm and to determine if other

unruptured aneurysms exist (Fig. 330-7C). At some centers, the ruptured aneurysm can be treated using endovascular techniques at the time of the initial angiogram as a way to expedite treatment and minimize the number of invasive procedures. CT angiography is an alternative method for locating the aneurysm and may be sufficient to plan definitive therapy.

Close monitoring (daily or twice daily) of electrolytes is important because hyponatremia can occur precipitously during the first 2 weeks following SAH (see above).

The electrocardiogram (ECG) frequently shows ST-segment and T-wave changes similar to those associated with cardiac ischemia. A prolonged QRS complex, increased QT interval, and prominent “peaked” or deeply inverted symmetric T waves are usually secondary to the intracranial hemorrhage. There is evidence that structural myocardial lesions produced by circulating catecholamines and excessive discharge of sympathetic neurons may occur after SAH, causing these ECG changes and a reversible cardiomyopathy sufficient to cause shock or congestive heart failure. Echocardiography reveals a pattern of regional wall motion abnormalities that follow the distribution of sympathetic nerves rather than the major coronary arteries, with relative sparing of the ventricular wall apex. The sympathetic nerves themselves appear to be injured by direct toxicity from the excessive catecholamine release. An asymptomatic troponin elevation is common. Serious ventricular dysrhythmias occurring in-hospital are unusual.

TREATMENT SUBARACHNOID HEMORRHAGE

Early aneurysm repair prevents rerupture and allows the safe application of techniques to improve blood flow (e.g., induced hypertension) should symptomatic vasospasm develop. An aneurysm can be “clipped” by a neurosurgeon or “coiled” by an endovascular surgeon. Surgical repair involves placing a metal clip across the aneurysm neck, thereby immediately eliminating the risk of rebleeding. This approach requires craniotomy and brain retraction, which is associated with neurologic morbidity. Endovascular techniques involve placing platinum coils, or other embolic material, within the aneurysm via a catheter that is passed from the femoral artery. The aneurysm is packed tightly to enhance thrombosis and over time is walled off from the circulation (Fig. 330-7D). There have been two prospective randomized trials of surgery versus endovascular treatment for ruptured aneurysms: the first was the International Subarachnoid Aneurysm Trial (ISAT), which was terminated early when 24% of patients treated with endovascular therapy were dead or dependent at 1 year compared to 31% treated with surgery, a significant 23% relative reduction. After 5 years, risk of death was lower in the coiling group, although the proportion of survivors who were independent was the same in both groups. Risk of rebleeding was low, but more common in the coiling group. These results favoring coiling at 1 year were confirmed in a second trial, although the differences in functional outcome were no longer significant at 3 years. Because some aneurysms have a morphology that is not amenable to endovascular treatment, surgery remains an important treatment option. Centers that combine both endovascular and neurosurgical expertise likely offer the best outcomes for patients, and there are reliable data showing that specialized aneurysm treatment centers can improve mortality rates.

The medical management of SAH focuses on protecting the airway, managing blood pressure before and after aneurysm treatment, preventing rebleeding prior to treatment, managing vasospasm, treating hydrocephalus, treating hyponatremia, limiting secondary brain insults, and preventing pulmonary embolus (PE).

Intracranial hypertension following aneurysmal rupture occurs secondary to subarachnoid blood, parenchymal hematoma, acute hydrocephalus, or loss of vascular autoregulation. Patients who are stuporous should undergo emergent ventriculostomy to measure ICP and to treat high ICP in order to prevent cerebral ischemia. Medical therapies designed to combat raised ICP (e.g., osmotic therapy and sedation) can also be used as needed. High ICP refractory to treatment is a poor prognostic sign.

Prior to definitive treatment of the ruptured aneurysm, care is required to maintain adequate cerebral perfusion pressure while avoiding excessive elevation of arterial pressure. If the patient is alert, it is reasonable to lower the systolic blood pressure to below 160 mmHg using nicardipine, labetalol, or esmolol. If the patient has a depressed level of consciousness, ICP should be measured and the cerebral perfusion pressure targeted to 60–70 mmHg. If headache or neck pain is severe, mild sedation and analgesia are prescribed. Extreme sedation is avoided if possible because it can obscure the ability to clinically detect changes in neurologic status. Adequate hydration is necessary to avoid a decrease in blood volume predisposing to brain ischemia.

Seizures are uncommon at the onset of aneurysmal rupture. The quivering, jerking, and extensor posturing that often accompany loss of consciousness with SAH are probably related to the sharp rise in ICP rather than seizures. However, anticonvulsants are sometimes given as prophylactic therapy because a seizure could theoretically promote rebleeding.

Glucocorticoids may help reduce the head and neck ache caused by the irritative effect of the subarachnoid blood. There is no good evidence that they reduce cerebral edema, are neuroprotective, or reduce vascular injury, and their routine use therefore is not recommended.

Antifibrinolytic agents are not routinely prescribed but may be considered in patients in whom aneurysm treatment cannot proceed immediately. They are associated with a reduced incidence of aneurysmal rerupture but may also increase the risk of delayed cerebral infarction and deep vein thrombosis (DVT). Several recent studies suggest that a shorter duration of use (until the aneurysm is secured or for the first 3 days) may decrease rerupture and be safer than found in earlier studies of longer duration treatment.

Vasospasm remains the leading cause of morbidity and mortality following aneurysmal SAH. Treatment with the calcium channel antagonist nimodipine (60 mg PO every 4 h) improves outcome, perhaps by preventing ischemic injury rather than reducing the risk of vasospasm. Nimodipine can cause significant hypotension in some patients, which may worsen cerebral ischemia in patients with vasospasm. Symptomatic cerebral vasospasm can also be treated by increasing the cerebral perfusion pressure by raising mean arterial pressure through plasma volume expansion and the judicious use of IV vasopressor agents, usually phenylephrine or norepinephrine. Raised perfusion pressure has been associated with clinical improvement in many patients, but high arterial pressure may promote rebleeding in unprotected aneurysms. Treatment with

induced hypertension and hypervolemia generally requires monitoring of arterial and central venous pressures; it is best to infuse pressors through a central venous line as well. Volume expansion helps prevent hypotension and augments cardiac output.

If symptomatic vasospasm persists despite optimal medical therapy, intraarterial vasodilators and percutaneous transluminal angioplasty are considered. Vasodilatation by direct angioplasty appears to be permanent, allowing hypertensive therapy to be tapered sooner. The pharmacologic vasodilators (verapamil and nicardipine) do not last more than about 24 h, and therefore multiple treatments may be required until the subarachnoid blood is reabsorbed. Although intraarterial papaverine is an effective vasodilator, there is evidence that papaverine may be neurotoxic, so its use should generally be avoided.

Acute hydrocephalus can cause stupor or coma. It may clear spontaneously or require temporary ventricular drainage. When chronic hydrocephalus develops, ventricular shunting is the treatment of choice.

Free-water restriction is contraindicated in patients with SAH at risk for vasospasm because hypovolemia and hypotension may occur and precipitate cerebral ischemia. Many patients continue to experience a decline in serum sodium despite receiving parenteral fluids containing normal saline. Frequently, supplemental oral salt coupled with normal saline will mitigate hyponatremia, but often patients also require intravenous hypertonic saline. Care must be taken not to correct serum sodium too quickly in patients with marked hyponatremia of several days' duration, as central pontine myelinolysis may occur.

All patients should have pneumatic compression stockings applied to prevent pulmonary embolism. Unfractionated heparin administered subcutaneously for DVT prophylaxis can be initiated immediately following endovascular treatment and within days following craniotomy with surgical clipping and is a useful adjunct to pneumatic compression stockings. Treatment of pulmonary embolus depends on whether the aneurysm has been treated and whether or not the patient has had a craniotomy. Systemic anticoagulation with heparin is contraindicated in patients with ruptured and untreated aneurysms. It is a relative contraindication following craniotomy for several days, and it may delay thrombosis of a coiled aneurysm. If DVT or PE occurs within the first days following craniotomy, use of an inferior vena cava filter may be considered to prevent additional pulmonary emboli, whereas systemic anticoagulation with heparin is preferred following successful endovascular treatment.

SECTION 4 ONCOLOGIC EMERGENCIES

331 Oncologic Emergencies

Rasim Gucalp, Janice P. Dutcher

Emergencies in patients with cancer may be classified into three groups: pressure or obstruction caused by a space-occupying lesion, metabolic or hormonal problems (paraneoplastic syndromes, [Chap. 121](#)), and treatment-related complications.

STRUCTURAL-OBSTRUCTIVE ONCOLOGIC EMERGENCIES

SUPERIOR VENA CAVA SYNDROME

Superior vena cava syndrome (SVCS) is the clinical manifestation of superior vena cava (SVC) obstruction, with severe reduction in venous return from the head, neck, and upper extremities. Malignant

tumors, such as lung cancer, lymphoma, and metastatic tumors, are responsible for the majority of SVCS cases. With the expanding use of intravascular devices (e.g., permanent central venous access catheters, pacemaker/defibrillator leads), the prevalence of benign causes of SVCS is increasing now, accounting for at least 40% of cases. Lung cancer, particularly of small-cell and squamous cell histologies, accounts for approximately 85% of all cases of malignant origin. In young adults, malignant lymphoma is a leading cause of SVCS. Hodgkin's lymphoma involves the mediastinum more commonly than other lymphomas but rarely causes SVCS. When SVCS is noted in a young man with a mediastinal mass, the differential diagnosis is lymphoma versus primary mediastinal germ cell tumor. Metastatic cancers to the mediastinal lymph nodes, such as testicular and breast carcinomas, account for a small proportion of cases. Other causes include benign tumors, aortic aneurysm, thyromegaly, thrombosis, and fibrosing mediastinitis from prior irradiation, histoplasmosis, or Behçet's syndrome. SVCS as

1788 the initial manifestation of Behçet's syndrome may be due to inflammation of the SVC associated with thrombosis.

Patients with SVCS usually present with neck and facial swelling (especially around the eyes), dyspnea, and cough. Other symptoms include hoarseness, tongue swelling, headaches, nasal congestion, epistaxis, hemoptysis, dysphagia, pain, dizziness, syncope, and lethargy. Bending forward or lying down may aggravate the symptoms. The characteristic physical findings are dilated neck veins; an increased number of collateral veins covering the anterior chest wall; cyanosis; and edema of the face, arms, and chest. Facial swelling and plethora are typically exacerbated when the patient is supine. More severe cases include proptosis, glossal and laryngeal edema, and obtundation. The clinical picture is milder if the obstruction is located above the azygos vein. Symptoms are usually progressive, but in some cases, they may improve as collateral circulation develops.

Signs and symptoms of cerebral and/or laryngeal edema, though rare, are associated with a poorer prognosis and require urgent evaluation. Seizures are more likely related to brain metastases than to cerebral edema from venous occlusion. Patients with small-cell lung cancer and SVCS have a higher incidence of brain metastases than those without SVCS.

Cardiorespiratory symptoms at rest, particularly with positional changes, suggest significant airway and vascular obstruction and limited physiologic reserve. Cardiac arrest or respiratory failure can occur, particularly in patients receiving sedatives or undergoing general anesthesia.

Rarely, esophageal varices may develop. These are "downhill" varices based on the direction of blood flow from cephalad to caudad (in contrast to "uphill" varices associated with caudad to cephalad flow from portal hypertension). If the obstruction to the SVC is proximal to the azygos vein, varices develop in the upper one-third of the esophagus. If the obstruction involves or is distal to the azygos vein, varices occur in the entire length of the esophagus. Variceal bleeding may be a late complication of chronic SVCS.

Superior vena cava obstruction may lead to bilateral breast edema with bilateral enlarged breast. Unilateral breast dilatation may be seen as a consequence of axillary or subclavian vein blockage.

The diagnosis of SVCS is a clinical one. The most significant chest radiographic finding is widening of the superior mediastinum, most commonly on the right side. Pleural effusion occurs in only 25% of patients, often on the right side. The majority of these effusions are exudative and occasionally chylous. However, a normal chest radiograph is still compatible with the diagnosis if other characteristic findings are present. Computed tomography (CT) provides the most reliable view of the mediastinal anatomy. The diagnosis of SVCS requires diminished or absent opacification of central venous structures with prominent collateral venous circulation. Magnetic resonance imaging (MRI) has no advantages over CT. Invasive procedures, including bronchoscopy, percutaneous needle biopsy, mediastinoscopy, and even thoracotomy, can be performed by a skilled clinician without any major risk of bleeding. Endobronchial or esophageal ultrasound-guided needle aspiration may establish the diagnosis safely. For patients with a known cancer, a detailed workup usually is not necessary, and appropriate treatment may be started after obtaining a CT scan of the thorax. For those with no history of malignancy, a detailed evaluation is essential to rule out benign causes and determine a specific diagnosis to direct the appropriate therapy.

TREATMENT SUPERIOR VENA CAVA SYNDROME

The one potentially life-threatening complication of a superior mediastinal mass is tracheal obstruction. Upper airway obstruction demands emergent therapy. Diuretics with a low-salt diet, head elevation, and oxygen may produce temporary symptomatic relief. Glucocorticoids may be useful at shrinking lymphoma masses; they are of no benefit in patients with lung cancer.

Radiation therapy is the primary treatment for SVCS caused by non-small-cell lung cancer and other metastatic solid tumors. Chemotherapy is effective when the underlying cancer is small-cell

carcinoma of the lung, lymphoma, or germ cell tumor. SVCS recurs in 10–30% of patients; it may be palliated with the use of intravascular self-expanding stents (**Fig. 331-1**). Early stenting may be necessary in patients with severe symptoms; however, the prompt increase in venous return after stenting may precipitate heart failure and pulmonary edema. Other complications of stent placement include hematoma at the insertion site, SVC perforation, stent migration in the right ventricle, stent fracture, and pulmonary embolism. Surgery may provide immediate relief for patients in whom a benign process is the cause.

Clinical improvement occurs in most patients, although this improvement may be due to the development of adequate collateral circulation. The mortality associated with SVCS does not relate to caval obstruction but rather to the underlying cause.

SVCS AND CENTRAL VENOUS CATHETERS IN ADULTS

The use of long-term central venous catheters has become common practice in patients with cancer. Major vessel thrombosis may occur. In these cases, catheter removal should be combined with anticoagulation to prevent embolization. SVCS in this setting, if detected early, can be treated by fibrinolytic therapy without sacrificing the catheter. The routine use of low-dose warfarin or low-molecular-weight heparin to prevent thrombosis related to permanent central venous access catheters in cancer patients is not recommended.

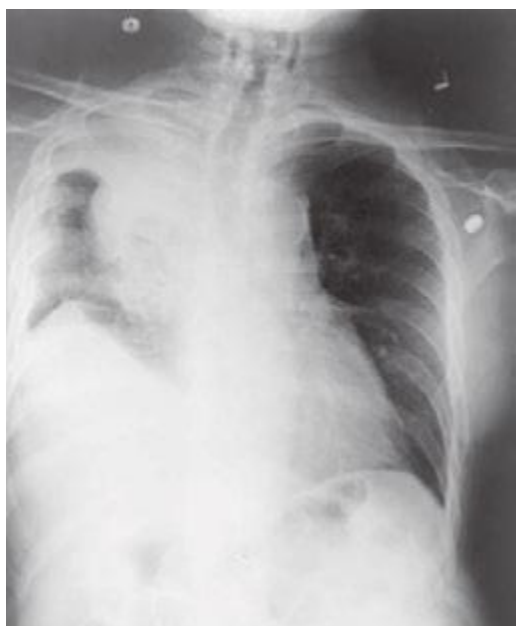
PERICARDIAL EFFUSION/TAMPONADE

Malignant pericardial disease is found at autopsy in 5–10% of patients with cancer, most frequently with lung cancer, breast cancer, leukemias, and lymphomas. Cardiac tamponade as the initial presentation of extrathoracic malignancy is rare. The origin is not malignancy in about 50% of cancer patients with symptomatic pericardial disease, but it can be related to irradiation, drug-induced pericarditis, hypothyroidism, idiopathic pericarditis, infection, or autoimmune diseases. Two types of radiation pericarditis occur: an acute inflammatory, effusive pericarditis occurring within months of irradiation, which usually resolves spontaneously, and a chronic effusive pericarditis that may appear up to 20 years after radiation therapy and is accompanied by a thickened pericardium.

Most patients with pericardial metastasis are asymptomatic. However, the common symptoms are dyspnea, cough, chest pain, orthopnea, and weakness. Pleural effusion, sinus tachycardia, jugular venous distention, hepatomegaly, peripheral edema, and cyanosis are the most frequent physical findings. Relatively specific diagnostic findings, such as paradoxical pulse, diminished heart sounds, pulsus alternans (pulse waves alternating between those of greater and lesser amplitude with successive beats), and friction rub are less common than with nonmalignant pericardial disease. Chest radiographs and electrocardiogram (ECG) reveal abnormalities in 90% of patients, but half of these abnormalities are nonspecific. Echocardiography is the most helpful diagnostic test. Pericardial fluid may be serous, serosanguineous, or hemorrhagic, and cytologic examination of pericardial fluid is diagnostic in most patients. Measurements of tumor markers in the pericardial fluid are not helpful in the diagnosis of malignant pericardial fluid. Pericardioscopy (not widely available) with targeted pericardial and epicardial biopsy may differentiate neoplastic and benign pericardial disease. A combination of cytology, pericardial and epicardial biopsy, and guided pericardioscopy gives the best diagnostic yield. CT scan findings of irregular pericardial thickening and mediastinal lymphadenopathy suggest this is a malignant pericardial effusion. Cancer patients with pericardial effusion containing malignant cells on cytology have a very poor survival, about 7 weeks.

TREATMENT PERICARDIAL EFFUSION/TAMPONADE

Pericardiocentesis with or without the introduction of sclerosing agents, the creation of a pericardial window, complete pericardial stripping, cardiac irradiation, or systemic chemotherapy are effective treatments. Acute pericardial tamponade with life-threatening hemodynamic instability requires immediate drainage of fluid. This



A



B



C

FIGURE 331-1 Superior vena cava syndrome (SVCS). **A.** Chest radiographs of a 59-year-old man with recurrent SVCS caused by non-small-cell lung cancer showing right paratracheal mass with right pleural effusion. **B.** Computed tomography of same patient demonstrating obstruction of the superior vena cava with thrombosis (*arrow*) by the lung cancer (*square*) and collaterals (*arrowheads*). **C.** Balloon angioplasty (*arrowhead*) with Wallstent (*arrow*) in same patient.

can be quickly achieved by pericardiocentesis. The recurrence rate after percutaneous catheter drainage is about 20%. Sclerotherapy (pericardial instillation of bleomycin, mitomycin C, or tetracycline) may decrease recurrences. Alternatively, subxiphoid pericardiotomy can be performed in 45 min under local anesthesia. Thoracoscopic pericardial fenestration can be employed for benign causes; however, 60% of malignant pericardial effusions recur after this procedure. In a subset of patients, drainage of the pericardial effusion is paradoxically followed by worsening hemodynamic instability. This so-called “postoperative low cardiac output syndrome” occurs in up to 10% of patients undergoing surgical drainage and carries poor short-term survival.

INTESTINAL OBSTRUCTION

Intestinal obstruction and reobstruction are common problems in patients with advanced cancer, particularly colorectal or ovarian carcinoma. However, other cancers, such as lung or breast cancer and melanoma, can metastasize within the abdomen, leading to intestinal obstruction. Metastatic disease from colorectal, ovarian, pancreatic, gastric, and occasionally breast cancer can lead to peritoneal carcinomatosis, with infiltration of the omentum and peritoneal surface, thus limiting bowel motility. Typically, obstruction occurs at multiple sites in peritoneal carcinomatosis. Melanoma has a predilection to involve the small bowel; this involvement may be isolated, and resection may result in prolonged survival. Intestinal pseudoobstruction is caused by infiltration of the mesentery or bowel muscle by tumor, involvement of the celiac plexus, or paraneoplastic neuropathy in patients with small-cell lung cancer. Paraneoplastic neuropathy is associated with IgG antibodies reactive to neurons of the myenteric and submucosal plexuses of the jejunum and stomach. Ovarian cancer can lead to authentic luminal obstruction or to pseudoobstruction that results when circumferential invasion of a bowel segment arrests the forward progression of peristaltic contractions.

The onset of obstruction is usually insidious. Pain is the most common symptom and is usually colicky in nature. Pain can also be due to abdominal distention, tumor masses, or hepatomegaly. Vomiting can be intermittent or continuous. Patients with complete obstruction usually have constipation. Physical examination may reveal abdominal distention with tympany, ascites, visible peristalsis, high-pitched bowel sounds, and tumor masses. Erect plain abdominal films may reveal multiple air-fluid levels and dilation of the small or large bowel. Acute cecal dilation to >12–14 cm is considered a surgical emergency because of the high likelihood of rupture. CT scan is useful in defining the extent of disease and the exact nature of the obstruction and differentiating benign from malignant causes of obstruction in patients who have undergone surgery for malignancy. Malignant obstruction is suggested by a mass at the site of obstruction or prior surgery, adenopathy, or an abrupt transition zone and irregular bowel thickening at the obstruction site. Benign obstruction is more likely when CT shows mesenteric vascular changes, a large volume of ascites, or a smooth transition zone and smooth bowel thickening at the obstruction site. In challenging patients with obstructive symptoms, particularly low-grade small-bowel obstruction (SBO), CT enteroclysis often can help establish the diagnosis by providing distention of small-bowel loops. In this technique, water-soluble contrast is infused through a nasenteric tube into the duodenum or proximal small bowel followed by CT images. The prognosis for the patient with cancer who develops intestinal obstruction is poor; median survival is 3–4 months. About 25–30% of patients are found to have intestinal obstruction due to causes other than cancer. Adhesions from previous operations are a common benign cause. Ileus induced by vinca alkaloids, narcotics, or other drugs is another reversible cause.

TREATMENT INTestinal OBSTRUCTION

The management of intestinal obstruction in patients with advanced malignancy depends on the extent of the underlying malignancy, options for further antineoplastic therapy, estimated life expectancy,

1790 the functional status of the major organs, and the extent of the obstruction. The initial management should include surgical evaluation. Operation is not always successful and may lead to further complications with a substantial mortality rate (10–20%). Laparoscopy can diagnose and treat malignant bowel obstruction in some cases. Self-expanding metal stents placed in the gastric outlet, duodenum, proximal jejunum, colon, or rectum may palliate obstructive symptoms at those sites without major surgery. Patients known to have advanced intraabdominal malignancy should receive a prolonged course of conservative management, including nasogastric decompression. Percutaneous endoscopic or surgical gastrostomy tube placement is an option for palliation of nausea and vomiting, the so-called “venting gastrostomy.” Treatment with antiemetics, antispasmodics, and analgesics may allow patients to remain outside the hospital. Octreotide may relieve obstructive symptoms through its inhibitory effect on gastrointestinal secretion. Glucocorticoids have anti-inflammatory effects and may help the resolution of bowel obstruction. They also have antiemetic effects.

URINARY OBSTRUCTION

Urinary obstruction may occur in patients with prostatic or gynecologic malignancies, particularly cervical carcinoma; metastatic disease from other primary sites such as carcinomas of the breast, stomach, lung, colon, and pancreas; or lymphomas. Radiation therapy to pelvic tumors may cause fibrosis and subsequent ureteral obstruction. Bladder outlet obstruction is usually due to prostate and cervical cancers and may lead to bilateral hydronephrosis and renal failure.

Flank pain is the most common symptom. Persistent urinary tract infection, persistent proteinuria, or hematuria in patients with cancer should raise suspicion of ureteral obstruction. Total anuria and/or anuria alternating with polyuria may occur. A slow, continuous rise in the serum creatinine level necessitates immediate evaluation. Renal ultrasound is the safest and cheapest way to identify hydronephrosis. The function of an obstructed kidney can be evaluated by a nuclear scan. CT scan can reveal the point of obstruction and identify a retroperitoneal mass or adenopathy.

TREATMENT URINARY OBSTRUCTION

Obstruction associated with flank pain, sepsis, or fistula formation is an indication for immediate palliative urinary diversion. Internal ureteral stents can be placed under local anesthesia. Percutaneous nephrostomy offers an alternative approach for drainage. The placement of a nephrostomy is associated with a significant rate of pyelonephritis. In the case of bladder outlet obstruction due to malignancy, a suprapubic cystostomy can be used for urinary drainage. An aggressive intervention with invasive approaches to improve the obstruction should be weighed against the likelihood of antitumor response, and the ability to reverse renal insufficiency should be evaluated.

MALIGNANT BILIARY OBSTRUCTION

This common clinical problem can be caused by a primary carcinoma arising in the pancreas, ampulla of Vater, bile duct, or liver or by metastatic disease to the periductal lymph nodes or liver parenchyma. The most common metastatic tumors causing biliary obstruction are gastric, colon, breast, and lung cancers. Jaundice, light-colored stools, dark urine, pruritus, and weight loss due to malabsorption are usual symptoms. Pain and secondary infection are uncommon in malignant biliary obstruction. Ultrasound, CT scan, or percutaneous transhepatic or endoscopic retrograde cholangiography will identify the site and nature of the biliary obstruction.

TREATMENT MALIGNANT BILIARY OBSTRUCTION

Palliative intervention is indicated only in patients with disabling pruritus resistant to medical treatment, severe malabsorption, or infection. Stenting under radiographic control, surgical bypass,

or radiation therapy with or without chemotherapy may alleviate the obstruction. The choice of therapy should be based on the site of obstruction (proximal vs distal), the type of tumor (sensitive to radiotherapy, chemotherapy, or neither), and the general condition of the patient. In the absence of pruritus, biliary obstruction may be a largely asymptomatic cause of death.

SPINAL CORD COMPRESSION

Malignant spinal cord compression (MSCC) is defined as compression of the spinal cord and/or cauda equina by an extradural tumor mass. The minimum radiologic evidence for cord compression is indentation of the theca at the level of clinical features. Spinal cord compression occurs in 5–10% of patients with cancer. Epidural tumor is the first manifestation of malignancy in about 10% of patients. The underlying cancer is usually identified during the initial evaluation; lung cancer is the most common cause of MSCC.

Metastatic tumor involves the vertebral column more often than any other part of the bony skeleton. Lung, breast, and prostate cancer are the most frequent offenders. Multiple myeloma also has a high incidence of spine involvement. Lymphomas, melanoma, renal cell cancer, and genitourinary cancers also cause cord compression. The thoracic spine is the most common site (70%), followed by the lumbosacral spine (20%) and the cervical spine (10%). Involvement of multiple sites is most frequent in patients with breast and prostate carcinoma. Cord injury develops when metastases to the vertebral body or pedicle enlarge and compress the underlying dura. Another cause of cord compression is direct extension of a paravertebral lesion through the intervertebral foramen. These cases usually involve a lymphoma, myeloma, or pediatric neoplasm. Parenchymal spinal cord metastasis due to hematogenous spread is rare. Intramedullary metastases can be seen in lung cancer, breast cancer, renal cancer, melanoma, and lymphoma and are frequently associated with brain metastases and leptomeningeal disease.

Expanding extradural tumors induce injury through several mechanisms. Obstruction of the epidural venous plexus leads to edema. Local production of inflammatory cytokines enhances blood flow and edema formation. Compression compromises blood flow, leading to ischemia. Production of vascular endothelial growth factor is associated with spinal cord hypoxia and has been implicated as a potential cause of damage after spinal cord injury.

The most common initial symptom in patients with spinal cord compression is localized back pain and tenderness due to involvement of vertebrae by tumor. Pain is usually present for days or months before other neurologic findings appear. It is exacerbated by movement and by coughing or sneezing. It can be differentiated from the pain of disk disease by the fact that it worsens when the patient is supine. Radicular pain is less common than localized back pain and usually develops later. Radicular pain in the cervical or lumbosacral areas may be unilateral or bilateral. Radicular pain from the thoracic roots is often bilateral and is described by patients as a feeling of tight, band-like constriction around the thorax and abdomen. Typical cervical radicular pain radiates down the arm; in the lumbar region, the radiation is down the legs. *Lhermitte’s sign*, a tingling or electric sensation down the back and upper and lower limbs upon flexing or extending the neck, may be an early sign of cord compression. Loss of bowel or bladder control may be the presenting symptom but usually occurs late in the course. Occasionally patients present with ataxia of gait without motor and sensory involvement due to involvement of the spinocerebellar tract.

On physical examination, pain induced by straight leg raising, neck flexion, or vertebral percussion may help to determine the level of cord compression. Patients develop numbness and paresthesias in the extremities or trunk. Loss of sensibility to pinprick is as common as loss of sensibility to vibration or position. The upper limit of the zone of sensory loss is often one or two vertebrae below the site of compression. Motor findings include weakness, spasticity, and abnormal muscle stretching. An extensor plantar reflex reflects significant compression. Deep tendon reflexes may be brisk. Motor and sensory loss usually precedes sphincter disturbance. Patients with autonomic dysfunction may present with decreased anal tonus, decreased perineal sensibility,

and a distended bladder. The absence of the anal wink reflex or the bulbocavernosus reflex confirms cord involvement. In doubtful cases, evaluation of postvoiding urinary residual volume can be helpful. A residual volume of >150 mL suggests bladder dysfunction. Autonomic dysfunction is an unfavorable prognostic factor. Patients with progressive neurologic symptoms should have frequent neurologic examinations and rapid therapeutic intervention. Other illnesses that may mimic cord compression include osteoporotic vertebral collapse, disk disease, pyogenic abscess or vertebral tuberculosis, radiation myelopathy, neoplastic leptomeningitis, benign tumors, epidural hematoma, and spinal lipomatosis.

Cauda equina syndrome is characterized by low back pain; diminished sensation over the buttocks, posterior-superior thighs, and perineal area in a saddle distribution; rectal and bladder dysfunction; sexual impotence; absent bulbocavernosus, patellar, and Achilles' reflexes; and variable amount of lower-extremity weakness. This reflects compression of nerve roots as they form the cauda equina after leaving the spinal cord. The majority of cauda equine tumors are primary tumors of glial or nerve sheath origin; metastases are very rare.

Patients with cancer who develop back pain should be evaluated for spinal cord compression as quickly as possible (Fig. 331-2). Treatment is more often successful in patients who are ambulatory and still have sphincter control at the time treatment is initiated. Patients should have a neurologic examination and plain films of the spine. Those whose physical examination suggests cord compression should receive dexamethasone (6 mg intravenously every 6 h), starting immediately.

Erosion of the pedicles (the "winking owl" sign) is the earliest radiologic finding of vertebral tumor. Other radiographic changes include increased intrapedicular distance, vertebral destruction, lytic or sclerotic lesions, scalloped vertebral bodies, and vertebral body collapse. Vertebral collapse is not a reliable indicator of the presence of tumor; about 20% of cases of vertebral collapse, particularly those in

older patients and postmenopausal women, are due not to cancer but to osteoporosis. Also, a normal appearance on plain films of the spine does not exclude the diagnosis of cancer. The role of bone scans in the detection of cord compression is not clear; this method is sensitive but less specific than spinal radiography.

The full-length image of the cord provided by MRI is the imaging procedure of choice. Multiple epidural metastases are noted in 25% of patients with cord compression, and their presence influences treatment plans. On T1-weighted images, good contrast is noted between the cord, cerebrospinal fluid, and extradural lesions. Owing to its sensitivity in demonstrating the replacement of bone marrow by tumor, MRI can show which parts of a vertebra are involved by tumor. MRI also visualizes intraspinal extradural masses compressing the cord. T2-weighted images are most useful for the demonstration of intramedullary pathology. Gadolinium-enhanced MRI can help to delineate intramedullary disease. MRI is as good as or better than myelography plus postmyelogram CT scan in detecting metastatic epidural disease with cord compression. Myelography should be reserved for patients who have poor MRIs or who cannot undergo MRI promptly. CT scan in conjunction with myelography enhances the detection of small areas of spinal destruction.

In patients with cord compression and an unknown primary tumor, a simple workup including chest radiography, mammography, measurement of prostate-specific antigen, and abdominal CT usually reveals the underlying malignancy.

TREATMENT SPINAL CORD COMPRESSION

The treatment of patients with spinal cord compression is aimed at relief of pain and restoration/preservation of neurologic function (Fig. 331-2). Management of MSCC requires a multidisciplinary approach.

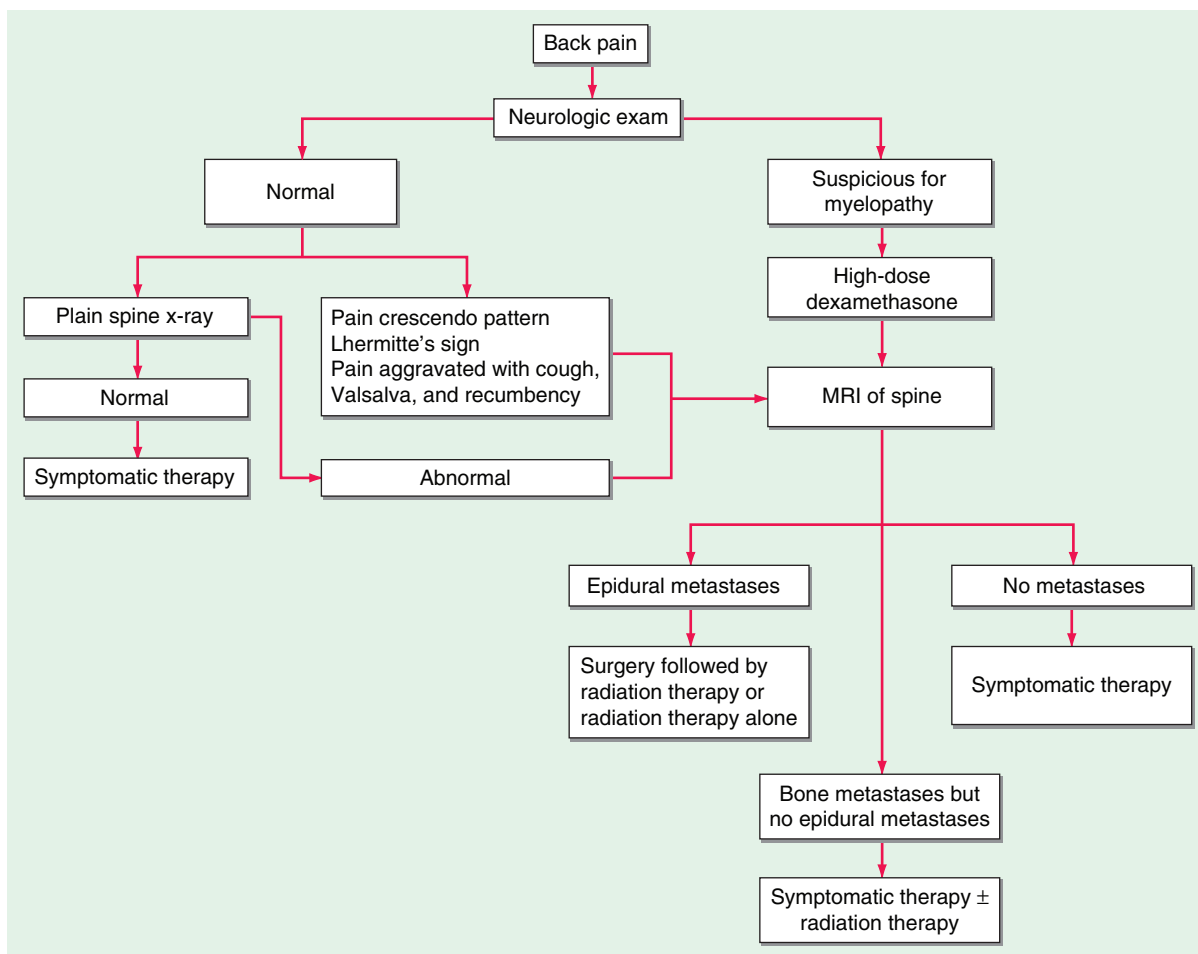


FIGURE 331-2 Management of cancer patients with back pain.

Radiation therapy plus glucocorticoids is generally the initial treatment of choice for most patients with spinal cord compression. Up to 75% of patients treated when still ambulatory remain ambulatory, but only 10% of patients with paraplegia recover walking capacity. Indications for surgical intervention include unknown etiology, failure of radiation therapy, a radioresistant tumor type (e.g., melanoma or renal cell cancer), pathologic fracture dislocation, and rapidly evolving neurologic symptoms. Laminectomy is done for tissue diagnosis and for the removal of posteriorly localized epidural deposits in the absence of vertebral body disease. Because most cases of epidural spinal cord compression are due to anterior or anterolateral extradural disease, resection of the anterior vertebral body along with the tumor, followed by spinal stabilization, has achieved good results. A randomized trial showed that patients who underwent an operation followed by radiotherapy (within 14 days) retained the ability to walk significantly longer than those treated with radiotherapy alone. Surgically treated patients also maintained continence and neurologic function significantly longer than patients in the radiation group. The length of survival was not significantly different in the two groups, although there was a trend toward longer survival in the surgery group. The study drew some criticism for the poorer than expected results in the patients who did not go to surgery. The benefit of surgery over radiotherapy decreased in patients over age 65 years. However, patients should be evaluated for surgery if they are expected to survive longer than 3 months. Conventional radiotherapy has a role after surgery. Chemotherapy may have a role in patients with chemosensitive tumors who have had prior radiotherapy to the same region and who are not candidates for surgery. Most patients with prostate cancer who develop cord compression have already had hormonal therapy; however, for those who have not, androgen deprivation is combined with surgery and radiotherapy. Patients who previously received radiotherapy for MSCC with an in-field tumor progression can be treated with reirradiation if they are not surgical candidates.

Patients with metastatic vertebral tumors may benefit from percutaneous vertebroplasty or kyphoplasty, the injection of acrylic cement into a collapsed vertebra to stabilize the fracture. Pain palliation is common, and local antitumor effects have been noted. Cement leakage may cause symptoms in about 10% of patients. Bisphosphonates may be helpful in prevention of SCC in patients with bony involvement.

The histology of the tumor is an important determinant of both recovery and survival. Rapid onset and progression of signs and symptoms are poor prognostic features.

INCREASED INTRACRANIAL PRESSURE

About 25% of patients with cancer die with intracranial metastases. The cancers that most often metastasize to the brain are lung and breast cancers and melanoma. Brain metastases often occur in the presence of systemic disease, and they frequently cause major symptoms, disability, and early death. The initial presentation of brain metastases from a previously unknown primary cancer is common. Lung cancer is most commonly the primary malignancy. Chest/abdomen CT scans and brain MRI as the initial diagnostic studies can identify a biopsy site in most patients.

The signs and symptoms of a metastatic brain tumor are similar to those of other intracranial expanding lesions: headache, nausea, vomiting, behavioral changes, seizures, and focal, progressive neurologic changes. Occasionally the onset is abrupt, resembling a stroke, with the sudden appearance of headache, nausea, vomiting, and neurologic deficits. This picture is usually due to hemorrhage into the metastasis. Melanoma, germ cell tumors, and renal cell cancers have a particularly high incidence of intracranial bleeding. The tumor mass and surrounding edema may cause obstruction of the circulation of cerebrospinal fluid, with resulting hydrocephalus. Patients with increased intracranial pressure may have papilledema with visual disturbances and neck stiffness. As the mass enlarges, brain tissue may be displaced

through the fixed cranial openings, producing various herniation syndromes.

CT scan and MRI are equally effective in the diagnosis of brain metastases. CT scan with contrast should be used as a screening procedure. The CT scan shows brain metastases as multiple enhancing lesions of various sizes with surrounding areas of low-density edema. If a single lesion or no metastases are visualized by contrast-enhanced CT, MRI of the brain should be performed. Gadolinium-enhanced MRI is more sensitive than CT at revealing meningeal involvement and small lesions, particularly in the brainstem or cerebellum.

Intracranial hypertension ("pseudotumor cerebri") secondary to tretinoin therapy has been reported.

TREATMENT INCREASED INTRACRANIAL PRESSURE

Dexamethasone is the best initial treatment for all symptomatic patients with brain metastases. Patients with multiple lesions should usually receive whole-brain radiation. Patients with a single brain metastasis and with controlled extracranial disease may be treated with surgical excision followed by whole-brain radiation therapy, especially if they are younger than 60 years. Radioresistant tumors should be resected if possible. Stereotactic radiosurgery (SRS) is recommended in patients with a limited number of brain metastases (one to four) who have stable, systemic disease or reasonable systemic treatment options and for patients who have a small number of metastatic lesions in whom whole-brain radiation therapy has failed. With a gamma knife or linear accelerator, multiple small, well-collimated beams of ionizing radiation destroy lesions seen on MRI. Some patients with increased intracranial pressure associated with hydrocephalus may benefit from shunt placement. If neurologic deterioration is not reversed with medical therapy, ventriculotomy to remove cerebrospinal fluid (CSF) or craniotomy to remove tumors or hematomas may be necessary.

NEOPLASTIC MENINGITIS

Tumor involving the leptomeninges is a complication of both primary central nervous system (CNS) tumors and tumors that metastasize to the CNS. The incidence is estimated at 3–8% of patients with cancer. Melanoma, breast and lung cancer, lymphoma (including AIDS-associated), and acute leukemia are the most common causes. Synchronous intraparenchymal brain metastases are evident in 11–31% of patients with neoplastic meningitis. Leptomeningeal seeding is frequent in patients undergoing resection of brain metastases or receiving stereotactic radiotherapy for brain metastases.

Patients typically present with multifocal neurologic signs and symptoms, including headache, gait abnormality, mental changes, nausea, vomiting, seizures, back or radicular pain, and limb weakness. Signs include cranial nerve palsies, extremity weakness, paresthesia, and decreased deep tendon reflexes.

Diagnosis is made by demonstrating malignant cells in the CSF; however, up to 40% of patients may have false-negative CSF cytology. An elevated CSF protein level is nearly always present (except in HTLV-1-associated adult T cell leukemia). Patients with neurologic signs and symptoms consistent with neoplastic meningitis who have a negative CSF cytology but an elevated CSF protein level should have the spinal tap repeated at least three times for cytologic examination before the diagnosis is rejected. MRI findings suggestive of neoplastic meningitis include leptomeningeal, subependymal, dural, or cranial nerve enhancement; superficial cerebral lesions; intradural nodules; and communicating hydrocephalus. Spinal cord imaging by MRI is a necessary component of the evaluation of nonleukemia neoplastic meningitis because ~20% of patients have cord abnormalities, including intradural enhancing nodules that are diagnostic for leptomeningeal involvement. Cauda equina lesions are common, but lesions may be seen anywhere in the spinal canal. The value of MRI for the diagnosis of leptomeningeal disease is limited in patients with hematopoietic malignancy. Radiolabeled CSF flow studies are abnormal in up to 70% of patients with neoplastic meningitis; ventricular outlet

obstruction, abnormal flow in the spinal canal, or impaired flow over the cerebral convexities may affect distribution of intrathecal chemotherapy, resulting in decreased efficacy or increased toxicity. Radiation therapy may correct CSF flow abnormalities before use of intrathecal chemotherapy. Neoplastic meningitis can also lead to intracranial hypertension and hydrocephalus. Placement of a ventriculoperitoneal shunt may effectively palliate symptoms in these patients.

The development of neoplastic meningitis usually occurs in the setting of uncontrolled cancer outside the CNS; thus, prognosis is poor (median survival 10–12 weeks). However, treatment of the neoplastic meningitis may successfully alleviate symptoms and control the CNS spread.

TREATMENT NEOPLASTIC MENINGITIS

Intrathecal chemotherapy, usually methotrexate, cytarabine, or thiotepa, is delivered by lumbar puncture or by an intraventricular reservoir (Ommaya). An extended-release preparation of cytarabine (Depocyte) has a longer half-life and is more effective than other formulations. Among solid tumors, breast cancer responds best to therapy. Epidermal growth factor receptor (EGFR) tyrosine kinase inhibitors (TKIs) may be effective in non-small-cell lung cancer patients with *EGFR* mutations and leptomeningeal involvement. Patients with neoplastic meningitis from either acute leukemia or lymphoma may be cured of their CNS disease if the systemic disease can be eliminated.

SEIZURES

Seizures occurring in a patient with cancer can be caused by the tumor itself, by metabolic disturbances, by radiation injury, by cerebral infarctions, by chemotherapy-related encephalopathies, or by CNS infections. Metastatic disease to the CNS is the most common cause of seizures in patients with cancer. However, seizures occur more frequently in primary brain tumors than in metastatic brain lesions. Seizures are a presenting symptom of CNS metastasis in 6–29% of cases. Approximately 10% of patients with CNS metastasis eventually develop seizures. Tumors that affect the frontal, temporal, and parietal lobes are more commonly associated with seizures than are occipital lesions. The presence of frontal lesions correlates with early seizures, and the presence of hemispheric symptoms increases the risk for late seizures. Both early and late seizures are uncommon in patients with posterior fossa and sellar lesions. Seizures are common in patients with CNS metastases from melanoma and low-grade primary brain tumors. Very rarely, cytotoxic drugs such as etoposide, busulfan, ifosfamide, and chlorambucil cause seizures. Another cause of seizures related to drug therapy is reversible posterior leukoencephalopathy syndrome (RPLS). RPLS is associated rarely with administration of cisplatin, 5-fluorouracil, bleomycin, vinblastine, vincristine, etoposide, paclitaxel, ifosfamide, cyclophosphamide, doxorubicin, cytarabine, methotrexate, oxaliplatin, cyclosporine, tacrolimus, and vascular endothelial growth factor inhibitors including bevacizumab, aflibercept, sunitinib, sorafenib, pazopanib, and axitinib. RPLS occurs in patients undergoing allogeneic bone marrow or solid-organ transplantation. RPLS is characterized by headache, altered consciousness, generalized seizures, visual disturbances, hypertension, and posterior cerebral white matter vasogenic edema on CT/MRI. Seizures may begin focally but are typically generalized.

TREATMENT SEIZURES

Patients in whom seizures due to CNS metastases have been demonstrated should receive anticonvulsive treatment with phenytoin or levetiracetam. If this is not effective, valproic acid can be added. Prophylactic anticonvulsant therapy is not recommended. In postcraniotomy patients, prophylactic antiepileptic drugs should be withdrawn during the first week after surgery. Most antiseizure medications including phenytoin induce cytochrome P450 (CYP450), which alters the metabolism of many antitumor agents,

including irinotecan, taxanes, and etoposide as well as molecular targeted agents, including imatinib, gefitinib, erlotinib, tipifarnib, sorafenib, sunitinib, temsirolimus, everolimus, and vemurafenib. Levetiracetam and topiramate are anticonvulsant agents not metabolized by the hepatic CYP450 system and do not alter the metabolism of antitumor agents. They have become the preferred drugs. Surgical resection and other antitumor treatments such as radiotherapy and chemotherapy may improve seizure control.

PULMONARY AND INTRACEREBRAL LEUKOSTASIS

Hyperleukocytosis and the leukostasis syndrome associated with it is a potentially fatal complication of acute leukemia (particularly myeloid leukemia) that can occur when the peripheral blast cell count is $>100,000/\text{mL}$. The frequency of hyperleukocytosis is 5–13% in acute myeloid leukemia (AML) and 10–30% in acute lymphoid leukemia; however, leukostasis is rare in lymphoid leukemia. At such high blast cell counts, blood viscosity is increased, blood flow is slowed by aggregates of tumor cells, and the primitive myeloid leukemic cells are capable of invading through the endothelium and causing hemorrhage. Brain and lung are most commonly affected. Patients with brain leukostasis may experience stupor, headache, dizziness, tinnitus, visual disturbances, ataxia, confusion, coma, or sudden death. On examination, papilledema, retinal vein distension, retinal hemorrhages, and focal deficit may be present. Administration of 600 cGy of whole-brain irradiation can protect against this complication and can be followed by rapid institution of antileukemic therapy. Hydroxyurea, 3–5 g, can rapidly reduce a high blast cell count while the accurate diagnostic workup is in progress. Pulmonary leukostasis may present as respiratory distress and hypoxemia, and progress to respiratory failure. Chest radiographs may be normal but usually show interstitial or alveolar infiltrates. Hyperleukocytosis rarely may cause acute leg ischemia, renal vein thrombosis, myocardial ischemia, bowel infarction, and priapism. Arterial blood gas results should be interpreted cautiously. Rapid consumption of plasma oxygen by the markedly increased number of white blood cells can cause spuriously low arterial oxygen tension. Pulse oximetry is the most accurate way of assessing oxygenation in patients with hyperleukocytosis. Leukapheresis may be helpful in decreasing circulating blast counts. Treatment of the leukemia can result in pulmonary hemorrhage from lysis of blasts in the lung, called *leukemic cell lysis pneumopathy*. Intravascular volume depletion and unnecessary blood transfusions may increase blood viscosity and worsen the leukostasis syndrome. Leukostasis is very rarely a feature of the high white cell counts associated with chronic lymphoid or chronic myeloid leukemia.

When acute promyelocytic leukemia is treated with differentiating agents like tretinoin and arsenic trioxide, cerebral or pulmonary leukostasis may occur as tumor cells differentiate into mature neutrophils. This complication can be largely avoided by using cytotoxic chemotherapy or arsenic together with the differentiating agents.

HEMOPTYSIS

Hemoptysis may be caused by nonmalignant conditions, but lung cancer accounts for a large proportion of cases. Up to 20% of patients with lung cancer have hemoptysis some time in their course. Endobronchial metastases from carcinoid tumors, breast cancer, colon cancer, kidney cancer, and melanoma may also cause hemoptysis. The volume of bleeding is often difficult to gauge. Massive hemoptysis is defined as $>200\text{--}600\text{ mL}$ of blood produced in 24 h. However, any hemoptysis should be considered massive if it threatens life. When respiratory difficulty occurs, hemoptysis should be treated emergently. The first priorities are to maintain the airway, optimize oxygenation, and stabilize the hemodynamic status. If the bleeding side is known, the patient should be placed in a lateral decubitus position, with the bleeding side down to prevent aspiration into the unaffected lung, and given supplemental oxygen. If large-volume bleeding continues or the airway is compromised, the patient should be intubated and undergo emergency bronchoscopy. If the site of bleeding is detected, either the patient undergoes a definitive surgical procedure or the lesion is treated with a neodymium:yttrium-aluminum-garnet (Nd:YAG) laser, argon plasma coagulation, or electrocautery. In stable

1794 patients, multidetector CT angiography delineates bronchial and non-bronchial systemic arteries and identifies the source of bleeding and underlying pathology with high sensitivity. Massive hemoptysis usually originates from the high-pressure bronchial circulation. Bronchial artery embolization is considered a first-line definite procedure for managing hemoptysis. Bronchial artery embolization may control brisk bleeding in 75–90% of patients, permitting the definitive surgical procedure to be done more safely.

Embolization without definitive surgery is associated with rebleeding in 20–50% of patients. Recurrent hemoptysis usually responds to a second embolization procedure. A postembolization syndrome characterized by pleuritic pain, fever, dysphagia, and leukocytosis may occur; it lasts 5–7 days and resolves with symptomatic treatment. Bronchial or esophageal wall necrosis, myocardial infarction, and spinal cord infarction are rare complications. Surgery, as a salvage strategy, is indicated after failure of embolization and is associated with better survival when performed in a nonurgent setting.

Pulmonary hemorrhage with or without hemoptysis in hematologic malignancies is often associated with fungal infections, particularly *Aspergillus* sp. After granulocytopenia resolves, the lung infiltrates in aspergillosis may cavitate and cause massive hemoptysis. Thrombocytopenia and coagulation defects should be corrected, if possible. Surgical evaluation is recommended in patients with aspergillosis-related cavitory lesions.

Bevacizumab, an antibody to vascular endothelial growth factor (VEGF) that inhibits angiogenesis, has been associated with life-threatening hemoptysis in patients with non-small-cell lung cancer, particularly of squamous cell histology. Non-small-cell lung cancer patients with cavitory lesions or previous hemoptysis (≥ 2.5 mL) within the past 3 months have higher risk for pulmonary hemorrhage.

AIRWAY OBSTRUCTION

Airway obstruction refers to a blockage at the level of the mainstem bronchi or above. It may result either from intraluminal tumor growth or from extrinsic compression of the airway. The most common cause of malignant upper airway obstruction is invasion from an adjacent primary tumor, most commonly lung cancer, followed by esophageal, thyroid, and mediastinal malignancies including lymphomas. Extrathoracic primary tumors such as renal, colon, or breast cancer can cause airway obstruction through endobronchial and/or mediastinal lymph node metastases. Patients may present with dyspnea, hemoptysis, stridor, wheezing, intractable cough, postobstructive pneumonia, or hoarseness. Chest radiographs usually demonstrate obstructing lesions. CT scans reveal the extent of tumor. Cool, humidified oxygen, glucocorticoids, and ventilation with a mixture of helium and oxygen (Heliox) may provide temporary relief. If the obstruction is proximal to the larynx, a tracheostomy may be lifesaving. For more distal obstructions, particularly intrinsic lesions incompletely obstructing the airway, bronchoscopy with mechanical debulking and dilatation or ablational treatments including laser treatment, photodynamic therapy, argon plasma coagulation, electrocautery, or stenting can produce immediate relief in most patients (Fig. 331-3). However, radiation therapy (either external-beam irradiation or brachytherapy) given together with glucocorticoids may also open the airway. Symptomatic extrinsic compression may be palliated by stenting. Patients with primary airway tumors such as squamous cell carcinoma, carcinoid tumor, adenocystic carcinoma, or non-small-cell lung cancer, if resectable, should have surgery.

METABOLIC EMERGENCIES

HYPERCALCEMIA

Hypercalcemia is the most common paraneoplastic syndrome. **Its pathogenesis and management are discussed fully in Chaps. 121 and 424.**

SYNDROME OF INAPPROPRIATE SECRETION OF ANTIDIURETIC HORMONE (SIADH)

Hyponatremia is a common electrolyte abnormality in cancer patients, and SIADH is the most common cause among patients with cancer. **SIADH is discussed fully in Chaps. 121 and 401e.**



A



B

FIGURE 331-3 Airway obstruction. A. Computed tomography scan of a 62-year-old man with tracheal obstruction caused by renal carcinoma showing paratracheal mass with tracheal invasion/obstruction (arrow). B. Chest x-ray of same patient after stent (arrows) placement.

LACTIC ACIDOSIS

Lactic acidosis is a rare and potentially fatal metabolic complication of cancer. Lactic acidosis associated with sepsis and circulatory failure is a common preterminal event in many malignancies. Lactic acidosis in the absence of hypoxemia may occur in patients with leukemia, lymphoma, or solid tumors. In some cases, hypoglycemia also is present. Extensive involvement of the liver by tumor is often present. In most cases, decreased metabolism and increased production by the tumor both contribute to lactate accumulation. Tumor cell overexpression of certain glycolytic enzymes and mitochondrial dysfunction can contribute to its increased lactate production. HIV-infected patients have an increased risk of aggressive lymphoma; lactic acidosis that occurs in such patients may be related either to the rapid growth of the tumor or from toxicity of nucleoside reverse transcriptase inhibitors. Symptoms of lactic acidosis include tachypnea, tachycardia, change of mental status, and hepatomegaly. The serum level of lactic acid may reach 10–20 mmol/L (90–180 mg/dL). Treatment is aimed at the underlying disease. *The danger from lactic acidosis is from the acidosis, not the lactate.* Sodium bicarbonate should be added if acidosis is very severe or if hydrogen ion production is very rapid and uncontrolled. Other treatment options include renal replacement therapy, such as hemodialysis, and thiamine replacement. The prognosis is poor regardless of the treatment offered.

HYPOGLYCEMIA

Persistent hypoglycemia is occasionally associated with tumors other than pancreatic islet cell tumors. Usually these tumors are large; tumors of mesenchymal origin, hepatomas, or adrenocortical tumors

may cause hypoglycemia. Mesenchymal tumors are usually located in the retroperitoneum or thorax. Obtundation, confusion, and behavioral aberrations occur in the postabsorptive period and may precede the diagnosis of the tumor. These tumors often secrete incompletely processed insulin-like growth factor II (IGF-II), a hormone capable of activating insulin receptors and causing hypoglycemia. Tumors secreting incompletely processed big IGF-II are characterized by an increased IGF-II to IGF-I ratio, suppressed insulin and C-peptide level, and inappropriately low growth hormone and β -hydroxybutyrate concentrations. Rarely, hypoglycemia is due to insulin secretion by a non-islet cell carcinoma. The development of hepatic dysfunction from liver metastases and increased glucose consumption by the tumor can contribute to hypoglycemia. If the tumor cannot be resected, hypoglycemia symptoms may be relieved by the administration of glucose, glucocorticoids, or glucagon.

Hypoglycemia can be artifactual; hyperleukocytosis from leukemia, myeloproliferative diseases, leukemoid reactions, or colony-stimulating factor treatment can increase glucose consumption in the test tube after blood is drawn, leading to pseudohypoglycemia.

ADRENAL INSUFFICIENCY

In patients with cancer, adrenal insufficiency may go unrecognized because the symptoms, such as nausea, vomiting, anorexia, and orthostatic hypotension, are nonspecific and may be mistakenly attributed to progressive cancer or to therapy. Primary adrenal insufficiency may develop owing to replacement of both glands by metastases (lung, breast, colon, or kidney cancer; lymphoma), to removal of both glands, or to hemorrhagic necrosis in association with sepsis or anticoagulation. Impaired adrenal steroid synthesis occurs in patients being treated for cancer with mitotane, ketoconazole, or aminoglutethimide or undergoing rapid reduction in glucocorticoid therapy. Rarely, metastatic replacement causes primary adrenal insufficiency as the first manifestation of an occult malignancy. Metastasis to the pituitary or hypothalamus is found at autopsy in up to 5% of patients with cancer, but associated secondary adrenal insufficiency is rare. On the other hand, ipilimumab, an anti-CTLA-4 antibody used for treatment of malignant melanoma, may cause autoimmunity including autoimmune-like enterocolitis, hypophysitis, and hepatitis. Autoimmune hypophysitis may present with headache, visual field defects, and pituitary hormone deficiencies manifesting as hypopituitarism, adrenal insufficiency (including adrenal crisis), or hypothyroidism. Anti-CTLA-4-associated hypophysitis symptoms occur at an average of 6–12 weeks after initiation of therapy. The treatment of severe autoimmune toxicity is glucocorticoids. Almost all patients with hypophysitis respond to withdrawal of ipilimumab and glucocorticoid therapy in several days. However, pituitary dysfunction may resolve or may be permanent, requiring long-term therapy and thyroid and testosterone replacement. Peripheral Addison's disease can also be observed with anti-CTLA-4 antibodies. Megestrol acetate, used to manage cancer and HIV-related cachexia, may suppress plasma levels of cortisol and adrenocorticotropic hormone (ACTH). Patients taking megestrol may develop adrenal insufficiency, and even those whose adrenal dysfunction is not symptomatic may have inadequate adrenal reserve if they become seriously ill. Paradoxically, some patients may develop Cushing's syndrome and/or hyperglycemia because of the glucocorticoid-like activity of megestrol acetate. Cranial irradiation for childhood brain tumors may affect the hypothalamus-pituitary-adrenal axis, resulting in secondary adrenal insufficiency.

Acute adrenal insufficiency is potentially lethal. Treatment of suspected adrenal crisis is initiated after the sampling of serum cortisol and ACTH levels (Chap. 406).

TREATMENT-RELATED EMERGENCIES

TUMOR LYSIS SYNDROME

Tumor lysis syndrome (TLS) is characterized by hyperuricemia, hyperkalemia, hyperphosphatemia, and hypocalcemia and is caused by the destruction of a large number of rapidly proliferating neoplastic cells. Acidosis may also develop. Acute renal failure occurs frequently.

TLS is most often associated with the treatment of Burkitt's lymphoma, acute lymphoblastic leukemia, and other rapidly proliferating lymphomas, but it also may be seen with chronic leukemias and, rarely, with solid tumors. This syndrome has been seen in patients with chronic lymphocytic leukemia after treatment with nucleosides like fludarabine. TLS has been observed with administration of glucocorticoids, hormonal agents such as letrozole and tamoxifen, and monoclonal antibodies such as rituximab and gemtuzumab. TLS usually occurs during or shortly (1–5 days) after chemotherapy. Rarely, spontaneous necrosis of malignancies causes TLS.

Hyperuricemia may be present at the time of chemotherapy. Effective treatment kills malignant cells and leads to increased serum uric acid levels from the turnover of nucleic acids. Owing to the acidic local environment, uric acid can precipitate in the tubules, medulla, and collecting ducts of the kidney, leading to renal failure. Lactic acidosis and dehydration may contribute to the precipitation of uric acid in the renal tubules. The finding of uric acid crystals in the urine is strong evidence for uric acid nephropathy. The ratio of urinary uric acid to urinary creatinine is >1 in patients with acute hyperuricemic nephropathy and <1 in patients with renal failure due to other causes.

Hyperphosphatemia, which can be caused by the release of intracellular phosphate pools by tumor lysis, produces a reciprocal depression in serum calcium, which causes severe neuromuscular irritability and tetany. Deposition of calcium phosphate in the kidney and hyperphosphatemia may cause renal failure. Potassium is the principal intracellular cation, and massive destruction of malignant cells may lead to hyperkalemia. Hyperkalemia in patients with renal failure may rapidly become life-threatening by causing ventricular arrhythmias and sudden death.

The likelihood that TLS will occur in patients with Burkitt's lymphoma is related to the tumor burden and renal function. Hyperuricemia and high serum levels of lactate dehydrogenase (LDH >1500 U/L), both of which correlate with total tumor burden, also correlate with the risk of TLS. In patients at risk for TLS, pretreatment evaluations should include a complete blood count, serum chemistry evaluation, and urine analysis. High leukocyte and platelet counts may artificially elevate potassium levels ("pseudohyperkalemia") due to lysis of these cells after the blood is drawn. In these cases, plasma potassium instead of serum potassium should be followed. In pseudohyperkalemia, no electrocardiographic abnormalities are present. In patients with abnormal baseline renal function, the kidneys and retroperitoneal area should be evaluated by sonography and/or CT to rule out obstructive uropathy. Urine output should be watched closely.

TREATMENT TUMOR LYSIS SYNDROME

Recognition of risk and prevention are the most important steps in the management of this syndrome (Fig. 331-4). The standard preventive approach consists of allopurinol, urinary alkalinization, and aggressive hydration. Urinary alkalization with sodium bicarbonate is controversial. It increases uric acid solubility, but decreases calcium phosphate solubility. If it is used, it should be discontinued when hyperphosphatemia develops. Intravenous allopurinol may be given in patients who cannot tolerate oral therapy. In some cases, uric acid levels cannot be lowered sufficiently with the standard preventive approach. Rasburicase (recombinant urate oxidase) can be effective in these instances, particularly when renal failure is present. Urate oxidase is missing from primates and catalyzes the conversion of poorly soluble uric acid to readily soluble allantoin. Rasburicase acts rapidly, decreasing uric acid levels within hours; however, it may cause hypersensitivity reactions such as bronchospasm, hypoxemia, and hypotension. Rasburicase should also be administered to high-risk patients for TLS prophylaxis. Rasburicase is contraindicated in patients with glucose-6-phosphate dehydrogenase deficiency who are unable to break down hydrogen peroxide, an end product of the urate oxidase reaction. Rasburicase is known to cause *ex vivo* enzymatic degradation of uric acid in test tube at room temperature. This leads to spuriously low uric acid levels

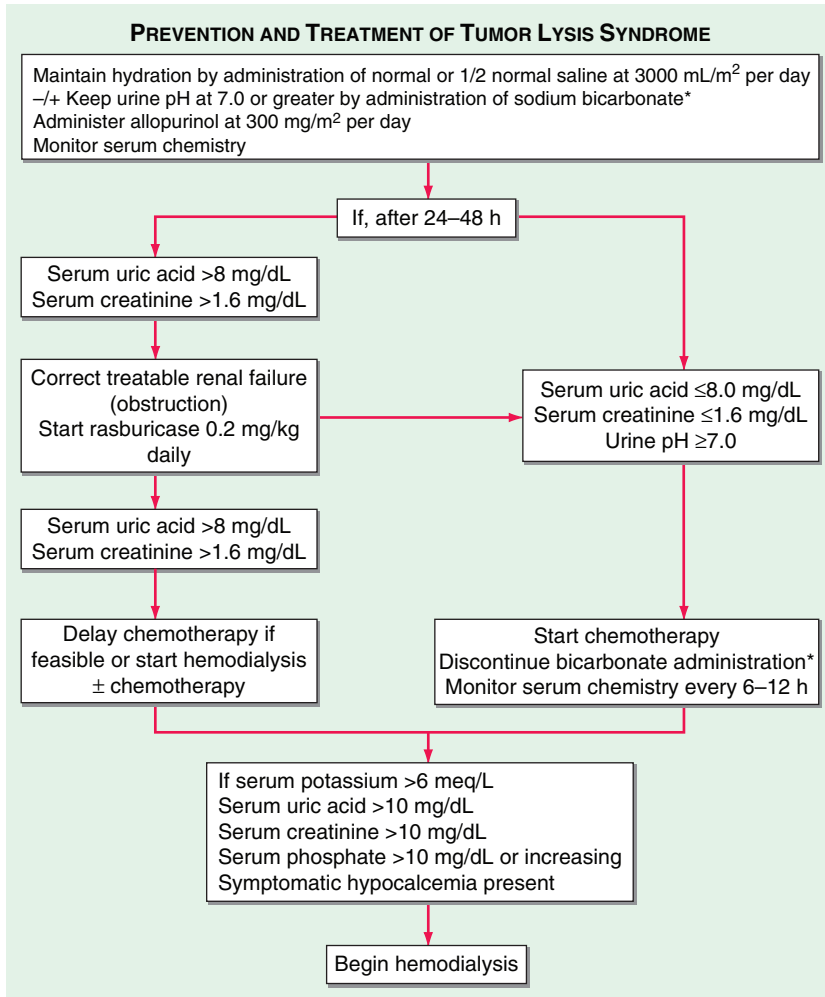


FIGURE 331-4 Management of patients at high risk for the tumor lysis syndrome.

*See text.

during laboratory monitoring of the patient with TLS. Samples must be cooled immediately to deactivate the urate oxidase. Despite aggressive prophylaxis, TLS and/or oliguric or anuric renal failure may occur. Care should be taken to prevent worsening of symptomatic hypocalcemia by induction of alkalosis during bicarbonate infusion. Administration of sodium bicarbonate may also lead to urinary precipitation of calcium phosphate, which is less soluble at alkaline pH. Dialysis is often necessary and should be considered early in the course. Hemodialysis is preferred. Hemofiltration offers a gradual, continuous method of removing cellular by-products and fluid. The prognosis is excellent, and renal function recovers after the uric acid level is lowered to ≤ 10 mg/dL.

HUMAN ANTIBODY INFUSION REACTIONS

The initial infusion of human or humanized antibodies (e.g., rituximab, gemtuzumab, trastuzumab, alemtuzumab, panitumumab, brentuximab vedotin) is associated with fever, chills, nausea, asthenia, and headache in up to half of treated patients. Bronchospasm and hypotension occur in 1% of patients. Severe manifestations including pulmonary infiltrates, acute respiratory distress syndrome, and cardiogenic shock occur rarely. Laboratory manifestations include elevated hepatic aminotransferase levels, thrombocytopenia, and prolongation of prothrombin time. The pathogenesis is thought to be activation of immune effector processes (cells and complement) and release of inflammatory cytokines, such as tumor necrosis factor α , interferon gamma, interleukin 6, and interleukin 10 (cytokine release syndrome [CRS]). Although its origins are not completely understood, CRS is believed to be due to activation of a variety of cell types including

monocytes/macrophages and T and B lymphocytes. Severe reactions from rituximab have occurred with high numbers ($>50 \times 10^9$ lymphocytes) of circulating cells bearing the target antigen (CD20) and have been associated with a rapid fall in circulating tumor cells, mild electrolyte evidence of TLS, and very rarely, death. In addition, increased liver enzymes, D-dimer, and LDH and prolongation of the prothrombin time may occur. Diphenhydramine, hydrocortisone, and acetaminophen can often prevent or suppress the infusion-related symptoms. If they occur, the infusion is stopped and restarted at half the initial infusion rate after the symptoms have abated. Severe CRS may require intensive support for acute respiratory distress syndrome (ARDS) and resistant hypotension.

HEMOLYTIC-UREMIC SYNDROME

Hemolytic-uremic syndrome (HUS) and, less commonly, thrombotic thrombocytopenic purpura (TTP) (Chap. 341) may rarely occur after treatment with antineoplastic drugs, including mitomycin, gemcitabine, cisplatin, and bleomycin, and with VEGF inhibitors. It occurs most often in patients with gastric, lung, colorectal, pancreatic, and breast carcinoma. In one series, 35% of patients were without evident cancer at the time this syndrome appeared. Secondary HUS/TTP has also been reported as a rare but sometimes fatal complication of bone marrow transplantation.

HUS usually has its onset 4–8 weeks after the last dose of chemotherapy, but it is not rare to detect it several months later. HUS is characterized by microangiopathic hemolytic anemia, thrombocytopenia, and renal failure. Dyspnea, weakness, fatigue, oliguria, and purpura are also common initial symptoms and findings. Systemic hypertension and pulmonary edema frequently occur. Severe hypertension, pulmonary edema, and rapid worsening of hemolysis and renal function may occur after a blood or blood product transfusion. Cardiac findings include atrial arrhythmias, pericardial friction rub, and pericardial effusion. Raynaud's phenomenon is part of the syndrome in patients treated with bleomycin.

Laboratory findings include severe to moderate anemia associated with red blood cell fragmentation and numerous schistocytes on peripheral smear. Reticulocytosis, decreased plasma haptoglobin, and an LDH level document hemolysis. The serum bilirubin level is usually normal or slightly elevated. The Coombs' test is negative. The white cell count is usually normal, and thrombocytopenia ($<100,000/\mu\text{L}$) is almost always present. Most patients have a normal coagulation profile, although some have mild elevations in thrombin time and in levels of fibrin degradation products. The serum creatinine level is elevated at presentation and shows a pattern of subacute worsening within weeks of the initial azotemia. The urinalysis reveals hematuria, proteinuria, and granular or hyaline casts; and circulating immune complexes may be present.

The basic pathologic lesion appears to be deposition of fibrin in the walls of capillaries and arterioles, and these deposits are similar to those seen in HUS due to other causes. These microvascular abnormalities involve mainly the kidneys and rarely occur in other organs. The pathogenesis of cancer treatment-related HUS is not completely understood, but probably the most important factor is endothelial damage. Primary forms of HUS/TTP are related to a decrease in processing of von Willebrand factor by a protease called ADAMTS13.

The case fatality rate is high; most patients die within a few months. There is no consensus on the optimal treatment for chemotherapy-induced HUS. Treatment modalities for HUS/TTP including immunocomplex removal (plasmapheresis, immunoadsorption, or exchange transfusion), antiplatelet/anticoagulant therapies, immunosuppressive

therapies, and plasma exchange have varying degrees of success. The outcome with plasma exchange is generally poor, as in many other cases of secondary TTP. Rituximab is successfully used in patients with chemotherapy-induced HUS as well as in ADAMTS13-deficient TTP.

NEUTROPENIA AND INFECTION

These remain the most common serious complications of cancer therapy. **They are covered in detail in Chap. 104.**

PULMONARY INFILTRATES

Patients with cancer may present with dyspnea associated with diffuse interstitial infiltrates on chest radiographs. Such infiltrates may be due to progression of the underlying malignancy, treatment-related toxicities, infection, and/or unrelated diseases. The cause may be multifactorial; however, most commonly they occur as a consequence of treatment. Infiltration of the lung by malignancy has been described in patients with leukemia, lymphoma, and breast and other solid cancers. Pulmonary lymphatics may be involved diffusely by neoplasm (pulmonary lymphangitic carcinomatosis), resulting in a diffuse increase in interstitial markings on chest radiographs. The patient is often mildly dyspneic at the onset, but pulmonary failure develops over a period of weeks. In some patients, dyspnea precedes changes on the chest radiographs and is accompanied by a nonproductive cough. This syndrome is characteristic of solid tumors. In patients with leukemia, diffuse microscopic neoplastic peribronchial and peribronchiolar infiltration is frequent but may be asymptomatic. However, some patients present with diffuse interstitial infiltrates, an alveolar capillary block syndrome, and respiratory distress. In these situations, glucocorticoids can provide symptomatic relief, but specific chemotherapy should always be started promptly.

Several cytotoxic agents, such as bleomycin, methotrexate, busulfan, nitrosoureas, gemcitabine, mitomycin, vinorelbine, docetaxel, paclitaxel, fludarabine, pentostatin, and ifosfamide may cause pulmonary damage. The most frequent presentations are interstitial pneumonitis, alveolitis, and pulmonary fibrosis. Some cytotoxic agents, including methotrexate and procarbazine, may cause an acute hypersensitivity reaction. Cytosine arabinoside has been associated with noncardiogenic pulmonary edema. Administration of multiple cytotoxic drugs, as well as radiotherapy and preexisting lung disease, may potentiate the pulmonary toxicity. Supplemental oxygen may potentiate the effects of drugs and radiation injury. Patients should always be managed with the lowest FIO₂ that is sufficient to maintain hemoglobin saturation.

The onset of symptoms may be insidious, with symptoms including dyspnea, nonproductive cough, and tachycardia. Patients may have bibasilar crepitant rales, end-inspiratory crackles, fever, and cyanosis. The chest radiograph generally shows an interstitial and sometimes an intraalveolar pattern that is strongest at the lung bases and may be symmetric. A small effusion may occur. Hypoxemia with decreased carbon monoxide diffusing capacity is always present. Glucocorticoids may be helpful in patients in whom pulmonary toxicity is related to radiation therapy or to chemotherapy. Treatment is otherwise supportive.

Molecular targeted agents, imatinib, erlotinib, and gefitinib are potent inhibitors of tyrosine kinases. These drugs may cause interstitial lung disease (ILD). In the case of gefitinib, preexisting fibrosis, poor performance status, and prior thoracic irradiation are independent risk factors; this complication has a high fatality rate. In Japan, incidence of interstitial lung disease associated with gefitinib was about 4.5% compared to 0.5% in the United States. Temsirolimus and everolimus, both esters a derivative of rapamycin, are agents that block the effects of mammalian target of rapamycin (mTOR), an enzyme that has an important role in regulating the synthesis of proteins that control cell division. It may cause ground-glass opacities in the lung with or without diffuse interstitial disease and lung parenchymal consolidation. Patients may be asymptomatic with only radiologic findings or may be symptomatic. Symptoms include cough, dyspnea, and/or hypoxemia, and sometimes patients present with systemic symptoms such as fever and fatigue. The incidence of everolimus-induced

interstitial lung disease also appears to be higher in Japanese patients. Treatment includes dose reduction or withdrawal and, in some cases, the addition of glucocorticoids.

Radiation pneumonitis and/or fibrosis is a relatively frequent side effect of thoracic radiation therapy. It may be acute or chronic. Radiation-induced lung toxicity is a function of the irradiated lung volume, dose per fraction, and radiation dose. The larger the irradiated lung field, the higher is the risk for radiation pneumonitis. The use of concurrent chemoradiation, particularly regimens including paclitaxel, increases pulmonary toxicity. Radiation pneumonitis usually develops 2–6 months after completion of radiotherapy. The clinical syndrome, which varies in severity, consists of dyspnea, cough with scanty sputum, low-grade fever, and an initial hazy infiltrate on chest radiographs. The infiltrate and tissue damage usually are confined to the radiation field. The patients subsequently may develop a patchy alveolar infiltrate and air bronchograms, which may progress to acute respiratory failure that is sometimes fatal. A lung biopsy may be necessary to make the diagnosis. Asymptomatic infiltrates found incidentally after radiation therapy need not be treated. However, prednisone should be administered to patients with fever or other symptoms. The dosage should be tapered slowly after the resolution of radiation pneumonitis, because abrupt withdrawal of glucocorticoids may cause an exacerbation of pneumonia. Delayed radiation fibrosis may occur years after radiation therapy and is signaled by dyspnea on exertion. Often it is mild, but it can progress to chronic respiratory failure. Therapy is supportive.

Classical radiation pneumonitis that leads to pulmonary fibrosis is due to radiation-induced production of local cytokines such as platelet-derived growth factor β , tumor necrosis factor, interleukins, and transforming growth factor β in the radiation field. An immunologically mediated sporadic radiation pneumonitis occurs in about 10% of patients; bilateral alveolitis mediated by T cells results in infiltrates outside the radiation field. This form of radiation pneumonitis usually resolves without sequelae.

Pneumonia is a common problem in patients undergoing treatment for cancer. Bacterial pneumonia typically causes a localized infiltrate on chest radiographs. Therapy is tailored to the causative organism. When diffuse interstitial infiltrates appear in a febrile patient, the differential diagnosis is extensive and includes pneumonia due to infection with *Pneumocystis carinii*; viral infections including cytomegalovirus, adenovirus, herpes simplex virus, herpes zoster, respiratory syncytial virus, or intracellular pathogens such as *Mycoplasma* and *Legionella*; effects of drugs or radiation; tumor progression; nonspecific pneumonitis; and fungal disease. Detection of opportunistic pathogens in pulmonary infections is still a challenge. Diagnostic tools include chest radiographs, CT scans, bronchoscopy with bronchoalveolar lavage, brush cytology, transbronchial biopsy, fine-needle aspiration, and open lung biopsy. In addition to the culture, evaluation of bronchoalveolar lavage fluid for *P. carinii* by polymerase chain reaction (PCR) and serum galactomannan test improve the diagnostic yield. Patients with cancer who are neutropenic and have fever and local infiltrates on chest radiograph should be treated initially with broad-spectrum antibiotics. A new or persistent focal infiltrate not responding to broad-spectrum antibiotics argues for initiation of empiric antifungal therapy. When diffuse bilateral infiltrates develop in patients with febrile neutropenia, broad-spectrum antibiotics plus trimethoprim-sulfamethoxazole, with or without erythromycin, should be initiated. Addition of an antiviral agent is necessary in some settings, such as patients undergoing allogeneic hematopoietic stem cell transplantation. If the patient does not improve in 4 days, open lung biopsy is the procedure of choice. Bronchoscopy with bronchoalveolar lavage may be used in patients who are poor candidates for surgery.

In patients with pulmonary infiltrates who are afebrile, heart failure and multiple pulmonary emboli are in the differential diagnosis.

NEUTROPENIC ENTEROCOLITIS

Neutropenic enterocolitis (typhlitis) is the inflammation and necrosis of the cecum and surrounding tissues that may complicate the treatment



FIGURE 331-5 Abdominal computed tomography (CT) scans of a 72-year-old woman with neutropenic enterocolitis secondary to chemotherapy. **A.** Air in inferior mesenteric vein (arrow) and bowel wall with pneumatosis intestinalis. **B.** CT scan of upper abdomen demonstrating air in portal vein (arrows).

of acute leukemia. Nevertheless, it may involve any segment of the gastrointestinal tract including small intestine, appendix, and colon. This complication has also been seen in patients with other forms of cancer treated with taxanes, 5-fluorouracil, irinotecan, vinorelbine, cisplatin, carboplatin, and high-dose chemotherapy (Fig. 331-5). It also has been reported in patients with AIDS, aplastic anemia, cyclic neutropenia, idiosyncratic drug reactions involving antibiotics, and immunosuppressive therapies. The patient develops right lower quadrant abdominal pain, often with rebound tenderness and a tense, distended abdomen, in a setting of fever and neutropenia. Watery diarrhea (often containing sloughed mucosa) and bacteremia are common, and bleeding may occur. Plain abdominal films are generally of little value in the diagnosis; CT scan may show marked bowel wall thickening, particularly in the cecum, with bowel wall edema, mesenteric stranding, and ascites, and may help to differentiate neutropenic colitis from other abdominal disorders such as appendicitis, diverticulitis, and *Clostridium difficile*-associated colitis in this high-risk population. Patients with bowel wall thickness >10 mm on ultrasonogram have higher mortality rates. However, bowel wall thickening is significantly more prominent in patients with *C. difficile* colitis. Pneumatosis intestinalis is a more specific finding, seen only in those with neutropenic enterocolitis and ischemia. The combined involvement of the small and large bowel suggests a diagnosis of neutropenic enterocolitis.

Rapid institution of broad-spectrum antibiotics, bowel rest, and nasogastric suction may reverse the process. Use of myeloid growth factors improved outcome significantly. Surgical intervention is reserved for severe cases of neutropenic enterocolitis with evidence of perforation, peritonitis, gangrenous bowel, or gastrointestinal hemorrhage despite correction of any coagulopathy.

C. difficile colitis is increasing in incidence. Newer strains of *C. difficile* produce about 20 times more of toxins A and B compared to previously studied strains. *C. difficile* risk is also increased with chemotherapy. Antibiotic coverage for *C. difficile* should be added if pseudomembranous colitis cannot be excluded.

HEMORRHAGIC CYSTITIS

Hemorrhagic cystitis can develop in patients receiving cyclophosphamide or ifosfamide. Both drugs are metabolized to acrolein, which is a strong chemical irritant that is excreted in the urine. Prolonged contact or high concentrations may lead to bladder irritation and hemorrhage. Symptoms include gross hematuria, frequency, dysuria, burning, urgency, incontinence, and nocturia. The best management is prevention. Maintaining a high rate of urine flow minimizes exposure. In addition, 2-mercaptoethanesulfonate (mesna) detoxifies the metabolites and can be coadministered with the instigating drugs. Mesna usually is given three times on the day of ifosfamide administration in doses that are each 20% of the total ifosfamide dose. If hemorrhagic cystitis develops, the maintenance of a high urine flow may be sufficient supportive care. If conservative management is not effective, irrigation of the bladder with a 0.37–0.74% formalin solution for 10 min stops the bleeding in most cases. N-Acetylcysteine may also be an effective irrigant. Prostaglandin (carboprost) can inhibit the process. In extreme cases, ligation of the hypogastric arteries, urinary diversion, or cystectomy may be necessary.

Hemorrhagic cystitis also occurs in patients who undergo bone marrow transplantation (BMT). In the BMT setting, early-onset hemorrhagic cystitis is related to drugs in the treatment regimen (e.g., cyclophosphamide), and late-onset hemorrhagic cystitis is usually due to the polyoma virus BKV or adenovirus type 11. BKV load in urine alone or in combination with acute graft-versus-host disease correlates with development of hemorrhagic cystitis. Viral causes are usually detected by PCR-based diagnostic tests. Treatment of viral hemorrhagic cystitis is largely supportive, with reduction in doses of immunosuppressive agents, if possible. No antiviral therapy is approved, although cidofovir is reported to be effective in a small series. Hyperbaric oxygen therapy has been used successfully in patients with BKV-associated and cyclophosphamide-induced hemorrhagic cystitis during hematopoietic stem cell transplantation, as well as in hemorrhagic radiation cystitis.

HYPERSENSITIVITY REACTIONS TO ANTINEOPLASTIC DRUGS

Many antineoplastic drugs may cause hypersensitivity reaction. These reactions are unpredictable and potentially life-threatening. Most reactions occur during or within hours of parenteral drug administration. Taxanes, platinum compounds, asparaginase, etoposide, procarbazine, and biologic agents, including rituximab, bevacizumab, trastuzumab, gemtuzumab, cetuximab, and alemtuzumab, are more commonly associated with acute hypersensitivity reactions than are other agents. Acute hypersensitivity reactions to some drugs, such as taxanes, occur during the first or second dose administered. Hypersensitivity to platinum compounds occurs after prolonged exposure. Skin testing may identify patients with high risk for hypersensitivity after carboplatin exposure. Premedication with histamine H₁ and H₂ receptor antagonists and glucocorticoids reduces the incidence of hypersensitivity reaction to taxanes, particularly paclitaxel. Despite premedication, hypersensitivity reactions may still occur. In these cases, rapid desensitization in the intensive care unit setting or re-treatment may be attempted with care, but the use of alternative agents may be required. Candidate patients for desensitization include those who have mild to severe hypersensitivity type I, with mast cell-mediated and IgE-dependent reactions occurring during a chemotherapy infusion or shortly thereafter.

332e Cellular and Molecular Biology of the Kidney

Alfred L. George, Jr., Eric G. Neilson

The kidney is one of the most highly differentiated organs in the body. At the conclusion of embryologic development, nearly 30 different cell types form a multitude of filtering capillaries and segmented nephrons enveloped by a dynamic interstitium. This cellular diversity modulates a variety of complex physiologic processes. Endocrine functions, the regulation of blood pressure and intraglomerular hemodynamics, solute and water transport, acid-base balance, and removal of drug metabolites are all accomplished by intricate mechanisms of renal response. This breadth of physiology hinges on the clever ingenuity of nephron architecture that evolved as complex organisms came out of water to live on land.

EMBRYOLOGIC DEVELOPMENT

Kidneys develop from intermediate mesoderm under the timed or sequential control of a growing number of genes, described in [Fig. 332e-1](#). The transcription of these genes is guided by morphogenic cues that invite two ureteric buds to each penetrate bilateral metanephric blastema, where they induce primary mesenchymal cells to form early nephrons. The two ureteric buds emerge from posterior nephric ducts and mature into separate collecting systems that eventually form a renal pelvis and ureter. Induced mesenchyme undergoes mesenchymal epithelial transitions to form comma-shaped bodies at the proximal end of each ureteric bud leading to the formation of S-shaped nephrons that cleft and enjoin with penetrating endothelial cells derived from sprouting angioblasts. Under the influence of vascular endothelial growth factor A (VEGF-A), these penetrating cells form capillaries with surrounding mesangial cells that differentiate into a glomerular filter for plasma water and solute. The ureteric buds branch, and each branch produce a new set of nephrons. The number of branching events ultimately determines the total number of nephrons in each kidney. There are approximately 900,000 glomeruli in each kidney in normal birth weight adults and as few as 225,000 in low-birth-weight adults, with the latter producing numerous comorbid risks.

Glomeruli evolve as complex capillary filters with fenestrated endothelia under the guiding influence of VEGF-A and angiotensin-1

secreted by adjacently developing podocytes. Epithelial podocytes facing the urinary space envelop the exterior basement membrane supporting these emerging endothelial capillaries. Podocytes are partially polarized and periodically fall off into the urinary space by epithelial-mesenchymal transition, and to a lesser extent apoptosis, only to be replenished by migrating parietal epithelia from Bowman capsule. Impaired replenishment results in heavy proteinuria. Podocytes attach to the basement membrane by special foot processes and share a slit-pore membrane with their neighbor. The slit-pore membrane forms a filter for plasma water and solute by the synthetic interaction of nephrin, annexin-4, CD2AP, FAT, ZO-1, P-cadherin, podocin, TRPC6, PLCE1, and Neph 1-3 proteins. Mutations in many of these proteins also result in heavy proteinuria. The glomerular capillaries are embedded in a mesangial matrix shrouded by parietal and proximal tubular epithelia forming Bowman capsule. Mesangial cells have an embryonic lineage consistent with arteriolar or juxtaglomerular cells and contain contractile actin-myosin fibers. These mesangial cells make contact with glomerular capillary loops, and their local matrix holds them in condensed arrangement.

Between nephrons lies the renal interstitium. This region forms a functional space surrounding glomeruli and their downstream tubules, which are home to resident and trafficking cells such as fibroblasts, dendritic cells, occasional lymphocytes, and lipid-laden macrophages. The cortical and medullary capillaries, which siphon off solute and water following tubular reclamation of glomerular filtrate, are also part of the interstitial fabric as well as a web of connective tissue that supports the kidney's emblematic architecture of folding tubules. The relational precision of these structures determines the unique physiology of the kidney.

Each nephron is partitioned during embryologic development into a proximal tubule, descending and ascending limbs of the loop of Henle, distal tubule, and the collecting duct. These classic tubular segments build from subsegments lined by highly unique epithelia serving regional physiology. All nephrons have the same structural components, but there are two types whose structures depend on their location within the kidney. The majority of nephrons are cortical, with glomeruli located in the mid-to-outer cortex. Fewer nephrons are juxtamedullary, with glomeruli at the boundary of the cortex and outer medulla. Cortical nephrons have short loops of Henle, whereas juxtamedullary nephrons have long loops of Henle. There are critical differences in blood supply as well. The peritubular capillaries surrounding cortical nephrons are shared among adjacent nephrons. By contrast, juxtamedullary nephrons depend on individual capillaries

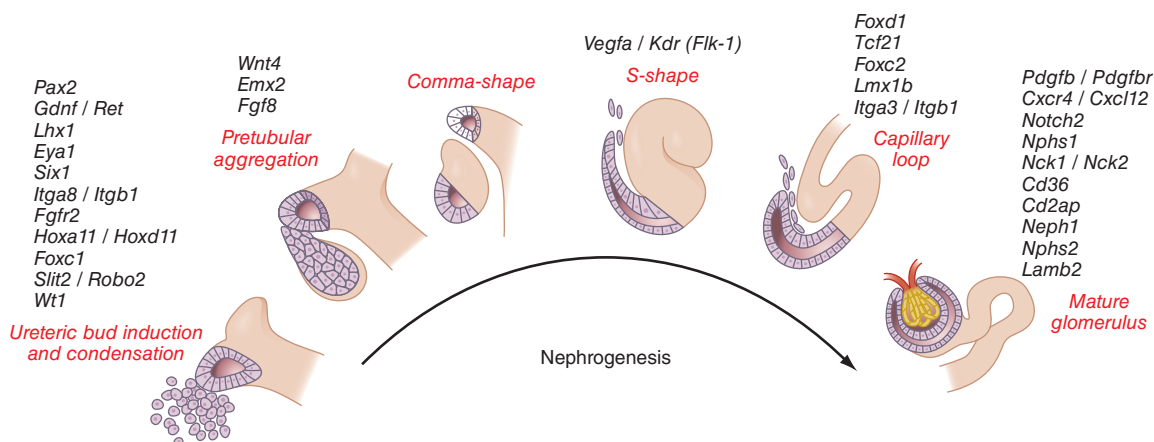


FIGURE 332e-1 Genes controlling renal nephrogenesis. A growing number of genes have been identified at various stages of glomerulotubular development in the mammalian kidney. The genes listed have been tested in various genetically modified mice, and their location corresponds to the classical stages of kidney development postulated by Saxen in 1987.

called *vasa recta*. Cortical nephrons perform most of the glomerular filtration because there are more of them and because their afferent arterioles are larger than their respective efferent arterioles. The juxtamedullary nephrons, with longer loops of Henle, create an osmotic gradient for concentrating urine. How developmental instructions specify the differentiation of all these unique epithelia among various tubular segments is still unknown.

DETERMINANTS AND REGULATION OF GLOMERULAR FILTRATION

Renal blood flow normally drains approximately 20% of the cardiac output, or 1000 mL/min. Blood reaches each nephron through the afferent arteriole leading into a glomerular capillary where large amounts of fluid and solutes are filtered to form the tubular fluid. The distal ends of the glomerular capillaries coalesce to form an efferent arteriole leading to the first segment of a second capillary network (cortical peritubular capillaries or medullary vasa recta) surrounding the tubules (Fig. 332e-2A). Thus, nephrons have two capillary beds arranged in a series separated by the efferent arteriole that regulates the hydrostatic pressure in both capillary beds. The distal capillaries empty into small venous branches that coalesce into larger veins to eventually form the renal vein.

The hydrostatic pressure gradient across the glomerular capillary wall is the primary driving force for glomerular filtration. Oncotic pressure within the capillary lumen, determined by the concentration of unfiltered plasma proteins, partially offsets the hydrostatic pressure gradient and opposes filtration. As the oncotic pressure rises along the length of the glomerular capillary, the driving force for filtration falls to zero on reaching the efferent arteriole. Approximately 20% of the renal plasma flow is filtered into Bowman space, and the ratio of glomerular filtration rate (GFR) to renal plasma flow determines the filtration fraction. Several factors, mostly hemodynamic, contribute to the regulation of filtration under physiologic conditions.

Although glomerular filtration is affected by renal artery pressure, this relationship is not linear across the range of physiologic blood pressures due to autoregulation of GFR. Autoregulation of glomerular filtration is the result of three major factors that modulate either afferent or efferent arteriolar tone: these include an autonomous vaso-reactive (myogenic) reflex in the afferent arteriole, *tubuloglomerular feedback*, and angiotensin II-mediated vasoconstriction of the efferent arteriole. The myogenic reflex is a first line of defense against fluctuations in renal blood flow. Acute changes in renal perfusion pressure evoke reflex constriction or dilatation of the afferent arteriole in response to increased or decreased pressure, respectively. This phenomenon helps protect the glomerular capillary from sudden changes in systolic pressure.

Tubuloglomerular feedback (TGF) changes the rate of filtration and tubular flow by reflex vasoconstriction or dilatation of the afferent arteriole. TGF is mediated by specialized cells in the thick ascending limb of the loop of Henle called the *macula densa* that act as sensors of solute concentration and tubular flow rate. With high tubular flow rates, a proxy for an inappropriately high filtration rate, there is increased solute delivery to the macula densa (Fig. 332e-2B) that evokes vasoconstriction of the afferent arteriole causing GFR to return toward normal. One component of the soluble signal from the macula densa is adenosine triphosphate (ATP) released by the cells during increased NaCl reabsorption. ATP is metabolized in the extracellular space to generate adenosine, a potent vasoconstrictor of the afferent arteriole. During conditions associated with a fall in filtration rate, reduced solute delivery to the macula densa attenuates TGF, allowing afferent arteriolar dilatation and restoring glomerular filtration to normal levels. Angiotensin II and reactive oxygen species enhance, while nitric oxide (NO) blunts, TGF.

The third component underlying autoregulation of GFR involves angiotensin II. During states of reduced renal blood flow, renin is released from granular cells within the wall of the afferent arteriole near the macula densa in a region called the juxtaglomerular apparatus (Fig. 332e-2B). Renin, a proteolytic enzyme, catalyzes the conversion of angiotensinogen to angiotensin I, which is subsequently converted to angiotensin II by angiotensin-converting enzyme (ACE) (Fig. 332e-2C).

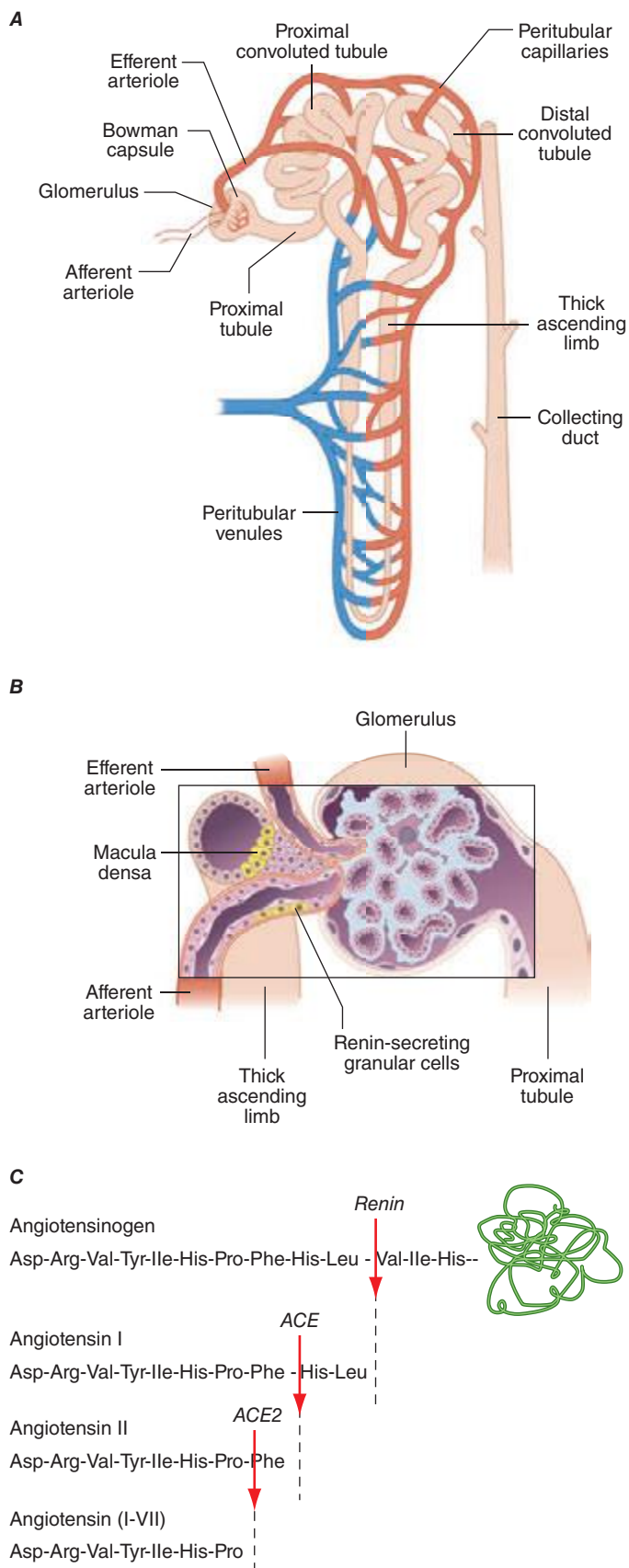
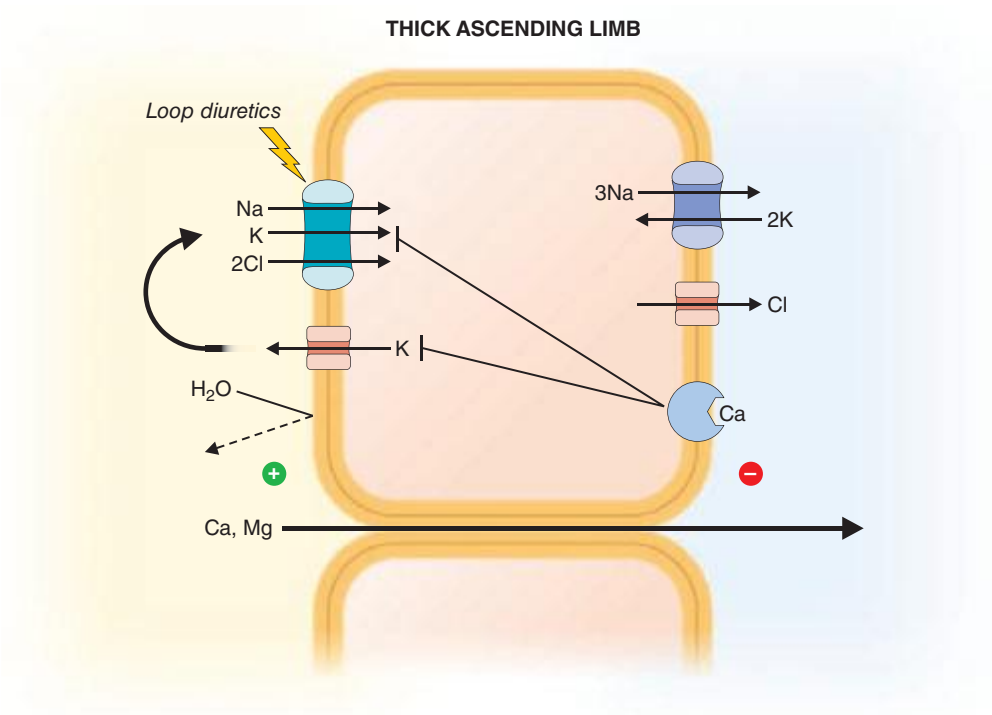
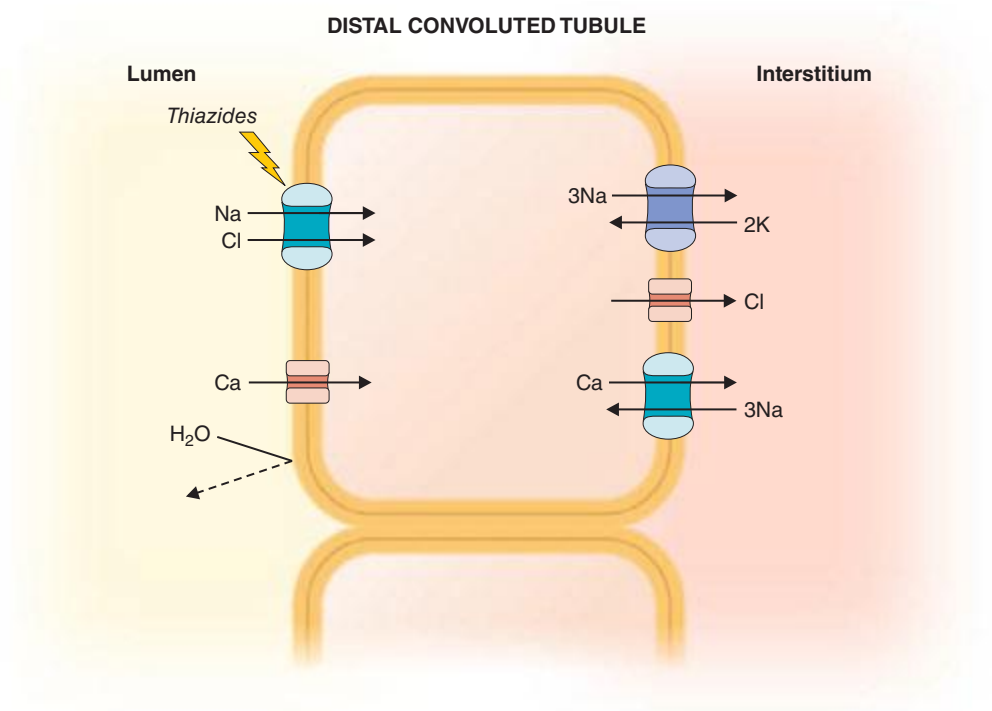


FIGURE 332e-2 Renal microcirculation and the renin-angiotensin system. **A.** Diagram illustrating relationships of the nephron with glomerular and peritubular capillaries. **B.** Expanded view of the glomerulus with its juxtaglomerular apparatus including the macula densa and adjacent afferent arteriole. **C.** Proteolytic processing steps in the generation of angiotensins.

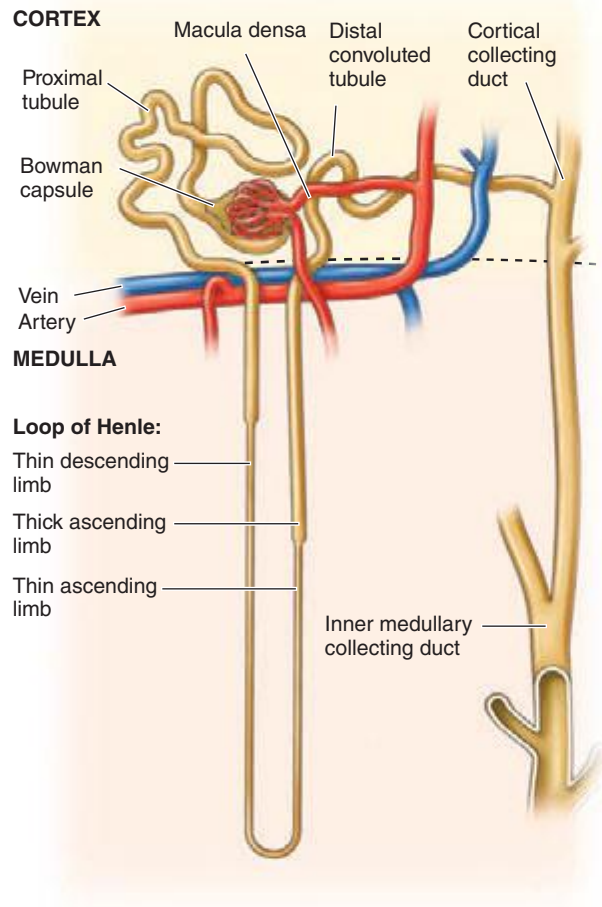


B

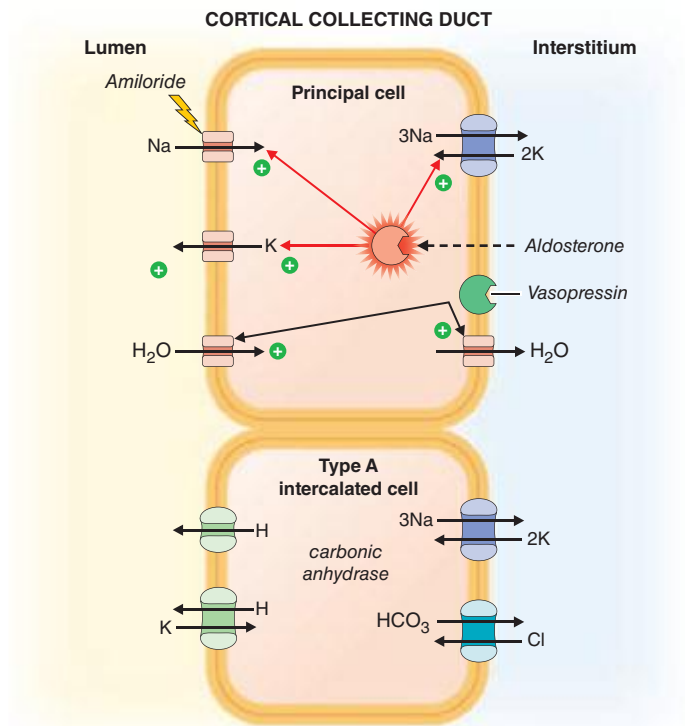


C

FIGURE 332e-3 (Continued)

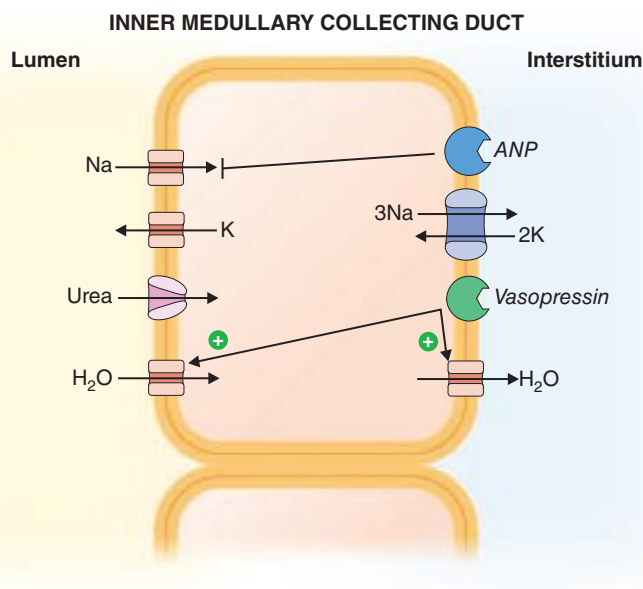


D



E

FIGURE 332e-3 (Continued)



F

FIGURE 332e-3 (Continued)

contains leaky epithelia, whereas distal nephron segments, such as the collecting duct, contain tight epithelia. Leaky epithelia are most well suited for bulk fluid reabsorption, whereas tight epithelia allow for more refined control and regulation of transport.

MEMBRANE TRANSPORT

Cell membranes are composed of hydrophobic lipids that repel water and aqueous solutes. The movement of solutes and water across cell membranes is made possible by discrete classes of integral membrane proteins, including channels, pumps, and transporters. These different mechanisms mediate specific types of transport activities, including *active transport* (pumps), *passive transport* (channels), *facilitated diffusion* (transporters), and *secondary active transport* (cotransporters). Active transport requires metabolic energy generated by the hydrolysis of ATP. Active transport pumps are ion-translocating ATPases, including the ubiquitous Na⁺/K⁺-ATPase, the H⁺-ATPases, and Ca²⁺-ATPases. Active transport creates asymmetric ion concentrations across a cell membrane and can move ions against a chemical gradient. The potential energy stored in a concentration gradient of an ion such as Na⁺ can be used to drive transport through other mechanisms (secondary active transport). Pumps are often *electrogenic*, meaning they can create an asymmetric distribution of electrostatic charges across the membrane and establish a voltage or membrane potential. The movement of solutes through a membrane protein by simple diffusion is called passive transport. This activity is mediated by channels created by selectively permeable membrane proteins, and it allows solute or water to move across a membrane driven by favorable *concentration gradients* or *electrochemical potential*. Facilitated diffusion is a specialized type of passive transport mediated by simple transporters called *carriers* or *uniporters*. For example, hexose transporters such as GLUT2 mediate glucose transport by tubular cells. These transporters are driven by the concentration gradient for glucose that is highest in extracellular fluids and lowest in the cytoplasm due to rapid metabolism. Many other transporters operate by translocating two or more ions/solutes in concert either in the same direction (*symporters* or *cotransporters*) or in opposite directions (*antiporters* or *exchangers*) across the cell membrane. The movement of two or more ions/solutes may produce no net change in the balance of electrostatic charges across the membrane (*electroneutral*), or a transport event may alter the balance of charges (*electrogenic*). Several inherited disorders of renal tubular solute and water transport occur as a consequence of mutations in genes encoding a variety of channels, transporter proteins, and their regulators ([Table 332e-1](#)).

SEGMENTAL NEPHRON FUNCTIONS

Each anatomic segment of the nephron has unique characteristics and specialized functions enabling selective transport of solutes and water (Fig. 332e-3). Through sequential events of reabsorption and secretion along the nephron, tubular fluid is progressively conditioned into urine. Knowledge of the major tubular mechanisms responsible for solute and water transport is critical for understanding hormonal regulation of kidney function and the pharmacologic manipulation of renal excretion.

PROXIMAL TUBULE

The proximal tubule is responsible for reabsorbing ~60% of filtered NaCl and water, as well as ~90% of filtered bicarbonate and most critical nutrients such as glucose and amino acids. The proximal tubule uses both cellular and paracellular transport mechanisms. The apical membrane of proximal tubular cells has an expanded surface area available for reabsorptive work created by a dense array of microvilli called the *brush border*, and leaky tight junctions enable high-capacity fluid reabsorption.

Solute and water pass through these tight junctions to enter the lateral intercellular space where absorption by the peritubular capillaries occurs. Bulk fluid reabsorption by the proximal tubule is driven by high oncotic pressure and low hydrostatic pressure within the peritubular capillaries. Cellular transport of most solutes by the proximal tubule is coupled to the Na⁺ concentration gradient established by the activity of a basolateral Na⁺/K⁺-ATPase (Fig. 332e-3A). This active transport mechanism maintains a steep Na⁺ gradient by keeping intracellular Na⁺ concentrations low. Solute reabsorption is coupled to the Na⁺ gradient by Na⁺-dependent transporters such as Na⁺-glucose and Na⁺-phosphate cotransporters. In addition to the paracellular route, water reabsorption also occurs through the cellular pathway enabled by constitutively active water channels (aquaporin-1) present on both apical and basolateral membranes.

Proximal tubular cells reclaim bicarbonate by a mechanism dependent on carbonic anhydrases. Filtered bicarbonate is first titrated by protons delivered to the lumen by Na⁺/H⁺ exchange. The resulting carbonic acid (H₂CO₃) is metabolized by brush border carbonic anhydrase to water and carbon dioxide. Dissolved carbon dioxide then diffuses into the cell, where it is enzymatically hydrated by cytoplasmic carbonic anhydrase to re-form carbonic acid. Finally, intracellular carbonic acid dissociates into free protons and bicarbonate anions, and bicarbonate exits the cell through a basolateral Na⁺/HCO₃⁻ cotransporter.

TABLE 332e-1 INHERITED DISORDERS AFFECTING RENAL TUBULAR ION AND SOLUTE TRANSPORT

Disease or Syndrome	Gene	OMIM ^a
Disorders Involving the Proximal Tubule		
Proximal renal tubular acidosis	Sodium bicarbonate cotransporter (<i>SLC4A4</i> , 4q21)	604278
Fanconi-Bickel syndrome	Glucose transporter, GLUT2 (<i>SLC2A2</i> , 3q26.2)	227810
Isolated renal glycosuria	Sodium glucose cotransporter (<i>SLC5A2</i> , 16p11.2)	233100
Cystinuria		
Type I	Cystine, dibasic and neutral amino acid transporter (<i>SLC3A1</i> , 2p16.3)	220100
Non-type I	Amino acid transporter, light subunit (<i>SLC7A9</i> , 19q13.1)	600918
Lysinuric protein intolerance	Amino acid transporter (<i>SLC7A7</i> , 4q11.2)	222700
Hartnup disorder	Neutral amino acid transporter (<i>SLC6A19</i> , 5p15.33)	34500
Hereditary hypophosphatemic rickets with hypercalcemia	Sodium phosphate cotransporter (<i>SLC34A3</i> , 9q34)	241530
Renal hypouricemia		
Type 1	Urate-anion exchanger (<i>SLC22A12</i> , 11q13)	220150
Type 2	Urate transporter, GLUT9 (<i>SLC2A9</i> , 4p16.1)	612076
Dent disease	Chloride channel, ClC-5 (<i>CLCN5</i> , Xp11.22)	300009
X-linked recessive nephrolithiasis with renal failure	Chloride channel, ClC-5 (<i>CLCN5</i> , Xp11.22)	310468
X-linked recessive hypophosphatemic rickets	Chloride channel, ClC-5 (<i>CLCN5</i> , Xp11.22)	307800
Disorders Involving the Loop of Henle		
Bartter syndrome		
Type 1	Sodium, potassium chloride cotransporter (<i>SLC12A1</i> , 15q21.1)	241200
Type 2	Potassium channel, ROMK (<i>KCNJ1</i> , 11q24)	601678
Type 3	Chloride channel, ClC-Kb (<i>CLCNKB</i> , 1p36)	602023
with sensorineural deafness	Chloride channel accessory subunit, Barttin (<i>BSND</i> , 1p31)	602522
Autosomal dominant hypocalcemia with Bartter-like syndrome	Calcium-sensing receptor (<i>CASR</i> , 3q13.33)	601199
Familial hypocalciuric hypercalcemia	Calcium-sensing receptor (<i>CASR</i> , 3q13.33)	145980
Primary hypomagnesemia	Claudin-16 or paracellin-1 (<i>CLDN16</i> or <i>PCLN1</i> , 3q27)	248250
Isolated renal magnesium loss	Sodium potassium ATPase, γ_1 -subunit (<i>ATP1G1</i> , 11q23)	154020
Disorders Involving the Distal Tubule and Collecting Duct		
Gitelman syndrome	Sodium chloride cotransporter (<i>SLC12A3</i> , 16q13)	263800
Primary hypomagnesemia with secondary hypocalcemia	Melastatin-related transient receptor potential cation channel 6 (<i>TRPM6</i> , 9q22)	602014
Pseudoaldosteronism (Liddle's syndrome)	Epithelial sodium channel β and γ subunits (<i>SCNN1B</i> , <i>SCNN1G</i> , 16p12.1)	177200
Recessive pseudohypoaldosteronism type 1	Epithelial sodium channel, α , β , and γ subunits (<i>SCNN1A</i> , 12p13; <i>SCNN1B</i> , <i>SCNN1G</i> , 16pp12.1)	264350
Pseudohypoaldosteronism type 2 (Gordon's hyperkalemia-hypertension syndrome)	Kinases WNK-1, WNK-4 (<i>WNK1</i> , 12p13; <i>WNK4</i> , 17q21.31)	145260
X-linked nephrogenic diabetes insipidus	Vasopressin V2 receptor (<i>AVPR2</i> , Xq28)	304800
Nephrogenic diabetes insipidus (autosomal)	Water channel, aquaporin-2 (<i>AQP2</i> , 12q13)	125800
Distal renal tubular acidosis		
autosomal dominant	Anion exchanger-1 (<i>SLC4A1</i> , 17q21.31)	179800
autosomal recessive	Anion exchanger-1 (<i>SLC4A1</i> , 17q21.31)	602722
with neural deafness	Proton ATPase, β 1 subunit (<i>ATP6V1B1</i> , 2p13.3)	192132
with normal hearing	Proton ATPase, 116-kD subunit (<i>ATP6V0A4</i> , 7q34)	602722

^aOnline Mendelian Inheritance in Man database (<http://www.ncbi.nlm.nih.gov/Omim>).

This process is saturable, resulting in urinary bicarbonate excretion when plasma levels exceed the physiologically normal range (24–26 meq/L). Carbonic anhydrase inhibitors such as acetazolamide, a class of weak diuretic agents, block proximal tubule reabsorption of bicarbonate and are useful for alkalinizing the urine.

The proximal tubule contributes to acid secretion by two mechanisms involving the titration of the urinary buffers ammonia (NH_3) and phosphate. Renal NH_3 is produced by glutamine metabolism in the proximal tubule. Subsequent diffusion of NH_3 out of the proximal tubular cell enables trapping of H^+ secreted by sodium-proton exchange in the lumen as ammonium ion (NH_4^+). Cellular K^+ levels inversely modulate proximal tubular ammoniogenesis, and in the setting of high serum K^+ from hypoaldosteronism, reduced ammoniogenesis facilitates the appearance of type IV renal tubular acidosis.

Filtered hydrogen phosphate ion (HPO_4^{2-}) is also titrated in the proximal tubule by secreted H^+ to form H_2PO_4^- , and this reaction constitutes a major component of the urinary buffer referred to as titratable acid. Most filtered phosphate ion is reabsorbed by the proximal tubule through a sodium-coupled cotransport process that is regulated by parathyroid hormone.

Chloride is poorly reabsorbed throughout the first segment of the proximal tubule, and a rise in Cl^- concentration counterbalances the removal of bicarbonate anion from tubular fluid. In later proximal tubular segments, cellular Cl^- reabsorption is initiated by apical exchange of cellular formate for higher luminal concentrations of Cl^- . Once in the lumen, formate anions are titrated by H^+ (provided by Na^+/H^+ exchange) to generate neutral formic acid, which can diffuse passively across the apical membrane back into the cell where it

dissociates a proton and is recycled. Basolateral Cl^- exit is mediated by a K^+/Cl^- cotransporter.

Reabsorption of glucose is nearly complete by the end of the proximal tubule. Cellular transport of glucose is mediated by apical Na^+ -glucose cotransport coupled with basolateral, facilitated diffusion by a glucose transporter. This process is also saturable, leading to glycosuria when plasma levels exceed 180–200 mg/dL, as seen in untreated diabetes mellitus.

The proximal tubule possesses specific transporters capable of secreting a variety of organic acids (carboxylate anions) and bases (mostly primary amine cations). Organic anions transported by these systems include urate, dicarboxylic acid anions (succinate), ketoacid anions, and several protein-bound drugs not filtered at the glomerulus (penicillins, cephalosporins, and salicylates). Probenecid inhibits renal organic anion secretion and can be clinically useful for raising plasma concentrations of certain drugs like penicillin and oseltamivir. Organic cations secreted by the proximal tubule include various biogenic amine neurotransmitters (dopamine, acetylcholine, epinephrine, norepinephrine, and histamine) and creatinine. The ATP-dependent transporter P-glycoprotein is highly expressed in brush border membranes and secretes several medically important drugs, including cyclosporine, digoxin, tacrolimus, and various cancer chemotherapeutic agents. Certain drugs like cimetidine and trimethoprim compete with endogenous compounds for transport by the organic cation pathways. Although these drugs elevate serum creatinine levels, there is no change in the actual GFR.

The proximal tubule, through distinct classes of Na^+ -dependent and Na^+ -independent transport systems, reabsorbs amino acids efficiently. These transporters are specific for different groups of amino acids. For example, cystine, lysine, arginine, and ornithine are transported by a system comprising two proteins encoded by the *SLC3A1* and *SLC7A9* genes. Mutations in either *SLC3A1* or *SLC7A9* impair reabsorption of these amino acids and cause the disease cystinuria. Peptide hormones, such as insulin and growth hormone, β_2 -microglobulin, albumin, and other small proteins, are taken up by the proximal tubule through a process of absorptive endocytosis and are degraded in acidified endocytic lysosomes. Acidification of these vesicles depends on a vacuolar H^+ -ATPase and Cl^- channel. Impaired acidification of endocytic vesicles because of mutations in a Cl^- channel gene (*CLCN5*) causes low-molecular-weight proteinuria in Dent disease.

LOOP OF HENLE

The loop of Henle consists of three major segments: descending thin limb, ascending thin limb, and ascending thick limb. These divisions are based on cellular morphology and anatomic location, but also correlate with specialization of function. Approximately 15–25% of filtered NaCl is reabsorbed in the loop of Henle, mainly by the thick ascending limb. The loop of Henle has an important role in urinary concentration by contributing to the generation of a hypertonic medullary interstitium in a process called *countercurrent multiplication*. The loop of Henle is the site of action for the most potent class of diuretic agents (loop diuretics) and also contributes to reabsorption of calcium and magnesium ions.

The descending thin limb is highly water permeable owing to dense expression of constitutively active aquaporin-1 water channels. By contrast, water permeability is negligible in the ascending limb. In the thick ascending limb, there is a high level of secondary active salt transport enabled by the $\text{Na}^+/\text{K}^+/2\text{Cl}^-$ cotransporter on the apical membrane in series with basolateral Cl^- channels and Na^+/K^+ -ATPase (Fig. 332e-3B). The $\text{Na}^+/\text{K}^+/2\text{Cl}^-$ cotransporter is the primary target for loop diuretics. Tubular fluid K^+ is the limiting substrate for this cotransporter (tubular concentration of K^+ is similar to plasma, about 4 meq/L), but transporter activity is maintained by K^+ recycling through an apical potassium channel. The cotransporter also enables reabsorption of NH_4^+ in lieu of K^+ , and this leads to accumulation of both NH_4^+ and NH_3 in the medullary interstitium. An inherited disorder of the thick ascending limb, Bartter syndrome, also results in a salt-wasting renal disease associated with hypokalemia and metabolic alkalosis;

loss-of-function mutations in one of five distinct genes encoding components of the $\text{Na}^+/\text{K}^+/2\text{Cl}^-$ cotransporter (*NKCC2*), apical K^+ channel (*KCNJ1*), basolateral Cl^- channel (*CLCNKB*, *BSND*), or calcium-sensing receptor (*CASR*) can cause Bartter syndrome.

Potassium recycling also contributes to a positive electrostatic charge in the lumen relative to the interstitium that promotes divalent cation (Mg^{2+} and Ca^{2+}) reabsorption through a paracellular pathway. A Ca^{2+} -sensing, G-protein-coupled receptor (CaSR) on basolateral membranes regulates NaCl reabsorption in the thick ascending limb through dual signaling mechanisms using either cyclic AMP or eicosanoids. This receptor enables a steep relationship between plasma Ca^{2+} levels and renal Ca^{2+} excretion. Loss-of-function mutations in CaSR cause familial hypercalcemic hypocalciuria because of a blunted response of the thick ascending limb to extracellular Ca^{2+} . Mutations in *CLDN16* encoding paracellin-1, a transmembrane protein located within the tight junction complex, leads to familial hypomagnesemia with hypercalciuria and nephrocalcinosis, suggesting that the ion conductance of the paracellular pathway in the thick limb is regulated.

The loop of Henle contributes to urine-concentrating ability by establishing a *hypertonic medullary interstitium* that promotes water reabsorption by the downstream inner medullary collecting duct. *Countercurrent multiplication* produces a hypertonic medullary interstitium using two countercurrent systems: the loop of Henle (opposing descending and ascending limbs) and the vasa recta (medullary peritubular capillaries enveloping the loop). The countercurrent flow in these two systems helps maintain the hypertonic environment of the inner medulla, but NaCl reabsorption by the thick ascending limb is the primary initiating event. Reabsorption of NaCl without water dilutes the tubular fluid and adds new osmoles to medullary interstitial fluid. Because the descending thin limb is highly water permeable, osmotic equilibrium occurs between the descending limb tubular fluid and the interstitial space, leading to progressive solute trapping in the inner medulla. Maximum medullary interstitial osmolality also requires partial recycling of urea from the collecting duct.

DISTAL CONVOLUTED TUBULE

The distal convoluted tubule reabsorbs ~5% of the filtered NaCl . This segment is composed of a tight epithelium with little water permeability. The major NaCl -transporting pathway uses an apical membrane, electroneutral thiazide-sensitive Na^+/Cl^- cotransporter in tandem with basolateral Na^+/K^+ -ATPase and Cl^- channels (Fig. 332e-3C). Apical Ca^{2+} -selective channels (TRPV5) and basolateral $\text{Na}^+/\text{Ca}^{2+}$ exchange mediate calcium reabsorption in the distal convoluted tubule. Ca^{2+} reabsorption is inversely related to Na^+ reabsorption and is stimulated by parathyroid hormone. Blocking apical Na^+/Cl^- cotransport will reduce intracellular Na^+ , favoring increased basolateral $\text{Na}^+/\text{Ca}^{2+}$ exchange and passive apical Ca^{2+} entry. Loss-of-function mutations of *SLC12A3* encoding the apical Na^+/Cl^- cotransporter cause Gitelman syndrome, a salt-wasting disorder associated with hypokalemic alkalosis and hypocalciuria. Mutations in genes encoding WNK kinases, WNK-1 and WNK-4, cause pseudohypoaldosteronism type II or Gordon syndrome characterized by familial hypertension with hyperkalemia. WNK kinases influence the activity of several tubular ion transporters. Mutations in this disorder lead to overactivity of the apical Na^+/Cl^- cotransporter in the distal convoluted tubule as the primary stimulus for increased salt reabsorption, extracellular volume expansion, and hypertension. Hyperkalemia may be caused by diminished activity of apical K^+ channels in the collecting duct, a primary route for K^+ secretion. Mutations in *TRPM6* encoding Mg^{2+} permeable ion channels also cause familial hypomagnesemia with hypocalcemia. A molecular complex of TRPM6 and TRPM7 proteins is critical for Mg^{2+} reabsorption in the distal convoluted tubule.

COLLECTING DUCT

The collecting duct modulates the final composition of urine. The two major divisions, the cortical collecting duct and inner medullary collecting duct, contribute to reabsorbing ~4–5% of filtered Na^+ and are important for hormonal regulation of salt and water balance. The

cortical collecting duct contains *high-resistance epithelia* with two cell types. Principal cells are the main water, Na^+ -reabsorbing, and K^+ -secreting cells, and the site of action of aldosterone, K^+ -sparing diuretics, and mineralocorticoid receptor antagonists such as spironolactone. The other cells are type A and B intercalated cells. Type A intercalated cells mediate acid secretion and bicarbonate reabsorption also under the influence of aldosterone. Type B intercalated cells mediate bicarbonate secretion and acid reabsorption.

Virtually all transport is mediated through the cellular pathway for both principal cells and intercalated cells. In principal cells, passive apical Na^+ entry occurs through the amiloride-sensitive, epithelial Na^+ channel (ENaC) with basolateral exit via the Na^+/K^+ -ATPase (Fig. 332e-3E). This Na^+ reabsorptive process is tightly regulated by aldosterone and is physiologically activated by a variety of proteolytic enzymes that cleave extracellular domains of ENaC; plasmin in the tubular fluid of nephrotic patients, for example, activates ENaC, leading to sodium retention. Aldosterone enters the cell across the basolateral membrane, binds to a cytoplasmic mineralocorticoid receptor, and then translocates into the nucleus, where it modulates gene transcription, resulting in increased Na^+ reabsorption and K^+ secretion. Activating mutations in ENaC increase Na^+ reclamation and produce hypokalemia, hypertension, and metabolic alkalosis (Liddle's syndrome). The potassium-sparing diuretics amiloride and triamterene block ENaC, causing reduced Na^+ reabsorption.

Principal cells secrete K^+ through an apical membrane potassium channel. Several forces govern the secretion of K^+ . Most importantly, the high intracellular K^+ concentration generated by Na^+/K^+ -ATPase creates a favorable concentration gradient for K^+ secretion into tubular fluid. With reabsorption of Na^+ without an accompanying anion, the tubular lumen becomes negative relative to the cell interior, creating a favorable electrical gradient for secretion of potassium. When Na^+ reabsorption is blocked, the electrical component of the driving force for K^+ secretion is blunted, and this explains lack of excess urinary K^+ loss during treatment with potassium-sparing diuretics or mineralocorticoid receptor antagonists. K^+ secretion is also promoted by aldosterone actions that increase regional Na^+ transport favoring more electronegativity and by increasing the number and activity of potassium channels. Fast tubular fluid flow rates that occur during volume expansion or diuretics acting "upstream" of the cortical collecting duct also increase K^+ secretion, as does the presence of relatively nonreabsorbable anions (including bicarbonate and semisynthetic penicillins) that contribute to the lumen-negative potential. Off-target effects of certain antibiotics, such as trimethoprim and pentamidine, block ENaCs and predispose to hyperkalemia, especially when renal K^+ handling is impaired for other reasons. Principal cells, as described below, also participate in water reabsorption by increased water permeability in response to vasopressin.

Intercalated cells do not participate in Na^+ reabsorption but, instead, mediate acid-base secretion. These cells perform two types of transport: active H^+ transport mediated by H^+ -ATPase (proton pump), and $\text{Cl}^-/\text{HCO}_3^-$ exchange. Intercalated cells arrange the two transport mechanisms on opposite membranes to enable either acid or base secretion. Type A intercalated cells have an apical proton pump that mediates acid secretion and a basolateral $\text{Cl}^-/\text{HCO}_3^-$ anion exchanger for bicarbonate reabsorption (Fig. 332e-3E); aldosterone increases the number of H^+ -ATPase pumps, sometimes contributing to the development of metabolic alkalosis. Secreted H^+ is buffered by NH_3 that has diffused into the collecting duct lumen from the surrounding interstitium. By contrast, type B intercalated cells have the anion exchanger on the apical membrane to mediate bicarbonate secretion while the proton pump resides on the basolateral membrane to enable acid reabsorption. Under conditions of acidemia, the kidney preferentially uses type A intercalated cells to secrete the excess H^+ and generate more HCO_3^- . The opposite is true in states of bicarbonate excess with alkalemia where the type B intercalated cells predominate. An extracellular protein called *hensin* mediates this adaptation.

Inner medullary collecting duct cells share many similarities with principal cells of the cortical collecting duct. They have apical Na^+ and K^+ channels that mediate Na^+ reabsorption and K^+ secretion,

respectively (Fig. 332e-3F). Inner medullary collecting duct cells also have vasopressin-regulated water channels (aquaporin-2 on the apical membrane, aquaporin-3 and -4 on the basolateral membrane). The antidiuretic hormone vasopressin binds to the V2 receptor on the basolateral membrane and triggers an intracellular signaling cascade through G-protein-mediated activation of adenylyl cyclase, resulting in an increase in the cellular levels of cyclic AMP. This signaling cascade stimulates the insertion of water channels into the apical membrane of the inner medullary collecting duct cells to promote increased water permeability. This increase in permeability enables water reabsorption and production of concentrated urine. In the absence of vasopressin, inner medullary collecting duct cells are water impermeable, and urine remains dilute.

Sodium reabsorption by inner medullary collecting duct cells is also inhibited by the natriuretic peptides called *atrial natriuretic peptide* or *renal natriuretic peptide* (urodilatin); the same gene encodes both peptides but uses different posttranslational processing of a common prohormone to generate different proteins. Atrial natriuretic peptides are secreted by atrial myocytes in response to volume expansion, whereas urodilatin is secreted by renal tubular epithelia. Natriuretic peptides interact with either apical (urodilatin) or basolateral (atrial natriuretic peptides) receptors on inner medullary collecting duct cells to stimulate guanylyl cyclase and increase levels of cytoplasmic cGMP. This effect in turn reduces the activity of the apical Na^+ channel in these cells and attenuates net Na^+ reabsorption, producing natriuresis.

The inner medullary collecting duct transports urea out of the lumen, returning urea to the interstitium, where it contributes to the hypertonicity of the medullary interstitium. Urea is recycled by diffusing from the interstitium into the descending and ascending limbs of the loop of Henle.

HORMONAL REGULATION OF SODIUM AND WATER BALANCE

The balance of solute and water in the body is determined by the amounts ingested, distributed to various fluid compartments, and excreted by skin, bowel, and kidneys. *Tonicity*, the osmolar state determining the volume behavior of cells in a solution, is regulated by water balance (Fig. 332e-4A), and *extracellular blood volume* is regulated by Na^+ balance (Fig. 332e-4B). The kidney is a critical modulator of both physiologic processes.

WATER BALANCE

Tonicity depends on the variable concentration of *effective osmoles* inside and outside the cell causing water to move in either direction across its membrane. Classic effective osmoles, like Na^+ , K^+ , and their anions, are solutes trapped on either side of a cell membrane, where they collectively partition and obligate water to move and find equilibrium in proportion to retained solute; Na^+/K^+ -ATPase keeps most K^+ inside cells and most Na^+ outside. Normal tonicity (~280 mosmol/L) is rigorously defended by osmoregulatory mechanisms that control water balance to protect tissues from inadvertent *dehydration* (cell shrinkage) or *water intoxication* (cell swelling), both of which are deleterious to cell function (Fig. 332e-4A).

The mechanisms that control osmoregulation are distinct from those governing extracellular volume, although there is some shared physiology in both processes. While cellular concentrations of K^+ have a determinant role in any level of tonicity, the routine surrogate marker for assessing clinical tonicity is the concentration of serum Na^+ . Any reduction in total body water, which raises the Na^+ concentration, triggers a brisk sense of thirst and conservation of water by decreasing renal water excretion mediated by release of vasopressin from the posterior pituitary. Conversely, a decrease in plasma Na^+ concentration triggers an increase in renal water excretion by suppressing the secretion of vasopressin. Whereas all cells expressing mechanosensitive TRPV1, 2, or 4 channels, among potentially other sensors, respond to changes in tonicity by altering their volume and Ca^{2+} concentration, only TRPV⁺ neuronal cells connected to the organum vasculosum of the lamina terminalis are *osmoreceptive*. Only these cells, because of their neural

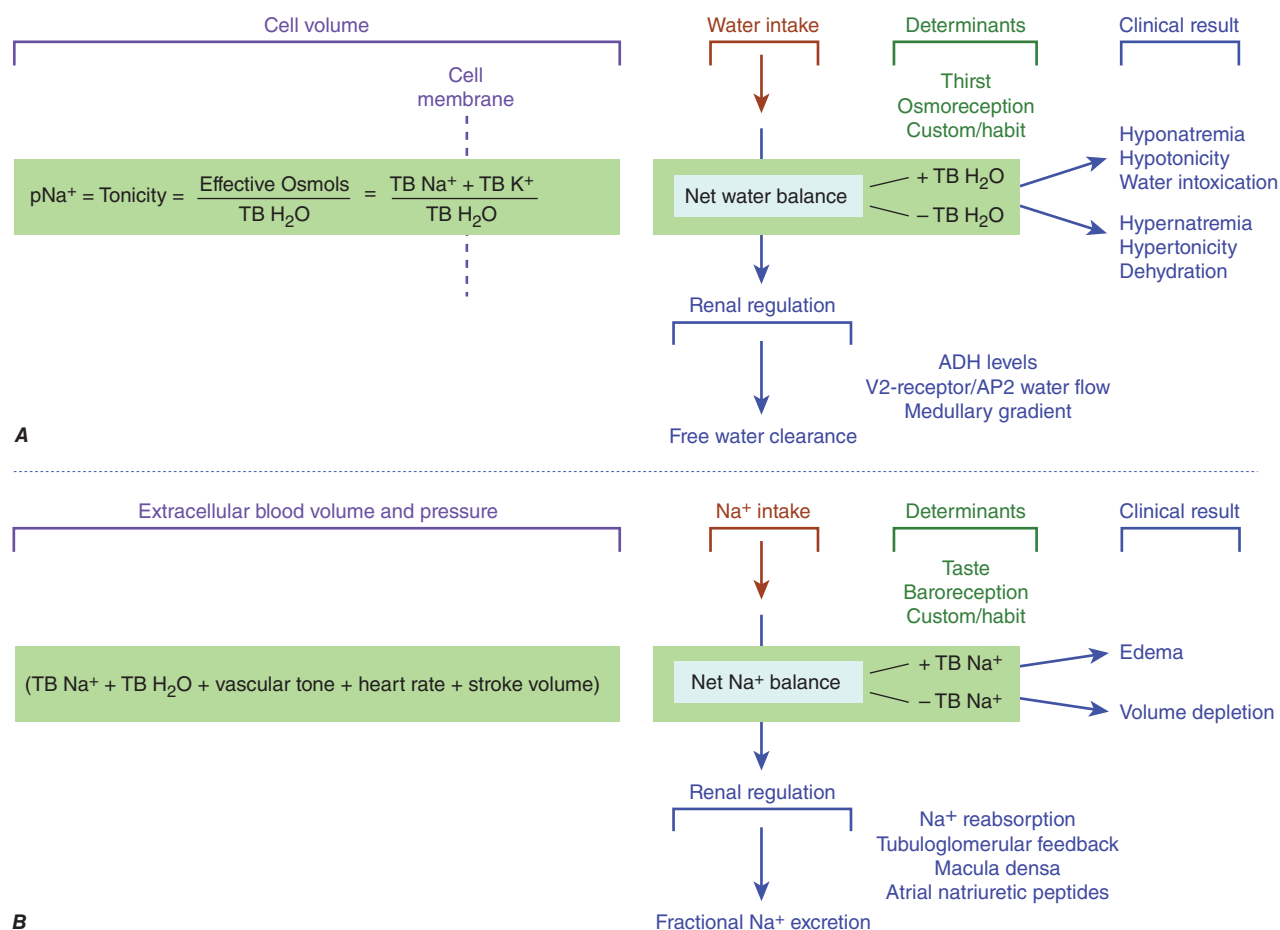


FIGURE 332e-4 Determinants of sodium and water balance. A. Plasma Na⁺ concentration is a surrogate marker for plasma tonicity, the volume behavior of cells in a solution. Tonicity is determined by the number of effective osmoles in the body divided by the total body H₂O (TB H₂O), which translates simply into the total body Na (TB Na⁺) and anions outside the cell separated from the total body K (TB K⁺) inside the cell by the cell membrane. Net water balance is determined by the integrated functions of thirst, osmoreception, Na reabsorption, vasopressin release, and the strength of the medullary gradient in the kidney, keeping tonicity within a narrow range of osmolality around 280 mosmol/L. When water metabolism is disturbed and total body water increases, hyponatremia, hypotonicity, and water intoxication occur; when total body water decreases, hypernatremia, hypertonicity, and dehydration occur. **B.** Extracellular blood volume and pressure are an integrated function of total body Na⁺ (TB Na⁺), total body H₂O (TB H₂O), vascular tone, heart rate, and stroke volume that modulates volume and pressure in the vascular tree of the body. This extracellular blood volume is determined by net Na balance under the control of taste, baroreception, habit, Na⁺ reabsorption, macula densa/tubuloglomerular feedback, and natriuretic peptides. When Na⁺ metabolism is disturbed and total body Na⁺ increases, edema occurs; when total body Na⁺ is decreased, volume depletion occurs. ADH, antidiuretic hormone; AQP2, aquaporin-2.

connectivity and adjacency to a minimal blood-brain barrier, modulate the downstream release of vasopressin by the posterior lobe of the pituitary gland. Secretion is stimulated primarily by changing tonicity and secondarily by other nonosmotic signals such as variable blood volume, stress, pain, nausea, and some drugs. The release of vasopressin by the posterior pituitary increases linearly as plasma tonicity rises above normal, although this varies, depending on the perception of extracellular volume (one form of cross-talk between mechanisms that adjudicate blood volume and osmoregulation). Changing the intake or excretion of water provides a means for adjusting plasma tonicity; thus, osmoregulation governs water balance.

The kidneys play a vital role in maintaining water balance through the regulation of renal water excretion. The ability to concentrate urine to an osmolality exceeding that of plasma enables water conservation, whereas the ability to produce urine more dilute than plasma promotes excretion of excess water. For water to enter or exit a cell, the cell membrane must express aquaporins. In the kidney, aquaporin-1 is constitutively active in all water-permeable segments of the proximal and distal tubules, whereas vasopressin-regulated aquaporin-2, -3, and -4 in the inner medullary collecting duct promote rapid water permeability. Net water reabsorption is ultimately driven by the osmotic gradient between dilute tubular fluid and a hypertonic medullary interstitium.

SODIUM BALANCE

The perception of *extracellular blood volume* is determined, in part, by the integration of arterial tone, cardiac stroke volume, heart rate, and the water and solute content of extracellular fluid. Na⁺ and accompanying anions are the most abundant extracellular effective osmoles and together support a blood volume around which pressure is generated. Under normal conditions, this volume is regulated by sodium balance (Fig. 332e-4B), and the balance between daily Na⁺ intake and excretion is under the influence of *baroreceptors* in regional blood vessels and vascular hormone sensors modulated by atrial natriuretic peptides, the renin-angiotensin-aldosterone system, Ca²⁺ signaling, adenosine, vasopressin, and the neural adrenergic axis. If Na⁺ intake exceeds Na⁺ excretion (positive Na⁺ balance), then an increase in blood volume will trigger a proportional increase in urinary Na⁺ excretion. Conversely, when Na⁺ intake is less than urinary excretion (negative Na⁺ balance), blood volume will decrease and trigger enhanced renal Na⁺ reabsorption, leading to decreased urinary Na⁺ excretion.

The renin-angiotensin-aldosterone system is the best-understood hormonal system modulating renal Na⁺ excretion. Renin is synthesized and secreted by granular cells in the wall of the afferent arteriole. Its secretion is controlled by several factors, including β₁-adrenergic

stimulation to the afferent arteriole, input from the macula densa, and prostaglandins. Renin and ACE activity eventually produce angiotensin II that directly or indirectly promotes renal Na^+ and water reabsorption. Stimulation of proximal tubular Na^+/H^+ exchange by angiotensin II directly increases Na^+ reabsorption. Angiotensin II also promotes Na^+ reabsorption along the collecting duct by stimulating aldosterone secretion by the adrenal cortex. Constriction of the efferent glomerular arteriole by angiotensin II indirectly increases the filtration fraction and raises peritubular capillary oncotic pressure to promote tubular Na^+ reabsorption. Finally, angiotensin II inhibits renin secretion through a negative feedback loop. Alternative metabolism of angiotensin by ACE2 generates the vasodilatory peptide angiotensin 1-7 that acts through Mas receptors to counterbalance several actions of angiotensin II on blood pressure and renal function (Fig. 332e-2C).

Aldosterone is synthesized and secreted by granulosa cells in the adrenal cortex. It binds to cytoplasmic mineralocorticoid receptors in the collecting duct principal cells that increase activity of ENaC, apical

membrane K^+ channel, and basolateral Na^+/K^+ -ATPase. These effects are mediated in part by aldosterone-stimulated transcription of the gene encoding serum/glucocorticoid-induced kinase 1 (SGK1). The activity of ENaC is increased by SGK1-mediated phosphorylation of Nedd4-2, a protein that promotes recycling of the Na^+ channel from the plasma membrane. Phosphorylated Nedd4-2 has impaired interactions with ENaC, leading to increased channel density at the plasma membrane and increased capacity for Na^+ reabsorption by the collecting duct.

Chronic exposure to aldosterone causes a decrease in urinary Na^+ excretion lasting only a few days, after which Na^+ excretion returns to previous levels. This phenomenon, called *aldosterone escape*, is explained by decreased proximal tubular Na^+ reabsorption following blood volume expansion. Excess Na^+ that is not reabsorbed by the proximal tubule overwhelms the reabsorptive capacity of more distal nephron segments. This escape may be facilitated by atrial natriuretic peptides that lose their effectiveness in the clinical settings of heart failure, nephrotic syndrome, and cirrhosis, leading to severe Na^+ retention and volume overload.

333e Adaptation of the Kidney to Injury

Joseph V. Bonventre

Many years ago Claude Bernard (1878) introduced the concepts of *milieu extérieur* (the environment where an organism lives) and *milieu intérieur* (the environment in which the tissues of that organism live). He argued that the *milieu intérieur* varied very little and that there were vital mechanisms that functioned to maintain this internal environment constant. Walter B. Cannon later extended these concepts by recognizing that the constancy of the internal state, which he termed the *homeostatic state*, was evidence of physiologic mechanisms that act to maintain this minimal variability. In higher animals, the plasma is maintained remarkably constant in composition both within an individual and among individuals. The kidney plays a vital role in this constancy. The kidney changes the composition of the urine to maintain electrolyte and acid-base balance and can produce hormones that can maintain constancy of blood hemoglobin and mineral metabolism. When the kidney is injured, the remaining functional mass responds and attempts to continue to maintain the *milieu intérieur*. It is remarkable how well the residual nephrons can perform in this task so that in many cases homeostasis is maintained until the glomerular filtration rate (GFR) drops to very low levels. At this point, the functional tissue can no longer compensate. In this chapter, we will discuss a number of these compensatory adaptations that the kidney makes in response to injury in an attempt to protect itself and protect the *milieu intérieur*. A theme that permeates, however, is that these adaptive processes can often be maladaptive and contribute to enhanced renal dysfunction, facilitating a positive feedback process that is inherently unstable.

RESPONSES OF THE KIDNEY TO REDUCED NUMBERS OF NEPHRONS DURING DEVELOPMENT

Renal disease is associated with a reduction in functional nephrons. The rest of the kidney adapts to this reduction by increasing blood flow to and the size of the remaining glomeruli and increasing size and function of the remaining tubules. Robert Platt, in 1936, argued that "...a high glomerular pressure, together with loss of nephrons (destroyed by disease) [is] an explanation of the peculiarities of renal function in this stage of kidney disease." The raised glomerular pressure will increase the amount of filtrate produced by each nephron and thus compensate for a time for the destruction of part of the kidney. But eventually there are too few nephrons remaining to produce an adequate filtrate, even though they may work under the highest possible pressure, associated with a high systemic blood pressure. The responses to kidney injury can be both adaptive and maladaptive, and in many cases, the early adaptive responses can become maladaptive over time, leading to progressive decline in the anatomic and functional integrity of the kidney. As described previously, the early responses are likely in many cases motivated by attempts to maintain the constancy of the *milieu intérieur* for the survival of the organism (Claude Bernard).

Barry Brenner in the 1960s and 1970s carried out micropuncture experiments to define the pressures in glomerular capillaries as well as afferent and efferent resistances and modeled the behavior of the factors that governed glomerular filtration in health and disease. According to the Brenner Hyperfiltration Hypothesis, a reduction in the number of nephrons results in glomerular hypertension, hyperfiltration, and enlargement of glomeruli and this hyperfiltration results in damage to those glomeruli over time and ultimately decreased kidney function. According to this hypothesis, a positive feedback process is set into motion whereby injury to the glomeruli will result in further hyperfiltration to other glomeruli and hence more accelerated injury to those glomeruli. Since nephrons are not generated after 34–36 weeks of gestation or after birth (if earlier than 34–36 weeks) in humans, this hypothesis implies a deterministic effect of low nephron numbers at birth. There is over a 10-fold variation in the number of nephrons per

kidney in the population (200,000 to over 2.5 million). This variation is not explained by kidney size in the adult. Children born with low birth weights would be more prone to kidney disease as adults. There are many reasons why there might be reduced nephron numbers at birth: developmental abnormalities, genetic predisposition, and environmental factors, such as malnutrition. There are thought to be interactions between these various factors. Reduced nephron mass can also occur with chronic kidney disease (CKD) in the adult, and the response of the kidney is similar qualitatively with hyperfiltration of the remaining nephrons.

Developmental Abnormalities There are many congenital abnormalities of the kidney and urinary tract (CAKUT). Dysplastic kidneys have varying degrees of abnormalities that interfere with their function. Anatomically abnormal kidneys can be associated with abnormalities of the lower urinary tract. Urinary tract abnormalities resulting in obstruction or vesicoureteric reflux can dramatically alter the normal development of the kidney nephrons. Dysplastic or hypoplastic kidneys can be cystic in patterns that are distinct from polycystic kidney disease. Of course, autosomal recessive kidney disease can result in widespread cyst formation.

Hypoplastic kidneys are characterized by a reduced number of functional nephrons. One definition of hypoplastic kidneys is as follows: "Kidney mass below two standard deviations of that of age-matched normal [individuals] or a combined kidney mass of less than half normal for the patient's age." Renal agenesis and cystic dysplasia often affects only one kidney. This results in hypertrophy of the other kidney if it is unaffected by any congenital abnormality itself. Although there is hypertrophy in size, it is not clear if this is associated with an increase in the number of nephrons on the contralateral side.

The prevalence of CAKUT has been generally found to be between 0.003 and 0.2%, depending on the population studied. This excludes fetuses with transient upper renal tract dilatation likely related to the high rate of fetal urine flow rate. In the adult U.S. Renal Data System (USRDS) of patients with end-stage kidney disease, approximately 0.6% are listed as having dysplastic or hypoplastic kidneys as a primary cause of the disease. This is likely an underestimate, however, because many patients with "small kidneys" may be misdiagnosed with chronic glomerulonephritis or chronic pyelonephritis.

Environmental Contributions to Reduced Nephron Mass The most important environmental factor responsible for reduced nephron number is growth restriction within the uterus. This has been associated with disease processes such as diabetes mellitus in the mother, but there also is a strong genetic disposition. Low-birth-weight children are more likely to be born to mothers who, themselves, were born with low birth weight. There are clearly other environmental factors. Caloric restriction during pregnancy in humans has been associated with altered glucose as adults and increased risk for hypertension. In one study, it was found that if women were calorie restricted in midgestation, the time of most rapid nephrogenesis, there was a threefold incidence of albuminuria in their children when they were tested as adults. Factors such as deficiency in vitamin A, sodium, zinc, or iron have been implicated as predisposing to abnormal kidney development. Other environmental factors that can influence kidney development are medications taken by the mother, such as dexamethasone, angiotensin-converting enzyme inhibitors, and angiotensin receptor antagonists (Table 333e-1). Protein restriction in mice during pregnancy can reduce lifespan of the offspring by 200 days. Obesity may play an important role in determining kidney outcome long term in patients with reduced kidney mass. It has been shown in mice fed a high-fat diet that the rodents that had reduced nephron number had a greater incidence of hypertension and renal fibrosis.

TABLE 333e-1 DRUGS THAT INHIBIT NEPHROGENESIS

Dexamethasone
Angiotensin-converting enzyme inhibitors
Angiotensin receptor blockers
Gentamicin
Nonsteroidal anti-inflammatory drugs

Implications of Low Nephron Number at Birth David Barker was the first to describe the association between low birth weight and later cardiovascular death. This was followed by studies relating low birth weight to risk for diabetes, stroke, hypertension, and CKD. It has been found that there is an inverse relationship between nephron number and blood pressure in adults. This relationship was found in Caucasians but not in African Americans. Approximately one-third of children with a single functioning kidney at the age of 10 years had signs of renal injury as determined by the presence of hypertension, albuminuria, or the use of renoprotective drugs. Another study revealed that 20–40% of patients born with a single functional kidney had renal failure requiring dialysis by 30 years of age.

ADAPTIVE RESPONSES OF THE KIDNEY TO REDUCED KIDNEY MASS THAT CHARACTERIZES CHRONIC KIDNEY DISEASE

In the early stages of CKD, there are many adaptations structurally and functionally that limit the consequences of the loss of nephrons on total-body homeostasis. In later stages of disease, however, these adaptations are insufficient to counteract the consequences of nephron loss and in fact often become maladaptive.

Counterbalance Renal counterbalance was defined by Hinman in 1923 as “an attempt on the part of the less injured or uninjured portion (of the kidney) to take over the work of the more injured portion.” Hinman defined “renal reserve” to be of two types: “native reserve, which is the normal physiological response to stimulation . . . and acquired reserve, which involves growth or compensation due to overstimulation.” It was known that removal of one kidney results in an increase in size of the contralateral kidney. If, instead of nephrectomy, one kidney is rendered ischemic and the other left intact, there is a resultant atrophy of the postischemic kidney. If the contralateral kidney is removed, however, before the atrophy becomes too severe, then the postischemic kidney increases markedly in size. With the contralateral kidney in place, there is vasoconstriction and reduced renal blood flow to the postischemic kidney. This is rapidly reversed, however, when the contralateral normal kidney is removed. The factors responsible for the persistent initial (prenephrectomy) vasoconstriction and those responsible for the rapid vasodilation and enhanced growth after contralateral nephrectomy are unknown.

Hypertrophy Because nephrons of mammals, in contrast to those of fish, cannot regenerate, the loss of functional units of the kidney, either due to disease or surgery, results in anatomic and functional changes in the remaining nephrons. As described above, there is increased blood flow to remaining glomeruli with potentially adverse effects over time of the resultant increased size of the remaining glomeruli and hyperfiltration (Fig. 333e-1). In addition, there is hypertrophy of the tubules. Some of the mediators of this hypertrophy of the remaining functional tubules are listed in Table 333e-2. In the adult, within a few weeks after unilateral nephrectomy for donation of a kidney, the GFR is approximately 70% of the pre-nephrectomy value. It then remains relatively stable for most patients over 15–20 years. The hyperfiltration is related to an increase in renal blood flow likely secondary to dilatation of the afferent arterioles potentially due to increases in nitric oxide (NO) production. The rate of increase in GFR is slower in the adult than it is in the young after nephrectomy. There are a number of factors that have been implicated at the cellular and nephron level to account for the compensatory hypertrophy that ensues after removal of functional nephrons (Table 333e-2).

With increased blood flow to the kidney, there is glomerular hypertension (i.e., an increase in glomerular capillary pressure). There is increased wall tension and force on the capillary wall that is counteracted by contractile properties of the endothelium and elastic properties of the glomerular basement membrane. The force is conveyed to podocytes, which adapt by reinforcing cell cycle arrest and increasing cell adhesion in an adaptive attempt to maintain the delicate architecture of the interdigitating foot processes. Over time, however, these increased forces due to glomerular hypertension lead to podocyte damage and glomerulosclerosis.

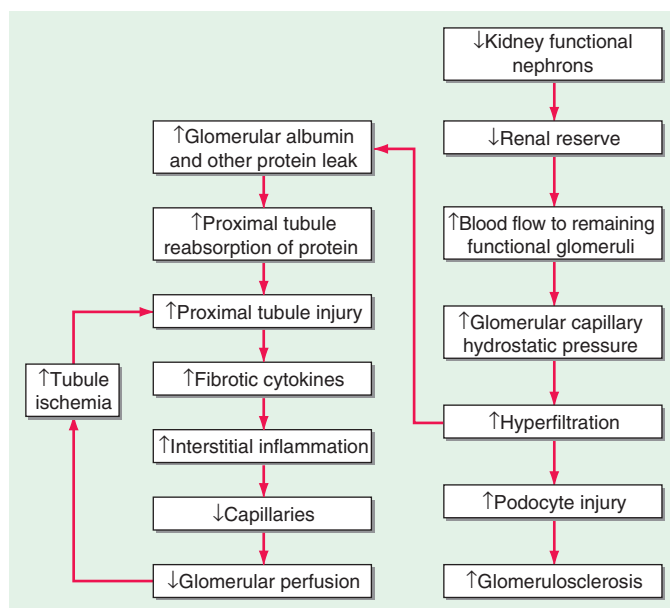


FIGURE 333e-1 Some of the pathophysiologic mechanisms involved with the maladaptive response to a reduction in the number of functional nephrons due to prenatal factors or postnatal disease processes.

Other Systemic and Renal Adaptations to Reduced Nephron Function With reduced functional nephrons, as is seen in CKD, there are many other systemic adaptations that occur to preserve the *milieu intérieur* because the kidney is involved in so many regulatory networks that are then stressed when there is dysfunction. In the 1960s, Neil Bricker introduced the “intact nephron hypothesis.” According to his concept, with decreases in the number of functioning nephrons, each remaining nephron has to adapt to carry a larger burden of transport, synthetic function, and regulatory function.

POTASSIUM Under normal and abnormal conditions, most of the filtered potassium is reabsorbed in the proximal tubule so that excretion is determined by secretion by the distal nephron. Potassium handling is altered in CKD protecting the organism somewhat from lethal hyperkalemia. Hyperkalemia is a common feature of individuals with CKD. Hyperkalemia (if not severe and dangerous) is adaptive in that it promotes potassium secretion by the principal cells of the collecting duct. When patients with CKD are given a potassium load, they can excrete it at the same rate as patients with normal renal function except that they do so at a higher serum potassium, consistent with the view that the hyperkalemia facilitates potassium excretion. The direct effect of hyperkalemia on potassium secretion by the distal nephron is independent of changes in aldosterone levels, but “normal” levels of aldosterone are necessary to see the effect of hyperkalemia on potassium excretion. Elevated potassium stimulates the production of aldosterone, and this effect is also seen in patients with CKD. Aldosterone increases the density and activity of the basolateral $\text{Na}^+\text{-K}^+$ ATPase and

TABLE 333e-2 FACTORS IMPLICATED IN COMPENSATORY RENAL GROWTH AFTER NEPHRON LOSS

Increased renal blood flow
Increased tubular absorption of Na with decreased distal delivery and decreased afferent arterial resistance due to adaptive tubuloglomerular feedback
Hepatocyte growth factor
Glucose transporters
Increased renal nerve activity
Insulin-like growth factor
Mammalian target of rapamycin (mTOR) signaling pathway activation
p21 ^{Waf1} , p27 ^{kip1} , and p57 ^{kip2}
Transforming growth factor β

the number of Na⁺ channels in the apical membrane of the collecting duct. In CKD, the excretion of the dietary load of potassium occurs at the expense of an elevation in serum potassium concentrations.

SODIUM As renal function is reduced with CKD, there is a reduced ability to excrete sodium. Thus, patients with advanced kidney disease are often fluid overloaded. In early disease, however, there are functional adaptations that the kidney assumes to help to maintain the *milieu intérieur*. With loss of functional nephrons, the remaining nephrons are hyperperfused and are hyperfiltering in a manner that can be influenced by dietary protein intake. Although protein restriction can decrease this compensatory hyperperfusion, there is generally more sodium and water filtered and delivered to the remaining nephrons. There is some preservation of glomerulotubular balance with increased proximal tubule sodium and water reabsorption associated with increased levels of the Na/H exchanger in apical membranes of the tubule. The tubuloglomerular feedback (TGF) of the remaining nephrons is sensitive to sodium intake. With high sodium intake in normal renal function, a negative feedback process occurs by which increased distal delivery results in reduced GFR and hence filtration of sodium. In CKD, the TGF becomes a positive feedback process by which increased distal delivery results in increased filtration so that the need to excrete an increased amount of sodium per nephron is achieved. This conversion from a negative feedback process to a positive feedback process may be due to conversion of an adenosine-dominated vasoconstrictive feedback on the afferent arteriole of the glomerulus to a NO-dominated vasodilatory feedback. Like so many of these adaptive responses, this one may turn maladaptive, resulting in higher intraglomerular hydrostatic pressures with increased mechanical strain on the glomerular capillary wall and podocytes and increased glomerulosclerosis as a consequence.

ACID-BASE HOMEOSTASIS The kidneys excrete approximately 1 mEq/kg per day of dietary acid load under normal dietary conditions. With decreased kidney functional mass, there is an adaptive response to increase H⁺ excretion by the remaining functional nephrons. This takes the form of enhanced nephron ammoniogenesis and increased distal nephron H⁺ ion secretion, which is mediated by the renin-angiotensin system and endothelin-1. NH₃ is produced by deamidation of glutamine in the proximal tubule. NH₃ is converted to NH₄⁺ in the collecting duct, where it buffers the secreted H⁺. It has been argued, however, that these mechanistic attempts to enhance H⁺ secretion can be maladaptive in that they can contribute to kidney inflammation and fibrosis and hence facilitate the progression of CKD.

MINERAL METABOLISM In CKD, there is a decrease in the ability of the kidney to excrete phosphate and produce 1,25-dihydroxyvitamin D₃ [1,25(OH)₂D₃]. There is a resultant increase in serum phosphate and reduction in serum calcium (Fig. 333e-2). In response, the body adapts by increasing production of parathyroid hormone (PTH) and fibroblast growth factor-23 (FGF-23) in an attempt to increase phosphaturia. The elevated levels of PTH act on bone to increase bone resorption and on osteocytes to increase FGF-23 expression. Elevated levels of PTH increase FGF-23 expression by activating protein kinase A and wnt signaling in osteoblast-like cells. There are a number of other factors that increase bone FGF-23 production in CKD including systemic acidosis, altered hydroxyapatite metabolism, changes in bone matrix, and release of low-molecular-weight FGFs. Although the production of PTH and FGF-23 initially are adaptive attempts to maintain body phosphate levels by enhancing excretion by the kidney, they become maladaptive due to systemic effects on the cardiovascular system and bone, as renal function continues to deteriorate. PTH and FGF-23 decrease the kidney's ability to reabsorb phosphate by decreasing the levels of the sodium-phosphate cotransporters NaPi2a and NaPi2c on the apical and basolateral membranes of the renal tubule. FGF-23 also reduces the ability of the kidney to generate 1,25(OH)₂D₃. In the parathyroid gland, the FGF-23 receptor, the klotho-fibroblast growth factor 1 complex, is downregulated with a consequent loss of the normal action of FGF-23 to downregulate PTH production. PTH and FGF-23 have been implicated in the cardiovascular disease that is so characteristic of patients with CKD. With CKD, there is less klotho expression in

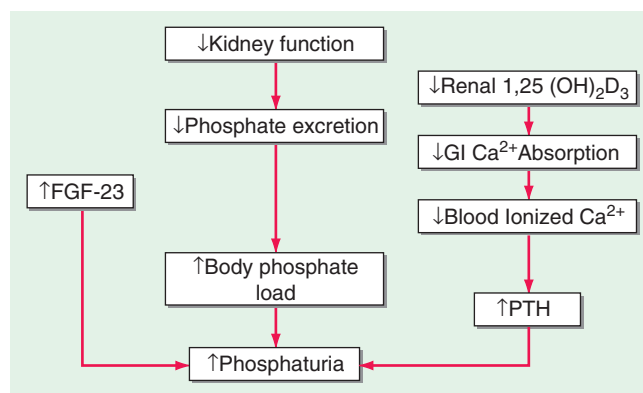


FIGURE 333e-2 Modification of the trade-off hypothesis of Slatopolsky and Bricker as it relates to the adaptation of the body to decreased functional renal mass in an attempt to maintain calcium and phosphate stores and serum levels. 1,25(OH)₂D₃, 1,25-dihydroxyvitamin D₃; FGF-23, fibroblast growth factor-23; GI, gastrointestinal; PTH, parathyroid hormone.

the kidney and the parathyroid glands. Klotho deficiency contributes to soft tissue calcifications in CKD. FGF-23 has been associated with increased mortality in CKD and has been reported to be involved causally in the development of left ventricular hypertrophy. PTH also has been reported to directly affect rat myocardial cells, increasing calcium entry into the cells and contributing to death of the cells.

THE EFFECTS OF ACUTE KIDNEY INJURY ON SUSCEPTIBILITY TO SUBSEQUENT INJURY (PRECONDITIONING)

Preconditioning represents activation by the organism of intrinsic defense mechanisms to cope with pathologic conditions. Ischemic preconditioning is the phenomenon whereby a prior ischemic insult renders the organ resistant to a subsequent ischemic insult. Renal protection afforded by prior renal injury was described approximately 100 years ago, in 1912, by Suzuki, who noted that the kidney became resistant to uranium nephrotoxicity if the animal had previously been exposed to a sublethal dose of uranium. This resistance of the renal epithelium to recurrent toxic injury was proposed to be a defense mechanism of the kidney. There have been a number of studies over the years demonstrating that preconditioning with a number of renal toxicants leads to protection against injury associated with a second exposure to the same toxicant or to another nephrotoxicant. It is not, however, a universal finding that toxins confer resistance to subsequent insults.

Kidney ischemic preconditioning is the conveyance of protection against ischemia due to prior exposure of the kidney to sublethal episodes of ischemia. In some experiments in rodents, these prior exposures were short (e.g., 5 min) and repeated or longer. Subsequent protection was generally found at 1–2 h or up to 48 h, but there has been a report of protection in the mouse for up to 12 weeks after the preconditioning exposures. Unilateral ischemia, with the contralateral kidney left alone, was also protective against a subsequent ischemic insult to the postischemic kidney, revealing that systemic uremia was not necessary for protection.

Remote Ischemic Preconditioning Remote ischemic preconditioning is a therapeutic strategy by which protection can be afforded in one vascular bed by ischemia to another vascular bed in the same organ or a different organ. A large number of studies have demonstrated that ischemia to one organ protects against ischemia to another. There are very few mechanistic studies of remote preconditioning in the kidney. In one study, naloxone blocked preconditioning in the kidney, implicating opiates as effectors. Remote preconditioning induced by ischemia to the muscle of the arm induced by a blood pressure cuff can result in protection of the kidney against a subsequent insult, such as one related to contrast agents in humans. Some of the cellular processes and signaling mechanisms proposed to explain preconditioning in the kidney and other organs are listed in Table 333e-3. These protective

TABLE 333e-3 FACTORS AND PROCESSES IMPLICATED AS PROTECTIVE MEDIATORS OF ISCHEMIC PRECONDITIONING

Adenosine
AKT (protein kinase B)
Antioxidants
Autophagy
Bradykinin
Decrease in genes regulating inflammation (cytokine synthesis, leukocyte chemotaxis, adhesion, exocytosis, innate immune signaling pathways)
Extracellular signal-related kinase (ERK)
Heat shock proteins
Hypoxia-inducible factors (HIFs)
JAK-STAT pathway
Jun N-terminal kinase (JNK)
Mitochondrial ATP-sensitive potassium channel (K ⁺ ATP channel)
Mitochondrial connexin 43
Nitric oxide
Opioids
Protein kinase C (PKC)
Sirtuin activity (SIRT1)

processes, most of which have been identified in the heart, involve multiple signaling pathways that affect decreased apoptosis, inhibition of mitochondrial permeability transition pores, activation of survival pathways, autophagy, and other pathways involved in reducing energy consumption or reactive oxygen production. In a study from our laboratory, inducible NO synthase was found to be an important contributor to the adaptive response to kidney injury, which results in protection against a subsequent insult. Identification of the responsible protective factor(s) mediating the advantageous adaptive response to remote ischemic preconditioning would provide a therapeutic approach for prevention of acute kidney injury or facilitation of a protective adaptation to kidney injury.

ADAPTIVE RESPONSE OF THE KIDNEY TO ACUTE INJURY

Adaptive Response to Hypoxic Injury Hypoxia plays a role in ischemic, septic, and toxic acute kidney injury. Many conditions result in a global or regional impairment of oxygen delivery. This is particularly important in the outer medulla where there is baseline reduced oxygen tension and a complex capillary network that, by its nature, is susceptible to interruption. In addition, the S3 segment of the proximal tubule is very dependent on oxidative metabolism, whereas the medullary thick ascending limb of the nephron that also traverses the outer medulla can adapt to hypoxia by converting to glycolysis as a primary energy source.

One proposed adaptive response to hypoxia is a reduction in glomerular filtration with consequent reduction in “work” requirement for reabsorption of solutes by the tubule. This was termed *acute renal success* by Thureau many years ago. The importance of this has been questioned, however, because there is no significant reduction in renal oxygen consumption in post–cardiac surgery patients with acute kidney injury in the setting of reduced GFR and renal blood flow.

If hypoxia or other influences, such as toxins, damage the proximal tubule and interfere with reabsorption of sodium and water, it is important that the kidney adapt in such a way so that there is not a large natriuresis that might compromise intravascular volume and blood pressure. This is accomplished, at least in part, by tubuloglomerular feedback (TGF). The increased distal delivery of salt and water results in a homeostatic adaptation to decrease glomerular filtration and hence decrease tubular delivery of salt and water through the glomerulus and reduce the delivery to the distal nephron. This adaptive response to acute injury is different from the role of TGF in CKD, as we have discussed previously in this chapter. In chronic disease with reduced nephron function, there is a steady-state need to increase excretion of sodium, whereas with acute injury, excretion of sodium is reduced.

Many genes are activated by hypoxia that are adaptive in serving to protect the cell and organ. With hypoxia, hypoxia-inducible factor (HIF) 1 α rapidly accumulates due to the inhibition of the HIF prolyl-hydroxylases, which normally promote HIF1 α proteasomal degradation. HIF1 α then dimerizes with HIF1 β and the dimer moves to the nucleus, where it upregulates a number of genes whose protein products are involved in energy metabolism, angiogenesis, and apoptosis, enhancing oxygen delivery and metabolic adaptation to hypoxia. This takes the form of a complex interplay among factors that regulate perfusion, cellular redox state, and mitochondrial function. For example, upregulation of NO production by sepsis results in vasodilatation and reduction in mitochondrial respiration and oxygen consumption. In addition, HIF1 activation in endothelial cells may be important for adaptive preservation of the microvasculature during and after hypoxia. Better understanding of the role that the HIFs play in protective adaptation has led to an aggressive development of HIF prolyl-hydroxylase inhibitors by biotechnology and pharmaceutical companies for clinical use.

Adaptive Response to Toxic Injury Specific to the Proximal Tubule One can model an acute kidney injury by genetically inserting a Simian diphtheria toxin (DT) receptor into the proximal tubule and then adding either a single dose of DT or multiple doses of the toxin. Repair of the kidney after a single dose of DT can be shown to be adaptive with few longer term sequelae. There is a very robust proliferative response of the proximal tubule cells to replace the cells that die as a result of the DT. Ultimately the inflammation resolves, and there is little, if any, residual interstitial inflammation, expansion, or matrix deposition.

Maladaptive Response of the Kidney to Acute Injury By contrast to the above adaptive repair that occurs after a single insult, after three doses of DT administered at weekly intervals, there is maladaptive repair with development over time of a chronic interstitial infiltrate, increased myofibroblast proliferation, tubulointerstitial fibrosis, and tubular atrophy, as well as an increase in serum creatinine (0.6 ± 0.1 mg/dL vs 0.18 ± 0.02 mg/dL in control mice) by week 5, 2 weeks after the last dose in the thrice-treated animals. There is a dramatic increase in the number of interstitial cells that expressed the platelet-derived growth factor receptor β (pericytes/perivascular fibroblasts), α SMA (myofibroblasts), FSP-1/S100A4 (fibroblast specific protein-1), and F4/80 (macrophages). In addition, there is loss of endothelial cells, interstitial capillaries, and development of focal global and segmental glomerulosclerosis.

It has become increasingly recognized as a result of large epidemiologic studies that even mild forms of acute kidney injury are associated with adverse short- and long-term outcomes including onset or progression of CKD and more rapid progression to end-stage kidney disease. Experimental models in animals, such as the DT model described above, provide pathophysiologic explanations for how the effects of acute injury can lead to chronic inflammation, vascular rarefaction, tubular cell atrophy, interstitial fibrosis, and glomerulosclerosis. Recurrent specific tubular injury leads to a pattern very typical of CKD in humans: tubular atrophy, interstitial chronic inflammation and fibrosis, vascular rarefaction, and glomerulosclerosis. The mechanisms involved in the development of glomerulosclerosis evoked by primary tubular injury may be multifactorial. Damage to nephron segments may lead to sluffing of cells into the lumen and to tubular obstruction. Progressive narrowing of the early proximal tubule near the glomerular tuft can lead to a sclerotic atubular glomerulus like those that are seen with ureteral obstruction. There may be paracrine signaling from injured and regenerating/undifferentiated epithelium to directly impact the glomerulus. Alternatively, a progressive tubulointerstitial reaction originating around atrophic and undifferentiated tubules may directly encroach upon the glomerular tuft. The loss of interstitial capillaries may lead to a progressive reduction of glomerular blood flow with ischemia to the glomerulus and to the kidney regions perfused by the postglomerular capillaries. This speaks to the fact that primary tubular injury can trigger a response that adversely affects multiple compartments of the kidney and leads to a positive feedback process, involving loss of capillaries, glomerulosclerosis, persistent ischemia, tubular atrophy, increased fibrosis, and ultimately kidney failure.

334 Acute Kidney Injury


Sushrut S. Waikar, Joseph V. Bonventre

Acute kidney injury (AKI), previously known as acute renal failure, is characterized by the sudden impairment of kidney function resulting in the retention of nitrogenous and other waste products normally cleared by the kidneys. AKI is not a single disease but, rather, a designation for a heterogeneous group of conditions that share common diagnostic features: specifically, an increase in the blood urea nitrogen (BUN) concentration and/or an increase in the plasma or serum creatinine (SCr) concentration, often associated with a reduction in urine volume. It is important to recognize that AKI is a clinical diagnosis and not a structural one. A patient may have AKI without injury to the kidney parenchyma. AKI can range in severity from asymptomatic and transient changes in laboratory parameters of glomerular filtration rate (GFR), to overwhelming and rapidly fatal derangements in effective circulating volume regulation and electrolyte and acid-base composition of the plasma.

EPIDEMIOLOGY

AKI complicates 5–7% of acute care hospital admissions and up to 30% of admissions to the intensive care unit, particularly in the setting of diarrheal illnesses, infectious diseases like malaria and leptospirosis, and natural disasters such as earthquakes. The incidence of AKI has grown by more than fourfold in the United States since 1988 and is estimated to have a yearly incidence of 500 per 100,000 population, higher than the yearly incidence of stroke. AKI is associated with a markedly increased risk of death in hospitalized individuals, particularly in those admitted to the ICU where in-hospital mortality rates may exceed 50%. AKI increases the risk for the development or worsening of chronic kidney disease. Patients who survive and recover from an episode of severe AKI requiring dialysis are at increased risk for the later development of dialysis-requiring end-stage kidney disease. AKI may be community-acquired or hospital-acquired. Common causes of community-acquired AKI include volume depletion, adverse effects of medications, and obstruction of the urinary tract. The most common clinical settings for hospital-acquired AKI are sepsis, major surgical procedures, critical illness involving heart or liver failure, intravenous iodinated contrast administration, and nephrotoxic medication administration.

AKI IN THE DEVELOPING WORLD

 AKI is also a major medical complication in the developing world, where the epidemiology differs from that in developed countries due to differences in demographics, economics, geography, and comorbid disease burden. While certain features of AKI are common to both—particularly since urban centers of some developing countries increasingly resemble those in the developed world—many etiologies for AKI are region-specific such as envenomations from snakes, spiders, caterpillars, and bees; infectious causes such as malaria and leptospirosis; and crush injuries and resultant rhabdomyolysis from earthquakes.

ETIOLOGY AND PATHOPHYSIOLOGY

The causes of AKI have traditionally been divided into three broad categories: prerenal azotemia, intrinsic renal parenchymal disease, and postrenal obstruction ([Fig. 334-1](#)).

PRERENAL AZOTEMIA

Prerenal azotemia (from “azo,” meaning nitrogen, and “-emia”) is the most common form of AKI. It is the designation for a rise in SCr or BUN concentration due to inadequate renal plasma flow and intraglomerular hydrostatic pressure to support normal glomerular

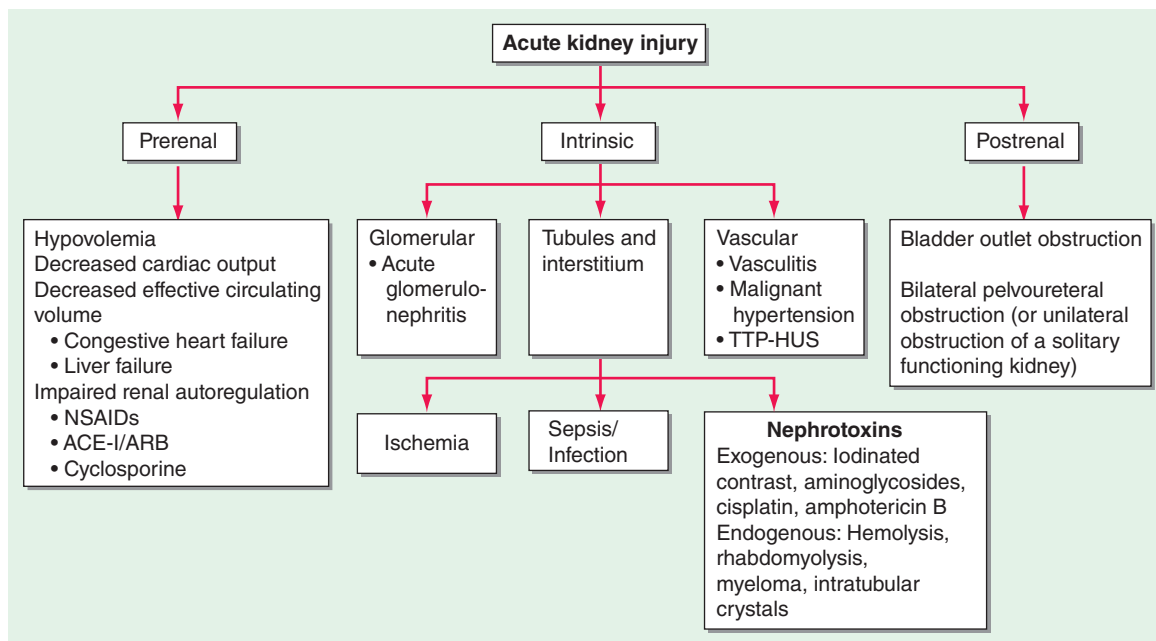


FIGURE 334-1 Classification of the major causes of acute kidney injury. ACE-I, angiotensin-converting enzyme inhibitor-I; ARB, angiotensin receptor blocker; NSAIDs, nonsteroidal anti-inflammatory drugs; TTP-HUS, thrombotic thrombocytopenic purpura–hemolytic-uremic syndrome.

filtration. The most common clinical conditions associated with prerenal azotemia are hypovolemia, decreased cardiac output, and medications that interfere with renal autoregulatory responses such as nonsteroidal anti-inflammatory drugs (NSAIDs) and inhibitors of angiotensin II (Fig. 334-2). Prerenal azotemia may coexist with other forms of intrinsic AKI associated with processes acting directly on the renal parenchyma. Prolonged periods of prerenal azotemia may lead to ischemic injury, often termed acute tubular necrosis (ATN). By definition, prerenal azotemia involves no parenchymal damage to the kidney and is rapidly reversible once intraglomerular hemodynamics are restored.

Normal GFR is maintained in part by the relative resistances of the afferent and efferent renal arterioles, which determine the glomerular plasma flow and the transcapillary hydraulic pressure gradient that drive glomerular ultrafiltration. Mild degrees of hypovolemia and reductions in cardiac output elicit compensatory renal physiologic changes. Because renal blood flow accounts for 20% of the cardiac output, renal vasoconstriction and salt and water reabsorption occur as homeostatic responses to decreased effective circulating volume or cardiac output in order to maintain blood pressure and increase intravascular volume to sustain perfusion to the cerebral and coronary vessels. Mediators of this response include angiotensin II, norepinephrine, and vasopressin (also termed antidiuretic hormone). Glomerular filtration can be maintained despite reduced renal blood flow by angiotensin II–mediated renal efferent vasoconstriction, which maintains glomerular capillary hydrostatic pressure closer to normal and thereby prevents marked reductions in GFR if renal blood flow reduction is not excessive.

In addition, a myogenic reflex within the afferent arteriole leads to dilation in the setting of low perfusion pressure, thereby maintaining glomerular perfusion. Intrarenal biosynthesis of vasodilator prostaglandins (prostacyclin, prostaglandin E_2), kallikrein and kinins, and possibly nitric oxide (NO) also increase in response to low renal perfusion pressure. Autoregulation is also accomplished by tubuloglomerular feedback, in which decreases in solute delivery to the macula densa (specialized cells within the distal tubule) elicit dilation of the juxta-posed afferent arteriole in order to maintain glomerular perfusion, a mechanism mediated, in part, by NO. There is a limit, however, to the ability of these counterregulatory mechanisms to maintain GFR in the face of systemic hypotension. Even in healthy adults, renal autoregulation usually fails once the systolic blood pressure falls below 80 mmHg.

A number of factors determine the robustness of the autoregulatory response and the risk of prerenal azotemia. Atherosclerosis,

long-standing hypertension, and older age can lead to hyalinosis and myointimal hyperplasia, causing structural narrowing of the intrarenal arterioles and impaired capacity for renal afferent vasodilation. In chronic kidney disease, renal afferent vasodilation may be operating at maximal capacity in order to maximize GFR in response to reduced functional renal mass. Drugs can affect the compensatory changes evoked to maintain GFR. NSAIDs inhibit renal prostaglandin production, limiting renal afferent vasodilation. Angiotensin-converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) limit renal efferent vasoconstriction; this effect is particularly pronounced in patients with bilateral renal artery stenosis or unilateral renal artery stenosis (in the case of a solitary functioning kidney) because renal efferent vasoconstriction is needed to maintain GFR due to low renal perfusion. The combined use of NSAIDs with ACE inhibitors or ARBs poses a particularly high risk for developing prerenal azotemia.

Many individuals with advanced cirrhosis exhibit a unique hemodynamic profile that resembles prerenal azotemia despite total-body volume overload. Systemic vascular resistance is markedly reduced due to primary arterial vasodilation in the splanchnic circulation, resulting ultimately in activation of vasoconstrictor responses similar to those seen in hypovolemia. AKI is a common complication in this setting, and it can be triggered by volume depletion and spontaneous bacterial peritonitis. A particularly poor prognosis is seen in the case of type 1 hepatorenal syndrome, in which AKI without an alternate cause (e.g., shock and nephrotoxic drugs) persists despite volume administration and withholding of diuretics. Type 2 hepatorenal syndrome is a less severe form characterized mainly by refractory ascites.

INTRINSIC AKI

The most common causes of intrinsic AKI are sepsis, ischemia, and nephrotoxins, both endogenous and exogenous (Fig. 334-3). In many cases, prerenal azotemia advances to tubular injury. Although classically termed “acute tubular necrosis,” human biopsy confirmation of tubular necrosis is, in general, often lacking in cases of sepsis and ischemia; indeed, processes such as inflammation, apoptosis, and altered regional perfusion may be important contributors pathophysiologically. Other causes of intrinsic AKI are less common and can be conceptualized anatomically according to the major site of renal parenchymal damage: glomeruli, tubulointerstitium, and vessels.

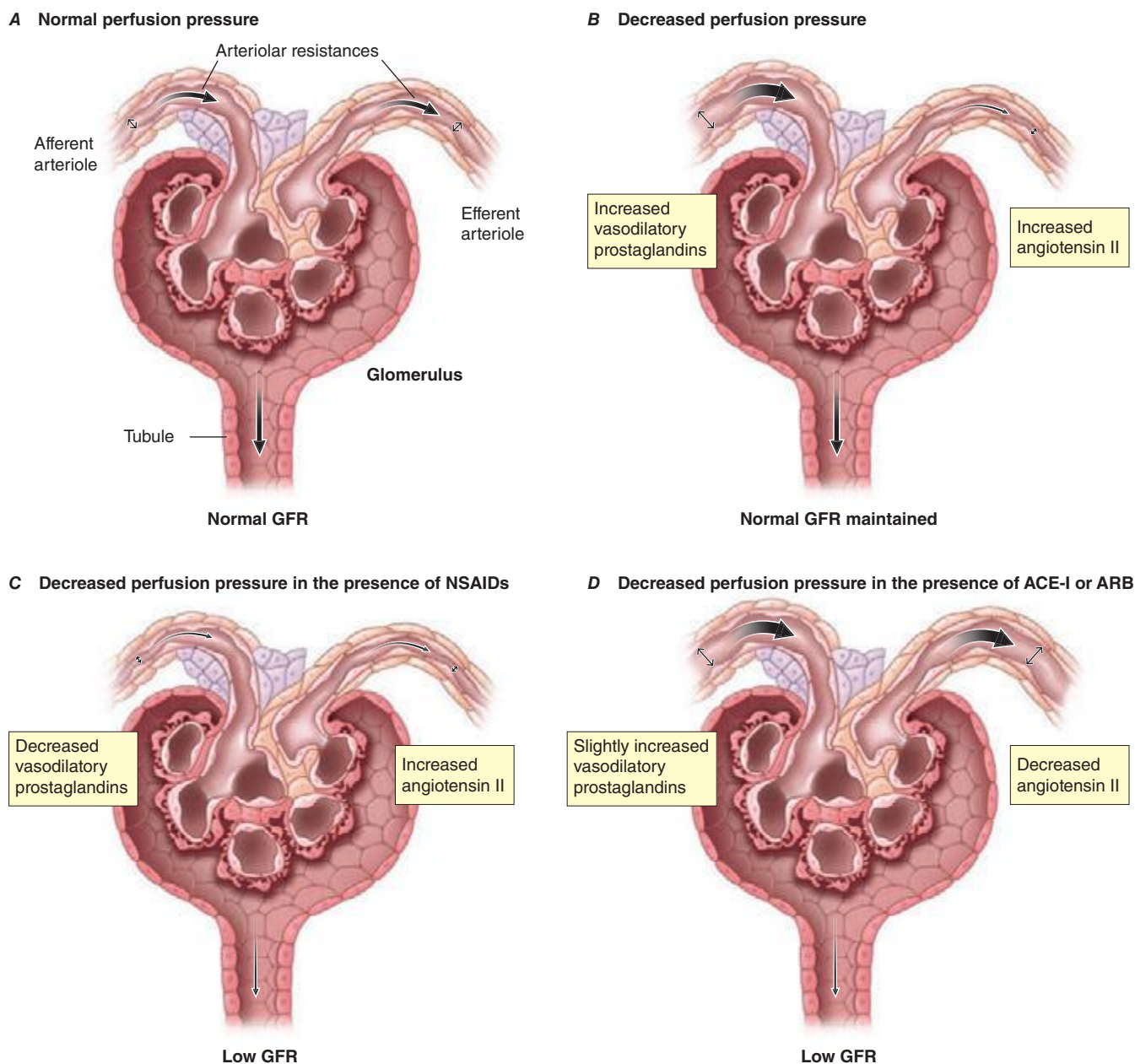


FIGURE 334-2 Intrarenal mechanisms for autoregulation of the glomerular filtration rate (GFR) under decreased perfusion pressure and reduction of the GFR by drugs. **A.** Normal conditions and a normal GFR. **B.** Reduced perfusion pressure within the autoregulatory range. Normal glomerular capillary pressure is maintained by afferent vasodilatation and efferent vasoconstriction. **C.** Reduced perfusion pressure with a nonsteroidal anti-inflammatory drug (NSAID). Loss of vasodilatory prostaglandins increases afferent resistance; this causes the glomerular capillary pressure to drop below normal values and the GFR to decrease. **D.** Reduced perfusion pressure with an angiotensin-converting enzyme inhibitor (ACE-I) or an angiotensin receptor blocker (ARB). Loss of angiotensin II action reduces efferent resistance; this causes the glomerular capillary pressure to drop below normal values and the GFR to decrease. (From JG Abuelo: *N Engl J Med* 357:797-805, 2007; with permission.)

SEPSIS-ASSOCIATED AKI

In the United States, more than 700,000 cases of sepsis occur each year. AKI complicates more than 50% of cases of severe sepsis and greatly increases the risk of death. Sepsis is also a very important cause of AKI in the developing world. Decreases in GFR with sepsis can occur even in the absence of overt hypotension, although most cases of severe AKI typically occur in the setting of hemodynamic collapse requiring vasopressor support. While there is clearly tubular injury associated with AKI in sepsis as manifest by the presence of tubular debris and casts in the urine, postmortem examinations of kidneys from individuals with severe sepsis suggest that other factors, perhaps related to inflammation, mitochondrial dysfunction, and interstitial edema, must be considered in the pathophysiology of sepsis-induced AKI.

The hemodynamic effects of sepsis—arising from generalized arterial vasodilation, mediated in part by cytokines that upregulate the

expression of inducible NO synthase in the vasculature—can lead to a reduction in GFR. The operative mechanisms may be excessive efferent arteriole vasodilation, particularly early in the course of sepsis, or renal vasoconstriction from activation of the sympathetic nervous system, the renin-angiotensin-aldosterone system, vasopressin, and endothelin. Sepsis may lead to endothelial damage, which results in microvascular thrombosis, activation of reactive oxygen species, and leukocyte adhesion and migration, all of which may injure renal tubular cells.

ISCHEMIA-ASSOCIATED AKI

Healthy kidneys receive 20% of the cardiac output and account for 10% of resting oxygen consumption, despite constituting only 0.5% of the human body mass. The kidneys are also the site of one of the most hypoxic regions in the body, the renal medulla. The outer medulla is particularly vulnerable to ischemic damage because of the

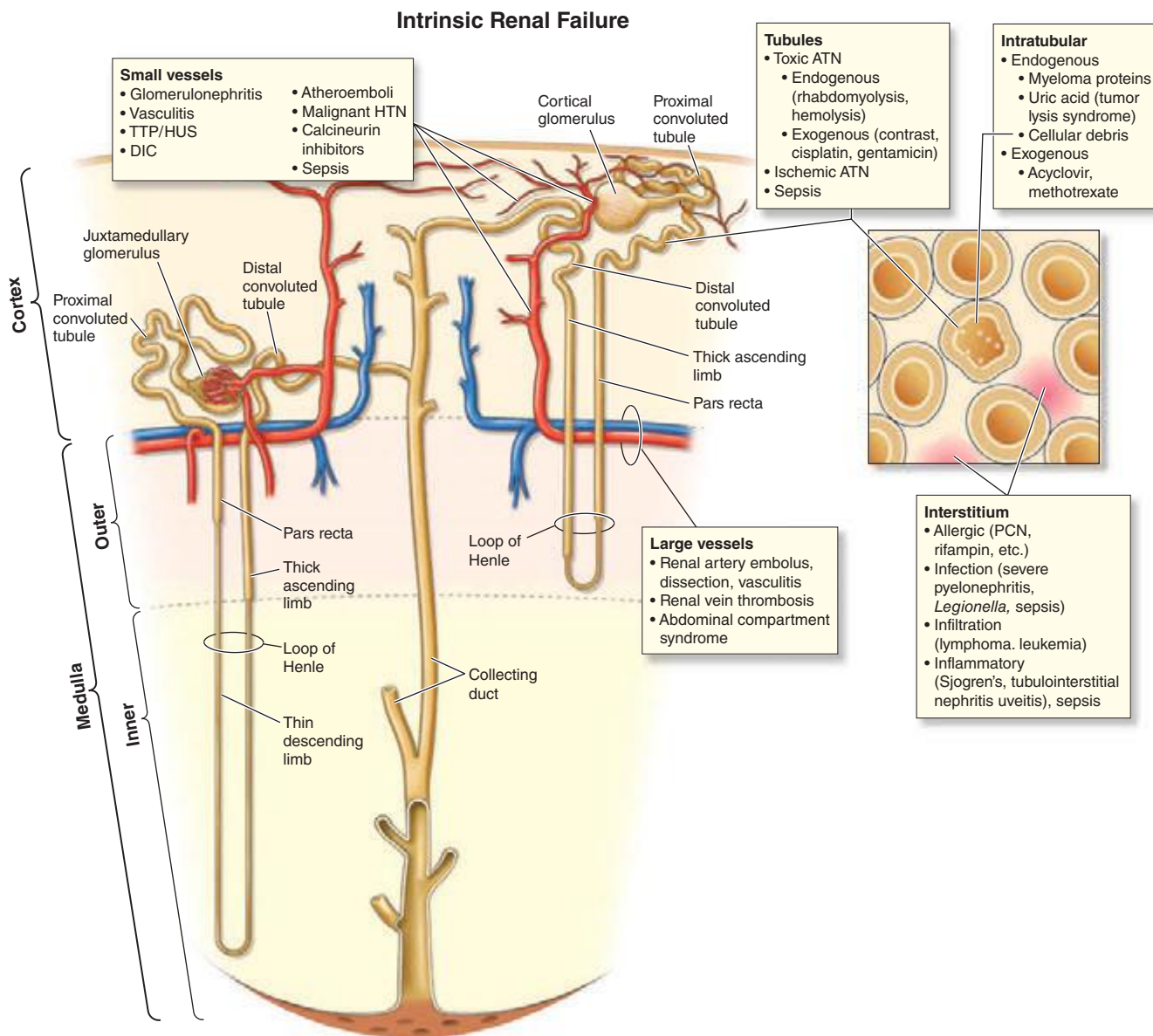


FIGURE 334-3 Major causes of intrinsic acute kidney injury. ATN, acute tubular necrosis; DIC, disseminated intravascular coagulation; HTN, hypertension; PCN, penicillin; TTP/HUS, thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome; TINU, tubulointerstitial nephritis-uveitis.

architecture of the blood vessels that supply oxygen and nutrients to the tubules. Enhanced leukocyte-endothelial interactions in the small vessels lead to inflammation and reduced local blood flow to the metabolically very active S3 segment of the proximal tubule, which depends on oxidative metabolism for survival. Ischemia alone in a normal kidney is usually not sufficient to cause severe AKI, as evidenced by the relatively low risk of severe AKI even after total interruption of renal blood flow during suprarenal aortic clamping or cardiac arrest. Clinically, AKI more commonly develops when ischemia occurs in the context of limited renal reserve (e.g., chronic kidney disease or older age) or coexisting insults such as sepsis, vasoactive or nephrotoxic drugs, rhabdomyolysis, or the systemic inflammatory states associated with burns and pancreatitis. Prerenal azotemia and ischemia-associated AKI represent a continuum of the manifestations of renal hypoperfusion. Persistent prerenal vasoconstriction may be a common underlying cause of the reduction in GFR seen in AKI; implicated factors for vasoconstriction include activation of tubuloglomerular feedback from enhanced delivery of solute to the macula densa following proximal tubule injury, increased basal vascular tone and reactivity to vasoconstrictive agents, and decreased vasodilator responsiveness. Other contributors to low GFR include backleak of filtrate across damaged and

denuded tubular epithelium and mechanical obstruction of tubules from necrotic debris (Fig. 334-4).

Postoperative AKI Ischemia-associated AKI is a serious complication in the postoperative period, especially after major operations involving significant blood loss and intraoperative hypotension. The procedures most commonly associated with AKI are cardiac surgery with cardiopulmonary bypass (particularly for combined valve and bypass procedures), vascular procedures with aortic cross clamping, and intraperitoneal procedures. Severe AKI requiring dialysis occurs in approximately 1% of cardiac and vascular surgery procedures. The risk of severe AKI has been less well studied for major intraperitoneal procedures but appears to be of comparable magnitude. Common risk factors for postoperative AKI include underlying chronic kidney disease, older age, diabetes mellitus, congestive heart failure, and emergency procedures. The pathophysiology of AKI following cardiac surgery is multifactorial. Major AKI risk factors are common in the population undergoing cardiac surgery. The use of nephrotoxic agents including iodinated contrast for cardiac imaging prior to surgery may increase the risk of AKI. Cardiopulmonary bypass is a unique hemodynamic state characterized by nonpulsatile flow and exposure of the circulation to extracorporeal circuits. Longer duration of cardiopulmonary bypass is a risk factor for AKI. In addition to ischemic

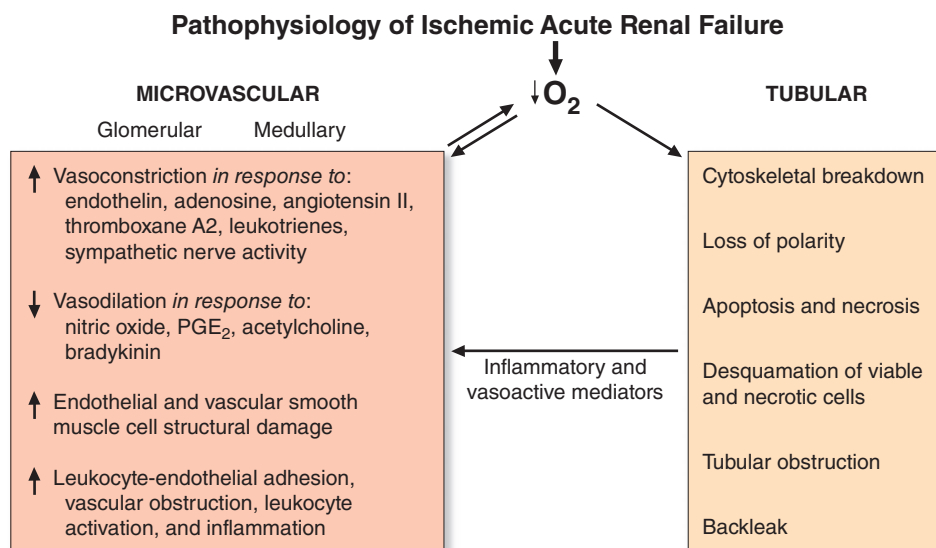


FIGURE 334-4 Interacting microvascular and tubular events contributing to the pathophysiology of ischemic acute kidney injury.

PGE₂, prostaglandin E₂. (From JV Bonventre, JM Weinberg: *J Am Soc Nephrol* 14:2199, 2003.)

injury from sustained hypoperfusion, cardiopulmonary bypass may cause AKI through a number of mechanisms including extracorporeal circuit activation of leukocytes and inflammatory processes, hemolysis with resultant pigment nephropathy (see below), and aortic injury with resultant atheroemboli. AKI from atheroembolic disease, which can also occur following percutaneous catheterization of the aorta, or spontaneously, is due to cholesterol crystal embolization resulting in partial or total occlusion of multiple small arteries within the kidney. Over time, a foreign body reaction can result in intimal proliferation, giant cell formation, and further narrowing of the vascular lumen, accounting for the generally subacute (over a period of weeks rather than days) decline in renal function.

Burns and Acute Pancreatitis Extensive fluid losses into the extravascular compartments of the body frequently accompany severe burns and acute pancreatitis. AKI is an ominous complication of burns, affecting 25% of individuals with more than 10% total body surface area involvement. In addition to severe hypovolemia resulting in decreased cardiac output and increased neurohormonal activation, burns and acute pancreatitis both lead to dysregulated inflammation and an increased risk of sepsis and acute lung injury, all of which may facilitate the development and progression of AKI. Individuals undergoing massive fluid resuscitation for trauma, burns, and acute pancreatitis can also develop the abdominal compartment syndrome, where markedly elevated intraabdominal pressures, usually higher than 20 mmHg, lead to renal vein compression and reduced GFR.

Diseases of the Microvasculature Leading to Ischemia Microvascular causes of AKI include the thrombotic microangiopathies (antiphospholipid antibody syndrome, radiation nephritis, malignant nephrosclerosis, and thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome [TTP-HUS]), scleroderma, and atheroembolic disease. Large-vessel diseases associated with AKI include renal artery dissection, thromboembolism, thrombosis, and renal vein compression or thrombosis.

NEPHROTOXIN-ASSOCIATED AKI

The kidney has very high susceptibility to nephrotoxicity due to extremely high blood perfusion and concentration of circulating substances along the nephron where water is reabsorbed and in the medullary interstitium; this results in high-concentration exposure of toxins to tubular, interstitial, and endothelial cells. Nephrotoxic injury occurs in response to a number of pharmacologic compounds with diverse structures, endogenous substances, and environmental exposures. All structures of the kidney are vulnerable to toxic injury, including the tubules, interstitium, vasculature, and collecting system. As with other forms of AKI, risk factors for nephrotoxicity include older age, chronic kidney disease (CKD), and prerenal azotemia. Hypoalbuminemia may

increase the risk of some forms of nephrotoxin-associated AKI due to increased free circulating drug concentrations.

Contrast Agents Iodinated contrast agents used for cardiovascular and computed tomography (CT) imaging are a leading cause of AKI. The risk of AKI, or “contrast nephropathy,” is negligible in those with normal renal function but increases markedly in the setting of CKD, particularly diabetic nephropathy. The most common clinical course of contrast nephropathy is characterized by a rise in SCr beginning 24–48 h following exposure, peaking within 3–5 days, and resolving within 1 week. More severe, dialysis-requiring AKI is uncommon except in the setting of significant preexisting CKD, often in association with congestive heart failure or other coexisting causes for ischemia-associated AKI. Patients with multiple myeloma and renal disease are particularly susceptible. Low fractional excretion of sodium and relatively benign urinary sediment without features of tubular necrosis (see below) are common findings. Contrast nephropathy is thought to occur from a combination of factors, including (1) hypoxia in the renal outer medulla due to perturbations in renal microcirculation and occlusion of small vessels; (2) cytotoxic damage to the tubules directly or via the generation of oxygen free radicals, especially because the concentration of the agent within the tubule is markedly increased; and (3) transient tubule obstruction with precipitated contrast material. Other diagnostic agents implicated as a cause of AKI are high-dose gadolinium used for magnetic resonance imaging (MRI) and oral sodium phosphate solutions used as bowel purgatives.

Antibiotics Several antimicrobial agents are commonly associated with AKI. *Aminoglycosides* and *amphotericin B* both cause tubular necrosis. Nonoliguric AKI (i.e., without a significant reduction in urine volume) accompanies 10–30% of courses of aminoglycoside antibiotics, even when plasma levels are in the therapeutic range. Aminoglycosides are freely filtered across the glomerulus and then accumulate within the renal cortex, where concentrations can greatly exceed those of the plasma. AKI typically manifests after 5–7 days of therapy and can present even after the drug has been discontinued. Hypomagnesemia is a common finding.

Amphotericin B causes renal vasoconstriction from an increase in tubuloglomerular feedback as well as direct tubular toxicity mediated by reactive oxygen species. Nephrotoxicity from amphotericin B is dose and duration dependent. This drug binds to tubular membrane cholesterol and introduces pores. Clinical features of amphotericin B nephrotoxicity include polyuria, hypomagnesemia, hypocalcemia, and nongap metabolic acidosis.

Vancomycin may be associated with AKI, particularly when trough levels are high, but a causal relationship with AKI has not been definitively

established. *Acyclovir* can precipitate in tubules and cause AKI by tubular obstruction, particularly when given as an intravenous bolus at high doses (500 mg/m²) or in the setting of hypovolemia. *Foscarnet*, *pentamidine*, *tenofovir*, and *cidofovir* are also frequently associated with AKI due to tubular toxicity. AKI secondary to acute interstitial nephritis can occur as a consequence of exposure to many antibiotics, including *penicillins*, *cephalosporins*, *quinolones*, *sulfonamides*, and *rifampin*.

Chemotherapeutic Agents *Cisplatin* and *carboplatin* are accumulated by proximal tubular cells and cause necrosis and apoptosis. Intensive hydration regimens have reduced the incidence of cisplatin nephrotoxicity, but it remains a dose-limiting toxicity. *Ifosfamide* may cause hemorrhagic cystitis and tubular toxicity, manifested as type II renal tubular acidosis (Fanconi's syndrome), polyuria, hypokalemia, and a modest decline in GFR. Antiangiogenesis agents, such as *bevacizumab*, can cause proteinuria and hypertension via injury to the glomerular microvasculature (thrombotic microangiopathy). Other antineoplastic agents such as mitomycin C and gemcitabine may cause thrombotic microangiopathy with resultant AKI.

Toxic Ingestions Ethylene glycol, present in automobile antifreeze, is metabolized to oxalic acid, glycolaldehyde, and glyoxylate, which may cause AKI through direct tubular injury. Diethylene glycol is an industrial agent that has been the cause of outbreaks of severe AKI around the world due to adulteration of pharmaceutical preparations. The metabolite 2-hydroxyethoxyacetic acid (HEAA) is thought to be responsible for tubular injury. Melamine contamination of foodstuffs has led to nephrolithiasis and AKI, either through intratubular obstruction or possibly direct tubular toxicity. Aristolochic acid was found to be the cause of "Chinese herb nephropathy" and "Balkan nephropathy" due to contamination of medicinal herbs or farming. The list of environmental toxins is likely to grow and contribute to a better understanding of previously catalogued "idiopathic" chronic tubular interstitial disease, a common diagnosis in both the developed and developing world.

Endogenous Toxins AKI may be caused by a number of endogenous compounds, including myoglobin, hemoglobin, uric acid, and myeloma light chains. Myoglobin can be released by injured muscle cells, and hemoglobin can be released during massive hemolysis leading to pigment nephropathy. Rhabdomyolysis may result from traumatic crush injuries, muscle ischemia during vascular or orthopedic surgery, compression

during coma or immobilization, prolonged seizure activity, excessive exercise, heat stroke or malignant hyperthermia, infections, metabolic disorders (e.g., hypophosphatemia, severe hypothyroidism), and myopathies (drug-induced, metabolic, or inflammatory). Pathogenic factors for AKI include intrarenal vasoconstriction, direct proximal tubular toxicity, and mechanical obstruction of the distal nephron lumen when myoglobin or hemoglobin precipitates with Tamm-Horsfall protein (uromodulin, the most common protein in urine and produced in the thick ascending limb of the loop of Henle), a process favored by acidic urine. Tumor lysis syndrome may follow initiation of cytotoxic therapy in patients with high-grade lymphomas and acute lymphoblastic leukemia; massive release of uric acid (with serum levels often exceeding 15 mg/dL) leads to precipitation of uric acid in the renal tubules and AKI ([Chap. 331](#)). Other features of tumor lysis syndrome include hyperkalemia and hyperphosphatemia. The tumor lysis syndrome can also occasionally occur spontaneously or with treatment for solid tumors or multiple myeloma. Myeloma light chains can also cause AKI by direct tubular toxicity and by binding to Tamm-Horsfall protein to form obstructing intratubular casts. Hypercalcemia, which can also be seen in multiple myeloma, may cause AKI by intense renal vasoconstriction and volume depletion.

Allergic Acute Tubulointerstitial Disease and Other Causes of Intrinsic AKI

While many of the ischemic and toxic causes of AKI previously described result in tubulointerstitial disease, many drugs are also associated with the development of an allergic response characterized by an inflammatory infiltrate and often peripheral and urinary eosinophilia. AKI may be caused by severe infections and infiltrative diseases. Diseases of the glomeruli or vasculature can lead to AKI by compromising blood flow within the renal circulation. Glomerulonephritis and vasculitis are less common causes of AKI. It is particularly important to recognize these diseases early because they require timely treatment with immunosuppressive agents or therapeutic plasma exchange.

POSTRENAL ACUTE KIDNEY INJURY

([See also Chap. 343](#)) Postrenal AKI occurs when the normally unidirectional flow of urine is acutely blocked either partially or totally, leading to increased retrograde hydrostatic pressure and interference with glomerular filtration. Obstruction to urinary flow may be caused by functional or structural derangements anywhere from the renal pelvis to the tip of the urethra ([Fig. 334-5](#)). Normal urinary flow rate

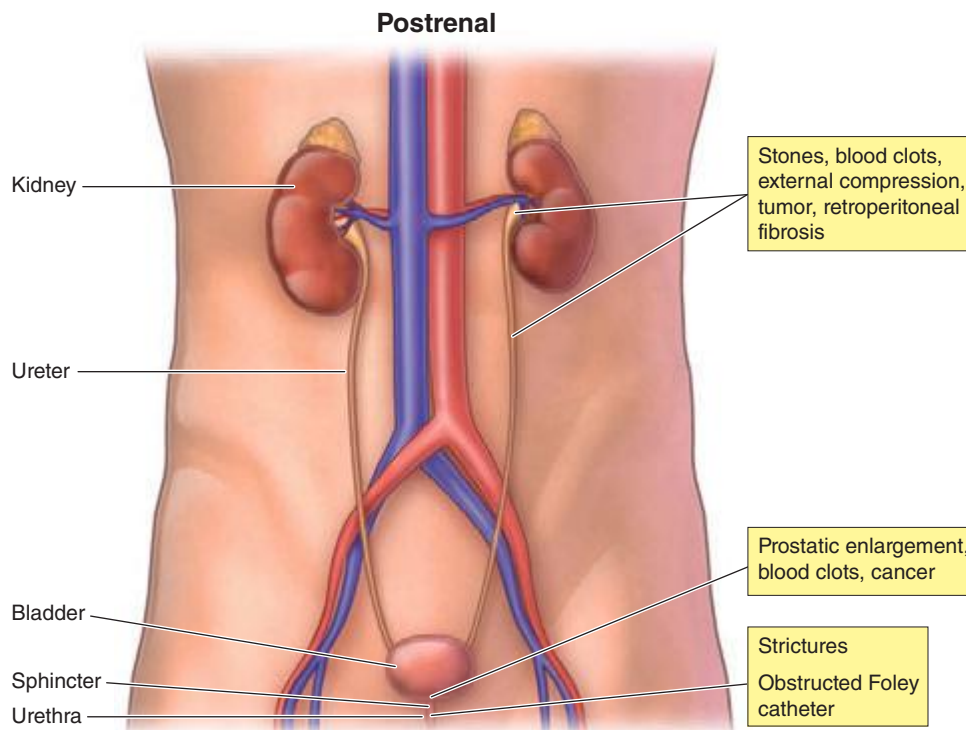


FIGURE 334-5 Anatomic sites and causes of obstruction leading to postrenal acute kidney injury.

does not rule out the presence of partial obstruction, because the GFR is normally two orders of magnitude higher than the urinary flow rate. For AKI to occur in healthy individuals, obstruction must affect both kidneys unless only one kidney is functional, in which case unilateral obstruction can cause AKI. Unilateral obstruction may cause AKI in the setting of significant underlying CKD or, in rare cases, from reflex vasospasm of the contralateral kidney. Bladder neck obstruction is a common cause of postrenal AKI and can be due to prostate disease (benign prostatic hypertrophy or prostate cancer), neurogenic bladder, or therapy with anticholinergic drugs. Obstructed Foley catheters can cause postrenal AKI if not recognized and relieved. Other causes of lower tract obstruction are blood clots, calculi, and urethral strictures. Ureteric obstruction can occur from intraluminal obstruction (e.g., calculi, blood clots, sloughed renal papillae), infiltration of the ureteric wall (e.g., neoplasia), or external compression (e.g., retroperitoneal fibrosis, neoplasia, abscess, or inadvertent surgical damage). The pathophysiology of postrenal AKI involves hemodynamic alterations triggered by an abrupt increase in intratubular pressures. An initial period of hyperemia from afferent arteriolar dilation is followed by intrarenal vasoconstriction from the generation of angiotensin II, thromboxane A₂, and vasopressin, and a reduction in NO production. Reduced GFR is due to underperfusion of glomeruli and, possibly, changes in the glomerular ultrafiltration coefficient.

DIAGNOSTIC EVALUATION (TABLE 334-1)

The presence of AKI is usually inferred by an elevation in the SCr concentration. AKI is currently defined by a rise from baseline of at least 0.3 mg/dL within 48 h or at least 50% higher than baseline within 1 week, or a reduction in urine output to less than 0.5 mL/kg per hour for longer than 6 h. It is important to recognize that given this definition, some patients with AKI will not have tubular or glomerular damage (e.g., prerenal azotemia). The distinction between AKI and CKD is important for proper diagnosis and treatment. The distinction is straightforward when a recent baseline SCr concentration is available, but more difficult in the many instances in which the baseline is unknown. In such cases, clues suggestive of CKD can come from radiologic studies (e.g., small, shrunken kidneys with cortical thinning on renal ultrasound, or evidence of renal osteodystrophy) or laboratory tests such as normocytic anemia in the absence of blood loss or secondary hyperparathyroidism with hyperphosphatemia and hypocalcemia, consistent with CKD. No set of tests, however, can rule out AKI superimposed on CKD because AKI is a frequent complication in patients with CKD, further complicating the distinction. Serial blood tests showing continued substantial rise of SCr represents clear evidence of AKI. Once the diagnosis of AKI is established, its cause needs to be determined.

HISTORY AND PHYSICAL EXAMINATION

The clinical context, careful history taking, and physical examination often narrow the differential diagnosis for the cause of AKI. Prerenal azotemia should be suspected in the setting of vomiting, diarrhea, glycosuria causing polyuria, and several medications including diuretics, NSAIDs, ACE inhibitors, and ARBs. Physical signs of orthostatic hypotension, tachycardia, reduced jugular venous pressure, decreased skin turgor, and dry mucous membranes are often present in prerenal azotemia. A history of prostatic disease, nephrolithiasis, or pelvic or paraaortic malignancy would suggest the possibility of postrenal AKI. Whether or not symptoms are present early during obstruction of the urinary tract depends on the location of obstruction. Colicky flank pain radiating to the groin suggests acute ureteric obstruction. Nocturia and urinary frequency or hesitancy can be seen in prostatic disease. Abdominal fullness and suprapubic pain can accompany massive bladder enlargement. Definitive diagnosis of obstruction requires radiologic investigations.

A careful review of all medications is imperative in the evaluation of an individual with AKI. Not only are medications frequently a cause of AKI, but doses of administered medications must be adjusted for estimated GFR. Idiosyncratic reactions to a wide variety of medications

can lead to allergic interstitial nephritis, which may be accompanied by fever, arthralgias, and a pruritic erythematous rash. The absence of systemic features of hypersensitivity, however, does not exclude the diagnosis of interstitial nephritis.

AKI accompanied by palpable purpura, pulmonary hemorrhage, or sinusitis raises the possibility of systemic vasculitis with glomerulonephritis. Atheroembolic disease can be associated with livedo reticularis and other signs of emboli to the legs. A tense abdomen should prompt consideration of acute abdominal compartment syndrome, which requires measurement of bladder pressure. Signs of limb ischemia may be clues to the diagnosis of rhabdomyolysis.

URINE FINDINGS

Complete anuria early in the course of AKI is uncommon except in the following situations: complete urinary tract obstruction, renal artery occlusion, overwhelming septic shock, severe ischemia (often with cortical necrosis), or severe proliferative glomerulonephritis or vasculitis. A reduction in urine output (oliguria, defined as <400 mL/24 h) usually denotes more severe AKI (i.e., lower GFR) than when urine output is preserved. Oliguria is associated with worse clinical outcomes. Preserved urine output can be seen in nephrogenic diabetes insipidus characteristic of longstanding urinary tract obstruction, tubulointerstitial disease, or nephrotoxicity from cisplatin or aminoglycosides, among other causes. Red or brown urine may be seen with or without gross hematuria; if the color persists in the supernatant after centrifugation, then pigment nephropathy from rhabdomyolysis or hemolysis should be suspected.

The urinalysis and urine sediment examination are invaluable tools, but they require clinical correlation because of generally limited sensitivity and specificity (Fig. 334-6) (Chap. 62e). In the absence of preexisting proteinuria from CKD, AKI from ischemia or nephrotoxins leads to mild proteinuria (<1 g/d). Greater proteinuria in AKI suggests damage to the glomerular ultrafiltration barrier or excretion of myeloma light chains; the latter are not detected with conventional urine dipsticks (which detect albumin) and require the sulfosalicylic acid test or immunoelectrophoresis. Atheroemboli can cause a variable degree of proteinuria. Extremely heavy proteinuria (“nephrotic range,” >3.5 g/d) can occasionally be seen in glomerulonephritis, vasculitis, or interstitial nephritis (particularly from NSAIDs). AKI can also complicate cases of minimal change disease, a cause of the nephrotic syndrome (Chap. 332e). If the dipstick is positive for hemoglobin but few red blood cells are evident in the urine sediment, then rhabdomyolysis or hemolysis should be suspected.

Prerenal azotemia may present with hyaline casts or an unremarkable urine sediment exam. Postrenal AKI may also lead to an unremarkable sediment, but hematuria and pyuria may be seen depending on the cause of obstruction. AKI from ATN due to ischemic injury, sepsis, or certain nephrotoxins has characteristic urine sediment findings: pigmented “muddy brown” granular casts and tubular epithelial cell casts. These findings may be absent in more than 20% of cases, however. Glomerulonephritis may lead to dysmorphic red blood cells or red blood cell casts. Interstitial nephritis may lead to white blood cell casts. The urine sediment findings overlap somewhat in glomerulonephritis and interstitial nephritis, and a diagnosis is not always possible on the basis of the urine sediment alone. Urine eosinophils have a limited role in differential diagnosis; they can be seen in interstitial nephritis, pyelonephritis, cystitis, atheroembolic disease, or glomerulonephritis. Crystalluria may be important diagnostically. The finding of oxalate crystals in AKI should prompt an evaluation for ethylene glycol toxicity. Abundant uric acid crystals may be seen in the tumor lysis syndrome.

BLOOD LABORATORY FINDINGS

Certain forms of AKI are associated with characteristic patterns in the rise and fall of SCr. Prerenal azotemia typically leads to modest rises in SCr that return to baseline with improvement in hemodynamic status. Contrast nephropathy leads to a rise in SCr within 24–48 h, peak within 3–5 days, and resolution within 5–7 days. In comparison, atheroembolic disease usually manifests with more subacute rises in

TABLE 334-1 MAJOR CAUSES, CLINICAL FEATURES, AND DIAGNOSTIC STUDIES FOR PRERENAL AND INTRINSIC ACUTE KIDNEY INJURY

Etiology	Clinical Features	Laboratory Features	Comments
Prerenal azotemia	History of poor fluid intake or fluid loss (hemorrhage, diarrhea, vomiting, sequestration into extravascular space); NSAID/ACE-I/ARB; heart failure; evidence of volume depletion (tachycardia, absolute or postural hypotension, low jugular venous pressure, dry mucous membranes), decreased effective circulatory volume (cirrhosis, heart failure)	BUN/creatinine ratio above 20, FeNa <1%, hyaline casts in urine sediment, urine specific gravity >1.018, urine osmolality >500 mOsm/kg	Low FeNa, high specific gravity and osmolality may not be seen in the setting of CKD, diuretic use; BUN elevation out of proportion to creatinine may alternatively indicate upper GI bleed or increased catabolism. Response to restoration of hemodynamics is most diagnostic.
Sepsis-associated AKI	Sepsis, sepsis syndrome, or septic shock. Overt hypotension not always seen in mild to moderate AKI	Positive culture from normally sterile body fluid; urine sediment often contains granular casts, renal tubular epithelial cell casts	FeNa may be low (<1%), particularly early in the course, but is usually >1% with osmolality <500 mOsm/kg
Ischemia-associated AKI	Systemic hypotension, often superimposed upon sepsis and/or reasons for limited renal reserve such as older age, CKD	Urine sediment often contains granular casts, renal tubular epithelial cell casts. FeNa typically >1%.	
Nephrotoxin-Associated AKI: Endogenous			
Rhabdomyolysis	Traumatic crush injuries, seizures, immobilization	Elevated myoglobin, creatine kinase; urine heme positive with few red blood cells	FeNa may be low (<1%)
Hemolysis	Recent blood transfusion with transfusion reaction	Anemia, elevated LDH, low haptoglobin	FeNa may be low (<1%); evaluation for transfusion reaction
Tumor lysis	Recent chemotherapy	Hyperphosphatemia, hypocalcemia, hyperuricemia	
Multiple myeloma	Age >60 years, constitutional symptoms, bone pain	Monoclonal spike in urine or serum electrophoresis; low anion gap; anemia	Bone marrow or renal biopsy can be diagnostic
Nephrotoxin-Associated AKI: Exogenous			
Contrast nephropathy	Exposure to iodinated contrast	Characteristic course is rise in SCr within 1–2 d, peak within 3–5 d, recovery within 7 d	FeNa may be low (<1%)
Tubular injury	Aminoglycoside antibiotics, cisplatin, tenofovir, zoledronate, ethylene glycol, aristolochic acid, and melamine (to name a few)	Urine sediment often contains granular casts, renal tubular epithelial cell casts. FeNa typically >1%.	Can be oliguric or nonoliguric
Interstitial nephritis	Recent medication exposure; can have fever, rash, arthralgias	Eosinophilia, sterile pyuria; often nonoliguric	Urine eosinophils have limited diagnostic accuracy; systemic signs of drug reaction often absent; kidney biopsy may be helpful
Other Causes of Intrinsic AKI			
Glomerulonephritis/vasculitis	Variable (Chap. 338) features include skin rash, arthralgias, sinusitis (AGBM disease), lung hemorrhage (AGBM, ANCA, lupus), recent skin infection or pharyngitis (poststreptococcal)	ANA, ANCA, AGBM antibody, hepatitis serologies, cryoglobulins, blood culture, decreased complement levels, ASO titer (abnormalities of these tests depending on etiology)	Kidney biopsy may be necessary
Interstitial nephritis	Nondrug-related causes include tubulointerstitial nephritis-uveitis (TINU) syndrome, <i>Legionella</i> infection	Eosinophilia, sterile pyuria; often nonoliguric	Urine eosinophils have limited diagnostic accuracy; kidney biopsy may be necessary
TTP/HUS	Neurologic abnormalities and/or AKI; recent diarrheal illness; use of calcineurin inhibitors; pregnancy or postpartum; spontaneous	Schistocytes on peripheral blood smear, elevated LDH, anemia, thrombocytopenia	“Typical HUS” refers to AKI with a diarrheal prodrome, often due to Shiga toxin released from <i>Escherichia coli</i> or other bacteria; “atypical HUS” is due to inherited or acquired complement dysregulation. “TTP-HUS” refers to sporadic cases in adults. Diagnosis may involve screening for ADAMTS13 activity, Shiga toxin-producing <i>E. coli</i> , genetic evaluation of complement regulatory proteins, and kidney biopsy.
Atheroembolic disease	Recent manipulation of the aorta or other large vessels; may occur spontaneously or after anticoagulation; retinal plaques, palpable purpura, livedo reticularis, GI bleed	Hypocomplementemia, eosinophiluria (variable), variable amounts of proteinuria	Skin or kidney biopsy can be diagnostic
Postrenal AKI	History of kidney stones, prostate disease, obstructed bladder catheter, retroperitoneal or pelvic neoplasm	No specific findings other than AKI; may have pyuria or hematuria	Imaging with computed tomography or ultrasound

Abbreviations: ACE-I, angiotensin-converting enzyme inhibitor-I; AGBM, antglomerular basement membrane; AKI, acute kidney injury; ANA, antinuclear antibody; ANCA, antineutrophilic cytoplasmic antibody; ARB, angiotensin receptor blocker; ASO, antistreptolysin O; BUN, blood urea nitrogen; CKD, chronic kidney disease; FeNa, fractional excretion of sodium; GI, gastrointestinal; LDH, lactate dehydrogenase; NSAID, nonsteroidal anti-inflammatory drug; TTP/HUS, thrombotic thrombocytopenic purpura/hemolytic-uremic syndrome.

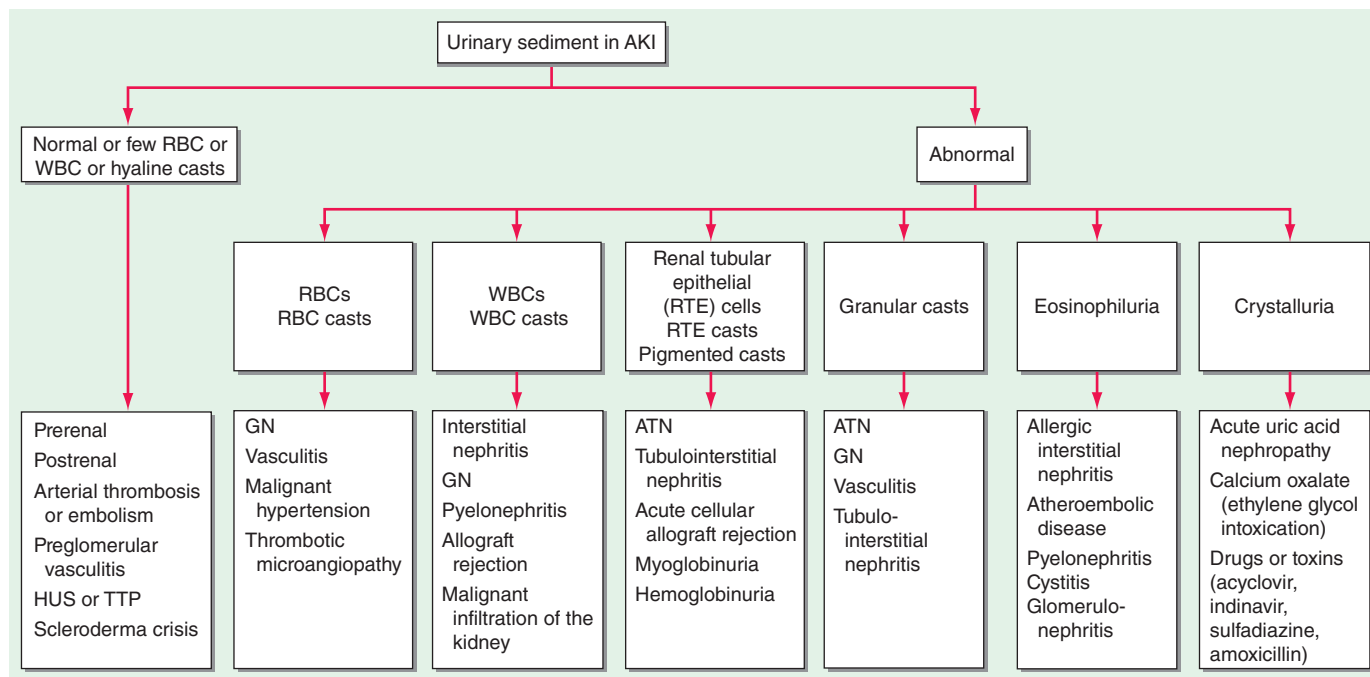


FIGURE 334-6 Interpretation of urinary sediment findings in acute kidney injury (AKI). ATN, acute tubular necrosis; GN, glomerulonephritis; HUS, hemolytic-uremic syndrome; RBCs, red blood cells; RTE, renal tubular epithelial; TTP, thrombotic thrombocytopenic purpura; WBCs, white blood cells. (Adapted from L Yang, JV Bonventre: *Diagnosis and clinical evaluation of acute kidney injury*. In *Comprehensive Nephrology*, 4th ed. J Floege et al [eds]. Philadelphia, Elsevier, 2010.)

SCr, although severe AKI with rapid increases in SCr can occur in this setting. With many of the epithelial cell toxins such as aminoglycoside antibiotics and cisplatin, the rise in SCr is characteristically delayed for 3–5 days to 2 weeks after initial exposure.

A complete blood count may provide diagnostic clues. Anemia is common in AKI and is usually multifactorial in origin. It is not related to an effect of AKI solely on production of red blood cells because this effect in isolation takes longer to manifest. Peripheral eosinophilia can accompany interstitial nephritis, atheroembolic disease, polyarteritis nodosa, and Churg-Strauss vasculitis. Severe anemia in the absence of bleeding may reflect hemolysis, multiple myeloma, or thrombotic microangiopathy (e.g., HUS or TTP). Other laboratory findings of thrombotic microangiopathy include thrombocytopenia, schistocytes on peripheral blood smear, elevated lactate dehydrogenase level, and low haptoglobin content. Evaluation of patients suspected of having TTP-HUS includes measurement of levels of the von Willebrand factor cleaving protease (ADAMTS13) and testing for Shiga toxin-producing *Escherichia coli*. “Atypical HUS” constitutes the majority of adult cases of HUS; genetic testing is important because it is estimated that 60–70% of atypical HUS patients have mutations in genes encoding proteins that regulate the alternative complement pathway.

AKI often leads to hyperkalemia, hyperphosphatemia, and hypocalcemia. Marked hyperphosphatemia with accompanying hypocalcemia, however, suggests rhabdomyolysis or the tumor lysis syndrome. Creatine phosphokinase levels and serum uric acid are elevated in rhabdomyolysis, while tumor lysis syndrome shows normal or marginally elevated creatine kinase and markedly elevated serum uric acid. The anion gap may be increased with any cause of uremia due to retention of anions such as phosphate, hippurate, sulfate, and urate. The co-occurrence of an increased anion gap and an osmolal gap may suggest ethylene glycol poisoning, which may also cause oxalate crystalluria. Low anion gap may provide a clue to the diagnosis of multiple myeloma due to the presence of unmeasured cationic proteins. Laboratory blood tests helpful for the diagnosis of glomerulonephritis and vasculitis include depressed complement levels and high titers of antinuclear antibodies (ANAs), antineutrophilic cytoplasmic antibodies (ANCAs), antiglomerular basement membrane (AGBM) antibodies, and cryoglobulins.

RENAL FAILURE INDICES

Several indices have been used to help differentiate prerenal azotemia from intrinsic AKI when the tubules are malfunctioning. The low tubular flow rate and increased renal medullary recycling of urea seen in prerenal azotemia may cause a disproportionate elevation of the BUN compared to creatinine. Other causes of disproportionate BUN elevation need to be kept in mind, however, including upper gastrointestinal bleeding, hyperalimentation, increased tissue catabolism, and glucocorticoid use.

The fractional excretion of sodium (FeNa) is the fraction of the filtered sodium load that is reabsorbed by the tubules, and is a measure of both the kidney’s ability to reabsorb sodium as well as endogenously and exogenously administered factors that affect tubular reabsorption. As such, it depends on sodium intake, effective intravascular volume, GFR, diuretic intake, and intact tubular reabsorptive mechanisms. With prerenal azotemia, the FeNa may be below 1%, suggesting avid tubular sodium reabsorption. In patients with CKD, a FeNa significantly above 1% can be present despite a superimposed prerenal state. The FeNa may also be above 1% despite hypovolemia due to treatment with diuretics. Low FeNa is often seen early in glomerulonephritis and other disorders and, hence, should not be taken as prima facie evidence of prerenal azotemia. Low FeNa is therefore suggestive, but not synonymous, with effective intravascular volume depletion, and should not be used as the sole guide for volume management. The response of urine output to crystalloid or colloid fluid administration may be both diagnostic and therapeutic in prerenal azotemia. In ischemic AKI, the FeNa is frequently above 1% because of tubular injury and resultant inability to reabsorb sodium. Several causes of ischemia-associated and nephrotoxin-associated AKI can present with FeNa below 1%, however, including sepsis (often early in the course), rhabdomyolysis, and contrast nephropathy.

The ability of the kidney to produce a concentrated urine is dependent upon many factors and reliant on good tubular function in multiple regions of the kidney. In the patient not taking diuretics and with good baseline kidney function, urine osmolality may be above 500 mOsm/kg in prerenal azotemia, consistent with an intact medullary gradient and elevated serum vasopressin levels causing water reabsorption resulting in concentrated urine. In elderly patients and those

with CKD, however, baseline concentrating defects may exist, making urinary osmolality unreliable in many instances. Loss of concentrating ability is common in septic or ischemic AKI, resulting in urine osmolality below 350 mOsm/kg, but the finding is not specific.

RADIOLOGIC EVALUATION

Postrenal AKI should always be considered in the differential diagnosis of AKI because treatment is usually successful if instituted early. Simple bladder catheterization can rule out urethral obstruction. Imaging of the urinary tract with renal ultrasound or CT should be undertaken to investigate obstruction in individuals with AKI unless an alternate diagnosis is apparent. Findings of obstruction include dilation of the collecting system and hydroureteronephrosis. Obstruction can be present without radiologic abnormalities in the setting of volume depletion, retroperitoneal fibrosis, encasement with tumor, and also early in the course of obstruction. If a high clinical index of suspicion for obstruction persists despite normal imaging, antegrade or retrograde pyelography should be performed. Imaging may also provide additional helpful information about kidney size and echogenicity to assist in the distinction between acute versus CKD. In CKD, kidneys are usually smaller unless the patient has diabetic nephropathy, HIV-associated nephropathy, or infiltrative diseases. Normal sized kidneys are expected in AKI. Enlarged kidneys in a patient with AKI suggests the possibility of acute interstitial nephritis. Vascular imaging may be useful if venous or arterial obstruction is suspected, but the risks of contrast administration should be kept in mind. MRI with gadolinium-based contrast agents should be avoided if possible in severe AKI due to the possibility of inducing nephrogenic system fibrosis, a rare but serious complication seen most commonly in patients with end-stage renal disease.

KIDNEY BIOPSY

If the cause of AKI is not apparent based on the clinical context, physical examination, laboratory studies, and radiologic evaluation, kidney biopsy should be considered. The kidney biopsy can provide definitive diagnostic and prognostic information about acute kidney disease and CKD. The procedure is most often used in AKI when prerenal azotemia, postrenal AKI, and ischemic or nephrotoxic AKI have been deemed unlikely, and other possible diagnoses are being considered such as glomerulonephritis, vasculitis, interstitial nephritis, myeloma kidney, HUS and TTP, and allograft dysfunction. Kidney biopsy is associated with a risk of bleeding, which can be severe and organ- or life-threatening in patients with thrombocytopenia or coagulopathy.

NOVEL BIOMARKERS

BUN and creatinine are functional biomarkers of glomerular filtration rather than tissue injury biomarkers and, therefore, may be suboptimal for the diagnosis of actual parenchymal kidney damage. BUN and creatinine are also relatively slow to rise after kidney injury. Several novel kidney injury biomarkers have been investigated and show promise for earlier and accurate diagnosis of AKI. *Kidney injury molecule-1* (KIM-1) is a type 1 transmembrane protein that is abundantly expressed in proximal tubular cells injured by ischemia or nephrotoxins such as cisplatin. KIM-1 is not expressed in appreciable quantities in the absence of tubular injury or in extrarenal tissues. KIM-1's functional role may be to confer phagocytic properties to tubular cells, enabling them to clear debris from the tubular lumen after kidney injury. KIM-1 can be detected shortly after ischemic or nephrotoxic injury in the urine and, therefore, may be an easily tested biomarker in the clinical setting. *Neutrophil gelatinase associated lipocalin* (NGAL, also known as lipocalin-2 or siderocalin) is another novel biomarker of AKI. NGAL was first discovered as a protein in granules of human neutrophils. NGAL can bind to iron siderophore complexes and may have tissue-protective effects in the proximal tubule. NGAL is highly upregulated after inflammation and kidney injury and can be detected in the plasma and urine within 2 h of cardiopulmonary bypass-associated AKI. Other candidate biomarkers of AKI include interleukin (IL) 18, a proinflammatory cytokine of the IL-1 superfamily that may mediate ischemic proximal tubular injury, and L-type fatty acid binding protein, which is expressed

in ischemic proximal tubule cells and may be renoprotective by binding free fatty acids and lipid peroxidation products. A number of other biomarkers are under investigation for early and accurate identification of AKI and for risk stratification to identify individuals at increased risk. The optimal use of novel AKI biomarkers in clinical settings is an area of ongoing investigation.

COMPLICATIONS

The kidney plays a central role in homeostatic control of volume status, blood pressure, plasma electrolyte composition, and acid-base balance, and for excretion of nitrogenous and other waste products. Complications associated with AKI are, therefore, protean, and depend on the severity of AKI and other associated conditions. Mild to moderate AKI may be entirely asymptomatic, particularly early in the course.

UREMIA

Buildup of nitrogenous waste products, manifested as an elevated BUN concentration, is a hallmark of AKI. BUN itself poses little direct toxicity at levels below 100 mg/dL. At higher concentrations, mental status changes and bleeding complications can arise. Other toxins normally cleared by the kidney may be responsible for the symptom complex known as uremia. Few of the many possible uremic toxins have been definitively identified. The correlation of BUN and SCr concentrations with uremic symptoms is extremely variable, due in part to differences in urea and creatinine generation rates across individuals.

HYPERVOLEMIA AND HYPOVOLEMIA

Expansion of extracellular fluid volume is a major complication of oliguric and anuric AKI, due to impaired salt and water excretion. The result can be weight gain, dependent edema, increased jugular venous pressure, and pulmonary edema; the latter can be life threatening. Pulmonary edema can also occur from volume overload and hemorrhage in pulmonary renal syndromes. AKI may also induce or exacerbate acute lung injury characterized by increased vascular permeability and inflammatory cell infiltration in lung parenchyma. Recovery from AKI can sometimes be accompanied by polyuria, which, if untreated, can lead to significant volume depletion. The polyuric phase of recovery may be due to an osmotic diuresis from retained urea and other waste products as well as delayed recovery of tubular reabsorptive functions.

HYPONATREMIA

Administration of excessive hypotonic crystalloid or isotonic dextrose solutions can result in hyposmolality and hyponatremia, which, if severe, can cause neurologic abnormalities, including seizures.

HYPERKALEMIA

Abnormalities in plasma electrolyte composition can be mild or life threatening. Frequently the most concerning complication of AKI is hyperkalemia. Marked hyperkalemia is particularly common in rhabdomyolysis, hemolysis, and tumor lysis syndrome due to release of intracellular potassium from damaged cells. Potassium affects the cellular membrane potential of cardiac and neuromuscular tissues. Muscle weakness may be a symptom of hyperkalemia. The more serious complication of hyperkalemia is due to effects on cardiac conduction, leading to potentially fatal arrhythmias.

ACIDOSIS

Metabolic acidosis, usually accompanied by an elevation in the anion gap, is common in AKI, and can further complicate acid-base and potassium balance in individuals with other causes of acidosis, including sepsis, diabetic ketoacidosis, or respiratory acidosis.

HYPERPHOSPHATEMIA AND HYPOCALCEMIA

AKI can lead to hyperphosphatemia, particularly in highly catabolic patients or those with AKI from rhabdomyolysis, hemolysis, and tumor lysis syndrome. Metastatic deposition of calcium phosphate can lead to hypocalcemia. AKI-associated hypocalcemia may also arise from derangements in the vitamin D–parathyroid hormone–fibroblast

growth factor-23 axis. Hypocalcemia is often asymptomatic but can lead to perioral paresthesias, muscle cramps, seizures, carpopedal spasms, and prolongation of the QT interval on electrocardiography. Calcium levels should be corrected for the degree of hypoalbuminemia, if present, or ionized calcium levels should be followed. Mild, asymptomatic hypocalcemia does not require treatment.

BLEEDING

Hematologic complications of AKI include anemia and bleeding, both of which are exacerbated by coexisting disease processes such as sepsis, liver disease, and disseminated intravascular coagulation. Direct hematologic effects from AKI-related uremia include decreased erythropoiesis and platelet dysfunction.

INFECTIONS

Infections are a common precipitant of AKI and also a dreaded complication of AKI. Impaired host immunity has been described in end-stage renal disease and may be operative in severe AKI.

CARDIAC COMPLICATIONS

The major cardiac complications of AKI are arrhythmias, pericarditis, and pericardial effusion.

MALNUTRITION

AKI is often a severely hypercatabolic state, and therefore, malnutrition is a major complication.

TREATMENT ACUTE KIDNEY INJURY

PREVENTION AND TREATMENT

The management of individuals with and at risk for AKI varies according to the underlying cause (Table 334-2). Common to all are several principles. Optimization of hemodynamics, correction of fluid and electrolyte imbalances, discontinuation of nephrotoxic medications, and dose adjustment of administered medications are all critical. Common causes of AKI such as sepsis and ischemic ATN do not yet have specific therapies once injury is established, but meticulous clinical attention is needed to support the patient until (if) AKI resolves. The kidney possesses remarkable capacity to repair itself after even severe, dialysis-requiring AKI. However, many patients with AKI do not recover fully and may remain dialysis dependent. It has become increasingly apparent that AKI predisposes to accelerated progression of CKD, and CKD is an important risk factor for AKI.

Prerenal Azotemia Prevention and treatment of prerenal azotemia require optimization of renal perfusion. The composition of replacement fluids should be targeted to the type of fluid lost. Severe acute blood loss should be treated with packed red blood cells. Isotonic crystalloid and/or colloid should be used for less severe acute hemorrhage or plasma loss in the case of burns and pancreatitis. Crystalloid solutions are less expensive and probably equally efficacious as colloid solutions. Hydroxyethyl starch solutions increase the risk of severe AKI and are contraindicated. Crystalloid has been reported to be preferable to albumin in the setting of traumatic brain injury. Isotonic crystalloid (e.g., 0.9% saline) or colloid should be used for volume resuscitation in severe hypovolemia, whereas hypotonic crystalloids (e.g., 0.45% saline) suffice for less severe hypovolemia. Excessive chloride administration from 0.9% saline may lead to hyperchloremic metabolic acidosis and may impair GFR. Bicarbonate-containing solutions (e.g., dextrose water with 150 mEq sodium bicarbonate) should be used if metabolic acidosis is a concern.

Optimization of cardiac function in AKI may require use of inotropic agents, preload- and afterload-reducing agents, antiarrhythmic drugs, and mechanical aids such as an intraaortic balloon pump. Invasive hemodynamic monitoring to guide therapy may be necessary.

Cirrhosis and Hepatorenal Syndrome Fluid management in individuals with cirrhosis, ascites, and AKI is challenging because of the frequent

TABLE 334-2 MANAGEMENT OF ACUTE KIDNEY INJURY

General Issues	
1.	Optimization of systemic and renal hemodynamics through volume resuscitation and judicious use of vasopressors
2.	Elimination of nephrotoxic agents (e.g., ACE inhibitors, ARBs, NSAIDs, aminoglycosides) if possible
3.	Initiation of renal replacement therapy when indicated
Specific Issues	
1.	Nephrotoxin-specific
a.	Rhabdomyolysis: aggressive intravenous fluids; consider forced alkaline diuresis
b.	Tumor lysis syndrome: aggressive intravenous fluids and allopurinol or rasburicase
2.	Volume overload
a.	Salt and water restriction
b.	Diuretics
c.	Ultrafiltration
3.	Hyponatremia
a.	Restriction of enteral free water intake, minimization of hypotonic intravenous solutions including those containing dextrose
b.	Hypertonic saline is rarely necessary in AKI. Vasopressin antagonists are generally not needed.
4.	Hyperkalemia
a.	Restriction of dietary potassium intake
b.	Discontinuation of potassium-sparing diuretics, ACE inhibitors, ARBs, NSAIDs
c.	Loop diuretics to promote urinary potassium loss
d.	Potassium binding ion-exchange resin (sodium polystyrene sulfonate)
e.	Insulin (10 units regular) and glucose (50 mL of 50% dextrose) to promote entry of potassium intracellularly
f.	Inhaled beta-agonist therapy to promote entry of potassium intracellularly
g.	Calcium gluconate or calcium chloride (1 g) to stabilize the myocardium
5.	Metabolic acidosis
a.	Sodium bicarbonate (if pH <7.2 to keep serum bicarbonate >15 mmol/L)
b.	Administration of other bases, e.g., THAM
c.	Renal replacement therapy
6.	Hyperphosphatemia
a.	Restriction of dietary phosphate intake
b.	Phosphate binding agents (calcium acetate, sevelamer hydrochloride, aluminum hydroxide—taken with meals)
7.	Hypocalcemia
a.	Calcium carbonate or calcium gluconate if symptomatic
8.	Hypermagnesemia
a.	Discontinue Mg ²⁺ containing antacids
9.	Hyperuricemia
a.	Acute treatment is usually not required except in the setting of tumor lysis syndrome (see above)
10.	Nutrition
a.	Sufficient protein and calorie intake (20–30 kcal/kg per day) to avoid negative nitrogen balance. Nutrition should be provided via the enteral route if possible.
11.	Drug dosing
a.	Careful attention to dosages and frequency of administration of drugs, adjustment for degree of renal failure
b.	Note that serum creatinine concentration may overestimate renal function in the non–steady state characteristic of patients with AKI

Abbreviations: ACE, angiotensin-converting enzyme; ARBs, angiotensin receptor blocker; NSAIDs, nonsteroidal anti-inflammatory drug; THAM, tris (hydroxymethyl) aminomethane.

difficulty in ascertaining intravascular volume status. Administration of intravenous fluids as a volume challenge may be required diagnostically as well as therapeutically. Excessive volume administration may, however, result in worsening ascites and pulmonary compromise in

the setting of hepatorenal syndrome or AKI due to superimposed spontaneous bacterial peritonitis. Peritonitis should be ruled out by culture of ascitic fluid. Albumin may prevent AKI in those treated with antibiotics for spontaneous bacterial peritonitis. The definitive treatment of the hepatorenal syndrome is orthotopic liver transplantation. Bridge therapies that have shown promise include terlipressin (a vasopressin analog), combination therapy with octreotide (a somatostatin analog) and midodrine (an α_1 -adrenergic agonist), and norepinephrine, in combination with intravenous albumin (25–50 g, maximum 100 g/d).

Intrinsic AKI Several agents have been tested and have failed to show benefit in the treatment of acute tubular injury. These include atrial natriuretic peptide, low-dose dopamine, endothelin antagonists, loop diuretics, calcium channel blockers, α -adrenergic receptor blockers, prostaglandin analogs, antioxidants, antibodies against leukocyte adhesion molecules, and insulin-like growth factor, among many others. Most studies have enrolled patients with severe and well-established AKI, and treatment may have been initiated too late. Novel kidney injury biomarkers may provide an opportunity to test agents earlier in the course of AKI.

AKI due to acute glomerulonephritis or vasculitis may respond to immunosuppressive agents and/or plasmapheresis (**Chap. 332e**). Allergic interstitial nephritis due to medications requires discontinuation of the offending agent. Glucocorticoids have been used, but not tested in randomized trials, in cases where AKI persists or worsens despite discontinuation of the suspected medication. AKI due to scleroderma (scleroderma renal crisis) should be treated with ACE inhibitors. Idiopathic TTP-HUS is a medical emergency and should be treated promptly with plasma exchange. Pharmacologic blockade of complement activation may be effective in atypical HUS.

Early and aggressive volume repletion is mandatory in patients with rhabdomyolysis, who may initially require 10 L of fluid per day. Alkaline fluids (e.g., 75 mmol/L sodium bicarbonate added to 0.45% saline) may be beneficial in preventing tubular injury and cast formation, but carry the risk of worsening hypocalcemia. Diuretics may be used if fluid repletion is adequate but unsuccessful in achieving urinary flow rates of 200–300 mL/h. There is no specific therapy for established AKI in rhabdomyolysis, other than dialysis in severe cases or general supportive care to maintain fluid and electrolyte balance and tissue perfusion. Careful attention must be focused on calcium and phosphate status because of precipitation in damaged tissue and release when the tissue heals.

Postrenal AKI Prompt recognition and relief of urinary tract obstruction can forestall the development of permanent structural damage induced by urinary stasis. The site of obstruction defines the treatment approach. Transurethral or suprapubic bladder catheterization may be all that is needed initially for urethral strictures or functional bladder impairment. Ureteric obstruction may be treated by percutaneous nephrostomy tube placement or ureteral stent placement. Relief of obstruction is usually followed by an appropriate diuresis for several days. In rare cases, severe polyuria persists due to tubular dysfunction and may require continued administration of intravenous fluids and electrolytes for a period of time.

SUPPORTIVE MEASURES

Volume Management Hypervolemia in oliguric or anuric AKI may be life threatening due to acute pulmonary edema, especially because many patients have coexisting pulmonary disease, and AKI likely increases pulmonary vascular permeability. Fluid and sodium should be restricted, and diuretics may be used to increase the urinary flow rate. There is no evidence that increasing urine output itself improves the natural history of AKI, but diuretics may help to avoid the need for dialysis in some cases. In severe cases of volume overload, furosemide may be given as a bolus (200 mg) followed by an intravenous drip (10–40 mg/h), with or without a thiazide diuretic. In decompensated heart failure, stepped diuretic therapy was found to be superior to ultrafiltration in preserving

renal function. Diuretic therapy should be stopped if there is no response. Dopamine in low doses may transiently increase salt and water excretion by the kidney in prerenal states, but clinical trials have failed to show any benefit in patients with intrinsic AKI. Because of the risk of arrhythmias and potential bowel ischemia, it has been argued that the risks of dopamine outweigh the benefits in the treatment or prevention of AKI.

Electrolyte and Acid-Base Abnormalities The treatment of **dysnatremias and hyperkalemia is described in Chap. 63**. Metabolic acidosis is generally not treated unless severe (pH <7.20 and serum bicarbonate <15 mmol/L). Acidosis can be treated with oral or intravenous sodium bicarbonate (**Chap. 66**), but overcorrection should be avoided because of the possibility of metabolic alkalosis, hypocalcemia, hypokalemia, and volume overload. Hyperphosphatemia is common in AKI and can usually be treated by limiting intestinal absorption of phosphate using phosphate binders (calcium carbonate, calcium acetate, lanthanum, sevelamer, or aluminum hydroxide). Hypocalcemia does not usually require therapy unless symptoms are present. Ionized calcium should be monitored rather than total calcium when hypoalbuminemia is present.

Malnutrition Protein energy wasting is common in AKI, particularly in the setting of multisystem organ failure. Inadequate nutrition may lead to starvation ketoacidosis and protein catabolism. Excessive nutrition may increase the generation of nitrogenous waste and lead to worsening azotemia. Total parenteral nutrition requires large volumes of fluid administration and may complicate efforts at volume control. According to the Kidney Disease Improving Global Outcomes (KDIGO) guidelines, patients with AKI should achieve a total energy intake of 20–30 kcal/kg per day. Protein intake should vary depending on the severity of AKI: 0.8–1.0 g/kg per day in noncatabolic AKI without the need for dialysis; 1.0–1.5 g/kg per day in patients on dialysis; and up to a maximum of 1.7 g/kg per day if hypercatabolic and receiving continuous renal replacement therapy. Trace elements and water-soluble vitamins should also be supplemented in AKI patients treated with dialysis and continuous renal replacement therapy.

Anemia The anemia seen in AKI is usually multifactorial and is not improved by erythropoiesis-stimulating agents, due to their delayed onset of action and the presence of bone marrow resistance in critically ill patients. Uremic bleeding may respond to desmopressin or estrogens, but may require dialysis for treatment in the case of long-standing or severe uremia. Gastrointestinal prophylaxis with proton pump inhibitors or histamine (H_2) receptor blockers is required. Venous thromboembolism prophylaxis is important and should be tailored to the clinical setting; low-molecular-weight heparins and factor Xa inhibitors have unpredictable pharmacokinetics in severe AKI and should be avoided.

Dialysis Indications and Modalities (See also Chap. 336) Dialysis is indicated when medical management fails to control volume overload, hyperkalemia, or acidosis; in some toxic ingestions; and when there are severe complications of uremia (asterixis, pericardial rub or effusion, encephalopathy, uremic bleeding). The timing of dialysis is still a matter of debate. Late initiation of dialysis carries the risk of avoidable volume, electrolyte, and metabolic complications of AKI. On the other hand, initiating dialysis too early may unnecessarily expose individuals to intravenous lines and invasive procedures, with the attendant risks of infection, bleeding, procedural complications, and hypotension. The initiation of dialysis should not await the development of a life-threatening complication of renal failure. Many nephrologists initiate dialysis for AKI empirically when the BUN exceeds a certain value (e.g., 100 mg/dL) in patients without clinical signs of recovery of kidney function. The available modes for renal replacement therapy in AKI require either access to the peritoneal cavity (for peritoneal dialysis) or the large blood vessels (for hemodialysis, hemofiltration, and other hybrid procedures). Small solutes are removed across a semipermeable membrane down their concentration

335 Chronic Kidney Disease

Joanne M. Bargman, Karl Skorecki

gradient (“diffusive” clearance) and/or along with the movement of plasma water (“convective” clearance). The choice of modality is often dictated by the immediate availability of technology and the expertise of medical staff. Peritoneal dialysis is performed through a temporary intraperitoneal catheter. It is rarely used in the United States for AKI in adults but has enjoyed widespread use internationally, particularly when hemodialysis technology is not available. Dialysate solution is instilled into and removed from the peritoneal cavity at regular intervals in order to achieve diffusive and convective clearance of solutes across the peritoneal membrane; ultrafiltration of water is achieved by the presence of an osmotic gradient across the peritoneal membrane achieved by high concentrations of dextrose in the dialysate solution. Because of its continuous nature, it is often better tolerated than intermittent procedures like hemodialysis in hypotensive patients. Peritoneal dialysis may not be sufficient for hypercatabolic patients due to inherent limitations in dialysis efficacy.

Hemodialysis can be used intermittently or continuously and can be done through convective clearance, diffusive clearance, or a combination of the two. Vascular access is through the femoral, internal jugular, or subclavian veins. Hemodialysis is an intermittent procedure that removes solutes through diffusive and convective clearance. Hemodialysis is typically performed 3–4 h per day, three to four times per week, and is the most common form of renal replacement therapy for AKI. One of the major complications of hemodialysis is hypotension, particularly in the critically ill.

Continuous intravascular procedures were developed in the early 1980s to treat hemodynamically unstable patients without inducing the rapid shifts of volume, osmolarity, and electrolytes characteristic of intermittent hemodialysis. Continuous renal replacement therapy (CRRT) can be performed by convective clearance (continuous venovenous hemofiltration [CVVH]), in which large volumes of plasma water (and accompanying solutes) are forced across the semipermeable membrane by means of hydrostatic pressure; the plasma water is then replaced by a physiologic crystalloid solution. CRRT can also be performed by diffusive clearance (continuous venovenous hemodialysis [CVVHD]), a technology similar to hemodialysis except at lower blood flow and dialysate flow rates. A hybrid therapy combines both diffusive and convective clearance (continuous venovenous hemodiafiltration [CVVHDF]). To achieve some of the advantages of CRRT without the need for 24-h staffing of the procedure, some physicians favor slow low-efficiency dialysis (SLED) or extended daily dialysis (EDD). In this therapy, blood flow and dialysate flow are higher than in CVVHD, but the treatment time is reduced to 12 h or less.

The optimal dose of dialysis for AKI is not clear. Daily intermittent hemodialysis and high-dose CRRT do not confer a demonstrable survival or renal recovery advantage, but care should be taken to avoid undertreatment. Studies have failed to show that continuous therapies are superior to intermittent therapies. If available, CRRT is often preferred in patients with severe hemodynamic instability, cerebral edema, or significant volume overload.

OUTCOME AND PROGNOSIS

The development of AKI is associated with a significantly increased risk of in-hospital and long-term mortality, longer length of stay, and increased costs. Prerenal azotemia, with the exception of the cardiorenal and hepatorenal syndromes, and postrenal azotemia carry a better prognosis than most cases of intrinsic AKI. The kidneys may recover even after severe, dialysis-requiring AKI. Survivors of an episode of AKI requiring temporary dialysis, however, are at extremely high risk for progressive CKD, and up to 10% may develop end-stage renal disease. Postdischarge care under the supervision of a nephrologist for aggressive secondary prevention of kidney disease is prudent. Patients with AKI are more likely to die prematurely after they leave the hospital even if their kidney function has recovered.

Chronic kidney disease (CKD) encompasses a spectrum of different pathophysiologic processes associated with abnormal kidney function and a progressive decline in glomerular filtration rate (GFR). **Figure 335-1** provides a recently updated classification, in which stages of CKD are stratified by both estimated GFR and the degree of albuminuria, in order to predict risk of progression of CKD. Previously, CKD had been staged solely by the GFR. However, the risk of worsening of kidney function is closely linked to the amount of albuminuria, and so it has been incorporated into the classification.

The pathophysiologic processes, adaptations, clinical presentations, assessment, and therapeutic interventions associated with CKD will be the focus of this chapter. The dispiriting term *end-stage renal disease* represents a stage of CKD where the accumulation of toxins, fluid, and electrolytes normally excreted by the kidneys results in the *uremic syndrome*. This syndrome leads to death unless the toxins are removed by renal replacement therapy, using dialysis or kidney transplantation. These interventions are discussed in **Chaps. 336 and 337**. *End-stage renal disease* will be supplanted in this chapter by the term *stage 5 CKD*.

PATHOPHYSIOLOGY OF CHRONIC KIDNEY DISEASE

The pathophysiology of CKD involves two broad sets of mechanisms of damage: (1) initiating mechanisms specific to the underlying etiology (e.g., genetically determined abnormalities in kidney development or integrity, immune complex deposition and inflammation in certain types of glomerulonephritis, or toxin exposure in certain diseases of the renal tubules and interstitium) and (2) a set of progressive mechanisms, involving hyperfiltration and hypertrophy of the remaining viable nephrons, that are a common consequence following long-term reduction of renal mass, irrespective of underlying etiology (**Chap. 333e**). The responses to reduction in nephron number are mediated by vasoactive hormones, cytokines, and growth factors. Eventually, these short-term adaptations of hypertrophy and hyperfiltration become maladaptive as the increased pressure and flow within the nephron predisposes to distortion of glomerular architecture, abnormal podocyte function, and disruption of the filtration barrier leading to sclerosis and dropout of the remaining nephrons (**Fig. 335-2**). Increased intrarenal activity of the renin-angiotensin system (RAS) appears to contribute both to the initial adaptive hyperfiltration and to the subsequent maladaptive hypertrophy and sclerosis. This process explains why a reduction in renal mass from an isolated insult may lead to a progressive decline in renal function over many years (**Fig. 335-3**).

IDENTIFICATION OF RISK FACTORS AND STAGING OF CKD

It is important to identify factors that increase the risk for CKD, even in individuals with normal GFR. Risk factors include small for gestation birth weight, childhood obesity, hypertension, diabetes mellitus, autoimmune disease, advanced age, African ancestry, a family history of kidney disease, a previous episode of acute kidney injury, and the presence of proteinuria, abnormal urinary sediment, or structural abnormalities of the urinary tract.

Many rare inherited forms of CKD follow a Mendelian inheritance pattern, often as part of a systemic syndrome, with the most common in this category being autosomal dominant polycystic kidney disease. In addition, recent research in the genetics of predisposition to common complex diseases (**Chap. 82**) has revealed DNA sequence variants at a number of genetic loci that are associated with common forms of CKD. A striking example is the finding of allelic versions of the *APOL1* gene, of West African population ancestry, which contributes to the several-fold higher frequency of certain common etiologies of nondiabetic CKD (e.g., focal segmental glomerulosclerosis) observed among African and Hispanic Americans. The prevalence in West African populations seems to have arisen as an evolutionary adaptation conferring protection from tropical pathogens. As in other common diseases

Prognosis of CKD by GFR and albuminuria categories: KDIGO 2012				Persistent albuminuria categories description and range		
				A1	A2	A3
				Normal to mildly increased	Moderately increased	Severely increased
				<30 mg/g <3 mg/mmol	30–300 mg/g 3–30 mg/mmol	>300 mg/g >30 mg/mmol
GFR categories (ml/min/1.73 m ²) description and range	G1	Normal or high	≥90			
	G2	Mildly decreased	60–89			
	G3a	Mildly to moderately decreased	45–59			
	G3b	Moderately to severely decreased	30–44			
	G4	Severely decreased	15–29			
	G5	Kidney failure	<15			

FIGURE 335-1 Kidney Disease Improving Global Outcome (KDIGO) classification of chronic kidney disease (CKD). Gradation of color from green to red corresponds to increasing risk and progression of CKD. GFR, glomerular filtration rate. (Reproduced with permission from *Kidney Int Suppl* 3:5-14, 2013.)

with a heritable component, an environmental trigger (such as a viral pathogen) is required to transform genetic risk into disease.

To stage CKD, it is necessary to estimate the GFR rather than relying on serum creatinine concentration (Table 335-1). Many laboratories now report an estimated GFR, or eGFR, using one of these equations.

The normal annual mean decline in GFR with age from the peak GFR (~120 mL/min per 1.73 m²) attained during the third decade of life is ~1 mL/min per year per 1.73 m², reaching a mean value of

70 mL/min per 1.73 m² at age 70. Although reduced GFR occurs with human aging, the lower GFR signifies a true loss of kidney function, with all of the implications that apply to the corresponding stage of CKD. The mean GFR is lower in women than in men. For example, a woman in her 80s with a normal serum creatinine may have a GFR of just 50 mL/min per 1.73 m². Thus, even a mild elevation in serum creatinine concentration (e.g., 130 μmol/L [1.5 mg/dL]) often signifies a substantial reduction in GFR in most individuals.

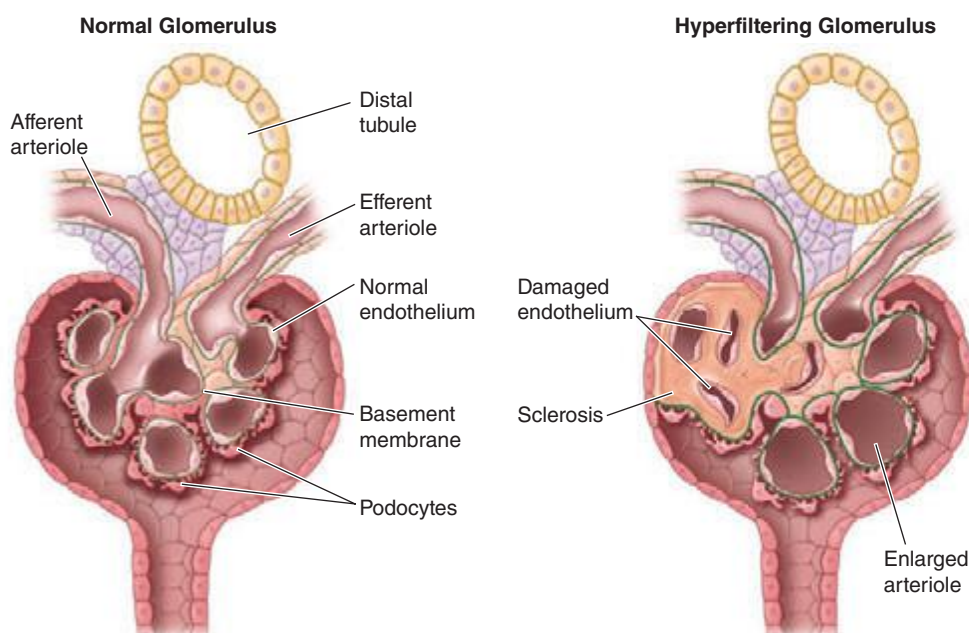


FIGURE 335-2 Left: Schema of the normal glomerular architecture. Right: Secondary glomerular changes associated with a reduction in nephron number, including enlargement of capillary lumens and focal adhesions, which are thought to occur consequent to compensatory hyperfiltration and hypertrophy in the remaining nephrons. (Modified from *JR Ingelfinger: N Engl J Med* 348:99, 2003.)

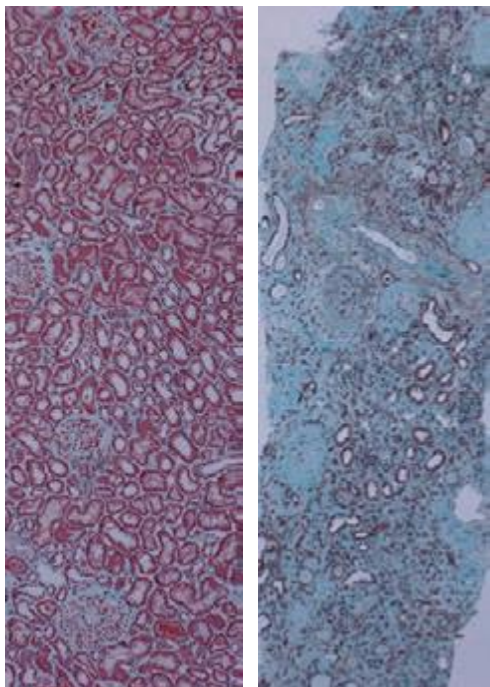


FIGURE 335-3 *Left:* Low-power photomicrograph of a normal kidney showing normal glomeruli and healthy tubulointerstitium without fibrosis. *Right:* Low-power photomicrograph of chronic kidney disease with sclerosis of many glomeruli and severe tubulointerstitial fibrosis (Masson trichrome, $\times 40$ magnification). (Slides courtesy of the late Dr. Andrew Herzenberg.)

The equations for estimating GFR are valid only if the patient is in steady state, that is, the serum creatinine is neither rising nor falling over days.

Measurement of albuminuria is also helpful for monitoring nephron injury and the response to therapy in many forms of CKD, especially chronic glomerular diseases. Although an accurate 24-h urine collection is the standard for measurement of albuminuria, the measurement of protein-to-creatinine ratio in a spot first-morning urine sample is often more practical to obtain and correlates well, but not perfectly, with 24-h urine collections. *Microalbuminuria* (Fig. 335-1, stage A2) refers to the excretion of amounts of albumin too small to detect by urinary dipstick or conventional measures of urine protein. It is a good screening test for early detection of renal disease, and may be a marker for the presence of microvascular disease in general. If a patient has a large amount of excreted albumin, there is no reason to test for microalbuminuria.

Stages 1 and 2 CKD are usually not associated with any symptoms arising from the decrement in GFR. If the decline in GFR progresses to stages 3 and 4, clinical and laboratory complications of CKD become

TABLE 335-1 RECOMMENDED EQUATIONS FOR ESTIMATION OF GLOMERULAR FILTRATION RATE (GFR) USING SERUM CREATININE CONCENTRATION (S_{Cr}), AGE, SEX, RACE, AND BODY WEIGHT

1. Equation from the Modification of Diet in Renal Disease study

$$\text{Estimated GFR (mL/min per } 1.73 \text{ m}^2) = 1.86 \times (S_{Cr})^{-1.154} \times (\text{age})^{-0.203}$$

Multiply by 0.742 for women

Multiply by 1.21 for African ancestry

2. CKD-EPI equation

$$\text{GFR} = 141 \times \min(S_{Cr}/\kappa, 1)^{\alpha} \times \max(S_{Cr}/\kappa, 1)^{-1.209} \times 0.993^{\text{Age}}$$

Multiply by 1.018 for women

Multiply by 1.159 for African ancestry

where S_{Cr} is serum creatinine in mg/dL, κ is 0.7 for females and 0.9 for males, α is -0.329 for females and -0.411 for males, min indicates the minimum of S_{Cr}/κ or 1, and max indicates the maximum of S_{Cr}/κ or 1.

Abbreviation: CKD-EPI, Chronic Kidney Disease Epidemiology Collaboration.

more prominent. Virtually all organ systems are affected, but the most evident complications include anemia and associated easy fatigability; decreased appetite with progressive malnutrition; abnormalities in calcium, phosphorus, and mineral-regulating hormones, such as $1,25(\text{OH})_2\text{D}_3$ (calcitriol), parathyroid hormone (PTH), and fibroblast growth factor 23 (FGF-23); and abnormalities in sodium, potassium, water, and acid-base homeostasis. Many patients, especially the elderly, will have eGFR values compatible with stage 2 or 3 CKD. However, the majority of these patients will show no further deterioration of renal function. The primary care physician is advised to recheck kidney function, and if it is stable and not associated with proteinuria, the patient can usually be managed in this setting. However, if there is evidence of decline of GFR, uncontrolled hypertension, or proteinuria, referral to a nephrologist is appropriate. If the patient progresses to stage 5 CKD, toxins accumulate such that patients usually experience a marked disturbance in their activities of daily living, well-being, nutritional status, and water and electrolyte homeostasis, eventuating in the *uremic syndrome*.

ETIOLOGY AND EPIDEMIOLOGY

It has been estimated from population survey data that at least 6% of the adult population in the United States has CKD at stages 1 and 2. An additional 4.5% of the U.S. population is estimated to have stages 3 and 4 CKD. **Table 335-2** lists the five most frequent categories of causes of CKD, cumulatively accounting for greater than 90% of the CKD disease burden worldwide. The relative contribution of each category varies among different geographic regions. The most frequent cause of CKD in North America and Europe is diabetic nephropathy, most often secondary to type 2 diabetes mellitus. Patients with newly diagnosed CKD often also present with hypertension. When no overt evidence for a primary glomerular or tubulointerstitial kidney disease process is present, CKD is often attributed to hypertension. However, it is now appreciated that such individuals can be considered in two categories. The first includes patients with a silent primary glomerulopathy, such as focal segmental glomerulosclerosis, without the overt nephrotic or nephritic manifestations of glomerular disease (**Chap. 338**). The second includes patients in whom progressive nephrosclerosis and hypertension is the renal correlate of a systemic vascular disease, often also involving large- and small-vessel cardiac and cerebral pathology. This latter combination is especially common in the elderly, in whom chronic renal ischemia as a cause of CKD may be underdiagnosed. The increasing incidence of CKD in the elderly has been ascribed, in part, to decreased mortality rate from the cardiac and cerebral complications of atherosclerotic vascular disease, enabling a greater segment of the population to eventually manifest the renal component of generalized vascular disease. Nevertheless, it should be appreciated that the vast majority of such patients with early stages of CKD will succumb to the cardiovascular and cerebrovascular consequences of the vascular disease before they can progress to the most advanced stages of CKD. Indeed, even a minor decrement in GFR or the presence of albuminuria is now recognized as a major risk factor for cardiovascular disease.

PATHOPHYSIOLOGY AND BIOCHEMISTRY OF UREMIA

Although serum urea and creatinine concentrations are used to measure the excretory capacity of the kidneys, accumulation of these two molecules themselves does not account for the many symptoms and signs that characterize the uremic syndrome in advanced renal failure.

TABLE 335-2 LEADING CATEGORIES OF ETIOLOGIES OF CKD^a

- Diabetic nephropathy
- Glomerulonephritis
- Hypertension-associated CKD (includes vascular and ischemic kidney disease and primary glomerular disease with associated hypertension)
- Autosomal dominant polycystic kidney disease
- Other cystic and tubulointerstitial nephropathy

^aRelative contribution of each category varies with geographic region and race.

Hundreds of toxins that accumulate in renal failure have been implicated in the uremic syndrome. These include water-soluble, hydrophobic, protein-bound, charged, and uncharged compounds. Additional categories of nitrogenous excretory products include guanidino compounds, urates and hippurates, products of nucleic acid metabolism, polyamines, myoinositol, phenols, benzoates, and indoles. It is thus evident that the serum concentrations of urea and creatinine should be viewed as being readily measured, but incomplete, surrogate markers for these compounds, and monitoring the levels of urea and creatinine in the patient with impaired kidney function represents a vast oversimplification of the uremic state.

The uremic syndrome and the disease state associated with advanced renal impairment involve more than renal excretory failure. A host of metabolic and endocrine functions normally performed by the kidneys is also impaired or suppressed, and this results in anemia, malnutrition, and abnormal metabolism of carbohydrates, fats, and proteins. Furthermore, plasma levels of many hormones, including PTH, FGF-23, insulin, glucagon, steroid hormones including vitamin D and sex hormones, and prolactin, change with CKD as a result of reduced excretion, decreased degradation, or abnormal regulation. Finally, CKD is associated with worsening systemic inflammation. Elevated levels of C-reactive protein are detected along with other acute-phase reactants, whereas levels of so-called negative acute-phase reactants, such as albumin and fetuin, decline with progressive reduction in GFR. Thus, the inflammation associated with CKD is important in the *malnutrition-inflammation-atherosclerosis/calcification syndrome*, which contributes in turn to the acceleration of vascular disease and comorbidity associated with advanced kidney disease.

In summary, the pathophysiology of the uremic syndrome can be divided into manifestations in three spheres of dysfunction: (1) those consequent to the accumulation of toxins that normally undergo renal excretion, including products of protein metabolism; (2) those consequent to the loss of other kidney functions, such as fluid and electrolyte homeostasis and hormone regulation; and (3) progressive systemic inflammation and its vascular and nutritional consequences.

CLINICAL AND LABORATORY MANIFESTATIONS OF CHRONIC KIDNEY DISEASE AND UREMIA

Uremia leads to disturbances in the function of virtually every organ system. Chronic dialysis can reduce the incidence and severity of many of these disturbances, so that the overt and florid manifestations of uremia have largely disappeared in the modern health setting. However, even optimal dialysis therapy is not completely effective as renal replacement therapy, because some disturbances resulting from impaired kidney function fail to respond to dialysis.

FLUID, ELECTROLYTE, AND ACID-BASE DISORDERS

Sodium and Water Homeostasis In most patients with stable CKD, the total-body content of sodium and water is modestly increased, although this may not be apparent on clinical examination. With normal renal function, the tubular reabsorption of filtered sodium and water is adjusted so that urinary excretion matches intake. Many forms of kidney disease (e.g., glomerulonephritis) disrupt this balance such that dietary intake of sodium exceeds its urinary excretion, leading to sodium retention and attendant extracellular fluid volume (ECFV) expansion. This expansion may contribute to hypertension, which itself can accelerate the nephron injury. As long as water intake does not exceed the capacity for water clearance, the ECFV expansion will be isotonic and the patient will have a normal plasma sodium concentration (Chap. 333e). Hyponatremia is not commonly seen in CKD patients but, when present, often responds to water restriction. The patient with ECFV expansion (peripheral edema, sometimes hypertension poorly responsive to therapy) should be counseled regarding salt restriction. Thiazide diuretics have limited utility in stages 3–5 CKD, such that administration of loop diuretics, including furosemide, bumetanide, or torsemide, may also be needed. Resistance to loop diuretics in CKD often mandates use of higher doses than those used in patients with more normal kidney function. The combination of

loop diuretics with metolazone, which inhibits the sodium chloride co-transporter of the distal convoluted tubule, can promote renal salt excretion. Diuretic resistance with intractable edema and hypertension in advanced CKD may serve as an indication to initiate dialysis.

In addition to problems with salt and water excretion, some patients with CKD may instead have impaired renal conservation of sodium and water. When an extrarenal cause for fluid loss, such as gastrointestinal (GI) loss, is present, these patients may be prone to ECFV depletion because of the inability of the failing kidney to reclaim filtered sodium adequately. Furthermore, depletion of ECFV, whether due to GI losses or overzealous diuretic therapy, can further compromise kidney function through underperfusion, or a “prerenal” basis, leading to acute-on-chronic kidney failure. In this setting, cautious volume repletion with normal saline may return the ECFV to normal and restore renal function to baseline without having to intervene with dialysis.

Potassium Homeostasis In CKD, the decline in GFR is not necessarily accompanied by a parallel decline in urinary potassium excretion, which is predominantly mediated by aldosterone-dependent secretion in the distal nephron. Another defense against potassium retention in these patients is augmented potassium excretion in the GI tract. Notwithstanding these two homeostatic responses, hyperkalemia may be precipitated in certain settings. These include increased dietary potassium intake, protein catabolism, hemolysis, hemorrhage, transfusion of stored red blood cells, and metabolic acidosis. In addition, a host of medications can inhibit renal potassium excretion and lead to hyperkalemia. The most important medications in this respect include the RAS inhibitors and spironolactone and other potassium-sparing diuretics such as amiloride, eplerenone, and triamterene.

Certain causes of CKD can be associated with earlier and more severe disruption of potassium-secretory mechanisms in the distal nephron, out of proportion to the decline in GFR. These include conditions associated with hyporeninemic hypoaldosteronism, such as diabetes, and renal diseases that preferentially affect the distal nephron, such as obstructive uropathy and sickle cell nephropathy.

Hypokalemia is not common in CKD and usually reflects markedly reduced dietary potassium intake, especially in association with excessive diuretic therapy or concurrent GI losses. The use of potassium supplements and potassium-sparing diuretics may be risky in patients with impaired renal function, and should be constantly reevaluated as GFR declines.

Metabolic Acidosis Metabolic acidosis is a common disturbance in advanced CKD. The majority of patients can still acidify the urine, but they produce less ammonia and, therefore, cannot excrete the normal quantity of protons in combination with this urinary buffer. Hyperkalemia, if present, further depresses ammonia production. The combination of hyperkalemia and hyperchloremic metabolic acidosis is often present, even at earlier stages of CKD (stages 1–3), in patients with diabetic nephropathy or in those with predominant tubulointerstitial disease or obstructive uropathy; this is a non-anion-gap metabolic acidosis.

With worsening renal function, the total urinary net daily acid excretion is usually limited to 30–40 mmol, and the anions of retained organic acids can then lead to an anion-gap metabolic acidosis. Thus, the non-anion-gap metabolic acidosis that can be seen in earlier stages of CKD may be complicated by the addition of an anion-gap metabolic acidosis as CKD progresses. In most patients, the metabolic acidosis is mild; the pH is rarely <7.35 and can usually be corrected with oral sodium bicarbonate supplementation. Animal and human studies have suggested that even modest degrees of metabolic acidosis may be associated with the development of protein catabolism. Alkali supplementation may attenuate the catabolic state and possibly slow CKD progression and accordingly is recommended when the serum bicarbonate concentration falls below 20–23 mmol/L. The concomitant sodium load mandates careful attention to volume status and the need for diuretic agents.

Dietary salt restriction and the use of loop diuretics, occasionally in combination with metolazone, may be needed to maintain euolemia. In contrast, overzealous salt restriction or diuretic use can lead to ECFV depletion and precipitate a further decline in GFR. The rare patient with salt-losing nephropathy may require a sodium-rich diet or salt supplementation. Water restriction is indicated only if there is a problem with hyponatremia. Intractable ECFV expansion, despite dietary salt restriction and diuretic therapy, may be an indication to start renal replacement therapy. Hyperkalemia often responds to dietary restriction of potassium, the use of kaliuretic diuretics, and avoidance of both potassium supplements (including occult sources, such as dietary salt substitutes) and potassium-retaining medications (especially angiotensin-converting enzyme [ACE] inhibitors or angiotensin receptor blockers [ARBs]). Kaliuretic diuretics promote urinary potassium excretion, whereas potassium-binding resins, such as calcium resonium or sodium polystyrene, can promote potassium loss through the GI tract and may reduce the incidence of hyperkalemia. Intractable hyperkalemia is an indication (although uncommon) to consider institution of dialysis in a CKD patient. The renal tubular acidosis and subsequent anion-gap metabolic acidosis in progressive CKD will respond to alkali supplementation, typically with sodium bicarbonate. Recent studies suggest that this replacement should be considered when the serum bicarbonate concentration falls below 20–23 mmol/L to avoid the protein catabolic state seen with even mild degrees of metabolic acidosis and to slow the progression of CKD.

DISORDERS OF CALCIUM AND PHOSPHATE METABOLISM

The principal complications of abnormalities of calcium and phosphate metabolism in CKD occur in the skeleton and the vascular bed, with occasional severe involvement of extraosseous soft tissues. It is likely that disorders of bone turnover and disorders of vascular and soft tissue calcification are related to each other (Fig. 335-3).

Bone Manifestations of CKD The major disorders of bone disease can be classified into those associated with high bone turnover with increased PTH levels (including osteitis fibrosa cystica, the classic lesion of secondary hyperparathyroidism) and low bone turnover with low or normal PTH levels (adynamic bone disease and osteomalacia).

The pathophysiology of secondary hyperparathyroidism and the consequent high-turnover bone disease is related to abnormal mineral metabolism through the following events: (1) declining GFR leads to reduced excretion of phosphate and, thus, phosphate retention; (2) the retained phosphate stimulates increased synthesis of both FGF-23 by osteocytes and PTH and stimulates growth of parathyroid gland mass; and (3) decreased levels of ionized calcium, resulting from suppression of calcitriol production by FGF-23 and by the failing kidney, as well as phosphate retention, also stimulate PTH production. Low calcitriol levels contribute to hyperparathyroidism, both by leading to hypocalcemia and also by a direct effect on PTH gene transcription. These changes start to occur when the GFR falls below 60 mL/min.

FGF-23 is part of a family of phosphatonins that promotes renal phosphate excretion. Recent studies have shown that levels of this hormone, secreted by osteocytes, increase early in the course of CKD, even before phosphate retention and hyperphosphatemia. FGF-23 may defend normal serum phosphorus in at least three ways: (1) increased renal phosphate excretion; (2) stimulation of PTH, which also increases renal phosphate excretion; and (3) suppression of the formation of $1,25(\text{OH})_2\text{D}_3$, leading to diminished phosphorus absorption from the GI tract. Interestingly, high levels of FGF-23 are also an independent risk factor for left ventricular hypertrophy and mortality in CKD, dialysis, and renal transplant patients. Moreover, elevated levels of FGF-23 may indicate the need for therapeutic intervention (e.g., phosphate restriction), even when serum phosphate levels are within the normal range.



FIGURE 335-4 Tumoral calcinosis. This patient was on hemodialysis for many years and was nonadherent to dietary phosphorus restriction or the use of phosphate binders. He was chronically severely hyperphosphatemic. He developed an enlarging painful mass on his arm that was extensively calcified.

Hyperparathyroidism stimulates bone turnover and leads to *osteitis fibrosa cystica*. Bone histology shows abnormal osteoid, bone and bone marrow fibrosis, and in advanced stages, the formation of bone cysts, sometimes with hemorrhagic elements so that they appear brown in color, hence the term *brown tumor*. Clinical manifestations of severe hyperparathyroidism include bone pain and fragility, brown tumors, compression syndromes, and erythropoietin resistance in part related to the bone marrow fibrosis. Furthermore, PTH itself is considered a uremic toxin, and high levels are associated with muscle weakness, fibrosis of cardiac muscle, and nonspecific constitutional symptoms.

Low-turnover bone disease can be grouped into two categories—adynamic bone disease and osteomalacia. Adynamic bone disease is increasing in prevalence, especially among diabetics and the elderly. It is characterized by reduced bone volume and mineralization and may result from excessive suppression of PTH production, chronic inflammation, or both. Suppression of PTH can result from the use of vitamin D preparations or from excessive calcium exposure in the form of calcium-containing phosphate binders or high-calcium dialysis solutions. Complications of adynamic bone disease include an increased incidence of fracture and bone pain and an association with increased vascular and cardiac calcification. Occasionally the calcium will precipitate in the soft tissues into large concretions termed “tumoral calcinosis” (Fig. 335-4).

Calcium, Phosphorus, and the Cardiovascular System Recent epidemiologic evidence has shown a strong association between hyperphosphatemia and increased cardiovascular mortality rate in patients with stage 5 CKD and even in patients with earlier stages of CKD. Hyperphosphatemia and hypercalcemia are associated with increased vascular calcification, but it is unclear whether the excessive mortality rate is mediated by this mechanism. Studies using computed tomography (CT) and electron-beam CT scanning show that CKD patients have calcification of the media in coronary arteries and even heart valves that appear to be orders of magnitude greater than that in patients without renal disease. The magnitude of the calcification is proportional to age and hyperphosphatemia and is also associated with low PTH levels and low bone turnover. It is possible that in patients with advanced kidney disease, ingested calcium cannot be deposited in bones with low turnover and, therefore, is deposited at extraosseous sites, such as the vascular bed and soft tissues. It is interesting in this regard that there is also an association between osteoporosis and vascular calcification in the general population. Finally, hyperphosphatemia can induce a change in gene expression in vascular cells to an osteoblast-like profile, leading to vascular calcification and even ossification.



FIGURE 335-5 Calciphylaxis. This peritoneal dialysis patient was on chronic warfarin therapy for atrial fibrillation. She noticed a small painful nodule on the abdomen that was followed by progressive skin necrosis and ulceration of the anterior abdominal wall. She was treated with hyperbaric oxygen, intravenous thiosulfate, and discontinuation of warfarin, with slow resolution of the ulceration.

Other Complications of Abnormal Mineral Metabolism Calciphylaxis (calcific uremic arteriolopathy) is a devastating condition seen almost exclusively in patients with advanced CKD. It is heralded by livedo reticularis and advances to patches of ischemic necrosis, especially on the legs, thighs, abdomen, and breasts (Fig. 335-5). Pathologically, there is evidence of vascular occlusion in association with extensive vascular and soft tissue calcification. It appears that this condition is increasing in incidence. Originally it was ascribed to severe abnormalities in calcium and phosphorus control in dialysis patients, usually associated with advanced hyperparathyroidism. However, more recently, calciphylaxis has been seen with increasing frequency in the absence of severe hyperparathyroidism. Other etiologies have been suggested, including the increased use of oral calcium as a phosphate binder. Warfarin is commonly used in hemodialysis patients, and one of the effects of warfarin therapy is to decrease the vitamin K-dependent regeneration of matrix GLA protein. This latter protein is important in preventing vascular calcification. Thus, warfarin treatment is considered a risk factor for calciphylaxis, and if a patient develops this syndrome, this medication should be discontinued and replaced with alternative forms of anticoagulation.

TREATMENT

DISORDERS OF CALCIUM AND PHOSPHATE METABOLISM

The optimal management of secondary hyperparathyroidism and osteitis fibrosa is prevention. Once the parathyroid gland mass is very large, it is difficult to control the disease. Careful attention should be paid to the plasma phosphate concentration in CKD patients, who should be counseled on a low-phosphate diet as well as the appropriate use of phosphate-binding agents. These are agents that are taken with meals and complex the dietary phosphate to limit its GI absorption. Examples of phosphate binders are calcium acetate and calcium carbonate. A major side effect of calcium-based phosphate binders is calcium accumulation and hypercalcemia, especially in patients with low-turnover bone disease. Sevelamer and lanthanum are non-calcium-containing polymers that also function as phosphate binders; they do not predispose CKD patients to hypercalcemia and may attenuate calcium deposition in the vascular bed.

Calcitriol exerts a direct suppressive effect on PTH secretion and also indirectly suppresses PTH secretion by raising the concentration of ionized calcium. However, calcitriol therapy may result in hypercalcemia and/or hyperphosphatemia through increased GI absorption of these minerals. Certain analogues of calcitriol are available (e.g., paricalcitol) that suppress PTH secretion with less attendant hypercalcemia.

Recognition of the role of the extracellular calcium-sensing receptor has led to the development of calcimimetic agents that enhance the sensitivity of the parathyroid cell to the suppressive effect of calcium. This class of drug, which includes cinacalcet, produces a dose-dependent reduction in PTH and plasma calcium concentration in some patients.

Current National Kidney Foundation Kidney Disease Outcomes Quality Initiative guidelines recommend a target PTH level between 150 and 300 pg/mL, recognizing that very low PTH levels are associated with adynamic bone disease and possible consequences of fracture and ectopic calcification.

CARDIOVASCULAR ABNORMALITIES

Cardiovascular disease is the leading cause of morbidity and mortality in patients at every stage of CKD. The incremental risk of cardiovascular disease in those with CKD compared to the age- and sex-matched general population ranges from 10- to 200-fold, depending on the stage of CKD. Between 30 and 45% of patients reaching stage 5 CKD already have advanced cardiovascular complications. As a result, most patients with CKD succumb to cardiovascular disease (Fig. 335-6) before ever reaching stage 5 CKD. Thus, the focus of patient care in earlier CKD stages should be directed to prevention of cardiovascular complications.

Ischemic Vascular Disease The presence of any stage of CKD is a major risk factor for ischemic cardiovascular disease, including occlusive coronary, cerebrovascular, and peripheral vascular disease. The increased prevalence of vascular disease in CKD patients derives from both traditional (“classic”) and nontraditional (CKD-related) risk factors. Traditional risk factors include hypertension, hypervolemia, dyslipidemia, sympathetic overactivity, and hyperhomocysteinemia. The CKD-related risk factors comprise anemia, hyperphosphatemia, hyperparathyroidism, increased FGF-23, sleep apnea, and generalized inflammation. The inflammatory state associated with a reduction in kidney function is reflected in increased circulating acute-phase reactants, such as inflammatory cytokines and C-reactive protein, with a corresponding fall in the “negative acute-phase reactants,” such as serum albumin and fetuin. The inflammatory state appears to accelerate vascular occlusive disease, and low levels of fetuin may permit more rapid vascular calcification, especially in the face of hyperphosphatemia. Other abnormalities seen in CKD may augment

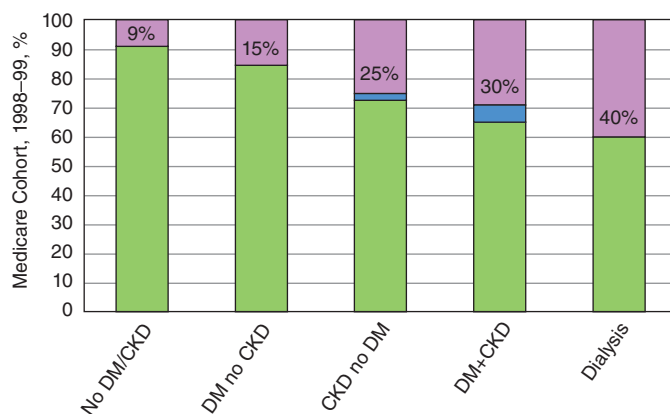


FIGURE 335-6 U.S. Renal Data System showing increased likelihood of dying rather than starting dialysis or reaching stage 5 chronic kidney disease (CKD). 1, Death; 2, ESRD; 3, event-free. DM; diabetes mellitus. (Adapted from RN Foley et al: *J Am Soc Nephrol* 16:489-495, 2005.)

TREATMENT **CARDIOVASCULAR ABNORMALITIES**

myocardial ischemia, including left ventricular hypertrophy and microvascular disease. In addition, hemodialysis, with its attendant episodes of hypotension and hypovolemia, may further aggravate coronary ischemia and repeatedly stun the myocardium. Interestingly, however, the largest increment in cardiovascular mortality rate in dialysis patients is not necessarily directly associated with documented acute myocardial infarction but, instead, presents with congestive heart failure and all of its manifestations and sudden death.

Cardiac troponin levels are frequently elevated in CKD without evidence of acute ischemia. The elevation complicates the diagnosis of acute myocardial infarction in this population. Serial measurements may be needed, and if the level is unchanged, it is possible that there is no acute myocardial ischemia. Therefore, the trend in levels over the hours after presentation may be more informative than a single, elevated level. Interestingly, consistently elevated levels are an independent prognostic factor for adverse cardiovascular events in this population.

Heart Failure Abnormal cardiac function secondary to myocardial ischemia, left ventricular hypertrophy, and frank cardiomyopathy, in combination with the salt and water retention that can be seen with CKD, often results in heart failure or even pulmonary edema. Heart failure can be a consequence of diastolic or systolic dysfunction, or both. A form of “low-pressure” pulmonary edema can also occur in advanced CKD, manifesting as shortness of breath and a “bat wing” distribution of alveolar edema fluid on the chest x-ray. This finding can occur even in the absence of ECFV overload and is associated with normal or mildly elevated pulmonary capillary wedge pressure. This process has been ascribed to increased permeability of alveolar capillary membranes as a manifestation of the uremic state, and it responds to dialysis. Other CKD-related risk factors, including anemia and sleep apnea, may contribute to the risk of heart failure.

Hypertension and Left Ventricular Hypertrophy Hypertension is one of the most common complications of CKD. It usually develops early during the course of CKD and is associated with adverse outcomes, including the development of ventricular hypertrophy and a more rapid loss of renal function. Many studies have shown a relationship between the level of blood pressure and the rate of progression of diabetic and nondiabetic kidney disease. Left ventricular hypertrophy and dilated cardiomyopathy are among the strongest risk factors for cardiovascular morbidity and mortality in patients with CKD and are thought to be related primarily, but not exclusively, to prolonged hypertension and ECFV overload. In addition, anemia and the placement of an arteriovenous fistula for hemodialysis can generate a high cardiac output state and consequent heart failure.

The absence of hypertension may signify poor left ventricular function. Indeed, in epidemiologic studies of dialysis patients, low blood pressure actually carries a worse prognosis than does high blood pressure. This mechanism, in part, accounts for the “reverse causation” seen in dialysis patients, wherein the presence of traditional risk factors, such as hypertension, hyperlipidemia, and obesity, appear to portend a better prognosis. Importantly, these observations derive from cross-sectional studies of late-stage CKD patients and should not be interpreted to discourage appropriate management of these risk factors in CKD patients, especially at early stages. In contrast to the general population, it is possible that in late-stage CKD, low blood pressure, reduced body mass index, and hypolipidemia indicate the presence of an advanced malnutrition-inflammation state, with poor prognosis.

The use of exogenous erythropoiesis-stimulating agents can increase blood pressure and the requirement for antihypertensive drugs. Chronic ECFV overload is also a contributor to hypertension, and improvement in blood pressure can often be seen with the use of dietary sodium restriction, diuretics, and fluid removal with dialysis. Nevertheless, because of activation of the RAS and other disturbances in the balance of vasoconstrictors and vasodilators, some patients remain hypertensive despite careful attention to ECFV status.

MANAGEMENT OF HYPERTENSION

The overarching goal of hypertension therapy in CKD is to prevent the extrarenal complications of high blood pressure, such as cardiovascular disease and stroke. Although a clear-cut generalizable benefit in slowing progression of CKD remains as yet unproven, the benefit for cardiac and neurologic health is compelling. In all patients with CKD, blood pressure should be controlled to levels recommended by national guideline panels. In CKD patients with diabetes or proteinuria >1 g per 24 h, blood pressure should be reduced to 130/80 mmHg, if achievable without prohibitive adverse effects. Salt restriction should be the first line of therapy. When volume management alone is not sufficient, the choice of antihypertensive agent is similar to that in the general population. ACE inhibitors and ARBs appear to slow the rate of decline of kidney function in a manner that extends beyond reduction of systemic arterial pressure and that involves correction of the intraglomerular hyperfiltration and hypertension involved in progression of CKD described above. Occasionally, introduction of ACE inhibitors and ARBs can actually precipitate an episode of acute kidney injury, especially when used in combination in patients with ischemic renovascular disease. The use of ACE inhibitors and ARBs may also be complicated by the development of hyperkalemia. Often the concomitant use of a kaliuretic diuretic, such as metolazone, can improve potassium excretion in addition to improving blood pressure control. Potassium-sparing diuretics should be used with caution or avoided altogether in most patients.

MANAGEMENT OF CARDIOVASCULAR DISEASE

There are many strategies available to treat the traditional and nontraditional risk factors in CKD patients. Although these have proved effective in the general population, there is little evidence for their benefit in patients with advanced CKD, especially those on dialysis. Certainly hypertension, elevated serum levels of homocysteine, and dyslipidemia promote atherosclerotic disease and are treatable complications of CKD. Renal disease complicated by nephrotic syndrome is associated with a very atherogenic lipid profile and hypercoagulability, which increases the risk of occlusive vascular disease. Because diabetes mellitus and hypertension are the two most frequent causes of advanced CKD, it is not surprising that cardiovascular disease is the most frequent cause of death in dialysis patients. The role of “inflammation” may be quantitatively more important in patients with kidney disease, and the treatment of more traditional risk factors may result in only modest success. However, modulation of traditional risk factors may be the only weapon in the therapeutic armamentarium for these patients until the nature of inflammation in CKD and its treatment are better understood.

Lifestyle changes, including regular exercise, should be advocated. Hyperlipidemia in patients with CKD should be managed according to national guidelines. If dietary measures are not sufficient, preferred lipid-lowering medications, such as statins, should be used. Again, the use of these agents has not been of proven benefit for patients with advanced CKD.

Pericardial Disease Chest pain with respiratory accentuation, accompanied by a friction rub, is diagnostic of pericarditis. Classic electrocardiographic abnormalities include PR-interval depression and diffuse ST-segment elevation. Pericarditis can be accompanied by pericardial effusion that is seen on echocardiography and can rarely lead to tamponade. However, the pericardial effusion can be asymptomatic, and pericarditis can be seen without significant effusion.

Pericarditis is observed in advanced uremia, and with the advent of timely initiation of dialysis, is not as common as it once was. It is now more often observed in underdialyzed, nonadherent patients than in those starting dialysis.

TREATMENT PERICARDIAL DISEASE

Uremic pericarditis is an absolute indication for the urgent initiation of dialysis or for intensification of the dialysis prescription in those already receiving dialysis. Because of the propensity to hemorrhage in pericardial fluid, hemodialysis should be performed without heparin. A pericardial drainage procedure should be considered in patients with recurrent pericardial effusion, especially with echocardiographic signs of impending tamponade. Nonuremic causes of pericarditis and effusion include viral, malignant, tuberculous, and autoimmune etiologies. It may also be seen after myocardial infarction and as a complication of treatment with the antihypertensive drug minoxidil.

HEMATOLOGIC ABNORMALITIES

Anemia A normocytic, normochromic anemia is observed as early as stage 3 CKD and is almost universal by stage 4. The primary cause in patients with CKD is insufficient production of erythropoietin (EPO) by the diseased kidneys. Additional factors are reviewed in [Table 335-3](#).

The anemia of CKD is associated with a number of adverse pathophysiologic consequences, including decreased tissue oxygen delivery and utilization, increased cardiac output, ventricular dilation, and ventricular hypertrophy. Clinical manifestations include fatigue and diminished exercise tolerance, angina, heart failure, decreased cognition and mental acuity, and impaired host defense against infection. In addition, anemia may play a role in growth restriction in children with CKD. Although many studies in CKD patients have found that anemia and resistance to exogenous erythropoietic-stimulating agents (ESA) are associated with a poor prognosis, the relative contribution to a poor outcome of the low hematocrit itself, versus inflammation as a cause of the anemia and ESA resistance, remains unclear.

TREATMENT ANEMIA

The availability of recombinant human ESA has been one of the most significant advances in the care of renal patients since the introduction of dialysis and renal transplantation. The routine use of these recombinant hormones has obviated the need for regular blood transfusions in severely anemic CKD patients, thus dramatically reducing the incidence of transfusion-associated infections and iron overload. Frequent blood transfusions in dialysis patients also lead to the development of alloantibodies that can sensitize the patient to donor kidney antigens and make renal transplantation more problematic.

Adequate bone marrow iron stores should be available before treatment with ESA is initiated. Iron supplementation is usually essential to ensure an optimal response to ESA in patients with CKD because the demand for iron by the marrow frequently exceeds the amount of iron that is immediately available for erythropoiesis (measured by percent transferrin saturation), as well as the amount in iron stores (measured by serum ferritin). For the CKD patient not yet on dialysis or the patient treated with peritoneal dialysis, oral iron supplementation should be attempted. If there is GI

intolerance, the patient may have to undergo IV iron infusion. For patients on hemodialysis, IV iron can be administered during dialysis, keeping in mind that iron therapy can increase the susceptibility to bacterial infections. In addition to iron, an adequate supply of other major substrates and cofactors for red cell production must be ensured, including vitamin B₁₂ and folate. Anemia resistant to recommended doses of ESA in the face of adequate iron stores may be due to some combination of the following: acute or chronic inflammation, inadequate dialysis, severe hyperparathyroidism, chronic blood loss or hemolysis, chronic infection, or malignancy. Blood transfusions increase the risk of hepatitis, iron overload, and transplant sensitization; they should be avoided unless the anemia fails to respond to ESA and the patient is symptomatic.

Randomized, controlled trials of ESA in CKD have failed to show an improvement in cardiovascular outcomes with this therapy. Indeed, there has been an indication that the use of ESA in CKD may be associated with an increased risk of stroke in those with type 2 diabetes, an increase in thromboembolic events, and perhaps a faster progression to the need for dialysis. Therefore, any benefit in terms of improvement of anemic symptoms needs to be balanced against the potential cardiovascular risk. Although further studies are needed, it is quite clear that complete normalization of the hemoglobin concentration has not been demonstrated to be of incremental benefit to CKD patients. Current practice is to target a hemoglobin concentration of 100–115 g/L.

Abnormal Hemostasis Patients with later stages of CKD may have a prolonged bleeding time, decreased activity of platelet factor III, abnormal platelet aggregation and adhesiveness, and impaired prothrombin consumption. Clinical manifestations include an increased tendency to bleeding and bruising, prolonged bleeding from surgical incisions, menorrhagia, and GI bleeding. Interestingly, CKD patients also have a greater susceptibility to thromboembolism, especially if they have renal disease that includes nephrotic-range proteinuria. The latter condition results in hypoalbuminemia and renal loss of anticoagulant factors, which can lead to a thrombophilic state.

TREATMENT ABNORMAL HEMOSTASIS

Abnormal bleeding time and coagulopathy in patients with renal failure may be reversed temporarily with desmopressin (DDAVP), cryoprecipitate, IV conjugated estrogens, blood transfusions, and ESA therapy. Optimal dialysis will usually correct a prolonged bleeding time.

Given the coexistence of bleeding disorders and a propensity to thrombosis that is unique in the CKD patient, decisions about anticoagulation that have a favorable risk-benefit profile in the general population may not be applicable to the patient with advanced CKD. One example is warfarin anticoagulation for atrial fibrillation; the decision to anticoagulate should be made on an individual basis in the CKD patient because there appears to be a greater risk of bleeding complications.

Certain anticoagulants, such as fractionated low-molecular-weight heparin, may need to be avoided or dose-adjusted in these patients, with monitoring of factor Xa activity where available. It is often more prudent to use conventional unfractionated heparin, titrated to the measured partial thromboplastin time, in hospitalized patients requiring an alternative to warfarin anticoagulation. The new classes of oral anticoagulants are all, in part, renally eliminated and need dose adjustment in the face of decreased GFR ([Chap. 143](#)).

NEUROMUSCULAR ABNORMALITIES

Central nervous system (CNS), peripheral, and autonomic neuropathy as well as abnormalities in muscle structure and function are all well-recognized complications of CKD. Subtle clinical manifestations of uremic neuromuscular disease usually become evident at stage 3 CKD. Early manifestations of CNS complications include mild disturbances in memory and concentration and sleep disturbance. Neuromuscular irritability, including hiccups, cramps, and twitching, becomes evident

TABLE 335-3 CAUSES OF ANEMIA IN CKD

Relative deficiency of erythropoietin
Diminished red blood cell survival
Bleeding diathesis
Iron deficiency
Hyperparathyroidism/bone marrow fibrosis
Chronic inflammation
Folate or vitamin B ₁₂ deficiency
Hemoglobinopathy
Comorbid conditions: hypo-/hyperthyroidism, pregnancy, HIV-associated disease, autoimmune disease, immunosuppressive drugs

at later stages. In advanced untreated kidney failure, asterixis, myoclonus, seizures, and coma can be seen.

Peripheral neuropathy usually becomes clinically evident after the patient reaches stage 4 CKD, although electrophysiologic and histologic evidence occurs earlier. Initially, sensory nerves are involved more than motor, lower extremities more than upper, and distal parts of the extremities more than proximal. The “restless leg syndrome” is characterized by ill-defined sensations of sometimes debilitating discomfort in the legs and feet relieved by frequent leg movement. If dialysis is not instituted soon after onset of sensory abnormalities, motor involvement follows, including muscle weakness. Evidence of peripheral neuropathy without another cause (e.g., diabetes mellitus) is an indication for starting renal replacement therapy. Many of the complications described above will resolve with dialysis, although subtle nonspecific abnormalities may persist.

GASTROINTESTINAL AND NUTRITIONAL ABNORMALITIES

Uremic fetor, a urine-like odor on the breath, derives from the breakdown of urea to ammonia in saliva and is often associated with an unpleasant metallic taste (dysgeusia). Gastritis, peptic disease, and mucosal ulcerations at any level of the GI tract occur in uremic patients and can lead to abdominal pain, nausea, vomiting, and GI bleeding. These patients are also prone to constipation, which can be worsened by the administration of calcium and iron supplements. The retention of uremic toxins also leads to anorexia, nausea, and vomiting.

Protein restriction may be useful to decrease nausea and vomiting; however, it may put the patient at risk for malnutrition and should be carried out, if possible, in consultation with a registered dietitian specializing in the management of CKD patients. Protein-energy malnutrition, a consequence of low protein and caloric intake, is common in advanced CKD and is often an indication for initiation of renal replacement therapy. Metabolic acidosis and the activation of inflammatory cytokines can promote protein catabolism. Assessment for protein-energy malnutrition should begin at stage 3 CKD. A number of indices are useful in this assessment and include dietary history, including food diary and subjective global assessment; edema-free body weight; and measurement of urinary protein nitrogen appearance. Dual-energy x-ray absorptiometry is now widely used to estimate lean body mass versus ECFV. Adjunctive tools include clinical signs, such as skinfold thickness, mid-arm muscle circumference, and additional laboratory tests such as serum pre-albumin and cholesterol levels. Nutritional guidelines for patients with CKD are summarized in the “Treatment” section.

ENDOCRINE-METABOLIC DISTURBANCES

Glucose metabolism is impaired in CKD, as evidenced by a slowing of the rate at which blood glucose levels decline after a glucose load. However, fasting blood glucose is usually normal or only slightly elevated, and the mild glucose intolerance does not require specific therapy. Because the kidney contributes to insulin removal from the circulation, plasma levels of insulin are slightly to moderately elevated in most uremic patients, both in the fasting and postprandial states. Because of this diminished renal degradation of insulin, patients on insulin therapy may need progressive reduction in dose as their renal function worsens. Many hypoglycemic agents, including the gliptins, require dose reduction in renal failure, and some, such as metformin, are contraindicated when the GFR is less than half of normal.

In women with CKD, estrogen levels are low, and menstrual abnormalities, infertility, and inability to carry pregnancies to term are common. When the GFR has declined to ~40 mL/min, pregnancy is associated with a high rate of spontaneous abortion, with only ~20% of pregnancies leading to live births, and pregnancy may hasten the progression of the kidney disease itself. Women with CKD who are contemplating pregnancy should consult first with a nephrologist in conjunction with an obstetrician specializing in high-risk pregnancy. Men with CKD have reduced plasma testosterone levels, and sexual dysfunction and oligospermia may supervene. Sexual maturation may be delayed or impaired in adolescent children with CKD, even among those treated with dialysis. Many of these

abnormalities improve or reverse with intensive dialysis or with successful renal transplantation.

DERMATOLOGIC ABNORMALITIES

Abnormalities of the skin are prevalent in progressive CKD. Pruritus is quite common and one of the most vexing manifestations of the uremic state. In advanced CKD, even on dialysis, patients may become more pigmented, and this is felt to reflect the deposition of retained pigmented metabolites, or *urochromes*. Although many of the cutaneous abnormalities improve with dialysis, pruritus is often tenacious. The first lines of management are to rule out unrelated skin disorders, such as scabies, and to treat hyperphosphatemia, which can cause itch. Local moisturizers, mild topical glucocorticoids, oral antihistamines, and ultraviolet radiation have been reported to be helpful.

A skin condition unique to CKD patients called *nephrogenic fibrosing dermatopathy* consists of progressive subcutaneous induration, especially on the arms and legs. The condition is similar to scleromyxedema and is seen very rarely in patients with CKD who have been exposed to the magnetic resonance contrast agent gadolinium. Current recommendations are that patients with CKD stage 3 (GFR 30–59 mL/min) should minimize exposure to gadolinium, and those with CKD stages 4–5 (GFR <30 mL/min) should avoid the use of gadolinium agents unless it is medically necessary. Concomitant liver disease appears to be a risk factor. However, no patient should be denied an imaging investigation that is critical to management, and under such circumstances, rapid removal of gadolinium by hemodialysis (even in patients not yet receiving renal replacement therapy) shortly after the procedure may mitigate this sometimes devastating complication.

EVALUATION AND MANAGEMENT OF PATIENTS WITH CKD

INITIAL APPROACH

History and Physical Examination Symptoms and overt signs of kidney disease are often subtle or absent until renal failure supervenes. Thus, the diagnosis of kidney disease often surprises patients and may be a cause of skepticism and denial. Particular aspects of the history that are germane to renal disease include a history of hypertension (which can cause CKD or more commonly be a consequence of CKD), diabetes mellitus, abnormal urinalyses, and problems with pregnancy such as preeclampsia or early pregnancy loss. A careful drug history should be elicited: patients may not volunteer use of analgesics, for example. Other drugs to consider include nonsteroidal anti-inflammatory agents, cyclooxygenase-2 (COX-2) inhibitors, antimicrobials, chemotherapeutic agents, antiretroviral agents, proton pump inhibitors, phosphate-containing bowel cathartics, and lithium. In evaluating the uremic syndrome, questions about appetite, weight loss, nausea, hiccups, peripheral edema, muscle cramps, pruritus, and restless legs are especially helpful. A careful family history of kidney disease, together with assessment of manifestations in other organ systems such as auditory, visual, and integumentary, may lead to the diagnosis of a heritable form of CKD (e.g., Alport or Fabry disease, cystinosis) or shared environmental exposure to nephrotoxic agents (e.g., heavy metals, aristolochic acid). It should be noted that clustering of CKD, sometimes of different etiologies, is often observed within families.

The physical examination should focus on blood pressure and target organ damage from hypertension. Thus, funduscopy and precordial examination (left ventricular heave, a fourth heart sound) should be carried out. Funduscopy is important in the diabetic patient, because it may show evidence of diabetic retinopathy, which is associated with nephropathy. Other physical examination manifestations of CKD include edema and sensory polyneuropathy. The finding of asterixis or a pericardial friction rub not attributable to other causes usually signifies the presence of the uremic syndrome.

Laboratory Investigation Laboratory studies should focus on a search for clues to an underlying causative or aggravating disease process and on the degree of renal damage and its consequences. Serum and urine protein electrophoresis, looking for multiple myeloma, should be obtained in all patients >35 years with unexplained CKD, especially if there is associated anemia and elevated, or even inappropriately

normal, serum calcium concentration in the face of renal insufficiency. In the presence of glomerulonephritis, autoimmune diseases such as lupus and underlying infectious etiologies such as hepatitis B and C and HIV should be tested. Serial measurements of renal function should be obtained to determine the pace of renal deterioration and ensure that the disease is truly chronic rather than acute or subacute and hence potentially reversible. Serum concentrations of calcium, phosphorus, vitamin D, and PTH should be measured to evaluate metabolic bone disease. Hemoglobin concentration, iron, vitamin B₁₂, and folate should also be evaluated. A 24-h urine collection may be helpful, because protein excretion >300 mg may be an indication for therapy with ACE inhibitors or ARBs.

Imaging Studies The most useful imaging study is a renal ultrasound, which can verify the presence of two kidneys, determine if they are symmetric, provide an estimate of kidney size, and rule out renal masses and evidence of obstruction. Because it takes time for kidneys to shrink as a result of chronic disease, the finding of bilaterally small kidneys supports the diagnosis of CKD of long-standing duration, with an irreversible component of scarring. If the kidney size is normal, it is possible that the renal disease is acute or subacute. The exceptions are diabetic nephropathy (where kidney size is increased at the onset of diabetic nephropathy before CKD supervenes), amyloidosis, and HIV nephropathy, where kidney size may be normal in the face of CKD. Polycystic kidney disease that has reached some degree of renal failure will almost always present with enlarged kidneys with multiple cysts (Chap. 339). A discrepancy >1 cm in kidney length suggests either a unilateral developmental abnormality or disease process or renovascular disease with arterial insufficiency affecting one kidney more than the other. The diagnosis of renovascular disease can be undertaken with different techniques, including Doppler sonography, nuclear medicine studies, or CT or magnetic resonance imaging (MRI) studies. If there is a suspicion of reflux nephropathy (recurrent childhood urinary tract infection, asymmetric renal size with scars on the renal poles), a voiding cystogram may be indicated. However, in most cases, by the time the patient has CKD, the reflux has resolved, and even if still present, repair does not improve renal function. Radiographic contrast imaging studies are not particularly helpful in the investigation of CKD. Intravenous or intraarterial dye should be avoided where possible in the CKD patient, especially with diabetic nephropathy, because of the risk of radiographic contrast dye-induced renal failure. When unavoidable, appropriate precautionary measures include avoidance of hypovolemia at the time of contrast exposure, minimization of the dye load, and choice of radiographic contrast preparations with the least nephrotoxic potential. Additional measures thought to attenuate contrast-induced worsening of renal function include judicious administration of sodium bicarbonate-containing solutions and *N*-acetylcysteine.

Kidney Biopsy In the patient with bilaterally small kidneys, renal biopsy is not advised because (1) it is technically difficult and has a greater likelihood of causing bleeding and other adverse consequences, (2) there is usually so much scarring that the underlying disease may not be apparent, and (3) the window of opportunity to render disease-specific therapy has passed. Other contraindications to renal biopsy include uncontrolled hypertension, active urinary tract infection, bleeding diathesis (including ongoing anticoagulation), and severe obesity. Ultrasound-guided percutaneous biopsy is the favored approach, but a surgical or laparoscopic approach can be considered, especially in the patient with a single kidney where direct visualization and control of bleeding are crucial. In the CKD patient in whom a kidney biopsy is indicated (e.g., suspicion of a concomitant or superimposed active process such as interstitial nephritis or in the face of accelerated loss of GFR), the bleeding time should be measured, and if increased, desmopressin should be administered immediately prior to the procedure.

A brief run of hemodialysis (without heparin) may also be considered prior to renal biopsy to normalize the bleeding time.

ESTABLISHING THE DIAGNOSIS AND ETIOLOGY OF CKD

The most important initial diagnostic step is to distinguish newly diagnosed CKD from acute or subacute renal failure, because the latter two

conditions may respond to targeted therapy. Previous measurements of serum creatinine concentration are particularly helpful in this regard. Normal values from recent months or even years suggest that the current extent of renal dysfunction could be more acute, and hence reversible, than might otherwise be appreciated. In contrast, elevated serum creatinine concentration in the past suggests that the renal disease represents a chronic process. Even if there is evidence of chronicity, there is the possibility of a superimposed acute process (e.g., ECFV depletion, urinary infection or obstruction, or nephrotoxin exposure) supervening on the chronic condition. If the history suggests multiple systemic manifestations of recent onset (e.g., fever, polyarthritis, rash), it should be assumed that renal insufficiency is part of an acute systemic illness.

Although renal biopsy can usually be performed in early CKD (stages 1–3), it is not always indicated. For example, in a patient with a history of type 1 diabetes mellitus for 15–20 years with retinopathy, nephrotic-range proteinuria, and absence of hematuria, the diagnosis of diabetic nephropathy is very likely and biopsy is usually not necessary. However, if there were some other finding not typical of diabetic nephropathy, such as hematuria or white blood cell casts, or absence of diabetic retinopathy, some other disease may be present and a biopsy may be indicated.

In the absence of a clinical diagnosis, renal biopsy may be the only recourse to establish an etiology in early-stage CKD. However, as noted above, once the CKD is advanced and the kidneys are small and scarred, there is little utility and significant risk in attempting to arrive at a specific diagnosis. Genetic testing is increasingly entering the repertoire of diagnostic tests, since the patterns of injury and kidney morphologic abnormalities often reflect overlapping causal mechanisms, whose origins can sometimes be attributed to a genetic predisposition or cause.

TREATMENT CHRONIC KIDNEY DISEASE

Treatments aimed at specific causes of CKD are discussed elsewhere. Among others, these include optimized glucose control in diabetes mellitus, immunosuppressive agents for glomerulonephritis, and emerging specific therapies to retard cystogenesis in polycystic kidney disease. The optimal timing of both specific and nonspecific therapy is usually well before there has been a measurable decline in GFR and certainly before CKD is established. It is helpful to measure sequentially and plot the rate of decline of GFR in all patients. Any acceleration in the rate of decline should prompt a search for superimposed acute or subacute processes that may be reversible. These include ECFV depletion, uncontrolled hypertension, urinary tract infection, new obstructive uropathy, exposure to nephrotoxic agents (such as nonsteroidal anti-inflammatory drugs [NSAIDs] or radiographic dye), and reactivation or flare of the original disease, such as lupus or vasculitis.

SLOWING THE PROGRESSION OF CKD

There is variation in the rate of decline of GFR among patients with CKD. However, the following interventions should be considered in an effort to stabilize or slow the decline of renal function.

Reducing Intraglomerular Hypertension and Proteinuria Increased intraglomerular filtration pressures and glomerular hypertrophy develop as a response to loss of nephron number from different kidney diseases. This response is maladaptive, as it promotes the ongoing decline of kidney function even if the inciting process has been treated or spontaneously resolved. Control of glomerular hypertension is important in slowing the progression of CKD. Moreover, elevated blood pressure increases proteinuria by increasing its flux across the glomerular capillaries. Conversely, the renoprotective effect of antihypertensive medications is gauged through the consequent reduction of proteinuria. Thus, the more effective a given treatment is in lowering protein excretion, the greater the subsequent impact on protection from decline in GFR. This observation is the basis for the treatment guideline establishing 130/80 mmHg as the target blood pressure in proteinuric CKD patients.

ACE inhibitors and ARBs inhibit the angiotensin-induced vasoconstriction of the efferent arterioles of the glomerular microcirculation. This inhibition leads to a reduction in both intraglomerular filtration pressure and proteinuria. Several controlled studies have shown that these drugs are effective in slowing the progression of renal failure in patients with advanced stages of both diabetic and nondiabetic CKD. This slowing in progression of CKD is strongly associated with the proteinuria-lowering effect. In the absence of an antiproteinuric response with either agent alone, combined treatment with both ACE inhibitors and ARBs has been considered. The combination is associated with a greater reduction in proteinuria compared to either agent alone. Insofar as reduction in proteinuria is a surrogate for improved renal outcome, the combination would appear to be advantageous. However, there is a greater incidence of acute kidney injury and adverse cardiac events from such combination therapy. It is uncertain, therefore, whether the ACE inhibitor plus ARB therapy can be advised routinely. Adverse effects from these agents include cough and angioedema with ACE inhibitors and anaphylaxis and hyperkalemia with either class. A progressive increase in serum creatinine concentration with these agents may suggest the presence of renovascular disease within the large or small arteries. Development of these side effects may mandate the use of second-line antihypertensive agents instead of the ACE inhibitors or ARBs. Among the calcium channel blockers, diltiazem and verapamil may exhibit superior antiproteinuric and renoprotective effects compared to the dihydropyridines. At least two different categories of response can be considered: one in which progression is strongly associated with systemic and intraglomerular hypertension and proteinuria (e.g., diabetic nephropathy, glomerular diseases) and in which ACE inhibitors and ARBs are likely to be the first choice; and another in which proteinuria is mild or absent initially (e.g., adult polycystic kidney disease and other tubulointerstitial diseases), where the contribution of intraglomerular hypertension is less prominent and other antihypertensive agents can be useful for control of systemic hypertension.

SLOWING THE PROGRESSION OF DIABETIC NEPHROPATHY

See Chap. 418.

MANAGING OTHER COMPLICATIONS OF CHRONIC KIDNEY DISEASE

Medication Dose Adjustment Although the loading dose of most drugs is not affected by CKD because no renal elimination is used in the calculation, the maintenance doses of many drugs will need to be adjusted. For those agents in which >70% excretion is by a nonrenal route, such as hepatic elimination, dose adjustment may not be needed. Some drugs that should be avoided include metformin, meperidine, and oral hypoglycemics that are eliminated by the kidney. NSAIDs should be avoided because of the risk of further worsening of kidney function. Many antibiotics, antihypertensives, and antiarrhythmics may require a reduction in dosage or change in the dose interval. Several online Web-based databases for dose adjustment of medications according to stage of CKD or estimated GFR are available (e.g., <http://www.globalrph.com/renaldosing2.htm>). Nephrotoxic radiocontrast agents and gadolinium should be avoided or used according to strict guidelines when medically necessary as described above.

PREPARATION FOR RENAL REPLACEMENT THERAPY (See also Chap. 337)

Temporary relief of symptoms and signs of impending uremia, such as anorexia, nausea, vomiting, lassitude, and pruritus, may sometimes be achieved with protein restriction. However, this carries a significant risk of malnutrition, and thus plans for more long-term management should be in place.

Maintenance dialysis and kidney transplantation have extended the lives of hundreds of thousands of patients with CKD worldwide. Clear indications for initiation of renal replacement therapy for patients with CKD include uremic pericarditis, encephalopathy, intractable muscle cramping, anorexia, and nausea not attributable to reversible causes such as peptic ulcer disease, evidence of malnutrition, and fluid and electrolyte abnormalities, principally hyperkalemia or ECFV overload, that are refractory to other measures.

Recommendations for the Optimal Time for Initiation of Renal Replacement Therapy

Because of the individual variability in the severity of uremic symptoms and renal function, it is ill-advised to assign an arbitrary urea nitrogen or creatinine level to the need to start dialysis. Moreover, patients may become accustomed to chronic uremia and deny symptoms, only to find that they feel better with dialysis and realize in retrospect how poorly they were feeling before its initiation.

Previous studies suggested that starting dialysis before the onset of severe symptoms and signs of uremia was associated with prolongation of survival. This led to the concept of “healthy” start and is congruent with the philosophy that it is better to keep patients feeling well all along rather than allowing them to become ill with uremia before trying to return them to better health with dialysis or transplantation. Although recent studies have not confirmed an association of early-start dialysis with improved patient survival, there may be merit in this approach for some patients. On a practical level, advanced preparation may help to avoid problems with the dialysis process itself (e.g., a poorly functioning fistula for hemodialysis or malfunctioning peritoneal dialysis catheter) and, thus, preempt the morbidity associated with resorting to the insertion of temporary hemodialysis access with its attendant risks of sepsis, bleeding, thrombosis, and association with accelerated mortality.

Patient Education Social, psychological, and physical preparation for the transition to renal replacement therapy and the choice of the optimal initial modality are best accomplished with a gradual approach involving a multidisciplinary team. Along with conservative measures discussed in the sections above, it is important to prepare patients with an intensive educational program, explaining the likelihood and timing of initiation of renal replacement therapy and the various forms of therapy available, and the option of nondialytic maximum conservative care. The more knowledgeable that patients are about hemodialysis (both in-center and home-based), peritoneal dialysis, and kidney transplantation, the easier and more appropriate will be their decisions. Patients who are provided with educational programs are more likely to choose home-based dialysis therapy. This approach is of societal benefit because home-based therapy is less expensive and is associated with improved quality of life. The educational programs should be commenced no later than stage 4 CKD so that the patient has sufficient time and cognitive function to learn the important concepts, make informed choices, and implement preparatory measures for renal replacement therapy.

Exploration of social support is also important. In those who may perform home dialysis or undergo preemptive renal transplantation, early education of family members for selection and preparation of a home dialysis helper or a biologically or emotionally related potential living kidney donor should occur long before the onset of symptomatic renal failure.

Kidney transplantation (Chap. 337) offers the best potential for complete rehabilitation, because dialysis replaces only a small fraction of the kidneys’ filtration function and none of the other renal functions, including endocrine and anti-inflammatory effects. Generally, kidney transplantation follows a period of dialysis treatment, although preemptive kidney transplantation (usually from a living donor) can be carried out if it is certain that the renal failure is irreversible.

IMPLICATIONS FOR GLOBAL HEALTH

In contrast to the natural decline and successful eradication of many devastating infectious diseases, there is rapid growth in the prevalence of metabolic and vascular disease in developing countries. Diabetes mellitus is becoming increasingly prevalent in these countries, perhaps due in part to change in dietary habits, diminished physical activity, and weight gain. Therefore, it follows that there will be a proportionate increase in vascular and renal disease. Health care agencies must plan for improved screening for early detection, prevention, and treatment plans in these nations and must start considering options for improved availability of renal replacement therapies.

336 Dialysis in the Treatment of Renal Failure

Kathleen D. Liu, Glenn M. Chertow

Dialysis may be required for the treatment of either acute or chronic kidney disease. The use of continuous renal replacement therapies (CRRTs) and slow low-efficiency dialysis (SLED) is specific to the management of acute renal failure and is discussed in [Chap. 334](#). These modalities are performed continuously (CRRT) or over 6–12 h per session (SLED), in contrast to the 3–4 h of an intermittent hemodialysis session. [Advantages and disadvantages of CRRT and SLED are discussed in Chap. 334.](#)

Peritoneal dialysis is rarely used in developed countries for the treatment of acute renal failure because of the increased risk of infection and (as will be discussed in more detail below) less efficient clearance per unit of time. The focus of this chapter will be on the use of peritoneal and hemodialysis for end-stage renal disease (ESRD).

With the widespread availability of dialysis, the lives of hundreds of thousands of patients with ESRD have been prolonged. In the United States alone, there are now approximately 615,000 patients with ESRD, the vast majority of whom require dialysis. The incidence rate for ESRD is 357 cases per million population per year. The incidence of ESRD is disproportionately higher in African Americans (940 per million population per year) as compared with white Americans (280 per million population per year). In the United States, the leading cause of ESRD is diabetes mellitus, currently accounting for nearly 45% of newly diagnosed cases of ESRD. Approximately 30% of patients have ESRD that has been attributed to hypertension, although it is unclear whether in these cases hypertension is the cause or a consequence of vascular disease or other unknown causes of kidney failure. Other prevalent causes of ESRD include glomerulonephritis, polycystic kidney disease, and obstructive uropathy.



Globally, mortality rates for patients with ESRD are lowest in Europe and Japan but very high in the developing world because of the limited availability of dialysis. In the United States, the mortality rate of patients on dialysis has decreased slightly but remains extremely high, with a 5-year survival rate of approximately 35–40%. Deaths are due mainly to cardiovascular diseases and infections (approximately 40 and 10% of deaths, respectively). Older age, male sex, nonblack race, diabetes mellitus, malnutrition, and underlying heart disease are important predictors of death.

TREATMENT OPTIONS FOR ESRD PATIENTS

Commonly accepted criteria for initiating patients on maintenance dialysis include the presence of uremic symptoms, the presence of hyperkalemia unresponsive to conservative measures, persistent extracellular volume expansion despite diuretic therapy, acidosis refractory to medical therapy, a bleeding diathesis, and a creatinine clearance or estimated glomerular filtration rate (GFR) below 10 mL/min per 1.73 m² (see [Chap. 335](#) for estimating equations). Timely referral to a nephrologist for advanced planning and creation of a dialysis access, education about ESRD treatment options, and management of the complications of advanced chronic kidney disease (CKD), including hypertension, anemia, acidosis, and secondary hyperparathyroidism, are advisable. Recent data have suggested that a sizable fraction of ESRD cases result following episodes of acute renal failure, particularly among persons with underlying CKD. Furthermore, there is no benefit to initiating dialysis preemptively at a GFR of 10–14 mL/min per 1.73 m² compared to initiating dialysis for symptoms of uremia.

In ESRD, treatment options include hemodialysis (in center or at home); peritoneal dialysis, as either continuous ambulatory peritoneal dialysis (CAPD) or continuous cyclic peritoneal dialysis (CCPD); or transplantation ([Chap. 337](#)). Although there are significant geographic variations and differences in practice patterns, hemodialysis remains the most common therapeutic modality for ESRD (>90% of

patients) in the United States. In contrast to hemodialysis, peritoneal dialysis is continuous, but much less efficient, in terms of solute clearance. Although no large-scale clinical trials have been completed comparing outcomes among patients randomized to either hemodialysis or peritoneal dialysis, outcomes associated with both therapies are similar in most reports, and the decision of which modality to select is often based on personal preferences and quality-of-life considerations.

HEMODIALYSIS

Hemodialysis relies on the principles of solute diffusion across a semipermeable membrane. Movement of metabolic waste products takes place down a concentration gradient from the circulation into the dialysate. The rate of diffusive transport increases in response to several factors, including the magnitude of the concentration gradient, the membrane surface area, and the mass transfer coefficient of the membrane. The latter is a function of the porosity and thickness of the membrane, the size of the solute molecule, and the conditions of flow on the two sides of the membrane. According to laws of diffusion, the larger the molecule, the slower is its rate of transfer across the membrane. A small molecule, such as urea (60 Da), undergoes substantial clearance, whereas a larger molecule, such as creatinine (113 Da), is cleared less efficiently. In addition to diffusive clearance, movement of waste products from the circulation into the dialysate may occur as a result of ultrafiltration. Convective clearance occurs because of solvent drag, with solutes being swept along with water across the semipermeable dialysis membrane.

THE DIALYZER

There are three essential components to hemodialysis: the dialyzer, the composition and delivery of the dialysate, and the blood delivery system ([Fig. 336-1](#)). The dialyzer is a plastic chamber with the ability to perfuse blood and dialysate compartments simultaneously at very high flow rates. The hollow-fiber dialyzer is the most common in use in the United States. These dialyzers are composed of bundles of capillary tubes through which blood circulates while dialysate travels on the outside of the fiber bundle. The majority of dialyzers now manufactured in the United States are “biocompatible” synthetic membranes derived from polysulfone or related compounds (versus older cellulose “bioincompatible” membranes that activated the complement cascade). The frequency of reprocessing and reuse of hemodialyzers and blood lines varies across the world. In general, as the cost of disposable supplies has decreased, their use has increased. Formaldehyde, peracetic acid–hydrogen peroxide, glutaraldehyde, and bleach have all been used as reprocessing agents.

DIALYSATE

The potassium concentration of dialysate may be varied from 0 to 4 mmol/L depending on the predialysis serum potassium concentration. The usual dialysate calcium concentration is 1.25 mmol/L (2.5 meq/L), although modification may be required in selected settings (e.g., higher dialysate calcium concentrations may be used in patients with hypocalcemia associated with secondary hyperparathyroidism or following parathyroidectomy). The usual dialysate sodium concentration is 136–140 mmol/L. In patients who frequently develop hypotension during their dialysis run, “sodium modeling” to counterbalance urea-related osmolar gradients is often used. With sodium modeling, the dialysate sodium concentration is gradually lowered from the range of 145–155 mmol/L to isotonic concentrations (136–140 mmol/L) near the end of the dialysis treatment, typically declining either in steps or in a linear or exponential fashion. Higher dialysate sodium concentrations and sodium modeling may predispose patients to positive sodium balance and increased thirst; thus, these strategies to ameliorate intradialytic hypotension may be undesirable in hypertensive patients or in patients with large interdialytic weight gains. Because patients are exposed to approximately 120 L of water during each dialysis treatment, water used for the dialysate is subjected to filtration, softening, deionization, and, ultimately, reverse osmosis to remove microbiologic contaminants and dissolved ions.

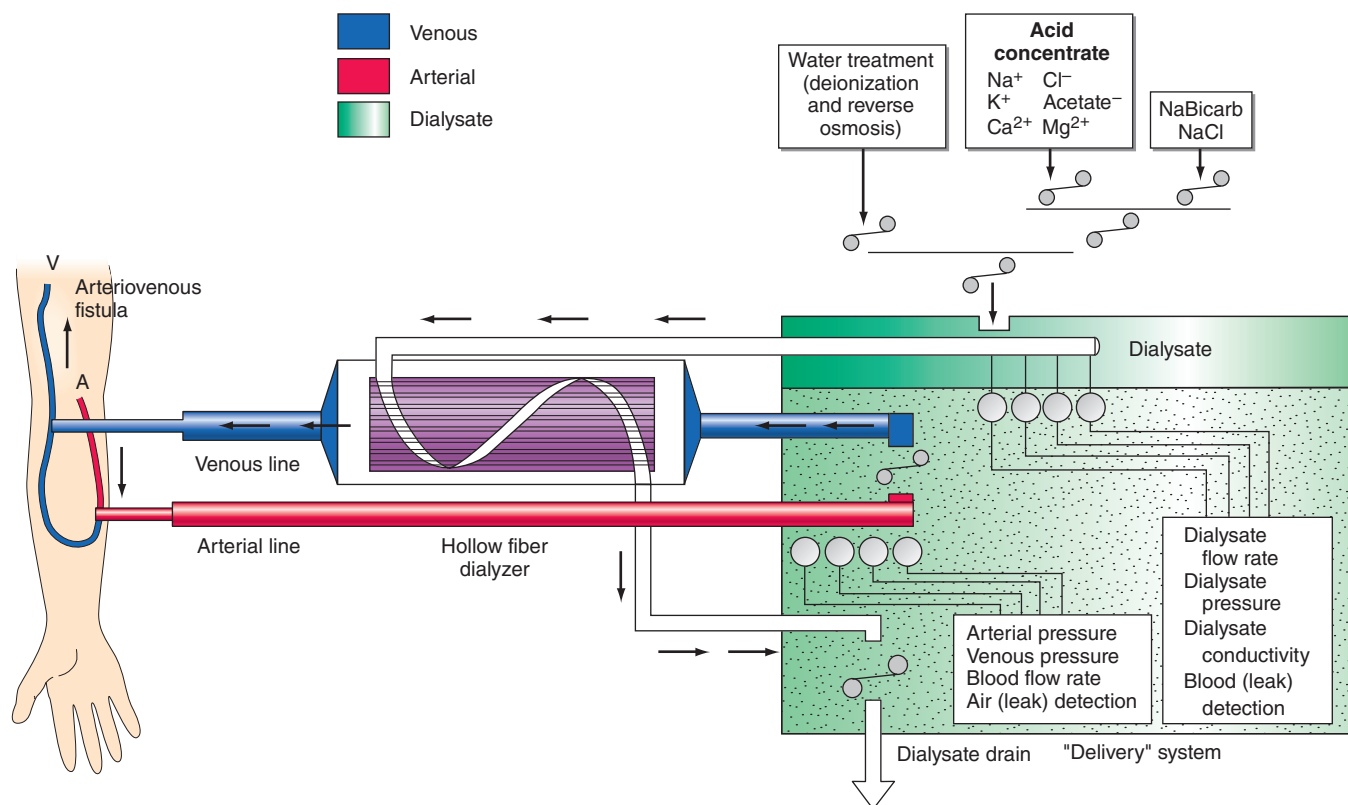


FIGURE 336-1 Schema for hemodialysis. A, artery; V, vein.

BLOOD DELIVERY SYSTEM

The blood delivery system is composed of the extracorporeal circuit and the dialysis access. The dialysis machine consists of a blood pump, dialysis solution delivery system, and various safety monitors. The blood pump moves blood from the access site, through the dialyzer, and back to the patient. The blood flow rate may range from 250–500 mL/min, depending on the type and integrity of the vascular access. Negative hydrostatic pressure on the dialysate side can be manipulated to achieve desirable fluid removal or *ultrafiltration*. Dialysis membranes have different ultrafiltration coefficients (i.e., mL removed/min per mmHg) so that along with hydrostatic changes, fluid removal can be varied. The dialysis solution delivery system dilutes the concentrated dialysate with water and monitors the temperature, conductivity, and flow of dialysate.

DIALYSIS ACCESS

The fistula, graft, or catheter through which blood is obtained for hemodialysis is often referred to as a *dialysis access*. A native fistula created by the anastomosis of an artery to a vein (e.g., the Brescia-Cimino fistula, in which the cephalic vein is anastomosed end-to-side to the radial artery) results in arterialization of the vein. This facilitates its subsequent use in the placement of large needles (typically 15 gauge) to access the circulation. Although fistulas have the highest long-term patency rate of all dialysis access options, fistulas are created in a minority of patients in the United States. Many patients undergo placement of an arteriovenous graft (i.e., the interposition of prosthetic material, usually polytetrafluoroethylene, between an artery and a vein) or a tunneled dialysis catheter. In recent years, nephrologists, vascular surgeons, and health care policy makers in the United States have encouraged creation of arteriovenous fistulas in a larger fraction of patients (the “fistula first” initiative). Unfortunately, even when created, arteriovenous fistulas may not mature sufficiently to provide reliable access to the circulation, or they may thrombose early in their development.

Grafts and catheters tend to be used among persons with smaller-caliber veins or persons whose veins have been damaged by repeated venipuncture, or after prolonged hospitalization. The most important complication of arteriovenous grafts is thrombosis of the graft and

graft failure, due principally to intimal hyperplasia at the anastomosis between the graft and recipient vein. When grafts (or fistulas) fail, catheter-guided angioplasty can be used to dilate stenoses; monitoring of venous pressures on dialysis and of access flow, although not routinely performed, may assist in the early recognition of impending vascular access failure. In addition to an increased rate of access failure, grafts and (in particular) catheters are associated with much higher rates of infection than fistulas.

Intravenous large-bore catheters are often used in patients with acute and chronic kidney disease. For persons on maintenance hemodialysis, tunneled catheters (either two separate catheters or a single catheter with two lumens) are often used when arteriovenous fistulas and grafts have failed or are not feasible due to anatomic considerations. These catheters are tunneled under the skin; the tunnel reduces bacterial translocation from the skin, resulting in a lower infection rate than with nontunneled temporary catheters. Most tunneled catheters are placed in the internal jugular veins; the external jugular, femoral, and subclavian veins may also be used.

Nephrologists, interventional radiologists, and vascular surgeons generally prefer to avoid placement of catheters into the subclavian veins; while flow rates are usually excellent, subclavian stenosis is a frequent complication and, if present, will likely prohibit permanent vascular access (i.e., a fistula or graft) in the ipsilateral extremity. Infection rates may be higher with femoral catheters. For patients with multiple vascular access complications and no other options for permanent vascular access, tunneled catheters may be the last “lifeline” for hemodialysis. Translumbar or transhepatic approaches into the inferior vena cava may be required if the superior vena cava or other central veins draining the upper extremities are stenosed or thrombosed.

GOALS OF DIALYSIS

The hemodialysis procedure consists of pumping heparinized blood through the dialyzer at a flow rate of 300–500 mL/min, while dialysate flows in an opposite *counter-current* direction at 500–800 mL/min. The efficiency of dialysis is determined by blood and dialysate flow through the dialyzer as well as dialyzer characteristics (i.e., its efficiency in removing solute). The *dose* of dialysis, which is currently defined as a derivation of the fractional urea clearance during a single

treatment, is further governed by patient size, residual kidney function, dietary protein intake, the degree of anabolism or catabolism, and the presence of comorbid conditions.

Since the landmark studies of Sargent and Gotch relating the measurement of the dose of dialysis using urea concentrations with morbidity in the National Cooperative Dialysis Study, the *delivered* dose of dialysis has been measured and considered as a quality assurance and improvement tool. Although the fractional removal of urea nitrogen and derivations thereof are considered to be the standard methods by which “adequacy of dialysis” is measured, a large multicenter randomized clinical trial (the HEMO Study) failed to show a difference in mortality associated with a large difference in urea clearance. Current targets include a urea reduction ratio (the fractional reduction in blood urea nitrogen per hemodialysis session) of >65–70% and a body water-indexed clearance \times time product (KT/V) above 1.2 or 1.05, depending on whether urea concentrations are “equilibrated.” For the majority of patients with ESRD, between 9 and 12 h of dialysis are required each week, usually divided into three equal sessions. Several studies have suggested that longer hemodialysis session lengths may be beneficial (independent of urea clearance), although these studies are confounded by a variety of patient characteristics, including body size and nutritional status. Hemodialysis “dose” should be individualized, and factors other than the urea nitrogen should be considered, including the adequacy of ultrafiltration or fluid removal and control of hyperkalemia, hyperphosphatemia, and metabolic acidosis. A recent randomized clinical trial (the Frequent Hemodialysis Network Trial) demonstrated improved control of hypertension and hyperphosphatemia, reduced left ventricular mass, and improved self-reported physical health with six times per week hemodialysis compared to the usual three times per week therapy. A companion trial in which frequent nocturnal hemodialysis was compared to conventional hemodialysis at home showed no significant effect on left ventricular mass or self-reported physical health. Finally, an evaluation of the U.S. Renal Data System registry showed a significant increase in mortality and hospitalization for heart failure after the longer interdialytic interval that occurs over the dialysis “weekend.”

COMPLICATIONS DURING HEMODIALYSIS

Hypotension is the most common acute complication of hemodialysis, particularly among patients with diabetes mellitus. Numerous factors appear to increase the risk of hypotension, including excessive ultrafiltration with inadequate compensatory vascular filling, impaired vasoactive or autonomic responses, osmolar shifts, overzealous use of antihypertensive agents, and reduced cardiac reserve. Patients with arteriovenous fistulas and grafts may develop high-output cardiac failure due to shunting of blood through the dialysis access; on rare occasions, this may necessitate ligation of the fistula or graft. Because of the vasodilatory and cardiodepressive effects of acetate, its use as the buffer in dialysate was once a common cause of hypotension. Since the introduction of bicarbonate-containing dialysate, dialysis-associated hypotension has become less common. The management of hypotension during dialysis consists of discontinuing ultrafiltration, the administration of 100–250 mL of isotonic saline or 10 mL of 23% saturated hypertonic saline, or administration of salt-poor albumin. Hypotension during dialysis can frequently be prevented by careful evaluation of the dry weight and by ultrafiltration modeling, such that more fluid is removed at the beginning rather than the end of the dialysis procedure. Additional maneuvers include the performance of sequential ultrafiltration followed by dialysis, cooling of the dialysate during dialysis treatment, and avoiding heavy meals during dialysis. Midodrine, an oral selective α_1 adrenergic agent, has been advocated by some practitioners, although there is insufficient evidence of its safety and efficacy to support its routine use.

Muscle cramps during dialysis are also a common complication. The etiology of dialysis-associated cramps remains obscure. Changes in muscle perfusion because of excessively rapid volume removal (e.g., >10–12 mL/kg per hour) or targeted removal below the patient’s estimated dry weight often precipitate dialysis-associated cramps. Strategies that may be used to prevent cramps include reducing volume removal during dialysis, ultrafiltration profiling, and the use of sodium modeling (see above).

Anaphylactoid reactions to the dialyzer, particularly on its first use, have been reported most frequently with the bioincompatible cellulosic-containing membranes. Dialyzer reactions can be divided into two types, A and B. Type A reactions are attributed to an IgE-mediated intermediate hypersensitivity reaction to ethylene oxide used in the sterilization of new dialyzers. This reaction typically occurs soon after the initiation of a treatment (within the first few minutes) and can progress to full-blown anaphylaxis if the therapy is not promptly discontinued. Treatment with steroids or epinephrine may be needed if symptoms are severe. The type B reaction consists of a symptom complex of nonspecific chest and back pain, which appears to result from complement activation and cytokine release. These symptoms typically occur several minutes into the dialysis run and typically resolve over time with continued dialysis.

PERITONEAL DIALYSIS

In peritoneal dialysis, 1.5–3 L of a dextrose-containing solution is infused into the peritoneal cavity and allowed to dwell for a set period of time, usually 2–4 h. As with hemodialysis, toxic materials are removed through a combination of convective clearance generated through ultrafiltration and diffusive clearance down a concentration gradient. The clearance of solutes and water during a peritoneal dialysis exchange depends on the balance between the movement of solute and water into the peritoneal cavity versus absorption from the peritoneal cavity. The rate of diffusion diminishes with time and eventually stops when equilibration between plasma and dialysate is reached. Absorption of solutes and water from the peritoneal cavity occurs across the peritoneal membrane into the peritoneal capillary circulation and via peritoneal lymphatics into the lymphatic circulation. The rate of peritoneal solute transport varies from patient to patient and may be altered by the presence of infection (peritonitis), drugs, and physical factors such as position and exercise.

FORMS OF PERITONEAL DIALYSIS

Peritoneal dialysis may be carried out as CAPD, CCPD, or a combination of both. In CAPD, dialysate is manually infused into the peritoneal cavity and exchanged three to five times during the day. A nighttime dwell is frequently instilled at bedtime and remains in the peritoneal cavity through the night. In CCPD, exchanges are performed in an automated fashion, usually at night; the patient is connected to an automatedycler that performs a series of exchange cycles while the patient sleeps. The number of exchange cycles required to optimize peritoneal solute clearance varies by the peritoneal membrane characteristics; as with hemodialysis, solute clearance should be tracked to ensure dialysis “adequacy.”

Peritoneal dialysis solutions are available in volumes typically ranging from 1.5 to 3 L. The major difference between the dialysate used for peritoneal dialysis rather than hemodialysis is that the hypertonicity of peritoneal dialysis solutions drives solute and fluid removal, whereas solute removal in hemodialysis depends on concentration gradients, and fluid removal requires transmembrane pressure. Typically, dextrose at varying concentrations contributes to the hypertonicity of peritoneal dialysate. Icodextrin is a nonabsorbable carbohydrate that can be used in place of dextrose. Studies have demonstrated more efficient ultrafiltration with icodextrin than with dextrose-containing solutions. Icodextrin is typically used as the “last fill” for patients on CCPD or for the longest dwell in patients on CAPD. The most common additives to peritoneal dialysis solutions are heparin to prevent obstruction of the dialysis catheter lumen with fibrin and antibiotics during an episode of acute peritonitis. Insulin may also be added in patients with diabetes mellitus.

ACCESS TO THE PERITONEAL CAVITY

Access to the peritoneal cavity is obtained through a peritoneal catheter. Catheters used for maintenance peritoneal dialysis are flexible, being made of silicone rubber with numerous side holes at the distal end. These catheters usually have two Dacron cuffs. The scarring that occurs around the cuffs anchors the catheter and seals it from bacteria tracking from the skin surface into the peritoneal cavity; it also prevents the

external leakage of fluid from the peritoneal cavity. The cuffs are placed in the preperitoneal plane and ~2 cm from the skin surface.

The *peritoneal equilibrium test* is a formal evaluation of peritoneal membrane characteristics that measures the transfer rates of creatinine and glucose across the peritoneal membrane. Patients are classified as low, low-average, high-average, and high transporters. Patients with rapid equilibration (i.e., high transporters) tend to absorb more glucose and lose efficiency of ultrafiltration with long daytime dwells. High transporters also tend to lose larger quantities of albumin and other proteins across the peritoneal membrane. In general, patients with rapid transporting characteristics require more frequent, shorter dwell time exchanges, nearly always obligating use of a cycler. Slower (low and low-average) transporters tend to do well with fewer exchanges. The efficiency of solute clearance also depends on the volume of dialysate infused. Larger volumes allow for greater solute clearance, particularly with CAPD in patients with low and low-average transport characteristics.

As with hemodialysis, the optimal dose of peritoneal dialysis is unknown. Several observational studies have suggested that higher rates of urea and creatinine clearance (the latter generally measured in liters per week) are associated with lower mortality rates and fewer uremic complications. However, a randomized clinical trial (Adequacy of Peritoneal Dialysis in Mexico [ADEMEX]) failed to show a significant reduction in mortality or complications with a relatively large increment in urea clearance. In general, patients on peritoneal dialysis do well when they retain residual kidney function. The rates of technique failure increase with years on dialysis and have been correlated with loss of residual function to a greater extent than loss of peritoneal membrane capacity. For some patients in whom CCPD does not provide sufficient solute clearance, a hybrid approach can be adopted where one or more daytime exchanges are added to the CCPD regimen. Although this approach can enhance solute clearance and prolong a patient's capacity to remain on peritoneal dialysis, the burden of the hybrid approach can be overwhelming.

COMPLICATIONS DURING PERITONEAL DIALYSIS

The major complications of peritoneal dialysis are peritonitis, catheter-associated nonperitonitis infections, weight gain and other metabolic disturbances, and residual uremia (especially among patients with no residual kidney function).

Peritonitis typically develops when there has been a break in sterile technique during one or more of the exchange procedures. Peritonitis is usually defined by an elevated peritoneal fluid leukocyte count (100/ μ L, of which at least 50% are polymorphonuclear neutrophils); these cutoffs are lower than in spontaneous bacterial peritonitis because of the presence of dextrose in peritoneal dialysis solutions and rapid bacterial proliferation in this environment without antibiotic therapy. The clinical presentation typically consists of pain and cloudy dialysate, often with fever and other constitutional symptoms. The most common culprit organisms are gram-positive cocci, including *Staphylococcus*, reflecting the origin from the skin. Gram-negative rod infections are less common; fungal and mycobacterial infections can be seen in selected patients, particularly after antibacterial therapy. Most cases of peritonitis can be managed either with intraperitoneal or oral antibiotics, depending on the organism; many patients with peritonitis do not require hospitalization. In cases where peritonitis is due to hydrophilic gram-negative rods (e.g., *Pseudomonas* sp.) or yeast, antimicrobial therapy is usually not sufficient, and catheter removal is required to ensure complete eradication of infection. Nonperitonitis catheter-associated infections (often termed *tunnel infections*) vary widely in severity. Some cases can be managed with local antibiotic or silver nitrate administration, whereas others are severe enough to require peritoneal antibiotic therapy and catheter removal.

Peritoneal dialysis is associated with a variety of metabolic complications. Albumin and other proteins can be lost across the peritoneal membrane in concert with the loss of metabolic wastes. Hypoproteinemia obligates a higher dietary protein intake in order to maintain nitrogen balance. Hyperglycemia and weight gain are also common complications of peritoneal dialysis. Several hundred calories in the form

of dextrose are absorbed each day, depending on the concentration employed. Peritoneal dialysis patients, particularly those with diabetes mellitus, are then prone to other complications of insulin resistance, including hypertriglyceridemia. On the positive side, the continuous nature of peritoneal dialysis usually allows for a more liberal diet, due to continuous removal of potassium and phosphorus—two major dietary components whose accumulation can be hazardous in ESRD.

LONG-TERM OUTCOMES IN ESRD

Cardiovascular disease constitutes the major cause of death in patients with ESRD. Cardiovascular mortality and event rates are higher in dialysis patients than in patients after transplantation, although rates are extraordinarily high in both populations. The underlying cause of cardiovascular disease is unclear but may be related to shared risk factors (e.g., diabetes mellitus, hypertension, atherosclerotic and arteriosclerotic vascular disease), chronic inflammation, massive changes in extracellular volume (especially with high interdialytic weight gains), inadequate treatment of hypertension, dyslipidemia, anemia, dystrophic vascular calcification, hyperhomocysteinemia, and, perhaps, alterations in cardiovascular dynamics during the dialysis treatment. Few studies have targeted cardiovascular risk reduction in ESRD patients; none have demonstrated consistent benefit. Two clinical trials of statin agents in ESRD demonstrated significant reductions in low-density lipoprotein (LDL) cholesterol concentrations, but no significant reductions in death or cardiovascular events (Die Deutsche Diabetes Dialyse Studie [4D] and A Study to Evaluate the Use of Rosuvastatin in Subjects on Regular Hemodialysis [AURORA]). The Study of Heart and Renal Protection (SHARP), which included patients on dialysis- and non-dialysis-requiring CKD, showed a 17% reduction in the rate of major cardiovascular events or cardiovascular death with simvastatin-ezetimibe treatment. Most experts recommend conventional cardioprotective strategies (e.g., lipid-lowering agents, aspirin, inhibitors of the renin-angiotensin-aldosterone system, and β -adrenergic antagonists) in dialysis patients based on the patients' cardiovascular risk profile, which appears to be increased by more than an order of magnitude relative to persons unaffected by kidney disease. Other complications of ESRD include a high incidence of infection, progressive debility and frailty, protein-energy malnutrition, and impaired cognitive function.

GLOBAL PERSPECTIVE



The incidence of ESRD is increasing worldwide with longer life expectancies and improved care of infectious and cardiovascular diseases. The management of ESRD varies widely by country and within country by region, and it is influenced by economic and other major factors. In general, peritoneal dialysis is more commonly performed in poorer countries owing to its lower expense and the high cost of establishing in-center hemodialysis units.

337 Transplantation in the Treatment of Renal Failure

Jamil Azzi, Edgar L. Milford, Mohamed H. Sayegh, Anil Chandraker

Transplantation of the human kidney is the treatment of choice for advanced chronic renal failure. Worldwide, tens of thousands of these procedures have been performed with more than 180,000 patients bearing functioning kidney transplants in the United States today. When azathioprine and prednisone initially were used as immunosuppressive drugs in the 1960s, the results with properly matched familial donors were superior to those with organs from deceased donors: 75–90% compared with 50–60% graft survival rates at 1 year. During the 1970s and 1980s, the success rate at the 1-year mark for deceased-donor

TABLE 337-1 DEFINITION OF AN EXPANDED CRITERIA DONOR AND A NON-HEART-BEATING DONOR (DONATION AFTER CARDIAC DEATH)

Expanded Criteria Donor (ECD)
Deceased donor >60 years
Deceased donor >50 years and hypertension and creatinine >1.5 mg/dL
Deceased donor >50 years and hypertension and death caused by cerebrovascular accident (CVA)
Deceased donor >50 years and death caused by CVA and creatinine >1.5 mg/dL
Donation after Cardiac Death ^a (DCD)
I: Brought in dead
II: Unsuccessful resuscitation
III: Awaiting cardiac arrest
IV: Cardiac arrest after brainstem death
V: Cardiac arrest in a hospital patient

^aKidneys can be used for transplantation from categories II–V but are commonly only used from categories III and IV. The survival of these kidneys has not been shown to be inferior to that of deceased-donor kidneys.

Note: Kidneys can be both ECD and DCD. ECD kidneys have been shown to have a poorer survival, and there is a separate shorter waiting list for ECD kidneys. They are generally used for patients for whom the benefits of being transplanted earlier outweigh the associated risks of using an ECD kidney.

transplants rose progressively. Currently, deceased-donor grafts have a 92% 1-year survival and living-donor grafts have a 96% 1-year survival. Although there has been improvement in long-term survival, it has not been as impressive as the short-term survival, and currently the “average” ($t_{1/2}$) life expectancy of a living-donor graft is around 20 years and that of a deceased-donor graft is close to 14 years.

Mortality rates after transplantation are highest in the first year and are age-related: 2% for ages 18–34 years, 3% for ages 35–49 years, and 6.8% for ages ≥50–60 years. These rates compare favorably with those in the chronic dialysis population even after risk adjustments for age, diabetes, and cardiovascular status. Although the loss of kidney transplant due to acute rejection is currently rare, most allografts succumb at varying rates to a chronic process consisting of interstitial fibrosis, tubular atrophy, vasculopathy, and glomerulopathy, the pathogenesis of which is incompletely understood. Overall, transplantation returns most patients to an improved lifestyle and an improved life expectancy compared with patients on dialysis.

RECENT ACTIVITY AND RESULTS

In 2011, there were more than 11,835 deceased-donor kidney transplants and 5772 living-donor transplants in the United States, with the ratio of deceased to living donors remaining stable over the last few years. The backlog of patients with end-stage renal disease (ESRD) has been increasing every year, and it always lags behind the number of available donors. As the number of patients with end-stage kidney disease increases, the demand for kidney transplants continues to increase. In 2011, there were 55,371 active adult candidates on the waiting list, and less than 18,000 patients were transplanted. This imbalance is set to worsen over the coming years with the predicted increased rates of obesity and diabetes worldwide. In an attempt to increase utilization of deceased-donor kidneys and reduce discard rates of organs, criteria for the use of so-called expanded criteria donor (ECD) kidneys and kidneys from donors after cardiac death (DCD) have been developed (**Table 337-1**). ECD

kidneys are usually used for older patients who are expected to fare less well on dialysis.

The overall results of transplantation are presented in **Table 337-2** as the survival of grafts and of patients. At the 1-year mark, graft survival is higher for living-donor recipients, most likely because those grafts are not subject to as much ischemic injury. The more effective drugs now in use for immunosuppression have almost equalized the risk of graft rejection in all patients for the first year. At 5 and 10 years, however, there has been a steeper decline in survival of those with deceased-donor kidneys.

RECIPIENT SELECTION

There are few absolute contraindications to renal transplantation. The transplant procedure is relatively noninvasive, as the organ is placed in the inguinal fossa without entering the peritoneal cavity. Recipients without perioperative complications often can be discharged from the hospital in excellent condition within 5 days of the operation.

Virtually all patients with ESRD who receive a transplant have a higher life expectancy than do risk-matched patients who remain on dialysis. Even though diabetic patients and older candidates have a higher mortality rate than other transplant recipients, their survival is improved with transplantation compared with those remaining on dialysis. This global benefit of transplantation as a treatment modality poses substantial ethical issues for policy makers, as the number of deceased kidneys available is far from sufficient to meet the current needs of the candidates. The current standard of care is that the candidate should have a life expectancy of >5 years to be put on a deceased organ wait list. Even for living donation, the candidate should have >5 years of life expectancy. This standard has been established because the benefits of kidney transplantation over dialysis are realized only after a perioperative period in which the mortality rate is higher in transplanted patients than in dialysis patients with comparable risk profiles.

All candidates must have a thorough risk-versus-benefit evaluation before being approved for transplantation. In particular, an aggressive approach to diagnosis of correctable coronary artery disease, presence of latent or indolent infection (HIV, hepatitis B or C, tuberculosis), and neoplasm should be a routine part of the candidate workup. Most transplant centers consider overt AIDS and active hepatitis absolute contraindications to transplantation because of the high risk of opportunistic infection. Some centers are now transplanting individuals with hepatitis and even HIV infection under strict protocols to determine whether the risks and benefits favor transplantation over dialysis.

Among the few absolute “immunologic” contraindications to transplantation is the presence of antibodies against the donor kidney at the time of the anticipated transplant that can cause hyperacute rejection. Those harmful antibodies include natural antibodies against the ABO blood group antigens and antibodies against human leukocyte antigen (HLA) class I (A, B, C) or class II (DR) antigens. These antibodies are routinely excluded by proper screening of the candidate’s ABO compatibility and direct cytotoxic cross-matching of candidate serum with lymphocytes of the donor.

TISSUE TYPING AND CLINICAL IMMUNOGENETICS

Matching for antigens of the HLA major histocompatibility gene complex (**Chap. 373e**) is an important criterion for selection of donors for renal allografts. Each mammalian species has a single chromosomal

TABLE 337-2 MEAN RATES OF GRAFT AND PATIENT SURVIVAL FOR KIDNEYS TRANSPLANTED IN THE UNITED STATES FROM 1998 TO 2008^a

	1-Year Follow-Up		5-Year Follow-Up		10-Year Follow-Up	
	Grafts, %	Patients, %	Grafts, %	Patients, %	Grafts, %	Patients, %
Deceased donor	92	96	72	84	46	64
Living donor	96	99	81	91	59	77

^aAll patients transplanted are included, and the follow-up unadjusted survival data from the 1-, 5-, and 10-year periods are presented to show the attrition rates over time within the two types of organ donors.

Source: Data from Summary Tables, 2009 Annual Reports, Scientific Registry of Transplant Recipients.

region that encodes the strong, or major, transplantation antigens, and this region on the human sixth chromosome is called *HLA*. HLA antigens have been classically defined by serologic techniques, but methods to define specific nucleotide sequences in genomic DNA are increasingly being used. Other “minor” antigens may play crucial roles, in addition to the ABH(O) blood groups and endothelial antigens that are not shared with lymphocytes. The Rh system is not expressed on graft tissue. Evidence for designation of HLA as the genetic region that encodes major transplantation antigens comes from the success rate in living related donor renal and bone marrow transplantation, with superior results in HLA-identical sibling pairs. Nevertheless, 5% of HLA-identical renal allografts are rejected, often within the first weeks after transplantation. These failures represent states of prior sensitization to non-HLA antigens. Non-HLA minor antigens are relatively weak when initially encountered and are, therefore, suppressible by conventional immunosuppressive therapy. Once priming has occurred, however, secondary responses are much more refractory to treatment.

DONOR SELECTION

Donors can be deceased or volunteer living donors. When first-degree relatives are donors, graft survival rates at 1 year are 5–7% greater than those for deceased-donor grafts. The 5-year survival rates still favor a partially matched (3/6 HLA mismatched) family donor over a randomly selected cadaver donor. In addition, living donors provide the advantage of immediate availability. For both living and deceased donors, the 5-year outcomes are poor if there is a complete (6/6) HLA mismatch.

The survival rate of living unrelated renal allografts is as high as that of perfectly HLA-matched cadaver renal transplants and comparable to that of kidneys from living relatives. This outcome is probably a consequence of both short cold ischemia time and the extra care taken to document that the condition and renal function of the donor are optimal before proceeding with a living unrelated donation. It is illegal in the United States to purchase organs for transplantation.

Living volunteer donors should be cleared of any medical conditions that may cause morbidity and mortality after kidney transplantation. Concern has been expressed about the potential risk to a volunteer kidney donor of premature renal failure after several years of increased blood flow and hyperfiltration per nephron in the remaining kidney. There are a few reports of the development of hypertension, proteinuria, and even lesions of focal segmental sclerosis in donors over long-term follow-up. It is also desirable to consider the risk of development of type 1 diabetes mellitus in a family member who is a potential donor to a diabetic renal failure patient. Anti-insulin and anti-islet cell antibodies should be measured and glucose tolerance tests should be performed in such donors to exclude a prediabetic state. Selective renal arteriography should be performed on donors to rule out the presence of multiple or abnormal renal arteries, because the surgical procedure is difficult and the ischemic time of the transplanted kidney is long when there are vascular abnormalities. Transplant surgeons are now using a laparoscopic method to isolate and remove the living donor's kidney. This operation has the advantage of less evident surgical scars, and, because there is less tissue trauma, the laparoscopic donors have a substantially shorter hospital stay and less discomfort than those who have the traditional surgery.

Deceased donors should be free of malignant neoplastic disease, hepatitis, and HIV due to possible transmission to the recipient, although there is increasing interest in using hepatitis C- and HIV-positive organs in previously infected recipients. Increased risk of graft failure exists when the donor is elderly or has renal failure and when the kidney has a prolonged period of ischemia and storage.

In the United States, there is a coordinated national system of regulations, allocation support, and outcomes analysis for kidney transplantation called the Organ Procurement Transplant Network. It is now possible to remove deceased-donor kidneys and maintain them for up to 48 h on cold pulsatile perfusion or with simple flushing

and cooling. This approach permits adequate time for typing, cross-matching, transportation, and selection problems to be solved.

PRESENSITIZATION

A positive cytotoxic cross-match of recipient serum with donor T lymphocytes indicates the presence of preformed donor-specific anti-HLA class I antibodies and is usually predictive of an acute vasculitic event termed *hyperacute rejection*. This finding, along with ABO incompatibility, represents the only absolute immunologic contraindication for kidney transplantation. Recently, more tissue typing laboratories have shifted to a flow cytometric–based cross-match assay, which detects the presence of anti-HLA antibodies that are not necessarily detected on a cytotoxic cross-match assay and may not be an absolute contraindication to transplantation. The known sources of such sensitization are blood transfusion, a prior transplant, pregnancy, and vaccination/infection. Patients sustained by dialysis often show fluctuating antibody titers and specificity patterns. At the time of assignment of a cadaveric kidney, cross-matches are performed with at least a current serum. Previously analyzed antibody specificities and additional cross-matches are performed accordingly. Flow cytometry detects binding of anti-HLA antibodies of a candidate's serum by a recipient's lymphocytes. This highly sensitive test can be useful for avoidance of accelerated, and often untreatable, early graft rejection in patients receiving second or third transplants.

For the purposes of cross-matching, donor T lymphocytes, which express class I but not class II antigens, are used as targets for detection of anti-class I (HLA-A and -B) antibodies that are expressed on all nucleated cells. Preformed anti-class II (HLA-DR and -DQ) antibodies against the donor also carry a higher risk of graft loss, particularly in recipients who have suffered early loss of a prior kidney transplant. B lymphocytes, which express both class I and class II antigens, are used as targets in these assays.

Some non-HLA antigens restricted in expression to endothelium and monocytes have been described, but clinical relevance is not well established. A series of minor histocompatibility antigens do not elicit antibodies, and sensitization to these antigens is detectable only by cytotoxic T cells, an assay too cumbersome for routine use.

Desensitization before transplantation by reducing the level of anti-donor antibodies using plasmapheresis and administration of pooled immunoglobulin, or both, has been useful in reducing the risk of hyperacute rejection following transplantation.

IMMUNOLOGY OF REJECTION

Both cellular and humoral (antibody-mediated) effector mechanisms can play roles in kidney transplant rejection.

Cellular rejection is mediated by lymphocytes that respond to HLA antigens expressed within the organ. The CD4+ lymphocyte responds to class II (HLA-DR) incompatibility by proliferating and releasing pro-inflammatory cytokines that augment the proliferative response of the immune system. CD8+ cytotoxic lymphocyte precursors respond primarily to class I (HLA-A, -B) antigens and mature into cytotoxic effector cells that cause organ damage through direct contact and lysis of donor target cells. Full T cell activation requires not only T cell receptor binding to the alloantigens presented by self or donor HLA molecules (indirect and direct presentation, respectively), but also engaging costimulatory molecules such as CD28 on T cells and CD80 and CD86 ligands on antigen-presenting cells (Fig. 337-1). Signaling through both of these pathways induces activation of the kinase activity of calcineurin, which, in turn, activates transcription factors, leading to upregulation of multiple genes, including interleukin 2 (IL-2) and interferon gamma. IL-2 signals through the target of rapamycin (TOR) to induce cell proliferation in an autocrine fashion. There is evidence that non-HLA antigens can also play a role in renal transplant rejection episodes. Recipients who receive a kidney from an HLA-identical sibling can have rejection episodes and require maintenance immunosuppression, whereas identical twin transplants require no immunosuppression. There are documented non-HLA antigens, such as an endothelial-specific antigen system with limited polymorphism and a tubular antigen, which can act as targets of humoral or cellular rejection responses, respectively.

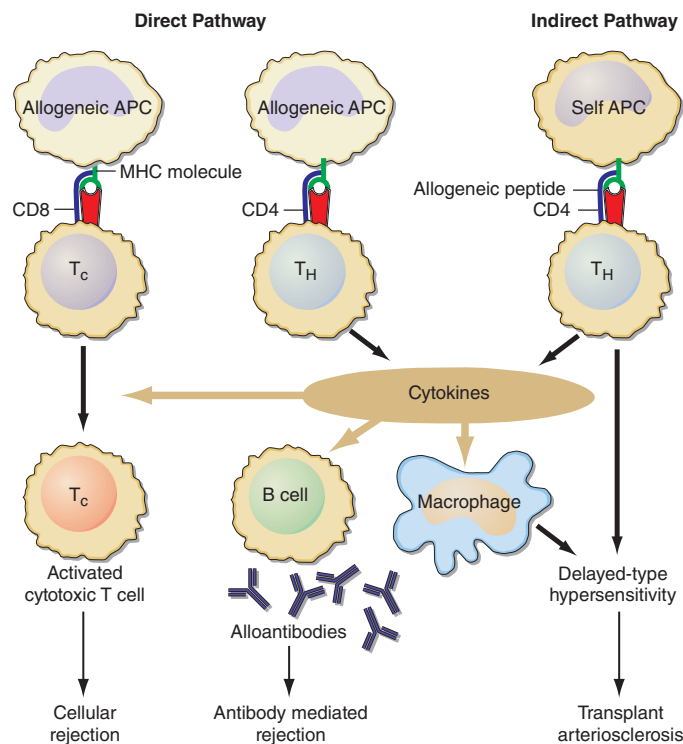


FIGURE 337-1 Recognition pathways for major histocompatibility complex (MHC) antigens. Graft rejection is initiated by CD4 helper T lymphocytes (T_H) having antigen receptors that bind to specific complexes of peptides and MHC class II molecules on antigen-presenting cells (APC). In transplantation, in contrast to other immunologic responses, there are two sets of T cell clones involved in rejection. In the direct pathway, the class II MHC of donor allogeneic APCs is recognized by CD4 T_H cells that bind to the intact MHC molecule, and class I MHC allogeneic cells are recognized by CD8 T cells. The latter generally proliferate into cytotoxic cells (T_C). In the indirect pathway, the incompatible MHC molecules are processed into peptides that are presented by the self-APCs of the recipient. The indirect, but not the direct, pathway is the normal physiologic process in T cell recognition of foreign antigens. Once T_H cells are activated, they proliferate and, by secretion of cytokines and direct contact, exert strong helper effects on macrophages, T_C , and B cells. (From MH Sayegh, LH Turka: *N Engl J Med*, 338:1813, 1998. Copyright 1998, Massachusetts Medical Society. All rights reserved.)

IMMUNOSUPPRESSIVE TREATMENT

Immunosuppressive therapy, as currently available, generally suppresses all immune responses, including those to bacteria, fungi, and even malignant tumors. In general, all clinically useful drugs are more selective to primary than to memory immune responses. Agents to suppress the immune response are classically divided into induction and maintenance agents and will be discussed in the following paragraphs. Those currently in clinical use are listed in [Table 337-3](#).

INDUCTION THERAPY

Induction therapy is currently given to most kidney transplant recipients in the United States at the time of transplant to reduce the risk of early acute rejection and to minimize or eliminate the use of either steroids or calcineurin inhibitors and their associated toxicities. Induction therapy consists of antibodies that could be monoclonal or polyclonal and depletion or nondepletional.

Depleting Agents Peripheral human lymphocytes, thymocytes, or lymphocytes from spleens or thoracic duct fistulas are injected into horses, rabbits, or goats to produce antilymphocyte serum, from which the globulin fraction is then separated, resulting in antithymocyte globulin. Those polyclonal antibodies induce lymphocyte depletion, and the immune system may take several months to recover.

Monoclonal antibodies against defined lymphocyte subsets offer a more precise and standardized form of therapy. Alemtuzumab is directed to the CD52 protein, widely distributed on immune cells such as B and T cells, natural killer cells, macrophages, and some granulocytes.

Nondepleting Agents Another approach to more selective therapy is to target the 55-kDa alpha chain of the IL-2 receptor, which is expressed only on T cells that have been recently activated. This approach is used as prophylaxis for acute rejection in the immediate posttransplant period and is effective at decreasing the early acute rejection rate with few adverse side effects.

The next step in the evolution of this therapeutic strategy, which has already been achieved in the short term in small numbers of immunologically well-matched patients, is the elimination of all maintenance immunosuppression therapy.

MAINTENANCE THERAPY

All kidney transplant recipients should receive maintenance immunosuppressive therapies except identical twins. The most frequently used combination is triple therapy with prednisone, a calcineurin inhibitor, and an antimetabolite; mammalian TOR (mTOR) inhibitors can replace one of the last two agents. More recently, the U.S. Food and Drug Administration (FDA) approved a new costimulatory blocking antibody, belatacept, as a new strategy to prevent long-term calcineurin inhibitor toxicity.

Antimetabolites *Azathioprine*, an analogue of mercaptopurine, was for two decades the keystone to immunosuppressive therapy in humans, but has given way to more effective agents. This agent can inhibit synthesis of DNA, RNA, or both. Azathioprine is administered in doses of 1.5–2 mg/kg per day. Reduction in the dose is required because of leukopenia and occasionally thrombocytopenia. Excessive amounts of azathioprine may also cause jaundice, anemia, and alopecia. If it is essential to administer allopurinol concurrently, the azathioprine dose must be reduced. Because inhibition of xanthine oxidase delays degradation, this combination is best avoided.

Mycophenolate mofetil or *mycophenolate sodium*, both of which are metabolized to mycophenolic acid, is now used in place of azathioprine in most centers. It has a similar mode of action and a mild degree of gastrointestinal toxicity but produces less bone marrow suppression. Its advantage is its increased potency in preventing or reversing rejection.

Steroids *Glucocorticoids* are important adjuncts to immunosuppressive therapy. Among all the agents employed, prednisone has effects that are easiest to assess, and in large doses it is usually effective for the reversal of rejection. In general, 200–300 mg prednisone is given immediately before or at the time of transplantation, and the dose is reduced to 30 mg within a week. The side effects of the glucocorticoids, particularly impairment of wound healing and predisposition to infection, make it desirable to taper the dose as rapidly as possible in the immediate postoperative period. Many centers now have protocols for early discontinuance or avoidance of steroids because of long-term adverse effects on bone, skin, and glucose metabolism. For treatment of acute rejection, methylprednisolone, 0.5–1 g IV, is administered immediately upon diagnosis of beginning rejection and continued once daily for 3 days. Such “pulse” doses are not effective in chronic rejection. Most patients whose renal function is stable after 6 months or a year do not require large doses of prednisone; maintenance doses of 5–10 mg/d are the rule. A major effect of steroids is preventing the release of IL-6 and IL-1 by monocytes-macrophages.

Calcineurin Inhibitors *Cyclosporine* is a fungal peptide with potent immunosuppressive activity. It acts on the calcineurin pathway to block transcription of mRNA for IL-2 and other proinflammatory cytokines, thereby inhibiting T cell proliferation. Although it works alone, cyclosporine is more effective in conjunction with glucocorticoids and mycophenolate. Clinical results with tens of thousands of renal transplants have been impressive. Among its toxic effects (nephrotoxicity, hepatotoxicity, hirsutism, tremor, gingival hyperplasia,

TABLE 337-3 MAINTENANCE IMMUNOSUPPRESSIVE DRUGS

Agent	Pharmacology	Mechanisms	Side Effects
Glucocorticoids	Increased bioavailability with hypoalbuminemia and liver disease; prednisone/prednisolone generally used	Binds cytosolic receptors and heat shock proteins. Blocks transcription of IL-1, -2, -3, -6, TNF- α , and IFN- γ	Hypertension, glucose intolerance, dyslipidemia, osteoporosis
Cyclosporine (CsA)	Lipid-soluble polypeptide, variable absorption, microemulsion more predictable	Trimolecular complex with cyclophilin and calcineurin \rightarrow block in cytokine (e.g., IL-2) production; however, stimulates TGF- β production	Nephrotoxicity, hypertension, dyslipidemia, glucose intolerance, hirsutism/hyperplasia of gums
Tacrolimus	Macrolide, well absorbed	Trimolecular complex with FKBP-12 and calcineurin \rightarrow block in cytokine (e.g., IL-2) production; may stimulate TGF- β production	Similar to CsA, but hirsutism/hyperplasia of gums unusual and diabetes more likely
Azathioprine	Mercaptopurine analogue	Hepatic metabolites inhibit purine synthesis	Marrow suppression (WBC > RBC > platelets)
Mycophenolate mofetil/sodium	Metabolized to mycophenolic acid	Inhibits purine synthesis via inosine monophosphate dehydrogenase	Diarrhea/cramps; dose-related liver and marrow suppression is uncommon
Sirolimus/everolimus	Macrolide, poor oral bioavailability	Complexes with FKBP-12 and then blocks p70 S6 kinase in the IL-2 receptor pathway for proliferation	Hyperlipidemia, thrombocytopenia
Belatacept	Fusion protein, intravenous injections	Binds CD80 and CD86, prevents CD28 binding and T cell activation	Posttransplant lymphoproliferative disease

Abbreviations: FKBP-12, FK506 binding protein 12; IFN, interferon; IL, interleukin; RBC, red blood cells; TGF, transforming growth factor; TNF, tumor necrosis factor; WBC, white blood cells.

diabetes), only nephrotoxicity presents a serious management problem and is further discussed below.

Tacrolimus (previously called FK506) is a fungal macrolide that has the same mode of action as cyclosporine as well as a similar side effect profile; it does not, however, produce hirsutism or gingival hyperplasia. De novo diabetes mellitus is more common with tacrolimus. The drug was first used in liver transplantation and may substitute for cyclosporine entirely or as an alternative in renal patients whose rejections are poorly controlled by cyclosporine.

mTOR Inhibitors **Sirolimus** (previously called rapamycin) is another fungal macrolide but has a different mode of action; i.e., it inhibits T cell growth factor signaling pathways, preventing the response to IL-2 and other cytokines. Sirolimus can be used in conjunction with cyclosporine or tacrolimus, or with mycophenolic acid, to avoid the use of calcineurin inhibitors.

Everolimus is another mTOR inhibitor with similar mechanism of action as sirolimus but with better bioavailability.

Belatacept **Belatacept** is a fusion protein that binds costimulatory ligands (CD80 and CD86) present on antigen-presenting cells, interrupting their binding to CD28 on T cells. This inhibition leads to T cell anergy and apoptosis. **Belatacept** is FDA approved for kidney transplant recipients and is given monthly as an intravenous infusion.

CLINICAL COURSE AND MANAGEMENT OF THE RECIPIENT

Adequate hemodialysis should be performed within 48 h of surgery, and care should be taken that the serum potassium level is not markedly elevated so that intraoperative cardiac arrhythmias can be averted. The diuresis that commonly occurs postoperatively must be carefully monitored. In some instances, it may be massive, reflecting the inability of ischemic tubules to regulate sodium and water excretion; with large diureses, massive potassium losses may occur. Most chronically uremic patients have some excess of extracellular fluid, and it is useful to maintain an expanded fluid volume in the immediate postoperative period. Acute tubular necrosis (ATN) due to ischemia may cause immediate oliguria or may follow an initial short period of graft function. Recovery usually occurs within 3 weeks, although periods as long as 6 weeks have been reported. Superimposition of rejection on ATN is common, and the differential diagnosis may be difficult without a graft biopsy. Cyclosporine therapy prolongs ATN, and some patients do not diurese until the dose is reduced drastically. Many centers avoid starting cyclosporine for the first several days, using antilymphocyte globulin (ALG) or a monoclonal antibody along with mycophenolic acid and prednisone until renal function is established. **Figure 337-2** illustrates an algorithm followed by many transplant centers for early posttransplant management of recipients at high or low risk of early renal dysfunction.

THE REJECTION EPISODE

Early diagnosis of rejection allows prompt institution of therapy to preserve renal function and prevent irreversible damage. Clinical evidence of rejection is rarely characterized by fever, swelling, and tenderness over the allograft. Rejection may present only with a rise in serum creatinine, with or without a reduction in urine volume. The focus should be on ruling out other causes of functional deterioration.

Doppler ultrasonography may be useful in ascertaining changes in the renal vasculature and in renal blood flow. Thrombosis of the renal vein occurs rarely; it may be reversible if it is caused by technical factors and intervention is prompt. Diagnostic ultrasound is the procedure of choice to rule out urinary obstruction or to confirm the presence of perirenal collections of urine, blood, or lymph. A rise in the serum creatinine level is a late marker of rejection, but it may be the only sign. Novel biomarkers are needed for early noninvasive detection of allograft rejection.

Calcineurin inhibitors (cyclosporine and tacrolimus) have an afferent arteriolar constrictor effect on the kidney and may produce permanent vascular and interstitial injury after sustained high-dose therapy. This action will lead to a deterioration in renal function difficult to distinguish from rejection without a renal biopsy. Interstitial fibrosis, isometric tubular vacuolization, and thickening of arteriolar walls are suggestive of this side effect, but not diagnostic. Hence, if no rejection is detected on the biopsy, serum creatinine may respond to a reduction in dose. However, if rejection activity is present in the biopsy, appropriate therapy is indicated. The first rejection episode is usually treated with IV administration of methylprednisolone, 500–1000 mg daily for 3 days. Failure to respond is an indication for antibody therapy, usually with antithymocyte globulin.

Evidence of antibody-mediated injury is present when endothelial injury and deposition of complement component c4d is detected by fluorescence labeling. This is usually accompanied by detection of the antibody in the recipient blood. The prognosis is poor, and aggressive use of plasmapheresis, immunoglobulin infusions, anti-CD20 monoclonal antibody (rituximab) to target B lymphocytes, bortezomib to target antibody-producing plasma cells, and eculizumab to inhibit complement is indicated.

MANAGEMENT PROBLEMS

The typical times after transplantation when the most common opportunistic infections occur are shown in **Table 337-4**. Prophylaxis for cytomegalovirus (CMV) and *Pneumocystis jiroveci* pneumonia is given for 6–12 months after transplantation.

The signs and symptoms of infection may be masked or distorted. Fever without obvious cause is common, and only after days or weeks may it become apparent that it has a viral or fungal origin. Bacterial infections are most common during the first month after transplantation.

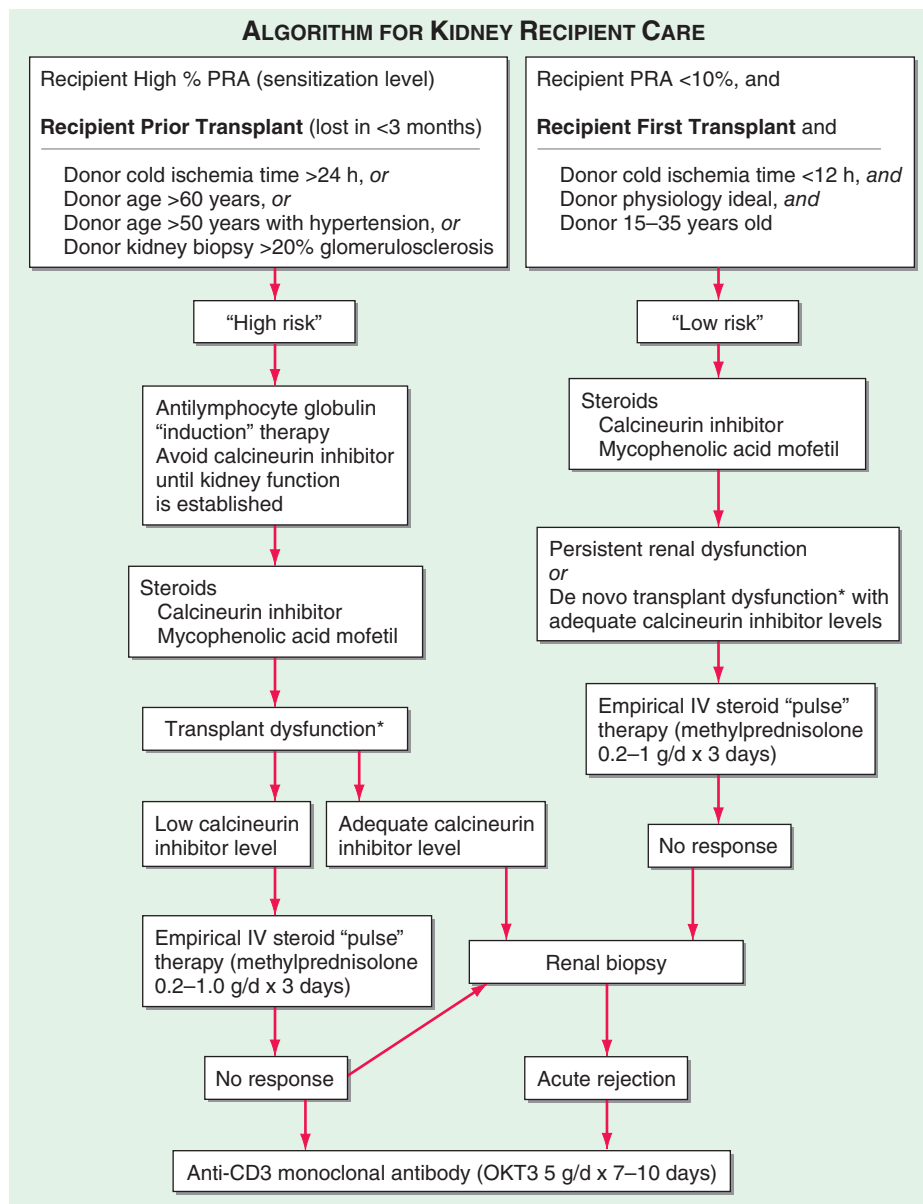


FIGURE 337-2 A typical algorithm for early posttransplant care of a kidney recipient. If any of the recipient or donor “high-risk” factors exist, more aggressive management is called for. Low-risk patients can be treated with a standard immunosuppressive regimen. Patients at higher risk of rejection or early ischemic and nephrotoxic transplant dysfunction are often induced with an antilymphocyte globulin to provide more potent early immunosuppression or to spare calcineurin nephrotoxicity. *When there is early transplant dysfunction, prerenal, obstructive, and vascular causes must be ruled out by ultrasonographic examination. The panel reactive antibody (PRA) is a quantitation of how much antibody is present in a candidate against a panel of cells representing the distribution of antigens in the donor pool.

TABLE 337-4 THE MOST COMMON OPPORTUNISTIC INFECTIONS IN RENAL TRANSPLANT RECIPIENTS

Peritransplant (<1 month)	Late (>6 months)
Wound infections	<i>Aspergillus</i>
Herpesvirus	<i>Nocardia</i>
Oral candidiasis	BK virus (polyoma)
Urinary tract infection	Herpes zoster
Early (1–6 months)	Hepatitis B
<i>Pneumocystis jiroveci</i>	Hepatitis C
Cytomegalovirus	
<i>Legionella</i>	
<i>Listeria</i>	
Hepatitis B	
Hepatitis C	

The importance of blood cultures in such patients cannot be overemphasized because systemic infection without obvious foci is common. Particularly ominous are rapidly occurring pulmonary lesions, which may result in death within 5 days of onset. When these lesions become apparent, immunosuppressive agents should be discontinued, except for maintenance doses of prednisone.

Aggressive diagnostic procedures, including transbronchial and open-lung biopsy, are frequently indicated. In the case of *P. jiroveci* (Chap. 244) infection, trimethoprim-sulfamethoxazole (TMP-SMX) is the treatment of choice; amphotericin B has been used effectively in systemic fungal infections. Prophylaxis against *P. jiroveci* with daily or alternate-day low-dose TMP-SMX is very effective. Involvement of the oropharynx with *Candida* (Chap. 240) may be treated with local nystatin. Tissue-invasive fungal infections require treatment with systemic agents such as fluconazole. Small doses (a total of 300 mg) of amphotericin given over a period of 2 weeks may be effective in fungal infections refractory to fluconazole. Macrolide antibiotics, especially

ketoconazole and erythromycin, and some calcium channel blockers (diltiazem, verapamil) compete with calcineurin inhibitors for P450 catabolism and cause elevated levels of these immunosuppressive drugs. Analeptics, such as phenytoin and carbamazepine, will increase catabolism to result in low levels. *Aspergillus* (Chap. 241), *Nocardia* (Chap. 199), and especially CMV (Chap. 219) infections also occur.

CMV is a common and dangerous DNA virus in transplant recipients. It does not generally appear until the end of the first post-transplant month. Active CMV infection is sometimes associated, or occasionally confused, with rejection episodes. Patients at highest risk for severe CMV disease are those without anti-CMV antibodies who receive a graft from a CMV antibody-positive donor (15% mortality). Valganciclovir is a cost-effective and bioavailable oral form of ganciclovir that has been proved effective in both prophylaxis and treatment of CMV disease. Early diagnosis in a febrile patient with clinical suspicion of CMV disease can be made by determining CMV viral load in the blood. A rise in IgM antibodies to CMV is also diagnostic. Culture of CMV from blood may be less sensitive. Tissue invasion of CMV is common in the gastrointestinal tract and lungs. CMV retinopathy occurs late in the course, if untreated. Treatment of active CMV disease with valganciclovir is always indicated. In many patients immune to CMV, viral activation can occur with major immunosuppressive regimens.

The polyoma group (BK, JC, SV40) is another class of DNA viruses that can become dormant in kidneys and can be activated by immunosuppression. When reactivation occurs with BK, there is a 50% chance of progressive fibrosis and loss of the graft within 1 year by the activated virus. Risk of infection is associated with the overall degree of immunosuppression rather than the individual immunosuppressive drugs used. Renal biopsy is necessary for the diagnosis. There have been variable results with leflunomide, cidofovir, and quinolone antibiotics (which are effective against polyoma helicase), but it is most important to reduce the immunosuppressive load.

The complications of glucocorticoid therapy are well known and include gastrointestinal bleeding, impairment of wound healing, osteoporosis, diabetes mellitus, cataract formation, and hemorrhagic pancreatitis. The treatment of unexplained jaundice in transplant patients should include cessation or reduction of immunosuppressive drugs if hepatitis or drug toxicity is suspected. Therapy in such circumstances often does not result in rejection of a graft, at least for several weeks. Acyclovir is effective in therapy for herpes simplex virus infections.

CHRONIC LESIONS OF THE TRANSPLANTED KIDNEY

Although 1-year transplant survival is excellent, most recipients experience progressive decline in kidney function over time thereafter. Chronic renal transplant dysfunction can be caused by recurrent disease, hypertension, cyclosporine or tacrolimus nephrotoxicity, chronic immunologic rejection, secondary focal glomerulosclerosis, or a combination of these pathophysiologies. Chronic vascular changes with intimal proliferation and medial hypertrophy are commonly found. Control of systemic and intrarenal hypertension with angiotensin-converting enzyme (ACE) inhibitors is thought to have a beneficial influence on the rate of progression of chronic renal transplant dysfunction. Renal biopsy can distinguish subacute cellular rejection from recurrent disease or secondary focal sclerosis.

MALIGNANCY

The incidence of tumors in patients on immunosuppressive therapy is 5–6%, or approximately 100 times greater than that in the general population in the same age range. The most common lesions are cancer of the skin and lips and carcinoma in situ of the cervix, as well as lymphomas such as non-Hodgkin's lymphoma. The risks are increased in proportion to the total immunosuppressive load administered and the time elapsed since transplantation. Surveillance for skin and cervical cancers is necessary.

OTHER COMPLICATIONS

Both chronic dialysis and renal transplant patients have a higher incidence of death from myocardial infarction and stroke than does the

population at large, and this is particularly true in diabetic patients. Contributing factors are the use of glucocorticoids and sirolimus and hypertension. Recipients of renal transplants have a high prevalence of coronary artery and peripheral vascular diseases. The percentage of deaths from these causes has been slowly rising as the numbers of transplanted diabetic patients and the average age of all recipients increase. More than 50% of renal recipient mortality is attributable to cardiovascular disease. In addition to strict control of blood pressure and blood lipid levels, close monitoring of patients for indications of further medical or surgical intervention is an important part of management.

Hypertension may be caused by (1) native kidney disease, (2) rejection activity in the transplant, (3) renal artery stenosis if an end-to-end anastomosis was constructed with an iliac artery branch, and (4) renal calcineurin inhibitor toxicity, which may improve with reduction in dose. Whereas ACE inhibitors may be useful, calcium channel blockers are more frequently used initially. Amelioration of hypertension to the range of 120–130/70–80 mmHg should be the goal in all patients.

Hypercalcemia after transplantation may indicate failure of hyperplastic parathyroid glands to regress. Aseptic necrosis of the head of the femur is probably due to preexisting hyperparathyroidism, with aggravation by glucocorticoid treatment. With improved management of calcium and phosphorus metabolism during chronic dialysis, the incidence of parathyroid-related complications has fallen dramatically. Persistent hyperparathyroid activity may require subtotal parathyroidectomy.

Although most transplant patients have robust production of erythropoietin and normalization of hemoglobin, *anemia* is commonly seen in the posttransplant period. Often the anemia is attributable to bone marrow-suppressant immunosuppressive medications such as azathioprine, mycophenolic acid, and sirolimus. Gastrointestinal bleeding is a common side effect of high-dose and long-term steroid administration. Many transplant patients have creatinine clearances of 30–50 mL/min and can be considered in the same way as other patients with chronic renal insufficiency for anemia management, including supplemental erythropoietin.

Chronic hepatitis, particularly when due to hepatitis B virus, can be a progressive, fatal disease over a decade or so. Patients who are persistently hepatitis B surface antigen-positive are at higher risk, according to some studies, but the presence of hepatitis C virus is also a concern when one embarks on a course of immunosuppression in a transplant recipient.

338 Glomerular Diseases

Julia B. Lewis, Eric G. Neilson

Two human kidneys harbor nearly 1.8 million glomerular capillary tufts. Each glomerular tuft resides within Bowman's space. The capsule circumscribing this space is lined by parietal epithelial cells that transition into tubular epithelia forming the proximal nephron or migrate into the tuft to replenish podocytes. The glomerular capillary tuft derives from an afferent arteriole that forms a branching capillary bed embedded in mesangial matrix (Fig. 338-1). This capillary network funnels into an efferent arteriole, which passes filtered blood into cortical peritubular capillaries or medullary vasa recta that supply and exchange with a folded tubular architecture. Hence the glomerular capillary tuft, fed and drained by arterioles, represents an arteriolar portal system. Fenestrated endothelial cells resting on a glomerular basement membrane (GBM) line glomerular capillaries. Delicate foot processes extending from epithelial podocytes shroud the outer surface of these capillaries, and podocytes interconnect to each other by slit-pore membranes forming a selective filtration barrier.

The glomerular capillaries filter 120–180 L/d of plasma water containing various solutes for reclamation or discharge by downstream

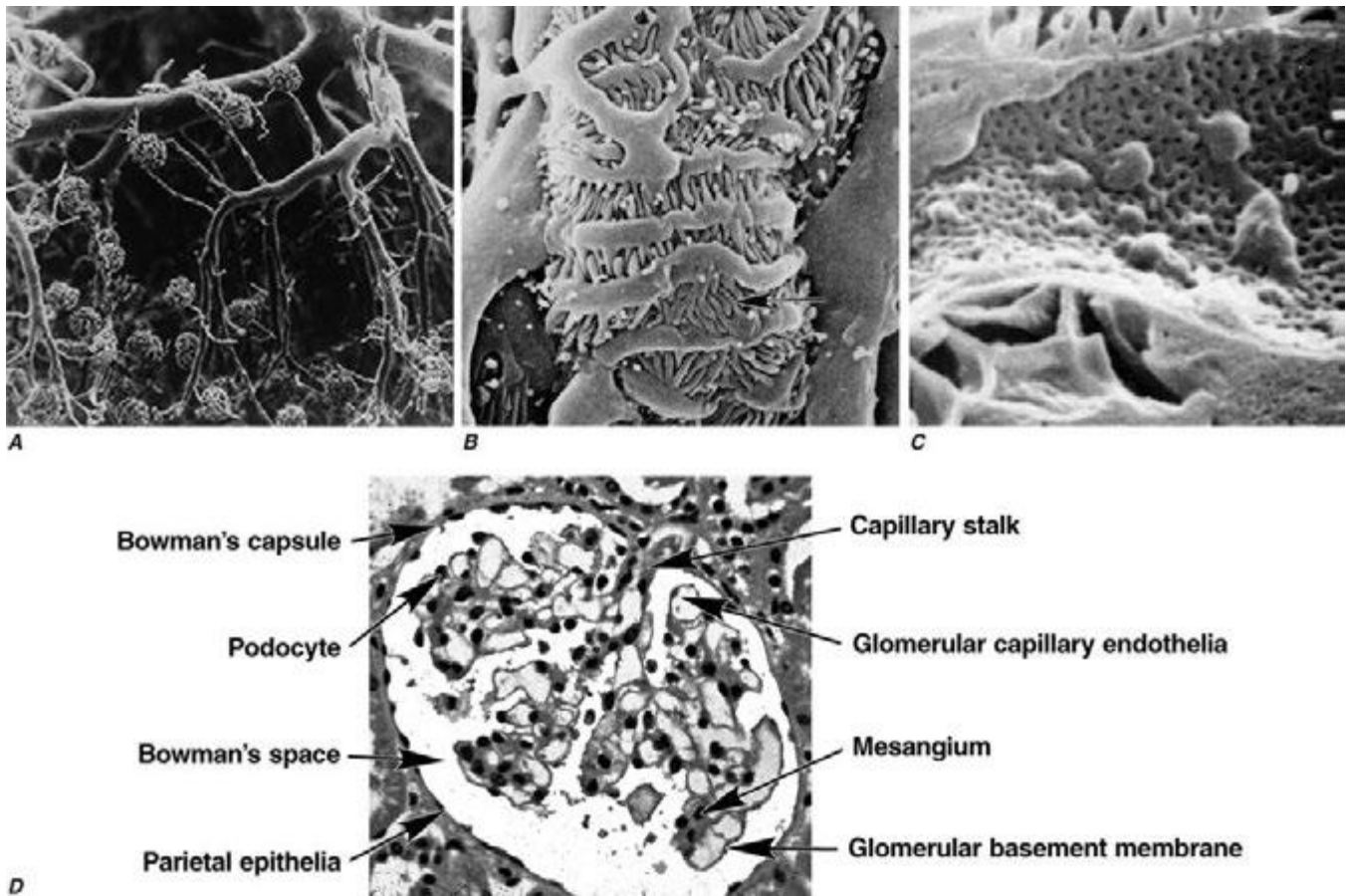


FIGURE 338-1 Glomerular architecture. **A.** The glomerular capillaries form from a branching network of renal arteries, arterioles, leading to an afferent arteriole, glomerular capillary bed (tuft), and a draining efferent arteriole. (From VH Gattone II et al: *Hypertension* 5:8, 1983.) **B.** Scanning electron micrograph of podocytes that line the outer surface of the glomerular capillaries (arrow shows foot process). **C.** Scanning electron micrograph of the fenestrated endothelia lining the glomerular capillary. **D.** The various normal regions of the glomerulus on light microscopy. (A–C: Courtesy of Dr. Vincent Gattone, Indiana University; with permission.)

tubules. Most large proteins and all cells are excluded from filtration by a physicochemical barrier governed by pore size and negative electrostatic charge. The mechanics of filtration and reclamation are quite complicated for many solutes (Chap. 325). For example, in the case of serum albumin, the glomerulus is an imperfect barrier. Although albumin has a negative charge, which would tend to repel the negatively charged GBM, it only has a physical radius of 3.6 nm, while pores in the GBM and slit-pore membranes have a radius of 4 nm. Consequently, variable amounts of albumin inevitably cross the filtration barrier to be reclaimed by megalin and cubilin receptors along the proximal tubule. Remarkably, humans with normal nephrons excrete on average 8–10 mg of albumin in daily voided urine, approximately 20–60% of total excreted protein. This amount of albumin, and other proteins, can rise to gram quantities following glomerular injury.

The breadth of diseases affecting the glomerulus is expansive because the glomerular capillaries can be injured in a variety of ways, producing many different lesions. Some order to this vast subject is brought by grouping all of these diseases into a smaller number of clinical syndromes.

PATHOGENESIS OF GLOMERULAR DISEASE

There are many forms of glomerular disease with pathogenesis variably linked to the presence of genetic mutations, infection, toxin exposure, autoimmunity, atherosclerosis, hypertension, emboli, thrombosis, or diabetes mellitus. Even after careful study, however, the cause often remains unknown, and the lesion is called *idiopathic*. Specific or unique features of pathogenesis are mentioned with the description of each of the glomerular diseases later in this chapter.

Some glomerular diseases result from genetic mutations producing familial disease or a founder effect: congenital nephrotic syndrome from mutations in *NPHS1* (nephrin) and *NPHS2* (podocin) affect the slit-pore membrane at birth, and *TRPC6* cation channel mutations produce *focal segmental glomerulosclerosis* (FSGS) in adulthood; polymorphisms in the gene encoding apolipoprotein L1, *APOL1*, are a major risk for nearly 70% of African Americans with nondiabetic end-stage renal disease, particularly FSGS; mutations in complement factor H associate with *membranoproliferative glomerulonephritis* (MPGN) or *atypical hemolytic uremic syndrome* (aHUS), type II partial lipodystrophy from mutations in genes encoding lamin A/C, or *PPAR γ* cause a metabolic syndrome associated with MPGN, which is sometimes accompanied by dense deposits and C3 nephritic factor; Alport's syndrome, from mutations in the genes encoding for the $\alpha 3$, $\alpha 4$, or $\alpha 5$ chains of type IV collagen, produces *split-basement membranes* with *glomerulosclerosis*; and lysosomal storage diseases, such as α -galactosidase A deficiency causing Fabry's disease and *N*-acetylneuraminic acid hydrolase deficiency causing nephrosialidosis, produce FSGS.

Systemic hypertension and atherosclerosis can produce pressure stress, ischemia, or lipid oxidants that lead to *chronic glomerulosclerosis*. *Malignant hypertension* can quickly complicate glomerulosclerosis with fibrinoid necrosis of arterioles and glomeruli, thrombotic microangiopathy, and acute renal failure. *Diabetic nephropathy* is an acquired sclerotic injury associated with thickening of the GBM secondary to the long-standing effects of hyperglycemia, advanced glycosylation end products, and reactive oxygen species.

Inflammation of the glomerular capillaries is called *glomerulonephritis*. Most glomerular or mesangial antigens involved in *immune-mediated glomerulonephritis* are unknown (Fig. 338-2). Glomerular epithelial or mesangial cells may shed or express epitopes that mimic

other immunogenic proteins made elsewhere in the body. Bacteria, fungi, and viruses can directly infect the kidney producing their own antigens. Autoimmune diseases like idiopathic *membranous glomerulonephritis* (MGN) or MPGN are confined to the kidney, whereas systemic inflammatory diseases like *lupus nephritis* or *granulomatosis with polyangiitis* (Wegener's) spread to the kidney, causing secondary glomerular injury. *Antiglomerular basement membrane disease* producing Goodpasture's syndrome primarily injures both the lung and kidney because of the narrow distribution of the $\alpha 3$ NC1 domain of type IV collagen that is the target antigen.

Local activation of Toll-like receptors on glomerular cells, deposition of immune complexes, or complement injury to glomerular structures induces mononuclear cell infiltration, which subsequently leads to an adaptive immune response attracted to the kidney by local release of chemokines. Neutrophils, macrophages, and T cells are drawn by

chemokines into the glomerular tuft, where they react with antigens and epitopes on or near somatic cells or their structures, producing more cytokines and proteases that damage the mesangium, capillaries, and/or the GBM. While the adaptive immune response is similar to that of other tissues, early T cell activation plays an important role in the mechanism of glomerulonephritis. Antigens presented by class II major histocompatibility complex (MHC) molecules on macrophages and dendritic cells in conjunction with associative recognition molecules engage the CD4/8 T cell repertoire.

Mononuclear cells by themselves can injure the kidney, but autoimmune events that damage glomeruli classically produce a humoral immune response. *Poststreptococcal glomerulonephritis*, *lupus nephritis*, and *idiopathic membranous nephritis* typically are associated with immune deposits along the GBM, while anti-GBM antibodies produce the linear binding of anti-GBM disease. Preformed circulating

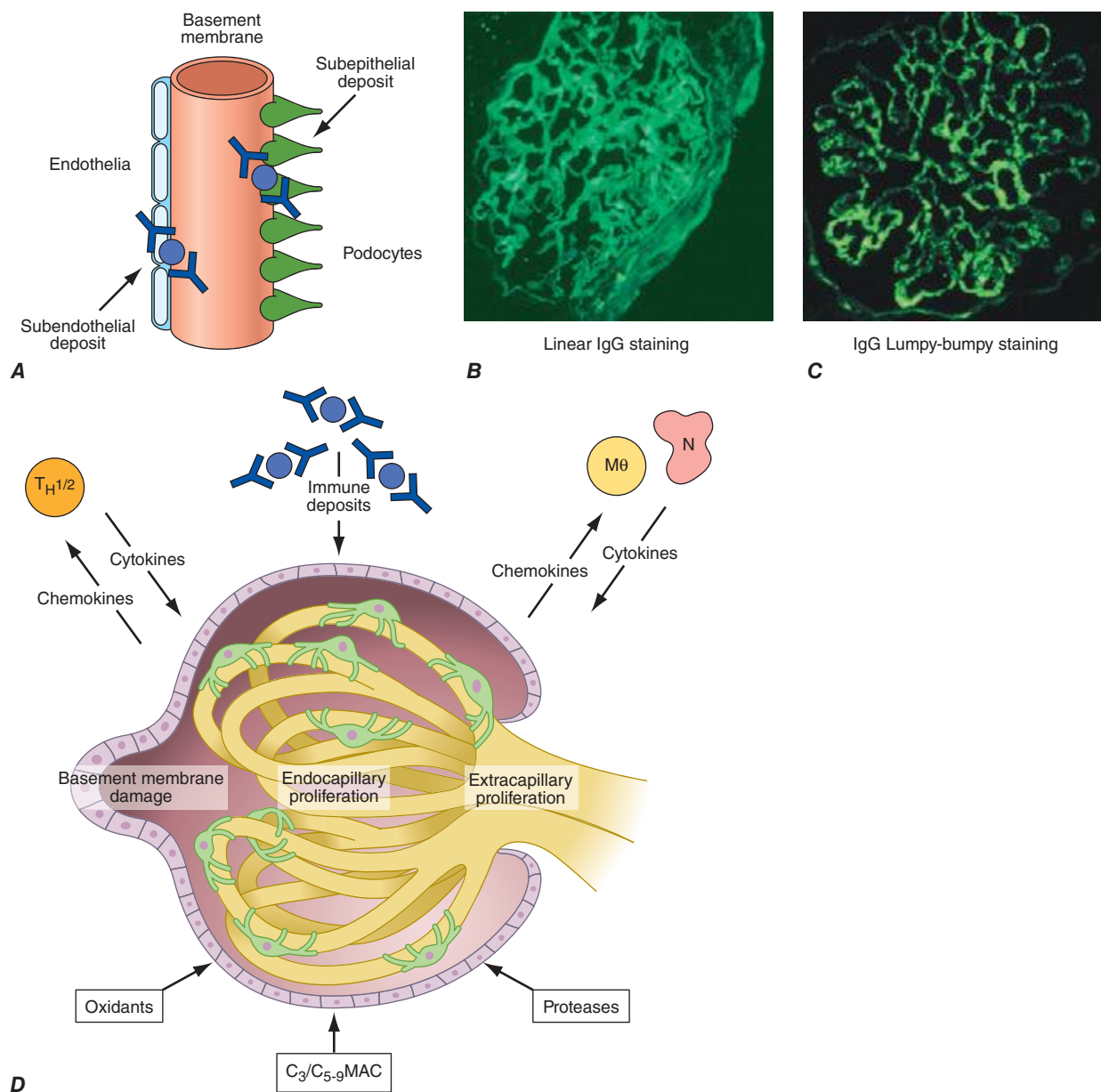


FIGURE 338-2 The glomerulus is injured by a variety of mechanisms. **A.** Preformed immune deposits can precipitate from the circulation and collect along the glomerular basement membrane (GBM) in the subendothelial space or can form in situ along the subepithelial space. **B.** Immunofluorescent staining of glomeruli with labeled anti-IgG demonstrating linear staining from a patient with anti-GBM disease or immune deposits from a patient with membranous glomerulonephritis. **C.** The mechanisms of glomerular injury have a complicated pathogenesis. Immune deposits and complement deposition classically draw macrophages and neutrophils into the glomerulus. T lymphocytes may follow to participate in the injury pattern as well. **D.** Amplification mediators as locally derived oxidants and proteases expand this inflammation, and, depending on the location of the target antigen and the genetic polymorphisms of the host, basement membranes are damaged with either endocapillary or extracapillary proliferation.

immune complexes can precipitate along the subendothelial side of the GBM, while other immune deposits form in situ on the subepithelial side. These latter deposits accumulate when circulating autoantibodies find their antigen trapped along the subepithelial edge of the GBM. Immune deposits in the glomerular mesangium may result from the deposition of preformed circulating complexes or in situ antigen-antibody interactions. Immune deposits stimulate the release of local proteases and activate the complement cascade, producing C_{5-9} attack complexes. In addition, local oxidants damage glomerular structures, producing proteinuria and effacement of the podocytes. Overlapping etiologies or pathophysiologic mechanisms can produce similar glomerular lesions, suggesting that downstream molecular and cellular responses often converge toward common patterns of injury.

PROGRESSION OF GLOMERULAR DISEASE

Persistent glomerulonephritis that worsens renal function is always accompanied by interstitial nephritis, renal fibrosis, and tubular atrophy (see Fig. 62e-27). What is not so obvious, however, is that renal failure in glomerulonephritis best correlates histologically with the appearance of tubulointerstitial nephritis rather than with the type of inciting glomerular injury.

Loss of renal function due to interstitial damage is explained hypothetically by several mechanisms. The simplest explanation is that urine flow is impeded by tubular obstruction as a result of interstitial inflammation and fibrosis. Thus, obstruction of the tubules with debris or by extrinsic compression results in aglomerular nephrons. A second mechanism suggests that interstitial changes, including interstitial edema or fibrosis, alter tubular and vascular architecture and thereby compromise the normal tubular transport of solutes and water from tubular lumen to vascular space. This failure increases the solute and water content of the tubule fluid, resulting in isosthenuria and polyuria. Adaptive mechanisms related to tubuloglomerular feedback also fail, resulting in a reduction of renin output from the juxtaglomerular apparatus trapped by interstitial inflammation. Consequently, the local vasoconstrictive influence of angiotensin II on the glomerular arterioles decreases, and filtration drops owing to a generalized decrease in arteriolar tone. A third mechanism involves changes in vascular resistance due to damage of peritubular capillaries. The cross-sectional volume of these capillaries is decreased by interstitial inflammation, edema, or fibrosis. These structural alterations in vascular resistance affect renal function through two mechanisms. First, tubular cells are very metabolically active, and, as a result, decreased perfusion leads to ischemic injury. Second, impairment of glomerular arteriolar outflow leads to increased intraglomerular hypertension in less-involved glomeruli; this selective intraglomerular hypertension aggravates and extends *mesangial sclerosis* and *glomerulosclerosis* to less-involved glomeruli. Regardless of the exact mechanism, early *acute tubulointerstitial nephritis* (see Fig. 62e-27) suggests potentially recoverable renal function, whereas the development of *chronic interstitial fibrosis* prognosticates permanent loss (see Fig. 62e-30).

Persistent damage to glomerular capillaries spreads to the tubulointerstitium in association with proteinuria. There is a hypothesis that efferent arterioles leading from inflamed glomeruli carry forward inflammatory mediators, which induces downstream interstitial nephritis, resulting in fibrosis. Glomerular filtrate from injured glomerular capillaries adherent to Bowman's capsule may also be misdirected to the periglomerular interstitium. Most nephrologists believe, however, that proteinuric glomerular filtrate forming tubular fluid is

the primary route to downstream tubulointerstitial injury, although none of these hypotheses are mutually exclusive.

The simplest explanation for the effect of proteinuria on the development of interstitial nephritis is that increasingly severe proteinuria, carrying activated cytokines and lipoproteins producing reactive oxygen species, triggers a downstream inflammatory cascade in and around epithelial cells lining the tubular nephron. These effects induce T lymphocyte and macrophage infiltrates in the interstitial spaces along with fibrosis and tubular atrophy.

Tubules disaggregate following direct damage to their basement membranes, leading to epithelial-mesenchymal transitions forming more interstitial fibroblasts at the site of injury. Transforming growth factor β (TGF- β), fibroblast growth factor 2 (FGF-2), hypoxemia-inducible factor 1 α (HIF-1 α), and platelet-derived growth factor (PDGF) are particularly active in this transition. With persistent nephritis, fibroblasts multiply and lay down tenascin and a fibronectin scaffold for the polymerization of new interstitial collagen types I/III. These events form scar tissue through a process called fibrogenesis. In experimental studies, bone morphogenetic protein 7 and hepatocyte growth factor can reverse early fibrogenesis and preserve tubular architecture. When fibroblasts outdistance their survival factors, apoptosis occurs, and the permanent renal scar becomes acellular, leading to irreversible renal failure.

APPROACH TO THE PATIENT: Glomerular Disease

HEMATURIA, PROTEINURIA, AND PYURIA

Patients with glomerular disease usually have some hematuria with varying degrees of proteinuria. Hematuria is typically asymptomatic. As few as three to five red blood cells in the spun sediment from first-voided morning urine is suspicious. The diagnosis of glomerular injury can be delayed because patients will not realize they have *microscopic hematuria*, and only rarely with the exception of IgA nephropathy and sickle cell disease is *gross hematuria* present. When working up microscopic hematuria, perhaps accompanied by minimal proteinuria (<500 mg/24 h), it is important to exclude anatomic lesions, such as malignancy of the urinary tract, particularly in older men. Microscopic hematuria may also appear with the onset of benign prostatic hypertrophy, interstitial nephritis, papillary necrosis, hypercalciuria, renal stones, cystic kidney diseases, or renal vascular injury. However, when red blood cell casts (see Fig. 62e-34) or dysmorphic red blood cells are found in the sediment, glomerulonephritis is likely.

Sustained proteinuria >1–2 g/24 h is also commonly associated with glomerular disease. Patients often will not know they have proteinuria unless they become edematous or notice foaming urine on voiding. *Sustained proteinuria* has to be distinguished from lesser amounts of so-called *benign proteinuria* in the normal population (Table 338-1). This latter class of proteinuria is nonsustained, generally <1 g/24 h, and is sometimes called *functional* or *transient proteinuria*. Fever, exercise, obesity, sleep apnea, emotional stress, and congestive heart failure can explain transient proteinuria. Proteinuria only seen with upright posture is called *orthostatic proteinuria* and has a benign prognosis. Isolated proteinuria sustained over multiple clinic visits is found in many glomerular lesions. Proteinuria in most adults with glomerular disease is *nonselective*, containing albumin and a mixture of other serum proteins, whereas in children with *minimal change disease*, the proteinuria is *selective* and composed largely of albumin.

TABLE 338-1 URINE ASSAYS FOR ALBUMINURIA/PROTEINURIA

	24-Hour Albumin ^a (mg/24 h)	Albumin ^a /Creatinine Ratio (mg/g)	Dipstick Proteinuria	24-Hour Urine Protein ^b (mg/24 h)
Normal	8–10	<30	–	<150
Microalbuminuria	30–300	30–300	–/Trace/1+	–
Proteinuria	>300	>300	Trace–3+	>150

^aAlbumin detected by radioimmunoassay. ^bAlbumin represents 20–60% of the total protein excreted in the urine.

Some patients with inflammatory glomerular disease, such as acute poststreptococcal glomerulonephritis or MPGN, have *pyuria* characterized by the presence of considerable numbers of leukocytes. This latter finding has to be distinguished from urine infected with bacteria.

CLINICAL SYNDROMES

Various forms of glomerular injury can also be parsed into several distinct syndromes on clinical grounds (Table 338-2). These syndromes, however, are not always mutually exclusive. There is an *acute nephritic syndrome* producing 1–2 g/24 h of proteinuria,

TABLE 338-2 PATTERNS OF CLINICAL GLOMERULONEPHRITIS

Glomerular Syndromes	Proteinuria	Hematuria	Vascular Injury
Acute Nephritic Syndromes			
Poststreptococcal glomerulonephritis ^a	+ / ++	+ / +++	–
Subacute bacterial endocarditis ^a	+ / ++	++	–
Lupus nephritis ^a	+ / ++	+ / +++	+
Antiglomerular basement membrane disease ^a	++	+ / +++	–
IgA nephropathy ^a	+ / ++	+++ ^c	–
ANCA small-vessel vasculitis ^a			
Granulomatosis with polyangiitis (Wegener's)	+ / ++	+ / +++	++++
Microscopic polyangiitis	+ / ++	+ / +++	++++
Churg-Strauss syndrome	+ / ++	+ / +++	++++
Henoch-Schönlein purpura ^a	+ / ++	+ / +++	++++
Cryoglobulinemia ^a	+ / ++	+ / +++	++++
Membranoproliferative glomerulonephritis ^a	++	+ / +++	–
Mesangioproliferative glomerulonephritis	+	+ / ++	–
Pulmonary-Renal Syndromes			
Goodpasture's syndrome ^a	++	+ / +++	–
ANCA small-vessel vasculitis ^a			
Granulomatosis with polyangiitis (Wegener's)	+ / ++	+ / +++	++++
Microscopic polyangiitis	+ / ++	+ / +++	++++
Churg-Strauss syndrome	+ / ++	+ / +++	++++
Henoch-Schönlein purpura ^a	+ / ++	+ / +++	++++
Cryoglobulinemia ^a	+ / ++	+ / +++	++++
Nephrotic Syndromes			
Minimal change disease	++++	–	–
Focal segmental glomerulosclerosis	+++ / +++++	+	–
Membranous glomerulonephritis	++++	+	–
Diabetic nephropathy	++ / +++++	– / +	–
AL and AA amyloidosis	+++ / +++++	+	+ / ++
Light-chain deposition disease	+++	+	–
Fibrillary-immunotactoid disease	+++ / +++++	+	+
Fabry's disease	+	+	–
Basement Membrane Syndromes			
Anti-GBM disease ^a	++	+ / +++	–
Alport's syndrome	++	++	–
Thin basement membrane disease	+	++	–
Nail-patella syndrome	++ / +++	++	–
Glomerular Vascular Syndromes			
Atherosclerotic nephropathy	+	+	+++
Hypertensive nephropathy ^b	+ / ++	+ / ++	++
Cholesterol emboli	+ / ++	++	+++
Sickle cell disease	+ / ++	+++ ^c	+++
Thrombotic microangiopathies	++	++	+++
Antiphospholipid syndrome	++	++	+++
ANCA small-vessel vasculitis ^a			
Granulomatosis with polyangiitis (Wegener's)	+ / ++	+ / +++	++++
Microscopic polyangiitis	+ / ++	+ / +++	++++
Churg-Strauss syndrome	+++	+ / +++	++++
Henoch-Schönlein purpura ^a	+ / ++	+ / +++	++++
Cryoglobulinemia ^a	+ / ++	+ / +++	++++
AL and AA amyloidosis	+++ / +++++	+	+ / ++

(Continued)

TABLE 338-2 PATTERNS OF CLINICAL GLOMERULONEPHRITIS (CONTINUED)

Infectious Disease–Associated Syndromes			
Poststreptococcal glomerulonephritis ^a	+/++	++/+++	–
Subacute bacterial endocarditis ^a	+/++	++	–
HIV	+++	+ / ++	–
Hepatitis B and C	+++	+ / ++	–
Syphilis	+++	+	–
Leprosy	+++	+	–
Malaria	+++	+ / ++	–
Schistosomiasis	+++	+ / ++	–

^aCan present as rapidly progressive glomerulonephritis (RPGN); sometimes called crescentic glomerulonephritis. ^bCan present as a malignant hypertensive crisis producing an aggressive fibrinoid necrosis in arterioles and small arteries with microangiopathic hemolytic anemia. ^cCan present with gross hematuria.

Abbreviations: AA, amyloid A; AL, amyloid L; ANCA, antineutrophil cytoplasmic antibodies; GBM, glomerular basement membrane.

hematuria with red blood cell casts, pyuria, hypertension, fluid retention, and a rise in serum creatinine associated with a reduction in glomerular filtration. If glomerular inflammation develops slowly, the serum creatinine will rise gradually over many weeks, but if the serum creatinine rises quickly, particularly over a few days, acute nephritis is sometimes called *rapidly progressive glomerulonephritis* (RPGN); the histopathologic term *crescentic glomerulonephritis* is the pathologic equivalent of the clinical presentation of RPGN. When patients with RPGN present with lung hemorrhage from Goodpasture's syndrome, antineutrophil cytoplasmic antibodies (ANCA)-associated small-vessel vasculitis, lupus erythematosus, or cryoglobulinemia, they are often diagnosed as having a *pulmonary-renal syndrome*. *Nephrotic syndrome* describes the onset of heavy proteinuria (>3.0 g/24 h), hypertension, hypercholesterolemia, hypoalbuminemia, edema/anasarca, and microscopic hematuria; if only large amounts of proteinuria are present without clinical manifestations, the condition is sometimes called *nephrotic-range proteinuria*. The glomerular filtration rate (GFR) in these patients may initially be normal or, rarely, higher than normal, but with persistent hyperfiltration and continued nephron loss, it typically declines over months to years. Patients with a *basement membrane syndrome* either have genetically abnormal basement membranes (Alport's syndrome) or an autoimmune response to basement membrane collagen IV (Goodpasture's syndrome) associated with microscopic hematuria, mild to heavy proteinuria, and hypertension with variable elevations in serum creatinine. *Glomerular-vascular syndrome* describes patients with vascular injury producing hematuria and moderate proteinuria. Affected individuals can have vasculitis, thrombotic microangiopathy, antiphospholipid syndrome, or, more commonly, a systemic disease such as atherosclerosis, cholesterol emboli, hypertension, sickle cell anemia, and autoimmunity. *Infectious disease-associated syndrome* is most important if one has a global perspective. Save for subacute bacterial endocarditis in the Western Hemisphere, malaria and schistosomiasis may be the most common causes of glomerulonephritis throughout the world, closely followed by HIV and chronic hepatitis B and C. These infectious diseases produce a variety of inflammatory reactions in glomerular capillaries, ranging from nephrotic syndrome to acute nephritic injury, and urinalyses that demonstrate a combination of hematuria and proteinuria.

These six general categories of syndromes are usually determined at the bedside with the help of a history and physical examination, blood chemistries, renal ultrasound, and urinalysis. These initial studies help frame further diagnostic workup that typically involves testing of the serum for the presence of various proteins (HIV and hepatitis B and C antigens), antibodies (anti-GBM, antiphospholipid, antistreptolysin O [ASO], anti-DNAse, antihyaluronidase, ANCA, anti-DNA, cryoglobulins, anti-HIV, and anti-hepatitis B and C antibodies) or depletion of complement components (C₃ and C₄). The bedside history and

physical examination can also help determine whether the glomerulonephritis is isolated to the kidney (*primary glomerulonephritis*) or is part of a systemic disease (*secondary glomerulonephritis*).

When confronted with an abnormal urinalysis and elevated serum creatinine, with or without edema or congestive heart failure, one must consider whether the glomerulonephritis is *acute* or *chronic*. This assessment is best made by careful history (last known urinalysis or serum creatinine during pregnancy or insurance physical, evidence of infection, or use of medication or recreational drugs); the size of the kidneys on renal ultrasound examination; and how the patient feels at presentation. Chronic glomerular disease often presents with decreased kidney size. Patients who quickly develop renal failure are fatigued and weak and often have uremic symptoms associated with nausea, vomiting, fluid retention, and somnolence. Primary glomerulonephritis presenting with renal failure that has progressed slowly, however, can be remarkably asymptomatic, as are patients with acute glomerulonephritis without much loss in renal function. Once this initial information is collected, selected patients who are clinically stable, have adequate blood clotting parameters, and are willing and able to receive treatment are encouraged to have a renal biopsy.

RENAL PATHOLOGY

A renal biopsy in the setting of glomerulonephritis quickly identifies the type of glomerular injury and often suggests a course of treatment. The biopsy is processed for light microscopy using stains for *hematoxylin and eosin (H&E)* to assess cellularity and architecture, *periodic acid-Schiff (PAS)* to stain carbohydrate moieties in the membranes of the glomerular tuft and tubules, *Jones-methenamine silver* to enhance basement membrane structure, *Congo red* for amyloid deposits, and *Masson's trichrome* to identify collagen deposition and assess the degree of glomerulosclerosis and interstitial fibrosis. Biopsies are also processed for direct immunofluorescence using conjugated antibodies against IgG, IgM, and IgA to detect the presence of "lumpy-bumpy" immune deposits or "linear" IgG or IgA antibodies bound to GBM, antibodies against trapped complement proteins (C₃ and C₄), or specific antibodies against a relevant antigen. High-resolution electron microscopy can clarify the principal location of immune deposits and the status of the basement membrane.

Each region of a renal biopsy is assessed separately. By light microscopy, glomeruli (at least 10 and ideally 20) are reviewed individually for discrete lesions; <50% involvement is considered *focal*, and >50% is *diffuse*. Injury in each glomerular tuft can be *segmental*, involving a portion of the tuft, or *global*, involving most of the glomerulus. Glomeruli having *proliferative* characteristics show increased cellularity. When cells in the capillary tuft proliferate, it is called *endocapillary*, and when cellular proliferation extends into Bowman's space, it is called *extracapillary*. *Synechiae* are formed when epithelial podocytes attach to Bowman's capsule in the setting of glomerular injury; *crescents*, which in some cases may be the extension of synechiae, develop

when fibrocellular/fibrin collections fill all or part of Bowman's space; and *sclerotic* glomeruli show acellular, amorphous accumulations of proteinaceous material throughout the tuft with loss of functional capillaries and normal mesangium. Since *age-related glomerulosclerosis* is common in adults, one can estimate the background percentage of sclerosis by dividing the patient's age in half and subtracting 10. Immunofluorescent and electron microscopy can detect the presence and location of *subepithelial*, *subendothelial*, or *mesangial* immune deposits, or *reduplication* or *splitting* of the basement membrane. In the other regions of the biopsy, the vasculature surrounding glomeruli and tubules can show *angiopathy*, *vasculitis*, the presence of *fibrils*, or *thrombi*. The tubules can be assessed for adjacency to one another; separation can be the result of edema, tubular dropout, or collagen deposition resulting from interstitial fibrosis. Interstitial fibrosis is an ominous sign of irreversibility and progression to renal failure.

ACUTE NEPHRITIC SYNDROMES

Acute nephritic syndromes classically present with hypertension, hematuria, red blood cell casts, pyuria, and mild to moderate proteinuria. Extensive inflammatory damage to glomeruli causes a fall in GFR and eventually produces uremic symptoms with salt and water retention, leading to edema and hypertension.

POSTSTREPTOCOCCAL GLOMERULONEPHRITIS

Poststreptococcal glomerulonephritis is prototypical for *acute endocapillary proliferative glomerulonephritis*. The incidence of poststreptococcal glomerulonephritis has dramatically decreased in developed countries and in these locations is typically sporadic. Acute poststreptococcal glomerulonephritis in underdeveloped countries is epidemic and usually affects children between the ages of 2 and 14 years, but in developed countries is more typical in the elderly, especially in association with debilitating conditions. It is more common in males, and the familial or cohabitant incidence is as high as 40%. Skin and throat infections with particular M types of streptococci (nephritogenic strains) antedate glomerular disease; M types 47, 49, 55, 2, 60, and 57 are seen following impetigo and M types 1, 2, 4, 3, 25, 49, and 12 with pharyngitis. Poststreptococcal glomerulonephritis due to impetigo develops 2–6 weeks after skin infection and 1–3 weeks after streptococcal pharyngitis.

The renal biopsy in poststreptococcal glomerulonephritis demonstrates hypercellularity of mesangial and endothelial cells, glomerular infiltrates of polymorphonuclear leukocytes, granular subendothelial immune deposits of IgG, IgM, C₃, C₄, and C₅₋₉, and subepithelial deposits (which appear as “humps”) (see Fig. 62e-6). (See [Glomerular Schematic 1](#).) Poststreptococcal glomerulonephritis is an immune-mediated disease involving putative streptococcal antigens, circulating

immune complexes, and activation of complement in association with cell-mediated injury. Many candidate antigens have been proposed over the years; candidates from nephritogenic streptococci of interest at the moment are: a cationic cysteine proteinase known as streptococcal pyrogenic exotoxin B (SPEB) that is generated by proteolysis of a zymogen precursor (zSPEB), and NAP1r, the nephritis-associated plasmin receptor. These two antigens have biochemical affinity for plasmin, bind as complexes facilitated by this relationship, and activate the alternate complement pathway. The nephritogenic antigen, SPEB, has been demonstrated inside the subepithelial “humps” on biopsy.

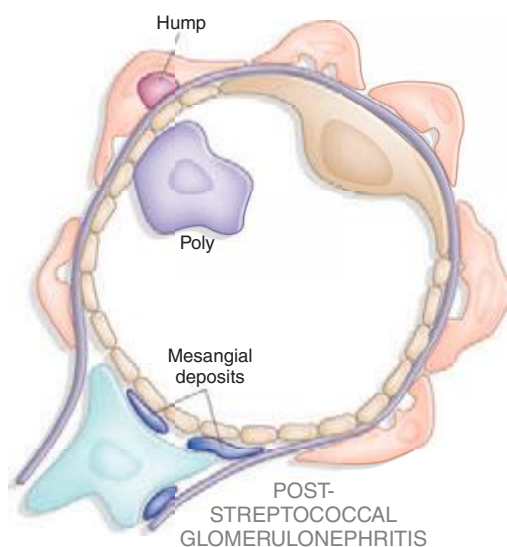
The classic presentation is an acute nephritic picture with hematuria, pyuria, red blood cell casts, edema, hypertension, and oliguric renal failure, which may be severe enough to appear as RPGN. Systemic symptoms of headache, malaise, anorexia, and flank pain (due to swelling of the renal capsule) are reported in as many as 50% of cases. Five percent of children and 20% of adults have proteinuria in the nephrotic range. In the first week of symptoms, 90% of patients will have a depressed CH₅₀ and decreased levels of C₃ with normal levels of C₄. Positive rheumatoid factor (30–40%), cryoglobulins and circulating immune complexes (60–70%), and ANCA against myeloperoxidase (10%) are also reported. Positive cultures for streptococcal infection are inconsistently present (10–70%), but increased titers of ASO (30%), anti-DNAse, (70%), or antihyaluronidase antibodies (40%) can help confirm the diagnosis. Consequently, the diagnosis of poststreptococcal glomerulonephritis rarely requires a renal biopsy. A subclinical disease is reported in some series to be four to five times as common as clinical nephritis, and these latter cases are characterized by asymptomatic microscopic hematuria with low serum C₃ complement levels.

Treatment is supportive, with control of hypertension, edema, and dialysis as needed. Antibiotic treatment for streptococcal infection should be given to all patients and their cohabitants. There is no role for immunosuppressive therapy, even in the setting of crescents. Recurrent poststreptococcal glomerulonephritis is rare despite repeated streptococcal infections. Early death is rare in children but does occur in the elderly. Overall, the prognosis is good, with permanent renal failure being very uncommon, less than 1% in children. Complete resolution of the hematuria and proteinuria in the majority of children occurs within 3–6 weeks of the onset of nephritis but 3–10% of children may have persistent microscopic hematuria, nonnephrotic proteinuria, or hypertension. The prognosis in elderly patients is worse with a high incidence of azotemia (up to 60%), nephrotic-range proteinuria, and end-stage renal disease.

SUBACUTE BACTERIAL ENDOCARDITIS

Endocarditis-associated glomerulonephritis is typically a complication of subacute bacterial endocarditis, particularly in patients who remain untreated for a long time, have negative blood cultures, or have right-sided endocarditis. Glomerulonephritis is unusual in acute bacterial endocarditis because it takes 10–14 days to develop immune complex-mediated injury, by which time the patient has been treated, often with emergent surgery. Grossly, the kidneys in subacute bacterial endocarditis have subcapsular hemorrhages with a “flea-bitten” appearance, and microscopy on renal biopsy reveals focal proliferation around foci of necrosis associated with abundant mesangial, subendothelial, and subepithelial immune deposits of IgG, IgM, and C₃. Patients who present with a clinical picture of RPGN have crescents. Embolic infarcts or septic abscesses may also be present. The pathogenesis hinges on the renal deposition of circulating immune complexes in the kidney with complement activation. Patients present with gross or microscopic hematuria, pyuria, and mild proteinuria or, less commonly, RPGN with rapid loss of renal function. A normocytic anemia, elevated erythrocyte sedimentation rate, hypocomplementemia, high titers of rheumatoid factor, type III cryoglobulins, circulating immune complexes, and ANCAs may be present. Levels of serum creatinine may be elevated at diagnosis, but with modern therapy there is little progression to chronic renal failure. Primary treatment is eradication of the infection with 4–6 weeks of antibiotics, and if accomplished expeditiously, the prognosis for renal recovery is good. ANCA-associated vasculitis sometimes accompanies or is confused

Glomerular schematic 1



As variants of persistent bacterial infection in blood-associated glomerulonephritis, postinfectious glomerulonephritis can occur in patients with ventriculoatrial and ventriculoperitoneal shunts; pulmonary, intraabdominal, pelvic, or cutaneous infections; and infected vascular prostheses. In developed countries, a significant proportion of cases afflict adults, especially the immunocompromised, and the predominant organism is *Staphylococcus*. The clinical presentation of these conditions is variable and includes proteinuria, microscopic hematuria, acute renal failure, and hypertension. Serum complement levels are low, and there may be elevated levels of C-reactive proteins, rheumatoid factor, antinuclear antibodies, and cryoglobulins. Renal lesions include membranoproliferative glomerulonephritis (MPGN), diffuse proliferative and exudative glomerulonephritis (DPGN), or mesangioproliferative glomerulonephritis, sometimes leading to RPGN. Treatment focuses on eradicating the infection, with most patients treated as if they have endocarditis. The prognosis is guarded.

LUPUS NEPHRITIS

Lupus nephritis is a common and serious complication of systemic lupus erythematosus (SLE) and most severe in African-American female adolescents. Thirty to 50% of patients will have clinical manifestations of renal disease at the time of diagnosis, and 60% of adults and 80% of children develop renal abnormalities at some point in the course of their disease. Lupus nephritis results from the deposition of circulating immune complexes, which activate the complement cascade leading to complement-mediated damage, leukocyte infiltration, activation of procoagulant factors, and release of various cytokines. In situ immune complex formation following glomerular binding of nuclear antigens, particularly necrotic nucleosomes, also plays a role in renal injury. The presence of antiphospholipid antibodies may also trigger a thrombotic microangiopathy in a minority of patients.

The clinical manifestations, course of disease, and treatment of lupus nephritis are closely linked to renal pathology. The most common clinical sign of renal disease is proteinuria, but hematuria, hypertension, varying degrees of renal failure, and active urine sediment with red blood cell casts can all be present. Although significant renal pathology can be found on biopsy even in the absence of major abnormalities in the urinalysis, most nephrologists do not biopsy patients until the urinalysis is convincingly abnormal. The extrarenal manifestations of lupus are important in establishing a firm diagnosis of systemic lupus because, while serologic abnormalities are common in lupus nephritis, they are not diagnostic. Anti-dsDNA antibodies that fix complement correlate best with the presence of renal disease. Hypocomplementemia is common in patients with acute lupus nephritis (70–90%) and declining complement levels may herald a flare. Although urinary biomarkers of lupus nephritis are being identified to assist in predicting renal flares, renal biopsy is the only reliable method of identifying the morphologic variants of lupus nephritis.

The World Health Organization (WHO) workshop in 1974 first outlined several distinct patterns of lupus-related glomerular injury; these were modified in 1982. In 2004 the International Society of Nephrology in conjunction with the Renal Pathology Society again updated the classification. This latest version of lesions seen on biopsy (Table 338-3) best defines clinicopathologic correlations, provides valuable prognostic information, and forms the basis for modern treatment recommendations. Class I nephritis describes normal glomerular histology by any technique or normal light microscopy with minimal mesangial deposits on immunofluorescent or electron microscopy. Class II designates mesangial immune complexes with *mesangial proliferation*. Both class I and II lesions are typically associated with minimal renal manifestation and normal renal function; nephrotic syndrome is rare. Patients with lesions limited to the renal mesangium have an excellent prognosis and generally do not need therapy for their lupus nephritis.

The subject of lupus nephritis is presented under acute nephritic syndromes because of the aggressive and important proliferative lesions seen in class III–V renal disease. Class III describes *focal lesions*

TABLE 338-3 CLASSIFICATION FOR LUPUS NEPHRITIS

Class I	Minimal mesangial	Normal histology with mesangial deposits
Class II	Mesangial proliferation	Mesangial hypercellularity with expansion of the mesangial matrix
Class III	Focal nephritis	Focal endocapillary ± extracapillary proliferation with focal subendothelial immune deposits and mild mesangial expansion
Class IV	Diffuse nephritis	Diffuse endocapillary ± extracapillary proliferation with diffuse subendothelial immune deposits and mesangial alterations
Class V	Membranous nephritis	Thickened basement membranes with diffuse subepithelial immune deposits; may occur with class III or IV lesions and is sometimes called mixed membranous and proliferative nephritis
Class VI	Sclerotic nephritis	Global sclerosis of nearly all glomerular capillaries

Note: Revised in 2004 by the International Society of Nephrology-Renal Pathology Society Study Group.

with *proliferation or scarring*, often involving only a segment of the glomerulus (see Fig. 62e-12). Class III lesions have the most varied course. Hypertension, an active urinary sediment, and proteinuria are common with nephrotic-range proteinuria in 25–33% of patients. Elevated serum creatinine is present in 25% of patients. Patients with mild proliferation involving a small percentage of glomeruli respond well to therapy with steroids alone, and fewer than 5% progress to renal failure over 5 years. Patients with more severe proliferation involving a greater percentage of glomeruli have a far worse prognosis and lower remission rates. Treatment of those patients is the same as that for class IV lesions. Many nephrologists believe that class III lesions are simply an early presentation of class IV disease. Others believe severe class III disease is a discrete lesion requiring aggressive therapy. Class IV describes *global, diffuse proliferative lesions* involving the vast majority of glomeruli. Patients with class IV lesions commonly have high anti-DNA antibody titers, low serum complement, hematuria, red blood cell casts, proteinuria, hypertension, and decreased renal function; 50% of patients have nephrotic-range proteinuria. Patients with crescents on biopsy often have a rapidly progressive decline in renal function (see Fig. 62e-12). Without treatment, this aggressive lesion has the worst renal prognosis. However, if a remission—defined as a return to near-normal renal function and proteinuria ≤330 mg/dL per day—is achieved with treatment, renal outcomes are excellent. Current evidence suggests that inducing a remission with administration of high-dose steroids and either cyclophosphamide or mycophenolate mofetil for 2–6 months, followed by maintenance therapy with lower doses of steroids and mycophenolate mofetil or azathioprine, best balances the likelihood of successful remission with the side effects of therapy. There is no consensus on use of high-dose intravenous methylprednisolone versus oral prednisone, monthly intravenous cyclophosphamide versus daily oral cyclophosphamide, or other immunosuppressants such as cyclosporine, tacrolimus, rituximab, or belimumab. Nephrologists tend to avoid prolonged use of cyclophosphamide in patients of childbearing age without first banking eggs or sperm.

The class V lesion describes subepithelial immune deposits producing a *membranous pattern*; a subcategory of class V lesions is associated with proliferative lesions and is sometimes called *mixed membranous and proliferative disease* (see Fig. 62e-11); this category of injury is treated like class IV glomerulonephritis. Sixty percent of patients present with nephrotic syndrome or lesser amounts of proteinuria. Patients with lupus nephritis class V, like patients with *idiopathic membranous nephropathy*, are predisposed to renal-vein thrombosis and other thrombotic complications. A minority of patients with class V will present with hypertension and renal dysfunction. There are conflicting data on the clinical course, prognosis, and appropriate therapy for

patients with class V disease, which may reflect the heterogeneity of this group of patients. Patients with severe nephrotic syndrome, elevated serum creatinine, and a progressive course will probably benefit from therapy with steroids in combination with other immunosuppressive agents. Therapy with inhibitors of the renin-angiotensin system also may attenuate the proteinuria. Antiphospholipid antibodies present in lupus may result in glomerular microthromboses and complicate the course in up to 20% of lupus nephritis patients. The renal prognosis is worse even with anticoagulant therapy.

Patients with any of the above lesions also can transform to another lesion; hence patients often require reevaluation, including repeat renal biopsy. Lupus patients with class VI lesions have greater than 90% *sclerotic glomeruli* and end-stage renal disease with interstitial fibrosis. As a group, approximately 20% of patients with lupus nephritis will reach end-stage disease, requiring dialysis or transplantation. Systemic lupus tends to become quiescent once there is renal failure, perhaps due to the immunosuppressant effects of uremia. However, patients with lupus nephritis have a markedly increased mortality compared with the general population. Renal transplantation in renal failure from lupus, usually performed after approximately 6 months of inactive disease, results in allograft survival rates comparable to patients transplanted for other reasons.

ANTIGLOMERULAR BASEMENT MEMBRANE DISEASE

Patients who develop autoantibodies directed against glomerular basement antigens frequently develop a glomerulonephritis termed *antiglomerular basement membrane (anti-GBM) disease*. When they present with lung hemorrhage and glomerulonephritis, they have a pulmonary-renal syndrome called *Goodpasture's syndrome*. The target epitopes for this autoimmune disease lie in the quaternary structure of $\alpha 3$ NC1 domain of collagen IV. Indeed, anti-GBM disease may be considered an autoimmune "conformeropathy" that involves the perturbation of quaternary structure of the $\alpha 345$ NC1 hexamer. MHC-restricted T cells initiate the autoantibody response because humans are not tolerant to the epitopes created by this quaternary structure. The epitopes are normally sequestered in the collagen IV hexamer and can be exposed by infection, smoking, oxidants, or solvents. Goodpasture's syndrome appears in two age groups: in young men in their late twenties and in men and women in their sixties and seventies. Disease in the younger age group is usually explosive, with hemoptysis, a sudden fall in hemoglobin, fever, dyspnea, and hematuria. Hemoptysis is largely confined to smokers, and those who present with lung hemorrhage as a group do better than older populations who have prolonged, asymptomatic renal injury; presentation with oliguria is often associated with a particularly bad outcome. The performance of an urgent kidney biopsy is important in suspected cases of Goodpasture's syndrome to confirm the diagnosis and assess prognosis. Renal biopsies typically show *focal or segmental necrosis* that later, with aggressive destruction of the capillaries by cellular proliferation, leads to crescent formation in Bowman's space (see Fig. 62e-14). As these lesions progress, there is concomitant interstitial nephritis with fibrosis and tubular atrophy.

The presence of anti-GBM antibodies and complement is recognized on biopsy by linear immunofluorescent staining for IgG (rarely IgA). In testing serum for anti-GBM antibodies, it is particularly important that the $\alpha 3$ NC1 domain of collagen IV alone be used as the target. This is because nonnephritic antibodies against the $\alpha 1$ NC1 domain are seen in paraneoplastic syndromes and cannot be discerned from assays that use whole basement membrane fragments as the binding target. Between 10 and 15% of sera from patients with Goodpasture's syndrome also contain ANCA antibodies against myeloperoxidase. This subset of patients has a vasculitis-associated variant, which has a surprisingly good prognosis with treatment. Prognosis at presentation is worse if there are >50% crescents on renal biopsy with advanced fibrosis, if serum creatinine is >5–6 mg/dL, if oliguria is present, or if there is a need for acute dialysis. Although frequently attempted, most of these latter patients will not respond to plasmapheresis and steroids. Patients with advanced renal failure who present with hemoptysis should still be treated for their lung hemorrhage, as it responds to

plasmapheresis and can be lifesaving. Treated patients with less severe disease typically respond to 8–10 treatments of plasmapheresis accompanied by oral prednisone and cyclophosphamide in the first 2 weeks. Kidney transplantation is possible, but because there is risk of recurrence, experience suggests that patients should wait for 6 months and until serum antibodies are undetectable.

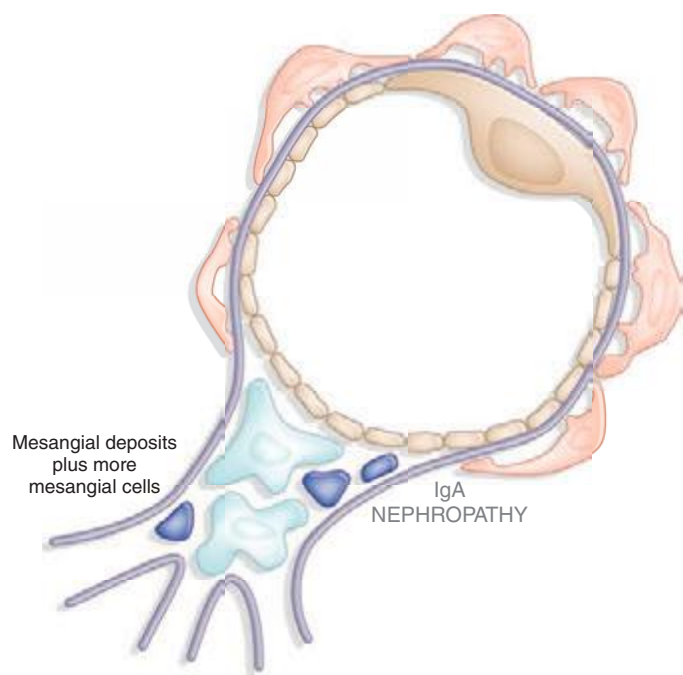
IgA NEPHROPATHY

Berger first described the glomerulonephritis now termed *IgA nephropathy*. It is classically characterized by episodic hematuria associated with the deposition of IgA in the mesangium. IgA nephropathy is one of the most common forms of glomerulonephritis worldwide. There is a male preponderance, a peak incidence in the second and third decades of life, and rare familial clustering. There are geographic differences in the prevalence of IgA nephropathy, with 30% prevalence along the Asian and Pacific Rim and 20% in southern Europe, compared to a much lower prevalence in northern Europe and North America. It was initially hypothesized that variation in detection, in part, accounted for regional differences. With clinical care in nephrology becoming more uniform, this variation in prevalence more likely reflects true differences among racial and ethnic groups.

IgA nephropathy is predominantly a sporadic disease but susceptibility to it has been shown uncommonly to have a genetic component depending on geography and the existence of "founder effects." Familial forms of IgA nephropathy are more common in northern Italy and eastern Kentucky. No single causal gene has been identified. Clinical and laboratory evidence suggests close similarities between Henoch-Schönlein purpura and IgA nephropathy. Henoch-Schönlein purpura is distinguished clinically from IgA nephropathy by prominent systemic symptoms, a younger age (<20 years old), preceding infection, and abdominal complaints. Deposits of IgA are also found in the glomerular mesangium in a variety of systemic diseases, including chronic liver disease, Crohn's disease, gastrointestinal adenocarcinoma, chronic bronchiectasis, idiopathic interstitial pneumonia, dermatitis herpetiformis, mycosis fungoides, leprosy, ankylosing spondylitis, relapsing polychondritis, and Sjögren's syndrome. IgA deposition in these entities is not usually associated with clinically significant glomerular inflammation or renal dysfunction and thus is not called IgA nephropathy.

IgA nephropathy is an immune complex-mediated glomerulonephritis defined by the presence of diffuse mesangial IgA deposits often associated with mesangial hypercellularity. (See **Glomerular Schematic 2.**) IgM, IgG, C₃, or immunoglobulin light chains may be codistributed with IgA. IgA deposited in the mesangium is typically polymeric and of the IgA1 subclass, the pathogenic significance of which is not clear. Abnormalities have been described in IgA production by plasma cells, particularly secretory IgA; in IgA clearance, predominately by the liver; in mesangial IgA clearance and receptors for IgA; and in growth factor and cytokine-mediated events. Currently, however, abnormalities in the O-glycosylation of the hinge region of IgA seem to best account for the pathogenesis of sporadic IgA nephropathy. Despite the presence of elevated serum IgA levels in 20–50% of patients, IgA deposition in skin biopsies in 15–55% of patients, or elevated levels of secretory IgA and IgA-fibronectin complexes, a renal biopsy is necessary to confirm the diagnosis. Although the immunofluorescent pattern of IgA on renal biopsy defines IgA nephropathy in the proper clinical context, a variety of histologic lesions may be seen on light microscopy (see Fig. 62e-8), including DPGN; *segmental sclerosis*; and, rarely, *segmental necrosis with cellular crescent formation*, which typically presents as RPGN.

The two most common presentations of IgA nephropathy are recurrent episodes of macroscopic hematuria during or immediately following an upper respiratory infection often accompanied by proteinuria or persistent asymptomatic microscopic hematuria. Nephrotic syndrome is uncommon. Proteinuria can also first appear late in the course of the disease. Rarely patients present with acute renal failure and a rapidly progressive clinical picture. IgA nephropathy is a benign disease for the majority of patients, and 5–30% of patients may go into a complete remission, with others having hematuria but well



preserved renal function. In the minority of patients who have progressive disease, progression is slow, with renal failure seen in only 25–30% of patients with IgA nephropathy over 20–25 years. This risk varies considerably among populations. Cumulatively, risk factors for the loss of renal function identified thus far account for less than 50% of the variation in observed outcome but include the presence of hypertension or proteinuria, the absence of episodes of macroscopic hematuria, male sex, older age of onset, and extensive glomerulosclerosis or interstitial fibrosis on renal biopsy. Several analyses in large populations of patients found persistent proteinuria for 6 months or longer to have the greatest predictive power for adverse renal outcomes.

There is no agreement on optimal treatment. Both large studies that include patients with multiple glomerular diseases and small studies of patients with IgA nephropathy support the use of angiotensin-converting enzyme (ACE) inhibitors in patients with proteinuria or declining renal function. Tonsillectomy, steroid therapy, and fish oil have all been suggested in small studies to benefit select patients with IgA nephropathy. When presenting as RPGN, patients typically receive steroids, cytotoxic agents, and plasmapheresis.

ANCA SMALL-VESSEL VASCULITIS

A group of patients with small-vessel vasculitis (arterioles, capillaries, and venules; rarely small arteries) and glomerulonephritis have serum ANCA; the antibodies are of two types, anti-proteinase 3 (PR3) or anti-myeloperoxidase (MPO) (**Chap. 385**); Lamp-2 antibodies have also been reported experimentally as potentially pathogenic. ANCA are produced with the help of T cells and activate leukocytes and monocytes, which together damage the walls of small vessels. Endothelial injury also attracts more leukocytes and extends the inflammation. Granulomatosis with polyangiitis, microscopic polyangiitis, and Churg-Strauss syndrome belong to this group because they are ANCA-positive and have a *pauci-immune glomerulonephritis* with few immune complexes in small vessels and glomerular capillaries. Patients with any of these three diseases can have any combination of the above serum antibodies, but anti-PR3 antibodies are more common in granulomatosis with polyangiitis and anti-MPO antibodies are more common in microscopic polyangiitis or Churg-Strauss. Although each of these diseases has some unique clinical features, most features do not predict relapse or progression, and as a group, they are generally treated in the same way. Since mortality is high without treatment, virtually all patients

receive urgent treatment. Induction therapy usually includes some combination of plasmapheresis, methylprednisolone, and cyclophosphamide. Monthly “pulse” IV cyclophosphamide to induce remission of ANCA-associated vasculitis is as effective as daily oral cyclophosphamide but may be associated with increased relapses. Steroids are tapered soon after acute inflammation subsides, and patients are maintained on cyclophosphamide or azathioprine for up to a year to minimize the risk of relapse. Benefit with using mycophenolate mofetil or rituximab has not been proven.

Granulomatosis with Polyangiitis Patients with this disease classically present with fever, purulent rhinorrhea, nasal ulcers, sinus pain, polyarthralgias/arthritis, cough, hemoptysis, shortness of breath, microscopic hematuria, and 0.5–1 g/24 h of proteinuria; occasionally there may be cutaneous purpura and mononeuritis multiplex. Presentation without renal involvement is termed *limited granulomatosis with polyangiitis*, although some of these patients will show signs of renal injury later. Chest x-ray often reveals nodules and persistent infiltrates, sometimes with cavities. Biopsy of involved tissue will show a small-vessel vasculitis and adjacent noncaseating granulomas. Renal biopsies during active disease demonstrate *segmental necrotizing glomerulonephritis* without immune deposits (see Fig. 62e-13). The disease is more common in patients exposed to silica dust and those with α_1 -antitrypsin deficiency, which is an inhibitor of PR3. Relapse after achieving remission is common and is more common in patients with granulomatosis with polyangiitis than the other ANCA-associated vasculitis, necessitating diligent follow-up care. Although associated with an unacceptable high mortality rate without treatment, the greatest threat to patients, especially elderly patients in the first year of therapy, is from adverse events, which are often secondary to treatment, rather than active vasculitis.

Microscopic Polyangiitis Clinically, these patients look somewhat similar to those with granulomatosis with polyangiitis, except they rarely have significant lung disease or destructive sinusitis. The distinction is made on biopsy, where the vasculitis in microscopic polyangiitis is without granulomas. Some patients will also have injury limited to the capillaries and venules.

Churg-Strauss Syndrome When small-vessel vasculitis is associated with peripheral eosinophilia, cutaneous purpura, mononeuritis, asthma, and allergic rhinitis, a diagnosis of Churg-Strauss syndrome is considered. Hypergammaglobulinemia, elevated levels of serum IgE, or the presence of rheumatoid factor sometimes accompanies the allergic state. Lung inflammation, including fleeting cough and pulmonary infiltrates, often precedes the systemic manifestations of disease by years; lung manifestations are rarely absent. A third of patients may have exudative pleural effusions associated with eosinophils. Small-vessel vasculitis and *focal segmental necrotizing glomerulonephritis* can be seen on renal biopsy, usually absent eosinophils or granulomas. The cause of Churg-Strauss syndrome is autoimmune, but the inciting factors are unknown.

MEMBRANOPROLIFERATIVE GLOMERULONEPHRITIS

MPGN is sometimes called *mesangiocapillary glomerulonephritis* or *lobar glomerulonephritis*. It is an immune-mediated glomerulonephritis characterized by thickening of the GBM with mesangioproliferative changes; 70% of patients have hypocomplementemia. MPGN is rare in African Americans, and idiopathic disease usually presents in childhood or young adulthood. MPGN is subdivided pathologically into type I, type II, and type III disease. *Type I MPGN* is commonly associated with persistent hepatitis C infections, autoimmune diseases like lupus or cryoglobulinemia, or neoplastic diseases (**Table 338-4**). *Types II and III MPGN* are usually idiopathic, except in patients with complement factor H deficiency, in the presence of C_3 nephritic factor and/or in partial lipodystrophy producing type II disease, or complement receptor deficiency in type III disease. MPGN has been proposed to be reclassified into immunoglobulin-mediated disease (driven by the classical complement pathway) and non-immunoglobulin-mediated disease (driven by the alternative complement pathway).

TABLE 338-4 MEMBRANOPROLIFERATIVE GLOMERULONEPHRITIS**Type I Disease (Most Common)**

Idiopathic
 Subacute bacterial endocarditis
 Systemic lupus erythematosus
 Hepatitis C ± cryoglobulinemia
 Mixed cryoglobulinemia
 Hepatitis B
 Cancer: Lung, breast, and ovary (germinal)

Type II Disease (Dense Deposit Disease)

Idiopathic
 C₃ nephritic factor–associated
 Partial lipodystrophy

Type III Disease

Idiopathic
 Complement receptor deficiency

Type I MPGN, the most proliferative of the three types, shows mesangial proliferation with lobular segmentation on renal biopsy and mesangial interposition between the capillary basement membrane and endothelial cells, producing a double contour sometimes called *tram-tracking* (see Fig. 62e-9). (See **Glomerular Schematic 3**.) Subendothelial deposits with low serum levels of C₃ are typical, although 50% of patients have normal levels of C₃ and occasional intramesangial deposits. Low serum C₃ and a dense thickening of the GBM containing ribbons of dense deposits and C₃ characterize type II MPGN, sometimes called *dense deposit disease* (see Fig. 62e-10). Classically, the glomerular tuft has a lobular appearance; intramesangial deposits are rarely present and subendothelial deposits are generally absent. Proliferation in type III MPGN is less common than the other two types and is often focal; mesangial interposition is rare, and subepithelial deposits can occur along widened segments of the GBM that appear laminated and disrupted.

Type I MPGN is secondary to glomerular deposition of circulating immune complexes or their in situ formation. Types II and III MPGN may be related to “nephritic factors,” which are autoantibodies that stabilize C₃ convertase and allow it to activate serum C₃; MPGN can also result from acquired or genetic abnormalities in the alternative complement pathway. Patients with MPGN present with proteinuria, hematuria, and pyuria (30%); systemic symptoms of fatigue and malaise that are most common in children with type I disease; or an acute nephritic picture with RPGN and a speedy deterioration in renal function in up to 25% of patients. Low serum C₃ levels are common. Fifty percent of patients with MPGN develop end-stage disease 10 years after

diagnosis, and 90% have renal insufficiency after 20 years. Nephrotic syndrome, hypertension, and renal insufficiency all predict poor outcome. In the presence of proteinuria, treatment with inhibitors of the renin-angiotensin system is prudent. Evidence for treatment with dipyridamole, Coumadin (warfarin), or cyclophosphamide is not strongly established. There is some evidence supporting the efficacy of treatment of *primary MPGN* with steroids, particularly in children, as well as reports of efficacy with plasma exchange and other immunosuppressive drugs. If defects in the complement pathway are found, treatment with eculizumab is of hypothetical but unproven benefit. In *secondary MPGN*, treating the associated infection, autoimmune disease, or neoplasms is of demonstrated benefit. In particular, pegylated interferon and ribavirin are useful in reducing viral load. Although all primary renal diseases can recur over time in transplanted renal allografts, patients with MPGN are well known to be at risk for not only a histologic recurrence but also a clinically significant recurrence with loss of graft function.

MESANGIOPROLIFERATIVE GLOMERULONEPHRITIS

Mesangioproliferative glomerulonephritis is characterized by expansion of the mesangium, sometimes associated with mesangial hypercellularity; thin, single contoured capillary walls; and mesangial immune deposits. Clinically, it can present with varying degrees of proteinuria and, commonly, hematuria. Mesangioproliferative disease may be seen in IgA nephropathy, *Plasmodium falciparum* malaria, resolving postinfectious glomerulonephritis, and class II nephritis from lupus, all of which can have a similar histologic appearance. With these secondary entities excluded, the diagnosis of *primary mesangioproliferative glomerulonephritis* is made in less than 15% of renal biopsies. As an immune-mediated renal lesion with deposits of IgM, C1q, and C₃, the clinical course is variable. Patients with isolated hematuria may have a very benign course, and those with heavy proteinuria occasionally progress to renal failure. There is little agreement on treatment, but some clinical reports suggest benefit from use of inhibitors of the renin-angiotensin system, steroid therapy, and even cytotoxic agents.

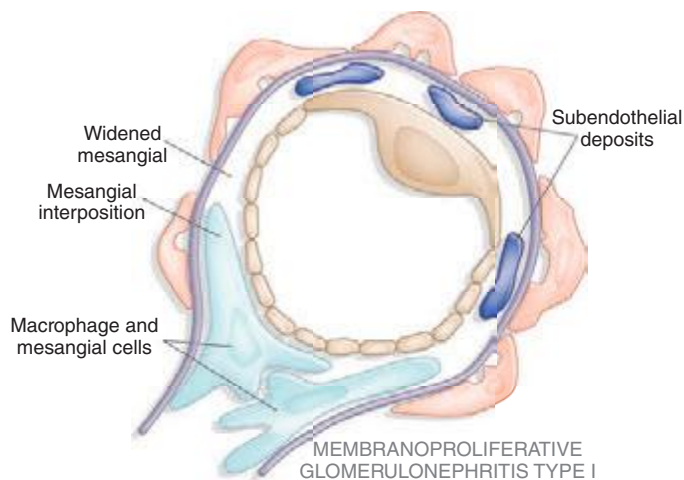
NEPHROTIC SYNDROME

Nephrotic syndrome classically presents with heavy proteinuria, minimal hematuria, hypoalbuminemia, hypercholesterolemia, edema, and hypertension. If left undiagnosed or untreated, some of these syndromes will progressively damage enough glomeruli to cause a fall in GFR, producing renal failure. Multiple studies have noted that the higher the 24-h urine protein excretion, the more rapid is the decline in GFR.

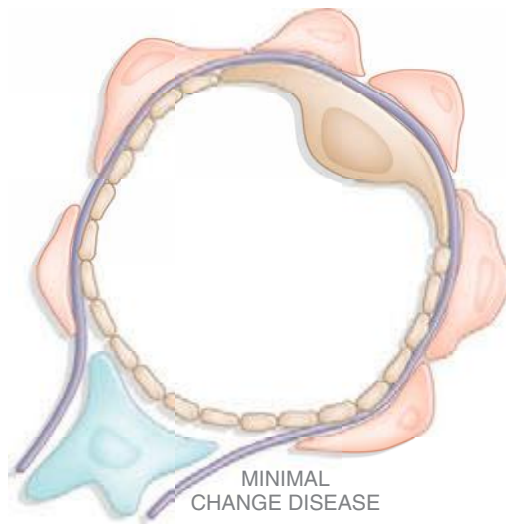
Therapies for various causes of nephrotic syndrome are noted under individual disease headings below. In general, all patients with hypercholesterolemia secondary to nephrotic syndrome should be treated with lipid-lowering agents because they are at increased risk for cardiovascular disease. Edema secondary to salt and water retention can be controlled with the judicious use of diuretics, avoiding intravascular volume depletion. Venous complications secondary to the hypercoagulable state associated with nephrotic syndrome can be treated with anticoagulants. The losses of various serum binding proteins, such as thyroid-binding globulin, lead to alterations in functional tests. Lastly, proteinuria itself is hypothesized to be nephrotoxic, and treatment of proteinuria with inhibitors of the renin-angiotensin system can lower urinary protein excretion.

MINIMAL CHANGE DISEASE

Minimal change disease (MCD), sometimes known as *nil lesion*, causes 70–90% of nephrotic syndrome in childhood but only 10–15% of nephrotic syndrome in adults. Minimal change disease usually presents as a primary renal disease but can be associated with several other conditions, including Hodgkin’s disease, allergies, or use of nonsteroidal anti-inflammatory agents; significant interstitial nephritis often accompanies cases associated with nonsteroidal drug use. Minimal change disease on renal biopsy shows no obvious glomerular lesion by light microscopy and is negative for deposits by immunofluorescent microscopy, or occasionally shows small amounts of IgM in the mesangium

Glomerular schematic 3

Glomerular schematic 4



(see Fig. 62e-1). (See Glomerular Schematic 4.) Electron microscopy, however, consistently demonstrates an effacement of the foot process supporting the epithelial podocytes with weakening of slit-pore membranes. The pathophysiology of this lesion is uncertain. Most agree there is a circulating cytokine, perhaps related to a T cell response that alters capillary charge and podocyte integrity. The evidence for cytokine-related immune injury is circumstantial and is suggested by the presence of preceding allergies, altered cell-mediated immunity during viral infections, and the high frequency of remissions with steroids.

Minimal change disease presents clinically with the abrupt onset of edema and nephrotic syndrome accompanied by acellular urinary sediment. Average urine protein excretion reported in 24 h is 10 g with severe hypoalbuminemia. Less common clinical features include hypertension (30% in children, 50% in adults), microscopic hematuria (20% in children, 33% in adults), atopy or allergic symptoms (40% in children, 30% in adults), and decreased renal function (<5% in children, 30% in adults). The appearance of acute renal failure in adults is often seen more commonly in patients with low serum albumin and intrarenal edema (nephrosarca) that is responsive to intravenous albumin and diuretics. This presentation must be distinguished from acute renal failure secondary to hypovolemia. Acute tubular necrosis and interstitial inflammation are also reported. In children, the abnormal urine principally contains albumin with minimal amounts of higher-molecular-weight proteins, and is sometimes called *selective proteinuria*. Although up to 30% of children have a spontaneous remission, all children today are treated with steroids; only children who are nonresponders are biopsied in this setting. Primary responders are patients who have a complete remission (<0.2 mg/24 h of proteinuria) after a single course of prednisone; steroid-dependent patients relapse as their steroid dose is tapered. Frequent relapsers have two or more relapses in the 6 months following taper, and steroid-resistant patients fail to respond to steroid therapy. Adults are not considered steroid-resistant until after 4 months of therapy. Ninety to 95% of children will develop a complete remission after 8 weeks of steroid therapy, and 80–85% of adults will achieve complete remission, but only after a longer course of 20–24 weeks. Patients with steroid resistance may have FSGS on repeat biopsy. Some hypothesize that if the first renal biopsy does not have a sample of deeper corticomedullary glomeruli, then the correct diagnosis of FSGS may be missed.

Relapses occur in 70–75% of children after the first remission, and early relapse predicts multiple subsequent relapses, as do high levels of basal proteinuria. The frequency of relapses decreases after puberty. There is an increased risk of relapse following the rapid tapering of steroids in all groups. Relapses are less common in adults but are more resistant to subsequent therapy. Prednisone is first-line therapy, either given daily or on alternate days. Other immunosuppressive drugs, such as cyclophosphamide, chlorambucil, and mycophenolate

mofetil, are saved for frequent relapsers, steroid-dependent patients, or steroid-resistant patients. Cyclosporine can induce remission, but relapse is also common when cyclosporine is withdrawn. The long-term prognosis in adults is less favorable when acute renal failure or steroid resistance occurs.

FOCAL SEGMENTAL GLOMERULOSCLEROSIS

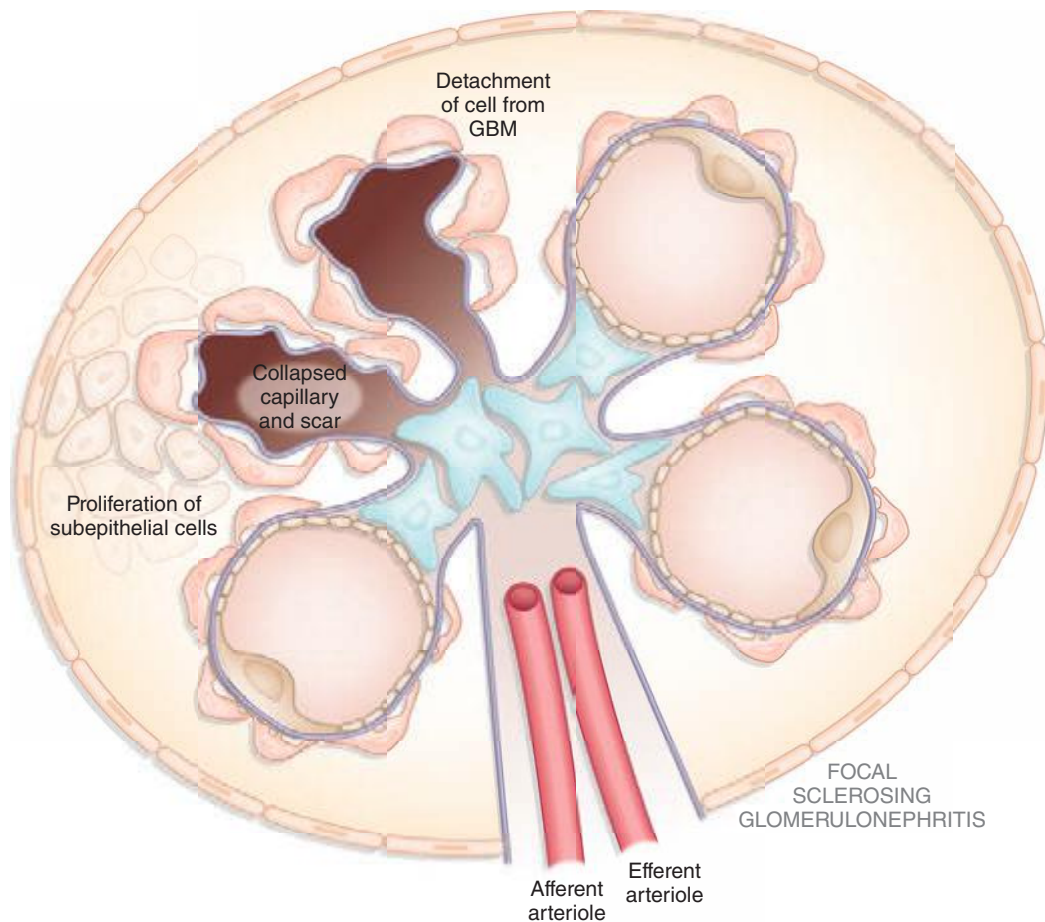
Focal segmental glomerulosclerosis (FSGS) refers to a pattern of renal injury characterized by segmental glomerular scars that involve some but not all glomeruli; the clinical findings of FSGS largely manifest as proteinuria. When the secondary causes of FSGS are eliminated (Table 338-5), the remaining patients are considered to have primary FSGS. The incidence of this disease is increasing, and it now represents up to one-third of cases of nephrotic syndrome in adults and one-half of cases of nephrotic syndrome in African Americans, in whom it is seen more commonly. The pathogenesis of FSGS is probably multifactorial. Possible mechanisms include a T cell-mediated circulating permeability factor, increased soluble urokinase receptor levels, TGF- β -mediated cellular proliferation and matrix synthesis, and podocyte abnormalities associated with genetic mutations. Risk polymorphisms at the *APOL1* locus encoding apolipoprotein L1 expressed in podocytes substantially explain the increased burden of FSGS among African Americans with or without HIV-associated disease.

The pathologic changes of FSGS are most prominent in glomeruli located at the corticomedullary junction (see Fig. 62e-2), so if the renal biopsy specimen is from superficial tissue, the lesions can be missed, which sometimes leads to a misdiagnosis of MCD. In addition to focal and segmental scarring, other variants have been described, including cellular lesions with *endocapillary hypercellularity* and heavy proteinuria; *collapsing glomerulopathy* (see Fig. 62e-3) with segmental or global glomerular collapse and a rapid decline in renal function; a hilar stalk lesion (see Fig. 62e-4); or the *glomerular tip lesion* (see Fig. 62e-5), which may have a better prognosis. (See Glomerular Schematic 5.)

FSGS can present with hematuria, hypertension, any level of proteinuria, or renal insufficiency. Nephrotic-range proteinuria, African-American race, and renal insufficiency are associated with a poor outcome, with 50% of patients reaching renal failure in 6–8 years. FSGS rarely remits spontaneously, but treatment-induced remission of proteinuria significantly improves prognosis. Treatment of patients with *primary FSGS* should include inhibitors of the renin-angiotensin system. Based on retrospective studies, patients with nephrotic-range proteinuria can be treated with steroids but respond far less often and

TABLE 338-5 FOCAL SEGMENTAL GLOMERULOSCLEROSIS

Primary focal segmental glomerulosclerosis
Secondary focal segmental glomerulosclerosis
Viruses: HIV/hepatitis B/parvovirus
Hypertensive nephropathy
Reflux nephropathy
Cholesterol emboli
Drugs: Heroin/analgesics/pamidronate
Oligomeganephronia
Renal dysgenesis
Alport's syndrome
Sickle cell disease
Lymphoma
Radiation nephritis
Familial podocytopathies
<i>NPHS1</i> mutation/nephrin
<i>NPHS2</i> mutation/podocin
<i>TRPC6</i> mutation/cation channel
<i>ACTN4</i> mutation/actinin
α -Galactosidase A deficiency/Fabry's disease
N-acetylneuraminic acid hydrolase deficiency/nephrosialidosis



after a longer course of therapy than patients with MCD. Proteinuria remits in only 20–45% of patients receiving a course of steroids over 6–9 months. Limited evidence suggests the use of cyclosporine in steroid-responsive patients helps ensure remissions. Relapse frequently occurs after cessation of cyclosporine therapy, and cyclosporine itself can lead to a deterioration of renal function due to its nephrotoxic effects. A role for other agents that suppress the immune system has not been established. Primary FSGS recurs in 25–40% of patients given allografts at end-stage disease, leading to graft loss in half of those cases. The treatment of *secondary FSGS* typically involves treating the underlying cause and controlling proteinuria. There is no role for steroids or other immunosuppressive agents in secondary FSGS.

MEMBRANOUS GLOMERULONEPHRITIS

Membranous glomerulonephritis (MGN), or *membranous nephropathy* as it is sometimes called, accounts for approximately 30% of cases of nephrotic syndrome in adults, with a peak incidence between the ages of 30 and 50 years and a male to female ratio of 2:1. It is rare in childhood and the most common cause of nephrotic syndrome in the elderly. In 25–30% of cases, MGN is associated with a malignancy (solid tumors of the breast, lung, colon), infection (hepatitis B, malaria, schistosomiasis), or rheumatologic disorders like lupus or rarely rheumatoid arthritis (**Table 338-6**).

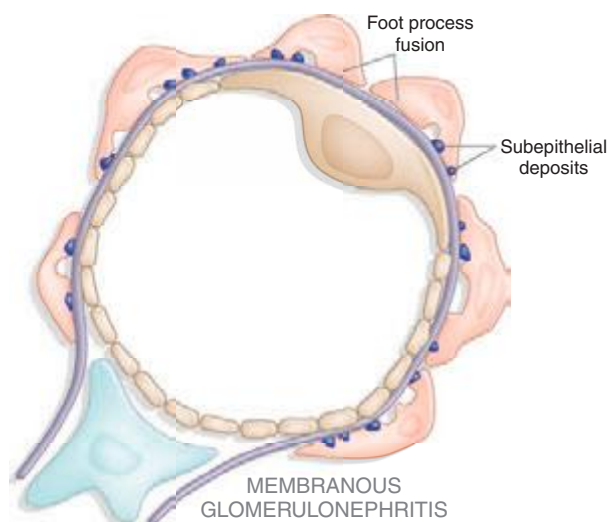
Uniform thickening of the basement membrane along the peripheral capillary loops is seen by light microscopy on renal biopsy (**see Fig. 62e-7**); this thickening needs to be distinguished from that seen in diabetes and amyloidosis. (**See Glomerular Schematic 6.**) Immunofluorescence demonstrates diffuse granular deposits of IgG and C₃, and electron microscopy typically reveals electron-dense subepithelial deposits. While different stages (I–V) of progressive membranous lesions have been described, some published analyses indicate the degree of tubular atrophy or interstitial fibrosis is more

predictive of progression than is the stage of glomerular disease. The presence of subendothelial deposits or the presence of tubuloreticular inclusions strongly points to a diagnosis of membranous lupus nephritis, which may precede the extrarenal manifestations of lupus. Work in Heyman nephritis, an animal model of MGN, suggests that glomerular lesions result from in situ formation of immune complexes with megalin receptor-associated protein as the putative antigen. This antigen is not found in human podocytes. Human antibodies have been described against neutral endopeptidase expressed by podocytes in infants whose mothers lack this protein. In most adults, autoantibodies against the M-type phospholipase A₂ receptor (PLA₂R) circulate and bind to a conformational epitope present in the receptor on human podocytes, producing in situ deposits characteristic of idiopathic membranous nephropathy. Other renal diseases and secondary membranous nephropathy do not appear to involve such

TABLE 338-6 MEMBRANOUS GLOMERULONEPHRITIS

Primary/idiopathic membranous glomerulonephritis
Secondary membranous glomerulonephritis
Infection: Hepatitis B and C, syphilis, malaria, schistosomiasis, leprosy, filariasis
Cancer: Breast, colon, lung, stomach, kidney, esophagus, neuroblastoma
Drugs: Gold, mercury, penicillamine, nonsteroidal anti-inflammatory agents, probenecid
Autoimmune diseases: Systemic lupus erythematosus, rheumatoid arthritis, primary biliary cirrhosis, dermatitis herpetiformis, bullous pemphigoid, myasthenia gravis, Sjögren's syndrome, Hashimoto's thyroiditis
Other systemic diseases: Fanconi's syndrome, sickle cell anemia, diabetes, Crohn's disease, sarcoidosis, Guillain-Barré syndrome, Weber-Christian disease, angiofollicular lymph node hyperplasia

Glomerular schematic 6



autoantibodies and levels of these autoantibodies have correlated with the severity of MGN. Eighty percent of patients with MGN present with nephrotic syndrome and nonselective proteinuria. Microscopic hematuria is seen but less commonly than in IgA nephropathy or FSGS. Spontaneous remissions occur in 20–33% of patients and often occur late in the course after years of nephrotic syndrome, which make treatment decisions difficult. One-third of patients continue to have relapsing nephrotic syndrome but maintain normal renal function, and approximately another third of patients develop renal failure or die from the complications of nephrotic syndrome. Male gender, older age, hypertension, and the persistence of proteinuria are associated with worse prognosis. Although thrombotic complications are a feature of all nephrotic syndromes, MGN has the highest reported incidences of renal vein thrombosis, pulmonary embolism, and deep vein thrombosis. Prophylactic anticoagulation is controversial but has been recommended for patients with severe or prolonged proteinuria in the absence of risk factors for bleeding.

In addition to the treatment of edema, dyslipidemia, and hypertension, inhibition of the renin-angiotensin system is recommended. Therapy with immunosuppressive drugs is also recommended for patients with primary MGN and persistent proteinuria (>3.0 g/24 h). The choice of immunosuppressive drugs for therapy is controversial, but current recommendations are to treat with steroids and cyclophosphamide, chlorambucil, mycophenolate mofetil, or cyclosporine. In patients who relapse or fail to respond to this therapy, the use of rituximab, an anti-CD20 antibody directed at B cells, or synthetic adrenocorticotropic hormone may be considered.

DIABETIC NEPHROPATHY

Diabetic nephropathy is the single most common cause of chronic renal failure in the United States, accounting for 45% of patients receiving renal replacement therapy, and is a rapidly growing problem worldwide. The dramatic increase in the number of patients with diabetic nephropathy reflects the epidemic increase in obesity, metabolic syndrome, and type 2 diabetes mellitus. Approximately 40% of patients with types 1 or 2 diabetes develop nephropathy, but due to the higher prevalence of type 2 diabetes (90%) compared to type 1 (10%), the majority of patients with diabetic nephropathy have type 2 disease. Renal lesions are more common in African-American, Native American, Polynesian, and Maori populations. Risk factors for the development of diabetic nephropathy include hyperglycemia, hypertension, dyslipidemia, smoking, a family history of diabetic nephropathy, and gene polymorphisms affecting the activity of the renin-angiotensin-aldosterone axis.

Within 1–2 years after the onset of clinical diabetes, morphologic changes appear in the kidney. Thickening of the GBM is a sensitive indicator for the presence of diabetes but correlates poorly with

the presence or absence of clinically significant nephropathy. The composition of the GBM is altered notably with a loss of heparan sulfate moieties that form the negatively charged filtration barrier. This change results in increased filtration of serum proteins into the urine, predominately negatively charged albumin. The expansion of the mesangium due to the accumulation of extracellular matrix correlates with the clinical manifestations of diabetic nephropathy (see stages in Fig. 62e-20). This expansion in mesangial matrix is associated with the development of *mesangial sclerosis*. Some patients also develop eosinophilic, PAS⁺ nodules called *nodular glomerulosclerosis* or *Kimmelstiel-Wilson nodules*. Immunofluorescence microscopy often reveals the nonspecific deposition of IgG (at times in a linear pattern) or complement staining without immune deposits on electron microscopy. Prominent vascular changes are frequently seen with hyaline and hypertensive arteriosclerosis. This is associated with varying degrees of chronic glomerulosclerosis and tubulointerstitial changes. Renal biopsies from patients with types 1 and 2 diabetes are largely indistinguishable.

These pathologic changes are the result of a number of postulated factors. Multiple lines of evidence support an important role for increases in glomerular capillary pressure (intraglomerular hypertension) in alterations in renal structure and function. Direct effects of hyperglycemia on the actin cytoskeleton of renal mesangial and vascular smooth-muscle cells as well as diabetes-associated changes in circulating factors such as atrial natriuretic factor, angiotensin II, and insulin-like growth factor (IGF) may account for this. Sustained glomerular hypertension increases matrix production, alterations in the GBM with disruption in the filtration barrier (and hence proteinuria), and glomerulosclerosis. A number of factors have also been identified that alter matrix production, including the accumulation of advanced glycosylation end products, circulating factors including growth hormone, IGF-I, angiotensin II, connective tissue growth factor, TGF- β , and dyslipidemia.

The natural history of diabetic nephropathy in patients with types 1 and 2 diabetes is similar. However, since the onset of type 1 diabetes is readily identifiable and the onset of type 2 diabetes is not, a patient newly diagnosed with type 2 diabetes may present with *advanced diabetic nephropathy*. At the onset of diabetes, renal hypertrophy and glomerular hyperfiltration are present. The degree of glomerular hyperfiltration correlates with the subsequent risk of clinically significant nephropathy. In the approximately 40% of patients with diabetes who develop diabetic nephropathy, the earliest manifestation is an increase in albuminuria detected by sensitive radioimmunoassay (Table 338-1). Albuminuria in the range of 30–300 mg/24 h is called *microalbuminuria*. Microalbuminuria appears 5–10 years after the onset of diabetes. It is currently recommended to test patients with type 1 disease for microalbuminuria 5 years after diagnosis of diabetes and yearly thereafter and, because the time of onset of type 2 diabetes is often unknown, to test type 2 patients at the time of diagnosis of diabetes and yearly thereafter.

Patients with small increases in albuminuria increase their levels of urinary albumin excretion, typically reaching dipstick positive levels of proteinuria (>300 mg albuminuria) 5–10 years after the onset of early albuminuria. Microalbuminuria is a potent risk factor for cardiovascular events and death in patients with type 2 diabetes. Many patients with type 2 diabetes and microalbuminuria succumb to cardiovascular events before they progress to proteinuria or renal failure. Proteinuria in frank diabetic nephropathy can be variable, ranging from 500 mg to 25 g/24 h, and is often associated with nephrotic syndrome. More than 90% of patients with type 1 diabetes and nephropathy have diabetic retinopathy, so the absence of retinopathy in type 1 patients with proteinuria should prompt consideration of a diagnosis other than diabetic nephropathy; only 60% of patients with type 2 diabetes with nephropathy have diabetic retinopathy. There is a significant correlation between the presence of retinopathy and the presence of Kimmelstiel-Wilson nodules (see Fig. 62e-20). Also, characteristically, patients with advanced diabetic nephropathy have normal to enlarged kidneys, in contrast to other glomerular diseases where kidney size is usually decreased. Using the above epidemiologic and clinical data, and

in the absence of other clinical or serologic data suggesting another disease, diabetic nephropathy is usually diagnosed without a renal biopsy. After the onset of proteinuria, renal function inexorably declines, with 50% of patients reaching renal failure over another 5–10 years; thus, from the earliest stages of microalbuminuria, it usually takes 10–20 years to reach end-stage renal disease. Once renal failure appears, however, survival on dialysis is shorter for patients with diabetes compared to other dialysis patients. Survival is best for patients with type 1 diabetes who receive a transplant from a living related donor.

Good evidence supports the benefits of blood sugar and blood pressure control as well as inhibition of the renin-angiotensin system in retarding the progression of diabetic nephropathy. In patients with type 1 diabetes, intensive control of blood sugar clearly prevents the development or progression of diabetic nephropathy. The evidence for benefit of intensive blood glucose control in patients with type 2 diabetes is less certain, with current studies reporting conflicting results.

Controlling systemic blood pressure decreases renal and cardiovascular adverse events in this high-risk population. The vast majority of patients with diabetic nephropathy require three or more antihypertensive drugs to achieve this goal. Drugs that inhibit the renin-angiotensin system, independent of their effects on systemic blood pressure, have been shown in numerous large clinical trials to slow the progression of diabetic nephropathy at early (microalbuminuria) and late (proteinuria with reduced glomerular filtration) stages, independent of any effect they may have on systemic blood pressure. Since angiotensin II increases efferent arteriolar resistance and, hence, glomerular capillary pressure, one key mechanism for the efficacy of ACE inhibitors or angiotensin receptor blockers (ARBs) is reducing glomerular hypertension. Patients with type 1 diabetes for 5 years who develop albuminuria or declining renal function should be treated with ACE inhibitors. Patients with type 2 diabetes and microalbuminuria or proteinuria may be treated with ACE inhibitors or ARBs. Evidence suggests increased risk for cardiovascular adverse events in some patients with a combination of two drugs (ACE inhibitors, ARBs, renin inhibitors, or aldosterone antagonists) that suppress several components of the renin-angiotensin system.

GLOMERULAR DEPOSITION DISEASES

Plasma cell dyscrasias producing excess light chain immunoglobulin sometimes lead to the formation of glomerular and tubular deposits that cause heavy proteinuria and renal failure; the same is true for the accumulation of serum amyloid A protein fragments seen in several inflammatory diseases. This broad group of proteinuric patients has *glomerular deposition disease*.

Light Chain Deposition Disease The biochemical characteristics of nephrotoxic light chains produced in patients with light chain malignancies often confer a specific pattern of renal injury; that of either *cast nephropathy* (see Fig. 62e-17), which causes renal failure but not heavy proteinuria or amyloidosis, or light chain deposition disease (see Fig. 62e-16), which produces nephrotic syndrome with renal failure. These latter patients produce kappa light chains that do not have the biochemical features necessary to form amyloid fibrils. Instead, they self-aggregate and form granular deposits along the glomerular capillary and mesangium, tubular basement membrane, and Bowman's capsule. When predominant in glomeruli, nephrotic syndrome develops, and about 70% of patients progress to dialysis. Light-chain deposits are not fibrillar and do not stain with Congo red, but they are easily detected with anti-light chain antibody using immunofluorescence or as granular deposits on electron microscopy. A combination of the light chain rearrangement, self-aggregating properties at neutral pH, and abnormal metabolism probably contribute to the deposition. Treatment for light chain deposition disease is treatment of the primary disease and, if possible, autologous stem cell transplantation.

Renal Amyloidosis Most *renal amyloidosis* is either the result of primary fibrillar deposits of immunoglobulin light chains known as amyloid L (AL), or secondary to fibrillar deposits of serum amyloid A (AA) protein fragments (Chap. 137). Even though both occur for

different reasons, their clinicopathophysiology is quite similar and will be discussed together. Amyloid infiltrates the liver, heart, peripheral nerves, carpal tunnel, upper pharynx, and kidney, producing restrictive cardiomyopathy, hepatomegaly, macroglossia, and heavy proteinuria sometimes associated with renal vein thrombosis. In systemic AL amyloidosis, also called *primary amyloidosis*, light chains produced in excess by clonal plasma cell dyscrasias are made into fragments by macrophages so they can self-aggregate at acid pH. A disproportionate number of these light chains (75%) are of the *lambda* class. About 10% of these patients have overt myeloma with lytic bone lesions and infiltration of the bone marrow with >30% plasma cells; nephrotic syndrome is common, and about 20% of patients progress to dialysis. AA amyloidosis is sometimes called *secondary amyloidosis* and also presents as nephrotic syndrome. It is due to deposition of β -pleated sheets of serum amyloid A protein, an acute phase reactant whose physiologic functions include cholesterol transport, immune cell attraction, and metalloproteases activation. Forty percent of patients with AA amyloid have rheumatoid arthritis, and another 10% have ankylosing spondylitis or psoriatic arthritis; the rest derive from other lesser causes. Less common in Western countries but more common in Mediterranean regions, particularly in Sephardic and Iraqi Jews, is familial Mediterranean fever (FMF). FMF is caused by a mutation in the gene encoding pyrin, whereas Muckle-Wells syndrome, a related disorder, results from a mutation in cryopyrin; both proteins are important in the apoptosis of leukocytes early in inflammation; such proteins with pyrin domains are part of a new pathway called the *inflammasome*. Receptor mutations in tumor necrosis factor receptor 1 (TNFR1)-associated periodic syndrome also produce chronic inflammation and secondary amyloidosis. Fragments of serum amyloid A protein increase and self-aggregate by attaching to receptors for advanced glycation end products in the extracellular environment; nephrotic syndrome is common, and about 40–60% of patients progress to dialysis. AA and AL amyloid fibrils are detectable with Congo red or in more detail with electron microscopy (see Fig. 62e-15). Currently developed serum free light chain nephelometry assays are useful in the early diagnosis and follow-up of disease progression. Biopsy of involved liver or kidney is diagnostic 90% of the time when the pretest probability is high; abdominal fat pad aspirates are positive about 70% of the time, but apparently less so when looking for AA amyloid. Amyloid deposits are distributed along blood vessels and in the mesangial regions of the kidney. The treatment for primary amyloidosis, melphalan and autologous hematopoietic stem cell transplantation, can delay the course of disease in about 30% of patients. Secondary amyloidosis is also relentless unless the primary disease can be controlled. Some new drugs in development that disrupt the formation of fibrils may be available in the future.

Fibrillary-Immunotactoid Glomerulopathy Fibrillary-immunotactoid glomerulopathy is a rare (<1.0% of renal biopsies), morphologically defined disease characterized by glomerular accumulation of nonbranching randomly arranged fibrils. Some classify amyloid and nonamyloid fibril-associated renal diseases all as fibrillary glomerulopathies with immunotactoid glomerulopathy reserved for nonamyloid fibrillary disease not associated with a systemic illness. Others define fibrillary glomerulonephritis as a nonamyloid fibrillary disease with fibrils 12–24 nm and immunotactoid glomerulonephritis with fibrils >30 nm. In either case, fibrillar/microtubular deposits of oligoclonal or oligotypic immunoglobulins and complement appear in the mesangium and along the glomerular capillary wall. Congo red stains are negative. The cause of this “nonamyloid” glomerulopathy is mostly idiopathic; reports of immunotactoid glomerulonephritis describe an occasional association with chronic lymphocytic leukemia or B cell lymphoma. Both disorders appear in adults in the fourth decade with moderate to heavy proteinuria, hematuria, and a wide variety of histologic lesions, including DPGN, MPGN, MGN, or mesangioproliferative glomerulonephritis. Nearly half of patients develop renal failure over a few years. There is no consensus on treatment of this uncommon disorder. The disease has been reported to recur following renal transplantation in a minority of cases.

Fabry's disease is an X-linked inborn error of globotriaosylceramide metabolism secondary to deficient lysosomal α -galactosidase A activity, resulting in excessive intracellular storage of globotriaosylceramide. Affected organs include the vascular endothelium, heart, brain, and kidneys. Classically, Fabry's disease presents in childhood in males with acroparesthesias, angiokeratoma, and hypohidrosis. Over time male patients develop cardiomyopathy, cerebrovascular disease, and renal injury, with an average age of death around 50 years of age. Hemizygotes with hypomorphic mutations sometimes present in the fourth to sixth decade with single-organ involvement. Rarely, dominant-negative α -galactosidase A mutations or female heterozygotes with unfavorable X inactivation present with mild single-organ involvement. Rare females develop severe manifestations including renal failure but do so later in life than males. Renal biopsy reveals enlarged glomerular visceral epithelial cells packed with small clear vacuoles containing globotriaosylceramide; vacuoles may also be found in parietal and tubular epithelia (see Fig. 62e-18). These vacuoles of electron-dense materials in parallel arrays (zebra bodies) are easily seen on electron microscopy. Ultimately, renal biopsies reveal FSGS. The nephropathy of Fabry's disease typically presents in the third decade as mild to moderate proteinuria, sometimes with microscopic hematuria or nephrotic syndrome. Urinalysis may reveal oval fat bodies and birefringent glycolipid globules under polarized light (Maltese cross). Renal biopsy is necessary for definitive diagnosis. Progression to renal failure occurs by the fourth or fifth decade. Treatment with inhibitors of the renin-angiotensin system is recommended. Treatment with recombinant α -galactosidase A clears microvascular endothelial deposits of globotriaosylceramide from the kidneys, heart, and skin. In patients with advanced organ involvement, progression of disease occurs despite enzyme replacement therapy. Variable responses to enzyme therapy may be due to the occurrence of neutralizing antibodies or differences in uptake of the

enzyme. Graft and patient survival following renal transplantation in patients with Fabry's are similar to other causes of end-stage renal disease.

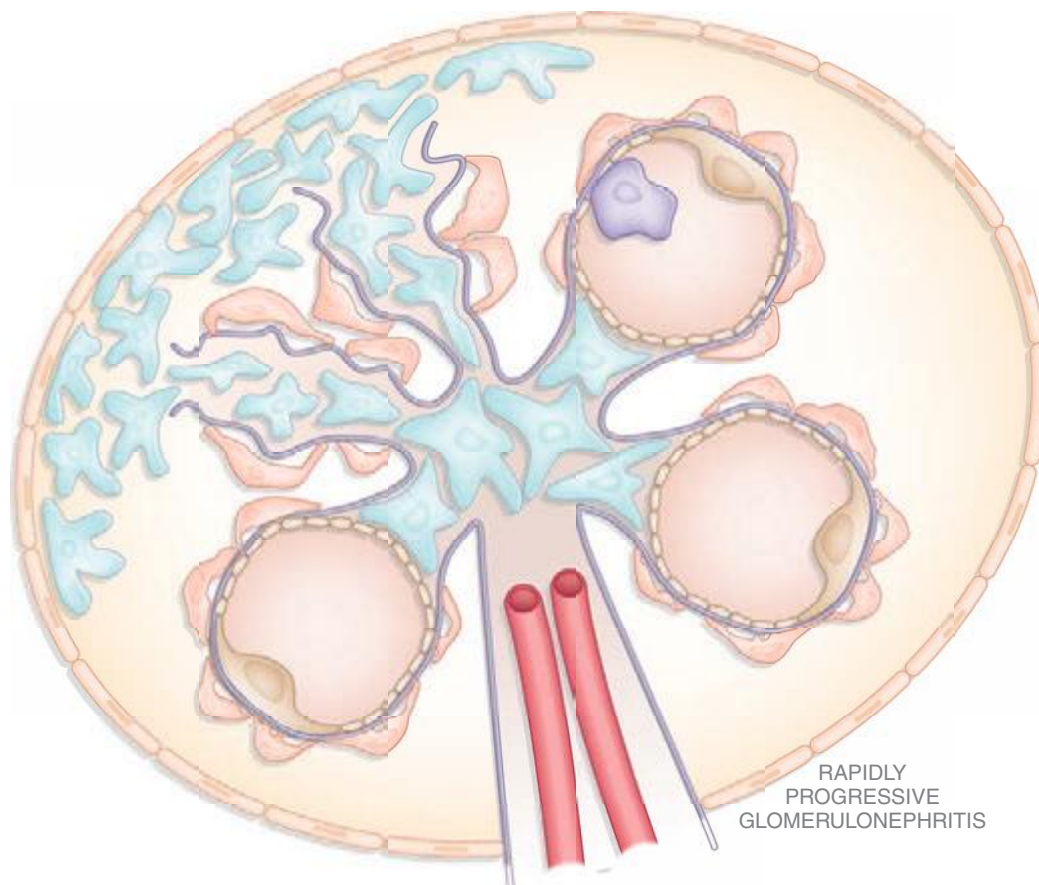
PULMONARY-RENAL SYNDROMES

Several diseases can present with catastrophic hemoptysis and glomerulonephritis associated with varying degrees of renal failure. The usual causes include Goodpasture's syndrome, granulomatosis with polyangiitis, microscopic polyangiitis, Churg-Strauss vasculitis, and, rarely, Henoch-Schönlein purpura or cryoglobulinemia. Each of these diseases can also present without hemoptysis and are discussed in detail earlier in "Acute Nephritic Syndromes." (See **Glomerular Schematic 7.**) Pulmonary bleeding in this setting is life-threatening and often results in airway intubation, and acute renal failure requires dialysis. Diagnosis is difficult initially because biopsies and serologic testing take time. Treatment with plasmapheresis and methylprednisolone is often empirical and temporizing until results of testing are available.

BASEMENT MEMBRANE SYNDROMES

All kidney epithelia, including podocytes, rest on basement membranes assembled into a planar surface through the interweaving of collagen IV with laminins, nidogen, and sulfated proteoglycans. Structural abnormalities in GBM associated with hematuria are characteristic of several familial disorders related to the expression of collagen IV genes. The extended family of collagen IV contains six chains, which are expressed in different tissues at different stages of embryonic development. All epithelial basement membranes early in human development are composed of interconnected triple-helical protomers rich in $\alpha 1(\text{IV})$ collagen. Some specialized tissues undergo a developmental switch replacing $\alpha 1(\text{IV})$ protomers with an $\alpha 3(\text{IV})$ collagen network; this switch occurs in the kidney (glomerular and tubular basement membrane), lung, testis, cochlea, and eye, while an

Glomerular schematic 7



$\alpha 5, \alpha 5, \alpha 6(IV)$ network appears in skin, smooth muscle, and esophagus and along Bowman's capsule in the kidney. This switch probably occurs because the $\alpha 3, \alpha 4, \alpha 5(IV)$ network is more resistant to proteases and ensures the structural longevity of critical tissues. When basement membranes are the target of glomerular disease, they produce moderate proteinuria, some hematuria, and progressive renal failure.

ANTI-GBM DISEASE

Autoimmune disease where antibodies are directed against the $\alpha 3$ NC1 domain of collagen IV produces an *anti-GBM disease* often associated with RPGN and/or a pulmonary-renal syndrome called *Goodpasture's syndrome*. Discussion of this disease is covered earlier in "Acute Nephritic Syndromes."

ALPORT'S SYNDROME

Classically, patients with Alport's syndrome develop hematuria, thinning and splitting of the GBMs, mild proteinuria ($<1-2$ g/24 h), which appears late in the course, followed by chronic glomerulosclerosis leading to renal failure in association with sensorineural deafness. Some patients develop lenticonus of the anterior lens capsule, "dot and fleck" retinopathy, and rarely, mental retardation or leiomyomatosis. Approximately 85% of patients with Alport's syndrome have an X-linked inheritance of mutations in the $\alpha 5(IV)$ collagen chain on chromosome Xq22-24. Female carriers have variable penetrance depending on the type of mutation or the degree of mosaicism created by X inactivation. Fifteen percent of patients have autosomal recessive disease of the $\alpha 3(IV)$ or $\alpha 4(IV)$ chains on chromosome 2q35-37. Rarely, some kindred have an autosomal dominant inheritance of dominant-negative mutations in $\alpha 3(IV)$ or $\alpha 4(IV)$ chains.

Pedigrees with the X-linked syndrome are quite variable in their rate and frequency of tissue damage leading to organ failure. Seventy percent of patients have the juvenile form with nonsense or missense mutations, reading frame shifts, or large deletions and generally develop renal failure and sensorineural deafness by age 30. Patients with splice variants, exon skipping, or missense mutations of α -helical glycines generally deteriorate after the age of 30 (adult form) with mild or late deafness. Early severe deafness, lenticonus, or proteinuria suggests a poorer prognosis. Usually females from X-linked pedigrees have only microhematuria, but up to 25% of carrier females have been reported to have more severe renal manifestations. Pedigrees with the autosomal recessive form of the disease have severe early disease in both females and males with asymptomatic parents.

Clinical evaluation should include a careful eye examination and hearing tests. However, the absence of extrarenal symptoms does not rule out the diagnosis. Since $\alpha 5(IV)$ collagen is expressed in the skin, some X-linked Alport's patients can be diagnosed with a skin biopsy revealing the lack of the $\alpha 5(IV)$ collagen chain on immunofluorescent analysis. Patients with mutations in $\alpha 3(IV)$ or $\alpha 4(IV)$ require a renal biopsy. Genetic testing can be used for the diagnosis of Alport's syndrome and the demonstration of the mode of inheritance. Early in their disease, Alport's patients typically have thin basement membranes on renal biopsy (see Fig. 62e-19), which thicken over time into multilamellations surrounding lucent areas that often contain granules of varying density—the so-called split basement membrane. In any Alport's kidney, there are areas of thinning mixed with splitting of the GBM. Tubules drop out, glomeruli scar, and the kidney eventually succumbs to interstitial fibrosis. All affected members of a family with X-linked Alport's syndrome should be identified and followed, including mothers of affected males. Primary treatment is control of systemic hypertension and use of ACE inhibitors to slow renal progression. Although patients who receive renal allografts usually develop anti-GBM antibodies directed toward the collagen epitopes absent in their native kidney, overt Goodpasture's syndrome is rare and graft survival is good.

THIN BASEMENT MEMBRANE DISEASE

Thin basement membrane disease (TBMD) characterized by persistent or recurrent hematuria is not typically associated with proteinuria, hypertension, or loss of renal function or extrarenal disease. Although

not all cases are familial (perhaps a founder effect), it usually presents in childhood in multiple family members and is also called *benign familial hematuria*. Cases of TBMD have genetic defects in type IV collagen but in contrast to Alport behave as an autosomal dominant disorder that in ~40% of families segregates with the *COL(IV) $\alpha 3$ /COL(IV) $\alpha 4$* loci. Mutations in these loci can result in a spectrum of disease ranging from TBMD to autosomal dominant or recessive Alport's. The GBM shows diffuse thinning compared to normal values for the patient's age in otherwise normal biopsies (see Fig. 62e-19). The vast majority of patients have a benign course.

NAIL-PATELLA SYNDROME

Patients with nail-patella syndrome develop iliac horns on the pelvis and dysplasia of the dorsal limbs involving the patella, elbows, and nails, variably associated with neural-sensory hearing impairment, glaucoma, and abnormalities of the GBM and podocytes, leading to hematuria, proteinuria, and FSGS. The syndrome is autosomal dominant, with haploinsufficiency for the LIM homeodomain transcription factor LMX1B; pedigrees are extremely variable in the penetrance for all features of the disease. LMX1B regulates the expression of genes encoding $\alpha 3$ and $\alpha 4$ chains of collagen IV, interstitial type III collagen, podocin, and CD2AP that help form the slit-pore membranes connecting podocytes. Mutations in the LIM domain region of LMX1B associate with glomerulopathy, and renal failure appears in as many as 30% of patients. Proteinuria or isolated hematuria is discovered throughout life, but usually by the third decade, and is inexplicably more common in females. On renal biopsy there is lucent damage to the lamina densa of the GBM, an increase in collagen III fibrils along glomerular capillaries and in the mesangium, and damage to the slit-pore membrane, producing heavy proteinuria not unlike that seen in congenital nephrotic syndrome. Patients with renal failure do well with transplantation.

GLOMERULAR-VASCULAR SYNDROMES

A variety of diseases result in classic vascular injury to the glomerular capillaries. Most of these processes also damage blood vessels elsewhere in the body. The group of diseases discussed here lead to vasculitis, renal endothelial injury, thrombosis, ischemia, and/or lipid-based occlusions.

ATHEROSCLEROTIC NEPHROPATHY

Aging in the developed world is commonly associated with the occlusion of coronary and systemic blood vessels. The reasons for this include obesity, insulin resistance, smoking, hypertension, and diets rich in lipids that deposit in the arterial and arteriolar circulation, producing local inflammation and fibrosis of small blood vessels. When the renal arterial circulation is involved, the glomerular microcirculation is damaged, leading to *chronic nephrosclerosis*. Patients with GFRs <60 mL/min have more cardiovascular events and hospitalizations than those with higher filtration rates. Several aggressive lipid disorders can accelerate this process, but most of the time atherosclerotic progression to chronic nephrosclerosis is associated with poorly controlled hypertension. Approximately 10% of glomeruli are normally sclerotic by age 40, rising to 20% by age 60 and 30% by age 80. Serum lipid profiles in humans are greatly affected by *apolipoprotein E* polymorphisms; the E4 allele is accompanied by increases in serum cholesterol and is more closely associated with atherogenic profiles in patients with renal failure. Mutations in E2 alleles, particularly in Japanese patients, produce a specific renal abnormality called *lipoprotein glomerulopathy* associated with glomerular lipoprotein thrombi and capillary dilation.

HYPERTENSIVE NEPHROSCLEROSIS

Uncontrolled systemic hypertension causes permanent damage to the kidneys in about 6% of patients with elevated blood pressure. As many as 27% of patients with end-stage kidney disease have hypertension as a primary cause. Although there is not a clear correlation between the extent or duration of hypertension and the risk of end-organ damage, *hypertensive nephrosclerosis* is fivefold more frequent in African Americans than whites. Risk alleles associated with *APOLL1*, a functional gene for apolipoprotein L1 expressed in podocytes

substantially explains the increased burden of end-stage renal disease among African Americans. Associated risk factors for progression to end-stage kidney disease include increased age, male gender, race, smoking, hypercholesterolemia, duration of hypertension, low birth weight, and preexisting renal injury. Kidney biopsies in patients with hypertension, microhematuria, and moderate proteinuria demonstrate arteriosclerosis, chronic nephrosclerosis, and interstitial fibrosis in the absence of immune deposits (see Fig. 62e-21). Today, based on a careful history, physical examination, urinalysis, and some serologic testing, the diagnosis of chronic nephrosclerosis is usually inferred without a biopsy. Treating hypertension is the best way to avoid progressive renal failure; most guidelines recommend lowering blood pressure to <130/80 mmHg if there is preexisting diabetes or kidney disease. In the presence of kidney disease, most patients begin antihypertensive therapy with two drugs, classically a thiazide diuretic and an ACE inhibitor; most will require three drugs. There is strong evidence in African Americans with hypertensive nephrosclerosis that therapy initiated with an ACE inhibitor can slow the rate of decline in renal function independent of effects on systemic blood pressure. Malignant acceleration of hypertension complicates the course of chronic nephrosclerosis, particularly in the setting of scleroderma or cocaine use (see Fig. 62e-24). The hemodynamic stress of malignant hypertension leads to fibrinoid necrosis of small blood vessels, thrombotic microangiopathy, a nephritic urinalysis, and acute renal failure. In the setting of renal failure, chest pain, or papilledema, the condition is treated as a hypertensive emergency. Slightly lowering the blood pressure often produces an immediate reduction in GFR that improves as the vascular injury attenuates and autoregulation of blood vessel tone is restored.

CHOLESTEROL EMBOLI

Aging patients with clinical complications from atherosclerosis sometimes shower cholesterol crystals into the circulation—either spontaneously or, more commonly, following an endovascular procedure with manipulation of the aorta—or with use of systemic anticoagulation. Spontaneous emboli may shower acutely or shower subacutely and somewhat more silently. Irregular emboli trapped in the microcirculation produce ischemic damage that induces an inflammatory reaction. Depending on the location of the atherosclerotic plaques releasing these cholesterol fragments, one may see cerebral transient ischemic attacks; livedo reticularis in the lower extremities; Hollenhorst plaques in the retina with visual field cuts; necrosis of the toes; and acute glomerular capillary injury leading to *focal segmental glomerulosclerosis* sometimes associated with hematuria, mild proteinuria, and loss of renal function, which typically progresses over a few years. Occasional patients have fever, eosinophilia, or eosinophiluria. A skin biopsy of an involved area may be diagnostic. Since tissue fixation dissolves the cholesterol, one typically sees only residual, biconvex clefts in involved vessels (see Fig. 62e-22). There is no therapy to reverse embolic occlusions, and steroids do not help. Controlling blood pressure and lipids and cessation of smoking are usually recommended for prevention.

SICKLE CELL DISEASE

Although individuals with SA-hemoglobin are usually asymptomatic, most will gradually develop hyposthenuria due to subclinical infarction of the renal medulla, thus predisposing them to volume depletion. There is an unexpectedly high prevalence of sickle trait among dialysis patients who are African American. Patients with homozygous SS-sickle cell disease develop chronic vasoocclusive disease in many organs. Polymers of deoxygenated SS-hemoglobin distort the shape of red blood cells. These cells attach to endothelia and obstruct small blood vessels, producing frequent and painful sickle cell crises over time. Vessel occlusions in the kidney produce glomerular hypertension, FSGS, interstitial nephritis, and renal infarction associated with hyposthenuria, microscopic hematuria, and even gross hematuria; some patients also present with MPGN. Renal function can be overestimated due to the increased tubular secretion of creatinine seen in many patients with SS-sickle cell. By the second or third decade of life, persistent vasoocclusive disease in the kidney leads to varying degrees of renal failure, and some patients end up on

dialysis. Treatment is directed to reducing the frequency of painful crises and administering ACE inhibitors in the hope of delaying a progressive decline in renal function. In sickle cell patients undergoing renal transplantation, renal graft survival is comparable to African Americans in the general transplant population.

THROMBOTIC MICROANGIOPATHIES


Thrombotic thrombocytopenic purpura (TTP) and *hemolytic-uremic syndrome* (HUS) represent a spectrum of thrombotic microangiopathies. Thrombotic thrombocytopenic purpura and hemolytic-uremic syndrome share the general features of idiopathic thrombocytopenic purpura, hemolytic anemia, fever, renal failure, and neurologic disturbances. When patients, particularly children, have more evidence of renal injury, their condition tends to be called HUS. In adults with neurologic disease, it is considered to be TTP. In adults there is often a mixture of both, which is why they are often referred to as having TTP/HUS. On examination of kidney tissue, there is evidence of *glomerular capillary endotheliosis* associated with platelet thrombi, damage to the capillary wall, and formation of fibrin material in and around glomeruli (see Fig. 62e-23). These tissue findings are similar to what is seen in preclampsia/HELLP (hemolysis, elevated liver enzymes, and low platelet count syndrome), malignant hypertension, and the antiphospholipid syndrome. TTP/HUS is also seen in pregnancy; with the use of oral contraceptives or quinine; in renal transplant patients given OKT3 for rejection; in patients taking the calcineurin inhibitors, cyclosporine and tacrolimus, or in patients taking the antiplatelet agents, ticlopidine and clopidogrel; or following HIV infection.

Although there is no agreement on how much they share a final common pathophysiology, two general groups of patients are recognized: childhood HUS associated with enterohemorrhagic diarrhea and TTP/HUS in adults. Childhood HUS is caused by a toxin released by *Escherichia coli* 0157:H7 and occasionally by *Shigella dysenteriae*. This shiga toxin (verotoxin) directly injures endothelia, enterocytes, and renal cells, causing apoptosis, platelet clumping, and intravascular hemolysis by binding to the glycolipid receptors (Gb3). These receptors are more abundant along endothelia in children compared to adults. Shiga toxin also inhibits the endothelial production of ADAMTS13. In familial cases of adult TTP/HUS, there is a genetic deficiency of the ADAMTS13 metalloprotease that cleaves large multimers of von Willebrand's factor. Absent ADAMTS13, these large multimers cause platelet clumping and intravascular hemolysis. An antibody to ADAMTS13 is found in many sporadic cases of adult TTP/HUS, but not all; many patients also have antibodies to the thrombospondin receptor on selected endothelial cells in small vessels or increased levels of plasminogen-activator inhibitor 1 (PAI-1). Some children with complement protein deficiencies express atypical HUS (aHUS), which can be treated with liver transplant. The treatment of adult TTP/HUS is daily plasmapheresis, which can be lifesaving. Plasmapheresis is given until the platelet count rises, but in relapsing patients it normally is continued well after the platelet count improves, and in resistant patients twice-daily exchange may be helpful. Most patients respond within 2 weeks of daily plasmapheresis. Since TTP/HUS often has an autoimmune basis, there is an anecdotal role in relapsing patients for using splenectomy, steroids, immunosuppressive drugs, bortezomib, or rituximab, an anti-CD20 antibody. Patients with childhood HUS from infectious diarrhea are not given antibiotics, because antibiotics are thought to accelerate the release of the toxin and the diarrhea is usually self-limited. No intervention appears superior to supportive therapy in children with postdiarrheal HUS.

ANTIPHOSPHOLIPID ANTIBODY SYNDROME (SEE CHAP. 379)

GLOBAL CONSIDERATIONS

INFECTIOUS DISEASE-ASSOCIATED SYNDROMES

 A number of infectious diseases will injure the glomerular capillaries as part of a systemic reaction producing an immune response or from direct infection of renal tissue. Evidence of this immune response is collected by glomeruli in the form of immune deposits that damage the kidney, producing moderate proteinuria and

hematuria. A high prevalence of many of these infectious diseases in undeveloped countries results in infection-associated renal disease being the most common cause of glomerulonephritis in many parts of the world.

Poststreptococcal Glomerulonephritis This form of glomerulonephritis is one of the classic complications of streptococcal infection. The discussion of this disease can be found earlier, in the section “Acute Nephritic Syndromes.”

Subacute Bacterial Endocarditis Renal injury from persistent bacteremia absent the continued presence of a foreign body, regardless of cause, is treated presumptively as if the patient has endocarditis. The discussion of this disease can be found earlier, in the section “Acute Nephritic Syndromes.”

Human Immunodeficiency Virus Renal disease is an important complication of HIV disease. The risk of development of end-stage renal disease is much higher in HIV-infected African Americans than in HIV-infected whites. About 50% of HIV-infected patients with kidney disease have HIV-associated nephropathy (HIVAN) on biopsy. The lesion in HIVAN is FSGS, characteristically revealing a collapsing glomerulopathy (see Fig. 62e-3) with visceral epithelial cell swelling, microcystic dilatation of renal tubules, and tubuloreticular inclusion. Renal epithelial cells express replicating HIV virus, but host immune responses also play a role in the pathogenesis. MPGN and DPGN have also been reported but more commonly in HIV-infected whites and in patients coinfecting with hepatitis B or C. HIV-associated TTP has also been reported. Other renal lesions include DPGN, IgA nephropathy, and MCD. Renal biopsy may be indicated to distinguish between these lesions.

HIV patients with FSGS typically present with nephrotic-range proteinuria and hypoalbuminemia, but unlike patients with other etiologies for nephrotic syndrome, they do not commonly have hypertension, edema, or hyperlipidemia. Renal ultrasound also reveals large, echogenic kidneys despite the finding that renal function in some patients declines rapidly. Treatment with inhibitors of the renin-angiotensin system decreases the proteinuria. Effective antiretroviral therapy benefits both the patient and the kidney and improves survival of HIV-infected patients with chronic kidney disease (CKD) or end-stage renal disease. In HIV-infected patients not yet on therapy, the presence of HIVAN is an indication to initiate therapy. Following the introduction of antiretroviral therapy, survival on dialysis for the HIV-infected patient has improved dramatically. Renal transplantations in HIV-infected patients without detectable viral loads or histories of opportunistic infections provide a better survival benefit over dialysis. Following transplantation, patient and graft survival are similar to the general transplant population despite frequent rejections.

Hepatitis B and C Typically infected patients present with microscopic hematuria, nonnephrotic or nephrotic-range proteinuria, and hypertension. There is a close association between hepatitis B infection and polyarteritis nodosa with vasculitis appearing generally in the first 6 months following infection. Renal manifestations include renal artery aneurysms, renal infarction, and ischemic scars. Alternatively, the hepatitis B carrier state can produce a MGN that is more common in children than adults, or MPGN that is more common in adults than in children. Renal histology is indistinguishable from idiopathic MGN or type I MPGN. Viral antigens are found in the renal deposits. There are no good treatment guidelines, but interferon α -2b and lamivudine have been used to some effect in small studies. Children have a good prognosis, with 60–65% achieving spontaneous remission within 4 years. In contrast, 30% of adults have renal insufficiency and 10% have renal failure 5 years after diagnosis.

Up to 30% of patients with chronic hepatitis C infection have some renal manifestations. Patients often present with type II mixed cryoglobulinemia, nephrotic syndrome, microscopic hematuria, abnormal liver function tests, depressed C3 levels, anti-hepatitis C virus (HCV) antibodies, and viral RNA in the blood. The renal lesions most commonly seen, in order of decreasing frequency, are *cryoglobulinemic glomerulonephritis*, MGN, and *type I MPGN*. Treatment with pegylated interferon and ribavirin is typical to reduce the viral load.

Other Viruses Other viral infections are occasionally associated with glomerular lesions, but cause and effect are not well established. These viral infections and their respective glomerular lesions include: cytomegalovirus producing MPGN; influenza and anti-GBM disease; measles-associated endocapillary proliferative glomerulonephritis, with measles antigen in the capillary loops and mesangium; parvovirus causing mild proliferative or mesangioproliferative glomerulonephritis or FSGS; mumps and mesangioproliferative glomerulonephritis; Epstein-Barr virus producing MPGN, diffuse proliferative nephritis, or IgA nephropathy; dengue hemorrhagic fever causing endocapillary proliferative glomerulonephritis; and coxsackievirus producing *focal glomerulonephritis* or DPGN.

Syphilis Secondary syphilis, with rash and constitutional symptoms, develops weeks to months after the chancre first appears and occasionally presents with the nephrotic syndrome from MGN caused by subepithelial immune deposits containing treponemal antigens. Other lesions have also rarely been described including interstitial syphilitic nephritis. The diagnosis is confirmed with nontreponemal and treponemal tests for *Treponema pallidum*. The renal lesion responds to treatment with penicillin or an alternative drug, if allergic. Additional testing for other sexually transmitted diseases is an important part of disease management.

Leprosy Despite aggressive eradication programs, approximately 400,000 new cases of leprosy appear annually worldwide. The diagnosis is best made in patients with multiple skin lesions accompanied by sensory loss in affected areas, using skin smears showing paucibacillary or multibacillary infection (WHO criteria). Leprosy is caused by infection with *Mycobacterium leprae* and can be classified by Ridley-Jopling criteria into various types: tuberculoid, borderline tuberculoid, mid-borderline and borderline lepromatous, and lepromatous. Renal involvement in leprosy is related to the quantity of bacilli in the body, and the kidney is one of the target organs during splanchnic localization. In some series, all cases with borderline lepromatous and lepromatous types of leprosy have various forms of renal involvement including FSGS, mesangioproliferative glomerulonephritis, or renal amyloidosis; much less common are the renal lesions of DPGN and MPGN. Treatment of the infection can cause remission of the renal disease.

Malaria There are 300–500 million incident cases of malaria each year worldwide, and the kidney is commonly involved. Glomerulonephritis is due to immune complexes containing malarial antigens that are implanted in the glomerulus. In malaria from *P. falciparum*, mild proteinuria is associated with subendothelial deposits, mesangial deposits, and mesangioproliferative glomerulonephritis that usually resolve with treatment. In quartan malaria from infection with *Plasmodium malariae*, children are more commonly affected and renal involvement is more severe. Transient proteinuria and microscopic hematuria can resolve with treatment of the infection. However, resistant nephrotic syndrome with progression to renal failure over 3–5 years does happen, as <50% of patients respond to steroid therapy. Affected patients with nephrotic syndrome have thickening of the glomerular capillary walls, with subendothelial deposits of IgG, IgM, and C₃ associated with a sparse membranoproliferative lesion. The rare mesangioproliferative glomerulonephritis reported with *Plasmodium vivax* or *Plasmodium ovale* typically has a benign course.

Schistosomiasis Schistosomiasis affects more than 300 million people worldwide and primarily involves the urinary and gastrointestinal tracts. Glomerular involvement varies with the specific strain of schistosomiasis; *Schistosoma mansoni* is most commonly associated with clinical renal disease, and the glomerular lesions can be classified: Class I is a *mesangioproliferative glomerulonephritis*; class II is an *extracapillary proliferative glomerulonephritis*; class III is a *membranoproliferative glomerulonephritis*; class IV is a *focal segmental glomerulonephritis*; and class V is *amyloidosis*. Classes I–II often remit with treatment of the infection, but classes III and IV lesions are associated with IgA immune deposits and progress despite antiparasitic and/or immunosuppressive therapy.

Other Parasites Renal involvement with toxoplasmosis infections is rare. When it occurs, patients present with nephrotic syndrome

and have a histologic picture of MPGN. Fifty percent of patients with leishmaniasis will have mild to moderate proteinuria and microscopic hematuria, but renal insufficiency is rare. Acute DPGN, MGN, and mesangioproliferative glomerulonephritis have all been observed on biopsy. Filariasis and trichinosis are caused by nematodes and are sometimes associated with glomerular injury presenting with proteinuria, hematuria, and a variety of histologic lesions that typically resolve with eradication of the infection.

339 Polycystic Kidney Disease and Other Inherited Disorders of Tubule Growth and Development

Jing Zhou, Martin R. Pollak

The polycystic kidney diseases are a group of genetically heterogeneous disorders and a leading cause of kidney failure. The autosomal dominant form of polycystic kidney disease (ADPKD) is the most common life-threatening monogenic disease, affecting 12 million people worldwide. The autosomal recessive form of polycystic kidney disease (ARPKD) is rarer but affects the pediatric population. Kidney cysts are often seen in a wide range of syndromic diseases. Recent studies have shown that defects in the structure or function of the primary cilia may underlie this group of genetic diseases collectively termed *ciliopathies* (Table 339-1).

AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE

Etiology and Pathogenesis (Fig. 339-1) ADPKD is characterized by progressive formation of epithelial-lined cysts in the kidney. Although cysts only occur in 5% of the tubules in the kidney, the enormous growth of these cysts ultimately leads to the loss of normal surrounding tissues and loss of renal function. The cellular defects in ADPKD that have been known for a long time are increased cell proliferation and fluid secretion, decreased cell differentiation, and abnormal extracellular matrix. ADPKD is caused by mutations in *PKD1* and *PKD2*, which, respectively, code for polycystin-1 (PC1) and polycystin-2 (PC2). PC1 is a large 11-transmembrane protein that functions like a G protein-coupled receptor. PC2 is a calcium-permeable six-transmembrane protein that structurally belongs to the transient receptor potential (TRP) cation channel family. PC1 and PC2 are widely expressed in almost all tissues and organs. PC1 expression is high in development and low in the adult, whereas PC2 expression is relatively constant. PC1 and PC2 are found on the primary cilium, a hair-like structure present on the apical membrane of a cell, in addition to the cell membranes and cell-cell junctions of tubular epithelial cells. Defects in the primary cilia are linked to a wide spectrum of human diseases, collectively termed *ciliopathies*. The most common phenotype shared by many *ciliopathies* is kidney cysts. PC1 and PC2 bind to each other via their respective C-terminal tails to form a receptor-channel complex and regulate each other's function. The PC1/2 protein complex serves as a mechanosensor or chemical sensor and regulates calcium and G-protein signaling. The PC1/2 protein complex may also directly regulate a number of cellular functions including the cell cycle, the actin cytoskeleton, planar cell polarity (PCP), and cell migration. This protein complex has also been implicated in regulating a number of signaling pathways, including Wnt, mammalian target of rapamycin (mTOR), STAT3, cMET, phosphoinositide 3-kinase (PI3K)/AKT, G protein-coupled receptor (GPCR), and epidermal growth factor receptor (EGFR), as well as in the localization and activity of cystic fibrosis transmembrane conductance (CFTR). One hypothesis is

TABLE 339-1 INHERITED DISEASES COMMONLY ASSOCIATED WITH A CYSTIC PHENOTYPE

Disease	Mode of Inheritance	Renal Abnormalities	Other Clinical Features	Genes
Autosomal dominant polycystic kidney disease	AD	Cortical and medullary cysts	Liver, pancreatic cysts, hypertension, subarachnoid hemorrhage	<i>PKD1, PKD2</i>
Autosomal recessive polycystic kidney disease	AR	Distal and collecting duct cysts	Oligohydramnios if severe, hypertension, ascending cholangitis, liver fibrosis	<i>PKHD1</i>
Medullary cystic kidney	AD	Small fibrotic kidneys; medullary cysts	In adults, gout	<i>MCKD1, MCKD2/UMOD</i>
Nephronophthisis	AR	Small fibrotic kidneys; medullary cysts	Growth retardation, anemia (visual loss, liver fibrosis, cerebellar ataxia if associated with another syndrome)	<i>NPHP1-4, IQCB1, CEP290, GLIS2, RPGRIP1L, NEK8, SDCCAG8, TMEM67, TTC21B</i>
Senior-Løken syndrome	AR	Renal cysts	Juvenile nephronophthisis, Leber's amaurosis	<i>NPHP1-6, SDCCAG8</i>
Leber's congenital amaurosis	AR	Renal cysts	Visual impairment in first year of life, pigmentary retinopathy	<i>GUCY2D, RPE65, LCA3-14</i> (including <i>LCA10, CEP290</i>)
Meckel-Gruber syndrome	AR	Cortical and medullary cysts	CNS anomalies, polydactyly, congenital heart defects	<i>MKS1, TMEM216, TMEM67, CEP290, RPGRIP1L, CC2D2A, TCTN2, B9D1, B9D2, NPHP3</i>
Bardet-Biedl syndrome	AR	Renal cysts	Obesity, polydactyly, retinitis pigmentosa, anosmia, congenital heart defects, mental retardation	<i>BBS1, 2, ARL6, BBS4,5, MKKS, BBS7, TTC8, BBS9, 10, TRIM32, BBS12, MKS1, CEP290, C2ORF86</i> ; modifiers <i>MKS1, MKS3, CCDC28B</i>
Oral-facial-digital syndrome type I	AR	Renal cysts	Oral cavity, face, and digit anomalies; CNS abnormalities; cystic kidney disease; X-linked with male lethality, primary ciliary dyskinesia	<i>OFD1</i>
Cranioectodermal dysplasia (Sensenbrenner's syndrome)	AR	Renal cysts	Skeletal dysplasia, thoracic deformities, polydactyly, renal cysts, retinitis pigmentosa	<i>IFT80</i>
Tuberous sclerosis	AD	Renal cysts	Angiomyolipomas, renal cell carcinoma, facial angiofibromas, CNS hamartomas	<i>TSC1, TSC2</i>
Von Hippel-Lindau disease	AD	Renal cysts	Renal cell carcinoma, retinal angiomas, CNS hemangioblastomas, pheochromocytomas	<i>VHL</i>

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; CNS, central nervous system.

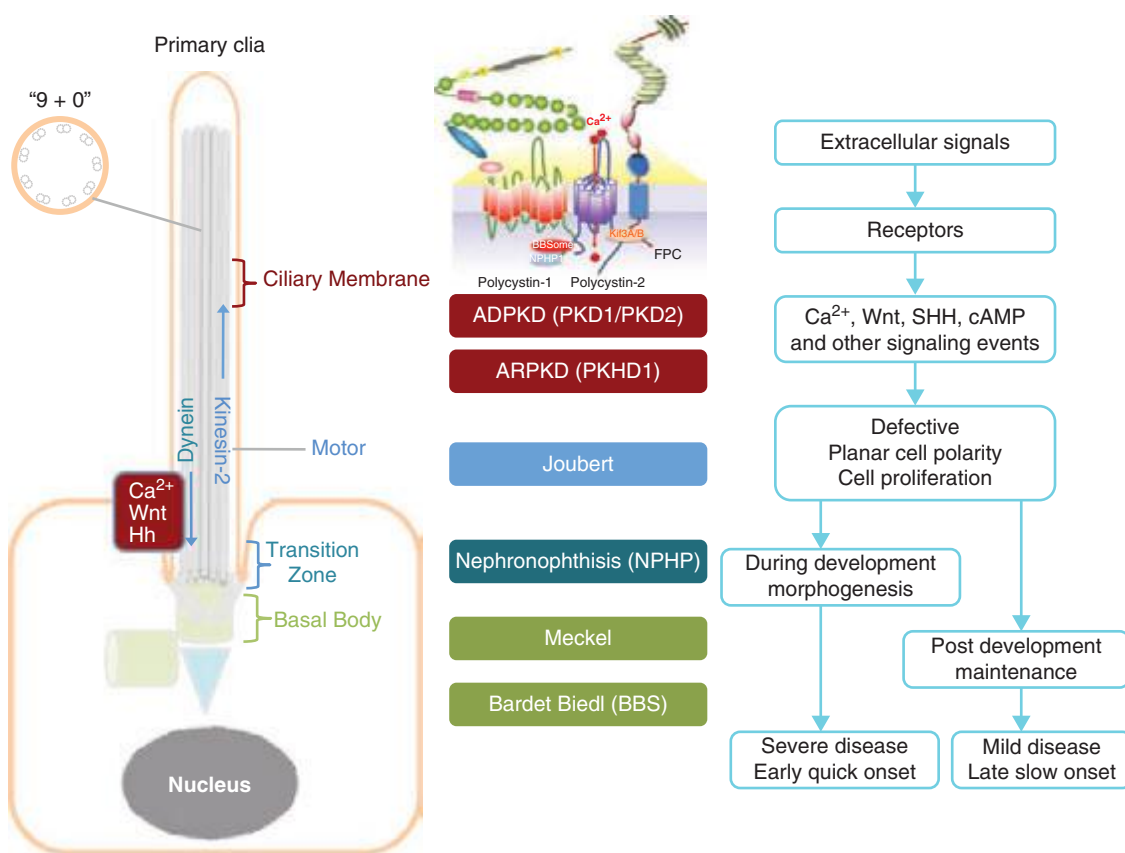


FIGURE 339-1 Scheme of the primary cilium and cystic kidney disease proteins. **Left.** A scheme of the primary cilium. Primary cilia share a “9+0” organization of microtubule doublets. Proteins are transported into the cilium by motor protein kinesin 2 and transported out of the cilium by dynein. The cilium is connected to the basal body through the transition zone. **Middle.** Topology of autosomal dominant polycystic kidney disease (ADPKD) and autosomal recessive polycystic kidney disease (ARPKD) proteins polycystin-1, polycystin-2, and fibrocystin/polyductin (FPC) are shown. PC1 also interacts with other proteins such as components of the BBSome and NPHP1. PC2 and FPC both interact with kinesin 2 (KIF 3A/B). Localization of disease proteins in the cilium, the transition zone, and the basal body is color coded. **Right.** Potential disease mechanisms due to cilium-mediated signaling events.

that loss of ciliary function of PC1 and PC2 leads to reduced calcium signaling and a subsequent increase of adenylyl cyclase activity and decrease of phosphodiesterase activity, which, in turn, causes increased cellular cyclic AMP (cAMP). Increased cAMP promotes protein kinase A activity, among other effectors, and, in turn, leads to cyst growth by promoting proliferation and fluid secretion of cyst-lining cells through chloride and aquaporin channels in ADPKD kidneys.

Genetic Considerations ADPKD is inherited as an autosomal dominant trait with complete penetrance but variable expressivity. The disease affects all ethnic groups worldwide with an estimated prevalence of 1:1000 to 1:400. Only half of the patients with ADPKD are clinically diagnosed during their lifetime. ADPKD is genetically heterogeneous. The first disease gene (*PKD1*) was localized to the region of the α -globin gene on chromosome 16p13 in 1985, and a second disease gene (*PKD2*) locus was mapped to chromosome 4q21-q23 in 1993. Mutations of *PKD1* and *PKD2* are responsible for ~85% and ~15% of ADPKD cases, respectively. However, patients with *PKD2* mutations may be higher than 15% because they tend to have milder clinical disease and, as a result, may be underdiagnosed. Embryonic lethality of *Pkd1* and *Pkd2* knockout mice suggests that human homozygotes may be lethal and thus not clinically recognized.

PKD1 is comprised of 46 exons occupying ~52 kb of genomic DNA. It produces an ~14-kb transcript that encodes PC1, a protein of ~4300 amino acids. A feature of the *PKD1* gene is that the 5' three-quarters of *PKD1* have been duplicated at six other sites on chromosome 16p, and many of them produce mRNA transcripts, which provides a major challenge for genetic analysis of the duplicated region. *PKD2* is a single-copy gene with 15 exons producing an ~5.3-kb mRNA transcript that encodes PC2, a protein of 968 amino acids. The presence of additional genes for ADPKD was suggested based on several families

linked to neither *PKD1* nor *PKD2* genes. However, careful analyses have excluded the existence of a third ADPKD gene.

In ADPKD patients, every cell carries a germline mutant allele of either *PKD1* or *PKD2*. However, cysts develop in only a small fraction of the nephrons. Cysts are thought to originate from clonal growth of single cells that have received a somatic “second hit” mutation in the “normal” allele of the *PKD1* or *PKD2* gene. Accumulating evidence in mouse models now shows that partial loss of function of the second allele of *Pkd1* in a proliferative environment is sufficient for cystogenesis, suggesting that a critical amount of *PKD1* is needed in a cell. Somatic inactivation of the second allele of *Pkd1* in adult mice results in very slow onset of cyst development in the kidney, but a “third hit,” such as an additional genetic or epigenetic event, the inactivation of a growth-suppressor gene, the activation of a growth-promoting gene(s), or an event like renal injury that activates the developmental program, may promote rapid cyst formation.

Clinical Manifestations ADPKD is characterized by the progressive bilateral formation of renal cysts. Focal renal cysts are typically detected in affected subjects before 30 years of age. Hundreds to thousands of cysts are usually present in the kidneys of most patients in the fifth decade (Fig. 339-2). Enlarged kidneys can each reach a fourfold increase in length and weigh up to 20 times the normal weight. The clinical presentations of ADPKD are highly variable. Although many patients are asymptomatic until the fourth to fifth decade of life and are diagnosed by incidental discoveries of hypertension or abdominal masses, back or flank pain is a frequent symptom in ~60% of patients with ADPKD. The pain may result from renal cyst infection, hemorrhage, or nephrolithiasis. Gross hematuria resulting from cyst rupture occurs in ~40% of patients during the course of their disease, and many of them will have recurrent episodes. Flank pain and hematuria



FIGURE 339-2 Photograph showing a kidney from a patient with autosomal dominant polycystic kidney disease. The kidney has been cut open to expose the parenchyma and internal aspects of cysts.

may coexist if the cyst that ruptures is connected with the collecting system. Proteinuria is usually a minor feature of ADPKD. Infection is the second most common cause of death for patients with ADPKD. Up to half of patients with ADPKD will have one or more episodes of renal infection during their lifetime. An infected cyst and acute pyelonephritis are the most common renal infections often due to gram-negative bacteria, which are associated with fever and flank pain, with or without bacteremia. These complications and renal insufficiency often correlate with structural abnormality of the renal parenchyma. Kidney stones occur in ~20% of patients with ADPKD. Different from the general population, more than half of the stones in patients with ADPKD are composed of uric acid, with the remainder due to calcium oxalate. Distal acidification defects, abnormal ammonium transport, low urine pH, and hypocitraturia may be important in the pathogenesis of renal stones in ADPKD. Renal cell carcinoma is a rare complication of ADPKD with no apparent increased frequency compared to the general population. However, in ADPKD, these tumors are more often bilateral at presentation, multicentric, and sarcomatoid in type. Radiologic imaging is often not helpful in distinguishing cyst infection and cyst hemorrhage because of their complexity. Computed tomography (CT) scan and magnetic resonance imaging (MRI) are often useful in distinguishing a malignancy from a complex cyst. Cardiovascular complications are the major cause of mortality in patients with ADPKD. Hypertension is common and typically occurs before any reduction in glomerular filtration rate (GFR). Hypertension is a risk factor for both cardiovascular and kidney disease progression in ADPKD. Notably, some normotensive patients with ADPKD may also have left ventricular hypertrophy. Hypertension in ADPKD may result from the increased activation of the renin-angiotensin-aldosterone

system, increased sympathetic nerve activity, and impaired endothelial cilium function-dependent relaxation of small resistant blood vessels.

The progression of ADPKD has striking inter- and intrafamilial variability. The disease can present as early as in utero, but end-stage renal disease typically occurs in late middle age. Risk factors include early diagnosis of ADPKD, hypertension, gross hematuria, multiple pregnancies, and large kidney size. Liver cysts derived from the biliary epithelia are the most common extrarenal complication. Polycystic liver disease associated with ADPKD is different from autosomal dominant polycystic liver disease (ADPLD), which is caused by mutations in at least two distinct genes (*PRKCSH* and *SEC63*) and does not progress to renal failure. Massive polycystic liver disease occurs almost exclusively in women with ADPKD, particularly those with multiple pregnancies.

Intracranial aneurysm (ICA) occurs four to five times more frequently in ADPKD patients than in the general population and causes high mortality. The disease gene products PC1 and PC2 may be directly responsible for defects in arterial smooth muscle cells and myofibroblasts. The focal nature and the natural history of ICA in ADPKD remain unclear. A family history of ICA is a risk factor of aneurysm rupture in ADPKD, but whether hypertension and cigarette smoking are independent risk factors is not clear. About 20–50% of patients may experience “warning headaches” preceding the index episode of subarachnoid hemorrhage due to ruptured ICA. A CT scan is generally used as the first diagnostic test. A lumbar puncture may be used to confirm the diagnosis. The role of radiologic screening for ICA in asymptomatic patients with ADPKD remains unclear. ADPKD patients with a positive family history of ICAs may undergo presymptomatic screening of ICAs by magnetic resonance angiography. Other vascular abnormalities in ADPKD patients include diffuse arterial dolichoectasias of the anterior and posterior cerebral circulation, which can predispose to arterial dissection and stroke. Mitral valve prolapse occurs in up to 30% of patients with ADPKD, and tricuspid valve prolapse is less common. Other valvular abnormalities occurring with increased frequency in ADPKD patients include insufficiency of the mitral, aortic, and tricuspid valves. Most patients are asymptomatic, but some may progress and require valve replacement. The prevalence of colonic diverticulae and abdominal wall hernias is also increased in ADPKD patients.

Diagnosis Diagnosis is typically made from a positive family history consistent with autosomal dominant inheritance and multiple kidney cysts bilaterally. Renal ultrasonography is often used for presymptomatic screening of at-risk subjects and for evaluation of potential living-related kidney donors from ADPKD families. The presence of *at least two renal cysts (unilateral or bilateral)* is sufficient for diagnosis among at-risk subjects between 15 and 29 years of age with a sensitivity of 96% and specificity of 100%. The presence of *at least two cysts in each kidney* and the presence *at least four cysts in each kidney* are required for the diagnosis of at-risk subjects age 30 to 59 years and age 60 years or older, respectively, with a sensitivity of 100% and specificity of 100%. This is because there is an increased frequency of developing simple renal cysts with age. Conversely, in subjects between age 30 and 59 years, the absence of *at least two cysts in each kidney*, which is associated with a false-negative rate of 0%, can be used for disease exclusion. These criteria have a lower sensitivity for patients with a *PKD2* mutation because of a late onset of ADPKD2. CT scan and T2-weighted MRI, with and without contrast enhancement, are more sensitive than ultrasonography and can detect cysts of smaller size. However, a CT scan exposes the patient to radiation and radiocontrast, which may cause serious allergic reactions and nephrotoxicity in patients with renal insufficiency. T2-weighted MRI, with gadolinium as a contrast agent, has minimal renal toxicity and can detect cysts of only 2–3 mm in diameter. However, a large majority of cysts may still be below the detection level. Genetic testing by linkage analyses and mutational analyses is available for ambiguous cases. Because of the large size of the *PKD1* gene and the presence of multiple highly homologous pseudogenes, mutational analysis of the *PKD1* gene is difficult and costly. Application of new technologies, such as paired-end

next-generation sequencing with multiplexing individually bar-coded long-range polymerase chain reaction libraries, may reduce the costs and improve the sensitivity for clinical genetic testing.

TREATMENT AUTOSOMAL DOMINANT POLYCYSTIC KIDNEY DISEASE

No specific treatment to prevent cyst growth or the decline of renal function has been approved by U.S. Food and Drug Administration. Blood pressure control to a target of 140/90 mmHg is recommended according to the guidelines from the eighth report of the Joint National Committee on Prevention, Detection, Evaluation, and Treatment of High Blood Pressure (JNC VIII report) for reducing cardiovascular complications in ADPKD and renal disease progression. More rigorous blood pressure control does not equal greater clinical benefits. Maintaining a target systolic blood pressure to 110 mmHg in patients with moderate or advanced disease may increase the risk of renal disease progression by reducing renal blood flow. Lipid-soluble antibiotics against common gram-negative enteric organisms, such as trimethoprim-sulfamethoxazole, quinolones, and chloramphenicol, are preferred for cyst infection because most renal cysts are not connected to glomerular filtration and antibiotics that are capable of penetrating the cyst walls are likely to be more effective. Treatment often requires 4–6 weeks. The treatment of kidney stones in ADPKD includes standard measures such as analgesics for pain relief and hydration to ensure adequate urine flow. Management of chronic flank, back, or abdominal pain due to renal enlargement may include both pharmacologic (non-narcotic and narcotic analgesics) and nonpharmacologic measures (transcutaneous electrical nerve stimulation, acupuncture, and biofeedback). Occasionally, surgical decompression of cysts may be necessary. More than half of ADPKD patients eventually require peritoneal dialysis, hemodialysis, or kidney transplantation. Peritoneal dialysis may not be suitable for some patients with massively enlarged polycystic kidneys due to the small intraabdominal space for efficient peritoneal exchange of fluid and solutes and increased chance of abdominal hernia and back pain. Patients with very large polycystic kidneys and recurrent renal cyst infection may require pretransplant nephrectomy or bilateral nephrectomy to accommodate the allograft and reduce the pain.

Specific treatment strategies for ADPKD have focused on slowing renal disease progression and lowering cardiovascular risk. For the latter, the main approach is to control blood pressure by inhibiting the renin-angiotensin-aldosterone system. The ongoing HALT PKD trial was set to evaluate the impact of intensive blockade of the renin-angiotensin-aldosterone system and levels of blood pressure control on progressive renal disease. Most approaches target the slowing of renal disease progression by inhibiting cell proliferation and fluid secretion. Several clinical trials have been conducted targeting cell proliferation, including studies on sirolimus and everolimus, inhibitors of the mTOR pathway; OPC31260 and tolvaptan, which inhibit cAMP pathways by antagonizing the activation of vasopressin V2 receptor (V2R) in collecting ducts and reduce cell proliferation by decreasing renal cAMP levels; and somatostatin analogues, which reduce cAMP levels by binding to several GPCRs. Both the V2R antagonists and somatostatin analogues appear to slow the decline of renal function, although with some side effects such as liver function impairment, polydipsia, and diarrhea. A combination of different growth inhibitors may enhance efficacy and reduce side effects. Additional preclinical studies in animal models include the use of inhibitors to nonreceptor tyrosine kinase Src, B-raf, cyclin-dependent kinase (CDK), transcription factors STAT3 and STAT6 (pyrimethamine and leflunomide), purinergic receptors, hepatocyte growth factor receptor, and glucosylceramide, and agonists to peroxisome proliferator activated receptor- γ (PPAR γ receptors (thiazolidinediones).

AUTOSOMAL RECESSIVE POLYCYSTIC KIDNEY DISEASE

Genetic Considerations ARPKD is a significant hereditary renal disease in childhood, with an estimated prevalence of 1 in 20,000 live births. A carrier frequency of up to 1:70 has been reported. Mutations in a single gene, *PKHD1*, are responsible for all the clinical presentations of ARPKD. *PKHD1*, localized on human chromosome region 6p21.1-6p12.2, is one of the largest genes in the genome, occupies ~450 kb of DNA, and contains at least 86 exons. It produces multiple alternatively spliced transcripts. The largest transcript encodes fibrocystin/polyductin (FPC), which is a large receptor-like integral membrane protein of 4074 amino acids. FPC has a single transmembrane, a large N-terminal extracellular region, and a short intracellular cytoplasmic domain. FPC is localized on the primary cilia of epithelial cells of cortical and medullary collecting ducts and cholangiocytes of bile ducts, similar to polycystins and several other ciliopathy proteins. FPC is also expressed on the basal body and plasma membrane. The large extracellular domain of FPC is presumed to bind to an as yet unknown ligand(s) and is involved in cell-cell and cell-matrix interactions. FPC interacts with ADPKD protein PC2 and may also participate in regulation of the mechanosensory function of the primary cilia, calcium signaling, and PCP, suggesting a common mechanism underlying cystogenesis between ADPKD and ARPKD. FPC is also found on the centrosomes and mitotic spindle and may regulate centrosome duplication and mitotic spindle assembly during cell division. A large number of various mutations have been found throughout *PKHD1* and are unique to individual families. Most patients are compound heterozygotes for *PKHD1* mutations. Patients with two truncation mutations appear to have an earlier onset of the disease.

Clinical Features Classic ARPKD is generally diagnosed in utero or within the neonatal period and characterized by greatly enlarged echogenic kidneys in diseased fetuses. Reduced fetal urine production may contribute to oligohydramnios and pulmonary hypoplasia. About 30% of affected neonates die shortly after birth due to respiratory insufficiency. Close to 60% of mortality occurs within the first month of life. In the classic group, most patients are born with renal insufficiency and ESRD. However, infants often have a transient improvement in their GFR; death from renal insufficiency at this stage is rare. Some patients are diagnosed after the neonatal stage and form the older group. Morbidity and mortality in this group often involve systemic hypertension, progressive renal insufficiency, and liver manifestations. The hallmarks of ARPKD liver disease are biliary dysgenesis due to a primary ductal plate malformation with associated periportal fibrosis, namely congenital hepatic fibrosis (CHF) and dilatation of intrahepatic bile ducts (Caroli's disease). CHF and Caroli's disease can then lead to portal hypertension exhibiting hepatosplenomegaly, variceal bleeding, and cholangitis. Some patients with the diagnosis of ARPKD at 1 year of age with nephromegaly exhibit slowly declining renal function over 20 years with only minimally enlarged kidneys at ESRD and markedly atrophic kidneys following renal transplantation. The slow progression of renal disease is likely due to increasing fibrosis rather than the development of cysts. Systemic hypertension is common in all ARPKD patients, even those with normal renal function.

Diagnosis Ultrasonography, CT, and MRI all can be used for diagnosis. Ultrasonography reveals large, echogenic kidneys with poor corticomedullary differentiation. The diagnosis can be made in utero after 24 weeks of gestation in severe cases. Macrocysts generally are not common at birth in ARPKD patients. The absence of renal cysts in either parent, particularly if they are more than 40 years of age on ultrasonography, helps distinguish ARPKD from ADPKD in older patients. Clinical, laboratory, or radiographic evidence of hepatic fibrosis, hepatic pathology demonstrating characteristic ductal plate abnormalities, family history of affected siblings, or parental consanguinity suggestive of autosomal recessive inheritance is helpful. The lack of mutational hotspots and the large and complex genomic structure of *PKHD1* make molecular diagnosis difficult; however, presymptomatic screening of other at-risk members in a family with already identified ARPKD mutations is straightforward and inexpensive.

TREATMENT AUTOSOMAL RECESSIVE POLYCYSTIC KIDNEY DISEASE

There is no specific therapy for ARPKD. Appropriate neonatal intensive care, blood pressure control, dialysis, and kidney transplantation increase survival into adulthood. Complications of hepatic fibrosis may necessitate liver transplantation. Patients with severe Caroli's disease may need portosystemic shunting. Upcoming therapies may target abnormal cell signaling mechanisms, as described above for ADPKD.

OTHER DISEASES CHARACTERIZED BY LARGE KIDNEY CYSTS

TUBEROUS SCLEROSIS

Tuberous sclerosis (TS) is a rare autosomal dominant syndrome caused by mutations in one of two genes, *TSC1*, encoding hamartin, or, *TSC2*, encoding tuberin. Published estimates of prevalence vary widely, but it certainly occurs in less than 1:5000 births. Kidney cysts are a frequent feature of this condition, as are two other abnormalities of kidney growth, renal cell carcinoma and renal angiomyolipomas. TS is a syndrome affecting multiple organ systems. Other features of TS include benign growths in the nervous system, eyes, heart, lung, liver, and skin. Essentially all TS patients have associated skin lesions, and a large proportion of patients have neurologic and cognitive manifestations. The *TSC2* gene is adjacent to *PKD1* in the human genome. Some patients have deletions in their genomic DNA that inactivate these two genes. Such individuals may have manifestations of both ADPKD and TS.

The most common kidney finding in TS is the presence of angiomyolipomas. These growths tend to be multiple and bilateral. Although they are usually benign, they may bleed. Surgical removal is often recommended as a prophylactic measure in people with angiomyolipomas larger than 4 cm in diameter. The cysts in TS are radiographically similar to those seen in ADPKD. In contrast to ADPKD, there is a clearly increased risk of renal cell carcinoma in TS patients. Regular periodic imaging is recommended in TS patients with kidney involvement to screen for the development of renal cell carcinoma.

Although not common, TS may lead to significant chronic kidney disease (CKD) and progress to end-stage kidney failure. Patients with TS and CKD typically have an unremarkable urine sediment and only minimal to mild amounts of proteinuria.

Mechanistically, the *TSC1* and *TSC2* gene products tuberin and hamartin interact physically. This protein complex is localized to the base of the cilia and inhibits intracellular signaling processes mediated by mTOR, leading to abnormal growth in a number of tissues. Investigation of mTOR inhibitors as therapy for TS is ongoing.

VON HIPPEL-LINDAU DISEASE

Von Hippel-Lindau disease (VHL) is an inherited cancer syndrome with renal manifestations. VHL is an autosomal dominant condition caused by mutations in the VHL tumor-suppressor gene. VHL is localized to the primary cilia and is necessary for the formation of primary cilia. Like many autosomal dominant cancer syndromes, VHL is recessive at the cellular level: a somatic mutation in the second VHL allele leads to loss of VHL in the cell and abnormal growth. Kidney manifestations of VHL include multiple bilateral kidney cysts and renal cell carcinomas. Kidney cysts and carcinoma affect the majority of VHL patients. Nonrenal features of VHL include pheochromocytomas, cerebellar hemangioblastomas, and retinal hemangiomas.

Annual screening of the kidneys by imaging with CT or MRI is recommended for early detection of renal cell carcinomas. Increasingly, nephron-sparing surgical approaches are being used for removal of cancerous lesions in order to preserve kidney function.

OTHER INHERITED DISEASES OF TUBULE GROWTH AND DEVELOPMENT

ADPKD is by far the most common adult-onset, single-gene form of kidney disease. The large cysts that are sometimes seen in VHL and

TS are similar in appearance to the cysts seen in ADPKD. A variety of other inherited disorders affecting primarily tubule and renal interstitial function can lead to CKD and eventual end-stage kidney disease in the absence of large tubule-derived cysts.

Inherited diseases affecting the tubulointerstitial compartment of the kidney can lead to secondary glomerular stress and glomerulosclerosis with some degree of concomitant proteinuria. Similarly, disorders of glomerular function will typically lead to secondary interstitial fibrosis and tubule atrophy. From a clinical perspective, therefore, distinguishing between a genetic disease of the renal tubules and a disease of the glomerulus may not be easy, particularly in the absence of a gross phenotype such as large kidney cysts.

MEDULLARY CYSTIC KIDNEY DISEASE (AUTOSOMAL DOMINANT INTERSTITIAL KIDNEY DISEASE)

The medullary cystic kidney diseases (MCKD) are autosomal dominant disorders. Despite the nosology, kidney cysts are not invariably present. Older literature often grouped MCKD together with the childhood-onset disorders known as the nephronophthoses, but these are distinct clinical and genetic entities.

Medullary Cystic Kidney Disease Type I Patients with MCKD type I (MCKD I) have mutations in the mucin 1 gene *MUC1*. In contrast to MCKD type II (MCKD II) patients, individuals with MCKD I do not have elevated uric acid levels. The disease-causing *MUC1* mutations that have been reported all alter a repeat region within the *MUC1* gene, leading to a large "neoprotein" fragment that may lead to toxic effects on the kidney tubule.

Clinically, patients with MCKD I exhibit slowly progressive CKD in adulthood, with only minimal amounts of increased urine protein and occasional renal cysts seen on ultrasound examination. Kidney histology shows tubulointerstitial fibrosis and tubular atrophy. The mechanisms by which *MUC1* mutations cause human kidney disease are not known.

Medullary Cystic Kidney Disease Type II MCKD II is caused by mutations in the *UMOD* gene, which encodes the protein uromodulin, also known as Tamm-Horsfall protein. Uromodulin is also found on the centrosome, the mitotic spindle, and the primary cilia; it colocalizes with nephrocystin-1 and KIF3A on the cilia. *UMOD* mutations also cause the conditions that have been referred to as familial juvenile hyperuricemic nephropathy (HNFJ1) and glomerulocystic kidney disease (GCKD), although it is not clear that these different names represent clearly distinct disorders. The term *uromodulin-associated kidney disease* (UAKD) has been suggested as a better name for MCKD II and the various other related *UMOD*-associated diseases. Despite the name, kidney cysts are not a common feature of MCKD II. MCKD II should be suspected clinically in patients with a family history of late-onset kidney disease, benign urine sediments, absence of significant proteinuria, and hyperuricemia. Large genome-wide association studies have suggested that certain common noncoding sequence variants in *UMOD* are associated with a moderately increased risk of CKD in the general population.

Other Forms of Familial Tubulointerstitial Kidney Disease A small number of families have been identified with autosomal dominant tubulointerstitial kidney disease and hyperuricemia who lack *UMOD* mutations. Some of these families carry disease-segregating mutations in the renin gene *REN*. There are other families who lack mutations in *UMOD*, *MUC1*, or *REN*. Thus, mutations in other yet-to-be identified genes are able to produce similar interstitial kidney disease, both with and without hyperuricemia.

Kidney biopsies in patients with any of the various forms of MCKD typically show interstitial fibrosis. These histologic features are not diagnostic of any particular genetic entity, and the specific diagnosis must be made by other means. Genetic tests for alterations in specific genes are increasingly available in the clinical setting.

Patients with autosomal dominant interstitial kidney disease, *UMOD* or *REN* mutations, or hyperuricemia and gout should be treated similarly to others with these findings, with uric acid-lowering agents, such as allopurinol or febuxostat.

NEPHRONOPHTHISIS

A large and growing number of genetically distinct but related autosomal recessive disorders are referred to as nephronophthyses. These should not be confused with the adult-onset autosomal dominant medullary cystic kidney diseases discussed above, despite the often confusing nomenclature seen in older medical literature. Nephronophthysis is quite rare but is nevertheless the most common inherited childhood form of kidney failure requiring kidney replacement therapy.

Like ADPKD and ARPKD, the various genetically heterogeneous entities that fall under the category of nephronophthysis (NPHP) are disorders of ciliary function. Mutations in a very large number of genes have been identified that lead to NPHP under an autosomal recessive pattern of inheritance. The various forms of NPHP share common features, including tubulointerstitial fibrosis, corticomedullary cysts, and progressive CKD, leading to renal failure. Proteinuria is absent or mild, and the urine sediment is not active.

NPHP is often divided into infantile, juvenile, and adolescent forms. The juvenile form is the most frequent and usually caused by mutations in the *NPHP2* gene. The infantile form, usually caused by *NPHP2* mutations, is associated with end-stage kidney failure in early childhood. Patients with the adolescent form of NPHP typically develop end-stage kidney failure in early adulthood. The products of the NPHP genes are referred to as nephrocystins. *NPHP1* through *NPHP16* have been reported; some are referred to by other names as well.

NPHP can present as an isolated finding or be part of several multiorgan syndromes. Neurologic abnormalities are present in a significant number of patients. Bone and liver abnormalities are seen in some NPHP patients. Senior-Løken syndrome is defined by the presence of NPHP with retinitis pigmentosa. Joubert's syndrome is defined by multiple neurologic findings, including hypoplasia of the cerebellar vermis. Some forms of this genetically heterogeneous syndrome include NPHP as a component.

The multisystem disease Bardet-Biedl syndrome (BBS) is defined clinically by a spectrum of features, including truncal obesity, cognitive impairment, retinal dystrophy, polydactyly, developmental urogenital abnormalities, and kidney cysts. The kidney phenotype is NPHP-like, with small cysts deriving from the tubules, tubulointerstitial and often secondary glomerular disease, and urine concentrating defects. There are 18 BBS genes cloned. BBS follows autosomal recessive inheritance. Like ADPKD, ARPKD, and NPHP, BBS is a disease of abnormal ciliary function.

The multiple genes and gene products (nephrocystins) that are responsible for NPHP are expressed in cilia, basal bodies, and the centrosomes of kidney tubule cells. It has been hypothesized that all of the NPHP gene defects lead to a clinical phenotype by interfering with the regulation of PCP.

There are no specific clinical tests that define NPHP. Genetic diagnosis is possible but cumbersome because of the large number of genes that can be responsible. There are no specific therapies for NPHP. Rather, therapy is aimed at treating signs of these diseases as well as the systemic abnormalities seen with all CKDs. Chronic dialysis or kidney transplantation is eventually required for NPHP-affected individuals.

KARYOMEGALIC TUBULOINTERSTITIAL NEPHRITIS

Karyomegalic tubulointerstitial nephritis is an exceptionally rare form of kidney disease with adult-onset progressive kidney failure. Kidney biopsy shows chronic tubulointerstitial nephritis, as well as interstitial fibrosis. This is a recessive disorder caused by inheritance of two mutant copies of the *FAN1* gene. *FAN1* encodes a component of a DNA repair machinery complex. Individuals with two mutant *FAN1* genes are genetically sensitized to the effect of DNA damage. Kidney histology shows karyomegaly in addition to the nonspecific findings of interstitial fibrosis and tubular atrophy.

MEDULLARY SPONGE KIDNEY

Medullary sponge kidney (MSK) is often grouped together with inherited disorders of the kidney affecting tubule growth and development, although it is usually a sporadic finding rather than an inherited

phenotype. MSK is caused by developmental malformation and cystic dilatation of the renal collecting ducts. The medullary cysts seen in this entity can be quite variable in size.

MSK is usually a benign entity. The diagnosis of MSK is often made incidentally. In the past, the diagnosis of MSK was often made by intravenous pyelography (IVP). CT scans, which have replaced IVPs for much routine kidney imaging, are not as sensitive in detecting MSK.

MSK is associated with an increased frequency of calcium phosphate and calcium oxalate kidney stones. Altered flow characteristics in the kidney tubules may lead to the development of formation of a nidus for stone formation. Kidney stones in this group are treated the same as are kidney stones in the general population. MSK patients also often exhibit reduced kidney concentrating ability and an increased frequency of urinary tract infections.

CONGENITAL ABNORMALITIES OF THE KIDNEY AND URINARY TRACT

The structural abnormalities known as the congenital abnormalities of the kidney and urinary tract (CAKUTs) are a group of etiologically and phenotypically heterogeneous disorders. Some form of CAKUT is estimated to occur in up to 1 in 500 live births. Specific abnormalities classified as part of the CAKUT spectrum include kidney hypoplasia, kidney agenesis, ureteropelvic junction obstruction, and vesicoureteral reflux.

CAKUT can be the cause of clinically significant problems in both adults and children. However, it is a major contributor to kidney failure in children, accounting for more than one-third of end-stage kidney disease in this group.

CAKUT is typically a sporadic finding but can also cluster in families. Familial forms can be observed as parts of multisystem developmental syndromes. A growing list of specific genes have been identified, which when mutated lead to syndromic forms of CAKUT. For example, the branchio-oto-renal syndrome, characterized by developmental abnormalities in the neck, ears, and kidney, can be caused by mutations in the *EYA1* and *SIX1* genes. Mutations in the *PAX2* transcription factor gene can cause the autosomal dominant renal coloboma syndrome, characterized by optic nerve malformations and hypoplastic kidneys.

In many instances, CAKUT is caused by environmental influences rather than genetic alterations. For example, renal tubular dysgenesis, defined by altered tubule development, can be caused by prenatal exposure to angiotensin-converting enzyme inhibitors or angiotensin receptor blockers.

MITOCHONDRIAL DISEASE

Inherited disorders of the mitochondrial genome (discussed elsewhere in this text [Chap. 85e]) commonly affect kidney function. Thirteen of the genes involved in encoding components of the mitochondrial respiratory chain are located on the mitochondrial genome that is inherited maternally. The remainder of these components is encoded by the nuclear genome. These defects of oxidative phosphorylation may affect multiple organs and tissues.

Neuromuscular disease is the best recognized part of this complex phenotype. Kidney disease is now recognized as a common component as well. Tubulointerstitial disease may be seen on kidney biopsy, and progression to kidney failure may occur. Glomerular involvement, manifest as proteinuria and glomerulosclerosis, can also develop. Changes in proximal tubule activity are the most common renal phenotype. Patients may have several defects in proximal tubule transport, including the Fanconi syndrome. Some patients may also have acidosis, hypophosphatemic rickets, hypercalciuria, glycosuria, and tubular proteinuria. Decreased urine concentrating ability is common.

GLOBAL CONSIDERATIONS



The disorders discussed above are all seen worldwide. In addition, a previously unrecognized epidemic of kidney disease is leading to very high rates of kidney failure in and near the

western coast of Central America. This Mesoamerican nephropathy is particularly common in Nicaragua and El Salvador. Mesoamerican nephropathy patients do not have significant proteinuria, suggesting that this is a disease of the kidney tubules and interstitium. The cause is unknown, but some have suggested that a combination of toxic environmental factors and heat stress underlies the development of this kidney disease, which has a striking male predominance. However, the fact that, in many families, a large fraction of the men are affected with kidney disease has suggested that a strong genetic component may be involved as well.

340 Tubulointerstitial Diseases of the Kidney

Laurence H. Beck, David J. Salant

Inflammation or fibrosis of the renal interstitium and atrophy of the tubular compartment are common consequences of diseases that target the glomeruli or vasculature. Distinct from these secondary phenomena, however, are a group of disorders that primarily affect the tubules and interstitium, with relative sparing of the glomeruli and renal vessels. Such disorders are conveniently divided into acute and chronic tubulointerstitial nephritis (TIN) (Table 340-1).

Acute TIN most often presents with acute renal failure (Chap. 334). The acute nature of this group of disorders may be caused by aggressive inflammatory infiltrates that lead to tissue edema, tubular cell injury, and compromised tubular flow, or by frank obstruction of the tubules with casts, cellular debris, or crystals. There is sometimes flank pain due to distention of the renal capsule. Urinary sediment is often active with leukocytes and cellular casts, but depends on the exact nature of the disorder in question.

The clinical features of chronic TIN are more indolent and may manifest with disorders of tubular function, including polyuria from impaired concentrating ability (nephrogenic diabetes insipidus), defective proximal tubular reabsorption leading to features of Fanconi's syndrome (glycosuria, phosphaturia, aminoaciduria, hypokalemia, and type II renal tubular acidosis [RTA] from bicarbonaturia), or non-anion-gap metabolic acidosis and hyperkalemia (type IV RTA) due to impaired ammoniogenesis, as well as progressive azotemia (rising creatinine and blood urea nitrogen [BUN]). There is often modest proteinuria (rarely >2 g/d) attributable to decreased tubular reabsorption of filtered proteins; however, nephrotic-range albuminuria may occur in some conditions due to the development of secondary focal segmental glomerulosclerosis (FSGS). Renal ultrasonography may reveal changes of "medical renal disease," such as increased echogenicity of the renal parenchyma with loss of corticomedullary differentiation, prominence of the renal pyramids, and cortical scarring in some conditions. The predominant pathology in chronic TIN is interstitial fibrosis with patchy mononuclear cell infiltration and widespread tubular atrophy, luminal dilation, and thickening of tubular basement membranes. Because of the nonspecific nature of the histopathology, biopsy specimens rarely provide a specific diagnosis. Thus, diagnosis relies on careful analysis of history, drug or toxin exposure, associated symptoms, and imaging studies.

ACUTE INTERSTITIAL NEPHRITIS

In 1897, Councilman reported on eight cases of acute interstitial nephritis (AIN) in the Medical and Surgical Reports of the Boston City Hospital; three as a postinfectious complication of scarlet fever and two from diphtheria. Later, he described the lesion as "an acute inflammation of the kidney characterized by cellular and fluid exudation in the interstitial tissue, accompanied by, but not dependant on, degeneration

TABLE 340-1 CLASSIFICATION OF THE CAUSES OF TUBULOINTERSTITIAL DISEASES OF THE KIDNEY

Acute Tubulointerstitial Disorders

Acute Interstitial Nephritis

Therapeutic agents

- Antibiotics (β -lactams, sulfonamides, quinolones, vancomycin, erythromycin, linezolid, minocycline, rifampin, ethambutol, acyclovir)
- Nonsteroidal anti-inflammatory drugs, COX-2 inhibitors
- Diuretics (rarely thiazides, loop diuretics, triamterene)
- Anticonvulsants (phenytoin, valproate, carbamazepine, phenobarbital)
- Miscellaneous (proton pump inhibitors, H₂ blockers, captopril, mesalazine, indinavir, allopurinol, lenalidomide)

Infection

- Bacteria (*Streptococcus*, *Staphylococcus*, *Legionella*, *Salmonella*, *Brucella*, *Yersinia*, *Corynebacterium diphtheriae*)
- Viruses (EBV, CMV, hantavirus, polyomavirus, HIV)
- Miscellaneous (*Leptospira*, *Rickettsia*, *Mycoplasma*, *Histoplasma*)

Autoimmune

- Tubulointerstitial nephritis with uveitis (TINU)
- Sjögren's syndrome
- Systemic lupus erythematosus
- Granulomatous interstitial nephritis
- IgG4-related systemic disease
- Idiopathic autoimmune interstitial nephritis

Acute obstructive disorders

- Light chain cast nephropathy ("myeloma kidney")
- Acute phosphate nephropathy
- Acute urate nephropathy

Chronic Tubulointerstitial Disorders

- Vesicoureteral reflux/reflux nephropathy
- Sickle cell disease
- Chronic exposure to toxins or therapeutic agents
- Analgesics, especially those containing phenacetin
- Lithium
- Heavy metals (lead, cadmium)
- Aristolochic acid (Chinese herbal and Balkan endemic nephropathies)
- Calcineurin inhibitors (cyclosporine, tacrolimus)

Metabolic Disturbances

- Hypercalcemia and/or nephrocalcinosis
- Hyperuricemia
- Prolonged hypokalemia
- Hyperoxaluria
- Cystinosis (see Chap. 339)

Cystic and Hereditary Disorders (see Chap. 339)

- Polycystic kidney disease
- Nephronophthisis
- Adult medullary cystic disease
- Medullary sponge kidney

Miscellaneous

- Aging
- Chronic glomerulonephritis
- Chronic urinary tract obstruction
- Ischemia and vascular disease
- Radiation nephritis (rare)

Abbreviations: CMV, cytomegalovirus; COX, cyclooxygenase; EBV, Epstein-Barr virus.

of the epithelium; the exudation is not purulent in character, and the lesions may be both diffuse and focal." Today AIN is far more often encountered as an allergic reaction to a drug (Table 340-1). Immune-mediated AIN may also occur as part of a known autoimmune syndrome, but in some cases there is no identifiable cause despite features suggestive of an immunologic etiology (Table 340-1).

ALLERGIC INTERSTITIAL NEPHRITIS

Although biopsy-proven AIN accounts for no more than ~15% of cases of unexplained acute renal failure, this is likely a substantial underestimate of the true incidence. This is because potentially offending medications are more often identified and empirically discontinued in a patient noted to have a rising serum creatinine, without the benefit of a renal biopsy to establish the diagnosis of AIN.

Clinical Features The classic presentation of AIN, namely, fever, rash, peripheral eosinophilia, and oliguric renal failure occurring after 7–10 days of treatment with methicillin or another β -lactam antibiotic, is the exception rather than the rule. More often, patients are found incidentally to have a rising serum creatinine or present with symptoms attributable to acute renal failure (Chap. 334). Atypical reactions can occur, most notably nonsteroidal anti-inflammatory drug (NSAID)-induced AIN, in which fever, rash, and eosinophilia are rare, but acute renal failure with heavy proteinuria is common. A particularly severe and rapid-onset AIN may occur upon reintroduction of rifampin after a drug-free period. More insidious reactions to the agents listed in Table 340-1 may lead to progressive tubulointerstitial damage. Examples include proton pump inhibitors and, rarely, sulfonamide and 5-aminosalicylate (mesalazine and sulfasalazine) derivatives and antiretrovirals.

Diagnosis Finding otherwise unexplained renal failure with or without oliguria and exposure to a potentially offending agent usually points to the diagnosis. Peripheral blood eosinophilia adds supporting evidence but is present in only a minority of patients. Urinalysis reveals pyuria with white blood cell casts and hematuria. Urinary eosinophils are neither sensitive nor specific for AIN; therefore, testing is not recommended. Renal biopsy is generally not required for diagnosis but reveals extensive interstitial and tubular infiltration of leukocytes, including eosinophils.

TREATMENT ALLERGIC INTERSTITIAL NEPHRITIS

Discontinuation of the offending agent often leads to reversal of the renal injury. However, depending on the duration of exposure and degree of tubular atrophy and interstitial fibrosis that has occurred, the renal damage may not be completely reversible. Glucocorticoid therapy may accelerate renal recovery, but does not appear to impact long-term renal survival. It is best reserved for those cases with severe renal failure in which dialysis is imminent or if renal function continues to deteriorate despite stopping the offending drug (Fig. 340-1 and Table 340-2).

SJÖGREN'S SYNDROME

Sjögren's syndrome is a systemic autoimmune disorder that primarily targets the exocrine glands, especially the lacrimal and salivary glands, and thus results in symptoms, such as dry eyes and mouth, that constitute the "sicca syndrome" (Chap. 383). Tubulointerstitial nephritis with a predominant lymphocytic infiltrate is the most common renal manifestation of Sjögren's syndrome and can be associated with distal RTA, nephrogenic diabetes insipidus, and moderate renal failure. Diagnosis is strongly supported by positive serologic testing for anti-Ro (SS-A) and anti-La (SS-B) antibodies. A large proportion of patients with Sjögren's syndrome also have polyclonal hypergammaglobulinemia. Treatment

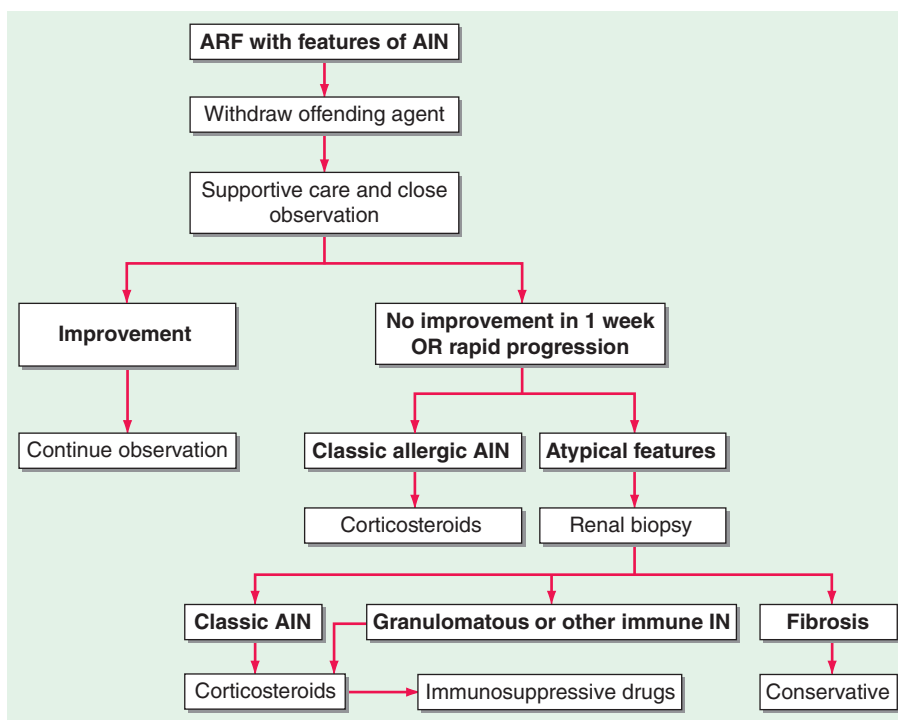


FIGURE 340-1 Algorithm for the treatment of allergic and other immune-mediated acute interstitial nephritis (AIN). ARF, acute renal failure; IN, interstitial nephritis. See text for immunosuppressive drugs used for refractory or relapsing AIN. (Modified from S Reddy, DJ Salant: *Ren Fail* 20:829, 1998.)

is initially with glucocorticoids, although patients may require maintenance therapy with azathioprine or mycophenolate mofetil to prevent relapse (Fig. 340-1 and Table 340-2).

TUBULOINTERSTITIAL NEPHRITIS WITH UVEITIS (TINU)

TINU is a systemic autoimmune disease of unknown etiology. It accounts for fewer than 5% of all cases of AIN, affects females three times more often than males, and has a median age of onset of 15 years. Its hallmark feature, in addition to a lymphocyte-predominant interstitial nephritis (Fig. 340-2), is a painful anterior uveitis, often bilateral and accompanied by blurred vision and photophobia. Diagnosis is

TABLE 340-2 INDICATIONS FOR CORTICOSTEROIDS AND IMMUNOSUPPRESSIVES IN INTERSTITIAL NEPHRITIS

Absolute Indications

- Sjögren's syndrome
- Sarcoidosis
- SLE interstitial nephritis
- Adults with TINU
- Idiopathic and other granulomatous interstitial nephritis

Relative Indications

- Drug-induced or idiopathic AIN with:
 - Rapid progression of renal failure
 - Diffuse infiltrates on biopsy
 - Impending need for dialysis
 - Delayed recovery
- Children with TINU
- Postinfectious AIN with delayed recovery (?)

Abbreviations: AIN, acute interstitial nephritis; SLE, systemic lupus erythematosus; TINU, tubulointerstitial nephritis with uveitis.

Source: Modified from S Reddy, DJ Salant: *Ren Fail* 20:829, 1998.

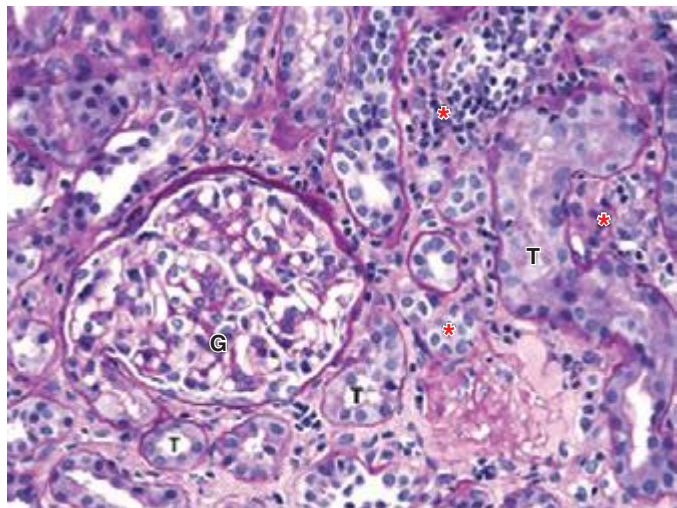


FIGURE 340-2 Acute interstitial nephritis (AIN) in a patient who presented with acute iritis, low-grade fever, erythrocyte sedimentation rate of 103, pyuria and cellular casts on urinalysis, and a newly elevated serum creatinine of 2.4 mg/dL. Both the iritis and AIN improved after intravenous methylprednisolone. This PAS-stained renal biopsy shows a mononuclear cell interstitial infiltrate (*asterisks*) and edema separating the tubules (T) and a normal glomerulus (G). Some of the tubules contain cellular debris and infiltrating inflammatory cells. The findings in this biopsy are indistinguishable from those that would be seen in a case of drug-induced AIN. PAS, Periodic acid–Schiff.

often confounded by the fact that the ocular symptoms precede or accompany the renal disease in only one-third of cases. Additional extrarenal features include fever, anorexia, weight loss, abdominal pain, and arthralgia. The presence of such symptoms as well as elevated creatinine, sterile pyuria, mild proteinuria, features of Fanconi's syndrome, and elevated erythrocyte sedimentation rate should raise suspicion for this disorder. Serologies suggestive of the more common autoimmune diseases are usually negative, and TINU is often a diagnosis of exclusion after other causes of uveitis and renal disease, such as Sjögren's syndrome, Behçet's disease, sarcoidosis, and systemic lupus erythematosus, have been considered. Clinical symptoms are typically self-limited in children, but are more apt to follow a relapsing course in adults. The renal and ocular manifestations generally respond well to oral glucocorticoids, although maintenance therapy with agents such as methotrexate, azathioprine, or mycophenolate may be necessary to prevent relapses (Fig. 340-1 and Table 340-2).

SYSTEMIC LUPUS ERYTHEMATOSUS

An interstitial mononuclear cell inflammatory reaction often accompanies the glomerular lesion in most cases of class III or IV lupus nephritis (Chap. 338), and deposits of immune complexes can be identified in tubule basement membranes in about 50% of cases. Occasionally, however, the tubulointerstitial inflammation predominates and may manifest with azotemia and type IV RTA rather than features of glomerulonephritis.

GRANULOMATOUS INTERSTITIAL NEPHRITIS

Some patients may present with features of AIN but follow a protracted and relapsing course. Renal biopsy in such patients reveals a more chronic inflammatory infiltrate with granulomas and multinucleated giant cells. Most often, no associated disease or cause is found; however, some of these cases may have or subsequently develop the pulmonary, cutaneous, or other systemic manifestations of *sarcoidosis* such as hypercalcemia. Most patients experience some improvement in renal function if treated early with glucocorticoids before the development of significant interstitial fibrosis and tubular atrophy (Table 340-2). Other immunosuppressive agents may be required for those who relapse frequently upon steroid withdrawal. Other immunosuppressive agents may be required for those who relapse frequently upon

steroid withdrawal (Fig. 340-1). Tuberculosis should be ruled out before starting treatment because this too is a rare cause of granulomatous interstitial nephritis.

IgG4-RELATED SYSTEMIC DISEASE

A form of AIN characterized by a dense inflammatory infiltrate containing IgG4-expressing plasma cells can occur as a part of a syndrome known as IgG4-related systemic disease. Autoimmune pancreatitis, sclerosing cholangitis, retroperitoneal fibrosis, and a chronic sclerosing sialadenitis (mimicking Sjögren's syndrome) may variably be present as well. Fibrotic lesions that form pseudotumors in the affected organs soon replace the initial inflammatory infiltrates and often lead to biopsy or excision for fear of true malignancy. Although the involvement of IgG4 in the pathogenesis is not understood, glucocorticoids have been successfully used as first-line treatment in this group of disorders, once they are correctly diagnosed.

IDIOPATHIC AIN

Some patients present with typical clinical and histologic features of AIN but have no evidence of drug exposure or clinical or serologic features of an autoimmune disease. The presence in some cases of autoantibodies to a tubular antigen, similar to that identified in rats with an induced form of interstitial nephritis, suggests that an autoimmune response may be involved. Like TINU and granulomatous interstitial nephritis, idiopathic AIN is responsive to glucocorticoid therapy but may follow a relapsing course requiring maintenance treatment with another immunosuppressive agent (Fig. 340-1 and Table 340-2).

INFECTION-ASSOCIATED AIN

AIN may also occur as a local inflammatory reaction to microbial infection (Table 340-1) and should be distinguished from acute bacterial pyelonephritis (Chap. 162). Acute bacterial pyelonephritis does not generally cause acute renal failure unless it affects both kidneys or causes septic shock. Presently, infection-associated AIN is most often seen in immunocompromised patients, particularly renal transplant recipients with reactivation of polyomavirus BK (Chaps. 169 and 337).

CRYSTAL DEPOSITION DISORDERS AND OBSTRUCTIVE TUBULOPATHIES

Acute renal failure may occur when crystals of various types are deposited in tubular cells and interstitium or when they obstruct tubules. Oliguric acute renal failure, often accompanied by flank pain from tubular obstruction, may occur in patients treated with sulfadiazine for toxoplasmosis, indinavir and atazanavir for HIV, and intravenous acyclovir for severe herpesvirus infections. Urinalysis reveals "sheaf of wheat" sulfonamide crystals, individual or parallel clusters of needle-shaped indinavir crystals, or red-green birefringent needle-shaped crystals of acyclovir. This adverse effect is generally precipitated by hypovolemia and is reversible with saline volume repletion and drug withdrawal. Distinct from the obstructive disease, a frank AIN from indinavir crystal deposition has also been reported.

Acute tubular obstruction is also the cause of oliguric renal failure in patients with *acute urate nephropathy*. It typically results from severe hyperuricemia from tumor lysis syndrome in patients with lympho- or myeloproliferative disorders treated with cytotoxic agents, but also may occur spontaneously before the treatment has been initiated (Chap. 331). Uric acid crystallization in the tubules and collecting system leads to partial or complete obstruction of the collecting ducts, renal pelvis, or ureter. A dense precipitate of birefringent uric acid crystals is found in the urine, usually in association with microscopic or gross hematuria. Prophylactic allopurinol reduces the risk of uric acid nephropathy but is of no benefit once tumor lysis has occurred. Once oliguria has developed, attempts to increase tubular flow and solubility of uric acid with alkaline diuresis may be of some benefit; however, emergent treatment with hemodialysis or rasburicase, a recombinant urate oxidase, is usually required to rapidly lower uric acid levels and restore renal function.

Calcium oxalate crystal deposition in tubular cells and interstitium may lead to permanent renal dysfunction in patients who survive ethylene glycol intoxication, in patients with enteric hyperoxaluria from

ileal resection or small-bowel bypass surgery, and in patients with hereditary hyperoxaluria (Chap. 342). *Acute phosphate nephropathy* is an uncommon but serious complication of oral Phospho-soda used as a laxative or for bowel preparation for colonoscopy. It results from calcium phosphate crystal deposition in tubules and interstitium and occurs especially in subjects with underlying renal impairment and hypovolemia. Consequently, Phospho-soda should be avoided in patients with chronic kidney disease.

LIGHT CHAIN CAST NEPHROPATHY

Patients with multiple myeloma may develop acute renal failure in the setting of hypovolemia, infection, or hypercalcemia or after exposure to NSAIDs or radiographic contrast media. The diagnosis of light chain cast nephropathy (LCCN)—commonly known as *myeloma kidney*—should be considered in patients who fail to recover when the precipitating factor is corrected or in any elderly patient with otherwise unexplained acute renal failure.

In this disorder, filtered monoclonal immunoglobulin light chains (Bence-Jones proteins) form intratubular aggregates with secreted Tamm-Horsfall protein in the distal tubule. Casts, in addition to obstructing the tubular flow in affected nephrons, incite a giant cell or foreign body reaction and can lead to tubular rupture, resulting in interstitial fibrosis (Fig. 340-3). Although LCCN generally occurs in patients with known multiple myeloma and a large plasma cell burden, the disorder should also be considered as a possible diagnosis in patients who have known monoclonal gammopathy even in the absence of frank myeloma. Filtered monoclonal light chains may also cause less pronounced renal manifestations in the absence of obstruction, due to direct toxicity to proximal tubular cells and intracellular crystal formation. This may result in isolated tubular disorders such as RTA or full Fanconi's syndrome.

Diagnosis Clinical clues to the diagnosis include anemia, bone pain, hypercalcemia, and an abnormally narrow anion gap due to hypoalbuminemia and hypergammaglobulinemia. Urinary dipsticks detect albumin but not immunoglobulin light chains; however, laboratory detection of increased amounts of protein in a spot urine specimen and a negative dipstick result are highly suggestive that the urine contains Bence-Jones protein. Serum and urine should both be sent for protein electrophoresis and for immunofixation for the detection and identification of a potential monoclonal band. A sensitive method is available to detect urine and serum free light chains.

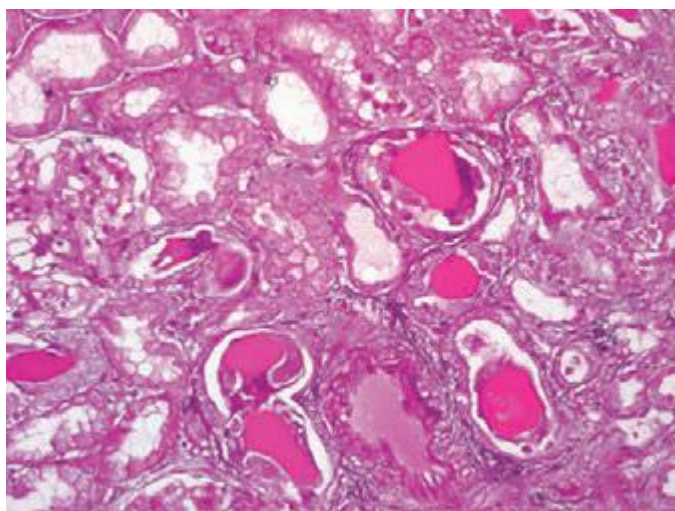


FIGURE 340-3 Histologic appearance of myeloma cast nephropathy. A hematoxylin-eosin–stained kidney biopsy shows many atrophic tubules filled with eosinophilic casts (consisting of Bence-Jones protein), which are surrounded by giant cell reactions. (Courtesy of Dr. Michael N. Koss, University of Southern California Keck School of Medicine; with permission.)

TREATMENT LIGHT CHAIN CAST NEPHROPATHY

The goals of treatment are to correct precipitating factors such as hypovolemia and hypercalcemia, discontinue potential nephrotoxic agents, and treat the underlying plasma cell dyscrasia (Chap. 136); plasmapheresis to remove light chains is of questionable value for LCCN.

LYMPHOMATOUS INFILTRATION OF THE KIDNEY

Interstitial infiltration by malignant B lymphocytes is a common autopsy finding in patients dying of chronic lymphocytic leukemia and non-Hodgkin's lymphoma; however, this is usually an incidental finding. Rarely, such infiltrates may cause massive enlargement of the kidneys and oliguric acute renal failure. Although high-dose glucocorticoids and subsequent chemotherapy often result in recovery of renal function, the prognosis in such cases is generally poor.

CHRONIC TUBULOINTERSTITIAL DISEASES

Improved occupational and public health measures, together with the banning of over-the-counter phenacetin-containing analgesics, has led to a dramatic decline in the incidence of chronic interstitial nephritis (CIN) from heavy metal—particularly lead and cadmium—exposure and analgesic nephropathy in North America. Today, CIN is most often the result of renal ischemia or secondary to a primary glomerular disease (Chap. 338). Other important forms of CIN are the result of developmental anomalies or inherited diseases such as reflux nephropathy or sickle cell nephropathy and may not be recognized until adolescence or adulthood. Although it is impossible to reverse damage that has already occurred, further deterioration may be prevented or at least slowed in such cases by treating glomerular hypertension, a common denominator in the development of secondary FSGS and progressive loss of functioning nephrons. Therefore, awareness and early detection of patients at risk may prevent them from developing end-stage renal disease (ESRD).

VESICoureTERAL REFLUX AND REFLUX NEPHROPATHY

Reflux nephropathy is the consequence of vesicoureteral reflux (VUR) or other urologic anomalies in early childhood. It was previously called *chronic pyelonephritis* because it was believed to result from recurrent urinary tract infections (UTIs) in childhood. VUR stems from abnormal retrograde urine flow from the bladder into one or both ureters and kidneys because of mislocated and incompetent ureterovesical valves (Fig. 340-4). Although high-pressure sterile reflux may impair normal growth of the kidneys, when coupled with recurrent UTIs in early childhood, the result is patchy interstitial scarring and tubular atrophy. Loss of functioning nephrons leads to hypertrophy of the remnant glomeruli and eventual secondary FSGS. Reflux nephropathy often goes unnoticed until early adulthood when chronic kidney disease is detected during routine evaluation or during pregnancy. Affected adults are frequently asymptomatic, but may give a history of prolonged bed-wetting or recurrent UTIs during childhood, and exhibit variable renal insufficiency, hypertension, mild to moderate proteinuria, and unremarkable urine sediment. When both kidneys are affected, the disease often progresses inexorably over several years to ESRD, despite the absence of ongoing urinary infections or reflux. A single affected kidney may go undetected, except for the presence of hypertension. Renal ultrasound in adults characteristically shows asymmetric small kidneys with irregular outlines, thinned cortices, and regions of compensatory hypertrophy (Fig. 340-4).

TREATMENT VESICoureTERAL REFLUX AND REFLUX NEPHROPATHY

Maintenance of sterile urine in childhood has been shown to limit scarring of the kidneys. Surgical reimplantation of the ureters into the bladder to restore competency is indicated in young children with persistent high-grade reflux but is ineffective and is

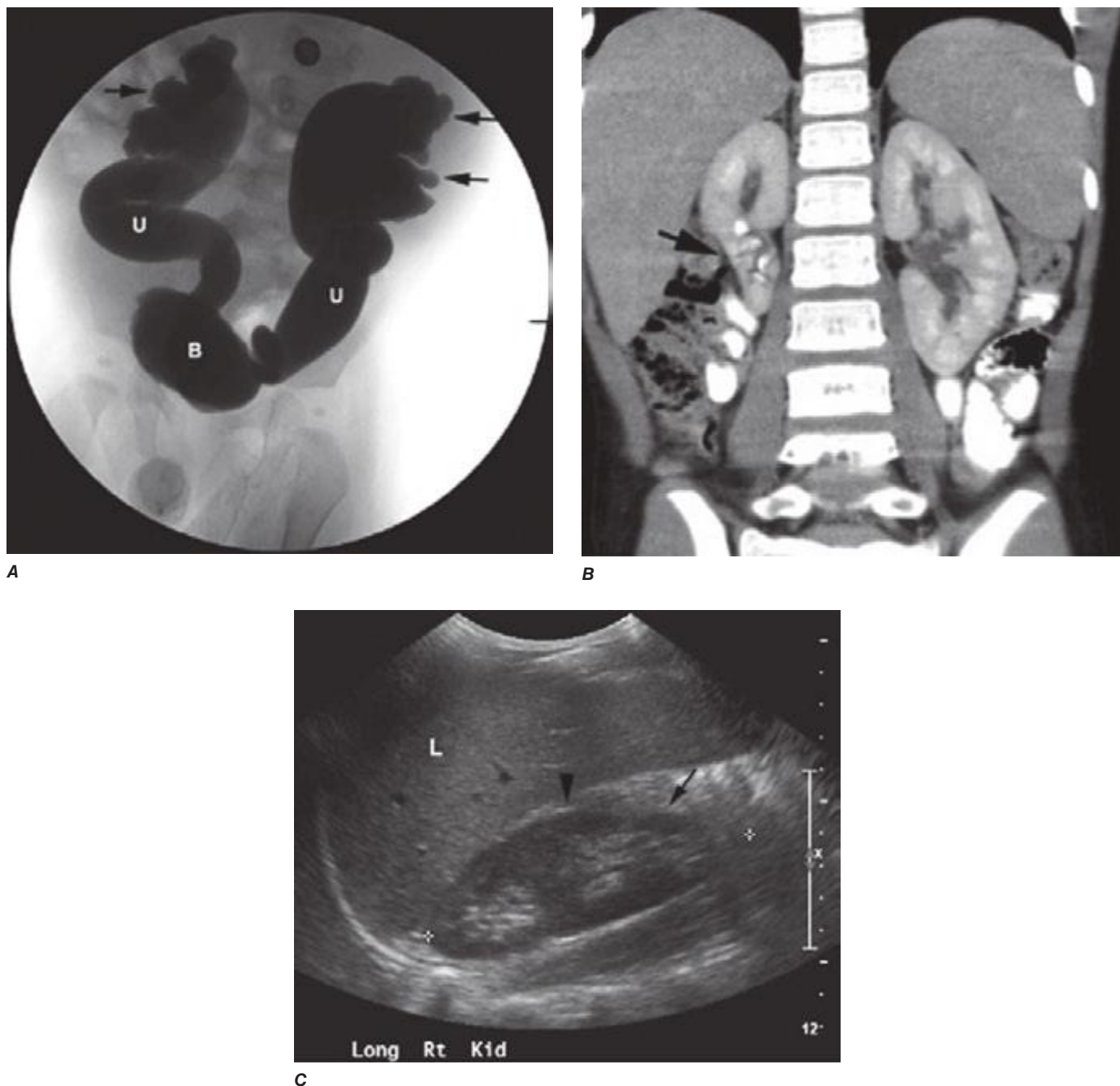


FIGURE 340-4 Radiographs of vesicoureteral reflux (VUR) and reflux nephropathy. **A.** Voiding cystourethrogram in a 7-month-old baby with bilateral high-grade VUR evidenced by clubbed calyces (*arrows*) and dilated tortuous ureters (U) entering the bladder (B). **B.** Abdominal computed tomography scan (coronal plane reconstruction) in a child showing severe scarring of the lower portion of the right kidney (*arrow*). **C.** Sonogram of the right kidney showing loss of parenchyma at the lower pole due to scarring (*arrow*) and hypertrophy of the mid-region (*arrowhead*). (Courtesy of Dr. George Gross, University of Maryland Medical Center; with permission.)

not indicated in adolescents or adults after scarring has occurred. Aggressive control of blood pressure with an angiotensin-converting enzyme inhibitor (ACEI) or angiotensin receptor blocker (ARB) and other agents is effective in reducing proteinuria and may significantly forestall further deterioration of renal function.

SICKLE CELL NEPHROPATHY

The pathogenesis and clinical manifestations of sickle cell nephropathy are described in [Chap. 341](#). Evidence of tubular injury may be evident in childhood and early adolescence in the form of polyuria due to decreased concentrating ability or type IV renal tubular acidosis years before there is significant nephron loss and proteinuria from secondary FSGS. Early recognition of these subtle renal abnormalities or

development of microalbuminuria in a child with sickle cell disease may warrant consultation with a nephrologist and/or therapy with low-dose ACEIs. Papillary necrosis may result from ischemia due to sickling of red cells in the relatively hypoxic and hypertonic medullary vasculature and present with gross hematuria and ureteric obstruction by sloughed ischemic papillae ([Table 340-3](#)).

TUBULOINTERSTITIAL ABNORMALITIES ASSOCIATED WITH GLOMERULONEPHRITIS

Primary glomerulopathies are often associated with damage to tubules and interstitium. This may occasionally be due to the same pathologic process affecting the glomerulus and tubulointerstitium, as is the case with immune-complex deposition in lupus nephritis. More often, however, chronic tubulointerstitial changes occur as a secondary

TABLE 340-3 MAJOR CAUSES OF PAPILLARY NECROSIS

Analgesic nephropathy
Sickle cell nephropathy
Diabetes with urinary tract infection
Prolonged NSAID use (rare)

Abbreviation: NSAID, nonsteroidal anti-inflammatory drug.

consequence of prolonged glomerular dysfunction. Potential mechanisms by which glomerular disease might cause tubulointerstitial injury include proteinuria-mediated damage to the epithelial cells, activation of tubular cells by cytokines and complement, or reduced peritubular blood flow leading to downstream tubulointerstitial ischemia, especially in the case of glomeruli that are globally obsolescent due to severe glomerulonephritis. It is often difficult to discern the initial cause of injury by renal biopsy in a patient who presents with advanced renal disease in this setting.

ANALGESIC NEPHROPATHY

Analgesic nephropathy results from the long-term use of compound analgesic preparations containing phenacetin (banned in the United States since 1983), aspirin, and caffeine. In its classic form, analgesic nephropathy is characterized by renal insufficiency, papillary necrosis (Table 340-3) attributable to the presumed concentration of the drug to toxic levels in the inner medulla, and a radiographic constellation of small, scarred kidneys with papillary calcifications best appreciated by computed tomography (Fig. 340-5). Patients may also have polyuria due to impaired concentrating ability and non-anion-gap metabolic acidosis from tubular damage. Shedding of a sloughed necrotic papilla can cause gross hematuria and ureteric colic due to ureteral obstruction. Individuals with ESRD as a result of analgesic nephropathy are at increased risk of a urothelial malignancy compared to patients with other causes of renal failure. Recent cohort studies in individuals with normal baseline renal function suggest that the moderate chronic use of current analgesic preparations available in the United States, including acetaminophen and NSAIDs, does not seem to cause the constellation of findings known as analgesic nephropathy, although volume-depleted individuals and those with chronic kidney disease are at higher risk of NSAID-related renal toxicity. Nonetheless, it is recommended that heavy users of acetaminophen and NSAIDs be screened for evidence of renal disease.

ARISTOLOCHIC ACID NEPHROPATHY

Two seemingly unrelated forms of CIN, Chinese herbal nephropathy and Balkan endemic nephropathy, have recently been linked by the underlying etiologic agent aristolochic acid and are now collectively termed aristolochic acid nephropathy (AAN). In Chinese herbal

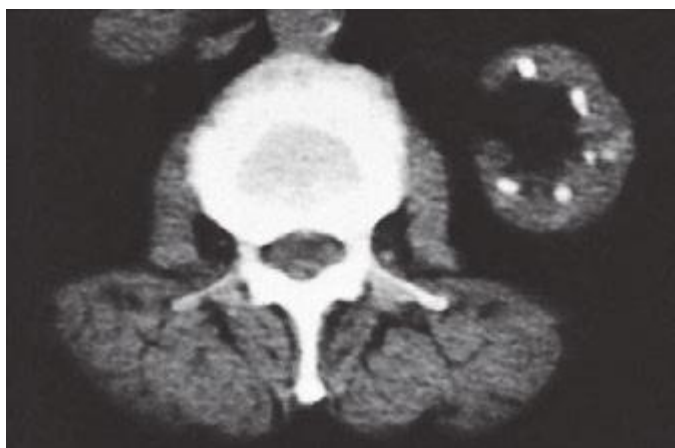


FIGURE 340-5 Radiologic appearance of analgesic nephropathy.

A noncontrast computed tomography scan shows an atrophic left kidney with papillary calcifications in a garland pattern. (Reprinted by permission from Macmillan Publishers, Ltd., MM Elseviers et al: *Kidney International* 48:1316, 1995.)

nephropathy, first described in the early 1990s in young women taking traditional Chinese herbal preparations as part of a weight-loss regimen, one of the offending agents has been identified as aristolochic acid, a known carcinogen from the plant *Aristolochia*. Multiple *Aristolochia* species have been used in traditional herbal remedies for centuries and continue to be available despite official bans on their use in many countries. Molecular evidence has also implicated aristolochic acid in Balkan endemic nephropathy, a chronic tubulointerstitial nephritis found primarily in towns along the tributaries of the Danube River and first described in the 1950s. Although the exact route of exposure is not known with certainty, contamination of local grain preparations with the seeds of *Aristolochia* species seems most likely. Aristolochic acid, after prolonged exposure, produces renal interstitial fibrosis with a relative paucity of cellular infiltrates. The urine sediment is bland, with rare leukocytes and only mild proteinuria. Anemia may be disproportionately severe relative to the level of renal dysfunction. Definitive diagnosis of AAN requires two of the following three features: characteristic histology on kidney biopsy; confirmation of aristolochic acid ingestion; and detection of aristolactam-DNA adducts in kidney or urinary tract tissue. These latter lesions represent a molecular signature of aristolochic acid-derived DNA damage and often consist of characteristic A:T-to-T:A transversions. Due to this mutagenic activity, AAN is associated with a very high incidence of upper urinary tract urothelial cancers, with risk related to cumulative dose. Surveillance with computed tomography, ureteroscopy, and urine cytology is warranted, and consideration should be given to bilateral nephroureterectomy once a patient has reached ESRD.

KARYOMEGALIC INTERSTITIAL NEPHRITIS

Karyomegalic interstitial nephritis is an unusual form of slowly progressive chronic kidney disease with mild proteinuria, interstitial fibrosis, tubular atrophy, and oddly enlarged nuclei of proximal tubular epithelial cells. It has been linked to mutations in *FANL1*, a nuclease involved in DNA repair, which may render carriers of the mutation susceptible to environmental DNA-damaging agents.

LITHIUM-ASSOCIATED NEPHROPATHY

The use of lithium salts for the treatment of manic-depressive illness may have several renal sequelae, the most common of which is nephrogenic diabetes insipidus manifesting as polyuria and polydipsia. Lithium accumulates in principal cells of the collecting duct by entering through the epithelial sodium channel (ENaC), where it inhibits glycogen synthase kinase 3 β and downregulates vasopressin-regulated aquaporin water channels. Less frequently, chronic tubulointerstitial nephritis develops after prolonged (>10–20 years) lithium use and is most likely to occur in patients who have experienced repeated episodes of toxic lithium levels. Findings on renal biopsy include interstitial fibrosis and tubular atrophy that are out of proportion to the degree of glomerulosclerosis or vascular disease, a sparse lymphocytic infiltrate, and small cysts or dilation of the distal tubule and collecting duct that are highly characteristic of this disorder. The degree of interstitial fibrosis correlates with both duration and cumulative dose of lithium. Individuals with lithium-associated nephropathy are typically asymptomatic, with minimal proteinuria, few urinary leukocytes, and normal blood pressure. Some patients develop more severe proteinuria due to secondary FSGS, which may contribute to further loss of renal function.

TREATMENT LITHIUM-ASSOCIATED NEPHROPATHY

Renal function should be followed regularly in patients taking lithium, and caution should be exercised in patients with underlying renal disease. The use of amiloride to inhibit lithium entry via ENaC has been effective to prevent and treat lithium-induced nephrogenic diabetes insipidus, but it is not clear if it will prevent lithium-induced CIN. Once lithium-associated nephropathy is detected, the discontinuation of lithium in attempt to forestall further renal deterioration can be problematic, as lithium is an effective mood stabilizer that is often incompletely substituted by other agents. Furthermore, despite discontinuation of lithium, chronic

renal disease in such patients is often irreversible and can slowly progress to ESRD. The most prudent approach is to monitor lithium levels frequently and adjust dosing to avoid toxic levels (preferably <1 meq/L). This is especially important because lithium is cleared less effectively as renal function declines. In patients who develop significant proteinuria, ACEI or ARB treatment should be initiated.

CALCINEURIN-INHIBITOR NEPHROTOXICITY

The calcineurin inhibitor (CNI) immunosuppressive agents cyclosporine and tacrolimus can cause both acute and chronic renal injury. Acute forms can result from vascular causes such as vasoconstriction or the development of thrombotic microangiopathy, or can be due to a toxic tubulopathy. Chronic CNI-induced renal injury is typically seen in solid organ (including heart-lung and liver) transplant recipients and manifests with a slow but irreversible reduction of glomerular filtration rate, with mild proteinuria and arterial hypertension. Hyperkalemia is a relatively common complication and is caused, in part, by tubular resistance to aldosterone. The histologic changes in renal tissue include patchy interstitial fibrosis and tubular atrophy, often in a “striped” pattern. In addition, the intrarenal vasculature often demonstrates hyalinosis, and focal glomerulosclerosis can be present as well. Similar changes may occur in patients receiving CNIs for autoimmune diseases, although the doses are generally lower than those used for organ transplantation. Dose reduction or CNI avoidance appears to mitigate the chronic tubulointerstitial changes, but may increase the risk of rejection and graft loss.

HEAVY METAL (LEAD) NEPHROPATHY

Heavy metals, such as lead or cadmium, can lead to a chronic tubulointerstitial process after prolonged exposure. The disease entity is no longer commonly diagnosed, because such heavy metal exposure has been greatly reduced due to the known health risks from lead and the consequent removal of lead from most commercial products and fuels. Nonetheless, occupational exposure is possible in workers involved in the manufacture or destruction of batteries, removal of lead paint, or manufacture of alloys and electrical equipment (cadmium) in countries where industrial regulation is less stringent. In addition, ingestion of moonshine whiskey distilled in lead-tainted containers has been one of the more frequent sources of lead exposure.

Early signs of chronic lead intoxication are attributable to proximal tubule dysfunction, particularly hyperuricemia as a result of diminished urate secretion. The triad of “saturnine gout,” hypertension, and renal insufficiency should prompt a practitioner to ask specifically about lead exposure. Unfortunately, evaluating lead burden is not as straightforward as ordering a blood test; the preferred methods involve measuring urinary lead after infusion of a chelating agent or by radiographic fluoroscopy of bone. Several recent studies have shown an association between chronic low-level lead exposure and decreased renal function, although either of these two factors may have been the primary event. In those patients who have CIN of unclear origin and an elevated total body lead burden, repeated treatments of lead chelation therapy have been shown to slow the decline in renal function.

METABOLIC DISORDERS

Disorders leading to excessively high or low levels of certain electrolytes and products of metabolism can also lead to chronic kidney disease if untreated.

CHRONIC URIC ACID NEPHROPATHY

The constellation of pathologic findings that represent *gouty nephropathy* are very uncommon nowadays and are more of historical interest than clinical importance, as gout is typically well managed with allopurinol and other agents. However, there is emerging evidence that hyperuricemia is an independent risk factor for the development of chronic kidney disease, perhaps through endothelial damage. The complex interactions of hyperuricemia, hypertension, and renal failure are still incompletely understood.

Presently, gouty nephropathy is most likely to be encountered in patients with severe tophaceous gout and prolonged hyperuricemia from a hereditary disorder of purine metabolism (**Chap. 431e**). This should be distinguished from juvenile hyperuricemic nephropathy, a form of medullary cystic kidney disease caused by mutations in uromodulin (UMOD) (**Chap. 339**). Histologically, the distinctive feature of gouty nephropathy is the presence of crystalline deposits of uric acid and monosodium urate salts in the kidney parenchyma. These deposits not only cause intrarenal obstruction but also incite an inflammatory response, leading to lymphocytic infiltration, foreign-body giant cell reaction, and eventual fibrosis, especially in the medullary and papillary regions of the kidney. Since patients with gout frequently suffer from hypertension and hyperlipidemia, degenerative changes of the renal arterioles may constitute a striking feature of the histologic abnormality, out of proportion to the other morphologic defects. Clinically, gouty nephropathy is an insidious cause of chronic kidney disease. Early in its course, glomerular filtration rate may be near normal, often despite morphologic changes in medullary and cortical interstitium, proteinuria, and diminished urinary concentrating ability. Treatment with allopurinol and urine alkalinization is generally effective in preventing uric acid nephrolithiasis and the consequences of recurrent kidney stones; however, gouty nephropathy may be intractable to such measures. Furthermore, the use of allopurinol in asymptomatic hyperuricemia has not been consistently shown to improve renal function.

HYPERCALCEMIC NEPHROPATHY

(See also **Chap. 424**) Chronic hypercalcemia, as occurs in primary hyperparathyroidism, sarcoidosis, multiple myeloma, vitamin D intoxication, or metastatic bone disease, can cause tubulointerstitial disease and progressive renal failure. The earliest lesion is a focal degenerative change in renal epithelia, primarily in collecting ducts, distal tubules, and loops of Henle. Tubular cell necrosis leads to nephron obstruction and stasis of intrarenal urine, favoring local precipitation of calcium salts and infection. Dilation and atrophy of tubules eventually occur, as do interstitial fibrosis, mononuclear leukocyte infiltration, and interstitial calcium deposition (nephrocalcinosis). Calcium deposition may also occur in glomeruli and the walls of renal arterioles.


Clinically, the most striking defect is an inability to maximally concentrate the urine, due to reduced collecting duct responsiveness to arginine vasopressin and defective transport of sodium and chloride in the loop of Henle. Reductions in both glomerular filtration rate and renal blood flow can occur, both in acute and in prolonged hypercalcemia. Eventually, uncontrolled hypercalcemia leads to severe tubulointerstitial damage and overt renal failure. Abdominal x-rays may demonstrate nephrocalcinosis as well as nephrolithiasis, the latter due to the hypercalciuria that often accompanies hypercalcemia.

Treatment consists of reducing the serum calcium concentration toward normal and correcting the primary abnormality of calcium metabolism (**Chap. 424**). Renal dysfunction of acute hypercalcemia may be completely reversible. Gradual progressive renal insufficiency related to chronic hypercalcemia, however, may not improve even with correction of the calcium disorder.

HYPOKALEMIC NEPHROPATHY

Patients with prolonged and severe hypokalemia from chronic laxative or diuretic abuse, surreptitious vomiting, or primary aldosteronism may develop a reversible tubular lesion characterized by vacuolar degeneration of proximal and distal tubular cells. Eventually, tubular atrophy and cystic dilation accompanied by interstitial fibrosis may ensue, leading to irreversible chronic kidney disease. Timely correction of the hypokalemia will prevent further progression, but persistent hypokalemia can cause ESRD.

GLOBAL PERSPECTIVE

 The causes of acute and chronic interstitial nephritis vary widely across the globe. Analgesic nephropathy continues to be seen in countries where phenacetin-containing compound analgesic preparations are readily available. Adulterants in unregulated herbal and traditional medicaments pose a threat of toxic interstitial nephritis, as

exemplified by aristolochic acid contamination of herbal slimming preparations. Contamination of food sources with toxins, such as the recent outbreak of nephrolithiasis and acute renal failure from melamine contamination of infant milk formula, poses a continuing risk. Large-scale exposure to aristolochic acid remains prevalent in many Asian countries where traditional herbal medicine use is common. Although industrial exposure to lead and cadmium has largely disappeared as a cause of chronic interstitial nephritis in developed nations, it remains a risk for nephrotoxicity in countries where such exposure is less well controlled. New endemic forms of chronic kidney disease continue to be described, such as the nephropathy found among Pacific coastal plantation workers in Central America, which may be related to repetitive heat exposure and fluid losses.

341 Vascular Injury to the Kidney

Nelson Leung, Stephen C. Textor

The renal circulation is complex and is characterized by a highly perfused arteriolar network, reaching cortical glomerular structures adjacent to lower-flow vasa recta that descend into medullary segments. Disorders of the larger vessels, including renal artery stenosis and atheroembolic disease, are discussed elsewhere (Chap. 354). This chapter examines primary disorders of the renal microvessels, many of which are associated with thrombosis and hemolysis.

THROMBOTIC MICROANGIOPATHY

Thrombotic microangiopathy (TMA) is characterized by injured endothelial cells that are thickened, swollen, or detached mainly from arterioles and capillaries. Platelet and hyaline thrombi causing partial or complete occlusion are integral to the histopathology of TMA. TMA is usually the result of microangiopathic hemolytic anemia (MAHA), with its typical features of thrombocytopenia and schistocytes. In the kidney, TMA is characterized by swollen endocapillary cells (endotheliosis), fibrin thrombi, platelet plugs, arterial intimal fibrosis, and a membranoproliferative pattern. Fibrin thrombi may extend into the arteriolar vascular pole, producing glomerular collapse and at times cortical necrosis. In kidneys that recover from acute TMA, secondary focal segmental glomerulosclerosis may be seen. Diseases associated with this lesion include thrombotic thrombocytopenic purpura (TTP), hemolytic-uremic syndrome (HUS), malignant hypertension, scleroderma renal crisis, antiphospholipid syndrome, preeclampsia/HELLP (hemolysis, elevated liver enzymes, low platelet count) syndrome, HIV infection, and radiation nephropathy.

HEMOLYTIC-UREMIC SYNDROME/THROMBOTIC THROMBOCYTOPENIC PURPURA

HUS and TTP are the prototypes for MAHA. Historically, HUS and TTP were distinguished mainly by their clinical and epidemiologic differences. TTP develops more commonly in adults and was thought to include neurologic involvement more often. HUS occurs more commonly in children, particularly when associated with hemorrhagic diarrhea. However, atypical HUS (aHUS) can first appear in adulthood, and better testing has revealed that neurologic involvement is as common in HUS as in TTP. Accordingly, HUS and TTP now should be differentiated and treated according to their specific pathophysiologic features.

Hemolytic-Uremic Syndrome HUS is loosely defined by the presence of MAHA and renal impairment. At least four variants are recognized. The most common is Shiga toxin-producing *Escherichia coli* (STEC) HUS, which is also known as D⁺HUS or enterohemorrhagic *E. coli* (EHEC) HUS. Most cases involve children <5 years of age, but adults also are susceptible, as evidenced by a 2011 outbreak in northern Europe. Diarrhea, often bloody, precedes MAHA within

1 week in >80% of cases. Abdominal pain, cramping, and vomiting are frequent, whereas fever is typically absent. Neurologic symptoms, including dysphasia, hyperreflexia, blurred vision, memory deficits, encephalopathy, perseveration, and agraphia, often develop, especially in adults. Seizures and cerebral infarction can occur in severe cases. STEC HUS is caused by the Shiga toxins (Stx1 and Stx2), which are also referred to as *verotoxins*. These toxins are produced by certain strains of *E. coli* and *Shigella dysenteriae*. In the United States and Europe, the most common STEC strain is O157:H7, but HUS due to other strains (O157:H⁻, O111:H⁻, O26:H11/H⁻, O145:H28, and O104:H4) has occurred. After entry into the circulation, Shiga toxin binds to the glycolipid surface receptor globotriaosylceramide (Gb3), which is richly expressed on cells of the renal microvasculature. Upon binding, the toxin enters the cells, inducing inflammatory cytokines (interleukin 8 [IL-8], monocyte chemoattractant protein 1 [MCP-1], and stromal cell-derived factor 1 [SDF-1]) and chemokine receptors (CXCR4 and CXCR7); this action results in platelet aggregation and the microangiopathic process. *Streptococcus pneumoniae* can also cause HUS. Certain strains produce a neuraminidase that cleaves the N-acetylneuraminic acid moieties covering the Thomsen-Friedenreich antigen on platelets and endothelial cells. Exposure of this normally cryptic antigen to preformed IgM results in severe MAHA.

Atypical HUS is the result of congenital complement dysregulation. The affected patients have the low C3 and normal C4 levels characteristic of alternative pathway activation. Factor H deficiency, the most common defect, has been linked to families with aHUS. Factor H competes with factor B to prevent the formation of C3bBb and acts as a cofactor for factor I, which proteolytically degrades C3b. More than 70 mutations of the factor H gene have been identified. Most are missense mutations that produce abnormalities in the C-terminus region, affecting its binding to C3b but not its concentration. Other mutations result in low levels or the complete absence of the protein. Deficiencies in other complement-regulatory proteins, such as factor I, factor B, membrane cofactor protein (CD46), C3, complement factor H-related protein 1 (CFHR1), CFHR3, CFHR5, and thrombomodulin, have also been reported. Finally, an autoimmune variant of aHUS has been discovered. DEAP (deficient for CFHR protein and positive for factor H autoantibody) HUS occurs when an autoantibody to factor H is formed. DEAP HUS is often associated with a deletion of an 84-kb fragment of the chromosome that encodes for CFHR1 and CFHR3. The autoantibody blocks the binding of factor H to C3b and surface-bound C3 convertase.

Thrombotic Thrombocytopenic Purpura Traditionally, TTP is characterized by the pentad: MAHA, thrombocytopenia, neurologic symptoms, fever, and renal failure. The pathophysiology of TTP involves the accumulation of ultra-large multimers of von Willebrand factor as a result of the absence or markedly decreased activity (<5–10%) of the plasma protease ADAMTS13, a disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13. These ultra-large multimers form clots and shear erythrocytes, resulting in MAHA; however, the absence of ADAMTS13 alone may not itself produce TTP. Often, an additional trigger (such as infection, surgery, pancreatitis, or pregnancy) is required to initiate clinical TTP.

Data from the Oklahoma TTP/HUS Registry suggest an incidence rate of 11.3 cases/10⁶ patients in the United States. The median age of onset is 40 years. The incidence is more than nine times higher among blacks than among non-blacks. Like that of systemic lupus erythematosus, the incidence of TTP is nearly three times higher among women than among men. If untreated, TTP has a mortality rate exceeding 90%. Even with modern therapy, 20% of patients die within the first month from complications of microvascular thrombosis.

The classic form of TTP is idiopathic TTP, which is usually the result of a deficiency in ADAMTS13. While TTP had traditionally been associated with infection, malignancy, and intense inflammation (e.g., pancreatitis), ADAMTS13 activity usually is not decreased in these conditions. In idiopathic TTP, the formation of an autoantibody to ADAMTS13 (IgG or IgM) either increases its clearance or inhibits its activity. Upshaw-Schulman syndrome is a hereditary condition characterized by congenital deficiency of ADAMTS13. TTP in these

patients can start within the first weeks of life but in some instances may not present until the patient is several years of age. Both environmental and genetic factors are thought to influence the development of TTP. Plasma transfusion is an effective strategy for prevention and treatment.

Drug-induced TMA is a recognized complication of treatment with some chemotherapeutic agents, immunosuppressive agents, antiplatelet agents, and quinine. Two different mechanisms have been described. Endothelial damage (pathologically similar to that in HUS) is the main cause of the TMA that develops in association with chemotherapeutic agents (e.g., mitomycin C, gemcitabine) and immunosuppressive agents (cyclosporine, tacrolimus, and sirolimus). This process is usually dose-dependent. Alternatively, TMA may develop as a result of drug-induced autoantibodies. This form is less likely to be dose-dependent and can, in fact, occur after a single dose in patients with previous exposure. Ticlopidine produces TTP by inducing an autoantibody to ADAMTS13, but ADAMTS13 deficiency is found in fewer than half of patients with clopidogrel-associated TTP. Quinine appears to induce autoantibodies to granulocytes, lymphocytes, endothelial cells, and platelet glycoprotein IIB/IX or IIB/IIIa complexes, but not to ADAMTS13. Quinine-associated TTP is more common among women. TMA has been reported with drugs that inhibit vascular endothelial growth factor, such as bevacizumab; the mechanism is not completely understood.

TREATMENT HUS/TTP

Treatment should be based on pathophysiology. Autoantibody-mediated TTP and DEAP HUS respond to plasma exchange or plasmapheresis. In addition to removing the autoantibodies, plasma exchange with fresh-frozen plasma replaces ADAMTS13. Twice-daily plasma exchanges with administration of vincristine and rituximab may be effective in refractory cases. Plasma infusion is usually sufficient to replace the ADAMTS13 in Upshaw-Schülman syndrome. Plasma exchange should be considered if larger volumes are necessary. Drug-induced TMA secondary to endothelial damage typically does not respond to plasma exchange and is treated primarily by discontinuing use of the agent and providing supportive care. Similarly, STEC HUS should be treated with supportive measures. Plasma exchange has not been found to be effective. Antimotility agents and antibiotics increase the incidence of HUS among children, but azithromycin was recently found to decrease the duration of bacterial shedding by adults. Eculizumab is a monoclonal antibody to C5 that is approved for use in aHUS, for which ongoing therapy may be necessary. Plasma infusion/exchange may play a role in aHUS by replacing complement-regulatory proteins. Antibiotics and washed red cells should be given in neuraminidase-associated HUS, and plasmapheresis may be helpful. However, plasma and whole-blood transfusion should be avoided since these products contain IgM, which may exacerbate MAHA. Finally, combined factor H and ADAMTS13 deficiency have been reported. The affected patients are generally less responsive to plasma infusion, a result illustrating the complexity of the management of these cases.

HEMATOPOIETIC STEM CELL TRANSPLANTATION–ASSOCIATED THROMBOTIC MICROANGIOPATHY (HSCT-TMA)

HSCT-TMA develops after HSCT, with an incidence of 8.2%. Etiologic factors include conditioning regimens, immunosuppression, infections, and graft-versus-host disease. Other risk factors include female sex and human leukocyte antigen (HLA)–mismatched donor grafts. HSCT-TMA usually occurs within the first 100 days of HSCT. **Table 341-1** lists definitions of HSCT-TMA currently used for clinical trials. Diagnosis may be difficult since thrombocytopenia, anemia, and renal insufficiency are common after HSCT. HSCT-TMA carries a high mortality rate (75% within 3 months). The majority of patients have >5% ADAMTS13 activity, and plasma exchange is beneficial in <50% of patients. Discontinuation of calcineurin inhibitors and substitution with daclizumab (antibody to

TABLE 341-1 CRITERIA FOR ESTABLISHING MICROANGIOPATHIC KIDNEY INJURY ASSOCIATED WITH HEMATOPOIETIC STEM CELL TRANSPLANTATION

International Working Group	Blood and Marrow Transplant Clinical Trials Network Toxicity Committee
>4% schistocytes in the blood	RBC fragmentation and at least 2 schistocytes per high-power field
De novo, prolonged, or progressive thrombocytopenia	Concurrent increase in LDH concentration above baseline
A sudden and persistent increase in LDH concentration	Negative direct and indirect Coombs test
Decrease in hemoglobin level or increased RBC transfusion requirement	Concurrent renal and/or neurologic dysfunction without other explanations
Decrease in haptoglobin concentration	

Note: These features underscore the need to identify pathways of hemolysis and thrombocytopenia that accompany deterioration of kidney function.

Abbreviations: LDH, lactate dehydrogenase; RBC, red blood cell.

the IL-2 receptor) are recommended. Treatment with rituximab and defibrotide may also be helpful.

HIV-RELATED TMA

HIV-related TMA is a complication encountered mainly before widespread use of highly active antiretroviral therapy. It is seen in patients with advanced AIDS and low CD4+ T cell counts although it can be the first manifestation of HIV infection. The presence of MAHA, thrombocytopenia, and renal failure are suggestive, but renal biopsy is required for diagnosis since other renal diseases are also associated with HIV infection. Thrombocytopenia may prohibit renal biopsy in some patients. The mechanism of injury is unclear, although HIV can induce apoptosis in endothelial cells. ADAMTS13 activity is not reduced in these patients. Cytomegalovirus co-infection may also be a risk factor. Effective antiviral therapy is key, while plasma exchange should be limited to patients who have evidence of TTP.

RADIATION NEPHROPATHY

Either local or total body irradiation can produce microangiopathic injury. The kidney is one of the most radiosensitive organs, and injury can result with as little as 4–5 Gy. Such injury is characterized by renal insufficiency, proteinuria, and hypertension usually developing ≥6 months after radiation exposure. Renal biopsy reveals classic TMA with damage to glomerular, tubular, and vascular cells, but systemic evidence of MAHA is uncommon. Because of its high incidence after allogeneic HSCT, radiation nephropathy is often referred to as *bone marrow transplant nephropathy*. No specific therapy is available, although observational evidence supports renin-angiotensin system blockade.

SCLEDERMA (PROGRESSIVE SYSTEMIC SCLEROSIS)

Kidney involvement is common (up to 52%) in patients with widespread scleroderma, with 20% of cases resulting directly from scleroderma renal crisis. Other renal manifestations in scleroderma include transient (prerenal) or medication-related forms of acute kidney injury (e.g., associated with D-penicillamine, nonsteroidal anti-inflammatory drugs, or cyclosporine). Scleroderma renal crisis occurs in 12% of patients with diffuse systemic sclerosis but in only 2% of those with limited systemic sclerosis. Scleroderma renal crisis is the most severe manifestation of renal involvement, and is characterized by accelerated hypertension, a rapid decline in renal function, nephrotic proteinuria, and hematuria. Retinopathy and encephalopathy may accompany the hypertension. Salt and water retention with microvascular injury can lead to pulmonary edema. Cardiac manifestations, including myocarditis, pericarditis, and arrhythmias, denote an especially poor prognosis. Although MAHA is present in more half of patients, coagulopathy is rare.

The renal lesion in scleroderma renal crisis is characterized by arcuate artery intimal and medial proliferation with luminal narrowing.

This lesion is described as “onion-skinning” and can be accompanied by glomerular collapse due to reduced blood flow. Histologically, scleroderma renal crisis is indistinguishable from malignant hypertension, with which it can coexist. Fibrinoid necrosis and thrombosis are common. Before the availability of angiotensin-converting enzyme (ACE) inhibitors, the mortality rate for scleroderma renal crisis was >90% at 1 month. Introduction of renin-angiotensin system blockade has lowered the mortality rate to 30% at 3 years. Nearly two-thirds of patients with scleroderma renal crisis may require dialysis support, with recovery of renal function in 50% (median time, 1 year). Glomerulonephritis and vasculitis associated with antineutrophil cytoplasmic antibodies and systemic lupus erythematosus have been described in patients with scleroderma. An association has been found with a speckled pattern of antinuclear antibodies and with antibodies to RNA polymerases I and III. Anti-U3-RNP may identify young patients at risk for scleroderma renal crisis. Anticentromere antibody, in contrast, is a negative predictor of this disorder. Because of the overlap between scleroderma renal crisis and other autoimmune disorders, a renal biopsy is recommended for patients with atypical renal involvement, especially if hypertension is absent.

Treatment with ACE inhibition is the first-line therapy unless contraindicated. The goal of therapy is to reduce systolic and diastolic blood pressure by 20 mmHg and 10 mmHg, respectively, every 24 h until blood pressure is normal. Additional antihypertensive therapy may be given once the dose of drug for ACE inhibition is maximized. Both ACE inhibitors and angiotensin II receptor antagonists are effective, although data suggest that treatment with ACE inhibitors is superior. ACE inhibition alone does not prevent scleroderma renal crisis, but it does reduce the impact of hypertension. Intravenous iloprost has been used in Europe for blood pressure management and improvement of renal perfusion. Kidney transplantation is not recommended for 2 years after the start of dialysis since delayed recovery may occur.

ANTIPHOSPHOLIPID SYNDROME

Antiphospholipid syndrome (**Chap. 379**) can be either primary or secondary to systemic lupus erythematosus. It is characterized by a predisposition to systemic thrombosis (arterial and venous) and fetal morbidity mediated by antiphospholipid antibodies—mainly anticardiolipin antibodies (IgG, IgM, or IgA), lupus anticoagulant, or anti- β -2 glycoprotein I antibodies (anti β 2GPI). Patients with both anticardiolipin antibodies and anti β 2GPI appear to have the highest risk of thrombosis. The vascular compartment within the kidney is the main site of renal involvement. Arteriosclerosis is commonly present in the arcuate and intralobular arteries. In the intralobular arteries, fibrous intimal hyperplasia characterized by intimal thickening secondary to intense myofibroblastic intimal cellular proliferation with extracellular matrix deposition is frequently seen along with onion-skinning. Arterial and arteriolar fibrous and fibrocellular occlusions are present in more than two-thirds of biopsy samples. Cortical necrosis and focal cortical atrophy may result from vascular occlusion. TMA is commonly present in renal biopsies, although signs of MAHA and platelet consumption are usually absent. TMA is especially common in the catastrophic variant of antiphospholipid syndrome. In patients with secondary antiphospholipid syndrome, other glomerulopathies may be present, including membranous nephropathy, minimal change disease, focal segmental glomerulosclerosis, and pauci-immune crescentic glomerulonephritis.

Large vessels can be involved in antiphospholipid syndrome and may form the proximal nidus near the ostium for thrombosis of the renal artery. Renal vein thrombosis can occur and should be suspected in patients with lupus anticoagulant who develop nephrotic-range proteinuria. Progression to end-stage renal disease can occur, and a thrombosis may form in the vascular access and the renal allografts. Hypertension is common. Treatment entails lifelong anticoagulation. Glucocorticoids may be beneficial in accelerated hypertension. Immunosuppression and plasma exchange may be helpful for catastrophic episodes of antiphospholipid syndrome but by themselves do not reduce recurrent thrombosis.

HELLP SYNDROME

HELLP (hemolysis, elevated liver enzymes, low platelets) syndrome is a dangerous complication of pregnancy associated with microvascular injury. Occurring in 0.2–0.9% of all pregnancies and in 10–20% of women with severe preeclampsia, this syndrome carries a mortality rate of 7.4–34%. Most commonly developing in the third trimester, 10% of cases occur before week 27 and 30% post-partum. Although a strong association exists between HELLP syndrome and preeclampsia, nearly 20% of cases are not preceded by recognized preeclampsia. Risk factors include abnormal placentation, family history, and elevated levels of fetal mRNA for FLT1 (vascular endothelial growth factor receptor 1) and endoglin. Patients with HELLP syndrome have higher levels of inflammatory markers (C-reactive protein, IL-1Ra, and IL-6) and soluble HLA-DR than do those with preeclampsia alone.

Renal failure occurs in half of patients with HELLP syndrome, although the etiology is not well understood. Limited data suggest that renal failure is the result of both preeclampsia and acute tubular necrosis. Renal histologic findings are those of TMA with endothelial cell swelling and occlusion of the capillary lumens, but luminal thrombi are typically absent. However, thrombi become more common in severe eclampsia and HELLP syndrome. Although renal failure is common, the organ that defines this syndrome is the liver. Subcapsular hepatic hematomas sometimes produce spontaneous rupture of the liver and can be life-threatening. Neurologic complications such as cerebral infarction, cerebral and brainstem hemorrhage, and cerebral edema are other potentially life-threatening complications. Nonfatal complications include placental abruption, permanent vision loss due to Purtscher-like (hemorrhagic and vaso-occlusive vasculopathy) retinopathy, pulmonary edema, bleeding, and fetal demise.

Many features are shared by HELLP syndrome and MAHA. Diagnosis of HELLP syndrome is complicated by the fact that aHUS and TTP also can be triggered by pregnancy. Patients with antiphospholipid syndrome also have an elevated risk of HELLP syndrome. A history of MAHA before pregnancy is of diagnostic value. Serum levels of ADAMTS13 activity are reduced (by 30–60%) in HELLP syndrome but not to the levels seen in TTP (<5%). Determination of the ratio of lactate dehydrogenase to aspartate aminotransferase may be helpful; this ratio is 13:1 in patients with HELLP syndrome and preeclampsia as opposed to 29:1 in patients without preeclampsia. Other markers, such as antithrombin III (decreased in HELLP syndrome but not in TTP) and D-dimer (elevated in HELLP syndrome but not in TTP), may also be useful. HELLP syndrome usually resolves spontaneously after delivery, although a small percentage of HELLP cases occur post-partum. Glucocorticoids may decrease inflammatory markers, although two randomized controlled trials failed to show much benefit. Plasma exchange should be considered if hemolysis is refractory to glucocorticoids and/or delivery, especially if TTP has not been ruled out.

SICKLE CELL NEPHROPATHY

Renal complications in sickle cell disease result from occlusion of the vasa recta in the renal medulla. The low partial pressure of oxygen and high osmolarity predispose to hemoglobin S polymerization and erythrocyte sickling. Sequelae include hyposthenuria, hematuria, and papillary necrosis (which can also occur in sickle trait). The kidney responds by increases in blood flow and glomerular filtration rate mediated by prostaglandins. This dependence on prostaglandins may explain the greater reduction of glomerular filtration rate by nonsteroidal anti-inflammatory drugs in these patients than in others. The glomeruli are typically enlarged. Intracapillary fragmentation and phagocytosis of sickled erythrocytes are thought to be responsible for the membranoproliferative glomerulonephritis-like lesion, and focal segmental glomerulosclerosis is seen in more advanced cases. Proteinuria is present in 20–30%, and nephrotic-range proteinuria is associated with progression to renal failure. ACE inhibitors reduce proteinuria, although data are lacking on prevention of renal failure. Patients with sickle cell disease are also more prone to acute renal failure. The cause is thought to reflect microvascular occlusion associated with nontraumatic rhabdomyolysis, high fever, infection, and

1866 generalized sickling. Chronic kidney disease is present in 12–20% of patients. Despite the frequency of renal disease, hypertension is uncommon in patients with sickle cell disease.

RENAL VEIN THROMBOSIS

Renal vein thrombosis either can present with flank pain, tenderness, hematuria, rapid decline in renal function, and proteinuria or can be silent. Occasionally, renal vein thrombosis is identified during a workup for pulmonary embolism. The left renal vein is more commonly involved, and two-thirds of cases are bilateral. Etiologies can be divided into three broad categories: endothelial damage, venous stasis, and hypercoagulability. Homocystinuria, endovascular intervention, and surgery can produce vascular endothelial damage. Dehydration, which is more common among male patients, is a common cause of stasis in the pediatric population. Stasis also can result from compression and kinking of the renal veins from retroperitoneal processes such as retroperitoneal fibrosis and abdominal neoplasms. Thrombosis can occur throughout the renal circulation, including the renal veins, with antiphospholipid antibody syndrome. Renal vein thrombosis can also be secondary to nephrotic syndrome, particularly membranous nephropathy. Other hypercoagulable states less commonly associated with renal vein thrombosis include proteins C and S, antithrombin deficiency, factor V Leiden, disseminated malignancy, and oral contraceptives. Severe nephrotic syndrome may also predispose patients to renal vein thrombosis.

Diagnostic screening can be performed with Doppler ultrasonography, which is more sensitive than ultrasonography alone. CT angiography is nearly 100% sensitive. Magnetic resonance angiography is another option but is more expensive. Treatment for renal vein thrombosis consists of anticoagulation and therapy for the underlying cause. Endovascular thrombolysis may be considered in severe cases. Occasionally, nephrectomy may be undertaken for life-threatening complications. Vena caval filters are often used to prevent migration of thrombi.

342 Nephrolithiasis

Gary C. Curhan


Nephrolithiasis, or kidney stone disease, is a common, painful, and costly condition. Each year, billions of dollars are spent on nephrolithiasis-related activity, with the majority of expenditures on surgical treatment of existing stones. While a stone may form due to crystallization of lithogenic factors in the upper urinary tract, it can subsequently move into the ureter and cause renal colic. Although nephrolithiasis is rarely fatal, patients who have had renal colic report that it is the worst pain they have ever experienced. The evidence on which to base clinical recommendations is not as strong as desired; nonetheless, most experts agree that the recurrence of most, if not all, types of stones can be prevented with careful evaluation and targeted recommendations. Preventive treatment may be lifelong; therefore, an in-depth understanding of this condition must inform the implementation of tailored interventions that are most appropriate for and acceptable to the patient.

There are various types of kidney stones. It is clinically important to identify the stone type, which informs prognosis and selection of the optimal preventive regimen. Calcium oxalate stones are most common (~75%); next, in order, are calcium phosphate (~15%), uric acid (~8%), struvite (~1%), and cystine (<1%) stones. Many stones are a mixture of crystal types (e.g., calcium oxalate and calcium phosphate) and also contain protein in the stone matrix. Rarely, stones are composed of medications, such as acyclovir, indinavir, and triamterene.

Infectious stones, if not appropriately treated, can have devastating consequences and lead to end-stage renal disease. Consideration

should be given to teaching practitioners strategies to prevent stone recurrence and its related morbidity.

EPIDEMIOLOGY

 Nephrolithiasis is a global disease. Data suggest an increasing prevalence, likely due to Westernization of lifestyle habits (e.g., dietary changes, increasing body mass index). National Health and Nutrition Examination Survey data for 2007–2010 indicate that up to 19% of men and 9% of women will develop at least one stone during their lifetime. The prevalence is ~50% lower among black individuals than among whites. The incidence of nephrolithiasis (i.e., the rate at which previously unaffected individuals develop their first stone) also varies by age, sex, and race. Among white men, the peak annual incidence is ~3.5 cases/1000 at age 40 and declines to ~2 cases/1000 by age 70. Among white women in their thirties, the annual incidence is ~2.5 cases/1000; the figure decreases to ~1.5/1000 at age 50 and beyond. In addition to the medical costs associated with nephrolithiasis, this condition also has a substantial economic impact, as those affected are often of working age. Once an individual has had a stone, the prevention of a recurrence is essential. Published recurrence rates vary by the definitions and diagnostic methods used. Some reports have relied on symptomatic events, while others have been based on imaging. Most experts agree that radiographic evidence of a second stone should be considered to represent a recurrence, even if the stone has not yet caused symptoms.

ASSOCIATED MEDICAL CONDITIONS

Nephrolithiasis is a systemic disorder. Several conditions predispose to stone formation, including gastrointestinal malabsorption (e.g., Crohn's disease, gastric bypass surgery), primary hyperparathyroidism, obesity, type 2 diabetes mellitus, and distal renal tubular acidosis. A number of other medical conditions are more likely to be present in individuals with a history of nephrolithiasis, including hypertension, gout, cholelithiasis, reduced bone mineral density, and chronic kidney disease.

Individuals with medullary sponge kidney (MSK), a condition designated by an anatomic description, often have metabolic abnormalities, such as higher levels of urine calcium and lower levels of urine citrate, and are more likely to form calcium phosphate stones. As intravenous urography is now rarely used, the diagnosis of MSK has become less frequent. Fortunately, the diagnosis of MSK does not change either the evaluation or the treatment recommendations; thus, it is not essential in pursuing the diagnosis of nephrolithiasis.

Although nephrolithiasis does not directly cause upper urinary tract infections (UTIs), a UTI in the setting of an obstructing stone is a urologic emergency (“pus under pressure”) and requires urgent intervention to reestablish drainage.

PATHOGENESIS

In the consideration of the processes involved in crystal formation, it is helpful to view urine as a complex solution. A clinically useful concept is *supersaturation* (the point at which the concentration product exceeds the solubility product). However, even though the urine in most individuals is supersaturated with respect to one or more types of crystals, the presence of inhibitors of crystallization prevents the majority of the population from continuously forming stones. The most clinically important inhibitor of calcium-containing stones is urine citrate. While supersaturation is a calculated value (rather than being directly measured) and does not perfectly predict stone formation, it is a useful guide as it integrates the multiple factors that are measured in a 24-h urine collection.

Recent studies have changed the paradigm for the site of initiation of stone formation. Renal biopsies of stone formers have revealed calcium phosphate in the renal interstitium. It is hypothesized that this calcium phosphate extends down to the papilla and erodes through the papillary epithelium, where it provides a site for deposition of calcium oxalate and calcium phosphate crystals. The majority of calcium oxalate stones grow on calcium phosphate at the tip of the renal papilla (*Randall's plaque*). Thus, the process of stone formation may begin

years before a clinically detectable stone is identified. The processes involved in interstitial deposition are under active investigation.

RISK FACTORS

Risk factors for nephrolithiasis can be categorized as dietary, nondietary, or urinary. These risk factors vary by stone type and by clinical characteristics.

Dietary Risk Factors Patients who develop stones often change their diet; therefore, studies that retrospectively assess diet may be hampered by recall bias. Some studies have examined the relation between diet and changes in the lithogenic composition of the urine, often using calculated supersaturation. However, the composition of the urine does not perfectly predict risk, and not all components that modify risk are included in the calculation of supersaturation. Thus, dietary associations are best investigated by prospective studies that examine actual stone formation as the outcome. Dietary factors that are associated with an increased risk of nephrolithiasis include animal protein, oxalate, sodium, sucrose, and fructose. Dietary factors associated with a lower risk include calcium, potassium, and phytate.

CALCIUM The role of dietary calcium deserves special attention. Although in the past dietary calcium had been suspected of increasing the risk of stone disease, several prospective observational studies and a randomized controlled trial have demonstrated that higher dietary calcium intake is related to a *lower* risk of stone formation. The reduction in risk associated with higher calcium intake may be due to a reduction in intestinal absorption of dietary oxalate that results in lower urine oxalate. Low calcium intake is contraindicated as it increases the risk of stone formation and may contribute to lower bone density in stone formers.

Despite similar bioavailability, supplemental calcium may increase the risk of stone formation. The discrepancy between the risks from dietary calcium and calcium supplements may be due to the timing of supplemental calcium intake or to higher total calcium consumption leading to higher urinary calcium excretion.

OXALATE Urinary oxalate is derived from both endogenous production and absorption of dietary oxalate. Owing to its low and often variable bioavailability, much of the oxalate in food may not be readily absorbed. However, absorption may be higher in stone formers. Although observational studies demonstrate that dietary oxalate is only a weak risk factor for stone formation, urinary oxalate is a strong risk factor for calcium oxalate stone formation, and efforts to avoid high oxalate intake should thus be beneficial.

OTHER NUTRIENTS Several other nutrients have been studied and implicated in stone formation. Higher intake of animal protein may lead to increased excretion of calcium and uric acid as well as to decreased urinary excretion of citrate, all of which increase the risk of stone formation. Higher sodium and sucrose intake increases calcium excretion independent of calcium intake. Higher potassium intake decreases calcium excretion, and many potassium-rich foods increase urinary citrate excretion due to their alkali content. Other dietary factors that have been inconsistently associated with lower stone risk include magnesium and phytate.

Vitamin C supplements are associated with an increased risk of calcium oxalate stone formation, possibly because of raised levels of oxalate in urine. Thus, calcium oxalate stone formers should be advised to avoid vitamin C supplements. Although high doses of supplemental vitamin B₆ may be beneficial in selected patients with type 1 primary hyperoxaluria, the benefit of supplemental vitamin B₆ in other patients is uncertain.

FLUIDS AND BEVERAGES The risk of stone formation increases as urine volume decreases. When the urine output is less than 1 L/d, the risk of stone formation more than doubles. Fluid intake is the main determinant of urine volume, and the importance of fluid intake in preventing stone formation has been demonstrated in observational studies and in a randomized controlled trial. Observational studies have found that coffee, tea, beer, and wine are associated with a reduced risk of stone

formation. Sugar-sweetened carbonated beverage consumption may increase risk.

Nondietary Risk Factors Age, race, body size, and environment are important risk factors for nephrolithiasis. The incidence of stone disease is highest in middle-aged white men, but stones can form in infants as well as in the elderly. There is geographic variability, with the highest prevalence in the southeastern United States. Weight gain increases the risk of stone formation, and the increasing prevalence of nephrolithiasis in the United States may be due in part to the increasing prevalence of obesity. Environmental and occupational influences that may lead to lower urine volume, such as working in a hot environment or lack of ready access to water or a bathroom, are important considerations.

Urinary Risk Factors • URINE VOLUME As mentioned above, lower urine volume results in increased concentrations of lithogenic factors and is a common and readily modifiable risk factor. A randomized trial has demonstrated the effectiveness of elevated fluid intake in increasing urine volume and reducing the risk of stone recurrence.


URINE CALCIUM Higher urine calcium excretion increases the likelihood of formation of calcium oxalate and calcium phosphate stones. While the term *hypercalciuria* is often used, there is no widely accepted cutoff that distinguishes between normal and abnormal urine calcium excretion. In fact, the relation between urine calcium and stone risk appears to be continuous; thus the use of an arbitrary threshold should be avoided. Levels of urine calcium excretion are higher in individuals with a history of nephrolithiasis; however, the mechanisms remain poorly understood. Greater gastrointestinal calcium absorption is one important contributor, and greater bone turnover (with a resultant reduction in bone mineral density) may be another. Primary renal calcium loss, with lower serum calcium concentrations and elevated serum levels of parathyroid hormone (PTH) (and a normal 25-hydroxy vitamin D level), is rare.

URINE OXALATE Higher urine oxalate excretion increases the likelihood of calcium oxalate stone formation. As for urine calcium, no definition for “abnormal” urine oxalate excretion is widely accepted. Given that the relation between urine oxalate and stone risk is continuous, simple dichotomization of urine oxalate excretion is not helpful in assessing risk. The two sources of urine oxalate are endogenous generation and dietary intake. Dietary oxalate is the major contributor and also the source that can be modified. Notably, higher dietary calcium intake reduces gastrointestinal oxalate absorption and thereby reduces urine oxalate.

URINE CITRATE Urine citrate is a natural inhibitor of calcium-containing stones; thus, lower urine citrate excretion increases the risk of stone formation. Citrate reabsorption is influenced by the intracellular pH of proximal tubular cells. Metabolic acidosis will lead to a reduction in citrate excretion by increasing reabsorption of filtered citrate. However, a notable proportion of patients have lower urine citrate for reasons that remain unclear.

URINE URIC ACID Higher urine levels of uric acid—a risk factor for uric acid stone formation—are found in individuals with excess purine consumption and rare genetic conditions that lead to overproduction of uric acid. This characteristic does not appear to be associated with the risk of calcium oxalate stone formation.

URINE pH Urine pH influences the solubility of some crystal types. Uric acid stones form only when the urine pH is consistently ≤ 5.5 or lower, whereas calcium phosphate stones are more likely to form when the urine pH is ≥ 6.5 or higher. Cystine is more soluble at higher urine pH. Calcium oxalate stones are not influenced by urine pH.

 **Genetic Risk Factors** The risk of nephrolithiasis is more than twofold greater in individuals with a family history of stone disease. This association is likely due to a combination of genetic predisposition and similar environmental exposures. While a number of monogenic disorders cause nephrolithiasis, the genetic contributors to common forms of stone disease remain to be determined.

The two most common and well-characterized rare monogenic disorders that lead to stone formation are primary hyperoxaluria and cystinuria. *Primary hyperoxaluria* is an autosomal recessive disorder that causes excessive endogenous oxalate generation by the liver, with consequent calcium oxalate stone formation and crystal deposition in organs. Intraparenchymal calcium oxalate deposition in the kidney can eventually lead to renal failure. *Cystinuria* is an autosomal recessive disorder that causes abnormal reabsorption of filtered dibasic amino acids. The excessive urinary excretion of cystine, which is poorly soluble, leads to cystine stone formation. Cystine stones are visible on plain radiographs and often manifest as staghorn calculi or multiple bilateral stones. Repeat episodes of obstruction and instrumentation can cause chronic renal impairment.

APPROACH TO THE PATIENT: Nephrolithiasis

At present, there are no widely accepted, evidence-based guidelines for the evaluation and treatment of nephrolithiasis. However, there are standard approaches to patients with acute and chronic presentations that can reasonably guide the clinical evaluation.

It typically requires weeks to months (and often much longer) for a kidney stone to grow to a clinically detectable size. Although the passage of a stone is a dramatic event, stone formation and growth are characteristically clinically silent. A stone can remain asymptomatic in the kidney for years or even decades before signs (e.g., hematuria) or symptoms (e.g., pain) become apparent. Thus, it is important to remember that the onset of symptoms, typically attributable to a stone moving into the ureter, does not provide insight into when the stone actually formed. The factors that induce stone movement are unknown.

Clinical Presentation and Differential Diagnosis There are two common presentations for individuals with an acute stone event: renal colic and painless gross hematuria. *Renal colic* is a misnomer because pain typically does not subside completely; rather, it varies in intensity. When a stone moves into the ureter, the discomfort often begins with a sudden onset of unilateral flank pain. The intensity of the pain can increase rapidly, and there are no alleviating factors. This pain, which is accompanied often by nausea and occasionally by vomiting, may radiate, depending on the location of the stone. If the stone lodges in the upper part of the ureter, pain may radiate anteriorly; if the stone is in the lower part of the ureter, pain can radiate to the ipsilateral testicle in men or

the ipsilateral labium in women. Occasionally, a patient has gross hematuria without pain.

Other diagnoses may be confused with acute renal colic. If the stone is lodged at the right ureteral pelvic junction, symptoms may mimic those of acute cholecystitis. If the stone blocks the ureter as it crosses over the right pelvic brim, symptoms may mimic acute appendicitis, whereas blockage at the left pelvic brim may be confused with acute diverticulitis. If the stone lodges in the ureter at the ureterovesical junction, the patient may experience urinary urgency and frequency. In female patients, the latter symptoms may lead to an incorrect diagnosis of bacterial cystitis; the urine will contain red and white blood cells, but the urine culture will be negative. An obstructing stone with proximal infection may present as acute pyelonephritis. A UTI in the setting of ureteral obstruction is a medical emergency that requires immediate restoration of drainage by placement of either a ureteral stent or a percutaneous nephrostomy tube. Other conditions to consider in the differential diagnosis include muscular or skeletal pain, *herpes zoster*, duodenal ulcer, abdominal aortic aneurysm, gynecologic conditions, ureteral stricture, and ureteral obstruction by materials other than a stone, such as a blood clot or sloughed papilla. Extraluminal processes can lead to ureteral compression and obstruction; however, because of the gradual onset, these conditions do not typically present with renal colic.

Diagnosis and Intervention Serum chemistry findings are typically normal, but the white blood cell count may be elevated. Examination of the urine sediment will usually reveal red and white blood cells and occasionally crystals (Fig. 342-1). The absence of hematuria does not exclude a stone, particularly when urine flow is completely obstructed by a stone.

The diagnosis is often made on the basis of the history, physical examination, and urinalysis. Thus, it may not be necessary to wait for radiographic confirmation before treating the symptoms. The diagnosis is confirmed by an appropriate imaging study—preferably helical CT, which is highly sensitive, allows visualization of uric acid stones (traditionally considered “radiolucent”), and is able to avoid radiocontrast (Fig. 342-2). Helical CT detects stones as small as 1 mm that may be missed by other imaging modalities. Typically, helical CT reveals a ureteral stone or evidence of recent passage (e.g., perinephric stranding or hydronephrosis), whereas a plain abdominal radiograph (kidney/ureter/bladder, or KUB) can miss a stone in the ureter or kidney, even if it is radiopaque, and does not provide information on obstruction. Abdominal ultrasound offers the advantage of avoiding radiation and provides

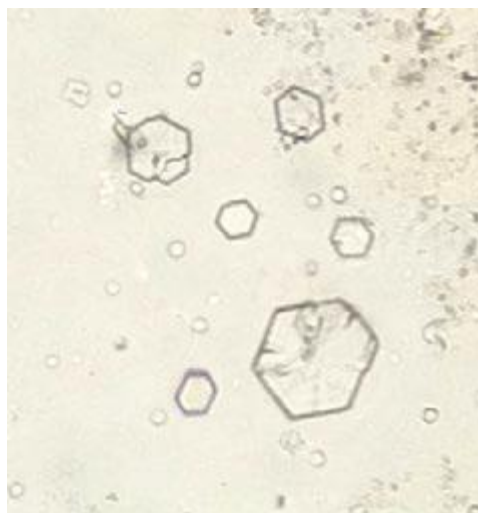
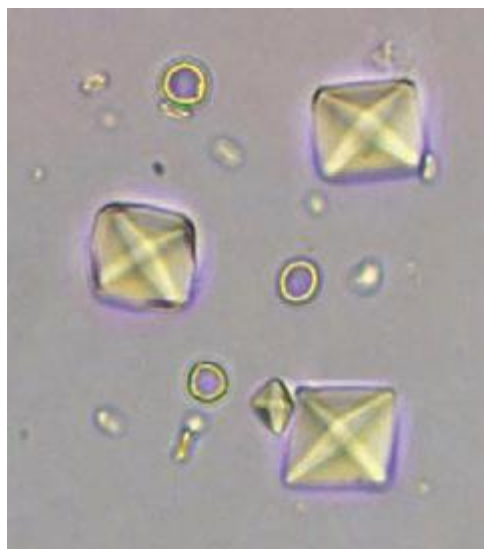


FIGURE 342-1 Urine sediment from a patient with calcium oxalate stones (*left*) and a patient with cystine stones (*right*). Calcium oxalate dihydrate crystals are bipyramidally shaped, and cystine crystals are hexagonal. (*Left panel image courtesy of Dr. John Lieske, Mayo Clinic.*)



FIGURE 342-2 Coronal noncontrast CT image from a patient who presented with left-sided renal colic. An obstructing calculus, present in the distal left ureter at the level of S1, measures 10 mm in maximal dimension. There is severe left hydronephrosis and associated left perinephric fat stranding. In addition, there is a non-obstructing 6-mm left renal calculus in the interpolar region. (Image courtesy of Dr. Stuart Silverman, Brigham and Women's Hospital.)

information on hydronephrosis, but it is not as sensitive as CT and images only the kidney and possibly the proximal segment of the ureter; thus most ureteral stones are not detectable by ultrasound.

Many patients who experience their first episode of colic seek emergent medical care. Randomized trials have demonstrated that parenterally administered nonsteroidal anti-inflammatory drugs (such as ketorolac) are just as effective as opioids in relieving symptoms and have fewer side effects. Excessive fluid administration has not been shown to be beneficial; therefore, the goal should be to maintain euolemia. If the pain can be adequately controlled and the patient is able to take fluids orally, hospitalization can be avoided. Use of an alpha-blocker may increase the rate of spontaneous stone passage.

Urologic intervention should be postponed unless there is evidence of UTI, a low probability of spontaneous stone passage (e.g., a stone measuring ≥ 6 mm or an anatomic abnormality), or intractable pain. A ureteral stent may be placed cystoscopically, but this procedure typically requires general anesthesia, and the stent can be quite uncomfortable, may cause gross hematuria, and may increase the risk of UTI.

If an intervention is indicated, the selection of the most appropriate intervention is determined by the size, location, and composition of the stone; the urinary tract anatomy; and the experience of the urologist. Extracorporeal shockwave lithotripsy, the least invasive option, uses shock waves generated outside the body to fragment the stone. An endourologic approach can remove a stone by basket extraction or laser fragmentation. For large upper-tract stones, percutaneous nephrostolithotomy has the highest likelihood of rendering the patient stone-free. Advances in urologic approaches and instruments have nearly eliminated the need for open surgical procedures such as ureterolithotomy or pyelolithotomy.

Evaluation for Stone Prevention More than half of first-time stone formers will have a recurrence within 10 years. A careful evaluation is indicated to identify predisposing factors, which can then be modified to reduce the risk of new stone formation. It is appropriate to proceed with an evaluation even after the first stone because recurrences are common and are usually preventable with inexpensive lifestyle modifications or other treatments.

History A detailed history, obtained from the patient and from a thorough review of medical records, should include the number and frequency of episodes (distinguishing stone passage from stone formation) and previous imaging studies, interventions, evaluations, and treatments. Inquiries about the patient's medical history should cover UTIs, bariatric surgery, gout, hypertension, and diabetes mellitus. A family history of stone disease may reveal a genetic predisposition. A complete list of current prescription and over-the-counter medications as well as vitamin and mineral supplements is essential. The review of systems should focus on identifying possible etiologic factors related to low urine volume (e.g., high insensible losses; low fluid intake) and gastrointestinal malabsorption as well as on ascertaining how frequently the patient voids during the day and overnight.

A large body of compelling evidence has demonstrated the important role of diet in stone disease. Thus, the dietary history should encompass information on usual dietary habits (meals and snacks), calcium intake, consumption of high-oxalate foods (spinach, rhubarb, potatoes), and fluid intake (including specific beverages typically consumed).

Physical Examination The physical examination should assess weight, blood pressure, costovertebral angle tenderness, and lower-extremity edema as well as signs of other systemic conditions such as primary hyperparathyroidism and gout.

Laboratory Evaluation If not recently measured, the following serum levels should be determined: electrolytes (to uncover hypokalemia or renal tubular acidosis), creatinine, calcium, and uric acid. The PTH level should be measured if indicated by high-normal or elevated serum and urine calcium concentrations. Often, 25-hydroxy vitamin D is measured in concert with PTH to investigate the possible role of secondarily elevated PTH levels in the setting of vitamin D insufficiency.

The urinalysis, including examination of the sediment, can provide useful information. In individuals with asymptomatic residual renal stones, red and white blood cells are frequently present in urine. If there is concern about the possibility of an infection, a urine culture should be performed. The sediment may also reveal crystals (Fig. 342-1), which may help identify the stone type and also provide prognostic information, as crystalluria is a strong risk factor for new stone formation.

The results from 24-h urine collections serve as the cornerstone on which therapeutic recommendations are based. Recommendations on lifestyle modification should be deferred until urine collection is complete. As a baseline assessment, patients should collect at least two 24-h urine samples while consuming their usual diet and usual volume of fluid. The following factors should be measured: total volume, calcium, oxalate, citrate, uric acid, sodium, potassium, phosphorus, pH, and creatinine. When available, the calculated supersaturation is also informative. There is substantial day-to-day variability in the 24-h excretion of many relevant factors; therefore, obtaining values from two collections is important before committing a patient to long-term lifestyle changes or medication. The interpretation of the 24-h urine results should take into account that the collections are usually performed on a weekend day when the patient is staying at home; an individual's habits may differ dramatically (beneficially or detrimentally) at work or outside the home. Specialized testing, such as calcium loading or restriction, is not recommended as it does not influence clinical recommendations.

Stone composition analysis is essential if a stone or fragment is available; patients should be encouraged to retrieve passed stones. The stone type cannot be determined with certainty from 24-h urine results.

Imaging The "gold standard" diagnostic test is helical CT without contrast. If not already performed during an acute episode, a CT should be considered to definitively establish the baseline stone burden. A suboptimal imaging study may not detect a residual stone that, if subsequently passed, would be mistaken for a new

stone. In this instance, the preventive medical regimen might be unnecessarily changed as the result of a preexisting stone.

Recommendations for follow-up imaging should be tailored to the individual patient. While CT provides the best information, the radiation dose is substantially higher than from other modalities; therefore, CT should be performed only if the results will lead to a change in clinical recommendations. Although they are less sensitive, renal ultrasound or a KUB exam are typically used to minimize radiation exposure, with recognition of the limitations.

Prevention of New Stone Formation Recommendations for preventing stone formation depend on the stone type and the results of metabolic evaluation. After remediable secondary causes of stone formation (e.g., primary hyperparathyroidism) are excluded, the focus should turn to modification of the urine composition to reduce the risk of new stone formation. The urinary constituents are continuous variables, and the associated risk is continuous; thus, there is no definitive threshold. Dichotomization into “normal” and “abnormal” can be misleading and should be avoided.

For all stone types, consistently diluted urine reduces the likelihood of crystal formation. The urine volume should be at least 2 L/d. Because of differences in insensible fluid losses and fluid intake from food sources, the required total fluid intake will vary from person to person. Rather than specify how much to drink, it is more helpful to educate patients about how much *more* they need to drink in light of their 24-h urine volume. For example, if the daily urine volume is 1.5 L, then the patient should be advised to drink at least 0.5 L more per day in order to increase the urine volume to the goal of 2 L/day.

RECOMMENDATIONS FOR SPECIFIC STONE TYPES

Calcium Oxalate Risk factors for calcium oxalate stones include higher urine calcium, higher urine oxalate, and lower urine citrate. This stone type is insensitive to pH in the physiologic range.

Individuals with higher urine calcium excretion tend to absorb a higher percentage of ingested calcium. Nevertheless, dietary calcium restriction is not beneficial and, in fact, is likely to be harmful (see “Dietary Risk Factors,” above). In a randomized trial in men with high urine calcium and recurrent calcium oxalate stones, a diet containing 1200 mg of calcium and a low intake of sodium and animal protein significantly reduced subsequent stone formation from that with a low-calcium diet (400 mg/d). Excessive calcium intake (>1200 mg/d) should be avoided.

A thiazide diuretic, in doses higher than those used to treat hypertension, can substantially lower urine calcium excretion. Several randomized controlled trials have demonstrated that thiazide diuretics can reduce calcium oxalate stone recurrence by ~50%. When a thiazide is prescribed, dietary sodium restriction is essential to obtain the desired reduction in urinary calcium excretion. While bisphosphonates may reduce urine calcium excretion in some individuals, there are no data on whether this class of medication can reduce stone formation; therefore, bisphosphonates cannot be recommended solely for stone prevention at present.

A reduction in urine oxalate will in turn reduce the supersaturation of calcium oxalate. In patients with the common form of nephrolithiasis, avoiding high-dose vitamin C supplements is the only known strategy that reduces endogenous oxalate production.

Oxalate is a metabolic end product; therefore, any dietary oxalate that is absorbed will be excreted in the urine. Reducing absorption of exogenous oxalate involves two approaches. First, the avoidance of foods that contain high amounts of oxalate, such as spinach, rhubarb, and potatoes, is prudent. However, extreme oxalate restriction has not been demonstrated to reduce stone recurrence and could be harmful to overall health, given other health benefits of many foods that are erroneously considered to be high in oxalate. Controversy exists regarding the most clinically relevant measure of the oxalate content of foods (e.g., bioavailability). Notably, the absorption of oxalate is reduced by higher calcium intake; therefore, individuals with higher-than-desired urinary oxalate should be counseled to consume adequate calcium. Oxalate absorption can be influenced by the

intestinal microbiota, depending on the presence of oxalate-degrading bacteria. Currently, however, there are no available therapies to alter the microbiota that beneficially affect urinary oxalate excretion over the long term.

Citrate is a natural inhibitor of calcium oxalate and calcium phosphate stones. Higher-level consumption of foods rich in alkali (i.e., fruits and vegetables) can increase urine citrate. For patients with lower urine citrate in whom dietary modification does not adequately increase urine citrate, the addition of supplemental alkali (typically potassium citrate) will lead to an increase in urinary citrate excretion. Sodium salts, such as sodium bicarbonate, while successful in raising urine citrate, are typically avoided due to the adverse effects of sodium on urine calcium excretion.

Past reports suggested that higher levels of urine uric acid may increase the risk of calcium oxalate stones, but more recent studies do not support this association. However, allopurinol reduced stone recurrence in one randomized controlled trial in patients with calcium oxalate stones and high urine uric acid levels. The lack of association between urine uric acid level and calcium oxalate stones suggests that a different mechanism underlies the observed beneficial effect of allopurinol.

Additional dietary modifications may be beneficial in reducing stone recurrence. Restriction of nondairy animal protein (e.g., meat, chicken, seafood) is a reasonable approach and may result in higher excretion of citrate and lower excretion of calcium. In addition, reducing sodium intake to <2.5 g/d may decrease urinary excretion of calcium. Sucrose and fructose intake should be minimized.

For adherence to a dietary pattern that is more manageable for patients than manipulating individual nutrients, the DASH (Dietary Approaches to Stop Hypertension) diet provides an appropriate and readily available option. Randomized trials have conclusively shown the DASH diet to reduce blood pressure. At present, only data from observational studies are available, but these demonstrate a strong and consistent inverse association between the DASH diet and risk of stone formation.

Calcium Phosphate Calcium phosphate stones share risk factors with calcium oxalate stones, including higher concentrations of urine calcium and lower concentrations of urine citrate, but additional factors deserve attention. Higher urine phosphate levels and higher urine pH (typically ≥ 6.5) are associated with an increased likelihood of calcium phosphate stone formation. Calcium phosphate stones are more common in patients with distal renal tubular acidosis and primary hyperparathyroidism.

There are no randomized trials on which to base preventive recommendations for calcium phosphate stone formers, so the interventions are focused on modification of the recognized risk factors. Thiazide diuretics (with sodium restriction) may be used to reduce urine calcium, as described above for calcium oxalate stones. In patients with low urine citrate levels, alkali supplements (e.g., potassium citrate) may be used to increase these concentrations. However, the urine pH of these patients should be monitored carefully because supplemental alkali can raise urine pH, thereby potentially increasing the risk of stone formation. Reduction of dietary phosphate may be beneficial by reducing urine phosphate excretion.

Uric Acid The two main risk factors for uric acid stones are persistently low urine pH and higher uric acid excretion. Urine pH is the predominant influence on uric acid solubility; therefore, the mainstay of prevention of uric acid stone formation entails increasing urine pH. While acidifying the urine is not easily done, alkalinizing the urine can be readily achieved by increasing the intake of foods rich in alkali (e.g., fruits and vegetables) and reducing the intake of foods that produce acid (e.g., animal flesh). If necessary, supplementation with bicarbonate or citrate salts (preferably potassium citrate) can be used to reach the recommended pH goal of 6 to 7 throughout the day and night.

Urine uric acid excretion is determined by uric acid generation. Uric acid is the end product of purine metabolism; thus reduced consumption of purine-containing foods can lower urine uric acid excretion. It is noteworthy that the serum uric acid level is dependent on the fractional excretion of uric acid and therefore does not provide

information on urine uric acid excretion. For example, an individual with high uric acid generation and concurrent high fractional excretion of uric acid will have high urine uric acid excretion with a normal (or even low) serum uric acid level. If alkalization of the urine alone is not successful and if dietary modifications do not reduce urine uric acid sufficiently, then the use of a xanthine oxidase inhibitor, such as allopurinol or febuxostat, can reduce urine uric acid excretion by 40–50%.

Cystine Cystine excretion is not easily modified. Long-term dietary cystine restriction is not feasible and is unlikely to be successful; thus the focus for cystine stone prevention is on increasing cystine solubility. This goal may be achieved by treatment with medication that covalently binds to cystine (tiopronin and penicillamine) and a medication that raises urine pH. Tiopronin is the preferred choice due to its better adverse event profile. The preferred alkalizing agent is potassium citrate as sodium salts may increase cystine excretion. As with all stone types, and especially in patients with cystinuria, maintaining a high urine volume is an essential component of the preventive regimen.

Struvite Struvite stones, also known as *infection stones* or *triple-phosphate stones*, form only when the upper urinary tract is infected with urease-producing bacteria such as *Proteus mirabilis*, *Klebsiella pneumoniae*, or *Providencia* species. Urease produced by these bacteria hydrolyzes urea and may elevate the urine pH to a supraphysiologic level (>8.0). Struvite stones may grow quickly and fill the renal pelvis (*staghorn calculi*).

Struvite stones require complete removal by a urologist. New stone formation can be avoided by the prevention of UTIs. In patients with recurrent upper UTIs (e.g., some individuals with surgically altered urinary drainage or spinal cord injury), the urease inhibitor acetohydroxamic acid can be considered; however, this agent should be used with caution because of potential side effects.

LONG-TERM FOLLOW-UP

In general, the preventive regimens described above do not cure the underlying pathophysiologic process. Thus these recommendations typically need to be followed for the patient's lifetime, and it is essential to tailor recommendations in a way that is acceptable to the patient. Because the memory of the acute stone event fades and patients often return to old habits (e.g., insufficient fluid intake), long-term follow-up is important to ensure that the preventive regimen has been implemented and has resulted in the desired reduction in the risk of new stone formation.

Follow-up imaging should be planned thoughtfully. Many patients with recurrent episodes of renal colic that lead to emergency room visits often undergo repeat CT studies. While CT does provide the best information, the radiation dose is substantially higher than that with plain abdominal radiography (KUB). Small stones may be missed by KUB, and ultrasound has a limited ability to determine size and number of stones. Minimizing radiation exposure should be a goal of the long-term follow-up plan and must be balanced against the gain in diagnostic information.

stone formation. Early diagnosis and prompt therapy are, therefore, essential to minimize the otherwise devastating effects of obstruction on kidney structure and function.

ETIOLOGY

Obstruction to urine flow can result from *intrinsic* or *extrinsic mechanical blockade* as well as from *functional defects* not associated with fixed occlusion of the urinary drainage system. Mechanical obstruction can occur at any level of the urinary tract, from the renal calyces to the external urethral meatus. Normal points of narrowing, such as the ureteropelvic and ureterovesical junctions, bladder neck, and urethral meatus, are common sites of obstruction. When obstruction is above the level of the bladder, unilateral dilatation of the ureter (*hydroureter*) and renal pyelocalyceal system (*hydronephrosis*) occurs; lesions at or below the level of the bladder cause bilateral involvement.

Common forms of obstruction are listed in [Table 343-1](#). Childhood causes include *congenital malformations*, such as narrowing of the ureteropelvic junction and abnormal insertion of the ureter into the bladder, the most common cause. Vesicoureteral reflux in the absence of urinary tract infection or bladder neck obstruction often resolves with age. Reinsertion of the ureter into the bladder is indicated if reflux is severe and unlikely to improve spontaneously, if renal function deteriorates, or if urinary tract infections recur despite chronic antimicrobial therapy. Vesicoureteral reflux may cause prenatal hydronephrosis and, if severe, can lead to recurrent urinary infections and renal scarring in childhood. Posterior urethral valves are the most common cause of bilateral hydronephrosis in boys. In adults, urinary tract obstruction (UTO) is due mainly to *acquired defects*. Pelvic tumors, calculi, and urethral stricture predominate. Ligation of, or injury to, the ureter during pelvic or colonic surgery can lead to hydronephrosis which, if unilateral, may remain undetected. Obstructive uropathy may also result from extrinsic neoplastic (carcinoma of

TABLE 343-1 COMMON MECHANICAL CAUSES OF URINARY TRACT OBSTRUCTION

Ureter	Bladder Outlet	Urethra
Congenital		
Ureteropelvic junction narrowing or obstruction	Bladder neck obstruction Ureterocele	Posterior urethral valves
Ureterovesical junction narrowing or obstruction and reflux		Anterior urethral valves
Ureterocele		Stricture
Retrocaval ureter		Meatal stenosis Phimosis
Acquired Intrinsic Defects		
Calculi	Benign prostatic hyperplasia	Stricture
Inflammation	Cancer of prostate	Tumor
Infection	Cancer of bladder	Calculi
Trauma	Calculi	Trauma
Sloughed papillae	Diabetic neuropathy	Phimosis
Tumor	Spinal cord disease	
Blood clots	Anticholinergic drugs and α -adrenergic antagonists	
Acquired Extrinsic Defects		
Pregnant uterus	Carcinoma of cervix, colon	Trauma
Retroperitoneal fibrosis		
Aortic aneurysm	Trauma	
Uterine leiomyomata		
Carcinoma of uterus, prostate, bladder, colon, rectum		
Lymphoma		
Pelvic inflammatory disease, endometriosis		
Accidental surgical ligation		

343 Urinary Tract Obstruction

Julian L. Seifter

Obstruction to the flow of urine, with attendant stasis and elevation in urinary tract pressure, impairs renal and urinary conduit functions and is a common cause of acute and chronic kidney disease (obstructive nephropathy). With early relief of obstruction, the defects in function usually disappear completely. However, chronic obstruction may produce permanent loss of renal mass (renal atrophy) and excretory capability, as well as enhanced susceptibility to local infection and

cervix or colon) or inflammatory disorders. Lymphomas and pelvic or colonic neoplasms with retroperitoneal involvement are causes of ureteral obstruction. As many as 50% of men over 40 years old may have lower urinary tract symptoms associated with benign prostatic hypertrophy, but these symptoms may occur without bladder outlet obstruction.

Functional impairment of urine flow occurs when voiding is altered by abnormal pontine or sacral centers of micturition control. It may be asymptomatic or associated with lower urinary tract symptoms such as frequency, urgency, urge and postmicturition incontinence, nocturia, straining to void, slow stream, hesitancy, or a feeling of incomplete emptying. A history should be sought for trauma, back injury, surgery, diabetes, neurologic or psychiatric conditions, and medications. Causes include neurogenic bladder, often with adynamic ureter, and vesicoureteral reflux. Reflux in children may result in severe unilateral or bilateral hydronephrosis and hydronephrosis. Urinary retention may be the consequence of α -adrenergic and anticholinergic agents, as well as opiates. Hydronephrosis in pregnancy is due to relaxational effects of progesterone on smooth muscle of the renal pelvis, as well as ureteral compression by the enlarged uterus.

Diagnostic tools to identify anatomic obstruction include urinary flow measurements and a postvoid residual. Cystourethroscopy and urodynamic studies may be reserved for the symptomatic patient to assess the filling phase (cystometry), pressure-volume relationship of the bladder, bladder compliance, and capacity. Pressure-flow analysis evaluates bladder contractility and bladder outlet resistance during voiding. Bladder obstruction is characterized by high pressures in women, whereas in men, a diagnosis of bladder outlet obstruction is based on flow rate and voiding pressures. A voiding cystourethrogram may be useful in evaluating incomplete emptying and bladder neck and urethral pathology.

CLINICAL FEATURES AND PATHOPHYSIOLOGY

The pathophysiology and clinical features of UTO are summarized in **Table 343-2**. *Pain*, the symptom that most commonly leads to medical attention, is due to distention of the collecting system or renal capsule. Pain severity is influenced more by the rate at which distention develops than by the degree of distention. Acute supravascular obstruction, as from a stone lodged in a ureter (**Chap. 342**), is associated with excruciating pain, known as *renal colic*. This pain often radiates to the

lower abdomen, testes, or labia. By contrast, more insidious causes of obstruction, such as chronic narrowing of the ureteropelvic junction, may produce little or no pain and yet result in total destruction of the affected kidney. Flank pain that occurs only with micturition is pathognomonic of vesicoureteral reflux.

Obstruction of urine flow results in an increase in hydrostatic pressures proximal to the site of obstruction. It is this buildup of pressure that leads to the accompanying pain, the distention of the collecting system in the kidney, and elevated intratubular pressures that initiate tubular dysfunction. As the increased hydrostatic pressure is expressed in the urinary space of the glomeruli, further filtration decreases or stops completely.

Azotemia develops when overall excretory function is impaired, often in the setting of bladder outlet obstruction, bilateral renal pelvic or ureteric obstruction, or unilateral disease in a patient with a solitary functioning kidney. Complete bilateral obstruction should be suspected when acute renal failure is accompanied by anuria. Any patient with renal failure otherwise unexplained, or with a history of nephrolithiasis, hematuria, diabetes mellitus, prostatic enlargement, pelvic surgery, trauma, or tumor should be evaluated for UTO.

In the acute setting, partial, bilateral obstruction may mimic prerenal azotemia with concentrated urine and sodium retention. However, with more prolonged obstruction, symptoms of *polyuria* and *nocturia* commonly accompany partial UTO and result from diminished renal concentrating ability. Impairment of transcellular salt reabsorption in the proximal tubule, medullary thick ascending limb of Henle, and collecting duct cells is due to downregulation of transport proteins including the Na^+ , K^+ adenosine triphosphatase (ATPase), NaK_2Cl cotransporter (NKCC) in the thick ascending limb, and the epithelial Na^+ channel (ENaC) in collecting duct cells. Consequences include failure to produce urine free of salt (natriuresis) and loss of medullary hypertonicity producing a urinary concentrating defect. In addition to direct effects on renal transport mechanisms, increased prostaglandin E_2 (PGE_2) (due to induction of cyclooxygenase-2 [COX-2]), angiotensin II (with its downregulation of Na^+ transporters), and atrial or B-type natriuretic peptides (ANP or BNP) (due to volume expansion in the azotemic patient) contribute to the decreased salt reabsorption along the nephron.

Dysregulation of aquaporin-2 water channels in the collecting duct contributes to the polyuria. The defect usually does not improve with administration of vasopressin and is therefore a form of acquired nephrogenic diabetes insipidus.

Wide fluctuations in urine output in a patient with azotemia should always raise the possibility of intermittent or partial UTO. If fluid intake is inadequate, severe dehydration and hyponatremia may develop. However, as with other causes of poor renal function, excesses of salt and water intake may result in edema and hyponatremia.

Partial bilateral UTO often results in *acquired distal renal tubular acidosis*, *hyperkalemia*, and *renal salt wasting*. The H^+ -ATPase, situated on the apical membrane of the intercalated cells of the collecting duct, is critical for distal H^+ secretion. The trafficking of intracellular H^+ pumps from the cytoplasm to the cell membrane is disrupted in UTO. The decreased function of the ENaC, in the apical membrane of neighboring collecting duct principal cells, contributes to decreased Na^+ reabsorption (salt-wasting), decreased electronegativity of the tubule lumen, and therefore decreased K^+ secretion via K^+ channels (hyperkalemia) and H^+ secretion via the H^+ -ATPases (distal renal tubular acidosis [RTA]). Proximal tubule ammoniogenesis, important to the elimination of H^+ as NH_4^+ , is impaired. These defects in tubule function are often accompanied by renal tubulointerstitial damage. Azotemia with hyperkalemia and metabolic acidosis should prompt consideration of UTO.

The renal interstitium becomes edematous and infiltrated with mononuclear inflammatory cells early in UTO. Later, interstitial fibrosis and atrophy of the papillae and medulla occur and precede these processes in the cortex. The increase in angiotensin II noted in UTO contributes to the inflammatory response and fibroblast accumulation through mechanisms involving profibrotic cytokines. With time, this process leads to chronic kidney damage.

TABLE 343-2 PATHOPHYSIOLOGY OF BILATERAL URETERAL OBSTRUCTION

Hemodynamic Effects	Tubule Effects	Clinical Features
Acute		
↑ Renal blood flow	↑ Ureteral and tubule pressures	Pain (capsule distention)
↓ GFR	↑ Reabsorption of Na^+ , urea, water	Azotemia
↓ Medullary blood flow		Oliguria or anuria
↑ Vasodilator prostaglandins, nitric oxide		
Chronic		
↓ Renal blood flow	↓ Medullary osmolarity	Azotemia
↓↓ GFR	↓ Concentrating ability	Hypertension
↑ Vasoconstrictor prostaglandins	Structural damage; parenchymal atrophy	AVP-insensitive polyuria
↑ Renin-angiotensin production	↓ Transport functions for Na^+ , K^+ , H^+	Natriuresis
		Hyperkalemic, hyperchloremic acidosis
Release of Obstruction		
Slow ↑ in GFR (variable)	↓ Tubule pressure	Postobstructive diuresis
	↑ Solute load per nephron (urea, NaCl)	Potential for volume depletion and electrolyte imbalance due to losses of Na^+ , K^+ , PO_4^{2-} , Mg^{2+} , and water
	Natriuretic factors present	

Abbreviations: AVP, arginine vasopressin; GFR, glomerular filtration rate.

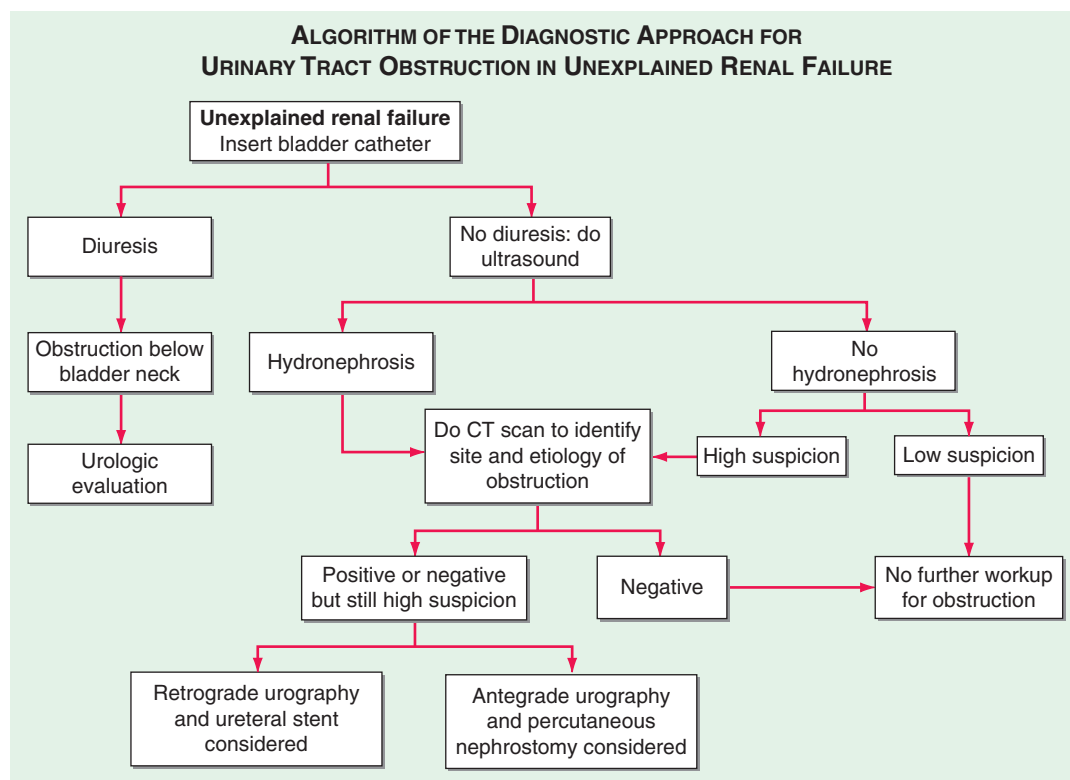


FIGURE 343-1 Diagnostic approach for urinary tract obstruction in unexplained renal failure. CT, computed tomography.

UTO must always be considered in patients with urinary tract infections or urolithiasis. Urinary stasis encourages the growth of organisms. Urea-splitting bacteria are associated with magnesium ammonium phosphate (struvite) calculi. Hypertension is frequent in acute and subacute unilateral obstruction and is usually a consequence of increased release of renin by the involved kidney. Chronic kidney disease from bilateral UTO, often associated with extracellular volume expansion, may result in significant hypertension. Erythrocytosis, an infrequent complication of obstructive uropathy, is secondary to increased erythropoietin production.

DIAGNOSIS

A history of difficulty in voiding, pain, infection, or change in urinary volume is common. Evidence for distention of the kidney or urinary bladder can often be obtained by palpation and percussion of the abdomen. A careful rectal and genital examination may reveal enlargement or nodularity of the prostate, abnormal rectal sphincter tone, or a rectal or pelvic mass.

Urinalysis may reveal hematuria, pyuria, and bacteriuria. The urine sediment is often normal, even when obstruction leads to marked azotemia and extensive structural damage. An abdominal scout film may detect nephrocalcinosis or a radiopaque stone. As indicated in Fig. 343-1, if UTO is suspected, a bladder catheter should be inserted. Abdominal ultrasonography should be performed to evaluate renal and bladder size, as well as pyelocalyceal contour. Ultrasonography is approximately 90% specific and sensitive for detection of hydronephrosis. False-positive results are associated with diuresis, renal cysts, or the presence of an extrarenal pelvis, a normal congenital variant. Congenital ureteropelvic junction (UPJ) obstruction may be mistaken for renal cystic disease. Hydronephrosis may be absent on ultrasound when obstruction is less than 48 h in duration or associated with volume contraction, staghorn calculi, retroperitoneal fibrosis, or infiltrative renal disease. Duplex Doppler ultrasonography may detect an increased resistive index in urinary obstruction.

Recent advances in technology have led to alternatives and have largely replaced the once standard intravenous urogram in the further evaluation of UTO. The high-resolution multidetector row computed

tomography (CT) scan in particular has advantages of visualizing the retroperitoneum, as well as identifying both intrinsic and extrinsic sites of obstruction. Noncontrast CT scans improve visualization of the urinary tract in the patient with renal impairment and are safer for patients at risk for contrast nephropathy. Magnetic resonance urography is a promising technique but, at this time, not superior to the CT scan and carries the risk of certain gadolinium agents in patients with renal insufficiency, i.e., nephrogenic systemic fibrosis. The intravenous urogram may define the site of obstruction and demonstrate dilatation of the calyces, renal pelvis, and ureter above the obstruction. The ureter may be tortuous in chronic obstruction. Radionuclide scans are able to give differential renal function but give less anatomic detail than CT or intravenous urography (IVU).

To facilitate visualization of a suspected lesion in a ureter or renal pelvis, retrograde or antegrade urography should be attempted. These procedures do not carry risk of contrast-induced acute renal failure in patients with renal insufficiency. The retrograde approach involves catheterization of the involved ureter under cystoscopic control, whereas the antegrade technique necessitates percutaneous placement of a catheter into the renal pelvis. Although the antegrade approach may provide immediate decompression of a unilateral obstructing lesion, many urologists initially attempt the retrograde approach unless the catheterization is unsuccessful.

Voiding cystourethrography is of value in the diagnosis of vesico-ureteral reflux and bladder neck and urethral obstructions. Postvoiding films reveal residual urine. Endoscopic visualization by the urologist often permits precise identification of lesions involving the urethra, prostate, bladder, and ureteral orifices.

TREATMENT URINARY TRACT OBSTRUCTION

UTO complicated by infection requires immediate relief of obstruction to prevent development of generalized sepsis and progressive renal damage. Sepsis necessitates prompt urologic intervention. Drainage may be achieved by nephrostomy, ureterostomy, or ureteral, urethral, or suprapubic catheterization. Prolonged antibiotic treatment may be necessary. Chronic or recurrent infections in a

1874 poorly functioning obstructed kidney may necessitate nephrectomy. When infection is not present, surgery is often delayed until acid-base, fluid, and electrolyte status is restored. Nevertheless, the site of obstruction should be ascertained as soon as feasible. Elective relief of obstruction is usually recommended in patients with urinary retention, recurrent urinary tract infections, persistent pain, or progressive loss of renal function. Benign prostatic hypertrophy may be treated medically with α -adrenergic blockers and 5 α -reductase inhibitors. Functional obstruction secondary to neurogenic bladder may be decreased with the combination of frequent voiding and cholinergic drugs.

PROGNOSIS

With relief of obstruction, the prognosis regarding return of renal function depends largely on whether irreversible renal damage has occurred. When obstruction is not relieved, the course will depend mainly on whether the obstruction is complete or incomplete and bilateral or unilateral, as well as whether or not urinary tract infection is also present. Complete obstruction with infection can lead to total destruction of the kidney within days. Partial return of glomerular filtration rate may follow relief of complete obstruction of 1 and 2 weeks' duration, but after 8 weeks of obstruction, recovery is unlikely. In the absence of definitive evidence of irreversibility, every effort should be made to decompress the obstruction in the hope of restoring renal function at least partially. A renal radionuclide scan, performed after a prolonged period of decompression, may be used to predict the reversibility of renal dysfunction.

POSTOBSTRUCTIVE DIURESIS

Relief of bilateral, but not unilateral, complete obstruction commonly results in polyuria, which may be massive. The urine is usually hypotonic and may contain large amounts of sodium chloride, potassium, phosphate, and magnesium. The natriuresis is due in part to the normal correction of extracellular volume expansion, the increase in natriuretic factors accumulated during the period of renal failure, and depressed salt and water reabsorption when urine flow is reestablished. The retained urea is excreted with improved GFR, resulting in an osmotic diuresis which increases the urine volume of electrolyte-free water. In the majority of patients, this diuresis results in the *appropriate* excretion of the excesses of retained salt and water. When extracellular volume and composition return to normal, the diuresis usually abates spontaneously. Occasionally, iatrogenic expansion of extracellular volume is responsible for, or sustains, the diuresis observed in the postobstructive period. Replacement with intravenous fluids in amounts less than urinary losses usually prevents this complication. More aggressive fluid management is required in the setting of hypovolemia, hypotension, or disturbances in serum electrolyte concentrations.

The loss of electrolyte-free water with urea may result in hypernatremia. Serum and urine sodium and osmolal concentrations should guide the use of appropriate intravenous replacement. Often replacement with 0.45% saline is required. Relief of obstruction may be followed by urinary salt and water losses severe enough to provoke profound dehydration and vascular collapse. In these patients, decreased tubule reabsorptive capacity is probably responsible for the marked diuresis. Appropriate therapy in such patients includes intravenous administration of salt-containing solutions to replace sodium and volume deficits.

SECTION 1 DISORDERS OF THE ALIMENTARY TRACT

344 Approach to the Patient with Gastrointestinal Disease

William L. Hasler, Chung Owyang

ANATOMIC CONSIDERATIONS

The gastrointestinal (GI) tract extends from the mouth to the anus and is composed of several organs with distinct functions. Specialized independently controlled thickened sphincters that assist in gut compartmentalization separate the organs. The gut wall is organized into well-defined layers that contribute to functional activities in each region. The mucosa is a barrier to luminal contents or a site for transfer of fluids or nutrients. Gut smooth muscle in association with the enteric nervous system mediates propulsion from one region to the next. Many GI organs possess a serosal layer that provides a supportive foundation but that also permits external input.

Interactions with other organ systems serve the needs both of the gut and the body. Pancreaticobiliary conduits deliver bile and enzymes into the duodenum. A rich vascular supply is modulated by GI tract activity. Lymphatic channels assist in gut immune activities. Intrinsic gut wall nerves provide the basic controls for propulsion and fluid regulation. Extrinsic neural input provides volitional or involuntary control to degrees that are specific for each gut region.

FUNCTIONS OF THE GASTROINTESTINAL TRACT

The GI tract serves two main functions—assimilating nutrients and eliminating waste. The gut anatomy is organized to serve these functions. In the mouth, food is processed, mixed with salivary amylase, and delivered to the gut lumen. The esophagus propels the bolus into the stomach; the lower esophageal sphincter prevents oral reflux of gastric contents. The esophageal mucosa has a protective squamous histology, which does not permit significant diffusion or absorption. Propulsive esophageal activities are exclusively aboral and coordinate with relaxation of the upper and lower esophageal sphincters on swallowing.

The stomach furthers food preparation by triturating and mixing the bolus with pepsin and acid. Gastric acid also sterilizes the upper gut. The proximal stomach serves a storage function by relaxing to accommodate the meal. The distal stomach exhibits phasic contractions that propel solid food residue against the pylorus, where it is repeatedly propelled proximally for further mixing before it is emptied into the duodenum. Finally, the stomach secretes intrinsic factor for vitamin B₁₂ absorption.

The small intestine serves most of the nutrient absorptive function of the gut. The intestinal mucosa exhibits villus architecture to provide maximal surface area for absorption and is endowed with specialized enzymes and transporters. Triturated food from the stomach mixes with pancreatic juice and bile in the duodenum to facilitate digestion. Pancreatic juice contains the main enzymes for carbohydrate, protein, and fat digestion as well as bicarbonate to optimize the pH for activation of these enzymes. Bile secreted by the liver and stored in the gallbladder is essential for intestinal lipid digestion. The proximal intestine is optimized for rapid absorption of nutrient breakdown products and most minerals, whereas the ileum is better suited for absorption of vitamin B₁₂ and bile acids. The small intestine also aids in waste elimination. Bile contains by-products of erythrocyte degradation, toxins, metabolized and unmetabolized medications, and cholesterol. Motor function of the small intestine delivers indigestible food residue and

sloughed enterocytes into the colon for further processing. The small intestine terminates in the ileocecal junction, a sphincteric structure that prevents coloileal reflux and maintains small-intestinal sterility.

The colon prepares the waste material for controlled evacuation. The colonic mucosa dehydrates the stool, decreasing daily fecal volumes from 1000–1500 mL delivered from the ileum to 100–200 mL expelled from the rectum. The colonic lumen possesses a dense bacterial colonization that ferments undigested carbohydrates and short-chain fatty acids. Whereas transit times in the esophagus are on the order of seconds and times in the stomach and small intestine range from minutes to a few hours, propagation through the colon takes more than 1 day in most individuals. Colonic motor patterns exhibit a to-and-fro character that facilitates slow fecal desiccation. The proximal colon serves to mix and absorb fluid, while the distal colon exhibits peristaltic contractions and mass actions that function to expel the stool. The colon terminates in the anus, a structure with volitional and involuntary controls to permit retention of the fecal bolus until it can be released in a socially convenient setting.

EXTRINSIC MODULATION OF GUT FUNCTION

GI function is modified by influences outside of the gut. Unlike other organ systems, the gut is in continuity with the outside environment. Thus, protective mechanisms are vigilant against deleterious effects of foods, medications, toxins, and infectious organisms. Mucosal immune mechanisms include chronic lymphocyte and plasma cell populations in the epithelial layer and lamina propria backed up by lymph node chains to prevent noxious agents from entering the circulation. Antimicrobial peptides secreted by Paneth cells in the intestine further contribute to the defense mechanisms against pathogens in the lumen. All substances absorbed into the bloodstream are filtered through the liver via the portal venous circulation. In the liver, many drugs and toxins are detoxified by a variety of mechanisms. Although intrinsic nerves control most basic gut activities, extrinsic neural input modulates many functions. Two activities under voluntary control are swallowing and defecation. Many normal GI reflexes involve extrinsic vagus or splanchnic nerve pathways. The brain-gut axis further alters function in regions not under volitional regulation. As an example, stress has potent effects on gut motor, secretory, and sensory functions.

OVERVIEW OF GASTROINTESTINAL DISEASES

GI diseases develop as a result of abnormalities within or outside of the gut and range in severity from those that produce mild symptoms and no long-term morbidity to those with intractable symptoms or adverse outcomes. Diseases may be localized to one organ or exhibit diffuse involvement at many sites.

CLASSIFICATION OF GI DISEASES

GI diseases are manifestations of alterations in nutrient assimilation or waste evacuation or in the activities supporting these main functions.

Impaired Digestion and Absorption Diseases of the stomach, intestine, biliary tree, and pancreas can disrupt digestion and absorption. The most common intestinal maldigestion syndrome, lactase deficiency, produces gas and diarrhea after ingestion of dairy products and has no adverse outcomes. Other intestinal enzyme deficiencies produce similar symptoms after ingestion of other simple sugars. Conversely, celiac disease, bacterial overgrowth, infectious enteritis, Crohn's ileitis, and radiation damage, which affect digestion and/or absorption more diffusely, produce anemia, dehydration, electrolyte disorders, or malnutrition. Gastric hypersecretory conditions such as Zollinger-Ellison

1876 syndrome damage the intestinal mucosa, impair pancreatic enzyme activation, and accelerate transit due to excess gastric acid. Biliary obstruction from stricture or neoplasm impairs fat digestion. Impaired pancreatic enzyme release in chronic pancreatitis or pancreatic cancer decreases intraluminal digestion and can lead to malnutrition.

Altered Secretion Selected GI diseases result from dysregulation of gut secretion. Gastric acid hypersecretion occurs in Zollinger-Ellison syndrome, G cell hyperplasia, retained antrum syndrome, and some individuals with duodenal ulcers. Conversely, patients with atrophic gastritis or pernicious anemia release little or no gastric acid. Inflammatory and infectious small-intestinal and colonic diseases produce fluid loss through impaired absorption or enhanced secretion. Common intestinal and colonic hypersecretory conditions cause diarrhea and include acute bacterial or viral infection, chronic *Giardia* or cryptosporidia infections, small-intestinal bacterial overgrowth, bile salt diarrhea, microscopic colitis, diabetic diarrhea, and abuse of certain laxatives. Less common causes include large colonic villus adenomas and endocrine neoplasias with tumor overproduction of secretagogue transmitters like vasoactive intestinal polypeptide.

Altered Gut Transit Impaired gut transit may be secondary to mechanical obstruction. Esophageal occlusion often results from acid-induced stricture or neoplasm. Gastric outlet obstruction develops from peptic ulcer disease or gastric cancer. Small-intestinal obstruction most commonly results from adhesions but may also occur with Crohn's disease, radiation- or drug-induced strictures, and less likely malignancy. The most common cause of colonic obstruction is colon cancer, although inflammatory strictures develop in patients with inflammatory bowel disease, after certain infections such as diverticulitis, or with some drugs.

Retardation of propulsion also develops from disordered motor function. Achalasia is characterized by impaired esophageal body peristalsis and incomplete lower esophageal sphincter relaxation. Gastroparesis is the symptomatic delay in gastric emptying of meals due to impaired gastric motility. Intestinal pseudoobstruction causes marked delays in small-bowel transit due to enteric nerve or intestinal smooth-muscle injury. Slow-transit constipation is produced by diffusely impaired colonic propulsion. Constipation also is produced by outlet abnormalities such as rectal prolapse, intussusception, or dys-synergia—a failure of anal or puborectalis relaxation upon attempted defecation.

Disorders of rapid propulsion are less common than those with delayed transit. Rapid gastric emptying occurs in postvagotomy dumping syndrome, with gastric hypersecretion, and in some cases of functional dyspepsia and cyclic vomiting syndrome. Exaggerated intestinal or colonic motor patterns may be responsible for diarrhea in irritable bowel syndrome. Accelerated transit with hyperdefecation is noted in hyperthyroidism.

Immune Dysregulation Many inflammatory GI conditions are consequences of altered gut immune function. The mucosal inflammation of celiac disease results from dietary ingestion of gluten-containing grains. Some patients with food allergy also exhibit altered immune populations. Eosinophilic esophagitis and eosinophilic gastroenteritis are inflammatory disorders with prominent mucosal eosinophils. Ulcerative colitis and Crohn's disease are disorders of uncertain etiology that produce mucosal injury primarily in the lower gut. The microscopic colitides, lymphocytic and collagenous colitis, exhibit colonic subepithelial infiltrates without visible mucosal damage. Bacterial, viral, and protozoal organisms may produce ileitis or colitis in selected patient populations.

Impaired Gut Blood Flow Different GI regions are at variable risk for ischemic damage from impaired blood flow. Rare cases of gastroparesis result from blockage of the celiac and superior mesenteric arteries. More commonly encountered are intestinal and colonic ischemia that are consequences of arterial embolus, arterial thrombosis, venous thrombosis, or hypoperfusion from dehydration, sepsis, hemorrhage, or reduced cardiac output. These may produce mucosal

injury, hemorrhage, or even perforation. Chronic ischemia may result in intestinal stricture. Some cases of radiation enterocolitis exhibit reduced mucosal blood flow.

Neoplastic Degeneration All GI regions are susceptible to malignant degeneration to varying degrees. In the United States, colorectal cancer is most common and usually presents after age 50 years. Worldwide, gastric cancer is prevalent especially in certain Asian regions. Esophageal cancer develops with chronic acid reflux or after an extensive alcohol or tobacco use history. Small-intestinal neoplasms are rare and occur with underlying inflammatory disease. Anal cancers arise after prior anal infection or inflammation. Pancreatic and biliary cancers elicit severe pain, weight loss, and jaundice and have poor prognoses. Hepatocellular carcinoma usually arises in the setting of chronic viral hepatitis or cirrhosis secondary to other causes. Most GI cancers exhibit carcinomatous histology; however, lymphomas and other cell types also are observed.

Disorders Without Obvious Organic Abnormalities The most common GI disorders show no abnormalities on biochemical or structural testing and include irritable bowel syndrome, functional dyspepsia, functional chest pain, and functional heartburn. These disorders exhibit altered gut motor function; however, the pathogenic relevance of these abnormalities is uncertain. Exaggerated visceral sensory responses to noxious stimulation may cause discomfort in these disorders. Symptoms in other patients result from altered processing of visceral pain sensations in the central nervous system. Functional bowel patients with severe symptoms may exhibit significant emotional disturbances on psychometric testing. Subtle immunologic defects may contribute to functional symptoms as well.

Genetic Influences Although many GI diseases result from environmental factors, others exhibit hereditary components. Family members of inflammatory bowel disease patients show a genetic predisposition to disease development themselves. Colonic and esophageal malignancies arise in certain inherited disorders. Rare genetic dysmotility syndromes are described. Familial clustering is even observed in the functional bowel disorders, although this may be secondary learned familial illness behavior rather than a true hereditary factor.

SYMPTOMS OF GASTROINTESTINAL DISEASE

The most common GI symptoms are abdominal pain, heartburn, nausea and vomiting, altered bowel habits, GI bleeding, and jaundice (Table 344-1). Others are dysphagia, anorexia, weight loss, fatigue, and extraintestinal symptoms.

Abdominal Pain Abdominal pain results from GI disease and extraintestinal conditions involving the genitourinary tract, abdominal wall, thorax, or spine. Visceral pain generally is midline in location and vague in character, whereas parietal pain is localized and precisely described. Common inflammatory diseases with pain include peptic ulcer, appendicitis, diverticulitis, inflammatory bowel disease, and infectious enterocolitis. Other intraabdominal causes of pain include gallstone disease and pancreatitis. Noninflammatory visceral sources include mesenteric ischemia and neoplasia. The most common causes of abdominal pain are irritable bowel syndrome and functional dyspepsia.

Heartburn Heartburn, a burning substernal sensation, is reported intermittently by at least 40% of the population. Classically, heartburn is felt to result from excess gastroesophageal reflux of acid. However, some cases exhibit normal esophageal acid exposure and may result from reflux of nonacidic material or heightened sensitivity of esophageal mucosal nerves.

Nausea and Vomiting Nausea and vomiting are caused by GI diseases, medications, toxins, acute and chronic infection, endocrine disorders, labyrinthine conditions, and central nervous system disease. The best-characterized GI etiologies relate to mechanical obstruction of the upper gut; however, disorders of propulsion including gastroparesis and intestinal pseudoobstruction also elicit prominent symptoms. Nausea and vomiting also are commonly reported by patients with

TABLE 344-1 COMMON CAUSES OF COMMON GASTROINTESTINAL (GI) SYMPTOMS

Abdominal Pain	Nausea and Vomiting	Diarrhea	GI Bleeding	Obstructive Jaundice
Appendicitis	Medications	Infection	Ulcer disease	Bile duct stones
Gallstone disease	GI obstruction	Poorly absorbed sugars	Esophagitis	Cholangiocarcinoma
Pancreatitis	Motor disorders	Inflammatory bowel disease	Varices	Cholangitis
Diverticulitis	Functional bowel disorder	Microscopic colitis	Vascular lesions	Sclerosing cholangitis
Ulcer disease	Enteric infection	Functional bowel disorder	Neoplasm	Ampullary stenosis
Esophagitis	Pregnancy	Celiac disease	Diverticula	Ampullary carcinoma
GI obstruction	Endocrine disease	Pancreatic insufficiency	Hemorrhoids	Pancreatitis
Inflammatory bowel disease	Motion sickness	Hyperthyroidism	Fissures	Pancreatic tumor
Functional bowel disorder	Central nervous system disease	Ischemia	Inflammatory bowel disease	
Vascular disease		Endocrine tumor	Infectious colitis	
Gynecologic causes				
Renal stone				

irritable bowel syndrome and functional disorders of the upper gut (including chronic idiopathic nausea and functional vomiting).

Altered Bowel Habits Altered bowel habits are common complaints of patients with GI disease. Constipation is reported as infrequent defecation, straining with defecation, passage of hard stools, or a sense of incomplete fecal evacuation. Causes of constipation include obstruction, motor disorders of the colon, medications, and endocrine diseases such as hypothyroidism and hyperparathyroidism. Diarrhea is reported as frequent defecation, passage of loose or watery stools, fecal urgency, or a similar sense of incomplete evacuation. The differential diagnosis of diarrhea is broad and includes infections, inflammatory causes, malabsorption, and medications. Irritable bowel syndrome produces constipation, diarrhea, or an alternating bowel pattern. Fecal mucus is common in irritable bowel syndrome, whereas pus characterizes inflammatory disease. Steatorrhea develops with malabsorption.

GI Bleeding Hemorrhage may develop from any gut organ. Most commonly, upper GI bleeding presents with melena or hematemesis, whereas lower GI bleeding produces passage of bright red or maroon stools. However, briskly bleeding upper sites can elicit voluminous red rectal bleeding, whereas slowly bleeding ascending colon sites may produce melena. Chronic slow GI bleeding may present with iron deficiency anemia. The most common upper GI causes of bleeding are ulcer disease, gastroduodenitis, and esophagitis. Other etiologies include portal hypertensive causes, malignancy, tears across the gastroesophageal junction, and vascular lesions. The most prevalent lower GI sources of hemorrhage include hemorrhoids, anal fissures, diverticula, ischemic colitis, and arteriovenous malformations. Other causes include neoplasm, inflammatory bowel disease, infectious colitis, drug-induced colitis, and other vascular lesions.

Jaundice Jaundice results from prehepatic, intrahepatic, or posthepatic disease. Posthepatic causes of jaundice include biliary diseases, such as choledocholithiasis, acute cholangitis, primary sclerosing cholangitis, other strictures, and neoplasm, and pancreatic disorders, such as acute and chronic pancreatitis, stricture, and malignancy.

Other Symptoms Other symptoms are manifestations of GI disease. Dysphagia,odynophagia, and unexplained chest pain suggest esophageal disease. A globus sensation is reported with esophagopharyngeal conditions, but also occurs with functional GI disorders. Weight loss, anorexia, and fatigue are nonspecific symptoms of neoplastic, inflammatory, gut motility, pancreatic, small-bowel mucosal, and psychiatric conditions. Fever is reported with inflammatory illness, but malignancies also evoke febrile responses. GI disorders also produce extraintestinal symptoms. Inflammatory bowel disease is associated with hepatobiliary dysfunction, skin and eye lesions, and arthritis. Celiac disease may present with dermatitis herpetiformis. Jaundice can produce pruritus. Conversely, systemic diseases can have GI consequences. Systemic lupus may cause gut ischemia, presenting with pain

or bleeding. Overwhelming stress or severe burns may lead to gastric ulcer formation.

EVALUATION OF THE PATIENT WITH GASTROINTESTINAL DISEASE

Evaluation of the patient with GI disease begins with a careful history and examination. Subsequent investigation with a variety of tools designed to test gut structure or function are indicated in selected cases. Some patients exhibit normal findings on diagnostic testing. In these individuals, validated symptom profiles are used to confidently diagnose a functional bowel disorder.

HISTORY

The history of the patient with suspected GI disease has several components. Symptom timing suggests specific etiologies. Symptoms of short duration commonly result from acute infection, toxin exposure, or abrupt inflammation or ischemia. Long-standing symptoms point to underlying chronic inflammatory or neoplastic conditions or functional bowel disorders. Symptoms from mechanical obstruction, ischemia, inflammatory bowel disease, and functional bowel disorders are worsened by meals. Conversely, ulcer symptoms may be relieved by eating or antacids. Symptom patterns and duration may suggest underlying etiologies. Ulcer pain occurs at intermittent intervals lasting weeks to months, whereas biliary colic has a sudden onset and lasts up to several hours. Pain from acute inflammation as with acute pancreatitis is severe and persists for days to weeks. Meals elicit diarrhea in some cases of inflammatory bowel disease and irritable bowel syndrome. Defecation relieves discomfort in inflammatory bowel disease and irritable bowel syndrome. Functional bowel disorders are exacerbated by stress. Sudden awakening from sound sleep suggests organic rather than functional disease. Diarrhea from malabsorption usually improves with fasting, whereas secretory diarrhea persists without oral intake.

Symptom relation to other factors narrows the list of diagnostic possibilities. Obstructive symptoms with prior abdominal surgery raise concern for adhesions, whereas loose stools after gastrectomy or gallbladder excision suggest dumping syndrome or postcholecystectomy diarrhea. Symptom onset after travel prompts a search for enteric infection. Medications may produce pain, altered bowel habits, or GI bleeding. Lower GI bleeding likely results from neoplasms, diverticula, or vascular lesions in an older person and from anorectal abnormalities or inflammatory bowel disease in a younger individual. Celiac disease is prevalent in people of northern European descent, whereas inflammatory bowel disease is more common in certain Jewish populations. A sexual history may raise concern for sexually transmitted diseases or immunodeficiency.

For more than two decades, working groups have been convened to devise symptom criteria to improve the confident diagnosis of functional bowel disorders and to minimize the numbers of unnecessary diagnostic tests performed. The most widely accepted symptom-based

1878 criteria are the Rome criteria. When tested against findings of structural investigations, the Rome criteria exhibit diagnostic specificities exceeding 90% for many of the functional bowel disorders.

PHYSICAL EXAMINATION

The physical exam complements information from the history. Abnormal vital signs provide diagnostic clues and determine the need for acute intervention. Fever suggests inflammation or neoplasm. Orthostasis is found with significant blood loss, dehydration, sepsis, or autonomic neuropathy. Skin, eye, or joint findings may point to specific diagnoses. Neck exam with swallowing assessment evaluates dysphagia. Cardiopulmonary disease may present with abdominal pain or nausea; thus lung and cardiac exams are important. Pelvic examination tests for a gynecologic source of abdominal pain. Rectal exam may detect blood, indicating gut mucosal injury or neoplasm or a palpable inflammatory mass in appendicitis. Metabolic conditions and gut motor disorders have associated peripheral neuropathy.

Inspection of the abdomen may reveal distention from obstruction, tumor, or ascites or vascular abnormalities with liver disease. Ecchymoses develop with severe pancreatitis. Auscultation can detect bruits or friction rubs from vascular disease or hepatic tumors. Loss of bowel sounds signifies ileus, whereas high-pitched, hyperactive sounds characterize intestinal obstruction. Percussion assesses liver size and can detect shifting dullness from ascites. Palpation assesses for hepatosplenomegaly as well as neoplastic or inflammatory masses. Abdominal exam is helpful in evaluating unexplained pain. Intestinal ischemia elicits severe pain but little tenderness. Patients with visceral pain may exhibit generalized discomfort, whereas those with parietal pain or peritonitis have directed pain, often with involuntary guarding, rigidity, or rebound. Patients with musculoskeletal abdominal wall pain may note tenderness exacerbated by Valsalva or straight-leg lift maneuvers.

TOOLS FOR PATIENT EVALUATION

Laboratory, radiographic, and functional tests can assist in diagnosis of suspected GI disease. The GI tract also is amenable to internal evaluation with upper and lower endoscopy and to examination of luminal contents. Histopathologic exams of GI tissues complement these tests.

Laboratory Selected laboratory tests facilitate the diagnosis of GI disease. Iron-deficiency anemia suggests mucosal blood loss, whereas vitamin B₁₂ deficiency results from small-intestinal, gastric, or pancreatic disease. Either also can result from inadequate oral intake. Leukocytosis and increased sedimentation rates and C-reactive protein levels are found in inflammatory conditions, whereas leukopenia is seen in viremic illness. Severe vomiting or diarrhea elicits electrolyte disturbances, acid-base abnormalities, and elevated blood urea nitrogen. Pancreaticobiliary or liver disease is suggested by elevated pancreatic or liver chemistries. Thyroid chemistries, cortisol, and calcium levels are obtained to exclude endocrinologic causes of GI symptoms. Pregnancy testing is considered for women with unexplained nausea. Serologic tests can screen for celiac disease, inflammatory bowel disease, rheumatologic diseases like lupus or scleroderma, and paraneoplastic dysmotility syndromes. Hormone levels are obtained for suspected endocrine neoplasia. Intraabdominal malignancies produce other tumor markers including the carcinoembryonic antigen CA 19-9 and α -fetoprotein. Blood testing also monitors medication therapy in some diseases, as with thiopurine metabolite levels in inflammatory bowel disease. Other body fluids are sampled under certain circumstances. Ascitic fluid is analyzed for infection, malignancy, or findings of portal hypertension. Cerebrospinal fluid is obtained for suspected central nervous system causes of vomiting. Urine samples screen for carcinoid, porphyria, and heavy metal intoxication.

Luminal Contents Luminal contents can be examined for diagnostic clues. Stool samples are cultured for bacterial pathogens, examined for leukocytes and parasites, or tested for *Giardia* antigen. Duodenal aspirates can be examined for parasites or cultured for bacterial overgrowth. Fecal fat is quantified in possible malabsorption. Stool electrolytes can be measured in diarrheal conditions. Laxative screens

are done when laxative abuse is suspected. Gastric acid is quantified to rule out Zollinger-Ellison syndrome. Esophageal pH testing is done for refractory symptoms of acid reflux, whereas impedance techniques assess for nonacidic reflux. Pancreatic juice is analyzed for enzyme or bicarbonate content to exclude pancreatic exocrine insufficiency.

Endoscopy The gut is accessible with endoscopy, which can provide the diagnosis of the causes of bleeding, pain, nausea and vomiting, weight loss, altered bowel function, and fever. **Table 344-2** lists the most common indications for the major endoscopic procedures. Upper endoscopy evaluates the esophagus, stomach, and duodenum, whereas colonoscopy assesses the colon and distal ileum. Upper endoscopy is advocated as the initial structural test performed in patients with suspected ulcer disease, esophagitis, neoplasm, malabsorption, and Barrett's metaplasia because of its ability to directly visualize as well as biopsy the abnormality. Colonoscopy is the procedure of choice for colon cancer screening and surveillance as well as diagnosis of colitis secondary to infection, ischemia, radiation, and inflammatory bowel disease. Sigmoidoscopy examines the colon up to the splenic flexure and is currently used to exclude distal colonic inflammation or obstruction in young patients not at significant risk for colon cancer. For elusive GI bleeding secondary to arteriovenous malformations or superficial ulcers, small-intestinal examination is performed with push enteroscopy, capsule endoscopy, or double-balloon enteroscopy. Capsule endoscopy also can visualize small-intestinal Crohn's disease in individuals with negative barium radiography. Endoscopic retrograde cholangiopancreatography (ERCP) provides diagnoses of pancreatic and biliary disease. Endoscopic ultrasound is useful for evaluating extent of disease in GI malignancy as well as exclusion of choledocholithiasis, evaluation of pancreatitis, drainage of pancreatic pseudocysts, and assessment of anal continuity.

Radiography/Nuclear Medicine Radiographic tests evaluate diseases of the gut and extraluminal structures. Oral or rectal contrast agents like barium provide mucosal definition from the esophagus to the rectum. Contrast radiography also assesses gut transit and pelvic floor dysfunction. Barium swallow is the initial procedure for evaluation of dysphagia to exclude subtle rings or strictures and assess for achalasia, whereas small-bowel contrast radiology reliably diagnoses intestinal tumors and Crohn's ileitis. Contrast enemas are performed when colonoscopy is unsuccessful or contraindicated. Ultrasound and computed tomography (CT) evaluate regions not accessible by endoscopy or contrast studies, including the liver, pancreas, gallbladder, kidneys, and retroperitoneum. These tests are useful for diagnosis of mass lesions, fluid collections, organ enlargement, and, in the case of ultrasound, gallstones. CT and magnetic resonance (MR) colonography are being evaluated as alternatives to colonoscopy for colon cancer screening. MR imaging assesses the pancreaticobiliary ducts to exclude neoplasm, stones, and sclerosing cholangitis, and the liver to characterize benign and malignant tumors. Specialized CT or MR enterography can assess intensity of inflammatory bowel disease. Angiography excludes mesenteric ischemia and determines spread of malignancy. Angiographic techniques also access the biliary tree in obstructive jaundice. CT and MR techniques can be used to screen for mesenteric occlusion, thereby limiting exposure to angiographic dyes. Positron emission tomography can facilitate distinguishing malignant from benign disease in several organ systems.

Scintigraphy both evaluates structural abnormalities and quantifies luminal transit. Radionuclide bleeding scans localize bleeding sites in patients with brisk hemorrhage so that therapy with endoscopy, angiography, or surgery may be directed. Radiolabeled leukocyte scans can search for intraabdominal abscesses not visualized on CT. Biliary scintigraphy is complementary to ultrasound in the assessment of cholecystitis. Scintigraphy to quantify esophageal and gastric emptying is well established, whereas techniques to measure small-intestinal or colonic transit are less widely used.

Histopathology Gut mucosal biopsies obtained at endoscopy evaluate for inflammatory, infectious, and neoplastic disease. Deep rectal biopsies assist with diagnosis of Hirschsprung's disease or amyloid. Liver

TABLE 344-2 COMMON INDICATIONS FOR ENDOSCOPY

Upper Endoscopy	Colonoscopy	Endoscopic Retrograde Cholangiopancreatography	Endoscopic Ultrasound	Capsule Endoscopy	Double-Balloon Endoscopy
Dyspepsia despite treatment	Cancer screening	Jaundice	Staging of malignancy	Obscure gastrointestinal (GI) bleeding	Ablation of small-intestinal bleeding sources
Dyspepsia with signs of organic disease	Lower GI bleeding	Postbiliary surgery complaints	Characterize and biopsy submucosal mass	Suspected Crohn's disease of the small intestine	Biopsy of suspicious small-intestinal masses/ulcers
Refractory vomiting	Anemia	Cholangitis			
Dysphagia	Diarrhea	Gallstone pancreatitis			
Upper GI bleeding	Polypectomy	Pancreatic/biliary/ampullary tumor	Bile duct stones		
Anemia	Obstruction	Unexplained pancreatitis	Chronic pancreatitis		
Weight loss	Biopsy radiologic abnormality	Pancreatitis with unremitting pain	Drain pseudocyst		
Malabsorption	Cancer surveillance: family history prior polyp/cancer, colitis	Fistulas	Anal continuity		
Biopsy radiologic abnormality	Palliate neoplasm	Biopsy radiologic abnormality			
Polypectomy	Remove foreign body	Pancreaticobiliary drainage			
Place gastrostomy	Place stent across stenosis	Sample bile			
Barrett's surveillance		Sphincter of Oddi manometry			
Palliate neoplasm					
Sample duodenal tissue/fluid					
Remove foreign body					
Endoscopic mucosal resection or ablation of dysplastic Barrett's mucosa					
Place stent across stenosis					

biopsy is indicated in cases with abnormal liver chemistries, in unexplained jaundice, following liver transplant to exclude rejection, and to characterize the degree of inflammation in patients with chronic viral hepatitis prior to initiating antiviral therapy. Biopsies obtained during CT or ultrasound can evaluate for other intraabdominal conditions not accessible by endoscopy.

Functional Testing Tests of gut function provide important data when structural testing is nondiagnostic. In addition to gastric acid and pancreatic function testing, functional testing of motor activity is provided by manometric techniques. Esophageal manometry is useful for suspected achalasia, whereas small-intestinal manometry tests for pseudoobstruction. A wireless motility capsule is now available to measure transit and contractile activity in the stomach, small intestine, and colon in a single test. Anorectal manometry with balloon expulsion testing is used for unexplained incontinence or constipation from outlet dysfunction. Anorectal manometry and electromyography also assess anal function in fecal incontinence. Biliary manometry tests for sphincter of Oddi dysfunction with unexplained biliary pain. Measurement of breath hydrogen while fasting and after oral mono- or oligosaccharide challenge can screen for carbohydrate intolerance and small-intestinal bacterial overgrowth.

TREATMENT GASTROINTESTINAL DISEASE

Management options for the patient with GI disease depend on the cause of symptoms. Available treatments include modifications in dietary intake, medications, interventional endoscopy or radiology techniques, surgery, and therapies directed to external influences.

NUTRITIONAL MANIPULATION

Dietary modifications for GI disease include treatments that only reduce symptoms, therapies that correct pathologic defects, and measures that replace normal food intake with enteral or parenteral formulations. Changes that improve symptoms but do not reverse an organic abnormality include lactose restriction for lactase deficiency, liquid meals in gastroparesis, carbohydrate restrictions

with dumping syndrome, and low-FODMAP (fermentable oligo-di-monosaccharides and polyols) diets in irritable bowel syndrome. The gluten-free diet for celiac disease exemplifies a modification that serves as primary therapy to reduce mucosal inflammation. Enteral medium-chain triglycerides replace normal fats in short-gut syndrome or severe ileal disease. Perfusion of liquid meals through a gastrostomy is performed in those who cannot swallow safely. Enteral feeding through a jejunostomy is considered for gastric dysmotility syndromes that preclude feeding into the stomach. Intravenous hyperalimentation is used for individuals with generalized gut malfunction who cannot tolerate or who cannot be sustained with enteral nutrition.

PHARMACOTHERAPY

Several medications are available to treat GI diseases. Considerable health care resources are expended on over-the-counter remedies. Many prescription drug classes are offered as short-term or continuous therapy of GI illness. A plethora of alternative treatments have gained popularity in GI conditions for which traditional therapies provide incomplete relief.

Over-the-Counter Agents Over-the-counter agents are reserved for mild GI symptoms. Antacids and histamine H₂ antagonists decrease symptoms in gastroesophageal reflux and dyspepsia, whereas antiflatulents and adsorbents reduce gaseous symptoms. More potent acid inhibitors such as proton pump inhibitors are now available over the counter for treatment of chronic gastroesophageal reflux disease (GERD). Fiber supplements, stool softeners, enemas, and laxatives are used for constipation. Laxatives are categorized as stimulants, osmotic agents (including isotonic preparations containing polyethylene glycol), and poorly absorbed sugars. Nonprescription antidiarrheal agents include bismuth subsalicylate, kaolin-pectin combinations, and loperamide. Supplemental enzymes include lactase pills for lactose intolerance and bacterial α -galactosidase to treat excess gas. In general, use of a nonprescription preparation for more than a short time for chronic persistent symptoms should be supervised by a health care provider.

1880 Prescription Drugs Prescription drugs for GI diseases are a major focus of attention from pharmaceutical companies. Potent acid suppressants, including drugs that inhibit the proton pump, are advocated for acid reflux when over-the-counter preparations are inadequate. Cytoprotective agents rarely are used for upper gut ulcers. Prokinetic drugs stimulate GI propulsion in gastroparesis and pseudoobstruction. Prosecretory drugs are prescribed for constipation refractory to other agents. Prescription antidiarrheals include opiate drugs, anticholinergic antispasmodics, tricyclics, bile acid binders, and serotonin antagonists. Antispasmodics and antidepressants also are useful for functional abdominal pain, whereas narcotics are used for pain control in organic conditions such as disseminated malignancy and chronic pancreatitis. Antiemetics in several classes reduce nausea and vomiting. Potent pancreatic enzymes decrease malabsorption and pain from pancreatic disease. Antisecretory drugs such as the somatostatin analogue octreotide treat hypersecretory states. Antibiotics treat ulcer disease secondary to *Helicobacter pylori*, infectious diarrhea, diverticulitis, intestinal bacterial overgrowth, and Crohn's disease. Some cases of irritable bowel syndrome (especially those with diarrhea) respond to nonabsorbable antibiotic therapy. Anti-inflammatory and immunosuppressive drugs are used in ulcerative colitis, Crohn's disease, microscopic colitis, refractory celiac disease, and gut vasculitis. Chemotherapy with or without radiotherapy is offered for GI malignancies. Most GI carcinomas respond poorly to such therapy, whereas lymphomas may be cured with such intervention.

Alternative Therapies Alternative treatments are marketed to treat selected GI symptoms. Ginger, acupuncture, and acustimulation have been advocated for nausea, whereas pyridoxine has been investigated for nausea of first-trimester pregnancy. Probiotics containing active bacterial cultures are used as adjuncts in some cases of infectious diarrhea and irritable bowel syndrome. Probiotics that selectively nourish benign commensal bacteria may ultimately show benefit in functional disorders as well. Low-potency pancreatic enzyme preparations are sold as general digestive aids but have little evidence to support their efficacy.

ENTERIC THERAPIES/INTERVENTIONAL ENDOSCOPY AND RADIOLOGY

Simple luminal interventions are commonly performed for GI diseases. Nasogastric tube suction decompresses the upper gut in ileus or mechanical obstruction. Nasogastric lavage of saline or water in the patient with upper GI hemorrhage determines the rate of bleeding and helps evacuate blood prior to endoscopy. Enteral feedings can be initiated through a nasogastric or nasoenteric tube. Enemas relieve fecal impaction or assist in gas evacuation in acute colonic pseudoobstruction. A rectal tube can be left in place to vent the distal colon in colonic pseudoobstruction and other colonic distention disorders.

In addition to its diagnostic role, endoscopy has therapeutic capabilities in certain settings. Cautery techniques can stop hemorrhage from ulcers, vascular malformations, and tumors. Injection with vasoconstrictor substances or sclerosants is used for bleeding ulcers, vascular malformations, varices, and hemorrhoids. Endoscopic encirclement of varices and hemorrhoids with constricting bands stops hemorrhage from these sites, whereas endoscopically placed clips can occlude arterial bleeding sites. Endoscopy can remove polyps or debulk lumen-narrowing malignancies. Endoscopic mucosal resection and radiofrequency techniques can remove or ablate some cases of Barrett's esophagus with dysplasia. Endoscopic sphincterotomy of the ampulla of Vater relieves symptoms of choledocholithiasis. Obstructions of the gut lumen and pancreaticobiliary tree are relieved by endoscopic dilatation or placement of plastic or expandable metal stents. In cases of acute colonic pseudoobstruction, colonoscopy is used to withdraw luminal gas. Finally, endoscopy is commonly used to insert feeding tubes.

Radiologic measures also are useful in GI disease. Angiographic embolization or vasoconstriction decreases bleeding from sites not amenable to endoscopic intervention. Dilatation or stenting with fluoroscopic guidance relieves luminal strictures. Contrast enemas can reduce volvulus and evacuate air in acute colonic pseudoobstruction. CT and ultrasound help drain abdominal fluid collections, in many cases obviating the need for surgery. Percutaneous transhepatic cholangiography relieves biliary obstruction when ERCP is contraindicated. Lithotripsy can fragment gallstones in patients who are not candidates for surgery. In some instances, radiologic approaches offer advantages over endoscopy for gastroenterostomy placement. Finally, central venous catheters for parenteral nutrition may be placed using radiographic techniques.

SURGERY

Surgery is performed to cure disease, control symptoms without cure, maintain nutrition, or palliate unresectable neoplasm. Medication-unresponsive ulcerative colitis, diverticulitis, cholecystitis, appendicitis, and intraabdominal abscess are curable with surgery, whereas only symptom control without cure is possible with Crohn's disease. Surgery is mandated for ulcer complications such as bleeding, obstruction, or perforation and intestinal obstructions that persist after conservative care. Fundoplication of the gastroesophageal junction is performed for severe ulcerative esophagitis and drug-refractory symptomatic acid reflux. Achalasia responds to operations to relieve lower esophageal sphincter pressure. Operations for motor disorders have been introduced including implanted electrical stimulators for gastroparesis and electrical devices and artificial sphincters for fecal incontinence. Surgery may be needed to place a jejunostomy for long-term enteral feedings. The threshold for performing surgery depends on the clinical setting. In all cases, the benefits of operation must be weighed against the potential for postoperative complications.

THERAPY DIRECTED TO EXTERNAL INFLUENCES

In some conditions, GI symptoms respond to treatments directed outside the gut. Psychological therapies including psychotherapy, behavior modification, hypnosis, and biofeedback have shown efficacy in functional bowel disorders. Patients with significant psychological dysfunction and those with little response to treatments targeting the gut are likely to benefit from this form of therapy.

345 Gastrointestinal Endoscopy

Louis Michel Wong Kee Song, Mark Topazian

Gastrointestinal endoscopy has been attempted for over 200 years, but the introduction of semirigid gastroscopes in the middle of the twentieth century marked the dawn of the modern endoscopic era. Since then, rapid advances in endoscopic technology have led to dramatic changes in the diagnosis and treatment of many digestive diseases. Innovative endoscopic devices and new endoscopic treatment modalities continue to expand the use of endoscopy in patient care.

Current flexible endoscopes provide an electronic video image generated by a charge-coupled device in the tip of the endoscope. Operator controls permit deflection of the endoscope tip; fiberoptic bundles or light-emitting diodes bring light to the tip of the endoscope; and working channels allow washing, suctioning, and the passage of instruments. Progressive changes in the diameter and stiffness of endoscopes have improved the ease and patient tolerance of endoscopy.

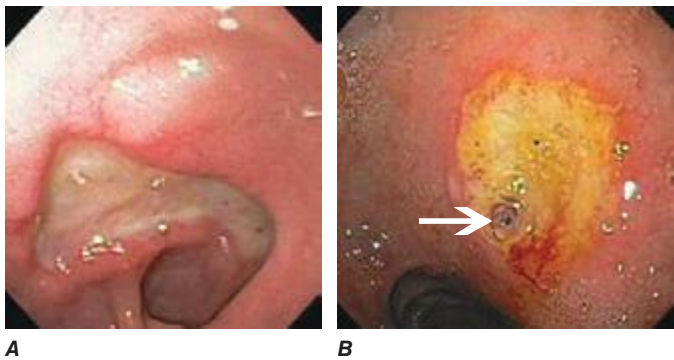


FIGURE 345-1 Duodenal ulcers. **A.** Ulcer with a clean base. **B.** Ulcer with a visible vessel (*arrow*) in a patient with recent hemorrhage.

ENDOSCOPIC PROCEDURES

UPPER ENDOSCOPY

Upper endoscopy, also referred to as esophagogastroduodenoscopy (EGD), is performed by passing a flexible endoscope through the mouth into the esophagus, stomach, and duodenum. The procedure is the best method for examining the upper gastrointestinal mucosa. While the upper gastrointestinal radiographic series has similar accuracy for diagnosis of duodenal ulcer (Fig. 345-1), EGD is superior for detection of gastric ulcers (Fig. 345-2) and flat mucosal lesions such as Barrett's esophagus (Fig. 345-3), and it permits directed biopsy and endoscopic therapy. Intravenous conscious sedation is given to most patients in the United States to ease the anxiety and discomfort of the procedure, although in many countries EGD is routinely performed with topical pharyngeal anesthesia only. Patient tolerance of unsedated EGD is improved by the use of an ultrathin, 5-mm diameter endoscope that can be passed transorally or transnasally.

COLONOSCOPY

Colonoscopy is performed by passing a flexible colonoscope through the anal canal into the rectum and colon. The cecum is reached in >95% of cases, and the terminal ileum can often be examined. Colonoscopy is the gold standard for imaging the colonic mucosa. Colonoscopy has greater sensitivity than barium enema for colitis (Fig. 345-4), polyps (Fig. 345-5), and cancer (Fig. 345-6). Computed tomography (CT) colonography is an emerging technology that rivals the accuracy of colonoscopy for detection of some polyps and cancer, although it may not be sensitive for the detection of flat lesions, such as serrated polyps (Fig. 345-7). Conscious sedation is usually given before colonoscopy in the United States, although a willing patient and a skilled examiner can complete the procedure without sedation in many cases.

FLEXIBLE SIGMOIDOSCOPY

Flexible sigmoidoscopy is similar to colonoscopy, but visualizes only the rectum and a variable portion of the left colon, typically to 60 cm

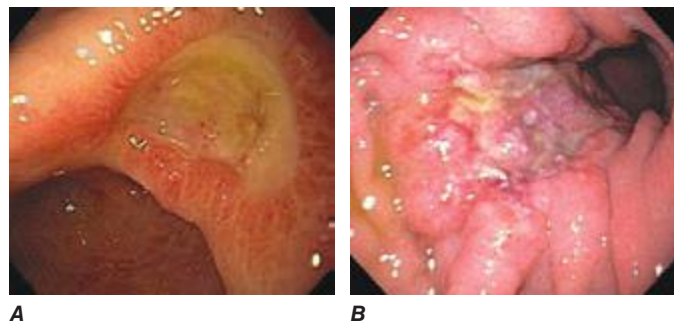


FIGURE 345-2 Gastric ulcers. **A.** Benign gastric ulcer. **B.** Malignant gastric ulcer involving greater curvature of stomach.

from the anal verge. This procedure causes abdominal cramping, but it is brief and is usually performed without sedation. Flexible sigmoidoscopy is primarily used for evaluation of diarrhea and rectal outlet bleeding.

SMALL-BOWEL ENDOSCOPY

Three endoscopic techniques are currently used to evaluate the small intestine, most often in patients presenting with presumed small-bowel bleeding. For *capsule endoscopy*, the patient swallows a disposable capsule that contains a complementary metal oxide silicon (CMOS) chip camera. Color still images (Fig. 345-8) are transmitted wirelessly to an external receiver at several frames per second until the capsule's battery is exhausted or it is passed into the toilet. Capsule endoscopy enables visualization of the small-bowel mucosa beyond the reach of a conventional endoscope and, at present, is solely a diagnostic procedure.

Push enteroscopy is performed with a long endoscope similar in design to an upper endoscope. The enteroscope is pushed down the small bowel, sometimes with the help of a stiffening overtube that extends from the mouth to the small intestine. The proximal to mid-junum is usually reached, and the instrument channel of the endoscope allows for biopsy or endoscopic therapy.

Deeper insertion into the small bowel can be accomplished by *single- or double-balloon enteroscopy* or *spiral enteroscopy* (Fig. 345-9). These instruments enable pleating of the small intestine onto an overtube (see Video 346e-1). With balloon-assisted enteroscopy, the entire intestinal tract can be visualized in some patients when both the oral and anal routes of insertion are used. Biopsies and endoscopic therapy can be performed throughout the visualized small bowel (Fig. 345-10).

ENDOSCOPIC RETROGRADE CHOLANGIOPANCREATOGRAPHY (ERCP)

During ERCP a side-viewing endoscope is passed through the mouth to the duodenum, the ampulla of Vater is identified and cannulated with a thin plastic catheter, and radiographic contrast material is injected into the bile duct and pancreatic duct under fluoroscopic guidance (Fig. 345-11). When indicated, the sphincter of Oddi can be opened using the technique of endoscopic sphincterotomy (Fig. 345-12). Stones can be retrieved from the ducts (see Video 346e-15), biopsies can be performed, strictures can be dilated and/or stented (Fig. 345-13), and ductal leaks can be stented (Fig. 345-14). ERCP is often performed for therapy but remains important in diagnosis, especially for sphincter of Oddi dysfunction and for tissue sampling of ductal strictures.

ENDOSCOPIC ULTRASOUND (EUS)

EUS utilizes high-frequency ultrasound transducers incorporated into the tip of a flexible endoscope. Ultrasound images are obtained of the gut wall and adjacent organs, vessels, and lymph nodes. By sacrificing depth of ultrasound penetration and bringing the ultrasound transducer close to the area of interest via endoscopy, high-resolution images are obtained. EUS provides the most accurate preoperative local staging of esophageal, pancreatic, and rectal malignancies (Fig. 345-15), although it does not detect most distant metastases. EUS is also useful for diagnosis of bile duct stones, gallbladder disease, submucosal gastrointestinal lesions, and chronic pancreatitis. Fine-needle aspirates and core biopsies of masses and lymph nodes in the posterior mediastinum, abdomen, pancreas, retroperitoneum, and pelvis can be obtained under EUS guidance (Fig. 345-16). EUS-guided therapeutic procedures are increasingly performed, including drainage of abscesses, pseudocysts, and pancreatic necrosis into the gut lumen (see Video 346e-2), celiac plexus neurolysis for treatment of pancreatic pain, ethanol ablation of pancreatic neuroendocrine tumors, treatment of gastrointestinal hemorrhage, and drainage of obstructed biliary and pancreatic ducts.

NATURAL ORIFICE TRANSLUMINAL ENDOSCOPIC SURGERY (NOTES)

NOTES is an evolving collection of endoscopic methods that entail passage of an endoscope or its accessories into or through the wall of the gastrointestinal tract to perform diagnostic or therapeutic

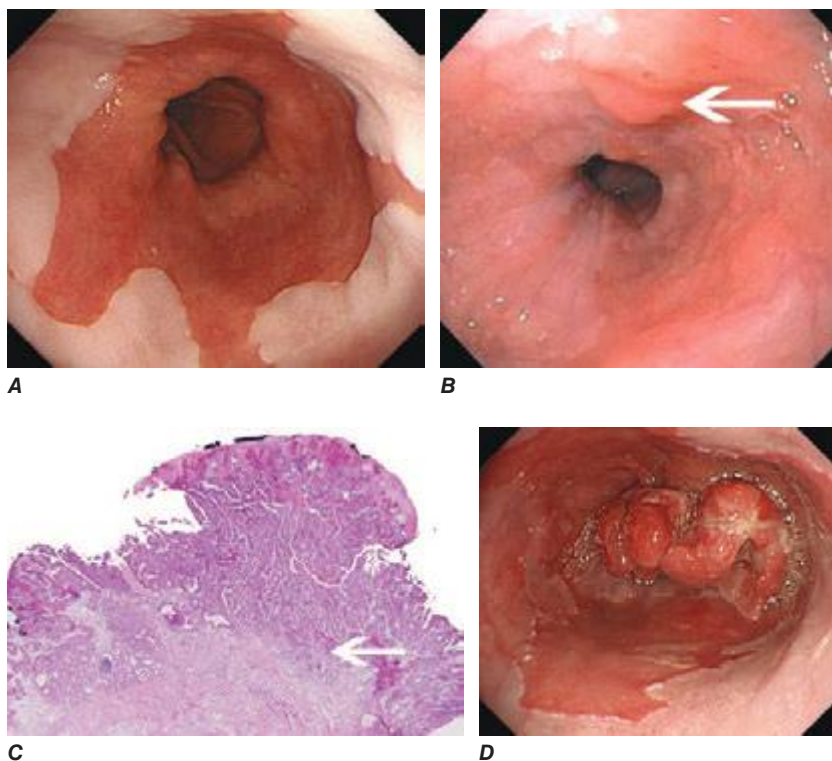


FIGURE 345-3 Barrett's esophagus. **A.** Pink tongues of Barrett's mucosa extending proximally from the gastroesophageal junction. **B.** Barrett's esophagus with a suspicious nodule (*arrow*) identified during endoscopic surveillance. **C.** Histologic finding of intramucosal adenocarcinoma in the endoscopically resected nodule. Tumor extends into the esophageal submucosa (*arrow*). **D.** Barrett's esophagus with locally advanced adenocarcinoma.

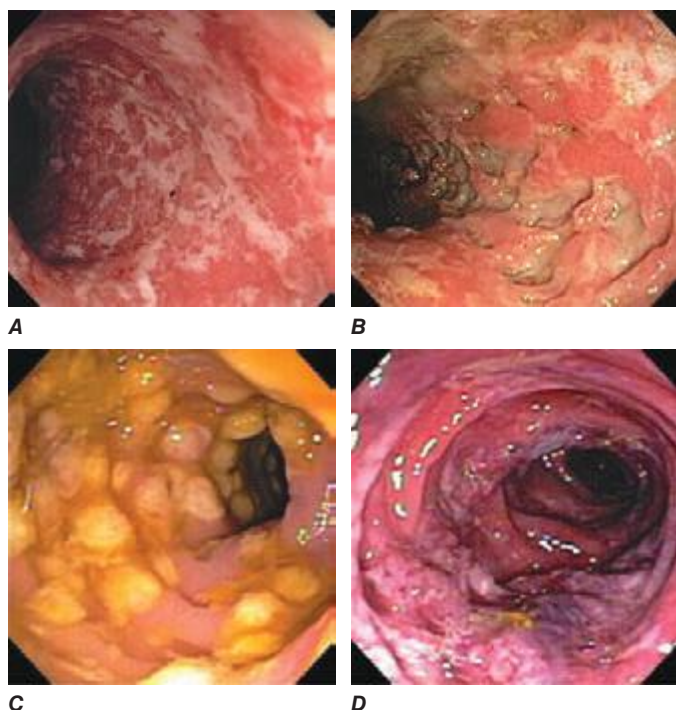


FIGURE 345-4 Causes of colitis. **A.** Chronic ulcerative colitis with diffuse ulcerations and exudates. **B.** Severe Crohn's colitis with deep ulcers. **C.** Pseudomembranous colitis with yellow, adherent pseudomembranes. **D.** Ischemic colitis with patchy mucosal edema, subepithelial hemorrhage, and cyanosis.

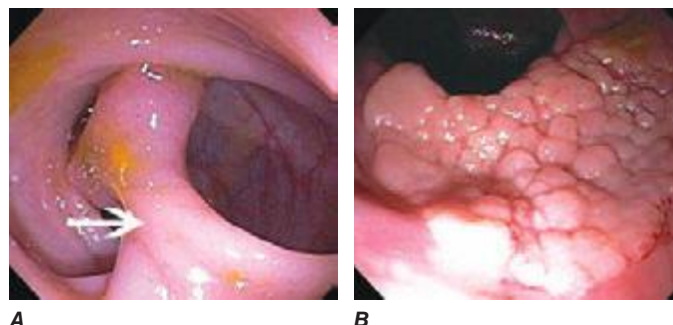


FIGURE 345-5 Colonic polyps. **A.** Pedunculated colon polyp on a thick stalk covered with normal mucosa (*arrow*). **B.** Sessile rectal polyp.



FIGURE 345-6 Colon adenocarcinoma growing into the lumen.



A



B



C

FIGURE 345-7 Flat serrated polyp in the cecum. **A.** Appearance of the lesion under conventional white-light imaging. **B.** Mucosal patterns and boundary of the lesion enhanced with narrow band imaging. **C.** Submucosal lifting of the lesion with dye (methylene blue) injection prior to resection.

interventions. Some NOTES procedures, such as percutaneous endoscopic gastrostomy (PEG) or endoscopic necrosectomy of pancreatic necrosis, are well-established clinical procedures (see Video 346e-2); others, such as per-oral endoscopic myotomy (POEM) and endoscopic full-thickness resection of gastrointestinal mural lesions (Fig. 345-17, see Video 346e-3), are emerging as viable clinical therapeutic options; and still others, such as endoscopic appendectomy, cholecystectomy, and tubal ligation, are in development, and their ultimate clinical



FIGURE 345-8 Capsule endoscopy image of jejunal vascular ectasia.

application is presently unclear. NOTES is currently an area of intense innovation and endoscopic research.

ENDOSCOPIC RESECTION AND CLOSURE TECHNIQUES

Endoscopic mucosal resection (EMR) (see Video 346e-4) and endoscopic submucosal dissection (ESD) (Fig. 345-18, see Video 346e-5) are two commonly used techniques for the resection of benign and early-stage malignant gastrointestinal neoplasms. In addition to providing larger specimens for more accurate histopathologic assessment and diagnosis, these techniques can be potentially

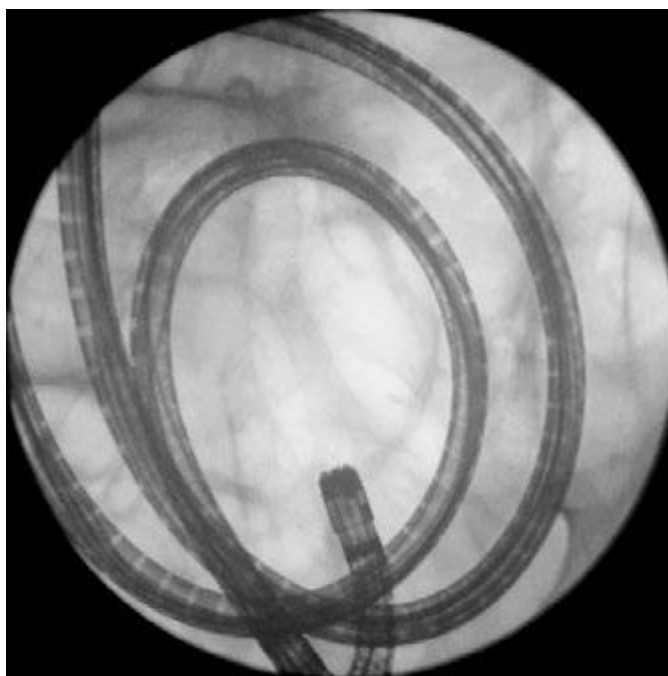


FIGURE 345-9 Radiograph of a double-balloon enteroscope in the small intestine.



FIGURE 345-10 Nonsteroidal anti-inflammatory drug (NSAID)-induced proximal ileal stricture diagnosed by double-balloon endoscopy. **A.** Ileal stricture causing obstructive symptoms. **B.** Balloon dilatation of the ileal stricture. **C.** Appearance of stricture after dilatation.

curative for certain dysplastic lesions and focal intramucosal carcinomas involving the esophagus, stomach, and colon. Several devices are also available for closure of EMR and ESD defects, as well as gastrointestinal fistulas and perforations. Endoscopic clips deployed through the working channel of an endoscope have been used for many years to treat bleeding lesions, but the development of more robust over-the-scope clips has facilitated endoscopic closure of gastrointestinal fistulas and perforations not previously amenable to endoscopic therapy (see Video 346e-6). Endoscopic suturing is also feasible, and the technique can be used to close perforations and large defects (Fig. 345-19, see Video 346e-7), anastomotic leaks, and fistulas. Other potential indications for endoscopic suturing include stent fixation to prevent its migration (Fig. 345-20), and endoscopic bariatric procedures. These technologies are likely to have an expanding role in patient care.

RISKS OF ENDOSCOPY

Medications used during conscious sedation may cause respiratory depression or allergic reactions. All endoscopic procedures carry some risk of bleeding and gastrointestinal perforation. The risk is small with diagnostic upper endoscopy and colonoscopy (<1:1000 procedures), but ranges from 0.5 to 5% when therapeutic procedures, such as EMR and ESD, control of hemorrhage, or stricture dilatation, are performed. Bleeding and perforation are rare adverse events with flexible sigmoidoscopy. The risk of adverse events for diagnostic EUS (without needle aspiration) is similar to that for diagnostic upper endoscopy.

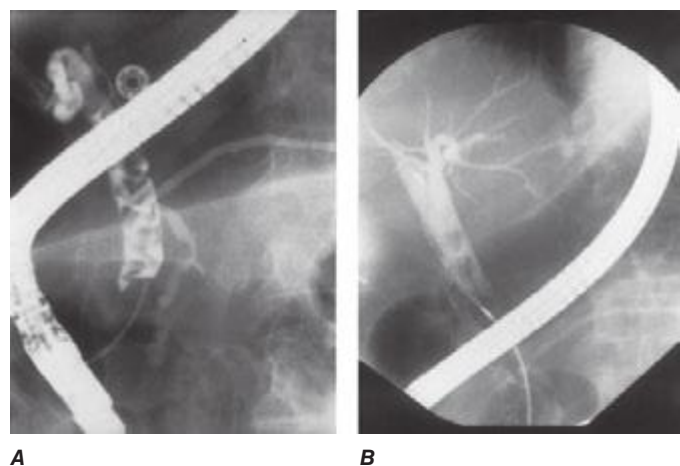


FIGURE 345-11 Endoscopic retrograde cholangiopancreatography (ERCP) for bile duct stones with cholangitis. **A.** Faceted bile duct stones are demonstrated in the common bile duct. **B.** After endoscopic sphincterotomy, the stones are extracted with a Dormia basket. A small abscess communicates with the left hepatic duct.

Infectious complications are uncommon with most endoscopic procedures. Some procedures carry a higher incidence of postprocedure bacteremia, and prophylactic antibiotics may be indicated (Table 345-1). Management of antithrombotic agents prior to endoscopic procedures should take into account the procedural risk of hemorrhage, the agent, and the patient condition, as summarized in Table 345-2.

ERCP carries additional risks. Pancreatitis occurs in about 5% of patients undergoing the procedure and in up to 30% of patients with sphincter of Oddi dysfunction. Young anicteric patients with normal ducts are at increased risk. Post-ERCP pancreatitis is usually mild and self-limited, but may result in prolonged hospitalization, surgery, diabetes, or death when severe. Bleeding occurs in 1% of endoscopic sphincterotomies. Ascending cholangitis, pseudocyst infection, retroperitoneal perforation, and abscess formation may occur as a result of ERCP.

Percutaneous gastrostomy tube placement during EGD is associated with a 10–15% incidence of adverse events, most often wound infections. Fasciitis, pneumonia, bleeding, buried bumper syndrome, and colonic injury may result from gastrostomy tube placement.

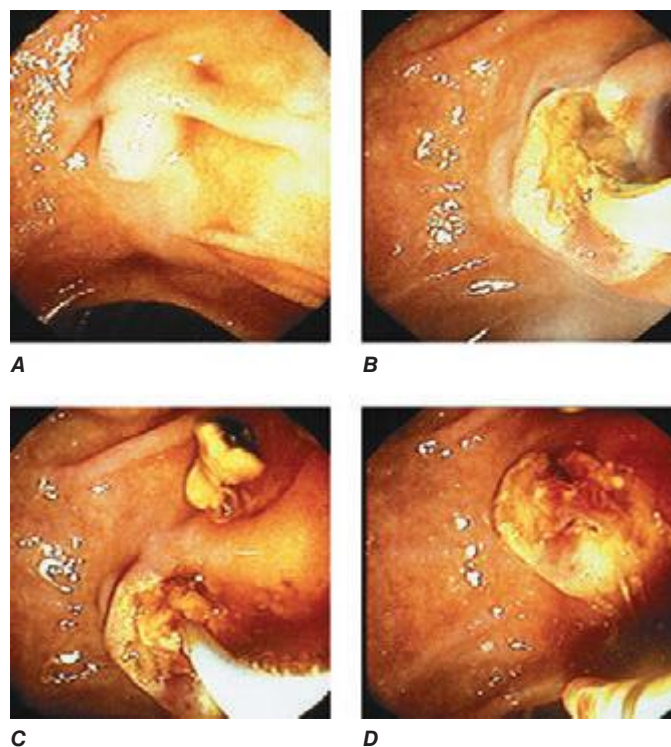


FIGURE 345-12 Endoscopic sphincterotomy. **A.** A normal-appearing ampulla of Vater. **B.** Sphincterotomy is performed with electrocautery. **C.** Bile duct stones are extracted with a balloon catheter. **D.** Final appearance of the sphincterotomy.

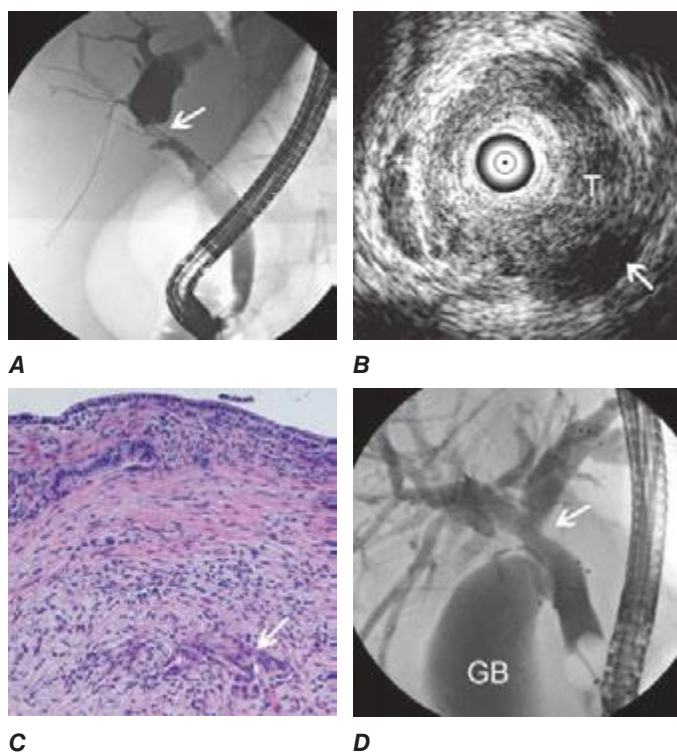


FIGURE 345-13 Endoscopic diagnosis, staging, and palliation of hilar cholangiocarcinoma. **A.** Endoscopic retrograde cholangiopancreatography (ERCP) in a patient with obstructive jaundice demonstrates a malignant-appearing stricture of the biliary confluence extending into the left and right intrahepatic ducts. **B.** Intraductal ultrasound of the biliary stricture demonstrates marked bile duct wall thickening due to tumor (T) with partial encasement of the hepatic artery (arrow). **C.** Intraductal biopsy obtained during ERCP demonstrates malignant cells infiltrating the submucosa of the bile duct wall (arrow). **D.** Endoscopic placement of bilateral self-expanding metal stents (arrow) relieves the biliary obstruction. GB, gallbladder. (Image C courtesy of Dr. Thomas Smyrk; with permission.)

URGENT ENDOSCOPY

ACUTE GASTROINTESTINAL HEMORRHAGE

Endoscopy is an important diagnostic and therapeutic technique for patients with acute gastrointestinal hemorrhage. Although gastrointestinal bleeding stops spontaneously in most cases, some patients will have persistent or recurrent hemorrhage that may be life-threatening. Clinical predictors of rebleeding help identify patients most likely

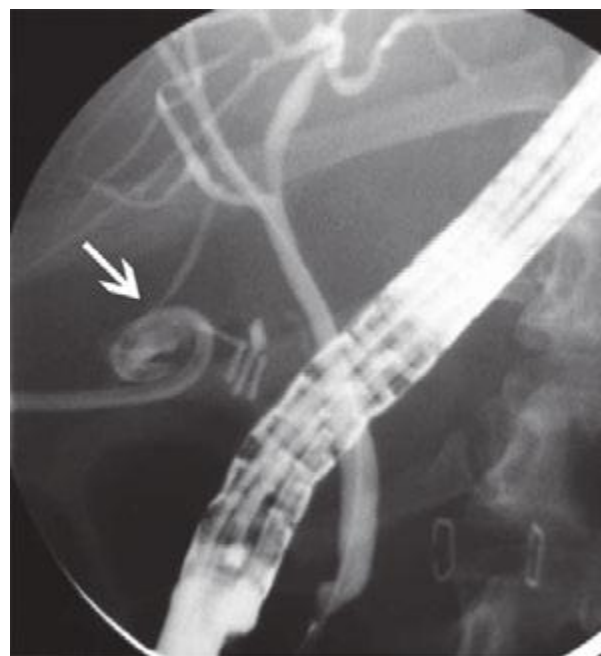


FIGURE 345-14 Bile leak (arrow) from a duct of Luschka after laparoscopic cholecystectomy. Contrast leaks from a small right intrahepatic duct into the gallbladder fossa and then flows into the pigtail of a percutaneous drainage catheter.

to benefit from urgent endoscopy and endoscopic, angiographic, or surgical hemostasis.

Initial Evaluation The initial evaluation of the bleeding patient focuses on the severity of hemorrhage as reflected by the postural vital signs, the frequency of hematemesis or melena, and (in some cases) findings on nasogastric lavage. Decreases in hematocrit and hemoglobin lag behind the clinical course and are not reliable gauges of the magnitude of acute bleeding. This initial evaluation, completed well before the bleeding source is confidently identified, guides immediate supportive care of the patient, triage to the ward or intensive care unit, and timing of endoscopy. The severity of the initial hemorrhage is the most important indication for urgent endoscopy, since a large initial bleed increases the likelihood of ongoing or recurrent bleeding. Patients with resting hypotension or orthostatic change in vital signs, repeated hematemesis, or bloody nasogastric aspirate that does not clear with large-volume lavage, or those requiring blood transfusions, should be considered for urgent endoscopy. In addition, patients with cirrhosis, coagulopathy, or respiratory or renal failure and those over 70 years of age are more likely to have significant rebleeding.

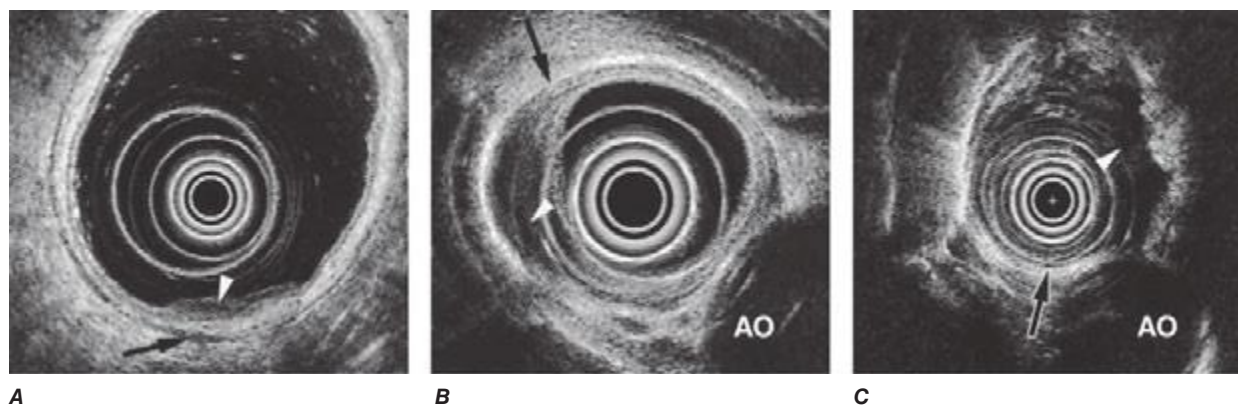


FIGURE 345-15 Local staging of gastrointestinal cancers with endoscopic ultrasound. In each example, the white arrowhead marks the primary tumor and the black arrow indicates the muscularis propria of the intestinal wall. **A.** T1 gastric cancer. The tumor does not invade the mp. **B.** T2 esophageal cancer. The tumor invades the muscularis propria. **C.** T3 esophageal cancer. The tumor extends through the muscularis propria into the surrounding tissue and focally abuts the aorta. AO, aorta.

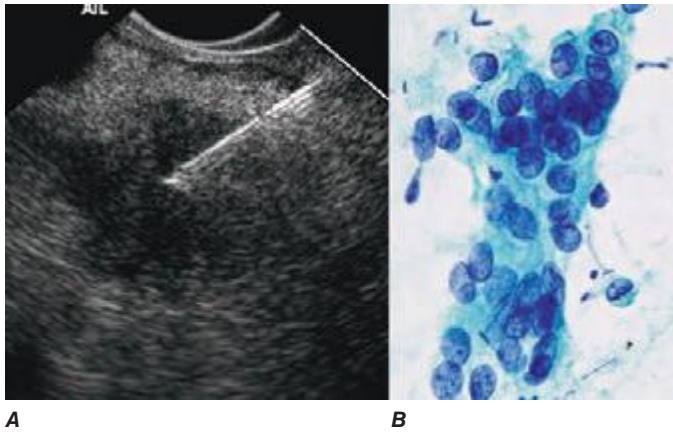


FIGURE 345-16 Endoscopic ultrasound (EUS)-guided fine-needle aspiration (FNA). **A.** Ultrasound image of a 22-gauge needle passed through the duodenal wall and positioned in a hypoechoic pancreatic head mass. **B.** Micrograph of aspirated malignant cells. (Image B courtesy of Dr. Michael R. Henry; with permission.)

Bedside evaluation also suggests an upper or lower gastrointestinal source of bleeding in most patients. Over 90% of patients with melena are bleeding proximal to the ligament of Treitz, and about 85% of patients with hematochezia are bleeding from the colon. Melena can result from bleeding in the small bowel or right colon, especially in older patients with slow colonic transit. Conversely, some patients with massive hematochezia may be bleeding from an upper gastrointestinal source, such as a gastric Dieulafoy lesion or

duodenal ulcer, with rapid intestinal transit. Early upper endoscopy should be considered in such patients.

Endoscopy should be performed after the patient has been resuscitated with intravenous fluids and transfusions, as necessary. Marked coagulopathy or thrombocytopenia is usually treated before endoscopy, since correction of these abnormalities may lead to resolution of bleeding, and techniques for endoscopic hemostasis are limited in such patients. Metabolic derangements should also be addressed. Tracheal intubation for airway protection should be considered before upper endoscopy in patients with repeated recent hematemesis, encephalopathy, and suspected variceal hemorrhage.

Most patients with significant hematochezia can undergo colonoscopy after a rapid colonic purge with a polyethylene glycol solution; the preparation fluid may be administered via a nasogastric tube. Colonoscopy has a higher diagnostic yield than radionuclide bleeding scans or angiography in lower gastrointestinal bleeding, and endoscopic therapy can be applied in some cases. In a minority of cases, endoscopic assessment is hindered by poor visualization due to persistent vigorous bleeding with recurrent hemodynamic instability, and other techniques (such as angiography or emergent subtotal colectomy) must be employed. In such patients, massive bleeding originating from an upper gastrointestinal source should also be considered and excluded by upper endoscopy. The anal and rectal mucosa should be visualized endoscopically early in the course of massive rectal bleeding, because bleeding lesions in or close to the anal canal may be identified that are amenable to endoscopic or surgical transanal hemostatic techniques.

Peptic Ulcer The endoscopic appearance of peptic ulcers provides useful prognostic information and guides the need for endoscopic therapy in patients with acute hemorrhage (Fig. 345-21). A clean-based ulcer is associated with a low risk (3–5%) of rebleeding; patients with melena and a clean-based ulcer are often discharged home from the emergency

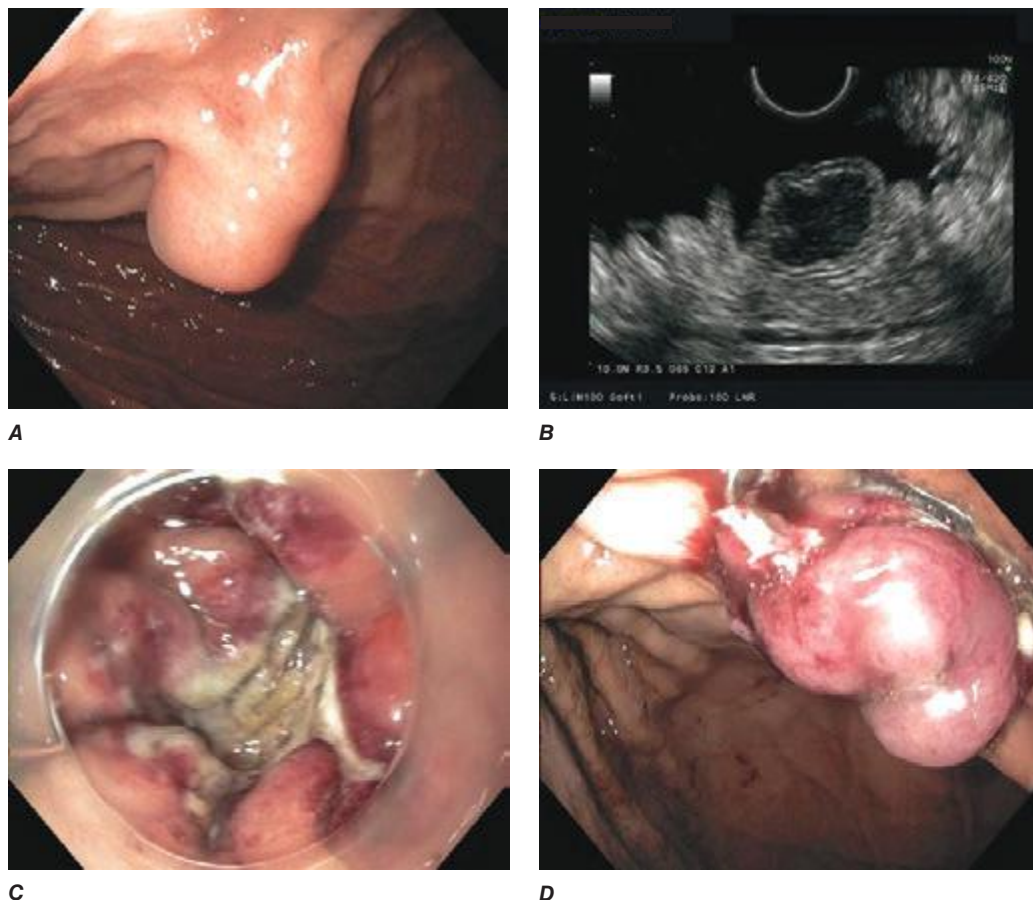
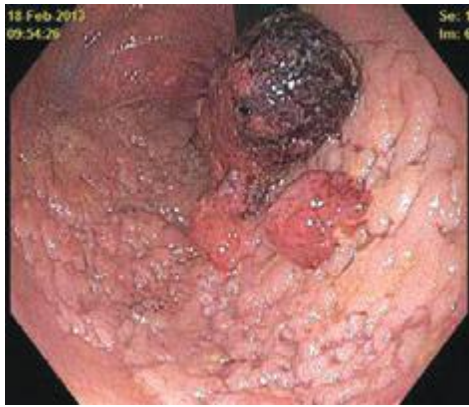


FIGURE 345-17 Endoscopic full-thickness resection of a gastrointestinal stromal tumor. **A.** Subepithelial lesion in the proximal stomach. **B.** Hypoechoic lesion arising from the fourth layer (muscularis propria) at endoscopic ultrasound. **C.** Full-thickness resection defect. **D.** Closure of defect using an over-the-scope clip.



A



B



C



D



A



B



C



D

FIGURE 345-18 Endoscopic submucosal dissection. **A.** Large flat distal rectal adenoma with central lobulation. **B.** Marking the periphery of the lesion with coagulation dots. **C.** Rectal defect following endoscopic submucosal dissection. **D.** Specimen resected en bloc.

FIGURE 345-19 Closure of large defect using an endoscopic suturing device. **A.** Ulcerated inflammatory fibroid polyp in the antrum. **B.** Large defect following endoscopic submucosal dissection of the lesion. **C.** Closure of the defect using endoscopic sutures (arrows). **D.** Resected specimen.

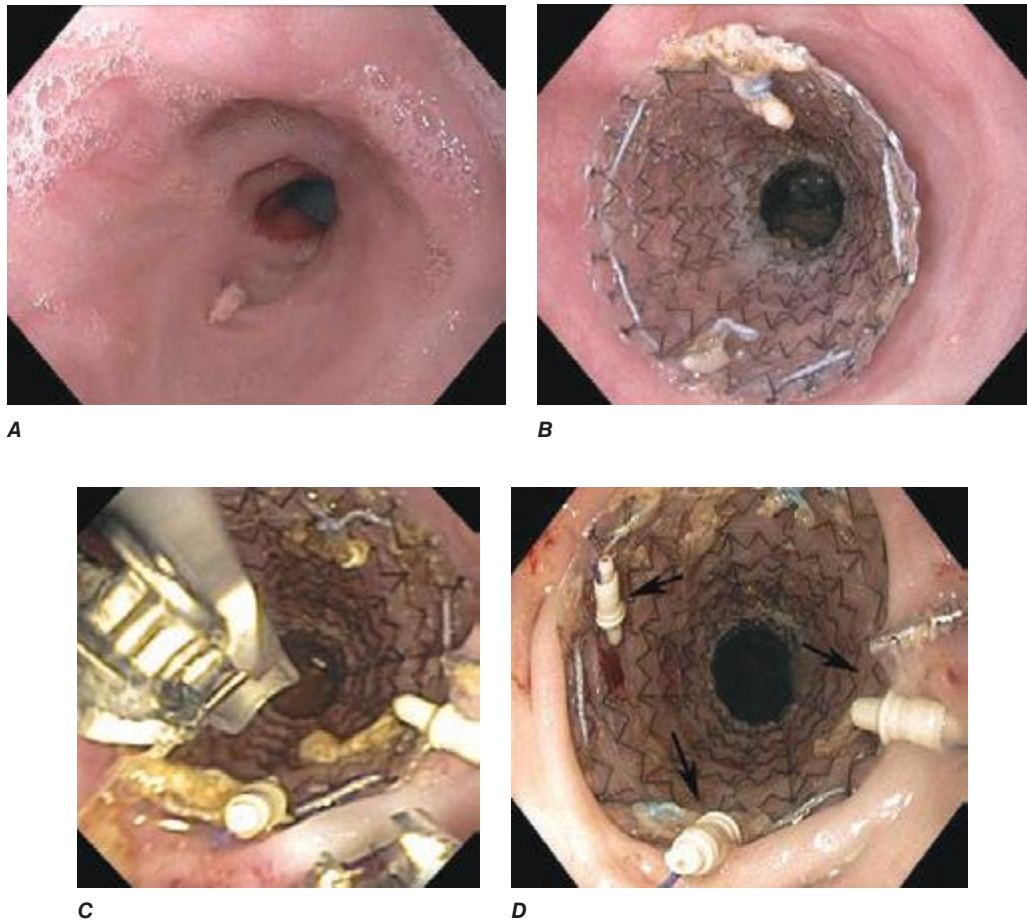


FIGURE 345-20 Prevention of stent migration using endoscopic sutures. **A.** Esophagogastric anastomotic stricture refractory to balloon dilation. **B.** Temporary placement of covered esophageal stent. **C.** Endoscopic suturing device to anchor stent to esophageal wall. **D.** Stent fixation with endoscopic sutures (*arrows*).

TABLE 345-1 ANTIBIOTIC PROPHYLAXIS FOR ENDOSCOPIC PROCEDURES

Patient Condition	Procedure Contemplated	Goal of Prophylaxis	Periprocedural Antibiotic Prophylaxis
All cardiac conditions	Any endoscopic procedure	Prevention of infective endocarditis	Not indicated
Bile duct obstruction in the absence of cholangitis	ERCP with complete drainage	Prevention of cholangitis	Not recommended
Bile duct obstruction in absence of cholangitis	ERCP with anticipated incomplete drainage (e.g., sclerosing cholangitis, hilar strictures)	Prevention of cholangitis	Recommended; continue antibiotics after the procedure
Sterile pancreatic fluid collection (e.g., pseudocyst, necrosis), which communicates with pancreatic duct	ERCP	Prevention of cyst infection	Recommended; continue antibiotics after the procedure
Sterile pancreatic fluid collection	Transmural drainage	Prevention of cyst infection	Recommended
Solid lesion along upper GI tract	EUS-FNA	Prevention of local infection	Not recommended ^a
Solid lesion along lower GI tract	EUS-FNA	Prevention of local infection	Insufficient data to make firm recommendation ^b
Cystic lesions along GI tract (including mediastinum)	EUS-FNA	Prevention of cyst infection	Recommended
All patients	Percutaneous endoscopic feeding tube placement	Prevention of peristomal infection	Recommended
Cirrhosis with acute GI bleeding	Required for all such patients, regardless of endoscopic procedures	Prevention of infectious complications and reduction of mortality	Recommended, upon admission ^c
Synthetic vascular graft and other nonvalvular cardiovascular devices	Any endoscopic procedure	Prevention of graft and device infection	Not recommended ^d
Prosthetic joints	Any endoscopic procedure	Prevention of septic arthritis	Not recommended ^e

^aLow rates of bacteremia and local infection. ^bEndoscopists may choose on a case-by-case basis. ^cRisk for bacterial infection associated with cirrhosis and GI bleeding is well established. ^dNo reported cases of infection associated with endoscopy. ^eVery low risk of infection.

Abbreviations: ERCP, endoscopic retrograde cholangiopancreatography; EUS-FNA, endoscopic ultrasound–fine-needle aspiration; GI, gastrointestinal.

Source: Adapted from S Banerjee et al: *Gastrointest Endosc* 67:719, 2008; with permission from Elsevier.

TABLE 345-2 MANAGEMENT OF ANTITHROMBOTIC DRUGS BEFORE ENDOSCOPIC PROCEDURES

Drug	Bleeding Risk of Procedure	Management	Interval Between Last Dose and Procedure	Comments
Warfarin	Low ^a	Continue	N/A	Ensure that INR is not supratherapeutic
	High ^b	Discontinue	3–7 days (usually 5), INR should be ≤ 1.5 for procedure	Consider bridging therapy with heparin ^c
New oral anticoagulants (dabigatran, rivaroxaban, apixaban)	Low ^a	Continue	N/A	
	High ^b	Discontinue	Varies from 1 to 5 days, depending on the drug and the patient's renal function	Bridging therapy generally unnecessary
Heparin	Low ^a	Continue	N/A	
	High ^b	Discontinue	4–6 h for unfractionated heparin	Skip one dose if using low-molecular-weight heparin
Aspirin	Any	Continue	N/A	Low-dose aspirin does not substantially increase the risk of endoscopic procedures
Aspirin with dipyridamole	Low ^a	Continue	N/A	
	High ^b	Discontinue	2–7 days	Consider continuing aspirin monotherapy
Thienopyridines	Low ^a	Continue	N/A	
	High ^b	Discontinue	5 days (clopidogrel or ticagrelor), 7 days (prasugrel), 10–14 days (ticlopidine)	Consider bridging therapy with aspirin

^aLow-risk endoscopic procedures include esophagogastroduodenoscopy (EGD) or colonoscopy with or without biopsy, endoscopic ultrasound (EUS) without fine-needle aspiration (FNA), and endoscopic retrograde cholangiopancreatography (ERCP) with stent exchange. ^bHigh-risk endoscopic procedures include EGD or colonoscopy with dilation, polypectomy, or thermal ablation; percutaneous endoscopic gastrostomy; EUS with FNA; and ERCP with sphincterotomy or pseudocyst drainage. ^cBridging therapy with heparin may be considered for patients discontinuing warfarin who are at high risk for thromboembolism, including those with mitral valve replacement or aortic valve replacement with other risk factors; those with nonvalvular atrial fibrillation with a history of stroke, embolic event, cardiac thrombus, or CHADS₂ score ≥ 4 ; and those with venous thromboembolism within the past 3 months or severe underlying thrombophilia.

Source: TH Baron et al: N Engl J Med 368:2113, 2013; MA Anderson et al: Gastrointest Endosc 70:1060, 2009; MJ Zuckerman et al: Gastrointest Endosc 61:189, 2005.

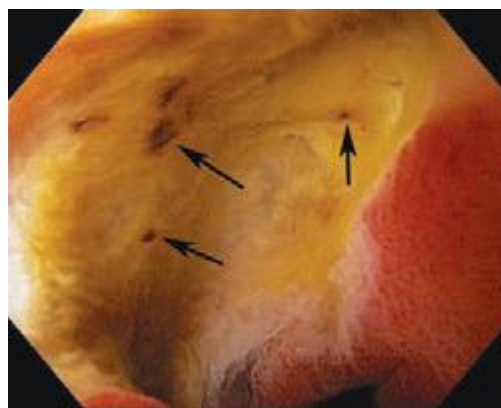
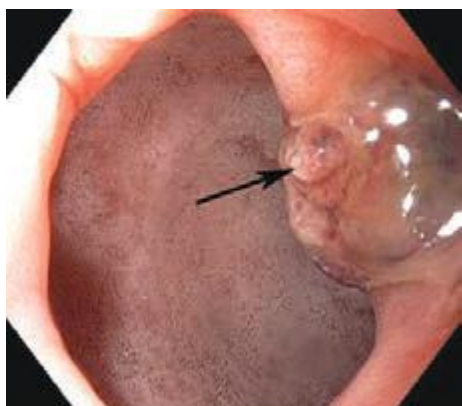
**A****B****C****D****E**

FIGURE 345-21 Stigmata of hemorrhage in peptic ulcers. **A.** Gastric antral ulcer with a clean base. **B.** Duodenal ulcer with flat pigmented spots (arrows). **C.** Duodenal ulcer with a dense adherent clot. **D.** Gastric ulcer with a pigmented protuberance/visible vessel. **E.** Duodenal ulcer with active spurting (arrow).



A



B

FIGURE 345-22 Endoscopic hemostasis of ulcer bleeding. *A.* Pyloric channel ulcer with visible vessel (*arrow*). *B.* Ulcer hemostasis with placement of an over-the-scope clip.

room or endoscopy suite if they are young, reliable, and otherwise healthy. Flat pigmented spots and adherent clots covering the ulcer base have a 10% and 20% risk of rebleeding, respectively. Endoscopic therapy is often considered for an ulcer with an adherent clot. When a fibrin plug is seen protruding from a vessel wall in the base of an ulcer (so-called sentinel clot or visible vessel), the risk of rebleeding from the ulcer is 40%. This finding generally leads to endoscopic therapy to decrease the rebleeding rate. Occasionally, active spurting from an ulcer is seen, with >90% risk of ongoing bleeding without therapy.

Endoscopic therapy of ulcers with high-risk stigmata typically lowers the rebleeding rate to 5–10%. Several hemostatic techniques are available, including injection of epinephrine or a sclerosant into and around the vessel, “coaptive coagulation” of the vessel in the base of the ulcer using a thermal probe that is pressed against the site of bleeding, placement of hemoclips (**Fig. 345-22**), or a combination of these modalities (**see Video 346e-8**). In conjunction with endoscopic therapy, the administration of a proton pump inhibitor decreases the risk of rebleeding and improves patient outcome.

Varices Two complementary strategies guide therapy of bleeding varices: local treatment of the bleeding varices and treatment of the underlying portal hypertension. Local therapies, including endoscopic variceal band ligation, endoscopic variceal sclerotherapy, and balloon tamponade with a Sengstaken-Blakemore tube, effectively control acute hemorrhage in most patients, although therapies that decrease portal pressure (pharmacologic treatment, surgical shunts, or radiologically placed intrahepatic portosystemic shunts) also play an important role.

Endoscopic variceal ligation (EVL) is indicated for the prevention of a first bleed (primary prophylaxis) from large esophageal varices (**Figs. 345-23 and 24**), particularly in patients in whom beta blockers are contraindicated or not tolerated. EVL is also the preferred endoscopic

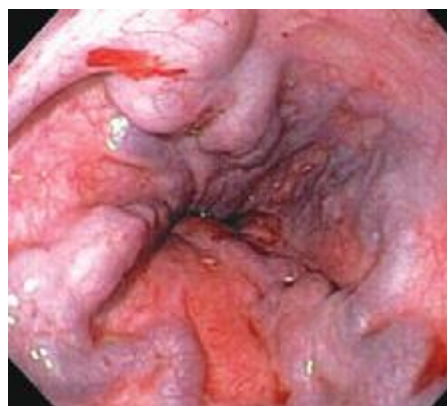


FIGURE 345-23 Esophageal varices.

therapy for control of active esophageal variceal bleeding and for subsequent eradication of esophageal varices (secondary prophylaxis). During EVL, a varix is suctioned into a cap fitted on the end of the endoscope, and a rubber band is released from the cap, ligating the varix (**Fig. 345-24**, **see Video 346e-9**). EVL controls acute hemorrhage in up to 90% of patients. Complications of EVL, such as postbanding ulcer bleeding and esophageal stenosis, are uncommon. Endoscopic variceal sclerotherapy (EVS) involves the injection of a sclerosing, thrombogenic solution into or next to esophageal varices. EVS also controls acute hemorrhage in most patients, but it is generally used as salvage therapy when band ligation fails because of its higher complication rate compared to EVL. These techniques are used when varices are actively bleeding during endoscopy or (more commonly) when



A



B

FIGURE 345-24 Endoscopic band ligation of esophageal varices. *A.* Large esophageal varices with stigmata of recent bleeding. *B.* Band ligation of varices.



A



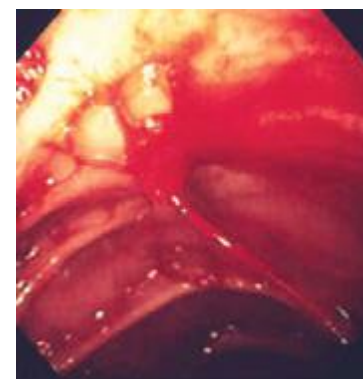
B

FIGURE 345-25 Gastric varices. **A.** Large gastric fundal varices. **B.** Stigmata of recent bleeding from the same gastric varices (arrow).

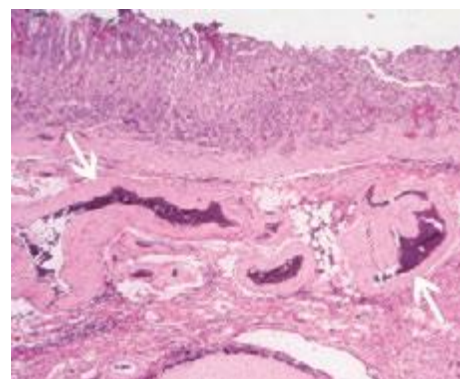
varices are the only identifiable cause of acute hemorrhage. Bleeding from large gastric fundic varices (Fig. 345-25) is best treated with endoscopic cyanoacrylate (“glue”) injection (see Video 346e-10), because EVL or EVS of these varices is associated with a high rebleeding rate. Complications of cyanoacrylate injection include infection and glue embolization to other organs, such as the lungs, brain, and spleen.

After treatment of the acute hemorrhage, an elective course of endoscopic therapy can be undertaken with the goal of eradicating esophageal varices and preventing rebleeding months to years later. However, this chronic therapy is less successful, preventing long-term rebleeding in ~50% of patients. Pharmacologic therapies that decrease portal pressure have similar efficacy, and the two modalities may be combined.

Dieulafoy’s Lesion This lesion, also called *persistent caliber artery*, is a large-caliber arteriole that runs immediately beneath the gastrointestinal mucosa and bleeds through a pinpoint mucosal erosion (Fig. 345-26). Dieulafoy’s lesion is seen most commonly on the lesser curvature of the proximal stomach, causes impressive arterial hemorrhage, and may be difficult to diagnose; it is often recognized only after repeated endoscopy for recurrent bleeding. Endoscopic therapy, such as thermal coagulation or band ligation, is typically effective for control of bleeding and ablation of the underlying vessel once the lesion has been identified (see Video 346e-11). Rescue therapies, such as angiographic embolization or surgical oversewing, are considered in situations where endoscopic therapy has failed.



A



B

FIGURE 345-26 Dieulafoy’s lesion. **A.** Actively spurting jejunal Dieulafoy’s lesion. There is no underlying mucosal lesion. **B.** Histology of a gastric Dieulafoy’s lesion. A persistent caliber artery (arrows) is present in the gastric submucosa, immediately beneath the mucosa.

Mallory-Weiss Tear A Mallory-Weiss tear is a linear mucosal rent near or across the gastroesophageal junction that is often associated with retching or vomiting (Fig. 345-27). When the tear disrupts a submucosal arteriole, brisk hemorrhage may result. Endoscopy is the best method of diagnosis, and an actively bleeding tear can be treated endoscopically with epinephrine injection, coaptive coagulation, band ligation, or hemoclips (see Video 346e-12). Unlike peptic ulcer, a Mallory-Weiss tear with a nonbleeding sentinel clot in its base rarely rebleeds and thus does not necessitate endoscopic therapy.

Vascular Ectasias Vascular ectasias are flat mucosal vascular anomalies that are best diagnosed by endoscopy. They usually cause slow intestinal blood loss and occur either in a sporadic fashion or in a well-defined pattern of distribution (e.g., gastric antral vascular ectasia [GAVE] or “watermelon stomach”) (Fig. 345-28). Cecal vascular ectasias, GAVE, and radiation-induced rectal ectasias are often responsive to local endoscopic ablative therapy, such as argon plasma coagulation (see Video 346e-13). Patients with diffuse small-bowel vascular ectasias (associated with chronic renal failure and with hereditary hemorrhagic telangiectasia) may continue to bleed despite endoscopic treatment of easily accessible lesions by conventional endoscopy. These patients may benefit from deep enteroscopy with endoscopic therapy, pharmacologic treatment with octreotide or estrogen/progesterone therapy, or intraoperative enteroscopy.

Colonic Diverticula Diverticula form where nutrient arteries penetrate the muscular wall of the colon en route to the colonic mucosa (Fig. 345-29). The artery found in the base of a diverticulum may bleed, causing painless and impressive hematochezia. Colonoscopy is indicated in patients with hematochezia and suspected diverticular hemorrhage, because other causes of bleeding (such as vascular ectasias, colitis, and colon cancer) must be excluded. In addition, an actively bleeding diverticulum may be seen and treated during colonoscopy (Fig. 345-30, see Video 346e-14).

GASTROINTESTINAL OBSTRUCTION AND PSEUDOObSTRUCTION

Endoscopy is useful for evaluation and treatment of some forms of gastrointestinal obstruction. An important exception is small-bowel obstruction due to surgical adhesions, which is generally not diagnosed or treated endoscopically. Esophageal, gastroduodenal, and colonic obstruction or pseudoobstruction can all be diagnosed and often managed endoscopically.

Acute Esophageal Obstruction

Esophageal obstruction by impacted food (Fig. 345-31) or an ingested foreign body is a potentially life-threatening event and represents an endoscopic emergency. Left untreated, the patient may develop esophageal ulceration, ischemia, and perforation. Patients with persistent esophageal obstruction often have hypersalivation and are usually

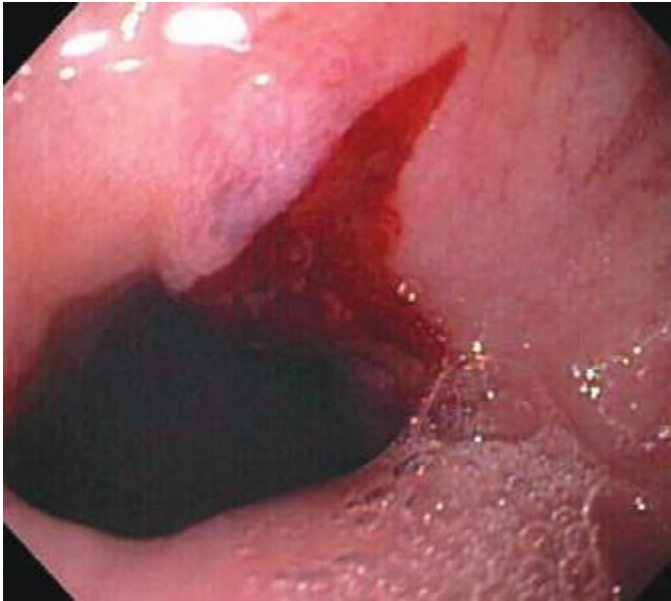


FIGURE 345-27 Mallory-Weiss tear at the gastroesophageal junction.

unable to swallow water; endoscopy is generally the best initial test in such patients, because endoscopic removal of the obstructing material is usually possible, and the presence of an underlying esophageal pathology can often be determined. Radiographs of the chest and neck should be considered before endoscopy in patients with fever, obstruction for ≥ 24 h, or ingestion of a sharp object, such as a fishbone. Radiographic contrast studies interfere with subsequent endoscopy and are not advisable in most patients with a clinical picture of esophageal obstruction. Sips of a carbonated beverage, sublingual nifedipine or nitrates, or intravenous glucagon may resolve an esophageal food impaction, but in most patients, an underlying web, ring, or stricture is present and endoscopic removal of the obstructing food bolus is necessary.

Gastric Outlet Obstruction Obstruction of the gastric outlet is commonly caused by gastric, duodenal, or pancreatic malignancy or chronic peptic ulceration with stenosis of the pylorus (Fig. 345-32). Patients vomit partially digested food many hours after eating. Gastric decompression with a nasogastric tube and subsequent lavage for removal of retained material is the first step in treatment. The diagnosis can then be confirmed with a saline load test, if desired. Endoscopy is useful for diagnosis and treatment. Patients with benign pyloric stenosis may be treated with endoscopic balloon dilatation of the pylorus, and a course of endoscopic dilatation results in long-term relief of symptoms in about 50% of patients. Malignant gastric outlet obstruction can be relieved with endoscopically placed expandable stents in patients with inoperable malignancy (Fig. 345-33).

Colonic Obstruction and Pseudoobstruction These both present with abdominal distention and discomfort; tympany; and a dilated, air-filled

colon on plain abdominal radiography. The radiographic appearance can be characteristic of a particular condition, such as sigmoid volvulus (Fig. 345-34). Both structural obstruction and pseudoobstruction may lead to colonic perforation if left untreated. Acute colonic pseudoobstruction is a form of colonic ileus that is usually attributable to electrolyte disorders, narcotic and anticholinergic medications, immobility (as after surgery), and retroperitoneal hemorrhage or mass. Multiple causative factors are often present. Colonoscopy, water-soluble contrast enema, or CT may be used to assess for an obstructing lesion and differentiate obstruction from pseudoobstruction. One of these diagnostic studies should be strongly considered if the patient does not have clear risk factors for pseudoobstruction, if radiographs do not show air in the rectum, or if the patient fails to improve when underlying causes of pseudoobstruction have been addressed. The risk of cecal perforation in pseudoobstruction rises when the cecal diameter exceeds 12 cm, and decompression of the colon may be achieved using intravenous neostigmine or via colonoscopic decompression (Fig. 345-35). Most patients should receive a trial of conservative therapy (with correction of electrolyte disorders, removal of offending medications, and increased mobilization) before undergoing an invasive decompressive procedure for colonic pseudoobstruction.

Colonic obstruction is an indication for urgent intervention. In the past, emergent diverting colostomy was usually performed with a subsequent second operation after bowel preparation to treat the underlying cause of obstruction. Colonoscopic placement of an expandable stent is now a widely used alternative that can relieve malignant colonic obstruction without emergency surgery and permit bowel preparation for an elective one-stage operation (Fig. 345-36, see Video 346e-15).

ACUTE BILIARY OBSTRUCTION

The steady, severe pain that occurs when a gallstone acutely obstructs the common bile duct often brings patients to a hospital. The diagnosis of a ductal stone is suspected when the patient is jaundiced or when serum liver tests or pancreatic enzyme levels are elevated; it is confirmed by EUS, magnetic resonance cholangiography (MRCP), or direct cholangiography (performed endoscopically, percutaneously, or during surgery). ERCP is currently the primary means of diagnosing and treating common bile duct stones in most hospitals in the United States (Figs. 345-11 and 345-12).

Bile Duct Imaging Whereas transabdominal ultrasound diagnoses only a minority of bile duct stones, MRCP and EUS are $>90\%$ accurate and have an important role in diagnosis. Examples of these modalities are shown in Fig. 345-37.

If the suspicion for a bile duct stone is high and urgent treatment is required (as in a patient with obstructive jaundice and biliary sepsis), ERCP is the procedure of choice, because it remains the gold standard for diagnosis and allows for immediate treatment (see Video 346e-16). If a persistent bile duct stone is relatively unlikely (as in a patient with gallstone pancreatitis), ERCP may be supplanted by less invasive imaging techniques, such as EUS, MRCP, or intraoperative cholangiography performed during cholecystectomy, sparing patients the risk and discomfort of ERCP.

Ascending Cholangitis Charcot's triad of jaundice, abdominal pain, and fever is present in about 70% of patients with ascending cholangitis and biliary sepsis. These patients are managed initially with fluid resuscitation and intravenous antibiotics. Abdominal ultrasound is often performed to assess for gallbladder stones and bile duct dilation. However, the bile duct may not be dilated early in the course of acute biliary obstruction. Medical management usually improves the patient's clinical status, providing a window of approximately 24 h during which biliary drainage should be established, typically by ERCP. Undue delay



FIGURE 345-28 Gastrointestinal vascular ectasias. **A.** Gastric antral vascular ectasia ("watermelon stomach") characterized by stripes of prominent flat or raised vascular ectasias. **B.** Cecal vascular ectasias. **C.** Radiation-induced vascular ectasias of the rectum in a patient previously treated for prostate cancer.

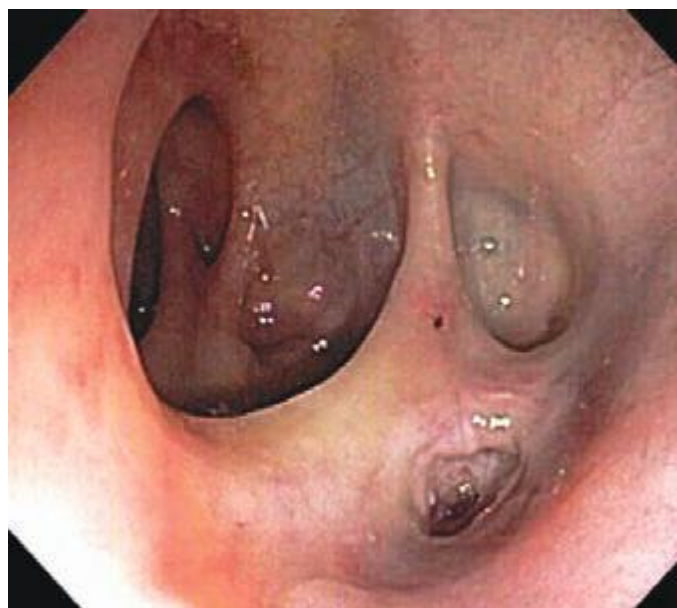


FIGURE 345-29 Colonic diverticula.

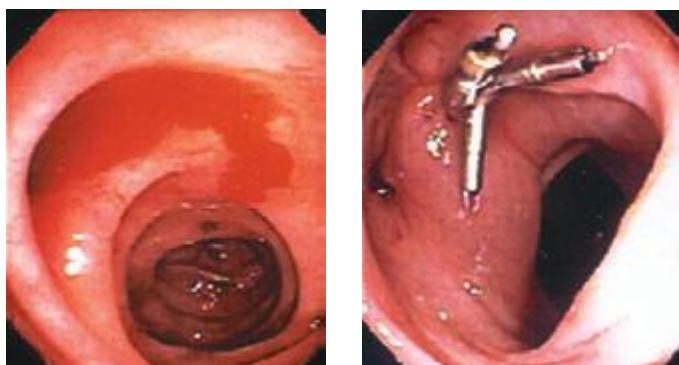


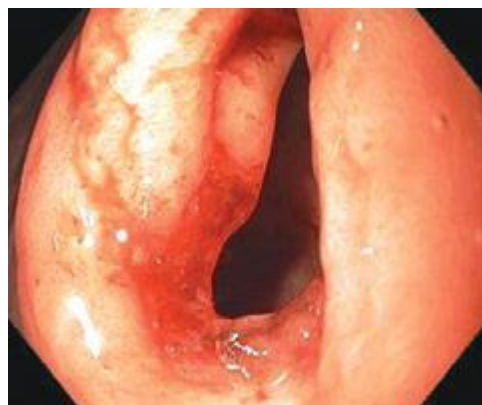
FIGURE 345-30 Diverticular hemorrhage. **A.** Actively bleeding sigmoid diverticulum. **B.** Hemostasis achieved using endoscopic clips.



A



B



C

FIGURE 345-32 Gastric outlet obstruction due to pyloric stenosis.

A. Sequela of nonsteroidal anti-inflammatory drug (NSAID)-induced ulcer disease with severe stenosis of the pylorus (*arrow*). **B.** Balloon dilation of the stenosis. **C.** Appearance of pyloric ring after dilation.



FIGURE 345-31 Esophageal food (meat) impaction.

can result in recrudescence of overt sepsis and increased morbidity and mortality rates. In addition to Charcot's triad, the additional presence of shock and confusion (Reynolds's pentad) is associated with high mortality rate and should prompt urgent intervention to restore biliary drainage.

Gallstone Pancreatitis Gallstones may cause acute pancreatitis as they pass through the ampulla of Vater. The occurrence of gallstone pancreatitis usually implies passage of a stone into the duodenum, and only about 20% of patients harbor a persistent stone in the ampulla or the common bile duct. Retained stones are more common in patients with jaundice, rising serum liver tests following hospitalization, severe pancreatitis, or superimposed ascending cholangitis.

Urgent ERCP decreases the morbidity rate of gallstone pancreatitis in a subset of patients with retained bile duct stones. It is unclear

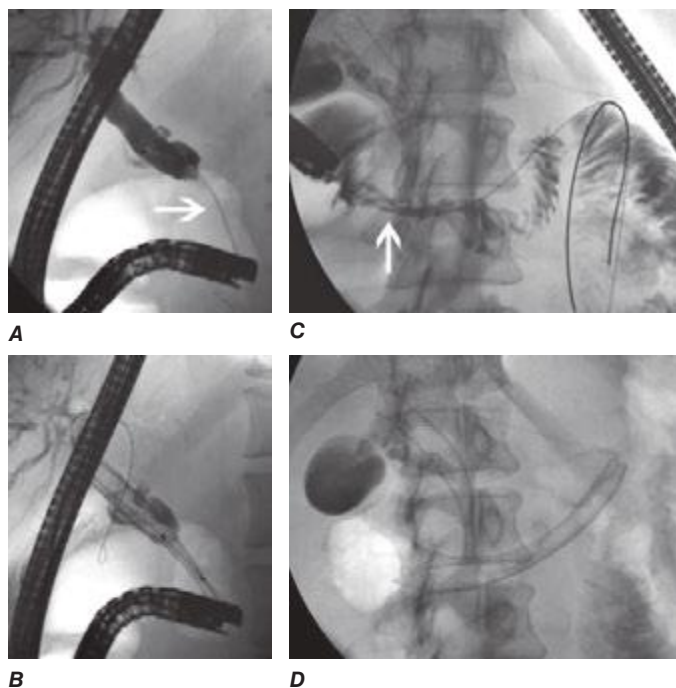


FIGURE 345-33 Biliary and duodenal self-expanding metal stents (SEMS) for obstruction caused by pancreatic cancer. **A.** Endoscopic retrograde cholangiopancreatography (ERCP) demonstrates a distal bile duct stricture (*arrow*). **B.** A biliary SEMS is placed. **C.** Contrast injection demonstrates a duodenal stricture (*arrow*). **D.** Biliary and duodenal SEMS in place.

whether the benefit of ERCP is mainly attributable to treatment and prevention of ascending cholangitis or to relief of pancreatic ductal obstruction. ERCP is warranted early in the course of gallstone pancreatitis if ascending cholangitis is suspected, especially in a jaundiced patient. Urgent ERCP may also benefit patients predicted to have severe pancreatitis using a clinical index of severity, such as the Glasgow or Ranson score. Because the benefit of ERCP is limited to patients with a retained bile duct stone, a strategy of initial MRCP or EUS for diagnosis decreases the utilization of ERCP in gallstone pancreatitis and improves clinical outcomes by limiting the occurrence of ERCP-related adverse events.



FIGURE 345-34 Sigmoid volvulus with the characteristic radiologic appearance of a “bent inner tube.”

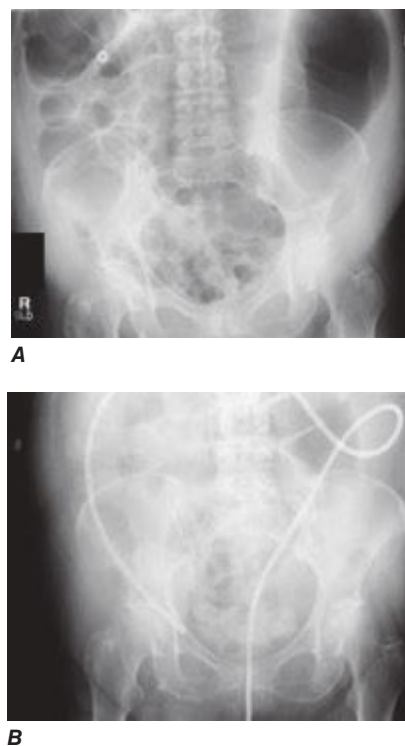


FIGURE 345-35 Acute colonic pseudoobstruction. **A.** Acute colonic dilatation occurring in a patient soon after knee surgery. **B.** Colonoscopic placement of decompression tube with marked improvement in colonic dilatation.

ELECTIVE ENDOSCOPY

DYSPEPSIA

Dyspepsia is a chronic or recurrent burning discomfort or pain in the upper abdomen that may be caused by diverse processes such as gastroesophageal reflux, peptic ulcer disease, and “nonulcer dyspepsia,” a heterogeneous category that includes disorders of motility, sensation, and somatization. Gastric and esophageal malignancies are less common causes of dyspepsia. Careful history-taking allows accurate differential diagnosis of dyspepsia in only about half of patients. In the remainder, endoscopy can be a useful diagnostic tool, especially in patients whose symptoms are not resolved by an empirical trial of symptomatic treatment. Endoscopy should be performed at the outset in patients with dyspepsia and alarm features, such as weight loss or iron-deficiency anemia.

GASTROESOPHAGEAL REFLUX DISEASE (GERD)

When classic symptoms of gastroesophageal reflux are present, such as water brash and substernal heartburn, presumptive diagnosis and empirical treatment are often sufficient. Endoscopy is a sensitive test for diagnosis of esophagitis (Fig. 345-38), but will miss nonerosive reflux disease (NERD) because some patients have symptomatic reflux without esophagitis. The most sensitive test for diagnosis of GERD is 24-h ambulatory pH monitoring. Endoscopy is indicated in patients with reflux symptoms refractory to antisecretory therapy; in those with alarm symptoms, such as dysphagia, weight loss, or gastrointestinal bleeding; and in those with recurrent dyspepsia after treatment that is not clearly due to reflux on clinical grounds alone. Endoscopy should be considered in patients with long-standing (≥ 10 years) GERD, because they have a sixfold increased risk of harboring Barrett’s esophagus compared to a patient with < 1 year of reflux symptoms. Patients with Barrett’s esophagus (Fig. 345-3) generally undergo a surveillance program of periodic endoscopy with biopsies to detect dysplasia or early carcinoma.

Barrett’s Esophagus Barrett’s esophagus is specialized columnar metaplasia that replaces the normal squamous mucosa of the distal esophagus in some persons with GERD. Barrett’s epithelium is a major risk factor

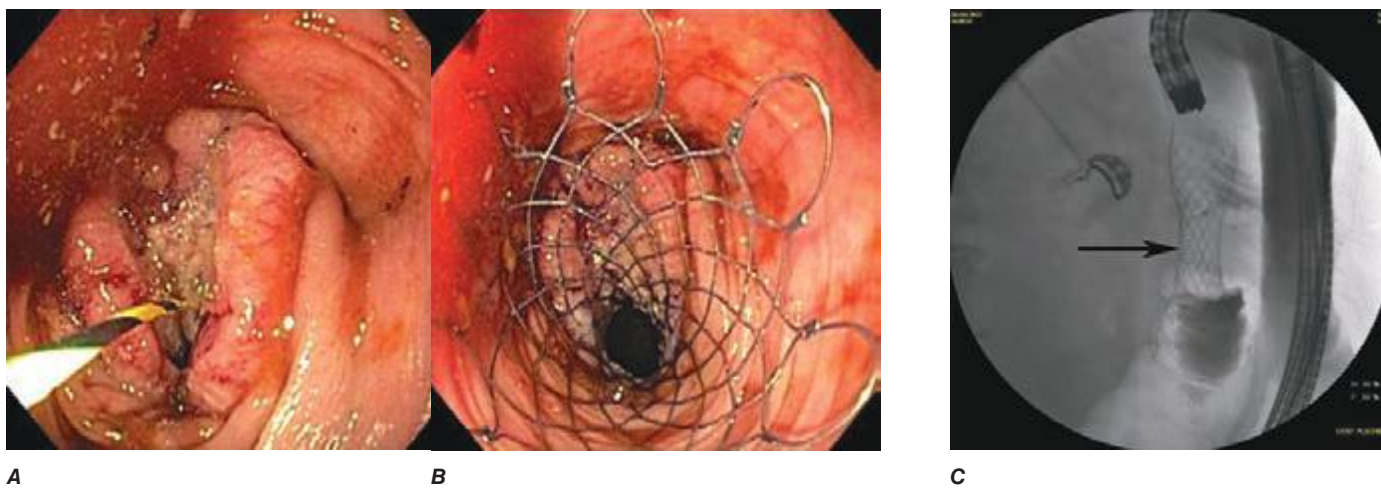


FIGURE 345-36 Obstructing colonic carcinoma. **A.** Colonic adenocarcinoma causing marked luminal narrowing of the distal transverse colon. **B.** Endoscopic placement of a self-expandable metal stent. **C.** Radiograph of expanded stent across the obstructing tumor with a residual waist (arrow).

for adenocarcinoma of the esophagus and is readily detected endoscopically, due to proximal displacement of the squamocolumnar junction (Fig. 345-3). A screening EGD for Barrett's esophagus should be considered in patients with a chronic (≥ 10 year) history of GERD symptoms. Endoscopic biopsy is the gold standard for confirmation of Barrett's esophagus and for dysplasia or cancer arising in Barrett's mucosa.

PEPTIC ULCER

Peptic ulcer classically causes epigastric gnawing or burning, often occurring nocturnally and promptly relieved by food or antacids. Although endoscopy is the most sensitive diagnostic test for peptic ulcer, it is not a cost-effective strategy in young patients with ulcer-like dyspeptic symptoms unless endoscopy is available at low cost. Patients with suspected peptic ulcer should be evaluated for *Helicobacter pylori* infection. Serology (past or present infection), urea breath testing (current infection), and stool tests are noninvasive and less costly than endoscopy with biopsy. Patients with alarm symptoms and those with persistent symptoms despite treatment should undergo endoscopy to exclude gastric malignancy and other etiologies.

NONULCER DYSPEPSIA

Nonulcer dyspepsia may be associated with bloating and, unlike peptic ulcer, tends not to remit and recur. Most patients describe marginal relief on acid-reducing, prokinetic, or anti-*Helicobacter* therapy, and are referred for endoscopy to exclude a refractory ulcer and assess for other causes. Although endoscopy is useful for excluding other diagnoses, its impact on the treatment of patients with nonulcer dyspepsia is limited.

DYSPHAGIA

About 50% of patients presenting with difficulty swallowing have a mechanical obstruction; the remainder has a motility disorder, such as achalasia or diffuse esophageal spasm. Careful history-taking often points to a presumptive diagnosis and leads to the appropriate use of diagnostic tests. Esophageal strictures (Fig. 345-39) typically cause progressive dysphagia, first for solids, then for liquids; motility disorders often cause intermittent dysphagia for both solids and liquids. Some underlying disorders have characteristic historic features: Schatzki's ring (Fig. 345-40) causes episodic dysphagia for solids, typically at the beginning of a meal; oropharyngeal motor disorders typically present with difficulty initiating deglutition (*transfer dysphagia*) and nasal reflux or coughing with swallowing; and achalasia may cause nocturnal regurgitation of undigested food.

When mechanical obstruction is suspected, endoscopy is a useful initial diagnostic test, because it permits immediate biopsy and/or dilatation of strictures, masses, or rings. The presence of linear furrows and multiple corrugated rings throughout a narrowed esophagus (*feline esophagus*) should raise suspicion for eosinophilic esophagitis, an increasingly recognized cause for recurrent dysphagia and food impaction (Fig. 345-41). Blind or forceful passage of an endoscope may lead to perforation in a patient with stenosis of the cervical esophagus or a Zenker's diverticulum, but gentle passage of an endoscope under direct visual guidance is reasonably safe. Endoscopy can miss a subtle stricture or ring in some patients.

When transfer dysphagia is evident or an esophageal motility disorder is suspected, esophageal radiography and/or a video-swallow study

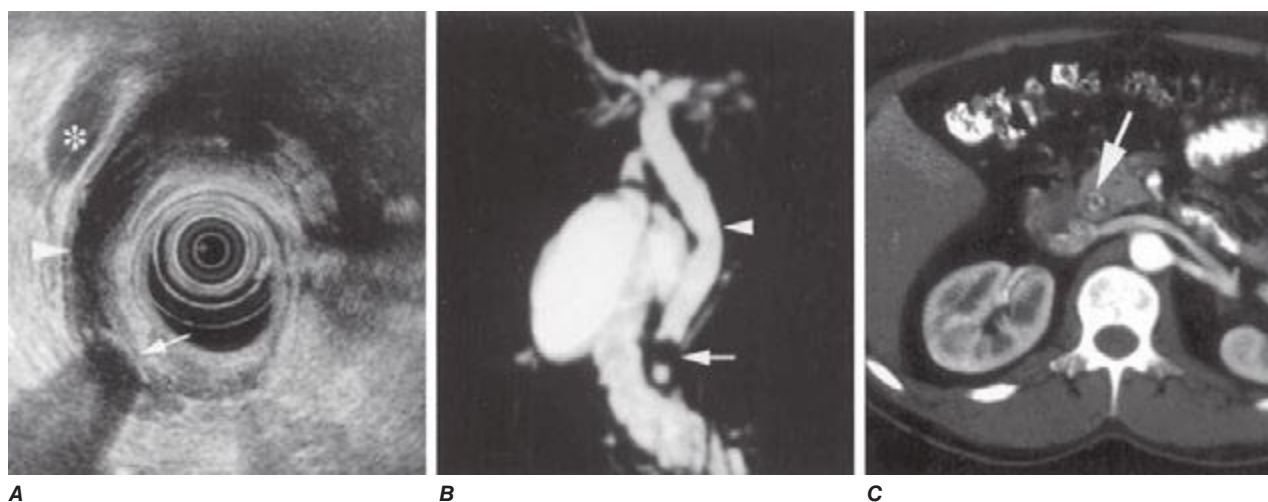


FIGURE 345-37 Methods of bile duct imaging. Arrows mark bile duct stones. Arrowheads indicate the common bile duct, and the asterisk marks the portal vein. **A.** Endoscopic ultrasound (EUS). **B.** Magnetic resonance cholangiopancreatography (MRCP). **C.** Helical computed tomography (CT).

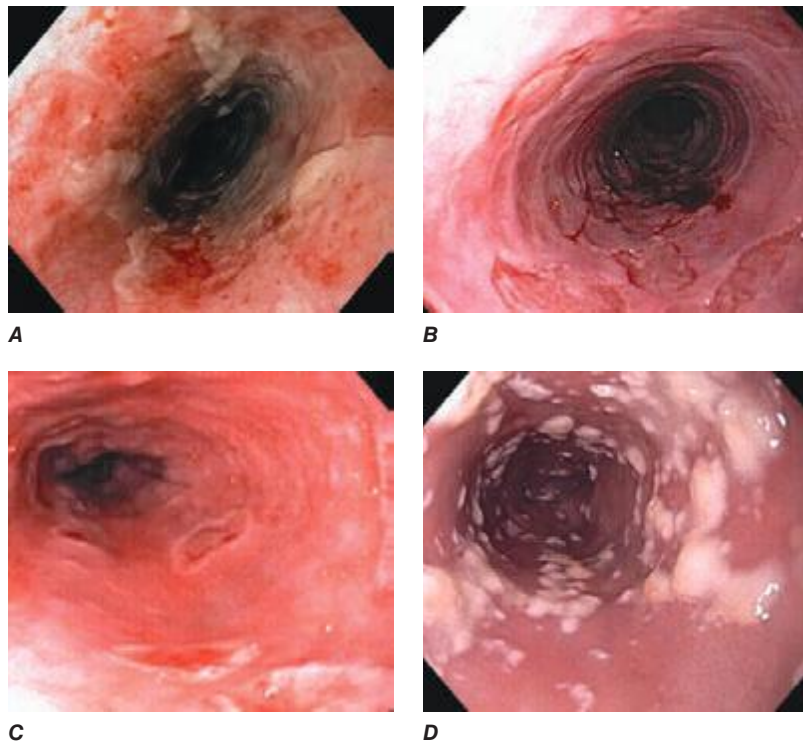


FIGURE 345-38 Causes of esophagitis. **A.** Severe reflux esophagitis with mucosal ulceration and friability. **B.** Cytomegalovirus esophagitis. **C.** Herpes simplex virus esophagitis with target-type shallow ulcerations. **D.** *Candida* esophagitis with white plaques adherent to the esophageal mucosa.

are the best initial diagnostic tests. The oropharyngeal swallowing mechanism, esophageal peristalsis, and the lower esophageal sphincter can all be assessed. In some disorders, subsequent esophageal manometry may also be important for diagnosis.

TREATMENT OF MALIGNANCIES

Endoscopy plays an important role in the treatment of gastrointestinal malignancies. Early-stage malignancies limited to the superficial layers of the gastrointestinal mucosa may be resected using the techniques of endoscopic mucosal resection (EMR) (see Video 346e-4) or endoscopic submucosal dissection (ESD) (see Video 346e-5). Photodynamic therapy (PDT) and radiofrequency ablation (RFA) are effective modalities for ablative treatment of high-grade dysplasia and intramucosal cancer in Barrett's esophagus. Gastrointestinal stromal tumors can be removed en bloc by endoscopic full-thickness resection

(see Video 346e-3). In general, endoscopic techniques offer the advantage of a minimally invasive approach to treatment, but rely on other imaging techniques (such as CT, magnetic resonance imaging [MRI], positron emission tomography [PET], and EUS) to exclude distant metastases or locally advanced disease better treated by surgery or other modalities. The decision to treat an early-stage gastrointestinal malignancy endoscopically is often made in collaboration with a surgeon and/or oncologist.

Endoscopic palliation of gastrointestinal malignancies relieves symptoms and in many cases prolongs survival. Malignant obstruction can be relieved by endoscopic stent placement (Figs. 345-13, 345-33, and 345-36; see Video 346e-15), and malignant gastrointestinal bleeding can often be palliated endoscopically as well. EUS-guided celiac plexus neurolysis may relieve pancreatic cancer pain.

ANEMIA AND OCCULT BLOOD IN THE STOOL

Iron-deficiency anemia may be attributed to poor iron absorption (as in celiac sprue) or, more commonly, chronic blood loss. Intestinal bleeding should be strongly suspected in men and postmenopausal women with iron-deficiency anemia, and colonoscopy is indicated in such patients, even in the absence of detectable occult blood in the stool. Approximately 30% will have large colonic polyps, 10% will have colorectal cancer, and a few additional patients will have colonic vascular lesions. When a convincing source of blood loss is not found in the colon, upper gastrointestinal endoscopy should be considered; if no lesion is found, duodenal biopsies should be obtained to exclude sprue (Fig. 345-42). Small-bowel evaluation with capsule endoscopy (Fig. 345-43), CT or magnetic resonance (MR) enterography, or balloon-assisted enteroscopy may be appropriate if both EGD and colonoscopy are unrevealing.

Tests for occult blood in the stool detect hemoglobin or the heme moiety and are most sensitive for colonic blood loss, although they will also detect larger amounts of upper gastrointestinal bleeding. Patients over age 50 with occult blood in normal-appearing stool should undergo colonoscopy to diagnose or exclude colorectal neoplasia. The diagnostic yield is lower than in iron-deficiency anemia. Whether upper endoscopy is also indicated depends on the patient's symptoms.

The small intestine may be the source of chronic intestinal bleeding, especially if colonoscopy and upper endoscopy are not diagnostic. The utility of small-bowel evaluation varies with the clinical setting and is most important in patients in whom bleeding causes chronic or recurrent anemia. In contrast to the low diagnostic yield



FIGURE 345-39 Peptic esophageal stricture associated with esophagitis.

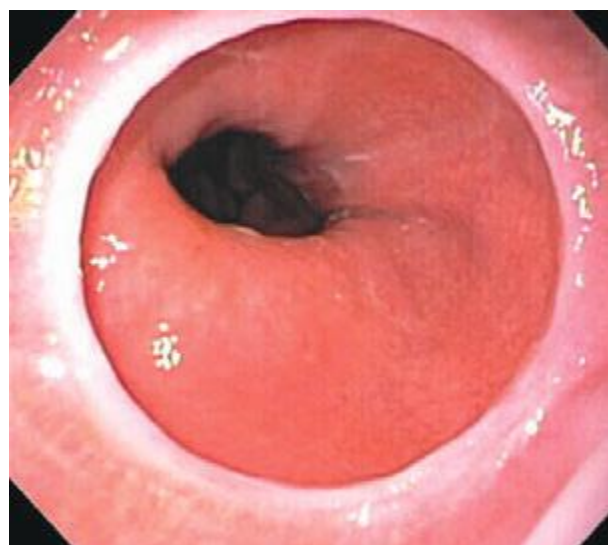


FIGURE 345-40 Schatzki's ring at the gastroesophageal junction.



FIGURE 345-41 Eosinophilic esophagitis with multiple circular rings of the esophagus creating a corrugated appearance, and an impacted grape at the narrowed esophagogastric junction. The diagnosis requires biopsy with histologic finding of > 15–20 eosinophils per high-power field.

of small-bowel radiography, positive findings on capsule endoscopy are seen in 50–70% of patients with suspected small intestinal bleeding. The most common finding is mucosal vascular ectasias. CT or MR enterography accurately detects small-bowel masses and inflammation and is also useful for initial small-bowel evaluation. Deep enteroscopy may follow capsule endoscopy for biopsy of lesions or to provide specific therapy, such as argon plasma coagulation of vascular ectasias (Fig. 345-44).

COLORECTAL CANCER SCREENING

The majority of colon cancers develop from preexisting colonic adenomas, and colorectal cancer can be largely prevented by the detection and removal of adenomatous polyps (see Video 346e-17). The choice of screening strategy for an asymptomatic person depends on personal and family history. Individuals with inflammatory bowel disease, a history of colorectal polyps or cancer, family members with adenomatous polyps or cancer, or certain familial cancer syndromes (Fig. 345-45)

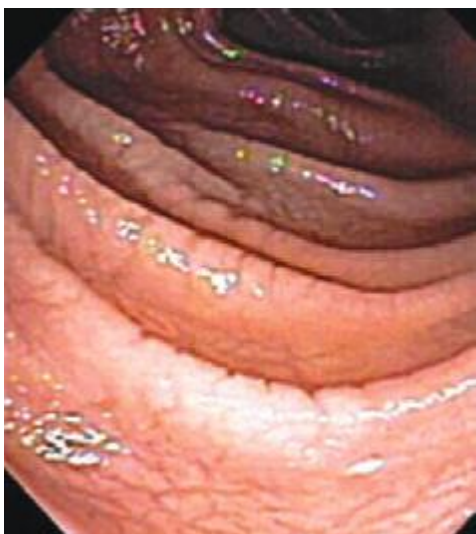


FIGURE 345-42 Scalloped duodenal folds in a patient with celiac sprue.

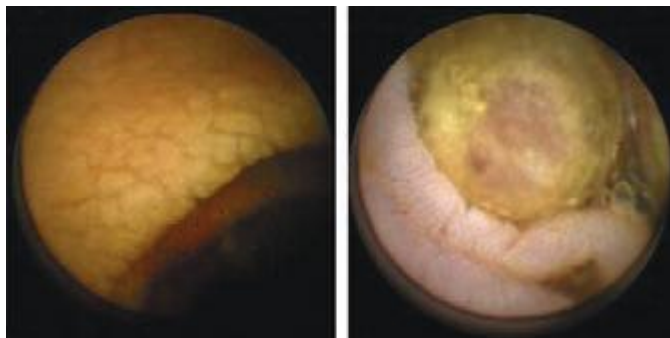


FIGURE 345-43 Capsule endoscopy images of a mildly scalloped jejunal fold (left) and an ileal tumor (right) in a patient with celiac sprue. (Images courtesy of Dr. Elizabeth Rajan; with permission.)



A



B

FIGURE 345-44 A. Mid-jejunal vascular ectasia identified by double-balloon endoscopy. B. Ablation of vascular ectasia with argon plasma coagulation.

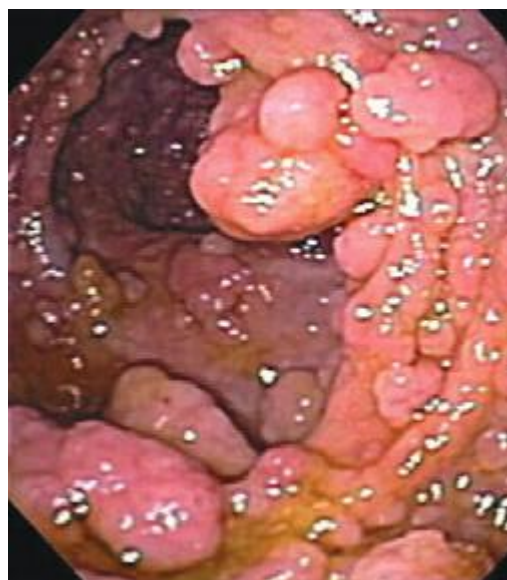


FIGURE 345-45 Innumerable colon polyps of various sizes in a patient with familial adenomatous polyposis syndrome.

TABLE 345-3 COLORECTAL CANCER SCREENING STRATEGIES

	Choices/Recommendations	Comments
Average-Risk Patients		
Asymptomatic individuals ≥ 50 years of age (≥ 45 years of age for African Americans)	Colonoscopy every 10 years ^a	Preferred cancer prevention strategy
	Annual fecal immunochemical test (FIT) or fecal occult blood test (FOBT), multiple take-home specimen cards	Cancer detection strategy; fails to detect most polyps; colonoscopy if results are positive
	Computed tomography (CT) colonography every 5 years	Colonoscopy if results are positive
	Flexible sigmoidoscopy every 5 years Double-contrast barium enema every 5 years	Fails to detect proximal colon polyps and cancers Less sensitive than colonoscopy or CT colonography; misses some cancers and polyps
Personal History of Polyps or Colorectal Cancer		
1 or 2 small (<1 cm) adenomas with low-grade dysplasia	Repeat colonoscopy in 5–10 years	Assuming complete polyp resection; interval may vary based on prior history, family history
3 to 9 adenomas, or any high-risk adenoma ^b	Repeat colonoscopy in 3 years; subsequent colonoscopy based on findings	Assuming complete polyp resection
≥ 10 adenomas	Repeat colonoscopy in <3 years based on clinical judgment	Consider evaluation for FAP or HNPCC; see recommendations below
Piecemeal removal of a sessile polyp	Exam in 2–6 months to verify complete removal	
Small (<1 cm) hyperplastic polyps of sigmoid and rectum	Repeat colonoscopy in 10 years	Those with hyperplastic polyposis syndrome merit more frequent follow-up
Sessile serrated adenoma/polyp <10 mm, without dysplasia	Repeat colonoscopy in 5 years	
Sessile serrated adenoma/polyp ≥ 10 mm or with dysplasia, or ≥ 2 serrated polyps	Repeat colonoscopy in 3 years	Serrated polyposis syndrome merits more frequent follow-up
Incompletely removed serrated polyp ≥ 1 cm	Exam in 2–6 months to verify complete removal	
Colon cancer	Evaluate entire colon around the time of resection, then repeat colonoscopy in 1 year	Subsequent colonoscopy in 3 years if the 1-year exam is normal
Inflammatory Bowel Disease		
Long-standing (>8 years) ulcerative pancolitis or Crohn's colitis, or left-sided ulcerative colitis of >15 years in duration	Colonoscopy with biopsies every 1–3 years	
Family History of Polyps or Colorectal Cancer		
First-degree relatives with only small tubular adenomas	Same as average risk	
Single first-degree relative with CRC or advanced adenoma at age ≥ 60 years	Colonoscopy every 10 years starting at age 40	
Single first-degree relative with CRC or advanced adenoma at age <60 years, OR two first-degree relatives with CRC or advanced adenomas at any age	Colonoscopy every 5 years beginning at age 40 years or 10 years younger than age at diagnosis of the youngest affected relative, whichever is earlier	
FAP	Sigmoidoscopy or colonoscopy annually, beginning at age 10–12 years	Consider genetic counseling and testing
HNPCC	Colonoscopy every 2 years beginning at age 20–25 years (or 10 years younger than the youngest affected first-degree relative) until age 40, then annually thereafter	Consider histologic evaluation for microsatellite instability in tumor specimens of patients who meet Bethesda criteria; consider genetic counseling and testing

^aAssumes good colonic preparation and complete exam to cecum. ^bHigh-risk adenoma: any adenoma ≥ 1 cm in size or containing high-grade dysplasia or villous features.

Abbreviations: CRC, colorectal cancer; FAP, familial adenomatous polyposis; HNPCC, hereditary nonpolyposis colorectal cancer.

Source: Adapted from DA Lieberman et al: *Gastroenterology* 143:844, 2012; B Levin et al: *CA Cancer J Clin* 58:130, 2008; American Cancer Society Guidelines (<http://www.cancer.org/cancer/colonandrectumcancer/moreinformation/colonandrectumcancerearlydetection/colorectal-cancer-early-detection-acs-recommendations>), accessed November 15, 2013.

are at increased risk for colorectal cancer. An individual without these factors is generally considered at average risk.

Screening strategies are summarized in Table 345-3. Although stool tests for occult blood have been shown to decrease mortality rate from colorectal cancer, they do not detect some cancers and many polyps, and direct visualization of the colon is a more effective screening strategy. Either sigmoidoscopy or colonoscopy may be used for cancer screening in asymptomatic average-risk individuals. The use of sigmoidoscopy was based on the historical finding that the majority of colorectal cancers occurred in the rectum and left colon and that patients with right-sided colon cancers had left-sided polyps. Over

the past several decades, however, the distribution of colon cancers has changed in the United States, with proportionally fewer rectal and left-sided cancers than in the past. Large American studies of colonoscopy for screening of average-risk individuals show that cancers are roughly equally distributed between left and right colon and half of patients with right-sided lesions have no polyps in the left colon. Visualization of the entire colon thus appears to be the optimal strategy for colorectal cancer screening and prevention.

Virtual colonoscopy (VC) is a radiologic technique that images the colon with CT following rectal insufflation of the colonic lumen. Computer rendering of CT images generates an electronic display of

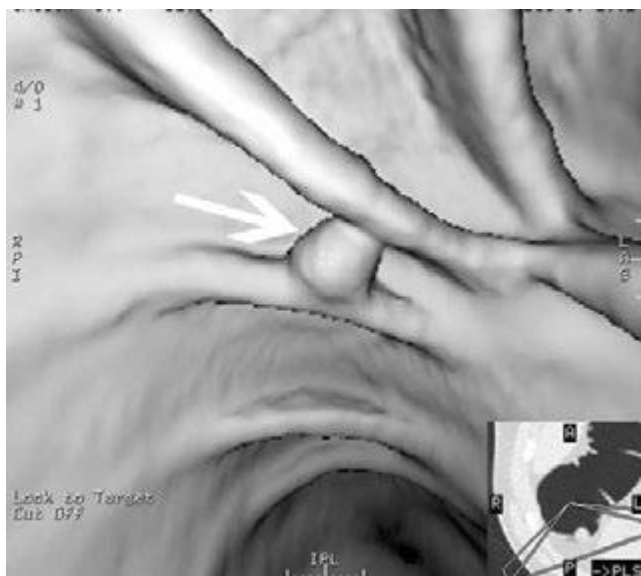


FIGURE 345-46 Virtual colonoscopy image of a colon polyp (arrow). (Image courtesy of Dr. Jeff Fidler; with permission.)

a virtual “flight” along the colonic lumen, simulating colonoscopy (Fig. 345-46). Comparative studies of virtual and routine colonoscopy have shown conflicting results, but technical refinements have improved the performance characteristics of VC. The use of VC for colorectal cancer screening may become more widespread in the future, particularly at institutions with demonstrated skill with this technique. Findings detected during virtual colonoscopy often require subsequent conventional colonoscopy for confirmation and treatment.

DIARRHEA

Most cases of diarrhea are acute, self-limited, and due to infections or medication. Chronic diarrhea (lasting >6 weeks) is more often due to a primary inflammatory, malabsorptive, or motility disorder; is less likely to resolve spontaneously; and generally requires diagnostic evaluation. Patients with chronic diarrhea or severe, unexplained acute diarrhea often undergo endoscopy if stool tests for pathogens are unrevealing. The choice of endoscopic testing depends on the clinical setting.

Patients with colonic symptoms and findings such as bloody diarrhea, tenesmus, fever, or leukocytes in stool generally undergo sigmoidoscopy or colonoscopy to assess for colitis (Fig. 345-4). Sigmoidoscopy is an appropriate initial test in most patients. Conversely, patients with symptoms and findings suggesting small-bowel disease, such as large-volume watery stools, substantial weight loss, and malabsorption of iron, calcium, or fat, may undergo upper endoscopy with duodenal aspirates for assessment of bacterial overgrowth and biopsies for assessment of mucosal diseases, such as celiac sprue.

Many patients with chronic diarrhea do not fit either of these patterns. In the setting of a long-standing history of alternating constipation and diarrhea dating to early adulthood, without findings such as blood in the stool or anemia, a diagnosis of irritable bowel syndrome may be made without direct visualization of the bowel. Steatorrhea and upper abdominal pain may prompt evaluation of the pancreas rather than the gut. Patients whose chronic diarrhea is not easily categorized often undergo initial colonoscopy to examine the entire colon and terminal ileum for inflammatory or neoplastic disease (Fig. 345-47).

MINOR HEMATOCHESIA

Bright red blood passed with or on formed brown stool usually has a rectal, anal, or distal sigmoid source (Fig. 345-48). Patients with even trivial amounts of hematochezia should be investigated with flexible sigmoidoscopy and anoscopy to exclude polyps or cancers in the distal colon. Patients reporting red blood on the toilet tissue only, without blood in the toilet or on the stool, are generally bleeding from a lesion



FIGURE 345-47 Ulcerated ileal carcinoid tumor.

in the anal canal. Careful external inspection, digital examination, and proctoscopy with anoscopy are sufficient for diagnosis in most cases.

PANCREATITIS

About 20% of patients with pancreatitis have no identified cause after routine clinical investigation (including a review of medication and alcohol use, measurement of serum triglyceride and calcium levels, abdominal ultrasonography, and CT). Endoscopic assessment leads to a specific diagnosis in the majority of such patients, often altering clinical management. Endoscopic investigation is particularly appropriate if the patient has had more than one episode of pancreatitis.

Microolithiasis, or the presence of microscopic crystals in bile, is a leading cause of previously unexplained acute pancreatitis and is sometimes seen during abdominal ultrasonography as layering sludge or flecks of floating, echogenic material in the gallbladder. Gallbladder bile can be obtained for microscopic analysis by administering a cholecystokinin analogue during endoscopy, causing contraction of the gallbladder. Bile is suctioned from the duodenum as it drains from the papilla, and the darkest fraction is examined for cholesterol crystals or bilirubinate granules. The combination of EUS of the gallbladder and bile microscopy is probably the most sensitive means of diagnosing microolithiasis.



FIGURE 345-48 Internal hemorrhoids with bleeding (arrow) as seen on a retroflexed view of the rectum.

Previously undetected chronic pancreatitis, pancreatic malignancy, or pancreas divisum may be diagnosed by either ERCP or EUS. Sphincter of Oddi dysfunction or stenosis is a potential cause for pancreatitis and can be diagnosed by manometric studies performed during ERCP. Autoimmune pancreatitis may require EUS-guided pancreatic biopsy for histologic diagnosis.

Severe pancreatitis often results in pancreatic fluid collections. Both pseudocysts and areas of walled-off pancreatic necrosis can be drained into the stomach or duodenum endoscopically, using transpapillary and transmural endoscopic techniques. Pancreatic necrosis can be treated by direct endoscopic necrosectomy (see Video 346e-2).

CANCER STAGING

Local staging of esophageal, gastric, pancreatic, bile duct, and rectal cancers can be obtained with EUS (Fig. 345-15). EUS with fine-needle aspiration (Fig. 345-16) currently provides the most accurate preoperative assessment of local tumor and nodal staging, but it does not detect most distant metastases. Details of the local tumor stage can guide treatment decisions including resectability and need for neoadjuvant therapy. EUS with transesophageal needle biopsy may also be used to assess the presence of non-small-cell lung cancer in mediastinal nodes.

OPEN-ACCESS ENDOSCOPY

Direct scheduling of endoscopic procedures by primary care physicians without preceding gastroenterology consultation, or *open-access endoscopy*, is common. When the indications for endoscopy are clear-cut and appropriate, the procedural risks are low, and the patient understands what to expect, open-access endoscopy streamlines patient care and decreases costs.

Patients referred for open-access endoscopy should have a recent history, physical examination, and medication review. A copy of such an evaluation should be available when the patient comes to the endoscopy suite. Patients with unstable cardiovascular or respiratory conditions should not be referred directly for open-access endoscopy. Patients with particular conditions and undergoing certain procedures should be prescribed prophylactic antibiotics prior to endoscopy (Table 345-1). In addition, patients taking anticoagulants and/or antiplatelet drugs may require adjustment of these agents before endoscopy based on the procedure risk for bleeding and condition risk for a thromboembolic event (Table 345-2).

Common indications for open-access EGD include dyspepsia resistant to a trial of appropriate therapy; dysphagia; gastrointestinal bleeding; and persistent anorexia or early satiety. Open-access colonoscopy is often requested in men or postmenopausal women with iron-deficiency anemia, in patients over age 50 with occult blood in the stool, in patients with a previous history of colorectal adenomatous polyps or cancer, and for colorectal cancer screening. Flexible sigmoidoscopy is commonly performed as an open-access procedure.

When patients are referred for open-access colonoscopy, the primary care provider may need to choose a colonic preparation. Commonly used oral preparations include polyethylene glycol lavage solution, with or without citric acid. A “split-dose” regimen improves the quality of colonic preparation. Sodium phosphate purgatives may cause fluid and electrolyte abnormalities and renal toxicity, especially in patients with renal failure or congestive heart failure and those over 70 years of age.

Gastrointestinal endoscopy is an increasingly important method for diagnosis and treatment of disease. This atlas demonstrates endoscopic findings in a variety of gastrointestinal infectious, inflammatory, vascular, and neoplastic conditions. Cancer screening and prevention are common indications for gastrointestinal endoscopy, and the premalignant conditions of Barrett's esophagus and colonic polyps are illustrated. Endoscopic treatment modalities for gastrointestinal bleeding, polyps, and biliary stones are demonstrated in video clips. The images shown in this atlas are also found in [Chap. 345](#) of the book.

VIDEO 346e-1 **Methods of deep enteroscopy.** (*Animations courtesy of Dr. Mark Stark and Dr. Jonathan Leighton; with permission.*)

VIDEO 346e-2 **Pancreatic necrosis** treated by transduodenal endoscopic drainage and necrosectomy.

VIDEO 346e-3 **Endoscopic full-thickness resection** of a gastric subepithelial lesion.

VIDEO 346e-4 **Endoscopic submucosal dissection** of a large rectal adenoma.

VIDEO 346e-5 **Over-the-scope clip closure** of a spontaneous esophageal perforation.

VIDEO 346e-6 **Endoscopic suturing** for stent fixation.

VIDEO 346e-7 **Actively bleeding duodenal ulcer** treated with dilute epinephrine injection, thermal probe application, and hemoclips. (*Video courtesy of Dr. Navtej Buttar; with permission.*)

VIDEO 346e-8 **Actively bleeding esophageal varices** treated with endoscopic band ligation.

VIDEO 346e-9 **Large, bleeding gastric varix** treated with endoscopic cyanoacrylate injection.

VIDEO 346e-10 **Dieulafoy's lesion** treated endoscopically.

VIDEO 346e-11 **Bleeding Mallory-Weiss tear** treated with hemoclip placement.

VIDEO 346e-12 **Radiation proctopathy** treated with argon plasma coagulation.

VIDEO 346e-13 **Actively bleeding colonic diverticulum** treated with dilute epinephrine injection and band ligation.

VIDEO 346e-14 **Stent placement** for palliation of malignant colonic obstruction.

VIDEO 346e-15 **Bile duct stones** removed after endoscopic sphincterotomy.

VIDEO 346e-16 **Barrett's esophagus with high-grade dysplasia** treated with endoscopic mucosal resection.

VIDEO 346e-17 **Pedunculated and sessile colonic polyps** removed with snare cautery during colonoscopy.

347 Diseases of the Esophagus

Peter J. Kahrilas, Ikuo Hirano

ESOPHAGEAL STRUCTURE AND FUNCTION

The esophagus is a hollow, muscular tube coursing through the posterior mediastinum joining the hypopharynx to the stomach with a sphincter at each end. It functions to transport food and fluid between these ends, otherwise remaining empty. The physiology of swallowing, esophageal motility, and oral and pharyngeal dysphagia are described in [Chap. 53](#). Esophageal diseases can be manifested by impaired function or pain. Key functional impairments are swallowing disorders and excessive gastroesophageal reflux. Pain, sometimes indistinguishable from cardiac chest pain, can result from inflammation, infection, dysmotility, or neoplasm.

SYMPTOMS OF ESOPHAGEAL DISEASE

The clinical history remains central to the evaluation of esophageal symptoms. A thoughtfully obtained history will often expedite management. Important details include weight gain or loss, gastrointestinal bleeding, dietary habits including the timing of meals, smoking, and alcohol consumption. The major esophageal symptoms are heartburn, regurgitation, chest pain, dysphagia, odynophagia, and globus sensation.

Heartburn (pyrosis), the most common esophageal symptom, is characterized by a discomfort or burning sensation behind the sternum that arises from the epigastrium and may radiate toward the neck. Heartburn is an intermittent symptom, most commonly experienced after eating, during exercise, and while lying recumbent. The discomfort is relieved with drinking water or antacid but can occur frequently interfering with normal activities including sleep. The association between heartburn and gastroesophageal reflux disease (GERD) is so strong that empirical therapy for GERD has become accepted management. However, the term “heartburn” is often misused and/or referred to with other terms such as “indigestion” or “repeating,” making it important to clarify the intended meaning.

Regurgitation is the effortless return of food or fluid into the pharynx without nausea or retching. Patients report a sour or burning fluid in the throat or mouth that may also contain undigested food particles. Bending, belching, or maneuvers that increase intraabdominal pressure can provoke regurgitation. A clinician needs to discriminate among regurgitation, vomiting, and rumination. *Vomiting* is preceded by nausea and accompanied by retching. *Rumination* is a behavior in which recently swallowed food is regurgitated and then reswallowed repetitively for up to an hour. Although there is some linkage between rumination and mental deficiency, the behavior is also exhibited by unimpaired individuals who sometimes even find it pleasurable.

Chest pain is a common esophageal symptom with characteristics similar to cardiac pain, sometimes making this distinction difficult. Esophageal pain is usually experienced as a pressure type sensation in the mid chest, radiating to the mid back, arms, or jaws. The similarity to cardiac pain is likely because the two organs share a nerve plexus and the nerve endings in the esophageal wall have poor discriminative ability among stimuli. Esophageal distention or even chemostimulation (e.g., with acid) will often be perceived as chest pain. Gastroesophageal reflux is the most common cause of esophageal chest pain.

Esophageal *dysphagia* (Chap. 53) is often described as a feeling of food “sticking” or even lodging in the chest. Important distinctions are between uniquely solid food dysphagia as opposed to liquid and solid, episodic versus constant dysphagia, and progressive versus static dysphagia. If the dysphagia is for liquids as well as solid food, it suggests a motility disorder such as achalasia. Conversely, uniquely solid food dysphagia is suggestive of a stricture, ring, or tumor. Of note, a patient’s localization of food hang-up in the esophagus is notoriously imprecise. Approximately 30% of distal esophageal obstructions are perceived as cervical dysphagia. In such instances, the absence of concomitant symptoms generally associated with oropharyngeal dysphagia such as aspiration, nasopharyngeal regurgitation, cough, drooling, or obvious neuromuscular compromise should suggest an esophageal etiology.

Odynophagia is pain either caused by or exacerbated by swallowing. Although typically considered distinct from dysphagia, odynophagia may manifest concurrently with dysphagia. Odynophagia is more common with pill or infectious esophagitis than with reflux esophagitis and should prompt a search for these entities. When odynophagia does occur in GERD, it is likely related to an esophageal ulcer or deep erosion.

Globus sensation, alternatively labeled “globus hystericus,” is the perception of a lump or fullness in the throat that is felt irrespective of swallowing. Although such patients are frequently referred for an evaluation of dysphagia, globus sensation is often relieved by the act of swallowing. As implied by its alternative name (globus hystericus), globus sensation often occurs in the setting of anxiety or obsessive-compulsive disorders. Clinical experience teaches that it is often attributable to GERD.

Water brash is excessive salivation resulting from a vagal reflex triggered by acidification of the esophageal mucosa. This is not a common symptom. Afflicted individuals will describe the unpleasant sensation of the mouth rapidly filling with salty thin fluid, often in the setting of concomitant heartburn.

DIAGNOSTIC STUDIES

ENDOSCOPY

Endoscopy, also known as esophagogastroduodenoscopy (EGD), is the most useful test for the evaluation of the proximal gastrointestinal tract. Modern instruments produce high-quality, color images of the esophageal, gastric, and duodenal lumen. Endoscopes also have an instrumentation channel through which biopsy forceps, injection catheters for local delivery of therapeutic agents, balloon dilators, or hemostatic devices can be used. The key advantages of endoscopy over barium radiography are: (1) increased sensitivity

for the detection of mucosal lesions, (2) vastly increased sensitivity for the detection of abnormalities mainly identifiable by color such as Barrett’s metaplasia or vascular lesions, (3) the ability to obtain biopsy specimens for histologic examination of suspected abnormalities, and (4) the ability to dilate strictures during the examination. The main disadvantages of endoscopy are cost and the utilization of sedatives or anesthetics.

RADIOGRAPHY

Contrast radiography of the esophagus, stomach, and duodenum can demonstrate reflux of the contrast media, hiatal hernia, mucosal granularity, erosions, ulcerations, and strictures. The sensitivity of radiography compared with endoscopy for detecting reflux esophagitis reportedly ranges from 22–95%, with higher grades of esophagitis (i.e., ulceration or stricture) exhibiting greater detection rates. Conversely, the sensitivity of barium radiography for detecting esophageal strictures is greater than that of endoscopy, especially when the study is done in conjunction with barium-soaked bread or a 13-mm barium tablet. Barium studies also provide an assessment of esophageal function and morphology that may be undetected on endoscopy. Tracheoesophageal fistula, altered postsurgical anatomy, and extrinsic esophageal compression are conditions where radiographic imaging complements endoscopic assessment. Hypopharyngeal pathology and disorders of the cricopharyngeus muscle are better appreciated on radiographic examination than with endoscopy, particularly with rapid sequence or video fluoroscopic recording. The major shortcoming of barium radiography is that it rarely obviates the need for endoscopy. Either a positive or a negative study is usually followed by an endoscopic evaluation either to obtain biopsies, provide therapy, or clarify findings in the case of a positive examination or to add a level of certainty in the case of a negative one.

ENDOSCOPIC ULTRASOUND

Endoscopic ultrasound (EUS) instruments combine an endoscope with an ultrasound transducer to create a transmural image of the tissue surrounding the endoscope tip. The key advantage of EUS over alternative radiologic imaging techniques is much greater resolution attributable to the proximity of the ultrasound transducer to the area being examined. Available devices can provide either radial imaging (360-degree, cross-sectional) or a curved linear image that can guide fine-needle aspiration of imaged structures such as lymph nodes or tumors. Major esophageal applications of EUS are to stage esophageal cancer, to evaluate dysplasia in Barrett’s esophagus, and to assess submucosal lesions.

ESOPHAGEAL MANOMETRY

Esophageal manometry, or motility testing, entails positioning a pressure-sensing catheter within the esophagus and then observing the contractility following test swallows. The upper and lower esophageal sphincters appear as zones of high pressure that relax on swallowing, while the intersphincteric esophagus exhibits peristaltic contractions. Manometry is used to diagnose motility disorders (achalasia, diffuse esophageal spasm) and to assess peristaltic integrity prior to the surgery for reflux disease. Technologic advances have enhanced esophageal manometry as high-resolution esophageal pressure topography (Fig. 347-1). Manometry can also be combined with intraluminal impedance monitoring. Impedance recordings use a catheter with a series of paired electrodes. Esophageal luminal contents in contact with the electrodes decrease (liquid) or increase (air) the impedance signal, allowing detection of antegrade or retrograde esophageal bolus transit.

REFLUX TESTING

GERD is often diagnosed in the absence of endoscopic esophagitis, which would otherwise define the disease. This occurs in the settings of partially treated disease, an abnormally sensitive esophageal mucosa, or without obvious explanation. In such instances, reflux

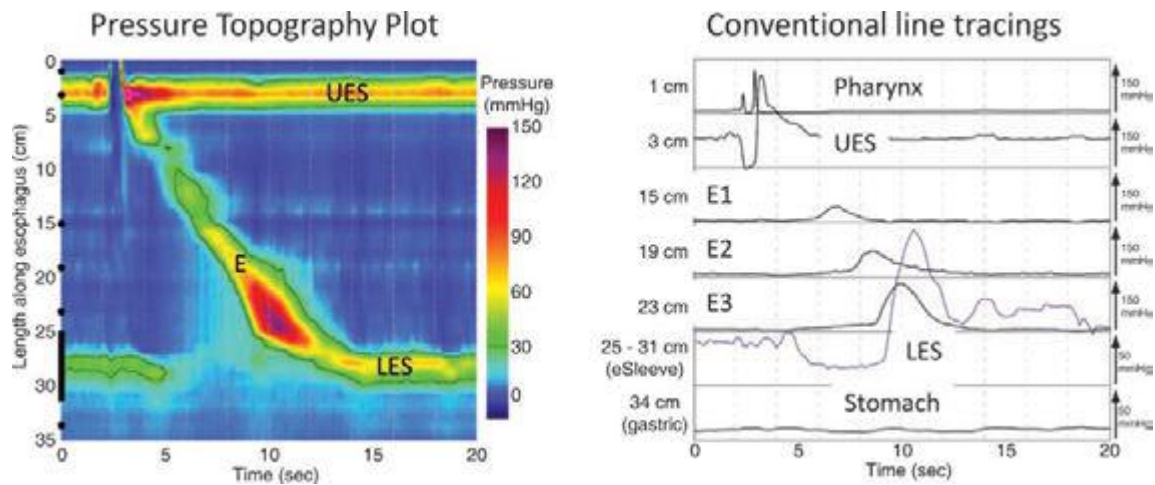


FIGURE 347-1 High-resolution esophageal pressure topography (*right*) and conventional manometry (*left*) of a normal swallow. E, esophageal body; LES, lower esophageal sphincter; UES, upper esophageal sphincter.

testing can demonstrate excessive esophageal exposure to refluxed gastric juice, the physiologic abnormality of GERD. This can be done by ambulatory 24- to 48-h esophageal pH recording using either a wireless pH-sensitive transmitter that is anchored to the esophageal mucosa or a transnasally positioned wire electrode with the tip stationed in the distal esophagus. Either way, the outcome is expressed as the percentage of the day that the pH was less than 4 (indicative of recent acid reflux), with values exceeding 5% indicative of GERD. Reflux testing is useful with atypical symptoms or an inexplicably poor response to therapy. Intraluminal impedance monitoring can be added to pH monitoring to detect reflux events irrespective of whether or not they are acidic, potentially increasing the sensitivity of the study.

STRUCTURAL DISORDERS

HIATAL HERNIA

Hiatus hernia is a herniation of viscera, most commonly the stomach, into the mediastinum through the esophageal hiatus of the diaphragm. Four types of hiatus hernia are distinguished with type I, or sliding hiatal hernia, comprising at least 95% of the overall total. A sliding hiatal hernia is one in which the gastroesophageal junction and gastric cardia translocate cephalad as a result of weakening of the phrenoesophageal ligament attaching the gastroesophageal junction to the diaphragm at the hiatus and dilatation of the diaphragmatic hiatus. The incidence of sliding hernia increases with age. True to its name, sliding hernias enlarge with increased intraabdominal pressure, swallowing, and respiration. Conceptually, sliding hernias are the result of wear and tear: increased intraabdominal pressure from abdominal obesity, pregnancy, etc., along with hereditary factors predisposing to the condition. The main significance of sliding hernias is the propensity of affected individuals to have GERD.

Types II, III, and IV hiatal hernias are all subtypes of paraesophageal hernia in which the herniation into the mediastinum includes a visceral structure other than the gastric cardia. With type II and III paraesophageal hernias, the gastric fundus also herniates with the distinction being that in type II, the gastroesophageal junction remains fixed at the hiatus, whereas type III is a combined sliding and paraesophageal hernia. With type IV hiatal hernias, viscera other than the stomach herniate into the mediastinum, most commonly the colon. With type II and III paraesophageal hernias, the stomach inverts as it herniates and large paraesophageal hernias can lead to an upside down stomach, gastric volvulus, and even strangulation of the stomach. Because of this risk, surgical repair is often advocated for large paraesophageal hernias.

RINGS AND WEBS

A lower esophageal mucosal ring, also called a *B ring*, is a thin membranous narrowing at the squamocolumnar mucosal junction (**Fig. 347-2**). Its origin is unknown, but B rings are demonstrable in about 10–15% of the general population and are usually asymptomatic. When the lumen diameter is less than 13 mm, distal rings are usually associated with episodic solid food dysphagia and are called *Schatzki rings*. Patients typically present older than 40 years, consistent with an acquired rather than congenital origin. Schatzki ring is one of the most common causes of intermittent food impaction, also known as “steakhouse syndrome” because meat is a typical instigator. Symptomatic rings are easily treated by dilation.

Web-like constrictions higher in the esophagus can be of congenital or inflammatory origin. Asymptomatic cervical esophageal webs are demonstrated in about 10% of people and typically originate along the anterior aspect of the esophagus. When circumferential, they can

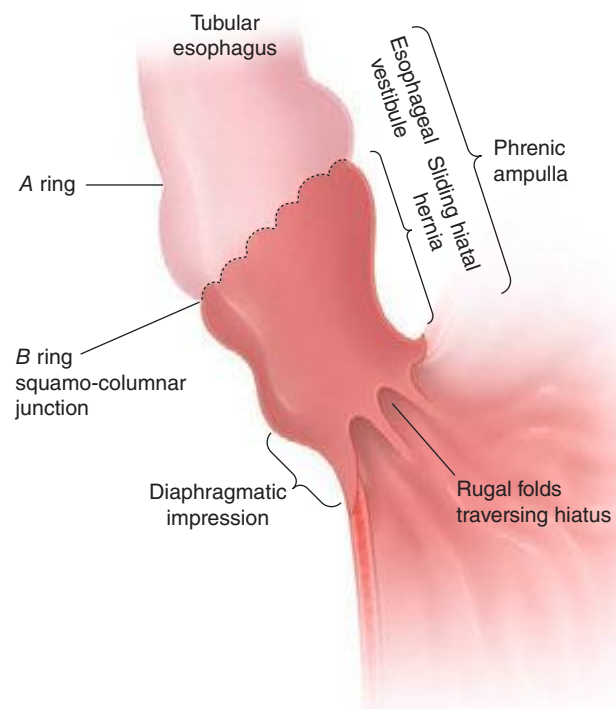


FIGURE 347-2 Radiographic anatomy of the gastroesophageal junction.

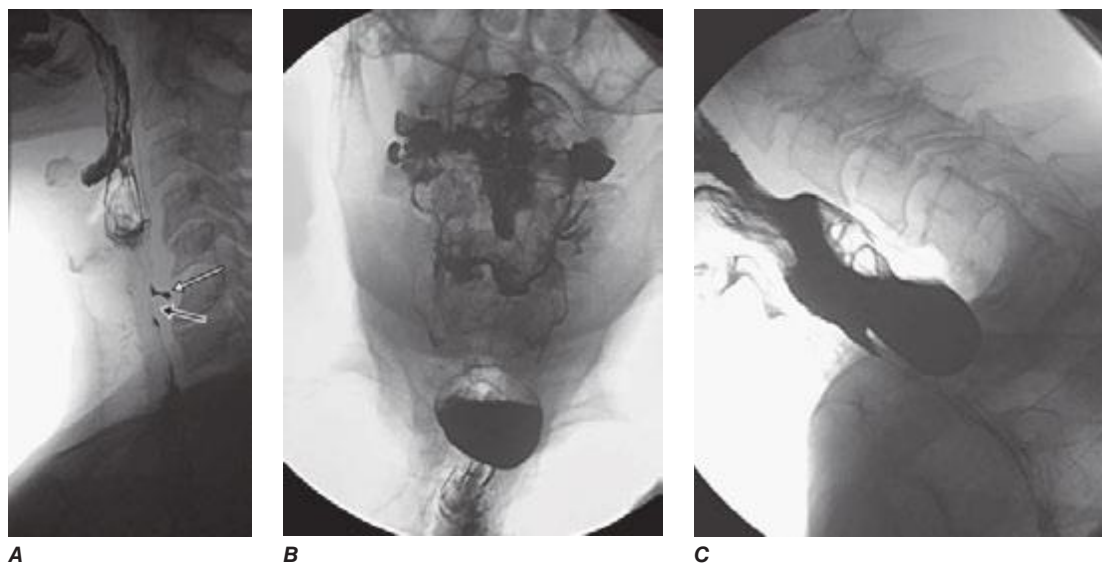


FIGURE 347-3 Examples of small (A) and large (B, C) Zenker's diverticula arising from Killian's triangle in the distal hypopharynx. Smaller diverticula are evident only during the swallow, whereas larger ones retain food and fluid.

cause intermittent dysphagia to solids similar to Schatzki rings and are similarly treated with dilatation. The combination of symptomatic proximal esophageal webs and iron-deficiency anemia in middle-aged women constitutes Plummer-Vinson syndrome.

DIVERTICULA

Esophageal diverticula are categorized by location with the most common being epiphrenic, hypopharyngeal (Zenker's), and midesophageal. Epiphrenic and Zenker's diverticula are false diverticula involving herniation of the mucosa and submucosa through the muscular layer of the esophagus. These lesions result from increased intraluminal pressure associated with distal obstruction. In the case of Zenker's, the obstruction is a stenotic cricopharyngeus muscle (upper esophageal sphincter), and the hypopharyngeal herniation most commonly occurs in an area of natural weakness proximal to the cricopharyngeus known as *Killian's triangle* (Fig. 347-3). Small Zenker's diverticula are usually asymptomatic, but when they enlarge sufficiently to retain food and saliva they can be associated with dysphagia, halitosis, and aspiration. Treatment is by surgical diverticulectomy and cricopharyngeal myotomy or a marsupialization procedure in which an endoscopic stapling device is used to divide the cricopharyngeus.

Epiphrenic diverticula are usually associated with achalasia or a distal esophageal stricture. Midesophageal diverticula may be caused by traction from adjacent inflammation (classically tuberculosis) in which case they are true diverticula involving all layers of the esophageal wall, or by pulsion associated with esophageal motor disorders. Midesophageal and epiphrenic diverticula are usually asymptomatic until they enlarge sufficiently to retain food and cause dysphagia and regurgitation. Symptoms attributable to the diverticula tend to correlate more with the underlying esophageal disorder than the size of the diverticula. Large diverticula can be removed surgically, usually in conjunction with a myotomy if the underlying cause is achalasia. Diffuse intramural esophageal diverticulosis is a rare entity that results from dilatation of the excretory ducts of submucosal esophageal glands (Fig. 347-4). Esophageal candidiasis and proximal esophageal strictures are commonly found in association with this disorder.

TUMORS

Esophageal cancer occurs in about 4.5:100,000 people in the United States with the associated mortality being only slightly less at 4.4:100,000. It is about 10 times less common than colorectal cancer but kills about one-quarter as many patients. These statistics



FIGURE 347-4 Intramural esophageal pseudodiverticulosis associated with chronic obstruction. Invaginations of contrast into the esophageal wall outline deep esophageal glands.

emphasize both the rarity and lethality of esophageal cancer. One notable trend is the shift of dominant esophageal cancer type from squamous cell to adenocarcinoma, strongly linked to reflux disease and Barrett's metaplasia. Other distinctions between cell types are the

predilection for adenocarcinoma to affect the distal esophagus in white males and squamous cell to affect the more proximal esophagus in black males with the added risk factors of smoking, alcohol consumption, caustic injury, and human papilloma virus infection (Chap. 109).

The typical presentation of esophageal cancer is of progressive solid food dysphagia and weight loss. Associated symptoms may include odynophagia, iron deficiency, and, with midesophageal tumors, hoarseness from left recurrent laryngeal nerve injury. Generally, these are indications of locally invasive or even metastatic disease manifest by tracheoesophageal fistulas and vocal cord paralysis. Even when detected as a small lesion, esophageal cancer has poor survival because of the abundant esophageal lymphatics leading to regional lymph node metastases.

Benign esophageal tumors are uncommon and usually discovered incidentally. In decreasing frequency of occurrence, cell types include leiomyoma, fibrovascular polyps, squamous papilloma, granular cell tumors, lipomas, neurofibromas, and inflammatory fibroid polyps. These generally become symptomatic only when they are associated with dysphagia and merit removal only under the same circumstances.

CONGENITAL ANOMALIES

The most common congenital esophageal anomaly is esophageal atresia, occurring in about 1 in 5000 live births. Atresia can occur in several permutations, the common denominator being developmental failure of fusion between the proximal and distal esophagus associated with a tracheoesophageal fistula, most commonly with the distal segment excluded. Alternatively, there can be an H-type configuration in which esophageal fusion has occurred, but with a tracheoesophageal fistula. Esophageal atresia is usually recognized and corrected surgically within the first few days of life. Later life complications include dysphagia from anastomotic strictures or absent peristalsis and reflux, which can be severe. Less common developmental anomalies include congenital esophageal stenosis, webs, and duplications.

Dysphagia can also result from congenital abnormalities that cause extrinsic compression of the esophagus. In dysphagia lusoria, the esophagus is compressed by an aberrant right subclavian artery arising from the descending aorta and passing behind the esophagus. Alternatively vascular rings may surround and constrict the esophagus.

Heterotopic gastric mucosa, also known as an esophageal inlet patch, is a focus of gastric type epithelium in the proximal cervical esophagus; the estimated prevalence is 4.5%. The inlet patch is thought to result from incomplete replacement of embryonic columnar epithelium with squamous epithelium. The majority of inlet patches are asymptomatic, but acid production can occur as most contain fundic type gastric epithelium with parietal cells.

ESOPHAGEAL MOTILITY DISORDERS

Esophageal motility disorders are diseases attributable to esophageal neuromuscular dysfunction commonly associated with dysphagia, chest pain, or heartburn. The major entities are achalasia, diffuse esophageal spasm (DES), and GERD. Motility disorders can also be secondary to broader disease processes as is the case with pseudoachalasia, Chagas' disease, and scleroderma. Not included in this discussion are diseases affecting the pharynx and proximal esophagus, the impairment of which is almost always part of a more global neuromuscular disease process.

ACHALASIA

Achalasia is a rare disease caused by loss of ganglion cells within the esophageal myenteric plexus with a population incidence of about 1:100,000 and usually presenting between age 25 and 60. With longstanding disease, aganglionosis is noted. The disease involves both excitatory (cholinergic) and inhibitory (nitric oxide) ganglionic neurons. Functionally, inhibitory neurons mediate deglutitive lower esophageal sphincter (LES) relaxation and the sequential propagation

of peristalsis. Their absence leads to impaired deglutitive LES relaxation and absent peristalsis. Increasing evidence suggests that the ultimate cause of ganglion cell degeneration in achalasia is an autoimmune process attributable to a latent infection with human herpes simplex virus 1 combined with genetic susceptibility.

Long-standing achalasia is characterized by progressive dilatation and sigmoid deformity of the esophagus with hypertrophy of the LES. Clinical manifestations may include dysphagia, regurgitation, chest pain, and weight loss. Most patients report solid and liquid food dysphagia. Regurgitation occurs when food, fluid, and secretions are retained in the dilated esophagus. Patients with advanced achalasia are at risk for bronchitis, pneumonia, or lung abscess from chronic regurgitation and aspiration. Chest pain is frequent early in the course of achalasia, thought to result from esophageal spasm. Patients describe a squeezing, pressure-like retrosternal pain, sometimes radiating to the neck, arms, jaw, and back. Paradoxically, some patients complain of heartburn that may be a chest pain equivalent. Treatment of achalasia is less effective in relieving chest pain than it is in relieving dysphagia or regurgitation.

The differential diagnosis of achalasia includes DES, Chagas' disease, and pseudoachalasia. Chagas' disease is endemic in areas of central Brazil, Venezuela, and northern Argentina and spread by the bite of the reduviid (kissing) bug that transmits the protozoan, *Trypanosoma cruzi*. The chronic phase of the disease develops years after infection and results from destruction of autonomic ganglion cells throughout the body, including the heart, gut, urinary tract, and respiratory tract. Tumor infiltration, most commonly seen with carcinoma in the gastric fundus or distal esophagus, can mimic idiopathic achalasia. The resultant "pseudoachalasia" accounts for up to 5% of suspected cases and is more likely with advanced age, abrupt onset of symptoms (<1 year), and weight loss. Hence, endoscopy is a necessary part of the evaluation of achalasia. When the clinical suspicion for pseudoachalasia is high and endoscopy nondiagnostic, computed tomography (CT) scanning or EUS may be of value. Rarely, pseudoachalasia can result from a paraneoplastic syndrome with circulating antineuronal antibodies.

Achalasia is diagnosed by barium swallow x-ray and/or esophageal manometry; endoscopy has a relatively minor role other than to exclude pseudoachalasia. The barium swallow x-ray appearance is of a dilated esophagus with poor emptying, an air-fluid level, and tapering at the LES giving it a beak-like appearance (Fig. 347-5). Occasionally, an epiphrenic diverticulum is observed. In long-standing achalasia, the esophagus may assume a sigmoid configuration. The diagnostic criteria for achalasia with esophageal manometry are impaired LES relaxation and absent peristalsis. High-resolution manometry has somewhat advanced this diagnosis; three subtypes of achalasia are differentiated based on the pattern of pressurization in the nonperistaltic esophagus (Fig. 347-6). Because manometry identifies early disease before esophageal dilatation and food retention, it is the most sensitive diagnostic test.

There is no known way of preventing or reversing achalasia. Therapy is directed at reducing LES pressure so that gravity and esophageal pressurization promote esophageal emptying. Peristalsis rarely, if ever, recovers. However, in many instances, remnants of peristalsis masked by esophageal pressurization and dilatation prior to therapy are demonstrable following effective treatment. LES pressure can be reduced by pharmacologic therapy, pneumatic balloon dilatation, or surgical myotomy. No large, controlled trials of the therapeutic alternatives exist, and the optimal approach is debated. Pharmacologic therapies are relatively ineffective but are often used as temporizing therapies. Nitrates or calcium channel blockers are administered before eating, advising caution because of their effects on blood pressure. Botulinum toxin, injected into the LES under endoscopic guidance, inhibits acetylcholine release from nerve endings and improves dysphagia in about 66% of cases for at least 6 months. Sildenafil and alternative phosphodiesterase inhibitors effectively decrease LES pressure, but practicalities limit their clinical use in achalasia.

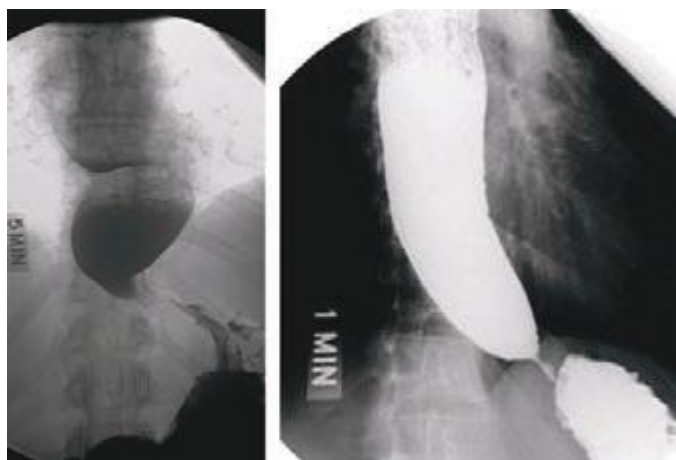


FIGURE 347-5 Achalasia with esophageal dilatation, tapering at the gastroesophageal junction, and an air-fluid level within the esophagus. The example on the left shows sigmoid deformity with very advanced disease.

The only durable therapies for achalasia are pneumatic dilatation and Heller myotomy. Pneumatic dilatation, with a reported efficacy ranging from 32–98%, is an endoscopic technique using a noncompliant, cylindrical balloon dilator positioned across the LES and inflated to a diameter of 3–4 cm. The major complication is perforation with a reported incidence of 0.5–5%. The most common surgical procedure for achalasia is laparoscopic Heller myotomy, usually performed in conjunction with an antireflux procedure (partial fundoplication); good to excellent results are reported in 62–100% of cases. A European randomized controlled trial demonstrated an equivalent response rate of approximately 90% for both pneumatic dilation and laparoscopic Heller myotomy at 2-year follow-up. Occasionally, patients with advanced disease fail to respond to pneumatic dilatation or Heller myotomy. In such refractory cases, esophageal resection with gastric pull-up or interposition of a segment of transverse colon may be the only option other than gastrostomy feeding.

An endoscopic approach to LES myotomy has been introduced, referred to as per oral esophageal myotomy. This technique involves the creation of a tunnel within the esophageal wall through which the circular muscle of the LES and distal esophagus are transected with electrocautery. Short-term studies of efficacy have been favorable. Potential advantages over the conventional laparoscopic approach include avoidance of surgical disruption of the diaphragmatic hiatus and more rapid recovery.

In untreated or inadequately treated achalasia, esophageal dilatation predisposes to stasis esophagitis. Prolonged stasis esophagitis is the likely explanation for the association between achalasia and esophageal squamous cell cancer. Tumors develop after years of achalasia, usually in the setting of a greatly dilated esophagus with the overall squamous cell cancer risk increased 17-fold compared to controls.

DIFFUSE ESOPHAGEAL SPASM (DES)

DES is manifested by episodes of dysphagia and chest pain attributable to abnormal esophageal contractions with normal deglutitive LES relaxation. Beyond that, there is little consensus. The pathophysiology and natural history of DES are ill defined. Radiographically, DES has been characterized by tertiary contractions or a “corkscrew esophagus” (Fig. 347-7), but in many instances, these abnormalities are actually indicative of achalasia. Manometrically, a variety of defining features have been proposed including uncoordinated (“spastic”) activity in the distal esophagus, spontaneous and repetitive contractions, or high-amplitude and prolonged contractions. The current consensus, derived from high-resolution manometry studies, is to define spasm by the occurrence of contractions in the distal esophagus with short latency relative to the time of the pharyngeal contraction, a dysfunction indicative of impairment of inhibitory myenteric plexus neurons.

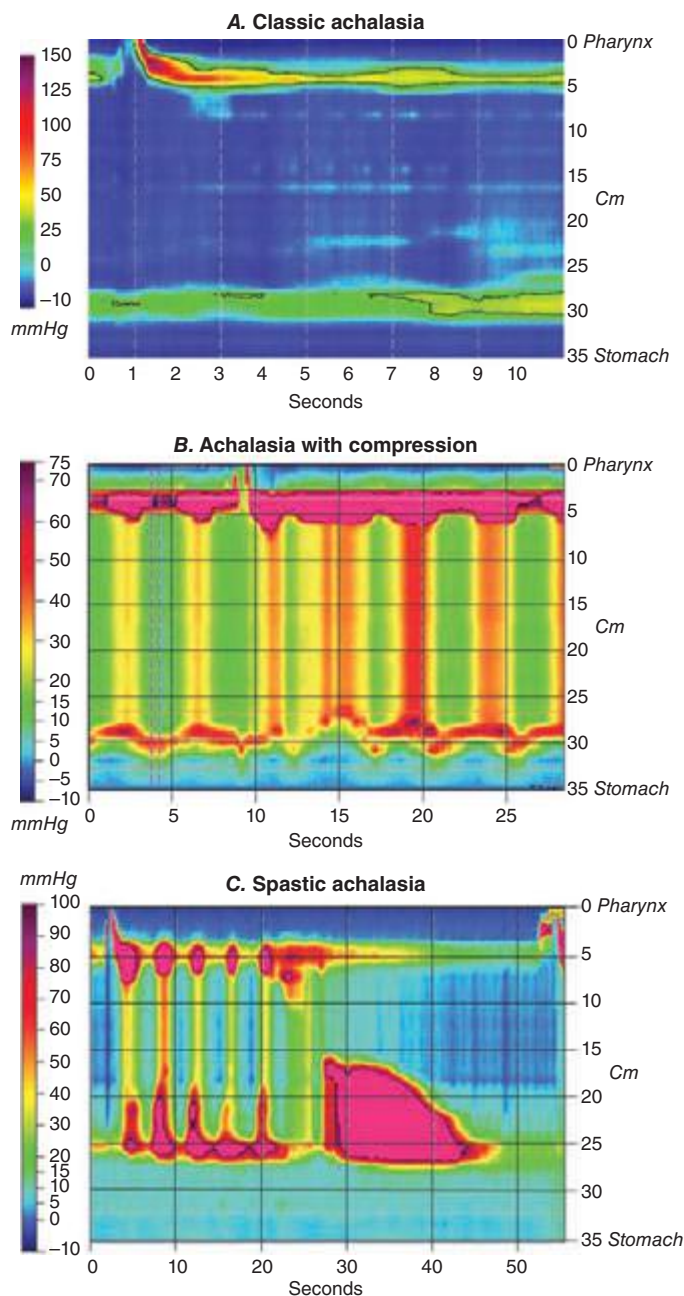


FIGURE 347-6 Three subtypes of achalasia: classic (A), with esophageal compression (B), and spastic achalasia (C) imaged with pressure topography. All are characterized by impaired lower esophageal sphincter (LES) relaxation and absent peristalsis. However, classic achalasia has minimal pressurization of the esophageal body, whereas substantial fluid pressurization is observed in achalasia with esophageal compression, and spastic esophageal contractions are observed with spastic achalasia.

When defined in this restrictive fashion (Fig. 347-8), DES is actually much less common than achalasia.

Esophageal chest pain closely mimics angina pectoris. Features suggesting esophageal pain include pain that is nonexertional, prolonged, interrupts sleep, meal-related, relieved with antacids, and accompanied by heartburn, dysphagia, or regurgitation. However, all of these features exhibit overlap with cardiac pain, which still must be the primary consideration. Furthermore, even within the spectrum of esophageal diseases, both chest pain and dysphagia are also characteristic of peptic or infectious esophagitis. Only after these more common entities have been excluded by evaluation and/or treatment should a diagnosis of DES be pursued.

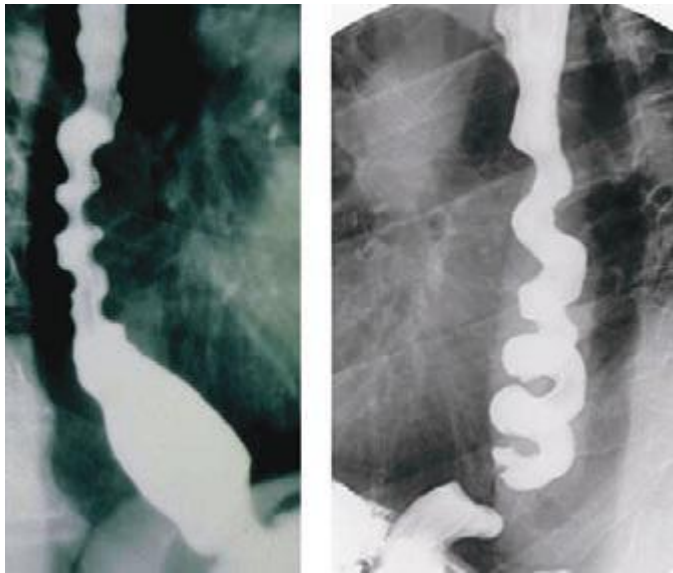


FIGURE 347-7 Diffuse esophageal spasm. The characteristic “corkscrew” esophagus results from spastic contraction of the circular muscle in the esophageal wall; more precisely, this is actually a helical array of muscle. These findings are also seen with spastic achalasia.

Although the defining criteria are in flux, DES is diagnosed by manometry. Endoscopy is useful to identify alternative structural and inflammatory lesions that may cause chest pain. Radiographically, a “corkscrew esophagus,” “rosary bead esophagus,” pseudodiverticula, or curling can be indicative of DES, but these are also found with spastic achalasia. Given these vagaries of defining DES, and the resultant heterogeneity of patients identified for inclusion in therapeutic trials, it is not surprising that trial results have been disappointing. Only small, uncontrolled trials exist, reporting response to nitrates, calcium channel blockers, hydralazine, botulinum toxin, and anxiolytics. The only controlled trial showing efficacy was with an anxiolytic. Surgical therapy (long myotomy or even esophagectomy) should be considered only with severe weight loss or unbearable pain. These indications are extremely rare.

NONSPECIFIC MANOMETRIC FINDINGS

Manometric studies done to evaluate chest pain and/or dysphagia often report minor abnormalities (e.g., hypertensive or hypotensive

peristalsis, hypertensive LES) that are insufficient to diagnose either achalasia or DES. These findings are of unclear significance. Reflux and psychiatric diagnoses, particularly anxiety and depression, are common among such individuals. A lower visceral pain threshold and symptoms of irritable bowel syndrome are noted in more than half of such patients. Consequently, therapy for these individuals should either target the most common esophageal disorder, GERD, or more global conditions such as depression or somatization neurosis that are found to be coexistent.

GASTROESOPHAGEAL REFLUX DISEASE (GERD)

The current conception of GERD is to encompass a family of conditions with the commonality that they are caused by gastroesophageal reflux resulting in either troublesome symptoms or an array of potential esophageal and extraesophageal manifestations. It is estimated that 15% of adults in the United States are affected by GERD, although such estimates are based only on population studies of self-reported chronic heartburn. With respect to the esophagus, the spectrum of injury includes esophagitis, stricture, Barrett’s esophagus, and adenocarcinoma (Fig. 347-9). Of particular concern is the rising incidence of esophageal adenocarcinoma, an epidemiologic trend that parallels the increasing incidence of GERD. There were about 8000 incident cases of esophageal adenocarcinoma in the United States in 2013 (half of all esophageal cancers); it is estimated that this disease burden has increased two- to sixfold in the last 20 years.

PATHOPHYSIOLOGY

The best-defined subset of GERD patients, albeit a minority overall, have esophagitis. Esophagitis occurs when refluxed gastric acid and pepsin cause necrosis of the esophageal mucosa causing erosions and ulcers. Note that some degree of gastroesophageal reflux is normal, physiologically intertwined with the mechanism of belching (transient LES relaxation), but esophagitis results from excessive reflux, often accompanied by impaired clearance of the refluxed gastric juice. Restricting reflux to that which is physiologically intended depends on the anatomic and physiologic integrity of the esophagogastric junction, a complex sphincter comprised of both the LES and the surrounding crural diaphragm. Three dominant mechanisms of esophagogastric junction incompetence are recognized: (1) transient LES relaxations (a vagovagal reflex in which LES relaxation is elicited by gastric distention), (2) LES hypotension, or (3) anatomic distortion of the esophagogastric junction inclusive of hiatus hernia. Of note, the third factor, esophagogastric junction anatomic disruption, is both significant unto itself and also because it interacts with the first two mechanisms. Transient LES relaxations account for about 90% of reflux in normal subjects or GERD patients without hiatus hernia, but patients with hiatus hernia have a more heterogeneous mechanistic profile. Factors tending to exacerbate reflux regardless of mechanism are abdominal obesity, pregnancy, gastric hypersecretory states, delayed gastric emptying, disruption of esophageal peristalsis, and gluttony.

After acid reflux, peristalsis returns the refluxed fluid to the stomach and acid clearance is completed by titration of the residual acid by bicarbonate contained in swallowed saliva. Consequently, two causes of prolonged acid clearance are impaired peristalsis and reduced salivation. Impaired peristaltic emptying can be attributable to disrupted peristalsis or superimposed reflux associated with a hiatal hernia. With superimposed reflux, fluid retained within a sliding hiatal hernia refluxes back into the

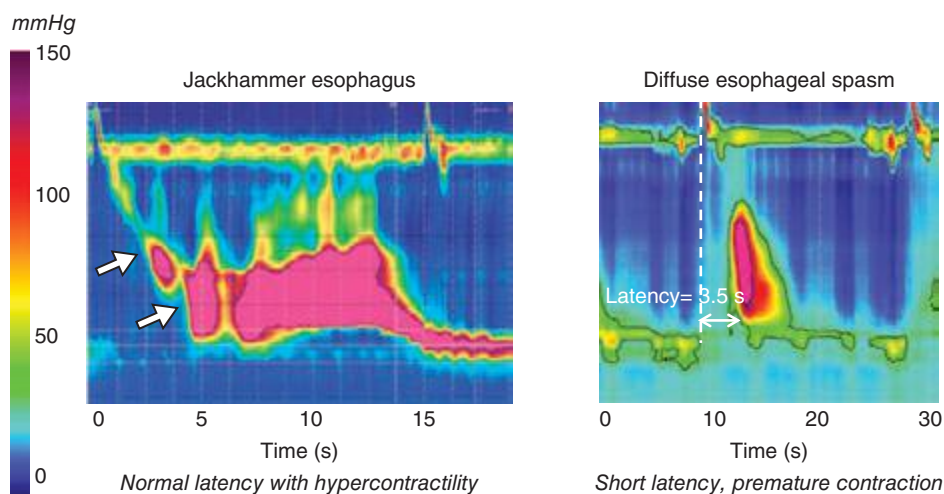


FIGURE 347-8 Esophageal pressure topography of the two major variants of esophageal spasm: jackhammer esophagus (left) and diffuse esophageal spasm (right). Jackhammer esophagus is defined by the extraordinarily vigorous and repetitive contractions with normal peristaltic onset and normal latency of the contraction. Diffuse esophageal spasm is similar but primarily defined by a short latency (premature) contraction.

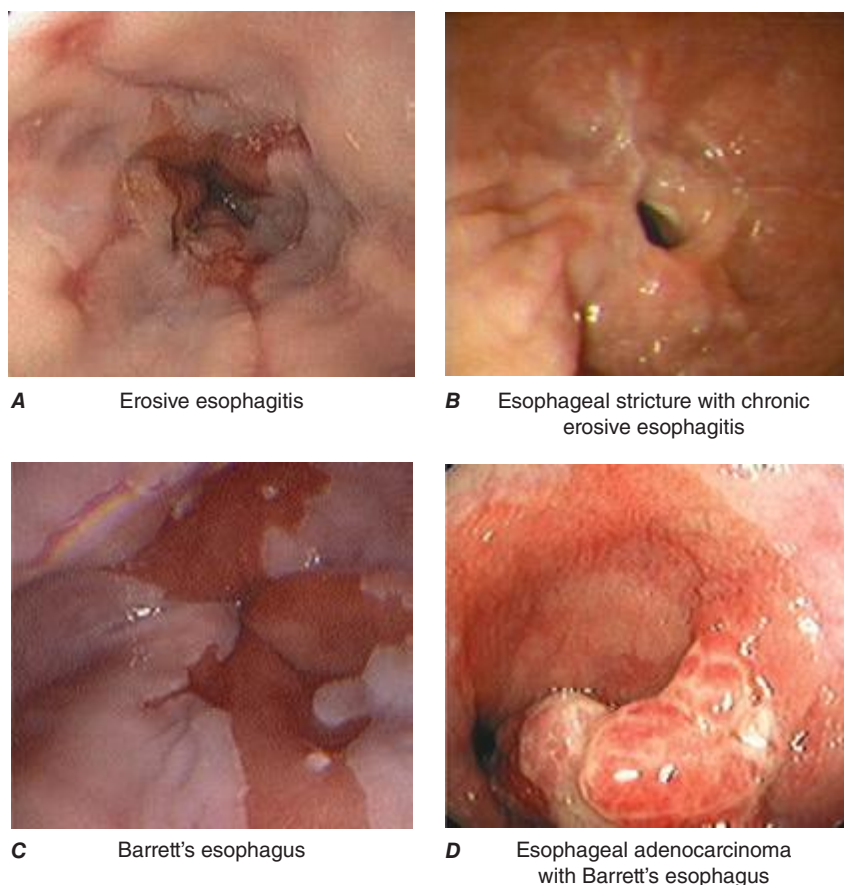


FIGURE 347-9 Endoscopic appearance of (A) peptic esophagitis, (B) a peptic stricture, (C) Barrett's metaplasia, and (D) adenocarcinoma developing within an area of Barrett's esophagus.

esophagus during swallow-related LES relaxation, a phenomenon that does not normally occur.

Inherent in the pathophysiologic model of GERD is that gastric juice is harmful to the esophageal epithelium. However, gastric acid hypersecretion is usually not a dominant factor in the development of esophagitis. An obvious exception is with Zollinger-Ellison syndrome, which is associated with severe esophagitis in about 50% of patients. Another caveat is with chronic *Helicobacter pylori* gastritis, which may have a protective effect by inducing atrophic gastritis with concomitant hypoacidity. Pepsin, bile, and pancreatic enzymes within gastric secretions can also injure the esophageal epithelium, but their noxious properties are either lessened without an acidic environment or dependent on acidity for activation. Bile warrants attention because it persists in refluxate despite acid-suppressing medications. Bile can transverse the cell membrane, imparting severe cellular injury in a weakly acidic environment, and has also been invoked as a cofactor in the pathogenesis of Barrett's metaplasia and adenocarcinoma. Hence, the causticity of gastric refluxate extends beyond hydrochloric acid.

SYMPTOMS

Heartburn and regurgitation are the typical symptoms of GERD. Somewhat less common are dysphagia and chest pain. In each case, multiple potential mechanisms for symptom genesis operate that extend beyond the basic concepts of mucosal erosion and activation of afferent sensory nerves. Specifically, hypersensitivity and functional pain are increasingly recognized as cofactors. Nonetheless, the dominant clinical strategy is empirical treatment with acid inhibitors, reserving further evaluation for those who fail to respond. Important exceptions to this are patients with chest pain or persistent dysphagia, each of which may be indicative of more morbid conditions. With chest pain, cardiac disease must be carefully considered. In the case of persistent dysphagia, chronic reflux can lead to the development of

a peptic stricture or adenocarcinoma, each of which benefits from early detection and/or specific therapy.

Extraesophageal syndromes with an established association to GERD include chronic cough, laryngitis, asthma, and dental erosions. A multitude of other conditions including pharyngitis, chronic bronchitis, pulmonary fibrosis, chronic sinusitis, cardiac arrhythmias, sleep apnea, and recurrent aspiration pneumonia have proposed associations with GERD. However, in both cases, it is important to emphasize the word *association* as opposed to *causation*. In many instances, the disorders likely coexist because of shared pathogenetic mechanisms rather than strict causality. Potential mechanisms for extraesophageal GERD manifestations are either regurgitation with direct contact between the refluxate and supraesophageal structures or via a vagovagal reflex wherein reflux activation of esophageal afferent nerves triggers efferent vagal reflexes such as bronchospasm, cough, or arrhythmias.

DIFFERENTIAL DIAGNOSIS

Although generally quite characteristic, symptoms from GERD need to be distinguished from symptoms related to infectious, pill, or eosinophilic esophagitis, peptic ulcer disease, dyspepsia, biliary colic, coronary artery disease, and esophageal motility disorders. It is especially important that coronary artery disease be given early consideration because of its potentially lethal implications. The remaining elements of the differential diagnosis can be addressed by endoscopy, upper gastrointestinal series, or biliary tract ultrasonography as appropriate. The distinction among etiologies of esophagitis is usually easily made by endoscopy with mucosal biopsies, which are necessary to evaluate for infection or eosinophilic inflammation. In terms of endoscopic appearance, infectious esophagitis is diffuse and tends to involve the proximal esophagus far more frequently than does reflux esophagitis. The ulcerations seen in peptic esophagitis are usually solitary and distal, whereas infectious ulcerations are punctate and diffuse. Eosinophilic esophagitis characteristically exhibits multiple esophageal rings, linear furrows, or white punctate exudate. Esophageal ulcerations from pill esophagitis are usually singular and deep at points of luminal narrowing, especially near the carina, with sparing of the distal esophagus.

COMPLICATIONS

The complications of GERD are related to chronic esophagitis (bleeding and stricture) and the relationship between GERD and esophageal adenocarcinoma. However, both esophagitis and peptic strictures have become increasingly rare in the era of potent antisecretory medications. Conversely, the most severe histologic consequence of GERD is Barrett's metaplasia with the associated risk of esophageal adenocarcinoma, and the incidence of these lesions has increased, not decreased, in the era of potent acid suppression. Barrett's metaplasia, endoscopically recognized by tongues of reddish mucosa extending proximally from the gastroesophageal junction (Fig. 347-9) or histopathologically by the finding of specialized columnar metaplasia, is associated with a substantially increased risk for development of esophageal adenocarcinoma.

Barrett's metaplasia can progress to adenocarcinoma through the intermediate stages of low- and high-grade dysplasia (Fig. 347-10). Owing to this risk, areas of Barrett's and especially any included areas of mucosal irregularity should be extensively biopsied. The rate of cancer development is estimated at 0.1–0.3% per year, but vagaries in definitional criteria and of the extent of Barrett's metaplasia requisite to establish the diagnosis have contributed to variability and inconsistency in this risk assessment. The group at greatest risk is obese white males in their sixth decade of life. However, despite common practice, the utility of endoscopic screening and surveillance programs intended

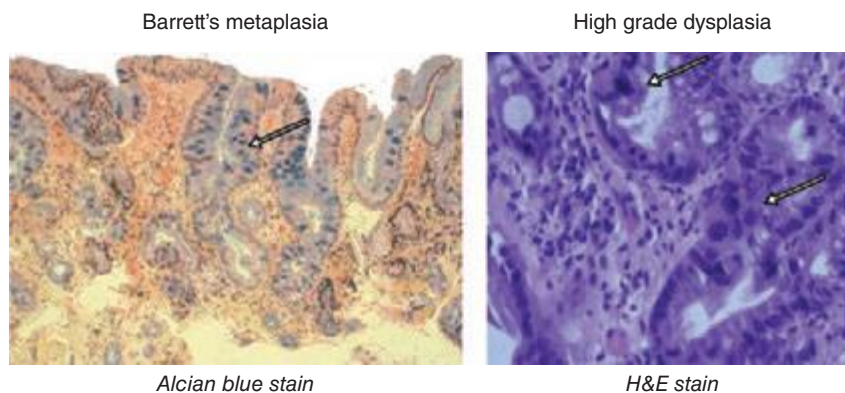


FIGURE 347-10 Histopathology of Barrett's metaplasia and Barrett's with high-grade dysplasia. H&E, hematoxylin and eosin.

to control the adenocarcinoma risk has not been established. Also of note, no high-level evidence confirms that aggressive antisecretory therapy or antireflux surgery causes regression of Barrett's esophagus or prevents adenocarcinoma.

Although the management of Barrett's esophagus remains controversial, the finding of dysplasia in Barrett's, particularly high-grade dysplasia, mandates further intervention. In addition to the high rate of progression to adenocarcinoma, there is also a high prevalence of unrecognized coexisting cancer with high-grade dysplasia. Nonetheless, treatment remains controversial. Esophagectomy, intensive endoscopic surveillance, and mucosal ablation have all been advocated. Currently, esophagectomy is the gold standard treatment for high-grade dysplasia in an otherwise healthy patient with minimal surgical risk. However, esophagectomy has a mortality ranging from 3–10%, along with substantial morbidity. That, along with increasing evidence of the effectiveness of endoscopic therapy with purpose-built radiofrequency ablation devices, has led many to favor this therapy as a preferable management strategy.

TREATMENT GASTROESOPHAGEAL REFLUX DISEASE (GERD)

Lifestyle modifications are routinely advocated as GERD therapy. Broadly speaking, these fall into three categories: (1) avoidance of foods that reduce LES pressure, making them "refluxogenic" (these commonly include fatty foods, alcohol, spearmint, peppermint, tomato-based foods, and possibly coffee and tea); (2) avoidance of acidic foods that are inherently irritating; and (3) adoption of behaviors to minimize reflux and/or heartburn. In general, minimal evidence supports the efficacy of these measures. However, clinical experience dictates that subsets of patients are benefitted by specific recommendations, based on their unique history and symptom profile. A patient with sleep disturbance from nighttime heartburn is likely to benefit from elevation of the head of the bed and avoidance of eating before retiring, but those recommendations are superfluous for a patient without nighttime symptoms. The most broadly applicable recommendation is for weight reduction. Even though the benefit with respect to reflux cannot be assured, the strong epidemiologic relationship between body mass index and GERD and the secondary health gains of weight reduction are beyond dispute.

The dominant pharmacologic approach to GERD management is with inhibitors of gastric acid secretion, and abundant data support the effectiveness of this approach. Pharmacologically reducing the acidity of gastric juice does not prevent reflux, but it ameliorates reflux symptoms and allows esophagitis to heal. The hierarchy of effectiveness among pharmaceuticals parallels their antisecretory potency. Proton pump inhibitors (PPIs) are more efficacious than histamine₂ receptor antagonists (H₂RAs), and both are superior to placebo. No major differences exist among PPIs, and only modest gain is achieved by increased dosage.

Paradoxically, the perceived frequency and severity of heartburn correlate poorly with the presence or severity of esophagitis. When GERD treatments are assessed in terms of resolving heartburn, both efficacy and differences among pharmaceuticals are less clear-cut than with the objective of healing esophagitis. Although the same overall hierarchy of effectiveness exists, observed efficacy rates are lower and vary widely, likely reflecting patient heterogeneity.

Reflux symptoms tend to be chronic, irrespective of esophagitis. Thus, a common management strategy is indefinite treatment with PPIs or H₂RAs as necessary for symptom control. The side effects of PPI therapy are generally minimal. Vitamin B₁₂ and iron absorption may be compromised and susceptibility to enteric infections, particularly *Clostridium difficile* colitis, increased with treatment. Population studies have also suggested a slight increased risk of bone fracture with chronic PPI use suggesting an impairment of calcium absorption, but prospective studies have failed to corroborate this. Nonetheless, as with any medication, PPI dosage should be minimized to that necessary for the clinical indication.

Laparoscopic Nissen fundoplication, wherein the proximal stomach is wrapped around the distal esophagus to create an antireflux barrier, is a surgical alternative to the management of chronic GERD. Just as with PPI therapy, evidence on the utility of fundoplication is strongest for treating esophagitis, and controlled trials suggest similar efficacy to PPI therapy. However, the benefits of fundoplication must be weighed against potential deleterious effects, including surgical morbidity and mortality, postoperative dysphagia, failure or breakdown requiring reoperation, an inability to belch, and increased bloating, flatulence, and bowel symptoms after surgery.

EOSINOPHILIC ESOPHAGITIS

Eosinophilic esophagitis (EoE) is increasingly recognized in adults and children around the world. Current prevalence estimates identified 4–6 cases per 10,000 with a predilection for white males. The increasing prevalence of EoE is attributable to a combination of an increasing incidence and a growing recognition of the condition. There is also an incompletely understood, but important, overlap between EoE and GERD that confuses diagnosis of the disease.

EoE is diagnosed based on the combination of typical esophageal symptoms and esophageal mucosal biopsies demonstrating squamous epithelial eosinophil-predominant inflammation. Alternative etiologies of esophageal eosinophilia include GERD, drug hypersensitivity, connective tissue disorders, hypereosinophilic syndrome, and infection. Current evidence indicates that EoE is an immunologic disorder induced by antigen sensitization in susceptible individuals. Dietary factors play an important role in both the pathogenesis and treatment of EoE. Aeroallergens may also contribute, but the evidence is weaker. The natural history of EoE is unclear, but an increased risk of esophageal stricture development paralleling the duration of untreated disease has been noted.

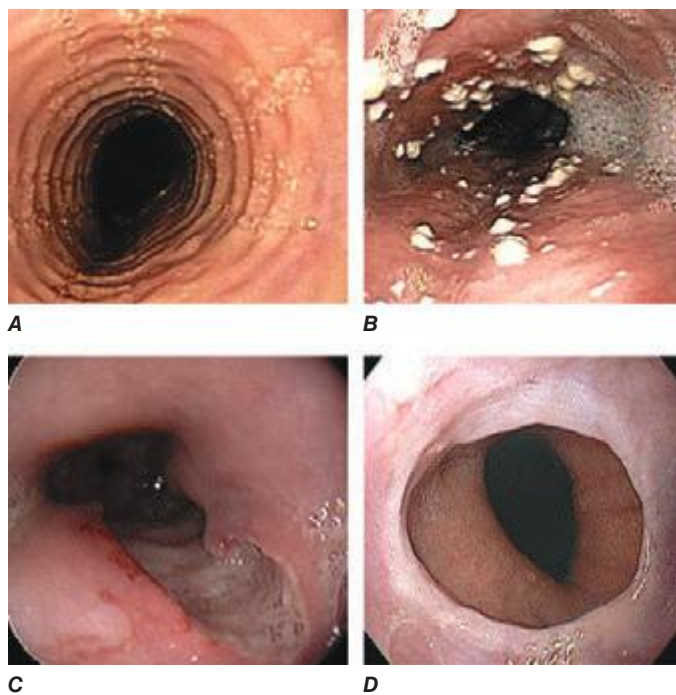


FIGURE 347-11 Endoscopic features of (A) eosinophilic esophagitis (EoE), (B) *Candida* esophagitis, (C) giant ulcer associated with HIV, (D) and a Schatzki ring.

EoE should be strongly considered in children and adults with dysphagia and esophageal food impactions. In preadolescent children, symptom presentations of EoE include chest or abdominal pain, nausea, vomiting, and food aversion. Other symptoms in adults may include atypical chest pain and heartburn, particularly heartburn that is refractory to PPI therapy. An atopic history of food allergy, asthma, eczema, or allergic rhinitis is present in the majority of patients. Peripheral blood eosinophilia is demonstrable in up to 50% of patients, but the specificity of this finding is problematic in the setting of concomitant atopy. The characteristic endoscopic esophageal findings are loss of vascular markings (edema), multiple esophageal rings, longitudinally oriented furrows, and punctate exudate (Fig. 347-11). Histologic confirmation is made with the demonstration of esophageal mucosal eosinophilia (greatest density ± 15 eosinophils per high-power field) (Fig. 347-12). Complications of EoE include esophageal stricture, narrow-caliber esophagus, food impaction, and esophageal perforation.

The goals of EoE management are symptom control and the prevention of complications. Once esophageal eosinophilia is demonstrated, patients typically undergo a trial of PPI therapy as a practical means of excluding a contribution of GERD to the esophageal mucosal inflammation. PPI-responsive esophageal eosinophilia, characterized by elimination of mucosal eosinophilia, occurs in 30–50% of cases of suspected EoE. Patients with persistent symptoms and eosinophilic inflammation following PPI therapy are subsequently considered for EoE treatments such as elimination diets or swallowed topical glucocorticoids. Elemental formula diets are a highly effective therapy that have primarily been studied in children but are limited by palatability. Notably, allergy testing by means of either serum IgE or skin prick testing has demonstrated poor sensitivity and specificity in the identification of foods that incite the esophageal inflammatory response. Allergy testing combining skin prick and atopy patch testing has been effective in children with EoE, but additional validation is needed. Empiric elimination of common food allergies (milk, wheat, egg, soy, nuts, and seafood) followed by systematic reintroduction has been an effective diet therapy in both children and adults with EoE. The intent of the elimination diet approach is the identification of a single food trigger or a small number of food triggers. Swallowed, topical

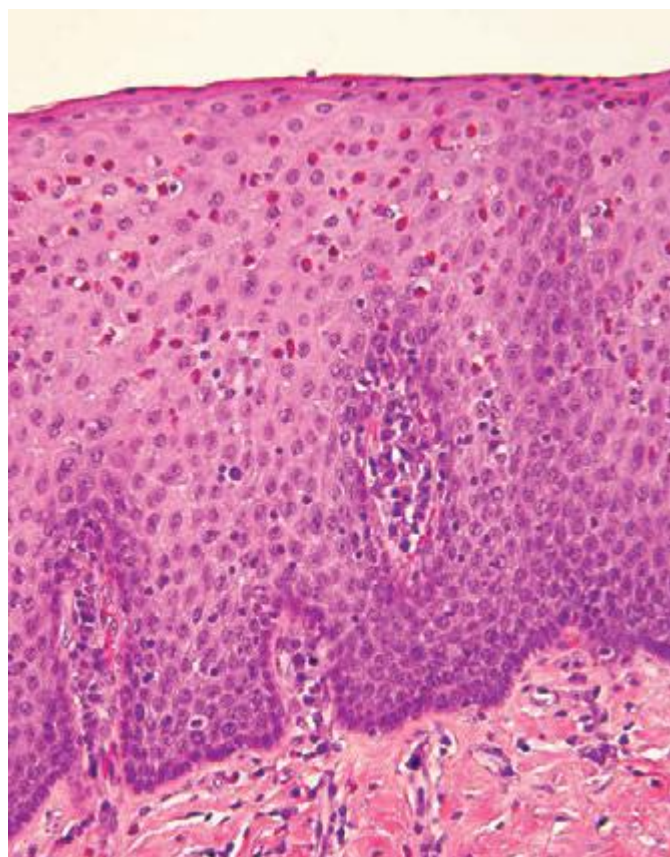


FIGURE 347-12 Histopathology of eosinophilic esophagitis (EoE) showing infiltration of the esophageal squamous epithelium with eosinophils. Additional features of basal cell hyperplasia and lamina propria fibrosis are present. Eosinophilic inflammation can also be seen with gastroesophageal reflux disease.

glucocorticoids (fluticasone propionate or budesonide) are highly effective, but recurrence of disease is common following the cessation of therapy. Systemic glucocorticoids are reserved for severely afflicted patients refractory to less morbid treatments. Esophageal dilation is very effective at relieving dysphagia in patients with fibrostenosis. Dilation should be approached conservatively because of the risk of deep, esophageal mural laceration or perforation in the stiff-walled esophagus that is characteristic of the disease.

INFECTIOUS ESOPHAGITIS

With the increased use of immunosuppression for organ transplantation as well as chronic inflammatory diseases and chemotherapy along with the AIDS epidemic, infections with *Candida* species, herpesvirus, and cytomegalovirus (CMV) have become relatively common. Although rare, infectious esophagitis also occurs among the nonimmunocompromised, with herpes simplex and *Candida albicans* being the most common pathogens. Among AIDS patients, infectious esophagitis becomes more common as the CD4 count declines; cases are rare with a CD4 count >200 and common when <100 . HIV itself may also be associated with a self-limited syndrome of acute esophageal ulceration with oral ulcers and a maculopapular skin rash at the time of seroconversion. Additionally, some patients with advanced disease have deep, persistent esophageal ulcers treated with oral glucocorticoids or thalidomide. However, with the widespread use of protease inhibitors, a reduction in these HIV complications has been noted.

Regardless of the infectious agent, odynophagia is a characteristic symptom of infectious esophagitis; dysphagia, chest pain, and hemorrhage are also common. Odynophagia is uncommon with reflux esophagitis, so its presence should always raise suspicion of an alternative etiology.

Candida is normally found in the throat, but can become pathogenic and produce esophagitis in a compromised host; *C. albicans* is most common. *Candida* esophagitis also occurs with esophageal stasis secondary to esophageal motor disorders and diverticula. Patients complain of odynophagia and dysphagia. If oral thrush is present, empirical therapy is appropriate, but co-infection is common, and persistent symptoms should lead to prompt endoscopy with biopsy, which is the most useful diagnostic evaluation. *Candida* esophagitis has a characteristic appearance of white plaques with friability. Rarely, *Candida* esophagitis is complicated by bleeding, perforation, stricture, or systemic invasion. Oral fluconazole (200–400 mg on the first day, followed by 100–200 mg daily) for 14–21 days is the preferred treatment. Patients refractory to fluconazole may respond to itraconazole, voriconazole, or posaconazole. Alternatively, poorly responsive patients or those who cannot swallow medications can be treated with an intravenous echinocandin (caspofungin 50 mg daily for 7–21 days).

HERPETIC ESOPHAGITIS

Herpes simplex virus type 1 or 2 may cause esophagitis. Vesicles on the nose and lips may coexist and are suggestive of a herpetic etiology. Varicella-zoster virus can also cause esophagitis in children with chickenpox or adults with zoster. The characteristic endoscopic findings are vesicles and small, punched-out ulcerations. Because herpes simplex infections are limited to squamous epithelium, biopsies from the ulcer margins are most likely to reveal the characteristic ground-glass nuclei, eosinophilic Cowdry's type A inclusion bodies, and giant cells. Culture or polymerase chain reaction (PCR) assays are helpful to identify acyclovir-resistant strains. Acyclovir (200 mg orally five times a day for 7–10 days) can be used for immunocompetent hosts, although the disease is typically self-limited after a 1- to 2-week period in such patients. Immunocompromised patients are treated with acyclovir (400 mg orally five times a day for 14–21 days), famciclovir (500 mg orally three times a day), or valacyclovir (1 g orally three times a day). In patients with severe odynophagia, intravenous acyclovir, 5 mg/kg every 8 h for 7–14 days, reduces this morbidity.

CYTOMEGALOVIRUS

CMV esophagitis occurs primarily in immunocompromised patients, particularly organ transplant recipients. CMV is usually activated from a latent stage. Endoscopically, CMV lesions appear as serpiginous ulcers in an otherwise normal mucosa, particularly in the distal esophagus. Biopsies from the ulcer bases have the greatest diagnostic yield for finding the pathognomonic large nuclear or cytoplasmic inclusion bodies. Immunohistology with monoclonal antibodies to CMV and in situ hybridization tests are useful for early diagnosis. Data on therapy for CMV esophagitis are limited. Treatment studies of CMV gastrointestinal disease have demonstrated effectiveness of both ganciclovir (5 mg/kg every 12 h intravenously) and foscarnet (90 mg/kg every 12 h intravenously). Valganciclovir (900 mg two times a day), an oral formulation of ganciclovir, can also be used. Therapy is continued until healing, which may take 3–6 weeks. Maintenance therapy may be needed for patients with relapsing disease.

MECHANICAL TRAUMA AND IATROGENIC INJURY

ESOPHAGEAL PERFORATION

Most cases of esophageal perforation are from instrumentation of the esophagus or trauma. Alternatively, forceful vomiting or retching can lead to spontaneous rupture at the gastroesophageal junction (Boerhaave's syndrome). More rarely, corrosive esophagitis or neoplasms lead to perforation. Instrument perforation from endoscopy or nasogastric tube placement typically occurs in the hypopharynx or at the gastroesophageal junction. Perforation may also occur at the site of a stricture in the setting of endoscopic food disimpaction or esophageal dilation. Esophageal perforation causes pleuritic retrosternal pain that can be associated with pneumomediastinum and subcutaneous emphysema. Mediastinitis is a major complication of esophageal perforation, and prompt recognition is key to optimizing outcome. CT

of the chest is most sensitive in detecting mediastinal air. Esophageal perforation is confirmed by a contrast swallow, usually Gastrografin followed by thin barium. Treatment includes nasogastric suction and parenteral broad-spectrum antibiotics with prompt surgical drainage and repair in noncontained leaks. Conservative therapy with NPO status and antibiotics without surgery may be appropriate in cases of contained perforation that are detected early. Endoscopic clipping or stent placement may be indicated in nonoperated iatrogenic perforations or nonoperable cases such as perforated tumors.

MALLORY-WEISS TEAR

Vomiting, retching, or vigorous coughing can cause a nontransmural tear at the gastroesophageal junction that is a common cause of upper gastrointestinal bleeding. Most patients present with hematemesis. Antecedent vomiting is anticipated but not always evident. Bleeding usually abates spontaneously, but protracted bleeding may respond to local epinephrine or cauterization therapy, endoscopic clipping, or angiographic embolization. Surgery is rarely needed.

RADIATION ESOPHAGITIS

Radiation esophagitis can complicate treatment for thoracic cancers, especially breast and lung, with the risk proportional to radiation dosage. Radiosensitizing drugs such as doxorubicin, bleomycin, cyclophosphamide, and cisplatin also increase the risk. Dysphagia and odynophagia may last weeks to months after therapy. The esophageal mucosa becomes erythematous, edematous, and friable. Submucosal fibrosis and degenerative tissue changes and stricturing may occur years after the radiation exposure. Radiation exposure in excess of 5000 cGy has been associated with increased risk of esophageal stricture. Treatment for acute radiation esophagitis is supportive. Chronic strictures are managed with esophageal dilation.

CORROSIVE ESOPHAGITIS

Caustic esophageal injury from ingestion of alkali or, less commonly, acid can be accidental or from attempted suicide. Absence of oral injury does not exclude possible esophageal involvement. Thus, early endoscopic evaluation is recommended to assess and grade the injury to the esophageal mucosa. Severe corrosive injury may lead to esophageal perforation, bleeding, stricture, and death. Glucocorticoids have not been shown to improve the clinical outcome of acute corrosive esophagitis and are not recommended. Healing of more severe grades of caustic injury is commonly associated with severe stricture formation and often requires repeated dilatation.

PILL ESOPHAGITIS

Pill-induced esophagitis occurs when a swallowed pill fails to traverse the entire esophagus and lodges within the lumen. Generally, this is attributed to poor "pill taking habits": inadequate liquid with the pill or lying down immediately after taking a pill. The most common location for the pill to lodge is in the mid-esophagus near the crossing of the aorta or carina. Extrinsic compression from these structures halts the movement of the pill or capsule. Since initially reported in 1970, more than 1000 cases of pill esophagitis have been reported, suggesting that this is not an unusual occurrence. A wide variety of medications are implicated with the most common being doxycycline, tetracycline, quinidine, phenytoin, potassium chloride, ferrous sulfate, nonsteroidal anti-inflammatory drugs (NSAIDs), and bisphosphonates. However, virtually any pill can result in pill esophagitis if taken carelessly.

Typical symptoms of pill esophagitis are the sudden onset of chest pain and odynophagia. Characteristically, the pain will develop over a period of hours or will awaken the individual from sleep. A classic history in the setting of ingestion of recognized pill offenders obviates the need for diagnostic testing in most patients. When endoscopy is performed, localized ulceration or inflammation is evident. Histologically, acute inflammation is typical. Chest CT imaging will sometimes reveal esophageal thickening consistent with transmural inflammation. Although the condition usually resolves within days to weeks, symptoms may persist for months and stricture can develop in severe cases. No specific therapy is known to hasten the healing process, but antisecretory medications are frequently prescribed to

remove concomitant reflux as an aggravating factor. When healing results in stricture formation, dilation is indicated.

FOREIGN BODIES AND FOOD IMPACTION

Food or foreign bodies may lodge in the esophagus causing complete obstruction, which in turn can cause an inability to handle secretions (foaming at the mouth) and severe chest pain. Food impaction may occur due to stricture, carcinoma, Schatzki ring, eosinophilic esophagitis, or simply inattentive eating. If it does not spontaneously resolve, impacted food can be dislodged endoscopically. Use of meat tenderizer enzymes to facilitate passage of a meat bolus is discouraged because of potential esophageal injury. Glucagon (1 mg IV) is sometimes tried before endoscopic dislodgement. After emergent treatment, patients should be evaluated for potential causes of the impaction with treatment rendered as indicated.

ESOPHAGEAL MANIFESTATIONS OF SYSTEMIC DISEASE

SCLERODERMA AND COLLAGEN VASCULAR DISEASES

Scleroderma esophagus (hypotensive LES and absent esophageal peristalsis) was initially described as a manifestation of scleroderma or other collagen vascular diseases and thought to be specific for these disorders. However, this nomenclature subsequently proved unfortunate and has been discarded because an estimated half of qualifying patients do not have an identifiable systemic disease, and reflux disease is often the only identifiable association. When scleroderma esophagus occurs as a manifestation of a collagen vascular disease, the histopathologic findings are of infiltration and destruction of the esophageal muscularis propria with collagen deposition and fibrosis. The pathogenesis of absent peristalsis and LES hypotension in the absence of a collagen vascular disease is unknown. Regardless of the underlying cause, the manometric abnormalities predispose patients to severe GERD due to inadequate LES barrier function combined with poor esophageal clearance of refluxed acid. Dysphagia may also be manifest but is generally mild and alleviated by eating in an upright position and using liquids to facilitate solid emptying.

DERMATOLOGIC DISEASES

A host of dermatologic disorders (pemphigus vulgaris, bullous pemphigoid, cicatricial pemphigoid, Behçet's syndrome, and epidermolysis bullosa) can affect the oropharynx and esophagus, particularly the proximal esophagus with blisters, bullae, webs, and strictures. Glucocorticoid treatment is usually effective. Erosive lichen planus, Stevens-Johnson syndrome, and graft-versus-host disease can also involve the esophagus. Esophageal dilatation may be necessary to treat strictures.

348 Peptic Ulcer Disease and Related Disorders

John Del Valle

PEPTIC ULCER DISEASE

Burning epigastric pain exacerbated by fasting and improved with meals is a symptom complex associated with peptic ulcer disease (PUD). An *ulcer* is defined as disruption of the mucosal integrity of the stomach and/or duodenum leading to a local defect or excavation due to active inflammation. Ulcers occur within the stomach and/or duodenum and are often chronic in nature. Acid peptic disorders are very common in the United States, with 4 million individuals (new cases and recurrences) affected per year. Lifetime prevalence of PUD in the United States is ~12% in men and 10% in women. PUD

significantly affects quality of life by impairing overall patient well-being and contributing substantially to work absenteeism. Moreover, an estimated 15,000 deaths per year occur as a consequence of complicated PUD. The financial impact of these common disorders has been substantial, with an estimated burden on direct and indirect health care costs of ~\$6 billion per year in the United States, with \$3 billion spent on hospitalizations, \$2 billion on physician office visits, and \$1 billion in decreased productivity and days lost from work.

GASTRIC PHYSIOLOGY

Despite the constant attack on the gastroduodenal mucosa by a host of noxious agents (acid, pepsin, bile acids, pancreatic enzymes, drugs, and bacteria), integrity is maintained by an intricate system that provides mucosal defense and repair.

Gastric Anatomy The gastric epithelial lining consists of rugae that contain microscopic gastric pits, each branching into four or five gastric glands made up of highly specialized epithelial cells. The makeup of gastric glands varies with their anatomic location. Glands within the gastric cardia comprise <5% of the gastric gland area and contain mucous and endocrine cells. The 75% of gastric glands are found within the oxyntic mucosa and contain mucous neck, parietal, chief, endocrine, enterochromaffin, and enterochromaffin-like (ECL) cells (Fig. 348-1). Pyloric glands contain mucous and endocrine cells (including gastrin cells) and are found in the antrum.

The parietal cell, also known as the oxyntic cell, is usually found in the neck, or isthmus, or in the oxyntic gland. The resting, or unstimulated, parietal cell has prominent cytoplasmic tubulovesicles and intracellular canaliculi containing short microvilli along its apical surface (Fig. 348-2). H^+,K^+ -adenosine triphosphatase (ATPase) is expressed in the tubulovesicle membrane; upon cell stimulation, this membrane, along with apical membranes, transforms into a dense network of apical intracellular canaliculi containing long microvilli. Acid secretion, a

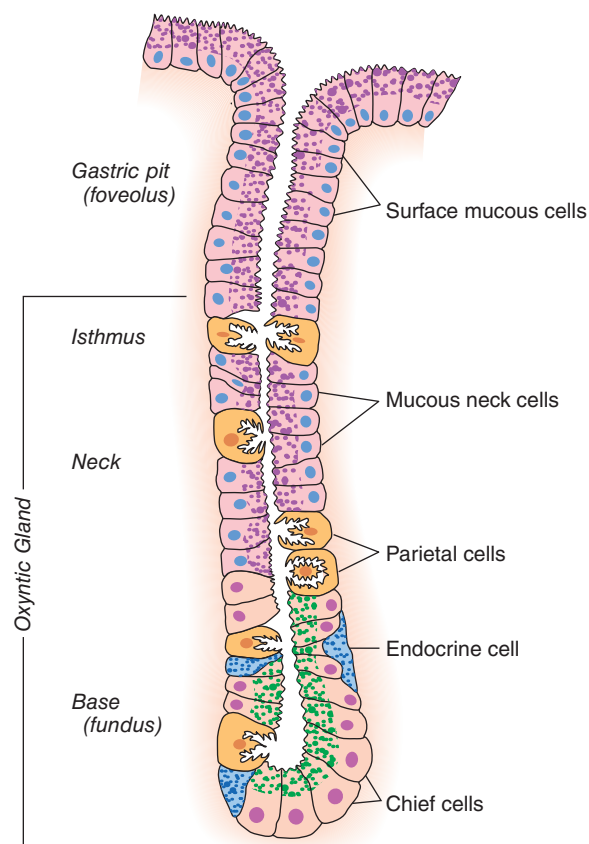


FIGURE 348-1 Diagrammatic representation of the oxyntic gastric gland. (Adapted from S Ito, RJ Winchester. *J Cell Biol* 16:541, 1963. doi:10.1083/jcb.16.3.541. © 1963 Ito and Winchester.)

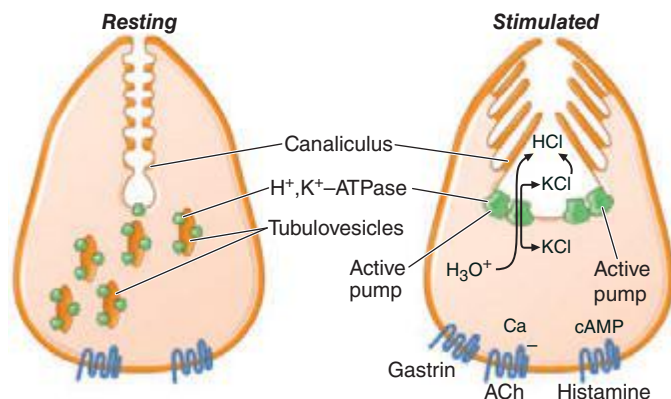


FIGURE 348-2 Gastric parietal cell undergoing transformation after secretagogue-mediated stimulation. cAMP, cyclic adenosine monophosphate. (Adapted from SJ Hersey, G Sachs: *Physiol Rev* 75:155, 1995.)

process requiring high energy, occurs at the apical canalicular surface. Numerous mitochondria (30–40% of total cell volume) generate the energy required for secretion.

Gastroduodenal Mucosal Defense The gastric epithelium is under constant assault by a series of endogenous noxious factors, including hydrochloric acid (HCl), pepsinogen/pepsin, and bile salts. In addition, a steady flow of exogenous substances such as medications, alcohol, and bacteria encounter the gastric mucosa. A highly intricate biologic system is in place to provide defense from mucosal injury and to repair any injury that may occur.

The mucosal defense system can be envisioned as a three-level barrier, composed of preepithelial, epithelial, and subepithelial elements (Fig. 348-3). The first line of defense is a mucus-bicarbonate-phospholipid layer, which serves as a physicochemical barrier to multiple molecules, including hydrogen ions. Mucus is secreted in a regulated fashion by gastroduodenal surface epithelial cells. It consists primarily of water (95%) and a mixture of phospholipids and glycoproteins (mucin). The mucous gel functions as a nonstirred water layer impeding diffusion of ions and molecules such as pepsin. Bicarbonate, secreted in a regulated manner by surface epithelial cells of the gastroduodenal mucosa into the mucous gel, forms a pH gradient ranging from 1 to 2 at the gastric luminal surface and reaching 6 to 7 along the epithelial cell surface.

Surface epithelial cells provide the next line of defense through several factors, including mucus production, epithelial cell ionic transporters that maintain intracellular pH and bicarbonate production, and intracellular tight junctions. Surface epithelial cells generate heat shock proteins that prevent protein denaturation and protect cells from certain factors such as increased temperature, cytotoxic agents, or oxidative stress. Epithelial cells also generate trefoil factor family peptides and cathelicidins, which also play a role in surface cell protection and regeneration. If the preepithelial barrier were breached, gastric epithelial cells bordering a site of injury can migrate to restore a damaged region (*restitution*). This process occurs independent of cell division and requires uninterrupted blood flow and an alkaline pH in the surrounding environment. Several growth factors, including epidermal growth factor (EGF), transforming growth factor (TGF) α , and basic fibroblast growth factor (FGF), modulate the process of restitution. Larger defects that are not effectively repaired by restitution require cell proliferation. Epithelial cell regeneration is regulated by prostaglandins and growth factors such as EGF and TGF- α . In tandem with epithelial cell renewal, formation of new vessels (*angiogenesis*) within the injured microvascular bed occurs. Both FGF and vascular endothelial growth factor (VEGF) are important in regulating angiogenesis in the gastric mucosa.

An elaborate microvascular system within the gastric submucosal layer is the key component of the subepithelial defense/repair system, providing HCO_3^- , which neutralizes the acid generated by the parietal cell. Moreover, this microcirculatory bed provides an adequate

supply of micronutrients and oxygen while removing toxic metabolic by-products.

Prostaglandins play a central role in gastric epithelial defense/repair (Fig. 348-4). The gastric mucosa contains abundant levels of prostaglandins that regulate the release of mucosal bicarbonate and mucus, inhibit parietal cell secretion, and are important in maintaining mucosal blood flow and epithelial cell restitution. Prostaglandins are derived from esterified arachidonic acid, which is formed from phospholipids (cell membrane) by the action of phospholipase A_2 . A key enzyme that controls the rate-limiting step in prostaglandin synthesis is cyclooxygenase (COX), which is present in two isoforms (COX-1, COX-2), each having distinct characteristics regarding structure, tissue distribution, and expression. COX-1 is expressed in a host of tissues, including the stomach, platelets, kidneys, and endothelial cells. This isoform is expressed in a constitutive manner and plays an important role in maintaining the integrity of renal function, platelet aggregation, and gastrointestinal (GI) mucosal integrity. In contrast, the expression of COX-2 is inducible by inflammatory stimuli, and it is expressed in macrophages, leukocytes, fibroblasts, and synovial cells. The beneficial effects of nonsteroidal anti-inflammatory drugs (NSAIDs) on tissue inflammation are due to inhibition of COX-2; the toxicity of these drugs (e.g., GI mucosal ulceration and renal dysfunction) is related to inhibition of the COX-1 isoform. The highly COX-2-selective NSAIDs have the potential to provide the beneficial effect of decreasing tissue inflammation while minimizing toxicity in the GI tract. Selective COX-2 inhibitors have had adverse effects on the cardiovascular system, leading to increased risk of myocardial infarction. Therefore, the U.S. Food and Drug Administration (FDA) has removed two of these agents (valdecoxib and rofecoxib) from the market (see below).

Nitric oxide (NO) is important in the maintenance of gastric mucosal integrity. The key enzyme NO synthase is constitutively expressed in the mucosa and contributes to cytoprotection by stimulating gastric mucus, increasing mucosal blood flow, and maintaining epithelial cell barrier function. The central nervous system (CNS) and hormonal factors also play a role in regulating mucosal defense through multiple pathways (Fig. 348-3).

Physiology of Gastric Secretion Hydrochloric acid and pepsinogen are the two principal gastric secretory products capable of inducing mucosal injury. Gastric acid and pepsinogen play a physiologic role in protein digestion; absorption of iron, calcium, magnesium, and vitamin B_{12} ; and killing ingested bacteria. Acid secretion should be viewed as occurring under basal and stimulated conditions. Basal acid production occurs in a circadian pattern, with highest levels occurring during the night and lowest levels during the morning hours. Cholinergic input via the vagus nerve and histaminergic input from local gastric sources are the principal contributors to basal acid secretion. Stimulated gastric acid secretion occurs primarily in three phases based on the site where the signal originates (cephalic, gastric, and intestinal). Sight, smell, and taste of food are the components of the cephalic phase, which stimulates gastric secretion via the vagus nerve. The gastric phase is activated once food enters the stomach. This component of secretion is driven by nutrients (amino acids and amines) that directly stimulate the G cell to release gastrin, which in turn activates the parietal cell via direct and indirect mechanisms. Distention of the stomach wall also leads to gastrin release and acid production. The last phase of gastric acid secretion is initiated as food enters the intestine and is mediated by luminal distention and nutrient assimilation. A series of pathways that inhibit gastric acid production are also set into motion during these phases. The GI hormone somatostatin is released from endocrine cells found in the gastric mucosa (D cells) in response to HCl. Somatostatin can inhibit acid production by both direct (parietal cell) and indirect mechanisms (decreased histamine release from ECL cells and gastrin release from G cells). Additional neural (central and peripheral) and humoral (amylin, atrial natriuretic peptide [ANP], cholecystokinin, ghrelin, interleukin 11 [IL-11], obestatin, secretin, and serotonin) factors play a role in counterbalancing acid secretion. Under physiologic circumstances, these phases occur simultaneously. Ghrelin, the appetite-regulating hormone expressed in Gr cells in the

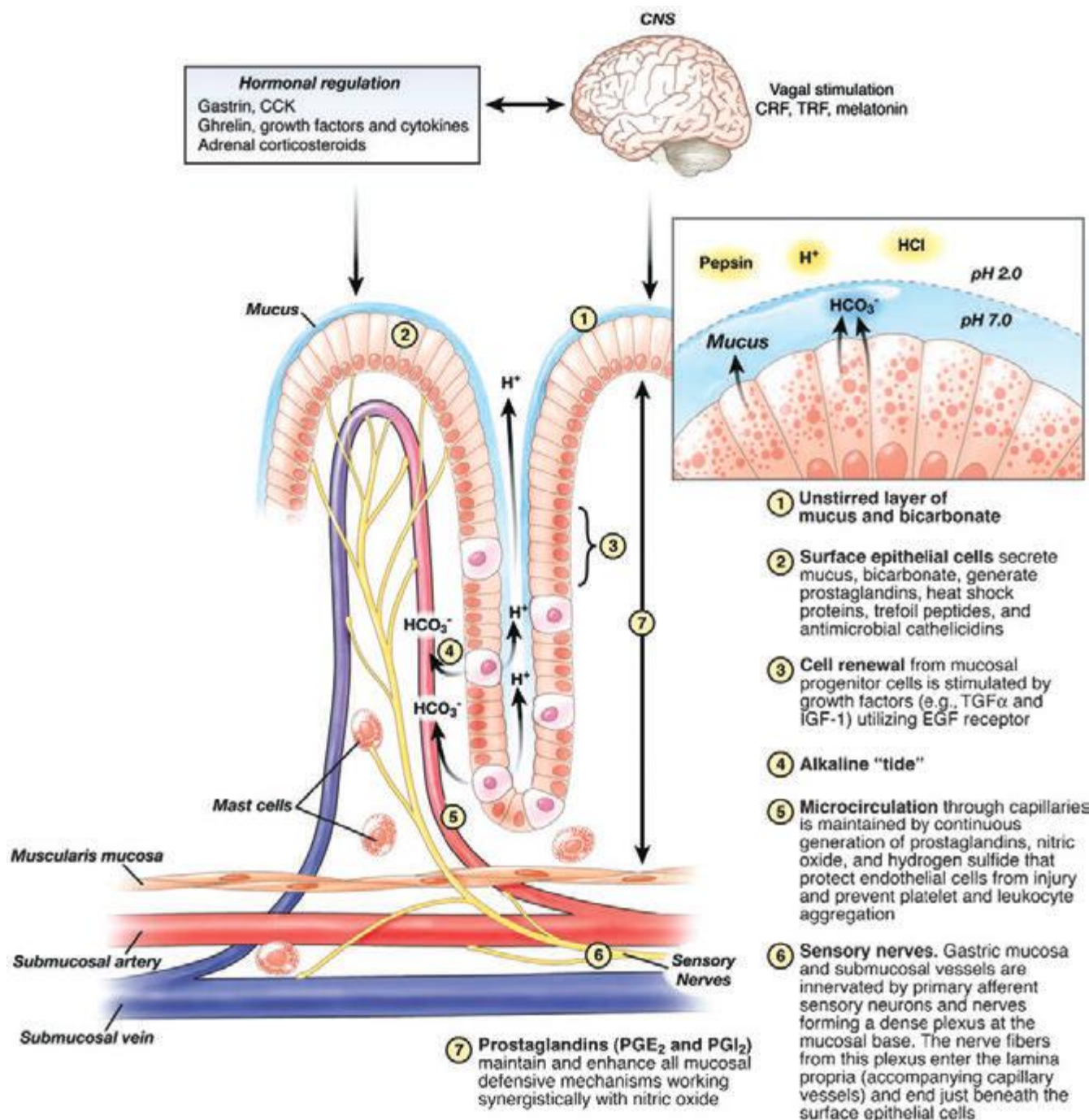


FIGURE 348-3 Components involved in providing gastroduodenal mucosal defense and repair. CCK, cholecystokinin; CRF, corticotropin-releasing factor; EGF, epidermal growth factor; HCl, hydrochloride; IGF, insulin-like growth factor; TGF α , transforming growth factor α ; TRF, thyrotropin releasing factor. (Modified and updated from Tarnawski A. Cellular and molecular mechanisms of mucosal defense and repair. In: Yoshikawa T, Arakawa T. Bioregulation and Its Disorders in the Gastrointestinal Tract. Tokyo, Japan: Blackwell Science, 1998:3–17.)

stomach, may increase gastric acid secretion through stimulation of histamine release from ECL cells, but this remains to be confirmed.

The acid-secreting parietal cell is located in the oxyntic gland, adjacent to other cellular elements (ECL cell, D cell) important in the gastric secretory process (Fig. 348-5). This unique cell also secretes intrinsic factor (IF) and IL-11. The parietal cell expresses receptors for several stimulants of acid secretion, including histamine (H $_2$), gastrin (cholecystokinin B/gastrin receptor), and acetylcholine (muscarinic, M $_3$). Binding of histamine to the H $_2$ receptor leads to activation of adenylate cyclase and an increase in cyclic adenosine monophosphate (AMP). Activation of the gastrin and muscarinic receptors results in activation of the protein kinase C/phosphoinositide signaling pathway. Each of these signaling pathways in turn regulates a series of downstream kinase cascades that control the acid-secreting pump, H $^+$,K $^+$ -

ATPase. The discovery that different ligands and their corresponding receptors lead to activation of different signaling pathways explains the potentiation of acid secretion that occurs when histamine and gastrin or acetylcholine are combined. More importantly, this observation explains why blocking one receptor type (H $_2$) decreases acid secretion stimulated by agents that activate a different pathway (gastrin, acetylcholine). Parietal cells also express receptors for ligands that inhibit acid production (prostaglandins, somatostatin, and EGF). Histamine also stimulates gastric acid secretion indirectly by activating the histamine H $_3$ receptor on D-cells, which inhibits somatostatin release.

The enzyme H $^+$,K $^+$ -ATPase is responsible for generating the large concentration of H $^+$. It is a membrane-bound protein that consists of two subunits, α and β . The active catalytic site is found within the α subunit; the function of the β subunit is unclear. This enzyme uses the

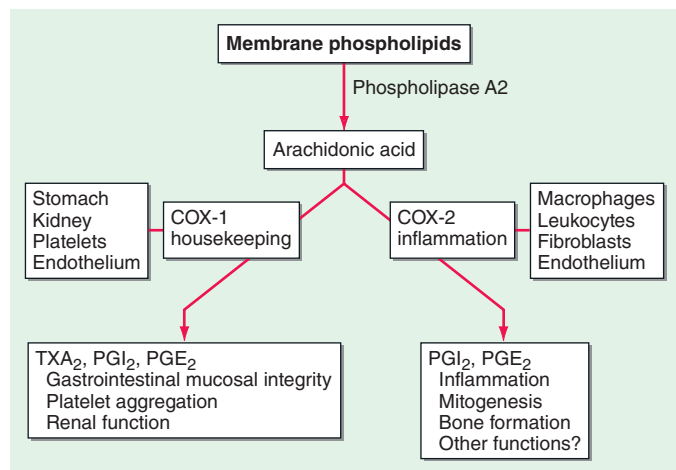


FIGURE 348-4 Schematic representation of the steps involved in synthesis of prostaglandin E₂ (PGE₂) and prostacyclin (PGI₂). Characteristics and distribution of the cyclooxygenase (COX) enzymes 1 and 2 are also shown. TXA₂, thromboxane A₂.

chemical energy of adenosine triphosphate (ATP) to transfer H⁺ ions from parietal cell cytoplasm to the secretory canaliculi in exchange for K⁺. The H⁺,K⁺-ATPase is located within the secretory canaliculus and in nonsecretory cytoplasmic tubulovesicles. The tubulovesicles are impermeable to K⁺, which leads to an inactive pump in this location. The distribution of pumps between the nonsecretory vesicles and the secretory canaliculus varies according to parietal cell activity (Fig. 348-2). Proton pumps are recycled back to the inactive state in cytoplasmic vesicles once parietal cell activation ceases. Small G proteins of the Rab family and secretory carrier membrane proteins (SCAMPS) are postulated to participate in parietal cell membrane translocation. In addition, acid secretion requires a number of apical and basolateral parietal cell membrane chloride and potassium channels.

The chief cell, found primarily in the gastric fundus, synthesizes and secretes pepsinogen, the inactive precursor of the proteolytic enzyme pepsin. The acid environment within the stomach leads to cleavage of the inactive precursor to pepsin and provides the low pH (<2) required for pepsin activity. Pepsin activity is significantly diminished at a pH of 4 and irreversibly inactivated and denatured at a pH of ≥7. Many of the

secretagogues that stimulate acid secretion also stimulate pepsinogen release. The precise role of pepsin in the pathogenesis of PUD remains to be established.

PATHOPHYSIOLOGIC BASIS OF PEPTIC ULCER DISEASE

PUD encompasses both gastric and duodenal ulcers. *Ulcers* are defined as breaks in the mucosal surface >5 mm in size, with depth to the submucosa. Duodenal ulcers (DUs) and gastric ulcers (GUs) share many common features in terms of pathogenesis, diagnosis, and treatment, but several factors distinguish them from one another. *Helicobacter pylori* and NSAIDs are the most common risk factors for PUD, with estimated odds ratios in the United States of 3.7 and 3.3, respectively. Additional risk factors (odds ratio) include chronic obstructive lung disease (2.34), chronic renal insufficiency (2.29), current tobacco use (1.99), former tobacco use (1.55), older age (1.67), three or more doctor visits in a year (1.49), coronary heart disease (1.46), former alcohol use (1.29), African-American race (1.20), obesity (1.18), and diabetes (1.13). The mechanisms by which some of these risk factors lead to ulcer disease are highlighted below.

Epidemiology • DUODENAL ULCERS DUs are estimated to occur in 6–15% of the Western population. The incidence of DUs declined steadily from 1960 to 1980 and has remained stable since then. The death rates, need for surgery, and physician visits have decreased by >50% over the past 30 years. The reason for the reduction in the frequency of DUs is likely related to the decreasing frequency of *H. pylori*. Before the discovery of *H. pylori*, the natural history of DUs was typified by frequent recurrences after initial therapy. Eradication of *H. pylori* has greatly reduced these recurrence rates.

GASTRIC ULCERS GUs tend to occur later in life than duodenal lesions, with a peak incidence reported in the sixth decade. More than one-half of GUs occur in males and are less common than DUs, perhaps due to the higher likelihood of GUs being silent and presenting only after a complication develops. Autopsy studies suggest a similar incidence of DUs and GUs.

Pathology • DUODENAL ULCERS DUs occur most often in the first portion of the duodenum (>95%), with ~90% located within 3 cm of the pylorus. They are usually ≤1 cm in diameter but can occasionally reach 3–6 cm (giant ulcer). Ulcers are sharply demarcated, with depth at times reaching the muscularis propria. The base of the ulcer often consists of a zone of eosinophilic necrosis with surrounding fibrosis. Malignant DUs are extremely rare.

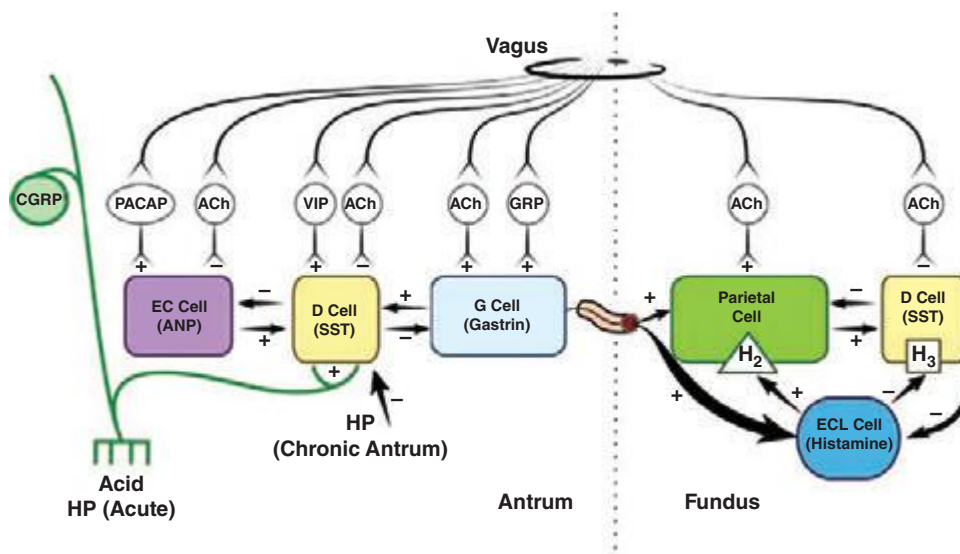


FIGURE 348-5 Regulation of gastric acid secretion at the cellular level. ACh, acetylcholine; ANP, atrial natriuretic peptide; CGRP, calcitonin gene-related peptide; EC, enterochromaffin; ECL, enterochromaffin-like; GRP, gastrin-releasing peptide; PACAP, pituitary adenylate-cyclase activating peptide; SST, somatostatin; VIP, vasoactive intestinal peptide.

GASTRIC ULCERS In contrast to DUs, GUs can represent a malignancy and should be biopsied upon discovery. Benign GUs are most often found distal to the junction between the antrum and the acid secretory mucosa. Benign GUs are quite rare in the gastric fundus and are histologically similar to DUs. Benign GUs associated with *H. pylori* are also associated with antral gastritis. In contrast, NSAID-related GUs are not accompanied by chronic active gastritis but may instead have evidence of a chemical gastropathy, typified by foveolar hyperplasia, edema of the lamina propria, and epithelial regeneration in the absence of *H. pylori*. Extension of smooth-muscle fibers into the upper portions of the mucosa, where they are not typically found, may also occur.

Pathophysiology • DUODENAL ULCERS *H. pylori* and NSAID-induced injury account for the majority of DUs. Many acid secretory abnormalities have been described in DU patients. Of these, average basal and nocturnal gastric acid secretion appears to be increased in DU patients as compared to controls; however, the level of overlap between DU patients and control subjects is substantial. The reason for this altered secretory process is unclear, but *H. pylori* infection may contribute. Bicarbonate secretion is significantly decreased in the duodenal bulb of patients with an active DU as compared to control subjects. *H. pylori* infection may also play a role in this process (see below).

GASTRIC ULCERS As in DUs, the majority of GUs can be attributed to either *H. pylori* or NSAID-induced mucosal damage. GUs that occur in the prepyloric area or those in the body associated with a DU or a duodenal scar are similar in pathogenesis to DUs. Gastric acid output (basal and stimulated) tends to be normal or decreased in GU patients. When GUs develop in the presence of minimal acid levels, impairment of mucosal defense factors may be present. GUs have been classified based on their location: Type I occur in the gastric body and tend to be associated with low gastric acid production; type II occur in the antrum and gastric acid can vary from low to normal; type III occur within 3 cm of the pylorus and are commonly accompanied by DUs and normal or high gastric acid production; and type IV are found in the cardia and are associated with low gastric acid production.

H. PYLORI AND ACID PEPTIC DISORDERS Gastric infection with the bacterium *H. pylori* accounts for the majority of PUD (Chap. 188). This organism also plays a role in the development of gastric mucosa-associated lymphoid tissue (MALT) lymphoma and gastric adenocarcinoma. Although the entire genome of *H. pylori* has been sequenced, it is still not clear how this organism, which resides in the stomach, causes ulceration in the duodenum, or whether its eradication will lead to a decrease in gastric cancer.

The bacterium The bacterium, initially named *Campylobacter pyloridis*, is a gram-negative microaerophilic rod found most commonly in the deeper portions of the mucous gel coating the gastric mucosa or between the mucous layer and the gastric epithelium. It may attach to gastric epithelium but under normal circumstances does not appear to invade cells. It is strategically designed to live within the aggressive environment of the stomach. It is S-shaped (~0.5–3 μm in size) and contains multiple sheathed flagella. Initially, *H. pylori* resides in the antrum but, over time, migrates toward the more proximal segments of the stomach. The organism is capable of transforming into a coccoid form, which represents a dormant state that may facilitate survival in adverse conditions. The genome of *H. pylori* (1.65 million base pairs) encodes ~1500 proteins. Among this multitude of proteins there are factors that are essential determinants of *H. pylori*-mediated pathogenesis and colonization such as the outer membrane protein (Hop proteins), urease, and the vacuolating cytotoxin (Vac A). Moreover, the majority of *H. pylori* strains contain a genomic fragment that encodes the cag pathogenicity island (cag-PAI). Several of the genes that make up cag-PAI encode components of a type IV secretion island that translocates Cag A into host cells. Once in the cell, Cag A activates a series of cellular events important in cell growth and cytokine production. *H. pylori* also has extensive genetic diversity that in turn enhances its ability to promote disease. The first step in infection by *H. pylori* is dependent on the bacteria's motility and its ability to

produce urease. Urease produces ammonia from urea, an essential step in alkalizing the surrounding pH. Additional bacterial factors include catalase, lipase, adhesins, platelet-activating factor, and pic B (induces cytokines). Multiple strains of *H. pylori* exist and are characterized by their ability to express several of these factors (Cag A, Vac A, etc.). It is possible that the different diseases related to *H. pylori* infection can be attributed to different strains of the organism with distinct pathogenic features.

Epidemiology The prevalence of *H. pylori* varies throughout the world and depends largely on the overall standard of living in the region. In developing parts of the world, 80% of the population may be infected by the age of 20, whereas the prevalence is 20–50% in industrialized countries. In contrast, in the United States this organism is rare in childhood. The overall prevalence of *H. pylori* in the United States is ~30%, with individuals born before 1950 having a higher rate of infection than those born later. About 10% of Americans <30 years of age are colonized with the bacteria. The rate of infection with *H. pylori* in industrialized countries has decreased substantially in recent decades. The steady increase in the prevalence of *H. pylori* noted with increasing age is due primarily to a cohort effect, reflecting higher transmission during a period in which the earlier cohorts were children. It has been calculated through mathematical models that improved sanitation during the latter half of the nineteenth century dramatically decreased transmission of *H. pylori*. Moreover, with the present rate of intervention, the organism will be ultimately eliminated from the United States. Two factors that predispose to higher colonization rates include poor socioeconomic status and less education. These factors, not race, are responsible for the rate of *H. pylori* infection in blacks and Hispanic Americans being double the rate seen in whites of comparable age. Other risk factors for *H. pylori* infection are (1) birth or residence in a developing country, (2) domestic crowding, (3) unsanitary living conditions, (4) unclean food or water, and (5) exposure to gastric contents of an infected individual.

Transmission of *H. pylori* occurs from person to person, following an oral-oral or fecal-oral route. The risk of *H. pylori* infection is declining in developing countries. The rate of infection in the United States has fallen by >50% when compared to 30 years ago.

Pathophysiology *H. pylori* infection is virtually always associated with a chronic active gastritis, but only 10–15% of infected individuals develop frank peptic ulceration. The basis for this difference is unknown, but is likely due to a combination of host and bacterial factors some of which are outlined below. Initial studies suggested that >90% of all DUs were associated with *H. pylori*, but *H. pylori* is present in only 30–60% of individuals with GUs and 50–70% of patients with DUs. The pathophysiology of ulcers not associated with *H. pylori* or NSAID ingestion (or the rare Zollinger-Ellison syndrome [ZES]) is becoming more relevant as the incidence of *H. pylori* is dropping, particularly in the Western world (see below).

The particular end result of *H. pylori* infection (gastritis, PUD, gastric MALT lymphoma, gastric cancer) is determined by a complex interplay between bacterial and host factors (Fig. 348-6).

1. **Bacterial factors:** *H. pylori* is able to facilitate gastric residence, induce mucosal injury, and avoid host defense. Different strains of *H. pylori* produce different virulence factors. A specific region of the bacterial genome, the pathogenicity island (cag-PAI), encodes the virulence factors Cag A and pic B. Vac A also contributes to pathogenicity, although it is not encoded within the pathogenicity island. These virulence factors, in conjunction with additional bacterial constituents, can cause mucosal damage, in part through their ability to target the host immune cells. For example, Vac A targets human CD4 T cells, inhibiting their proliferation and in addition can disrupt normal function of B cells, CD8 T cells, macrophages, and mast cells. Multiple studies have demonstrated that *H. pylori* strains that are cag-PAI positive are associated with a higher risk of PUD, premalignant gastric lesions, and gastric cancer than are strains that lack the cag-PAI. In addition, *H. pylori* may directly inhibit parietal cell H⁺K⁺-ATPase activity through a

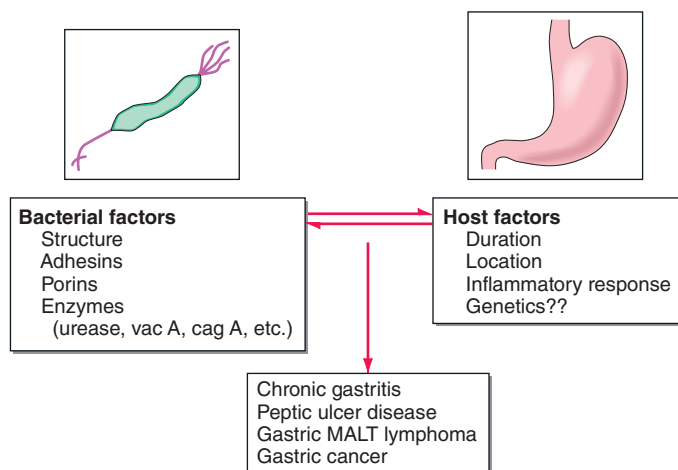


FIGURE 348-6 Outline of the bacterial and host factors important in determining *H. pylori*-induced gastrointestinal disease. MALT, mucosal-associated lymphoid tissue.

Cag A-dependent mechanism, leading in part to the low acid production observed after acute infection with the organism. Urease, which allows the bacteria to reside in the acidic stomach, generates NH_3 , which can damage epithelial cells. The bacteria produce surface factors that are chemotactic for neutrophils and monocytes, which in turn contribute to epithelial cell injury (see below). *H. pylori* makes proteases and phospholipases that break down the glycoprotein lipid complex of the mucous gel, thus reducing the efficacy of this first line of mucosal defense. *H. pylori* expresses adhesins (OMPs like BabA), which facilitate attachment of the bacteria to gastric epithelial cells. Although lipopolysaccharide (LPS) of gram-negative bacteria often plays an important role in the infection, *H. pylori* LPS has low immunologic activity compared to that of other organisms. It may promote a smoldering chronic inflammation.

2. **Host factors:** Studies in twins suggest that there may be genetic predisposition to acquire *H. pylori*. The inflammatory response to *H. pylori* includes recruitment of neutrophils, lymphocytes (T and B), macrophages, and plasma cells. The pathogen leads to local injury by binding to class II major histocompatibility complex (MHC) molecules expressed on gastric epithelial cells, leading to cell death (*apoptosis*). Moreover, bacterial strains that encode cag-PAI can introduce Cag A into the host cells, leading to further cell injury and activation of cellular pathways involved in cytokine production and repression of tumor-suppressor genes. Elevated concentrations of multiple cytokines are found in the gastric epithelium of *H. pylori*-infected individuals, including interleukin (IL) 1 α/β , IL-2, IL-6, IL-8, tumor necrosis factor (TNF) α , and interferon (IFN) γ . *H. pylori* infection also leads to both a mucosal and a systemic humoral response, which does not lead to eradication of the bacteria but further compounds epithelial cell injury. Additional mechanisms by which *H. pylori* may cause epithelial cell injury include (1) activated neutrophil-mediated production of reactive oxygen or nitrogen species and enhanced epithelial cell turnover and (2) apoptosis related to interaction with T cells (T helper 1, or T_H1 , cells) and IFN- γ . Finally, the human stomach can be colonized by a host of commensal organisms that may affect the likelihood of *H. pylori*-mediated mucosal injury.

The reason for *H. pylori*-mediated duodenal ulceration remains unclear. Studies suggest that *H. pylori* associated with duodenal ulceration may be more virulent. In addition, certain specific bacterial factors such as the DU-promoting gene A (*dupA*), may be associated with the development of DUs. Another potential contributing factor is that gastric metaplasia in the duodenum of DU patients, which may be due to high acid exposure (see below), permits *H. pylori* to bind to it and produce local injury secondary to the host response. Another hypothesis is that *H. pylori* antral infection could lead to increased

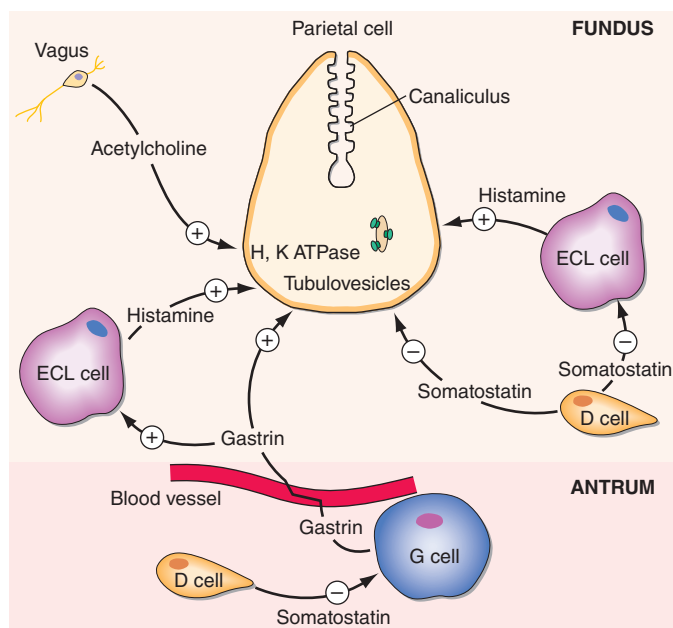


FIGURE 348-7 Summary of potential mechanisms by which *H. pylori* may lead to gastric secretory abnormalities. D, somatostatin cell; ECL, enterochromaffin-like cell; G, G cell. (Adapted from J Calam et al: *Gastroenterology* 113:543, 1997.)

acid production, increased duodenal acid, and mucosal injury. Basal and stimulated (meal, gastrin-releasing peptide [GRP]) gastrin release are increased in *H. pylori*-infected individuals, and somatostatin-secreting D cells may be decreased. *H. pylori* infection might induce increased acid secretion through both direct and indirect actions of *H. pylori* and proinflammatory cytokines (IL-8, TNF, and IL-1) on G, D, and parietal cells (Fig. 348-7). GUs, in contrast, are associated with *H. pylori*-induced pangastritis and normal or low gastric acid secretion. *H. pylori* infection has also been associated with decreased duodenal mucosal bicarbonate production. Data supporting and contradicting each of these interesting theories have been demonstrated. Thus, the mechanism by which *H. pylori* infection of the stomach leads to duodenal ulceration remains to be established.

In summary, the final effect of *H. pylori* on the GI tract is variable and determined by microbial and host factors. The type and distribution of gastritis correlate with the ultimate gastric and duodenal pathology observed. Specifically, the presence of antral-predominant gastritis is associated with DU formation; gastritis involving primarily the corpus predisposes to the development of GUs, gastric atrophy, and ultimately gastric carcinoma (Fig. 348-8).

NSAID-INDUCED DISEASE

Epidemiology NSAIDs represent a group of the most commonly used medications in the United States. More than 30 billion over-the-counter tablets and over 100 million prescriptions are sold yearly in the United States alone. In fact, after the introduction of COX-2 inhibitors in the year 2000, the number of prescriptions written for NSAIDs was >111 million at a cost of \$4.8 billion. Side effects and complications due to NSAIDs are considered the most common drug-related toxicities in the United States. The spectrum of NSAID-induced morbidity ranges from nausea and dyspepsia (prevalence reported as high as 50–60%) to a serious GI complication such as endoscopy-documented peptic ulceration (15–30% of individuals taking NSAIDs regularly) complicated by bleeding or perforation in as many as 1.5% of users per year. It is estimated that NSAID-induced GI bleeding accounts for 60,000–120,000 hospital admissions per year, and deaths related to NSAID-induced toxicity may be as high as 16,000 per year in the United States. Approximately 4–5% of patients develop symptomatic ulcers within 1 year. Unfortunately, dyspeptic symptoms do not correlate with NSAID-induced pathology. Over 80% of patients with serious NSAID-related complications did not have preceding dyspepsia.

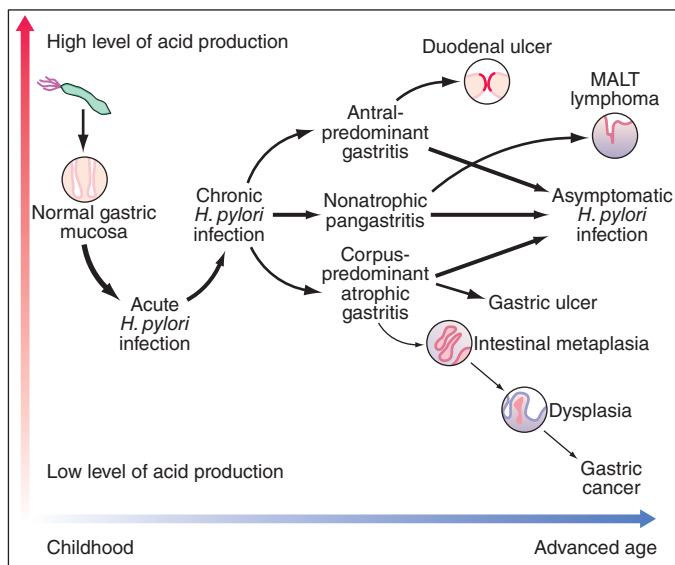


FIGURE 348-8 Natural history of *H. pylori* infection. MALT, mucosal-associated lymphoid tissue. (Used with permission from S Suerbaum, P Michetti; *N Engl J Med* 347:1175, 2002.)

In view of the lack of warning signs, it is important to identify patients who are at increased risk for morbidity and mortality related to NSAID usage. Even 75 mg/d of aspirin may lead to serious GI ulceration; thus, no dose of NSAID is completely safe. In fact, the incidence of mucosal injury (ulcers and erosions) in patients taking low-dose aspirin (75–325 mg) has been estimated to range from as low as 8% to as high as 60%. It appears that *H. pylori* infection increases the risk of PUD-associated GI bleeding in chronic users of low-dose aspirin. Established risk factors include advanced age, history of ulcer, concomitant use of glucocorticoids, high-dose NSAIDs, multiple NSAIDs, concomitant use of anticoagulants, clopidogrel, and serious or multisystem disease. Possible risk factors include concomitant infection with *H. pylori*, cigarette smoking, and alcohol consumption.

Pathophysiology Prostaglandins play a critical role in maintaining gastroduodenal mucosal integrity and repair. It therefore follows that interruption of prostaglandin synthesis can impair mucosal defense and repair, thus facilitating mucosal injury via a systemic mechanism. Animal studies have demonstrated that neutrophil adherence to the gastric microcirculation plays an essential role in the initiation of NSAID-induced mucosal injury. A summary of the pathogenetic pathways by which systemically administered NSAIDs may lead to mucosal injury is shown in Fig. 348-9. Single nucleotide polymorphisms (SNPs) have been found in several genes, including those encoding certain subtypes of cytochrome P450 (see below), interleukin-1 β (*IL-1 β*), angiotensinogen (*AGT*), and an organic ion transporting polypeptide (*SLCO1B1*), but these findings need confirmation in larger scale studies.

Injury to the mucosa also occurs as a result of the topical encounter with NSAIDs. Aspirin and many NSAIDs are weak acids that remain in a nonionized lipophilic form when found within the acid environment of the stomach. Under these conditions, NSAIDs migrate across lipid membranes of epithelial cells, leading to cell injury once trapped intracellularly in an ionized form. Topical NSAIDs can also alter the surface mucous layer, permitting back diffusion of H⁺ and pepsin, leading to further epithelial cell damage. Moreover, enteric-coated or buffered preparations are also associated with risk of peptic ulceration.

The interplay between *H. pylori* and NSAIDs in the pathogenesis of PUD is complex. Meta-analysis supports the conclusion that each of these aggressive factors is independent and synergistic risk factors for PUD and its complications such as GI bleeding. For example, eradication of *H. pylori* reduces the likelihood of GI complications in high-risk individuals to levels observed in individuals with average risk of NSAID-induced complications.

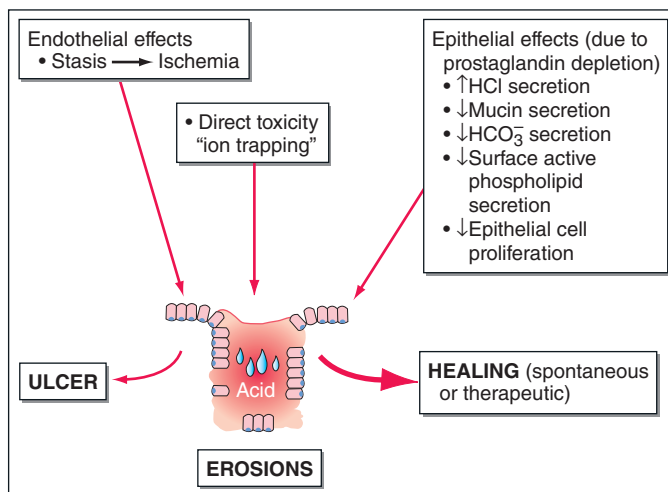


FIGURE 348-9 Mechanisms by which nonsteroidal anti-inflammatory drugs may induce mucosal injury. (Adapted from J Scheiman et al; *J Clin Outcomes Management* 3:23, 1996. Copyright 2003 Turner White Communications, Inc., www.turner-white.com. Used with permission.)

PATHOGENETIC FACTORS UNRELATED TO *H. PYLORI* AND NSAIDS IN ACID PEPTIC DISEASE Cigarette smoking has been implicated in the pathogenesis of PUD. Not only have smokers been found to have ulcers more frequently than do nonsmokers, but smoking appears to decrease healing rates, impair response to therapy, and increase ulcer-related complications such as perforation. The mechanism responsible for increased ulcer diathesis in smokers is unknown. Theories have included altered gastric emptying, decreased proximal duodenal bicarbonate production, increased risk for *H. pylori* infection, and cigarette-induced generation of noxious mucosal free radicals. Genetic predisposition may play a role in ulcer development. First-degree relatives of DU patients are three times as likely to develop an ulcer; however, the potential role of *H. pylori* infection in contacts is a major consideration. Increased frequencies of blood group O and of the nonsecretor status have also been implicated as genetic risk factors for peptic diathesis. However, *H. pylori* preferentially binds to group O antigens. Additional genetic factors have been postulated to predispose certain individuals to developing PUD and/or upper GI bleeding. Specifically, genes encoding the NSAID-metabolizing enzymes cytochrome P450 2C9 and 2C8 (*CYP2C9* and *CYP2C8*) are potential susceptibility genes for NSAID-induced PUD, but unfortunately, the studies have not been consistent in demonstrating this association. In a United Kingdom study, the *CYP2C19*17* gain-of-function polymorphism was associated with PUD in a Caucasian cohort, irrespective of ulcer etiology. These findings need to be confirmed in broader studies. Psychological stress has been thought to contribute to PUD, but studies examining the role of psychological factors in its pathogenesis have generated conflicting results. Although PUD is associated with certain personality traits (neuroticism), these same traits are also present in individuals with nonulcer dyspepsia (NUD) and other functional and organic disorders.

Diet has also been thought to play a role in peptic diseases. Certain foods and beverages can cause dyspepsia, but no convincing studies indicate an association between ulcer formation and a specific diet. Specific chronic disorders have been shown to have a strong association with PUD: (1) advanced age, (2) chronic pulmonary disease, (3) chronic renal failure, (4) cirrhosis, (5) nephrolithiasis, (6) α_1 -antitrypsin deficiency, and (7) systemic mastocytosis. Disorders with a possible association are (1) hyperparathyroidism, (2) coronary artery disease, (3) polycythemia vera, (4) chronic pancreatitis, (5) former alcohol use, (6) obesity, (7) African-American race, and (8) three or more doctor visits in a year.

Multiple factors play a role in the pathogenesis of PUD. The two predominant causes are *H. pylori* infection and NSAID ingestion. PUD

TABLE 348-1 CAUSES OF ULCERS NOT CAUSED BY *HELICOBACTER PYLORI* AND NSAIDs

Pathogenesis of Non-Hp and Non-NSAID Ulcer Disease	
Infection	
	Cytomegalovirus
	Herpes simplex virus
	<i>Helicobacter heilmannii</i>
Drug/Toxin	
	Bisphosphonates
	Chemotherapy
	Clopidogrel
	Crack cocaine
	Glucocorticoids (when combined with NSAIDs)
	Mycophenolate mofetil
	Potassium chloride
Miscellaneous	
	Basophilia in myeloproliferative disease
	Duodenal obstruction (e.g., annular pancreas)
	Infiltrating disease
	Ischemia
	Radiation therapy
	Eosinophilic infiltration
	Sarcoidosis
	Crohn's disease
	Idiopathic hypersecretory state

Abbreviations: Hp, *H. pylori*; NSAIDs, nonsteroidal anti-inflammatory drugs.

not related to *H. pylori* or NSAIDs is increasing. Other less common causes of PUD are shown in **Table 348-1**. These etiologic agents should be considered as the incidence of *H. pylori* is decreasing. Independent of the inciting or injurious agent, peptic ulcers develop as a result of an imbalance between mucosal protection/repair and aggressive factors. Gastric acid plays an important role in mucosal injury.

CLINICAL FEATURES

History Abdominal pain is common to many GI disorders, including DU and GU, but has a poor predictive value for the presence of either DU or GU. Up to 10% of patients with NSAID-induced mucosal disease can present with a complication (bleeding, perforation, and obstruction) without antecedent symptoms. Despite this poor correlation, a careful history and physical examination are essential components of the approach to a patient suspected of having peptic ulcers.

Epigastric pain described as a burning or gnawing discomfort can be present in both DU and GU. The discomfort is also described as an ill-defined, aching sensation or as hunger pain. The typical pain pattern in DU occurs 90 minutes to 3 hours after a meal and is frequently relieved by antacids or food. Pain that awakes the patient from sleep (between midnight and 3 A.M.) is the most discriminating symptom, with two-thirds of DU patients describing this complaint. Unfortunately, this symptom is also present in one-third of patients with NUD (see below). Elderly patients are less likely to have abdominal pain as a manifestation of PUD and may instead present with a complication such as ulcer bleeding or perforation. The pain pattern in GU patients may be different from that in DU patients, where discomfort may actually be precipitated by food. Nausea and weight loss occur more commonly in GU patients. Endoscopy detects ulcers in <30% of patients who have dyspepsia.

The mechanism for development of abdominal pain in ulcer patients is unknown. Several possible explanations include acid-induced activation of chemical receptors in the duodenum, enhanced duodenal sensitivity to bile acids and pepsin, or altered gastroduodenal motility.

Variation in the intensity or distribution of the abdominal pain, as well as the onset of associated symptoms such as nausea and/or

vomiting, may be indicative of an ulcer complication. Dyspepsia that becomes constant, is no longer relieved by food or antacids, or radiates to the back may indicate a penetrating ulcer (pancreas). Sudden onset of severe, generalized abdominal pain may indicate perforation. Pain worsening with meals, nausea, and vomiting of undigested food suggest gastric outlet obstruction. Tarry stools or coffee-ground emesis indicate bleeding.

Physical examination Epigastric tenderness is the most frequent finding in patients with GU or DU. Pain may be found to the right of the midline in 20% of patients. Unfortunately, the predictive value of this finding is rather low. Physical examination is critically important for discovering evidence of ulcer complication. Tachycardia and orthostasis suggest dehydration secondary to vomiting or active GI blood loss. A severely tender, board-like abdomen suggests a perforation. Presence of a succussion splash indicates retained fluid in the stomach, suggesting gastric outlet obstruction.

PUD-Related Complications • GASTROINTESTINAL BLEEDING GI bleeding is the most common complication observed in PUD. Bleeding is estimated to occur in 19.4–57 per 100,000 individuals in a general population or in approximately 15% of patients. Bleeding and complications of ulcer disease occur more often in individuals >60 years of age. The 30-day mortality rate is as high as 5–10%. The higher incidence in the elderly is likely due to the increased use of NSAIDs in this group. In addition, up to 80% of the mortality in PUD-related bleeding is due to nonbleeding causes such as multiorgan failure (24%), pulmonary complications (24%), and malignancy (34%).

Up to 20% of patients with ulcer-related hemorrhage bleed without any preceding warning signs or symptoms.

PERFORATION The second most common ulcer-related complication is perforation, being reported in as many as 6–7% of PUD patients with an estimated 30-day mortality of over 20%. As in the case of bleeding, the incidence of perforation in the elderly appears to be increasing secondary to increased use of NSAIDs. *Penetration* is a form of perforation in which the ulcer bed tunnels into an adjacent organ. DUs tend to penetrate posteriorly into the pancreas, leading to pancreatitis, whereas GUs tend to penetrate into the left hepatic lobe. Gastrocolic fistulas associated with GUs have also been described.

GASTRIC OUTLET OBSTRUCTION Gastric outlet obstruction is the least common ulcer-related complication, occurring in 1–2% of patients. A patient may have relative obstruction secondary to ulcer-related inflammation and edema in the peripyloric region. This process often resolves with ulcer healing. A fixed, mechanical obstruction secondary to scar formation in the peripyloric areas is also possible. The latter requires endoscopic (balloon dilation) or surgical intervention. Signs and symptoms relative to mechanical obstruction may develop insidiously. New onset of early satiety, nausea, vomiting, increase of postprandial abdominal pain, and weight loss should make gastric outlet obstruction a possible diagnosis.

Differential Diagnosis The list of GI and non-GI disorders that can mimic ulceration of the stomach or duodenum is quite extensive. The most commonly encountered diagnosis among patients seen for upper abdominal discomfort is NUD. NUD, also known as *functional dyspepsia* or *essential dyspepsia*, refers to a group of heterogeneous disorders typified by upper abdominal pain without the presence of an ulcer. Dyspepsia has been reported to occur in up to 30% of the U.S. population. Up to 60% of patients seeking medical care for dyspepsia have a negative diagnostic evaluation. The etiology of NUD is not established, and the potential role of *H. pylori* in NUD remains controversial.

Several additional disease processes that may present with “ulcer-like” symptoms include proximal GI tumors, gastroesophageal reflux, vascular disease, pancreaticobiliary disease (biliary colic, chronic pancreatitis), and gastroduodenal Crohn's disease.

Diagnostic Evaluation In view of the poor predictive value of abdominal pain for the presence of a gastroduodenal ulcer and the multiple disease

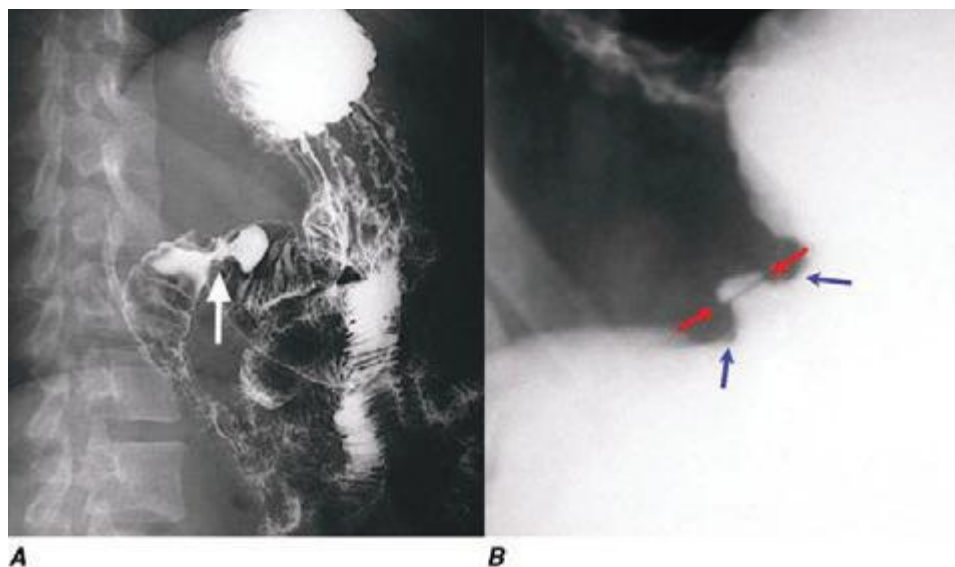


FIGURE 348-10 Barium study demonstrating (A) a benign duodenal ulcer and (B) a benign gastric ulcer.

processes that can mimic this disease, the clinician is often confronted with having to establish the presence of an ulcer. Documentation of an ulcer requires either a radiographic (barium study) or an endoscopic procedure. However, a large percentage of patients with symptoms suggestive of an ulcer have NUD; testing for *H. pylori* and antibiotic therapy (see below) is appropriate for individuals who are otherwise healthy and <45 years of age, before embarking on a diagnostic evaluation (Chap. 54).

Barium studies of the proximal GI tract are still occasionally used as a first test for documenting an ulcer. The sensitivity of older single-contrast barium meals for detecting a DU is as high as 80%, with a double-contrast study providing detection rates as high as 90%. Sensitivity for detection is decreased in small ulcers (<0.5 cm), with presence of previous scarring, or in postoperative patients. A DU appears as a well-demarcated crater, most often seen in the bulb (Fig. 348-10A). A GU may represent benign or malignant disease. Typically, a benign GU also appears as a discrete crater with radiating mucosal folds originating from the ulcer margin (Fig. 348-10B). Ulcers >3 cm in size or those associated with a mass are more often malignant. Unfortunately, up to 8% of GUs that appear to be benign by radiographic appearance are malignant by endoscopy or surgery. Radiographic studies that show a GU must be followed by endoscopy and biopsy.

Endoscopy provides the most sensitive and specific approach for examining the upper GI tract (Fig. 348-11). In addition to permitting direct visualization of the mucosa, endoscopy facilitates photographic documentation of a mucosal defect and tissue biopsy to rule out malignancy (GU) or *H. pylori*. Endoscopic examination is particularly helpful in identifying lesions too small to detect by radiographic examination, for evaluation of atypical radiographic abnormalities, or to determine if an ulcer is a source of blood loss.

Although the methods for diagnosing *H. pylori* are outlined in Chap. 181, a brief summary will be included here (Table 348-2). Several biopsy urease tests have been developed (PyloriTek, CLOtest, Hpfast, Pronto Dry) that have a sensitivity and specificity of >90–95%. Several noninvasive methods for detecting this organism have been developed. Three types of studies routinely used

include serologic testing, the ^{13}C - or ^{14}C -urea breath test, and the fecal *H. pylori* (Hp) antigen test. A urinary Hp antigen test, as well as a refined monoclonal antibody stool antigen test, appears promising.

Occasionally, specialized testing such as serum gastrin and gastric acid analysis or sham feeding may be needed in individuals with complicated or refractory PUD (see “Zollinger-Ellison Syndrome [ZES],” below). Screening for aspirin or NSAIDs (blood or urine) may also be necessary in refractory *H. pylori*-negative PUD patients.

TREATMENT PEPTIC ULCER DISEASE

Before the discovery of *H. pylori*, the therapy of PUD was centered on the old dictum by Schwartz of “no acid, no ulcer.” Although acid secretion is still important in the pathogenesis of PUD, eradication of *H. pylori* and therapy/prevention of NSAID-induced disease is the mainstay of treatment. A summary of commonly used drugs for treatment of acid peptic disorders is shown in Table 348-3.

ACID-NEUTRALIZING/INHIBITORY DRUGS

Antacids Before we understood the important role of histamine in stimulating parietal cell activity, neutralization of secreted acid with antacids constituted the main form of therapy for peptic

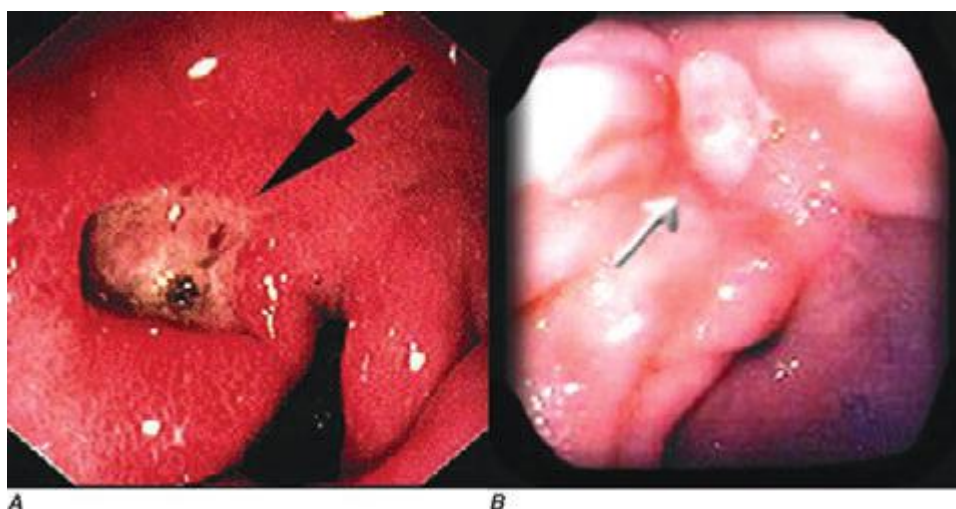


FIGURE 348-11 Endoscopy demonstrating (A) a benign duodenal ulcer and (B) a benign gastric ulcer.

TABLE 348-2 TESTS FOR DETECTION OF *H. PYLORI*

Test	Sensitivity/ Specificity, %	Comments
Invasive (Endoscopy/Biopsy Required)		
Rapid urease	80–95/95–100	Simple, false negative with recent use of PPIs, antibiotics, or bismuth compounds
Histology	80–90/>95	Requires pathology processing and staining; provides histologic information
Culture	—/—	Time-consuming, expensive, dependent on experience; allows determination of antibiotic susceptibility
Noninvasive		
Serology	>80/>90	Inexpensive, convenient; not useful for early follow-up
Urea breath test	>90/>90	Simple, rapid; useful for early follow-up; false negatives with recent therapy (see rapid urease test); exposure to low-dose radiation with ¹⁴ C test
Stool antigen	>90/>90	Inexpensive, convenient

Abbreviation: PPIs, proton pump inhibitors.

ulcers. They are now rarely, if ever, used as the primary therapeutic agent but instead are often used by patients for symptomatic relief of dyspepsia. The most commonly used agents are mixtures of aluminum hydroxide and magnesium hydroxide. Aluminum hydroxide can produce constipation and phosphate depletion; magnesium hydroxide may cause loose stools. Many of the commonly used antacids (e.g., Maalox, Mylanta) have a combination of both aluminum and magnesium hydroxide in order to avoid these side effects. The magnesium-containing preparation should not be used in chronic renal failure patients because of possible hypermagnesemia, and aluminum may cause chronic neurotoxicity in these patients.

Calcium carbonate and sodium bicarbonate are potent antacids with varying levels of potential problems. The long-term use of calcium carbonate (converts to calcium chloride in the stomach) can lead to milk-alkali syndrome (hypercalcemia, hyperphosphatemia with possible renal calcinosis and progression to renal insufficiency). Sodium bicarbonate may induce systemic alkalosis.

TABLE 348-3 DRUGS USED IN THE TREATMENT OF PEPTIC ULCER DISEASE

Drug Type/Mechanism	Examples	Dose
Acid-suppressing drugs		
Antacids	Mylanta, Maalox, Tums, Gaviscon	100–140 meq/L 1 and 3 h after meals and hs
H_2 receptor antagonists	Cimetidine	400 mg bid
	Ranitidine	300 mg hs
	Famotidine	40 mg hs
	Nizatidine	300 mg hs
Proton pump inhibitors	Omeprazole	20 mg/d
	Lansoprazole	30 mg/d
	Rabeprazole	20 mg/d
	Pantoprazole	40 mg/d
	Esomeprazole	20 mg/d
	Dexlansoprazole	30 mg/d
Mucosal protective agents		
Sucralfate	Sucralfate	1 g qid
Prostaglandin analogue	Misoprostol	200 μ g qid
Bismuth-containing compounds	Bismuth subsalicylate (BSS)	See anti- <i>H. pylori</i> regimens (Table 348-4)

Abbreviation: hs, at bedtime (*hora somni*).

H_2 Receptor Antagonists Four of these agents are presently available (cimetidine, ranitidine, famotidine, and nizatidine), and their structures share homology with histamine. Although each has different potency, all will significantly inhibit basal and stimulated acid secretion to comparable levels when used at therapeutic doses. Moreover, similar ulcer-healing rates are achieved with each drug when used at the correct dosage. Presently, this class of drug is often used for treatment of active ulcers (4–6 weeks) in combination with antibiotics directed at eradicating *H. pylori* (see below).

Cimetidine was the first H_2 receptor antagonist used for the treatment of acid peptic disorders. The initial recommended dosing profile for cimetidine was 300 mg qid. Subsequent studies have documented the efficacy of using 800 mg at bedtime for treatment of active ulcer, with healing rates approaching 80% at 4 weeks. Cimetidine may have weak antiandrogenic side effects resulting in reversible gynecomastia and impotence, primarily in patients receiving high doses for prolonged periods of time (months to years, as in ZES). In view of cimetidine's ability to inhibit cytochrome P450, careful monitoring of drugs such as warfarin, phenytoin, and theophylline is indicated with long-term usage. Other rare reversible adverse effects reported with cimetidine include confusion and elevated levels of serum aminotransferases, creatinine, and serum prolactin. Ranitidine, famotidine, and nizatidine are more potent H_2 receptor antagonists than cimetidine. Each can be used once a day at bedtime for ulcer prevention, which was commonly done before the discovery of *H. pylori* and the development of proton pump inhibitors (PPIs). Patients may develop tolerance to H_2 blockers, a rare event with PPIs (see below). Comparable nighttime dosing regimens are ranitidine 300 mg, famotidine 40 mg, and nizatidine 300 mg.

Additional rare, reversible systemic toxicities reported with H_2 receptor antagonists include pancytopenia, neutropenia, anemia, and thrombocytopenia, with a prevalence rate varying from 0.01–0.2%. Cimetidine and ranitidine (to a lesser extent) can bind to hepatic cytochrome P450; famotidine and nizatidine do not.

Proton Pump (H^+ , K^+ -ATPase) Inhibitors Omeprazole, esomeprazole, lansoprazole, rabeprazole, and pantoprazole are substituted benzimidazole derivatives that covalently bind and irreversibly inhibit H^+ , K^+ -ATPase. Esomeprazole, one of the newest members of this drug class, is the S-enantiomer of omeprazole, which is a racemic mixture of both S- and R-optical isomers. The R-isomer of lansoprazole, dexlansoprazole, is the most recent PPI approved for clinical use. Its reported advantage is a dual delayed-release system, aimed at improving treatment of gastroesophageal reflux disease (GERD). These are the most potent acid inhibitory agents available. Omeprazole and lansoprazole are the PPIs that have been used for the longest time. Both are acid-labile and are administered as enteric-coated granules in a sustained-release capsule that dissolves within the small intestine at a pH of 6. Lansoprazole is available in an orally disintegrating tablet that can be taken with or without water, an advantage for individuals who have significant dysphagia. Absorption kinetics are similar to the capsule. In addition, a lansoprazole-naproxen combination preparation that has been made available is targeted at decreasing NSAID-related GI injury (see below). Omeprazole is available as nonenteric-coated granules mixed with sodium bicarbonate in a powder form that can be administered orally or via gastric tube. The sodium bicarbonate has two purposes: to protect the omeprazole from acid degradation and to promote rapid gastric alkalization and subsequent proton pump activation, which facilitates rapid action of the PPI. Pantoprazole and rabeprazole are available as enteric-coated tablets. Pantoprazole is also available as a parenteral formulation for intravenous use. These agents are lipophilic compounds; upon entering the parietal cell, they are protonated and trapped within the acid environment of the tubulovesicular and canalicular system. These agents potently inhibit all phases of gastric acid secretion. Onset of action is rapid, with a maximum acid inhibitory effect between 2 and 6 h after administration and

duration of inhibition lasting up to 72–96 h. With repeated daily dosing, progressive acid inhibitory effects are observed, with basal and secretagogue-stimulated acid production being inhibited by >95% after 1 week of therapy. The half-life of PPIs is ~18 h; thus, it can take between 2 and 5 days for gastric acid secretion to return to normal levels once these drugs have been discontinued. Because the pumps need to be activated for these agents to be effective, their efficacy is maximized if they are administered before a meal (except for the immediate-release formulation of omeprazole) (e.g., in the morning before breakfast). Mild to moderate hypergastrinemia has been observed in patients taking these drugs. Carcinoid tumors developed in some animals given the drugs preclinically; however, extensive experience has failed to demonstrate gastric carcinoid tumor development in humans. Serum gastrin levels return to normal levels within 1–2 weeks after drug cessation. Rebound gastric acid hypersecretion has been described in *H. pylori*-negative individuals after discontinuation of PPIs. It occurs even after relatively short-term usage (2 months) and may last for up to 2 months after the PPI has been discontinued. The mechanism involves gastrin-induced hyperplasia and hypertrophy of histamine-secreting ECL cells. The clinical relevance of this observation is that individuals may have worsening symptoms of GERD or dyspepsia upon stopping the PPI. Gradual tapering of the PPI and switching to an H₂ receptor antagonist may prevent this from occurring. *H. pylori*-induced inflammation and concomitant decrease in acid production may explain why this does not occur in *H. pylori*-positive patients. IF production is also inhibited, but vitamin B₁₂-deficiency anemia is uncommon, probably because of the large stores of the vitamin. As with any agent that leads to significant hypochlorhydria, PPIs may interfere with absorption of drugs such as ketoconazole, ampicillin, iron, and digoxin. Hepatic cytochrome P450 can be inhibited by the earlier PPIs (omeprazole, lansoprazole). Rabeprazole, pantoprazole, and esomeprazole do not appear to interact significantly with drugs metabolized by the cytochrome P450 system. The overall clinical significance of this observation is not definitely established. Caution should be taken when using theophylline, warfarin, diazepam, atazanavir, and phenytoin concomitantly with PPIs. Long-term acid suppression, especially with PPIs, has been associated with a higher incidence of community-acquired pneumonia as well as community and hospital acquired *Clostridium difficile*-associated disease. These observations require confirmation but should alert the practitioner to take caution when recommending these agents for long-term use, especially in elderly patients at risk for developing pneumonia or *C. difficile* infection. A population-based study revealed that long-term use of PPIs was associated with the development of hip fractures in older women. The absolute risk of fracture remained low despite an observed increase associated with the dose and duration of acid suppression. The mechanism for this observation is not clear, and this finding must be confirmed before making broad recommendations regarding the discontinuation of these agents in patients who benefit from them. Long-term use of PPIs has also been implicated in the development of iron and magnesium deficiency, but here again, the studies are limited and inconclusive. PPIs may exert a negative effect on the antiplatelet effect of clopidogrel. Although the evidence is mixed and inconclusive, a small increase in mortality and readmission rate for coronary events was seen in patients receiving a PPI while on clopidogrel in earlier studies. Subsequently, three meta-analyses reported an inverse correlation between clopidogrel and PPI use; therefore, the influence of this drug interaction on mortality is not clearly established. The mechanism involves the competition of the PPI and clopidogrel with the same cytochrome P450 (CYP2C19). Whether this is a class effect of PPIs is unclear; there appears to be at least a theoretical advantage of pantoprazole over the other PPIs, but this has not been confirmed. This drug interaction is particularly relevant in light of the common use of aspirin and clopidogrel for prevention of coronary events and the efficacy of PPIs in preventing GI bleeding in these patients. The FDA has made several recommendations while awaiting further evidence to clarify the impact of PPI therapy on clopidogrel use. Health care providers should

continue to prescribe clopidogrel to patients who require it and should reevaluate the need for starting or continuing treatment with a PPI. From a practical standpoint, additional recommendations to consider include the following: Patients taking clopidogrel with aspirin, especially with other GI risk factors for bleeding, should receive GI protective therapy. Although high-dose H₂ blockers have been considered an option, these do not appear to be as effective as PPIs. If PPIs are to be given, some have recommended that there be a 12-h separation between administration of the PPI and clopidogrel to minimize competition of the two agents with the involved cytochrome P450. One option is to give the PPI 30 min before breakfast and the clopidogrel at bedtime. Insufficient data are available to firmly recommend one PPI over another. Patients 65 years of age or older have a higher risk for some of the long-term side effects of PPIs highlighted above, in part due to the higher prevalence of concomitant chronic diseases. It is therefore important to carefully select individuals, especially among the elderly, who need long-term PPI therapy and discontinue it in those individuals who do not need it.

Two new formulations of acid inhibitory agents are being developed. Tenatoprazole is a PPI containing an imidazopyridine ring instead of a benzimidazole ring, which promotes irreversible proton pump inhibition. This agent has a longer half-life than the other PPIs and may be beneficial for inhibiting nocturnal acid secretion, which has significant relevance in GERD. A second new class of agents is the potassium-competitive acid pump antagonists (P-CABs). These compounds inhibit gastric acid secretion via potassium competitive binding of the H⁺,K⁺-ATPase.

CYTOPROTECTIVE AGENTS

Sucralfate Sucralfate is a complex sucrose salt in which the hydroxyl groups have been substituted by aluminum hydroxide and sulfate. This compound is insoluble in water and becomes a viscous paste within the stomach and duodenum, binding primarily to sites of active ulceration. Sucralfate may act by several mechanisms: serving as a physicochemical barrier, promoting a trophic action by binding growth factors such as EGF, enhancing prostaglandin synthesis, stimulating mucus and bicarbonate secretion, and enhancing mucosal defense and repair. Toxicity from this drug is rare, with constipation being most common (2–3%). It should be avoided in patients with chronic renal insufficiency to prevent aluminum-induced neurotoxicity. Hypophosphatemia and gastric bezoar formation have also been reported rarely. Standard dosing of sucralfate is 1 g qid.

Bismuth-Containing Preparations Sir William Osler considered bismuth-containing compounds the drug of choice for treating PUD. The resurgence in the use of these agents is due to their effect against *H. pylori*. Colloidal bismuth subcitrate (CBS) and bismuth subsalicylate (BSS, Pepto-Bismol) are the most widely used preparations. The mechanism by which these agents induce ulcer healing is unclear. Adverse effects with short-term use include black stools, constipation, and darkening of the tongue. Long-term use with high doses, especially with the avidly absorbed CBS, may lead to neurotoxicity. These compounds are commonly used as one of the agents in an anti-*H. pylori* regimen (see below).

Prostaglandin Analogues In view of their central role in maintaining mucosal integrity and repair, stable prostaglandin analogues were developed for the treatment of PUD. The mechanism by which this rapidly absorbed drug provides its therapeutic effect is through enhancement of mucosal defense and repair. The most common toxicity noted with this drug is diarrhea (10–30% incidence). Other major toxicities include uterine bleeding and contractions; misoprostol is contraindicated in women who may be pregnant, and women of childbearing age must be made clearly aware of this potential drug toxicity. The standard therapeutic dose is 200 µg qid.

Miscellaneous Drugs A number of drugs including anticholinergic agents and tricyclic antidepressants were used for treating acid

1922 peptic disorders, but in light of their toxicity and the development of potent antisecretory agents, these are rarely, if ever, used today.

THErapy OF *H. PYLORI*

The physician's goal in treating PUD is to provide relief of symptoms (pain or dyspepsia), promote ulcer healing, and ultimately prevent ulcer recurrence and complications. The greatest influence of understanding the role of *H. pylori* in peptic disease has been the ability to prevent recurrence. Documented eradication of *H. pylori* in patients with PUD is associated with a dramatic decrease in ulcer recurrence to <10–20% as compared to 59% in GU patients and 67% in DU patients when the organism is not eliminated. Eradication of the organism may lead to diminished recurrent ulcer bleeding. The effect of its eradication on ulcer perforation is unclear.

Extensive effort has been made in determining who of the many individuals with *H. pylori* infection should be treated. The common conclusion arrived at by multiple consensus conferences around the world is that *H. pylori* should be eradicated in patients with documented PUD. This holds true independent of time of presentation (first episode or not), severity of symptoms, presence of confounding factors such as ingestion of NSAIDs, or whether the ulcer is in remission. Some have advocated treating patients with a history of documented PUD who are found to be *H. pylori*-positive by serology or breath testing. Over one-half of patients with gastric MALT lymphoma experience complete remission of the tumor in response to *H. pylori* eradication. The Maastricht IV/Florence Consensus Report recommends a test-and-treat approach for patients with uninvestigated dyspepsia if the local incidence of *H. pylori* is greater than 20%. In addition, recommendations from this consensus report include testing and eradicating *H. pylori* in patients who will be using NSAIDs (including low-dose aspirin) on a long-term basis, especially if there is a prior history of PUD. These individuals will require continued PPI treatment as well as eradication treatment, because eradication of the organism alone does not eliminate the risk of gastroduodenal ulcers in patients already receiving long-term NSAIDs. Treating patients with NUD to prevent gastric cancer or patients with GERD requiring long-term acid suppression remains controversial. Guidelines from the American College of Gastroenterology suggest eradication of *H. pylori* in patients who have undergone resection of early gastric cancer. The Maastricht IV/Florence Consensus Report also evaluated *H. pylori* treatment in gastric cancer prevention and recommends that eradication should be considered in the following situations: first-degree relatives of family members with gastric cancer; patients with previous gastric neoplasm treated by endoscopic or subtotal resection; individuals with a risk of gastritis (severe pangastritis or body-predominant gastritis) or severe atrophy; patients with gastric acid inhibition for more than 1 year; individuals with strong environmental risk factors for gastric cancer (heavy smoking; high exposure to dust, coal, quartz, or cement; and/or work in quarries); and *H. pylori*-positive patients with a fear of gastric cancer.

Multiple drugs have been evaluated in the therapy of *H. pylori*. No single agent is effective in eradicating the organism. Combination therapy for 14 days provides the greatest efficacy, although regimens based on sequential administration of antibiotics also appear promising (see below). A shorter administration course (7–10 days), although attractive, has not proved as successful as the 14-day regimens. The agents used with the greatest frequency include amoxicillin, metronidazole, tetracycline, clarithromycin, and bismuth compounds.

Suggested treatment regimens for *H. pylori* are outlined in **Table 348-4**. Choice of a particular regimen will be influenced by several factors, including efficacy, patient tolerance, existing antibiotic resistance, and cost of the drugs. The aim for initial eradication rates should be 85–90%. Dual therapy (PPI plus amoxicillin, PPI plus clarithromycin, ranitidine bismuth citrate [Tritec] plus clarithromycin) is not recommended in view of studies demonstrating eradication rates of <80–85%. The combination of bismuth, metronidazole, and tetracycline was the first triple regimen found effective against

TABLE 348-4 REGIMENS RECOMMENDED FOR ERADICATION OF *H. PYLORI* INFECTION

Drug	Dose
Triple Therapy	
1. Bismuth subsalicylate <i>plus</i> Metronidazole <i>plus</i> Tetracycline ^a	2 tablets qid 250 mg qid 500 mg qid
2. Ranitidine bismuth citrate <i>plus</i> Tetracycline <i>plus</i> Clarithromycin or metronidazole	400 mg bid 500 mg bid 500 mg bid
3. Omeprazole (lansoprazole) <i>plus</i> Clarithromycin <i>plus</i> Metronidazole ^b <i>or</i> Amoxicillin ^c	20 mg bid (30 mg bid) 250 or 500 mg bid 500 mg bid 1 g bid
Quadruple Therapy	
Omeprazole (lansoprazole)	20 mg (30 mg) daily
Bismuth subsalicylate	2 tablets qid
Metronidazole	250 mg qid
Tetracycline	500 mg qid

^aAlternative: use prepacked Helidac (see text). ^bAlternative: use prepacked Prevpac (see text). ^cUse either metronidazole or amoxicillin, not both.

H. pylori. The combination of two antibiotics plus either a PPI, H₂ blocker, or bismuth compound has comparable success rates. Addition of acid suppression assists in providing early symptom relief and enhances bacterial eradication.

Triple therapy, although effective, has several drawbacks, including the potential for poor patient compliance and drug-induced side effects. Compliance is being addressed by simplifying the regimens so that patients can take the medications twice a day. Simpler (dual therapy) and shorter regimens (7 and 10 days) are not as effective as triple therapy for 14 days. Two anti-*H. pylori* regimens are available in prepackaged formulation: Prevpac (lansoprazole, clarithromycin, and amoxicillin) and Helidac (BSS, tetracycline, and metronidazole). The contents of the Prevpac are to be taken twice per day for 14 days, whereas Helidac constituents are taken four times per day with an antisecretory agent (PPI or H₂ blocker), also for at least 14 days. Clarithromycin-based triple therapy should be avoided in settings where *H. pylori* resistance to this agent exceeds 15–20%.

Side effects have been reported in up to 20–30% of patients on triple therapy. Bismuth may cause black stools, constipation, or darkening of the tongue. The most feared complication with amoxicillin is pseudomembranous colitis, but this occurs in <1–2% of patients. Amoxicillin can also lead to antibiotic-associated diarrhea, nausea, vomiting, skin rash, and allergic reaction. Concomitant use of probiotics may ameliorate some of the antibiotic side effects (see below). Tetracycline has been reported to cause rashes and, very rarely, hepatotoxicity and anaphylaxis.

One important concern with treating patients who may not need therapy is the potential for development of antibiotic-resistant strains. The incidence and type of antibiotic-resistant *H. pylori* strains vary worldwide. Strains resistant to metronidazole, clarithromycin, amoxicillin, and tetracycline have been described, with the latter two being uncommon. Antibiotic-resistant strains are the most common cause for treatment failure in compliant patients. Unfortunately, in vitro resistance does not predict outcome in patients. Culture and sensitivity testing of *H. pylori* is not performed routinely. Although resistance to metronidazole has been found in as many as 30% of isolates in North America and 80% in developing countries, triple therapy is effective in eradicating the organism in >50% of patients infected with a resistant strain. Clarithromycin resistance is seen in 13% of individuals in the United States, with resistance to amoxicillin being <1% and resistance to both metronidazole and clarithromycin in the 5% range.

Failure of *H. pylori* eradication with triple therapy in a compliant patient is usually due to infection with a resistant organism. Quadruple therapy (Table 348-4), where clarithromycin is substituted for metronidazole (or vice versa), should be the next step. The combination of pantoprazole, amoxicillin, and rifabutin for 10 days has also been used successfully (86% cure rate) in patients infected with resistant strains. Additional regimens considered for second-line therapy include levofloxacin-based triple therapy (levofloxacin, amoxicillin, PPI) for 10 days and furazolidone-based triple therapy (furazolidone, amoxicillin, PPI) for 14 days. Unfortunately, there is no universally accepted treatment regimen recommended for patients who have failed two courses of antibiotics. If eradication is still not achieved in a compliant patient, then culture and sensitivity of the organism should be considered. Additional factors that may lower eradication rates include the patient's country of origin (higher in Northeast Asia than other parts of Asia or Europe) and cigarette smoking. In addition, meta-analysis suggests that even the most effective regimens (quadruple therapy including PPI, bismuth, tetracycline, and metronidazole and triple therapy including PPI, clarithromycin, and amoxicillin) may have suboptimal eradication rates (<80%), thus demonstrating the need for the development of more efficacious treatments.

In view of the observation that 15–25% of patients treated with first-line therapy may still remain infected with the organism, new approaches to treatment have been explored. One promising approach is sequential therapy. Regimens examined consist of 5 days of amoxicillin and a PPI, followed by an additional 5 days of PPI plus tinidazole and clarithromycin or levofloxacin. One promising regimen that has the benefit of being shorter in duration, easier to take, and less expensive is 5 days of concomitant therapy (PPI twice daily, amoxicillin 1 g twice daily, levofloxacin 500 mg twice daily, and tinidazole 500 mg twice daily). Initial studies have demonstrated eradication rates of >90% with good patient tolerance. Confirmation of these findings and applicability of this approach in the United States are needed, although some experts are recommending abandoning clarithromycin-based triple therapy in the United States for the concomitant therapy or the alternative sequential therapies highlighted above.

Innovative non-antibiotic-mediated approaches have been explored in an effort to improve eradication rates of *H. pylori*. Pretreatment of patients with *N*-acetylcysteine as a mucolytic agent to destroy the *H. pylori* biofilm and therefore impair antibiotic resistance has been examined, but more studies are needed to confirm the applicability of this approach. In vitro studies suggest that certain probiotics like *Lactobacillus* or its metabolites can inhibit *H. pylori*. Administration of probiotics has been attempted in several clinical studies in an effort to maximize antibiotic-mediated eradication with varying results. Overall, it appears that the use of certain probiotics, such as *Lactobacillus* spp., *Saccharomyces* spp., *Bifidobacterium* spp., and *Bacillus clausii*, did not alter eradication rates but importantly decreased antibiotic-associated side effects including nausea, dysgeusia, diarrhea, and abdominal discomfort/pain, resulting in enhanced tolerability of *H. pylori* therapies. Additional studies are needed to confirm the potential benefits of probiotics in this setting.

Reinfection after successful eradication of *H. pylori* is rare in the United States (<1% per year). If recurrent infection occurs within the first 6 months after completing therapy, the most likely explanation is recrudescence as opposed to reinfection.

THERAPY OF NSAID-RELATED GASTRIC OR DUODENAL INJURY

Medical intervention for NSAID-related mucosal injury includes treatment of an active ulcer and primary prevention of future injury. Recommendations for the treatment and primary prevention of NSAID-related mucosal injury are listed in Table 348-5. Ideally, the injurious agent should be stopped as the first step in the therapy of an active NSAID-induced ulcer. If that is possible, then treatment with one of the acid inhibitory agents (H_2 blockers, PPIs) is indicated. Cessation of NSAIDs is not always possible because of the patient's

TABLE 348-5 RECOMMENDATIONS FOR TREATMENT OF NSAID-RELATED MUCOSAL INJURY

Clinical Setting	Recommendation
Active ulcer	
NSAID discontinued	H_2 receptor antagonist or PPI
NSAID continued	PPI
Prophylactic therapy	Misoprostol PPI Selective COX-2 inhibitor
<i>H. pylori</i> infection	Eradication if active ulcer present or there is a past history of peptic ulcer disease

Abbreviations: COX-2, isoenzyme of cyclooxygenase; NSAID, nonsteroidal anti-inflammatory drug; PPI, proton pump inhibitor.

severe underlying disease. Only PPIs can heal GUs or DUs, independent of whether NSAIDs are discontinued.

The approach to primary prevention has included avoiding the agent, using the lowest possible dose of the agent, using NSAIDs that are theoretically less injurious, using newer topical NSAID preparations, and/or using concomitant medical therapy to prevent NSAID-induced injury. Several nonselective NSAIDs that are associated with a lower likelihood of GI toxicity include diclofenac, aceclofenac, and ibuprofen, although the beneficial effect may be eliminated if higher dosages of the agents are used. Primary prevention of NSAID-induced ulceration can be accomplished by misoprostol (200 μ g qid) or a PPI. High-dose H_2 blockers (famotidine, 40 mg bid) have also shown some promise in preventing endoscopically documented ulcers, although PPIs are superior. The highly selective COX-2 inhibitors, celecoxib and rofecoxib, are 100 times more selective inhibitors of COX-2 than standard NSAIDs, leading to gastric or duodenal mucosal injury that is comparable to placebo; their utilization led to an increase in cardiovascular events and withdrawal from the market. Additional caution was engendered when the CLASS study demonstrated that the advantage of celecoxib in preventing GI complications was offset when low-dose aspirin was used simultaneously. Therefore, gastric protection therapy is required in individuals taking COX-2 inhibitors and aspirin prophylaxis. Finally, much of the work demonstrating the benefit of COX-2 inhibitors and PPIs on GI injury has been performed in individuals of average risk; it is unclear if the same level of benefit will be achieved in high-risk patients. For example, concomitant use of warfarin and a COX-2 inhibitor was associated with rates of GI bleeding similar to those observed in patients taking nonselective NSAIDs. A combination of factors, including withdrawal of the majority of COX-2 inhibitors from the market, the observation that low-dose aspirin appears to diminish the beneficial effect of COX-2 selective inhibitors, and the growing use of aspirin for prophylaxis of cardiovascular events, have significantly altered the approach to gastric protective therapy during the use of NSAIDs. A set of guidelines for the approach to the use of NSAIDs was published by the American College of Gastroenterology and is shown in Table 348-6. Individuals who are not at risk for cardiovascular events, do not use aspirin, and are without risk for GI complications can receive nonselective NSAIDs without gastric protection. In those without cardiovascular risk factors but with a high potential risk (prior GI bleeding or multiple GI risk factors) for NSAID-induced GI toxicity, cautious use of a selective COX-2 inhibitor and co-therapy with misoprostol or high-dose PPI are recommended. Individuals at moderate GI risk without cardiac risk factors can be treated with a COX-2 inhibitor alone or with a nonselective NSAID with misoprostol or a PPI. Individuals with cardiovascular risk factors, who require low-dose aspirin and have low potential for NSAID-induced toxicity, should be considered for a non-NSAID agent or use of a traditional NSAID in combination with gastric protection, if warranted. Finally, individuals with cardiovascular and GI risks who require aspirin must be considered for non-NSAID therapy, but if that is not an option, then gastric protection with any type of NSAID must be considered. Any patient, regardless

	No/Low NSAID GI Risk	NSAID GI Risk
No CV risk (no aspirin)	Traditional NSAID	Coxib or Traditional NSAID + PPI or misoprostol Consider non-NSAID therapy
CV risk (consider aspirin)	Traditional NSAID + PPI or misoprostol if GI risk warrants gastroprotection Consider non-NSAID therapy	A gastroprotective agent must be added if a traditional NSAID is prescribed Consider non-NSAID therapy

Abbreviations: CV, cardiovascular; GI, gastrointestinal; NSAID, nonsteroidal anti-inflammatory drug; PPI, proton pump inhibitor.

Source: Adapted from AM Fendrick: Am J Manag Care 10:740, 2004. Reproduced with permission of INTELLISPHERE, LLC via Copyright Clearance Center.

of risk status, who is being considered for long-term traditional NSAID therapy, should also be considered for *H. pylori* testing and treatment if positive. Assuring the use of GI protective agents with NSAIDs is difficult, even in high-risk patients. This is in part due to underprescribing of the appropriate protective agent; other times the difficulty is related to patient compliance. The latter may be due to patients forgetting to take multiple pills or preferring not to take the extra pill, especially if they have no GI symptoms. Several NSAID gastroprotective-containing combination pills are now commercially available, including double-dose famotidine with ibuprofen, diclofenac with misoprostol, and naproxen with esomeprazole. Although initial studies suggested improved compliance and a cost advantage when taking these combination drugs, their clinical benefit over the use of separate pills has not been established. Efforts continue toward developing safer NSAIDs, including NO-releasing NSAIDs, hydrogen sulfide-releasing NSAIDs, dual COX/5-LOX inhibitors, NSAID prodrugs, or agents that can effectively sequester unbound NSAIDs without interfering with their efficacy.

APPROACH AND THERAPY: SUMMARY

Controversy continues regarding the best approach to the patient who presents with dyspepsia (Chap. 54). The discovery of *H. pylori* and its role in pathogenesis of ulcers has added a new variable to the equation. Previously, if a patient <50 years of age presented with dyspepsia and without alarming signs or symptoms suggestive of an ulcer complication or malignancy, an empirical therapeutic trial with acid suppression was commonly recommended. Although this approach is practiced by some today, an approach presently gaining approval for the treatment of patients with dyspepsia is outlined in Fig. 348-12. The referral to a gastroenterologist is for the potential need of endoscopy and subsequent evaluation and treatment if the endoscopy is negative.

Once an ulcer (GU or DU) is documented, the main issue at stake is whether *H. pylori* or an NSAID is involved. With *H. pylori* present, independent of the NSAID status, triple therapy is recommended for 14 days, followed by continued acid-suppressing drugs (H_2 receptor antagonist or PPIs) for a total of 4–6 weeks. Selection of patients for documentation of *H. pylori* eradication (organisms gone at least 4 weeks after completing antibiotics) is an area of some debate. The test of choice for documenting eradication is the laboratory-based validated monoclonal stool antigen test or a urea breath test (UBT). The patient must be off antisecretory agents when being tested for eradication of *H. pylori* with UBT or stool antigen. Serologic testing is not useful for the purpose of documenting eradication because antibody titers fall slowly and often do not become undetectable. Two approaches toward documentation of eradication exist: (1) Test for eradication only in individuals with a complicated course or in individuals who are frail or with multisystem disease who would do poorly with an ulcer recurrence, and (2) test all patients for

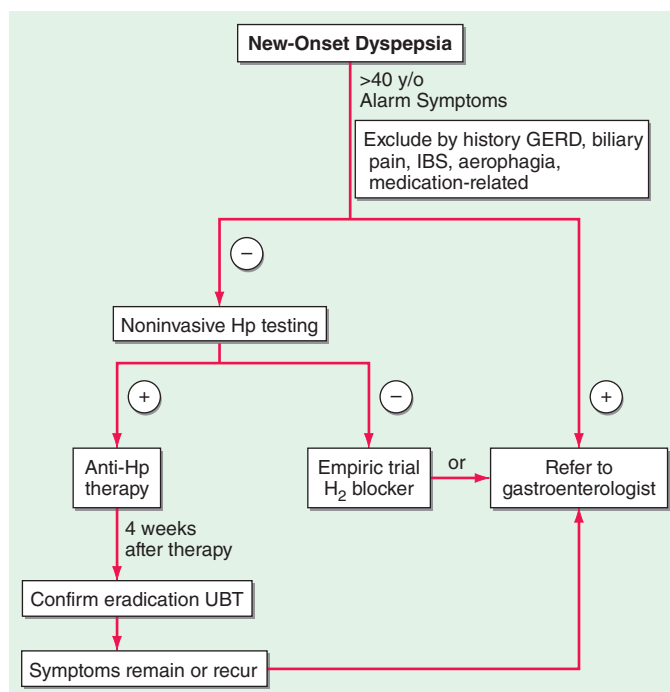


FIGURE 348-12 Overview of new-onset dyspepsia. GERD, gastroesophageal reflux disease; Hp, *Helicobacter pylori*; IBS, irritable bowel syndrome; UBT, urea breath test. (Adapted from BS Anand and DY Graham: Endoscopy 31:215, 1999.)

successful eradication. Some recommend that patients with complicated ulcer disease, or who are frail, should be treated with long-term acid suppression, thus making documentation of *H. pylori* eradication a moot point. In view of this discrepancy in practice, it would be best to discuss with the patient the different options available.

Several issues differentiate the approach to a GU versus a DU. GUs, especially of the body and fundus, have the potential of being malignant. Multiple biopsies of a GU should be taken initially; even if these are negative for neoplasm, repeat endoscopy to document healing at 8–12 weeks should be performed, with biopsy if the ulcer is still present. About 70% of GUs eventually found to be malignant undergo significant (usually incomplete) healing. Repeat endoscopy is warranted in patients with DU if symptoms persist despite medical therapy or a complication is suspected.

The majority (>90%) of GUs and DUs heal with the conventional therapy outlined above. A GU that fails to heal after 12 weeks and a DU that does not heal after 8 weeks of therapy should be considered refractory. Once poor compliance and persistent *H. pylori* infection have been excluded, NSAID use, either inadvertent or surreptitious, must be excluded. In addition, cigarette smoking must be eliminated. For a GU, malignancy must be meticulously excluded. Next, consideration should be given to a gastric acid hypersecretory state such as ZES (see "Zollinger-Ellison Syndrome," below) or the idiopathic form, which can be excluded with gastric acid analysis. Although a subset of patients have gastric acid hypersecretion of unclear etiology as a contributing factor to refractory ulcers, ZES should be excluded with a fasting gastrin or secretin stimulation test (see below). More than 90% of refractory ulcers (either DUs or GUs) heal after 8 weeks of treatment with higher doses of PPI (omeprazole, 40 mg/d; lansoprazole 30–60 mg/d). This higher dose is also effective in maintaining remission. Surgical intervention may be a consideration at this point; however, other rare causes of refractory ulcers must be excluded before recommending surgery. Rare etiologies of refractory ulcers that may be diagnosed by gastric or duodenal biopsies include ischemia, Crohn's disease, amyloidosis, sarcoidosis, lymphoma, eosinophilic gastroenteritis, or infection (cytomegalovirus [CMV], tuberculosis, or syphilis).

SURGICAL THERAPY

Surgical intervention in PUD can be viewed as being either elective, for treatment of medically refractory disease, or as urgent/emergent, for the treatment of an ulcer-related complication. The development of pharmacologic and endoscopic approaches for the treatment of peptic disease and its complications has led to a substantial decrease in the number of operations needed for this disorder with a drop of over 90% for elective ulcer surgery over the last four decades. Refractory ulcers are an exceedingly rare occurrence. Surgery is more often required for treatment of an ulcer-related complication.

Hemorrhage is the most common ulcer-related complication, occurring in ~15–25% of patients. Bleeding may occur in any age group but is most often seen in older patients (sixth decade or beyond). The majority of patients stop bleeding spontaneously, but endoscopic therapy (**Chap. 345**) is necessary in some. Parenterally and orally administered PPIs also decrease ulcer rebleeding in patients who have undergone endoscopic therapy. Patients unresponsive or refractory to endoscopic intervention will require surgery (~5% of transfusion-requiring patients).

Free peritoneal perforation occurs in ~2–3% of DU patients. As in the case of bleeding, up to 10% of these patients will not have antecedent ulcer symptoms. Concomitant bleeding may occur in up to 10% of patients with perforation, with mortality being increased substantially. Peptic ulcer can also penetrate into adjacent organs, especially with a posterior DU, which can penetrate into the pancreas, colon, liver, or biliary tree.

Pyloric channel ulcers or DUs can lead to gastric outlet obstruction in ~2–3% of patients. This can result from chronic scarring or from impaired motility due to inflammation and/or edema with pylorospasm. Patients may present with early satiety, nausea, vomiting of undigested food, and weight loss. Conservative management with nasogastric suction, intravenous hydration/nutrition, and antisecretory agents is indicated for 7–10 days with the hope that a functional obstruction will reverse. If a mechanical obstruction persists, endoscopic intervention with balloon dilation may be effective. Surgery should be considered if all else fails.

SPECIFIC OPERATIONS FOR DUODENAL ULCERS

Surgical treatment was originally designed to decrease gastric acid secretion. Operations most commonly performed include (1) vagotomy and drainage (by pyloroplasty, gastroduodenostomy, or gastrojejunostomy), (2) highly selective vagotomy (which does not require a drainage procedure), and (3) vagotomy with antrectomy. The specific procedure performed is dictated by the underlying circumstances: elective versus emergency, the degree and extent of duodenal ulceration, the etiology of the ulcer (*H. pylori*, NSAIDs, malignancy), and the expertise of the surgeon. Moreover, the trend has been toward a dramatic decrease in the need for surgery for treatment of refractory PUD, and when needed, minimally invasive and anatomy-preserving operations are preferred.

Vagotomy is a component of each of these procedures and is aimed at decreasing acid secretion through ablating cholinergic input to the stomach. Unfortunately, both truncal and selective vagotomy (preserves the celiac and hepatic branches) result in gastric atony despite successful reduction of both basal acid output (BAO; decreased by 85%) and maximal acid output (MAO; decreased by 50%). Drainage through pyloroplasty or gastroduodenostomy is required in an effort to compensate for the vagotomy-induced gastric motility disorder. This procedure has an intermediate complication rate and a 10% ulcer recurrence rate. To minimize gastric dysmotility, highly selective vagotomy (also known as parietal cell, super-selective, or proximal vagotomy) was developed. Only the vagal fibers innervating the portion of the stomach that contains parietal cells is transected, thus leaving fibers important for regulating gastric motility intact. Although this procedure leads to an immediate decrease in both BAO and stimulated acid output, acid secretion recovers over time. By the end of the first postoperative year, basal and stimulated acid output are ~30 and 50%,

respectively, of preoperative levels. Ulcer recurrence rates are higher with highly selective vagotomy ($\geq 10\%$), although the overall complication rates are the lowest of the three procedures.

The procedure that provides the lowest rates of ulcer recurrence (1%) but has the highest complication rate is vagotomy (truncal or selective) in combination with antrectomy. Antrectomy is aimed at eliminating an additional stimulant of gastric acid secretion, gastrin. Two principal types of reanastomoses are used after antrectomy: gastroduodenostomy (Billroth I) or gastrojejunostomy (Billroth II) (**Fig. 348-13**). Although Billroth I is often preferred over II, severe duodenal inflammation or scarring may preclude its performance. Prospective, randomized studies confirm that partial gastrectomy followed by Roux-en-Y reconstruction leads to a significantly better clinical, endoscopic, and histologic outcome than Billroth II reconstruction.

Of these procedures, highly selective vagotomy may be the one of choice in the elective setting, except in situations where ulcer recurrence rates are high (prepyloric ulcers and those refractory to medical therapy). Selection of vagotomy and antrectomy may be more appropriate in these circumstances.

These procedures have been traditionally performed by standard laparotomy. The advent of laparoscopic surgery has led several surgical teams to successfully perform highly selective vagotomy, truncal vagotomy/pyloroplasty, and truncal vagotomy/antrectomy through this approach. An increase in the number of laparoscopic procedures for treatment of PUD has occurred. Laparoscopic repair of perforated peptic ulcers is safe, feasible for the experienced surgeon and is associated with decreased postoperative pain, although it does take longer than an open approach. Moreover, no difference between the two approaches is noted in postoperative complications or length of hospital stay.

Specific Operations for Gastric Ulcers The location and the presence of a concomitant DU dictate the operative procedure performed for a GU. Antrectomy (including the ulcer) with a Billroth I

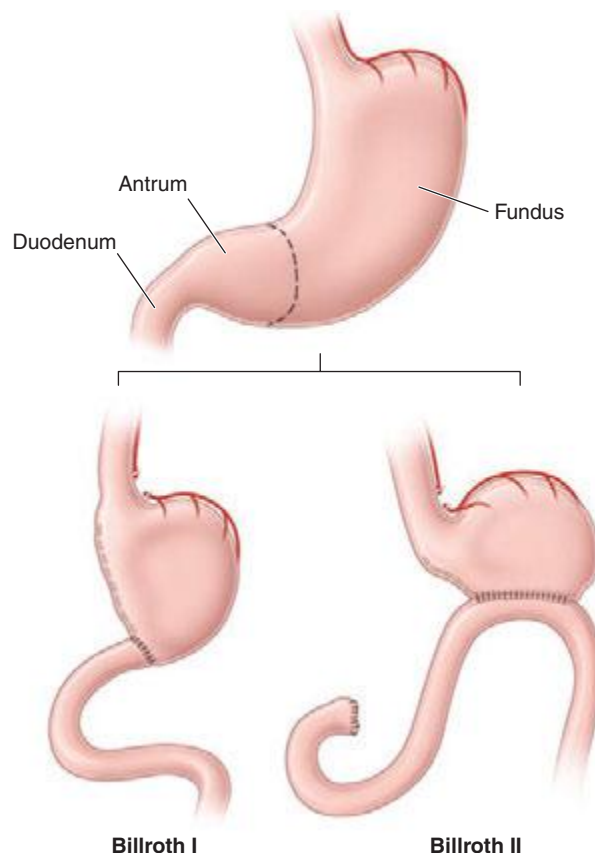


FIGURE 348-13 Schematic representation of Billroth I and II procedures.

anastomosis is the treatment of choice for an antral ulcer. Vagotomy is performed only if a DU is present. Although ulcer excision with vagotomy and drainage procedure has been proposed, the higher incidence of ulcer recurrence makes this a less desirable approach. Ulcers located near the esophagogastric junction may require a more radical approach, a subtotal gastrectomy with a Roux-en-Y esophagogastric anastomosis (Csendes' procedure). A less aggressive approach, including antrectomy, intraoperative ulcer biopsy, and vagotomy (Kelling-Madlener procedure), may be indicated in fragile patients with a high GU. Ulcer recurrence approaches 30% with this procedure.

Surgery-Related Complications Complications seen after surgery for PUD are related primarily to the extent of the anatomic modification performed. Minimal alteration (highly selective vagotomy) is associated with higher rates of ulcer recurrence and less GI disturbance. More aggressive surgical procedures have a lower rate of ulcer recurrence but a greater incidence of GI dysfunction. Overall, morbidity and mortality related to these procedures are quite low. Morbidity associated with vagotomy and antrectomy or pyloroplasty is $\leq 5\%$, with mortality $\sim 1\%$. Highly selective vagotomy has lower morbidity and mortality rates of 1 and 0.3%, respectively.

In addition to the potential early consequences of any intraabdominal procedure (bleeding, infection, thromboembolism), gastroparesis, duodenal stump leak, and afferent loop obstruction can be observed.

RECURRENT ULCERATION The risk of ulcer recurrence is directly related to the procedure performed. Ulcers that recur after partial gastric resection tend to develop at the anastomosis (stomal or marginal ulcer). Epigastric abdominal pain is the most frequent presenting complaint ($>90\%$). Severity and duration of pain tend to be more progressive than observed with DUs before surgery.

Ulcers may recur for several reasons, including incomplete vagotomy, inadequate drainage, retained antrum, and, less likely, persistent or recurrent *H. pylori* infection. ZES should have been excluded preoperatively. Surreptitious use of NSAIDs is an important reason for recurrent ulcers after surgery, especially if the initial procedure was done for an NSAID-induced ulcer. Once *H. pylori* and NSAIDs have been excluded as etiologic factors, the question of incomplete vagotomy or retained gastric antrum should be explored. For the latter, fasting plasma gastrin levels should be determined. If elevated, retained antrum or ZES (see below) should be considered. Incomplete vagotomy can be ruled out by gastric acid analysis coupled with sham feeding. In this test, gastric acid output is measured while the patient sees, smells, and chews a meal (without swallowing). The cephalic phase of gastric secretion, which is mediated by the vagus, is being assessed with this study. An increase in gastric acid output in response to sham feeding is evidence that the vagus nerve is intact. A rise in serum pancreatic polypeptide $>50\%$ within 30 min of sham feeding is also suggestive of an intact vagus nerve.

Medical therapy with H_2 blockers will heal postoperative ulceration in 70–90% of patients. The efficacy of PPIs has not been fully assessed in this group, but one may anticipate greater rates of ulcer healing compared to those obtained with H_2 blockers. Repeat operation (complete vagotomy, partial gastrectomy) may be required in a small subgroup of patients who have not responded to aggressive medical management.

AFFERENT LOOP SYNDROMES Although rarely seen today as a result of the decrease in the performance of Billroth II anastomosis, two types of afferent loop syndrome can occur in patients who have undergone this type of partial gastric resection. The more common of the two is bacterial overgrowth in the afferent limb secondary to stasis. Patients may experience postprandial abdominal pain, bloating, and diarrhea with concomitant malabsorption of fats and vitamin B_{12} . Cases refractory to antibiotics may require surgical revision of the loop. The less common afferent loop syndrome can present with severe abdominal pain and bloating that occur 20–60 min after meals. Pain is often followed by nausea and vomiting of

bile-containing material. The pain and bloating may improve after emesis. The cause of this clinical picture is theorized to be incomplete drainage of bile and pancreatic secretions from an afferent loop that is partially obstructed. Cases refractory to dietary measures may need surgical revision or conversion of the Billroth II anastomosis to a Roux-en-Y gastrojejunostomy.

DUMPING SYNDROME Dumping syndrome consists of a series of vasomotor and GI signs and symptoms and occurs in patients who have undergone vagotomy and drainage (especially Billroth procedures). Two phases of dumping, early and late, can occur. Early dumping takes place 15–30 min after meals and consists of crampy abdominal discomfort, nausea, diarrhea, belching, tachycardia, palpitations, diaphoresis, light-headedness, and, rarely, syncope. These signs and symptoms arise from the rapid emptying of hyperosmolar gastric contents into the small intestine, resulting in a fluid shift into the gut lumen with plasma volume contraction and acute intestinal distention. Release of vasoactive GI hormones (vasoactive intestinal polypeptide, neurotensin, motilin) is also theorized to play a role in early dumping.

The late phase of dumping typically occurs 90 min to 3 h after meals. Vasomotor symptoms (light-headedness, diaphoresis, palpitations, tachycardia, and syncope) predominate during this phase. This component of dumping is thought to be secondary to hypoglycemia from excessive insulin release.

Dumping syndrome is most noticeable after meals rich in simple carbohydrates (especially sucrose) and high osmolality. Ingestion of large amounts of fluids may also contribute. Up to 50% of postvagotomy and drainage patients will experience dumping syndrome to some degree early on. Signs and symptoms often improve with time, but a severe protracted picture can occur in up to 1% of patients.

Dietary modification is the cornerstone of therapy for patients with dumping syndrome. Small, multiple (six) meals devoid of simple carbohydrates coupled with elimination of liquids during meals is important. Antidiarrheals and anticholinergic agents are complementary to diet. Guar and pectin, which increase the viscosity of intraluminal contents, may be beneficial in more symptomatic individuals. Acarbose, an α -glucosidase inhibitor that delays digestion of ingested carbohydrates, has also been shown to be beneficial in the treatment of the late phases of dumping. The somatostatin analogue octreotide has been successful in diet-refractory cases. This drug is administered subcutaneously (50 μ g tid), titrated according to clinical response. A long-acting depot formulation of octreotide can be administered once every 28 days and provides symptom relief comparable to the short-acting agent. In addition, patient weight gain and quality of life appear to be superior with the long-acting form.

POSTVAGOTOMY DIARRHEA Up to 10% of patients may seek medical attention for the treatment of postvagotomy diarrhea. This complication is most commonly observed after truncal vagotomy, which is rarely performed today. Patients may complain of intermittent diarrhea that occurs typically 1–2 h after meals. Occasionally the symptoms may be severe and relentless. This is due to a motility disorder from interruption of the vagal fibers supplying the luminal gut. Other contributing factors may include decreased absorption of nutrients (see below), increased excretion of bile acids, and release of luminal factors that promote secretion. Diphenoxylate or loperamide is often useful in symptom control. The bile salt-binding agent cholestyramine may be helpful in severe cases. Surgical reversal of a 10-cm segment of jejunum may yield a substantial improvement in bowel frequency in a subset of patients.

BILE REFLUX GASTROPATHY A subset of post-partial gastrectomy patients who present with abdominal pain, early satiety, nausea, and vomiting will have mucosal erythema of the gastric remnant as the only finding. Histologic examination of the gastric mucosa reveals minimal inflammation but the presence of epithelial cell injury. This clinical picture is categorized as bile or alkaline reflux gastropathy/gastritis. Although reflux of bile is implicated as the

reason for this disorder, the mechanism is unknown. Prokinetic agents, cholestyramine, and sucralfate have been somewhat effective treatments. Severe refractory symptoms may require using either nuclear scanning with ^{99m}Tc -HIDA to document reflux or an alkaline challenge test, where 0.1 N NaOH is infused into the stomach in an effort to reproduce the patient's symptoms. Surgical diversion of pancreaticobiliary secretions away from the gastric remnant with a Roux-en-Y gastrojejunostomy consisting of a long (50–60 cm) Roux limb has been used in severe cases. Biliary vomiting improves, but early satiety and bloating may persist in up to 50% of patients.

MALDIGESTION AND MALABSORPTION Weight loss can be observed in up to 60% of patients after partial gastric resection. Patients can experience a 10% loss of body weight, which stabilizes 3 months postoperatively. A significant component of this weight reduction is due to decreased oral intake. However, mild steatorrhea can also develop. Reasons for maldigestion/malabsorption include decreased gastric acid production, rapid gastric emptying, decreased food dispersion in the stomach, reduced luminal bile concentration, reduced pancreatic secretory response to feeding, and rapid intestinal transit.

Decreased serum vitamin B₁₂ levels can be observed after partial gastrectomy. This is usually not due to deficiency of IF, since a minimal amount of parietal cells (source of IF) are removed during antrectomy. Reduced vitamin B₁₂ may be due to competition for the vitamin by bacterial overgrowth or inability to split the vitamin from its protein-bound source due to hypochlorhydria.

Iron-deficiency anemia may be a consequence of impaired absorption of dietary iron in patients with a Billroth II gastrojejunostomy. Absorption of iron salts is normal in these individuals; thus, a favorable response to oral iron supplementation can be anticipated. Folate deficiency with concomitant anemia can also develop in these patients. This deficiency may be secondary to decreased absorption or diminished oral intake.

Malabsorption of vitamin D and calcium resulting in osteoporosis and osteomalacia is common after partial gastrectomy and gastrojejunostomy (Billroth II). Osteomalacia can occur as a late complication in up to 25% of post-partial gastrectomy patients. Bone fractures occur twice as commonly in men after gastric surgery as in a control population. It may take years before x-ray findings demonstrate diminished bone density. Elevated alkaline phosphatase, reduced serum calcium, bone pain, and pathologic fractures may be seen in patients with osteomalacia. The high incidence of these abnormalities in this subgroup of patients justifies treating them with vitamin D and calcium supplementation indefinitely. Therapy is especially important in females. Copper deficiency has also been reported in patients undergoing surgeries that bypass the duodenum, where copper is primarily absorbed. Patients may present with a rare syndrome that includes ataxia, myelopathy, and peripheral neuropathy.

GASTRIC ADENOCARCINOMA The incidence of adenocarcinoma in the gastric stump is increased 15 years after resection. Some have reported a four- to fivefold increase in gastric cancer 20–25 years after resection. The pathogenesis is unclear but may involve alkaline reflux, bacterial proliferation, or hypochlorhydria. The role of endoscopic screening is not clear, and most guidelines do not support its use.

ADDITIONAL COMPLICATIONS Reflux esophagitis and a higher incidence of gallstones and cholecystitis have been reported to patients undergoing subtotal gastrectomy. The latter is thought to be due to decreased gallbladder contractility associated with vagotomy and bypass of the duodenum, leading to decreased postprandial release of cholecystokinin.

RELATED CONDITIONS

ZOLLINGER–ELLISON SYNDROME

Severe peptic ulcer diathesis secondary to gastric acid hypersecretion due to unregulated gastrin release from a non- β cell endocrine tumor (gastrinoma) defines the components of ZES. Initially, ZES was

typified by aggressive and refractory ulceration in which total gastrectomy provided the only chance for enhancing survival. Today it can be cured by surgical resection in up to 40% of patients.

Epidemiology The incidence of ZES varies from 0.1–1% of individuals presenting with PUD. Males are more commonly affected than females, and the majority of patients are diagnosed between ages 30 and 50. Gastrinomas are classified into sporadic tumors (more common) and those associated with multiple endocrine neoplasia (MEN) type 1 (see below). The widespread availability and use of PPIs has led to a decreased patient referral for gastrinoma evaluation, delay in diagnosis, and an increase in false-positive diagnoses of ZES. In fact, diagnosis may be delayed for 6 or more years after symptoms consistent with ZES are displayed.

Pathophysiology Hypergastrinemia originating from an autonomous neoplasm is the driving force responsible for the clinical manifestations in ZES. Gastrin stimulates acid secretion through gastrin receptors on parietal cells and by inducing histamine release from ECL cells. Gastrin also has a trophic action on gastric epithelial cells. Long-standing hypergastrinemia leads to markedly increased gastric acid secretion through both parietal cell stimulation and increased parietal cell mass. The increased gastric acid output leads to peptic ulcer diathesis, erosive esophagitis, and diarrhea.

Tumor Distribution Although early studies suggested that the vast majority of gastrinomas occurred within the pancreas, a significant number of these lesions are extrapancreatic. Over 80% of these tumors are found within the hypothetical gastrinoma triangle (confluence of the cystic and common bile ducts superiorly, junction of the second and third portions of the duodenum inferiorly, and junction of the neck and body of the pancreas medially). Duodenal tumors constitute the most common nonpancreatic lesion; between 50 and 75% of gastrinomas are found here. Duodenal tumors are smaller, slower growing, and less likely to metastasize than pancreatic lesions. Less common extrapancreatic sites include stomach, bones, ovaries, heart, liver, and lymph nodes. More than 60% of tumors are considered malignant, with up to 30–50% of patients having multiple lesions or metastatic disease at presentation. Histologically, gastrin-producing cells appear well-differentiated, expressing markers typically found in endocrine neoplasms (chromogranin, neuron-specific enolase).

Clinical Manifestations Gastric acid hypersecretion is responsible for the signs and symptoms observed in patients with ZES. Peptic ulcer is the most common clinical manifestation, occurring in >90% of gastrinoma patients. Initial presentation and ulcer location (duodenal bulb) may be indistinguishable from common PUD. Clinical situations that should create suspicion of gastrinoma are ulcers in unusual locations (second part of the duodenum and beyond), ulcers refractory to standard medical therapy, ulcer recurrence after acid-reducing surgery, ulcers presenting with frank complications (bleeding, obstruction, and perforation), or ulcers in the absence of *H. pylori* or NSAID ingestion. Symptoms of esophageal origin are present in up to two-thirds of patients with ZES, with a spectrum ranging from mild esophagitis to frank ulceration with stricture and Barrett's mucosa.

Diarrhea, the next most common clinical manifestation, is found in up to 50% of patients. Although diarrhea often occurs concomitantly with acid peptic disease, it may also occur independent of an ulcer. Etiology of the diarrhea is multifactorial, resulting from marked volume overload to the small bowel, pancreatic enzyme inactivation by acid, and damage of the intestinal epithelial surface by acid. The epithelial damage can lead to a mild degree of maldigestion and malabsorption of nutrients. The diarrhea may also have a secretory component due to the direct stimulatory effect of gastrin on enterocytes or the co-secretion of additional hormones from the tumor such as vasoactive intestinal peptide.

Gastrinomas can develop in the presence of MEN 1 syndrome (**Chaps. 113 and 408**) in ~25% of patients. This autosomal dominant disorder involves primarily three organ sites: the parathyroid glands (80–90%), pancreas (40–80%), and pituitary gland (30–60%). The

syndrome is caused by inactivating mutations of the *MEN1* tumor suppressor gene found on the long arm of chromosome 11q13. The gene encodes for Menin, which has an important role in DNA replication and transcriptional regulation. A genetic diagnosis is obtained by sequencing of the *MEN1* gene, which can reveal mutations in 70–90% of typical MEN 1 cases. A family may have an unknown mutation, making a genetic diagnosis impossible, and therefore certain individuals will require a clinical diagnosis, which is determined by whether a patient has tumors in two of the three endocrine organs (parathyroid, pancreas/duodenum, or pituitary) or has a family history of MEN 1 and one of the endocrine organ tumors. In view of the stimulatory effect of calcium on gastric secretion, the hyperparathyroidism and hypercalcemia seen in MEN 1 patients may have a direct effect on ulcer disease. Resolution of hypercalcemia by parathyroidectomy reduces gastrin and gastric acid output in gastrinoma patients. An additional distinguishing feature in ZES patients with MEN 1 is the higher incidence of gastric carcinoid tumor development (as compared to patients with sporadic gastrinomas). ZES presents and is diagnosed earlier in MEN 1 patients, and they have a more indolent course as compared to patients with sporadic gastrinoma. Gastrinomas tend to be smaller, multiple, and located in the duodenal wall more often than is seen in patients with sporadic ZES. Establishing the diagnosis of MEN 1 is critical in order to provide genetic counseling to the patient and his or her family and also to determine the recommended surgical approach.

Diagnosis Biochemical measurements of gastrin and acid secretion in patients suspected of ZES play an important role in establishing this rare diagnosis. Often, patients suspected of having ZES will be treated with a PPI in an effort to ameliorate symptoms and decrease the likelihood of possible acid-related complications. The presence of the PPI, which will lower acid secretion and potentially elevate fasting gastrin levels in normal individuals, will make the diagnostic approach in these individuals somewhat difficult. Significant morbidity related to peptic diathesis has been described when stopping PPIs in gastrinoma patients; therefore, a systematic approach in stopping these agents is warranted (see below). The first step in the evaluation of a patient suspected of having ZES is to obtain a fasting gastrin level. A list of clinical scenarios that should arouse suspicion regarding this diagnosis is shown in [Table 348-7](#). Fasting gastrin levels obtained using a dependable assay are usually <150 pg/mL. A normal fasting gastrin, on two separate occasions, especially if the patient is on a PPI, virtually excludes this diagnosis. Virtually all gastrinoma patients will have a gastrin level >150–200 pg/mL. Measurement of fasting gastrin should be repeated to confirm the clinical suspicion. Some of the commercial biochemical assays used for measuring serum gastrin may be inaccurate. Variable specificity of the antibodies used have led to both false-positive and false-negative fasting gastrin levels, placing in jeopardy the ability to make an accurate diagnosis of ZES.

Multiple processes can lead to an elevated fasting gastrin level, the most frequent of which are gastric hypochlorhydria and achlorhydria, with or without pernicious anemia. Gastric acid induces feedback inhibition of gastrin release. A decrease in acid production will

subsequently lead to failure of the feedback inhibitory pathway, resulting in net hypergastrinemia. Gastrin levels will thus be high in patients using antisecretory agents for the treatment of acid peptic disorders and dyspepsia. *H. pylori* infection can also cause hypergastrinemia. Additional causes of elevated gastrin include retained gastric antrum; G cell hyperplasia; gastric outlet obstruction; renal insufficiency; massive small-bowel obstruction; and conditions such as rheumatoid arthritis, vitiligo, diabetes mellitus, and pheochromocytoma. Although a fasting gastrin >10 times normal is highly suggestive of ZES, two-thirds of patients will have fasting gastrin levels that overlap with levels found in the more common disorders outlined above, especially if a PPI is being taken by the patient. The effect of the PPI on gastrin levels and acid secretion will linger several days after stopping the PPI; therefore, it should be stopped for a minimum of 7 days before testing. During this period, the patient should be placed on a histamine H₂ antagonist, such as famotidine, twice to three times per day. Although this type of agent has a short-term effect on gastrin and acid secretion, it needs to be stopped 24 h before repeating fasting gastrin levels or performing some of the tests highlighted below. The patient may take antacids for the final day, stopping them approximately 12 h before testing is performed. Heightened awareness of complications related to gastric acid hypersecretion during the period of PPI cessation is critical.

The next step in establishing a biochemical diagnosis of gastrinoma is to assess acid secretion. Nothing further needs to be done if decreased acid output in the absence of a PPI is observed. A pH can be measured on gastric fluid obtained either during endoscopy or through nasogastric aspiration; a pH <3 is suggestive of a gastrinoma, but a pH >3 is not helpful in excluding the diagnosis. In those situations where the pH is >3, formal gastric acid analysis should be performed if available. Normal BAO in nongastric surgery patients is typically <5 meq/h. A BAO >15 meq/h in the presence of hypergastrinemia is considered pathognomonic of ZES, but up to 12% of patients with common PUD may have elevated BAO to a lesser degree that can overlap with levels seen in ZES patients. In an effort to improve the sensitivity and specificity of gastric secretory studies, a BAO/MAO ratio was established using pentagastrin infusion as a way to maximally stimulate acid production, with a BAO/MAO ratio >0.6 being highly suggestive of ZES. Pentagastrin is no longer available in the United States, making measurement of MAO virtually impossible. An endoscopic method for measuring gastric acid output has been developed but requires further validation.

Gastrin provocative tests have been developed in an effort to differentiate between the causes of hypergastrinemia and are especially helpful in patients with indeterminate acid secretory studies. The tests are the secretin stimulation test and the calcium infusion study. The most sensitive and specific gastrin provocative test for the diagnosis of gastrinoma is the secretin study. An increase in gastrin of ≥120 pg within 15 min of secretin injection has a sensitivity and specificity of >90% for ZES. PPI-induced hypochlorhydria or achlorhydria may lead to a false-positive secretin test; thus, this agent must be stopped for 1 week before testing.

The calcium infusion study is less sensitive and specific than the secretin test, which, coupled with it being a more cumbersome study with greater potential for adverse effects, relegates it to rare utilization in the cases where the patient's clinical characteristics are highly suggestive of ZES but the secretin stimulation is inconclusive.

TABLE 348-7 WHEN TO OBTAIN A FASTING SERUM GASTRIN LEVEL

Multiple ulcers
Ulcers in unusual locations; associated with severe esophagitis; resistant to therapy with frequent recurrences; in the absence of nonsteroidal anti-inflammatory drug ingestion or <i>H. pylori</i> infection
Ulcer patients awaiting surgery
Extensive family history for peptic ulcer disease
Postoperative ulcer recurrence
Basal hyperchlorhydria
Unexplained diarrhea or steatorrhea
Hypercalcemia
Family history of pancreatic islet, pituitary, or parathyroid tumor
Prominent gastric or duodenal folds

Tumor Localization Once the biochemical diagnosis of gastrinoma has been confirmed, the tumor must be located. Multiple imaging studies have been used in an effort to enhance tumor localization ([Table 348-8](#)). The broad range of sensitivity is due to the variable success rates achieved by the different investigative groups. Endoscopic ultrasound (EUS) permits imaging of the pancreas with a high degree of resolution (<5 mm). This modality is particularly helpful in excluding small neoplasms within the pancreas and in assessing the presence of surrounding lymph nodes and vascular involvement, but it is not very sensitive for finding duodenal lesions. Several types of endocrine tumors express cell-surface receptors for somatostatin. This permits the localization of gastrinomas by measuring the uptake of the stable somatostatin analogue¹¹¹ In-pentetreotide (OctreoScan) with sensitivity and specificity rates of >85%.

TABLE 348-8 SENSITIVITY OF IMAGING STUDIES IN ZOLLINGER-ELLISON SYNDROME

Study	Sensitivity, %	
	Primary Gastrinoma	Metastatic Gastrinoma
Ultrasound	21–28	14
CT scan	55–70	>85
Selective angiography	35–68	33–86
Portal venous sampling	70–90	N/A
SASI	55–78	41
MRI	55–70	>85
OctreoScan	67–86	80–100
EUS	80–100	N/A

Abbreviations: CT, computed tomography; EUS, endoscopic ultrasonography; MRI, magnetic resonance imaging; N/A, not applicable; OctreoScan, imaging with ¹¹¹In-pentetreotide; SASI, selective arterial secretin injection.

Up to 50% of patients have metastatic disease at diagnosis. Success in controlling gastric acid hypersecretion has shifted the emphasis of therapy toward providing a surgical cure. Detecting the primary tumor and excluding metastatic disease are critical in view of this paradigm shift. Once a biochemical diagnosis has been confirmed, the patient should first undergo an abdominal computed tomography (CT) scan, magnetic resonance imaging (MRI), or OctreoScan (depending on availability) to exclude metastatic disease. In addition, the positron emitter ⁶⁸Ga has been used to label somatostatin analogues for positron emission tomography (PET) with some success. In addition, hybrid scanners combining CT scan with PET scan are also available in certain specialized centers. Once metastatic disease has been excluded, an experienced endocrine surgeon may opt for exploratory laparotomy with intraoperative ultrasound or transillumination. In other centers, careful examination of the peripancreatic area with EUS, accompanied by endoscopic exploration of the duodenum for primary tumors, will be performed before surgery. Selective arterial secretin injection may be a useful adjuvant for localizing tumors in a subset of patients. The extent of the diagnostic and surgical approach must be carefully balanced with the patient's overall physiologic condition and the natural history of a slow-growing gastrinoma.

TREATMENT ZOLLINGER-ELLISON SYNDROME

Treatment of functional endocrine tumors is directed at ameliorating the signs and symptoms related to hormone overproduction, curative resection of the neoplasm, and attempts to control tumor growth in metastatic disease.

PPIs are the treatment of choice and have decreased the need for total gastrectomy. Initial PPI doses tend to be higher than those used for treatment of GERD or PUD. The initial dose of omeprazole, lansoprazole, rabeprazole, or esomeprazole should be in the range of 60 mg in divided doses in a 24-h period. Dosing can be adjusted to achieve a BAO <10 meq/h (at the drug trough) in surgery-naïve patients and to <5 meq/h in individuals who have previously undergone an acid-reducing operation. Although the somatostatin analogue has inhibitory effects on gastrin release from receptor-bearing tumors and inhibits gastric acid secretion to some extent, PPIs have the advantage of reducing parietal cell activity to a greater degree. Despite this, octreotide may be considered as adjunctive therapy to the PPI in patients with tumors that express somatostatin receptors and have peptic symptoms that are difficult to control with high-dose PPI.

The ultimate goal of surgery would be to provide a definitive cure. Improved understanding of tumor distribution has led to immediate cure rates as high as 60% with 10-year disease-free intervals as high as 34% in sporadic gastrinoma patients undergoing surgery. A positive outcome is highly dependent on the experience of the surgical team treating these rare tumors. Surgical therapy of gastrinoma patients with MEN 1 remains controversial because of the difficulty in rendering these patients disease-free with surgery. In contrast to

the encouraging postoperative results observed in patients with sporadic disease, only 6% of MEN 1 patients are disease free 5 years after an operation. Moreover, in contrast to patients with sporadic ZES, the clinical course of MEN 1 patients is benign and rarely leads to disease-related mortality, recommending that early surgery be deferred. Some groups suggest surgery only if a clearly identifiable, nonmetastatic lesion is documented by structural studies. Others advocate a more aggressive approach, where all patients free of hepatic metastasis are explored and all detected tumors in the duodenum are resected; this is followed by enucleation of lesions in the pancreatic head, with a distal pancreatectomy to follow. The outcome of the two approaches has not been clearly defined. Laparoscopic surgical interventions may provide attractive approaches in the future but currently seem to be of some limited benefit in patients with gastrinoma because a significant percentage of the tumors may be extrapancreatic and difficult to localize with a laparoscopic approach. Finally, patients selected for surgery should be individuals whose health status would lead them to tolerate a more aggressive operation and obtain the long-term benefits from such aggressive surgery, which are often witnessed after 10 years.

Therapy of metastatic endocrine tumors in general remains suboptimal; gastrinomas are no exception. In light of the observation that in many instances tumor growth is indolent and that many individuals with metastatic disease remain relatively stable for significant periods of time, many advocate not instituting systemic tumor-targeted therapy until evidence of tumor progression or refractory symptoms not controlled with PPIs are noted. Medical approaches, including biological therapy (IFN- α , long-acting somatostatin analogues, peptide receptor radionuclides), systemic chemotherapy (streptozotocin, 5-fluorouracil, and doxorubicin), and hepatic artery embolization, may lead to significant toxicity without a substantial improvement in overall survival. ¹¹¹In-pentetreotide has been used in the therapy of metastatic neuroendocrine tumors; further studies are needed. Several novel therapies are being explored, including radiofrequency ablation or cryoablation of liver lesions and use of agents that block the vascular endothelial growth receptor pathway (bevacizumab, sunitinib) or the mammalian target of rapamycin (**Chap. 113**).

Surgical approaches, including debulking surgery and liver transplantation for hepatic metastasis, have also produced limited benefit.

The overall 5- and 10-year survival rates for gastrinoma patients are 62–75% and 47–53%, respectively. Individuals with the entire tumor resected or those with a negative laparotomy have 5- and 10-year survival rates >90%. Patients with incompletely resected tumors have 5- and 10-year survival rates of 43% and 25%, respectively. Patients with hepatic metastasis have <20% survival at 5 years. Favorable prognostic indicators include primary duodenal wall tumors, isolated lymph node tumor, the presence of MEN 1, and undetectable tumor upon surgical exploration. Poor outcome is seen in patients with shorter disease duration; higher gastrin levels (>10,000 pg/mL); large pancreatic primary tumors (>3 cm); metastatic disease to lymph nodes, liver, and bone; and Cushing's syndrome. Rapid growth of hepatic metastases is also predictive of poor outcome.

STRESS-RELATED MUCOSAL INJURY

Patients suffering from shock, sepsis, massive burns, severe trauma, or head injury can develop acute erosive gastric mucosal changes or frank ulceration with bleeding. Classified as stress-induced gastritis or ulcers, injury is most commonly observed in the acid-producing (fundus and body) portions of the stomach. The most common presentation is GI bleeding, which is usually minimal but can occasionally be life threatening. Respiratory failure requiring mechanical ventilation and underlying coagulopathy are risk factors for bleeding, which tends to occur 48–72 h after the acute injury or insult.

Histologically, stress injury does not contain inflammation or *H. pylori*; thus, "gastritis" is a misnomer. Although elevated gastric acid secretion may be noted in patients with stress ulceration after head

1930 trauma (Cushing's ulcer) and severe burns (Curling's ulcer), mucosal ischemia, breakdown of the normal protective barriers of the stomach, systemic release of cytokines, poor GI motility, and oxidative stress also play an important role in the pathogenesis. Acid must contribute to injury in view of the significant drop in bleeding noted when acid inhibitors are used as prophylaxis for stress gastritis.

Improvement in the general management of intensive care unit patients has led to a significant decrease in the incidence of GI bleeding due to stress ulceration. The estimated decrease in bleeding is from 20–30% to <5%. This improvement has led to some debate regarding the need for prophylactic therapy. The high mortality associated with stress-induced clinically important GI bleeding (>40%) and the limited benefit of medical (endoscopic, angiographic) and surgical therapy in a patient with hemodynamically compromising bleeding associated with stress ulcer/gastritis support the use of preventive measures in high-risk patients (mechanically ventilated, coagulopathy, multiorgan failure, or severe burns). Maintenance of gastric pH >3.5 with continuous infusion of H₂ blockers or liquid antacids administered every 2–3 h are viable options. Tolerance to the H₂ blocker is likely to develop; thus, careful monitoring of the gastric pH and dose adjustment are important if H₂ blockers are used. Sucralfate slurry (1 g every 4–6 h) has also been somewhat successful but requires a gastric tube and may lead to constipation and aluminum toxicity. Sucralfate use in endotracheal intubated patients has also been associated with aspiration pneumonia. Meta-analysis comparing H₂ blockers with PPIs for the prevention of stress-associated clinically important and overt GI bleeding demonstrates superiority of the latter without increasing the risk of nosocomial infections, increasing mortality, or prolonging intensive care unit length of stay. Therefore, PPIs are the treatment of choice for stress prophylaxis. Oral PPI is the best option if the patient can tolerate enteral administration. Pantoprazole is available as an intravenous formulation for individuals in whom enteral administration is not possible. If bleeding occurs despite these measures, endoscopy, intra-arterial vasopressin, and embolization are options. If all else fails, then surgery should be considered. Although vagotomy and antrectomy may be used, the better approach would be a total gastrectomy, which has an exceedingly high mortality rate in this setting.

GASTRITIS

The term *gastritis* should be reserved for histologically documented inflammation of the gastric mucosa. Gastritis is not the mucosal erythema seen during endoscopy and is not interchangeable with “dyspepsia.” The etiologic factors leading to gastritis are broad and heterogeneous. Gastritis has been classified based on time course (acute vs chronic), histologic features, and anatomic distribution or proposed pathogenic mechanism (Table 348-9).

The correlation between the histologic findings of gastritis, the clinical picture of abdominal pain or dyspepsia, and endoscopic findings noted on gross inspection of the gastric mucosa is poor. Therefore, there is no typical clinical manifestation of gastritis.

Acute Gastritis The most common causes of acute gastritis are infectious. Acute infection with *H. pylori* induces gastritis. However, *H. pylori* acute gastritis has not been extensively studied. It is reported as presenting with sudden onset of epigastric pain, nausea, and vomiting, and limited mucosal histologic studies demonstrate a marked infiltrate of neutrophils with edema and hyperemia. If not treated, this picture will evolve into one of chronic gastritis. Hypochlorhydria lasting for up to 1 year may follow acute *H. pylori* infection.

Bacterial infection of the stomach or phlegmonous gastritis is a rare, potentially life-threatening disorder characterized by marked and diffuse acute inflammatory infiltrates of the entire gastric wall, at times accompanied by necrosis. Elderly individuals, alcoholics, and AIDS patients may be affected. Potential iatrogenic causes include polypectomy and mucosal injection with India ink. Organisms associated with this entity include streptococci, staphylococci, *Escherichia coli*, *Proteus*, and *Haemophilus* species. Failure of supportive measures and antibiotics may result in gastrectomy.

TABLE 348-9 CLASSIFICATION OF GASTRITIS

- I. Acute gastritis
 - A. Acute *H. pylori* infection
 - B. Other acute Infectious gastritides
 1. Bacterial (other than *H. pylori*)
 2. *H. heilmannii*
 3. Phlegmonous
 4. Mycobacterial
 5. Syphilitic
 6. Viral
 7. Parasitic
 8. Fungal
- II. Chronic atrophic gastritis
 - A. Type A: Autoimmune, body-predominant
 - B. Type B: *H. pylori*-related, antral-predominant
 - C. Indeterminate
- III. Uncommon forms of gastritis
 - A. Lymphocytic
 - B. Eosinophilic
 - C. Crohn's disease
 - D. Sarcoidosis
 - E. Isolated granulomatous gastritis
 - F. Russell body gastritis

Other types of infectious gastritis may occur in immunocompromised individuals such as AIDS patients. Examples include herpetic (herpes simplex) or CMV gastritis. The histologic finding of intranuclear inclusions would be observed in the latter.

Chronic Gastritis Chronic gastritis is identified histologically by an inflammatory cell infiltrate consisting primarily of lymphocytes and plasma cells, with very scant neutrophil involvement. Distribution of the inflammation may be patchy, initially involving superficial and glandular portions of the gastric mucosa. This picture may progress to more severe glandular destruction, with atrophy and metaplasia. Chronic gastritis has been classified according to histologic characteristics. These include superficial atrophic changes and gastric atrophy. The association of atrophic gastritis with the development of gastric cancer has led to the development of endoscopic and serologic markers of severity. Some of these include gross inspection and classification of mucosal abnormalities during standard endoscopy, magnification endoscopy, endoscopy with narrow band imaging and/or autofluorescence imaging, and measurement of several serum biomarkers including pepsinogen I and II levels, gastrin-17, and anti-*H. pylori* serologies. The clinical utility of these tools is currently being explored.

The early phase of chronic gastritis is *superficial gastritis*. The inflammatory changes are limited to the lamina propria of the surface mucosa, with edema and cellular infiltrates separating intact gastric glands. The next stage is *atrophic gastritis*. The inflammatory infiltrate extends deeper into the mucosa, with progressive distortion and destruction of the glands. The final stage of chronic gastritis is *gastric atrophy*. Glandular structures are lost, and there is a paucity of inflammatory infiltrates. Endoscopically, the mucosa may be substantially thin, permitting clear visualization of the underlying blood vessels.

Gastric glands may undergo morphologic transformation in chronic gastritis. Intestinal metaplasia denotes the conversion of gastric glands to a small intestinal phenotype with small-bowel mucosal glands containing goblet cells. The metaplastic changes may vary in distribution from patchy to fairly extensive gastric involvement. Intestinal metaplasia is an important predisposing factor for gastric cancer (Chap. 109).

Chronic gastritis is also classified according to the predominant site of involvement. Type A refers to the body-predominant form (autoimmune), and type B is the antral-predominant form (*H. pylori*-related).

This classification is artificial in view of the difficulty in distinguishing between these two entities. The term *AB gastritis* has been used to refer to a mixed antral/body picture.

TYPE A GASTRITIS The less common of the two forms involves primarily the fundus and body, with antral sparing. Traditionally, this form of gastritis has been associated with pernicious anemia (Chap. 128) in the presence of circulating antibodies against parietal cells and IF; thus, it is also called *autoimmune gastritis*. *H. pylori* infection can lead to a similar distribution of gastritis. The characteristics of an autoimmune picture are not always present.

Antibodies to parietal cells have been detected in >90% of patients with pernicious anemia and in up to 50% of patients with type A gastritis. The parietal cell antibody is directed against H⁺,K⁺-ATPase. T cells are also implicated in the injury pattern of this form of gastritis. A subset of patients infected with *H. pylori* develop antibodies against H⁺,K⁺-ATPase, potentially leading to the atrophic gastritis pattern seen in some patients infected with this organism. The mechanism is thought to involve molecular mimicry between *H. pylori* LPS and H⁺,K⁺-ATPase.

Parietal cell antibodies and atrophic gastritis are observed in family members of patients with pernicious anemia. These antibodies are observed in up to 20% of individuals over age 60 and in ~20% of patients with vitiligo and Addison's disease. About one-half of patients with pernicious anemia have antibodies to thyroid antigens, and about 30% of patients with thyroid disease have circulating antiparietal cell antibodies. Anti-IF antibodies are more specific than parietal cell antibodies for type A gastritis, being present in ~40% of patients with pernicious anemia. Another parameter consistent with this form of gastritis being autoimmune in origin is the higher incidence of specific familial histocompatibility haplotypes such as HLA-B8 and HLA-DR3.

The parietal cell-containing gastric gland is preferentially targeted in this form of gastritis, and achlorhydria results. Parietal cells are the source of IF, the lack of which will lead to vitamin B₁₂ deficiency and its sequelae (megaloblastic anemia, neurologic dysfunction).

Gastric acid plays an important role in feedback inhibition of gastrin release from G cells. Achlorhydria, coupled with relative sparing of the antral mucosa (site of G cells), leads to hypergastrinemia. Gastrin levels can be markedly elevated (>500 pg/mL) in patients with pernicious anemia. ECL cell hyperplasia with frank development of gastric carcinoid tumors may result from gastrin trophic effects. Hypergastrinemia and achlorhydria may also be seen in nonpernicious anemia-associated type A gastritis.

TYPE B GASTRITIS Type B, or antral-predominant, gastritis is the more common form of chronic gastritis. *H. pylori* infection is the cause of this entity. Although described as "antral-predominant," this is likely a misnomer in view of studies documenting the progression of the inflammatory process toward the body and fundus of infected individuals. The conversion to a pangastritis is time-dependent and estimated to require 15–20 years. This form of gastritis increases with age, being present in up to 100% of persons over age 70. Histology improves after *H. pylori* eradication. The number of *H. pylori* organisms decreases dramatically with progression to gastric atrophy, and the degree of inflammation correlates with the level of these organisms. Early on, with antral-predominant findings, the quantity of *H. pylori* is highest and a dense chronic inflammatory infiltrate of the lamina propria is noted, accompanied by epithelial cell infiltration with polymorphonuclear leukocytes (Fig. 348-14).

Multifocal atrophic gastritis, gastric atrophy with subsequent metaplasia, has been observed in chronic *H. pylori*-induced gastritis. This may ultimately lead to development of gastric adenocarcinoma (Fig. 348-8; Chap. 109). *H. pylori* infection is now considered an independent risk factor for gastric cancer. Worldwide epidemiologic studies have documented a higher incidence of *H. pylori* infection in patients with adenocarcinoma of the stomach as compared to control subjects. Seropositivity for *H. pylori* is associated with a three- to sixfold increased risk of gastric cancer. This risk may be as high as ninefold after adjusting for the inaccuracy of serologic testing in the elderly. The mechanism by which *H. pylori* infection leads to cancer

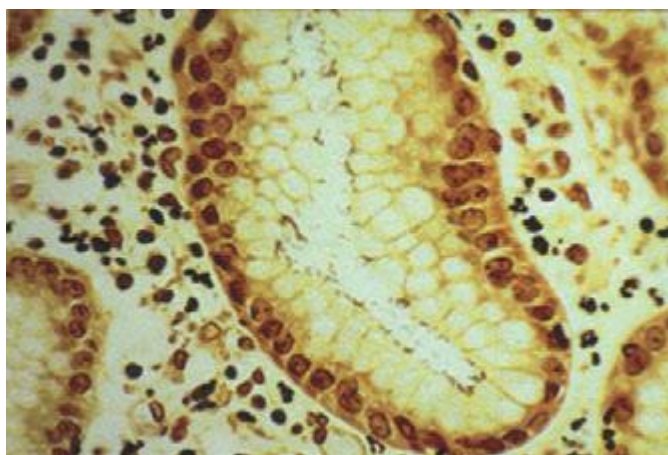


FIGURE 348-14 Chronic gastritis and *H. pylori* organisms. Steiner silver stain of superficial gastric mucosa showing abundant darkly stained microorganisms layered over the apical portion of the surface epithelium. Note that there is no tissue invasion.

is unknown, but it appears to be related to the chronic inflammation induced by the organism. Eradication of *H. pylori* as a general preventative measure for gastric cancer is being evaluated but is not yet recommended.

Infection with *H. pylori* is also associated with development of a low-grade B cell lymphoma, gastric MALT lymphoma (Chap. 134). The chronic T cell stimulation caused by the infection leads to production of cytokines that promote the B cell tumor. The tumor should be initially staged with a CT scan of the abdomen and EUS. Tumor growth remains dependent on the presence of *H. pylori*, and its eradication is often associated with complete regression of the tumor. The tumor may take more than a year to regress after treating the infection. Such patients should be followed by EUS every 2–3 months. If the tumor is stable or decreasing in size, no other therapy is necessary. If the tumor grows, it may have become a high-grade B cell lymphoma. When the tumor becomes a high-grade aggressive lymphoma histologically, it loses responsiveness to *H. pylori* eradication.

TREATMENT CHRONIC GASTRITIS

Treatment in chronic gastritis is aimed at the sequelae and not the underlying inflammation. Patients with pernicious anemia will require parenteral vitamin B₁₂ supplementation on a long-term basis. Eradication of *H. pylori* is often recommended even if PUD or a low-grade MALT lymphoma is not present.

Miscellaneous Forms of Gastritis *Lymphocytic gastritis* is characterized histologically by intense infiltration of the surface epithelium with lymphocytes. The infiltrative process is primarily in the body of the stomach and consists of mature T cells and plasmacytes. The etiology of this form of chronic gastritis is unknown. It has been described in patients with celiac sprue, but whether there is a common factor associating these two entities is unknown. No specific symptoms suggest lymphocytic gastritis. A subgroup of patients have thickened folds noted on endoscopy. These folds are often capped by small nodules that contain a central depression or erosion; this form of the disease is called *varioliform gastritis*. *H. pylori* probably plays no significant role in lymphocytic gastritis. Therapy with glucocorticoids or sodium cromoglycate has obtained unclear results.

Marked eosinophilic infiltration involving any layer of the stomach (mucosa, muscularis propria, and serosa) is characteristic of *eosinophilic gastritis*. Affected individuals will often have circulating eosinophilia with clinical manifestation of systemic allergy. Involvement may

1932 range from isolated gastric disease to diffuse eosinophilic gastroenteritis. Antral involvement predominates, with prominent edematous folds being observed on endoscopy. These prominent antral folds can lead to outlet obstruction. Patients can present with epigastric discomfort, nausea, and vomiting. Treatment with glucocorticoids has been successful.

Several systemic disorders may be associated with *granulomatous gastritis*. Gastric involvement has been observed in Crohn's disease. Involvement may range from granulomatous infiltrates noted only on gastric biopsies to frank ulceration and stricture formation. Gastric Crohn's disease usually occurs in the presence of small-intestinal disease. Several rare infectious processes can lead to granulomatous gastritis, including histoplasmosis, candidiasis, syphilis, and tuberculosis. Other unusual causes of this form of gastritis include sarcoidosis, idiopathic granulomatous gastritis, and eosinophilic granulomas involving the stomach. Establishing the specific etiologic agent in this form of gastritis can be difficult, at times requiring repeat endoscopy with biopsy and cytology. Occasionally, a surgically obtained full-thickness biopsy of the stomach may be required to exclude malignancy.

Russell body gastritis (RBG) is a mucosal lesion of unknown etiology that has a pseudotumoral endoscopic appearance. Histologically, it is defined by the presence of numerous plasma cells containing Russell bodies (RBs) that express kappa and lambda light chains. Only 10 cases have been reported, and 7 of these have been associated with *H. pylori* infection. The lesion can be confused with a neoplastic process, but it is benign in nature, and the natural history of the lesion is not known. There have been cases of resolution of the lesion when *H. pylori* was eradicated.

MÉNÉTRIER'S DISEASE

Ménétrier's disease (MD) is a very rare gastropathy characterized by large, tortuous mucosal folds. MD has an average age of onset of 40–60 years with a male predominance. The differential diagnosis of large gastric folds includes ZES, malignancy (lymphoma, infiltrating carcinoma), infectious etiologies (CMV, histoplasmosis, syphilis, tuberculosis), gastritis polyposa profunda, and infiltrative disorders such as sarcoidosis. MD is most commonly confused with large or multiple gastric polyps (prolonged PPI use) or familial polyposis syndromes. The mucosal folds in MD are often most prominent in the body and fundus, sparing the antrum. Histologically, massive foveolar hyperplasia (hyperplasia of surface and glandular mucous cells) and a marked reduction in oxyntic glands and parietal cells and chief cells are noted. This hyperplasia produces the prominent folds observed. The pits of the gastric glands elongate and may become extremely dilated and tortuous. Although the lamina propria may contain a mild chronic inflammatory infiltrate including eosinophils and plasma cells, MD is not considered a form of gastritis. The etiology of this unusual clinical picture in children is often CMV, but the etiology in adults is unknown. Overexpression of the growth factor TGF- α has been demonstrated in patients with MD. The overexpression of TGF- α in turn results in overstimulation of the epidermal growth factor receptor (EGFR) pathway and increased proliferation of mucus cells, resulting in the observed foveolar hyperplasia.

The clinical presentation in adults is usually insidious and progressive. Epigastric pain, nausea, vomiting, anorexia, peripheral edema, and weight loss are signs and symptoms in patients with MD. Occult GI bleeding may occur, but overt bleeding is unusual and, when present, is due to superficial mucosal erosions. In fact, bleeding is more often seen in one of the common mimics of MD, gastric polyposis. Twenty to 100% of patients (depending on time of presentation) develop a protein-losing gastropathy due to hypersecretion of gastric mucus accompanied by hypoalbuminemia and edema. Gastric acid secretion is usually reduced or absent because of the decreased parietal cells. Large gastric folds are readily detectable by either radiographic (barium meal) or endoscopic methods. Endoscopy with deep mucosal biopsy, preferably full thickness with a snare technique, is required to establish the diagnosis and exclude other entities that may present similarly. A nondiagnostic biopsy may lead to a surgically obtained full-thickness biopsy to exclude malignancy. Although MD is considered premalignant by some, the risk of neoplastic progression is not defined. Complete blood count, serum gastrin, serum

albumin, CMV and *H. pylori* serology, and pH testing of gastric aspirate during endoscopy should be included as part of the initial evaluation of patients with large gastric folds.

TREATMENT MÉNÉTRIER'S DISEASE

Medical therapy with anticholinergic agents, prostaglandins, PPIs, prednisone, somatostatin analogues (octreotide) and H₂ receptor antagonists yields varying results. Ulcers should be treated with a standard approach. The discovery that MD is associated with overstimulation of the EGFR pathway has led to the successful use of the EGF inhibitory antibody, cetuximab, in these patients. Specifically, four of seven patients who completed a 1-month trial with this agent demonstrated near complete histologic remission and improvement in symptoms. Cetuximab is now considered the first-line treatment for MD, leaving total gastrectomy for severe disease with persistent and substantial protein loss despite therapy with this agent.

349 Disorders of Absorption

Henry J. Binder

Disorders of absorption constitute a broad spectrum of conditions with multiple etiologies and varied clinical manifestations. Almost all of these clinical problems are associated with *diminished* intestinal absorption of one or more dietary nutrients and are often referred to as the *malabsorption syndrome*. This term is not ideal as it represents a pathophysiologic state, does *not* provide an etiologic explanation for the underlying problem, and should not be considered an adequate final diagnosis. The only clinical conditions in which absorption is *increased* are hemochromatosis and Wilson's disease, in which absorption of iron and copper, respectively, is elevated.

Most malabsorption syndromes are associated with *steatorrhea*, an increase in stool fat excretion to >6% of dietary fat intake. Some malabsorption disorders are not associated with steatorrhea: primary lactase deficiency, a congenital absence of the small-intestinal brush border disaccharidase enzyme lactase, is associated with lactose "malabsorption," and pernicious anemia is associated with a marked decrease in intestinal absorption of cobalamin (vitamin B₁₂) due to an absence of gastric parietal-cell intrinsic factor, which is required for cobalamin absorption.

Disorders of absorption must be included in the differential diagnosis of diarrhea (**Chap. 55**). First, diarrhea is frequently associated with and/or is a consequence of the diminished absorption of one or more dietary nutrients. The diarrhea may be secondary either to the intestinal process that is responsible for the steatorrhea or to steatorrhea per se. Thus, celiac disease (see below) is associated with both extensive morphologic changes in the small-intestinal mucosa and reduced absorption of several dietary nutrients; in contrast, the diarrhea of steatorrhea is the result of the effect of nonabsorbed dietary fatty acids on intestinal (usually colonic) ion transport. For example, oleic acid and ricinoleic acid (a bacterially hydroxylated fatty acid that is also the active ingredient in castor oil, a widely used laxative) induce active colonic Cl ion secretion, most likely secondary to increasing intracellular Ca. In addition, diarrhea per se may result in mild steatorrhea (<11 g of fat excretion while on a 100-g fat diet). Second, most patients will indicate that they have diarrhea, not that they have fat malabsorption. Third, many intestinal disorders that have diarrhea as a prominent symptom (e.g., ulcerative colitis, traveler's diarrhea secondary to an enterotoxin produced by *Escherichia coli*) do not necessarily have diminished absorption of any dietary nutrient.

Diarrhea as a *symptom* (i.e., when the term is used by patients to describe their bowel movement pattern) may reflect a decrease in stool consistency, an increase in stool volume, an increase in number of bowel movements, or any combination of these three changes. In contrast, diarrhea as a *sign* is a quantitative increase in stool water or weight of >200–225 mL or g per 24 h when a Western-type diet is consumed. Individuals consuming a diet with higher fiber content may normally have a stool weight of up to 400 g/24 h. Thus, the clinician must clarify what an individual patient means by diarrhea. Some 10% of patients referred to gastroenterologists for further evaluation of unexplained diarrhea do not have an increase in stool water when this variable is determined quantitatively. Such patients may have small, frequent, somewhat loose bowel movements with stool urgency that is indicative of proctitis but do not have an increase in stool weight or volume.

It is also critical to establish whether a patient's diarrhea is secondary to diminished absorption of one or more dietary nutrients rather than being due to small- and/or large-intestinal fluid and electrolyte secretion. The former has often been termed *osmotic diarrhea*, while the latter has been referred to as *secretory diarrhea*. Unfortunately, both secretory and osmotic elements can be present simultaneously in the same disorder; thus, this distinction is not always precise. Nonetheless, two studies—determination of stool electrolytes and observation of the effect of a fast on stool output—can help make this distinction.

The demonstration of the effect of prolonged (>24 h) fasting on stool output can suggest that a *dietary nutrient* is responsible for the individual's diarrhea. Secretory diarrhea associated with enterotoxin-induced traveler's diarrhea would not be affected by prolonged fasting, as enterotoxin-induced stimulation of intestinal fluid and electrolyte secretion is not altered by eating. In contrast, diarrhea secondary to lactose malabsorption in primary lactase deficiency would undoubtedly cease during a prolonged fast. Thus, a substantial decrease in stool output by a fasting patient during quantitative stool collection lasting at least 24 h is presumptive evidence that the diarrhea is related to malabsorption of a dietary nutrient. The persistence of stool output during fasting indicates that the diarrhea is likely secretory and that its cause is *not* a dietary nutrient. Either a luminal (e.g., *E. coli* enterotoxin) or a circulating (e.g., vasoactive intestinal peptide) secretagogue could be responsible for unaltered persistence of a patient's diarrhea during a prolonged fast. The observed effects of fasting can be compared and correlated with stool electrolyte and osmolality determinations.

Measurement of stool electrolytes and osmolality requires comparison of Na^+ and K^+ concentrations in liquid stool with the osmolality of the stool in order to determine the presence or absence of a so-called stool osmotic gap. The following formula is used:

$$2 \times (\text{stool } [\text{Na}^+] + \text{stool } [\text{K}^+]) \leq \text{stool osmolality}$$

The cation concentrations are doubled to estimate stool anion concentrations. The presence of a significant osmotic gap suggests the presence in stool water of a substance (or substances) other than Na^+/K^+ anions that is presumably responsible for the patient's diarrhea. Originally, stool osmolality was measured, but it is almost invariably greater than the required 290–300 mosmol/kg H_2O , reflecting bacterial degradation of nonabsorbed carbohydrate either immediately before defecation or in the stool jar while specimen awaits chemical analysis, even when the stool is refrigerated. As a result, the stool osmolality should be assumed to be 300 mosmol/kg H_2O . A low stool osmolality (<290 mosmol/kg H_2O) reflects the addition of either dilute urine or water, indicating either collection of urine and stool together or so-called factitious diarrhea, a form of Münchausen's syndrome. When the calculated difference in the formula above is >50, an osmotic gap exists; its presence suggests that the diarrhea is due to a nonabsorbed dietary nutrient—e.g., a fatty acid and/or a carbohydrate. When this difference is <25, it is presumed that a dietary nutrient is not responsible for the diarrhea. Since elements of both osmotic diarrhea (i.e., due to malabsorption of a dietary nutrient) and secretory diarrhea may be present, this distinction at times is less clear-cut at the bedside than when used as a teaching example. Ideally, the presence of an osmotic gap will be

associated with a marked decrease in stool output during a prolonged fast, while an osmotic gap will likely be absent in an individual whose stool output is not reduced substantially during a period of fasting.

NUTRIENT DIGESTION AND ABSORPTION

The lengths of the small intestine and the colon are ~300 cm and ~80 cm, respectively. However, the effective functional surface area is ~600-fold greater than that of a hollow tube as a result of folds, villi (in the small intestine), and microvilli. The functional surface area of the small intestine is somewhat greater than that of a doubles tennis court. In addition to nutrient digestion and absorption, the intestinal epithelia have several other functions:

1. *Barrier and immune defense.* The intestine is exposed to a large number of potential antigens and enteric and invasive microorganisms, and it is extremely effective at preventing the entry of almost all of these agents. The intestinal mucosa also synthesizes and secretes secretory IgA.
2. *Fluid and electrolyte absorption and secretion.* The intestine absorbs ~7–8 L of fluid daily, a volume comprising dietary fluid intake (1–2 L/d) and salivary, gastric, pancreatic, biliary, and intestinal fluid (6–7 L/d). Several stimuli, especially bacteria and bacterial enterotoxins, induce fluid and electrolyte secretion that may lead to diarrhea ([Chap. 160](#)).
3. *Synthesis and secretion of several proteins.* The intestinal mucosa is a major site for the production of proteins, including apolipoproteins.
4. *Production of several bioactive amines and peptides.* The intestine is one of the largest endocrine organs in the body and produces several amines (e.g., 5-hydroxytryptophan) and peptides that serve as paracrine and hormonal mediators of intestinal function.

The small and large intestines are distinct anatomically (villi are present in the small intestine but are absent in the colon) and functionally (nutrient digestion and absorption take place in the small intestine but not in the colon). No precise anatomic characteristics separate duodenum, jejunum, and ileum, although certain nutrients are absorbed exclusively in specific areas of the small intestine. However, villous cells in the small intestine (surface epithelial cells in the colon) and crypt cells have distinct anatomic and functional characteristics. Intestinal epithelial cells are continuously renewed; new proliferating epithelial cells at the base of the crypt migrate over 48–72 h to the tip of the villus (or surface of the colon), where they exist as well-developed epithelial cells with digestive and absorptive function. This high rate of cell turnover explains the relatively rapid resolution of diarrhea and other digestive-tract side effects during chemotherapy as new cells not exposed to these toxic agents are produced. Equally important is the paradigm of separation of villous/surface cell and crypt cell functions. Digestive hydrolytic enzymes are present primarily in the brush border of villous epithelial cells. Absorptive and secretory functions are also separate: villous/surface cells are primarily, but not exclusively, the site for absorptive function, while secretory function is located in crypts of both the small and large intestines.

Nutrients, minerals, and vitamins are absorbed by one or more active-transport mechanisms. These mechanisms are energy dependent and are mediated by membrane transport proteins. These processes will result in the *net* movement of a substance against or in the absence of an electrochemical concentration gradient. Intestinal absorption of amino acids and monosaccharides (e.g., glucose) is also a specialized form of active transport—*secondary active transport*. The movement of actively transported nutrients against a concentration gradient is Na^+ dependent and is due to a Na^+ gradient across the apical membrane. The Na^+ gradient is maintained by Na^+/K^+ -adenosine triphosphatase (ATPase), the so-called Na^+ pump located on the basolateral membrane, which extrudes Na^+ and maintains low intracellular $[\text{Na}^+]$ as well as the Na^+ gradient across the apical membrane. As a result, active glucose absorption and glucose-stimulated Na^+ absorption require both the apical membrane transport protein SGLT1 and the basolateral Na^+/K^+ -ATPase. In addition to exhibiting Na^+ for its absorption, glucose stimulates Na^+ and fluid absorption; this effect is

1934 the physiologic basis of oral rehydration therapy for the treatment of diarrhea (Chap. 55). The mechanisms of intestinal fluid and electrolyte absorption and secretion are discussed in Chap. 55.

Although the intestinal epithelial cells are crucial mediators of absorption and of ion and water flow, the several cell types in the lamina propria (e.g., mast cells, macrophages, myofibroblasts) and the enteric nervous system interact with the epithelium to regulate mucosal cell function. Intestinal function results from the integrated responses and interactions of intestinal epithelial cells and intestinal muscle.

ENTEROHEPATIC CIRCULATION OF BILE ACIDS

Bile acids are not present in the diet but are synthesized in the liver by a series of enzymatic steps that also represent cholesterol catabolism. Indeed, interruption of the enterohepatic circulation of bile acids can reduce serum cholesterol levels by 10% before a new steady state is established. Bile acids are either primary or secondary. Primary bile acids are synthesized in the liver from cholesterol, and secondary bile acids are synthesized from primary bile acids in the intestine by colonic bacterial enzymes. The two primary bile acids in humans are cholic acid and chenodeoxycholic acid; the two most abundant secondary bile acids are deoxycholic acid and lithocholic acid. The liver synthesizes ~500 mg of bile acids daily; the bile acids are conjugated to either taurine or glycine (to form tauroconjugated and glycoconjugated bile acids, respectively) and are secreted into the duodenum in bile. The primary functions of bile acids are (1) to promote bile flow, (2) to solubilize cholesterol and phospholipid in the gallbladder by mixed micelle formation, and (3) to enhance dietary lipid digestion and absorption by forming mixed micelles in the proximal small intestine.

Bile acids are primarily absorbed by an active, Na⁺-dependent process that takes place exclusively in the ileum; to a lesser extent, they are absorbed by non-carrier-mediated transport processes in the jejunum, ileum, and colon. Conjugated bile acids that enter the colon are deconjugated by colonic bacterial enzymes. The unconjugated bile acids are rapidly absorbed by nonionic diffusion. Colonic bacterial enzymes also dehydroxylate bile acids to secondary bile acids.

Bile acids absorbed from the intestine return to the liver via the portal vein and are then re-secreted (Fig. 349-1). Bile acid synthesis is largely autoregulated by 7 α -hydroxylase, the initial enzyme in cholesterol degradation. A decrease in the volume of bile acids returning to the liver from the intestine is associated with an increase in bile acid synthesis/cholesterol catabolism, which helps keep the bile-acid pool size relatively constant. However, the capacity to increase bile acid synthesis is limited to ~2- to 2.5-fold (see below). The bile-acid pool size is ~4 g. The pool is circulated via the enterohepatic circulation about twice during each meal, or six to eight times during a 24-h period. A relatively small quantity of bile acids is not absorbed and is excreted in stool daily; this fecal loss is matched by hepatic bile-acid synthesis.

Defects in any of the steps in enterohepatic circulation of bile acids can result in a decrease in the duodenal concentration of conjugated bile acids and consequently in the development of steatorrhea. Thus, steatorrhea can be caused by abnormalities in bile acid synthesis and excretion, their physical state in the intestinal lumen, and reabsorption (Table 349-1).

Synthesis Decreased bile acid synthesis and steatorrhea have been demonstrated in chronic liver disease, but steatorrhea often is not a major component of illness in these patients.

Secretion Although bile acid secretion may be reduced or absent in biliary obstruction, steatorrhea is rarely a significant medical problem in these patients. In contrast, primary biliary cirrhosis represents a defect in canalicular excretion of organic anions, including bile acids, and not infrequently is associated with steatorrhea and its consequences (e.g., chronic bone disease). Thus, the osteopenia/osteomalacia and other chronic bone abnormalities often present in patients with primary biliary cirrhosis and other cholestatic syndromes are secondary to steatorrhea that then leads to calcium and vitamin D malabsorption as well as to the effects of cholestasis (e.g., bile acids and inflammatory cytokines).

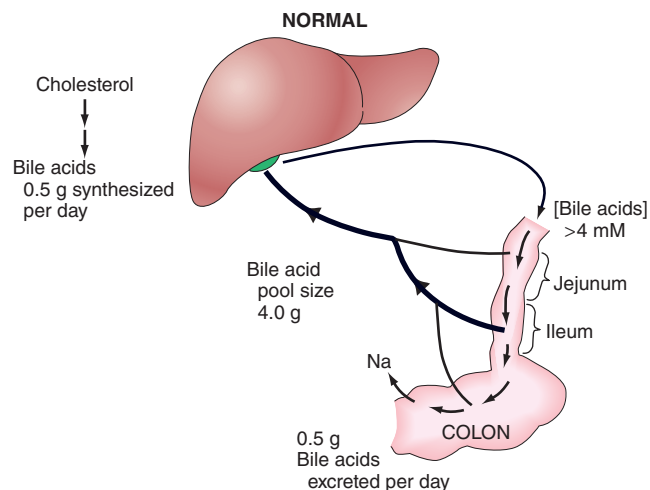


FIGURE 349-1 Schematic representation of the enterohepatic circulation of bile acids. Bile acid synthesis is cholesterol catabolism and occurs in the liver. Bile acids are secreted in bile and are stored in the gallbladder between meals and at night. Food in the duodenum induces the release of cholecystokinin, a potent stimulus for gallbladder contraction resulting in bile acid entry into the duodenum. Bile acids are primarily absorbed via an Na-dependent transport process that is located only in the ileum. A relatively small quantity of bile acids (~500 mg) is not absorbed in a 24-h period and is lost in stool. Fecal bile acid losses are matched by bile acid synthesis. The bile acid pool (the total amount of bile acids in the body) is ~4 g and is circulated twice during each meal or six to eight times in a 24-h period.

Maintenance of Conjugated Bile Acids In bacterial overgrowth syndromes associated with diarrhea, steatorrhea, and macrocytic anemia, a colonic type of bacterial flora is increased in the small intestine. Steatorrhea is primarily a result of the decrease in conjugated bile acids secondary to their deconjugation by colonic-type bacteria. Two complementary explanations account for the resulting impairment of micelle formation: (1) Unconjugated bile acids are rapidly absorbed in the jejunum by nonionic diffusion, and the result is a reduced concentration of duodenal bile acids. (2) The critical micellar concentration (CMC) of unconjugated bile acids is higher than that of conjugated bile acids; therefore, unconjugated bile acids are less effective than conjugated bile acids in micelle formation.

Reabsorption Ileal dysfunction caused by either Crohn's disease or surgical resection results in a decrease in bile acid reabsorption in the ileum and an increase in the delivery of bile acids to the large intestine. The resulting clinical consequences—diarrhea with or without steatorrhea—are determined by the degree of ileal dysfunction and the response of the enterohepatic circulation to bile acid losses (Table 349-2). Patients with limited ileal disease or resection often have diarrhea but not steatorrhea. The diarrhea, a result of stimulation of active Cl secretion by bile acids in the colon, has been called *bile acid diarrhea* or *cholenteric enteropathy* and responds promptly to cholestyramine, an anion-binding resin. Steatorrhea does not develop because hepatic synthesis of bile acids increases to compensate for

TABLE 349-1 DEFECTS IN ENTEROHEPATIC CIRCULATION OF BILE ACIDS

Process	Pathophysiologic Defect	Disease Example
Synthesis	Decreased hepatic function	Cirrhosis
Biliary secretion	Altered canalicular function	Primary biliary cirrhosis
Maintenance of conjugated bile acids	Bacterial overgrowth	Jejunal diverticulosis
Reabsorption	Abnormal ileal function	Crohn's disease

TABLE 349-2 COMPARISON OF BILE ACID AND FATTY ACID DIARRHEA

	Bile Acid Diarrhea	Fatty Acid Diarrhea
Extent of ileal disease	Limited	Extensive
Ileal bile-acid absorption	Reduced	Reduced
Fecal bile-acid excretion	Increased	Increased
Fecal bile-acid loss compensated by hepatic synthesis	Yes	No
Bile-acid pool size	Normal	Reduced
Intraduodenal (bile acid)	Normal	Reduced
Steatorrhea	None or mild	>20 g
Response to cholestyramine	Yes	No
Response to low-fat diet	No	Yes

the rate of fecal bile-acid losses, resulting in maintenance of both the bile-acid pool size and the intraduodenal concentrations of bile acids. In contrast, patients with greater degrees of ileal disease and/or resection often have diarrhea and steatorrhea that do not respond to cholestyramine. In this situation, ileal disease is also associated with increased volumes of bile acids entering the colon; however, hepatic synthesis can no longer increase sufficiently to maintain the bile-acid pool size. As a consequence, the intraduodenal concentration of bile acids is reduced to less than the CMC, and the result is impaired micelle formation and steatorrhea. This second situation is often called *fatty acid diarrhea*. Cholestyramine may not be effective (and may even exacerbate the diarrhea by further depleting the intraduodenal bile-acid concentration); however, a low-fat diet to reduce fatty acid entry into the colon can be effective. Two clinical features—the length of the ileal section removed and the degree of steatorrhea—can predict whether an individual patient will respond to cholestyramine. Unfortunately, these predictors are imperfect, and a therapeutic trial of cholestyramine is often necessary to establish whether an individual patient will benefit from cholestyramine. Table 349-2 contrasts the characteristics of bile acid diarrhea (small ileal dysfunction) and fatty acid diarrhea (large ileal dysfunction).

Bile acid diarrhea can also occur in the absence of ileal inflammation and/or resection and is characterized by an abnormal ⁷⁵SeHCAT retention study and reduced ileal release of fibroblast growth factor 19, a negative regulator of bile acid synthesis, with a consequent increase in bile acid synthesis and secretion that exceeds ileal bile-acid absorption. The diarrhea in these patients also responds to cholestyramine.

LIPIDS

Steatorrhea is caused by one or more defects in the digestion and absorption of dietary fat. The average intake of dietary fat in the United States is ~120–150 g/d, and fat absorption is linear to dietary fat intake. The total load of fat presented to the small intestine is considerably greater, as substantial amounts of lipid are secreted in bile each day (see “Enterohepatic Circulation of Bile Acids,” above). Three types of fatty acids compose fats: long-chain fatty acids (LCFAs), medium-chain fatty acids (MCFAs), and short-chain fatty acids (SCFAs) (Table 349-3). Dietary fat is exclusively composed of long-chain triglycerides (LCTs)—i.e., glycerol that is bound via ester linkages to three LCFAs. While the majority of dietary LCFAs have carbon chain lengths of 16 or 18, all fatty acids of carbon chain length >12 are metabolized in the same manner; saturated and unsaturated fatty acids are handled identically.

Assimilation of dietary lipid requires three integrated processes: (1) an intraluminal, or digestive, phase; (2) a mucosal, or absorptive, phase; and (3) a delivery, or postabsorptive, phase. An abnormality at any site involved in these processes can cause steatorrhea (Table 349-4). Therefore, it is essential that any patient with steatorrhea be evaluated to identify the specific physiologic defect in overall lipid digestion/absorption, as therapy will be determined by the specific etiology.

The digestive phase has two components, *lipolysis* and *micelle formation*. Although dietary lipid is in the form of LCTs, the intestinal mucosa does not absorb triglycerides; they must first be hydrolyzed

TABLE 349-3 COMPARISON OF DIFFERENT TYPES OF FATTY ACIDS

	Long-Chain	Medium-Chain	Short-Chain
Carbon chain length	>12	8–12	<8
Present in diet	In large amounts	In small amounts	No
Origin	In diet as triglycerides	Only in small amounts in diet as triglycerides	Bacterial degradation in colon of nonabsorbed carbohydrate to fatty acids
Primary site of absorption	Small intestine	Small intestine	Colon
Requires pancreatic lipolysis	Yes	No	No
Requires micelle formation	Yes	No	No
Present in stool	Minimal	No	Substantial

(Fig. 349-2). The initial step in lipid digestion is the formation of emulsions of finely dispersed lipid, which is accomplished by mastication and gastric contractions. Lipolysis, the hydrolysis of triglycerides to free fatty acids, monoglycerides, and glycerol by lipase, is initiated in the stomach by lingual and gastric lipases that have a pH optimum of 4.5–6.0. About 20–30% of total lipolysis occurs in the stomach. Lipolysis is completed in the duodenum and jejunum by pancreatic lipase, which is inactivated by a pH <7.0. Pancreatic lipolysis is greatly enhanced by the presence of a second pancreatic enzyme, colipase, which facilitates the movement of lipase to the triglyceride.

Impaired lipolysis can lead to steatorrhea and can occur in the presence of pancreatic insufficiency due to chronic pancreatitis in adults or cystic fibrosis in children and adolescents. Normal lipolysis can be maintained by ~5% of maximal pancreatic lipase secretion; thus, steatorrhea is a late manifestation of these disorders. A reduction in intraduodenal pH can also result in altered lipolysis, as pancreatic lipase is inactivated at pH <7. Thus, ~15% of patients who have gastrinoma (Chap. 348), with substantial increases in gastric acid secretion from ectopic production of gastrin (usually from an islet cell adenoma), have diarrhea, and some have steatorrhea believed to be secondary to acid inactivation of pancreatic lipase. Similarly, patients who have chronic pancreatitis (with reduced lipase secretion) often have a decrease in pancreatic bicarbonate secretion, which will also result in a lowering of intraduodenal pH and inactivation of endogenous pancreatic lipase or of therapeutically administered lipase.

Overlying the microvillus membrane of the small intestine is the so-called unstirred water layer, a relatively stagnant aqueous phase that must be traversed by the products of lipolysis that are primarily water insoluble. Water-soluble mixed micelles provide a mechanism

TABLE 349-4 DEFECTS IN LIPID DIGESTION AND ABSORPTION IN STEATORRHEA

Phase, Process	Pathophysiologic Defect	Disease Example
Digestive		
Lipolysis formation	Decreased lipase secretion	Chronic pancreatitis
Micelle formation	Decreased intraduodenal bile acids	See Table 349-1
Absorptive		
Mucosal uptake and re-esterification	Mucosal dysfunction	Celiac disease
Postabsorptive		
Chylomicron formation	Absent betalipoproteins	Abetalipoproteinemia
Delivery from intestine	Abnormal lymphatics	Intestinal lymphangiectasia

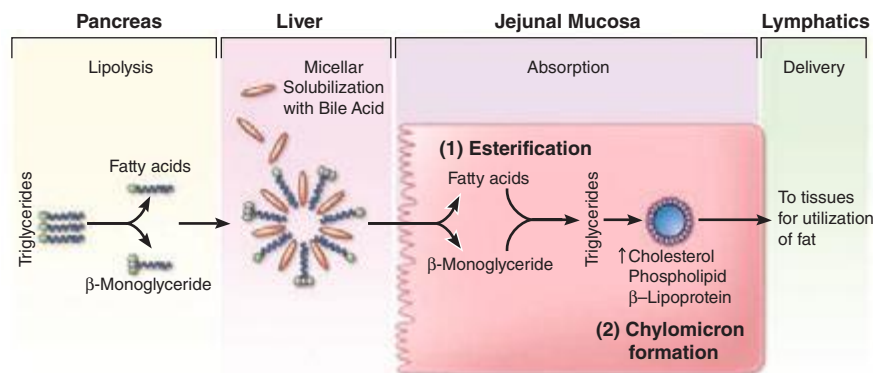


FIGURE 349-2 Schematic representation of lipid digestion and absorption. Dietary lipid is in the form of long-chain triglycerides. The overall process can be divided into (1) a digestive phase that includes both lipolysis and micelle formation requiring pancreatic lipase and conjugated bile acids, respectively, in the duodenum; (2) an absorptive phase for mucosal uptake and re-esterification; and (3) a postabsorptive phase that includes chylomicron formation and exit from the intestinal epithelial cell via lymphatics. (Courtesy of John M. Dietschy, MD; with permission.)

by which the water-insoluble products of lipolysis can reach the luminal plasma membrane of villous epithelial cells—the site for lipid absorption. Mixed micelles are molecular aggregates composed of fatty acids, monoglycerides, phospholipids, cholesterol, and conjugated bile acids. These mixed micelles are formed when the concentration of conjugated bile acids is greater than its CMC, which differs among the several bile acids present in the small-intestinal lumen. Conjugated bile acids, synthesized in the liver and excreted into the duodenum in bile, are regulated by the enterohepatic circulation (see above). Steatorrhea can result from impaired movement of fatty acids across the unstirred aqueous fluid layer in two situations: (1) an increase in the relative thickness of the unstirred water layer that occurs in bacterial overgrowth syndromes (see below) secondary to functional stasis (e.g., scleroderma); and (2) a decrease in the *duodenal* concentration of conjugated bile acids below the CMC, resulting in impaired micelle formation. Thus, steatorrhea can be caused by one or more defects in the enterohepatic circulation of bile acids.

Uptake and re-esterification constitute the *absorptive phase* of lipid digestion/absorption. Although passive diffusion has been thought to be responsible, a carrier-mediated process may mediate fatty acid and monoglyceride uptake. Regardless of the uptake process, fatty acids and monoglycerides are re-esterified by a series of enzymatic steps in the endoplasmic reticulum to form triglycerides, in which lipid exits from the intestinal epithelial cell. Impaired lipid absorption as a result of mucosal inflammation (e.g., celiac disease) and/or intestinal resection can also lead to steatorrhea.

The re-esterified triglycerides require the formation of *chylomicrons* to permit their exit from the small-intestinal epithelial cell and their delivery to the liver via the lymphatics. Chylomicrons are composed of β -lipoprotein and contain triglycerides, cholesterol, cholesterol esters, and phospholipids and enter the lymphatics, not the portal vein. Defects in the *postabsorptive phase* of lipid digestion/absorption can also result in steatorrhea, but these disorders are uncommon. Abetalipoproteinemia, or acanthocytosis, is a rare disorder of impaired synthesis of β -lipoprotein associated with abnormal erythrocytes (acanthocytes), neurologic problems, and steatorrhea (Chap. 421). Lipolysis, micelle formation, and lipid uptake are all normal in patients with abetalipoproteinemia, but the re-esterified triglyceride cannot exit the epithelial cell because of the failure to produce chylomicrons. Small-intestinal biopsy samples obtained from these rare patients in the postprandial state reveal lipid-laden small-intestinal epithelial cells that become perfectly normal in appearance after a 72- to 96-h fast. Similarly, abnormalities of intestinal lymphatics (e.g., intestinal

lymphangiectasia) may also be associated with steatorrhea as well as protein loss (see below). Steatorrhea can result from defects at any of the several steps in lipid digestion/absorption.

The mechanism of lipid digestion/absorption outlined above is limited to *dietary* lipid, which is almost exclusively in the form of LCTs (Table 349-3). Medium-chain triglycerides (MCTs), composed of fatty acids with carbon chain lengths of 8–12, are present in large amounts in coconut oil and are used as a nutritional supplement. MCTs can be digested and absorbed by a pathway different from that involved in LCT digestion and absorption; at one time, MCTs held promise as an important treatment for steatorrhea of almost all etiologies. Unfortunately, they have been less therapeutically effective than expected because, for reasons that are not completely understood, their use often is not associated with an increase in body weight.

In contrast to LCTs, MCTs do not require pancreatic lipolysis as they can be absorbed intact by the intestinal epithelial cell. Further, micelle formation is not necessary for the absorption of MCTs (or MCFAs, if hydrolyzed by pancreatic lipase). MCTs are absorbed more efficiently than LCTs for the following reasons: (1) The rate of absorption is greater for MCTs than for LCFAs. (2) After absorption, MCFAs are not re-esterified. (3) After absorption, MCTs are hydrolyzed to MCFAs. (4) MCTs do not require chylomicron formation to exit intestinal epithelial cells. (5) The route of MCT exit is via the portal vein and not via lymphatics. Thus, the absorption of MCTs is greater than that of LCTs in pancreatic insufficiency, conditions with reduced intraduodenal bile acid concentrations, small-intestinal mucosal disease, abetalipoproteinemia, and intestinal lymphangiectasia.

SCFAs are not dietary lipids but are synthesized by colonic bacterial enzymes from nonabsorbed carbohydrate and are the anions present at the highest concentration in stool (80–130 mM). The SCFAs in stool are primarily acetate, propionate, and butyrate, whose carbon chain lengths are 2, 3, and 4, respectively. Butyrate is the primary nutrient for colonic epithelial cells, and its deficiency can be associated with one or more colitides. SCFAs conserve calories and carbohydrate: carbohydrates that are not completely absorbed in the small intestine will not be absorbed in the large intestine because of the absence of both disaccharidases and SGLT1, the transport protein that mediates monosaccharide absorption. In contrast, SCFAs are rapidly absorbed and stimulate colonic NaCl and fluid absorption. Most antibiotic-associated diarrhea not caused by *Clostridium difficile* is due to antibiotic suppression of the colonic microbiota, with a resulting decrease in SCFA production. As *C. difficile* accounts for only ~15–20% of all antibiotic-associated diarrhea, a relative decrease in colonic production of SCFA is likely the cause of most antibiotic-associated diarrhea.

The clinical manifestations of steatorrhea are a consequence both of the underlying disorder responsible for its development and of steatorrhea per se. Depending on the degree of steatorrhea and the level of dietary intake, significant fat malabsorption may lead to weight loss. Steatorrhea per se can be responsible for diarrhea; if the primary cause of the steatorrhea has not been identified, a low-fat diet can often ameliorate the diarrhea by decreasing fecal fat excretion. Steatorrhea is commonly associated with fat-soluble vitamin deficiency, which requires replacement with water-soluble preparations of these vitamins.

Disorders of absorption may also be associated with malabsorption of other dietary nutrients—most often carbohydrates—with or without a decrease in dietary lipid digestion and absorption. Therefore, knowledge of the mechanisms of digestion and absorption of carbohydrates, proteins, and other minerals and vitamins is useful in the evaluation of patients with altered intestinal nutrient absorption.

TABLE 349-5 PRIMARY LACTASE DEFICIENCY IN ADULT ETHNIC GROUPS

Ethnic Group	Prevalence of Lactase Deficiency, %
Northern European	5–15
Mediterranean	60–85
African black	85–100
American black	45–80
American white	10–25
Native American	50–95
Mexican American	40–75
Asian	90–100

Source: From FJ Simoons: *Am J Dig Dis* 23:963, 1978.

CARBOHYDRATES

Carbohydrates in the diet are present in the form of starch, disaccharides (sucrose and lactose), and glucose. Carbohydrates are absorbed only in the small intestine and only in the form of monosaccharides. Therefore, before their absorption, starch and disaccharides must first be digested by pancreatic amylase and intestinal brush border disaccharidases to monosaccharides. Monosaccharide absorption occurs by a Na-dependent process mediated by the brush border transport protein SGLT1.

Lactose malabsorption is the only clinically important disorder of carbohydrate absorption. Lactose, the disaccharide present in milk, requires digestion by brush border lactase to its two constituent monosaccharides, glucose and galactose. Lactase is present in almost all species in the postnatal period but then disappears throughout the animal kingdom, except in humans. Lactase activity persists in many individuals throughout life. Two different types of lactase deficiency exist—primary and secondary. In *primary lactase deficiency*, a genetically determined decrease or absence of lactase is noted, while all other aspects of both intestinal absorption and brush border enzymes are normal. In a number of nonwhite groups, primary lactase deficiency is common in adulthood. In fact, Northern European and North American whites are the only groups to maintain small-intestinal lactase activity throughout adult life. **Table 349-5** presents the incidence of primary lactase deficiency in several ethnic groups. Lactase persistence in adults is an abnormality due to a defect in the regulation of its maturation. In contrast, *secondary lactase deficiency* occurs in association with small-intestinal mucosal disease, with abnormalities in both structure and function of other brush border enzymes and transport processes. Secondary lactase deficiency is often seen in celiac disease.

As lactose digestion is rate-limiting compared to glucose/galactose absorption, lactase deficiency is associated with significant lactose malabsorption. Some individuals with lactose malabsorption develop symptoms such as diarrhea, abdominal pain, cramps, and/or flatus. Most individuals with primary lactase deficiency do not have symptoms. Since lactose intolerance may be associated with symptoms suggestive of irritable bowel syndrome, persistence of such symptoms in an individual who exhibits lactose intolerance while on a strict lactose-free diet suggests that the person's symptoms were related to irritable bowel syndrome.

The development of symptoms of lactose intolerance is related to several factors:

1. *Amount of lactose in the diet.*
2. *Rate of gastric emptying.* Symptoms are more likely when gastric emptying is rapid than when it is slower. Therefore, skim milk is more likely to be associated with symptoms of lactose intolerance than whole milk, as the rate of gastric emptying after skim milk intake is more rapid. Similarly, diarrhea following subtotal gastrectomy is often a result of lactose intolerance, as gastric emptying is accelerated in patients with a gastrojejunostomy.
3. *Small-intestinal transit time.* Although the small and large intestines both contribute to the development of symptoms, many symptoms of lactase deficiency are related to the interaction of

colonic bacteria and nonabsorbed lactose. More rapid small-intestinal transit makes symptoms more likely.

4. *Colonic compensation by production of SCFAs from nonabsorbed lactose.* Reduced levels of colonic microflora, which can follow antibiotic use, are associated with increased symptoms after lactose ingestion, especially in a lactase-deficient individual.

Glucose-galactose or monosaccharide malabsorption may also be associated with diarrhea and is due to a congenital absence of SGLT1. Diarrhea develops when individuals with this disorder ingest carbohydrates that contain actively transported monosaccharides (e.g., glucose, galactose) but not when they ingest monosaccharides that are not actively transported (e.g., fructose). Fructose is absorbed by the brush border transport protein GLUT 5, a facilitated diffusion process that is not Na-dependent and is distinct from SGLT1. In contrast, some individuals develop diarrhea as a result of the consumption of large quantities of sorbitol, a sugar used in diabetic candy; sorbitol is only minimally absorbed because of the absence of an intestinal absorptive transport mechanism for this sugar.

PROTEINS

Protein is present in food almost exclusively as polypeptides and requires extensive hydrolysis to di- and tripeptides and amino acids before absorption. Proteolysis occurs in both the stomach and the small intestine; it is mediated by pepsin, which is secreted as pepsinogen by gastric chief cells, and by trypsinogen and other peptidases from pancreatic acinar cells. The proenzymes pepsinogen and trypsinogen must be activated to pepsin (by pepsin at a pH <5) and to trypsin (by the intestinal brush border enzyme enterokinase and subsequently by trypsin), respectively. Proteins are absorbed by separate transport systems for di- and tripeptides and for different types of amino acids—e.g., neutral and dibasic. Alterations in either protein or amino acid digestion and absorption are rarely observed clinically, even in the presence of extensive small-intestinal mucosal inflammation. However, three rare genetic disorders involve protein digestion/absorption: (1) *Enterokinase deficiency* is due to an absence of the brush border enzyme that converts the proenzyme trypsinogen to trypsin and is associated with diarrhea, growth retardation, and hypoproteinemia. (2) *Hartnup's syndrome*, a defect in neutral amino acid transport, is characterized by a pellagra-like rash and neuropsychiatric symptoms. (3) *Cystinuria*, a defect in dibasic amino acid transport, is associated with renal calculi and chronic pancreatitis.

APPROACH TO THE PATIENT: Malabsorption

The clues provided by the history, symptoms, and initial preliminary observations will serve to limit extensive, ill-focused, and expensive laboratory and imaging studies. For example, a clinician evaluating a patient who has symptoms suggestive of malabsorption and who has recently undergone extensive small-intestinal resection for mesenteric ischemia should direct the initial assessment almost exclusively to defining whether a short-bowel syndrome might explain the entire clinical picture. Similarly, the development of a pattern of bowel movements suggestive of steatorrhea in a patient with long-standing alcohol abuse and chronic pancreatitis should prompt an assessment of pancreatic exocrine function.

The classic picture of malabsorption is rarely seen today in most parts of the United States. As a consequence, diseases with malabsorption must be suspected in individuals who have less severe symptoms and signs and subtle evidence of the altered absorption of only a *single* nutrient rather than obvious evidence of the malabsorption of multiple nutrients.

Although diarrhea can be caused by changes in fluid and electrolyte movement in either the small or the large intestine, dietary nutrients are absorbed almost exclusively in the small intestine.

Therefore, the demonstration of diminished absorption of a dietary nutrient provides unequivocal evidence for small-intestinal disease, although colonic dysfunction may also be present (e.g., Crohn's disease may involve both the small and large intestines). Dietary nutrient absorption may be segmental or diffuse along the small intestine and is site specific. Thus, for example, calcium, iron, and folic acid are exclusively absorbed by active-transport processes in the proximal small intestine, especially the duodenum; in contrast, the active-transport mechanisms for both cobalamin and bile acids are operative only in the ileum. Therefore, in an individual who years previously has had an intestinal resection, the details of which are not presently available, a presentation with evidence of calcium, folic acid, and/or iron malabsorption but without cobalamin deficiency makes it likely that the duodenum and proximal jejunum, but not the ileum, were resected.

Some nutrients—e.g., glucose, amino acids, and lipids—are absorbed throughout the small intestine, although their rate of absorption is greater in the proximal than in the distal segments. However, after segmental resection of the small intestine, the remaining segments undergo both morphologic and functional “adaptation” to enhance absorption. Such adaptation is secondary to the presence of luminal nutrients and hormonal stimuli and may not be complete in humans for several months after resection. Adaptation is critical for the survival of individuals who have undergone massive resection of the small intestine and/or colon.

Establishing the diagnosis of steatorrhea and identifying its specific cause are often quite difficult. The “gold standard” remains a timed, quantitative stool-fat determination. From a practical standpoint, stool collections are invariably difficult and often incomplete, as nobody wants to handle stool. A qualitative test—Sudan III staining—has long been available to document an increase in stool fat. This test is rapid and inexpensive but, as a qualitative test, does not establish the degree of fat malabsorption and is best used as a preliminary screening study. Many of the blood, breath, and isotopic tests that have been developed (1) do not directly measure fat absorption; (2) exhibit excellent sensitivity when steatorrhea is obvious and severe but poor sensitivity when steatorrhea is mild (e.g., assays for stool chymotrypsin and elastase, which can potentially distinguish pancreatic from nonpancreatic etiologies of steatorrhea); or (3) have not survived the transition from the research laboratory to commercial application.

Nevertheless, routine laboratory studies (i.e., complete blood count, prothrombin time, serum protein determination, alkaline phosphatase) may suggest dietary nutrient depletion, especially deficiencies of iron, folate, cobalamin, and vitamins D and K. Additional studies include measurement of serum carotene, cholesterol, albumin, iron, folate, and cobalamin levels. The serum carotene level can also be reduced if the patient's dietary intake of leafy vegetables is poor.

If steatorrhea and/or altered absorption of other nutrients is suspected, then history, clinical observations, and laboratory testing can help detect deficiency of a nutrient, especially of a fat-soluble vitamin (A, D, E, or K). Thus, evidence of metabolic bone disease with elevated alkaline phosphatase concentrations and/or reduced serum calcium levels suggests vitamin D malabsorption. A deficiency of vitamin K is suggested by an elevated prothrombin time in an individual without liver disease who is not taking anticoagulants. Macrocytic anemia leads to an evaluation for possible cobalamin or folic acid malabsorption. Iron-deficiency anemia in the absence of occult bleeding from the gastrointestinal tract in either a male patient or a nonmenstruating female patient requires an evaluation for iron malabsorption and the exclusion of celiac disease, as iron is absorbed exclusively in the proximal small intestine.

At times, however, a timed (72-h) quantitative stool collection, preferably while the patient is on a defined diet, must be undertaken in order to determine stool fat content and establish the diagnosis of steatorrhea. The presence of steatorrhea then requires further assessment to identify the pathophysiologic process(es) responsible

for the defect in dietary lipid digestion/absorption (Table 349-4). Other studies include the Schilling test (Chap. 350e), the D-xylose test, duodenal mucosal biopsy, small-intestinal radiologic examination, and tests of pancreatic exocrine function.

THE SCHILLING TEST

This test (Chap. 350e) is performed to determine the cause of cobalamin malabsorption. An understanding of the physiology and pathophysiology of cobalamin absorption is very valuable, enhancing comprehension of aspects of gastric, pancreatic, and ileal function. Unfortunately, the Schilling test has not been available commercially in the United States for the past few years.

URINARY D-XYLOSE TEST

The urinary D-xylose test for carbohydrate absorption provides an assessment of proximal small-intestinal mucosal function. D-Xylose, a pentose, is absorbed almost exclusively in the proximal small intestine. The D-xylose test is usually performed by administering 25 g of D-xylose and collecting urine for 5 h. An abnormal test (excretion of <4.5 g) primarily reflects duodenal/jejunal mucosal disease. The D-xylose test can also be abnormal in patients with blind loop syndrome (as a consequence primarily of an abnormal intestinal mucosa) and, as a false-positive study, in patients with large collections of fluid in a third space (i.e., ascites, pleural fluid). The ease of obtaining a mucosal biopsy of the small intestine by endoscopy and the false-negative rate of the D-xylose test have led to its diminished use. When small-intestinal mucosal disease is suspected, a small-intestinal mucosal biopsy should be performed.

RADIOLOGIC EXAMINATION

Radiologic examination of the small intestine using barium contrast (small-bowel series or study) can provide important information in the evaluation of the patient with presumed or suspected malabsorption. This study is most often performed in conjunction with an examination of the esophagus, stomach, and duodenal bulb. Because insufficient barium is given to the patient to permit an adequate examination of the small-intestinal mucosa, especially in the ileum, many gastrointestinal radiologists alter the procedure by performing either a small-bowel series in which a large amount of barium is given by mouth, without concurrent examination of the esophagus and stomach, or an enteroclysis study in which a large amount of barium is introduced into the duodenum via a fluoroscopically placed tube. In addition, many of the diagnostic features initially described by radiologists to denote the presence of small-intestinal disease (e.g., flocculation, segmentation) are rarely seen with current barium suspensions. Nonetheless, in skilled hands, barium contrast examination of the small intestine can yield important information. For example, with extensive mucosal disease, intestinal dilation can be seen as a dilution of barium from increased intestinal fluid secretion (Fig. 349-3). A normal barium contrast study does *not* exclude the possibility of small-intestinal disease. However, a small-bowel series remains useful in the search for anatomic abnormalities, such as strictures and fistulas (as in Crohn's disease) or blind loop syndrome (e.g., multiple jejunal diverticula) and to define the extent of a previous surgical resection. Other imaging studies that assess the integrity of small-intestinal morphology are CT enterography and magnetic resonance enterography. Capsule endoscopy and double-balloon enteroscopy are other useful aids in the diagnostic assessment of small-intestinal pathology.

BIOPSY OF SMALL-INTESTINAL MUCOSA

A small-intestinal mucosal biopsy is essential in the evaluation of a patient with documented steatorrhea or chronic diarrhea (i.e., that lasting >3 weeks) (Chap. 55). The ready availability of endoscopic equipment to examine the stomach and duodenum has led to its almost uniform use as the preferred method of obtaining

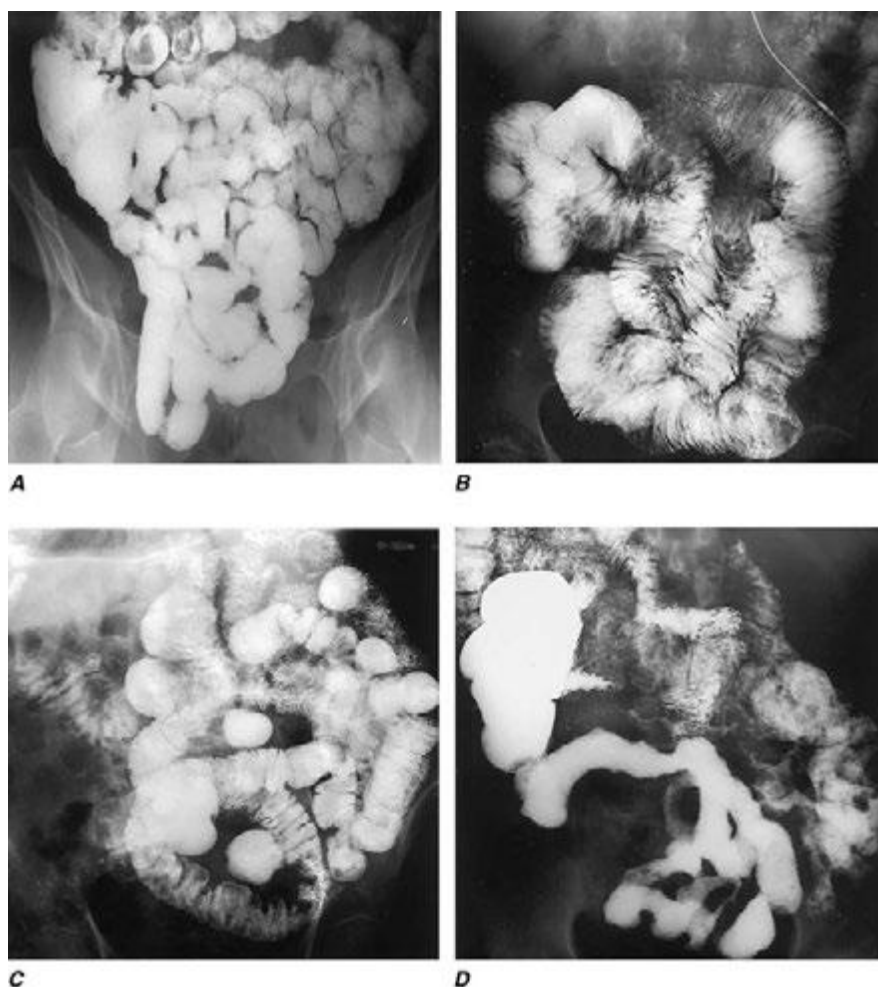


FIGURE 349-3 Barium contrast small-intestinal radiologic examinations. **A.** Normal individual. **B.** Celiac sprue. **C.** Jejunal diverticulosis. **D.** Crohn's disease. (Courtesy of Morton Burrell, MD, Yale University; with permission.)

histologic material from the proximal small-intestinal mucosa. The primary indications for a small-intestinal biopsy are evaluation of a patient (1) either with documented or suspected steatorrhea or with chronic diarrhea, and (2) with diffuse or focal abnormalities of the small intestine defined on a small-intestinal series. Lesions seen on small-bowel biopsy can be classified into three categories (Table 349-6):

1. *Diffuse, specific lesions.* Relatively few diseases associated with altered nutrient absorption have specific histopathologic abnormalities on small-intestinal mucosal biopsy, and these diseases are uncommon. *Whipple's disease* is characterized by the presence of periodic acid-Schiff (PAS)-positive macrophages in the lamina propria; the bacilli that are also present may require electron microscopic examination for identification (Fig. 349-4). *Abetalipoproteinemia* is characterized by a normal mucosal appearance except for the presence of mucosal absorptive cells that contain lipid postprandially and disappear after a prolonged period of either fat-free intake or fasting. *Immune globulin deficiency* is associated with a variety of histopathologic findings on small-intestinal mucosal biopsy. The characteristic feature is the absence of or substantial reduction in the number of plasma cells in the lamina propria; the mucosal architecture may be either perfectly normal or flat (i.e., villous atrophy). As patients with immune globulin deficiency are often infected with *Giardia lamblia*, *Giardia* trophozoites may also be seen in the biopsy.
2. *Patchy, specific lesions.* Several diseases feature an abnormal small-intestinal mucosa with a patchy distribution. As a result,

biopsy samples obtained randomly or in the absence of endoscopically visualized abnormalities may not reveal diagnostic features. Intestinal *lymphoma* can at times be diagnosed on mucosal biopsy by the identification of malignant lymphoma cells in the lamina propria and submucosa (Chap. 134). Dilated lymphatics in the submucosa and sometimes in the lamina propria indicate *lymphangiectasia* associated with hypoproteinemia secondary to protein loss into the intestine. *Eosinophilic gastroenteritis* comprises a heterogeneous group of disorders with a spectrum of presentations and symptoms, with an eosinophilic infiltrate of the lamina propria, and with or without peripheral eosinophilia. The patchy nature of the infiltrate and its presence in the submucosa often lead to an absence of histopathologic findings on mucosal biopsy. As the involvement of the duodenum in *Crohn's disease* is also submucosal and not necessarily continuous, mucosal biopsies are not the most direct approach to the diagnosis of duodenal *Crohn's disease* (Chap. 351). Amyloid deposition can be identified by Congo Red staining in some patients with *amyloidosis* involving the duodenum (Chap. 136).

3. *Diffuse, nonspecific lesions.* *Celiac disease* presents with a characteristic mucosal appearance on duodenal/proximal jejunal mucosal biopsy that is *not* diagnostic of the disease. The diagnosis of celiac disease is established by clinical, histologic, and immunologic responses to a gluten-free diet. *Tropical sprue* (see below) is associated with histologic findings similar to those of celiac disease after a tropical or subtropical exposure but does not respond to gluten restriction; most often symptoms improve with antibiotics and folate administration.

TABLE 349-6 DISEASES THAT CAN BE DIAGNOSED BY SMALL-INTESTINAL MUCOSAL BIOPSIES

Lesions	Pathologic Findings
Diffuse, Specific	
Whipple's disease	Lamina propria includes macrophages containing material positive on periodic acid–Schiff staining
Agammaglobulinemia	No plasma cells; either normal or absent villi ("flat mucosa")
Abetalipoproteinemia	Normal villi; epithelial cells vacuolated with fat postprandially
Patchy, Specific	
Intestinal lymphoma	Malignant cells in lamina propria and submucosa
Intestinal lymphangiectasia	Dilated lymphatics; clubbed villi
Eosinophilic gastroenteritis	Eosinophil infiltration of lamina propria and mucosa
Amyloidosis	Amyloid deposits
Crohn's disease	Noncaseating granulomas
Infection by one or more microorganisms (see text)	Specific organisms
Mastocytosis	Mast cell infiltration of lamina propria
Diffuse, Nonspecific	
Celiac disease	Short or absent villi; mononuclear infiltrate; epithelial cell damage; hypertrophy of crypts
Tropical sprue	Similar to celiac disease
Bacterial overgrowth	Patchy damage to villi; lymphocyte infiltration
Folate deficiency	Short villi; decreased mitosis in crypts; megalocytosis
Vitamin B ₁₂ deficiency	Similar to folate deficiency
Radiation enteritis	Similar to folate deficiency
Zollinger-Ellison syndrome	Mucosal ulceration and erosion from acid
Protein-calorie malnutrition	Villous atrophy; secondary bacterial overgrowth
Drug-induced enteritis	Variable histology


Several microorganisms can be identified in small-intestinal biopsy samples, establishing a correct diagnosis. At times, the biopsy is performed specifically to diagnose infection (e.g., Whipple's disease or giardiasis). In most other instances, the infection is detected incidentally during the workup for diarrhea or other abdominal symptoms. Many of these infections occur in immunocompromised patients with diarrhea; the etiologic agents include *Cryptosporidium*, *Isospora belli*, microsporidia, *Cyclospora*, *Toxoplasma*, cytomegalovirus, adenovirus, *Mycobacterium avium-intracellulare*, and *G. lamblia*. In immunocompromised patients, when *Candida*, *Aspergillus*, *Cryptococcus*, or *Histoplasma* organisms are seen on duodenal biopsy, their presence generally reflects systemic infection. Apart from Whipple's disease and infections in the immunocompromised host, small-bowel biopsy is seldom used as the primary mode of diagnosis of infection. Even giardiasis is more easily diagnosed by stool antigen studies and/or duodenal aspiration than by duodenal biopsy.

Patients with steatorrhea require assessment of *pancreatic exocrine function*, which is often abnormal in chronic pancreatitis. The secretin test that collects pancreatic secretions by duodenal intubation following intravenous administration of secretin is the only test that directly measures pancreatic exocrine function but is available only at a few specialized centers. Endoscopic approaches (endoscopic retrograde cholangiopancreatography, endoscopic ultrasound) provide an excellent assessment of pancreatic duct anatomy but do *not* assess exocrine function (Chap. 370).

Table 349-7 summarizes the results of the D-xylose test, the Schilling test, and small-intestinal mucosal biopsy in patients with steatorrhea of various etiologies.

SPECIFIC DISEASE ENTITIES

CELIAC DISEASE

 *Celiac disease* is a common cause of malabsorption of one or more nutrients. Although celiac disease was originally considered largely a disease of white individuals, especially persons of European descent, recent observations have established that it is a common disease with protean manifestations, a worldwide distribution, and an estimated incidence in the United States that is as high as 1 in 113 people. Its incidence has increased over the past 50 years. Celiac disease has had several other names, including nontropical sprue, celiac sprue, adult celiac disease, and gluten-sensitive enteropathy. The etiology of celiac disease is not known, but environmental, immunologic, and genetic factors are important. Celiac disease is considered an "iceberg" disease. A small number of individuals have classical symptoms and manifestations related to nutrient malabsorption along with a varied natural history; the onset of symptoms can occur at all points from the first year of life through the eighth decade. A much larger number of individuals have "atypical celiac disease", with manifestations that are not obviously related to intestinal malabsorption (e.g., anemia, osteopenia, infertility, and neurologic symptoms). Finally, an even larger number of persons have "silent celiac disease"; they are essentially asymptomatic despite abnormal small-intestinal histopathology and serologies (see below).

The hallmark of celiac disease is an abnormal small-intestinal biopsy (Fig. 349-4) and the response of the condition (including symptoms and histologic changes on small-intestinal biopsy) to the elimination of gluten from the diet. The histologic changes have a proximal-to-distal intestinal distribution of severity, which probably reflects the exposure of the intestinal mucosa to varied amounts of dietary gluten. The symptoms do not necessarily correlate with histologic changes, especially as many newly diagnosed patients with celiac disease may be asymptomatic or only minimally symptomatic (often with no gastrointestinal symptoms).

The symptoms of celiac disease may appear with the introduction of cereals into an infant's diet, although spontaneous remissions often occur during the second decade of life that may be either permanent or followed by the reappearance of symptoms over several years. Alternatively, the symptoms of celiac disease may first become evident at almost any age throughout adulthood. In many patients, frequent spontaneous remissions and exacerbations occur. The symptoms range from significant malabsorption of multiple nutrients, with diarrhea, steatorrhea, weight loss, and the consequences of nutrient depletion (i.e., anemia and metabolic bone disease), to the absence of gastrointestinal symptoms despite evidence of the depletion of a single nutrient (e.g., iron or folate deficiency, osteomalacia, edema from protein loss). Asymptomatic relatives of patients with celiac disease have been identified as having this disease either by small-intestinal biopsy or by serologic studies (e.g., antiendomysial antibodies, tissue transglutaminase [tTG], deamidated gliadin peptide). The availability of these "celiac serologies" has led to a substantial increase in the frequency of diagnosis of celiac disease, and the diagnosis is now being made primarily in patients without "classic" symptoms but with atypical and subclinical presentations.

Etiology The etiology of celiac disease is not known, but environmental, immunologic, and genetic factors all appear to contribute to the disease. One *environmental* factor is the clear association of the disease with gliadin, a component of gluten that is present in wheat, barley, and rye. In addition to the role of gluten restriction in treatment, the instillation of gluten into both the normal-appearing rectum and the distal ileum of patients with celiac disease results in morphologic changes within hours.

An *immunologic* component in the pathogenesis of celiac disease is critical and involves both adaptive and innate immune responses. Serum

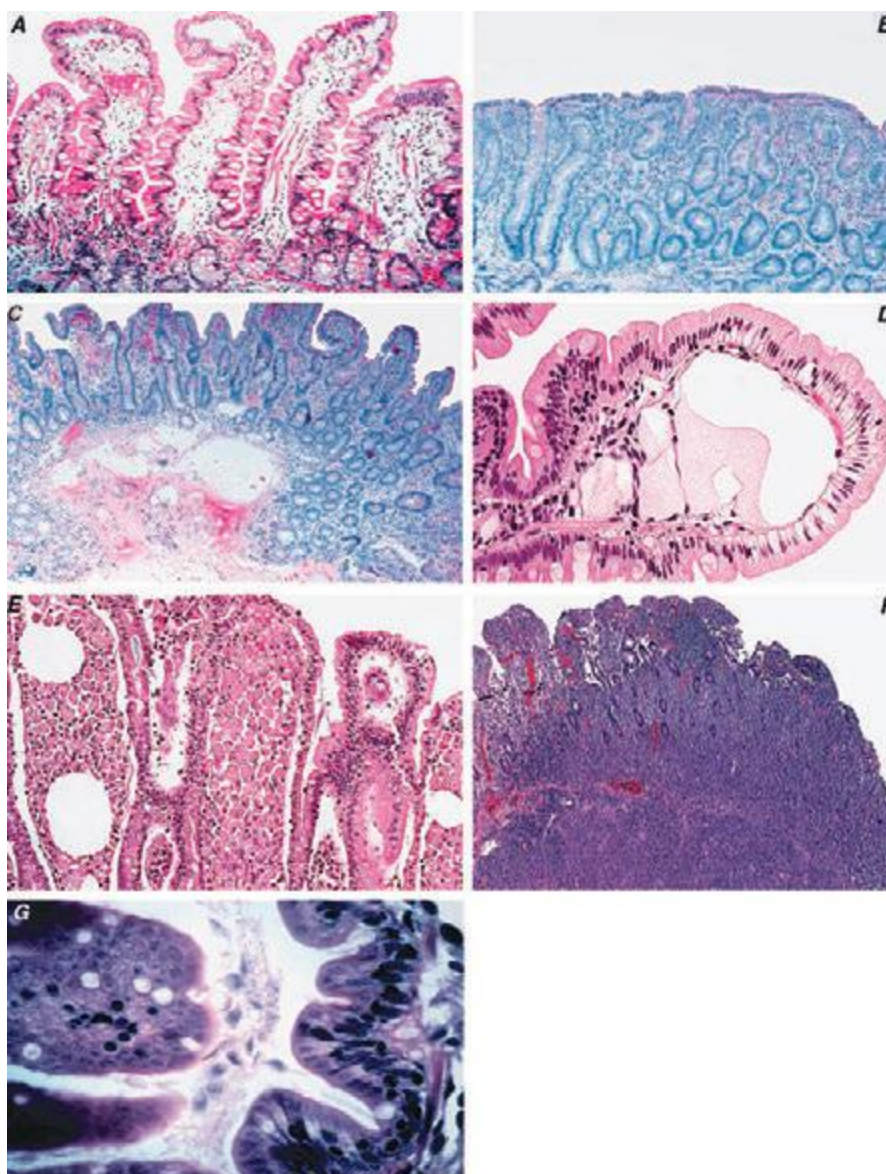


FIGURE 349-4 Small-intestinal mucosal biopsies. **A.** Normal individual. **B.** Untreated celiac sprue. **C.** Treated celiac sprue. **D.** Intestinal lymphangiectasia. **E.** Whipple's disease. **F.** Lymphoma. **G.** Giardiasis. (Courtesy of Marie Robert, MD, Yale University; with permission.)

antibodies—IgA antigliadin, antiendomysial, and anti-tTG antibodies—are present, but it is not known whether such antibodies are primary or secondary to the tissue damage. The presence of antiendomysial antibody is 90–95% sensitive and 90–95% specific; the antigen recognized by antiendomysial antibody is tTG, which deaminates gliadin, which is presented to HLA-DQ2 or HLA-DQ8 (see below). Antibody studies are frequently used to identify patients with celiac disease; patients with these antibodies should undergo duodenal biopsy. This autoantibody has not been linked to a pathogenetic mechanism (or mechanisms) responsible for celiac disease. Nonetheless, this antibody is useful in establishing the

true prevalence of celiac disease in the general population. A 4-week course of treatment with prednisolone induces a remission in a patient with celiac disease who continues to eat gluten and converts the “flat” abnormal duodenal biopsy to a more normal-appearing one. In addition, gliadin peptides interact with gliadin-specific T cells that mediate tissue injury and induce the release of one or more cytokines (e.g., interferon γ) that cause tissue injury.


 Genetic factor(s) are also involved in celiac disease. The incidence of symptomatic celiac disease varies widely in different population groups (high among whites, low among blacks and

TABLE 349-7 RESULTS OF DIAGNOSTIC STUDIES IN STEATORRHEA OF VARIOUS ETIOLOGIES

	D-Xylose Test	Schilling Test	Duodenal Mucosal Biopsy
Chronic pancreatitis	Normal	50% abnormal; if abnormal, normal with pancreatic enzyme treatment	Normal
Bacterial overgrowth syndromes	Normal or only modestly abnormal	Often abnormal; if abnormal, normal after antibiotic treatment	Usually normal
Ileal disease	Normal	Abnormal	Normal
Celiac disease	Decreased	Normal	Abnormal: probably “flat”
Intestinal lymphangiectasia	Normal	Normal	Abnormal: “dilated lymphatics”

1942 Asians) and is 10% among first-degree relatives of celiac disease patients. However, serologic studies provide clear evidence that celiac disease is present worldwide. Furthermore, all patients with celiac disease express the HLA-DQ2 or HLA-DQ8 allele, although only a minority of people expressing DQ2/DQ8 have celiac disease. Absence of DQ2/DQ8 excludes the diagnosis of celiac disease.

Diagnosis A small-intestinal biopsy is required to establish a diagnosis of celiac disease (Fig. 349-4). A biopsy should be performed when patients have symptoms and laboratory findings suggestive of nutrient malabsorption and/or deficiency as well as a positive tTG antibody test. Since the presentation of celiac disease is often subtle, without overt evidence of malabsorption or nutrient deficiency, a relatively low threshold for biopsy performance is important. It is more prudent to perform a biopsy than another test of intestinal absorption that can never completely exclude or establish this diagnosis.

The diagnosis of celiac disease requires the detection of characteristic histologic changes on small-intestinal biopsy together with a prompt clinical and histologic response after the institution of a gluten-free diet. If IgA antiendomysial or tTG antibodies have been detected in serologic studies, they too should disappear after a gluten-free diet is started. With the increase in the number of patients diagnosed with celiac disease (mostly by serologic studies), the spectrum of histologic changes seen on duodenal biopsy has increased and includes findings that are not as severe as the classic changes shown in Fig. 349-4. The classic changes seen on duodenal/jejunal biopsy are restricted to the mucosa and include (1) an increase in the number of intraepithelial lymphocytes; (2) absence or a reduced height of villi, which causes a flat appearance with increased crypt cell proliferation resulting in crypt hyperplasia and loss of villous structure, with consequent villous, but not mucosal, atrophy; (3) a cuboidal appearance and nuclei that are no longer oriented basally in surface epithelial cells; and (4) increased numbers of lymphocytes and plasma cells in the lamina propria (Fig. 349-4B). Although these features are characteristic of celiac disease, they are *not* diagnostic because a similar appearance can develop in tropical sprue, eosinophilic enteritis, and milk-protein intolerance in children and occasionally in lymphoma, bacterial overgrowth, Crohn's disease, and gastrinoma with acid hypersecretion. However, a characteristic histologic appearance that reverts toward normal after the initiation of a gluten-free diet establishes the diagnosis of celiac disease (Fig. 349-4C). Readministration of gluten, with or without an additional small-intestinal biopsy, is not necessary.

A number of patients exhibit *gluten sensitivity*; i.e., they have gastrointestinal symptoms that respond to gluten restriction but do not have celiac disease. The basis for such gluten sensitivity is not known.

Failure to Respond to Gluten Restriction The most common cause of persistent symptoms in a patient who fulfills all the criteria for the diagnosis of celiac disease is *continued intake of gluten*. Gluten is ubiquitous, and a significant effort must be made to exclude all gluten from the diet. Use of rice flour in place of wheat flour is very helpful, and several support groups provide important aid to patients with celiac disease and to their families. More than 90% of patients who have the characteristic findings of celiac disease respond to complete dietary gluten restriction. The remainder constitute a heterogeneous group (whose condition is often called *refractory celiac disease* or *refractory sprue*) that includes some patients who (1) respond to restriction of other dietary protein (e.g., soy); (2) respond to glucocorticoid treatment; (3) are "temporary" (i.e., whose clinical and morphologic findings disappear after several months or years); or (4) fail to respond to all measures and have a fatal outcome, with or without documented complications of celiac disease, such as the development of intestinal T cell lymphoma or autoimmune enteropathy.

Therapeutic approaches that do not include a gluten-free diet are being developed and include the use of peptidases to inactivate toxic gliadin peptides and of small molecules to block toxic peptide uptake across intestinal tight junctions.


Mechanism of Diarrhea The diarrhea in celiac disease has several pathogenetic mechanisms. Diarrhea may be secondary to (1) steatorrhea, which is primarily a result of changes in jejunal mucosal function; (2) secondary lactase deficiency, a consequence of changes in jejunal brush border enzymatic function; (3) bile acid malabsorption resulting in bile acid-induced fluid secretion in the colon (in cases with more extensive disease involving the ileum); and (4) endogenous fluid secretion resulting from crypt hyperplasia. Celiac disease patients with more severe involvement may improve temporarily with *dietary lactose and fat restriction* while awaiting the full effects of total gluten restriction, which constitutes primary therapy.

Associated Diseases Celiac disease is associated with dermatitis herpetiformis (DH), but this association has not been explained. Patients with DH have characteristic papulovesicular lesions that respond to dapsone. Almost all patients with DH have histologic changes in the small intestine consistent with celiac disease, although usually much milder and less diffuse in distribution. Most patients with DH have mild or no gastrointestinal symptoms. In contrast, relatively few patients with celiac disease have DH.

Celiac disease is also associated with diabetes mellitus type 1, IgA deficiency, Down syndrome, and Turner's syndrome. The clinical importance of the association with diabetes is that, although severe watery diarrhea without evidence of malabsorption is most often diagnosed as "diabetic diarrhea" (Chap. 417), assay of antiendomysial antibodies and/or a small-intestinal biopsy must be considered to exclude celiac disease.

Complications The most important complication of celiac disease is the development of cancer. The incidences of both gastrointestinal and nongastrointestinal neoplasms as well as intestinal lymphoma are elevated among patients with celiac disease. For unexplained reasons, the frequency of lymphoma in patients with celiac disease is higher in Ireland and the United Kingdom than in the United States. The possibility of lymphoma must be considered whenever a patient with celiac disease who has previously done well on a gluten-free diet is no longer responsive to gluten restriction or a patient who presents with clinical and histologic features consistent with celiac disease does not respond to a gluten-free diet. Other complications of celiac disease include the development of intestinal ulceration independent of lymphoma and so-called refractory sprue (see above) and collagenous sprue. In *collagenous sprue*, a layer of collagen-like material is present beneath the basement membrane; patients with collagenous sprue generally do not respond to a gluten-free diet and often have a poor prognosis.

TROPICAL SPRUE

 *Tropical sprue* is a poorly understood syndrome that affects both expatriates and natives in certain but not all tropical areas and is manifested by chronic diarrhea, steatorrhea, weight loss, and nutritional deficiencies, including those of both folate and cobalamin. This disease affects 5–10% of the population in some tropical areas.

Chronic diarrhea in a tropical environment is most often caused by infectious agents, including *G. lamblia*, *Yersinia enterocolitica*, *C. difficile*, *Cryptosporidium parvum*, and *Cyclospora cayetanensis*. Tropical sprue should not be entertained as a possible diagnosis until the presence of cysts and trophozoites has been excluded in three stool samples. **Chronic infections of the gastrointestinal tract and diarrhea in patients with or without AIDS are discussed in Chaps. 160, 161, and 226.**

The small-intestinal mucosa of individuals living in tropical areas is not identical to that of individuals who reside in temperate climates. In residents of tropical areas, biopsies reveal a mild alteration of villous architecture with a modest increase in mononuclear cells in the lamina propria, which on occasion can be as severe as that seen in celiac disease. These changes are observed both in native residents and in expatriates living in tropical regions and are usually associated with mild decreases in absorptive function, but they revert to "normal" when an individual moves or returns to a temperate area. Some have suggested that the changes seen in tropical enteropathy and in tropical

sprue represent different ends of the spectrum of a single entity, but convincing evidence to support this concept is lacking.

Etiology Because tropical sprue responds to antibiotics, the consensus is that it may be caused by one or more infectious agents. Nonetheless, the etiology and pathogenesis of tropical sprue are uncertain. First, its occurrence is not evenly distributed in all tropical areas; rather, it is found in specific locations, including southern India, the Philippines, and several Caribbean islands (e.g., Puerto Rico, Haiti), but is rarely observed in Africa, Jamaica, or Southeast Asia. Second, an occasional individual does not develop symptoms of tropical sprue until long after having left an endemic area. For this reason, celiac disease (often referred to as celiac sprue) was originally called *nontropical sprue* to distinguish it from tropical sprue. Third, multiple microorganisms have been identified in jejunal aspirates, with relatively little consistency among studies. *Klebsiella pneumoniae*, *Enterobacter cloacae*, and *E. coli* have been implicated in some studies of tropical sprue, while other studies have favored a role for a toxin produced by one or more of these bacteria. Fourth, the incidence of tropical sprue appears to have decreased substantially during the past two or three decades, perhaps in relation to improved sanitation in many tropical countries during this time. Some have speculated that the reduced occurrence is attributable to the wider use of antibiotics in acute diarrhea, especially in travelers to tropical areas from temperate countries. Fifth, the role of folic acid deficiency in the pathogenesis of tropical sprue requires clarification. Folic acid is absorbed exclusively in the duodenum and proximal jejunum, and most patients with tropical sprue have evidence of folate malabsorption and depletion. Although folate deficiency can cause changes in small-intestinal mucosa that are corrected by folate replacement, several earlier studies reporting that tropical sprue could be cured by folic acid did not provide an explanation for the “insult” that was initially responsible for folate malabsorption.

The clinical pattern of tropical sprue varies in different areas of the world (e.g., India vs. Puerto Rico). Not infrequently, individuals in southern India initially report the occurrence of acute enteritis before the development of steatorrhea and malabsorption. In contrast, in Puerto Rico, a more insidious onset of symptoms and a more dramatic response to antibiotics are seen than in some other locations. Tropical sprue in different areas of the world may not be the same disease, and similar clinical entities may have different etiologies.

Diagnosis The diagnosis of tropical sprue is best based on an abnormal small-intestinal mucosal biopsy in an individual with chronic diarrhea and evidence of malabsorption who is either residing or has recently lived in a tropical country. The small-intestinal biopsy in tropical sprue does not reveal pathognomonic features but resembles, and can often be indistinguishable from, that seen in celiac disease (Fig. 349-4). The biopsy sample in tropical sprue has less villous architectural alteration and more mononuclear cell infiltrate in the lamina propria. In contrast to those of celiac disease, the histologic features of tropical sprue manifest with a similar degree of severity throughout the small intestine, and a gluten-free diet does not result in either clinical or histologic improvement in tropical sprue.

TREATMENT TROPICAL SPRUE

Broad-spectrum antibiotics and folic acid are most often curative, especially if the patient leaves the tropical area and does not return. Tetracycline should be used for up to 6 months and may be associated with improvement within 1–2 weeks. Folic acid alone induces hematologic remission as well as improvement in appetite, weight gain, and some morphologic changes in small-intestinal biopsy. Because of marked folate deficiency, folic acid is most often given together with antibiotics.

SHORT-BOWEL SYNDROME

Short-bowel syndrome is a descriptive term for the myriad clinical problems that follow resection of various lengths of small intestine or,

on rare occasions, are congenital (e.g., microvillous inclusion disease). The factors that determine both the type and degree of symptoms include (1) the specific segment (jejunum vs. ileum) resected, (2) the length of the resected segment, (3) the integrity of the ileocecal valve, (4) whether any large intestine has also been removed, (5) residual disease in the remaining small and/or large intestine (e.g., Crohn’s disease, mesenteric artery disease), and (6) the degree of adaptation in the remaining intestine. Short-bowel syndrome can occur in persons of any age, from neonates to the elderly.

Three different situations in adults mandate intestinal resection: (1) mesenteric vascular disease, including atherosclerosis, thrombotic phenomena, and vasculitides; (2) primary mucosal and submucosal disease (e.g., Crohn’s disease); and (3) operations without preexisting small-intestinal disease (e.g., after trauma).

After resection of the small intestine, the residual intestine undergoes adaptation of both structure and function that may last for up to 6–12 months. Continued intake of dietary nutrients and calories is required to stimulate adaptation via direct contact with the intestinal mucosa, the release of one or more intestinal hormones, and pancreatic and biliary secretions. Thus, enteral nutrition with calorie administration must be maintained, especially in the early postoperative period, even if an extensive intestinal resection requiring parenteral nutrition (PN) has been performed. The subsequent ability of such patients to absorb nutrients will not be known for several months, until adaptation is complete.

Multiple factors besides the absence of intestinal mucosa (required for lipid, fluid, and electrolyte absorption) contribute to diarrhea and steatorrhea in these patients. Removal of the ileum, and especially the ileocecal valve, is often associated with more severe diarrhea than jejunal resection. Without part or all of the ileum, diarrhea can be caused by an increase in bile acids entering the colon; these acids stimulate colonic fluid and electrolyte secretion. Absence of the ileocecal valve is also associated with a decrease in intestinal transit time and bacterial overgrowth from the colon. The presence of the colon (or a major portion) is associated with substantially less diarrhea and a lower likelihood of *intestinal failure* (an inability to maintain nutrition without parenteral support) as a result of fermentation of nonabsorbed carbohydrates to SCFAs. The latter are absorbed in the colon and stimulate Na and water absorption, improving overall fluid balance. Lactose intolerance as a result of the removal of lactase-containing mucosa as well as gastric hypersecretion may also contribute to the diarrhea.

In addition to diarrhea and/or steatorrhea, a range of nonintestinal symptoms is observed in some patients. The frequency of renal calcium oxalate calculi increases significantly in patients with a small-intestinal resection and an intact colon; this greater frequency is due to an increase in oxalate absorption by the large intestine, with subsequent *enteric hyperoxaluria*. Two possible mechanisms for the increase in oxalate absorption in the colon have been suggested: (1) increased bile acids and fatty acids that augment colonic mucosal permeability, resulting in enhanced oxalate absorption; and (2) increased fatty acids that bind calcium, resulting in an enhanced amount of soluble oxalate that is then absorbed. Since oxalate is high in relatively few foods (e.g., spinach, rhubarb, tea), dietary restrictions alone do not constitute adequate treatment. Cholestyramine (an anion-binding resin) and calcium have proved useful in reducing hyperoxaluria. Similarly, an increase in cholesterol gallstones is related to a decrease in the bile-acid pool size, which results in the generation of cholesterol supersaturation in gallbladder bile. Gastric hypersecretion of acid occurs in many patients after large resections of the small intestine. The etiology is unclear but may be related to either reduced hormonal inhibition of acid secretion or increased gastrin levels due to reduced small-intestinal catabolism of circulating gastrin. The resulting gastric acid secretion may be an important factor contributing to diarrhea and steatorrhea. A reduced pH in the duodenum can inactivate pancreatic lipase and/or precipitate duodenal bile acids, thereby increasing steatorrhea, and an increase in gastric secretion can create a volume overload relative to the reduced small-intestinal absorptive capacity. Inhibition of gastric acid secretion with proton pump inhibitors can help reduce diarrhea and steatorrhea, but only for the first 6 months.

Treatment of short-bowel syndrome depends on the severity of symptoms and on whether the individual is able to maintain caloric and electrolyte balance with oral intake alone. Initial treatment includes judicious use of opiates (including codeine) to reduce stool output and to establish an effective diet. If the colon is in situ, the initial diet should be low in fat and high in carbohydrate in order to minimize diarrhea from fatty acid stimulation of colonic fluid secretion. MCTs (see Table 349-3), a low-lactose diet, and various soluble fiber-containing diets should also be tried. In the absence of an ileocecal valve, possible bacterial overgrowth must be considered and treated. If gastric acid hypersecretion is contributing to diarrhea and steatorrhea, a proton pump inhibitor may be helpful. Usually none of these therapeutic approaches provides an instant solution, but each can contribute to the reduction of disabling diarrhea.

The patient's vitamin and mineral status must also be monitored; replacement therapy should be initiated if indicated. Fat-soluble vitamins, folate, cobalamin, calcium, iron, magnesium, and zinc are the most critical factors to monitor on a regular basis. If these approaches are not successful, home PN is an established therapy that can be maintained for many years. Small-intestinal transplantation is becoming established as a possible approach for individuals with extensive intestinal resection who cannot be maintained without PN—i.e., those with intestinal failure. A recombinant analogue of glucagon-like peptide 2 (GLP-2; teduglutide) is approved for use in patients with PN-dependent short-bowel syndrome on the basis of its ability to increase intestinal growth and improve absorption.

BACTERIAL OVERGROWTH SYNDROMES

Bacterial overgrowth syndromes comprise a group of disorders with diarrhea, steatorrhea, and macrocytic anemia whose common feature is the proliferation of colonic-type bacteria within the small intestine. This bacterial proliferation is due to stasis caused by impaired peristalsis (*functional stasis*), changes in intestinal anatomy (*anatomic stasis*), or direct communication between the small and large intestine. These conditions have also been referred to as *stagnant bowel syndrome* or *blind loop syndrome*.

Pathogenesis The manifestations of bacterial overgrowth syndromes are a direct consequence of the presence of increased amounts of a colonic-type bacterial flora, such as *E. coli* or *Bacteroides*, in the small intestine. *Macrocytic anemia* is due to cobalamin—not folate—deficiency. Most bacteria require cobalamin for growth, and increasing concentrations of bacteria use up the relatively small amounts of dietary cobalamin. *Steatorrhea* is due to impaired micelle formation as a consequence of a reduced intraduodenal concentration of conjugated bile acids and the presence of unconjugated bile acids. Certain bacteria, including *Bacteroides*, deconjugate conjugated bile acids to unconjugated bile acids. Unconjugated bile acids are absorbed more rapidly than conjugated bile acids; as a result, the intraduodenal concentration of bile acids is reduced. In addition, the CMC of unconjugated bile acids is higher than that of conjugated bile acids, and the result is a decrease in micelle formation. *Diarrhea* is due, at least in part, to steatorrhea, when it is present. However, some patients manifest diarrhea *without* steatorrhea, and it is assumed that the colonic-type bacteria in these patients are producing one or more bacterial enterotoxins that are responsible for fluid secretion and diarrhea.

Etiology The etiology of these different disorders is bacterial proliferation in the small-intestinal lumen secondary to anatomic or functional stasis or to a communication between the relatively sterile small intestine and the colon, with its high levels of aerobic and anaerobic bacteria. Several examples of *anatomic* stasis have been identified: (1) one or more diverticula (both duodenal and jejunal) (Fig. 294-3C); (2) fistulas and strictures related to Crohn's disease (Fig. 349-3D); (3) a proximal

duodenal afferent loop following subtotal gastrectomy and gastrojejunostomy; (4) a bypass of the intestine (e.g., a jejunoileal bypass for obesity); and (5) dilation at the site of a previous intestinal anastomosis. These anatomic derangements are often associated with the presence of a segment (or segments) of intestine out of continuity of propagated peristalsis, with consequent stasis and bacterial proliferation. Bacterial overgrowth syndromes can also occur in the *absence* of an anatomic blind loop when *functional* stasis is present. Impaired peristalsis and bacterial overgrowth in the absence of a blind loop occur in scleroderma, where motility abnormalities exist in both the esophagus and the small intestine (Chap. 382). Functional stasis and bacterial overgrowth can also develop in association with diabetes mellitus and in the small intestine when a direct connection exists between the small and large intestines, including an ileocolonic resection, or occasionally after an enterocolic anastomosis that permits entry of bacteria into the small intestine as a result of bypassing the ileocecal valve.

Diagnosis The diagnosis may be suspected from the combination of a low serum cobalamin level and an elevated serum folate level, as enteric bacteria frequently produce folate compounds that are absorbed in the duodenum. Ideally, the bacterial overgrowth syndromes are diagnosed by the demonstration of increased levels of aerobic and/or anaerobic colonic-type bacteria in a jejunal aspirate obtained by intubation. However, this specialized test is rarely available. Breath hydrogen testing with administration of lactulose (a nondigestible disaccharide) has also been used to detect bacterial overgrowth. The Schilling test can diagnose bacterial overgrowth (see Chap. 350e) but is not available routinely. Often the diagnosis is suspected clinically and confirmed by the response to treatment.

TREATMENT BACTERIAL OVERGROWTH SYNDROMES

Primary treatment should be directed, if at all possible, to the surgical correction of an anatomic blind loop. In the absence of functional stasis, it is important to define the anatomic relationships responsible for stasis and bacterial overgrowth. For example, bacterial overgrowth secondary to strictures, one or more diverticula, or a proximal afferent loop can potentially be cured by surgical correction of the anatomic state. In contrast, the functional stasis of scleroderma or certain anatomic stasis states (e.g., multiple jejunal diverticula) cannot be corrected surgically, and these conditions should be treated with broad-spectrum antibiotics. Tetracycline used to be the initial drug of choice; because of increasing resistance, however, other antibiotics, such as metronidazole, amoxicillin/clavulanic acid, rifaximin and cephalosporins, have been employed. The antibiotic should be given for ~3 weeks or until symptoms remit. Although the natural history of these conditions is chronic, antibiotics should not be given continuously. Symptoms usually remit within 2–3 weeks of initial antibiotic therapy. Treatment need not be repeated until symptoms recur. For frequent recurrences, several treatment strategies exist, but the use of antibiotics for 1 week per month, whether or not symptoms are present, is often most effective.

Unfortunately, therapy for bacterial overgrowth syndromes is largely empirical, with an absence of clinical trials on which to base rational decisions regarding antibiotic choice, treatment duration, and/or the best approach to therapy for recurrences. Bacterial overgrowth may also occur as a component of another chronic disease, such as Crohn's disease, radiation enteritis, or short-bowel syndrome. Treatment of the bacterial overgrowth in these settings will not cure the underlying problem but may be very important in ameliorating a subset of clinical problems that are related to bacterial overgrowth.

WHIPPLE'S DISEASE

Whipple's disease is a chronic multisystemic disease associated with diarrhea, steatorrhea, weight loss, arthralgia, and central nervous

system (CNS) and cardiac problems; it is caused by the bacterium *Tropheryma whipplei*. Until the identification of *T. whipplei* by polymerase chain reaction, the hallmark of Whipple's disease had been the presence of PAS-positive macrophages in the small intestine (Fig. 349-4E) and other organs with evidence of disease.

Etiology *T. whipplei*, a small (50–500 nm) gram-positive bacillus in the group Actinobacteria, has low virulence but high infectivity. Symptoms of Whipple's disease are relatively minimal compared to the bacterial burden in multiple tissues.

Clinical presentation The onset of Whipple's disease is insidious and is characterized by diarrhea, steatorrhea, abdominal pain, weight loss, migratory large-joint arthropathy, and fever as well as ophthalmologic and CNS symptoms. Dementia is a relatively late symptom and an extremely poor prognostic sign, especially in patients who experience relapse after the induction of a remission with antibiotics. For unexplained reasons, the disease occurs primarily in middle-aged white men. The steatorrhea in these patients is generally believed to be secondary to both small-intestinal mucosal injury and lymphatic obstruction due to the increased number of PAS-positive macrophages in the lamina propria of the small intestine.

Diagnosis The diagnosis of Whipple's disease is suggested by a multisystemic disease in a patient with diarrhea and steatorrhea. Tissue biopsy of the small intestine and/or other organs that may be involved (e.g., liver, lymph nodes, heart, eyes, CNS, or synovial membranes), given the patient's symptoms, is the primary approach. The presence of PAS-positive macrophages containing the characteristic small bacilli is suggestive of this diagnosis. However, *T. whipplei*-containing macrophages can be confused with PAS-positive macrophages containing *M. avium* complex, which may be a cause of diarrhea in AIDS. The presence of the *T. whipplei* bacillus outside of macrophages is a more important indicator of active disease than is their presence within the macrophages. *T. whipplei* has now been successfully grown in culture.

TREATMENT WHIPPLE'S DISEASE

The treatment for Whipple's disease is prolonged use of antibiotics. The current regimen of choice is double-strength trimethoprim-sulfamethoxazole for ~1 year. PAS-positive macrophages can persist after successful treatment, and the presence of bacilli outside of macrophages is indicative of persistent infection or an early sign of recurrence. Recurrence of disease activity, especially with dementia, is an extremely poor prognostic sign and requires an antibiotic that crosses the blood-brain barrier. If trimethoprim-sulfamethoxazole is not tolerated, chloramphenicol is an appropriate second choice.

PROTEIN-LOSING ENTEROPATHY

Protein-losing enteropathy is not a specific disease but rather a group of gastrointestinal and nongastrointestinal disorders with hypoproteinemia and edema in the absence of either proteinuria or defects in protein synthesis (e.g., chronic liver disease). These diseases are characterized by excess protein loss into the gastrointestinal tract. Normally, ~10% of total protein catabolism occurs via the gastrointestinal tract. Evidence of increased protein loss into the gastrointestinal tract is found in more than 65 different diseases, which can be classified into three groups: (1) mucosal ulceration, such that the protein loss primarily represents exudation across damaged mucosa (e.g., ulcerative colitis, gastrointestinal carcinomas, and peptic ulcer); (2) nonulcerated mucosa, but with evidence of mucosal damage so that the protein loss represents loss across epithelia with altered permeability (e.g., celiac disease and Ménétrier's disease in the small intestine and stomach, respectively); and (3) lymphatic dysfunction, representing either primary lymphatic disease or lymphatic disease secondary to partial lymphatic obstruction that may occur as a result of enlarged lymph nodes or cardiac disease.

Diagnosis The diagnosis of protein-losing enteropathy is suggested by peripheral edema and low serum albumin and globulin levels in the absence of renal and hepatic disease. An individual with protein-losing enteropathy only rarely has selective loss of *only* albumin or *only* globulins. Therefore, marked reduction of serum albumin with normal serum globulins should not initiate an evaluation for protein-losing enteropathy but should suggest renal and/or hepatic disease. Likewise, reduced serum globulins with normal serum albumin levels are more likely a result of reduced globulin synthesis rather than enhanced globulin loss into the intestine. An increase in protein loss into the gastrointestinal tract has been documented by the administration of one of several radiolabeled proteins and its quantitation in stool during a 24- or 48-h period. Unfortunately, none of these radiolabeled proteins is available for routine clinical use. α_1 -Antitrypsin, a protein that accounts for ~4% of total serum proteins and is resistant to proteolysis, can be used to detect enhanced rates of serum protein loss into the intestinal tract but cannot be used to assess gastric protein loss because of its degradation in an acid milieu. α_1 -Antitrypsin clearance is measured by determining stool volume as well as both stool and plasma α_1 -antitrypsin concentrations. In addition to the loss of protein via abnormal and distended lymphatics, peripheral lymphocytes may be lost via lymphatics, with consequent relative lymphopenia. Thus, lymphopenia in a patient with hypoproteinemia indicates increased loss of protein into the gastrointestinal tract.

Patients with increased protein loss into the gastrointestinal tract from lymphatic obstruction often have steatorrhea and diarrhea. The steatorrhea is a result of altered lymphatic flow as lipid-containing chylomicrons exit from intestinal epithelial cells via intestinal lymphatics (Table 349-4; Fig. 349-4). In the absence of mechanical or anatomic lymphatic obstruction, intrinsic intestinal lymphatic dysfunction—with or without lymphatic dysfunction in the peripheral extremities—has been designated *intestinal lymphangiectasia*. Similarly, ~50% of individuals with intrinsic peripheral lymphatic disease (Milroy's disease) also have intestinal lymphangiectasia and hypoproteinemia. Other than steatorrhea and enhanced protein loss into the gastrointestinal tract, all other aspects of intestinal absorptive function are normal in intestinal lymphangiectasia.

Other Causes Patients who appear to have idiopathic protein-losing enteropathy without evidence of gastrointestinal disease should be examined for cardiac disease—especially right-sided valvular disease and chronic pericarditis (Chaps. 284 and 288). On occasion, hypoproteinemia can be the only presenting manifestation in these two types of heart disease. Ménétrier's disease (also called *hypertrophic gastropathy*) is an uncommon entity that involves the body and fundus of the stomach and is characterized by large gastric folds, reduced gastric acid secretion, and, at times, enhanced protein loss into the stomach.

TREATMENT PROTEIN-LOSING ENTEROPATHY

As excess protein loss into the gastrointestinal tract is most often secondary to a specific disease, treatment should be directed primarily to the underlying disease process and not to the hypoproteinemia. For example, if significant hypoproteinemia with resulting peripheral edema is secondary to celiac disease or ulcerative colitis, a gluten-free diet and mesalamine, respectively, would be the initial therapy. When enhanced protein loss is secondary to lymphatic obstruction, it is critical to establish the nature of this obstruction. Identification of mesenteric nodes or lymphoma may be possible by imaging studies. Similarly, it is important to exclude cardiac disease as a cause of protein-losing enteropathy, either by echosonography or, on occasion, by a right-heart catheterization.

The increased protein loss that occurs in intestinal lymphangiectasia is a result of distended lymphatics associated with lipid malabsorption. The hypoproteinemia is treated with a low-fat diet and the administration of MCTs (Table 349-3), which do not exit from the intestinal epithelial cells via lymphatics but are delivered to the body via the portal vein.

TABLE 349-8 CLASSIFICATION OF MALABSORPTION SYNDROMES

Inadequate digestion
Postgastrectomy ^a
Deficiency or inactivation of pancreatic lipase
Exocrine pancreatic insufficiency
Chronic pancreatitis
Pancreatic carcinoma
Cystic fibrosis
Pancreatic insufficiency—congenital or acquired
Gastrinoma—acid inactivation of lipase ^a
Drugs—orlistat
Reduced intraduodenal bile acid concentration/impaired micelle formation
Liver disease
Parenchymal liver disease
Cholestatic liver disease
Bacterial overgrowth in small intestine:
Anatomic stasis
Afferent loop
Stasis/blind loop/strictures/fistulae
Functional stasis
Diabetes ^a
Scleroderma ^a
Intestinal pseudo-obstruction
Interrupted enterohepatic circulation of bile salts
Ileal resection
Crohn's disease ^a
Drugs (binding or precipitating bile salts)—neomycin, cholestyramine, calcium carbonate
Impaired mucosal absorption/mucosal loss or defect
Intestinal resection or bypass ^a
Inflammation, infiltration, or infection:
Crohn's disease ^a
Amyloidosis
Scleroderma ^a
Lymphoma ^a
Eosinophilic enteritis
Mastocytosis
Tropical sprue
Celiac disease
Collagenous sprue
Whipple's disease ^a
Radiation enteritis ^a
Folate and vitamin B ₁₂ deficiency
Infections—giardiasis
Graft versus host disease
Genetic disorders
Disaccharidase deficiency
Agammaglobulinemia
Abetalipoproteinemia
Hartnup's disease
Cystinuria
Impaired nutrient delivery to and/or from intestine:
Lymphatic obstruction
Lymphoma ^a
Lymphangiectasia
Circulatory disorders
Congestive heart failure
Constrictive pericarditis
Mesenteric artery atherosclerosis
Vasculitis
Endocrine and metabolic disorders
Diabetes ^a
Hypoparathyroidism
Adrenal insufficiency
Hyperthyroidism
Carcinoid syndrome

^aMalabsorption caused by more than one mechanism.**TABLE 349-9 PATHOPHYSIOLOGY OF CLINICAL MANIFESTATIONS OF MALABSORPTION DISORDERS**

Symptom or Sign	Mechanism
Weight loss/malnutrition	Anorexia, malabsorption of nutrients
Diarrhea	Impaired absorption or secretion of water and electrolytes; colonic fluid secretion secondary to unabsorbed dihydroxy bile acids and fatty acids
Flatus	Bacterial fermentation of unabsorbed carbohydrate
Glossitis, cheilosis, stomatitis	Deficiency of iron, vitamin B ₁₂ , folate, and vitamin A
Abdominal pain	Bowel distention or inflammation, pancreatitis
Bone pain	Calcium, vitamin D malabsorption, protein deficiency, osteoporosis
Tetany, paresthesia	Calcium and magnesium malabsorption
Weakness	Anemia, electrolyte depletion (particularly K ⁺)
Azotemia, hypotension	Fluid and electrolyte depletion
Amenorrhea, decreased libido	Protein depletion, decreased calories, secondary hypopituitarism
Anemia	Impaired absorption of iron, folate, vitamin B ₁₂
Bleeding	Vitamin K malabsorption, hypoprothrombinemia
Night blindness/xerophthalmia	Vitamin A malabsorption
Peripheral neuropathy	Vitamin B ₁₂ and thiamine deficiency
Dermatitis	Deficiency of vitamin A, zinc, and essential fatty acid

SUMMARY

The many conditions that can produce malabsorption are classified by their pathophysiology in [Table 349-8](#). The pathophysiology of the various clinical manifestations of malabsorption is summarized in [Table 349-9](#).

350e The Schilling Test

Henry J. Binder

The Schilling test is performed to determine the cause of cobalamin malabsorption. Unfortunately, this test has not been available commercially in the United States for the last few years. Since an understanding of the physiology and pathophysiology of cobalamin absorption is very valuable in enhancing one's understanding of aspects of gastric, pancreatic, and ileal function, discussion of the Schilling test is provided as supplemental information to **Chap. 349**. Because cobalamin absorption requires multiple steps, including gastric, pancreatic, and ileal processes, the Schilling test also can be used to assess the integrity of the organs involved in those processes (**Chap. 128**).

Cobalamin is present primarily in meat. Except in strict vegans, dietary cobalamin deficiency is exceedingly uncommon. Dietary cobalamin is bound in the stomach to a glycoprotein called *R-binder protein*, which is synthesized in both the stomach and the salivary glands. This cobalamin-R binder complex is formed in the acid milieu of the stomach. Cobalamin absorption has an absolute requirement for *intrinsic factor*, another glycoprotein synthesized and released by gastric parietal cells, to promote its uptake by specific cobalamin receptors on the brush border of ileal enterocytes. Pancreatic protease enzymes split the cobalamin-R binder complex to release cobalamin in the proximal small intestine, where cobalamin then is bound by intrinsic factor.

As a consequence, cobalamin absorption may be abnormal in the following conditions:

1. *Pernicious anemia*. In this disease, immunologically mediated atrophy of gastric parietal cells leads to an absence of both gastric acid and intrinsic factor secretion.
2. *Chronic pancreatitis* can result from a deficiency of pancreatic proteases to split the cobalamin-R binder complex. Although 50% of patients with chronic pancreatitis reportedly have an abnormal Schilling test that is corrected by pancreatic enzyme replacement, cobalamin-responsive macrocytic anemia in chronic pancreatitis is extremely rare. Although this probably reflects a difference in the digestion/absorption of cobalamin in food versus that in a crystalline form, the Schilling test still can be used to assess pancreatic exocrine function.
3. *Achlorhydria* is the absence of hydrochloric acid; intrinsic factor is also secreted with acid which is responsible for splitting cobalamin away from the proteins in food to which it is bound. Up to one-third of individuals >60 years of age have marginal vitamin B₁₂ absorption because of an inability to release cobalamin from

food; these people have no defects in the absorption of crystalline vitamin B₁₂.

4. *Bacterial overgrowth syndromes*, which are most often secondary to stasis in the small intestine, lead to bacterial utilization of cobalamin (often referred to as *stagnant bowel syndrome*; see below).
5. *Ileal dysfunction* (as a result of either inflammation or prior intestinal resection) is due to impaired function of the mechanism of cobalamin-intrinsic factor uptake by ileal intestinal epithelial cells.

In the Schilling test, ⁵⁸Co-labeled cobalamin is administered orally, and urine is collected for 24 h. The test is dependent on normal renal and bladder function. Urinary excretion of cobalamin reflects cobalamin absorption, provided that intrahepatic binding sites for cobalamin are fully occupied. To ensure saturation of these binding sites so that all absorbed radiolabeled cobalamin will be excreted in urine, 1 mg of cobalamin is administered intramuscularly 1 h after ingestion of the radiolabeled cobalamin. The Schilling test may yield an abnormal result (usually defined as <10% excretion in 24 h) in pernicious anemia, chronic pancreatitis, blind loop syndrome, and ileal disease (**Table 350e-1**). Therefore, whenever an abnormal Schilling result is obtained, ⁵⁸Co-labeled cobalamin should be administered on another occasion, this time bound to intrinsic factor, with pancreatic enzymes, or after a 5-day course of antibiotic treatment (often with tetracycline). A variation of the Schilling test can detect failure to split cobalamin from food proteins. The labeled cobalamin is cooked together with a scrambled egg and administered orally. People with achlorhydria excrete <10% of the labeled cobalamin in the urine. In addition to establishing the etiology for cobalamin deficiency, the Schilling test can help delineate the pathologic process responsible for steatorrhea by assessing ileal, pancreatic, and small-intestinal luminal function. Unfortunately, the Schilling test is performed infrequently because of the unavailability of human intrinsic factor.

TABLE 350e-1 DIFFERENTIAL RESULTS OF THE SCHILLING TEST IN SEVERAL DISEASES ASSOCIATED WITH COBALAMIN MALABSORPTION


	⁵⁸ Co-Labeled Cobalamin	With Intrinsic Factor	With Pancreatic Enzymes	After 5 Days of Antibiotics
Pernicious anemia	Reduced	Normal	Reduced	Reduced
Chronic pancreatitis	Reduced	Reduced	Normal	Reduced
Bacterial overgrowth	Reduced	Reduced	Reduced	Normal
Ileal disease	Reduced	Reduced	Reduced	Reduced

351 Inflammatory Bowel Disease

Sonia Friedman, Richard S. Blumberg

Inflammatory bowel disease (IBD) is an immune-mediated chronic intestinal condition. Ulcerative colitis (UC) and Crohn's disease (CD) are the two major types of IBD.

GLOBAL CONSIDERATIONS: EPIDEMIOLOGY

 The incidence and prevalence of IBD are highest in Westernized nations, with UC incidence estimates ranging from 0.6 to 24.3 per 100,000 in Europe, 0 to 19.2 per 100,000 in North America, and 0.1 to 6.3 per 100,000 in the Middle East and Asia and CD estimates ranging from 0.3 to 12.7 per 100,000 in Europe, 0 to 20.2 per 100,000 in North America, and 0.04 to 5.0 per 100,000 in the Middle East and Asia (Table 351-1). For prevalence rates, the UC estimates range from 4.9 to 505 per 100,000 in Europe, 37.5 to 248.6 per 100,000 in North America, and 4.9 to 168.3 per 100,000 in the Middle East and Asia, and the CD estimates range from 0.6 to 322 per 100,000 in Europe, 16.7 to 318.5 per 100,000 in North America, and 0.88 to 67.9 per 100,000 in Asia and the Middle East. The highest reported incidence rates are in Canada (19.2 per 100,000 for UC and 20.2 per 100,000 for CD), with approximately 0.6% of the Canadian population having IBD. Countries in the Pacific, including New Zealand and Australia, which share many possible environmental risk factors and similar genetic background as northwest Europe and North America, have high incidence rates of IBD.

In countries that are becoming more Westernized, including China, South Korea, India, Lebanon, Iran, Thailand, and countries in the French West Indies and North Africa, IBD appears to be emerging, emphasizing the importance of environmental factors in disease pathogenesis. In Japan, the prevalence of CD has risen rapidly from 2.9 cases per 100,000 in 1986 to 13.5 per 100,000 in 1998, whereas in South Korea, the prevalence of UC has quadrupled from 7.6 per 100,000 in 1997 to 30.9 per 100,000 in 2005. In Hong Kong, the prevalence of UC almost tripled from 2.3 in 1997 to 6.3 per 100,000 over a 9-year period. In Singapore, the prevalence of CD increased from 1.3 in 1990 to 7.2 per 100,000 in 2004. In China the number of cases of UC has increased by fourfold between 1981–1990 and 1991–2000.

Increasing immigration to Western societies also has an impact on the incidence and prevalence of IBD. The prevalence of UC among southern Asians who immigrated to the United Kingdom (UK) was

higher in comparison to the European UK population (17 cases per 100,000 persons vs 7 per 100,000). Spanish patients who emigrated within Europe, but not those who immigrated to Latin America, developed IBD more frequently than controls. Individuals who have immigrated to Westernized countries and then returned to their country of birth also continue to demonstrate an increased risk of developing IBD.

Peak incidence of UC and CD is in the second to fourth decades, with 78% of CD studies and 51% of UC studies reporting the highest incidence among those age 20–29 years old. A second modest rise in incidence occurs between the seventh and ninth decades of life. The female-to-male ratio ranges from 0.51 to 1.58 for UC studies and 0.34 to 1.65 for CD studies, suggesting that the diagnosis of IBD is not gender specific. The greatest incidence of IBD is among white and Jewish people, but the incidence of IBD in Hispanic and Asian people is increasing, as noted above. Urban areas have a higher prevalence of IBD than rural areas, and high socioeconomic classes have a higher prevalence than lower socioeconomic classes.

Epidemiologic studies have identified a number of potential environmental factors that are associated with disease risk (Fig. 351-1). In Caucasian populations, smoking is an important risk factor in IBD with opposite effects on UC (odds ratio [OR] 0.58) and CD (OR 1.76), whereas in other ethnic groups with different genetic susceptibility, smoking may play a lesser role. There is a protective effect of previous appendectomy with confirmed appendicitis (reduction of 13–26%), particularly at a young age, on the development of UC across different geographical regions and populations. There is a modest association with the development of CD. Oral contraceptive use is associated with the risk of CD (OR 1.4). The association between oral contraceptive use and UC is limited to women with a history of smoking. There is an association between antibiotic use and the development of childhood IBD with children who received one or more dispensations of antibiotics during the first year of life having a 2.9-fold increase in the risk of developing IBD during childhood. Breastfeeding may also protect against the development of IBD. These factors are consistent with the rapid increase in IBD incidence recently noted during the first decade of life. Infectious gastroenteritis with pathogens (e.g., *Salmonella*, *Shigella*, *Campylobacter* spp., *Clostridium difficile*) increases IBD risk by two- to threefold. Diets high in animal protein, sugars, sweets, oils, fish and shellfish, and dietary fat, especially ω -6 fatty acids, and low in ω -3 fatty acids have been implicated in increasing the risk of IBD.

IBD is a familial disease in 5–10% of patients (Fig. 351-2). Some of these patients may exhibit early-onset disease during the first decade of life and, in CD, a concordance of anatomic site and clinical type within families. In the remainder of patients, IBD is observed in the absence of a family history (i.e., sporadic disease). If a patient has IBD, the lifetime risk that a first-degree relative will be affected is ~10%. If two parents have IBD, each child has a 36% chance of being affected. In twin studies, 38–58% of monozygotic twins are concordant for CD and 6–18% are concordant for UC, whereas 4% of dizygotic twins are concordant for CD and 0–2% are concordant for UC in Swedish and Danish cohorts. The risks of developing IBD are higher in first-degree relatives of Jewish versus non-Jewish patients: 7.8% versus 5.2% for CD and 4.5% versus 1.6% for UC.

GLOBAL CONSIDERATIONS: IBD PHENOTYPES


 There are racial differences in IBD location and behavior that may reflect underlying genetic variations and have important implications for diagnosis and management of disease. For example, African-American patients are more likely than non-Hispanic whites to develop esophagogastroduodenal CD, colorectal disease, and perianal disease and are less likely to have ileal involvement. They are also at higher risk for uveitis and sacroiliitis. Hispanics have a higher prevalence of perianal CD and erythema nodosum and a more proximal extent of disease. Fistulizing CD has been reported in nearly one-third of Hispanic patients, up to one-quarter of African-American patients, and up to one-half of Asian patients. Both African-American

TABLE 351-1 EPIDEMIOLOGY OF IBD

	Ulcerative Colitis	Crohn's Disease
Incidence (North America) per person-years	0–19.2 per 100,000	0–20.2 per 100,000
Age of onset	Second to fourth decades and seventh to ninth decades	Second to fourth decades and seventh to ninth decades
Ethnicity	Jewish > non-Jewish white > African American > Hispanic > Asian	
Female/male ratio	0.51–1.58	0.34–1.65
Smoking	May prevent disease (odds ratio 0.58)	May cause disease (odds ratio 1.76)
Oral contraceptives	No increased risk	Odds ratio 1.4
Appendectomy	Protective (risk reduction of 13–26%)	Not protective
Monozygotic twins	6–18% concordance	38–58% concordance
Dizygotic twins	0–2% concordance	4% concordance
Antibiotic use in the first year of life	2.9x the risk of developing childhood IBD	

Abbreviation: IBD, inflammatory bowel disease.

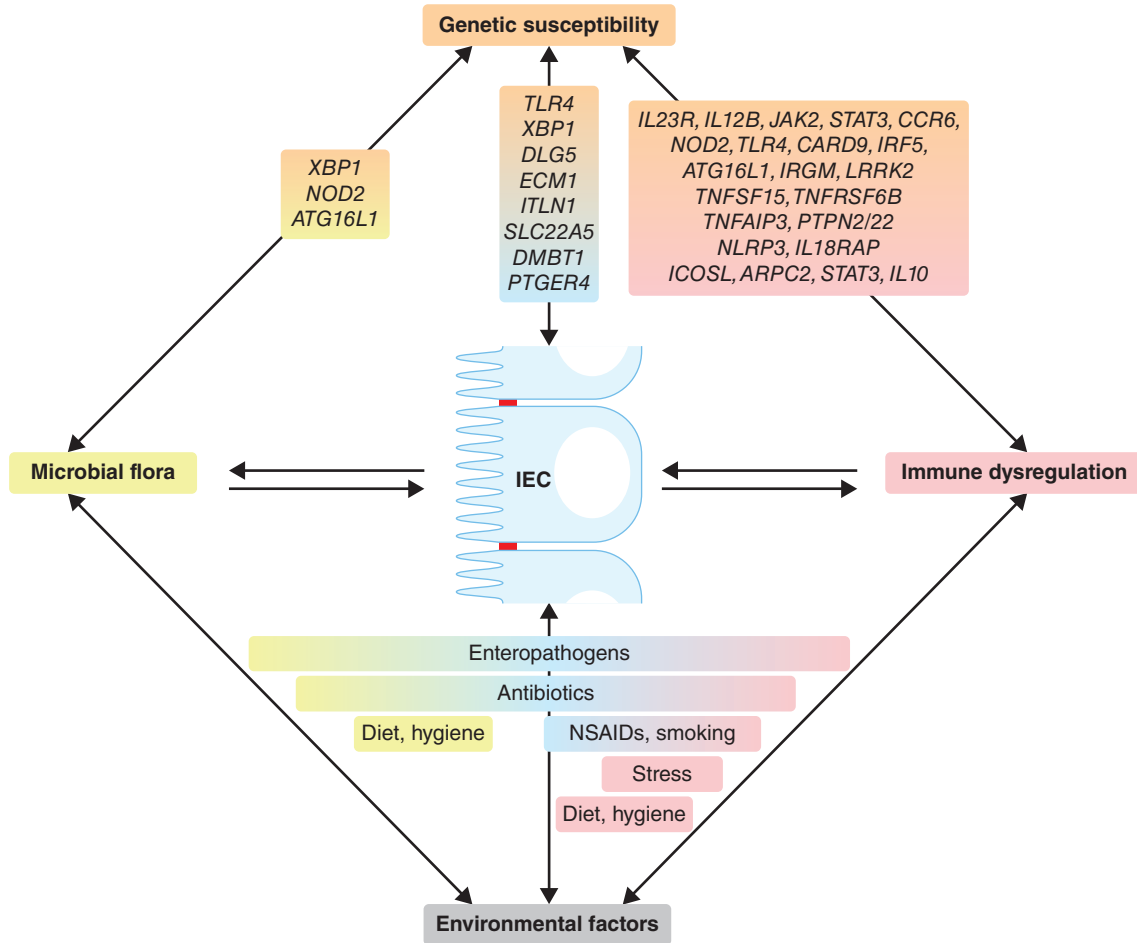


FIGURE 351-1 Pathogenesis of inflammatory bowel disease (IBD). In IBD, the tridirectional relationship between the commensal flora (microbiota), intestinal epithelial cells (IEC), and mucosal immune system is dysregulated, leading to chronic inflammation. Each of these three factors is affected by genetic and environmental factors that determine risk for the disease. NSAIDs, nonsteroidal anti-inflammatory drugs. (Adapted from A Kaser et al: *Annu Rev Immunol* 28:573, 2010.)

and Hispanic CD patients, but not UC patients, had a lower prevalence of family history of IBD than their white counterparts. There are few data on all aspects of disease in Hispanics, in the incidence and prevalence of IBD in African Americans, and in Asians with IBD outside Asia. These ethnic variations implicate the importance of different genetic and/or environmental factors in the pathogenesis of this disorder.

ETIOLOGY AND PATHOGENESIS

Under physiologic conditions, homeostasis normally exists between the commensal microbiota, epithelial cells that line the interior of the intestines (intestinal epithelial cells [IECs]) and immune cells within the tissues (Fig. 351-1). A consensus hypothesis is that each of these three major host compartments that function together as an integrated “supraorganism” (microbiota, IECs, and immune cells) are affected by specific environmental (e.g., smoking, antibiotics, enteropathogens) and genetic factors that, in a susceptible host, cumulatively and interactively disrupt homeostasis, which in so doing culminates in a chronic state of dysregulated inflammation; that is IBD. Although chronic activation of the mucosal immune system may represent an appropriate response to an infectious agent, a search for such an agent has thus far been unrewarding in IBD. As such, IBD is currently considered an inappropriate immune response to the endogenous (autochthonous) commensal microbiota within the intestines, with or without some component of autoimmunity. Importantly, the normal, uninfamed intestines contain a large number of immune cells that are in a unique state of activation, in which the gut is restrained from full immunologic responses to the commensal microbiota and dietary antigens by very powerful regulatory pathways that function within the immune system (e.g., T regulatory cells that express the FoxP3 transcription factor and suppress inflammation). During the course of infections or other environmental stimuli in the normal host, full activation of the gut-associated lymphoid tissues occurs but is rapidly superseded by dampening of the immune response and tissue repair. In IBD such processes may not be regulated normally.

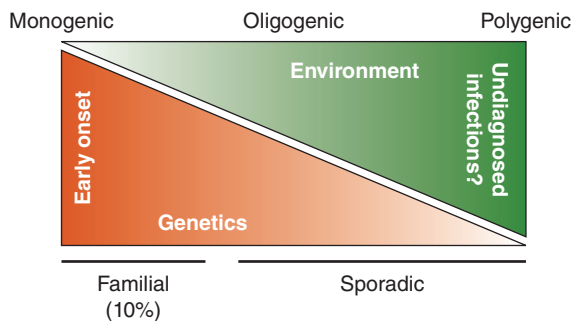


FIGURE 351-2 A model for the syndromic nature of inflammatory bowel disease. Genetic and environmental factors variably influence the development and phenotypic manifestations of IBD. At the one extreme, IBD is exemplified as a simple Mendelian disorder as observed in “early-onset IBD” due to single gene defects such as *IL10*, *IL10RA*, and *IL10RB*; and at the other extreme, it may be exemplified by as yet to be described emerging infectious diseases. (Adapted from A Kaser et al: *Dig Dis* 28:395, 2010.)

GENETIC CONSIDERATIONS



The genetic underpinning of IBD is known from its occurrence in the context of several genetic syndromes and the development of severe, refractory IBD in early life in the setting of single gene defects that affect the immune system (Table 351-2). In addition, IBD has a familial origin in at least 10% of afflicted individuals (Fig. 351-2). In the majority of patients, IBD is considered to be a polygenic disorder that gives rise to multiple clinical subgroups within UC and CD. A variety of genetic approaches including candidate gene studies, linkage analysis, and genome-wide association studies (GWASs) that focus on the identification of disease-associated, single-nucleotide polymorphisms (SNPs) within the human genome and, more recently, whole-genome sequencing have elucidated many of the genetic factors that affect risk for these diseases. GWASs have, to date, identified 163 genetic loci with 100 of these loci observed to be associated with both disease phenotypes (Table 351-3). The remainder are specific for either CD (30 loci) or UC (20 loci). These genetic similarities account for the overlapping immunopathogenesis and consequently epidemiologic observations of both diseases in the same families and similarities in response to therapies. Because the specific causal variants for each identified gene or locus are largely unknown, it is not clear whether the similarities in the genetic risk factors associated with CD and UC that are observed are shared at structural or functional levels. The risk conferred by each identified gene or locus is unequal and generally small, such that only ~20% of the genetic variance is considered to be explained by the current genetic information. Further, many of the genetic risk factors identified are also observed to be associated with risk for other immune-mediated diseases, suggesting that related immunogenetic pathways are involved in the pathogenesis of multiple different disorders accounting for the common responsiveness to similar types of biologic therapies (e.g., anti-tumor necrosis factor therapies) and possibly the simultaneous occurrence of these disorders. The diseases and the genetic risk factors that are shared with IBD include rheumatoid arthritis (*TNFAIP3*), psoriasis (*IL23R*, *IL12B*), ankylosing spondylitis (*IL23R*), type 1 diabetes mellitus (*IL10*, *PTPN2*), asthma (*ORMDL3*), and systemic lupus erythematosus (*TNFAIP3*, *IL10*) among others.

The genetic factors defined to date that are recognized to mediate risk for IBD have highlighted the importance of several common mechanisms of disease (Table 351-3). These include the following: those genes that are associated with fundamental cell biologic

processes such as endoplasmic reticulum (ER) and metabolic stress (e.g., *XBPI*, *ORMDL3*, *OCTN*), which serve to regulate the secretory activity of cells involved in responses to the commensal microbiota such as Paneth and goblet cells and the manner in which intestinal cells respond to the metabolic products of bacteria; those associated with innate immunity and autophagy (e.g., *NOD2*, *ATG16L1*, *IRGM*, *JAK2*, *STAT3*) that function in innate immune cells (both parenchymal and hematopoietic) to respond to and effectively clear bacteria, mycobacteria, and viruses; those that are associated with the regulation of adaptive immunity (e.g., *IL23R*, *IL12B*, *IL10*, *PTPN2*), which regulate the balance between inflammatory and anti-inflammatory (regulatory) cytokines; and, finally, those that are involved in the development and resolution of inflammation (e.g., *MST1*, *CCR6*, *TNFAIP3*, *PTGER4*) and ultimately leukocyte recruitment and inflammatory mediator production. Some of these loci are associated with specific subtypes of disease such as the association between *NOD2* polymorphisms and fibrostenosing CD or *ATG16L1* and fistulizing disease, especially within the ileum. However, the clinical utility of these genetic risk factors for the diagnosis or determination of prognosis and therapeutic responses remains to be defined.

COMMENSAL MICROBIOTA AND IBD

The endogenous commensal microbiota within the intestines plays a central role in the pathogenesis of IBD. Humans are born sterile and acquire their commensal microbiota initially from the mother during egress through the birth canal and subsequently from environmental sources. A stable configuration of up to 1000 species of bacteria that achieves a biomass of approximately 10^{12} colony-forming units per gram of feces is achieved by 3 years of age, which likely persists into adult life, with each individual human possessing a unique combination of species. In addition, the intestines contain other microbial life forms including archae, viruses, and protists. The microbiota is thus considered as a critical and sustaining component of the organism. The establishment and maintenance of the intestinal microbiota composition and function is under the control of host (e.g., immune and epithelial responses), environmental (e.g., diet and antibiotics), and likely genetic (e.g., *NOD2*) factors (Fig. 351-1). In turn, the microbiota, through its structural components and metabolic activity, has major influences on the epithelial and immune function of the host, which, through epigenetic effects, may have durable consequences. During early life when the commensal microbiota is being established, these microbial effects on the host may be particularly important in determining later life risk for IBD. Specific components of the microbiota can promote or protect from disease. The commensal microbiota in patients with both UC and CD is demonstrably different from nonafflicted individuals, a state of dysbiosis, suggesting the presence of microorganisms that drive disease (e.g., Proteobacteria such as enteroinvasive and adherent *Escherichia coli*) and to which the immune response is directed and/or the loss of microorganisms that hinder inflammation (e.g., Firmicutes such as *Faecalibacterium prausnitzii*). Many of the changes in the commensal microbiota occur as a consequence of the inflammation. In addition, agents that alter the intestinal microbiota such as metronidazole, ciprofloxacin, and elemental diets, may improve CD. CD may also respond to fecal diversion, demonstrating the ability of luminal contents to exacerbate disease.

DEFECTIVE IMMUNE REGULATION IN IBD

The mucosal immune system is normally unreactive to luminal contents due to oral (mucosal) tolerance. When soluble antigens are administered orally rather than subcutaneously or intramuscularly, antigen-specific nonresponsiveness is induced. Multiple mechanisms are involved in the induction of oral tolerance and include deletion or anergy of antigen-reactive T cells or induction of CD4+ T cells that suppress gut inflammation (e.g., T regulatory cells expressing the FoxP3 transcription factor) that secrete anti-inflammatory cytokines such as interleukin (IL) 10, IL-35, and transforming growth factor β (TGF- β). Oral tolerance may be responsible for the lack of immune responsiveness to dietary antigens and the commensal microbiota

TABLE 351-2 PRIMARY GENETIC DISORDERS ASSOCIATED WITH IBD

Name	Genetic Association	Phenotype
Turner's syndrome	Loss of part or all of X chromosome	Associated with UC and colonic CD
Hermansky-Pudlak	Autosomal recessive chromosome 10q23	Granulomatous colitis, oculocutaneous albinism, platelet dysfunction, pulmonary fibrosis
Wiskott-Aldrich syndrome (WAS)	X-linked recessive disorder, loss of WAS protein function	Colitis, immunodeficiency, severely dysfunctional platelets, and thrombocytopenia
Glycogen storage disease	Deficiency of the glucose-6-phosphate transport protein type B1	Granulomatous colitis, presents in infancy with hypoglycemia, growth failure, hepatomegaly, and neutropenia
Immune dysregulation polyendocrinopathy, enteropathy X-linked (IPEX)	Loss of FoxP3 transcription factor and T regulatory cell function	UC-like autoimmune enteropathy, with endocrinopathy (neonatal type 1 diabetes or thyroiditis), dermatitis
Early-onset IBD	Deficient IL-10 and IL-10 receptor function	Severe, refractory IBD in early life

Abbreviations: CD, Crohn's disease; IBD, inflammatory bowel disease; IL, interleukin; UC, ulcerative colitis.

TABLE 351-3 EXAMPLES OF GENETIC LOCI ASSOCIATED WITH CD AND/OR UC

Chromosome	Putative Gene	Gene Name	Protein Function	CD	UC
ER Stress and Metabolism					
5q31	<i>SLC22A5</i>	Solute carrier family 22, member 5	β carnitine transporter	+	
7p21	<i>AGR2</i>	Anterior gradient 2	ER stress	+	+
17q21	<i>ORMDL3</i>	Orosomucoid related member 1-like 3	ER stress and lipid synthesis	+	+
22q12	<i>XBP1</i>	X-box binding protein 1	ER stress	+	+
1q23	<i>ITLN1</i>	Intelectin 1	Bacterial binding	+	
2q37	<i>ATG16L1</i>	ATG16 autophagy related 16-like 1	Autophagy	+	
5q33	<i>IRGM</i>	Immunity-related GTPase family, M	Autophagy	+	
9p24	<i>JAK2</i>	Janus kinase 2	IL-6R and IL-23R signaling	+	+
12q12	<i>LRRK2</i>	Leucine-rich repeat kinase 2	Autophagy?	+	
16q12	<i>NOD2</i>	Nucleotide-binding oligomerization domain containing 2	Bacterial sensing and autophagy activation	+	
17q21	<i>STAT3</i>	Signal transducer and activator of transcription 3	IL-6R, IL-23R, and IL-10R signaling	+	+
Adaptive Immunity					
1p31	<i>IL23R</i>	Interleukin 23 receptor	Th17 cell stimulation	+	+
1q32	<i>IL10</i>	Interleukin 10	Treg-associated cytokine		+
5q33	<i>IL12B</i>	Interleukin 12B	IL-12 p40 chain of IL-12/IL-23	+	+
18p11	<i>PTPN2</i>	Protein tyrosine phosphatase, nonreceptor type 2	T cell regulation	+	
Inflammation					
3p21	<i>MST1</i>	Macrophage stimulating 1	Macrophage activation	+	+
5p13	<i>PTGER4</i>	Prostaglandin E receptor 4	PGE ₂ receptor	+	+
6q23	<i>TNFAIP3</i>	Tumor necrosis factor, alpha-induced protein 3 (A20)	Toll-like receptor regulation	+	
6q27	<i>CCR6</i>	Chemokine (C-C motif) receptor 6	Dendritic cell migration	+	

Abbreviations: CD, Crohn's disease; ER, endoplasmic reticulum; GTPase, guanosine triphosphatase; IL, interleukin; PGE₂, prostaglandin E₂; UC, ulcerative colitis.

Source: Adapted from A Kaser et al: *Ann Rev Immunol* 28:573, 2010; B Khor et al: *Nature* 474:307, 2011; and L Jostins et al: *Nature* 491:119, 2012.

in the intestinal lumen. In IBD this suppression of inflammation is altered, leading to uncontrolled inflammation. The mechanisms of this regulated immune suppression are incompletely known.

Gene knockout (^{-/-}) or transgenic (Tg) mouse models of IBD, which include those that are directed at genes demonstrated to be associated with risk for the human disease, have revealed that deleting specific cytokines (e.g., IL-2, IL-10, TGF- β) or their receptors, deleting molecules associated with T cell antigen recognition (e.g., T cell antigen receptors), or interfering with IEC barrier function and the regulation of responses to commensal bacteria (e.g., XBP1, N-cadherin, mucus glycoprotein, or nuclear factor- κ B [NF- κ B]) leads to spontaneous colitis or enteritis. In the majority of circumstances, intestinal inflammation in these animal models requires the presence of the commensal microbiota. Thus, a variety of specific alterations can lead to immune activation by commensal microbiota and inflammation directed at the intestines in mice. How these relate to human IBD remains to be defined, but they are consistent with inappropriate responses of the genetically susceptible host to the commensal microbiota.

In both UC and CD, an inflammatory pathway thus likely emerges from the genetic predisposition that is associated with inappropriate innate immune and epithelial sensing and reactivity to commensal bacteria that secrete inflammatory mediators together with inadequate regulatory pathways that lead to activated CD4⁺ and CD8⁺ T cells within the epithelium and lamina propria that altogether secrete excessive quantities of inflammatory cytokines relative to anti-inflammatory cytokines. Some cytokines activate other inflammatory cells (macrophages and B cells), and others act indirectly to recruit other lymphocytes, inflammatory leukocytes, and mononuclear cells from the bloodstream into the gut through interactions between homing receptors on leukocytes (e.g., α 4 β 7 integrin) and addressins on vascular endothelium (e.g., MadCAM1). Consistent with this, neutralization of tumor necrosis factor (TNF) or α 4 β 7 integrin demonstrate therapeutic efficacy in IBD. CD4⁺ T helper (T_H) cells that promote inflammation are of three major types, all of which may be associated with colitis in animal models and perhaps humans: T_H1 cells (secrete interferon [IFN] γ), T_H2 cells (secrete IL-4, IL-5, IL-13), and T_H17 cells (secrete IL-17, IL-21). T_H1 cells induce transmural granulomatous inflammation that

resembles CD; T_H2 cells, and related natural killer T cells that secrete IL-13, induce superficial mucosal inflammation resembling UC in animal models; and T_H17 cells may be responsible for neutrophilic recruitment. However, neutralization of the cytokines produced by these cells, such as IFN- γ or IL-17, has yet to show efficacy in therapeutic trials. Each of these T cell subsets cross-regulate each other. The T_H1 cytokine pathway is initiated by IL-12, a key cytokine in the pathogenesis of experimental models of mucosal inflammation. IL-4 and IL-23, together with IL-6 and TGF- β , induce T_H2 and T_H17 cells, respectively, and IL-23 inhibits the suppressive function of regulatory T cells. Activated macrophages secrete TNF and IL-6. These characteristics of the immune response in IBD explain the beneficial therapeutic effects of antibodies to block proinflammatory cytokines or the signaling by their receptors (e.g., anti-TNF, anti-IL-12, anti-IL-23, anti-IL-6, or Janus kinase [JAK] inhibitors) or molecules associated with leukocyte recruitment (e.g., anti- α 4 β 7), or the use of cytokines that inhibit inflammation and promote regulatory T cells (e.g., IL-10) or promote intestinal barrier function and may be beneficial to humans with intestinal inflammation.

THE INFLAMMATORY CASCADE IN IBD

Once initiated in IBD by abnormal innate immune sensing of bacteria by parenchymal cells (e.g., IECs) and hematopoietic cells (e.g., dendritic cells), the immune inflammatory response is perpetuated by T cell activation. A sequential cascade of inflammatory mediators extends the response; each step is a potential target for therapy. Inflammatory cytokines such as IL-1, IL-6, and TNF have diverse effects on tissues. They promote fibrogenesis, collagen production, activation of tissue metalloproteinases, and the production of other inflammatory mediators; they also activate the coagulation cascade in local blood vessels (e.g., increased production of von Willebrand's factor). These cytokines are normally produced in response to infection but are usually turned off or inhibited at the appropriate time to limit tissue damage. In IBD their activity is not regulated, resulting in an imbalance between the proinflammatory and anti-inflammatory mediators. Therapies such as the 5-aminosalicylic acid (5-ASA) compounds and glucocorticoids are potent inhibitors

of these inflammatory mediators through inhibition of transcription factors such as NF- κ B that regulate their expression.

PATHOLOGY

ULCERATIVE COLITIS: MACROSCOPIC FEATURES

UC is a mucosal disease that usually involves the rectum and extends proximally to involve all or part of the colon. About 40–50% of patients have disease limited to the rectum and rectosigmoid, 30–40% have disease extending beyond the sigmoid but not involving the whole colon, and 20% have a total colitis. Proximal spread occurs in continuity without areas of uninvolved mucosa. When the whole colon is involved, the inflammation extends 2–3 cm into the terminal ileum in 10–20% of patients. The endoscopic changes of *backwash ileitis* are superficial and mild and are of little clinical significance. Although variations in macroscopic activity may suggest skip areas, biopsies from normal-appearing mucosa are usually abnormal. Thus, it is important to obtain multiple biopsies from apparently uninvolved mucosa, whether proximal or distal, during endoscopy. One caveat is that effective medical therapy can change the appearance of the mucosa such that either skip areas or the entire colon can be microscopically normal.

With mild inflammation, the mucosa is erythematous and has a fine granular surface that resembles sandpaper. In more severe disease, the mucosa is hemorrhagic, edematous, and ulcerated (Fig. 351-3). In long-standing disease, inflammatory polyps (pseudopolyps) may be present as a result of epithelial regeneration. The mucosa may appear normal in remission, but in patients with many years of disease it appears atrophic and featureless, and the entire colon becomes narrowed and shortened. Patients with fulminant disease can develop a toxic colitis or megacolon where the bowel wall thins and the mucosa is severely ulcerated; this may lead to perforation.

ULCERATIVE COLITIS: MICROSCOPIC FEATURES

Histologic findings correlate well with the endoscopic appearance and clinical course of UC. The process is limited to the mucosa and superficial submucosa, with deeper layers unaffected except in fulminant disease. In UC, two major histologic features suggest chronicity and help distinguish it from infectious or acute self-limited colitis. First, the crypt architecture of the colon is distorted; crypts may be bifid and reduced in number, often with a gap between the crypt bases and the muscularis mucosae. Second, some patients have basal plasma cells and multiple basal lymphoid aggregates. Mucosal vascular congestion, with edema and focal hemorrhage, and an inflammatory cell infiltrate of neutrophils, lymphocytes, plasma cells, and macrophages may be present.

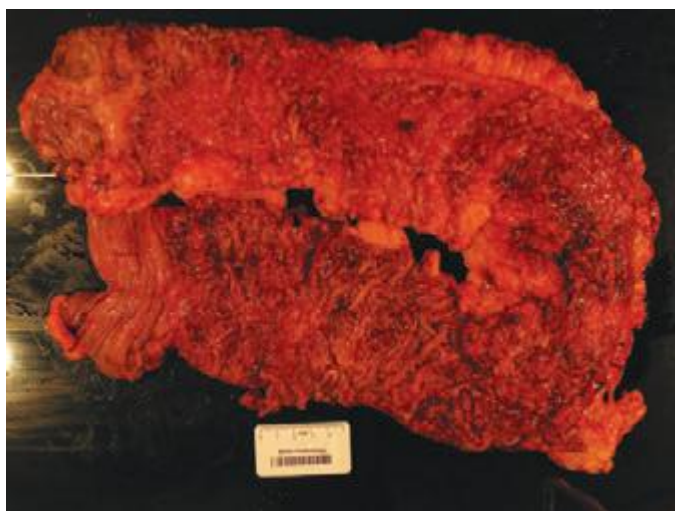


FIGURE 351-3 Ulcerative colitis. Diffuse (nonsegmental) mucosal disease, with broad areas of ulceration. The bowel wall is not thickened, and there is no cobblestoning. (Courtesy of Dr. R. Odze, Division of Gastrointestinal Pathology, Department of Pathology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

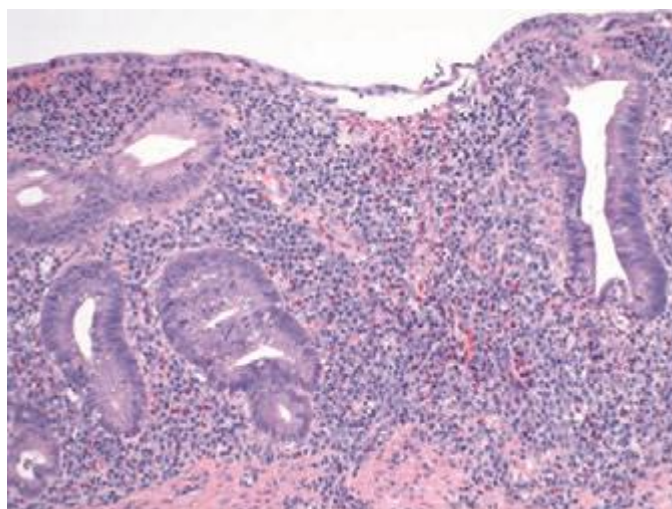


FIGURE 351-4 Medium-power view of colonic mucosa in ulcerative colitis showing diffuse mixed inflammation, basal lymphoplasmacytosis, crypt atrophy and irregularity, and superficial erosion. These features are typical of chronic active ulcerative colitis. (Courtesy of Dr. R. Odze, Division of Gastrointestinal Pathology, Department of Pathology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

The neutrophils invade the epithelium, usually in the crypts, giving rise to cryptitis and, ultimately, to crypt abscesses (Fig. 351-4). Ileal changes in patients with backwash ileitis include villous atrophy and crypt regeneration with increased inflammation, increased neutrophil and mononuclear inflammation in the lamina propria, and patchy cryptitis and crypt abscesses.

CROHN'S DISEASE: MACROSCOPIC FEATURES

CD can affect any part of the gastrointestinal (GI) tract from the mouth to the anus. Some 30–40% of patients have small bowel disease alone, 40–55% have disease involving both the small and large intestines, and 15–25% have colitis alone. In the 75% of patients with small intestinal disease, the terminal ileum is involved in 90%. Unlike UC, which almost always involves the rectum, the rectum is often spared in CD. CD is segmental with skip areas in the midst of diseased intestine (Fig. 351-5). Perirectal fistulas, fissures, abscesses, and anal stenosis are present in one-third of patients with CD, particularly those with



FIGURE 351-5 Crohn's disease of the colon showing thickening of the wall, with stenosis, linear serpiginous ulcers and cobblestoning of the mucosa. (Courtesy of Dr. R. Odze, Division of Gastrointestinal Pathology, Department of Pathology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

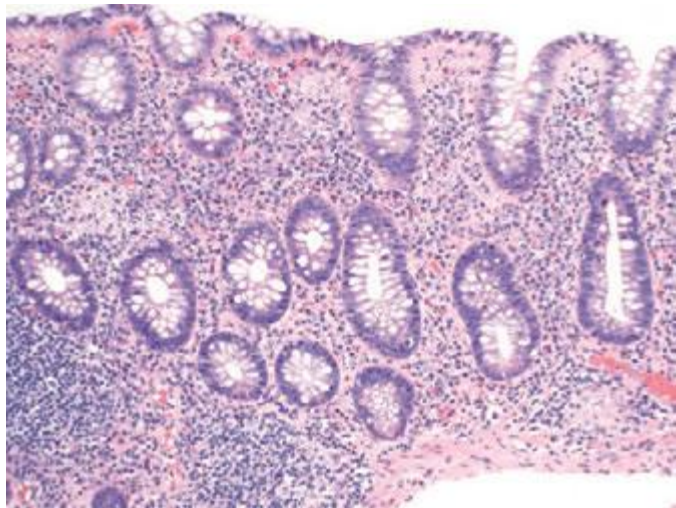


FIGURE 351-6 Medium-power view of Crohn's colitis showing mixed acute and chronic inflammation, crypt atrophy, and multiple small epithelioid granulomas in the mucosa. (Courtesy of Dr. R Odze, Division of Gastrointestinal Pathology, Department of Pathology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

colonic involvement. Rarely, CD may also involve the liver and the pancreas.

Unlike UC, CD is a transmural process. Endoscopically, aphthous or small superficial ulcerations characterize mild disease; in more active disease, stellate ulcerations fuse longitudinally and transversely to demarcate islands of mucosa that frequently are histologically normal. This “cobblestone” appearance is characteristic of CD, both endoscopically and by barium radiography. As in UC, pseudopolyps can form in CD.

Active CD is characterized by focal inflammation and formation of fistula tracts, which resolve by fibrosis and stricturing of the bowel. The bowel wall thickens and becomes narrowed and fibrotic, leading to chronic, recurrent bowel obstructions. Projections of thickened mesentery encase the bowel (“creeping fat”), and serosal and mesenteric inflammation promotes adhesions and fistula formation.

CROHN'S DISEASE: MICROSCOPIC FEATURES

The earliest lesions are aphthoid ulcerations and focal crypt abscesses with loose aggregations of macrophages, which form noncaseating granulomas in all layers of the bowel wall (Fig. 351-6). Granulomas can be seen in lymph nodes, mesentery, peritoneum, liver, and pancreas. Although granulomas are a pathognomonic feature of CD, they are rarely found on mucosal biopsies. Surgical resection reveals granulomas in about one-half of cases. Other histologic features of CD include submucosal or subserosal lymphoid aggregates, particularly away from areas of ulceration, gross and microscopic skip areas, and transmural inflammation that is accompanied by fissures that penetrate deeply into the bowel wall and sometimes form fistulous tracts or local abscesses.

CLINICAL PRESENTATION

ULCERATIVE COLITIS

Signs and Symptoms The major symptoms of UC are diarrhea, rectal bleeding, tenesmus, passage of mucus, and crampy abdominal pain. The severity of symptoms correlates with the extent of disease. Although UC can present acutely, symptoms usually have been present for weeks to months. Occasionally, diarrhea and bleeding are so intermittent and mild that the patient does not seek medical attention.

Patients with proctitis usually pass fresh blood or blood-stained mucus, either mixed with stool or streaked onto the surface of a normal or hard stool. They also have tenesmus, or urgency with a feeling of incomplete evacuation, but rarely have abdominal pain. With proctitis or proctosigmoiditis, proximal transit slows, which

TABLE 351-4 ULCERATIVE COLITIS: DISEASE PRESENTATION

	Mild	Moderate	Severe
Bowel movements	<4 per day	4–6 per day	>6 per day
Blood in stool	Small	Moderate	Severe
Fever	None	<37.5°C mean (<99.5°F)	>37.5°C mean (>99.5°F)
Tachycardia	None	<90 mean pulse	>90 mean pulse
Anemia	Mild	>75%	≤75%
Sedimentation rate	<30 mm		>30 mm
Endoscopic appearance	Erythema, decreased vascular pattern, fine granularity	Marked erythema, coarse granularity, absent vascular markings, contact bleeding, no ulcerations	Spontaneous bleeding, ulcerations

may account for the constipation commonly seen in patients with distal disease.

When the disease extends beyond the rectum, blood is usually mixed with stool or grossly bloody diarrhea may be noted. Colonic motility is altered by inflammation with rapid transit through the inflamed intestine. When the disease is severe, patients pass a liquid stool containing blood, pus, and fecal matter. Diarrhea is often nocturnal and/or postprandial. Although severe pain is not a prominent symptom, some patients with active disease may experience vague lower abdominal discomfort or mild central abdominal cramping. Severe cramping and abdominal pain can occur with severe attacks of the disease. Other symptoms in moderate to severe disease include anorexia, nausea, vomiting, fever, and weight loss.

Physical signs of proctitis include a tender anal canal and blood on rectal examination. With more extensive disease, patients have tenderness to palpation directly over the colon. Patients with a toxic colitis have severe pain and bleeding, and those with megacolon have hepatic tympany. Both may have signs of peritonitis if a perforation has occurred. The classification of disease activity is shown in Table 351-4.

Laboratory, Endoscopic, and Radiographic Features Active disease can be associated with a rise in acute-phase reactants (C-reactive protein [CRP]), platelet count, and erythrocyte sedimentation rate (ESR), and a decrease in hemoglobin. Fecal lactoferrin is a highly sensitive and specific marker for detecting intestinal inflammation. Fecal calprotectin levels correlate well with histologic inflammation, predict relapses, and detect pouchitis. Both fecal lactoferrin and calprotectin are becoming an integral part of IBD management and are used frequently to rule out active inflammation versus symptoms of irritable bowel or bacterial overgrowth. In severely ill patients, the serum albumin level will fall rather quickly. Leukocytosis may be present but is not a specific indicator of disease activity. Proctitis or proctosigmoiditis rarely causes a rise in CRP. Diagnosis relies on the patient's history; clinical symptoms; negative stool examination for bacteria, *C. difficile* toxin, and ova and parasites; sigmoidoscopic appearance (see Fig. 345-4A); and histology of rectal or colonic biopsy specimens.

Sigmoidoscopy is used to assess disease activity and is usually performed before treatment. If the patient is not having an acute flare, colonoscopy is used to assess disease extent and activity (Fig. 351-7). Endoscopically mild disease is characterized by erythema, decreased vascular pattern, and mild friability. Moderate disease is characterized by marked erythema, absent vascular pattern, friability and erosions, and severe disease by spontaneous bleeding and ulcerations. Histologic features change more slowly than clinical features but can also be used to grade disease activity.

The earliest radiologic change of UC seen on single-contrast barium enema is a fine mucosal granularity. With increasing severity, the mucosa becomes thickened, and superficial ulcers are seen. Deep ulcerations can appear as “collar-button” ulcers, which indicate that the ulceration has penetrated the mucosa. Haustral folds may be normal in mild disease, but as activity progresses they become edematous and thickened. Loss of haustration can occur, especially in patients



FIGURE 351-7 Colonoscopy with acute ulcerative colitis: severe colon inflammation with erythema, friability, and exudates. (Courtesy of Dr. M. Hamilton, Gastroenterology Division, Department of Medicine, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

with long-standing disease. In addition, the colon becomes shortened and narrowed. Polyps in the colon may be postinflammatory polyps or pseudopolyps, adenomatous polyps, or carcinoma.

Computed tomography (CT) scanning or magnetic resonance imaging (MRI) is not as helpful as endoscopy in making the diagnosis of UC, but typical findings include mild mural thickening (<1.5 cm), inhomogeneous wall density, absence of small bowel thickening, increased perirectal and presacral fat, target appearance of the rectum, and adenopathy.

Complications Only 15% of patients with UC present initially with catastrophic illness. Massive hemorrhage occurs with severe attacks of disease in 1% of patients, and treatment for the disease usually stops the bleeding. However, if a patient requires 6–8 units of blood within 24–48 h, colectomy is indicated. *Toxic megacolon* is defined as a transverse or right colon with a diameter of >6 cm, with loss of haustration in patients with severe attacks of UC. It occurs in about 5% of attacks and can be triggered by electrolyte abnormalities and narcotics. About 50% of acute dilations will resolve with medical therapy alone, but urgent colectomy is required for those that do not improve. Perforation is the most dangerous of the local complications, and the physical signs of peritonitis may not be obvious, especially if the patient is receiving glucocorticoids. Although perforation is rare, the mortality rate for perforation complicating a toxic megacolon is about 15%. In addition, patients can develop a toxic colitis and such severe ulcerations that the bowel may perforate without first dilating.

Strictures occur in 5–10% of patients and are always a concern in UC because of the possibility of underlying neoplasia. Although benign strictures can form from the inflammation and fibrosis of UC, strictures that are impassable with the colonoscope should be presumed malignant until proven otherwise. A stricture that prevents passage of the colonoscope is an indication for surgery. UC patients occasionally develop anal fissures, perianal abscesses, or hemorrhoids, but the occurrence of extensive perianal lesions should suggest CD.

CROHN'S DISEASE

Signs and Symptoms Although CD usually presents as acute or chronic bowel inflammation, the inflammatory process evolves toward one of two patterns of disease: a fibrostenotic obstructing pattern or a penetrating fistulous pattern, each with different treatments and prognoses. The site of disease influences the clinical manifestations.

ILEOCOLITIS Because the most common site of inflammation is the terminal ileum, the usual presentation of ileocolitis is a chronic history of recurrent episodes of right lower quadrant pain and diarrhea.

Sometimes the initial presentation mimics acute appendicitis with pronounced right lower quadrant pain, a palpable mass, fever, and leukocytosis. Pain is usually colicky; it precedes and is relieved by defecation. A low-grade fever is usually noted. High-spiking fever suggests intraabdominal abscess formation. Weight loss is common—typically 10–20% of body weight—and develops as a consequence of diarrhea, anorexia, and fear of eating.

An inflammatory mass may be palpated in the right lower quadrant of the abdomen. The mass is composed of inflamed bowel, adherent and indurated mesentery, and enlarged abdominal lymph nodes. Extension of the mass can cause obstruction of the right ureter or bladder inflammation, manifested by dysuria and fever. Edema, bowel wall thickening, and fibrosis of the bowel wall within the mass account for the radiographic “string sign” of a narrowed intestinal lumen.

Bowel obstruction may take several forms. In the early stages of disease, bowel wall edema and spasm produce intermittent obstructive manifestations and increasing symptoms of postprandial pain. Over several years, persistent inflammation gradually progresses to fibrostenotic narrowing and stricture. Diarrhea will decrease and be replaced by chronic bowel obstruction. Acute episodes of obstruction occur as well, precipitated by bowel inflammation and spasm or sometimes by impaction of undigested food or medication. These episodes usually resolve with intravenous fluids and gastric decompression.

Severe inflammation of the ileocecal region may lead to localized wall thinning, with microperforation and fistula formation to the adjacent bowel, the skin, or the urinary bladder, or to an abscess cavity in the mesentery. Enterovesical fistulas typically present as dysuria or recurrent bladder infections or, less commonly, as pneumaturia or fecaluria. Enterocutaneous fistulas follow tissue planes of least resistance, usually draining through abdominal surgical scars. Enterovaginal fistulas are rare and present as dyspareunia or as a feculent or foul-smelling, often painful vaginal discharge. They are unlikely to develop without a prior hysterectomy.

JEJUNOILEITIS Extensive inflammatory disease is associated with a loss of digestive and absorptive surface, resulting in malabsorption and steatorrhea. Nutritional deficiencies can also result from poor intake and enteric losses of protein and other nutrients. Intestinal malabsorption can cause anemia, hypoalbuminemia, hypocalcemia, hypomagnesemia, coagulopathy, and hyperoxaluria with nephrolithiasis in patients with an intact colon. Many patients need to take oral and often intravenous iron. Vertebral fractures are caused by a combination of vitamin D deficiency, hypocalcemia, and prolonged glucocorticoid use. Pellagra from niacin deficiency can occur in extensive small-bowel disease, and malabsorption of vitamin B₁₂ can lead to megaloblastic anemia and neurologic symptoms. Other important nutrients to measure and replete if low are folate and vitamins A, E, and K. Levels of minerals such as zinc, selenium, copper, and magnesium are often low in patients with extensive small-bowel inflammation or resections, and these should be repleted as well. Most patients should take a daily multivitamin, calcium, and vitamin D supplements.

Diarrhea is characteristic of active disease; its causes include (1) bacterial overgrowth in obstructive stasis or fistulization, (2) bile-acid malabsorption due to a diseased or resected terminal ileum, and (3) intestinal inflammation with decreased water absorption and increased secretion of electrolytes.

COLITIS AND PERIANAL DISEASE Patients with colitis present with low-grade fevers, malaise, diarrhea, crampy abdominal pain, and sometimes hematochezia. Gross bleeding is not as common as in UC and appears in about one-half of patients with exclusively colonic disease. Only 1–2% bleed massively. Pain is caused by passage of fecal material through narrowed and inflamed segments of the large bowel. Decreased rectal compliance is another cause for diarrhea in Crohn's colitis patients. Toxic megacolon is rare but may be seen with severe inflammation and short duration disease.

Strictureing can occur in the colon in 4–16% of patients and produce symptoms of bowel obstruction. If the endoscopist is unable to traverse a stricture in Crohn's colitis, surgical resection should be considered, especially if the patient has symptoms of chronic obstruction.

1954 Colonic disease may fistulize into the stomach or duodenum, causing feculent vomiting, or to the proximal or mid-small bowel, causing malabsorption by “short circuiting” and bacterial overgrowth. Ten percent of women with Crohn’s colitis will develop a rectovaginal fistula.

Perianal disease affects about one-third of patients with Crohn’s colitis and is manifested by incontinence, large hemorrhoidal tags, anal strictures, anorectal fistulae, and perirectal abscesses. Not all patients with perianal fistula will have endoscopic evidence of colonic inflammation.

GASTRODUODENAL DISEASE Symptoms and signs of upper GI tract disease include nausea, vomiting, and epigastric pain. Patients usually have an *Helicobacter pylori*-negative gastritis. The second portion of the duodenum is more commonly involved than the bulb. Fistulas involving the stomach or duodenum arise from the small or large bowel and do not necessarily signify the presence of upper GI tract involvement. Patients with advanced gastroduodenal CD may develop a chronic gastric outlet obstruction.

Laboratory, Endoscopic, and Radiographic Features Laboratory abnormalities include elevated ESR and CRP. In more severe disease, findings include hypoalbuminemia, anemia, and leukocytosis.

Endoscopic features of CD include rectal sparing, aphthous ulcerations, fistulas, and skip lesions. Colonoscopy allows examination and biopsy of mass lesions or strictures and biopsy of the terminal ileum. Upper endoscopy is useful in diagnosing gastroduodenal involvement in patients with upper tract symptoms. Ileal or colonic strictures may be dilated with balloons introduced through the colonoscope. Strictures ≤ 4 cm and those at anastomotic sites respond better to endoscopic dilation. The perforation rate is as high as 10%. Most endoscopists dilate only fibrotic strictures and not those associated with active inflammation. Wireless capsule endoscopy (WCE) allows direct visualization of the entire small-bowel mucosa (Fig. 351-8). The diagnostic yield of detecting lesions suggestive of active CD is higher with WCE than CT or magnetic resonance (MR) enterography or small-bowel series. WCE cannot be used in the setting of a small-bowel stricture. Capsule retention occurs in $<1\%$ of patients with suspected CD, but retention rates of 4–6% are seen in patients with

established CD. It is helpful to give the patient with CD a patency capsule, which is made of barium and starts to dissolve 30 h after ingestion. An abdominal x-ray can be taken at around 30 h after ingestion to see if the capsule is still present in the small bowel, which would indicate a stricture.

In CD, early radiographic findings in the small bowel include thickened folds and aphthous ulcerations. “Cobblestoning” from longitudinal and transverse ulcerations most frequently involves the small bowel. In more advanced disease, strictures, fistulas, inflammatory masses, and abscesses may be detected. The earliest macroscopic findings of colonic CD are aphthous ulcers. These small ulcers are often multiple and separated by normal intervening mucosa. As the disease progresses, aphthous ulcers become enlarged, deeper, and occasionally connected to one another, forming longitudinal stellate, serpiginous, and linear ulcers (see Fig. 345-4B).

The transmural inflammation of CD leads to decreased luminal diameter and limited distensibility. As ulcers progress deeper, they can lead to fistula formation. The radiographic “string sign” represents long areas of circumferential inflammation and fibrosis, resulting in long segments of luminal narrowing. The segmental nature of CD results in wide gaps of normal or dilated bowel between involved segments.

Both CT and MRI of the small bowel can be performed by enterography (CTE or MRE), using oral and IV contrast, as well as enteroclysis. Although institutional preference guides technique selection, CTE and MRE tend to be preferred over enteroclysis due to ease and patient preference. Although CTE, MRE, and small-bowel follow-through (SBFT) have been shown to be equally accurate in the identification of active small-bowel inflammation, CTE and MRE have been shown to be superior to SBFT in the detection of extraluminal complications, including fistulas, sinus tracts, and abscesses. Currently, the use of CT scans is more common than MRI due to institutional availability and expertise. However, MRI is thought to offer superior soft tissue contrast and has the added advantage of avoiding radiation exposure changes (Figs. 351-9 and 351-10). The lack of ionizing radiation is particularly appealing in younger patients and when monitoring response to therapy where serial images will be obtained. Either CTE or MRE is the first-line test for the evaluation of suspected CD and its complications. Pelvic MRI is superior to CT for demonstrating pelvic lesions such as ischioanal abscesses and perianal fistulae (Fig. 351-11).

Complications Because CD is a transmural process, serosal adhesions develop that provide direct pathways for fistula formation and reduce the incidence of free perforation. Perforation occurs in 1–2% of patients, usually in the ileum but occasionally in the jejunum or as a complication of toxic megacolon. The peritonitis of free perforation, especially colonic, may be fatal. Intraabdominal and pelvic abscesses occur in 10–30% of patients with CD at some time in the course of their illness. CT-guided percutaneous drainage of the abscess is standard therapy. Despite adequate drainage, most patients need resection of the offending bowel segment. Percutaneous drainage has an especially high failure rate in abdominal wall abscesses. Systemic glucocorticoid therapy increases the risk of intraabdominal and pelvic abscesses in CD patients who have never had an operation. Other complications include intestinal obstruction in 40%, massive hemorrhage, malabsorption, and severe perianal disease.

Serologic Markers Patients with CD show a wide variation in the way they present and progress over time. Some patients present with mild disease activity and do well with generally safe and mild medications, but many others exhibit more severe disease and can develop serious complications that will require surgery. Current and developing biologic therapies can help halt progression of disease and give patients with moderate to severe CD a better quality of life. There are potential risks of biologic therapies such as infection and malignancy, and it would be optimal to determine at the time of diagnosis which patients will require more aggressive medical therapy. This same argument holds true for UC patients as well.

Subsets of patients with differing immune responses to microbial antigens have been described, and serology is often tested for



FIGURE 351-8 Wireless capsule endoscopy image in a patient with Crohn’s disease of the ileum shows ulcerations and narrowing of the intestinal lumen. (Courtesy of Dr. S. Reddy, Gastroenterology Division, Department of Medicine, Brigham and Women’s Hospital, Boston, Massachusetts; with permission.)



FIGURE 351-9 A coronal magnetic resonance image was obtained using a half Fourier single-shot T2-weighted acquisition with fat saturation in a 27-year-old pregnant (23 weeks' gestation) woman. The patient had Crohn's disease and was maintained on 6-mercaptopurine and prednisone. She presented with abdominal pain, distension, vomiting, and small-bowel obstruction. The image reveals a 7- to 10-cm long stricture at the terminal ileum (white arrows) causing obstruction and significant dilatation of the proximal small bowel (white asterisk). A fetus is seen in the uterus (dashed white arrows). (Courtesy of Drs. J. F. B. Chick and P. B. Shyn, *Abdominal Imaging and Intervention*, Department of Radiology, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts; with permission.)

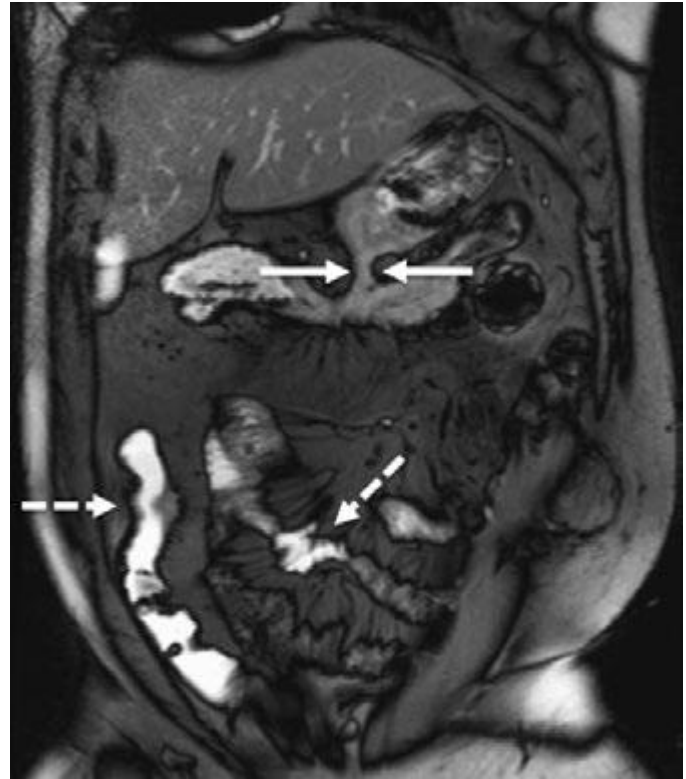


FIGURE 351-10 A coronal balanced, steady-state, free precession, T2-weighted image with fat saturation was obtained in a 32-year-old man with Crohn's disease and prior episodes of bowel obstruction, fistulas, and abscesses. He was being treated with 6-mercaptopurine and presented with abdominal distention and diarrhea. The image demonstrates a new gastrocolic fistula (solid white arrows). Multifocal involvement of the small bowel and terminal ileum is also present (dashed white arrows). (Courtesy of Drs. J. F. B. Chick and P. B. Shyn, *Abdominal Imaging and Intervention*, Department of Radiology, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts; with permission.)

perinuclear antineutrophil cytoplasmic antibodies (pANCA) and anti-*Saccharomyces cerevisiae* antibodies (ASCAs). Unfortunately, these serologic markers are only marginally useful in helping to make the diagnosis of UC or CD and in predicting the course of disease. For success in diagnosing IBD and in differentiating between CD and UC, the efficacy of these serologic tests depends on the prevalence of IBD in a specific population. pANCA positivity is found in about 60–70% of UC patients and 5–10% of CD patients; 5–15% of first-degree relatives of UC patients are pANCA positive, whereas only 2–3% of the general population is pANCA positive. Sixty to 70% of CD patients, 10–15% of UC patients, and up to 5% of non-IBD controls are ASCA positive. In a patient population with a combined prevalence of UC and CD of 62%, pANCA/ASCA serology showed a sensitivity of 64% and a specificity of 94%. Positive and negative predictive values (PPVs and NPVs) for pANCA/ASCA also vary based on the prevalence of IBD in a given population. For the patient population with a prevalence of IBD of 62%, the PPV is 94%, and the NPV is 63%.

Other serologic tests include antibodies to *Escherichia coli* outer membrane porin protein C (OmpC), which is found in 55% of CD patients; antibodies to I₂, a homologue of the bacterial transcription factor families from a *Pseudomonas fluorescens*-associated sequence that is found in 50–54% of CD patients; and anti-flagellin (anti-CBir1) antibodies, which have been identified in approximately 50% of CD patients.

Children with CD positive for all four immune responses (ASCA+, OmpC+, I₂+, and anti-Cbir1+) may have more aggressive disease and a shorter time to progression to internal perforating

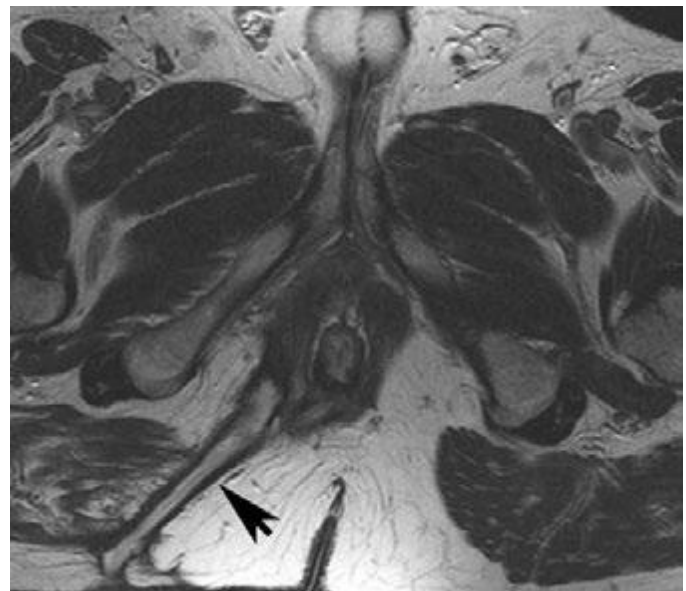


FIGURE 351-11 Axial T2-weighted magnetic resonance image obtained in a 37-year-old man with Crohn's disease shows a linear fluid-filled perianal fistula (arrow) in the right ischioanal fossa. (Courtesy of Dr. K. Morteale, *Gastrointestinal Radiology*, Department of Radiology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

1956 and/or stricturing disease. However, larger prospective studies in both children and adults have not yet been performed and compared to CRP or other markers.

Clinical factors described at diagnosis are more helpful than serologies at predicting the natural history of CD. The initial requirements for glucocorticoid use, an age at diagnosis below 40 years and the presence of perianal disease at diagnosis, have been shown to be independently associated with subsequent disabling CD after 5 years. Except in special circumstances (such as before consideration of an ileoanal pouch anastomosis [IPAA] in a patient with indeterminate colitis), serologic markers have only minimal clinical utility.

DIFFERENTIAL DIAGNOSIS OF UC AND CD

UC and CD have similar features to many other diseases. In the absence of a key diagnostic test, a combination of features is used (Table 351-5). Once a diagnosis of IBD is made, distinguishing between UC and CD is impossible initially in up to 15% of cases. These are termed *indeterminate colitis*. Fortunately, in most cases, the true nature of the underlying colitis becomes evident later in the course of the patient's disease. Approximately 5% (range 1–20%) of colon resection specimens are difficult to classify as either UC or CD because they exhibit overlapping histologic features.

INFECTIOUS DISEASES

Infections of the small intestines and colon can mimic CD or UC. They may be bacterial, fungal, viral, or protozoal in origin (Table 351-6). *Campylobacter* colitis can mimic the endoscopic appearance of severe UC and can cause a relapse of established UC. *Salmonella* can cause watery or bloody diarrhea, nausea, and vomiting. Shigellosis causes watery diarrhea, abdominal pain, and fever followed by rectal tenesmus and by the passage of blood and mucus per rectum. All three are usually self-limited, but 1% of patients infected with *Salmonella* become asymptomatic carriers. *Yersinia enterocolitica* infection occurs mainly in the terminal ileum and causes mucosal ulceration, neutrophil invasion, and thickening of the ileal wall. Other bacterial infections that may mimic IBD include *C. difficile*, which presents with watery

TABLE 351-6 DISEASES THAT MIMIC IBD

Infectious Etiologies		
Bacterial	Mycobacterial	Viral
<i>Salmonella</i>	Tuberculosis	Cytomegalovirus
<i>Shigella</i>	<i>Mycobacterium avium</i>	Herpes simplex
Toxicogenic		HIV
<i>Escherichia coli</i>	Parasitic	Fungal
<i>Campylobacter</i>	Amebiasis	Histoplasmosis
<i>Yersinia</i>	<i>Isospora</i>	<i>Candida</i>
<i>Clostridium difficile</i>	<i>Trichuris trichiura</i>	<i>Aspergillus</i>
Gonorrhea	Hookworm	
<i>Chlamydia trachomatis</i>	<i>Strongyloides</i>	
Noninfectious Etiologies		
Inflammatory	Neoplastic	Drugs and Chemicals
Appendicitis	Lymphoma	NSAIDs
Diverticulitis	Metastatic	Phosphosoda
Diversion colitis	Carcinoma	Cathartic colon
Collagenous/lymphocytic colitis	Carcinoma of the ileum	Gold
Ischemic colitis	Carcinoid	Oral contraceptives
Radiation colitis/enteritis	Familial polyposis	Cocaine
Solitary rectal ulcer syndrome		Ipilimumab
Eosinophilic gastroenteritis		Mycophenolate mofetil
Neutropenic colitis		
Behçet's syndrome		
Graft-versus-host disease		

Abbreviation: IBD, inflammatory bowel disease; NSAIDs, nonsteroidal anti-inflammatory drugs.

diarrhea, tenesmus, nausea, and vomiting; and *E. coli*, three categories of which can cause colitis. These are enterohemorrhagic, enteroinvasive, and enteroadherent *E. coli*, all of which can cause bloody diarrhea and abdominal tenderness. Diagnosis of bacterial colitis is made by sending stool specimens for bacterial culture and *C. difficile* toxin analysis. Gonorrhea, *Chlamydia*, and syphilis can also cause proctitis.

GI involvement with mycobacterial infection occurs primarily in the immunosuppressed patient but may occur in patients with normal immunity. Distal ileal and cecal involvement predominates, and patients present with symptoms of small-bowel obstruction and a tender abdominal mass. The diagnosis is made most directly by colonoscopy with biopsy and culture. *Mycobacterium avium-intracellulare* complex infection occurs in advanced stages of HIV infection and in other immunocompromised states; it usually manifests as a systemic infection with diarrhea, abdominal pain, weight loss, fever, and malabsorption. Diagnosis is established by acid-fast smear and culture of mucosal biopsies.

Although most of the patients with viral colitis are immunosuppressed, cytomegalovirus (CMV) and herpes simplex proctitis may occur in immunocompetent individuals. CMV occurs most commonly in the esophagus, colon, and rectum but may also involve the small intestine. Symptoms include abdominal pain, bloody diarrhea, fever, and weight loss. With severe disease, necrosis and perforation can occur. Diagnosis is made by identification of characteristic intranuclear inclusions in mucosal cells on biopsy. Herpes simplex infection of the GI tract is limited to the oropharynx, anorectum, and perianal areas. Symptoms include anorectal pain, tenesmus, constipation, inguinal adenopathy, difficulty with urinary voiding, and sacral paresthesias. Diagnosis is made by rectal biopsy with identification of characteristic cellular inclusions and viral culture. HIV itself can cause diarrhea, nausea, vomiting, and anorexia. Small intestinal biopsies show partial villous atrophy; small bowel bacterial overgrowth and fat malabsorption may also be noted.

TABLE 351-5 DIFFERENT CLINICAL, ENDOSCOPIC, AND RADIOGRAPHIC FEATURES

	Ulcerative Colitis	Crohn's Disease
Clinical		
Gross blood in stool	Yes	Occasionally
Mucus	Yes	Occasionally
Systemic symptoms	Occasionally	Frequently
Pain	Occasionally	Frequently
Abdominal mass	Rarely	Yes
Significant perineal disease	No	Frequently
Fistulas	No	Yes
Small intestinal obstruction	No	Frequently
Colonic obstruction	Rarely	Frequently
Response to antibiotics	No	Yes
Recurrence after surgery	No	Yes
Endoscopic		
Rectal sparing	Rarely	Frequently
Continuous disease	Yes	Occasionally
"Cobblestoning"	No	Yes
Granuloma on biopsy	No	Occasionally
Radiographic		
Small bowel significantly abnormal	No	Yes
Abnormal terminal ileum	No	Yes
Segmental colitis	No	Yes
Asymmetric colitis	No	Yes
Stricture	Occasionally	Frequently

Protozoan parasites include *Isospora belli*, which can cause a self-limited infection in healthy hosts but causes a chronic profuse, watery diarrhea, and weight loss in AIDS patients. *Entamoeba histolytica* or related species infect about 10% of the world's population; symptoms include abdominal pain, tenesmus, frequent loose stools containing blood and mucus, and abdominal tenderness. Colonoscopy reveals focal punctate ulcers with normal intervening mucosa; diagnosis is made by biopsy or serum amebic antibodies. Fulminant amebic colitis is rare but has a mortality rate of >50%.

Other parasitic infections that may mimic IBD include hookworm (*Necator americanus*), whipworm (*Trichuris trichiura*), and *Strongyloides stercoralis*. In severely immunocompromised patients, *Candida* or *Aspergillus* can be identified in the submucosa. Disseminated histoplasmosis can involve the ileocecal area.

NONINFECTIOUS DISEASES

Diverticulitis can be confused with CD clinically and radiographically. Both diseases cause fever, abdominal pain, tender abdominal mass, leukocytosis, elevated ESR, partial obstruction, and fistulas. Perianal disease or ileitis on small-bowel series favors the diagnosis of CD. Significant endoscopic mucosal abnormalities are more likely in CD than in diverticulitis. Endoscopic or clinical recurrence following segmental resection favors CD. Diverticular-associated colitis is similar to CD, but mucosal abnormalities are limited to the sigmoid and descending colon.

Ischemic colitis is commonly confused with IBD. The ischemic process can be chronic and diffuse, as in UC, or segmental, as in CD. Colonic inflammation due to ischemia may resolve quickly or may persist and result in transmural scarring and stricture formation. Ischemic bowel disease should be considered in the elderly following abdominal aortic aneurysm repair or when a patient has a hypercoagulable state or a severe cardiac or peripheral vascular disorder. Patients usually present with sudden onset of left lower quadrant pain, urgency to defecate, and the passage of bright red blood per rectum. Endoscopic examination often demonstrates a normal-appearing rectum and a sharp transition to an area of inflammation in the descending colon and splenic flexure.

The effects of radiotherapy on the GI tract can be difficult to distinguish from IBD. Acute symptoms can occur within 1–2 weeks of starting radiotherapy. When the rectum and sigmoid are irradiated, patients develop bloody, mucoid diarrhea and tenesmus, as in distal UC. With small-bowel involvement, diarrhea is common. Late symptoms include malabsorption and weight loss. Strictureing with obstruction and bacterial overgrowth may occur. Fistulas can penetrate the bladder, vagina, or abdominal wall. Flexible sigmoidoscopy reveals mucosal granularity, friability, numerous telangiectasias, and occasionally discrete ulcerations. Biopsy can be diagnostic.

Solitary rectal ulcer syndrome is uncommon and can be confused with IBD. It occurs in persons of all ages and may be caused by impaired evacuation and failure of relaxation of the puborectalis muscle. Single or multiple ulcerations may arise from anal sphincter overactivity, higher intrarectal pressures during defecation, and digital removal of stool. Patients complain of constipation with straining and pass blood and mucus per rectum. Other symptoms include abdominal pain, diarrhea, tenesmus, and perineal pain. Ulceration as large as 5 cm in diameter is usually seen anteriorly or anterior-laterally 3–15 cm from the anal verge. Biopsies can be diagnostic.

Several types of colitis are associated with nonsteroidal anti-inflammatory drugs (NSAIDs), including de novo colitis, reactivation of IBD, and proctitis caused by use of suppositories. Most patients with NSAID-related colitis present with diarrhea and abdominal pain, and complications include stricture, bleeding, obstruction, perforation, and fistulization. Withdrawal of these agents is crucial, and in cases of reactivated IBD, standard therapies are indicated.

There are complications of two drugs used in a hospital setting that mimic IBD. The first is ipilimumab, a drug that targets cytotoxic T lymphocyte antigen 4 (CTLA-4) and reverses T cell inhibition and is used to treat metastatic melanoma; ipilimumab has an incidence of IBD in 0.0017 cases per 100 person-years. Ipilimumab-induced

colitis is typically treated with glucocorticoids or infliximab. The second is mycophenolate mofetil (MMF), an immunosuppressive agent commonly used to prevent posttransplant rejection. The colitis associated with MMF is common and can occur in more than one-third of patients taking the drug. Treatment is dose reduction or cessation of the drug.

THE ATYPICAL COLITIDES

Two atypical colitides—collagenous colitis and lymphocytic colitis—have completely normal endoscopic appearances. Collagenous colitis has two main histologic components: increased subepithelial collagen deposition and colitis with increased intraepithelial lymphocytes. The female to male ratio is 9:1, and most patients present in the sixth or seventh decades of life. The main symptom is chronic watery diarrhea. Treatments range from sulfasalazine or mesalamine and diphenoxylate/atropine (Lomotil) to bismuth to budesonide to prednisone or azathioprine/6-mercaptopurine for refractory disease. Risk factors include smoking; use of NSAIDs, proton pump inhibitors, or beta blockers; and a history of autoimmune disease.

Lymphocytic colitis has features similar to collagenous colitis, including age at onset and clinical presentation, but it has almost equal incidence in men and women and no subepithelial collagen deposition on pathologic section. However, intraepithelial lymphocytes are increased. Use of sertraline (but not beta blockers) is an additional risk factor. The frequency of celiac disease is increased in lymphocytic colitis and ranges from 9 to 27%. Celiac disease should be excluded in all patients with lymphocytic colitis, particularly if diarrhea does not respond to conventional therapy. Treatment is similar to that of collagenous colitis with the exception of a gluten-free diet for those who have celiac disease.

Diversion colitis is an inflammatory process that arises in segments of the large intestine that are excluded from the fecal stream. It usually occurs in patients with ileostomy or colostomy when a mucus fistula or a Hartmann's pouch has been created. Clinically, patients have mucus or bloody discharge from the rectum. Erythema, granularity, friability, and, in more severe cases, ulceration can be seen on endoscopy. Histopathology shows areas of active inflammation with foci of cryptitis and crypt abscesses. Crypt architecture is normal, which differentiates it from UC. It may be impossible to distinguish from CD. Short-chain fatty acid enemas may help in diversion colitis, but the definitive therapy is surgical reanastomosis.

EXTRAIESTINAL MANIFESTATIONS

Up to one-third of IBD patients have at least one extraintestinal disease manifestation.

DERMATOLOGIC

Erythema nodosum (EN) occurs in up to 15% of CD patients and 10% of UC patients. Attacks usually correlate with bowel activity; skin lesions develop after the onset of bowel symptoms, and patients frequently have concomitant active peripheral arthritis. The lesions of EN are hot, red, tender nodules measuring 1–5 cm in diameter and are found on the anterior surface of the lower legs, ankles, calves, thighs, and arms. Therapy is directed toward the underlying bowel disease.

Pyoderma gangrenosum (PG) is seen in 1–12% of UC patients and less commonly in Crohn's colitis. Although it usually presents after the diagnosis of IBD, PG may occur years before the onset of bowel symptoms, run a course independent of the bowel disease, respond poorly to colectomy, and even develop years after proctocolectomy. It is usually associated with severe disease. Lesions are commonly found on the dorsal surface of the feet and legs but may occur on the arms, chest, stoma, and even the face. PG usually begins as a pustule and then spreads concentrically to rapidly undermine healthy skin. Lesions then ulcerate, with violaceous edges surrounded by a margin of erythema. Centrally, they contain necrotic tissue with blood and exudates. Lesions may be single or multiple and grow as large as 30 cm. They are sometimes very difficult to treat and often require IV antibiotics, IV glucocorticoids, dapsone, azathioprine, thalidomide, IV cyclosporine, or infliximab.

Other dermatologic manifestations include pyoderma vegetans, which occurs in intertriginous areas; pyostomatitis vegetans, which involves the mucous membranes; Sweet syndrome, a neutrophilic dermatosis; and metastatic CD, a rare disorder defined by cutaneous granuloma formation. Psoriasis affects 5–10% of patients with IBD and is unrelated to bowel activity consistent with the potential shared immunogenetic basis of these diseases. Perianal skin tags are found in 75–80% of patients with CD, especially those with colon involvement. Oral mucosal lesions, seen often in CD and rarely in UC, include aphthous stomatitis and “cobblestone” lesions of the buccal mucosa.

RHEUMATOLOGIC

Peripheral arthritis develops in 15–20% of IBD patients, is more common in CD, and worsens with exacerbations of bowel activity. It is asymmetric, polyarticular, and migratory and most often affects large joints of the upper and lower extremities. Treatment is directed at reducing bowel inflammation. In severe UC, colectomy frequently cures the arthritis.

Ankylosing spondylitis (AS) occurs in about 10% of IBD patients and is more common in CD than UC. About two-thirds of IBD patients with AS express the HLA-B27 antigen. The AS activity is not related to bowel activity and does not remit with glucocorticoids or colectomy. It most often affects the spine and pelvis, producing symptoms of diffuse low-back pain, buttock pain, and morning stiffness. The course is continuous and progressive, leading to permanent skeletal damage and deformity. Anti-TNF therapy reduces spinal inflammation and improves functional status and quality of life.

Sacroiliitis is symmetric, occurs equally in UC and CD, is often asymptomatic, does not correlate with bowel activity, and does not always progress to AS. Other rheumatic manifestations include hypertrophic osteoarthropathy, pelvic/femoral osteomyelitis, and relapsing polycondritis.

OCULAR

The incidence of ocular complications in IBD patients is 1–10%. The most common are conjunctivitis, anterior uveitis/iritis, and episcleritis. Uveitis is associated with both UC and Crohn’s colitis, may be found during periods of remission, and may develop in patients following bowel resection. Symptoms include ocular pain, photophobia, blurred vision, and headache. Prompt intervention, sometimes with systemic glucocorticoids, is required to prevent scarring and visual impairment. Episcleritis is a benign disorder that presents with symptoms of mild ocular burning. It occurs in 3–4% of IBD patients, more commonly in Crohn’s colitis, and is treated with topical glucocorticoids.

HEPATOBIILIARY

Hepatic steatosis is detectable in about one-half of the abnormal liver biopsies from patients with CD and UC; patients usually present with hepatomegaly. Fatty liver usually results from a combination of chronic debilitating illness, malnutrition, and glucocorticoid therapy. Cholelithiasis occurs in 10–35% of CD patients with ileitis or ileal resection. Gallstone formation is caused by malabsorption of bile acids, resulting in depletion of the bile salt pool and the secretion of lithogenic bile.

Primary sclerosing cholangitis (PSC) is a disorder characterized by both intrahepatic and extrahepatic bile duct inflammation and fibrosis, frequently leading to biliary cirrhosis and hepatic failure; approximately 5% of patients with UC have PSC, but 50–75% of patients with PSC have IBD. PSC occurs less often in patients with CD. Although it can be recognized after the diagnosis of IBD, PSC can be detected earlier or even years after proctocolectomy. Consistent with this, the immunogenetic basis for PSC appears to be overlapping but distinct from UC based on GWAS, although both IBD and PSC are commonly pANCA positive. Most patients have no symptoms at the time of diagnosis; when symptoms are present, they consist of fatigue, jaundice, abdominal pain, fever, anorexia, and malaise. The traditional gold standard diagnostic test is endoscopic retrograde cholangiopancreatography (ERCP), but magnetic resonance cholangiopancreatography (MRCP) is also sensitive and specific. MRCP is reasonable as an initial diagnostic test in children and can visualize irregularities, multifocal

strictures, and dilatations of all levels of the biliary tree. In patients with PSC, both ERCP and MRCP demonstrate multiple bile duct strictures alternating with relatively normal segments.

The bile acid ursodeoxycholic acid (ursodiol) may reduce alkaline phosphatase and serum aminotransferase levels, but histologic improvement has been marginal. High doses (25–30 mg/kg per day) may decrease the risk of colorectal dysplasia and cancer in patients with UC and PSC. Endoscopic stenting may be palliative for cholestasis secondary to bile duct obstruction. Patients with symptomatic disease develop cirrhosis and liver failure over 5–10 years and eventually require liver transplantation. PSC patients have a 10–15% lifetime risk of developing cholangiocarcinoma and then cannot be transplanted. Patients with IBD and PSC are at increased risk of colon cancer and should be surveyed yearly by colonoscopy and biopsy.

In addition, cholangiography is normal in a small percentage of patients who have a variant of PSC known as *small duct primary sclerosing cholangitis*. This variant (sometimes referred to as “pericholangitis”) is probably a form of PSC involving small-caliber bile ducts. It has similar biochemical and histologic features to classic PSC. It appears to have a significantly better prognosis than classic PSC, although it may evolve into classic PSC. Granulomatous hepatitis and hepatic amyloidosis are much rarer extraintestinal manifestations of IBD.

UROLOGIC

The most frequent genitourinary complications are calculi, ureteral obstruction, and ileal bladder fistulas. The highest frequency of nephrolithiasis (10–20%) occurs in patients with CD following small bowel resection. Calcium oxalate stones develop secondary to hyperoxaluria, which results from increased absorption of dietary oxalate. Normally, dietary calcium combines with luminal oxalate to form insoluble calcium oxalate, which is eliminated in the stool. In patients with ileal dysfunction, however, nonabsorbed fatty acids bind calcium and leave oxalate unbound. The unbound oxalate is then delivered to the colon, where it is readily absorbed, especially in the presence of inflammation.

METABOLIC BONE DISORDERS

Low bone mass occurs in 3–30% of IBD patients. The risk is increased by glucocorticoids, cyclosporine, methotrexate, and total parenteral nutrition (TPN). Malabsorption and inflammation mediated by IL-1, IL-6, TNF, and other inflammatory mediators also contribute to low bone density. An increased incidence of hip, spine, wrist, and rib fractures has been noted: 36% in CD and 45% in UC. The absolute risk of an osteoporotic fracture is about 1% per person per year. Fracture rates, particularly in the spine and hip, are highest among the elderly (age >60). One study noted an OR of 1.72 for vertebral fracture and an OR of 1.59 for hip fracture. The disease severity predicted the risk of a fracture. Only 13% of IBD patients who had a fracture were on any kind of antifracture treatment. Up to 20% of bone mass can be lost per year with chronic glucocorticoid use. The effect is dosage-dependent. Budesonide may also suppress the pituitary-adrenal axis and thus carries a risk of causing osteoporosis.

Osteonecrosis is characterized by death of osteocytes and adipocytes and eventual bone collapse. The pain is aggravated by motion and swelling of the joints. It affects the hips more often than knees and shoulders, and in one series, 4.3% of patients developed osteonecrosis within 6 months of starting glucocorticoids. Diagnosis is made by bone scan or MRI, and treatment consists of pain control, cord decompression, osteotomy, and joint replacement.

THROMBOEMBOLIC DISORDERS

Patients with IBD have an increased risk of both venous and arterial thrombosis even if the disease is not active. Factors responsible for the hypercoagulable state have included abnormalities of the platelet-endothelial interaction, hyperhomocysteinemia, alterations in the coagulation cascade, impaired fibrinolysis, involvement of tissue factor-bearing microvesicles, disruption of the normal coagulation system by autoantibodies, and a genetic predisposition. A spectrum of vasculitides involving small, medium, and large vessels has also been observed.

OTHER DISORDERS

More common cardiopulmonary manifestations include endocarditis, myocarditis, pleuropericarditis, and interstitial lung disease. A secondary or reactive amyloidosis can occur in patients with long-standing IBD, especially in patients with CD. Amyloid material is deposited systemically and can cause diarrhea, constipation, and renal failure. The renal disease can be successfully treated with colchicine. Pancreatitis is a rare extraintestinal manifestation of IBD and results from duodenal fistulas; ampullary CD; gallstones; PSC; drugs such as 6-mercaptopurine, azathioprine, or, very rarely, 5-ASA agents; autoimmune pancreatitis; and primary CD of the pancreas.

TREATMENT INFLAMMATORY BOWEL DISEASE

TREATMENT

5-ASA AGENTS

The mainstay of therapy for mild to moderate UC is sulfasalazine and the other 5-ASA agents. These agents are effective at inducing and maintaining remission in UC. They may have a limited role in inducing remission in CD but no clear role in maintenance of CD. Newer sulfa-free aminosalicylate preparations deliver increased amounts of the pharmacologically active ingredient of sulfasalazine (5-ASA, mesalamine) to the site of active bowel disease while limiting systemic toxicity. Peroxisome proliferator activated receptor γ (PPAR- γ) may mediate 5-ASA therapeutic action by decreasing nuclear localization of NF- κ B. Sulfa-free aminosalicylate formulations include alternative azo-bonded carriers, 5-ASA dimers, and delayed-release and controlled-release preparations. Each has the same efficacy as sulfasalazine when equimolar concentrations are used.

Sulfasalazine was originally developed to deliver both antibacterial (sulfapyridine) and anti-inflammatory (5-ASA) therapy into the connective tissues of joints and the colonic mucosa. The molecular structure provides a convenient delivery system to the colon by allowing the intact molecule to pass through the small intestine after only partial absorption and to be broken down in the colon by bacterial azo reductases that cleave the azo bond linking the sulfa and 5-ASA moieties. Sulfasalazine is effective treatment for mild to moderate UC and is occasionally used in Crohn's colitis, but its high rate of side effects limits its use. Although sulfasalazine is more effective at higher doses, at 6 or 8 g/d up to 30% of patients experience allergic reactions or intolerable side effects such as headache, anorexia, nausea, and vomiting that are attributable to the sulfapyridine moiety. Hypersensitivity reactions, independent of sulfapyridine levels, include rash, fever, hepatitis, agranulocytosis, hypersensitivity pneumonitis, pancreatitis, worsening of colitis, and reversible sperm abnormalities. Sulfasalazine can also impair folate absorption, and patients should be given folic acid supplements.

Balsalazide contains an azo bond binding mesalamine to the carrier molecule 4-aminobenzoyl- β -alanine; it is effective in the colon.

Olsalazine is composed of two 5-ASA radicals linked by an azo bond, which is split in the colon by bacterial reduction, and two 5-ASA molecules are released. Olsalazine is similar in effectiveness to sulfasalazine in treating UC, but up to 17% of patients experience nonbloody diarrhea caused by increased secretion of fluid in the small bowel.

Delzicol and *Asacol HD* (high dose) are enteric-coated forms of mesalamine with the 5-ASA being released at pH >7. They disintegrate with complete breakup of the tablet occurring in many different parts of the gut ranging from the small intestine to the splenic flexure; they have increased gastric residence when taken with a meal. *Asacol* has recently been discontinued and replaced with *Delzicol*, which lacks dibutyl phthalate (DBP), an inactive ingredient in *Asacol*'s enteric coating. DBP has been associated with adverse effects on the male reproductive system in animals at very high doses. *Asacol HD* with the same chemical in its coating is still on the market, but the human doses of DBP are within acceptable limits of toxicity.

Lialda is a once-a-day formulation of mesalamine (Multi-Matrix System [MMX]) designed to release mesalamine in the colon. The MMX technology incorporates mesalamine into a lipophilic matrix within a hydrophilic matrix encapsulated in a polymer resistant to degradation at a low pH (<7) to delay release throughout the colon. The safety profile appears to be comparable to other 5-ASA formulations.

Apriso is a formulation containing encapsulated mesalamine granules that delivers mesalamine to the terminal ileum and colon via a proprietary extended-release mechanism (Intellicor). The outer coating (Eudragit L) dissolves at a pH >6. In addition, there is a polymer matrix core that aids in sustained release throughout the colon. Because *Lialda* and *Apriso* are given once daily, an anticipated benefit is improved compliance compared with two to four daily doses required for other mesalamine preparations.

Pentasa is another mesalamine formulation that uses an ethylcellulose coating to allow water absorption into small beads containing the mesalamine. Water dissolves the 5-ASA, which then diffuses out of the bead into the lumen. Disintegration of the capsule occurs in the stomach. The microspheres then disperse throughout the entire GI tract from the small intestine through the distal colon in both fasted and fed conditions.

Salofalk® Granu-Stix, an unencapsulated version of mesalamine, has been in use in Europe for induction and maintenance of remission for several years.

Appropriate doses of the 5-ASA compounds are shown in **Table 351-7**. Some 50–75% of patients with mild to moderate UC improve when treated with 5-ASA doses equivalent to 2 g/d of

TABLE 351-7 ORAL 5-ASA PREPARATIONS

Preparation	Formulation	Delivery	Dosing Per Day
Azo-Bond			
Sulfasalazine (500 mg) (Azulfidine)	Sulfapyridine-5-ASA	Colon	3–6 g (acute) 2–4 g (maintenance)
Olsalazine (250 mg) (Dipentum)	5-ASA–5-ASA	Colon	1–3 g
Balsalazide (750 mg) (Colazal)	Aminobenzoyl-alanine–5-ASA	Colon	6.75–9 g
Delayed-Release			
Mesalamine (400, 800 mg) (Delzicol, Asacol HD)	Eudragit S (pH 7)	Distal ileum-colon	2.4–4.8 g (acute) 1.6–4.8 g (maintenance)
Mesalamine (1.2 g) (Lialda)	MMX mesalamine (SPD476)	Ileum-colon	2.4–4.8 g
Controlled-Release			
Mesalamine (250, 500, 1000 mg) (Pentasa)	Ethylcellulose microgranules	Stomach-colon	2–4 g (acute) 1.5–4 g (maintenance)
Delayed- and Extended-Release			
Mesalamine (0.375 g) (Apriso)	Intellicor extended-release mechanism	Ileum-colon	1.5 g (maintenance)

1960 mesalamine; the dose response continues up to at least 4.8 g/d. As a general rule, 5-ASA agents act within 2–4 weeks. 5-ASA doses equivalent to 1.5–4 g/d of mesalamine maintain remission in 50–75% of patients with UC.

More common side effects of the 5-ASA medications include headaches, nausea, hair loss, and abdominal pain. Rare side effects of the 5-ASA medications include renal impairment, hematuria, pancreatitis, and paradoxical worsening of colitis. Renal function tests and urinalysis should be checked yearly.

Topical *Rowasa* enemas are composed of mesalamine and are effective in mild-to-moderate distal UC. Clinical response occurs in up to 80% of UC patients with colitis distal to the splenic flexure. Combination therapy with mesalamine in both oral and enema form is more effective than either treatment alone for both distal and extensive UC.

Canasa suppositories composed of mesalamine are effective in treating proctitis.

GLUCOCORTICOIDS

The majority of patients with moderate to severe UC benefit from oral or parenteral glucocorticoids. Prednisone is usually started at doses of 40–60 mg/d for active UC that is unresponsive to 5-ASA therapy. Parenteral glucocorticoids may be administered as hydrocortisone, 300 mg/d, or methylprednisolone, 40–60 mg/d. A new glucocorticoid for UC, budesonide (Uceris), is released entirely in the colon and has minimal to no glucocorticoid side effects. The dose is 9 mg/d for 8 weeks, and no taper is required. Topically applied glucocorticoids are also beneficial for distal colitis and may serve as an adjunct in those who have rectal involvement plus more proximal disease. Hydrocortisone enemas or foam may control active disease, although they have no proven role as maintenance therapy. These glucocorticoids are significantly absorbed from the rectum and can lead to adrenal suppression with prolonged administration. Topical 5-ASA therapy is more effective than topical steroid therapy in the treatment of distal UC.

Glucocorticoids are also effective for treatment of moderate to severe CD and induce a 60–70% remission rate compared to a 30% placebo response. The systemic effects of standard glucocorticoid formulations have led to the development of more potent formulations that are less well-absorbed and have increased first-pass metabolism. Controlled ileal-release budesonide has been nearly equal to prednisone for ileocolonic CD with fewer glucocorticoid side effects. Budesonide is used for 2–3 months at a dose of 9 mg/d, and then tapered. Budesonide 6 mg/d is effective in reducing relapse rates at 3–6 months but not at 12 months in CD patients with a medically induced remission.

Glucocorticoids play no role in maintenance therapy in either UC or CD. Once clinical remission has been induced, they should be tapered according to the clinical activity, normally at a rate of no more than 5 mg/week. They can usually be tapered to 20 mg/d within 4–5 weeks but often take several months to be discontinued altogether. The side effects are numerous, including fluid retention, abdominal striae, fat redistribution, hyperglycemia, subcapsular cataracts, osteonecrosis, osteoporosis, myopathy, emotional disturbances, and withdrawal symptoms. Most of these side effects, aside from osteonecrosis, are related to the dose and duration of therapy.

ANTIBIOTICS

Antibiotics have no role in the treatment of active or quiescent UC. However, pouchitis, which occurs in about a third of UC patients after colectomy and IPAA, usually responds to treatment with metronidazole and/or ciprofloxacin.

Metronidazole is effective in active inflammatory, fistulous, and perianal CD and may prevent recurrence after ileal resection. The most effective dose is 15–20 mg/kg per day in three divided doses; it is usually continued for several months. Common side effects include nausea, metallic taste, and disulfiram-like reaction. Peripheral neuropathy can occur with prolonged administration (several months) and on rare occasions is permanent despite discontinuation. *Ciprofloxacin* (500 mg bid) is also beneficial for

inflammatory, perianal, and fistulous CD but has been associated with Achilles tendinitis and rupture. Both ciprofloxacin and metronidazole antibiotics can be used as first-line drugs for short periods of time in active inflammatory, fistulizing, and perianal CD.

AZATHIOPRINE AND 6-MERCAPTOPYRINE

Azathioprine and 6-mercaptopurine (6-MP) are purine analogues commonly employed in the management of glucocorticoid-dependent IBD. Azathioprine is rapidly absorbed and converted to 6-MP, which is then metabolized to the active end product, thioguanine, an inhibitor of purine ribonucleotide synthesis and cell proliferation. These agents also inhibit the immune response. Efficacy can be seen as early as 3–4 weeks but can take up to 4–6 months. Adherence can be monitored by measuring the levels of 6-thioguanine and 6-methylmercaptopurine, end products of 6-MP metabolism. Azathioprine (2–3 mg/kg per day) and 6-MP (1–1.5 mg/kg per day) have been used successfully as glucocorticoid-sparing agents in up to two-thirds of UC and CD patients previously unable to be weaned from glucocorticoids. They are also used as maintenance therapy in UC and CD and for treating active perianal disease and fistulas in CD. In addition, 6-MP or azathioprine is effective for postoperative prophylaxis of CD.

Although azathioprine and 6-MP are usually well tolerated, pancreatitis occurs in 3–4% of patients, typically presents within the first few weeks of therapy, and is completely reversible when the drug is stopped. Other side effects include nausea, fever, rash, and hepatitis. Bone marrow suppression (particularly leukopenia) is dose-related and often delayed, necessitating regular monitoring of the complete blood cell count (CBC). Additionally, 1 in 300 individuals lacks thiopurine methyltransferase, the enzyme responsible for drug metabolism to inactive end-products (6-methylmercaptopurine); an additional 11% of the population are heterozygotes with intermediate enzyme activity. Both are at increased risk of toxicity because of increased accumulation of active 6-thioguanine metabolites. Although 6-thioguanine and 6-methylmercaptopurine levels can be followed to determine correct drug dosing and reduce toxicity, weight-based dosing is an acceptable alternative. CBCs and liver function tests should be monitored frequently regardless of dosing strategy. IBD patients treated with azathioprine/6-MP are at approximately a fourfold increased risk of developing a lymphoma. This increased risk could be a result of the medications, the underlying disease, or both.

METHOTREXATE

Methotrexate (MTX) inhibits dihydrofolate reductase, resulting in impaired DNA synthesis. Additional anti-inflammatory properties may be related to decreased IL-1 production. Intramuscular (IM) or subcutaneous (SC) MTX (25 mg/week) is effective in inducing remission and reducing glucocorticoid dosage; 15 mg/week is effective in maintaining remission in active CD. Potential toxicities include leukopenia and hepatic fibrosis, necessitating periodic evaluation of CBCs and liver enzymes. The role of liver biopsy in patients on long-term MTX is uncertain but is probably limited to those with increased liver enzymes. Hypersensitivity pneumonitis is a rare but serious complication of therapy.

CYCLOSPORINE

Cyclosporine (CSA) is a lipophilic peptide with inhibitory effects on both the cellular and humoral immune systems. CSA blocks the production of IL-2 by T helper lymphocytes. CSA binds to cyclophilin, and this complex inhibits calcineurin, a cytoplasmic phosphatase enzyme involved in the activation of T cells. CSA also indirectly inhibits B cell function by blocking helper T cells. CSA has a more rapid onset of action than 6-MP and azathioprine.

CSA is most effective when given at 2–4 mg/kg per day IV in severe UC that is refractory to IV glucocorticoids, with 82% of patients responding. CSA can be an alternative to colectomy. The long-term success of oral CSA is not as dramatic, but if patients are started on 6-MP or azathioprine at the time of hospital discharge, remission can be maintained. For the 2 mg/kg dose, levels as

measured by monoclonal radioimmunoassay or by the high-performance liquid chromatography assay should be maintained between 150 and 350 ng/mL.

CSA may cause significant toxicity; renal function should be monitored frequently. Hypertension, gingival hyperplasia, hypertrichosis, paresthesias, tremors, headaches, and electrolyte abnormalities are common side effects. Creatinine elevation calls for dose reduction or discontinuation. Seizures may also complicate therapy, especially if the patient is hypomagnesemic or if serum cholesterol levels are <3.1 mmol/L (<120 mg/dL). Opportunistic infections, most notably *Pneumocystis carinii* pneumonia, may occur with combination immunosuppressive treatment; prophylaxis should be given. Major adverse events occurred in 15% of patients in one large study, including nephrotoxicity not responding to dose adjustment, serious infections, seizures, anaphylaxis, and death of two patients. This high incidence suggests that vigorous monitoring by experienced clinicians at tertiary care centers may be required. To compare IV cyclosporine versus infliximab, a large trial was conducted in Europe by the GETAID group. The results indicated identical 7-day response rates between cyclosporine 2 mg/kg (with doses adjusted for levels of 150–250 ng/mL) and infliximab 5 mg/kg, with both groups achieving response rates of 85%. Serious infections occurred in 5 of 55 cyclosporine patients and 4 of 56 infliximab patients. Response rates were similar in the two groups at day 98 among patients treated with oral cyclosporine versus infliximab at the usual induction dose and maintenance dose regimen (40% and 46%, respectively). In light of data showing equal efficacy of CSA and infliximab in severe UC, more physicians are relying on infliximab rather than CSA in these patients.

TACROLIMUS

Tacrolimus is a macrolide antibiotic with immunomodulatory properties similar to CSA. It is 100 times as potent as CSA and is not dependent on bile or mucosal integrity for absorption. These pharmacologic properties enable tacrolimus to have good oral absorption despite proximal small bowel Crohn's involvement. It has shown efficacy in children with refractory IBD and in adults with extensive involvement of the small bowel. It is also effective in adults with glucocorticoid-dependent or refractory UC and CD as well as refractory fistulizing CD.

BIOLOGIC THERAPIES

Biologic therapy was traditionally reserved for moderately to severely ill patients with CD who had failed other therapies. However, it is now commonly given as an initial therapy for patients with moderate to severe CD in order to prevent future disease complications. Patients who respond to biologic therapies enjoy an improvement in clinical symptoms; a better quality of life; less disability, fatigue, and depression; and fewer surgeries and hospitalizations.

Anti-TNF Therapies The first biologic therapy approved for CD was *infliximab*, a chimeric IgG1 antibody against TNF- α , which is now also approved for treatment of moderately to severely active UC. Of active CD patients refractory to glucocorticoids, 6-MP, or 5-ASA, 65% will respond to IV infliximab (5 mg/kg); one-third will enter complete remission. The ACCENT I (A Crohn's Disease Clinical Trial Evaluating Infliximab in a New Long-Term Treatment Regimen) study showed that of the patients who experience an initial response, 40% will maintain remission for at least 1 year with repeated infusions of infliximab every 8 weeks.

Infliximab is also effective in CD patients with refractory perianal and enterocutaneous fistulas, with the ACCENT II trial showing a 68% response rate (50% reduction in fistula drainage) and a 50% complete remission rate. Reinfusion, typically every 8 weeks, is necessary to continue therapeutic benefits in many patients.

The SONIC (Study of Biologic and Immunomodulator-Naive Patients with Crohn's Disease) trial compared infliximab plus azathioprine, infliximab alone, and azathioprine alone in immunomodulator- and biologic-naive patients with moderate to severe CD. At 1 year, the infliximab plus azathioprine group had a glucocorticoid-free

remission rate of 46% compared with 35% for infliximab alone and 24% for azathioprine alone. There was also complete mucosal healing at week 26 with the combined approach relative to either infliximab or azathioprine alone (44% vs 30% vs 17%). The adverse events were equal between groups.

Two large trials of infliximab in moderate to severe UC also showed efficacy with a response rate of 37–49%, with about one-fifth of patients maintaining remission after 54 weeks. Dosing for UC and CD are identical, with induction dosing at 0, 2, and 6 weeks and every 8 weeks thereafter. There is a similar study to SONIC in patients with moderate to severe UC. After 16 weeks of therapy, UC patients taking azathioprine plus infliximab had a glucocorticoid-free remission rate of 40% compared to 24% (article now published) and 22% of those on azathioprine and infliximab alone, respectively. This is even further evidence for “top-down” or more aggressive therapy for both moderate to severe CD and UC.

Adalimumab is a recombinant human monoclonal IgG1 antibody containing only human peptide sequences and is injected subcutaneously. Adalimumab binds TNF and neutralizes its function by blocking the interaction between TNF and its cell-surface receptor. Therefore, it seems to have a similar mechanism of action to infliximab but with less immunogenicity. Adalimumab has been approved for treatment of moderate to severe CD. CHARM (Crohn's Trial of the Fully Human Adalimumab for Remission Maintenance) is an adalimumab maintenance study in patients who responded to adalimumab induction therapy. About 50% of the patients in this trial were previously treated with infliximab. Remission rates ranged from 42–48% of infliximab-naïve patients at 1 year compared with remission rates of 31–34% in patients who had previously received infliximab. Another trial showed a remission rate of 21% at 4 weeks in patients who had initially responded to and then failed infliximab. In clinical practice, the remission rate in patients taking adalimumab increases with a dose increase to 40 mg weekly instead of every other week. Adalimumab is now also approved for the treatment of moderately to severely active UC.

Certolizumab pegol is a pegylated form of an anti-TNF Fab portion of an antibody administered SC once monthly. SC certolizumab pegol was effective for induction of clinical response in patients with active inflammatory CD. In the PRECISE II (Pegylated Antibody Fragment Evaluation in Crohn's Disease) trial of maintenance therapy with certolizumab in patients who responded to certolizumab induction, the results were similar to the CHARM trial. At week 26, the subgroup of patients who were infliximab naïve had a response of 69% as compared to 44% in patients who had previously received infliximab.

Golimumab is another fully human IgG1 antibody against TNF- α and is currently approved for the treatment of moderately to severely active UC. All of the patients in the golimumab trial were infliximab-naïve. Like adalimumab and certolizumab, golimumab is injected SC.

Side Effects of Anti-TNF Therapies • DEVELOPMENT OF ANTIBODIES The development of antibodies to infliximab (ATIs) is associated with an increased risk of infusion reactions and a decreased response to treatment. Current practice does not include giving on-demand or episodic infusions in contrast to periodic (every 8 week) infusions because patients are most likely to develop ATIs. ATIs are generally present when the quality of response or the response duration to infliximab infusion decreases. Decreasing the dosing intervals or increasing the dosage to 10 mg/kg may restore the efficacy. There are commercial assays for both infliximab and adalimumab antibodies and trough levels to determine optimal dosing. If a patient has high ATIs and a low trough level of infliximab, it is best to switch to another anti-TNF therapy. Most acute infusion reactions and serum sickness can be managed with glucocorticoids and antihistamines. Some reactions can be serious and would necessitate a change in therapy, especially if a patient has ATIs.

NON-HODGKIN'S LYMPHOMA (NHL) The baseline risk of NHL in CD patients is 2:10,000, which is slightly higher than in the general population.

1962 Azathioprine and/or 6-MP therapy increases the risk to about 4:10,000. The highest risk for thiopurine-associated NHL is in patients over 65 years old, with a moderate risk in those between the ages of 50 and 65. Anti-TNF therapy increases the risk to approximately 6:10,000.

HEPATOSPLENIC T CELL LYMPHOMA (HSTCL) HSTCL is a nearly universally fatal lymphoma in patients with or without CD. In patients with CD, events reported to the Food and Drug Administration Adverse Event Reporting System (FDA AERS) and search of PubMed and Embase published case reports demonstrate a total of 37 unique cases. Eighty-six percent of the patients were male, with a median age of 26 years. Patients had CD for a mean of 10 years before the diagnosis of HSTCL. Thirty-six cases had used either 6-MP or azathioprine, and 28 cases had used infliximab. Of these 28 cases, 27 had also used 6-MP or azathioprine. The other case had a history of both infliximab and adalimumab exposure.

SKIN LESIONS New-onset psoriasiform skin lesions develop in nearly 5% of IBD patients treated with anti-TNF therapy. Most often, these can be treated topically, and rarely, anti-TNF therapy must be decreased, switched, or stopped. The risk of melanoma is increased almost twofold with anti-TNF and not thiopurine use. The risk of nonmelanoma skin cancer is increased with thiopurines and biologics, especially with 1 year of follow-up or greater. Patients on these medications should have a skin check at least once a year.

INFECTIONS All of the anti-TNF drugs are associated with an increased risk of infections, particularly reactivation of latent tuberculosis and opportunistic fungal infections including disseminated histoplasmosis and coccidioidomycosis. It is recommended that patients have a purified protein derivative (PPD) or a QuantiFERON-TB gold test as well as a chest x-ray before initiation of anti-TNF therapy. Patients over 65 have a higher rate of infections and death on infliximab or adalimumab than those younger than 65 years of age.

OTHER Acute liver injury due to reactivation of hepatitis B virus and to autoimmune effects and cholestasis has been reported. Rarely, infliximab and the other anti-TNF drugs have been associated with optic neuritis, seizures, new onset or exacerbation of clinical symptoms, and radiographic evidence of central nervous system demyelinating disorders, including multiple sclerosis. They may exacerbate symptoms in patients with New York Heart Association functional class III/IV heart failure.

ANTI-INTEGRINS Integrins are expressed on the cell surface of leukocytes and serve as mediators of leukocyte adhesion to vascular endothelium. $\alpha 4$ -Integrin along with its $\beta 1$ or $\beta 7$ subunit interact with endothelial ligands termed adhesion molecules. Interaction between $\alpha 4\beta 7$ and mucosal addressin cellular adhesion molecule (MAdCAM-1) is important in lymphocyte trafficking to gut mucosa.

Natalizumab is a recombinant humanized IgG4 antibody against $\alpha 4$ -integrin that has been shown to be effective in induction and maintenance of patients with CD. It has been approved since February 2008 for the treatment of patients with CD refractory or intolerant to anti-TNF therapy. The rates of response and remission at 3 months are about 60% and 40%, respectively, with a sustained remission rate of about 40% at 36 weeks.

One case of progressive multifocal leukoencephalopathy (PML) after eight infusions of natalizumab was observed among 1043 patients in the clinical trials for CD, and two patients developed PML in the multiple sclerosis (MS) trials after a median of 120 weeks. There were 410 postmarketing cases of PML, 408 in MS and 2 in CD. The most important risk factor for development of PML is exposure to the John Cunningham (JC) polyomavirus, seen in 50–55% of the adult population. The other two risk factors for development of PML are longer duration of treatment, especially beyond 2 years, and prior treatment with an immunosuppressant medication. Patients with all three risk factors have an estimated risk of 11:1000.

The FDA approved a commercial enzyme-linked immunosorbent assay (ELISA) kit to assay anti-JC viral antibodies (Stratify JCV

Antibody ELISA; Focus Diagnostics, Cypress, CA) in early 2012. The test is 99% accurate in stratifying risk of PML. It is recommended that all patients be tested prior to initiating natalizumab therapy. JC virus serologies are then measured every 6 months because 1–2% of patients will seroconvert yearly. All patients taking natalizumab and their providers must be enrolled in the TOUCH (Tysabri Outreach Unified Commitment for Health) pharmacovigilance program. Natalizumab is administered IV, 300 mg every 4 weeks. Labeling requirements mandate that it not be used in combination with any immunosuppressant medications.

Vedolizumab, another leukocyte trafficking inhibitor, is indicated for patients who have had an inadequate response or lost response to, or were intolerant of a TNF blocker or immunomodulator; or had an inadequate response or were intolerant to, or demonstrated dependence on glucocorticoids. It is an option for patients who are JC antibody positive since it does not cross the blood-brain barrier. Vedolizumab is a monoclonal antibody directed against $\alpha 4\beta 7$ integrin specifically and has the ability to convey gut-selective immunosuppression.

THERAPIES IN DEVELOPMENT

Ustekinumab, a fully human IgG1 monoclonal antibody, blocks the biologic activity of IL-12 and IL-23 through their common p40 subunit by inhibiting the interaction of these cytokines with their receptors on T cells, natural killer cells, and antigen presenting cells. It shows efficacy in moderate to severe CD in clinical trials.

Tofacitinib is an oral inhibitor of Janus kinases 1, 3, and, to a lesser extent, 2. It is expected to block signaling involving common gamma chain-containing cytokines including IL-2, IL-4, IL-7, IL-9, IL-15, and IL-21. These cytokines are integral to lymphocyte activation, function, and proliferation. It is effective in moderate to severe UC in clinical trials.

NUTRITIONAL THERAPIES

Dietary antigens may stimulate the mucosal immune response. Patients with active CD respond to bowel rest, along with TPN. Bowel rest and TPN are as effective as glucocorticoids at inducing remission of active CD but are not effective as maintenance therapy. Enteral nutrition in the form of elemental or peptide-based preparations is also as effective as glucocorticoids or TPN, but these diets are not palatable. Enteral diets may provide the small intestine with nutrients vital to cell growth and do not have the complications of TPN. In contrast to CD, dietary intervention does not reduce inflammation in UC. Standard medical management of UC and CD is shown in [Fig. 351-12](#).

SURGICAL THERAPY

Ulcerative Colitis Nearly one-half of patients with extensive chronic UC undergo surgery within the first 10 years of their illness. The indications for surgery are listed in [Table 351-8](#). Morbidity is about 20% for elective, 30% for urgent, and 40% for emergency proctocolectomy. The risks are primarily hemorrhage, contamination and sepsis, and neural injury. The operation of choice is an ileoanal J pouch anastomosis (IPAA).

Because UC is a mucosal disease, the rectal mucosa can be dissected and removed down to the dentate line of the anus or about 2 cm proximal to this landmark. The ileum is fashioned into a pouch that serves as a neorectum. This ileal pouch is then sutured circumferentially to the anus in an end-to-end fashion. If performed carefully, this operation preserves the anal sphincter and maintains continence. The overall operative morbidity is 10%, with the major complication being bowel obstruction. Pouch failure necessitating conversion to permanent ileostomy occurs in 5–10% of patients. Some inflamed rectal mucosa is usually left behind, and thus endoscopic surveillance is necessary. Primary dysplasia of the ileal mucosa of the pouch has occurred rarely.

Patients with IPAA usually have about 6–10 bowel movements a day. On validated quality-of-life indices, they report better performance in sports and sexual activities than ileostomy patients. The most frequent complication of IPAA is pouchitis in about 30–50%

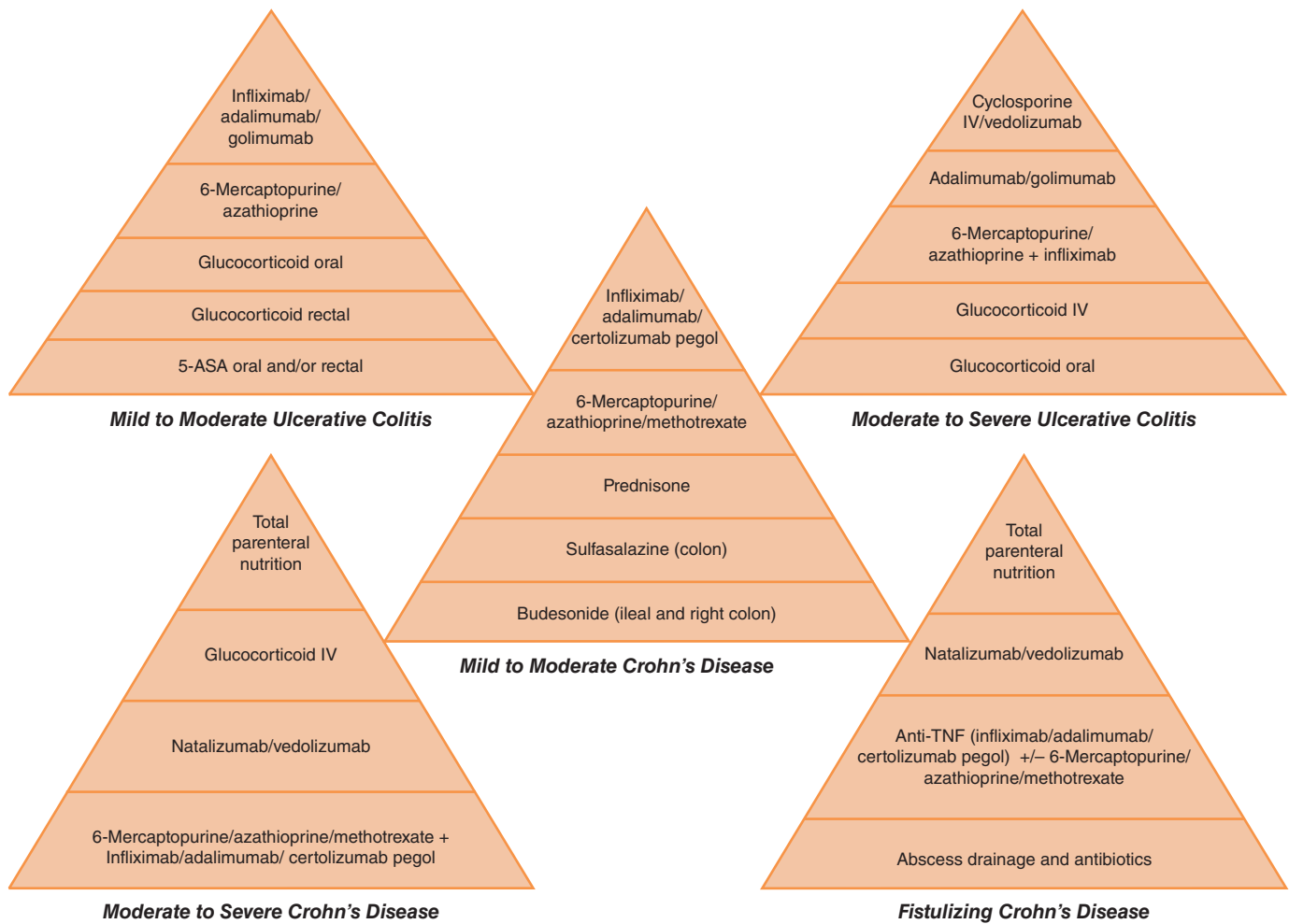


FIGURE 351-12 Medical management of inflammatory bowel disease. 5-ASA, 5-aminosalicylic acid; CD, Crohn's disease; UC, ulcerative colitis.

of patients with UC. This syndrome consists of increased stool frequency, watery stools, cramping, urgency, nocturnal leakage of stool, arthralgias, malaise, and fever. Pouch biopsies may distinguish true pouchitis from underlying CD. Although pouchitis usually responds to antibiotics, 3–5% of patients remain refractory and may require glucocorticoids, immunomodulators, anti-TNF therapy, or even pouch removal. A highly concentrated probiotic preparation with four strains of *Lactobacillus*, three strains of *Bifidobacterium*, and one strain of *Streptococcus salivarius* can prevent the recurrence of pouchitis when taken daily.

Crohn's Disease Most patients with CD require at least one operation in their lifetime. The need for surgery is related to duration of disease and the site of involvement. Patients with small-bowel disease have an 80% chance of requiring surgery. Those with colitis alone have a 50% chance. Surgery is an option only when medical treatment has failed or complications dictate its necessity. The indications for surgery are shown in Table 351-8.

SMALL INTESTINAL DISEASE Because CD is chronic and recurrent, with no clear surgical cure, as little intestine as possible is resected. Current surgical alternatives for treatment of obstructing CD include resection of the diseased segment and stricturoplasty. Surgical resection of the diseased segment is the most frequently performed operation, and in most cases, primary anastomosis can be done to restore continuity. If much of the small bowel has already been resected and the strictures are short, with intervening areas of normal mucosa, stricturoplasties should be done to avoid a functionally insufficient length of bowel. The strictured area of intestine is incised longitudinally and the incision sutured transversely, thus widening the narrowed area. Complications of stricturoplasty include prolonged ileus, hemorrhage, fistula, abscess, leak, and resticture.

There is evidence that mesalamine, nitroimidazole antibiotics, 6-MP/azathioprine, infliximab, and adalimumab are all superior to placebo for the prevention of postoperative recurrence of CD. Mesalamine is the least effective, and the side effects of the nitroimidazole antibiotics limit their use. Risk factors for early recurrence of disease include cigarette smoking, penetrating disease (internal fistulas, abscesses, or other evidence of penetration through the wall of the bowel), early recurrence since a previous surgery, multiple surgeries, and a young age at the time of the first surgery. Aggressive postoperative treatment with 6-MP/

TABLE 351-8 INDICATIONS FOR SURGERY

Ulcerative Colitis	Crohn's Disease
Intractable disease	Small Intestine
Fulminant disease	Stricture and obstruction
Toxic megacolon	unresponsive to medical therapy
Colonic perforation	Massive hemorrhage
Massive colonic hemorrhage	Refractory fistula
Extracolonic disease	Abscess
Colonic obstruction	Colon and rectum
Colon cancer prophylaxis	Intractable disease
Colon dysplasia or cancer	Fulminant disease
	Perianal disease unresponsive to medical therapy
	Refractory fistula
	Colonic obstruction
	Cancer prophylaxis
	Colon dysplasia or cancer

1964 azathioprine, infliximab, or adalimumab should be considered for this group of patients. It is also recommended to evaluate for endoscopic recurrence of CD via a colonoscopy, if possible, 6 months after surgery.

COLORECTAL DISEASE A greater percentage of patients with Crohn's colitis require surgery for intractability, fulminant disease, and anorectal disease. Several alternatives are available, ranging from the use of a temporary loop ileostomy to resection of segments of diseased colon or even the entire colon and rectum. For patients with segmental involvement, segmental colon resection with primary anastomosis can be performed. In 20–25% of patients with extensive colitis, the rectum is spared sufficiently to consider rectal preservation. Most surgeons believe that an IPAA is contraindicated in CD due to the high incidence of pouch failure. A diverting colostomy may help heal severe perianal disease or rectovaginal fistulas, but disease almost always recurs with reanastomosis. These patients often require a total proctocolectomy and ileostomy.

INFLAMMATORY BOWEL DISEASE AND PREGNANCY

Patients with quiescent UC and CD have normal fertility rates; the fallopian tubes can be scarred by the inflammatory process of CD, especially on the right side because of the proximity of the terminal ileum. In addition, perirectal, perineal, and rectovaginal abscesses and fistulae can result in dyspareunia. Infertility in men can be caused by sulfasalazine but reverses when treatment is stopped. In women who have an ileoanal J pouch anastomosis, most studies show that the fertility rate is reduced to about 50–80% of normal. This is due to scarring or occlusion of the fallopian tubes secondary to pelvic inflammation.

In mild or quiescent UC and CD, fetal outcome is nearly normal. Spontaneous abortions, stillbirths, and developmental defects are increased with increased disease activity, not medications. The courses of CD and UC during pregnancy mostly correlate with disease activity at the time of conception. Patients should be in remission for 6 months before conceiving. Most CD patients can deliver vaginally, but cesarean delivery may be the preferred route of delivery for patients with anorectal and perirectal abscesses and fistulas to reduce the likelihood of fistulas developing or extending into the episiotomy scar. Unless they desire multiple children, UC patients with an IPAA should consider a cesarean delivery due to an increased risk of future fecal incontinence.

Sulfasalazine, Lialda, Apriso, Delzicol, and balsalazide are safe for use in pregnancy and nursing with the caveat that additional folate supplementation must be given with sulfasalazine. Asacol HD and olsalazine are considered by the FDA to be class C agents in pregnancy and thus not recommended. Topical 5-ASA agents are also safe during pregnancy and nursing. Glucocorticoids are generally safe for use during pregnancy and are indicated for patients with moderate to severe disease activity. The amount of glucocorticoids received by the nursing infant is minimal. The safest antibiotics to use for CD in pregnancy for short periods of time (weeks, not months) are ampicillin and cephalosporins. Metronidazole can be used in the second or third trimester. Ciprofloxacin causes cartilage lesions in immature animals and should be avoided because of the absence of data on its effects on growth and development in humans.

6-MP and azathioprine pose minimal or no risk during pregnancy, but experience is limited. If the patient cannot be weaned from the drug or has an exacerbation that requires 6-MP/azathioprine during pregnancy, she should continue the drug with informed consent. Breast milk has been shown to contain negligible levels of 6-MP/azathioprine when measured in a limited number of patients.

Little data exist on CSA in pregnancy. In a small number of patients with severe IBD treated with IV CSA during pregnancy, 80% of pregnancies were successfully completed without development of renal toxicity or congenital malformations. However, because of the lack of data, CSA should probably be avoided unless the patient would otherwise require surgery.

MTX is contraindicated in pregnancy and nursing. In a large prospective study, no increased risk of stillbirths, miscarriages, or spontaneous abortions was seen with infliximab, adalimumab, or certolizumab, which are all class B drugs. Infliximab and adalimumab are IgG1 antibodies and are actively transported across the placenta in the late second and third trimester. Infants can have serum levels of both infliximab and adalimumab up to 7 months of age, and live vaccines should be avoided during this time. Certolizumab crosses the placenta by passive diffusion, and infant serum and cord blood levels are minimal. The anti-TNF drugs are relatively safe in nursing. Miniscule levels of both infliximab and adalimumab, but not certolizumab, have been reported in breast milk. These levels are of no clinical significance. It is recommended that drugs not be switched during pregnancy unless necessitated by the medical condition of the IBD. Natalizumab is considered as a class C drug because there is limited data in pregnancy.

Surgery in UC should be performed only for emergency indications, including severe hemorrhage, perforation, and megacolon refractory to medical therapy. Total colectomy and ileostomy carry a 50% risk of postoperative spontaneous abortion. Fetal mortality is also high in CD requiring surgery. Patients with IPAA have increased nighttime stool frequency during pregnancy that resolves postpartum. Transient small-bowel obstruction or ileus has been noted in up to 8% of patients with ileostomies.

CANCER IN INFLAMMATORY BOWEL DISEASE

ULCERATIVE COLITIS

Patients with long-standing UC are at increased risk for developing colonic epithelial dysplasia and carcinoma (**Fig. 351-13**).

The risk of neoplasia in chronic UC increases with duration and extent of disease. From one large meta-analysis, the risk of cancer in patients with UC is estimated at 2% after 10 years, 8% after 20 years, and 18% after 30 years of disease. Data from a 30-year surveillance program in the United Kingdom calculated the risk of colorectal cancer to be 7.7% at 20 years and 15.8% at 30 years of disease. The rates of colon cancer are higher than in the general population, and colonoscopic surveillance is the standard of care.

Annual or biennial colonoscopy with multiple biopsies is recommended for patients with >8–10 years of extensive colitis (greater than one-third of the colon involved) or 12–15 years of proctosigmoiditis (less than one-third but more than just the rectum) and has been widely used to screen and survey for subsequent dysplasia and carcinoma. Risk factors for cancer in UC include long-duration disease, extensive disease, family history of colon cancer, PSC, a colon stricture, and the presence of postinflammatory pseudopolyps on colonoscopy.

CROHN'S DISEASE

Risk factors for developing cancer in Crohn's colitis are long-duration and extensive disease, bypassed colon segments, colon strictures, PSC,

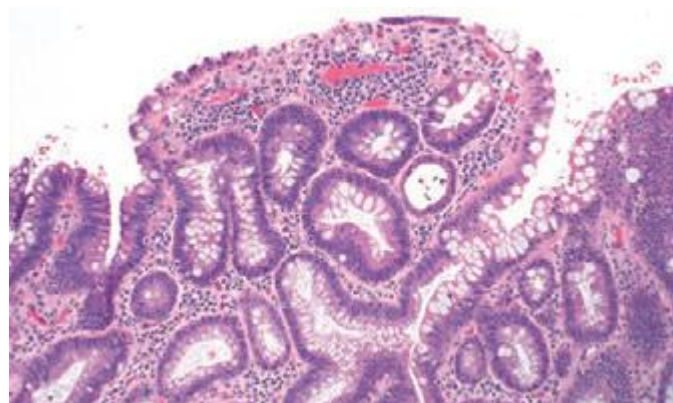


FIGURE 351-13 Medium-power view of low-grade dysplasia in a patient with chronic ulcerative colitis. Low-grade dysplastic crypts are interspersed among regenerating crypts. (Courtesy of Dr. R. Odze, Division of Gastrointestinal Pathology, Department of Pathology, Brigham and Women's Hospital, Boston, Massachusetts; with permission.)

and family history of colon cancer. The cancer risks in CD and UC are probably equivalent for similar extent and duration of disease. In the CESAME study, a prospective observational cohort of IBD patients in France, the standardized incidence ratios of colorectal cancer were 2.2 for all IBD patients (95% confidence interval [CI], 1.5–3.0; $p < .001$) and 7.0 for patients with long-standing extensive colitis (both Crohn's and UC) (95% CI, 4.4–10.5; $p < .001$). Thus, the same endoscopic surveillance strategy used for UC is recommended for patients with chronic Crohn's colitis. A pediatric colonoscope can be used to pass narrow strictures in CD patients, but surgery should be considered in symptomatic patients with impassable strictures.

MANAGEMENT OF DYSPLASIA AND CANCER

Dysplasia can be flat or polypoid. If flat high-grade dysplasia is encountered on colonoscopic surveillance, the usual treatment is colectomy for UC and either colectomy or segmental resection for CD. If flat low-grade dysplasia is found (Fig. 351-13), most investigators recommend immediate colectomy. Adenomas may occur coincidentally in UC and CD patients with chronic colitis and can be removed endoscopically provided that biopsies of the surrounding mucosa are free of dysplasia. High-definition and high-magnification colonoscopes and dye sprays have increased the rate of dysplasia detection.

IBD patients are also at greater risk for other malignancies. Patients with CD may have an increased risk of non-Hodgkin's lymphoma, leukemia, and myelodysplastic syndromes. Severe, chronic, complicated perianal disease in CD patients may be associated with an increased risk of cancer in the lower rectum and anal canal (squamous cell cancers). Although the absolute risk of small-bowel adenocarcinoma in CD is low (2.2% at 25 years in one study), patients with long-standing, extensive, small-bowel disease should consider screening.

352 Irritable Bowel Syndrome

Chung Owyang

Irritable bowel syndrome (IBS) is a functional bowel disorder characterized by abdominal pain or discomfort and altered bowel habits in the absence of detectable structural abnormalities. No clear diagnostic markers exist for IBS; thus the diagnosis of the disorder is based on clinical presentation. In 2006, the Rome II criteria for the diagnosis of IBS were revised (Table 352-1). Throughout the world, about 10–20% of adults and adolescents have symptoms consistent with IBS, and most studies show a female predominance. IBS symptoms tend to come and go over time and often overlap with other functional disorders such as fibromyalgia, headache, backache, and genitourinary symptoms. Severity of symptoms varies and can significantly impair quality of life, resulting in high health care costs. Advances in basic, mechanistic, and clinical investigations have improved our understanding of this disorder and its physiologic and psychosocial determinants. Altered

TABLE 352-1 DIAGNOSTIC CRITERIA FOR IRRITABLE BOWEL SYNDROME^a

Recurrent abdominal pain or discomfort^b at least 3 days per month in the last 3 months associated with two or more of the following:

1. Improvement with defecation
2. Onset associated with a change in frequency of stool
3. Onset associated with a change in form (appearance) of stool

^aCriteria fulfilled for the last 3 months with symptom onset at least 6 months prior to diagnosis. ^bDiscomfort means an uncomfortable sensation not described as pain. In pathophysiology research and clinical trials, a pain/discomfort frequency of at least 2 days a week during screening evaluation is required for subject eligibility.

Source: Adapted from GF Longstreth et al: *Gastroenterology* 130:1480, 2006.

gastrointestinal (GI) motility, visceral hyperalgesia, disturbance of brain-gut interaction, abnormal central processing, autonomic and hormonal events, genetic and environmental factors, and psychosocial disturbances are variably involved, depending on the individual. This progress may result in improved methods of treatment.

CLINICAL FEATURES

IBS is a disorder that affects all ages, although most patients have their first symptoms before age 45. Older individuals have a lower reporting frequency. Women are diagnosed with IBS two to three times as often as men and make up 80% of the population with severe IBS. As indicated in Table 352-1, pain or abdominal discomfort is a key symptom for the diagnosis of IBS. These symptoms should be improved with defecation and/or have their onset associated with a change in frequency or form of stool. Painless diarrhea or constipation does not fulfill the diagnostic criteria to be classified as IBS. Supportive symptoms that are not part of the diagnostic criteria include defecation straining, urgency or a feeling of incomplete bowel movement, passing mucus, and bloating.

Abdominal Pain According to the current IBS diagnostic criteria, abdominal pain or discomfort is a prerequisite clinical feature of IBS. Abdominal pain in IBS is highly variable in intensity and location. It is frequently episodic and crampy, but it may be superimposed on a background of constant ache. Pain may be mild enough to be ignored or it may interfere with daily activities. Despite this, malnutrition due to inadequate caloric intake is exceedingly rare with IBS. Sleep deprivation is also unusual because abdominal pain is almost uniformly present only during waking hours. However, patients with severe IBS frequently wake repeatedly during the night; thus, nocturnal pain is a poor discriminating factor between organic and functional bowel disease. Pain is often exacerbated by eating or emotional stress and improved by passage of flatus or stools. In addition, female patients with IBS commonly report worsening symptoms during the premenstrual and menstrual phases.

Altered Bowel Habits Alteration in bowel habits is the most consistent clinical feature in IBS. The most common pattern is constipation alternating with diarrhea, usually with one of these symptoms predominating. At first, constipation may be episodic, but eventually it becomes continuous and increasingly intractable to treatment with laxatives. Stools are usually hard with narrowed caliber, possibly reflecting excessive dehydration caused by prolonged colonic retention and spasm. Most patients also experience a sense of incomplete evacuation, thus leading to repeated attempts at defecation in a short time span. Patients whose predominant symptom is constipation may have weeks or months of constipation interrupted with brief periods of diarrhea. In other patients, diarrhea may be the predominant symptom. Diarrhea resulting from IBS usually consists of small volumes of loose stools. Most patients have stool volumes of <200 mL. Nocturnal diarrhea does not occur in IBS. Diarrhea may be aggravated by emotional stress or eating. Stool may be accompanied by passage of large amounts of mucus. Bleeding is not a feature of IBS unless hemorrhoids are present, and malabsorption or weight loss does not occur.

Bowel pattern subtypes are highly unstable. In a patient population with ~33% prevalence rates of IBS-diarrhea predominant (IBS-D), IBS-constipation predominant (IBS-C), and IBS-mixed (IBS-M) forms, 75% of patients change subtypes and 29% switch between IBS-C and IBS-D over 1 year. The heterogeneity and variable natural history of bowel habits in IBS increase the difficulty of conducting pathophysiology studies and clinical trials.

Gas and Flatulence Patients with IBS frequently complain of abdominal distention and increased belching or flatulence, all of which they attribute to increased gas. Although some patients with these symptoms actually may have a larger amount of gas, quantitative measurements reveal that most patients who complain of increased gas generate no more than a normal amount of intestinal gas. Most IBS patients have impaired transit and tolerance of intestinal gas loads. In addition, patients with IBS tend to reflux gas from the distal to the more proximal intestine, which may explain the belching.

Some patients with bloating may also experience visible distention with increase in abdominal girth. Both symptoms are more common among female patients and in those with higher overall Somatic Symptom Checklist scores. IBS patients who experienced bloating alone have been shown to have lower thresholds for pain and desire to defecate compared to those with concomitant distention irrespective of bowel habit. When patients were grouped according to sensory threshold, hyposensitive individuals had distention significantly more than those with hypersensitivity and this was observed more in the constipation subgroup. This suggests that the pathogenesis of bloating and distention may not be the same.

Upper Gastrointestinal Symptoms Between 25 and 50% of patients with IBS complain of dyspepsia, heartburn, nausea, and vomiting. This suggests that other areas of the gut apart from the colon may be involved. Prolonged ambulant recordings of small-bowel motility in patients with IBS show a high incidence of abnormalities in the small bowel during the diurnal (waking) period; nocturnal motor patterns are not different from those of healthy controls. The overlap between dyspepsia and IBS is great. The prevalence of IBS is higher among patients with dyspepsia (31.7%) than among those who reported no symptoms of dyspepsia (7.9%). Conversely, among patients with IBS, 55.6% reported symptoms of dyspepsia. In addition, the functional abdominal symptoms can change over time. Those with predominant dyspepsia or IBS can flux between the two. Although the prevalence of functional gastrointestinal disorders is stable over time, the turnover in symptom status is high. Many episodes of symptom disappearance are due to subjects changing symptoms rather than total symptom resolution. Thus it is conceivable that functional dyspepsia and IBS are two manifestations of a single, more extensive digestive system disorder. Furthermore, IBS symptoms are prevalent in noncardiac chest pain patients, suggesting overlap with other functional gut disorders.

PATHOPHYSIOLOGY

The pathogenesis of IBS is poorly understood, although roles of abnormal gut motor and sensory activity, central neural dysfunction, psychological disturbances, mucosal inflammation, stress, and luminal factors have been proposed.

Gastrointestinal Motor Abnormalities Studies of colonic myoelectrical and motor activity under unstimulated conditions have not shown consistent abnormalities in IBS. In contrast, colonic motor abnormalities are more prominent under stimulated conditions in IBS. IBS patients may exhibit increased rectosigmoid motor activity for up to 3 h after eating. Similarly, inflation of rectal balloons both in IBS-D and IBS-C patients leads to marked and prolonged distention-evoked contractile activity. Recordings from the transverse, descending, and sigmoid colon showed that the motility index and peak amplitude of high-amplitude propagating contractions (HAPCs) in diarrhea-prone IBS patients were greatly increased compared to those in healthy subjects and were associated with rapid colonic transit and accompanied by abdominal pain.

Visceral Hypersensitivity As with studies of motor activity, IBS patients frequently exhibit exaggerated sensory responses to visceral stimulation. The frequency of perceptions of food intolerance is at least two-fold more common than in the general population. Postprandial pain has been temporally related to entry of the food bolus into the cecum in 74% of patients. On the other hand, prolonged fasting in IBS patients is often associated with significant improvement in symptoms. Rectal balloon inflation produces nonpainful and painful sensations at lower volumes in IBS patients than in healthy controls without altering rectal tension, suggestive of visceral afferent dysfunction in IBS. Similar studies show gastric and esophageal hypersensitivity in patients with nonulcer dyspepsia and noncardiac chest pain, raising the possibility that these conditions have a similar pathophysiologic basis. Lipids lower the thresholds for the first sensation of gas, discomfort, and pain in IBS patients. Hence, postprandial symptoms in IBS patients may be explained in part by a nutrient-dependent exaggerated sensory component of the gastrocolonic response. In contrast to enhanced gut sensitivity, IBS patients do not exhibit heightened sensitivity

TABLE 352-2 PROPOSED MECHANISMS FOR VISCERAL HYPERSENSITIVITY

End-organ sensitivity	Long-term hyperalgesia
“Silent” nociceptors	Tonic cortical regulation
CNS modulation	Neuroplasticity
Cortex	
Brainstem	

Abbreviation: CNS, central nervous system.

elsewhere in the body. Thus, the afferent pathway disturbances in IBS appear to be selective for visceral innervation with sparing of somatic pathways. The mechanisms responsible for visceral hypersensitivity are still under investigation. It has been proposed that these exaggerated responses may be due to (1) increased end-organ sensitivity with recruitment of “silent” nociceptors; (2) spinal hyperexcitability with activation of nitric oxide and possibly other neurotransmitters; (3) endogenous (cortical and brainstem) modulation of caudad nociceptive transmission; and (4) over time, the possible development of long-term hyperalgesia due to development of neuroplasticity, resulting in permanent or semipermanent changes in neural responses to chronic or recurrent visceral stimulation (**Table 352-2**).

Central Neural Dysregulation The role of central nervous system (CNS) factors in the pathogenesis of IBS is strongly suggested by the clinical association of emotional disorders and stress with symptom exacerbation and the therapeutic response to therapies that act on cerebral cortical sites. Functional brain imaging studies such as magnetic resonance imaging (MRI) have shown that in response to distal colonic stimulation, the mid-cingulate cortex—a brain region concerned with attention processes and response selection—shows greater activation in IBS patients. Modulation of this region is associated with changes in the subjective unpleasantness of pain. In addition, IBS patients also show preferential activation of the prefrontal lobe, which contains a vigilance network within the brain that increases alertness. These may represent a form of cerebral dysfunction leading to the increased perception of visceral pain.

Abnormal Psychological Features Abnormal psychiatric features are recorded in up to 80% of IBS patients, especially in referral centers; however, no single psychiatric diagnosis predominates. Most of these patients demonstrated exaggerated symptoms in response to visceral distention, and this abnormality persists even after exclusion of psychological factors.

Psychological factors influence pain thresholds in IBS patients, as stress alters sensory thresholds. An association between prior sexual or physical abuse and development of IBS has been reported. Abuse is associated with greater pain reporting, psychological distress, and poor health outcome. Brain functional MRI studies show greater activation of the posterior and middle dorsal cingulate cortex, which is implicated in affect processing in IBS patients with a past history of sexual abuse.

Thus, patients with IBS frequently demonstrate increased motor reactivity of the colon and small bowel to a variety of stimuli and altered visceral sensation associated with lowered sensation thresholds. These may result from CNS–enteric nervous system dysregulation (**Fig. 352-1**).

Postinfectious IBS IBS may be induced by GI infection. In an investigation of 544 patients with confirmed bacterial gastroenteritis, one-quarter developed IBS subsequently. Conversely, about a third of IBS patients experienced an acute “gastroenteritis-like” illness at the onset of their chronic IBS symptomatology. This group of “postinfective” IBS occurs more commonly in females and affects younger rather than older patients. Risk factors for developing postinfectious IBS include, in order of importance, prolonged duration of initial illness, toxicity of infecting bacterial strain, smoking, mucosal markers of inflammation, female gender, depression, hypochondriasis, and adverse life events in the preceding 3 months. Age older than 60 years might protect against postinfectious IBS, whereas treatment with antibiotics has been

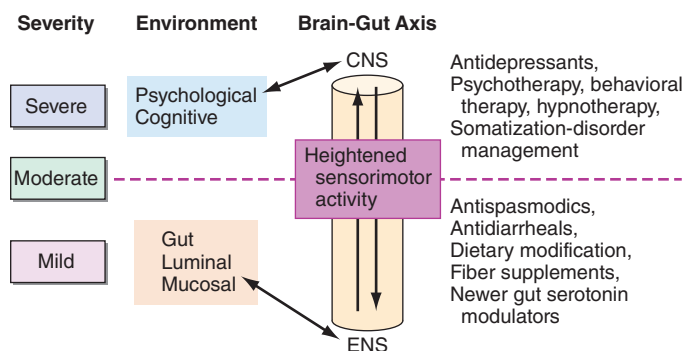


FIGURE 352-1 Therapeutic targets for irritable bowel syndrome.

Patients with mild to moderate symptoms usually have intermittent symptoms that correlate with altered gut physiology. Treatments include gut-acting pharmacologic agents such as antispasmodics, antidiarrheals, fiber supplements, and gut serotonin modulators. Patients who have severe symptoms usually have constant pain and psychosocial difficulties. This group of patients is best managed with antidepressants and other psychosocial treatments. CNS, central nervous system; ENS, enteric nervous system.

associated with increased risk. The microbes involved in the initial infection are *Campylobacter*, *Salmonella*, and *Shigella*. Those patients with *Campylobacter* infection who are toxin-positive are more likely to develop postinfective IBS. Increased rectal mucosal enteroendocrine cells, T lymphocytes, and increased gut permeability are acute changes following *Campylobacter* enteritis that could persist for more than a year and may contribute to postinfective IBS.

Immune Activation and Mucosal Inflammation Some patients with IBS display persistent signs of low-grade mucosal inflammation with activated lymphocytes, mast cells, and enhanced expression of proinflammatory cytokines. These abnormalities may contribute to abnormal epithelial secretion and visceral hypersensitivity. There is increasing evidence that some members of the superfamily of transient receptor potential (TRP) cation channels such as TRPV1 (vanilloid) channels are central to the initiation and persistence of visceral hypersensitivity. Mucosal inflammation can lead to increased expression of TRPV1 in the enteric nervous system. Enhanced expression of TRPV1 channels in the sensory neurons of the gut has been observed in IBS, and such expression appears to correlate with visceral hypersensitivity and abdominal pain. Interestingly, clinical studies have also shown increased intestinal permeability in patients with IBS-D. Psychological stress and anxiety can increase the release of proinflammatory cytokine, and this in turn may alter intestinal permeability. This provides a functional link between psychological stress, immune activation, and symptom generation in patients with IBS.

Altered Gut Flora A high prevalence of small intestinal bacterial overgrowth in IBS patients has been noted based on positive lactulose hydrogen breath test. This finding, however, has been challenged by a number of other studies that found no increased incidence of bacterial overgrowth based on jejunal aspirate culture. Abnormal H_2 breath

test can occur because of small-bowel rapid transit and may lead to erroneous interpretation. Hence, the role of testing for small intestinal bacterial overgrowth in IBS patients remains unclear.

Studies using culture-independent approaches such as 16S rRNA gene-based analysis found significant differences between the molecular profile of the fecal microbiota of IBS patients and that of healthy subjects. IBS patients had decreased proportions of the genera *Bifidobacterium* and *Lactobacillus* and increased ratios of Firmicutes:Bacteroidetes. It has been speculated that these changes may be related to stress and diet. A temporary reduction in lactobacilli has been reported in animal models of early-life stress. On the other hand, Firmicutes is the dominant phylum in adults consuming a diet high in animal fat and protein. However, it is still unclear whether such changes in fecal microbiota are causal, consequential, or merely the result of constipation and diarrhea. In addition, the stability of the change in the microbiota needs to be determined.

Abnormal Serotonin Pathways The serotonin (5-HT)-containing enterochromaffin cells in the colon are increased in a subset of IBS-D patients compared to healthy individuals or patients with ulcerative colitis. Furthermore, postprandial plasma 5-HT levels were significantly higher in this group of patients compared to healthy controls. Because serotonin plays an important role in the regulation of GI motility and visceral perception, the increased release of serotonin may contribute to the postprandial symptoms of these patients and provides a rationale for the use of serotonin antagonists in the treatment of this disorder.

APPROACH TO THE PATIENT: Irritable Bowel Syndrome

Because IBS is a disorder for which no pathognomonic abnormalities have been identified, its diagnosis relies on recognition of positive clinical features and elimination of other organic diseases. Symptom-based criteria have been developed for the purpose of differentiating patients with IBS from those with organic diseases. These include the Manning, Rome I, Rome II, and Rome III criteria (Table 352-1). The diagnostic values of these criteria are shown in Table 352-3. In a validation study, Rome III performed less well than either the Rome I and II criteria and all criteria studied to date showed positive predictive values of <50%, which underscores the need for developing diagnostic strategies for IBS that are more cost-effective than the current approaches. A careful history and physical examination are frequently helpful in establishing the diagnosis. Clinical features suggestive of IBS include the following: recurrence of lower abdominal pain with altered bowel habits over a period of time without progressive deterioration, onset of symptoms during periods of stress or emotional upset, absence of other systemic symptoms such as fever and weight loss, and small-volume stool without any evidence of blood.

On the other hand, the appearance of the disorder for the first time in old age, progressive course from time of onset, persistent diarrhea after a 48-h fast, and presence of nocturnal diarrhea or steatorrheal stools argue against the diagnosis of IBS.

TABLE 352-3 SENSITIVITY, SPECIFICITY, POSITIVE AND NEGATIVE PREDICTIVE VALUES, AND POSITIVE AND NEGATIVE LIKELIHOOD RATIOS FOR THE ROME AND MANNING CRITERIA FOR IRRITABLE BOWEL SYNDROME^a

	Sensitivity, % (95% CI)	Specificity, % (95% CI)	Positive Predictive Value, % (95% CI)	Negative Predictive Value, % (95% CI)	Positive Likelihood Ratio (95% CI)	Negative Likelihood Ratio (95% CI)
Rome III criteria	17.4 (13.9–21.5)	95.6 (94.4–96.5)	49.6 (42.0–58.7)	82.1 (80.0–83.6)	3.92 (2.85–5.38)	0.86 (0.83–0.91)
Rome II criteria	23.3 (19.4–27.8)	94.5 (93.2–95.5)	51.7 (44.9–59.5)	82.9 (80.8–84.4)	4.21 (3.20–5.53)	0.81 (0.77–0.86)
Rome I criteria	24.3 (20.3–28.8)	93.9 (92.6–95.0)	50.5 (44.0–58.1)	83.0 (80.9–84.4)	4.01 (3.08–5.22)	0.81 (0.76–0.85)
Manning criteria (3 criteria)	13.7 (10.6–17.6)	97.1 (96.1–97.8)	54.1 (45.3–64.6)	81.6 (79.6–83.1)	4.66 (3.18–6.82)	0.89 (0.85–0.93)

^aExcluding individuals reporting lower gastrointestinal alarm symptoms from the definition of irritable bowel syndrome.

Because the major symptoms of IBS—abdominal pain, abdominal bloating, and alteration in bowel habits—are common complaints of many GI organic disorders, the list of differential diagnoses is a long one. The quality, location, and timing of pain may be helpful to suggest specific disorders. Pain due to IBS that occurs in the epigastric or periumbilical area must be differentiated from biliary tract disease, peptic ulcer disorders, intestinal ischemia, and carcinoma of the stomach and pancreas. If pain occurs mainly in the lower abdomen, the possibility of diverticular disease of the colon, inflammatory bowel disease (including ulcerative colitis and Crohn's disease), and carcinoma of the colon must be considered. Postprandial pain accompanied by bloating, nausea, and vomiting suggests gastroparesis or partial intestinal obstruction. Intestinal infestation with *Giardia lamblia* or other parasites may cause similar symptoms. When diarrhea is the major complaint, the possibility of lactase deficiency, laxative abuse, malabsorption, celiac sprue, hyperthyroidism, inflammatory bowel disease, and infectious diarrhea must be ruled out. On the other hand, constipation may be a side effect of many different drugs, such as anticholinergic, antihypertensive, and antidepressant medications. Endocrinopathies such as hypothyroidism and hypoparathyroidism must also be considered in the differential diagnosis of constipation, particularly if other systemic signs or symptoms of these endocrinopathies are present. In addition, acute intermittent porphyria and lead poisoning may present in a fashion similar to IBS, with painful constipation as the major complaint. These possibilities are suspected on the basis of their clinical presentations and are confirmed by appropriate serum and urine tests.

Few tests are required for patients who have typical IBS symptoms and no alarm features. Unnecessary investigations may be costly and even harmful. The American Gastroenterological Association has delineated factors to be considered when determining the aggressiveness of the diagnostic evaluation. These include the duration of symptoms, the change in symptoms over time, the age and sex of the patient, the referral status of the patient, prior diagnostic studies, a family history of colorectal malignancy, and the degree of psychosocial dysfunction. Thus, a younger individual with mild symptoms requires a minimal diagnostic evaluation, while an older person or an individual with rapidly progressive symptoms should undergo a more thorough exclusion of organic disease. Most patients should have a complete blood count and sigmoidoscopic examination; in addition, stool specimens should be examined for ova and parasites in those who have diarrhea. In patients with persistent diarrhea not responding to simple

antidiarrheal agents, a sigmoid colon biopsy should be performed to rule out microscopic colitis. In those age >40 years, an air-contrast barium enema or colonoscopy should also be performed. If the main symptoms are diarrhea and increased gas, the possibility of lactase deficiency should be ruled out with a hydrogen breath test or with evaluation after a 3-week lactose-free diet. Some patients with IBS-D may have undiagnosed celiac sprue. Because the symptoms of celiac sprue respond to a gluten-free diet, testing for celiac sprue in IBS may prevent years of morbidity and attendant expense. Decision-analysis studies show that serology testing for celiac sprue in patients with IBS-D has an acceptable cost when the prevalence of celiac sprue is >1% and is the dominant strategy when the prevalence is >8%. In patients with concurrent symptoms of dyspepsia, upper GI radiographs or esophagogastroduodenoscopy may be advisable. In patients with postprandial right upper quadrant pain, an ultrasonogram of the gallbladder should be obtained. Laboratory features that argue against IBS include evidence of anemia, elevated sedimentation rate, presence of leukocytes or blood in stool, and stool volume >200–300 mL/d. These findings would necessitate other diagnostic considerations.

TREATMENT IRRITABLE BOWEL SYNDROME

PATIENT COUNSELING AND DIETARY ALTERATIONS

Reassurance and careful explanation of the functional nature of the disorder and of how to avoid obvious food precipitants are important first steps in patient counseling and dietary change. Occasionally, a meticulous dietary history may reveal substances (such as coffee, disaccharides, legumes, and cabbage) that aggravate symptoms. Excessive fructose and artificial sweeteners, such as sorbitol or mannitol, may cause diarrhea, bloating, cramping, or flatulence. As a therapeutic trial, patients should be encouraged to eliminate any foodstuffs that appear to produce symptoms. However patients should avoid nutritionally depleted diets. A diet low in fermentable oligosaccharides, disaccharides, monosaccharides, and polyols (FODMAPs) (Table 352-4) has been shown to be helpful in IBS patients. FODMAPs are poorly absorbed by the small intestine and fermented by bacteria in the colon to produce gas and osmotically active carbohydrates. Clinical studies demonstrate that in IBS patients, ingestion of FODMAPs such as lactose, fructose, or sorbitol, alone or in combination, produce gut symptoms such as gas and diarrhea. On the other hand, a randomized controlled study showed that a diet low in FODMAPs reduced symptoms in IBS patients. This

TABLE 352-4 SOME COMMON FOOD SOURCES OF FODMAPs

Food Type	Free Fructose	Lactose	Fructans	Galacto-oligosaccharides	Polyols
Fruits	Apple, cherry, mango, pear, watermelon		Peach, persimmon, watermelon		Apple, apricot, pear, avocado, blackberries, cherry, nectarine, plum, prune
Vegetables	Asparagus, artichokes, sugar snap peas		Artichokes, beetroot, Brussels sprout, chicory, fennel, garlic, leek, onion, peas		Cauliflower, mushroom, snow peas
Grains and cereals			Wheat, rye, barley		
Nuts and seeds			Pistachios		
Milk and milk products		Milk, yogurt, ice cream, custard, soft cheeses			
Legumes			Legumes, lentils, chickpeas	Legumes, chickpeas, lentils	
Other	Honey, high-fructose corn syrup		Chicory drinks		
Food additives			Inulin, FOS		Sorbitol, mannitol, maltitol, xylitol, isomalt

Abbreviations: FODMAPs, fermentable oligosaccharides, disaccharides, monosaccharides, and polyols; FOS, fructo-oligosaccharides

Source: Adapted from PR Gibson et al: Am J Gastroenterol 107:657, 2012.

approach may be used in diarrhea-predominant IBS patients with severe gas and bloating. Durable adherence can be expected in up to 75% of patients.

Stool-Bulking Agents High-fiber diets and bulking agents, such as bran or hydrophilic colloid, are frequently used in treating IBS. The water-holding action of fibers may contribute to increased stool bulk because of the ability of fiber to increase fecal output of bacteria. Fiber also speeds up colonic transit in most persons. In diarrhea-prone patients, whole-colonic transit is faster than average; however, dietary fiber can delay transit. Furthermore, because of their hydrophilic properties, stool-bulking agents bind water and thus prevent both excessive hydration and dehydration of stool. The latter observation may explain the clinical experience that a high-fiber diet relieves diarrhea in some IBS patients. Fiber supplementation with psyllium has been shown to reduce perception of rectal distention, indicating that fiber may have a positive effect on visceral afferent function.

The beneficial effects of dietary fiber on colonic physiology suggest that dietary fiber should be an effective treatment for IBS patients, but controlled trials of dietary fiber have produced variable results. This is not surprising since IBS is a heterogeneous disorder, with some patients being constipated and other having predominant diarrhea. Most investigations report increases in stool weight, decreases in colonic transit times, and improvement in constipation. Others have noted benefits in patients with alternating diarrhea and constipation, pain, and bloating. However, most studies observe no responses in patients with diarrhea- or pain-predominant IBS. It is possible that different fiber preparations may have dissimilar effects on selected symptoms in IBS. A cross-over comparison of different fiber preparations found that psyllium produced greater improvements in stool pattern and abdominal pain than bran. Furthermore, psyllium preparations tend to produce less bloating and distention. Despite the equivocal data regarding efficacy, most gastroenterologists consider stool-bulking agents worth trying in patients with IBS-C. Fiber should be started at a nominal dose and slowly titrated up as tolerated over the course of several weeks to a targeted dose of 20–30 g of total dietary and supplementary fiber per day. Even when used judiciously, fiber can exacerbate bloating, flatulence, constipation, and diarrhea.

Antispasmodics Clinicians have observed that anticholinergic drugs may provide temporary relief for symptoms such as painful cramps related to intestinal spasm. Although controlled clinical trials have produced mixed results, evidence generally supports beneficial effects of anticholinergic drugs for pain. A meta-analysis of 26 double-blind clinical trials of antispasmodic agents in IBS reported better global improvement (62%) and abdominal pain reductions (64%) compared to placebo (35% and 45%, respectively), suggesting efficacy in some patients. The drugs are most effective when prescribed in anticipation of predictable pain. Physiologic studies demonstrate that anticholinergic drugs inhibit the gastrocolic reflex; hence, postprandial pain is best managed by giving antispasmodics 30 min before meals so that effective blood levels are achieved shortly before the anticipated onset of pain. Most anticholinergics contain natural belladonna alkaloids, which may cause xerostomia, urinary hesitancy and retention, blurred vision, and drowsiness. They should be used in the elderly with caution. Some physicians prefer to use synthetic anticholinergics such as dicyclomine that have less effect on mucous membrane secretions and produce fewer undesirable side effects.

Antidiarrheal Agents Peripherally acting opiate-based agents are the initial therapy of choice for IBS-D. Physiologic studies demonstrate increases in segmenting colonic contractions, delays in fecal transit, increases in anal pressures, and reductions in rectal perception with these drugs. When diarrhea is severe, especially in the painless diarrhea variant of IBS, small doses of loperamide, 2–4 mg every 4–6 h up to a maximum of 12 g/d, can be prescribed. These agents are less addictive than paregoric, codeine, or tincture of opium. In general, the intestines do not become tolerant of the antidiarrheal effect of opiates, and increasing doses are not required to maintain antidiarrheal

potency. These agents are most useful if taken before anticipated stressful events that are known to cause diarrhea. However, not infrequently, a high dose of loperamide may cause cramping because of increases in segmenting colonic contractions. Another antidiarrheal agent that may be used in IBS patients is the bile acid binder cholestyramine resin.

Antidepressant Drugs In addition to their mood-elevating effects, antidepressant medications have several physiologic effects that suggest they may be beneficial in IBS. In IBS-D patients, the tricyclic antidepressant imipramine slows jejunal migrating motor complex transit propagation and delays orocecal and whole-gut transit, indicative of a motor inhibitory effect. Some studies also suggest that tricyclic agents may alter visceral afferent neural function.

A number of studies indicate that tricyclic antidepressants may be effective in some IBS patients. In a 2-month study of desipramine, abdominal pain improved in 86% of patients compared to 59% given placebo. Another study of desipramine in 28 IBS patients showed improvement in stool frequency, diarrhea, pain, and depression. When stratified according to the predominant symptoms, improvements were observed in IBS-D patients, with no improvement being noted in IBS-C patients. The beneficial effects of the tricyclic compounds in the treatment of IBS appear to be independent of their effects on depression. The therapeutic benefits for the bowel symptoms occur faster and at a lower dosage. The efficacy of antidepressant agents in other chemical classes in the management of IBS is less well evaluated. In contrast to tricyclic agents, the selective serotonin reuptake inhibitor (SSRI) paroxetine accelerates orocecal transit, raising the possibility that this drug class may be useful in IBS-C patients. The SSRI citalopram blunts perception of rectal distention and reduces the magnitude of the gastrocolonic response in healthy volunteers. A small placebo-controlled study of citalopram in IBS patients reported reductions in pain. However, these findings could not be confirmed in another randomized controlled trial that showed that citalopram at 20 mg/d for 4 weeks was not superior to placebo in treating nondepressed IBS patients. Hence, the efficacy of SSRIs in the treatment of IBS needs further confirmation.

Antiflatulence Therapy The management of excessive gas is seldom satisfactory, except when there is obvious aerophagia or disaccharidase deficiency. Patients should be advised to eat slowly and not chew gum or drink carbonated beverages. Bloating may decrease if an associated gut syndrome such as IBS or constipation is improved. If bloating is accompanied by diarrhea and worsens after ingesting dairy products, fresh fruits, vegetables, or juices, further investigation or a dietary exclusion trial may be worthwhile. Avoiding flatogenic foods, exercising, losing excess weight, and taking activated charcoal are safe but unproven remedies. Data regarding the use of surfactants such as simethicone are conflicting. Antibiotics may help in a subgroup of IBS patients with predominant symptoms of bloating. Beano, an over-the-counter oral β -glycosidase solution, may reduce rectal passage of gas without decreasing bloating and pain. Pancreatic enzymes reduce bloating, gas, and fullness during and after high-calorie, high-fat meal ingestion.

Modulation of Gut Flora Antibiotic treatment benefits a subset of IBS patients. In a double-blind, randomized, placebo-controlled study, neomycin dosed at 500 mg twice daily for 10 days was more effective than placebo at improving symptom scores among IBS patients. The nonabsorbed oral antibiotic rifaximin is the most thoroughly studied antibiotic for the treatment of IBS. In a double-blind, placebo-controlled study, patients receiving rifaximin at a dose of 550 mg two times daily for 2 weeks experienced substantial improvement of global IBS symptoms over placebo. Rifaximin is the only antibiotic with demonstrated sustained benefit beyond therapy cessation in IBS patients. The drug has a favorable safety and tolerability profile compared with systemic antibiotics. A systematic review and meta-analysis of five studies of IBS patients found that rifaximin is more effective than placebo for global symptoms and bloating (odds ratio 1.57) with a number needed to treat (NNT) of

1970 10.2. The modest therapeutic gain was similar to that yielded by other current available therapies for IBS. However, currently there are still insufficient data to recommend routine use of this antibiotic in the treatment of IBS.

Because altered colonic flora may contribute to the pathogenesis of IBS, this has led to great interest in using probiotics to naturally alter the flora. A meta-analysis of 10 probiotic studies in IBS patients found significant relief of pain and bloating with the use of *Bifidobacterium breve*, *B. longum*, and *Lactobacillus acidophilus* species compared to placebo. However, there was no change in stool frequency or consistency. Large-scale studies of well-phenotyped IBS patients are needed to establish the efficacy of these probiotics.

Serotonin Receptor Agonist and Antagonists Serotonin receptor antagonists have been evaluated as therapies for IBS-D. Serotonin acting on 5-HT₃ receptors enhances the sensitivity of afferent neurons projecting from the gut. In humans, a 5-HT₃ receptor antagonist such as alosetron reduces perception of painful visceral stimulation in IBS. It also induces rectal relaxation, increases rectal compliance, and delays colonic transit. Meta-analysis of 14 randomized controlled trials of alosetron or cilansetron showed that these antagonists are more effective than placebo in achieving global improvement in IBS symptoms and relief of abdominal pain and discomfort. These agents are more likely to cause constipation in IBS patients with diarrhea alternating with constipation. Also, 0.2% of patients using 5-HT₃ antagonists developed ischemic colitis versus none in the control group. In postrelease surveillance, 84 cases of ischemic colitis were observed, including 44 cases that required surgery and 4 deaths. As a consequence, the medication was voluntarily withdrawn by the manufacturer in 2000. Alosetron has been reintroduced under a new risk-management program where patients have to sign a patient-physician agreement. This has significantly limited its usage.

Novel 5-HT₄ receptor agonists such as tegaserod exhibit prokinetic activity by stimulating peristalsis. In IBS patients with constipation, tegaserod accelerated intestinal and ascending colon transit. Clinical trials involving >4000 IBS-C patients reported reductions in discomfort and improvements in constipation and bloating, compared to placebo. Diarrhea is the major side effect. However, tegaserod has been withdrawn from the market; a meta-analysis revealed an increase in serious cardiovascular events.

Chloride Channel Activators Lubiprostone is a bicyclic fatty acid that stimulates chloride channels in the apical membrane of intestinal epithelial cells. Chloride secretion induces passive movement of sodium and water into the bowel lumen and improves bowel function. Oral lubiprostone was effective in the treatment of patients with constipation-predominant IBS in large phase II and phase III randomized, double-blinded, placebo-controlled multicenter trials. Responses were significantly greater in patients receiving lubiprostone 8 µg twice daily for 3 months than in those receiving placebo. In general, the drug was quite well tolerated. The major side effects are nausea and diarrhea. Lubiprostone is a new class of compounds for treatment of chronic constipation with or without IBS.

Guanylate Cyclase-C Agonist Linaclotide is a minimally absorbed 14-amino-acid peptide guanylate cyclase-C (GC-C) agonist that binds to and activates GC-C on the luminal surface of intestinal epithelium. Activation of GC-C results in generation of cyclic guanosine monophosphate (cGMP), which triggers secretion of fluid, sodium, and bicarbonate. In animal models, linaclotide accelerates GI transit and reduces visceral nociception. The analgesic action of linaclotide appears to be mediated by cGMP acting on afferent pain fibers innervating the GI tract. A phase III, double-blind, controlled trial showed that linaclotide, 290 µg given once daily, significantly improved abdominal pain, bloating, and spontaneous bowel movement. The only significant side effect was diarrhea, which occurred in 4.5% of the patients. The drug has been approved for treatment of constipation in IBS-C patients.

TABLE 352-5 SPECTRUM OF SEVERITY IN IBS

	Mild	Moderate	Severe
Clinical Features			
Prevalence	70%	25%	5%
Correlations with gut physiology	+++	++	+
Symptoms constant	0	+	+++
Psychosocial difficulties	0	+	+++
Health care issues	+	++	+++
Practice type	Primary	Specialty	Referral

SUMMARY

The treatment strategy of IBS depends on the severity of the disorder (Table 352-5). Most IBS patients have mild symptoms. They are usually cared for in primary care practices, have little or no psychosocial difficulties, and do not seek health care often. Treatment usually involves education, reassurance, and dietary/lifestyle changes. A smaller portion have moderate symptoms that are usually intermittent and correlate with altered gut physiology, e.g., worsened with eating or stress and relieved by defecation. For IBS-D patients, treatments include gut-acting pharmacologic agents such as antispasmodics, antidiarrheals, bile acid binders, and the newer gut serotonin modulators (Table 352-6). In IBS-C patients, increased fiber intake and the use of osmotic agents such as polyethylene glycol may achieve satisfactory results. For patients with more severe constipation, a chloride channel opener (lubiprostone) or GC-C agonist (linaclotide) may be considered. For IBS patients with predominant gas and bloating, a low-FODMAP diet may provide significant relief. Some patients may benefit from probiotics and rifaximin treatment. A small proportion of IBS patients have severe and refractory symptoms, are usually seen in referral centers, and frequently have constant pain and psychosocial difficulties (Fig. 352-1). This group of patients is best managed with antidepressants and other psychological treatments (Table 352-6).

TABLE 352-6 POSSIBLE DRUGS FOR A DOMINANT SYMPTOM IN IBS

Symptom	Drug	Dose
Diarrhea	Loperamide	2–4 mg when necessary/ maximum 12 g/d
	Cholestyramine resin	4 g with meals
	Alosetron ^a	0.5–1 mg bid (for severe IBS, women)
Constipation	Psyllium husk	3–4 g bid with meals, then adjust
	Methylcellulose	2 g bid with meals, then adjust
	Calcium polycarbophil	1 g qd to qid
	Lactulose syrup	10–20 g bid
	70% sorbitol	15 mL bid
	Polyethylene glycol 3350	17 g in 250 mL water qd
	Lubiprostone (Amitiza)	24 mg bid
Abdominal pain	Magnesium hydroxide	30–60 mL qd
	Linaclotide	290 µg qd
	Smooth-muscle relaxant	qd to qid ac
	Tricyclic antidepressants	Start 25–50 mg hs, then adjust
Gas and bloating	Selective serotonin reuptake inhibitors	Begin small dose, increase as needed
	Low FODMAP diet	
	Probiotics	qd
	Rifaximin	550 mg bid

^aAvailable only in the United States.

Abbreviation: FODMAP, fermentable oligosaccharides, disaccharides, monosaccharides, and polyols.

Source: Adapted from GF Longstreth et al: Gastroenterology 130:1480, 2006.

353 Diverticular Disease and Common Anorectal Disorders

Rizwan Ahmed, Susan L. Gearhart

DIVERTICULAR DISEASE



Incidence and Epidemiology In the United States, diverticulosis affects 70% of the population above the age of 80. Fortunately, only 20% of patients with diverticulosis develop symptomatic disease, 1–2% require hospitalization, and <1% will require surgery. Diverticular disease has become the fifth most costly gastrointestinal disorder in the United States. Previously overlooked, the majority of patients with diverticular disease report a lower health-related quality of life and more depression as compared to matched controls, thus adding to health care costs. Formerly, diverticular disease was confined to developed countries; however, with the adoption of westernized diets in underdeveloped countries, diverticulosis is on the rise across the globe. Immigrants to the United States develop diverticular disease at the same rate as U.S. natives. Although the prevalence among females and males is similar, males tend to present at a younger age. The mean age at presentation of the disease is 59 years and is now shifting to affect younger populations.

Anatomy and Pathophysiology Two types of diverticula occur in the intestine: true and false (or pseudo diverticula). A true diverticulum is a saclike herniation of the entire bowel wall, whereas a pseudo diverticulum involves only a protrusion of the mucosa and submucosa through the muscularis propria of the colon (Fig. 353-1). The type of diverticulum affecting the colon is the pseudodiverticulum. Diverticula commonly affect the left and sigmoid colon; the rectum is always spared. However, in Asian populations, 70% of diverticula are seen in the right colon and cecum as well. *Diverticulitis* is inflammation of a diverticulum. Previous understanding of the pathogenesis of diverticulosis attributed a low-fiber diet as the sole culprit, and onset of diverticulitis would occur acutely when these diverticula become obstructed. However, evidence now suggests that the pathogenesis is more complex and multifactorial. The diverticula occur at the point where the nutrient artery, or *vasa recti*, penetrates through the muscularis propria, resulting in a break in the integrity of the colonic wall. This anatomic restriction may be a result of the relative high-pressure zone within the muscular sigmoid colon. Thus, higher-amplitude contractions combined with constipated, high-fat-content stool within the sigmoid lumen in an area of weakness in the colonic wall results in the creation of these diverticula. Consequently, the *vasa recti* is either compressed or eroded, leading to either perforation or bleeding. Chronic low-grade inflammation is thought to play a key role. Furthermore, better understanding of the gut microbiota suggests that dysbiosis is an important aspect of disease.

Presentation, Evaluation, and Management of Diverticular Bleeding

Hemorrhage from a colonic diverticulum is the most common cause of hematochezia in patients >60 years, yet only 20% of patients with diverticulosis will have gastrointestinal bleeding. Patients at increased risk for bleeding tend to be hypertensive, have atherosclerosis, and regularly use aspirin and nonsteroidal anti-inflammatory agents. Most bleeds are self-limited and stop spontaneously with bowel rest. The lifetime risk of rebleeding is 25%.

Initial localization of diverticular bleeding may include colonoscopy, multiplanar computed tomography (CT) angiogram, or nuclear medicine tagged red cell scan. If the patient is stable, ongoing bleeding is best managed by angiography. If mesenteric angiography can localize the bleeding site, the vessel can be occluded successfully with a coil in 80% of cases. The patient can then be followed closely with repetitive colonoscopy, if necessary, looking for evidence of colonic ischemia. Alternatively, a segmental resection of the colon can be undertaken to eliminate the risk of further bleeding. This may be advantageous in patients on chronic anticoagulation. However, with highly selective coil embolization, the rate of colonic ischemia is <10% and the risk of

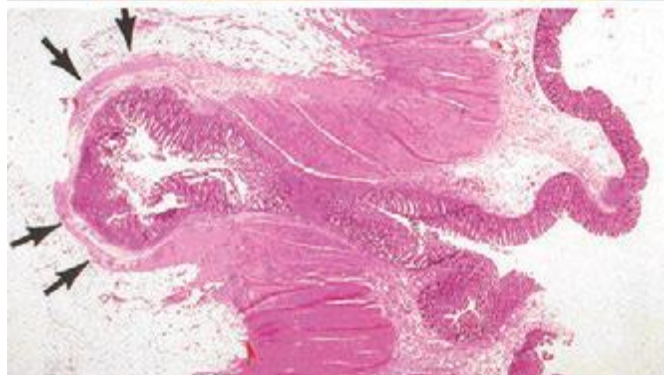


FIGURE 353-1 Gross and microscopic view of sigmoid diverticular disease. Arrows mark an inflamed diverticulum with the diverticular wall made up only of mucosa.

acute rebleeding is <25%. Long-term results (40 months) indicate that more than 50% of patients with acute diverticular bleeds treated with highly selective angiography have had definitive treatment. As another alternative, a selective infusion of vasopressin can be given to stop the hemorrhage, although this has been associated with significant complications, including myocardial infarction and intestinal ischemia. Furthermore, bleeding recurs in 50% of patients once the infusion is stopped.

If the patient is unstable or has had a 6-unit bleed within 24 h, current recommendations are that surgery should be performed. If the bleeding has been localized, a segmental resection can be performed. If the site of bleeding has not been definitively identified, a subtotal colectomy may be required. In patients without severe comorbidities, surgical resection can be performed with a primary anastomosis. A higher anastomotic leak rate has been reported in patients who received >10 units of blood.

Presentation, Evaluation, and Staging of Diverticulitis Acute uncomplicated diverticulitis characteristically presents with fever, anorexia, left lower quadrant abdominal pain, and obstipation (Table 353-1). In <25% of cases, patients may present with generalized peritonitis indicating the presence of a diverticular perforation. If a pericolonic abscess has formed, the patient may have abdominal distention and signs of localized peritonitis. Laboratory investigations will demonstrate a

Uncomplicated Diverticular Disease—75%

Abdominal pain
Fever
Leukocytosis
Anorexia/obstipation

Complicated Diverticular Disease—25%

Abscess 16%
Perforation 10%
Stricture 5%
Fistula 2%

leukocytosis. Rarely, a patient may present with an air-fluid level in the left lower quadrant on plain abdominal film. This is a giant diverticulum of the sigmoid colon and is managed with resection to avoid impending perforation.

The diagnosis of diverticulitis is best made on CT with the following findings: sigmoid diverticula, thickened colonic wall >4 mm, and inflammation within the pericolic fat ± the collection of contrast material or fluid. In 16% of patients, an abdominal abscess may be present. Symptoms of irritable bowel syndrome (Chap. 352) may mimic those of diverticulitis. Therefore, suspected diverticulitis that does not meet CT criteria or is not associated with a leukocytosis or fever is not diverticular disease. Other conditions that can mimic diverticular disease include an ovarian cyst, endometriosis, acute appendicitis, and pelvic inflammatory disease.

Although the benefit of colonoscopy in the evaluation of patients with diverticular disease has been called into question, its use is still considered important in the exclusion of colorectal cancer. The parallel epidemiology of colorectal cancer and diverticular disease provides enough concern for an endoscopic evaluation before operative management. Therefore, a colonoscopy should be performed ~6 weeks after an attack of diverticular disease.

Complicated diverticular disease is defined as diverticular disease associated with an abscess or perforation and less commonly with a fistula (Table 353-1). Perforated diverticular disease is staged using the Hinchey classification system (Fig. 353-2). This staging system was developed to predict outcomes following the surgical management of complicated diverticular disease. In complicated diverticular disease with fistula formation, common locations include cutaneous, vaginal, or vesicle fistulas. These conditions present with either passage of stool through the skin or vagina or the presence of air in the urinary stream (pneumaturia). Colovaginal fistulas are more common in women who have undergone a hysterectomy.

TREATMENT DIVERTICULAR DISEASE**MEDICAL MANAGEMENT**

Asymptomatic diverticular disease discovered on imaging studies or at the time of colonoscopy is best managed by diet alterations. Patients should be instructed to eat a fiber-enriched diet that includes 30 g of fiber each day. Supplementary fiber products such as Metamucil, Fibercon, or Citrucel are useful. The incidence of complicated diverticular disease appears to be increased in patients who smoke. Therefore, patients should be encouraged to refrain from smoking. The historical recommendation to avoid eating nuts is not based on more than anecdotal data.

Symptomatic uncomplicated diverticular disease with confirmation of inflammation and infection within the colon should be treated initially with antibiotics and bowel rest. Nearly 75% of patients hospitalized for acute diverticulitis will respond to nonoperative treatment with a suitable antimicrobial regimen. The current recommended antimicrobial coverage is trimethoprim/sulfamethoxazole or ciprofloxacin and metronidazole targeting aerobic gram-negative rods and anaerobic bacteria. Unfortunately, these agents do not cover enterococci, and the addition of ampicillin

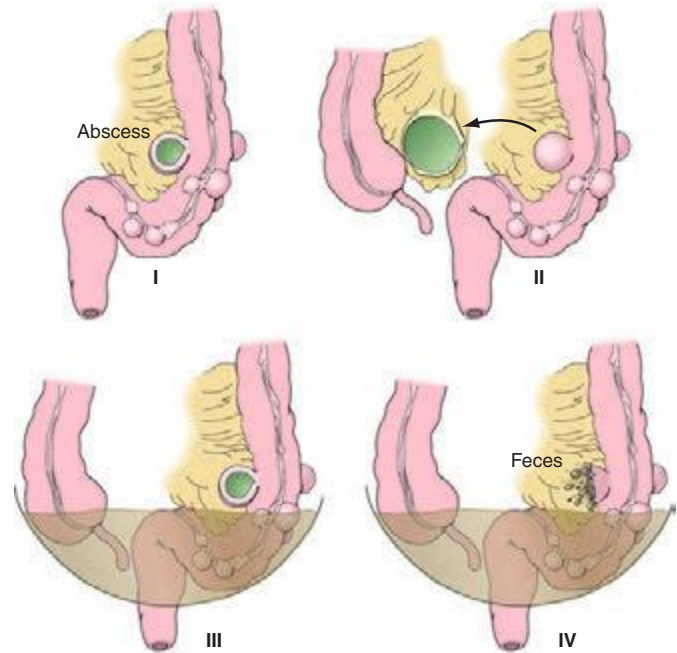


FIGURE 353-2 Hinchey classification of diverticulitis. Stage I: Perforated diverticulitis with a confined paracolic abscess. Stage II: Perforated diverticulitis that has closed spontaneously with distant abscess formation. Stage III: Noncommunicating perforated diverticulitis with fecal peritonitis (the diverticular neck is closed off, and therefore, contrast will not freely expel on radiographic images). Stage IV: Perforation and free communication with the peritoneum, resulting in fecal peritonitis.

to this regimen for nonresponders is recommended. Alternatively, single-agent therapy with a third-generation penicillin such as IV piperacillin or oral penicillin/clavulanic acid may be effective. The usual course of antibiotics is 7–10 days, although this length of time is being investigated. Patients should remain on a limited diet until their pain resolves.

Once the acute attack has resolved, the mainstay medical management of diverticular disease to prevent symptoms has evolved. Newer directions are targeted at colonic inflammation and dysbiosis. Diverticular disease is now considered a functional bowel disorder associated with low-grade inflammation. Therefore, the use of anti-inflammatory medications such as mesalazine has become popular. Patients treated with mesalazine have a decreased recurrence of symptomatic disease. Randomized trials of anti-inflammatory medications are ongoing.

Treatment strategies targeting dysbiosis in diverticular disease are also beneficial. Use of the polymerase chain reaction (PCR) on stool specimens from consumers of a high-fiber diet has shown different bacterial content than stool of consumers of a low-fiber, high-fat diet. Probiotics are being increasingly used by gastroenterologists for multiple bowel disorders and have been shown to prevent recurrence of diverticulitis. Specifically probiotics containing *Lactobacillus acidophilus* and *Bifidobacterium* strains have been shown to be beneficial. Furthermore, rifaximin (a poorly absorbed broad-spectrum antibiotic), when compared to fiber alone, is associated with 30% less frequent recurrent symptoms from uncomplicated diverticular disease.

SURGICAL MANAGEMENT

Preoperative risk factors influencing postoperative mortality rates include higher American Society of Anesthesiologists (ASA) physical status class (Table 353-2) and preexisting organ failure. In patients who are low risk (ASA P1 and P2), surgical therapy can be offered to those who do not rapidly improve on medical therapy. For uncomplicated diverticular disease, medical therapy can be continued beyond two attacks without an increased risk of perforation requiring a

TABLE 353-2 AMERICAN SOCIETY OF ANESTHESIOLOGISTS PHYSICAL STATUS CLASSIFICATION SYSTEM

P1	A normal healthy patient
P2	A patient with mild systemic disease
P3	A patient with severe systemic disease
P4	A patient with severe systemic disease that is a constant threat to life
P5	A moribund patient who is not expected to survive without the operation
P6	A declared brain-dead patient whose organs are being removed for donor purposes

colostomy. However, patients on immunosuppressive therapy, in chronic renal failure, or with a collagen-vascular disease have a fivefold greater risk of perforation during recurrent attacks. Surgical therapy is indicated in all low-surgical-risk patients with complicated diverticular disease.

The goals of surgical management of diverticular disease include controlling sepsis, eliminating complications such as fistula or obstruction, removing the diseased colonic segment, and restoring intestinal continuity. These goals must be obtained while minimizing morbidity rate, length of hospitalization, and cost in addition to maximizing survival and quality of life. **Table 353-3** lists the operations most commonly indicated based on the Hinchey classification and the predicted morbidity and mortality rates. Surgical objectives include removal of the diseased sigmoid down to the rectosigmoid junction. Failure to do this may result in recurrent disease. The current options for uncomplicated diverticular disease include an open sigmoid resection or a laparoscopic sigmoid resection. The benefits of laparoscopic resection over open surgical techniques include early discharge (by at least 1 day), less narcotic use, less postoperative complications, and an earlier return to work.

The options for the surgical management of complicated diverticular disease (**Fig. 353-3**) include the following: (1) proximal diversion of the fecal stream with an ileostomy or colostomy and sutured omental patch with drainage, (2) resection with colostomy and mucous fistula or closure of distal bowel with formation of a Hartmann's pouch, (3) resection with anastomosis (coloproctostomy), or (4) resection with anastomosis and diversion (coloproctostomy with loop ileostomy or colostomy). Laparoscopic techniques have been used for complicated diverticular disease; however, higher conversion rates to open techniques have been reported.

Patients with Hinchey stages I and II disease are managed with percutaneous drainage followed by resection with anastomosis about 6 weeks later. Current guidelines put forth by the American Society of Colon and Rectal Surgeons suggest, in addition to antibiotic therapy, CT-guided percutaneous drainage of diverticular abscesses that are greater than 3 cm and have a well-defined wall. Abscesses that are less than 3 cm may resolve with antibiotic therapy alone. Contraindications to percutaneous drainage are no percutaneous access route, pneumoperitoneum, and fecal peritonitis.

TABLE 353-3 OUTCOME FOLLOWING SURGICAL THERAPY FOR COMPLICATED DIVERTICULAR DISEASE

Hinchey Stage	Operative Procedure	Anastomotic Leak Rate, %	Overall Morbidity Rate, %
I	Resection with primary anastomosis without diverting stoma	3.8	22
II	Resection with primary anastomosis +/- diversion	3.8	30
III	Hartmann's procedure vs diverting colostomy and omental pedicle graft	—	0 vs. 6 mortality
IV	Hartmann's procedure vs diverting colostomy and omental pedicle graft	—	6 vs. 2 mortality

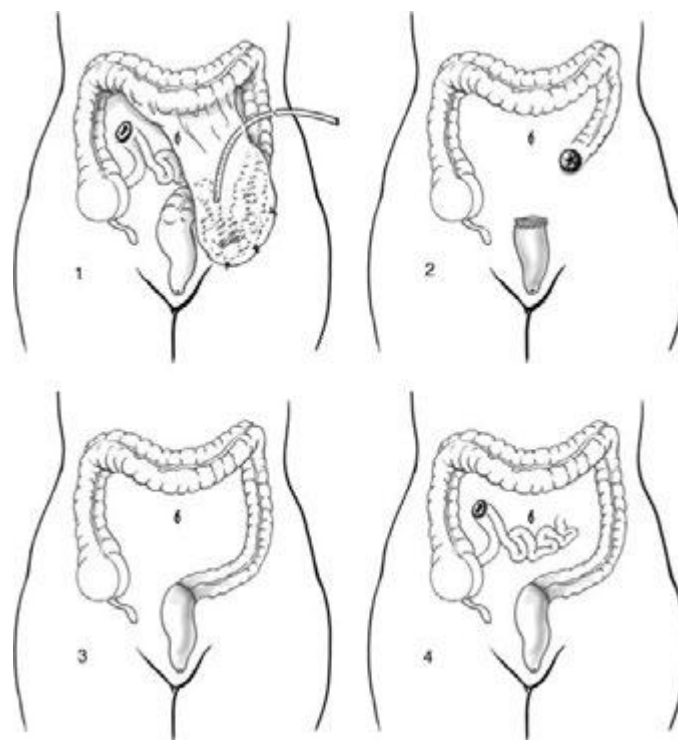


FIGURE 353-3 Methods of surgical management of complicated diverticular disease. (1) Drainage, omental pedicle graft, and proximal diversion. (2) Hartmann's procedure. (3) Sigmoid resection with coloproctostomy. (4) Sigmoid resection with coloproctostomy and proximal diversion.

Urgent operative intervention is undertaken if patients develop generalized peritonitis, and most will need to be managed with a Hartmann's procedure (resection of the sigmoid colon with end colostomy and rectal stump). In selected cases, nonoperative therapy may be considered. In one nonrandomized study, nonoperative management of isolated paracolic abscesses (Hinchey stage I) was associated with only a 20% recurrence rate at 2 years. More than 80% of patients with distant abscesses (Hinchey stage II) required surgical resection for recurrent symptoms.

Hinchey stage III disease is managed with a Hartmann's procedure or with primary anastomosis and proximal diversion. If the patient has significant comorbidities, making operative intervention risky, a limited procedure including intraoperative peritoneal lavage (irrigation), omental patch to the oversewn perforation, and proximal diversion of the fecal stream with either an ileostomy or transverse colostomy can be performed. No anastomosis of any type should be attempted in Hinchey stage IV disease. A limited approach to these patients is associated with a decreased mortality rate.

Recurrent Symptoms Recurrent abdominal symptoms following surgical resection for diverticular disease occur in 10% of patients. Recurrent diverticular disease develops in patients following inadequate surgical resection. A retained segment of diseased rectosigmoid colon is associated with twice the incidence of recurrence. The presence of irritable bowel syndrome may also cause recurrence of initial symptoms. Patients undergoing surgical resection for presumed diverticulitis and symptoms of chronic abdominal cramping and irregular loose bowel movements consistent with irritable bowel syndrome have poorer functional outcomes.

COMMON DISEASES OF THE ANORECTUM

RECTAL PROLAPSE (PROCIDENTIA)

Incidence and Epidemiology Rectal prolapse is six times more common in women than in men. The incidence of rectal prolapse peaks in women >60 years. Women with rectal prolapse have a higher incidence

1974 of associated pelvic floor disorders including urinary incontinence, rectocele, cystocele, and enterocele. About 20% of children with rectal prolapse will have cystic fibrosis. All children presenting with prolapse should undergo a sweat chloride test. Less common associations include Ehlers-Danlos syndrome, solitary rectal ulcer syndrome, congenital hypothyroidism, Hirschsprung's disease, dementia, mental retardation, and schizophrenia.

Anatomy and Pathophysiology Rectal prolapse (proctidentia) is a circumferential, full-thickness protrusion of the rectal wall through the anal orifice. It is often associated with a redundant sigmoid colon, pelvic laxity, and a deep rectovaginal septum (pouch of Douglas). Initially, rectal prolapse was felt to be the result of early internal rectal intussusception, which occurs in the upper to mid rectum. This was considered to be the first step in an inevitable progression to full-thickness external prolapse. However, only 1 of 38 patients with internal prolapse followed for >5 years developed full-thickness prolapse. Others have suggested that full-thickness prolapse is the result of damage to the nerve supply to the pelvic floor muscles or pudendal nerves from repeated stretching with straining to defecate. Damage to the pudendal nerves would weaken the pelvic floor muscles, including the external anal sphincter muscles. Bilateral pudendal nerve injury is more significantly associated with prolapse and incontinence than unilateral injury.

Presentation and Evaluation In external prolapse, the majority of patient complaints include anal mass, bleeding per rectum, and poor perianal hygiene. Prolapse of the rectum usually occurs following defecation and will spontaneously reduce or require the patient to manually reduce the prolapse. Constipation occurs in ~30–67% of patients with rectal prolapse. Differing degrees of fecal incontinence occur in 50–70% of patients. Patients with internal rectal prolapse will present with symptoms of both constipation and incontinence. Other associated findings include outlet obstruction (anismus) in 30%, colonic inertia in 10%, and solitary rectal ulcer syndrome in 12%.

Office evaluation is best performed after the patient has been given an enema, which enables the prolapse to protrude. An important distinction should be made between full-thickness rectal prolapse and isolated mucosal prolapse associated with hemorrhoidal disease (Fig. 353-4). Mucosal prolapse is known for radial grooves rather than circumferential folds around the anus and is due to increased laxity of the connective tissue between the submucosa and underlying muscle of the anal canal. The evaluation of prolapse should also include cystoproctography and colonoscopy. These examinations evaluate for associated pelvic floor disorders and rule out a malignancy or a polyp as the lead point for prolapse. If rectal prolapse is associated with chronic constipation, the patient should undergo a defecating proctogram and a sitzmark study. This will evaluate for the presence of anismus or colonic inertia. Anismus is the result of attempting to defecate against a closed pelvic floor and is also known as *nonrelaxing puborectalis*. This can be seen when straightening of the rectum fails to occur on fluoroscopy while the patient is attempting to defecate. In colonic inertia, a sitzmark study will demonstrate retention of >20% of markers on abdominal x-ray 5 days after swallowing. For patients with fecal incontinence, endoanal ultrasound and manometric evaluation, including pudendal nerve testing of their anal sphincter muscles, may be performed before surgery for prolapse (see “Fecal Incontinence,” below).

TREATMENT RECTAL PROLAPSE

The medical approach to the management of rectal prolapse is limited and includes stool-bulking agents or fiber supplementation to ease the process of evacuation. Surgical correction of rectal prolapse is the mainstay of therapy. Two approaches are commonly considered, transabdominal and transperineal. Transabdominal approaches have been associated with lower recurrence rates, but some patients with significant comorbidities are better served by a transperineal approach.

Common transperineal approaches include a transanal proctectomy (Altmeier procedure), mucosal proctectomy (Delorme

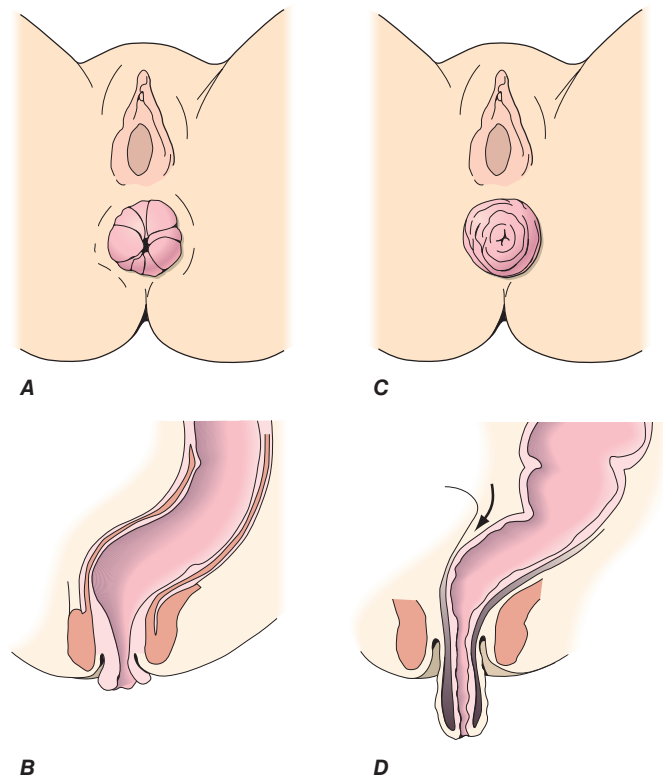


FIGURE 353-4 Degrees of rectal prolapse. Mucosal prolapse only (A, B, sagittal view). Full-thickness prolapse associated with redundant rectosigmoid and deep pouch of Douglas (C, D, sagittal view).

procedure), or placement of a Tirsch wire encircling the anus. The goal of the transperineal approach is to remove the redundant rectosigmoid colon. Common transabdominal approaches include presacral suture or mesh rectopexy (Ripstein) with (Frykman-Goldberg) or without resection of the redundant sigmoid. Colon resection, in general, is reserved for patients with constipation and outlet obstruction. Ventral rectopexy is an effective method of abdominal repair of full-thickness prolapse that does not require sigmoid resection (see description below). This repair may have improved functional results over other abdominal repairs. Transabdominal procedures can be performed effectively with laparoscopic and, more recently, robotic techniques without increased incidence of recurrence. The goal of the transabdominal approach is to restore normal anatomy by removing redundant bowel and reattaching the supportive tissue of the rectum to the presacral fascia. The final alternative is abdominal proctectomy with end-sigmoid colostomy. If total colonic inertia is present, as defined by a history of constipation and a positive sitzmark study, a subtotal colectomy with an ileosigmoid or rectal anastomosis may be required at the time of rectopexy.

Previously, the presence of internal rectal prolapse identified on imaging studies has been considered a nonsurgical disorder and biofeedback was recommended. However, only one-third of patients will have successful resolution of symptoms from biofeedback. Two surgical procedures more effective than biofeedback are the Stapled Transanal Rectal Resection (STARR) and the Laparoscopic Ventral Rectopexy (LVR). The STARR procedure (Fig. 353-5) is performed through the anus in patients with internal prolapse. A circular stapling device is inserted through the anus; the internal prolapse is identified and ligated with the stapling device. LVR (Fig. 353-6) is performed through an abdominal approach. An opening in the peritoneum is created on the left side of the rectosigmoid junction, and this opening continues down anterior on the rectum into the pouch of Douglas. No rectal mobilization is performed, thus avoiding any autonomic nerve injury. Mesh is secured to the anterior and lateral portion of the rectum, the vaginal fornix,

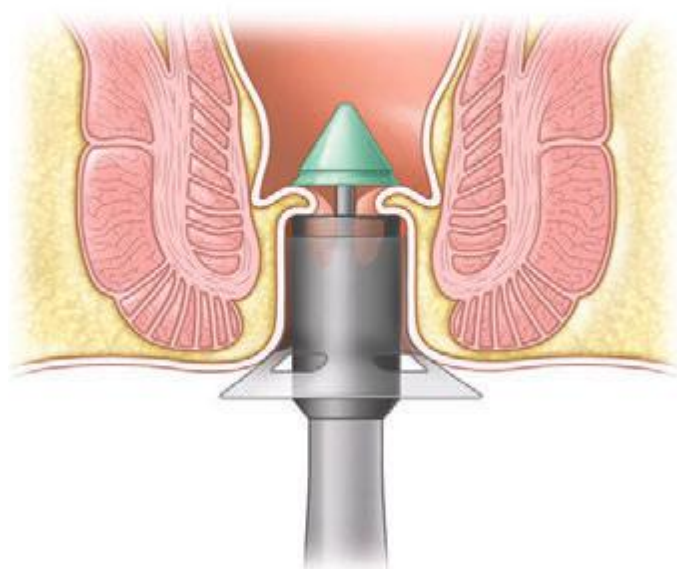


FIGURE 353-5 Stapled transanal rectal resection. Schematic of placement of the circular stapling device.

and the sacral promontory, allowing for closure of the rectovaginal septum and correction of the internal prolapse. In both procedures, recurrence at 1 year was low (<10%) and symptoms improved in more than three-fourths of patients.

FECAL INCONTINENCE

Incidence and Epidemiology Fecal incontinence is the involuntary passage of fecal material for at least 1 month in an individual with a developmental age of at least 4 years. The prevalence of fecal incontinence in the United States is 0.5–11%. The majority of patients are women and

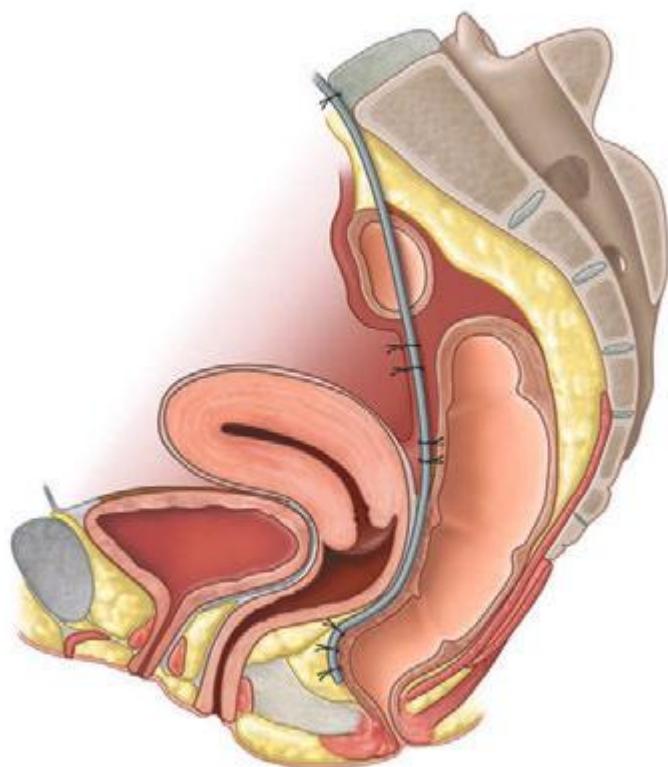


FIGURE 353-6 Laparoscopic ventral rectopexy (LVR). To reduce the internal prolapse and close any rectovaginal septal defect, the pouch of Douglas is opened and mesh is secured to the anterolateral rectum, vaginal fornix, and sacrum. (From A D'Hoore et al: *Br J Surg* 91:1500, 2004.)

TABLE 353-4 MEDICAL CONDITIONS THAT CONTRIBUTE TO SYMPTOMS OF FECAL INCONTINENCE

Neurologic Disorders

- Dementia
- Brain tumor
- Stroke
- Multiple sclerosis
- Tabes dorsalis
- Cauda equina lesions

Skeletal Muscle Disorders

- Myasthenia gravis
- Myopathies, muscular dystrophy

Miscellaneous

- Hypothyroidism
- Irritable bowel syndrome
- Diabetes
- Severe diarrhea
- Scleroderma

above the age of 65. A higher incidence of incontinence is seen among parous women. One-half of patients with fecal incontinence also suffer from urinary incontinence. The majority of incontinence is a result of obstetric injury to the pelvic floor, either while carrying a fetus or during the delivery. An anatomic sphincter defect may occur in up to 32% of women following childbirth regardless of visible damage to the perineum. Risk factors at the time of delivery include prolonged labor, the use of forceps, and the need for an episiotomy. Symptoms of incontinence can present after two or more decades following obstetric injury. Medical conditions known to contribute to the development of fecal incontinence are listed in [Table 353-4](#).

Anatomy and Pathophysiology The anal sphincter complex is made up of the internal and external anal sphincter. The internal sphincter is smooth muscle and a continuation of the circular fibers of the rectal wall. It is innervated by the intestinal myenteric plexus and is therefore not under voluntary control. The external anal sphincter is formed in continuation with the levator ani muscles and is under voluntary control. The pudendal nerve supplies motor innervation to the external anal sphincter. Obstetric injury may result in tearing of the muscle fibers anteriorly at the time of the delivery. This results in an obvious anterior defect on endoanal ultrasound. Injury may also be the result of stretching of the pudendal nerves during pregnancy or delivery of the fetus through the birth canal.

Presentation and Evaluation Patients may suffer with varying degrees of fecal incontinence. Minor incontinence includes incontinence to flatus and occasional seepage of liquid stool. Major incontinence is frequent inability to control solid waste. As a result of fecal incontinence, patients suffer from poor perianal hygiene. Beyond the immediate problems associated with fecal incontinence, these patients are often withdrawn and suffer from depression. For this reason, quality-of-life measures are an important component in the evaluation of patients with fecal incontinence.

The evaluation of fecal incontinence should include a thorough history and physical exam including digital rectal examination (DRE). Weak sphincter tone on DRE and loss of the “anal wink” reflex (S1-level control) may indicate a neurogenic dysfunction. Perianal scars may represent surgical injury. Other studies helpful in the diagnosis of fecal incontinence include anal manometry, pudendal nerve terminal motor latency (PNTML), and endoanal ultrasound. Centers that care for patients with fecal incontinence will have an anorectal physiology laboratory that uses standardized methods of evaluating anorectal physiology. Anorectal manometry (ARM) measures resting and squeeze pressures within the anal canal using an intraluminal water-perfused catheter. Current methods of ARM include use of a three-dimensional, high-resolution system with a 12-catheter perfusion system, which allows physiologic delineation of anatomic abnormalities.

1976 Pudendal nerve studies evaluate the function of the nerves innervating the anal canal using a finger electrode placed in the anal canal. Stretch injuries to these nerves will result in a delayed response of the sphincter muscle to a stimulus, indicating a prolonged latency. Finally, endoanal ultrasound will evaluate the extent of the injury to the sphincter muscles before surgical repair. Unfortunately, all of these investigations are user-dependent, and very few studies demonstrate that these studies predict outcome following an intervention. Magnetic resonance imaging (MRI) has been used, but its routine use for imaging in fecal incontinence is not well established.

Rarely does a pelvic floor disorder exist alone. The majority of patients with fecal incontinence will have some degree of urinary incontinence. Similarly, fecal incontinence is a part of the spectrum of pelvic organ prolapse. For this reason, patients may present with symptoms of obstructed defecation as well as fecal incontinence. Careful evaluation including dynamic MRI or cinedefecography should be performed to search for other associated defects. Surgical repair of incontinence without attention to other associated defects may decrease the success of the repair.

TREATMENT **FECAL INCONTINENCE**

Medical management of fecal incontinence includes strategies to bulk up the stool, which help in increasing fecal sensation. These include fiber supplementation, loperamide, diphenoxylate, and bile acid binders. These agents harden the stool and delay frequency of bowel movements and are helpful in patients with minimal to mild symptoms. Furthermore, patients can be offered a form of physical therapy called biofeedback. This therapy helps strengthen the external sphincter muscle while training the patient to relax with defecation to avoid unnecessary straining and further injury to the sphincter muscles. Biofeedback has had variable success and is dependent on the motivation of the patient. At a minimum, biofeedback is risk free and safe. Most patients will have some improvement. For this reason, it should be incorporated into the initial recommendation to all patients with fecal incontinence.

The “gold standard” for the treatment of fecal incontinence with an isolated sphincter defect has been the overlapping sphincteroplasty. The external anal sphincter muscle and scar tissue as well as any identifiable internal sphincter muscle are dissected free from the surrounding adipose and connective tissue and then an overlapping repair is performed in an attempt to rebuild the muscular ring and restore its function. Long-term results following overlapping sphincteroplasty show about a 50% failure rate over 5 years. Poorer outcome has been seen in patients with prolonged pudendal nerve terminal motor latency.

Sacral neuromodulation, collagen-enhancing injectables, radiofrequency therapy, and the artificial bowel sphincter are other options. Sacral nerve stimulation and the artificial bowel sphincter are both adaptations of procedures developed for the management of urinary incontinence. Sacral nerve stimulation is ideally suited for patients with intact but weak anal sphincters. A temporary nerve stimulator is placed on the third sacral nerve. If there is at least a 50% improvement in symptoms, a permanent nerve stimulator is placed under the skin. The artificial bowel sphincter is a cuff and reservoir apparatus that allows for manual inflation of a cuff placed around the anus, increasing anal tone. This allows the patient to manually close off the anal canal until defecation is necessary. Long-term results for sacral stimulation have been promising, with nearly 80% of patients having a reduction in incontinence episodes by at least 50%. This reduction has been sustainable in studies out to 5 years. Unfortunately, the artificial bowel sphincter has been associated with a 30% infection rate. Accordingly, implantation is performed less often.

Collagen-enhancing injectables have been around for several years. The largest open trial involved 115 incontinent patients treated with nonanimal stabilized hyaluronic acid (NASHA/DX) gel. In this study, patients underwent injections of NASHA/DX (Solesta)

into the anal mucosa and were followed for 12 months. The results were promising, with over 50% achieving greater than 50% reduction in incontinence episodes, and these results were sustainable up to 2 years. This method is another less invasive therapy for patients with fecal incontinence.

Radiofrequency energy delivery to the anal canal in patients with fecal incontinence aids in the development and restructuring of collagen fibers and provides tensile strength to the sphincter muscles. The radiofrequency is delivered as an office procedure with sedation. The results have been variable, with 20–50% of patients having a sustained reduction in incontinence episodes for 5 years.

Finally, the use of stem cells to increase the bulk of the sphincter muscles is currently being tested. Stem cells can be harvested from the patient’s own muscle, grown, and then implanted into their sphincter complex. Concern for cost and the need for an additional procedure dampen enthusiasm. Trial results are awaited.

HEMORRHOIDAL DISEASE

Incidence and Epidemiology Symptomatic hemorrhoids affect >1 million individuals in the Western world per year. The prevalence of hemorrhoidal disease is not selective for age or sex. However, age is known to be a risk factor. The prevalence of hemorrhoidal disease is less in underdeveloped countries. The typical low-fiber, high-fat Western diet is associated with constipation and straining and the development of symptomatic hemorrhoids.

Anatomy and Pathophysiology Hemorrhoidal cushions are a normal part of the anal canal. The vascular structures contained within this tissue aid in continence by preventing damage to the sphincter muscle. Three main hemorrhoidal complexes traverse the anal canal—the left lateral, the right anterior, and the right posterior. Engorgement and straining lead to prolapse of this tissue into the anal canal. Over time, the anatomic support system of the hemorrhoidal complex weakens, exposing this tissue to the outside of the anal canal where it is susceptible to injury. Hemorrhoids are commonly classified as external or internal. External hemorrhoids originate below the dentate line and are covered with squamous epithelium and are associated with an internal component. External hemorrhoids are painful when thrombosed. Internal hemorrhoids originate above the dentate line and are covered with mucosa and transitional zone epithelium and represent majority of hemorrhoids. The standard classification of hemorrhoidal disease is based on the progression of the disease from their normal internal location to the prolapsing external position ([Table 353-5](#)).

Presentation and Evaluation Patients commonly present to a physician for two reasons: bleeding and protrusion. Pain is less common than with fissures and, if present, is described as a dull ache from engorgement of the hemorrhoidal tissue. Severe pain may indicate a thrombosed hemorrhoid. Hemorrhoidal bleeding is described as painless

TABLE 353-5 THE STAGING AND TREATMENT OF HEMORRHOIDS

Stage	Description of Classification	Treatment
I	Enlargement with bleeding	Fiber supplementation Cortisone suppository Sclerotherapy
II	Protrusion with spontaneous reduction	Fiber supplementation Cortisone suppository
III	Protrusion requiring manual reduction	Fiber supplementation Cortisone suppository Banding Operative hemorrhoidectomy
IV	Irreducible protrusion	Fiber supplementation Cortisone suppository Operative hemorrhoidectomy

bright red blood seen either in the toilet or upon wiping. Occasional patients can present with significant bleeding, which may be a cause of anemia; however, the presence of a colonic neoplasm must be ruled out in anemic patients. Patients who present with a protruding mass complain about inability to maintain perianal hygiene and are often concerned about the presence of a malignancy.

The diagnosis of hemorrhoidal disease is made on physical examination. Inspection of the perianal region for evidence of thrombosis or excoriation is performed, followed by a careful digital examination. Anoscopy is performed paying particular attention to the known position of hemorrhoidal disease. The patient is asked to strain. If this is difficult for the patient, the maneuver can be performed while sitting on a toilet. The physician is notified when the tissue prolapses. It is important to differentiate the circumferential appearance of a full-thickness rectal prolapse from the radial nature of prolapsing hemorrhoids (see “Rectal Prolapse,” above). The stage and location of the hemorrhoidal complexes are defined.

TREATMENT HEMORRHOIDAL DISEASE

The treatment for bleeding hemorrhoids is based on the stage of the disease (Table 353-5). In all patients with bleeding, the possibility of other causes must be considered. In young patients without a family history of colorectal cancer, the hemorrhoidal disease may be treated first and a colonoscopic examination performed if the bleeding continues. Older patients who have not had colorectal cancer screening should undergo colonoscopy or flexible sigmoidoscopy.

With rare exceptions, the acutely thrombosed hemorrhoid can be excised within the first 72 h by performing an elliptical excision. Sitz baths, fiber, and stool softeners are prescribed. Additional therapy for bleeding hemorrhoids includes the office procedures of banding and sclerotherapy. Sensation begins at the dentate line; therefore, banding or sclerotherapy can be performed without discomfort in the office. Bands are placed around the engorged tissue, causing ischemia and fibrosis. This aids in fixing the tissue proximally in the anal canal. Patients may complain of a dull ache for 24 h following band application. During sclerotherapy, 1–2 mL of a sclerosant (usually sodium tetradecyl sulfate) is injected using a 25-gauge needle into the submucosa of the hemorrhoidal complex. Care must be taken not to inject the anal canal circumferentially, or stenosis may occur.

For surgical management of hemorrhoidal disease, excisional hemorrhoidectomy, transhemorrhoidal dearterialization (THD), or stapled hemorrhoidectomy (“the procedure for prolapse or hemorrhoids” [PPH]) is the procedure of choice. All surgical methods of management are equally effective in the treatment of symptomatic third- and fourth-degree hemorrhoids. However, because the sutured hemorrhoidectomy involves the removal of redundant tissue down to the anal verge, unpleasant anal skin tags are removed as well. The stapled hemorrhoidectomy is associated with less discomfort; however, this procedure does not remove anal skin tags. THD uses ultrasound guidance to ligate the blood supply to the anal tissue, hence reducing hemorrhoidal engorgement. No procedures on hemorrhoids should be done in patients who are immunocompromised or who have active proctitis. Furthermore, emergent hemorrhoidectomy for bleeding hemorrhoids is associated with a higher complication rate.

Acute complications associated with the treatment of hemorrhoids include pain, infection, recurrent bleeding, and urinary retention. Care should be taken to place bands properly and to avoid overhydration in patients undergoing operative hemorrhoidectomy. Late complications include fecal incontinence as a result of injury to the sphincter during the dissection. Anal stenosis may develop from overzealous excision, with loss of mucosal skin bridges for reepithelialization. Finally, an *ectropion* (prolapse of rectal mucosa from the anal canal) may develop. Patients with an ectropion complain of a “wet” anus as a result of inability to prevent soiling once the rectal mucosa is exposed below the dentate line.

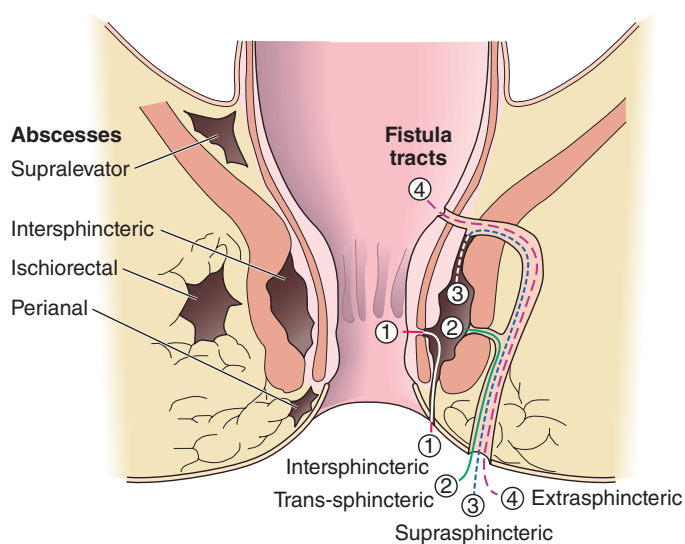


FIGURE 353-7 Common locations of anorectal abscess (left) and fistula in ano (right).

ANORECTAL ABSCESS

Incidence and Epidemiology The development of a perianal abscess is more common in men than women by a ratio of 3:1. The peak incidence is in the third to fifth decade of life. Perianal pain associated with the presence of an abscess accounts for 15% of office visits to a colorectal surgeon. The disease is more prevalent in immunocompromised patients such as those with diabetes, hematologic disorders, or inflammatory bowel disease and persons who are HIV positive. These disorders should be considered in patients with recurrent perianal infections.

Anatomy and Pathophysiology An anorectal abscess is an abnormal fluid-containing cavity in the anorectal region. Anorectal abscess results from an infection involving the glands surrounding the anal canal. Normally, these glands release mucus into the anal canal, which aids in defecation. When stool accidentally enters the anal glands, the glands become infected and an abscess develops. Anorectal abscesses are perianal in 40–50% of patients, ischiorectal in 20–25%, intersphincteric in 2–5%, and supralelevator in 2.5% (Fig. 353-7).

Presentation and Evaluation Perianal pain and fever are the hallmarks of an abscess. Patients may have difficulty voiding and have blood in the stool. A prostatic abscess may present with similar complaints, including dysuria. Patients with a prostatic abscess will often have a history of recurrent sexually transmitted diseases. On physical examination, a large fluctuant area is usually readily visible. Routine laboratory evaluation shows an elevated white blood cell count. Diagnostic procedures are rarely necessary unless evaluating a recurrent abscess. A CT scan or MRI has an accuracy of 80% in determining incomplete drainage. If there is a concern about the presence of inflammatory bowel disease, a rigid or flexible sigmoidoscopic examination may be done at the time of drainage to evaluate for inflammation within the rectosigmoid region. A more complete evaluation for Crohn’s disease would include a full colonoscopy and small-bowel series.

TREATMENT ANORECTAL ABSCESS

As with all abscesses, the “gold standard” is drainage. Office drainage of an uncomplicated anorectal abscess may suffice. A small incision close to the anal verge is made, and a Mallenkot drain is advanced into the abscess cavity. For patients who have a complicated abscess or who are diabetic or immunocompromised, drainage should be performed in an operating room under anesthesia. These patients are at greater risk for developing necrotizing fasciitis. There is limited role of antibiotics in management of anorectal abscesses. The antibiotics are only warranted in patients who are

1978 immunocompromised or have prosthetic heart valves, artificial joints, diabetes, or inflammatory bowel disease.

FISTULA IN ANO

Incidence and Epidemiology The incidence and prevalence of fistulating perianal disease parallels the incidence of anorectal abscess, estimating to be 1 in 10,000 individuals. Some 30–40% of abscesses will give rise to fistula in ano. Although the majority of the fistulas are cryptoglandular in origin, 10% are associated with IBD, tuberculosis, malignancy, and radiation.

Anatomy and Pathophysiology A fistula in ano is defined as a communication of an abscess cavity with an identifiable internal opening within the anal canal. This identifiable opening is most commonly located at the dentate line where the anal glands enter the anal canal. Patients experiencing continuous drainage following the treatment of a perianal abscess likely have a fistula in ano. These fistulas are classified by their relationship to the anal sphincter muscles, with 70% being intersphincteric, 23% transsphincteric, 5% suprasphincteric, and 2% extrasphincteric (Fig. 353-7).

Presentation and Evaluation A patient with a fistula in ano will complain of constant drainage from the perianal region associated with a firm mass. The drainage may increase with defecation. Perianal hygiene is difficult to maintain. Examination under anesthesia is the best way to evaluate a fistula. At the time of the examination, anoscopy is performed to look for an internal opening. Diluted hydrogen peroxide will aid in identifying such an opening. In lieu of anesthesia, MRI with an endoanal coil will also identify tracts in 80% of the cases. After drainage of an abscess with insertion of a Mallenkot catheter, a fistulagram through the catheter can be obtained in search of an occult fistula tract. Goodsall's rule states that a posterior external fistula will enter the anal canal in the posterior midline, whereas an anterior fistula will enter at the nearest crypt. A fistula exiting >3 cm from the anal verge may have a complicated upward extension and may not obey Goodsall's rule.

TREATMENT FISTULA IN ANO

A newly diagnosed draining fistula is best managed with placement of a seton, a vessel loop or silk tie placed through the fistula tract, which maintains the tract open and quiets down the surrounding inflammation that occurs from repeated blockage of the tract. Once the inflammation is less, the exact relationship of the fistula tract to the anal sphincters can be ascertained. A simple fistulotomy can be performed for intersphincteric and low (less than one-third of the muscle) transsphincteric fistulas without compromising continence. For a higher transsphincteric fistula, an anorectal advancement flap in combination with a drainage catheter or fibrin glue may be used. Very long (>2 cm) and narrow tracts respond better to fibrin glue than shorter tracts. Simple ligation of the internal fistula tract (LIFT procedure) has also been used in the management of simple fistula with good success.

Patients should be maintained on stool-bulking agents, nonnarcotic pain medication, and sitz baths following surgery for a fistula. Early complications from these procedures include urinary retention and bleeding. Later complications are rare (<10%) and include temporary and permanent incontinence. Recurrence is 0–18% following fistulotomy and 20–30% following anorectal advancement flap and the LIFT procedure,

ANAL FISSURE

Incidence and Epidemiology Anal fissures occur at all ages but are more common in the third through the fifth decades. A fissure is the most common cause of rectal bleeding in infancy. The prevalence is equal in males and females. It is associated with constipation, diarrhea, infectious etiologies, perianal trauma, and Crohn's disease.

Anatomy and Pathophysiology Trauma to the anal canal occurs following defecation. This injury occurs in the anterior or, more commonly, the posterior anal canal. Irritation caused by the trauma to the anal canal results in an increased resting pressure of the internal sphincter. The blood supply to the sphincter and anal mucosa enters laterally. Therefore, increased anal sphincter tone results in a relative ischemia in the region of the fissure and leads to poor healing of the anal injury. A fissure that is not in the posterior or anterior position should raise suspicion for other causes, including tuberculosis, syphilis, Crohn's disease, and malignancy.

Presentation and Evaluation A fissure can be easily diagnosed on history alone. The classic complaint is pain, which is strongly associated with defecation and is relentless. The bright red bleeding that can be associated with a fissure is less extensive than that associated with hemorrhoids. On examination, most fissures are located in either the posterior or anterior position. A lateral fissure is worrisome because it may have a less benign nature, and systemic disorders should be ruled out. A chronic fissure is indicated by the presence of a hypertrophied anal papilla at the proximal end of the fissure and a sentinel pile or skin tag at the distal end. Often the circular fibers of the hypertrophied internal sphincter are visible within the base of the fissure. If anal manometry is performed, elevation in anal resting pressure and a sawtooth deformity with paradoxical contractions of the sphincter muscles are pathognomonic.

TREATMENT ANAL FISSURE

The management of the acute fissure is conservative. Stool softeners for those with constipation, increased dietary fiber, topical anesthetics, glucocorticoids, and sitz baths are prescribed and will heal 60–90% of fissures. Chronic fissures are those present for >6 weeks. These can be treated with modalities aimed at decreasing the anal canal resting pressure including nifedipine or nitroglycerin ointment applied three times a day and botulinum toxin type A, up to 20 units, injected into the internal sphincter on each side of the fissure. Surgical management includes anal dilatation and lateral internal sphincterotomy. Usually, one-third of the internal sphincter muscle is divided; it is easily identified because it is hypertrophied. Recurrence rates from medical therapy are higher, but this is offset by a risk of incontinence following sphincterotomy. Lateral internal sphincterotomy may lead to incontinence more commonly in women.

ACKNOWLEDGMENT

We would like to thank Cory Sandore for providing some illustrations for this chapter. Gregory Bulkley, MD, contributed to this chapter in an earlier edition and some of that material has been retained here.

354 Mesenteric Vascular Insufficiency

Rizwan Ahmed, Mahmoud Malas

INTESTINAL ISCHEMIA

INCIDENCE AND EPIDEMIOLOGY

Intestinal ischemia occurs when splanchnic perfusion fails to meet the metabolic demands of the intestines, resulting in ischemic tissue injury. Mesenteric ischemia affects 2–3 people per 100,000, and the incidence of mesenteric ischemia is bound to increase in the aging population. Delay in diagnosis and management results in a high mortality, and prompt interventions may be life-saving. Intestinal ischemia is further classified based on etiology, which dictates management: (1) arterioocclusive mesenteric ischemia, (2) nonocclusive mesenteric ischemia, and (3) mesenteric venous thrombosis.

Risk factors for arterioocclusive mesenteric ischemia are generally acute in onset and include atrial fibrillation, recent myocardial infarction, valvular heart disease, and recent cardiac or vascular catheterization, all of which result in embolic clots reaching the mesenteric circulation. Nonocclusive mesenteric ischemia, also known as “intestinal angina,” is generally more insidious and most often seen in the aging population affected by atherosclerotic disease. Patients with chronic atherosclerotic disease could also suffer an acute insult from emboli leading to complete occlusion. Nonocclusive mesenteric ischemia is also seen in patients receiving high-dose vasopressor infusions, patients with cardiogenic or septic shock, and patients with cocaine overdose. Nonocclusive mesenteric ischemia is the most prevalent gastrointestinal disease complicating cardiovascular surgery. The incidence of ischemic colitis following elective aortic repair is 5–9%, and the incidence triples in patients following emergent repair. Mesenteric venous thrombosis is less common and is associated with the presence of a hypercoagulable state including protein C or S deficiency, antithrombin III deficiency, polycythemia vera, and carcinoma.

ANATOMY AND PATHOPHYSIOLOGY

The blood supply to the intestines is depicted in **Fig. 354-1**. To prevent ischemic injury, extensive collateralization occurs between major mesenteric trunks and branches of the mesenteric arcades. Collateral vessels within the small bowel are numerous and meet within the duodenum and the bed of the pancreas. Collateral vessels within the colon meet at the splenic flexure and descending/sigmoid colon. These areas, which are inherently at risk for decreased blood flow, are known as *Griffiths’ point* and *Sudeck’s point*, respectively, and are the most common locations for colonic ischemia (**Fig. 354-1**, shaded areas). The splanchnic circulation can receive up to 30% of the cardiac output. Protective responses to prevent intestinal ischemia include abundant

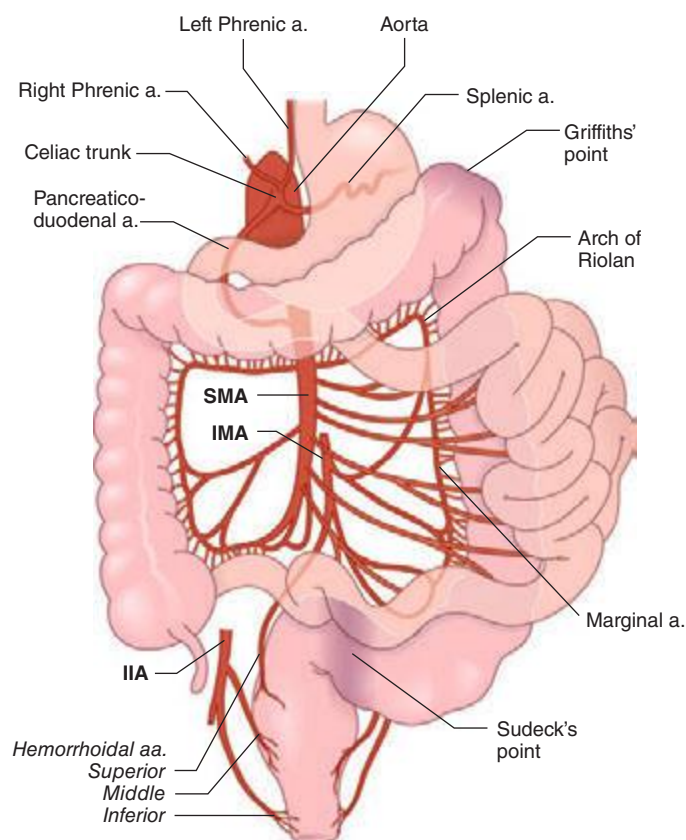


FIGURE 354-1 Blood supply to the intestines includes the celiac artery, superior mesenteric artery (SMA), inferior mesenteric artery (IMA), and branches of the internal iliac artery (IIA). Griffiths’ and Sudeck’s points, indicated by shaded areas, are watershed areas within the colonic blood supply and common locations for ischemia.

collateralization, autoregulation of blood flow, and the ability to increase oxygen extraction from the blood.

Occlusive ischemia is a result of disruption of blood flow by an embolus or progressive thrombosis in a major artery supplying the intestine. Emboli originate from the heart in more than 75% of cases and lodge preferentially in the superior mesenteric artery just distal to the origin of the middle colic artery. Progressive thrombosis of at least two of the major vessels supplying the intestine is required for the development of chronic intestinal angina. Nonocclusive ischemia is disproportionate mesenteric vasoconstriction (arteriolar vasospasm) in response to a severe physiologic stress such as shock. If left untreated, early mucosal stress ulceration will progress to full-thickness injury. Even in the early stages of ischemia, there is translocation of bacteria across the intestinal mucosa, resulting in bacteremia that can lead to sepsis.

PRESENTATION, EVALUATION, AND MANAGEMENT

Intestinal ischemia remains one of the most challenging diagnoses. The mortality rate is greater than 50%. The most significant indicator of survival is the timeliness of diagnosis and treatment. An overview of diagnosis and management of each form of intestinal ischemia is given in **Table 354-1**.

Acute mesenteric ischemia resulting from arterial embolus or thrombosis presents with severe acute, nonremitting abdominal pain strikingly out of proportion to the physical findings. Associated symptoms may include nausea and vomiting, transient diarrhea, anorexia, and bloody stools. With the exception of minimal abdominal distention and hypoactive bowel sounds, early abdominal examination is unimpressive. Later findings will demonstrate peritonitis and cardiovascular collapse. In the evaluation of acute intestinal ischemia, routine laboratory tests should be obtained, including complete blood count, serum chemistry, coagulation profile, arterial blood gas, amylase, lipase, lactic acid, blood type and cross match, and cardiac enzymes. Regardless of the need for urgent surgery, emergent admission to a monitored bed or intensive care unit is recommended for resuscitation and further evaluation. If the diagnosis of intestinal ischemia is being considered, consultation with a surgical service is necessary. Often the decision to operate is made on a high index of suspicion from the history and physical exam despite normal laboratory findings.

Other diagnostic modalities that may be useful in diagnosis but should not delay surgical therapy include electrocardiogram (ECG), echocardiogram, abdominal radiographs, computed tomography (CT), and mesenteric angiography. More recently, mesentery duplex scanning and visible light spectroscopy during colonoscopy have been demonstrated to be beneficial. The ECG may demonstrate an arrhythmia, indicating the possible source of the emboli. A plain abdominal film may show evidence of free intraperitoneal air, indicating a perforated viscus and the need for emergent exploration. Earlier features of intestinal ischemia seen on abdominal radiographs include bowel-wall edema, known as “thumbprinting.” If the ischemia progresses, air can be seen within the bowel wall (*pneumatosis intestinalis*) and within the portal venous system. Other features include calcifications of the aorta and its tributaries, indicating atherosclerotic disease. With the administration of oral and IV contrast, dynamic CT angiography with three-dimensional reconstruction is a highly sensitive test for intestinal ischemia. In acute embolic disease, mesenteric angiography is best performed intraoperatively. A mesenteric duplex scan demonstrating a high peak velocity of flow in the superior mesenteric artery (SMA) is associated with an approximately 80% positive predictive value of mesenteric ischemia. More significantly, a negative duplex scan virtually precludes the diagnosis of mesenteric ischemia. Duplex imaging serves as a screening test; further investigations with angiography are needed. The biggest limitation of duplex scanning is body habitus; in obese patients, imaging is poor yield. However, in patients with chronic disease, “food fear” often leads to a decreased appetite and therefore less abdominal fat, and duplex imaging is very high yield. The endoscopic techniques using visible light spectroscopy can be used in the diagnosis of chronic ischemia. When suspecting mesenteric ischemia involving the colon, performing an endoscopy to evaluate up to the splenic

TABLE 354-1 OVERVIEW OF THE MANAGEMENT OF ACUTE INTESTINAL ISCHEMIA

Condition	Key to Early Diagnosis	Treatment of Underlying Cause	Treatment of Specific Lesion	Treatment of Systemic Consequence
Arterioocclusive mesenteric ischemia	Computed tomography (CT) angiography	Anticoagulation	Laparotomy	Ensure hydration
1. Arterial embolus	Early laparotomy	Cardioversion	Embolectomy	Give antibiotics
		Proximal thrombectomy	Vascular bypass	Reverse acidosis
			Assess viability and resect dead bowel	Optimize oxygen delivery
				Avoid vasoconstrictors
2. Arterial thrombosis	Duplex ultrasound Angiography	Anticoagulation	Endovascular approach: thrombolysis, angioplasty and stenting	Give antibiotics
		Hydration	Endarterectomy/thrombectomy or vascular bypass	Reverse acidosis
			Assess viability and resect dead bowel	Optimize oxygen delivery
				Support cardiac output
				Avoid vasoconstrictors
Mesenteric venous thrombosis	Spiral CT	Anticoagulation	Anticoagulation ± laparotomy/thrombectomy/catheter-directed thrombolysis	Give antibiotics
Venous thrombosis	Angiography with venous phase	Massive hydration	Assess viability and resect dead bowel	Reverse acidosis
				Optimize oxygen delivery
				Support cardiac output
				Avoid vasoconstrictors
Nonocclusive mesenteric ischemia	Vasospasm: Angiography	Ensure hydration	Vasospasm	Ensure hydration
	Hypoperfusion: Spiral CT or colonoscopy	Support cardiac output	Intraarterial vasodilators	Give antibiotics
		Avoid vasoconstrictors	Hypoperfusion	Reverse acidosis
			Delayed laparotomy	Optimize oxygen delivery
			Assess viability and resect dead bowel	Support cardiac output
				Avoid vasoconstrictors

Source: Modified from GB Bulkley, in JL Cameron (ed): *Current Surgical Therapy*, 2nd ed. Toronto, BC Decker, 1986.

flexure is high yield. This is often an excellent diagnostic tool in patients with chronic renal insufficiency who cannot tolerate IV contrast.

The “gold standard” for the diagnosis of acute arterial occlusive disease is angiography, and management is laparotomy. Surgical exploration should not be delayed if suspicion of acute occlusive mesenteric ischemia is high or evidence of clinical deterioration or frank peritonitis is present. The goal of operative exploration is to resect compromised bowel and restore blood supply. The entire length of the small and large bowel beginning at the ligament of Treitz should be evaluated. The pattern of intestinal ischemia may indicate the level of arterial occlusion. In the case of SMA occlusion where the embolus usually lies just proximal to the origin of the middle colic artery, the proximal jejunum is often spared while the remainder of the small bowel to the transverse colon will be ischemic. The surgical management of acute mesenteric ischemia of the small bowel is embolectomy via arteriotomy; a small incision is made in the artery through which the clot is retrieved. Another way to manage acute thrombosis is thrombolysis therapy and angioplasty, with stent placement. However, this approach is more commonly applied to treat chronic mesenteric ischemia. If this is unsuccessful, a bypass from the aorta or iliac artery to the SMA is performed.

Nonocclusive or vasospastic mesenteric ischemia presents with generalized abdominal pain, anorexia, bloody stools, and abdominal distention. Often these patients are obtunded, and physical findings may not assist in the diagnosis. The presence of a leukocytosis, metabolic acidosis, elevated amylase or creatinine phosphokinase levels, and/or lactic acidosis is useful in support of the diagnosis of advanced intestinal ischemia; however, these markers may not be indicative of either reversible ischemia or frank necrosis. Investigational markers for intestinal ischemia include D-dimer, glutathione S-transferase, platelet-activating factor (PAF), and mucosal pH monitoring. Regardless of the need for urgent surgery, emergent admission to a monitored bed or intensive care unit is recommended for resuscitation and further evaluation. Early manifestations of intestinal ischemia include fluid sequestration within the bowel wall leading to a loss of interstitial volume. Aggressive fluid resuscitation may be necessary. To optimize oxygen delivery, nasal O₂ and blood transfusions may be given. Broad-spectrum antibiotics should be given to provide sufficient coverage for

enteric pathogens, including gram-negative and anaerobic organisms. Frequent monitoring of the patient’s vital signs, urine output, blood gases, and lactate levels is paramount, as is frequent abdominal examination. All vasoconstricting agents should be avoided; fluid resuscitation is the intervention of choice to maintain hemodynamics.

If ischemic colitis is a concern, colonoscopy should be performed to assess the integrity of the colon mucosa. Visualization of the rectosigmoid region may demonstrate decreased mucosal integrity, associated more commonly with nonocclusive mesenteric ischemia, or, on occasion, occlusive disease as a result of acute loss of inferior mesenteric arterial flow following aortic surgery. Ischemia of the colonic mucosa is graded as *mild* with minimal mucosal erythema or as *moderate* with pale mucosal ulcerations and evidence of extension to the muscular layer of the bowel wall. *Severe* ischemic colitis presents with severe ulcerations resulting in black or green discoloration of the mucosa, consistent with full-thickness bowel-wall necrosis. The degree of reversibility can be predicted from the mucosal findings: mild erythema is nearly 100% reversible, moderate is approximately 50% reversible, and frank necrosis is simply dead bowel. Follow-up colonoscopy can be performed to rule out progression of ischemic colitis.

Laparotomy for nonocclusive mesenteric ischemia is warranted for signs of peritonitis or worsening endoscopic findings and if the patient’s condition does not improve with aggressive resuscitation. Ischemic colitis is optimally treated with resection of the ischemic bowel and formation of a proximal stoma. Primary anastomosis should not be performed in patients with acute intestinal ischemia.

Patients with mesenteric venous thrombosis may present with a gradual or sudden onset. Symptoms include vague abdominal pain, nausea, and vomiting. Examination findings include abdominal distention with mild to moderate tenderness and signs of dehydration. The diagnosis of mesenteric thrombosis is frequently made on abdominal spiral CT with oral and IV contrast. Findings on CT angiography with venous phase include bowel-wall thickening and ascites. Intravenous contrast will demonstrate a delayed arterial phase and clot within the superior mesenteric vein. The goal of management is to optimize hemodynamics and correct electrolyte abnormalities with massive fluid resuscitation. Intravenous antibiotics as well as

355 Acute Intestinal Obstruction

Danny O. Jacobs

anticoagulation should be initiated. If laparotomy is performed and mesenteric venous thrombosis is suspected, heparin anticoagulation is immediately initiated, and compromised bowel is resected. Of all acute intestinal disorders, mesenteric venous insufficiency is associated with the best prognosis.

Chronic intestinal ischemia presents with intestinal angina or postprandial abdominal pain associated with need for increased blood flow to the intestine following meals. Patients report abdominal cramping and pain following ingestion of a meal. Weight loss and chronic diarrhea may also be noted. Abdominal pain without weight loss is not chronic mesenteric angina. Physical examination will often reveal a malnourished patient with an abdominal bruit as well as other manifestations of atherosclerosis. Duplex ultrasound evaluation of the mesenteric vessels has gained in popularity. It is important to perform the test fasting because the presence of increased bowel gas prevents adequate visualization of flow disturbances within the vessels or the lack of a vasodilation response to feeding during the test. This tool is frequently used as a screening test for patients with symptoms suggestive of chronic mesenteric ischemia. The gold standard for confirmation of mesenteric arterial occlusion is mesenteric angiography. Evaluation with mesenteric angiography allows for identification and possible intervention for the treatment of atherosclerosis within the vessel lumen and will also evaluate the patency of remaining mesenteric vessels. The use of mesenteric angiography may be limited in the presence of renal failure or contrast allergy. Magnetic resonance angiography is an alternative if the administration of contrast dye is contraindicated.

The management of chronic intestinal ischemia includes medical management of atherosclerotic disease by exercise, cessation of smoking, and antiplatelet and lipid-lowering medications. A full cardiac evaluation should be performed before intervention on chronic mesenteric ischemia. Newer endovascular procedures may avoid an operative intervention in selected patient populations. Angioplasty with endovascular stenting in the treatment of chronic mesenteric ischemia is associated with an 80% long-term success rate. In patients requiring surgical exploration, the approach used is determined by findings of the mesenteric angiogram. The entire length of the small and large bowel should be evaluated, beginning at the ligament of Treitz. Restoration of blood flow at the time of laparotomy is accomplished with mesenteric vessel endarterectomy or bypass.

Determination of intestinal viability intraoperatively in patients with suspected intestinal ischemia can be challenging. After revascularization, the bowel wall should be observed for return of a pink color and peristalsis. Palpation of major arterial mesenteric vessels can be performed, as well as applying a Doppler flowmeter to the antimesenteric border of the bowel wall, but neither is a definitive indicator of viability. In equivocal cases, 1 g of IV sodium fluorescein is administered, and the pattern of bowel reperfusion is observed under ultraviolet illumination with a standard (3600 Å) Wood's lamp. An area of nonfluorescence >5 mm in diameter suggests nonviability. If doubt persists, reexploration performed 24–48 h following surgery will allow demarcation of nonviable bowel. Primary intestinal anastomosis in patients with ischemic bowel is always worrisome; thus, delayed bowel reconstruction and reanastomosis should be deferred to the time of second-look laparotomy.

ACKNOWLEDGMENTS

We thank Cory Sandore for providing the illustration for this chapter. Susan Gearhart contributed to this chapter in the 18th edition.

EPIDEMIOLOGY

Morbidity and mortality from acute intestinal obstruction have been decreasing over the past several decades. Nevertheless, the diagnosis can still be challenging, and the type of complications that patients suffer has not changed significantly. The extent of mechanical obstruction is typically described as partial, high-grade, or complete—generally correlating with the risk of complications and the urgency with which the underlying disease process must be addressed. Obstruction is also commonly described as being either “simple” or, alternatively, “strangulated” if vascular insufficiency and intestinal ischemia are evident.

Acute intestinal obstruction occurs either *mechanically* from blockage or from intestinal dysmotility when there is no blockage. In the latter instance, the abnormality is described as being *functional*. Mechanical bowel obstruction may be caused by extrinsic processes, intrinsic abnormalities of the bowel wall, or intraluminal abnormalities (Table 355-1). Within each of these broad categories are many diseases that can impede intestinal propulsion. Intrinsic diseases that can cause intestinal obstruction are usually congenital, inflammatory, neoplastic, or traumatic in origin, although intussusception and radiation injury can also be etiologic. Primary small-bowel cancers rarely cause acute obstruction.

Acute intestinal obstruction accounts for approximately 1–3 % of all hospitalizations and a quarter of all urgent or emergent general surgery admissions. Approximately 80% of cases involve the small bowel, and about one-third of these patients show evidence of significant ischemia. The mortality rate for patients with strangulation who are operated on within 24–30 h of the onset of symptoms is approximately 8% but triples shortly thereafter.

Extrinsic diseases most commonly cause mechanical obstruction of the small intestine. In the United States and Europe, almost all cases are caused by postoperative adhesions (>50%), carcinomatosis, or herniation of the anterior abdominal wall. Carcinomatosis most often originates from the ovary, pancreas, stomach, or colon, although rarely, metastasis from distant organs like the breast and skin can occur. Adhesions are responsible for >90% of cases of early postoperative obstruction that require intervention.

Operations of the lower abdomen, including appendectomy and colorectal and gynecologic procedures, are especially likely to create

TABLE 355-1 MOST COMMON CAUSES OF ACUTE INTESTINAL OBSTRUCTION

Extrinsic Disease
Adhesions (especially due to previous abdominal surgery), internal or external hernias, neoplasms (including carcinomatosis and extraintestinal malignancies, mostly commonly ovarian), endometriosis or intraperitoneal abscesses, and idiopathic sclerosis
Intrinsic Disease
Congenital (e.g., malrotation, atresia, stenosis, intestinal duplication, cyst formation, and congenital bands—the latter rarely in adults)
Inflammation (e.g., inflammatory bowel disease, especially Crohn's disease, but also diverticulitis, radiation, tuberculosis, lymphogranuloma venereum, and schistosomiasis)
Neoplasia (note: primary small-bowel cancer is rare; obstructive colon cancer may mimic small-bowel obstruction if the ileocecal valve is incompetent)
Traumatic (e.g., hematoma formation, anastomotic strictures)
Other, including intussusception (where the lead point is typically a polyp or tumor in adults), volvulus, obstruction of duodenum by superior mesenteric artery, radiation or ischemic injury, and aganglionosis, which is Hirschsprung's disease
Intraluminal Abnormalities
Bezoars, feces, foreign bodies including inspissated barium, gallstones (entering the lumen via a cholecystoenteric fistula), enteroliths

TABLE 355-2 ACUTE SMALL-INTESTINAL AND COLONIC OBSTRUCTION INCIDENCES

Cause	Incidence
Postoperative adhesions	>50%
Neoplasms	~20%
Hernias (especially ventral or internal types, where the risk of strangulation is increased)	~10%
Inflammatory bowel disease, other inflammation (obstruction may resolve if acute inflammation and edema subside)	~5%
Intussusception, volvulus, other miscellaneous diseases	<15%

adhesions that can cause bowel obstruction (Table 355-2). Overall, small-bowel obstruction is slightly more common in women. The risk of internal herniation is increased by abdominal procedures such as laparoscopic or open Roux-en-Y gastric bypass. Although laparoscopic procedures may generate fewer postoperative adhesions compared with open surgery, the risk of obstructive adhesion formation is not eliminated.

In many patients who are successfully treated for adhesive small-bowel obstruction, obstruction will recur. The rate varies according to how patients were initially managed. Approximately 20% of patients who were treated conservatively and between 5 and 30% of patients who were managed operatively will require readmission within 10 years.

Volvulus, which occurs when bowel twists on its mesenteric axis, can cause partial or complete obstruction and vascular insufficiency. The sigmoid colon is most commonly affected, accounting for approximately two-thirds of all cases of volvulus and 4% of all cases of large-bowel obstruction. The cecum and terminal ileum can also volvulize, or the cecum alone may be involved as a cecal bascule. Risk factors include institutionalization, the presence of neuropsychiatric conditions requiring psychotropic medication, chronic constipation, and aging; patients typically present in their seventies or eighties. Colonic volvulus is more common in Eastern Europe, Russia, and Africa than it is in the United States. It is rare for adhesions or hernias to obstruct the colon. Cancer of the descending colon and rectum is responsible for approximately two-thirds of all cases, followed by diverticulitis and volvulus.

Functional obstruction, also known as *ileus* and *pseudo-obstruction*, is present when dysmotility prevents intestinal contents from being propelled distally and no mechanical blockage exists. Ileus that occurs after intraabdominal surgery is the most commonly identified form of functional bowel obstruction, although there are many other causes (Table 355-3). Although postoperative ileus is most often transient,

TABLE 355-3 MOST COMMON CAUSES OF ILEUS (FUNCTIONAL OR PSEUDO-OBSTRUCTION OF THE INTESTINE)

Intraabdominal procedures, lumbar spinal injuries, or surgical procedures on the lumbar spine and pelvis
Metabolic or electrolyte abnormalities, especially hypokalemia and hypomagnesemia, but also hyponatremia, uremia, and severe hyperglycemia
Drugs such as opiates, antihistamines, and some psychotropic (e.g., haloperidol, tricyclic antidepressants) and anticholinergic agents
Intestinal ischemia
Intraabdominal or retroperitoneal inflammation or hemorrhage
Lower lobe pneumonias
Intraoperative radiation (likely due to muscle damage)
Systemic sepsis
Hyperparathyroidism
Pseudo-obstruction (Ogilvie's syndrome)
Ileus secondary to hereditary or acquired visceral myopathies and neuropathies that disrupt myocellular neural coordination
Some collagen vascular diseases such as lupus erythematosus or scleroderma

it is the most common reason why hospital discharge is delayed. Pseudo-obstruction of the colon, also known as Ogilvie's syndrome, is a relatively rare disease. Some patients with Ogilvie's syndrome have colonic dysmotility due to abnormalities of their autonomic nervous system that may be inherited.

PATHOPHYSIOLOGY

The manifestations of acute intestinal obstruction depend on the nature of the underlying disease process, its location, and changes in blood flow (Fig. 355-1). Increased intestinal contractility, which occurs proximally and distal to the obstruction, is a characteristic response. Subsequently, intestinal peristalsis slows as the intestine or stomach proximal to the point of obstruction dilates and fills with gastrointestinal secretions and swallowed air. Although swallowed air is the primary contributor to intestinal distension, intraluminal air may also accumulate from fermentation, local carbon dioxide production, and altered gaseous diffusion.

Intraluminal dilation also increases intraluminal pressure. When luminal pressure exceeds venous pressure, venous and lymphatic drainage is impeded. Edema ensues, and the bowel wall proximal to the site of blockage may become hypoxic. Epithelial necrosis can be identified within 12 h of obstruction. Ultimately, arterial blood supply may become so compromised that full-thickness ischemia, necrosis, and perforation result. Stasis increases the bacteria counts within the jejunum and ileum. The most commonly cultured intraluminal organisms are *Escherichia coli*, *Streptococcus faecalis*, and *Klebsiella*, which may also be recovered from mesenteric lymph nodes and other more distant sites.

Other manifestations depend on the degree of hypovolemia, the patient's metabolic response, and the presence or absence of associated intestinal ischemia. Inflammatory edema eventually increases the production of reactive oxygen species and activates neutrophils and macrophages, which accumulate within the bowel wall. Their accumulation, along with changes in innate immunity, disrupts secretory and neuro-motor processes. Dehydration is caused by loss of the normal intestinal absorptive capacity as well as fluid accumulation in the gastric or intestinal wall and intraperitoneally.

Anorexia and emesis tend to exacerbate intravascular volume depletion. In the worst case scenario that is most commonly identified after distal obstruction, emesis leads to losses of gastric potassium, hydrogen, and chloride, while dehydration stimulates proximal renal tubule bicarbonate reabsorption. Intraperitoneal fluid accumulation, especially in patients with severe distal bowel obstruction, may increase intraabdominal pressure enough to elevate the diaphragm and inhibit respiration and to impede systemic venous return and promote vascular instability. Severe hemodynamic compromise may elicit a systemic inflammatory response and generalized microvascular leakage.

Closed-loop obstruction results when the proximal and distal openings of a given bowel segment are both occluded, e.g., due to volvulus or a hernia. It is the most common precursor for strangulation, but not every closed loop strangulates. The risk of vascular insufficiency, systemic inflammation, hemodynamic compromise, and irreversible intestinal ischemia is much greater in patients with closed-loop obstruction. Pathologic changes may occur more rapidly, and emergency intervention is indicated. Irreversible bowel ischemia progresses to transmural necrosis even if the obstruction is relieved. It is also important to remember that patients with high-grade distal colonic obstruction who have competent ileocecal valves may present with closed-loop obstruction. In the latter instance, the cecum may progressively dilate such that ischemic necrosis results in cecal perforation. This risk is generally greatest when the cecal diameter exceeds 12 cm, as informed by Laplace's law. Patients with distal colonic obstruction whose ileocecal valves are incompetent tend to present later in the course of disease and mimic patients with distal small-bowel obstruction.

HISTORY AND PHYSICAL FINDINGS

Even though the presenting signs and symptoms can be misleading, many patients with acute obstruction can be accurately diagnosed after a thorough history and physical examination is performed. Early

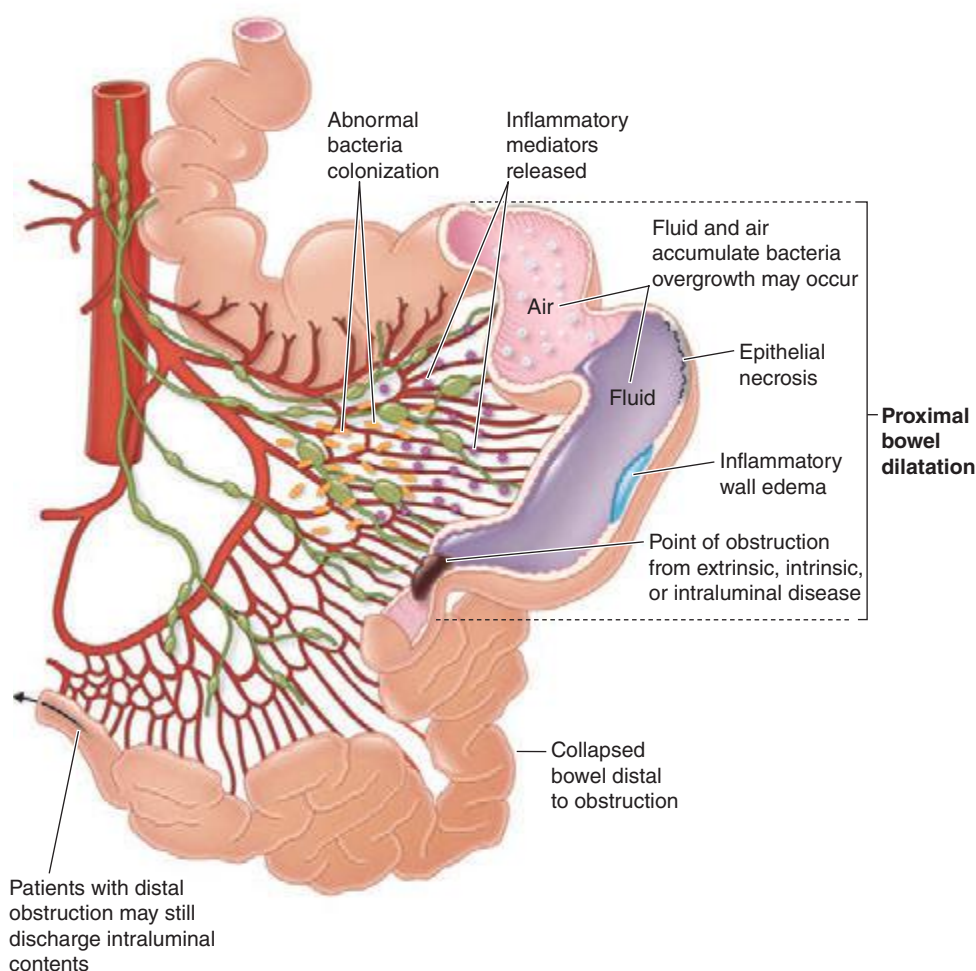


FIGURE 355-1 Pathophysiologic changes of small-bowel obstruction.

recognition allows earlier treatment that decreases the risk of progression or other excess morbidity. Small-bowel obstruction with strangulation can be especially difficult to diagnosis promptly.

The cardinal signs are colicky abdominal pain, abdominal distention, emesis, and obstipation. More intraluminal fluid accumulates in patients with distal obstruction, which typically leads to greater distention, more discomfort, and delayed emesis. This emesis is feculent when there is bacterial overgrowth. Patients with more proximal obstruction commonly present with less abdominal distention but more pronounced vomiting. Elements of the history that might be helpful include any prior history of surgery, including herniorrhaphy, as well as any history of cancer or inflammatory bowel disease.

Most patients, even with simple obstruction, appear to be critically ill. Many may be oliguric, hypotensive, and tachycardic because of severe intravascular volume depletion. Fever is worrisome for strangulation or systemic inflammatory changes. Bowel sounds and bowel functional activity are notoriously difficult to interpret. Classically, many patients with early small-bowel obstruction will have high-pitched, “musical” tinkling bowel sounds and peristaltic “rushes” known as borborygmi. Later in the course of disease, the bowel sounds may be absent or hypoactive as peristaltic activity decreases. This is in contrast to the common findings in patients with ileus or pseudo-obstruction where bowel sounds are typically absent or hypoactive from the beginning. Lastly, patients with partial blockage may continue to pass flatus and stool, and those with complete blockage may evacuate bowel contents present downstream beyond their obstruction.

All surgical incisions should be examined. The presence of a tender abdominal or groin mass strongly suggests that an incarcerated hernia may be the cause of obstruction. The presence of tenderness

should increase the concern about the presence of complications such as ischemia, necrosis, or peritonitis. Severe pain with localization or signs of peritoneal irritation is suspicious for strangulated or closed-loop obstruction. It is important to remember that the discomfort may be out of proportion to physical findings mimicking the complaints of patients with acute mesenteric ischemia. Every patient should have a rectal examination. Patients with colonic volvulus present with the classic manifestations of closed-loop obstruction: severe abdominal pain, vomiting, and obstipation. Asymmetrical abdominal distension and a tympanic mass may be evident.

Patients with ileus or pseudo-obstruction may have signs and symptoms similar to those of bowel obstruction. Although abdominal distention is present, colicky abdominal pain is typically absent, and patients may not have nausea or emesis. Ongoing, regular discharge of stool or flatus can sometimes help distinguish patients with ileus from those with complete mechanical bowel obstruction.

LABORATORY AND IMAGING STUDIES

Laboratory testing should include a complete blood count and serum electrolyte and creatinine measurements. Serial assessments are often useful. Mild hemoconcentration and slight elevation of the white blood cell count commonly occur after simple bowel obstruction. Emesis and dehydration may cause hypokalemia, hypochloremia, elevated blood urea nitrogen-to-creatinine ratios, and metabolic alkalosis. Patients may be hyponatremic on admission because many have attempted to rehydrate themselves with hypotonic fluids. The presence of guaiac-positive stools and iron-deficiency anemia are strongly suggestive of malignancy.

Higher white blood cell counts with the presence of immature forms or the presence of metabolic acidosis are worrisome for severe volume

1984 depletion or ischemic necrosis and sepsis. At this time, there are no laboratory tests that are especially useful for identifying the presence of simple or strangulated obstruction, although increases in serum D-lactate, creatine kinase bb isoenzymes, or intestinal fatty acid binding protein levels may be suggestive of the latter.

In all cases, when considering diagnostic imaging, the key is not to delay surgical consultation and operative intervention when the patient's signs or symptoms strongly suggest that high-grade or complete obstruction or bowel compromise is present. Plain films of the abdomen, which must include upright or cross-table lateral views, can be completed quickly and may confirm the clinical suspicion 60% of the time. Interpretation immediately after operation is difficult. A "staircasing" pattern of dilated air and fluid-filled small-bowel loops >2.5 cm in diameter with little or no air seen in the colon are classical findings in patients with small-bowel obstruction, although findings may be equivocal in some patients with documented disease. Little bowel gas appears in patients with proximal bowel obstruction or in patients whose intestinal lumens are filled with fluid. Upright plain films of the abdomen of patients with large-bowel obstruction typically show colon dilatation. Small-bowel air-fluid levels will not be obvious if the ileocecal valve is competent. Although it can be difficult to distinguish from ileus, small-bowel obstruction is more likely when air-fluid levels are seen without significant colonic distension. Free air suggests that perforation has occurred in patients who have not recently undergone surgical procedures. Radiopaque foreign bodies or enteroliths may be visualized. A gas-filled, "coffee bean"-shaped dilated shadow may be seen in patients with volvulus.

More sophisticated imaging can be beneficial when the diagnosis is unclear. Magnetic resonance imaging has been used to diagnose small-bowel obstruction, but it is more expensive and, typically, provides less spatial resolution. Ultrasonographic evaluations are especially difficult to interpret but may be sensitive and appropriate studies to evaluate patients who are pregnant or for whom x-ray exposure is otherwise contraindicated or inappropriate.

Computed tomography (CT) is the most commonly used imaging modality. Its sensitivity for detecting bowel obstruction is approximately 95% (78–100%) in patients with high-grade obstruction, with a specificity of 96% and an accuracy of $\geq 95\%$. Its accuracy in diagnosing closed-loop obstruction is much lower (60%). Examples of some CT images are reproduced in Fig. 355-2. It may also provide useful information regarding location or identify particular circumstances where surgical intervention is needed urgently. Patients who have evidence of contrast appearing within the cecum within 4–24 h of oral administration can be expected to improve with high sensitivity and specificity (~95% each). For example, contrast studies may demonstrate a "bird's beak," a "c-loop," or "whorl" deformity on CT imaging at the site where twisting obstructs the lumen when a colonic volvulus is present.

CT imaging with enteral and IV contrast can also identify ischemia. Altered bowel wall enhancement is the most specific early finding, but its sensitivity is low. Mesenteric venous gas, pneumoperitoneum, and pneumatosis intestinalis are late findings indicating the presence of bowel necrosis. CT scanning after a water-soluble contrast enema may help distinguish ileus or pseudo-obstruction from distal large-bowel obstruction in patients who present with evidence of small-bowel and colonic distention. CT enteroclysis can accurately identify neoplasia as a cause of bowel obstruction. Contrast enemas or colonoscopies are almost always needed to identify causes of acute colonic obstruction.

Barium studies are generally contraindicated in patients with firm evidence of complete or high-grade bowel obstruction, especially when they present acutely. Barium should never be given orally to a patient with possible obstruction until that diagnosis has been excluded. In every other case, such investigations should only be performed in exceptional circumstances and with great caution because patients with significant obstruction may develop barium concretions as an additional source of blockage and some who would have otherwise recovered will require operative intervention. Barium opacification also renders cross-sectional imaging studies or angiography uninterpretable.

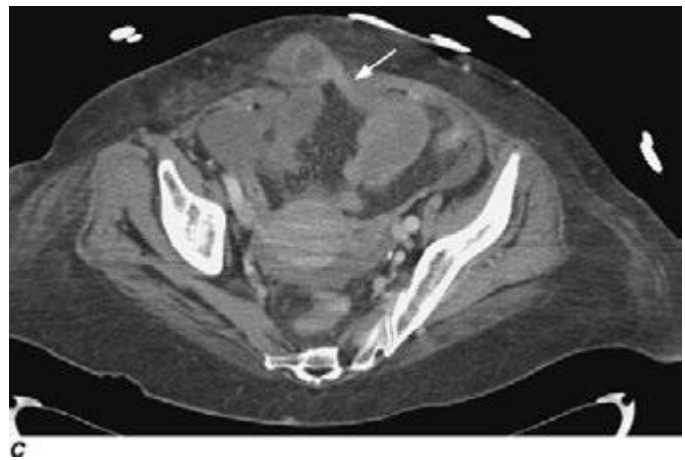
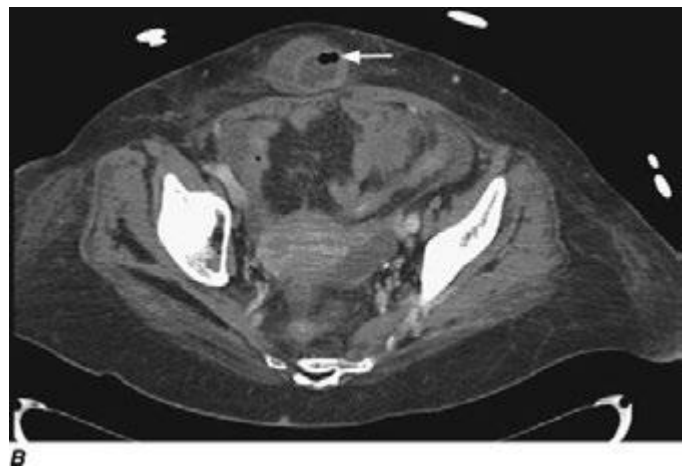
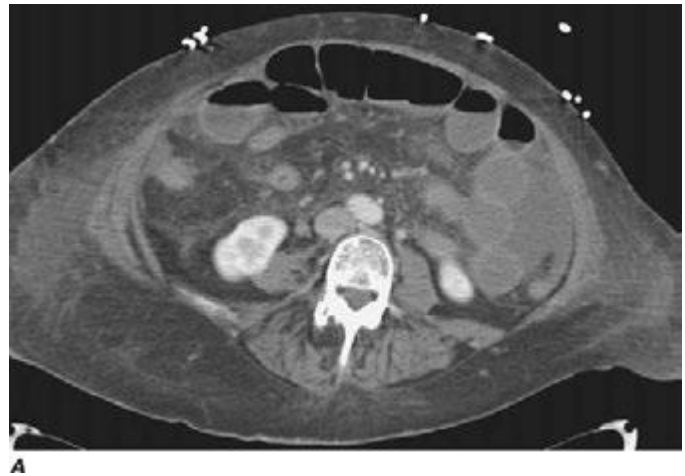


FIGURE 355-2 Computed tomography with oral and intravenous contrast demonstrating (A) evidence of small-bowel dilatation with air-fluid levels consistent with a small-bowel obstruction; (B) a partial small-bowel obstruction from an incarcerated ventral hernia (arrow); and (C) decompressed bowel seen distal to the hernia (arrow). (From W Silen: *Acute intestinal obstruction*, in DL Longo et al [eds]: *Harrison's Principles of Internal Medicine*, 18th ed. New York, McGraw-Hill, 2012.)

TREATMENT ACUTE INTESTINAL OBSTRUCTION

An improved understanding of the pathophysiology of bowel obstruction and the importance of fluid resuscitation, electrolyte repletion, intestinal decompression, and the selected use of antibiotics have likely contributed to a reduction in the mortality from acute bowel obstruction. Every patient should be stabilized as quickly

as possible. Nasogastric tube suction decompresses the stomach, minimizes further distention from swallowed air, improves patient comfort, and reduces the risk of aspiration. Urine output should be assessed using a Foley catheter. In some cases, for example, in patients with cardiac disease, central venous pressures should be monitored. The use of antibiotics is controversial, although prophylactic administration is warranted if surgery is required. Complete bowel obstruction is an indication for intervention. Stenting may be possible and warranted for some patients with high-grade obstruction due to unresectable stage IV malignancy. Stenting may also allow elective mechanical bowel preparation before surgery is undertaken. Because treatment options are so variable, it is helpful to make as precise a diagnosis as possible preoperatively.

ILEUS

Patients with ileus are treated supportively with intravenous fluids and nasogastric decompression while any underlying pathology is treated. Pharmacologic therapy is not yet proven to be efficacious or cost-effective. However, peripherally active μ -opioid receptor antagonists (e.g., alvimopan and methylnaltrexone) may accelerate gastrointestinal recovery in some patients who have undergone abdominal surgery.

COLONIC PSEUDO-OBSTRUCTION (OGILVIE'S DISEASE)

Neostigmine is an acetylcholinesterase inhibitor that increases cholinergic (parasympathetic) activity, which can stimulate colonic motility. Some studies have shown it to be moderately effective in alleviating acute colonic pseudo-obstruction. It is the most common therapeutic approach and can be used once it is certain that there is no mechanical obstruction. Cardiac monitoring is required, and atropine should be immediately available. Intravenous administration induces defecation and flatus within 10 min in the majority of patients who will respond. Sympathetic blockade by epidural anesthesia can successfully ameliorate pseudo-obstruction in some patients.

VOLVULUS

Patients with sigmoid volvulus can often be decompressed using a flexible tube inserted through a rigid proctoscope or using a flexible sigmoidoscope. Successful decompression results in sudden release of gas and fluid with evidence of decreased abdominal distension and allows definitive correction to be scheduled electively. Cecal volvulus most often requires laparotomy or laparoscopic correction.

INTRAOPERATIVE STRATEGIES

Approximately 60–80% of selected patients with mechanical bowel obstruction can be successfully treated conservatively. Indeed, most cases of radiation-induced obstruction should also be managed nonoperatively if possible. In most circumstances, early consultation with a general surgeon is prudent when there is concern about strangulation obstruction or other abnormality that needs to be addressed urgently. Deterioration signifies a need for intervention. At this time, the decision as to whether the patient can continue to be treated nonoperatively can only be based on clinical judgment, although, as described earlier, imaging studies can sometimes be helpful. The frequency of major complications after operation ranges from 12 to 47%, with greater risk being attributed to resection therapies and the patient's overall health. Risk is increased for patients with American Society of Anesthesiologists (ASA) class III or higher.

At operation, dilation proximal to the site of blockage with distal collapse is a defining feature of bowel obstruction. Intraoperative strategies depend on the underlying problem and range from lysis of adhesions to resection with or without diverting ostomy to primary resection with anastomosis. Resection is warranted when there is concern about the bowel's viability after the obstructive process is relieved. Laparoscopic approaches can be useful for patients with early obstruction when extensive adhesions are not expected to be present. Some patients with high-grade obstruction

secondary to malignant disease that is not amendable to resection will benefit from bypass procedures.

ADULT INTUSSUSCEPTION AND GALLSTONE ILEUS

Primary resection is prudent. Careful manual reduction of any involved bowel may limit the amount of intestine that needs to be removed. A proximal ostomy may be required if unprepped colon is involved. Only 60% of patients with gallstone ileus obstruct in the ileum. The most common site of intestinal obstruction in patients with gallstone "ileus" is the ileum (60% of patients). The gallstone enters the intestinal tract most often via a cholecystoduodenal fistula. It can usually be removed by operative enterolithotomy. Addressing the gallbladder disease during urgent or emergent surgery is not recommended.

POSTOPERATIVE BOWEL OBSTRUCTION

Early postoperative mechanical bowel obstruction is that which occurs within the first 6 weeks of operation. Most are partial and can be expected to resolve spontaneously. It tends to respond and behave differently from classic mechanical bowel obstruction and may be very difficult to distinguish from postoperative ileus. A higher index of suspicion for a definitive site of obstruction is warranted for patients who undergo laparoscopic surgical procedures. Patients who first had ileus and then subsequently develop obstructive symptoms after an initial return of normal bowel function are more likely to have true postoperative small-bowel obstruction. The longer it takes for a patient's obstructive symptoms to resolve after hospitalization, the more likely the patient is to require surgical intervention.

ACKNOWLEDGMENT

The wisdom and expertise of Dr. William Silen are gratefully acknowledged.

356

Acute Appendicitis and Peritonitis

Danny O. Jacobs

ACUTE APPENDICITIS

INCIDENCE AND EPIDEMIOLOGY

Appendicitis occurs more frequently in Westernized societies. Although its incidence is decreasing for uncertain reasons, acute appendicitis remains the most common emergency general surgical disease affecting the abdomen, with a rate of approximately 100 per 100,000 person-years in Europe and the Americas or about 11 cases per 10,000 people annually. Approximately 9% of men and 7% of women will experience an episode during their lifetime. Appendicitis occurs most commonly in 10- to 19-year-olds, although the average age at diagnosis appears to be gradually increasing, as is the frequency of the disease in African Americans, Asians, and Native Americans. Overall, 70% of patients are less than 30 years old and most are men; the male-to-female ratio is 1.4:1.

One of the more common complications and most important causes of excess morbidity and mortality is perforation, whether it is contained and localized or unconstrained within the peritoneal cavity. In contrast to the trend observed for appendicitis and appendectomy, the incidence of perforated appendicitis (~20 cases per 100,000 person-years) is increasing. The explanation for this phenomenon is unknown. Approximately 20% of all patients have evidence of perforation at presentation, but the percentage risk is much higher in patients under 5 or over 65 years of age.

Appendicitis was first described in 1886 by Reginald Fitz. Its etiology is still not completely understood. Fecaliths, incompletely digested food residue, lymphoid hyperplasia, intraluminal scarring, tumors, bacteria, viruses, and inflammatory bowel disease have all been associated with inflammation of the appendix and appendicitis.

Although not proven, obstruction of the appendiceal lumen is believed to be an important step in the development of appendicitis. In some cases, obstruction leads to bacterial overgrowth and luminal distension, with an increase in intraluminal pressure that can inhibit the flow of lymph and blood in some cases. Then, vascular thrombosis and ischemic necrosis with perforation of the distal appendix may occur. Any perforation that occurs near the base of the appendix should raise concerns about another disease process. Most patients who will perforate do so before they are evaluated by surgeons.

Appendiceal fecaliths (or appendicoliths) are found in approximately 50% of patients with gangrenous appendicitis who perforate but are rarely identified in those who have simple disease. As mentioned earlier, the incidence of perforated, but not simple, appendicitis is increasing. The rate of perforated and nonperforated appendicitis is correlated in men but not in women. Together these observations suggest that the underlying pathophysiologic processes are different and that simple appendicitis does not always progress to perforation. Furthermore, some cases of simple acute appendicitis may resolve spontaneously or with antibiotic therapy, and recurrent disease is remotely possible. The relative frequency of these events is unknown.

When perforation occurs, the resultant leak may be contained by the omentum or other surrounding tissues to form an abscess. Free perforation normally causes severe peritonitis. These patients may also develop infective suppurative thrombosis of the portal vein and its tributaries along with intrahepatic abscesses. The prognosis of the very unfortunate patients who develop this dreaded complication is very poor.

CLINICAL MANIFESTATIONS

More refined approaches to diagnosis, supportive care, and surgical intervention are likely responsible for the remarkable decrease in the risk of mortality from simple appendicitis to currently less than 1%. Nevertheless, it is still important to identify patients who might have appendicitis as early as possible to minimize their risk of developing complications. Patients who have had symptoms for more than 48 h are more likely to perforate.

Appendicitis should be included in the differential diagnosis of abdominal pain for every patient in any age group unless it is certain that the organ has been previously removed (Table 356-1).

The appendix's anatomical location, which varies, directly influences how the patient presents for care. Where the appendix can be "found" ranges from local differences in how the appendiceal body and tip lie relative to its attachment to the cecum (Figs. 356-1 and 356-2), to where the appendix is actually situated in the peritoneal cavity—for example, from its typical location in the right lower quadrant, to the pelvis, right flank, right upper quadrant (as may be observed during

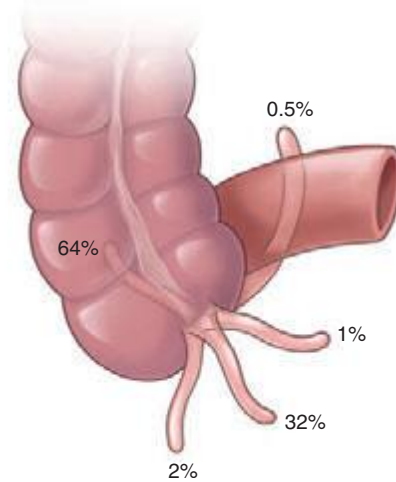


FIGURE 356-1 Regional anatomical variations of the appendix.

pregnancy), or even the left side of the abdomen for patients with malrotation or who have severely redundant colons.

Because the differential diagnosis of appendicitis is so extensive, deciding if a patient has appendicitis can be difficult (Table 356-2). Soliciting an appropriate history requires detecting symptoms that might suggest alternative diagnoses. Patients with appendicitis may not have any abdominal discomfort early in the disease process. Furthermore, many patients may not present with the classically described history or physical findings.

What is the classic history? Nonspecific complaints occur first. Patients may notice changes in bowel habits or malaise and vague, perhaps intermittent, crampy, abdominal pain in the epigastric or periumbilical region. The pain subsequently migrates to the right lower quadrant over 12–24 h, where it is sharper and can be definitively localized as transmural inflammation when the appendix irritates the parietal peritoneum. Parietal peritoneal irritation may be associated with local muscle rigidity and stiffness. Patients with appendicitis

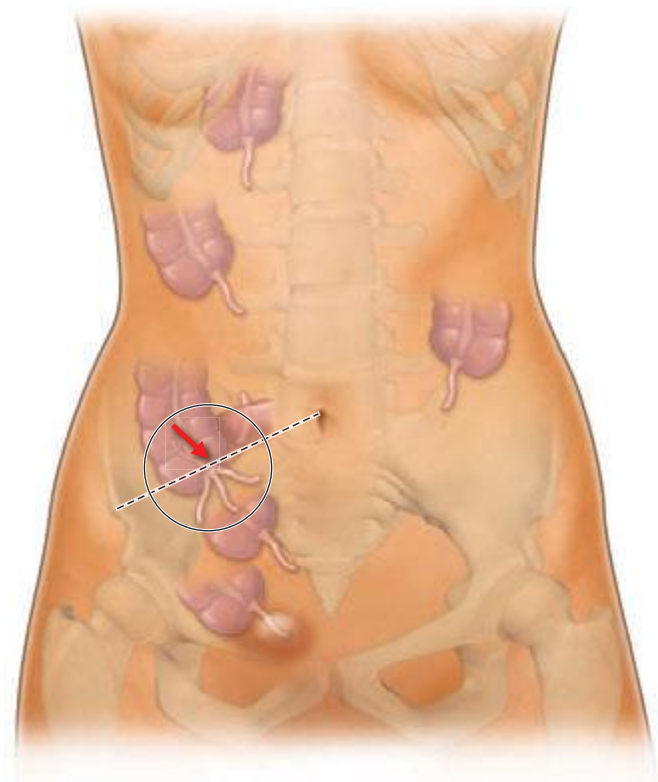


FIGURE 356-2 Locations of the appendix and cecum.

TABLE 356-1 SOME CONDITIONS THAT MIMIC APPENDICITIS

Crohn's disease	Meckel's diverticulitis
Cholecystitis or other gallbladder disease	Mittelschmerz
Diverticulitis	Mesenteric adenitis
Ectopic pregnancy	Omental torsion
Endometriosis	Pancreatitis
Gastroenteritis or colitis	Lower lobe pneumonia
Gastric or duodenal ulceration	Pelvic inflammatory disease
Hepatitis	Ruptured ovarian cyst or other cystic disease of the ovaries
Kidney disease, including nephrolithiasis	Small-bowel obstruction
Liver abscess	Urinary tract infection

TABLE 356-2 RELATIVE FREQUENCY OF COMMON PRESENTING SYMPTOMS

Symptoms	Frequency
Abdominal pain	>95%
Anorexia	>70%
Constipation	4–16%
Diarrhea	4–16%
Fever	10–20%
Migration of pain to right lower quadrant	50–60%
Nausea	>65%
Vomiting	50–75%

will most often observe that their nausea, if present, followed the development of abdominal pain, which can help distinguish them from patients with gastroenteritis, for example, where nausea occurs first. Emesis, if present, also occurs after the onset of pain and is typically mild and scant. Thus, timing of the onset of symptoms and the characteristics of the patient's pain and any associated findings must be rigorously assessed. Anorexia is so common that the diagnosis of appendicitis should be questioned in its absence.

Arriving at the correct diagnosis is even more challenging when the appendix is not located in the right lower quadrant, in women of childbearing age, and in the very young or elderly. Because the differential diagnosis of appendicitis is so broad, often the key question to answer expeditiously is whether the patient has appendicitis or some other condition that requires immediate operative intervention. A major concern is that the likelihood of a delay in diagnosis is greater if the appendix is unusually positioned. All patients should undergo a rectal examination. An inflamed appendix located behind the cecum or below the pelvic brim may prompt very little tenderness of the anterior abdominal wall.

Patients with pelvic appendicitis are more likely to present with dysuria, urinary frequency, diarrhea, or tenesmus. They may only experience pain in the suprapubic region on palpation or on rectal or pelvic examination. A pelvic examination in women is mandatory to rule out conditions affecting urogynecologic organs that can cause abdominal pain and mimic appendicitis such as pelvic inflammatory disease, ectopic pregnancy, and ovarian torsion. The relative frequencies of some presenting signs are displayed in [Table 356-3](#).

Patients with simple appendicitis normally only appear mildly ill with a pulse and temperature that are usually only slightly above normal. The provider should be concerned about other disease processes beside appendicitis or the presence of complications such as perforation, phlegmon, or abscess formation if the temperature is $>38.3^{\circ}\text{C}$ ($\sim 101^{\circ}\text{F}$) and if there are rigors.

Patients with appendicitis will be found to lie quite still to avoid peritoneal irritation caused by movement, and some will report discomfort caused by a bumpy car ride on the way to the hospital or clinic, coughing, sneezing, or other actions that replicate a Valsalva maneuver. The entire abdomen should be examined systematically starting in an area where the patient does not report discomfort if

TABLE 356-3 RELATIVE FREQUENCY OF SOME PRESENTING SIGNS

Signs	Frequency (%)
Abdominal tenderness	>95%
Right lower quadrant tenderness	>90%
Rebound tenderness	30–70%
Rectal tenderness	30–40%
Cervical motion tenderness	30%
Rigidity	$\sim 10\%$
Psoas sign	3–5%
Obturator sign	5–10%
Rovsing's sign	5%
Palpable mass	<5%

TABLE 356-4 CLASSIC SIGNS OF APPENDICITIS IN PATIENTS WITH ABDOMINAL PAIN

Maneuver	Findings
Rovsing's sign	Palpating in the left lower quadrant causes pain in the right lower quadrant
Obturator sign	Internal rotation of the hip causes pain, suggesting the possibility of an inflamed appendix located in the pelvis
Iliopsoas sign	Extending the right hip causes pain along posterolateral back and hip, suggesting retrocecal appendicitis

possible. Classically, maximal tenderness is identified in the right lower quadrant at or near McBurney's point, which is located approximately one-third of the way along a line originating at the anterior iliac spine and running to the umbilicus. Gentle pressure in the left lower quadrant may elicit pain in the right lower quadrant if the appendix is located there. This is Rovsing's sign ([Table 356-4](#)). Evidence of parietal peritoneal irritation is often best elicited by gentle abdominal percussion, jiggling the patient's gurney or bed, or mildly bumping the feet.

Atypical presentation and pain patterns are common, especially in the very old or the very young. Diagnosing appendicitis in children can be especially challenging because they tend to respond so dramatically to stimulation and obtaining an accurate history may be difficult. In addition, it is important to remember that the smaller omentum found in children may be less likely to wall off an appendiceal perforation. Observing the child in a quiet surrounding may be helpful.

Signs and symptoms of appendicitis can be subtle in the elderly who may not react as vigorously to appendicitis as younger people. Pain, if noticed, may be minimal and have originated in the right lower quadrant or, otherwise, where the appendix is located. It may never have been noticed to be intermittent, or there may only be significant discomfort with deep palpation. Nausea, anorexia, and emesis may be the predominant complaints. The rare patient may even present with signs and symptoms of distal bowel obstruction secondary to appendiceal inflammation and phlegmon or abscess formation.

LABORATORY TESTING

Laboratory testing does not identify patients with appendicitis but can help the clinician work through the differential diagnosis. The white blood cell count is only mildly to moderately elevated in approximately 70% of patients with simple appendicitis (with a leukocytosis of 10,000–18,000 cells/ μL). A "left shift" toward immature polymorphonuclear leukocytes is present in $>95\%$ of cases. A sickle cell preparation may be prudent to obtain in those of African, Spanish, Mediterranean, or Indian ancestry. Serum amylase and lipase levels should be measured.

Urinalysis is indicated to help exclude genitourinary conditions that may mimic acute appendicitis, but a few red or white blood cells may be present as a nonspecific finding. However, an inflamed appendix that abuts the ureter or bladder may cause sterile pyuria or hematuria. Every woman of childbearing age should have a pregnancy test. Cervical cultures are indicated if pelvic inflammatory disease is suspected. Anemia and guaiac-positive stools should raise concern about the presence of other diseases or complications such as cancer.

IMAGING

Plain films of the abdomen are rarely helpful and so are not routinely obtained unless the clinician is worried about other conditions such as intestinal obstruction, perforated viscus, or ureterolithiasis. Less than 5% of patients will present with an opaque fecalith in the right lower quadrant. The presence of a fecalith is not diagnostic of appendicitis, although its presence in an appropriate location where the patient complains of pain is suggestive.

The effectiveness of ultrasonography as a tool to diagnosis appendicitis is highly operator dependent. Even in very skilled hands, the appendix may not be visualized. Its overall sensitivity is 0.86, with a

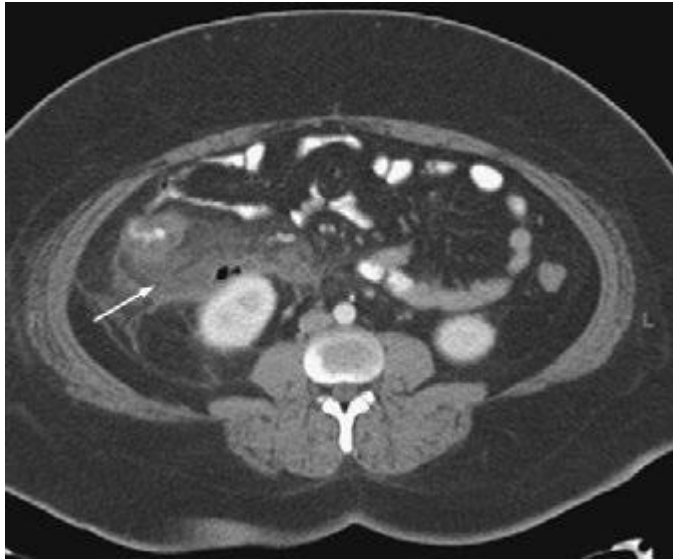


FIGURE 356-3 Computed tomography with oral and intravenous contrast of acute appendicitis. There is thickening of the wall of the appendix and periappendiceal stranding (*arrow*).

specificity of 0.81. Ultrasonography, especially intravaginal techniques, appears to be most useful for identifying pelvic pathology in women. Ultrasonographic findings suggesting the presence of appendicitis include wall thickening, an increased appendiceal diameter, and the presence of free fluid.

The sensitivity and specificity of computed tomography (CT) are 0.94 and 0.95, respectively. Thus, CT imaging, given its high negative predictive value, may be helpful if the diagnosis is in doubt, although studies performed early in the course of disease may not have any typical radiographic findings. Suggestive findings on CT examination include dilatation >6 mm with wall thickening, a lumen that does not fill with enteric contrast, and fatty tissue stranding or air surrounding the appendix, which suggests inflammation (**Figs. 356-3 and 356-4**). The presence of luminal air or contrast is not consistent with a diagnosis of appendicitis. Furthermore, nonvisualization of the appendix is a nonspecific finding that should not be used to rule out the presence of appendiceal or periappendiceal inflammation.

SPECIAL PATIENT POPULATIONS

Appendicitis is the most common extrauterine general surgical emergency observed during pregnancy. Early symptoms of appendicitis



FIGURE 356-4 Appendiceal fecalith (*arrow*).

such as nausea and anorexia may be overlooked. Diagnosing appendicitis in pregnant patients may be especially difficult because as the uterus enlarges the appendix may be pushed higher along the right flank even to the right upper quadrant or because the gravid uterus may obscure typical physical findings. Ultrasonography may facilitate early diagnosis. A high index of suspicion is required because of the effects of unrecognized and untreated appendicitis on the fetus. For example, the fetal mortality rate is four times greater (from 5 to 20%) in patients with perforation.

Immunocompromised patients may present with only mild tenderness and may have many other disease processes in their differential diagnosis, including atypical infections from mycobacteria, *Cytomegalovirus*, or other fungi. Enterocolitis is a concern and may be present in patients who present with abdominal pain, fever, and neutropenia due to chemotherapy. CT imaging may be very helpful, although it is important not to be overly cautious and delay operative intervention for those patients who are believed to have appendicitis.

TREATMENT ACUTE APPENDICITIS

In the absence of contraindications, a patient who has a strongly suggestive medical history and physical examination with supportive laboratory findings should undergo appendectomy urgently. In this instance, imaging studies are not required. In patients in whom the evaluation is suggestive but not convincing, imaging and further study are appropriate. Pelvic ultrasonography is indicated in women of childbearing age. Thereafter, CT may accurately indicate the presence of appendicitis or other intraabdominal processes that warrant intervention. Whenever the diagnosis is uncertain, it is prudent to observe the patient and repeat the abdominal examination over 6–8 h. Any evidence of progression is an indication for operation. Narcotics can be given to patients with severe discomfort, especially if the first abdominal examination is completed before drugs are administered.

All patients should be fully prepared for surgery and have any fluid and electrolyte abnormalities corrected. Either laparoscopic or open appendectomy is a satisfactory choice for patients with uncomplicated appendicitis. Management of those who present with a mass representing a phlegmon or abscess can be more difficult. Such patients are best served by treatment with broad-spectrum antibiotics, drainage if there is an abscess >3 cm in diameter, and parenteral fluids and bowel rest if they appear to respond to conservative management. The appendix can then be more safely removed 6–12 weeks later when inflammation has diminished.

Laparoscopic appendectomy now accounts for approximately 60% of all appendectomies. Laparoscopic appendectomy is associated with less postoperative pain and, possibly, a shorter length of stay and faster return to normal activity. Patients who undergo laparoscopic appendectomy also appear to have fewer wound infections, although the risk of intraabdominal abscess formation may be higher. A laparoscopic approach may also be useful when the exact diagnosis is uncertain, yet direct visualization and exploration of the abdomen are needed. A laparoscopic approach may also facilitate exposure in those who are very obese. A thorough examination of the abdomen is indicated if the appendix appears normal at operation, which can be expected to occur in up to 15–20% of cases.

Absent complications, most patients can be discharged within 24–40 h of operation. The most common postoperative complications are fever and leukocytosis. Continuation of these findings beyond 5 days should raise concern for the presence of an intraabdominal abscess. The mortality rate for uncomplicated, nonperforated appendicitis is 0.1–0.5%, which approximates the overall risk of general anesthesia. The mortality rate for perforated appendicitis or other complicated disease is much higher, ranging from 3% overall to a high as 15% in the elderly.

ACUTE PERITONITIS

Acute peritonitis, or inflammation of the visceral and parietal peritoneum, is most often but not always infectious in origin, resulting from perforation of a hollow viscus. This is called *secondary peritonitis*, as opposed to *primary* or *spontaneous peritonitis*, when a specific intraabdominal source cannot be identified. In either instance, the inflammation can be localized or diffuse.

ETIOLOGY

Infective organisms may contaminate the peritoneal cavity after spillage from a hollow viscus, because of a penetrating wound of the abdominal wall, or because of the introduction of a foreign object like a peritoneal dialysis catheter or port that becomes infected. Secondary peritonitis most commonly results from perforation of the appendix, colonic diverticuli, or the stomach and duodenum. It may also occur as a complication of bowel infarction or incarceration, cancer, inflammatory bowel disease, and intestinal obstruction or volvulus. Conditions that may cause secondary bacterial peritonitis and their mechanisms are listed in **Table 356-5**. Over 90% of the cases of primary or spontaneous bacterial peritonitis occur in patients with ascites or hyponatremia (<1 g/L).

TABLE 356-5 CONDITIONS LEADING TO SECONDARY BACTERIAL PERITONITIS

Bowel Perforation	Perforation or Leakage of Other Organs
Appendicitis trauma (blunt or penetrating)	Biliary leakage (e.g., after liver biopsy)
Anastomotic leakage	Cholecystitis
Adhesion	Intraperitoneal bleeding
Diverticulitis	Pancreatitis
Iatrogenic (including endoscopic perforation)	Salpingitis
Ingested foreign body	Traumatic or other rupture of urinary bladder
Inflammation	
Intussusception	Loss of peritoneal integrity
Neoplasms	Intraperitoneal chemotherapy
Obstruction	Iatrogenic (e.g., postoperative foreign body)
Peptic ulcer disease	Perinephric abscess
Strangulated hernia	Peritoneal dialysis or other indwelling devices
Vascular (including ischemia or embolus)	Trauma

Aseptic peritonitis is most commonly caused by the abnormal presence of physiologic fluids like gastric juice, bile, pancreatic enzymes, blood, or urine. It can also be caused by the effects of normally sterile foreign bodies like surgical sponges or instruments. More rarely, it occurs as a complication of systemic diseases like lupus erythematosus, porphyria, and familial Mediterranean fever. The chemical irritation caused by stomach acid and activated pancreatic enzymes is extreme and secondary bacterial infection may occur.

CLINICAL FEATURES

The cardinal signs and symptoms of peritonitis are acute, typically severe, abdominal pain with tenderness and fever. How the patient's complaints of pain are manifested depends on their overall physical health and whether the inflammation is diffuse or localized. Elderly and immunosuppressed patients may not respond as aggressively to the irritation. Diffuse, generalized peritonitis is most often recognized as diffuse abdominal tenderness with local guarding, rigidity, and other evidence of parietal peritoneal irritation. Physical findings may only be identified in a specific region of the abdomen if the intraperitoneal inflammatory process is limited or otherwise contained as may occur in patients with uncomplicated appendicitis or diverticulitis. Bowel sounds are usually absent to hypoactive.

Most patients present with tachycardia and signs of volume depletion with hypotension. Laboratory testing typically reveals a significant leukocytosis, and patients may be severely acidotic. Radiographic studies may show dilatation of the bowel and associated bowel wall edema. Free air, or other evidence of leakage, requires attention and could represent a surgical emergency. In stable patients in whom ascites is present, diagnostic paracentesis is indicated, where the fluid is tested for protein and lactate dehydrogenase and the cell count is measured.

THERAPY AND PROGNOSIS

Whereas mortality rates can be less than 10% for reasonably healthy patients with relatively uncomplicated, localized peritonitis, mortality rates >40% have been reported for the elderly or immunocompromised. Successful treatment depends on correcting any electrolyte abnormalities, restoration of fluid volume and stabilization of the cardiovascular system, appropriate antibiotic therapy, and surgical correction of any underlying abnormalities.

ACKNOWLEDGMENT

The wisdom and expertise of Dr. William Silen is gratefully acknowledged in this updated chapter on acute appendicitis and peritonitis.

SECTION 2 LIVER AND BILIARY TRACT DISEASE

357 Approach to the Patient with Liver Disease

Marc G. Ghany, Jay H. Hoofnagle

A diagnosis of liver disease usually can be made accurately by careful elicitation of the patient's history, physical examination, and application of a few laboratory tests. In some circumstances, radiologic examinations are helpful or, indeed, diagnostic. Liver biopsy is considered the criterion standard in evaluation of liver disease but is now needed less for diagnosis than for grading and staging of disease. This chapter provides an introduction to diagnosis and management of liver disease, briefly reviewing the structure and function of the liver; the major clinical manifestations of liver disease; and the use of clinical

history, physical examination, laboratory tests, imaging studies, and liver biopsy.

LIVER STRUCTURE AND FUNCTION

The liver is the largest organ of the body, weighing 1–1.5 kg and representing 1.5–2.5% of the lean body mass. The size and shape of the liver vary and generally match the general body shape—long and lean or squat and square. This organ is located in the right upper quadrant of the abdomen under the right lower rib cage against the diaphragm and projects for a variable extent into the left upper quadrant. It is held in place by ligamentous attachments to the diaphragm, peritoneum, great vessels, and upper gastrointestinal organs. The liver receives a dual blood supply; ~20% of the blood flow is oxygen-rich blood from the hepatic artery, and 80% is nutrient-rich blood from the portal vein arising from the stomach, intestines, pancreas, and spleen.

The majority of cells in the liver are hepatocytes, which constitute two-thirds of the organ's mass. The remaining cell types are Kupffer cells (members of the reticuloendothelial system), stellate (Ito or fat-storing) cells, endothelial and blood vessel cells, bile ductular cells, and cells of supporting structures. Viewed by light microscopy, the liver appears to be organized in lobules, with portal areas at the periphery and central veins in the center of each lobule. However, from a functional point of view, the liver is organized into acini, with both hepatic arterial and portal venous blood entering the acinus from the portal areas (zone 1) and then flowing through the sinusoids to the terminal hepatic veins (zone 3); the intervening hepatocytes constitute zone 2. The advantage of viewing the acinus as the physiologic unit of the liver is that this perspective helps to explain the morphologic patterns and zonality of many vascular and biliary diseases not explained by the lobular arrangement.

Portal areas of the liver consist of small veins, arteries, bile ducts, and lymphatics organized in a loose stroma of supporting matrix and small amounts of collagen. Blood flowing into the portal areas is distributed through the sinusoids, passing from zone 1 to zone 3 of the acinus and draining into the terminal hepatic veins ("central veins"). Secreted bile flows in the opposite direction—i.e., in a counter-current pattern from zone 3 to zone 1. The sinusoids are lined by unique endothelial cells that have prominent fenestrae of variable sizes, allowing the free flow of plasma but not of cellular elements. The plasma is thus in direct contact with hepatocytes in the subendothelial space of Disse.

Hepatocytes have distinct polarity. The basolateral side of the hepatocyte lines the space of Disse and is richly lined with microvilli; it exhibits endocytotic and pinocytotic activity, with passive and active uptake of nutrients, proteins, and other molecules. The apical pole of the hepatocyte forms the canalicular membranes through which bile components are secreted. The canaliculi of hepatocytes form a fine network, which fuses into the bile ductular elements near the portal areas. Kupffer cells usually lie within the sinusoidal vascular space and represent the largest group of fixed macrophages in the body. The stellate cells are located in the space of Disse but are not usually prominent unless activated, when they produce collagen and matrix. Red blood cells stay in the sinusoidal space as blood flows through the lobules, but white blood cells can migrate through or around endothelial cells into the space of Disse and from there to portal areas, where they can return to the circulation through lymphatics.

Hepatocytes perform numerous and vital roles in maintaining homeostasis and health. These functions include the synthesis of most essential serum proteins (albumin, carrier proteins, coagulation factors, many hormonal and growth factors), the production of bile and its carriers (bile acids, cholesterol, lecithin, phospholipids), the regulation of nutrients (glucose, glycogen, lipids, cholesterol, amino acids), and the metabolism and conjugation of lipophilic compounds (bilirubin, anions, cations, drugs) for excretion in the bile or urine. Measurement of these activities to assess liver function is complicated by the multiplicity and variability of these functions. The most commonly used liver "function" tests are measurements of serum bilirubin, serum albumin, and prothrombin time. The serum bilirubin level is a measure of hepatic conjugation and excretion; the serum albumin level and prothrombin time are measures of protein synthesis. Abnormalities of bilirubin, albumin, and prothrombin time are typical of hepatic dysfunction. Frank liver failure is incompatible with life, and the functions of the liver are too complex and diverse to be subserved by a mechanical pump; a dialysis membrane; or a concoction of infused hormones, proteins, and growth factors.

LIVER DISEASES

While there are many causes of liver disease (Table 357-1), these disorders generally present clinically in a few distinct patterns and are usually classified as hepatocellular, cholestatic (obstructive), or mixed. In *hepatocellular diseases* (such as viral hepatitis and alcoholic liver disease), features of liver injury, inflammation, and necrosis predominate. In *cholestatic diseases* (such as gallstone or malignant obstruction, primary biliary cirrhosis, and some drug-induced liver

TABLE 357-1 LIVER DISEASES

Inherited hyperbilirubinemia	Liver involvement in systemic diseases
Gilbert's syndrome	Sarcoidosis
Crigler-Najjar syndrome, types I and II	Amyloidosis
Dubin-Johnson syndrome	Glycogen storage diseases
Rotor syndrome	Celiac disease
Viral hepatitis	Tuberculosis
Hepatitis A	<i>Mycobacterium avium-intracellulare</i> infection
Hepatitis B	Cholestatic syndromes
Hepatitis C	Benign postoperative cholestasis
Hepatitis D	Jaundice of sepsis
Hepatitis E	Total parenteral nutrition-induced jaundice
Others (Epstein-Barr virus [mononucleosis] herpesvirus, adenovirus hepatitis)	Cholestasis of pregnancy
Cryptogenic hepatitis	Cholangitis and cholecystitis
Immune and autoimmune liver diseases	Extrahepatic biliary obstruction (stone, stricture, cancer)
Primary biliary cirrhosis	Biliary atresia
Autoimmune hepatitis	Caroli's disease
Sclerosing cholangitis	Cryptosporidiosis
Overlap syndromes	Drug-induced liver disease
Graft-versus-host disease	Hepatocellular patterns (isoniazid, acetaminophen)
Allograft rejection	Cholestatic patterns (methyltestosterone)
Genetic liver diseases	Mixed patterns (sulfonamides, phenytoin)
α_1 Antitrypsin deficiency	Micro- and macrovesicular steatosis (methotrexate, fialuridine)
Hemochromatosis	Vascular injury
Wilson's disease	Veno-occlusive disease
Benign recurrent intrahepatic cholestasis	Budd-Chiari syndrome
Progressive familial intrahepatic cholestasis, types I–III	Ischemic hepatitis
Others (galactosemia, tyrosinemia, cystic fibrosis, Newman-Pick disease, Gaucher's disease)	Passive congestion
Alcoholic liver disease	Portal vein thrombosis
Acute fatty liver	Nodular regenerative hyperplasia
Acute alcoholic hepatitis	Mass lesions
Laënnec's cirrhosis	Hepatocellular carcinoma
Nonalcoholic fatty liver	Cholangiocarcinoma
Steatosis	Adenoma
Steatohepatitis	Focal nodular hyperplasia
Acute fatty liver of pregnancy	Metastatic tumors
	Abscess
	Cysts
	Hemangioma

diseases), features of inhibition of bile flow predominate. In a mixed pattern, features of both hepatocellular and cholestatic injury are present (such as in cholestatic forms of viral hepatitis and many drug-induced liver diseases). The pattern of onset and prominence of symptoms can rapidly suggest a diagnosis, particularly if major risk factors are considered, such as the age and sex of the patient and a history of exposure or risk behaviors.

Typical presenting symptoms of liver disease include jaundice, fatigue, itching, right-upper-quadrant pain, nausea, poor appetite, abdominal distention, and intestinal bleeding. At present, however, many patients are diagnosed with liver disease who have no symptoms and who have been found to have abnormalities in biochemical liver tests as a part of a routine physical examination or screening for blood

donation or for insurance or employment. The wide availability of batteries of liver tests makes it relatively simple to demonstrate the presence of liver injury as well as to rule it out in someone in whom liver disease is suspected.

Evaluation of patients with liver disease should be directed at (1) establishing the etiologic diagnosis, (2) estimating disease severity (*grading*), and (3) establishing the disease stage (*staging*). *Diagnosis* should focus on the category of disease (hepatocellular, cholestatic, or mixed injury) as well as on the specific etiologic diagnosis. *Grading* refers to assessment of the severity or activity of disease—active or inactive as well as mild, moderate, or severe. *Staging* refers to estimation of the point in the course of the natural history of the disease, whether early or late; or precirrhotic, cirrhotic, or end-stage. This chapter introduces general, salient concepts in the evaluation of patients with liver disease that help lead to the diagnoses discussed in subsequent chapters.

CLINICAL HISTORY

The clinical history should focus on the symptoms of liver disease—their nature, patterns of onset, and progression—and on potential risk factors for liver disease. The manifestations of liver disease include constitutional symptoms such as fatigue, weakness, listlessness, malaise, increased need for sleep, lack of stamina, and poor energy. Symptoms can also suggest the presence of cirrhosis, end-stage liver disease, or complications of cirrhosis such as portal hypertension. Generally, the constellation of symptoms and their patterns of onset rather than a specific symptom points to an etiology.

Fatigue is the most common and most characteristic symptom of liver disease. It is variously described as lethargy, weakness, listlessness, malaise, increased need for sleep, lack of stamina, and poor energy. The fatigue of liver disease typically arises after activity or exercise and is rarely present or severe after adequate rest; i.e., it is “afternoon” rather than “morning” fatigue. Fatigue in liver disease is often intermittent and variable in severity from hour to hour and day to day. In some patients, it may not be clear whether fatigue is due to the liver disease or to other problems such as stress, anxiety, sleep disturbance, or a concurrent illness.

Nausea occurs with more severe liver disease and may accompany fatigue or be provoked by smelling food odors or eating fatty foods. Vomiting can occur but is rarely persistent or prominent. Poor appetite with weight loss occurs frequently in acute liver disease but is rare in chronic disease except when cirrhosis is present and advanced. Diarrhea is uncommon in liver disease except with severe jaundice, in which a lack of bile acids reaching the intestine can lead to steatorrhea.

Right-upper-quadrant discomfort or ache (“liver pain”) occurs in many liver diseases and is usually marked by tenderness over the liver area. The pain arises from stretching or irritation of Glisson’s capsule, which surrounds the liver and is rich in nerve endings. Severe pain is most typical of gallbladder disease, liver abscess, and severe veno-occlusive disease but is also an occasional accompaniment of acute hepatitis.

Itching occurs with acute liver disease, appearing early in obstructive jaundice (from biliary obstruction or drug-induced cholestasis) and somewhat later in hepatocellular disease (acute hepatitis). Itching also occurs in chronic liver diseases—typically the cholestatic forms such as primary biliary cirrhosis and sclerosing cholangitis, in which it is often the presenting symptom, preceding the onset of jaundice. However, itching can occur in any liver disease, particularly once cirrhosis develops.

Jaundice is the hallmark symptom of liver disease and perhaps the most reliable marker of severity. Patients usually report darkening of the urine before they notice scleral icterus. Jaundice is rarely detectable with a bilirubin level $<43 \mu\text{mol/L}$ (2.5 mg/dL). With severe cholestasis, there will also be lightening of the color of the stools and steatorrhea. Jaundice without dark urine usually indicates indirect (unconjugated) hyperbilirubinemia and is typical of hemolytic anemia and the genetic disorders of bilirubin conjugation, the common and benign form being Gilbert’s syndrome and the rare and severe form being

Crigler-Najjar syndrome. Gilbert’s syndrome affects up to 5% of the general population; the jaundice in this condition is more noticeable after fasting and with stress.

Major risk factors for liver disease that should be sought in the clinical history include details of alcohol use, medication use (including herbal compounds, birth control pills, and over-the-counter medications), personal habits, sexual activity, travel, exposure to jaundiced or other high-risk persons, injection drug use, recent surgery, remote or recent transfusion of blood or blood products, occupation, accidental exposure to blood or needlestick, and familial history of liver disease.

For assessing the risk of viral hepatitis, a careful history of sexual activity is of particular importance and should include the number of lifetime sexual partners and, for men, a history of having sex with men. Sexual exposure is a common mode of spread of hepatitis B but is rare for hepatitis C. A family history of hepatitis, liver disease, and liver cancer is also important. Maternal-infant transmission occurs with both hepatitis B and C. Vertical spread of hepatitis B can now be prevented by passive and active immunization of the infant at birth. Vertical spread of hepatitis C is uncommon, but there are no reliable means of prevention. Transmission is more common among HIV-co-infected mothers and is also linked to prolonged and difficult labor and delivery, early rupture of membranes, and internal fetal monitoring. A history of injection drug use, even in the remote past, is of great importance in assessing the risk for hepatitis B and C. Injection drug use is now the single most common risk factor for hepatitis C. Transfusion with blood or blood products is no longer an important risk factor for acute viral hepatitis. However, blood transfusions received before the introduction of sensitive enzyme immunoassays for antibody to hepatitis C virus in 1992 is an important risk factor for chronic hepatitis C. Blood transfusion before 1986, when screening for antibody to hepatitis B core antigen was introduced, is also a risk factor for hepatitis B. Travel to a developing area of the world, exposure to persons with jaundice, and exposure to young children in day-care centers are risk factors for hepatitis A. Tattooing and body piercing (for hepatitis B and C) and eating shellfish (for hepatitis A) are frequently mentioned but are actually types of exposure that quite rarely lead to the acquisition of hepatitis.



Hepatitis E is one of the more common causes of jaundice in Asia and Africa but is uncommon in developed nations. Recently, non-travel-related (*autochthonous*) cases of hepatitis E have been described in developed countries, including the United States. These cases appear to be due to strains of hepatitis E virus that are endemic in swine and some wild animals (genotypes 3 and 4). While occasional cases are associated with eating raw or undercooked pork or game (deer and wild boars), most cases of hepatitis E occur without known exposure, predominantly in elderly man without typical risk factors for viral hepatitis. Hepatitis E infection can become chronic in immunosuppressed individuals (such as transplant recipients, patients receiving chemotherapy, or patients with HIV infection), in whom it presents with abnormal serum enzymes in the absence of markers of hepatitis B or C.

A history of alcohol intake is important in assessing the cause of liver disease and also in planning management and recommendations. In the United States, for example, at least 70% of adults drink alcohol to some degree, but significant alcohol intake is less common; in population-based surveys, only 5% of individuals have more than two drinks per day, the average drink representing 11–15 g of alcohol. Alcohol consumption associated with an increased rate of alcoholic liver disease is probably more than two drinks (22–30 g) per day in women and three drinks (33–45 g) in men. Most patients with alcoholic cirrhosis have a much higher daily intake and have drunk excessively for ≥ 10 years before onset of liver disease. In assessing alcohol intake, the history should also focus on whether alcohol abuse or dependence is present. Alcoholism is usually defined by the behavioral patterns and consequences of alcohol intake, not by the amount. *Abuse* is defined by a repetitive pattern of drinking alcohol that has adverse effects on social, family, occupational, or health status. *Dependence* is defined by alcohol-seeking behavior, despite its adverse effects. Many

TABLE 357-2 CAGE QUESTIONS^a

Acronym	Question
C	Have you ever felt you ought to cut down on your drinking?
A	Have people annoyed you by criticizing your drinking?
G	Have you ever felt guilty or bad about your drinking?
E	Have you ever had a drink first thing in the morning to steady your nerves or get rid of a hangover (eye-opener)?

^aOne “yes” response should raise suspicion of an alcohol use problem, and more than one is a strong indication of abuse or dependence.

alcoholics demonstrate both dependence and abuse, and dependence is considered the more serious and advanced form of alcoholism. A clinically helpful approach to diagnosis of alcohol dependence and abuse is the use of the CAGE questionnaire (Table 357-2), which is recommended for all medical history-taking.

Family history can be helpful in assessing liver disease. Familial causes of liver disease include Wilson’s disease; hemochromatosis and α_1 antitrypsin deficiency; and the more uncommon inherited pediatric liver diseases—i.e., familial intrahepatic cholestasis, benign recurrent intrahepatic cholestasis, and Alagille syndrome. Onset of severe liver disease in childhood or adolescence in conjunction with a family history of liver disease or neuropsychiatric disturbance should lead to investigation for Wilson’s disease. A family history of cirrhosis, diabetes, or endocrine failure and the appearance of liver disease in adulthood suggests hemochromatosis and should prompt investigation of iron status. Abnormal iron studies in adult patients warrant genotyping of the *HFE* gene for the C282Y and H63D mutations typical of genetic hemochromatosis. In children and adolescents with iron overload, other non-*HFE* causes of hemochromatosis should be sought. A family history of emphysema should provoke investigation of α_1 antitrypsin levels and, if levels are low, for protease inhibitor (Pi) genotype.

PHYSICAL EXAMINATION

The physical examination rarely uncovers evidence of liver dysfunction in a patient without symptoms or laboratory findings, nor are most signs of liver disease specific to one diagnosis. Thus, the physical examination complements rather than replaces the need for other diagnostic approaches. In many patients, the physical examination is normal unless the disease is acute or severe and advanced. Nevertheless, the physical examination is important in that it can yield the first evidence of hepatic failure, portal hypertension, and liver decompensation. In addition, the physical examination can reveal signs—related either to risk factors or to associated diseases or findings—that point to a specific diagnosis.

Typical physical findings in liver disease are icterus, hepatomegaly, hepatic tenderness, splenomegaly, spider angiomas, palmar erythema, and excoriations. Signs of advanced disease include muscle wasting, ascites, edema, dilated abdominal veins, hepatic fetor, asterixis, mental confusion, stupor, and coma. In male patients with cirrhosis, particularly that related to alcohol use, signs of hyperestrogenemia such as gynecomastia, testicular atrophy, and loss of male-pattern hair distribution may be found.

Icterus is best appreciated when the sclera is inspected under natural light. In fair-skinned individuals, a yellow tinge to the skin may be obvious. In dark-skinned individuals, examination of the mucous membranes below the tongue can demonstrate jaundice. Jaundice is rarely detectable if the serum bilirubin level is $<43 \mu\text{mol/L}$ (2.5 mg/dL) but may remain detectable below this level during recovery from jaundice (because of protein and tissue binding of conjugated bilirubin).

Spider angiomas and palmar erythema occur in both acute and chronic liver disease; these manifestations may be especially prominent in persons with cirrhosis but can develop in normal individuals and are frequently found during pregnancy. Spider angiomas are superficial, tortuous arterioles and—unlike simple telangiectases—typically fill from the center outward. Spider angiomas occur only on the arms, face, and upper torso; they can be pulsatile and may be difficult to detect in dark-skinned individuals.

Hepatomegaly is not a highly reliable sign of liver disease because of variability in the liver’s size and shape and the physical impediments to assessment of liver size by percussion and palpation. Marked hepatomegaly is typical of cirrhosis, veno-occlusive disease, infiltrative disorders such as amyloidosis, metastatic or primary cancers of the liver, and alcoholic hepatitis. Careful assessment of the liver edge may also reveal unusual firmness, irregularity of the surface, or frank nodules. Perhaps the most reliable physical finding in the liver examination is hepatic tenderness. Discomfort when the liver is touched or pressed upon should be carefully sought with percussive comparison of the right and left upper quadrants.

Splenomegaly, which occurs in many medical conditions, can be a subtle but significant physical finding in liver disease. The availability of ultrasound methods for assessment of the spleen allows confirmation of the physical finding.

Signs of advanced liver disease include muscle wasting and weight loss as well as hepatomegaly, bruising, ascites, and edema. Ascites is best appreciated by attempts to detect shifting dullness by careful percussion. Ultrasound examination will confirm the finding of ascites in equivocal cases. Peripheral edema can occur with or without ascites. In patients with advanced liver disease, other factors frequently contribute to edema formation, including hypoalbuminemia, venous insufficiency, heart failure, and medications.

Hepatic failure is defined as the occurrence of signs or symptoms of hepatic encephalopathy in a person with severe acute or chronic liver disease. The first signs of hepatic encephalopathy can be subtle and nonspecific—change in sleep patterns, change in personality, irritability, and mental dullness. Thereafter, confusion, disorientation, stupor, and eventually coma supervene. In acute liver failure, excitability and mania may be present. Physical findings include asterixis and flapping tremors of the body and tongue. *Fetor hepaticus* refers to the slightly sweet, ammoniacal odor that can develop in patients with liver failure, particularly if there is portal-venous shunting of blood around the liver. Other causes of coma and disorientation should be excluded, mainly electrolyte imbalances, sedative use, and renal or respiratory failure. The appearance of hepatic encephalopathy during acute hepatitis is the major criterion for diagnosis of fulminant hepatitis and indicates a poor prognosis. In chronic liver disease, encephalopathy is usually triggered by a medical complication such as gastrointestinal bleeding, over-diuresis, uremia, dehydration, electrolyte imbalance, infection, constipation, or use of narcotic analgesics.

A helpful measure of hepatic encephalopathy is a careful mental-status examination and use of the trail-making test, which consists of a series of 25 numbered circles that the patient is asked to connect as rapidly as possible using a pencil. The normal range for the connect-the-dot test is 15–30 sec; it is considerably longer in patients with early hepatic encephalopathy. Other tests include drawing of abstract objects or comparison of a signature to previous examples. More sophisticated testing—e.g., with electroencephalography and visual evoked potentials—can detect mild forms of encephalopathy but are rarely clinically useful.

Other signs of advanced liver disease include umbilical hernia from ascites, hydrothorax, prominent veins over the abdomen, and *caput medusa*, a condition that consists of collateral veins radiating from the umbilicus and results from recanalization of the umbilical vein. Widened pulse pressure and signs of a hyperdynamic circulation can occur in patients with cirrhosis as a result of fluid and sodium retention, increased cardiac output, and reduced peripheral resistance. Patients with long-standing cirrhosis and portal hypertension are prone to develop the hepatopulmonary syndrome, which is defined by the triad of liver disease, hypoxemia, and pulmonary arteriovenous shunting. The hepatopulmonary syndrome is characterized by platypnea and orthodeoxia: shortness of breath and oxygen desaturation that occur paradoxically upon the assumption of an upright position. Measurement of oxygen saturation by pulse oximetry is a reliable screening test for hepatopulmonary syndrome.

Several skin disorders and changes are common in liver disease. Hyperpigmentation is typical of advanced chronic cholestatic diseases such as primary biliary cirrhosis and sclerosing cholangitis. In these

same conditions, xanthelasma and tendon xanthomata occur as a result of retention and high serum levels of lipids and cholesterol. Slate-gray pigmentation of the skin is also seen with hemochromatosis if iron levels are high for a prolonged period. Mucocutaneous vasculitis with palpable purpura, especially on the lower extremities, is typical of cryoglobulinemia of chronic hepatitis C but can also occur in chronic hepatitis B.

Some physical signs point to specific liver diseases. Kayser-Fleischer rings occur in Wilson's disease and consist of a golden-brown copper pigment deposited in Descemet's membrane at the periphery of the cornea; they are best seen by slit-lamp examination. Dupuytren contracture and parotid enlargement are suggestive of chronic alcoholism and alcoholic liver disease. In metastatic liver disease or primary hepatocellular carcinoma, signs of cachexia and wasting as well as firm hepatomegaly and a hepatic bruit may be prominent.

DIAGNOSIS OF LIVER DISEASE

The major causes of liver disease and key diagnostic features are outlined in [Table 357-3](#), and an algorithm for evaluation of the patient with suspected liver disease is shown in [Fig. 357-1](#). Specifics of diagnosis are discussed in later chapters. The most common causes of acute liver disease are viral hepatitis (particularly hepatitis A, B, and C), drug-induced liver injury, cholangitis, and alcoholic liver disease. Liver biopsy usually is not needed in the diagnosis and management of acute liver disease, exceptions being situations where the diagnosis remains unclear despite thorough clinical and laboratory investigation. Liver biopsy can be helpful in diagnosing drug-induced liver disease and acute alcoholic hepatitis.

The most common causes of chronic liver disease, in general order of frequency, are chronic hepatitis C, alcoholic liver disease, nonalcoholic steatohepatitis, chronic hepatitis B, autoimmune hepatitis, sclerosing cholangitis, primary biliary cirrhosis, hemochromatosis, and Wilson's disease. Hepatitis E virus is a rare cause of chronic hepatitis, with cases occurring mostly in persons who are immunosuppressed or immunodeficient. Strict diagnostic criteria have not been developed for most

liver diseases, but liver biopsy plays an important role in the diagnosis of autoimmune hepatitis, primary biliary cirrhosis, nonalcoholic and alcoholic steatohepatitis, and Wilson's disease (with a quantitative hepatic copper level in the last instance).

Laboratory Testing Diagnosis of liver disease is greatly aided by the availability of reliable and sensitive tests of liver injury and function. A typical battery of blood tests used for initial assessment of liver disease includes measurement of levels of serum alanine and aspartate aminotransferases, alkaline phosphatase, direct and total serum bilirubin and albumin, and prothrombin time. The pattern of abnormalities generally points to hepatocellular versus cholestatic liver disease and helps determine whether the disease is acute or chronic and whether cirrhosis and hepatic failure are present. On the basis of these results, further testing over time may be necessary. Other laboratory tests may be helpful, such as γ -glutamyl transpeptidase to define whether alkaline phosphatase elevations are due to liver disease; hepatitis serology to define the type of viral hepatitis; and autoimmune markers to diagnose primary biliary cirrhosis (antimitochondrial antibody), sclerosing cholangitis (peripheral antineutrophil cytoplasmic antibody), and autoimmune hepatitis (antinuclear, smooth-muscle, and liver-kidney microsomal antibody). A simple delineation of laboratory abnormalities and common liver diseases is given in [Table 357-3](#).

The use and interpretation of liver function tests are summarized in [Chap. 358](#).

Diagnostic Imaging Great advances have been made in hepatobiliary imaging, although no method is adequately accurate in demonstrating underlying cirrhosis. Of the many modalities available for imaging the liver, ultrasound, CT, and MRI are the most commonly employed and are complementary to one another. In general, ultrasound and CT are highly sensitive for detecting biliary duct dilation and are the first-line options for investigating cases of suspected obstructive jaundice. All three modalities can detect a fatty liver, which appears bright on imaging studies. Modifications of CT and MRI can be used to quantify liver fat, and this information may ultimately be valuable in monitoring therapy in patients with fatty liver disease. Magnetic resonance cholangiopancreatography (MRCP) and endoscopic retrograde cholangiopancreatography (ERCP) are the procedures of choice for visualization of the biliary tree. MRCP offers several advantages over ERCP: there is no need for contrast media or ionizing radiation, images can be acquired faster, the procedure is less operator dependent, and it carries no risk of pancreatitis. MRCP is superior to ultrasound and CT for detecting choledocholithiasis but is less specific. MRCP is useful in the diagnosis of bile duct obstruction and congenital biliary abnormalities, but ERCP is more valuable in evaluating ampullary lesions and primary sclerosing cholangitis. ERCP permits biopsy, direct visualization of the ampulla and common bile duct, and intraductal ultrasonography. It also provides several therapeutic options in patients with obstructive jaundice, such as sphincterotomy, stone extraction, and placement of nasobiliary catheters and biliary stents. Doppler ultrasound and MRI are used to assess hepatic vasculature and hemodynamics and to monitor surgically or radiologically placed vascular shunts, including transjugular intrahepatic portosystemic shunts. Multidetector or spiral CT and MRI with contrast-enhancement are the procedures of choice for the identification and evaluation of hepatic masses, the staging of liver tumors, and preoperative assessment. With regard to mass lesions, the sensitivity of hepatic imaging continues to increase; unfortunately, specificity remains a problem, and often two and sometimes three studies are needed before a diagnosis can be reached. Recently, ultrasound transient elastography has been approved for the measurement of hepatic stiffness—providing an indirect assessment of cirrhosis; this technique can eliminate the need for liver biopsy if the only indication is the assessment of disease stage. Magnetic resonance elastography is now undergoing evaluation for its ability to detect different degrees of hepatic fibrosis. Studies are ongoing to determine whether hepatic elastography is an appropriate means of monitoring fibrosis and disease progression. Finally, interventional radiologic techniques allow the biopsy of solitary lesions, the radiofrequency

TABLE 357-3 IMPORTANT DIAGNOSTIC TESTS IN COMMON LIVER DISEASES

Disease	Diagnostic Test
Hepatitis A	Anti-HAV IgM
Hepatitis B	
Acute	HBsAg and anti-HBc IgM
Chronic	HBsAg and HBeAg and/or HBV DNA
Hepatitis C	Anti-HCV and HCV RNA
Hepatitis D (delta)	HBsAg and anti-HDV
Hepatitis E	Anti-HEV IgM and HEV RNA
Autoimmune hepatitis	ANA or SMA, elevated IgG levels, and compatible histology
Primary biliary cirrhosis	Mitochondrial antibody, elevated IgM levels, and compatible histology
Primary sclerosing cholangitis	P-ANCA, cholangiography
Drug-induced liver disease	History of drug ingestion
Alcoholic liver disease	History of excessive alcohol intake and compatible histology
Nonalcoholic steatohepatitis	Ultrasound or CT evidence of fatty liver and compatible histology
α_1 Antitrypsin disease	Reduced α_1 antitrypsin levels, phenotype PiZZ or PiSZ
Wilson's disease	Decreased serum ceruloplasmin and increased urinary copper; increased hepatic copper level
Hemochromatosis	Elevated iron saturation and serum ferritin; genetic testing for <i>HFE</i> gene mutations
Hepatocellular cancer	Elevated α -fetoprotein level (to >500 ng/mL); ultrasound or CT image of mass

Abbreviations: HAV, HBV, HCV, HDV, HEV: hepatitis A, B, C, D, E virus; HBsAg, hepatitis B surface antigen; anti-HBc, antibody to hepatitis B core (antigen); HBeAg, hepatitis B e antigen; ANA, antinuclear antibody; SMA, smooth-muscle antibody; P-ANCA, peripheral antineutrophil cytoplasmic antibody.

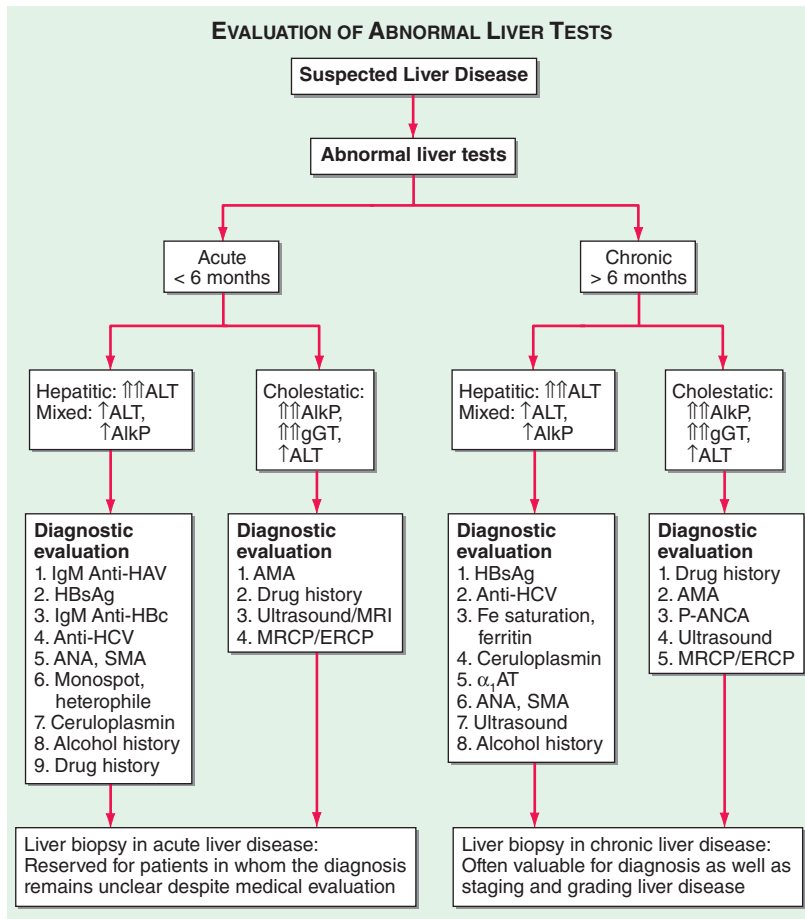


FIGURE 357-1 Algorithm for evaluation of abnormal liver tests. For patients with suspected liver disease, an appropriate approach to evaluation is initial routine liver testing—e.g., measurement of serum bilirubin, albumin, alanine aminotransferase (ALT), aspartate aminotransferase (AST), and alkaline phosphatase (AlkP). These results (sometimes complemented by testing of γ -glutamyl transpeptidase; gGT) will establish whether the pattern of abnormalities is hepatic, cholestatic, or mixed. In addition, the duration of symptoms or abnormalities will indicate whether the disease is acute or chronic. If the disease is acute and if history, laboratory tests, and imaging studies do not reveal a diagnosis, liver biopsy is appropriate to help establish the diagnosis. If the disease is chronic, liver biopsy can be helpful not only for diagnosis but also for grading of the activity and staging the progression of disease. This approach is generally applicable to patients without immune deficiency. In patients with HIV infection or recipients of bone marrow or solid organ transplants, the diagnostic evaluation should also include evaluation for opportunistic infections (e.g., with adenovirus, cytomegalovirus, *Coccidioides*, hepatitis E virus) as well as for vascular and immunologic conditions (veno-occlusive disease, graft-versus-host disease). HAV, hepatitis A virus; HCV, hepatitis C virus; HBsAg, hepatitis B surface antigen; anti-HBc, antibody to hepatitis B core (antigen); ANA, antinuclear antibody; SMA, smooth-muscle antibody; MRCP, magnetic resonance cholangiopancreatography; ERCP, endoscopic retrograde cholangiopancreatography; α_1 AT, α_1 antitrypsin; AMA, antimitochondrial antibody; P-ANCA, peripheral antineutrophil cytoplasmic antibody.

ablation and chemoembolization of cancerous lesions, the insertion of drains into hepatic abscesses, the measurement of portal pressure, and the creation of vascular shunts in patients with portal hypertension. Which modality to use depends on factors such as availability, cost, and experience of the radiologist with each technique.

Liver Biopsy Liver biopsy remains the criterion standard in the evaluation of patients with liver disease, particularly chronic liver disease. Liver biopsy is necessary for diagnosis in selected instances but is more often useful for assessment of the severity (grade) and stage of liver damage, prediction of prognosis, and monitoring of the response to treatment. The size of the liver biopsy sample is an important determinant of reliability; a length of 1.5–2 cm is necessary for accurate

assessment of fibrosis. In the future, noninvasive means of assessing disease activity (batteries of blood tests) and fibrosis (elastography and fibrosis markers) may replace liver biopsy for the staging and grading of disease.

GRADING AND STAGING OF LIVER DISEASE

Grading refers to an assessment of the severity or activity of liver disease, whether acute or chronic; active or inactive; and mild, moderate, or severe. Liver biopsy is the most accurate means of assessing severity, particularly in chronic liver disease. Serum aminotransferase levels serve as convenient and noninvasive markers for disease activity but do not always reliably reflect disease severity. Thus, normal serum aminotransferase levels in patients with hepatitis B surface antigen in serum may indicate the inactive carrier state or may reflect mild chronic hepatitis B or hepatitis B with fluctuating disease activity. Serum testing for hepatitis B e antigen and hepatitis B virus DNA can help sort out these different patterns, but these markers can also fluctuate and change over time. Similarly, in chronic hepatitis C, serum aminotransferase levels can be normal despite moderate disease activity. Finally, in both alcoholic and nonalcoholic steatohepatitis, aminotransferase levels are quite unreliable in reflecting severity. In these conditions, liver biopsy is helpful in guiding management and identifying appropriate therapy, particularly if treatment is difficult, prolonged, and expensive, as is often the case in chronic viral hepatitis. Of the several well-verified numerical scales for grading activity in chronic liver disease, the most commonly used are the histology activity index and the Ishak histology scale.

Liver biopsy is also the most accurate means of assessing stage of disease as early or advanced, precirrhotic, and cirrhotic. Staging of disease pertains largely to chronic liver diseases in which progression to cirrhosis and end-stage disease can occur but may require years or decades. Clinical features, biochemical tests, and hepatic imaging studies are helpful in assessing stage but generally become abnormal only in the middle to late stages of cirrhosis. Noninvasive tests that suggest advanced fibrosis include mild elevations of bilirubin, prolongation of prothrombin time, slight decreases in serum albumin, and mild thrombocytopenia (which is often the first indication of worsening fibrosis). Combinations of blood test results have been used to create models for predicting advanced liver disease, but these models are not reliable enough to use on a regular basis and only separate advanced from early disease. Recently, elastography and noninvasive breath tests using ^{13}C -labeled compounds have been proposed as a means of detecting early stages of fibrosis and liver dysfunction, but their reliability and reproducibility remain to be proven. Thus, at present, mild to moderate stages of hepatic fibrosis are detectable only by liver biopsy.

In the assessment of stage, the degree of fibrosis is usually used as the quantitative measure. The amount of fibrosis is generally staged on a scale of 0 to 4+ (Metavir scale) or 0 to 6+ (Ishak scale). The importance of staging relates primarily to prognosis and to optimal management of complications. Patients with cirrhosis are candidates for screening and surveillance for esophageal varices and hepatocellular carcinoma. Patients without advanced fibrosis need not undergo screening.

Cirrhosis can also be staged clinically. A reliable staging system is the modified Child-Pugh classification, with a scoring system of 5–15: scores of 5 and 6 represent Child-Pugh class A (consistent with “compensated cirrhosis”), scores of 7–9 represent class B, and scores of 10–15 represent class C (Table 357-4). This scoring system was initially devised to stratify patients into risk groups before portal

TABLE 357-4 CHILD-PUGH CLASSIFICATION OF CIRRHOSIS

Factor	Units	Points Toward Total Score		
		1	2	3
Serum bilirubin	μmol/L	<34	34–51	>51
	mg/dL	<2.0	2.0–3.0	>3.0
Serum albumin	g/L	>35	30–35	<30
	g/dL	>3.5	3.0–3.5	<3.0
Prothrombin time	seconds prolonged	<4	4–6	>6
	INR ^a	<1.7	1.7–2.3	>2.3
Ascites		None	Easily controlled	Poorly controlled
Hepatic encephalopathy		None	Minimal	Advanced

^aInternational normalized ratio.

Note: The Child-Pugh score is calculated by adding the scores for the five factors and can range from 5 to 15. The resulting Child-Pugh class can be A (a score of 5–6), B (7–9), or C (≥10). Decompensation indicates cirrhosis, with a Child-Pugh score of ≥7 (class B). This level has been the accepted criterion for listing a patient for liver transplantation.

decompressive surgery. The Child-Pugh score is a reasonably reliable predictor of survival in many liver diseases and predicts the likelihood of major complications of cirrhosis, such as bleeding from varices and spontaneous bacterial peritonitis. This classification scheme was used to assess prognosis in cirrhosis and to provide standard criteria for listing a patient as a candidate for liver transplantation (Child-Pugh class B). Recently, the Child-Pugh system has been replaced by the Model for End-Stage Liver Disease (MELD) system for the latter purpose. The MELD score is a prospectively derived system designed to predict the prognosis of patients with liver disease and portal hypertension. This score is calculated from three noninvasive variables: the prothrombin time expressed as the international normalized ratio (INR), the serum bilirubin level, and the serum creatinine concentration. (<http://optn.transplant.hrsa.gov/resources/MeldPeldCalculator.asp?index=98>).

The MELD system provides a more objective means of assessing disease severity and has less center-to-center variation than the Child-Pugh score as well as a wider range of values. MELD is currently used to establish priority listing for liver transplantation in the United States. A similar system, PELD (pediatric end-stage liver disease), is based on bilirubin, INR, serum albumin, age, and nutritional status and is used for children <12 years of age.

Thus, liver biopsy is helpful not only in diagnosis but also in management of chronic liver disease and assessment of prognosis. Because liver biopsy is an invasive procedure and not without complications, it should be used only when it will contribute materially to decisions about management and therapy.

NONSPECIFIC ISSUES IN THE MANAGEMENT OF PATIENTS WITH LIVER DISEASE

Specifics on the management of different forms of acute or chronic liver disease are supplied in subsequent chapters, but certain issues are applicable to any patient with liver disease. These issues include advice regarding alcohol use, medication use, vaccination, and surveillance for complications of liver disease. Alcohol should be used sparingly, if at all, by patients with liver disease. Abstinence from alcohol should be encouraged for all patients with alcohol-related liver disease, patients with cirrhosis, and patients receiving interferon-based therapy for hepatitis B or C. With regard to vaccinations, all patients with liver disease should receive hepatitis A vaccine, and those with risk factors should receive hepatitis B vaccine as well. Influenza and pneumococcal vaccination should also be encouraged, with adherence to the recommendations of the Centers for Disease Control and Prevention. Patients with liver disease should exercise caution in using any medications other than those that are most necessary. Drug-induced hepatotoxicity can mimic many forms of liver disease and can cause exacerbations of chronic hepatitis and cirrhosis; drugs should be suspected in any situation in which the cause of exacerbation is unknown. Finally,

consideration should be given to surveillance for complications of chronic liver disease such as variceal hemorrhage and hepatocellular carcinoma. Cirrhosis warrants upper endoscopy to assess the presence of varices, and the patient should receive chronic therapy with beta blockers or should be offered endoscopic obliteration if large varices are found. Moreover, cirrhosis warrants screening and long-term surveillance for development of hepatocellular carcinoma. While the optimal regimen for such surveillance has not been established, an appropriate approach is ultrasound of the liver at 6- to 12-month intervals.

358 Evaluation of Liver Function

Daniel S. Pratt

Several biochemical tests are useful in the evaluation and management of patients with hepatic dysfunction. These tests can be used to (1) detect the presence of liver disease, (2) distinguish among different types of liver disorders, (3) gauge the extent of known liver damage, and (4) follow the response to treatment.

Liver tests have shortcomings. They can be normal in patients with serious liver disease and abnormal in patients with diseases that do not affect the liver. Liver tests rarely suggest a specific diagnosis; rather, they suggest a general category of liver disease, such as hepatocellular or cholestatic, which then further directs the evaluation.

The liver carries out thousands of biochemical functions, most of which cannot be easily measured by blood tests. Laboratory tests measure only a limited number of these functions. In fact, many tests, such as the aminotransferases or alkaline phosphatase, do not measure liver function at all. Rather, they detect liver cell damage or interference with bile flow. Thus, no one test enables the clinician to accurately assess the liver's total functional capacity.

To increase both the sensitivity and the specificity of laboratory tests in the detection of liver disease, it is best to use them as a battery. Tests usually employed in clinical practice include the bilirubin, aminotransferases, alkaline phosphatase, albumin, and prothrombin time tests. When more than one of these tests provide abnormal findings or the findings are persistently abnormal on serial determinations, the probability of liver disease is high. When all test results are normal, the probability of missing occult liver disease is low.

When evaluating patients with liver disorders, it is helpful to group these tests into general categories as outlined below.

TESTS BASED ON DETOXIFICATION AND EXCRETORY FUNCTIONS

Serum Bilirubin (See also Chap. 58) Bilirubin, a breakdown product of the porphyrin ring of heme-containing proteins, is found in the blood in two fractions—conjugated and unconjugated. The unconjugated fraction, also termed the *indirect fraction*, is insoluble in water and is bound to albumin in the blood. The conjugated (direct) bilirubin fraction is water soluble and can therefore be excreted by the kidney. When measured by modifications of the original van den Bergh method, normal values of total serum bilirubin are reported between 1 and 1.5 mg/dL with 95% of a normal population falling between 0.2 and 0.9 mg/dL. If the direct-acting fraction is less than 15% of the total, the bilirubin can be considered to all be indirect. The most frequently reported upper limit of normal for conjugated bilirubin is 0.3 mg/dL.

Elevation of the unconjugated fraction of bilirubin is rarely due to liver disease. An isolated elevation of unconjugated bilirubin is seen primarily in hemolytic disorders and in a number of genetic conditions such as Crigler-Najjar and Gilbert's syndromes (Chap. 58). Isolated unconjugated hyperbilirubinemia (bilirubin elevated but <15% direct) should prompt a workup for hemolysis (Fig. 358-1). In the absence of hemolysis, an isolated, unconjugated hyperbilirubinemia in an otherwise healthy patient can be attributed to Gilbert's syndrome, and no further evaluation is required.

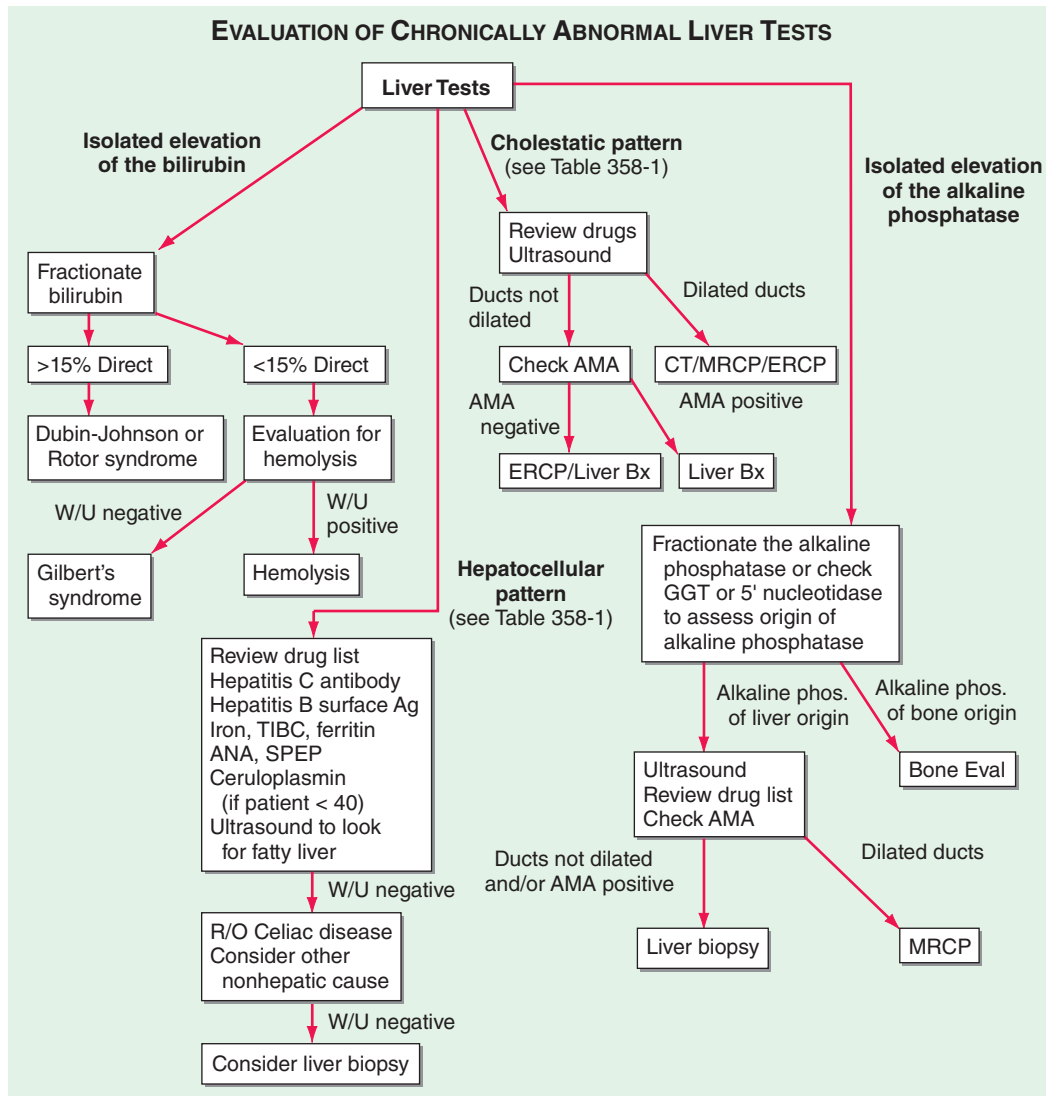


FIGURE 358-1 Algorithm for the evaluation of chronically abnormal liver tests. AMA, antimitochondrial antibody; ANA, antinuclear antibody; Bx, biopsy; CT, computed tomography; ERCP, endoscopic retrograde cholangiopancreatography; GGT, γ glutamyl transpeptidase; MRCP, magnetic resonance cholangiopancreatography; R/O, rule out; SPEP, serum protein electrophoresis; TIBC, total iron-binding capacity; W/U, workup.

In contrast, conjugated hyperbilirubinemia almost always implies liver or biliary tract disease. The rate-limiting step in bilirubin metabolism is not conjugation of bilirubin, but rather the transport of conjugated bilirubin into the bile canaliculi. Thus, elevation of the conjugated fraction may be seen in any type of liver disease. In most liver diseases, both conjugated and unconjugated fractions of the bilirubin tend to be elevated. Except in the presence of a purely unconjugated hyperbilirubinemia, fractionation of the bilirubin is rarely helpful in determining the cause of jaundice.

Although the degree of elevation of the serum bilirubin has not been critically assessed as a prognostic marker, it is important in a number of conditions. In viral hepatitis, the higher the serum bilirubin, the greater is the hepatocellular damage. Total serum bilirubin correlates with poor outcomes in alcoholic hepatitis. It is also a critical component of the Model for End-Stage Liver Disease (MELD) score, a tool used to estimate survival of patients with end-stage liver disease and assess operative risk of patients with cirrhosis. An elevated total serum bilirubin in patients with drug-induced liver disease indicates more severe injury.

Urine Bilirubin Unconjugated bilirubin always binds to albumin in the serum and is not filtered by the kidney. Therefore, any bilirubin found in the urine is conjugated bilirubin; the presence of bilirubinuria implies the presence of liver disease. A urine dipstick test can theoretically give the same information as fractionation of the

serum bilirubin. This test is almost 100% accurate. Phenothiazines may give a false-positive reading with the Ictotest tablet. In patients recovering from jaundice, the urine bilirubin clears prior to the serum bilirubin.

Blood Ammonia Ammonia is produced in the body during normal protein metabolism and by intestinal bacteria, primarily those in the colon. The liver plays a role in the detoxification of ammonia by converting it to urea, which is excreted by the kidneys. Striated muscle also plays a role in detoxification of ammonia, where it is combined with glutamic acid to form glutamine. Patients with advanced liver disease typically have significant muscle wasting, which likely contributes to hyperammonemia in these patients. Some physicians use the blood ammonia for detecting encephalopathy or for monitoring hepatic synthetic function, although its use for either of these indications has problems. There is very poor correlation between either the presence or the severity of acute encephalopathy and elevation of blood ammonia; it can be occasionally useful for identifying occult liver disease in patients with mental status changes. There is also a poor correlation of the blood serum ammonia and hepatic function. The ammonia can be elevated in patients with severe portal hypertension and portal blood shunting around the liver even in the presence of normal or near-normal hepatic function. Elevated arterial ammonia levels have been shown to correlate with outcome in fulminant hepatic failure.

Serum Enzymes The liver contains thousands of enzymes, some of which are also present in the serum in very low concentrations. These enzymes have no known function in the serum and behave like other serum proteins. They are distributed in the plasma and in interstitial fluid and have characteristic half-lives, which are usually measured in days. Very little is known about the catabolism of serum enzymes, although they are probably cleared by cells in the reticuloendothelial system. The elevation of a given enzyme activity in the serum is thought to primarily reflect its increased rate of entrance into serum from damaged liver cells.

Serum enzyme tests can be grouped into three categories: (1) enzymes whose elevation in serum reflects damage to hepatocytes, (2) enzymes whose elevation in serum reflects cholestasis, and (3) enzyme tests that do not fit precisely into either pattern.

ENZYMES THAT REFLECT DAMAGE TO HEPATOCYTES The aminotransferases (transaminases) are sensitive indicators of liver cell injury and are most helpful in recognizing acute hepatocellular diseases such as hepatitis. They include aspartate aminotransferase (AST) and alanine aminotransferase (ALT). AST is found in the liver, cardiac muscle, skeletal muscle, kidneys, brain, pancreas, lungs, leukocytes, and erythrocytes in decreasing order of concentration. ALT is found primarily in the liver and is therefore a more specific indicator of liver injury. The aminotransferases are normally present in the serum in low concentrations. These enzymes are released into the blood in greater amounts when there is damage to the liver cell membrane resulting in increased permeability. Liver cell necrosis is not required for the release of the aminotransferases, and there is a poor correlation between the degree of liver cell damage and the level of the aminotransferases. Thus, the absolute elevation of the aminotransferases is of no prognostic significance in acute hepatocellular disorders.

The normal range for aminotransferases varies widely among laboratories, but generally ranges from 10–40 IU/L. The interlaboratory variation in normal range is due to technical reasons; no reference standards exist to establish upper limits of normal for ALT and AST. Some have recommended revisions of normal limits of the aminotransferases to adjust for sex and body mass index, but others have noted the potential costs and unclear benefits of implementing this change.

Any type of liver cell injury can cause modest elevations in the serum aminotransferases. Levels of up to 300 IU/L are nonspecific and may be found in any type of liver disorder. Minimal ALT elevations in asymptomatic blood donors rarely indicate severe liver disease; studies have shown that fatty liver disease is the most likely explanation. Striking elevations—i.e., aminotransferases >1000 IU/L—occur almost exclusively in disorders associated with extensive hepatocellular injury such as (1) viral hepatitis, (2) ischemic liver injury (prolonged hypotension or acute heart failure), or (3) toxin- or drug-induced liver injury.

The pattern of the aminotransferase elevation can be helpful diagnostically. In most acute hepatocellular disorders, the ALT is higher than or equal to the AST. Whereas the AST:ALT ratio is typically <1 in patients with chronic viral hepatitis and nonalcoholic fatty liver disease, a number of groups have noted that as cirrhosis develops, this ratio rises to >1. An AST:ALT ratio >2:1 is suggestive, whereas a ratio >3:1 is highly suggestive, of alcoholic liver disease. The AST in alcoholic liver disease is rarely >300 IU/L, and the ALT is often normal. A low level of ALT in the serum is due to an alcohol-induced deficiency of pyridoxal phosphate.

The aminotransferases are usually not greatly elevated in obstructive jaundice. One notable exception occurs during the acute phase of biliary obstruction caused by the passage of a gallstone into the common bile duct. In this setting, the aminotransferases can briefly be in the 1000–2000 IU/L range. However, aminotransferase levels decrease quickly, and the liver function tests rapidly evolve into those typical of cholestasis.

ENZYMES THAT REFLECT CHOLESTASIS The activities of three enzymes—alkaline phosphatase, 5'-nucleotidase, and γ -glutamyl transpeptidase (GGT)—are usually elevated in cholestasis. Alkaline phosphatase and

5'-nucleotidase are found in or near the bile canalicular membrane of hepatocytes, whereas GGT is located in the endoplasmic reticulum and in bile duct epithelial cells. Reflecting its more diffuse localization in the liver, GGT elevation in serum is less specific for cholestasis than are elevations of alkaline phosphatase or 5'-nucleotidase. Some have advocated the use of GGT to identify patients with occult alcohol use. Its lack of specificity makes its use in this setting questionable.

The normal serum alkaline phosphatase consists of many distinct isoenzymes found in the liver; bone; placenta; and, less commonly, small intestine. Patients over age 60 can have a mildly elevated alkaline phosphatase (1–1.5 times normal), whereas individuals with blood types O and B can have an elevation of the serum alkaline phosphatase after eating a fatty meal due to the influx of intestinal alkaline phosphatase into the blood. It is also nonpathologically elevated in children and adolescents undergoing rapid bone growth because of bone alkaline phosphatase, and late in normal pregnancies due to the influx of placental alkaline phosphatase.

Elevation of liver-derived alkaline phosphatase is not totally specific for cholestasis, and a less than threefold elevation can be seen in almost any type of liver disease. Alkaline phosphatase elevations greater than four times normal occur primarily in patients with cholestatic liver disorders, infiltrative liver diseases such as cancer and amyloidosis, and bone conditions characterized by rapid bone turnover (e.g., Paget's disease). In bone diseases, the elevation is due to increased amounts of the bone isoenzymes. In liver diseases, the elevation is almost always due to increased amounts of the liver isoenzyme.

If an elevated serum alkaline phosphatase is the only abnormal finding in an apparently healthy person, or if the degree of elevation is higher than expected in the clinical setting, identification of the source of elevated isoenzymes is helpful (Fig. 358-1). This problem can be approached in two ways. First, and most precise, is the fractionation of the alkaline phosphatase by electrophoresis. The second, best substantiated, and most available approach involves the measurement of serum 5'-nucleotidase or GGT. These enzymes are rarely elevated in conditions other than liver disease.

In the absence of jaundice or elevated aminotransferases, an elevated alkaline phosphatase of liver origin often, but not always, suggests early cholestasis and, less often, hepatic infiltration by tumor or granulomata. Other conditions that cause isolated elevations of the alkaline phosphatase include Hodgkin's disease, diabetes, hyperthyroidism, congestive heart failure, amyloidosis, and inflammatory bowel disease.

The level of serum alkaline phosphatase elevation is not helpful in distinguishing between intrahepatic and extrahepatic cholestasis. There is essentially no difference among the values found in obstructive jaundice due to cancer, common duct stone, sclerosing cholangitis, or bile duct stricture. Values are similarly increased in patients with intrahepatic cholestasis due to drug-induced hepatitis; primary biliary cirrhosis; rejection of transplanted livers; and, rarely, alcohol-induced steatohepatitis. Values are also greatly elevated in hepatobiliary disorders seen in patients with AIDS (e.g., AIDS cholangiopathy due to cytomegalovirus or cryptosporidial infection and tuberculosis with hepatic involvement).

TESTS THAT MEASURE BIOSYNTHETIC FUNCTION OF THE LIVER

Serum Albumin Serum albumin is synthesized exclusively by hepatocytes. Serum albumin has a long half-life: 18–20 days, with ~4% degraded per day. Because of this slow turnover, the serum albumin is not a good indicator of acute or mild hepatic dysfunction; only minimal changes in the serum albumin are seen in acute liver conditions such as viral hepatitis, drug-related hepatotoxicity, and obstructive jaundice. In hepatitis, albumin levels <3 g/dL should raise the possibility of chronic liver disease. Hypoalbuminemia is more common in chronic liver disorders such as cirrhosis and usually reflects severe liver damage and decreased albumin synthesis. One exception is the patient with ascites in whom synthesis may be normal or even increased, but levels are low because of the increased volume of distribution. However, hypoalbuminemia is not specific for liver disease and may occur in protein malnutrition of any cause, as well as protein-losing enteropathies, nephrotic syndrome, and chronic infections that are

1998 associated with prolonged increases in levels of serum interleukin 1 and/or tumor necrosis factor, cytokines that inhibit albumin synthesis. Serum albumin should not be measured for screening in patients in whom there is no suspicion of liver disease. A general medical clinic study of consecutive patients in whom no indications were present for albumin measurement showed that although 12% of patients had abnormal test results, the finding was of clinical importance in only 0.4%.

Serum Globulins Serum globulins are a group of proteins made up of γ globulins (immunoglobulins) produced by B lymphocytes and α and β globulins produced primarily in hepatocytes. γ globulins are increased in chronic liver disease, such as chronic hepatitis and cirrhosis. In cirrhosis, the increased serum γ globulin concentration is due to the increased synthesis of antibodies, some of which are directed against intestinal bacteria. This occurs because the cirrhotic liver fails to clear bacterial antigens that normally reach the liver through the hepatic circulation.

Increases in the concentration of specific isotypes of γ globulins are often helpful in the recognition of certain chronic liver diseases. Diffuse polyclonal increases in IgG levels are common in autoimmune hepatitis; increases >100% should alert the clinician to this possibility. Increases in the IgM levels are common in primary biliary cirrhosis, whereas increases in the IgA levels occur in alcoholic liver disease.

COAGULATION FACTORS

With the exception of factor VIII, which is produced by vascular endothelial cells, the blood clotting factors are made exclusively in hepatocytes. Their serum half-lives are much shorter than albumin, ranging from 6 h for factor VII to 5 days for fibrinogen. Because of their rapid turnover, measurement of the clotting factors is the single best acute measure of hepatic synthetic function and helpful in both diagnosis and assessing the prognosis of acute parenchymal liver disease. Useful for this purpose is the *serum prothrombin time*, which collectively measures factors II, V, VII, and X. Biosynthesis of factors II, VII, IX, and X depends on vitamin K. The international normalized ratio (INR) is used to express the degree of anticoagulation on warfarin therapy. The INR standardizes prothrombin time measurement according to the

characteristics of the thromboplastin reagent used in a particular lab, which is expressed as an International Sensitivity Index (ISI); the ISI is then used in calculating the INR.

The prothrombin time may be elevated in hepatitis and cirrhosis as well as in disorders that lead to vitamin K deficiency such as obstructive jaundice or fat malabsorption of any kind. Marked prolongation of the prothrombin time, >5 s above control and not corrected by parenteral vitamin K administration, is a poor prognostic sign in acute viral hepatitis and other acute and chronic liver diseases. The INR, along with the total serum bilirubin and creatinine, are components of the MELD score, which is used as a measure of hepatic decompensation and to allocate organs for liver transplantation.

OTHER DIAGNOSTIC TESTS

Although tests may direct the physician to a category of liver disease, additional radiologic testing and procedures are often necessary to make the proper diagnosis, as shown in Fig. 358-1. The most commonly used ancillary tests are reviewed here, as are the noninvasive tests available for assessing hepatic fibrosis.

Percutaneous Liver Biopsy Percutaneous biopsy of the liver is a safe procedure that can be easily performed at the bedside with local anesthesia and ultrasound guidance. Liver biopsy is of proven value in the following situations: (1) hepatocellular disease of uncertain cause, (2) prolonged hepatitis with the possibility of autoimmune hepatitis, (3) unexplained hepatomegaly, (4) unexplained splenomegaly, (5) hepatic filling defects by radiologic imaging, (6) fever of unknown origin, (7) and staging of malignant lymphoma. Liver biopsy is most accurate in disorders causing diffuse changes throughout the liver and is subject to sampling error in focal infiltrative disorders such as hepatic metastases. Liver biopsy should not be the initial procedure in the diagnosis of cholestasis. The biliary tree should first be assessed for signs of obstruction. Contraindications to performing a percutaneous liver biopsy include significant ascites and prolonged INR. Under these circumstances, the biopsy can be performed via the transjugular approach.

TABLE 358-1 LIVER TEST PATTERNS IN HEPATOBILIARY DISORDERS

Type of Disorder	Bilirubin	Aminotransferases	Alkaline Phosphatase	Albumin	Prothrombin Time
Hemolysis/Gilbert's syndrome	Normal to 86 μ mol/L (5 mg/dL) 85% due to indirect fractions No bilirubinuria	Normal	Normal	Normal	Normal
Acute hepatocellular necrosis (viral and drug hepatitis, hepatotoxins, acute heart failure)	Both fractions may be elevated Peak usually follows aminotransferases Bilirubinuria	Elevated, often >500 IU, ALT > AST	Normal to <3 \times normal elevation	Normal	Usually normal. If >5 \times above control and not corrected by parenteral vitamin K, suggests poor prognosis
Chronic hepatocellular disorders	Both fractions may be elevated Bilirubinuria	Elevated, but usually <300 IU	Normal to <3 \times normal elevation	Often decreased	Often prolonged Fails to correct with parenteral vitamin K
Alcoholic hepatitis, cirrhosis	Both fractions may be elevated Bilirubinuria	AST:ALT >2 suggests alcoholic hepatitis or cirrhosis	Normal to <3 \times normal elevation	Often decreased	Often prolonged Fails to correct with parenteral vitamin K
Intra- and extrahepatic cholestasis	Both fractions may be elevated	Normal to moderate elevation	Elevated, often >4 \times normal elevation	Normal, unless chronic	Normal If prolonged, will correct with parenteral vitamin K
(Obstructive jaundice)	Bilirubinuria	Rarely >500 IU		Normal	Normal
Infiltrative diseases (tumor, granulomata); partial bile duct obstruction	Usually normal	Normal to slight elevation	Elevated, often >4 \times normal elevation Fractionate, or confirm liver origin with 5'-nucleotidase or γ glutamyl transpeptidase		

Noninvasive Tests to Detect Hepatic Fibrosis Although liver biopsy is the standard for the assessment of hepatic fibrosis, noninvasive measures of hepatic fibrosis have been developed and show promise. These measures include multiparameter tests aimed at detecting and staging the degree of hepatic fibrosis and imaging techniques. FibroTest (marketed as FibroSure in the United States) is the best evaluated of the multiparameter blood tests. The test incorporates haptoglobin, bilirubin, GGT, apolipoprotein A-I, and α 2-macroglobulin and has been found to have high positive and negative predictive values for diagnosing advanced fibrosis in patients with chronic hepatitis C, chronic hepatitis B, and alcoholic liver disease and patients taking methotrexate for psoriasis. Transient elastography (TE), marketed as FibroScan, and magnetic resonance elastography (MRE) both have gained U.S. Food and Drug Administration approval for use in the management of patients with liver disease. TE uses ultrasound waves to measure hepatic stiffness noninvasively. TE has been shown to be accurate for identifying advanced fibrosis in patients with chronic hepatitis C, primary biliary cirrhosis, hemochromatosis, nonalcoholic fatty liver disease, and recurrent chronic hepatitis after liver transplantation. MRE has been found to be superior to TE for staging liver fibrosis in patients with a variety of chronic liver diseases, but requires access to a magnetic resonance imaging scanner.

Ultrasonography Ultrasonography is the first diagnostic test to use in patients whose liver tests suggest cholestasis, to look for the presence of a dilated intrahepatic or extrahepatic biliary tree or to identify gallstones. In addition, it shows space-occupying lesions within the liver, enables the clinician to distinguish between cystic and solid masses, and helps direct percutaneous biopsies. Ultrasonography with Doppler imaging can detect the patency of the portal vein, hepatic artery, and hepatic veins and determine the direction of blood flow. This is the first test ordered in patients suspected of having Budd-Chiari syndrome.

USE OF LIVER TESTS

As previously noted, the best way to increase the sensitivity and specificity of laboratory tests in the detection of liver disease is to employ a battery of tests that includes the aminotransferases, alkaline phosphatase, bilirubin, albumin, and prothrombin time along with the judicious use of the other tests described in this chapter. **Table 358-1** shows how patterns of liver tests can lead the clinician to a category of disease that will direct further evaluation. However, it is important to remember that no single set of liver tests will necessarily provide a diagnosis. It is often necessary to repeat these tests on several occasions over days to weeks for a diagnostic pattern to emerge. Figure 358-1 is an algorithm for the evaluation of chronically abnormal liver tests.

GLOBAL CONSIDERATIONS

The tests and principles presented in this chapter are applicable worldwide. The causes of liver test abnormalities vary according to region. In developing nations, infectious diseases are more commonly the etiology of abnormal serum liver tests than in developed nations.

ACKNOWLEDGMENT

This chapter represents a revised version of a chapter in previous editions of *Harrison's* in which Marshall M. Kaplan was a co-author.

BILIRUBIN METABOLISM

The details of bilirubin metabolism are presented in **Chap. 58**. However, the hyperbilirubinemias are best understood in terms of perturbations of specific aspects of bilirubin metabolism and transport, and these will be briefly reviewed here as depicted in **Fig. 359-1**.

Bilirubin is the end product of heme degradation. Some 70–90% of bilirubin is derived from degradation of the hemoglobin of senescent red blood cells. Bilirubin produced in the periphery is transported to the liver within the plasma, where, due to its insolubility in aqueous solutions, it is tightly bound to albumin. Under normal circumstances, bilirubin is removed from the circulation rapidly and efficiently by hepatocytes. Transfer of bilirubin from blood to bile involves four distinct but interrelated steps (Fig. 359-1).

1. **Hepatocellular uptake:** Uptake of bilirubin by the hepatocyte has carrier-mediated kinetics. Although a number of candidate bilirubin transporters have been proposed, the actual transporter remains elusive.
2. **Intracellular binding:** Within the hepatocyte, bilirubin is kept in solution by binding as a nonsubstrate ligand to several of the glutathione-S-transferases, formerly called ligandins.
3. **Conjugation:** Bilirubin is conjugated with one or two glucuronic acid moieties by a specific UDP-glucuronosyltransferase to form bilirubin mono- and diglucuronide, respectively. Conjugation disrupts the internal hydrogen bonding that limits aqueous solubility of bilirubin, and the resulting glucuronide conjugates are highly soluble in water. Conjugation is obligatory for excretion of bilirubin across the bile canalicular membrane into bile. The UDP-glucuronosyltransferases have been classified into gene families based on the degree of homology among the mRNAs for the various isoforms. Those that conjugate bilirubin and certain

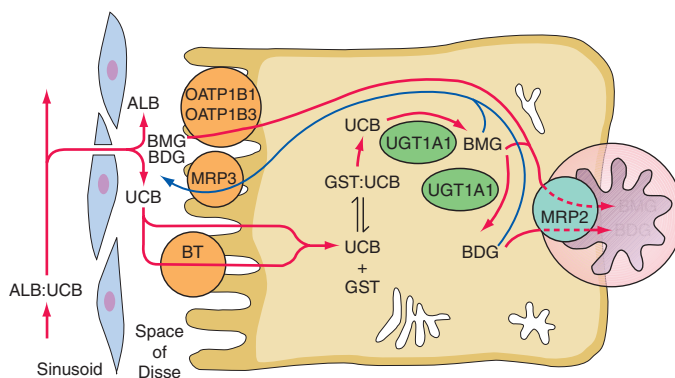


FIGURE 359-1 Hepatocellular bilirubin transport. Albumin-bound bilirubin in sinusoidal blood passes through endothelial cell fenestrae to reach the hepatocyte surface, entering the cell by both facilitated and simple diffusional processes. Within the cell, it is bound to glutathione-S-transferases and conjugated by bilirubin-UDP-glucuronosyltransferase (UGT1A1) to mono- and diglucuronides, which are actively transported across the canalicular membrane into the bile. In addition to this direct excretion of bilirubin glucuronides, a portion are transported into the portal circulation by MRP3 and subjected to reuptake into the hepatocyte by OATP1B1 and OATP1B3. ALB, albumin; BDG, bilirubin diglucuronide; BMG, bilirubin monoglucuronide; BT, proposed bilirubin transporter; GST, glutathione-S-transferase; MRP2 and MRP3, multidrug resistance-associated proteins 2 and 3; OATP1B1 and OATP1B3, organic anion transport proteins 1B1 and 1B3; UCB, unconjugated bilirubin; UGT1A1, bilirubin-UDP-glucuronosyltransferase.

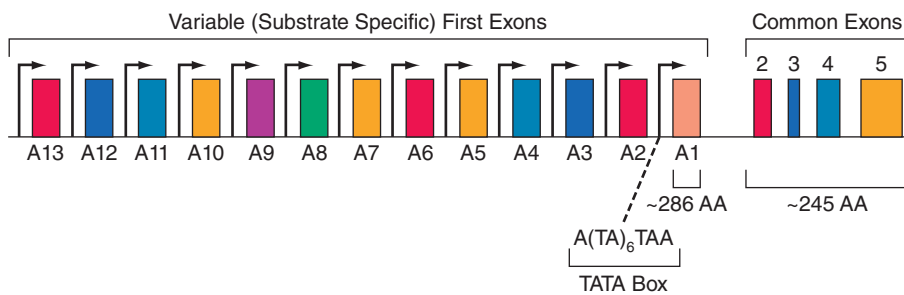


FIGURE 359-2 Structural organization of the human *UGT1* gene complex. This large complex on chromosome 2 contains at least 13 substrate-specific first exons (A1, A2, etc.). Since four of these are pseudogenes, nine *UGT1* isoforms with differing substrate specificities are expressed. Each exon 1 has its own promoter and encodes the amino-terminal substrate-specific ~286 amino acids of the various *UGT1*-encoded isoforms, and common exons 2–5 that encode the 245 carboxyl-terminal amino acids common to all of the isoforms. mRNAs for specific isoforms are assembled by splicing a particular first exon such as the bilirubin-specific exon A1 to exons 2 to 5. The resulting message encodes a complete enzyme, in this particular case bilirubin-UDP-glucuronosyltransferase (*UGT1A1*). Mutations in a first exon affect only a single isoform. Those in exons 2–5 affect all enzymes encoded by the *UGT1* complex.

other substrates have been designated the *UGT1* family. These are expressed from a single gene complex by alternative promoter usage. This gene complex contains multiple substrate-specific first exons, designated A1, A2, etc. (Fig. 359-2), each with its own promoter and each encoding the amino-terminal half of a specific isoform. In addition, there are four common exons (exons 2–5) that encode the shared carboxyl-terminal half of all of the *UGT1* isoforms. The various first exons encode the specific aglycone substrate binding sites for each isoform, while the shared exons encode the binding site for the sugar donor, UDP-glucuronic acid, and the transmembrane domain. Exon A1 and the four common exons, collectively designated the *UGT1A1* gene (Fig. 359-2), encode the physiologically critical enzyme bilirubin-UDP-glucuronosyltransferase (*UGT1A1*). A functional corollary of the organization of the *UGT1* gene is that a mutation in one of the first exons will affect only a single enzyme isoform. By contrast, a mutation in exons 2–5 will alter all isoforms encoded by the *UGT1* gene complex.

4. **Biliary excretion:** It has been thought until recently that bilirubin mono- and diglucuronides are excreted directly across the canalicular plasma membrane into the bile canaliculus by an ATP-dependent transport process mediated by a canalicular membrane protein called *multidrug resistance-associated protein 2* (*MRP2*). Mutations of *MRP2* result in the Dubin-Johnson syndrome (see below). However, studies in patients with Rotor syndrome (see below) indicate that after formation, a portion of the glucuronides are transported into the portal circulation by a sinusoidal membrane protein called *multidrug resistance-associated protein 3* (*MRP3*) and subjected to reuptake into the hepatocyte by the sinusoidal membrane uptake transporters *organic anion transport protein 1B1* (*OATP1B1*) and *OATP1B3*.

EXTRAHEPATIC ASPECTS OF BILIRUBIN DISPOSITION

Bilirubin in the Gut Following secretion into bile, conjugated bilirubin reaches the duodenum and passes down the gastrointestinal tract without reabsorption by the intestinal mucosa. An appreciable fraction is converted by bacterial metabolism in the gut to the water-soluble colorless compound urobilinogen. Urobilinogen undergoes enterohepatic cycling. Urobilinogen not taken up by the liver reaches the systemic circulation, from which some is cleared by the kidneys. Unconjugated bilirubin ordinarily does not reach the gut except in neonates or, by ill-defined alternative pathways, in the presence of severe unconjugated

hyperbilirubinemia (e.g., Crigler-Najjar syndrome, type I [CN-I]). Unconjugated bilirubin that reaches the gut is partly reabsorbed, amplifying any underlying hyperbilirubinemia. Recent reports suggest that oral administration of calcium phosphate with or without the lipase inhibitor orlistat may be an efficient means to interrupt bilirubin enterohepatic cycling to reduce serum bilirubin levels in this situation. Although orlistat administration for 4–6 weeks to 16 patients with Crigler-Najjar syndrome was associated with a 10–20% decrease in serum bilirubin in 7 patients, the cost and side effects (i.e., diarrhea) may obviate the small benefit achievable with this treatment.

Renal Excretion of Bilirubin Conjugates

Unconjugated bilirubin is not excreted in urine, as it is too tightly bound to albumin for effective glomerular filtration and there is no tubular mechanism for its renal secretion. In contrast, the bilirubin conjugates are readily filtered

at the glomerulus and can appear in urine in disorders characterized by increased bilirubin conjugates in the circulation.

DISORDERS OF BILIRUBIN METABOLISM LEADING TO UNCONJUGATED HYPERBILIRUBINEMIA

INCREASED BILIRUBIN PRODUCTION

Hemolysis Increased destruction of erythrocytes leads to increased bilirubin turnover and unconjugated hyperbilirubinemia; the hyperbilirubinemia is usually modest in the presence of normal liver function. In particular, the bone marrow is only capable of a sustained eightfold increase in erythrocyte production in response to a hemolytic stress. Therefore, hemolysis alone cannot result in a sustained hyperbilirubinemia of more than ~68 $\mu\text{mol/L}$ (4 mg/dL). Higher values imply concomitant hepatic dysfunction. When hemolysis is the only abnormality in an otherwise healthy individual, the result is a purely unconjugated hyperbilirubinemia, with the direct-reacting fraction as measured in a typical clinical laboratory being $\leq 15\%$ of the total serum bilirubin. In the presence of systemic disease, which may include a degree of hepatic dysfunction, hemolysis may produce a component of conjugated hyperbilirubinemia in addition to an elevated unconjugated bilirubin concentration. Prolonged hemolysis may lead to the precipitation of bilirubin salts within the gallbladder or biliary tree, resulting in the formation of gallstones in which bilirubin, rather than cholesterol, is the major component. Such pigment stones may lead to acute or chronic cholecystitis, biliary obstruction, or any other biliary tract consequence of calculous disease.

Ineffective Erythropoiesis During erythroid maturation, small amounts of hemoglobin may be lost at the time of nuclear extrusion, and a fraction of developing erythroid cells is destroyed within the marrow. These processes normally account for a small proportion of bilirubin that is produced. In various disorders, including thalassemia major, megaloblastic anemias due to folate or vitamin B₁₂ deficiency, congenital erythropoietic porphyria, lead poisoning, and various congenital and acquired dyserythropoietic anemias, the fraction of total bilirubin production derived from ineffective erythropoiesis is increased, reaching as much as 70% of the total. This may be sufficient to produce modest degrees of unconjugated hyperbilirubinemia.

Miscellaneous Degradation of the hemoglobin of extravascular collections of erythrocytes, such as those seen in massive tissue infarctions or large hematomas, may lead transiently to unconjugated hyperbilirubinemia.

DECREASED HEPATIC BILIRUBIN CLEARANCE

Decreased Hepatic Uptake Decreased hepatic bilirubin uptake is believed to contribute to the unconjugated hyperbilirubinemia of Gilbert syndrome (GS), although the molecular basis for this finding remains unclear (see below). Several drugs, including flavaspidic acid, novobiocin, and rifampin, as well as various cholecystographic contrast agents, have been reported to inhibit bilirubin uptake. The resulting unconjugated hyperbilirubinemia resolves with cessation of the medication.

Impaired Conjugation • PHYSIOLOGIC NEONATAL JAUNDICE Bilirubin produced by the fetus is cleared by the placenta and eliminated by the maternal liver. Immediately after birth, the neonatal liver must assume responsibility for bilirubin clearance and excretion. However, many hepatic physiologic processes are incompletely developed at birth. Levels of UGT1A1 are low, and alternative excretory pathways allow passage of unconjugated bilirubin into the gut. Since the intestinal flora that convert bilirubin to urobilinogen are also undeveloped, an enterohepatic circulation of unconjugated bilirubin ensues. As a consequence, most neonates develop mild unconjugated hyperbilirubinemia between days 2 and 5 after birth. Peak levels are typically <85–170 $\mu\text{mol/L}$ (5–10 mg/dL) and decline to normal adult concentrations within 2 weeks, as mechanisms required for bilirubin disposition mature. Prematurity, often associated with more profound immaturity of hepatic function and hemolysis, can result in higher levels of unconjugated hyperbilirubinemia. A rapidly rising unconjugated bilirubin concentration, or absolute levels >340 $\mu\text{mol/L}$ (20 mg/dL), puts the infant at risk for bilirubin encephalopathy, or kernicterus. Under these circumstances, bilirubin crosses an immature blood-brain barrier and precipitates in the basal ganglia and other areas of the brain. The consequences range from appreciable neurologic deficits to death. Treatment options include phototherapy, which converts bilirubin into water-soluble photoisomers that are excreted directly into bile, and exchange transfusion. The canalicular mechanisms responsible for bilirubin excretion are also immature at birth, and their maturation may lag behind that of UGT1A1; this can lead to transient conjugated neonatal hyperbilirubinemia, especially in infants with hemolysis.

ACQUIRED CONJUGATION DEFECTS A modest reduction in bilirubin conjugating capacity may be observed in advanced hepatitis or cirrhosis. However, in this setting, conjugation is better preserved than other aspects of bilirubin disposition, such as canalicular excretion. Various drugs, including pregnanediol, novobiocin, chloramphenicol, and gentamicin, may produce unconjugated hyperbilirubinemia by inhibiting UGT1A1 activity. Bilirubin conjugation may be inhibited by certain fatty acids that are present in breast milk but not serum of mothers whose infants have excessive neonatal hyperbilirubinemia

(*breast milk jaundice*). Alternatively, there may be increased enterohepatic circulation of bilirubin in these infants. A recent study has correlated epidermal growth factor (EGF) content of breast milk with elevated bilirubin levels in these infants; however, a cause-and-effect relationship remains to be established. The pathogenesis of breast milk jaundice appears to differ from that of transient familial neonatal hyperbilirubinemia (Lucey-Driscoll syndrome), in which there is a UGT1A1 inhibitor in maternal serum.

HEREDITARY DEFECTS IN BILIRUBIN CONJUGATION

Three familial disorders characterized by differing degrees of unconjugated hyperbilirubinemia have long been recognized. The defining clinical features of each are described below (Table 359-1). While these disorders have been recognized for decades to reflect differing degrees of deficiency in the ability to conjugate bilirubin, recent advances in the molecular biology of the *UGT1* gene complex have elucidated their interrelationships and clarified previously puzzling features.

Crigler-Najjar Syndrome, Type I CN-I is characterized by striking unconjugated hyperbilirubinemia of about 340–765 $\mu\text{mol/L}$ (20–45 mg/dL) that appears in the neonatal period and persists for life. Other conventional hepatic biochemical tests such as serum aminotransferases and alkaline phosphatase are normal, and there is no evidence of hemolysis. Hepatic histology is also essentially normal except for the occasional presence of bile plugs within canaliculi. Bilirubin glucuronides are virtually absent from the bile, and there is no detectable constitutive expression of UGT1A1 activity in hepatic tissue. Neither UGT1A1 activity nor the serum bilirubin concentration responds to administration of phenobarbital or other enzyme inducers. In the absence of conjugation, unconjugated bilirubin accumulates in plasma, from which it is eliminated very slowly by alternative pathways that include direct passage into the bile and small intestine. These account for the small amounts of urobilinogen found in feces. No bilirubin is found in the urine. First described in 1952, the disorder is rare (estimated prevalence, 0.6–1.0 per million). Many patients are from geographically or socially isolated communities in which consanguinity is common, and pedigree analyses show an autosomal recessive pattern of inheritance. The majority of patients (type IA) exhibit defects in the glucuronide conjugation of a spectrum of substrates in addition to bilirubin, including various drugs and other xenobiotics. These individuals have mutations in one of the common exons (2–5) of the *UGT1* gene (Fig. 359-2). In a smaller subset (type IB), the defect is limited largely to bilirubin conjugation, and the causative mutation is in the bilirubin-specific exon A1. Estrogen glucuronidation is mediated by UGT1A1 and is defective in all CN-I patients. More than 30 different genetic lesions of *UGT1A1* responsible for CN-I have been identified,

TABLE 359-1 PRINCIPAL DIFFERENTIAL CHARACTERISTICS OF GILBERT AND CRIGLER-NAJJAR SYNDROMES

Feature	Crigler-Najjar Syndrome		Gilbert Syndrome
	Type I	Type II	
Total serum bilirubin, $\mu\text{mol/L}$ (mg/dL)	310–755 (usually >345) (18–45 [usually >20])	100–430 (usually \leq 345) (6–25 [usually \leq 20])	Typically \leq 70 $\mu\text{mol/L}$ (\leq 4 mg/dL) in absence of fasting or hemolysis
Routine liver tests	Normal	Normal	Normal
Response to phenobarbital	None	Decreases bilirubin by >25%	Decreases bilirubin to normal
Kernicterus	Usual	Rare	No
Hepatic histology	Normal	Normal	Usually normal; increased lipofuscin pigment in some
Bile characteristics			
Color	Pale or colorless	Pigmented	Normal dark color
Bilirubin fractions	>90% unconjugated	Largest fraction (mean: 57%) monoconjugates	Mainly diconjugates but monoconjugates increased (mean: 23%)
Bilirubin UDP-glucuronosyltransferase activity	Typically absent; traces in some patients	Markedly reduced: 0–10% of normal	Reduced: typically 10–33% of normal
Inheritance (all autosomal)	Recessive	Predominantly recessive	Promoter mutation: recessive Missense mutations: 7 of 8 dominant; 1 reportedly recessive

2002 including deletions, insertions, alterations in intron splice donor and acceptor sites, exon skipping, and point mutations that introduce premature stop codons or alter critical amino acids. Their common feature is that they all encode proteins with absent or, at most, traces of bilirubin-UDP-glucuronosyltransferase enzymatic activity.

Prior to the availability of phototherapy, most patients with CN-I died of bilirubin encephalopathy (*kernicterus*) in infancy or early childhood. A few lived as long as early adult life without overt neurologic damage, although more subtle testing usually indicated mild but progressive brain damage. In the absence of liver transplantation, death eventually supervened from late-onset bilirubin encephalopathy, which often followed a nonspecific febrile illness. Although isolated hepatocyte transplantation has been used in a small number of cases of CN-I, early liver transplantation (**Chap. 368**) remains the best hope to prevent brain injury and death.

Crigler-Najjar Syndrome, Type II (CN-II) This condition was recognized as a distinct entity in 1962 and is characterized by marked unconjugated hyperbilirubinemia in the absence of abnormalities of other conventional hepatic biochemical tests, hepatic histology, or hemolysis. It differs from CN-I in several specific ways (Table 359-1): (1) Although there is considerable overlap, average bilirubin concentrations are lower in CN-II; (2) accordingly, CN-II is only infrequently associated with kernicterus; (3) bile is deeply colored, and bilirubin glucuronides are present, with a striking, characteristic increase in the proportion of monoglucuronides; (4) UGT1A1 in liver is usually present at reduced levels (typically $\leq 10\%$ of normal) but may be undetectable by older, less sensitive assays; and (5) while typically detected in infancy, hyperbilirubinemia was not recognized in some cases until later in life and, in one instance, at age 34. As with CN-I, most CN-II cases exhibit abnormalities in the conjugation of other compounds, such as salicylamide and menthol, but in some instances, the defect appears limited to bilirubin. Reduction of serum bilirubin concentrations by $>25\%$ in response to enzyme inducers such as phenobarbital distinguishes CN-II from CN-I, although this response may not be elicited in early infancy and often is not accompanied by measurable UGT1A1 induction. Bilirubin concentrations during phenobarbital administration do not return to normal but are typically in the range of 51–86 $\mu\text{mol/L}$ (3–5 mg/dL). Although the incidence of kernicterus in CN-II is low, instances have occurred, not only in infants but also in adolescents and adults, often in the setting of an intercurrent illness, fasting, or another factor that temporarily raises the serum bilirubin concentration above baseline and reduces serum albumin levels. For this reason, phenobarbital therapy is widely recommended, a single bedtime dose often sufficing to maintain clinically safe serum bilirubin concentrations.

Over 77 different mutations in the *UGT1* gene have been identified as causing CN-I or CN-II. It was found that missense mutations are more common in CN-II patients, as would be expected in this less severe phenotype. Their common feature is that they encode for a bilirubin-UDP-glucuronosyltransferase with markedly reduced, but detectable, enzymatic activity. The spectrum of residual enzyme activity explains the spectrum of phenotypic severity of the resulting hyperbilirubinemia. Molecular analysis has established that a large majority of CN-II patients are either homozygotes or compound heterozygotes for CN-II mutations and that individuals carrying one mutated and one entirely normal allele have normal bilirubin concentrations.

Gilbert Syndrome (GS) This syndrome is characterized by mild unconjugated hyperbilirubinemia, normal values for standard hepatic biochemical tests, and normal hepatic histology other than a modest increase of lipofuscin pigment in some patients. Serum bilirubin concentrations are most often $<51 \mu\text{mol/L}$ ($<3 \text{ mg/dL}$), although both higher and lower values are frequent. The clinical spectrum of hyperbilirubinemia fades into that of CN-II at serum bilirubin concentrations of 86–136 $\mu\text{mol/L}$ (5–8 mg/dL). At the other end of the scale, the distinction between mild cases of GS and a normal state is often blurred. Bilirubin concentrations may fluctuate substantially in any given individual, and at least 25% of patients will exhibit temporarily normal values during prolonged follow-up. More elevated values

are associated with stress, fatigue, alcohol use, reduced caloric intake, and intercurrent illness, while increased caloric intake or administration of enzyme-inducing agents produces lower bilirubin levels. GS is most often diagnosed at or shortly after puberty or in adult life during routine examinations that include multichannel biochemical analyses. UGT1A1 activity is typically reduced to 10–35% of normal, and bile pigments exhibit a characteristic increase in bilirubin monoglucuronides. Studies of radiobilirubin kinetics indicate that hepatic bilirubin clearance is reduced to an average of one-third of normal. Administration of phenobarbital normalizes both the serum bilirubin concentration and hepatic bilirubin clearance; however, failure of UGT1A1 activity to improve in many such instances suggests the possible coexistence of an additional defect. Compartmental analysis of bilirubin kinetic data suggests that GS patients have a defect in bilirubin uptake as well as in conjugation. Defect(s) in the hepatic uptake of other organic anions that at least partially share an uptake mechanism with bilirubin, such as sulfobromophthalein and indocyanine green (ICG), are observed in a minority of patients. The metabolism and transport of bile acids that do not utilize the bilirubin uptake mechanism are normal. The magnitude of changes in the serum bilirubin concentration induced by provocation tests such as 48 hours of fasting or the IV administration of nicotinic acid have been reported to be of help in separating GS patients from normal individuals. Other studies dispute this assertion. Moreover, on theoretical grounds, the results of such studies should provide no more information than simple measurements of the baseline serum bilirubin concentration. Family studies indicate that GS and hereditary hemolytic anemias such as hereditary spherocytosis, glucose-6-phosphate dehydrogenase deficiency, and β -thalassemia trait sort independently. Reports of hemolysis in up to 50% of GS patients are believed to reflect better case finding, since patients with both GS and hemolysis have higher bilirubin concentrations, and are more likely to be jaundiced, than patients with either defect alone.

GS is common, with many series placing its prevalence at $\geq 8\%$. Males predominate over females by reported ratios ranging from 1.5:1 to $>7:1$. However, these ratios may have a large artifactual component since normal males have higher mean bilirubin levels than normal females, but the diagnosis of GS is often based on comparison to normal ranges established in men. The high prevalence of GS in the general population may explain the reported frequency of mild unconjugated hyperbilirubinemia in liver transplant recipients. The disposition of most xenobiotics metabolized by glucuronidation appears to be normal in GS, as is oxidative drug metabolism in the majority of reported studies. The principal exception is the metabolism of the antitumor agent irinotecan (CPT-11), whose active metabolite (SN-38) is glucuronidated specifically by bilirubin-UDP-glucuronosyltransferase. Administration of CPT-11 to patients with GS has resulted in several toxicities, including intractable diarrhea and myelosuppression. Some reports also suggest abnormal disposition of menthol, estradiol benzoate, acetaminophen, tolbutamide, and rifamycin SV. Although some of these studies have been disputed, and there have been no reports of clinical complications from use of these agents in GS, prudence should be exercised in prescribing them, or any agents metabolized primarily by glucuronidation, in this condition. It should also be noted that the HIV protease inhibitors indinavir and atazanavir (**Chap. 226**) can inhibit UGT1A1, resulting in hyperbilirubinemia that is most pronounced in patients with preexisting GS.

Most older pedigree studies of GS were consistent with autosomal dominant inheritance with variable expressivity. However, studies of the *UGT1* gene in GS have indicated a variety of molecular genetic bases for the phenotypic picture and several different patterns of inheritance. Studies in Europe and the United States found that nearly all patients had normal coding regions for UGT1A1 but were homozygous for the insertion of an extra TA (i.e., $A[\text{TA}]_7\text{TAA}$ rather than $A[\text{TA}]_6\text{TAA}$) in the promoter region of the first exon. This appeared to be necessary, but not sufficient, for clinically expressed GS, since 15% of normal controls were also homozygous for this variant. While normal by standard criteria, these individuals had somewhat higher bilirubin concentrations

than the rest of the controls studied. Heterozygotes for this abnormality had bilirubin concentrations identical to those homozygous for the normal A[TA]₆TAA allele. The prevalence of the A[TA]₆TAA allele in a general Western population is 30%, in which case 9% would be homozygotes. This is slightly higher than the prevalence of GS based on purely phenotypic parameters. It was suggested that additional variables, such as mild hemolysis or a defect in bilirubin uptake, might be among the factors enhancing phenotypic expression of the defect.

Phenotypic expression of GS due solely to the A[TA]₇TAA promoter abnormality is inherited as an autosomal recessive trait. A number of CN-II kindreds have been identified in whom there is also an allele containing a normal coding region but the A[TA]₇TAA promoter abnormality. CN-II heterozygotes who have the A[TA]₆TAA promoter are phenotypically normal, whereas those with the A[TA]₇TAA promoter express the phenotypic picture of GS. GS in such kindreds may also result from homozygosity for the A[TA]₇TAA promoter abnormality. Seven different missense mutations in the *UGT1* gene that reportedly cause GS with dominant inheritance have been found in Japanese individuals. Another Japanese patient with mild unconjugated hyperbilirubinemia was homozygous for a missense mutation in exon 5. GS in her family appeared to be recessive. Missense mutations causing GS have not been reported outside of certain Asian populations.

DISORDERS OF BILIRUBIN METABOLISM LEADING TO MIXED OR PREDOMINANTLY CONJUGATED HYPERBILIRUBINEMIA

In hyperbilirubinemia due to acquired liver disease (e.g., acute hepatitis, common bile duct stone), there are usually elevations in the serum concentrations of both conjugated and unconjugated bilirubin. Although biliary tract obstruction or hepatocellular cholestatic injury may present on occasion with a predominantly conjugated hyperbilirubinemia, it is generally not possible to differentiate intrahepatic from extrahepatic causes of jaundice based on the serum levels or relative proportions of unconjugated and conjugated bilirubin. The major reason for determining the amounts of conjugated and unconjugated bilirubin in the serum is for the initial differentiation of hepatic parenchymal and obstructive disorders (mixed conjugated and unconjugated hyperbilirubinemia) from the inheritable and hemolytic disorders discussed above that are associated with unconjugated hyperbilirubinemia.

FAMILIAL DEFECTS IN HEPATIC EXCRETORY FUNCTION

Dubin-Johnson Syndrome (DJS) This benign, relatively rare disorder is characterized by low-grade, predominantly conjugated hyperbilirubinemia (Table 359-2). Total bilirubin concentrations are typically between 34 and 85 $\mu\text{mol/L}$ (2 and 5 mg/dL) but on occasion can be in the normal range or as high as 340–430 $\mu\text{mol/L}$ (20–25 mg/dL) and

can fluctuate widely in any given patient. The degree of hyperbilirubinemia may be increased by intercurrent illness, oral contraceptive use, and pregnancy. Because the hyperbilirubinemia is due to a predominant rise in conjugated bilirubin, bilirubinuria is characteristically present. Aside from elevated serum bilirubin levels, other routine laboratory tests are normal. Physical examination is usually normal except for jaundice, although an occasional patient may have hepatosplenomegaly.

Patients with DJS are usually asymptomatic, although some may have vague constitutional symptoms. These latter patients have usually undergone extensive and often unnecessary diagnostic examinations for unexplained jaundice and have high levels of anxiety. In women, the condition may be subclinical until the patient becomes pregnant or receives oral contraceptives, at which time chemical hyperbilirubinemia becomes frank jaundice. Even in these situations, other routine liver function tests, including serum alkaline phosphatase and transaminase activities, are normal.

A cardinal feature of DJS is the accumulation in the lysosomes of centrilobular hepatocytes of dark, coarsely granular pigment. As a result, the liver may be grossly black in appearance. This pigment is thought to be derived from epinephrine metabolites that are not excreted normally. The pigment may disappear during bouts of viral hepatitis, only to reaccumulate slowly after recovery.

Biliary excretion of a number of anionic compounds is compromised in DJS. These include various cholecystographic agents, as well as sulfobromophthalein (Bromsulphalein, BSP), a synthetic dye formerly used in a test of liver function. In this test, the rate of disappearance of BSP from plasma was determined following bolus IV administration. BSP is conjugated with glutathione in the hepatocyte; the resulting conjugate is normally excreted rapidly into the bile canaliculus. Patients with DJS exhibit characteristic rises in plasma concentrations at 90 minutes after injection, due to reflux of conjugated BSP into the circulation from the hepatocyte. Dyes such as ICG that are taken up by hepatocytes but are not further metabolized prior to biliary excretion do not show this reflux phenomenon. Continuous BSP infusion studies suggest a reduction in the time to maximum plasma concentration (t_{max}) for biliary excretion. Bile acid disposition, including hepatocellular uptake and biliary excretion, is normal in DJS. These patients have normal serum and biliary bile acid concentrations and do not have pruritus.

By analogy with findings in several mutant rat strains, the selective defect in biliary excretion of bilirubin conjugates and certain other classes of organic compounds, but not of bile acids, that characterizes DJS in humans was found to reflect defective expression of MRP2, an ATP-dependent canalicular membrane transporter. Several different mutations in the *MRP2* gene produce the Dubin-Johnson phenotype, which has an autosomal recessive pattern of inheritance. Although

TABLE 359-2 PRINCIPAL DIFFERENTIAL CHARACTERISTICS OF INHERITABLE DISORDERS OF BILE CANALICULAR FUNCTION

	DJS	Rotor	PFIC1	BRIC1	PFIC2	BRIC2	PFIC3
Gene	<i>ABCCA</i>	<i>SLCO1B1/SLCO1B3</i>	<i>ATP8B1</i>	<i>ATP8B1</i>	<i>ABCB11</i>	<i>ABCB11</i>	<i>ABCB4</i>
Protein	MRP2	OATP1B1/1B3	FIC1	FIC1	BSEP	BSEP	MDR3
Cholestasis	No	No	Yes	Episodic	Yes	Episodic	Yes
Serum γ -GT	Normal	Normal	Normal	Normal	Normal	Normal	↑↑
Serum bile acids	Normal	Normal	↑↑	↑↑ during episodes	↑↑	↑↑ during episodes	↑↑
Clinical features	Mild conjugated hyperbilirubinemia; otherwise normal liver function; dark pigment in liver; characteristic pattern of urinary coproporphyrins	Mild conjugated hyperbilirubinemia; otherwise normal liver function; liver without abnormal pigmentation	Severe cholestasis beginning in childhood	Recurrent episodes of cholestasis beginning at any age	Severe cholestasis beginning in childhood	Recurrent episodes of cholestasis beginning at any age	Severe cholestasis beginning in childhood; decreased phospholipids in bile

Abbreviations: BRIC, benign recurrent intrahepatic cholestasis; BSEP, bile salt excretory protein; DJS, Dubin-Johnson syndrome; γ -GT, γ -glutamyltransferase; MRP2, multidrug resistance-associated protein 2; OATP1A/1B, organic anion transport proteins 1B1 and 1B3; PFIC, progressive familial intrahepatic cholestasis; ↑↑, increased.

2004 MRP2 is undoubtedly important in the biliary excretion of conjugated bilirubin, the fact that this pigment is still excreted in the absence of MRP2 suggests that other, as yet uncharacterized, transport proteins may serve in a secondary role in this process.

Patients with DJS also have a diagnostic abnormality in urinary coproporphyrin excretion. There are two naturally occurring coproporphyrin isomers, I and III. Normally, ~75% of the coproporphyrin in urine is isomer III. In urine from DJS patients, total coproporphyrin content is normal, but >80% is isomer I. Heterozygotes for the syndrome show an intermediate pattern. The molecular basis for this phenomenon remains unclear.

Rotor Syndrome This benign, autosomal recessive disorder is clinically similar to DJS (Table 359-2), although it is seen even less frequently. A major phenotypic difference is that the liver in patients with Rotor syndrome has no increased pigmentation and appears totally normal. The only abnormality in routine laboratory tests is an elevation of total serum bilirubin, due to a predominant rise in conjugated bilirubin. This is accompanied by bilirubinuria. Several additional features differentiate Rotor syndrome from DJS. In Rotor syndrome, the gallbladder is usually visualized on oral cholecystography, in contrast to the nonvisualization that is typical of DJS. The pattern of urinary coproporphyrin excretion also differs. The pattern in Rotor syndrome resembles that of many acquired disorders of hepatobiliary function, in which coproporphyrin I, the major coproporphyrin isomer in bile, refluxes from the hepatocyte back into the circulation and is excreted in urine. Thus, total urinary coproporphyrin excretion is substantially increased in Rotor syndrome, in contrast to the normal levels seen in DJS. Although the fraction of coproporphyrin I in urine is elevated, it is usually <70% of the total, compared with ≥80% in DJS. The disorders also can be distinguished by their patterns of BSP excretion. Although clearance of BSP from plasma is delayed in Rotor syndrome, there is no reflux of conjugated BSP back into the circulation as seen in DJS. Kinetic analysis of plasma BSP infusion studies suggests the presence of a defect in intrahepato cellular storage of this compound. This has never been demonstrated directly. Recent studies indicate that the molecular basis of Rotor syndrome results from simultaneous deficiency of the plasma membrane transporters OATP1B1 and OATP1B3. This results in reduced reuptake of conjugated bilirubin that has been pumped out of the cell into the portal circulation by MRP3 (Fig. 359-1).

Benign Recurrent Intrahepatic Cholestasis (BRIC) This rare disorder is characterized by recurrent attacks of pruritus and jaundice. The typical episode begins with mild malaise and elevations in serum aminotransferase levels, followed rapidly by rises in alkaline phosphatase and conjugated bilirubin and onset of jaundice and itching. The first one or two episodes may be misdiagnosed as acute viral hepatitis. The cholestatic episodes, which may begin in childhood or adulthood, can vary in duration from several weeks to months, followed by a complete clinical and biochemical resolution. Intervals between attacks may vary from several months to years. Between episodes, physical examination is normal, as are serum levels of bile acids, bilirubin, transaminases, and alkaline phosphatase. The disorder is familial and has an autosomal recessive pattern of inheritance. BRIC is considered a benign disorder in that it does not lead to cirrhosis or end-stage liver disease. However, the episodes of jaundice and pruritus can be prolonged and debilitating, and some patients have undergone liver transplantation to relieve the intractable and disabling symptoms. Treatment during the cholestatic episodes is symptomatic; there is no specific treatment to prevent or shorten the occurrence of episodes.

A gene termed *FIC1* was recently identified and found to be mutated in patients with BRIC. Curiously, this gene is expressed strongly in the small intestine but only weakly in the liver. The protein encoded by *FIC1* shows little similarity to those that have been shown to play a role in bile canalicular excretion of various compounds. Rather, it appears to be a member of a P-type ATPase family that transports aminophospholipids from the outer to the inner leaflet of a variety of cell membranes. Its relationship to the pathobiology of this disorder

remains unclear. A second phenotypically identical form of BRIC, termed BRIC type 2, has been described resulting from mutations in the bile salt excretory protein (BSEP), the protein that is defective in progressive familial intrahepatic cholestasis type 2 (Table 359-2). How some mutations in this protein result in the episodic BRIC phenotype is unknown.

Progressive Familial Intrahepatic Cholestasis (FIC) This name is applied to three phenotypically related syndromes (Table 359-2). Progressive FIC type 1 (Byler disease) presents in early infancy as cholestasis that may be initially episodic. However, in contrast to BRIC, Byler disease progresses to malnutrition, growth retardation, and end-stage liver disease during childhood. This disorder is also a consequence of a *FIC1* mutation. The functional relationship of the *FIC1* protein to the pathogenesis of cholestasis in these disorders is unknown. Two other types of progressive FIC (types 2 and 3) have been described. Progressive FIC type 2 is associated with a mutation in the protein originally named *sister of p-glycoprotein*, now known as *bile salt excretory protein*, which is the major bile canalicular exporter of bile acids. As noted above, some mutations of this protein are associated with BRIC type 2, rather than the progressive FIC type 2 phenotype. Progressive FIC type 3 has been associated with a mutation of MDR3, a protein that is essential for normal hepatocellular excretion of phospholipids across the bile canaliculus. Although all three types of progressive FIC have similar clinical phenotypes, only type 3 is associated with high serum levels of γ -glutamyltransferase activity. In contrast, activity of this enzyme is normal or only mildly elevated in symptomatic BRIC and progressive FIC types 1 and 2.

360

Acute Viral Hepatitis

Jules L. Dienstag

Acute viral hepatitis is a systemic infection affecting the liver predominantly. Almost all cases of acute viral hepatitis are caused by one of five viral agents: hepatitis A virus (HAV), hepatitis B virus (HBV), hepatitis C virus (HCV), the HBV-associated delta agent or hepatitis D virus (HDV), and hepatitis E virus (HEV). All these human hepatitis viruses are RNA viruses, except for hepatitis B, which is a DNA virus but replicates like a retrovirus. Although these agents can be distinguished by their molecular and antigenic properties, all types of viral hepatitis produce clinically similar illnesses. These range from asymptomatic and inapparent to fulminant and fatal acute infections common to all types, on the one hand, and from subclinical persistent infections to rapidly progressive chronic liver disease with cirrhosis and even hepatocellular carcinoma, common to the bloodborne types (HBV, HCV, and HDV), on the other.

VIROLOGY AND ETIOLOGY

Hepatitis A HAV is a nonenveloped 27-nm, heat-, acid-, and ether-resistant RNA virus in the *Hepatovirus* genus of the picornavirus family (Fig. 360-1). Its virion contains four capsid polypeptides, designated VP1 to VP4, which are cleaved posttranslationally from the polyprotein product of a 7500-nucleotide genome. Inactivation of viral activity can be achieved by boiling for 1 min, by contact with formaldehyde and chlorine, or by ultraviolet irradiation. Despite nucleotide sequence variation of up to 20% among isolates of HAV, and despite the recognition of four genotypes affecting humans, all strains of this virus are immunologically indistinguishable and belong to one serotype. Hepatitis A has an incubation period of ~4 weeks. Its replication is limited to the liver, but the virus is present in the liver, bile, stools, and blood during the late incubation period and acute preicteric/presymptomatic phase of illness. Despite slightly longer persistence of virus in the liver, fecal shedding, viremia, and infectivity diminish rapidly once jaundice becomes apparent. HAV can be cultivated reproducibly in vitro.

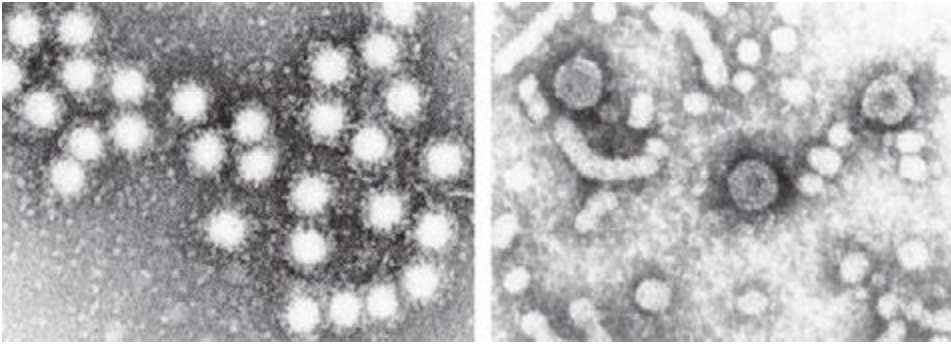


FIGURE 360-1 Electron micrographs of hepatitis A virus particles and serum from a patient with hepatitis B. **Left:** 27-nm hepatitis A virus particles purified from stool of a patient with acute hepatitis A and aggregated by antibody to hepatitis A virus. **Right:** Concentrated serum from a patient with hepatitis B, demonstrating the 42-nm virions, tubular forms, and spherical 22-nm particles of hepatitis B surface antigen. 132,000 \times . (Hepatitis D resembles 42-nm virions of hepatitis B but is smaller, 35–37 nm; hepatitis E resembles hepatitis A virus but is slightly larger, 32–34 nm; hepatitis C has been visualized as a 55-nm particle.)

Antibodies to HAV (anti-HAV) can be detected during acute illness when serum aminotransferase activity is elevated and fecal HAV shedding is still occurring. This early antibody response is predominantly of the IgM class and persists for several (~3) months, rarely for 6–12 months. During convalescence, however, anti-HAV of the IgG class becomes the predominant antibody (Fig. 360-2). Therefore, the diagnosis of hepatitis A is made during acute illness by demonstrating anti-HAV of the IgM class. After acute illness, anti-HAV of the IgG class remains detectable indefinitely, and patients with serum anti-HAV are immune to reinfection. Neutralizing antibody activity parallels the appearance of anti-HAV, and the IgG anti-HAV present in immune globulin accounts for the protection it affords against HAV infection.

Hepatitis B HBV is a DNA virus with a remarkably compact genomic structure; despite its small, circular, 3200-bp size, HBV DNA codes for four sets of viral products with a complex, multiparticulate structure. HBV achieves its genomic economy by relying on an efficient strategy of encoding proteins from four overlapping genes: S, C, P, and X (Fig. 360-3), as detailed below. Once thought to be unique among viruses, HBV is now recognized as one of a family of animal viruses, hepadnaviruses (hepatotropic DNA viruses), and is classified as hepadnavirus type 1. Similar viruses infect certain species of woodchucks, ground and tree squirrels, and Pekin ducks, to mention the most carefully characterized. Like HBV, all have the same distinctive three morphologic forms, have counterparts to the envelope and nucleocapsid virus antigens of HBV, replicate in the liver but exist in extrahepatic sites, contain their own endogenous DNA polymerase, have partially double-strand and partially single-strand genomes, are associated with acute and chronic hepatitis and hepatocellular carcinoma, and rely on a replicative strategy unique among DNA viruses but typical of retroviruses. Instead of DNA replication directly from a DNA

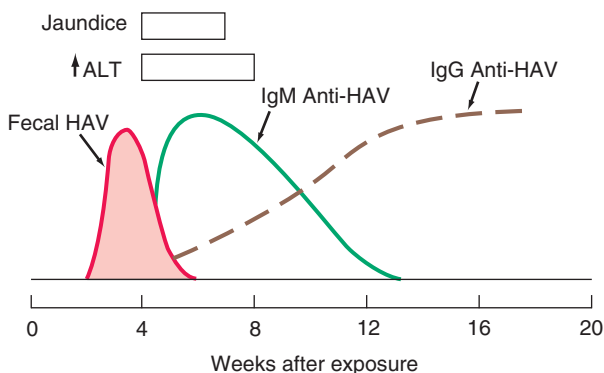


FIGURE 360-2 Scheme of typical clinical and laboratory features of hepatitis A virus (HAV). ALT, alanine aminotransferase.

template, hepadnaviruses rely on reverse transcription (effected by the DNA polymerase) of minus-strand DNA from a “pregenomic” RNA intermediate. Then plus-strand DNA is transcribed from the minus-strand DNA template by the DNA-dependent DNA polymerase and converted in the hepatocyte nucleus to a covalently closed circular DNA, which serves as a template for messenger RNA and pregenomic RNA. Viral proteins are translated by the messenger RNA, and the proteins and genome are packaged into virions and secreted from the hepatocyte. Although HBV is difficult to cultivate in vitro in the conventional sense from clinical material, several cell lines have been transfected with HBV DNA. Such transfected cells support in vitro replication of the intact virus and its component proteins.

VIRAL PROTEINS AND PARTICLES Of the three particulate forms of HBV (Table 360-1), the most numerous are the 22-nm particles, which appear as spherical or long filamentous forms; these are antigenically indistinguishable from the outer surface or envelope protein of HBV and are thought to represent excess viral envelope protein. Outnumbered in serum by a factor of 100 or 1000 to 1 compared with the spheres and tubules are large, 42-nm, double-shelled spherical particles, which represent the intact hepatitis B virion (Fig. 360-1). The envelope protein expressed on the outer surface of the virion and on the smaller spherical and tubular structures is referred to as *hepatitis B surface antigen* (HBsAg). The concentration of HBsAg and virus particles in the blood may reach 500 $\mu\text{g}/\text{mL}$ and 10 trillion particles per milliliter, respectively. The envelope protein, HBsAg, is the product of the S gene of HBV.

Envelope HBsAg subdeterminants include a common group-reactive antigen, *a*, shared by all HBsAg isolates and one of several subtype-specific antigens—*d* or *y*, *w* or *r*—as well as other specificities. Hepatitis B isolates fall into one of at least eight subtypes and ten

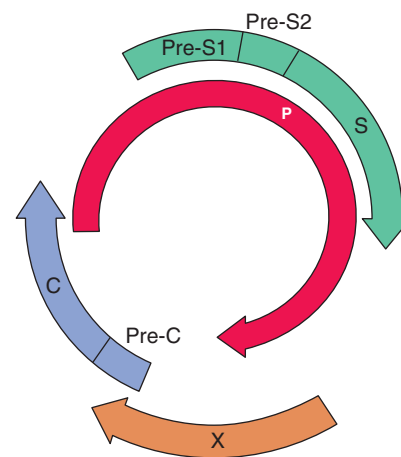


FIGURE 360-3 Compact genomic structure of hepatitis B virus (HBV). This structure, with overlapping genes, permits HBV to code for multiple proteins. The S gene codes for the “major” envelope protein, HBsAg. Pre-S1 and pre-S2, upstream of S, combine with S to code for two larger proteins, “middle” protein, the product of pre-S2 + S, and “large” protein, the product of pre-S1 + pre-S2 + S. The largest gene, P, codes for DNA polymerase. The C gene codes for two nucleocapsid proteins, HBeAg, a soluble, secreted protein (initiation from the pre-C region of the gene), and HBcAg, the intracellular core protein (initiation after pre-C). The X gene codes for HBxAg, which can transactivate the transcription of cellular and viral genes; its clinical relevance is not known, but it may contribute to carcinogenesis by binding to p53.

TABLE 360-1 NOMENCLATURE AND FEATURES OF HEPATITIS VIRUSES

Hepatitis Type	Virus Particle, nm	Morphology	Genome ^a	Classification	Antigen(s)	Antibodies	Remarks
HAV	27	Icosahedral nonenveloped	7.5-kb RNA, linear, ss, +	Hepatovirus	HAV	Anti-HAV	Early fecal shedding Diagnosis: IgM anti-HAV Previous infection: IgG anti-HAV
HBV	42	Double-shelled virion (surface and core) spherical	3.2-kb DNA, circular, ss/ds	Hepadnavirus	HBsAg HBcAg HBeAg	Anti-HBs Anti-HBc Anti-HBe	Bloodborne virus; carrier state Acute diagnosis: HBsAg, IgM anti-HBc Chronic diagnosis: IgG anti-HBc, HBsAg Markers of replication: HBeAg, HBV DNA Liver, lymphocytes, other organs
	27	Nucleocapsid core			HBcAg HBeAg	Anti-HBc Anti-HBe	Nucleocapsid contains DNA and DNA polymerase; present in hepatocyte nucleus; HBcAg does not circulate; HBeAg (soluble, nonparticulate) and HBV DNA circulate—correlate with infectivity and complete virions
	22	Spherical and filamentous; represents excess virus coat material			HBsAg	Anti-HBs	HBsAg detectable in >95% of patients with acute hepatitis B; found in serum, body fluids, hepatocyte cytoplasm; anti-HBs appears following infection—protective antibody
HCV	Approx. 50–80	Enveloped	9.4-kb RNA, linear, ss, +	Hepacivirus	HCV C100-3 C33c C22-3 NS5	Anti-HCV	Bloodborne agent, formerly labeled non-A, non-B hepatitis Acute diagnosis: anti-HCV (C33c, C22-3, NS5), HCV RNA Chronic diagnosis: anti-HCV (C100-3, C33c, C22-3, NS5) and HCV RNA; cytoplasmic location in hepatocytes
HDV	35–37	Enveloped hybrid particle with HBsAg coat and HDV core	1.7-kb RNA, circular, ss, –	Resembles viroids and plant satellite viruses (genus Deltavirus)	HBsAg HDAG	Anti-HBs Anti-HDV	Defective RNA virus, requires helper function of HBV (hepadnaviruses); HDV antigen (HDAG) present in hepatocyte nucleus Diagnosis: anti-HDV, HDV RNA; HBV/HDV co-infection—IgM anti-HBc and anti-HDV; HDV superinfection—IgG anti-HBc and anti-HDV
HEV	32–34	Nonenveloped icosahedral	7.6-kb RNA, linear, ss, +	Hepevirus	HEV antigen	Anti-HEV	Agent of enterically transmitted hepatitis; rare in United States; occurs in Asia, Mediterranean countries, Central America Diagnosis: IgM/IgG anti-HEV (assays not routinely available); virus in stool, bile, hepatocyte cytoplasm

^ass, single-strand; ss/ds, partially single-strand, partially double-strand; –, minus-strand; +, plus-strand.

Note: See text for abbreviations.

genotypes (A–J). Geographic distribution of genotypes and subtypes varies; genotypes A (corresponding to subtype *adw*) and D (*ayw*) predominate in the United States and Europe, whereas genotypes B (*adw*) and C (*adr*) predominate in Asia. Clinical course and outcome are independent of subtype, but genotype B appears to be associated with less rapidly progressive liver disease and cirrhosis and a lower likelihood, or delayed appearance, of hepatocellular carcinoma than genotype C or D. Patients with genotype A are more likely to clear circulating viremia and to achieve HBeAg and HBsAg seroconversion, both spontaneously and in response to antiviral therapy. In addition, “precore” mutations are favored by certain genotypes (see below).

Upstream of the S gene are the pre-S genes (Fig. 360-3), which code for pre-S gene products, including receptors on the HBV surface for polymerized human serum albumin and for hepatocyte membrane proteins. The pre-S region actually consists of both pre-S1 and pre-S2. Depending on where translation is initiated, three potential HBsAg gene products are synthesized. The protein product of the S gene is HBsAg (*major protein*), the product of the S region plus the adjacent pre-S2 region is the *middle protein*, and the product of the pre-S1 plus pre-S2 plus S regions is the *large protein*. Compared with the smaller spherical and tubular particles of HBV, complete 42-nm virions are enriched in the large protein. Both pre-S proteins and their respective antibodies can be detected during HBV infection, and the period of pre-S antigenemia appears to coincide with other markers of virus replication, as detailed below; however, pre-S proteins have little clinical relevance and are not included in routine serologic testing repertoires.

The intact 42-nm virion contains a 27-nm nucleocapsid core particle. Nucleocapsid proteins are coded for by the C gene. The

antigen expressed on the surface of the nucleocapsid core is *hepatitis B core antigen* (HBcAg), and its corresponding antibody is anti-HBc. A third HBV antigen is *hepatitis B e antigen* (HBeAg), a soluble, nonparticulate, nucleocapsid protein that is immunologically distinct from intact HBcAg but is a product of the same C gene. The C gene has two initiation codons, a precore and a core region (Fig. 360-3). If translation is initiated at the precore region, the protein product is HBeAg, which has a signal peptide that binds it to the smooth endoplasmic reticulum, the secretory apparatus of the cell, leading to its secretion into the circulation. If translation begins at the core region, HBcAg is the protein product; it has no signal peptide, it is not secreted, but it assembles into nucleocapsid particles, which bind to and incorporate RNA, and which, ultimately, contain HBV DNA. Also packaged within the nucleocapsid core is a DNA polymerase, which directs replication and repair of HBV DNA. When packaging within viral proteins is complete, synthesis of the incomplete plus strand stops; this accounts for the single-strand gap and for differences in the size of the gap. HBcAg particles remain in the hepatocyte, where they are readily detectable by immunohistochemical staining and are exported after encapsidation by an envelope of HBsAg. Therefore, naked core particles do not circulate in the serum. The secreted nucleocapsid protein, HBeAg, provides a convenient, readily detectable, qualitative marker of HBV replication and relative infectivity.

HBsAg-positive serum containing HBeAg is more likely to be highly infectious and to be associated with the presence of hepatitis B virions (and detectable HBV DNA, see below) than HBeAg-negative or anti-HBe-positive serum. For example, HBsAg-positive mothers who are HBeAg-positive almost invariably (>90%) transmit hepatitis

B infection to their offspring, whereas HBsAg-positive mothers with anti-HBe rarely (10–15%) infect their offspring.

Early during the course of acute hepatitis B, HBeAg appears transiently; its disappearance may be a harbinger of clinical improvement and resolution of infection. Persistence of HBeAg in serum beyond the first 3 months of acute infection may be predictive of the development of chronic infection, and the presence of HBeAg during chronic hepatitis B tends to be associated with ongoing viral replication, infectivity, and inflammatory liver injury (except during the early decades after perinatally acquired HBV infection; see below).

The third and largest of the HBV genes, the P gene (Fig. 360-3), codes for HBV DNA polymerase; as noted above, this enzyme has both DNA-dependent DNA polymerase and RNA-dependent reverse transcriptase activities. The fourth gene, X, codes for a small, non-particulate protein, *hepatitis B x antigen* (HBxAg), that is capable of transactivating the transcription of both viral and cellular genes (Fig. 360-3). In the cytoplasm, HBxAg effects calcium release (possibly from mitochondria), which activates signal-transduction pathways that lead to stimulation of HBV reverse transcription and HBV DNA replication. Such transactivation may enhance the replication of HBV, leading to the clinical association observed between the expression of HBxAg and antibodies to it in patients with severe chronic hepatitis and hepatocellular carcinoma. The transactivating activity can enhance the transcription and replication of other viruses besides HBV, such as HIV. Cellular processes transactivated by X include the human interferon γ gene and class I major histocompatibility genes; potentially, these effects could contribute to enhanced susceptibility of HBV-infected hepatocytes to cytolytic T cells. The expression of X can also induce programmed cell death (apoptosis). The clinical relevance of HBxAg is limited, however, and testing for it is not part of routine clinical practice.

SEROLOGIC AND VIROLOGIC MARKERS After a person is infected with HBV, the first virologic marker detectable in serum within 1–12 weeks, usually between 8 and 12 weeks, is HBsAg (Fig. 360-4). Circulating HBsAg precedes elevations of serum aminotransferase activity and clinical symptoms by 2–6 weeks and remains detectable during the entire icteric or symptomatic phase of acute hepatitis B and beyond. In typical cases, HBsAg becomes undetectable 1–2 months after the onset of jaundice and rarely persists beyond 6 months. After HBsAg disappears, antibody to HBsAg (anti-HBs) becomes detectable in serum and remains detectable indefinitely thereafter. Because HBcAg is intracellular and, when in the serum, sequestered within an HBsAg coat, naked core particles do not circulate in serum, and therefore, HBcAg is not detectable routinely in the serum of patients with HBV infection. By contrast, anti-HBc is readily demonstrable in serum, beginning within the first 1–2 weeks after the appearance of HBsAg and preceding detectable levels of anti-HBs by weeks to months.

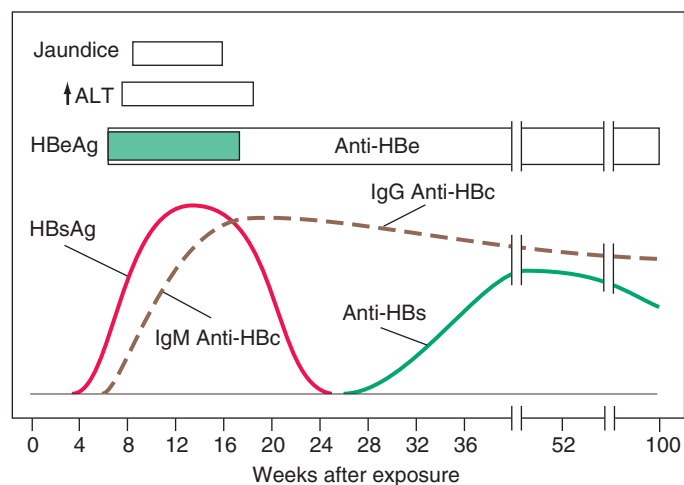


FIGURE 360-4 Scheme of typical clinical and laboratory features of acute hepatitis B. ALT, alanine aminotransferase.

Because variability exists in the time of appearance of anti-HBs after HBV infection, occasionally a gap of several weeks or longer may separate the disappearance of HBsAg and the appearance of anti-HBs. During this “gap” or “window” period, anti-HBc may represent the only serologic evidence of current or recent HBV infection, and blood containing anti-HBc in the absence of HBsAg and anti-HBs has been implicated in transfusion-associated hepatitis B. In part because the sensitivity of immunoassays for HBsAg and anti-HBs has increased, however, this window period is rarely encountered. In some persons, years after HBV infection, anti-HBc may persist in the circulation longer than anti-HBs. Therefore, isolated anti-HBc does not necessarily indicate active virus replication; most instances of isolated anti-HBc represent hepatitis B infection in the remote past. Rarely, however, isolated anti-HBc represents low-level hepatitis B viremia, with HBsAg below the detection threshold, and, occasionally, isolated anti-HBc represents a cross-reacting or false-positive immunologic specificity. Recent and remote HBV infections can be distinguished by determination of the immunoglobulin class of anti-HBc. Anti-HBc of the IgM class (IgM anti-HBc) predominates during the first 6 months after acute infection, whereas IgG anti-HBc is the predominant class of anti-HBc beyond 6 months. Therefore, patients with current or recent acute hepatitis B, including those in the anti-HBc window, have IgM anti-HBc in their serum. In patients who have recovered from hepatitis B in the remote past as well as those with chronic HBV infection, anti-HBc is predominantly of the IgG class. Infrequently, in ≤ 1 –5% of patients with acute HBV infection, levels of HBsAg are too low to be detected; in such cases, the presence of IgM anti-HBc establishes the diagnosis of acute hepatitis B. When isolated anti-HBc occurs in the rare patient with chronic hepatitis B whose HBsAg level is below the sensitivity threshold of contemporary immunoassays (a low-level carrier), anti-HBc is of the IgG class. Generally, in persons who have recovered from hepatitis B, anti-HBs and anti-HBc persist indefinitely.

The temporal association between the appearance of anti-HBs and resolution of HBV infection as well as the observation that persons with anti-HBs in serum are protected against reinfection with HBV suggests that *anti-HBs is the protective antibody*. Therefore, strategies for prevention of HBV infection are based on providing susceptible persons with circulating anti-HBs (see below). Occasionally, in $\sim 10\%$ of patients with chronic hepatitis B, low-level, low-affinity anti-HBs can be detected. This antibody is directed against a subtype determinant different from that represented by the patient’s HBsAg; its presence is thought to reflect the stimulation of a related clone of antibody-forming cells, but it has no clinical relevance and does not signal imminent clearance of hepatitis B. These patients with HBsAg and such nonneutralizing anti-HBs should be categorized as having chronic HBV infection.

The other readily detectable serologic marker of HBV infection, HBeAg, appears concurrently with or shortly after HBsAg. Its appearance coincides temporally with high levels of virus replication and reflects the presence of circulating intact virions and detectable HBV DNA (with the notable exception of patients with precore mutations who cannot synthesize HBeAg—see “Molecular Variants”). Pre-S1 and pre-S2 proteins are also expressed during periods of peak replication, but assays for these gene products are not routinely available. In self-limited HBV infections, HBeAg becomes undetectable shortly after peak elevations in aminotransferase activity, before the disappearance of HBsAg, and anti-HBe then becomes detectable, coinciding with a period of relatively lower infectivity (Fig. 360-4). Because markers of HBV replication appear transiently during acute infection, testing for such markers is of little clinical utility in typical cases of acute HBV infection. In contrast, markers of HBV replication provide valuable information in patients with protracted infections.

Departing from the pattern typical of acute HBV infections, in chronic HBV infection, HBsAg remains detectable beyond 6 months, anti-HBc is primarily of the IgG class, and anti-HBs is either undetectable or detectable at low levels (see “Laboratory Features”) (Fig. 360-5). During early chronic HBV infection, HBV DNA can be detected both in serum and in hepatocyte nuclei, where it is present in free or episomal form. This relatively highly *replicative stage* of HBV infection is

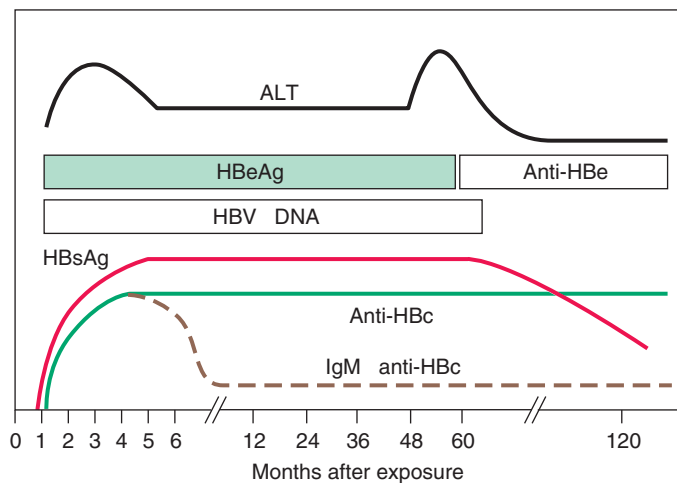


FIGURE 360-5 Scheme of typical laboratory features of wild-type chronic hepatitis B. HBeAg and hepatitis B virus (HBV) DNA can be detected in serum during the relatively *replicative phase* of chronic infection, which is associated with infectivity and liver injury. Seroconversion from the replicative phase to the relatively *nonreplicative phase* occurs at a rate of ~10% per year and is heralded by an acute hepatitis-like elevation of alanine aminotransferase (ALT) activity; during the nonreplicative phase, infectivity and liver injury are limited. In HBeAg-negative chronic hepatitis B associated with mutations in the precore region of the HBV genome, replicative chronic hepatitis B occurs in the absence of HBeAg.

the time of maximal infectivity and liver injury; HBeAg is a qualitative marker and HBV DNA a quantitative marker of this replicative phase, during which all three forms of HBV circulate, including intact virions. Over time, the relatively replicative phase of chronic HBV infection gives way to a relatively *nonreplicative phase*. This occurs at a rate of ~10% per year and is accompanied by seroconversion from HBeAg to anti-HBe. In many cases, this seroconversion coincides with a transient, usually mild, acute hepatitis-like elevation in aminotransferase activity, believed to reflect cell-mediated immune clearance of virus-infected hepatocytes. In the nonreplicative phase of chronic infection, when HBV DNA is demonstrable in hepatocyte nuclei, it tends to be integrated into the host genome. In this phase, only spherical and tubular forms of HBV, *not intact virions*, circulate, and liver injury tends to subside. Most such patients would be characterized as *inactive HBV carriers*. In reality, the designations *replicative* and *nonreplicative* are only relative; even in the so-called nonreplicative phase, HBV replication can be detected at levels of approximately $\leq 10^3$ virions with highly sensitive amplification probes such as the polymerase chain reaction (PCR); below this replication threshold, liver injury and infectivity of HBV are limited to negligible. Still, the distinctions are pathophysiologically and clinically meaningful. Occasionally, nonreplicative HBV infection converts back to replicative infection. Such spontaneous reactivations are accompanied by reexpression of HBeAg and HBV DNA, and sometimes of IgM anti-HBc, as well as by exacerbations of liver injury. Because high-titer IgM anti-HBc can reappear during acute exacerbations of chronic hepatitis B, relying on IgM anti-HBc versus IgG anti-HBc to distinguish between acute and chronic hepatitis B infection, respectively, may not always be reliable; in such cases, patient history is invaluable in helping to distinguish *de novo* acute hepatitis B infection from acute exacerbation of chronic hepatitis B infection.

MOLECULAR VARIANTS Variation occurs throughout the HBV genome, and clinical isolates of HBV that do not express typical viral proteins have been attributed to mutations in individual or even multiple gene locations. For example, variants have been described that lack nucleocapsid proteins (commonly), envelope proteins (very rarely), or both. Two categories of naturally occurring HBV variants have attracted the most attention. One of these was identified initially in Mediterranean countries among patients with severe chronic HBV infection and

detectable HBV DNA but with anti-HBe instead of HBeAg. These patients were found to be infected with an HBV mutant that contained an alteration in the precore region rendering the virus incapable of encoding HBeAg. Although several potential mutation sites exist in the pre-C region, the region of the C gene necessary for the expression of HBeAg (see “Virology and Etiology”), the most commonly encountered in such patients is a single base substitution, from G to A in the second to last codon of the pre-C gene at nucleotide 1896. This substitution results in the replacement of the TGG tryptophan codon by a stop codon (TAG), which prevents the translation of HBeAg. Another mutation, in the core-promoter region, prevents transcription of the coding region for HBeAg and yields an HBeAg-negative phenotype. Patients with such mutations in the precore region and who are unable to secrete HBeAg may have severe liver disease that progresses more rapidly to cirrhosis, or alternatively, they are identified clinically later in the course of the natural history of chronic hepatitis B, when the disease is more advanced. Both “wild-type” HBV and precore-mutant HBV can coexist in the same patient, or mutant HBV may arise late during wild-type HBV infection. In addition, clusters of fulminant hepatitis B in Israel and Japan were attributed to common-source infection with a precore mutant. Fulminant hepatitis B in North America and western Europe, however, occurs in patients infected with wild-type HBV, in the absence of precore mutants, and both precore mutants and other mutations throughout the HBV genome occur commonly, even in patients with typical, self-limited, milder forms of HBV infection. HBeAg-negative chronic hepatitis with mutations in the precore region is now the most frequently encountered form of hepatitis B in Mediterranean countries and in Europe. In the United States, where HBV genotype A (less prone to G1896A mutation) is prevalent, precore-mutant HBV is much less common; however, as a result of immigration from Asia and Europe, the proportion of HBeAg-negative hepatitis B-infected individuals has increased in the United States, and they now represent approximately 30–40% of patients with chronic hepatitis B. Characteristic of such HBeAg-negative chronic hepatitis B are lower levels of HBV DNA (usually $\leq 10^5$ IU/mL) and one of several patterns of aminotransferase activity—persistent elevations, periodic fluctuations above the normal range, and periodic fluctuations between the normal and elevated range.

The second important category of HBV mutants consists of *escape mutants*, in which a single amino acid substitution, from glycine to arginine, occurs at position 145 of the immunodominant *a* determinant common to all HBsAg subtypes. This HBsAg alteration leads to a critical conformational change that results in a loss of neutralizing activity by anti-HBs. This specific HBV/*a* mutant has been observed in two situations, active and passive immunization, in which humoral immunologic pressure may favor evolutionary change (“escape”) in the virus—in a small number of hepatitis B vaccine recipients who acquired HBV infection despite the prior appearance of neutralizing anti-HBs and in HBV-infected liver transplant recipients treated with a high-potency human monoclonal anti-HBs preparation. Although such mutants have not been recognized frequently, their existence raises a concern that may complicate vaccination strategies and serologic diagnosis.

Different types of mutations emerge during antiviral therapy of chronic hepatitis B with nucleoside analogues; such “YMDD” and similar mutations in the polymerase motif of HBV are described in Chap. 362.

EXTRAHEPATIC SITES Hepatitis B antigens and HBV DNA have been identified in extrahepatic sites, including lymph nodes, bone marrow, circulating lymphocytes, spleen, and pancreas. Although the virus does not appear to be associated with tissue injury in any of these extrahepatic sites, its presence in these “remote” reservoirs has been invoked (but is not necessary) to explain the recurrence of HBV infection after orthotopic liver transplantation. The clinical relevance of such extrahepatic HBV is limited.

Hepatitis D The delta hepatitis agent, or HDV, the only member of the genus *Deltavirus*, is a defective RNA virus that co-infects with and requires the helper function of HBV (or other hepadnaviruses)

for its replication and expression. Slightly smaller than HBV, HDV is a formalin-sensitive, 35- to 37-nm virus with a hybrid structure. Its nucleocapsid expresses HDV antigen (HDAG), which bears no antigenic homology with any of the HBV antigens, and contains the virus genome. The HDV core is “encapsidated” by an outer envelope of HBsAg, indistinguishable from that of HBV except in its relative compositions of major, middle, and large HBsAg component proteins. The genome is a small, 1700-nucleotide, circular, single-strand RNA of negative polarity that is nonhomologous with HBV DNA (except for a small area of the polymerase gene) but that has features and the rolling circle model of replication common to genomes of plant satellite viruses or viroids. HDV RNA contains many areas of internal complementarity; therefore, it can fold on itself by internal base pairing to form an unusual, very stable, rodlike structure that contains a very stable, self-cleaving and self-ligating ribozyme. HDV RNA requires host RNA polymerase II for its replication in the hepatocyte nucleus via RNA-directed RNA synthesis by transcription of genomic RNA to a complementary antigenomic (plus strand) RNA; the antigenomic RNA, in turn, serves as a template for subsequent genomic RNA synthesis effected by host RNA polymerase I. HDV RNA has only one open reading frame, and HDAG, a product of the antigenomic strand, is the only known HDV protein; HDAG exists in two forms: a small, 195-amino-acid species, which plays a role in facilitating HDV RNA replication, and a large, 214-amino-acid species, which appears to suppress replication but is required for assembly of the antigen into virions. HDV antigens have been shown to bind directly to RNA polymerase II, resulting in stimulation of transcription. Although complete hepatitis D virions and liver injury require the cooperative helper function of HBV, intracellular replication of HDV RNA can occur without HBV. Genomic heterogeneity among HDV isolates has been described; however, pathophysiologic and clinical consequences of this genetic diversity have not been recognized. The clinical spectrum of hepatitis D is common to all eight genotypes identified, the predominant of which is genotype 1.

HDV can either infect a person simultaneously with HBV (*co-infection*) or superinfect a person already infected with HBV (*super-infection*); when HDV infection is transmitted from a donor with one HBsAg subtype to an HBsAg-positive recipient with a different subtype, HDV assumes the HBsAg subtype of the recipient, rather than the donor. Because HDV relies absolutely on HBV, the duration of HDV infection is determined by the duration of (and cannot outlast) HBV infection. HDV replication tends to suppress HBV replication; therefore, patients with hepatitis D tend to have lower levels of HBV replication. HDV antigen is expressed primarily in hepatocyte nuclei and is occasionally detectable in serum. During acute HDV infection, anti-HDV of the IgM class predominates, and 30–40 days may elapse after symptoms appear before anti-HDV can be detected. In self-limited infection, anti-HDV is low-titer and transient, rarely remaining detectable beyond the clearance of HBsAg and HDV antigen. In chronic HDV infection, anti-HDV circulates in high titer, and both IgM and IgG anti-HDV can be detected. HDV antigen in the liver and HDV RNA in serum and liver can be detected during HDV replication.

Hepatitis C Hepatitis C virus, which, before its identification was labeled “non-A, non-B hepatitis,” is a linear, single-strand, positive-sense, 9600-nucleotide RNA virus, the genome of which is similar in organization to that of flaviviruses and pestiviruses; HCV is the only member of the genus *Hepacivirus* in the family Flaviviridae. The HCV genome contains a single, large open reading frame (gene) that codes for a

virus polyprotein of ~3000 amino acids, which is cleaved after translation to yield 10 viral proteins. The 5′ end of the genome consists of an untranslated region (containing an internal ribosomal entry site, IRES) adjacent to the genes for three structural proteins, the nucleocapsid core protein, C, and two structural envelope glycoproteins, E1 and E2. The 5′ untranslated region and core gene are highly conserved among genotypes, but the envelope proteins are coded for by the hypervariable region, which varies from isolate to isolate and may allow the virus to evade host immunologic containment directed at accessible virus-envelope proteins. The 3′ end of the genome also includes an untranslated region and contains the genes for seven nonstructural (NS) proteins, p7, NS2, NS3, NS4A, NS4B, NS5A, and NS5B. p7 is a membrane ion channel protein necessary for efficient assembly and release of HCV. The NS2 cysteine protease cleaves NS3 from NS2, and the NS3-4A serine protease cleaves all the downstream proteins from the polyprotein. Important NS proteins involved in virus replication include the NS3 helicase; NS3-4A serine protease; the multifunctional membrane-associated phosphoprotein NS5A, an essential component of the viral replication membranous web (along with NS4B); and the NS5B RNA-dependent RNA polymerase (Fig. 360-6). Because HCV does not replicate via a DNA intermediate, it does not integrate into the host genome. Because HCV tends to circulate in relatively low titer, 10^3 – 10^7 virions/mL, visualization of the 50- to 80-nm virus particles remains difficult. Still, the replication rate of HCV is very high, 10^{12} virions per day; its half-life is 2.7 h. The chimpanzee is a helpful but cumbersome animal model. Although a robust, reproducible, small animal model is lacking, HCV replication has been documented in an immunodeficient mouse model containing explants of human liver and in transgenic mouse and rat models. Although in vitro replication is difficult, replicons in hepatocellular carcinoma-derived cell lines support replication of genetically manipulated, truncated, or full-length HCV RNA (but not intact virions); infectious pseudotyped retroviral HCV particles have been shown to yield functioning envelope proteins. In 2005, complete replication of HCV and intact 55-nm virions were described in cell culture systems. HCV entry into the hepatocyte occurs via the nonliver-specific CD81 receptor and the liver-specific tight junction protein claudin-1. A growing list of additional host receptors to which HCV binds on cell entry includes occludin, low-density lipoprotein receptors, glycosaminoglycans, scavenger receptor B1, and epidermal growth factor receptor, among others. Relying on the same assembly and secretion pathway as low-density and very-low-density lipoproteins, HCV is a lipovirion and masquerades as a lipoprotein, which may limit its visibility to the adaptive immune system and which may explain its ability to evade immune containment and clearance. After viral entry and uncoating, translation is initiated by the IRES on the endoplasmic reticulum

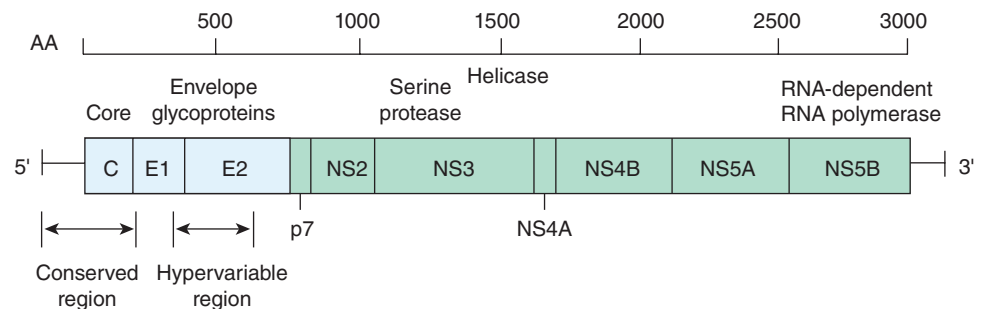


FIGURE 360-6 Organization of the hepatitis C virus genome and its associated, 3000-amino-acid (AA) proteins. The three structural genes at the 5′ end are the core region, C, which codes for the nucleocapsid, and the envelope regions, E1 and E2, which code for envelope glycoproteins. The 5′ untranslated region and the C region are highly conserved among isolates, whereas the envelope domain E2 contains the hypervariable region. At the 3′ end are seven nonstructural (NS) regions—p7, a membrane protein adjacent to the structural proteins that appears to function as an ion channel; NS2, which codes for a cysteine protease; NS3, which codes for a serine protease and an RNA helicase; NS4 and NS4B; NS5A, a multifunctional membrane-associated phosphoprotein, an essential component of the viral replication membranous web; and NS5B, which codes for an RNA-dependent RNA polymerase. After translation of the entire polyprotein, individual proteins are cleaved by both host and viral proteases.

2010 membrane, and the HCV polyprotein is cleaved during translation and posttranslationally by host cellular proteases as well as HCV NS2-3 and NS3-4A proteases. Host cofactors involved in HCV replication include cyclophilin A, which binds to NS5A and yields conformational changes required for viral replication, and liver-specific host microRNA miR-122.

At least six distinct major genotypes (and a minor genotype 7), as well as >50 subtypes within genotypes, of HCV have been identified by nucleotide sequencing. Genotypes differ from one another in sequence homology by $\geq 30\%$, and subtypes differ by approximately 20%. Because divergence of HCV isolates within a genotype or subtype and within the same host may vary insufficiently to define a distinct genotype, these intragenotypic differences are referred to as *quasispecies* and differ in sequence homology by only a few percent. The genotypic and quasispecies diversity of HCV, resulting from its high mutation rate, interferes with effective humoral immunity. Neutralizing antibodies to HCV have been demonstrated, but they tend to be short lived, and HCV infection does not induce lasting immunity against reinfection with different virus isolates or even the same virus isolate. Thus, neither *heterologous* nor *homologous* immunity appears to develop commonly after acute HCV infection. Some HCV genotypes are distributed worldwide, whereas others are more geographically confined (see “Epidemiology and Global Features”). In addition, differences exist among genotypes in responsiveness to antiviral therapy but not in pathogenicity or clinical progression (except for genotype 3, in which hepatic steatosis and clinical progression are more likely).

Currently available, third-generation immunoassays, which incorporate proteins from the core, NS3, and NS5 regions, detect anti-HCV antibodies during acute infection. The most sensitive indicator of HCV infection is the presence of HCV RNA, which requires molecular amplification by PCR or transcription-mediated amplification (TMA) (Fig. 360-7). To allow standardization of the quantification of HCV RNA among laboratories and commercial assays, HCV RNA is reported as international units (IUs) per milliliter; quantitative assays with a broad dynamic range are available that allow detection of HCV RNA with a sensitivity as low as 5 IU/mL. HCV RNA can be detected within a few days of exposure to HCV—well before the appearance of anti-HCV—and tends to persist for the duration of HCV infection. Application of sensitive molecular probes for HCV RNA has revealed the presence of replicative HCV in peripheral blood lymphocytes of infected persons; however, as is the case for HBV in lymphocytes, the clinical relevance of HCV lymphocyte infection is not known.

Hepatitis E Previously labeled *epidemic* or *enterically transmitted non-A, non-B hepatitis*, HEV is an enterically transmitted virus that causes clinically apparent hepatitis primarily in India, Asia, Africa, and Central America; in those geographic areas, HEV is the most common cause of acute hepatitis; one-third of the global population appears to have been infected. This agent, with epidemiologic features resembling those of hepatitis A, is a 27- to 34-nm, nonenveloped, HAV-like virus with a 7200-nucleotide, single-strand, positive-sense RNA genome. HEV has three open reading frames (ORF) (genes), the largest of

which, *ORF1*, encodes nonstructural proteins involved in virus replication. A middle-sized gene, *ORF2*, encodes the nucleocapsid protein, the major nonstructural protein, and the smallest, *ORF3*, encodes a structural protein whose function remains undetermined. All HEV isolates appear to belong to a single serotype, despite genomic heterogeneity of up to 25% and the existence of five genotypes, only four of which have been detected in humans; genotypes 1 and 2 appear to be more virulent, whereas genotypes 3 and 4 are more attenuated and account for subclinical infections. Contributing to the perpetuation of this virus are animal reservoirs, most notably in swine. No genomic or antigenic homology, however, exists between HEV and HAV or other picornaviruses; and HEV, although resembling caliciviruses, is sufficiently distinct from any known agent to merit its own classification as a unique genus, *Hepevirus*, within the family Hepeviridae. The virus has been detected in stool, bile, and liver and is excreted in the stool during the late incubation period. Both IgM anti-HEV during early acute infection and IgG anti-HEV predominating after the first 3 months can be detected. Currently, availability and reliability of serologic/virologic testing for HEV infection is limited but can be done in specialized laboratories (e.g., the Centers for Disease Control and Prevention).

PATHOGENESIS

Under ordinary circumstances, none of the hepatitis viruses is known to be directly cytopathic to hepatocytes. Evidence suggests that the clinical manifestations and outcomes after acute liver injury associated with viral hepatitis are determined by the immunologic responses of the host. Among the viral hepatitis, the immunopathogenesis of hepatitis B and C has been studied most extensively.

Hepatitis B For HBV, the existence of inactive hepatitis B carriers with normal liver histology and function suggests that the virus is not directly cytopathic. The fact that patients with defects in cellular immune competence are more likely to remain chronically infected rather than to clear HBV supports the role of cellular immune responses in the pathogenesis of hepatitis B–related liver injury. The model that has the most experimental support involves cytolytic T cells sensitized specifically to recognize host and hepatitis B viral antigens on the liver cell surface. Nucleocapsid proteins (HBcAg and possibly HBsAg), present on the cell membrane in minute quantities, are the viral target antigens that, with host antigens, invite cytolytic T cells to destroy HBV-infected hepatocytes. Differences in the robustness and broad polyclonality of CD8+ cytolytic T cell responsiveness; in the level of HBV-specific helper CD4+ T cells; in attenuation, depletion, and exhaustion of virus-specific T cells; in viral T cell epitope escape mutations that allow the virus to evade T cell containment; and in the elaboration of antiviral cytokines by T cells have been invoked to explain differences in outcomes between those who recover after acute hepatitis and those who progress to chronic hepatitis, or between those with mild and those with severe (fulminant) acute HBV infection.

Although a robust cytolytic T cell response occurs and eliminates virus-infected liver cells during acute hepatitis B, >90% of HBV DNA has been found in experimentally infected chimpanzees to disappear from the liver and blood before maximal T cell infiltration of the liver and before most of the biochemical and histologic evidence of liver injury. This observation suggests that components of the innate immune system and inflammatory cytokines, independent of cytopathic antiviral mechanisms, participate in the early immune response to HBV infection; this effect has been shown to represent elimination of HBV replicative intermediates from the cytoplasm and covalently closed circular viral DNA from the nucleus of infected hepatocytes. In turn, the innate immune response to HBV infection is mediated largely by natural killer (NK) cell cytotoxicity, activated by immunosuppressive cytokines (e.g., interleukin [IL] 10 and transforming growth factor [TGF] β), reduced signals from inhibitory receptor expression (e.g., major histocompatibility complex), or increased signals from activating receptor expression on infected hepatocytes. In addition, NK cells reduce helper CD4+ cells, which results in reduced CD8+ cells and exhaustion of the virus-specific T cell response to HBV infection. Ultimately, HBV-HLA-specific cytolytic T cell responses of the adaptive immune system are felt to be responsible for recovery from HBV infection.

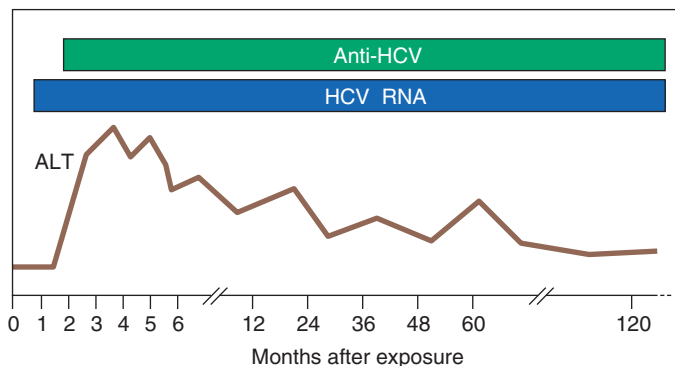


FIGURE 360-7 Scheme of typical laboratory features during acute hepatitis C progressing to chronicity. Hepatitis C virus (HCV) RNA is the first detectable event, preceding alanine aminotransferase (ALT) elevation and the appearance of anti-HCV.

Debate continues over the relative importance of viral and host factors in the pathogenesis of HBV-associated liver injury and its outcome. As noted above, precore genetic mutants of HBV have been associated with the more severe outcomes of HBV infection (severe chronic and fulminant hepatitis), suggesting that, under certain circumstances, relative pathogenicity is a property of the virus, not the host. The fact that concomitant HDV and HBV infections are associated with more severe liver injury than HBV infection alone and the fact that cells transfected *in vitro* with the gene for HDV antigen express HDV antigen and then become necrotic in the absence of any immunologic influences are also consistent with a viral effect on pathogenicity. Similarly, in patients who undergo liver transplantation for end-stage chronic hepatitis B, occasionally, rapidly progressive liver injury appears in the new liver. This clinical pattern is associated with an unusual histologic pattern in the new liver, *fibrosing cholestatic hepatitis*, which, ultrastructurally, appears to represent a choking of the cell with overwhelming quantities of HBsAg. This observation suggests that, under the influence of the potent immunosuppressive agents required to prevent allograft rejection, HBV may have a direct cytopathic effect on liver cells, independent of the immune system.

Although the precise mechanism of liver injury in HBV infection remains elusive, studies of nucleocapsid proteins have shed light on the profound immunologic tolerance to HBV of babies born to mothers with highly replicative (HBeAg-positive), chronic HBV infection. In HBeAg-expressing transgenic mice, *in utero* exposure to HBeAg, which is sufficiently small to traverse the placenta, induces T cell tolerance to both nucleocapsid proteins. This, in turn, may explain why, when infection occurs so early in life, immunologic clearance does not occur, and protracted, lifelong infection ensues.

An important distinction should be drawn between HBV infection acquired at birth, common in endemic areas, such as East Asia, and infection acquired in adulthood, common in the West. Infection in the neonatal period is associated with the acquisition of high-level immunologic tolerance to HBV and absence of an acute hepatitis illness, but the almost invariable establishment of chronic, often lifelong infection. Neonatally acquired HBV infection can culminate decades later in cirrhosis and hepatocellular carcinoma (see “Complications and Sequelae”). In contrast, when HBV infection is acquired during adolescence or early adulthood, the host immune response to HBV-infected hepatocytes tends to be robust, an acute hepatitis-like illness is the rule, and failure to recover is the exception. After adulthood-acquired infection, chronicity is uncommon, and the risk of hepatocellular carcinoma is very low. Based on these observations, some authorities categorize HBV infection into an “immunotolerant” phase, an “immunoreactive” phase, and an “inactive” phase. This somewhat simplistic formulation does not apply at all to the typical adult in the West with self-limited acute hepatitis B, in whom no period of immunologic tolerance occurs. Even among those with neonatally acquired HBV infection, in whom immunologic tolerance is established definitively, intermittent bursts of hepatic necroinflammatory activity punctuate the early decades of life during which liver injury appears to be quiescent (labeled by some as the “immunotolerant” phase). In addition, even when clinically apparent liver injury and progressive fibrosis emerge during later decades (the so-called immunoreactive, or immunointolerant, phase), the level of immunologic tolerance to HBV remains substantial. More accurately, in patients with neonatally acquired HBV infection, a dynamic equilibrium exists between tolerance and intolerance, the outcome of which determines the clinical expression of chronic infection. Persons infected as neonates tend to have a relatively higher level of immunologic tolerance during the early decades of life and a relatively lower level (but only rarely a loss) of tolerance in the later decades of life.

Hepatitis C Cell-mediated immune responses and elaboration by T cells of antiviral cytokines contribute to the multicellular innate and adaptive immune responses involved in the containment of infection and pathogenesis of liver injury associated with hepatitis C. The fact that HCV is so efficient in evading these immune mechanisms is a testament to its highly evolved ability to disrupt host immune responses

at multiple levels. After exposure to HCV, the host cell identifies viral product motifs (pattern recognition receptors) that distinguish the virus from “self,” resulting in the elaboration of interferons and other cytokines that result in activation of innate and adaptive immune responses. Intrahepatic HLA class I restricted cytolytic T cells directed at nucleocapsid, envelope, and nonstructural viral protein antigens have been demonstrated in patients with chronic hepatitis C; however, such virus-specific cytolytic T cell responses do not correlate adequately with the degree of liver injury or with recovery. Yet, a consensus has emerged supporting a role in the pathogenesis of HCV-associated liver injury of virus-activated CD4+ helper T cells that stimulate, via the cytokines they elaborate, HCV-specific CD8+ cytotoxic T cells. These responses appear to be more robust (higher in number, more diverse in viral antigen specificity, more functionally effective, and more long lasting) in those who recover from HCV than in those who have chronic infection. Contributing to chronic infection are a CD4+ proliferative defect that results in rapid contraction of CD4+ responses, mutations in CD8+ T cell–targeted viral epitopes that allow HCV to escape immune-mediated clearance, and upregulation of inhibitory receptors on functionally impaired, exhausted T cells. Although attention has focused on adaptive immunity, HCV proteins have been shown to interfere with innate immunity by resulting in blocking of type I interferon responses and inhibition of interferon signaling and effector molecules in the interferon signaling cascade. Several HLA alleles have been linked with self-limited hepatitis C, the most convincing of which is the CC haplotype of the *IL28B* gene, which codes for interferon λ 3, a component of innate immune antiviral defense. The *IL28B* association is even stronger when combined with HLA class II *DQB1*03:01*. The link between non-CC *IL28B* polymorphisms and failure to clear HCV infection has been explained by a chromosome 19q13.13 frameshift variant upstream of *IL28B*, the Δ G polymorphism of which creates an ORF in a novel interferon gene (*IFN- λ 4*) associated with impaired HCV clearance. Also shown to contribute to limiting HCV infection are NK cells of the innate immune system that function when HLA class I molecules required for successful adaptive immunity are underexpressed. Both peripheral and intrahepatic NK cell cytotoxicity are dysfunctional in persistent HCV infection. Adding to the complexity of the immune response, HCV core, NS4B, and NS5B have been shown to suppress the immunoregulatory nuclear factor (NF)- κ B pathway, resulting in reduced antiapoptotic proteins and a resultant increased vulnerability to tumor necrosis factor (TNF) α -mediated cell death. Patients with hepatitis C and unfavorable (non-CC, associated with reduced HCV clearance) *IL28B* alleles have been shown to have depressed NK cell/innate immune function. Of note, the emergence of substantial viral quasispecies diversity and HCV sequence variation allow the virus to evade attempts by the host to contain HCV infection by both humoral and cellular immunity.

Finally, cross-reactivity between viral antigens (HCV NS3 and NS5A) and host autoantigens (cytochrome P450 2D6) has been invoked to explain the association between hepatitis C and a subset of patients with autoimmune hepatitis and antibodies to liver-kidney microsomal (LKM) antigen (anti-LKM) ([Chap. 362](#)).

EXTRAHEPATIC MANIFESTATIONS

Immune complex–mediated tissue damage appears to play a pathogenetic role in the extrahepatic manifestations of acute hepatitis B. The occasional prodromal serum sickness–like syndrome observed in acute hepatitis B appears to be related to the deposition in tissue blood vessel walls of HBsAg-anti-HBs circulating immune complexes, leading to activation of the complement system and depressed serum complement levels.

In patients with chronic hepatitis B, other types of immune-complex disease may be seen. Glomerulonephritis with the nephrotic syndrome is observed occasionally; HBsAg, immunoglobulin, and C3 deposition has been found in the glomerular basement membrane. Whereas generalized vasculitis (polyarteritis nodosa) develops in considerably fewer than 1% of patients with chronic HBV infection, 20–30% of patients with polyarteritis nodosa have HBsAg in serum ([Chap. 385](#)). In these patients, the affected small- and medium-size arterioles contain

2012 HBsAg, immunoglobulins, and complement components. Another extrahepatic manifestation of viral hepatitis, essential mixed cryoglobulinemia (EMC), was reported initially to be associated with hepatitis B. The disorder is characterized clinically by arthritis, cutaneous vasculitis (palpable purpura), and occasionally, glomerulonephritis and serologically by the presence of circulating cryoprecipitable immune complexes of more than one immunoglobulin class (Chaps. 338 and 385). Many patients with this syndrome have chronic liver disease, but the association with HBV infection is limited; instead, a substantial proportion has chronic HCV infection, with circulating immune complexes containing HCV RNA. Immune-complex glomerulonephritis is another recognized extrahepatic manifestation of chronic hepatitis C.

PATHOLOGY

The typical morphologic lesions of all types of viral hepatitis are similar and consist of panlobular infiltration with mononuclear cells, hepatic cell necrosis, hyperplasia of Kupffer cells, and variable degrees of cholestasis. Hepatic cell regeneration is present, as evidenced by numerous mitotic figures, multinucleated cells, and “rosette” or “pseudoacinar” formation. The mononuclear infiltration consists primarily of small lymphocytes, although plasma cells and eosinophils occasionally are present. Liver cell damage consists of hepatic cell degeneration and necrosis, cell dropout, ballooning of cells, and acidophilic degeneration of hepatocytes (forming so-called Councilman or apoptotic bodies). Large hepatocytes with a ground-glass appearance of the cytoplasm may be seen in chronic but not in acute HBV infection; these cells contain HBsAg and can be identified histochemically with orcein or aldehyde fuchsin. In uncomplicated viral hepatitis, the reticulin framework is preserved.

In hepatitis C, the histologic lesion is often remarkable for a relative paucity of inflammation, a marked increase in activation of sinusoidal lining cells, lymphoid aggregates, the presence of fat (more frequent in genotype 3 and linked to increased fibrosis), and, occasionally, bile duct lesions in which biliary epithelial cells appear to be piled up without interruption of the basement membrane. Occasionally, microvesicular steatosis occurs in hepatitis D. In hepatitis E, a common histologic feature is marked cholestasis. A cholestatic variant of slowly resolving acute hepatitis A also has been described.

A more severe histologic lesion, *bridging hepatic necrosis*, also termed *subacute* or *confluent necrosis* or *interface hepatitis*, is observed occasionally in acute hepatitis. “Bridging” between lobules results from large areas of hepatic cell dropout, with collapse of the reticulin framework. Characteristically, the bridge consists of condensed reticulum, inflammatory debris, and degenerating liver cells that span adjacent portal areas, portal to central veins, or central vein to central vein. This lesion had been thought to have prognostic significance; in many of the originally described patients with this lesion, a subacute course terminated in death within several weeks to months, or severe chronic hepatitis and cirrhosis developed; however, the association between bridging necrosis and a poor prognosis in patients with acute hepatitis has not been upheld. Therefore, although demonstration of this lesion in patients with chronic hepatitis has prognostic significance (Chap. 362), its demonstration during acute hepatitis is less meaningful, and liver biopsies to identify this lesion are no longer undertaken routinely in patients with acute hepatitis. In *massive hepatic necrosis* (fulminant hepatitis, “acute yellow atrophy”), the striking feature at postmortem examination is the finding of a small, shrunken, soft liver. Histologic examination reveals massive necrosis and dropout of liver cells of most lobules with extensive collapse and condensation of the reticulin framework. When histologic documentation is required in the management of fulminant or very severe hepatitis, a biopsy can be done by the angiographically guided transjugular route, which permits the performance of this invasive procedure in the presence of severe coagulopathy.

Immunohistochemical and electron-microscopic studies have localized HBsAg to the cytoplasm and plasma membrane of infected liver cells. In contrast, HBeAg predominates in the nucleus, but, occasionally, scant amounts are also seen in the cytoplasm and on the cell membrane. HDV antigen is localized to the hepatocyte nucleus, whereas HAV, HCV, and HEV antigens are localized to the cytoplasm.

EPIDEMIOLOGY AND GLOBAL FEATURES



Before the availability of serologic tests for hepatitis viruses, all viral hepatitis cases were labeled either as “infectious” or “serum” hepatitis. Modes of transmission overlap, however, and a clear distinction among the different types of viral hepatitis cannot be made solely on the basis of clinical or epidemiologic features (Table 360-2). The most accurate means to distinguish the various types of viral hepatitis involves specific serologic testing.

Hepatitis A This agent is transmitted almost exclusively by the fecal-oral route. Person-to-person spread of HAV is enhanced by poor personal hygiene and overcrowding; large outbreaks as well as sporadic cases have been traced to contaminated food, water, milk, frozen raspberries and strawberries, green onions imported from Mexico, and shellfish. Intrafamily and intrainstitutional spread are also common. Early epidemiologic observations supported a predilection for hepatitis A to occur in late fall and early winter. In temperate zones, epidemic waves have been recorded every 5–20 years as new segments of nonimmune population appeared; however, in developed countries, the incidence of hepatitis A has been declining, presumably as a function of improved sanitation, and these cyclic patterns are no longer observed. No HAV carrier state has been identified after acute hepatitis A; perpetuation of the virus in nature depends presumably on nonepidemic, inapparent subclinical infection, ingestion of contaminated food or water in, or imported from, endemic areas, and/or contamination linked to environmental reservoirs.

In the general population, anti-HAV, a marker for previous HAV infection, increases in prevalence as a function of increasing age and of decreasing socioeconomic status. In the 1970s, serologic evidence of prior hepatitis A infection occurred in ~40% of urban populations in the United States, most of whose members never recalled having had a symptomatic case of hepatitis. In subsequent decades, however, the prevalence of anti-HAV has been declining in the United States. In developing countries, exposure, infection, and subsequent immunity are almost universal in childhood. As the frequency of subclinical childhood infections declines in developed countries, a susceptible cohort of adults emerges. Hepatitis A tends to be more symptomatic in adults; therefore, paradoxically, as the frequency of HAV infection declines, the likelihood of clinically apparent, even severe, HAV illnesses increases in the susceptible adult population. Travel to endemic areas is a common source of infection for adults from nonendemic areas. More recently recognized epidemiologic foci of HAV infection include child care centers, neonatal intensive care units, promiscuous men who have sex with men, injection drug users, and unvaccinated close contacts of newly arrived international adopted children, most of whom emanate from countries with intermediate-to-high hepatitis A endemicity. Although hepatitis A is rarely bloodborne, several outbreaks have been recognized in recipients of clotting-factor concentrates. In the United States, the introduction of hepatitis A vaccination programs among children from high-incidence states has resulted in a >70% reduction in the annual incidence of new HAV infections and has shifted the burden of new infections from children to young adults. In the most recent, 1999–2006 U.S. Public Health Service National Health and Nutrition Examination Survey (NHANES), the prevalence of anti-HAV in the U.S. population was 35%, representing (compared to the 1988–1994 survey) a stable frequency of infection and natural immunity in adults >19 years old but an increase in vaccine-induced immunity for children age 6–19 years.

Hepatitis B Percutaneous inoculation has long been recognized as a major route of hepatitis B transmission, but the outmoded designation “serum hepatitis” is an inaccurate label for the epidemiologic spectrum of HBV infection. As detailed below, most of the hepatitis transmitted by blood transfusion is not caused by HBV; moreover, in approximately two-thirds of patients with acute type B hepatitis, no history of an identifiable percutaneous exposure can be elicited. We now recognize that many cases of hepatitis B result from less obvious modes of nonpercutaneous or covert percutaneous transmission. HBsAg has been identified in almost every body fluid from

TABLE 360-2 CLINICAL AND EPIDEMIOLOGIC FEATURES OF VIRAL HEPATITIS

Feature	HAV	HBV	HCV	HDV	HEV
Incubation (days)	15–45, mean 30	30–180, mean 60–90	15–160, mean 50	30–180, mean 60–90	14–60, mean 40
Onset	Acute	Insidious or acute	Insidious	Insidious or acute	Acute
Age preference	Children, young adults	Young adults (sexual and percutaneous), babies, toddlers	Any age, but more common in adults	Any age (similar to HBV)	Epidemic cases: young adults (20–40 years); sporadic cases: older adults (>60)
Transmission					
Fecal-oral	+++	–	–	–	+++
Percutaneous	Unusual	+++	+++	+++	–
Perinatal	–	+++	± ^a	+	–
Sexual	±	++	± ^a	++	–
Clinical					
Severity	Mild	Occasionally severe	Moderate	Occasionally severe	Mild
Fulminant	0.1%	0.1–1%	0.1%	5–20% ^b	1–2% ^e
Progression to chronicity	None	Occasional (1–10%) (90% of neonates)	Common (85%)	Common ^d	None ^f
Carrier	None	0.1–30% ^c	1.5–3.2%	Variable ^g	None
Cancer	None	+ (neonatal infection)	+	±	None
Prognosis	Excellent	Worse with age, debility	Moderate	Acute, good Chronic, poor	Good
Prophylaxis					
	Ig, inactivated vaccine	HBIG, recombinant vaccine	None	HBV vaccine (none for HBV carriers)	Vaccine
Therapy					
	None	Interferon Lamivudine Adefovir Pegylated interferon Entecavir Telbivudine Tenofovir	Pegylated interferon plus ribavirin, telaprevir, boceprevir	Pegylated interferon ±	None ^h

^aPrimarily with HIV co-infection and high-level viremia in index case; risk ~5%. ^bUp to 5% in acute HBV/HDV co-infection; up to 20% in HDV superinfection of chronic HBV infection. ^cVaries considerably throughout the world and in subpopulations within countries; see text. ^dIn acute HBV/HDV co-infection, the frequency of chronicity is the same as that for HBV; in HDV superinfection, chronicity is invariable. ^e10–20% in pregnant women. ^fExcept as observed in immunosuppressed liver allograft recipients or other immunosuppressed hosts. ^gCommon in Mediterranean countries; rare in North America and western Europe. ^hAnecdotal reports and retrospective studies suggest that pegylated interferon and/or ribavirin are effective in treating chronic hepatitis E, observed in immunocompromised persons.

Abbreviation: HBIG, hepatitis B immunoglobulin. See text for other abbreviations.

infected persons, and at least some of these body fluids—most notably semen and saliva—are infectious, albeit less so than serum, when administered percutaneously or nonpercutaneously to experimental animals. Among the nonpercutaneous modes of HBV transmission, oral ingestion has been documented as a potential but inefficient route of exposure. By contrast, the two nonpercutaneous routes considered to have the greatest impact are intimate (especially sexual) contact and perinatal transmission.

In sub-Saharan Africa, intimate contact among toddlers is considered instrumental in contributing to the maintenance of the high frequency of hepatitis B in the population. Perinatal transmission occurs primarily in infants born to mothers with chronic hepatitis B or (rarely) mothers with acute hepatitis B during the third trimester of pregnancy or during the early postpartum period. Perinatal transmission is uncommon in North America and western Europe but occurs with great frequency and is the most important mode of HBV perpetuation in East Asia and developing countries. Although the precise mode of perinatal transmission is unknown, and although ~10% of infections may be acquired in utero, epidemiologic evidence suggests that most infections occur approximately at the time of delivery and are not related to breast-feeding. The likelihood of perinatal transmission of HBV correlates with the presence of HBeAg and high-level viral replication; 90% of HBeAg-positive mothers but only 10–15% of anti-HBe-positive mothers transmit HBV infection to their offspring. In most cases, acute infection in the neonate is clinically asymptomatic, but the child is very likely to remain chronically infected.

The >350–400 million HBsAg carriers in the world constitute the main reservoir of hepatitis B in human beings. Whereas serum HBsAg is infrequent (0.1–0.5%) in normal populations in the United States and western Europe, a prevalence of up to 5–20% has been found in East Asia and in some tropical countries; in persons with Down's

syndrome, lepromatous leprosy, leukemia, Hodgkin's disease, or polyarteritis nodosa; in patients with chronic renal disease on hemodialysis; and in injection drug users.

Other groups with high rates of HBV infection include spouses of acutely infected persons; sexually promiscuous persons (especially promiscuous men who have sex with men); health care workers exposed to blood; persons who require repeated transfusions especially with pooled blood-product concentrates (e.g., hemophiliacs); residents and staff of custodial institutions for the developmentally handicapped; prisoners; and, to a lesser extent, family members of chronically infected patients. In volunteer blood donors, the prevalence of anti-HBs, a reflection of previous HBV infection, ranges from 5–10%, but the prevalence is higher in lower socioeconomic strata, older age groups, and persons—including those mentioned above—exposed to blood products. Because of highly sensitive virologic screening of donor blood, the risk of acquiring HBV infection from a blood transfusion is 1 in 230,000.

Prevalence of infection, modes of transmission, and human behavior conspire to mold geographically different epidemiologic patterns of HBV infection. In East Asia and Africa, hepatitis B, a disease of the newborn and young children, is perpetuated by a cycle of maternal-neonatal spread. In North America and western Europe, hepatitis B is primarily a disease of adolescence and early adulthood, the time of life when intimate sexual contact and recreational and occupational percutaneous exposures tend to occur. To some degree, however, this dichotomy between high-prevalence and low-prevalence geographic regions has been minimized by immigration from high-prevalence to low-prevalence areas. The introduction of hepatitis B vaccine in the early 1980s and adoption of universal childhood vaccination policies in many countries resulted in a dramatic, ~90% decline in the incidence of new HBV infections in those countries as well as in the dire

TABLE 360-3 HIGH-RISK POPULATIONS FOR WHOM HBV INFECTION SCREENING IS RECOMMENDED

Persons born in countries/regions with a high ($\geq 8\%$) and intermediate ($\geq 2\%$) prevalence of HBV infection including immigrants and adopted children and including persons born in the United States who were not vaccinated as infants and whose parents emigrated from areas of high HBV endemicity

Household and sexual contacts of persons with hepatitis B

Babies born to HBsAg-positive mothers

Persons who have used injection drugs

Persons with multiple sexual contacts or a history of sexually transmitted disease

Men who have sex with men

Inmates of correctional facilities

Persons with elevated alanine or aspartate aminotransferase levels

Blood/plasma/organ/tissue/semen donors

Persons with HCV or HIV infection

Hemodialysis patients

Pregnant women

Persons who are the source of blood or body fluids that would be an indication for postexposure prophylaxis (e.g., needlestick, mucosal exposure, sexual assault)

Persons who require immunosuppressive or cytotoxic therapy (including anti-tumor necrosis factor α therapy for rheumatologic or inflammatory bowel disorders)

consequences of chronic infection, including hepatocellular carcinoma. Populations and groups for whom HBV infection screening is recommended are listed in [Table 360-3](#).

Hepatitis D Infection with HDV has a worldwide distribution, but two epidemiologic patterns exist. In Mediterranean countries (northern Africa, southern Europe, the Middle East), HDV infection is endemic among those with hepatitis B, and the disease is transmitted predominantly by nonpercutaneous means, especially close personal contact. In nonendemic areas, such as the United States and northern Europe, HDV infection is confined to persons exposed frequently to blood and blood products, primarily injection drug users and hemophiliacs. HDV infection can be introduced into a population through drug users or by migration of persons from endemic to nonendemic areas. Thus, patterns of population migration and human behavior facilitating percutaneous contact play important roles in the introduction and amplification of HDV infection. Occasionally, the migrating epidemiology of hepatitis D is expressed in explosive outbreaks of severe hepatitis, such as those that have occurred in remote South American villages as well as in urban centers in the United States. Ultimately, such outbreaks of hepatitis D—either of co-infections with acute hepatitis B or of superinfections in those already infected with HBV—may blur the distinctions between endemic and nonendemic areas. On a global scale, HDV infection declined at the end of the 1990s. Even in Italy, an HDV-endemic area, public health measures introduced to control HBV infection resulted during the 1990s in a 1.5%/year reduction in the prevalence of HDV infection. Still, the frequency of HDV infection during the first decade of the twenty-first century has not fallen below levels reached during the 1990s; the reservoir has been sustained by survivors infected during 1970–1980 and recent immigrants from still-endemic to less-endemic countries.

Hepatitis C Routine screening of blood donors for HBsAg and the elimination of commercial blood sources in the early 1970s reduced the frequency of, but did not eliminate, transfusion-associated hepatitis. During the 1970s, the likelihood of acquiring hepatitis after transfusion of voluntarily donated, HBsAg-screened blood was $\sim 10\%$ per patient (up to 0.9% per unit transfused); 90–95% of these cases were classified, based on serologic exclusion of hepatitis A and B, as “non-A, non-B” hepatitis. For patients requiring transfusion of pooled products, such as clotting factor concentrates, the risk was even higher, up to 20–30%.

During the 1980s, voluntary self-exclusion of blood donors with risk factors for AIDS and then the introduction of donor screening

for anti-HIV reduced further the likelihood of transfusion-associated hepatitis to $< 5\%$. During the late 1980s and early 1990s, the introduction first of “surrogate” screening tests for non-A, non-B hepatitis (alanine aminotransferase [ALT] and anti-HBc, both shown to identify blood donors with a higher likelihood of transmitting non-A, non-B hepatitis to recipients) and, subsequently, after the discovery of HCV, first-generation immunoassays for anti-HCV reduced the frequency of transfusion-associated hepatitis even further. A prospective analysis of transfusion-associated hepatitis conducted between 1986 and 1990 showed that the frequency of transfusion-associated hepatitis at one urban university hospital fell from a baseline of 3.8% per patient (0.45% per unit transfused) to 1.5% per patient (0.19% per unit) after the introduction of surrogate testing and to 0.6% per patient (0.03% per unit) after the introduction of first-generation anti-HCV assays. The introduction of second-generation anti-HCV assays reduced the frequency of transfusion-associated hepatitis C to almost imperceptible levels—1 in 100,000—and these gains were reinforced by the application of third-generation anti-HCV assays and of automated PCR testing of donated blood for HCV RNA, which has resulted in a reduction in the risk of transfusion-associated HCV infection to 1 in 2.3 million transfusions.

In addition to being transmitted by transfusion, hepatitis C can be transmitted by other percutaneous routes, such as injection drug use. In addition, this virus can be transmitted by occupational exposure to blood, and the likelihood of infection is increased in hemodialysis units. Although the frequency of transfusion-associated hepatitis C fell as a result of blood-donor screening, the overall frequency of hepatitis C remained the same until the early 1990s, when the overall frequency fell by 80%, in parallel with a reduction in the number of new cases in injection drug users. After the exclusion of anti-HCV-positive plasma units from the donor pool, rare, sporadic instances have occurred of hepatitis C among recipients of immunoglobulin (Ig) preparations for intravenous (but not intramuscular) use.

Serologic evidence for HCV infection occurs in 90% of patients with a history of transfusion-associated hepatitis (almost all occurring before 1992, when second-generation HCV screening tests were introduced); hemophiliacs and others treated with clotting factors; injection drug users; 60–70% of patients with sporadic “non-A, non-B” hepatitis who lack identifiable risk factors; 0.5% of volunteer blood donors; and, in a survey conducted in the United States between 1999 and 2002, 1.6% of the general population in the United States, which translates into 4.1 million persons (3.2 million with viremia), the majority of whom are unaware of their infections. Moreover, such population surveys do not include higher-risk groups such as incarcerated prisoners and active injection drug users, indicating that the actual prevalence is even higher. Comparable frequencies of HCV infection occur in most countries around the world, with 170 million persons infected worldwide, but extraordinarily high prevalences of HCV infection occur in certain countries such as Egypt, where $> 20\%$ of the population (as high as 50% in persons born prior to 1960) in some cities is infected. The high frequency in Egypt is attributable to contaminated equipment used for medical procedures and unsafe injection practices in the 1950s to 1980s (during a campaign to eradicate schistosomiasis with intravenous tartar emetic). In the United States, African Americans and Mexican Americans have higher frequencies of HCV infection than whites. Between 1988 and 1994, 30- to 40-year-old adult males had the highest prevalence of HCV infection; however, in a survey conducted between 1999 and 2002, the peak age decile had shifted to those age 40–49 years; an increase in hepatitis C–related mortality has paralleled this secular trend, increasing since 1995 predominantly in the 45- to 65-year age group. Thus, despite an 80% reduction in new HCV infections during the 1990s, the prevalence of HCV infection in the population was sustained by an aging cohort that had acquired their infections three to four decades earlier, during the 1960s and 1970s, as a result predominantly of self-inoculation with recreational drugs. As death resulting from HIV infection fell after 1999, age-adjusted mortality associated with HCV infection surpassed that of HIV infection in 2007; $> 70\%$ of HCV-associated deaths occurred in the “baby boomer” cohort born between 1945 and 1965. Compared to the 1.6%

prevalence of HCV infection in the population at large, the prevalence in the 1945–1965 birth cohort was 3.2%, representing three-quarters of all infected persons. Therefore, in 2012, the Centers for Disease Control and Prevention recommended that all persons born between 1945 and 1965 be screened for hepatitis C, without ascertainment of risk, a recommendation shown to be cost-effective and predicted to identify 800,000 infected persons. Because of the availability of highly effective antiviral therapy, such screening would have the potential to avert 200,000 cases of cirrhosis and 47,000 cases of hepatocellular carcinoma and to prevent 120,000 hepatitis-related deaths.

Hepatitis C accounts for 40% of chronic liver disease, is the most frequent indication for liver transplantation, and is estimated to account for 8000–10,000 deaths per year in the United States. The distribution of HCV genotypes varies in different parts of the world. Worldwide, genotype 1 is the most common. In the United States, genotype 1 accounts for 70% of HCV infections, whereas genotypes 2 and 3 account for the remaining 30%; among African Americans, the frequency of genotype 1 is even higher (i.e., 90%). Genotype 4 predominates in Egypt; genotype 5 is localized to South Africa, genotype 6 to Hong Kong, and genotype 7 to Central Africa. Most asymptomatic blood donors found to have anti-HCV and ~20–30% of persons with reported cases of acute hepatitis C do not fall into a recognized risk group; however, many such blood donors do recall risk-associated behaviors when questioned carefully.

As a bloodborne infection, HCV potentially can be transmitted sexually and perinatally; however, both of these modes of transmission are inefficient for hepatitis C. Although 10–15% of patients with acute hepatitis C report having potential sexual sources of infection, most studies have failed to identify sexual transmission of this agent. The chances of sexual and perinatal transmission have been estimated to be ~5% but shown in a prospective study to be only 1% between monogamous sexual partners, well below comparable rates for HIV and HBV infections. Moreover, sexual transmission appears to be confined to such subgroups as persons with multiple sexual partners and sexually transmitted diseases. Breast-feeding does not increase the risk of HCV infection between an infected mother and her infant. Infection of health workers is not dramatically higher than among the general population; however, health workers are more likely to acquire HCV infection through accidental needle punctures, the efficiency of which is ~3%. Infection of household contacts is rare as well.

Besides persons born between 1945 and 1965, other groups with an increased frequency of HCV infection are listed in [Table 360-4](#). In immunosuppressed individuals, levels of anti-HCV may be undetectable, and a diagnosis may require testing for HCV RNA. Although new acute cases of hepatitis C are rare, newly diagnosed cases are common among otherwise healthy persons who experimented briefly with injection drugs, as noted above, three or four decades earlier. Such instances usually remain unrecognized for years, until unearthed by laboratory screening for routine medical examinations, insurance applications, and attempted blood donation. Although, overall, the annual incidence of new HCV infections has continued to fall, the rate of new infections has been increasing since 2002 in a new cohort

of young injection drug users, age 15–24 years (accounting for more than two-thirds of all acute cases), who, unlike older cohorts, had not learned to take precautions to prevent bloodborne infections.

Hepatitis E This type of hepatitis, identified in India, Asia, Africa, the Middle East, and Central America, resembles hepatitis A in its primarily enteric mode of spread. The commonly recognized cases occur after contamination of water supplies such as after monsoon flooding, but sporadic, isolated cases occur. An epidemiologic feature that distinguishes HEV from other enteric agents is the rarity of secondary person-to-person spread from infected persons to their close contacts. Large waterborne outbreaks in endemic areas are linked to genotypes 1 and 2, arise in populations that are immune to HAV, favor young adults, and account for antibody prevalences of 30–80%. In nonendemic areas of the world, such as the United States, clinically apparent acute hepatitis E is extremely rare; however, during the 1988–1994 NHANES survey conducted by the U.S. Public Health Service, the prevalence of anti-HEV was 21%, reflecting subclinical infections, infection with genotypes 3 and 4, predominantly in older males (>60 years). In nonendemic areas, HEV accounts hardly at all for cases of sporadic hepatitis; however, cases imported from endemic areas have been found in the United States. Evidence supports a zoonotic reservoir for HEV primarily in swine, which may account for the mostly subclinical infections in nonendemic areas.

CLINICAL AND LABORATORY FEATURES

Symptoms and Signs Acute viral hepatitis occurs after an incubation period that varies according to the responsible agent. Generally, incubation periods for hepatitis A range from 15–45 days (mean, 4 weeks), for hepatitis B and D from 30–180 days (mean, 8–12 weeks), for hepatitis C from 15–160 days (mean, 7 weeks), and for hepatitis E from 14–60 days (mean, 5–6 weeks). The *prodromal symptoms* of acute viral hepatitis are systemic and quite variable. Constitutional symptoms of anorexia, nausea and vomiting, fatigue, malaise, arthralgias, myalgias, headache, photophobia, pharyngitis, cough, and coryza may precede the onset of jaundice by 1–2 weeks. The nausea, vomiting, and anorexia are frequently associated with alterations in olfaction and taste. A low-grade fever between 38° and 39°C (100°–102°F) is more often present in hepatitis A and E than in hepatitis B or C, except when hepatitis B is heralded by a serum sickness–like syndrome; rarely, a fever of 39.5°–40°C (103°–104°F) may accompany the constitutional symptoms. Dark urine and clay-colored stools may be noticed by the patient from 1–5 days before the onset of clinical jaundice.

With the onset of *clinical jaundice*, the constitutional prodromal symptoms usually diminish, but in some patients, mild weight loss (2.5–5 kg) is common and may continue during the entire icteric phase. The liver becomes enlarged and tender and may be associated with right upper quadrant pain and discomfort. Infrequently, patients present with a cholestatic picture, suggesting extrahepatic biliary obstruction. Splenomegaly and cervical adenopathy are present in 10–20% of patients with acute hepatitis. Rarely, a few spider angiomas appear during the icteric phase and disappear during convalescence. During the *recovery phase*, constitutional symptoms disappear, but usually some liver enlargement and abnormalities in liver biochemical tests are still evident. The duration of the posticteric phase is variable, ranging from 2–12 weeks, and is usually more prolonged in acute hepatitis B and C. Complete clinical and biochemical recovery is to be expected 1–2 months after all cases of hepatitis A and E and 3–4 months after the onset of jaundice in three-quarters of uncomplicated, self-limited cases of hepatitis B and C (among healthy adults, acute hepatitis B is self-limited in 95–99%, whereas hepatitis C is self-limited in only ~15%). In the remainder, biochemical recovery may be delayed. A substantial proportion of patients with viral hepatitis never become icteric.

Infection with HDV can occur in the presence of acute or chronic HBV infection; the duration of HBV infection determines the duration of HDV infection. When acute HDV and HBV infection occur simultaneously, clinical and biochemical features may be indistinguishable from those of HBV infection alone, although occasionally they are

TABLE 360-4 HIGH-RISK POPULATIONS FOR WHOM HCV-INFECTION SCREENING IS RECOMMENDED

Persons born between 1945 and 1965
Persons who have ever used injection drugs
Persons with HIV infection
Hemophiliacs treated with clotting factor concentrates prior to 1987
Persons who have ever undergone long-term hemodialysis
Persons with unexplained elevations of aminotransferase levels
Transfusion or transplantation recipients prior to July 1992
Recipients of blood or organs from a donor found to be positive for hepatitis C
Children born to women with hepatitis C
Health care, public safety, and emergency medical personnel following needle injury or mucosal exposure to HCV-contaminated blood
Sexual partners of persons with hepatitis C infection

more severe. As opposed to patients with *acute* HBV infection, patients with *chronic* HBV infection can support HDV replication indefinitely, as when acute HDV infection occurs in the presence of a nonresolving acute HBV infection or, more commonly, when acute hepatitis D is superimposed on underlying chronic hepatitis B. In such cases, the HDV superinfection appears as a clinical exacerbation or an episode resembling acute viral hepatitis in someone already chronically infected with HBV. Superinfection with HDV in a patient with chronic hepatitis B often leads to clinical deterioration (see below).

In addition to superinfections with other hepatitis agents, acute hepatitis-like clinical events in persons with chronic hepatitis B may accompany spontaneous HBeAg to anti-HBe seroconversion or spontaneous reactivation (i.e., reversion from relatively nonreplicative to replicative infection). Such reactivations can occur as well in therapeutically immunosuppressed patients with chronic HBV infection when cytotoxic/immunosuppressive drugs are withdrawn; in these cases, restoration of immune competence is thought to allow resumption of previously checked cell-mediated immune cytolysis of HBV-infected hepatocytes. Occasionally, acute clinical exacerbations of chronic hepatitis B may represent the emergence of a precore mutant (see “Virology and Etiology”), and the subsequent course in such patients may be characterized by periodic exacerbations. Cytotoxic chemotherapy can lead to reactivation of chronic hepatitis C as well, and anti-TNF- α therapy can lead to reactivation of both hepatitis B and C.

Laboratory Features The serum aminotransferases aspartate aminotransferase (AST) and alanine aminotransferase (ALT) (previously designated SGOT and SGPT) increase to a variable degree during the prodromal phase of acute viral hepatitis and precede the rise in bilirubin level (Figs. 360-2 and 360-4). The level of these enzymes, however, does not correlate well with the degree of liver cell damage. Peak levels vary from 400–4000 IU or more; these levels are usually reached at the time the patient is clinically icteric and diminish progressively during the recovery phase of acute hepatitis. The diagnosis of anicteric hepatitis is based on clinical features and on aminotransferase elevations.

Jaundice is usually visible in the sclera or skin when the serum bilirubin value is $>43 \mu\text{mol/L}$ (2.5 mg/dL). When jaundice appears, the serum bilirubin typically rises to levels ranging from 85–340 $\mu\text{mol/L}$ (5–20 mg/dL). The serum bilirubin may continue to rise despite falling serum aminotransferase levels. In most instances, the total bilirubin is equally divided between the conjugated and unconjugated fractions. Bilirubin levels $>340 \mu\text{mol/L}$ (20 mg/dL) extending and persisting late into the course of viral hepatitis are more likely to be associated with severe disease. In certain patients with underlying hemolytic anemia, however, such as glucose-6-phosphate dehydrogenase deficiency and sickle cell anemia, a high serum bilirubin level is common, resulting from superimposed hemolysis. In such patients, bilirubin levels $>513 \mu\text{mol/L}$ (30 mg/dL) have been observed and are not necessarily associated with a poor prognosis.

Neutropenia and lymphopenia are transient and are followed by a relative lymphocytosis. Atypical lymphocytes (varying between 2 and 20%) are common during the acute phase. Measurement of the prothrombin time (PT) is important in patients with acute viral hepatitis, because a prolonged value may reflect a severe hepatic synthetic defect, signify extensive hepatocellular necrosis, and indicate a worse prognosis. Occasionally, a prolonged PT may occur with only mild increases in the serum bilirubin and aminotransferase levels. Prolonged nausea and vomiting, inadequate carbohydrate intake, and poor hepatic glycogen reserves may contribute to hypoglycemia noted occasionally in patients with severe viral hepatitis. Serum alkaline phosphatase may be normal or only mildly elevated, whereas a fall in serum albumin is uncommon in uncomplicated acute viral hepatitis. In some patients, mild and transient steatorrhea has been noted, as well as slight microscopic hematuria and minimal proteinuria.

A diffuse but mild elevation of the γ globulin fraction is common during acute viral hepatitis. Serum IgG and IgM levels are elevated in about one-third of patients during the acute phase of viral hepatitis, but the serum IgM level is elevated more characteristically during acute hepatitis A. During the acute phase of viral hepatitis, antibodies

to smooth muscle and other cell constituents may be present, and low titers of rheumatoid factor, nuclear antibody, and heterophile antibody can also be found occasionally. In hepatitis C and D, antibodies to LKM may occur; however, the species of LKM antibodies in the two types of hepatitis are different from each other as well as from the LKM antibody species characteristic of autoimmune hepatitis type 2 (Chap. 362). The autoantibodies in viral hepatitis are nonspecific and can also be associated with other viral and systemic diseases. In contrast, virus-specific antibodies, which appear during and after hepatitis virus infection, are serologic markers of diagnostic importance.

As described above, serologic tests are available routinely with which to establish a diagnosis of hepatitis A, B, D, and C. Tests for fecal or serum HAV are not routinely available. Therefore, a diagnosis of hepatitis A is based on detection of IgM anti-HAV during acute illness (Fig. 360-2). Rheumatoid factor can give rise to false-positive results in this test.

A diagnosis of HBV infection can usually be made by detection of HBsAg in serum. Infrequently, levels of HBsAg are too low to be detected during acute HBV infection, even with contemporary, highly sensitive immunoassays. In such cases, the diagnosis can be established by the presence of IgM anti-HBc.

The titer of HBsAg bears little relation to the severity of clinical disease. Indeed, an inverse correlation exists between the serum concentration of HBsAg and the degree of liver cell damage. For example, titers are highest in immunosuppressed patients, lower in patients with chronic liver disease (but higher in mild chronic than in severe chronic hepatitis), and very low in patients with acute fulminant hepatitis. These observations suggest that, in hepatitis B, the degree of liver cell damage and the clinical course are related to variations in the patient's immune response to HBV rather than to the amount of circulating HBsAg. In immunocompetent persons, however, a correlation exists between markers of HBV replication and liver injury (see below).

Another important serologic marker in patients with hepatitis B is HBeAg. Its principal clinical usefulness is as an indicator of relative infectivity. Because HBeAg is invariably present during early acute hepatitis B, HBeAg testing is indicated primarily in chronic infection.

In patients with hepatitis B surface antigenemia of unknown duration (e.g., blood donors found to be HBsAg-positive) testing for IgM anti-HBc may be useful to distinguish between acute or recent infection (IgM anti-HBc-positive) and chronic HBV infection (IgM anti-HBc-negative, IgG anti-HBc-positive). A false-positive test for IgM anti-HBc may be encountered in patients with high-titer rheumatoid factor. Also, IgM anti-HBc may be reexpressed during acute reactivation of chronic hepatitis B.

Anti-HBs is rarely detectable in the presence of HBsAg in patients with *acute* hepatitis B, but 10–20% of persons with *chronic* HBV infection may harbor low-level anti-HBs. This antibody is directed not against the common group determinant, *a*, but against the heterotypic subtype determinant (e.g., HBsAg of subtype *ad* with anti-HBs of subtype *y*). In most cases, this serologic pattern cannot be attributed to infection with two different HBV subtypes, and the presence of this antibody is not a harbinger of imminent HBsAg clearance. When such antibody is detected, its presence is of no recognized clinical significance (see “Virology and Etiology”).

After immunization with hepatitis B vaccine, which consists of HBsAg alone, anti-HBs is the only serologic marker to appear. The commonly encountered serologic patterns of hepatitis B and their interpretations are summarized in Table 360-5. Tests for the detection of HBV DNA in liver and serum are now available. Like HBeAg, serum HBV DNA is an indicator of HBV replication, but tests for HBV DNA are more sensitive and quantitative. First-generation hybridization assays for HBV DNA had a sensitivity of 10^5 – 10^6 virions/mL, a relative threshold below which infectivity and liver injury are limited and HBeAg is usually undetectable. Currently, testing for HBV DNA has shifted from insensitive hybridization assays to amplification assays (e.g., the PCR-based assay, which can detect as few as 10 or 100 virions/mL); among the commercially available PCR assays, the most useful are those with the highest sensitivity (5–10 IU/mL) and the largest dynamic range (10^0 – 10^9 IU/mL). With increased sensitivity,

TABLE 360-5 COMMONLY ENCOUNTERED SEROLOGIC PATTERNS OF HEPATITIS B INFECTION

HBsAg	Anti-HBs	Anti-HBc	HBeAg	Anti-HBe	Interpretation
+	–	IgM	+	–	Acute hepatitis B, high infectivity ^a
+	–	IgG	+	–	Chronic hepatitis B, high infectivity
+	–	IgG	–	+	1. Late acute or chronic hepatitis B, low infectivity 2. HBeAg-negative (“precore-mutant”) hepatitis B (chronic or, rarely, acute)
+	+	+	+/-	+/-	1. HBsAg of one subtype and heterotypic anti-HBs (common) 2. Process of seroconversion from HBsAg to anti-HBs (rare)
–	–	IgM	+/-	+/-	1. Acute hepatitis B ^a 2. Anti-HBc “window”
–	–	IgG	–	+/-	1. Low-level hepatitis B carrier 2. Hepatitis B in remote past
–	+	IgG	–	+/-	Recovery from hepatitis B
–	+	–	–	–	1. Immunization with HBsAg (after vaccination) 2. Hepatitis B in the remote past (?) 3. False-positive

^aIgM anti-HBc may reappear during acute reactivation of chronic hepatitis B.

Note: See text for abbreviations.

amplification assays remain reactive well below the current 10^3 – 10^4 IU/mL threshold for infectivity and liver injury. These markers are useful in following the course of HBV replication in patients with chronic hepatitis B receiving antiviral chemotherapy (Chap. 362). Except for the early decades of life after perinatally acquired HBV infection (see above), in immunocompetent adults with chronic hepatitis B, a general correlation exists between the level of HBV replication, as reflected by the level of serum HBV DNA, and the degree of liver injury. High-serum HBV DNA levels, increased expression of viral antigens, and necroinflammatory activity in the liver go hand in hand unless immunosuppression interferes with cytolytic T cell responses to virus-infected cells; reduction of HBV replication with antiviral drugs tends to be accompanied by an improvement in liver histology. Among patients with chronic hepatitis B, high levels of HBV DNA increase the risk of cirrhosis, hepatic decompensation, and hepatocellular carcinoma (see “Complications and Sequelae”).

In patients with hepatitis C, an episodic pattern of aminotransferase elevation is common. A specific serologic diagnosis of hepatitis C can be made by demonstrating the presence in serum of anti-HCV. When contemporary immunoassays are used, anti-HCV can be detected in acute hepatitis C during the initial phase of elevated aminotransferase activity and remains detectable after recovery (rare) and during chronic infection (common). Nonspecificity can confound immunoassays for anti-HCV, especially in persons with a low prior probability of infection, such as volunteer blood donors, or in persons with circulating rheumatoid factor, which can bind nonspecifically to assay reagents; testing for HCV RNA can be used in such settings to distinguish between true-positive and false-positive anti-HCV determinations. Assays for HCV RNA are the most sensitive tests for HCV infection and represent the “gold standard” in establishing a diagnosis of hepatitis C. HCV RNA can be detected even before acute elevation of aminotransferase activity and before the appearance of anti-HCV in patients with acute hepatitis C. In addition, HCV RNA remains detectable indefinitely, continuously in most but intermittently in some, in patients with chronic hepatitis C (detectable as well in some persons with normal liver tests, i.e., inactive carriers). In the very small minority of patients with hepatitis C who lack anti-HCV, a diagnosis can be supported by detection of HCV RNA. If all these tests are negative and the patient has a well-characterized case of hepatitis after percutaneous exposure to blood or blood products, a diagnosis of hepatitis caused by an unidentified agent can be entertained.

Amplification techniques are required to detect HCV RNA, and two types are available. One is a branched-chain complementary DNA (bDNA) assay, in which the detection signal (a colorimetrically

detectable enzyme bound to a complementary DNA probe) is amplified. The other involves target amplification (i.e., synthesis of multiple copies of the viral genome) by PCR or TMA, in which the viral RNA is reverse transcribed to complementary DNA and then amplified by repeated cycles of DNA synthesis. Both can be used as quantitative assays and a measurement of relative “viral load”; PCR and TMA, with a sensitivity of 10 – 10^2 IU/mL, are more sensitive than bDNA, with a sensitivity of 10^3 IU/mL; assays are available with a wide dynamic range (10 – 10^7 IU/mL). Determination of HCV RNA level is not a reliable marker of disease severity or prognosis but is helpful in predicting relative responsiveness to antiviral therapy. The same is true for determinations of HCV genotype (Chap. 362).

A proportion of patients with hepatitis C have isolated anti-HBc

in their blood, a reflection of a common risk in certain populations of exposure to multiple bloodborne hepatitis agents. The anti-HBc in such cases is almost invariably of the IgG class and usually represents HBV infection in the remote past (HBV DNA undetectable); it rarely represents current HBV infection with low-level virus carriage.

The presence of HDV infection can be identified by demonstrating intrahepatic HDV antigen or, more practically, an anti-HDV seroconversion (a rise in titer of anti-HDV or de novo appearance of anti-HDV). Circulating HDV antigen, also diagnostic of acute infection, is detectable only briefly, if at all. Because anti-HDV is often undetectable once HBsAg disappears, retrospective serodiagnosis of acute self-limited, simultaneous HBV and HDV infection is difficult. Early diagnosis of acute infection may be hampered by a delay of up to 30–40 days in the appearance of anti-HDV.

When a patient presents with acute hepatitis and has HBsAg and anti-HDV in serum, determination of the class of anti-HBc is helpful in establishing the relationship between infection with HBV and HDV. Although IgM anti-HBc does not distinguish *absolutely* between acute and chronic HBV infection, its presence is a reliable indicator of recent infection and its absence a reliable indicator of infection in the remote past. In simultaneous acute HBV and HDV infections, IgM anti-HBc will be detectable, whereas in acute HDV infection superimposed on chronic HBV infection, anti-HBc will be of the IgG class. Tests for the presence of HDV RNA are useful for determining the presence of ongoing HDV replication and relative infectivity.

The serologic/virologic course of events during acute hepatitis E is entirely analogous to that of acute hepatitis A, with brief fecal shedding of virus and viremia and an early IgM anti-HEV response that predominates during approximately the first 3 months but is eclipsed thereafter by long-lasting IgG anti-HEV. Diagnostic tests of varying reliability for hepatitis E are commercially available but used routinely primarily outside the United States; in the United States, diagnostic serologic/virologic assays can be performed at the Centers for Disease Control and Prevention or other specialized reference laboratories.

Liver biopsy is rarely necessary or indicated in acute viral hepatitis, except when the diagnosis is questionable or when clinical evidence suggests a diagnosis of chronic hepatitis.

A diagnostic algorithm can be applied in the evaluation of cases of acute viral hepatitis. A patient with acute hepatitis should undergo four serologic tests, HBsAg, IgM anti-HAV, IgM anti-HBc, and anti-HCV (Table 360-6). The presence of HBsAg, with or without IgM anti-HBc, represents HBV infection. If IgM anti-HBc is present, the HBV infection is considered acute; if IgM anti-HBc is absent, the HBV infection is considered chronic. A diagnosis of acute hepatitis B can be made in

TABLE 360-6 SIMPLIFIED DIAGNOSTIC APPROACH IN PATIENTS PRESENTING WITH ACUTE HEPATITIS

Serologic Tests of Patient's Serum				
HBsAg	IgM Anti-HAV	IgM Anti-HBc	Anti-HCV	Diagnostic Interpretation
+	-	+	-	Acute hepatitis B
+	-	-	-	Chronic hepatitis B
+	+	-	-	Acute hepatitis A superimposed on chronic hepatitis B
+	+	+	-	Acute hepatitis A and B
-	+	-	-	Acute hepatitis A
-	+	+	-	Acute hepatitis A and B (HBsAg below detection threshold)
-	-	+	-	Acute hepatitis B (HBsAg below detection threshold)
-	-	-	+	Acute hepatitis C

Note: See text for abbreviations.

the absence of HBsAg when IgM anti-HBc is detectable. A diagnosis of acute hepatitis A is based on the presence of IgM anti-HAV. If IgM anti-HAV coexists with HBsAg, a diagnosis of simultaneous HAV and HBV infections can be made; if IgM anti-HBc (with or without HBsAg) is detectable, the patient has simultaneous acute hepatitis A and B, and if IgM anti-HBc is undetectable, the patient has acute hepatitis A superimposed on chronic HBV infection. The presence of anti-HCV supports a diagnosis of acute hepatitis C. Occasionally, testing for HCV RNA or repeat anti-HCV testing later during the illness is necessary to establish the diagnosis. Absence of all serologic markers is consistent with a diagnosis of "non-A, non-B, non-C" hepatitis, if the epidemiologic setting is appropriate.

In patients with chronic hepatitis, initial testing should consist of HBsAg and anti-HCV. Anti-HCV supports and HCV RNA testing establishes the diagnosis of chronic hepatitis C. If a serologic diagnosis of chronic hepatitis B is made, testing for HBeAg and anti-HBe is indicated to evaluate relative infectivity. Testing for HBV DNA in such patients provides a more quantitative and sensitive measure of the level of virus replication and, therefore, is very helpful during antiviral therapy (Chap. 362). In patients with chronic hepatitis B and normal aminotransferase activity in the absence of HBeAg, serial testing over time is often required to distinguish between inactive carriage and HBeAg-negative chronic hepatitis B with fluctuating virologic and necroinflammatory activity. In persons with hepatitis B, testing for anti-HDV is useful in those with severe and fulminant disease, with severe chronic disease, with chronic hepatitis B and acute hepatitis-like exacerbations, with frequent percutaneous exposures, and from areas where HDV infection is endemic.

PROGNOSIS

Virtually all previously healthy patients with hepatitis A recover completely with no clinical sequelae. Similarly, in acute hepatitis B, 95–99% of previously healthy adults have a favorable course and recover completely. Certain clinical and laboratory features, however, suggest a more complicated and protracted course. Patients of advanced age and with serious underlying medical disorders may have a prolonged course and are more likely to experience severe hepatitis. Initial presenting features such as ascites, peripheral edema, and symptoms of hepatic encephalopathy suggest a poorer prognosis. In addition, a prolonged PT, low serum albumin level, hypoglycemia, and very high serum bilirubin values suggest severe hepatocellular disease. Patients with these clinical and laboratory features deserve prompt hospital admission. The case fatality rate in hepatitis A and B is very low (~0.1%) but is increased by advanced age and underlying debilitating disorders. Among patients ill enough to be hospitalized for acute hepatitis B, the fatality rate is 1%. Hepatitis C is less severe during the acute phase than hepatitis B and is more likely to be anicteric; fatalities are rare, but the precise case fatality rate is not known. In outbreaks of waterborne hepatitis E in India and Asia, the case fatality rate is 1–2%

and up to 10–20% in pregnant women. Contributing to fulminant hepatitis E in endemic countries are instances of acute hepatitis E superimposed on underlying chronic liver disease ("acute-on-chronic" liver disease). Patients with simultaneous acute hepatitis B and hepatitis D do not necessarily experience a higher mortality rate than do patients with acute hepatitis B alone; however, in several outbreaks of acute simultaneous HBV and HDV infection among injection drug users, the case fatality rate was ~5%. When HDV superinfection occurs in a person with chronic hepatitis B, the likelihood of fulminant hepatitis and death is increased substantially. Although the case fatality rate for hepatitis D is not known definitively, in outbreaks of severe HDV superinfection in isolated populations with a high hepatitis B carrier rate, a mortality rate >20% has been recorded.

COMPLICATIONS AND SEQUELAE

A small proportion of patients with hepatitis A experience *relapsing hepatitis* weeks to months after apparent recovery from acute hepatitis. Relapses are characterized by recurrence of symptoms, aminotransferase elevations, occasionally jaundice, and fecal excretion of HAV. Another unusual variant of acute hepatitis A is *cholestatic hepatitis*, characterized by protracted cholestatic jaundice and pruritus. Rarely, liver test abnormalities persist for many months, even up to a year. Even when these complications occur, hepatitis A remains self-limited and does not progress to chronic liver disease. During the prodromal phase of acute hepatitis B, a serum sickness–like syndrome characterized by arthralgia or arthritis, rash, angioedema, and rarely, hematuria and proteinuria may develop in 5–10% of patients. This syndrome occurs before the onset of clinical jaundice, and these patients are often diagnosed erroneously as having rheumatologic diseases. The diagnosis can be established by measuring serum aminotransferase levels, which are almost invariably elevated, and serum HBsAg. As noted above, EMC is an immune-complex disease that can complicate chronic hepatitis C and is part of a spectrum of B cell lymphoproliferative disorders, which, in rare instances, can evolve to B cell lymphoma (Chap. 134). Attention has been drawn as well to associations between hepatitis C and such cutaneous disorders as porphyria cutanea tarda and lichen planus. A mechanism for these associations is unknown. Finally, related to the reliance of HCV on lipoprotein secretion and assembly pathways and on interactions of HCV with glucose metabolism, HCV infection may be complicated by hepatic steatosis, hypercholesterolemia, insulin resistance (and other manifestations of the metabolic syndrome), and type 2 diabetes mellitus; both hepatic steatosis and insulin resistance appear to accelerate hepatic fibrosis and blunt responsiveness to antiviral therapy (Chap. 362).

The most feared complication of viral hepatitis is *fulminant hepatitis* (massive hepatic necrosis); fortunately, this is a rare event. Fulminant hepatitis is seen primarily in hepatitis B, D, and E, but rare fulminant cases of hepatitis A occur primarily in older adults and in persons with underlying chronic liver disease, including, according to some reports, chronic hepatitis B and C. Hepatitis B accounts for >50% of fulminant cases of viral hepatitis, a sizable proportion of which are associated with HDV infection and another proportion with underlying chronic hepatitis C. Fulminant hepatitis is hardly ever seen in hepatitis C, but hepatitis E, as noted above, can be complicated by fatal fulminant hepatitis in 1–2% of all cases and in up to 20% of cases in pregnant women. Patients usually present with signs and symptoms of encephalopathy that may evolve to deep coma. The liver is usually small and the PT excessively prolonged. The combination of rapidly shrinking liver size, rapidly rising bilirubin level, and marked prolongation of the PT, even as aminotransferase levels fall, together with clinical signs of confusion, disorientation, somnolence, ascites, and edema, indicates that the patient has hepatic failure with encephalopathy. Cerebral edema is common; brainstem compression, gastrointestinal bleeding, sepsis, respiratory failure, cardiovascular collapse, and renal failure are terminal events. The mortality rate is exceedingly high (>80% in patients with deep coma), but patients who survive may have a complete biochemical and histologic recovery. If a donor liver can be located in time, liver transplantation may be life-saving in patients with fulminant hepatitis (Chap. 368).

Documenting the disappearance of HBsAg after apparent clinical recovery from acute hepatitis B is particularly important. Before laboratory methods were available to distinguish between acute hepatitis and acute hepatitis-like exacerbations (*spontaneous reactivations*) of chronic hepatitis B, observations suggested that ~10% of previously healthy patients remained HBsAg-positive for >6 months after the onset of clinically apparent acute hepatitis B. One-half of these persons cleared the antigen from their circulations during the next several years, but the other 5% remained chronically HBsAg-positive. More recent observations suggest that the true rate of chronic infection after clinically apparent acute hepatitis B is as low as 1% in normal, immunocompetent, young adults. Earlier, higher estimates may have been confounded by inadvertent inclusion of acute exacerbations in chronically infected patients; these patients, chronically HBsAg-positive before exacerbation, were unlikely to seroconvert to HBsAg-negative thereafter. Whether the rate of chronicity is 10% or 1%, such patients have IgG anti-HBc in serum; anti-HBs is either undetected or detected at low titer against the opposite subtype specificity of the antigen (see “Laboratory Features”). These patients may (1) be inactive carriers; (2) have low-grade, mild chronic hepatitis; or (3) have moderate to severe chronic hepatitis with or without cirrhosis. The likelihood of remaining chronically infected after acute HBV infection is especially high among neonates, persons with Down’s syndrome, chronically hemodialyzed patients, and immunosuppressed patients, including persons with HIV infection.

Chronic hepatitis is an important late complication of acute hepatitis B occurring in a small proportion of patients with acute disease but more common in those who present with chronic infection without having experienced an acute illness, as occurs typically after neonatal infection or after infection in an immunosuppressed host (**Chap. 362**). The following clinical and laboratory features suggest progression of acute hepatitis to chronic hepatitis: (1) lack of complete resolution of clinical symptoms of anorexia, weight loss, fatigue, and the persistence of hepatomegaly; (2) the presence of bridging/interface or multilobular hepatic necrosis on liver biopsy during protracted, severe acute viral hepatitis; (3) failure of the serum aminotransferase, bilirubin, and globulin levels to return to normal within 6–12 months after the acute illness; and (4) the persistence of HBeAg for >3 months or HBsAg for >6 months after acute hepatitis.

Although acute hepatitis D infection does not increase the likelihood of chronicity of simultaneous acute hepatitis B, hepatitis D has the potential for contributing to the severity of chronic hepatitis B. Hepatitis D superinfection can transform inactive or mild chronic hepatitis B into severe, progressive chronic hepatitis and cirrhosis; it also can accelerate the course of chronic hepatitis B. Some HDV superinfections in patients with chronic hepatitis B lead to fulminant hepatitis. As defined in longitudinal studies over three decades, the annual rates of cirrhosis and hepatocellular carcinoma in patients with chronic hepatitis D are 4% and 2.8%, respectively. Although HDV and HBV infections are associated with severe liver disease, mild hepatitis and even inactive carriage have been identified in some patients, and the disease may become indolent beyond the early years of infection.

After acute HCV infection, the likelihood of remaining chronically infected approaches 85–90%. Although many patients with chronic hepatitis C have no symptoms, cirrhosis may develop in as many as 20% within 10–20 years of acute illness; in some series of cases reported by referral centers, cirrhosis has been reported in as many as 50% of patients with chronic hepatitis C. Although chronic hepatitis C accounts for at least 40% of cases of chronic liver disease and of patients undergoing liver transplantation for end-stage liver disease in the United States and Europe, in the majority of patients with chronic hepatitis C, morbidity and mortality are limited during the initial 20 years after the onset of infection. Progression of chronic hepatitis C may be influenced by advanced age of acquisition, long duration of infection, immunosuppression, coexisting excessive alcohol use, concomitant hepatic steatosis, other hepatitis virus infection, or HIV coinfection. In fact, instances of severe and rapidly progressive chronic hepatitis B and C are being recognized with increasing frequency in patients with HIV infection (**Chap. 226**). In contrast, neither HAV

nor HEV causes chronic liver disease in immunocompetent hosts; however, cases of chronic hepatitis E have been observed in immunosuppressed organ-transplant recipients, persons receiving cytotoxic chemotherapy, and persons with HIV infection.

Rare complications of viral hepatitis include pancreatitis, myocarditis, atypical pneumonia, aplastic anemia, transverse myelitis, and peripheral neuropathy. Persons with chronic hepatitis B, particularly those infected in infancy or early childhood and especially those with HBeAg and/or high-level HBV DNA, have an enhanced risk of hepatocellular carcinoma. The risk of hepatocellular carcinoma is increased as well in patients with chronic hepatitis C, almost exclusively in patients with cirrhosis, and almost always after at least several decades, usually after three decades of disease (**Chap. 111**). In children, hepatitis B may present rarely with anicteric hepatitis, a nonpruritic papular rash of the face, buttocks, and limbs, and lymphadenopathy (papular acrodermatitis of childhood or Gianotti-Crosti syndrome).

Rarely, autoimmune hepatitis (**Chap. 362**) can be triggered by a bout of otherwise self-limited acute hepatitis, as reported after acute hepatitis A, B, and C.

DIFFERENTIAL DIAGNOSIS

Viral diseases such as infectious mononucleosis; those due to cytomegalovirus, herpes simplex, and coxsackieviruses; and toxoplasmosis may share certain clinical features with viral hepatitis and cause elevations in serum aminotransferase and, less commonly, in serum bilirubin levels. Tests such as the differential heterophile and serologic tests for these agents may be helpful in the differential diagnosis if HBsAg, anti-HBc, IgM anti-HAV, and anti-HCV determinations are negative. Aminotransferase elevations can accompany almost any systemic viral infection; other rare causes of liver injury confused with viral hepatitis are infections with *Leptospira*, *Candida*, *Brucella*, *Mycobacteria*, and *Pneumocystis*. A complete drug history is particularly important because many drugs and certain anesthetic agents can produce a picture of either acute hepatitis or cholestasis (**Chap. 361**). Equally important is a past history of unexplained “repeated episodes” of acute hepatitis. This history should alert the physician to the possibility that the underlying disorder is chronic hepatitis. Alcoholic hepatitis must also be considered, but usually the serum aminotransferase levels are not as markedly elevated, and other stigmata of alcoholism may be present. The finding on liver biopsy of fatty infiltration, a neutrophilic inflammatory reaction, and “alcoholic hyaline” would be consistent with alcohol-induced rather than viral liver injury. Because acute hepatitis may present with right upper quadrant abdominal pain, nausea and vomiting, fever, and icterus, it is often confused with acute cholecystitis, common duct stone, or ascending cholangitis. Patients with acute viral hepatitis may tolerate surgery poorly; therefore, it is important to exclude this diagnosis, and in confusing cases, a percutaneous liver biopsy may be necessary before laparotomy. Viral hepatitis in the elderly is often misdiagnosed as obstructive jaundice resulting from a common duct stone or carcinoma of the pancreas. Because acute hepatitis in the elderly may be quite severe and the operative mortality high, a thorough evaluation including biochemical tests, radiographic studies of the biliary tree, and even liver biopsy may be necessary to exclude primary parenchymal liver disease. Another clinical constellation that may mimic acute hepatitis is right ventricular failure with passive hepatic congestion or hypoperfusion syndromes, such as those associated with shock, severe hypotension, and severe left ventricular failure. Also included in this general category is any disorder that interferes with venous return to the heart, such as right atrial myxoma, constrictive pericarditis, hepatic vein occlusion (Budd-Chiari syndrome), or venoocclusive disease. Clinical features are usually sufficient to distinguish among these vascular disorders and viral hepatitis. Acute fatty liver of pregnancy, cholestasis of pregnancy, eclampsia, and the HELLP (*h* emolysis, *e* levated *l* iver tests, and *l* ow *p* latelets) syndrome can be confused with viral hepatitis during pregnancy. Very rarely, malignancies metastatic to the liver can mimic acute or even fulminant viral hepatitis. Occasionally, genetic or metabolic liver disorders (e.g., Wilson’s disease, α_1 antitrypsin deficiency) and nonalcoholic fatty liver disease are confused with acute viral hepatitis.

In hepatitis B, among previously healthy adults who present with clinically apparent acute hepatitis, recovery occurs in ~99%; therefore, antiviral therapy is not likely to improve the rate of recovery and is not required. In rare instances of severe acute hepatitis B, treatment with a nucleoside analogue at oral doses used to treat chronic hepatitis B (**Chap. 362**) has been attempted successfully. Although clinical trials have not been done to establish the efficacy or duration of this approach, most authorities would recommend institution of antiviral therapy with a nucleoside analogue (entecavir or tenofovir, the most potent and least resistance-prone agents) for severe, but not mild–moderate, acute hepatitis B. Treatment should continue until 3 months after HBsAg seroconversion or 6 months after HBeAg seroconversion.

In typical cases of acute hepatitis C, recovery is rare, progression to chronic hepatitis is the rule, and meta-analyses of small clinical trials suggest that antiviral therapy with interferon α monotherapy (3 million units SC three times a week) is beneficial, reducing the rate of chronicity considerably by inducing sustained responses in 30–70% of patients. In a German multicenter study of 44 patients with acute symptomatic hepatitis C, initiation of intensive interferon α therapy (5 million units SC daily for 4 weeks, then three times a week for another 20 weeks) within an average of 3 months after infection resulted in a sustained virologic response rate of 98%. Although treatment of acute hepatitis C is recommended, the optimum regimen, duration of therapy, and time to initiate therapy remain to be determined. Many authorities now opt for a 24-week course (beginning within 2–3 months after onset) of long-acting pegylated interferon plus the nucleoside analogue ribavirin, although the value of adding ribavirin has not been demonstrated (see **Chap. 362** for doses). Patients with jaundice and women are more likely to recover from acute hepatitis C, and now that genetic markers associated with spontaneous recovery (*IL28B* CC haplotype) versus persistence (non-CC haplotypes) have been defined, such genetic testing can help determine the need for and immediacy of treating acute hepatitis C—maintaining a high threshold for treating patients with CC and a very low threshold for early intervention in patients with non-CC genotypes. Protease inhibitor–based antiviral therapy with telaprevir or boceprevir, now approved for chronic hepatitis C, genotype 1 (**Chap. 362**), has not been approved for acute hepatitis C. Moreover, given the high efficacy of pegylated interferon–based therapy for acute hepatitis C, in all likelihood, the addition of a protease inhibitor would add costs and side effects without incremental efficacy. When, however, after 2014, all-oral, brief-duration, low-resistance antiviral regimens replace the current standard of care, the new approaches will be applied to acute hepatitis C and, potentially (pending the outcome of clinical trials), could even be used immediately after exposure (e.g., occupational) to prevent infection and the onset of hepatitis. Because of the marked reduction over the past two decades in the frequency of acute hepatitis C, opportunities to identify and treat patients with acute hepatitis C are rare, except in injection drug users and health workers who sustain hepatitis C–contaminated needle sticks. After such occupational accidents, when monitoring for ALT elevations and the presence of HCV RNA identifies acute hepatitis C (risk only ~3%), therapy should be initiated.

Notwithstanding these specific therapeutic considerations, in most cases of typical acute viral hepatitis, specific treatment generally is not necessary. Although hospitalization may be required for clinically severe illness, most patients do not require hospital care. Forced and prolonged bed rest is not essential for full recovery, but many patients will feel better with restricted physical activity. A high-calorie diet is desirable, and because many patients may experience nausea late in the day, the major caloric intake is best tolerated in the morning. Intravenous feeding is necessary in the acute stage if the patient has persistent vomiting and cannot maintain oral intake. Drugs capable of producing adverse reactions such as cholestasis and drugs metabolized by the liver should be avoided. If

severe pruritus is present, the use of the bile salt-sequestering resin cholestyramine is helpful. Glucocorticoid therapy has no value in acute viral hepatitis, even in severe cases associated with *bridging necrosis*, and may be deleterious, even increasing the risk of chronicity (e.g., of acute hepatitis B).

Physical isolation of patients with hepatitis to a single room and bathroom is rarely necessary except in the case of fecal incontinence for hepatitis A and E or uncontrolled, voluminous bleeding for hepatitis B (with or without concomitant hepatitis D) and hepatitis C. Because most patients hospitalized with hepatitis A excrete little, if any, HAV, the likelihood of HAV transmission from these patients during their hospitalization is low. Therefore, burdensome *enteric precautions* are no longer recommended. Although gloves should be worn when the bed pans or fecal material of patients with hepatitis A are handled, these precautions do not represent a departure from sensible procedure and contemporary universal precautions for all hospitalized patients. For patients with hepatitis B and hepatitis C, emphasis should be placed on blood precautions (i.e., avoiding direct, ungloved hand contact with blood and other body fluids). Enteric precautions are unnecessary. The importance of simple hygienic precautions such as hand washing cannot be overemphasized. Universal precautions that have been adopted for all patients apply to patients with viral hepatitis. Hospitalized patients may be discharged following substantial symptomatic improvement, a significant downward trend in the serum aminotransferase and bilirubin values, and a return to normal of the PT. Mild aminotransferase elevations should not be considered contraindications to the gradual resumption of normal activity.

In *fulminant hepatitis*, the goal of therapy is to support the patient by maintenance of fluid balance, support of circulation and respiration, control of bleeding, correction of hypoglycemia, and treatment of other complications of the comatose state in anticipation of liver regeneration and repair. Protein intake should be restricted, and oral lactulose or neomycin administered. Glucocorticoid therapy has been shown in controlled trials to be ineffective. Likewise, exchange transfusion, plasmapheresis, human cross-circulation, porcine liver cross-perfusion, hemoperfusion, and extracorporeal liver-assist devices have not been proven to enhance survival. Meticulous intensive care that includes prophylactic antibiotic coverage is the one factor that does appear to improve survival. Orthotopic liver transplantation is resorted to with increasing frequency, with excellent results, in patients with fulminant hepatitis (**Chap. 368**).

PROPHYLAXIS

Because application of therapy for acute viral hepatitis is limited and because antiviral therapy for chronic viral hepatitis is cumbersome, costly, and not effective in all patients (**Chap. 362**), emphasis is placed on prevention through immunization. The prophylactic approach differs for each of the types of viral hepatitis. In the past, immunoprophylaxis relied exclusively on passive immunization with antibody-containing globulin preparations purified by cold ethanol fractionation from the plasma of hundreds of normal donors. Currently, for hepatitis A, B, and E, active immunization with vaccines is the preferable approach to prevention.

Hepatitis A Both passive immunization with IG and active immunization with killed vaccines are available. All preparations of IG contain anti-HAV concentrations sufficient to be protective. When administered before exposure or during the early incubation period, IG is effective in preventing clinically apparent hepatitis A. For postexposure prophylaxis of intimate contacts (household, sexual, institutional) of persons with hepatitis A, the administration of 0.02 mL/kg is recommended as early after exposure as possible; it may be effective even when administered as late as 2 weeks after exposure. Prophylaxis is not necessary for those who have already received hepatitis A vaccine, for casual contacts (office, factory, school, or hospital), for most elderly persons, who are very likely to be immune, or for those known to have anti-HAV in their serum. In day care centers, recognition of hepatitis A in children or staff should provide a stimulus for immunoprophylaxis in the center and in the children's family members. By the time most common-source outbreaks of hepatitis A are recognized, it is

usually too late in the incubation period for IG to be effective; however, prophylaxis may limit the frequency of secondary cases. For travelers to tropical countries, developing countries, and other areas outside standard tourist routes, IG prophylaxis had been recommended before a vaccine became available. When such travel lasted <3 months, 0.02 mL/kg was given; for longer travel or residence in these areas, a dose of 0.06 mL/kg every 4–6 months was recommended. Administration of plasma-derived globulin is safe; all contemporary lots of IG are subjected to viral inactivation steps and must be free of HCV RNA as determined by PCR testing. Administration of IM lots of IG has not been associated with transmission of HBV, HCV, or HIV.

Formalin-inactivated vaccines made from strains of HAV attenuated in tissue culture have been shown to be safe, immunogenic, and effective in preventing hepatitis A. Hepatitis A vaccines are approved for use in persons who are at least 1 year old and appear to provide adequate protection beginning 4 weeks after a primary inoculation. If it can be given within 4 weeks of an expected exposure, such as by travel to an endemic area, hepatitis A vaccine is the preferred approach to *pre-exposure* immunoprophylaxis. If travel is more imminent, IG (0.02 mL/kg) should be administered at a different injection site, along with the first dose of vaccine. Because vaccination provides long-lasting protection (protective levels of anti-HAV should last 20 years after vaccination), persons whose risk will be sustained (e.g., frequent travelers or those remaining in endemic areas for prolonged periods) should be vaccinated, and vaccine should supplant the need for repeated IG injections. Shortly after its introduction, hepatitis A vaccine was recommended for children living in communities with a high incidence of HAV infection; in 1999, this recommendation was extended to include all children living in states, counties, and communities with high rates of HAV infection. As of 2006, the Advisory Committee on Immunization Practices of the U.S. Public Health Service recommended *routine hepatitis A vaccination of all children*. Other groups considered to be at increased risk for HAV infection and who are candidates for hepatitis A vaccination include military personnel, populations with cyclic outbreaks of hepatitis A (e.g., Alaskan natives), employees of day care centers, primate handlers, laboratory workers exposed to hepatitis A or fecal specimens, and patients with chronic liver disease. Because of an increased risk of fulminant hepatitis A—observed in some experiences but not confirmed in others—among patients with chronic hepatitis C, patients with chronic hepatitis C are candidates for hepatitis A vaccination, as are persons with chronic hepatitis B. Other populations whose recognized risk of hepatitis A is increased should be vaccinated, including men who have sex with men, injection drug users, persons with clotting disorders who require frequent administration of clotting-factor concentrates, persons traveling from the United States to countries with high or intermediate hepatitis A endemicity, postexposure prophylaxis for contacts of persons with hepatitis A, and household members and other close contacts of adopted children arriving from countries with high and moderate hepatitis A endemicity. Recommendations for dose and frequency differ for the two approved vaccine preparations (Table 360-7); all injections are IM. Hepatitis A vaccine has been reported to be effective in preventing secondary household and day care center-associated cases of acute hepatitis A. Because the vaccine

provides long-lasting protection and is simpler to use, in 2006, the Immunization Practices Advisory Committee of the U.S. Public Health Service favored hepatitis A vaccine to IG for postexposure prophylaxis of healthy persons age 2–40 years; for younger or older persons, for immunosuppressed patients, and for patients with chronic liver disease, IG should continue to be used. In the United States, reported mortality resulting from hepatitis A declined in parallel with hepatitis A vaccine-associated reductions in the annual incidence of new infections.

Hepatitis B Until 1982, prevention of hepatitis B was based on *passive* immunoprophylaxis either with standard Ig, containing modest levels of anti-HBs, or hepatitis B immunoglobulin (HBIG), containing high-titer anti-HBs. The efficacy of standard IG has never been established and remains questionable; even the efficacy of HBIG, demonstrated in several clinical trials, has been challenged, and its contribution appears to be in reducing the frequency of clinical *illness*, not in preventing *infection*. The first vaccine for *active* immunization, introduced in 1982, was prepared from purified, noninfectious 22-nm spherical HBsAg particles derived from the plasma of healthy HBsAg carriers. In 1987, the plasma-derived vaccine was supplanted by a genetically engineered vaccine derived from recombinant yeast. The latter vaccine consists of HBsAg particles that are nonglycosylated but are otherwise indistinguishable from natural HBsAg; two recombinant vaccines are licensed for use in the United States. Current recommendations can be divided into those for pre-exposure and postexposure prophylaxis.

For *pre-exposure* prophylaxis against hepatitis B in settings of frequent exposure (health workers exposed to blood; first-responder public safety workers; hemodialysis patients and staff; residents and staff of custodial institutions for the developmentally handicapped; injection drug users; inmates of long-term correctional facilities; persons with multiple sexual partners or who have had a sexually transmitted disease; men who have sex with men; persons such as hemophiliacs who require long-term, high-volume therapy with blood derivatives; household and sexual contacts of persons with chronic HBV infection; persons living in or traveling extensively in endemic areas; unvaccinated children under the age of 18; unvaccinated children who are Alaskan natives, Pacific Islanders, or residents in households of first-generation immigrants from endemic countries; persons born in countries with a prevalence of HBV infection $\geq 2\%$; patients with chronic liver disease; persons <age 60 with diabetes mellitus [those ≥ 60 at the discretion of their physicians]; persons with end-stage renal disease; persons with HIV infection), three IM (deltoid, not gluteal) injections of hepatitis B vaccine are recommended at 0, 1, and 6 months (other, optional schedules are summarized in Table 360-8). Pregnancy is *not* a contraindication to vaccination. In areas of low HBV endemicity such as the United States, despite the availability of safe and effective hepatitis B vaccines, a strategy of vaccinating persons in high-risk groups was not effective. The incidence of new hepatitis B cases continued to increase in the United States after the introduction of vaccines; <10% of all targeted persons in high-risk groups were actually vaccinated, and ~30% of persons with sporadic acute hepatitis B did not fall into any high-risk-group category. Therefore, to have an impact on the frequency of HBV infection in an area of low endemicity such as the United States, universal hepatitis B vaccination in childhood has been recommended. For unvaccinated children born after the implementation of universal infant vaccination, vaccination during early adolescence, at age 11–12 years, was recommended, and this recommendation has been extended to include all unvaccinated children age 0–19 years. In HBV-hyperendemic areas (e.g., Asia), universal vaccination of children has resulted in a marked 10- to 15-year decline in hepatitis B and its complications, including hepatocellular carcinoma.

The two available recombinant hepatitis B vaccines are comparable, one containing 10 μg of HBsAg (Recombivax-HB) and the other containing 20 μg of HBsAg (Engerix-B), and recommended doses for each injection vary for the two preparations (Table 360-8). Combinations of hepatitis B vaccine with other childhood vaccines are available as well (Table 360-8).

For unvaccinated persons sustaining an exposure to HBV, *post-exposure* prophylaxis with a combination of HBIG (for rapid

TABLE 360-7 HEPATITIS A VACCINATION SCHEDULES

Age, years	No. of Doses	Dose	Schedule, months
HAVRIX (GlaxoSmithKline)^a			
1–18	2	720 ELU ^b (0.5 mL)	0, 6–12
≥ 19	2	1440 ELU (1 mL)	0, 6–12
VAQTA (Merck)			
1–18	2	25 units (0.5 mL)	0, 6–18
≥ 19	2	50 units (1 mL)	0, 6–18

^aA combination of this hepatitis A vaccine and hepatitis B vaccine, TWINRIX, is licensed for simultaneous protection against both of these viruses among adults (age ≥ 18 years). Each 1-mL dose contains 720 ELU of hepatitis A vaccine and 20 μg of hepatitis B vaccine. These doses are recommended at months 0, 1, and 6. ^bEnzyme-linked immunoassay units.

Abbreviation: ELU, enzyme-linked immunoassay unit.

TABLE 360-8 PRE-EXPOSURE HEPATITIS B VACCINATION SCHEDULES

Target Group	No. of Doses	Dose	Schedule, months
RECOMBIVAX-HB (Merck)^a			
Infants, children (<1–10 years)	3	5 µg (0.5 mL)	0, 1–2, 4–6
Adolescents (11–19 years)	3 or 4	5 µg (0.5 mL)	0–2, 1–4, 4–6 or 0, 12, 24 or 0, 1, 2, 12
	<i>Or</i>		
	2	10 µg (1 mL)	0, 4–6 (age 11–15)
Adults (≥20 years)	3	10 µg (1 mL)	0–2, 1–4, 4–6
Hemodialysis patients ^b			
<20 years	3	5 µg (0.5 mL)	0, 1, 6
≥20 years	3	40 µg (4 mL)	0, 1, 6
ENGERIX-B (GlaxoSmithKline)^c			
Infants, children (<1–10 years)	3 or 4	10 µg (0.5 mL)	0, 1–2, 4–6 or 0, 1, 2, 12
Adolescents (10–19 years)	3 or 4	10 µg (0.5 mL)	0, 1–2, 4–6 or 0, 12, 24 or 0, 1, 2, 12
Adults (≥20 years)	3 or 4	20 µg (1 mL)	0–2, 1–4, 4–6 or 0, 1, 2, 12
Hemodialysis patients ^b			
<20 years	4	10 µg (0.5 mL)	0, 1, 2, 6
≥20 years	4	40 µg (2 mL)	0, 1, 2, 6

^aThis manufacturer produces a licensed combination of hepatitis B vaccine and vaccines against *Haemophilus influenzae* type b and *Neisseria meningitidis*, Comvax, for use in infants and young children. Please consult product insert for dose and schedule. ^bThis group also includes other immunocompromised persons. ^cThis manufacturer produces two licensed combination hepatitis B vaccines: (1) Twinrix, recombinant hepatitis B vaccine plus inactivated hepatitis A vaccine, is licensed for simultaneous protection against both of these viruses among adults (age ≥18 years). Each 1-mL dose contains 720 ELU of hepatitis A vaccine and 20 µg of hepatitis B vaccine. These doses are recommended at months 0, 1, and 6. (2) Pediarix, recombinant hepatitis B vaccine plus diphtheria and tetanus toxoid, pertussis, and inactivated poliovirus, is licensed for use in infants and young children. Please consult product insert for doses and schedules.

achievement of high-titer circulating anti-HBs) and hepatitis B vaccine (for achievement of long-lasting immunity as well as its apparent efficacy in attenuating clinical illness after exposure) is recommended. For *perinatal* exposure of infants born to HBsAg-positive mothers, a single dose of HBIG, 0.5 mL, should be administered IM in the thigh *immediately after birth*, followed by a complete course of three injections of recombinant hepatitis B vaccine (see doses above) to be started within the first 12 h of life. For those experiencing a direct percutaneous inoculation or transmucosal exposure to HBsAg-positive blood or body fluids (e.g., accidental *needle stick*, other mucosal penetration, or ingestion), a single IM dose of HBIG, 0.06 mL/kg, administered as soon after exposure as possible, is followed by a complete course of hepatitis B vaccine to begin within the first week. For those exposed by *sexual* contact to a patient with acute hepatitis B, a single IM dose of HBIG, 0.06 mL/kg, should be given within 14 days of exposure, to be followed by a complete course of hepatitis B vaccine. When both HBIG and hepatitis B vaccine are recommended, they may be given at the same time but at separate sites. Testing adults for anti-HBs after a course of vaccine is advisable to document the acquisition of immunity, but, because hepatitis B vaccine immunogenicity is nearly universal in infants, postvaccination anti-HBs testing of children is not recommended.

The precise duration of protection afforded by hepatitis B vaccine is unknown; however, ~80–90% of immunocompetent adult vaccinees retain protective levels of anti-HBs for at least 5 years, and 60–80% for 10 years, and protective antibody has been documented to last for at least two decades after vaccination in infancy. Thereafter and even after anti-HBs becomes undetectable, protection persists against clinical hepatitis B, hepatitis B surface antigenemia, and chronic HBV infection. Currently, *booster* immunizations are not recommended routinely, except in immunosuppressed persons who have lost

detectable anti-HBs or immunocompetent persons who sustain percutaneous HBsAg-positive inoculations after losing detectable antibody. Specifically, for hemodialysis patients, annual anti-HBs testing is recommended after vaccination; booster doses are recommended when anti-HBs levels fall to <10 mIU/mL. As noted above, for persons at risk of both hepatitis A and B, a combined vaccine is available containing 720 enzyme-linked immunoassay units (ELUs) of inactivated HAV and 20 µg of recombinant HBsAg (at 0, 1, and 6 months).

Hepatitis D Infection with hepatitis D can be prevented by vaccinating susceptible persons with hepatitis B vaccine. No product is available for immunoprophylaxis to prevent HDV superinfection in HBsAg carriers; for them, avoidance of percutaneous exposures and limitation of intimate contact with persons who have HDV infection are recommended.

Hepatitis C IG is ineffective in preventing hepatitis C and is no longer recommended for postexposure prophylaxis in cases of perinatal, needle stick, or sexual exposure. Although prototype vaccines that induce antibodies to HCV envelope proteins have been developed, currently, hepatitis C vaccination is not feasible practically. Genotype and quasi-species viral heterogeneity as well as rapid evasion of neutralizing antibodies by this rapidly mutating virus, conspire to render HCV a difficult target for immunoprophylaxis with a vaccine. Prevention of transfusion-associated hepatitis C has been accomplished by the following successively introduced measures: exclusion of commercial blood donors and reliance on a volunteer blood supply; screening donor blood with surrogate markers such as ALT (no longer recommended) and anti-HBc, markers that identify segments of the blood donor population with an increased risk of bloodborne infections; exclusion of blood donors in high-risk groups for AIDS and the introduction of anti-HIV screening tests; and progressively sensitive serologic and virologic screening tests for HCV infection.

In the absence of active or passive immunization, prevention of hepatitis C includes behavior changes and precautions to limit exposures to infected persons. Recommendations designed to identify patients with clinically inapparent hepatitis as candidates for medical management have as a secondary benefit the identification of persons whose contacts could be at risk of becoming infected. A so-called look-back program has been recommended to identify persons who were transfused before 1992 with blood from a donor found subsequently to have hepatitis C. In addition, anti-HCV testing is recommended for persons born between 1945 and 1965, anyone who received a blood transfusion or a transplanted organ before the introduction of second-generation screening tests in 1992, those who ever used injection drugs (or took other illicit drugs by noninjection routes), chronically hemodialyzed patients, persons with clotting disorders who received clotting factors made before 1987 from pooled blood products, persons with elevated aminotransferase levels, health workers exposed to HCV-positive blood or contaminated needles, recipients of blood or organs from a donor found to be positive for hepatitis C, persons with HIV infection, health care and public safety personnel following a needle stick or other nonpercutaneous exposure to HCV-infected material, sexual partners of persons with hepatitis C, and children born to HCV-positive mothers (Table 360-4).

For stable, monogamous sexual partners, sexual transmission of hepatitis C is unlikely, and sexual barrier precautions are not recommended. For persons with multiple sexual partners or with sexually transmitted diseases, the risk of sexual transmission of hepatitis C is increased, and barrier precautions (latex condoms) are recommended. A person with hepatitis C should avoid sharing such items as razors, toothbrushes, and nail clippers with sexual partners and family members. No special precautions are recommended for babies born to mothers with hepatitis C, and breast-feeding does not have to be restricted.

Hepatitis E Whether IG prevents hepatitis E remains undetermined. Safe and effective recombinant genotype 1 vaccines, which protect against other genotypes as well, have been developed and are available in endemic areas but not in the United States.

Liver injury is a possible consequence of ingestion of any xenobiotic, including industrial toxins, pharmacologic agents, and complementary and alternative medications (CAMs). Among patients with acute liver failure, drug-induced liver injury is the most common cause, and evidence for hepatotoxicity detected during clinical trials for drug development is the most common reason for failure of compounds to reach approval status. Drug-induced liver injury requires careful history taking to identify unrecognized exposure to chemicals used in work or at home, drugs taken by prescription or bought over the counter, and herbal or dietary supplement medicines. Hepatotoxic drugs can injure the hepatocyte directly, e.g., via a free-radical or metabolic intermediate that causes peroxidation of membrane lipids and that results in liver cell injury. Alternatively, a drug or its metabolite may activate components of the innate or adaptive immune system, stimulate apoptotic pathways, or initiate damage to bile excretory pathways (Fig. 361-1). Interference with bile canalicular pumps can allow endogenous bile acids, which can injure the liver, to accumulate. Such secondary injury, in turn, may lead to necrosis of hepatocytes; injure bile ducts, producing cholestasis; or block pathways of lipid movement, inhibit protein synthesis, or impair mitochondrial oxidation of fatty acids, resulting in lactic acidosis and intracellular triglyceride accumulation (expressed histologically as microvesicular steatosis). In other instances, drug metabolites sensitize hepatocytes to toxic cytokines. The differences observed between susceptible and nonsusceptible drug recipients may be attributable to HLA haplotypes that determine binding of drug-related haptens on the cell surface as well as to polymorphisms in elaboration of competing, protective cytokines, as has been suggested for acetaminophen hepatotoxicity (see below). Immune mechanisms may include cytotoxic lymphocytes or antibody-mediated cellular cytotoxicity. In addition, a role has been shown for activation of nuclear transporters, such as the constitutive androstane receptor (CAR) or, more recently, the pregnane X receptor (PXR), in the induction of drug hepatotoxicity.

DRUG METABOLISM

Most drugs, which are water-insoluble, undergo a series of metabolic steps, culminating in a water-soluble form appropriate for renal or biliary excretion. This process begins with oxidation or methylation mediated initially by the microsomal mixed-function oxygenases, cytochrome P450 (phase I reaction), followed by glucuronidation or sulfation (phase II reaction) or inactivation by glutathione. Most drug hepatotoxicity is mediated by a phase I toxic metabolite, but glutathione depletion, precluding inactivation of harmful compounds by glutathione S-transferase, can contribute as well.

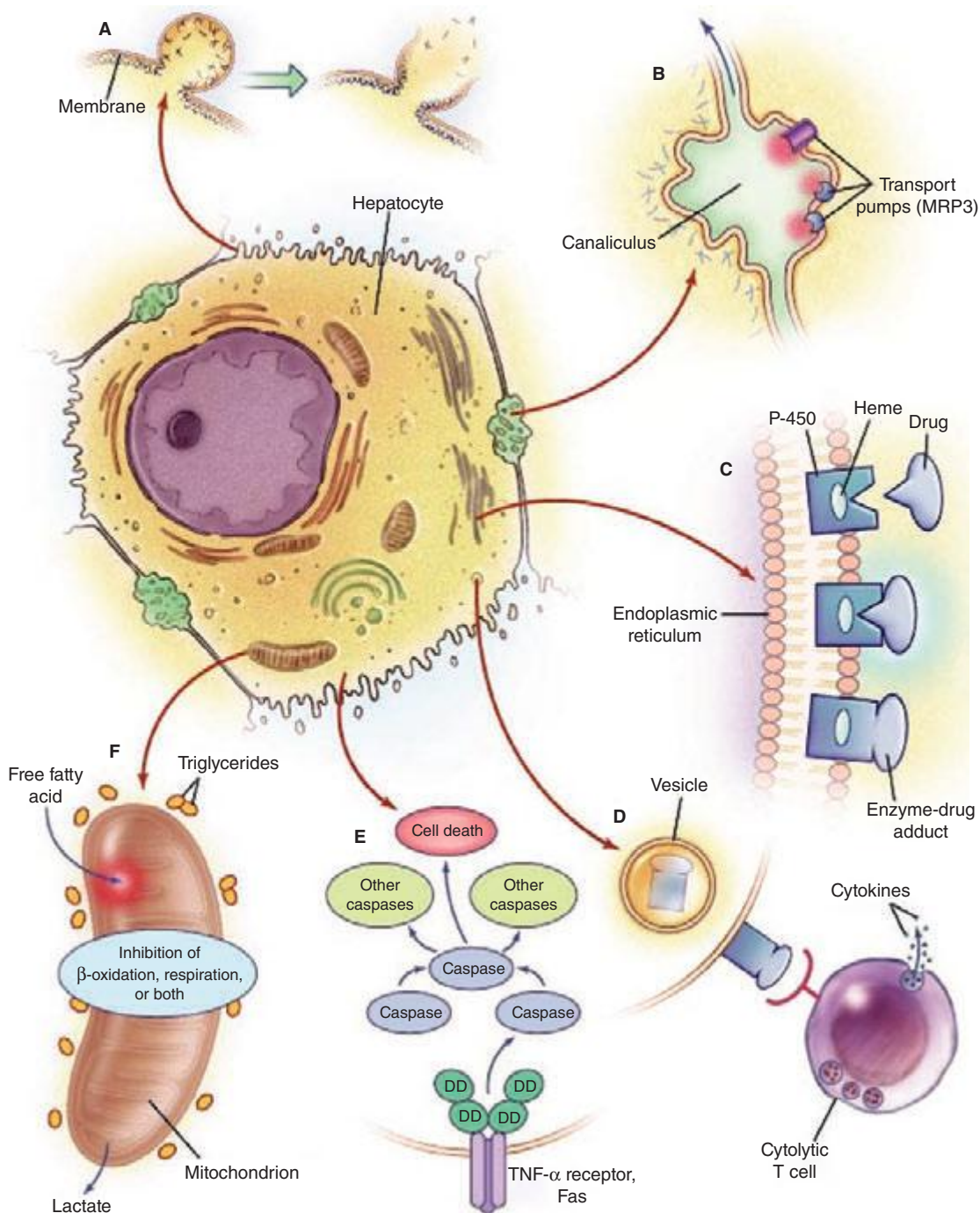
LIVER INJURY CAUSED BY DRUGS

In general, two major types of chemical hepatotoxicity have been recognized: (1) direct toxic and (2) idiosyncratic. As shown in Table 361-1, direct toxic hepatitis occurs with predictable regularity in individuals exposed to the offending agent and is dose-dependent. The latent period between exposure and liver injury is usually short (often several hours), although clinical manifestations may be delayed for 24–48 h. Agents producing toxic hepatitis are generally systemic poisons or are converted in the liver to toxic metabolites. The direct hepatotoxins result in morphologic abnormalities that are reasonably characteristic and reproducible for each toxin. For example, carbon tetrachloride and trichloroethylene characteristically produce a centrilobular zonal necrosis, whereas yellow phosphorus poisoning typically results in periportal injury. The hepatotoxic octapeptides of *Amanita phalloides* usually produce massive hepatic necrosis; the lethal dose of the toxin is ~10 mg, the amount found in a single deathcap mushroom. Liver injury, which is often only one facet of the toxicity produced by the direct hepatotoxins, may go unrecognized until jaundice appears.

In idiosyncratic drug reactions, the occurrence of hepatitis is usually infrequent (1 in 10^3 – 10^5 patients) and unpredictable; the response is not as clearly dose-dependent as is injury associated with direct hepatotoxins, and liver injury may occur at any time during or shortly after exposure to the drug. That said, recent data suggest that most agents causing idiosyncratic toxicity are given at a daily dose exceeding 100 mg, suggesting a role for dose—drugs with low potency must be given in higher doses that engender greater chances for “off-target” effects. Adding to the difficulty of predicting or identifying idiosyncratic drug hepatotoxicity is the occurrence of mild, transient, nonprogressive serum aminotransferase elevations that resolve with continued drug use. Such “adaptation,” the mechanism of which is unknown, is well recognized for drugs such as isoniazid, valproate, phenytoin, and HMG-CoA reductase inhibitors (statins). Extrahepatic manifestations of hypersensitivity, such as rash, arthralgias, fever, leukocytosis, and eosinophilia, occur in about one-quarter of patients with idiosyncratic hepatotoxic drug reactions but are characteristic for certain drugs and not others. Both primary immunologic injury and direct hepatotoxicity related to idiosyncratic differences in generation of toxic metabolites have been invoked to explain idiosyncratic drug reactions. The most current data appear to implicate the adaptive immune system responding to the formation of immune stimulatory compounds resulting from phase I metabolic activation of the offending drug. Differences in host susceptibility may result from varying kinetics of toxic metabolite generation and genetic polymorphisms in downstream drug-metabolizing pathways or cytokine activation; in addition, certain HLA haplotypes have been associated with hepatotoxicity of certain drugs such as amoxicillin-clavulanate and flucloxacillin. Occasionally, however, the clinical features of an allergic reaction (prominent tissue eosinophilia, autoantibodies, etc.) are difficult to ignore and suggest activation of IgE pathways. A few instances of drug hepatotoxicity are observed to be associated with autoantibodies, including a class of antibodies to liver-kidney microsomes, anti-LKM2, directed against a cytochrome P450 enzyme.

Idiosyncratic reactions lead to a morphologic pattern that is more variable than those produced by direct toxins; a single agent is often capable of causing a variety of lesions, although certain patterns tend to predominate. Depending on the agent involved, idiosyncratic hepatitis may result in a clinical and morphologic picture indistinguishable from that of viral hepatitis (e.g., isoniazid or ciprofloxacin). So-called hepatocellular injury is the most common form, featuring spotty necrosis in the liver lobule with a predominantly lymphocytic infiltrate resembling that observed in acute hepatitis A, B, or C. Drug-induced cholestasis ranges from mild to increasingly severe: (1) bland cholestasis with limited hepatocellular injury (e.g., estrogens, 17, α -substituted androgens); (2) inflammatory cholestasis (e.g., amoxicillin-clavulanic acid [the most frequently implicated antibiotic among cases of drug-induced liver injury], oxacillin, erythromycin estolate); (3) sclerosing cholangitis (e.g., after intrahepatic infusion of the chemotherapeutic agent floxuridine for hepatic metastases from a primary colonic carcinoma); and (4) disappearance of bile ducts, “ductopenic” cholestasis, similar to that observed in chronic rejection (Chap. 368) following liver transplantation (e.g., carbamazepine, levofloxacin). Cholestasis may result from binding of drugs to canalicular membrane transporters, accumulation of toxic bile acids resulting from canalicular pump failure, or genetic defects in canalicular transporter proteins. Clinically, the distinction between a hepatocellular and a cholestatic reaction is indicated by the R value, the ratio of alanine aminotransferase (ALT) to alkaline phosphatase values, both expressed as multiples of the upper limit of normal. An R value of >5.0 is associated with hepatocellular injury, $R < 2.0$ with cholestatic injury, and R between 2.0 and 5.0 with mixed hepatocellular-cholestatic injury.

Morphologic alterations may also include bridging hepatic necrosis (e.g., methyl dopa) or, infrequently, hepatic granulomas (e.g., sulfonamides). Some drugs result in macrovesicular or microvesicular steatosis or steatohepatitis, which, in some cases, has been linked to mitochondrial dysfunction and lipid peroxidation. Severe hepatotoxicity associated with steatohepatitis, most likely a result of mitochondrial toxicity, is being recognized with increasing frequency among patients receiving antiretroviral therapy with reverse transcriptase inhibitors for HIV infection (e.g., zidovudine, didanosine), although



- A.** Rupture of cell membrane.
B. Injury of bile canaliculus (disruption of transport pumps).
C. P-450-drug covalent binding (drug adducts).
D. Drug adducts targeted by CTLs/cytokines.
E. Activation of apoptotic pathway by TNF α /Fas.
F. Inhibition of mitochondrial function.

FIGURE 361-1 Potential mechanisms of drug-induced liver injury. The normal hepatocyte may be affected adversely by drugs through **(A)** disruption of intracellular calcium homeostasis that leads to the disassembly of actin fibrils at the surface of the hepatocyte, resulting in blebbing of the cell membrane, rupture, and cell lysis; **(B)** disruption of actin filaments next to the canaliculus (the specialized portion of the cell responsible for bile excretion), leading to loss of villous processes and interruption of transport pumps such as multidrug resistance-associated protein 3 (MRP3), which, in turn, prevents the excretion of bilirubin and other organic compounds; **(C)** covalent binding of the heme-containing cytochrome P450 enzyme to the drug, thus creating nonfunctioning adducts; **(D)** migration of these enzyme-drug adducts to the cell surface in vesicles to serve as target immunogens for cytolytic attack by T cells, stimulating an immune response involving cytolytic T cells and cytokines; **(E)** activation of apoptotic pathways by tumor necrosis factor α (TNF- α) receptor or Fas (DD denotes death domain), triggering the cascade of intercellular caspases, resulting in programmed cell death; or **(F)** inhibition of mitochondrial function by a dual effect on both β -oxidation and the respiratory-chain enzymes, leading to failure of free fatty acid metabolism, a lack of aerobic respiration, and accumulation of lactate and reactive oxygen species (which may disrupt mitochondrial DNA). Toxic metabolites excreted in bile may damage bile-duct epithelium (not shown). CTLs, cytolytic T lymphocytes. (Reproduced from WM Lee: *Drug-induced hepatotoxicity*. *N Engl J Med* 349:474, 2003, with permission.)

TABLE 361-1 SOME FEATURES OF TOXIC AND DRUG-INDUCED HEPATIC INJURY

Features	Direct Toxic Effect ^a		Idiosyncratic ^a			Other ^a
	Carbon Tetrachloride	Acetaminophen	Amoxicillin-Clavulanate	Isoniazid	Ciprofloxacin	Estrogens/Androgenic Steroids
Predictable and dose-related toxicity	+	+	0	0	0	+
Latent period	Short	Short	Delayed onset	Variable	May be short	Variable
Arthralgia, fever, rash, eosinophilia	0	0	0	0	0	0
Liver morphology	Necrosis, fatty infiltration	Centrilobular necrosis	Mixed hepatocellular/cholestatic	Hepatocellular injury resembling viral hepatitis	Hepatocellular injury resembling viral hepatitis	Cholestasis <i>without</i> portal inflammation

^aThe drugs listed are typical examples.

many of these drugs have been withdrawn because of such hepatotoxicity (Chap. 226). Generally, such mitochondrial hepatotoxicity of these antiretroviral agents is reversible, but dramatic, nonreversible hepatotoxicity associated with mitochondrial injury (inhibition of DNA polymerase γ) was the cause of acute liver failure encountered during early clinical trials of now-abandoned fialuridine, a fluorinated pyrimidine analogue with potent antiviral activity against hepatitis B virus. Another potential target for idiosyncratic drug hepatotoxicity is sinusoidal lining cells; when these are injured, such as by high-dose chemotherapeutic agents (e.g., cyclophosphamide, melphalan, busulfan) administered prior to bone marrow transplantation, venoocclusive disease can result. Nodular regenerative hyperplasia, a subtle form of portal hypertension, may also result from vascular injury to portal venous endothelium following systemic chemotherapy, such as with oxaliplatin, as part of adjuvant treatment for colon cancer.

Not all adverse hepatic drug reactions can be classified as either toxic or idiosyncratic. For example, oral contraceptives, which combine estrogenic and progestational compounds, may result in impairment of hepatic tests and, occasionally, jaundice; however, they do not produce necrosis or fatty change, manifestations of hypersensitivity are generally absent, and susceptibility to the development of oral contraceptive-induced cholestasis appears to be genetically determined. Such estrogen-induced cholestasis is more common in women with cholestasis of pregnancy, a disorder linked to genetic defects in multi-drug resistance-associated canalicular transporter proteins.

Any idiosyncratic reaction that occurs in <1:10,000 recipients will go unrecognized in most clinical trials, which involve only several thousand recipients. The U.S. Food and Drug Administration (FDA) and pharmaceutical companies have learned to look for even subtle indications of serious toxicity and monitor regularly the number of trial subjects in whom any aminotransferase elevations develop, as a possible surrogate for more serious toxicity. Even more valid as a predictor of severe hepatotoxicity is the occurrence of jaundice in patients enrolled in a clinical drug trial, so called “Hy’s Law,” named after Hyman Zimmerman, one of the pioneers of the field of drug hepatotoxicity. He recognized that, if jaundice occurred during a phase III trial, more serious liver injury was likely, with a 10:1 ratio between cases of jaundice and liver failure—10 patients with jaundice to 1 patient with acute liver failure. Thus, the finding of such Hy’s Law cases during drug development often portends failure of approval, particularly if any of the subjects sustains a bad outcome. Troglitazone, a peroxisome proliferator-activated receptor γ agonist, was the first in its class of thiazolidinedione insulin-sensitizing agents. Although in retrospect, Hy’s Law cases of jaundice had occurred during phase III trials, no instances of liver failure were recognized until well after the drug was introduced, underlining the importance of postmarketing surveillance in identifying toxic drugs and in leading to their withdrawal from use. Fortunately, such hepatotoxicity is not characteristic of the second-generation thiazolidinedione insulin-sensitizing agents rosiglitazone and pioglitazone; in clinical trials, the frequency of aminotransferase elevations in patients treated with these medications did not differ from that in placebo recipients, and isolated reports of liver injury among recipients are extremely rare.

Proving that an episode of liver injury is caused by a drug is difficult in many cases. Drug-induced liver injury is nearly always a presumptive diagnosis, and many other disorders produce a similar clinicopathologic picture. Thus, causality may be difficult to establish and requires several separate supportive assessment variables to lead to a high level of certainty, including temporal association (time of onset, time to resolution), clinical-biochemical features, type of injury (hepatocellular versus cholestatic), extrahepatic features, likelihood that a given agent is to blame based on its past record, and exclusion of other potential causes. Scoring systems such as the Roussel-Uclaf Causality Assessment Method (RUCAM) yield residual uncertainty and have not been adopted widely. Currently, the U.S. Drug-Induced Liver Injury Network (DILIN) relies on a structured expert opinion process requiring detailed data on each case and a comprehensive review by three experts who arrive at a consensus on a five-degree scale of likelihood (definite, highly likely, probable, possible, unlikely); however, this approach is not practical for routine clinical application.

Generally, drug hepatotoxicity is not more frequent in persons with underlying chronic liver disease, although the severity of the outcome may be amplified. Reported exceptions include hepatotoxicity of aspirin, methotrexate, isoniazid (only in certain experiences), antiretroviral therapy for HIV infection, and certain drugs such as conditioning regimens for bone marrow transplantation in the presence of hepatitis C.

TREATMENT TOXIC AND DRUG-INDUCED HEPATIC DISEASE

Treatment is largely supportive, except in acetaminophen hepatotoxicity (see below). In patients with fulminant hepatitis resulting from drug hepatotoxicity, liver transplantation may be lifesaving (Chap. 368). Withdrawal of the suspected agent is indicated at the first sign of an adverse reaction. A number of studies have suggested that lethal outcomes follow continued use of an agent in the face of symptoms and signs of liver injury. In the case of the direct toxins, liver involvement should not divert attention from renal or other organ involvement, which may also threaten survival. A number of agents are occasionally used but are of questionable value: glucocorticoids for drug hepatotoxicity with allergic features, silibinin for hepatotoxic mushroom poisoning, and ursodeoxycholic acid for cholestatic drug hepatotoxicity have never been shown to be effective and are not recommended.

In Table 361-2, several classes of chemical agents are listed together with examples of the pattern of liver injury produced by them. Certain drugs appear to be responsible for the development of chronic as well as acute hepatic injury. For example, nitrofurantoin, minocycline, hydralazine, and methyldopa have been associated with moderate to severe chronic hepatitis with autoimmune features. Methotrexate, tamoxifen, and amiodarone have been implicated in the development of cirrhosis. Portal hypertension in the absence of cirrhosis may result from alterations in hepatic architecture produced by vitamin A or arsenic intoxication, industrial exposure to vinyl chloride, or

TABLE 361-2 PRINCIPAL ALTERATIONS OF HEPATIC MORPHOLOGY PRODUCED BY SOME COMMONLY USED DRUGS AND CHEMICALS^a

Principal Morphologic Change	Class of Agent	Example
Cholestasis	Anabolic steroid	Methyl testosterone, many other body-building supplements
	Antibiotic	Erythromycin estolate, nitrofurantoin, rifampin, amoxicillin-clavulanic acid, oxacillin
	Anticonvulsant	Carbamazepine
	Antidepressant	Duloxetine, mirtazapine, tricyclic antidepressants
	Anti-inflammatory	Sulindac
	Antiplatelet	Clopidogrel
	Antihypertensive	Irbesartan, fosinopril
	Antithyroid	Methimazole
	Calcium channel blocker	Nifedipine, verapamil
	Immunosuppressive	Cyclosporine
	Lipid-lowering	Ezetimibe
	Oncotherapeutic	Anabolic steroids, busulfan, tamoxifen, irinotecan, cytarabine, temozolomide
	Oral contraceptive	Norethynodrel with mestranol
	Oral hypoglycemic	Chlorpropamide
Tranquilizer	Chlorpromazine ^b	
Fatty liver	Antiarrhythmic	Amiodarone
	Antibiotic	Tetracycline (high-dose, IV)
	Anticonvulsant	Valproic acid
	Antiviral	Dideoxynucleosides (e.g., zidovudine), protease inhibitors (e.g., indinavir, ritonavir)
	Oncotherapeutic	Asparaginase, methotrexate, tamoxifen
Hepatitis	Anesthetic	Halothane, fluothane
	Antiandrogen	Flutamide
	Antibiotic	Isoniazid, ^c rifampicin, nitrofurantoin, telithromycin, minocycline, ^d pyrazinamide, trovafloxacin ^e
	Anticonvulsant	Phenytoin, carbamazepine, valproic acid, phenobarbital
	Antidepressant	Ipriproniazid, amitriptyline, trazodone, venlafaxine, fluoxetine, paroxetine, duloxetine, sertraline, nefazodone ^e
	Antifungal	Ketoconazole, fluconazole, itraconazole
	Antihypertensive	Methyldopa, ^f captopril, enalapril, lisinopril, losartan
	Anti-inflammatory	Ibuprofen, indomethacin, diclofenac, sulindac, bromfenac
	Antipsychotic	Risperidone
	Antiviral	Zidovudine, didanosine, stavudine, nevirapine, ritonavir, indinavir, tipranavir, zalcitabine
	Calcium channel blocker	Nifedipine, verapamil, diltiazem
	Cholinesterase inhibitor	Tacrine
	Diuretic	Chlorothiazide
	Laxative	Oxyphenisatin ^g
	Norepinephrine-reuptake inhibitor	Atomoxetine
	Oral hypoglycemic	Troglitazone, ^h acarbose
Mixed hepatitis/cholestatic	Antibiotic	Amoxicillin-clavulanic acid, trimethoprim-sulfamethoxazole
	Antibacterial	Clindamycin
	Antifungal	Terbinafine
	Antihistamine	Cyproheptadine
	Immunosuppressive	Azathioprine
	Lipid-lowering	Nicotinic acid, lovastatin, ezetimibe
Toxic (necrosis)	Analgesic	Acetaminophen
	Hydrocarbon	Carbon tetrachloride
	Metal	Yellow phosphorus
	Mushroom	<i>Amanita phalloides</i>
	Solvent	Dimethylformamide
Granulomas	Antiarrhythmic	Quinidine, diltiazem
	Antibiotic	Sulfonamides
	Anticonvulsant	Carbamazepine
	Anti-inflammatory	Phenylbutazone
	Xanthine oxidase inhibitor	Allopurinol
Vascular injury	Chemotherapeutic	Oxaliplatin, melphalan

^aSeveral agents cause more than one type of liver lesion and appear under more than one category. ^bRarely associated with primary biliary cirrhosis–like lesion. ^cOccasionally associated with chronic hepatitis or bridging hepatic necrosis or cirrhosis. ^dAssociated with an autoimmune hepatitis–like syndrome. ^eWithdrawn from use because of severe hepatotoxicity.

administration of thorium dioxide. The latter three agents have also been associated with angiosarcoma of the liver. Oral contraceptives have been implicated in the development of hepatic adenoma and, rarely, hepatocellular carcinoma and hepatic vein occlusion (Budd-Chiari syndrome). Another unusual lesion, peliosis hepatis (blood cysts of the liver), has been observed in some patients treated with anabolic or contraceptive steroids. The existence of these hepatic disorders expands the spectrum of liver injury induced by chemical agents and emphasizes the need for a thorough drug history in all patients with liver dysfunction. A helpful LiverTox website that contains up-to-date information on drug-induced liver injury is available through the National Institute of Diabetes and Digestive and Kidney Diseases and the National Library of Medicine (www.livertox.nih.gov).

The following are patterns of adverse hepatic reactions for some prototypic agents.

ACETAMINOPHEN HEPATOTOXICITY (DIRECT TOXIN)

Acetaminophen represents the most prevalent cause of acute liver failure in the Western world; up to 72% of patients with acetaminophen hepatotoxicity in Scandinavia—somewhat lower frequencies in the United Kingdom and the United States—progress to encephalopathy and coagulopathy. Acetaminophen causes dose-related centrilobular hepatic necrosis after single-time-point ingestions, as intentional self-harm, or over extended periods, as unintentional overdoses, when multiple drug preparations or inappropriate drug amounts are used daily for several days, e.g., for relief of pain or fever. In these instances, 8 g/d, twice the daily recommended maximum dose, over several days can readily lead to liver failure. Use of opioid-acetaminophen combinations appears to be particularly harmful, because habituation to the opioid may occur with a gradual increase in opioid-acetaminophen combination dosing over days or weeks. A single dose of 10–15 g, occasionally less, may produce clinical evidence of liver injury. Fatal fulminant disease is usually (although not invariably) associated with ingestion of ≥ 25 g. Blood levels of acetaminophen correlate with severity of hepatic injury (levels >300 $\mu\text{g/mL}$ 4 h after ingestion are predictive of the development of severe damage; levels <150 $\mu\text{g/mL}$ suggest that hepatic injury is highly unlikely). Nausea, vomiting, diarrhea, abdominal pain, and shock are early manifestations occurring 4–12 h after ingestion. Then 24–48 h later, when these features are abating, hepatic injury becomes apparent. Maximal abnormalities and hepatic failure are evident 3–5 days after ingestion, and aminotransferase levels exceeding 10,000 IU/L are not uncommon (i.e., levels far exceeding those in patients with viral hepatitis). Renal failure and myocardial injury may be present. Whether or not a clear history of overdose can be elicited, clinical suspicion of acetaminophen hepatotoxicity should be raised by the presence of the extremely high aminotransferase levels in association with low bilirubin levels that are characteristic of this hyperacute injury. This biochemical signature should trigger further questioning of the subject if possible; however, denial or altered mentation may confound diagnostic efforts. In this setting, a presumptive diagnosis is reasonable, and the proven antidote, *N*-acetylcysteine—both safe and presumed to be effective even when injury has already begun to evolve—should be instituted.

Acetaminophen is metabolized predominantly by a phase II reaction to innocuous sulfate and glucuronide metabolites; however, a small proportion of acetaminophen is metabolized by a phase I reaction to a hepatotoxic metabolite formed from the parent compound by the cytochrome P450 CYP2E1. This metabolite, *N*-acetyl-*p*-benzoquinoneimine (NAPQI), is detoxified by binding to “hepatoprotective” glutathione to become harmless, water-soluble mercapturic acid, which undergoes renal excretion. When excessive amounts of NAPQI are formed, or when glutathione levels are low, glutathione levels are depleted and overwhelmed, permitting covalent binding to nucleophilic hepatocyte macromolecules forming acetaminophen-protein “adducts.” These adducts, which can be measured in serum by high-performance liquid chromatography, hold promise as diagnostic markers of acetaminophen hepatotoxicity, and a point-of-care assay for acetaminophen-Cys adducts is under development. The binding of acetaminophen to hepatocyte macromolecules is believed to

lead to hepatocyte necrosis; the precise sequence and mechanism are unknown. Hepatic injury may be potentiated by prior administration of alcohol, phenobarbital, isoniazid, or other drugs; by conditions that stimulate the mixed-function oxidase system; or by conditions such as starvation (including inability to maintain oral intake during severe febrile illnesses) that reduce hepatic glutathione levels. Alcohol induces cytochrome P450 CYP2E1; consequently, increased levels of the toxic metabolite NAPQI may be produced in chronic alcoholics after acetaminophen ingestion, but the role of alcohol in potentiating acute acetaminophen injury is still debated. Alcohol also suppresses hepatic glutathione production. Therefore, in chronic alcoholics, the toxic dose of acetaminophen may be as low as 2 g, and alcoholic patients should be warned specifically about the dangers of even standard doses of this commonly used drug. In a 2006 study, aminotransferase elevations were identified in 31–44% of normal subjects treated for 14 days with the maximal recommended dose of acetaminophen, 4 g daily (administered alone or as part of an acetaminophen-opioid combination); because these changes were transient and never associated with bilirubin elevation, the clinical relevance of these findings remains to be determined. Although underlying hepatitis C virus (HCV) infection was found to be associated with an increased risk of acute liver injury in patients hospitalized for acetaminophen overdose, generally, in patients with nonalcoholic liver disease, acetaminophen taken in recommended doses is well tolerated. Acetaminophen use in cirrhotic patients has not been associated with hepatic decompensation. On the other hand, because of the link between acetaminophen use and liver injury, and because of the limited safety margin between safe and toxic doses, the FDA has recommended that the daily dose of acetaminophen be reduced from 4 g to 3 g (even lower for persons with chronic alcohol use), that all acetaminophen-containing products be labeled prominently as containing acetaminophen, and that the potential for liver injury be prominent in the packaging of acetaminophen and acetaminophen-containing products. Within opioid combination products, the limit for the acetaminophen component has been lowered to 325 mg per tablet.

TREATMENT ACETAMINOPHEN OVERDOSAGE

Treatment includes gastric lavage, supportive measures, and oral administration of activated charcoal or cholestyramine to prevent absorption of residual drug. Neither charcoal nor cholestyramine appears to be effective if given >30 min after acetaminophen ingestion; if they are used, the stomach lavage should be done before other agents are administered orally. The chances of possible, probable, and high-risk hepatotoxicity can be derived from a nomogram plot (Fig. 361-2), readily available in emergency departments, as a function of measuring acetaminophen plasma levels 8 h after ingestion. In patients with high acetaminophen blood levels (>200 $\mu\text{g/mL}$ measured at 4 h or >100 $\mu\text{g/mL}$ at 8 h after ingestion), the administration of *N*-acetylcysteine reduces the severity of hepatic necrosis. This agent provides sulfhydryl donor groups to replete glutathione, which is required to render harmless toxic metabolites that would otherwise bind covalently via sulfhydryl linkages to cell proteins, resulting in the formation of drug metabolite-protein adducts. Therapy should be begun within 8 h of ingestion but may be at least partially effective when given as late as 24–36 h after overdose. Later administration of sulfhydryl compounds is of uncertain value. Routine use of *N*-acetylcysteine has substantially reduced the occurrence of fatal acetaminophen hepatotoxicity. *N*-acetylcysteine may be given orally but is more commonly used as an IV solution, with a loading dose of 140 mg/kg over 1 h, followed by 70 mg/kg every 4 h for 15–20 doses. Whenever a patient with potential acetaminophen hepatotoxicity is encountered, a local poison control center should be contacted. Treatment can be stopped when plasma acetaminophen levels indicate that the risk of liver damage is low. If signs of hepatic failure (e.g., progressive jaundice, coagulopathy, confusion) occur despite *N*-acetylcysteine therapy for acetaminophen hepatotoxicity, liver transplantation may be the only option. Early arterial blood lactate levels among such patients with acute liver failure

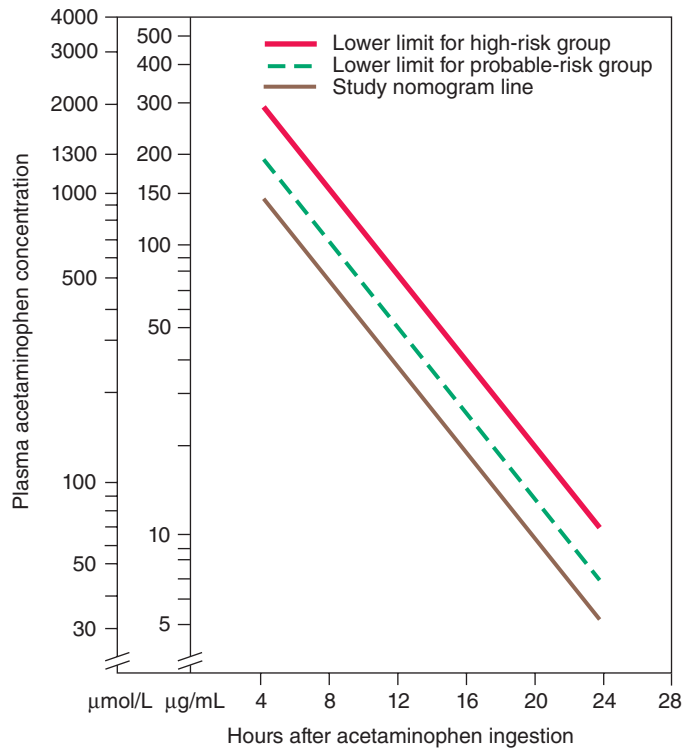


FIGURE 361-2 Nomogram to define risk of acetaminophen hepatotoxicity according to initial plasma acetaminophen concentration. (After BH Rumack, H Matthew: *Pediatrics* 55:871, 1975.)

may distinguish patients highly likely to require liver transplantation (lactate levels >3.5 mmol/L) from those likely to survive without liver replacement. Acute renal injury occurs in nearly 75% of patients with severe acetaminophen injury but is virtually always self-limited.

Survivors of acute acetaminophen overdose rarely, if ever, have ongoing liver injury or sequelae.

ISONIAZID HEPATOTOXICITY (TOXIC AND IDIOSYNCRATIC REACTION)

Isoniazid (INH) remains central to most antituberculous prophylactic and therapeutic regimens, despite its long-standing recognition as a hepatotoxin. In 10% of patients treated with INH, elevated serum aminotransferase levels develop during the first few weeks of therapy; however, these elevations in most cases are self-limited, mild (values for ALT <200 IU/L), and resolve despite continued drug use. This adaptive response allows continuation of the agent if symptoms and progressive enzyme elevations do not follow the initial elevations. Acute hepatocellular drug-induced liver injury secondary to INH is evident with a variable latency period up to 6 months and is more frequent in alcoholics and patients taking certain other medications, such as barbiturates, rifampin, and pyrazinamide. If the clinical threshold of encephalopathy is reached, severe hepatic injury is likely to be fatal or to require liver transplantation. Liver biopsy reveals morphologic changes similar to those of viral hepatitis or bridging hepatic necrosis. Substantial liver injury appears to be age-related, increasing substantially after age 35; the highest frequency is in patients over age 50, and the lowest is in patients under the age of 20. Even for patients >50 years of age monitored carefully during therapy, hepatotoxicity occurs in only ~2%, well below the risk estimate derived from earlier experiences. Fever, rash, eosinophilia, and other manifestations of drug allergy are distinctly unusual. Recently, antibodies to INH have been detected in INH recipients, but a link to causality of liver injury remains unclear. A clinical picture resembling chronic hepatitis has been observed in a few patients. Many public health programs that require INH prophylaxis for a positive tuberculin skin test or Quantiferon test include monthly monitoring of aminotransferase levels, although this practice has been called into question. Even more effective in limiting serious outcomes

may be encouraging patients to be alert for symptoms such as nausea, fatigue, or jaundice, because most fatalities occur in the setting of continued INH use despite clinically apparent illness.

SODIUM VALPROATE HEPATOTOXICITY (TOXIC AND IDIOSYNCRATIC REACTION)

Sodium valproate, an anticonvulsant useful in the treatment of petit mal and other seizure disorders, has been associated with the development of severe hepatic toxicity and, rarely, fatalities, predominantly in children but also in adults. Among children listed as candidates for liver transplantation, valproate is the most common antiepileptic drug implicated. Asymptomatic elevations of serum aminotransferase levels have been recognized in as many as 45% of treated patients. These “adaptive” changes, however, appear to have no clinical importance, because major hepatotoxicity is not seen in the majority of patients despite continuation of drug therapy. In the rare patients in whom jaundice, encephalopathy, and evidence of hepatic failure are found, examination of liver tissue reveals microvesicular fat and bridging hepatic necrosis, predominantly in the centrilobular zone. Bile duct injury may also be apparent. Most likely, sodium valproate is not directly hepatotoxic, but its metabolite, 4-pentenoic acid, may be responsible for hepatic injury. Valproate hepatotoxicity is more common in persons with mitochondrial enzyme deficiencies and may be ameliorated by IV administration of carnitine, which valproate therapy can deplete. Recently, valproate toxicity has been linked to HLA haplotypes (*DR4* and *B*1502*) and to mutations in mitochondrial DNA polymerase gamma 1.

NITROFURANTOIN HEPATOTOXICITY (IDIOSYNCRATIC REACTION)

This commonly used antibiotic for urinary tract infections may cause an acute hepatitis leading to fatal outcome or, more frequently, chronic hepatitis of varying severity but indistinguishable from autoimmune chronic hepatitis. These two scenarios may reflect the frequent use and reuse of the drug for treatment of recurrent cystitis in women. Although most toxic agents manifest injury within 6 months of first ingestion, nitrofurantoin may have a longer latency period, in part perhaps because of its intermittent, recurrent use. Autoantibodies to nuclear components, smooth muscle, and mitochondria are seen and may subside after resolution of infection; however, glucocorticoid or other immunosuppressive medication may be necessary to resolve the autoimmune injury, and cirrhosis may be seen in cases that are not recognized quickly. Interstitial pulmonary fibrosis presenting as chronic cough and dyspnea may be present and resolve slowly with medication withdrawal. Histologic findings are identical to those of autoimmune hepatitis. A similar disease pattern can be observed with minocycline that is used repeatedly for the treatment of acne in teenagers as well as with hydralazine and alpha methyl dopa.

AMOXICILLIN-CLAVULANATE HEPATOTOXICITY (IDIOSYNCRATIC MIXED REACTION)

Currently, the most common agent implicated as causing drug-induced liver injury in the United States and in Europe is amoxicillin-clavulanate (most frequent brand name: Augmentin). This medication causes a very specific syndrome of mixed or primarily cholestatic injury. Because hepatotoxicity may follow amoxicillin-clavulanate therapy after a relatively long latency period, the liver injury may begin to manifest at the time of drug withdrawal or after the drug has been withdrawn. The high prevalence of hepatotoxicity reflects in part the very frequent use of this drug for respiratory tract infections, including community-acquired pneumonia. The mechanism of hepatotoxicity is unclear, but the liver injury is thought to be caused by amoxicillin toxicity that is potentiated in some way by clavulanate, which itself appears not to be toxic. Symptoms include nausea, anorexia, fatigue, and jaundice—which may be prolonged—with pruritus. Rash is quite uncommon. On occasion, amoxicillin-clavulanate, like other cholestatic hepatotoxic drugs, causes permanent injury to small bile ducts, leading to the so-called “vanishing bile duct syndrome.” In vanishing bile duct syndrome, initially, liver injury is minimal except for severe cholestasis; however, over time, histologic evidence of bile duct

abnormalities is replaced by a paucity and eventual absence of discernible ducts on subsequent biopsies.

PHENYTOIN HEPATOTOXICITY (IDIOSYNCRATIC REACTION)

Phenytoin, formerly diphenylhydantoin, a mainstay in the treatment of seizure disorders, has been associated in rare instances with the development of severe hepatitis-like liver injury leading to fulminant hepatic failure. In many patients, the hepatitis is associated with striking fever, lymphadenopathy, rash (Stevens-Johnson syndrome or exfoliative dermatitis), leukocytosis, and eosinophilia, suggesting an immunologically mediated hypersensitivity mechanism. Despite these observations, evidence suggests that metabolic idiosyncrasy may be responsible for hepatic injury. In the liver, phenytoin is converted by cytochrome P450 to metabolites, including the highly reactive electrophilic arene oxides. These metabolites are normally metabolized further by epoxide hydrolases. A defect (genetic or acquired) in epoxide hydrolase activity could permit covalent binding of arene oxides to hepatic macromolecules, thereby leading to hepatic injury. Hepatic injury is usually manifest within the first 2 months after beginning phenytoin therapy. With the exception of an abundance of eosinophils in the liver, the clinical, biochemical, and histologic picture resembles that of viral hepatitis. In rare instances, bile duct injury may be the salient feature of phenytoin hepatotoxicity, with striking features of intrahepatic cholestasis. Asymptomatic elevations of aminotransferase and alkaline phosphatase levels have been observed in a sizable proportion of patients receiving long-term phenytoin therapy. These liver changes are believed by some authorities to represent the potent hepatic enzyme-inducing properties of phenytoin and are accompanied histologically by swelling of hepatocytes in the absence of necro-inflammatory activity or evidence of chronic liver disease.

AMIODARONE HEPATOTOXICITY (TOXIC AND IDIOSYNCRATIC REACTION)

Therapy with this potent antiarrhythmic drug is accompanied in 15–50% of patients by modest elevations of serum aminotransferase levels that may remain stable or diminish despite continuation of the drug. Such abnormalities may appear days to many months after beginning therapy. A proportion of those with elevated aminotransferase levels have detectable hepatomegaly, and clinically important liver disease develops in <5% of patients. Features that represent a direct effect of the drug on the liver and that are common to the majority of long-term recipients are ultrastructural phospholipidosis, unaccompanied by clinical liver disease, and interference with hepatic mixed-function oxidase metabolism of other drugs. The cationic amphiphilic drug and its major metabolite desethylamiodarone accumulate in hepatocyte lysosomes and mitochondria and in bile duct epithelium. The relatively common elevations in aminotransferase levels are also considered a predictable, dose-dependent, direct hepatotoxic effect. On the other hand, in the rare patient with clinically apparent, symptomatic liver disease, liver injury resembling that seen in alcoholic liver disease is observed. The so-called pseudoalcoholic liver injury can range from steatosis to alcoholic hepatitis-like neutrophilic infiltration and Mallory's hyaline to cirrhosis. Electron-microscopic demonstration of phospholipid-laden lysosomal lamellar bodies can help to distinguish amiodarone hepatotoxicity from typical alcoholic hepatitis. This category of liver injury appears to be a metabolic idiosyncrasy that allows hepatotoxic metabolites to be generated. Rarely, an acute idiosyncratic hepatocellular injury resembling viral hepatitis or cholestatic hepatitis occurs. Hepatic granulomas have occasionally been observed. Because amiodarone has a long half-life, liver injury may persist for months after the drug is stopped.

ERYTHROMYCIN HEPATOTOXICITY (CHOLESTATIC IDIOSYNCRATIC REACTION)

The most important adverse effect associated with erythromycin, more common in children than adults, is the infrequent occurrence of a cholestatic reaction. Although most of these reactions have been associated with erythromycin estolate, other erythromycins may also be responsible. The reaction usually begins during the first 2 or 3 weeks of therapy and includes nausea, vomiting, fever, right upper quadrant abdominal pain, jaundice, leukocytosis, and moderately elevated

aminotransferase and alkaline phosphatase levels. The clinical picture can resemble acute cholecystitis or bacterial cholangitis. Liver biopsy reveals variable cholestasis; portal inflammation comprising lymphocytes, polymorphonuclear leukocytes, and eosinophils; and scattered foci of hepatocyte necrosis. Symptoms and laboratory findings usually subside within a few days of drug withdrawal, and evidence of chronic liver disease has not been found on follow-up. The precise mechanism remains ill-defined.

ORAL CONTRACEPTIVE HEPATOTOXICITY (CHOLESTATIC REACTION)

The administration of oral contraceptive combinations of estrogenic and progestational steroids leads to intrahepatic cholestasis with pruritus and jaundice in a small number of patients weeks to months after taking these agents. Especially susceptible seem to be patients with recurrent idiopathic jaundice of pregnancy, severe pruritus of pregnancy, or a family history of these disorders. With the exception of liver biochemical tests, laboratory studies are normal, and extrahepatic manifestations of hypersensitivity are absent. Liver biopsy reveals cholestasis with bile plugs in dilated canaliculi and striking bilirubin staining of liver cells. In contrast to chlorpromazine-induced cholestasis, portal inflammation is absent. The lesion is reversible on withdrawal of the agent. The two steroid components appear to act synergistically on hepatic function, although the estrogen may be primarily responsible. Oral contraceptives are contraindicated in patients with a history of recurrent jaundice of pregnancy. Primarily benign, but rarely malignant, neoplasms of the liver, hepatic vein occlusion, and peripheral sinusoidal dilatation have also been associated with oral contraceptive therapy. Focal nodular hyperplasia of the liver is not more frequent among users of oral contraceptives.

ANABOLIC STEROIDS (CHOLESTATIC REACTION)

The most common form of liver injury caused by complementary and alternative medications is the profound cholestasis associated with anabolic steroids used by body builders. Unregulated agents sold in gyms and health food stores as diet supplements, which are taken by athletes to improve their performance, may contain anabolic steroids. Jaundice in a young male that is accompanied by a cholestatic, rather than a hepatitic, laboratory profile almost invariably will turn out to be caused by the use of one of a variety of androgen congeners. Such agents have the potential to injure bile transport pumps and to cause intense cholestasis; the time to onset is variable, and resolution, which is the rule, may require many weeks to months. Initially, anorexia, nausea, and malaise may occur, followed by pruritus in some but not all patients. Serum aminotransferase levels are usually <100 IU/L and serum alkaline phosphatase levels are generally moderately elevated with bilirubin levels frequently exceeding 342 $\mu\text{mol/L}$ (20 mg/dL). Examination of liver tissue reveals cholestasis without substantial inflammation or necrosis. Anabolic steroids have also been used by prescription to treat bone marrow failure. In this setting, hepatic sinusoidal dilatation and peliosis hepatis have been reported in rare patients, as have hepatic adenomas and hepatocellular carcinoma.

TRIMETHOPRIM-SULFAMETHOXAZOLE HEPATOTOXICITY (IDIOSYNCRATIC REACTION)

This antibiotic combination is used routinely for urinary tract infections in immunocompetent persons and for prophylaxis against and therapy of *Pneumocystis carinii* pneumonia in immunosuppressed persons (transplant recipients, patients with AIDS). With its increasing use, its occasional hepatotoxicity is being recognized with growing frequency. Its likelihood is unpredictable, but when it occurs, trimethoprim-sulfamethoxazole hepatotoxicity follows a relatively uniform latency period of several weeks and is often accompanied by eosinophilia, rash, and other features of a hypersensitivity reaction. Biochemically and histologically, acute hepatocellular necrosis predominates, but cholestatic features are quite frequent. Occasionally, cholestasis without necrosis occurs, and, very rarely, a severe cholangiolytic pattern of liver injury is observed. In most cases, liver injury is self-limited, but rare fatalities have been recorded. The hepatotoxicity is attributable to the sulfamethoxazole component of the drug and is similar in features to

2030 that seen with other sulfonamides; tissue eosinophilia and granulomas may be seen. The risk of trimethoprim-sulfamethoxazole hepatotoxicity is increased in persons with HIV infection.

HMG-COA REDUCTASE INHIBITORS (STATINS) (IDIOSYNCRATIC MIXED HEPATOCELLULAR AND CHOLESTATIC REACTION)

Between 1 and 2% of patients taking lovastatin, simvastatin, pravastatin, fluvastatin, or one of the newer statin drugs for the treatment of hypercholesterolemia experience asymptomatic, reversible elevations (>threefold) of aminotransferase activity. Acute hepatitis-like histologic changes, centrilobular necrosis, and centrilobular cholestasis have been described in a very small number of cases. In a larger proportion, minor aminotransferase elevations appear during the first several weeks of therapy. Careful laboratory monitoring can distinguish between patients with minor, transitory changes, who may continue therapy and those with more profound and sustained abnormalities, who should discontinue therapy. Because clinically meaningful aminotransferase elevations are so rare after statin use and do not differ in meta-analyses from the frequency of such laboratory abnormalities in placebo recipients, a panel of liver experts recommended to the National Lipid Association's Safety Task Force that liver test monitoring was not necessary in patients treated with statins and that statin therapy need not be discontinued in patients found to have asymptomatic isolated aminotransferase elevations during therapy. Statin hepatotoxicity is not increased in patients with chronic hepatitis C, hepatic steatosis, or other underlying liver diseases, and statins can be used safely in these patients.

TOTAL PARENTERAL NUTRITION (STEATOSIS, CHOLESTASIS)

Total parenteral nutrition (TPN) is often complicated by cholestatic hepatitis attributable to steatosis, cholestasis, or gallstones (or gallbladder sludge). Steatosis or steatohepatitis may result from the excess carbohydrate calories in these nutritional supplements and is the predominant form of TPN-associated liver disorder in adults. The frequency of this complication has been reduced substantially by the introduction of balanced TPN formulas that rely on lipid as an alternative caloric source. Cholestasis and cholelithiasis, caused by the absence of stimulation of bile flow and secretion resulting from the lack of oral intake, is the predominant form of TPN-associated liver disease in infants, especially in premature neonates. Often, cholestasis in such neonates is multifactorial, contributed to by other factors such as sepsis, hypoxemia, and hypotension; occasionally, TPN-induced cholestasis in neonates culminates in chronic liver disease and liver failure. When TPN-associated liver test abnormalities occur in adults, balancing the TPN formula with more lipid is the intervention of first recourse. In infants with TPN-associated cholestasis, the addition of oral feeding may ameliorate the problem. Therapeutic interventions suggested, but not shown, to be of proven benefit, include cholecystokinin, ursodeoxycholic acid, S-adenosyl methionine, and taurine.

ALTERNATIVE AND COMPLEMENTARY MEDICINES (IDIOSYNCRATIC HEPATITIS, STEATOSIS)

Herbal medications that are of scientifically unproven efficacy and that lack prospective safety oversight by regulatory agencies currently account for more than 20% of drug-induced liver injury in the United States. Besides anabolic steroids, the most common category of dietary or herbal products is weight loss agents. Included among the herbal remedies associated with toxic hepatitis are Jin Bu Huan, xiao-chai-hu-tang, germander, chaparral, senna, mistletoe, skullcap, gentian, comfrey (containing pyrrolizidine alkaloids), ma huang, bee pollen, valerian root, pennyroyal oil, kava, celandine, Impila (*Callilepis laureola*), LipoKinetix, Hydroxycut, herbal nutritional supplements, and herbal teas containing *Camellia sinensis* (green tea extract). Well characterized

are the acute hepatitis-like histologic lesions following Jin Bu Huan use: focal hepatocellular necrosis, mixed mononuclear portal tract infiltration, coagulative necrosis, apoptotic hepatocyte degeneration, tissue eosinophilia, and microvesicular steatosis. Megadoses of vitamin A can injure the liver, as can pyrrolizidine alkaloids, which often contaminate Chinese herbal preparations and can cause a venoocclusive injury leading to sinusoidal hepatic vein obstruction. Because some alternative medicines induce toxicity via active metabolites, alcohol and drugs that stimulate cytochrome P450 enzymes may enhance the toxicity of some of these products. Conversely, some alternative medicines also stimulate cytochrome P450 and may result in or amplify the toxicity of recognized drug hepatotoxins. Given the widespread use of such poorly defined herbal preparations, hepatotoxicity is likely to be encountered with increasing frequency; therefore, a drug history in patients with acute and chronic liver disease should include use of "alternative medicines" and other nonprescription preparations sold in so-called health food stores.

HIGHLY ACTIVE ANTIRETROVIRAL THERAPY (HAART) FOR HIV INFECTION (MITOCHONDRIAL TOXIC, IDIOSYNCRATIC, STEATOSIS; HEPATOCELLULAR, CHOLESTATIC, AND MIXED)

The recognition of drug hepatotoxicity in persons with HIV infection is complicated in this population by the many alternative causes of liver injury (chronic viral hepatitis, fatty infiltration, infiltrative disorders, mycobacterial infection, etc.), but drug hepatotoxicity associated with HAART is an emerging and common type of liver injury in HIV-infected persons (**Chap. 226**). Although no one antiviral agent is recognized as a potent hepatotoxin, combination regimens including reverse transcriptase and protease inhibitors cause hepatotoxicity in ~10% of treated patients. Implicated most frequently are combinations including nucleoside analogue reverse transcriptase inhibitors zidovudine, didanosine, and, to a lesser extent, stavudine; protease inhibitors ritonavir and indinavir (and amprenavir when used together with ritonavir), as well as tipranavir; and nonnucleoside reverse transcriptase inhibitors nevirapine and, to a lesser extent, efavirenz. These drugs cause predominantly hepatocellular injury but cholestatic injury as well, and prolonged (>6 months) use of reverse transcriptase inhibitors has been associated with mitochondrial injury, steatosis, and lactic acidosis. Indirect hyperbilirubinemia, resulting from direct inhibition of bilirubin-conjugating activity by UDP-glucuronosyltransferase, usually without elevation of aminotransferase or alkaline phosphatase activities, occurs in ~10% of patients treated with the protease inhibitor indinavir. Distinguishing the impact of HAART hepatotoxicity in patients with HIV and hepatitis virus co-infection is made challenging by the following: (1) both chronic hepatitis B and hepatitis C can affect the natural history of HIV infection and the response to HAART, and (2) HAART can have an impact on chronic viral hepatitis. For example, immunologic reconstitution with HAART can result in immunologically mediated liver-cell injury in patients with chronic hepatitis B co-infection if treatment with an antiviral agent for hepatitis B (e.g., the nucleoside analogue lamivudine) is withdrawn or if nucleoside analogue resistance emerges. Infection with HIV, especially with low CD4+ T cell counts, has been reported to increase the rate of hepatic fibrosis associated with chronic hepatitis C, and HAART therapy can increase levels of serum aminotransferases and HCV RNA in patients with hepatitis C co-infection. Didanosine or stavudine should not be used with ribavirin in patients with HIV/HCV co-infection because of an increased risk of severe mitochondrial toxicity and lactic acidosis.

ACKNOWLEDGMENT

Kurt J. Isselbacher, MD, contributed to this chapter in previous editions of Harrison's.

362 Chronic Hepatitis

Jules L. Dienstag

Chronic hepatitis represents a series of liver disorders of varying causes and severity in which hepatic inflammation and necrosis continue for at least 6 months. Milder forms are nonprogressive or only slowly progressive, while more severe forms may be associated with scarring and architectural reorganization, which, when advanced, lead ultimately to cirrhosis. Several categories of chronic hepatitis have been recognized. These include chronic viral hepatitis, drug-induced chronic hepatitis (Chap. 361), and autoimmune chronic hepatitis. In many cases, clinical and laboratory features are insufficient to allow assignment into one of these three categories; these “idiopathic” cases are also believed to represent autoimmune chronic hepatitis. Finally, clinical and laboratory features of chronic hepatitis are observed occasionally in patients with such hereditary/metabolic disorders as Wilson’s disease (copper overload), α_1 antitrypsin deficiency (Chaps. 365 and 429), and nonalcoholic fatty liver disease (Chap. 367e) and even occasionally in patients with alcoholic liver injury (Chap. 363). Although all types of chronic hepatitis share certain clinical, laboratory, and histopathologic features, chronic viral and chronic autoimmune hepatitis are sufficiently distinct to merit separate discussions. For discussion of acute hepatitis, see Chap. 360.

CLASSIFICATION OF CHRONIC HEPATITIS

Common to all forms of chronic hepatitis are histopathologic distinctions based on localization and extent of liver injury. These vary from the milder forms, previously labeled *chronic persistent hepatitis* and *chronic lobular hepatitis*, to the more severe form, formerly called *chronic active hepatitis*. When first defined, these designations were believed to have prognostic implications, which were not corroborated by subsequent observations. Categorization of chronic hepatitis based primarily on histopathologic features has been replaced by a more informative classification based on a combination of clinical, serologic, and histologic variables. Classification of chronic hepatitis is based on (1) its cause; (2) its histologic activity, or *grade*; and (3) its degree of progression, or *stage*. Thus, neither clinical features alone nor histologic features—requiring liver biopsy—alone are sufficient to characterize and distinguish among the several categories of chronic hepatitis.

CLASSIFICATION BY CAUSE

Clinical and serologic features allow the establishment of a diagnosis of *chronic viral hepatitis*, caused by hepatitis B, hepatitis B plus D, or hepatitis C; *autoimmune hepatitis*, including several subcategories, I and II (perhaps III), based on serologic distinctions; *drug-associated chronic hepatitis*; and a category of unknown cause, or *cryptogenic chronic hepatitis* (Table 362-1). These are addressed in more detail below.

CLASSIFICATION BY GRADE

Grade, a histologic assessment of necroinflammatory activity, is based on examination of the liver biopsy. An assessment of important histologic features includes the degree of *periportal necrosis* and the disruption of the limiting plate of periportal hepatocytes by inflammatory cells (so-called *piecemeal necrosis* or *interface hepatitis*); the degree of confluent necrosis that links or forms bridges between vascular structures—between portal tract and portal tract or even more important bridges between portal tract and central vein—referred to as *bridging necrosis*; the degree of hepatocyte degeneration and focal necrosis within the lobule; and the degree of *portal inflammation*. Several scoring systems that take these histologic features into account have been devised, and the most popular are the histologic activity index (HAI), used commonly in the United States, and the METAVIR score, used in Europe (Table 362-2). Based on the presence and degree of these features of histologic activity, chronic hepatitis can be graded as mild, moderate, or severe.

TABLE 362-1 CLINICAL AND LABORATORY FEATURES OF CHRONIC HEPATITIS

Type of Hepatitis	Diagnostic Test(s)	Autoantibodies	Therapy
Chronic hepatitis B	HBsAg, IgG anti-HBc, HBeAg, HBV DNA	Uncommon	IFN- α , PEG IFN- α Oral agents: First-line: entecavir, tenofovir Second-line: lamivudine, adefovir, telbivudine
Chronic hepatitis C	Anti-HCV, HCV RNA	Anti-LKM1 ^a	PEG IFN- α plus ribavirin Telaprevir ^b Boceprevir ^b
Chronic hepatitis D	Anti-HDV, HDV RNA, HBsAg, IgG anti-HBc	Anti-LKM3	IFN- α , PEG IFN- α ^c
Autoimmune hepatitis	ANA ^d (homogeneous), anti-LKM1 (\pm) Hyperglobulinemia	ANA, anti-LKM1, anti-SLA ^e	Prednisone, azathioprine
Drug-associated	—	Uncommon	Withdraw drug
Cryptogenic	All negative	None	Prednisone (?), azathioprine (?)

^aAntibodies to liver-kidney microsomes type 1 (autoimmune hepatitis type II and some cases of hepatitis C). ^bAdministered as a triple-drug combination with PEG IFN and ribavirin. Between the writing and publication of this chapter, two additional drugs were approved for hepatitis C, simeprevir and sofosbuvir (see www.hcvguidelines.org). ^cEarly clinical trials suggested benefit of IFN- α therapy; PEG IFN- α is as effective, if not more so, and has supplanted standard IFN- α . ^dAntinuclear antibody (autoimmune hepatitis type I). ^eAntibodies to soluble liver antigen (autoimmune hepatitis type III).

Abbreviations: HBc, hepatitis B core; HBeAg, hepatitis B e antigen; HBsAg, hepatitis B surface antigen; HBV, hepatitis B virus; HCV, hepatitis C virus; HDV, hepatitis D virus; IFN- α , interferon α ; IgG, immunoglobulin G; LKM, liver-kidney microsome; PEG IFN- α , pegylated interferon α ; SLA, soluble liver antigen.

CLASSIFICATION BY STAGE

The stage of chronic hepatitis, which reflects the level of progression of the disease, is based on the degree of hepatic fibrosis. When fibrosis is so extensive that fibrous septa surround parenchymal nodules and alter the normal architecture of the liver lobule, the histologic lesion is defined as *cirrhosis*. Staging is based on the degree of fibrosis as categorized on a numerical scale from 0–6 (HAI) or 0–4 (METAVIR) (Table 362-2). Several noninvasive approaches have been introduced to provide approximations of hepatic histologic stage, including serum biomarkers of fibrosis and imaging determinations of liver elasticity.

CHRONIC VIRAL HEPATITIS

Both the enterically transmitted forms of viral hepatitis, hepatitis A and E, are self-limited and do not cause chronic hepatitis (rare reports notwithstanding in which acute hepatitis A serves as a trigger for the onset of autoimmune hepatitis in genetically susceptible patients or in which hepatitis E (Chap. 360) can cause chronic liver disease in immunosuppressed hosts, e.g., after liver transplantation). In contrast, the entire clinicopathologic spectrum of chronic hepatitis occurs in patients with chronic viral hepatitis B and C as well as in patients with chronic hepatitis D superimposed on chronic hepatitis B.

CHRONIC HEPATITIS B

The likelihood of chronicity after acute hepatitis B varies as a function of age. Infection at birth is associated with clinically silent acute infection but a 90% chance of chronic infection, whereas infection in young adulthood in immunocompetent persons is typically associated with clinically apparent acute hepatitis but a risk of chronicity of only approximately 1%. Most cases of chronic hepatitis B among adults, however, occur in patients who never had a recognized episode of clinically apparent acute viral hepatitis. The degree of liver injury (grade) in patients with chronic hepatitis B is variable, ranging from none in inactive carriers to mild to moderate to severe. Among adults

TABLE 362-2 HISTOLOGIC GRADING AND STAGING OF CHRONIC HEPATITIS

Histologic Feature	Histologic Activity Index (HAI) ^a		METAVIR ^b		
	Severity	Score	Severity	Score	
Necroinflammatory Activity (grade)					
Periportal necrosis, including piecemeal necrosis and/or bridging necrosis (BN)	None	0	None	0	
	Mild	1	Mild	1	
	Mild/moderate	2	Moderate	2	
	Moderate	3	Severe	3	
	Severe	4			
Intralobular necrosis	Confluent	—None	None or mild	0	
		—Focal	Moderate	1	
		—Zone 3 some	Severe	2	
		—Zone 3 most			
		—Zone 3 + BN few			
	Focal	—Zone 3 + BN multiple	5		
		—Panacinar/multiacinar	6		
		—None	0		
		—≤1 focus/10× field	1		
		—2–4 foci/10× field	2		
Portal Inflammation	—5–10 foci/10× field	3			
	—>10 foci/10× field	4			
	None	0			
	Mild	1			
	Moderate	2			
	Moderate/marked	3			
	Marked	4			
	Total	0–18		A0–A3 ^c	
Fibrosis (stage)					
None		0		F0	
Portal fibrosis—some		1		F1	
Portal fibrosis—most		2		F1	
Bridging fibrosis—few		3		F2	
Bridging fibrosis—many		4		F3	
Incomplete cirrhosis		5		F4	
Cirrhosis		6		F4	
	Total	6		4	

^aIshak K, Baptista A, Bianchi L, et al: Histologic grading and staging of chronic hepatitis. *J Hepatol* 22:696, 1995. ^bBedossa P, Poynard T, French METAVIR Cooperative Study Group: An algorithm for grading activity in chronic hepatitis C. *Hepatology* 24:289, 1996. ^cNecroinflammatory grade: A0 = none; A1 = mild; A2 = moderate; A3 = severe.

with chronic hepatitis B, histologic features are of prognostic importance. In one long-term study of patients with chronic hepatitis B, investigators found a 5-year survival rate of 97% for patients with mild chronic hepatitis, 86% for patients with moderate to severe chronic hepatitis, and only 55% for patients with chronic hepatitis and postnecrotic cirrhosis. The 15-year survival in these cohorts was 77%, 66%, and 40%, respectively. On the other hand, more recent observations do not allow us to be so sanguine about the prognosis in patients with mild chronic hepatitis; among such patients followed for 1–13 years, progression to more severe chronic hepatitis and cirrhosis has been observed in more than a quarter of cases.

More important to consider than histology alone in patients with chronic hepatitis B is the degree of hepatitis B virus (HBV) replication. As reviewed in [Chap. 360](#), chronic HBV infection can occur in the presence or absence of serum hepatitis B e antigen (HBeAg), and generally, for both HBeAg-reactive and HBeAg-negative chronic hepatitis B, the level of HBV DNA correlates with the level of liver injury and risk of progression. In *HBeAg-reactive chronic hepatitis B*, two phases have been recognized based on the relative level of HBV replication.

The relatively *replicative phase* is characterized by the presence in the serum of HBeAg and HBV DNA levels well in excess of 10^3 – 10^4 IU/mL, sometimes exceeding 10^9 IU/mL; by the presence in the liver of detectable intrahepatocyte nucleocapsid antigens (primarily hepatitis B core antigen [HBeAg]); by high infectivity; and by accompanying liver injury. In contrast, the relatively *nonreplicative phase* is characterized by the absence of the conventional serum marker of HBV replication (HBeAg), the appearance of anti-HBe, levels of HBV DNA below a threshold of $\sim 10^3$ IU/mL, the absence of intrahepatocytic HBeAg, limited infectivity, and minimal liver injury. Patients in the replicative phase tend to have more severe chronic hepatitis, whereas those in the nonreplicative phase tend to have minimal or mild chronic hepatitis or to be inactive hepatitis B carriers. The likelihood in a patient with HBeAg-reactive chronic hepatitis B of converting spontaneously from relatively replicative to nonreplicative infection is approximately 10% per year. Distinctions in HBV replication and in histologic category, however, do not always coincide. In patients with HBeAg-reactive chronic HBV infection, especially when acquired at birth or in early childhood, as recognized commonly in Asian countries, a dichotomy is common between very high levels of HBV replication during the early decades of life (when the level of host tolerance of HBV is relatively high) and negligible levels of liver injury. Yet despite the relatively immediate, apparently benign nature of liver disease for many decades in this population, in the middle decades, activation of liver injury emerges as relative tolerance of the host to HBV declines, and these patients with childhood-acquired HBV infection are ultimately at increased risk later in life of cirrhosis, hepatocellular carcinoma (HCC) ([Chap. 111](#)), and liver-related death. **A discussion of the pathogenesis of liver injury in patients with chronic hepatitis B appears in [Chap. 360](#).**



HBeAg-negative chronic hepatitis B (i.e., chronic HBV infection with active virus replication, readily detectable HBV DNA but without HBeAg [anti-HBe-reactive]), is more common than HBeAg-reactive chronic hepatitis B in Mediterranean and European countries and in Asia (and, correspondingly, in HBV genotypes other than A). Compared to patients with HBeAg-reactive chronic hepatitis B, patients with HBeAg-negative chronic hepatitis B have levels of HBV DNA that are several orders of magnitude lower (no more than 10^5 – 10^6 IU/mL) than those observed in the HBeAg-reactive subset. Most such

cases represent precore or core-promoter mutations acquired late in the natural history of the disease (mostly early-life onset; age range 40–55 years, older than that for HBeAg-reactive chronic hepatitis B); these mutations prevent translation of HBeAg from the precore component of the HBV genome (precore mutants) or are characterized by downregulated transcription of precore mRNA (core-promoter mutants; [Chap. 360](#)). Although their levels of HBV DNA tend to be lower than among patients with HBeAg-reactive chronic hepatitis B, patients with HBeAg-negative chronic hepatitis B can have progressive liver injury (complicated by cirrhosis and HCC) and experience episodic reactivation of liver disease reflected in fluctuating levels of aminotransferase activity (“flares”). The biochemical and histologic activity of HBeAg-negative disease tends to correlate closely with levels of HBV replication, unlike the case mentioned above of Asian patients with HBeAg-reactive chronic hepatitis B during the early decades of their HBV infection. An important point worth reiterating is the observation that the level of HBV replication is the most important risk factor for the ultimate development of cirrhosis and HCC in both HBeAg-reactive and HBeAg-negative patients. Although levels of HBV DNA are lower and more

readily suppressed by therapy to undetectable levels in HBeAg-negative (compared to HBeAg-reactive) chronic hepatitis B, achieving sustained responses that permit discontinuation of antiviral therapy is less likely in HBeAg-negative patients (see below). Inactive carriers are patients with circulating hepatitis B surface antigen (HBsAg), normal serum aminotransferase levels, undetectable HBeAg, and levels of HBV DNA that are either undetectable or present at a threshold of $\leq 10^3$ IU/mL. This serologic profile can occur not only in inactive carriers but also in patients with HBeAg-negative chronic hepatitis B during periods of relative inactivity; distinguishing between the two requires sequential biochemical and virologic monitoring over many months.

The spectrum of *clinical features* of chronic hepatitis B is broad, ranging from asymptomatic infection to debilitating disease or even end-stage, fatal hepatic failure. As noted above, the onset of the disease tends to be insidious in most patients, with the exception of the very few in whom chronic disease follows failure of resolution of clinically apparent acute hepatitis B. **The clinical and laboratory features associated with progression from acute to chronic hepatitis B are discussed in Chap. 360.**

Fatigue is a common symptom, and persistent or intermittent *jaundice* is a common feature in severe or advanced cases. Intermittent deepening of jaundice and recurrence of malaise and anorexia, as well as worsening fatigue, are reminiscent of acute hepatitis; such exacerbations may occur spontaneously, often coinciding with evidence of virologic reactivation; may lead to progressive liver injury; and, when superimposed on well-established cirrhosis, may cause hepatic decompensation. Complications of cirrhosis occur in end-stage chronic hepatitis and include ascites, edema, bleeding gastroesophageal varices, hepatic encephalopathy, coagulopathy, or hypersplenism. Occasionally, these complications bring the patient to initial clinical attention. Extrahepatic complications of chronic hepatitis B, similar to those seen during the prodromal phase of acute hepatitis B, are associated with deposition of circulating hepatitis B antigen-antibody immune complexes. These include arthralgias and arthritis, which are common, and the more rare purpuric cutaneous lesions (leukocytoclastic vasculitis), immune-complex glomerulonephritis, and generalized vasculitis (polyarteritis nodosa) (**Chaps. 360 and 385**).

Laboratory features of chronic hepatitis B do not distinguish adequately between histologically mild and severe hepatitis. Aminotransferase elevations tend to be modest for chronic hepatitis B but may fluctuate in the range of 100–1000 units. As is true for acute viral hepatitis B, alanine aminotransferase (ALT) tends to be more elevated than aspartate aminotransferase (AST); however, once cirrhosis is established, AST tends to exceed ALT. Levels of alkaline phosphatase activity tend to be normal or only marginally elevated. In severe cases, moderate elevations in serum bilirubin (51.3–171 $\mu\text{mol/L}$ [3–10 mg/dL]) occur. Hypoalbuminemia and prolongation of the prothrombin time occur in severe or end-stage cases. Hyperglobulinemia and detectable circulating autoantibodies are distinctly absent in chronic hepatitis B (in contrast to autoimmune hepatitis). **Viral markers of chronic HBV infection are discussed in Chap. 360.**

TREATMENT CHRONIC HEPATITIS B

Although progression to cirrhosis is more likely in severe than in mild or moderate chronic hepatitis B, all forms of chronic hepatitis B can be progressive, and progression occurs primarily in patients with active HBV replication. Moreover, in populations of patients with chronic hepatitis B who are at risk for HCC (**Chap. 111**), the risk is highest for those with continued, high-level HBV replication and lower for persons in whom initially high-level HBV DNA falls spontaneously over time. Therefore, management of chronic hepatitis B is directed at suppressing the level of virus replication. Although clinical trials tend to focus on clinical endpoints achieved over 1–2 years (e.g., suppression of HBV DNA to undetectable levels, loss of HBeAg/HBsAg, improvement in histology, normalization of ALT), these short-term gains translate into reductions in the risk of clinical progression, hepatic decompensation, and death. To date, seven drugs have been approved for treatment of chronic hepatitis B:

injectable interferon (IFN) α ; pegylated interferon (long-acting IFN bound to polyethylene glycol, PEG [PEG IFN]); and the oral agents lamivudine, adefovir dipivoxil, entecavir, telbivudine, and tenofovir.

Antiviral therapy for hepatitis B has evolved rapidly since the mid-1990s, as has the sensitivity of tests for HBV DNA. When IFN and lamivudine were evaluated in clinical trials, HBV DNA was measured by insensitive hybridization assays with detection thresholds of 10^5 – 10^6 virions/mL; when adefovir, entecavir, telbivudine, tenofovir, and PEG IFN were studied in clinical trials, HBV DNA was measured by sensitive amplification assays (polymerase chain reaction [PCR]) with detection thresholds of 10^1 – 10^3 viral copies/mL or IU/mL. Recognition of these distinctions is helpful when comparing results of clinical trials that established the efficacy of these therapies (reviewed below in chronological order of publication of these efficacy trials).

INTERFERON

IFN- α was the first approved therapy for chronic hepatitis B. Although it is no longer used to treat hepatitis B, standard IFN is important historically, having provided important lessons about antiviral therapy in general. For immunocompetent adults with HBeAg-reactive chronic hepatitis B (who tend to have high-level HBV DNA [$>10^5$ – 10^6 virions/mL] and histologic evidence of chronic hepatitis on liver biopsy), a 16-week course of IFN given subcutaneously at a daily dose of 5 million units, or three times a week at a dose of 10 million units, results in a loss of HBeAg and hybridization-detectable HBV DNA (i.e., a reduction to levels below 10^5 – 10^6 virions/mL) in ~30% of patients, with a concomitant improvement in liver histology. Seroconversion from HBeAg to anti-HBe occurred in approximately 20%, and, in early trials, approximately 8% lost HBsAg. Successful IFN therapy and seroconversion are often accompanied by an acute hepatitis-like elevation in aminotransferase activity, which has been postulated to result from enhanced cytolytic T cell clearance of HBV-infected hepatocytes. Relapse after successful therapy is rare (1 or 2%). The likelihood of responding to IFN is higher in patients with lower levels of HBV DNA and substantial elevations of ALT. Although children can respond as well as adults, IFN therapy has not been effective in very young children infected at birth. Similarly, IFN therapy has not been effective in immunosuppressed persons, Asian patients with neonatal acquisition of infection and minimal-to-mild ALT elevations, or patients with decompensated chronic hepatitis B (in whom such therapy can actually be detrimental, sometimes precipitating decompensation, often associated with severe adverse effects). Among patients with HBeAg loss during therapy, long-term follow-up has demonstrated that 80% experience eventual loss of HBsAg (i.e., all serologic markers of infection, and normalization of ALT over a 9-year posttreatment period). In addition, improved long-term and complication-free survival as well as a reduction in the frequency of HCC have been documented among IFN responders, supporting the conclusion that successful antiviral therapy improves the natural history of chronic hepatitis B.

Initial trials of brief-duration IFN therapy in patients with *HBeAg-negative chronic hepatitis B* were disappointing, suppressing HBV replication transiently during therapy but almost never resulting in sustained antiviral responses. In subsequent IFN trials among patients with HBeAg-negative chronic hepatitis B, however, more protracted courses, lasting up to 1.5 years, have been reported to result in sustained remissions documented to last for several years, with suppressed HBV DNA and aminotransferase activity, in ~20%.

Complications of IFN therapy include systemic “flu-like” symptoms; marrow suppression; emotional lability (irritability, depression, anxiety); autoimmune reactions (especially autoimmune thyroiditis); and miscellaneous side effects such as alopecia, rashes, diarrhea, and numbness and tingling of the extremities. With the possible exception of autoimmune thyroiditis, all these side effects are reversible upon dose lowering or cessation of therapy.

Although no longer competitive with the newer generation of antivirals, IFN did represent the first successful antiviral approach and set a standard against which to measure subsequent drugs in the achievement of durable virologic, serologic, biochemical, and

2034 histologic responses; consolidation of virologic and biochemical benefit in the ensuing years after therapy; and improvement in the natural history of chronic hepatitis B. Standard IFN has been supplanted by long-acting PEG IFN (see below), and IFN nonresponders are now treated with one of the newer oral nucleoside analogues.

LAMIVUDINE

The first of the nucleoside analogues to be approved, the dideoxynucleoside lamivudine inhibits reverse transcriptase activity of both HIV and HBV and is a potent and effective agent for patients with chronic hepatitis B. Although generally superseded by newer, more potent agents, lamivudine is still used in regions of the world where newer agents are not yet approved or are not affordable. In clinical trials among patients with HBeAg-reactive chronic hepatitis B, lamivudine therapy at daily doses of 100 mg for 48–52 weeks suppressed HBV DNA by a median of approximately 5.5 log₁₀ copies/mL and to undetectable levels, as measured by PCR amplification assays, in approximately 40% of patients. Therapy was associated with HBeAg loss in 32–33%; HBeAg seroconversion (i.e., conversion from HBeAg-reactive to anti-HBe-reactive) in 16–21%; normalization of ALT in 40–75%; improvement in histology in 50–60%; retardation in fibrosis in 20–30%; and prevention of progression to cirrhosis. HBeAg responses can occur even in subgroups who are resistant to IFN (e.g., those with high-level HBV DNA) or who failed in the past to respond to it. As is true for IFN therapy of chronic hepatitis B, patients with near-normal ALT activity tend not to experience HBeAg responses (despite suppression of HBV DNA), and those with ALT levels exceeding 5 × the upper limit of normal can expect 1-year HBeAg seroconversion rates of 50–60%. Generally, HBeAg seroconversions are confined to patients who achieve suppression of HBV DNA to <10⁴ copies/mL (equivalent to ~10³ IU/mL). Lamivudine-associated HBeAg responses are accompanied by a posttreatment HBsAg seroconversion rate comparable to that seen after IFN-induced HBeAg responses. Among Western patients who undergo HBeAg responses during a year-long course of therapy and in whom the response is sustained for 4–6 months after cessation of therapy, the response is durable thereafter in the vast majority (>80%); therefore, the achievement of an HBeAg response represents a viable stopping point in therapy. Reduced durability has been reported in Asian patients; therefore, to support the durability of HBeAg responses, patients should receive a period of consolidation therapy of ≥6 months in Western patients and ≥1 year in Asian patients after HBeAg seroconversion. Close posttreatment monitoring is necessary to identify HBV reactivation promptly and to resume therapy. If HBeAg is unaffected by lamivudine therapy, the current approach is to continue therapy until an HBeAg response occurs, but long-term therapy may be required to suppress HBV replication and, in turn, limit liver injury; HBeAg seroconversions can increase to a level of 50% after 5 years of therapy. Histologic improvement continues to accrue with therapy beyond the first year; after a cumulative course of 3 years of lamivudine therapy, necroinflammatory activity is reduced in the majority of patients, and even cirrhosis has been shown to regress to precirrhotic stages in as many as three-quarters of patients.

Losses of HBsAg have been few during the first year of lamivudine therapy, and this observation had been cited as an advantage of IFN-based over lamivudine therapy; however, in head-to-head comparisons between standard IFN and lamivudine monotherapy, HBsAg losses were rare in both groups. Trials in which lamivudine and IFN were administered in combination failed to show a benefit of combination therapy over lamivudine monotherapy for either treatment-naïve patients or prior IFN nonresponders.

In patients with *HBeAg-negative chronic hepatitis B* (i.e., in those with precore and core-promoter HBV mutations), 1 year of lamivudine therapy results in HBV DNA suppression and normalization of ALT in three-quarters of patients and in histologic improvement in approximately two-thirds. Therapy has been shown to suppress HBV DNA by approximately 4.5 log₁₀ copies/mL (baseline HBV DNA levels are lower than in patients with HBeAg-reactive hepatitis B)

and to undetectable levels in approximately 70%, as measured by sensitive PCR amplification assays. Lacking HBeAg at the outset, patients with HBeAg-negative chronic hepatitis B cannot achieve an HBeAg response—a stopping point in HBeAg-reactive patients; almost invariably, when therapy is discontinued, reactivation is the rule. Therefore, these patients require long-term therapy; with successive years, the proportion with suppressed HBV DNA and normal ALT increases.

Clinical and laboratory side effects of lamivudine are negligible and indistinguishable from those observed in placebo recipients. Still, lamivudine doses should be reduced in patients with reduced creatinine clearance. During lamivudine therapy, transient ALT elevations, resembling those seen during IFN therapy and during spontaneous HBeAg-to-anti-HBe seroconversions, occur in one-fourth of patients. These ALT elevations may result from restored cytolytic T cell activation permitted by suppression of HBV replication. Similar ALT elevations, however, occur at an identical frequency in placebo recipients, but ALT elevations associated with HBeAg seroconversion are confined to lamivudine-treated patients. When therapy is stopped after a year of therapy, two- to threefold ALT elevations occur in 20–30% of lamivudine-treated patients, representing renewed liver-cell injury as HBV replication returns. Although these posttreatment flares are almost always transient and mild, rare severe exacerbations, especially in cirrhotic patients, have been observed, mandating close and careful clinical and virologic monitoring after discontinuation of treatment. Many authorities caution against discontinuing therapy in patients with cirrhosis, in whom posttreatment flares could precipitate decompensation.

Long-term monotherapy with lamivudine is associated with methionine-to-valine (M204V) or methionine-to-isoleucine (M204I) mutations, primarily at amino acid 204 in the tyrosine-methionine-aspartate-aspartate (YMDD) motif of HBV DNA polymerase, analogous to mutations that occur in HIV-infected patients treated with this drug. During a year of therapy, YMDD mutations occur in 15–30% of patients; the frequency increases with each year of therapy, reaching 70% at year 5. Ultimately, patients with YMDD mutants experience degradation of clinical, biochemical, and histologic responses; therefore, if treatment is begun with lamivudine monotherapy, the emergence of lamivudine resistance, reflected clinically by a breakthrough from suppressed levels of HBV DNA and ALT, is managed by adding another antiviral to which YMDD variants are sensitive (e.g., adefovir, tenofovir; see below).

Currently, although lamivudine is very safe and still used widely in other parts of the world, in the United States and Europe, lamivudine has been eclipsed by more potent antivirals that have superior resistance profiles (see below); it is no longer recommended as first-line therapy. Still, as the first successful oral antiviral agent for use in hepatitis B, lamivudine has provided proof of the concept that polymerase inhibitors can achieve virologic, serologic, biochemical, and histologic benefits. In addition, lamivudine has been shown to be effective in the treatment of patients with decompensated hepatitis B (for whom IFN is contraindicated), in some of whom decompensation can be reversed. Moreover, among patients with cirrhosis or advanced fibrosis, lamivudine has been shown to be effective in reducing the risk of progression to hepatic decompensation and, marginally, the risk of HCC. In the half decade following the introduction in the United States of lamivudine therapy for hepatitis B, referral of patients with HBV-associated end-stage liver disease for liver transplantation was reduced by ~30%, supporting further the beneficial impact of oral antiviral therapy on the natural history of chronic hepatitis B.

Because lamivudine monotherapy can result universally in the rapid emergence of YMDD variants in persons with HIV infection, patients with chronic hepatitis B should be tested for anti-HIV prior to therapy; if HIV infection is identified, lamivudine monotherapy at the HBV daily dose of 100 mg is contraindicated. These patients should be treated for both HIV and HBV with an HIV drug regimen that includes or is supplemented by at least two drugs active against HBV; antiretroviral therapy (ART) often contains two drugs

with antiviral activity against HBV (e.g., tenofovir and emtricitabine), but if lamivudine is part of the regimen, the daily dose should be 300 mg (Chap. 226). The safety of lamivudine during pregnancy has not been established; however, the drug is not teratogenic in rodents and has been used safely in pregnant women with HIV infection and with HBV infection. Limited data even suggest that administration of lamivudine during the last months of pregnancy to mothers with high-level hepatitis B viremia ($\geq 10^8$ IU/mL) can reduce the likelihood of perinatal transmission of hepatitis B.

ADEFOVIR DIPIVOXIL

At an oral daily dose of 10 mg, the acyclic nucleotide analogue adefovir dipivoxil, the prodrug of adefovir, reduces HBV DNA by approximately $3.5\text{--}4 \log_{10}$ copies/mL and is equally effective in treatment-naïve patients and IFN nonresponders. In HBeAg-reactive chronic hepatitis B, a 48-week course of adefovir dipivoxil was shown to achieve histologic improvement (and reduce the progression of fibrosis) and normalization of ALT in just over one-half of patients, HBeAg seroconversion in 12%, HBeAg loss in 23%, and suppression to an undetectable level of HBV DNA in 13–21%, as measured by PCR. Similar to IFN and lamivudine, adefovir dipivoxil is more likely to achieve an HBeAg response in patients with high baseline ALT (e.g., among adefovir-treated patients with ALT level $>5 \times$ the upper limit of normal), and HBeAg seroconversions occurred in 25%. The durability of adefovir-induced HBeAg responses is high (91% in one study); therefore, HBeAg response can be relied upon as a stopping point for adefovir therapy, after a period of consolidation therapy, as outlined above. Although data on the impact of additional therapy beyond 1 year are limited, biochemical, serologic, and virologic outcomes improve progressively as therapy is continued.

In patients with *HBeAg-negative chronic hepatitis B*, a 48-week course of 10 mg/d of adefovir dipivoxil resulted in histologic improvement in two-thirds, normalization of ALT in three-fourths, and suppression of HBV DNA to PCR-undetectable levels in one-half to two-thirds. As was true for lamivudine, because HBeAg responses—a potential stopping point—cannot be achieved in this group, reactivation is the rule when adefovir therapy is discontinued, and indefinite, long-term therapy is required. Treatment beyond the first year consolidates the gain of the first year; after 5 years of therapy, improvement in hepatic inflammation and regression of fibrosis were observed in three-fourths of patients, ALT was normal in 70%, and HBV DNA was undetectable in almost 70%. In one study, stopping adefovir after 5 years was followed by sustained suppression of HBV DNA and ALT, but most HBeAg-negative patients are treated indefinitely unless HBsAg loss, albeit very rare, is achieved.

Adefovir contains a flexible acyclic linker instead of the L-nucleoside ring of lamivudine, avoiding steric hindrance by mutated amino acids. In addition, the molecular structure of phosphorylated adefovir is very similar to that of its natural substrate; therefore, mutations to adefovir would also affect binding of the natural substrate, dATP. Hypothetically, these are among the reasons that resistance to adefovir dipivoxil is much less likely than resistance to lamivudine; no resistance was encountered in 1 year of clinical trial therapy. In subsequent years, however, adefovir resistance begins to emerge (asparagine to threonine at amino acid 236 [N236T] and alanine to valine or threonine at amino acid 181 [A181V/T], primarily), occurring in 2.5% after 2 years, but in 29% after 5 years of therapy (reported in HBeAg-negative patients). Among patients co-infected with HBV and HIV and who have normal CD4+ T cell counts, adefovir dipivoxil is effective in suppressing HBV dramatically (by $5 \log_{10}$ in one study). Moreover, adefovir dipivoxil is effective in lamivudine-resistant, YMDD-mutant HBV and can be used when such lamivudine-induced variants emerge. When lamivudine resistance occurs, adding adefovir (i.e., maintaining lamivudine to preempt the emergence of adefovir resistance) is superior to switching to adefovir. Almost invariably, patients with adefovir-mutant HBV respond to lamivudine (or newer agents, such as entecavir, see below). When, in the past, adefovir had been evaluated as therapy for HIV infection, doses of 60–120 mg were

required to suppress HIV, and, at these doses, the drug was nephrotoxic. Even at 30 mg/d, creatinine elevations of 44 $\mu\text{mol/L}$ (0.5 mg/dL) occurred in 10% of patients; however, at the HBV-effective dose of 10 mg, such elevations of creatinine are rarely encountered. If any nephrotoxicity does occur, it rarely appears before 6–8 months of therapy. Although renal tubular injury is a rare potential side effect, and although creatinine monitoring is recommended during treatment, the therapeutic index of adefovir dipivoxil is high, and the nephrotoxicity observed in clinical trials at higher doses was reversible. For patients with underlying renal disease, frequency of administration of adefovir dipivoxil should be reduced to every 48 h for creatinine clearances of 30–49 mL/min; to every 72 h for creatinine clearances of 10–29 mL/min; and once a week, following dialysis, for patients undergoing hemodialysis. Adefovir dipivoxil is very well tolerated, and ALT elevations during and after withdrawal of therapy are similar to those observed and described above in clinical trials of lamivudine. An advantage of adefovir is its relatively favorable resistance profile; however, it is not as potent as the other approved oral agents, it does not suppress HBV DNA as rapidly or as uniformly as the others, it is the least likely of all agents to result in HBeAg seroconversion, and 20–50% of patients fail to suppress HBV DNA by $2 \log_{10}$ (“primary nonresponders”). For these reasons, adefovir, which has been supplanted in both treatment-naïve and lamivudine-resistant patients by the more potent, less resistance-prone nucleotide analogue tenofovir (see below), is no longer recommended as first-line therapy.

PEGYLATED INTERFERON

After long-acting PEG IFN was shown to be effective in the treatment of hepatitis C (see below), this more convenient drug was evaluated in the treatment of chronic hepatitis B. Once-a-week PEG IFN is more effective than the more frequently administered, standard IFN, and several large-scale trials of PEG IFN versus oral nucleoside analogues have been conducted among patients with HBeAg-reactive and HBeAg-negative chronic hepatitis B.

In HBeAg-reactive chronic hepatitis B, two large-scale studies were done. One study evaluated PEG IFN- α 2b (100 μg weekly for 32 weeks, then 50 μg weekly for another 20 weeks for a total of 52 weeks, with a comparison arm of combination PEG IFN with oral lamivudine) in 307 subjects. The other study involved PEG IFN- α 2a (180 μg weekly for 48 weeks) in 814 primarily Asian patients, three-fourths of whom had ALT $\geq 2 \times$ the upper limit of normal, with comparison arms of lamivudine monotherapy and combination PEG IFN plus lamivudine. At the end of therapy (48–52 weeks) in the PEG IFN monotherapy arms, HBeAg loss occurred in approximately 30%, HBeAg seroconversion in 22–27%, undetectable HBV DNA (<400 copies/mL by PCR) in 10–25%, normal ALT in 34–39%, and a mean reduction in HBV DNA of $2 \log_{10}$ copies/mL (PEG IFN- α 2b) to $4.5 \log_{10}$ copies/mL (PEG IFN- α 2a). Six months after completing PEG IFN monotherapy in these trials, HBeAg losses were present in approximately 35%, HBeAg seroconversion in approximately 30%, undetectable HBV DNA in 7–14%, normal ALT in 32–41%, and a mean reduction in HBV DNA of $2\text{--}2.4 \log_{10}$ copies/mL. Although the combination of PEG IFN and lamivudine was superior at the end of therapy in one or more serologic, virologic, or biochemical outcomes, neither the combination arm (in both studies) nor the lamivudine monotherapy arm (in the PEG IFN- α 2a trial) demonstrated any benefit compared to the PEG IFN monotherapy arms 6 months after therapy. Moreover, HBsAg seroconversion occurred in 3–7% of PEG IFN recipients (with or without lamivudine); some of these seroconversions were identified by the end of therapy, but many were identified during the posttreatment follow-up period. The likelihood of HBeAg loss in PEG IFN-treated HBeAg-reactive patients is associated with HBV genotype A $>$ B $>$ C $>$ D (shown for PEG IFN- α 2b but not for α -2a).

Based on these results, some authorities concluded that PEG IFN monotherapy should be the first-line therapy of choice in HBeAg-reactive chronic hepatitis B; however, this conclusion has been challenged. Although a finite, 1-year course of PEG IFN results in a higher

rate of sustained response (6 months after treatment) than is achieved with oral nucleoside/nucleotide analogue therapy, the comparison is confounded by the fact that oral agents are not discontinued at the end of 1 year. Instead, taken orally and free of side effects, therapy with oral agents is extended indefinitely or until after the occurrence of an HBeAg response. The rate of HBeAg responses after 2 years of oral-agent nucleoside analogue therapy is at least as high as, if not higher than, that achieved with PEG IFN after 1 year; favoring oral agents is the absence of injections, difficult-to-tolerate side effects, and laboratory monitoring as well as lower direct and indirect medical care costs and inconvenience. The association of HBsAg responses with PEG IFN therapy occurs in such a small proportion of patients that subjecting everyone to PEG IFN for the marginal gain of HBsAg responses during or immediately after therapy in such a very small minority is questionable. Moreover, HBsAg responses occur in a comparable proportion of patients treated with early-generation nucleoside/nucleotide analogues in the years after therapy, and, with the newer, more potent nucleoside analogues, the frequency of HBsAg loss during the first year of therapy equals that of PEG IFN and is exceeded during year 2 and beyond (see below). Of course, resistance is not an issue during PEG IFN therapy, but the risk of resistance is much lower with new agents ($\leq 1\%$ up to 3–6 years in previously treatment-naïve, entecavir-treated and tenofovir-treated patients; see below). Finally, the level of HBV DNA inhibition that can be achieved with the newer agents, and even with lamivudine, exceeds that which can be achieved with PEG IFN, in some cases by several orders of magnitude.

In HBeAg-negative chronic hepatitis B, a trial of PEG IFN- α 2a (180 μg weekly for 48 weeks versus comparison arms of lamivudine monotherapy and of combination therapy) in 564 patients showed that PEG IFN monotherapy resulted at the end of therapy in suppression of HBV DNA by a mean of 4.1 \log_{10} copies/mL, undetectable HBV DNA (<400 copies/mL by PCR) in 63%, normal ALT in 38%, and loss of HBsAg in 4%. Although lamivudine monotherapy and combination lamivudine–PEG IFN therapy were both superior to PEG IFN at the end of therapy, no advantage of lamivudine monotherapy or combination therapy was apparent over PEG IFN monotherapy 6 months after therapy—suppression of HBV DNA by a mean of 2.3 \log_{10} copies/mL, undetectable HBV DNA in 19%, and normal ALT in 59%. In subjects involved in this trial followed for up to 5 years, among the two-thirds followed who had been treated initially with PEG IFN, 17% maintained HBV DNA suppression to <400 copies/mL, but ALT remained normal in only 22%; HBsAg loss increased gradually to 12%. Among the half followed who had been treated initially with lamivudine monotherapy, HBV DNA remained <400 copies/mL in 7% and ALT normal in 16%; by year 5, 3.5% had lost HBsAg. As was the case for standard IFN therapy in HBeAg-negative patients, only a small proportion maintained responsiveness after completion of PEG IFN therapy, raising questions about the relative value of a finite period of PEG IFN, versus a longer course with a potent, low-resistance oral nucleoside analogue in these patients. Moreover, the value of PEG IFN for HBeAg-negative chronic hepatitis B has not been confirmed. In the only other controlled clinical trial of PEG IFN for HBeAg-negative chronic hepatitis B, the hepatitis C regimen of PEG IFN plus ribavirin was compared to PEG IFN monotherapy. In this trial, HBV DNA suppression (<400 copies/mL) occurred in only 7.5% of the two groups combined, and no study subject lost HBsAg.

In patients treated with PEG IFN, HBeAg and HBsAg responses have been associated with *IL28B* genotype CC, the favorable genotype identified in trials of PEG IFN for chronic hepatitis C. Also, reductions in quantitative HBsAg levels have been shown to correlate with and to be predictive of responsiveness to PEG IFN in chronic hepatitis B. If HBsAg levels fail to fall within the first 12–24 weeks or to reach $<20,000$ IU/mL by week 24, PEG IFN therapy is unlikely to be effective and should be discontinued.

ENTECAVIR

Entecavir, an oral cyclopentyl guanosine analogue polymerase inhibitor, appears to be the most potent of the HBV antivirals and is just as well tolerated as lamivudine. In a 709-subject clinical trial among

HBeAg-reactive patients, oral entecavir, 0.5 mg daily, was compared to lamivudine, 100 mg daily. At 48 weeks, entecavir was superior to lamivudine in suppression of HBV DNA (mean 6.9 versus 5.5 \log_{10} copies/mL), percentage with undetectable HBV DNA (<300 copies/mL by PCR; 67% versus 36%), histologic improvement (≥ 2 -point improvement in necroinflammatory HAI score; 72% versus 62%), and normal ALT (68% versus 60%). The two treatments were indistinguishable in percentage with HBeAg loss (22% versus 20%) and seroconversion (21% versus 18%). Among patients treated with entecavir for 96 weeks, HBV DNA was undetectable cumulatively in 80% (versus 39% for lamivudine), and HBeAg seroconversions had occurred in 31% (versus 26% for lamivudine). After 3–6 years of entecavir, HBeAg seroconversions have been observed in 39–44% and HBsAg loss in 5–6%. Similarly, in a 638-subject clinical trial among HBeAg-negative patients, at week 48, oral entecavir, 0.5 mg daily, was superior to lamivudine, 100 mg daily, in suppression of HBV DNA (mean 5.0 versus 4.5 \log_{10} copies/mL) and in percentage with undetectable HBV DNA (90% versus 72%), histologic improvement (70% versus 61%), and normal ALT (78% versus 71%). No resistance mutations were encountered in previously treatment-naïve, entecavir-treated patients during 96 weeks of therapy, and in a cohort of subjects treated for up to 6 years, resistance emerged in only 1.2%. Entecavir-induced HBeAg seroconversions are as durable as those achieved with other antivirals. Its high barrier to resistance coupled with its high potency renders entecavir a first-line drug for patients with chronic hepatitis B.

Entecavir is also effective against lamivudine-resistant HBV infection. In a trial of 286 lamivudine-resistant patients, entecavir, at a higher daily dose of 1 mg, was superior to lamivudine, as measured at week 48, in achieving suppression of HBV DNA (mean 5.1 versus 0.48 \log_{10} copies/mL), undetectable HBV DNA (72% versus 19%), normal ALT (61% versus 15%), HBeAg loss (10% versus 3%), and HBeAg seroconversion (8% versus 3%). In this population of lamivudine-experienced patients, however, entecavir resistance emerged in 7% at 48 weeks. Although entecavir resistance requires both a YMDD mutation and a second mutation at one of several other sites (e.g., T184A, S202G/I, or M250V), resistance to entecavir in lamivudine-resistant chronic hepatitis B has been recorded to increase progressively to 43% at 4 years; therefore, entecavir is not as attractive a choice as adefovir or tenofovir for patients with lamivudine-resistant hepatitis B.

In clinical trials, entecavir has an excellent safety profile; in addition, on-treatment and posttreatment ALT flares are relatively uncommon and relatively mild in entecavir-treated patients. Doses should be reduced for patients with reduced creatinine clearance. Entecavir does have low-level antiviral activity against HIV and cannot be used as monotherapy to treat HBV infection in HIV/HBV co-infected persons.

TELBIVUDINE

Telbivudine, a cytosine analogue, is similar in efficacy to entecavir but slightly less potent in suppressing HBV DNA (a slightly less profound median 6.4 \log_{10} reduction in HBeAg-reactive disease and a similar 5.2 \log_{10} reduction in HBeAg-negative disease). In its registration trial, telbivudine at an oral daily dose of 600 mg suppressed HBV DNA to <300 copies/mL in 60% of HBeAg-positive and 88% of HBeAg-negative patients, reduced ALT to normal in 77% of HBeAg-positive and 74% of HBeAg-negative patients, and improved histology in 65% of HBeAg-positive and 67% of HBeAg-negative patients. Although resistance to telbivudine (M204I, not M204V, mutations) was less frequent than resistance to lamivudine at the end of 1 year, resistance mutations after 2 years of treatment occurred in up to 22%. Generally well tolerated, telbivudine has been associated with a low frequency of asymptomatic creatine kinase elevations and with a very low frequency of peripheral neuropathy; frequency of administration should be reduced for patients with impaired creatinine clearance. Its excellent potency notwithstanding, the inferior resistance profile of telbivudine has limited its appeal; telbivudine is neither recommended as first-line therapy nor widely used.

TENOFOVIR

Tenofovir disoproxil fumarate, an acyclic nucleotide analogue and potent antiretroviral agent used to treat HIV infection, is similar to adefovir but more potent in suppressing HBV DNA and inducing HBeAg responses; it is highly active against both wild-type and lamivudine-resistant HBV and active in patients whose response to adefovir is slow and/or limited. At an oral once-daily dose of 300 mg for 48 weeks, tenofovir suppressed HBV DNA by 6.2 log₁₀ (to undetectable levels [<400 copies/mL] in 76%) in HBeAg-positive patients and by 4.6 log₁₀ (to undetectable levels in 93%) in HBeAg-negative patients; reduced ALT to normal in 68% of HBeAg-positive and 76% of HBeAg-negative patients; and improved histology in 74% of HBeAg-positive and 72% of HBeAg-negative patients. In HBeAg-positive patients, HBeAg seroconversions occurred in 21% by the end of year 1, 27% by year 2, 34% by year 3, and 40% by year 5 of tenofovir treatment; HBsAg loss occurred in 3% by the end of year 1 and 6% at year 2, and 8% by year 5. After

5 years of tenofovir therapy, 87% of patients experienced histologic improvement, including reduction in fibrosis score (51%) and regression of cirrhosis (71%). The 5-year safety (negligible renal toxicity, in 1%, and mild reduction in bone density, in ~0.5%) and resistance profiles (none recorded through 5 years) of tenofovir are very favorable as well; therefore, tenofovir has supplanted adefovir both as first-line therapy for chronic hepatitis B and as add-on therapy for lamivudine-resistant chronic hepatitis B. Frequency of tenofovir administration should be reduced for patients with impaired creatinine clearance.

A comparison of the six antiviral therapies in current use appears in Table 362-3; their relative potencies in suppressing HBV DNA are shown in Fig. 362-1.

COMBINATION THERAPY

Although the combination of lamivudine and PEG IFN suppresses HBV DNA more profoundly during therapy than does monotherapy

TABLE 362-3 COMPARISON OF PEGYLATED INTERFERON (PEG IFN), LAMIVUDINE, ADEFOVIR, ENTECAVIR, TELBIVUDINE, AND TENOFOVIR THERAPY FOR CHRONIC HEPATITIS B^a

Feature	PEG IFN ^b	Lamivudine	Adefovir	Entecavir	Telbivudine	Tenofovir
Route of administration	Subcutaneous injection	Oral	Oral	Oral	Oral	Oral
Duration of therapy ^c	48–52 weeks	≥52 weeks	≥48 weeks	≥48 weeks	≥52 weeks	≥48 weeks
Tolerability	Poorly tolerated	Well tolerated	Well tolerated; creatinine monitoring recommended	Well tolerated	Well tolerated	Well tolerated; creatinine monitoring recommended
HBeAg seroconversion						
1 yr Rx	18–20%	16–21%	12%	21%	22%	21%
>1 yr Rx	NA	up to 50% @ 5 yrs	43% @ 3 yrs ^d	31% @ 2 yrs 44% @ 6 yrs	30% @ 2 yrs	40% @ 5 yrs
Log ₁₀ HBV DNA reduction (mean copies/mL)						
HBeAg-reactive	4.5	5.5	median 3.5–5	6.9	6.4	6.2
HBeAg-negative	4.1	4.4–4.7	median 3.5–3.9	5.0	5.2	4.6
HBV DNA PCR negative (<300–400 copies/mL; <1000 copies/mL for adefovir) at end of yr 1						
HBeAg-reactive	10–25%	36–44%	13–21%	67% (91% @ 4 yrs)	60%	76%
HBeAg-negative	63%	60–73%	48–77%	90%	88%	93%
ALT normalization at end of yr 1						
HBeAg-reactive	39%	41–75%	48–61%	68%	77%	68%
HBeAg-negative	34–38%	62–79%	48–77%	78%	74%	76%
HBsAg loss yr 1	3–4%	≤1%	0%	2%	<1%	3%
>yr 1	12% 5 yr after 1 yr of Rx	No data	5% at yr 5	6% at yr 6	No data	8% at yr 5
Histologic improvement (≥2 point reduction in HAI) at yr 1						
HBeAg-reactive	38% 6 months after	49–62%	53–68%	72%	65%	74%
HBeAg-negative	48% 6 months after	61–66%	64%	70%	67%	72%
Viral resistance	None	15–30% @ 1 yr 70% @ 5 yrs	None @ 1 yr 29% @ 5 yrs	≤1% @ 1 yr ^e 1.2% @ 6 yrs ^e	Up to 5% @ yr 1 Up to 22% @ yr 2	0% @ yr 1 0% through yr 5
Pregnancy category	C	C ^f	C	C	B	B
Cost (US\$) for 1 yr	~\$18,000	~\$2,500	~\$6,500	~\$8,700 ^g	~\$6,000	~\$6,000

^aGenerally, these comparisons are based on data on each drug tested individually versus placebo in registration clinical trials; because, with rare exception, these comparisons are not based on head-to-head testing of these drugs, relative advantages and disadvantages should be interpreted cautiously. ^bAlthough standard interferon α administered daily or three times a week is approved as therapy for chronic hepatitis B, it has been supplanted by PEG IFN, which is administered once a week and is more effective. Standard interferon has no advantages over PEG IFN. ^cDuration of therapy in clinical efficacy trials; use in clinical practice may vary. ^dBecause of a computer-generated randomization error that resulted in misallocation of drug versus placebo during the second year of clinical trial treatment, the frequency of HBeAg seroconversion beyond the first year is an estimate (Kaplan-Meier analysis) based on the small subset in whom adefovir was administered correctly. ^e7% during a year of therapy (43% at year 4) in lamivudine-resistant patients. ^fDespite its Class C designation, lamivudine has an extensive pregnancy safety record in women with HIV/AIDS. ^gApproximately \$17,400 for lamivudine-refractory patients.

Abbreviations: ALT, alanine aminotransferase; HAI, histologic activity index; HBeAg, hepatitis B e antigen; HBsAg, hepatitis B surface antigen; HBV, hepatitis B virus; NA, not applicable; PEG IFN, pegylated interferon; PCR, polymerase chain reaction; Rx, therapy; yr, year.

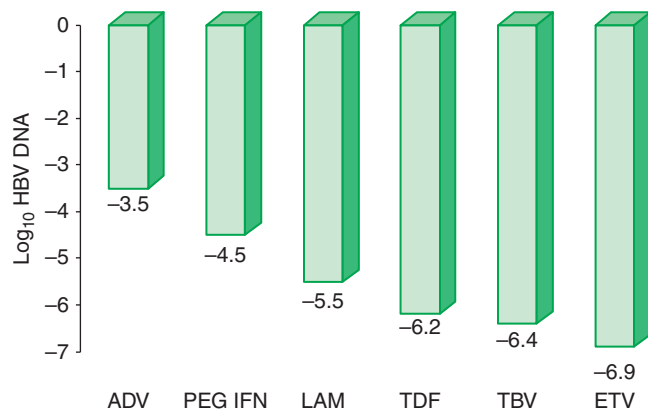


FIGURE 362-1 Relative potency of antiviral drugs for hepatitis B, as reflected by median \log_{10} HBV DNA reduction in HBeAg-positive chronic hepatitis B. These data are from individual reports of large, randomized controlled registration trials that were the basis for approval of the drugs. In most instances, these data do not represent direct comparisons among the drugs, because study populations were different, baseline patient variables were not always uniform, and the sensitivity and dynamic range of the HBV DNA assays used in the trials varied. ADV, adefovir dipivoxil; ETV, entecavir; LAM, lamivudine; PEG IFN, pegylated interferon α 2a; TBV, telbivudine; TDF, tenofovir disoproxil fumarate.

with either drug alone (and is much less likely to be associated with lamivudine resistance), this combination used for a year is no better than a year of PEG IFN in achieving sustained responses. To date, combinations of oral nucleoside/nucleotide agents have not achieved an enhancement in virologic, serologic, or biochemical efficacy over that achieved by the more potent of the combined drugs given individually. In a 2-year trial of combination entecavir and tenofovir versus entecavir monotherapy, for a small subgroup of patients with very high HBV DNA levels ($\geq 10^8$ IU/mL), a reduction in HBV DNA to < 50 IU/mL was higher in the combination group (79% versus 62%); however, no differences in HBeAg responses or any other endpoint were observed between the combination-therapy and monotherapy groups, even in the high-HBV DNA subgroup. On the other hand, combining agents that are not cross-resistant (e.g., lamivudine and adefovir or tenofovir) has the potential to reduce the risk or perhaps even to preempt entirely the emergence of drug resistance. In the future, the treatment paradigm may shift from the current approach of sequential monotherapy to preemptive combination therapy, perhaps not for all patients but for subsets (e.g., patients with very high levels of HBV DNA, immunosuppressed patients); however, designing and executing clinical trials that demonstrate superior efficacy and resistance profile of combination therapy over monotherapy with entecavir or tenofovir will remain challenging.

NOVEL ANTIVIRALS AND STRATEGIES

In addition to the seven approved antiviral drugs for hepatitis B, emtricitabine, a fluorinated cytosine analogue very similar to lamivudine in structure, efficacy, and resistance profile, offers no advantage over lamivudine. A combination of emtricitabine and tenofovir is approved for the treatment of HIV infection and is an appealing combination therapy for hepatitis B, especially for lamivudine-resistant disease; however, neither emtricitabine nor the combination is approved yet for hepatitis B. Several initially promising antiviral agents have been abandoned because of toxicity (e.g., clevudine, which was linked to myopathy during its clinical development). Because direct-acting antivirals have been so successful in the management of chronic hepatitis B, more unconventional approaches—e.g., immunologic (e.g., toll receptor agonists) or genetic manipulation (e.g., RNA interference—gene silencing—to reduce HBV DNA transcription)—are not likely to be competitive, unless they can be shown to go beyond current antivirals in achieving recovery (HBsAg

seroconversion) from HBV infection. Finally, initial emphasis in the development of antiviral therapy for hepatitis B was placed on monotherapy; whether combination regimens will yield additive or synergistic efficacy remains to be determined.

TREATMENT RECOMMENDATIONS

Several learned societies and groups of expert physicians have issued treatment recommendations for patients with chronic hepatitis B; the most authoritative and updated (and free of financial support by pharmaceutical companies) are those of the American Association for the Study of Liver Diseases (AASLD) and of the European Association for the Study of the Liver (EASL). Although the recommendations differ slightly, a consensus has emerged on most of the important points (Table 362-4). No treatment is recommended or available for inactive “nonreplicative” hepatitis B carriers (undetectable HBeAg with normal ALT and HBV DNA $\leq 10^3$ IU/mL documented serially over time). In patients with detectable HBeAg and HBV DNA levels $> 2 \times 10^4$ IU/mL, treatment is recommended by the AASLD for those with ALT levels above $2 \times$ the upper limit of normal. (The EASL recommends treatment in HBeAg-positive patients for HBV DNA levels $> 2 \times 10^3$ IU/mL and ALT above the upper limit of normal.) For HBeAg-positive patients with ALT $\leq 2 \times$ the upper limit of normal, in whom sustained responses are not likely and who would require multiyear therapy, antiviral therapy is not recommended currently. This pattern is common during the early decades of life among Asian patients infected at birth; even in this group, therapy would be considered for those > 40 years of age, ALT persistently at the high end of the twofold range, and/or with a family history of HCC, especially if the liver biopsy shows moderate to severe necroinflammatory activity or fibrosis. In this group, when, eventually, ALT becomes elevated later in life, antiviral therapy should be instituted. For patients with HBeAg-negative chronic hepatitis B, ALT $> 2 \times$ the upper limit of normal (above the upper limit of normal according to EASL), and HBV DNA $> 2 \times 10^3$ IU/mL, antiviral therapy is recommended. If HBV DNA is $> 2 \times 10^3$ IU/mL and ALT is 1 to $> 2 \times$ the upper limit of normal, liver biopsy should be considered to help in arriving at a decision to treat if substantial liver injury is present (treatment in this subset would be recommended according to EASL guidelines, because ALT is elevated).

For patients with compensated cirrhosis, because antiviral therapy has been shown to retard clinical progression, treatment is recommended regardless of HBeAg status and ALT as long as HBV DNA is detectable at $> 2 \times 10^3$ IU/mL (detectable at any level according to the EASL); monitoring without therapy is recommended for those with HBV DNA $< 2 \times 10^3$ IU/mL, unless ALT is elevated. For patients with decompensated cirrhosis, treatment is recommended regardless of serologic and biochemical status, as long as HBV DNA is detectable. Patients with decompensated cirrhosis should be evaluated as candidates for liver transplantation.

Among the seven available drugs for hepatitis B, PEG IFN has supplanted standard IFN, entecavir has supplanted lamivudine, and tenofovir has supplanted adefovir. PEG IFN, entecavir, or tenofovir is recommended as first-line therapy (Table 362-3). PEG IFN requires finite-duration therapy, achieves the highest rate of HBeAg responses after a year of therapy, and does not support viral mutations, but it requires subcutaneous injections and is associated with inconvenience, more intensive clinical and laboratory monitoring, and intolerance. Oral nucleoside analogues require long-term therapy in most patients, and when used alone, lamivudine and telbivudine foster the emergence of viral mutations, adefovir somewhat less so, and entecavir (except in lamivudine-experienced patients) and tenofovir rarely at all. Oral agents do not require injections or cumbersome laboratory monitoring, are very well tolerated, lead to improved histology in 50–90% of patients, suppress HBV DNA more profoundly than PEG IFN, and are effective even in patients who fail to respond to IFN-based therapy. Although oral agents are less likely to result in HBeAg responses during the first year of therapy, as compared to PEG IFN, treatment with oral agents tends to be extended beyond the first year and, by the end of the second year,

TABLE 362-4 RECOMMENDATIONS FOR TREATMENT OF CHRONIC HEPATITIS B^a

HBeAg status	Clinical	HBV DNA (IU/mL)	ALT	Recommendation
HBeAg-reactive	^b	>2 × 10 ⁴	≤2 × ULN ^{c,d}	No treatment; monitor. In patients >40, with family history of hepatocellular carcinoma, and/or ALT persistently at the high end of the twofold range, liver biopsy may help in decision to treat
	Chronic hepatitis	>2 × 10 ^{4d}	>2 × ULN ^d	Treat ^e
	Cirrhosis compensated	>2 × 10 ³	< or > ULN	Treat ^e with oral agents, not PEG IFN
	Cirrhosis decompensated	<2 × 10 ³	>ULN	Consider treatment ^f
HBeAg-negative	^b	≤2 × 10 ³	≤ULN	Inactive carrier; treatment not necessary
	Chronic hepatitis	>10 ³	1 to >2 × ULN ^d	Consider liver biopsy; treat ^h if biopsy shows moderate to severe inflammation or fibrosis
	Chronic hepatitis	>10 ⁴	>2 × ULN ^d	Treat ^{h,i}
	Cirrhosis compensated	>2 × 10 ³	< or > ULN	Treat ^e with oral agents, not PEG IFN
	Cirrhosis decompensated	<2 × 10 ³	>ULN	Consider treatment ^f
		Detectable	< or > ULN	Treat ^h with oral agents ^g , not PEG IFN; refer for liver transplantation
	Undetectable	< or > ULN	Observe; refer for liver transplantation	

^aBased on practice guidelines of the American Association for the Study of Liver Diseases (AASLD). Except as indicated in footnotes, these guidelines are similar to those issued by the European Association for the Study of the Liver (EASL). ^bLiver disease tends to be mild or inactive clinically; most such patients do not undergo liver biopsy. ^cThis pattern is common during early decades of life in Asian patients infected at birth. ^dAccording to the EASL guidelines, treat if HBV DNA is >2 × 10³ IU/mL and ALT >ULN. ^eOne of the potent oral drugs with a high barrier to resistance (entecavir or tenofovir) or PEG IFN can be used as first-line therapy (see text). These oral agents, but not PEG IFN, should be used for interferon-refractory/intolerant and immunocompromised patients. PEG IFN is administered weekly by subcutaneous injection for a year; the oral agents are administered daily for at least a year and continued indefinitely or until at least 6 months after HBeAg seroconversion. ^fAccording to EASL guidelines, patients with compensated cirrhosis and detectable HBV DNA at any level, even with normal ALT, are candidates for therapy. Most authorities would treat indefinitely, even in HBeAg-positive disease after HBeAg seroconversion. ^gBecause the emergence of resistance can lead to loss of antiviral benefit and further deterioration in decompensated cirrhosis, a low-resistance regimen is recommended—entecavir or tenofovir monotherapy or combination therapy with the more resistance-prone lamivudine (or telbivudine) plus adefovir. Therapy should be instituted urgently. ^hBecause HBeAg seroconversion is not an option, the goal of therapy is to suppress HBV DNA and maintain a normal ALT. PEG IFN is administered by subcutaneous injection weekly for a year; caution is warranted in relying on a 6-month posttreatment interval to define a sustained response, because the majority of such responses are lost thereafter. Oral agents, entecavir or tenofovir, are administered daily, usually indefinitely or until, as very rarely occurs, virologic and biochemical responses are accompanied by HBsAg seroconversion. ⁱFor older patients and those with advanced fibrosis, consider lowering the HBV DNA threshold to >2 × 10³ IU/mL.

Abbreviations: AASLD, American Association for the Study of Liver Diseases; ALT, alanine aminotransferase; EASL, European Association for the Study of the Liver; HBeAg, hepatitis B e antigen; HBsAg, hepatitis B surface antigen; HBV, hepatitis B virus; PEG IFN, pegylated interferon; ULN, upper limit of normal.

yields HBeAg responses (and even HBsAg responses) comparable in frequency to those achieved after 1 year of PEG IFN (and without the associated side effects) (Table 362-5). Although adefovir and tenofovir are safe, creatinine monitoring is recommended. Substantial experience with lamivudine during pregnancy (see above) has identified no teratogenicity. Although interferons do not appear to cause congenital anomalies, interferons have antiproliferative properties and should be avoided during pregnancy. Adefovir during pregnancy has not been associated with birth defects; however, there may be an increased risk of spontaneous abortion. Data on the safety of entecavir during pregnancy have not been published. Sufficient data in animals and limited data in humans suggest that telbivudine and tenofovir can be used safely during pregnancy. In general, except perhaps for lamivudine, and until additional data become available, the other antivirals for hepatitis B should be avoided or used with extreme caution during pregnancy.

As noted above, some physicians prefer to begin with PEG IFN, while other physicians and patients prefer oral agents as first-line therapy. For patients with decompensated cirrhosis, the emergence of resistance can result in further deterioration and loss of antiviral effectiveness. Therefore, in this patient subset, the threshold for relying on therapy with a very favorable resistance profile (e.g., entecavir or tenofovir) or on combination therapy is low. PEG IFN should not be used in patients with compensated or decompensated cirrhosis.

For patients with end-stage chronic hepatitis B who undergo liver transplantation, reinfection of the new liver is almost universal in the absence of antiviral therapy. The majority of patients become high-level viremic carriers with minimal liver injury. Before the availability of antiviral therapy, an unpredictable proportion experienced severe hepatitis B–related liver injury, sometimes a fulminant-like hepatitis and sometimes a rapid recapitulation of the original severe chronic hepatitis B (Chap. 360). Currently, however, prevention of

recurrent hepatitis B after liver transplantation has been achieved definitively by combining hepatitis B immune globulin with one of the oral nucleoside or nucleotide analogues (Chap. 368); preliminary data suggest that the newer, more potent, and less resistance-prone oral agents may be used instead of hepatitis B immune globulin for posttransplantation therapy.

Patients with HBV-HIV co-infection can have progressive HBV-associated liver disease and, occasionally, a severe exacerbation of hepatitis B resulting from immunologic reconstitution following ART. Lamivudine should never be used as monotherapy in patients with HBV-HIV infection because HIV resistance emerges rapidly to both viruses. Adefovir has been used successfully to treat chronic hepatitis B in HBV-HIV co-infected patients but is no longer considered a first-line agent for HBV. Entecavir has low-level activity against HIV and can result in selection of HIV resistance; therefore, it should be avoided in HBV-HIV co-infection. Tenofovir and the combination of tenofovir and emtricitabine in one pill are approved therapies for HIV and represent excellent choices for treating HBV infection in HBV-HIV co-infected patients. Generally, even for HBV-HIV co-infected patients who do not yet meet treatment criteria for HIV infection, treating for both HBV and HIV is recommended.

Patients with chronic hepatitis B who undergo cytotoxic chemotherapy for treatment of malignancies as well as patients treated with immunosuppressive, anticytokine, or antitumor necrosis factor therapies experience enhanced HBV replication and viral expression on hepatocyte membranes during chemotherapy coupled with suppression of cellular immunity. When chemotherapy is withdrawn, such patients are at risk for reactivation of hepatitis B, often severe and occasionally fatal. Such rebound reactivation represents restoration of cytolytic T cell function against a target organ enriched in HBV expression. Preemptive treatment with lamivudine prior to the initiation of chemotherapy has been shown to reduce the risk of such reactivation. The newer, more potent oral antiviral agents are

TABLE 362-5 PEGYLATED INTERFERON VERSUS ORAL NUCLEOSIDE ANALOGUES FOR THE TREATMENT OF CHRONIC HEPATITIS B

	PEG IFN	Nucleoside Analogues
Administration	Weekly injection	Daily, orally
Tolerability	Poorly tolerated, intensive monitoring	Well tolerated, limited monitoring
Duration of therapy	Finite 48 weeks	≥1 year, indefinite in most patients
Maximum mean HBV DNA suppression	4.5 log ₁₀	6.9 log ₁₀
Effective in high-level HBV DNA (≥10 ⁹ IU/mL)	No	Yes
HBeAg seroconversion		
During 1 year of therapy	~30%	~20%
During >1 year of therapy	Not applicable	30% (year 2) to up to 50% (year 5)
HBeAg-negative posttreatment HBV DNA suppression	17% @ 5 years	7% @ 4 years (lamivudine)
HBsAg loss		
During 1 year of therapy	3–4%	0–3%
During >1 year of therapy	Not applicable	3–8% @ 5 years of therapy
After 1 year of therapy–HBeAg-negative	12% @ 5 years	3.5% @ 5 years
Antiviral resistance	None	Lamivudine: ~30% year 1, ~70% year 5 Adefovir: 0% year 1, ~30% year 5 Telbivudine: up to 4% year 1, 22% year 2 Entecavir: ≤1.2% through year 6 Tenofovir: 0% through year 5
Use in cirrhosis, transplantation, immunosuppressed	No	Yes
Cost, 1 year of therapy	++++	+ to ++

Abbreviations: HBV, hepatitis B virus; HBeAg, hepatitis B e antigen; HBsAg, hepatitis B surface antigen; PEG IFN, pegylated interferon.

even more effective in preventing hepatitis B reactivation and with a lower risk of antiviral drug resistance. The optimal duration of antiviral therapy after completion of chemotherapy is not known, but a suggested approach is 6 months for inactive hepatitis B carriers and longer-duration therapy in patients with baseline HBV DNA levels >2 × 10³ IU/mL, until standard clinical endpoints are met (Table 362-4).

CHRONIC HEPATITIS D (DELTA HEPATITIS)

Chronic hepatitis D virus (HDV) may follow acute co-infection with HBV but at a rate no higher than the rate of chronicity of acute hepatitis B. That is, although HDV co-infection can increase the severity of acute hepatitis B, HDV does not increase the likelihood of progression to chronic hepatitis B. When, however, HDV superinfection occurs in a person who is already chronically infected with HBV, long-term HDV infection is the rule, and a worsening of the liver disease is the expected consequence. Except for severity, chronic hepatitis B plus D has similar clinical and laboratory features to those seen in chronic hepatitis B alone. Relatively severe and progressive chronic hepatitis, with or without cirrhosis, is the rule, and mild chronic hepatitis is the exception. Occasionally, however, mild hepatitis or even, rarely, inactive carriage occurs in patients with chronic hepatitis B plus D, and the disease may become indolent after several years of infection. A distinguishing serologic feature of chronic hepatitis D is the presence in the circulation of antibodies to liver-kidney microsomes (anti-LKM); however, the anti-LKM seen in hepatitis D, anti-LKM3, are directed against uridine diphosphate glucuronosyltransferase and are distinct from anti-LKM1 seen in patients with autoimmune hepatitis and in a subset of patients

with chronic hepatitis C (see below). **The clinical and laboratory features of chronic HDV infection are summarized in Chap. 360.**

TREATMENT CHRONIC HEPATITIS D

Management is not well defined. Glucocorticoids are ineffective and are not used. Preliminary experimental trials of IFN-α suggested that conventional doses and durations of therapy lower levels of HDV RNA and aminotransferase activity only transiently during treatment but have no impact on the natural history of the disease. In contrast, high-dose IFN-α (9 million units three times a week) for 12 months may be associated with a sustained loss of HDV replication and clinical improvement in up to 50% of patients. Moreover, the beneficial impact of treatment has been observed to persist for 15 years and to be associated with a reduction in grade of hepatic necrosis and inflammation, reversion of advanced fibrosis (improved stage), and clearance of HDV RNA in some patients. A suggested approach to therapy has been high-dose, long-term IFN for at least a year and, in responders, extension of therapy until HDV RNA and HBsAg clearance. PEG IFN has also been shown to be effective in the treatment of chronic hepatitis D (e.g., after 48 weeks of therapy, associated with undetectable HDV RNA, durable for at least 24 posttreatment weeks, in a quarter of patients) and is a more convenient replacement for standard IFN. None of the nucleoside analogue antiviral agents for hepatitis B are effective in hepatitis D. In patients with end-stage liver disease secondary to chronic hepatitis D, liver transplantation has been effective. If hepatitis D recurs in the new liver without the expression of hepatitis B (an unusual serologic profile in immunocompetent persons but common in transplant patients), liver injury is limited. In fact, the outcome of transplantation for chronic hepatitis D is superior to that for chronic hepatitis B; in such patients, combination hepatitis B immune globulin and nucleoside analogue therapy for hepatitis B is indicated (**Chap. 368**).

CHRONIC HEPATITIS C

Regardless of the epidemiologic mode of acquisition of hepatitis C virus (HCV) infection, chronic hepatitis follows acute hepatitis C in 50–70% of cases; chronic infection is common even in those with a return to normal in aminotransferase levels after acute hepatitis C, adding up to an 85% likelihood of chronic HCV infection after acute hepatitis C. Few clues had emerged to explain host differences associated with chronic infection until recently, when variation in a single nucleotide polymorphism (SNP) on chromosome 19, *IL28B* (which codes for IFN-λ3), was identified that distinguished between responders and nonresponders to antiviral therapy (see below). The same variants correlated with spontaneous resolution after acute infection: 53% in genotype C/C, 30% in genotype C/T, but only 23% in genotype T/T. The association with HCV clearance after acute infection is even stronger when *IL28B* haplotype is combined with haplotype G/G of an SNP near HLA class II *DBQ1*03:01*.

In patients with chronic hepatitis C followed for 20 years, progression to cirrhosis occurs in about 20–25%. Such is the case even for patients with relatively clinically mild chronic hepatitis, including those without symptoms, with only modest elevations of aminotransferase activity, and with mild chronic hepatitis on liver biopsy. Even in cohorts of well-compensated patients with chronic hepatitis C referred for clinical research trials (no complications of chronic liver disease and with normal hepatic synthetic function), the prevalence of cirrhosis may be as high as 50%. Most cases of hepatitis C are identified initially in asymptomatic patients who have no history of acute hepatitis C (e.g., those discovered while attempting to donate blood, while undergoing lab testing as part of an application for life insurance, or as a result of routine laboratory tests). The source of HCV infection in many of these cases is not defined, although a long-forgotten percutaneous exposure (e.g., injection drug use) in the remote past can be elicited in a substantial proportion and probably accounts for most infections; most of these infections were acquired in the 1960s and 1970s, coming to clinical attention decades later.

Approximately one-third of patients with chronic hepatitis C have normal or near-normal aminotransferase activity; although one-third to one-half of these patients have chronic hepatitis on liver biopsy, the grade of liver injury and stage of fibrosis tend to be mild in the vast majority. In some cases, more severe liver injury has been reported—even, rarely, cirrhosis, most likely the result of previous histologic activity. Among patients with persistent normal aminotransferase activity sustained over ≥ 5 –10 years, histologic progression has been shown to be rare; however, approximately one-fourth of patients with normal aminotransferase activity experience subsequent aminotransferase elevations, and histologic injury can be progressive once abnormal biochemical activity resumes. Therefore, continued clinical monitoring is indicated, even for patients with normal aminotransferase activity.

Despite this substantial rate of progression of chronic hepatitis C, and despite the fact that liver failure can result from end-stage chronic hepatitis C, the long-term prognosis for chronic hepatitis C in a majority of patients is relatively benign. Mortality over 10–20 years among patients with transfusion-associated chronic hepatitis C has been shown not to differ from mortality in a matched population of transfused patients in whom hepatitis C did not develop. Although death in the hepatitis group is more likely to result from liver failure, and although hepatic decompensation may occur in $\sim 15\%$ of such patients over the course of a decade, the majority (almost 60%) of patients remain asymptomatic and well compensated, with no clinical sequelae of chronic liver disease. Overall, chronic hepatitis C tends to be very slowly and insidiously progressive, if at all, in the vast majority of patients, whereas in approximately one-fourth of cases, chronic hepatitis C will progress eventually to end-stage cirrhosis. In fact, because HCV infection is so prevalent, and because a proportion of patients progress inexorably to end-stage liver disease, hepatitis C is the most frequent indication for liver transplantation (**Chap. 368**). In the United States, hepatitis C accounts for up to 40% of all chronic liver disease, and, as of 2007, mortality caused by hepatitis C surpassed that associated with HIV/AIDS. Moreover, because the prevalence of HCV infection is so much higher in the “baby boomer” cohort borne between 1945 and 1965, three-quarters of the mortality associated with hepatitis C occurs in this age cohort. Referral bias may account for the more severe outcomes described in cohorts of patients reported from tertiary care centers (20-year progression of $\geq 20\%$) versus the more benign outcomes in cohorts of patients monitored from initial blood-product-associated acute hepatitis or identified in community settings (20-year progression of only 4–7%). Still unexplained, however, are the wide ranges in reported progression to cirrhosis, from 2% over 17 years in a population of women with hepatitis C infection acquired from contaminated anti-D immune globulin to 30% over ≤ 11 years in recipients of contaminated intravenous immune globulin.

Progression of liver disease in patients with chronic hepatitis C has been reported to be more likely in patients with older age, longer duration of infection, advanced histologic stage and grade, more complex quasispecies diversity, increased hepatic iron, concomitant other liver disorders (alcoholic liver disease, chronic hepatitis B, hemochromatosis, α_1 antitrypsin deficiency, and steatohepatitis), HIV infection, and obesity. Among these variables, however, duration of infection appears to be one of the most important, and some of the others probably reflect disease duration to some extent (e.g., quasispecies diversity, hepatic iron accumulation). No other epidemiologic or clinical features of chronic hepatitis C (e.g., severity of acute hepatitis, level of aminotransferase activity, level of HCV RNA, presence or absence of jaundice during acute hepatitis) are predictive of eventual outcome. Despite the relatively benign nature of chronic hepatitis C over time in many patients, cirrhosis following chronic hepatitis C has been associated with the late development, after several decades, of HCC (**Chap. 111**); the annual rate of HCC in cirrhotic patients with hepatitis C is 1–4%, occurring primarily in patients who have had HCV infection for 30 years or more.

Perhaps the best prognostic indicator in chronic hepatitis C is liver histology; the rate of hepatic fibrosis may be slow, moderate, or rapid. Patients with mild necrosis and inflammation as well as those with limited fibrosis have an excellent prognosis and limited progression to cirrhosis. In contrast, among patients with moderate to severe

necroinflammatory activity or fibrosis, including septal or bridging fibrosis, progression to cirrhosis is highly likely over the course of 10–20 years. The pace of fibrosis progression may be accelerated by such factors as concomitant HIV infection, other causes of liver disease, excessive alcohol use, and hepatic steatosis. Among patients with compensated cirrhosis associated with hepatitis C, the 10-year survival rate is close to 80%; mortality occurs at a rate of 2–6% per year; decompensation at a rate of 4–5% per year; and, as noted above, HCC at a rate of 1–4% per year. **A discussion of the pathogenesis of liver injury in patients with chronic hepatitis C appears in Chap. 360.**

Clinical features of chronic hepatitis C are similar to those described above for chronic hepatitis B. Generally, fatigue is the most common symptom; jaundice is rare. Immune complex–mediated extrahepatic complications of chronic hepatitis C are less common than in chronic hepatitis B (despite the fact that assays for immune complexes are often positive in patients with chronic hepatitis C), with the exception of essential mixed cryoglobulinemia (**Chap. 360**), which is linked to cutaneous vasculitis and membranoproliferative glomerulonephritis as well as lymphoproliferative disorders such as B-cell lymphoma and unexplained monoclonal gammopathy. In addition, chronic hepatitis C has been associated with extrahepatic complications unrelated to immune-complex injury. These include Sjögren’s syndrome, lichen planus, porphyria cutanea tarda, type 2 diabetes mellitus, and the metabolic syndrome (including insulin resistance and steatohepatitis).

Laboratory features of chronic hepatitis C are similar to those in patients with chronic hepatitis B, but aminotransferase levels tend to fluctuate more (the characteristic episodic pattern of aminotransferase activity) and to be lower, especially in patients with long-standing disease. An interesting and occasionally confusing finding in patients with chronic hepatitis C is the presence of autoantibodies. Rarely, patients with autoimmune hepatitis (see below) and hyperglobulinemia have false-positive immunoassays for anti-HCV. On the other hand, some patients with serologically confirmable chronic hepatitis C have circulating anti-LKM. These antibodies are anti-LKM1, as seen in patients with autoimmune hepatitis type 2 (see below), and are directed against a 33-amino-acid sequence of cytochrome P450 IID6. The occurrence of anti-LKM1 in some patients with chronic hepatitis C may result from the partial sequence homology between the epitope recognized by anti-LKM1 and two segments of the HCV polyprotein. In addition, the presence of this autoantibody in some patients with chronic hepatitis C suggests that autoimmunity may be playing a role in the pathogenesis of chronic hepatitis C.

Histopathologic features of chronic hepatitis C, especially those that distinguish hepatitis C from hepatitis B, are described in Chap. 360.

TREATMENT CHRONIC HEPATITIS C

Therapy for chronic hepatitis C has evolved substantially in the two decades since IFN- α was introduced for this indication. The therapeutic armamentarium has grown to include PEG IFN with ribavirin and, in 2011, the introduction of protease inhibitors telaprevir and boceprevir used in combination with PEG IFN and ribavirin in patients with HCV genotype 1. When first approved, IFN- α was administered via subcutaneous injection three times a week for 6 months but achieved a sustained virologic response (SVR) (**Fig. 362-2**) (a reduction of HCV RNA to undetectable levels by PCR when measured ≥ 6 months after completion of therapy) below 10%. Doubling the duration of therapy—but not increasing the dose or changing IFN preparations—increased the SVR rate to $\sim 20\%$, and addition to the regimen of daily ribavirin, an oral guanosine nucleoside, increased the SVR rate to 40%. When used alone, ribavirin is ineffective and does not reduce HCV RNA levels appreciably, but ribavirin enhances the efficacy of IFN by reducing the likelihood of virologic relapse after the achievement of an end-treatment response (**Fig. 362-2**) (response measured during, and maintained to the end of, treatment). Proposed mechanisms to explain the role of ribavirin include subtle direct reduction of HCV replication, inhibition of host inosine monophosphate dehydrogenase activity (and associated depletion

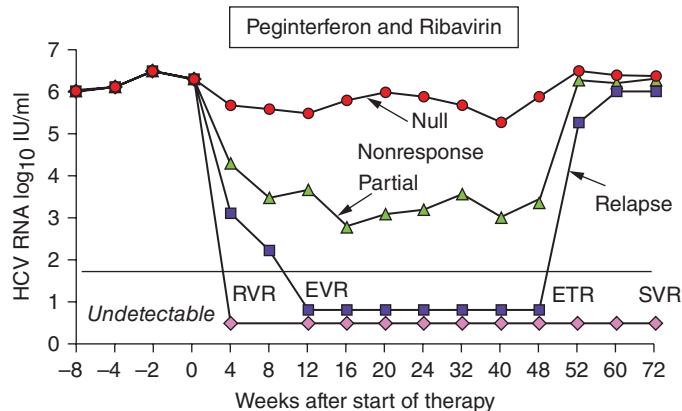


FIGURE 362-2 Classification of virologic responses based on outcomes during and after a 48-week course of pegylated interferon (PEG IFN) plus ribavirin antiviral therapy in patients with hepatitis C, genotype 1 or 4 (for genotype 2 or 3, the course would be 24 weeks). Nonresponders can be classified as null responders (hepatitis C virus [HCV] RNA reduction of $<2 \log_{10}$ IU/mL) or partial responders (HCV RNA reduction $\geq 2 \log_{10}$ IU/mL but not suppressed to undetectable) by week 24 of the therapy. In responders, HCV RNA can become undetectable, as shown with sensitive amplification assays, within 4 weeks (RVR, rapid virologic response); can be reduced by $\geq 2 \log_{10}$ IU/mL within 12 weeks (early virologic response, EVR; if HCV RNA is undetectable at 12 weeks, the designation is “complete” EVR); or at the end of therapy, 48 weeks (ETR, end-treatment response). In responders, if HCV RNA remains undetectable for 24 weeks after ETR, week 72, the patient has a sustained virologic response (SVR), but if HCV RNA becomes detectable again, the patient is considered to have relapsed. In patients treated with protease inhibitor–based therapy, several additional milestones are monitored: (1) among boceprevir-treated patients, the level of HCV RNA reduction ($>1 \log_{10}$ or $\geq 1 \log_{10}$ IU/mL) during the 4-week PEG IFN–ribavirin lead-in phase; (2) during boceprevir therapy, undetectable HCV RNA at week 8 (week 4 of triple-drug therapy; RVR); and (3) among telaprevir-treated patients, undetectable HCV RNA at week 4 and 12 (extended RVR). (Reproduced with permission, courtesy of Marc G. Ghany, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health and the American Association for the Study of Liver Diseases. *Hepatology* 49:1335, 2009.)

of guanosine pools), immune modulation, induction of virologic mutational catastrophe, and enhancement of IFN-stimulated gene expression. IFN therapy results in activation of the JAK-STAT signal transduction pathway, which culminates in the intracellular elaboration of genes and their protein products that have antiviral properties. Hepatitis C proteins inhibit JAK-STAT signaling at several steps along the pathway, and exogenous IFN restores expression of IFN-stimulated genes and their antiviral effects.

Treatment with the combination of PEG IFN and ribavirin increased responsiveness (frequency of SVR) to as high as 55% overall, to $>40\%$ in genotypes 1 and 4, and to $>80\%$ in genotypes 2 and 3. Still, many important lessons about antiviral therapy for chronic hepatitis C were learned from the experience with IFN monotherapy and combination IFN-ribavirin therapy. Even in the absence of biochemical and virologic responses, histologic improvement occurs in approximately three-fourths of all treated patients. In chronic hepatitis C, unlike the case in hepatitis B, responses to therapy are not accompanied by transient, acute hepatitis-like aminotransferase elevations. Instead, ALT levels fall precipitously during therapy. Up to 90% of virologic responses are achieved within the first 12 weeks of therapy; responses thereafter are rare. Most relapses occur within the first 12 weeks after treatment; therefore, an SVR at week 12 posttreatment is roughly equivalent to a 24-week SVR. SVRs are very durable; normal ALT, improved histology, and absence of HCV RNA in serum and liver have been documented a decade after successful

therapy, and “relapses” 2 years after sustained responses are almost unheard of. Thus, an SVR to antiviral therapy of chronic hepatitis C is tantamount to a cure.

Patient variables that tend to correlate with sustained virologic responsiveness to IFN-based therapy include favorable genotype (genotypes 2 and 3 as opposed to genotypes 1 and 4); low baseline HCV RNA level (<2 million copies/mL, which is equivalent to $<800,000$ IU/mL, the current convention of quantitation); histologically mild hepatitis and minimal fibrosis; age <40 ; female gender; and absence of obesity, insulin resistance, and type 2 diabetes mellitus. Patients with cirrhosis can respond, but they are less likely to do so. For patients treated with combination IFN-ribavirin, therapy for those with genotype 1 should last a full 48 weeks, whereas in those with genotypes 2 and 3, a 24-week course of therapy suffices (although refined tailoring of treatment duration may be indicated based on rapidity of response or associated cofactors, see below). The response rate in African Americans is disappointingly low for reasons that are not fully understood. Potentially contributing to, but not explaining entirely, low responsiveness in African Americans are a higher proportion with genotype 1, slower early viral kinetics during therapy, impaired HCV-specific immunity, and recently recognized host genetic differences in *IL28B* alleles, described below. The response rate in Latino patients is also low, despite the fact that the frequency of the favorable *IL28B* C allele is as common in Hispanic patients as in whites. Moreover, the likelihood of a sustained response is best if adherence to the treatment regimen is high (i.e., if patients receive $\geq 80\%$ of the IFN and ribavirin doses and if they continue treatment for $\geq 80\%$ of the anticipated duration of therapy). Other variables reported to correlate with increased responsiveness include brief duration of infection, low HCV quasispecies diversity, immunocompetence, absence of hepatic steatosis, and low liver iron levels. High levels of HCV RNA, more histologically advanced liver disease, and high quasispecies diversity all go hand in hand with advanced duration of infection, which is one of the most important clinical variables determining IFN responsiveness. The ironic fact, then, is that patients whose disease is least likely to progress are the ones *most* likely to respond to IFN and vice versa.

Genetic changes in the virus may explain differences in treatment responsiveness in some patients (e.g., among patients with genotype 1b, responsiveness to IFN is enhanced in those with amino-acid-substitution mutations in the nonstructural protein 5A gene). As described above in the discussion of spontaneous recovery from acute hepatitis C, IFN gene variants discovered recently in genome-wide association studies have been shown to have a substantial impact on responsiveness of patients with genotype 1 to antiviral therapy. In studies of patients treated with PEG IFN and ribavirin, variants of the *IL28B* SNP that code for IFN- $\lambda 3$ (a type III IFN, the receptors for which are more discretely distributed than IFN- α receptors and more concentrated in hepatocytes) correlate significantly with responsiveness. Patients homozygous for the C allele at this locus have the highest frequency of achieving an SVR ($\sim 80\%$), those homozygous for the T allele at this locus are least likely to achieve an SVR ($\sim 25\%$), and those heterozygous at this locus (C/T) have an intermediate level of responsiveness (SVRs in $\sim 35\%$). The fact that C/C is common in whites of European ancestry and even more so in Japanese persons but rare in African Americans helps explain the differences in observed responsiveness among these population groups.

Side effects of IFN therapy are described above in the section on treatment of chronic hepatitis B. The most pronounced side effect of ribavirin therapy is hemolysis; a reduction in hemoglobin of up to 2–3 g or in hematocrit of 5–10% can be anticipated. A small, unpredictable proportion of patients experience profound, brisk hemolysis, resulting in symptomatic anemia; therefore, close monitoring of blood counts is crucial, and ribavirin should be avoided in patients with anemia or hemoglobinopathies and in patients with coronary artery disease or cerebrovascular disease, in whom anemia can precipitate an ischemic event. When symptomatic anemia occurs,

ribavirin dose reductions or addition of erythropoietin to boost red blood cell levels may be required; erythropoietin has been shown to improve patients' quality of life but not the likelihood of achieving an SVR. If ribavirin is stopped during therapy, SVR rates fall, but responsiveness can be maintained as long as ribavirin is not stopped and the total ribavirin dose exceeds 60% of the planned dose. In addition, ribavirin, which is excreted renally, should not be used in patients with renal insufficiency; the drug is teratogenic, precluding its use during pregnancy and mandating the scrupulous use of efficient contraception during therapy (IFNs, too, because of their antiproliferative properties, are contraindicated during pregnancy).

Ribavirin can also cause nasal and chest congestion, pruritus, and precipitation of gout. Combination IFN-ribavirin therapy is more difficult to tolerate than IFN monotherapy. In one large clinical trial of combination therapy versus monotherapy, among those in the 1-year treatment group, 21% of the combination group (but only 14% of the monotherapy group) had to discontinue treatment, whereas 26% of the combination group (but only 9% of the monotherapy group) required dose reductions.

Studies of viral kinetics have shown that despite a virion half-life in serum of only 2–3 h, the level of HCV is maintained by a high replication rate of 10^{12} hepatitis C virions per day. IFN- α blocks virion production or release with an efficacy that increases with increasing drug doses; moreover, the calculated death rate for infected cells during IFN therapy is inversely related to level of HCV RNA; patients with the most rapid death rate of infected hepatocytes are more likely to achieve undetectable HCV RNA at 3 months; in practice, failure to achieve an early virologic response (EVR), a $\geq 2\text{-log}_{10}$ reduction in HCV RNA by week 12, predicts failure to experience a subsequent SVR. Similarly, patients in whom HCV RNA becomes undetectable within 4 weeks (i.e., who achieve a rapid virologic response [RVR]) have a very high likelihood of achieving an SVR (Fig. 362-2). Therefore, to achieve rapid viral clearance from serum and the liver, *high-dose induction therapy* has been advocated. In practice, however, high-dose induction with IFN-based therapy has not yielded higher sustained response rates.

For the treatment of chronic hepatitis C, standard IFNs were supplanted by PEG IFNs. These have elimination times up to sevenfold longer than standard IFNs (i.e., a substantially longer half-life) and achieve prolonged concentrations, permitting administration once (rather than three times) a week. Instead of the frequent drug peaks (linked to side effects) and troughs (when drug is absent) associated with frequent administration of short-acting IFNs, administration of PEG IFNs results in drug concentrations that are more stable and sustained over time. Once-a-week PEG IFN monotherapy is twice as effective as monotherapy with its standard IFN counterpart, approaches the efficacy of combination standard IFN plus ribavirin, and is as well tolerated as standard IFNs, without more difficult-to-manage thrombocytopenia and leukopenia than standard IFNs. For most of the decade prior to 2011, when protease inhibitors were introduced for HCV genotype 1 (see below), the standard of care was a combination of PEG IFN plus ribavirin for all HCV genotypes.

Two PEG IFNs are available: PEG IFN- $\alpha 2b$ and - $\alpha 2a$. PEG IFN- $\alpha 2b$ consists of a 12-kD, linear PEG molecule bound to IFN- $\alpha 2b$, whereas PEG IFN- $\alpha 2a$ consists of a larger, 40-kD, branched PEG molecule bound to IFN- $\alpha 2a$; because of its larger size and smaller volume of extravascular distribution, PEG IFN- $\alpha 2a$ can be given at a uniform dose independent of weight, whereas the dose of the smaller PEG IFN- $\alpha 2b$, which has a much wider volume distribution, must be weight-based (Table 362-6). In the registration trial for PEG IFN- $\alpha 2b$ plus ribavirin, the best regimen was 48 weeks of 1.5 $\mu\text{g}/\text{kg}$ of PEG IFN once a week plus 800 mg of ribavirin daily. A post hoc analysis suggested that weight-based dosing of ribavirin would have been more effective than the fixed 800-mg dose used in the study (a broader dose/weight range was approved subsequently; see below). In the first registration trial for PEG IFN- $\alpha 2a$ plus ribavirin, the best regimen was 48 weeks of 180 μg of PEG IFN plus 1000 mg (for patients <75 kg) to 1200 mg (for patients ≥ 75 kg) of ribavirin. SVRs of 54 and 56% were reported in these two studies, respectively. A subsequent

TABLE 362-6 PEGYLATED INTERFERON (PEG IFN) $\alpha 2a$ AND $\alpha 2b$ FOR CHRONIC HEPATITIS C

	PEG IFN- $\alpha 2b$	PEG IFN- $\alpha 2a$
PEG size	12 kD linear	40 kD branched
Elimination half-life	54 h	65 h
Clearance	725 mL/h	60 mL/h
Dose	1.5 $\mu\text{g}/\text{kg}$ (weight-based)	180 μg
Storage	Room temperature	Refrigerated
Ribavirin dose		
Genotype 1	800–1400 mg ^a	1000–1200 mg ^b
Genotype 2/3	800 mg	800 mg
Duration of therapy		
Genotype 1	48 weeks	48 weeks
Genotype 2/3	48 weeks ^c	24 weeks
Efficacy of combination therapy ^d	54%	56%
Genotype 1	40–42%	41–51%
Genotype 2/3	82%	76–78%

^aIn the registration trial for PEG IFN- $\alpha 2b$ plus ribavirin, the optimal regimen was 1.5 μg of PEG IFN plus 800 mg of ribavirin; however, a post hoc analysis of this study suggested that higher ribavirin doses are better. In subsequent trials of PEG IFN- $\alpha 2b$ with ribavirin in patients with genotype 1, the following daily ribavirin doses have been validated: 800 mg for patients weighing <65 kg, 1000 mg for patients weighing >65–85 kg, 1200 mg for patients weighing >85–105 kg, and 1400 mg for patients weighing >105 kg. ^b1000 mg for patients weighing <75 kg; 1200 mg for patients weighing ≥ 75 kg. ^cIn the registration trial for PEG IFN- $\alpha 2b$ plus ribavirin, all patients were treated for 48 weeks; however, data from other trials of standard interferons and the other PEG IFN demonstrated that 24 weeks suffices for patients with genotypes 2 and 3. For patients with genotype 3 who have advanced fibrosis/cirrhosis and/or high-level HCV RNA, a full 48 weeks is preferable. ^dAttempts to compare the two PEG IFN preparations based on the results of registration clinical trials are confounded by differences between trials of the two agents in methodologic details (different ribavirin doses, different methods for recording depression, and other side effects) and study-population composition (different proportion with bridging fibrosis/cirrhosis, proportion from the United States versus international, mean weight, proportion with genotype 1, and proportion with high-level HCV RNA). In the head-to-head comparison of the two PEG IFN preparations in the IDEAL trial reported in 2009, the two drugs were comparable in tolerability and efficacy. PEG IFN- $\alpha 2b$ was administered at a weekly weight-based dose of 1.0 $\mu\text{g}/\text{kg}$ or 1.5 $\mu\text{g}/\text{kg}$, and PEG IFN- $\alpha 2a$ at a weekly fixed dose of 180 μg . For PEG IFN- $\alpha 2b$, daily ribavirin weight-based doses ranged between 800 and 1400 mg based on weight criteria (see footnote a, above), whereas for PEG IFN- $\alpha 2a$, daily ribavirin weight-based doses ranged between 1000 and 1200 mg (see footnote b, above). For the two PEG IFN- $\alpha 2b$ study arms, ribavirin dose reductions for ribavirin-associated adverse effects were done in 200- to 400-mg decrements; for PEG IFN- $\alpha 2a$, the ribavirin dose was reduced to 600 mg for intolerability. Sustained virologic responses occurred in 38.0% of the low-dose PEG IFN- $\alpha 2b$ group, 39.8% of the standard, full-dose PEG IFN- $\alpha 2b$ group, and 40.9% of the PEG IFN- $\alpha 2a$ group.

Abbreviations: HCV RNA, hepatitis C virus RNA; PEG, polyethylene glycol.

study of PEG IFN- $\alpha 2a$ plus ribavirin showed that, for patients with genotypes 2 and 3, a duration of 24 weeks and a ribavirin dose of 800 mg were sufficient. Among the three studies, for patients in the optimal treatment arm, SVR rates for patients with genotype 1 were 42–51%, and for patients with genotypes 2 and 3, rates were 76–82%. Between genotypes 2 and 3, genotype 3 is somewhat more refractory, and some authorities would extend therapy for a full 48 weeks in patients with genotype 3, especially if they have advanced hepatic fibrosis or cirrhosis and/or high-level HCV RNA.

In the initial registration trials for combination PEG IFN plus ribavirin, both combination PEG IFN regimens were compared to standard IFN- $\alpha 2b$ plus ribavirin. Side effects of the combination PEG IFN- $\alpha 2b$ regimen were comparable to those for the combination standard IFN regimen; however, when the combination PEG IFN- $\alpha 2a$ regimen was compared with the combination standard IFN- $\alpha 2b$ regimen, flu-like symptoms and depression were less common in the combination PEG IFN group. Although ascertainment of side effects differed between studies of the two drugs, when each was tested against standard IFN- $\alpha 2b$ plus ribavirin, combination PEG IFN- $\alpha 2a$ plus ribavirin appeared to be better tolerated. In a head-to-head trial of the two PEG IFNs (the IDEAL trial), the two PEG IFNs were found to be comparable in efficacy (achievement of SVR) and tolerability, although headache, nausea, fever, myalgia, depression, and drug

discontinuation for any reason were less frequent in patients treated with PEG IFN- α 2a than standard-dose PEG IFN- α 2b. In contrast, neutropenia and rash were more frequent in patients treated with PEG IFN- α 2a than standard-dose PEG IFN- α 2b. In two subsequent head-to-head trials and a systematic review of randomized trials, PEG IFN- α 2a was more effective than PEG IFN- α 2b (SVR in genotypes 1–4: 48–55% versus 32–40%, respectively). In trials of PEG IFN- α 2b among patients with HCV genotype 1, a broader range of weight-based daily ribavirin doses has been validated: 800 mg for weight <65 kg, 1000 mg for weight 65–85 kg, 1200 mg for weight >85–105 kg, and 1400 mg for weight >105 kg. Recommended doses for the two PEG IFNs plus ribavirin and other comparisons between the two therapies are shown in Table 362-6.

Until the 2011 introduction of protease inhibitors, unless ribavirin was contraindicated (see above), combination PEG IFN plus ribavirin was the recommended course of therapy—24 weeks for genotypes 2 and 3 and 48 weeks for genotype 1. For patients with genotypes 1 and 4, the standard of care now includes protease inhibitors or other direct-acting antiviral agents (see below); however, PEG IFN-ribavirin remained the standard of care for patients with genotypes 2 and 3 until late 2013. For patients treated with combination PEG IFN-ribavirin, measurement of quantitative HCV RNA levels at 12 weeks is helpful in guiding therapy; if a 2- \log_{10} drop in HCV RNA has not been achieved by this time, chances for an SVR are negligible, and additional therapy is futile. If the 12-week HCV RNA has fallen by 2 \log_{10} (EVR), the chances for an SVR at the end of therapy are approximately two-thirds; if the 12-week HCV RNA is undetectable (“complete” EVR), the chances for an SVR exceed 80% (Fig. 362-2). Because absence of an EVR is such a strong predictor of the absence of an ultimate SVR, therapy is discontinued for failure to achieve a 12-week 2- \log_{10} drop in HCV RNA (EVR).

Studies have suggested that the frequency of an SVR to PEG IFN-ribavirin therapy can be increased in patients with baseline variables weighing against a response (e.g., HCV RNA $>8 \times 10^5$ IU/mL, weight >85 kg) by raising the dose of PEG IFN (e.g., to as high as 270 μ g of PEG IFN- α 2a) and/or the dose of ribavirin to as high as 1600 mg daily (if tolerated or supplemented by erythropoietin) or by tailoring treatment based on viral response to prolong the duration of viral clearance before discontinuing therapy, i.e., extending therapy from 48 to 72 weeks for patients with genotype 1 and a slow virologic response (i.e., those whose HCV RNA has not fallen rapidly to undetectable levels within 4 weeks [absence of RVR]). Tailoring therapy based on the kinetics of HCV RNA reduction has also been applied to abbreviating the duration of therapy in patients with genotype 1 (and 4). The results of several clinical trials suggest that, in patients with genotype 1 (and 4) who have a 4-week RVR (which occurs in $\leq 20\%$), especially in the subset with low baseline HCV RNA, 24 weeks of therapy with PEG IFN and weight-based ribavirin suffices, yielding SVR rates of $\sim 90\%$ and comparable to those achieved in this cohort with 48 weeks of therapy. Although initial reports suggested that, for patients with genotype 2 and (somewhat less so) genotype 3, in rapid virologic responders with undetectable HCV RNA at week 4, the total duration of therapy required to achieve an SVR could be as short as 12–16 weeks, a very sizable, definitive subsequent trial showed that relapse is increased if treatment duration is curtailed and that a full 24 weeks is superior for these genotypes (except for the minority with very low baseline levels of HCV RNA).

Persons with chronic HCV infection have been shown to suffer increased liver-related mortality. On the other hand, successful antiviral therapy of chronic hepatitis C resulting in an SVR has been shown to improve survival (and to reduce the need for liver transplantation), to lower the risk of liver failure and liver-related death and all-cause deaths, to slow the progression of chronic hepatitis C, and to reverse fibrosis and even cirrhosis. Although successful treatment reduces mortality in cirrhotic patients (and those with advanced fibrosis) and reduces the likelihood of HCC, the risk of liver-related death and HCC persists, albeit at a much reduced level, necessitating continued clinical monitoring and cancer surveillance after SVR in cirrhotics. On the other hand, in the absence of an SVR,

routine-dose/duration IFN-based therapy does not reduce the risk of HCC. Similarly, for nonresponders to PEG IFN-ribavirin therapy, three trials of long-term maintenance therapy with PEG IFN have shown no benefit in reducing the risk of histologic progression or clinical decompensation, including the development of HCC. For PEG IFN-ribavirin nonresponders who have had a full, adequate course of therapy, the benefit of retreatment—with higher doses or a longer course of the original PEG IFN regimen or the alternative PEG IFN regimen or with a different type of IFN preparation (e.g., consensus IFN)—is marginal at best. Fortunately, such nonresponders can now be retreated with protease inhibitor-based therapy (see following).

FIRST-GENERATION PROTEASE INHIBITORS (2011–2013)

The HCV RNA genome encodes a single polyprotein, which is cleaved during and after translation by host and viral-encoded proteases. One protease involved in the cleavage of the viral polyprotein is an NS3-4A viral protein that has serine protease activity. Telaprevir and boceprevir are serine protease inhibitors that target NS3-4A. In 2011, telaprevir and boceprevir used in combination with PEG IFN and ribavirin were approved by the U.S. Food and Drug Administration (FDA) for the treatment of hepatitis C genotype 1 in adults with stable liver disease, both in patients who have not been treated before or who have failed previous treatment. Because the presently available HCV protease inhibitors have not been studied comprehensively in patients with genotypes other than 1, their use in these populations is not recommended.

Because resistance develops rapidly, both telaprevir and boceprevir must be used in combination with a PEG IFN and ribavirin-based regimen and should never be used alone. Ribavirin in particular appears to reduce relapse rates significantly in protease inhibitor-based regimens, such that those who cannot take or are intolerant to ribavirin are unlikely to benefit from the addition of these agents. All current telaprevir and boceprevir regimens consist of periods of triple therapy (protease inhibitor plus PEG IFN plus ribavirin) and periods of dual therapy (PEG IFN plus ribavirin). Telaprevir regimens begin with 12 weeks of triple therapy followed by dual therapy of a duration based on HCV RNA status at weeks 4 and 12 (“response-guided therapy”) and prior treatment status. Boceprevir-based regimens consist of a 4-week lead-in period of dual (PEG IFN-ribavirin) therapy followed by triple therapy and, in some instances, a further extension of dual therapy, with duration of response-guided therapy based on HCV RNA status at weeks 4, 8, and 24 and prior treatment status (Table 362-7).

For patients with HCV genotype 1, protease inhibitors have significantly improved the frequency of RVRs and SVRs as compared to PEG IFN plus ribavirin alone. In treatment-naïve patients treated with telaprevir, an SVR was seen in up to 79% of patients who received 12 weeks of triple therapy followed by 12–36 weeks of dual therapy, and among those with EVRs (undetectable HCV RNA at weeks 4 and 12) and response-guided therapy stopped at week 24 (12 weeks of triple therapy, then 12 weeks of dual therapy), the rate of SVRs was 83–89% (92% in a subsequent study). In studies with boceprevir in treatment-naïve patients, SVRs were seen in 59–66% of patients, and among those with undetectable HCV RNA at 8 weeks, the SVR rate increased to 86–88%.

Protease inhibitors have also been studied in patients previously treated unsuccessfully with PEG IFN plus ribavirin. In studies with telaprevir, SVRs were seen in 83–88% of patients who had a previous relapse, 54–59% of partial responders (HCV RNA reduced by $\geq 2 \log_{10}$ IU/mL but not to undetectable levels), and 29–33% of null responders (HCV RNA reduced by $< 2 \log_{10}$ IU/mL). With boceprevir, SVRs occurred in 75% of prior relapsers and in 40–52% of previous partial responders; response rates in null responders are similar to those achieved with telaprevir-based therapy. In a substantial proportion of protease inhibitor nonresponders, resistance-associated variants can be identified, but these variants are not archived, and wild-type HCV reemerges in almost all cases within 1.5 to 2 years. SVRs to these protease inhibitors are highest in prior relapsers and

TABLE 362-7 INDICATIONS AND RECOMMENDATIONS FOR ANTIVIRAL THERAPY OF CHRONIC HEPATITIS C^a**Standard Indications for Therapy**

Detectable HCV RNA (with or without elevated ALT)

Portal/bridging fibrosis or moderate to severe hepatitis on liver biopsy (the necessity of a pretreatment biopsy is no longer embraced universally)

Indications for IFN/ribavirin-based therapy apply to adults as well as to children age 2–17, in whom treatment may be considered at reduced weight-based doses (see product inserts); protease inhibitors are not recommended for children age <18 years

Retreatment Recommended**Genotype 1**

Relapsers, partial responders, or nonresponders after a previous course of standard IFN monotherapy or combination standard IFN/ribavirin therapy or PEG IFN/ribavirin

A course of PEG IFN/ribavirin plus protease inhibitor as below

Genotypes 2, 3, 4

Relapsers after a previous course of standard IFN monotherapy or combination standard IFN/ribavirin therapy

A course of PEG IFN plus ribavirin

Nonresponders to a previous course of standard IFN monotherapy or combination standard IFN/ribavirin therapy

A course of PEG IFN plus ribavirin—more likely to achieve a sustained virologic response in white patients without previous ribavirin therapy, with low baseline HCV RNA levels, with a ≥ 2 -log₁₀ reduction in HCV RNA during previous therapy, with genotypes 2 and 3, and without reduction in ribavirin dose

Antiviral Therapy Management Decisions Made on an Individual Basis

Children (age <18 years)—protease inhibitors not recommended.

Age >70 (in protease inhibitor trials, telaprevir trials included patients age 18–70; boceprevir trials included patients >18 years of age [no upper age cutoff])

Mild hepatitis on liver biopsy

Persons with severe renal insufficiency (require reduced doses of PEG IFN and ribavirin)

Long-Term Maintenance Therapy Recommended

Cutaneous vasculitis and glomerulonephritis associated with chronic hepatitis C

Long-Term Maintenance Therapy in Nonresponders Not Recommended**Antiviral Therapy Not Recommended**

Decompensated cirrhosis (except, perhaps, in transplantation centers with experience in graded escalation, low-dose treatment to achieve undetectable HCV RNA prior to transplantation; results are mixed)

Pregnancy (teratogenicity of ribavirin)

Contraindications to use of antiviral medications

Therapeutic Regimens**HCV genotype 1****TREATMENT-NAÏVE (telaprevir and boceprevir) and PRIOR RELAPSE (telaprevir)**

PEG IFN- α 2a 180 μ g weekly plus weight-based ribavirin 1000 mg/d (<75 kg) to 1200 mg/d (\geq 75 kg) or

PEG IFN- α 2b 1.5 μ g/kg weekly plus weight-based ribavirin 800 mg/d (\leq 65 kg), 1000 mg/d (>65–85 kg), 1200 mg/d (>85–105 kg), or 1400 mg/d (>105 kg)

Plus response-guided therapy with a protease inhibitor consisting of either:

Boceprevir 800 mg three times daily with food started after a lead-in treatment of 4 weeks with PEG IFN–ribavirin

- Patients with undetectable HCV RNA at 8 and 24 weeks should receive triple-drug therapy (PEG IFN, ribavirin, boceprevir) through week 28 (4 weeks of PEG IFN–ribavirin then 24 weeks of triple-drug therapy). If HCV RNA is detectable at 4 weeks, continuing therapy through 48 weeks (4 weeks of PEG IFN–ribavirin then 44 weeks of triple-drug therapy) may increase the sustained response rate.
- Patients with detectable HCV RNA at 8 weeks and undetectable at 24 weeks should receive triple-drug therapy (PEG IFN, ribavirin, boceprevir) through week 36 (4 weeks of PEG IFN–ribavirin then 32 weeks of triple-drug therapy) followed by a return to PEG IFN–ribavirin for 12 more weeks, for a total treatment duration of 48 weeks.
- Patients with cirrhosis who are treatment-naïve and have undetectable HCV RNA at weeks 8 and 24 should continue triple-drug therapy (PEG IFN, ribavirin, boceprevir) through 48 weeks (4 weeks of PEG IFN–ribavirin then 44 weeks of triple-drug therapy).
- Stopping rules for futility: HCV RNA \geq 100 IU/mL at week 12 or any detectable HCV RNA at week 24

or

Telaprevir 750 mg three times daily with fatty food started at the beginning of therapy without a PEG IFN–ribavirin lead-in

- Patients with undetectable HCV RNA at 4 and 12 weeks should receive triple-drug therapy (PEG IFN, ribavirin, telaprevir) for 12 weeks then PEG IFN and ribavirin for another 12 weeks, for a total of 24 weeks.
- Patients with detectable HCV RNA at 4 or 12 weeks and undetectable at 24 weeks should receive triple-drug therapy (PEG IFN, ribavirin, telaprevir) for 12 weeks, then PEG IFN–ribavirin for another 36 weeks, for a total treatment duration of 48 weeks.
- Patients with cirrhosis who are treatment-naïve and have undetectable HCV RNA at 4 and 12 weeks should receive triple-drug therapy for 12 weeks then PEG IFN–ribavirin for another 36 weeks, for a total treatment duration of 48 weeks.
- Stopping rules for futility: HCV RNA >1000 IU/mL at week 4 or 12 or any detectable HCV RNA at week 24

TREATMENT-EXPERIENCED

PEG IFN- α 2a 180 μ g weekly plus weight-based ribavirin 1000 mg/d (<75 kg) to 1200 mg/d (\geq 75 kg) or

PEG IFN- α 2b 1.5 μ g/kg weekly plus weight-based ribavirin 800 mg/d (\leq 65 kg), 1000 mg/d (>65–85 kg), 1200 mg/d (>85–105 kg), or 1400 mg/d (>105 kg)

Plus a protease inhibitor consisting of either:

Response-guided therapy with boceprevir 800 mg three times daily with food started after a lead-in treatment of 4 weeks with PEG IFN–ribavirin

- For prior relapsers and partial responders (HCV RNA reduction of ≥ 2 log₁₀ during previous therapy), follow the response-guided algorithm below; for prior null responders (HCV RNA reduction <2 log₁₀ during previous therapy), a full 48-week course of therapy (4-week PEG IFN–ribavirin lead-in, followed by 44 weeks of triple-drug therapy [PEG IFN, ribavirin, boceprevir]) is recommended.

(Continued)

- Patients with undetectable HCV RNA at 8 and 24 weeks should receive triple-drug therapy (PEG IFN, ribavirin, boceprevir) through week 36 (4 weeks of PEG IFN–ribavirin then 32 weeks of triple-drug therapy). If HCV RNA is detectable at 4 weeks, continuing therapy through 48 weeks (4 weeks of PEG IFN–ribavirin then 44 weeks of triple therapy) may increase the sustained response rate.
- Patients with detectable HCV RNA at 8 weeks and undetectable at week 24 should receive triple-drug therapy (PEG IFN, ribavirin, boceprevir) through week 36 (4 weeks of PEG IFN–ribavirin then 32 weeks of triple-drug therapy), followed by a return to PEG IFN–ribavirin for 12 more weeks, for a total treatment duration of 48 weeks.
- Patients with cirrhosis who are treatment-experienced and have undetectable HCV RNA at weeks 8 and 24 should continue triple-drug therapy (PEG IFN, ribavirin, boceprevir) through 48 weeks (4 weeks of PEG IFN–ribavirin then 44 weeks of triple-drug therapy).
- Stopping rules for futility: HCV RNA ≥ 100 IU/mL at week 12 or any detectable HCV RNA at week 24

or

Telaprevir 750 mg three times daily with fatty food started at the beginning of therapy without a PEG IFN–ribavirin lead-in and without a response-guided approach, i.e., all patients receive a full 48-week course, independent of early responsiveness.

- For prior relapsers, follow guidelines for treatment-naïve patients above.
- Prior partial responders and null responders should receive triple-drug therapy (PEG IFN, ribavirin, telaprevir) for 12 weeks then PEG IFN and ribavirin for another 36 weeks, for a total of 48 weeks.
- Stopping rules for futility: HCV RNA > 1000 IU/mL at week 4 or 12 or any detectable HCV RNA at week 24

HCV genotype 1 but protease inhibitors unavailable or contraindicated: 48 weeks of therapy

PEG IFN- $\alpha 2a$ 180 μ g weekly plus weight-based ribavirin 1000 mg/d (< 75 kg) to 1200 mg/d (≥ 75 kg) or

PEG IFN- $\alpha 2b$ 1.5 μ g/kg weekly plus weight-based ribavirin 800 mg/d (≤ 65 kg), 1000 mg/d (> 65 –85 kg), 1200 mg/d (> 85 –105 kg), or 1400 mg/d (> 105 kg)

HCV genotype 4: 48 weeks of PEG IFN–ribavirin therapy

PEG IFN- $\alpha 2a$ 180 μ g weekly plus weight-based ribavirin 1000 mg/d (< 75 kg) to 1200 mg/d (≥ 75 kg) or

PEG IFN- $\alpha 2b$ 1.5 μ g/kg weekly plus weight-based ribavirin 800 mg/d (≤ 65 kg), 1000 mg/d (> 65 –85 kg), 1200 mg/d (> 85 –105 kg), or 1400 mg/d (> 105 kg)

- Treatment should be discontinued in patients who do not achieve an early virologic response at week 12.

- Patients who do achieve an early virologic response should be retested at week 24, and treatment should be discontinued if HCV RNA remains detectable.

HCV genotypes 2 and 3: 24 weeks of therapy

PEG IFN- $\alpha 2a$ 180 μ g weekly plus ribavirin 800 mg/d or

PEG IFN- $\alpha 2b$ 1.5 μ g/kg weekly plus ribavirin 800 mg/d (for patients with genotype 3 who have advanced fibrosis and/or high-level HCV RNA, a full 48 weeks of therapy may be preferable)

For HCV/HIV co-infected patients: 48 weeks, regardless of genotype, of weekly PEG IFN- $\alpha 2a$ (180 μ g) or PEG IFN- $\alpha 2b$ (1.5 μ g/kg) plus a daily ribavirin dose of at least 600–800 mg, up to full weight-based 1000–1400 mg dosing if tolerated. Protease inhibitors may be used for genotype 1; however, because of potential drug-drug interactions between HCV protease inhibitors and HIV antiretroviral drugs, HCV protease inhibitors should be used cautiously in HCV/HIV co-infected patients. If protease inhibitors are used, a full 48-week course is recommended without response-guided therapy. For boceprevir, 4 weeks of PEG IFN–ribavirin lead-in, followed by 44 weeks of triple-drug therapy (PEG IFN, ribavirin, boceprevir). For telaprevir, 12 weeks of triple-drug therapy (PEG IFN, ribavirin, telaprevir), followed by 36 weeks of PEG IFN–ribavirin therapy. Stopping rules for futility are as noted above.

Features Associated with Reduced Responsiveness

Single nucleotide polymorphism (SNP) T allele (as opposed to C allele) at *IL28B* locus

Genotype 1a (compared to 1b)

High-level HCV RNA ($> 800,000$ IU/mL)^b

Advanced fibrosis (bridging fibrosis, cirrhosis)

Long-duration disease

Age $> 40^b$

High HCV quasispecies diversity

Immunosuppression

African-American ethnicity

Latino ethnicity

Obesity

Hepatic steatosis

Insulin resistance, type 2 diabetes mellitus^b

Reduced adherence (lower drug doses and reduced duration of therapy)

For boceprevir, $< 1 \log_{10}$ reduction in HCV RNA during 4-week PEG IFN–ribavirin lead-in

For protease inhibitor therapy, absence of extended rapid virologic response (eRVR), i.e., detectable HCV RNA, at weeks 4 and 12 for telaprevir; at weeks 8 and 24 for boceprevir

^aAs this chapter was going to press, two additional drugs were approved for hepatitis C, simeprevir and sofosbuvir. Rapidly evolving new recommendations are supplanting the recommendations in this table; for up-to-date treatment recommendations, please see www.hcvguidelines.org. ^bLess influential in patients treated with protease inhibitors.

Abbreviations: ALT, alanine aminotransferase; HCV, hepatitis C virus; IFN, interferon; PEG IFN, pegylated interferon; IU, international units (1 IU/mL is equivalent to ~ 2.5 copies/mL).

treatment-naïve patients (white $>$ black ethnicity), lower in prior partial responders, lower still in prior null responders, and lowest in cirrhotic prior null responders (Fig. 362-3). Responses to protease inhibitor triple-drug regimens are higher in patients with *IL28B* C than non-C genotypes, HCV genotype 1b than genotype 1a, less advanced than more advanced fibrosis stage, whites than blacks, lower body mass index (BMI) than elevated BMI, and, for boceprevir, achievement of a $> 1 \log_{10}$ HCV RNA reduction during 4 weeks of

PEG IFN–ribavirin lead-in therapy. Age and HCV RNA level are less influential and insulin resistance is noninfluential on response to these antiviral agents.

Both protease inhibitors have potential toxicities. Telaprevir is associated with a severe, generalized (trunk and extremities), often confluent, maculopapular, pruritic rash in $\sim 6\%$ of treated patients. Other common side effects include pruritus, rectal burning, nausea, diarrhea, fatigue, and anemia, which may be relatively refractory,

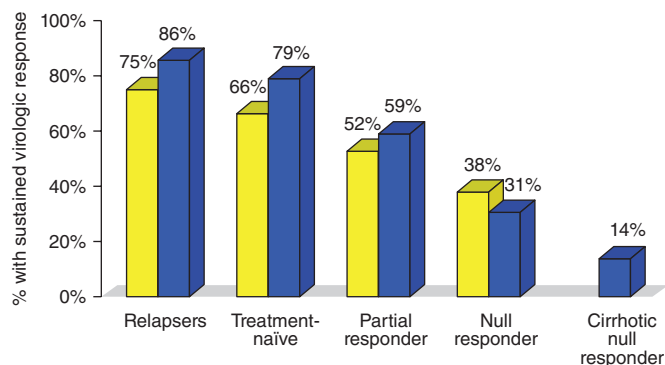


FIGURE 362-3 Maximal efficacy (sustained virologic responses, SVR) of telaprevir (blue bars) and boceprevir (yellow bars) reported in phase III clinical trials. (Figure created using data from Bacon BR et al: *N Engl J Med* 364:1207, 2011; Jacobson IM et al: *N Engl J Med* 364:2405, 2011; Poordad F et al: *N Engl J Med* 364:1195, 2011; Zeuzem S et al: *N Engl J Med* 364:2417, 2011; Vierling JM et al: *Hepatology* 54 [Suppl 1]:796A-797A, 2011; Ghany MG et al: *Hepatology* 54:1433, 2011.)

occasionally requiring transfusion. Complete blood counts should be obtained at baseline and then at 2, 4, 8, and 12 weeks after starting telaprevir. Anemia can occur in half of boceprevir-treated patients, as can neutropenia in up to 30% and thrombocytopenia in 3–4%. Complete blood counts should be obtained at baseline and then at 4, 8, and 12 weeks after starting boceprevir. Other side effects of boceprevir include fatigue, nausea, headache, dysgeusia (altered or unpleasant taste), dry mouth, vomiting, and diarrhea.

Use of protease inhibitors is further complicated by numerous drug-drug interactions. As telaprevir and boceprevir are both eliminated by and inhibit CYP3A4, these agents should not be administered with other medications that induce CYP3A4 or are dependent on CYP3A4 for elimination. Care should be taken to examine for any potential interactions between protease inhibitors and other medications the patient may be taking, because serious adverse events can occur. A convenient website is available to check for such drug-drug interactions (www.hep-druginteractions.org).

TREATMENT RECOMMENDATIONS*

Prior to therapy, HCV genotype should be determined, because the genotype dictates the duration of therapy and potentially the agents to be used. PEG IFN plus ribavirin represents the foundation of treatment for all HCV genotypes; patients infected with genotype 1 should also receive a protease inhibitor (telaprevir or boceprevir) when these are available and not contraindicated (Table 362-7). For chronic HCV genotype 1 infection, the AASLD and EASL published treatment guidelines in 2011 reflecting FDA-approved indications for the new protease inhibitors, and in 2012, United Kingdom and French consensus guidelines were published. For treatment-naïve patients and prior relapsers, response-guided therapy with telaprevir or boceprevir is recommended. For telaprevir, the regimen consists of 12 weeks of triple therapy, followed by 12 or 36 weeks of PEG IFN–ribavirin consolidation, depending on whether extended RVR milestones (HCV RNA undetectable at weeks 4 and 12) are met.

*As this chapter was going to press, two additional antiviral drugs, a second-generation protease inhibitor simeprevir and nucleoside analogue polymerase inhibitor sofosbuvir were approved for the treatment of hepatitis C. Simeprevir, which is effective for genotype 1, must be administered, like first-generation protease inhibitors, for 12 weeks with PEG IFN and ribavirin, followed by another 12 weeks of PEG IFN and ribavirin (no response-guided therapy). Sofosbuvir, the more convenient and broadly applicable of the two new drugs, must be administered with PEG IFN and ribavirin but for only 12 weeks in patients with genotypes 1, 4–6; for patients with genotypes 2 and 3, PEG IFN is not required. Sofosbuvir plus ribavirin are administered for 12 weeks in genotype 2 and for 24 weeks in genotype 3. Antiviral therapy is evolving very rapidly; by the end of 2014, all-oral, interferon-free combinations (e.g., sofosbuvir plus the NS5a inhibitor ledipasvir) will supplant earlier treatment regimens. For updated treatment recommendations, please consult www.hcvguidelines.org.

For boceprevir, the regimen consists of a 4-week PEG IFN–ribavirin lead-in period, followed by 24–32 weeks of triple-drug therapy, depending on whether HCV RNA milestones (undetectable at weeks 8 and 24) are met; if HCV RNA is detectable at week 8 but undetectable at week 24, after 36 weeks of therapy (4-week PEG IFN–ribavirin lead-in plus 32 weeks of triple-drug therapy), an additional 12 weeks of PEG IFN–ribavirin consolidation is recommended. For prior partial and null responders, a full 48-week course of telaprevir (no lead-in period, no response-guided therapy) is recommended; for boceprevir, a 4-week PEG IFN–ribavirin lead-in period is followed by response-guided therapy (32 weeks of triple-drug therapy if HCV RNA is undetectable at weeks 8 and 24 or, if HCV RNA is still detectable at week 8 [but undetectable at week 24], 32 weeks of triple-drug therapy followed by 12 weeks of PEG IFN–ribavirin consolidation). For cirrhotic patients (and for any boceprevir-treated patient whose HCV RNA does not fall by $>1 \log_{10}$ by week 4), a full 48-week course without response-guided therapy should be considered.

Monitoring of HCV plasma RNA is crucial in assessing response to therapy. The goal of treatment is to eradicate HCV RNA, which is predicted by the absence of HCV RNA by PCR 6 months after stopping treatment (SVR). When therapy relied on PEG IFN and ribavirin, failure to achieve a 2-log_{10} drop in HCV RNA by week 12 of therapy (EVR) rendered it unlikely that further therapy would result in an SVR. When PEG IFN and ribavirin are part of a protease inhibitor regimen, HCV RNA should be measured at baseline and at weeks 4, 8 (for boceprevir), 12, and 24 to assess response to treatment and to aid in decisions regarding treatment duration (response-guided therapy), as well as 12 and 24 weeks after therapy. Stopping rules are important to prevent the emergence of resistance; if HCV RNA is >1000 IU/mL at 4 or 12 weeks of telaprevir (or still detectable at week 24), or if HCV RNA is ≥ 100 IU/mL at week 12 of boceprevir (or detectable at week 24), all treatment should be stopped.

INDICATIONS FOR ANTIVIRAL THERAPY*

Patients with chronic hepatitis C who have detectable HCV RNA in serum, whether or not aminotransferase levels are increased, and chronic hepatitis of at least moderate grade and stage (portal or bridging fibrosis) are candidates for antiviral therapy with PEG IFN plus ribavirin. Most authorities recommend 800 mg of ribavirin for patients with genotypes 2 and 3 for both types of PEG IFN and weight-based 1000–1200 mg (when used with PEG IFN- α 2a) or 800–1400 mg (when used with PEG IFN- α 2b) ribavirin for patients with genotype 1 (and 4), unless ribavirin is contraindicated (Table 362-7). These PEG IFN and ribavirin doses are used with protease inhibitors for patients with genotype 1 (Table 362-7). Although patients with persistently normal ALT activity tend to progress histologically very slowly or not at all, they respond to antiviral therapy just as well as do patients with elevated ALT levels; therefore, although observation without therapy is an option, such patients are potential candidates for antiviral therapy. As noted above, therapy with IFN has been shown to improve survival and complication-free survival and to slow progression of (and to reverse) fibrosis.

HCV genotype determines the duration of PEG IFN and ribavirin therapy: 24 weeks for those with genotypes 2 and 3 and 48 weeks for patients with genotypes 4 and 1 (in patients for whom protease inhibitors are not available or contraindicated). For patients with genotype 4, treatment should be discontinued in patients who do not achieve an EVR at week 12. For patients with genotypes 2 and 3, a full, 24-week course is most effective, although the duration may be reduced to 12–16 weeks for patients with genotype 2, a low baseline level of viremia, and an RVR, especially to be considered for patients who tolerate therapy poorly. Also, consideration should be given to increasing the duration of therapy to 48 weeks for patients with genotype 3 who have advanced fibrosis and/or a high baseline level of viremia. As noted above, the absence of a 2-log_{10} drop in HCV RNA at week 12 (EVR) weighs heavily against the likelihood of an SVR; therefore, measuring HCV RNA at 12 weeks is recommended routinely (Fig. 362-2), and therapy can be discontinued if an EVR is not achieved. Among patients with genotype 4 who achieve an

2048 EVR ($\geq 2\text{-log}_{10}$ HCV RNA reduction) but in whom HCV RNA remains detectable at week 24, an SVR is unlikely, and therapy can be discontinued. Although response rates are lower in patients with certain pretreatment variables, selection for treatment should not be based on symptoms, genotype, HCV RNA level, mode of acquisition of hepatitis C, or advanced hepatic fibrosis. Patients with cirrhosis can respond and should not be excluded as candidates for therapy. For patients being treated with telaprevir and boceprevir, treating physicians should explain the negative impact of non-C *IL28B* genotype and advanced fibrosis on outcome.

Patients who have relapsed after, or failed to respond to (Fig. 362-2), a course of IFN monotherapy are potential candidates for retreatment with PEG IFN plus ribavirin (i.e., a more effective treatment regimen is required), and this approach remains current for patients with genotypes 2, 3, or 4; however, for patients with genotype 1, combination protease inhibitor/PEG IFN/ribavirin therapy is indicated. For patients with genotypes 2, 3, or 4 who were nonresponders to a prior course of IFN monotherapy, retreatment with IFN monotherapy or combination IFN plus ribavirin therapy is unlikely to achieve an SVR; however, a trial of combination PEG IFN plus ribavirin may be worthwhile, although an SVR is the outcome in <15–20% of patients. SVRs to retreatment of nonresponders are more frequent in those who had never received ribavirin in the past, those with genotypes 2 and 3, those with low pretreatment HCV RNA levels, and noncirrhotics, but less frequent in African Americans, those who failed to achieve a substantial reduction in HCV RNA during their previous course of therapy (null responders, Fig. 362-2), and those who required ribavirin dose reductions. Potential approaches to improving responsiveness to PEG IFN–ribavirin in prior nonresponders include longer duration of treatment; higher doses of PEG IFN, ribavirin, or both; and switching to a different IFN preparation; however, as noted above, none of these approaches achieves more than a marginal benefit. Treatment with a protease inhibitor–based regimen should be pursued in patients with genotype 1 who have relapsed after or not responded to prior treatment with IFN monotherapy or PEG IFN plus ribavirin, unless these protease inhibitors are not available or contraindicated (Table 362-7).

Early PEG IFN treatment is indicated for persons with acute hepatitis C; ribavirin, which is used frequently in such instances, has not been shown to improve efficacy over that of PEG IFN alone, and the new protease inhibitors have not been approved for acute hepatitis C (Chap. 360). In patients with biochemically and histologically mild chronic hepatitis C, the rate of progression is slow, and monitoring without therapy is an option; however, such patients respond just as well to combination PEG IFN plus ribavirin therapy or triple-drug, protease-based therapy (for genotype 1) as those with elevated ALT and more histologically severe hepatitis. Therefore, therapy for these patients should be considered and the decision made based on such factors as patient motivation, genotype, stage of fibrosis, age, and comorbid conditions. A pretreatment liver biopsy to assess histologic grade and stage provides substantial information about progression of hepatitis C in the past, has prognostic value for future progression, and can identify such histologic factors as steatosis and stage of fibrosis, which can influence responsiveness to therapy. As therapy has improved for patients with a broad range of histologic severity, and as noninvasive laboratory markers and imaging correlates of fibrosis have gained popularity, some authorities, especially in Europe, place less value on, and do not recommend, pretreatment liver biopsies. On the other hand, serum markers of fibrosis are not considered sufficiently accurate, and histologic findings provide important prognostic information to physician and patient. Therefore, although the contemporary role of a pretreatment liver biopsy commands less of a consensus, a pretreatment liver biopsy still provides useful information and should be considered.

Patients with compensated cirrhosis can respond to therapy, although their likelihood of a sustained response is lower than in noncirrhotics; moreover, survival has been shown to improve after successful antiviral therapy in cirrhotics. Similarly, although several retrospective studies have suggested that antiviral therapy in

cirrhotics with chronic hepatitis C, independent of treatment outcome per se, reduces the frequency of HCC, less advanced disease in the treated cirrhotics, not treatment itself (i.e., lead-time bias), may have accounted for the reduced frequency of HCC observed in the treated cohorts in these reports; prospective studies to address this question have failed to demonstrate benefit, unless an SVR is achieved. Patients with decompensated cirrhosis are not candidates for IFN-based antiviral therapy but should be referred for liver transplantation. Some liver transplantation centers have evaluated progressively escalated, low-dose antiviral therapy in an attempt to eradicate hepatitis C viremia prior to transplantation; however, such therapy has been shown to reduce but not to prevent the risk of HCV reinfection after transplantation. After liver transplantation for end-stage liver disease caused by hepatitis C, recurrent hepatitis C is the rule, and the pace of disease progression is more accelerated than in immunocompetent patients (Chap. 368). Current therapy with PEG IFN and ribavirin after liver transplantation is unsatisfactory in most patients, but attempts to minimize immunosuppression are beneficial. Early experience with protease inhibitor–based therapy is encouraging, but inhibition of CYP3A4 by protease inhibitors can lead to markedly increased levels of immunosuppressive calcineurin inhibitors (especially tacrolimus), which requires intensive monitoring and can be very challenging. The cutaneous and renal vasculitis of HCV-associated essential mixed cryoglobulinemia (Chap. 360) may respond to antiviral therapy, but sustained responses are rare after discontinuation of therapy; therefore, prolonged, perhaps indefinite, therapy (as reported with IFN-based therapy) is recommended in this group (no indication for prolonged protease inhibitor therapy exists currently). Anecdotal reports suggest that antiviral therapy may be effective in porphyria cutanea tarda or lichen planus associated with hepatitis C.

In patients with HCV/HIV co-infection, hepatitis C is more progressive and severe than in HCV-monoinfected patients. Although patients with HCV/HIV co-infection respond to antiviral therapy for hepatitis C, they do not respond as well as patients with HCV infection alone. Four large national and international trials of antiviral therapy among patients with HCV/HIV co-infection have shown that PEG IFN (both $\alpha 2a$ and $\alpha 2b$) plus ribavirin (daily doses ranging from flat-dosed 600–800 mg to weight-based 1000/1200 mg) is superior to standard IFN regimens; however, SVR rates were lower than in HCV-monoinfected patients, ranging from 14 to 38% for patients with genotypes 1 and 4 and from 44 to 73% for patients with genotypes 2 and 3. In the three largest trials, all patients, including those with genotypes 2 and 3, were treated for a full 48 weeks. In addition, tolerability of therapy was lower than in HCV-monoinfected patients; therapy was discontinued because of side effects in 12–39% of patients in these clinical trials. Based on these trials, weekly PEG IFN plus daily ribavirin at a daily dose of at least 600–800 mg, up to full weight-based doses, at doses recommended for HCV-monoinfected patients, if tolerated, is recommended for a full 48 weeks, regardless of genotype. An alternative recommendation for ribavirin doses was issued by a European Consensus Conference and consisted of standard, weight-based 1000–1200 mg for genotypes 1 and 4, but 800 mg for genotypes 2 and 3. A head-to-head trial of combination PEG IFN–ribavirin therapy in HCV/HIV co-infection demonstrated statistically indistinguishable efficacy of the two types of PEG IFN, despite a small advantage for PEG IFN– $\alpha 2a$: For PEG IFN– $\alpha 2b$ and – $\alpha 2a$, SVRs occurred in 28% versus 32%, respectively, of patients with genotypes 1 and 4 and in 62% versus 71%, respectively, of patients with genotypes 2 and 3.

Although data are limited and recommendations pending, protease inhibitors may be used for genotype 1; however, because of potential drug-drug interactions between HCV protease inhibitors and HIV antiretroviral drugs (especially in ritonavir-boosted HIV protease inhibitors), HCV protease inhibitors should be used cautiously in HCV-HIV co-infected patients. If protease inhibitors are used, a full 48-week course is recommended without response-guided therapy: for boceprevir, 4 weeks of PEG IFN–ribavirin lead-in, followed by 44 weeks of triple-drug therapy (PEG IFN, ribavirin, boceprevir), and

for telaprevir, 12 weeks of triple-drug therapy (PEG IFN, ribavirin, telaprevir), followed by 36 weeks of PEG IFN–ribavirin therapy. In preliminary trials among HIV-HCV co-infected patients, telaprevir-based triple-drug therapy (independent of whether they were receiving antiretroviral therapy [no antiretroviral drugs, efavirenz-tenofovir-emtricitabine, or ritonavir-boosted atazanavir-tenofovir-emtricitabine or lamivudine]) resulted in an SVR in 28 of 38 patients (74%), compared with 10 of 22 control patients (45%) treated with PEG IFN–ribavirin (60 study subjects); boceprevir-based triple-drug therapy (all were also receiving antiretroviral therapy) resulted in an SVR in 40 of 64 patients (63%), compared with 10 of 34 control patients (29%) treated with PEG IFN–ribavirin (98 study subjects). Thus, for the prior standard of care, PEG IFN plus ribavirin, although the likelihood of an SVR is lower for HIV-HCV co-infected patients than for HCV-monoinfected patients, for protease inhibitor–based regimens, rates of SVR are comparable in HIV-HCV co-infected and HCV-monoinfected patients.

In HCV/HIV-infected patients, ribavirin can potentiate the toxicity of didanosine (e.g., lactic acidosis) and the lipoatrophy of stavudine, and zidovudine can exacerbate ribavirin-associated hemolytic anemia; therefore, these drug combinations should be avoided.

Patients with a history of injection drug use and alcoholism can be treated successfully for chronic hepatitis C, preferably in conjunction with drug and alcohol treatment programs. Because ribavirin is excreted renally, patients with end-stage renal disease, including those undergoing dialysis (which does not clear ribavirin), are not ideal candidates for ribavirin therapy. Rare reports suggest that reduced-dose ribavirin can be used, but the frequency of anemia is very high, and data on efficacy are limited. If patients with renal failure (glomerular filtration rate <60 mL/min) are treated, the PEG IFN- α 2a dose should be reduced from 180 to 135 μ g weekly and the PEG IFN- α 2b dose reduced from 1.5 to 1 μ g/kg weekly; similarly, the daily ribavirin dose in this population should be reduced to 200–800 mg (but not used or used cautiously at very low doses) if hemodialysis is required. Neither the optimal regimen nor the efficacy of therapy is well established in this population.

NOVEL ANTIVIRALS*

To date, attempts to develop better-tolerated ribavirin successors or improved types of IFN- α or longer acting IFNs than PEG IFN have not been successful. The demonstration that responsiveness to antiviral therapy is influenced by genetic variation in *IL28B*, which codes for IFN- λ (as noted above), raises the possibility that IFN- λ might be an effective or even more effective IFN for treating hepatitis C; early trials are in progress, but elevations of aminotransferase levels in treated subjects have raised concerns and delayed development. Beyond telaprevir and boceprevir, other direct antivirals that target HCV polymerase, protease, or NS5A (a membrane phosphoprotein component of the viral replication complex) are being investigated, as well as agents that can target host-encoded proteins. Among the novel antivirals are drugs with improved pharmacokinetic and resistance profiles, less treatment complexity, pangenotypic activity, fewer side effects, and fewer drug-drug interactions.* The pace of successful trials of all-oral regimens has accelerated. All-oral combinations of a second-generation protease inhibitor (asunaprevir) plus an NS5A inhibitor (daclatasvir); of a uridine nucleoside polymerase inhibitor (sofosbuvir)* plus ribavirin; of a polymerase inhibitor (sofosbuvir) plus an NS5A inhibitor (ledipasvir or daclatasvir) and ribavirin; and of combinations of a ritonavir-boosted protease inhibitor (ABT-450) plus a nonnucleoside polymerase inhibitor (ABT-333) plus an NS5A inhibitor (ABT-267) with or without ribavirin have been studied in clinical trials. Several of these drug combinations have achieved SVR rates exceeding 90%, even approaching 100%, for both treatment-naïve and treatment-experienced patients (including patients who failed to respond to first-generation protease inhibitors), across all HCV genotypes and independent of host *IL28B* genotype, and with treatment durations of 12–24 weeks or even shorter (8 weeks). Potentially, as early as 2014 or 2015, such combinations of direct antiviral agents will

be used in drug cocktails that may replace IFN-based regimens entirely.

Less advanced is development of inhibitors of host proteins, such as oral, nonimmunosuppressive inhibitors of cyclophilin A (which interacts with NS5A during HCV replication) and subcutaneous antisense antagonists of host liver-expressed micro-RNA-122 (which promotes HCV replication). Given the accelerated progress of all-oral, short-treatment-duration, high-efficacy, direct-acting antivirals, these alternative approaches may not be practical or competitive.

AUTOIMMUNE HEPATITIS

DEFINITION

Autoimmune hepatitis is a chronic disorder characterized by continuing hepatocellular necrosis and inflammation, usually with fibrosis, which can progress to cirrhosis and liver failure. When fulfilling criteria of severity, this type of chronic hepatitis, when untreated, may have a 6-month mortality of as high as 40%. Based on contemporary estimates of the natural history of autoimmune hepatitis, the 10-year survival is 80–98% for treated and 67% for untreated patients. The prominence of extrahepatic features of autoimmunity and seroimmunologic abnormalities in this disorder supports an autoimmune process in its pathogenesis; this concept is reflected in the prior labels *lupoid* and *plasma cell hepatitis*. Autoantibodies and other typical features of autoimmunity, however, do not occur in all cases; among the broader categories of “idiopathic” or cryptogenic chronic hepatitis, many, perhaps the majority, are probably autoimmune in origin. Cases in which hepatotropic viruses, metabolic/genetic derangements (including nonalcoholic fatty liver disease), and hepatotoxic drugs have been excluded represent a spectrum of heterogeneous liver disorders of unknown cause, a proportion of which are most likely autoimmune hepatitis.

IMMUNOPATHOGENESIS

The weight of evidence suggests that the progressive liver injury in patients with autoimmune hepatitis is the result of a cell-mediated immunologic attack directed against liver cells. In all likelihood, predisposition to autoimmunity is inherited, whereas the liver specificity of this injury is triggered by environmental (e.g., chemical, drug [e.g., minocycline], or viral) factors. For example, patients have been described in whom apparently self-limited cases of acute hepatitis A, B, or C led to autoimmune hepatitis, presumably because of genetic susceptibility or predisposition. Evidence to support an autoimmune pathogenesis in this type of hepatitis includes the following: (1) In the liver, the histopathologic lesions are composed predominantly of cytotoxic T cells and plasma cells; (2) circulating autoantibodies (nuclear, smooth muscle, thyroid, etc.; see below), rheumatoid factor, and hyperglobulinemia are common; (3) other autoimmune disorders—such as thyroiditis, rheumatoid arthritis, autoimmune hemolytic anemia, ulcerative colitis, membranoproliferative glomerulonephritis, juvenile diabetes mellitus, celiac disease, and Sjögren’s syndrome—occur with increased frequency in patients and in their relatives who have autoimmune hepatitis; (4) histocompatibility haplotypes associated with autoimmune diseases, such as HLA-B1, -B8, -DR3, and -DR4 as well as extended haplotype *DRB1*0301* and *DRB1*0401* alleles, are common in patients with autoimmune hepatitis; and (5) this type of chronic hepatitis is responsive to glucocorticoid/immunosuppressive therapy, effective in a variety of autoimmune disorders.

Cellular immune mechanisms appear to be important in the pathogenesis of autoimmune hepatitis. In vitro studies have suggested that in patients with this disorder, CD4⁺ T lymphocytes are capable of becoming sensitized to hepatocyte membrane proteins and of destroying liver cells. Molecular mimicry by cross-reacting antigens that contain epitopes similar to liver antigens is postulated to activate these T cells, which infiltrate, and result in injury to, the liver. Abnormalities of immunoregulatory control over cytotoxic lymphocytes (impaired regulatory CD4⁺CD25⁺ T cell influences) may play a role as well. Studies of genetic predisposition to autoimmune hepatitis demonstrate that certain haplotypes are associated with the disorder, as enumerated above,

2050 as are polymorphisms in cytotoxic T lymphocyte antigens (*CTLA-4*) and tumor necrosis factor α (*TNFA*2*). The precise triggering factors, genetic influences, and cytotoxic and immunoregulatory mechanisms involved in this type of liver injury remain incompletely defined.

Intriguing clues into the pathogenesis of autoimmune hepatitis come from the observation that circulating autoantibodies are prevalent in patients with this disorder. Among the autoantibodies described in these patients are antibodies to nuclei (so-called antinuclear antibodies [ANAs], primarily in a homogeneous pattern) and smooth muscle (so-called anti-smooth-muscle antibodies, directed at actin, vimentin, and skeleton), antibodies to F-actin, antibodies to liver-kidney microsomes (anti-LKM, see below), antibodies to “soluble liver antigen” (directed against a uracil-guanine-adenine transfer RNA suppressor protein), antibodies to α -actinin, and antibodies to the liver-specific asialoglycoprotein receptor (or “hepatic lectin”) and other hepatocyte membrane proteins. Although some of these provide helpful diagnostic markers, their involvement in the pathogenesis of autoimmune hepatitis has not been established.

Humoral immune mechanisms have been shown to play a role in the extrahepatic manifestations of autoimmune and idiopathic hepatitis. Arthralgias, arthritis, cutaneous vasculitis, and glomerulonephritis occurring in patients with autoimmune hepatitis appear to be mediated by the deposition of circulating immune complexes in affected tissue vessels, followed by complement activation, inflammation, and tissue injury. While specific viral antigen-antibody complexes can be identified in acute and chronic viral hepatitis, the nature of the immune complexes in autoimmune hepatitis has not been defined.

CLINICAL FEATURES

Many of the *clinical features* of autoimmune hepatitis are similar to those described for chronic viral hepatitis. The onset of disease may be insidious or abrupt; the disease may present initially like, and be confused with, acute viral hepatitis; a history of recurrent bouts of what had been labeled *acute hepatitis* is not uncommon. In approximately a quarter of patients, the diagnosis is made in the absence of symptoms, based on abnormal liver laboratory tests. A subset of patients with autoimmune hepatitis has distinct features. Such patients are predominantly young to middle-aged women with marked hyperglobulinemia and high-titer circulating ANAs. This is the group with positive lupus erythematosus (LE) preparations (initially labeled “*lupoid*” hepatitis) in whom other autoimmune features are common. Fatigue, malaise, anorexia, amenorrhea, acne, arthralgias, and jaundice are common. Occasionally, arthritis, maculopapular eruptions (including cutaneous vasculitis), erythema nodosum, colitis, pleurisy, pericarditis, anemia, azotemia, and sicca syndrome (keratoconjunctivitis, xerostomia) occur. In some patients, complications of cirrhosis, such as ascites and edema (associated with portal hypertension and hypoalbuminemia), encephalopathy, hypersplenism, coagulopathy, or variceal bleeding may bring the patient to initial medical attention.

The course of autoimmune hepatitis may be variable. In patients with mild disease or limited histologic lesions (e.g., piecemeal necrosis without bridging), progression to cirrhosis is limited, but, even in this subset, clinical monitoring is important to identify progression; up to half left untreated can progress to cirrhosis over the course of 15 years. In North America, cirrhosis at presentation is more common in African Americans than in whites. In those with severe symptomatic autoimmune hepatitis (aminotransferase levels >10 times normal, marked hyperglobulinemia, “aggressive” histologic lesions—bridging necrosis or multilobular collapse, cirrhosis), the 6-month mortality without therapy may be as high as 40%. Such severe disease accounts for only 20% of cases; the natural history of milder disease is variable, often accentuated by spontaneous remissions and exacerbations. Especially poor prognostic signs include the presence histologically of multilobular collapse at the time of initial presentation and failure of serum bilirubin to improve after 2 weeks of therapy. Death may result from hepatic failure, hepatic coma, other complications of cirrhosis (e.g., variceal hemorrhage), and intercurrent infection. In patients with established cirrhosis, HCC may be a late complication

(**Chap. 111**) but occurs less frequently than in cirrhosis associated with viral hepatitis.

Laboratory features of autoimmune hepatitis are similar to those seen in chronic viral hepatitis. Liver biochemical tests are invariably abnormal but may not correlate with the clinical severity or histopathologic features in individual cases. Many patients with autoimmune hepatitis have normal serum bilirubin, alkaline phosphatase, and globulin levels with only minimal aminotransferase elevations. Serum AST and ALT levels are increased and fluctuate in the range of 100–1000 units. In severe cases, the serum bilirubin level is moderately elevated (51–171 $\mu\text{mol/L}$ [3–10 mg/dL]). Hypoalbuminemia occurs in patients with very active or advanced disease. Serum alkaline phosphatase levels may be moderately elevated or near normal. In a small proportion of patients, marked elevations of alkaline phosphatase activity occur; in such patients, clinical and laboratory features overlap with those of primary biliary cirrhosis (**Chap. 365**). The prothrombin time is often prolonged, particularly late in the disease or during active phases.

Hypergammaglobulinemia (>2.5 g/dL) is common in autoimmune hepatitis, as is the presence of rheumatoid factor. As noted above, circulating autoantibodies are also prevalent, most characteristically ANAs in a homogeneous staining pattern. Smooth-muscle antibodies are less specific, seen just as frequently in chronic viral hepatitis. Because of the high levels of globulins achieved in the circulation of some patients with autoimmune hepatitis, occasionally the globulins may bind nonspecifically in solid-phase binding immunoassays for viral antibodies. This has been recognized most commonly in tests for antibodies to hepatitis C virus, as noted above. In fact, studies of autoantibodies in autoimmune hepatitis have led to the recognition of new categories of autoimmune hepatitis. *Type I autoimmune hepatitis* is the classic syndrome prevalent in North America and northern Europe occurring in young women, associated with marked hyperglobulinemia, lupoid features, circulating ANAs, and HLA-DR3 or HLA-DR4 (especially *B8-DRB1*03*). Also associated with type I autoimmune hepatitis are autoantibodies against actin and atypical perinuclear antineutrophilic cytoplasmic antibodies (pANCA).

Type II autoimmune hepatitis, often seen in children, more common in Mediterranean populations, and linked to HLA-DRB1 and HLA-DQB1 haplotypes, is associated not with ANA but with anti-LKM. Actually, anti-LKM represent a heterogeneous group of antibodies. In type II autoimmune hepatitis, the antibody is anti-LKM1, directed against cytochrome P450 2D6. This is the same anti-LKM seen in some patients with chronic hepatitis C. Anti-LKM2 is seen in drug-induced hepatitis, and anti-LKM3 (directed against uridine diphosphate glucuronyltransferases) is seen in patients with chronic hepatitis D. Another autoantibody observed in type II autoimmune hepatitis is directed against liver cytosol formiminotransferase cyclo-deaminase (anti-liver cytosol 1). More controversial is whether or not a third category of autoimmune hepatitis exists, *type III autoimmune hepatitis*. These patients lack ANA and anti-LKM1 but have circulating antibodies to soluble liver antigen. Most of these patients are women and have clinical features similar to, perhaps more severe than, those of patients with type I autoimmune hepatitis. Type III autoimmune hepatitis does not appear to represent a distinct category but, instead, is part of the spectrum of type I autoimmune hepatitis; this subcategory has not been adopted by a consensus of international experts.

Liver biopsy abnormalities are similar to those described for chronic viral hepatitis. Expanding portal tracts and extending beyond the plate of periportal hepatocytes into the parenchyma (designated *interface hepatitis* or *piecemeal necrosis*) is a mononuclear cell infiltrate that, in autoimmune hepatitis, may include the presence of plasma cells. Necroinflammatory activity characterizes the lobular parenchyma, and evidence of hepatocellular regeneration is reflected by “rosette” formation, the occurrence of thickened liver cell plates, and regenerative “pseudolobules.” Septal fibrosis, bridging fibrosis, and cirrhosis are frequent. In patients with early autoimmune hepatitis presenting as an acute-hepatitis-like illness, lobular and centrilobular (as opposed to the more common periportal) necrosis has been reported. Bile duct injury and granulomas are uncommon; however,

a subgroup of patients with autoimmune hepatitis has histologic, biochemical, and serologic features overlapping those of primary biliary cirrhosis (Chap. 365).

DIAGNOSTIC CRITERIA

An international group has suggested a set of criteria for establishing a diagnosis of autoimmune hepatitis. Exclusion of liver disease caused by genetic disorders, viral hepatitis, drug hepatotoxicity, and alcohol are linked with such inclusive diagnostic criteria as hyperglobulinemia, autoantibodies, and characteristic histologic features. This international group has also suggested a comprehensive diagnostic scoring system that, rarely required for typical cases, may be helpful when typical features are not present. Factors that weigh in favor of the diagnosis include female gender; predominant aminotransferase elevation; presence and level of globulin elevation; presence of nuclear, smooth muscle, LKM1, and other autoantibodies; concurrent other autoimmune diseases; characteristic histologic features (interface hepatitis, plasma cells, rosettes); HLA-DR3 or -DR4 markers; and response to treatment (see below). A more simplified, more specific scoring system relies on four variables: autoantibodies, serum IgG level, typical or compatible histologic features, and absence of viral hepatitis markers. Weighing against the diagnosis are predominant alkaline phosphatase elevation, mitochondrial antibodies, markers of viral hepatitis, history of hepatotoxic drugs or excessive alcohol, histologic evidence of bile duct injury, or such atypical histologic features as fatty infiltration, iron overload, and viral inclusions.

DIFFERENTIAL DIAGNOSIS

Early during the course of chronic hepatitis, autoimmune hepatitis may resemble typical *acute viral hepatitis* (Chap. 360). Without histologic assessment, severe chronic hepatitis cannot be readily distinguished based on clinical or biochemical criteria from mild chronic hepatitis. In adolescence, *Wilson's disease* (Chaps. 365 and 429) may present with features of chronic hepatitis long before neurologic manifestations become apparent and before the formation of Kayser-Fleischer rings (copper deposition in Descemet's membrane in the periphery of the cornea). In this age group, serum ceruloplasmin and serum and urinary copper determinations plus measurement of liver copper levels establish the correct diagnosis. *Postnecrotic* or *cryptogenic cirrhosis* and *primary biliary cirrhosis* (Chap. 365) share clinical features with autoimmune hepatitis, and both alcoholic hepatitis (Chap. 363) and nonalcoholic steatohepatitis (Chap. 367e) may present with many features common to autoimmune hepatitis; historic, biochemical, serologic, and histologic assessments are usually sufficient to allow these entities to be distinguished from autoimmune hepatitis. Of course, the distinction between autoimmune and chronic viral hepatitis is not always straightforward, especially when viral antibodies occur in patients with autoimmune disease or when autoantibodies occur in patients with viral disease. Furthermore, the presence of extrahepatic features such as arthritis, cutaneous vasculitis, or pleuritis—not to mention the presence of circulating autoantibodies—may cause confusion with *rheumatologic disorders* such as rheumatoid arthritis and systemic lupus erythematosus. The existence of clinical and biochemical features of progressive necroinflammatory liver disease distinguishes chronic hepatitis from these other disorders, which are not associated with severe liver disease. Rarely, hepatic venous outflow obstruction (Budd-Chiari syndrome) may present with features suggestive of autoimmune hepatitis, but painful hepatomegaly, ascites, and vascular imaging provide distinguishing diagnostic clues. Other diagnostic considerations would include celiac disease and ischemic liver disease, which would be readily distinguishable by clinical and laboratory features from autoimmune hepatitis.

Finally, occasionally, features of autoimmune hepatitis overlap with features of autoimmune biliary disorders such as primary biliary cirrhosis, primary sclerosing cholangitis (Chaps. 365 and 369), or, even more rarely, mitochondrial antibody-negative autoimmune cholangitis. Such overlap syndromes are difficult to categorize, and often response to therapy may be the distinguishing factor that establishes the diagnosis.

The mainstay of management in autoimmune hepatitis is glucocorticoid therapy. Several controlled clinical trials have documented that such therapy leads to symptomatic, clinical, biochemical, and histologic improvement as well as increased survival. A therapeutic response can be expected in up to 80% of patients. Unfortunately, therapy has not been shown in clinical trials to prevent ultimate progression to cirrhosis; however, instances of reversal of fibrosis and cirrhosis have been reported in patients responding to treatment, and rapid treatment responses within 1 year do translate into a reduction in progression to cirrhosis. Although some advocate the use of prednisolone (the hepatic metabolite of prednisone), prednisone is just as effective and is favored by most authorities. Therapy may be initiated at 20 mg/d, but a popular regimen in the United States relies on an initiation dose of 60 mg/d. This high dose is tapered successively over the course of a month down to a maintenance level of 20 mg/d. An alternative, but equally effective, approach is to begin with half the prednisone dose (30 mg/d) along with azathioprine (50 mg/d). With azathioprine maintained at 50 mg/d, the prednisone dose is tapered over the course of a month down to a maintenance level of 10 mg/d. The advantage of the combination approach is a reduction, over the span of an 18-month course of therapy, in serious, life-threatening complications of steroid therapy (e.g., cushingoid features, hypertension, diabetes, osteoporosis) from 66% down to under 20%. Genetic analysis for thiopurine S-methyltransferase allelic variants does not correlate with azathioprine-associated cytopenias or efficacy and is not assessed routinely in patients with autoimmune hepatitis. In combination regimens, 6-mercaptopurine may be substituted for its prodrug azathioprine, but this is rarely required. Azathioprine alone, however, is not effective in achieving remission, nor is alternate-day glucocorticoid therapy. Limited experience with budesonide in noncirrhotic patients suggests that this steroid side effect-sparing drug may be effective. Although therapy has been shown to be effective for severe autoimmune hepatitis (AST $\geq 10 \times$ the upper limit of normal or $\geq 5 \times$ the upper limit of normal in conjunction with serum globulin greater than or equal to twice normal; bridging necrosis or multilobular necrosis on liver biopsy; presence of symptoms), therapy is not indicated for mild forms of chronic hepatitis, and the efficacy of therapy in mild or asymptomatic autoimmune hepatitis has not been established.

Improvement of fatigue, anorexia, malaise, and jaundice tends to occur within days to several weeks; biochemical improvement occurs over the course of several weeks to months, with a fall in serum bilirubin and globulin levels and an increase in serum albumin. Serum aminotransferase levels usually drop promptly, but improvements in AST and ALT alone do not appear to be reliable markers of recovery in individual patients; histologic improvement, characterized by a decrease in mononuclear infiltration and in hepatocellular necrosis, may be delayed for 6–24 months. Still, if interpreted cautiously, aminotransferase levels are valuable indicators of relative disease activity, and many authorities do not advocate for serial liver biopsies to assess therapeutic success or to guide decisions to alter or stop therapy. Rapidity of response is more common in older patients (≥ 69 years) and those with HLA *DBR1*04*; although rapid responders may progress less slowly to cirrhosis and liver transplantation, they are no less likely than slower responders to relapse after therapy. Therapy should continue for at least 12–18 months. After tapering and cessation of therapy, the likelihood of relapse is at least 50%, even if posttreatment histology has improved to show mild chronic hepatitis, and the majority of patients require therapy at maintenance doses indefinitely. Continuing azathioprine alone (2 mg/kg body weight daily) after cessation of prednisone therapy has been shown to reduce the frequency of relapse. Long-term maintenance with low-dose prednisone (≤ 10 mg daily) has also been shown to keep autoimmune hepatitis in check, but maintenance azathioprine is more effective in maintaining remission.

In medically refractory cases, an attempt should be made to intensify treatment with high-dose glucocorticoid monotherapy (60 mg daily) or combination glucocorticoid (30 mg daily) plus high-dose azathioprine (150 mg daily) therapy. After a month, doses of prednisone can be reduced by 10 mg a month, and doses of azathioprine can be reduced by 50 mg a month toward ultimate, conventional maintenance doses. Patients refractory to this regimen may be treated with cyclosporine, tacrolimus, or mycophenolate mofetil; however, to date, only limited anecdotal reports support these approaches. If medical therapy fails, or when chronic hepatitis progresses to cirrhosis and is associated with life-threatening complications of liver decompensation, liver transplantation is the only recourse (**Chap. 368**); failure of the bilirubin to improve after 2 weeks of therapy should prompt early consideration of the patient for liver transplantation. Recurrence of autoimmune hepatitis in the new liver occurs rarely in most experiences but in as many as 35–40% of cases in others.

Like all patients with chronic liver disease, patients with autoimmune hepatitis should be vaccinated against hepatitis A and B, ideally before immunosuppressive therapy is begun, if practical.

ACKNOWLEDGMENT

Kurt J. Isselbacher, MD, contributed to this chapter in previous editions of *Harrison's*.

363 Alcoholic Liver Disease

Mark E. Mailliard, Michael F. Sorrell

Chronic and excessive alcohol ingestion is one of the major causes of liver disease. The pathology of alcoholic liver disease consists of three major lesions, with the progressive injury rarely existing in a pure form: (1) fatty liver, (2) alcoholic hepatitis, and (3) cirrhosis. Fatty liver is present in >90% of daily as well as binge drinkers. A much smaller percentage of heavy drinkers will progress to alcoholic hepatitis, thought to be a precursor to cirrhosis. The prognosis of severe alcoholic liver disease is dismal; the mortality of patients with alcoholic hepatitis concurrent with cirrhosis is nearly 60% at 4 years. Although alcohol is considered a direct hepatotoxin, only between 10 and 20% of alcoholics will develop alcoholic hepatitis. The explanation for this apparent paradox is unclear but involves the complex interaction of facilitating factors, such as drinking patterns, diet, obesity, and gender. There are no diagnostic tools that can predict individual susceptibility to alcoholic liver disease.

GLOBAL CONSIDERATIONS



Alcohol is the world's third largest risk factor for disease burden. The harmful use of alcohol results in 2.5 million deaths each year. Most of the mortality attributed to alcohol is secondary to cirrhosis. Mortality from cirrhosis is declining in most Western countries, concurrent with a reduction in alcohol consumption, with the exceptions of the United Kingdom, Russia, Romania, and Hungary. These increases in cirrhosis and its complications are closely correlated with increased volume of alcohol consumed per capita population and are regardless of gender.

ETIOLOGY AND PATHOGENESIS

Quantity and duration of alcohol intake are the most important risk factors involved in the development of alcoholic liver disease (**Table 363-1**). The roles of beverage type(s), i.e. wine, beer, or spirits, and pattern of drinking (daily versus binge drinking) are less clear. Progress beyond the fatty liver stage seems to require additional risk

TABLE 363-1 RISK FACTORS FOR ALCOHOLIC LIVER DISEASE

Risk Factor	Comment
Quantity	In men, 40–80 g/d of ethanol produces fatty liver; 160 g/d for 10–20 years causes hepatitis or cirrhosis. Only 15% of alcoholics develop alcoholic liver disease.
Gender	Women exhibit increased susceptibility to alcoholic liver disease at amounts >20 g/d; two drinks per day is probably safe.
Hepatitis C	HCV infection concurrent with alcoholic liver disease is associated with younger age for severity, more advanced histology, and decreased survival.
Genetics	Patatin-like phospholipase domain-containing protein 3 (PNPLA3) has been associated with alcoholic cirrhosis.
Fatty liver	Alcohol injury does not require malnutrition, but obesity and nonalcoholic fatty liver are risk factors. Patients should receive vigorous attention to nutritional support.

factors that remain incompletely defined. Although there are genetic predispositions for alcoholism (**Chap. 467**), gender is a strong determinant for alcoholic liver disease. Women are more susceptible to alcoholic liver injury when compared to men. They develop advanced liver disease with substantially less alcohol intake. In general, the time it takes to develop liver disease is directly related to the amount of alcohol consumed. It is useful in estimating alcohol consumption to understand that one beer, four ounces of wine, or one ounce of 80% spirits all contain ~12 g of alcohol. The threshold for developing alcoholic liver disease is higher in men, while women are at increased risk for developing similar degrees of liver injury by consuming significantly less. Gender-dependent differences result from poorly understood effects of estrogen, proportion of body fat, and the gastric metabolism of alcohol. Obesity, a high-fat diet, and the protective effect of coffee have been postulated to play a part in the development of the pathogenic process.

Chronic infection with hepatitis C virus (HCV) (**Chap. 362**) is an important comorbidity in the progression of alcoholic liver disease to cirrhosis in chronic and excessive drinkers. Even moderate alcohol intake of 20–50 g/d increases the risk of cirrhosis and hepatocellular cancer in HCV-infected individuals. Patients with both alcoholic liver injury and HCV infection develop decompensated liver disease at a younger age and have poorer overall survival. Increased liver iron stores and, rarely, porphyria cutanea tarda can occur as a consequence of the overlapping injurious processes secondary to alcohol abuse and HCV infection. In addition, alcohol intake of >50 g/d by HCV-infected patients decreases the efficacy of interferon-based antiviral therapy.

The pathogenesis of alcoholic liver injury is unclear. The present conceptual foundation is that alcohol acts as a direct hepatotoxin and that malnutrition does not have a major role. Ingestion of alcohol initiates an inflammatory cascade by its metabolism to acetaldehyde, resulting in a variety of metabolic responses. Steatosis from lipogenesis, fatty acid synthesis, and depression of fatty acid oxidation appears secondary to effects on sterol regulatory transcription factor and peroxisome proliferator-activated receptor α (PPAR- α). Intestinal-derived endotoxin initiates a pathogenic process through toll-like receptor 4 and tumor necrosis factor α (TNF- α) that facilitates hepatocyte apoptosis and necrosis. The cell injury and endotoxin release initiated by ethanol and its metabolites also activate innate and adaptive immunity pathways releasing proinflammatory cytokines (e.g., TNF- α), chemokines, and proliferation of T and B cells. The production of toxic protein-aldehyde adducts, generation of reducing equivalents, and oxidative stress also contribute to the liver injury. Hepatocyte injury and impaired regeneration following chronic alcohol ingestion are ultimately associated with stellate cell activation and collagen production, which are key events in fibrogenesis. The resulting fibrosis from

continuing alcohol use determines the architectural derangement of the liver and associated pathophysiology.

PATHOLOGY

The liver has a limited repertoire in response to injury. Fatty liver is the initial and most common histologic response to hepatotoxic stimuli, including excessive alcohol ingestion. The accumulation of fat within the perivenular hepatocytes coincides with the location of alcohol dehydrogenase, the major enzyme responsible for alcohol metabolism. Continuing alcohol ingestion results in fat accumulation throughout the entire hepatic lobule. Despite extensive fatty change and distortion of the hepatocytes with macrovesicular fat, the cessation of drinking results in normalization of hepatic architecture and fat content. Alcoholic fatty liver has traditionally been regarded as entirely benign, but similar to the spectrum of nonalcoholic fatty liver disease (Chap. 367e), the appearance of steatohepatitis and certain pathologic features such as giant mitochondria, perivenular fibrosis, and macrovesicular fat may be associated with progressive liver injury.

The transition between fatty liver and the development of alcoholic hepatitis is blurred. The hallmark of alcoholic hepatitis is hepatocyte injury characterized by ballooning degeneration, spotty necrosis, polymorphonuclear infiltrate, and fibrosis in the perivenular and perisinusoidal space of Disse. Mallory-Denk bodies are often present in florid cases but are neither specific nor necessary to establish the diagnosis. Alcoholic hepatitis is thought to be a precursor to the development of cirrhosis. However, like fatty liver, it is potentially reversible with cessation of drinking. Cirrhosis is present in up to 50% of patients with biopsy-proven alcoholic hepatitis, and its regression is uncertain, even with abstinence.

CLINICAL FEATURES

The clinical manifestations of alcoholic fatty liver are subtle and characteristically detected as a consequence of the patient's visit for a seemingly unrelated matter. Previously unsuspected hepatomegaly is often the only clinical finding. Occasionally, patients with fatty liver will present with right upper quadrant discomfort, nausea, and, rarely, jaundice. Differentiation of alcoholic fatty liver from nonalcoholic fatty liver is difficult unless an accurate drinking history is ascertained. In every instance where liver disease is present, a thoughtful and sensitive drinking history should be obtained. Standard, validated questions accurately detect alcohol-related problems (Chap. 467). Alcoholic hepatitis is associated with a wide gamut of clinical features. Fever, spider nevi, jaundice, and abdominal pain simulating an acute abdomen represent the extreme end of the spectrum, while many patients will be entirely asymptomatic. Portal hypertension, ascites, or variceal bleeding can occur in the absence of cirrhosis. Recognition of the clinical features of alcoholic hepatitis is central to the initiation of an effective and appropriate diagnostic and therapeutic strategy. It is important to recognize that patients with alcoholic cirrhosis often exhibit clinical features identical to other causes of cirrhosis.

LABORATORY FEATURES

Patients with alcoholic liver disease are often identified through routine screening tests. The typical laboratory abnormalities seen in fatty liver are nonspecific and include modest elevations of aspartate aminotransferase (AST), alanine aminotransferase (ALT), and γ -glutamyl transpeptidase (GGTP), often accompanied by hypertriglyceridemia and hyperbilirubinemia. In alcoholic hepatitis and in contrast to other causes of fatty liver, AST and ALT are usually elevated two- to sevenfold. They are rarely >400 IU, and the AST/ALT ratio is >1 (Table 363-2). Hyperbilirubinemia is accompanied by modest increases in the alkaline phosphatase level. Derangement in hepatocyte synthetic function indicates more serious disease. Hypoalbuminemia and coagulopathy are common in advanced liver injury. Ultrasonography is useful in detecting fatty infiltration of the liver and determining liver size. The demonstration by ultrasound of portal vein flow reversal, ascites, and intraabdominal venous collaterals indicates serious liver injury with less potential for complete reversal.

TABLE 363-2 LABORATORY DIAGNOSIS OF ALCOHOLIC FATTY LIVER AND ALCOHOLIC HEPATITIS

Test	Comment
AST	Increased two- to sevenfold, <400 IU/L, greater than ALT
ALT	Increased two- to sevenfold, <400 IU/L
AST/ALT	Usually >1
GGTP	Not specific to alcohol, easily inducible, elevated in all forms of fatty liver
Bilirubin	May be markedly increased in alcoholic hepatitis despite modest elevation in alkaline phosphatase

Abbreviations: ALT, alanine aminotransferase; AST, aspartate aminotransferase; GGTP, γ -glutamyl transpeptidase.

PROGNOSIS

Critically ill patients with alcoholic hepatitis have short-term (30-day) mortality rates $>50\%$. Severe alcoholic hepatitis is heralded by coagulopathy (prothrombin time increased >5 s), anemia, serum albumin concentrations <25 g/L (2.5 mg/dL), serum bilirubin levels >137 $\mu\text{mol/L}$ (8 mg/dL), renal failure, and ascites. A discriminant function calculated as $4.6 \times$ (the prolongation of the prothrombin time above control [seconds]) + serum bilirubin (mg/dL) can identify patients with a poor prognosis (discriminant function >32). A Model for End-Stage Liver Disease (MELD) score (Chap. 368) ≥ 21 also is associated with significant mortality in alcoholic hepatitis. The presence of ascites, variceal hemorrhage, deep encephalopathy, or hepatorenal syndrome predicts a dismal prognosis. The pathologic stage of the injury can be helpful in predicting prognosis. Liver biopsy should be performed whenever possible to establish the diagnosis and to guide the therapeutic decisions.

TREATMENT ALCOHOLIC LIVER DISEASE

Complete abstinence from alcohol is the cornerstone in the treatment of alcoholic liver disease. Improved survival and the potential for reversal of histologic injury regardless of the initial clinical presentation are associated with total avoidance of alcohol ingestion. Referral of patients to experienced alcohol counselors and/or alcohol treatment programs should be routine in the management of patients with alcoholic liver disease. Attention should be directed to the nutritional and psychosocial states during the evaluation and treatment periods. Because of data suggesting that the pathogenic mechanisms in alcoholic hepatitis involve cytokine release and the perpetuation of injury by immunologic processes, glucocorticoids have been extensively evaluated in the treatment of alcoholic hepatitis. Patients with severe alcoholic hepatitis, defined as a discriminant function >32 or MELD >20 , should be given prednisone, 40 mg/d, or prednisolone, 32 mg/d, for 4 weeks, followed by a steroid taper (Fig. 363-1). Exclusion criteria include active gastrointestinal bleeding, renal failure, or pancreatitis. Women with encephalopathy from severe alcoholic hepatitis may be particularly good candidates for glucocorticoids. A Lille score >0.45 , at <http://www.lillemodel.com>, uses pretreatment variables plus the change in total bilirubin at day 7 of glucocorticoids to identify patients unresponsive to therapy.

The role of TNF- α expression and receptor activity in alcoholic liver injury has led to an examination of TNF inhibition as an alternative to glucocorticoids for severe alcoholic hepatitis. The nonspecific TNF inhibitor, pentoxifylline, demonstrated improved survival in the therapy of severe alcoholic hepatitis, primarily due to a decrease in hepatorenal syndrome (Fig. 363-2). Monoclonal antibodies that neutralize serum TNF- α should not be used in alcoholic hepatitis because of studies reporting increased deaths secondary to infection and renal failure.

Liver transplantation is an accepted indication for treatment in selected and motivated patients with end-stage cirrhosis. Outcomes are equal or superior to other indications for transplantation. In general, transplant candidacy should be reevaluated after a defined

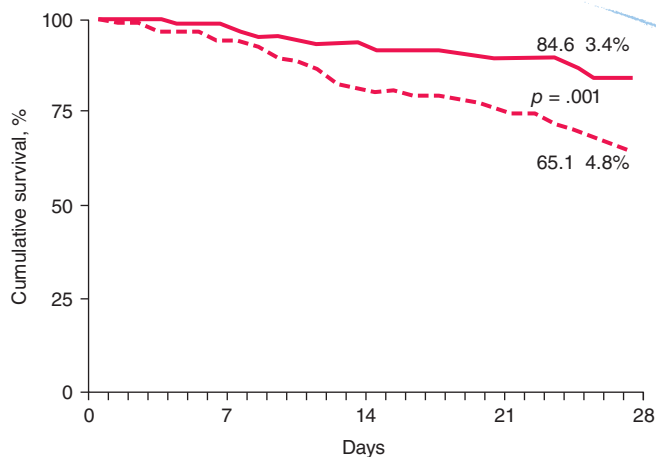


FIGURE 363-1 Effect of glucocorticoid therapy of severe alcoholic hepatitis on short-term survival: the result of a meta-analysis of individual data from three studies. Prednisolone, solid line; placebo, dotted line. (Adapted from P Mathurin et al: *J Hepatol* 36:480, 2002, with permission from Elsevier Science.)

period of sobriety. Patients presenting with alcoholic hepatitis have been largely excluded from transplant candidacy because of the perceived risk of increased surgical mortality and high rates of recidivism following transplantation. Recently, a European multidisciplinary group has reported excellent long-term transplant outcomes in highly selected patients with florid alcoholic hepatitis. General application of transplantation in such patients must await confirmatory outcomes by others.

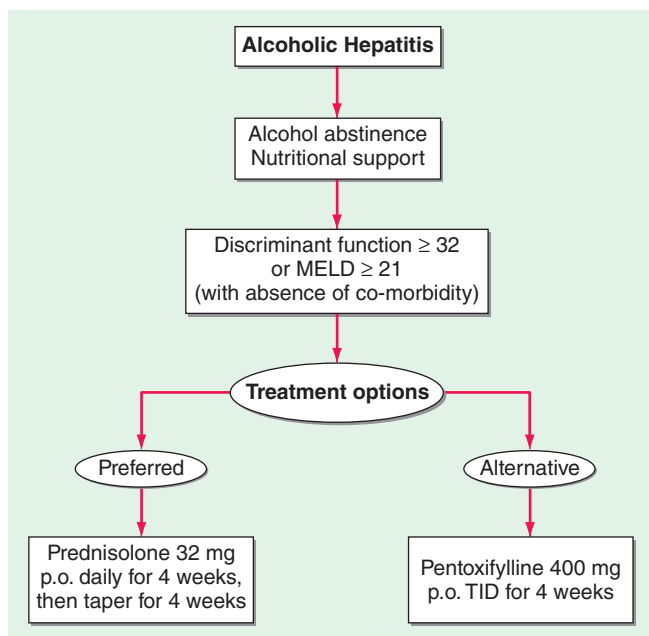


FIGURE 363-2 Treatment algorithm for alcoholic hepatitis. As identified by a calculated discriminant function >32 (see text), patients with severe alcoholic hepatitis, without the presence of gastrointestinal bleeding or infection, would be candidates for either glucocorticoids or pentoxifylline administration.

364 Nonalcoholic Fatty Liver Diseases and Nonalcoholic Steatohepatitis

Manal F. Abdelmalek, Anna Mae Diehl

INCIDENCE, PREVALENCE, AND NATURAL HISTORY

Nonalcoholic fatty liver disease (NAFLD) is the most common chronic liver disease in many parts of the world, including the United States. Population-based abdominal imaging studies have demonstrated fatty liver in at least 25% of American adults. Because the vast majority of these subjects deny hazardous levels of alcohol consumption (defined as greater than one drink per day in women or two drinks per day in men), they are considered to have NAFLD. NAFLD is strongly associated with overweight/obesity and insulin resistance. However, it can also occur in lean individuals and is particularly common in those with a paucity of adipose depots (i.e., lipodystrophy). Ethnic/racial factors also appear to influence liver fat accumulation; the documented prevalence of NAFLD is lowest in African Americans (~25%), highest in Americans of Hispanic ancestry (~50%), and intermediate in American whites (~33%).

NAFLD encompasses a spectrum of liver pathology with different clinical prognoses. The simple accumulation of triglyceride within hepatocytes (hepatic steatosis) is on the most clinically benign extreme of the spectrum. On the opposite, most clinically ominous extreme, are cirrhosis (Chap. 365) and primary liver cancer (Chap. 111). The risk of developing cirrhosis is extremely low in individuals with chronic hepatic steatosis, but increases as steatosis becomes complicated by histologically conspicuous hepatocyte death and inflammation (i.e., nonalcoholic steatohepatitis [NASH]). NASH itself is also a heterogeneous condition; sometimes it improves to steatosis or normal histology, sometimes it remains relatively stable for years, but sometimes it results in progressive accumulation of fibrous scar that eventuates in cirrhosis. Once NAFLD-related cirrhosis develops, the annual incidence of primary liver cancer is 1%.

Abdominal imaging is not able to determine which individuals with NAFLD have associated liver cell death and inflammation (i.e., NASH), and specific blood tests to diagnose NASH are not yet available. However, population-based studies that have used elevated serum ALT as a marker of liver injury indicate that about 6–8% of American adults have serum ALT elevations that cannot be explained by excessive alcohol consumption, other known causes of fatty liver disease (Table 364-1), viral hepatitis, or drug-induced or congenital liver diseases. Because the prevalence of such “cryptogenic” ALT elevations increases with body mass index, it is presumed that they are due to NASH. Hence, at any given point in time, NASH is present in about 25% of individuals who have NAFLD (i.e., about 6% of the general U.S. adult population has NASH). Smaller cross-sectional studies in which liver biopsies have been performed on NASH patients at tertiary referral centers consistently demonstrate advanced fibrosis or cirrhosis in about 25% of those cohorts. By extrapolation, therefore, cirrhosis develops in about 6% of individuals with NAFLD (i.e., in about 1.5–2% of the general U.S. population). The risk for advanced liver fibrosis is highest in individuals with NASH who are older than 45–50 years of age and overweight/obese or afflicted with type 2 diabetes.

To put these data in perspective, it is helpful to recall that the prevalence of hepatitis C–related cirrhosis in the United States is about 0.5%. Thus, NAFLD-related cirrhosis is about three to four times more common than cirrhosis caused by chronic hepatitis C infection. Consistent with these data, experts have predicted that NAFLD will surpass hepatitis C as the leading indication for liver transplantation in the United States within the next decade. Similar to cirrhosis caused by other liver diseases, cirrhosis caused by NAFLD increases the risk for primary liver cancer. Both hepatocellular carcinoma and intrahepatic cholangiocarcinoma (ICC) have also been reported to occur in NAFLD patients without cirrhosis, suggesting that NAFLD per se may be a premalignant condition. NAFLD, NASH, and NAFLD-related cirrhosis are not limited to adults. All have been well documented in children. As in adults, obesity and insulin resistance are the main risk

TABLE 364-1 ALTERNATIVE CAUSES OF HEPATIC STEATOSIS

- Alcoholic liver disease
- Hepatitis C (particularly genotype 3)
- Inborn errors of metabolism
 - Abetalipoproteinemia
 - Cholesterol ester storage disease
 - Galactosemia
 - Glycogen storage disease
 - Hereditary fructose intolerance
 - Homocystinuria
 - Systemic carnitine deficiency
 - Tyrosinemia
 - Weber-Christian syndrome
 - Wilson's disease
 - Wolman's disease
- Medications (see [Table 364-2](#))
- Miscellaneous
 - Industrial exposure to petrochemical
 - Inflammatory bowel disease
 - Lipodystrophy
 - Bacterial overgrowth
 - Starvation
 - Parenteral nutrition
- Surgical procedures
 - Bilopancreatic diversion
 - Extensive small-bowel resection
 - Gastric bypass
 - Jejunioleal bypass
- Reye's syndrome
- Acute fatty liver of pregnancy
- HELLP syndrome (*hemolytic anemia, elevated liver enzymes, low platelet count*)

factors for pediatric NAFLD. Thus, the rising incidence and prevalence of childhood obesity suggests that NAFLD is likely to become an even greater contributor to society's burden of liver disease in the future.

PATHOGENESIS

The mechanisms underlying the pathogenesis and progression of NAFLD are not entirely clear. The best-understood mechanisms pertain to hepatic steatosis. This is proven to result when hepatocyte mechanisms for triglyceride synthesis (e.g., lipid uptake and *de novo* lipogenesis) overwhelm mechanisms for triglyceride disposal (e.g., degradative metabolism and lipoprotein export), leading to accumulation of fat (i.e., triglyceride) within hepatocytes. Obesity stimulates hepatocyte triglyceride accumulation by altering the intestinal microbiota to enhance both energy harvest from dietary sources and intestinal permeability. Reduced intestinal barrier function increases hepatic exposure to gut-derived products, which stimulate liver cells to generate inflammatory mediators that inhibit insulin actions. Obese adipose depots also produce excessive soluble factors (adipokines) that inhibit tissue insulin sensitivity. Insulin resistance promotes hyperglycemia. This drives the pancreas to produce more insulin to maintain glucose homeostasis. However, hyperinsulinemia also promotes lipid uptake, fat synthesis, and fat storage. The net result is hepatic triglyceride accumulation (i.e., steatosis).

Triglyceride *per se* is not hepatotoxic. However, its precursors (e.g., fatty acids and diacylglycerols) and metabolic by-products (e.g., reactive oxygen species) may damage hepatocytes, leading to hepatocyte lipotoxicity. Lipotoxicity also triggers the generation of other factors (e.g., inflammatory cytokines, hormonal mediators) that deregulate systems that normally maintain hepatocyte viability. The net result is increased hepatocyte death. Dying hepatocytes, in turn, release various factors that trigger wound healing responses that aim to

replace (regenerate) lost hepatocytes. Such repair involves transient expansion of other cell types, such as myofibroblasts and progenitor cells, that make and degrade matrix, remodel the vasculature, and generate replacement hepatocytes, as well as the recruitment of immune cells that release factors that modulate liver injury and repair. NASH is the morphologic manifestation of lipotoxicity and resultant wound healing responses. Because the severity and duration of lipotoxic liver injury dictate the intensity and duration of repair, the histologic features and outcomes of NASH are variable. Cirrhosis and liver cancer are potential outcomes of chronic NASH. Cirrhosis results from futile repair, i.e., progressive accumulation of wound healing cells, fibrous matrix, and abnormal vasculature (scarring), rather than efficient reconstruction/regeneration of healthy hepatic parenchyma. Primary liver cancers develop when malignantly transformed liver cells escape mechanisms that normally control regenerative growth. The mechanisms responsible for futile repair (cirrhosis) and liver carcinogenesis are not well understood. Because normal liver regeneration is a very complex process, there are multiple opportunities for deregulation and, thus, pathogenic heterogeneity. To date, this heterogeneity has confounded development of both diagnostic tests and treatments for defective/deregulated liver repair (i.e., cirrhosis and cancer). Hence, current strategies focus on circumventing misrepair by preventing and/or reducing lipotoxic liver injury.

DIAGNOSIS

Diagnosing NAFLD requires demonstration of increased liver fat in the absence of hazardous levels of alcohol consumption. Thresholds for potentially dangerous alcohol ingestion have been set at more than one drink per day in women and two drinks per day in men based on epidemiologic evidence that the prevalence of serum aminotransferase elevations increases when alcohol consumption habitually exceeds these levels. In those studies, one drink was defined as having 10 g of ethanol and, thus, is equivalent to one can of beer, 4 ounces of wine, or 1.5 ounces (one shot) of distilled spirits. Other causes of liver fat accumulation (particularly exposure to certain drugs; [Table 364-2](#)) and liver injury (e.g., viral hepatitis, autoimmune liver disease, iron or copper overload, α_1 antitrypsin deficiency) must also be excluded. Thus, establishing the diagnosis of NAFLD does not require invasive testing: it can be accomplished by history and physical examination, liver imaging (ultrasound is an acceptable first-line test; computed tomography [CT] or magnetic resonance imaging [MRI] enhances sensitivity for liver fat detection but adds expense), and blood tests to exclude other liver diseases. It is important to emphasize that the liver may not be enlarged, and serum aminotransferases and liver function tests (e.g., bilirubin, albumin, prothrombin time) may be completely normal, in individuals with NAFLD. Because there is yet no one specific blood test for NAFLD, confidence in the diagnosis of NAFLD is increased by identification of NAFLD risk factors. The latter include increased body mass index, insulin resistance/type 2 diabetes mellitus, and other parameters indicative of the metabolic syndrome (e.g., systemic hypertension, dyslipidemia, hyperuricemia/gout, cardiovascular disease; [Chap. 422](#)) in the patient or family members.

Establishing the severity of NAFLD-related liver injury and related scarring (i.e., staging NAFLD) is more difficult than simply diagnosing NAFLD. Staging is critically important, however, because it is necessary to define prognosis and thereby determine treatment recommendations. The goal of staging is to distinguish patients with NASH from those with simple steatosis and to identify which of the NASH patients have advanced fibrosis. The 10-year probability of developing liver-related morbidity or mortality in steatosis is negligible, and hence, this subgroup of NAFLD patients tends to be managed conservatively (see below). In contrast, more intensive follow-up and therapy are justified in NASH patients, and the subgroup with advanced fibrosis merits the most intensive scrutiny and intervention because their 10-year risk of liver-related morbidity and mortality is clearly increased.

Staging approaches can be separated into noninvasive testing (i.e., blood testing, physical examination, and imaging) and invasive approaches (i.e., liver biopsy). Blood test evidence of hepatic dysfunction (e.g., hyperbilirubinemia, hypoalbuminemia, prothrombin time

- Cytotoxic and cytostatic drugs
 - L-Asparaginase
 - Azacitidine
 - Azaserine
 - Bleomycin
 - Methotrexate
 - Puromycin
 - Tetracycline
 - Doxycycline
- Metals
 - Antimony
 - Barium salts
 - Chromates
 - Phosphorus
 - Rare earths of low atomic number
 - Thallium compounds
 - Uranium compounds
- Other drugs and toxins
 - Amiodarone
 - 4,4'-Diethylaminoethoxyhexesterol
 - Ethionine
 - Ethyl bromide
 - Estrogens
 - Glucocorticoids
 - Highly active antiretroviral therapy
 - Hydralazine
 - Hypoglycin
 - Orotate
 - Perhexiline maleate
 - Safrole
 - Tamoxifen

prolongation) or portal hypertension (e.g., thrombocytopenia) and stigmata of portal hypertension on physical examination (e.g., spider angiomas, palmar erythema, splenomegaly, ascites, clubbing, encephalopathy) suggest a diagnosis of advanced NAFLD. Currently, however, liver biopsy is the gold standard for establishing the severity of liver injury and fibrosis because it is both more sensitive and specific than these other tests for establishing NAFLD severity. Although invasive, liver biopsy is seldom complicated by serious adverse sequelae such as significant bleeding, pain, or inadvertent puncture of other organs and thus is relatively safe. However, biopsy suffers from potential sampling error unless tissue cores of 2 cm or longer are acquired. Also, examination of tissue at a single point in time is not reliable for determining whether the pathologic processes are progressing or regressing. The risk of serial liver biopsies within short time intervals is generally deemed as unacceptable outside of research studies. These limitations of liver biopsy have stimulated efforts to develop noninvasive approaches to stage NAFLD. As is true for many other types of chronic liver disease, in NAFLD the levels of serum aminotransferases (aspartate aminotransferase [AST] and alanine aminotransferase [ALT]) do not reliably reflect the severity of liver cell injury, extent of liver cell death, or related liver inflammation and fibrosis. Thus, they are imperfect for determining which individuals with NAFLD have NASH. This has stimulated research to identify superior markers of liver injury. Serum levels of keratin 8 and keratin 18 appear to be promising surrogates. Keratins 8 and 18 (K8/18) are epithelial cytoskeletal proteins that undergo cleavage during programmed cell death (apoptosis). Both cleaved and full-length K8/18 are released into the blood as hepatocytes die, and studies suggest that serum levels of K8/18 differentiate individuals with NASH from those with simple steatosis or normal livers more reliably than do serum aminotransferase levels. Moreover, K8/18 levels appear to parallel the severity of liver fibrosis, with higher levels marking individuals who are likely to have worse

scarring (i.e., advanced liver fibrosis or cirrhosis). While promising, testing for K8/18 has not yet become standard clinical practice. Other blood tests and imaging approaches that quantify liver fibrosis are also being developed. Recently, the U.S. Food and Drug Administration (FDA) approved an ultrasound-based test that measures liver stiffness as a surrogate marker of fibrosis (FibroScan®) (Chap. 358). This new tool will likely be used serially to monitor fibrosis progression and regression in NAFLD patients. Studies that compare the receiver operator characteristics of K8/18 plus FibroScan® versus liver biopsy for monitoring NAFLD evolution are forthcoming.

CLINICAL FEATURES OF NAFLD

Most subjects with NAFLD are asymptomatic. The diagnosis is often made when abnormal liver aminotransferases or features of fatty liver are noted during an evaluation performed for other reasons. NAFLD may also be diagnosed during the workup of vague right upper quadrant abdominal pain, hepatomegaly, or an abnormal-appearing liver at time of abdominal surgery. Obesity is present in 50–90% of subjects. Most patients with NAFLD also have other features of the metabolic syndrome (Chap. 422). Some have subtle stigmata of chronic liver disease, such as spider angiomas, palmer erythema, or splenomegaly. In a small minority of patients with advanced NAFLD, complications of end-stage liver disease (e.g., jaundice, features of portal hypertension such as ascites or variceal hemorrhage) may be the initial findings.

The association of NAFLD with obesity, diabetes, hypertriglyceridemia, hypertension, and cardiovascular disease is well known. Other associations include chronic fatigue, mood alterations, obstructive sleep apnea, thyroid dysfunction, and chronic pain syndrome. NAFLD is an independent risk factor for metabolic syndrome (Chap. 422). Longitudinal studies suggest that patients with NASH are at two- to threefold increased risk for the development of metabolic syndrome. Similarly, studies have shown that patients with NASH have a higher risk for the development of hypertension and diabetes mellitus. The presence of NAFLD is also independently associated with endothelial dysfunction, increased carotid intimal thickness, and the number of plaques in carotid and coronary arteries. Such data indicate that NAFLD has many deleterious effects on health in general.

TREATMENT OF NAFLD

Treatment of NAFLD can be divided into three components: (1) specific therapy of NAFLD-related liver disease; (2) treatment of NAFLD-associated comorbidities; and (3) treatment of the complications of advanced NAFLD. The subsequent discussion focuses on specific therapies for NAFLD, with some mention of their impact on major NAFLD comorbidities (insulin resistance/diabetes, obesity, and dyslipidemia). Treatment of the complications of advanced NAFLD involves management of the complications of cirrhosis and portal hypertension, including primary liver cancers. Approaches to accomplish these objectives are similar to those used in other chronic liver diseases and are covered elsewhere in the textbook (Chaps. 365 and 111).

At present, there are no FDA-approved therapies for the treatment of NAFLD. Thus, the current approach to NAFLD management focuses on treatment to improve the risk factors for NASH (i.e., obesity, insulin resistance, metabolic syndrome, dyslipidemia). Based on our understanding of the natural history of NAFLD, only patients with NASH or those with features of hepatic fibrosis on liver biopsy are considered currently for targeted pharmacologic therapies. This approach may change as our understanding of disease pathophysiology improves and potential targets of therapy evolve.

Diet and Exercise Lifestyle changes and dietary modification are the foundation for NAFLD treatment. Many studies indicate that lifestyle modification can improve serum aminotransferases and hepatic steatosis, with loss of at least 3–5% of body weight improving steatosis, but greater weight loss (up to 10%) necessary to improve steatohepatitis. The benefits of different dietary macronutrient contents (e.g., low-carbohydrate vs low-fat diets, saturated vs unsaturated fat diets) and different intensities of calorie restriction appear to be comparable. In adults with NAFLD, exercise regimens that improve fitness may be sufficient to reduce hepatic steatosis, but their impact on other aspects

of liver histology remains unknown. Unfortunately, most NAFLD patients are unable to achieve sustained weight loss. Although pharmacologic therapies such as orlistat, topiramate, and phentermine to facilitate weight loss are available, their role in the treatment of NAFLD remains experimental.

Pharmacologic Therapies Several drug therapies have been tried in both research and clinical settings. No agent has yet been approved by the FDA for the treatment of NAFLD. Hence, this remains an area of active research. Because NAFLD is strongly associated with the metabolic syndrome and type 2 diabetes (Chaps. 417 and 418), the efficacy of various insulin-sensitizing agents has been examined. *Metformin*, an agent that mainly improves hepatic insulin sensitivity, has been evaluated in several small, open-label studies in adults and a recent larger, prospectively randomized trial in children (dubbed the TONIC study). Although several of the adult NASH studies suggested improvements in aminotransferases and/or liver histology, metformin did not improve liver histology in the TONIC study of children with NASH. Thus, it is not currently recommended as a treatment for NASH. Uncontrolled open-label studies have also investigated *thiazolidinediones* (*pioglitazone* and *rosiglitazone*) in adults with NASH. This class of drugs is known to improve systemic insulin resistance. Both pioglitazone and rosiglitazone reduced aminotransferases and improved some of the histologic features of NASH in small, uncontrolled studies. A large, National Institutes of Health–sponsored, randomized placebo-controlled clinical trial, the PIVENs Study (Pioglitazone vs Vitamin E vs Placebo for the Treatment of 247 Nondiabetic Adults with NASH), demonstrated that resolution of histologic NASH occurred more often in subjects treated with pioglitazone (30 mg/d) than with placebo for 18 months (47 vs 21%, $p = .001$). However, many subjects in the pioglitazone group gained weight, and liver fibrosis did not improve. Also, it should be noted that the long-term safety and efficacy of thiazolidinediones in patients with NASH has not been established. Five-year follow-up of subjects treated with rosiglitazone demonstrated no reduction in liver fibrosis, and rosiglitazone has been associated with increased long-term risk for cardiovascular mortality. Hence, it is not recommended as a treatment for NAFLD. Pioglitazone may be safer because in a recent large meta-analysis it was associated with reduced overall mortality, myocardial infarction, and stroke. However, caution must be exercised when considering its use in patients with impaired myocardial function.

Antioxidants have also been evaluated for the treatment of NAFLD because oxidant stress is thought to contribute to the pathogenesis of NASH. *Vitamin E*, an inexpensive yet potent antioxidant, has been examined in several small pediatric and adult studies with varying results. In all of those studies, vitamin E was well tolerated, and most showed modest improvements in aminotransferase levels, radiographic features of hepatic steatosis, and/or histologic features of NASH. Vitamin E (800 IU/d) was also compared to placebo in the PIVENs and TONIC studies. In PIVENs, vitamin E was the only agent that achieved the predetermined primary endpoint (i.e., improvement in steatohepatitis, lobular inflammation, and steatosis score, without an increase in the fibrosis score). This endpoint was met in 43% of patients in the vitamin E group ($p = .001$ vs placebo), 34% in the pioglitazone group ($p = .04$ vs placebo), and 19% in the placebo group. Vitamin E also improved NASH histology in pediatric patients with NASH (TONIC trial). However, a recent population-based study suggested that chronic vitamin E therapy may increase the risk for cardiovascular mortality. Thus, vitamin E should only be considered as a first-line pharmacotherapy for nondiabetic NASH patients. Also, given its potentially negative effects on cardiovascular health, caution should be exercised until the risk-to-benefit ratio and long-term therapeutic efficacy of vitamin E are better defined. Ursodeoxycholic acid (a bile acid that improves certain cholestatic liver diseases) and *betaine* (metabolite of choline that raises SAM levels and decreases cellular oxidative damage) offer no histologic benefit over placebo in patients with NASH. Experimental evidence to support the use of *omega-3 fatty acids* in NAFLD exists; however, a recent large, multicenter, placebo-controlled study failed to demonstrate a histologic benefit. Other pharmacotherapies are also being evaluated in NAFLD (e.g., *probiotics*, *farnesoid X receptor agonists*, *anticytokine agents*, *glucagon-like peptide agonists*,

dipeptidyl IV antagonists); however, sufficient data do not yet exist to justify their use as NASH treatments in standard clinical practice.

Statins are an important class of agents to treat dyslipidemia and decrease cardiovascular risk. There is no evidence to suggest that statins cause liver failure in patients with any chronic liver disease, including NAFLD. The incidence of liver enzyme elevations in NAFLD patients taking statins is also no different than that of healthy controls or patients with other chronic liver diseases. Moreover, several studies have suggested that statins may improve aminotransferases and histology in patients with NASH. Yet, there is continued reluctance to use statins in patients with NAFLD. The lack of evidence that statins harm the liver in NAFLD patients, combined with the increase risk for cardiovascular morbidity and mortality in NAFLD patients, warrants the use of statins to treat dyslipidemia in patients with NAFLD/NASH.

Bariatric Surgery Although interest in bariatric surgery as a treatment for NAFLD exists, a recently published Cochrane review concluded that lack of randomized clinical trials or adequate clinical studies prevents definitive assessment of benefits and harms of bariatric surgery as a treatment for NASH. Most studies of bariatric surgery have shown that bariatric surgery is generally safe in individuals with well-compensated chronic liver disease and improves hepatic steatosis and necroinflammation (i.e., features of NAFLD/NASH); however, effects on hepatic fibrosis have been variable. Concern lingers because some of the largest prospective studies suggest that hepatic fibrosis might progress after bariatric surgery. Thus, the Cochrane review deemed it premature to recommend bariatric surgery as a primary treatment for NASH. There is also general agreement that patients with NAFLD-related cirrhosis and portal hypertension should be excluded as candidates for bariatric surgery. However, given growing evidence for the benefits of bariatric surgery on metabolic syndrome complications in individuals with refractory obesity, it is not contraindicated in otherwise eligible patients with NAFLD or NASH.

Liver Transplantation Patients with NAFLD in whom end-stage liver disease develops should be evaluated for liver transplantation (Chap. 368). The outcomes of liver transplantation in well-selected patients with NAFLD are generally good, but comorbid medical conditions associated with NAFLD, such as diabetes mellitus, obesity, and cardiovascular disease, often limit transplant candidacy. NAFLD may recur after liver transplantation. The risk factors for recurrent or de novo NAFLD after liver transplantation are multifactorial and include hypertriglyceridemia, obesity, diabetes mellitus, and immunosuppressive therapies, particularly glucocorticoids.

GLOBAL HEALTH CONSIDERATIONS



The epidemic of obesity is now a global and accelerating phenomenon. Worldwide, there are over 1 billion overweight adults, of whom at least 300 million are obese. In the wake of the obesity epidemic follow numerous comorbidities, including NAFLD. NAFLD is the most common liver disease identified in Western countries and the fastest rising form of chronic liver disease worldwide. Present understanding of NAFLD natural history is based mainly on studies in whites who became overweight/obese and developed the metabolic syndrome in adulthood. The impact of the global childhood obesity epidemic on NAFLD pathogenesis/progression is unknown. Emerging evidence demonstrates that advanced NAFLD, including cirrhosis and primary liver cancer, can occur in children, prompting concerns that childhood-onset NAFLD might follow a more aggressive course than typical adult-acquired NAFLD. Some of the most populated parts of the world are in the midst of industrial revolutions, and certain environmental pollutants seem to exacerbate NAFLD. Some studies also suggest that the risk for NASH and NAFLD-related cirrhosis may be higher in certain ethnic groups such as Asians, certain Hispanics, and Native Americans and lower in others such as African Americans, compared with whites. Although all of these variables confound efforts to predict the net impact of this obesity-related liver disease on global health, it seems likely that NAFLD will remain a major cause of chronic liver disease worldwide for the foreseeable future.

365 Cirrhosis and Its Complications

Bruce R. Bacon

Cirrhosis is a condition that is defined histopathologically and has a variety of clinical manifestations and complications, some of which can be life-threatening. In the past, it has been thought that cirrhosis was never reversible; however, it has become apparent that when the underlying insult that has caused the cirrhosis has been removed, there can be reversal of fibrosis. This is most apparent with the successful treatment of chronic hepatitis C; however, reversal of fibrosis is also seen in patients with hemochromatosis who have been successfully treated and in patients with alcoholic liver disease who have discontinued alcohol use.

Regardless of the cause of cirrhosis, the pathologic features consist of the development of fibrosis to the point that there is architectural distortion with the formation of regenerative nodules. This results in a decrease in hepatocellular mass, and thus function, and an alteration of blood flow. The induction of fibrosis occurs with activation of hepatic stellate cells, resulting in the formation of increased amounts of collagen and other components of the extracellular matrix.

Clinical features of cirrhosis are the result of pathologic changes and mirror the severity of the liver disease. Most hepatic pathologists provide an assessment of grading and staging when evaluating liver biopsy samples. These grading and staging schemes vary between disease states and have been developed for most conditions, including chronic viral hepatitis, nonalcoholic fatty liver disease, and primary biliary cirrhosis. Advanced fibrosis usually includes bridging fibrosis with nodularity designated as stage 3 and cirrhosis designated as stage 4. Patients who have cirrhosis have varying degrees of compensated liver function, and clinicians need to differentiate between those who have stable, compensated cirrhosis and those who have decompensated cirrhosis. Patients who have developed complications of their liver disease and have become decompensated should be considered for liver transplantation. Many of the complications of cirrhosis will require specific therapy. *Portal hypertension* is a significant complicating feature of decompensated cirrhosis and is responsible for the development of ascites and bleeding from esophagogastric varices, two complications that signify decompensated cirrhosis. Loss of hepatocellular function results in jaundice, coagulation disorders, and hypoalbuminemia and contributes to the causes of portosystemic encephalopathy. The complications of cirrhosis are basically the same regardless of the etiology. Nonetheless, it is useful to classify patients by the cause of their liver disease (Table 365-1); patients can be divided into broad groups with alcoholic cirrhosis, cirrhosis due to chronic viral hepatitis, biliary cirrhosis, and other, less common causes such as cardiac cirrhosis, cryptogenic cirrhosis, and other miscellaneous causes.

ALCOHOLIC CIRRHOSIS

Excessive chronic alcohol use can cause several different types of chronic liver disease, including alcoholic fatty liver, alcoholic hepatitis, and alcoholic cirrhosis. Furthermore, use of excessive alcohol can

contribute to liver damage in patients with other liver diseases, such as hepatitis C, hemochromatosis, and fatty liver disease related to obesity. Chronic alcohol use can produce fibrosis in the absence of accompanying inflammation and/or necrosis. Fibrosis can be centrilobular, pericellular, or periportal. When fibrosis reaches a certain degree, there is disruption of the normal liver architecture and replacement of liver cells by regenerative nodules. In alcoholic cirrhosis, the nodules are usually <3 mm in diameter; this form of cirrhosis is referred to as *micronodular*. With cessation of alcohol use, larger nodules may form, resulting in a mixed micronodular and macronodular cirrhosis.

Pathogenesis Alcohol is the most commonly used drug in the United States, and more than two-thirds of adults drink alcohol each year. Thirty percent have had a binge within the past month, and over 7% of adults regularly consume more than two drinks per day. Unfortunately, more than 14 million adults in the United States meet the diagnostic criteria for alcohol abuse or dependence. In the United States, chronic liver disease is the tenth most common cause of death in adults, and alcoholic cirrhosis accounts for approximately 40% of deaths due to cirrhosis.

Ethanol is mainly absorbed by the small intestine and, to a lesser degree, through the stomach. Gastric alcohol dehydrogenase (ADH) initiates alcohol metabolism. Three enzyme systems account for metabolism of alcohol in the liver. These include cytosolic ADH, the microsomal ethanol oxidizing system (MEOS), and peroxisomal catalase. The majority of ethanol oxidation occurs via ADH to form acetaldehyde, which is a highly reactive molecule that may have multiple effects. Ultimately, acetaldehyde is metabolized to acetate by aldehyde dehydrogenase (ALDH). Intake of ethanol increases intracellular accumulation of triglycerides by increasing fatty acid uptake and by reducing fatty acid oxidation and lipoprotein secretion. Protein synthesis, glycosylation, and secretion are impaired. Oxidative damage to hepatocyte membranes occurs due to the formation of reactive oxygen species; acetaldehyde is a highly reactive molecule that combines with proteins to form protein-acetaldehyde adducts. These adducts may interfere with specific enzyme activities, including microtubular formation and hepatic protein trafficking. With acetaldehyde-mediated hepatocyte damage, certain reactive oxygen species can result in Kupffer cell activation. As a result, profibrogenic cytokines are produced that initiate and perpetuate stellate cell activation, with the resultant production of excess collagen and extracellular matrix. Connective tissue appears in both periportal and pericentral zones and eventually connects portal triads with central veins forming regenerative nodules. Hepatocyte loss occurs, and with increased collagen production and deposition, together with continuing hepatocyte destruction, the liver contracts and shrinks in size. This process generally takes from years to decades to occur and requires repeated insults.

Clinical Features The diagnosis of alcoholic liver disease requires an accurate history regarding both amount and duration of alcohol consumption. Patients with alcoholic liver disease can present with nonspecific symptoms such as vague right upper quadrant abdominal pain, fever, nausea and vomiting, diarrhea, anorexia, and malaise. Alternatively, they may present with more specific complications of chronic liver disease, including ascites, edema, or upper gastrointestinal (GI) hemorrhage. Many cases present incidentally at the time of autopsy or elective surgery. Other clinical manifestations include the development of jaundice or encephalopathy. The abrupt onset of any of these complications may be the first event prompting the patient to seek medical attention. Other patients may be identified in the course of an evaluation of routine laboratory studies that are found to be abnormal. On physical examination, the liver and spleen may be enlarged, with the liver edge being firm and nodular. Other frequent findings include scleral icterus, palmar erythema (Fig. 365-1), spider angiomas (Fig. 365-2), parotid gland enlargement, digital clubbing, muscle wasting, or the development of edema and ascites. Men may have decreased body hair and gynecomastia as well as testicular atrophy, which may be a consequence of hormonal abnormalities or a direct toxic effect of alcohol on the testes. In women with advanced

TABLE 365-1 CAUSES OF CIRRHOSIS

Alcoholism	Cardiac cirrhosis
Chronic viral hepatitis	Inherited metabolic liver disease
Hepatitis B	Hemochromatosis
Hepatitis C	Wilson's disease
Autoimmune hepatitis	α_1 Antitrypsin deficiency
Nonalcoholic steatohepatitis	Cystic fibrosis
Biliary cirrhosis	Cryptogenic cirrhosis
Primary biliary cirrhosis	
Primary sclerosing cholangitis	
Autoimmune cholangiopathy	



FIGURE 365-1 Palmar erythema. This figure shows palmar erythema in a patient with alcoholic cirrhosis. The erythema is peripheral over the palm with central pallor.

alcoholic cirrhosis, menstrual irregularities usually occur, and some women may be amenorrheic. These changes are often reversible following cessation of alcohol.

Laboratory tests may be completely normal in patients with early compensated alcoholic cirrhosis. Alternatively, in advanced liver disease, many abnormalities usually are present. Patients may be anemic either from chronic GI blood loss, nutritional deficiencies, or hypersplenism related to portal hypertension, or as a direct suppressive effect of alcohol on the bone marrow. A unique form of hemolytic anemia (with spur cells and acanthocytes) called *Zieve's syndrome* can occur in patients with severe alcoholic hepatitis. Platelet counts are often reduced early in the disease, reflective of portal hypertension with hypersplenism. Serum total bilirubin can be normal or elevated with advanced disease. Direct bilirubin is frequently mildly elevated in patients with a normal total bilirubin, but the abnormality typically progresses as the disease worsens. Prothrombin times are often prolonged and usually do not respond to administration of parenteral vitamin K. Serum sodium levels are usually normal unless patients have ascites and then can be depressed, largely due to ingestion of excess free water. Serum alanine and aspartate aminotransferases (ALT, AST) are typically elevated, particularly in patients who continue to drink, with AST levels being higher than ALT levels, usually by a 2:1 ratio.

Diagnosis Patients who have any of the above-mentioned clinical features, physical examination findings, or laboratory studies should be considered to have alcoholic liver disease. The diagnosis, however, requires accurate knowledge that the patient is continuing to use and

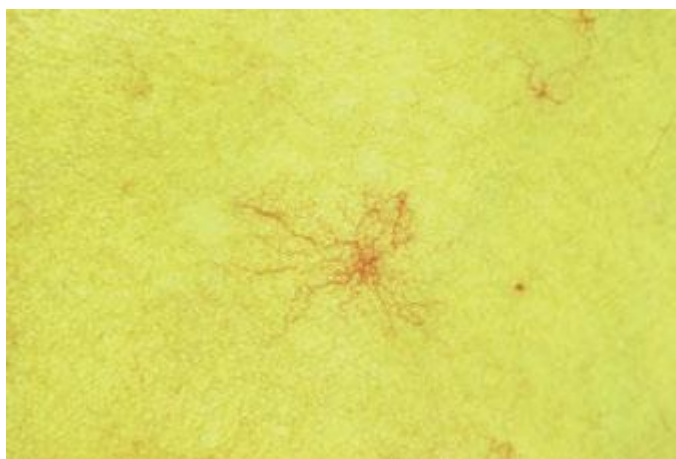


FIGURE 365-2 Spider angioma. This figure shows a spider angioma in a patient with hepatitis C cirrhosis. With release of central compression, the arteriole fills from the center and spreads out peripherally.

abuse alcohol. Furthermore, other forms of chronic liver disease (e.g., chronic viral hepatitis or metabolic or autoimmune liver diseases) must be considered or ruled out, or if present, an estimate of relative causality along with the alcohol use should be determined. Liver biopsy can be helpful to confirm a diagnosis, but generally when patients present with alcoholic hepatitis and are still drinking, liver biopsy is withheld until abstinence has been maintained for at least 6 months to determine residual, nonreversible disease.

In patients who have had complications of cirrhosis and who continue to drink, there is a <50% 5-year survival. In contrast, in patients who are able to remain abstinent, the prognosis is significantly improved. In patients with advanced liver disease, the prognosis remains poor; however, in individuals who are able to remain abstinent, liver transplantation is a viable option.

TREATMENT ALCOHOLIC CIRRHOSIS

Abstinence is the cornerstone of therapy for patients with alcoholic liver disease. In addition, patients require good nutrition and long-term medical supervision to manage underlying complications that may develop. Complications such as the development of ascites and edema, variceal hemorrhage, or portosystemic encephalopathy all require specific management and treatment. Glucocorticoids are occasionally used in patients with severe alcoholic hepatitis in the absence of infection. Survival has been shown to improve in certain studies. Treatment is restricted to patients with a discriminant function (DF) value of >32. The DF is calculated as the serum total bilirubin plus the difference in the patient's prothrombin time compared to control (in seconds) multiplied by 4.6. In patients for whom this value is >32, there is improved survival at 28 days with the use of glucocorticoids.

Other therapies that have been used include oral pentoxifylline, which decreases the production of tumor necrosis factor α (TNF- α) and other proinflammatory cytokines. In contrast to glucocorticoids, with which complications can occur, pentoxifylline is relatively easy to administer and has few, if any, side effects. A variety of nutritional therapies have been tried with either parenteral or enteral feedings; however, it is unclear whether any of these modalities have significantly improved survival.

Recent studies have used parenterally administered inhibitors of TNF- α such as infliximab or etanercept. Early results have shown no adverse events; however, there was no clear-cut improvement in survival. Anabolic steroids, propylthiouracil, antioxidants, colchicine, and penicillamine have all been used but do not show clear-cut benefits and are not recommended.

As mentioned above, the cornerstone to treatment is cessation of alcohol use. Recent experience with medications that reduce craving for alcohol, such as acamprosate calcium, has been favorable. Patients may take other necessary medications even in the presence of cirrhosis. Acetaminophen use is often discouraged in patients with liver disease; however, if no more than 2 g of acetaminophen per day are consumed, there generally are no problems.

CIRRHOSIS DUE TO CHRONIC VIRAL HEPATITIS B OR C



Of patients exposed to the hepatitis C virus (HCV), approximately 80% develop chronic hepatitis C, and of those, about 20–30% will develop cirrhosis over 20–30 years. Many of these patients have had concomitant alcohol use, and the true incidence of cirrhosis due to hepatitis C alone is unknown. Nonetheless, this represents a significant number of patients. It is expected that an even higher percentage will go on to develop cirrhosis over longer periods of time. In the United States, approximately 5 to 6 million people have been exposed to HCV, with about 4 million who are chronically viremic. Worldwide, about 170 million individuals have hepatitis C, with some areas of the world (e.g., Egypt) having up to 15% of the population infected. HCV is a noncytotoxic virus, and liver damage is probably immune-mediated. Progression of liver disease due to chronic

2060 hepatitis C is characterized by portal-based fibrosis with bridging fibrosis and nodularity developing, ultimately culminating in the development of cirrhosis. In cirrhosis due to chronic hepatitis C, the liver is small and shrunken with characteristic features of a mixed micro- and macronodular cirrhosis seen on liver biopsy. In addition to the increased fibrosis that is seen in cirrhosis due to hepatitis C, an inflammatory infiltrate is found in portal areas with interface hepatitis and occasionally some lobular hepatocellular injury and inflammation. In patients with HCV genotype 3, steatosis is often present.



Similar findings are seen in patients with cirrhosis due to chronic hepatitis B. Of adult patients exposed to hepatitis B, about 5% develop chronic hepatitis B, and about 20% of those patients will go on to develop cirrhosis. Special stains for hepatitis B core (HBc) and hepatitis B surface (HBs) antigen will be positive, and ground-glass hepatocytes signifying hepatitis B surface antigen (HBsAg) may be present. In the United States, there are about 2 million carriers of hepatitis B, whereas in other parts of the world where hepatitis B virus (HBV) is endemic (i.e., Asia, Southeast Asia, sub-Saharan Africa), up to 15% of the population may be infected, having acquired the infection vertically at the time of birth. Thus, over 300–400 million individuals are thought to have hepatitis B worldwide. Approximately 25% of these individuals may ultimately develop cirrhosis.

Clinical Features and Diagnosis Patients with cirrhosis due to either chronic hepatitis C or B can present with the usual symptoms and signs of chronic liver disease. Fatigue, malaise, vague right upper quadrant pain, and laboratory abnormalities are frequent presenting features. Diagnosis requires a thorough laboratory evaluation, including quantitative HCV RNA testing and analysis for HCV genotype, or hepatitis B serologies to include HBsAg, anti-HBs, HBeAg (hepatitis B e antigen), anti-HBe, and quantitative HBV DNA levels.

TREATMENT **CIRRHOSIS DUE TO CHRONIC VIRAL HEPATITIS B OR C**

Management of complications of cirrhosis revolves around specific therapy for treatment of whatever complications occur (e.g., esophageal variceal hemorrhage, development of ascites and edema, or encephalopathy). In patients with chronic hepatitis B, numerous studies have shown beneficial effects of antiviral therapy, which is effective at viral suppression, as evidenced by reducing aminotransferase levels and HBV DNA levels, and improving histology by reducing inflammation and fibrosis. Several clinical trials and case series have demonstrated that patients with decompensated liver disease can become compensated with the use of antiviral therapy directed against hepatitis B. Currently available therapy includes lamivudine, adefovir, telbivudine, entecavir, and tenofovir. Interferon α can also be used for treating hepatitis B, but it should not be used in cirrhotics.

Treatment of patients with cirrhosis due to hepatitis C is a little more difficult because the side effects of pegylated interferon and ribavirin therapy are often difficult to manage. Dose-limiting cytopenias (platelets, white blood cells, red blood cells) or severe side effects can result in discontinuation of treatment. Nonetheless, if patients can tolerate treatment, and if it is successful, the benefit is great and disease progression is reduced. Recent studies have shown that if platelets are $<100,000$, albumin is <3.5 g/dL, and Model for End-Stage Liver Disease (MELD) score is >10 , the risk of severe complications of interferon-based antiviral therapy is significant. Recent approval of Direct Acting Antivirals (DAAs) has led to improved efficacy of treatment with regimens that are safe and well tolerated.

CIRRHOSIS FROM AUTOIMMUNE HEPATITIS AND NONALCOHOLIC FATTY LIVER DISEASE

Other causes of posthepatic cirrhosis include autoimmune hepatitis and cirrhosis due to nonalcoholic steatohepatitis. Many patients with autoimmune hepatitis (AIH) present with cirrhosis that is already

established. Typically, these patients will not benefit from immunosuppressive therapy with glucocorticoids or azathioprine because the AIH is “burned out.” In this situation, liver biopsy does not show a significant inflammatory infiltrate. Diagnosis in this setting requires positive autoimmune markers such as antinuclear antibody (ANA) or anti-smooth-muscle antibody (ASMA). When patients with AIH present with cirrhosis and active inflammation accompanied by elevated liver enzymes, there can be considerable benefit from the use of immunosuppressive therapy.

Patients with nonalcoholic steatohepatitis are increasingly being found to have progressed to cirrhosis. With the epidemic of obesity that continues in Western countries, more and more patients are identified with nonalcoholic fatty liver disease (**Chap. 364**). Of these, a significant subset has nonalcoholic steatohepatitis and can progress to increased fibrosis and cirrhosis. Over the past several years, it has been increasingly recognized that many patients who were thought to have cryptogenic cirrhosis in fact have nonalcoholic steatohepatitis. As their cirrhosis progresses, they become catabolic and then lose the telltale signs of steatosis seen on biopsy. Management of complications of cirrhosis due to either AIH or nonalcoholic steatohepatitis is similar to that for other forms of cirrhosis.

BILIARY CIRRHOSIS

Biliary cirrhosis has pathologic features that are different from either alcoholic cirrhosis or posthepatic cirrhosis, yet the manifestations of end-stage liver disease are the same. Cholestatic liver disease may result from necroinflammatory lesions, congenital or metabolic processes, or external bile duct compression. Thus, two broad categories reflect the anatomic sites of abnormal bile retention: *intrahepatic* and *extrahepatic*. The distinction is important for obvious therapeutic reasons. Extrahepatic obstruction may benefit from surgical or endoscopic biliary tract decompression, whereas intrahepatic cholestatic processes will not improve with such interventions and require a different approach.

The major causes of chronic cholestatic syndromes are primary biliary cirrhosis (PBC), autoimmune cholangitis (AIC), primary sclerosing cholangitis (PSC), and idiopathic adulthood ductopenia. These syndromes are usually clinically distinguished from each other by antibody testing, cholangiographic findings, and clinical presentation. However, they all share the histopathologic features of chronic cholestasis, such as cholate stasis; copper deposition; xanthomatous transformation of hepatocytes; and irregular, so-called biliary fibrosis. In addition, there may be chronic portal inflammation, interface activity, and chronic lobular inflammation. Ductopenia is a result of this progressive disease as patients develop cirrhosis.

PRIMARY BILIARY CIRRHOSIS

PBC is seen in about 100–200 individuals per million, with a strong female preponderance and a median age of around 50 years at the time of diagnosis. The cause of PBC is unknown; it is characterized by portal inflammation and necrosis of cholangiocytes in small- and medium-sized bile ducts. Cholestatic features prevail, and biliary cirrhosis is characterized by an elevated bilirubin level and progressive liver failure. Liver transplantation is the treatment of choice for patients with decompensated cirrhosis due to PBC. A variety of therapies have been proposed, but ursodeoxycholic acid (UDCA) is the only approved treatment that has some degree of efficacy by slowing the rate of progression of the disease.

Antimitochondrial antibodies (AMA) are present in about 90% of patients with PBC. These autoantibodies recognize intermitochondrial membrane proteins that are enzymes of the pyruvate dehydrogenase complex (PDC), the branched-chain 2-oxoacid dehydrogenase complex, and the 2-oxoglutarate dehydrogenase complex. Most relate to pyruvate dehydrogenase. These autoantibodies are not pathogenic but rather are useful markers for making a diagnosis of PBC.

Pathology Histopathologic analyses of liver biopsies of patients with PBC have resulted in identifying four distinct stages of the disease as it progresses. The earliest lesion is termed *chronic nonsuppurative*

destructive cholangitis and is a necrotizing inflammatory process of the portal tracts. Medium and small bile ducts are infiltrated with lymphocytes and undergo duct destruction. Mild fibrosis and sometimes bile stasis can occur. With progression, the inflammatory infiltrate becomes less prominent, but the number of bile ducts is reduced and there is proliferation of smaller bile ductules. Increased fibrosis ensues with the expansion of periportal fibrosis to bridging fibrosis. Finally, cirrhosis, which may be micronodular or macronodular, develops.

Clinical Features Currently, most patients with PBC are diagnosed well before the end-stage manifestations of the disease are present, and, as such, most patients are actually asymptomatic. When symptoms are present, they most prominently include a significant degree of fatigue out of proportion to what would be expected for either the severity of the liver disease or the age of the patient. Pruritus is seen in approximately 50% of patients at the time of diagnosis, and it can be debilitating. It might be intermittent and usually is most bothersome in the evening. In some patients, pruritus can develop toward the end of pregnancy, and there are examples of patients having been diagnosed with cholestasis of pregnancy rather than PBC. Pruritus that presents prior to the development of jaundice indicates severe disease and a poor prognosis.

Physical examination can show jaundice and other complications of chronic liver disease, including hepatomegaly, splenomegaly, ascites, and edema. Other features that are unique to PBC include hyperpigmentation, xanthelasma, and xanthomata, which are related to the altered cholesterol metabolism seen in this disease. Hyperpigmentation is evident on the trunk and the arms and is seen in areas of exfoliation and lichenification associated with progressive scratching related to the pruritus. Bone pain resulting from osteopenia or osteoporosis is occasionally seen at the time of diagnosis.

Laboratory Findings Laboratory findings in PBC show cholestatic liver enzyme abnormalities with an elevation in γ -glutamyl transpeptidase and alkaline phosphatase (ALP) along with mild elevations in aminotransferases (ALT and AST). Immunoglobulins, particularly IgM, are typically increased. Hyperbilirubinemia usually is seen once cirrhosis has developed. Thrombocytopenia, leukopenia, and anemia may be seen in patients with portal hypertension and hypersplenism. Liver biopsy shows characteristic features as described above and should be evident to any experienced hepatopathologist. Up to 10% of patients with characteristic PBC will have features of AIH as well and are defined as having “overlap” syndrome. These patients are usually treated as PBC patients and may progress to cirrhosis with the same frequency as typical PBC patients. Some patients require immunosuppressive medications as well.

Diagnosis PBC should be considered in patients with chronic cholestatic liver enzyme abnormalities. It is most often seen in middle-aged women. AMA testing may be negative, and it should be remembered that as many as 10% of patients with PBC may be AMA-negative. Liver biopsy is most important in this setting of AMA-negative PBC. In patients who are AMA-negative with cholestatic liver enzymes, PSC should be ruled out by way of cholangiography.

TREATMENT PRIMARY BILIARY CIRRHOSIS

Treatment of the typical manifestations of cirrhosis are no different for PBC than for other forms of cirrhosis. UDCA has been shown to improve both biochemical and histologic features of the disease. Improvement is greatest when therapy is initiated early; the likelihood of significant improvement with UDCA is low in patients with PBC who present with manifestations of cirrhosis. UDCA is given in doses of 13–15 mg/kg per day; the medication is usually well-tolerated, although some patients have worsening pruritus with initiation of therapy. A small proportion of patients may have diarrhea or headache as a side effect of the drug. UDCA has been shown to slow the rate of progression of PBC, but it does not reverse or cure the disease. Patients with PBC require long-term follow-up by a physician experienced with the disease. Certain patients may need

to be considered for liver transplantation should their liver disease decompensate.

The main symptoms of PBC are fatigue and pruritus, and symptom management is important. Several therapies have been tried for treatment of fatigue, but none of them have been successful; frequent naps should be encouraged. Pruritus is treated with antihistamines, narcotic receptor antagonists (naltrexone), and rifampin. Cholestyramine, a bile salt–sequestering agent, has been helpful in some patients but is somewhat tedious and difficult to take. Plasmapheresis has been used rarely in patients with severe intractable pruritus. There is an increased incidence of osteopenia and osteoporosis in patients with cholestatic liver disease, and bone density testing should be performed. Treatment with a bisphosphonate should be instituted when bone disease is identified.

PRIMARY SCLEROSING CHOLANGITIS

As in PBC, the cause of PSC remains unknown. PSC is a chronic cholestatic syndrome that is characterized by diffuse inflammation and fibrosis involving the entire biliary tree, resulting in chronic cholestasis. This pathologic process ultimately results in obliteration of both the intra- and extrahepatic biliary tree, leading to biliary cirrhosis, portal hypertension, and liver failure. The cause of PSC remains unknown despite extensive investigation into various mechanisms related to bacterial and viral infections, toxins, genetic predisposition, and immunologic mechanisms, all of which have been postulated to contribute to the pathogenesis and progression of this syndrome.

Pathologic changes that can occur in PSC show bile duct proliferation as well as ductopenia and fibrous cholangitis (pericholangitis). Often, liver biopsy changes in PSC are not pathognomonic, and establishing the diagnosis of PSC must involve imaging of the biliary tree. Periductal fibrosis is occasionally seen on biopsy specimens and can be quite helpful in making the diagnosis. As the disease progresses, biliary cirrhosis is the final, end-stage manifestation of PSC.

Clinical Features The usual clinical features of PSC are those found in cholestatic liver disease, with fatigue, pruritus, steatorrhea, deficiencies of fat-soluble vitamins, and the associated consequences. As in PBC, the fatigue is profound and nonspecific. Pruritus can often be debilitating and is related to the cholestasis. The severity of pruritus does not correlate with the severity of the disease. Metabolic bone disease, as seen in PBC, can occur with PSC and should be treated (see above).

Laboratory Findings Patients with PSC typically are identified in the course of an evaluation of abnormal liver enzymes. Most patients have at least a twofold increase in ALP and may have elevated aminotransferases as well. Albumin levels may be decreased, and prothrombin times are prolonged in a substantial proportion of patients at the time of diagnosis. Some degree of correction of a prolonged prothrombin time may occur with parenteral vitamin K. A small subset of patients have aminotransferase elevations greater than five times the upper limit of normal and may have features of AIH on biopsy. These individuals are thought to have an overlap syndrome between PSC and AIH. Autoantibodies are frequently positive in patients with the overlap syndrome but are typically negative in patients who only have PSC. One autoantibody, the perinuclear antineutrophil cytoplasmic antibody (p-ANCA), is positive in about 65% of patients with PSC. Over 50% of patients with PSC also have ulcerative colitis (UC); accordingly, once a diagnosis of PSC is established, colonoscopy should be performed to look for evidence of UC.

Diagnosis The definitive diagnosis of PSC requires cholangiographic imaging. Over the last several years, magnetic resonance imaging (MRI) with magnetic resonance cholangiopancreatography (MRCP) has been used as the imaging technique of choice for initial evaluation. Once patients are screened in this manner, some investigators feel that endoscopic retrograde cholangiopancreatography (ERCP) should also be performed to be certain whether or not a dominant stricture is present. Typical cholangiographic findings in PSC are multifocal stricturing and beading involving both the intrahepatic and extrahepatic biliary tree. However, although involvement may be of

2062 the intrahepatic bile ducts alone or of the extrahepatic bile ducts alone, more commonly, both are involved. These strictures are typically short and with intervening segments of normal or slightly dilated bile ducts that are distributed diffusely, producing the classic beaded appearance. The gallbladder and cystic duct can be involved in up to 15% of cases. Patients with high-grade, diffuse stricturing of the intrahepatic bile ducts have an overall poor prognosis. Gradually, biliary cirrhosis develops, and patients will progress to decompensated liver disease with all the manifestations of ascites, esophageal variceal hemorrhage, and encephalopathy.

TREATMENT PRIMARY SCLEROSING CHOLANGITIS

There is no specific proven treatment for PSC. A recently completed study of high-dose (20 mg/kg per day) UDCA was found to be harmful. Some clinicians use UDCA at “PBC dosages” of 13–15 mg/kg per day with anecdotal improvement. Endoscopic dilatation of dominant strictures can be helpful, but the ultimate treatment is liver transplantation. A dreaded complication of PSC is the development of cholangiocarcinoma, which is a relative contraindication to liver transplantation. Symptoms of pruritus are common, and the approach is as mentioned previously for this problem in patients with PBC (see above).

CARDIAC CIRRHOSIS

Definition Patients with long-standing right-sided congestive heart failure may develop chronic liver injury and cardiac cirrhosis. This is an increasingly uncommon, if not rare, cause of chronic liver disease given the advances made in the care of patients with heart failure.

Etiology and Pathology In the case of long-term right-sided heart failure, there is an elevated venous pressure transmitted via the inferior vena cava and hepatic veins to the sinusoids of the liver, which become dilated and engorged with blood. The liver becomes enlarged and swollen, and with long-term passive congestion and relative ischemia due to poor circulation, centrilobular hepatocytes can become necrotic, leading to pericentral fibrosis. This fibrotic pattern can extend to the periphery of the lobule outward until a unique pattern of fibrosis causing cirrhosis can occur.

Clinical Features Patients typically have signs of congestive heart failure and will manifest an enlarged firm liver on physical examination. ALP levels are characteristically elevated, and aminotransferases may be normal or slightly increased with AST usually higher than ALT. It is unlikely that patients will develop variceal hemorrhage or encephalopathy.

Diagnosis The diagnosis is usually made in someone with clear-cut cardiac disease who has an elevated ALP and an enlarged liver. Liver biopsy shows a pattern of fibrosis that can be recognized by an experienced hepatopathologist. Differentiation from Budd-Chiari syndrome (BCS) can be made by seeing extravasation of red blood cells in BCS, but not in cardiac hepatopathy. Venooclusive disease can also affect hepatic outflow and has characteristic features on liver biopsy. Venooclusive disease can be seen under the circumstances of conditioning for bone marrow transplant with radiation and chemotherapy; it can also be seen with the ingestion of certain herbal teas as well as pyrrolizidine alkaloids. This is typically seen in Caribbean countries and rarely in the United States. Treatment is based on management of the underlying cardiac disease.

OTHER TYPES OF CIRRHOSIS

There are several other less common causes of chronic liver disease that can progress to cirrhosis. These include inherited metabolic liver diseases such as hemochromatosis, Wilson’s disease, α_1 antitrypsin (α_1 AT) deficiency, and cystic fibrosis. For all of these disorders, the manifestations of cirrhosis are similar, with some minor variations, to those seen in other patients with other causes of cirrhosis.

Hemochromatosis is an inherited disorder of iron metabolism that results in a progressive increase in hepatic iron deposition, which, over time, can lead to a portal-based fibrosis progressing to cirrhosis, liver failure, and hepatocellular cancer. While the frequency of hemochromatosis is relatively common, with genetic susceptibility occurring in 1 in 250 individuals, the frequency of end-stage manifestations due to the disease is relatively low, and fewer than 5% of those patients who are genotypically susceptible will go on to develop severe liver disease from hemochromatosis. Diagnosis is made with serum iron studies showing an elevated transferrin saturation and an elevated ferritin level, along with abnormalities identified by *HFE* mutation analysis. Treatment is straightforward, with regular therapeutic phlebotomy.

Wilson’s disease is an inherited disorder of copper homeostasis with failure to excrete excess amounts of copper, leading to an accumulation in the liver. This disorder is relatively uncommon, affecting 1 in 30,000 individuals. Wilson’s disease typically affects adolescents and young adults. Prompt diagnosis before end-stage manifestations become irreversible can lead to significant clinical improvement. Diagnosis requires determination of ceruloplasmin levels, which are low; 24-h urine copper levels, which are elevated; typical physical examination findings, including Kayser-Fleischer corneal rings; and characteristic liver biopsy findings. Treatment consists of copper-chelating medications.

α_1 AT deficiency results from an inherited disorder that causes abnormal folding of the α_1 AT protein, resulting in failure of secretion of that protein from the liver. It is unknown how the retained protein leads to liver disease. Patients with α_1 AT deficiency at greatest risk for developing chronic liver disease have the ZZ phenotype, but only about 10–20% of such individuals will develop chronic liver disease. Diagnosis is made by determining α_1 AT levels and phenotype. Characteristic periodic acid–Schiff (PAS)-positive, diastase-resistant globules are seen on liver biopsy. The only effective treatment is liver transplantation, which is curative.

Cystic fibrosis is an uncommon inherited disorder affecting whites of northern European descent. A biliary-type cirrhosis can occur, and some patients derive benefit from the chronic use of UDCA.

MAJOR COMPLICATIONS OF CIRRHOSIS

The clinical course of patients with advanced cirrhosis is often complicated by a number of important sequelae that can occur regardless of the underlying cause of the liver disease. These include portal hypertension and its consequences of gastroesophageal variceal hemorrhage, splenomegaly, ascites, hepatic encephalopathy, spontaneous bacterial peritonitis (SBP), hepatorenal syndrome, and hepatocellular carcinoma (Table 365-2).

PORTAL HYPERTENSION

Portal hypertension is defined as the elevation of the hepatic venous pressure gradient (HVPG) to >5 mmHg. Portal hypertension is caused by a combination of two simultaneously occurring hemodynamic processes: (1) increased intrahepatic resistance to the passage of blood

TABLE 365-2 COMPLICATIONS OF CIRRHOSIS

Portal hypertension	Coagulopathy
Gastroesophageal varices	Factor deficiency
Portal hypertensive gastropathy	Fibrinolysis
Splenomegaly, hypersplenism	Thrombocytopenia
Ascites	Bone disease
Spontaneous bacterial peritonitis	Osteopenia
Hepatorenal syndrome	Osteoporosis
Type 1	Osteomalacia
Type 2	Hematologic abnormalities
Hepatic encephalopathy	Anemia
Hepatopulmonary syndrome	Hemolysis
Portopulmonary hypertension	Thrombocytopenia
Malnutrition	Neutropenia

flow through the liver due to cirrhosis and regenerative nodules, and (2) increased splanchnic blood flow secondary to vasodilation within the splanchnic vascular bed. Portal hypertension is directly responsible for the two major complications of cirrhosis: variceal hemorrhage and ascites. *Variceal hemorrhage* is an immediate life-threatening problem with a 20–30% mortality rate associated with each episode of bleeding. The portal venous system normally drains blood from the stomach, intestines, spleen, pancreas, and gallbladder, and the portal vein is formed by the confluence of the superior mesenteric and splenic veins. Deoxygenated blood from the small bowel drains into the superior mesenteric vein along with blood from the head of the pancreas, the ascending colon, and part of the transverse colon. Conversely, the splenic vein drains the spleen and the pancreas and is joined by the inferior mesenteric vein, which brings blood from the transverse and descending colon as well as from the superior two-thirds of the rectum. Thus, the portal vein normally receives blood from almost the entire GI tract.

The causes of portal hypertension are usually subcategorized as prehepatic, intrahepatic, and posthepatic (Table 365-3). Prehepatic causes of portal hypertension are those affecting the portal venous system before it enters the liver; they include portal vein thrombosis and splenic vein thrombosis. Posthepatic causes encompass those affecting the hepatic veins and venous drainage to the heart; they include BCS, venoocclusive disease, and chronic right-sided cardiac congestion. Intrahepatic causes account for over 95% of cases of portal hypertension and are represented by the major forms of cirrhosis. Intrahepatic causes of portal hypertension can be further subdivided into presinusoidal, sinusoidal, and postsinusoidal causes. Postsinusoidal causes include venoocclusive disease, whereas presinusoidal causes include congenital hepatic fibrosis and schistosomiasis. Sinusoidal causes are related to cirrhosis from various causes.

Cirrhosis is the most common cause of portal hypertension in the United States, and clinically significant portal hypertension is present in >60% of patients with cirrhosis. Portal vein obstruction may be idiopathic or can occur in association with cirrhosis or with infection, pancreatitis, or abdominal trauma.

Coagulation disorders that can lead to the development of portal vein thrombosis include polycythemia vera; essential thrombocytosis; deficiencies in protein C, protein S, antithrombin 3, and factor V Leiden; and abnormalities in the gene-regulating prothrombin production. Some patients may have a subclinical myeloproliferative disorder.

Clinical Features The three primary complications of portal hypertension are gastroesophageal varices with hemorrhage, ascites, and

hypersplenism. Thus, patients may present with upper GI bleeding, which, on endoscopy, is found to be due to esophageal or gastric varices; with the development of ascites along with peripheral edema; or with an enlarged spleen with associated reduction in platelets and white blood cells on routine laboratory testing.

ESOPHAGEAL VARICES Over the last decade, it has become common practice to screen known cirrhotics with endoscopy to look for esophageal varices. Such screening studies have shown that approximately one-third of patients with histologically confirmed cirrhosis have varices. Approximately 5–15% of cirrhotics per year develop varices, and it is estimated that the majority of patients with cirrhosis will develop varices over their lifetimes. Furthermore, it is anticipated that roughly one-third of patients with varices will develop bleeding. Several factors predict the risk of bleeding, including the severity of cirrhosis (Child's class, MELD score); the height of wedged-hepatic vein pressure; the size of the varix; the location of the varix; and certain endoscopic stigmata, including red wale signs, hematocystic spots, diffuse erythema, bluish color, cherry red spots, or white-nipple spots. Patients with tense ascites are also at increased risk for bleeding from varices.

Diagnosis In patients with cirrhosis who are being followed chronically, the development of portal hypertension is usually revealed by the presence of thrombocytopenia; the appearance of an enlarged spleen; or the development of ascites, encephalopathy, and/or esophageal varices with or without bleeding. In previously undiagnosed patients, any of these features should prompt further evaluation to determine the presence of portal hypertension and liver disease. Varices should be identified by endoscopy. Abdominal imaging, either by computed tomography (CT) or MRI, can be helpful in demonstrating a nodular liver and in finding changes of portal hypertension with intraabdominal collateral circulation. If necessary, interventional radiologic procedures can be performed to determine wedged and free hepatic vein pressures that will allow for the calculation of a wedged-to-free gradient, which is equivalent to the portal pressure. The average normal wedged-to-free gradient is 5 mmHg, and patients with a gradient >12 mmHg are at risk for variceal hemorrhage.

TREATMENT VARICEAL HEMORRHAGE

Treatment for variceal hemorrhage as a complication of portal hypertension is divided into two main categories: (1) primary prophylaxis and (2) prevention of rebleeding once there has been an initial variceal hemorrhage. Primary prophylaxis requires routine screening by endoscopy of all patients with cirrhosis. Once varices that are at increased risk for bleeding are identified, primary prophylaxis can be achieved either through nonselective beta blockade or by variceal band ligation. Numerous placebo-controlled clinical trials of either propranolol or nadolol have been reported in the literature. The most rigorous studies were those that only included patients with significantly enlarged varices or with hepatic vein pressure gradients >12 mmHg. Patients treated with beta blockers have a lower risk of variceal hemorrhage than those treated with placebo over 1 and 2 years of follow-up. There is also a decrease in mortality related to variceal hemorrhage. Unfortunately, overall survival was improved in only one study. Further studies have demonstrated that the degree of reduction of portal pressure is a significant feature to determine success of therapy. Therefore, it has been suggested that repeat measurements of hepatic vein pressure gradients may be used to guide pharmacologic therapy; however, this may be cost-prohibitive. Several studies have evaluated variceal band ligation and variceal sclerotherapy as methods for providing primary prophylaxis.

Endoscopic variceal ligation (EVL) has achieved a level of success and comfort with most gastroenterologists who see patients with these complications of portal hypertension. Thus, in patients with cirrhosis who are screened for portal hypertension and are found to have large varices, it is recommended that they receive either beta blockade or primary prophylaxis with EVL.

TABLE 365-3 CLASSIFICATION OF PORTAL HYPERTENSION

Prehepatic
Portal vein thrombosis
Splenic vein thrombosis
Massive splenomegaly (Banti's syndrome)
Hepatic
Presinusoidal
Schistosomiasis
Congenital hepatic fibrosis
Sinusoidal
Cirrhosis—many causes
Alcoholic hepatitis
Postsinusoidal
Hepatic sinusoidal obstruction (venoocclusive syndrome)
Posthepatic
Budd-Chiari syndrome
Inferior vena caval webs
Cardiac causes
Restrictive cardiomyopathy
Constrictive pericarditis
Severe congestive heart failure

The approach to patients once they have had a variceal bleed is first to treat the acute bleed, which can be life-threatening, and then to prevent further bleeding. Prevention of further bleeding is usually accomplished with repeated variceal band ligation until varices are obliterated. Treatment of acute bleeding requires both fluid and blood-product replacement as well as prevention of subsequent bleeding with EVL.

The medical management of acute variceal hemorrhage includes the use of vasoconstricting agents, usually somatostatin or octreotide. Vasopressin was used in the past but is no longer commonly used. Balloon tamponade (Sengstaken-Blakemore tube or Minnesota tube) can be used in patients who cannot get endoscopic therapy immediately or who need stabilization prior to endoscopic therapy. Control of bleeding can be achieved in the vast majority of cases; however, bleeding recurs in the majority of patients if definitive endoscopic therapy has not been instituted. Octreotide, a direct splanchnic vasoconstrictor, is given at dosages of 50–100 µg/h by continuous infusion. Endoscopic intervention is used as first-line treatment to control bleeding acutely. Some endoscopists will use variceal injection therapy (sclerotherapy) as initial therapy, particularly when bleeding is vigorous. Variceal band ligation is used to control acute bleeding in over 90% of cases and should be repeated until obliteration of all varices is accomplished. When esophageal varices extend into the proximal stomach, band ligation is less successful. In these situations, when bleeding continues from gastric varices, consideration for a transjugular intrahepatic portosystemic shunt (TIPS) should be made. This technique creates a portosystemic shunt by a percutaneous approach using an expandable metal stent, which is advanced under angiographic guidance to the hepatic veins and then through the substance of the liver to create a direct portocaval shunt. This offers an alternative to surgery for acute decompression of portal hypertension. Encephalopathy can occur in as many as 20% of patients after TIPS and is particularly problematic in elderly patients and in patients with preexisting encephalopathy. TIPS should be reserved for individuals who fail endoscopic or medical management or who are poor surgical risks. TIPS can sometimes be used as a bridge to transplantation. Surgical esophageal transection is a procedure that is rarely used and generally is associated with a poor outcome.

PREVENTION OF RECURRENT BLEEDING (FIG. 365-3)

Once patients have had an acute bleed and have been managed successfully, attention should be paid to preventing recurrent bleeding. This usually requires repeated variceal band ligation until varices are obliterated. Beta blockade may be of adjunctive benefit in patients who are having recurrent variceal band ligation; however, once varices have been obliterated, the need for beta blockade is lessened. Despite successful variceal obliteration, many patients will still have portal hypertensive gastropathy from which bleeding can occur. Nonselective beta blockade may be helpful to prevent further bleeding from portal hypertensive gastropathy once varices have been obliterated.

Portosystemic shunt surgery is less commonly performed with the advent of TIPS; nonetheless, this procedure should be considered for patients with good hepatic synthetic function who could benefit by having portal decompressive surgery.

SPLENOMEGALY AND HYPERSPLENISM

Congestive splenomegaly is common in patients with portal hypertension. Clinical features include the presence of an enlarged spleen on physical examination and the development of thrombocytopenia and leukopenia in patients who have cirrhosis. Some patients will have fairly significant left-sided and left upper quadrant abdominal pain related to an enlarged and engorged spleen. Splenomegaly itself usually requires no specific treatment, although splenectomy can be successfully performed under very special circumstances.

Hypersplenism with the development of thrombocytopenia is a common feature of patients with cirrhosis and is usually the first indication of portal hypertension.

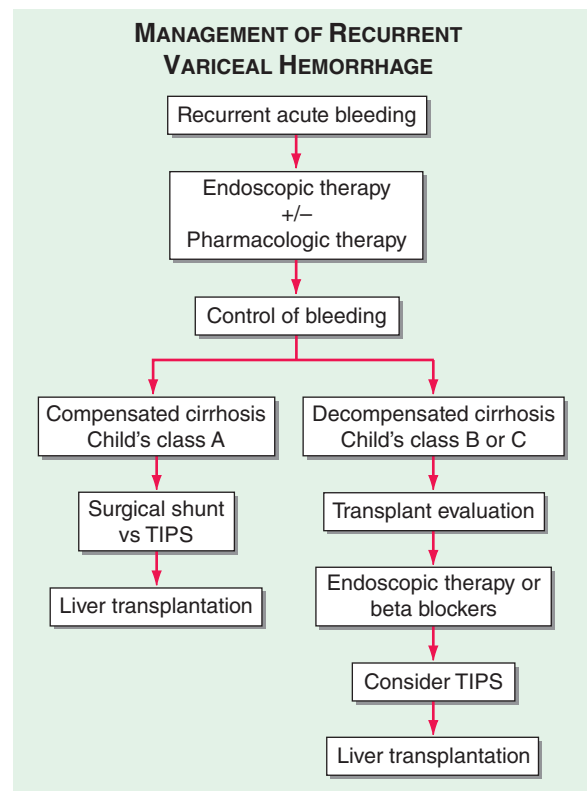


FIGURE 365-3 Management of recurrent variceal hemorrhage.

This algorithm describes an approach to management of patients who have recurrent bleeding from esophageal varices. Initial therapy is generally with endoscopic therapy often supplemented by pharmacologic therapy. With control of bleeding, a decision needs to be made as to whether patients should go on to a surgical shunt or TIPS (if they are Child's class A) and be considered for transplant, or if they should have TIPS and be considered for transplant (if they are Child's class B or C). TIPS, transjugular intrahepatic portosystemic shunt.

ASCITES

Definition Ascites is the accumulation of fluid within the peritoneal cavity. Overwhelmingly, the most common cause of ascites is portal hypertension related to cirrhosis; however, clinicians should remember that malignant or infectious causes of ascites can be present as well, and careful differentiation of these other causes are obviously important for patient care.

Pathogenesis The presence of portal hypertension contributes to the development of ascites in patients who have cirrhosis (Fig. 365-4). There is an increase in intrahepatic resistance, causing increased portal pressure, but there is also vasodilation of the splanchnic arterial system, which, in turn, results in an increase in portal venous inflow. Both of these abnormalities result in increased production of splanchnic lymph. Vasodilating factors such as nitric oxide are responsible for the vasodilatory effect. These hemodynamic changes result in sodium retention by causing activation of the renin-angiotensin-aldosterone system with the development of hyperaldosteronism. The renal effects of increased aldosterone leading to sodium retention also contribute to the development of ascites. Sodium retention causes fluid accumulation and expansion of the extracellular fluid volume, which results in the formation of peripheral edema and ascites. Sodium retention is the consequence of a homeostatic response caused by underfilling of the arterial circulation secondary to arterial vasodilation in the splanchnic vascular bed. Because the retained fluid is constantly leaking out of the intravascular compartment into the peritoneal cavity, the sensation of vascular filling is not achieved, and the process continues. Hypoalbuminemia and reduced plasma oncotic pressure also contribute to the loss of fluid from the vascular compartment into the

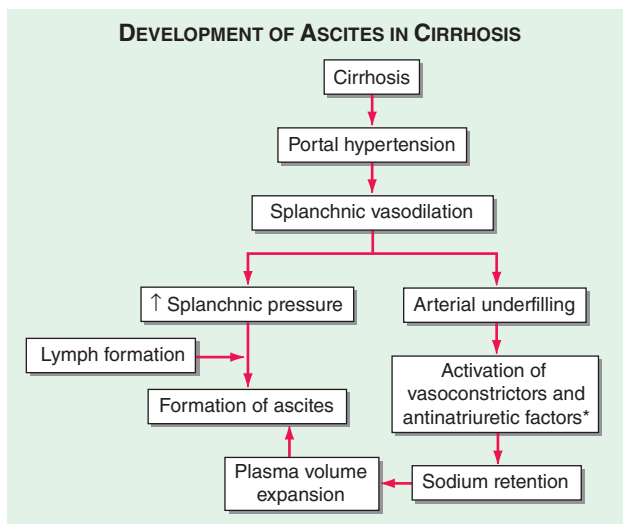


FIGURE 365-4 Development of ascites in cirrhosis. This flow diagram illustrates the importance of portal hypertension with splanchnic vasodilation in the development of ascites. *Antinatriuretic factors include the renin-angiotensin-aldosterone system and the sympathetic nervous system.

peritoneal cavity. Hypoalbuminemia is due to decreased synthetic function in a cirrhotic liver.

Clinical Features Patients typically note an increase in abdominal girth that is often accompanied by the development of peripheral edema. The development of ascites is often insidious, and it is surprising that some patients wait so long and become so distended before seeking medical attention. Patients usually have at least 1–2 L of fluid in the abdomen before they are aware that there is an increase. If ascitic fluid is massive, respiratory function can be compromised, and patients will complain of shortness of breath. Hepatic hydrothorax may also occur in this setting, contributing to respiratory symptoms. Patients with massive ascites are often malnourished and have muscle wasting and excessive fatigue and weakness.

Diagnosis Diagnosis of ascites is by physical examination and is often aided by abdominal imaging. Patients will have bulging flanks, may have a fluid wave, or may have the presence of shifting dullness. This is determined by taking patients from a supine position to lying on either their left or right side and noting the movement of the dullness to percussion. Subtle amounts of ascites can be detected by ultrasound or CT scanning. Hepatic hydrothorax is more common on the right side and implicates a rent in the diaphragm with free flow of ascitic fluid into the thoracic cavity.

When patients present with ascites for the first time, it is recommended that a diagnostic paracentesis be performed to characterize the fluid. This should include the determination of total protein and albumin content, blood cell counts with differential, and cultures. In the appropriate setting, amylase may be measured and cytology performed. In patients with cirrhosis, the protein concentration of the ascitic fluid is quite low, with the majority of patients having an ascitic fluid protein concentration <1 g/dL. The development of the serum ascites-to-albumin gradient (SAAG) has replaced the description of exudative or transudative fluid. When the gradient between the serum albumin level and the ascitic fluid albumin level is >1.1 g/dL, the cause of the ascites is most likely due to portal hypertension; this is usually in the setting of cirrhosis. When the gradient is <1.1 g/dL, infectious or malignant causes of ascites should be considered. When levels of ascitic fluid proteins are very low, patients are at increased risk for developing SBP. A high level of red blood cells in the ascitic fluid signifies a traumatic tap or perhaps a hepatocellular cancer or a ruptured omental varix. When the absolute level of polymorphonuclear leukocytes is >250/μL, the question of ascitic fluid infection should be strongly considered. Ascitic fluid cultures should be obtained using bedside inoculation of culture media.

TREATMENT ASCITES

Patients with small amounts of ascites can usually be managed with dietary sodium restriction alone. Most average diets in the United States contain 6–8 g of sodium per day, and if patients eat at restaurants or fast-food outlets, the amount of sodium in their diet can exceed this amount. Thus, it is often extremely difficult to get patients to change their dietary habits to ingest <2 g of sodium per day, which is the recommended amount. Patients are frequently surprised to realize how much sodium is in the standard U.S. diet; thus, it is important to make educational pamphlets available to the patient. Often, a simple recommendation is to eat fresh or frozen foods, avoiding canned or processed foods, which are usually preserved with sodium. When a moderate amount of ascites is present, diuretic therapy is usually necessary. Traditionally, spironolactone at 100–200 mg/d as a single dose is started, and furosemide may be added at 40–80 mg/d, particularly in patients who have peripheral edema. In patients who have never received diuretics before, the failure of the above-mentioned dosages suggests that they are not being compliant with a low-sodium diet. If compliance is confirmed and ascitic fluid is not being mobilized, spironolactone can be increased to 400–600 mg/d and furosemide increased to 120–160 mg/d. If ascites is still present with these dosages of diuretics in patients who are compliant with a low-sodium diet, then they are defined as having *refractory ascites*, and alternative treatment modalities including repeated large-volume paracentesis or a TIPS procedure should be considered (Fig. 365-5). Recent studies have shown that TIPS, while managing the ascites, does not improve survival in these patients. Unfortunately, TIPS is often associated with an increased frequency of hepatic encephalopathy and must be considered carefully on a case-by-case basis. The prognosis for patients with cirrhosis with ascites is poor, and some studies have shown that <50% of patients survive 2 years after the onset of ascites. Thus, there should be consideration for liver transplantation in patients with the onset of ascites.

SPONTANEOUS BACTERIAL PERITONITIS

SBP is a common and severe complication of ascites characterized by spontaneous infection of the ascitic fluid without an intraabdominal source. In patients with cirrhosis and ascites severe enough for hospitalization, SBP can occur in up to 30% of individuals and can have a 25% in-hospital mortality rate. Bacterial translocation is the presumed mechanism for development of SBP, with gut flora traversing the intestine

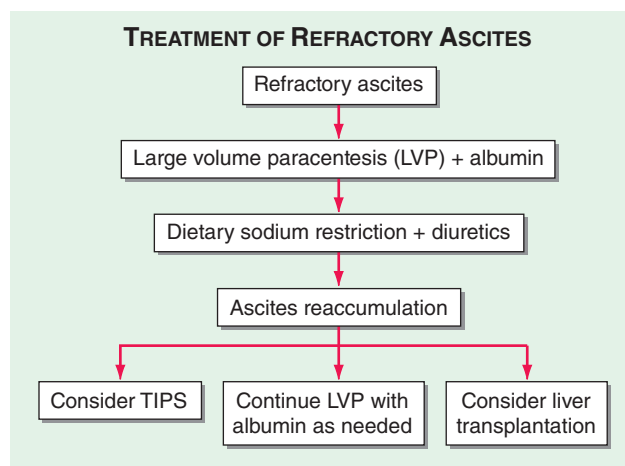


FIGURE 365-5 Treatment of refractory ascites. In patients who develop azotemia in the course of receiving diuretics in the management of their ascites, some will require repeated large-volume paracentesis (LVP), some may be considered for transjugular intrahepatic portosystemic shunt (TIPS), and some would be good candidates for liver transplantation. These decisions are all individualized.

into mesenteric lymph nodes, leading to bacteremia and seeding of the ascitic fluid. The most common organisms are *Escherichia coli* and other gut bacteria; however, gram-positive bacteria, including *Streptococcus viridans*, *Staphylococcus aureus*, and *Enterococcus* sp., can also be found. If more than two organisms are identified, secondary bacterial peritonitis due to a perforated viscus should be considered. The diagnosis of SBP is made when the fluid sample has an absolute neutrophil count $>250/\mu\text{L}$. Bedside cultures should be obtained when ascitic fluid is tapped. Patients with ascites may present with fever, altered mental status, elevated white blood cell count, and abdominal pain or discomfort, or they may present without any of these features. Therefore, it is necessary to have a high degree of clinical suspicion, and peritoneal taps are important for making the diagnosis. Treatment is with a second-generation cephalosporin, with cefotaxime being the most commonly used antibiotic. In patients with variceal hemorrhage, the frequency of SBP is significantly increased, and prophylaxis against SBP is recommended when a patient presents with upper GI bleeding. Furthermore, in patients who have had an episode(s) of SBP and recovered, once-weekly administration of antibiotics is used as prophylaxis for recurrent SBP.

HEPATORENAL SYNDROME

The hepatorenal syndrome (HRS) is a form of functional renal failure without renal pathology that occurs in about 10% of patients with advanced cirrhosis or acute liver failure. There are marked disturbances in the arterial renal circulation in patients with HRS; these include an increase in vascular resistance accompanied by a reduction in systemic vascular resistance. The reason for renal vasoconstriction is most likely multifactorial and is poorly understood. The diagnosis is made usually in the presence of a large amount of ascites in patients who have a stepwise progressive increase in creatinine. Type 1 HRS is characterized by a progressive impairment in renal function and a significant reduction in creatinine clearance within 1–2 weeks of presentation. Type 2 HRS is characterized by a reduction in glomerular filtration rate with an elevation of serum creatinine level, but it is fairly stable and is associated with a better outcome than that of type 1 HRS.

HRS is often seen in patients with refractory ascites and requires exclusion of other causes of acute renal failure. Treatment has, unfortunately, been difficult, and in the past, dopamine or prostaglandin analogues were used as renal vasodilating medications. Carefully performed studies have failed to show clear-cut benefit from these therapeutic approaches. Currently, patients are treated with midodrine, an α -agonist, along with octreotide and intravenous albumin. The best therapy for HRS is liver transplantation; recovery of renal function is typical in this setting. In patients with either type 1 or type 2 HRS, the prognosis is poor unless transplant can be achieved within a short period of time.

HEPATIC ENCEPHALOPATHY

Portosystemic encephalopathy is a serious complication of chronic liver disease and is broadly defined as an alteration in mental status and cognitive function occurring in the presence of liver failure. In acute liver injury with fulminant hepatic failure, the development of encephalopathy is a requirement for a diagnosis of fulminant failure. Encephalopathy is much more commonly seen in patients with chronic liver disease. Gut-derived neurotoxins that are not removed by the liver because of vascular shunting and decreased hepatic mass get to the brain and cause the symptoms that we know of as hepatic encephalopathy. Ammonia levels are typically elevated in patients with hepatic encephalopathy, but the correlation between severity of liver disease and height of ammonia levels is often poor, and most hepatologists do not rely on ammonia levels to make a diagnosis. Other compounds and metabolites that may contribute to the development of encephalopathy include certain false neurotransmitters and mercaptans.

Clinical Features In acute liver failure, changes in mental status can occur within weeks to months. Brain edema can be seen in these patients, with severe encephalopathy associated with swelling of the gray matter. Cerebral herniation is a feared complication of brain edema in acute liver failure, and treatment is meant to decrease edema with mannitol and judicious use of intravenous fluids.

In patients with cirrhosis, encephalopathy is often found as a result of certain precipitating events such as hypokalemia, infection, an increased dietary protein load, or electrolyte disturbances. Patients may be confused or exhibit a change in personality. They may actually be quite violent and difficult to manage; alternatively, patients may be very sleepy and difficult to rouse. Because precipitating events are so commonly found, they should be sought carefully. If patients have ascites, this should be tapped to rule out infection. Evidence of GI bleeding should be sought, and patients should be appropriately hydrated. Electrolytes should be measured and abnormalities corrected. In patients presenting with encephalopathy, asterixis is often present. Asterixis can be elicited by having patients extend their arms and bend their wrists back. In this maneuver, patients who are encephalopathic have a “liver flap”—i.e., a sudden forward movement of the wrist. This requires patients to be able to cooperate with the examiner and obviously cannot be elicited in patients who are severely encephalopathic or in hepatic coma.

The diagnosis of hepatic encephalopathy is clinical and requires an experienced clinician to recognize and put together all of the various features. Often when patients have encephalopathy for the first time, they are unaware of what is transpiring, but once they have been through the experience for the first time, they can identify when this is developing in subsequent situations and can often self-medicate to impair the development or worsening of encephalopathy.

TREATMENT HEPATIC ENCEPHALOPATHY

Treatment is multifactorial and includes management of the above-mentioned precipitating factors. Sometimes hydration and correction of electrolyte imbalance are all that is necessary. In the past, restriction of dietary protein was considered for patients with encephalopathy; however, the negative impact of that maneuver on overall nutrition is thought to outweigh the benefit when treating encephalopathy, and it is thus discouraged. There may be some benefit to replacing animal-based protein with vegetable-based protein in some patients with encephalopathy that is difficult to manage. The mainstay of treatment for encephalopathy, in addition to correcting precipitating factors, is to use lactulose, a nonabsorbable disaccharide, which results in colonic acidification. Catharsis ensues, contributing to the elimination of nitrogenous products in the gut that are responsible for the development of encephalopathy. The goal of lactulose therapy is to promote 2–3 soft stools per day. Patients are asked to titrate their amount of ingested lactulose to achieve the desired effect. Poorly absorbed antibiotics are often used as adjunctive therapies for patients who have had a difficult time with lactulose. The alternating administration of neomycin and metronidazole has commonly been used to reduce the individual side effects of each: neomycin for renal insufficiency and ototoxicity and metronidazole for peripheral neuropathy. More recently, rifaximin at 550 mg twice daily has been very effective in treating encephalopathy without the known side effects of neomycin or metronidazole. Zinc supplementation is sometimes helpful in patients with encephalopathy and is relatively harmless. The development of encephalopathy in patients with chronic liver disease is a poor prognostic sign, but this complication can be managed in the vast majority of patients.

MALNUTRITION IN CIRRHOSIS

Because the liver is principally involved in the regulation of protein and energy metabolism in the body, it is not surprising that patients with advanced liver disease are commonly malnourished. Once patients become cirrhotic, they are more catabolic, and muscle protein is metabolized. There are multiple factors that contribute to the malnutrition of cirrhosis, including poor dietary intake, alterations in gut nutrient absorption, and alterations in protein metabolism. Dietary supplementation for patients with cirrhosis is helpful in preventing patients from becoming catabolic.

ABNORMALITIES IN COAGULATION

Coagulopathy is almost universal in patients with cirrhosis. There is decreased synthesis of clotting factors and impaired clearance of anticoagulants. In addition, patients may have thrombocytopenia from hypersplenism due to portal hypertension. Vitamin K–dependent clotting factors are factors II, VII, IX, and X. Vitamin K requires biliary excretion for its subsequent absorption; thus, in patients with chronic cholestatic syndromes, vitamin K absorption is frequently diminished. Intravenous or intramuscular vitamin K can quickly correct this abnormality. More commonly, the synthesis of vitamin K–dependent clotting factors is diminished because of a decrease in hepatic mass, and, under these circumstances, administration of parenteral vitamin K does not improve the clotting factors or the prothrombin time. Platelet function is often abnormal in patients with chronic liver disease, in addition to decreases in platelet levels due to hypersplenism.

BONE DISEASE IN CIRRHOSIS

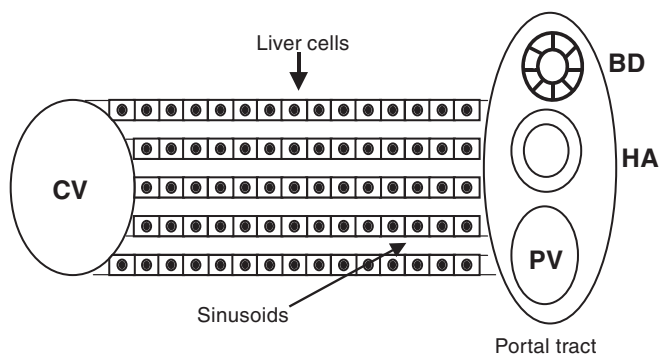
Osteoporosis is common in patients with chronic cholestatic liver disease because of malabsorption of vitamin D and decreased calcium ingestion. The rate of bone resorption exceeds that of new bone formation in patients with cirrhosis, resulting in bone loss. Dual x-ray absorptiometry (DEXA) is a useful method for determining osteoporosis or osteopenia in patients with chronic liver disease. When a DEXA scan shows decreased bone mass, treatment should be administered with bisphosphonates that are effective at inhibiting resorption of bone and efficacious in the treatment of osteoporosis.

HEMATOLOGIC ABNORMALITIES IN CIRRHOSIS

Numerous hematologic manifestations of cirrhosis are present, including anemia from a variety of causes including hypersplenism, hemolysis, iron deficiency, and perhaps folate deficiency from malnutrition. Macrocytosis is a common abnormality in red blood cell morphology seen in patients with chronic liver disease, and neutropenia may be seen as a result of hypersplenism.

Although clinical and laboratory features yield clues to the extent of inflammatory processes (disease grade), the degree of scarring and architectural distortion (disease stage), and the nature of the disease process, the liver biopsy is felt to represent the gold standard for assessing the degree of liver injury and fibrosis. Examination of liver histology provides not only a basis for quantitative scoring of disease activity and progression but also a wealth of qualitative information that can direct and inform diagnosis and management.

A normal liver lobule consists of portal (zone 1), lobular (midzonal or zone 2), and central (zone 3) zones. The portal tract contains the hepatic artery (HA) and portal vein (PV), which represent the dual vascular supply to the liver, as well as the bile duct (BD). The lobular area contains cords of liver cells surrounded by vascular sinusoids, and the central zone consists of the central vein (CV), the terminal branch of the hepatic vein (see figure below).



Included in this atlas of liver biopsies are examples of common morphologic features of acute and chronic liver disorders, some involving the lobular areas (e.g., the lobular inflammatory changes of acute hepatitis, apoptotic hepatocyte degeneration in acute and chronic hepatitis, virus antigen localization in hepatocyte cytoplasm

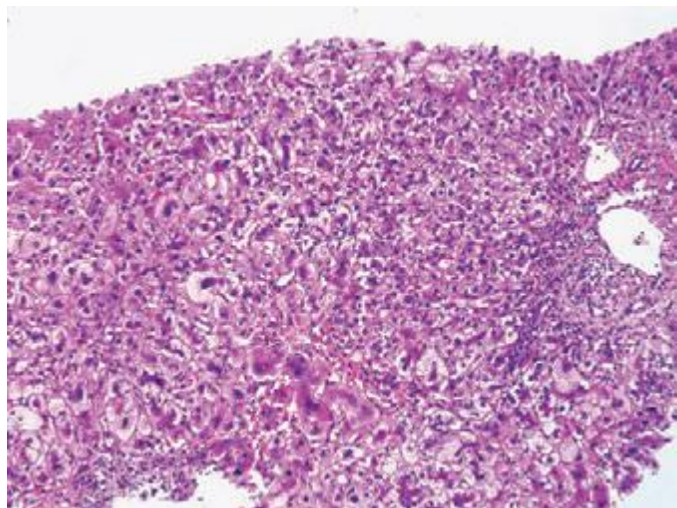


FIGURE 366e-1 Acute hepatitis with lobular inflammation and hepatocellular ballooning (hematoxylin and eosin [H&E], 10 \times).

and/or nuclei, viral inclusion bodies, copper or iron deposition, other inclusion bodies) and others involving the portal tracts (e.g., the portal mononuclear infiltrate that expands and spills over beyond the border of periportal hepatocytes in chronic hepatitis C, autoimmune hepatitis, and liver allograft rejection) or centrilobular areas (e.g., acute acetaminophen hepatotoxicity). Other histologic features of importance include hepatic steatosis (observed in alcoholic liver injury, nonalcoholic fatty liver disorders, and metabolic disorders—including mitochondrial injury—and in patients with chronic viral hepatitis); injury of bile ducts in the portal tract, an important diagnostic hallmark of primary biliary cirrhosis, primary sclerosing cholangitis, and liver allograft rejection; cholestasis in intrahepatic or extrahepatic biliary obstruction or in infiltrative disorders; ductular proliferation in the setting of marked hepatocellular necrosis; plasma cell infiltration common in autoimmune hepatitis; portal inflammation affecting portal veins (“endothelialitis”) in liver allograft rejection; and mild-to-severe fibrosis, in varying distribution and pattern, as a consequence of liver injury common to many disorders. (All magnifications reflect the objective lens used.)

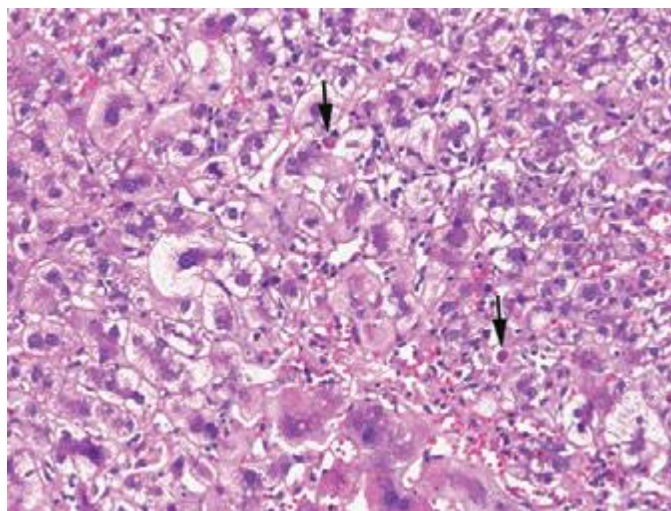


FIGURE 366e-2 Acute hepatitis, higher magnification, showing lobular inflammation, hepatocellular ballooning, and acidophilic bodies (arrows) (H&E, 20 \times).

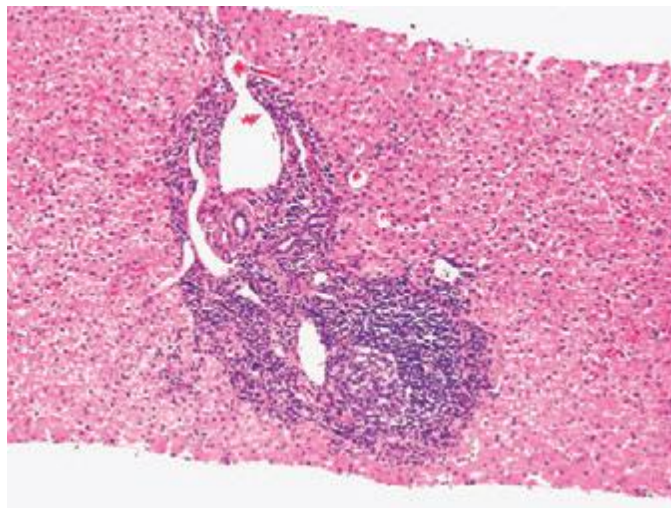


FIGURE 366e-3 Chronic hepatitis C with portal lymphoid infiltrate and lymphoid follicle containing germinal center (H&E, 10 \times).

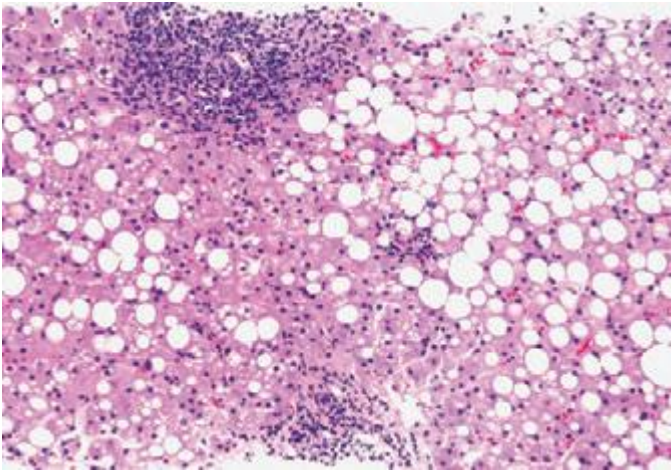


FIGURE 366e-4 Chronic hepatitis C with portal and lobular inflammation and steatosis (H&E, 10 \times).

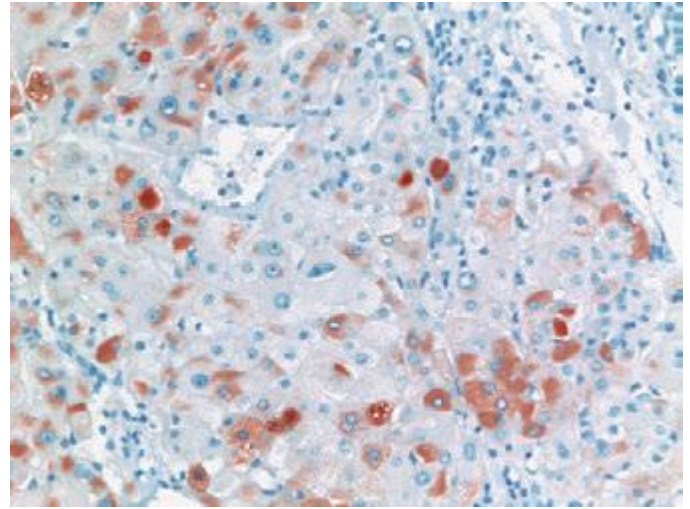


FIGURE 366e-7 Chronic hepatitis B with hepatocellular cytoplasmic staining for hepatitis B surface antigen (immunoperoxidase, 20 \times).

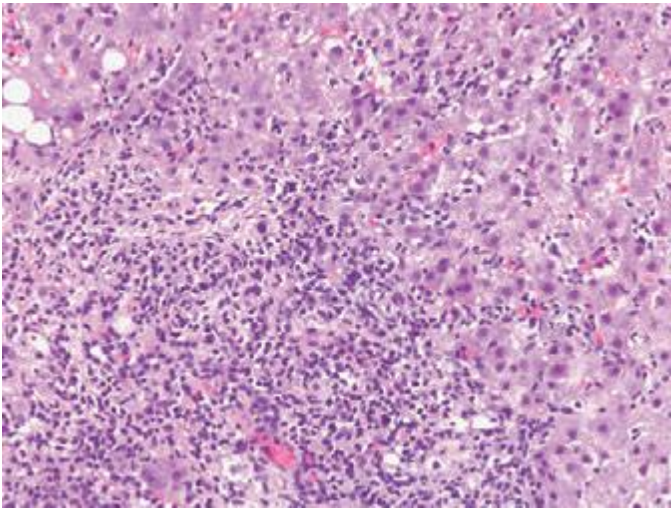


FIGURE 366e-5 Chronic hepatitis C with portal inflammation and interface hepatitis (erosion of the limiting plate of periportal hepatocytes by infiltrating mononuclear cells) (H&E, 20 \times).

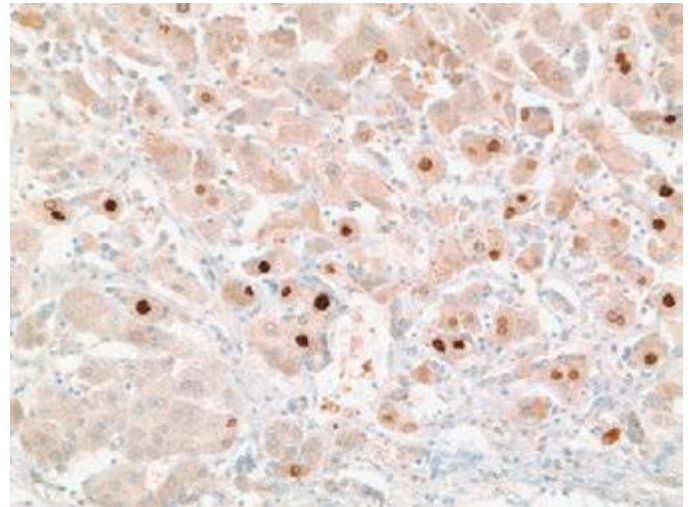


FIGURE 366e-8 Chronic hepatitis B with hepatocellular nuclear staining for hepatitis B core antigen (immunoperoxidase, 20 \times).

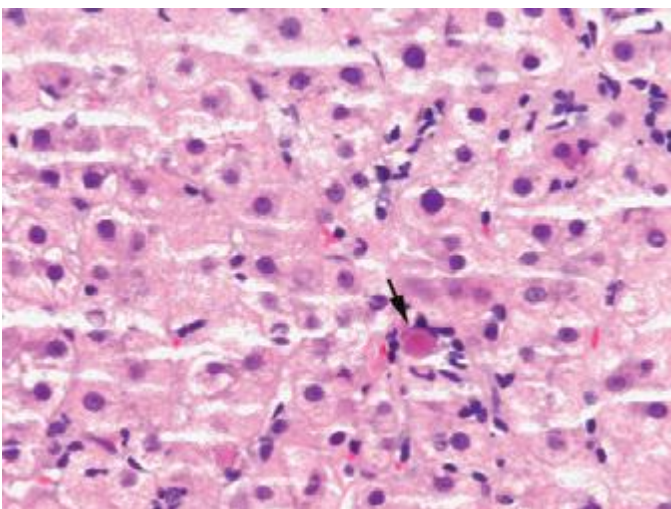


FIGURE 366e-6 Lobular inflammation with acidophilic body (apoptotic body) surrounded by lymphoid cells (H&E, 40 \times).

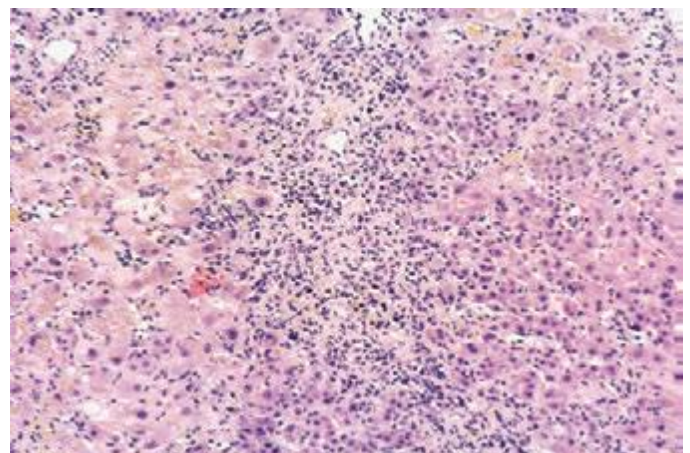


FIGURE 366e-9 Autoimmune hepatitis with portal and lobular inflammation, interface hepatitis, and cholestasis (H&E, 10 \times).

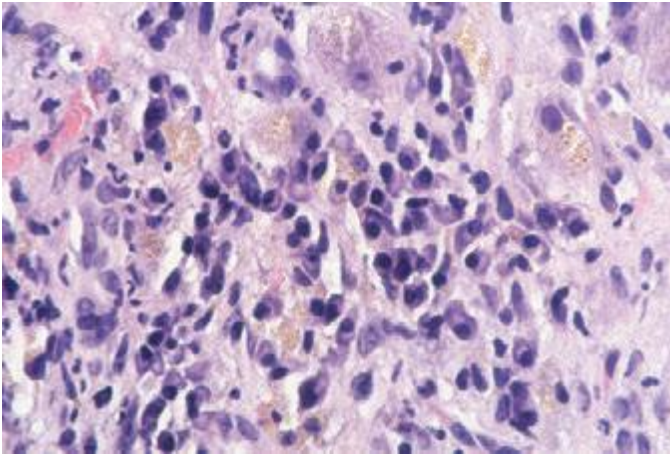


FIGURE 366e-10 Autoimmune hepatitis, higher magnification, showing dense plasma cell infiltrate in the portal and periportal regions (H&E, 40 \times).

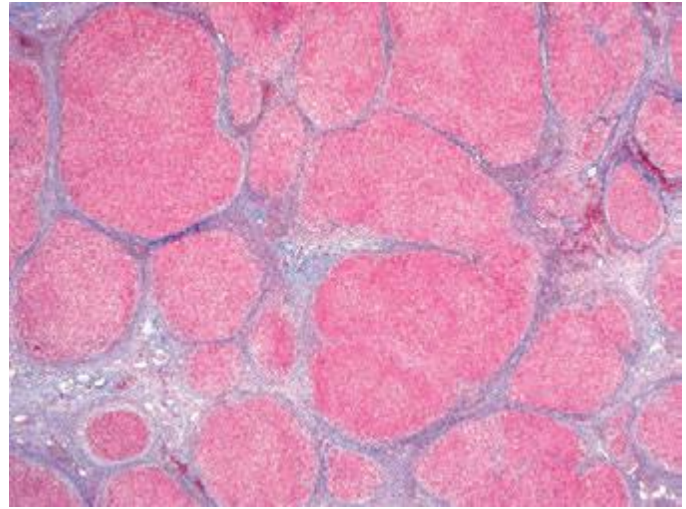


FIGURE 366e-13 Cirrhosis with architectural alteration resulting from fibrosis and nodular hepatocellular regeneration (Masson trichrome, 2 \times).

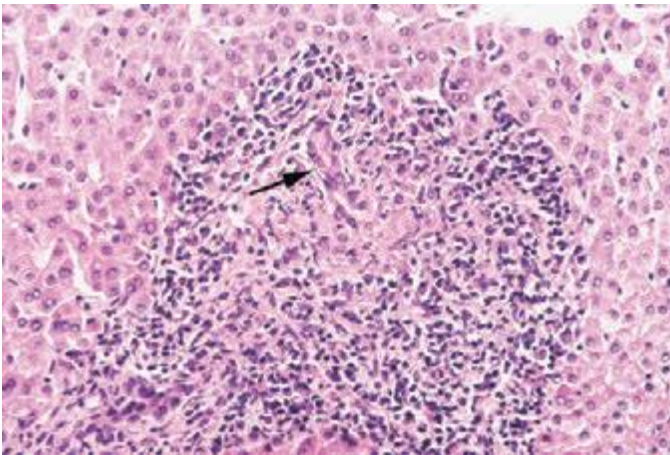


FIGURE 366e-11 Primary biliary cirrhosis with degenerating bile duct epithelium (“florid ductular lesion”) (*arrow*) surrounded by epithelioid granulomatous reaction and lymphoplasmacytic infiltrate (H&E, 40 \times).

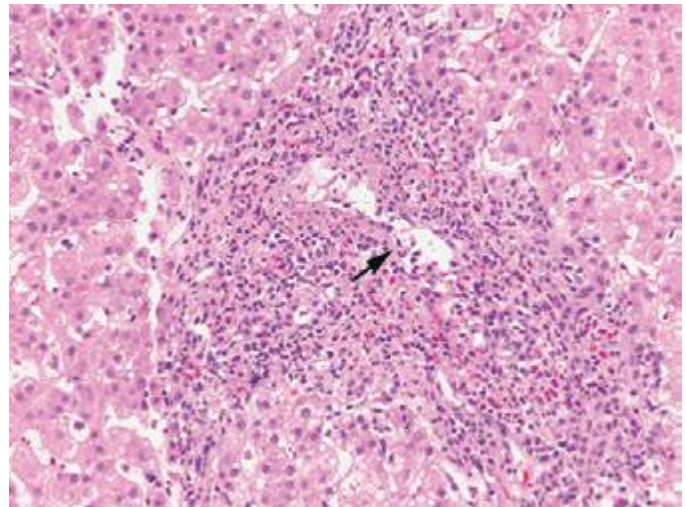


FIGURE 366e-14 Acute cellular rejection of orthotopic liver allograft demonstrating a mixed inflammatory cell infiltrate (lymphoid cells, eosinophils, neutrophils) of the portal tract as well as endothelialitis of the portal vein (*arrow*) and bile duct injury (H&E, 10 \times).

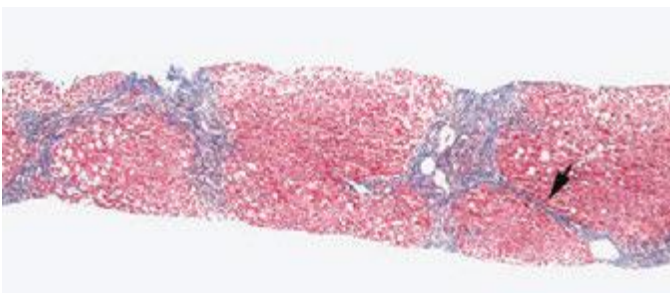


FIGURE 366e-12 Chronic hepatitis C with bridging fibrosis (*arrow*) (Masson trichrome, 10 \times).

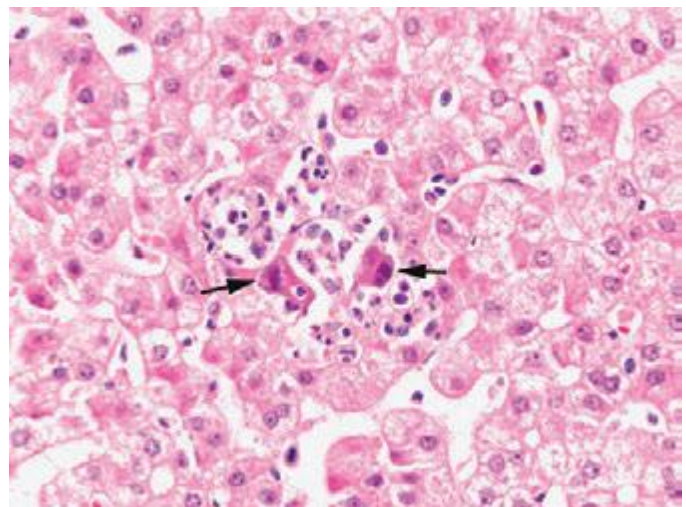


FIGURE 366e-15 Liver allograft with cytomegalovirus infection showing hepatocytes with nuclear inclusions (*arrows*) surrounded by a neutrophilic and lymphoid infiltrate (H&E, 10 \times).

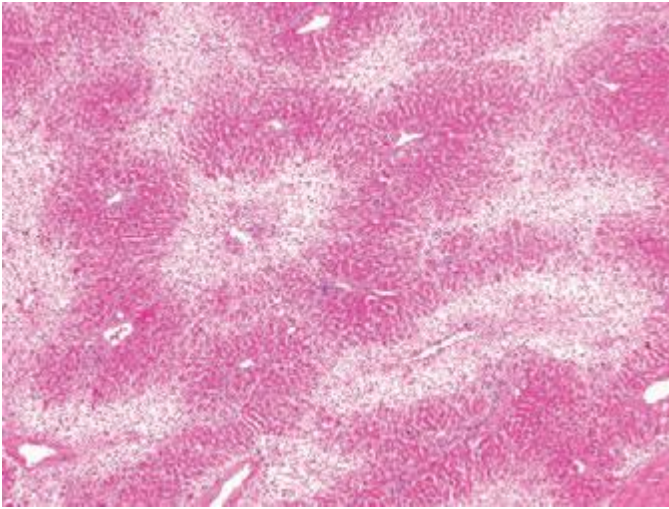


FIGURE 366e-16 Combined acetaminophen hepatotoxicity and alcoholic liver injury with extensive centrilobular areas of necrosis (H&E, 4 \times).

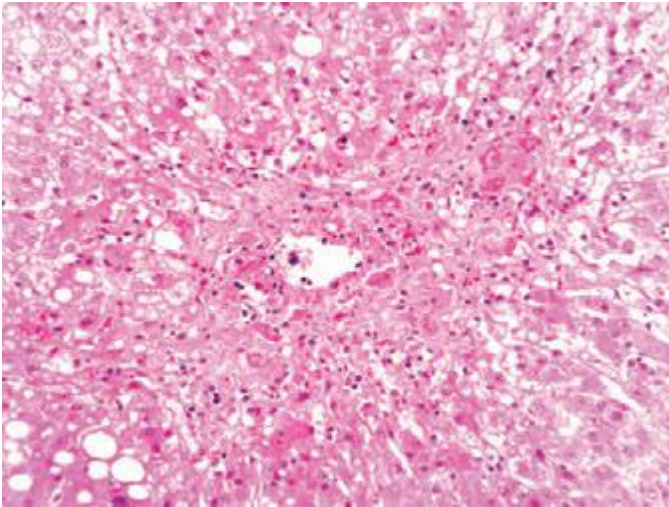


FIGURE 366e-17 Combined acetaminophen hepatotoxicity and alcoholic liver injury at higher magnification showing necrotic centrilobular area with Mallory bodies (H&E 20 \times).

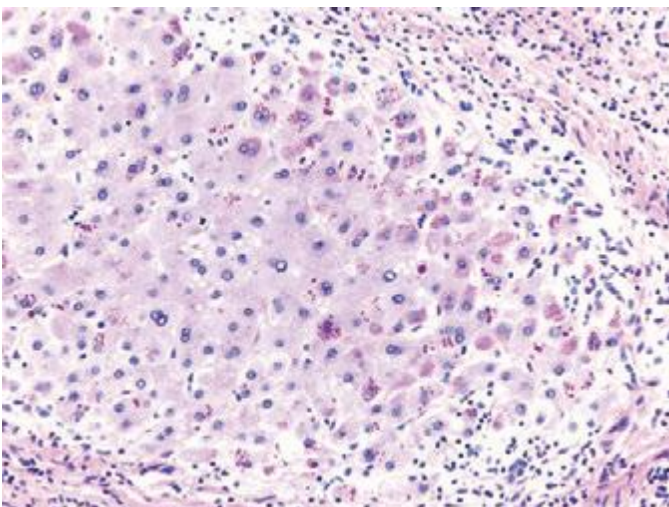


FIGURE 366e-18 α_1 Antitrypsin deficiency with cytoplasmic periodic acid–Schiff (PAS)-positive, diastase-resistant globules in many hepatocytes, predominantly at the periphery of a cirrhotic nodule (PAS, 20 \times).

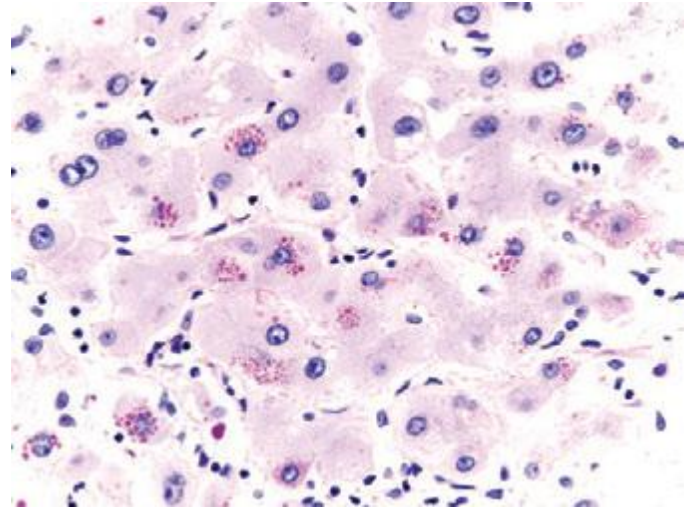


FIGURE 366e-19 α_1 Antitrypsin deficiency with higher magnification of PAS-positive, diastase-resistant globules (PAS, 40 \times).

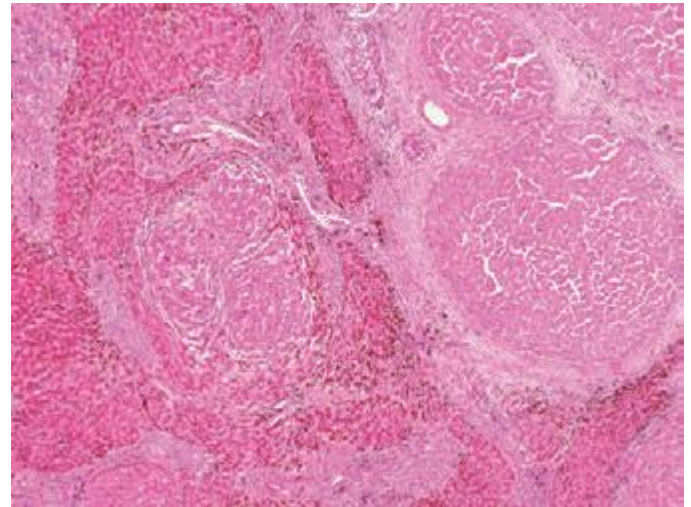


FIGURE 366e-20 Cirrhosis secondary to hemochromatosis with hepatocellular carcinoma; brown hemosiderin pigment (iron) is present in the cirrhotic liver, while the hepatocellular carcinoma nodules are hemosiderin-free (H&E, 4 \times).

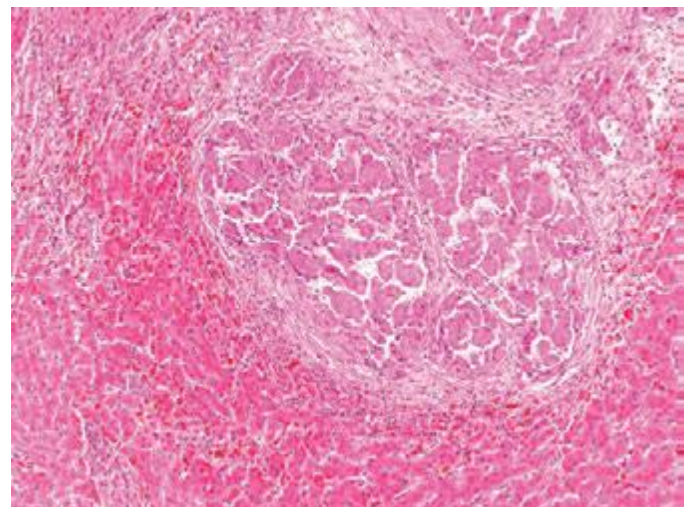


FIGURE 366e-21 Cirrhosis secondary to hemochromatosis with hepatocellular carcinoma at higher magnification, demonstrating nodules of large malignant cells with highly disorganized architecture (H&E, 10 \times).

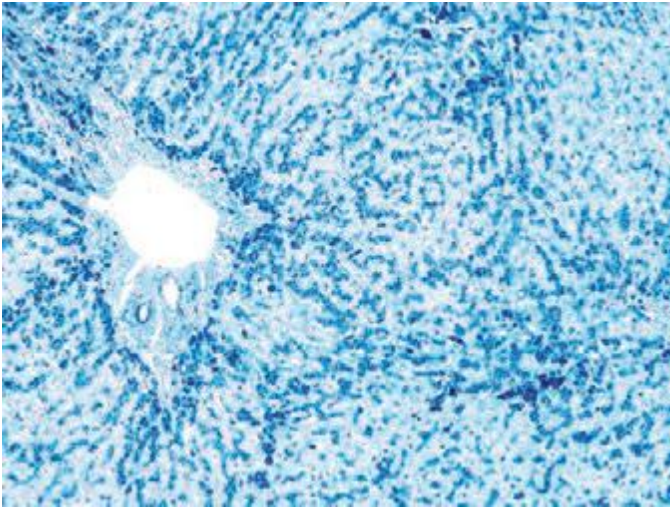


FIGURE 366e-22 Hemochromatosis with iron stain demonstrating extensive iron deposition and characteristic pattern of pericanalicular distribution of iron (iron stain, 10 \times).

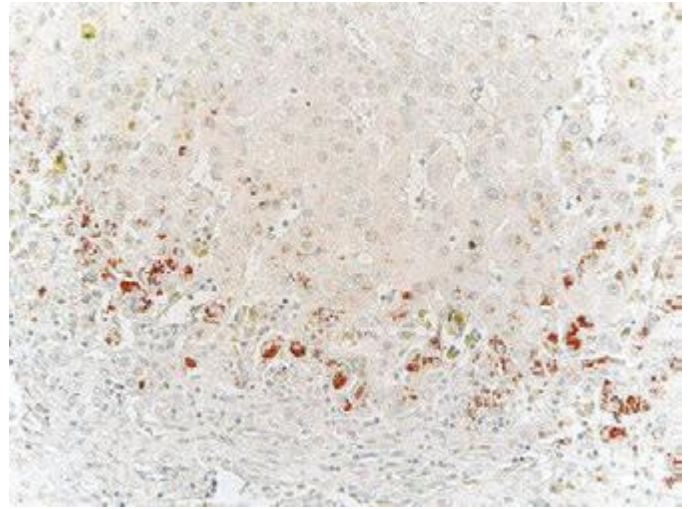


FIGURE 366e-25 Primary sclerosing cholangitis showing peripheral cholestasis (green) and cytoplasmic red granular staining of hepatocytes for copper (rhodamine copper stain, 20 \times).

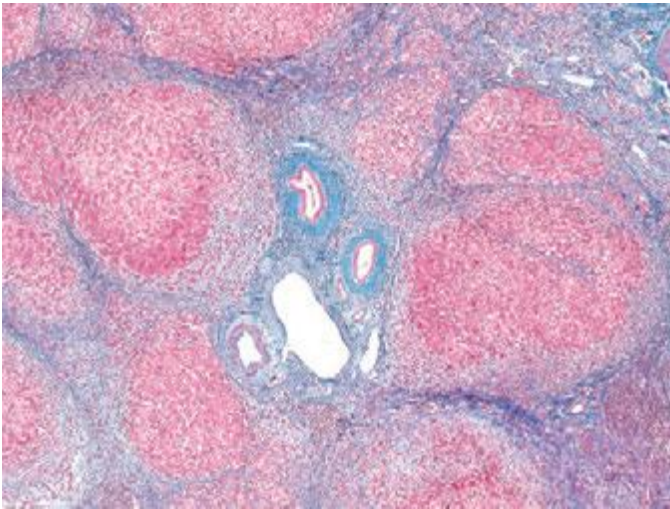


FIGURE 366e-23 Primary sclerosing cholangitis showing cirrhosis and periductular fibrosis (Masson trichrome, 4 \times).

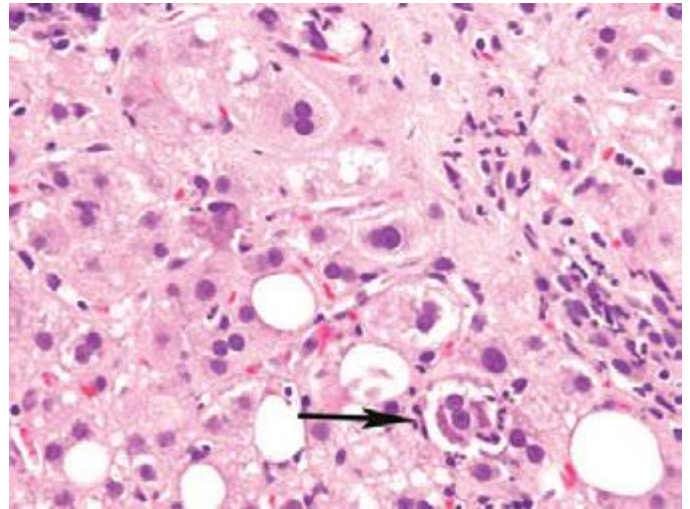


FIGURE 366e-26 Nonalcoholic steatohepatitis (NASH) showing steatosis, ballooned hepatocytes, and Mallory bodies with surrounding polymorphonuclear leukocytes (arrow) (H&E, 20 \times).

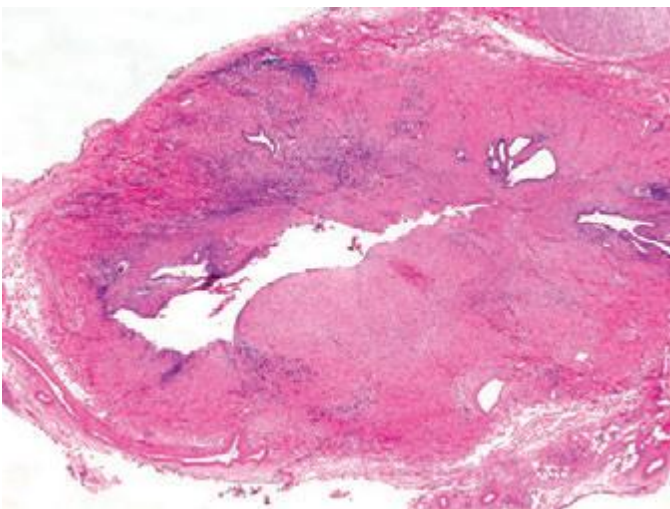


FIGURE 366e-24 Primary sclerosing cholangitis showing the extrahepatic bile duct (in a liver explant obtained at the time of hepatectomy for orthotopic liver transplantation) with marked mural chronic inflammation and fibrosis as well as peribiliary glands (H&E, 2 \times).

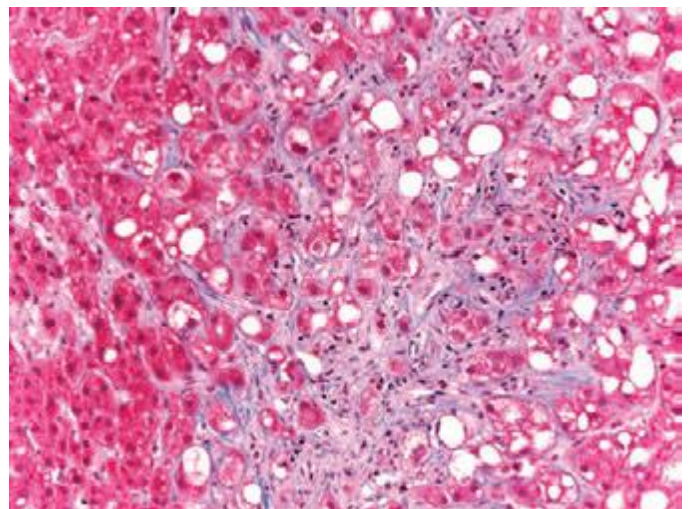


FIGURE 366e-27 Nonalcoholic steatohepatitis (NASH) showing steatosis with perisinusoidal and pericellular fibrosis (H&E, 20 \times).

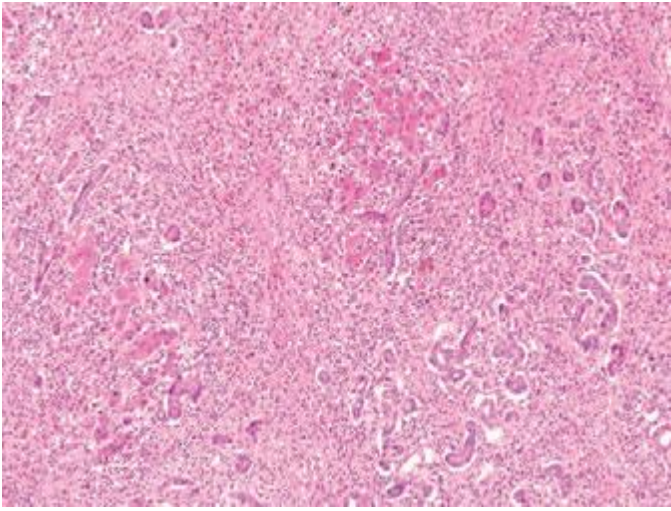


FIGURE 366e-28 Acute hepatitis with submassive hepatic necrosis with marked parenchymal collapse, remnant islands of surviving hepatocytes, and a marked ductular reaction (H&E, 10 \times).

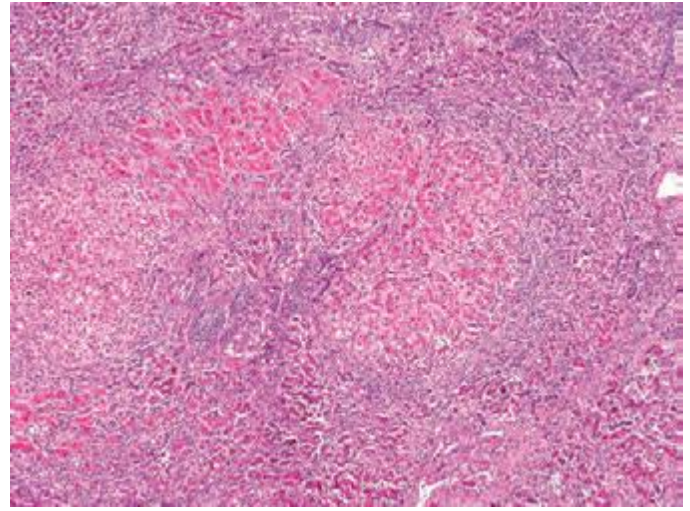


FIGURE 366e-29 Wilson's disease showing cirrhosis, extensive collapse, and ductular reaction in a teenager with an acute presentation (H&E, 4 \times).

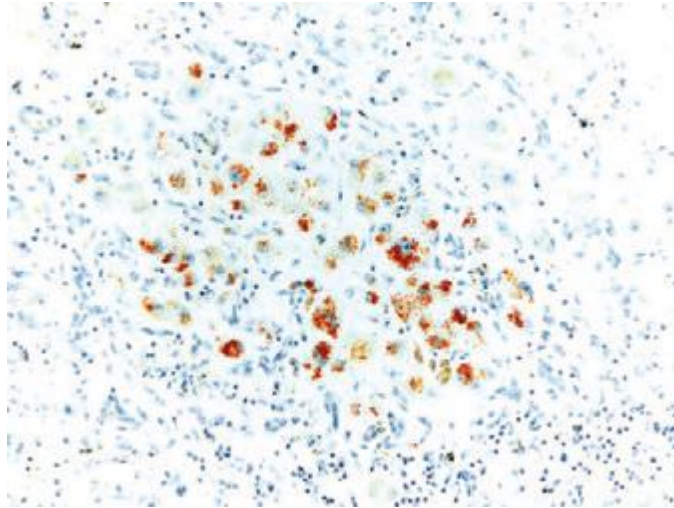


FIGURE 366e-30 Wilson's disease showing extensive hepatocyte cytoplasmic red granular staining for copper in a cirrhotic nodule (rhodanine copper stain, 20 \times).

367e Genetic, Metabolic, and Infiltrative Diseases Affecting the Liver

Bruce R. Bacon

There are a number of disorders of the liver that fit within the categories of genetic, metabolic, and infiltrative disorders (Table 367e-1). Inherited disorders include hemochromatosis, Wilson's disease, α_1 antitrypsin (α_1 AT) deficiency, and cystic fibrosis (CF). Hemochromatosis is the most common inherited disorder affecting white populations, with the genetic susceptibility for the disease being identified in 1 in 250 individuals. Over the past 15 years, it has become increasingly apparent that nonalcoholic fatty liver disease (NAFLD) is the most common cause of elevated liver enzymes found in the U.S. population. This disorder is discussed in greater detail in Chap. 364. Infiltrative disorders of the liver are relatively rare.

GENETIC LIVER DISEASES

Hereditary Hemochromatosis Hereditary hemochromatosis (HH) is a common inherited disorder of iron metabolism (Chap. 428). Our knowledge of the disease and its phenotypic expression has changed since 1996, when the gene for HH, called *HFE*, was identified, allowing for genetic testing for the two major mutations (C282Y and H63D) that are responsible for *HFE*-related HH. Subsequently, several additional genes/proteins involved in the regulation of iron homeostasis have been identified, contributing to a better understanding of cellular iron uptake and release and the characterization of additional causes of inherited iron overload (Table 367e-2).

Most patients with HH are asymptomatic; however, when patients present with symptoms, they are frequently nonspecific and include weakness, fatigue, lethargy, and weight loss. Specific, organ-related symptoms include abdominal pain, arthralgias, and symptoms and signs of chronic liver disease. Increasingly, most patients are now identified before they have symptoms, either through family studies or from the performance of screening iron studies. Several prospective population studies have shown that C282Y homozygosity is found in about 1 in 250 individuals of northern European descent, with the heterozygote frequency seen in approximately 1 in 10 individuals. It is important to consider HH in patients who present with the symptoms

TABLE 367e-1 GENETIC, METABOLIC, AND INFILTRATIVE DISEASES AFFECTING THE LIVER

Genetic
• Hereditary hemochromatosis
• Wilson's disease
• α_1 Antitrypsin deficiency
• Cystic fibrosis
Metabolic
• Nonalcoholic fatty liver disease
• Lipid storage diseases
• Gaucher's
• Niemann-Pick
• Tangier
• Fabry's
• Porphyrias
• Porphyria cutanea tarda
Infiltrative disorders
• Amyloidosis
• Granulomas
• Sarcoidosis
• Lymphoma

TABLE 367e-2 CLASSIFICATION OF IRON OVERLOAD SYNDROMES

Hereditary Hemochromatosis (HH)

<i>HFE</i> -related (type 1)
C282Y/C282Y
C282Y/H63D
Other <i>HFE</i> mutations
Non- <i>HFE</i> -related
Juvenile HH
HJV—hemojuvelin (type 2a)
HAMP—hepcidin (type 2b)
TfR2-related HH (type 3)
Ferroportin-related HH (type 4)
African iron overload

Secondary Iron Overload

Iron-loading anemias
Parenteral iron overload
Chronic liver disease

Miscellaneous

Neonatal iron overload
Aceruloplasminemia
Congenital atransferrinemia

Abbreviations: HAMP, hepcidin; HJV, hemojuvelin; TfR2, transferrin receptor 2.

and signs known to occur in established HH. When confronted with abnormal serum iron studies, clinicians should not wait for typical symptoms or findings of HH to appear before considering the diagnosis. However, once the diagnosis of HH is considered, either by an evaluation of abnormal screening iron studies in the context of family studies, in a patient with an abnormal genetic test, or in the evaluation of a patient with any of the typical symptoms (Table 367e-3) or clinical findings (Table 367e-4), definitive diagnosis is relatively straightforward. Transferrin saturation (serum iron divided by total iron-binding capacity [TIBC] or transferrin, times 100%) and ferritin levels should be obtained. Both of these will be elevated in a symptomatic patient. It must be remembered that ferritin is an acute-phase reactant and can be elevated in a number of other inflammatory disorders, such as rheumatoid arthritis, or in various neoplastic diseases, such as lymphoma or other cancers. Also, serum ferritin is elevated in a majority of patients with nonalcoholic steatohepatitis (NASH), hepatitis C, and alcoholic liver disease in the absence of iron overload.

At present, if patients have an elevated transferrin saturation or ferritin level, genetic testing should be performed; if they are a C282Y homozygote or a compound heterozygote (C282Y/H63D), the diagnosis is confirmed. If liver enzymes (alanine aminotransferase [ALT], aspartate aminotransferase [AST]) are elevated or the ferritin is >1000 $\mu\text{g/L}$, the patient should be considered for liver biopsy because there is an increased frequency of advanced fibrosis in these individuals. If liver biopsy is performed, iron deposition is found in a periportal distribution with a periportal to pericentral gradient; iron is found predominantly in parenchymal cells, and Kupffer cells are spared.

TABLE 367e-3 SYMPTOMS OF HEREDITARY HEMOCHROMATOSIS

Symptom	%
Weakness, lethargy, fatigue	40–85
Apathy, lack of interest	40–85
Abdominal pain	30–60
Weight loss	30–60
Arthralgias	40–60
Loss of libido, impotence	30–60
Amenorrhea	20–60
Congestive heart failure symptoms	0–40

TABLE 367e-4 PHYSICAL FINDINGS IN HEREDITARY HEMOCHROMATOSIS

Finding	%
Hepatomegaly	60–85
Cirrhosis	50–95
Skin pigmentation	40–80
Arthritis (second, third metacarpophalangeal joints)	40–60
Clinical diabetes	10–60
Splenomegaly	10–40
Loss of body hair	10–30
Testicular atrophy	10–30
Dilated cardiomyopathy	0–30

TREATMENT HEREDITARY HEMOCHROMATOSIS

Treatment of HH is relatively straightforward with weekly phlebotomy aimed to reduce iron stores, recognizing that each unit of blood contains 200–250 mg of iron. If patients are diagnosed and treated before the development of hepatic fibrosis, all complications of the disease can be avoided. Maintenance phlebotomy is required in most patients and usually can be achieved with 1 unit of blood removed every 2–3 months. Family studies should be performed with transferrin saturation, ferritin, and genetic testing offered to all first-degree relatives.

Wilson's Disease Wilson's disease is an inherited disorder of copper homeostasis first described in 1912 (Chap. 429). The Wilson's disease gene was discovered in 1993, with the identification of *ATP7B*. This P-type ATPase is involved in copper transport and is necessary for the export of copper from the hepatocyte. Thus, in patients with mutations in *ATP7B*, copper is retained in the liver, leading to increased copper storage and ultimately liver disease as a result.

The clinical presentation of Wilson's disease is variable and includes chronic hepatitis, hepatic steatosis, and cirrhosis in adolescents and young adults. Neurologic manifestations indicate that liver disease is present and include speech disorders and various movement disorders. Diagnosis includes the demonstration of a reduced ceruloplasmin level, increased urinary excretion of copper, the presence of Kayser-Fleischer rings in the corneas of the eyes, and an elevated hepatic copper level, in the appropriate clinical setting. The genetic diagnosis of Wilson's disease is difficult because >500 mutations in *ATP7B* have been described with different degrees of frequency and penetration in certain populations.

TREATMENT WILSON'S DISEASE

Treatment consists of copper-chelating medications such as D-penicillamine and trientine. A role for zinc acetate has also been established. Medical treatment is lifelong, and severe relapses leading to liver failure and death can occur with cessation of therapy. Liver transplantation is curative with respect to the underlying metabolic defect and restores the normal phenotype with respect to copper homeostasis.

α_1 Antitrypsin Deficiency α_1 AT deficiency was first described in the late 1960s in patients with severe pulmonary disease. α_1 AT is a 52-kDa glycoprotein produced in hepatocytes, phagocytes, and epithelial cells in the lungs, which inhibits serine proteases, primarily neutrophil elastase. In α_1 AT deficiency, increased amounts of neutrophil elastase can result in progressive lung injury from degradation of elastin, leading to premature emphysema. In the 1970s, α_1 AT deficiency was discovered as a cause of neonatal liver disease, so-called "neonatal hepatitis." It is now known to be a cause of liver disease in infancy, early childhood, and adolescence, and in adults.

In α_1 AT deficiency, variants in the proteinase inhibitor (Pi) gene located on chromosome 14 alter α_1 AT structure, interfering with hepatocellular export. Aggregated, deformed polymers of α_1 AT accumulate in the hepatocyte endoplasmic reticulum. There are over 75 different α_1 AT variants. Conventional nomenclature identifies normal variants as PiMM; these individuals have normal blood levels of α_1 AT. The most common abnormal variants are called S and Z. Individuals homozygous for the Z mutation (PiZZ) have low levels of α_1 AT (about 15% of normal), and these patients are susceptible to liver and/or lung disease, yet only a proportion (about 25%) of PiZZ patients develop disease manifestations. Null variants have undetectable levels of α_1 AT and are susceptible to premature lung disease.

α_1 AT deficiency has been identified in all populations; however, the disorder is most common in patients of northern European and Iberian descent. The disorder affects about 1 in 1500 to 2000 individuals in North America. The natural history of α_1 AT deficiency is quite variable because many individuals with the PiZZ variant never develop disease, whereas others can develop childhood cirrhosis leading to liver transplantation.

In adults, the diagnosis often comes in the course of evaluation of patients with abnormal liver test abnormalities or in a workup for cirrhosis. A hint to diagnosis may be coexistent lung disease at a relatively young age or a family history of liver and/or lung disease. Patients may have symptoms of pulmonary disease with cough and dyspnea. Liver disease may be asymptomatic other than fatigue, or patients may present with complications of decompensated liver disease.

Diagnosis of α_1 AT deficiency is confirmed by blood tests showing reduced levels of serum α_1 AT, accompanied by Pi determinations. Most patients with liver disease have either PiZZ or PiSZ; occasionally, patients with PiMZ have reduced levels of α_1 AT, but they usually do not have a low enough level to cause disease. Liver biopsy is often performed to determine stage of hepatic fibrosis and shows characteristic PAS-positive, diastase-resistant globules in the periphery of the hepatic lobule.

TREATMENT α_1 ANTITRYPSIN DEFICIENCY

Treatment of α_1 AT deficiency is usually nonspecific and supportive. For patients with liver involvement, other sources of liver injury, such as alcohol, should be avoided. Evidence for other liver diseases (e.g., viral hepatitis B and C, hemochromatosis, NAFLD) should be sought and treated if possible. Smoking can worsen lung disease progression in α_1 AT deficiency and should be discontinued. Patients with lung disease may be eligible to receive infusions of α_1 AT, which has been shown to halt further damage to the lungs. If liver disease becomes decompensated, transplantation should be pursued and is curative. Following transplant, patients express the Pi phenotype of the donor. Finally, risk of hepatocellular carcinoma is significantly increased in patients with cirrhosis due to α_1 AT deficiency.

Cystic Fibrosis CF should also be considered as an inherited form of chronic liver disease, although the principal manifestations of CF include chronic lung disease and pancreatic insufficiency (Chap. 313). A small percentage of patients with CF who survive to adulthood have a form of biliary cirrhosis characterized by cholestatic liver enzyme abnormalities and the development of chronic liver disease. Ursodeoxycholic acid is occasionally helpful in improving liver test abnormalities and in reducing symptoms. The disease is slowly progressive.

METABOLIC LIVER DISEASES

Nonalcoholic Fatty Liver Disease NAFLD and NASH are common liver diseases causing abnormal liver test results and progressing to cirrhosis. **NAFLD and NASH are discussed in detail in Chap. 364.**

Lipid Storage Diseases There are a number of rare lipid storage diseases that involve the liver, including the inherited disorders of Gaucher's disease and Niemann-Pick disease (Chap. 433e). Other rare disorders

TABLE 367e-5 LIPID STORAGE DISEASES

Gaucher's
Niemann-Pick
Abetalipoproteinemia
Tangier
Fabry's
Types I and IV hyperlipoproteinemia

include abetalipoproteinemia, Tangier disease, Fabry's disease, and types I and V hyperlipoproteinemia (Table 367e-5). Hepatomegaly is present due to increased fat deposition, and increased glycogen is found in the liver.

Porphyrias The porphyrias are a group of metabolic disorders in which there are defects in the biosynthesis of heme necessary for incorporation into numerous hemoproteins such as hemoglobin, myoglobin, catalase, and the cytochromes (Chap. 430). Porphyrias can present as either acute or chronic diseases, with the acute disorder causing recurring bouts of abdominal pain, and the chronic disorders characterized by painful skin lesions. Porphyria cutanea tarda (PCT) is the most commonly encountered porphyria. Patients present with characteristic vesicular lesions on sun-exposed areas of the skin, principally the dorsum of the hands, the tips of the ears, or the cheeks. About 40% of patients with PCT have mutations in the gene for hemochromatosis (*HFE*), and ~50% have hepatitis C; thus, iron studies and *HFE* mutation analysis as well as hepatitis C testing should be considered in all patients who present with PCT. PCT is also associated with excess alcohol use and some medications, most notably estrogens.

TREATMENT PORPHYRIAS

The mainstay of treatment of PCT is iron reduction by therapeutic phlebotomy, which is successful in reversing the skin lesions in the majority of patients. If hepatitis C is present, this should be treated as well. Acute intermittent porphyria presents with abdominal pain, with the diagnosis made by avoidance of certain precipitating factors such as starvation or certain diets. Intravenous heme as hematin has been used for treatment.

INFILTRATIVE DISORDERS

Amyloidosis Amyloidosis is a metabolic storage disease that results from deposition of insoluble proteins that are aberrantly folded and assembled and then deposited in a variety of tissues (Chap. 136).

Amyloidosis is divided into two types, primary and secondary, based on the broad concepts of association with myeloma (primary) or chronic inflammatory illnesses (secondary). The disease is generally considered rare, although, in certain disease states or in certain populations, it can be more common. For example, when associated with familial Mediterranean fever, it is seen in high frequency in Sephardic Jews and Armenians living in Armenia and less frequently in Ashkenazi Jews, Turks, and Arabs. Amyloidosis frequently affects patients suffering from tuberculosis and leprosy and can be seen in upwards of 10–15% of patients with ankylosing spondylitis, rheumatoid arthritis, or Crohn's disease. In one surgical pathology series, amyloid was found in <1% of cases. The liver is commonly involved in cases of systemic amyloidosis, but it is frequently not clinically apparent and only documented at autopsy. Pathologic findings in the liver include positive staining with the Congo red histochemical stain where there is an apple-green birefringence noted under polarizing light.

Granulomas Granulomas are frequently found in the liver when patients are being evaluated for cholestatic liver enzyme abnormalities. Granulomas can be seen in primary biliary cirrhosis, but there are other characteristic clinical (e.g., pruritus, fatigue) and laboratory findings (cholestatic liver tests, antimitochondrial antibody) that allow for a definitive diagnosis of that disorder. Granulomatous infiltration can also be seen as the principal hepatic manifestation of sarcoidosis, and this is the most common presentation of hepatic granulomas (Chap. 390). The vast majority of these patients do not require any specific treatment other than what would normally be used for treatment of their sarcoidosis. A small subset, however, can develop a particularly bothersome desmoplastic reaction with a significant increase in fibrosis, which can progress to cirrhosis and liver failure. These patients may require treatment with immunosuppressive therapy and may require liver transplantation. In patients who have granulomas in the liver not associated with sarcoidosis, treatment is rarely needed.

Diagnosis requires liver biopsy, and it is important to establish a diagnosis so that a cause for the elevated liver enzymes is carefully identified. Some medications can cause granulomatous infiltration of the liver, the most notable of which is allopurinol.

Lymphoma Involvement of the liver with lymphoma can sometimes be with bulky mass lesions but can also be as a difficult-to-diagnose infiltrative disorder that does not show any characteristic findings on abdominal imaging studies (Chap. 134). Patients may present with severe liver disease, jaundice, hypoalbuminemia, mild to moderately elevated aminotransferases, and an elevated alkaline phosphatase.

A liver biopsy is required for diagnosis and should be considered when routine blood testing does not lead to a diagnosis of the liver dysfunction.

368 Liver Transplantation

Raymond T. Chung, Jules L. Dienstag

Liver transplantation—the replacement of the native, diseased liver by a normal organ (allograft)—has matured from an experimental procedure reserved for desperately ill patients to an accepted, lifesaving operation applied more optimally in the natural history of end-stage liver disease. The preferred and technically most advanced approach is *orthotopic transplantation*, in which the native organ is removed and the donor organ is inserted in the same anatomic location. Pioneered in the 1960s by Thomas Starzl at the University of Colorado and, later, at the University of Pittsburgh and by Roy Calne in Cambridge, England, liver transplantation is now performed routinely worldwide. Success measured as 1-year survival has improved from ~30% in the 1970s to >90% today. These improved prospects for prolonged survival resulted from refinements in operative technique, improvements in organ procurement and preservation, advances in immunosuppressive therapy, and, perhaps most influentially, more enlightened patient selection and timing. Despite the perioperative morbidity and mortality, the technical and management challenges of the procedure, and its costs, liver transplantation has become the approach of choice for selected patients whose chronic or acute liver disease is progressive, life-threatening, and unresponsive to medical therapy. Based on the current level of success, the number of liver transplants has continued to grow each year; in 2012, 6256 patients received liver allografts in the United States. Still, the demand for new livers continues to outpace availability; as of mid-2013, 15,806 patients in the United States were on a waiting list for a donor liver. In response to this drastic shortage of donor organs, many transplantation centers supplement cadaver-organ liver transplantation with living-donor transplantation.

INDICATIONS

Potential candidates for liver transplantation are children and adults who, in the absence of contraindications (see below), suffer from severe, irreversible liver disease for which alternative medical or surgical treatments have been exhausted or are unavailable. *Timing of the operation is of critical importance.* Indeed, improved timing and better patient selection are felt to have contributed more to the increased

success of liver transplantation in the 1980s and beyond than all the impressive technical and immunologic advances combined. Although the disease should be advanced, and although opportunities for spontaneous or medically induced stabilization or recovery should be allowed, the procedure should be done sufficiently early to give the surgical procedure a fair chance for success. Ideally, transplantation should be considered in patients with end-stage liver disease who are experiencing or have experienced a life-threatening complication of hepatic decompensation or whose quality of life has deteriorated to unacceptable levels. Although patients with well-compensated cirrhosis can survive for many years, many patients with quasi-stable chronic liver disease have much more advanced disease than may be apparent. As discussed below, the better the status of the patient prior to transplantation, the higher will be its anticipated success rate. The decision about *when* to transplant is complex and requires the combined judgment of an experienced team of hepatologists, transplant surgeons, anesthesiologists, and specialists in support services, not to mention the well-informed consent of the patient and the patient's family.

TRANSPLANTATION IN CHILDREN

Indications for transplantation in children are listed in [Table 368-1](#). The most common is *biliary atresia*. *Inherited or genetic disorders of metabolism* associated with liver failure constitute another major indication for transplantation in children and adolescents. In Crigler-Najjar disease type I and in certain hereditary disorders of the urea cycle and of amino acid or lactate-pyruvate metabolism, transplantation may be the only way to prevent impending deterioration of central nervous system function, despite the fact that the native liver is structurally normal. Combined heart and liver transplantation has yielded dramatic improvement in cardiac function and in cholesterol levels in children with homozygous familial hypercholesterolemia; combined liver and kidney transplantation has been successful in patients with primary hyperoxaluria type I. In hemophiliacs with transfusion-associated hepatitis and liver failure, liver transplantation has been associated with recovery of normal factor VIII synthesis.

TRANSPLANTATION IN ADULTS

Liver transplantation is indicated for end-stage *cirrhosis* of all causes ([Table 368-1](#)). In *sclerosing cholangitis* and *Caroli's disease* (multiple cystic dilatations of the intrahepatic biliary tree), recurrent infections and sepsis associated with inflammatory and fibrotic obstruction of the biliary tree may be an indication for transplantation. Because prior biliary surgery complicates and is a relative contraindication for liver transplantation, surgical diversion of the biliary tree has been all but abandoned for patients with sclerosing cholangitis. In patients who

undergo transplantation for *hepatic vein thrombosis* (*Budd-Chiari syndrome*), postoperative anticoagulation is essential; underlying myeloproliferative disorders may have to be treated but are not a contraindication to liver transplantation. If a donor organ can be located quickly, before life-threatening complications—including cerebral edema—set in, patients with acute liver failure are candidates for liver transplantation. Routine candidates for liver transplantation are patients with *alcoholic cirrhosis*, *chronic viral hepatitis*, and *primary hepatocellular malignancies*. Although all three of these categories are considered to be high risk, liver transplantation can be offered to carefully selected patients. Currently, chronic hepatitis C and alcoholic liver disease are the most common indications for liver transplantation, accounting for over 40% of all adult candidates who undergo the procedure. Patients with alcoholic cirrhosis can be considered as candidates for transplantation if they meet strict criteria for abstinence and reform; however, these criteria still do not prevent recidivism in up to a quarter of cases. In highly selected cases in a limited number of centers, transplantation for severe *acute* alcoholic hepatitis has been performed with success; however, because patients with acute alcoholic hepatitis are still actively using alcohol, and because continued alcohol abuse remains a concern, acute alcoholic hepatitis is not a routine indication for liver transplantation. Patients with chronic hepatitis C have early allograft and patient survival comparable to those of other subsets of patients after transplantation; however, reinfection in the donor organ is universal, recurrent hepatitis C is insidiously progressive, allograft cirrhosis develops in 20–30% at 5 years, and cirrhosis and late organ failure occur at a higher frequency beyond 5 years. With the introduction of highly effective direct acting antiviral agents targeting HCV, it is expected that allograft outcomes will improve significantly in the coming years. In patients with chronic hepatitis B, in the absence of measures to prevent recurrent hepatitis B, survival after transplantation is reduced by approximately 10–20%; however, prophylactic use of hepatitis B immune globulin (HBIG) during and after transplantation increases the success of transplantation to a level comparable to that seen in patients with nonviral causes of liver decompensation. Specific oral antiviral drugs (e.g., entecavir, tenofovir disoproxil fumarate) ([Chap. 362](#)) can be used both for prophylaxis against and for treatment of recurrent hepatitis B, facilitating further the management of patients undergoing liver transplantation for end-stage hepatitis B; most transplantation centers rely on antiviral drugs with or without HBIG to manage patients with hepatitis B. Issues of disease recurrence are discussed in more detail below. Patients with nonmetastatic primary hepatobiliary tumors—primary hepatocellular carcinoma (HCC), cholangiocarcinoma, hepatoblastoma, angiosarcoma, epithelioid hemangioendothelioma, and multiple or massive hepatic adenomata—have undergone liver transplantation; however, for some hepatobiliary malignancies, overall survival is significantly lower than that for other categories of liver disease. Most transplantation centers have reported 5-year recurrence-free survival rates in patients with unresectable HCC for single tumors <5 cm in diameter or for three or fewer lesions all <3 cm comparable to those seen in patients undergoing transplantation for nonmalignant indications. Consequently, liver transplantation is currently restricted to patients whose hepatic malignancies meet these criteria. Expanded criteria for patients with HCC continue to be evaluated. Because the likelihood of recurrent cholangiocarcinoma is very high, only highly selected patients with limited disease are being evaluated for transplantation after intensive chemotherapy and radiation.

CONTRAINDICATIONS

Absolute contraindications for transplantation include life-threatening systemic diseases, uncontrolled extrahepatic bacterial or fungal infections, preexisting advanced cardiovascular or pulmonary disease, multiple uncorrectable life-threatening congenital anomalies, metastatic malignancy, and active drug or alcohol abuse ([Table 368-2](#)). Because carefully selected patients in their sixties and even seventies have undergone transplantation successfully, advanced age per se is no longer considered an absolute contraindication; however, in older patients a more thorough preoperative evaluation should be undertaken to

TABLE 368-1 INDICATIONS FOR LIVER TRANSPLANTATION

Children	Adults
Biliary atresia	Primary biliary cirrhosis
Neonatal hepatitis	Secondary biliary cirrhosis
Congenital hepatic fibrosis	Primary sclerosing cholangitis
Alagille's syndrome ^a	Autoimmune hepatitis
Byler's disease ^b	Caroli's disease ^c
α_1 -Antitrypsin deficiency	Cryptogenic cirrhosis
Inherited disorders of metabolism	Chronic hepatitis with cirrhosis
Wilson's disease	Hepatic vein thrombosis
Tyrosinemia	Fulminant hepatitis
Glycogen storage diseases	Alcoholic cirrhosis
Lysosomal storage diseases	Chronic viral hepatitis
Protoporphyrria	Primary hepatocellular malignancies
Crigler-Najjar disease type I	Hepatic adenomas
Familial hypercholesterolemia	Nonalcoholic steatohepatitis
Primary hyperoxaluria type I	Familial amyloid polyneuropathy
Hemophilia	

^aArteriohepatic dysplasia, with paucity of bile ducts, and congenital malformations, including pulmonary stenosis. ^bIntrahepatic cholestasis, progressive liver failure, and mental and growth retardation. ^cMultiple cystic dilatations of the intrahepatic biliary tree.

TABLE 368-2 CONTRAINDICATIONS TO LIVER TRANSPLANTATION

Absolute	Relative
Uncontrolled extrahepatic infection	Age >70
Active, untreated sepsis	Prior extensive hepatobiliary surgery
Uncorrectable, life-limiting congenital anomalies	Portal vein thrombosis
Active substance or alcohol abuse	Renal failure not attributable to liver disease
Advanced cardiopulmonary disease	Previous extrahepatic malignancy (not including nonmelanoma skin cancer)
Extrahepatic malignancy (not including nonmelanoma malignancy skin cancer)	Severe obesity
Metastatic malignancy to the liver	Severe malnutrition/wasting
Cholangiocarcinoma	Medical noncompliance
AIDS	HIV seropositivity with failure to control HIV viremia or CD4 <100/ μ L
Life-threatening systemic diseases	Intrahepatic sepsis
	Severe hypoxemia secondary to right-to-left intrapulmonary shunts (P_{O_2} <50 mmHg)
	Severe pulmonary hypertension (mean pulmonary artery pressure >35 mmHg)
	Uncontrolled psychiatric disorder

exclude ischemic cardiac disease and other comorbid conditions. Advanced age (>70 years), however, should be considered a *relative contraindication*—that is, a factor to be taken into account with other relative contraindications. Other relative contraindications include portal vein thrombosis, HIV infection, preexisting renal disease not associated with liver disease (which may prompt consideration of combined liver and kidney transplantation), intrahepatic or biliary sepsis, severe hypoxemia (P_{O_2} <50 mmHg) resulting from right-to-left intrapulmonary shunts, portopulmonary hypertension with high mean pulmonary artery pressures (>35 mmHg), previous extensive hepatobiliary surgery, any uncontrolled serious psychiatric disorder, and lack of sufficient social supports. Any one of these relative contraindications is insufficient in and of itself to preclude transplantation. For example, the problem of portal vein thrombosis can be overcome by constructing a graft from the donor liver portal vein to the recipient's superior mesenteric vein. Now that highly active antiretroviral therapy has dramatically improved the survival of persons with HIV infection (Chap. 226), and because end-stage liver disease caused by chronic hepatitis C and B has emerged as a serious source of morbidity and mortality in the HIV-infected population, liver transplantation has now been performed successfully in selected HIV-positive persons who have excellent control of HIV infection. Selected patients with CD4 \pm T cell counts >100/ μ L and with pharmacologic suppression of HIV viremia have undergone transplantation for end-stage liver disease. HIV-infected persons who have received liver allografts for end-stage liver disease resulting from chronic hepatitis B have experienced survival rates compared to those of HIV-negative persons undergoing transplantation for the same indication. In contrast, recurrent hepatitis C virus (HCV) in the allograft has limited long-term success in persons with HCV-related end-stage liver disease. Again, it is expected that the availability of direct acting antiviral agents targeting HCV, will significantly improve allograft outcomes.

TECHNICAL CONSIDERATIONS

CADAVER DONOR SELECTION

Cadaver donor livers for transplantation are procured primarily from victims of head trauma. Organs from brain-dead donors up to age 60 are acceptable if the following criteria are met: hemodynamic stability,

adequate oxygenation, absence of bacterial or fungal infection, absence of abdominal trauma, absence of hepatic dysfunction, and serologic exclusion of hepatitis B (HBV) and C viruses and HIV. Occasionally, organs from donors with hepatitis B and C are used (e.g., for recipients with prior hepatitis B and C, respectively). Organs from donors with antibodies to hepatitis B core antigen (anti-HBc) can also be used when the need is especially urgent, and recipients of these organs are treated prophylactically with antiviral drugs. Cardiovascular and respiratory functions are maintained artificially until the liver can be removed. Transplantation of organs procured from deceased donors who have succumbed to cardiac death can be performed successfully under selected circumstances, when ischemic time is minimized and liver histology preserved. Compatibility in ABO blood group and organ size between donor and recipient are important considerations in donor selection; however, ABO-incompatible, split liver, or reduced-donor-organ transplants can be performed in emergencies or marked donor scarcity. Tissue typing for human leukocyte antigen (HLA) matching is not required, and preformed cytotoxic HLA antibodies do not preclude liver transplantation. Following perfusion with cold electrolyte solution, the donor liver is removed and packed in ice. The use of University of Wisconsin (UW) solution, rich in lactobionate and raffinose, has permitted the extension of cold ischemic time up to 20 h; however, 12 h may be a more reasonable limit. Improved techniques for harvesting multiple organs from the same donor have increased the availability of donor livers, but the availability of donor livers is far outstripped by the demand. Currently in the United States, all donor livers are distributed through a nationwide organ-sharing network (United Network for Organ Sharing [UNOS]) designed to allocate available organs based on regional considerations and recipient acuity. Recipients who have the highest disease severity generally have the highest priority, but allocation strategies that balance highest urgency against best outcomes continue to evolve to distribute cadaver organs most effectively. Allocation based on the Child-Turcotte-Pugh (CTP) score, which uses five clinical variables (encephalopathy stage, ascites, bilirubin, albumin, and prothrombin time) and waiting time, has been replaced by allocation based on urgency alone, calculated by the Model for End-Stage Liver Disease (MELD) score. The MELD score is based on a mathematical model that includes bilirubin, creatinine, and prothrombin time expressed as international normalized ratio (INR) (Table 368-3). Neither waiting time (except as a tie breaker between two potential recipients with the same MELD scores) nor posttransplantation outcome is taken into account, but use of the MELD score

TABLE 368-3 UNITED NETWORK FOR ORGAN SHARING (UNOS) LIVER TRANSPLANTATION WAITING LIST CRITERIA

Status 1	Fulminant hepatic failure (including primary graft nonfunction and hepatic artery thrombosis within 7 days after transplantation as well as acute decompensated Wilson's disease) ^a
The Model for End-Stage Liver Disease (MELD) score, on a continuous scale, ^b determines allocation of the remainder of donor organs. This model is based on the following calculation:	
$3.78 \times \log_e \text{bilirubin (mg/100 mL)} \pm 11.2 \times \log_e \text{international normalized ratio (INR)} \pm 9.57 \times \log_e \text{creatinine (mg/100 mL)} \pm 6.43$ (v 0 for alcoholic and cholestatic liver disease, $\times 1$ for all other types of liver disease). ^{c,d,e}	
Online calculators to determine MELD scores are available, such as the following: http://optn.transplant.hrsa.gov/resources/professionalresources.asp?index=9 .	

^aFor children <18 years of age, status 1 includes acute or chronic liver failure plus hospitalization in an intensive care unit or inborn errors of metabolism. Status 1 is retained for those persons with fulminant hepatic failure and supersedes the MELD score. ^bThe MELD scale is continuous, with 34 levels ranging between 6 and 40. Donor organs usually do not become available unless the MELD score exceeds 20. ^cPatients with stage T2 hepatocellular carcinoma receive 22 disease-specific points. ^dCreatinine is included because renal function is a validated predictor of survival in patients with liver disease. For adults undergoing dialysis twice a week, the creatinine in the equation is set to 4 mg/100 mL. ^eFor children <18 years of age, the Pediatric End-Stage Liver Disease (PELD) scale is used. This scale is based on albumin, bilirubin, INR, growth failure, and age. Status 1 is retained.

2070 has been shown to reduce waiting list mortality, to reduce waiting time prior to transplantation, to be the best predictor of pretransplantation mortality, to satisfy the prevailing view that medical need should be the decisive determinant, and to eliminate both the subjectivity inherent in the CTP scoring system (presence and degree of ascites and hepatic encephalopathy) and the differences in waiting times among different regions of the country. Recent data indicate that liver recipients with MELD scores <15 experienced higher posttransplantation mortality rates than similarly classified patients who remained on the wait list. This observation led to the modification of UNOS policy to allocate donor organs to candidates with MELD scores exceeding 15 within the local or regional procurement organization before offering the organ to local patients whose scores are <15. In addition, serum sodium, another important predictor of survival in liver transplantation candidates, is taken into consideration in allocating donor livers.

The highest priority (status 1) continues to be reserved for patients with fulminant hepatic failure or primary graft nonfunction. Because candidates for liver transplantation who have HCC may not be sufficiently decompensated to compete for donor organs based on urgency criteria alone, and because protracted waiting for cadaver donor organs often results in tumor growth beyond acceptable limits for transplantation, such patients are assigned disease-specific MELD points (Table 368-3). Other disease-specific MELD exceptions include portopulmonary hypertension, hepatopulmonary syndrome, familial amyloid polyneuropathy, primary hyperoxaluria (necessitating liver-kidney transplantation), cystic fibrosis liver disease, and highly selected cases of hilar cholangiocarcinoma.

LIVING DONOR TRANSPLANTATION

Occasionally, especially for liver transplantation in children, one cadaver organ can be split between two recipients (one adult and one child). A more viable alternative, transplantation of the right lobe of the liver from a healthy adult donor into an adult recipient, has gained increased popularity. Living donor transplantation of the left lobe (left lateral segment), introduced in the early 1990s to alleviate the extreme shortage of donor organs for small children, accounts currently for approximately one-third of all liver transplantation procedures in children. Driven by the shortage of cadaver organs, living donor transplantation involving the more sizable right lobe is being considered with increasing frequency in adults; however, living donor liver transplantation cannot be expected to solve the donor organ shortage; 246 such procedures were done in 2012, representing only about 4% of all liver transplant operations done in the United States.

Living donor transplantation can reduce waiting time and cold-ischemia time; is done under elective, rather than emergency, circumstances; and may be lifesaving in recipients who cannot afford to wait for a cadaver donor. The downside, of course, is the risk to the healthy donor (a mean of 10 weeks of medical disability; biliary complications in ~5%; postoperative complications such as wound infection, small-bowel obstruction, and incisional hernias in 9-19%; and even, in 0.2-0.4%, death) as well as the increased frequency of biliary (15-32%) and vascular (10%) complications in the recipient. Potential donors must participate voluntarily without coercion, and transplantation teams should go to great lengths to exclude subtle coercive or inappropriate psychological factors as well as outline carefully to both donor and recipient the potential benefits and risks of the procedure. Donors for the procedure should be 18-60 years old; have a compatible blood type with the recipient; have no chronic medical problems or history of major abdominal surgery; be related genetically or emotionally to the recipient; and pass an exhaustive series of clinical, biochemical, and serologic evaluations to unearth disqualifying medical disorders. The recipient should meet the same UNOS criteria for liver transplantation as recipients of a cadaver donor allograft. Comprehensive outcome data on adult-to-adult living donor liver transplantation are being collected (www.nih-a2all.org).

SURGICAL TECHNIQUE

Removal of the recipient's native liver is technically difficult, particularly in the presence of portal hypertension with its associated

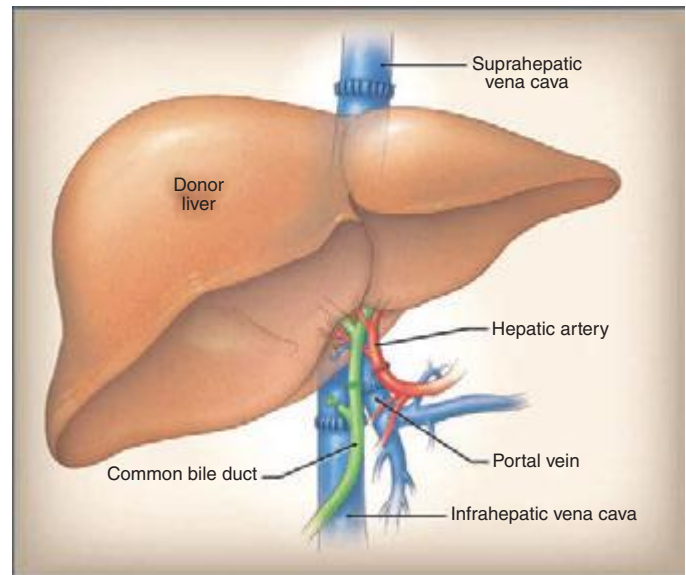


FIGURE 368-1 The anastomoses in orthotopic liver transplantation. The anastomoses are performed in the following sequence: (1) suprahepatic and infrahepatic vena cava, (2) portal vein, (3) hepatic artery, and (4) common bile duct-to-duct anastomosis. (Adapted from JL Dienstag, AB Cosimi: *N Engl J Med* 367:1483, 2012.)

collateral circulation and extensive varices and especially in the presence of scarring from previous abdominal operations. The combination of portal hypertension and coagulopathy (elevated prothrombin time and thrombocytopenia) may translate into large blood product transfusion requirements. After the portal vein and infrahepatic and suprahepatic inferior vena cavae are dissected, the hepatic artery and common bile duct are dissected. Then the native liver is removed and the donor organ inserted. During the anhepatic phase, coagulopathy, hypoglycemia, hypocalcemia, and hypothermia are encountered and must be managed by the anesthesiology team. Caval, portal vein, hepatic artery, and bile duct anastomoses are performed in succession, the last by end-to-end suturing of the donor and recipient common bile ducts (Fig. 368-1) or by choledochojejunostomy to a Roux-en-Y loop if the recipient common bile duct cannot be used for reconstruction (e.g., in sclerosing cholangitis). A typical transplant operation lasts 8 h, with a range of 6-18 h. Because of excessive bleeding, large volumes of blood, blood products, and volume expanders may be required during surgery; however, blood requirements have fallen sharply with improvements in surgical technique, blood-salvage interventions, and experience.

As noted above, emerging alternatives to orthotopic liver transplantation include split-liver grafts, in which one donor organ is divided and inserted into two recipients; and living donor procedures, in which part of the left (for children), the left (for children or small adults), or the right (for adults) lobe of the liver is harvested from a living donor for transplantation into the recipient. In the adult procedure, once the right lobe is removed from the donor, the donor right hepatic vein is anastomosed to the recipient right hepatic vein remnant, followed by donor-to-recipient anastomoses of the portal vein and then the hepatic artery. Finally, the biliary anastomosis is performed, duct-to-duct if practical or via Roux-en-Y anastomosis. Heterotopic liver transplantation, in which the donor liver is inserted without removal of the native liver, has met with very limited success and acceptance, except in a very small number of centers. In attempts to support desperately ill patients until a suitable donor organ can be identified, several transplantation centers are studying extracorporeal perfusion with bioartificial liver cartridges constructed from hepatocytes bound to hollow fiber systems and used as temporary hepatic-assist devices, but their efficacy remains to be established. Areas of research with the potential to overcome the shortage of donor organs include hepatocyte transplantation and

xenotransplantation with genetically modified organs of nonhuman origin (e.g., swine).

POSTOPERATIVE COURSE AND MANAGEMENT

IMMUNOSUPPRESSIVE THERAPY

The introduction in 1980 of cyclosporine as an immunosuppressive agent contributed substantially to the improvement in survival after liver transplantation. Cyclosporine, a calcineurin inhibitor, blocks early activation of T cells and is specific for T cell functions that result from the interaction of the T cell with its receptor and that involve the calcium-dependent signal transduction pathway. As a result, the activity of cyclosporine leads to inhibition of lymphokine gene activation, blocking interleukins 2, 3, and 4, tumor necrosis factor α , and other lymphokines. Cyclosporine also inhibits B cell functions. This process occurs without affecting rapidly dividing cells in the bone marrow, which may account for the reduced frequency of posttransplantation systemic infections. The most common and important side effect of cyclosporine therapy is nephrotoxicity. Cyclosporine causes dose-dependent renal tubular injury and direct renal artery vasospasm. Following renal function is therefore important in monitoring cyclosporine therapy, perhaps even a more reliable indicator than blood levels of the drug. Nephrotoxicity is reversible and can be managed by dose reduction. Other adverse effects of cyclosporine therapy include hypertension, hyperkalemia, tremor, hirsutism, glucose intolerance, and gingival hyperplasia.

Tacrolimus, a macrolide lactone antibiotic isolated from a Japanese soil fungus, *Streptomyces tsukubaensis*, has the same mechanism of action as cyclosporine but is 10–100 times more potent. Initially applied as “rescue” therapy for patients in whom rejection occurred despite the use of cyclosporine, tacrolimus was shown to be associated with a reduced frequency of acute, refractory, and chronic rejection. Although patient and graft survival are the same with these two drugs, the advantage of tacrolimus in minimizing episodes of rejection, reducing the need for additional glucocorticoid doses, and reducing the likelihood of bacterial and cytomegalovirus (CMV) infection has simplified the management of patients undergoing liver transplantation. In addition, the oral absorption of tacrolimus is more predictable than that of cyclosporine, especially during the early postoperative period when T-tube drainage interferes with the enterohepatic circulation of cyclosporine. As a result, in most transplantation centers, tacrolimus has now supplanted cyclosporine for primary immunosuppression, and many centers rely on oral rather than IV administration from the outset. For transplantation centers that prefer cyclosporine, a better-absorbed microemulsion preparation is available.

Although more potent than cyclosporine, tacrolimus is also more toxic and more likely to be discontinued for adverse events. The toxicity of tacrolimus is similar to that of cyclosporine; nephrotoxicity and neurotoxicity are the most commonly encountered adverse effects, and neurotoxicity (tremor, seizures, hallucinations, psychoses, coma) is more likely and more severe in tacrolimus-treated patients. Both drugs can cause diabetes mellitus, but tacrolimus does not cause hirsutism or gingival hyperplasia. Because of overlapping toxicity between cyclosporine and tacrolimus, especially nephrotoxicity, and because tacrolimus reduces cyclosporine clearance, these two drugs should not be used together. Because 99% of tacrolimus is metabolized by the liver, hepatic dysfunction reduces its clearance; in primary graft nonfunction (when, for technical reasons or because of ischemic damage prior to its insertion, the allograft is defective and does not function normally from the outset), tacrolimus doses have to be reduced substantially, especially in children. Both cyclosporine and tacrolimus are metabolized by the cytochrome P450 IIIA system, and, therefore, drugs that induce cytochrome P450 (e.g., phenytoin, phenobarbital, carbamazepine, rifampin) reduce available levels of cyclosporine and tacrolimus; and drugs that inhibit cytochrome P450 (e.g., erythromycin, fluconazole, ketoconazole, clotrimazole, itraconazole, verapamil, diltiazem, danazol, metoclopramide, the HIV protease inhibitor ritonavir, and the HCV protease inhibitors telaprevir and boceprevir) increase cyclosporine and tacrolimus blood levels. Indeed, itraconazole is used occasionally to help boost tacrolimus levels. Like

azathioprine, cyclosporine and tacrolimus appear to be associated with a risk of lymphoproliferative malignancies (see below), which may occur earlier after cyclosporine or tacrolimus than after azathioprine therapy. Because of these side effects, combinations of cyclosporine or tacrolimus with prednisone and an antimetabolite (azathioprine or mycophenolic acid, see below)—all at reduced doses—are preferable regimens for immunosuppressive therapy.

Mycophenolic acid, a nonnucleoside purine metabolism inhibitor derived as a fermentation product from several *Penicillium* species, is another immunosuppressive drug being used for patients undergoing liver transplantation. Mycophenolate has been shown to be better than azathioprine, when used with other standard immunosuppressive drugs, in preventing rejection after renal transplantation and has been adopted widely as well for use in liver transplantation. The most common adverse effects of mycophenolate are bone marrow suppression and gastrointestinal complaints.

In patients with pretransplantation renal dysfunction or renal deterioration that occurs intraoperatively or immediately postoperatively, tacrolimus or cyclosporine therapy may not be practical; under these circumstances, induction or maintenance of immunosuppression with antithymocyte globulin (ATG, thymoglobulin) or monoclonal antibodies to T cells, OKT3, may be appropriate. Therapy with these agents has been especially effective in reversing acute rejection in the posttransplantation period and is the standard treatment for acute rejection that fails to respond to methylprednisolone boluses. Available data support the use of thymoglobulin induction to delay calcineurin inhibitor use and its attendant nephrotoxicity. IV infusions of thymoglobulin may be complicated by fever and chills, which can be ameliorated by premedication with antipyretics and a low dose of glucocorticoids. Infusions of OKT3 may be complicated by fever, chills, and diarrhea, or by pulmonary edema, which can be fatal. Because OKT3 is such a potent immunosuppressive agent, its use is also more likely to be complicated by opportunistic infection or lymphoproliferative disorders; therefore, because of the availability of alternative immunosuppressive drugs, OKT3 is now used sparingly.

Sirolimus, an inhibitor of the mammalian target of rapamycin (mTOR), blocks later events in T cell activation, is approved for use in kidney transplantation, but is not approved for use in liver transplant recipients because of the reported association with an increased frequency of hepatic artery thrombosis in the first month posttransplantation. In patients with calcineurin inhibitor-related nephrotoxicity, conversion to sirolimus has been demonstrated to be effective in preventing rejection with accompanying improvements in renal function. Because of its profound antiproliferative effects, sirolimus has also been suggested to be a useful immunosuppressive agent in patients with a prior or current history of malignancy, such as HCC. Side effects include hyperlipidemia, peripheral edema, oral ulcers, and interstitial pneumonitis. Everolimus is a hydroxyethyl derivative of sirolimus that, when used in conjunction with low-dose tacrolimus, also provides successful protection against acute rejection, with decreased renal impairment compared to that associated with standard tacrolimus dosing. Everolimus and sirolimus share a similar adverse events profile; therefore, neither of these agents is approved for routine use in liver allograft recipients.

The most important principle of immunosuppression is that the ideal approach strikes a balance between immunosuppression and immunologic competence. In general, given sufficient immunosuppression, acute liver allograft rejection is nearly always reversible. On one hand, incompletely treated acute rejection predisposes to the development of chronic rejection, which can threaten graft survival. On the other hand, if the cumulative dose of immunosuppressive therapy is too large, the patient may succumb to opportunistic infection. In hepatitis C, pulse glucocorticoids or OKT3 use accelerate recurrent allograft hepatitis. Further complicating matters, acute rejection can be difficult to distinguish histologically from recurrent hepatitis C. Therefore, immunosuppressive drugs must be used judiciously, with strict attention to the infectious consequences of such therapy and careful confirmation of the diagnosis of acute rejection. In this vein, efforts have been made to minimize the use of glucocorticoids, a

2072 mainstay of immunosuppressive regimens, and steroid-free immunosuppression can be achieved in some instances. Patients who undergo liver transplantation for autoimmune diseases such as primary biliary cirrhosis, autoimmune hepatitis, and primary sclerosing cholangitis are less likely to achieve freedom from glucocorticoids.

POSTOPERATIVE COMPLICATIONS

Complications of liver transplantation can be divided into nonhepatic and hepatic categories (Tables 368-4 and 368-5). In addition, both immediate postoperative and late complications are encountered. As a rule, patients who undergo liver transplantation have been chronically ill for protracted periods and may be malnourished and wasted. The impact of such chronic illness and the multisystem failure that accompanies liver failure continue to require attention in the postoperative period. Because of the massive fluid losses and fluid shifts that occur during the operation, patients may remain fluid-overloaded during the immediate postoperative period, straining cardiovascular reserve; this effect can be amplified in the face of transient renal dysfunction and pulmonary capillary vascular permeability. Continuous monitoring of cardiovascular and pulmonary function, measures to maintain the integrity of the intravascular compartment and to treat extravascular volume overload, and scrupulous attention to potential sources and sites of infection are of paramount importance. Cardiovascular instability may also result from the electrolyte imbalance that may accompany reperfusion of the donor liver as well as from restoration of systemic vascular resistance following implantation. Pulmonary function may be compromised further by paralysis of the right hemidiaphragm associated with phrenic nerve injury. The hyperdynamic state with increased cardiac output that is characteristic of patients with liver failure reverses rapidly after successful liver transplantation.

Other immediate management issues include renal dysfunction. Prerenal azotemia, acute kidney injury associated with hypoperfusion (acute tubular necrosis), and renal toxicity caused by antibiotics,

TABLE 368-5 HEPATIC COMPLICATIONS OF LIVER TRANSPLANTATION

Hepatic Dysfunction Common After Major Surgery	
Prehepatic	Pigment load Hemolysis Blood collections (hematomas, abdominal collections)
Intrahepatic	
Early	Hepatotoxic drugs and anesthesia Hypoperfusion (hypotension, shock, sepsis) Benign postoperative cholestasis
Late	Transfusion-associated hepatitis Exacerbation of primary hepatic disease
Posthepatic	Biliary obstruction ↓ Renal clearance of conjugated bilirubin (renal dysfunction)
Hepatic Dysfunction Unique to Liver Transplantation	
Primary graft nonfunction	
Vascular compromise	Portal vein obstruction Hepatic artery thrombosis Anastomotic leak with intraabdominal bleeding
Bile duct disorder	Stenosis, obstruction, leak
Rejection	
Recurrent primary hepatic disease	

tacrolimus, or cyclosporine are encountered frequently in the postoperative period, sometimes necessitating dialysis. Hemolytic-uremic syndrome can be associated with cyclosporine, tacrolimus, or OKT3. Occasionally, postoperative intraperitoneal bleeding may be sufficient to increase intraabdominal pressure, which, in turn, may reduce renal blood flow; this effect is rapidly reversible when abdominal distention is relieved by exploratory laparotomy to identify and ligate the bleeding site and to remove intraperitoneal clot.

Anemia may also result from acute upper gastrointestinal bleeding or from transient hemolytic anemia, which may be autoimmune, especially when blood group O livers are transplanted into blood group A or B recipients. This autoimmune hemolytic anemia is mediated by donor intrahepatic lymphocytes that recognize red blood cell A or B antigens on recipient erythrocytes. Transient in nature, this process resolves once the donor liver is repopulated by recipient bone marrow-derived lymphocytes; the hemolysis can be treated by transfusing blood group O red blood cells and/or by administering higher doses of glucocorticoids. Transient thrombocytopenia is also commonly encountered. Aplastic anemia, a late occurrence, is rare but has been reported in almost 30% of patients who underwent liver transplantation for acute, severe hepatitis of unknown cause.

Bacterial, fungal, or viral infections are common and may be life-threatening postoperatively. Early after transplant surgery, common postoperative infections predominate—pneumonia, wound infections, infected intraabdominal collections, urinary tract infections, and IV line infections—rather than opportunistic infections; these infections may involve the biliary tree and liver as well. Beyond the first postoperative month, the toll of immunosuppression becomes evident, and opportunistic infections—CMV, herpes viruses, fungal infections (*Aspergillus*, *Candida*, cryptococcal disease), mycobacterial infections, parasitic infections (*Pneumocystis*, *Toxoplasma*), bacterial infections (*Nocardia*, *Legionella*, *Listeria*)—predominate. Rarely, early infections represent those transmitted with the donor liver, either infections present in the donor or infections acquired during procurement processing. De novo viral hepatitis infections acquired from the donor organ or, almost unheard of now, from transfused blood products occur after typical incubation periods for these agents (well beyond the first month). Obviously, infections in an immunosuppressed host demand early recognition and prompt management; prophylactic antibiotic therapy is administered routinely in the immediate postoperative period. Use

TABLE 368-4 NONHEPATIC COMPLICATIONS OF LIVER TRANSPLANTATION

Fluid overload	
Cardiovascular instability	Arrhythmias Congestive heart failure Cardiomyopathy
Pulmonary compromise	Pneumonia Pulmonary capillary vascular permeability Fluid overload
Renal dysfunction	Prerenal azotemia Hypoperfusion injury (acute tubular necrosis) Drug nephrotoxicity ↓ Renal blood flow secondary to ↑ intraabdominal pressure
Hematologic	Anemia secondary to gastrointestinal and/or intraabdominal bleeding Hemolytic anemia, aplastic anemia Thrombocytopenia
Infection	Bacterial: early, common postoperative infections Fungal/parasitic: late, opportunistic infections Viral: late, opportunistic infections, recurrent hepatitis
Neuropsychiatric	Seizures Metabolic encephalopathy Depression Difficult psychosocial adjustment
Diseases of donor	Infectious Malignant
Malignancy	B cell lymphoma (posttransplantation lymphoproliferative disorders) De novo neoplasms (particularly squamous cell skin carcinoma)

of sulfamethoxazole with trimethoprim reduces the incidence of postoperative *Pneumocystis carinii* pneumonia. Antiviral prophylaxis for CMV with ganciclovir should be administered in patients at high risk (e.g., when a CMV-seropositive donor organ is implanted into a CMV-seronegative recipient).

Neuropsychiatric complications include seizures (commonly associated with cyclosporine and tacrolimus toxicity), metabolic encephalopathy, depression, and difficult psychosocial adjustment. Rarely, diseases are transmitted by the allograft from the donor to the recipient. In addition to viral and bacterial infections, malignancies of donor origin have occurred. Posttransplantation lymphoproliferative disorders, especially B cell lymphoma, are a recognized complication associated with immunosuppressive drugs such as azathioprine, tacrolimus, and cyclosporine (see above). Epstein-Barr virus has been shown to play a contributory role in some of these tumors, which may regress when immunosuppressive therapy is reduced. De novo neoplasms appear at increased frequency after liver transplantation, particularly squamous cell carcinomas of the skin. Routine screening should be performed.

Long-term complications after liver transplantation attributable primarily to immunosuppressive medications include diabetes mellitus and osteoporosis (associated with glucocorticoids and calcineurin inhibitors) as well as hypertension, hyperlipidemia, and chronic renal insufficiency (associated with cyclosporine and tacrolimus). Monitoring and treating these disorders are routine components of posttransplantation care; in some cases, they respond to changes in immunosuppressive regimen, while in others, specific treatment of the disorder is introduced. Data from a large U.S. database showed that the prevalence of renal failure was 18% at year 5 and 25% at year 10 after liver transplantation. Similarly, the high frequency of diabetes, hypertension, hyperlipidemia, obesity, and the metabolic syndrome renders patients susceptible to cardiovascular disease after liver transplantation; although hepatic complications account for most of the mortality after liver transplantation, renal failure and cardiovascular disease are the other leading causes of late mortality after liver transplantation.

HEPATIC COMPLICATIONS

Hepatic dysfunction after liver transplantation is similar to the hepatic complications encountered after major abdominal and cardiothoracic surgery; however, in addition, hepatic complications include primary graft failure, vascular compromise, failure or stricture of the biliary anastomoses, and rejection. As in nontransplantation surgery, postoperative jaundice may result from prehepatic, intrahepatic, and posthepatic sources. *Prehepatic* sources represent the massive hemoglobin pigment load from transfusions, hemolysis, hematomas, ecchymoses, and other collections of blood. *Early intrahepatic* liver injury includes effects of hepatotoxic drugs and anesthesia; hypoperfusion injury associated with hypotension, sepsis, and shock; and benign postoperative cholestasis. *Late intrahepatic* sources of liver injury include exacerbation of primary disease. *Posthepatic* sources of hepatic dysfunction include biliary obstruction and reduced renal clearance of conjugated bilirubin. Hepatic complications unique to liver transplantation include primary graft failure associated with ischemic injury to the organ during harvesting; vascular compromise associated with thrombosis or stenosis of the portal vein or hepatic artery anastomoses; vascular anastomotic leak; stenosis, obstruction, or leakage of the anastomosed common bile duct; recurrence of primary hepatic disorder (see below); and rejection.

TRANSPLANT REJECTION

Despite the use of immunosuppressive drugs, rejection of the transplanted liver still occurs in a proportion of patients, beginning 1–2 weeks after surgery. Clinical signs suggesting rejection are fever, right upper quadrant pain, and reduced bile pigment and volume. Leukocytosis may occur, but the most reliable indicators are increases in serum bilirubin and aminotransferase levels. Because these tests lack specificity, distinguishing among rejection, biliary obstruction, primary graft nonfunction, vascular compromise, viral hepatitis, CMV infection, drug hepatotoxicity, and recurrent primary disease may be difficult. Radiographic visualization of the biliary tree and/or

percutaneous liver biopsy often help to establish the correct diagnosis. Morphologic features of acute rejection include a mixed portal cellular infiltrate, bile duct injury, and/or endothelial inflammation (“endothelialitis”); some of these findings are reminiscent of graft-versus-host disease, primary biliary cirrhosis, or recurrent allograft hepatitis C. As soon as transplant rejection is suspected, treatment consists of IV methylprednisolone in repeated boluses; if this fails to abort rejection, many centers use thymoglobulin or OKT3. Caution should be exercised when managing acute rejection with pulse glucocorticoids or OKT3 in patients with HCV infection, because of the high risk of triggering recurrent allograft hepatitis C.

Chronic rejection is a relatively rare outcome that can follow repeated bouts of acute rejection or that occurs unrelated to preceding rejection episodes. Morphologically, chronic rejection is characterized by progressive cholestasis, focal parenchymal necrosis, mononuclear infiltration, vascular lesions (intimal fibrosis, subintimal foam cells, fibrinoid necrosis), and fibrosis. This process may be reflected as ductopenia—the vanishing bile duct syndrome, which is more common in patients undergoing liver transplantation for autoimmune liver disease. Reversibility of chronic rejection is limited; in patients with therapy-resistant chronic rejection, retransplantation has yielded encouraging results.

OUTCOME

SURVIVAL

The survival rate for patients undergoing liver transplantation has improved steadily since 1983. One-year survival rates have increased from ~70% in the early 1980s to 85–90% from 2003 to the present time. Currently, the 5-year survival rate exceeds 60%. An important observation is the relationship between clinical status before transplantation and outcome. For patients who undergo liver transplantation when their level of compensation is high (e.g., still working or only partially disabled), a 1-year survival rate of >85% is common. For those whose level of decompensation mandates continuous in-hospital care prior to transplantation, the 1-year survival rate is ~70%, whereas for those who are so decompensated that they require life support in an intensive care unit, the 1-year survival rate is ~50%. Since the adoption by UNOS in 2002 of the MELD system for organ allocation, posttransplantation survival has been found to be affected adversely for candidates with MELD scores >25, considered high disease severity. Thus, irrespective of allocation scheme, high disease severity before transplantation corresponds to diminished posttransplantation survival. Another important distinction in survival has been drawn between high- and low-risk patient categories. For patients who do not fit any “high-risk” designations, 1-year and 5-year survival rates of 85 and 80%, respectively, have been recorded. In contrast, among patients in high-risk categories—cancer, fulminant hepatitis, age >65, concurrent renal failure, respirator dependence, portal vein thrombosis, and history of a portacaval shunt or multiple right upper quadrant operations—survival statistics fall into the range of 60% at 1 year and 35% at 5 years. Survival after retransplantation for primary graft nonfunction is ~50%. Causes of failure of liver transplantation vary with time. Failures within the first 3 months result primarily from technical complications, postoperative infections, and hemorrhage. Transplant failures after the first 3 months are more likely to result from infection, rejection, or recurrent disease (such as malignancy or viral hepatitis).

RECURRENCE OF PRIMARY DISEASE

Features of autoimmune hepatitis, primary sclerosing cholangitis, and primary biliary cirrhosis overlap with those of rejection or posttransplantation bile duct injury. Whether autoimmune hepatitis and sclerosing cholangitis recur after liver transplantation is controversial; data supporting recurrent autoimmune hepatitis (in up to one-third of patients in some series) are more convincing than those supporting recurrent sclerosing cholangitis. Similarly, reports of recurrent primary biliary cirrhosis after liver transplantation have appeared; however, the histologic features of primary biliary cirrhosis and chronic rejection are virtually indistinguishable and occur as frequently in patients with

2074 primary biliary cirrhosis as in patients undergoing transplantation for other reasons. The presence of a florid inflammatory bile duct lesion is highly suggestive of the recurrence of primary biliary cirrhosis, but even this lesion can be observed in acute rejection. Hereditary disorders such as Wilson's disease and α_1 -antitrypsin deficiency have not recurred after liver transplantation; however, recurrence of disordered iron metabolism has been observed in some patients with hemochromatosis. Hepatic vein thrombosis (Budd-Chiari syndrome) may recur; this can be minimized by treating underlying myeloproliferative disorders and by anticoagulation. Because cholangiocarcinoma recurs almost invariably, few centers now offer transplantation to such patients; however, a few highly selected patients with operatively confirmed stage I or II cholangiocarcinoma who undergo liver transplantation combined with neoadjuvant chemoradiation may experience excellent outcomes. In patients with intrahepatic HCC who meet criteria for transplantation, 1- and 5-year survivals are similar to those observed in patients undergoing liver transplantation for non-malignant disease. Finally, metabolic disorders such as nonalcoholic steatohepatitis recur frequently, especially if the underlying metabolic predisposition is not altered. The metabolic syndrome occurs commonly after liver transplantation as a result of recurrent nonalcoholic fatty liver, immunosuppressive medications, and/or, in patients with hepatitis C related to the impact of HCV infection on insulin resistance, diabetes and fatty liver.

Hepatitis A can recur after transplantation for fulminant hepatitis A, but such acute reinfection has no serious clinical sequelae. In fulminant hepatitis B, recurrence is not the rule; however, in the absence of any prophylactic measures, hepatitis B usually recurs after transplantation for end-stage chronic hepatitis B. Before the introduction of prophylactic antiviral therapy, immunosuppressive therapy sufficient to prevent allograft rejection led inevitably to marked increases in hepatitis B viremia, regardless of pretransplantation levels. Overall graft and patient survival were poor, and some patients experienced a rapid recapitulation of severe injury—severe chronic hepatitis or even fulminant hepatitis—after transplantation. Also recognized in the era before availability of antiviral regimens was *fibrosing cholestatic hepatitis*, rapidly progressive liver injury associated with marked hyperbilirubinemia, substantial prolongation of the prothrombin time (both out of proportion to relatively modest elevations of aminotransferase activity), and rapidly progressive liver failure. This lesion has been suggested to represent a “choking off” of the hepatocyte by an overwhelming density of HBV proteins. Complications such as sepsis and pancreatitis were also observed more frequently in patients undergoing liver transplantation for hepatitis B prior to the introduction of antiviral therapy. The introduction of long-term prophylaxis with HBIg revolutionized liver transplantation for chronic hepatitis B. Preoperative hepatitis B vaccination, preoperative or postoperative interferon (IFN) therapy, or short-term (≤ 2 months) HBIg prophylaxis has not been shown to be effective, but a retrospective analysis of data from several hundred European patients followed for 3 years after transplantation has shown that long-term (≥ 6 months) prophylaxis with HBIg is associated with a lowering of the risk of HBV reinfection from ~ 75 to 35% and a reduction in mortality from ~ 50 to 20%.

As a result of long-term HBIg use following liver transplantation for chronic hepatitis B, similar improvements in outcome have been observed in the United States, with 1-year survival rates between 75% and 90%. Currently, with HBIg prophylaxis, the outcome of liver transplantation for chronic hepatitis B is indistinguishable from that for chronic liver disease unassociated with chronic hepatitis B; essentially, medical concerns regarding liver transplantation for chronic hepatitis B have been eliminated. Passive immunoprophylaxis with HBIg is begun during the anhepatic stage of surgery, repeated daily for the first 6 postoperative days, and then continued with infusions that are given either at regular intervals of 4–6 weeks or, alternatively, when anti-hepatitis B surface (HBs) levels fall below a threshold of 100 mIU/mL. The current approach in most centers is to continue HBIg indefinitely, which can add approximately \$20,000 per year to the cost of care; some centers are evaluating regimens that shift to less frequent administration or to IM administration in the late posttransplantation

period or, in low-risk patients, maintenance with antiviral therapy (see below) alone. Still, “breakthrough” HBV infection occasionally occurs.

Further improving the outcome of liver transplantation for chronic hepatitis B is the current availability of such antiviral drugs as lamivudine, adefovir, entecavir, and tenofovir disoproxil fumarate (**Chap. 362**). When these drugs are administered to patients with decompensated liver disease, a proportion improve sufficiently to postpone imminent liver transplantation. In addition, antiviral therapy can be used to prevent recurrence of HBV infection when administered *prior to* transplantation; to treat hepatitis B that recurs *after* transplantation, including in patients who break through HBIg prophylaxis; and to reverse the course of otherwise fatal fibrosing cholestatic hepatitis. Clinical trials have shown that lamivudine antiviral therapy reduces the level of HBV replication substantially, sometimes even resulting in clearance of hepatitis B surface antigen (HBsAg); reduces alanine aminotransferase (ALT) levels; and improves histologic features of necrosis and inflammation. Long-term use of lamivudine is safe and effective, but after several months, a proportion of patients become resistant to lamivudine, resulting from YMDD (tyrosine-methionine-aspartate-aspartate) mutations in the HBV polymerase motif (**Chap. 362**). In approximately one-half of such resistant patients, hepatic deterioration may ensue. Fortunately, adefovir and tenofovir disoproxil fumarate are available as well and can be used to treat lamivudine-associated YMDD variants, effectively “rescuing” patients experiencing hepatic decompensation after lamivudine breakthrough. Currently, most liver transplantation centers combine HBIg plus lamivudine, adefovir, entecavir, or tenofovir disoproxil fumarate. In low-risk patients with no detectable hepatitis B viremia at the time of transplantation, a number of clinical trials have suggested that antiviral prophylaxis can suffice, without HBIg or with a finite duration of HBIg, to prevent recurrent HBV infection of the allograft. Antiviral prophylactic approaches applied to patients undergoing liver transplantation for chronic hepatitis B are being used as well for patients without hepatitis B who receive organs from donors with antibody to hepatitis B core antigen (anti-HBc). Patients who undergo liver transplantation for chronic hepatitis B plus D are less likely to experience recurrent liver injury than patients undergoing liver transplantation for hepatitis B alone; still, such co-infected patients would also be offered standard posttransplantation prophylactic therapy for hepatitis B.

Accounting for up to 40% of all liver transplantation procedures, the most common indication for liver transplantation is end-stage liver disease resulting from chronic hepatitis C. Recurrence of HCV infection after liver transplantation can be documented in almost every patient. The clinical consequences of recurrent hepatitis C are limited during the first 5 years after transplantation. Nonetheless, despite the relative clinical benignity of recurrent hepatitis C in the early years after liver transplantation, and despite the negligible impact on patient survival during these early years, histologic studies have documented the presence of moderate to severe chronic hepatitis in more than one-half of all patients and cirrhosis in $\sim 20\%$ at 5 years. Allograft cirrhosis is even more common, occurring in up to two-thirds of patients at 5 years, if moderate hepatitis is detected in a 1-year liver biopsy. Not surprisingly, then, for patients undergoing liver transplantation for hepatitis C, allograft and patient survival are diminished substantially between 5 and 10 years after transplantation.

In a proportion of patients, even during the early posttransplantation period, recurrent hepatitis C may be sufficiently severe biochemically and histologically to merit antiviral therapy. Treatment with pegylated IFN can *suppress* HCV-associated liver injury but rarely leads to *sustained* benefit. Sustained virologic responses are the exception, and reduced tolerability is often dose-limiting. Preemptive combination antiviral therapy with pegylated IFN and the nucleoside analogue ribavirin immediately after transplantation does not appear to provide any advantage over therapy introduced after clinical hepatitis has occurred. Similarly, although IFN-based antiviral therapy is not recommended for patients with decompensated liver disease, some centers have experimented with pretransplantation antiviral therapy in an attempt to eradicate HCV replication prior to transplantation; preliminary results are promising, but IFN treatment of patients with

369 Diseases of the Gallbladder and Bile Ducts

Norton J. Greenberger, Gustav Paumgartner

end-stage liver disease can lead to worsening of hepatic decompensation, and HCV infection has recurred after transplantation in some of these recipients. Trials of hepatitis C immune globulin preparations to prevent recurrent hepatitis C after liver transplantation have not been successful. Similarly, a trial of a high-dose monoclonal antibody to the HCV E2 envelope glycoprotein delayed but did not prevent reappearance of viremia.

Although the current standard-of-care treatment of allograft hepatitis C is pegylated IFN and ribavirin, in a number of studies, the safety and efficacy of the addition of the approved HCV protease inhibitors telaprevir or boceprevir to pegylated IFN and ribavirin in genotype 1–infected patients with recurrent hepatitis C have been examined. Because of the profound inhibitory effects of the HCV protease inhibitors on the metabolism of the calcineurin inhibitors (increasing cyclosporine levels almost 5-fold and tacrolimus levels 70-fold), calcineurin inhibitor doses must be reduced to safe levels in these patients. In one multicenter study, treatment with a telaprevir- or boceprevir-based triple-drug regimen (with pegylated IFN and ribavirin) achieved rates of HCV clearance similar to those achieved in patients with chronic hepatitis C who had not undergone transplantation. Unfortunately, tolerability of these protease inhibitor–based regimens remains problematic in this population, particularly in persons with allograft cirrhosis, in whom the frequency of hepatic decompensation is increased. The approval of several new direct-acting antiviral (DAA) agents and of IFN-free DAA regimens against HCV will have a major impact on the management and outcome of both pretransplantation and posttransplantation HCV infection. Such therapeutic approaches (1) permit the clearance of viremia in a substantial proportion of decompensated cirrhotics, thereby preventing recurrent allograft infection and, possibly, even improving the clinical status of these patients, delaying or obviating the need for liver replacement; and (2) achieve sustained virologic responses in a much higher proportion of persons with allograft HCV infection, because of improvements in antiviral treatment efficacy and tolerability.

A small number of allograft recipients succumb to early HCV-associated liver injury, and a syndrome reminiscent of fibrosing cholestatic hepatitis (see above) has been observed rarely. Because patients with more episodes of rejection receive more immunosuppressive therapy, and because immunosuppressive therapy enhances HCV replication, patients with severe or multiple episodes of rejection are more likely to experience early recurrence of hepatitis C after transplantation. Both high viral levels and older donor age have been linked to recurrent HCV-induced liver disease and to earlier disease recurrence after transplantation.

Patients who undergo liver transplantation for end-stage alcoholic cirrhosis are at risk of resorting to drinking again after transplantation, a potential source of recurrent alcoholic liver injury. Currently, alcoholic liver disease is one of the more common indications for liver transplantation, accounting for 20–25% of all liver transplantation procedures, and most transplantation centers screen candidates carefully for predictors of continued abstinence. Recidivism is more likely in patients whose sobriety prior to transplantation was <6 months. For abstinent patients with alcoholic cirrhosis, liver transplantation can be undertaken successfully, with outcomes comparable to those for other categories of patients with chronic liver disease, when coordinated by a team approach that includes substance abuse counseling.

POSTTRANSPLANTATION QUALITY OF LIFE

Full rehabilitation is achieved in the majority of patients who survive the early postoperative months and escape chronic rejection or unmanageable infection. Psychosocial maladjustment interferes with medical compliance in a small number of patients, but most manage to adhere to immunosuppressive regimens, which must be continued indefinitely. In one study, 85% of patients who survived their transplant operations returned to gainful activities. In fact, some women have conceived and carried pregnancies to term after transplantation without demonstrable injury to their infants.

PHYSIOLOGY OF BILE PRODUCTION AND FLOW

BILE SECRETION AND COMPOSITION

Bile formed in the hepatic lobules is secreted into a complex network of canaliculi, small bile ductules, and larger bile ducts that run with lymphatics and branches of the portal vein and hepatic artery in portal tracts situated between hepatic lobules. These interlobular bile ducts coalesce to form larger septal bile ducts that join to form the right and left hepatic ducts, which in turn, unite to form the common hepatic duct. The common hepatic duct is joined by the cystic duct of the gallbladder to form the common bile duct (CBD), which enters the duodenum (often after joining the main pancreatic duct) through the ampulla of Vater.

Hepatic bile is an isotonic fluid with an electrolyte composition resembling blood plasma. The electrolyte composition of gallbladder bile differs from that of hepatic bile because most of the inorganic anions, chloride and bicarbonate, have been removed by reabsorption across the gallbladder epithelium. As a result of water reabsorption, total solute concentration of bile increases from 3–4 g/dL in hepatic bile to 10–15 g/dL in gallbladder bile.

Major solute components of bile by moles percent include bile acids (80%), lecithin and traces of other phospholipids (16%), and unesterified cholesterol (4.0%). In the lithogenic state, the cholesterol value can be as high as 8–10%. Other constituents include conjugated bilirubin; proteins (all immunoglobulins, albumin, metabolites of hormones, and other proteins metabolized in the liver); electrolytes; mucus; and, often, drugs and their metabolites.

The total daily basal secretion of hepatic bile is ~500–600 mL. Many substances taken up or synthesized by the hepatocyte are secreted into the bile canaliculi. The canalicular membrane forms microvilli and is associated with microfilaments of actin, microtubules, and other contractile elements. Prior to their secretion into the bile, many substances are taken up into the hepatocyte, while others, such as phospholipids, a portion of primary bile acids, and some cholesterol, are synthesized *de novo* in the hepatocyte. Three mechanisms are important in regulating bile flow: (1) active transport of bile acids from hepatocytes into the bile canaliculi, (2) active transport of other organic anions, and (3) cholangiocellular secretion. The last is a secretin-mediated and cyclic AMP–dependent mechanism that results in the secretion of a sodium- and bicarbonate-rich fluid into the bile ducts.

Active vectorial secretion of biliary constituents from the portal blood into the bile canaliculi is driven by a set of polarized transport systems at the basolateral (sinusoidal) and the canalicular apical plasma membrane domains of the hepatocyte. Two sinusoidal bile salt uptake systems have been cloned in humans, the Na⁺/taurocholate cotransporter (NTCP, SLC10A1) and the organic anion–transporting proteins (OATPs), which also transport a large variety of non-bile salt organic anions. Several ATP-dependent canalicular transport systems, “export pumps,” (ATP-binding cassette transport proteins, also known as ABC transporters) have been identified, the most important of which are: the bile salt export pump (BSEP, ABCB11); the anionic conjugate export pump (MRP2, ABCC2), which mediates the canalicular excretion of various amphiphilic conjugates formed by phase II conjugation (e.g., bilirubin mono- and diglucuronides and drugs); the multidrug export pump (MDR1, ABCB1) for hydrophobic cationic compounds; and the phospholipid export pump (MDR3, ABCB4). Two hemitransporters ABCG5/G8, functioning as a couple, constitute the canalicular cholesterol and phytosterol transporter. FIC1 (ATP8B1) is an aminophospholipid transferase (“flippase”) essential for maintaining the lipid asymmetry of the canalicular membrane. The canalicular membrane also contains ATP-independent transport systems such as the Cl/HCO₃ anion exchanger isoform 2 (AE2, SLC4A2)

2076 for canalicular bicarbonate secretion. For most of these transporters, genetic defects have been identified that are associated with various forms of cholestasis or defects of biliary excretion. FIC1 is defective in progressive familial intrahepatic cholestasis type 1 (PFIC1) and benign recurrent intrahepatic cholestasis type 1 (BRIC1) and results in ablation of all other ATP-dependent transporter functions. BSEP is defective in PFIC2 and BRIC2. Mutations of MRP2 (ABCC2) cause the Dubin-Johnson syndrome, an inherited form of conjugated hyperbilirubinemia (Chap. 359). A defective MDR3 (ABCB4) results in PFIC3. ABCG5/G8, the canalicular half transporters for cholesterol and other neutral sterols, are defective in sitosterolemia. The cystic fibrosis transmembrane regulator (CFTR, ABCC7) located on bile duct epithelial cells but not on canalicular membranes is defective in cystic fibrosis, which is associated with impaired cholangiocellular pH regulation during ductular bile formation and chronic cholestatic liver disease, occasionally resulting in biliary cirrhosis.

THE BILE ACIDS

The primary bile acids, cholic acid and chenodeoxycholic acid (CDCA), are synthesized from cholesterol in the liver, conjugated with glycine or taurine, and secreted into the bile. Secondary bile acids, including deoxycholate and lithocholate, are formed in the colon as bacterial metabolites of the primary bile acids. However, lithocholic acid is much less efficiently absorbed from the colon than deoxycholic acid. Another secondary bile acid, found in low concentration, is ursodeoxycholic acid (UDCA), a stereoisomer of CDCA. In healthy subjects, the ratio of glycine to taurine conjugates in bile is ~3:1.

Bile acids are detergent-like molecules that in aqueous solutions and above a critical concentration of about 2 mM form molecular aggregates called *micelles*. Cholesterol alone is sparingly soluble in aqueous environments, and its solubility in bile depends on both the total lipid concentration and the relative molar percentages of bile acids and lecithin. Normal ratios of these constituents favor the formation of solubilizing *mixed micelles*, while abnormal ratios promote the precipitation of cholesterol crystals in bile via an intermediate liquid crystal phase.

In addition to facilitating the biliary excretion of cholesterol, bile acids facilitate the normal intestinal absorption of dietary fats, mainly cholesterol and fat-soluble vitamins, via a micellar transport mechanism (Chap. 349). Bile acids also serve as a major physiologic driving force for hepatic bile flow and aid in water and electrolyte transport in the small bowel and colon.

ENTEROHEPATIC CIRCULATION

Bile acids are efficiently conserved under normal conditions. Unconjugated, and to a lesser degree also conjugated, bile acids are absorbed by *passive diffusion* along the entire gut. Quantitatively much more important for bile salt recirculation, however, is the *active transport* mechanism for conjugated bile acids in the distal ileum (Chap. 349). The reabsorbed bile acids enter the portal bloodstream and are taken up rapidly by hepatocytes, reconstituted, and resecreted into bile (enterohepatic circulation).

The normal bile acid pool size is approximately 2–4 g. During digestion of a meal, the bile acid pool undergoes at least one or more enterohepatic cycles, depending on the size and composition of the meal. Normally, the bile acid pool circulates ~5–10 times daily. Intestinal reabsorption of the pool is about 95% efficient; therefore, fecal loss of bile acids is in the range of 0.2–0.4 g/d. In the steady state, this fecal loss is compensated by an equal daily synthesis of bile acids by the liver, and, thus, the size of the bile acid pool is maintained. Bile acids in the intestine release fibroblast growth factor 19 (FGF19) into the circulation, which is transported to the liver where it suppresses synthesis of bile acids from cholesterol by inhibiting the rate-limiting enzyme cytochrome P450 7A1 (CYP7A1) and also promotes gallbladder relaxation. While the loss of bile salts in stool is usually matched by increased hepatic synthesis, the maximum rate of synthesis is ~5 g/d, which may be insufficient to replete the bile acid pool size when there is pronounced impairment of intestinal bile salt reabsorption.

The expression of ABC transporters in the enterohepatic circulation and of the rate-limiting enzymes of bile acid and cholesterol synthesis

are regulated in a coordinated fashion by nuclear receptors, which are ligand-activated transcription factors. The hepatic BSEP (ABCB11) is upregulated by the farnesoid X receptor (FXR), a bile acid sensor that also represses bile acid synthesis. The expression of the cholesterol transporter, ABCG5/G8, is upregulated by the liver X receptor (LXR), which is an oxysterol sensor.

GALLBLADDER AND SPHINCTERIC FUNCTIONS

In the fasting state, the sphincter of Oddi offers a high-pressure zone of resistance to bile flow from the CBD into the duodenum. Its tonic contraction serves to (1) prevent reflux of duodenal contents into the pancreatic and bile ducts and (2) promote filling of the gallbladder. The major factor controlling the evacuation of the gallbladder is the peptide hormone cholecystokinin (CCK), which is released from the duodenal mucosa in response to the ingestion of fats and amino acids. CCK produces (1) powerful contraction of the gallbladder, (2) decreased resistance of the sphincter of Oddi, and (3) enhanced flow of biliary contents into the duodenum.

Hepatic bile is “concentrated” within the gallbladder by energy-dependent transmucosal absorption of water and electrolytes. Almost the entire bile acid pool may be sequestered in the gallbladder following an overnight fast for delivery into the duodenum with the first meal of the day. The normal capacity of the gallbladder is ~30 mL of bile.

DISEASES OF THE GALLBLADDER

CONGENITAL ANOMALIES

Anomalies of the biliary tract are not uncommon and include abnormalities in number, size, and shape (e.g., agenesis of the gallbladder, duplications, rudimentary or oversized “giant” gallbladders, and diverticula). *Phrygian cap* is a clinically innocuous entity in which a partial or complete septum (or fold) separates the fundus from the body. Anomalies of position or suspension are not uncommon and include left-sided gallbladder, intrahepatic gallbladder, retrodisplacement of the gallbladder, and “floating” gallbladder. The latter condition predisposes to acute torsion, volvulus, or herniation of the gallbladder.

GALLSTONES

Epidemiology and Pathogenesis Gallstones are quite prevalent in most Western countries. Gallstone formation increases after age 50. In the United States, the third National Health and Nutrition Examination Survey (NHANES III) has revealed an overall prevalence of gallstones of 7.9% in men and 16.6% in women. The prevalence was high in Mexican Americans (8.9% in men, 26.7% in women), intermediate for non-Hispanic whites (8.6% in men, 16.6% in women), and low for African Americans (5.3% in men, 13.9% in women).

Gallstones are formed because of abnormal bile composition. They are divided into two major types: cholesterol stones and pigment stones. Cholesterol stones account for more than 90% of all gallstones in Western industrialized countries. Cholesterol gallstones usually contain >50% cholesterol monohydrate plus an admixture of calcium salts, bile pigments, proteins, and fatty acids. Pigment stones are composed primarily of calcium bilirubinate; they contain <20% cholesterol and are classified into “black” and “brown” types, the latter forming secondary to chronic biliary infection.

CHOLESTEROL STONES AND BILIARY SLUDGE Cholesterol is essentially water insoluble and requires aqueous dispersion into either micelles or vesicles, both of which require the presence of a second lipid to solubilize the cholesterol. Cholesterol and phospholipids are secreted into bile as unilamellar bilayered vesicles, which are converted into mixed micelles consisting of bile acids, phospholipids, and cholesterol by the action of bile acids. If there is an excess of cholesterol in relation to phospholipids and bile acids, unstable, cholesterol-rich vesicles remain, which aggregate into large multilamellar vesicles from which cholesterol crystals precipitate (Fig. 369-1).

There are several important mechanisms in the formation of lithogenic (stone-forming) bile. The most important is increased biliary secretion of cholesterol. This may occur in association with obesity, the metabolic syndrome, high-caloric and cholesterol-rich diets,

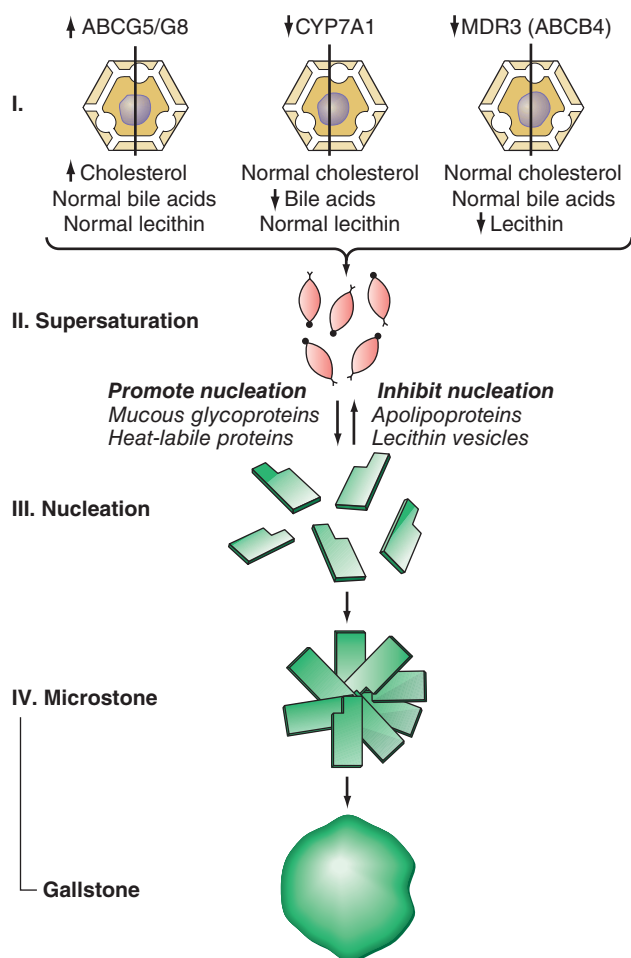


FIGURE 369-1 Scheme showing pathogenesis of cholesterol gallstone formation. Conditions or factors that increase the ratio of cholesterol to bile acids and phospholipids (lecithin) favor gallstone formation. ABCB4, ATP-binding cassette transporter; ABCG5/8, ATP-binding cassette (ABC) transporter G5/G8; CYP7A1, cytochrome P450 7A1; MDR3, multidrug resistance protein 3, also called phospholipid export pump.

or drugs (e.g., clofibrate) and may result from increased activity of hydroxymethylglutaryl-coenzyme A (HMG-CoA) reductase, the rate-limiting enzyme of hepatic cholesterol synthesis, and increased hepatic uptake of cholesterol from blood. In patients with gallstones, dietary cholesterol *increases* biliary cholesterol secretion. This does not occur in non-gallstone patients on high-cholesterol diets. In addition to environmental factors such as high-caloric and cholesterol-rich diets, genetic factors play an important role in gallstone disease. A large study of symptomatic gallstones in Swedish twins provided strong evidence for a role of genetic factors in gallstone pathogenesis. Genetic factors accounted for 25%, shared environmental factors for 13%, and individual environmental factors for 62% of the phenotypic variation among monozygotic twins. A single nucleotide polymorphism of the gene encoding the hepatic cholesterol transporter ABCG5/G8 has been found in 21% of patients with gallstones, but only in 9% of the general population. It is thought to cause a gain of function of the cholesterol transporter and to contribute to cholesterol hypersecretion. A high prevalence of gallstones is found among first-degree relatives of gallstone carriers and in certain ethnic populations such as American Indians, Chilean Indians, and Chilean Hispanics. A common genetic trait has been identified for some of these populations by mitochondrial DNA analysis. In some patients, impaired hepatic conversion of cholesterol to bile acids may also occur, resulting in an increase of the lithogenic cholesterol/bile acid ratio. Although most cholesterol stones have a polygenic basis, there are rare monogenic (Mendelian) causes.

Recently, a mutation in the *CYP7A1* gene has been described that results in a deficiency of the enzyme cholesterol 7-hydroxylase, which catalyzes the initial step in cholesterol catabolism and bile acid synthesis. The homozygous state is associated with hypercholesterolemia and gallstones. Because the phenotype is expressed in the heterozygote state, mutations in the *CYP7A1* gene may contribute to the susceptibility to cholesterol gallstone disease in the population. Mutations in the *MDR3* (*ABCB4*) gene, which encodes the phospholipid export pump in the canalicular membrane of the hepatocyte, may cause defective phospholipid secretion into bile, resulting in cholesterol supersaturation of bile and formation of cholesterol gallstones in the gallbladder and in the bile ducts. Thus, an excess of biliary cholesterol in relation to bile acids and phospholipids is primarily due to hypersecretion of cholesterol, but hyposecretion of bile acids or phospholipids may contribute. An additional disturbance of bile acid metabolism that is likely to contribute to supersaturation of bile with cholesterol is enhanced conversion of cholic acid to deoxycholic acid, with replacement of the cholic acid pool by an expanded deoxycholic acid pool. It may result from enhanced dehydroxylation of cholic acid and increased absorption of newly formed deoxycholic acid. An increased deoxycholate secretion is associated with hypersecretion of cholesterol into bile.

While supersaturation of bile with cholesterol is an important prerequisite for gallstone formation, it is generally not sufficient by itself to produce cholesterol precipitation *in vivo*. Most individuals with supersaturated bile do not develop stones because the time required for cholesterol crystals to nucleate and grow is longer than the time bile remains in the gallbladder.

An important mechanism is *nucleation* of cholesterol monohydrate crystals, which is greatly accelerated in human lithogenic bile. Accelerated nucleation of cholesterol monohydrate in bile may be due to either an *excess of pronucleating factors* or a *deficiency of antinucleating factors*. Mucin and certain nonmucin glycoproteins, principally immunoglobulins, appear to be pronucleating factors, while apolipoproteins A-I and A-II and other glycoproteins appear to be antinucleating factors. Pigment particles may possibly play a role as nucleating factors. In a genome-wide analysis of serum bilirubin levels, the uridine diphosphate-glucuronyltransferase 1A1 (*UGT1A1*) Gilbert's syndrome gene variant was associated with the presence of gallstone disease. Because most gallstones associated with the *UGT1A1* variant were cholesterol stones, this finding points to the role of pigment particles in the pathogenesis of gallbladder stones. Cholesterol monohydrate crystal nucleation and crystal growth probably occur within the mucin gel layer. Vesicle fusion leads to liquid crystals, which, in turn, nucleate into solid cholesterol monohydrate crystals. Continued growth of the crystals occurs by direct nucleation of cholesterol molecules from supersaturated unilamellar or multilamellar biliary vesicles.

A third important mechanism in cholesterol gallstone formation is *gallbladder hypomotility*. If the gallbladder emptied all supersaturated or crystal-containing bile completely, stones would not be able to grow. A high percentage of patients with gallstones exhibit abnormalities of gallbladder emptying. Ultrasonographic studies show that gallstone patients display an increased gallbladder volume during fasting and also after a test meal (residual volume) and that fractional emptying after gallbladder stimulation is decreased. The incidence of gallstones is increased in conditions associated with infrequent or impaired gallbladder emptying such as fasting, parenteral nutrition, or pregnancy and in patients using drugs that inhibit gallbladder motility.

Biliary sludge is a thick, mucous material that, upon microscopic examination, reveals lecithin-cholesterol liquid crystals, cholesterol monohydrate crystals, calcium bilirubinate, and mucin gels. Biliary sludge typically forms a crescent-like layer in the most dependent portion of the gallbladder and is recognized by characteristic echoes on ultrasonography (see below). The presence of biliary sludge implies two abnormalities: (1) the normal balance between gallbladder mucin secretion and elimination has become deranged, and (2) nucleation of biliary solutes has occurred. That biliary sludge may be a precursor form of gallstone disease is evident from several observations. In one study, 96 patients with gallbladder sludge were followed prospectively by serial ultrasound studies. In 18%, biliary sludge disappeared and did

2078 not recur for at least 2 years. In 60%, biliary sludge disappeared and reappeared; in 14%, gallstones (8% asymptomatic, 6% symptomatic) developed; and in 6%, severe biliary pain with or without acute pancreatitis occurred. In 12 patients, cholecystectomies were performed, 6 for gallstone-associated biliary pain and 3 in symptomatic patients with sludge but without gallstones who had prior attacks of pancreatitis; the latter did not recur after cholecystectomy. It should be emphasized that biliary sludge can develop with disorders that cause gallbladder hypomotility; i.e., surgery, burns, total parenteral nutrition, pregnancy, and oral contraceptives—all of which are associated with gallstone formation. However, the presence of biliary sludge implies supersaturation of bile with either cholesterol or calcium bilirubinate.

Two other conditions are associated with cholesterol-stone or biliary-sludge formation: pregnancy and rapid weight reduction through a very-low-calorie diet. There appear to be two key changes during pregnancy that contribute to a “cholelithogenic state”: (1) a marked increase in cholesterol saturation of bile during the third trimester and (2) sluggish gallbladder contraction in response to a standard meal, resulting in impaired gallbladder emptying. That these changes are related to pregnancy per se is supported by several studies that show reversal of these abnormalities quite rapidly after delivery. During pregnancy, gallbladder sludge develops in 20–30% of women and gallstones in 5–12%. Although biliary sludge is a common finding during pregnancy, it is usually asymptomatic and often resolves spontaneously after delivery. Gallstones, which are less common than sludge and frequently associated with biliary colic, may also disappear after delivery because of spontaneous dissolution related to bile becoming unsaturated with cholesterol postpartum.

Approximately 10–20% of persons with rapid weight reduction achieved through very-low-calorie dieting develop gallstones. In a study involving 600 patients who completed a 3-month, 520-kcal/d diet, UDCA in a dosage of 600 mg/d proved highly effective in preventing gallstone formation; gallstones developed in only 3% of UDCA recipients, compared to 28% of placebo-treated patients. In obese patients treated by gastric banding, 500 mg/d of UDCA reduced the risk of gallstone formation from 30% to 8% within a follow-up of 6 months.

To summarize, cholesterol gallstone disease occurs because of several defects, which include (1) bile supersaturation with cholesterol, (2) nucleation of cholesterol monohydrate with subsequent crystal retention and stone growth, and (3) abnormal gallbladder motor function with delayed emptying and stasis. Other important factors known to predispose to cholesterol-stone formation are summarized in **Table 369-1**.

PIGMENT STONES Black pigment stones are composed of either pure calcium bilirubinate or polymer-like complexes with calcium and mucin glycoproteins. They are more common in patients who have chronic hemolytic states (with increased conjugated bilirubin in bile), liver cirrhosis, Gilbert’s syndrome, or cystic fibrosis. Gallbladder stones in patients with ileal diseases, ileal resection, or ileal bypass generally are also black pigment stones. Enterohepatic recycling of bilirubin in ileal disease states contributes to their pathogenesis. Brown pigment stones are composed of calcium salts of unconjugated bilirubin with varying amounts of cholesterol and protein. They are caused by the presence of increased amounts of unconjugated, insoluble bilirubin in bile that precipitates to form stones. Deconjugation of an excess of soluble bilirubin mono- and diglucuronides may be mediated by endogenous β -glucuronidase but may also occur by spontaneous hydrolysis. Sometimes, the enzyme is also produced when bile is chronically infected by bacteria, and such stones are brown. Pigment stone formation is frequent in Asia and is often associated with infections in the gallbladder and biliary tree (Table 369-1).

Diagnosis Procedures of potential use in the diagnosis of cholelithiasis and other diseases of the gallbladder are detailed in **Table 369-2**. Ultrasonography of the gallbladder is very accurate in the identification of cholelithiasis and has replaced oral cholecystography (**Fig. 369-2A**). Stones as small as 1.5 mm in diameter may be confidently identified provided that firm criteria are used (e.g., acoustic “shadowing” of opacities that are within the gallbladder lumen and that change with the patient’s position [by gravity]). In major medical centers, the false-negative and

TABLE 369-1 PREDISPOSING FACTORS FOR CHOLESTEROL AND PIGMENT GALLSTONE FORMATION

Cholesterol Stones	
1.	Demographic/genetic factors: Prevalence highest in North American Indians, Chilean Indians, and Chilean Hispanics, greater in Northern Europe and North America than in Asia, lowest in Japan; familial disposition; hereditary aspects
2.	Obesity, metabolic syndrome: Normal bile acid pool and secretion but increased biliary secretion of cholesterol
3.	Weight loss: Mobilization of tissue cholesterol leads to increased biliary cholesterol secretion while enterohepatic circulation of bile acids is decreased
4.	Female sex hormones <ol style="list-style-type: none"> Estrogens stimulate hepatic lipoprotein receptors, increase uptake of dietary cholesterol, and increase biliary cholesterol secretion Natural estrogens, other estrogens, and oral contraceptives lead to decreased bile salt secretion and decreased conversion of cholesterol to cholesteryl esters
5.	Pregnancy: Impaired gallbladder emptying caused by progesterone combined with the influence of estrogens, which increase biliary cholesterol secretion
6.	Increasing age: Increased biliary secretion of cholesterol, decreased size of bile acid pool, decreased secretion of bile salts
7.	Gallbladder hypomotility leading to stasis and formation of sludge <ol style="list-style-type: none"> Prolonged parenteral nutrition Fasting Pregnancy Drugs such as octreotide
8.	Clofibrate therapy: Increased biliary secretion of cholesterol
9.	Decreased bile acid secretion <ol style="list-style-type: none"> Primary biliary cirrhosis Genetic defect of the <i>CYP7A1</i> gene
10.	Decreased phospholipid secretion: Genetic defect of the <i>MDR3</i> gene
11.	Miscellaneous <ol style="list-style-type: none"> High-calorie, high-fat diet Spinal cord injury
Pigment Stones	
1.	Demographic/genetic factors: Asia, rural setting
2.	Chronic hemolysis
3.	Alcoholic liver cirrhosis
4.	Pernicious anemia
5.	Cystic fibrosis
6.	Chronic biliary tract infection, parasite infections
7.	Increasing age
8.	Ileal disease, ileal resection or bypass

false-positive rates for ultrasound in gallstone patients are ~2–4%. Biliary sludge is material of low echogenic activity that typically forms a layer in the most dependent position of the gallbladder. This layer shifts with postural changes but fails to produce acoustic shadowing; these two characteristics distinguish sludges from gallstones. Ultrasound can also be used to assess the emptying function of the gallbladder.

The plain abdominal film may detect gallstones containing sufficient calcium to be radiopaque (10–15% of cholesterol and ~50% of pigment stones). Plain radiography may also be of use in the diagnosis of emphysematous cholecystitis, porcelain gallbladder, limey bile, and gallstone ileus.

Oral cholecystography (OCG) has historically been a useful procedure for the diagnosis of gallstones but has been replaced by ultrasound and is regarded as obsolete. It may be used to assess the patency of the cystic duct and gallbladder emptying function. Further, OCG can also delineate the size and number of gallstones and determine whether they are calcified.

Radiopharmaceuticals such as ^{99m}Tc -labeled *N*-substituted iminodiacetic acids (HIDA, DIDA, DISIDA, etc.) are rapidly extracted from

TABLE 369-2 DIAGNOSTIC EVALUATION OF THE GALLBLADDER

Diagnostic Advantages	Diagnostic Limitations	Comment
Gallbladder Ultrasound		
Rapid	Bowel gas	Procedure of choice for detection of stones
Accurate identification of gallstones (>95%)	Massive obesity	
Simultaneous scanning of GB, liver, bile ducts, pancreas	Ascites	
"Real-time" scanning allows assessment of GB volume, contractility		
Not limited by jaundice, pregnancy		
May detect very small stones		
Plain Abdominal X-Ray		
Low cost	Relatively low yield	Pathognomonic findings in: calcified gallstones
Readily available	? Contraindicated in pregnancy	Limey bile, porcelain GB Emphysematous cholecystitis Gallstone ileus
Radioisotope Scans (HIDA, DIDA, etc.)		
Accurate identification of cystic duct obstruction	? Contraindicated in pregnancy	Indicated for confirmation of suspected acute cholecystitis; less sensitive and less specific in chronic cholecystitis; useful in diagnosis of acalculous cholecystopathy, especially if given with CCK to assess gallbladder emptying
Simultaneous assessment of bile ducts	Serum bilirubin >103–205 μmol/L (6–12 mg/dL)	
	Cholecystogram of low resolution	

Abbreviations: CCK, cholecystokinin; GB, gallbladder.

the blood and are excreted into the biliary tree in high concentration even in the presence of mild to moderate serum bilirubin elevations. Failure to image the gallbladder in the presence of biliary ductal visualization may indicate cystic duct obstruction, acute or chronic cholecystitis, or surgical absence of the organ. Such scans have some application in the diagnosis of acute cholecystitis.

Symptoms of Gallstone Disease Gallstones usually produce symptoms by causing inflammation or obstruction following their migration into the cystic duct or CBD. The most specific and characteristic symptom of gallstone disease is biliary colic that is a constant and often long-lasting pain (see below). Obstruction of the cystic duct or CBD by a stone produces increased intraluminal pressure and distention of the viscus that cannot be relieved by repetitive biliary contractions. The resultant visceral pain is characteristically a severe, steady ache or fullness in

the epigastrium or right upper quadrant (RUQ) of the abdomen with frequent radiation to the interscapular area, right scapula, or shoulder.

Biliary colic begins quite suddenly and may persist with severe intensity for 30 min to 5 h, subsiding gradually or rapidly. It is steady rather than intermittent, as would be suggested by the word *colic*, which must be regarded as a misnomer, although it is in widespread use. An episode of biliary pain persisting beyond 5 h should raise the suspicion of acute cholecystitis (see below). Nausea and vomiting frequently accompany episodes of biliary pain. An elevated level of serum bilirubin and/or alkaline phosphatase suggests a common duct stone. Fever or chills (rigors) with biliary pain usually imply a complication, i.e., cholecystitis, pancreatitis, or cholangitis. Complaints of short-lasting, vague epigastric fullness, dyspepsia, eructation, or flatulence, especially following a fatty meal, should not be confused with biliary pain. Such symptoms are frequently elicited from patients with or without gallstone disease

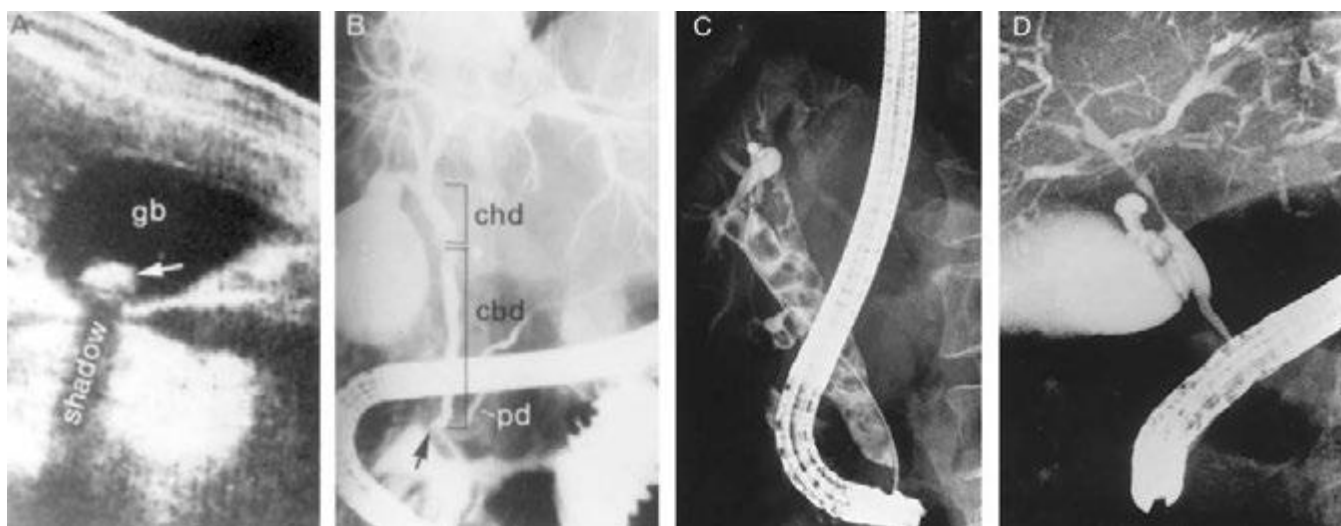


FIGURE 369-2 Examples of ultrasound and radiologic studies of the biliary tract. **A.** An ultrasound study showing a distended gallbladder (GB) containing a single large stone (*arrow*), which casts an acoustic shadow. **B.** Endoscopic retrograde cholangiopancreatogram (ERCP) showing normal biliary tract anatomy. In addition to the endoscope and large vertical gallbladder filled with contrast dye, the common hepatic duct (CHD), common bile duct (CBD), and pancreatic duct (PD) are shown. The *arrow* points to the ampulla of Vater. **C.** Endoscopic retrograde cholangiogram (ERC) showing choledocholithiasis. The biliary tract is dilated and contains multiple radiolucent calculi. **D.** ERCP showing sclerosing cholangitis. The common bile duct shows areas that are strictured and narrowed.

2080 but are not specific for biliary calculi. Biliary colic may be precipitated by eating a fatty meal, by consumption of a large meal following a period of prolonged fasting, or by eating a normal meal; it is frequently nocturnal, occurring within a few hours of retiring.

Natural History Gallstone disease discovered in an asymptomatic patient or in a patient whose symptoms are not referable to cholelithiasis is a common clinical problem. Sixty to 80% of persons with asymptomatic gallstones remain asymptomatic over follow-up periods of up to 25 years. The probability of developing symptoms within 5 years after diagnosis is 2–4% per year and decreases in the years thereafter to 1–2%. The yearly incidence of complications is about 0.1–0.3%. Patients remaining asymptomatic for 15 years were found to be unlikely to develop symptoms during further follow-up, and most patients who did develop complications from their gallstones experienced *prior* warning symptoms. Similar conclusions apply to diabetic patients with silent gallstones. Decision analysis has suggested that (1) the cumulative risk of death due to gallstone disease while on expectant management is small, and (2) prophylactic cholecystectomy is not warranted.

Complications requiring cholecystectomy are much more common in gallstone patients who have developed symptoms of biliary pain. Patients found to have gallstones at a young age are more likely to develop symptoms from cholelithiasis than are patients >60 years at the time of initial diagnosis. Patients with diabetes mellitus and gallstones may be somewhat more susceptible to septic complications, but the magnitude of risk of septic biliary complications in diabetic patients is incompletely defined.

TREATMENT GALLSTONES

SURGICAL THERAPY

In asymptomatic gallstone patients, the risk of developing symptoms or complications requiring surgery is quite small (see above). Thus, a recommendation for cholecystectomy in a patient with gallstones should probably be based on assessment of three factors: (1) the presence of symptoms that are frequent enough or severe enough to interfere with the patient's general routine; (2) the presence of a prior complication of gallstone disease, i.e., history of acute cholecystitis, pancreatitis, gallstone fistula, etc.; or (3) the presence of an underlying condition predisposing the patient to increased risk of gallstone complications (e.g., calcified or porcelain gallbladder and/or a previous attack of acute cholecystitis regardless of current symptomatic status). Patients with very large gallstones (>3 cm in diameter) and patients harboring gallstones in a congenitally anomalous gallbladder might also be considered for prophylactic cholecystectomy. Although young age is a worrisome factor in asymptomatic gallstone patients, few authorities would now recommend routine cholecystectomy in all young patients with silent stones. Laparoscopic cholecystectomy is a minimal-access approach for the removal of the gallbladder together with its stones. Its advantages include a markedly shortened hospital stay, minimal disability, and decreased cost, and it is the procedure of choice for most patients referred for elective cholecystectomy.

From several studies involving >4000 patients undergoing laparoscopic cholecystectomy, the following key points emerge: (1) complications develop in ~4% of patients, (2) conversion to laparotomy occurs in 5%, (3) the death rate is remarkably low (i.e., <0.1%), and (4) the rate of bile duct injuries is low (i.e., 0.2–0.6%) and comparable with open cholecystectomy. These data indicate why laparoscopic cholecystectomy has become the "gold standard" for treating symptomatic cholelithiasis.

MEDICAL THERAPY—GALLSTONE DISSOLUTION

In carefully selected patients with a functioning gallbladder and with radiolucent stones <10 mm in diameter, complete dissolution can be achieved in ~50% of patients within 6 months to 2 years. For good results within a reasonable time period, this therapy should be limited to radiolucent stones smaller than 5 mm in diameter. The dose of UDCA should be 10–15 mg/kg per day. Stones larger than

10 mm in size rarely dissolve. Pigment stones are not responsive to UDCA therapy. Probably ≤10% of patients with *symptomatic* cholelithiasis are candidates for such treatment. However, in addition to the vexing problem of recurrent stones (30–50% over 3–5 years of follow-up), there is also the factor of taking an expensive drug for up to 2 years. The advantages and success of laparoscopic cholecystectomy have largely reduced the role of gallstone dissolution to patients who wish to avoid or are not candidates for elective cholecystectomy. However, patients with cholesterol gallstone disease who develop recurrent choledocholithiasis after cholecystectomy should be on long-term treatment with UDCA.

ACUTE AND CHRONIC CHOLECYSTITIS

Acute Cholecystitis Acute inflammation of the gallbladder wall usually follows obstruction of the cystic duct by a stone. Inflammatory response can be evoked by three factors: (1) *mechanical inflammation* produced by increased intraluminal pressure and distention with resulting ischemia of the gallbladder mucosa and wall, (2) *chemical inflammation* caused by the release of lysolecithin (due to the action of phospholipase on lecithin in bile) and other local tissue factors, and (3) *bacterial inflammation*, which may play a role in 50–85% of patients with acute cholecystitis. The organisms most frequently isolated by culture of gallbladder bile in these patients include *Escherichia coli*, *Klebsiella* spp., *Streptococcus* spp., and *Clostridium* spp.

Acute cholecystitis often begins as an attack of biliary pain that progressively worsens. Approximately 60–70% of patients report having experienced prior attacks that resolved spontaneously. As the episode progresses, however, the pain of acute cholecystitis becomes more generalized in the right upper abdomen. As with biliary colic, the pain of cholecystitis may radiate to the interscapular area, right scapula, or shoulder. Peritoneal signs of inflammation such as increased pain with jarring or on deep respiration may be apparent. The patient is anorectic and often nauseated. Vomiting is relatively common and may produce symptoms and signs of vascular and extracellular volume depletion. Jaundice is unusual early in the course of acute cholecystitis but may occur when edematous inflammatory changes involve the bile ducts and surrounding lymph nodes.

A low-grade fever is characteristically present, but shaking chills or rigors are not uncommon. The RUQ of the abdomen is almost invariably tender to palpation. An enlarged, tense gallbladder is palpable in 25–50% of patients. Deep inspiration or cough during subcostal palpation of the RUQ usually produces increased pain and inspiratory arrest (Murphy's sign). Localized rebound tenderness in the RUQ is common, as are abdominal distention and hypoactive bowel sounds from paralytic ileus, but generalized peritoneal signs and abdominal rigidity are usually lacking, in the absence of perforation.

The diagnosis of acute cholecystitis is usually made on the basis of a characteristic history and physical examination. The triad of sudden onset of RUQ tenderness, fever, and leukocytosis is highly suggestive. Typically, leukocytosis in the range of 10,000–15,000 cells per microliter with a left shift on differential count is found. The serum bilirubin is mildly elevated (<85.5 μmol/L [5 mg/dL]) in fewer than half of patients, whereas about one-fourth have modest elevations in serum aminotransferases (usually less than a fivefold elevation). Ultrasound will demonstrate calculi in 90–95% of cases and is useful for detection of signs of gallbladder inflammation including thickening of the wall, pericholecystic fluid, and dilatation of the bile duct. The radionuclide (e.g., HIDA) biliary scan may be confirmatory if bile duct imaging is seen without visualization of the gallbladder.

Approximately 75% of patients treated medically have remission of acute symptoms within 2–7 days following hospitalization. In 25%, however, a complication of acute cholecystitis will occur despite conservative treatment (see below). In this setting, prompt surgical intervention is required. Of the 75% of patients with acute cholecystitis who undergo remission of symptoms, ~25% will experience a recurrence of cholecystitis within 1 year, and 60% will have at least one recurrent bout within 6 years. In view of the natural history of the disease, acute cholecystitis is best treated by early surgery whenever possible.

Mirizzi's syndrome is a rare complication in which a gallstone becomes impacted in the cystic duct or neck of the gallbladder causing compression of the CBD, resulting in CBD obstruction and jaundice. Ultrasound shows gallstone(s) lying outside the hepatic duct. Endoscopic retrograde cholangiopancreatography (ERCP) (Fig. 369-2B), percutaneous transhepatic cholangiography (PTC), or magnetic resonance cholangiopancreatography (MRCP) will usually demonstrate the characteristic extrinsic compression of the CBD. Surgery consists of removing the cystic duct, diseased gallbladder, and the impacted stone. The preoperative diagnosis of Mirizzi's syndrome is important to avoid CBD injury.

ACALCULOUS CHOLECYSTITIS In 5–10% of patients with acute cholecystitis, calculi obstructing the cystic duct are not found at surgery. In >50% of such cases, an underlying explanation for acalculous inflammation is not found. An increased risk for the development of acalculous cholecystitis is especially associated with serious trauma or burns, with the postpartum period following prolonged labor, and with orthopedic and other nonbiliary major surgical operations in the postoperative period. It may possibly complicate periods of prolonged parenteral hyperalimentation. For some of these cases, biliary sludge in the cystic duct may be responsible. Other precipitating factors include vasculitis, obstructing adenocarcinoma of the gallbladder, diabetes mellitus, torsion of the gallbladder, "unusual" bacterial infections of the gallbladder (e.g., *Leptospira*, *Streptococcus*, *Salmonella*, or *Vibrio cholerae*), and parasitic infestation of the gallbladder. Acalculous cholecystitis may also be seen with a variety of other systemic disease processes (e.g., sarcoidosis, cardiovascular disease, tuberculosis, syphilis, actinomycosis).

Although the clinical manifestations of acalculous cholecystitis are indistinguishable from those of calculous cholecystitis, the setting of acute gallbladder inflammation complicating severe underlying illness is characteristic of acalculous disease. Ultrasound or computed tomography (CT) examinations demonstrating a large, tense, static gallbladder without stones and with evidence of poor emptying over a prolonged period may be diagnostically useful in some cases. The complication rate for acalculous cholecystitis exceeds that for calculous cholecystitis. Successful management of acute acalculous cholecystitis appears to depend primarily on early diagnosis and surgical intervention, with meticulous attention to postoperative care.

ACALCULOUS CHOLECYSTOPATHY Disordered motility of the gallbladder can produce recurrent biliary pain in patients without gallstones. Infusion of an octapeptide of CCK can be used to measure the gallbladder ejection fraction during cholescintigraphy. The surgical findings have included abnormalities such as chronic cholecystitis, gallbladder muscle hypertrophy, and/or a markedly narrowed cystic duct. Some of these patients may well have had antecedent gallbladder disease. The following criteria can be used to identify patients with acalculous cholecystopathy: (1) recurrent episodes of typical RUQ pain characteristic of biliary tract pain, (2) abnormal CCK cholescintigraphy demonstrating a gallbladder ejection fraction of <40%, and (3) infusion of CCK reproducing the patient's pain. An additional clue would be the identification of a large gallbladder on ultrasound examination. Finally, it should be noted that sphincter of Oddi dysfunction can also give rise to recurrent RUQ pain and CCK-scintigraphic abnormalities.

EMPHYSEMATOUS CHOLECYSTITIS So-called emphysematous cholecystitis is thought to begin with acute cholecystitis (calculous or acalculous) followed by ischemia or gangrene of the gallbladder wall and infection by gas-producing organisms. Bacteria most frequently cultured in this setting include anaerobes, such as *Clostridium welchii* or *Clostridium perfringens*, and aerobes, such as *E. coli*. This condition occurs most frequently in elderly men and in patients with diabetes mellitus. The clinical manifestations are essentially indistinguishable from those of nongaseous cholecystitis. The diagnosis is usually made on plain abdominal film by finding gas within the gallbladder lumen, dissecting within the gallbladder wall to form a gaseous ring, or in the pericholecystic tissues. The morbidity and mortality rates with emphysematous cholecystitis are considerable. Prompt surgical intervention coupled with appropriate antibiotics is mandatory.

Chronic Cholecystitis Chronic inflammation of the gallbladder wall is almost always associated with the presence of gallstones and is thought to result from repeated bouts of subacute or acute cholecystitis or from persistent mechanical irritation of the gallbladder wall by gallstones. The presence of bacteria in the bile occurs in >25% of patients with chronic cholecystitis. The presence of infected bile in a patient with chronic cholecystitis undergoing elective cholecystectomy probably adds little to the operative risk. Chronic cholecystitis may be asymptomatic for years, may progress to symptomatic gallbladder disease or to acute cholecystitis, or may present with complications (see below).

Complications of Cholecystitis • EMPYEMA AND HYDROPS Empyema of the gallbladder usually results from progression of acute cholecystitis with persistent cystic duct obstruction to superinfection of the stagnant bile with a pus-forming bacterial organism. The clinical picture resembles that of cholangitis with high fever; severe RUQ pain; marked leukocytosis; and often, prostration. Empyema of the gallbladder carries a high risk of gram-negative sepsis and/or perforation. Emergency surgical intervention with proper antibiotic coverage is required as soon as the diagnosis is suspected.

Hydrops or mucocele of the gallbladder may also result from prolonged obstruction of the cystic duct, usually by a large solitary calculus. In this instance, the obstructed gallbladder lumen is progressively distended, over a period of time, by mucus (mucocele) or by a clear transudate (hydrops) produced by mucosal epithelial cells. A visible, easily palpable, nontender mass sometimes extending from the RUQ into the right iliac fossa may be found on physical examination. The patient with hydrops of the gallbladder frequently remains asymptomatic, although chronic RUQ pain may also occur. Cholecystectomy is indicated, because empyema, perforation, or gangrene may complicate the condition.

GANGRENE AND PERFORATION Gangrene of the gallbladder results from ischemia of the wall and patchy or complete tissue necrosis. Underlying conditions often include marked distention of the gallbladder, vasculitis, diabetes mellitus, empyema, or torsion resulting in arterial occlusion. Gangrene usually predisposes to perforation of the gallbladder, but perforation may also occur in chronic cholecystitis without premonitory warning symptoms. *Localized perforations* are usually contained by the omentum or by adhesions produced by recurrent inflammation of the gallbladder. Bacterial superinfection of the walled-off gallbladder contents results in abscess formation. Most patients are best treated with cholecystectomy, but some seriously ill patients may be managed with cholecystostomy and drainage of the abscess. *Free perforation* is less common but is associated with a mortality rate of ~30%. Such patients may experience a sudden transient relief of RUQ pain as the distended gallbladder decompresses; this is followed by signs of generalized peritonitis.

FISTULA FORMATION AND GALLSTONE ILEUS Fistula formation into an adjacent organ adherent to the gallbladder wall may result from inflammation and adhesion formation. Fistulas into the duodenum are most common, followed in frequency by those involving the hepatic flexure of the colon, stomach or jejunum, abdominal wall, and renal pelvis. Clinically "silent" biliary-enteric fistulas occurring as a complication of acute cholecystitis have been found in up to 5% of patients undergoing cholecystectomy. Asymptomatic cholecystoenteric fistulas may sometimes be diagnosed by finding gas in the biliary tree on plain abdominal films. Barium contrast studies or endoscopy of the upper gastrointestinal tract or colon may demonstrate the fistula. Treatment in the symptomatic patient usually consists of cholecystectomy, CBD exploration, and closure of the fistulous tract.

Gallstone ileus refers to mechanical intestinal obstruction resulting from the passage of a large gallstone into the bowel lumen. The stone customarily enters the duodenum through a cholecystoenteric fistula at that level. The site of obstruction by the impacted gallstone is usually at the ileocecal valve, provided that the more proximal small bowel is of normal caliber. The majority of patients do not give a history of either prior biliary tract symptoms or complaints suggestive of acute cholecystitis or fistula formation. Large stones, >2.5 cm in diameter, are

2082 thought to predispose to fistula formation by gradual erosion through the gallbladder fundus. Diagnostic confirmation may occasionally be found on the plain abdominal film (e.g., small-intestinal obstruction with gas in the biliary tree and a calcified, ectopic gallstone) or following an upper gastrointestinal series (cholecystoduodenal fistula with small-bowel obstruction at the ileocecal valve). Laparotomy with stone extraction (or propulsion into the colon) remains the procedure of choice to relieve obstruction. Evacuation of large stones within the gallbladder should also be performed. In general, the gallbladder and its attachment to the intestines should be left alone.

LIMEY (MILK OF CALCIUM) BILE AND PORCELAIN GALLBLADDER Calcium salts in the lumen of the gallbladder in sufficient concentration may produce calcium precipitation and diffuse, hazy opacification of bile or a layering effect on plain abdominal roentgenography. This so-called limey bile, or milk of calcium bile, is usually clinically innocuous, but cholecystectomy is recommended, especially when it occurs in a hydropic gallbladder. In the entity called *porcelain gallbladder*, calcium salt deposition within the wall of a chronically inflamed gallbladder may be detected on the plain abdominal film. Cholecystectomy is advised in all patients with porcelain gallbladder because in a high percentage of cases this finding appears to be associated with the development of carcinoma of the gallbladder.

TREATMENT ACUTE CHOLECYSTITIS

MEDICAL THERAPY

Although surgical intervention remains the mainstay of therapy for acute cholecystitis and its complications, a period of in-hospital stabilization may be required before cholecystectomy. Oral intake is eliminated, nasogastric suction may be indicated, and extracellular volume depletion and electrolyte abnormalities are repaired. Meperidine or nonsteroidal anti-inflammatory drugs (NSAIDs) are usually employed for analgesia because they may produce less spasm of the sphincter of Oddi than drugs such as morphine. Intravenous antibiotic therapy is usually indicated in patients with severe acute cholecystitis, even though bacterial superinfection of bile may not have occurred in the early stages of the inflammatory process. Antibiotic therapy is guided by the most common organisms likely to be present, which are *E. coli*, *Klebsiella* spp., and *Streptococcus* spp. Effective antibiotics include ureidopenicillins such as piperacillin or mezlocillin, ampicillin sulbactam, ciprofloxacin, moxifloxacin, and third-generation cephalosporins. Anaerobic coverage by a drug such as metronidazole should be added if gangrenous or emphysematous cholecystitis is suspected. Imipenem and meropenem represent potent parenteral antibiotics that cover the whole spectrum of bacteria causing ascending cholangitis. They should, however, be reserved for the most severe, life-threatening infections when other regimens have failed (**Chap. 186**). Postoperative complications of wound infection, abscess formation, and sepsis are reduced in antibiotic-treated patients.

SURGICAL THERAPY

The optimal timing of surgical intervention in patients with acute cholecystitis depends on stabilization of the patient. The clear trend is toward earlier surgery, and this is due in part to requirements for shorter hospital stays. Urgent (emergency) cholecystectomy or cholecystostomy is probably appropriate in most patients in whom a complication of acute cholecystitis such as empyema, emphysematous cholecystitis, or perforation is suspected or confirmed. Patients with uncomplicated acute cholecystitis should undergo early elective laparoscopic cholecystectomy, ideally within 48–72 h after diagnosis. The complication rate is not increased in patients undergoing early as opposed to delayed (>6 weeks after diagnosis) cholecystectomy. Delayed surgical intervention is probably best reserved for (1) patients in whom the overall medical condition imposes an unacceptable risk for early surgery and (2) patients in whom the diagnosis of acute cholecystitis is in doubt. Thus, early cholecystectomy (within 72 h) is the treatment of choice for most

patients with acute cholecystitis. Mortality figures for emergency cholecystectomy in most centers range from 1–3%, whereas the mortality risk for early elective cholecystectomy is ~0.5% in patients under age 60. Of course, the operative risks increase with age-related diseases of other organ systems and with the presence of long- or short-term complications of gallbladder disease. Seriously ill or debilitated patients with cholecystitis may be managed with cholecystostomy and tube drainage of the gallbladder. Elective cholecystectomy may then be done at a later date.

Postcholecystectomy Complications Early complications following cholecystectomy include atelectasis and other pulmonary disorders, abscess formation (often subphrenic), external or internal hemorrhage, biliary-enteric fistula, and bile leaks. Jaundice may indicate absorption of bile from an intraabdominal collection following a biliary leak or mechanical obstruction of the CBD by retained calculi, intraductal blood clots, or extrinsic compression.

Overall, cholecystectomy is a very successful operation that provides total or near-total relief of preoperative symptoms in 75–90% of patients. The most common cause of persistent postcholecystectomy symptoms is an overlooked symptomatic nonbiliary disorder (e.g., reflux esophagitis, peptic ulceration, pancreatitis, or—most often—irritable bowel syndrome). In a small percentage of patients, however, a disorder of the extrahepatic bile ducts may result in persistent symptomatology. These so-called postcholecystectomy syndromes may be due to (1) biliary strictures, (2) retained biliary calculi, (3) cystic duct stump syndrome, (4) stenosis or dyskinesia of the sphincter of Oddi, or (5) bile salt–induced diarrhea or gastritis.

CYSTIC DUCT STUMP SYNDROME In the absence of cholangiographically demonstrable retained stones, symptoms resembling biliary pain or cholecystitis in the postcholecystectomy patient have frequently been attributed to disease in a long (>1 cm) cystic duct remnant (cystic duct stump syndrome). Careful analysis, however, reveals that postcholecystectomy complaints are attributable to other causes in almost all patients in whom the symptom complex was originally thought to result from the existence of a long cystic duct stump. Accordingly, considerable care should be taken to investigate the possible role of other factors in the production of postcholecystectomy symptoms before attributing them to cystic duct stump syndrome.

PAPILLARY DYSFUNCTION, PAPILLARY STENOSIS, SPASM OF THE SPHINCTER OF ODDI, AND BILIARY DYSKINESIA Symptoms of biliary colic accompanied by signs of recurrent, intermittent biliary obstruction may be produced by acalculous cholecystopathy, papillary stenosis, papillary dysfunction, spasm of the sphincter of Oddi, and biliary dyskinesia. Papillary stenosis is thought to result from acute or chronic inflammation of the papilla of Vater or from glandular hyperplasia of the papillary segment. Five criteria have been used to define papillary stenosis: (1) upper abdominal pain, usually RUQ or epigastric; (2) abnormal liver tests; (3) dilatation of the CBD upon ERCP examination; (4) delayed (>45 min) drainage of contrast material from the duct; and (5) increased basal pressure of the sphincter of Oddi, a finding that may be of only minor significance. An alternative to ERCP is magnetic resonance cholangiography (MRC) if ERCP and/or biliary manometry are either unavailable or not feasible. After exclusion of acalculous cholecystopathy, treatment consists of endoscopic or surgical sphincteroplasty to ensure wide patency of the distal portions of both the bile and pancreatic ducts. The greater the number of the preceding criteria present, the greater is the likelihood that a patient does have a degree of papillary stenosis sufficient to justify correction. The factors usually considered as indications for sphincterotomy include (1) prolonged duration of symptoms, (2) lack of response to symptomatic treatment, (3) presence of severe disability, and (4) the patient's choice of sphincterotomy over surgery (given a clear understanding on his or her part of the risks involved in both procedures).

Criteria for diagnosing dyskinesia of the sphincter of Oddi are even more controversial than those for papillary stenosis. Proposed mechanisms include spasm of the sphincter, denervation sensitivity resulting

in hypertonicity, and abnormalities of the sequencing or frequency rates of sphincter-contraction waves. When thorough evaluation has failed to demonstrate another cause for the pain, and when cholangiographic and manometric criteria suggest a diagnosis of biliary dyskinesia, medical treatment with nitrates or anticholinergics to attempt pharmacologic relaxation of the sphincter has been proposed. Endoscopic biliary sphincterotomy (EBS) or surgical sphincteroplasty may be indicated in patients who fail to respond to a 2- to 3-month trial of medical therapy, especially if basal sphincter of Oddi pressures are elevated. EBS has become the procedure of choice for removing bile duct stones and for other biliary and pancreatic problems.

BILE SALT-INDUCED DIARRHEA AND GASTRITIS Postcholecystectomy patients may develop symptoms of dyspepsia, which have been attributed to duodenogastric reflux of bile. However, firm data linking these symptoms to bile gastritis after surgical removal of the gallbladder are lacking. Cholecystectomy induces persistent changes in gut transit, and these changes effect a noticeable modification of bowel habits. Cholecystectomy shortens gut transit time by accelerating passage of the fecal bolus through the colon with marked acceleration in the right colon, thus causing an increase in colonic bile acid output and a shift in bile acid composition toward the more diarrheagenic secondary bile acids, i.e. deoxycholic acid. Diarrhea that is severe enough, i.e., three or more watery movements per day, can be classified as postcholecystectomy diarrhea, and this occurs in 5–10% of patients undergoing elective cholecystectomy. Treatment with bile acid-sequestering agents such as cholestyramine or colestipol is often effective in ameliorating troublesome diarrhea.

THE HYPERPLASTIC CHOLECYSTOSES

The term *hyperplastic cholecystoses* is used to denote a group of disorders of the gallbladder characterized by excessive proliferation of normal tissue components.

Adenomyomatosis is characterized by a benign proliferation of gallbladder surface epithelium with glandlike formations, extramural sinuses, transverse strictures, and/or fundal nodule (“adenoma” or “adenomyoma”) formation.

Cholesterolosis is characterized by abnormal deposition of lipid, especially cholesteryl esters, within macrophages in the lamina propria of the gallbladder wall. In its diffuse form (“strawberry gallbladder”), the gallbladder mucosa is brick red and speckled with bright yellow flecks of lipid. The localized form shows solitary or multiple “cholesterol polyps” studding the gallbladder wall. Cholesterol stones of the gallbladder are found in nearly half the cases. Cholecystectomy is indicated in both adenomyomatosis and cholesterolosis when symptomatic or when cholelithiasis is present.

The prevalence of gallbladder polyps in the adult population is ~5%, with a marked male predominance. Few significant changes have been found over a 5-year period in asymptomatic patients with gallbladder polyps <10 mm in diameter. Cholecystectomy is recommended in symptomatic patients, as well as in asymptomatic patients >50 years of age, or in those whose polyps are >10 mm in diameter or associated with gallstones or polyp growth on serial ultrasonography.

DISEASES OF THE BILE DUCTS

CONGENITAL ANOMALIES

Biliary Atresia and Hypoplasia Atretic and hypoplastic lesions of the extrahepatic and large intrahepatic bile ducts are the most common biliary anomalies of clinical relevance encountered in infancy. The clinical picture is one of severe obstructive jaundice during the first month of life, with pale stools. When biliary atresia is suspected on the basis of clinical, laboratory, and imaging findings, the diagnosis is confirmed by surgical exploration and operative cholangiography. Approximately 10% of cases of biliary atresia are treatable with Roux-en-Y choledochojejunostomy, with the Kasai procedure (hepatic portoenterostomy) being attempted in the remainder in an effort to restore some bile flow. Most patients, even those having successful biliary-enteric anastomoses, eventually develop chronic cholangitis, extensive hepatic fibrosis, and portal hypertension.

Choledochal Cysts Cystic dilatation may involve the free portion of the CBD, i.e., choledochal cyst, or may present as diverticulum formation in the intraduodenal segment. In the latter situation, chronic reflux of pancreatic juice into the biliary tree can produce inflammation and stenosis of the extrahepatic bile ducts leading to cholangitis or biliary obstruction. Because the process may be gradual, ~50% of patients present with onset of symptoms after age 10. The diagnosis may be made by ultrasound, abdominal CT, MRC, or cholangiography. Only one-third of patients show the classic triad of abdominal pain, jaundice, and an abdominal mass. Ultrasonographic detection of a cyst separate from the gallbladder should suggest the diagnosis of choledochal cyst, which can be confirmed by demonstrating the entrance of extrahepatic bile ducts into the cyst. Surgical treatment involves excision of the “cyst” and biliary-enteric anastomosis. Patients with choledochal cysts are at increased risk for the subsequent development of cholangiocarcinoma.

Congenital Biliary Ectasia Dilatation of intrahepatic bile ducts may involve either the major intrahepatic radicles (Caroli’s disease), the inter- and intralobular ducts (congenital hepatic fibrosis), or both. In Caroli’s disease, clinical manifestations include recurrent cholangitis, abscess formation in and around the affected ducts, and, often, brown pigment gallstone formation within portions of ectatic intrahepatic biliary radicles. Ultrasound, MRC, and CT are of great diagnostic value in demonstrating cystic dilatation of the intrahepatic bile ducts. Treatment with ongoing antibiotic therapy is usually undertaken in an effort to limit the frequency and severity of recurrent bouts of cholangitis. Progression to secondary biliary cirrhosis with portal hypertension, extrahepatic biliary obstruction, cholangiocarcinoma, or recurrent episodes of sepsis with hepatic abscess formation is common.

CHOLEDOCHOLITHIASIS

Pathophysiology and Clinical Manifestations Passage of gallstones into the CBD occurs in ~10–15% of patients with cholelithiasis. The incidence of common duct stones increases with increasing age of the patient, so that up to 25% of elderly patients may have calculi in the common duct at the time of cholecystectomy. Undetected duct stones are left behind in ~1–5% of cholecystectomy patients. The overwhelming majority of bile duct stones are cholesterol stones formed in the gallbladder, which then migrate into the extrahepatic biliary tree through the cystic duct. Primary calculi arising de novo in the ducts are usually brown pigment stones developing in patients with (1) hepatobiliary parasitism or chronic, recurrent cholangitis; (2) congenital anomalies of the bile ducts (especially Caroli’s disease); (3) dilated, sclerosed, or strictured ducts; or (4) an *MDR3* (*ABCB4*) gene defect leading to impaired biliary phospholipids secretion (low phospholipid-associated cholesterol cholelithiasis). Common duct stones may remain asymptomatic for years, may pass spontaneously into the duodenum, or (most often) may present with biliary colic or a complication.

Complications • CHOLANGITIS Cholangitis may be acute or chronic, and symptoms result from inflammation, which usually is caused by at least partial obstruction to the flow of bile. Bacteria are present on bile culture in ~75% of patients with acute cholangitis early in the symptomatic course. The characteristic presentation of acute cholangitis involves biliary pain, jaundice, and spiking fevers with chills (Charcot’s triad). Blood cultures are frequently positive, and leukocytosis is typical. *Nonsuppurative acute cholangitis* is most common and may respond relatively rapidly to supportive measures and to treatment with antibiotics. In *suppurative acute cholangitis*, however, the presence of pus under pressure in a completely obstructed ductal system leads to symptoms of severe toxicity—mental confusion, bacteremia, and septic shock. Response to antibiotics alone in this setting is relatively poor, multiple hepatic abscesses are often present, and the mortality rate approaches 100% unless prompt endoscopic or surgical relief of the obstruction and drainage of infected bile are carried out. Endoscopic management of bacterial cholangitis is as effective as surgical intervention. ERCP with endoscopic sphincterotomy is safe and the preferred initial procedure for both establishing a definitive diagnosis and providing effective therapy.

2084 OBSTRUCTIVE JAUNDICE Gradual obstruction of the CBD over a period of weeks or months usually leads to initial manifestations of jaundice or pruritus without associated symptoms of biliary colic or cholangitis. Painless jaundice may occur in patients with choledocholithiasis, but is much more characteristic of biliary obstruction secondary to malignancy of the head of the pancreas, bile ducts, or ampulla of Vater.

In patients whose obstruction is secondary to choledocholithiasis, associated chronic calculous cholecystitis is very common, and the gallbladder in this setting may be unable to distend. The absence of a palpable gallbladder in most patients with biliary obstruction from duct stones is the basis for Courvoisier's law, i.e., that the presence of a palpably enlarged gallbladder suggests that the biliary obstruction is secondary to an underlying malignancy rather than to calculous disease. Biliary obstruction causes progressive dilatation of the intrahepatic bile ducts as intrabiliary pressures rise. Hepatic bile flow is suppressed, and reabsorption and regurgitation of conjugated bilirubin into the bloodstream lead to jaundice accompanied by dark urine (bilirubinuria) and light-colored (acholic) stools.

CBD stones should be suspected in any patient with cholecystitis whose serum bilirubin level is $>85.5 \mu\text{mol/L}$ (5 mg/dL). The maximum bilirubin level is seldom $>256.5 \mu\text{mol/L}$ (15.0 mg/dL) in patients with choledocholithiasis unless concomitant hepatic or renal disease or another factor leading to marked hyperbilirubinemia exists. Serum bilirubin levels $\geq 342.0 \mu\text{mol/L}$ (20 mg/dL) should suggest the possibility of neoplastic obstruction. The serum alkaline phosphatase level is almost always elevated in biliary obstruction. A rise in alkaline phosphatase often precedes clinical jaundice and may be the only abnormality in routine liver function tests. There may be a two- to tenfold elevation of serum aminotransferases, especially in association with acute obstruction. Following relief of the obstructing process, serum aminotransferase elevations usually return rapidly to normal, while the serum bilirubin level may take 1–2 weeks to return to normal. The alkaline phosphatase level usually falls slowly, lagging behind the decrease in serum bilirubin.

PANCREATITIS The most common associated entity discovered in patients with nonalcoholic acute pancreatitis is biliary tract disease. Biochemical evidence of pancreatic inflammation complicates acute cholecystitis in 15% of cases and choledocholithiasis in $>30\%$, and the common factor appears to be the passage of gallstones through the common duct. Coexisting pancreatitis should be suspected in patients with symptoms of cholecystitis who develop (1) back pain or pain to the left of the abdominal midline, (2) prolonged vomiting with paralytic ileus, or (3) a pleural effusion, especially on the left side. Surgical treatment of gallstone disease is usually associated with resolution of the pancreatitis.

SECONDARY BILIARY CIRRHOSIS Secondary biliary cirrhosis may complicate prolonged or intermittent duct obstruction with or without recurrent cholangitis. Although this complication may be seen in patients with choledocholithiasis, it is more common in cases of prolonged obstruction from stricture or neoplasm. Once established, secondary biliary cirrhosis may be progressive even after correction of the obstructing process, and increasingly severe hepatic cirrhosis may lead to portal hypertension or to hepatic failure and death. Prolonged biliary obstruction may also be associated with clinically relevant deficiencies of the fat-soluble vitamins A, D, E, and K.

Diagnosis and Treatment The diagnosis of choledocholithiasis is usually made by cholangiography (Table 369-3), either preoperatively by endoscopic retrograde cholangiogram (ERC) (Fig. 369-2C) or MRCP or intraoperatively at the time of cholecystectomy. As many as 15% of patients undergoing cholecystectomy will prove to have CBD stones. When CBD stones are suspected prior to laparoscopic cholecystectomy, preoperative ERCP with endoscopic papillotomy and stone extraction is the preferred approach. It not only provides stone clearance but also defines the anatomy of the biliary tree in relationship to the cystic duct. CBD stones should be suspected in gallstone patients who have any of the following risk factors: (1) a history of jaundice or pancreatitis, (2) abnormal tests of liver function, and

(3) ultrasonographic or MRCP evidence of a dilated CBD or stones in the duct. Alternatively, if intraoperative cholangiography reveals retained stones, postoperative ERCP can be carried out. The need for preoperative ERCP is expected to decrease further as laparoscopic techniques for bile duct exploration improve.

The widespread use of laparoscopic cholecystectomy and ERCP has decreased the incidence of complicated biliary tract disease and the need for choledocholithotomy and T-tube drainage of the bile ducts. EBS followed by spontaneous passage or stone extraction is the treatment of choice in the management of patients with common duct stones, especially in elderly or poor-risk patients.

TRAUMA, STRICTURES, AND HEMOBILIA

Most benign strictures of the extrahepatic bile ducts result from surgical trauma and occur in about 1 in 500 cholecystectomies. Strictures may present with bile leak or abscess formation in the immediate postoperative period or with biliary obstruction or cholangitis as long as 2 years or more following the inciting trauma. The diagnosis is established by percutaneous or endoscopic cholangiography. Endoscopic brushing of biliary strictures may be helpful in establishing the nature of the lesion and is more accurate than bile cytology alone. When positive exfoliative cytology is obtained, the diagnosis of a neoplastic stricture is established. This procedure is especially important in patients with primary sclerosing cholangitis (PSC) who are predisposed to the development of cholangiocarcinomas. Successful operative correction of non-PSC bile duct strictures by a skillful surgeon with duct-to-bowel anastomosis is usually possible, although mortality rates from surgical complications, recurrent cholangitis, or secondary biliary cirrhosis are high.

Hemobilia may follow traumatic or operative injury to the liver or bile ducts, intraductal rupture of a hepatic abscess or aneurysm of the hepatic artery, biliary or hepatic tumor hemorrhage, or mechanical complications of choledocholithiasis or hepatobiliary parasitism. Diagnostic procedures such as liver biopsy, PTC, and transhepatic biliary drainage catheter placement may also be complicated by hemobilia. Patients often present with a classic triad of biliary pain, obstructive jaundice, and melena or occult blood in the stools. The diagnosis is sometimes made by cholangiographic evidence of blood clot in the biliary tree, but selective angiographic verification may be required. Although minor episodes of hemobilia may resolve without operative intervention, surgical ligation of the bleeding vessel is frequently required.

EXTRINSIC COMPRESSION OF THE BILE DUCTS

Partial or complete biliary obstruction may be produced by extrinsic compression of the ducts. The most common cause of this form of obstructive jaundice is carcinoma of the head of the pancreas. Biliary obstruction may also occur as a complication of either acute or chronic pancreatitis or involvement of lymph nodes in the porta hepatis by lymphoma or metastatic carcinoma. The latter should be distinguished from cholestasis resulting from massive replacement of the liver by tumor.

HEPATOBIILIARY PARASITISM

Infestation of the biliary tract by adult helminths or their ova may produce a chronic, recurrent pyogenic cholangitis with or without multiple hepatic abscesses, ductal stones, or biliary obstruction. This condition is relatively rare but does occur in inhabitants of southern China and elsewhere in Southeast Asia. The organisms most commonly involved are trematodes or flukes, including *Clonorchis sinensis*, *Opisthorchis viverrini* or *Opisthorchis felineus*, and *Fasciola hepatica*. The biliary tract also may be involved by intraductal migration of adult *Ascaris lumbricoides* from the duodenum or by intrabiliary rupture of hydatid cysts of the liver produced by *Echinococcus* spp. The diagnosis is made by cholangiography and the presence of characteristic ova on stool examination. When obstruction is present, the treatment of choice is laparotomy under antibiotic coverage, with common duct exploration and a biliary drainage procedure.

SCLEROSING CHOLANGITIS

Primary or idiopathic sclerosing cholangitis is characterized by a progressive, inflammatory, sclerosing, and obliterative process

TABLE 369-3 DIAGNOSTIC EVALUATION OF THE BILE DUCTS

Diagnostic Advantages	Diagnostic Limitations	Contraindications	Complications	Comment
Hepatobiliary Ultrasound				
Rapid	Bowel gas	None	None	Initial procedure of choice in investigating possible biliary tract obstruction
Simultaneous scanning of GB, liver, bile ducts, pancreas	Massive obesity Ascites			
Accurate identification of dilated bile ducts	Barium			
Not limited by jaundice, pregnancy	Partial bile duct obstruction			
Guidance for fine-needle biopsy	Poor visualization of distal CBD			
Computed Tomography				
Simultaneous scanning of GB, liver, bile ducts, pancreas	Extreme cachexia Movement artifact	Pregnancy	Reaction to iodinated contrast, if used	Indicated for evaluation of hepatic or pancreatic masses Procedure of choice in investigating possible biliary obstruction if diagnostic limitations prevent HBUS
Accurate identification of dilated bile ducts, masses	Ileus			
Not limited by jaundice, gas, obesity, ascites	Partial bile duct obstruction			
High-resolution image				
Guidance for fine-needle biopsy				
Magnetic Resonance Cholangiopancreatography				
Useful modality for visualizing pancreatic and biliary ducts	Cannot offer therapeutic intervention	Claustrophobia Certain metals (iron)	None	
Has excellent sensitivity for bile duct dilatation, biliary stricture, and intraductal abnormalities	High cost			
Can identify pancreatic duct dilatation or stricture, pancreatic duct stenosis, and pancreas divisum				
Endoscopic Retrograde Cholangiopancreatography				
Simultaneous pancreatography	Gastroduodenal obstruction	Pregnancy	Pancreatitis	Cholangiogram of choice in: Absence of dilated ducts ? Pancreatic, ampullary or gastroduodenal disease Prior biliary surgery Endoscopic sphincterotomy a treatment possibility
Best visualization of distal biliary tract		? Acute pancreatitis	Cholangitis, sepsis	
Bile or pancreatic cytology	? Roux-en-Y biliary-enteric anastomosis	? Severe cardiopulmonary disease	Infected pancreatic pseudocyst	
Endoscopic sphincterotomy and stone removal			Perforation (rare)	
Biliary manometry			Hypoxemia, aspiration	
Percutaneous Transhepatic Cholangiogram				
Extremely successful when bile ducts dilated	Nondilated or sclerosed ducts	Pregnancy Uncorrectable coagulopathy Massive ascites ? Hepatic abscess	Bleeding Hemobilia Bile peritonitis Bacteremia, sepsis	Indicated when ERCP is contraindicated or failed
Best visualization of proximal biliary tract				
Bile cytology/culture				
Percutaneous transhepatic drainage				
Endoscopic Ultrasound				
Most sensitive method to detect ampullary stones				

Abbreviations: CBD, common bile duct; ERCP, endoscopic retrograde cholangiopancreatography; GB, gallbladder; HBUS, hepatobiliary ultrasound.

affecting the extrahepatic and/or the intrahepatic bile ducts. The disorder occurs up to 75% in association with inflammatory bowel disease, especially ulcerative colitis. It may also be associated with autoimmune pancreatitis; multifocal fibrosclerosis syndromes such as retroperitoneal, mediastinal, and/or periureteral fibrosis; Riedel's struma; or pseudotumor of the orbit.

Immunoglobulin G4 (IgG4)-associated cholangitis is a recently described biliary disease of unknown etiology that presents with biochemical and cholangiographic features indistinguishable from PSC, is often associated with autoimmune pancreatitis and other fibrosing conditions, and is characterized by elevated serum IgG4 and infiltration of IgG4-positive plasma cells in bile ducts and liver tissue. In contrast to PSC, it is not associated with inflammatory bowel disease and should be suspected if associated with increased serum IgG4 and

unexplained pancreatic disease. Glucocorticoids are regarded as the initial treatment of choice. Relapse is common after steroid withdrawal, especially with proximal strictures. Long-term treatment with glucocorticoids and/or azathioprine may be needed after relapse or for inadequate response (**Chap. 371**).

Patients with primary sclerosing cholangitis often present with signs and symptoms of chronic or intermittent biliary obstruction: RUQ abdominal pain, pruritus, jaundice, or acute cholangitis. Late in the course, complete biliary obstruction, secondary biliary cirrhosis, hepatic failure, or portal hypertension with bleeding varices may occur. The diagnosis is usually established by finding multifocal, diffusely distributed strictures with intervening segments of normal or dilated ducts, producing a beaded appearance on cholangiography (**Fig. 369-2D**). The cholangiographic techniques of choice in suspected

2086 cases are MRCP and ERCP. When a diagnosis of sclerosing cholangitis has been established, a search for associated diseases, especially for chronic inflammatory bowel disease, should be carried out.

A recent study describes the natural history and outcome for 305 patients of Swedish descent with primary sclerosing cholangitis; 134 (44%) of the patients were asymptomatic at the time of diagnosis and, not surprisingly, had a significantly higher survival rate. The independent predictors of a bad prognosis were advanced age, serum bilirubin concentration, and liver histologic changes. Cholangiocarcinoma was found in 24 patients (8%). Inflammatory bowel disease was closely associated with primary sclerosing cholangitis and had a prevalence of 81% in this study population.

Small duct PSC is defined by the presence of chronic cholestasis and hepatic histology consistent with PSC but with normal findings on cholangiography. Small duct PSC is found in ~5% of patients with PSC and may represent an earlier stage of PSC associated with a significantly better long-term prognosis. However, such patients may progress to classic PSC and/or end-stage liver disease with consequent necessity of liver transplantation.

In patients with AIDS, cholangiopancreatography may demonstrate a broad range of biliary tract changes as well as pancreatic duct obstruction and occasionally pancreatitis (Chap. 226). Further, biliary tract lesions in AIDS include infection and cholangiopancreatographic changes similar to those of PSC. Changes noted include: (1) diffuse involvement of intrahepatic bile ducts alone, (2) involvement of both intra- and extrahepatic bile ducts, (3) ampullary stenosis, (4) stricture of the intrapancreatic portion of the CBD, and (5) pancreatic duct involvement. Associated infectious organisms include *Cryptosporidium*, *Mycobacterium avium-intracellulare*, cytomegalovirus, *Microsporidia*,

and *Isospora*. In addition, acalculous cholecystitis occurs in up to 10% of patients. ERCP sphincterotomy, while not without risk, provides significant pain reduction in patients with AIDS-associated papillary stenosis. Secondary sclerosing cholangitis may occur as a long-term complication of choledocholithiasis, cholangiocarcinoma, operative or traumatic biliary injury, or contiguous inflammatory processes.

TREATMENT SCLEROSING CHOLANGITIS

Therapy with cholestyramine may help control symptoms of pruritus, and antibiotics are useful when cholangitis complicates the clinical picture. Vitamin D and calcium supplementation may help prevent the loss of bone mass frequently seen in patients with chronic cholestasis. Glucocorticoids, methotrexate, and cyclosporine have not been shown to be efficacious in PSC. UDCA in high dosage (20 mg/kg) improves serum liver tests, but an effect on survival has not been documented. In cases where high-grade biliary obstruction (dominant strictures) has occurred, balloon dilatation or stenting may be appropriate. Only rarely is surgical intervention indicated. Efforts at biliary-enteric anastomosis or stent placement may, however, be complicated by recurrent cholangitis and further progression of the stenosing process. The prognosis is unfavorable, with a median survival of 9–12 years following the diagnosis, regardless of therapy. Four variables (age, serum bilirubin level, histologic stage, and splenomegaly) predict survival in patients with PSC and serve as the basis for a risk score. PSC is one of the most common indications for liver transplantation.

SECTION 3 DISORDERS OF THE PANCREAS

370 Approach to the Patient with Pancreatic Disease

Darwin L. Conwell, Norton J. Greenberger,
Peter A. Banks

GENERAL CONSIDERATIONS

As emphasized in Chap. 371, the etiologies as well as clinical manifestations of pancreatitis are quite varied. Although it is well-appreciated that pancreatitis is frequently secondary to biliary tract disease and alcohol abuse, it can also be caused by drugs, genetic mutations, trauma, and viral infections and is associated with metabolic and connective tissue disorders. In ~30% of patients with acute pancreatitis and 25–40% of patients with chronic pancreatitis, the etiology initially can be obscure.

The incidence of acute pancreatitis is about 5–35/100,000 new cases per year worldwide, with a mortality rate of about 3%. The incidence of chronic pancreatitis is about 4–8 new cases per 100,000 per year with a prevalence of 26–42 cases per 100,000. The number of patients admitted to the hospital who suffer with both acute and chronic pancreatitis in the United States is largely increasing and is now estimated to be 274,119 for acute pancreatitis and 19,724 for chronic pancreatitis. Acute pancreatitis is now the most common gastrointestinal diagnosis requiring hospitalization in the United States. Acute and chronic pancreatic disease costs an estimated 3 billion dollars annually in health care expenditures. These numbers may underestimate the true incidence and prevalence, because non-alcohol-induced pancreatitis has been largely ignored. At autopsy, the prevalence of chronic pancreatitis ranges from 0.04 to 5%.

The diagnosis of acute pancreatitis is generally clearly defined based on a combination of laboratory, imaging, and clinical symptoms. The diagnosis of chronic pancreatitis, especially in mild disease, is hampered by the relative inaccessibility of the pancreas to direct examination and the nonspecificity of the abdominal pain associated with chronic pancreatitis. Many patients with chronic pancreatitis do not have elevated blood amylase or lipase levels. Some patients with chronic pancreatitis develop signs and symptoms of pancreatic exocrine insufficiency, and thus, objective evidence for pancreatic disease can be demonstrated. However, there is a very large reservoir of pancreatic exocrine function. More than 90% of the pancreas must be damaged before maldigestion of fat and protein is manifested. Noninvasive, indirect tests of pancreatic exocrine function (fecal elastase) are much more likely to give abnormal results in patients with obvious advanced pancreatic disease (i.e., pancreatic calcification, steatorrhea, or diabetes mellitus) than in patients with occult disease. Invasive, direct tests of pancreatic secretory function (secretin tests) are the most sensitive and specific tests to detect early chronic pancreatic disease when imaging is equivocal or normal.

TESTS USEFUL IN THE DIAGNOSIS OF PANCREATIC DISEASE

Several tests have proved of value in the evaluation of pancreatic disease. Examples of specific tests and their usefulness in the diagnosis of acute and chronic pancreatitis are summarized in Table 370-1 and Fig. 370-1. At some institutions, pancreatic function tests are available and performed if the diagnosis of chronic pancreatic disease remains a possibility after noninvasive tests (ultrasound, computed tomography [CT], magnetic resonance cholangiopancreatography [MRCP]) or invasive tests (endoscopic retrograde cholangiopancreatography [ERCP], endoscopic ultrasonography [EUS]) have given normal or

TABLE 370-1 TESTS USEFUL IN THE DIAGNOSIS OF ACUTE AND CHRONIC PANCREATITIS AND PANCREATIC TUMORS

Test	Principle	Comment
Pancreatic Enzymes in Body Fluids		
Serum lipase	Pancreatic inflammation leads to increased serum enzyme levels	Enzyme measurement of choice for diagnosis of acute pancreatitis
Amylase		
1. Serum	Pancreatic inflammation leads to increased serum enzyme levels	Simple; reliable if test results are three times the upper limit of normal
2. Urine	Renal clearance of amylase is increased in acute pancreatitis	Infrequently used
3. Ascitic fluid	Disruption of gland or main pancreatic duct leads to increased amylase concentration	Can help establish source of ascites; false positives occur with intestinal obstruction and perforated ulcer; can also measure lipase
4. Pleural fluid	Exudative pleural effusion with pancreatitis	False positives occur with carcinoma of the lung and esophageal perforation
Studies Pertaining to Pancreatic Structure		
Radiologic and radionuclide tests		
1. Plain film of the abdomen	Can be abnormal in acute and chronic pancreatitis	Infrequently used
2. Upper gastrointestinal x-rays		Infrequently used
3. Ultrasonography (US)	Can provide information on edema, inflammation, calcification, pseudocysts, and mass lesions	Simple, noninvasive; sequential studies quite feasible; useful in diagnosis of gallstones; pancreas visualization limited by interference from overlying bowel gas
4. Computed tomography (CT) scan	Permits detailed visualization of pancreas and surrounding structures, pancreatic fluid collection, pseudocyst; assessment of necrosis or interstitial disease	Useful in the diagnosis of pancreatic calcification, dilated pancreatic ducts, and pancreatic tumors; may not be able to distinguish between inflammatory and neoplastic mass lesions
5. Magnetic resonance cholangiopancreatography (MRCP)	Three-dimensional imaging has been used to produce very good images of the pancreatic-biliary ductal system by a noninvasive technique	Has replaced ERCP as a diagnostic test; noninvasive
6. Endoscopic ultrasonography (EUS)	High-frequency transducer used with EUS can produce very high-resolution images and depict changes in the pancreatic duct and parenchyma with great detail	Can be used to assess gallstones, chronic pancreatitis, and pancreatic carcinoma
7. Endoscopic retrograde cholangiopancreatography (ERCP)	Cannulation of pancreatic and common bile duct permits visualization of pancreatic-biliary ductal system	Primarily a therapeutic procedure; invasive
Pancreatic biopsy with US or CT guidance	Percutaneous aspiration biopsy of mass-forming lesions of the pancreas	High diagnostic yield; laparotomy avoided; can be done with EUS for the evaluation of chronic pancreatitis, autoimmune pancreatitis, and pancreatic carcinoma
Tests of Exocrine Pancreatic Function		
Direct stimulation of the pancreas with analysis of duodenal contents		
1. Secretin test	Secretin leads to increased output of pancreatic juice and HCO ₃ ⁻ ; pancreatic secretory response is related to the functional mass of pancreatic tissue	Sensitive enough to detect occult disease; involves duodenal intubation and fluoroscopic placement of gastroduodenal tube; poorly defined normal enzyme response; overlap in chronic pancreatitis; large secretory reserve capacity of the pancreas; currently done at only a few medical centers
2. Endoscopic secretin test	Replaces need for tube placement duodenum	Sensitive enough to detect occult disease; high negative predictive value; avoids intubation and fluoroscopy; requires sedation
Measurement of intraluminal digestion products		
1. Quantitative stool fat determination	Lack of lipolytic enzymes brings about impaired fat digestion	Reliable reference standard for defining severity of malabsorption; does not distinguish between maldigestion and malabsorption
Measurement of pancreatic enzymes in feces		
1. Elastase	Pancreatic secretion of proteolytic enzymes; not degraded in intestine	Diagnostic accuracy best if value is <100 µg/g performed on a solid stool

inconclusive results. In this regard, tests using *direct* stimulation of the pancreas with secretin are the most sensitive.

Pancreatic Enzymes in Body Fluids The serum amylase and lipase levels are widely used as screening tests for acute pancreatitis in the patient with acute abdominal pain or back pain. Values greater than three times the upper limit of normal in combination with epigastric pain strongly suggest the diagnosis if gut perforation or infarction is excluded. In acute pancreatitis, the serum amylase and lipase are usually elevated within 24 h of onset and remain so for 3–7 days. Levels usually return to normal within 7 days unless there is pancreatic ductal disruption, ductal obstruction, or pseudocyst formation. Approximately 85% of patients with acute pancreatitis have a threefold or greater elevated serum lipase and amylase levels. The values may be

normal if (1) there is a delay (of 2–5 days) before blood samples are obtained, (2) the underlying disorder is chronic pancreatitis rather than acute pancreatitis, or (3) hypertriglyceridemia is present. Patients with hypertriglyceridemia and proven pancreatitis have been found to have spuriously low levels of amylase and perhaps lipase activity. In the absence of objective evidence of pancreatitis by abdominal ultrasound, CT scan, MRCP, or EUS, mild to moderate elevations of amylase and/or lipase are not helpful in making a diagnosis of chronic pancreatitis.

The serum amylase can be elevated in other conditions (Table 370-2), in part because the enzyme is found in many organs. In addition to the pancreas and salivary glands, small quantities of amylase are found in the tissues of the fallopian tubes, lung, thyroid, and tonsils and can be produced by various tumors (carcinomas of the lung, esophagus, breast, and ovary). Isoamylase determinations do not accurately

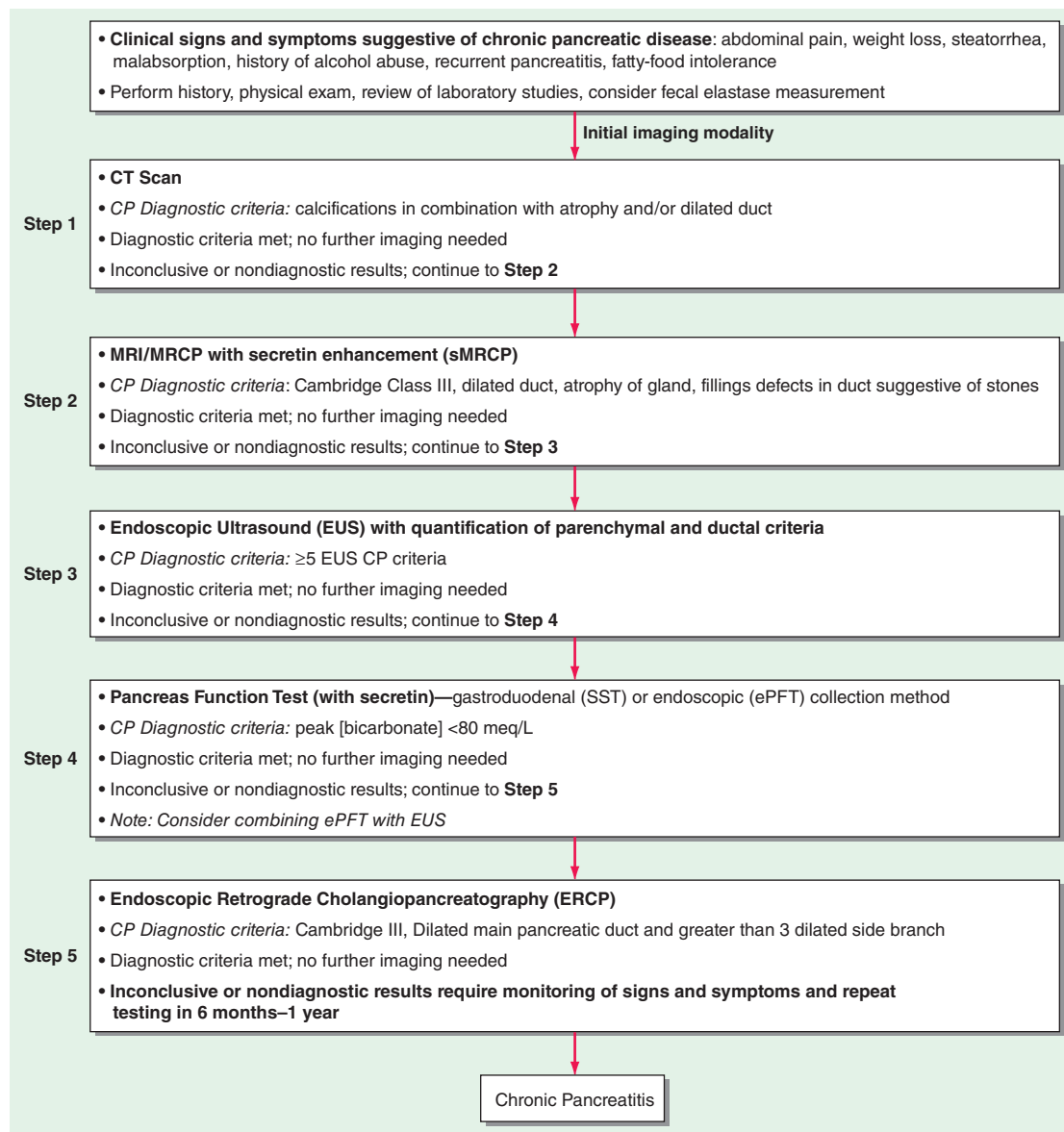


FIGURE 370-1 A stepwise diagnostic approach to the patient with suspected chronic pancreatitis (CP). Endoscopic ultrasonography (EUS) and magnetic resonance cholangiopancreatography (sMRCP/MRCP) are appropriate diagnostic alternatives to endoscopic retrograde cholangiopancreatography (ERCP). CT, computed tomography.

distinguish elevated blood amylase levels due to bona fide pancreatitis from elevated blood amylase levels due to a nonpancreatic source of amylase, especially when the blood amylase level is only moderately elevated. In patients with unexplained hyperamylasemia, measurement of macroamylase can avoid numerous tests in patients with this rare disorder.

Elevation of ascitic fluid amylase occurs in acute pancreatitis as well as in (1) ascites due to disruption of the main pancreatic duct or a leaking pseudocyst and (2) other abdominal disorders that simulate pancreatitis (e.g., intestinal obstruction, intestinal infarction, or perforated peptic ulcer). Elevation of pleural fluid amylase can occur in acute pancreatitis, chronic pancreatitis, carcinoma of the lung, and esophageal perforation. Lipase is the single best enzyme to measure for the diagnosis of acute pancreatitis. No single blood test is reliable for the diagnosis of acute pancreatitis in patients with renal failure. Pancreatic enzyme elevations are usually less than three times the upper limit of normal. Determining whether a patient with renal failure and abdominal pain has pancreatitis remains a difficult clinical problem. One study found that serum amylase levels were elevated in patients with renal dysfunction only when creatinine clearance was < 0.8 mL/s (< 50 mL/min). In such patients, the serum amylase level was invariably < 500 IU/L in the absence of objective evidence of acute pancreatitis. In that study, serum lipase and trypsin levels paralleled serum amylase

values. With these limitations in mind, the recommended screening test for acute pancreatitis in renal disease is serum lipase.

Studies Pertaining to Pancreatic Structure • RADIOLOGIC TESTS Plain films of the abdomen, which once provided useful information in patients with acute and chronic pancreatitis, have been superseded by other more detailed imaging procedures (ultrasound, EUS, CT, MRCP).

Ultrasonography (US) can provide important information in patients with acute pancreatitis, chronic pancreatitis, pseudocysts, and pancreatic carcinoma. Echographic appearances can indicate the presence of edema, inflammation, and calcification (not obvious on plain films of the abdomen), as well as pseudocysts, mass lesions, and gallstones. In acute pancreatitis, the pancreas is characteristically enlarged. In pancreatic pseudocyst, the usual appearance is primarily that of smooth, round fluid collection. Pancreatic carcinoma distorts the usual landmarks, and mass lesions > 3.0 cm are usually detected as localized, solid lesions. US is often the initial investigation for most patients with suspected pancreatic disease. However, obesity and excess small- and large-bowel gas can interfere with pancreatic imaging by US studies.

Computed tomography (CT) is the best imaging study for initial evaluation of a suspected pancreatic disorder and for the assessment of

TABLE 370-2 CAUSES OF HYPERAMYLASEMIA AND HYPERAMYLASURIA

Pancreatic Disease	
I.	Pancreatitis
A.	Acute
B.	Chronic: ductal obstruction
C.	Complications of pancreatitis
1.	Pancreatic pseudocyst
2.	Ascites caused by pancreatic duct disruption
3.	Pancreatic necrosis
II.	Pancreatic trauma
III.	Pancreatic carcinoma
Nonpancreatic Disorders	
I.	Renal insufficiency
II.	Salivary gland lesions
A.	Mumps
B.	Calculus
C.	Irradiation sialadenitis
D.	Maxillofacial surgery
III.	"Tumor" hyperamylasemia
A.	Carcinoma of the lung, esophagus, breast, or ovary
IV.	Macroamylasemia
V.	Burns
VI.	Diabetic ketoacidosis
VII.	Pregnancy
VIII.	Renal transplantation
IX.	Cerebral trauma
X.	Drugs: opiates
Other Abdominal Disorders	
I.	Biliary tract disease: cholecystitis, choledocholithiasis
II.	Intraabdominal disease
A.	Perforated or penetrating peptic ulcer
B.	Intestinal obstruction or inflammation
C.	Ruptured ectopic pregnancy
D.	Peritonitis
E.	Aortic aneurysm
F.	Postoperative hyperamylasemia

complications of acute and chronic pancreatitis. It is especially useful in the detection of pancreatic and peripancreatic acute fluid collections, fluid-containing lesions such as pseudocysts, walled-off necrosis, calcium deposits (see Chap. 371, Figs. 371-1, 371-2, and 371-4), and pancreatic neoplasms. Acute pancreatitis is characterized by (1) enlargement of the pancreatic outline, (2) distortion of the pancreatic contour, and/or (3) a pancreatic fluid that has a different attenuation coefficient than normal pancreas. Oral, water-soluble contrast agents are used to opacify the stomach and duodenum during CT scans; this strategy permits more precise delineation of various organs as well as mass lesions. Dynamic CT (using rapid IV administration of contrast) is useful in estimating the extent of pancreatic necrosis and in predicting morbidity and mortality. CT provides clear images much more rapidly and essentially negates artifact caused by patient movement. If acute pancreatitis is confirmed with serology and physical examination findings, CT scan in the first 3 days is not recommended to avoid overuse and minimize costs.

Endoscopic ultrasonography (EUS) produces high-resolution images of the pancreatic parenchyma and pancreatic duct with a transducer fixed to an endoscope that can be directed onto the surface of the pancreas through the stomach or duodenum. EUS and MRCP have largely replaced ERCP for diagnostic purposes in many centers. EUS allows one to obtain information about the pancreatic duct as well as the parenchyma and has few procedure-related complications associated with it, in contrast to the 5–10% of post-ERCP pancreatitis observed. EUS is also helpful in detecting common bile duct stones in acute

TABLE 370-3 ENDOSCOPIC ULTRASONOGRAPHIC CRITERIA FOR CHRONIC PANCREATITIS (TOTAL CRITERIA = 9)

Ductal	Parenchymal
Stones	Echogenic strands
Hyperechoic main duct margins	Echogenic foci
Main duct irregularity	Lobular contour
Main duct dilatation	Cyst
Visible side branches	

pancreatitis. Pancreatic masses can also be biopsied via EUS in cases with suspected pancreas cancer, and one can deliver nerve-blocking agents through EUS fine-needle injection in patients suffering from pancreatic pain from chronic pancreatitis or cancer. EUS has been studied as a diagnostic modality for chronic pancreatitis. Criteria for abnormalities on EUS in severe chronic pancreatic disease have been developed. There is general agreement that the presence of five or more of the nine criteria listed in Table 370-3 is highly predictive of chronic pancreatitis. Recent studies comparing EUS and ERCP to the secretin test in patients with unexplained abdominal pain suspected of having chronic pancreatitis show similar diagnostic accuracy in detecting early changes of chronic pancreatitis. The exact role of EUS versus CT, ERCP, or function testing in the early diagnosis of chronic pancreatitis has yet to be clearly defined.

Magnetic resonance imaging (MRI) and *magnetic resonance cholangiopancreatography* (MRCP) are now being used to view the bile ducts, pancreatic duct, and the pancreas parenchyma in both acute pancreatitis and chronic pancreatitis. For diagnostic imaging in chronic pancreatitis, non-breath-holding and three-dimensional turbo spin-echo techniques are being used to produce superb MRCP images. The main pancreatic duct and common bile duct can be seen well, but there is still a question as to whether changes can be detected consistently in the secondary ducts. The secondary ducts are not visualized in a normal pancreas. Secretin-enhanced MRCP is currently under investigation but is emerging as a method to better evaluate ductal changes. In anteroposterior imaging, T2 imaging of fluid collections can differentiate necrotic debris from fluid in suspected walled-off necrosis, and T1 imaging can diagnose hemorrhage in suspected pseudoaneurysm rupture.

Both EUS and MRCP have largely replaced ERCP in the diagnostic evaluation of pancreatic disease. As these techniques become more refined, especially with the administration of secretin, they may well be the diagnostic tests of choice to evaluate the pancreatic duct. ERCP is still needed for treatment of bile duct and pancreatic duct lesions. ERCP is primarily of therapeutic value after CT, EUS, or MRCP has detected abnormalities requiring invasive endoscopic treatment. ERCP can also be helpful at clarification of equivocal findings discovered with other imaging techniques (see Chap. 371, Fig. 371-1). Pancreatic carcinoma is characterized by stenosis or obstruction of either the pancreatic duct or the common bile duct; both ductal systems are often abnormal (double-duct sign). In chronic pancreatitis, ERCP abnormalities in the main pancreatic duct and side branches have been outlined by the Cambridge classification. The presence of ductal stenosis and irregularity can make it difficult to distinguish chronic pancreatitis from carcinoma. It is important to be aware that ERCP changes interpreted as indicating chronic pancreatitis actually may be due to the effects of aging on the pancreatic duct or sequelae of a recent attack of acute pancreatitis. Although aging may cause impressive ductal alterations, it does not affect the results of pancreatic function tests (i.e., the secretin test). Elevated serum amylase levels after ERCP have been reported in the majority of patients, and clinical pancreatitis in 5–10% of patients. Recent data suggest that pancreatic duct stenting and rectal indomethacin can decrease the incidence of ERCP-induced pancreatitis. ERCP should rarely be done for diagnostic purposes and should especially be avoided in high-risk patients.

PANCREATIC BIOPSY WITH RADIOLOGIC GUIDANCE Percutaneous aspiration biopsy or a trucut biopsy of a pancreatic mass often distinguishes a pancreatic inflammatory mass from a pancreatic neoplasm.

Pancreatic function tests (Table 370-1) can be divided into the following:

1. *Direct stimulation of the pancreas* by IV infusion of secretin followed by collection and measurement of duodenal contents

The secretin test, used to detect diffuse pancreatic disease, is based on the physiologic principle that the pancreatic secretory response is directly related to the functional mass of pancreatic tissue. In the standard assay, secretin is given IV in a dose of 0.2 $\mu\text{g}/\text{kg}$ of synthetic human secretin as a bolus. Normal values for the standard secretin test are (1) volume output >2 mL/kg per hour, (2) bicarbonate (HCO_3^-) concentration >80 mmol/L, and (3) HCO_3^- output >10 mmol/L in 1 h. The most reproducible measurement, giving the highest level of discrimination between normal subjects and patients with chronic pancreatic exocrine insufficiency, appears to be the maximal bicarbonate concentration. A cutoff point below 80 mmol/L is considered abnormal and suggestive of abnormal secretory function that is most commonly observed in early chronic pancreatitis.

There may be a dissociation between the results of the secretin test and other tests of absorptive function. For example, patients with chronic pancreatitis often have abnormally low outputs of HCO_3^- after secretin but have normal fecal fat excretion. Thus the secretin test measures the secretory capacity of ductular epithelium, whereas fecal fat excretion indirectly reflects intraluminal lipolytic activity. Steatorrhea does not occur until intraluminal levels of lipase are markedly reduced, underscoring the fact that only small amounts of enzymes are necessary for intraluminal digestive activities. It must be emphasized that an abnormal secretin test result suggests only that chronic pancreatic damage is present.

2. *Measurement of fecal pancreatic enzymes* such as elastase

Measurement of *intraluminal digestion products* (i.e., undigested muscle fibers, stool fat, and fecal nitrogen) is discussed in [Chap. 349](#). The amount of human elastase in stool reflects the pancreatic output of this proteolytic enzyme. Decreased elastase-1 activity (FE-1) in stool is an excellent test to detect severe pancreatic exocrine insufficiency (PEI) in patients with chronic pancreatitis and cystic fibrosis. FE-1 levels >200 $\mu\text{g}/\text{g}$ are normal; levels of 100–200 $\mu\text{g}/\text{g}$ are considered mild, and levels <100 $\mu\text{g}/\text{g}$ are severe for PEI. Although the test is simple and noninvasive, it can give false-positive results and has a low sensitivity. Fecal levels <50 $\mu\text{g}/\text{g}$ are definitive for PEI provided that the stool specimen is solid.

Tests useful in the diagnosis of exocrine pancreatic insufficiency and the differential diagnosis of malabsorption are also discussed in [Chaps. 349 and 371](#).

of secretin from the duodenal mucosa (S cells), which stimulates the secretion of water and electrolytes from pancreatic ductal cells. Release of cholecystokinin (CCK) from the duodenal and proximal jejunal mucosa (Ito cells) is largely triggered by long-chain fatty acids, essential amino acids (tryptophan, phenylalanine, valine, methionine), and gastric acid itself. CCK evokes an enzyme-rich secretion from acinar cells in the pancreas. The *parasympathetic nervous system* (via the vagus nerve) exerts significant control over pancreatic secretion. Secretion evoked by secretin and CCK depends on permissive roles of vagal afferent and efferent pathways. This is particularly true for enzyme secretion, whereas water and bicarbonate secretions are heavily dependent on the hormonal effects of secretin and to a lesser extent CCK. Also, vagal stimulation affects the release of vasoactive intestinal peptide (VIP), a secretin agonist. Pancreatic exocrine secretion is also influenced by inhibitory neuropeptides such as somatostatin, pancreatic polypeptide, peptide YY, neuropeptide Y, enkephalin, pancreastatin, calcitonin gene-related peptides, glucagon, and galanin. Although pancreatic polypeptide and peptide YY may act primarily on nerves outside the pancreas, somatostatin acts at multiple sites. Nitric oxide (NO) is also an important neurotransmitter.

WATER AND ELECTROLYTE SECRETION

Bicarbonate is the ion of primary physiologic importance within pancreatic secretion. The ductal cells secrete bicarbonate predominantly derived from plasma (93%) more than from intracellular metabolism (7%). Bicarbonate enters the duct lumen through the sodium bicarbonate cotransporter with depolarization caused by chloride efflux through the cystic fibrosis transmembrane conductance regulator (CFTR). Secretin and VIP bind at the basolateral surface and cause an increase in secondary messenger intracellular cyclic AMP, and act on the apical surface of the ductal cells opening the CFTR in promoting secretion. CCK, acting as a neuromodulator, markedly potentiates the stimulatory effects of secretin. Acetylcholine also plays an important role in ductal cell secretion. Intraluminal bicarbonate secreted from the ductal cells helps neutralize gastric acid and creates the appropriate pH for the activity of pancreatic enzymes and bile salts on ingested food.

ENZYME SECRETION

The acinar cell is highly compartmentalized and is concerned with the secretion of pancreatic enzymes. Proteins synthesized by the rough endoplasmic reticulum are processed in the Golgi and then targeted to the appropriate site, whether that be zymogen granules, lysosomes, or other cell compartments. The zymogen granules migrate to the apical region of the acinar cell awaiting the appropriate neural or hormonal stimulatory response. The pancreas secretes amylolytic, lipolytic, and proteolytic enzymes into the duct lumen. *Amylolytic enzymes*, such as amylase, hydrolyze starch to oligosaccharides and to the disaccharide maltose. The *lipolytic enzymes* include lipase, phospholipase A_2 , and cholesterol esterase. Bile salts inhibit lipase in isolation, but colipase, another constituent of pancreatic secretion, binds to lipase and prevents this inhibition. Bile salts activate phospholipase A and cholesterol esterase. *Proteolytic enzymes* include endopeptidases (trypsin, chymotrypsin), which act on internal peptide bonds of proteins and polypeptides; exopeptidases (carboxypeptidases, aminopeptidases), which act on the free carboxyl- and amino-terminal ends of peptides, respectively; and elastase. The proteolytic enzymes are secreted as inactive zymogen precursors. Ribonucleases (deoxyribonucleases, ribonuclease) are also secreted. *Enterokinase*, an enzyme found in the duodenal mucosa, cleaves the lysine-isoleucine bond of trypsinogen to form trypsin. Trypsin then activates the other proteolytic zymogens and phospholipase A_2 in a cascade phenomenon. All pancreatic enzymes have pH optima in the alkaline range. The nervous system initiates pancreatic enzyme secretion. The neurologic stimulation is cholinergic, involving extrinsic innervation by the vagus nerve and subsequent innervation by intrapancreatic cholinergic nerves. The stimulatory neurotransmitters are acetylcholine and gastrin-releasing peptides. These neurotransmitters activate calcium-dependent secondary messenger systems, resulting in the release of zymogens into the pancreas duct. VIP is present in intrapancreatic nerves and potentiates the effect of acetylcholine. In

371 Acute and Chronic Pancreatitis

Darwin L. Conwell, Peter Banks, Norton J. Greenberger

BIOCHEMISTRY AND PHYSIOLOGY OF PANCREATIC EXOCRINE SECRETION

GENERAL CONSIDERATIONS

The pancreas secretes 1500–3000 mL of isosmotic alkaline (pH >8) fluid per day containing about 20 enzymes. The pancreatic secretions provide the enzymes and bicarbonate needed to affect the major digestive activity of the gastrointestinal tract and provide an optimal pH for the function of these enzymes.

REGULATION OF PANCREATIC SECRETION

The exocrine pancreas is influenced by intimately interacting hormonal and neural systems. *Gastric acid* is the stimulus for the release

contrast to other species, there are no CCK receptors on acinar cells in humans. CCK in physiologic concentrations stimulates pancreatic secretion by stimulating afferent vagal and intrapancreatic nerves.

AUTOPROTECTION OF THE PANCREAS

Autodigestion of the pancreas is prevented by (1) the packaging of pancreatic proteases in precursor (proenzyme) form, (2) intracellular calcium homeostasis (low intracellular calcium in the cytosol of the acinar cell promotes the destruction of spontaneously activated trypsin), (3) acid-base balance, and (4) the synthesis of protective protease inhibitors (pancreatic secretory trypsin inhibitor [PSTI] or SPINK1), which can bind and inactivate about 20% of intracellular trypsin activity. Chymotrypsin C can also lyse and inactivate trypsin. These protease inhibitors are found in the acinar cell, the pancreatic secretions, and the α_1 - and α_2 -globulin fractions of plasma. Loss of any of these four protective mechanisms leads to premature enzyme activation, autodigestion, and acute pancreatitis.

ENTEROPANCREATIC AXIS AND FEEDBACK INHIBITION

Pancreatic enzyme secretion is controlled, at least in part, by a negative feedback mechanism induced by the presence of active serine proteases in the duodenum. To illustrate, perfusion of the duodenal lumen with phenylalanine (stimulates early digestion) causes a prompt increase in plasma CCK levels as well as increased secretion of chymotrypsin and other pancreatic enzymes. However, simultaneous perfusion with trypsin (stimulates late digestion) blunts both responses. Conversely, perfusion of the duodenal lumen with protease inhibitors actually leads to enzyme hypersecretion. The available evidence supports the concept that the duodenum contains a peptide called *CCK-releasing factor* (CCK-RF) that is involved in stimulating CCK release. It appears that serine proteases inhibit pancreatic secretion by inactivating a CCK-releasing peptide in the lumen of the small intestine. Thus, the integrative result of both bicarbonate and enzyme secretion depends on a feedback process for both bicarbonate and pancreatic enzymes. Acidification of the duodenum releases secretin, which stimulates vagal and other neural pathways to activate pancreatic duct cells, which secrete bicarbonate. This bicarbonate then neutralizes the duodenal acid, and the feedback loop is completed. Dietary proteins bind proteases, thereby leading to an increase in free CCK-RF. CCK is then released into the blood in physiologic concentrations, acting primarily through the neural pathways (vagal-vagal). This leads to acetylcholine-mediated pancreatic enzyme secretion. Proteases continue to be secreted from the pancreas until the protein within the duodenum is digested. At this point, pancreatic protease secretion is reduced to basic levels, thus completing this step in the feedback process.

ACUTE PANCREATITIS

GENERAL CONSIDERATIONS

Recent U.S. estimates from the National Inpatient Sample report that acute pancreatitis is the most common inpatient principal gastrointestinal diagnosis. The incidence of acute pancreatitis also varies in different countries and depends on cause (e.g., alcohol, gallstones, metabolic factors, drugs [Table 371-1]). The annual incidence ranges from 13–45/100,000 persons. Acute pancreatitis results in >250,000 hospitalizations per year. The median length of hospital stay is 4 days, with a median hospital cost of \$6,096 and a mortality of 1%. The estimated cost annually approaches \$2.6 billion. Hospitalization rates increase with age, are 88% higher among blacks, and are higher among males than females. The age-adjusted rate of hospital discharges with an acute pancreatitis diagnosis increased 62% between 1988 and 2004. From 2000 to 2009, the rate increased 30%. Thus, acute pancreatitis is increasing and is a significant burden on health care costs and resource utilization.

ETIOLOGY AND PATHOGENESIS

There are many causes of acute pancreatitis (Table 371-1), but the mechanisms by which these conditions trigger pancreatic inflammation have not been fully elucidated. Gallstones continue to be the leading cause of acute pancreatitis in most series (30–60%). The risk

TABLE 371-1 CAUSES OF ACUTE PANCREATITIS

Common Causes
Gallstones (including microlithiasis)
Alcohol (acute and chronic alcoholism)
Hypertriglyceridemia
Endoscopic retrograde cholangiopancreatography (ERCP), especially after biliary manometry
Drugs (azathioprine, 6-mercaptopurine, sulfonamides, estrogens, tetracycline, valproic acid, anti-HIV medications, 5-aminosalicylic acid [5-ASA])
Trauma (especially blunt abdominal trauma)
Postoperative (abdominal and nonabdominal operations)
Uncommon Causes
Vascular causes and vasculitis (ischemic-hypoperfusion states after cardiac surgery)
Connective tissue disorders and thrombotic thrombocytopenic purpura (TTP)
Cancer of the pancreas
Hypercalcemia
Periampullary diverticulum
Pancreas divisum
Hereditary pancreatitis
Cystic fibrosis
Renal failure
Infections (mumps, coxsackievirus, cytomegalovirus, echovirus, parasites)
Autoimmune (e.g., type 1 and type 2)
Causes to Consider in Patients with Recurrent Bouts of Acute Pancreatitis Without an Obvious Etiology
Occult disease of the biliary tree or pancreatic ducts, especially microlithiasis, biliary sludge
Drugs
Alcohol abuse
Metabolic: Hypertriglyceridemia, hypercalcemia
Anatomic: Pancreas divisum
Pancreatic cancer
Intraductal papillary mucinous neoplasm (IPMN)
Hereditary pancreatitis
Cystic fibrosis
Autoimmune
Idiopathic

of acute pancreatitis in patients with at least one gallstone <5 mm in diameter is fourfold greater than that in patients with larger stones. Alcohol is the second most common cause, responsible for 15–30% of cases in the United States. The incidence of pancreatitis in alcoholics is surprisingly low (5/100,000), indicating that in addition to the amount of alcohol ingested, other factors affect a person's susceptibility to pancreatic injury such as cigarette smoking. Acute pancreatitis occurs in 5–10% of patients following endoscopic retrograde cholangiopancreatography (ERCP). Use of a prophylactic pancreatic duct stent and rectal nonsteroidal anti-inflammatory drugs (NSAIDs) has been shown to reduce pancreatitis after ERCP. Risk factors for post-ERCP pancreatitis include minor papilla sphincterotomy, sphincter of Oddi dysfunction, prior history of post-ERCP pancreatitis, age <60 years, >2 contrast injections into the pancreatic duct, and endoscopic trainee involvement.

Hypertriglyceridemia is the cause of acute pancreatitis in 1.3–3.8% of cases; serum triglyceride levels are usually >11.3 mmol/L (>1000 mg/dL). Most patients with hypertriglyceridemia, when subsequently examined, show evidence of an underlying derangement in lipid metabolism, probably unrelated to pancreatitis. Such patients are prone to recurrent episodes of pancreatitis. Any factor (e.g., drugs or alcohol) that causes an abrupt increase in serum triglycerides can precipitate a bout of acute pancreatitis. Patients with a deficiency of apolipoprotein CII have an increased incidence of pancreatitis; apolipoprotein CII

2092 activates lipoprotein lipase, which is important in clearing chylomicrons from the bloodstream. Patients with diabetes mellitus who have developed ketoacidosis and patients who are on certain medications such as oral contraceptives may also develop high triglyceride levels. Approximately 0.1–2% of cases of acute pancreatitis are drug related. Drugs cause pancreatitis either by a hypersensitivity reaction or by the generation of a toxic metabolite, although in some cases, it is not clear which of these mechanisms is operative (Table 371-1).

Pathologically, acute pancreatitis varies from *interstitial pancreatitis* (pancreas blood supply maintained), which is generally self-limited to *necrotizing pancreatitis* (pancreas blood supply interrupted), in which the extent of necrosis may correlate with the severity of the attack and its systemic complications. Autodigestion is a currently accepted pathogenic theory; according to this theory, pancreatitis results when proteolytic enzymes (e.g., trypsinogen, chymotrypsinogen, proelastase, and lipolytic enzymes such as phospholipase A₂) are activated in the pancreas acinar cell rather than in the intestinal lumen. A number of factors (e.g., endotoxins, exotoxins, viral infections, ischemia, oxidative stress, lysosomal calcium, and direct trauma) are believed to facilitate premature activation of trypsin. Activated proteolytic enzymes, especially trypsin, not only digest pancreatic and peripancreatic tissues but also can activate other enzymes, such as elastase and phospholipase A₂. Spontaneous activation of trypsin also can occur.

ACTIVATION OF PANCREATIC ENZYMES IN THE PATHOGENESIS OF ACUTE PANCREATITIS

Several recent studies have suggested that pancreatitis is a disease that evolves in three phases. The *initial phase* is characterized by intrapancreatic digestive enzyme activation and acinar cell injury. Trypsin activation appears to be mediated by lysosomal hydrolases such as cathepsin B that become colocalized with digestive enzymes in intracellular organelles; it is currently believed that acinar cell injury is the consequence of trypsin activation. The *second phase* of pancreatitis involves the activation, chemoattraction, and sequestration of leukocytes and macrophages in the pancreas, resulting in an enhanced intrapancreatic inflammatory reaction. Neutrophil depletion induced by prior administration of an antineutrophil serum has been shown to reduce the severity of experimentally induced pancreatitis. There is also evidence to support the concept that neutrophils can activate trypsinogen. Thus, intrapancreatic acinar cell activation of trypsinogen could be a two-step process (i.e., an early neutrophil-independent and a later neutrophil-dependent phase). The *third phase* of pancreatitis is due to the effects of activated proteolytic enzymes and cytokines, released by the inflamed pancreas, on distant organs. Activated proteolytic enzymes, especially trypsin, not only digest pancreatic and peripancreatic tissues but also activate other enzymes such as elastase and phospholipase A₂. The active enzymes and cytokines then digest cellular membranes and cause proteolysis, edema, interstitial hemorrhage, vascular damage, coagulation necrosis, fat necrosis, and parenchymal cell necrosis. Cellular injury and death result in the liberation of bradykinin peptides, vasoactive substances, and histamine that can produce vasodilation, increased vascular permeability, and edema with profound effects on many organs. The systemic inflammatory response syndrome (SIRS) and acute respiratory distress syndrome (ARDS), as well as multiorgan failure, may occur as a result of this cascade of local and distant effects.

A number of genetic factors can increase the susceptibility and/or modify the severity of pancreatic injury in acute pancreatitis, recurrent pancreatitis, and chronic pancreatitis. All of the major genetic susceptibility factors center on the control of trypsin activity within the pancreatic acinar cell, in part because they were identified as candidate genes linked to intrapancreatic trypsin control. Five genetic variants have been identified as being associated with susceptibility to pancreatitis. The genes that have been identified include (1) cationic trypsinogen gene (*PRSS1*), (2) pancreatic secretory trypsin inhibitor (*SPINK1*), (3) the cystic fibrosis transmembrane conductance regulator gene (*CFTR*), (4) the chymotrypsin C gene (*CTRC*), and (5) the calcium-sensing receptor (*CASR*). Investigations of other genetic variants are currently under way, and new genes will be added to this list

in the future. Multiple medical, ethical, and psychological issues arise when these genes are discovered, and referral to genetic counselors is recommended.

APPROACH TO THE PATIENT:

Abdominal Pain

Abdominal pain is the major symptom of acute pancreatitis. Pain may vary from a mild discomfort to severe, constant, and incapacitating distress. Characteristically, the pain, which is steady and boring in character, is located in the epigastrium and periumbilical region, and may radiate to the back, chest, flanks, and lower abdomen. Nausea, vomiting, and abdominal distention due to gastric and intestinal hypomotility and chemical peritonitis are also frequent complaints.

Physical examination frequently reveals a distressed and anxious patient. Low-grade fever, tachycardia, and hypotension are fairly common. Shock is not unusual and may result from (1) hypovolemia secondary to exudation of blood and plasma proteins into the retroperitoneal space; (2) increased formation and release of kinin peptides, which cause vasodilation and increased vascular permeability; and (3) systemic effects of proteolytic and lipolytic enzymes released into the circulation. Jaundice occurs infrequently; when present, it usually is due to edema of the head of the pancreas with compression of the intrapancreatic portion of the common bile duct or passage of a biliary stone or sludge. Erythematous skin nodules due to subcutaneous fat necrosis may rarely occur. In 10–20% of patients, there are pulmonary findings, including basilar rales, atelectasis, and pleural effusion, the latter most frequently left sided. Abdominal tenderness and muscle rigidity are present to a variable degree, but compared with the intense pain, these signs may be less impressive. Bowel sounds are usually diminished or absent. An enlarged pancreas from acute fluid collection, walled off necrosis, or a pseudocyst may be palpable in the upper abdomen later in the course of the disease (i.e., 4–6 weeks). A faint blue discoloration around the umbilicus (Cullen's sign) may occur as the result of hemoperitoneum, and a blue-red-purple or green-brown discoloration of the flanks (Turner's sign) reflects tissue catabolism of hemoglobin from severe necrotizing pancreatitis with hemorrhage.

LABORATORY DATA

Serum amylase and lipase values threefold or more above normal virtually clinch the diagnosis if gut perforation, ischemia, and infarction are excluded. Serum lipase is the preferred test. However, it should be noted that there is no correlation between the severity of pancreatitis and the degree of serum lipase and amylase elevations. After 3–7 days, even with continuing evidence of pancreatitis, total serum amylase values tend to return toward normal. However, pancreatic isoamylase and lipase levels may remain elevated for 7–14 days. It should be recognized that amylase elevations in serum and urine occur in many conditions other than pancreatitis (see Chap. 370, Table 370-2). Importantly, patients with *acidemia* (arterial pH ≤ 7.32) may have spurious elevations in serum amylase. This finding explains why patients with diabetic ketoacidosis may have marked elevations in serum amylase without any other evidence of acute pancreatitis. Serum lipase activity increases in parallel with amylase activity and is more specific than amylase. A serum lipase measurement can be instrumental in differentiating a pancreatic or nonpancreatic cause for hyperamylasemia. *Leukocytosis* (15,000–20,000 leukocytes/ μ L) occurs frequently. Patients with more severe disease may show hemoconcentration with hematocrit values $>44\%$ and/or prerenal azotemia with a blood urea nitrogen (BUN) level >22 mg/dL resulting from loss of plasma into the retroperitoneal space and peritoneal cavity.

Hemoconcentration may be the harbinger of more severe disease (i.e., pancreatic necrosis), whereas azotemia is a significant risk factor for mortality. *Hyperglycemia* is common and is due to multiple factors, including decreased insulin release, increased glucagon release, and

TABLE 371-2 REVISED ATLANTA DEFINITIONS OF MORPHOLOGIC FEATURES OF ACUTE PANCREATITIS

Morphologic Feature	Definition	Computed Tomography Criteria
Interstitial pancreatitis	Acute inflammation of the pancreatic parenchyma and peripancreatic tissues, but without recognizable tissue necrosis	Pancreatic parenchyma enhancement by IV contrast agent No findings of peripancreatic necrosis
Necrotizing pancreatitis	Inflammation associated with pancreatic parenchymal necrosis and/or peripancreatic necrosis	Lack of pancreatic parenchymal enhancement by IV contrast agent and/or presence of findings of peripancreatic necrosis (see below—ANC and WON)
Acute pancreatic fluid collection	Peripancreatic fluid associated with interstitial edematous pancreatitis with no associated peripancreatic necrosis. This term applies only to areas of peripancreatic fluid seen within the first 4 weeks after onset of interstitial edematous pancreatitis and without the features of a pseudocyst.	Occurs in the setting of interstitial edematous pancreatitis Homogeneous collection with fluid density Confined by normal peripancreatic fascial planes No definable wall encapsulating the collection Adjacent to pancreas (no intrapancreatic extension)
Pancreatic pseudocyst	An encapsulated collection of fluid with a well-defined inflammatory wall usually outside the pancreas with minimal or no necrosis. This entity usually occurs >4 weeks after onset of interstitial edematous pancreatitis.	Well circumscribed, usually round or oval Homogeneous fluid density No nonliquid component Well-defined wall; that is, completely encapsulated Maturation usually requires >4 weeks after onset of acute pancreatitis; occurs after interstitial edematous pancreatitis
Acute necrotic collection (ANC)	A collection containing variable amounts of both fluid and necrosis associated with necrotizing pancreatitis; the necrosis can involve the pancreatic parenchyma and/or the peripancreatic tissues.	Occurs only in the setting of acute necrotizing pancreatitis Heterogeneous and nonliquid density of varying degrees in different locations (some appear homogeneous early in their course) No definable wall encapsulating the collection Location—intrapancreatic and/or extrapancreatic
Walled-off necrosis (WON)	A mature, encapsulated collection of pancreatic and/or peripancreatic necrosis that has developed a well-defined inflammatory wall. WON usually occurs >4 weeks after onset of necrotizing pancreatitis.	Heterogeneous with liquid and nonliquid density with varying degrees of loculations (some may appear homogeneous) Well-defined wall; that is, completely encapsulated Location—intrapancreatic and/or extrapancreatic Maturation usually requires 4 weeks after onset of acute necrotizing pancreatitis

Source: Modified from P Banks et al: Gut 62:102, 2013.

an increased output of adrenal glucocorticoids and catecholamines. *Hypocalcemia* occurs in ~25% of patients, and its pathogenesis is incompletely understood. Although earlier studies suggested that the response of the parathyroid gland to a decrease in serum calcium is impaired, subsequent observations have failed to confirm this phenomenon. Intraperitoneal saponification of calcium by fatty acids in areas of fat necrosis occurs occasionally, with large amounts (up to 6.0 g) dissolved or suspended in ascitic fluid. Such “soap formation” may also be significant in patients with pancreatitis, mild hypocalcemia, and

little or no obvious ascites. *Hyperbilirubinemia* (serum bilirubin >68 mmol/L or >4.0 mg/dL) occurs in ~10% of patients. However, jaundice is transient, and serum bilirubin levels return to normal in 4–7 days. Serum alkaline phosphatase and aspartate aminotransferase levels are also transiently elevated, and they parallel serum bilirubin values and may point to gallbladder-related disease or inflammation in the pancreatic head. *Hypertriglyceridemia* occurs in 5–10% of patients, and serum amylase levels in these individuals are often spuriously normal (Chap. 370). Approximately 5–10% of patients have *hypoxemia* (arterial PO₂ ≤60 mmHg), which may herald the onset of ARDS. Finally, the electrocardiogram is occasionally abnormal in acute pancreatitis with ST-segment and T-wave abnormalities simulating myocardial ischemia.

An abdominal ultrasound is recommended in the emergency ward as the initial diagnostic imaging modality and is most useful to evaluate for gallstone disease and the pancreatic head.

The revised Atlanta criteria have clearly outlined the morphologic features of acute pancreatitis on computed tomography (CT) scan as follows: (1) interstitial pancreatitis, (2) necrotizing pancreatitis, (3) acute pancreatic fluid collection, (4) pancreatic pseudocyst, (5) acute necrotic collection (ANC), and (6) walled-off pancreatic necrosis (WON) (Table 371-2 and Fig. 371-1). **Radiologic studies useful in the diagnosis of acute pancreatitis are discussed in Chap. 370 and listed in Table 370-1.**

DIAGNOSIS

Any severe acute pain in the abdomen or back should suggest the possibility of acute pancreatitis. The diagnosis is established by two of the following three criteria: (1) typical abdominal pain in the epigastrium that may radiate to the back, (2) threefold or greater elevation in serum lipase and/or amylase, and (3) confirmatory findings of acute pancreatitis on cross-sectional abdominal imaging. Patients also have associated nausea, emesis, fever, tachycardia, and abnormal findings on abdominal examination. Laboratory studies may reveal leukocytosis, hypocalcemia, and hyperglycemia. Although not required for diagnosis, markers of severity may include hemoconcentration (hematocrit >44%), admission azotemia (BUN >22 mg/dL), SIRS, and signs of organ failure (Table 371-3).

The *differential diagnosis* should include the following disorders: (1) perforated viscus, especially peptic ulcer; (2) acute cholecystitis and biliary colic; (3) acute intestinal obstruction; (4) mesenteric vascular occlusion; (5) renal colic; (6) inferior myocardial infarction; (7) dissecting aortic aneurysm; (8) connective tissue disorders with vasculitis; (9) pneumonia; and (10) diabetic ketoacidosis. It may be difficult to differentiate acute cholecystitis from acute pancreatitis, because an elevated serum amylase may be found in both disorders. Pain of biliary tract origin is more right sided or epigastric than periumbilical or left upper quadrant and can be more severe; ileus is usually absent. Ultrasound is helpful in establishing the diagnosis of cholelithiasis and cholecystitis. Intestinal obstruction due to mechanical factors can be differentiated from pancreatitis by the history of crescendo-decrescendo pain, findings on abdominal examination, and CT of the abdomen showing changes characteristic of mechanical obstruction. Acute mesenteric vascular occlusion is usually suspected in elderly debilitated patients with brisk leukocytosis, abdominal distention, and bloody diarrhea, confirmed by CT or magnetic resonance angiography. Vasculitides secondary to systemic lupus erythematosus and polyarteritis nodosa may be confused with pancreatitis, especially because pancreatitis may develop as a complication of these diseases. Diabetic ketoacidosis is often accompanied by abdominal pain and elevated total serum amylase levels, thus closely mimicking acute pancreatitis. However, the serum lipase level is not elevated in diabetic ketoacidosis.

CLINICAL COURSE, DEFINITIONS, AND CLASSIFICATIONS

The Revised Atlanta Classification (1) defines phases of acute pancreatitis, (2) defines severity of acute pancreatitis, and (3) clarifies imaging definitions as outlined below.

Phases of Acute Pancreatitis Two phases of acute pancreatitis have been defined, early (<2 weeks) and late (>2 weeks), which primarily



FIGURE 371-1 Acute pancreatitis: computed tomography (CT) evolution. **A.** Contrast-enhanced CT scan of the abdomen performed on admission for a patient with clinical and biochemical parameters suggestive of acute pancreatitis. Note the abnormal enhancement of the pancreatic parenchyma (*arrow*) suggestive of interstitial pancreatitis. **B.** Contrast-enhanced CT scan of the abdomen performed on the same patient 6 days later for persistent fever and systemic inflammatory response syndrome. The pancreas now demonstrates significant areas of nonenhancement consistent with development of necrosis, particularly in the body and neck region (*arrow*). Note that an early CT scan obtained within the first 48 h of hospitalization may underestimate or miss necrosis. **C.** Contrast-enhanced CT scan of the abdomen performed on the same patient 2 months after the initial episode of acute pancreatitis. CT now demonstrates evidence of a fluid collection consistent with walled-off pancreatic necrosis (*arrow*). (Courtesy of Dr. KJ Mortelet, Brigham and Women's Hospital; with permission.)

describes the hospital course of the disease. In the *early phase* of acute pancreatitis, which lasts 1–2 weeks, severity is defined by clinical parameters rather than morphologic findings. Most patients exhibit SIRS, and if this persists, patients are predisposed to organ failure. Three organ systems should be assessed to define organ failure: respiratory, cardiovascular, and renal. Organ failure is defined as a score of 2 or more for one of these three organ systems using the modified Marshall scoring system. Persistent organ failure (>48 h) is the most important clinical finding in regard to severity of the acute pancreatitis

episode. Organ failure that affects more than one organ is considered multisystem organ failure. CT imaging is usually not needed or recommended during the first 48 h of admission in acute pancreatitis.

The *late phase* is characterized by a protracted course of illness and may require imaging to evaluate for local complications. The important clinical parameter of severity, as in the early phase, is persistent organ failure. These patients may require supportive measures such as renal dialysis, ventilator support, or need for supplemental nutrition via the nasojejunal or parenteral route. The radiographic feature of greatest importance to recognize in this phase is the development of necrotizing pancreatitis on CT imaging. Necrosis generally prolongs hospitalization and, if infected, may require operative, endoscopic, or percutaneous intervention.

TABLE 371-3 SEVERE ACUTE PANCREATITIS

Risk Factors for Severity

- Age >60 years
- Obesity, BMI >30
- Comorbid disease (Charlson Comorbidity Index)

Markers of Severity at Admission or Within 24 h

- SIRS—defined by presence of 2 or more criteria:
- Core temperature <36° or >38°C
- Heart rate >90 beats/min
- Respirations >20/min or Pco₂ <32 mmHg
- White blood cell count >12,000/μL, <4000/μL, or 10% bands
- APACHE II
- Hemoconcentration (hematocrit >44%)
- Admission BUN (>22 mg/dL)
- BISAP Score
 - (B) BUN >25 mg/dL
 - (I) Impaired mental status
 - (S) SIRS: ≥2 of 4 present
 - (A) Age >60 years
 - (P) Pleural effusion
- Organ failure (Modified Marshall Score)
- Cardiovascular: systolic BP <90 mmHg, heart rate >130 beats/min
- Pulmonary: Pao₂ <60 mmHg
- Renal: serum creatinine >2.0 mg%

Markers of Severity During Hospitalization

- Persistent organ failure
- Pancreatic necrosis

Abbreviations: APACHE II, Acute Physiology and Chronic Health Evaluation II; BMI, body mass index; BISAP, Bedside Index of Severity in Acute Pancreatitis; BP, blood pressure; BUN, blood urea nitrogen; SIRS, systemic inflammatory response syndrome.

Severity of Acute Pancreatitis Three severity classifications have also been defined: mild, moderately severe, and severe. *Mild acute pancreatitis* is without local complications or organ failure. Most patients with interstitial acute pancreatitis have mild pancreatitis. In mild acute pancreatitis, the disease is self-limited and subsides spontaneously, usually within 3–7 days after treatment is instituted. Oral intake can be resumed if the patient is hungry, has normal bowel function, and is without nausea and vomiting. Typically, a clear or full liquid diet has been recommended for the initial meal; however, a low-fat solid diet is a reasonable choice following recovery from mild acute pancreatitis.

Moderately severe acute pancreatitis is characterized by transient organ failure (resolves in <48 h) or local or systemic complications in the absence of persistent organ failure. These patients may or may not have necrosis, but may develop a local complication such as a fluid collection that requires a prolonged hospitalization greater than 1 week.

Severe acute pancreatitis is characterized by persistent organ failure (>48 h). Organ failure can be single or multiple. A CT scan or magnetic resonance imaging (MRI) should be obtained to assess for necrosis and/or complications. If a local complication is encountered, management is dictated by clinical symptoms, evidence of infection, maturity of fluid collection, and clinical stability of the patient. Prophylactic antibiotics are not recommended.

Imaging in Acute Pancreatitis Two types of pancreatitis are recognized on imaging as *interstitial* or *necrotizing* based on pancreatic perfusion. CT imaging is best evaluated 3–5 days into hospitalization when patients are not responding to supportive care to look for local complications such as necrosis. Recent studies report the overutilization of CT imaging in acute pancreatitis and its inability to be better than clinical judgment in the early days of acute pancreatitis management. The revised criteria also outline the terminology for local

complications and fluid collections along with a CT imaging template to guide reporting of findings. Local morphologic features are summarized in Table 371-1. *Interstitial pancreatitis* occurs in 90–95% of admissions for acute pancreatitis and is characterized by diffuse gland enlargement, homogenous contrast enhancement, and mild inflammatory changes or peripancreatic stranding. Symptoms generally resolve with a week of hospitalization. *Necrotizing pancreatitis* occurs in 5–10% of acute pancreatitis admissions and does not evolve until several days of hospitalization. It is characterized by lack of pancreatic parenchymal enhancement by intravenous contrast agent and/or presence of findings of peripancreatic necrosis. According to the revised Atlanta criteria, the natural history of pancreatic and peripancreatic necrosis is variable because it may remain solid or liquefy, remain sterile or become infected, and persist or disappear over time. CT identification of local complications, particularly necrosis, is critical in patients who are not responding to therapy because patients with infected and sterile necrosis are at greatest risk of mortality (Figs. 371-1B, 371-2, and 371-3). The median prevalence of organ failure is 54% in necrotizing pancreatitis. The prevalence of organ failure is perhaps slightly higher in infected versus sterile necrosis. With single-organ system failure, the mortality is 3–10% but increases to 47% with multisystem organ failure.

ACUTE PANCREATITIS MANAGEMENT

We will briefly describe the management of patients with acute pancreatitis from the time of diagnosis in the emergency ward to ongoing hospital admission and, finally, to time of discharge, highlighting salient features based on severity and complications. It is important to note that 85–90% of cases of acute pancreatitis are self-limited and subside spontaneously, usually within 3–7 days after initiation of treatment, and do not exhibit organ failure or local complications.

The management of acute pancreatitis begins in the emergency ward. After a diagnosis has been

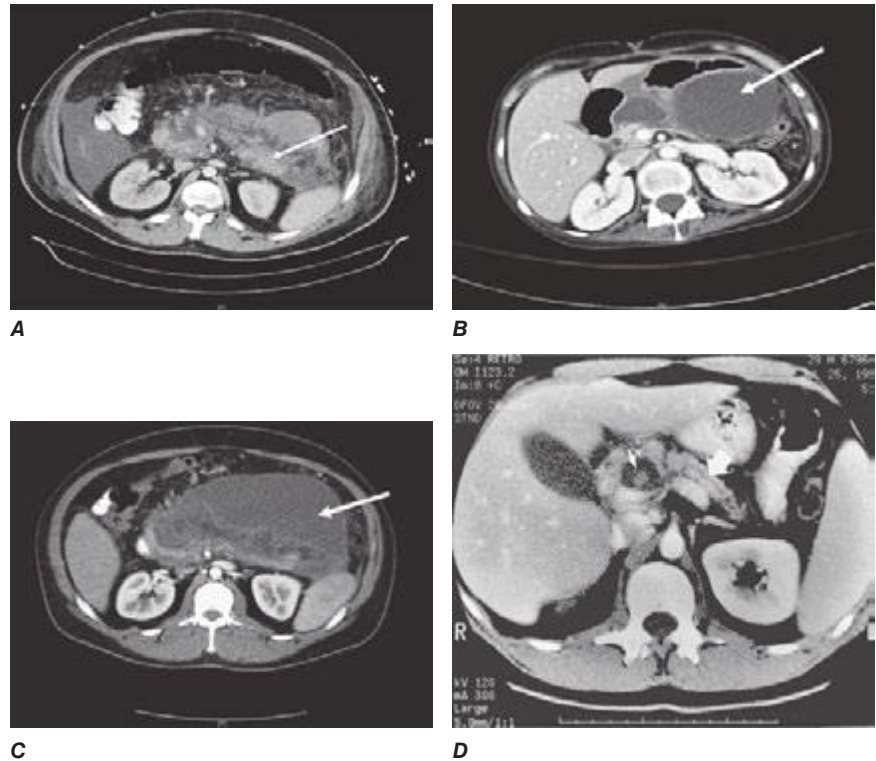


FIGURE 371-2 **A.** Acute necrotizing pancreatitis: computed tomography (CT) scan. Contrast-enhanced CT scan showing acute pancreatitis with necrosis. Arrow shows partially enhancing body/tail of pancreas surrounded by fluid with decreased enhancement in the neck/body of the pancreas. **B.** Acute fluid collection: CT scan. Contrast-enhanced CT scan showing fluid collection in the retroperitoneum (arrow) compressing the air-filled stomach arising from the pancreas in a patient with asparaginase-induced acute necrotizing pancreatitis. **C.** Walled-off pancreatic necrosis: CT scan. CT scan showing marked walled-off necrosis of the pancreas and peripancreatic area (arrow) in a patient with necrotizing pancreatitis. Addendum: In past years, both of these CT findings (Figs. 371-2B and 371-2C) would have been misinterpreted as pseudocysts. **D.** Spiral CT showing a pseudocyst (small arrow) with a pseudoaneurysm (light area in pseudocyst). Note the demonstration of the main pancreatic duct (big arrow), even though this duct is minimally dilated by endoscopic retrograde cholangiopancreatography. (A, B, C, courtesy of Dr. KJ Morteale, Brigham and Women's Hospital; D, courtesy of Dr. PR Ros, Brigham and Women's Hospital; with permission.)



FIGURE 371-3 **A.** Pancreaticopleural fistula: pancreatic duct leak on endoscopic retrograde cholangiopancreatography. Pancreatic duct leak (arrow) demonstrated at the time of retrograde pancreatogram in a patient with acute exacerbation of alcohol-induced acute or chronic pancreatitis. **B.** Pancreaticopleural fistula: computed tomography (CT) scan. Contrast-enhanced CT scan (coronal view) with arrows showing fistula tract from pancreatic duct disruption in the pancreatic pleural fistula. **C.** Pancreaticopleural fistula: chest x-ray. Large pleural effusion in the left hemithorax from a disrupted pancreatic duct. Analysis of pleural fluid revealed elevated amylase concentration. (Courtesy of Dr. KJ Morteale, Brigham and Women's Hospital; with permission.)

confirmed, aggressive fluid resuscitation is initiated, intravenous analgesics are administered, severity is assessed, and a search for etiologies that may impact acute care is begun. Patients who do not respond to aggressive fluid resuscitation in the emergency ward should be considered for admission to a step-down or intensive care unit for aggressive fluid resuscitation, hemodynamic monitoring, and management of necrosis or organ failure.

Fluid Resuscitation and Monitoring Response to Therapy The most important treatment intervention for acute pancreatitis is safe, aggressive intravenous fluid resuscitation. The patient is made NPO to rest the pancreas and is given intravenous narcotic analgesics to control abdominal pain and supplemental oxygen (2 L) via nasal cannula.

Intravenous fluids of lactated Ringer's or normal saline are initially bolused at 15–20 cc/kg (1050–1400 mL), followed by 3 mg/kg per hour (200–250 mL/h), to maintain urine output >0.5 cc/kg per hour. Serial bedside evaluations are required every 6–8 h to assess vital signs, oxygen saturation, and change in physical examination. Lactated Ringer's solution has been shown to decrease systemic inflammation and may be a better crystalloid than normal saline. A *targeted resuscitation strategy* with measurement of hematocrit and BUN every 8–12 h is recommended to ensure adequacy of fluid resuscitation and monitor response to therapy, noting less aggressive resuscitation strategy may be needed in milder forms of pancreatitis. A rising BUN during hospitalization is not only associated with inadequate hydration but also higher in-hospital mortality.

A decrease in hematocrit and BUN during the first 12–24 h is strong evidence that sufficient fluids are being administered. Serial measurements and bedside assessment for fluid overload are continued, and fluid rates are maintained at the current rate. Adjustments in fluid resuscitation may be required in patients with cardiac, pulmonary, or renal disease. A rise in hematocrit or BUN during serial measurement should be treated with a repeat volume challenge with a 2-L crystalloid bolus followed by increasing the fluid rate by 1.5 mg/kg per hour. If the BUN or hematocrit fails to respond (i.e., remains elevated or does not decrease) to this bolus challenge and increase in fluid rate, consideration of transfer to an intensive care unit is strongly recommended for hemodynamic monitoring.

Assessment of Severity and Hospital Triage Severity of acute pancreatitis should be determined in the emergency ward to assist in patient triage to a regular hospital ward or step-down unit or direct admission to an intensive care unit. The Bedside Index of Severity in Acute Pancreatitis (BISAP) incorporates five clinical and laboratory parameters obtained within the first 24 h of hospitalization (Table 371-3)—BUN >25 mg/dL, impaired mental status (Glasgow coma score <15), SIRS, age >60 years, and pleural effusion on radiography—that can be useful in assessing severity. Presence of three or more of these factors was associated with substantially increased risk for in-hospital mortality among patients with acute pancreatitis. In addition, an elevated hematocrit >44% and admission BUN >22 mg/dL are also associated with more severe acute pancreatitis. Incorporating these indices with the overall patient response to initial fluid resuscitation in the emergency ward can be useful at triaging patients to the appropriate hospital acute care setting.

In general, patients with lower BISAP scores, hematocrits, and admission BUNs tend to respond to initial management and are triaged to a regular hospital ward for ongoing care. If SIRS is not present at 24 h, the patient is unlikely to develop organ failure or necrosis. Therefore, patients with persistent SIRS at 24 h or underlying comorbid illnesses (e.g., chronic obstructive pulmonary disease, congestive heart failure) should be considered for a step-down unit setting if available. Patients with higher BISAP scores and elevations in hematocrit and admission BUN that do not respond to initial fluid resuscitation and exhibit evidence of respiratory failure, hypotension, or organ failure should be considered for direct admission to an intensive care unit.

Special Considerations Based on Etiology A careful history, review of medications, selected laboratory studies (liver profile, serum triglycerides, serum calcium), and an abdominal ultrasound are recommended in the emergency ward to assess for etiologies that may impact acute

management. An abdominal ultrasound is the initial imaging modality of choice and will evaluate the gallbladder and common duct and assess the pancreatic head.

GALLSTONE PANCREATITIS Patients with evidence of ascending cholangitis (rising white blood cell count, increasing liver enzymes) should undergo ERCP within 24–48 h of admission. Patients with gallstone pancreatitis are at increased risk of recurrence, and consideration should be given to performing a cholecystectomy during the same admission or within 4–6 weeks of discharge. An alternative for patients who are not surgical candidates would be to perform an endoscopic biliary sphincterotomy before discharge.

HYPERTRIGLYCERIDEMIA Serum triglycerides >1000 mg/dL are associated with acute pancreatitis. Initial therapy may include insulin, heparin, or plasmapheresis. Outpatient therapies include control of diabetes if present, administration of lipid-lowering agents, weight loss, and avoidance of drugs that elevate lipid levels.

Other potential etiologies that may impact acute hospital care include *hypercalcemia*, *autoimmune pancreatitis*, *post-ERCP pancreatitis*, and *drug-induced pancreatitis*. Treatment of hyperparathyroidism or malignancy is effective at reducing serum calcium. Autoimmune pancreatitis is responsive to glucocorticoid administration. Pancreatic duct stenting and rectal indomethacin administration are effective at decreasing pancreatitis after ERCP. Drugs that cause pancreatitis should be discontinued. Multiple drugs have been implicated, but only about 30 have been challenged (Class 1A) and found to be causative.

Nutritional Therapy A low-fat solid diet can be administered to subjects with mild acute pancreatitis after the abdominal pain has resolved. Enteral nutrition should be considered 2–3 days after admission in subjects with more severe pancreatitis instead of total parenteral nutrition (TPN). Enteral feeding maintains gut barrier integrity, limits bacterial translocation, is less expensive, and has fewer complications than TPN. The choice of gastric versus nasojejunal enteral feeding is currently under investigation.

Management of Local Complications (Table 371-4) Patients exhibiting signs of clinical deterioration despite aggressive fluid resuscitation and hemodynamic monitoring should be assessed for local complications, which may include necrosis, pseudocyst formation, pancreas duct disruption, peripancreatic vascular complications, and extrapancreatic infections. A multidisciplinary team approach is recommended including gastroenterology, surgery, interventional radiology, and intensive care specialists, and consideration should also be made for transfer to a pancreas center.

NECROSIS The management of necrosis requires a multidisciplinary team approach. Percutaneous aspiration of necrosis with Gram stain and culture should be performed if there are ongoing signs of possible pancreatic infection such as sustained leukocytosis, fever, or organ failure. There is currently no role for *prophylactic antibiotics* in necrotizing pancreatitis. It is reasonable to start broad-spectrum antibiotics in a patient who appears septic while awaiting the results of Gram stain and cultures. If cultures are negative, the antibiotics should be discontinued to minimize the risk of developing opportunistic or fungal superinfection. Repeated fine-needle aspiration and Gram stain with culture of pancreatic necrosis may be done every 5–7 days in the presence of persistent fever. Repeated CT or MRI imaging should also be considered with any change in clinical course to monitor for complications (e.g., thromboses, hemorrhage, abdominal compartment syndrome).

In general, *sterile necrosis* is most often managed conservatively unless complications arise. Once a diagnosis of *infected necrosis* is established and an organism identified, targeted antibiotics should be instituted. Pancreatic debridement (necrosectomy) should be considered for definitive management of *infected necrosis*, but clinical decisions are generally influenced by response to antibiotic treatment and overall clinical condition. Symptomatic local complications as outlined in the revised Atlanta criteria may require definitive therapy.

TABLE 371-4 COMPLICATIONS OF ACUTE PANCREATITIS

Local
Necrosis
Sterile
Infected
Walled-off necrosis
Pancreatic fluid collections
Pancreatic pseudocyst
Disruption of main pancreatic duct or secondary branches
Pancreatic ascites
Involvement of contiguous organs by necrotizing pancreatitis
Thrombosis of blood vessels (splenic vein, portal vein)
Pancreatic enteric fistula
Bowel infarction
Obstructive jaundice
Systemic
Pulmonary
Pleural effusion
Atelectasis
Mediastinal fluid
Pneumonitis
Acute respiratory distress syndrome
Hypoxemia (unrecognized)
Cardiovascular
Hypotension
Hypovolemia
Nonspecific ST-T changes in electrocardiogram simulating myocardial infarction
Pericardial effusion
Hematologic
Disseminated intravascular coagulation
Gastrointestinal hemorrhage
Peptic ulcer disease
Erosive gastritis
Hemorrhagic pancreatic necrosis with erosion into major blood vessels
Portal vein thrombosis, splenic vein thrombosis, variceal hemorrhage
Renal
Oliguria (<300 mL/d)
Azotemia
Renal artery and/or renal vein thrombosis
Acute tubular necrosis
Metabolic
Hyperglycemia
Hypertriglyceridemia
Hypocalcemia
Encephalopathy
Sudden blindness (Purtscher's retinopathy)
Central nervous system
Psychosis
Fat emboli
Fat necrosis
Subcutaneous tissues (erythematous nodules)
Bone
Miscellaneous (mediastinum, pleura, nervous system)

A step-up approach (percutaneous or endoscopic transgastric drainage followed, if necessary, by open necrosectomy) has been successfully reported by some pancreatic centers. One-third of the patients successfully treated with the step-up approach did not require major abdominal surgery. A recent randomized trial reported advantages

to an initial endoscopic approach compared to an initial surgical necrosectomy approach in select patients requiring intervention for symptomatic WON. Taken together, a more conservative approach to the management of infected pancreatic necrosis has evolved over the years under the close supervision of a multidisciplinary team. If conservative therapy can be safely implemented for 4–6 weeks, to allow the pancreatic collections to resolve or “wall-off,” surgical or endoscopic intervention is generally much safer and better tolerated by the patient.

PSEUDOCYST The incidence of pseudocyst is low, and most acute collections resolve over time. Less than 10% of patients have persistent fluid collections after 6 weeks that would meet the definition of a pseudocyst. Only symptomatic collections should be drained with surgery or endoscopy or by percutaneous route.

PANCREATIC DUCT DISRUPTION Pancreatic duct disruption may present with symptoms of increasing abdominal pain or shortness of breath in the setting of an enlarging fluid collection. Diagnosis can be confirmed on magnetic resonance cholangiopancreatography (MRCP) or ERCP. Placement of a bridging pancreatic stent for at least 6 weeks is >90% effective at resolving the leak. Nonbridging stents are less effective.

PERIVASCULAR COMPLICATIONS Perivascular complications may include *splenic vein thrombosis* with gastric varices and pseudoaneurysms. *Gastric varices* bleed less than 5% of the time. Life-threatening bleeding from a ruptured *pseudoaneurysm* can be diagnosed and treated with mesenteric angiography and embolization.

EXTRAPANCREATIC INFECTIONS Hospital-acquired infections occur in up to 20% of patients with acute pancreatitis. Patients should be continually monitored for the development pneumonia, urinary tract infection, and line infection. Continued culturing of urine, monitoring of chest x-rays, and routine changing of intravenous lines are important during hospitalization.

Follow-Up Care Hospitalizations for moderately severe and severe acute pancreatitis can last weeks to months and often involve a period of intensive care unit admission and outpatient rehabilitation or subacute nursing care. Follow-up evaluation should assess for development of diabetes, exocrine insufficiency, recurrent cholangitis, or development of infected fluid collections. As mentioned previously, cholecystectomy should be performed within 4–6 weeks of discharge if possible for patients with uncomplicated gallstone pancreatitis.

RECURRENT PANCREATITIS

Approximately 25% of patients who have had an attack of acute pancreatitis have a recurrence. The two most common etiologic factors are alcohol and cholelithiasis. In patients with recurrent pancreatitis without an obvious cause, the differential diagnosis should encompass occult biliary tract disease including microlithiasis, hypertriglyceridemia, drugs, pancreatic cancer, pancreas divisum, and cystic fibrosis (Table 371-1). In one series of 31 patients diagnosed initially as having idiopathic or recurrent acute pancreatitis, 23 were found to have occult gallstone disease. Thus, approximately two-thirds of patients with recurrent acute pancreatitis without an obvious cause actually have occult gallstone disease due to microlithiasis. Genetic defects as in hereditary pancreatitis and cystic fibrosis mutations can result in recurrent pancreatitis. Other diseases of the biliary tree and pancreatic ducts that can cause acute pancreatitis include choledochocoele; ampullary tumors; pancreas divisum; and pancreatic duct stones, stricture, and tumor. Approximately 2–4% of patients with pancreatic carcinoma present with acute pancreatitis.

PANCREATITIS IN PATIENTS WITH AIDS

The incidence of acute pancreatitis is increased in patients with AIDS for two reasons: (1) the high incidence of infections involving the pancreas such as infections with cytomegalovirus, *Cryptosporidium*, and the *Mycobacterium avium* complex; and (2) the frequent use by patients with AIDS of medications such as didanosine, pentamidine,

2098 trimethoprim-sulfamethoxazole, and protease inhibitors. Incidence has been markedly reduced due to advances in therapy (Chap. 226).

CHRONIC PANCREATITIS AND PANCREATIC EXOCRINE INSUFFICIENCY

PATHOPHYSIOLOGY

Chronic pancreatitis is a disease process characterized by irreversible damage to the pancreas as distinct from the reversible changes noted in acute pancreatitis (Table 371-4). The events that initiate and then perpetuate the inflammatory process in the pancreas are becoming more clearly understood. Irrespective of the mechanism of injury, it is becoming apparent that stellate cell activation that results in cytokine expression and production of extracellular matrix proteins cause acute and chronic inflammation and collagen deposition in the pancreas. Thus, the condition is defined by the presence of histologic abnormalities, including chronic inflammation, fibrosis, and progressive destruction of both exocrine and eventually endocrine tissue (atrophy). A number of etiologies have been associated with chronic pancreatitis resulting in the cardinal manifestations of the disease such as abdominal pain, steatorrhea, weight loss, and diabetes mellitus (Table 371-5).

Although alcohol has been believed to be the primary cause of chronic pancreatitis, other factors contribute to the disease because not all heavy consumers of alcohol develop pancreatic disease. There is also a strong association between smoking and chronic pancreatitis. Cigarette smoke leads to an increased susceptibility to pancreatic auto-digestion and predisposes to dysregulation of duct cell CFTR function.

Smoking is an independent, dose-dependent risk factor for chronic pancreatitis and recurrent acute pancreatitis. Both continued alcohol and smoking exposure are associated with pancreatic fibrosis, calcifications, and progression of disease

Recent characterization of pancreatic stellate cells (PSCs) has added insight into the underlying cellular responses behind development of chronic pancreatitis. Specifically, PSCs are believed to play a role in maintaining normal pancreatic architecture that can shift toward fibrogenesis in the case of chronic pancreatitis. The sentinel acute pancreatitis event (SAPE) hypothesis uniformly describes the events in the pathogenesis of chronic pancreatitis. It is believed that alcohol or additional stimuli lead to matrix metalloproteinase-mediated destruction of normal collagen in pancreatic parenchyma, which later allows for pancreatic remodeling. Proinflammatory cytokines, tumor necrosis factor α (TNF- α), interleukin 1 (IL-1), and interleukin 6 (IL-6), as well as oxidant complexes, are able to induce PSC activity with subsequent new collagen synthesis. In addition to being stimulated by cytokines, oxidants, or growth factors, PSCs also possess transforming growth factor β (TGF- β)-mediated self-activating autocrine pathways that may explain disease progression in chronic pancreatitis even after removal of noxious stimuli.

ETIOLOGIC CONSIDERATIONS

Among adults in the United States, alcoholism is the most common cause of clinically apparent chronic pancreatitis, whereas cystic fibrosis is the most frequent cause in children. As many as 25% of adults in the United States with chronic pancreatitis have the *idiopathic* form. Recent investigations have indicated that up to 15% of patients with idiopathic pancreatitis may have pancreatitis due to genetic defects (Table 371-5).

Whitcomb and associates studied several large families with hereditary chronic pancreatitis and were able to identify a genetic defect that affects the gene encoding for trypsinogen. Several additional defects of this gene have also been described. The defect prevents the destruction of prematurely activated trypsin and allows it to be resistant to the intracellular protective effect of trypsin inhibitor. It is hypothesized that this continual activation of digestive enzymes within the gland leads to acute injury and, finally, chronic pancreatitis. Since the initial discovery of the *PRSS1* mutation defect, other genetic diseases have been detected (Table 371-5).

Several other groups of investigators have documented mutations of *CFTR*. This gene functions as a cyclic AMP-regulated chloride channel. In patients with cystic fibrosis, the high concentration of macromolecules can block the pancreatic ducts. It must be appreciated, however, that there is a great deal of heterogeneity in relationship to the *CFTR* gene defect. More than 1000 putative mutations of the *CFTR* gene have been identified. Attempts to elucidate the relationship between the genotype and pancreatic manifestations have been hampered by the number of mutations. The ability to detect *CFTR* mutations has led to the recognition that the clinical spectrum of the disease is broader than previously thought. Two studies have clarified the association between mutations of the *CFTR* gene and another monosymptomatic form of cystic fibrosis (i.e., chronic pancreatitis). It is estimated that in patients with idiopathic pancreatitis, the frequency of a single *CFTR* mutation is 11 times the expected frequency and the frequency of two mutant alleles is 80 times the expected frequency. In these studies, the patients were adults when the diagnosis of pancreatitis was made; none had any clinical evidence of pulmonary disease, and sweat test results were not diagnostic of cystic fibrosis. The prevalence of such mutations is unclear, and further studies are certainly needed. In addition, the therapeutic and prognostic implication of these findings with respect to managing pancreatitis remains to be determined. Long-term follow-up of affected patients is needed. *CFTR* mutations are common in the general population. It is unclear whether the *CFTR* mutation alone can lead to pancreatitis as an autosomal recessive disease. A study evaluated 39 patients with idiopathic chronic pancreatitis to assess the risk associated with these mutations. Patients with two *CFTR* mutations (compound heterozygotes) demonstrated *CFTR* function at a level between that seen in typical cystic fibrosis and cystic

TABLE 371-5 CHRONIC PANCREATITIS AND PANCREATIC EXOCRINE INSUFFICIENCY: TIGAR-O CLASSIFICATION SYSTEM

Toxic-metabolic

- Alcoholic
- Tobacco smoking
- Hypercalcemia
- Hyperlipidemia
- Chronic renal failure
- Medications—phenacetin abuse
- Toxins—organotin compounds (e.g., dibutyltin dichloride, DBTC)

Idiopathic

- Early onset
- Late onset
- Tropical

Genetic

- Cationic trypsinogen (*PRSS1*)
- Cystic fibrosis transmembrane conductance regulator gene (*CFTR*)
- Calcium-sensing receptor (*CASR*)
- Chymotrypsin C gene (*CTRC*)
- Pancreatic secretory trypsin inhibitor gene (*SPINK1*)

Autoimmune

- Type 1 autoimmune chronic pancreatitis
- IgG4 systemic
- Type 2 autoimmune chronic pancreatitis

Recurrent and severe acute pancreatitis

- Postnecrotic (severe acute pancreatitis)
- Recurrent acute pancreatitis
- Vascular diseases/ischemia
- Radiation induced

Obstructive

- Pancreas divisum
- Duct obstruction (e.g., tumor)
- Preampullary duodenal wall cysts
- Posttraumatic pancreatic duct scars

Abbreviations: DBTC, dibutyltin dichloride; TIGAR-O, toxic-metabolic, idiopathic, genetic, autoimmune, recurrent and severe acute pancreatitis, obstructive.

fibrosis carriers and had a 40-fold increased risk of pancreatitis. The presence of an *N34S SPINK1* mutation increased the risk 20-fold. A combination of two *CFTR* mutations and an *N34S SPINK1* mutation increased the risk of pancreatitis 900-fold. Knowledge of the genetic defects and downstream alterations in protein expression has led to the development of novel genetic therapy in cystic fibrosis children that potentiates the *CFTR* channel resulting in improvement in lung function, quality of life, and weight gain. Table 371-5 lists recognized causes of chronic pancreatitis and pancreatic exocrine insufficiency.

AUTOIMMUNE PANCREATITIS (TABLE 371-6)

Autoimmune pancreatitis (AIP) is an uncommon disorder of presumed autoimmune causation with characteristic laboratory, histologic, and morphologic findings. In type I AIP, the pancreas is involved as part of an IgG4 systemic disease (Chap. 391e) and meets HISORt criteria as defined below. The characteristic pancreatic histopathologic findings include lymphoplasmacytic infiltrate, storiform fibrosis, and abundant IgG4 cells. AIP type 2 is histologically confirmed idiopathic duct centric pancreatitis with granulocytic infiltration of the duct wall (termed GEL), but without IgG4 positive cells and systemic involvement. Although AIP was initially described as a primary pancreatic disorder, it is now recognized that it is associated with other disorders of presumed autoimmune etiology, and this has been termed IgG4 systemic disease (Chap. 391e). The clinical features include IgG4-associated cholangitis, rheumatoid arthritis, Sjögren's syndrome, ulcerative colitis, mediastinal fibrosis and adenopathy, autoimmune thyroiditis, tubulointerstitial nephritis, retroperitoneal fibrosis, chronic periaortitis, chronic sclerosing sialadenitis, and Mikulicz's disease. Mild symptoms, usually abdominal pain, and recurrent acute pancreatitis are unusual. Furthermore, AIP is not a common cause of idiopathic recurrent pancreatitis.

Weight loss and new onset of diabetes may also occur. An obstructive pattern on liver tests is common (i.e., disproportionately elevated serum alkaline phosphatase and minimally elevated serum aminotransferases). Elevated serum levels of IgG4 provide a marker for the disease, particularly in Western populations. Serum IgG4 normally accounts for only 5–6% of the total IgG4 in healthy patients but is elevated to values >280 mg/dL in those with AIP. CT scans reveal abnormalities in the majority of patients and include diffuse enlargement, focal enlargement, and a distinct enlargement at the head of the pancreas. ERCP or MRCP reveals strictures in the bile duct in more than one-third of patients with AIP; these may include common bile duct strictures, intrahepatic bile duct strictures, or proximal bile duct strictures, with accompanying narrowing of the pancreatic portion of the bile duct. This has been termed autoimmune IgG4 cholangitis. Characteristic histologic findings include extensive lymphoplasmacytic infiltrates

with dense fibrosis around pancreatic ducts, as well as a lymphoplasmacytic infiltration, resulting in an obliterative phlebitis.

The Mayo Clinic HISORt criteria indicate that AIP can be diagnosed by the presence of at least two of the following: (1) histology; (2) imaging; (3) serology (elevated serum IgG4 levels); (4) other organ involvement; and (5) response to glucocorticoid therapy, with improvement in pancreatic and extrapancreatic manifestations.

Glucocorticoids have shown efficacy in alleviating symptoms, decreasing the size of the pancreas, and reversing histopathologic features in patients with AIP. Patients may respond dramatically to glucocorticoid therapy within a 2- to 4-week period. Prednisone is usually administered at an initial dose of 40 mg/d for 4 weeks followed by a taper of the daily dosage by 5 mg/wk based on monitoring of clinical parameters. Relief of symptoms, serial changes in abdominal imaging of the pancreas and bile ducts, decreased serum γ -globulin and IgG4 levels, and improvements in liver tests are parameters to follow. A poor response to glucocorticoids over a 2- to 4-week period should raise suspicion of pancreatic cancer or other forms of chronic pancreatitis. A recent multicenter international report reviewed 1064 patients with AIP. Clinical remission was achieved in 99% of type I and 92% of type II AIP patients with steroids. However, disease relapse occurred in 31% of type I and 9% of type II AIP patients. For treatment of disease relapse in type I AIP, glucocorticoids were successful in 201 of 295 (68%) patients, and azathioprine was successful in 52 of 58 patients (85%). A small number of patients responded favorably to 6-mercaptopurine, rituximab, cyclosporine, and cyclophosphamide. Types 1 and 2 AIP are highly responsive to initial glucocorticoid treatment. Relapse is common in type 1 patients, especially those with biliary tract strictures. Most relapses occur after glucocorticoids are discontinued. Patients with refractory symptoms and strictures generally require immunomodulator therapy as noted above. Appearance of interval cancers following a diagnosis of AIP is uncommon.

Clinical Features of Chronic Pancreatitis Patients with chronic pancreatitis seek medical attention predominantly because of two symptoms: abdominal pain or maldigestion and weight loss. The abdominal pain may be quite variable in location, severity, and frequency. The pain can be constant or intermittent with frequent pain-free intervals. Eating may exacerbate the pain, leading to a fear of eating with consequent weight loss. The spectrum of abdominal pain ranges from mild to quite severe, with narcotic dependence as a frequent consequence. Maldigestion is manifested as chronic diarrhea, steatorrhea, weight loss, and fatigue. Patients with chronic abdominal pain may or may not progress to maldigestion, and ~20% of patients will present with symptoms of maldigestion without a history of abdominal pain. Patients with chronic pancreatitis have significant morbidity and mortality and use appreciable amounts of societal resources. Despite steatorrhea, clinically apparent deficiencies of fat-soluble vitamins are surprisingly uncommon. Physical findings in these patients are usually unimpressive, so that there is a disparity between the severity of abdominal pain and the physical signs that usually consist of some mild tenderness.

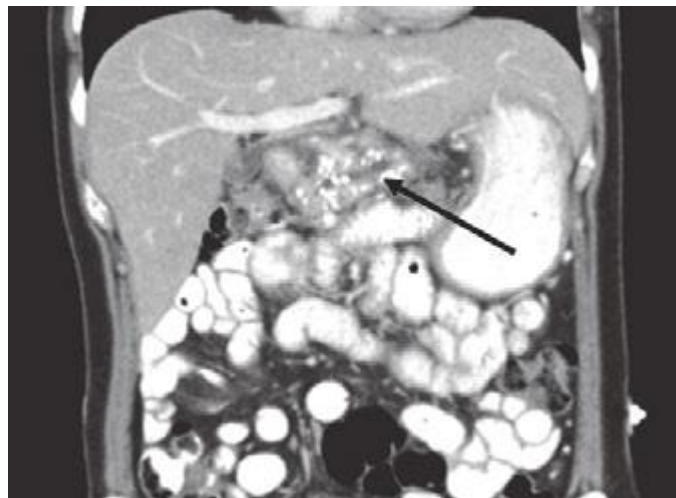
The diagnosis of early or mild chronic pancreatitis can be challenging because there is no biomarker for the disease. In contrast to acute pancreatitis, the serum amylase and lipase levels are usually not strikingly elevated in chronic pancreatitis. Elevation of serum bilirubin and alkaline phosphatase may indicate cholestasis secondary to common bile duct stricture caused by chronic inflammation. Many patients have impaired glucose tolerance with elevated fasting blood glucose levels. The fecal elastase-1 and small-bowel biopsy are useful in the evaluation of patients with suspected pancreatic steatorrhea. The fecal elastase level will be abnormal and small-bowel histology will be normal in such patients. A decrease of fecal elastase level to <100 μ g per gram of stool strongly suggests severe pancreatic exocrine insufficiency.

The radiographic evaluation of a patient with suspected chronic pancreatitis usually proceeds from a noninvasive to more invasive approach. Abdominal CT imaging (Fig. 371-4A,B) is the initial modality of choice, followed by MRI (Fig. 371-4C), endoscopic ultrasound, and pancreas function testing. In addition to excluding a

TABLE 371-6 CLINICAL FEATURES OF AUTOIMMUNE PANCREATITIS (AIP)

- Mild symptoms, usually abdominal pain, but without frequent attacks of acute pancreatitis
- Diffuse swelling and enlargement of the pancreas
- Two-thirds of patients present with either obstructive jaundice or a "mass" in the head of the pancreas mimicking carcinoma
- Diffuse irregular narrowing of the pancreatic duct (MRCP or ERCP)
- Increased levels of serum gamma globulins, especially IgG4
- Presence of other autoantibodies (ANA), rheumatoid factor (RF)
- Can occur with other autoimmune diseases: Sjögren's syndrome, primary sclerosing cholangitis, ulcerative colitis, rheumatoid arthritis
- Extrapancreatic bile duct changes such as stricture of the common bile duct and intrahepatic ducts
- Pancreatic calcifications (rare)
- Pancreatic biopsies reveal extensive fibrosis and lymphoplasmacytic infiltration
- Glucocorticoids are effective in alleviating symptoms, decreasing size of the pancreas, and reversing histopathologic changes

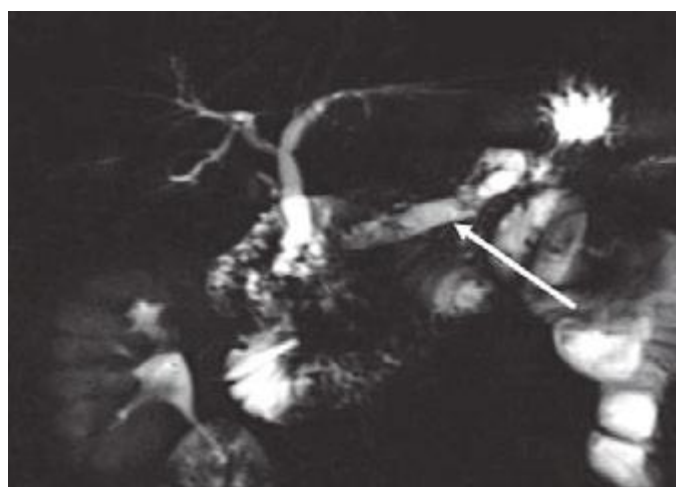
Abbreviations: ERCP, endoscopic retrograde cholangiopancreatography; MRCP, magnetic resonance cholangiopancreatography.



A



B



C

FIGURE 371-4 **A.** Chronic pancreatitis and pancreatic calculi: computed tomography (CT) scan. In this contrast-enhanced CT scan of the abdomen, there is evidence of an atrophic pancreas with multiple calcifications and stones in the parenchyma and dilated pancreatic duct (*arrow*). **B.** In this contrast-enhanced CT scan of the abdomen, there is evidence of an atrophic pancreas with multiple calcifications (*arrows*). Note the markedly dilated pancreatic duct seen in this section through the body and tail (*open arrows*). **C.** Chronic pancreatitis on magnetic resonance cholangiopancreatography (MRCP): dilated duct with filling defects. Gadolinium-enhanced magnetic resonance imaging/MRCP reveals a dilated pancreatic duct (*arrow*) in chronic pancreatitis with multiple filling defects suggestive of pancreatic duct calculi. (*A, C, courtesy of Dr. KJ Morteale, Brigham and Women's Hospital; with permission.*)

pseudocyst and pancreatic cancer, CT may show calcification, dilated ducts, or an atrophic pancreas. Although abdominal CT scanning and MRCP greatly aid in the diagnosis of pancreatic disease, the diagnostic test with the best sensitivity and specificity is the hormone stimulation test using secretin. The secretin test becomes abnormal when $\geq 60\%$ of the pancreatic exocrine function has been lost. This usually correlates well with the onset of chronic abdominal pain. The role of endoscopic ultrasonography (EUS) in diagnosing early chronic pancreatitis is still being defined. A total of nine endosonographic features have been described in chronic pancreatitis. The presence of five or more features is considered diagnostic of chronic pancreatitis. EUS is not a sensitive enough test for detecting early chronic pancreatitis alone (**Chap. 370**) and may show positive features in patients who have dyspepsia or even normal aging individuals. Recent data suggest that EUS can be combined with endoscopic pancreatic function testing (EUS-ePFT) during a single endoscopy to screen for chronic pancreatitis in patients with chronic abdominal pain. Diffuse calcifications noted on plain film of the abdomen usually indicate significant damage to the pancreas and are pathognomonic for chronic pancreatitis (**Fig. 371-4A**). Although alcohol is by far the most common cause of pancreatic calcification, such calcification may also be noted in hereditary pancreatitis, post-traumatic pancreatitis, hypercalcemic pancreatitis, idiopathic chronic pancreatitis, and tropical pancreatitis.

Complications of Chronic Pancreatitis The complications of chronic pancreatitis are protean and are listed in **Table 371-7**. Although most patients have impaired glucose tolerance, diabetic ketoacidosis and diabetic coma are uncommon. Likewise, end-organ damage (retinopathy, neuropathy, nephropathy) is also uncommon. A nondiabetic retinopathy may be due to either vitamin A and/or zinc deficiency. Gastrointestinal bleeding may occur from peptic ulceration, gastritis, a pseudocyst eroding into the duodenum, arterial bleeding into the pancreatic duct (hemorrhagic pancreatitis), or ruptured varices secondary to splenic vein thrombosis due to chronic inflammation of the tail of the pancreas. Jaundice, cholestasis, and biliary cirrhosis may occur from the chronic inflammatory reaction around the intrapancreatic portion of the common bile duct. Twenty years after the diagnosis of calcific chronic pancreatitis, the cumulative risk of pancreatic carcinoma is 4%. Patients with hereditary pancreatitis are at a 10-fold higher risk for pancreatic cancer.

TREATMENT CHRONIC PANCREATITIS

STEATORRHEA

The treatment of steatorrhea with pancreatic enzymes is straightforward even though complete correction of steatorrhea is unusual. Enzyme therapy usually brings diarrhea under control and restores absorption of fat to an acceptable level and affects weight gain. Thus, pancreatic enzyme replacement has been the cornerstone of therapy. In treating steatorrhea, it is important to use a potent pancreatic formulation that will deliver sufficient lipase into the duodenum to correct maldigestion and decrease steatorrhea. In an attempt to standardize the enzyme activity, potency, and bioavailability, the U.S. Food and Drug Administration (FDA) required that all pancreas enzyme drugs in the United States obtain a New Drug Application (NDA) by April 2008. **Table 371-8** lists frequently used formulations, but availability will be based on compliance with the FDA mandate. Recent data suggest that dosages up to

TABLE 371-7 COMPLICATIONS OF CHRONIC PANCREATITIS

Chronic abdominal pain	Jaundice
Narcotic addiction	Retinopathy
Diabetes mellitus/impaired glucose tolerance	Biliary stricture and/or biliary cirrhosis
Gastroparesis	Pseudocyst
Malabsorption/maldigestion	Metabolic bone disease
	Pancreatic cancer

TABLE 371-8 FDA-APPROVED PANCREATIC ENZYME (PANCRELIPASE) PREPARATIONS

Product	Enzyme Content/Unit Dose, U.S. Pharmacopeia Units		
	Lipase ^a	Amylase ^a	Protease ^a
Immediate-Release Capsule			
Non-enteric-coated			
Viokace 10,440	10,440	391,550	39,150
Viokace 20,880	20,880	78,300	78,300
Delayed-Release Capsules			
Enteric-coated mini-microspheres			
Creon 3000	3000	15,000	9500
Creon 6000	6000	30,000	19,000
Creon 12,000	12,000	60,000	38,000
Creon 24,000	24,000	120,000	76,000
Enteric-Coated Mini-Tablets			
Ultresa 13,800	13,800	27,600	27,600
Ultresa 20,700	20,700	41,400	41,400
Ultresa 23,000	23,000	46,000	46,000
Enteric-Coated Beads			
Zenpep 3000	3000	16,000	10,000
Zenpep 5000	5000	27,000	17,000
Zenpep 10,000	10,000	55,000	34,000
Zenpep 15,000	15,000	82,000	51,000
Zenpep 20,000	20,000	109,000	68,000
Zenpep 25,000	25,000	136,000	85,000
Enteric-Coated Micro-Tablets			
Pancreaze 4200	4200	17,500	10,000
Pancreaze 10,500	10,500	43,750	25,000
Pancreaze 16,800	16,800	70,000	40,000
Pancreaze 21,000	21,000	61,000	37,000
Bicarbonate-Buffered Enteric-Coated Microspheres			
Pertzye 8000	8000	30,250	28,750
Pertzye 16,000	16,000	60,500	57,500

^aU.S. Pharmacopeia (USP) units per tablet or capsule

Note: The FDA has mandated all enzyme manufacturers to submit New Drug Applications (NDAs) for all pancreatic extract drug products after reviewing data that showed substantial variations among currently marketed products. Numerous manufacturers have investigations under way to seek FDA approval for the treatment of exocrine pancreatic insufficiency due to cystic fibrosis or other conditions under the new guidelines for this class of drugs (www.fda.gov).

80,000–100,000 units of lipase taken during the meal may be necessary to normalize nutritional parameters in malnourished chronic pancreatitis patients, and some may require acid suppression with proton pump inhibitors.

ABDOMINAL PAIN

The management of pain in patients with chronic pancreatitis is problematic.

Recent meta-analyses have shown no consistent benefit of *enzyme therapy* at reducing pain in chronic pancreatitis. In some patients with idiopathic chronic pancreatitis, conventional non-enteric-coated enzyme preparations containing high concentrations of serine proteases may relieve mild abdominal pain or discomfort. The pain relief experienced by these patients actually may be due to improvements in the dyspepsia from maldigestion.

Gastroparesis is also quite common in patients with chronic pancreatitis. It is important to recognize and treat with prokinetic drugs because treatment with enzymes may fail simply because gastric dysmotility is interfering with the delivery of enzymes into the upper intestine. A recent prospective study reported that pregabalin can improve pain in chronic pancreatitis and lower pain medication requirement.

Endoscopic treatment of chronic pancreatitis pain may involve sphincterotomy, stenting, stone extraction, and drainage of a pancreatic pseudocyst. Therapy directed to the pancreatic duct would seem to be most appropriate in the setting of a dominant stricture, especially if a ductal stone has led to obstruction. The use of endoscopic stenting for patients with chronic pain, but without a dominant stricture, has not been subjected to any controlled trials. It is now appreciated that significant complications can occur from stenting (i.e., bleeding, cholangitis, stent migration, pancreatitis, and stent clogging). In patients with large-duct disease usually from alcohol-induced chronic pancreatitis, ductal decompression with *surgical therapy* has been the therapy of choice. Among such patients, 80% seem to obtain immediate relief; however, at the end of 3 years, one-half of the patients have recurrence of pain. Two randomized prospective trials comparing endoscopic to surgical therapy for chronic pancreatitis demonstrated that surgical therapy was superior to endoscopy at decreasing pain and improving quality of life in selected patients with dilated ducts and abdominal pain. This would suggest that chronic pancreatitis patients with dilated ducts and pain should be considered for surgical intervention. The role of preoperative stenting prior to surgery as a predictor of response has yet to be proven.

A Whipple procedure, total pancreatectomy, and autologous islet cell transplantation have been used in selected patients with chronic pancreatitis and abdominal pain refractory to conventional therapy. The patients who have benefited the most from total pancreatectomy have chronic pancreatitis without prior pancreatic surgery or evidence of islet cell insufficiency. The role of this procedure remains to be fully defined but may be an option in lieu of ductal decompression surgery or pancreatic resection in patients with intractable, painful small-duct disease, particularly as the standard surgical procedures tend to decrease islet cell yield. Celiac plexus block has not resulted in long-lasting pain relief.

HEREDITARY PANCREATITIS

Hereditary pancreatitis is a rare disease that is similar to chronic pancreatitis except for an early age of onset and evidence of hereditary factors. A genomewide search using genetic linkage analysis identified the hereditary pancreatitis gene on chromosome 7. Mutations in ion codons 29 (exon 2) and 122 (exon 3) of the cationic trypsinogen gene cause autosomal dominant forms of hereditary pancreatitis. The codon 122 mutations lead to a substitution of the corresponding arginine with another amino acid, usually histidine. This substitution, when it occurs, eliminates a fail-safe trypsin self-destruction site necessary to eliminate trypsin that is prematurely activated within the acinar cell. These patients have recurring attacks of severe abdominal pain that may last from a few days to a few weeks. The serum amylase and lipase levels may be elevated during acute attacks but are usually normal. Patients frequently develop pancreatic calcification, diabetes mellitus, and steatorrhea; in addition, they have an increased incidence of pancreatic carcinoma, with the cumulative incidence being as high as 40% by age 70 years. A recent natural history study of hereditary pancreatitis in more than 200 patients from France reported that abdominal pain started in childhood at age 10 years, steatorrhea developed at age 29 years, diabetes at age 38 years, and pancreatic carcinoma at age 55 years. Such patients often require surgical ductal decompression for pain relief. Abdominal complaints in relatives of patients with hereditary pancreatitis should raise the question of pancreatic disease.

PSTI, or SPINK1, is a 56-amino-acid peptide that specifically inhibits trypsin by physically blocking its active site. SPINK1 acts as the first line of defense against prematurely activated trypsinogen in the acinar cell. Recently, it has been shown that the frequency of SPINK1 mutations in patients with idiopathic chronic pancreatitis is markedly increased, suggesting that these mutations may be associated with pancreatitis.

PANCREATIC ENDOCRINE TUMORS

Pancreatic endocrine tumors are discussed in Chap. 113.

ANNULAR PANCREAS

When the ventral pancreatic anlage fails to migrate correctly to make contact with the dorsal anlage, the result may be a ring of pancreatic tissue encircling the duodenum. Such an annular pancreas may cause intestinal obstruction in the neonate or the adult. Symptoms of postprandial fullness, epigastric pain, nausea, and vomiting may be present for years before the diagnosis is entertained. The radiographic findings are symmetric dilation of the proximal duodenum with bulging of the recesses on either side of the annular band, effacement but not destruction of the duodenal mucosa, accentuation of the findings in the right anterior oblique position, and lack of change on repeated examinations. The differential diagnosis should include duodenal webs, tumors of the pancreas or duodenum, postbulbar peptic ulcer, regional enteritis, and adhesions. Patients with annular pancreas have an increased incidence of pancreatitis and peptic ulcer. Because of these and other potential complications, the treatment is surgical even if the condition has been present for years. Retrocolic duodenojejunostomy is the procedure of choice, although some surgeons advocate Billroth II gastrectomy, gastroenterostomy, and vagotomy.

PANCREAS DIVISUM

Pancreas divisum is present in 7–10% of the population and occurs when the embryologic ventral and dorsal pancreatic anlagen fail to fuse, so that pancreatic drainage is accomplished mainly through the accessory papilla. Pancreas divisum is the most common congenital anatomic variant of the human pancreas. Current evidence indicates that this anomaly does not predispose to the development of pancreatitis in the great majority of patients who harbor it. However, the combination of pancreas divisum and a small accessory orifice could result in dorsal duct obstruction. The challenge is to identify this

subset of patients with dorsal duct pathology. Cannulation of the dorsal duct by ERCP is not as easily done as is cannulation of the ventral duct. Patients with pancreatitis and pancreas divisum demonstrated by MRCP or ERCP should be treated with conservative measures. In many of these patients, pancreatitis is idiopathic and unrelated to the pancreas divisum. Endoscopic or surgical intervention is indicated only if pancreatitis recurs and no other cause can be found. If marked dilation of the dorsal duct can be demonstrated, surgical ductal decompression should be performed. It should be stressed that the ERCP/MRCP appearance of pancreas divisum (i.e., a small-caliber ventral duct with an arborizing pattern) may be mistaken as representing an obstructed main pancreatic duct secondary to a mass lesion.

MACROAMYLAEMIA

In macroamylasemia, amylase circulates in the blood in a polymer form too large to be easily excreted by the kidney. Patients with this condition demonstrate an elevated serum amylase value and a low urinary amylase value. The presence of macroamylase can be documented by chromatography of the serum. The prevalence of macroamylasemia is 1.5% of the nonalcoholic general adult hospital population. Usually macroamylasemia is an incidental finding and is not related to disease of the pancreas or other organs. Macrolipasemia has now been documented in a few patients with cirrhosis or non-Hodgkin's lymphoma. In these patients, the pancreas appeared normal on ultrasound and CT examination. Lipase was shown to be complexed with immunoglobulin A. Thus, the possibility of *both* macroamylasemia and macrolipasemia should be considered in patients with elevated blood levels of these enzymes.

ACKNOWLEDGMENTS

This chapter represents a revised version of chapters by Drs. Norton J. Greenberger, Phillip P. Toskes, and Bechien Wu that were in previous editions of Harrison's.

SECTION 1 THE IMMUNE SYSTEM IN HEALTH AND DISEASE

372e Introduction to the Immune System

Barton F. Haynes, Kelly A. Soderberg,
Anthony S. Fauci

DEFINITIONS

- **Adaptive immune system**—recently evolved system of immune responses mediated by T and B lymphocytes. Immune responses by these cells are based on specific antigen recognition by clonotypic receptors that are products of genes that rearrange during development and throughout the life of the organism. Additional cells of the adaptive immune system include various types of antigen-presenting cells.
- **Antibody**—B cell–produced molecules encoded by genes that rearrange during B cell development consisting of immunoglobulin heavy and light chains that together form the central component of the B cell receptor for antigen. Antibody can exist as B cell–surface antigen-recognition molecules or as secreted molecules in plasma and other body fluids.
- **Antigens**—foreign or self-molecules that are recognized by the adaptive and innate immune systems resulting in immune cell triggering, T cell activation, and/or B cell antibody production.
- **Antimicrobial peptides**—small peptides <100 amino acids in length that are produced by cells of the innate immune system and have anti-infectious agent activity.
- **Apoptosis**—the process of *programmed cell death* whereby signaling through various “death receptors” on the surface of cells (e.g., tumor necrosis factor [TNF] receptors, CD95) leads to a signaling cascade that involves activation of the caspase family of molecules and leads to DNA cleavage and cell death. Apoptosis, which does not lead to induction of inordinate inflammation, is to be contrasted with *cell necrosis*, which does lead to induction of inflammatory responses.
- **Autoimmune diseases**—diseases such as systemic lupus erythematosus and rheumatoid arthritis in which cells of the adaptive immune system such as autoreactive T and B cells become over-reactive and produce self-reactive T cell and antibody responses.
- **Autoinflammatory diseases**—hereditary disorders such as hereditary periodic fevers (HPFs) characterized by recurrent episodes of severe inflammation and fever due to mutations in controls of the innate inflammatory response, i.e., the inflammasome (see below and Table 372e-6). Patients with HPFs also have rashes and serosal and joint inflammation, and some can have neurologic symptoms. Autoinflammatory diseases are different from autoimmune diseases in that evidence for activation of adaptive immune cells such as autoreactive B cells is not present.
- **B cell receptor for antigen**—complex of surface molecules that rearrange during postnatal B cell development, made up of surface immunoglobulin (Ig) and associated Ig $\alpha\beta$ chain molecules that recognize nominal antigen via Ig heavy- and light-chain variable regions, and signal the B cell to terminally differentiate to make antigen-specific antibody.
- **B lymphocytes**—bone marrow-derived or bursal-equivalent lymphocytes that express surface immunoglobulin (the B cell receptor for antigen) and secrete specific antibody after interaction with antigen.
- **CD classification of human lymphocyte differentiation antigens**—the development of monoclonal antibody technology led to the discovery of a large number of new leukocyte surface molecules. In 1982, the First International Workshop on Leukocyte Differentiation Antigens was held to establish a nomenclature for cell-surface molecules of human leukocytes. From this and subsequent leukocyte differentiation workshops has come the *cluster of differentiation* (CD) classification of leukocyte antigens.
- **Chemokines**—soluble molecules that direct and determine immune cell movement and circulation pathways.
- **Complement**—cascading series of plasma enzymes and effector proteins whose function is to lyse pathogens and/or target them to be phagocytized by neutrophils and monocyte/macrophage lineage cells of the reticuloendothelial system.
- **Co-stimulatory molecules**—molecules of antigen-presenting cells (such as B7-1 and B7-2 or CD40) that lead to T cell activation when bound by ligands on activated T cells (such as CD28 or CD40 ligand).
- **Cytokines**—soluble proteins that interact with specific cellular receptors that are involved in the regulation of the growth and activation of immune cells and mediate normal and pathologic inflammatory and immune responses.
- **Dendritic cells**—myeloid and/or lymphoid lineage antigen-presenting cells of the adaptive immune system. Immature dendritic cells, or dendritic cell precursors, are key components of the innate immune system by responding to infections with production of high levels of cytokines. Dendritic cells are key initiators both of innate immune responses via cytokine production and of adaptive immune responses via presentation of antigen to T lymphocytes.
- **Ig Fc receptors**—receptors found on the surface of certain cells including B cells, natural killer cells, macrophages, neutrophils, and mast cells. Fc receptors bind to antibodies that have attached to invading pathogen-infected cells. They stimulate cytotoxic cells to destroy microbe-infected cells through a mechanism known as antibody-dependent cell-mediated cytotoxicity (ADCC). Examples of important Fc receptors include CD16 (Fc γ RIIIa), CD23 (Fc ϵ R), CD32 (Fc γ RII), CD64 (Fc γ RI), and CD89 (Fc α R).
- **Inflammasome**—large cytoplasmic complexes of intracellular proteins that link the sensing of microbial products and cellular stress to the proteolytic activation of interleukin (IL)-1 β and IL-18 inflammatory cytokines. Activation of molecules in the inflammasome is a key step in the response of the innate immune system for intracellular recognition of microbial and other danger signals in both health and pathologic states.
- **Innate immune system**—ancient immune recognition system of host cells bearing germline-encoded pattern recognition receptors that recognize pathogens and trigger a variety of mechanisms of pathogen elimination. Cells of the innate immune system include natural killer cell lymphocytes, monocytes/macrophages, dendritic cells, neutrophils, basophils, eosinophils, tissue mast cells, and epithelial cells.
- **Large granular lymphocytes**—lymphocytes of the innate immune system with azurophilic cytotoxic granules that have natural killer cell activity capable of killing foreign and host cells with few or no self-major histocompatibility complex (MHC) class I molecules.
- **Natural killer (NK) cells**—large granular lymphocytes that kill target cells expressing few or no human leukocyte antigen (HLA) class I molecules, such as malignantly transformed cells and virally infected cells. NK cells express receptors that inhibit killer cell function when self-MHC class I is present.

- *NK T cells*—innate-like lymphocytes that use an invariant T cell receptor (TCR)- α chain combined with a limited set of TCR- β chains and coexpress receptors commonly found on NK cells. NK T cells recognize lipid antigens of bacterial, viral, fungal, and protozoal infectious agents.
- *Pathogen-associated molecular patterns* (PAMPs)—Invariant molecular structures expressed by large groups of microorganisms that are recognized by host cellular pattern recognition receptors in the mediation of innate immunity.
- *Pattern recognition receptors* (PRRs)—germline-encoded receptors expressed by cells of the innate immune system that recognize PAMPs.
- *Polyreactive natural antibodies*—preexisting low-affinity antibodies produced by B cells that cross-react with multiple antigens and are available at the time of infection to bind to and coat the invading pathogen and harness innate responses to slow the infection until an adaptive high-affinity protective antibody response can be made.
- *T cell exhaustion*—state of T cells when the persistence of antigen disrupts memory T cell function, resulting in defects in memory T cell responses. Most frequently occurs in malignancies and in chronic viral infections such as HIV-1 and hepatitis C.
- *T cell receptor (TCR) for antigen*—complex of surface molecules that rearrange during postnatal T cell development made up of clonotypic TCR- α and - β chains that are associated with the CD3 complex composed of invariant γ , δ , ϵ , ζ , and η chains. TCR- α and - β chains recognize peptide fragments of protein antigen physically bound in antigen-presenting cell MHC class I or II molecules, leading to signaling via the CD3 complex to mediate effector functions.
- *T follicular helper T cells (T_{fh})*—CD4 T cells in B cell follicle germinal centers that produce IL-4 and IL-21 and drive B cell development and affinity maturation in peripheral lymphoid tissues such as lymph node and spleen.
- *T_H17 T cells*—CD4 T cells that secrete IL-17, IL-22, and IL-26 and play roles in autoimmune inflammatory disorders as well as defend against bacterial and fungal pathogens.
- *T lymphocytes*—thymus-derived lymphocytes that mediate adaptive cellular immune responses including T helper, T regulatory, and cytotoxic T lymphocyte effector cell functions.
- *Tolerance*—B and T cell nonresponsiveness to antigens that results from encounter with foreign or self-antigens by B and T lymphocytes in the absence of expression of antigen-presenting cell co-stimulatory molecules. Tolerance to antigens may be induced and maintained by multiple mechanisms either centrally (in the thymus for T cells or bone marrow for B cells) or peripherally at sites throughout the peripheral immune system.

INTRODUCTION

The human immune system has evolved over millions of years from both invertebrate and vertebrate organisms to develop sophisticated defense mechanisms to protect the host from microbes and their virulence factors. The normal immune system has three key properties: a highly diverse repertoire of antigen receptors that enables recognition of a nearly infinite range of pathogens; immune memory, to mount rapid recall immune responses; and immunologic tolerance, to avoid immune damage to normal self-tissues. From invertebrates, humans have inherited the *innate immune system*, an ancient defense system that uses germline-encoded proteins to recognize pathogens. Cells of the innate immune system, such as macrophages, dendritic cells, and NK lymphocytes, recognize PAMPs that are highly conserved among many microbes and use a diverse set of PRR molecules. Important components of the recognition of microbes by the innate immune system include recognition by germline-encoded host molecules, recognition of key microbe virulence factors but not recognition of self-molecules, and nonrecognition of benign foreign molecules or microbes. Upon contact with pathogens, macrophages and NK cells may kill pathogens directly or, in concert with dendritic cells, may activate a series of events that both slow the infection and recruit the more recently evolved arm of the human immune system, the *adaptive immune system*.

Adaptive immunity is found only in vertebrates and is based on the generation of antigen receptors on T and B lymphocytes by gene

rearrangements, such that individual T or B cells express unique antigen receptors on their surface capable of specifically recognizing diverse antigens of the myriad infectious agents in the environment. Coupled with finely tuned specific recognition mechanisms that maintain tolerance (nonreactivity) to self-antigens, T and B lymphocytes bring both *specificity* and *immune memory* to vertebrate host defenses.

This chapter describes the cellular components, key molecules (Table 372e-1), and mechanisms that make up the innate and adaptive immune systems and describes how adaptive immunity is recruited to the defense of the host by innate immune responses. An appreciation of the cellular and molecular bases of innate and adaptive immune responses is critical to understanding the pathogenesis of inflammatory, autoimmune, infectious, and immunodeficiency diseases.

THE INNATE IMMUNE SYSTEM

All multicellular organisms, including humans, have developed the use of a limited number of surface and intracellular germline-encoded molecules that recognize large groups of pathogens. Because of the myriad human pathogens, host molecules of the human innate immune system sense “danger signals” and either recognize PAMPs, the common molecular structures shared by many pathogens, or recognize host cell molecules produced in response to infection such as heat shock proteins and fragments of the extracellular matrix. PAMPs must be conserved structures vital to pathogen virulence and survival, such as bacterial endotoxin, so that pathogens cannot mutate molecules of PAMPs to evade human innate immune responses. PRRs are host proteins of the innate immune system that recognize PAMPs as host danger signal molecules (Tables 372e-2 and 372e-3). Thus, recognition of pathogen molecules by hematopoietic and nonhematopoietic cell types leads to activation/production of the complement cascade, cytokines, and antimicrobial peptides as effector molecules. In addition, pathogen PAMPs as host danger signal molecules activate dendritic cells to mature and to express molecules on the dendritic cell surface that optimize antigen presentation to respond to foreign antigens.

PATTERN RECOGNITION

Major PRR families of proteins include transmembrane proteins, such as the Toll-like receptors (TLRs) and C-type lectin receptors (CLRs), and cytoplasmic proteins, such as the retinoic acid-inducible gene (RIG)-1-like receptors (RLRs) and NOD-like receptors (NLRs) (Table 372e-3). A major group of PRR collagenous glycoproteins with C-type lectin domains are termed *collectins* and include the serum protein mannose-binding lectin (MBL). MBL and other collectins, as well as two other protein families—the pentraxins (such as C-reactive protein and serum amyloid P) and macrophage scavenger receptors—all have the property of opsonizing (coating) bacteria for phagocytosis by macrophages and can also activate the complement cascade to lyse bacteria. Integrins are cell-surface adhesion molecules that affect attachment between cells and the extracellular matrix and mediate signal transduction that reflects the chemical composition of the cell environment. For example, integrins signal after cells bind bacterial lipopolysaccharide (LPS) and activate phagocytic cells to ingest pathogens.

There are multiple connections between the innate and adaptive immune systems; these include (1) a plasma protein, LPS-binding protein, that binds and transfers LPS to the macrophage LPS receptor, CD14; (2) a human family of proteins called *Toll-like receptor proteins* (TLRs), some of which are associated with CD14, bind LPS, and signal epithelial cells, dendritic cells, and macrophages to produce cytokines and upregulate cell-surface molecules that signal the initiation of adaptive immune responses (Fig. 372e-1, Tables 372e-3 and 372e-4), and (3) families of intracellular microbial sensors called NLRs and RLRs. Proteins in the Toll family can be expressed on macrophages, dendritic cells, and B cells as well as on a variety of nonhematopoietic cell types, including respiratory epithelial cells. Eleven TLRs have been identified in humans, and 13 TLRs have been identified in mice (Tables 372e-4 and 372e-5). Upon ligation, TLRs activate a series of intracellular events that lead to the killing of bacteria- and viral-infected cells as well as to the recruitment and ultimate activation of antigen-specific T and B lymphocytes (Fig. 372e-1). Importantly, signaling by massive

TABLE 372e-1 HUMAN LEUKOCYTE SURFACE ANTIGENS—THE CD CLASSIFICATION OF LEUKOCYTE DIFFERENTIATION ANTIGENS

Surface Antigen (Other Names)	Family	Molecular Mass, kDa	Distribution	Ligand(s)	Function
CD1a (T6, HTA-1)	Ig	49	CD, cortical thymocytes, Langerhans type of DCs	TCR $\gamma\delta$ T cells	CD1 molecules present lipid antigens of intracellular bacteria such as <i>Mycobacterium leprae</i> and <i>M. tuberculosis</i> to TCR $\gamma\delta$ T cells
CD1b	Ig	45	CD, cortical thymocytes, Langerhans type of DCs	TCR $\gamma\delta$ T cells	
CD1c	Ig	43	DC, cortical thymocytes, subset of B cells, Langerhans type of DCs	TCR $\gamma\delta$ T cells	
CD1d	Ig	37	Cortical thymocytes, intestinal epithelium, Langerhans type of DCs	TCR $\gamma\delta$ T cells	
CD2 (T12, LFA-2)	Ig	50	T, NK	CD58, CD48, CD59, CD15	Alternative T cell activation, T cell anergy, T cell cytokine production, T- or NK-mediated cytotoxicity, T cell apoptosis, cell adhesion
CD3 (T3, Leu-4)	Ig	γ :25–28, δ :21–28, ϵ :20–25, η :21–22, ζ :16	T	Associates with the TCR	T cell activation and function; ζ is the signal transduction component of the CD3 complex
CD4 (T4, Leu-3)	Ig	55	T, myeloid	MHC-II, HIV, gp120, IL-16, SABP	T cell selection, T cell activation, signal transduction with p56/lck, primary receptor for HIV
CD7 (3A1, Leu-9)	Ig	40	T, NK	K-12 (CD7L)	T and NK cell signal transduction and regulation of IFN- γ , TNF- α production
CD8 (T8, Leu-2)	Ig	34	T	MHC-I	T cell selection, T cell activation, signal transduction with p56/lck
CD14 (LPS-receptor)	LRG	53–55	M, G (weak), not by myeloid progenitors	Endotoxin (lipopolysaccharide), lipoteichoic acid, PI	TLR4 mediates with LPS and other PAMP activation of innate immunity
CD16 (FcyRIIIa)	Ig	50–80	NK, macrophages, neutrophils	Fc portion of IgG	Mediates phagocytosis and ADCC
CD19 B4	Ig	95	B (except plasma cells), FDC	Not known	Associates with CD21 and CD81 to form a complex involved in signal transduction in B cell development, activation, and differentiation
CD20 (B1)	Unassigned	33–37	B (except plasma cells)	Not known	Cell signaling, may be important for B cell activation and proliferation
CD21 (B2, CR2, EBV-R, C3dR)	RCA	145	Mature B, FDC, subset of thymocytes	C3d, C3dg, iC3b, CD23, EBV	Associates with CD19 and CD81 to form a complex involved in signal transduction in B cell development, activation, and differentiation; Epstein-Barr virus receptor
CD22 (BL-CAM)	Ig	130–140	Mature B	CDw75	Cell adhesion, signaling through association with p72/sky, p53/56lyn, PI3 kinase, SHP1, fLc γ
CD23 (Fc ϵ RII, B6, Leu-20, BLAST-2)	C-type lectin	45	B, M, FDC	IgE, CD21, CD11b, CD11c	Regulates IgE synthesis, cytokine release by monocytes
CD28	Ig	44	T, plasma cells	CD80, CD86	Co-stimulatory for T cell activation; involved in the decision between T cell activation and anergy
CD32 (FcyRII)	Ig	40	NK, macrophages, neutrophils	Fc portion of IgG	Mediates phagocytosis and ADCC
CD40	TNFR	48–50	B, DC, EC, thymic epithelium, MP, cancers	CD154	B cell activation, proliferation, and differentiation; formation of GCs; isotype switching; rescue from apoptosis
CD45 (LCA, T200, B220)	PTP	180, 200, 210, 220	All leukocytes	Galectin-1, CD2, CD3, CD4	T and B activation, thymocyte development, signal transduction, apoptosis
CD45RA	PTP	210, 220	Subset T, medullary thymocytes, "naive" T	Galectin-1, CD2, CD3, CD4	Isoforms of CD45 containing exon 4 (A), restricted to a subset of T cells
CD45RB	PTP	200, 210, 220	All leukocytes	Galectin-1, CD2, CD3, CD4	Isoforms of CD45 containing exon 5 (B)
CD45RC	PTP	210, 220	Subset T, medullary thymocytes, "naive" T	Galectin-1, CD2, CD3, CD4	Isoforms of CD45 containing exon 6 (C), restricted to a subset of T cells
CD45RO	PTP	180	Subset T, cortical thymocytes, "memory" T	Galectin-1, CD2, CD3, CD4	Isoforms of CD45 containing no differentially spliced exons, restricted to a subset of T cells

(Continued)

TABLE 372e-1 HUMAN LEUKOCYTE SURFACE ANTIGENS—THE CD CLASSIFICATION OF LEUKOCYTE DIFFERENTIATION ANTIGENS (CONTINUED)

Surface Antigen (Other Names)	Family	Molecular Mass, kDa	Distribution	Ligand(s)	Function
CD64 (FcγRI)	Ig	45–55	Macrophages and monocytes	Fc portion of IgG	Mediates phagocytosis and ADCC
CD80 (B7-1, BB1)	Ig	60	Activated B and T, MP, DC	CD28, CD152	Co-regulator of T cell activation; signaling through CD28 stimulates and through CD152 inhibits T cell activation
CD86 (B7-2, B70)	Ig	80	Subset B, DC, EC, activated T, thymic epithelium	CD28, CD152	Co-regulator of T cell activation; signaling through CD28 stimulates and through CD152 inhibits T cell activation
CD89 (FCαR)	Ig	55–100	Neutrophils, eosinophils, monocytes, and MP	Fc portion of IgG	Mediates phagocytosis and ADCC of IgA-coated pathogens
CD95 (APO-1, Fas)	TNFR	43	Activated T and B	Fas ligand	Mediates apoptosis
CD152 (CTLA-4)	Ig	30–33	Activated T	CD80, CD86	Inhibits T cell proliferation
CD154 (CD40L)	TNF	33	Activated CD4+ T, subset CD8+ T, NK, M, basophil	CD40	Co-stimulatory for T cell activation, B cell proliferation and differentiation
CD279 (PD-1)	Ig	50–55	B, T, Tfh	PD-L1, PD-L2	Inhibits T cell proliferation

Abbreviations: ADCC, antibody-dependent cell-mediated cytotoxicity; CTLA, cytotoxic T lymphocyte-associated protein; DC, dendritic cells; EBV, Epstein-Barr virus; EC, endothelial cells; ECM, extracellular matrix; FcγRIII, low-affinity IgG receptor isoform A; FDC, follicular dendritic cells; G, granulocytes; GC, germinal center; GPI, glycosyl phosphatidylinositol; HTA, human thymocyte antigen; Ig, immunoglobulin; IgG, immunoglobulin G; LCA, leukocyte common antigen; LPS, lipopolysaccharide; MHC-I, major histocompatibility complex class I; MP, macrophages; Mr, relative molecular mass; NK, natural killer cells; P, platelets; PBT, peripheral blood T cells; PD-1, programmed cell death-1; PI, phosphatidylinositol; PI3K, phosphatidylinositol 3-kinase; PLC, phospholipase C; PTP, protein tyrosine phosphatase; TCR, T cell receptor; Tfh, T follicular helper cells; TNF, tumor necrosis factor; TNFR, tumor necrosis factor receptor. For an expanded list of cluster of differentiation (CD) human antigens, see Harrison's Online at <http://www.accessmedicine.com>; and for a full list of CD human antigens from the most recent Human Workshop on Leukocyte Differentiation Antigens (VII), see <http://mpr.nci.nih.gov/prowl/>.

Source: Compiled from T Kishimoto et al (eds): *Leukocyte Typing VI*. New York: Garland Publishing, 1997; R Brines et al: *Immunology Today* 18S:1, 1997; and S Shaw (ed): Protein reviews on the Web. <http://mpr.nci.nih.gov/prowl/>.

TABLE 372e-2 MAJOR COMPONENTS OF THE INNATE IMMUNE SYSTEM

Pattern recognition receptors (PRRs)	Toll-like receptors (TLRs), C-type lectin receptors (CLRs), retinoic acid-inducible gene (RIG)-1-like receptors (RLRs), and NOD-like receptors (NLRs)
Antimicrobial peptides	α-Defensins, β-defensins, cathelin, protegrin, granulysin, histatin, secretory leukoprotease inhibitor, and probiotics
Cells	Macrophages, dendritic cells, NK cells, NK-T cells, neutrophils, eosinophils, mast cells, basophils, and epithelial cells
Complement components	Classic and alternative complement pathway, and proteins that bind complement components
Cytokines	Autocrine, paracrine, endocrine cytokines that mediate host defense and inflammation, as well as recruit, direct, and regulate adaptive immune responses

Abbreviation: NK, natural killer.

amounts of LPS through TLR4 leads to the release of large amounts of cytokines that mediate LPS-induced shock. Mutations in TLR4 proteins in mice protect from LPS shock, and TLR mutations in humans protect from LPS-induced inflammatory diseases such as LPS-induced asthma (Fig. 372e-1).

Two other families of cytoplasmic PRRs are the NLRs and the RLRs. These families, unlike the TLRs, are composed primarily of soluble intracellular proteins that scan host cell cytoplasm for intracellular pathogens (Tables 372e-2 and 372e-3).

The intracellular microbial sensors, NLRs, after triggering, form large cytoplasmic complexes termed *inflammasomes*, which are aggregates of molecules including NOD-like receptor pyrin (NLRP) proteins that are members of the NLR family (Table 372e-3). Inflammasomes activate inflammatory caspases and IL-1β in the presence of nonbacterial danger signals (cell stress) and bacterial PAMPs. Mutations in inflammasome proteins can lead to chronic inflammation in a group of periodic febrile diseases called *autoinflammatory syndromes* (Table 372e-6).

EFFECTOR CELLS OF INNATE IMMUNITY

Cells of the innate immune system and their roles in the first line of host defense are listed in Table 372e-5. Equally important as their roles in the mediation of innate immune responses are the roles that each cell type plays in recruiting T and B lymphocytes of the adaptive immune system to engage in specific pathogen responses.

Monocytes-Macrophages Monocytes arise from precursor cells within bone marrow (Fig. 372e-2) and circulate with a half-life ranging from 1 to 3 days. Monocytes leave the peripheral circulation via capillaries and migration into a vast extravascular cellular pool. Tissue macrophages arise from monocytes that have migrated out of the circulation and by in situ proliferation of macrophage precursors in tissue. Common locations where tissue macrophages (and certain of their specialized forms) are found are lymph node, spleen, bone marrow, perivascular connective tissue, serous cavities such as the peritoneum, pleura, skin connective tissue, lung (alveolar macrophages), liver (Kupffer cells), bone (osteoclasts), central nervous system (microglia cells), and synovium (type A lining cells).

In general, monocytes-macrophages are on the first line of defense associated with innate immunity and ingest and destroy microorganisms through the release of toxic products such as hydrogen peroxide (H₂O₂) and nitric oxide (NO). Inflammatory mediators produced by macrophages attract additional effector cells such as neutrophils to the site of infection. Macrophage mediators include prostaglandins; leukotrienes; platelet activating factor; cytokines such as IL-1, TNF-α, IL-6, and IL-12; and chemokines (Tables 372e-7 to 372e-9).

Although monocytes-macrophages were originally thought to be the major antigen-presenting cells (APCs) of the immune system, it is now clear that cell types called *dendritic cells* are the most potent and effective APCs in the body (see below). Monocytes-macrophages mediate innate immune effector functions such as destruction of antibody-coated bacteria, tumor cells, or even normal hematopoietic cells in certain types of autoimmune cytopenias. Monocytes-macrophages ingest bacteria or are infected by viruses, and in doing so, they frequently undergo programmed cell death or *apoptosis*. Macrophages that are infected by intracellular infectious agents are recognized by dendritic cells as infected and apoptotic cells and are phagocytosed by dendritic cells. In this manner, dendritic cells “cross-present” infectious agent antigens of macrophages

TABLE 372e-3 PATTERN RECOGNITION RECEPTORS (PRRs) AND THEIR LIGANDS

PRRs	Localization	Ligand	Origin of the Ligand
TLR			
TLR1	Plasma membrane	Triacyl lipoprotein	Bacteria
TLR2	Plasma membrane	Lipoprotein	Bacteria, viruses, parasite, self
TLR3	Endolysosome	dsRNA	Virus
TLR4	Plasma membrane	LPS	Bacteria, viruses, self
TLR5	Plasma membrane	Flagellin	Bacteria
TLR6	Plasma membrane	Diacyl lipoprotein	Bacteria, viruses
TLR7 (human TLR8)	Endolysosome	ssRNA	Virus, bacteria, self
TLR9	Endolysosome	CpG-DNA	Virus, bacteria, protozoa, self
TLR10	Endolysosome	Unknown	Unknown
TLR11	Plasma membrane	Profilin-like molecule	Protozoa
RLR			
RIG-I	Cytoplasm	Short dsRNA, triphosphate dsRNA	RNA viruses, DNA virus
MDA5	Cytoplasm	Long dsRNA	RNA viruses (Picornaviridae)
LGP2	Cytoplasm	Unknown	RNA viruses
NLR			
NOD1	Cytoplasm	iE-DAP	Bacteria
NOD2	Cytoplasm	MDP	Bacteria
CLR			
Dectin-1	Plasma membrane	β^2 -Glucan	Fungi
Dectin-2	Plasma membrane	β^2 -Glucan	Fungi
MINCLE	Plasma membrane	SAP130	Self, fungi

Abbreviations: CLR, C-type lectin receptors; dsRNA, double-strand RNA; iE-DAP, D-glutamyl-meso-diaminopimelic acid moiety; LGP2, Laboratory of Genetics and Physiology 2 protein encoded by the gene *DHX58*; MDA5, melanoma differentiation-associated protein 5; MDP, MurNAc-L-Ala-D-isoGln, also known as muramyl dipeptide; MINCLE, macrophage-inducible C-type lectin; NLR, NOD-like receptor; NOD, NOTCH protein domain; RIG, retinoic acid-inducible gene; RLR, RIG-like receptors; TLR, Toll-like receptor.

Source: Adapted from O Takeuchi, S Akira: *Cell* 140:805, 2010, with permission.

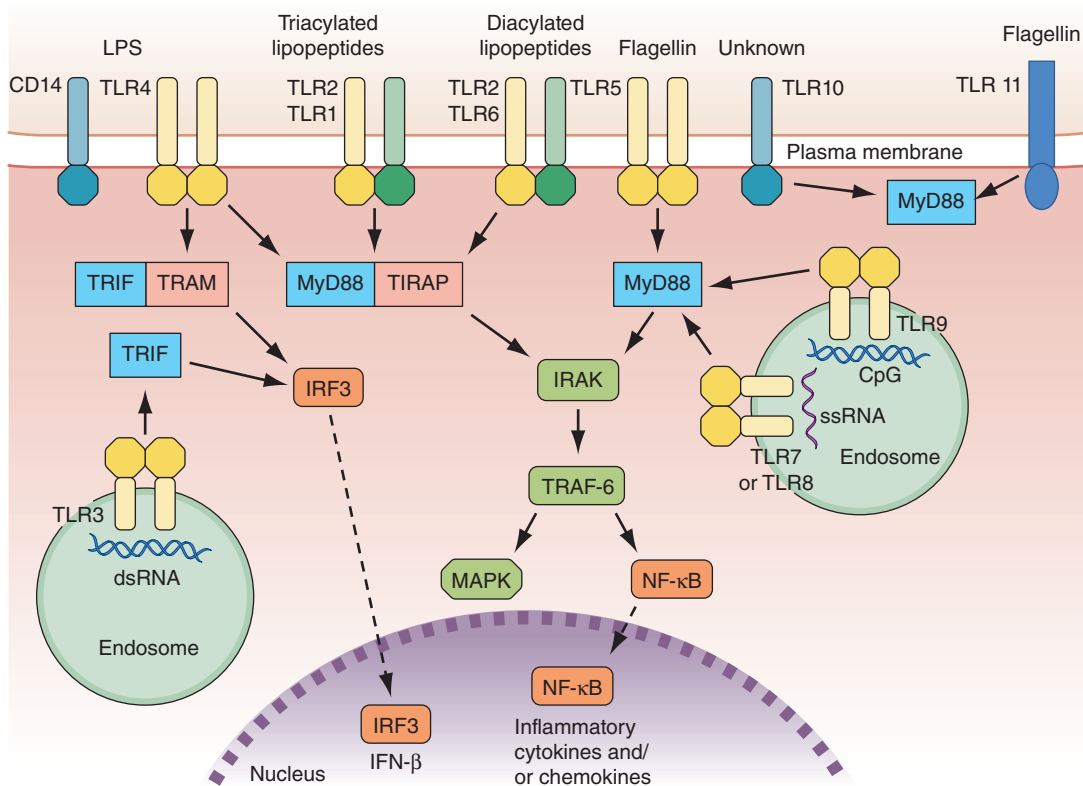


FIGURE 372e-1 Overview of major TLR signaling pathways. All TLRs signal through MyD88, with the exception of TLR3. TLR4 and the TLR2 subfamily (TLR1, TLR2, TLR6) also engage TIRAP. TLR3 signals through TRIF. TRIF is also used in conjunction with TRAM in the TLR4-MyD88-independent pathway. *Dashed arrows* indicate translocation into the nucleus. dsRNA, double-strand RNA; IFN, interferon; IRF3, interferon regulatory factor 3; LPS, lipopolysaccharide; MAPK, mitogen-activated protein kinases; NF- κ B, nuclear factor- κ B; ssRNA, single-strand RNA; TLR, Toll-like receptor. (Adapted from D van Duin et al: *Trends Immunol* 27:49, 2006, with permission.)

TABLE 372e-4 THE ROLE OF PATTERN RECOGNITION RECEPTORS (PRRs) IN MODULATION OF ADAPTIVE IMMUNE RESPONSES

PRR Family	PRRs	Ligand	DC or Macrophage Cytokine Response	Adaptive Immune Response
TLRs	TLR2 (heterodimer with TLR1 or 6)	Lipopeptides	Low IL-12p70	T _H 1
		Pam-3-cys (TLR 2/1) MALP (TLR 2/6)	High IL-10 IL-6	T _H 2 T regulatory
	TLR3	dsRNA	IL-12p70 IFN- α IL-6	T _H 1
	TLR4	<i>Escherichia coli</i> LPS	High IL-12p70 Intermediate IL-10 IL-6	T _H 1
	TLR5	Flagellin	High IL-12p70 Low IL-12p70	T _H 1 T _H 2
	TLR7/8	ssRNA Imidazoquinolines	High IL-12p70 IFN- α IL-6	T _H 1
	TLR9	CpG DNA	High IL-12p70 Low IL-10 IL-6 IFN- α	T _H 1
	TLR10	?	?	?
	TLR11	Profilin-like molecule Uropathogenic bacteria	?	?
C-type lectins	DC-SIGN	Env of HIV; core protein of HCV; components of <i>Mycobacterium tuberculosis</i> ; <i>Helicobacter pylori</i> , Lewis Ag	<i>H. pylori</i> , Lewis Ag Suppresses IL-12p70 Suppresses TLR signaling in DCs	T _H 2 T regulatory
NOD	NOD2	Muramyl dipeptide of peptidoglycan of bacteria	Induces IL-10 in DCs	Weak T cell response (tolerogenic?)
Mannose receptor	Mannose receptor	Mannosylated lipoarabinomannans from bacillus Calmette-Guérin and <i>M. tuberculosis</i>	Suppresses IL-12 and TLR signaling in DCs	Weak T cell response? (tolerogenic?)

Abbreviations: CpG, sequences in DNA recognized by TLR-9; DC, dendritic cell; DC-SIGN, DC-specific C-type lectin; dsRNA, double-strand RNA; HCV, hepatitis C; HIV, human immunodeficiency virus; LPS, lipopolysaccharide; MALP, macrophage-activating lipopeptide; NOD, NOTCH protein domain; ssRNA, single-strand RNA; T_H1, helper T cell; T_H2, helper T cell; TLR, Toll-like receptor.

Source: B Pulendran: *J Immunol* 174:2457, 2005. Copyright 2005 The American Association of Immunologists, Inc., with permission.

to T cells. Activated macrophages can also mediate antigen-nonspecific lytic activity and eliminate cell types such as tumor cells in the absence of antibody. This activity is largely mediated by cytokines (i.e., TNF- α and IL-1). Monocytes-macrophages express lineage-specific molecules (e.g., the cell-surface LPS receptor, CD14) as well as surface receptors for a number of molecules, including the Fc region of IgG, activated complement components, and various cytokines (Table 372e-7).

Dendritic Cells Human dendritic cells (DCs) contain several subsets, including myeloid DCs and plasmacytoid DCs. Myeloid DCs can differentiate into either macrophages-monocytes or tissue-specific DCs. In contrast to myeloid DCs, plasmacytoid DCs are inefficient APCs but are potent producers of type I interferon (IFN) (e.g., IFN- α) in response to viral infections. The maturation of DCs is regulated through cell-to-cell contact and soluble factors, and DCs attract immune effectors through secretion of chemokines. When DCs come in contact with bacterial products, viral proteins, or host proteins released as danger signals from distressed host cells (Figs. 372e-2 and 372e-3), infectious agent molecules bind to various TLRs and activate DCs to release cytokines and chemokines that drive cells of the innate immune system to become activated to respond to the invading organism, and recruit T and B cells of the adaptive immune system to respond. Plasmacytoid DCs produce antiviral IFN- α that activates NK cell killing of pathogen-infected cells; IFN- α also activates T cells to mature into antipathogen cytotoxic (killer) T cells. Following contact with pathogens, both plasmacytoid and myeloid DCs produce chemokines that attract helper and cytotoxic T cells, B cells, polymorphonuclear cells, and naïve and memory T cells as well as regulatory T cells to ultimately dampen the immune response once the pathogen is controlled. TLR engagement on

DCs upregulates MHC class II, B7-1 (CD80), and B7-2 (CD86), which enhance DC-specific antigen presentation and induce cytokine production (Table 372e-7). Thus, DCs are important bridges between early (innate) and later (adaptive) immunity. DCs also modulate and determine the types of immune responses induced by pathogens via the TLRs expressed on DCs (TLR7–9 on plasmacytoid DCs, TLR4 on monocytoïd DCs) and via the TLR adapter proteins that are induced to associate with TLRs (Fig. 372e-1, Table 372e-4). In addition, other PRRs, such as C-type lectins, NLRs, and mannose receptors, upon ligation by pathogen products, activate cells of the adaptive immune system and, like TLR stimulation, by a variety of factors, determine the type and quality of the adaptive immune response that is triggered (Table 372e-4).

Large Granular Lymphocytes/Natural Killer Cells Large granular lymphocytes (LGLs) or NK cells account for ~5–15% of peripheral blood lymphocytes. NK cells are nonadherent, nonphagocytic cells with large azurophilic cytoplasmic granules. NK cells express surface receptors for the Fc portion of IgG (FcR) (CD16) and for NCAM-I (CD56), and many NK cells express T lineage markers, particularly CD8, and proliferate in response to IL-2. NK cells arise in both bone marrow and thymic microenvironments.

Functionally, NK cells share features with both monocytes-macrophages and neutrophils in that they mediate both ADCC and NK cell activity. ADCC is the binding of an opsonized (antibody-coated) target cell to an Fc receptor-bearing effector cell via the Fc region of antibody, resulting in lysis of the target by the effector cell. NK cell cytotoxicity is the nonimmune (i.e., effector cell never having had previous contact with the target), MHC-unrestricted, non-antibody-mediated killing of target cells, which are usually malignant cell types, transplanted foreign cells, or virus-infected cells. Thus, NK cell cytotoxicity may play

TABLE 372e-5 CELLS OF THE INNATE IMMUNE SYSTEM AND THEIR MAJOR ROLES IN TRIGGERING ADAPTIVE IMMUNITY

Cell Type	Major Role in Innate Immunity	Major Role in Adaptive Immunity
Macrophages	Phagocytose and kill bacteria; produce antimicrobial peptides; bind LPS; produce inflammatory cytokines	Produce IL-1 and TNF- α to upregulate lymphocyte adhesion molecules and chemokines to attract antigen-specific lymphocyte. Produce IL-12 to recruit T _H 1 T helper cell responses; upregulate co-stimulatory and MHC molecules to facilitate T and B lymphocyte recognition and activation. Macrophages and dendritic cells, after LPS signaling, upregulate co-stimulatory molecules B7-1 (CD80) and B7-2 (CD86) that are required for activation of pathogen-specific T cells. There are also Toll-like proteins on B cells and dendritic cells that, after LPS ligation, induce CD80 and CD86 on these cells for T cell antigen presentation.
Plasmacytoid dendritic cells (DCs) of lymphoid lineage	Produce large amounts of interferon- α (IFN- α), which has antitumor and antiviral activity, and are found in T cell zones of lymphoid organs; they circulate in blood	IFN- α is a potent activator of macrophage and mature DCs to phagocytose invading pathogens and present pathogen antigens to T and B cells
Myeloid dendritic cells are of two types: interstitial and Langerhans-derived	Interstitial DCs are strong producers of IL-12 and IL-10 and are located in T cell zones of lymphoid organs, circulate in blood, and are present in the interstices of the lung, heart, and kidney; Langerhans DCs are strong producers of IL-12; are located in T cell zones of lymph nodes, skin epithelia, and the thymic medulla; and circulate in blood	Interstitial DCs are potent activators of macrophage and mature DCs to phagocytose invading pathogens and present pathogen antigens to T and B cells
Natural killer (NK) cells	Kill foreign and host cells that have low levels of MHC+ self-peptides. Express NK receptors that inhibit NK function in the presence of high expression of self-MHC.	Produce TNF- α and IFN- γ , which recruit T _H 1 helper T cell responses
NK-T cells	Lymphocytes with both T cell and NK surface markers that recognize lipid antigens of intracellular bacteria such as <i>Mycobacterium tuberculosis</i> by CD1 molecules and kill host cells infected with intracellular bacteria.	Produce IL-4 to recruit T _H 2 helper T cell responses, IgG1 and IgE production
Neutrophils	Phagocytose and kill bacteria, produce antimicrobial peptides	Produce nitric oxide synthase and nitric oxide, which inhibit apoptosis in lymphocytes and can prolong adaptive immune responses
Eosinophils	Kill invading parasites	Produce IL-5, which recruits Ig-specific antibody responses
Mast cells and basophils	Release TNF- α , IL-6, and IFN- γ in response to a variety of bacterial PAMPs	Produce IL-4, which recruits T _H 2 helper T cell responses, and recruit IgG1- and IgE-specific antibody responses
Epithelial cells	Produce antimicrobial peptides; tissue-specific epithelia produce mediator of local innate immunity; e.g., lung epithelial cells produce surfactant proteins (proteins within the collectin family) that bind and promote clearance of lung-invading microbes	Produces TGF- β , which triggers IgA-specific antibody responses

Abbreviations: IL-4, IL-5, IL-6, IL-10, and IL-12, interleukin 4, 5, 6, 10, and 12, respectively; MHC, major histocompatibility complex; LPS, lipopolysaccharide; PAMP, pathogen-associated molecular patterns; T_H, helper T cell; TNF- α , tumor necrosis factor- α .

Source: Adapted from R Medzhitov, CA Janeway: *Curr Opin Immunol* 9:4, 1997. Copyright 1997, with permission from Elsevier.

an important role in immune surveillance and destruction of malignant and virus-infected host cells. NK cell hyporesponsiveness is also observed in patients with *Chédiak-Higashi syndrome*, an autosomal recessive disease associated with fusion of cytoplasmic granules and defective degranulation of neutrophil lysosomes.

NK cells have a variety of surface receptors that have inhibitory or activating functions and belong to two structural families. These families include the immunoglobulin superfamily and the lectin-like type II transmembrane proteins. NK immunoglobulin superfamily receptors include the killer cell immunoglobulin-like activating or inhibitory receptors (KIRs), many of which have been shown to have HLA class I ligands. The KIRs are made up proteins with either two (KIR2D) or three (KIR3D) extracellular immunoglobulin domains (D). Moreover, their nomenclature designates their function as either inhibitory KIRs with a long (L) cytoplasmic tail and immunoreceptor tyrosine-based inhibitory motif (ITIM) (KIRDL) or activating KIRs with a short (S) cytoplasmic tail (KIRDS). NK cell inactivation by KIRs is a central mechanism to prevent damage to normal host cells. Genetic studies have demonstrated the association of KIRs with viral infection outcome and autoimmune disease (**Table 372e-10**).

In addition to the KIRs, a second set of immunoglobulin superfamily receptors includes the natural cytotoxicity receptors (NCRs), which include NKp46, NKp30, and NKp44. These receptors help to mediate NK cell activation against target cells. The ligands to which NCRs bind on target cells have been recently recognized to be comprised of molecules of pathogens such as influenza, vaccinia, and malaria as well as host molecules expressed on tumor cells.

NK cell signaling is, therefore, a highly coordinated series of inhibiting and activating signals that prevent NK cells from responding

to uninfected, nonmalignant self-cells; however, they are activated to attack malignant and virally infected cells (**Fig. 372e-4**). Recent evidence suggests that NK cells, although not possessing rearranging immune recognition genes, may be able to mediate recall for NK cell responses to viruses and for immune responses such as contact hypersensitivity.

Some NK cells express CD3 and invariant TCR- α chains and are termed *NK T cells*. TCRs of NK T cells recognize lipid molecules of intracellular bacteria when presented in the context of CD1d molecules on APCs. Upon activation, NK T cells secrete effector cytokines such as IL-4 and IFN- γ . This mode of recognition of intracellular bacteria such as *Listeria monocytogenes* and *Mycobacterium tuberculosis* by NK T cells leads to induction of activation of DCs and is thought to be an important innate defense mechanism against these organisms.

The receptors for the Fc portion of IgG (Fc γ Rs) are present on NK cells, B cells, macrophages, neutrophils, and mast cells and mediate interactions of IgG with antibody-coated target cells, such as virally infected cells. Antibody-NK interaction via antibody Fc and NK cell FcR links the adaptive and innate immune systems and regulates the mediation of IgG antibody effector functions such as ADCC. There are both activation and inhibitory Fc γ Rs. Activation FcRs, such as Fc γ RI (CD64), Fc γ RII (CD32), and Fc γ RIII (CD64), are characterized by the presence of an immunoreceptor tyrosine-based activating motif (ITAM) sequence, whereas inhibitory FcRs, such as Fc γ RIIb, contain an immunoreceptor tyrosine-based inhibitory motif (ITIM) sequence. There is evidence that dysregulation in IgG-Fc γ R interactions plays a role in arthritis, multiple sclerosis, and systemic lupus erythematosus.

TABLE 372e-6 DISEASES ASSOCIATED WITH INFLAMMASOME ACTIVITY

Disease	Clinical Features	Gene Mutated	Etiologic Agent	Inflammasome Involvement	Anakinra Response ^a
Familial cold autoinflammatory syndrome (FCAS)	Fever, arthralgia, cold-induced urticaria	<i>NALP3</i>		Overactive	Yes
Muckle-Wells syndrome (MWS)	Fever, arthralgia, urticaria, sensorineural deafness, amyloidosis	<i>NAPL3</i>		Overactive	Yes
Chronic infantile neurologic cutaneous and articular syndrome (CINCA, NOMID)	Fever, severe arthralgia, urticaria, neurologic problems, severe amyloidosis	<i>NALP3</i>		Overactive	Yes
Familial Mediterranean fever (FMF)	Fever, peritonitis, pleuritis, amyloidosis	Pyrin		Overactive	Partial
Pyogenic arthritis, pyoderma gangrenosum, and acne syndrome (PAPA)	Pyogenic sterile arthritis	<i>PSTPIP1</i>		Overactive	Yes
Hyperimmunoglobulin D syndrome (HIDS)	Arthralgia, abdominal pain, lymphadenopathy	Mevalonate kinase		To be demonstrated	Yes
Tumor necrosis factor receptor-1-associated syndrome (TRAPS)	Fever, abdominal pain, skin lesions	<i>TNF-R1</i>		To be demonstrated	Yes
Systemic-onset juvenile idiopathic arthritis (SOJIA)	Chronic joint inflammation		Unknown	To be demonstrated	Yes
Adult-onset Still's disease (AOSD)	Arthralgia, fever		Unknown	To be demonstrated	Yes
Behçet's disease	Arthralgia, uveitis, ulcers		Unknown	To be demonstrated	Yes
Schnitzler's syndrome	Urticaria, fever, arthralgia		Unknown	To be demonstrated	Yes
Gout	Metabolic arthritis, pain		Uric acid (MSU)	Activated	Yes
Pseudogout	Arthritis		CPPD	Activated	Yes
Contact dermatitis	Urticaria		Irritants	Activated	Unknown
Fever syndrome	Fever	<i>NALP12</i>		Unknown	Unknown
Hydatidiform mole	Hydatid mole	<i>NALP7</i>		Unknown	Unknown
Vitiligo	Skin depigmentation, autoimmunity	<i>NLRP1</i>		Overactive	Unknown
Crohn's disease		<i>NLRP3</i>		Underactive	Unknown
Multiple Sclerosis		<i>NLPR3</i>		Activated	Unknown
Psoriatic arthritis		<i>NLRP3</i>		Activated	Yes

^aAnakinra is a recombinant interleukin-1 (IL-1) receptor antagonist that functions to block the biologic activity of naturally occurring IL-1.

Abbreviation: CPPD, calcium pyrophosphate dehydrate.

Source: Adapted from F Martinon et al: *Ann Rev Immunol* 27:229, 2009. Copyright 2009. Reproduced with permission from Annual Reviews Inc.

Neutrophils, Eosinophils, and Basophils Granulocytes are present in nearly all forms of inflammation and are amplifiers and effectors of innate immune responses (Figs. 372e-2 and 372e-3). Unchecked accumulation and activation of granulocytes can lead to host tissue damage, as seen in neutrophil- and eosinophil-mediated *systemic necrotizing vasculitis*. Granulocytes are derived from stem cells in bone marrow. Each type of granulocyte (neutrophil, eosinophil, or basophil) is derived from a different subclass of progenitor cell that is stimulated to proliferate by colony-stimulating factors (Table 372e-7). During terminal maturation of granulocytes, class-specific nuclear morphology and cytoplasmic granules appear that allow for histologic identification of granulocyte type.

Neutrophils express Fc receptor IIIa for IgG (CD16) as well as receptors for activated complement components (C3b or CD35). Upon interaction of neutrophils with antibody-coated (opsonized) bacteria or immune complexes, azurophilic granules (containing myeloperoxidase, lysozyme, elastase, and other enzymes) and specific granules (containing lactoferrin, lysozyme, collagenase, and other enzymes) are released, and microbicidal superoxide radicals (O_2^-) are generated at the neutrophil surface. The generation of superoxide leads to inflammation by direct injury to tissue and by alteration of macromolecules such as collagen and DNA.

Eosinophils express Fc receptor II for IgG (CD32) and are potent cytotoxic effector cells for various parasitic organisms. In *Nippostrongylus*

brasiliensis helminth infection, eosinophils are important cytotoxic effector cells for removal of these parasites. Key to regulation of eosinophil cytotoxicity to *N. brasiliensis* worms are antigen-specific T helper cells that produce IL-4, thus providing an example of regulation of innate immune responses by adaptive immunity antigen-specific T cells. Intracytoplasmic contents of eosinophils, such as major basic protein, eosinophil cationic protein, and eosinophil-derived neurotoxin, are capable of directly damaging tissues and may be responsible in part for the organ system dysfunction in the *hypereosinophilic syndromes* (Chap. 80). Because the eosinophil granule contains anti-inflammatory types of enzymes (histaminase, arylsulfatase, phospholipase D), eosinophils may homeostatically downregulate or terminate ongoing inflammatory responses.

Basophils and tissue mast cells are potent reservoirs of cytokines such as IL-4 and can respond to bacteria and viruses with antipathogen cytokine production through multiple TLRs expressed on their surface. Mast cells and basophils can also mediate immunity through the binding of antipathogen antibodies. This is a particularly important host defense mechanism against parasitic diseases. Basophils express high-affinity surface receptors for IgE (FcεRI) (CD23) and, upon cross-linking of basophil-bound IgE by antigen, can release histamine, eosinophil chemotactic factor of anaphylaxis, and neutral protease—all mediators of allergic immediate (anaphylaxis) hypersensitivity responses (Table 372e-11). In addition, basophils express surface receptors for activated complement components (C3a, C5a), through

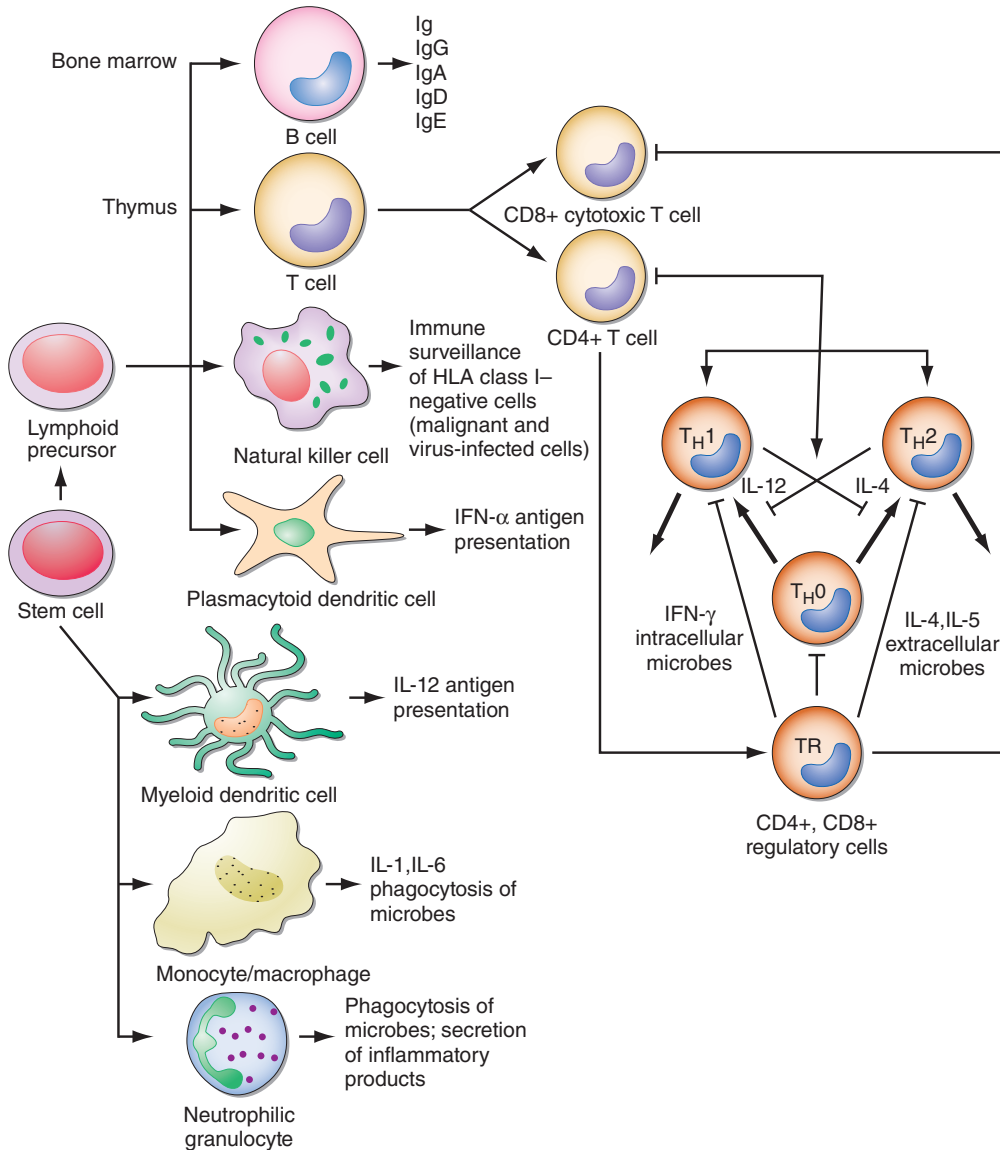


FIGURE 372e-2 Schematic model of intercellular interactions of adaptive immune system cells.

In this figure, the *arrows* denote that cells develop from precursor cells or produce cytokines or antibodies; *lines ending with bars* indicate suppressive intercellular interactions. Stem cells differentiate into either T cells, antigen-presenting dendritic cells, natural killer cells, macrophages, granulocytes, or B cells. Foreign antigen is processed by dendritic cells, and peptide fragments of foreign antigen are presented to CD4+ and/or CD8+ T cells. CD8+ T cell activation leads to induction of cytotoxic T lymphocyte (CTL) or killer T cell generation, as well as induction of cytokine-producing CD8+ cytotoxic T cells. For antibody production against the same antigen, active antigen is bound to sIg within the B cell receptor complex and drives B cell maturation into plasma cells that secrete Ig. T_H1 or T_H2 CD4+ T cells producing interleukin (IL) 4, IL-5, or interferon (IFN) γ regulate the Ig class switching and determine the type of antibody produced. T_H17 cells secrete IL-17, IL-22, IL-26, which contribute to host defense against extracellular bacteria and fungi, particularly at mucosal surfaces. CD4+, CD25+ T regulatory cells produce IL-10 and downregulate T and B cell responses once the microbe has been eliminated. GM-CSF, granulocyte-macrophage colony-stimulating factor; TNF, tumor necrosis factor.

which mediator release can be directly affected. Thus, basophils, like most cells of the immune system, can be activated in the service of host defense against pathogens, or they can be activated for mediation release and cause pathogenic responses in allergic and inflammatory diseases. **For further discussion of tissue mast cells, see Chap. 376.**

The Complement System The complement system, an important soluble component of the innate immune system, is a series of plasma enzymes, regulatory proteins, and proteins that are activated in a cascading fashion, resulting in cell lysis. There are four pathways of the complement system: the classic activation pathway activated by antigen/antibody immune complexes, the mannose-binding lectin (MBL) (a serum

collectin; Table 372e-3) activation pathway activated by microbes with terminal mannose groups, the alternative activation pathway activated by microbes or tumor cells, and the terminal pathway that is common to the first three pathways and leads to the membrane attack complex that lyses cells (Fig. 372e-5). The series of enzymes of the complement system are serine proteases.

Activation of the classic complement pathway via immune complex binding to C1q links the innate and adaptive immune systems via specific antibody in the immune complex. The alternative complement activation pathway is antibody-independent and is activated by binding of C3 directly to pathogens and “altered self” such as tumor cells. In the renal glomerular inflammatory disease *IgA nephropathy*, IgA activates the alternative complement pathway and causes glomerular damage and decreased renal function. Activation of the classic complement pathway via C1, C4, and C2 and activation of the alternative pathway via factor D, C3, and factor B both lead to cleavage and activation of C3. C3 activation fragments, when bound to target surfaces such as bacteria and other foreign antigens, are critical for opsonization (coating by antibody and complement) in preparation for phagocytosis. The MBL pathway substitutes MBL-associated serine proteases (MASPs) 1 and 2 for C1q, C1r, and C1s to activate C4. The MBL activation pathway is activated by mannose on the surface of bacteria and viruses.

The three pathways of complement activation all converge on the final common terminal pathway. C3 cleavage by each pathway results in activation of C5, C6, C7, C8, and C9, resulting in the membrane attack complex that physically inserts into the membranes of target cells or bacteria and lyses them.

Thus, complement activation is a critical component of innate immunity for responding to microbial infection. The functional consequences of complement activation by the three initiating pathways and

the terminal pathway are shown in Fig. 372e-5. In general the cleavage products of complement components facilitate microbe or damaged cell clearance (C1q, C4, C3), promote activation and enhancement of inflammation (anaphylatoxins, C3a, C5a), and promote microbe or opsonized cell lysis (membrane attack complex).

CYTOKINES

Cytokines are soluble proteins produced by a wide variety of cell types (Tables 372e-7 to 372e-9). They are critical for both normal innate and adaptive immune responses, and their expression may be perturbed in most immune, inflammatory, and infectious disease states.

TABLE 372e-7 CYTOKINES AND CYTOKINE RECEPTORS

Cytokine	Receptor	Cell Source	Cell Target	Biologic Activity
IL-1 α , β	Type I IL-1r, Type II IL-1r	Monocytes/macrophages, B cells, fibroblasts, most epithelial cells including thymic epithelium, endothelial cells	All cells	Upregulates adhesion molecule expression, neutrophil and macrophage emigration, mimics shock, fever, upregulates hepatic acute-phase protein production, facilitates hematopoiesis
IL-2	IL-2r α , β , common γ	T cells	T cells, B cells, NK cells, monocytes-macrophages	Promotes T cell activation and proliferation, B cell growth, NK cell proliferation and activation, enhanced monocyte/macrophage cytolytic activity
IL-3	IL-3r, common β	T cells, NK cells, mast cells	Monocytes-macrophages, mast cells, eosinophils, bone marrow progenitors	Stimulates hematopoietic progenitors
IL-4	IL-4r α , common γ	T cells, mast cells, basophils	T cells, B cells, NK cells, monocytes-macrophages, neutrophils, eosinophils, endothelial cells, fibroblasts	Stimulates T _H 2 helper T cell differentiation and proliferation; stimulates B cell Ig class switch to IgG1 and IgE anti-inflammatory action on T cells, monocytes; produced by T follicular helper cells in B cell germinal centers that stimulate B cell maturation.
IL-5	IL-5r α , common γ	T cells, mast cells, eosinophils	Eosinophils, basophils, murine B cells	Regulates eosinophil migration and activation
IL-6	IL-6r, gp130	Monocytes-macrophages, B cells, fibroblasts, most epithelium including thymic epithelium, endothelial cells	T cells, B cells, epithelial cells, hepatocytes, monocytes-macrophages	Induces acute-phase protein production, T and B cell differentiation and growth, myeloma cell growth, and osteoclast growth and activation
IL-7	IL-7r α , common γ	Bone marrow, thymic epithelial cells	T cells, B cells, bone marrow cells	Differentiates B, T, and NK cell precursors, activates T and NK cells
IL-8	CXCR1, CXCR2	Monocytes-macrophages, T cells, neutrophils, fibroblasts, endothelial cells, epithelial cells	Neutrophils, T cells, monocytes-macrophages, endothelial cells, basophils	Induces neutrophil, monocyte, and T cell migration, induces neutrophil adherence to endothelial cells and histamine release from basophils, and stimulates angiogenesis; suppresses proliferation of hepatic precursors
IL-9	IL-9r α , common γ	T cells	Bone marrow progenitors, B cells, T cells, mast cells	Induces mast cell proliferation and function, synergizes with IL-4 in IgG and IgE production and T cell growth, activation, and differentiation
IL-10	IL-10r	Monocytes-macrophages, T cells, B cells, keratinocytes, mast cells	Monocytes-macrophages, T cells, B cells, NK cells, mast cells	Inhibits macrophage proinflammatory cytokine production, downregulates cytokine class II antigen and B7-1 and B7-2 expression, inhibits differentiation of T _H 1 helper T cells, inhibits NK cell function, stimulates mast cell proliferation and function, B cell activation, and differentiation
IL-11	IL-11r α , gp130	Bone marrow stromal cells	Megakaryocytes, B cells, hepatocytes	Induces megakaryocyte colony formation and maturation, enhances antibody responses, stimulates acute-phase protein production
IL-12 (35-kDa and 40-kDa subunits)	IL-12r	Activated macrophages, dendritic cells, neutrophils	T cells, NK cells	Induces T _H 1 T helper cell formation and lymphokine-activated killer cell formation; increases CD8+ CTL cytolytic activity; \downarrow IL-17, \uparrow IFN- γ
IL-13	IL-13r/IL-4r α	T cells (T _H 2)	Monocytes-macrophages, B cells, endothelial cells, keratinocytes	Upregulates VCAM-1 and C-C chemokine expression on endothelial cells and B cell activation and differentiation, and inhibits macrophage proinflammatory cytokine production
IL-14	Unknown	T cells	Normal and malignant B cells	Induces B cell proliferation, inhibits antibody secretion, and expands selected B cell subgroups
IL-15	IL-15r α , common γ , IL2r β	Monocytes-macrophages, epithelial cells, fibroblasts	T cells, NK cells	Promotes T cell activation and proliferation, angiogenesis, and NK cells
IL-16	CD4	Mast cells, eosinophils, CD8+ T cells, respiratory epithelium	CD4+ T cells, monocytes-macrophages, eosinophils	Promotes chemoattraction of CD4+ T cells, monocytes, and eosinophils; inhibits HIV replication; inhibits T cell activation through CD3/T cell receptor
IL-17	IL-17r	CD4+ T cells	Fibroblasts, endothelium, epithelium, macrophages	Enhances cytokine/chemokine secretion; promotes delayed-type reactions
IL-18	IL-18r (IL-1R-related protein)	Keratinocytes, macrophages	T cells, B cells, NK cells	Upregulates IFN- γ production, enhances NK cell cytotoxicity
IL-21	IL- $\delta\gamma$ chain/IL-21R	CD4 T cells	NK cells	Downregulates NK cell-activating molecules, NKG2D/DAP10; produced by T follicular helper cells in B cell germinal centers that stimulate B cell maturation.
IL-22	IL-22 R1/IL-10R2	DC, T cells	Epithelial cells	Innate responses against bacterial pathogens; promotes hepatocyte survival
IL-23	IL-12R β 1/IL23R	Macrophages, other cell types	T cells	Opposite effects of IL-12 (\uparrow IL-17, \uparrow IFN- γ)

(Continued)

TABLE 372e-7 CYTOKINES AND CYTOKINE RECEPTORS (CONTINUED)

Cytokine	Receptor	Cell Source	Cell Target	Biologic Activity
IL-24	IL-20 R1/IL-20R2	Macrophages,	Nonhematopoietic cells such as fibroblasts	Promotes wound healing
	IL-22R1/IL-20 R2	T _H 2 cells		
IL-25 (also called IL-17E)	IL-17RB	CD4 T cells, mast cells	Fibroblasts, endothelium, epithelium, macrophages	Proinflammatory; induces cytokine production
IL-26	IL-20 R1/IL-10R2	T _H 1, T _H 17 T cells, synovial cells	Epithelial cells	Proinflammatory; induces cytokine production
IFN- α	Type I interferon receptor	All cells	All cells	Promotes antiviral activity; stimulates T cell, macrophage, and NK cell activity; direct antitumor effects; upregulates MHC class I antigen expression; used therapeutically in viral and autoimmune conditions
IFN- β	Type I interferon receptor	All cells	All cells	Antiviral activity; stimulates T cell, macrophage, and NK cell activity; direct antitumor effects; upregulates MHC class I antigen expression; used therapeutically in viral and autoimmune conditions
IFN- γ	Type II interferon receptor	T cells, NK cells	All cells	Regulates macrophage and NK cell activations; stimulates immunoglobulin secretion by B cells; induction of class II histocompatibility antigens; T _H 1 T cell differentiation
TNF- α	TNFR _I , TNFR _{II}	Monocytes-macrophages, mast cells, basophils, eosinophils, NK cells, B cells, T cells, keratinocytes, fibroblasts, thymic epithelial cells	All cells except erythrocytes	Fever, anorexia, shock, capillary leak syndrome, enhanced leukocyte cytotoxicity, enhanced NK cell function, acute phase protein synthesis, proinflammatory cytokine induction
TNF- β	TNFR _I , TNFR _{II}	T cells, B cells	All cells except erythrocytes	Cell cytotoxicity, lymph node and spleen development
LT- β	LT β R	T cells	All cells except erythrocytes	Cell cytotoxicity, normal lymph node development
G-CSF	G-CSFr; gp130	Monocytes-macrophages, fibroblasts, endothelial cells, thymic epithelial cells, stromal cells	Myeloid cells, endothelial cells	Regulates myelopoiesis; enhances survival and function of neutrophils; clinical use in reversing neutropenia after cytotoxic chemotherapy
GM-CSF	GM-CSFr, common β	T cells, monocytes-macrophages, fibroblasts, endothelial cells, thymic epithelial cells	Monocytes-macrophages, neutrophils, eosinophils, fibroblasts, endothelial cells	Regulates myelopoiesis; enhances macrophage bactericidal and tumoricidal activity; mediator of dendritic cell maturation and function; upregulates NK cell function; clinical use in reversing neutropenia after cytotoxic chemotherapy
M-CSF	M-CSFr (<i>c-fms</i> protooncogene)	Fibroblasts, endothelial cells, monocytes-macrophages, T cells, B cells, epithelial cells including thymic epithelium	Monocytes-macrophages	Regulates monocyte-macrophage production and function
LIF	LIFr- α ; gp130	Activated T cells, bone marrow stromal cells, thymic epithelium	Megakaryocytes, monocytes, hepatocytes, possibly lymphocyte subpopulations	Induces hepatic acute-phase protein production; stimulates macrophage differentiation; promotes growth of myeloma cells and hematopoietic progenitors; stimulates thrombopoiesis
OSM	OSMr; LIFr; gp130	Activated monocytes-macrophages and T cells, bone marrow stromal cells, some breast carcinoma cell lines, myeloma cells	Neurons, hepatocytes, monocytes-macrophages, adipocytes, alveolar epithelial cells, embryonic stem cells, melanocytes, endothelial cells, fibroblasts, myeloma cells	Induces hepatic acute-phase protein production; stimulates macrophage differentiation; promotes growth of myeloma cells and hematopoietic progenitors; stimulates thrombopoiesis; stimulates growth of Kaposi's sarcoma cells
SCF	SCFr (<i>c-kit</i> protooncogene)	Bone marrow stromal cells and fibroblasts	Embryonic stem cells, myeloid and lymphoid precursors, mast cells	Stimulates hematopoietic progenitor cell growth, mast cell growth; promotes embryonic stem cell migration
TGF- β (3 isoforms)	Type I, II, III TGF- β receptor	Most cell types	Most cell types	Downregulates T cell, macrophage, and granulocyte responses; stimulates synthesis of matrix proteins; stimulates angiogenesis
Lymphotoxin/SCM-1	XCR1	NK cells, mast cells, double-negative thymocytes, activated CD8+ T cells	T cells, NK cells	Chemoattractant for lymphocytes; only known chemokine of C class
MCP-1	CCR2	Fibroblasts, smooth-muscle cells, activated PBMCs	Monocytes-macrophages, NK cells, memory T cells, basophils	Chemoattractant for monocytes, activated memory T cells, and NK cells; induces granule release from CD8+ T cells and NK cells; potent histamine-releasing factor for basophils; suppresses proliferation of hematopoietic precursors; regulates monocyte protease production
MCP-2	CCR1, CCR2	Fibroblasts, activated PBMCs	Monocytes-macrophages, T cells, eosinophils, basophils, NK cells	Chemoattractant for monocytes, memory and naïve T cells, eosinophils, ?NK cells; activates basophils and eosinophils; regulates monocyte protease production

(Continued)

TABLE 372e-7 CYTOKINES AND CYTOKINE RECEPTORS (CONTINUED)

Cytokine	Receptor	Cell Source	Cell Target	Biologic Activity
MCP-3	CCR1, CCR2	Fibroblasts, activated PBMCs	Monocytes-macrophages, T cells, eosinophils, basophils, NK cells, dendritic cells	Chemoattractant for monocytes, memory and naïve T cells, dendritic cells, eosinophils, ?NK cells; activates basophils and eosinophils; regulates monocyte protease production
MCP-4	CCR2, CCR3	Lung, colon, small intestinal epithelial cells, activated endothelial cells	Monocytes-macrophages, T cells, eosinophils, basophils	Chemoattractant for monocytes, T cells, eosinophils, and basophils
Eotaxin	CCR3	Pulmonary epithelial cells, heart	Eosinophils, basophils	Potent chemoattractant for eosinophils and basophils; induces allergic airways disease; acts in concert with IL-5 to activate eosinophils; antibodies to eotaxin inhibit airway inflammation
TARC	CCR4	Thymus, dendritic cells, activated T cells	T cells, NK cells	Chemoattractant for T and NK cells
MDC	CCR4	Monocytes-macrophages, dendritic cells, thymus	Activated T cells	Chemoattractant for activated T cells; inhibits infection with T cell tropic HIV
MIP-1 α	CCR1, CCR5	Monocytes-macrophages, T cells	Monocytes-macrophages, T cells, dendritic cells, NK cells, eosinophils, basophils	Chemoattractant for monocytes, T cells, dendritic cells, and NK cells, and weak chemoattractant for eosinophils and basophils; activates NK cell function; suppresses proliferation of hematopoietic precursors; necessary for myocarditis associated with coxsackievirus infection; inhibits infection with monocyotropic HIV
MIP-1 β	CCR5	Monocytes-macrophages, T cells	Monocytes-macrophages, T cells, NK cells, dendritic cells	Chemoattractant for monocytes, T cells, and NK cells; activates NK cell function; inhibits infection with monocyotropic HIV
RANTES	CCR1, CCR2, CCR5	Monocytes-macrophages, T cells, fibroblasts, eosinophils	Monocytes-macrophages, T cells, NK cells, dendritic cells, eosinophils, basophils	Chemoattractant for monocytes-macrophages, CD4+, CD45Ro+ T cells, CD8+ T cells, NK cells, eosinophils, and basophils; induces histamine release from basophils; inhibits infections with monocyotropic HIV
LARC/MIP-3 α /Exodus-1	CCR6	Dendritic cells, fetal liver cells, activated T cells	T cells, B cells	Chemoattractant for lymphocytes
ELC/MIP-3 β	CCR7	Thymus, lymph node, appendix	Activated T cells and B cells	Chemoattractant for B and T cells; receptor upregulated on EBV-infected B cells and HSV-infected T cells
I-309/TCA-3	CCR8	Activated T cells	Monocytes-macrophages, T cells	Chemoattractant for monocytes; prevents glucocorticoid-induced apoptosis in some T cell lines
SLC/TCA-4/Exodus-2	CCR7	Thymic epithelial cells, lymph node, appendix, and spleen	T cells	Chemoattractant for T lymphocytes; inhibits hematopoiesis
DC-CK1/PARC	Unknown	Dendritic cells in secondary lymphoid tissues	Naïve T cells	May have a role in induction of immune responses
TECK	CCR9	Dendritic cells, thymus, liver, small intestine	T cells, monocytes-macrophages, dendritic cells	Thymic dendritic cell-derived cytokine, possibly involved in T cell development
GRO- α /MGSA	CXCR2	Activated granulocytes, monocyte-macrophages, and epithelial cells	Neutrophils, epithelial cells, ?endothelial cells	Neutrophil chemoattractant and activator; mitogenic for some melanoma cell lines; suppresses proliferation of hematopoietic precursors; angiogenic activity
GRO- β /MIP-2 α	CXCR2	Activated granulocytes and monocyte-macrophages	Neutrophils and ?endothelial cells	Neutrophil chemoattractant and activator; angiogenic activity
NAP-2	CXCR2	Platelets	Neutrophils, basophils	Derived from platelet basic protein; neutrophil chemoattractant and activator
IP-10	CXCR3	Monocytes-macrophages, T cells, fibroblasts, endothelial cells, epithelial cells	Activated T cells, tumor-infiltrating lymphocytes, ?endothelial cells, ?NK cells	IFN- γ -inducible protein that is a chemoattractant for T cells; suppresses proliferation of hematopoietic precursors
MIG	CXCR3	Monocytes-macrophages, T cells, fibroblasts	Activated T cells, tumor-infiltrating lymphocytes	IFN- γ -inducible protein that is a chemoattractant for T cells; suppresses proliferation of hematopoietic precursors
SDF-1	CXCR4	Fibroblasts	T cells, dendritic cells, ?basophils, ?endothelial cells	Low-potency, high-efficacy T cell chemoattractant; required for B-lymphocyte development; prevents infection of CD4+, CXCR4+ cells by T cell tropic HIV

(Continued)

TABLE 372e-7 CYTOKINES AND CYTOKINE RECEPTORS (CONTINUED)

Cytokine	Receptor	Cell Source	Cell Target	Biologic Activity
Fractalkine	CX3CR1	Activated endothelial cells	NK cells, T cells, monocytes-macrophages	Cell-surface chemokine/mucin hybrid molecule that functions as a chemoattractant, leukocyte activator, and cell adhesion molecule
PF-4	Unknown	Platelets, megakaryocytes	Fibroblasts, endothelial cells	Chemoattractant for fibroblasts; suppresses proliferation of hematopoietic precursors; inhibits endothelial cell proliferation and angiogenesis

Abbreviations: B7-1, CD80; B7-2, CD86; CCR, CC-type chemokine receptor; CXCR, CXC-type chemokine receptor; DC-CK, dendritic cell chemokine; EBV, Epstein-Barr virus; ELC, EB11 ligand chemokine (MIP-1b); G-CSF, granulocyte colony-stimulating factor; GM-CSF, granulocyte-macrophage colony-stimulating factor; GRP, growth-related peptide; HSV, herpes simplex virus; IFN, interferon; Ig, immunoglobulin; IL, interleukin; IP-10, IFN- γ -inducible protein-10; LARC, liver- and activation-regulated chemokine; LIF, leukemia inhibitory factor; MCP, monocyte chemotactic protein; M-CSF, macrophage colony-stimulating factor; MDC, macrophage-derived chemokine; MGSA, melanoma growth-stimulating activity; MHC, major histocompatibility complex; MIG, monokine induced by IFN- γ ; MIP, macrophage inflammatory protein; NAP, neutrophil-activating protein; NK, natural killer; OSM, oncostatin M; PARC, pulmonary- and activation-regulated chemokine; PBMC, peripheral blood mononuclear cells; PF, platelet factor; RANTES, regulated on activation, normally T cell-expressed and -secreted; SCF, stem cell factor; SDF, stromal cell-derived factor; SLC, secondary lymphoid tissue chemokine; TARC, thymus- and activation-regulated chemokine; TCA, T cell activation protein; TECK, thymus expressed chemokine; TGF, transforming growth factor; T_H1 and T_H2, helper T cell subsets; TNF, tumor necrosis factor; VCAM, vascular cell adhesion molecule.

Source: Data from JS Sundy et al: Appendix B, in *Inflammation, Basic Principles and Clinical Correlates*, 3rd ed, J Gallin and R Snyderman (eds). Philadelphia, Lippincott Williams and Wilkins, 1999.

TABLE 372e-8 CC, CXC, CX₃, C₁, AND XC FAMILIES OF CHEMOKINES AND CHEMOKINE RECEPTORS

Chemokine Receptor	Chemokine Ligands	Cell Types	Disease Connection
CCR1	CCL3 (MIP-1 α), CCL5 (RANTES), CCL7 (MCP-3), CCL14 (HCC1)	T cells, monocytes, eosinophils, basophils	Rheumatoid arthritis, multiple sclerosis
CCR2	CCL2 (MCP-1), CCL8 (MCP-2), CCL7 (MCP-3), CCL13 (MCP-4), CCL16 (HCC4)	Monocytes, dendritic cells (immature), memory T cells	Atherosclerosis, rheumatoid arthritis, multiple sclerosis, resistance to intracellular pathogens, type 2 diabetes mellitus
CCR3	CCL11 (eotaxin), CCL13 (eotaxin-2), CCL7 (MCP-3), CCL5 (RANTES), CCL8 (MCP-2), CCL13 (MCP-4)	Eosinophils, basophils, mast cells, T _H 2, platelets	Allergic asthma and rhinitis
CCR4	CCL17 (TARC), CCL22 (MDC)	T cells (T _H 2), dendritic cells (mature), basophils, macrophages, platelets	Parasitic infection, graft rejection, T cell homing to skin
CCR5	CCL3 (MIP-1 α), CCL4 (MIP-1 α), CCL5 (RANTES), CCL11 (eotaxin), CCL14 (HCC1), CCL16 (HCC4)	T cells, monocytes	HIV-1 co-receptor (T cell-tropic strains), transplant rejection
CCR6	CCL20 (MIP-3 α , LARC)	T cells (T regulatory and memory), B cells, dendritic cells	Mucosal humoral immunity, allergic asthma, intestinal T cell homing
CCR7	CCL19 (ELC), CCL21 (SLC)	T cells, dendritic cells (mature)	Transport of T cells and dendritic cells to lymph nodes, antigen presentation, and cellular immunity
CCR8	CCL1 (1309)	T cells (T _H 2), monocytes, dendritic cells	Dendritic cell migration to lymph node, type 2 cellular immunity, granuloma formation
CCR9	CCL25 (TECK)	T cells, IgA+ plasma cells	Homing of T cells and IgA+ plasma cells to the intestine, inflammatory bowel disease
CCR10	CCL27 (CTACK), CCL28 (MEC)	T cells	T cell homing to intestine and skin
CXCR1	CXCL8 (interleukin-8), CXCL6 (GCP2)	Neutrophils, monocytes	Inflammatory lung disease, COPD
CXCR2	CXCL8, CXCL1 (GRO α), CXCL2 (GRO α), CXCL3 (GRO α), CXCL5 (ENA-78), CXCL6	Neutrophils, monocytes, microvascular endothelial cells	Inflammatory lung disease, COPD, angiogenic for tumor growth
CXCR3-A	CXCL9 (MIG), CXCL10 (IP-10), CXCL11 (I-TAC)	Type 1 helper cells, mast cells, mesangial cells	Inflammatory skin disease, multiple sclerosis, transplant rejection
CXCR3-B	CXCL4 (PF4), CXCL9 (MIG), CXCL10 (IP-10), CXCL11 (I-TAC)	Microvascular endothelial cells, neoplastic cells	Angiostatic for tumor growth
CXCR4	CXCL12 (SDF-1)	Widely expressed	HIV-1 co-receptor (T cell-tropic), tumor metastases, hematopoiesis
CXCR5	CXCL13 (BCA-1)	B cells, follicular helper T cells	Formation of B cell follicles
CXCR6	CXCL16 (SR-PSOX)	CD8+ T cells, natural killer cells, and memory CD4+ T cells	Inflammatory liver disease, atherosclerosis (CXCL16)
CX ₃ CR1	CX3CL1 (fractalkine)	Macrophages, endothelial cells, smooth-muscle cells	Atherosclerosis
XCRI	XCL1 (lymphotactin), XCL2	T cells, natural killer cells	Rheumatoid arthritis, IgA nephropathy, tumor response

Abbreviations: BCA-1, B-cell chemoattractant 1; COPD, chronic obstructive pulmonary disease; CTACK, cutaneous T cell-attracting chemokine; ELC, Epstein-Barr 11-ligand chemokine; ENA, epithelial cell-derived neutrophil-activating peptide; GCP, granulocyte chemotactic protein; GRO, growth-regulated oncogene; HCC, hemofiltrate chemokine; IP-10, interferon inducible 10; I-TAC, interferon-inducible T cell alpha chemoattractant; LARC, liver- and activation-regulated chemokine; MCP, monocyte chemoattractant protein; MDC, macrophage-derived chemokine; MEC, mammary-enriched chemokine; MIG, monokine induced by interferon- γ ; MIP, macrophage inflammatory protein; PF, platelet factor; SDF, stromal cell-derived factor; SLC, secondary lymphoid-tissue chemokine; SR-PSOX, scavenger receptor for phosphatidylserine-containing oxidized lipids; TARC, thymus- and activation-regulated chemokine; TECK, thymus-expressed chemokine; T_H2, type 2 helper T cells.

Source: From IF Charo, RM Ransohoff: *N Engl J Med* 354:610, 2006, with permission. Copyright Massachusetts Medical Society. All rights reserved.

TABLE 372e-9 CYTOKINE FAMILIES GROUPED BY STRUCTURAL SIMILARITY

Hematopoietins	IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-9, IL-11, IL-12, IL-15, IL-16, IL-17, IL-21, IL-23, EPO, LIF, GM-CSF, G-CSF, OSM, CNTF, GH, and TPO
	TNF- α , LT- α , LT- β , CD40L, CD30L, CD27L, 4-1BBL, OX40, OPG, and FasL
IL-1	IL-1 α , IL-1 β , IL-1ra, IL-18, bFGF, aFGF, and ECGF
PDGF	PDGF A, PDGF B, and M-CSF
TGF- β	TGF- β and BMPs (1, 2, 4, etc.)
C-X-C chemokines	IL-8, Gro- $\alpha/\beta/\gamma$, NAP-2, ENA78, GCP-2, PF4, CTAP-3, MIG, and IP-10
C-C chemokines	MCP-1, MCP-2, MCP-3, MIP-1 α , MIP-1 β , RANTES

Abbreviations: aFGF, acidic fibroblast growth factor; 4-1BBL, 401 BB ligand; bFGF, basic fibroblast growth factor; BMP, bone marrow morphogenetic proteins; C-C, cysteine-cysteine; CD, cluster of differentiation; CNTF, ciliary neurotrophic factor; CTAP, connective tissue-activating peptide; C-X-C, cysteine-x-cysteine; ECGF, endothelial cell growth factor; EPO, erythropoietin; FasL, Fas ligand; GCP-2, granulocyte chemotactic protein 2; G-CSF, granulocyte colony-stimulating factor; GH, growth hormone; GM-CSF, granulocyte-macrophage colony-stimulating factor; Gro, growth-related gene products; IFN, interferon; IL, interleukin; IP, interferon- γ inducible protein; LIF, leukemia inhibitory factor; LT, lymphotoxin; MCP, monocyte chemoattractant; M-CSF, macrophage colony-stimulating factor; MIG, monokine induced by interferon- γ ; MIP, macrophage inflammatory protein; NAP-2, neutrophil activating protein 2; OPG, osteoprotegerin; OSM, oncostatin M; PDGF, platelet-derived growth factor; PF, platelet factor; R, receptor; RANTES, regulated on activation, normal T cell-expressed and -secreted; TGF, transforming growth factor; TNF, tumor necrosis factor; TPO, thyroperoxidase.

Cytokines are involved in the regulation of the growth, development, and activation of immune system cells and in the mediation of the inflammatory response. In general, cytokines are characterized by considerable redundancy; different cytokines have similar functions. In addition, many cytokines are pleiotropic in that they are capable of acting on many different cell types. This pleiotropism results from the expression on multiple cell types of receptors for the same cytokine (see below), leading to the formation of “cytokine networks.” The action of cytokines may be (1) autocrine when the target cell is the same cell that secretes the cytokine, (2) paracrine when the target cell is nearby, and (3) endocrine when the cytokine is secreted into the circulation and acts distal to the source.

Cytokines have been named based on presumed targets or based on presumed functions. Those cytokines that are thought to primarily target leukocytes have been named *interleukins* (IL-1, -2, -3, etc.). Many cytokines that were originally described as having a certain function have retained those names (e.g., granulocyte colony-stimulating factor [G-CSF]). Cytokines belong in general to three major structural families: the hematopoietin family; the TNF, IL-1, platelet-derived growth factor (PDGF), and transforming growth factor (TGF) β families; and the CXC and C-C chemokine families (Table 372e-9). Chemokines are cytokines that regulate cell movement and trafficking; they act through G protein-coupled receptors and have a distinctive three-dimensional structure. IL-8 is the only chemokine that early on was named an IL (Table 372e-7).

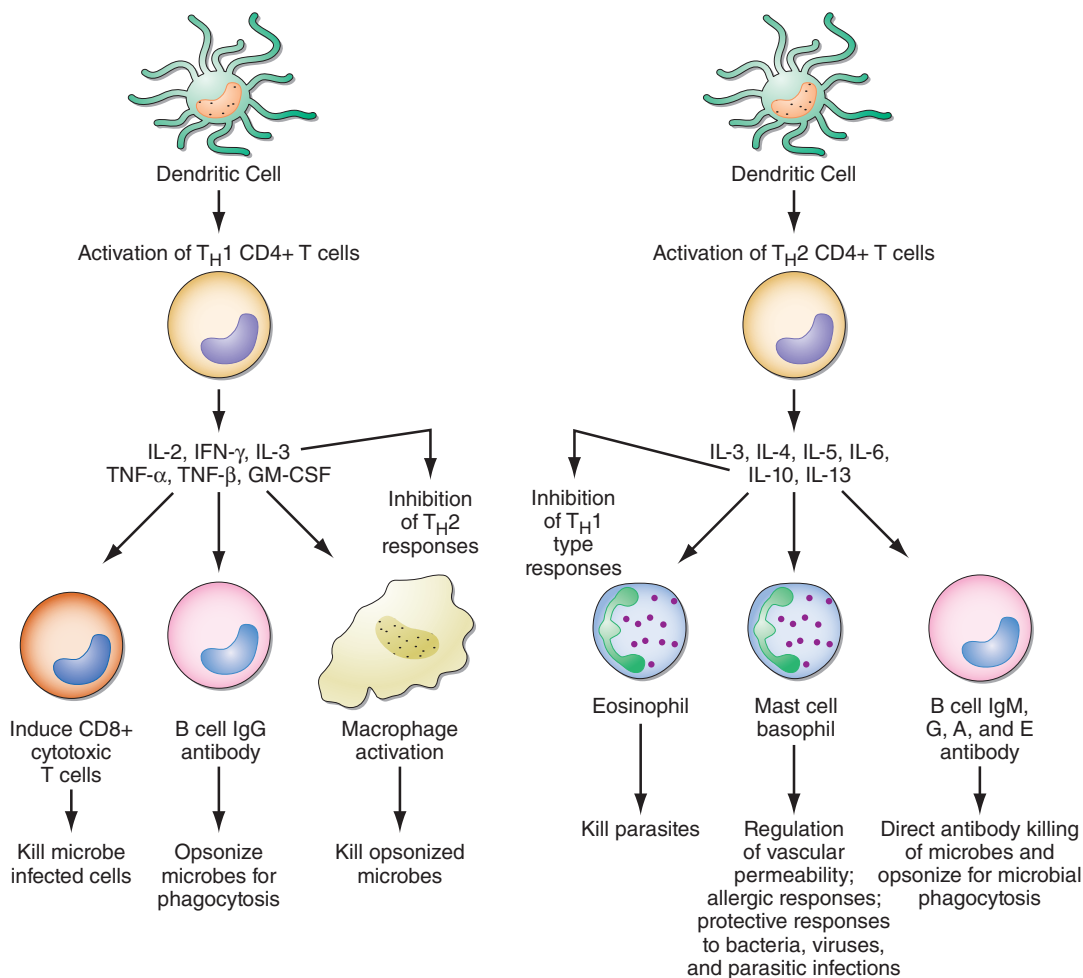


FIGURE 372e-3 CD4+ helper T1 (T_H1) cells and T_H2 T cells secrete distinct but overlapping sets of cytokines. T_H1 CD4+ cells are frequently activated in immune and inflammatory reactions against intracellular bacteria or viruses, whereas T_H2 CD4+ cells are frequently activated for certain types of antibody production against parasites and extracellular encapsulated bacteria; they are also activated in allergic diseases. GM-CSF, granulocyte-macrophage colony-stimulating factor; IFN, interferon; IL, interleukin; TNF, tumor necrosis factor. (Adapted from S Romagnani: CD4 effector cells, in *Inflammation: Basic Principles and Clinical Correlates*, 3rd ed, J Gallin, R Snyderman [eds]. Philadelphia, Lippincott Williams & Wilkins, 1999, p 177; with permission.)

TABLE 372e-10 ASSOCIATION OF KIRs WITH DISEASE

Disease	KIR Association	Observation
Psoriatic arthritis	KIR2DS1/KIR2DS2; HLA-Cw group homozygosity	Susceptibility
Spondylarthritides	Increased KIR3DL2 expression	May contribute to disease pathology
	Interaction HLA-B27 homodimers with KIR3DL1/KIR3DL2; independent of peptide	May contribute to disease pathogenesis
Ankylosing spondylitis	KIR3DL1/3DS1; HLA-B27 genotypes	Susceptibility
Rheumatoid vasculitis	KIR2DS2; HLA-Cw*03	Susceptibility
	Increased KIR2L2/2DS2 in patients with extraarticular manifestations	Clinical manifestations may have different genetic backgrounds with respect to KIR genotype
Rheumatoid arthritis	Decreased KIR2DS1/3DS1 in patients without bone erosions	Susceptibility
	KIR2DS4; HLA-Cw4	Susceptibility
Scleroderma	KIR2DS2+/KIR2DL2-	Susceptibility
Behçet's disease	Altered KIR3DL1 expression	Associated with severe eye disease
Psoriasis vulgaris	2DS1; HLA-Cw*06	Susceptibility
	2DS1; 2DL5; haplotype B	Susceptibility
IDDM	KIR2DS2; HLA-C1	Susceptibility
Type 1 diabetes	KIR2DS2; HLA-C1 and no HLA-C2, no HLA-Bw4	Increased disease progression
Preeclampsia	KIR2DL1 with fewer KIR2DS (mother); HLA-C2 (fetus)	Increased disease progression
AIDS	KIR3DS1; HLA-Bw4Ile80	Decreased disease progression
	KIR3DS1 homozygous; no HLA-Bw4Ile ⁸⁰	Increased disease progression
HCV infection	KIR2DL3 homozygous; HLA-C1 homozygous	Decreased disease progression
Cervical neoplasia (HPV induced)	KIR3DS1; HLA-C1 homozygous and no HLA-Bw4	Increased disease progression
Malignant melanoma	KIR2DL2 and/or KIR2DL3; HLA-C1	Increased disease progression

Abbreviations: HCV, hepatitis C virus; HLA, human leukocyte antigen; HPV, human papillomavirus; IDDM, insulin-dependent diabetes mellitus; KIR, killer cell immunoglobulin-like receptor.

Source: Adapted from R Diaz-Pena et al: *Adv Exp Med Biol* 649:286, 2009.

In general, cytokines exert their effects by influencing gene activation that results in cellular activation, growth, differentiation, functional cell-surface molecule expression, and cellular effector function. In this regard, cytokines can have dramatic effects on the regulation of immune responses and the pathogenesis of a variety of diseases. Indeed, T cells have been categorized on the basis of the pattern of cytokines that they secrete, which results in either humoral immune response (T_H2) or cell-mediated immune response (T_H1). A third type of T helper cell is the T_H17 cell that contributes to host defense against extracellular bacteria and fungi, particularly at mucosal sites (Fig. 372e-2).

Cytokine receptors can be grouped into five general families based on similarities in their extracellular amino acid sequences and conserved structural domains. The *immunoglobulin (Ig) superfamily* represents a large number of cell-surface and secreted proteins. The IL-1 receptors (type 1, type 2) are examples of cytokine receptors with extracellular Ig domains.

The hallmark of the *hematopoietic growth factor (type 1) receptor* family is that the extracellular regions of each receptor contain two conserved motifs. One motif, located at the N terminus, is rich in cysteine residues. The other motif is located at the C terminus proximal to the transmembrane region and comprises five amino acid residues,

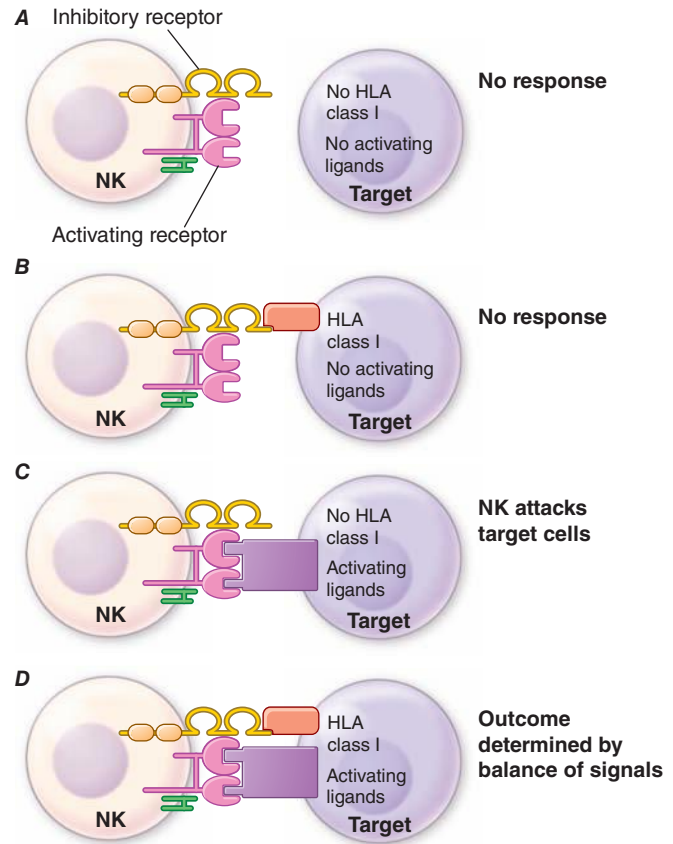


FIGURE 372e-4 Encounters between NK cells: potential targets and possible outcomes. The amount of activating and inhibitory receptors on the NK cells and the amount of ligands on the target cell, as well as the qualitative differences in the signals transduced, determine the extent of the NK response. **A.** When target cells have no HLA class I or activating ligands, NK cells cannot kill target cells. **B.** When target cells bear self-HLA, NK cells cannot kill targets. **C.** When target cells are pathogen infected and have downregulated HLA and express activating ligands, NK cells kill target cells. **D.** When NK cells encounter targets with both self-HLA and activating receptors, then the level of target killing is determined by the balance of inhibitory and activating signals to the NK cell. HLA, human leukocyte antigen; NK, natural killer. (Adapted from L Lanier: *Annu Rev Immunol* 23:225, 2005; reproduced with permission from Annual Reviews Inc. Copyright 2011 by Annual Reviews Inc.)

TABLE 372e-11 EXAMPLES OF MEDIATORS RELEASED FROM IMMUNE CELLS AND BASOPHILS

Mediator	Actions
Histamine	Smooth-muscle contraction, increased vascular permeability
Slow reacting substance of anaphylaxis (SRSA) (leukotriene C_4 , D_4 , E_4)	Smooth-muscle contraction
Eosinophil chemotactic factor of anaphylaxis (ECF-A)	Chemotactic attraction of eosinophils
Platelet-activating factor	Activates platelets to secrete serotonin and other mediators; smooth-muscle contraction; induces vascular permeability
Neutrophil chemotactic factor (NCF)	Chemotactic attraction of neutrophils
Leukotactic activity (leukotriene B_4)	Chemotactic attraction of neutrophils
Heparin	Anticoagulant
Basophil kallikrein of anaphylaxis (BK-A)	Cleaves kininogen to form bradykinin

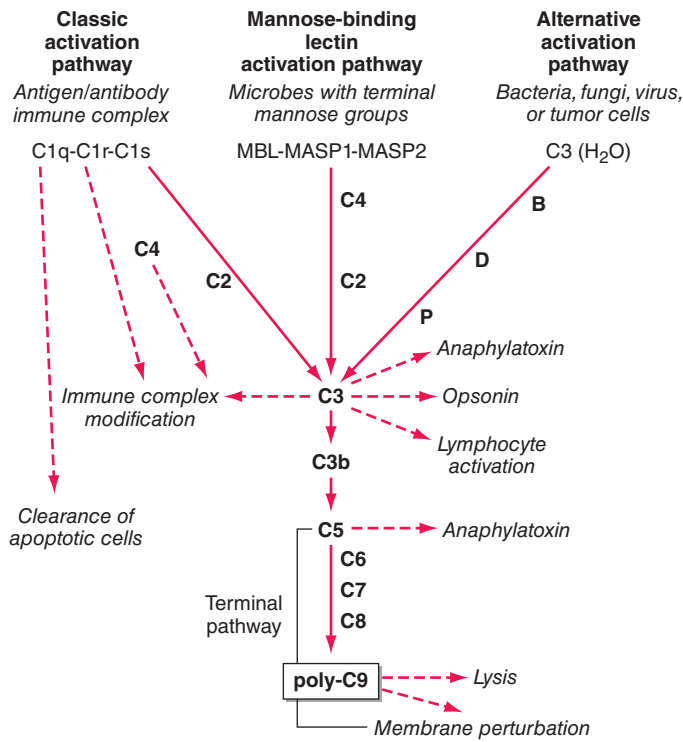


FIGURE 372e-5 The four pathways and the effector mechanisms of the complement system. Dashed arrows indicate the functions of pathway components. (After BJ Morley, MJ Walport: *The Complement Facts Books*. London, Academic Press, 2000, Chap. 2; with permission. Copyright Academic Press, London, 2000.)

tryptophan-serine-X-tryptophan-serine (WSXWS). This family can be grouped on the basis of the number of receptor subunits they have and on the utilization of shared subunits. A number of cytokine receptors, i.e., IL-6, IL-11, IL-12, and leukemia inhibitory factor, are paired with gp130. There is also a common 150-kDa subunit shared by IL-3, IL-5, and granulocyte-macrophage colony-stimulating factor (GM-CSF) receptors. The gamma chain (γ_c) of the IL-2 receptor is common to the IL-2, IL-4, IL-7, IL-9, and IL-15 receptors. Thus, the specific cytokine receptor is responsible for ligand-specific binding, whereas the subunits such as gp130, the 150-kDa subunit, and γ_c are important in signal transduction. The γ_c gene is on the X chromosome, and mutations in the γ_c protein result in the *X-linked form of severe combined immune deficiency syndrome (X-SCID)* (Chap. 374).

The members of the *interferon (type II) receptor* family include the receptors for IFN- γ and - β , which share a similar 210-amino-acid binding domain with conserved cysteine pairs at both the amino and carboxy termini. The members of the *TNF (type III) receptor family* share a common binding domain composed of repeated cysteine-rich regions. Members of this family include the p55 and p75 receptors for TNF (TNF-R1 and TNF-R2, respectively); CD40 antigen, which is an important B cell-surface marker involved in immunoglobulin isotype switching; fas/Apo-1, whose triggering induces apoptosis; CD27 and CD30, which are found on activated T cells and B cells; and nerve growth factor receptor.

The common motif for the *seven transmembrane helix family* was originally found in receptors linked to GTP-binding proteins. This family includes receptors for chemokines (Table 372e-8), β -adrenergic receptors, and retinal rhodopsin. It is important to note that two members of the chemokine receptor family, CXC chemokine receptor type 4 (CXCR4) and β chemokine receptor type 5 (CCR5), have been found to serve as the two major co-receptors for binding and entry of HIV into CD4-expressing host cells (Chap. 226).

Significant advances have been made in defining the signaling pathways through which cytokines exert their intracellular effects. The Janus family of protein tyrosine kinases (JAK) is a critical element involved in signaling via the hematopoietin receptors. Four JAK

kinases, JAK1, JAK2, JAK3, and Tyk2, preferentially bind different cytokine receptor subunits. Cytokine binding to its receptor brings the cytokine receptor subunits into apposition and allows a pair of JAKs to transphosphorylate and activate one another. The JAKs then phosphorylate the receptor on the tyrosine residues and allow signaling molecules to bind to the receptor, whereby the signaling molecules become phosphorylated. Signaling molecules bind the receptor because they have domains (SH2, or src homology 2 domains) that can bind phosphorylated tyrosine residues. There are a number of these important signaling molecules that bind the receptor, such as the adapter molecule SHC, which can couple the receptor to the activation of the mitogen-activated protein kinase pathway. In addition, an important class of substrate of the JAKs is the signal transducers and activators of transcription (STAT) family of transcription factors. STATs have SH2 domains that enable them to bind to phosphorylated receptors, where they are then phosphorylated by the JAKs. It appears that different STATs have specificity for different receptor subunits. The STATs then dissociate from the receptor and translocate to the nucleus, bind to DNA motifs that they recognize, and regulate gene expression. The STATs preferentially bind DNA motifs that are slightly different from one another and thereby control transcription of specific genes. The importance of this pathway is particularly relevant to lymphoid development. Mutations of JAK3 itself also result in a disorder identical to X-SCID; however, because JAK3 is found on chromosome 19 and not on the X chromosome, JAK3 deficiency occurs in boys and girls (Chap. 374).

THE ADAPTIVE IMMUNE SYSTEM

Adaptive immunity is characterized by antigen-specific responses to a foreign antigen or pathogen. A key feature of adaptive immunity is that following the initial contact with antigen (*immunologic priming*), subsequent antigen exposure leads to more rapid and vigorous immune responses (*immunologic memory*). The adaptive immune system consists of dual limbs of cellular and humoral immunity. The principal effectors of cellular immunity are T lymphocytes, whereas the principal effectors of humoral immunity are B lymphocytes. Both B and T lymphocytes derive from a common stem cell (Fig. 372e-6).

The proportion and distribution of immunocompetent cells in various tissues reflect cell traffic, homing patterns, and functional capabilities. Bone marrow is the major site of maturation of B cells, monocytes-macrophages, DCs, and granulocytes and contains pluripotent stem cells that, under the influence of various colony-stimulating factors, are capable of giving rise to all hematopoietic cell types. T cell precursors also arise from hematopoietic stem cells and home to the thymus for maturation. Mature T lymphocytes, B lymphocytes, monocytes, and DCs enter the circulation and home to peripheral lymphoid organs (lymph nodes, spleen) and mucosal surface-associated lymphoid tissue (gut, genitourinary, and respiratory tracts) as well as the skin and mucous membranes and await activation by foreign antigen.

T Cells The pool of effector T cells is established in the thymus early in life and is maintained throughout life both by new T cell production in the thymus and by antigen-driven expansion of virgin peripheral T cells into “memory” T cells that reside in peripheral lymphoid organs. The thymus exports ~2% of the total number of thymocytes per day throughout life, with the total number of daily thymic emigrants decreasing by ~3% per year during the first four decades of life.

Mature T lymphocytes constitute 70–80% of normal peripheral blood lymphocytes (only 2% of the total-body lymphocytes are contained in peripheral blood), 90% of thoracic duct lymphocytes, 30–40% of lymph node cells, and 20–30% of spleen lymphoid cells. In lymph nodes, T cells occupy deep paracortical areas around B cell germinal centers, and in the spleen, they are located in periarteriolar areas of white pulp (Chap. 79). T cells are the primary effectors of cell-mediated immunity, with subsets of T cells maturing into CD8+ cytotoxic T cells capable of lysis of virus-infected or foreign cells (short-lived effector T cells) and CD4+ T cells capable of T cell help for CD8+ T cell and B cell development. Two populations of long-lived memory T cells are triggered by infections: effector memory and central memory T cells.

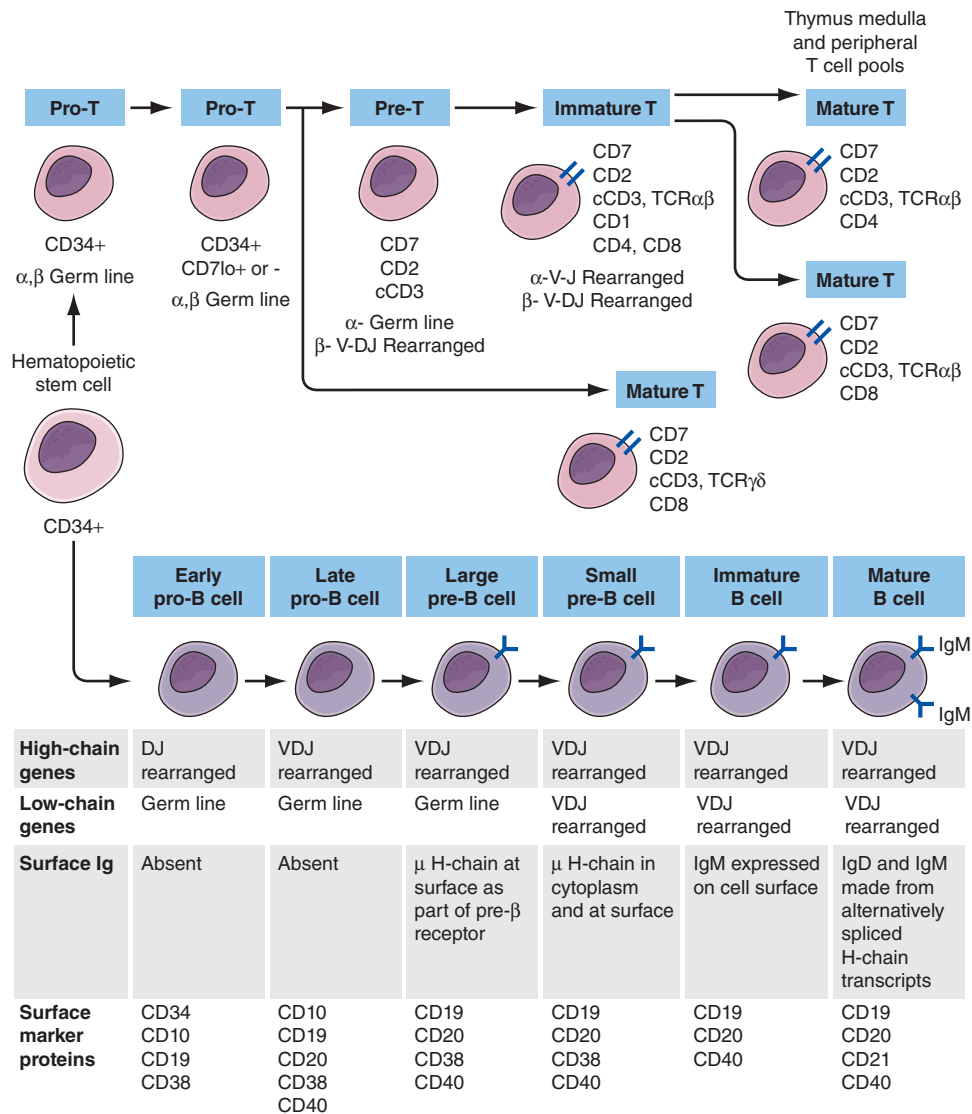


FIGURE 372e-6 Development stages of T and B cells. Elements of the developing T and B cell receptor for antigen are shown schematically. The classification into the various stages of B cell development is primarily defined by rearrangement of the immunoglobulin (Ig) heavy (H) and light (L) chain genes and by the absence or presence of specific surface markers. The classification of stages of T cell development is primarily defined by cell-surface marker protein expression (sCD3, surface CD3 expression; cCD3, cytoplasmic CD3 expression; TCR, T cell receptor). (Adapted from CA Janeway et al [eds]: *Immunobiology. The Immune Systemic Health and Disease, 4th ed.* New York, Garland, 1999; with permission.)

Effector memory T cells reside in nonlymphoid organs and respond rapidly to repeated pathogenic infections with cytokine production and cytotoxic functions to kill virus-infected cells. Central memory T cells home to lymphoid organs where they replenish long- and short-lived and effector memory T cells as needed.

In general, CD4⁺ T cells are the primary regulatory cells of T and B lymphocyte and monocyte function by the production of cytokines and by direct cell contact (Fig. 372e-2). In addition, T cells regulate erythroid cell maturation in bone marrow and, through cell contact (CD40 ligand), have an important role in activation of B cells and induction of Ig isotype switching. Considerable evidence now exists that colonization of the gut by commensal bacteria (the gut microbiome) is responsible for expansion of the peripheral CD4⁺ T cell compartment in normal children and adults.

Human T cells express cell-surface proteins that mark stages of intrathymic T cell maturation or identify specific functional subpopulations of mature T cells. Many of these molecules mediate or participate in important T cell functions (Table 372e-1, Fig. 372e-6).

The earliest identifiable T cell precursors in bone marrow are CD34⁺ pro-T cells (i.e., cells in which TCR genes are neither

rearranged nor expressed). In the thymus, CD34⁺ T cell precursors begin cytoplasmic (c) synthesis of components of the CD3 complex of TCR-associated molecules (Fig. 372e-6). Within T cell precursors, TCR for antigen gene rearrangement yields two T cell lineages, expressing either TCR- $\alpha\beta$ chains or TCR- $\gamma\delta$ chains. T cells expressing the TCR- $\alpha\beta$ chains constitute the majority of peripheral T cells in blood, lymph node, and spleen and terminally differentiate into either CD4⁺ or CD8⁺ cells. Cells expressing TCR- $\gamma\delta$ chains circulate as a minor population in blood; their functions, although not fully understood, have been postulated to be those of immune surveillance at epithelial surfaces and cellular defenses against mycobacterial organisms and other intracellular bacteria through recognition of bacterial lipids.

In the thymus, the recognition of self-peptides on thymic epithelial cells, thymic macrophages, and DCs plays an important role in shaping the T cell repertoire to recognize foreign antigen (*positive selection*) and in eliminating highly autoreactive T cells (*negative selection*). As immature cortical thymocytes begin to express surface TCR for antigen, autoreactive thymocytes are destroyed (negative selection), thymocytes with TCRs capable of interacting with foreign antigen peptides in the context of self-MHC antigens are activated and develop

372e-18 to maturity (positive selection), and thymocytes with TCRs that are incapable of binding to self-MHC antigens die of attrition (*no selection*). Mature thymocytes that are positively selected are either CD4+ helper T cells or MHC class II-restricted cytotoxic (killer) T cells, or they are CD8+ T cells destined to become MHC class I-restricted cytotoxic T cells. *MHC class I- or class II-restricted* means that T cells recognize antigen peptide fragments only when they are presented in the antigen-recognition site of a class I or class II MHC molecule, respectively (Chap. 373e).

After thymocyte maturation and selection, CD4 and CD8 thymocytes leave the thymus and migrate to the peripheral immune system. The thymus continues to be a contributor to the peripheral immune system well into adult life, both normally and when the peripheral T cell pool is damaged, such as occurs in AIDS and cancer chemotherapy.

MOLECULAR BASIS OF T CELL RECOGNITION OF ANTIGEN The TCR for antigen is a complex of molecules consisting of an antigen-binding heterodimer of either $\alpha\beta$ or $\gamma\delta$ chains noncovalently linked with five CD3 subunits (γ , δ , ϵ , ζ , and η) (Fig. 372e-7). The CD3 ζ chains are either disulfide-linked homodimers (CD3- ζ_2) or disulfide-linked heterodimers composed of one ζ chain and one η chain. TCR- $\alpha\beta$ or TCR- $\gamma\delta$ molecules must be associated with CD3 molecules to be inserted into the T cell-surface membrane, TCR α being paired with TCR- β and TCR- γ being paired with TCR- δ . Molecules of the CD3 complex mediate transduction of T cell activation signals via TCRs, whereas TCR- α and - β or - γ and - δ molecules combine to form the TCR antigen-binding site.

The α , β , γ , and δ TCR for antigen molecules have amino acid sequence homology and structural similarities to immunoglobulin

heavy and light chains and are members of the *immunoglobulin gene superfamily* of molecules. The genes encoding TCR molecules are encoded as clusters of gene segments that rearrange during the course of T cell maturation. This creates an efficient and compact mechanism for housing the diversity requirements of antigen receptor molecules. The TCR- α chain is on chromosome 14 and consists of a series of V (variable), J (joining), and C (constant) regions. The TCR- β chain is on chromosome 7 and consists of multiple V, D (diversity), J, and C TCR- β loci. The TCR- γ chain is on chromosome 7, and the TCR- δ chain is in the middle of the TCR- α locus on chromosome 14. Thus, molecules of the TCR for antigen have constant (framework) and variable regions, and the gene segments encoding the α , β , γ , and δ chains of these molecules are recombined and selected in the thymus, culminating in synthesis of the completed molecule. In both T and B cell precursors (see below), DNA rearrangements of antigen receptor genes involve the same enzymes, recombinase activating gene (RAG) 1 and RAG2, both DNA-dependent protein kinases.

TCR diversity is created by the different V, D, and J segments that are possible for each receptor chain by the many permutations of V, D, and J segment combinations, by “N-region diversification” due to the addition of nucleotides at the junction of rearranged gene segments, and by the pairing of individual chains to form a TCR dimer. As T cells mature in the thymus, the repertoire of antigen-reactive T cells is modified by selection processes that eliminate many autoreactive T cells, enhance the proliferation of cells that function appropriately with self-MHC molecules and antigen, and allow T cells with nonproductive TCR rearrangements to die.

TCR- $\alpha\beta$ cells do not recognize native protein or carbohydrate antigens. Instead, T cells recognize only short (~9–13 amino acids) peptide

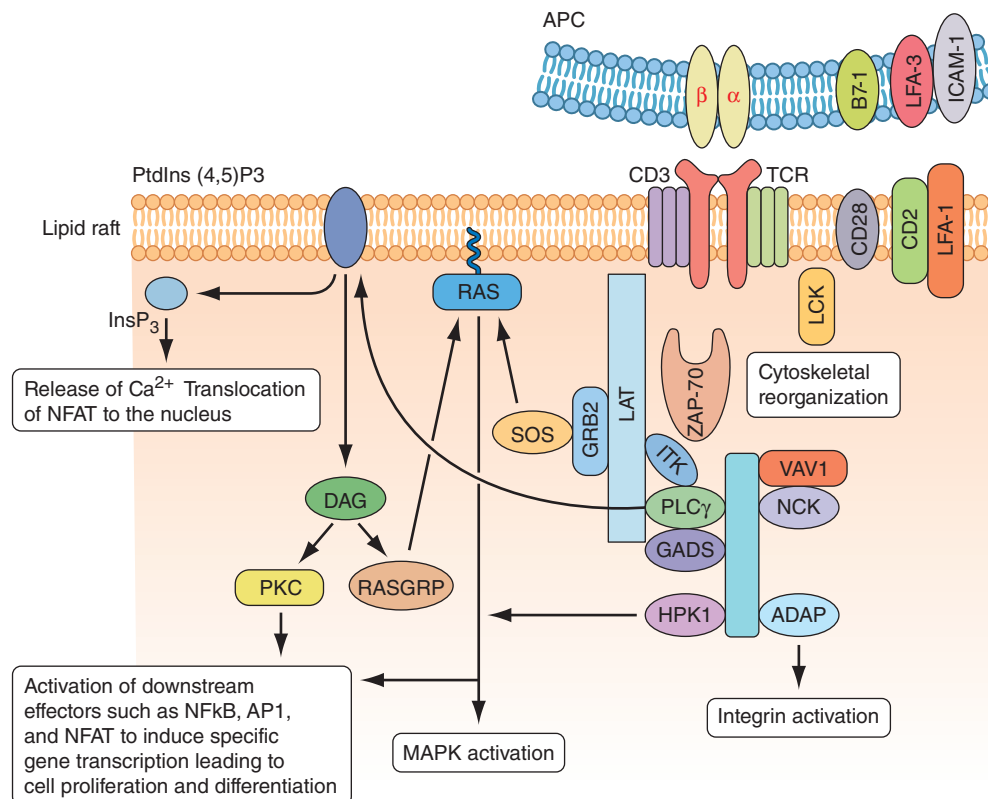


FIGURE 372e-7 Signaling through the T cell receptor. Activation signals are mediated via immunoreceptor tyrosine-based activation (ITAM) sequences in LAT and CD3 chains (blue bars) that bind to enzymes and transduce activation signals to the nucleus via the indicated intracellular activation pathways. Ligation of the T cell receptor (TCR) by MHC complexed with antigen results in sequential activation of LCK and γ -chain-associated protein kinase of 70 kDa (ZAP-70). ZAP-70 phosphorylates several downstream targets, including LAT (linker for activation of T cells) and SLP76 (SCR homology 2 [SH2] domain-containing leukocyte protein of 76 kDa). SLP76 is recruited to membrane-bound LAT through its constitutive interaction with GADS (GRB2-related adaptor protein). Together, SLP76 and LAT nucleate a multimolecular signaling complex, which induces a host of downstream responses, including calcium flux, mitogen-activated protein kinase (MAPK) activation, integrin activation, and cytoskeletal reorganization. APC, antigen-presenting cell. (Adapted from GA Koretzky et al: *Nat Rev Immunol* 6:67, 2006; with permission from Macmillan Publishers Ltd. Copyright 2006.)

fragments derived from protein antigens taken up or produced in APCs. Foreign antigens may be taken up by endocytosis into acidified intracellular vesicles or by phagocytosis and degraded into small peptides that associate with MHC class II molecules (exogenous antigen-presentation pathway). Other foreign antigens arise endogenously in the cytosol (such as from replicating viruses) and are broken down into small peptides that associate with MHC class I molecules (endogenous antigen-presenting pathway). Thus, APCs proteolytically degrade foreign proteins and display peptide fragments embedded in the MHC class I or II antigen-recognition site on the MHC molecule surface, where foreign peptide fragments are available to bind to TCR- $\alpha\beta$ or TCR- $\gamma\delta$ chains of reactive T cells. CD4 molecules act as adhesives and, by direct binding to MHC class II (DR, DQ, or DP) molecules, stabilize the interaction of TCR with peptide antigen (Fig. 372e-7). Similarly, CD8 molecules also act as adhesives to stabilize the TCR-antigen interaction by direct CD8 molecule binding to MHC class I (A, B, or C) molecules.

Antigens that arise in the cytosol and are processed via the endogenous antigen-presentation pathway are cleaved into small peptides by a complex of proteases called the *proteasome*. From the proteasome, antigen peptide fragments are transported from the cytosol into the lumen of the endoplasmic reticulum by a heterodimeric complex termed *transporters associated with antigen processing*, or TAP proteins. There, MHC class I molecules in the endoplasmic reticulum membrane physically associate with processed cytosolic peptides. Following peptide association with class I molecules, peptide-class I complexes are exported to the Golgi apparatus, and then to the cell surface, for recognition by CD8+ T cells.

Antigens taken up from the extracellular space via endocytosis into intracellular acidified vesicles are degraded by vesicle proteases into peptide fragments. Intracellular vesicles containing MHC class II molecules fuse with peptide-containing vesicles, thus allowing peptide fragments to physically bind to MHC class II molecules. Peptide-MHC class II complexes are then transported to the cell surface for recognition by CD4+ T cells (Chap. 373e).

Whereas it is generally agreed that the TCR- $\alpha\beta$ receptor recognizes peptide antigens in the context of MHC class I or class II molecules, lipids in the cell wall of intracellular bacteria such as *M. tuberculosis* can also be presented to a wide variety of T cells, including subsets of TCR- $\gamma\delta$ T cells, and a subset of CD8+ TCR- $\alpha\beta$ T cells. Importantly, bacterial lipid antigens are not presented in the context of MHC class I or II molecules, but rather are presented in the context of MHC-related CD1 molecules. Some $\gamma\delta$ T cells that recognize lipid antigens via CD1 molecules have very restricted TCR usage, do not need antigen priming to respond to bacterial lipids, and may actually be a form of innate rather than acquired immunity to intracellular bacteria.

Just as foreign antigens are degraded and their peptide fragments presented in the context of MHC class I or class II molecules on APCs, endogenous self-proteins also are degraded, and self-peptide fragments are presented to T cells in the context of MHC class I or class II molecules on APCs. In peripheral lymphoid organs, there are T cells that are capable of recognizing self-protein fragments but normally are *anergic* or *tolerant*, i.e., nonresponsive to self-antigenic stimulation, due to lack of self-antigen upregulating APC *co-stimulatory molecules* such as B7-1 (CD80) and B7-2 (CD86) (see below).

Once engagement of mature T cell TCR by foreign peptide occurs in the context of self-MHC class I or class II molecules, binding of non-antigen-specific adhesion ligand pairs such as CD54-CD11/CD18 and CD58-CD2 stabilizes MHC peptide-TCR binding, and the expression of these adhesion molecules is upregulated (Fig. 372e-7). Once antigen ligation of the TCR occurs, the T cell membrane is partitioned into *lipid membrane microdomains*, or *lipid rafts*, that coalesce the key signaling molecules TCR/CD3 complex, CD28, CD2, LAT (linker for activation of T cells), intracellular activated (dephosphorylated) src family protein tyrosine kinases (PTKs), and the key CD3 ζ -associated protein-70 (ZAP-70) PTK (Fig. 372e-7). Importantly, during T cell activation, the CD45 molecule, with protein tyrosine phosphatase activity, is partitioned away from the TCR complex to allow activating phosphorylation events to occur. The coalescence of signaling

molecules of activated T lymphocytes in *microdomains* has suggested that T cell-APC interactions can be considered *immunologic synapses*, analogous in function to neuronal synapses.

After TCR-MHC binding is stabilized, activation signals are transmitted through the cell to the nucleus and lead to the expression of gene products important in mediating the wide diversity of T cell functions such as the secretion of IL-2. The TCR does not have intrinsic signaling activity but is linked to a variety of signaling pathways via immunoreceptor tyrosine-based activation motifs (ITAMs) expressed on the various CD3 chains that bind to proteins that mediate signal transduction. Each of the pathways results in the activation of particular transcription factors that control the expression of cytokine and cytokine receptor genes. Thus, antigen-MHC binding to the TCR induces the activation of the src family of PTKs, fyn and lck (lck is associated with CD4 or CD8 co-stimulatory molecules); phosphorylation of CD3 ζ chain; activation of the related tyrosine kinases ZAP-70 and syk; and downstream activation of the calcium-dependent calcineurin pathway, the ras pathway, and the protein kinase C pathway. Each of these pathways leads to activation of specific families of transcription factors (including NF-AT, fos and jun, and rel/NF- κ B) that form heteromultimers capable of inducing expression of IL-2, IL-2 receptor, IL-4, TNF- α , and other T cell mediators.

In addition to the signals delivered to the T cell from the TCR complex and CD4 and CD8, molecules on the T cell, such as CD28 and inducible co-stimulator (ICOS), and molecules on DCs, such as B7-1 (CD80) and B7-2 (CD86), also deliver important co-stimulatory signals that upregulate T cell cytokine production and are essential for T cell activation. If signaling through CD28 or ICOS does not occur, or if CD28 is blocked, the T cell becomes anergic rather than activated (see “Immune Tolerance and Autoimmunity” below). CTLA-4 (CD152) is similar to CD28 in its ability to bind CD80 and CD86. Unlike CD28, CTLA-4 transmits an inhibitory signal to T cells, acting as an off switch.

T CELL EXHAUSTION IN VIRAL INFECTIONS AND CANCER In chronic viral infections such as HIV-1, hepatitis C virus, and hepatitis B virus and in chronic malignancies, the persistence of antigen disrupts memory T cell function, resulting in defects in memory T cell responses. This has been defined as *T cell exhaustion* and is associated with T cell programmed cell death protein 1 (PD-1) (CD279) expression. Exhausted T cells have compromised proliferation and lose the ability to produce effector molecules, like IL-2, TNF- α , and IFN- γ . PD-1 downregulates T cell responses and is associated with T cell exhaustion and disease progression. For this reason, inhibition of T cell PD-1 activity to enhance effector T cell function is being explored as a target for immunotherapy in both viral infections and certain malignancies.

T CELL SUPERANTIGENS Conventional antigens bind to MHC class I or II molecules in the groove of the $\alpha\beta$ heterodimer and bind to T cells via the V regions of the TCR- α and - β chains. In contrast, superantigens bind directly to the lateral portion of the TCR- β chain and MHC class II β chain and stimulate T cells based solely on the V β gene segment used independent of the D, J, and Va sequences present. *Superantigens* are protein molecules capable of activating up to 20% of the peripheral T cell pool, whereas conventional antigens activate <1 in 10,000 T cells. T cell superantigens include staphylococcal enterotoxins and other bacterial products. Superantigen stimulation of human peripheral T cells occurs in the clinical setting of *staphylococcal toxic shock syndrome*, leading to massive overproduction of T cell cytokines that leads to hypotension and shock (Chap. 172).

B CELLS Mature B cells constitute 10–15% of human peripheral blood lymphocytes, 20–30% of lymph node cells, 50% of splenic lymphocytes, and ~10% of bone marrow lymphocytes. B cells express on their surface intramembrane immunoglobulin (Ig) molecules that function as B cell receptors (BCRs) for antigen in a complex of Ig-associated α and β signaling molecules with properties similar to those described in T cells (Fig. 372e-8). Unlike T cells, which recognize only processed peptide fragments of conventional antigens embedded in the notches of MHC class I and class II antigens of APCs, B cells are capable of

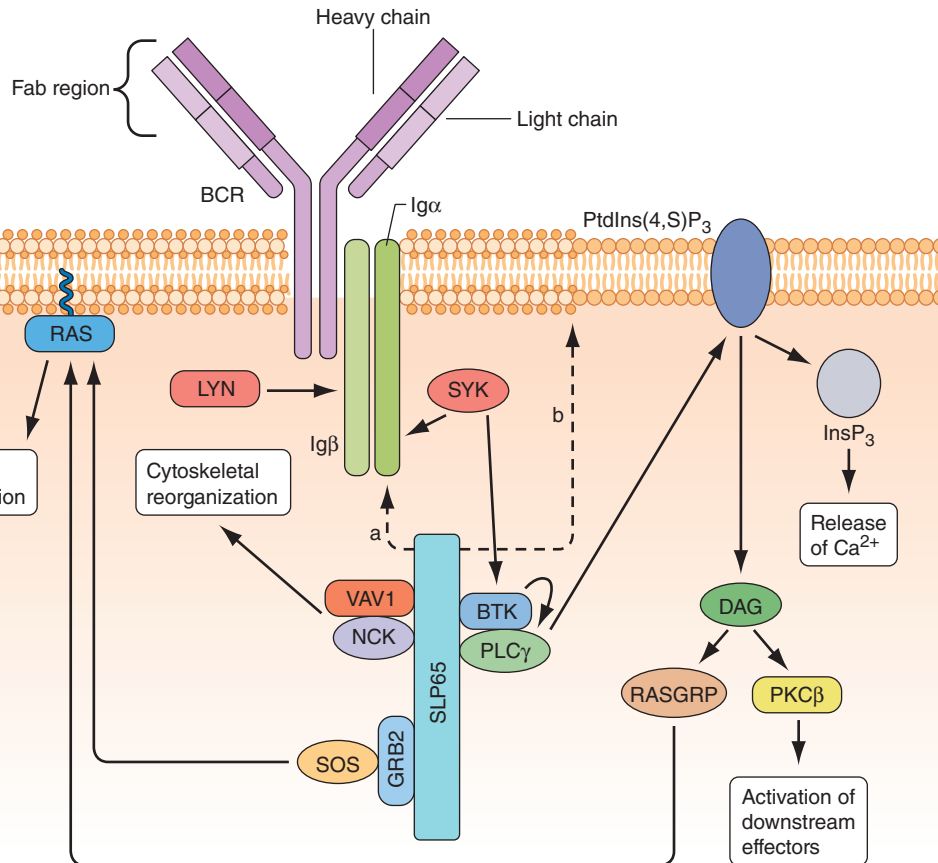


FIGURE 372e-8 B cell receptor (BCR) activation results in the sequential activation of protein tyrosine kinases, which results in the formation of a signaling complex and activation of downstream pathways as shown. Whereas SLP76 is recruited to the membrane through GADS and LAT, the mechanism of SLP65 recruitment is unclear. Studies have indicated two mechanisms: (a) direct binding by the SH2 domain of SLP65 to immunoglobulin (Ig) of the BCR complex or (b) membrane recruitment through a leucine zipper in the amino terminus of SLP65 and an unknown binding partner. ADAP, adhesion- and degranulation-promoting adaptor protein; AP1, activator protein 1; BTK, Bruton's tyrosine kinase; DAG, diacylglycerol; GRB2, growth factor receptor-bound protein 2; HPK1, hematopoietic progenitor kinase 1; InsP_3 , inositol-1,4,5-trisphosphate; ITK, interleukin-2-inducible T cell kinase; NCK, noncatalytic region of tyrosine kinase; NF- κ B, nuclear factor B; PKC, protein kinase C; PLC, phospholipase C; $\text{PtdIns}(4,5)\text{P}_2$, phosphatidylinositol-4,5-bisphosphate; RASGRP, RAS guanyl-releasing protein; SOS, son of sevenless homologue; SYK, spleen tyrosine kinase. (Adapted from GA Koretzky et al: *Nat Rev Immunol* 6:67, 2006; with permission from Macmillan Publishers Ltd. Copyright 2006.)

recognizing and proliferating to whole unprocessed native antigens via antigen binding to B cell-surface Ig (sIg) receptors. B cells also express surface receptors for the Fc region of IgG molecules (CD32) as well as receptors for activated complement components (C3d or CD21, C3b or CD35). The primary function of B cells is to produce antibodies. B cells also serve as APCs and are highly efficient at antigen processing. Their antigen-presenting function is enhanced by a variety of cytokines. Mature B cells are derived from bone marrow precursor cells that arise continuously throughout life (Fig. 372e-6).

B lymphocyte development can be separated into antigen-independent and antigen-dependent phases. Antigen-independent B cell development occurs in primary lymphoid organs and includes all stages of B cell maturation up to the sIg⁺ mature B cell. Antigen-dependent B cell maturation is driven by the interaction of antigen with the mature B cell sIg, leading to memory B cell induction, Ig class switching, and plasma cell formation. Antigen-dependent stages of B cell maturation occur in secondary lymphoid organs, including lymph node, spleen, and gut Peyer's patches. In contrast to the T cell repertoire that is generated intrathymically before contact with foreign antigen, the repertoire of B cells expressing diverse antigen-reactive sites is modified by further alteration of Ig genes after stimulation by antigen—a process called *somatic hypermutation*—that occurs in lymph node germinal centers.

During B cell development, diversity of the antigen-binding variable region of Ig is generated by an ordered set of Ig gene rearrangements

that are similar to the rearrangements undergone by TCR α , β , γ , and δ genes. For the heavy chain, there is first a rearrangement of D segments to J segments, followed by a second rearrangement between a V gene segment and the newly formed D-J sequence; the C segment is aligned to the V-D-J complex to yield a functional Ig heavy chain gene (V-D-J-C). During later stages, a functional κ or γ light chain gene is generated by rearrangement of a V segment to a J segment, ultimately yielding an intact Ig molecule composed of heavy and light chains.

The process of Ig gene rearrangement is regulated and results in a single antibody specificity produced by each B cell, with each Ig molecule comprising one type of heavy chain and one type of light chain. Although each B cell contains two copies of Ig light and heavy chain genes, only one gene of each type is productively rearranged and expressed in each B cell, a process termed *allelic exclusion*.

There are ~ 300 V_κ genes and 5 J_κ genes, resulting in the pairing of V_κ and J_κ genes to create >1500 different kappa light chain combinations. There are ~ 70 V_λ genes and 4 J_λ genes for >280 different lambda light chain combinations. The number of distinct light chains that can be generated is increased by somatic mutations within the V and J genes, thus creating large numbers of possible specificities from a limited amount of germline genetic information. As noted above, in heavy chain Ig gene rearrangement, the V_H domain is created by the joining of three types of germline genes called V_H , D_H , and J_H , thus allowing for even greater diversity in the variable region of heavy chains than of light chains.

The most immature B cell precursors (early pro-B cells) lack cytoplasmic Ig (cIg) and sIg (Fig. 372e-6). The large pre-B cell is marked by the acquisition of the surface pre-BCR composed of μ heavy (H) chains and a pre-B light chain, termed ψ LC. ψ LC is a surrogate light chain receptor encoded by the nonrearranged V pre-B and the $\gamma 5$ light chain locus (the pre-BCR). Pro- and pre-B cells are driven to proliferate and mature by signals from bone marrow stroma—in particular, IL-7. Light chain rearrangement occurs in the small pre-B cell stage such that the full BCR is expressed at the immature B cell stage. Immature B cells have rearranged Ig light chain genes and express sIgM. As immature B cells develop into mature B cells, sIgD is expressed as well as sIgM. At this point, B lineage development in bone marrow is complete, and B cells exit into the peripheral circulation and migrate to secondary lymphoid organs to encounter specific antigens.

Random rearrangements of Ig genes occasionally generate self-reactive antibodies, and mechanisms must be in place to correct these mistakes. One such mechanism is BCR editing, whereby autoreactive BCRs are mutated to not react with self-antigens. If receptor editing is unsuccessful in eliminating autoreactive B cells, then autoreactive B cells undergo negative selection in the bone marrow through induction of apoptosis after BCR engagement of self-antigen.

After leaving the bone marrow, B cells populate peripheral B cell sites, such as lymph node and spleen, and await contact with foreign antigens that react with each B cell's clonotypic receptor. Antigen-driven B cell activation occurs through the BCR, and a process known as *somatic hypermutation* takes place whereby point mutations in rearranged H- and L-genes give rise to mutant sIg molecules, some of which bind antigen better than the original sIg molecules. Somatic hypermutation, therefore, is a process whereby memory B cells in peripheral lymphoid organs have the best binding, or the highest-affinity antibodies. This overall process of generating the best antibodies is called *affinity maturation of antibody*.

Lymphocytes that synthesize IgG, IgA, and IgE are derived from sIgM⁺, sIgD⁺ mature B cells. Ig class switching occurs in lymph node and other peripheral lymphoid tissue germinal centers. CD40 on B cells and CD40 ligand on T cells constitute a critical co-stimulatory receptor-ligand pair of immune-stimulatory molecules. Pairs of CD40⁺ B cells and CD40 ligand⁺ T cells bind and drive B cell Ig class switching via T cell-produced cytokines such as IL-4 and TGF- β .

IL-1, -2, -4, -5, and -6 synergize to drive mature B cells to proliferate and differentiate into Ig-secreting cells.

Humoral Mediators of Adaptive Immunity: Immunoglobulins Immunoglobulins are the products of differentiated B cells and mediate the humoral arm of the immune response. The primary functions of antibodies are to bind specifically to antigen and bring about the inactivation or removal of the offending toxin, microbe, parasite, or other foreign substance from the body. The structural basis of Ig molecule function and Ig gene organization has provided insight into the role of antibodies in normal protective immunity, pathologic immune-mediated damage by immune complexes, and autoantibody formation against host determinants.

All immunoglobulins have the basic structure of two heavy and two light chains (Fig. 372e-8). Immunoglobulin isotype (i.e., G, M, A, D, E) is determined by the type of Ig heavy chain present. IgG and IgA isotypes can be divided further into subclasses (G1, G2, G3, G4, and A1, A2) based on specific antigenic determinants on Ig heavy chains. The characteristics of human immunoglobulins are outlined in **Table 372e-12**. The four chains are covalently linked by disulfide bonds. Each chain is made up of a V region and C regions (also called *domains*), themselves made up of units of ~110 amino acids. Light chains have one variable (V_L) and one constant (C_L) unit; heavy chains have one variable unit (V_H) and three or four constant (C_H) units, depending on isotype. As the name suggests, the constant, or C, regions of Ig molecules are made up of homologous sequences and share the same primary structure as all other Ig chains of the same isotype and subclass. Constant regions are involved in biologic functions of Ig molecules. The C_{H2} domain of IgG and the C_{H4} units of IgM are involved with the binding of the C1q portion of C1 during complement activation. The C_H region at the carboxy-terminal end of the IgG molecule, the Fc region, binds to surface Fc receptors (CD16, CD32, CD64) of macrophages, DCs, NK cells, B cells, neutrophils, and eosinophils. The Fc of IgA binds to Fc α R (CD89), and the Fc of IgE binds to Fc ϵ R (CD23).

Variable regions (V_L and V_H) constitute the antibody-binding (Fab) region of the molecule. Within the V_L and V_H regions are hypervariable regions (extreme sequence variability) that constitute the antigen-binding site unique to each Ig molecule. The idiotype is defined as the specific region of the Fab portion of the Ig molecule to which antigen

TABLE 372e-12 PHYSICAL, CHEMICAL, AND BIOLOGIC PROPERTIES OF HUMAN IMMUNOGLOBULINS

Property	IgG	IgA	IgM	IgD	IgE
Usual molecular form	Monomer	Monomer, dimer	Pentamer, hexamer	Monomer	Monomer
Other chains	None	J chain, SC	J chain	None	None
Subclasses	G1, G2, G3, G4	A1, A2	None	None	None
Heavy chain allotypes	Gm (=30)	No A1, A2m (2)	None	None	None
Molecular mass, kDa	150	160, 400	950, 1150	175	190
Serum level in average adult, mg/mL	9.5–12.5	1.5–2.6	0.7–1.7	0.04	0.0003
Percentage of total serum Ig	75–85	7–15	5–10	0.3	0.019
Serum half-life, days	23	6	5	3	2.5
Synthesis rate, mg/kg per day	33	65	7	0.4	0.016
Antibody valence	2	2, 4	10, 12	2	2
Classical complement activation	+(G1, 2?, 3)	–	++	–	–
Alternate complement activation	+(G4)	+	–	+	–
Binding cells via Fc	Macrophages, neutrophils, large granular lymphocytes	Lymphocytes	Lymphocytes	None	Mast cells, basophils, B cells
Biologic properties	Placental transfer, secondary Ab for most antipathogen responses	Secretory immunoglobulin	Primary Ab responses	Marker for mature B cells	Allergy, antiparasite responses

Source: After L Carayannopoulos, JD Capra, in WE Paul (ed): *Fundamental Immunology*, 3rd ed. New York, Raven, 1993; with permission.

372e-22 binds. Antibodies against the idiotype portion of an antibody molecule are called *anti-idiotypic antibodies*. The formation of such antibodies in vivo during a normal B cell antibody response may generate a negative (or “off”) signal to B cells to terminate antibody production.

IgG constitutes ~75–85% of total serum immunoglobulin. The four IgG subclasses are numbered in order of their level in serum, IgG1 being found in greatest amounts and IgG4 the least. IgG subclasses have clinical relevance in their varying ability to bind macrophage and neutrophil Fc receptors and to activate complement (Table 372e-12). Moreover, selective deficiencies of certain IgG subclasses give rise to clinical syndromes in which the patient is inordinately susceptible to bacterial infections. IgG antibodies are frequently the predominant antibody made after rechallenge of the host with antigen (secondary antibody response).

IgM antibodies normally circulate as a 950-kDa pentamer with 160-kDa bivalent monomers joined by a molecule called the *J chain*, a 15-kDa nonimmunoglobulin molecule that also effects polymerization of IgA molecules. IgM is the first immunoglobulin to appear in the immune response (primary antibody response) and is the initial type of antibody made by neonates. Membrane IgM in the monomeric form also functions as a major antigen receptor on the surface of mature B cells (Table 372e-12). IgM is an important component of immune complexes in autoimmune diseases. For example, IgM antibodies against IgG molecules (rheumatoid factors) are present in high titers in *rheumatoid arthritis*, other collagen diseases, and some infectious diseases (*subacute bacterial endocarditis*).

IgA constitutes only 7–15% of total serum immunoglobulin but is the predominant class of immunoglobulin in secretions. IgA in secretions (tears, saliva, nasal secretions, gastrointestinal tract fluid, and human milk) is in the form of secretory IgA (sIgA), a polymer consisting of two IgA monomers, a joining molecule, again called the *J chain*, and a glycoprotein called the *secretory protein*. Of the two IgA subclasses, IgA1 is primarily found in serum, whereas IgA2 is more prevalent in secretions. IgA fixes complement via the alternative complement pathway and has potent antiviral activity in humans by prevention of virus binding to respiratory and gastrointestinal epithelial cells.

IgD is found in minute quantities in serum and, together with IgM, is a major receptor for antigen on the naïve B cell surface. IgE, which is present in serum in very low concentrations, is the major class of immunoglobulin involved in arming mast cells and basophils by binding to these cells via the Fc region. Antigen cross-linking of IgE molecules on basophil and mast cell surfaces results in release of mediators of the immediate hypersensitivity (allergic) response (Table 372e-12).

CELLULAR INTERACTIONS IN REGULATION OF NORMAL IMMUNE RESPONSES

The net result of activation of the humoral (B cell) and cellular (T cell) arms of the adaptive immune system by foreign antigen is the elimination of antigen directly by specific effector T cells or in concert with specific antibody. Figure 372e-2 is a simplified schematic diagram of the T and B cell responses indicating some of these cellular interactions.

The expression of adaptive immune cell function is the result of a complex series of immunoregulatory events that occur in phases. Both T and B lymphocytes mediate immune functions, and each of these cell types, when given appropriate signals, passes through stages, from activation and induction through proliferation, differentiation, and ultimately effector functions. The effector function expressed may be at the end point of a response, such as secretion of antibody by a differentiated plasma cell, or it might serve a regulatory function that modulates other functions, such as is seen with CD4+ and CD8+ T lymphocytes that modulate both differentiation of B cells and activation of CD8+ cytotoxic T cells.

CD4 helper T cells can be subdivided on the basis of cytokines produced (Fig. 372e-2). Activated T_H1 -type helper T cells secrete IL-2, IFN- γ , IL-3, TNF- α , GM-CSF, and TNF- β , whereas activated T_H2 -type helper T cells secrete IL-3, -4, -5, -6, -10, and -13. T_H1 CD4+ T cells, through elaboration of IFN- γ , have a central role in mediating intracellular killing by a variety of pathogens. T_H1 CD4+ T cells also provide T cell help for generation of cytotoxic T cells and some types of opsonizing antibody, and they generally respond to antigens that lead to

delayed hypersensitivity types of immune responses for many intracellular viruses and bacteria (such as HIV or *M. tuberculosis*). In contrast, T_H2 cells have a primary role in regulatory humoral immunity and isotype switching. T_H2 cells, through production of IL-4 and IL-10, have a regulatory role in limiting proinflammatory responses mediated by T_H1 cells (Fig. 372e-2). In addition, T_H2 CD4+ T cells provide help to B cells for specific Ig production and respond to antigens that require high antibody levels for foreign antigen elimination (extracellular encapsulated bacteria such as *Streptococcus pneumoniae* and certain parasite infections). A new subset of the T_H family has been described, termed T_H17 , characterized as cells that secrete cytokines such as IL-17, -22, and -26. T_H17 cells have been shown to play a role in autoimmune inflammatory disorders in addition to defense against extracellular bacteria and fungi, particularly at mucosal surfaces. In summary, the type of T cell response generated in an immune response is determined by the microbe PAMPs presented to the DCs, the TLRs on the DCs that become activated, the types of DCs that are activated, and the cytokines that are produced (Table 372e-4). Commonly, myeloid DCs produce IL-12 and activate T_H1 T cell responses that result in IFN- γ and cytotoxic T cell induction, and plasmacytoid DCs produce IFN- α and lead to T_H2 responses that result in IL-4 production and enhanced antibody responses.

As shown in Figs. 372e-2 and 372e-3, upon activation by DCs, T cell subsets that produce IL-2, IL-3, IFN- γ , and/or IL-4, -5, -6, -10, and -13 are generated and exert positive and negative influences on effector T and B cells. For B cells, trophic effects are mediated by a variety of cytokines, particularly T cell-derived IL-3, -4, -5, and -6, that act at sequential stages of B cell maturation, resulting in B cell proliferation, differentiation, and ultimately antibody secretion. For cytotoxic T cells, trophic factors include inducer T cell secretion of IL-2, IFN- γ , and IL-12.

An important type of immunomodulatory T cell that controls immune responses is *CD4+ and CD8+ T regulatory cells*. These cells constitutively express the α chain of the IL-2 receptor (CD25), produce large amounts of IL-10, and can suppress both T and B cell responses. T regulatory cells are induced by immature DCs and play key roles in maintaining tolerance to self-antigens in the periphery. Loss of T regulatory cells is the cause of organ-specific autoimmune disease in mice such as autoimmune thyroiditis, adrenalitis, and oophoritis (see “Immune Tolerance and Autoimmunity” below). T regulatory cells also play key roles in controlling the magnitude and duration of immune responses to microbes. Normally, after the initial immune response to a microbe has eliminated the invader, T regulatory cells are activated to suppress the antimicrobe response and prevent host injury. Some microbes have adapted to induce T regulatory cell activation at the site of infection to promote parasite infection and survival. In *Leishmania* infection, the parasite locally induces T regulatory cell accumulation at skin infection sites that dampens anti-*Leishmania* T cell responses and prevents elimination of the parasite. It is thought that many chronic infections such as by *M. tuberculosis* are associated with abnormal T regulatory cell activation that prevents elimination of the microbe.

Although B cells recognize native antigen via B cell-surface Ig receptors, B cells require T cell help to produce high-affinity antibody of multiple isotypes that are the most effective in eliminating foreign antigen. This T cell dependence likely functions in the regulation of B cell responses and in protection against excessive autoantibody production. T cell–B cell interactions that lead to high-affinity antibody production require (1) processing of native antigen by B cells and expression of peptide fragments on the B cell surface for presentation to T_H cells, (2) the ligation of B cells by both the TCR complex and the CD40 ligand, (3) induction of the process termed *antibody isotype switching* in antigen-specific B cell clones, and (4) induction of the process of affinity maturation of antibody in the germinal centers of B cell follicles of lymph node and spleen.

Naïve B cells express cell-surface IgD and IgM, and initial contact of naïve B cells with antigen is via binding of native antigen to B cell-surface IgM. T cell cytokines, released following T_H2 cell contact with B cells or by a “bystander” effect, induce changes in Ig gene conformation

that promote recombination of Ig genes. These events then result in the “switching” of expression of heavy chain exons in a triggered B cell, leading to the secretion of IgG, IgA, or, in some cases, IgE antibody with the same V region antigen specificity as the original IgM antibody, for response to a wide variety of extracellular bacteria, protozoa, and helminths. CD40 ligand expression by activated T cells is critical for induction of B cell antibody isotype switching and for B cell responsiveness to cytokines. Patients with mutations in T cell CD40 ligand have B cells that are unable to undergo isotype switching, resulting in lack of memory B cell generation and the immunodeficiency syndrome of *X-linked hyper-IgM syndrome* (Chap. 374).

IMMUNE TOLERANCE AND AUTOIMMUNITY

Immune tolerance is defined as the absence of activation of pathogenic autoreactivity. *Autoimmune diseases* are syndromes caused by the activation of T or B cells or both, with no evidence of other causes such as infections or malignancies (Chap. 377e). Once thought to be mutually exclusive, immune tolerance and autoimmunity are now both recognized to be present normally in health; when abnormal, they represent extremes from the normal state. For example, it is now known that low levels of autoreactivity of T and B cells with self-antigens in the periphery are critical to their survival. Similarly, low levels of autoreactivity and thymocyte recognition of self-antigens in the thymus are the mechanisms whereby (1) normal T cells are positively selected to survive and leave the thymus to respond to foreign microbes in the periphery and (2) T cells highly reactive to self-antigens are negatively selected and die to prevent overly self-reactive T cells from getting into the periphery (central tolerance). However, not all self-antigens are expressed in the thymus to delete highly self-reactive T cells, and there are mechanisms for peripheral tolerance induction of T cells as well. Unlike the presentation of microbial antigens by mature DCs, the presentation of self-antigens by immature DCs neither activates nor matures the DCs to express high levels of co-stimulatory molecules such as B7-1 (CD80) or B7-2 (CD86). When peripheral T cells are stimulated by DCs expressing self-antigens in the context of HLA molecules, sufficient stimulation of T cells occurs to keep them alive, but otherwise they remain anergic, or nonresponsive, until they contact a DC with high levels of co-stimulatory molecules expressing microbial antigens. In the latter setting, normal T cells then become activated to respond to the microbe. If B cells have high-self-reactivity BCRs, they normally undergo either deletion in the bone marrow or receptor editing to express a less autoreactive receptor. Although many autoimmune diseases are characterized by abnormal or pathogenic autoantibody production (Table 372e-13), most autoimmune diseases are caused by a combination of excess T and B cell reactivity.

Multiple factors contribute to the genesis of clinical autoimmune disease syndromes, including genetic susceptibility (Table 372e-13), environmental immune stimulants such as drugs (e.g., procainamide and phenytoin [Dilantin] with drug-induced systemic lupus erythematosus), infectious agent triggers (such as Epstein-Barr virus and autoantibody production against red blood cells and platelets), and loss of T regulatory cells (leading to thyroiditis, adrenalitis, and oophoritis).

Immunity at Mucosal Surfaces Mucosa covering the respiratory, digestive, and urogenital tracts; the eye conjunctiva; the inner ear; and the ducts of all exocrine glands contain cells of the innate and adaptive mucosal immune system that protect these surfaces against pathogens. In the healthy adult, mucosa-associated lymphoid tissue (MALT) contains 80% of all immune cells within the body and constitutes the largest mammalian lymphoid organ system.

MALT has three main functions: (1) to protect the mucous membranes from invasive pathogens; (2) to prevent uptake of foreign antigens from food, commensal organisms, and airborne pathogens and particulate matter; and (3) to prevent pathologic immune responses from foreign antigens if they do cross the mucosal barriers of the body (Fig. 372e-9).

MALT is a compartmentalized system of immune cells that functions independently from systemic immune organs. Whereas the systemic immune organs are essentially sterile under normal conditions

and respond vigorously to pathogens, MALT immune cells are continuously bathed in foreign proteins and commensal bacteria, and they must select those pathogenic antigens that must be eliminated. MALT contains anatomically defined foci of immune cells in the intestine, tonsil, appendix, and peribronchial areas that are inductive sites for mucosal immune responses. From these sites, immune T and B cells migrate to effector sites in mucosal parenchyma and exocrine glands where mucosal immune cells eliminate pathogen-infected cells. In addition to mucosal immune responses, all mucosal sites have strong mechanical and chemical barriers and cleansing functions to repel pathogens.

Key components of MALT include specialized epithelial cells called “membrane” or “M” cells that take up antigens and deliver them to DCs or other APCs. Effector cells in MALT include B cells producing antipathogen neutralizing antibodies of secretory IgA as well as IgG isotype, T cells producing similar cytokines as in systemic immune system response, and T helper and cytotoxic T cells that respond to pathogen-infected cells.

Secretory IgA is produced in amounts of >50 mg/kg of body weight per 24 h and functions to inhibit bacterial adhesion, inhibit macromolecule absorption in the gut, neutralize viruses, and enhance antigen elimination in tissue through binding to IgA and receptor-mediated transport of immune complexes through epithelial cells.

Recent studies have demonstrated the importance of commensal gut and other mucosal bacteria to the health of the human immune system. Normal commensal flora induces anti-inflammatory events in the gut and protects epithelial cells from pathogens through TLRs and other PRR signaling. When the gut is depleted of normal commensal flora, the immune system becomes abnormal, with loss of T_H1 T cell function. Restoration of the normal gut flora can reestablish the balance in T helper cell ratios characteristic of the normal immune system. When the gut barrier is intact, either antigens do not transverse the gut epithelium or, when pathogens are present, a self-limited, protective MALT immune response eliminates the pathogen (Fig. 372e-9). However, when the gut barrier breaks down, immune responses to commensal flora antigens can cause inflammatory bowel diseases such as *Crohn’s disease* and, perhaps, *ulcerative colitis* (Fig. 372e-9) (Chap. 351). Uncontrolled MALT immune responses to food antigens, such as gluten, can cause *celiac disease* (Chap. 351).

THE CELLULAR AND MOLECULAR CONTROL OF PROGRAMMED CELL DEATH

The process of apoptosis (programmed cell death) plays a crucial role in regulating normal immune responses to antigen. In general, a wide variety of stimuli trigger one of several apoptotic pathways to eliminate microbe-infected cells, eliminate cells with damaged DNA, or eliminate activated immune cells that are no longer needed (Fig. 372e-10). The largest known family of “death receptors” is the TNF receptor (TNF-R) family (TNF-R1, TNF-R2, Fas [CD95], death receptor 3 [DR3], death receptor 4 [DR4; TNF-related apoptosis-inducing ligand receptor 1, or TRAIL-R1], and death receptor 5 [DR5, TRAIL-R2]); their ligands are all in the TNF- α family. Binding of ligands to these death receptors leads to a signaling cascade that involves activation of the *caspase* family of molecules that leads to DNA cleavage and cell death. Two other pathways of programmed cell death involve nuclear *p53* in the elimination of cells with abnormal DNA and *mitochondrial cytochrome c* to induce cell death in damaged cells (Fig. 372e-10). A number of human diseases have now been described that result from, or are associated with, mutated apoptosis genes (Table 372e-14). These include mutations in the Fas and Fas ligand genes in autoimmune and lymphoproliferation syndromes, and multiple associations of mutations in genes in the apoptotic pathway with malignant syndromes.

MECHANISMS OF IMMUNE-MEDIATED DAMAGE TO MICROBES OR HOST TISSUES

Several responses by the host innate and adaptive immune systems to foreign microbes culminate in rapid and efficient elimination of microbes. In these scenarios, the classic weapons of the adaptive immune system (T cells, B cells) interface with cells (macrophages, DCs, NK cells, neutrophils, eosinophils, basophils) and soluble products

TABLE 372e-13 RECOMBINANT OR PURIFIED AUTOANTIGENS RECOGNIZED BY AUTOANTIBODIES ASSOCIATED WITH HUMAN AUTOIMMUNE DISORDERS

Autoantigen	Autoimmune Diseases	Autoantigen	Autoimmune Diseases
Cell- or Organ-Specific Autoimmunity			
Acetylcholine receptor	Myasthenia gravis	Insulin receptor	Type B insulin resistance, acanthosis, systemic lupus erythematosus (SLE)
Actin	Chronic active hepatitis, primary biliary cirrhosis	Intrinsic factor type 1	Pernicious anemia
Adenine nucleotide translator (ANT)	Dilated cardiomyopathy, myocarditis	Leukocyte function-associated antigen (LFA-1)	Treatment-resistant Lyme arthritis
β-Adrenoreceptor	Dilated cardiomyopathy	Myelin-associated glycoprotein (MAG)	Polyneuropathy
Aromatic L-amino acid decarboxylase	Autoimmune polyendocrine syndrome type 1 (APS-1)	Myelin-basic protein	Multiple sclerosis, demyelinating diseases
Asialoglycoprotein receptor	Autoimmune hepatitis	Myelin oligodendrocyte glycoprotein (MOG)	Multiple sclerosis
Bactericidal/permeability-increasing protein (Bpi)	Cystic fibrosis vasculitides	Myosin	Rheumatic fever
Calcium-sensing receptor	Acquired hypoparathyroidism	p-80-Collin	Atopic dermatitis
Cholesterol side-chain cleavage enzyme (CYP11a)	Autoimmune polyglandular syndrome-1	Pyruvate dehydrogenase complex-E2 (PDC-E2)	Primary biliary cirrhosis
Collagen type IV-α3-chain	Goodpasture's syndrome	Sodium iodide symporter (NIS)	Graves' disease, autoimmune hypothyroidism
Cytochrome P450 2D6 (CYP2D6)	Autoimmune hepatitis	SOX-10	Vitiligo
Desmin	Crohn's disease, coronary artery disease	Thyroid and eye muscle shared protein	Thyroid-associated ophthalmopathy
Desmoglein 1	Pemphigus foliaceus	Thyroglobulin	Autoimmune thyroiditis
Desmoglein 3	Pemphigus vulgaris	Thyroid peroxidase	Autoimmune Hashimoto's thyroiditis
F-actin	Autoimmune hepatitis	Thyrotropin receptor	Graves' disease
GM gangliosides	Guillain-Barré syndrome	Tissue transglutaminase	Celiac disease
Glutamate decarboxylase (GAD65)	Type 1 diabetes, stiff-person syndrome	Transcription coactivator p75	Atopic dermatitis
Glutamate receptor (GLUR)	Rasmussen encephalitis	Tryptophan hydroxylase	Autoimmune polyglandular syndrome-1
H/K ATPase	Autoimmune gastritis	Tyrosinase	Vitiligo, metastatic melanoma
17-α-Hydroxylase (CYP17)	Autoimmune polyglandular syndrome-1	Tyrosine hydroxylase	Autoimmune polyglandular syndrome-1
21-Hydroxylase (CYP21)	Addison's disease		
IA-2 (ICA512)	Type 1 diabetes		
Insulin	Type 1 diabetes, insulin hypoglycemic syndrome (Hirata's disease)		
Systemic Autoimmunity			
ACTH	ACTH deficiency	Histone H2A-H2B-DNA	Systemic lupus erythematosus
Aminoacyl-tRNA histidyl synthetase	Myositis, dermatomyositis	IgE receptor	Chronic idiopathic urticaria
Aminoacyl-tRNA synthetase (several)	Polymyositis, dermatomyositis	Keratin	Rheumatoid arthritis
Cardiolipin	Systemic lupus erythematosus, antiphospholipid syndrome	Ku-DNA-protein kinase	Systemic lupus erythematosus
Carbonic anhydrase II	Systemic lupus erythematosus, Sjögren's syndrome, systemic sclerosis	Ku-nucleoprotein La phosphoprotein (La 55-B)	Connective tissue syndrome Sjögren's syndrome
Collagen (multiple types)	Rheumatoid arthritis, systemic lupus erythematosus, progressive systemic sclerosis	Myeloperoxidase	Necrotizing and crescentic glomerulonephritis (NCGN), systemic vasculitis
Centromere-associated proteins	Systemic sclerosis	Proteinase 3 (PR3)	Granulomatosis with polyangiitis (Wegener's), Churg-Strauss syndrome
DNA-dependent nucleoside-stimulated ATPase	Dermatomyositis	RNA polymerase I-III (RNP)	Systemic sclerosis, systemic lupus erythematosus
Fibrillarlin	Scleroderma	Signal recognition protein (SRP54)	Polymyositis
Fibronectin	Systemic lupus erythematosus, rheumatoid arthritis, morphea	Topoisomerase-1 (Scl-70)	Scleroderma, Raynaud's syndrome
Glucose-6-phosphate isomerase	Rheumatoid arthritis	Tublin	Chronic liver disease, visceral leishmaniasis
β2-Glycoprotein I (B2-GPI)	Primary antiphospholipid syndrome	Vimentin	Systemic autoimmune disease
Golgin (95, 97, 160, 180) Heat shock protein	Sjögren's syndrome, systemic lupus erythematosus, rheumatoid arthritis		
Hemidesmosomal protein 180	Bullous pemphigoid, herpes gestationis, cicatricial pemphigoid		

(Continued)

TABLE 372e-13 RECOMBINANT OR PURIFIED AUTOANTIGENS RECOGNIZED BY AUTOANTIBODIES ASSOCIATED WITH HUMAN AUTOIMMUNE DISORDERS (CONTINUED)

Autoantigen	Autoimmune Diseases	Autoantigen	Autoimmune Diseases
Plasma Protein and Cytokine Autoimmunity			
C1 inhibitor	Autoimmune C1 deficiency	Glycoprotein IIb/IIIg and Ib/IX	Autoimmune thrombocytopenia purpura
C1q	Systemic lupus erythematosus, membrane proliferative glomerulonephritis (MPGN)	IgA	Immunodeficiency associated with systemic lupus erythematosus, pernicious anemia, thyroiditis, Sjögren's syndrome, and chronic active hepatitis
Cytokines (IL-1 α , IL-1 β , IL-6, IL-10, LIF)	Rheumatoid arthritis, systemic sclerosis, normal subjects	Oxidized LDL (OxLDL)	Atherosclerosis
Factor II, factor V, factor VII, factor VIII, factor IX, factor X, factor XI, thrombin vWF	Prolonged coagulation time		
Cancer and Paraneoplastic Autoimmunity			
Amphiphysin	Neuropathy, small-cell lung cancer	p62 (IGF-II mRNA-binding protein)	Hepatocellular carcinoma (China)
Cyclin B1	Hepatocellular carcinoma	Recoverin	Cancer-associated retinopathy
DNA topoisomerase II	Liver cancer	Ri protein	Paraneoplastic opsoclonus myoclonus ataxia
Desmoplakin	Paraneoplastic pemphigus	β IV spectrin	Lower motor neuron syndrome
Gephyrin	Paraneoplastic stiff-person syndrome	Synaptotagmin	Lambert-Eaton myasthenic syndrome
Hu proteins	Paraneoplastic encephalomyelitis	Voltage-gated calcium channels	Lambert-Eaton myasthenic syndrome
Neuronal nicotinic acetylcholine receptor	Subacute autonomic neuropathy, cancer	Yo protein	Paraneoplastic cerebellar degeneration
p53	Cancer, systemic lupus erythematosus		

Source: From A Lernmark et al: J Clin Invest 108:1091, 2001; with permission.

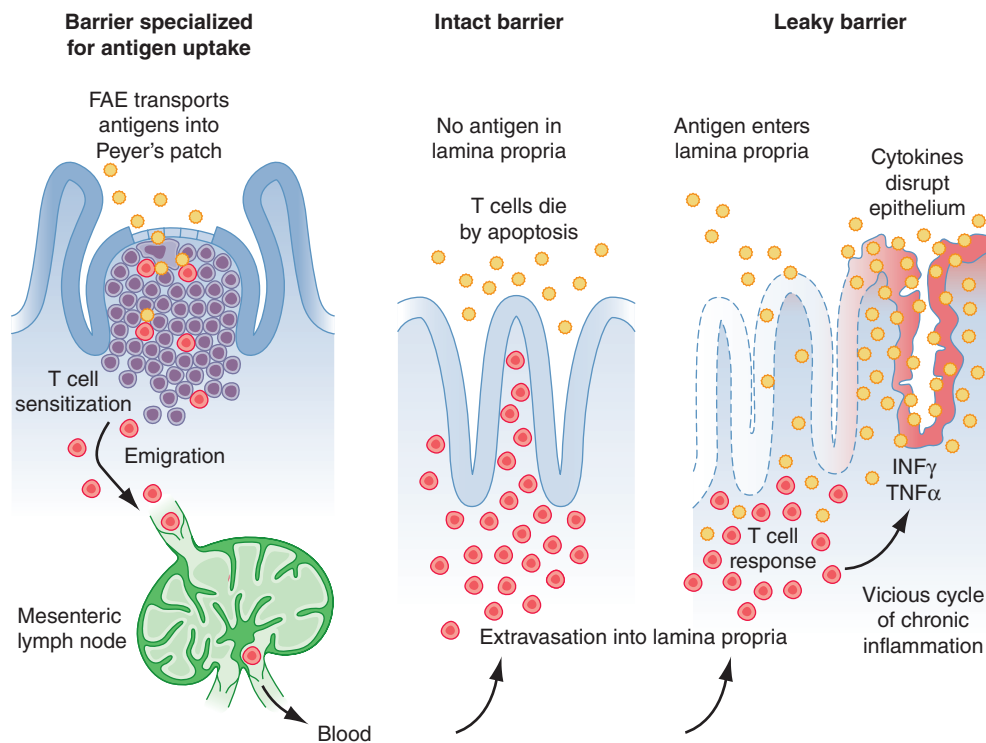


FIGURE 372e-9 Increased epithelial permeability may be important in the development of chronic gut T cell-mediated inflammation. CD4 T cells activated by gut antigens in Peyer's patches migrate to the lamina propria (LP). In healthy individuals, these cells die by apoptosis. Increased epithelial permeability may allow sufficient antigen to enter the LP to trigger T cell activation, breaking tolerance mediated by immunosuppressive cytokines and perhaps T regulatory cells. Proinflammatory cytokines then further increase epithelial permeability, setting up a vicious cycle of chronic inflammation. (From TT MacDonald et al: Science 307:1920, 2005; with permission.)

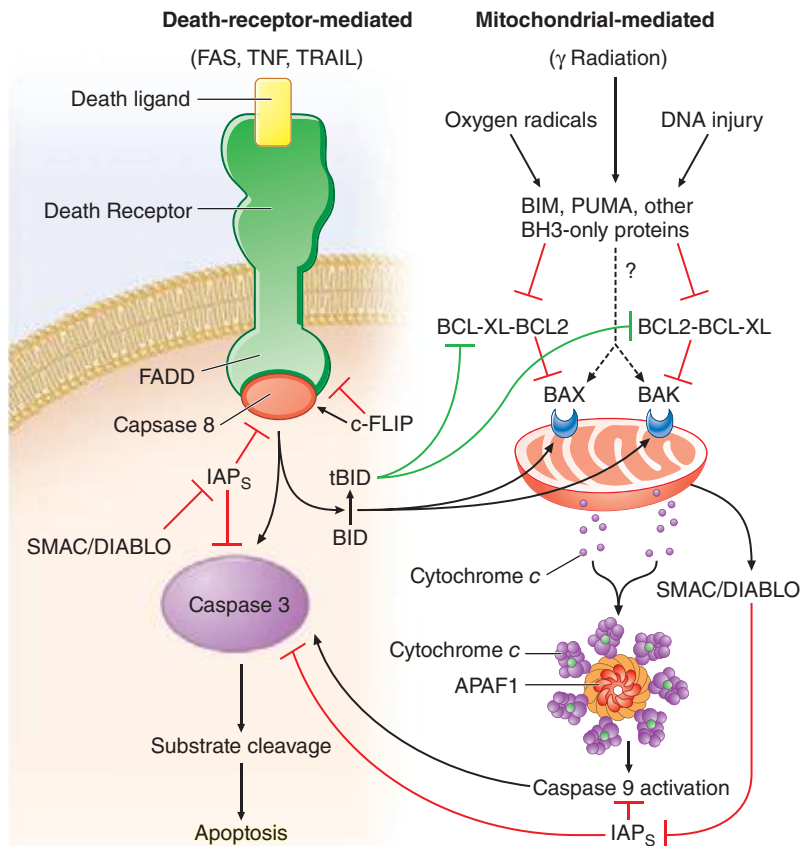


FIGURE 372e-10 Pathways of cellular apoptosis. There are two major pathways of apoptosis: the death-receptor pathway, which is mediated by activation of death receptors, and the BCL2-regulated mitochondrial pathway, which is mediated by noxious stimuli that ultimately lead to mitochondrial injury. Ligation of death receptors recruits the adaptor protein FAS-associated death domain (FADD). FADD in turn recruits caspase 8, which ultimately activates caspase 3, the key “executioner” caspase. Cellular FLICE-inhibitory protein (c-FLIP) can either inhibit or potentiate binding of FADD and caspase 8, depending on its concentration. In the intrinsic pathway, proapoptotic BH3 proteins are activated by noxious stimuli, which interact with and inhibit antiapoptotic BCL2 or BCL-XL. Thus, BAX and BAK are free to induce mitochondrial permeabilization with release of cytochrome c, which ultimately results in the activation of caspase 9 through the apoptosome. Caspase 9 then activates caspase 3. SMAC/DIABLO is also released after mitochondrial permeabilization and acts to block the action of inhibitors of apoptosis protein (IAPs), which inhibit caspase activation. There is potential cross-talk between the two pathways, which is mediated by the truncated form of BID (tBID) that is produced by caspase 8–mediated BID cleavage; tBID acts to inhibit the BCL2-BCL-XL pathway and to activate BAX and BAK. There is debate (indicated by the *question mark*) as to whether proapoptotic BH3 molecules (e.g., BIM and PUMA) act directly on BAX and BAK to induce mitochondrial permeability or whether they act only on BCL2-BCL-XL. APAF1, apoptotic protease-activating factor 1; BH3, BCL homologue; TNF, tumor necrosis factor; TRAIL, TNF-related apoptosis-inducing ligand. (From RS Hotchkiss et al: *N Engl J Med* 361:1570, 2009; with permission.)

(microbial peptides, pentraxins, complement and coagulation systems) of the innate immune system (Chaps. 80 and 376).

There are five general phases of host defenses: (1) migration of leukocytes to sites of antigen localization; (2) antigen-nonspecific recognition of pathogens by macrophages and other cells and systems of the innate immune system; (3) specific recognition of foreign antigens mediated by T and B lymphocytes; (4) amplification of the inflammatory response with recruitment of specific and nonspecific effector cells by complement components, cytokines, kinins, arachidonic acid metabolites, and mast cell–basophil products; and (5) macrophage, neutrophil, and lymphocyte participation in destruction of antigen with ultimate removal

of antigen particles by phagocytosis (by macrophages or neutrophils) or by direct cytotoxic mechanisms (involving macrophages, neutrophils, DCs, and lymphocytes). Under normal circumstances, orderly progression of host defenses through these phases results in a well-controlled immune and inflammatory response that protects the host from the offending antigen. However, dysfunction of any of the host defense systems can damage host tissue and produce clinical disease. Furthermore, for certain pathogens or antigens, the normal immune response itself might contribute substantially to the tissue damage. For example, the immune and inflammatory response in the brain to certain pathogens such as *M. tuberculosis* may be responsible for much of the morbidity rate of this disease in that organ system (Chap. 202). In addition, the morbidity rate associated with certain pneumonias such as that caused by *Pneumocystis jiroveci* may be associated more with inflammatory infiltrates than with the tissue-destructive effects of the microorganism itself (Chap. 244).

Molecular Basis of Lymphocyte–Endothelial Cell Interactions

The control of lymphocyte circulatory patterns between the bloodstream and peripheral lymphoid organs operates at the level of lymphocyte–endothelial cell interactions to control the specificity of lymphocyte subset entry into organs. Similarly, lymphocyte–endothelial cell interactions regulate the entry of lymphocytes into inflamed tissue. Adhesion molecule expression on lymphocytes and endothelial cells regulates the retention and subsequent egress of lymphocytes within tissue sites of antigenic stimulation, delaying cell exit from tissue and preventing reentry into the circulating lymphocyte pool (Fig. 372e-11). All types of lymphocyte migration begin with lymphocyte attachment to specialized regions of vessels, termed *high endothelial venules* (HEVs). An important concept is that adhesion molecules do not generally bind their ligand until a conformational change (ligand activation) occurs in the adhesion molecule that allows ligand binding. Induction of a conformation-dependent determinant on an adhesion molecule can be accomplished by cytokines or via ligation of other adhesion molecules on the cell.

The first stage of lymphocyte–endothelial cell interactions, *attachment and rolling*, occurs when lymphocytes leave the stream of flowing blood cells in a postcapillary venule and roll along venule endothelial cells (Fig. 372e-11). Lymphocyte rolling is mediated by the L-selectin molecule (LECAM-1, LAM-1, CD62L) and slows cell transit time through venules, allowing time for activation of adherent cells.

The second stage of lymphocyte–endothelial cell interactions, *firm adhesion with activation-dependent stable arrest*, requires stimulation of lymphocytes by chemoattractants or by endothelial cell–derived cytokines. Cytokines thought to participate in adherent cell activation include members of the IL-8 family, platelet-activation factor, leukotriene B₄, and C5a. In addition, HEVs express chemokines, SLC (CCL21) and ELC (CCL19), which participate in this process. Following activation by chemoattractants, lymphocytes shed L-selectin from the cell surface and upregulate cell CD11b/18 (MAC-1) or CD11a/18 (LFA-1) molecules, resulting in firm attachment of lymphocytes to HEVs.

Lymphocyte homing to peripheral lymph nodes involves adhesion of L-selectin to glycoprotein HEV ligands collectively referred to as *peripheral node addressin* (PNAd), whereas homing of lymphocytes to intestine Peyer’s patches primarily involves adhesion of the α4β7 integrin to mucosal addressin cell adhesion molecule-1 (MAdCAM-1) on the Peyer’s patch HEVs. However, for migration to mucosal

TABLE 372e-14 IMMUNE SYSTEM MOLECULE DEFECTS IN ANIMALS OR HUMANS THAT CAUSE AUTOIMMUNE OR MALIGNANT SYNDROMES

Protein	Defect	Disease or Syndrome	Observation in Animal Models or Humans
Cytokines and Signaling Proteins			
Tumor necrosis factor (TNF) α	Overexpression	Inflammatory bowel disease (IBD), arthritis, vasculitis	Mice
TNF- α	Underexpression	Systemic lupus erythematosus (SLE)	Mice
Interleukin (IL)-1-receptor antagonist	Underexpression	Arthritis	Mice
IL-2	Overexpression	IBD	Mice
IL-7	Overexpression	IBD	Mice
IL-10	Overexpression	IBD	Mice
IL-2 receptor	Overexpression	IBD	Mice
IL-10 receptor	Overexpression	IBD	Mice
IL-3	Overexpression	Demyelinating syndrome	Mice
Interferon- δ	Overexpression in skin	SLE	Mice
STAT-3	Underexpression	IBD	Mice
STAT-4	Overexpression	IBD	Mice
Transforming growth factor (TGF) β	Underexpression	Systemic wasting syndrome and IBD	Mice
TGF- β receptor in T cells	Underexpression	SLE	Mice
Programmed death (CD279, PD-1)	Underexpression	SLE-like syndrome	Mice
Cytotoxic T lymphocyte, antigen-4 (CTLA-4)	Underexpression	Systemic lymphoproliferative disease	Mice
IL-10	Underexpression	IBD (mouse), type 1 diabetes, thyroid disease, primary (human)	Mice and humans
Major Histocompatibility Locus Molecules^a			
HLA-B27	Allele expression or overexpression	Inflammatory bowel disease	Rats and humans
Complement deficiency of C1, 2, 3 or 4	Underexpression		Humans
LIGHT (TNF superfamily 14)	Overexpression	Systemic lymphoproliferative (mouse) and autoimmunity	Mice
HLA class II DQB10301, DQB10302	Allele expression	Juvenile-onset diabetes	Humans
HLA class II DQB10401, DQB10402	Allele expression	Rheumatoid arthritis	Humans
HLA class I B27	Allele expression	Ankylosing spondylitis, IBD	Rats and humans
Apoptosis Proteins			
TNF receptor 1 (TNF-R1)	Underexpression	Familial periodic fever syndrome	Humans
Fas (CD95; Apo-1)	Underexpression	Autoimmune lymphoproliferative syndrome type 1 (ALPS 1); malignant lymphoma; bladder cancer	Humans
Fas ligand	Underexpression	SLE (only one case identified)	Humans
Perforin	Underexpression	Familial hemophagocytic lymphohistiocytosis (FHL)	Humans
Caspase 10	Underexpression	Autoimmune lymphoproliferative syndrome type II (ALPS II)	Humans
bcl-10	Underexpression	Non-Hodgkin's lymphoma	Humans
P53	Underexpression	Various malignant neoplasms	Humans
Bax	Underexpression	Colon cancer; hematopoietic malignancies	Humans
bcl-2	Underexpression	Non-Hodgkin's lymphoma	Humans
c-IAP2	Underexpression	Low-grade MALT lymphoma	Humans
NAIP1	Underexpression	Spinal muscular atrophy	Humans

^aMany autoimmune diseases are associated with a myriad of major histocompatibility complex gene allele (HLA) types. They are presented here as examples.

Abbreviation: MALT, mucosa-associated lymphoid tissue.

Source: Adapted from L Mullauer: *Mutat Res* 488:211, 2001 and A Davidson, B Diamond: *N Engl J Med* 345:340, 2001.

Peyer's patch lymphoid aggregates, naïve lymphocytes primarily use L-selectin, whereas memory lymphocytes use $\alpha 4\beta 7$ integrin. $\alpha 4\beta 1$ integrin (CD49d/CD29, VLA-4)–VCAM-1 interactions are important in the initial interaction of memory lymphocytes with HEVs of multiple organs in sites of inflammation (Table 372e-15).

The third stage of leukocyte emigration in HEVs is *sticking and arrest*. Sticking of the lymphocyte to endothelial cells and arrest at the site of sticking are mediated predominantly by ligation of $\alpha 1\beta 2$

integrin LFA-1 to the integrin ligand ICAM-1 on HEVs. Whereas the first three stages of lymphocyte attachment to HEVs take only a few seconds, the fourth stage of lymphocyte emigration, *transendothelial migration*, takes ~10 min. Although the molecular mechanisms that control lymphocyte transendothelial migration are not fully characterized, the HEV CD44 molecule and molecules of the HEV glycocalyx (extracellular matrix) are thought to play important regulatory roles in this process (Fig. 372e-11). Finally, expression of matrix metalloproteases

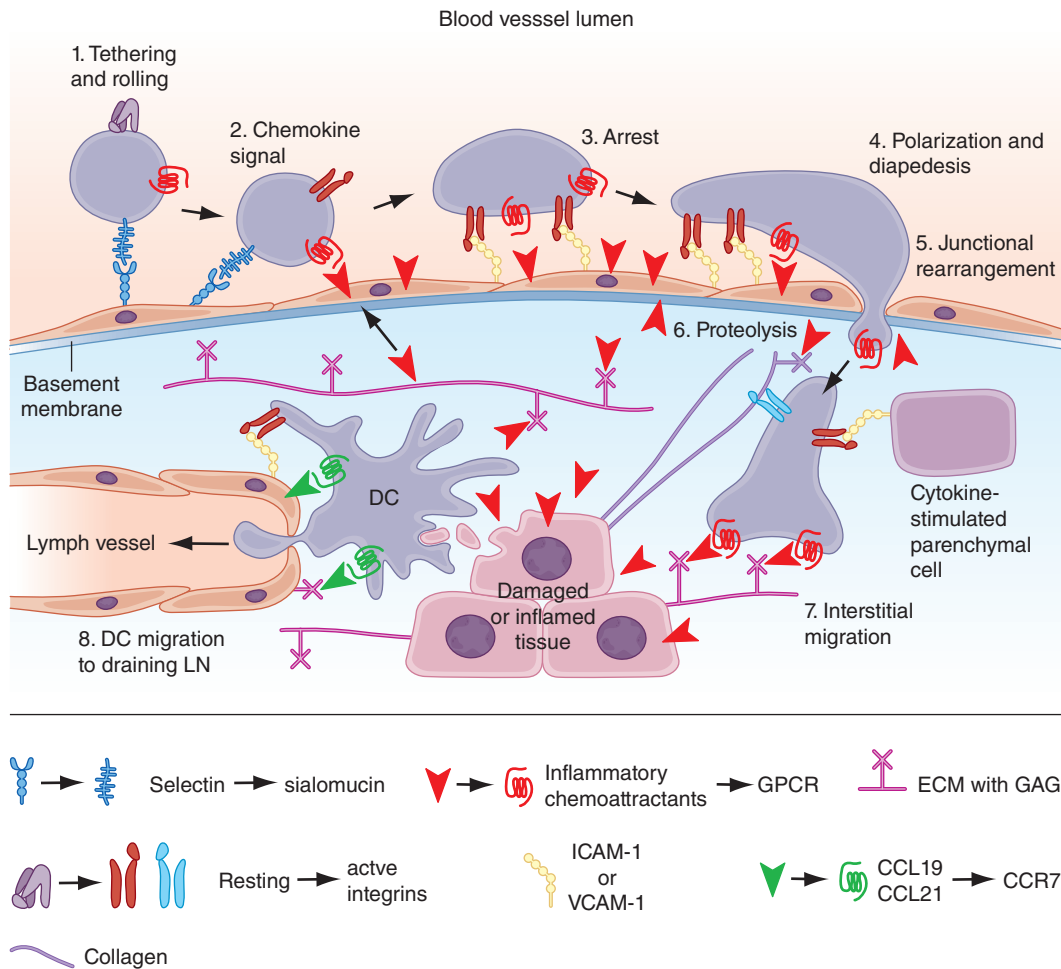


FIGURE 372e-11 Key migration steps of immune cells at sites of inflammation. Inflammation due to tissue damage or infection induces the release of cytokines (not shown) and inflammatory chemoattractants (red arrowheads) from distressed stromal cells and “professional” sentinels, such as mast cells and macrophages (not shown). The inflammatory signals induce upregulation of endothelial selectins and immunoglobulin “superfamily” members, particularly ICAM-1 and/or VCAM-1. Chemoattractants, particularly chemokines, are produced by or translocated across venular endothelial cells (red arrow) and are displayed in the lumen to rolling leukocytes. Those leukocytes that express the appropriate set of trafficking molecules undergo a multistep adhesion cascade (steps 1–3) and then polarize and move by diapedesis across the venular wall (steps 4 and 5). Diapedesis involves transient disassembly of endothelial junctions and penetration through the underlying basement membrane (step 6). Once in the extravascular (interstitial) space, the migrating cell uses different integrins to gain “footholds” on collagen fibers and other ECM molecules, such as laminin and fibronectin, and on inflammation-induced ICAM-1 on the surface of parenchymal cells (step 7). The migrating cell receives guidance cues from distinct sets of chemoattractants, particularly chemokines, which may be immobilized on glycosaminoglycans (GAG) that “decorate” many ECM molecules and stromal cells. Inflammatory signals also induce tissue dendritic cells (DCs) to undergo maturation. Once DCs process material from damaged tissues and invading pathogens, they upregulate CCR7, which allows them to enter draining lymph vessels that express the CCR7 ligand CCL21 (and CCL19). In lymph nodes (LNs), these antigen-loaded mature DCs activate naïve T cells and expand pools of effector lymphocytes, which enter the blood and migrate back to the site of inflammation. T cells in tissue also use this CCR7-dependent route to migrate from peripheral sites to draining lymph nodes through afferent lymphatics. (Adapted from AD Luster et al: *Nat Immunol* 6:1182, 2005; with permission from Macmillan Publishers Ltd. Copyright 2005.)

capable of digesting the subendothelial basement membrane, rich in nonfibrillar collagen, appears to be required for the penetration of lymphoid cells into the extravascular sites.

Abnormal induction of HEV formation and use of the molecules discussed above have been implicated in the induction and maintenance of inflammation in a number of chronic inflammatory diseases. In animal models of type 1 diabetes mellitus, MAdCAM-1 and GlyCAM-1 have been shown to be highly expressed on HEVs in inflamed pancreatic islets, and treatment of these animals with inhibitors of L-selectin and $\alpha 4$ integrin function blocked the development of type 1 diabetes mellitus (Chap. 417). A similar role for abnormal induction of the adhesion molecules of lymphocyte emigration has been suggested in rheumatoid arthritis (Chap. 380), Hashimoto’s thyroiditis (Chap. 405), Graves’ disease (Chap. 405), multiple sclerosis (Chap. 458), Crohn’s disease (Chap. 351), and ulcerative colitis (Chap. 351).

Immune-Complex Formation Clearance of antigen by immune-complex formation between antigen, complement, and antibody is a highly effective mechanism of host defense. However, depending on the level of immune complexes formed and their physicochemical properties, immune complexes may or may not result in host and foreign cell damage. After antigen exposure, certain types of soluble antigen-antibody complexes freely circulate and, if not cleared by the reticuloendothelial system, can be deposited in blood vessel walls and in other tissues such as renal glomeruli and cause *vasculitis* or *glomerulonephritis* syndromes (Chaps. 338 and 385). Deficiencies of early complement components are associated with inefficient clearance of immune complexes and immune complex mediated tissue damage in autoimmune syndromes, whereas deficiencies of the later complement components are associated with susceptibility to recurrent *Neisseria* infections (Table 372e-16).

TABLE 372e-15 TRAFFICKING MOLECULES INVOLVED IN INFLAMMATORY DISEASE PROCESSES

Disease	Key Effector Cell	Proposed Leukocyte Receptors for Endothelial Traffic Signals		
		L-Selectin, Ligand	GPCR	Integrin ^a
Acute Inflammation				
Myocardial infarction	Neutrophil	PSGL-1	CXCR1, CXCR2, PAFR, BLT1	LFA-1, Mac-1
Stroke	Neutrophil	L-Selectin, PSGL-1	CXCR1, CXCR2, PAFR, BLT1	LFA-1, Mac-1
Ischemia-reperfusion	Neutrophil	PSGL-1	CXCR1, CXCR2, PAFR, BLT1	LFA-1, Mac-1
T_H1 Inflammation				
Atherosclerosis	Monocyte	PSGL-1	CCR1, CCR2, BLT1, CXCR2, CX3CR1	VLA-4
Multiple sclerosis	T _H 1	PSGL-1	CXCR3, CCR5	VLA-4
	T _H 1	PSGL-1 (?)	CXCR3, CXCR6	VLA-4, LFA-1
Rheumatoid arthritis	Monocyte	PSGL-1 (?)	CCR2, CCR1	VLA-4, LFA-1
	Monocyte	PSGL-1	CCR1, CCR2	VLA-1, VLA-2, VLA-4, LFA-1
Psoriasis	T _H 1	PSGL-1	CXCR3, CXCR6	VLA-1, VLA-2, VLA-4, LFA-1
	Neutrophil	L-Selectin, PSGL-1	CXCR2, BLT1	LFA-1 ^b
Crohn's disease	Skin-homing T _H 1	CLA	CCR4, CCR10, CXCR3	VLA-4 ^c , LFA-1
Crohn's disease	Gut-homing T _H 1	PSGL-1	CCR9, CXCR3	α4, β7, LFA-1
Type 1 diabetes	T _H 1	PSGL-1 (?)	CCR4, CCR5	VLA-4, LFA-1
Allograft rejection	CD8	L-Selectin (?), PSGL-1 (?)	CXCR3	VLA-4, LFA-1
	CD8	PSGL-1	CXCR3, CX3CR1, BLT1	VLA-4, LFA-1
Hepatitis	B cell	L-Selectin, PSGL-1	CXCR5, CXCR4	VLA-4, LFA-1
	CD8	PSGL-1	CXCR3, CCR5, CXCR6	VLA-4
Lupus	T _H 1	None	CXCR6	VLA-4 ^d
	Plasmacytoid DC	L-Selectin, CLA	CCR7, CXCR3, ChemR23	LFA-1, Mac-1
	B cell	CLA (?)	CXCR5, CXCR4	LFA-1
T_H2 Inflammation				
Asthma	T _H 2	PSGL-1	CCR4, CCR8, BLT1	LFA-1
	Eosinophil	PSGL-1	CCR3, PAFR, BLT1	VLA-4, LFA-1
Atopic dermatitis	Mast cells	PSGL-1	CCR2, CCR3, BLT1	VLA-4, LFA-1
	Skin-homing T _H 2	CLA	CCR4, CCR10	VLA-4, LFA-1

^aVarious β₁ integrins have been linked in different ways in basal lamina and interstitial migration of distinct cell types and inflammatory settings. ^bIn some settings, Mac-1 has been linked to transmigration. ^cCD44 can act in concert with VLA-4 in particular models of leukocyte arrest. ^dT_H2 cells require VAP-1 to traffic to inflamed liver.

Source: From AD Luster et al: Nat Immunol 6:1182, 2005; with permission from Macmillan Publishers Ltd. Copyright 2005.

TABLE 372e-16 COMPLEMENT DEFICIENCIES AND ASSOCIATED DISEASES

Component	Associated Diseases
Classic Pathway	
Cl _q , Cl _r , Cl _s , C4	Immune-complex syndromes, ^a pyogenic infections
C2	Immune-complex syndromes, ^a few with pyogenic infections
C1 inhibitor	Rare immune-complex disease, few with pyogenic infections
C3 and Alternative Pathway C3	
C3	Immune-complex syndromes, ^a pyogenic infections
D	Pyogenic infections
Properdin	<i>Neisseria</i> infections
I	Pyogenic infections
H	Hemolytic-uremic syndrome
Membrane Attack Complex	
C5, C6, C7, C8	Recurrent <i>Neisseria</i> infections, immune-complex disease
C9	Rare <i>Neisseria</i> infections

^aImmune-complex syndromes include systemic lupus erythematosus (SLE) and SLE-like syndromes, glomerulonephritis, and vasculitis syndromes.

Source: After JA Schifferli, DK Peters: Lancet 322:957, 1983. Copyright 1983, with permission from Elsevier.

Immediate-Type Hypersensitivity Helper T cells that drive anti-allergen IgE responses are usually T_H2-type inducer T cells that secrete IL-4, IL-5, IL-6, and IL-10. Mast cells and basophils have high-affinity receptors for the Fc portion of IgE (FcRI), and cell-bound anti-allergen IgE effectively “arms” basophils and mast cells. Mediator release is triggered by antigen (allergen) interaction with Fc receptor-bound IgE, and the mediators released are responsible for the pathophysiologic changes of *allergic diseases* (Table 372e-11). Mediators released from mast cells and basophils can be divided into three broad functional types: (1) those that increase vascular permeability and contract smooth muscle (histamine, platelet-activating factor, SRS-A, BK-A), (2) those that are chemotactic for or activate other inflammatory cells (ECF-A, NCF, leukotriene B₄), and (3) those that modulate the release of other mediators (BK-A, platelet-activating factor) (**Chap. 376**).

Cytotoxic Reactions of Antibody In this type of immunologic injury, complement-fixing (C1-binding) antibodies against normal or foreign cells or tissues (IgM, IgG1, IgG2, IgG3) bind complement via the classic pathway and initiate a sequence of events similar to that initiated by immune-complex deposition, resulting in cell lysis or tissue injury. Examples of antibody-mediated cytotoxic reactions include red cell lysis in *transfusion reactions*, *Goodpasture's syndrome* with anti-glomerular basement membrane antibody formation, and *pemphigus vulgaris* with anti-epidermal antibodies inducing blistering skin disease.

Classic Delayed-Type Hypersensitivity Reactions Inflammatory reactions initiated by mononuclear leukocytes and not by antibody alone have

372e-30 been termed *delayed-type hypersensitivity reactions*. The term *delayed* has been used to contrast a secondary cellular response that appears 48–72 h after antigen exposure with an *immediate* hypersensitivity response generally seen within 12 h of antigen challenge and initiated by basophil mediator release or preformed antibody. For example, in an individual previously infected with *M. tuberculosis* organisms, intradermal placement of tuberculin purified protein derivative as a skin test challenge results in an indurated area of skin at 48–72 h, indicating previous exposure to tuberculosis.

The cellular events that result in classic delayed-type hypersensitivity responses are centered on T cells (predominantly, although not exclusively, IFN- γ , IL-2, and TNF- α -secreting T_H1-type helper T cells) and macrophages. Recently NK cells have been suggested to play a major role in the form of delayed hypersensitivity that occurs following skin contact with immunogens. First, local immune and inflammatory responses at the site of foreign antigen upregulate endothelial cell adhesion molecule expression, promoting the accumulation of lymphocytes at the tissue site. In the general schemes outlined in Figs. 372e-2 and 372e-3, antigen is processed by DCs and presented to small numbers of CD4⁺ T cells expressing a TCR specific for the antigen. IL-12 produced by APCs induces T cells to produce IFN- γ (T_H1 response). Macrophages frequently undergo epithelioid cell transformation and fuse to form multinucleated giant cells in response to IFN- γ . This type of mononuclear cell infiltrate is termed *granulomatous inflammation*. Examples of diseases in which delayed-type hypersensitivity plays a major role are fungal infections (*histoplasmosis*; Chap. 236), mycobacterial infections (*tuberculosis*, *leprosy*; Chaps. 202 and 203), chlamydial infections (*lymphogranuloma venereum*; Chap. 213), helminth infections (*schistosomiasis*; Chap. 259), reactions to toxins (*berylliosis*; Chap. 311), and hypersensitivity reactions to organic dusts (*hypersensitivity pneumonitis*; Chap. 310). In addition, delayed-type hypersensitivity responses play important roles in tissue damage in autoimmune diseases such as *rheumatoid arthritis*, *temporal arteritis*, and *granulomatosis with polyangiitis* (Wegener's) (Chaps. 380 and 385).

CLINICAL EVALUATION OF IMMUNE FUNCTION

Clinical assessment of immunity requires investigation of the four major components of the immune system that participate in host defense and in the pathogenesis of autoimmune diseases: (1) humoral immunity (B cells); (2) cell-mediated immunity (T cells, monocytes); (3) phagocytic cells of the reticuloendothelial system (macrophages), as well as polymorphonuclear leukocytes; and (4) complement. Clinical problems that require an evaluation of immunity include chronic infections, recurrent infections, unusual infecting agents, and certain autoimmune syndromes. The type of clinical syndrome under evaluation can provide information regarding possible immune defects (Chap. 374). Defects in cellular immunity generally result in viral, mycobacterial, and fungal infections. An extreme example of deficiency in cellular immunity is AIDS (Chap. 226). Antibody deficiencies result in recurrent bacterial infections, frequently with organisms such as *S. pneumoniae* and *Haemophilus influenzae* (Chap. 374). Disorders of phagocyte function are frequently manifested by recurrent skin infections, often due to *Staphylococcus aureus* (Chap. 80). Finally, deficiencies of early and late complement components are associated with autoimmune phenomena and recurrent *Neisseria* infections (Table 372e-16). **For further discussion of useful initial screening tests of immune function, see Chap. 374.**

IMMUNOTHERAPY

Many therapies for autoimmune and inflammatory diseases involve the use of nonspecific immune-modulating or immunosuppressive agents such as glucocorticoids or cytotoxic drugs. The goal of development of new treatments for immune-mediated diseases is to design ways to specifically interrupt pathologic immune responses, leaving nonpathologic immune responses intact. Novel ways to interrupt pathologic immune responses that are under investigation include the use of anti-inflammatory cytokines or specific cytokine inhibitors as anti-inflammatory agents, the use of monoclonal antibodies against

T or B lymphocytes as therapeutic agents, the use of intravenous Ig for certain infections and immune complex-mediated diseases, the use of specific cytokines to reconstitute components of the immune system, and bone marrow transplantation to replace the pathogenic immune system with a more normal immune system (Chaps. 80, 374, and 226). In particular, the use of a monoclonal antibody to B cells (rituximab, anti-CD20 MAb) is approved in the United States for the treatment of non-Hodgkin's lymphoma (Chap. 134) and, in combination with methotrexate, for treatment of adult patients with severe rheumatoid arthritis resistant to TNF- α inhibitors (Chap. 380). The U.S. Food and Drug Administration (FDA) approved the use of CTLA-4 antibodies in 2010 to block T cell anergy for use in cancer immunotherapy, and it was the first agent to demonstrate survival benefit in patients with advanced melanoma. Early-stage clinical trials have now shown that PD-1 blockade to reverse T cell exhaustion can induce tumor regression.

Cell-based therapies have been studied for many years, including ex vivo activation of NK cells for reinfusion into patients with malignancies and DC therapy of ex vivo priming of DCs for enhanced presentation of cancer antigens, with reinfusion of primed DCs into the patient. One such strategy for DC therapy has been approved by the FDA for treatment of advanced prostate cancer.

Cytokines and Cytokine Inhibitors Several TNF inhibitors are used as biological therapies in the treatment of rheumatoid arthritis; these include monoclonal antibodies, TNF-R Fc fusion proteins, and Fab fragments. Use of anti-TNF- α antibody therapies such as adalimumab, infliximab, and golimumab has resulted in clinical improvement in patients with these diseases and has opened the way for targeting TNF- α to treat other severe forms of autoimmune and/or inflammatory disease. Blockage of TNF- α has been effective in *rheumatoid arthritis*, *psoriasis*, *Crohn's disease*, and *ankylosing spondylitis*. Other cytokine inhibitors are recombinant soluble TNF- α receptor (R) fused to human Ig and anakinra (soluble IL-1 receptor antagonist, or IL-1ra). The treatment of autoinflammatory syndromes (Table 372e-6) with recombinant IL-1 receptor antagonist can prevent symptoms in these syndromes, because the overproduction of IL-1 β is a hallmark of these diseases.

TNF- α R-Fc fusion protein (etanercept) and IL-1ra act to inhibit the activity of pathogenic cytokines in rheumatoid arthritis, i.e., TNF- α and IL-1, respectively. Similarly, anti-IL-6, IFN- β , and IL-11 act to inhibit pathogenic proinflammatory cytokines. Anti-IL-6 (tocilizumab) inhibits IL-6 activity, whereas IFN- β and IL-11 decrease IL-1 and TNF- α production.

Of particular note has been the successful use of IFN- γ in the treatment of the phagocytic cell defect in *chronic granulomatous disease* (Chap. 80).

Monoclonal Antibodies to T and B Cells The OKT3 MAb against human T cells has been used for several years as a T cell-specific immunosuppressive agent that can substitute for horse antithymocyte globulin (ATG) in the treatment of solid organ transplant rejection. OKT3 produces fewer allergic reactions than ATG but does induce human anti-mouse Ig antibody—thus limiting its use. Anti-CD4 MAb therapy has been used in trials to treat patients with rheumatoid arthritis. While inducing profound immunosuppression, anti-CD4 MAb treatment also induces susceptibility to severe infections. Treatment of patients with a MAb against the T cell molecule CD40 ligand (CD154) is under investigation to induce tolerance to organ transplants, with promising results reported in animal studies. Monoclonal antibodies to the CD25 (IL-2a) receptor (basiliximab) are being used for treatment of graft-versus-host disease in bone marrow transplantation, and anti-CD20 MAb (rituximab) is used to treat hematologic neoplasms, autoimmune diseases, kidney transplant rejection, and rheumatoid arthritis. The anti-IgE monoclonal antibody (omalizumab) is used for blocking antigen-specific IgE that causes *hay fever* and *allergic rhinitis* (Chap. 376); however, side effects of anti-IgE include increased risk of anaphylaxis. Studies have shown that T_H17 cells, in addition to T_H1, are mediators of inflammation in Crohn's disease, and anti-IL-12/IL-23p40 antibody therapy has been studied as a treatment.

It is important to realize the potential risks of these immunosuppressive monoclonal antibodies. Natalizumab is a humanized IgG antibody against an $\alpha 4$ integrin that inhibits leukocyte migration into tissues and has been approved for treatment of multiple sclerosis in the United States. Both it and anti-CD20 (rituximab) have been associated with the onset of progressive multifocal leukoencephalopathy (PML)—a serious and usually fatal CNS infection caused by JC polyomavirus. Efalizumab, a humanized IgG monoclonal antibody previously approved for treatment of plaque psoriasis, has now been taken off the market due to reactivation of JC virus leading to fatal PML. Thus, use of any currently approved immunosuppressant immunotherapies should be undertaken with caution and with careful monitoring of patients according to FDA guidelines.

Tolerance Induction Specific immunotherapy has moved into a new era with the introduction of soluble CTLA-4 protein into clinical trials. Use of this molecule to block T cell activation via TCR/CD28 ligation during organ or bone marrow transplantation has showed promising results in animals and in early human clinical trials. Specifically, treatment of bone marrow with CTLA-4 protein reduces rejection of the graft in HLA-mismatched bone marrow transplantation. In addition, promising results with soluble CTLA-4 have been reported in the downmodulation of autoimmune T cell responses in the treatment of psoriasis; and it is being studied for treatment of systemic lupus erythematosus (Chap. 378).

Intravenous Immunoglobulin (IVIg) IVIg has been used successfully to block reticuloendothelial cell function and immune complex clearance in various immune cytopenias such as immune thrombocytopenia (Chap. 140). In addition, IVIg is useful for prevention of tissue damage in certain inflammatory syndromes such as Kawasaki disease

(Chap. 385) and as Ig replacement therapy for certain types of immunoglobulin deficiencies (Chap. 374). In addition, controlled clinical trials support the use of IVIg in selected patients with graft-versus-host disease, multiple sclerosis, myasthenia gravis, Guillain-Barré syndrome, and chronic demyelinating polyneuropathy.

Stem Cell Transplantation Hematopoietic stem cell transplantation (SCT) is now being comprehensively studied to treat several autoimmune diseases, including systemic lupus erythematosus, multiple sclerosis, and scleroderma. The goal of immune reconstitution in autoimmune disease syndromes is to replace a dysfunctional immune system with a normally reactive immune cell repertoire. Preliminary results in patients with scleroderma and lupus have showed encouraging results. Controlled clinical trials in these three diseases are now being launched in the United States and Europe to compare the toxicity and efficacy of conventional immunosuppression therapy with that of myeloablative autologous SCT. Recently, SCT was used in the setting of HIV-1 infection. HIV-1 infection of CD4+ T cells requires the presence of surface CD4 receptor and the chemokine receptor 5 (CCR5) co-receptor. Studies have demonstrated that patients who are homozygous for a 32-bp deletion in the CCR5 allele do not express CD4+ T cell CCR5 and thus are resistant to HIV-1 infection with HIV-1 strains that use this co-receptor. Stem cells from a homozygous CCR5 delta32 donor were transplanted to an HIV-infected patient following standard conditioning for such transplants, and the patient has maintained long-term control of the virus without antiretrovirals. Thus, a number of recent insights into immune system function have spawned a new field of interventional immunotherapy and have enhanced the prospect for development of more specific and nontoxic therapies for immune and inflammatory diseases.

373e The Major Histocompatibility Complex

Gerald T. Nepom

THE HLA COMPLEX AND ITS PRODUCTS

The human major histocompatibility complex (MHC), commonly called the human leukocyte antigen (HLA) complex, is a 4-megabase (Mb) region on chromosome 6 (6p21.3) that is densely packed with expressed genes. The best known of these genes are the HLA class I and class II genes, whose products are critical for immunologic specificity and transplantation histocompatibility, and they play a major role in susceptibility to a number of autoimmune diseases. Many other genes in the HLA region are also essential to the innate and antigen-specific functioning of the immune system. The HLA region shows extensive conservation with the MHC of other mammals in terms of genomic organization, gene sequence, and protein structure and function.

The *HLA class I genes* are located in a 2-Mb stretch of DNA at the telomeric end of the HLA region (Fig. 373e-1). The classic (MHC class Ia) HLA-A, -B, and -C loci, the products of which are integral participants in the immune response to intracellular infections, tumors, and allografts, are expressed in all nucleated cells and are highly polymorphic in the population. *Polymorphism* refers to a high degree of allelic variation within a genetic locus that leads to extensive variation between different individuals expressing different alleles. More than 2000 alleles at HLA-A, nearly 3000 alleles at HLA-B, and more than 1700 at HLA-C have been identified in different human populations, making this the most highly polymorphic segment known within the human genome. Each of the alleles at these loci encodes a *heavy chain* (also called an α chain) that associates noncovalently with the nonpolymorphic light chain β_2 -microglobulin, encoded on chromosome 15.

The nomenclature of HLA genes and their products is based on a revised World Health Organization (WHO) nomenclature, in which alleles are given a single designation that indicates locus, allotype, and sequence-based subtype. For example, *HLA-A*02:01* indicates subtype 1 of a group of alleles that encode HLA-A2 molecules. Subtypes that differ from each other at the nucleotide but not the amino acid sequence level are designated by an extra numeral (e.g., *HLA-B*07:02:01* and *HLA-B*07:02:02* are two variants of *HLA-B*07:02*, both encoding the same HLA-B7 molecule). The nomenclature of class II genes, discussed below, is made more complicated by the fact that both chains of a class II molecule are encoded by closely linked HLA-encoded loci, each of which may be polymorphic, and by the presence of differing numbers of isotypic DRB loci in different

individuals. It has become clear that accurate HLA genotyping requires DNA sequence analysis, and the identification of alleles at the DNA sequence level has contributed greatly to the understanding of the role of HLA molecules as peptide-binding ligands, to the analysis of associations of HLA alleles with certain diseases, to the study of the population genetics of HLA, and to a clearer understanding of the contribution of HLA differences to allograft rejection and graft-versus-host disease. Current databases of HLA class I and class II sequences can be accessed by the Internet (e.g., from the IMGT/HLA Database, <http://www.ebi.ac.uk/imgt/hla>), and frequent updates of HLA gene lists are published in several journals.

The biologic significance of this MHC genetic diversity, resulting in extreme variation in the human population, is evident from the perspective of the structure of MHC molecules. As shown in Fig. 373e-2, the MHC class I and class II genes encode MHC molecules that bind small peptides, and together this complex (pMHC; peptide-MHC) forms the ligand for recognition by T lymphocytes, through the antigen-specific T cell receptor (TCR). There is a direct link between the genetic variation and this structural interaction: The allelic changes in genetic sequence result in diversification of the peptide-binding capabilities of each MHC molecule and in differences for specific TCR binding. Thus, different pMHC complexes bind different antigens and are targets for recognition by different T cells.

The class I MHC and class II MHC structures, shown in Fig. 373e-2 B, C, are structurally closely related; however, there are a few key differences. While both bind peptides and present them to T cells, the binding pockets have different shapes, which influence the types of immune responses that result (discussed below). In addition, there are structural contact sites for T cell molecules known as CD8 and CD4, expressed on the class I or class II membrane-proximal domains, respectively. This ensures that when peptide antigens are presented by class I molecules, the responding T cells are predominantly of the CD8 class, and similarly, that T cells responding to class II pMHC complexes are predominantly CD4.

The nonclassic, or class Ib, MHC molecules, HLA-E, -F, and -G, are much less polymorphic than MHC Ia and appear to have distinct functions. The HLA-E molecule has a peptide repertoire displaying signal peptides cleaved from classic MHC class I molecules and is the major self-recognition target for the natural killer (NK) cell-inhibitory receptors NKG2A or NKG2C paired with CD94 (see below and Chap. 372e). This appears to be a function of immune surveillance, because loss of MHC class I signal peptides serves as a surrogate marker for injured or infected cells, leading to release of the inhibitory signal and subsequent activation of NK cells. HLA-E can also bind and present peptides to CD8 T cells, albeit with a limited scope, as only three HLA-E alleles are known. HLA-G is expressed mainly

in stem cells and in extravillous trophoblasts, the fetal cell population directly in contact with maternal tissues. It binds a wide array of peptides, is expressed in six different alternatively spliced forms, and provides inhibitory signals to both NK cells and T cells, presumably in the service of maintaining maternofetal tolerance. Pathologic expression in cancer and infections may also deliver a similar inhibitory immunologic function; 16 HLA-G alleles have been identified. The protein product of HLA-F is found mainly intracellularly, and the function of this locus, which encodes four alleles but has multiple transcriptional variations, remains largely unknown.

Additional class I-like genes have been identified, some HLA-linked and some encoded on other chromosomes, that show only distant homology to the class Ia and Ib molecules but share the three-dimensional class I structure.

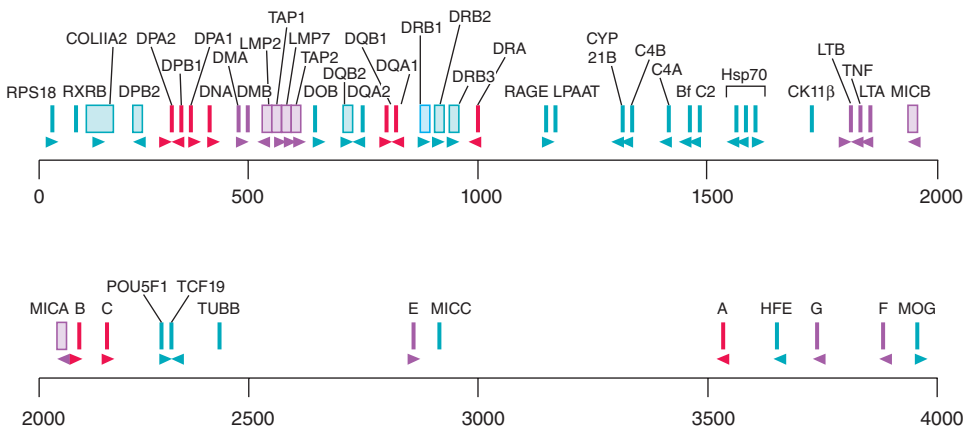


FIGURE 373e-1 Physical map of the HLA region, showing the class I and class II loci, other immunologically important loci, and a sampling of other genes mapped to this region. Gene orientation is indicated by arrowheads. Scale is in kilobase (kb). The approximate genetic distance from DP to A is 3.2 cM. This includes 0.8 cM between A and B (including 0.2 cM between C and B), 0.4–0.8 cM between B and DR-DQ, and 1.6–2.0 cM between DR-DQ and DP.

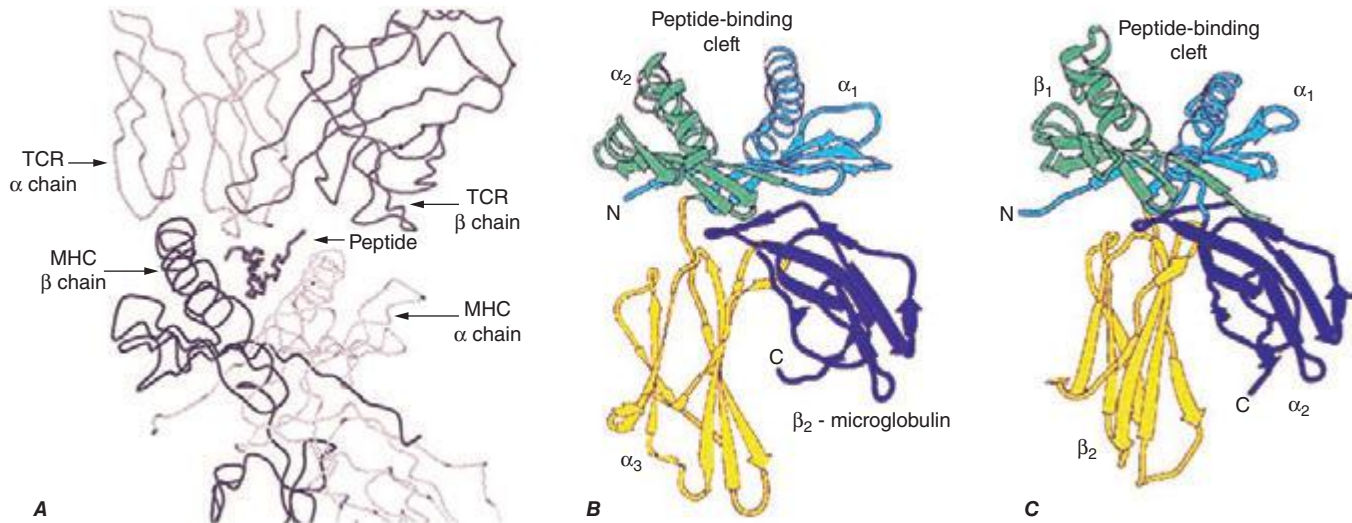


FIGURE 373e-2 A. The trimolecular complex of TCR (*top*), MHC molecule (*bottom*), and a bound peptide form the structural determinants of specific antigen recognition. Other panels (**B** and **C**) show the domain structure of MHC class I (**B**) and class II (**C**) molecules. The α_1 and α_2 domains of class I and the α_1 and β_1 domains of class II form a β -sheet platform that forms the floor of the peptide-binding groove, and α helices that form the sides of the groove. The α_3 (**B**) and β_2 domains (**C**) project from the cell surface and form the contact sites for CD8 and CD4, respectively. (Adapted from EL Reinherz et al: *Science* 286:1913, 1999; and C Janeway et al: *Immunobiology Bookshelf*, 2nd ed. Garland Publishing, New York, 1997; with permission.)

Those on chromosome 6p21 include MIC-A and MIC-B, which are encoded centromeric to HLA-B, and HLA-HFE, located 3 to 4 cM (centi-Morgan) telomeric of HLA-F. MIC-A and MIC-B do not bind peptide but are expressed on gut and other epithelium in a stress-inducible manner and serve as activation signals for certain $\gamma\delta$ T cells, NK cells, CD8 T cells, and activated macrophages, acting through the activating NKG2D receptors. Ninety-one MIC-A and 40 MIC-B alleles are known, and additional diversification comes from variable alanine repeat sequences in the transmembrane domain. Due to this structural diversity, MIC-A can be recognized as a foreign tissue target during organ transplantation, contributing to graft failure. HLA-HFE encodes the gene defective in hereditary hemochromatosis (**Chap. 428**). Among the non-HLA, class I-like genes, CD1 refers to a family of molecules that present glycolipids or other nonpeptide ligands to certain T cells, including T cells with NK activity; FcRn binds IgG within lysosomes and protects it from catabolism (**Chap. 372e**); and Zn- α_2 -glycoprotein 1 binds a nonpeptide ligand and promotes catabolism of triglycerides in adipose tissue. Like the HLA-A, -B, -C, -E, -F, and -G heavy chains, each of which forms a heterodimer with β_2 -microglobulin (Fig. 373e-2), the class I-like molecules, HLA-HFE, FcRn, and CD1 also bind to β_2 -microglobulin, but MIC-A, MIC-B, and Zn- α_2 -glycoprotein 1 do not.

The HLA class II region is also illustrated in Fig. 373e-1. Multiple class II genes are arrayed within the centromeric 1 Mb of the HLA region, forming distinct haplotypes. A *haplotype* refers to an array of alleles at polymorphic loci along a chromosomal segment. Multiple class II genes are present on a single haplotype, clustered into three major subregions: HLA-DR, -DQ, and -DP. Each of these subregions contains at least one functional alpha (A) locus and one functional beta (B) locus. Together these encode proteins that form the α and β polypeptide chains of a mature class II HLA molecule. Thus, the DRA and DRB genes encode an HLA-DR molecule; DQA and DQB genes encode HLA-DQ molecules; and DPA and DPB genes encode HLA-DP molecules. There are several DRB genes (*DRB1*, *DRB2*, *DRB3*, etc.), so that two expressed DR molecules are encoded on most haplotypes by combining the α -chain product of the DRA gene with separate β chains. More than 1000 alleles have been identified at the HLA-DRB1 locus, with most of the variation occurring within limited segments encoding residues that interact with antigens. Detailed analysis of sequences and population distribution of these alleles strongly suggest that this diversity is actively selected by environmental pressures associated with pathogen diversity. In the DQ region, both

DQA1 and DQB1 are polymorphic, with 50 DQA1 alleles and over 300 DQB1 alleles. The current nomenclature is largely analogous to that discussed above for class I, using the convention “locus + allele.”

In addition to allelic polymorphism, products of different DQA alleles can, with some limitations, pair with products of different DQB alleles through both *cis* and *trans* pairing to create combinatorial complexity and expand the number of expressed class II molecules. Because of the enormous allelic diversity in the general population, most individuals are heterozygous at all of the class I and class II loci. Thus, most individuals express six classic class I molecules (two each of HLA-A, -B, and -C) and many class II molecules—two DP, two to four DR, and multiple DQ (both *cis* and *trans* dimers).

OTHER GENES IN THE MHC

In addition to the class I and class II genes themselves, there are numerous genes interspersed among the HLA loci that have interesting and important immunologic functions. Our current concept of the function of MHC genes now encompasses many of these additional genes, some of which are also highly polymorphic. Indeed, direct comparison of the complete DNA sequences for eight of the entire 4-Mb MHC regions from different haplotypes show >44,000 nucleotide variations, encoding an extremely high potential for biologic diversity, and at least 97 genes located in this region are known to have coding region sequence variation. Specific examples include the TAP and LMP genes, as discussed in more detail below, which encode molecules that participate in intermediate steps in the HLA class I biosynthetic pathway. Another set of HLA genes, DMA and DMB, perform an analogous function for the class II pathway. These genes encode an intracellular molecule that facilitates the proper complexing of HLA class II molecules with antigen (see below). The *HLA class III region* is a name given to a cluster of genes between the class I and class II complexes, which includes genes for the two closely related cytokines tumor necrosis factor (TNF)- α and lymphotoxin (TNF- β); the complement components C2, C4, and Bf; heat shock protein (HSP) 70; and the enzyme 21-hydroxylase.

The class I genes HLA-A, -B, and -C are expressed in all nucleated cells, although generally to a higher degree on leukocytes than on nonleukocytes. In contrast, the class II genes show a more restricted distribution: HLA-DR and HLA-DP genes are constitutively expressed on most cells of the myeloid cell lineage, whereas all three class II gene families (HLA-DR, -DQ, and -DP) are inducible by certain stimuli provided by inflammatory cytokines such as interferon γ . Within the

lymphoid lineage, expression of these class II genes is constitutive on B cells and inducible on human T cells. Most endothelial and epithelial cells in the body, including the vascular endothelium and the intestinal epithelium, are also inducible for class II gene expression, and some cells show specialized expression, such as HLA-DQA2 and HLA-DQB2 on Langerhans cells. While somatic tissues normally express only class I and not class II genes, during times of local inflammation, they are recruited by cytokine stimuli to express class II genes as well, thereby becoming active participants in ongoing immune responses. Class II expression is controlled largely at the transcriptional level through a conserved set of promoter elements that interact with a protein known as *CIITA*. Cytokine-mediated induction of *CIITA* is a principal method by which tissue-specific expression of HLA gene expression is controlled. Other HLA genes involved in the immune response, such as TAP and LMP, are also susceptible to upregulation by signals such as interferon γ .

LINKAGE DISEQUILIBRIUM

In addition to extensive polymorphism at the class I and class II loci, another characteristic feature of the HLA complex is *linkage disequilibrium*. This is formally defined as a deviation from Hardy-Weinberg equilibrium for alleles at linked loci. This is reflected in the very low recombination rates between certain loci within the HLA complex. For example, recombination between DR and DQ loci is almost never observed in family studies, and characteristic haplotypes with particular arrays of DR and DQ alleles are found in every population. Similarly, the complement components C2, C4, and Bf are almost invariably inherited together, and the alleles at these loci are found in characteristic haplotypes. In contrast, there is a recombinational hotspot between DQ and DP, which are separated by 1–2 cM of genetic distance, despite their close physical proximity. Certain extended haplotypes encompassing the interval from DQ into the class I region are commonly found, the most notable being the haplotype DR3-B8-A1, which is found, in whole or in part, in 10–30% of northern European whites. It has been hypothesized that selective pressures may maintain linkage disequilibrium in HLA, but this remains to be determined. As discussed below under HLA and immunologic disease, one consequence of the phenomenon of linkage disequilibrium has been the resulting difficulty in assigning HLA-disease associations to a single allele at a single locus.

MHC STRUCTURE AND FUNCTION

Class I and class II molecules display a distinctive structural architecture, which contains specialized functional domains responsible for the unique genetic and immunologic properties of the HLA complex. The principal known function of both class I and class II HLA molecules is to bind antigenic peptides in order to present antigen to an appropriate T cell. The ability of a particular peptide to satisfactorily bind to an individual HLA molecule is a direct function of the molecular fit between the amino acid residues on the peptide with respect to the amino acid residues of the HLA molecule. The bound peptide forms a tertiary structure called the *MHC-peptide complex*, which communicates with T lymphocytes through binding to the TCR molecule. The first site of TCR-MHC-peptide interaction in the life of a T cell occurs in the thymus, where self-peptides are presented to developing thymocytes by MHC molecules expressed on thymic epithelium and hematopoietically derived antigen-presenting cells, which are primarily responsible for positive and negative selection, respectively (Chap. 372e). Thus, the population of MHC-T cell complexes expressed in the thymus shapes the TCR repertoire. Mature T cells encounter MHC molecules in the periphery both in the maintenance of tolerance (Chap. 377e) and in the initiation of immune responses. The MHC-peptide-TCR interaction is the central event in the initiation of most antigen-specific immune responses, since it is the structural determinant of the specificity. For potentially immunogenetic peptides, the ability of a given peptide to be generated and bound by an HLA molecule is a primary feature of whether or not an immune response to that peptide can be generated, and the repertoire of peptides that a particular individual's HLA molecules can bind exerts a major influence over the specificity of that individual's immune response.

When a TCR molecule binds to an HLA-peptide complex, it forms intermolecular contacts with both the antigenic peptide and with the HLA molecule itself. The outcome of this recognition event depends on the density and duration of the binding interaction, accounting for a dual specificity requirement for activation of the T cell. That is, the TCR must be specific both for the antigenic peptide and for the HLA molecule. The polymorphic nature of the presenting molecules, and the influence that this exerts on the peptide repertoire of each molecule, results in the phenomenon of *MHC restriction* of the T cell specificity for a given peptide. The binding of CD8 or CD4 molecules to the class I or class II molecule, respectively, also contributes to the interaction between T cell and the HLA-peptide complex, by providing for the selective activation of the appropriate T cell.

CLASS I STRUCTURE

(Fig. 373e-2B) As noted above, MHC class I molecules provide a cell-surface display of peptides derived from intracellular proteins, and they also provide the signal for self-recognition by NK cells. Surface-expressed class I molecules consist of an MHC-encoded 44-kD glycoprotein heavy chain, a non-MHC-encoded 12-kD light chain β_2 -microglobulin, and an antigenic peptide, typically 8–11 amino acids in length and derived from intracellularly produced protein. The heavy chain displays a prominent peptide-binding groove. In HLA-A and -B molecules, the groove is ~3 nm in length by 1.2 nm in maximum width (30 Å \times 12 Å), whereas it is apparently somewhat wider in HLA-C. Antigenic peptides are noncovalently bound in an extended conformation within the peptide-binding groove, with both N- and C-terminal ends anchored in pockets within the groove (A and F pockets, respectively) and, in many cases, with a prominent kink, or arch, approximately one-third of the way from the N-terminus that elevates the peptide main chain off the floor of the groove.

A remarkable property of peptide binding by MHC molecules is the ability to form highly stable complexes with a wide array of peptide sequences. This is accomplished by a combination of peptide sequence-independent and peptide sequence-dependent bonding. The former consists of hydrogen bond and van der Waals interactions between conserved residues in the peptide-binding groove and charged or polar atoms along the peptide backbone. The latter is dependent upon the six side pockets that are formed by the irregular surface produced by protrusion of amino acid side chains from within the binding groove. The side chains lining the pockets interact with some of the peptide side chains. The sequence polymorphism among different class I alleles and isotypes predominantly affects the residues that line these pockets, and the interactions of these residues with peptide residues constitute the sequence-dependent bonding that confers a particular sequence "motif" on the range of peptides that can bind any given MHC molecule.

CLASS I BIOSYNTHESIS

(Fig. 373e-3A) The biosynthesis of the classic MHC class I molecules reflects their role in presenting endogenous peptides. The heavy chain is cotranslationally inserted into the membrane of the endoplasmic reticulum (ER), where it becomes glycosylated and associates sequentially with the chaperone proteins calnexin and ERp57. It then forms a complex with β_2 -microglobulin, and this complex associates with the chaperone calreticulin and the MHC-encoded molecule tapasin, which physically links the class I complex to TAP, the MHC-encoded transporter associated with antigen processing. Meanwhile, peptides generated within the cytosol from intracellular proteins by the multisubunit, multicatalytic proteasome complex are actively transported into the ER by TAP, where they are trimmed by enzymes known as ER aminopeptidases. At this point, peptides with appropriate sequence complementarity bind specific class I molecules to form complete, folded heavy chain- β_2 -microglobulin-peptide trimer complexes. These are transported rapidly from the ER, through the *cis*- and *trans*-Golgi where the N-linked oligosaccharide is further processed, and thence to the cell surface.

Most of the peptides transported by TAP are produced in the cytosol by proteolytic cleavage of intracellular proteins by the multisubunit, multicatalytic proteasome, and inhibitors of the proteasome dramatically reduce expression of class I-presented antigenic

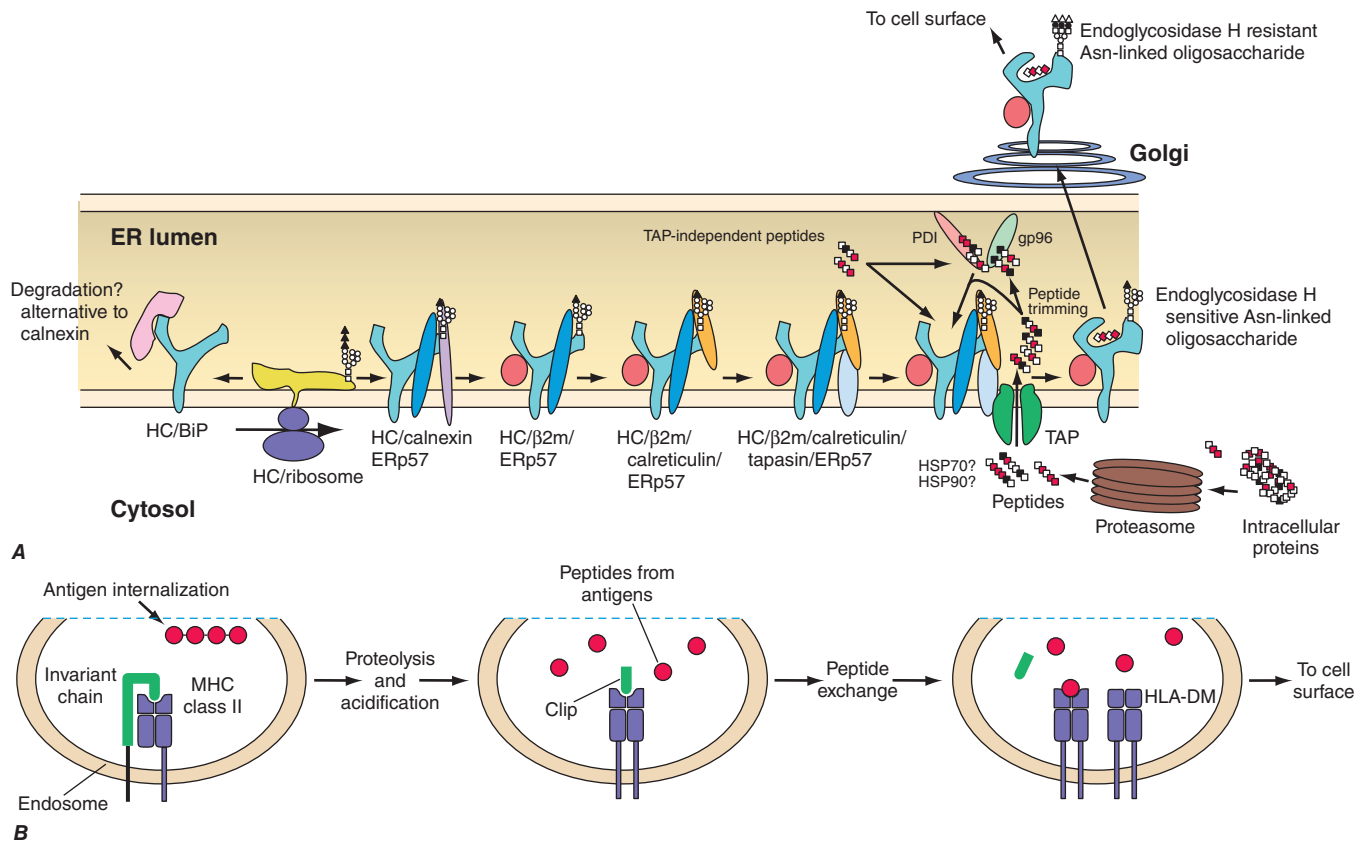


FIGURE 373e-3 Biosynthesis of class I (A) and class II (B) molecules. A. Nascent heavy chain (HC) becomes associated with β_2 -microglobulin (β_2m) and peptide through interactions with a series of chaperones. Peptides generated by the proteasome are transported into the endoplasmic reticulum (ER) by TAP. Peptides undergo N-terminal trimming in the ER and become associated with chaperones, including gp96 and PDI. Once peptide binds to HC- β_2m , the HC- β_2m -peptide trimeric complex exits the ER and is transported by the secretory pathway to the cell surface. In the Golgi, the N-linked oligosaccharide undergoes maturation, with addition of sialic acid residues. Molecules are not necessarily drawn to scale. **B.** Pathway of HLA class II molecule assembly and antigen processing. After transport through the Golgi and post-Golgi compartment, the class II-invariant chain complex moves to an acidic endosome, where the invariant chain is proteolytically cleaved into fragments and displaced by antigenic peptides, facilitated by interactions with the DMA-DMB chaperone protein. This class II molecule-peptide complex is then transported to the cell surface.

peptides. A thiol-dependent oxidoreductase ERp57, which mediates disulfide bond rearrangements, also appears to play an important role in folding the class I-peptide complex into a stable multicomponent molecule. The MHC-encoded proteasome subunits LMP2 and LMP7 may influence the spectrum of peptides produced but are not essential for proteasome function.

CLASS I FUNCTION

Peptide Antigen Presentation On any given cell, a class I molecule occurs in 100,000–200,000 copies and binds several hundred to several thousand distinct peptide species. The vast majority of these peptides are self-peptides to which the host immune system is tolerant by one or more of the mechanisms that maintain tolerance (e.g., clonal deletion in the thymus or clonal anergy or clonal ignorance in the periphery [Chaps. 372e and 377e]). However, class I molecules bearing foreign peptides expressed in a permissive immunologic context activate CD8 T cells, which, if naive, will then differentiate into cytolytic T lymphocytes (CTLs). These T cells and their progeny, through their $\alpha\beta$ TCRs, are then capable of Fas/CD95- and/or perforin-mediated cytotoxicity and/or cytokine secretion (Chap. 372e) upon further encounter with the class I-peptide combination that originally activated it, or other structurally related class I-peptide complexes. As alluded to above, this phenomenon by which T cells recognize foreign antigens in the context of specific MHC alleles is termed *MHC restriction*, and the specific MHC molecule is termed the *restriction element*. The most common source of foreign peptides presented by class I molecules is viral infection, in the course of which peptides from viral proteins

enter the class I pathway. The generation of a strong CTL response that destroys virally infected cells represents an important antigen-specific defense against many viral infections (Chap. 372e). In the case of some viral infections—hepatitis B, for example—CTL-induced target cell apoptosis is thought to be a more important mechanism of tissue damage than any direct cytopathic effect of the virus itself. The importance of the class I pathway in the defense against viral infection is underscored by the identification of a number of viral products that interfere with the normal class I biosynthetic pathway and thus block the immunogenetic expression of viral antigens.

Other examples of intracellularly generated peptides that can be presented by class I molecules in an immunogenic manner include peptides derived from nonviral intracellular infectious agents (e.g., *Listeria*, *Plasmodium*), tumor antigens, minor histocompatibility antigens, and certain autoantigens. There are also situations in which cell surface-expressed class I molecules are thought to acquire and present exogenously derived peptides.

HLA Class I Receptors and NK Cell Recognition (Chap. 372e) NK cells, which play an important role in innate immune responses, are activated to cytotoxicity and cytokine secretion by contact with cells that lack MHC class I expression, and NK cell activation is inhibited by cells that express MHC class I. In humans, the recognition of class I molecules by NK cells is carried out by three classes of receptor families, the killer cell-inhibitory cell receptor (KIR) family, the leukocyte Ig-like receptor (LIR) family, and the CD94/NKG2 family. The KIR family, also called CD158, is encoded on chromosome 19q13.4. KIR gene nomenclature is based on the number of domains (2D or 3D)

and the presence of long (L) or short (S) cytoplasmic domains. The KIR2DL1 and S1 molecules primarily recognize alleles of HLA-C, which possess a lysine at position 80 (HLA-Cw2, -4, -5, and -6), whereas the KIR2DL2/S2 and KIR2DL3/S3 families primarily recognize alleles of HLA-C with asparagine at this position (HLA-Cw1, -3, -7, and -8). The KIR3DL1 and S1 molecules predominantly recognize HLA-B alleles that fall into the HLA-Bw4 class determined by residues 77–83 in the α_1 domain of the heavy chain, whereas the KIR3DL2 molecule is an inhibitory receptor for HLA-A*03. One of the KIR products, KIR2DL4, is known to be an activating receptor for HLA-G. The most common KIR haplotype in whites contains one activating KIR and six inhibitory KIR genes, although there is a great deal of diversity in the population, with >100 different combinations. It appears that most individuals have at least one inhibitory KIR for a self-HLA class I molecule, providing a structural basis for NK cell target specificity, which helps prevent NK cells from attacking normal cells. The importance of KIR-HLA interactions to many immune responses is illustrated by studies associating KIR3DL1 or S1 with multiple sclerosis (Chap. 458), an autoimmune disease, but also with partial protection against HIV (Chap. 226), in both cases consistent with a role for HLA-KIR mediated NK activation. Studies also show an association of KIR2DS1 with protection from relapse following allogeneic bone marrow transplantation in acute myeloid leukemia when these inhibitory receptors in the donors do not recognize the recipient HLA-C.

The LIR gene family (CD85, also called ILT) is encoded centromeric of the KIR locus on 19q13.4, and it encodes a variety of inhibitory immunoglobulin-like receptors expressed on many lymphocyte and other hematopoietic lineages. Interaction of LIR-1 (ILT2) with NK or T cells inhibits activation and cytotoxicity, mediated by many different HLA class I molecules, including HLA-G. HLA-F also appears to interact with LIR molecules, although the functional context for this is not understood.

The third family of NK receptors for HLA is encoded in the NK complex on chromosome 12p12.3-13.1 and consists of CD94 and five NKG2 genes, A/B, C, E/H, D, and F. These molecules are C-type (calcium-binding) lectins, and most function as disulfide-bonded heterodimers between CD94 and one of the NKG2 glycoproteins. The principal ligand of CD94/NKG2A receptors is the HLA-E molecule, complexed to a peptide derived from the signal sequence of classic HLA class I molecules and HLA-G. Thus, analogous to the way in which KIR receptors recognize HLA-C, the NKG2 receptor monitors self-class I expression, albeit indirectly through peptide recognition in the context of HLA-E. NKG2C, -E, and -H appear to have similar specificities but act as activating receptors. NKG2D is expressed as a homodimer and functions as an activating receptor expressed on NK cells, $\gamma\delta$ TCR T cells, and activated CD8 T cells. When complexed with an adaptor called DAP10, NKG2D recognizes MIC-A and MIC-B molecules and activates the cytolytic response. NKG2D also binds a class of molecules known as *ULBP*, structurally related to class I molecules but not encoded in the MHC. **The function of NK cells in immune responses is discussed in Chap. 372e.**

CLASS II STRUCTURE

(Fig. 373e-2C) A specialized functional architecture similar to that of the class I molecules can be seen in the example of a class II molecule depicted in Fig. 373e-2C, with an antigen-binding cleft arrayed above a supporting scaffold that extends the cleft toward the external cellular environment. However, in contrast to the HLA class I molecular structure, β_2 -microglobulin is not associated with class II molecules. Rather, the class II molecule is a heterodimer, composed of a 29-kD α chain and a 34-kD β chain. The amino-terminal domains of each chain form the antigen-binding elements that, like the class I molecule, cradle a bound peptide in a groove bounded by extended α -helical loops, one encoded by the A (α chain) gene and one by the B (β chain) gene. Like the class I groove, the class II antigen-binding groove is punctuated by pockets that contact the side chains of amino acid residues of the bound peptide, but unlike the class I groove, it is open at both ends. Therefore, peptides bound by class II molecules vary greatly in length, since both the N- and C-terminal ends of the peptides can extend

through the open ends of this groove. Approximately 11 amino acids within the bound peptide form intimate contacts with the class II molecule itself, with backbone hydrogen bonds and specific side chain interactions combining to provide, respectively, stability and specificity to the binding (Fig. 373e-4).

The genetic polymorphisms that distinguish different class II genes correspond to changes in the amino acid composition of the class II molecule, and these variable sites are clustered predominantly around the pocket structures within the antigen-binding groove. As with class I, this is a critically important feature of the class II molecule, which explains how genetically different individuals have functionally different HLA molecules.

BIOSYNTHESIS AND FUNCTION OF CLASS II MOLECULES

(Fig. 373e-3B) The intracellular assembly of class II molecules occurs within a specialized compartmentalized pathway that differs dramatically from the class I pathway described above. As illustrated in Fig. 373e-3B, the class II molecule assembles in the ER in association with a chaperone molecule, known as the *invariant chain*. The invariant chain performs at least two roles. First, it binds to the class II molecule and blocks the peptide-binding groove, thus preventing antigenic peptides from binding. This role of the invariant chain appears to account for one of the important differences between class I and class II MHC pathways, since it can explain why class I molecules present endogenous peptides from proteins newly synthesized in the ER but class II molecules generally do not. Second, the invariant chain contains molecular localization signals that direct the class II molecule to traffic into post-Golgi compartments known as *endosomes*, which develop into specialized acidic compartments where proteases cleave the invariant chain, and antigenic peptides can now occupy the class II groove. The specificity and tissue distribution of these proteases appear to be an important way in which the immune system regulates access to the peptide-binding groove and T cells become exposed to specific self-antigens. Differences in protease expression in the thymus and in the periphery may in part determine which specific peptide sequences comprise the peripheral repertoire for T cell recognition. It is at this stage in the intracellular pathway, after cleavage of the invariant chain, that the MHC-encoded DM molecule catalytically facilitates the exchange of peptides within the class II groove to help optimize the specificity and stability of the MHC-peptide complex.

Once this MHC-peptide complex is deposited in the outer cell membrane, it becomes the target for T cell recognition via a specific TCR expressed on lymphocytes. Because the endosome environment contains internalized proteins retrieved from the extracellular environment, the class II-peptide complex often contains bound antigens that were originally derived from extracellular proteins. In this way, the class II peptide-loading pathway provides a mechanism for immune surveillance of the extracellular space. This appears to be an important feature that permits the class II molecule to bind foreign peptides, distinct from the endogenous pathway of class I-mediated presentation.

ROLE OF HLA IN TRANSPLANTATION

The development of modern clinical transplantation in the decades since the 1950s provided a major impetus for elucidation of the HLA system, as allograft survival is highest when donor and recipient are HLA-identical. Although many molecular events participate in transplantation rejection, allogeneic differences at class I and class II loci play a major role. Class I molecules can promote T cell responses in several different ways. In the cases of allografts in which the host and donor are mismatched at one or more class I loci, host T cells can be activated by classic *direct alloreactivity*, in which the antigen receptors on the host T cells react with the foreign class I molecule expressed on the allograft. In this situation, the response of any given TCR may be dominated by the allogeneic MHC molecule, the peptide bound to it, or some combination of the two. Another type of host anti-graft T cell response involves the uptake and processing of donor MHC antigens by host antigen-presenting cells and the subsequent presentation of the resulting peptides by host MHC molecules. This mechanism is termed *indirect alloreactivity*.

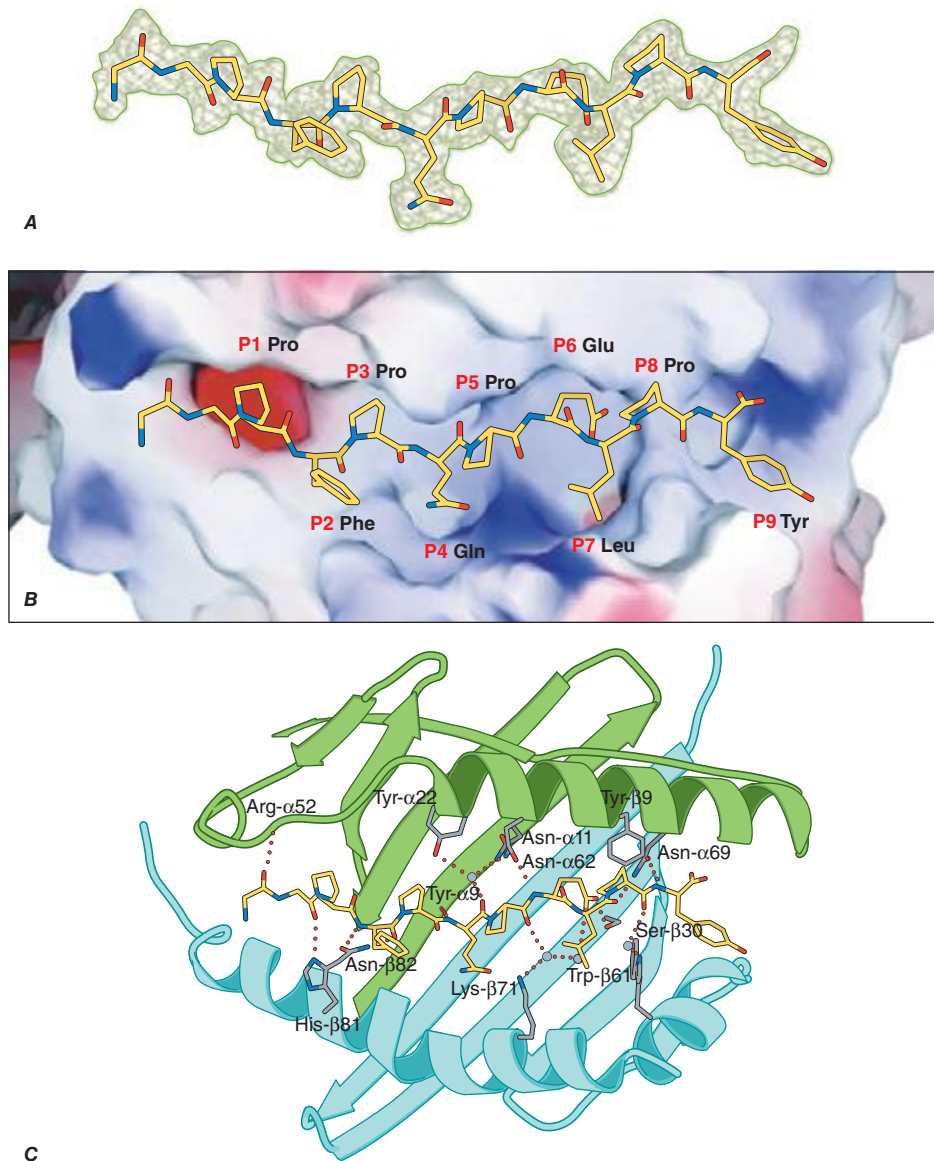


FIGURE 373e-4 Specific intermolecular interactions determine peptide binding to MHC class II molecules. A short peptide sequence derived from alpha-gliadin (**A**) is accommodated within the MHC class II binding groove by specific interactions between peptide side chains (the P1–P9 residues illustrated in **B**) and corresponding pockets in the MHC class II structure. The latter are determined by the genetic polymorphisms of the MHC gene, in this case encoding an HLA-DQ2 molecule (**C**). This shows the extensive hydrogen bond and salt bridge network, which tightly constrains the pMHC complex and presents the complex of antigen and restriction element for CD4 T cell recognition. (From C Kim et al: Structural basis for HLA-DQ2-mediated presentation of gluten epitopes in celiac disease. *Proc Natl Acad Sci USA* 101:4175, 2004.)

In the case of class I molecules on allografts that are shared by the host and the donor, a host T cell response may still be triggered because of peptides that are presented by the class I molecules of the graft but not of the host. The most common basis for the existence of these endogenous antigen peptides, called *minor histocompatibility antigens*, is a genetic difference between donor and host at a non-MHC locus encoding the structural gene for the protein from which the peptide is derived. These loci are termed *minor histocompatibility loci*, and nonidentical individuals typically differ at many such loci. CD4 T cells react to analogous class II variation, both direct and indirect, and class II differences alone are sufficient to drive allograft rejection.

ASSOCIATION OF HLA ALLELES WITH SUSCEPTIBILITY TO DISEASE

It has long been postulated that infectious agents provide the driving force for the allelic diversification seen in the HLA system. An important corollary of this hypothesis is that resistance to specific pathogens may differ between individuals, based on HLA genotype. Observations of specific HLA genes associated with resistance to malaria or dengue

fever, persistence of hepatitis B, and to disease progression in HIV infection are consistent with this model. For example, failure to clear persistent hepatitis B or C viral infection may reflect the inability of particular HLA molecules to present viral antigens effectively to T cells. Similarly, both protective and susceptible HLA allelic associations have been described for human papilloma virus-associated cervical neoplasia, implicating the MHC as an influence in mediating viral clearance in this form of cancer.

Pathogen diversity is probably also the major selective pressure favoring HLA heterozygosity. The extraordinary scope of HLA allelic diversity increases the likelihood that most new pathogens will be recognized by some HLA molecules, helping to ensure immune fitness to the host. However, another consequence of diversification is that some alleles may become capable of recognition of “innocent bystander” molecules, including drugs, environmental molecules, and tissue-derived self-antigens. In a few instances, single HLA alleles display a strong selectivity for binding of a particular agent that accounts for a genetically determined response: Hypersensitivity to

abacavir, an antiretroviral therapeutic, is directly linked to binding of abacavir in the antigen-binding pockets of HLA-B*57:01, where it is buried underneath antigenic peptides and distorts the landscape, changing T cell recognition specificity; an adverse drug reaction to abacavir is more than 500 times more likely to occur in persons with HLA-B*57:01 than in individuals without this HLA allele. Another example is chronic beryllium toxicity, which is linked to binding of beryllium by HLA-DP molecules with a specific glutamic acid polymorphic residue on the class II beta chain. Even in the case of more complex diseases, particular HLA alleles are strongly associated with certain inappropriate immune-mediated disease states, particularly for some common autoimmune disorders (Chap. 377e). By comparing allele frequencies in patients with any particular disease and in control populations, >100 such associations have been identified, some of which are listed in Table 373e-1. The strength of genetic association is reflected in the term *relative risk*, which is a statistical odds ratio representing the risk of disease for an individual carrying a particular genetic marker compared with the risk for individuals in that population without that marker. The nomenclature shown in Table 373e-1 reflects both the HLA serotype (e.g., DR3, DR4) and the HLA genotype (e.g., *DRB1*03:01*, *DRB1*04:01*). It is very likely the class I and class II alleles themselves are the true susceptibility alleles for most of these associations. However, because of the extremely strong linkage disequilibrium between the DR and DQ loci, in some cases it has been difficult to determine the specific locus or combination of class II loci involved. In some cases, the susceptibility gene may be one of the HLA-linked genes located near the class I or class II region, but not the HLA gene itself, and in other cases, the susceptibility gene may be a non-HLA gene such as TNF- α , which is nearby. Indeed, since linkage disequilibrium of some haplotypes extends across large segments of the MHC region, it is quite possible that combinations of genes may account for the particular associations of HLA haplotypes with disease. For example, on some haplotypes associated with rheumatoid arthritis, both HLA-DRB1 alleles and a particular polymorphism associated with the TNF locus may be contributory to disease risk. Other candidates for similar epistatic effects include the IKBL gene and the MICA locus, potentially in combination with classic HLA class II risk alleles.

As might be predicted from the known function of the class I and class II gene products, almost all of the diseases associated with specific HLA alleles have an immunologic component to their pathogenesis. The recent development of soluble HLA-peptide recombinant molecules as biological probes of T cell function, often in multivalent complexes referred to as “MHC tetramers,” represents an opportunity to use HLA genetic associations to develop biomarkers for detection of early disease progression. However, it should be stressed that even the strong HLA associations with disease (those associations with relative risk of ≥ 10) implicate normal, rather than defective, alleles. Most individuals who carry these susceptibility genes do not express the associated disease; in this way, the particular HLA gene is permissive for disease but requires other environmental (e.g., the presence of specific antigens) or genetic factors for full penetrance. In each case studied, even in diseases with very strong HLA associations, the concordance of disease in monozygotic twins is higher than in HLA-identical dizygotic twins or other sibling pairs, indicating that non-HLA genes contribute to susceptibility and can significantly modify the risk attributable to HLA.

Another group of diseases is genetically linked to HLA, not because of the immunologic function of HLA alleles but rather because they are caused by autosomal dominant or recessive abnormal alleles at loci that happen to reside in or near the HLA region. Examples of these are 21-hydroxylase deficiency (Chap. 406), hemochromatosis (Chap. 428), and spinocerebellar ataxia (Chap. 452).

CLASS I ASSOCIATIONS WITH DISEASE

Although the associations of human disease with particular HLA alleles or haplotypes predominantly involve the class II region, there are also several prominent disease associations with class I alleles. These include the association of Behçet’s disease (Chap. 387) with HLA-B51, psoriasis vulgaris (Chap. 71) with HLA-Cw6, and, most notably, the spondyloarthritides (Chap. 384) with HLA-B27. Twenty-five HLA-B

locus alleles, designated *HLA-B*27:01–B*27:25*, encode the family of B27 class I molecules. All of the subtypes share a common B pocket in the peptide-binding groove—a deep, negatively charged pocket that shows a strong preference for binding the arginine side chain. In addition, B27 is among the most negatively charged of HLA class I heavy chains, and the overall preference is for positively charged peptides. *HLA-B*27:05* is the predominant subtype in whites and most other non-Asian populations, and this subtype is very highly associated with ankylosing spondylitis (AS) (Chap. 384), both in its idiopathic form and in association with chronic inflammatory bowel disease or psoriasis vulgaris. It is also associated with reactive arthritis (ReA) (Chap. 384), with other idiopathic forms of peripheral arthritis (undifferentiated spondyloarthropathy), and with recurrent acute anterior uveitis. B27 is found in 50–90% of individuals with these conditions, compared with a prevalence of ~7% in North American whites.

It can be concluded that the B27 molecule itself is involved in disease pathogenesis, based on strong evidence from clinical epidemiology and on the occurrence of a spondyloarthropathy-like disease in HLA-B27 transgenic rats. The association of B27 with these diseases may derive from the specificity of a particular peptide or family of peptides bound to B27 or through another mechanism that is independent of the peptide specificity of B27. In particular, HLA-B27 has been shown to form heavy chain homodimers, utilizing the cysteine residue at position 67 of the B57 α chain, in the absence of β_2 -microglobulin. These homodimers are expressed on the surface of lymphocytes and monocytes from patients with AS, and receptors including KIR3DL1, KIR3DL2, and ILT4 (LILRB2) are capable of binding to them, promoting the activation and survival of cells expressing these receptors. Alternatively, this dimerization “misfolding” of B27 may initiate an intracellular stress signaling response, called the unfolded protein response (UPR), capable of modulating immune cell function, possibly in enthesial-resident T cells that act as sensors of damage and environmental stress.

CLASS II DISEASE ASSOCIATIONS

As can be seen in Table 373e-1, the majority of associations of HLA and disease are with class II alleles. Several diseases have complex HLA genetic associations.

Celiac Disease In the case of celiac disease (Chap. 349), it is probable that the HLA-DQ genes are the primary basis for the disease association. HLA-DQ genes present on both the celiac-associated DR3 and DR7 haplotypes include the *DQB1*02:01* gene, and further detailed studies have documented a specific class II $\alpha\beta$ dimer encoded by the *DQA1*05:01* and *DQB1*02:01* genes, which appears to account for most of the HLA genetic contribution to celiac disease susceptibility. This specific HLA association with celiac disease may have a straightforward explanation: Peptides derived from the wheat gluten component gliadin are bound to the molecule encoded by *DQA1*05:01* and *DQB1*02:01* and presented to T cells. Gliadin-derived peptides that are implicated in this immune activation bind the DQ class II dimer best when the peptide contains a glutamine to glutamic acid substitution. It has been proposed that tissue transglutaminase, an enzyme present at increased levels in the intestinal cells of celiac patients, converts glutamine to glutamic acid in gliadin, creating peptides that are capable of being bound by the DQ2 molecule and presented to T cells.

Pemphigus Vulgaris In the case of pemphigus vulgaris (Chap. 73), there are two HLA genes associated with disease, *DRB1*04:02* and *DQB1*05:03*. Peptides derived from desmoglein-3, an epidermal autoantigen, bind to the *DRB1*04:02*- and *DQB1*05:03*-encoded HLA molecules, and this combination of specific peptide binding and disease-associated class II molecule is sufficient to stimulate desmoglein-specific T cells. A bullous pemphigoid clinical variant, not involving desmoglein recognition, has been found to be associated with *HLA-DQB1*03:01*.

Juvenile Arthritis Pauciarticular juvenile arthritis (Chap. 380) is an autoimmune disease associated with genes at the DRB1 locus and also with genes at the DPB1 locus. Patients with both *DPB1*02:01* and a

TABLE 373e-1 SIGNIFICANT HLA CLASS I AND CLASS II ASSOCIATIONS WITH DISEASE

	Marker	Gene	Strength of Association
Spondyloarthropathies			
Ankylosing spondylitis	B27	<i>B*27:02, -04, -05</i>	++++
Reactive arthritis (Reiter's)	B27		++++
Acute anterior uveitis	B27		+++
Reactive arthritis (<i>Yersinia</i> , <i>Salmonella</i> , <i>Shigella</i> , <i>Chlamydia</i>)	B27		+++
Psoriatic spondylitis	B27		+++
Collagen-Vascular Diseases			
Juvenile arthritis, pauciarticular	DR8		++
	DR5		++
Rheumatoid arthritis	DR4	<i>DRB1*04:01, -04, -05</i>	+++
Sjögren's syndrome	DR3		++
Systemic lupus erythematosus			
White	DR3		+
Japanese	DR2		++
Autoimmune Gut and Skin			
Gluten-sensitive enteropathy (celiac disease)	DQ2	<i>DQA1*05:01</i> <i>DQB1*02:01</i>	+++
Chronic active hepatitis	DR3		++
Dermatitis herpetiformis	DR3		+++
Psoriasis vulgaris	Cw6		++
Pemphigus vulgaris	DR4	<i>DRB1*04:02</i>	+++
	DQ1	<i>DQB1*05:03</i>	
Bullous pemphigoid variant	DQ7	<i>DQB1*03:01</i>	+
Autoimmune Endocrine			
Type 1 diabetes mellitus	DQ8	<i>DQB1*03:02</i>	+++
	DR4	<i>DRB1*04:01, -04</i>	++
	DR3		
	DR2	<i>DQB1*06:02</i>	— ^a
Hyperthyroidism (Graves')	B8		+
	DR3		+
Hyperthyroidism (Japanese)	B35		+
Adrenal insufficiency	DR3		++
Autoimmune Neurologic			
Myasthenia gravis	B8		+
Multiple sclerosis	DR2	<i>DRB1*15:01</i>	+
	DR2	<i>DRB5*01:01</i>	++
Other			
Behçet's disease	B51		++
Congenital adrenal hyperplasia	B47	<i>21-OH (Cyp21B)</i>	+++
Narcolepsy	DR2	<i>DQB1*06:02</i>	++++
Goodpasture's syndrome (anti-GBM)	DR2		++
Abacavir hypersensitivity	B57	<i>B*57:01</i>	++++

^aStrong negative association, i.e., genetic association with protection from diabetes.

Abbreviation: GBM, glomerular basement membrane.

DRB1 susceptibility allele (usually *DRB1*08* or *-05*) have a higher relative risk than expected from the additive effect of those genes alone. In juvenile patients with rheumatoid factor–positive polyarticular disease, heterozygotes carrying both *DRB1*04:01* and *-04:04* have a relative risk >100, reflecting an apparent synergy in individuals inheriting both of these susceptibility genes.

Type 1 Diabetes Mellitus Type 1 (autoimmune) diabetes mellitus (Chap. 417) is associated with MHC genes on more than one haplotype. The presence of both the DR3 and DR4 haplotypes in one individual confers a twentyfold increased risk for type 1 diabetes; the strongest single association is with *DQB1*03:02*, and all haplotypes that carry a *DQB1*03:02* gene are associated with type 1 diabetes, whereas

related haplotypes that carry a different *DQB1* gene are not. However, the relative risk associated with inheritance of this gene can be modified, depending on other HLA genes present either on the same or a second haplotype. For example, the presence of a DR2–positive haplotype containing a *DQB1*06:02* gene is associated with decreased risk. This gene, *DQB1*06:02*, is considered “protective” for type 1 diabetes. Even some DRB1 genes that can occur on the same haplotype as *DQB1*03:02* may modulate risk, so that individuals with the DR4 haplotype that contains *DRB1*04:03* are less susceptible to type 1 diabetes than individuals with other DR4–*DQB1*03:02* haplotypes. There are some characteristic structural features of the diabetes-associated DQ molecule encoded by *DQB1*03:02*, particularly the capability for binding peptides that have negatively charged amino acids near their C-termini. This may indicate a role for specific antigenic peptides or T cell interactions in the immune response to islet-associated proteins.

Although the presence of a DR3 haplotype in combination with the DR4–*DQB1*0302* haplotype is a very high-risk combination for diabetes susceptibility, the specific gene on the DR3 haplotype that is responsible for this synergy is not yet identified.

Rheumatoid Arthritis The HLA genes associated with rheumatoid arthritis (RA) (Chap. 380) encode a distinctive sequence of amino acids from codons 67–74 of the DRβ molecule: RA-associated class II molecules carry the sequence LeuLeuGluGlnArgArgAlaAla or LeuLeuGluGlnLysArgAlaAla in this region, whereas non-RA-associated genes carry one or more differences in this region. These residues form a portion of the molecule that lies in the middle of the α-helical portion of the DRB1-encoded class II molecule, termed the *shared epitope*.

The highest risk for susceptibility to RA comes in individuals who carry both a *DRB1*04:01* and *DRB1*04:04* gene. These DR4-positive RA-associated alleles with the *shared epitope* are most frequent among patients with more severe, erosive disease. Several mechanisms have been proposed that link the shared epitope to immune reactivity in RA. This portion of the class II molecule may allow preferential binding of an arthritogenic peptide, it may favor the expansion of a type of self-reactive T lymphocyte, or it may itself form part of the pMHC ligand recognized by TCR that initiates synovial tissue recognition.

MOLECULAR MECHANISMS FOR HLA-DISEASE ASSOCIATIONS

As noted above, HLA molecules play a key role in the selection and establishment of the antigen-specific T cell repertoire and a major role in the subsequent activation of those T cells during the initiation of an immune response. Precise genetic polymorphisms characteristic of individual alleles dictate the specificity of these interactions and thereby instruct and guide antigen-specific immune events. These same genetically determined pathways are therefore implicated in disease pathogenesis when specific HLA genes are responsible for autoimmune disease susceptibility.

The fate of developing T cells within the thymus is determined by the affinity of interaction between TCR and HLA molecules bearing self-peptides, and thus the particular HLA types of each individual control the precise specificity of the T cell repertoire (Chap. 372e). The primary basis for HLA-associated disease susceptibility may well lie within this thymic maturation pathway. The positive selection of potentially autoreactive T cells, based on the presence of specific HLA susceptibility genes, may establish the threshold for disease risk in a particular individual.

At the time of onset of a subsequent immune response, the primary role of the HLA molecule is to bind peptide and present it to antigen-specific T cells. The HLA complex can therefore be viewed as encoding genetic determinants of precise immunologic activation events. Antigenic peptides that bind particular HLA molecules are capable of stimulating T cell immune responses; peptides that do not bind are not presented to T cells and are not immunogenic. This genetic control of the immune response is mediated by the polymorphic sites within the HLA antigen-binding groove that interact with the bound peptides. In autoimmune and immune-mediated diseases, it is likely that specific tissue antigens that are targets for pathogenic lymphocytes are

complexed with the HLA molecules encoded by specific susceptibility alleles. In autoimmune diseases with an infectious etiology, it is likely that immune responses to peptides derived from the initiating pathogen are bound and presented by particular HLA molecules to activate T lymphocytes that play a triggering or contributory role in disease pathogenesis. The concept that early events in disease initiation are triggered by specific HLA-peptide complexes offers some prospects for therapeutic intervention, since it may be possible to design compounds that interfere with the formation or function of specific HLA-peptide-TCR interactions.

When considering mechanisms of HLA associations with immune response and disease, it is well to remember that just as HLA genetics are complex, so are the mechanisms likely to be heterogeneous. Immune-mediated disease is a multistep process in which one of the HLA-associated functions is to establish a repertoire of potentially reactive T cells, whereas another HLA-associated function is to provide the essential peptide-binding specificity for T cell recognition. For diseases with multiple HLA genetic associations, it is possible that both of these interactions occur and synergize to advance an accelerated pathway of disease.

SECTION 1 THE IMMUNE SYSTEM IN HEALTH AND DISEASE

374 Primary Immune Deficiency Diseases

Alain Fischer

Immunity is intrinsic to life and an important tool in the fight for survival against pathogenic microorganisms. The human immune system can be divided into two major components: the innate immune system and the adaptive immune system (Chap. 372e). The innate immune system provides the rapid triggering of inflammatory responses based on the recognition (at the cell surface or within cells) of either molecules expressed by microorganisms or molecules that serve as “danger signals” released by cells under attack. These receptor/ligand interactions trigger signaling events that ultimately lead to inflammation. Virtually all cell lineages (not just immune cells) are involved in innate immune responses; however, myeloid cells (i.e., neutrophils and macrophages) play a major role because of their phagocytic capacity. The adaptive immune system operates by clonal recognition of antigens followed by a dramatic expansion of antigen-reactive cells and execution of an immune effector program. Most of the effector cells die off rapidly, whereas memory cells persist. Although both T and B lymphocytes recognize distinct chemical moieties and execute distinct adaptive immune responses, the latter is largely dependent on the former in generating long-lived humoral immunity. Adaptive responses utilize components of the innate immune system; for example, the antigen-presentation capabilities of dendritic cells help to determine the type of effector response. Not surprisingly, immune responses are controlled by a series of regulatory mechanisms.

Hundreds of gene products have been characterized as effectors or mediators of the immune system (Chap. 372e). Whenever the expression or function of one of these products is genetically impaired (provided the function is nonredundant), a primary immunodeficiency (PID) occurs.

PIDs are genetic diseases with primarily Mendelian inheritance. More than 250 conditions have now been described, and deleterious mutations in approximately 210 genes have been identified. The overall prevalence of PIDs has been estimated in various countries at 5 per 100,000 individuals; however, given the difficulty in diagnosing these rare and complex diseases, this figure is probably an underestimate. PIDs can involve all possible aspects of immune responses, from innate through adaptive, cell differentiation, and effector function and regulation. For the sake of clarity, PIDs should be classified according to (1) the arm of the immune system that is defective and (2) the mechanism of the defect (when known). Table 374-1 classifies the most prevalent PIDs according to this manner of classification; however, one should bear in mind that the classification of PIDs sometimes involves arbitrary decisions because of overlap and, in some cases, lack of data.

The consequences of PIDs vary widely as a function of the molecules that are defective. This concept translates into multiple levels of vulnerability to infection by pathogenic and opportunistic microorganisms, ranging from extremely broad (as in severe combined immunodeficiency [SCID]) to narrowly restricted to a single microorganism (as in Mendelian susceptibility to mycobacterial disease [MSMD]). The locations of the sites of infection and the causal microorganisms involved will thus help physicians arrive at proper diagnoses. PIDs can also lead to immunopathologic responses such as allergy (as in Wiskott-Aldrich syndrome), lymphoproliferation, and autoimmunity. A combination

Deficiencies of the Innate Immune System

- Phagocytic cells:
 - Impaired production: severe congenital neutropenia (SCN)
 - Asplenia
 - Impaired adhesion: leukocyte adhesion deficiency (LAD)
 - Impaired killing: chronic granulomatous disease (CGD)
- Innate immunity receptors and signal transduction:
 - Defects in Toll-like receptor signaling
 - Mendelian susceptibility to mycobacterial disease
- Complement deficiencies:
 - Classical, alternative, and lectin pathways
 - Lytic phase

Deficiencies of the Adaptive Immune System

- T lymphocytes:
 - Impaired development Severe combined immune deficiencies (SCIDs)
DiGeorge syndrome
 - Impaired survival, migration, function Combined immunodeficiencies
Hyper-IgE syndrome (autosomal dominant)
DOCK8 deficiency
CD40 ligand deficiency
Wiskott-Aldrich syndrome
Ataxia-telangiectasia and other DNA repair deficiencies
- B lymphocytes:
 - Impaired development XL and AR agammaglobulinemia
 - Impaired function Hyper-IgM syndrome
Common variable immunodeficiency (CVID)
IgA deficiency

Regulatory Defects

- Innate immunity Autoinflammatory syndromes (outside the scope of this chapter)
Severe colitis
- Adaptive immunity Hemophagocytic lymphohistiocytosis (HLH)
Autoimmune lymphoproliferation syndrome (ALPS)
Autoimmunity and inflammatory diseases (IPEX, APECED)

Abbreviations: APECED, autoimmune polyendocrinopathy candidiasis ectodermal dysplasia; AR, autosomal recessive; IPEX, immunodysregulation polyendocrinopathy enteropathy X-linked syndrome; XL, X-linked.

of recurrent infections, inflammation, and autoimmunity can be observed in a number of PIDs, thus creating obvious therapeutic challenges. Finally, some PIDs increase the risk of cancer, notably but not exclusively lymphocytic cancers, e.g., lymphoma.

DIAGNOSIS OF PRIMARY IMMUNODEFICIENCIES

The most frequent symptom prompting the diagnosis of a PID is the presence of recurrent or unusually severe infections. As mentioned above, recurrent allergic or autoimmune manifestations may also alert the physician to a possible diagnosis of PID. In such cases, a detailed account of the subject's personal and family medical history should be obtained. It is of the utmost importance to gather as much medical information as possible on relatives and up to several generations of ancestors. In addition to the obvious focus on primary symptoms, the clinical examination should evaluate the size of lymphoid organs and, when appropriate, look for the characteristic signs of a number of complex syndromes that may be associated with a PID.

The performance of laboratory tests should be guided to some extent by the clinical findings. Infections of the respiratory tract (bronchi, sinuses) mostly suggest a defective antibody response. In general, invasive bacterial infections can result from complement deficiencies, signaling defects of innate immune responses, asplenia, or defective antibody responses. Viral infections, recurrent *Candida*

infections, and opportunistic infections are generally suggestive of impaired T cell immunity. Skin infections and deep-seated abscesses primarily reflect innate immune defects (such as chronic granulomatous disease); however, they may also appear in the autosomal dominant hyper-IgE syndrome. Table 374-2 summarizes the laboratory tests that are most frequently used to diagnose a PID. More specific tests (notably genetic tests) are then used to make a definitive diagnosis.

The PIDs discussed below have been grouped together according to the affected cells and the mechanisms involved (Table 374-1, Fig. 374-1).

PRIMARY IMMUNODEFICIENCIES OF THE INNATE IMMUNE SYSTEM

PIDs of the innate immune system are relatively rare and account for approximately 10% of all PIDs.

SEVERE CONGENITAL NEUTROPENIA

Severe congenital neutropenia (SCN) consists of a group of inherited diseases that are characterized by severely impaired neutrophil counts (<500 polymorphonuclear leukocytes [PMN]/ μ L of blood). The condition is usually manifested from birth. SCN may also be cyclic (with a 3-week periodicity), and other neutropenia syndromes can also be intermittent. Although the most frequent inheritance pattern for SCN is autosomal dominant, autosomal recessive and X-linked recessive conditions also exist. Bacterial infections at the interface between the body and the external milieu (e.g., the orifices, wounds, and the respiratory tract) are common manifestations. Bacterial infections can rapidly progress through soft tissue and are followed by dissemination in the bloodstream. Severe visceral fungal infections can also ensue. The absence of pus is a hallmark of this condition.

Diagnosis of SCN requires examination of the bone marrow. Most SCNs are associated with a block in granulopoiesis at the promyelocytic stage (Fig. 374-1). SCN has multiple etiologies, and to date, mutations in 11 different genes have been identified. Most of these mutations result in isolated SCN, whereas others are syndromic (Chap. 80). The most frequent forms of SCN are caused by the premature cell death of granulocyte precursors, as observed in deficiencies of GFI1, HAX1, and elastase 2 (*ELANE*), with the latter accounting for 50% of SCN sufferers. Certain *ELANE* mutations cause cyclic neutropenia syndrome. A gain-of-function mutation in the *WASP* gene (see the section on "Wiskott-Aldrich syndrome" below) causes X-linked SCN, which is also associated with monocytopenia.

As mentioned above, SCN exposes the patient to life-threatening, disseminated bacterial and fungal infections. Treatment requires careful hygiene measures, notably in infants. Later in life, special oral and dental care is essential, along with the prevention of bacterial infection by prophylactic administration of trimethoprim/sulfamethoxazole. Subcutaneous injection of the cytokine granulocyte colony-stimulating factor (G-CSF) usually improves neutrophil development and thus prevents infection in most SCN diseases. However, there are two caveats: (1) a few cases of SCN with *ELANE* mutation are refractory to G-CSF and may require curative treatment via allogeneic hematopoietic stem cell transplantation (HSCT); and (2) a subset of G-CSF-treated patients carrying *ELANE* mutations are at a greater risk of developing acute myelogenous leukemia associated (in most cases) with somatic gain-of-function mutations of the G-CSF receptor gene.

ASPLENIA

Primary failure of the development of a spleen is an extremely rare disease that can be either syndromic (in Ivemark syndrome) or isolated with an autosomal dominant expression; in the latter case, mutations in the ribosomal protein SA gene were recently found. Due to the

TABLE 374-2 TESTS MOST FREQUENTLY USED TO DIAGNOSE A PRIMARY IMMUNE DEFICIENCY (PID)

Test	Information	PID Disease
• Blood cell counts and cell morphology	Neutrophil counts ^a	↓ Severe congenital neutropenia, ↑↑ LAD
	Lymphocyte counts ^a	T cell ID
	Eosinophilia	WAS, hyper-IgE syndrome
	Howell-Jolly bodies	Asplenia
• Chest x-ray	Thymic shadow	SCID, DiGeorge syndrome
	Costochondral junctions	Adenosine deaminase deficiency
• Bone x-ray	Metaphyseal ends	Cartilage hair hypoplasia
• Immunoglobulin serum levels	IgG, IgA, IgM	B cell ID
	IgE	Hyper-IgE syndrome, WAS, T cell ID
• Lymphocyte phenotype	T, B lymphocyte counts	T cell ID, agammaglobulinemia
• Dihydrorhodamine fluorescence (DHR) assay Nitroblue tetrazolium (NBT) assay	Reactive oxygen species production by PMNs	Chronic granulomatous disease
• CH50, AP50	Classic and alternative complement pathways	Complement deficiencies
• Ultrasonography of the abdomen	Spleen size	Asplenia

^aNormal counts vary with age. For example, the lymphocyte count is between 3000 and 9000/μL of blood below the age of 3 months and between 1500 and 2500/μL in adults.

Abbreviations: ID, immunodeficiency; LAD, leukocyte adhesion deficiency; PMNs, polymorphonuclear leukocytes; SCID, severe combined immunodeficiency; WAS, Wiskott-Aldrich syndrome.

absence of natural filtration of microbes in the blood, asplenia predisposes affected individuals to fulminant infections by encapsulated bacteria. Although most infections occur in the first years of life, cases may also arise in adulthood. The diagnosis is confirmed by abdominal ultrasonography and the detection of Howell-Jolly bodies in red blood cells. Effective prophylactic measures (twice-daily oral penicillin and appropriate vaccination programs) usually prevent fatal outcomes.

GATA2 DEFICIENCY

Recently an immunodeficiency combining monocytopenia and dendritic and lymphoid (B and natural killer [NK]) cell deficiency (DCML), also called monocytopenia with nontuberculous mycobacterial infections (mono-MAC), has been described as a consequence of a dominant mutation in the gene *GATA2*, a transcription factor involved in hematopoiesis. This condition also predisposes to lymphedema,

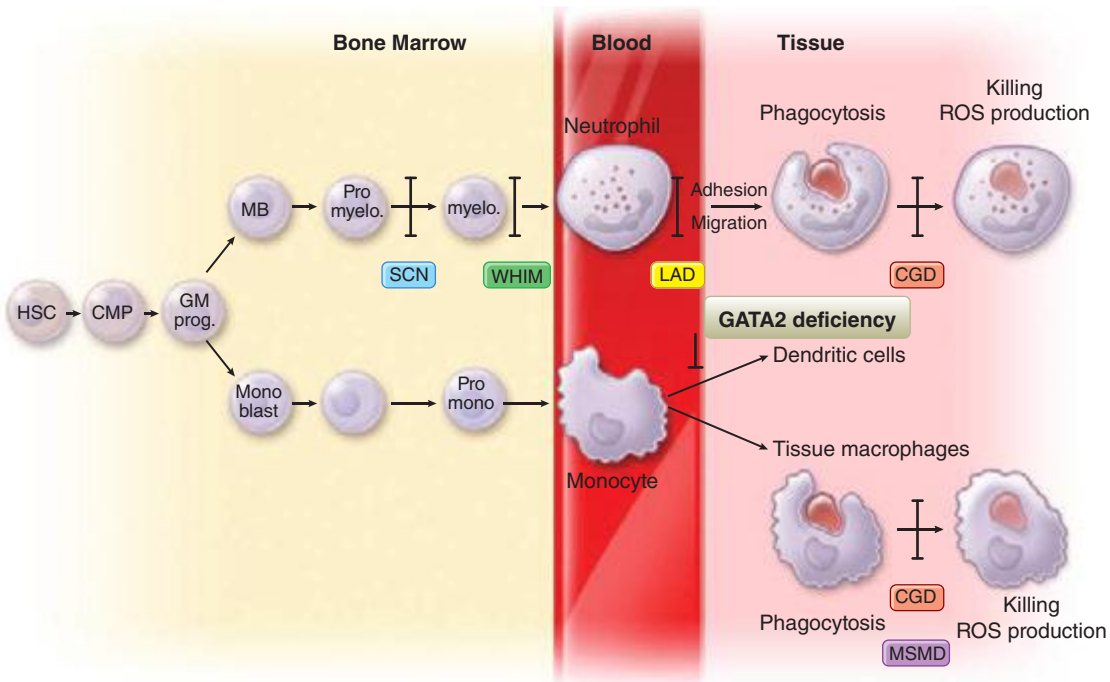


FIGURE 374-1 Differentiation of phagocytic cells and related primary immunodeficiencies (PIDs). Hematopoietic stem cells (HSCs) differentiate into common myeloid progenitors (CMPs) and then granulocyte-monocyte progenitors (GM-prog.), which, in turn, differentiate into neutrophils (MB: myeloblasts; Pro myelo: promyelocytes; myelo: myelocytes) or monocytes (monoblasts and promonocytes). Upon activation, neutrophils adhere to the vascular endothelium, transmigrate, and phagocytose the targets. Reactive oxygen species (ROS) are delivered to the microorganism-containing phagosomes. Macrophages in tissues kill using the same mechanism. Following activation by interferon γ (not shown here), macrophages can be armed to kill intracellular pathogens such as mycobacteria. For sake of simplicity, not all cell differentiation stages are shown. The abbreviations for PIDs are contained in boxes placed at corresponding stages of the pathway. CGD, chronic granulomatous diseases; GATA2, zinc finger transcription factor; LAD, leukocyte adhesion deficiencies; MSMD, Mendelian susceptibility to mycobacterial disease; SCN, severe congenital neutropenia; WHIM, warts, hypogammaglobulinemia, infections, and myelokathexis.

2106 myelodysplasia, and acute myeloid leukemia. Infections (bacterial and viral) are life-threatening, thus indicating, together with the malignant risk, HSCT.

LEUKOCYTE ADHESION DEFICIENCY (LAD)

Leukocyte adhesion deficiency (LAD) consists of three autosomal recessive conditions (LAD I, II, and III) (**Chap. 80**). The most frequent condition (LAD I) is caused by mutations in the $\beta 2$ integrin gene; following leukocyte activation, $\beta 2$ integrins mediate adhesion to inflamed endothelium expressing cognate ligands. LAD III results from a defect in a regulatory protein (kindlin, also known as Fermt 3) involved in activating the ligand affinity of $\beta 2$ integrins. The extremely rare LAD II condition is the end result of a defect in selectin-mediated leukocyte rolling that occurs prior to $\beta 2$ integrin binding. There is a primary defect in fucose transporter such that oligosaccharide selectin ligands are missing in this syndromic condition.

Given that neutrophils are not able to reach infected tissues, LAD renders the individual susceptible to bacterial and fungal infections in a way that is similar to that of patients with SCN. LAD also causes impaired wound healing and delayed loss of the umbilical cord. A diagnosis can be suspected in cases of pus-free skin/tissue infections and massive hyperleukocytosis ($>30,000/\mu\text{L}$) in the blood (mostly granulocytes). Patients with LAD III also develop bleeding because the $\beta 2$ integrin in platelets is not functional. Use of immunofluorescence and functional assays to detect $\beta 2$ integrin can help form a diagnosis. Severe forms of LAD may require HSCT, although gene therapy is also now being considered. Neutrophil-specific granule deficiency (a very rare condition caused by a mutation in the gene for transcription factor C/EBP α) results in a condition that is clinically similar to LAD.

CHRONIC GRANULOMATOUS DISEASES

Chronic granulomatous diseases (CGDs) are characterized by impaired phagocytic killing of microorganisms by neutrophils and macrophages (**Chap. 80**). The incidence is approximately 1 per 200,000 live births. About 70% of cases are associated with X-linked recessive inheritance versus autosomal inheritance in the remaining 30%. CGD causes deep-tissue bacterial and fungal abscesses in macrophage-rich organs such as the lymph nodes, liver, and lungs. Recurrent skin infections (such as folliculitis) are common and can prompt an early diagnosis of CGD. The infectious agents are typically catalase-positive bacteria (such as *Staphylococcus aureus* and *Serratia marcescens*) but also include *Burkholderia cepacia*, pathogenic mycobacteria (in certain regions of the world), and fungi (mainly filamentous molds, such as *Aspergillus*).

CGD is caused by defective production of reactive oxygen species (ROS) in the phagolysosome membrane following phagocytosis of microorganisms. It results from the lack of a component of NADPH oxidase (gp91phox or p22phox) or of the associated adapter/activating proteins (p47phox, p67phox, or p40phox) that mediate the transport of electrons into the phagolysosome for creating ROS by interaction with O_2 . Under normal circumstances, these ROS either directly kill engulfed microorganisms or enable the rise in pH needed to activate the phagosomal proteases that contribute to microbial killing. Diagnosis of CGD is based on assays of ROS production in neutrophils and monocytes (Table 374-2). As its name suggests, CGD is also a granulomatous disease. Macrophage-rich granulomas can often arise in the liver, spleen, and other organs. These are sterile granulomas that cause disease by obstruction (bladder, pylorus, etc.) or inflammation (colitis, restrictive lung disease).

The management of infections in patients with CGD can be a complex process. The treatment of bacterial infections is generally based on combination therapy with antibiotics that are able to penetrate into cells. The treatment of fungal infections requires aggressive, long-term use of antifungals. Inflammatory/granulomatous lesions are usually steroid-sensitive; however, glucocorticoids often contribute to the spread of infections. Hence, there is strong need for new therapeutic options in what is still a poorly understood disease.

The treatment of CGD mostly relies on preventing infections. It has been unambiguously demonstrated that prophylactic usage of trimethoprim/sulfamethoxazole is both well tolerated and highly effective

in reducing the risk of bacterial infection. Daily administration of azole derivatives (notably itraconazole) also reduces the frequency of fungal complications. It has long been suggested that interferon γ administration is helpful, although medical experts continue to disagree over this controversial issue. Patients may do reasonably well with prophylaxis and careful management. However, other patients develop severe and persistent fungal infections and/or chronic inflammatory complications that ultimately require HSCT. The latter is an established curative approach for CGD; however, the risk-versus-benefit ratio must be carefully assessed on a case-by-case basis. Gene therapy approaches are also being evaluated.

MENDELIAN SUSCEPTIBILITY TO MYCOBACTERIAL DISEASE (MSMD)

This group of diseases is characterized by a defect in the interleukin-12 (IL-12)–interferon (IFN) γ axis (including IL-12p40, IL-12 receptor [R] β_1 , IFN- γ R $_1$ and R $_2$, STAT1, IRF8 and ISG515 deficiencies), which ultimately leads to impaired IFN- γ -dependent macrophage activation. Both recessive and dominant inheritance modes have been observed. The hallmark of this PID is a specific and narrow vulnerability to tuberculous and nontuberculous mycobacteria. The most severe phenotype (as observed in complete IFN- γ receptor deficiency) is characterized by disseminated infection that can be fatal even when aggressive and appropriate antimycobacterial therapy is applied. In addition to mycobacterial infections, MSMD patients (and particularly those with an IL-12/IL-12 R deficiency) are prone to developing *Salmonella* infections. Although MSMDs are very rare, they should be considered in any patient with persistent mycobacterial infection. Treatment with IFN- γ may efficiently bypass an IL-12/IL-12R deficiency.

TOLL-LIKE RECEPTOR (TLR) PATHWAY DEFICIENCIES

In a certain group of patients with early-onset, invasive *Streptococcus pneumoniae* infections or (less frequently) *Staphylococcus aureus* or other pyogenic infections, conventional screening for PIDs does not identify the cause of the defect in host defense. It has been established that these patients carry recessive mutations in genes that encode essential adaptor molecules (IRAK4 and MYD88) involved in the signaling pathways of the majority of known Toll-like receptors (TLRs) (**Chap. 372e**). Remarkably, susceptibility to infection appears to decrease after the first few years of life—perhaps an indication that adaptive immunity (once triggered by an initial microbial challenge) is then able to prevent recurrent infections.

Certain TLRs (TLR-3, -7, -8, and -9) are involved in the recognition of RNA and DNA and usually become engaged during viral infections. Very specific susceptibility to herpes simplex encephalitis has been described in patients with a deficiency in Unc93b (a molecule associated with TLR-3, -7, -8, and -9 required for correct subcellular localization), TLR-3, or associated signaling molecules TRIF, TBK1, and TRAF3, resulting in defective type I IFN production. The fact that no other TLR deficiencies have been found—despite extensive screening of patients with unexplained, recurrent infections—strongly suggests that these receptors are functionally redundant. Hypomorphic mutations in NEMO/IKK- γ (a member of the NF- κ B complex, which is activated downstream of TLR receptors) lead to a complex, variable immunodeficiency and a number of associated features. Susceptibility to both invasive, pyogenic infections and mycobacteria may be observed in this particular setting.

COMPLEMENT DEFICIENCY

The complement system is composed of a complex cascade of plasma proteins (**Chap. 372e**) that leads to the deposition of C3b fragments on the surface of particles and the formation of immune complexes that can culminate in the activation of a lytic complex at the bacterial surface. C3 cleavage can be mediated via three pathways: the classic, alternate, and lectin pathways. C3b coats particles as part of the opsonization process that facilitates phagocytosis following binding to cognate receptors. A deficiency in any component of the classic pathway (C1q, C1r, C1s, C4, and C2) can predispose an individual to bacterial infections that are tissue-invasive or that occur in the respiratory tract. Likewise, a C3 deficiency or a deficiency in factor I (a protein that

regulates C3 consumption, thus leading to a C3 deficiency due to its absence) also results in the same type of vulnerability to infection. It has recently been reported that a very rare deficiency in ficolin-3 predisposes affected individuals to bacterial infections. Deficiencies in the alternative pathway (factors D and properdin) are associated with the occurrence of invasive *Neisseria* infections.

Lastly, deficiencies of any complement component involved in the lytic phase (C5, C6, C7, C8, and, to a lesser extent, C9) predispose affected individuals to systemic infection by *Neisseria*. This is explained by the critical role of complement in the lysis of the thick cell wall possessed by this class of bacteria.

Diagnosis of a complement deficiency relies primarily on testing the status of the classic and alternate pathway via functional assays, i.e., the CH50 and AP50 tests, respectively. When either pathway is profoundly impaired, determination of the status of the relevant components in that pathway enables a precise diagnosis. Appropriate vaccinations and daily administration of oral penicillin are efficient means of preventing recurrent infections. It is noteworthy that several complement deficiencies (in the classic pathway and the lytic phase) may also predispose affected individuals to autoimmune diseases (notably systemic lupus erythematosus; [Chap. 378](#)).

PRIMARY IMMUNODEFICIENCIES OF THE ADAPTIVE IMMUNE SYSTEM

T LYMPHOCYTE DEFICIENCIES (TABLE 374-1, FIGS. 374-2 AND 374-3)

Given the central role of T lymphocytes in adaptive immune responses ([Chap. 372e](#)), PIDs involving T cells generally have severe pathologic consequences; this explains the poor overall prognosis and the need for early diagnosis and the early intervention with appropriate therapy. Several differentiation pathways of T cell effectors have been described, one or all of which may be affected by a given PID (Fig. 374-2). Follicular helper CD4⁺ T cells in germinal centers are required for T-dependent antibody production, including the generation of Ig class-switched, high-affinity antibodies. CD4⁺ T_H1 cells provide cytokine-dependent (mostly IFN- γ -dependent) help to macrophages for intracellular killing of various microorganisms, including mycobacteria and *Salmonella*. CD4⁺ T_H2 cells produce IL-4, IL-5, and IL-13 and thus recruit and activate eosinophils and other cells required to fight helminth infections. CD4⁺ T_H17 cells produce IL-17 and IL-22 cytokines that recruit neutrophils to the skin and lungs to fight bacterial and fungal infections. Cytotoxic CD8⁺ T cells can kill infected cells, notably in the context of viral infections. In addition, certain T cell deficiencies predispose affected individuals to *Pneumocystis jiroveci* lung infections early in life and to chronic gut/biliary duct/liver infections by *Cryptosporidium* and related genera later on in life. Lastly, naturally occurring or induced regulatory T cells are essential for controlling inflammation (notably reactivity to commensal bacteria in the gut) and autoimmunity. The role of other T cell subsets with limited T cell receptor (TCR) diversity (such as $\gamma\delta$ TCR T cells or natural killer T [NKT] cells) in PIDs is less well known; however, these subsets can be defective in certain PIDs, and this finding can sometimes contribute to the diagnosis (e.g., NKT cell deficiency in X-linked proliferative syndrome). T cell deficiencies account for approximately 20% of all cases of PID.

Severe Combined Immunodeficiencies Severe combined immunodeficiencies (SCIDs) constitute a group of rare PIDs characterized by a profound block in T cell development and thus the complete absence of these cells. The developmental block is always the consequence of an intrinsic deficiency. The incidence of SCID is estimated to be 1 in 50,000 live births. Given the severity of the T cell deficiency, clinical consequences occur early in life (usually within 3 to 6 months of birth). The most frequent clinical manifestations are recurrent oral candidiasis, failure to thrive, and protracted diarrhea and/or acute interstitial pneumonitis caused by *Pneumocystis jiroveci* (although the latter can also be observed in the first year of life in children with B cell deficiencies). Severe viral infections or invasive bacterial infections can also occur. Patients may also experience complications related to infections caused by live vaccines (notably bacille Calmette-Guérin [BCG]) that may lead not only to local and regional infection but also to disseminated infection manifested by fever, splenomegaly, and skin and lytic bone lesions. A scaly skin eruption can be observed in a context of maternal T cell engraftment (see below). A diagnosis of SCID can be suspected based on the patient's clinical history and, possibly, a family history of deaths in very young children (suggestive of either X-linked or recessive inheritance). Lymphocytopenia is strongly suggestive of SCID in more than 90% of cases ([Table 316-2](#)). The absence of a thymic shadow on a chest x-ray can also be suggestive of SCID. An accurate diagnosis relies on precise determination of the number of circulating T, B, and NK lymphocytes and their subsets. T cell lymphopenia may be masked in some patients by the presence of maternal T cells

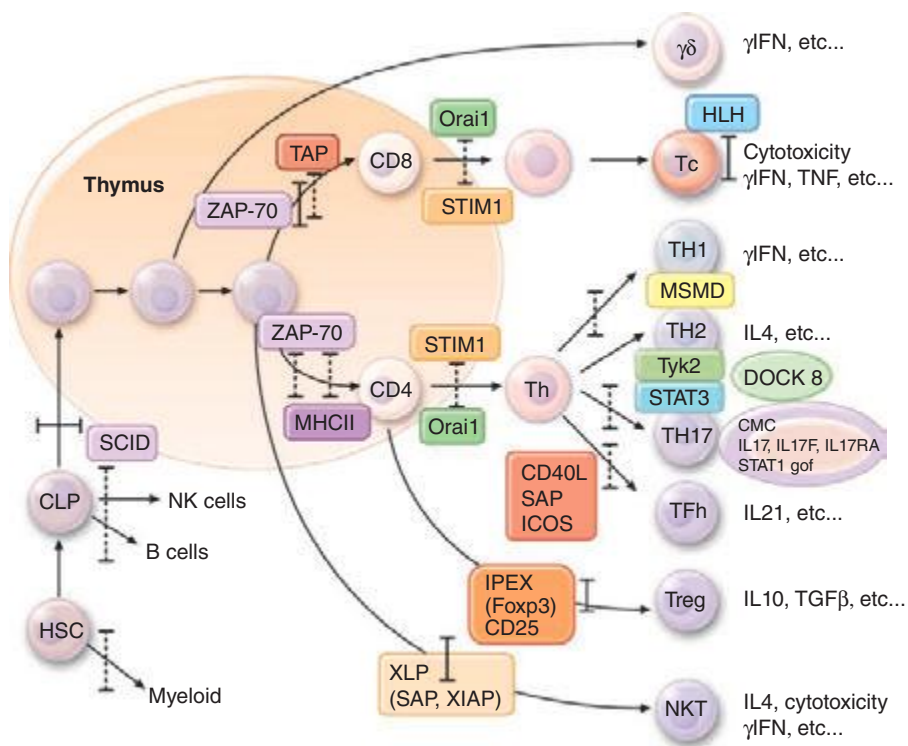


FIGURE 374-2 T cell differentiation, effector pathways, and related primary immunodeficiencies (PIDs). Hematopoietic stem cells (HSCs) differentiate into common lymphoid progenitors (CLPs), which, in turn, give rise to the T cell precursors that migrate to the thymus. The development of CD4⁺ and CD8⁺ T cells is shown. Known T cell effector pathways are indicated, i.e., $\gamma\delta$ cells, cytotoxic T cells (Tc), T_H1, T_H2, T_H17, Tfh (follicular helper) CD4 effector T cells, regulatory T cells (Treg), and natural killer T cells (NKTs); abbreviations for PIDs are contained in boxes. Vertical bars indicate a complete deficiency; broken bars indicate a partial deficiency. SCID, severe combined immunodeficiency; ZAP-70, zeta-associated protein deficiency; MHCII, major histocompatibility complex class II deficiency; TAP, TAP1 and TAP2 deficiencies; Orai1, STIM1 deficiencies; HLH, hematopoietic lymphohistiocytosis; MSMD, Mendelian susceptibility to mycobacterial disease; Tyk2, DOCK8, autosomal recessive form of hyper-IgE syndrome; STAT3, autosomal dominant form of hyper-IgE syndrome; IL17F, IL17RA, STAT1 (gof: gain of function), CMC (chronic mucocutaneous candidiasis), CD40L, ICOS, SAP deficiencies; IPEX, immunodysregulation polyendocrinopathy enteropathy X-linked syndrome; XLP, X-linked proliferative syndromes.

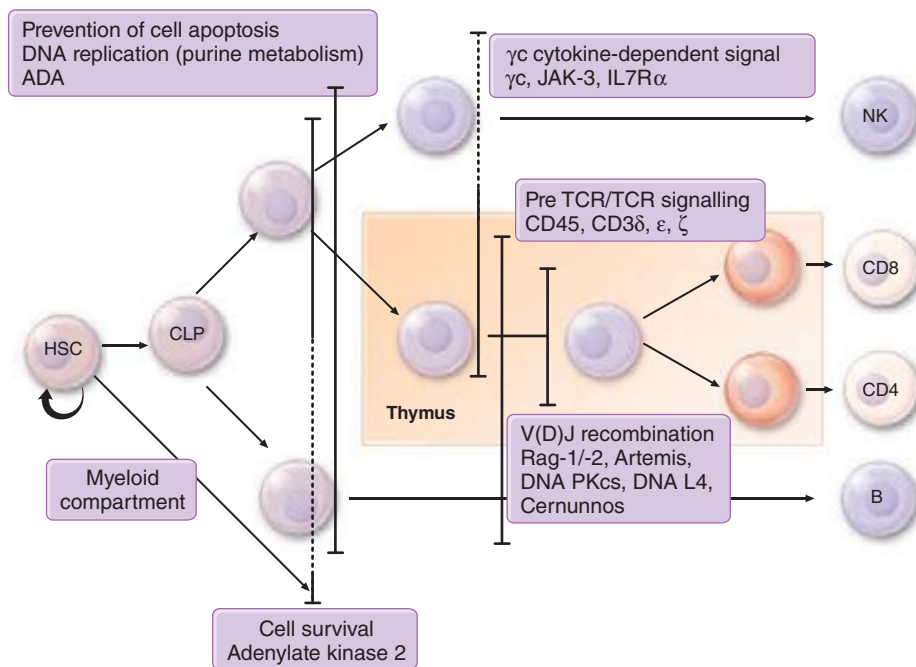


FIGURE 374-3 T cell differentiation and severe combined immunodeficiencies (SCIDs). The vertical bars indicate the five mechanisms currently known to lead to SCID. The names of deficient proteins are indicated in the boxes adjacent to the vertical bars. A broken line means that deficiency is partial or involves only some of the indicated immunodeficiencies. ADA, adenosine deaminase deficiency; CLPs, common lymphoid progenitors; DNAL4, DNA ligase 4; HSCs, hematopoietic stem cells; NKs, natural killer cells; TCR, T cell receptor.

(derived from maternal-fetal blood transfers) that cannot be eliminated. Although counts are usually low (<500/ μ L of blood), higher maternal T cell counts may, under some circumstances, initially mask the presence of SCID. Thus, screening for maternal cells by using adequate genetic markers should be performed whenever necessary. Inheritance pattern analysis and lymphocyte phenotyping can discriminate between various forms of SCID and provide guidance in the choice of accurate molecular diagnostic tests (see below). To date, five distinct causative mechanisms for SCID (Fig. 374-3) have been identified:

SEVERE COMBINED IMMUNODEFICIENCY CAUSED BY A CYTOKINE-SIGNALING DEFICIENCY The most frequent SCID phenotype (accounting for 40–50% of all cases) is the absence of both T and NK cells. This outcome results from a deficiency in either the common γ chain (γ c) receptor that is shared by several cytokine receptors (the IL-2, -4, -7, -9, -15, and -21 receptors) or Jak-associated kinase (JAK) 3 that binds to the cytoplasmic portion of the γ c chain receptor and induces signal transduction following cytokine binding. The former form of SCID (γ c deficiency) has an X-linked inheritance mode, while the second is autosomal recessive. A lack of the IL-7R α chain (which, together with γ c, forms the IL-7 receptor) induces a selective T cell deficiency.

PURINE METABOLISM DEFICIENCY Ten to 20% of SCID patients exhibit a deficiency in adenosine deaminase (ADA), an enzyme of purine metabolism that deaminates adenosine (ado) and deoxyadenosine (dAdo). An ADA deficiency results in the accumulation of ado and dAdo metabolites that induce premature cell death of lymphocyte progenitors. The condition results in the absence of B and NK lymphocytes as well as T cells. The clinical expression of complete ADA deficiency typically occurs very early in life. Since ADA is a ubiquitous enzyme, its deficiency can also cause bone dysplasia with abnormal costochondral junctions and metaphyses (found in 50% of cases) and neurologic defects. The very rare purine nucleoside phosphorylase (PNP) deficiency causes a profound although incomplete T cell deficiency that is often associated with severe neurologic impairments.

DEFECTIVE REARRANGEMENTS OF T AND B CELL RECEPTORS A series of SCID conditions are characterized by a selective deficiency in T and B lymphocytes with autosomal recessive inheritance. These conditions

account for 20–30% of SCID cases and result from mutations in genes encoding proteins that mediate the recombination of V(D)J gene elements in T and B cell antigen receptor genes (required for the generation of diversity in antigen recognition). The main deficiencies involve RAG-1, RAG-2, DNA-dependent protein kinase, and Artemis. A less severe (albeit variable) immunologic phenotype can result from other deficiencies in the same pathway, i.e., DNA ligase 4 and Cernunnos deficiencies. Given that these latter factors are involved in DNA repair, these deficiencies also cause developmental defects.

DEFECTIVE (PRE-)T CELL RECEPTOR SIGNALING IN THE THYMUS A selective T cell defect can be caused by a series of rare deficiencies in molecules involved in signaling via the pre-TCR or the TCR. These include deficiencies in CD3 subunits associated with the (pre-)TCR (i.e., CD3 δ , ϵ , and ζ) and CD45.

RETICULAR DYSGENESIS Reticular dysgenesis is an extremely rare form of SCID that causes T and NK deficiencies with severe neutropenia and sensorineural deafness. It results from an adenylate kinase 2 deficiency.

Patients with SCID require appropriate care with aggressive anti-infective therapies, immunoglobulin replacement, and (when necessary) parenteral nutrition support. In most cases, curative treatment relies on HSCT. Today, HSCT provides a very high curative potential for SCID patients who are otherwise in reasonably good condition. In this regard, neonatal screening, based on quantification of T cell receptor excision circles (TRECs) on a Guthrie card sample, is being developed. Gene therapy has been found to be successful for cases of X-linked SCID (γ c deficiency) and SCID caused by an ADA deficiency, although toxicity has become an issue in the treatment of the former disease that may now be overcome by use of newly generated vectors. Lastly, a third option for the treatment of ADA deficiency consists of enzyme substitution with a pegylated enzyme.

Thymic Defects A profound T cell defect can also result from faulty development of the thymus, as is most often observed in rare cases of DiGeorge syndrome—a relatively common condition leading to a constellation of developmental defects. In approximately 1% of such cases, the thymus is completely absent, leading to virtually no mature T cells. However, expansion of oligoclonal T cells can occur and is associated with skin lesions. Diagnosis (using immunofluorescence in situ hybridization) is based on the identification of a hemizygous deletion in the long arm of chromosome 22. To recover the capability for T cell differentiation, these cases require a thymic graft. CHARGE (coloboma of the eye, heart anomaly, choanal atresia, retardation, genital and ear anomalies) syndrome (CHD7 deficiency) is a less frequent cause of impaired thymus development. Lastly, the very rare “nude” defect is characterized by the absence of both hair and the thymus.

Omenn Syndrome Omenn syndrome consists of a subset of T cell deficiencies that present with a unique phenotype, including early-onset erythrodermia, alopecia, hepatosplenomegaly, and failure to thrive. These patients usually display T cell lymphocytosis, eosinophilia, and low B cell counts. It has been found that the T cells of these patients exhibit a low TCR heterogeneity. This peculiar syndrome is the consequence of hypomorphic mutations in genes usually associated with SCID, i.e., RAG-1, RAG-2, or (less frequently) Artemis or IL-7R α . The impaired homeostasis of differentiating T cells thus causes this immune system-associated disease. These patients are very fragile,

requiring simultaneous anti-infective therapy, nutritional support, and immunosuppression. HSCT provides a curative approach.

Functional T Cell Defects (Fig. 374-2) A subset of T cell PIDs with autosomal inheritance is characterized by partially preserved T cell differentiation but defective activation resulting in abnormal effector function. There are many causes of these defects, but all lead to susceptibility to viral and opportunistic infections, chronic diarrhea, and failure to thrive, with onset during childhood. Careful phenotyping and in vitro functional assays are required to identify these diseases, the best characterized of which are the following.

ZETA-ASSOCIATED PROTEIN 70 (ZAP70) DEFICIENCY Zeta-associated protein 70 (ZAP70) is recruited to the TCR following antigen recognition. A ZAP70 deficiency leads typically to an almost complete absence of CD8⁺ T cells; CD4⁺ T cells are present but cannot be activated in vitro by TCR stimulation.

CALCIUM SIGNALING DEFECTS A small number of patients have been reported who exhibit a profound defect in in vitro T and B cell activation as a result of defective antigen receptor-mediated Ca²⁺ influx. This defect is caused by a mutation in the calcium channel gene (*ORAI-1*) or its activator (*STIM-1*). It is noteworthy that these patients are also prone to autoimmune manifestations (blood cytopenias) and exhibit a nonprogressive muscle disease.

HUMAN LEUKOCYTE ANTIGEN (HLA) CLASS II DEFICIENCY Defective expression of HLA class II molecules is the hallmark of a group of four recessive genetic defects all of which affect molecules (RFX5, RFXAP, RFXANK, and CIITA) involved in the transactivation of the genes coding for HLA class II. As a result, low but variable CD4⁺ T cell counts are observed in addition to defective antigen-specific T and B cell responses. These patients are particularly susceptible to herpesvirus, adenovirus, and enterovirus infections and chronic gut/liver *Cryptosporidium* infections.

HLA CLASS I DEFICIENCY Defective expression of molecules involved in antigen presentation by HLA class I molecules (i.e., TAP-1, TAP-2, and Tapasin) leads to reduced CD8⁺ T cell counts, loss of HLA class I antigen expression, and a particular phenotype consisting of chronic obstructive pulmonary disease and severe vasculitis.

OTHER DEFECTS A variety of other T cell PIDs have been described, some of which are associated with a precise molecular defect (e.g., IL-2-inducible T cell kinase [ITK] deficiency, IL-21 receptor deficiency, CARD11 deficiency). These conditions are also characterized by profound vulnerability to infections, such as severe Epstein-Barr virus (EBV)-induced B cell proliferation and autoimmune disorders in ITK deficiency. Milder phenotypes are associated with CD8 and CD3 γ deficiencies.

HSCT is indicated for most of these diseases, although the prognosis is worse than in SCID because many patients are chronically infected at the time of diagnosis. Fairly aggressive immunosuppression and myeloablation may be necessary to achieve engraftment of allogeneic stem cells.

T Cell Primary Immunodeficiencies with DNA Repair Defects This is a group of PIDs characterized by a combination of T and B cell defects of variable intensity, together with a number of nonimmunologic features resulting from DNA fragility. The autosomal recessive disorder *ataxia-telangiectasia* (AT) is the most frequently encountered condition in this group. It has an incidence of 1:40,000 live births and causes B cell defects (low IgA, IgG2 deficiency, and low antibody production), which often require immunoglobulin replacement. AT is associated with a progressive T cell immunodeficiency. As the name suggests, the hallmark features of AT are telangiectasia and cerebellar ataxia. The latter manifestations may not be detectable before the age of 3–4 years, so that AT should be considered in young children with IgA deficiency and recurrent and problematic infections. Diagnosis is based on a cytogenetic analysis showing excessive chromosomal rearrangements (mostly affecting chromosomes 7 and 14) in lymphocytes. AT is caused by a mutation in the gene encoding the ATM protein—a kinase that plays an important role in the detection and repair of DNA lesions (or cell death if the lesions are too numerous) by triggering

several different pathways. Overall, AT is a progressive disease that carries a very high risk of lymphoma, leukemia, and (during adulthood) carcinomas. A variant of AT (“AT-like disease”) is caused by mutation in the *MRE11* gene.

Nijmegen breakage syndrome (NBS) is a less common condition that also results from chromosome instability (with the same cytogenetic abnormalities as in AT). NBS is characterized by a severe T and B cell combined immune deficiency with autosomal recessive inheritance. Individuals with NBS exhibit microcephaly and a bird-like face, but have neither ataxia nor telangiectasia. The risk of malignancies is very high. NBS results from a deficiency in nibrin (NBS1, a protein associated with MRE11 and Rad50 that is involved in checking DNA lesions) caused by hypomorphic mutations.

Severe forms of *dyskeratosis congenita* (also known as Hoyeraal-Hreidarsson syndrome) combine a progressive immunodeficiency that can also include an absence of B and NK lymphocytes, progressive bone marrow failure, microcephaly, in utero growth retardation, and gastrointestinal disease. The disease can be X-linked or, more rarely, autosomal recessive. It is caused by the mutation of genes encoding telomere maintenance proteins, including dyskerin (DKC1).

Finally, *immunodeficiency with centromeric and facial anomalies* (ICF) is a complex syndrome of autosomal recessive inheritance that variably combines a mild T cell immune deficiency with a more severe B cell immune deficiency, coarse face, digestive disease, and mild mental retardation. A diagnostic feature is the detection by cytogenetic analysis of multiradial aspects in multiple chromosomes (most frequently 1, 9, and 16) corresponding to an abnormal DNA structure secondary to defective DNA methylation. It is the consequence of a deficiency in the DNA methyltransferase DNMT3B, or ZBTB24.

T Cell Primary Immunodeficiencies with Hyper-IgE Several T cell PIDs are associated with elevated serum IgE levels (as in Omenn syndrome). A condition sometimes referred to as *autosomal recessive hyper-IgE syndrome* is notably characterized by recurrent bacterial infections in the skin and respiratory tract and severe skin and mucosal infections by pox viruses and human papillomaviruses, together with severe allergic manifestations. T and B lymphocyte counts are low. Mutations in the *DOCK8* gene have been found in most of these patients. This condition is an indication for HSCT.

A very rare, related condition with autosomal recessive inheritance that causes a similar susceptibility to infection with various microbes (see above), including mycobacteria, reportedly results from a deficiency in Tyk-2, a JAK family kinase involved in the signaling of many different cytokine receptors.

Autosomal Dominant Hyper-IgE Syndrome This unique condition, the *autosomal dominant hyper-IgE syndrome*, is usually diagnosed by the combination of recurrent skin and lung infections that can be complicated by pneumatoceles. Infections are caused by pyogenic bacteria and fungi. Several other manifestations characterize hyper-IgE syndrome, including facial dysmorphism, defective loss of primary teeth, hyperextensibility, scoliosis, and osteoporosis. Elevated serum IgE levels are typical of this syndrome. Defective T_H17 effector responses have been shown to account at least in part for the specific patterns of susceptibility to particular microbes. This condition is caused by a heterozygous (dominant) mutation in the gene encoding the transcription factor STAT3 that is required in a number of signaling pathways following binding of cytokine to cytokine receptors (such as that of IL-6 and the IL-6 receptor). It also results in partially defective antibody production because of defective IL-21R signaling. Hence, immunoglobulin substitution can be considered as prophylaxis of bacterial infections.

Cartilage Hair Hypoplasia The autosomal recessive *cartilage hair hypoplasia* (CHH) disease is characterized by short-limb dwarfism, metaphyseal dysostosis, and sparse hair, together with a combined T and B cell PID of extremely variable intensity (ranging from quasi-SCID to no clinically significant immune defects). The condition can predispose to erythroblastopenia, autoimmunity, and tumors. It is caused by mutations in the *RMRP* gene for a noncoding ribosome-associated RNA.

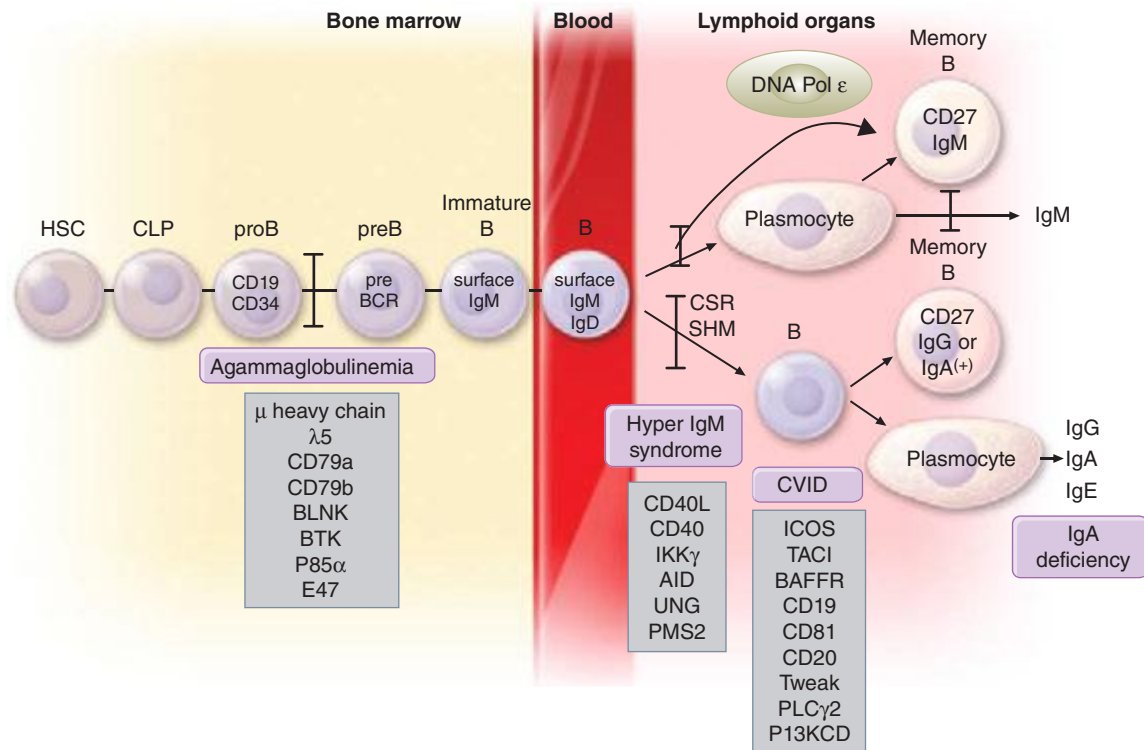


FIGURE 374-4 B cell differentiation and related primary immunodeficiencies (PIDs). Hematopoietic stem cells (HSCs) differentiate into common lymphoid progenitors (CLPs), which give rise to pre-B cells. The B cell differentiation pathway goes through the pre-B cell stage (expression of the μ heavy chain and surrogate light chain), the immature B cell stage (expression of surface IgM), and the mature B cell stage (expression of surface IgM and IgD). The main phenotypic characteristics of these cells are indicated. In lymphoid organs, B cells can differentiate into plasma cells and produce IgM or undergo (in germinal centers) Ig class switch recombination (CSR) and somatic mutation of the variable region of V genes (SHM) that enable selection of high-affinity antibodies. These B cells produce antibodies of various isotypes and generate memory B cells. PIDs are indicated in the purple boxes. CVID, common variable immunodeficiency.

CD40 Ligand and CD40 Deficiencies *Hyper-IgM syndrome* (HIGM) is a well-known PID that is usually classified as a B cell immune deficiency (see Fig. 374-4 and below). It results from defective immunoglobulin class switch recombination (CSR) in germinal centers and leads to profound deficiency in production of IgG, IgA, and IgE (although IgM production is maintained). Approximately half of HIGM sufferers are also prone to opportunistic infections, e.g., interstitial pneumonitis caused by *Pneumocystis jiroveci* (in young children), protracted diarrhea and cholangitis caused by *Cryptosporidium*, and infection of the brain with *Toxoplasma gondii*.

In the majority of cases, this condition has an X-linked inheritance and is caused by a deficiency in CD40 ligand (L). CD40L induces signaling events in B cells that are necessary for both CSR and adequate activation of other CD40-expressing cells that are involved in innate immune responses against the above-mentioned microorganisms. More rarely, the condition is caused by a deficiency in CD40 itself. The poorer prognosis of CD40L and CD40 deficiencies (relative to most other HIGM conditions) implies that (1) thorough investigations have to be performed in all cases of HIGM and (2) potentially curative HSCT should be discussed on a case-by-case basis for this group of patients.

Wiskott-Aldrich Syndrome *Wiskott-Aldrich syndrome* (WAS) is a complex, recessive, X-linked disease with an incidence of approximately 1 in 200,000 live births. It is caused by mutations in the *WASP* gene that affect not only T lymphocytes but also the other lymphocyte subsets, dendritic cells, and platelets. WAS is typically characterized by the following clinical manifestations: recurrent bacterial infections, eczema, and bleeding caused by thrombocytopenia. However, these manifestations are highly variable—mostly as a consequence of the many different *WASP* mutations that have been observed. Null mutations predispose affected individuals to invasive and bronchopulmonary infections, viral infections, severe eczema, and autoimmune manifestations. The latter include autoantibody-mediated blood cytopenia,

glomerulonephritis, skin and visceral vasculitis (including brain vasculitis), erythema nodosum, and arthritis. Another possible consequence of WAS is lymphoma, which may be virally induced (e.g., by EBV or Kaposi's sarcoma-associated herpesvirus). Thrombocytopenia can be severe and compounded by the peripheral destruction of platelets associated with autoimmune disorders. Hypomorphic mutations usually lead to milder outcomes that are generally limited to thrombocytopenia. It is noteworthy that even patients with "isolated" X-linked thrombocytopenia can develop severe autoimmune disease or lymphoma later in life. The immunologic workup is not very informative; there can be a relative CD8+ T cell deficiency, frequently accompanied by low serum IgM levels and decreased antigen-specific antibody responses. A typical feature is reduced-sized platelets on a blood smear. Diagnosis is based on intracellular immunofluorescence analysis of WAS protein (WASp) expression in blood cells. WASp regulates the actin cytoskeleton and thus plays an important role in many lymphocyte functions, including cell adhesion and migration and the formation of synapses between antigen-presenting and target cells. Predisposition to autoimmune disorders is in part related to defective regulatory T cells. The treatment of WAS should match the severity of disease expression. Prophylactic antibiotics, immunoglobulin G (IgG) supplementation, and careful topical treatment of eczema are indicated. Although splenectomy improves platelet count in a majority of cases, this intervention is associated with a significant risk of infection (both before and after HSCT). Allogeneic HSCT is curative, with fairly good results overall. Gene therapy trials are also under way. A similar condition has been reported in a girl with a deficiency in the Wiskott-Aldrich interacting protein (WIP).

A few other complex PIDs are worth mentioning. *Sp110 deficiency* causes a T cell PID with liver venoocclusive disease and hypogammaglobulinemia. *Chronic mucocutaneous candidiasis* (CMC) is a heterogeneous disease, considering the different inheritance patterns that have been observed. In some cases, chronic candidiasis is associated with

late-onset bronchopulmonary infections, bronchiectasis, and brain aneurysms. Moderate forms of CMC are related to autoimmunity and AIRE deficiency (see below). In this setting, predisposition to *Candida* infection is associated with the detection of autoantibodies to T_H17 cytokines. Recently, deficiencies in IL-17F and IL-17 receptor A and, above all, gain-of-function mutations in *STAT1* have been found to be associated with CMC. In all cases, CMC is related to defective T_H17 function. Innate immunodeficiency in *CARD9* also predisposes to chronic invasive fungal infection.

B LYMPHOCYTE DEFICIENCIES (TABLE 374-1, FIG. 374-4)

Deficiencies that predominantly affect B lymphocytes are the most frequent PIDs and account for 60–70% of all cases. B lymphocytes make antibodies. Pentameric IgMs are found in the vascular compartment and are also secreted at mucosal surfaces. IgG antibodies diffuse freely into extravascular spaces, whereas IgA antibodies are produced and secreted predominantly from mucosa-associated lymphoid tissues. Although Ig isotypes have distinct effector functions, including Fc receptor–mediated and (indirectly) C_3 receptor–dependent phagocytosis of microorganisms, they share the ability to recognize and neutralize a given pathogen. Defective antibody production therefore allows the establishment of invasive, pyogenic bacterial infections as well as recurrent sinus and pulmonary infections (mostly caused by *Streptococcus pneumoniae*, *Haemophilus influenzae*, *Moraxella catarrhalis*, and, less frequently, gram-negative bacteria). If left untreated, recurrent bronchial infections lead to bronchiectasis and, ultimately, cor pulmonale and death. Parasitic infections such as caused by *Giardia lamblia* and bacterial infections caused by *Helicobacter* and *Campylobacter* of the gut are also observed. A complete lack of antibody production (namely agammaglobulinemia) can also predispose affected individuals to severe, chronic, disseminated enteroviral infections causing meningoencephalitis, hepatitis, and a dermatomyositis-like disease.

Even with the most profound of B cell deficiencies, infections rarely occur before the age of 6 months; this is because of transient protection provided by the transplacental passage of immunoglobulins during the last trimester of pregnancy. Conversely, a genetically nonimmunodeficient child born to a mother with hypogammaglobulinemia is, in the absence of maternal Ig substitution, usually prone to severe bacterial infections in utero and for several months after birth.

Diagnosis of B cell PIDs relies on the determination of serum Ig levels (Table 316-2). Determination of antibody production following immunization with tetanus toxoid vaccine or nonconjugated pneumococcal polysaccharide antigens can also help diagnose more subtle deficiencies. Another useful test is B cell phenotype determination in switched μ - δ -CD27+ and nonswitched memory B cells (μ + δ +CD27+). In agammaglobulinemic patients, examination of bone marrow B cell precursors (Fig. 374-4) can help obtain a precise diagnosis and guide the choice of genetic tests.

Agammaglobulinemia Agammaglobulinemia is characterized by a profound defect in B cell development (<1% of the normal B cell blood count). In most patients, very low residual Ig isotypes can be detected in the serum. In 85% of cases, agammaglobulinemia is caused by a mutation in the *BTK* gene that is located on the X chromosome. The *BTK* gene product is a kinase that participates in (pre) B cell receptor signaling. When the kinase is defective, there is a block (albeit a leaky one) at the pre-B to B cell stage (Fig. 374-4). Detection of *BTK* by intracellular immunofluorescence of monocytes, and lack thereof in patients with X-linked agammaglobulinemia, is a useful diagnostic test. Not all of the mutations in *BTK* result in agammaglobulinemia, since some patients have a milder form of hypogammaglobulinemia and low but detectable B cell counts. These cases should not be confused with common variable immunodeficiency (CVID, see below). About 10% of agammaglobulinemia cases are caused by alterations in genes encoding elements of the pre-B cell receptor, i.e., the μ heavy chain, the $\lambda 5$ surrogate light chain, Iga or Ig β , the scaffold protein BLNK, and the p85 α subunit of phosphatidylinositol 3 phosphate kinase (P13K). In 5% of cases, the defect is unknown. It is noteworthy that agammaglobulinemia can be observed in patients with ICF

syndrome, despite the presence of normal peripheral B cell counts. Lastly, agammaglobulinemia can be a manifestation of a myelodysplastic syndrome (associated or not with neutropenia). Treatment of agammaglobulinemic patients is based on immunoglobulin replacement (see below). Profound hypogammaglobulinemia is also observed in adults, in association with thymoma.

Hyper-IgM (HIGM) Syndromes HIGM is a rare B cell PID characterized by defective Ig CSR. It results in very low serum levels of IgG and IgA and elevated or normal serum IgM levels. The clinical severity is similar to that seen in agammaglobulinemia, although chronic lung disease and sinusitis are less frequent and enteroviral infections are uncommon. As discussed above, a diagnosis of HIGM involves screening for an X-linked CD40L deficiency and an autosomal recessive CD40 deficiency, which affect both B and T cells. In 50% of cases affecting only B cells, these isolated HIGM syndromes result from mutations in the gene encoding activation-induced deaminase, the protein that induces CSR in B cell germinal centers. These patients usually have enlarged lymphoid organs. In the other 50% of cases, the etiology is unknown (except for rare UNG and PMS2 deficiencies). Furthermore, IgM-mediated autoimmunity and lymphomas can occur in HIGM syndrome. It is noteworthy that HIGM can result from fetal rubella syndrome or can be a predominant immunologic feature of other PIDs, such as the immunodeficiency associated with ectodermic anhydrotic hypoplasia X-linked NEMO deficiency and the combined T and B cell PIDs caused by DNA repair defects such as AT and Cernunnos deficiency.

Common Variable Immunodeficiency (CVID) CVID is an ill-defined condition characterized by low serum levels of one or more Ig isotypes. Its prevalence is estimated to be 1 in 20,000. The condition is recognized predominantly in adults, although clinical manifestations can occur earlier in life. Hypogammaglobulinemia is associated with at least partially defective antibody production in response to vaccine antigens. B lymphocyte counts are often normal but can be low. Besides infections, CVID patients may develop lymphoproliferation (splenomegaly), granulomatous lesions, colitis, antibody-mediated autoimmune disease, and lymphomas. A family history is found in 10% of cases. A clear-cut dominant inheritance pattern is found in some families, whereas recessive inheritance is observed more rarely. In most cases, no molecular cause can be identified. A small number of patients in Germany were found to carry mutations in the *ICOS* gene encoding a T cell-membrane protein that contributes to B cell activation and survival. In 10% of patients with CVID, monoallelic or biallelic mutations of the gene encoding TACI (a member of the tumor necrosis factor [TNF] receptor family that is expressed on B cells) have been found. In fact, heterozygous TACI mutations correspond to a genetic susceptibility factor, since similar heterozygous mutations are found in 1% of controls. The BAFF receptor was found to be defective in a kindred with CVID, although not all individuals carrying the mutation have CVID.

Recently a group of patients with hypogammaglobulinemia and lymphoproliferation were shown to exhibit dominant gain of function mutations in the *PIK3CD* gene encoding the p110 δ form of P13 kinase. A diagnosis of CVID should be made after excluding the presence of hypomorphic mutations associated with agammaglobulinemia or more subtle T cell defects; this is particularly the case in children. It is possible that many cases of CVID result from a constellation of factors, rather than a single genetic defect. Recently, rare cases of hypogammaglobulinemia were found to be associated with CD19 and CD81 deficiencies. These patients have B cells that can be identified by typing for other B cell markers. Hypogammaglobulinemia can be associated with neutropenia and lymphopenia in the warts, hypogammaglobulinemia, infections, and myelokathexis syndrome (WHIM) caused by dominant gain-of-function mutation of *CXCR4*, resulting in cell retention in the bone marrow.

Selective Ig Isotype Deficiencies IgA deficiency and CVID represent polar ends of a clinical spectrum due to the same underlying gene defect(s) in a large subset of these patients. IgA deficiency is the most common PID; it can be found in 1 in every 600 individuals. It is asymptomatic in most cases; however, individuals may present

2112 with increased numbers of acute and chronic respiratory infections that may lead to bronchiectasis. In addition, over their lifetime, these patients experience an increased susceptibility to drug allergies, atopic disorders, and autoimmune diseases. Symptomatic IgA deficiency is probably related to CVID, since it can be found in relatives of patients with CVID. Furthermore, IgA deficiency may progress to CVID. It is thus important to assess serum Ig levels in IgA-deficient patients (especially when infections occur frequently) in order to detect changes that should prompt the initiation of immunoglobulin replacement. Selective IgG2 (+G4) deficiency (which in some cases may be associated with IgA deficiency) can also result in recurrent sinopulmonary infections and should thus be specifically sought in this clinical setting. These conditions are ill-defined and often transient during childhood. A pathophysiologic explanation has not been found.

Selective Antibody Deficiency to Polysaccharide Antigens Some patients with normal serum Ig levels are prone to *S. pneumoniae* and *H. influenzae* infections of the respiratory tract. Defective production of antibodies against polysaccharide antigens (such as those in the *S. pneumoniae* cell wall) can be observed and is probably causative. This condition may correspond to a defect in marginal zone B cells, a B cell subpopulation involved in T-independent antibody responses.

Immunoglobulin Replacement IgG antibodies have a half-life of 21–28 days. Thus, injection of plasma-derived polyclonal IgG containing a myriad of high-affinity antibodies can provide protection against disease-causing microorganisms in patients with defective IgG antibody production. This form of therapy should not be based on laboratory data alone (i.e., IgG and/or antibody deficiency) but should be guided by the occurrence or not of infections; otherwise, patients might be subjected to unjustified IgG infusions. Immunoglobulin replacement can be performed by IV or subcutaneous routes. In the former case, injections have to be repeated every 3–4 weeks, with a residual target level of 800 mg/mL in patients who had very low IgG levels prior to therapy. Subcutaneous injections are typically performed once a week, although the frequency can be adjusted on a case-by-case basis. A trough level of 800 mg/mL is desirable. Whatever the mode of administration, the main goal is to reduce the frequency of the respiratory tract infections and prevent chronic lung and sinus disease. The two routes appear to be equally safe and efficacious, and so the choice should be left to the preference of the patient.

In patients with chronic lung disease, chest physical therapy with good pulmonary toilet and the cyclic use of antibiotics are also needed. Immunoglobulin replacement is well tolerated by most patients, although the selection of the best-tolerated Ig preparation may be necessary in certain cases. Since IgG preparations contain a small proportion of IgAs, caution should be taken in patients with residual antibody production capacity and a complete IgA deficiency, as these subjects may develop anti-IgA antibodies that can trigger anaphylactic shock. These patients should be treated with IgA-free IgG preparations. Immunoglobulin replacement is a lifelong therapy; its rationale and procedures have to be fully understood and mastered by the patient and his or her family in order to guarantee the strict observance required for efficacy.

PRIMARY IMMUNODEFICIENCIES AFFECTING REGULATORY PATHWAYS (TABLE 374-1)

An increasing number of PIDs have been found to cause homeostatic dysregulation of the immune system, either alone or in association with increased vulnerability to infections. Defects of this type affecting the innate immune system and autoinflammatory syndromes will not be covered in this chapter. However, three specific entities (hemophagocytic lymphohistiocytosis, lymphoproliferation, and autoimmunity) will be described below.

HEMOPHAGOCYtic LYMPHOHISTIOCYTOSIS

Hemophagocytic lymphohistiocytosis (HLH) is characterized by an unremitting activation of CD8+ T lymphocytes and macrophages that leads to organ damage (notably in the liver, bone marrow, and central nervous system). This syndrome results from a broad set of inherited diseases, all of which impair T and NK lymphocyte cytotoxicity. The

manifestations of HLH are often induced by a viral infection. EBV is the most frequent trigger. In severe forms of HLH, disease onset may start during the first year of life or even (in rare cases) at birth.

Diagnosis relies on the identification of the characteristic symptoms of HLH (fever, hepatosplenomegaly, edema, neurologic diseases, blood cytopenia, increased liver enzymes, hypofibrinogenemia, high triglyceride levels, elevated markers of T cell activation, and hemophagocytic features in the bone marrow or cerebrospinal fluid). Functional assays of postactivation cytotoxic granule exocytosis (CD107 fluorescence at the cell membrane) can suggest genetically determined HLH. The conditions can be classified into three subsets:

1. Familial HLH with autosomal recessive inheritance, including perforin deficiency (30% of cases) that can be recognized by assessing intracellular perforin expression; Munc13-4 deficiency (30% of cases); syntaxin 11 deficiency (10% of cases); Munc18-2 deficiency (20% of cases); and a few residual cases that lack a known molecular defect.
2. HLH with partial albinism. Three conditions combine HLH and abnormal pigmentation, where hair examination can help in the diagnosis: Chédiak-Higashi syndrome, Griscelli syndrome, and Hermansky Pudlak syndrome type II. Chédiak-Higashi syndrome is also characterized by the presence of giant lysosomes within leukocytes (**Chap. 80**), in addition to a primary neurologic disorder with slow progression of symptoms over time.
3. X-linked proliferative syndrome (XLP) is characterized in most patients by the induction of HLH following EBV infection, while other patients develop progressive hypogammaglobulinemia similar to what is observed in CVID and/or certain lymphomas. XLP is caused by a mutation in the *SH2DIA* gene that encodes the adaptor protein SAP (associated with a SLAM family receptor). Several immunologic abnormalities have been described, including low 2B4-mediated NK cell cytotoxicity, impaired differentiation of NKT cells, defective antigen-induced T cell death, and defective T cell helper activity for B cells. A related disorder (XLP2) has recently been described. It is also X-linked and induces HLH (frequently after EBV infection), although the clinical manifestation may be less pronounced. The condition is associated with a deficiency of the antiapoptotic molecule XIAP. The pathophysiology of XLP2 and its relationship to XLP1 remain unclear.

HLH is a life-threatening complication. The treatment of this condition requires aggressive immunosuppression with either the cytotoxic agent etoposide or anti-T cell antibodies. Once remission has been achieved, HSCT should be performed, since it provides the only curative form of therapy.

AUTOIMMUNE LYMPHOPROLIFERATIVE SYNDROME

Autoimmune lymphoproliferative syndrome (ALPS) is characterized by nonmalignant T and B lymphoproliferation causing splenomegaly and enlarged lymph nodes; 70% of patients also display autoimmune manifestations such as autoimmune cytopenias, Guillain-Barré syndrome, uveitis, and hepatitis (**Chaps. 79 and 372e**). A hallmark of ALPS is the presence of CD4–CD8– TCRαβ+ T cells (2–50%) in the blood of affected individuals. Hypergammaglobulinemia involving IgG and IgA is also frequently observed. The syndrome is caused by a defect in Fas-mediated apoptosis of lymphocytes, which can thus accumulate and mediate autoimmunity. Furthermore, ALPS can lead to malignancies.

Most patients carry a heterozygous mutation in the gene encoding Fas that is characterized by dominant inheritance and variable penetrance, depending on the nature of the mutation. A rare and severe form of the disease with early onset can be observed in patients carrying a biallelic mutation of Fas, which profoundly impairs the protein's expression and/or function. Fas-ligand, caspase 10, caspase 8, and neuroblastoma RAS viral oncogene homologue (NRAS) mutations have also been reported in a few cases of ALPS. Many cases of ALPS have not been precisely delineated at the molecular level. A B cell–predominant ALPS has recently been found associated with a protein kinase Cδ gene mutation. Treatment of ALPS is essentially based on the use of proapoptotic drugs, which need to be carefully administered in order to avoid toxicity.

COLITIS, AUTOIMMUNITY, AND PRIMARY IMMUNODEFICIENCIES

Several PIDs (most of which are T cell–related) can cause severe gut inflammation. The prototypic example is *immunodysregulation polyendocrinopathy enteropathy X-linked syndrome* (IPEX), characterized by a widespread inflammatory enteropathy, food intolerance, skin rashes, autoimmune cytopenias, and diabetes. The syndrome is caused by loss-of-function mutations in the gene encoding the transcription factor FOXP3, which is required for the acquisition of effector function by regulatory T cells. In most cases of IPEX, CD4+CD25+ regulatory T cells are absent from the blood. This condition has a poor prognosis and requires aggressive immunosuppression. The only possible curative approach is allogeneic HSCT. IPEX-like syndromes that lack a FOXP3 mutation have also been described. In some cases, a CD25 deficiency has been found. Defective CD25 expression also impairs regulatory cell expansion/function. This functional T cell deficiency means that CD25-deficient patients are also at increased risk of opportunistic infections. It is noteworthy that abnormalities in regulatory T cells have been described in other PID settings, such as in Omenn syndrome, STAT5b deficiency, STIM1 (Ca flux) deficiency, and WAS; these abnormalities may account (at least in part) for the occurrence of inflammation and autoimmunity. The autoimmune features observed in a small fraction of patients with DiGeorge syndrome may have the same cause. Recently, severe inflammatory gut disease has been described in patients with a deficiency in the IL-10 receptor or IL-10.

A distinct autoimmune entity is observed in *autoimmune polyendocrinopathy candidiasis ectodermal dysplasia* (APECED) syndrome, which is characterized by autosomal recessive inheritance. It consists of multiple autoimmune manifestations that can affect solid organs in general and endocrine glands in particular. Mild, chronic *Candida* infection is often associated with this syndrome. The condition is due to mutations in the autoimmune regulator (*AIRE*) gene and results in impaired thymic expression of self-antigens by medullary epithelial cells and impaired negative selection of self-reactive T cells that leads to autoimmune manifestations.

A combination of hypogammaglobulinemia, autoantibody production, cold-induced urticaria or skin granulomas, or autoinflammation has been reported, and has been termed the *PLCβ2-associated antibody deficiency and immune dysregulation* (PLAID or APLAID).

CONCLUSION

The variety and complexity of the clinical manifestations of the many different PIDs strongly indicate that it is important to raise awareness of these diseases. Indeed, early diagnosis is essential for establishing an appropriate therapeutic regimen. Hence, patients with suspected PIDs must always be referred to experienced clinical centers that are able to

perform appropriate molecular and genetic tests. A precise molecular diagnosis is not only necessary for initiating the most suitable treatment, but is also important for genetic counseling and prenatal diagnosis.

One pitfall that may hamper diagnosis is the high variability that is associated with many PIDs. Variable disease expression can result from the differing consequences of various mutations associated with a given condition, as exemplified by WAS and, to a lesser extent, X-linked agammaglobulinemia (XLA). There can also be effects of modifier genes (as also suspected in XLA) and environmental factors such as EBV infection that can be the main trigger of disease in XLP conditions. Furthermore, it has recently been established that somatic mutations in an affected gene can attenuate the phenotype of a number of T cell PIDs. This has been described for ADA deficiency, X-linked SCID, RAG deficiencies, NF-κB essential modulator (NEMO) deficiency, and, most frequently, WAS. In contrast, somatic mutations can create disease states analogous to PID, as reported for ALPS. Lastly, cytokine-neutralizing autoantibodies can mimic a PID, as shown for IFN-γ.

Many aspects of the pathophysiology of PIDs are still unknown, and the disease-causing gene mutations have not been identified in all cases (as illustrated by CVID and IgA deficiency). However, our medical understanding of PIDs has now reached the stage where scientifically based approaches to the diagnosis and treatment of these diseases can be implemented.

There are an increasing number of conditions in which a primary immunodeficiency (PID) has been described as one facet of a more complex disease setting. It is essential to consider associated diseases when a PID is identified as the primary manifestation and, conversely, not neglect the potentially harmful consequences of a PID that could be masked by other manifestations of a particular syndrome.

Below is a short description of these syndromes in which the PID is classified according to the arm of the immune system that is affected.

1. Primary Immunodeficiencies of the Innate Immune System

- a) Several severe congenital neutropenia (SCN) syndromes can be associated with malformations. The recently described SCN disease caused by glucose-6-phosphatase deficiency (G6PC3) can be associated with heart and urogenital malformations. The related glycogenesis Ib disease combines SCN with hypoglycemia and hepatosplenomegaly. Some *HAX-1* gene mutations lead to neurocognitive impairments as well as SCN. Barth syndrome combines SCN with cardiomyopathy. Lastly, Shwachman syndrome is a known autosomal recessive entity (caused by mutation of the *SBDS* gene) in which the defect in granulopoiesis can extend to the other hematopoietic lineages; short stature, bone metaphyseal dysplasia, and exocrine pancreatic insufficiency are known hallmarks of this condition.
- b) Syndromic asplenia combines the risk of infection with heart defects and situs inversus.
- c) Leukocyte adhesion deficiency (LAD) type II includes growth retardation and impairment of cognitive development.
- d) A few patients with X-linked chronic granulomatous disease present with a contiguous gene deletion syndrome that can include the McLeod phenotype, which is characterized by anemia, acanthocytosis, and a severe risk of immune reaction against donor red cells because the patient's red cells do not express the Kell antigen. The McLeod phenotype also can result in a neurologic disease.
- e) X-linked nuclear factor- κ B (NF- κ B) essential modulator (NEMO) deficiency provokes not only a variable set of deficiencies of both innate and adaptive immunity but also mild osteopetrosis, lymphedema, and, more frequently, anhydrotic ectodermal dysplasia, dysmorphic facies, and abnormal conical teeth. The last finding is often helpful in the diagnosis of that condition.

2. Primary Immunodeficiencies of the Adaptive Immune System

- a) **T cell primary immunodeficiencies.** Reticular dysgenesis, a rare severe combined immunodeficiency (SCID) characterized by T lymphopenia and agranulocytosis, can cause sensorineural deafness. Coronin A deficiency is another SCID variant that can be associated with behavioral disorders because the Coronin A gene is located in a genome area known to have been deleted in some patients with this disorder. The lack of enzymes of purine metabolism (adenosine deaminase and purine nucleoside phosphorylase) provokes not only profound T cell lymphocytopenia but also neurologic impairment, including dysautonomia and abnormalities of cognitive development of variable intensity, in many patients. The neurologic impairment can persist after hematopoietic stem cell transplantation (HSCT). Mild chondrodysplasia is a common finding in adenosine deaminase (ADA) deficiency and, indeed, can help the physician arrive at a final diagnosis.
- b) **Primary thymic defects.** DiGeorge syndrome is a complex embryopathy that is caused by hemizygous interstitial deletion of chromosome 22, leading to multiple developmental defects,

including conotruncal defects, hypoparathyroidism, and dysmorphic syndrome. Although a profound T cell immunodeficiency is rare in DiGeorge syndrome (~1% of cases), failure to recognize this feature is likely to have a fatal outcome. Similarly, some forms of the related CHARGE syndrome (mutation of the *CHD7* gene) also cause a profound T cell immunodeficiency.

- c) **T cell primary immunodeficiencies related to calcium influx defects.** Recently, rare T cell PIDs were found to be caused by defective store-operated entry of calcium ions into T and B lymphocytes after antigen stimulation. These defects (caused by *ORA-1* and *STIM-1* deficiencies) also lead to anhydrotic ectodermal dysplasia, abnormal teeth, and, above all, a nonprogressive muscle disease characterized by excessive fatigue.
- d) **DNA repair defects.** Several genetic defects impair DNA repair pathways. Many lead to combined T and B lymphocyte PIDs in a syndromal setting of varying complexity. The most common is ataxia-telangiectasia (AT), an autosomal recessive disorder with an incidence of 1 in 40,000 live births; AT causes a B cell immunodeficiency (low IgA, IgG2 deficiency, and low antibody production) that often requires immunoglobulin replacement therapy.

AT is associated with a progressive T cell immunodeficiency. As the condition's name suggests, the hallmark features are telangiectasia and cerebellar ataxia.

These manifestations may not be detectable before age 3–4 years, and so AT should be considered in young children with IgA deficiency and problematic infections. Diagnosis is based on a cytogenetic analysis showing excessive chromosomal rearrangements (mostly affecting chromosomes 7 and 14) in lymphocytes. AT is caused by mutation of the gene encoding the ATM protein, a kinase that plays a major role in the detection of DNA lesions and the organization of DNA repair (or cell death if the lesions are too numerous) by triggering several different pathways. Overall, AT is a progressive disease that carries a very high risk of lymphoma, leukemia, and (during adulthood) carcinomas. A variant of AT (AT-like disease) is caused by mutation of the *MRE11* gene.

Nijmegen breakage syndrome (NBS) is a less common condition that also results from chromosome instability (and the same cytogenetic abnormalities as in AT). It is characterized by a severe T and B cell combined immunodeficiency with autosomal recessive inheritance. Subjects with NBS exhibit microcephaly and a birdlike face but neither ataxia nor telangiectasia. The risk of malignancies is also very high. Nijmegen breakage syndrome results from a deficiency in Nibrin (NBS1, a protein associated with MRE11 and Rad50 that is involved in checking DNA lesions) caused by hypomorphic mutations.

Severe forms of dyskeratosis congenita (also known as Hoyeraal-Hreiderdsson syndrome) combine a progressive immunodeficiency that can include an absence of B and natural killer cell (NK) lymphocytes, progressive bone marrow failure, microcephaly, in utero growth retardation, and gut disease. The disease can be X-linked or, more rarely, autosomal recessive. It is caused by the mutation of genes encoding telomere maintenance proteins, including dyskerin (DKC1).

Bloom syndrome (helicase deficiency) combines a typical dysmorphic syndrome with growth retardation, skin lesions, and a mild immunodeficiency that also can be found in some patients with Fanconi's anemia.

Rare forms of combined T and B cell immunodeficiencies with autosomal recessive inheritance are associated in more complex syndromes with microcephaly, failure to grow, and a variable dysmorphic syndrome. These disorders are caused by mutation of the genes that encode DNA ligase 4 and Cernunnos (XLF), both of which are members of the nonhomologous end-joining DNA repair pathway.

The Vici syndrome combines callosal agenesis, cataracts, cardiomyopathy, hyperpigmentation, and a combined immunodeficiency. It is caused by biallelic *EPG5* gene mutation that results in defective autophagy.

Lastly, immunodeficiency, centromere instability, and facial anomalies (ICF) syndrome is a complex autosomal recessive syndrome that variably combines a mild T cell immunodeficiency and a more severe B cell immunodeficiency with a coarse face, intestinal disease, and mild mental retardation. A cytogenetic diagnostic feature is the presence of multiradial chromosomes (most frequently chromosomes 1, 9, and 16) caused by DNA defective methylation. The syndrome is a result of either DNA methyltransferase DNMT3B deficiency or ZBTB24 deficiency.

- e) **Growth hormone insensitivity syndrome (Laron dwarfism) with combined primary immunodeficiency.** Mutations in the *STAT5b* gene, which encodes a transcription factor involved in signaling downstream of the growth hormone receptor and the interleukin 2 (IL-2) receptor, lead to susceptibility to infection because of a partial, functional T cell immunodeficiency associated with autoimmune manifestations. The autoimmune manifestations probably result from defective generation/activation of regulatory T cells.
- f) **Hyper-IgE syndrome (autosomal dominant form).** Hyper-IgE syndrome is a complex disorder that combines skin infections, inflammation, and susceptibility to bacterial and fungal infections of skin and lungs, often with pneumatoceles, with characteristic syndromic signs such as facial dysmorphism, defective loss of primary teeth, hyperextensibility, scoliosis, and osteoporosis. Elevated serum IgE levels are typical of hyper-IgE syndrome. The recently reported defects in TH17 effector responses account, at least in part, for the vulnerability to specific infections. This condition is caused by heterozygous (dominant) mutation of the gene encoding the transcription factor STAT3, which is required in a number of signaling pathways downstream of cytokine/cytokine receptor interactions (notably for IL-6 and IL21).
- g) **Primary immunodeficiencies with bone disease.** The autosomal recessive cartilage hair hypoplasia (CHH) disease is characterized by short-limb dwarfism, metaphyseal dysostosis, and sparse hair, together with a combined T and B cell PID of variable intensity, ranging from quasi-SCID to an absence of clinically significant immunodeficiency. The condition can predispose to erythroblastopenia, autoimmunity, and tumors. It is caused by mutations in the *RMRP* gene for a noncoding ribosome-associated RNA.
- Schimke immunoosseous dysplasia is a rare autosomal recessive condition characterized by severe T and B cell immunodeficiency with spondyloepiphyseal dysplasia, growth retardation, and kidney and vascular diseases. It is the consequence of mutations in the *SMARCA1* gene. The function of the gene product may be related to DNA repair.
- h) **Venoocclusive disease with immunodeficiency (VODI syndrome)** is a rare autosomal recessive condition predominantly found in populations originating from Lebanon. It combines severe hepatic venoocclusive disease with usually mild T cell immunodeficiency and panhypogammaglobulinemia. It is caused by a deficiency in a nuclear protein, Sp110.

3. B Cell Primary Immunodeficiencies Hypogammaglobulinemia can be associated with chromosomal defects such as trisomy 18 and Jacobsen syndrome (hemizygous deletion of part of the long arm of chromosome 11). A rare biallelic deficiency of the mismatch repair protein PMS2 leads to a partial deficiency in Ig class switch recombination in patients at a very high risk of cancer in general and colon carcinomas and lymphomas in particular. Transcobalamin deficiency disturbs vitamin B₁₂ transport and therefore impairs hematopoiesis. Hypogammaglobulinemia is easily corrected by vitamin B₁₂ administration and can be a characteristic of this very rare disorder.

4. Primary Immunodeficiencies Affecting Regulatory Pathways Several inherited disorders that lead to hemophagocytic lymphohistiocytosis (HLH) also have features that are important in terms of both diagnosis and prognosis. Three of these disorders—Griscelli syndrome, Chédiak-Higashi syndrome, and the Hermansky-Pudlak type II syndrome—are characterized by partial albinism and silvery hair appearance that can facilitate diagnosis. Hermansky-Pudlak type II also can be a bleeding disorder if platelet aggregation is defective. Chédiak-Higashi syndrome also is characterized by an early-onset progressive neurologic disorder with impaired cognitive development and motor and sensory deficiencies, culminating in a generalized encephalopathy. The encephalopathy is not prevented or arrested by allogeneic HSCT even when the HLH risk is controlled.

5. Primary Immunodeficiencies Associated with Other Conditions Predisposition to infection, notably severe disseminated opportunistic infections including nontuberculous mycobacterial infections, can be associated with autoantibodies against interferon γ as observed in Asia.

A number of conditions can cause PIDs indirectly. For example, hypercatabolism in patients with Steinert's disease may cause hypogammaglobulinemia. Intestinal lymphangiectasia that includes both immunoglobulin and naive T cell loss and can expose the patient to a significant infectious risk. Urinary IgG loss may result from severe nephritic syndromes.

A number of drugs, including antimalarials, captopril, penicillamine, phenytoin, and sulfasalazine, can induce predominantly IgA hypogammaglobulinemia in (probably predisposed) adults.

One also should also consider (1) diseases that are not thought to be PIDs but include the occurrence of recurrent infections and (2) genetic defects of the immune system that lead to other clinical manifestations. A very good example of the first group is cystic fibrosis (CF). Despite having a functionally normal immune system, patients with CF develop protracted bacterial respiratory tract infections, notably *Pseudomonas aeruginosa* colonization. This bacterium can incapacitate innate immune responses and cause unremitting inflammation that further facilitates infection. An example of the second group is primary alveolar proteinosis, which is caused by a defect in surfactant clearance by alveolar macrophages. The condition results from mutation of the gene encoding the granulocyte-macrophage colony-stimulating factor receptor α .

SECTION 2 DISORDERS OF IMMUNE-MEDIATED INJURY

376 Allergies, Anaphylaxis, and Systemic Mastocytosis

Joshua A. Boyce, K. Frank Austen

The term *atopy* implies a tendency to manifest asthma, rhinitis, urticaria, and atopic dermatitis alone or in combination, in association with the presence of allergen-specific IgE. However, individuals without an atopic background may also develop hypersensitivity reactions, particularly urticaria and anaphylaxis, associated with the presence of IgE. Inasmuch as the mast cell is a key effector cell in allergic rhinitis and asthma, and the dominant effector in urticaria, anaphylaxis, and systemic mastocytosis, its developmental biology, activation pathway,

product profile, and target tissues will be considered in the introduction to these clinical disorders.

The binding of IgE to human mast cells and basophils, a process termed *sensitization*, prepares these cells for subsequent antigen-specific activation. The high-affinity Fc receptor for IgE, designated FcεRI, is composed of one α, one β, and two disulfide-linked γ chains, which together cross the plasma membrane seven times. The α chain is responsible for IgE binding, and the β and γ chains provide for signal transduction that follows the aggregation of the sensitized tetrameric receptors by polymeric antigen. The binding of IgE stabilizes the α chain at the plasma membrane, thus increasing the density of FcεRI receptors at the cell surface while sensitizing the cell for effector responses. This accounts for the correlation between serum IgE levels and the numbers of FcεRI receptors detected on circulating basophils.

Signal transduction is initiated through the action of a Src family-related tyrosine kinase termed Lyn that is constitutively associated with the β chain. Lyn transphosphorylates the canonical immunoreceptor tyrosine-based activation motifs (ITAMs) of the β and γ chains of the receptor, resulting in recruitment of more active Lyn to the β chain and of Syk tyrosine kinase. The phosphorylated tyrosines in the ITAMs function as binding sites for the tandem *src* homology two (SH2) domains within Syk. Syk activates not only phospholipase C γ , which associates with the linker of activated T cells at the plasma membrane, but also phosphatidylinositol 3-kinase to provide phosphatidylinositol-3,4,5-trisphosphate, which allows membrane targeting of the Tec family kinase Btk and its activation by Lyn. In addition, the Src family tyrosine kinase Fyn becomes activated after aggregation of IgE receptors and phosphorylates the adapter protein Gab2 that enhances activation of phosphatidylinositol 3-kinase. Indeed, this additional input is essential for mast cell activation, but it can be partially inhibited by Lyn, indicating that the extent of mast cell activation is in part regulated by the interplay between these Src family kinases. Activated phospholipase C γ cleaves phospholipid membrane substrates to provide inositol-1,4,5-trisphosphate (IP $_3$) and 1,2-diaclyglycerols (1,2-DAGs) so as to mobilize intracellular calcium and activate protein kinase C, respectively. The subsequent opening of calcium-regulated activated channels provides the sustained elevations of intracellular calcium required to recruit the mitogen-activated protein kinases, ERK, JNK, and p38 (serine/threonine kinases), which provide cascades to augment arachidonic acid release and to mediate nuclear translocation of transcription factors for various cytokines. The calcium ion-dependent activation of phospholipases cleaves membrane phospholipids to generate lysophospholipids, which, like 1,2-DAG, may facilitate the fusion of the secretory granule perigranular membrane with the cell membrane, a step that releases the membrane-free granules containing the preformed mediators of mast cell effects.

The secretory granule of the human mast cell has a crystalline structure, unlike mast cells of lower species. IgE-dependent cell activation results in solubilization and swelling of the granule contents within the first minute of receptor perturbation; this reaction is followed by the ordering of intermediate filaments about the swollen granule, movement of the granule toward the cell surface, and fusion of the perigranular membrane with that of other granules and with the plasmalemma to form extracellular channels for mediator release while maintaining cell viability.

In addition to exocytosis, aggregation of Fc ϵ RI initiates two other pathways for generation of bioactive products, namely, lipid mediators and cytokines. The biochemical steps involved in expression of such cytokines as tumor necrosis factor α (TNF- α), interleukin (IL) 1, IL-6, IL-4, IL-5, IL-13, granulocyte-macrophage colony-stimulating factor (GM-CSF), and others, including an array of chemokines, have not been specifically defined for mast cells. Inhibition studies of cytokine production (IL-1 β , TNF- α , and IL-6) in mouse mast cells with cyclosporine or FK506 reveal binding to the ligand-specific immunophilin and attenuation of the calcium ion- and calmodulin-dependent serine/threonine phosphatase, calcineurin.

Lipid mediator generation (Fig. 376-1) involves translocation of calcium ion-dependent cytosolic phospholipase A $_2$ to the outer nuclear membrane, with subsequent release of arachidonic acid for metabolic processing by the distinct prostanoid and leukotriene pathways. The constitutive prostaglandin endoperoxide synthase-1 (PGHS-1/cyclooxygenase-1) and the de novo inducible PGHS-2 (cyclooxygenase-2) convert released arachidonic acid to the sequential intermediates, prostaglandins G $_2$ and H $_2$. The glutathione-dependent hematopoietic prostaglandin D $_2$ (PGD $_2$) synthase then converts PGH $_2$ to PGD $_2$, the predominant mast cell prostanoid. The PGD $_2$ receptor DP $_1$ is expressed by platelets and epithelial cells, whereas DP $_2$ is expressed by T $_H$ 2 lymphocytes, eosinophils, and basophils. Mast cells also generate thromboxane A $_2$ (TXA $_2$), a short lived but powerful mediator that induces bronchoconstriction and platelet activation through the T prostanoid (TP) receptor.

For leukotriene biosynthesis, the released arachidonic acid is metabolized by 5-lipoxygenase (5-LO) in the presence of an integral nuclear membrane protein, 5-LO activating protein (FLAP). The calcium

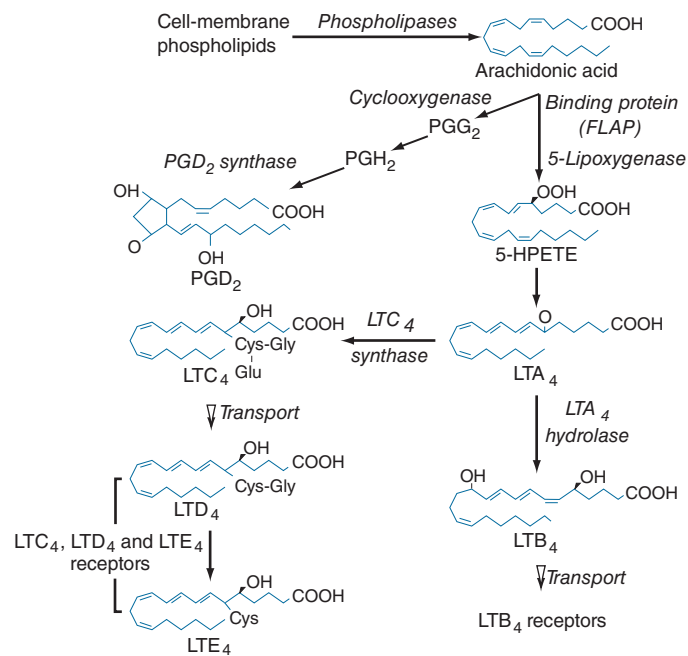


FIGURE 376-1 Pathways for biosynthesis and release of membrane-derived lipid mediators from mast cells. In the 5-lipoxygenase pathway, leukotriene A $_4$ (LTA $_4$) is the intermediate from which the terminal-pathway enzymes generate the distinct final products, leukotriene C $_4$ (LTC $_4$) and leukotriene B $_4$ (LTB $_4$), which leave the cell by separate saturable transport systems. Gamma glutamyl transpeptidase and a dipeptidase then cleave glutamic acid and glycine from LTC $_4$ to form LTD $_4$ and LTE $_4$, respectively. The major mast cell product of the cyclooxygenase system is PGD $_2$.

ion-dependent translocation of 5-LO to the nuclear membrane converts the arachidonic acid to the sequential intermediates, 5-hydroperoxyeicosatetraenoic acid (5-HPETE) and leukotriene (LT) A $_4$. LTA $_4$ is conjugated with reduced glutathione by LTC $_4$ synthase, an integral nuclear membrane protein homologous to FLAP. Intracellular LTC $_4$ is released by a carrier-specific export step for extracellular metabolism to the additional cysteinyl leukotrienes, LTD $_4$ and LTE $_4$, by the sequential removal of glutamic acid and glycine. Alternatively, cytosolic LTA $_4$ hydrolase converts some LTA $_4$ to the dihydroxy leukotriene LTB $_4$, which also undergoes specific export. Two receptors for LTB $_4$, BLT $_1$ and BLT $_2$, mediate chemotaxis of human neutrophils. Two receptors for the cysteinyl leukotrienes, CysLT $_1$ and CysLT $_2$, are present on smooth muscle of the airways and the microvasculature and on hematopoietic cells such as macrophages, eosinophils, and mast cells. Whereas the CysLT $_1$ receptor has a preference for LTD $_4$ and is blocked by the receptor antagonists in clinical use, the CysLT $_2$ receptor is equally responsive to LTD $_4$ and LTC $_4$, is unaffected by these antagonists, and is a negative regulator of the function of the CysLT $_1$ receptor. LTD $_4$, acting at CysLT $_1$ receptors, is the most potent known bronchoconstrictor, whereas LTE $_4$ induces a vascular leak and mediates the recruitment of eosinophils to the bronchial mucosa. Studies in gene-deleted mice indicate the existence of additional receptors for LTE $_4$. The lysophospholipid formed during the release of arachidonic acid from 1-O-alkyl-2-acyl-*sn*-glyceryl-3-phosphorylcholine can be acetylated in the second position to form platelet-activating factor (PAF). Serum levels of PAF correlated positively with the severity of anaphylaxis to peanut in a recent study, whereas the levels of PAF acetyl hydrolase (a PAF-degrading enzyme) were inversely related to the same outcome.

Unlike most other cells of bone marrow origin, mast cells circulate as committed progenitors lacking their characteristic secretory granules. These committed progenitors express *c-kit*, the receptor for stem cell factor (SCF). Unlike most other lineages, they retain and

increase *c-kit* expression with maturation. The SCF interaction with *c-kit* is an absolute requirement for the development of constitutive tissue mast cells residing in skin and connective tissue sites and for the accumulation of mast cells at mucosal surfaces during T_H2 -type immune responses. Several T cell-derived cytokines (IL-3, IL-4, IL-5, and IL-9) can potentiate SCF-dependent mast cell proliferation and/or survival in vitro in mice and humans. Indeed, mast cells are absent from the intestinal mucosa in clinical T cell deficiencies, but are present in the submucosa. Based on the immunodetection of secretory granule neutral proteases, mast cells in the lung parenchyma and intestinal mucosa selectively express tryptase, and those in the intestinal and airway submucosa, perivascular spaces, skin, lymph nodes, and breast parenchyma express tryptase, chymase, and carboxypeptidase A (CPA). In the mucosal epithelium of severe asthmatics, mast cells can express tryptase and CPA without chymase. The secretory granules of mast cells selectively positive for tryptase exhibit closed scrolls with a periodicity suggestive of a crystalline structure by electron microscopy, whereas the secretory granules of mast cells with multiple proteases are scroll-poor, with an amorphous or lattice-like appearance.

Mast cells are distributed at cutaneous and mucosal surfaces and in submucosal tissues about venules and could influence the entry of foreign substances by their rapid response capability (Fig. 376-2). Upon stimulus-specific activation and secretory granule exocytosis, histamine and acid hydrolases are solubilized, whereas the neutral proteases, which are cationic, remain largely bound to the anionic proteoglycans, heparin and chondroitin sulfate E, with which they function as a complex. Histamine and the various lipid mediators (PGD₂, LTC₄/D₄/E₄, PAF) alter venular permeability, thereby allowing influx of plasma proteins such as complement and immunoglobulins, whereas LTB₄ mediates leukocyte-endothelial cell adhesion and subsequent directed migration (chemotaxis). The accumulation of leukocytes and plasma opsonins facilitates defense of the microenvironment. The inflammatory response can also be detrimental, as in asthma, where the smooth-muscle constrictor activity of the cysteinyl leukotrienes is evident and much more potent than that of histamine.

The cellular component of the mast cell-mediated inflammatory response is augmented and sustained by cytokines and chemokines. IgE-dependent activation of human skin mast cells in situ elicits TNF- α production and release, which in turn induces endothelial cell responses favoring leukocyte adhesion. Similarly, activation of purified human lung mast cells or cord blood-derived cultured mast cells in vitro results in substantial production of proinflammatory (TNF- α) and immunomodulatory cytokines (IL-4, IL-5, IL-13) and

chemokines. Bronchial biopsy specimens from patients with asthma reveal that mast cells are immunohistochemically positive for IL-4 and IL-5, but that the predominant localization of IL-4, IL-5, and GM-CSF is to T cells, defined as T_H2 by this profile. IL-4 modulates the T cell phenotype to the T_H2 subtype, determines the isotype switch to IgE (as does IL-13), and upregulates Fc ϵ RI-mediated expression of cytokines by mast cells based on in vitro studies.

An immediate and late cellular phase of allergic inflammation can be induced in the skin, nose, or lung of some allergic humans with local allergen challenge. The immediate phase in the nose involves pruritus and watery discharge; in the lung, it involves bronchospasm and mucus secretion; and in the skin, it involves a wheal-and-flare response with pruritus. The reduced nasal patency, reduced pulmonary function, or erythema with swelling at the skin site in a late-phase response at 6–8 h is associated with biopsy findings of infiltrating and activated T_H2 cells, eosinophils, basophils, and some neutrophils. The progression from early mast cell activation to late cellular infiltration has been used as an experimental surrogate of rhinitis or asthma. However, in asthma, there is an intrinsic hyperreactivity of the airways independent of the associated inflammation. Moreover, early- and late-phase responses (at least in the lung) are far more sensitive to blockade of IgE-dependent mast cell activation (or actions of histamine and cysteinyl leukotrienes) than are spontaneous or virally induced asthma exacerbations.

Consideration of the mechanism of immediate-type hypersensitivity diseases in the human has focused largely on the IgE-dependent recognition of otherwise innocuous substances. A region of chromosome 5 (5q23-31) contains genes implicated in the control of IgE levels including IL-4 and IL-13, as well as IL-3 and IL-9, which are involved in mucosal mast cell hyperplasia, and IL-5 and GM-CSF, which are central to eosinophil development and their enhanced tissue viability. Genes with linkage to the specific IgE response to particular allergens include those encoding the major histocompatibility complex (MHC) and certain chains of the T cell receptor (TCR- $\alpha\delta$). The complexity of atopy and the associated diseases includes susceptibility, severity, and therapeutic responses, each of which is among the separate variables modulated by both innate and adaptive immune stimuli.

The induction of allergic disease requires sensitization of a pre-disposed individual to a specific allergen. The greatest propensity for the development of atopic allergy occurs in childhood and early adolescence. The allergen is processed by antigen-presenting cells of the monocytic lineage (particularly dendritic cells) located throughout the body at surfaces that contact the outside environment, such as the nose, lungs, eyes, skin, and intestine. These antigen-presenting cells present the epitope-bearing peptides via their MHC to T helper cells and their subsets. The T cell response depends both on cognate recognition and on the cytokine microenvironment provided by the antigen-presenting dendritic cells, with IL-4 directing a T_H2 subset, interferon (IFN) γ a T_H1 profile, and IL-6 with transforming growth factor β (TGF- β) a T_H17 subset. Allergens not only present antigenic epitopes via dendritic cells but also contain pattern recognition ligands that facilitate the immune response by direct initiation of cytokine generation from innate cell types such as basophils, mast cells, eosinophils, and others. The T_H2 response is associated with activation of specific B cells that can also present allergens or that transform into plasma cells for antibody production. Synthesis and release into the plasma of allergen-specific IgE results in sensitization of Fc ϵ R1-bearing cells such as mast cells and basophils, which become activated on exposure to the specific allergen. In certain diseases, including those associated with atopy, the monocyte and eosinophil populations can express a trimeric Fc ϵ R1, which lacks the β chain, and yet respond to its aggregation. An additional recently recognized class of *c-kit*-expressing innate cells

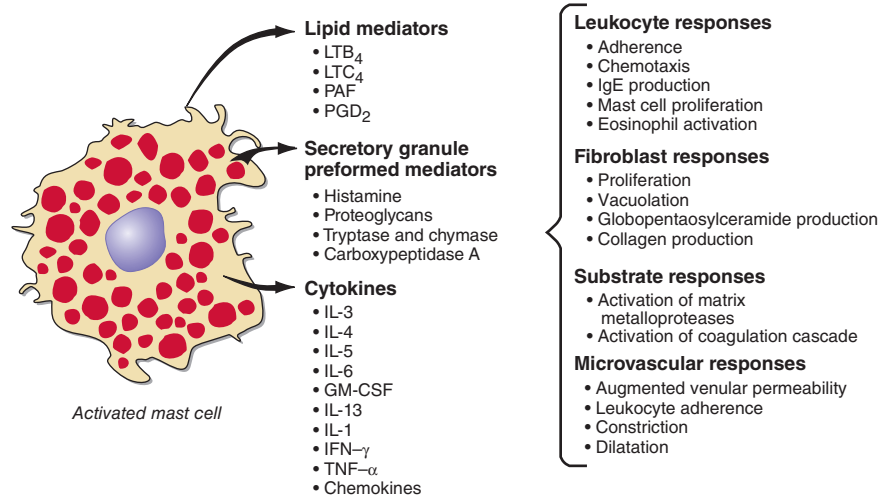


FIGURE 376-2 Bioactive mediators of three categories generated by IgE-dependent activation of murine mast cells can elicit common but sequential target cell effects leading to acute and sustained inflammatory responses. GM-CSF, granulocyte-macrophage colony-stimulating factor; IL, interleukin; IFN, interferon; LT, leukotriene; PAF, platelet-activating factor; PGD₂, prostaglandin D₂; TNF, tumor necrosis factor.

2116 (termed nuocytes, natural helper cells, or group 2 innate lymphoid cells) can generate large quantities of IL-5 and IL-13 during antihelminth responses, are prominent in nasal polyps from humans, and could well contribute to inflammation in allergic diseases.

ANAPHYLAXIS

DEFINITION

Life-threatening anaphylactic responses of sensitized humans occur within minutes after systemic exposure to specific antigen. They are manifested by respiratory distress due to laryngeal edema and/or intense bronchospasm, often followed by vascular collapse, or by shock without antecedent respiratory difficulty. Cutaneous manifestations exemplified by pruritus and urticaria with or without angioedema are characteristic of such systemic anaphylactic reactions. Gastrointestinal manifestations include nausea, vomiting, crampy abdominal pain, and diarrhea.

PREDISPOSING FACTORS AND ETIOLOGY

There is no convincing evidence that age, sex, race, or geographic location predisposes a human to anaphylaxis except through exposure to specific immunogens. According to most studies, atopy does not predispose individuals to anaphylaxis from penicillin therapy or venom of a stinging insect but is a risk factor for allergens in food or latex. Risk factors for a poor outcome, however, include older age, use of beta blockers, and the presence of preexisting asthma. Severe hymenoptera anaphylaxis (generally with prominent hypotension) can be a presenting feature of underlying systemic mastocytosis. Additionally, some individuals suffering from recurrent episodes of idiopathic anaphylaxis possess morphologically aberrant mast cells in their bone marrow that express a mutant, constitutively active form of *c-kit*, even without evidence of frank mastocytosis.

The materials capable of eliciting the systemic anaphylactic reaction in humans include the following: heterologous proteins in the form of hormones (insulin, vasopressin, parathormone); enzymes (trypsin, chymotrypsin, penicillinase, streptokinase); pollen extracts (ragweed, grass, trees); nonpollen allergen extracts (dust mites, dander of cats, dogs, horses, and laboratory animals); food (peanuts, milk, eggs, seafood, nuts, grains, beans, gelatin in capsules); monoclonal antibodies; occupation-related products (latex rubber products); Hymenoptera venom (yellow jacket, yellow and white-faced hornets, paper wasp, honey bee, imported fire ants); polysaccharides such as dextran and thiomersal as a vaccine preservative; drugs such as protamine; antibiotics (penicillins, cephalosporins, amphotericin B, nitrofurantoin, quinolones); chemotherapy agents (carboplatin, paclitaxel, doxorubicin); local anesthetics (procaine, lidocaine); muscle relaxants (suxamethonium, gallamine, pancuronium); vitamins (thiamine, folic acid); diagnostic agents (sodium dehydrocholate, sulfobromophthalein); biologics (omalizumab, rituximab, etanercept); and occupation-related chemicals (ethylene oxide). Drugs function as haptens that form immunogenic conjugates with host proteins. The conjugating hapten may be the parent compound, a nonenzymatically derived storage product, or a metabolite formed in the host. Recombinant biologics can also induce the formation of IgE against the proteins or against glycosylated structures that serve as immunogens. Most recently, outbreaks of anaphylaxis to the anti-epidermal growth factor antibody cetuximab were reported in association with elevated titers of serum IgE to alpha-1,3-galactose, an oligosaccharide found on certain nonprimate proteins. Alpha-galactose antibodies also account for some episodes of delayed anaphylaxis to beef, lamb, and pork.

PATHOPHYSIOLOGY AND MANIFESTATIONS

Individuals differ in the time of appearance of symptoms and signs, but the hallmark of the anaphylactic reaction is the onset of some manifestation within seconds to minutes after introduction of the antigen (with the exception of alpha-galactose allergy), generally by injection or less commonly by ingestion. There may be upper or lower airway obstruction or both. Laryngeal edema may be experienced as a "lump" in the throat, hoarseness, or stridor, whereas bronchial obstruction

is associated with a feeling of tightness in the chest and/or audible wheezing. Patients with asthma are predisposed to severe involvement of the lower airways and increased mortality. Flushing with diffuse erythema and a feeling of warmth may occur. A characteristic feature is the eruption of well-circumscribed, discrete cutaneous wheals with erythematous, raised, serpiginous borders and blanched centers. These urticarial eruptions are intensely pruritic and may be localized or disseminated. They may coalesce to form giant hives, and they seldom persist beyond 48 h. A localized, nonpitting, deeper edematous cutaneous process, angioedema, may also be present. It may be asymptomatic or cause a burning or stinging sensation. Angioedema of the bowel wall may cause sufficient intravascular volume depletion to precipitate cardiovascular collapse.

In fatal cases with clinical bronchial obstruction, the lungs show marked hyperinflation on gross and microscopic examination. The microscopic findings in the bronchi, however, are limited to luminal secretions, peribronchial congestion, submucosal edema, and eosinophilic infiltration, and the acute emphysema is attributed to intractable bronchospasm that subsides with death. The angioedema resulting in death by mechanical obstruction occurs in the epiglottis and larynx, but the process also is evident in the hypopharynx and to some extent in the trachea. On microscopic examination, there is wide separation of the collagen fibers and the glandular elements; vascular congestion and eosinophilic infiltration also are present. Patients dying of vascular collapse without antecedent hypoxia from respiratory insufficiency have visceral congestion with a presumptive loss of intravascular fluid volume. The associated electrocardiographic abnormalities, with or without infarction, in some patients may reflect a primary cardiac event mediated by mast cells (which are prominent near the coronary vessels) or may be secondary to a critical reduction in blood volume.

The angioedematous and urticarial manifestations of anaphylaxis have been attributed to the release of endogenous histamine. A role for the cysteinyl leukotrienes in causing marked bronchiolar constriction seems likely. Vascular collapse without respiratory distress in response to experimental challenge with the sting of a hymenopteran was associated with marked and prolonged elevations in blood histamine and intravascular coagulation and kinin generation. The finding that patients with systemic mastocytosis and episodic vascular collapse excrete large amounts of PGD₂ metabolites in addition to histamine suggests that PGD₂ is also of importance in the hypotensive anaphylactic reactions. As noted, serum PAF levels correlate with severity of anaphylaxis and are inversely proportional to the constitutive level of the acetylhydrolase involved in PAF inactivation. The actions of the array of mast cell-derived mediators are likely additive or synergistic at the target tissues.

DIAGNOSIS

The diagnosis of an anaphylactic reaction depends on a history revealing the onset of symptoms and signs within minutes after the responsible material is encountered. It is appropriate to rule out a complement-mediated immune complex reaction, an idiosyncratic response to a nonsteroidal anti-inflammatory drug (NSAID), or the direct effect of certain drugs or diagnostic agents on mast cells. Intravenous administration of a chemical mast cell-degranulating agent, including opiate derivatives and radiographic contrast media, may elicit generalized urticaria, angioedema, and a sensation of retrosternal oppression with or without clinically detectable bronchoconstriction or hypotension. In the transfusion anaphylactic reaction that occurs in patients with IgA deficiency, the responsible specificity resides in IgG or IgE anti-IgA; the mechanism of the reaction mediated by IgG anti-IgA is presumed to be complement activation with secondary mast cell participation.

The presence of specific IgE in the blood of patients with systemic anaphylaxis was demonstrated historically by passive transfer of the serum intradermally into a normal recipient, followed 24 h later by antigen challenge into the same site, with subsequent development of a wheal and flare (the *Prausnitz-Küstner reaction*). In current clinical practice, immunoassays using purified or recombinant antigens can demonstrate the presence of specific IgE in the serum of patients with

anaphylactic reactions, and skin testing may be performed after the patient has recovered to elicit a local wheal and flare in response to the putative antigen. Elevations of tryptase levels in serum implicate mast cell activation in a systemic reaction and are particularly informative for anaphylaxis with episodes of hypotension during general anesthesia or when there has been a fatal outcome. However, because of the short half-life of tryptase, elevated levels are best detected within 4 h of a systemic reaction. Moreover, anaphylactic reactions to foods characteristically are not associated with elevations in serum tryptase.

TREATMENT ANAPHYLAXIS

Early recognition of an anaphylactic reaction is mandatory, since death can occur within minutes to hours after the first symptoms. Mild symptoms such as pruritus and urticaria can be controlled by administration of 0.3–0.5 mL of 1:1000 (1 mg/mL) epinephrine SC or IM, with repeated doses as required at 5- to 20-min intervals for a severe reaction. The failure to use epinephrine within the first 20 min of symptoms is a risk factor for poor outcome in studies of anaphylaxis to food. If the antigenic material was injected into an extremity, the rate of absorption may be reduced by prompt application of a tourniquet proximal to the reaction site, administration of 0.2 mL of 1:1000 epinephrine into the site, and removal without compression of an insect stinger, if present. An IV infusion should be initiated to provide a route for administration of 2.5 mL epinephrine, diluted 1:10,000, at 5- to 10-min intervals, volume expanders such as normal saline, and vasopressor agents such as dopamine if intractable hypotension occurs. Replacement of intravascular volume due to postcapillary venular leakage may require several liters of saline. Epinephrine provides both α - and β -adrenergic effects, resulting in vasoconstriction, bronchial smooth-muscle relaxation, and attenuation of enhanced venular permeability. Oxygen alone via a nasal catheter or with nebulized albuterol may be helpful, but either endotracheal intubation or a tracheostomy is mandatory for oxygen delivery if progressive hypoxia develops. Ancillary agents such as the antihistamine diphenhydramine, 50–100 mg IM or IV, and aminophylline, 0.25–0.5 g IV, are appropriate for urticaria-angioedema and bronchospasm, respectively. Intravenous glucocorticoids, 0.5–1 mg/kg of methylprednisolone, are not effective for the acute event but may alleviate later recurrence of bronchospasm, hypotension, or urticaria.

PREVENTION

Prevention of anaphylaxis must take into account the sensitivity of the individual, the dose and character of the diagnostic or therapeutic agent, and the effect of the route of administration on the rate of absorption. Beta blockers are relatively contraindicated in persons at risk for anaphylactic reactions, especially those sensitive to Hymenoptera venom or those undergoing immunotherapy for respiratory system allergy. If there is a definite history of a past anaphylactic reaction to a medication, it is advisable to select a structurally unrelated agent. A knowledge of cross-reactivity among agents is critical since, for example, cephalosporins have a cross-reactive ring structure with the penicillins. When skin testing, a prick or scratch skin test should precede an intradermal test, since the latter has a higher risk of causing anaphylaxis. These tests should be performed before the administration of certain materials that are likely to elicit anaphylactic reactions, such as allergenic extracts. Skin testing for antibiotics or chemotherapeutic agents should be performed only on patients with a positive clinical history consistent with an IgE-mediated reaction and in imminent need of the antibiotic in question; skin testing is of no value for non-IgE-mediated eruptions. With regard to penicillin, two-thirds of patients with a positive reaction history and positive skin tests to benzylpenicilloyl-polylysine (BPL) and/or the minor determinant mixture (MDM) of benzylpenicillin products experience allergic reactions with treatment, and these reactions are almost uniformly of the anaphylactic type in those patients with minor determinant reactivity. Even patients without a history of previous clinical reactions have a 2–6% incidence of positive skin tests to the two test materials, and

about 3 per 1000 with a negative history experience anaphylaxis with therapy, with a mortality of about 1 per 100,000.

If an agent carrying a risk of eliciting an anaphylactic response is required because a non-cross-reactive alternative is not available, desensitization can be performed with most antibiotics and other classes of therapeutic agents by the IV, SC, or PO route. Typically, graded quantities of the drug are given by the selected route starting below the threshold dose for an adverse reaction and then doubling each dose until a therapeutic dosage is achieved. Due to the risk of systemic anaphylaxis during the course of desensitization, such a procedure should be performed under the supervision of a specialist and in a setting in which resuscitation equipment is at hand and an IV line is in place. Once a desensitized state is achieved, it is critical to continue administration of the therapeutic agent at regular intervals throughout the treatment period to prevent the reestablishment of a significant pool of sensitized cells.

A different form of protection involves the development of blocking antibody of the IgG class, which protects against Hymenoptera venom-induced anaphylaxis by interacting with antigen so that less reaches the sensitized tissue mast cells. The maximal risk for systemic anaphylactic reactions in persons with Hymenoptera sensitivity occurs in association with a currently positive skin test. Although there is little cross-reactivity between honey bee and yellow jacket venoms, there is a high degree of cross-reactivity between yellow jacket venom and the rest of the vespid venoms (yellow or white-faced hornets and wasps). Prevention involves modification of outdoor activities to exclude bare feet, wearing perfumed toiletries, eating in areas attractive to insects, clipping hedges or grass, and hauling away trash or fallen fruit. As with each anaphylactic sensitivity, the individual should wear an informational bracelet and have immediate access to an unexpired autoinjectable epinephrine kit. Venom immunotherapy for 5 years can induce a state of resistance to sting reactions that is independent of serum levels of specific IgG or IgE. For children under the age of 10 with a systemic reaction limited to skin, the likelihood of progression to more serious respiratory or vascular manifestations is low, and thus immunotherapy is not recommended.

URTICARIA AND ANGIOEDEMA

DEFINITION

Urticaria and angioedema may appear separately or together as cutaneous manifestations of localized nonpitting edema; a similar process may occur at mucosal surfaces of the upper respiratory or gastrointestinal tract. *Urticaria* involves only the superficial portion of the dermis, presenting as well-circumscribed wheals with erythematous raised serpiginous borders and blanched centers that may coalesce to become giant wheals. *Angioedema* is a well-demarcated localized edema involving the deeper layers of the skin, including the subcutaneous tissue, and can also involve the bowel wall. Recurrent episodes of urticaria and/or angioedema of less than 6 weeks' duration are considered acute, whereas attacks persisting beyond this period are designated chronic.

PREDISPOSING FACTORS AND ETIOLOGY

Urticaria and angioedema probably occur more frequently than reported because of the evanescent, self-limited nature of such eruptions, which seldom require medical attention when limited to the skin. Although persons in any age group may experience acute or chronic urticaria and/or angioedema, these lesions increase in frequency after adolescence, with the highest incidence occurring in persons in the third decade of life; indeed, one survey of college students indicated that 15–20% had experienced a pruritic wheal reaction.

The classification of urticaria-angioedema presented in [Table 376-1](#) focuses on the different mechanisms for eliciting clinical disease and can be useful for differential diagnosis; nonetheless, most cases of chronic urticaria are idiopathic. Urticaria and/or angioedema occurring during the appropriate season in patients with seasonal respiratory allergy or as a result of exposure to animals or molds is attributed to inhalation or physical contact with pollens, animal dander, and mold spores, respectively. However, urticaria and angioedema secondary to inhalation are relatively uncommon compared to urticaria and

1. **IgE-dependent**
 - a. Specific antigen sensitivity (pollens, foods, drugs, fungi, molds, Hymenoptera venom, helminths)
 - b. Physical: dermatographism, cold, solar, pressure, cholinergic
 - c. Autoimmune
2. **Bradykinin-mediated**
 - a. Hereditary angioedema: C1 inhibitor deficiency: null (type 1) and dysfunctional (type 2); mutated factor XII (type 3)
 - b. Acquired angioedema: C1 inhibitor deficiency: anti-idiotype and anti-C1 inhibitor
 - c. Angiotensin-converting enzyme inhibitors
3. **Complement-mediated**
 - a. Necrotizing vasculitis
 - b. Serum sickness
 - c. Reactions to blood products
4. **Nonimmunologic**
 - a. Direct mast cell–releasing agents (opiates, antibiotics, curare, D-tubocurarine, radiocontrast media)
 - b. Agents that alter arachidonic acid metabolism (aspirin and nonsteroidal anti-inflammatory agents, azo dyes, and benzoates)
5. **Idiopathic**

angioedema elicited by ingestion of fresh fruits, shellfish, fish, milk products, chocolate, legumes including peanuts, and various drugs that may elicit not only the anaphylactic syndrome with prominent gastrointestinal complaints but also urticaria alone.

Additional etiologies include physical stimuli such as cold, heat, solar rays, exercise, and mechanical irritation. The physical urticarias can be distinguished by the precipitating event and other aspects of the clinical presentation. *Dermatographism*, which occurs in 1–4% of the population, is defined by the appearance of a linear wheal at the site of a brisk stroke with a firm object or by any configuration appropriate to the eliciting event (Fig. 376-3). Dermatographism has a prevalence that peaks in the second to third decades. It is not influenced by atopy and has a duration generally of <5 years. *Pressure urticaria*, which often accompanies chronic idiopathic urticaria, presents in response to a sustained stimulus such as a shoulder strap or belt, running (feet), or manual labor (hands). *Cholinergic urticaria* is distinctive in that the pruritic wheals are of small size (1–2 mm) and are surrounded by a large area of erythema; attacks are precipitated by fever, a hot bath or shower, or exercise and are presumptively attributed to a rise in core

Dermatographism



FIGURE 376-3 Dermatographic urticarial lesion induced by stroking the forearm lightly with the edge of a tongue blade. The photograph, taken after 2 minutes, demonstrates a prominent wheal-and-flare reaction in the shape of an X. (From LA Goldsmith et al [eds]: *Fitzpatrick's Dermatology in General Medicine*, 8th ed. New York, McGraw-Hill, 2012. Photograph provided by Allen P. Kaplan, MD, Medical University of South Carolina.)

body temperature. *Exercise-induced anaphylaxis* can be precipitated by exertion alone or can be dependent on prior food ingestion. There is an association with the presence of IgE specific for α -5 gliadin, a component of wheat. The clinical presentation can be limited to flushing, erythema, and pruritic urticaria but may progress to angioedema of the face, oropharynx, larynx, or intestine or to vascular collapse; it is distinguished from cholinergic urticaria by presenting with wheals of conventional size and by not occurring with fever or a hot bath. *Cold urticaria* is local at body areas exposed to low ambient temperature or cold objects but can progress to vascular collapse with immersion in cold water (swimming). *Solar urticaria* is subdivided into six groups by the response to specific portions of the light spectrum. *Vibratory angioedema* may occur after years of occupational exposure or can be idiopathic; it may be accompanied by cholinergic urticaria. Other rare forms of physical allergy, always defined by stimulus-specific elicitation, include *local heat urticaria*, *aquagenic urticaria* from contact with water of any temperature (sometimes associated with polycythemia vera), and *contact urticaria* from direct interaction with some chemical substance.

Angioedema without urticaria due to the generation of bradykinin occurs with C1 inhibitor (C1INH) deficiency that may be inborn as an autosomal dominant characteristic or may be acquired through the appearance of an autoantibody. The angiotensin-converting enzyme (ACE) inhibitors can provoke a similar clinical presentation in 0.1–0.5% of hypertensive patients due to attenuated degradation of bradykinin. The urticaria and angioedema associated with classic serum sickness or with hypocomplementemic cutaneous necrotizing angitis are believed to be immune-complex diseases. The drug reactions to mast cell granule–releasing agents and to NSAIDs may be systemic, resembling anaphylaxis, or limited to cutaneous sites.

PATHOPHYSIOLOGY AND MANIFESTATIONS

Urticarial eruptions are distinctly pruritic, may involve any area of the body from the scalp to the soles of the feet, and appear in crops of 12- to 36-h duration, with old lesions fading as new ones appear. Most of the physical urticarias (cold, cholinergic, dermatographism) are an exception, with individual lesions lasting less than 2 h. The most common sites for urticaria are the extremities and face, with angioedema often being periorbital and in the lips. Although self-limited in duration, angioedema of the upper respiratory tract may be life-threatening due to laryngeal obstruction, whereas gastrointestinal involvement may present with abdominal colic, with or without nausea and vomiting, and may result in unnecessary surgical intervention. No residual discoloration occurs with either urticaria or angioedema unless there is an underlying vasculitic process leading to superimposed extravasation of erythrocytes.

The pathology is characterized by edema of the superficial dermis in urticaria and of the subcutaneous tissue and deep dermis in angioedema. Collagen bundles in affected areas are widely separated, and the venules are sometimes dilated. Any perivenular infiltrate consists of lymphocytes, monocytes, eosinophils, and neutrophils that are present in varying combination and numbers.

Perhaps the best-studied example of IgE- and mast cell–mediated urticaria and angioedema is *cold urticaria*. Cryoglobulins or cold agglutinins are present in up to 5% of these patients. Immersion of an extremity in an ice bath precipitates angioedema of the distal portion with urticaria at the air interface within minutes of the challenge. Histologic studies reveal marked mast cell degranulation with associated edema of the dermis and subcutaneous tissues. The histamine level in the plasma of venous effluent of the cold-challenged and angioedematous extremity is markedly increased, but no such increase appears in the plasma of effluent of the contralateral normal extremity. Elevated levels of histamine have been found in the plasma of venous effluent and in the fluid of suction blisters at experimentally induced lesional sites in patients with dermatographism, pressure urticaria, vibratory angioedema, light urticaria, and heat urticaria. By ultrastructural analysis, the pattern of mast cell degranulation in cold urticaria resembles an IgE-mediated response with solubilization of granule contents, fusion of the perigranular and cell membranes, and discharge of granule contents, whereas in a dermatographic lesion, there is additional superimposed zonal (piecemeal) degranulation. There are several reports of

resolution of cold urticaria by treatment with monoclonal anti-human IgE (omalizumab). Elevations of plasma histamine levels with biopsy-proven mast cell degranulation have also been demonstrated with generalized attacks of *cholinergic urticaria* and *exercise-related anaphylaxis* precipitated experimentally in subjects exercising on a treadmill while wearing a wet suit; however, only subjects with cholinergic urticaria have a concomitant decrease in pulmonary function.

Up to 40% of patients with chronic urticaria have an autoimmune cause for their disease including autoantibodies to IgE (5–10%) or, more commonly, to the α chain of Fc ϵ RI (35–45%). In these patients, autologous serum injected into their own skin can induce a wheal-and-flare reaction involving mast cell activation. The presence of these antibodies can also be recognized by their capacity to release histamine or induce activation markers such as CD63 or CD203 on basophils. An association with antibodies to microsomal peroxidase and/or thyroglobulin has been observed often with clinically significant Hashimoto's thyroiditis. In vitro studies reveal that these autoantibodies can mediate basophil degranulation with enhancement by serum as a source of the anaphylatoxic fragment, C5a.

Hereditary angioedema is an autosomal dominant disease due to a deficiency of C1INH (type 1) in about 85% of patients and to a dysfunctional protein (type 2) in the remainder. A third type of hereditary angioedema has been described in which C1INH function is normal, and the causal lesion is a mutant form of factor XII, which leads to generation of excessive bradykinin. In the acquired form of C1INH deficiency, there is excessive consumption due either to immune complexes formed between anti-idiotypic antibody and monoclonal IgG presented by B cell lymphomas or to an autoantibody directed to C1INH. C1INH blocks the catalytic function of activated factor XII (Hageman factor) and of kallikrein, as well as the C1r/C1s components of C1. During clinical attacks of angioedema, C1INH-deficient patients have elevated plasma levels of bradykinin, particularly in the venous effluent of an involved extremity, and reduced levels of prekallikrein and high-molecular-weight kininogen, from which bradykinin is cleaved. The parallel decline in the complement substrates C4 and C2 reflects the action of activated C1 during such attacks. Mice with targeted disruption of the gene for C1INH exhibit a chronic increase in vascular permeability. The pathobiology is aggravated by administration of an ACE inhibitor (captopril) and is attenuated by breeding the C1INH null strain to a bradykinin 2 receptor (Bk2R) null strain. As ACE is also described as kininase II, the use of blockers results in impaired bradykinin degradation and explains the angioedema that occurs idiosyncratically in hypertensive patients with a normal C1INH. Bradykinin-mediated angioedema, whether caused by ACE inhibitors or by C1INH deficiency, is noteworthy for the conspicuous absence of concomitant urticaria.

DIAGNOSIS

The rapid onset and self-limited nature of urticarial and angioedematous eruptions are distinguishing features. Additional characteristics are the occurrence of the urticarial crops in various stages of evolution and the asymmetric distribution of the angioedema. Urticaria and/or angioedema involving IgE-dependent mechanisms are often appreciated by historic considerations implicating specific allergens or physical stimuli, by seasonal incidence, and by exposure to certain environments. Direct reproduction of the lesion with physical stimuli is particularly valuable because it so often establishes the cause of the lesion. The diagnosis of an environmental allergen based on the clinical history can be confirmed by skin testing or assay for allergen-specific IgE in serum. IgE-mediated urticaria and/or angioedema may or may not be associated with an elevation of total IgE or with peripheral eosinophilia. Fever, leukocytosis, and an elevated sedimentation rate are absent.

The classification of urticarial and angioedematous states presented in Table 376-1 in terms of possible mechanisms necessarily includes some differential diagnostic points. Hypocomplementemia is not observed in IgE-mediated mast cell disease and may reflect either an acquired abnormality generally attributed to the formation of immune complexes or a genetic or acquired deficiency of C1INH. Chronic recurrent urticaria, generally in females, associated with arthralgias, an elevated sedimentation rate, and normo- or hypocomplementemia

suggests an underlying cutaneous necrotizing angitis. Vasculitic urticaria typically persists longer than 72 h, whereas conventional urticaria often has a duration of 12–36 h. Confirmation depends on a biopsy that reveals cellular infiltration, nuclear debris, and fibrinoid necrosis of the venules. The same pathobiologic process accounts for the urticaria in association with such diseases as systemic lupus erythematosus or viral hepatitis with or without associated arteritis. Serum sickness per se or a similar clinical entity due to drugs includes not only urticaria but also pyrexia, lymphadenopathy, myalgia, and arthralgia or arthritis. Urticarial reactions to blood products or intravenous administration of immunoglobulin are defined by the event and generally are not progressive unless the recipient is IgA-deficient in the former case or the reagent is aggregated in the latter.

The diagnosis of hereditary angioedema is suggested not only by family history but also by the lack of pruritus and of urticarial lesions, the prominence of recurrent gastrointestinal attacks of colic, and episodes of laryngeal edema. Laboratory diagnosis depends on demonstrating a deficiency of C1INH antigen (type 1) or a nonfunctional protein (type 2) by a catalytic inhibition assay. While levels of C1 are normal, its substrates, C4 and C2, are chronically depleted and fall further during attacks due to the activation of additional C1. Patients with the acquired forms of C1INH deficiency have the same clinical manifestations but differ in the lack of a familial element. Furthermore, their sera exhibit a reduction of C1 function and C1q protein as well as C1INH, C4, and C2. Inborn C1INH deficiency and ACE inhibitor-elicited angioedema are associated with elevated levels of bradykinin. Lastly, type 3 hereditary angioedema is associated with normal levels of complement proteins.

Urticaria and angioedema are distinct from contact sensitivity, a vesicular eruption that progresses to chronic thickening of the skin with continued allergenic exposure. They also differ from atopic dermatitis, a condition that may present as erythema, edema, papules, vesiculation, and oozing proceeding to a subacute and chronic stage in which vesiculation is less marked or absent and scaling, fissuring, and lichenification predominate in a distribution that characteristically involves the flexor surfaces. In cutaneous mastocytosis, the reddish brown macules and papules, characteristic of urticaria pigmentosa, urticate with pruritus upon trauma; and in systemic mastocytosis, without or with urticaria pigmentosa, there is episodic systemic flushing with or without urtication but no angioedema.

TREATMENT URTICARIA AND ANGIOEDEMA

Identification and subsequent elimination of the etiologic factor(s) provide the most satisfactory therapeutic program; this approach is feasible to varying degrees with IgE-mediated reactions to allergens or physical stimuli. For most forms of urticaria, H₁ antihistamines such as chlorpheniramine or diphenhydramine effectively attenuate both urtication and pruritus, but because of their side effects, non-sedating agents such as loratadine, desloratadine, and fexofenadine, or low-sedating agents such as cetirizine or levocetirizine generally are used first. Cyproheptadine in dosages beginning at 8 mg and ranging up to 32 mg daily and especially hydroxyzine in dosages beginning at 40 mg and ranging up to 200 mg daily have proven effective when H₁ antihistamines fail. The addition of an H₂ antagonist such as cimetidine, ranitidine, or famotidine in conventional dosages may add benefit when H₁ antihistamines are inadequate. Doxepin, a dibenzoxepin tricyclic compound with both H₁ and H₂ receptor antagonist activity, is yet another alternative. A CysLT₁ receptor antagonist such as montelukast, 10 mg/d, or zafirlukast, 20 mg twice a day, can be important add-on therapy.

Topical glucocorticoids are of no value, and systemic glucocorticoids are generally avoided in idiopathic, allergen-induced, or physical urticarias due to their long-term toxicity. Systemic glucocorticoids are useful in the management of patients with pressure urticaria, vasculitic urticaria (especially with eosinophil prominence), idiopathic angioedema with or without urticaria, or chronic urticaria that responds poorly to conventional treatment. With persistent vasculitic urticaria, hydroxychloroquine, dapsone, or colchicine may be added to the regimen after hydroxyzine and before or

2120 along with systemic glucocorticoids. Cyclosporine can be efficacious for patients with chronic idiopathic or chronic autoimmune urticaria that is severe and poorly responsive to other modalities and/or where a glucocorticoid requirement is excessive. For chronic urticaria induced by autoantibody activation of mast cells and basophils or cold urticaria, monoclonal anti-IgE antibodies such as omalizumab may be considered.

The therapy of inborn C1INH deficiency has been simplified by the finding that attenuated androgens correct the biochemical defect and afford prophylactic protection; their efficacy is attributed to production by the normal gene of an amount of functional C1INH sufficient to control the spontaneous activation of C1. The antifibrinolytic agent ε-aminocaproic acid may be used for preoperative prophylaxis but is contraindicated in patients with thrombotic tendencies or ischemia due to arterial atherosclerosis. Infusion of isolated C1INH protein may be used for prophylaxis or treatment of an acute attack; a bradykinin 2 receptor antagonist and ecallantide, a kallikrein inhibitor, which are administered SC, are each being assessed for amelioration of attacks. Treatment of the underlying hematologic malignancy is indicated for acquired C1INH deficiency.

SYSTEMIC MASTOCYTOSIS

DEFINITION

Systemic mastocytosis is defined by a clonal expansion of mast cells that in most instances is indolent and nonmalignant. The mast cell expansion is generally recognized only in bone marrow and in the normal peripheral distribution sites of the cells, such as skin, gastrointestinal mucosa, liver, and spleen. Mastocytosis occurs at any age and has a slight preponderance in males. The prevalence of systemic mastocytosis is not known, a familial occurrence is rare, and atopy is not increased.

CLASSIFICATION AND PATHOPHYSIOLOGY

A consensus classification for mastocytosis recognizes cutaneous mastocytosis with variants and four systemic forms (Table 376-2). Cutaneous mastocytosis is the most common diagnosis in children, whereas the form designated as *indolent systemic mastocytosis* (ISM) accounts for the majority of adult patients; it implies that there is no evidence of an associated hematologic disorder, liver disease, or lymphadenopathy and is not known to alter life expectancy. In *systemic mastocytosis associated with clonal hematologic non-mast cell lineage disease* (SM-AHNMD), the prognosis is determined by the nature of the associated disorder, which can range from dysmyelopoiesis to leukemia. In *aggressive systemic mastocytosis* (ASM), mast cell infiltration/proliferation in multiple organs such as liver, spleen, gut, and/or bone results in a poor prognosis; a subset of patients with this form has prominent eosinophilia with hepatosplenomegaly and lymphadenopathy. *Mast cell leukemia* (MCL) is the rarest form of the disease and is invariably fatal at present; the peripheral blood contains circulating, metachromatically staining,

atypical mast cells. An aleukemic form of MCL is recognized without circulating mast cells when the percentage of high-grade immature mast cells in bone marrow smears exceeds 20% in a nonspicular area. Mast cell sarcoma and extracutaneous mastocytomas are rare solid mast cell tumors with malignant and benign features, respectively.

A point mutation of A to T at codon 816 of *c-kit* that causes an aspartic acid to valine substitution is found in multiple cell lineages in patients with mastocytosis, resulting in a somatic gain-in-function mutation. This substitution, as well as other rare mutations of *c-kit*, is characteristic of patients with all forms of systemic mastocytosis but is also present in some children with cutaneous mastocytosis, as might be anticipated because mast cells are of bone marrow lineage. The prognosis for patients with cutaneous mastocytosis and for almost all with ISM is a normal life expectancy, whereas that for patients with SM-AHNMD is determined by a non-mast cell component. ASM and MCL carry a poorer prognosis. In infants and children with cutaneous manifestations, namely, urticaria pigmentosa or bullous lesions, visceral involvement is usually lacking, and resolution is common.

CLINICAL MANIFESTATIONS

The clinical manifestations of systemic mastocytosis, distinct from a leukemic complication, are due to tissue occupancy by the mast cell mass, the tissue response to that mass, and the release of bioactive substances acting at both local and distal sites. The pharmacologically induced manifestations are pruritus, flushing, palpitations and vascular collapse, gastric distress, lower abdominal crampy pain, and recurrent headache. The increase in local cell burden is evidenced by the lesions of urticaria pigmentosa at skin sites and may be a direct local cause of bone pain and/or malabsorption. Mast cell-mediated fibrotic changes occur in liver, spleen, and bone marrow but not in gastrointestinal tissue or skin. Immunofluorescent analysis of bone marrow and skin lesions in ISM and of spleen, lymph node, and skin in ASM has revealed only one mast cell phenotype, namely, scroll-poor cells expressing tryptase, chymase, and CPA.

The cutaneous lesions of urticaria pigmentosa are reddish-brown macules or papules that respond to trauma with urtication and erythema (Darier's sign). The apparent incidence of these lesions is ≥80% in patients with ISM and <50% in those with SM-AHNMD or ASM. Approximately 1% of patients with ISM have skin lesions that appear as tan-brown macules with striking patchy erythema and associated telangiectasia (telangiectasia macularis eruptiva perstans). In the upper gastrointestinal tract, gastritis and peptic ulcer are significant problems. In the lower intestinal tract, the occurrence of diarrhea and abdominal pain is attributed to increased motility due to mast cell mediators; this problem can be aggravated by malabsorption, which can also cause secondary nutritional insufficiency and osteomalacia. The periportal fibrosis associated with mast cell infiltration and a prominence of eosinophils may lead to portal hypertension and ascites. In some patients, flushing and recurrent vascular collapse are markedly aggravated by an idiosyncratic response to a minimal dosage of NSAIDs. The neuropsychiatric disturbances are clinically most evident as impaired recent memory, decreased attention span, and "migraine-like" headaches. Patients may experience exacerbation of a specific clinical sign or symptom with alcohol ingestion, temperature changes, stress, use of mast cell-interactive narcotics, or ingestion of NSAIDs.

DIAGNOSIS

Although the diagnosis of mastocytosis is generally suspected on the basis of the clinical history and physical findings, and can be supported by laboratory procedures, it can be established only by a tissue diagnosis. By convention, the diagnosis of systemic mastocytosis depends heavily on bone marrow biopsy to meet the criteria of one major plus one minor or three minor findings (Table 376-3). The bone marrow provides the major criterion by revealing aggregates of mast cells, often in paratrabecular and perivascular locations with lymphocytes and eosinophils, as well as the minor criteria of an abnormal mast cell morphology, an aberrant mast cell membrane immunophenotype, or a codon 816 mutation in any cell type. A serum total tryptase level

TABLE 376-2 CLASSIFICATION OF MASTOCYTOSIS

Cutaneous mastocytosis (CM)
Urticaria pigmentosa (UP)/maculopapular cutaneous mastocytosis (MPCM)
Variants: plaque form, nodular form; telangiectasia macularis eruptiva perstans (TMEP)
Solitary mastocytoma of skin
Diffuse cutaneous mastocytosis
Indolent systemic mastocytosis (ISM)
Systemic mastocytosis with an associated clonal hematologic non-mast cell lineage disease (SM-AHNMD)
Aggressive systemic mastocytosis (ASM)
Mast cell leukemia (MCL)
Mast cell sarcoma (MCS)
Extracutaneous mastocytoma

Source: Modified from SH Swerdlow et al (eds): *World Health Organization Classification of Tumors: Pathology and Genetics in Tumors of Hematopoietic and Lymphoid Tissues*. Lyon, IARC Press, 2008.

TABLE 376-3 DIAGNOSTIC CRITERIA FOR SYSTEMIC MASTOCYTOSIS^a

Major: Multifocal dense infiltrates of mast cells in bone marrow or other extracutaneous tissues with confirmation by immunodetection of tryptase or metachromasia

Minor: Abnormal mast cell morphology with a spindle shape and/or multilobed or eccentric nucleus

Aberrant mast cell surface phenotype with expression of CD25 (IL-2 receptor) and CD2 in addition to C117 (*c-kit*)

Detection of codon 816 mutation in peripheral blood cells, bone marrow cells, or lesional tissue

Total serum tryptase >20 ng/mL

^aDiagnosis requires either the major criterion and one minor criterion or three minor criteria.

and/or a 24-h urine collection for measurement of histamine, histamine metabolites, or metabolites of PGD₂ are noninvasive approaches to consider before bone marrow biopsy. The pro-β and α forms of tryptase are elevated in more than one-half of patients with systemic mastocytosis and provide a minor criterion; the fully processed (“mature”) β form is increased in patients undergoing an anaphylactic reaction. Additional studies directed by the presentation include a bone densitometry, bone scan, or skeletal survey; contrast studies of the upper gastrointestinal tract with small-bowel follow-through, computed tomography scan, or endoscopy; and a neuropsychiatric evaluation. Osteoporosis is increased in mastocytosis and may lead to pathologic fractures.

The differential diagnosis requires the exclusion of other flushing disorders. The 24-h urine assessment of 5-hydroxy-indoleacetic acid and metanephrines should exclude a carcinoid tumor or a pheochromocytoma. Some patients presenting with recurrent mast cell activation symptoms without an obvious increase in mast cell burden in skin or bone marrow have been shown to carry aberrant mast cells with clonality markers of D816C *c-kit* mutation or surface CD25 expression. Most patients with recurrent anaphylaxis, including the idiopathic group, present with angioedema and/or wheezing, which are not manifestations of systemic mastocytosis.

TREATMENT SYSTEMIC MASTOCYTOSIS

The management of systemic mastocytosis uses a stepwise and symptom/sign-directed approach that includes an H₁ antihistamine for flushing and pruritus, an H₂ antihistamine or proton pump inhibitor for gastric acid hypersecretion, oral cromolyn sodium for diarrhea and abdominal pain, and aspirin for severe flushing with or without associated vascular collapse, despite use of H₁ and H₂ antihistamines, to block biosynthesis of PGD₂. Systemic glucocorticoids appear to alleviate the malabsorption. Mast cell cyto-reductive therapy consisting of IFN-α or cladribine is generally reserved for advanced, nonindolent variants of systemic mastocytosis. Their efficacy in ASM is variable, perhaps because of dosage limitations due to side effects. Chemotherapy is appropriate for the frank leukemias. A self-injectable epinephrine prescription is recommended for most patients due to increased incidence of anaphylaxis. Although *c-kit* is a receptor tyrosine kinase, the gain-in-function mutation of codon 816 is not susceptible to inhibition by imatinib mesylate.

ALLERGIC RHINITIS

DEFINITION

Allergic rhinitis is characterized by sneezing; rhinorrhea; obstruction of the nasal passages; conjunctival, nasal, and pharyngeal itching; and lacrimation, all occurring in a temporal relationship to allergen exposure. Although commonly seasonal due to elicitation by airborne pollens, it can be perennial in an environment of chronic exposure to house dust mites, animal danders, or insect products. In North America, the incidence of allergic rhinitis is about 7%. The overall prevalence in North America is nearly 20%, with the peak prevalence of nearly 40% occurring in childhood and adolescence.

PREDISPOSING FACTORS AND ETIOLOGY

Allergic rhinitis generally occurs in atopic individuals, often in association with atopic dermatitis, food allergy, urticaria, and/or asthma (Chap. 309). Up to 40% of patients with rhinitis manifest asthma, whereas ~70% of individuals with asthma experience rhinitis. Symptoms generally appear before the fourth decade of life and tend to diminish gradually with aging, although complete spontaneous remissions are uncommon. A relatively small number of weeds that depend on wind rather than insects for pollination, as well as grasses and some trees, produce sufficient quantities of pollen suitable for wide distribution by air currents to elicit seasonal allergic rhinitis. The dates of pollination of these species generally vary little from year to year in a particular locale but may be quite different in another climate. In the temperate areas of North America, trees typically pollinate from March through May, grasses in June and early July, and ragweed from mid-August to early October. Molds, which are widespread in nature because they occur in soil or decaying organic matter, propagate spores in a pattern that depends on climatic conditions. Perennial allergic rhinitis occurs in response to allergens that are present throughout the year, including animal dander, cockroach-derived proteins, mold spores, or dust mites such as *Dermatophagoides farinae* and *Dermatophagoides pteronyssinus*. Dust mites are scavengers of human skin and excrete cysteine protease allergens in their feces. In up to one-half of patients with perennial rhinitis, no clear-cut allergen can be demonstrated as causative. The ability of many allergens to cause rhinitis rather than lower respiratory tract symptoms (particularly pollens) may be attributed to their large size, 10–100 μm, and retention within the nose.

PATHOPHYSIOLOGY AND MANIFESTATIONS

Episodic rhinorrhea, sneezing, obstruction of the nasal passages with lacrimation, and pruritus of the conjunctiva, nasal mucosa, and oropharynx are the hallmarks of allergic rhinitis. The nasal mucosa is pale and boggy, the conjunctiva congested and edematous, and the pharynx generally unremarkable. Swelling of the turbinates and mucous membranes with obstruction of the sinus ostia and eustachian tubes precipitates secondary infections of the sinuses and middle ear, respectively. Nasal polyps, representing mucosal protrusions containing edema fluid with variable numbers of eosinophils and degranulated mast cells, can increase obstructive symptoms and can concurrently arise within the nasopharynx or sinuses. However, atopy is not a risk factor for nasal polyps, which instead may occur in the setting of the aspirin-intolerant triad of rhinosinusitis and asthma and in patients with chronic staphylococcal colonization, which produces superantigens leading to an intense T_H2 inflammatory response.

The nose presents a large mucosal surface area through the folds of the turbinates and serves to adjust the temperature and moisture content of inhaled air and to filter out particulate materials >10 μm in size by impingement in a mucous blanket; ciliary action moves the entrapped particles toward the pharynx. Entrapment of pollen and digestion of the outer coat by mucosal enzymes such as lysozymes release protein allergens generally of 10,000–40,000 molecular weight. The initial interaction occurs between the allergen and intraepithelial mast cells and then proceeds to involve deeper perivascular mast cells, both of which are sensitized with specific IgE. During the symptomatic season when the mucosae are already swollen and hyperemic, there is enhanced adverse reactivity to the seasonal pollen. Biopsy specimens of nasal mucosa during seasonal rhinitis show submucosal edema with infiltration by eosinophils, along with some basophils and neutrophils.

The mucosal surface fluid contains IgA that is present because of its secretory piece and also IgE, which apparently arrives by diffusion from plasma cells in proximity to mucosal surfaces. IgE fixes to mucosal and submucosal mast cells, and the intensity of the clinical response to inhaled allergens is quantitatively related to the naturally occurring pollen dose. In sensitive individuals, the introduction of allergen into the nose is associated with sneezing, “stiffness,” and discharge, and the fluid contains histamine, PGD₂, and leukotrienes. Thus the mast cells of the nasal mucosa and submucosa generate and release mediators

2122 through IgE-dependent reactions that are capable of producing tissue edema and eosinophilic infiltration.

DIAGNOSIS

The diagnosis of seasonal allergic rhinitis depends largely on an accurate history of occurrence coincident with the pollination of the offending weeds, grasses, or trees. The continuous character of perennial allergic rhinitis due to contamination of the home or place of work makes historic analysis difficult, but there may be variability in symptoms that can be related to exposure to animal dander, dust mite and/or cockroach allergens, fungal spores, or work-related allergens such as latex. Patients with perennial rhinitis commonly develop the problem in adult life, and manifest nasal congestion and a postnasal discharge, often associated with thickening of the sinus membranes demonstrated by radiography. Perennial nonallergic rhinitis with eosinophilia syndrome (NARES) occurs in the middle decades of life and is characterized by nasal obstruction, anosmia, chronic sinusitis, and frequent aspirin intolerance. The term *vasomotor rhinitis* or *perennial nonallergic rhinitis* designates a condition of enhanced reactivity of the nasopharynx in which a symptom complex resembling perennial allergic rhinitis occurs with nonspecific stimuli, including chemical odors, temperature and humidity variations, and position changes but occurs without tissue eosinophilia or an allergic etiology. Other entities to be excluded are structural abnormalities of the nasopharynx; exposure to irritants; gustatory rhinitis associated with cholinergic activation that occurs while eating or ingesting alcohol; hypothyroidism; upper respiratory tract infection; pregnancy with prominent nasal mucosal edema; prolonged topical use of α -adrenergic agents in the form of nose drops (rhinitis medicamentosa); and the use of certain therapeutic agents such as rauwolfia, β -adrenergic antagonists, estrogens, progesterone, ACE inhibitors, aspirin and other NSAIDs, and drugs for erectile dysfunction (phosphodiesterase-5 inhibitors).

The nasal secretions of allergic patients are rich in eosinophils, and a modest peripheral eosinophilia is a common feature. Local or systemic neutrophilia implies infection. Total serum IgE is frequently elevated, but the demonstration of immunologic specificity for IgE is critical to an etiologic diagnosis. A skin test by the intracutaneous route (puncture or prick) with the allergens of interest provides a rapid and reliable approach to identifying allergen-specific IgE that has sensitized cutaneous mast cells. A positive intracutaneous skin test with 1:10–1:20 weight/volume of extract has a high predictive value for the presence of allergy. An intradermal test with a 1:500–1:1000 dilution of 0.05 mL may follow if indicated by history when the intracutaneous test is negative, but while more sensitive, it is less reliable due to the reactivity of some asymptomatic individuals at the test dose. Skin testing by the intracutaneous route for food allergens can be supportive of the clinical history. A double-blind, placebo-controlled challenge may document a food allergy, but such a procedure does bear the risk of an anaphylactic reaction. An elimination diet is safer but is tedious and less definitive. Food allergy is uncommon as a cause of allergic rhinitis.

Newer methodology for detecting total IgE, including the development of enzyme-linked immunosorbent assays (ELISA) employing anti-IgE bound to either a solid-phase or a liquid-phase particle, provides rapid and cost-effective determinations. Measurements of specific anti-IgE in serum are obtained by its binding to an allergen and quantitation by subsequent uptake of labeled anti-IgE. As compared to the skin test, the assay of specific IgE in serum is less sensitive but has high specificity.

PREVENTION

Avoidance of exposure to the offending allergen is the most effective means of controlling allergic diseases; removal of pets from the home to avoid animal danders, utilization of air-filtration devices to minimize the concentrations of airborne pollens, elimination of cockroach-derived proteins by chemical destruction of the pest and careful food storage, travel to areas where the allergen is not being generated, and even a change of domicile to eliminate a mold spore problem may be necessary. Control of dust mites by allergen avoidance includes use of plastic-lined covers for mattresses, pillows, and comforters; using a

filter-equipped vacuum cleaner; washing bedding and clothes at temperatures $>54.5^{\circ}\text{C}$ (above 130°F); and elimination of carpets and drapes.

TREATMENT ALLERGIC RHINITIS

Although allergen avoidance is the most cost-effective means of managing allergic rhinitis, treatment with pharmacologic agents represents the standard approach to seasonal or perennial allergic rhinitis. Oral H_1 antihistamines are effective for nasopharyngeal itching, sneezing, and watery rhinorrhea and for such ocular manifestations as itching, tearing, and erythema, but they are less efficacious for the nasal congestion. The older antihistamines are sedating, and they induce psychomotor impairment, including reduced eye-hand coordination and impaired automobile driving skills. Their anticholinergic (muscarinic) effects include visual disturbance, urinary retention, and constipation. Because the newer H_1 antihistamines such as fexofenadine, loratadine, desloratadine, cetirizine, levocetirizine, olopatadine, bilastine, and azelastine are less lipophilic and more H_1 selective, their ability to cross the blood-brain barrier is reduced, and thus their sedating and anticholinergic side effects are minimized. These newer antihistamines do not differ appreciably in efficacy for relief of rhinitis and/or sneezing. Azelastine nasal spray may benefit individuals with nonallergic vasomotor rhinitis, but it has an adverse effect of dysgeusia (taste perversion) in some patients. Because antihistamines have little effect on congestion, α -adrenergic agents such as phenylephrine or oxymetazoline are generally used topically to alleviate nasal congestion and obstruction. However, the duration of their efficacy is limited because of rebound rhinitis (i.e., 7- to 14-day use can lead to rhinitis medicamentosa) and such systemic responses as hypertension. Oral α -adrenergic agonist decongestants containing pseudoephedrine are standard for the management of nasal congestion, generally in combination with an antihistamine. While oral antihistamines typically reduce nasal and ocular symptoms by about one-third, pseudoephedrine must be added to achieve a similar reduction in nasal congestion. These pseudoephedrine combination products can cause insomnia and are precluded from use in patients with narrow angle glaucoma, urinary retention, severe hypertension, marked coronary artery disease, or a first-trimester pregnancy. The CysLT_1 blocker montelukast is approved for treatment of both seasonal and perennial rhinitis, and it reduces both nasal and ocular symptoms by about 20%. Cromolyn sodium, a nasal spray, is essentially without side effects and is used prophylactically on a continuous basis during the season. Intranasal high-potency glucocorticoids are the most potent drugs available for the relief of established rhinitis, seasonal or perennial, and are effective in relieving nasal congestion. They provide efficacy with substantially reduced side effects as compared with this same class of agent administered orally. Their most frequent side effect is local irritation, with *Candida* overgrowth being a rare occurrence. The currently available intranasal glucocorticoids—beclomethasone, flunisolide, triamcinolone, budesonide, fluticasone propionate, fluticasone furoate, ciclesonide, and mometasone furoate—are equally effective for nasal symptom relief, including nasal congestion; these agents all achieve up to 70% overall symptom relief with some variation in the time period for onset of benefit. Topical ipratropium is an anticholinergic agent effective in reducing rhinorrhea, including that in patients with perennial symptoms, and it can be additionally efficacious when combined with intranasal glucocorticoids. Local treatment with cromolyn sodium is effective in treating mild allergic conjunctivitis. Topical antihistamines such as olopatadine, azelastine, ketotifen, or epinastine administered to the eye provide rapid relief of itching and redness and are more effective than oral antihistamines.

Immunotherapy, often termed *hyposensitization*, consists of repeated subcutaneous injections of gradually increasing concentrations of the allergen(s) considered to be specifically responsible for the symptom complex. Controlled studies of ragweed, grass, dust mite, and cat dander allergens administered for treatment of allergic rhinitis have demonstrated at least partial relief of symptoms and

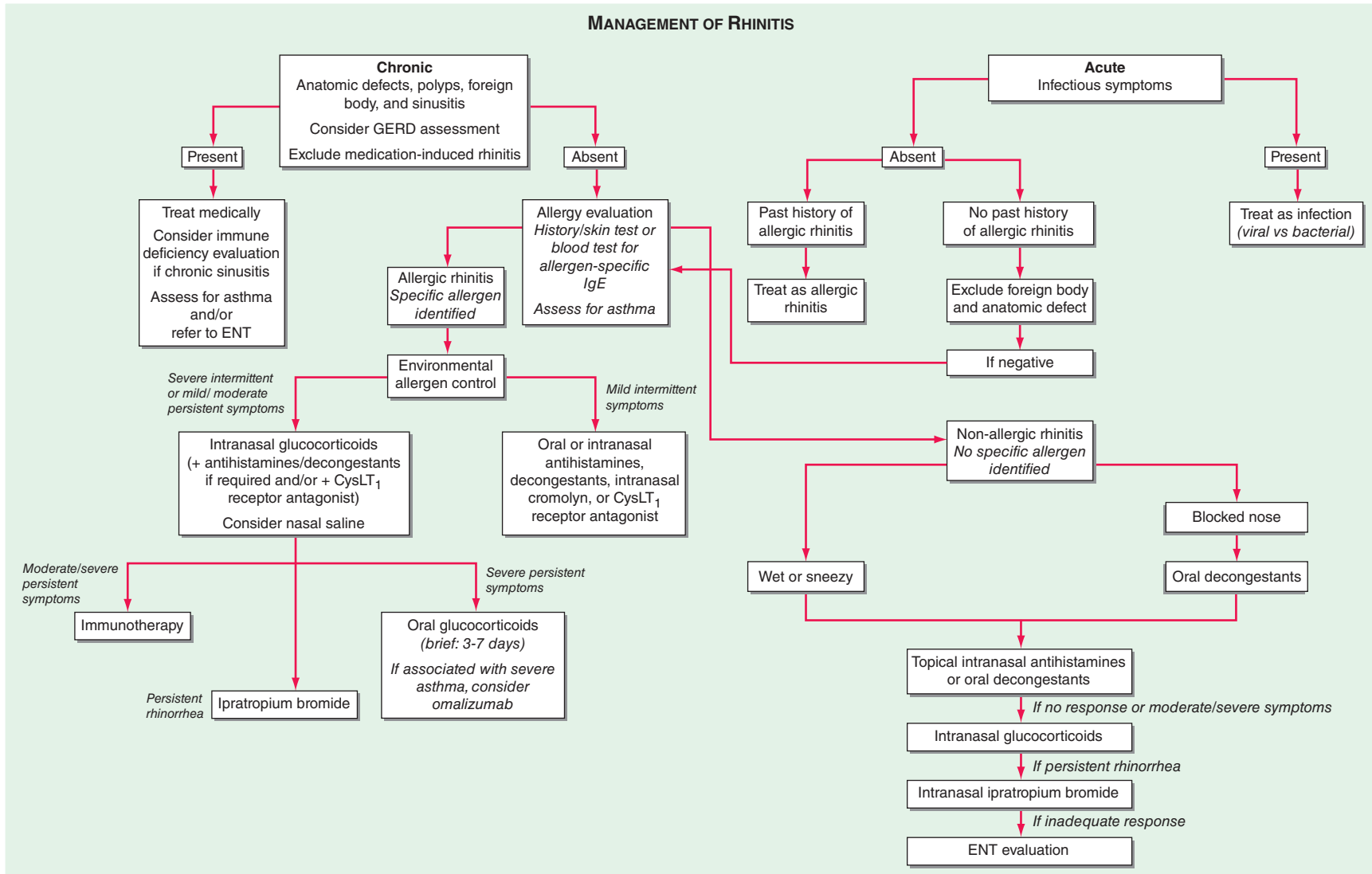


FIGURE 376-4 Algorithm for the diagnosis and management of rhinitis. ENT, ear, nose, and throat; GERD, gastroesophageal reflux disease.

2124 signs. The duration of such immunotherapy is 3–5 years, with discontinuation being based on minimal symptoms over two consecutive seasons of exposure to the allergen. Clinical benefit appears related to the administration of a high dose of relevant allergen, advancing from weekly to monthly intervals. Patients should remain at the treatment site for at least 20 minutes after allergen administration so that any anaphylactic consequence can be managed. Local reactions with erythema and induration are not uncommon and may persist for 1–3 days. Immunotherapy is contraindicated in patients with significant cardiovascular disease or unstable asthma and should be conducted with particular caution in any patient requiring β -adrenergic blocking therapy because of the difficulty in managing an anaphylactic complication. The response to immunotherapy is associated with a complex of cellular and humoral effects that includes a modulation in T cell cytokine production. Immunotherapy should be reserved for clearly documented seasonal or perennial rhinitis that is clinically related to defined allergen exposure with confirmation by the presence of allergen-specific IgE. Systemic treatment with a monoclonal antibody to IgE (omalizumab) that blocks mast cell and basophil sensitization has efficacy for allergic rhinitis and can be used with immunotherapy to enhance safety and efficacy. However, current approval is only for treatment of patients with persistent allergic asthma not controlled by inhaled glucocorticoid therapy. A sequence for the management of allergic or perennial rhinitis based on an allergen-specific diagnosis and stepwise management as required for symptom control would include the following: (1) identification of the offending allergen(s) by history with confirmation of the presence of allergen-specific IgE by skin test and/or serum assay; (2) avoidance of the offending allergen; and (3) medical management in a stepwise fashion (**Fig. 376-4**). Mild intermittent symptoms of allergic rhinitis are treated with oral antihistamines, oral CysLT₁ receptor antagonists, intranasal antihistamines, or intranasal cromolyn prophylaxis. Moderate to more severe allergic rhinitis is managed with intranasal glucocorticoids plus oral antihistamines, oral CysLT₁ receptor antagonists, or antihistamine-decongestant combinations. Persistent allergic rhinitis requiring the daily use of intranasal glucocorticoids with add-on interventions such as oral antihistamines, decongestant combinations, or topical ipratropium merits consideration of allergen-specific immunotherapy. Even a brief course of oral prednisone can be indicated for rapid relief of severe allergic rhinitis symptoms.

377e Autoimmunity and Autoimmune Diseases

Betty Diamond, Peter E. Lipsky

One of the central features of the immune system is the capacity to mount an inflammatory response to potentially harmful foreign materials while avoiding damage to self-tissues. Whereas recognition of self plays an important role in shaping the repertoires of immune receptors on both T and B cells and in clearing apoptotic and other tissue debris from sites throughout the body, the development of potentially harmful immune responses to self-antigens is, in general, prohibited. The essential feature of an *autoimmune disease* is that tissue injury is caused by the immunologic reaction of the organism against its own tissues. *Autoimmunity*, on the other hand, refers merely to the presence of antibodies or T lymphocytes that react with self-antigens and does not necessarily imply that the self-reactivity has pathogenic consequences. Autoimmunity is present in all individuals; however, autoimmune disease occurs only in those individuals in whom the breakdown of one or more of the basic mechanisms regulating immune tolerance results in self-reactivity that can cause tissue damage.

Autoimmunity is seen in normal individuals, with a higher frequency among normal older people. Polyreactive autoantibodies that recognize many host antigens are present throughout life. Expression of these autoantibodies may be increased after some inciting events. These antibodies are usually of the IgM heavy chain isotype and are encoded by nonmutated germline immunoglobulin variable region genes. When autoimmunity is induced by an inciting event, such as infection or tissue damage from trauma or ischemia, the autoreactivity is in general self-limited. When such autoimmunity does persist, however, pathology may or may not result. Even in the presence of organ pathology, it may be difficult to determine whether the damage is mediated by autoreactivity. After an inciting event, the development of self-reactivity may be the consequence of an ongoing pathologic process, may be nonpathogenic, or may contribute to tissue inflammation and damage. Individuals with autoimmune disease may have numerous autoantibodies, only some or even none of which may be pathogenic. Patients with systemic sclerosis may have a wide array of antinuclear antibodies that are important in disease classification but are not clearly pathogenic; patients with pemphigus may also exhibit a wide array of autoantibodies, only one of which (antibody to desmoglein) is known to be pathogenic.

MECHANISMS OF AUTOIMMUNITY

Since Ehrlich first postulated the existence of mechanisms to prevent the generation of self-reactivity in the early 1900s, ideas concerning the nature of this prohibition have developed in parallel with a progressive increase in understanding of the immune system. Burnet's clonal selection theory included the idea that interaction of lymphoid cells with their specific antigens during fetal or early postnatal life would lead to elimination of such "forbidden clones." This idea was refuted, however, when it was shown that autoimmune diseases could be induced in experimental animals by simple immunization procedures, that autoantigen-binding cells could be demonstrated easily in the circulation of normal individuals, and that self-limited autoimmune phenomena frequently developed after tissue damage from infection or trauma. These observations indicated that clones of cells capable of responding to autoantigens were present in the repertoire of antigen-reactive cells in normal adults and suggested that mechanisms in addition to clonal deletion were responsible for preventing their activation.

Currently, three general processes are thought to be involved in the maintenance of selective unresponsiveness to autoantigens (**Table 377e-1**): (1) sequestration of self-antigens, rendering them inaccessible to the immune system; (2) specific unresponsiveness (tolerance or anergy) of relevant T or B cells; and (3) limitation of potential reactivity by regulatory mechanisms. Derangements of these normal

TABLE 377e-1 MECHANISMS PREVENTING AUTOIMMUNITY

1. Sequestration of self-antigens
2. Generation and maintenance of tolerance
 - a. Central deletion of autoreactive lymphocytes
 - b. Peripheral anergy of autoreactive lymphocytes
 - c. Receptor replacement in autoreactive lymphocytes
3. Regulatory mechanisms
 - a. Regulatory T cells
 - b. Regulatory B cells
 - c. Regulatory mesenchymal cells
 - d. Regulatory cytokines
 - e. Idiotype network

processes may predispose to the development of autoimmunity (**Table 377e-2**). In general, these abnormal responses require both an exogenous trigger, such as bacterial or viral infection or cigarette smoking, and the presence of endogenous abnormalities in the cells of the immune system. Microbial superantigens, such as staphylococcal protein A and staphylococcal enterotoxins, are substances that can stimulate a broad range of T and B cells through specific interactions with selected families of immune receptors, irrespective of their antigen specificity. If autoantigen-reactive T and/or B cells express these receptors, autoimmunity may develop. Alternatively, molecular mimicry or cross-reactivity between a microbial product and a self-antigen may lead to activation of autoreactive lymphocytes. One of the best examples of autoreactivity and autoimmune disease resulting from molecular mimicry is rheumatic fever, in which antibodies to the M protein of streptococci cross-react with myosin, laminin, and other matrix proteins as well as with neuronal antigens. Deposition of these autoantibodies in the heart initiates an inflammatory response, whereas their penetration into the brain can result in Sydenham's chorea. Molecular mimicry between microbial proteins and host tissues has been reported in type 1 diabetes mellitus, rheumatoid arthritis, celiac disease, and multiple sclerosis. It is presumed that infectious agents may be able to overcome self-tolerance because they possess *pathogen-associated molecular patterns* (PAMPs). These molecules (e.g., bacterial endotoxin, RNA, or DNA) exert adjuvant-like effects on the immune system by interacting with *Toll-like receptors* (TLRs) and other *pattern recognition receptors* (PRRs)

TABLE 377e-2 MECHANISMS OF AUTOIMMUNITY

- I. Exogenous
 - A. Molecular mimicry
 - B. Superantigenic stimulation
 - C. Microbial and tissue damage-associated adjuvanticity
- II. Endogenous
 - A. Altered antigen presentation
 1. Loss of immunologic privilege
 2. Presentation of novel or cryptic epitopes (epitope spreading)
 3. Alteration of self-antigen
 4. Enhanced function of antigen-presenting cells
 - a. Costimulatory molecule expression
 - b. Cytokine production
 - B. Increased T cell help
 1. Cytokine production
 2. Costimulatory molecules
 - C. Increased B cell function
 1. B cell activating factor
 2. Costimulatory molecules
 - D. Apoptotic defects or defects in clearance of apoptotic material
 - E. Cytokine imbalance
 - F. Altered immunoregulation
 - G. Endocrine abnormalities

377e-2 that increase the immunogenicity and immunostimulatory capacity of the microbial material. The adjuvants activate dendritic cells through TLRs, which in turn stimulate the activation of previously quiescent lymphocytes that recognize both microbial antigens and self-antigens. Similarly, cellular and tissue damage due to the release of *damage-associated molecular patterns* (DAMPs), including DNA, RNA nucleosomes, and other tissue debris, may activate cells of the inflammatory and immune systems through engagement of the same array of PRRs.

Endogenous derangements of the immune system may also contribute to the loss of immunologic tolerance to self-antigens and the development of autoimmunity (Table 377e-2). Some autoantigens reside in immunologically privileged sites, such as the brain or the anterior chamber of the eye. These sites are characterized by the inability of engrafted tissue to elicit immune responses. Immunologic privilege results from a number of events, including the limited entry of proteins from those sites into lymphatics, the local production of immunosuppressive cytokines such as transforming growth factor β , and the local expression of molecules (including Fas ligand) that can induce apoptosis of activated T cells. Lymphoid cells remain in a state of immunologic ignorance (neither activated nor anergized) with regard to proteins expressed uniquely in immunologically privileged sites. If the privileged site is damaged by trauma or inflammation or if T cells are activated elsewhere, proteins expressed at this site can become immunogenic and also be the targets of immunologic assault. In multiple sclerosis and sympathetic ophthalmia, for example, antigens uniquely expressed in the brain and eye, respectively, become the target of activated T cells.

Alterations in antigen presentation may also contribute to autoimmunity. Peptide determinants (*epitopes*) of a self-antigen that are not routinely presented to lymphocytes may be recognized as a result of altered proteolytic processing of the molecule and the ensuing presentation of novel peptides (*cryptic epitopes*). When B cells rather than dendritic cells present self-antigen, they may also present cryptic epitopes that can activate autoreactive T cells. These cryptic epitopes will not previously have been available to effect the silencing of autoreactive lymphocytes. Furthermore, once there is immunologic recognition of one protein component of a multimolecular complex, reactivity may be induced to other components of the complex after internalization and presentation of all molecules within the complex (epitope spreading). Finally, inflammation, environmental agents, drug exposure, or normal senescence may cause a post-translational alteration in proteins, resulting in the generation of immune responses that cross-react with normal self-proteins. For example, the induction and/or release of protein arginine deiminase enzymes results in the conversion of arginine residues to citrullines in a variety of proteins, thereby altering their capacity to induce immune responses. Production of antibodies to citrullinated proteins has been observed in rheumatoid arthritis and chronic lung disease as well as in normal smokers and may contribute to organ pathology. Alterations in the availability and presentation of autoantigens may be important components of immunoreactivity in certain models of organ-specific autoimmune diseases. In addition, these factors may be relevant to an understanding of the pathogenesis of various drug-induced autoimmune conditions. However, the diversity of autoreactivity manifesting in non-organ-specific systemic autoimmune diseases suggests that these conditions may result from a more general activation of the immune system rather than from an alteration in individual self-antigens.

Many autoimmune diseases are characterized by the presence of antibodies that react with apoptotic material. Defects in the clearance of apoptotic material have been shown to elicit auto-immunity and autoimmune disease in a number of animal models. Moreover, such defects have been found in patients with systemic lupus erythematosus (SLE). Apoptotic debris that is not cleared quickly by the immune system can function as endogenous ligands for a number of PRRs on dendritic cells and B cells. Under such circumstances, dendritic cells and/or B cells are activated, and an immune response to apoptotic debris can develop. In addition, the presence of extracellular apoptotic material within germinal centers of secondary lymphoid organs in patients with SLE may facilitate the direct activation of autoimmune

B cell clones or may function to select such clones during immune responses.

Studies in a number of experimental models have suggested that intense stimulation of T lymphocytes can produce nonspecific signals that bypass the need for antigen-specific helper T cells and lead to polyclonal B cell activation with the formation of multiple autoantibodies. For example, antinuclear, antierythrocyte, and antilymphocyte antibodies are produced during the chronic graft-versus-host reaction. In addition, true autoimmune diseases, including autoimmune hemolytic anemia and immune complex-mediated glomerulonephritis, can be induced in this manner. While such diffuse activation of helper T cell activity clearly can cause autoimmunity, nonspecific stimulation of B lymphocytes can also lead to the production of autoantibodies. Thus, the administration of polyclonal B cell activators, such as bacterial endotoxin, to normal mice leads to the production of a number of autoantibodies, including those to DNA and IgG (rheumatoid factor). A variety of genetic modifications resulting in hyperresponsiveness of B cells also can lead to the production of autoantibodies and, in animals of appropriate genetic background, a lupus-like syndrome. Moreover, excess B cell activating factor (BAFF), a B cell survival factor, can cause T cell-independent B cell activation and the development of autoimmunity. SLE can also be induced in mice through exuberant dendritic cell activation, through a redundancy of TLR7 on the Y chromosome (as in BXSB-Yaa mice), or through exposure to CpG, a ligand for TLR9. The ensuing induction of inflammatory mediators can cause a switch from the production of nonpathogenic IgM autoantibodies to the production of pathogenic IgG autoantibodies in the absence of antigen-specific T cell help. Aberrant selection of the B or T cell repertoire at the time of antigen receptor expression can also predispose to autoimmunity. For example, B cell immunodeficiency caused by an absence of the B cell receptor-associated kinase (Bruton's tyrosine kinase) leads to X-linked agammaglobulinemia. This syndrome is characterized by reduced B cell numbers. This leads to high levels of BAFF which alter B cell selection and lead to greater survival of autoreactive B cells. Likewise, negative selection of autoreactive T cells in the thymus requires expression of the autoimmune regulator (AIRE) gene that enables the expression of tissue-specific proteins in thymic medullary epithelial cells. Peptides from these proteins are expressed in the context of major histocompatibility complex (MHC) molecules and mediate the central deletion of autoreactive T cells. The absence of AIRE gene expression leads to a failure of negative selection of autoreactive cells, autoantibody production, and severe inflammatory destruction of multiple organs. Individuals deficient in AIRE gene expression develop autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED).

Primary alterations in the activity of T and/or B cells, cytokine imbalances, or defective immunoregulatory circuits may also contribute to the emergence of autoimmunity. Diminished production of tumor necrosis factor (TNF) and interleukin (IL) 10 has been reported to be associated with the development of autoimmunity. Overproduction or therapeutic administration of type 1 interferon has also been associated with autoimmunity. Overexpression of costimulatory molecules on T cells similarly can lead to autoantibody production.

Autoimmunity may also result from an abnormality of immunoregulatory mechanisms. Observations made in both human autoimmune disease and animal models suggest that defects in the generation and expression of regulatory T cell (Treg) activity may allow the production of autoimmunity. It has recently been appreciated that the IPEX (immunodysregulation, polyendocrinopathy, enteropathy X-linked) syndrome results from the failure to express the FOXP3 gene, which encodes a molecule critical in the differentiation of Tregs. Administration of normal Tregs or of factors derived from them can prevent the development of autoimmune disease in rodent models of autoimmunity, and allogeneic stem cell transplantation ameliorates human IPEX. Abnormalities in the function of Tregs have been noted in a number of human autoimmune diseases, including rheumatoid arthritis and SLE, although it remains uncertain whether these functional abnormalities are causative or are secondary to inflammation. One of the mechanisms by which Tregs control immune/inflammatory

responses is by the production of the cytokine IL-10. In this regard, children with a deficiency in the expression of IL-10 or the IL-10 receptor develop inflammatory bowel disease that mimics Crohn's disease and that can be cured by allogeneic stem cell transplantation. Finally, recent data indicate that B cells may also exert regulatory function, largely through the production of IL-10. Deficiency of IL-10-producing regulatory B cells can prolong the course of multiple sclerosis in an animal model, and such cells are thought to be functionally diminished in human SLE.

It should be apparent that no single mechanism can explain all the varied manifestations of autoimmunity or autoimmune disease. Furthermore, genetic evaluation has shown that convergence of a number of abnormalities is often required for the induction of an autoimmune disease. Additional factors that appear to be important determinants in the induction of autoimmunity include age, sex (many autoimmune diseases are far more common in women), exposure to infectious agents, and environmental contacts. How all of these disparate factors affect the capacity to develop self-reactivity is currently being investigated intensively.

GENETIC CONSIDERATIONS



Evidence in humans that there are susceptibility genes for autoimmunity comes from family studies and especially from studies of twins. Studies in type 1 diabetes mellitus, rheumatoid arthritis, multiple sclerosis, and SLE have shown that ~15–30% of pairs of monozygotic twins show disease concordance, whereas the figure is <5% for dizygotic twins. The occurrence of different autoimmune diseases within the same family has suggested that certain susceptibility genes may predispose to a variety of autoimmune diseases. Genome-wide association studies have begun to identify polymorphisms in individual genes that are associated with specific autoimmune diseases. More than 50 genetic polymorphisms associated with one or more autoimmune diseases have been identified to date. It is notable that some genes are associated with multiple autoimmune diseases, whereas others are specifically associated with only one autoimmune condition. Moreover, recent genetic evidence suggests that clusters of genetic risk factors can commonly be found in groups of autoimmune diseases. Four general clusters have been identified: one group of 6 genetic polymorphisms most frequently associated with Crohn's disease, psoriasis, and multiple sclerosis; a second cluster of 8 polymorphisms most strongly associated with celiac disease, rheumatoid arthritis, and SLE; a third cluster of 7 polymorphisms most strongly associated with type 1 diabetes, multiple sclerosis, and rheumatoid arthritis; and a fourth cluster of more than 12 polymorphisms most strongly associated with type 1 diabetes, rheumatoid arthritis, celiac disease, Crohn's disease, and SLE. These results imply that autoimmune diseases with widely different clinical presentations and patterns of organ involvement could involve similar immunopathogenic pathways. For example, the same allele of the gene encoding PTPN22 is associated with multiple autoimmune diseases. Its product is a phosphatase expressed by a variety of hematopoietic cells that down-regulates antigen receptor-mediated stimulation of T and B cells. The risk allele is associated with type 1 diabetes mellitus, rheumatoid arthritis, and SLE in some populations. The explanation for the association of this polymorphism with autoimmune disease is uncertain, but it is likely that it diminishes antigen receptor signaling during lymphocyte development, permitting escape of autoreactive clones or decreased activation-induced apoptosis of autoantigen-reactive lymphocytes in the periphery. In recent

years, genome-wide association studies have demonstrated a variety of other genes that are involved in human autoimmune diseases. Most genes individually confer a relatively low risk for autoimmune diseases and are found in normal individuals. No gene has been identified that is essential for autoimmune diseases. In addition to this evidence from humans, certain inbred mouse strains reproducibly develop specific spontaneous or experimentally induced autoimmune diseases, whereas others do not. These findings have led to an extensive search for genes that determine susceptibility to autoimmune disease and for genes that might be protective.

The strongest consistent association for susceptibility to autoimmune disease is with particular MHC alleles. It has been suggested that the association of MHC genotype with autoimmune disease relates to differences in the ability of different allelic variations of MHC molecules to present autoantigenic peptides to autoreactive T cells. An alternative hypothesis involves the role of MHC alleles in shaping the T cell receptor repertoire during T cell ontogeny in the thymus. In addition, specific MHC gene products may themselves be the source of peptides that can be recognized by T cells. Cross-reactivity between such MHC peptides and peptides derived from proteins produced by common microbes may trigger autoimmunity by molecular mimicry. However, MHC genotype alone does not determine the development of autoimmunity. Identical twins are far more likely to develop the same autoimmune disease than MHC-identical nontwin siblings; this observation suggests that genetic factors other than the MHC affect disease susceptibility. Studies of the genetics of type 1 diabetes mellitus, SLE, rheumatoid arthritis, and multiple sclerosis in humans and mice have identified several independently segregating disease susceptibility loci in addition to the MHC. Genes that encode molecules of the innate immune response are also involved in autoimmunity. In humans, inherited homozygous deficiency of the early proteins of the classic pathway of complement (C1q, C4, or C2) as well as genes involved in the type 1 interferon pathway are very strongly associated with the development of SLE.

IMMUNOPATHOGENIC MECHANISMS IN AUTOIMMUNE DISEASES

The mechanisms of tissue injury in autoimmune diseases can be divided into antibody-mediated and cell-mediated processes. Representative examples are listed in [Table 377e-3](#).

TABLE 377e-3 MECHANISMS OF TISSUE DAMAGE IN AUTOIMMUNE DISEASE

Effector	Mechanism	Target	Disease
Autoantibody	Blocking or inactivation	α Chain of the nicotinic acetylcholine receptor	Myasthenia gravis
		Phospholipid- β_2 -glycoprotein I complex	Antiphospholipid syndrome
		Insulin receptor	Insulin-resistant diabetes mellitus
		Intrinsic factor	Pernicious anemia
	Stimulation	TSH receptor (LATS)	Graves' disease
		Proteinase-3 (ANCA)	Granulomatosis with polyangiitis
		Epidermal cadherin	Pemphigus vulgaris
		Desmoglein 3	
	Complement activation	α_3 Chain of collagen IV	Goodpasture's syndrome
	Immune complex formation	Double-stranded DNA	Systemic lupus erythematosus
Opsonization		Immunoglobulin	Rheumatoid arthritis
		Platelet GpIIb/IIIa	Autoimmune thrombocytopenic purpura
		Rh antigens, I antigen	Autoimmune hemolytic anemia
Antibody-dependent cellular cytotoxicity		Thyroid peroxidase, thyroglobulin	Hashimoto's thyroiditis
T cells	Cytokine production	?	Rheumatoid arthritis, multiple sclerosis, type 1 diabetes mellitus
	Cellular cytotoxicity	?	Type 1 diabetes mellitus

Abbreviations: ANCA, antineutrophil cytoplasmic antibody; LATS, long-acting thyroid stimulator; TSH, thyroid-stimulating hormone.

The pathogenicity of autoantibodies can be mediated through several mechanisms, including opsonization of soluble factors or cells, activation of an inflammatory cascade via the complement system, and interference with the physiologic function of soluble molecules or cells.

In autoimmune thrombocytopenic purpura, opsonization of platelets targets them for elimination by phagocytes. Likewise, in autoimmune hemolytic anemia, binding of immunoglobulin to red cell membranes leads to phagocytosis and lysis of the opsonized cell. Goodpasture's syndrome, a disease characterized by lung hemorrhage and severe glomerulonephritis, represents an example of antibody binding leading to local activation of complement and neutrophil accumulation and activation. The autoantibody in this disease binds to the α_3 chain of type IV collagen in the basement membrane. In SLE, activation of the complement cascade at sites of immunoglobulin deposition in renal glomeruli is considered to be a major mechanism of renal damage. Moreover, the DNA- and RNA-containing immune complexes in SLE activate TLR9 and TLR7, respectively, in dendritic cells and promote a proinflammatory, immunogenic milieu conducive to amplification of the autoimmune response.

Autoantibodies can also interfere with normal physiologic functions of cells or soluble factors. Autoantibodies to hormone receptors can lead to stimulation of cells or to inhibition of cell function through interference with receptor signaling. For example, long-acting thyroid stimulators—autoantibodies that bind to the receptor for thyroid-stimulating hormone (TSH)—are present in Graves' disease and function as agonists, causing the thyroid to respond as if there were an excess of TSH. Alternatively, antibodies to the insulin receptor can cause insulin-resistant diabetes mellitus through receptor blockade. In myasthenia gravis, autoantibodies to the acetylcholine receptor can be detected in 85–90% of patients and are responsible for muscle weakness. The exact location of the antigenic epitope, the valence and affinity of the antibody, and perhaps other characteristics determine whether activation or blockade results from antibody binding.

Antiphospholipid antibodies are associated with thromboembolic events in primary and secondary antiphospholipid syndrome and have also been associated with fetal wastage. The major antibody is directed to the phospholipid- β_2 -glycoprotein I complex and appears to exert a procoagulant effect. In pemphigus vulgaris, autoantibodies bind to desmoglein 3, a component of the epidermal cell desmosome, and play a role in the induction of the disease. These antibodies exert their pathologic effect by disrupting cell–cell junctions through stimulation of the production of epithelial proteases, with consequent blister formation. Cytoplasmic antineutrophil cytoplasmic antibody (c-ANCA), found in granulomatosis with polyangiitis, is an antibody to an intracellular antigen, the 29-kDa serine protease (proteinase-3). In vitro experiments have shown that IgG anti-c-ANCA causes cellular activation and degranulation of primed neutrophils.

It is important to note that autoantibodies of a given specificity may cause disease only in genetically susceptible hosts, as has been shown in experimental models of myasthenia gravis, SLE, rheumatic fever, and rheumatoid arthritis. Furthermore, once organ damage is initiated, new inflammatory cascades are initiated that can sustain and amplify the autoimmune process. Finally, some autoantibodies seem to be markers for disease but have, as yet, no known pathogenic potential.

AUTOIMMUNE DISEASES

Manifestations of autoimmunity are found in a large number of pathologic conditions. However, their presence does not necessarily imply that the pathologic process is an autoimmune disease. A number of attempts to establish formal criteria for the classification of diseases as autoimmune have been made, but none is universally accepted. One set of criteria is shown in [Table 377e-4](#); however, this scheme should be viewed merely as a guide in consideration of the problem.

To classify a disease as autoimmune, it is necessary to demonstrate that the immune response to a self-antigen causes the observed pathology. Initially, the detection of antibodies to the affected tissue in the serum of patients suffering from various diseases was taken as evidence that these diseases had an autoimmune basis. However, such

TABLE 377e-4 HUMAN AUTOIMMUNE DISEASE: PRESUMPTIVE EVIDENCE FOR IMMUNOLOGIC PATHOGENESIS

Major Criteria
1. Presence of autoantibodies or evidence of cellular reactivity to self
2. Documentation of relevant autoantibody or lymphocytic infiltrate in the pathologic lesion
3. Demonstration that relevant autoantibody or T cells can cause tissue pathology <ol style="list-style-type: none"> Transplacental transmission Adaptive transfer into animals In vitro impact on cellular function
Supportive Evidence
1. Reasonable animal model
2. Beneficial effect from immunosuppressive agents
3. Association with other evidence of autoimmunity
4. No evidence of infection or other obvious cause

autoantibodies are also found when tissue damage is caused by trauma or infection and in these cases are secondary to tissue damage. Thus, autoimmunity must be shown to be pathogenic before a disease is categorized as autoimmune.

To confirm autoantibody pathogenicity, it may be possible to transfer disease to experimental animals by the administration of autoantibodies from a patient, with the subsequent development of pathology in the recipient similar to that seen in the patient. This scenario has been documented, for example, in Graves' disease. Some autoimmune diseases can be transferred from mother to fetus and are observed in the newborn babies of diseased mothers. The symptoms of the disease in the newborn usually disappear as the levels of maternal antibody decrease. An exception, however, is congenital heart block, in which damage to the developing conducting system of the heart follows in utero transfer of anti-Ro antibody from the mother to the fetus. This antibody transfer can result in a permanent developmental defect in the heart.

In most situations, the critical factors that determine when the development of autoimmunity results in autoimmune disease have not been delineated. The relationship of autoimmunity to the development of autoimmune disease may be associated with the fine specificity of the antibodies or T cells or their specific effector capabilities. In many circumstances, a mechanistic understanding of the pathogenic potential of autoantibodies has not been established. In some autoimmune diseases, biased production of cytokines by helper T (T_H) cells may play a role in pathogenesis. In this regard, T cells can differentiate into specialized effector cells that predominantly produce interferon γ (T_H1), IL-4 (T_H2), or IL-17 (T_H17) or that provide help to B cells (T follicular helper, T_{FH}) ([Chap. 372e](#)). T_H1 cells facilitate macrophage activation and classic cell-mediated immunity, whereas T_H2 cells are thought to have regulatory functions and are involved in the resolution of normal immune responses as well as in the development of responses to a variety of parasites. T_H17 cells produce a number of inflammatory cytokines, including IL-17 and IL-22, and seem to be prominently involved in host resistance to certain fungal infections. T_{FH} cells help B cells by constitutively producing IL-21. In a number of autoimmune diseases, such as rheumatoid arthritis, multiple sclerosis, type 1 diabetes mellitus, and Crohn's disease, there appears to be biased differentiation of T_H1 and T_H17 cells, with resultant organ damage. Studies suggest an accentuated differentiation of T_H17 cells associated with animal models of inflammatory arthritis, whereas increased differentiation of T_{FH} cells has been associated with animal models of SLE.

ORGAN-SPECIFIC VERSUS SYSTEMIC AUTOIMMUNE DISEASES

The spectrum of autoimmune diseases ranges from conditions specifically affecting a single organ to systemic disorders that involve many organs ([Table 377e-5](#)). Hashimoto's autoimmune thyroiditis is an example of an organ-specific autoimmune disease ([Chap. 405](#)). In this

TABLE 377e-5 DISEASES ON THE AUTOIMMUNE SPECTRUM

Organ Specific	
Graves' disease	Vitiligo
Hashimoto's thyroiditis	Autoimmune hemolytic anemia
Autoimmune polyglandular syndrome	Autoimmune thrombocytopenic purpura
Type 1 diabetes mellitus	Pernicious anemia
Insulin-resistant diabetes mellitus	Myasthenia gravis
Immune-mediated infertility	Multiple sclerosis
Autoimmune Addison's disease	Guillain-Barré syndrome
Pemphigus vulgaris	Stiff-man syndrome
Pemphigus foliaceus	Acute rheumatic fever
Dermatitis herpetiformis	Sympathetic ophthalmia
Autoimmune alopecia	Goodpasture's syndrome
Organ Nonspecific (Systemic)	
Systemic lupus erythematosus	Granulomatosis with polyangiitis
Rheumatoid arthritis	Antiphospholipid syndrome
Systemic necrotizing vasculitis	Sjögren's syndrome

disorder, a specific lesion in the thyroid is associated with infiltration of mononuclear cells and damage to follicular cells. Antibody to thyroid constituents can be demonstrated in nearly all cases. Other organ- or tissue-specific autoimmune disorders include pemphigus vulgaris, autoimmune hemolytic anemia, idiopathic thrombocytopenic purpura, Goodpasture's syndrome, myasthenia gravis, and sympathetic ophthalmia. One important feature of some organ-specific autoimmune diseases is the tendency for overlap, such that an individual with one specific syndrome is more likely to develop a second syndrome. For example, there is a high incidence of pernicious anemia in individuals with autoimmune thyroiditis. More striking is the tendency for individuals with an organ-specific autoimmune disease to develop multiple other manifestations of autoimmunity without the development of associated organ pathology. Thus, as many as 50% of individuals with pernicious anemia have non-cross-reacting antibodies to thyroid constituents, whereas patients with myasthenia gravis may develop antinuclear antibodies, antithyroid antibodies, rheumatoid factor, antilymphocyte antibodies, and polyclonal hypergammaglobulinemia. Part of the explanation may relate to the genetic elements shared by individuals with these different diseases.

Systemic autoimmune diseases differ from organ-specific diseases in that pathologic lesions are found in multiple diverse organs and tissues. The hallmark of these conditions is the demonstration of associated relevant autoimmune manifestations that are likely to have an

etiologic role in organ pathology. SLE represents the prototype of these disorders because of its abundant autoimmune manifestations. SLE is a disease of protean manifestations that characteristically involves the kidneys, joints, skin, serosal surfaces, blood vessels, and central nervous system (**Chap. 378**). The disease is associated with a vast array of autoantibodies whose production appears to be a part of a generalized hyperreactivity of the humoral immune system. Other features of SLE include generalized B cell hyperresponsiveness and polyclonal hypergammaglobulinemia. Current evidence suggests that both hypo- and hyperresponsiveness to antigen can lead to survival and activation of autoreactive B cells in SLE. The autoantibodies in SLE are thought to arise as part of an accentuated T cell-dependent B cell response since most pathogenic anti-DNA autoantibodies exhibit evidence of extensive somatic hypermutation.

TREATMENT AUTOIMMUNE DISEASES

Treatment of autoimmune diseases can focus on suppressing the induction of autoimmunity, restoring normal regulatory mechanisms, or inhibiting the effector mechanisms. To decrease the number or function of autoreactive cells, immunosuppressive or ablative therapies are most commonly used. In recent years, cytokine blockade has been demonstrated to be effective in preventing immune activation in some diseases or in inhibiting the extensive inflammatory effector mechanisms characteristic of these diseases. New therapies have also been developed to target lymphoid cells more specifically by blocking a costimulatory signal needed for T or B cell activation, by blocking the migratory capacity of lymphocytes, or by eliminating the effector T cells or B cells. The efficacy of these therapies in some diseases—e.g., SLE (belimumab), rheumatoid arthritis (TNF neutralization, IL-6 receptor blockade, CD28 competition, B cell depletion, IL-1 competition), psoriasis (IL-12/23 depletion, TNF neutralization), and inflammatory bowel disease (TNF neutralization, IL-12 neutralization)—has been demonstrated. One major advance in inhibiting effector mechanisms has been the introduction of cytokine blockade that appears to limit organ damage in some diseases, including rheumatoid arthritis, inflammatory bowel disease, psoriasis, and the spondyloarthritides. Small molecules that block cytokine signaling pathways have recently been introduced into the clinic. Biologicals that interface with T cell activation (CTLA-4Ig) or delete B cells (anti-CD20 antibody) have recently been approved for the treatment of rheumatoid arthritis. Therapies that prevent target-organ damage or support target-organ function remain important in the management of autoimmune disease.

378 Systemic Lupus Erythematosus

Bevra Hannahs Hahn

DEFINITION AND PREVALENCE

Systemic lupus erythematosus (SLE) is an autoimmune disease in which organs and cells undergo damage initially mediated by tissue-binding autoantibodies and immune complexes. In most patients, autoantibodies are present for a few years before the first clinical symptom appears. Ninety percent of patients are women of child-bearing years; people of all genders, ages, and ethnic groups are susceptible. Prevalence of SLE in the United States is 20 to 150 per 100,000 women depending on race and gender; highest prevalence is in African-American and Afro-Caribbean women, and lowest prevalence is in white men.

PATHOGENESIS AND ETIOLOGY

The proposed pathogenic mechanisms of SLE are illustrated in [Fig. 378-1](#). Interactions between susceptibility genes and environmental factors result in abnormal immune responses, which vary between different patients. Those responses may include (1) activation of innate immunity (dendritic cells, monocyte/macrophages) by CpG DNA, DNA in immune complexes, viral DNA or RNA, and RNA in RNA/protein self-antigens; (2) lowered activation thresholds and abnormal activation pathways in adaptive immunity cells (mature T and B lymphocytes); (3) ineffective regulatory CD4+ and CD8+ T cells, B cells, and myeloid-derived suppressor cells; and (4) reduced clearance of immune complexes and apoptotic cells. Self-antigens (nucleosomal DNA/protein; RNA/protein in Sm, Ro, and La; phospholipids) are recognized by the immune system in surface blebs of apoptotic cells; thus autoantigens, autoantibodies, and immune complexes persist for prolonged periods of time, allowing inflammation and disease to develop. Immune cell activation is accompanied by increased secretion of proinflammatory type 1 and 2 interferons (IFNs), tumor necrosis factor α (TNF- α), interleukin (IL) 17 and B cell-maturation/survival cytokines B lymphocyte stimulator (BLyS/BAFF), and IL-10. Upregulation of genes induced by

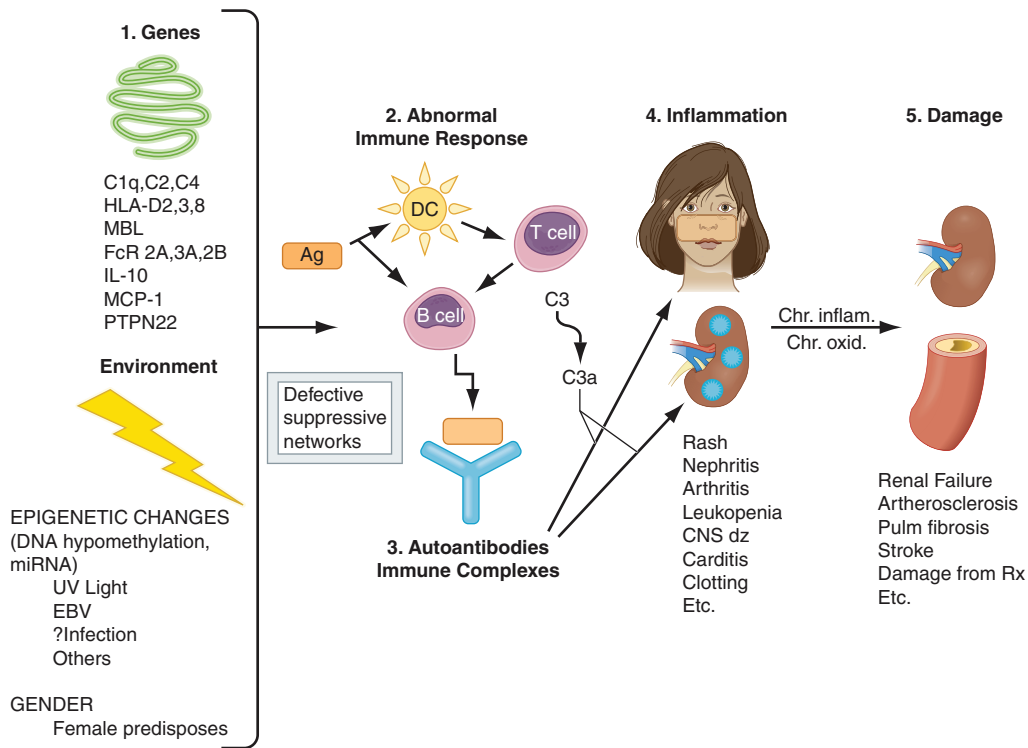


FIGURE 378-1 Pathogenesis of systemic lupus erythematosus (SLE). Genes confirmed in more than one genome-wide association analysis in northern European whites (several confirmed in Asians as well) as increasing susceptibility to SLE or lupus nephritis are listed (reviewed in SG Guerra et al: *Arthritis Res Ther* 14:211, 2012). Gene-environment interactions (reviewed in KH Costenbader et al: *Autoimmune Rev* 11:604, 2012) result in abnormal immune responses that generate pathogenic autoantibodies and immune complexes that deposit in tissue, activate complement, cause inflammation, and over time lead to irreversible organ damage (reviewed in GC Tsokos: *N Engl J Med* 365:2110, 2011; and BH Hahn, in DJ Wallace, BH Hahn [eds]: *Dubois' Lupus Erythematosus and Related Syndromes*, 8th ed. New York, Elsevier, 2013). Ag, antigen; C1q, complement system; C3, complement component; CNS, central nervous system; DC, dendritic cell; EBV, Epstein-Barr virus; HLA, human leukocyte antigen; FcR, immunoglobulin Fc-binding receptor; IL, interleukin; MCP, monocyte chemotactic protein; PTPN, phosphotyrosine phosphatase; UV, ultraviolet.

IFNs is a genetic “signature” in peripheral blood cells of 50–60% of SLE patients. Decreased production of other cytokines also contributes to SLE: lupus T and natural killer (NK) cells fail to produce enough IL-2 and transforming growth factor beta (TGF- β) to induce and sustain regulatory CD4+ and CD8+ T cells. The result of these abnormalities is sustained production of autoantibodies (referred to in Fig. 378-1 and described in Table 378-1) and immune complexes; pathogenic subsets bind target tissues, with activation of complement, leading to release of cytokines, chemokines, vasoactive peptides, oxidants, and proteolytic enzymes. This results in activation of multiple tissue cells (endothelial cells, tissue-fixed macrophages, mesangial cells, podocytes, renal tubular epithelial cells) and influx into target tissues of T and B cells, monocyte/macrophages, and dendritic cells. In the setting of chronic inflammation, accumulation of growth factors and products of chronic oxidation contribute to irreversible tissue damage, including fibrosis/sclerosis in glomeruli, arteries, lungs, and other tissues.

SLE is a multigenic disease. Rare single-gene defects confer high hazard ratios (HRs) for SLE (5 to 25), including homozygous deficiencies of early components of complement (C1q, r, s; C2; C4) and a mutation in *TREX1* on the X chromosome. In most genetically susceptible individuals, normal alleles of multiple genes each contribute a small amount to abnormal immune/inflammation/tissue damage responses; if enough predisposing variations are present, disease results. Approximately 45 predisposing genes (examples listed in Fig. 378-1) have been identified in recent genome-wide association studies in different racial groups. Individually, they confer an HR for SLE of 1.5–3 and account for approximately 18% of disease susceptibility, suggesting that environmental exposures and epigenetics play major roles. Predisposing, antigen-presenting human leukocyte antigen (HLA) molecules are most commonly found, in multiple ethnic groups (HLA

DRB1 *0301 and *1501, as well as multiple genes across the major histocompatibility complex (MHC) 120-gene region). Other genetic factors in whites include innate immunity pathway gene polymorphisms, especially associated with IFN- α (*STAT4*, *IRF5*, *IRAK1*, *TNFAIP3*, *PTPN22*), genes in lymphocyte signaling pathways (*PTPN22*, *PDCD-1*, *Ox40L*, *BANK-1*, *LYN*, *BLK*), genes that affect clearance of apoptotic cells or immune complexes (*C1q*, *FCRGIIA*, *FCRGIII A*, *CRP*, *ITGAM*), genes that influence neutrophil adherence (*ITGAM*), and genes that influence DNA repair (*TREX-1*). Some polymorphisms influence clinical manifestations; such as single nucleotide polymorphisms (SNPs) of *STAT4* that associate with severe disease, anti-DNA, nephritis, and antiphospholipid syndrome, and an allele of *FCGRIIA* encoding a receptor that binds immune complexes poorly and predisposes to nephritis. Some gene effects are in promoter regions (e.g., IL-10), and others are conferred by copy numbers (e.g., C4A). In addition to genome-encoded susceptibility and protective genes, the influence of certain microRNAs (miRNAs) on gene transcription, as well as post-transcriptional epigenetic modification of DNA (which is hypomethylated in T cells of SLE patients), probably play major roles in disease susceptibility.

Some gene polymorphisms contribute to several autoimmune diseases, such as *STAT4* and *CTLA4*. All of these gene polymorphisms/transcription/epigenetic combinations influence immune responses to the external and internal environment; when such responses are too high and/or too prolonged and/or inadequately regulated, autoimmune disease results.

Female sex is permissive for SLE with evidence for hormone effects, genes on the X chromosome, and epigenetic differences between genders playing a role. Females of many mammalian species make higher antibody responses than males. Women exposed to

TABLE 378-1 AUTOANTIBODIES IN SYSTEMIC LUPUS ERYTHEMATOSUS (SLE)

Antibody	Prevalence, %	Antigen Recognized	Clinical Utility
Antinuclear antibodies	98	Multiple nuclear	Best screening test; repeated negative tests make SLE unlikely
Anti-dsDNA	70	DNA (double-stranded)	High titers are SLE-specific and in some patients correlate with disease activity, nephritis, vasculitis
Anti-Sm	25	Protein complexed to 6 species of nuclear U1 RNA	Specific for SLE; no definite clinical correlations; most patients also have anti-RNP; more common in blacks and Asians than whites
Anti-RNP	40	Protein complexed to U1 RNA	Not specific for SLE; high titers associated with syndromes that have overlap features of several rheumatic syndromes including SLE; more common in blacks than whites
Anti-Ro (SS-A)	30	Protein complexed to hY RNA, primarily 60 kDa and 52 kDa	Not specific for SLE; associated with sicca syndrome, predisposes to subacute cutaneous lupus, and to neonatal lupus with congenital heart block; associated with decreased risk for nephritis
Anti-La (SS-B)	10	47-kDa protein complexed to hY RNA	Usually associated with anti-Ro; associated with decreased risk for nephritis
Antihistone	70	Histones associated with DNA (in nucleosome, chromatin)	More frequent in drug-induced lupus than in SLE
Antiphospholipid	50	Phospholipids, β_2 glycoprotein 1 (β_2 G1) cofactor, prothrombin	Three tests available—ELISAs for cardiolipin and β_2 G1, sensitive prothrombin time (DRVVT); predisposes to clotting, fetal loss, thrombocytopenia
Antierthrocyte	60	Erythrocyte membrane	Measured as direct Coombs test; a small proportion develops overt hemolysis
Antiplatelet	30	Surface and altered cytoplasmic antigens on platelets	Associated with thrombocytopenia, but sensitivity and specificity are not good; this is not a useful clinical test
Antineuronal (includes antiglutamate receptor)	60	Neuronal and lymphocyte surface antigens	In some series, a positive test in CSF correlates with active CNS lupus
Antiribosomal P	20	Protein in ribosomes	In some series, a positive test in serum correlates with depression or psychosis due to CNS lupus

Abbreviations: CNS, central nervous system; CSF, cerebrospinal fluid; DRVVT, dilute Russell viper venom time; ELISA, enzyme-linked immunosorbent assay.

estrogen-containing oral contraceptives or hormone replacement have an increased risk of developing SLE (1.2- to 2-fold). Estradiol binds to receptors on T and B lymphocytes, increasing activation and survival of those cells, thus favoring prolonged immune responses. Genes on the X chromosome that influence SLE, such as *TREX-1*, may play a role in gender predisposition, possibly because some genes on the second X in females are not silent. People with XXY karyotype (Klinefelter's syndrome) have a significantly increased risk for SLE.

Several environmental stimuli may influence SLE (Fig. 378-1). Exposure to ultraviolet light causes flares of SLE in approximately 70% of patients, possibly by increasing apoptosis in skin cells or by altering DNA and intracellular proteins to make them antigenic. Some infections induce normal immune responses that involve certain T and B cells that recognize self-antigens; such cells are not appropriately regulated, and autoantibody production occurs. Most SLE patients have autoantibodies for 3 years or more before the first symptoms of disease, suggesting that regulation controls the degree of autoimmunity for years before quantities and qualities of autoantibodies and pathogenic B and T cells cause clinical disease. Epstein-Barr virus (EBV) may be one infectious agent that can trigger SLE in susceptible individuals. Children and adults with SLE are more likely to be infected by EBV than age-, sex-, and ethnicity-matched controls. EBV contains amino acid sequences that mimic sequences on human spliceosomes (RNA/protein antigens) often recognized by autoantibodies in people with SLE. Current tobacco smoking increases risk for SLE (odds ratio [OR] 1.5). Prolonged occupational exposure to silica (e.g., inhalation of soap powder dust or soil in farming activities) increases risk (OR 4.3) in African-American women. Thus, interplay between genetic susceptibility, environment, gender, and abnormal immune responses results in autoimmunity (Chap. 377e).

PATHOLOGY

In SLE, biopsies of affected skin show deposition of Ig at the dermal-epidermal junction (DEJ), injury to basal keratinocytes, and inflammation dominated by T lymphocytes in the DEJ and around blood vessels and dermal appendages. Clinically unaffected skin may also show Ig deposition at the DEJ.

In renal biopsies, the pattern and severity of injury are important in diagnosis and in selecting the best therapy. Most recent clinical studies of

lupus nephritis have used the International Society of Nephrology (ISN) and the Renal Pathology Society (RPS) classification (Table 378-2). In the ISN/RPS classification, the addition of “a” for active and “c” for chronic changes gives physicians information regarding the potential reversibility of disease. The system focuses on glomerular disease, although the presence of tubular interstitial and vascular disease is important to clinical outcomes. In general, class III and IV disease, as well as class V accompanied by III or IV disease, should be treated with aggressive immunosuppression if possible, because there is a high risk for end-stage renal disease (ESRD) if patients are untreated or undertreated. In contrast, treatment for lupus nephritis is not recommended in patients with class I or II disease or with extensive irreversible changes. In the recent Systemic Lupus International Collaborating Clinic (SLICC) criteria for classification of SLE, a diagnosis can be established on the basis of renal histology without meeting additional criteria (Table 378-3).

Histologic abnormalities in blood vessels may also determine therapy. Patterns of vasculitis are not specific for SLE but may indicate active disease: leukocytoclastic vasculitis is most common (Chap. 385).

Lymph node biopsies are usually performed to rule out infection or malignancies. In SLE, they show nonspecific diffuse chronic inflammation.

DIAGNOSIS

The diagnosis of SLE is based on characteristic clinical features and autoantibodies. Current criteria for classification are listed in Table 378-3, and an algorithm for diagnosis and initial therapy is shown in Fig. 378-2. The criteria are intended for confirming the diagnosis of SLE in patients included in studies; the author uses them in individual patients for estimating the probability that a disease is SLE. Any combination of four or more criteria, with at least one in the clinical and one in the immunologic category, well documented at any time during an individual's history, makes it likely that the patient has SLE. (Specificity and sensitivity are ~93% and ~92%, respectively.) In many patients, criteria accrue over time. Antinuclear antibodies (ANA) are positive in >98% of patients during the course of disease; repeated negative tests by immunofluorescent methods suggest that the diagnosis is not SLE, unless other autoantibodies are present (Fig. 378-2). High-titer IgG antibodies to double-stranded DNA and antibodies to the Sm antigen are both specific for SLE and, therefore, favor the diagnosis in the presence of compatible clinical manifestations. The

TABLE 378-2 CLASSIFICATION OF LUPUS NEPHRITIS (INTERNATIONAL SOCIETY OF NEPHROLOGY AND RENAL PATHOLOGY SOCIETY)**Class I: Minimal Mesangial Lupus Nephritis**

Normal glomeruli by light microscopy, but mesangial immune deposits by immunofluorescence.

Class II: Mesangial Proliferative Lupus Nephritis

Purely mesangial hypercellularity of any degree or mesangial matrix expansion by light microscopy, with mesangial immune deposits. A few isolated subepithelial or subendothelial deposits may be visible by immunofluorescence or electron microscopy, but not by light microscopy.

Class III: Focal Lupus Nephritis

Active or inactive focal, segmental or global endo- or extracapillary glomerulonephritis involving $\leq 50\%$ of all glomeruli, typically with focal subendothelial immune deposits, with or without mesangial alterations.

Class III (A): Active lesions—focal proliferative lupus nephritis

Class III (A/C): Active and chronic lesions—focal proliferative and sclerosing lupus nephritis

Class III (C): Chronic inactive lesions with glomerular scars—focal sclerosing lupus nephritis

Class IV: Diffuse Lupus Nephritis

Active or inactive diffuse, segmental or global endo- or extracapillary glomerulonephritis involving $\geq 50\%$ of all glomeruli, typically with diffuse subendothelial immune deposits, with or without mesangial alterations. This class is divided into diffuse segmental (IV-S) lupus nephritis when $\geq 50\%$ of the involved glomeruli have segmental lesions, and diffuse global (IV-G) lupus nephritis when $\geq 50\%$ of the involved glomeruli have global lesions. Segmental is defined as a glomerular lesion that involves less than one-half of the glomerular tuft. This class includes cases with diffuse wire loop deposits but with little or no glomerular proliferation.

Class IV-S (A): Active lesions—diffuse segmental proliferative lupus nephritis

Class IV-G (A): Active lesions—diffuse global proliferative lupus nephritis

Class IV-S (A/C): Active and chronic lesions—diffuse segmental proliferative and sclerosing lupus nephritis

Class IV-G (A/C): Active and chronic lesions—diffuse global proliferative and sclerosing lupus nephritis

Class IV-S (C): Chronic inactive lesions with scars—diffuse segmental sclerosing lupus nephritis

Class IV-G (C): Chronic inactive lesions with scars—diffuse global sclerosing lupus nephritis

Class V: Membranous Lupus Nephritis

Global or segmental subepithelial immune deposits or their morphologic sequelae by light microscopy and by immunofluorescence or electron microscopy, with or without mesangial alterations. Class V lupus nephritis may occur in combination with class III or IV, in which case both will be diagnosed. Class V lupus nephritis may show advanced sclerosis.

Class VI: Advanced Sclerotic Lupus Nephritis

$\geq 90\%$ of glomeruli globally sclerosed without residual activity.

Note: Indicate and grade (mild, moderate, severe) tubular atrophy, interstitial inflammation and fibrosis, and severity of arteriosclerosis or other vascular lesions.

Source: JJ Weening et al: *Kidney Int* 65:521, 2004. Reprinted by permission from Macmillan Publishers Ltd., Copyright 2004.

presence in an individual of multiple autoantibodies without clinical symptoms should not be considered diagnostic for SLE, although such persons are at increased risk.

INTERPRETATION OF CLINICAL MANIFESTATIONS

When a diagnosis of SLE is made, it is important to establish the severity and potential reversibility of the illness and to estimate the possible consequences of various therapeutic interventions. In the following paragraphs, descriptions of some disease manifestations begin with relatively mild problems and progress to those more life-threatening.

OVERVIEW AND SYSTEMIC MANIFESTATIONS

At its onset, SLE may involve one or several organ systems; over time, additional manifestations may occur (Tables 378-3 and 378-4). Most of the autoantibodies characteristic of each person are present at the time clinical manifestations appear (Tables 378-1 and 378-3). Severity

TABLE 378-3 SYSTEMIC LUPUS INTERNATIONAL COLLABORATING CLINIC CRITERIA FOR CLASSIFICATION OF SYSTEMIC LUPUS ERYTHEMATOSUS

Clinical Manifestations	Immunologic Manifestations
Skin	ANA > reference negative value
Acute, subacute cutaneous LE	Anti-dsDNA
Chronic cutaneous LE	Anti-Sm
Oral ulcers	Antiphospholipid
Alopecia	Low serum complement
Synovitis	Positive direct Coombs test
Renal	
Prot/Cr ≥ 0.5	
RBC casts	
Biopsy ^a	
Neurologic	
Seizures, psychosis, mononeuritis, myelitis, peripheral or cranial neuropathies, acute confusional state	
Hemolytic anemia	
Leukopenia (<4000) or	
Lymphopenia (<1000)	
Thrombocytopenia (<100,000)	

^aRenal biopsy read as systemic lupus qualifies for classification as SLE even if none of the other above features are present.

Interpretation: Presence of any 4 criteria (must have at least 1 in each category) qualifies patient to be classified as having SLE with 93% specificity and 92% sensitivity.

Abbreviations: ANA, antinuclear antibody; Cr, creatinine; LE, lupus erythematosus; Prot, protein.

Source: M Petri et al: *Arthritis Rheum* 64:2677, 2012. Because these criteria are new, currently ongoing clinical studies use prior American College of Rheumatology Criteria; see EM Tan et al: *Arthritis Rheum* 25:1271, 1982; update MC Hochberg: *Arthritis Rheum* 40:1725, 1997.

of SLE varies from mild and intermittent to severe and fulminant. Approximately 85% of patients have either continuing active disease (while being treated) or one or more flares of active disease annually. Permanent complete remissions (absence of symptoms with no treatment) are rare. Systemic symptoms, particularly fatigue and myalgias/arthralgias, are present most of the time. Severe systemic illness requiring glucocorticoid therapy can occur with fever, prostration, weight loss, and anemia with or without other organ-targeted manifestations.

MUSCULOSKELETAL MANIFESTATIONS

Most people with SLE have intermittent polyarthritis, varying from mild to disabling, characterized by soft tissue swelling and tenderness in joints and/or tendons, most commonly in hands, wrists, and knees. Joint deformities (hands and feet) develop in only 10%. Erosions on joint x-rays are rare but can be identified by ultrasound in almost half of patients. Some individuals have rheumatoid-like arthritis with erosions and fulfill criteria for both RA and SLE (“rhupus”); they may be coded as having both diseases. If pain persists in a single joint, such as knee, shoulder, or hip, a diagnosis of ischemic necrosis of bone should be considered, particularly if there are no other manifestations of active SLE because its prevalence is increased in SLE, especially in patients treated with systemic glucocorticoids. Myositis with clinical muscle weakness, elevated creatine kinase levels, positive magnetic resonance imaging (MRI) scan, and muscle necrosis and inflammation on biopsy can occur, although most patients have myalgias without frank myositis. Glucocorticoid therapies (commonly) and antimalarial therapies (rarely) can cause muscle weakness; these adverse effects must be distinguished from active inflammatory disease.

CUTANEOUS MANIFESTATIONS

Lupus dermatitis can be classified as acute, subacute, or chronic, and there are many different types of lesions encompassed within these groups. Discoid lupus erythematosus (DLE) is the most common

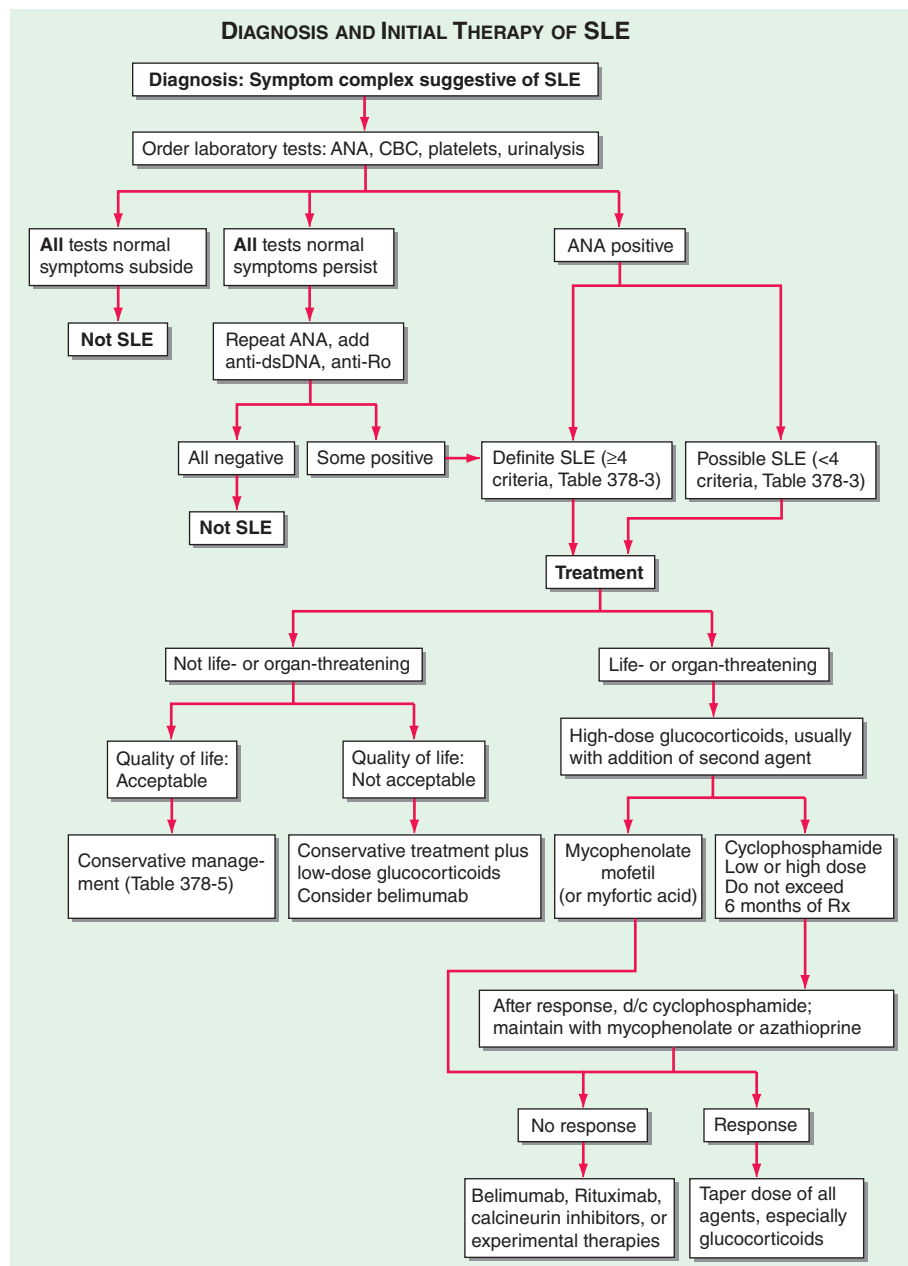


FIGURE 378-2 Algorithm for diagnosis and initial therapy of systemic lupus erythematosus (SLE). For guidelines on management of lupus and lupus nephritis, see BH Hahn et al: *Arthritis Care Res (Hoboken)* 64:797, 2012; GK Bertsias et al: *Ann Rheum Dis* 71:1771, 2012; and G Bertsias et al: *Ann Rheum Dis* 67:195, 2008. For details on mycophenolate and cyclophosphamide induction and maintenance therapies, see L Henderson et al: *Cochrane Database Syst Rev* 12:CD002922, 2012; Z Touma et al: *J Rheumatol* 38:69, 2011; EM Ginzler et al: *Arthritis Rheum* 62:211, 2010; FA Houssiau et al: *Ann Rheum Dis* 69:61, 2010; and MA Dooley et al: *N Engl J Med* 365:1886, 2011. For belimumab in treatment, see BH Hahn: *N Eng J Med* 368:1528, 2013. For rituximab, see L Lightstone: *Lupus* 22:390, 2013; and BH Rovin et al: *Arthritis Rheum* 64:1215, 2012. ANA, antinuclear antibodies; CBC, complete blood count.

chronic dermatitis in lupus; lesions are roughly circular with slightly raised, scaly hyperpigmented erythematous rims and depigmented, atrophic centers in which all dermal appendages are permanently destroyed. Lesions can be disfiguring, particularly on the face and scalp. Treatment consists primarily of topical or locally injected glucocorticoids and systemic antimalarials. Only 5% of people with DLE have SLE (although half have positive ANA); however, among individuals with SLE, as many as 20% have DLE. The most common acute SLE rash is a photosensitive, slightly raised erythema, occasionally scaly, on the face (particularly the cheeks and nose—the “butterfly” rash), ears, chin, V region of the neck and chest, upper back, and

extensor surfaces of the arms. Worsening of this rash often accompanies flare of systemic disease. Subacute cutaneous lupus erythematosus (SCLE) consists of scaly red patches similar to psoriasis, or circular flat red-rimmed lesions. Patients with these manifestations are exquisitely photosensitive; most have antibodies to Ro (SS-A). Other SLE rashes include recurring urticaria, lichen planus-like dermatitis, bullae, and panniculitis (“lupus profundus”). Rashes can be minor or severe; they may be the major disease manifestation. Small ulcerations on the oral or nasal mucosa are common in SLE; the lesions resemble aphthous ulcers.

RENAL MANIFESTATIONS

Nephritis is usually the most serious manifestation of SLE, particularly because nephritis and infection are the leading causes of mortality in the first decade of disease. Because nephritis is asymptomatic in most lupus patients, urinalysis should be ordered in any person suspected of having SLE. The classification of lupus nephritis is primarily histologic (see “Pathology,” above, and Table 378-2). Renal biopsy is recommended for every SLE patient with any clinical evidence of nephritis; results are used to plan current and near-future therapies. Patients with dangerous proliferative forms of glomerular damage (ISN III and IV) usually have microscopic hematuria and proteinuria (>500 mg per 24 h); approximately one-half develop nephrotic syndrome, and most develop hypertension. If diffuse proliferative glomerulonephritis (DPGN) is inadequately treated, virtually all patients develop ESRD within 2 years of diagnosis. Therefore, aggressive immunosuppression is indicated (usually systemic glucocorticoids plus a cytotoxic drug), unless damage is irreversible (Fig. 378-2, Table 378-5). African Americans are more likely to develop ESRD than are whites, even with the most current therapies. Overall in the United States, ~20% of individuals with lupus DPGN die or develop ESRD within 10 years of diagnosis. Such individuals require aggressive control of SLE and of the complications of renal disease and of therapy. Approximately 20% of SLE patients with proteinuria (usually nephrotic) have membranous glomerular changes without proliferative changes on renal biopsy. Their outcome is better than for those with DPGN, but patients with class V and nephrotic range proteinuria should be treated in the same way as those with classes III or IV proliferative disease. Lupus nephritis tends to be an ongoing disease, with flares requiring re-treatment or increased treatment over many years. For most people with lupus nephritis, accelerated atherosclerosis becomes important after several years of disease; attention must be given to control of systemic inflammation, blood pressure, hyperlipidemia, and hyperglycemia.

NERVOUS SYSTEM MANIFESTATIONS

There are many central nervous system (CNS) and peripheral nervous system manifestations of SLE; in some patients, these are the major cause of morbidity and mortality. It is useful to approach this diagnostically by asking first whether the symptoms result from SLE or another condition (such as infection in immunosuppressed

TABLE 378-4 CLINICAL MANIFESTATIONS OF SLE AND PREVALENCE OVER THE ENTIRE COURSE OF DISEASE^a

Manifestation	Prevalence, %
Systemic: Fatigue, malaise, fever, anorexia, weight loss	95
Musculoskeletal	95
Arthralgias/myalgias	95
Nonerosive polyarthritis	60
Hand deformities	10
Myopathy/myositis	25/5
Ischemic necrosis of bone	15
Cutaneous	80
Photosensitivity	70
Malar rash	50
Oral ulcers	40
Alopecia	40
Discoid rash	20
Vasculitis rash	20
Other (e.g., urticaria, subacute cutaneous lupus)	15
Hematologic	85
Anemia (chronic disease)	70
Leukopenia (<4000/ μ L)	65
Lymphopenia (<1500/ μ L)	50
Thrombocytopenia (<100,000/ μ L)	15
Lymphadenopathy	15
Splenomegaly	15
Hemolytic anemia	10
Neurologic	60
Cognitive disorder	50
Mood disorder	40
Headache	25
Seizures	20
Mono-, polyneuropathy	15
Stroke, TIA	10
Acute confusional state or movement disorder	2–5
Aseptic meningitis, myelopathy	<1
Cardiopulmonary	60
Pleurisy, pericarditis, effusions	30–50
Myocarditis, endocarditis	10
Lupus pneumonitis	10
Coronary artery disease	10
Interstitial fibrosis	5
Pulmonary hypertension, ARDS, hemorrhage	<5
Shrinking lung syndrome	<5
Renal	30–50
Proteinuria \geq 500 mg/24 h, cellular casts	30–50
Nephrotic syndrome	25
End-stage renal disease	5–10
Gastrointestinal	40
Nonspecific (nausea, mild pain, diarrhea)	30
Abnormal liver enzymes	40
Vasculitis	5
Thrombosis	15
Venous	10
Arterial	5
Ocular	15
Sicca syndrome	15
Conjunctivitis, episcleritis	10
Vasculitis	5

^aNumbers indicate percentage of patients who have the manifestation at some time during the course of illness.

Abbreviations: ARDS, acute respiratory distress syndrome; TIA, transient ischemic attack.

individuals or side effects of therapies). If symptoms are related to SLE, it should be determined whether they are caused by a diffuse process (requiring immunosuppression) or vascular occlusive disease (requiring anticoagulation). The most common manifestation of diffuse CNS lupus is cognitive dysfunction, including difficulties with memory and reasoning. Headaches are also common. When exacerbating, they often indicate SLE flare; when milder, they are difficult to distinguish from migraine or tension headaches. Seizures of any type may be caused by lupus; treatment often requires both antiseizure and immunosuppressive therapies. Psychosis can be the dominant manifestation of SLE; it must be distinguished from glucocorticoid-induced psychosis. The latter usually occurs in the first weeks of glucocorticoid therapy, at daily doses of \geq 40 mg of prednisone or equivalent; psychosis resolves over several days after glucocorticoids are decreased or stopped. Myelopathy is not rare and is often disabling; rapid initiation of immunosuppressive therapy starting with high-dose glucocorticoids is standard of care.

VASCULAR OCCLUSIONS

The prevalence of transient ischemic attacks, strokes, and myocardial infarctions is increased in patients with SLE. These vascular events are increased in, but not exclusive to, SLE patients with antibodies to phospholipids (antiphospholipid antibodies), which are associated with hypercoagulability and acute thrombotic events (Chap. 379). Chronic SLE with or without antiphospholipid antibodies is associated with accelerated atherosclerosis. Ischemia in the brain can be caused by focal occlusion (either noninflammatory or associated with vasculitis) or by embolization from carotid artery plaque or from fibrinous vegetations of Libman-Sacks endocarditis. Appropriate tests for antiphospholipid antibodies (see below) and for sources of emboli should be ordered in such patients to estimate the need for, intensity of, and duration of anti-inflammatory and/or anticoagulant therapies. In SLE, myocardial infarctions are primarily manifestations of accelerated atherosclerosis. The increased risk for vascular events is three- to tenfold overall, and is highest in women <49 years old. Characteristics associated with increased risk for atherosclerosis include older age, hypertension, dyslipidemia, dysfunctional proinflammatory high-density lipoproteins, repeated high scores for disease activity, high cumulative or daily doses of glucocorticoids, and high levels of homocysteine. When it is most likely that an event results from clotting, long-term anticoagulation is the therapy of choice. Two processes can occur at once—vasculitis plus bland vascular occlusions—in which case it is appropriate to treat with anticoagulation plus immunosuppression. Statin therapies reduce levels of low-density lipoproteins (LDL) in SLE patients; reduction of cardiac events by statins has been shown in SLE patients with renal transplants but not in other SLE cohorts to date.

PULMONARY MANIFESTATIONS

The most common pulmonary manifestation of SLE is pleuritis with or without pleural effusion. This manifestation, when mild, may respond to treatment with nonsteroidal anti-inflammatory drugs (NSAIDs); when more severe, patients require a brief course of glucocorticoid therapy. Pulmonary infiltrates also occur as a manifestation of active SLE and are difficult to distinguish from infection on imaging studies. Life-threatening pulmonary manifestations include interstitial inflammation leading to fibrosis, shrinking lung syndrome, and intraalveolar hemorrhage; all of these probably require early aggressive immunosuppressive therapy as well as supportive care.

CARDIAC MANIFESTATIONS

Pericarditis is the most frequent cardiac manifestation; it usually responds to anti-inflammatory therapy and infrequently leads to tamponade. More serious cardiac manifestations are myocarditis and fibrinous endocarditis of Libman-Sacks. The endocardial involvement can lead to valvular insufficiencies, most commonly of the mitral or aortic valves, or to embolic events. It has not been proven that glucocorticoid or other immunosuppressive therapies lead to improvement of lupus myocarditis or endocarditis, but it is usual practice to administer a trial of high-dose steroids along with appropriate supportive therapy for

TABLE 378-5 MEDICATIONS FOR THE MANAGEMENT OF SLE

Medication	Dose Range	Drug Interactions	Serious or Common Adverse Effects
NSAIDs, salicylates (Ecotrin ^a and St. Joseph's aspirin ^a approved by FDA for use in SLE)	Doses toward upper limit of recommended range usually required	A2R/ACE inhibitors, glucocorticoids, fluconazole, methotrexate, thiazides	NSAIDs: Higher incidence of aseptic meningitis, elevated liver enzymes, decreased renal function, vasculitis of skin; entire class, especially COX-2-specific inhibitors, may increase risk for myocardial infarction Salicylates: ototoxicity, tinnitus Both: GI events and symptoms, allergic reactions, dermatitis, dizziness, acute renal failure, edema, hypertension
Topical glucocorticoids	Mid potency for face; mid to high potency for other areas	None known	Atrophy of skin, contact dermatitis, folliculitis, hypopigmentation, infection
Topical sunscreens	SPF 15 at least; 30+ preferred	None known	Contact dermatitis
Hydroxychloroquine ^a (quinacrine can be added or substituted)	200–400 mg qd (100 mg qd)	None known	Retinal damage, agranulocytosis, aplastic anemia, ataxia, cardiomyopathy, dizziness, myopathy, ototoxicity, peripheral neuropathy, pigmentation of skin, seizures, thrombocytopenia. Quinacrine usually causes diffuse yellow skin coloration.
DHEA (dehydroepiandrosterone)	200 mg qd	Unclear	Acne, menstrual irregularities, high serum levels of testosterone
Methotrexate (for dermatitis, arthritis)	10–25 mg once a week, PO or SC, with folic acid; decrease dose if CrCl <60 mL/min	Acitretin, leflunomide, NSAIDs and salicylates, penicillins, probenecid, sulfonamides, trimethoprim	Anemia, bone marrow suppression, leukopenia, thrombocytopenia, hepatotoxicity, nephrotoxicity, infections, neurotoxicity, pulmonary fibrosis, pneumonitis, severe dermatitis, seizures.
Glucocorticoids, oral ^a (several specific brands are approved by FDA for use in SLE)	Prednisone, prednisolone: 0.5–1 mg/kg per day for severe SLE 0.07–0.3 mg/kg per day or qod for milder disease	A2R/ACE antagonists, antiarrhythmics class III, cyclosporine, NSAIDs and salicylates, phenothiazines, phenytoins, quinolones, rifampin, risperidone, thiazides, sulfonamides, warfarin	Infection, VZV infection, hypertension, hyperglycemia, hypokalemia, acne, allergic reactions, anxiety, aseptic necrosis of bone, cushingoid changes, CHF, fragile skin, insomnia, menstrual irregularities, mood swings, osteoporosis, psychosis
Methylprednisolone sodium succinate, IV ^a (FDA approved for lupus nephritis)	For severe disease, 1 g IV qd × 3 days	As for oral glucocorticoids	As for oral glucocorticoids (if used repeatedly); anaphylaxis
Cyclophosphamide ^b IV	Low dose (for whites of northern European backgrounds): 500 mg every 2 weeks for 6 doses, then begin maintenance with MMF or AZA. High dose: 7–25 mg/kg q month × 6; consider mesna administration with dose	Allopurinol, bone marrow suppressants, colony-stimulating factors, doxorubicin, rituximab, succinylcholine, zidovudine	Infection, VZV infection, bone marrow suppression, leukopenia, anemia, thrombocytopenia, hemorrhagic cystitis (less with IV), carcinoma of the bladder, alopecia, nausea, diarrhea, malaise, malignancy, ovarian and testicular failure. Ovarian failure is probably not a problem with low dose.
Oral	1.5–3 mg/kg per day; decrease dose for CrCl <25 mL/min		
Mycophenolate mofetil (MMF) ^b or mycophenolic acid (MPA)	MMF: 2–3 g/d PO for induction therapy, 1–2 g/d for maintenance therapy; max 1 g bid if CrCl <25 mL/min MPA: 360–1080 mg bid; caution if CrCl <25 mL/min	Acyclovir, antacids, azathioprine, bile acid-binding resins, ganciclovir, iron, salts, probenecid, oral contraceptives	Infection, leukopenia, anemia, thrombocytopenia, lymphoma, lymphoproliferative disorders, malignancy, alopecia, cough, diarrhea, fever, GI symptoms, headache, hypertension, hypercholesterolemia, hypokalemia, insomnia, peripheral edema, elevated liver enzymes, tremor, rash
Azathioprine (AZA) ^b	2–3 mg/kg per day PO for induction; 1–2 mg/kg per day for maintenance; decrease frequency of dose if CrCl <50 mL/min	ACE inhibitors, allopurinol, bone marrow suppressants, interferons, mycophenolate mofetil, rituximab, warfarin, zidovudine	Infection, VZV infection, bone marrow suppression, leukopenia, anemia, thrombocytopenia, pancreatitis, hepatotoxicity, malignancy, alopecia, fever, flulike illness, GI symptoms
Belimumab	10 mg/kg IV wks 0, 2, and 4, then monthly	IVIg	Infusion reactions, allergy, infections probable
Rituximab (for patients resistant to above therapies)	375 mg/m ² q wk × 4 or 1 g q 2 wks × 2	IVIg	Infection (including PML), infusion reactions, headache, arrhythmias, allergic responses

^aIndicates medication is approved for use in SLE by the U.S. Food and Drug Administration. ^bIndicates the medication has been used with glucocorticoids in the trials showing efficacy.

Abbreviations: A2R, angiotensin II receptor; ACE, angiotensin-converting enzyme; CHF, congestive heart failure; CrCl, creatinine clearance; FDA, U.S. Food and Drug Administration; GI, gastrointestinal; IVIg, intravenous immunoglobulin; NSAIDs, nonsteroidal anti-inflammatory drugs; PML, progressive multifocal leukoencephalopathy; SLE, systemic lupus erythematosus; SPF, sun protection factor; VZV, varicella-zoster virus.

heart failure, arrhythmia, or embolic events. As discussed above, patients with SLE are at increased risk for myocardial infarction, usually due to accelerated atherosclerosis, which probably results from immune attack, chronic inflammation, and/or chronic oxidative damage to arteries.

HEMATOLOGIC MANIFESTATIONS

The most frequent hematologic manifestation of SLE is anemia, usually normochromic normocytic, reflecting chronic illness. Hemolysis can be rapid in onset and severe, requiring high-dose glucocorticoid therapy, which is effective in most patients. Leukopenia is also common and almost always consists of lymphopenia, not granulocytopenia; lymphopenia rarely predisposes to infections and by itself usually does not require therapy. Thrombocytopenia may be a recurring problem. If platelet counts are $>40,000/\mu\text{L}$ and abnormal bleeding is absent, therapy may not be required. High-dose glucocorticoid therapy (e.g., 1 mg/kg per day of prednisone or equivalent) is usually effective for the first few episodes of severe thrombocytopenia. Recurring or prolonged hemolytic anemia or thrombocytopenia, or disease requiring an unacceptably high dose of daily glucocorticoids, should be treated with an additional strategy (see “Management of Systemic Lupus Erythematosus” below).

GASTROINTESTINAL MANIFESTATIONS

Nausea, sometimes with vomiting, and diarrhea can be manifestations of an SLE flare, as can diffuse abdominal pain probably caused by autoimmune peritonitis and/or intestinal vasculitis. Increases in serum aspartate aminotransferase (AST) and alanine aminotransferase (ALT) are common when SLE is active. These manifestations usually improve promptly during systemic glucocorticoid therapy. Vasculitis involving the intestine may be life-threatening; perforations, ischemia, bleeding, and sepsis are frequent complications. Aggressive immunosuppressive therapy with high-dose glucocorticoids is recommended for short-term control; evidence of recurrence is an indication for additional therapies.

OCULAR MANIFESTATIONS

Sicca syndrome (Sjögren’s syndrome; [Chap. 383](#)) and nonspecific conjunctivitis are common in SLE and rarely threaten vision. In contrast, retinal vasculitis and optic neuritis are serious manifestations: blindness can develop over days to weeks. Aggressive immunosuppression is recommended, although there are no controlled trials to prove effectiveness. Complications of systemic and intraorbital glucocorticoid therapy include cataracts (common) and glaucoma.

LABORATORY TESTS

Laboratory tests serve (1) to establish or rule out the diagnosis; (2) to follow the course of disease, particularly to suggest that a flare is occurring or organ damage is developing; and (3) to identify adverse effects of therapies.

TESTS FOR AUTOANTIBODIES (TABLES 378-1 AND 378-3)

Diagnostically, the most important autoantibodies to detect are ANA because the test is positive in $>95\%$ of patients, usually at the onset of symptoms. A few patients develop ANA within 1 year of symptom onset; repeated testing may thus be useful. ANA tests using immunofluorescent methods are more reliable than enzyme-linked immunosorbent assays (ELISAs) and/or bead assays, which have less specificity. ANA-negative lupus exists but is rare in adults and is usually associated with other autoantibodies (anti-Ro or anti-DNA). High-titer IgG antibodies to double-stranded DNA (dsDNA) (but not to single-stranded DNA) are specific for SLE. ELISA and immunofluorescent reactions of sera with the dsDNA in the flagellate *Crithidia luciliae* have $\sim 60\%$ sensitivity for SLE; identification of high-avidity anti-dsDNA in the Farr assay is not as sensitive but may correlate better with risk for nephritis. Titers of anti-dsDNA vary over time. In some patients, increases in quantities of anti-dsDNA herald a flare, particularly of nephritis or vasculitis, especially when associated with declining levels of C3 or C4 complement. Antibodies to Sm are also specific for SLE and assist in diagnosis; anti-Sm antibodies do not usually correlate with disease activity or clinical manifestations.

Antiphospholipid antibodies are not specific for SLE, but their presence fulfills one classification criterion, and they identify patients at increased risk for venous or arterial clotting, thrombocytopenia, and fetal loss. There are three widely accepted tests that measure different antibodies (anticardiolipin, anti- β_2 -glycoprotein, and the lupus anticoagulant). ELISA is used for anticardiolipin and anti- β_2 -glycoprotein (both internationally standardized with good reproducibility); a sensitive phospholipid-based activated prothrombin time such as the dilute Russell venom viper test is used to identify the lupus anticoagulant. The higher the titers of IgG anticardiolipin (>40 IU is considered high), and the greater the number of different antiphospholipid antibodies that are detected, the greater is the risk for a clinical episode of clotting. Quantities of antiphospholipid antibodies may vary markedly over time; repeated testing is justified if clinical manifestations of the antiphospholipid syndrome (APS) appear ([Chap. 379](#)). To classify a patient as having APS, with or without SLE, by international criteria requires the presence of one or more clotting episodes and/or repeated fetal losses plus at least two positive tests for antiphospholipid antibodies, at least 12 weeks apart; however, many patients with APS do not meet these stringent criteria, which are intended for inclusion of patients into studies.

An additional autoantibody test with predictive value (not used for diagnosis) detects anti-Ro/SS-A, which indicates increased risk for neonatal lupus, sicca syndrome, and SCLE. Women with child-bearing potential and SLE should be screened for antiphospholipid antibodies and anti-Ro, because both antibodies have the potential to cause fetal harm.

STANDARD TESTS FOR DIAGNOSIS

Screening tests for complete blood count, platelet count, and urinalysis may detect abnormalities that contribute to the diagnosis and influence management decisions.

TESTS FOR FOLLOWING DISEASE COURSE

It is useful to follow tests that indicate the status of organ involvement known to be present during SLE flares. These might include urinalysis for hematuria and proteinuria, hemoglobin levels, platelet counts, and serum levels of creatinine or albumin. There is great interest in identification of additional markers of disease activity. Candidates include levels of anti-DNA and anti-C1q antibodies, several components of complement (C3 is most widely available), activated complement products (including those that bind to the C4d receptor on erythrocytes), IFN-inducible gene expression in peripheral blood cells, serum levels of BLYS (B lymphocyte stimulator, also called BAFF), and urinary levels of TNF-like weak inducer of apoptosis (TWEAK), neutrophil gelatinase-associated lipocalin (NGAL), or monocyte chemoattractant protein 1 (MCP-1). None is uniformly agreed upon as a reliable indicator of flare or of response to therapeutic interventions. It is likely that a panel of multiple proteins will be developed to predict both impending flare and response to recently instituted therapies. For now, the physician should determine for each patient whether certain laboratory test changes predict flare. If so, altering therapy in response to these changes may be advisable (30 mg of prednisone daily for 2 weeks has been shown to prevent flares in patients with rising anti-DNA plus falling complement). In addition, given the increased prevalence of atherosclerosis in SLE, it is advisable to follow the recommendations of the National Cholesterol Education Program for testing and treatment, including scoring of SLE as an independent risk factor, similar to diabetes mellitus.

MANAGEMENT OF SYSTEMIC LUPUS ERYTHEMATOSUS

There is no cure for SLE, and complete sustained remissions are rare. Therefore, the physician should plan to induce remissions of acute flares and then maintain improvements with strategies that suppress symptoms to an acceptable level and prevent organ damage. Usually patients will endure some adverse effects of medications. Therapeutic choices depend on (1) whether disease manifestations are life-threatening or likely to cause organ damage, justifying aggressive therapies; (2) whether

2132 manifestations are potentially reversible; and (3) the best approaches to preventing complications of disease and its treatments. Therapies, doses, and adverse effects are listed in Table 378-5.

CONSERVATIVE THERAPIES FOR MANAGEMENT OF NON-LIFE-THREATENING DISEASE

Among patients with fatigue, pain, and autoantibodies indicative of SLE, but without major organ involvement, management can be directed to suppression of symptoms. Analgesics and antimalarials are mainstays. NSAIDs are useful analgesics/anti-inflammatories, particularly for arthritis/arthralgias. However, two major issues indicate caution in using NSAIDs. First, SLE patients compared with the general population are at increased risk for NSAID-induced aseptic meningitis, elevated serum transaminases, hypertension, and renal dysfunction. Second, all NSAIDs, particularly those that inhibit cyclooxygenase-2 specifically, may increase risk for myocardial infarction. Acetaminophen to control pain may be a good strategy, but NSAIDs are more effective in some patients. The relative hazards of NSAIDs compared with low-dose glucocorticoid therapy have not been established. Antimalarials (hydroxychloroquine, chloroquine, and quinacrine) often reduce dermatitis, arthritis, and fatigue. A randomized, placebo-controlled, prospective trial has shown that withdrawal of hydroxychloroquine results in increased numbers of disease flares; hydroxychloroquine also reduces accrual of tissue damage, including renal damage, over time. Because of potential retinal toxicity, patients receiving antimalarials should undergo ophthalmologic examinations annually. A placebo-controlled prospective trial suggests that administration of dehydroepiandrosterone may reduce disease activity. If quality of life is inadequate despite these conservative measures, treatment with low doses of systemic glucocorticoids may be necessary. The clinician may also consider treatment with belimumab (anti-BLyS) in these patients, although published clinical trials enrolled patients who had failed to respond to conservative therapies. Lupus dermatitis should be managed with topical sunscreens, antimalarials, topical glucocorticoids, and/or tacrolimus, and if severe or unresponsive, systemic glucocorticoids with or without mycophenolate mofetil.

LIFE-THREATENING SLE: PROLIFERATIVE FORMS OF LUPUS NEPHRITIS

Guidelines for management of lupus nephritis have been published recently by the American College of Rheumatology and the European League Against Rheumatism (encompassed and referenced in Fig. 378-2 and Table 378-5). The mainstay of treatment for any inflammatory life-threatening or organ-threatening manifestations of SLE is systemic glucocorticoids (0.5–1 mg/kg per day PO or 500–1000 mg of methylprednisolone sodium succinate IV daily for 3 days followed by 0.5–1 mg/kg of daily prednisone or equivalent). Evidence that glucocorticoid therapy is life-saving comes from retrospective studies from the predialysis era; survival was significantly better in people with DPGN treated with high-dose daily glucocorticoids (40–60 mg of prednisone daily for 4–6 months) versus lower doses. Currently, high doses are recommended for much shorter periods; recent trials of interventions for severe SLE use 4–6 weeks of 0.5–1 mg/kg per day of prednisone or equivalent. Thereafter, doses are tapered as rapidly as the clinical situation permits, usually to a maintenance dose ranging from 5 to 10 mg of prednisone or equivalent per day. Most patients with an episode of severe SLE require many years of maintenance therapy with low-dose glucocorticoids, which can be increased to prevent or treat disease flares. Frequent attempts to gradually reduce the glucocorticoid requirement are recommended because virtually everyone develops important adverse effects (Table 378-5). High-quality clinical studies regarding initiating therapy for severe, active SLE with IV pulses of high-dose glucocorticoids are not available. Most recent clinical trials in lupus nephritis have initiated therapy with high-dose IV glucocorticoid pulses (500–1000 mg daily for 3–5 days). This approach must be tempered by safety considerations, such as the presence of conditions adversely affected by glucocorticoids (e.g., infection, hyperglycemia, hypertension, osteoporosis).

Cytotoxic/immunosuppressive agents added to glucocorticoids are recommended to treat serious SLE. Almost all prospective controlled

trials in SLE involving cytotoxic agents have been conducted in combination with glucocorticoids in patients with lupus nephritis. Therefore, the following recommendations apply to treatment of nephritis. Either cyclophosphamide (an alkylating agent) or mycophenolate mofetil (a relatively lymphocyte-specific inhibitor of inosine monophosphatase and therefore of purine synthesis) is an acceptable choice for induction of improvement in severely ill patients; azathioprine (a purine analogue and cycle-specific antimetabolite) may be effective but is slower to influence response and associated with more flares. In patients whose renal biopsies show ISN grade III or IV disease, early treatment with combinations of glucocorticoids and cyclophosphamide reduces progression to ESRD and death. Shorter-term studies with glucocorticoids plus mycophenolate mofetil (prospective randomized trials of 6 months, follow-up studies of 36 months) show that this regimen is similar to cyclophosphamide in achieving improvement. Comparisons are complicated by effects of race, since higher proportions of African Americans (and other non-Asian, nonwhite races) respond to mycophenolate than to cyclophosphamide, whereas similar proportions of whites and Asians respond to each drug. Regarding toxicity, diarrhea is more common with mycophenolate mofetil; amenorrhea, leukopenia, and nausea are more common with cyclophosphamide. Importantly, rates of severe infections and death are similar in meta-analyses. Two different regimens of IV cyclophosphamide are available. For white patients with northern European backgrounds, low doses of cyclophosphamide (500 mg every 2 weeks for six total doses, followed by azathioprine or mycophenolate maintenance) are as effective as standard high doses, with less toxicity. Ten-year follow-up has shown no differences between the high-dose and low-dose groups (death or ESRD in 9–20% of patients in each group). The majority of the European patients were white; it is not clear whether the data apply to U.S. populations. High-dose cyclophosphamide (500–1000 mg/m² body surface area given monthly IV for 6 months, followed by azathioprine or mycophenolate maintenance) is an acceptable approach for patients with severe nephritis (e.g., multiple cellular crescents and/or fibrinoid necrosis on renal biopsy, or rapidly progressive glomerulonephritis). Cyclophosphamide and mycophenolate responses begin 3–16 weeks after treatment is initiated, whereas glucocorticoid responses may begin within 24 h.

For maintenance therapy, mycophenolate and azathioprine probably are similar in efficacy and toxicity; both are safer than cyclophosphamide. In a recently published multicenter study, mycophenolate was superior to azathioprine in maintaining renal function and survival in patients who responded to induction therapy with either cyclophosphamide or mycophenolate. The incidence of ovarian failure, a common effect of high-dose cyclophosphamide therapy (but probably not of low-dose therapy), can be reduced by treatment with a gonadotropin-releasing hormone agonist (e.g., leuprolide 3.75 mg intramuscularly) prior to each monthly cyclophosphamide dose. Patients with high serum creatinine levels (e.g., ≥ 265 $\mu\text{mol/L}$ [≥ 3.0 mg/dL]) many months in duration and high chronicity scores on renal biopsy are not likely to respond to any of these therapies. In general, it may be better to induce improvement in an African-American or Hispanic patient with proliferative glomerulonephritis with mycophenolate mofetil (2–3 g daily) rather than cyclophosphamide, with the option to switch if no evidence of response is detectable after 3–6 months of treatment. For whites and Asians, induction with either mycophenolate mofetil or cyclophosphamide is acceptable. Cyclophosphamide may be discontinued when it is clear that a patient is improving. The number of SLE flares is reduced by maintenance therapy with mycophenolate mofetil (1.5–2 g daily) or azathioprine (1–2.5 mg/kg per day). Both cyclophosphamide and mycophenolate mofetil are potentially teratogenic; patients should be off either medication for at least 3 months before attempting to conceive. Azathioprine can be used if necessary to control active SLE in patients who are pregnant. If azathioprine is used either for induction or maintenance therapy, patients may be prescreened for homozygous deficiency of the TMPT enzyme (which is required to metabolize the 6-mercaptopurine product of azathioprine) because they are at higher risk for bone marrow suppression.

Good improvement occurs in ~80% of lupus nephritis patients receiving either cyclophosphamide or mycophenolate at 1–2 years of follow-up. However, in some studies, at least 50% of these individuals have flares of nephritis over the next 5 years, and re-treatment is required; such individuals are more likely to progress to ESRD. Long-term outcome of lupus nephritis to most interventions is better in whites than in African Americans. Methotrexate (a folinic acid antagonist) may have a role in the treatment of arthritis and dermatitis but probably not in nephritis or other life-threatening disease. Small controlled trials (in Asia) of leflunomide, a relatively lymphocyte-specific pyrimidine antagonist licensed for use in rheumatoid arthritis, have suggested it can suppress disease activity in some SLE patients. Cyclosporine and tacrolimus, which inhibit production of IL-2 and T lymphocyte functions, have not been studied in prospective controlled trials in SLE in the United States; several studies in Asia have shown they are effective in lupus nephritis. Because they have potential nephrotoxicity but little bone marrow toxicity, the author uses them for periods of a few months in patients with steroid-resistant cytopenias of SLE or in steroid-resistant patients who have developed bone marrow suppression from standard cytotoxic agents.

Use of biologics directed against B cells for active SLE is under intense study. Use of anti-CD20 (rituximab), particularly in patients with SLE who are resistant to the more standard combination therapies discussed above, is controversial. Several open trials have shown efficacy in a majority of such patients, both for nephritis and for extrarenal lupus. However, recent prospective placebo-controlled randomized trials, one in renal and one in nonrenal SLE, did not show a difference between anti-CD20 and placebo when added to standard combination therapies. In contrast, recent trials of standard therapy plus belimumab (anti-BLyS, which binds soluble BLyS/BAFF, which is required for maturation of naïve and transitional B cells to plasma cells and memory B cells) showed improvement in 51% of SLE patients compared to 36% of those on placebo; these differences were statistically significant. The U.S. Food and Drug Administration (FDA) has approved belimumab for treatment of seropositive patients with SLE who have failed standard treatments. The belimumab trial did not include patients with active nephritis or CNS disease. Post hoc analyses have shown that the SLE patient most likely to respond to belimumab has fairly robust clinical activity (a Systemic Lupus Erythematosus Disease Activity Index [SLEDAI] score of ≥ 10), positive anti-DNA, and low serum complement. SLEDAI is a widely used measure of SLE disease activity; scores >3 reflect clinically active disease. At this time, it is useful to add belimumab to the therapeutic armamentarium in SLE, and it is clear that some patients benefit. However, its role in management of lupus nephritis is not yet known.

SPECIAL CONDITIONS IN SLE THAT MAY REQUIRE ADDITIONAL OR DIFFERENT THERAPIES

Crescentic Lupus Nephritis The presence of cellular or fibrotic crescents in glomeruli with proliferative glomerulonephritis indicates a worse prognosis than in patients without this feature. There are no large prospective multinational controlled trials showing efficacy of cyclophosphamide, mycophenolate, cyclosporine, or tacrolimus in such cases. Most authorities currently recommend that high-dose cyclophosphamide is the induction therapy of choice, in addition to high-dose glucocorticoids. One prospective trial from China showed superiority of mycophenolate to cyclophosphamide.

Membranous Lupus Nephritis Most SLE patients with membranous (INS-V) nephritis also have proliferative changes and should be treated for proliferative disease. However, some have pure membranous changes. Treatment for this group is less well defined. Some authorities do not recommend immunosuppression unless proteinuria is in the nephrotic range (although treatment with angiotensin-converting enzyme inhibitors or angiotensin II receptor blockers is recommended). In those patients, recent prospective controlled trials suggest that alternate-day glucocorticoids plus cyclophosphamide or mycophenolate mofetil or cyclosporine are all effective in the majority of

patients in reducing proteinuria. It is more controversial whether they preserve renal function over the long term.

Pregnancy and Lupus Fertility rates for men and women with SLE are probably normal. However, rate of fetal loss is increased (approximately two- to threefold) in women with SLE. Fetal demise is higher in mothers with high disease activity, antiphospholipid antibodies, and/or active nephritis. Suppression of disease activity can be achieved by administration of systemic glucocorticoids. A placental enzyme, 11- β -dehydrogenase 2, deactivates glucocorticoids; it is more effective in deactivating prednisone and prednisolone than the fluorinated glucocorticoids dexamethasone and betamethasone. Glucocorticoids are listed by the FDA as pregnancy category A (no evidence of teratogenicity in human studies); cyclosporine, tacrolimus, and rituximab are listed as category C (may be teratogenic in animals but no good evidence in humans); azathioprine, hydroxychloroquine, mycophenolate mofetil, and cyclophosphamide are category D (there is evidence of teratogenicity in humans, but benefits might outweigh risks in certain situations); and methotrexate is category X (risks outweigh benefits). Therefore, active SLE in pregnant women should be controlled with hydroxychloroquine and, if necessary, prednisone/prednisolone at the lowest effective doses for the shortest time required. Azathioprine may be added if these treatments do not suppress disease activity. Adverse effects of prenatal glucocorticoid exposure (primarily betamethasone) on offspring may include low birth weight, developmental abnormalities in the CNS, and predilection toward adult metabolic syndrome. It is likely that each of these glucocorticoids and immunosuppressive medications gets into breast milk, at least in low levels; patients should consider not breastfeeding if they need therapy for SLE. In SLE patients with antiphospholipid antibodies (on at least two occasions) and prior fetal losses, treatment with heparin (usually low-molecular-weight) plus low-dose aspirin has been shown in prospective controlled trials to increase significantly the proportion of live births; however, a recent prospective trial showed no differences in fetal outcomes in women taking aspirin compared to those taking aspirin plus low-molecular-weight heparin. An additional potential problem for the fetus is the presence of antibodies to Ro, sometimes associated with neonatal lupus consisting of rash and congenital heart block with or without cardiomyopathy. The cardiac manifestations can be life-threatening; therefore the presence of anti-Ro requires vigilant monitoring of fetal heart rates with prompt intervention (delivery if possible) if distress occurs. Recent evidence shows that hydroxychloroquine treatment of an anti-Ro-positive mother whose infant develops congenital heart block significantly reduces the chance that subsequent fetuses will develop heart block. There is some evidence that dexamethasone treatment of a mother in whom first- or second-degree heart block is detected in utero may sometimes prevent progression of heart block. Women with SLE usually tolerate pregnancy without disease flares. However, a small proportion develops severe flares requiring aggressive glucocorticoid therapy or early delivery. Poor maternal outcomes are highest in women with active nephritis or irreversible organ damage in kidneys, brain, or heart.

Lupus and Antiphospholipid Syndrome (APS) Patients with SLE who have venous or arterial clotting and/or repeated fetal losses and at least two positive tests for antiphospholipid antibodies have APS and should be managed with long-term anticoagulation ([Chap. 379](#)). A target international normalized ratio (INR) of 2.0–2.5 is recommended for patients with one episode of venous clotting; an INR of 3.0–3.5 is recommended for patients with recurring clots or arterial clotting, particularly in the CNS. Recommendations are based on both retrospective and prospective studies of posttreatment clotting events and adverse effects from anticoagulation.

Microvascular Thrombotic Crisis (Thrombotic Thrombocytopenic Purpura, Hemolytic-Uremic Syndrome) This syndrome of hemolysis, thrombocytopenia, and microvascular thrombosis in kidneys, brain, and other tissues carries a high mortality rate and occurs most commonly in young individuals with lupus nephritis. The most useful laboratory tests are identification of schistocytes on peripheral blood smears, elevated serum levels of lactate dehydrogenase, and antibodies to

2134 ADAMS13. Plasma exchange or extensive plasmapheresis is usually life-saving; most authorities recommend concomitant glucocorticoid therapy; there is no evidence that cytotoxic drugs are effective.

Lupus Dermatitis Patients with any form of lupus dermatitis should minimize exposure to ultraviolet light, using appropriate clothing and sunscreens with a sun protection factor of at least 30. Topical glucocorticoids and antimalarials (such as hydroxychloroquine) are effective in reducing lesion severity in most patients and are relatively safe. Systemic treatment with retinoic acid is a useful strategy in patients with inadequate improvement on topical glucocorticoids and antimalarials; adverse effects are potentially severe (particularly fetal abnormalities), and there are stringent reporting requirements for its use in the United States. Extensive, pruritic, bullous, or ulcerating dermatitides usually improve promptly after institution of systemic glucocorticoids; tapering may be accompanied by flare of lesions, thus necessitating use of a second medication such as hydroxychloroquine, retinoids, or cytotoxic medications such as methotrexate, azathioprine, or mycophenolate mofetil. In therapy-resistant lupus dermatitis there are reports of success with topical tacrolimus (caution must be exerted because of the possible increased risk for malignancies) or with systemic dapsone or thalidomide (the extreme danger of fetal deformities from thalidomide requires permission from and supervision by the supplier).

PREVENTIVE THERAPIES

Prevention of complications of SLE and its therapy include providing appropriate vaccinations (the administration of influenza and pneumococcal vaccines has been studied in patients with SLE; flare rates are similar to those receiving placebo) and suppressing recurrent urinary tract infections. Vaccination with attenuated live viruses is generally discouraged in patients who are immunosuppressed. Strategies to prevent osteoporosis should be initiated in most patients likely to require long-term glucocorticoid therapy and/or with other predisposing factors. Postmenopausal women can be protected from steroid-induced osteoporosis with either bisphosphonates or denosumab. Safety of long-term use of these strategies in premenopausal women is not well established. Control of hypertension and appropriate prevention strategies for atherosclerosis, including monitoring and treatment of dyslipidemias, management of hyperglycemia, and management of obesity, are recommended.

EXPERIMENTAL THERAPIES

Studies of highly targeted experimental therapies for SLE are in progress. They include targeting (1) activated B lymphocytes with anti-CD22 or TACI-Ig, (2) inhibition of IFN- α , (3) inhibition of B/T cell second signal coactivation with CTLA-Ig, (4) inhibition of innate immune activation via TLR7 or TLR7 and 9, (5) induction of regulatory T cells with peptides from immunoglobulins or autoantigens; (6) suppression of T cells, B cells, and monocyte/macrophages with laquinimod; and (7) inhibition of lymphocyte activation by blockade of Jak/Stat. A few studies have used vigorous untargeted immunosuppression with high-dose cyclophosphamide plus anti-T cell strategies, with rescue by transplantation of autologous hematopoietic stem cells for the treatment of severe and refractory SLE. One U.S. report showed an estimated mortality rate over 5 years of 15% and sustained remission in 50%. It is hoped that in the next edition of this text, we will be able to recommend more effective and less toxic approaches to treatment of SLE based on some of these strategies.

PATIENT OUTCOMES, PROGNOSIS, AND SURVIVAL



Survival in patients with SLE in the United States, Canada, Europe, and China is approximately 95% at 5 years, 90% at 10 years, and 78% at 20 years. In the United States, African Americans and Hispanic Americans with a mestizo heritage have a worse prognosis than whites, whereas Africans in Africa and Hispanic Americans with a Puerto Rican origin do not. The relative importance of gene mixtures and environmental differences accounting for ethnic differences is not known. Poor prognosis (~50% mortality in 10 years) in most series is associated with (at the time of diagnosis) high serum

creatinine levels ($>124 \mu\text{mol/L}$ [$>1.4 \text{ mg/dL}$]), hypertension, nephrotic syndrome (24-h urine protein excretion $>2.6 \text{ g}$), anemia (hemoglobin $<124 \text{ g/L}$ [$<12.4 \text{ g/dL}$]), hypoalbuminemia, hypocomplementemia, antiphospholipid antibodies, male sex, ethnicity (African American, Hispanic with mestizo heritage), and low socioeconomic status. Data regarding outcomes in SLE patients with renal transplants show mixed results: some series show a twofold increase in graft rejection compared to patients with other causes of ESRD, whereas others show no differences. Overall patient survival is comparable (85% at 2 years). Lupus nephritis occurs in approximately 10% of transplanted kidneys. Disability in patients with SLE is common due primarily to chronic fatigue, arthritis, and pain, as well as renal disease. As many as 25% of patients may experience remissions, sometimes for a few years, but these are rarely permanent. The leading causes of death in the first decade of disease are systemic disease activity, renal failure, and infections; subsequently, thromboembolic events become increasingly frequent causes of mortality.

DRUG-INDUCED LUPUS

This is a syndrome of positive ANA associated with symptoms such as fever, malaise, arthritis or intense arthralgias/myalgias, serositis, and/or rash. The syndrome appears during therapy with certain medications and biologic agents, is predominant in whites, has less female predilection than SLE, rarely involves kidneys or brain, is rarely associated with anti-dsDNA, is commonly associated with antibodies to histones, and usually resolves over several weeks after discontinuation of the offending medication. The list of substances that can induce lupus-like disease is long. Among the most frequent are the antiarrhythmics procainamide, disopyramide, and propafenone; the antihypertensive hydralazine; several angiotensin-converting enzyme inhibitors and beta blockers; the antithyroid propylthiouracil; the antipsychotics chlorpromazine and lithium; the anticonvulsants carbamazepine and phenytoin; the antibiotics isoniazid, minocycline, and nitrofurantoin (Macrochantin); the antirheumatic sulfasalazine; the diuretic hydrochlorothiazide; the antihyperlipidemics lovastatin and simvastatin; and IFNs and TNF inhibitors. ANA usually appears before symptoms; however, many of the medications mentioned above induce ANA in patients who never develop symptoms of drug-induced lupus. It is appropriate to test for ANA at the first hint of relevant symptoms and to use test results to help decide whether to withdraw the suspect agent.

379 Antiphospholipid Syndrome

Haralampos M. Moutsopoulos, Panayiotis G. Vlachoyiannopoulos

DEFINITIONS

Antiphospholipid syndrome (APS) is an autoantibody-mediated acquired thrombophilia characterized by recurrent arterial or venous thrombosis and/or pregnancy morbidity. The major autoantibodies detected in the patient's sera are directed against phospholipid (PL)-binding plasma proteins, mainly against a 43-kDa plasma apolipoprotein known as β_2 glycoprotein I ($\beta_2\text{GPI}$) and prothrombin. The plasma concentration of $\beta_2\text{GPI}$ is 50–200 $\mu\text{g/mL}$. $\beta_2\text{GPI}$ consists of 326 amino acids arranged in five domains (I through V). Domain V forms a positively charged patch, suitable to interact with negatively charged PL. In plasma, $\beta_2\text{GPI}$ has a circular conformation with domain V binding to and concealing the B cell epitopes lying on domain I. Another group of antibodies termed *lupus anticoagulant* (LA) elongate clotting times in vitro; this elongation is not corrected by adding normal plasma to the detection system (Table 379-1). Patients with APS often possess antibodies recognizing *Treponema pallidum* PL/cholesterol complexes, which are detected as biologic false-positive serologic tests for syphilis

TABLE 379-1 CLASSIFICATION AND NOMENCLATURE OF ANTIPHOSPHOLIPID ANTIBODIES

Name	Assay for their Detection	Comments
Antibodies against cardiolipin (aCL)	Enzyme-linked immunosorbent assay (ELISA) using as antigen cardiolipin (CL), a negatively charged phospholipid	aCL from patients with APS recognize β 2GPI existing in the human serum as well as in bovine serum, which is used to block the nonspecific bindings sites on the ELISA plate. CL simply stabilizes β 2GPI at high concentration on the polystyrene surface.
Antibodies against β 2GPI (anti- β 2GPI)	ELISA using as antigen affinity purified or recombinant β 2GPI in the absence of PL	Antibodies recognize β 2GPI bound in the absence of CL to an oxidized polystyrene surface, where oxygen atoms in the moieties C-O or C=O were introduced by γ -irradiation.
Lupus anticoagulant (LA)	Activated partial thromboplastin time (aPTT) Kaolin clotting time (KCT) Dilute Russel viper venom test (DRVVT)	Antibodies recognize β 2GPI or prothrombin (PT) and elongate aPTT, implying that they interfere with the generation of thrombin by prothrombin. Prolongation of the clotting times is an in vitro phenomenon, and LA induces thromboses in vivo.

Abbreviations: APL, antiphospholipid syndrome; β 2GPI, β 2 glycoprotein I; PL, phospholipid.

(BFP-ST5) and Venereal Disease Research Laboratory (VDRL) tests. APS may occur alone (primary) or in association with any other autoimmune disease (secondary). Catastrophic APS (CAPS) is defined as a rapidly progressive thromboembolic disease involving simultaneously three or more organs, organ systems, or tissues leading to corresponding functional defects.

EPIDEMIOLOGY

Anti-PL (aPL)-binding plasma protein antibodies occur in 1–5% of the general population. Their prevalence increases with age; however, it is questionable whether they induce thrombotic events in elderly individuals. One-third of patients with systemic lupus erythematosus (SLE) (Chap. 378) possess these antibodies, whereas their prevalence in other autoimmune connective tissue disorders, such as systemic sclerosis (scleroderma), Sjögren's syndrome, dermatomyositis, rheumatoid arthritis, and early undifferentiated connective tissue disease, ranges from 6 to 15%. One-third of aPL-positive individuals experience thrombotic events or pregnancy morbidity.

PATHOGENESIS

The trigger for the induction of antibodies to PL-binding proteins is not known. However infections, oxidative stress, major physical stresses such as surgery, and discontinuation of anticoagulant treatment may induce the exacerbation of the disease. Experimental data have shown that these phenomena are induced via (1) conformational changes of β 2GPI either complexed with microbial antigens or dimerization through interaction with endothelial cell surface receptor annexin 2/TLR4, the platelet receptors apolipoprotein E receptor 2' (apoER2') and/or GPIb/IX/V receptor, and/or the chemokine platelet factor 4 (PF4); or (2) impaired defensive mechanisms such as reduced generation of endothelial nitric oxide synthase. Adherence of β 2GPI to apoER2', GPIb/IX/V receptor, and/or PF4 induces activation of endothelial cells, platelets, and monocytes. This process activates downstream pathways such as p38 mitogen-activated protein (p38 MAP) kinase and nuclear factor (NF)- κ B, leading to the following events: secretion of proinflammatory cytokines, such as interleukin (IL) 1, IL-6, and IL-8; the expression of adhesion molecules; inhibition of cell-surface plasminogen activation; and expression of tissue factor. The above events change the phenotype of these cells to a prothrombotic form. In addition, anti- β 2GPI antibodies induce fetal injury in mice through complement activation, as shown by the evidence that C4-deficient mice were protected from fetal injury.

CLINICAL MANIFESTATIONS AND LABORATORY FINDINGS

Clinical manifestations represent mainly a direct or indirect expression of venous or arterial thrombosis and/or pregnancy morbidity (Table 379-2). Clinical features associated with venous thrombosis are superficial and deep vein thrombosis, cerebral venous thrombosis, signs and symptoms of intracranial hypertension, retinal vein thrombosis, pulmonary emboli, pulmonary arterial hypertension, and Budd-Chiari syndrome. Livedo reticularis consists of a mottled reticular vascular pattern that appears as a lace-like, purplish discoloration of the skin. It is probably caused by swelling of the venules owing to obstruction of capillaries by thrombi. This clinical manifestation

correlates with vascular lesions such as those in the central nervous system as well as aseptic bone necrosis. Arterial thrombosis is manifested as migraines, cognitive dysfunction, transient ischemic attacks, stroke, myocardial infarction, arterial thrombosis of upper and lower

TABLE 379-2 CLINICAL FEATURES OF ANTIPHOSPHOLIPID SYNDROME

Manifestation	%
Venous Thrombosis and Related Consequences	
Deep vein thrombosis	39
Livedo reticularis	24
Pulmonary embolism	14
Superficial thrombophlebitis	12
Thrombosis in various other sites	11
Arterial Thrombosis and Related Consequences	
Stroke	20
Cardiac valve thickening/dysfunction and/or Libman-Sacks vegetations	14
Transient ischemic attack	11
Myocardial ischemia (infarction or angina) and coronary bypass thrombosis	10
Leg ulcers and/or digital gangrene	9
Arterial thrombosis in the extremities	7
Retinal artery thrombosis/amaurosis fugax	7
Ischemia of visceral organs or avascular necrosis of bone	6
Multi-infarct dementia	3
Neurologic Manifestations of Uncertain Etiology	
Migraine	20
Epilepsy	7
Chorea	1
Cerebellar ataxia	1
Transverse myelopathy	0.5
Renal Manifestations Due to Various Reasons (Renal Artery/Renal Vein/Glomerular Thrombosis, Fibrous Intima Hyperplasia)	
	3
Osteoarticular Manifestations	
Arthralgia	39
Arthritis	27
Obstetric Manifestations (Referred to the Number of Pregnancies)	
Preeclampsia	10
Eclampsia	4
Fetal Manifestations (Referred to the Number of Pregnancies)	
Early fetal loss (<10 weeks)	35
Late fetal loss (\geq 10 weeks)	17
Premature birth among the live births	11
Hematologic Manifestations	
Thrombocytopenia	30
Autoimmune hemolytic anemia	10

Source: Adapted from R Cervera et al: Arthritis Rheum 46:1019, 2002.

2136 extremities, ischemic leg ulcers, digital gangrene, avascular necrosis of bone, retinal artery occlusion leading to painless transient vision loss, renal artery stenosis, and glomerular lesions, as well as infarcts of spleen, pancreas, and adrenals. Libman-Sacks endocarditis consists of very small vegetations, histologically characterized by organized platelet-fibrin microthrombi surrounded by growing fibroblasts and macrophages. Glomerular lesions are manifested with hypertension, mildly elevated serum creatinine levels, proteinuria, and mild hematuria. Histologically, these lesions are characterized in an acute phase by thrombotic microangiopathy involving glomerular capillaries, and in a chronic phase with fibrous intima hyperplasia, fibrous and/or fibrocellular occlusions of arterioles, and focal cortical atrophy (Table 379-2). Premature atherosclerosis has been recognized as a rare feature of APS. Coombs-positive hemolytic anemia and thrombocytopenia are laboratory findings associated with APS. Discontinuation of therapy, major surgery, infection, and trauma may trigger CAPS.

DIAGNOSIS AND DIFFERENTIAL DIAGNOSIS

The diagnosis of APS should be seriously considered in cases of thrombosis, cerebral vascular accidents in individuals younger than 55 years of age, or pregnancy morbidity in the presence of livedo reticularis or thrombocytopenia. In these cases, aPL antibodies should be measured. The presence of at least one clinical and one laboratory criterion ensures the diagnosis even in the presence of other causes of thrombophilia. Clinical criteria include: (1) vascular thrombosis defined as one or more clinical episodes of arterial, venous, or small vessel thrombosis in any tissue or organ; and (2) pregnancy morbidity, defined as (a) one or more unexplained deaths of a morphologically normal fetus at or beyond the tenth week of gestation; (b) one or more premature births of a morphologically normal neonate before the thirty-fourth week of gestation because of eclampsia, severe preeclampsia, or placental insufficiency; or (c) three or more unexplained consecutive spontaneous abortions before the tenth week of gestation. Laboratory criteria include (1) LA, (2) anticardiolipin (aCL), and/or (3) anti- β 2GPI antibodies, at intermediate or high titers on two occasions, 12 weeks apart.

Differential diagnosis is based on the exclusion of other inherited or acquired causes of thrombophilia (Chap. 141), Coombs-positive hemolytic anemia (Chap. 129), and thrombocytopenia (Chap. 140). Livedo reticularis with or without a painful ulceration on the lower extremities also may be a manifestation of disorders affecting (1) the vascular wall, such as polyarteritis nodosa, SLE, cryoglobulinemia, and lymphomas; or (2) the vascular lumen, such as myeloproliferative disorders, atherosclerosis, hypercholesterolemia, or other causes of thrombophilia.

TREATMENT ANTIPHOSPHOLIPID SYNDROME

After the first thrombotic event, APS patients should be placed on warfarin for life, aiming to achieve an international normalized ratio (INR) ranging from 2.5 to 3.5, alone or in combination with 80 mg of aspirin daily. Pregnancy morbidity is prevented by a combination of heparin with aspirin 80 mg daily. IV immunoglobulin (IVIg) 400 mg/kg every day for 5 days may also prevent abortions, whereas glucocorticoids are ineffective. Patients with aPL in the absence of any clinical event who are simultaneously positive for aCL, anti- β 2GPI, and LA or have SLE are at risk to develop thrombotic events and can be protected by aspirin 80 mg daily.

Some patients with APS and patients with CAPS have recurrent thrombotic events despite appropriate anticoagulation. In these cases, IVIg 400 mg/kg every day for 5 days may be of benefit. Patients with CAPS, who are treated in the intensive care unit, are unable to receive warfarin; in this situation, therapeutic doses of low-molecular-weight heparin should be administered. In cases of heparin-induced thrombocytopenia and thrombosis syndrome, inhibitors of phospholipid-bound activated factor X (FXa), such as fondaparinux 7.5 mg SC daily or rivaroxaban 10 mg PO daily, are effective. The above drugs are administered by fixed doses and do not require close monitoring; their safety during the first trimester of pregnancy has not been clearly established.

380 Rheumatoid Arthritis

Ankoor Shah, E. William St. Clair

INTRODUCTION

Rheumatoid arthritis (RA) is a chronic inflammatory disease of unknown etiology marked by a symmetric, peripheral polyarthritis. It is the most common form of chronic inflammatory arthritis and often results in joint damage and physical disability. Because it is a systemic disease, RA may result in a variety of extraarticular manifestations, including fatigue, subcutaneous nodules, lung involvement, pericarditis, peripheral neuropathy, vasculitis, and hematologic abnormalities.

Insights gained by a wealth of basic and clinical research over the past two decades have revolutionized the contemporary paradigms for the diagnosis and management of RA. Serum antibodies to cyclic citrullinated peptides (anti-CCPs) are routinely used along with rheumatoid factor as a biomarker of diagnostic and prognostic significance. Advances in imaging modalities have improved our ability to detect joint inflammation and destruction in RA. The science of RA has taken a major leap forward with the identification of new disease-related genes and further deciphering of the molecular pathways of disease pathogenesis. The relative importance of these different mechanisms has been highlighted by the observed benefits of the new class of highly targeted biologic and small-molecule therapies. Despite these gains, incomplete understanding of the initiating pathogenic pathways of RA remains a sizable barrier to its cure and prevention.

The last two decades have witnessed a remarkable improvement in the outcomes of RA. The historic descriptions of crippling arthritis are currently encountered much less frequently. Much of this progress can be traced to the expanded therapeutic armamentarium and the adoption of early treatment intervention. The shift in treatment strategy dictates a new mind-set for primary care practitioners—namely, one that demands early referral of patients with inflammatory arthritis to a rheumatologist for prompt diagnosis and initiation of therapy. Only then will patients achieve their best outcomes.

CLINICAL FEATURES

The incidence of RA increases between 25 and 55 years of age, after which it plateaus until the age of 75 and then decreases. The presenting symptoms of RA typically result from inflammation of the joints, tendons, and bursae. Patients often complain of early morning joint stiffness lasting more than 1 h that eases with physical activity. The earliest involved joints are typically the small joints of the hands and feet. The initial pattern of joint involvement may be monoarticular, oligoarticular (≤ 4 joints), or polyarticular (>5 joints), usually in a symmetric distribution. Some patients with inflammatory arthritis will present with too few affected joints to be classified as having RA—so-called undifferentiated inflammatory arthritis. Those with an undifferentiated arthritis who are most likely to be diagnosed later with RA have a higher number of tender and swollen joints, test positive for serum rheumatoid factor (RF) or anti-CCP antibodies, and have higher scores for physical disability.

Once the disease process of RA is established, the wrists, metacarpophalangeal (MCP), and proximal interphalangeal (PIP) joints stand out as the most frequently involved joints (Fig. 380-1). Distal interphalangeal (DIP) joint involvement may occur in RA, but it usually is a manifestation of coexistent osteoarthritis. Flexor tendon tenosynovitis is a frequent hallmark of RA and leads to decreased range of motion, reduced grip strength, and “trigger” fingers. Progressive destruction of the joints and soft tissues may lead to chronic, irreversible deformities. Ulnar deviation results from subluxation of the MCP joints, with subluxation of the proximal phalanx to the volar side of the hand. Hyperextension of the PIP joint with flexion of the DIP joint (“swan-neck deformity”), flexion of the PIP joint with hyperextension of the DIP joint (“boutonnière deformity”), and subluxation of the first MCP joint with hyperextension of the first interphalangeal (IP) joint (“Z-line



FIGURE 380-1 Metacarpophalangeal and proximal interphalangeal joint swelling in rheumatoid arthritis. (Courtesy of the American College of Rheumatology Image Bank.)

deformity”) also may result from damage to the tendons, joint capsule, and other soft tissues in these small joints. Inflammation about the ulnar styloid and tenosynovitis of the extensor carpi ulnaris may cause subluxation of the distal ulna, resulting in a “piano-key movement” of the ulnar styloid. Although metatarsophalangeal (MTP) joint involvement in the feet is an early feature of disease, chronic inflammation of the ankle and midtarsal regions usually comes later and may lead to pes planovalgus (“flat feet”). Large joints, including the knees and shoulders, are often affected in established disease, although these joints may remain asymptomatic for many years after onset.

Atlantoaxial involvement of the cervical spine is clinically noteworthy because of its potential to cause compressive myelopathy and neurologic dysfunction. Neurologic manifestations are rarely a presenting sign or symptom of atlantoaxial disease, but they may evolve over time with progressive instability of C1 on C2. The prevalence of atlantoaxial subluxation has been declining in recent years, and occurs now in less than 10% of patients. Unlike the spondyloarthritides (Chap. 384), RA rarely affects the thoracic and lumbar spine. Radiographic abnormalities of the temporomandibular joint occur commonly in patients with RA, but they are generally not associated with significant symptoms or functional impairment.

Extraarticular manifestations may develop during the clinical course of RA, even prior to the onset of arthritis (Fig. 380-2). Patients most likely to develop extraarticular disease have a history of smoking, have early onset of significant physical disability, and test positive for serum RF. Subcutaneous nodules, secondary Sjögren’s syndrome, pulmonary nodules, and anemia are among the most frequently observed

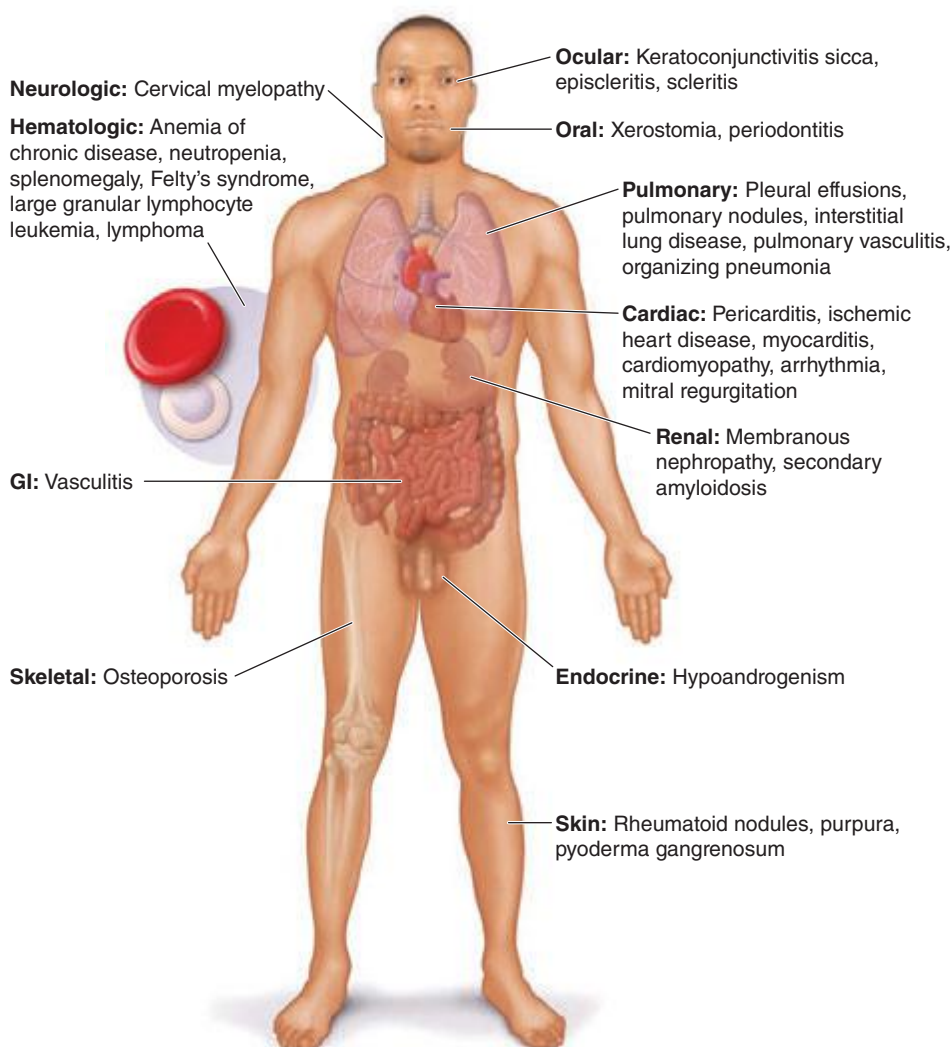


FIGURE 380-2 Extraarticular manifestations of rheumatoid arthritis.

2138 extraarticular manifestations. Recent studies have shown a decrease in the incidence and severity of at least some extraarticular manifestations, particularly Felty's syndrome and vasculitis.

The most common systemic and extraarticular features of RA are described in more detail in the sections below.

CONSTITUTIONAL

These signs and symptoms include weight loss, fever, fatigue, malaise, depression, and in the most severe cases, cachexia; they generally reflect a high degree of inflammation and may even precede the onset of joint symptoms. In general, the presence of a fever of $>38.3^{\circ}\text{C}$ (101°F) at any time during the clinical course should raise suspicion of systemic vasculitis (see below) or infection.

NODULES

Subcutaneous nodules occur in 30–40% of patients and more commonly in those with the highest levels of disease activity, the disease-related shared epitope (see below), a positive test for serum RF, and radiographic evidence of joint erosions. When palpated, the nodules are generally firm; nontender; and adherent to periosteum, tendons, or bursae; developing in areas of the skeleton subject to repeated trauma or irritation such as the forearm, sacral prominences, and Achilles tendon. They may also occur in the lungs, pleura, pericardium, and peritoneum. Nodules are typically benign, although they can be associated with infection, ulceration, and gangrene.

SJÖGREN'S SYNDROME

Secondary Sjögren's syndrome (**Chap. 383**) is defined by the presence of either keratoconjunctivitis sicca (dry eyes) or xerostomia (dry mouth) in association with another connective tissue disease, such as RA. Approximately 10% of patients with RA have secondary Sjögren's syndrome.

PULMONARY

Pleuritis, the most common pulmonary manifestation of RA, may produce pleuritic chest pain and dyspnea, as well as a pleural friction rub and effusion. Pleural effusions tend to be exudative with increased numbers of monocytes and neutrophils. Interstitial lung disease (ILD) may also occur in patients with RA and is heralded by symptoms of dry cough and progressive shortness of breath. ILD can be associated with cigarette smoking and is generally found in patients with higher disease activity, although it may be diagnosed in up to 3.5% of patients prior to the onset of joint symptoms. Diagnosis is readily made by high-resolution chest computed tomography (CT) scan. Pulmonary function testing shows a restrictive pattern (e.g., reduced total lung capacity) with a reduced diffusing capacity for carbon monoxide (DL_{CO}). The presence of ILD confers a poor prognosis. The prognosis is not quite as poor as that of idiopathic pulmonary fibrosis (e.g., usual interstitial pneumonitis) because ILD secondary to RA responds more favorably than idiopathic ILD to immunosuppressive therapy (**Chap. 315**). Pulmonary nodules may be solitary or multiple. Caplan's syndrome is a rare subset of pulmonary nodulosis characterized by the development of nodules and pneumoconiosis following silica exposure. Other less common pulmonary findings include respiratory bronchiolitis and bronchiectasis.

CARDIAC

The most frequent site of cardiac involvement in RA is the pericardium. However, clinical manifestations of pericarditis occur in less than 10% of patients with RA despite the fact that pericardial involvement may be detected in nearly one-half of the these patients by echocardiogram or autopsy studies. Cardiomyopathy, another clinically important manifestation of RA, may result from necrotizing or granulomatous myocarditis, coronary artery disease, or diastolic dysfunction. This involvement too may be subclinical and only identified by echocardiography or cardiac magnetic resonance imaging (MRI). Rarely, the heart muscle may contain rheumatoid nodules or be infiltrated with amyloid. Mitral regurgitation is the most common

valvular abnormality in RA, occurring at a higher frequency than the general population.

VASCULITIS

Rheumatoid vasculitis (**Chap. 385**) typically occurs in patients with long-standing disease, a positive test for serum RF, and hypocomplementemia. The overall incidence has decreased significantly in the last decade to be less than 1% of patients. The cutaneous signs vary and include petechiae, purpura, digital infarcts, gangrene, livedo reticularis, and in severe cases large, painful lower extremity ulcerations. Vasculitic ulcers, which may be difficult to distinguish from those caused by venous insufficiency, may be treated successfully with immunosuppressive agents (requiring cytotoxic treatment in severe cases) as well as skin grafting. Sensorimotor polyneuropathies, such as mononeuritis multiplex, may occur in association with systemic rheumatoid vasculitis.

HEMATOLOGIC

A normochromic, normocytic anemia often develops in patients with RA and is the most common hematologic abnormality. The degree of anemia parallels the degree of inflammation, correlating with the levels of serum C-reactive protein (CRP) and erythrocyte sedimentation rate (ESR). Platelet counts may also be elevated in RA as an acute-phase reactant. Immune-mediated thrombocytopenia is rare in this disease.

Felty's syndrome is defined by the clinical triad of neutropenia, splenomegaly, and nodular RA and is seen in less than 1% of patients, although its incidence appears to be declining in the face of more aggressive treatment of the joint disease. It typically occurs in the late stages of severe RA and is more common in whites than other racial groups. T cell large granular lymphocyte leukemia (T-LGL) may have a similar clinical presentation and often occurs in association with RA. T-LGL is characterized by a chronic, indolent clonal growth of LGL cells, leading to neutropenia and splenomegaly. As opposed to Felty's syndrome, T-LGL may develop early in the course of RA. Leukopenia apart from these disorders is uncommon and most often due to drug therapy.

LYMPHOMA

Large cohort studies have shown a two- to fourfold increased risk of lymphoma in RA patients compared with the general population. The most common histopathologic type of lymphoma is a diffuse large B cell lymphoma. The risk of developing lymphoma increases if the patient has high levels of disease activity or Felty's syndrome.

ASSOCIATED CONDITIONS

In addition to extraarticular manifestations, several conditions associated with RA contribute to disease morbidity and mortality rates. They are worthy of mention because they affect chronic disease management.

Cardiovascular Disease The most common cause of death in patients with RA is cardiovascular disease. The incidence of coronary artery disease and carotid atherosclerosis is higher in RA patients than in the general population even when controlling for traditional cardiac risk factors, such as hypertension, obesity, hypercholesterolemia, diabetes, and cigarette smoking. Furthermore, congestive heart failure (including both systolic and diastolic dysfunction) occurs at an approximately twofold higher rate in RA than in the general population. The presence of elevated serum inflammatory markers appears to confer an increased risk of cardiovascular disease in this population.

Osteoporosis Osteoporosis is more common in patients with RA than an age- and sex-matched population, with prevalence rates of 20–30%. The inflammatory milieu of the joint probably spills over into the rest of the body and promotes generalized bone loss by activating osteoclasts. Chronic use of glucocorticoids and disability-related immobility also contributes to osteoporosis. Hip fractures are more likely to occur in patients with RA and are significant predictors of increased disability and mortality rate in this disease.

Hypoandrogenism Men and postmenopausal women with RA have lower mean serum testosterone, luteinizing hormone (LH), and dehydroepiandrosterone (DHEA) levels than control populations. It has thus been hypothesized that hypoandrogenism may play a role in the pathogenesis of RA or arise as a consequence of the chronic inflammatory response. It is also important to realize that patients receiving chronic glucocorticoid therapy may develop hypoandrogenism owing to inhibition of LH and follicle-stimulating hormone (FSH) secretion from the pituitary gland. Because low testosterone levels may lead to osteoporosis, men with hypoandrogenism should be considered for androgen replacement therapy.

EPIDEMIOLOGY

RA affects approximately 0.5–1% of the adult population worldwide. There is evidence that the overall incidence of RA has been decreasing in recent decades, whereas the prevalence has remained the same because individuals with RA are living longer. The incidence and prevalence of RA varies based on geographic location, both globally and among certain ethnic groups within a country (Fig. 380-3). For example, the Native American Yakima, Pima, and Chippewa tribes of North America have reported prevalence rates in some studies of nearly 7%. In contrast, many population studies from Africa and Asia show lower prevalence rates for RA in the range of 0.2–0.4%.

Like many other autoimmune diseases, RA occurs more commonly in females than in males, with a 2–3:1 ratio. Interestingly, studies of RA from some of the Latin American and African countries show an even greater predominance of disease in females compared to males, with ratios of 6–8:1. Given this preponderance of females, various theories have been proposed to explain the possible role of estrogen in disease pathogenesis. Most of the theories center on the role of estrogens in enhancing the immune response. For example, some experimental studies have shown that estrogen can stimulate production of tumor necrosis factor α (TNF- α), a major cytokine in the pathogenesis of RA.

GENETIC CONSIDERATIONS

It has been recognized for over 30 years that genetic factors contribute to the occurrence of RA as well as to its severity. The likelihood that a first-degree relative of a patient will share the diagnosis of RA is 2–10 times greater than in the general population.

There remains, however, some uncertainty in the extent to which genetics plays a role in the causative mechanisms of RA. Although twin studies imply that genetic factors may explain up to 60% of the occurrence of RA, the more commonly stated estimate falls in the range of 10–25%. The estimate of genetic influence may vary across studies due to gene–environment interactions.

The alleles known to confer the greatest risk of RA are located within the major histocompatibility complex (MHC). It has been estimated that one-third of the genetic risk for RA resides within this locus. Most, but probably not all, of this risk is associated with allelic variation in the HLA-DRB1 gene, which encodes the MHC II β -chain molecule. The disease-associated HLA-DRB1 alleles share an amino acid sequence at positions 70–74 in the third hypervariable regions of the HLA-DR β -chain, termed the *shared epitope (SE)*. Carriership of the SE alleles is associated with production of anti-CCP antibodies and worse disease outcomes. Some of these HLA-DRB1 alleles bestow a high risk of disease (*0401), whereas others confer a more moderate risk (*0101, *0404, *1001, and *0901). Additionally, there is regional variation. In Greece, for example, where RA tends to be milder than in western European countries, RA susceptibility has been associated with the *0101 SE allele. By comparison, the *0401 or *0404 alleles are found in approximately 50–70% of northern Europeans and are the predominant risk alleles in this group. The most common disease susceptibility SE alleles in Asians, namely the Japanese, Koreans, and Chinese, are *0405 and *0901. Lastly, disease susceptibility of Native American populations such as the Pima and Tlingit Indians, where the prevalence of RA can be as high as 7%, is associated with the SE allele *1042. The risk of RA conferred by these SE alleles is less in African and Hispanic Americans than in individuals of European ancestry.

Genome-wide association studies (GWAS) have made possible the identification of several non-MHC-related genes that contribute to RA susceptibility. GWAS are based on the detection of single-nucleotide polymorphisms (SNPs), which allow for examination of the genetic architecture of complex diseases such as RA. There are approximately 10 million common SNPs within a human genome consisting of 3 billion base pairs. As a rule, GWAS identify only common variants, namely, those with a frequency of more than 5% in the general population.

Overall, several themes have emerged from GWAS in RA. First, the non-MHC loci identified as risk alleles for RA have only a modest

European ancestry:

HLA-DRB1:

- *0401
- *0404
- *0301
- *0101

PTPN22: European

STAT4: North American
TNFAIP3: North American
TRAF1/CF: North American
CTLA4: European

Asian ancestry:

HLA-DRB1:

- *0401 (East Asian)
- *0405
- *0901 (Japanese, Malaysian, Korean)

PADI4

CD244

Other:

CD40

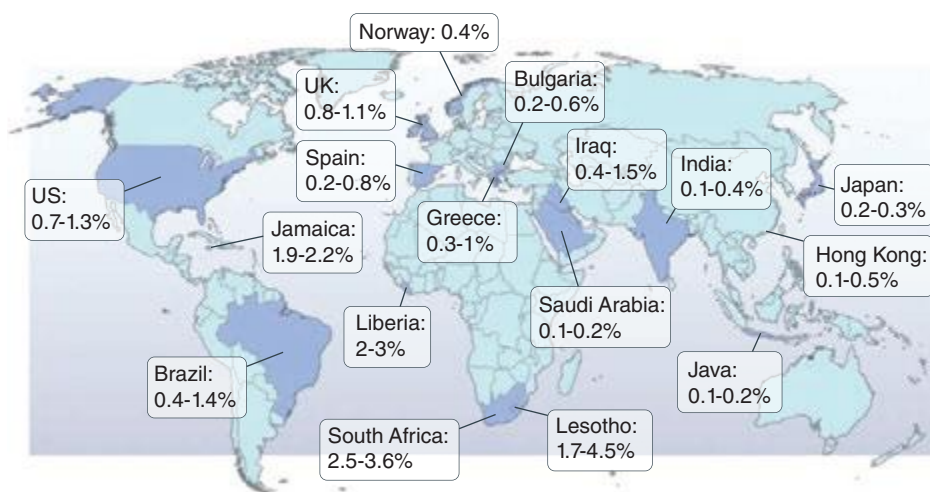


FIGURE 380-3 Global prevalence rates of rheumatoid arthritis (RA) with genetic associations. Listed are the major genetic alleles associated with RA. Although human leukocyte antigen (HLA)-DRB1 mutations are found globally, some alleles have been associated with RA in only certain ethnic groups.

2140 effect on risk; they also contribute to the risk for developing other autoimmune diseases, such as type 1 diabetes mellitus, systemic lupus erythematosus, and multiple sclerosis. Second, although most of the non-HLA associations are described in patients with anti-CCP antibody-positive disease, there are several risk loci that are unique to anti-CCP antibody-negative disease. Third, risk alleles vary among ethnic groups. And fourth, the risk loci mostly reside in genes encoding proteins involved in the regulation of the immune response. However, the risk alleles identified by GWAS only account at present for approximately 5% of the genetic risk, suggesting that rare variants or other classes of DNA variants, such as variants in copy number, may be yet found that significantly contribute to the overall risk model.

Recently, imputation of SNP data from a GWAS meta-analysis shows amino acid substitutions in the MHC locus independently associated with the risk for RA are at position 11, 71, and 74 in HLA-DR β 1, position 9 of HLA-B, and position 9 of HLA-DP β 1. The amino acids at position 11, 71, and 74 are located in the antigen-binding groove of the HLA-DR β 1 molecule, highlighting positions 71 and 74 that form part of the original shared epitope.

Among the best examples of the non-MHC genes contributing to the risk of RA is the gene encoding protein tyrosine phosphatase non-receptor 22 (*PTPN22*). This gene varies in frequency among patients from different parts of Europe (e.g., 3–10%), but is absent in patients of East Asian ancestry. *PTPN22* encodes lymphoid tyrosine phosphatase, a protein that regulates T and B cell function. Inheritance of the risk allele for *PTPN22* produces a gain-of-function in the protein that is hypothesized to result in the abnormal thymic selection of autoreactive T and B cells and appears to be associated exclusively with anti-CCP-positive disease. The peptidyl arginine deiminase type IV (*PADI4*) gene is another risk allele that encodes an enzyme involved in the conversion of arginine to citrulline and is postulated to play a role in the development of antibodies to citrullinated antigens. A polymorphism in *PADI4* has been associated with RA only in Asian populations.

Epigenetics is the study of heritable traits that affect gene expression but do not modify DNA sequence. It may provide a link between environmental exposure and predisposition to disease. The best-studied mechanisms include posttranslational histone modifications and DNA methylation. Although studies of epigenetic phenomena are limited, DNA methylation patterns have been shown to differ between RA patients and healthy controls, as well as patients with osteoarthritis.

ENVIRONMENTAL FACTORS

In addition to genetic predisposition, a host of environmental factors have been implicated in the pathogenesis of RA. The most reproducible of these environmental links is cigarette smoking. Numerous cohort and case control studies have demonstrated that smoking confers a relative risk for developing RA of 1.5–3.5. In particular, women who smoke cigarettes have a nearly 2.5 times greater risk of RA, a risk that persists even 15 years after smoking cessation. A twin who smokes will have a significantly higher risk for RA than his or her monozygotic co-twin, theoretically with the same genetic risk, who does not smoke. Interestingly, the risk from smoking is almost exclusively related to RF and anti-CCP antibody-positive disease. However, it has not been shown that smoking cessation, while having many health benefits, improves disease activity.

Researchers began to aggressively seek an infectious etiology for RA after the discovery in 1931 that sera from patients with this disease could agglutinate strains of streptococci. Certain viruses such as Epstein-Barr virus (EBV) have garnered the most interest over the past 30 years given their ubiquity, ability to persist for many years in the host, and frequent association with arthritic complaints. For example, titers of IgG antibodies against EBV antigens in the peripheral blood and saliva are significantly higher in patients with RA than the general population. EBV DNA has also been found in synovial fluid and synovial cells of RA patients. Because the evidence for these links is largely circumstantial, it has not been possible to directly implicate infection as a causative factor in RA.

PATHOLOGY

RA affects the synovial tissue and underlying cartilage and bone. The synovial membrane, which covers most articular surfaces, tendon sheaths, and bursae, normally is a thin layer of connective tissue. In joints, it faces the bone and cartilage, bridging the opposing bony surfaces and inserting at periosteal regions close to the articular cartilage. It consists primarily of two cell types—type A synoviocytes (macrophage-derived) and type B synoviocytes (fibroblast-derived). The synovial fibroblasts are the most abundant and produce the structural components of joints, including collagen, fibronectin, and laminin, as well as other extracellular constituents of the synovial matrix. The sublining layer consists of blood vessels and a sparse population of mononuclear cells within a loose network of connective tissue. Synovial fluid, an ultrafiltrate of blood, diffuses through the subsynovial lining tissue across the synovial membrane and into the joint cavity. Its main constituents are hyaluronan and lubricin. Hyaluronan is a glycosaminoglycan that contributes to the viscous nature of synovial fluid, which along with lubricin, lubricates the surface of the articular cartilage.

The pathologic hallmarks of RA are synovial inflammation and proliferation, focal bone erosions, and thinning of articular cartilage. Chronic inflammation leads to synovial lining hyperplasia and the formation of pannus, a thickened cellular membrane containing fibroblast-like synoviocytes and granulation-reactive fibrovascular tissue that invades the underlying cartilage and bone. The inflammatory infiltrate is made up of no less than six cell types: T cells, B cells, plasma cells, dendritic cells, mast cells, and, to a lesser extent, granulocytes. The T cells comprise 30–50% of the infiltrate, with the other cells accounting for the remainder. The topographical organization of these cells is complex and may vary among individuals with RA. Most often, the lymphocytes are diffusely organized among the tissue resident cells; however, in some cases, the B cells, T cells, and dendritic cells may form higher levels of organization, such as lymphoid follicles and germinal center-like structures. Growth factors secreted by synovial fibroblasts and macrophages promote the formation of new blood vessels in the synovial sublining that supply the increasing demands for oxygenation and nutrition required by the infiltrating leukocytes and expanding synovial tissue.

The structural damage to the mineralized cartilage and subchondral bone is mediated by the osteoclast. Osteoclasts are multinucleated giant cells that can be identified by their expression of CD68, tartrate-resistant acid phosphatase, cathepsin K, and the calcitonin receptor. They appear at the pannus-bone interface where they eventually form resorption lacunae. These lesions typically localize where the synovial membrane inserts into the periosteal surface at the edges of bones close to the rim of articular cartilage and at the attachment sites of ligaments and tendon sheaths. This process most likely explains why bone erosions usually develop at the radial sites of the MCP joints juxtaposed to the insertion sites of the tendons, collateral ligaments, and synovial membrane. Another form of bone loss is periarticular osteopenia that occurs in joints with active inflammation. It is associated with substantial thinning of the bony trabeculae along the metaphyses of bones, and likely results from inflammation of the bone marrow cavity. These lesions can be visualized on MRI scans, where they appear as signal alterations in the bone marrow adjacent to inflamed joints. Their signal characteristics show they are water-rich with a low fat content and are consistent with highly vascularized inflammatory tissue. These bone marrow lesions are often the forerunner of bone erosions.

The cortical bone layer that separates the bone marrow from the invading pannus is relatively thin and susceptible to penetration by the inflamed synovium. The bone marrow lesions seen on MRI scans are associated with an endosteal bone response characterized by the accumulation of osteoblasts and deposition of osteoid. Thus, in recent years, the concept of joint pathology in RA has been extended to include the bone marrow cavity. Finally, generalized osteoporosis, which results in the thinning of trabecular bone throughout the body, is a third form of bone loss found in patients with RA.

Articular cartilage is an avascular tissue comprised of a specialized matrix of collagens, proteoglycans, and other proteins. It is organized

in four distinct regions (superficial, middle, deep, and calcified cartilage zones)—chondrocytes constitute the unique cellular component in these layers. Originally, cartilage was considered to be an inert tissue, but it is now known to be a highly responsive tissue that reacts to inflammatory mediators and mechanical factors, which in turn, alter the balance between cartilage anabolism and catabolism. In RA, the initial areas of cartilage degradation are juxtaposed to the synovial pannus. The cartilage matrix is characterized by a generalized loss of proteoglycan, most evident in the superficial zones adjacent to the synovial fluid. Degradation of cartilage may also take place in the perichondrocytic zone and in regions adjacent to the subchondral bone.

PATHOGENESIS

The pathogenic mechanisms of synovial inflammation are likely to result from a complex interplay of genetic, environmental, and immunologic factors that produces dysregulation of the immune system and a breakdown in self-tolerance (Fig. 380-4). Precisely what triggers these initiating events and what genetic and environmental factors disrupt the immune system remains a mystery. However, a detailed molecular picture is emerging of the mechanisms underlying the chronic inflammatory response and the destruction of the articular cartilage and bone.

In RA, the preclinical stage appears to be characterized by a breakdown in self-tolerance. This idea is supported by the finding that autoantibodies, such as RF and anti-CCP antibodies, may be found in sera from patients many years before clinical disease can be detected. However, the antigenic targets of anti-CCP antibodies and RF are not restricted to the joint, and their role in disease pathogenesis remains speculative. Anti-CCP antibodies are directed against deaminated peptides, which result from posttranslational modification by the enzyme PADI4. They recognize citrulline-containing regions of several different matrix proteins, including filaggrin, keratin, fibrinogen, and vimentin, and are present at higher levels in the joint fluid compared to the serum. Other autoantibodies have been found in a minority of patients with RA, but they also occur in the setting of other types of arthritis. They bind to a diverse array of autoantigens, including type II collagen, human cartilage gp-39, aggrecan, calpastatin, BiP (immunoglobulin binding protein), and glucose-6-phosphate isomerase.

In theory, environmental stimulants may synergize with other factors to bring about inflammation in RA. People who smoke display higher citrullination of proteins in bronchoalveolar fluid than those who do not smoke. Thus, it has been speculated that long-term exposure to tobacco smoke might induce citrullination of cellular proteins in the lung and stimulate the expression of a neopeptide capable of inducing self-reactivity, which in turns, leads to formation of immune complexes and joint inflammation. Exposure to silicone dust and mineral oil, which has adjuvant effects, has also been linked to an increased risk for anti-CCP antibody-positive RA.

How might microbes or their products be involved in the initiating events of RA? The immune system is alerted to the presence of microbial infections through Toll-like receptors (TLRs). There are 10 TLRs in humans that recognize a variety of microbial products, including bacterial cell-surface lipopolysaccharides and heat-shock proteins (TLR4), lipoproteins (TLR2), double-strand RNA viruses (TLR3), and unmethylated CpG DNA from bacteria (TLR9). TLR2, -3, and -4 are abundantly expressed by synovial fibroblasts in early RA and, when bound by their ligands, upregulate production of proinflammatory cytokines. Although such events could amplify inflammatory pathways in RA, a specific role for TLRs in disease pathogenesis has not been elucidated.

The pathogenesis of RA is built upon the concept that self-reactive T cells drive the chronic inflammatory response. In theory, self-reactive T cells might arise in RA from abnormal central (thymic) selection due to defects in DNA repair leading to an imbalance of T cell death and life, or defects in the cell signaling apparatus lowering the threshold for T cell activation. Similarly, abnormal selection of the T cell repertoire in the periphery might lead to a breakdown in T cell tolerance. The support for these theories comes mainly from studies of arthritis

in mouse models. It has not been shown that patients with RA have abnormal thymic selection of T cells or defective apoptotic pathways regulating cell death. At least some antigen stimulation inside the joint seems likely, owing to the fact that T cells in the synovium express a cell-surface phenotype indicating prior antigen exposure and show evidence of clonal expansion. Of interest, peripheral blood T cells from patients with RA have been shown to display a fingerprint of premature aging that mostly affects inexperienced naïve T cells. In these studies, the most glaring findings have been the loss of telomeric sequences and a decrease in the thymic output of new T cells. Although intriguing, it is not clear how generalized T cell abnormalities might provoke a systemic disease dominated by synovitis.

There is substantial evidence supporting a role for CD4+ T cells in the pathogenesis of RA. First, the co-receptor CD4 on the surface of T cells binds to invariant sites on MHC class II molecules, stabilizing the MHC-peptide-T cell receptor complex during T cell activation. Because the SE on MHC class II molecules is a risk factor for RA, it follows that CD4+ T cell activation may play a role in the pathogenesis of this disease. Second, CD4+ memory T cells are enriched in the synovial tissue from patients with RA and can be implicated through “guilt by association.” Third, CD4+ T cells have been shown to be important in the initiation of arthritis in animal models. Fourth, some, but not all, T cell-directed therapies have shown clinical efficacy in this disease. Taken together, these lines of evidence suggest that CD4+ T cells play an important role in orchestrating the chronic inflammatory response in RA. However, other cell types, such as CD8+ T cells, natural killer (NK) cells, and B cells are present in synovial tissue and may also influence pathogenic responses.

In the rheumatoid joint, by mechanisms of cell-cell contact and release of soluble mediators, activated T cells stimulate macrophages and fibroblast-like synoviocytes to generate proinflammatory mediators and proteases that drive the synovial inflammatory response and destroy the cartilage and bone. CD4+ T cell activation is dependent on two signals: (1) T cell receptor binding to peptide-MHC on antigen-presenting cells; and (2) CD28 binding to CD80/86 on antigen-presenting cells. CD4+ T cells also provide help to B cells, which in turn, produce antibodies that may promote further inflammation in the joint. The previous T cell-centric model for the pathogenesis of RA was based on a T_H1 -driven paradigm, which came from studies indicating that CD4+ T helper (T_H) cells differentiated into T_H1 and T_H2 subsets, each with their distinctive cytokine profiles. T_H1 cells were found to mainly produce interferon γ (IFN- γ), lymphotoxin β , and TNF- α , whereas T_H2 cells predominantly secreted interleukin (IL)-4, IL-5, IL-6, IL-10, and IL-13. The recent discovery of another subset of T_H cells, namely the T_H17 lineage, has revolutionized our concepts concerning the pathogenesis of RA. In humans, naïve T cells are induced to differentiate into T_H17 cells by exposure to transforming growth factor β (TGF- β), IL-1, IL-6, and IL-23. Upon activation, T_H17 cells secrete a variety of proinflammatory mediators such as IL-17, IL-21, IL-22, TNF- α , IL-26, IL-6, and granulocyte-macrophage colony-stimulating factor (GM-CSF). Substantial evidence now exists from both animal models and humans that IL-17 plays an important role not only in promoting joint inflammation, but also in destroying cartilage and subchondral bone.

The immune system has evolved mechanisms to counterbalance the potential harmful immune-mediated inflammatory responses provoked by infectious agents and other triggers. Among these negative regulators are regulatory T (T_{reg}) cells, which are produced in the thymus and induced in the periphery to suppress immune-mediated inflammation. They are characterized by the surface expression of CD25 and the transcription factor forkhead box P3 (FOXP3) and orchestrate dominant tolerance through contact with other immune cells and secretion of inhibitory cytokines, such as TGF- β , IL-10, and IL-35. T_{reg} cells appear to be heterogeneous and capable of suppressing distinct classes (T_H1 , T_H2 , T_H17) of the immune response. In RA, the data that T_{reg} numbers are deficient compared to normal healthy controls are contradictory and inconclusive. Although some experimental evidence suggests that T_{reg} suppressive activity is lost due to dysfunctional expression of cytotoxic T lymphocyte antigen 4

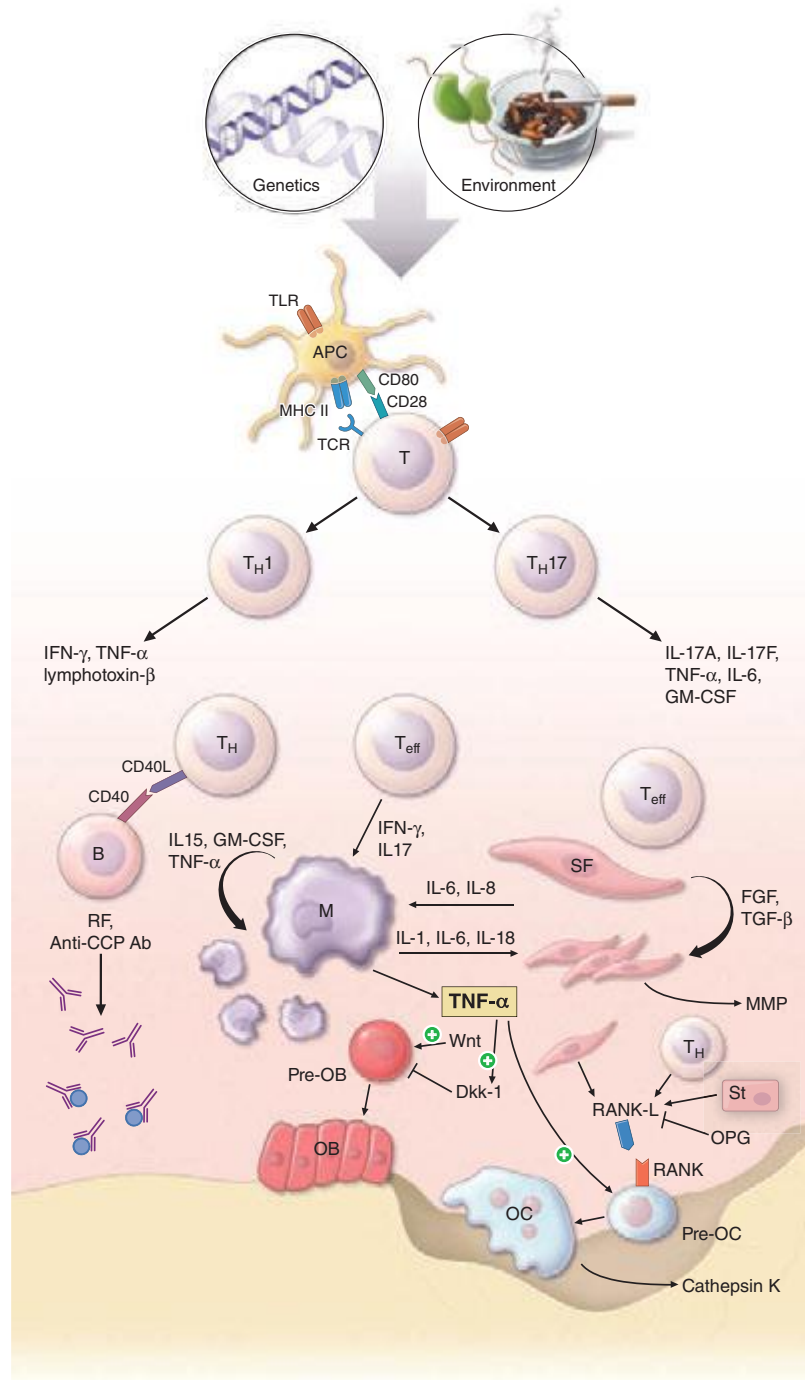


FIGURE 380-4 Pathophysiologic mechanisms of inflammation and joint destruction. Genetic predisposition along with environmental factors may trigger the development of rheumatoid arthritis (RA), with subsequent synovial T cell activation. CD4+ T cells become activated by antigen-presenting cells (APCs) through interactions between the T cell receptor and class II major histocompatibility complex (MHC)-peptide antigen (signal 1) with co-stimulation through the CD28-CD80/86 pathway, as well as other pathways (signal 2). In theory, ligands binding Toll-like receptors (TLRs) may further stimulate activation of APCs inside the joint. Synovial CD4+ T cells differentiate into T_H1 and T_H17 cells, each with their distinctive cytokine profile. CD4+ T_H cells in turn activate B cells, some of which are destined to differentiate into autoantibody-producing plasma cells. Immune complexes, possibly comprised of rheumatoid factors (RFs) and anti-cyclic citrullinated peptides (CCP) antibodies, may form inside the joint, activating the complement pathway and amplifying inflammation. T effector cells stimulate synovial macrophages (M) and fibroblasts (SF) to secrete proinflammatory mediators, among which is tumor necrosis factor α (TNF- α). TNF- α upregulates adhesion molecules on endothelial cells, promoting leukocyte influx into the joint. It also stimulates the production of other inflammatory mediators, such as interleukin 1 (IL-1), IL-6, and granulocyte-macrophage colony-stimulating factor (GM-CSF). TNF- α has a critically important function in regulating the balance between bone destruction and formation. It upregulates the expression of dickkopf-1 (DKK-1), which can then internalize Wnt receptors on osteoblast precursors. Wnt is a soluble mediator that promotes osteoblastogenesis and bone formation. In RA, bone formation is inhibited through the Wnt pathway, presumably due to the action of elevated levels of DKK-1. In addition to inhibiting bone formation, TNF- α stimulates osteoclastogenesis. However, it is not sufficient by itself to induce the differentiation of osteoclast precursors (Pre-OC) into activated osteoclasts capable of eroding bone. Osteoclast differentiation requires the presence of macrophage colony-stimulating factor (M-CSF) and receptor activator of nuclear factor- κ B (RANK) ligand (RANKL), which binds to RANK on the surface of Pre-OC. Inside the joint, RANKL is mainly derived from stromal cells, synovial fibroblasts, and T cells. Osteoprotegerin (OPG) acts as a decoy receptor for RANKL, thereby inhibiting osteoclastogenesis and bone loss. FGF, fibroblast growth factor; IFN, interferon; TGF, transforming growth factor.

(CTLA-4), the nature of T_{reg} defects in RA, if they exist, remains unclear.

Cytokines, chemokines, antibodies, and endogenous danger signals bind to receptors on the surface of immune cells and stimulate a cascade of intracellular signaling events that can amplify the inflammatory response. Signaling molecules and their binding partners in these pathways are the target of small-molecule drugs designed to interfere with signal transduction and block these reinforcing inflammatory loops. Examples of signal molecules in these critical inflammatory pathways include Janus kinase (JAK)/signal transducers and activators of transcription (STAT), spleen tyrosine kinase (Syk), mitogen-activated protein kinases (MAPKs), and nuclear factor- κ B (NF- κ B). These pathways exhibit significant cross-talk and are found in many cell types. Some signal transducers, such as JAK3, are primarily expressed in hematopoietic cells and play an important role in the inflammatory response in RA.

Activated B cells are also important players in the chronic inflammatory response. B cells give rise to plasma cells, which in turn, produce antibodies, including RF and anti-CCP antibodies. RFs may form large immune complexes inside the joint that contribute to the pathogenic process by fixing complement and promoting the release of proinflammatory chemokines and chemoattractants. In mouse models of arthritis, RF-containing immune complexes and anti-CCP-containing immune complexes synergize with other mechanisms to exacerbate the synovial inflammatory response.

RA is often considered to be a macrophage-driven disease because this cell type is the predominant source of proinflammatory cytokines inside the joint. Key proinflammatory cytokines released by synovial macrophages include TNF- α , IL-1, IL-6, IL-12, IL-15, IL-18, and IL-23. Synovial fibroblasts, the other major cell type in this microenvironment, produce the cytokines IL-1 and IL-6 as well as TNF- α . TNF- α is a pivotal cytokine in the pathobiology of synovial inflammation. It upregulates adhesion molecules on endothelial cells, promoting the influx of leukocytes into the synovial microenvironment; activates synovial fibroblasts; stimulates angiogenesis; promotes pain receptor sensitizing pathways; and drives osteoclastogenesis. Fibroblasts secrete matrix metalloproteinases (MMPs) as well as other proteases that are chiefly responsible for the breakdown of articular cartilage.

Osteoclast activation at the site of the pannus is closely tied to the presence of focal bone erosion. Receptor activator of nuclear factor- κ B ligand (RANKL) is expressed by stromal cells, synovial fibroblasts, and T cells. Upon binding to its receptor RANK on osteoclast progenitors, RANKL stimulates osteoclast differentiation and bone resorption. RANKL activity is regulated by osteoprotegerin (OPG), a decoy receptor of RANKL that blocks osteoclast formation. Monocytic cells in the synovium serve as the precursors of osteoclasts and, when exposed to macrophage colony-stimulating factor (M-CSF) and RANKL, fuse to form polykaryons termed *preosteoclasts*. These precursor cells undergo further differentiation into osteoclasts with the characteristic ruffled membrane. Cytokines such as TNF- α , IL-1, IL-6, and IL-17 increase the expression of RANKL in the joint and thus promote osteoclastogenesis. Osteoclasts also secrete cathepsin K, which is a cysteine protease that degrades the bone matrix by cleaving collagen. Stimulation of osteoclasts also contributes to generalized bone loss and osteoporosis.

Increased bone loss is only part of the story in RA, as decreased bone formation plays a crucial role in bone remodeling at sites of inflammation. Recent evidence shows that inflammation suppresses bone formation. The proinflammatory cytokine TNF- α plays a key role in actively suppressing bone formation by enhancing the expression of dickkopf-1 (DKK-1). DKK-1 is an important inhibitor of the Wnt pathway, which acts to promote osteoblast differentiation and bone formation. The Wnt system is a family of soluble glycoproteins that bind to cell-surface receptors known as frizzled (*fz*) and low-density lipoprotein (LDL) receptor-related proteins (LRPs) and promote cell growth. In animal models, increased levels of DKK-1 are associated with decreased bone formation, whereas inhibition of DKK-1 protects against structural damage in the joint. Wnt proteins also induce the formation of OPG and thereby shut down bone resorption, emphasizing their key role in tightly regulating the balance between bone resorption and formation.

DIAGNOSIS

The clinical diagnosis of RA is largely based on signs and symptoms of a chronic inflammatory arthritis, with laboratory and radiographic results providing important supplemental information. In 2010, a collaborative effort between the American College of Rheumatology (ACR) and the European League Against Rheumatism (EULAR) revised the 1987 ACR classification criteria for RA in an effort to improve early diagnosis with the goal of identifying patients who would benefit from early introduction of disease-modifying therapy (Table 380-1). Application of the newly revised criteria yields a score of 0–10, with a score of ≥ 6 fulfilling the requirements for definite RA. The new classification criteria differ in several ways from the older criteria set. The new criteria include a positive test for serum anti-CCP antibodies (also termed ACPA, anti-citrullinated peptide antibodies) as an item, which carries greater specificity for the diagnosis of RA than a positive test for RF. The newer classification criteria also do not take into account whether the patient has rheumatoid nodules or radiographic joint damage because these findings occur rarely in early RA. It is important to emphasize that the new 2010 ACR-EULAR criteria are “classification criteria” as opposed to “diagnostic criteria” and serve to distinguish patients at the onset of disease who have a high likelihood of evolution to chronic disease with persistent synovitis and joint damage. The presence of radiographic joint erosions or subcutaneous nodules may inform the diagnosis in the later stages of the disease.

LABORATORY FEATURES

Patients with systemic inflammatory diseases such as RA will often present with elevated nonspecific inflammatory markers such as an ESR or CRP. Detection of serum RF and anti-CCP antibodies is important in differentiating RA from other polyarticular diseases, although RF lacks diagnostic specificity and may be found in association with other chronic inflammatory diseases in which arthritis figures in the clinical manifestations.

IgM, IgG, and IgA isotypes of RF occur in sera from patients with RA, although the IgM isotype is the one most frequently measured by commercial laboratories. Serum IgM RF has been found in 75–80% of patients with RA; therefore, a negative result does not exclude the presence of this disease. It is also found in other connective tissue diseases, such as primary Sjögren’s syndrome, systemic lupus erythematosus, and type II mixed essential cryoglobulinemia, as well as chronic infections such as subacute bacterial endocarditis and hepatitis B and C. Serum RF may also be detected in 1–5% of the healthy population.

TABLE 380-1 CLASSIFICATION CRITERIA FOR RHEUMATOID ARTHRITIS

		Score
Joint involvement	1 large joint (shoulder, elbow, hip, knee, ankle)	0
	2–10 large joints	1
	1–3 small joints (MCP, PIP, thumb IP, MTP, wrists)	2
	4–10 small joints	3
	>10 joints (at least 1 small joint)	5
Serology	Negative RF and negative ACPA	0
	Low-positive RF or low-positive anti-CCP antibodies (≤ 3 times ULN)	2
	High-positive RF or high-positive anti-CCP antibodies (>3 times ULN)	3
Acute-phase reactants	Normal CRP and normal ESR	0
	Abnormal CRP or abnormal ESR	1
Duration of symptoms	<6 weeks	0
	≥ 6 weeks	1

Note: These criteria are aimed at classification of newly presenting patients who have at least one joint with definite clinical synovitis that is not better explained by another disease. A score of ≥ 6 fulfills requirements for definite RA.

Abbreviations: ACPA, anti-citrullinated peptide antibodies; CCP, cyclic citrullinated peptides; CRP, C-reactive protein; ESR, erythrocyte sedimentation rate; IP, interphalangeal joint; MCP, metacarpophalangeal joint; MTP, metatarsophalangeal joint; PIP, proximal interphalangeal joint; RF, rheumatoid factor; ULN, upper limit of normal.

Source: D Aletaha et al: Arthritis Rheum 62:2569, 2010.

2144 The presence of serum anti-CCP antibodies has about the same sensitivity as serum RF for the diagnosis of RA. However, its diagnostic specificity approaches 95%, so a positive test for anti-CCP antibodies in the setting of an early inflammatory arthritis is useful for distinguishing RA from other forms of arthritis. There is some incremental value in testing for the presence of both RF and anti-CCP, as some patients with RA are positive for RF but negative for anti-CCP and visa versa. The presence of RF or anti-CCP antibodies also has prognostic significance, with anti-CCP antibodies showing the most value for predicting worse outcomes.

SYNOVIAL FLUID ANALYSIS

Typically, synovial fluid from patients with RA reflects an inflammatory state. Synovial fluid white blood cell (WBC) counts can vary widely, but generally range between 5000 and 50,000 WBC/ μ L compared to <2000 WBC/ μ L for a noninflammatory condition such as osteoarthritis. In contrast to the synovial tissue, the overwhelming cell type in the synovial fluid is the neutrophil. Clinically, the analysis of synovial fluid is most useful for confirming an inflammatory arthritis (as opposed to osteoarthritis), while at the same time excluding infection or a crystal-induced arthritis such as gout or pseudogout (**Chap. 395**).

JOINT IMAGING

Joint imaging is a valuable tool not only for diagnosing RA, but also for tracking progression of any joint damage. Plain x-ray is the most common imaging modality, but it is limited to visualization of the bony structures and inferences about the state of the articular cartilage based on the amount of narrowing of the joint space. MRI and ultrasound techniques offer the added value of detecting changes in the soft tissues such as synovitis, tenosynovitis, and effusions as well as greater sensitivity for identifying bony abnormalities. Plain radiographs are usually relied upon in clinical practice for the purpose of diagnosis and monitoring of affected joints. However, in selected cases, MRI and ultrasound can provide additional diagnostic information that may guide clinical decision making. Musculoskeletal ultrasound with power Doppler is increasingly used in rheumatology clinical practice for detecting synovitis and bone erosion.

Plain Radiography Classically in RA, the initial radiographic finding is periarticular osteopenia. Practically speaking, however, this finding is difficult to appreciate on plain films and, in particular, on the newer digitalized x-rays. Other findings on plain radiographs include soft tissue swelling, symmetric joint space loss, and subchondral erosions, most frequently in the wrists and hands (MCPs and PIPs) and the feet (MTPs). In the feet, the lateral aspect of the fifth MTP is often targeted first, but other MTP joints may be involved at the same time. X-ray imaging of advanced RA may reveal signs of severe destruction, including joint subluxation and collapse (**Fig. 380-5**).



FIGURE 380-5 X-ray demonstrating progression of erosions on the proximal interphalangeal joint. (Courtesy of the American College of Rheumatology.)

MRI MRI offers the greatest sensitivity for detecting synovitis and joint effusions, as well as early bone and bone marrow changes. These soft tissue abnormalities often occur before osseous changes are noted on x-ray. Presence of bone marrow edema has been recognized to be an early sign of inflammatory joint disease and can predict the subsequent development of erosions on plain radiographs as well as MRI scans. Cost and availability of MRI are the main factors limiting its routine clinical use.

Ultrasound Ultrasound, including power color Doppler, has the ability to detect more erosions than plain radiography, especially in easily accessible joints. It can also reliably detect synovitis, including increased joint vascularity indicative of inflammation. The usefulness of ultrasound is dependent on the experience of the sonographer; however, it does offer the advantages of portability, lack of radiation, and low expense relative to MRI, factors that make it attractive as a clinical tool.

CLINICAL COURSE

The natural history of RA is complex and affected by a number of factors including age of onset, gender, genotype, phenotype (i.e., extraarticular manifestations or variants of RA), and comorbid conditions, which make for a truly heterogeneous disease. There is no simple way to predict the clinical course. It is important to realize that as many as 10% of patients with inflammatory arthritis fulfilling ACR classification criteria for RA will undergo a spontaneous remission within 6 months (particularly seronegative patients). However, the vast majority of patients will exhibit a pattern of persistent and progressive disease activity that waxes and wanes in intensity over time. A minority of patients will show intermittent and recurrent explosive attacks of inflammatory arthritis interspersed with periods of disease quiescence. Finally, an aggressive form of RA may occur in an unfortunate few with inexorable progression of severe erosive joint disease, although this highly destructive course is less common in the modern treatment era of biologics.

Disability, as measured by the Health Assessment Questionnaire (HAQ), shows gradual worsening of disability over time in the face of poorly controlled disease activity and disease progression. Disability may result from both a disease activity-related component that is potentially reversible with therapy and a joint damage-related component owing to the cumulative and largely irreversible effects of cartilage and bone breakdown. Early in the course of disease, the extent of joint inflammation is the primary determinant of disability, while in the later stages of disease, the amount of joint damage is the dominant contributing factor. Previous studies have shown that more than one-half of patients with RA are unable to work 10 years after the onset of their disease; however, increased employability and less work absenteeism has been reported recently with the use of newer therapies and earlier treatment intervention.

The overall mortality rate in RA is two times greater than the general population, with ischemic heart disease being the most common cause of death followed by infection. Median life expectancy is shortened by an average of 7 years for men and 3 years for women compared to control populations. Patients at higher risk for shortened survival are those with systemic extraarticular involvement, low functional capacity, low socioeconomic status, low education, and chronic prednisone use.

TREATMENT RHEUMATOID ARTHRITIS

The amount of clinical disease activity in patients with RA reflects the overall burden of inflammation and is the variable most influencing treatment decisions. Joint inflammation is the main driver of joint damage and is the most important cause of functional disability in the early stages of disease. Several composite indices have been developed to assess clinical disease activity. The ACR 20, 50, and 70 improvement criteria (which corresponds to a 20%, 50%, and 70% improvement, respectively, in joint counts, physician/patient assessment of disease severity, pain scale, serum levels of acute-phase

reactants [ESR or CRP], and a functional assessment of disability using a self-administered patient questionnaire) are a composite index with a dichotomous response variable. The ACR improvement criteria are commonly used in clinical trials as an endpoint for comparing the proportion of responders between treatment groups. In contrast, the Disease Activity Score (DAS), Simplified Disease Activity Index (SDAI), and the Clinical Disease Activity Index (CDAI) are continuous measures of disease activity. These scales are increasingly used in clinical practice for tracking disease status and, in particular, for documenting treatment response.

Several developments during the past two decades have changed the therapeutic landscape in RA. They include (1) the emergence of methotrexate as the disease-modifying antirheumatic drug (DMARD) of first choice for the treatment of early RA; (2) the development of novel highly efficacious biologicals that can be used alone or in combination with methotrexate; and (3) the proven superiority of combination DMARD regimens over methotrexate alone. The medications used for the treatment of RA may be divided into broad categories: nonsteroidal anti-inflammatory drugs (NSAIDs); glucocorticoids, such as prednisone and methylprednisolone; conventional DMARDs; and biologic DMARDs (Table 380-2). Although disease for some patients with RA is managed adequately with a single DMARD, such as methotrexate, the situation in most cases demands the use of a combination DMARD regimen that may vary in its components over the treatment course depending on fluctuations in disease activity and emergence of drug-related toxicities and comorbidities.

NSAIDs

NSAIDs were formerly viewed as the core of all other RA therapy, but they are now considered to be adjunctive therapy for management of symptoms uncontrolled by other measures. NSAIDs exhibit both analgesic and anti-inflammatory properties. The anti-inflammatory effects of NSAIDs derive from their ability to nonselectively inhibit cyclooxygenase (COX)-1 and COX-2. Although the results of clinical trials suggest NSAIDs are roughly equivalent in their efficacy, experience suggests that some individuals may preferentially respond to a particular NSAID. Chronic use should be minimized due to the possibility of side effects, including gastritis and peptic ulcer disease as well as impairment of renal function.

GLUCOCORTICOIDS

Glucocorticoids may serve in several ways to control disease activity in RA. First, they may be administered in low to moderate doses to achieve rapid disease control before the onset of fully effective DMARD therapy, which often takes several weeks or even months. Second, a 1- to 2-week burst of glucocorticoids may be prescribed for the management of acute disease flares, with dose and duration guided by the severity of the exacerbation. Chronic administration of low doses (5–10 mg/d) of prednisone (or its equivalent) may also be warranted to control disease activity in patients with an inadequate response to DMARD therapy. Low-dose prednisone therapy has been shown in prospective studies to retard radiographic progression of joint disease; however, the benefits of this approach must be carefully weighed against the risks. Best practices minimize chronic use of low-dose prednisone therapy owing to the risk of osteoporosis and other long-term complications; however, the use of chronic prednisone therapy is unavoidable in many cases. High-dose glucocorticoids may be necessary for treatment of severe extraarticular manifestations of RA, such as ILD. Finally, if a patient exhibits one or a few actively inflamed joints, the clinician may consider intraarticular injection of an intermediate-acting glucocorticoid such as triamcinolone acetonide. This approach may allow for rapid control of inflammation in the setting of a limited number of affected joints. Caution must be exercised to appropriately exclude joint infection, as it often mimics an RA flare.

Osteoporosis ranks as an important long-term complication of chronic prednisone use. The ACR recommends primary prevention of glucocorticoid-induced osteoporosis with a bisphosphonate in

any patient receiving 5 mg/d or more of prednisone for greater than 3 months. Although prednisone use is known to increase the risk of peptic ulcer disease, especially with concomitant NSAID use, no evidence-based guidelines have been published regarding the use of gastrointestinal ulcer prophylaxis in this situation.

DMARDs

DMARDs are so named because of their ability to slow or prevent structural progression of RA. The conventional DMARDs include hydroxychloroquine, sulfasalazine, methotrexate, and leflunomide; they exhibit a delayed onset of action of approximately 6–12 weeks. Methotrexate is the DMARD of choice for the treatment of RA and is the anchor drug for most combination therapies. It was approved for the treatment of RA in 1986 and remains the benchmark for the efficacy and safety of new disease-modifying therapies. At the dosages used for the treatment of RA, methotrexate has been shown to stimulate adenosine release from cells, producing an anti-inflammatory effect. The clinical efficacy of leflunomide, an inhibitor of pyrimidine synthesis, appears similar to that of methotrexate; it has been shown in well-designed trials to be effective for the treatment of RA as monotherapy or in combination with methotrexate and other DMARDs.

Although similar to the other DMARDs in its slow onset of action, hydroxychloroquine has not been shown to delay radiographic progression of disease and thus is not considered to be a true DMARD. In clinical practice, hydroxychloroquine is generally used for treatment of early, mild disease or as adjunctive therapy in combination with other DMARDs. Sulfasalazine is used in a similar manner and has been shown in randomized, controlled trials to reduce radiographic progression of disease. Minocycline, gold salts, penicillamine, azathioprine, and cyclosporine have all been used for the treatment of RA with varying degrees of success; however, they are used sparingly now due to their inconsistent clinical efficacy or unfavorable toxicity profile.

BIOLOGICALS

Biologic DMARDs have revolutionized the treatment of RA over the past decade (Table 380-2). They are protein therapeutics designed mostly to target cytokines and cell-surface molecules. The TNF inhibitors were the first biologicals approved for the treatment of RA. Anakinra, an IL-1 receptor antagonist, was approved shortly thereafter; however, its benefits have proved to be relatively modest compared with the other biologicals and is rarely used for the treatment of RA with the availability of other more effective agents. Abatacept, rituximab, and tocilizumab are the newest members of this class.

Anti-TNF Agents The development of TNF inhibitors was originally spurred by the experimental finding that TNF is a critical upstream mediator of joint inflammation. Currently, five agents that inhibit TNF- α are approved for the treatment of RA. There are three different anti-TNF monoclonal antibodies. Infliximab is a chimeric (part mouse and human) monoclonal antibody, whereas adalimumab and golimumab are humanized monoclonal antibodies. Certolizumab pegol is a pegylated Fc-free fragment of a humanized monoclonal antibody with binding specificity for TNF- α . Lastly, etanercept is a soluble fusion protein comprising the TNF receptor 2 in covalent linkage with the Fc portion of IgG1. All of the TNF inhibitors have been shown in randomized controlled clinical trials to reduce the signs and symptoms of RA, slow radiographic progression of joint damage, and improve physical function and quality of life. Anti-TNF drugs are typically used in combination with background methotrexate therapy. This combination regimen, which affords maximal benefit in many cases, is often the next step for treatment of patients with an inadequate response to methotrexate therapy. Etanercept, adalimumab, certolizumab pegol, and golimumab have also been approved for use as monotherapy.

Anti-TNF agents should be avoided in patients with active infection or a history of hypersensitivity to these agents and are contraindicated in patients with chronic hepatitis B infection or class III/IV congestive

TABLE 380-2 DMARDs USED FOR THE TREATMENT OF RHEUMATOID ARTHRITIS

Drug	Dosage	Serious Toxicities	Other Common Side Effects	Initial Evaluation	Monitoring
Hydroxychloroquine	200–400 mg/d orally (≤6.5 mg/kg)	Irreversible retinal damage Cardiotoxicity Blood dyscrasia	Nausea Diarrhea Headache Rash	Eye examination if >40 years old or prior ocular disease	Fundoscopic and visual field testing every 12 months
Sulfasalazine	Initial: 500 mg orally twice daily Maintenance: 1000–1500 mg twice daily	Granulocytopenia Hemolytic anemia (with G6PD deficiency)	Nausea Diarrhea Headache	CBC, LFTs G6PD level	CBC every 2–4 weeks for first 3 months, then every 3 months
Methotrexate	10–25 mg/week orally or SQ Folic acid 1 mg/d to reduce toxicities	Hepatotoxicity Myelosuppression Infection Interstitial pneumonitis Pregnancy category X	Nausea Diarrhea Stomatitis/ mouth ulcers Alopecia Fatigue	CBC, LFTs Viral hepatitis panel ^a Chest x-ray	CBC, creatinine, LFTs every 2–3 months
Leflunomide	10–20 mg/d	Hepatotoxicity Myelosuppression Infection Pregnancy category X	Alopecia Diarrhea	CBC, LFTs Viral hepatitis panel ^a	CBC, creatinine, LFTs every 2–3 months
TNF-α Inhibitors	Infliximab: 3 mg/kg IV at weeks 0, 2, 6, then every 8 weeks. May increase dose up to 10 mg/kg every 4 weeks Etanercept: 50 mg SQ weekly, or 25 mg SQ biweekly Adalimumab: 40 mg SQ every other week Golimumab: 50 mg SQ monthly Certolizumab: 400 mg SQ weeks 0, 2, 4, then 200 mg every other week	↑ Risk bacterial, fungal infections Reactivation of latent TB ↑ Lymphoma risk (controversial) Drug-induced lupus Neurologic deficits As above As above As above As above	Infusion reaction ↑ LFTs Injection site reaction Injection site reaction Injection site reaction Injection site reaction	PPD skin test PPD skin test PPD skin test PPD skin test	LFTs periodically Monitor for injection site reactions Monitor for injection site reactions Monitor for injection site reactions Monitor for injection site reactions
Abatacept	Weight based: <60 kg: 500 mg 60–100 kg: 750 mg >100 kg: 1000 mg IV dose at weeks 0, 2, and 4, and then every 4 weeks OR 125 mg SQ weekly	↑ Risk bacterial, viral infections	Headache Nausea	PPD skin test	Monitor for infusion reactions
Anakinra	100 mg SQ daily	↑ Risk bacterial, viral infections Reactivation of latent TB Neutropenia	Injection site reaction Headache	PPD skin test CBC with differential	CBC every month for 3 months, then every 4 months for 1 year Monitor for injection site reactions
Rituximab	1000 mg IV × 2, days 0 and 14 May repeat course every 24 weeks or more Premedicate with methylprednisolone 100 mg to decrease infusion reaction	↑ Risk bacterial, viral infections Infusion reaction Cytopenia Hepatitis B reactivation	Rash Fever	CBC Viral hepatitis panel ^a	CBC at regular intervals

(Continued)

TABLE 380-2 DMARDs USED FOR THE TREATMENT OF RHEUMATOID ARTHRITIS (CONTINUED)

Drug	Dosage	Serious Toxicities	Other Common Side Effects	Initial Evaluation	Monitoring
Tocilizumab	4–8 mg/kg	Risk of infection		PPD skin test	CBC and LFTs at regular intervals
	4–8 mg/kg IV monthly	Infusion reaction			
	OR	LFT elevation			
	162 mg SQ every other week (<100 kg weight)	Dyslipidemia			
	162 mg SQ every week (≥100 kg weight)	Cytopenias			
Tofacitinib	5 mg orally BID	Risk of infection	Upper respiratory tract infections	PPD skin test	CBC, LFTs, and lipids at regular intervals
		LFT elevation	Diarrhea		
		Dyslipidemia	Headache		
		Neutropenia	Nasopharyngitis		

^aViral hepatitis panel: hepatitis B surface antigen, hepatitis C viral antibody.

Abbreviations: CBC, complete blood count; DMARDs, disease-modifying antirheumatic drugs; G6PD, glucose-6-phosphate dehydrogenase; IV, intravenous; LFTs, liver function tests; PPD, purified protein derivative; SQ, subcutaneous; TB, tuberculosis.

heart failure. The major concern is the increased risk for infection, including serious bacterial infections, opportunistic fungal infection, and reactivation of latent tuberculosis. For this reason, all patients are screened for latent tuberculosis according to national guidelines prior to starting anti-TNF therapy (Chap. 202). In the United States, patients are skin tested using an intradermal injection of purified protein derivative (PPD); individuals with skin reactions of more than 5 mm are presumed to have had previous exposure to tuberculosis and are evaluated for active disease and treated accordingly. The QuantiFERON IFN- γ release assay may also be used in selected circumstances to screen for previous exposure to tuberculosis.

Anakinra Anakinra, the recombinant form of the naturally occurring IL-1 receptor antagonist. Although anakinra has seen limited use for the treatment of RA, it has enjoyed a resurgence of late as an effective therapy of some rare inherited syndromes dependent on IL-1 production, including neonatal-onset inflammatory disease, Muckle-Wells syndrome, and familial cold urticaria, as well as systemic juvenile-onset inflammatory arthritis and adult-onset Still's disease. Anakinra should not be combined with an anti-TNF drug due to the high rate of serious infections as observed with this regimen in a clinical trial.

Abatacept Abatacept is a soluble fusion protein consisting of the extracellular domain of human CTLA-4 linked to the modified portion of human IgG. It inhibits the co-stimulation of T cells by blocking CD28-CD80/86 interactions and may also inhibit the function of antigen-presenting cells by reverse signaling through CD80 and CD86. Abatacept has been shown in clinical trials to reduce disease activity, slow radiographic progression of damage, and improve functional disability. Many patients receive abatacept in combination with methotrexate or another DMARD such as leflunomide. Abatacept therapy has been associated with an increased risk of infection.

Rituximab Rituximab is a chimeric monoclonal antibody directed against CD20, a cell-surface molecule expressed by most mature B lymphocytes. It works by depleting B cells, which in turn, leads to a reduction in the inflammatory response by unknown mechanisms. These mechanisms may include a reduction in autoantibodies, inhibition of T cell activation, and alteration of cytokine production. Rituximab has been approved for the treatment of refractory RA in combination with methotrexate and has been shown to be more effective for patients with seropositive than seronegative disease. Rituximab therapy has been associated with mild to moderate infusion reactions as well as an increased risk of infection. Notably, there have been isolated reports of a potentially lethal brain disorder, progressive multifocal leukoencephalopathy (PML), in association with rituximab therapy, although the absolute risk of this complication appears to be very low in patients with RA. Most of these cases have occurred on a background of previous or current exposure to other potent immunosuppressive drugs.

Tocilizumab Tocilizumab is a humanized monoclonal antibody directed against the membrane and soluble forms of the IL-6 receptor. IL-6 is a proinflammatory cytokine implicated in the pathogenesis of RA, with detrimental effects on both joint inflammation and damage. IL-6 binding to its receptor activates intracellular signaling pathways that affect the acute-phase response, cytokine production, and osteoclast activation. Clinical trials attest to the clinical efficacy of tocilizumab therapy for RA, both as monotherapy and in combination with methotrexate and other DMARDs. Tocilizumab has been associated with an increased risk of infection, neutropenia, and thrombocytopenia; however, the hematologic abnormalities appear to be reversible upon stopping the drug. In addition, this agent has been shown to increase LDL cholesterol; however, it is not known as yet if this effect on lipid levels increases the risk for development of atherosclerotic disease.

SMALL-MOLECULE INHIBITORS

Because some patients do not adequately respond to conventional DMARDs or biologic therapy, other therapeutic targets have been investigated to fill this gap. Recently, drug development in RA has focused attention on the intracellular signaling pathways that transduce the positive signals of cytokines and other inflammatory mediators that create the positive feedback loops in the immune response. These synthetic DMARDs aim to provide the same efficacy as biological therapies in an oral formulation.

Tofacitinib Tofacitinib is a small-molecule inhibitor that primarily inhibits JAK1 and JAK3, which mediate signaling of the receptors for the common γ -chain-related cytokines IL-2, -4, -7, -9, -15, and -21 as well as IFN- γ and IL-6. These cytokines all play roles in promoting T and B cell activation as well as inflammation. Tofacitinib, an oral agent, has been shown in randomized, placebo-controlled clinical trials to improve the signs and symptoms of RA significantly over placebo. Major adverse events include elevated serum transaminases indicative of liver injury, neutropenia, increased cholesterol levels, and elevation in serum creatinine. Its use is also associated with an increased risk of infections. Tofacitinib can be used as monotherapy or in combination with methotrexate.

APPROACH TO THE PATIENT: Rheumatoid Arthritis

The original treatment pyramid for RA is now considered to be obsolete and has evolved into a new strategy that focuses on several goals: (1) early, aggressive therapy to prevent joint damage and disability; (2) frequent modification of therapy with utilization of combination therapy where appropriate; (3) individualization of therapy in an attempt to maximize response and minimize side effects; and (4) achieving, whenever possible, remission of clinical disease activity. A considerable amount of evidence supports this intensive treatment approach.

As mentioned earlier, methotrexate is the DMARD of first choice for initial treatment of moderate to severe RA. Failure to achieve adequate improvement with methotrexate therapy calls for a change in DMARD therapy, usually transition to an effective combination regimen. Effective combinations include: methotrexate, sulfasalazine, and hydroxychloroquine (oral triple therapy); methotrexate and leflunomide; and methotrexate plus a biological. The combination of methotrexate and an anti-TNF agent, for example, has been shown in randomized, controlled trials to be superior to methotrexate alone not only for reducing signs and symptoms of disease, but also for retarding the progression of structural joint damage. Predicting which patients will ultimately show radiologic joint damage is imprecise at best, although some factors such as an elevated serum level of acute-phase reactants, high burden of joint inflammation, and the presence of erosive disease are associated with increased likelihood of developing structural injury.

In 2012 a joint task force of the ACR and EULAR updated the treatment guidelines for RA. They do make a distinction between patients with early RA (<6 months of disease duration) and patients with established RA. These guidelines highlight the need to switch or add DMARD therapy after 3 months of worsening or persistent moderate/high disease activity. If disease still persists after 3 months of intense DMARD therapy, addition of a biologic agent is warranted. Treatment with a biologic agent or aggressive combination DMARD therapy was also recommended as initial therapy in certain patients with high disease activity and poor prognosis. However, it has not been clearly established that this more intensive initial approach is superior to starting with methotrexate alone and, in the absence of an inadequate therapeutic response, moving rapidly to combination therapy.

Some patients may not respond to an anti-TNF drug or may be intolerant of its side effects. Initial responders to an anti-TNF agent that later worsen may benefit from switching to another anti-TNF agent. The 2012 guidelines recommend that with loss or lack of effectiveness of anti-TNF after 3 months, one should switch to another anti-TNF or non-TNF biologic agent. In patients with high disease activity and a serious adverse event from an anti-TNF agent, a non-TNF drug should be used.

Studies have also shown that oral triple therapy (hydroxychloroquine, methotrexate, and sulfasalazine) is a reasonable first step for the treatment of early RA, including its use as a step-up strategy where treatment is initiated with methotrexate alone and then combined at 6 months with hydroxychloroquine and sulfasalazine if the disease is not adequately controlled.

A clinical state defined as low disease activity or remission is the optimal goal of therapy, although most patients never achieve remission despite every effort to achieve it. Composite indices, such as the Disease Activity Score-28 (DAS-28), are useful for classifying states of low disease activity and remission; however, they are imperfect tools due to the limitations of the clinical joint examination in which low-grade synovitis may escape detection. Complete remission has been stringently defined as the total absence of all articular and extraarticular inflammation and immunologic activity related to RA. However, evidence for this state can be difficult to demonstrate in clinical practice. In an effort to standardize and simplify the definition of remission for clinical trials, the ACR and EULAR developed two provisional operational definitions of remission in RA (Table 380-3). A patient may be considered in remission if he or she (1) meets all of the clinical and laboratory criteria listed in Table 380-3 or (2) has a composite SDAI score of <3.3. The SDAI is calculated by taking the sum of a tender joint and swollen joint count (using 28 joints), patient global assessment (0–10 scale), physician global assessment (0–10 scale), and CRP (in mg/dL). This definition of remission does not take into account the possibility of subclinical synovitis or that damage alone may produce a tender or swollen joint. Ignoring the semantics of these definitions, the aforementioned remission criteria are nonetheless useful for setting

TABLE 380-3 ACR/EULAR PROVISIONAL DEFINITION OF REMISSION IN RHEUMATOID ARTHRITIS

At any time point, patient must satisfy all of the following:

- Tender joint count ≤ 1
- Swollen joint count ≤ 1
- C-reactive protein ≤ 1 mg/dL
- Patient global assessment ≤ 1 (on a 0–10 scale)

OR

At any time point, patient must have a Simplified Disease Activity Index score of ≤ 3.3

Source: Adapted from DT Felson et al: *Arthritis Rheum* 63:573, 2011.

a level of disease control that will likely result in minimal or no progression of structural damage and disability.

PHYSICAL THERAPY AND ASSISTIVE DEVICES

All patients should receive a prescription for exercise and physical activity. Dynamic strength training, community-based comprehensive physical therapy, and physical-activity coaching (emphasizing 30 min of moderately intensive activity most days a week) have all been shown to improve muscle strength and perceived health status. Foot orthotics for painful valgus deformity decrease foot pain and resulting disability and functional limitations. Judicious use of wrist splints can also decrease pain; however, their benefits may be offset by decreased dexterity and a variable effect on grip strength.

SURGERY

Surgical procedures may improve pain and disability in RA—most notably the hands, wrists, and feet, typically after the failure of medical therapy with varying degrees of reported long-term success. For large joints, such as the knee, hip, shoulder, or elbow, total joint arthroplasty is an option for advanced joint disease. A few surgical options exist for dealing with the smaller hand joints. Silicone implants are the most common prosthetic for MCP arthroplasty and are generally implanted in patients with severe decreased arc of motion, marked flexion contractures, MCP joint pain with radiographic abnormalities, and severe ulnar drift. Arthrodesis and total wrist arthroplasty are reserved for patients with severe disease who have substantial pain and functional impairment. These two procedures appear to have equal efficacy in terms of pain control and patient satisfaction. Numerous surgical options exist for correction of hallux valgus in the forefoot, including arthrodesis and arthroplasty, as well as primarily arthrodesis for refractory hindfoot pain.


OTHER MANAGEMENT CONSIDERATIONS

Pregnancy Up to 75% of female RA patients will note overall improvement in symptoms during pregnancy, but often will flare after delivery. Flares during pregnancy are generally treated with low doses of prednisone; hydroxychloroquine and sulfasalazine are probably the safest DMARDs to use during pregnancy. Methotrexate and leflunomide therapy are contraindicated during pregnancy due to their teratogenicity in animals and humans. The experience with biologic agents has been insufficient to make specific recommendations for their use during pregnancy. Most rheumatologists avoid their use in this setting; however, exceptions are considered depending on the circumstances.

Elderly Patients RA presents in up to one-third of patients after the age of 60; however, older individuals may receive less aggressive treatment due to concerns about increased risks of drug toxicity. Studies suggest that conventional DMARDs and biologic agents are equally effective and safe in younger and older patients. Due to comorbidities, many elderly patients have an increased risk of infection. Aging also leads to a gradual decline in renal function that may raise the risk for side effects from NSAIDs and some DMARDs, such as methotrexate. Renal function must be taken into consideration before prescribing methotrexate, which is mostly cleared by the kidneys. To reduce the risks of side effects, methotrexate doses may need to

be adjusted downward for the drop in renal function that usually comes with the seventh and eighth decades of life. Methotrexate is usually not prescribed for patients with a serum creatinine greater than 2 mg/dL.

GLOBAL CHALLENGES

 Developing countries are finding an increase in the incidence of noncommunicable, chronic diseases such as diabetes, cardiovascular disease, and RA in the face of ongoing poverty, rampant infectious disease, and poor access to modern health care facilities. In these areas, patients tend to have a greater delay in diagnosis and limited access to specialists, and thus greater disease activity and disability at presentation. In addition, infection risk remains a significant issue for the treatment of RA in developing countries because of the immunosuppression associated with the use of glucocorticoids and most DMARDs. For example, in some developing countries, patients undergoing treatment for RA have a substantial increase in the incidence of tuberculosis, which demands the implementation of far more comprehensive screening practices and liberal use of isoniazid prophylaxis than in developed countries. The increased prevalence of hepatitis B and C, as well as human immunodeficiency virus (HIV), in these developing countries also poses challenges. Reactivation of viral hepatitis has been observed in association with some of the DMARDs, such as rituximab. Also, reduced access to anti-retroviral therapy may limit the control of HIV infection and therefore the choice of DMARD therapies.

Despite these challenges, one should attempt to initiate early treatment of RA in the developing countries with the resources at hand. Sulfasalazine and methotrexate are all reasonably accessible throughout the world where they can be used as both monotherapy and in combination with other drugs. The use of biologic agents is increasing in the developed countries as well as in other areas around the world, although their use is limited by high cost; national protocols restrict their use, and concerns remain about the risk for opportunistic infections.

SUMMARY

Improved understanding of the pathogenesis of RA and its treatment has dramatically revolutionized the management of this disease. The outcomes of patients with RA are vastly superior to those of the prebiologic modifier era; more patients than in years past are able to avoid significant disability and continue working, albeit with some job modifications in many cases. The need for early and aggressive treatment of RA as well as frequent follow-up visits for monitoring of drug therapy has implications for our health care system. Primary care physicians and rheumatologists must be prepared to work together as a team to reach the ambitious goals of best practice. In many settings, rheumatologists have reengineered their practice in a way that places high priority on consultations for any new patient with early inflammatory arthritis.

The therapeutic regimens for RA are becoming increasingly complex with the rapidly expanding therapeutic armamentarium. Patients receiving these therapies must be carefully monitored by both the primary care physician and the rheumatologist to minimize the risk of side effects and identify quickly any complications of chronic immunosuppression. Also, prevention and treatment of RA-associated conditions such as ischemic heart disease and osteoporosis will likely benefit from a team approach owing to the value of multidisciplinary care.


Research will continue to search for new therapies with superior efficacy and safety profiles and investigate treatment strategies that can bring the disease under control more rapidly and nearer to remission. However, prevention and cure of RA will likely require new breakthroughs in our understanding of disease pathogenesis. These insights may come from genetic studies illuminating critical pathways in the mechanisms of joint inflammation. Equally ambitious is the lofty goal of biomarker discovery that will open the door to personalized medicine for the care of patients with RA.

381 Acute Rheumatic Fever

Jonathan R. Carapetis

Acute rheumatic fever (ARF) is a multisystem disease resulting from an autoimmune reaction to infection with group A streptococcus. Although many parts of the body may be affected, almost all of the manifestations resolve completely. The major exception is cardiac valvular damage (rheumatic heart disease [RHD]), which may persist after the other features have disappeared.

GLOBAL CONSIDERATIONS

 ARF and RHD are diseases of poverty. They were common in all countries until the early twentieth century, when their incidence began to decline in industrialized nations. This decline was largely attributable to improved living conditions—particularly less crowded housing and better hygiene—which resulted in reduced transmission of group A streptococci. The introduction of antibiotics and improved systems of medical care had a supplemental effect.

The virtual disappearance of ARF and reduction in the incidence of RHD in industrialized countries during the twentieth century unfortunately was not replicated in developing countries, where these diseases continue unabated. RHD is the most common cause of heart disease in children in developing countries and is a major cause of mortality and morbidity in adults as well. It has been estimated that between 15 and 19 million people worldwide are affected by RHD, with approximately one-quarter of a million deaths occurring each year. Some 95% of ARF cases and RHD deaths now occur in developing countries, with particularly high rates in sub-Saharan Africa, Pacific nations, Australasia, and South and Central Asia. The pathogenetic pathway from exposure to group A streptococcus followed by pharyngeal infection and subsequent development of ARF, ARF recurrences, and development of RHD and its complications is associated with a range of risk factors and, therefore, potential interventions at each point (Fig. 381-1). In affluent countries, many of these risk factors are well controlled, and where needed, interventions are in place. Unfortunately, the greatest burden of disease is found in developing countries, most of which do not have the resources, capacity, and/or interest to tackle this multifaceted disease. In particular, almost none of the developing countries has a coordinated, register-based RHD control program, which is proven to be cost effective in reducing the burden of RHD. Enhancing awareness of RHD and mobilizing resources for its control in developing countries are issues requiring international attention.

EPIDEMIOLOGY

ARF is mainly a disease of children age 5–14 years. Initial episodes become less common in older adolescents and young adults and are rare in persons age >30 years. By contrast, recurrent episodes of ARF remain relatively common in adolescents and young adults. This pattern contrasts with the prevalence of RHD, which peaks between 25 and 40 years. There is no clear gender association for ARF, but RHD more commonly affects females, sometimes up to twice as frequently as males.

PATHOGENESIS

ORGANISM FACTORS

Based on currently available evidence, ARF is exclusively caused by infection of the upper respiratory tract with group A streptococci (Chap. 173). Although classically, certain M-serotypes (particularly types 1, 3, 5, 6, 14, 18, 19, 24, 27, and 29) were associated with ARF, in high-incidence regions, it is now thought that any strain of group A streptococcus has the potential to cause ARF. The potential role of skin infection and of groups C and G streptococci is currently being investigated.

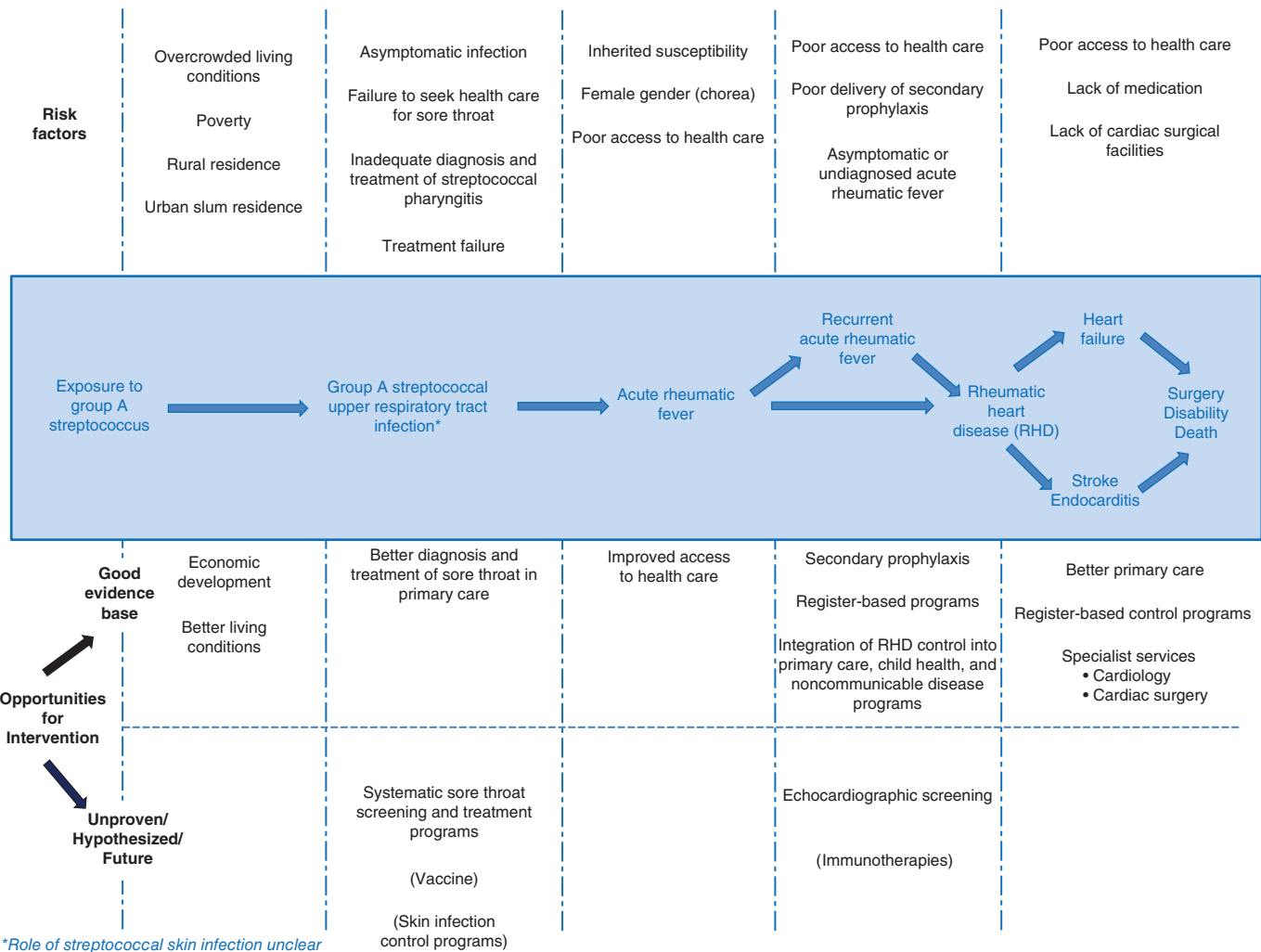


FIGURE 381-1 Pathogenetic pathway for acute rheumatic fever and rheumatic heart disease, with associated risk factors and opportunities for intervention at each step. Interventions in *parentheses* are either unproven or currently unavailable.

HOST FACTORS

Approximately 3–6% of any population may be susceptible to ARF, and this proportion does not vary dramatically between populations. Findings of familial clustering of cases and concordance in monozygotic twins—particularly for chorea—confirm that susceptibility to ARF is an inherited characteristic, with 44% concordance in monozygotic twins compared to 12% in dizygotic twins, and heritability more recently estimated at 60%. Most evidence for host factors focuses on immunologic determinants. Some human leukocyte antigen (HLA) class II alleles, particularly HLA-DR7 and HLA-DR4, appear to be associated with susceptibility, whereas other class II alleles have been associated with protection (HLA-DR5, HLA-DR6, HLA-DR51, HLA-DR52, and HLA-DQ). Associations have also been described with polymorphisms at the tumor necrosis factor α locus (TNF- α -308 and TNF- α -238), high levels of circulating mannose-binding lectin, and Toll-like receptors.

THE IMMUNE RESPONSE

The most widely accepted theory of rheumatic fever pathogenesis is based on the concept of molecular mimicry, whereby an immune response targeted at streptococcal antigens (mainly thought to be on the M protein and the *N*-acetylglucosamine of group A streptococcal carbohydrate) also recognizes human tissues. In this model, cross-reactive antibodies bind to endothelial cells on the heart valve, leading to activation of the adhesion molecule VCAM-1, with resulting recruitment of activated lymphocytes and lysis of endothelial cells in the presence of complement. The latter leads to release of peptides

including laminin, keratin, and tropomyosin, which, in turn, activates cross-reactive T cells that invade the heart, amplifying the damage and causing epitope spreading. An alternative hypothesis proposes that the initial damage is due to streptococcal invasion of epithelial surfaces, with binding of M protein to type IV collagen RHD allowing it to become immunogenic, but not through the mechanism of molecular mimicry.

CLINICAL FEATURES

There is a latent period of ~3 weeks (1–5 weeks) between the precipitating group A streptococcal infection and the appearance of the clinical features of ARF. The exceptions are chorea and indolent carditis, which may follow prolonged latent periods lasting up to 6 months. Although many patients report a prior sore throat, the preceding group A streptococcal infection is commonly subclinical; in these cases, it can only be confirmed using streptococcal antibody testing. The most common clinical features are polyarthritides (present in 60–75% of cases) and carditis (50–60%). The prevalence of chorea in ARF varies substantially between populations, ranging from <2 to 30%. Erythema marginatum and subcutaneous nodules are now rare, being found in <5% of cases.

HEART INVOLVEMENT

Up to 60% of patients with ARF progress to RHD. The endocardium, pericardium, or myocardium may be affected. Valvular damage is the hallmark of rheumatic carditis. The mitral valve is almost always

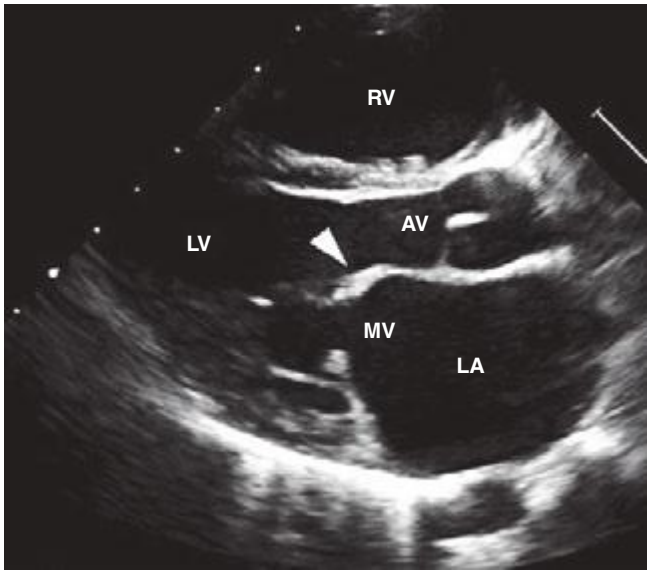


FIGURE 381-2 Transthoracic echocardiographic image from a 5-year-old boy with chronic rheumatic heart disease. This diastolic image demonstrates leaflet thickening, restriction of the anterior mitral valve leaflet tip and doming of the body of the leaflet toward the inter-ventricular septum. This appearance (marked by the *arrowhead*) is commonly described as a “hockey stick” or an “elbow” deformity. AV, aortic valve; LA, left atrium; LV, left ventricle; MV, mitral valve; RV, right ventricle. (Courtesy of Dr. Bo Remenyi, Department of Paediatric and Congenital Cardiac Services, Starship Children’s Hospital, Auckland, New Zealand.)

affected, sometimes together with the aortic valve; isolated aortic valve involvement is rare. Damage to the pulmonary or tricuspid valves is usually secondary to increased pulmonary pressures resulting from left-sided valvular disease. Early valvular damage leads to regurgitation. Over ensuing years, usually as a result of recurrent episodes, leaflet thickening, scarring, calcification, and valvular stenosis may develop (Fig. 381-2). See Videos 381-1 and 381-2 on the DVD. Therefore, the characteristic manifestation of carditis in previously unaffected individuals is mitral regurgitation, sometimes accompanied by aortic regurgitation. Myocardial inflammation may affect electrical conduction pathways, leading to P-R interval prolongation (first-degree atrioventricular block or rarely higher level block) and softening of the first heart sound.

People with RHD are often asymptomatic for many years before their valvular disease progresses to cause cardiac failure. Moreover, particularly in resource-poor settings, the diagnosis of ARF is often not made, so children, adolescents, and young adults may have RHD but not know it. These cases can be diagnosed using echocardiography; auscultation is poorly sensitive and specific for RHD diagnosis in asymptomatic patients. Echocardiographic screening of school-aged children in populations with high rates of RHD is becoming more widespread and has been facilitated by improving technologies in portable echocardiography and the availability of consensus guidelines for the diagnosis of RHD on echocardiography (Table 381-1). Although a diagnosis of definite RHD on screening echocardiography should lead to commencement of secondary prophylaxis, the clinical significance of borderline RHD has yet to be determined.

JOINT INVOLVEMENT

The most common form of joint involvement in ARF is arthritis, i.e., objective evidence of inflammation, with hot, swollen, red, and/or tender joints, and involvement of more than one joint (i.e., polyarthritis). Polyarthritis is typically migratory, moving from one joint to another over a period of hours. ARF almost always affects the large joints—most commonly the knees, ankles, hips, and elbows—and is asymmetric. The pain is severe and usually disabling until anti-inflammatory medication is commenced.

TABLE 381-1 WORLD HEART FEDERATION CRITERIA FOR ECHOCARDIOGRAPHIC DIAGNOSIS OF RHEUMATIC HEART DISEASE (RHD) IN INDIVIDUALS <20 YEARS OF AGE*

Definite RHD (either A, B, C, or D):	
(A)	Pathologic MR and at least two morphologic features of RHD of the mitral valve
(B)	MS mean gradient ≥ 4 mmHg (note: congenital MV anomalies must be excluded)
(C)	Pathologic AR and at least two morphologic features of RHD of the AV (note: bicuspid AV and dilated aortic root must be excluded)
(D)	Borderline disease of both the MV and AV
Borderline RHD (either A, B, or C):	
(A)	At least two morphologic features of RHD of the MV without pathologic MR or MS
(B)	Pathologic MR
(C)	Pathologic AR
Normal Echocardiographic Findings (all of A, B, C, and D):	
(A)	MR that does not meet all four Doppler criteria (physiologic MR)
(B)	AR that does not meet all four Doppler criteria (physiologic AR)
(C)	An isolated morphologic feature of RHD of the MV (e.g., valvular thickening), without any associated pathologic stenosis or regurgitation
(D)	Morphologic feature of RHD of the AV (e.g., valvular thickening), without any associated pathologic stenosis or regurgitation
Definitions of Pathologic Regurgitation and Morphologic Features of RHD:	
Pathologic MR: All of the following: seen in two views; in at least one view, jet length 2 cm; peak velocity ≥ 3 m/s; pansystolic jet in at least one envelope	
Pathologic AR: All of the following: seen in two views; in at least one view, jet length ≥ 1 cm; peak velocity ≥ 3 m/s; pandsystolic jet in at least one envelope	
Morphologic features of RHD in MV: anterior MV leaflet thickening ≥ 3 mm (age specific); chordal thickening; restricted leaflet motion; excessive leaflet tip motion during systole	
Morphologic features of RHD in AV: irregular or focal thickening; coaptation defect; restricted leaflet motion; prolapse	

*For criteria in individuals >20 years of age, see source document.

Abbreviations: AR, aortic regurgitation; AV, aortic valve; MR, mitral regurgitation; MS, mitral stenosis; MV, mitral valve.

Source: Adapted from Remenyi B et al: World Heart Federation criteria for echocardiographic diagnosis of rheumatic heart disease—an evidence-based guideline. *Nat Rev Cardiol* 9:297–309, 2012.

Less severe joint involvement is also relatively common and has been recognized as a potential major manifestation in high-risk populations in diagnostic guidelines from Australia, but at the time of writing, this was not reflected in the Jones criteria. Arthralgia without objective joint inflammation usually affects large joints in the same migratory pattern as polyarthritis. In some populations, aseptic monoarthritis may be a presenting feature of ARF, which may, in turn, result from early commencement of anti-inflammatory medication before the typical migratory pattern is established.

The joint manifestations of ARF are highly responsive to salicylates and other nonsteroidal anti-inflammatory drugs (NSAIDs). Indeed, joint involvement that persists for more than 1 or 2 days after starting salicylates is unlikely to be due to ARF.

CHOREA

Sydenham’s chorea commonly occurs in the absence of other manifestations, follows a prolonged latent period after group A streptococcal infection, and is found mainly in females. The choreiform movements affect particularly the head (causing characteristic darting movements of the tongue) and the upper limbs (Chap. 448). They may be generalized or restricted to one side of the body (hemi-chorea). In mild cases, chorea may be evident only on careful examination, whereas in the most severe cases, the affected individuals are unable to perform activities of daily living. There is often associated emotional lability or obsessive-compulsive traits,

2152 which may last longer than the choreiform movements (which usually resolve within 6 weeks but sometimes may take up to 6 months).

SKIN MANIFESTATIONS

The classic rash of ARF is *erythema marginatum* (Chap. 24), which begins as pink macules that clear centrally, leaving a serpiginous, spreading edge. The rash is evanescent, appearing and disappearing before the examiner's eyes. It occurs usually on the trunk, sometimes on the limbs, but almost never on the face.

Subcutaneous nodules occur as painless, small (0.5–2 cm), mobile lumps beneath the skin overlying bony prominences, particularly of the hands, feet, elbows, occiput, and occasionally the vertebrae. They are a delayed manifestation, appearing 2–3 weeks after the onset of disease, last for just a few days up to 3 weeks, and are commonly associated with carditis.

OTHER FEATURES

Fever occurs in most cases of ARF, although rarely in cases of pure chorea. Although high-grade fever ($\geq 39^{\circ}\text{C}$) is the rule, lower grade temperature elevations are not uncommon. Elevated acute-phase reactants are also present in most cases.

EVIDENCE OF A PRECEDING GROUP A STREPTOCOCCAL INFECTION

With the exception of chorea and low-grade carditis, both of which may become manifest many months later, evidence of a preceding group A streptococcal infection is essential in making the diagnosis of ARF. Because most cases do not have a positive throat swab culture or rapid antigen test, serologic evidence is usually needed. The most common serologic tests are the anti-streptolysin O (ASO) and anti-DNase B (ADB) titers. Where possible, age-specific reference ranges should be determined in a local population of healthy people without a recent group A streptococcal infection.

CONFIRMING THE DIAGNOSIS

Because there is no definitive test, the diagnosis of ARF relies on the presence of a combination of typical clinical features together with evidence of the precipitating group A streptococcal infection, and the exclusion of other diagnoses. This uncertainty led Dr. T. Duckett Jones in 1944 to develop a set of criteria (subsequently known as the *Jones criteria*) to aid in the diagnosis. At the time of writing, the Jones criteria were undergoing revision but had not yet been released. The existing diagnostic guideline is a World Health Organization update of the 1992 Jones Criteria (Table 381-2), though it should be noted that other guidelines, including those from Australia and New Zealand, suggest more sensitive criteria for making the diagnosis in patients from settings or populations at high risk of ARF.

TREATMENT ACUTE RHEUMATIC FEVER

Patients with possible ARF should be followed closely to ensure that the diagnosis is confirmed, treatment of heart failure and other symptoms is undertaken, and preventive measures including commencement of secondary prophylaxis, inclusion on an ARF registry, and health education are commenced. Echocardiography should be performed on all possible cases to aid in making the diagnosis and to determine the severity at baseline of any carditis. Other tests that should be performed are listed in Table 381-3.

There is no treatment for ARF that has been proven to alter the likelihood of developing, or the severity of, RHD. With the exception of treatment of heart failure, which may be life-saving in cases of severe carditis, the treatment of ARF is symptomatic.

ANTIBIOTICS

All patients with ARF should receive antibiotics sufficient to treat the precipitating group A streptococcal infection (Chap. 173). Penicillin is the drug of choice and can be given orally (as phenoxymethyl penicillin, 500 mg [250 mg for children ≤ 27 kg] PO twice daily, or amoxicillin, 50 mg/kg [maximum, 1 g] daily, for 10 days) or as a

TABLE 381-2 2002–2003 WORLD HEALTH ORGANIZATION CRITERIA FOR THE DIAGNOSIS OF RHEUMATIC FEVER AND RHEUMATIC HEART DISEASE (BASED ON THE 1992 REVISED JONES CRITERIA)

Diagnostic Categories	Criteria
Primary episode of rheumatic fever ^a	Two major or one major and two minor manifestations plus evidence of preceding group A streptococcal infection
Recurrent attack of rheumatic fever in a patient without established rheumatic heart disease	Two major or one major and two minor manifestations plus evidence of preceding group A streptococcal infection
Recurrent attack of rheumatic fever in a patient with established rheumatic heart disease ^b	Two minor manifestations plus evidence of preceding group A streptococcal infection ^c
Rheumatic chorea Insidious onset rheumatic carditis ^b	Other major manifestations or evidence of group A streptococcal infection not required
Chronic valve lesions of rheumatic heart disease (patients presenting for the first time with pure mitral stenosis or mixed mitral valve disease and/or aortic valve disease) ^d	Do not require any other criteria to be diagnosed as having rheumatic heart disease
Major manifestations	Carditis Polyarthritides Chorea Erythema marginatum Subcutaneous nodules
Minor manifestations	Clinical: fever, polyarthralgia Laboratory: elevated erythrocyte sedimentation rate or leukocyte count ^e Electrocardiogram: prolonged P-R interval
Supporting evidence of a preceding streptococcal infection within the last 45 days	Elevated or rising anti-streptolysin O or other streptococcal antibody, or A positive throat culture, or Rapid antigen test for group A streptococcus, or Recent scarlet fever ^f

^aPatients may present with polyarthritides (or with only polyarthralgia or monoarthritides) and with several (three or more) other minor manifestations, together with evidence of recent group A streptococcal infection. Some of these cases may later turn out to be rheumatic fever. It is prudent to consider them as cases of "probable rheumatic fever" (once other diagnoses are excluded) and advise regular secondary prophylaxis. Such patients require close follow-up and regular examination of the heart. This cautious approach is particularly suitable for patients in vulnerable age groups in high-incidence settings. ^bInfective endocarditis should be excluded. ^cSome patients with recurrent attacks may not fulfill these criteria. ^dCongenital heart disease should be excluded. ^e1992 Revised Jones criteria do not include elevated leukocyte count as a laboratory minor manifestation (but do include elevated C-reactive protein), and do not include recent scarlet fever as supporting evidence of a recent streptococcal infection.

Source: Reprinted with permission from WHO Expert Consultation on Rheumatic Fever and Rheumatic Heart Disease (2001: Geneva, Switzerland): *Rheumatic Fever and Rheumatic Heart Disease: Report of a WHO Expert Consultation* (WHO Tech Rep Ser, 923). Geneva, World Health Organization, 2004.

single dose of 1.2 million units (600,000 units for children ≤ 27 kg) IM benzathine penicillin G.

SALICYLATES AND NSAIDS

These may be used for the treatment of arthritis, arthralgia, and fever, once the diagnosis is confirmed. They are of no proven value in the treatment of carditis or chorea. Aspirin is the drug of choice, delivered at a dose of 50–60 mg/kg per day, up to a maximum of 80–100 mg/kg per day (4–8 g/d in adults) in four to five divided doses. At higher doses, the patient should be monitored for symptoms of salicylate toxicity such as nausea, vomiting, or tinnitus; if symptoms appear, lower doses should be used. When the acute

TABLE 381-3 RECOMMENDED TESTS IN CASES OF POSSIBLE ACUTE RHEUMATIC FEVER**Recommended for All Cases**

White blood cell count
Erythrocyte sedimentation rate
C-reactive protein
Blood cultures if febrile
Electrocardiogram (if prolonged P-R interval or other rhythm abnormality, repeat in 2 weeks and again at 2 months if still abnormal)
Chest x-ray if clinical or echocardiographic evidence of carditis
Echocardiogram (consider repeating after 1 month if negative)
Throat swab (preferably before giving antibiotics)—culture for group A streptococcus
Antistreptococcal serology: both anti-streptolysin O and anti-DNase B titers, if available (repeat 10–14 days later if first test not confirmatory)

Tests for Alternative Diagnoses, Depending on Clinical Features

Repeated blood cultures if possible endocarditis
Joint aspirate (microscopy and culture) for possible septic arthritis
Copper, ceruloplasmin, antinuclear antibody, drug screen for choreiform movements
Serology and autoimmune markers for arboviral, autoimmune, or reactive arthritis

Source: Reprinted with permission from Menzies School of Health Research.

symptoms are substantially resolved, usually within the first 2 weeks, patients on higher doses can have the dose reduced to 50–60 mg/kg per day for a further 2–4 weeks. Fever, joint manifestations, and elevated acute-phase reactants sometimes recur up to 3 weeks after the medication is discontinued. This does not indicate a recurrence and can be managed by recommencing salicylates for a brief period. Naproxen at a dose of 10–20 mg/kg per day is a suitable alternative to aspirin and has the advantage of twice-daily dosing.

CONGESTIVE HEART FAILURE

Glucocorticoids The use of glucocorticoids in ARF remains controversial. Two meta-analyses have failed to demonstrate a benefit of glucocorticoids compared to placebo or salicylates in improving the short- or longer term outcome of carditis. However, the studies included in these meta-analyses all took place >40 years ago and did not use medications in common usage today. Many clinicians treat cases of severe carditis (causing heart failure) with glucocorticoids in the belief that they may reduce the acute inflammation and result in more rapid resolution of failure. However, the potential benefits of this treatment should be balanced against the possible adverse effects. If used, prednisone or prednisolone is recommended at a dose of 1–2 mg/kg per day (maximum, 80 mg), usually for a few days or up to a maximum of 3 weeks.

MANAGEMENT OF HEART FAILURE

See Chap. 280.

BED REST

Traditional recommendations for long-term bed rest, once the cornerstone of management, are no longer widely practiced. Instead, bed rest should be prescribed as needed while arthritis and arthralgia are present and for patients with heart failure. Once symptoms are well controlled, gradual mobilization can commence as tolerated.

CHOREA

Medications to control the abnormal movements do not alter the duration or outcome of chorea. Milder cases can usually be managed by providing a calm environment. In patients with severe chorea, carbamazepine or sodium valproate is preferred to haloperidol. A response may not be seen for 1–2 weeks, and medication should be continued for 1–2 weeks after symptoms subside. There is recent

evidence that corticosteroids are effective and lead to more rapid symptom reduction in chorea. They should be considered in severe or refractory cases. Prednisone or prednisolone may be commenced at 0.5 mg/kg daily, with weaning as early as possible, preferably after 1 week if symptoms are reduced, although slower weaning or temporary dose escalation may be required if symptoms worsen.

INTRAVENOUS IMMUNOGLOBULIN (IVIG)

Small studies have suggested that IVIg may lead to more rapid resolution of chorea but have shown no benefit on the short- or long-term outcome of carditis in ARF without chorea. In the absence of better data, IVIg is *not* recommended except in cases of severe chorea refractory to other treatments.

PROGNOSIS

Untreated, ARF lasts on average 12 weeks. With treatment, patients are usually discharged from hospital within 1–2 weeks. Inflammatory markers should be monitored every 1–2 weeks until they have normalized (usually within 4–6 weeks), and an echocardiogram should be performed after 1 month to determine if there has been progression of carditis. Cases with more severe carditis need close clinical and echocardiographic monitoring in the longer term.

Once the acute episode has resolved, the priority in management is to ensure long-term clinical follow-up and adherence to a regimen of secondary prophylaxis. Patients should be entered onto the local ARF registry (if present) and contact made with primary care practitioners to ensure a plan for follow-up and administration of secondary prophylaxis before the patient is discharged. Patients and their families should also be educated about their disease, emphasizing the importance of adherence to secondary prophylaxis.

PREVENTION**PRIMARY PREVENTION**

Ideally, primary prevention would entail elimination of the major risk factors for streptococcal infection, particularly overcrowded housing. This is difficult to achieve in most places where ARF is common.

Therefore, the mainstay of primary prevention for ARF remains primary prophylaxis (i.e., the timely and complete treatment of group A streptococcal sore throat with antibiotics). If commenced within 9 days of sore throat onset, a course of penicillin (as outlined above for treatment of ARF) will prevent almost all cases of ARF that would otherwise have developed. In settings where ARF and RHD are common but microbiologic diagnosis of group A streptococcal pharyngitis is not available, such as in resource-poor countries, primary care guidelines often recommend that all patients with sore throat be treated with penicillin or, alternatively, that a clinical algorithm be used to identify patients with a higher likelihood of group A streptococcal pharyngitis. Although imperfect, such approaches recognize the importance of ARF prevention at the expense of overtreating many cases of sore throat that are not caused by group A streptococcus.

SECONDARY PREVENTION

The mainstay of controlling ARF and RHD is secondary prevention. Because patients with ARF are at dramatically higher risk than the general population of developing a further episode of ARF after a group A streptococcal infection, they should receive long-term penicillin prophylaxis to prevent recurrences. The best antibiotic for secondary prophylaxis is benzathine penicillin G (1.2 million units, or 600,000 units if ≤ 27 kg) delivered every 4 weeks. It can be given every 3 weeks, or even every 2 weeks, to persons considered to be at particularly high risk, although in settings where good compliance with an every-4-week dosing schedule can be achieved, more frequent dosing is rarely needed. Oral penicillin V (250 mg) can be given twice daily instead but is less effective than benzathine penicillin G. Penicillin-allergic patients can receive erythromycin (250 mg) twice daily.

The duration of secondary prophylaxis is determined by many factors, in particular the duration since the last episode of ARF

TABLE 381-4 AMERICAN HEART ASSOCIATION RECOMMENDATIONS FOR DURATION OF SECONDARY PROPHYLAXIS*

Category of Patient	Duration of Prophylaxis
Rheumatic fever without carditis	For 5 years after the last attack or 21 years of age (whichever is longer)
Rheumatic fever with carditis but no residual valvular disease	For 10 years after the last attack, or 21 years of age (whichever is longer)
Rheumatic fever with persistent valvular disease, evident clinically or on echocardiography	For 10 years after the last attack, or 40 years of age (whichever is longer); sometimes lifelong prophylaxis

*These are only recommendations and must be modified by individual circumstances as warranted. Note that some organizations recommend a minimum of 10 years of prophylaxis after the most recent episode, or until 21 years of age (whichever is longer), regardless of the presence of carditis with the initial episode.

Source: Adapted from AHA Scientific Statement Prevention of Rheumatic Fever and Diagnosis and Treatment of Acute Streptococcal Pharyngitis. *Circulation* 119:1541, 2009.

(recurrences become less likely with increasing time), age (recurrences are less likely with increasing age), and the severity of RHD (if severe, it may be prudent to avoid even a very small risk of recurrence because of the potentially serious consequences) (Table 381-4). Secondary prophylaxis is best delivered as part of a coordinated RHD control program, based around a registry of patients. Registries improve the ability to follow patients and identify those who default from prophylaxis and to institute strategies to improve adherence.

VIDEO 381-1A Transthoracic echocardiographic images of a 9-year-old girl with first episode of acute rheumatic fever. Images demonstrate the typical echocardiographic findings of acute rheumatic carditis. The valve leaflets are relatively thin and highly mobile. The failure of coaptation of the mitral valve leaflets is the result of chordal elongation and annular dilatation. The mitral valve regurgitation is moderate with a typical posterolaterally directed regurgitant jet of rheumatic carditis. **A.** Acute rheumatic carditis (apical four-chamber view echocardiogram).

VIDEO 381-1B Transthoracic echocardiographic images of a 9-year-old girl with first episode of acute rheumatic fever. Images demonstrate the typical echocardiographic findings of acute rheumatic carditis. The valve leaflets are relatively thin and highly mobile. The failure of coaptation of the mitral valve leaflets is the result of chordal elongation and annular dilatation. The mitral valve regurgitation is moderate with a typical posterolaterally directed regurgitant jet of rheumatic carditis. **B.** Acute rheumatic carditis (apical four-chamber view color Doppler echocardiogram).

VIDEO 381-1C Transthoracic echocardiographic images of a 9-year-old girl with first episode of acute rheumatic fever. Images demonstrate the typical echocardiographic findings of acute rheumatic carditis. The valve leaflets are relatively thin and highly mobile. The failure of coaptation of the mitral valve leaflets is the result of chordal elongation and annular dilatation. The mitral valve regurgitation is moderate with a typical posterolaterally directed regurgitant jet of rheumatic carditis. **C.** Acute rheumatic carditis (parasternal long-axis view echocardiogram).

VIDEO 381-1D Transthoracic echocardiographic images of a 9-year-old girl with first episode of acute rheumatic fever. Images demonstrate the typical echocardiographic findings of acute rheumatic carditis. The valve leaflets are relatively thin and highly mobile. The failure of coaptation of the mitral valve leaflets is the result of chordal elongation and annular dilatation. The mitral valve regurgitation is moderate with a typical posterolaterally directed regurgitant jet of rheumatic carditis. **D.** Acute rheumatic carditis (parasternal long-axis view color Doppler echocardiogram).

VIDEO 381-2A Transthoracic echocardiographic images are from a 5-year-old boy with chronic rheumatic heart disease with severe mitral valve regurgitation and moderate mitral valve stenosis. Images demonstrate the typical echocardiographic findings in advanced chronic rheumatic heart disease. Both the anterior and posterior mitral valve leaflets are markedly thickened. During diastole, the

motion of the anterior mitral valve leaflet tip is restricted with doming of the body of the leaflet toward the interventricular septum. This appearance is commonly described as a “hockey stick” or an “elbow” deformity. **A.** Chronic rheumatic heart disease (parasternal long-axis view). **VIDEO 381-2B** Transthoracic echocardiographic images are from a 5-year-old boy with chronic rheumatic heart disease with severe mitral valve regurgitation and moderate mitral valve stenosis. Images demonstrate the typical echocardiographic findings in advanced chronic rheumatic heart disease. Both the anterior and posterior mitral valve leaflets are markedly thickened. During diastole, the motion of the anterior mitral valve leaflet tip is restricted with doming of the body of the leaflet toward the interventricular septum. This appearance is commonly described as a “hockey stick” or an “elbow” deformity. **B.** Chronic rheumatic heart disease (apical two-chamber view echocardiogram).

382 Systemic Sclerosis (Scleroderma) and Related Disorders

John Varga

DEFINITION

Systemic sclerosis (SSc) is an uncommon connective tissue disorder characterized by multisystem involvement, heterogeneous clinical manifestations, a chronic and often progressive course, and significant disability and mortality. Multiple genes contribute to disease susceptibility; however, environmental exposures are likely to play a major role in causing SSc. The early stage of the disease is associated with prominent inflammatory features. Over time, functional and structural alterations in multiple vascular beds and progressive visceral organ dysfunction due to fibrosis dominate the clinical picture. Although thickened skin (scleroderma) is the distinguishing hallmark of SSc, skin induration can occur in localized forms of scleroderma and other disorders (Table 382-1). Patients with SSc can be broadly

TABLE 382-1 CONDITIONS ASSOCIATED WITH SKIN INDURATION

Systemic sclerosis (SSc)
Limited cutaneous SSc
Diffuse cutaneous SSc
Localized scleroderma
Guttate (plaque) morphea, diffuse (pansclerotic) morphea, bullous morphea
Linear scleroderma, coup de sabre, hemifacial atrophy
Pansclerotic morphea
Overlap syndromes
Mixed connective tissue disease
SSc/polymyositis
Stiff skin syndrome
Diabetic scleredema and scleredema of Buschke
Scleromyxedema (papular mucinosis)
Chronic graft-versus-host disease
Diffuse fasciitis with eosinophilia (Shulman's disease, eosinophilic fasciitis)
Chemically induced and drug-associated scleroderma-like conditions
Vinyl chloride–induced disease
Eosinophilia-myalgia syndrome (associated with L-tryptophan)
Nephrogenic systemic fibrosis (associated with gadolinium)
Paraneoplastic syndrome

TABLE 382-2 SUBSETS OF SYSTEMIC SCLEROSIS (SSc): FEATURES OF LIMITED CUTANEOUS SSc VERSUS DIFFUSE CUTANEOUS SSc

Characteristic Feature	Limited Cutaneous SSc	Diffuse Cutaneous SSc
Skin involvement	Indolent onset. Limited to fingers, distal to elbows, face; slow progression	Rapid onset. Diffuse: fingers, extremities, face, trunk; rapid progression
Raynaud's phenomenon	Antedates skin involvement, sometimes by years; may be associated with critical ischemia in the digits	Onset coincident with skin involvement; critical ischemia less common
Musculoskeletal	Mild arthralgia	Severe arthralgia, carpal tunnel syndrome, tendon friction rubs
Interstitial lung disease	Slowly progressive, generally mild	Frequent, early onset and progression, can be severe
Pulmonary arterial hypertension	Frequent, late, may occur as an isolated complication	Often occurs in association with interstitial lung disease
Scleroderma renal crisis	Very rare	Occurs in 15%; generally early (<4 years from disease onset)
Calcinosis cutis	Frequent, prominent	Less common, mild
Characteristic autoantibodies	Anticentromere	Anti-topoisomerase I (Scl-70), anti-RNA polymerase III

grouped into diffuse cutaneous and limited cutaneous subsets defined by the pattern of skin involvement, as well as clinical and laboratory features (Table 382-2). Diffuse cutaneous SSc (dcSSc) is associated with extensive skin induration, starting in the fingers and ascending from distal to proximal limbs and the trunk. These patients often have early interstitial lung disease and acute renal involvement. In contrast, in patients with limited cutaneous SSc (lcSSc), Raynaud's phenomenon may precede other manifestations of SSc by years. In these patients, skin involvement remains limited to the fingers (sclerodactyly), distal limbs, and face, and the trunk is not affected. The constellation of calcinosis cutis, Raynaud's phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia, seen in some lcSSc patients, is termed the *CREST syndrome*. Visceral organ involvement in lcSSc tends to show insidious progression, and pulmonary arterial hypertension (PAH), interstitial lung disease, hypothyroidism, and primary biliary cirrhosis may occur as late complications. In some patients, Raynaud's phenomenon and other characteristic features of SSc occur in the absence of skin thickening. This syndrome has been termed *SSc sine scleroderma*.

EPIDEMIOLOGY

SSc is an acquired sporadic disease with a worldwide distribution and affecting all races. In the United States, the incidence is estimated at 9–19 cases per million per year. The only community-based survey of SSc yielded a prevalence of 286 cases per million. There are an estimated 100,000 cases in the United States, although this number may be significantly higher if patients who do not meet strict classification criteria are also included. Rates of SSc in England, Australia, and Japan appear to be lower. Age, gender, and ethnicity are important in disease susceptibility. In common with other connective tissue diseases, SSc shows a strong female predominance (4.6:1), which is most pronounced in the childbearing years and declines after menopause. Although SSc can present at any age, the peak age of onset for both limited and diffuse cutaneous forms is 30–50 years. The incidence is higher in blacks than whites, and disease onset occurs at an earlier age. Furthermore, blacks with SSc are more likely to have diffuse cutaneous disease associated with interstitial lung involvement and a worse prognosis.

GENETIC CONSIDERATIONS



In general, SSc shows modest heritability, and the genetic associations identified to date make only a small contribution to disease susceptibility. Concordance rates for SSc are low (4.7%) in monozygotic twins, although concordance for antinuclear antibody (ANA) positivity is significantly higher. On the other hand, evidence for genetic contribution to disease susceptibility is provided by the observation that 1.6% of SSc patients have a first-degree relative with SSc, a prevalence rate markedly increased compared to the general population. The risk of Raynaud's phenomenon, interstitial lung disease, and other autoimmune diseases, including systemic lupus erythematosus (SLE) (Chap. 378), rheumatoid arthritis (Chap. 380), and autoimmune thyroiditis (Chap. 405), is also increased. Approaches to study the role of genetics in SSc use candidate gene single nucleotide polymorphism (SNP) analysis and genome-wide association studies (GWASs). Candidate gene studies in SSc have shown associations with multiple gene variants, many related to B and T lymphocyte activation and signaling (*BANK1*, *BLK*, *CD247*, *CSK*, *IRAK1*, *IL2RA*, *PTPN22*, and *TNIP1*). *IRAK1*, which codes for a gene involved in both innate and adaptive immunity, is the first X-linked gene associated with SSc and may contribute to female predominance. Other gene variants associated with SSc are involved in innate immunity and the interferon pathways (*IRF5*, *IRF7*, *STAT4*, *TNFAIP3*, and *TLR2*). In addition, candidate gene studies and GWAS both identified association with genes in the major histocompatibility complex (MHC), including *NOTCH4* and *PSORSC1*. In addition to disease susceptibility, some of these genetic loci are associated with particular SSc disease manifestations or serologic subsets, including interstitial lung disease (ILD) (*CTGF*, *CD226*), PAH (*TNIP1*), and scleroderma renal crisis (*HLA-DRB1**). Although the functional consequences of these gene variants are currently not well understood, they may result in altered immune function, leading to increased susceptibility to autoimmunity and inflammation. Of note, many of the genetic variants associated with SSc are also seen in other autoimmune disorders, including SLE, rheumatoid arthritis, and psoriasis, suggesting common pathways shared among these conditions. The genetic associations identified to date only explain a fraction of the heritability of SSc, and GWASs, fine mapping, and resequencing of DNA regions of interest to identify additional genetic susceptibility factors in SSc, particularly rare variants, are currently ongoing.

ENVIRONMENTAL AND OCCUPATIONAL RISK FACTORS

Given the relatively modest genetic contribution to disease susceptibility, environmental factors, such as infectious agents, intestinal microbiota, and occupational, dietary, and drug exposures, are likely to play a major role in causing SSc. Patients with SSc show evidence of chronic infection of lesional tissue with Epstein-Barr virus (EBV). They also have increased antibodies to human cytomegalovirus (hCMV), and anti-topoisomerase I (Scl-70) autoantibodies recognize hCMV-associated antigenic epitopes, suggesting molecular mimicry as a possible mechanistic link between hCMV infection and SSc. An epidemic of a novel syndrome with features suggestive of SSc occurred in Spain in the 1980s. The outbreak, termed *toxic oil syndrome*, was linked to contaminated rapeseed oils used for cooking. Another epidemic outbreak, termed *eosinophilia-myalgia syndrome* (EMS), occurred a decade later and was linked to the consumption of L-tryptophan-containing dietary supplements. Although both of these novel toxic-epidemic syndromes were characterized by scleroderma-like chronic skin changes and variable visceral organ involvement, they were associated with clinical, pathologic, and laboratory features distinguishing them from SSc. Occupational exposures tentatively linked with SSc include silica dust in miners, polyvinyl chloride, epoxy resins, and aromatic hydrocarbons including toluene and trichloroethylene. Drugs implicated in SSc-like illnesses include bleomycin, pentazocine, and cocaine, and appetite suppressants linked with pulmonary hypertension. Although case reports and series describing SSc in women with silicone breast implants had raised concern regarding a possible causal role of silicone

PATHOGENESIS

The following three cardinal pathophysiologic processes account for the protean clinical manifestations of SSc: (1) diffuse microangiopathy, (2) inflammation and autoimmunity, and (3) visceral and vascular fibrosis in multiple organs (Fig. 382-1). Autoimmunity and altered vascular reactivity are early manifestations. Complex and dynamic interplay between these processes initiates and then amplifies the fibrotic process.

ANIMAL MODELS OF DISEASE

No single animal model of SSc fully reproduces the three cardinal processes that underlie the pathogenesis, but some recapitulate selected aspects of the human disease, including fibrosis, microvascular involvement, and autoimmunity. Tight-skin mice (Tsk1) develop spontaneous skin thickening due to a mutation in the fibrillin-1 gene. The mutant fibrillin-1 protein disrupts extracellular matrix assembly and causes aberrant activation of transforming growth factor β (TGF- β). Fibrillin-1 mutations are associated with Marfan's disease as well as the stiff skin syndrome but not SSc. Skin and lung fibrosis can be induced in mice by injection of bleomycin, HOCl, or double-stranded RNA or by transplantation of human leukocyte antigen (HLA)-mismatched bone marrow or spleen cells. Targeted genetic modifications in mice give rise to new disease models for dissecting the pathogenetic roles of individual molecules, cell types, and networks. For example, mice lacking Smad3, an intracellular TGF- β signal transducer, adiponectin, or the nuclear receptor peroxisome proliferator-activated receptor (PPAR) γ or overexpressing Wnt10b

or adiponectin were either resistant or hypersensitive to chemically induced experimental scleroderma. These mouse models have potential utility in preclinical evaluation of potential therapies.

MICROANGIOPATHY

Involvement of small blood vessels in SSc affects multiple vascular beds and has important clinical sequelae including Raynaud's phenomenon, ischemic digital ulcers, scleroderma renal crisis, and PAH. Raynaud's phenomenon, an early disease manifestation, is characterized by an altered blood-flow response to cold challenge. This initially reversible functional abnormality is associated with autonomic and peripheral nervous system alterations, including impaired production of the neuropeptide calcitonin gene-related peptide from sensory afferent nerves and heightened sensitivity of α_2 -adrenergic receptors on vascular smooth-muscle cells. Isolated (primary) Raynaud's phenomenon is extremely common and generally benign and nonprogressive. In contrast, SSc-associated Raynaud's phenomenon is often progressive and complicated by irreversible structural changes, culminating in ischemic digital ulcers and loss of digits. Viruses, vascular cytotoxic factors, thrombogenic microparticles, complement and autoantibodies to phospholipids, β_2 glycoprotein I (β_2 GPI), and endothelial cells are suspected triggers of endothelial cell injury in SSc. Endothelial injury results in dysregulated production of endothelium-derived vasodilatory (nitric oxide and prostacyclin) and vasoconstricting (endothelin-1) substances, as well as increased expression of intercellular adhesion molecule 1 (ICAM-1) and other surface adhesion molecules. Microvessels show enhanced permeability and transendothelial leukocyte diapedesis, abnormal activation of coagulation cascades, elevated thrombin production, and impaired fibrinolysis. Spontaneous platelet aggregation causes release of serotonin, platelet-derived growth factor (PDGF), and platelet alpha granules including thromboxane, a potent vasoconstrictor. Smooth-muscle cell-like myointimal cells proliferate, the basement membrane is thickened and reduplicated, and fibrosis of the adventitial layers develops. The vasculopathic process affects capillaries, as well as arterioles, and even large vessels in many organs, resulting in reduced blood flow and tissue ischemia. Progressive luminal occlusion due to intimal and medial hypertrophy, combined with persistent endothelial cell damage and adventitial fibrosis, establish a vicious cycle that culminates in the striking absence of blood vessels (rarefaction) in late-stage disease. Recurrent ischemia-reperfusion generates reactive oxygen species (ROS) that further damage the endothelium through peroxidation of membrane lipids. Paradoxically, the process of revascularization that normally reestablishes blood flow to ischemic tissue is defective in SSc despite elevated levels of vascular endothelial growth factor (VEGF) and other angiogenic factors. Moreover, the number of bone marrow-derived circulating endothelial progenitor cells is reduced. Thus, widespread capillary loss, obliterative vasculopathy of small and medium-sized arteries, and failure to repair damaged vessels are hallmarks of SSc.

IMMUNE DYSREGULATION

Cellular Immunity The following observations highlight the autoimmune nature of SSc: presence of circulating autoantibodies; familial clustering of SSc with other autoimmune diseases; detection of immune cells, including T cells with oligoclonal antigen receptors, in target organs; elevated circulating levels and spontaneous secretion from blood mononuclear cells of inflammatory cytokines and chemokines such as interleukin (IL) 1, IL-4, IL-10, IL-17, IL-33, CCL2, and CXCL4; and the association with variants in genes functionally implicated in immune responses. Genetic studies in SSc reveal strong and consistent associations with major histocompatibility locus alleles, as well as non-HLA-linked genes encoding mediators of both adaptive and innate immune responses (*CD247*, *STAT4*, *IRF5*, *CD226*, and *TNFSF4*). In early SSc, mononuclear inflammatory cell infiltrates comprised of activated T cells, monocytes/macrophages, and dendritic cells can be seen in skin, lungs, and other affected organs prior to appearance of fibrosis or vascular damage. Dendritic cells and T cells can often be found in close proximity to activated fibroblasts and myofibroblasts. Tissue-infiltrating T cells express CD45 and HLA-DR activation markers and display restricted T cell receptor signatures indicative of oligoclonal expansion in response to (unknown) antigen.

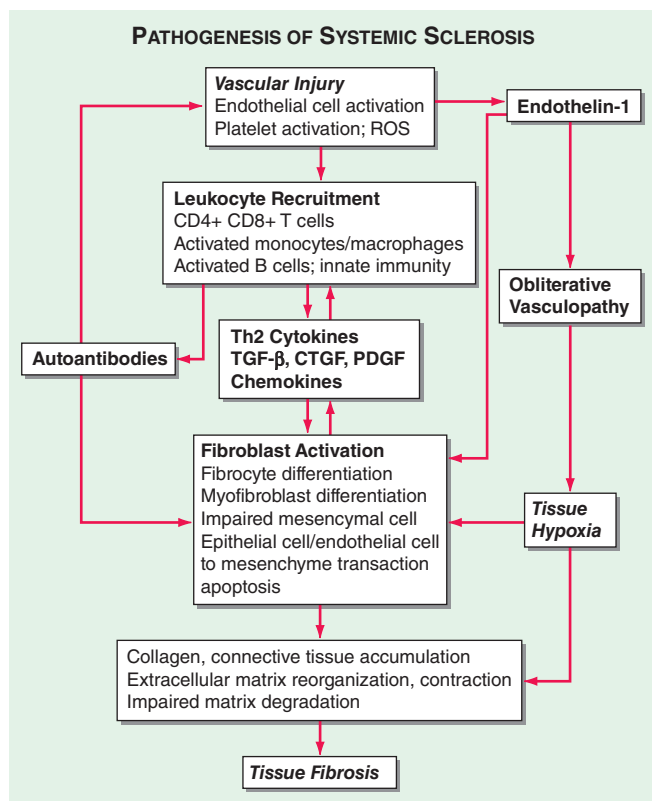


FIGURE 382-1 Initial vascular injury in a genetically susceptible individual leads to functional and structural vascular alterations, inflammation, and autoimmunity. The inflammatory and immune responses initiate and sustain fibroblast activation and differentiation, resulting in pathologic fibrogenesis and irreversible tissue damage. Vascular damage results in tissue ischemia that further contributes to progressive fibrosis and atrophy. CTGF, connective tissue growth factor; PDGF, platelet-derived growth factor; TGF- β , transforming growth factor β .

TABLE 382-3 AUTOANTIBODIES AND ASSOCIATED FEATURES IN SYSTEMIC SCLEROSIS (SSC)

Target Antigen	SSc Subset	Characteristic Clinical Association
Topoisomerase I	dcSSc	Tendon friction rubs, early ILD, cardiac involvement, scleroderma renal crisis
Centromere proteins	lcSSc	Digital ischemic ulcers, calcinosis cutis, isolated PAH, overlap syndromes; renal crisis rare
RNA polymerase III	dcSSc	Rapidly progressive skin involvement, tendon friction rubs, joint contractures, GAVE, renal crisis, contemporaneous cancers
U3-RNP (fibrillarin)	dcSSc	PAH, ILD, scleroderma renal crisis, myositis
Th/T0	lcSSc	ILD, PAH
PM/Scl	lcSSc	Calcinosis cutis, ILD, myositis overlap
Ku	Overlap	SLE, myositis
U1-RNP	MCTD	PAH, arthritis, myositis

Abbreviations: dcSSc, diffuse cutaneous SSc; GAVE, gastric antral vascular ectasia; ILD, interstitial lung disease; lcSSc, limited cutaneous SSc; MCTD, mixed connective tissue disease; PAH, pulmonary arterial hypertension; SLE, systemic lupus erythematosus.

Circulating T cells have elevated levels of chemokine receptors and α_1 integrin adhesion molecules, accounting for their enhanced binding to endothelium and to fibroblasts. Endothelial cells express ICAM-1 and other adhesion molecules that facilitate leukocyte diapedesis. Activated macrophages and T cells show a T_H2 -polarized type 2 immune response driven by dendritic cells and thymic stromal lymphopoietin. T_H2 cytokines such as IL-4 and IL-13 induce fibroblast activation and alternate M2 macrophage polarization, whereas the T_H1 cytokine interferon γ (IFN- γ) blocks cytokine-mediated fibroblast activation. Alternately activated M2 macrophages produce TGF- β and promote fibrosis. Although the frequency of circulating regulatory T cells that enforce immune tolerance is elevated in SSc, their immunosuppressive function is defective. Molecular characterization of SSc skin biopsies using DNA microarrays identifies a subset showing markedly elevated expression of inflammation-associated genes, particularly chemokines and their receptors, interferon response genes, and mediators of innate immunity. Evidence of activated innate immunity and toll-like receptor signaling, indicative of activation by type 1 interferon produced by plasmacytoid dendritic cells, is prominent in peripheral blood cells.

Humoral Autoimmunity Circulating ANAs can be detected in virtually all patients with SSc. In addition, a number of SSc-specific autoantibodies have been described. These SSc-specific antibodies show strong association with distinct disease endophenotypes (Table 382-3). While most are directed against intracellular proteins associated with cell proliferation, such as topoisomerase I and RNA polymerases I, II, and III, others are directed against cell-surface antigens, receptors, or secreted proteins. Autoantibodies have clinical utility as diagnostic and prognostic biomarkers in SSc, and some, such as antibodies directed against the angiotensin II receptor or the PDGF receptor, may have a direct pathogenic role.

A variety of mechanisms have been proposed for the development of autoantibodies in SSc. Proteolytic cleavage, increased expression, or altered subcellular localization of certain cellular proteins in SSc could lead to their recognition as neoepitopes by the immune system, resulting in breakdown of immune tolerance. B cells are implicated in both the autoimmune and fibrotic process in SSc. In addition to antibody production, B cells also present antigen, secrete IL-6 and TGF- β , and modulate T cell and dendritic cell function.

FIBROSIS

Fibrosis affecting multiple organs, a distinguishing feature of SSc, is characterized by progressive replacement of normal tissue architecture with dense, stiff, and acellular connective tissue. Fibrosis characteristically follows, and is thought to be a consequence of, inflammation,

autoimmunity, and microvascular damage. Fibroblasts are mesenchymal cells responsible for maintaining the functional and structural integrity of connective tissue. Upon activation by TGF- β and other extracellular cues, fibroblasts proliferate; migrate; secrete collagens, growth factors, chemokines, and cytokines; and transdifferentiate into contractile myofibroblasts. Under normal conditions, these fibrotic responses constitute self-limited physiologic remodeling necessary for tissue repair and regeneration. When these responses become sustained and amplified, pathologic fibrosis results. Autocrine stimulatory signaling by endogenously produced TGF- β and fibrotic mediators such as hypoxia, ROS, thrombin, Wnt ligands, connective tissue growth factor (CTGF), PDGF, lysophosphatidic acid, endothelin-1, mechanical forces, and endogenous ligands for toll-like receptors are responsible for maintaining sustained fibroblast activation underlying progressive fibrosis in SSc.

In addition to tissue-resident fibroblasts and transformation of epithelial cells into fibroblasts, bone marrow-derived circulating mesenchymal progenitor cells also contribute to fibrosis. The factors that regulate the differentiation of mesenchymal progenitor cells and their trafficking from the circulation into lesional tissue are unknown. Epithelial and endothelial cells, mesenchymal progenitor cells, and tissue fibroblasts can differentiate into smooth-muscle-like myofibroblasts. Although myofibroblasts can be detected transiently during normal wound healing, they persist in fibrotic tissue, possibly due to resistance to apoptosis, and contribute to scar formation via production of collagen and TGF- β and contraction of the surrounding extracellular matrix.

Explanted SSc fibroblasts may display an abnormally activated phenotype *ex vivo*, with variably increased rates of collagen gene transcription, spontaneous ROS generation, and constitutive expression of alpha smooth-muscle actin stress fibers. The persistence of the “scleroderma phenotype” of these cells during their serial passage *in vitro* may reflect autocrine TGF- β stimulatory loops, deregulated microRNA expressions, histone acetylation, and other epigenetic modifications.

PATHOLOGY

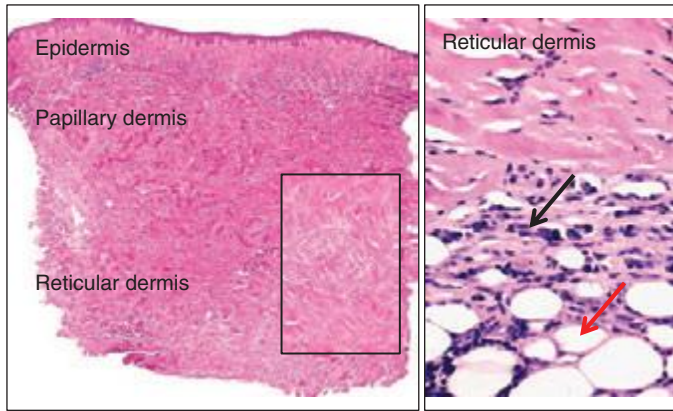
The distinguishing pathologic hallmark of SSc is the combination of widespread capillary loss and obliterative microangiopathy, together with fibrosis in the skin and internal organs. In early disease, perivascular inflammatory cell infiltrates composed of T lymphocytes, monocytes/macrophages, plasma cells, mast cells, and occasionally B cells may be detected in multiple organs. A bland noninflammatory obliterative vasculopathy as a late finding is prominent in the heart, lungs, kidneys, and intestinal tract. Fibrosis is found in the skin, lungs, gastrointestinal tract, heart, tendon sheaths, perifascicular tissue surrounding skeletal muscle, and some endocrine organs. In these tissues, accumulation of collagens, fibronectin, proteoglycans, tenascin, cartilage oligomeric matrix protein (COMP), and other structural macromolecules progressively disrupts normal architecture, resulting in impaired function of affected organs.

SKIN

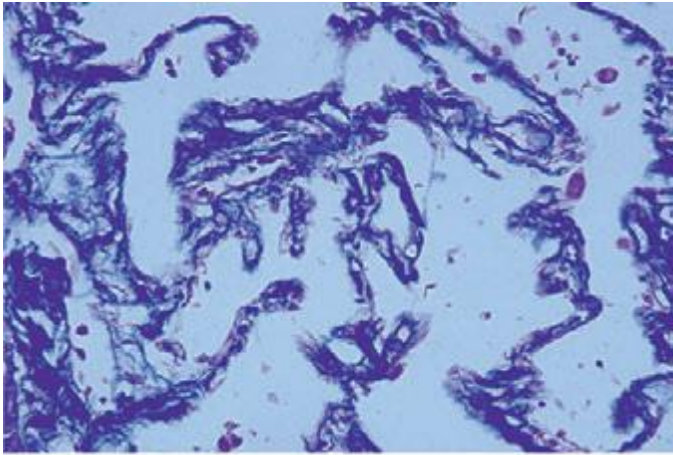
In the skin, fibrosis causes dermal expansion and obliteration of the hair follicles, eccrine glands, and other appendages (Fig. 382-2A). Collagen fiber accumulation is most prominent in the reticular dermis, and the fibrotic process invades the subjacent adipose layer with entrapment of adipocytes. With disease progression, the intradermal adipose layer is diminished and may completely disappear. The epidermis is atrophic, and the rete pegs are effaced.

LUNGS

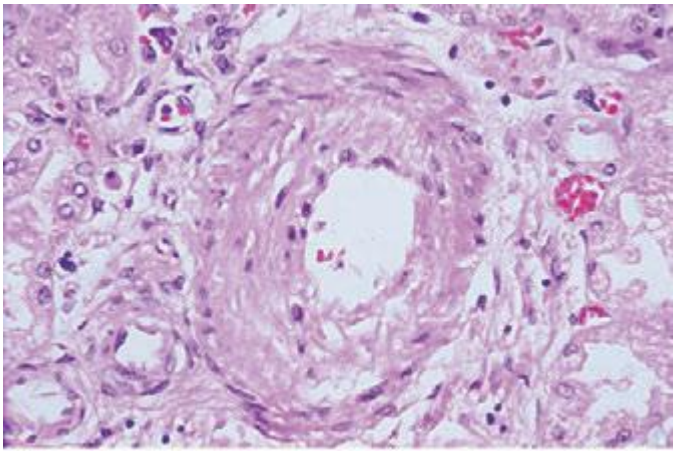
Patchy infiltration of the alveolar walls with T lymphocytes, macrophages, and eosinophils occurs in early disease. With progression, interstitial fibrosis and vascular damage dominate the pathologic picture, often coexisting within the same lesions in patients with dcSSc. Pulmonary fibrosis is characterized by expansion of the alveolar interstitium, with accumulation of collagen and other matrix proteins. The most common histologic pattern in SSc-associated ILD is nonspecific interstitial pneumonia (NSIP), distinct from the usual interstitial pneumonia (UIP) pattern characteristically seen in patients with



A



B



C

FIGURE 382-2 Pathologic findings in systemic sclerosis (SSc).

A. Left panel: The skin is thickened due to the marked expansion of the dermis. Inset, higher magnification showing thick hyalinized collagen bundles replace skin appendages. **Right panel:** Inflammation in the reticular dermis. Mononuclear inflammatory cells infiltrating the dermis and intradermal adipose tissue. **B.** Early interstitial lung disease. Diffuse fibrosis of the alveolar septae and a chronic inflammatory cell infiltrate. Trichrome stain. **C.** Pulmonary arterial obliterative vasculopathy. Striking intimal hyperplasia and narrowing of the lumen of a small pulmonary artery, with minimal interstitial fibrosis, in a patient with limited cutaneous SSc.

idiopathic pulmonary fibrosis (Fig. 382-2B). Progressive thickening of the alveolar septae results in obliteration of the airspaces and loss of pulmonary blood vessels. This process impairs gas exchange and contributes to pulmonary hypertension. Intimal thickening of the pulmonary arteries, best seen with elastin stain, underlies pulmonary hypertension (Fig. 382-2C) and, at autopsy, is often associated with multiple pulmonary emboli and evidence of myocardial fibrosis.

GASTROINTESTINAL TRACT

Pathologic changes can be found at any level from the mouth to the rectum. The lower esophagus shows prominent atrophy of the muscular layers and characteristic vascular lesions; striated muscle in the upper third of the esophagus is generally spared. Replacement of the normal intestinal tract architecture results in diminished peristaltic activity, with gastroesophageal reflux, dysmotility, and small-bowel obstruction. Chronic reflux is associated with esophageal inflammation, ulcerations, and stricture formation and may lead to Barrett's metaplasia.

KIDNEYS

In the kidneys, vascular lesions affecting the interlobular and arcuate arteries predominate. Chronic renal ischemia is associated with shrunken glomeruli. Acute scleroderma renal crisis is associated with a classic thrombotic microangiopathic pathology: reduplication of elastic lamina, marked intimal proliferation, and narrowing of the lumen in small renal arteries, commonly accompanied by thrombosis and hemolysis.

HEART

The heart is frequently affected, with prominent involvement of the myocardium and pericardium. The characteristic arteriolar lesions are concentric intimal hypertrophy and luminal narrowing, accompanied by contraction band necrosis reflecting ischemia-reperfusion injury and myocardial fibrosis. Fibrosis of the conduction system is common, especially at the sinoatrial node. Despite the prominent role of ischemia in SSc, the frequency of atherosclerotic coronary artery disease is comparable to the general population.

OTHER ORGANS

Synovitis may be found in early SSc; however, with disease progression, the synovium becomes fibrotic. Fibrosis of tendon sheaths and fascia produces palpable and sometimes audible tendon friction rubs. Inflammation and, in later stages, atrophy and fibrosis of the muscles are common findings. Fibrosis of the thyroid gland and of the minor salivary glands may be seen.

CLINICAL FEATURES

OVERVIEW

Virtually every organ can be clinically affected (Table 382-4). Most patients with SSc can be classified as lcSSc or dcSSc (Table 382-2). Although stratification of SSc patients into diffuse and limited cutaneous subsets is useful, disease expression is far more complex, and several distinct endophenotypes exist within each subset. For example, 10–15% of patients with lcSSc develop PAH without significant ILD. Other patients have systemic features of SSc without appreciable skin involvement (SSc sine scleroderma). Unique clinical phenotypes of SSc associate with specific autoantibodies (Table 382-3). Patients with “overlap” have typical SSc features coexisting with clinical and laboratory evidence of another autoimmune disease such as polymyositis, Sjögren's syndrome, polyarthritis, autoimmune liver disease, or SLE.

TABLE 382-4 FREQUENCY OF CLINICAL ORGAN INVOLVEMENT: LIMITED CUTANEOUS AND DIFFUSE CUTANEOUS FORMS OF SYSTEMIC SCLEROSIS (SSC)

Features	Limited Cutaneous SSc (%)	Diffuse Cutaneous SSc (%)
Skin involvement	90 ^a	100
Raynaud's phenomenon	99	98
Esophageal involvement	90	80
Pulmonary fibrosis	35	65
Pulmonary arterial hypertension	15	15
Myopathy	11	23
Cardiac involvement	9	12
Scleroderma renal crisis	2	15

^aApproximately 10% of limited cutaneous SSc patients have SSc sine scleroderma.

INITIAL CLINICAL PRESENTATION

The initial presentation is quite different in the diffuse and the limited cutaneous forms of the disease. In dcSSc, the interval between Raynaud's phenomenon and onset of other disease manifestations is typically brief (weeks to months). Soft tissue swelling and intense pruritus are signs of the early inflammatory "edematous" phase. The fingers, hands, distal limbs, and face are usually affected first. Diffuse skin hyperpigmentation and carpal tunnel syndrome can occur. Arthralgias, muscle weakness, fatigue, and decreased joint mobility are common. During the ensuing weeks to months, the inflammatory edematous phase evolves into the "fibrotic" phase, with skin induration associated with hair loss, reduced production of skin oils, and a decline in sweating capacity. Progressive flexion contractures of the fingers ensue. The wrists, elbows, shoulders, hip girdles, knees, and ankles become stiff due to fibrosis of the supporting joint structures. While advancing skin involvement is the most visible manifestation of early dcSSc, important and frequently clinically silent internal organ involvement develops during this stage. The initial 4 years from disease onset is the period of rapidly evolving pulmonary and renal damage. If organ failure does not occur during this period, the systemic process may stabilize.

Compared to dcSSc, the course of lcSSc is characteristically more indolent. In these patients, the interval between Raynaud's phenomenon and onset of manifestations such as gastroesophageal reflux, cutaneous telangiectasia, or soft tissue calcifications can be several years. On the other hand, scleroderma renal crisis and severe pulmonary fibrosis are uncommon in lcSSc. Clinically evident cardiac involvement and PAH develop in more than 15%. Overlap with keratoconjunctivitis sicca, polyarthritis, cutaneous vasculitis, and biliary cirrhosis is seen in some patients with lcSSc.

ORGAN INVOLVEMENT

RAYNAUD'S PHENOMENON

Raynaud's phenomenon, the most frequent extracutaneous complication of SSc, is characterized by episodes of reversible vasoconstriction in the fingers and toes. Vasoconstriction may also affect the tip of the nose and earlobes. Attacks are triggered by a decrease in temperature, as well as emotional stress and vibration. Typical attacks start with pallor, followed by cyanosis of variable duration. Hyperemia ensues spontaneously or with rewarming of the digit. The progression of the three color phases reflects the underlying vasoconstriction, ischemia, and reperfusion.

As much as 3–5% of the general population has Raynaud's phenomenon. In the absence of signs or symptoms of an underlying condition, Raynaud's phenomenon is classified as primary and represents an exaggerated physiologic response to cold. Secondary Raynaud's phenomenon can occur as a complication of SSc and other connective tissue diseases, hematologic and endocrine conditions, and occupational disorders, and can complicate the use of beta blockers and anticancer drugs such as cisplatin and bleomycin. Distinguishing primary versus secondary Raynaud's phenomenon presents a diagnostic challenge. Primary Raynaud's phenomenon is supported by the following: absence of an underlying cause; a family history of Raynaud's phenomenon; absence of digital tissue necrosis, ulceration, or gangrene; and a negative ANA test. Secondary Raynaud's phenomenon tends to develop at an older age (>30 years), is clinically more severe (episodes more frequent, prolonged, and painful), and is frequently associated with ischemic digital ulcers and loss of digits (Fig. 382-3). Nailfold capillaroscopy, where the cutaneous capillaries at the nail bed are viewed under a drop of grade B immersion oil using a low-power stereoscopic microscope, can be helpful in the evaluation of Raynaud's phenomenon. Primary Raynaud's phenomenon is associated with normal capillaries that appear as regularly spaced parallel vascular loops, whereas in patients with Raynaud's associated with SSc and other connective tissue diseases, nailfold capillaries are distorted with widened and irregular loops, dilated lumen, and areas of vascular "dropout." In addition to digits, cold-induced episodic Raynaud's-like vasospasm



FIGURE 382-3 Digital necrosis. Sharply demarcated necrosis of the fingertip in a patient with limited cutaneous systemic sclerosis (SSc) associated with severe Raynaud's phenomenon.

has been documented in the pulmonary, renal, gastrointestinal, and coronary circulations in SSc.

SKIN FEATURES

While early-stage SSc is associated with edematous skin changes, skin thickening is the hallmark that distinguishes SSc from other connective tissue diseases. The distribution of skin thickening is invariably symmetric and bilateral. It typically starts in the fingers and then characteristically advances from distal to proximal extremities in an ascending fashion. The involved skin is firm, coarse, and thickened, and the extremities and trunk may be darkly pigmented. In some patients, diffuse tanning in the absence of sun exposure is a very early manifestation of skin involvement. In dark-skinned patients, vitiligo-like hypopigmentation may occur. Because pigment loss spares the perifollicular areas, the skin may have a "salt-and-pepper" appearance, most prominently on the scalp, upper back, and chest. Dermal sclerosis due to collagen accumulation obliterates hair follicles, sweat glands, and eccrine and sebaceous glands, resulting in hair loss, decreased sweating, and dry skin. Transverse creases on the dorsum of the fingers disappear (Fig. 382-4). Fixed flexion contractures of the fingers cause reduced hand mobility and lead to muscle atrophy. Skin thickening in combination with fibrosis of the subjacent tendons accounts for contractures of the wrists, elbows, and knees. Thick ridges at the neck due



FIGURE 382-4 Sclerodactyly. Note skin induration on the fingers, and fixed flexion contractures at the proximal interphalangeal joints in a patient with limited cutaneous systemic sclerosis (SSc).



A



B

FIGURE 382-5 Cutaneous vascular changes. **A.** Capillary changes at the nailfold in a patient with limited cutaneous systemic sclerosis (lcSSc). **B.** Telangiectasia on the face.

to firm adherence of skin to the underlying platysma muscle interfere with neck extension. The face assumes a characteristic “mauskopf” appearance with taut and shiny skin, loss of wrinkles, and occasionally an expressionless facies due to reduced mobility of the eyelids, cheeks, and mouth. Thinning of the lips with accentuation of the central incisor teeth and fine wrinkles (radial furrowing) around the mouth complete the picture. Reduced oral aperture (microstomia) interferes with eating and oral hygiene. The nose assumes a pinched, beak-like appearance.

In established SSc, the skin is firmly bound to the subcutaneous fat (tethering) and undergoes thinning and atrophy. Telangiectasias are dilated skin capillaries 2–20 mm in diameter frequently seen in lcSSc. These lesions, reminiscent of hereditary hemorrhagic telangiectasia, are prominent on the face, hands, lips, and oral mucosa (Fig. 382-5). A greater number of telangiectasias correlates with the extent of microvascular complications, including PAH. Breakdown of atrophic skin leads to chronic ulcerations at the extensor surfaces of the proximal interphalangeal joints, the volar pads of the fingertips, and bony prominences such as the elbows and malleoli. Ulcers are painful and may become secondarily infected, resulting in osteomyelitis. Healing of ischemic fingertip ulcerations leaves characteristic fixed digital “pits.” Loss of soft tissue at the fingertips due to ischemia is frequent and may be associated with striking resorption of the terminal phalanges (acro-osteolysis) (Fig. 382-6).

Calcium deposits (calcinosis) in the skin and soft tissues occur in patients with lcSSc who are positive for anticentromere antibodies. The deposits, varying in size from tiny punctate lesions to large conglomerate masses, are composed of calcium hydroxyapatite crystals and can be readily visualized on plain x-rays. Frequent locations include the finger pads, palms, extensor surfaces of the forearms, and



FIGURE 382-6 Acro-osteolysis. Note dissolution of terminal phalanges in a patient with long-standing limited cutaneous systemic sclerosis (lcSSc) and Raynaud’s phenomenon.

the olecranon and prepatellar bursae (Fig. 382-7). They may occasionally ulcerate through the overlying skin, producing drainage of chalky white material, pain, and local inflammation. Paraspinal soft tissue calcifications may cause neurologic complications.

PULMONARY FEATURES

Pulmonary involvement is frequent in SSc and is the leading cause of death. The two principal forms are ILD and pulmonary vascular disease. Patients with SSc frequently develop some degree of both complications. Less frequent pulmonary manifestations include aspiration pneumonitis complicating chronic gastroesophageal reflux, pulmonary hemorrhage due to endobronchial telangiectasia, obliterative bronchiolitis, pleural reactions, restrictive ventilatory defect due to chest wall fibrosis, spontaneous pneumothorax, and drug-induced lung toxicity. The incidence of lung cancer is increased.

Interstitial Lung Disease (ILD) Evidence of ILD can be found in up to 90% of patients with SSc at autopsy and 85% by thin-section high-resolution computed tomography (HRCT). In contrast, clinically significant ILD develops in 16–43%; the frequency varies depending on the detection method used. Risk factors include male gender, African-American race, diffuse skin involvement, severe gastroesophageal reflux, and the presence of topoisomerase I autoantibodies, as well as a low forced vital capacity (FVC) or single-breath diffusing capacity of the lung for carbon monoxide (DLCO) at initial presentation. In these patients, the most rapid progression in lung disease occurs early in



FIGURE 382-7 Calcinosis cutis. Note large calcific deposit breaking through the skin in a patient with limited cutaneous systemic sclerosis (lcSSc).

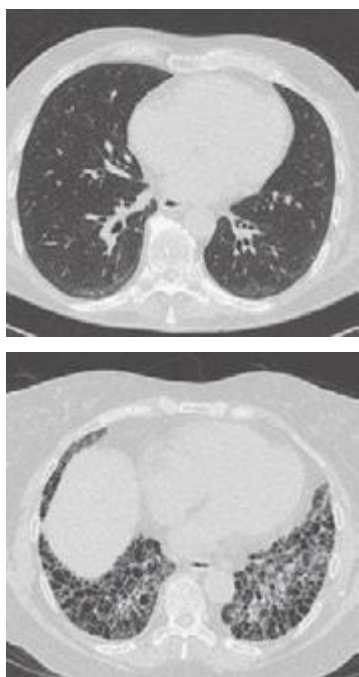


FIGURE 382-8 HRCT images of the chest from patients with systemic sclerosis. **Top panel:** Early interstitial lung disease. Mild changes with sub pleural reticulations and ground glass opacities in the lower lobes of the lung. Patient in supine position. **Bottom panel:** Extensive lung fibrosis with ground glass opacities, coarse reticular honeycombing, and traction bronchiectasis. (Courtesy of Rishi Agrawal, MD.)

the course of the disease (within the first 3 years), when the FVC can decline by 30% per year.

Pulmonary involvement can remain asymptomatic until it is advanced. The most common presenting respiratory symptoms—exertional dyspnea, fatigue, and reduced exercise tolerance—are subtle and slowly progressive. A chronic dry cough may be present. Physical examination may reveal “Velcro” crackles at the lung bases. Pulmonary function testing (PFT) is a sensitive method for detecting early pulmonary involvement. The most common abnormalities are reductions in FVC, total lung capacity (TLC), and DLCO. A reduction in DLCO that is significantly out of proportion to the reduction in lung volumes should raise suspicion for pulmonary vascular disease, but may also be due to anemia. Oxygen desaturation with exercise is common.

Chest radiography can rule out infection and other causes of pulmonary involvement, but compared to HRCT, it is relatively insensitive for detection of early ILD. HRCT shows subpleural reticular linear opacities and ground-glass opacifications, predominantly in the lower lobes, even in asymptomatic patients (Fig. 382-8). Additional HRCT findings include mediastinal lymphadenopathy, pulmonary nodules, traction bronchiectasis, and uncommonly, honeycomb changes. The extent of pulmonary interstitial changes on HRCT is a predictor of mortality in SSc. Bronchoalveolar lavage (BAL) can demonstrate inflammation in the lower respiratory tract and may be useful for ruling out infection. Although an elevated proportion of neutrophils (>2%) and/or eosinophils (>3%) in the BAL fluid is correlated with more extensive lung disease on HRCT and is associated with more rapid decline in FVC and reduced survival, BAL is not useful for identifying reversible alveolitis. Lung biopsy is indicated only in patients with atypical findings on chest radiographs and should be thoracoscopically guided. The histologic pattern on lung biopsy may predict the risk of progression of ILD. The most common pattern in SSc, NSIP, carries a better prognosis than UIP.

Pulmonary Arterial Hypertension (PAH) PAH, defined as a mean pulmonary arterial pressure ≥ 25 mmHg with a pulmonary capillary wedge pressure ≤ 15 mmHg, develops in approximately 15% of patients with SSc and can occur in association with ILD or as an isolated abnormality. The natural history of SSc-associated PAH is variable, but in many patients,

it follows a downhill course with development of right heart failure. The median survival of SSc patients with untreated PAH is 1 year following diagnosis. Risk factors for PAH include limited cutaneous disease, older age at disease onset, severe Raynaud’s phenomenon, and the presence of antibodies to centromere, U1-RNP, U3-RNP (fibrillarin), and B23.

The initial symptom of PAH is typically exertional dyspnea and reduced exercise capacity. With disease progression, angina, exertional near-syncope, and symptoms and signs of right-sided heart failure appear. Physical examination may show tachypnea, a prominent split S_2 heart sound, palpable right ventricular heave, elevated jugular venous pressure, and dependent edema. Doppler echocardiography provides a noninvasive method for estimating the pulmonary arterial pressure. In light of the poor prognosis of untreated PAH, all SSc patients should be screened for its presence at initial evaluation. Echocardiographic estimates of pulmonary arterial systolic pressures >40 mmHg at rest suggest PAH. Pulmonary function testing may show a reduced DLCO in isolation or out of proportion with the severity of restriction. Right heart catheterization is the gold standard for diagnosing PAH. Because echocardiography can result in over- or underestimation of pulmonary arterial pressures in SSc, cardiac catheterization is always required to confirm the presence of PAH; accurately assess its severity, including the degree of right heart dysfunction; and rule out venoocclusive disease and other cardiac causes of pulmonary hypertension. Yearly echocardiographic screening for PAH is recommended in most patients with SSc; an isolated decline in DLCO may also be indicative of developing PAH. Serum levels of brain natriuretic peptide (BNP) and N-terminal pro-BNP correlate with the presence and severity of PAH in SSc, as well as survival. While BNP measurements can be useful in screening for PAH and in monitoring the response to treatment, elevated BNP levels are not specific for PAH and also occur in other forms of right and left heart disease. The prognosis of SSc-associated PAH is worse, and treatment response poorer, than in idiopathic PAH.

GASTROINTESTINAL INVOLVEMENT

The gastrointestinal tract is affected in up to 90% of SSc patients with both limited and diffuse cutaneous forms of the disease. The pathologic features of atrophy of smooth muscle, intact mucosa, and obliterative small-vessel vasculopathy are similar throughout the length of the gastrointestinal tract.

Upper Gastrointestinal Tract Involvement Oropharyngeal manifestations due to a combination of xerostomia, reduced oral aperture, periodontal disease, and resorption of the mandibular condyles are frequent and cause much distress. The frenulum of the tongue may be shortened. Most patients have symptoms of gastroesophageal reflux disease (GERD): heartburn, regurgitation, and dysphagia. A combination of reduced lower esophageal sphincter pressure resulting in gastroesophageal reflux, impaired esophageal clearance of refluxed gastric contents due to diminished motility in the distal two-thirds of the esophagus, and delayed gastric emptying accounts for GERD. Manometry shows abnormal upper intestinal motility in most patients with SSc. Extragastric manifestations of GERD include hoarseness, chronic cough, and aspiration pneumonitis, which may aggravate underlying ILD. Chest computed tomography (CT) scan characteristically shows a dilated esophagus with intraluminal air. Endoscopy may be necessary to rule out opportunistic infections with *Candida*, herpes virus, and cytomegalovirus. Severe erosive esophagitis may be found on endoscopy in patients with minimal symptoms. Esophageal strictures and Barrett’s esophagus may complicate chronic GERD. Because Barrett’s esophagus is associated with an increased risk of adenocarcinoma, SSc patients with Barrett’s esophagus need periodic endoscopy and esophageal biopsy.

Gastroparesis with early satiety, abdominal distention, and aggravated reflux symptoms is common. The presence and severity of gastroparesis can be assessed by radionuclide gastric emptying studies. Gastric antral vascular ectasia (GAVE) in the antrum may occur. These subepithelial lesions, reflecting the diffuse small-vessel vasculopathy of SSc, are described as “watermelon stomach” due to their endoscopic appearance. Patients with GAVE can have recurrent episodes of gastrointestinal bleeding, resulting in chronic unexplained anemia.

2162 Lower Gastrointestinal Tract Involvement Impaired intestinal motility may result in malabsorption and chronic diarrhea secondary to bacterial overgrowth. Fat and protein malabsorption and vitamin B₁₂ and vitamin D deficiency ensue, sometimes culminating in severe malnutrition. Disturbed intestinal motor function can also cause intestinal pseudo-obstruction, with symptoms of nausea and abdominal distension that are indistinguishable from those of delayed gastric emptying. Patients present with recurrent episodes of acute abdominal pain, nausea, and vomiting. Radiographic studies show acute intestinal obstruction, and the major diagnostic challenge is to differentiate pseudoobstruction, which responds to supportive care and intravenous nutritional supplementation, from mechanical obstruction.

Colonic involvement may cause severe constipation, fecal incontinence, gastrointestinal bleeding from telangiectasia, and rectal prolapse. In late-stage SSc, wide-mouth sacculations or diverticula occur in the colon, occasionally causing perforation and bleeding. An occasional radiologic finding is pneumatosis cystoides intestinalis due to air trapping in the bowel wall that may rarely rupture and cause benign pneumoperitoneum. Although the liver is rarely affected, primary biliary cirrhosis may coexist with SSc.

RENAL INVOLVEMENT: SCLERODERMA RENAL CRISIS

Scleroderma renal crisis occurs in 10–15% of patients and generally within 4 years of the onset of the disease. Prior to the advent of angiotensin-converting enzyme (ACE) inhibitors, short-term survival in scleroderma renal crisis was <10%. The pathogenesis involves obliterative vasculopathy and luminal narrowing of the renal arcuate and interlobular arteries. Progressive reduction in renal blood flow, aggravated by vasospasm, leads to juxtaglomerular hyperplasia, increased renin secretion, and activation of angiotensin, with further renal vasoconstriction resulting in a vicious cycle that culminates in accelerated hypertension. Risk factors for scleroderma renal crisis include African-American race, male gender, and dcSSc with extensive and progressive skin involvement. Up to 50% of patients with scleroderma renal crisis have anti-RNA polymerase III antibodies. Palpable tendon friction rubs, pericardial effusion, new unexplained anemia, and thrombocytopenia may be harbingers of impending scleroderma renal crisis. High-risk patients with early SSc should be counseled to check their blood pressure daily. Patients with lcSSc or anticentromere antibodies rarely develop scleroderma renal crisis. Because there is an association between glucocorticoid use and scleroderma renal crisis, prednisone should be used in high-risk SSc patients only when absolutely required and at low doses (<10 mg/d).

Patients characteristically present with accelerated hypertension and progressive oliguric renal insufficiency. However, approximately 10% of patients with scleroderma renal crisis present with normal blood pressure. Normotensive renal crisis is generally associated with a poor outcome. Headache, blurred vision, and congestive heart failure may accompany elevation of blood pressure. Urinalysis typically shows mild proteinuria, granular casts, and microscopic hematuria; thrombocytopenia and microangiopathic hemolysis with fragmented red blood cells can be seen. Progressive oliguric renal failure over several days generally follows. In some cases, scleroderma renal crisis is misdiagnosed as thrombotic thrombocytopenic purpura or other forms of thrombotic microangiopathy. In these cases, a renal biopsy may be of some benefit. In addition, biopsy findings of vascular thrombosis and glomerular ischemic collapse predict poor renal outcomes. Oliguria or a creatinine >3 mg/dL at presentation predicts poor outcome, with permanent hemodialysis and high mortality. Rarely, crescentic glomerulonephritis occurs in the setting of SSc and may be associated with myeloperoxidase-specific antineutrophil cytoplasmic antibodies. Membranous glomerulonephritis may occur in patients treated with D-penicillamine. Asymptomatic renal function impairment occurs in up to half of SSc patients. Such subclinical renal involvement is associated with other vascular manifestations of SSc and rarely progresses.

CARDIAC INVOLVEMENT

Although it is often silent, cardiac involvement in SSc is frequently detected when patients are screened with sensitive diagnostic tools. Clinically evident cardiac involvement is associated with poor

outcomes. Cardiac disease in SSc may be primary or secondary to PAH, ILD, or renal involvement. It occurs more frequently in patients with dcSSc than in those with lcSSc and generally develops within 3 years of the onset of skin thickening. Clinically evident cardiac involvement in SSc is a poor prognostic factor. The endocardium, myocardium, and pericardium may each be affected separately or together. Manifestations of pericardial involvement include acute pericarditis, pericardial effusions, constrictive pericarditis, and cardiac tamponade. Conduction system fibrosis occurs commonly and may be silent or manifested by atrial and ventricular tachycardias or heart block. Recurrent vasospasm and ischemia-reperfusion injury contribute to myocardial fibrosis, resulting in asymptomatic systolic or diastolic left ventricular dysfunction that may progress to overt heart failure. Systemic and pulmonary hypertension and lung and renal involvement may also impact on the heart. Despite the presence of widespread obliterative vasculopathy, the frequency of clinical or pathologic epicardial coronary artery disease in SSc is not increased. While conventional echocardiography has low sensitivity for detecting SSc preclinical heart involvement, newer modalities such as tissue Doppler echocardiography (TDE), cardiac magnetic resonance imaging (cMRI), thallium perfusion, and nuclear imaging (single photon emission CT [SPECT]) reveal a high prevalence of abnormal myocardial function or perfusion in SSc patients. The serum level of N-terminal pro-BNP, a ventricular hormone, is a marker for PAH in SSc, but may also have utility as a marker of primary cardiac involvement.

MUSCULOSKELETAL COMPLICATIONS

Carpal tunnel syndrome occurs frequently and may be a presenting manifestation. Generalized arthralgia and stiffness are prominent in early disease. Mobility of small and large joints is progressively impaired, especially in dcSSc. Most commonly affected are the hands. Contractures develop at the proximal interphalangeal joints and wrists. Large joint contractures can be accompanied by tendon friction rubs, characterized by leathery crepitation that can be heard or palpated upon passive movement, that are due to extensive fibrosis and adhesion of the tendon sheaths and fascial planes at the affected joint. Presence of tendon friction rubs is associated with increased risk for renal and cardiac complications and reduced survival. True joint inflammation is uncommon; however, occasional patients develop erosive polyarthritis in the hands. Muscle weakness is common and may indicate deconditioning, disuse atrophy, and malnutrition. Less commonly, inflammatory myositis indistinguishable from idiopathic polymyositis may occur. A chronic noninflammatory myopathy characterized by atrophy and fibrosis in the absence of elevated muscle enzyme levels can be seen in late-stage SSc. Bone resorption occurs most commonly in the terminal phalanges, where it causes loss of the distal tufts (acroosteolysis) (Fig. 382-5). Resorption of the mandibular condyles can lead to bite difficulties. Osteolysis can also affect the ribs and distal clavicles.

OTHER DISEASE MANIFESTATIONS

Many SSc patients develop dry eyes and dry mouth (sicca complex). Biopsy of the minor salivary glands shows fibrosis rather than focal lymphocytic infiltration characteristic of primary Sjögren's syndrome (**Chap. 383**). Hypothyroidism is common and generally due to fibrosis of the thyroid gland. The frequency of macrovascular involvement, including peripheral vascular and coronary artery disease, may be increased. Whereas the central nervous system is generally spared, sensory trigeminal neuropathy due to fibrosis or vasculopathy can occur, presenting with gradual onset of pain and numbness. Pregnancy in women with SSc may be associated with an increased rate of adverse fetal outcomes. Furthermore, cardiopulmonary involvement may worsen during pregnancy, and new onset of scleroderma renal crisis has been described. Erectile dysfunction is frequent in men with SSc and may be the initial disease manifestation. Inability to attain or maintain penile erection is due to vascular insufficiency and fibrosis.

Malignancy in SSc Epidemiologic studies indicate an increased risk of cancer in SSc. Lung cancer and esophageal adenocarcinoma typically occur in the setting of long-standing ILD or gastroesophageal reflux disease and may be caused by chronic inflammation and repair. In contrast,

breast, lung, and ovarian carcinomas and lymphomas tend to occur in close temporal association with the clinical onset of SSc, particularly in patients who have autoantibodies to RNA polymerase III. In these cases, SSc may represent a paraneoplastic syndrome triggered by the anti-tumor immune response.

LABORATORY EVALUATION AND BIOMARKERS

A mild normocytic or microcytic anemia is frequent in patients with SSc and may indicate gastrointestinal bleeding caused by GAVE or chronic esophagitis. Macrocytic anemia may be caused by folate and vitamin B₁₂ deficiency due to small-bowel bacterial overgrowth and malabsorption or by drugs such as methotrexate or alkylating agents. Microangiopathic hemolytic anemia caused by mechanical fragmentation of red blood cells during their passage through microvessels coated with fibrin or platelet thrombi is a hallmark of the thrombotic microangiopathy associated with scleroderma renal crisis. Thrombocytopenia and leukopenia may indicate drug toxicity. In contrast to other connective tissue diseases, the erythrocyte sedimentation rate (ESR) is generally normal; an elevation may signal coexisting myositis or malignancy.

Antinuclear autoantibodies are present in almost all patients with SSc and can be detected at disease onset. Autoantibodies against topoisomerase I (Scl-70) and centromere are mutually exclusive and quite specific for SSc (Table 382-3). Topoisomerase I antibodies are detected in 31% of patients with dcSSc, but in only 13% of patients with lcSSc. They are associated with increased risk of ILD and poor outcomes. Anticentromere antibodies are detected in 38% of patients with lcSSc, but in only 2% of patients with dcSSc and rarely in patients with Raynaud's phenomenon and Sjögren's syndrome. Anticentromere antibodies in SSc are associated with PAH, but only infrequently with significant cardiac or renal involvement or ILD. Nucleolar immunofluorescence pattern on serologic testing reflects antibodies to U3-RNP (fibrillarin), Th/To, or PM/Scl, whereas a speckled immunofluorescence pattern indicates antibodies to RNA polymerase III. Although antibodies to β 2GPI occur in antiphospholipid antibody syndrome and are not specific for SSc, their presence in SSc is associated with an increased risk of ischemic lesions in the fingers.

DIAGNOSIS, STAGING, AND MONITORING

The diagnosis of SSc is made primarily on clinical grounds and is generally straightforward in patients with established disease. The presence of skin induration, with a characteristic symmetric distribution pattern associated with typical visceral organ manifestations, establishes the diagnosis with a high degree of certainty. Although the conditions listed in Table 382-1 can be associated with skin induration, the distribution pattern of skin lesions, together with the absence of Raynaud's phenomenon or typical visceral organ manifestations or SSc-specific autoantibodies, differentiates these conditions from SSc. Occasionally, full-thickness biopsy of the skin is required for establishing the diagnosis of scleredema, scleromyxedema, or nephrogenic systemic fibrosis. In lcSSc, a history of antecedent Raynaud's phenomenon and gastroesophageal reflux symptoms, coupled with the presence of sclerodactyly and capillary changes on nailfold capillaroscopy, often in combinations with cutaneous telangiectasia and calcinosis, helps to establish the diagnosis. The finding of digital tip pitting scars and radiologic evidence of pulmonary fibrosis in the lower lobes are particularly helpful diagnostically. Primary Raynaud's phenomenon is a common benign condition that must be differentiated from early or limited SSc. Nailfold microscopy is particularly helpful in this situation, because in primary Raynaud's phenomenon, the nailfold capillaries are normal, whereas in SSc, capillary abnormalities, as well as serum autoantibodies, can be detected even before other disease manifestations.

Establishing the diagnosis of SSc at an early stage of the disease may be a challenge. In dcSSc, initial symptoms are often nonspecific and relate to inflammation. Patients complain of fatigue, swelling, aching, and stiffness, and Raynaud's phenomenon may initially be absent. Physical examination may reveal diffuse upper extremity edema and puffy fingers. Patients at this stage are sometimes diagnosed as early rheumatoid arthritis, SLE, myositis, or, most commonly, undifferentiated connective tissue disease. Within weeks to months, Raynaud's phenomenon and characteristic clinical features appear accompanied by advancing induration of the skin. The presence of antinuclear and SSc-specific autoantibodies provides a high degree of diagnostic specificity. Raynaud's phenomenon

with fingertip ulcerations or other evidence of digital ischemia, coupled with telangiectasia, distal esophageal dysmotility, unexplained ILD or PAH, or accelerated hypertension with renal failure in the absence of clinically evident skin induration, suggests the diagnosis of SSc sine scleroderma. These patients may have anticentromere antibodies.

TREATMENT SYSTEMIC SCLEROSIS

OVERVIEW: MANAGEMENT PRINCIPLES To date, no therapy has been shown to significantly alter the natural history of SSc. In contrast, multiple interventions are highly effective in alleviating the symptoms, slowing the progression of the cumulative organ damage, and reducing disability. A significant reduction in disease-related mortality has been noted during the past 25 years. In light of the marked heterogeneity in disease manifestations, organ complications, and natural history, treatment must be tailored to each individual patient's unique needs.

A thorough investigation should be undertaken at baseline. Optimal management incorporates the following principles (Table 382-5): prompt and accurate diagnosis; classification and risk stratification based on clinical and laboratory evaluation; early recognition of organ-based complications and assessment of their extent, severity, and likelihood of deterioration; regular monitoring for disease progression, activity, new complications, and response to therapy; adjusting therapy; and continuing patient education. In order to minimize irreversible organ damage, the management of life-threatening complications must be proactive, with regular screening and initiation of appropriate intervention at the earliest possible opportunity. In light of the complex and multisystemic nature of the SSc, a team-based management approach integrating multiple specialists should be pursued whenever possible. Most patients are treated with combinations of drugs that impact different aspects of the disease. We encourage patients to become familiar with the spectrum of potential complications and understand therapeutic options and natural history, and empower them to partner with their treating physicians. This requires a long-term relationship between patient and physician, with ongoing counseling and encouragement.

DISEASE-MODIFYING THERAPY: IMMUNOSUPPRESSIVE AGENTS Immunosuppressive agents used in the treatment of other autoimmune or connective tissue diseases have generally shown modest or no benefit in SSc. Glucocorticoids may alleviate stiffness and aching in early-stage dcSSc but do not influence the progression of skin or internal organ involvement, and their use is associated with an increased risk of scleroderma renal crisis. Therefore, glucocorticoids should be given only when absolutely necessary, at the lowest dose possible, and for brief periods only. The use of cyclophosphamide has been extensively studied in light of its efficacy in the treatment of vasculitis (Chap. 385), SLE (Chap. 378), and other autoimmune diseases (Chap. 377e).

Both oral and intermittent IV cyclophosphamide were shown to reduce the progression of SSc-associated ILD, with stabilization and, rarely, modest improvement of pulmonary function and HRCT findings after 1 year of treatment. Improvement in respiratory symptoms and skin induration was also noted. These beneficial effects wane upon discontinuation of therapy. The benefits of cyclophosphamide need to be balanced against its potential toxicity, including bone marrow suppression, opportunistic infections, hemorrhagic cystitis and bladder cancer, premature ovarian failure, and late secondary malignancies.

TABLE 382-5 KEY PRINCIPLES IN MANAGEMENT

- Establish early and accurate diagnosis.
- Evaluate internal organ involvement.
- Define clinical disease stage and activity.
- Tailor individualized therapy to each patient's unique needs.
- Assess treatment response, and adjust therapy as needed; monitor for disease progression and new complications.

Methotrexate was associated with a modest skin improvement in small studies. Mycophenolate mofetil treatment was associated with improved skin induration in uncontrolled studies and was generally well tolerated. Small studies support the use of rituximab in SSc patients with skin involvement and ILD. The use of cyclosporine, azathioprine, extracorporeal photopheresis, thalidomide, rapamycin, imatinib, and IV immunoglobulin is currently not well supported by the literature. Intensive immune ablation using a conditioning regimen of high-dose chemotherapy with or without irradiation, followed by autologous stem cell reconstitution, has resulted in durable disease remission in some cases and is undergoing evaluation in randomized clinical trials. In light of its potential morbidity and mortality, as well as significant cost, autologous stem cell transplantation in SSc is still considered experimental.

Antifibrotic Therapy Because widespread tissue fibrosis in SSc causes progressive organ damage, drugs that interfere with the fibrotic process represent a rational therapeutic approach. D-Penicillamine has been extensively used as an antifibrotic agent. In retrospective studies, D-penicillamine stabilized and improved skin induration, prevented new internal organ involvement, and improved survival. However, a randomized controlled clinical trial in early active SSc found no difference in the extent of skin involvement between patients treated with standard-dose (750 mg/d) or very low-dose (125 mg every other day) D-penicillamine. Recent clinical trials show benefit of pirfenidone and of nintedanib in patients with idiopathic pulmonary fibrosis, with significant slowing of the loss of lung function. Whether these two new drugs will have comparable efficacy in the treatment of SSc-associated lung disease is still under investigation.

Vascular Therapy The goal of therapy is to control Raynaud's phenomenon, prevent the development and enhance the healing of ischemic complications, and slow the progression of obliterative vasculopathy. Patients should dress warmly, minimize cold exposure or stress, and avoid drugs that precipitate or exacerbate vasospastic episodes. Some patients with Raynaud's may respond to biofeedback therapy. Extended-release dihydropyridine calcium channel blockers such as nifedipine, amlodipine, or diltiazem can ameliorate Raynaud's phenomenon, but their use is often limited by side effects (palpitations, dependent edema, worsening gastroesophageal reflux). While ACE inhibitors do not reduce the frequency or severity of episodes, angiotensin II receptor blockers such as losartan are effective and generally well tolerated. Patients with Raynaud's phenomenon unresponsive to these therapies may require the addition of α_1 -adrenergic receptor blockers (e.g., prazosin), 5-phosphodiesterase inhibitors (e.g., sildenafil), serotonin reuptake inhibitors (e.g., fluoxetine), topical nitroglycerine, and intermittent infusions of IV prostaglandins. Low-dose aspirin and dipyridamole prevent platelet aggregation and may have a role as adjunctive agents. In patients with ischemic ulcers, the endothelin-1 receptor antagonist bosentan reduces the risk of new ulcers. Digital sympathectomy and local injections of botulinum type A (Botox) into the digits are options in patients with severe ischemia and impending loss of the digits. Empirical long-term therapy with statins and antioxidants may retard the progression of vascular damage and obliteration. Vasodilators such as ACE inhibitors, calcium channel blockers, and endothelin receptor blockers may also improve myocardial perfusion and left ventricular function.

TREATMENT OF GASTROINTESTINAL COMPLICATIONS Because oral problems including decreased oral aperture, decreased saliva production, gum recession and periodontal disease leading to teeth loss are common, regular dental care is recommended. Gastroesophageal reflux is very common and may occur in the absence of symptoms; therefore all patients with SSc should be treated. Patients should be instructed to elevate the head of the bed, eat frequent small meals, and avoid oral intake before bedtime. Proton pump inhibitors reduce acid reflux and may need to be given in relatively high doses. Prokinetic agents such as domperidone may be helpful, especially

if delayed gastric emptying is present. Episodic gastrointestinal bleeding from gastric antral vascular ectasia (watermelon stomach) may be amenable to treatment with endoscopic laser photocoagulation, although recurrence can occur. Bacterial overgrowth due to small-bowel dysmotility causes abdominal bloating and diarrhea and may lead to malabsorption and severe malnutrition. Treatment with short courses of rotating broad-spectrum antibiotics such as metronidazole, erythromycin, and tetracycline can eradicate bacterial overgrowth. Parenteral hyperalimentation is indicated if malnutrition develops. Chronic hypomotility of the small bowel may respond to octreotide, but pseudo-obstruction is difficult to treat. Fecal incontinence, a frequently underreported complication of SSc, may respond to anti-diarrheal medication and biofeedback therapy.

TREATMENT OF PULMONARY ARTERIAL HYPERTENSION (PAH) In patients with SSc, PAH carries an extremely poor prognosis and accounts for 30% of deaths. Because PAH is asymptomatic until advanced, patients with SSc should be screened for its presence at initial evaluation, and on a yearly basis thereafter. Treatment is generally started with an oral endothelin-1 receptor antagonist such as bosentan or a phosphodiesterase type 5 inhibitor such as sildenafil. Patients may also require diuretics and digoxin when appropriate. If hypoxemia is documented, supplemental oxygen should be prescribed in order to avoid hypoxia-induced secondary pulmonary vasoconstriction. Prostacyclin analogues such as epoprostenol or treprostinil can be given by continuous IV or SC infusion, or via intermittent nebulized inhalations. Combination therapy with different classes of agents, such as an endothelin-1 antagonist and a phosphodiesterase inhibitor, is often necessary. Lung transplantation remains an option for selected patients who fail medical therapy.

TREATMENT OF RENAL CRISIS Scleroderma renal crisis is a medical emergency. Since the outcome is largely determined by the extent of renal damage present at the time that aggressive therapy is initiated, prompt recognition of impending or early scleroderma renal crisis is essential, and efforts should be made to avoid its occurrence. High-risk SSc patients with early disease, extensive and progressive skin involvement, tendon friction rubs, and anti-RNA polymerase III antibodies should be instructed to monitor their blood pressure daily and report significant alterations immediately. Potentially nephrotoxic drugs should be avoided, and glucocorticoids should be used only when absolutely necessary and at low doses. Patients presenting with scleroderma renal crisis should be immediately hospitalized. Once other causes of renal disease are excluded, treatment should be started promptly with titration of short-acting ACE inhibitors, with the goal of achieving rapid normalization of the blood pressure. In patients with hypertension persisting despite ACE inhibitor therapy, addition of angiotensin II receptor blockers, calcium channel blockers, and direct renin inhibitors should be considered. Anecdotal evidence indicates responses to endothelin-1 receptor blockers and prostacyclins. Up to two-thirds of patients with scleroderma renal crisis go on to dialysis. The outcome of scleroderma renal crisis is worse in patients with antibodies to topoisomerase I compared to those with antibodies to RNA polymerase III. Substantial renal recovery can occur following scleroderma renal crisis, and dialysis can be discontinued, in 30–50% of the patients. Kidney transplantation is appropriate for those unable to discontinue dialysis after 2 years. Survival of transplanted SSc patients is comparable to that of patients with other connective tissue diseases, and recurrence of renal crisis is rare.

SKIN CARE Because skin involvement in SSc is never life-threatening and because it stabilizes and may even regress spontaneously, over time, the management of SSc should not be dictated by its cutaneous manifestations. The inflammatory symptoms of early skin involvement can be controlled with antihistamines and cautious short-term use of low-dose glucocorticoids (<5 mg/d of prednisone). Retrospective studies have shown that D-penicillamine

reduced the extent and progression of skin induration; however, these benefits could not be substantiated in a controlled prospective trial. Cyclophosphamide and methotrexate have modest effects on skin induration. Because the skin is dry, the use of hydrophilic ointments and bath oils is encouraged. Regular skin massage is helpful. Telangiectasia may present a cosmetic problem, especially on the face. Treatment with pulsed dye laser may have short-term benefit. Ischemic digital ulcers should be protected by occlusive dressing to promote healing and prevent infection. Infected skin ulcers are treated with topical antibiotics. Surgical debridement may be indicated. No therapy has been shown to be effective in preventing the formation of calcific soft tissue deposits or promoting their dissolution.

TREATMENT OF MUSCULOSKELETAL COMPLICATIONS Arthralgia and joint stiffness are common and distressing manifestations most prominent in early-stage disease. Short courses of nonsteroidal anti-inflammatory agents, weekly methotrexate, and cautious use of low-dose corticosteroids may alleviate these symptoms. Physical and occupational therapy can be effective for maintaining musculoskeletal function and improving long-term outcomes.

COURSE

The natural history of SSc is highly variable and difficult to predict, especially in early stages of the disease, when the specific subset—diffuse or limited cutaneous form—is not clear. Patient with dcSSc tend to have a more rapidly progressive course and worse prognosis than those with lcSSc.

In dcSSc, inflammatory symptoms such as fatigue, edema, arthralgia, and pruritus tend to subside, and the extent of skin thickening reaches a plateau at 2–4 years after disease onset, followed by slow regression. It is during the early edematous/inflammatory stage, generally lasting <3 years, that important visceral organ involvement occurs. While existing visceral organ involvement, such as pulmonary fibrosis, may progress even after skin involvement peaks, new organ involvement is rare. Scleroderma renal crisis almost invariably occurs within the first 4 years of disease. In late-stage disease (>6 years), the skin is usually soft and atrophic. Skin regression characteristically occurs in an order that is the reverse of initial involvement, with softening on the trunks followed by proximal and finally distal extremities; however, sclerodactyly and finger contractures generally persist. Relapse or recurrence of skin thickening after the peak of skin involvement has been reached is uncommon. Patients with lcSSc follow a clinical course that is markedly different than that of dcSSc. Raynaud's phenomenon typically precedes other disease manifestations by years or even decades. Visceral organ complications such as PAH and ILD generally develop late and progress slowly.

PROGNOSIS

SSc confers a substantial increase in the risk of premature death. Age- and gender-adjusted mortality rates are fivefold to eightfold higher compared to the general population, and more than half of all patients with SSc die from their disease. In one population-based study of SSc, the median survival was 11 years. In patients with dcSSc, 5- and 10-year survival rates are 70% and 55%, respectively, whereas in patients with lcSSc, 5- and 10-year survival rates are 90% and 75%, respectively. The prognosis correlates with the extent of skin involvement, which itself is a surrogate for visceral organ involvement. Major causes of death are PAH, pulmonary fibrosis, gastrointestinal involvement, and cardiac disease. Scleroderma renal crisis is associated with a 30% 3-year mortality. Lung cancer and excess cardiovascular deaths also contribute to increased mortality. Markers of poor prognosis include male gender, African-American race, older age at disease onset, extensive skin thickening with truncal involvement, palpable tendon friction rubs, and evidence of significant or progressive visceral organ involvement. Laboratory predictors of increased mortality at initial evaluation include an elevated ESR, anemia,

proteinuria and anti-topoisomerase I antibodies. In one study, SSc patients with extensive skin involvement, lung vital capacity <55% predicted, significant gastrointestinal involvement (pseudoostruction or malabsorption), evidence of cardiac involvement (arrhythmias or congestive heart failure), or scleroderma renal crisis had a cumulative 9-year survival <40%. The severity of PAH is strongly associated with mortality, and SSc patients who had a mean pulmonary arterial pressure ≥ 45 mmHg had a 33% 3-year survival. The advent of ACE inhibitors in scleroderma renal crisis had a dramatic impact on survival, increasing from <10% at 1 year in the pre-ACE inhibitor era to >70% 3-year survival at the present time. Moreover, 10-year survival in SSc has improved from <60% in the 1970s to >66–78% in the 1990s, a trend that reflects both earlier detection and better management of complications.

LOCALIZED SCLERODERMA

The term *scleroderma* is commonly used to describe a group of localized skin disorders (Table 382-1). These occur more commonly in children than in adults. In contrast to SSc, localized scleroderma is rarely complicated by Raynaud's phenomenon or significant internal organ involvement. Morphea presents as solitary or multiple circular patches of thickened skin or, rarely, as widespread induration (generalized or pansclerotic morphea); the fingers are spared. Linear scleroderma—streaks of thickened skin, typically in one or both lower extremities—may affect the subcutaneous tissues, leading to fibrosis and atrophy of supporting structures, muscle, and bone. In children, the growth of affected long bones can be retarded. When linear scleroderma lesions cross joints, significant contractures can develop.

MIXED CONNECTIVE TISSUE DISEASE

Patients who have lcSSc coexisting with features of SLE, polymyositis, and rheumatoid arthritis may have mixed connective tissue disease (MCTD). This overlap syndrome is generally associated with the presence of high titers of autoantibodies to U1-RNP. The characteristic initial presentation is Raynaud's phenomenon associated with puffy fingers and myalgia. Gradually, lcSSc features of sclerodactyly, calcinosis, and cutaneous telangiectasia develop. Skin rashes suggestive of SLE (malar rash, photosensitivity) or of dermatomyositis (heliotrope rash on the eyelids, erythematous rash on the knuckles) occur. Arthralgia is common, and some patients develop erosive polyarthritis. Pulmonary fibrosis and isolated or secondary PAH may develop. Other manifestations include esophageal dysmotility, pericarditis, Sjögren's syndrome, and renal disease, especially membranous glomerulonephritis. Laboratory evaluation indicates features of inflammation with elevated ESR and hypergammaglobulinemia. While anti-U1RNP antibodies are detected in the serum in high titers, SSc-specific autoantibodies are not found. In contrast to SSc, patients with MCTD often show a good response to treatment with glucocorticoids, and the long-term prognosis is better than that of SSc. Whether MCTD is a truly distinct entity or is, rather, a subset of SLE or SSc remains controversial.

EOSINOPHILIC FASCIITIS

Eosinophilic fasciitis is a rare idiopathic disorder associated with induration of the skin that generally develops rapidly. Adults are primarily affected. The skin has a coarse cobblestone "peau d'orange" appearance. In contrast to SSc, internal organ involvement is rare, and Raynaud's phenomenon and SSc-associated autoantibodies are absent. Furthermore, skin involvement spares the fingers. Full-thickness excisional biopsy of the lesional skin reveals fibrosis of the subcutaneous fascia and is generally required for diagnosis. Inflammation and eosinophil infiltration in the fascia are variably present. In the acute phase of the illness, peripheral blood eosinophilia may be prominent. MRI appears to be a sensitive tool for the diagnosis of eosinophilic fasciitis. In some patients, eosinophilic fasciitis occurs in association with, or preceding, myelodysplastic syndromes or multiple myeloma. Treatment with glucocorticoids leads to prompt resolution of the eosinophilia. In contrast, skin changes generally show slow and variable improvement. The prognosis of patients with eosinophilic fasciitis is good.

DEFINITION, INCIDENCE, AND PREVALENCE

Sjögren's syndrome is a chronic, slowly progressive autoimmune disease characterized by lymphocytic infiltration of the exocrine glands resulting in xerostomia and dry eyes. Approximately one-third of patients present with systemic manifestations; a small but significant number of patients develop malignant lymphoma. The disease presents alone (primary Sjögren's syndrome) or in association with other autoimmune rheumatic diseases (secondary Sjögren's syndrome) (Table 383-1).

Middle-aged women (female-to-male ratio, 9:1) are primarily affected, although Sjögren's syndrome may occur at any age, including childhood. The prevalence of primary Sjögren's syndrome is ~0.5–1%, while 30% of patients with autoimmune rheumatic diseases suffer from secondary Sjögren's syndrome.

PATHOGENESIS

Sjögren's syndrome is characterized by both lymphocytic infiltration of the exocrine glands and B lymphocyte hyperreactivity. An oligomonoclonal B cell process, which is characterized by cryoprecipitable monoclonal immunoglobulins (IgMκ) with rheumatoid factor activity, is evident in up to 25% of patients.

Sera from patients with Sjögren's syndrome often contain autoantibodies to non-organ-specific antigens such as immunoglobulins (rheumatoid factors) and extractable nuclear and cytoplasmic antigens (Ro/SS-A, La/SS-B). Ro/SS-A autoantigen consists of two polypeptides (52 and 60 kDa, respectively) in conjunction with cytoplasmic RNAs, whereas the 48-kDa La/SS-B protein is bound to RNA III polymerase transcripts. Autoantibodies to Ro/SS-A and La/SS-B antigens are usually detected at the time of diagnosis and are associated with earlier disease onset, longer disease duration, salivary gland enlargement, and more intense lymphocytic infiltration of minor salivary glands.

The major infiltrating cells in the affected exocrine glands are activated T and B lymphocytes. T cells predominate in mild lesions, whereas B cells are dominant in more severe lesions. Macrophages and dendritic cells also are found. The number of macrophages positive for interleukin (IL) 18 has been shown to correlate with parotid gland enlargement and low levels of the C4 component of complement, both of which are adverse predictors for lymphoma development.

Ductal and acinar epithelial cells appear to play a significant role in the initiation and perpetuation of autoimmune injury. These cells (1) express class II major histocompatibility complex (MHC) molecules, costimulatory molecules, and aberrant expression of intracellular autoantigens on cell membranes and thus are able to provide signals essential for lymphocytic activation; (2) inappropriately produce proinflammatory cytokines and lymphoattractant chemokines necessary for sustaining the autoimmune lesion and allowing progression to more sophisticated ectopic germinal center formation, which occurs in one-fifth of patients; and (3) express functional receptors of innate immunity, particularly Toll-like receptors (TLRs) 3, 7, and 9, that may account for the perpetuation of the autoimmune response.

TABLE 383-1 ASSOCIATION OF SJÖGREN'S SYNDROME WITH OTHER AUTOIMMUNE DISEASES

Rheumatoid arthritis
Systemic lupus erythematosus
Scleroderma
Mixed connective tissue disease
Primary biliary cirrhosis
Vasculitis
Chronic active hepatitis

Both infiltrating T and B cells have a tendency to be resistant to apoptosis. Levels of B cell-activating factor (BAFF) have been found to be elevated in patients with Sjögren's syndrome, especially those with hypergammaglobulinemia, and probably accounts for this antiapoptotic effect. Glandular epithelial cells seem to have an active role in the production of BAFF, which may be expressed and secreted after stimulation with type I interferon as well as with viral or synthetic double-stranded RNA. The triggering factor for epithelial activation appears to be a persistent enteroviral infection (possibly with coxsackievirus strains). Type I and type II interferon signatures have been described in ductal epithelial cells and T cells, respectively; their detection implies that interferons exert direct and cross-regulating effects on the pathogenic process.

A defect in cholinergic activity mediated through the M3 receptor and redistribution of the water-channel protein aquaporin 5, both leading to neuroepithelial dysfunction and diminished glandular secretions, have been proposed.

Molecular analysis of human leukocyte antigen (HLA) class II genes has revealed that Sjögren's syndrome, regardless of the patient's ethnic origin, is highly associated with the HLA DQA1*0501 allele. Genome-wide association studies have disclosed an increased prevalence of single-nucleotide polymorphisms in genes of IRF-5 and STAT-4, which participate in the activation of the type I interferon pathway.

CLINICAL MANIFESTATIONS

The majority of patients with Sjögren's syndrome have symptoms related to diminished lacrimal and salivary gland function. In most patients, the primary syndrome runs a slow and benign course. The initial manifestations can be mucosal or nonspecific dryness, and 8–10 years may elapse from the initial symptoms to full-blown development of the disease.

The principal oral symptom of Sjögren's syndrome is dryness (xerostomia). Patients report difficulty in swallowing dry food, an inability to speak continuously, a burning sensation, an increase in dental caries, and problems in wearing complete dentures. Physical examination shows a dry, erythematous, sticky oral mucosa. There is atrophy of the filiform papillae on the dorsum of the tongue, and saliva from the major glands is either not expressible or cloudy. Enlargement of the parotid or other major salivary glands occurs in two-thirds of patients with primary Sjögren's syndrome but is uncommon in those with the secondary syndrome. Diagnostic tests include sialometry, sialography, and scintigraphy. Newer imaging techniques, including ultrasound, MRI, and magnetic resonance sialography of the major salivary glands, are also being used. Biopsy of the labial minor salivary gland permits histopathologic confirmation of focal lymphocytic infiltrates.

Ocular involvement is the other major manifestation of Sjögren's syndrome. Patients usually describe a sandy or gritty feeling under the eyelids. Other symptoms include burning, accumulation of secretions in thick strands at the inner canthi, decreased tearing, redness, itching, eye fatigue, and increased photosensitivity. These symptoms, which define *keratoconjunctivitis sicca*, are attributed to the destruction of corneal and bulbar conjunctival epithelium. Diagnostic evaluation of keratoconjunctivitis sicca includes measurement of tear flow by Schirmer I test and determination of tear composition, with assessment of tear breakup time or tear lysozyme content. Slit-lamp examination of the cornea and conjunctiva after rose bengal staining reveals punctate corneal ulcerations and attached filaments of corneal epithelium.

Involvement of other exocrine glands, which occurs less frequently, includes a decrease in mucous gland secretions of the upper and lower respiratory tree, resulting in dry nose, throat, and trachea (xerotrachea). In addition, diminished secretion of the exocrine glands of the gastrointestinal tract leads to esophageal mucosal atrophy, atrophic gastritis, and subclinical pancreatitis. Dyspareunia due to dryness of the external genitalia and dry skin also may occur.

Extraglandular (systemic) manifestations are seen in one-third of patients with Sjögren's syndrome (Table 383-2) but are very rare in patients whose Sjögren's syndrome is associated with rheumatoid

TABLE 383-2 PREVALENCE OF EXTRAGLANDULAR MANIFESTATIONS IN PRIMARY SJÖGREN'S SYNDROME

Clinical Manifestation	Percent	Remarks
Arthralgias/arthritis	60	Usually non-erosive, leading to Jaccoud's arthropathy.
Raynaud's phenomenon	37	In one-third of patients, precedes sicca manifestations.
Lymphadenopathy	14	Lymphoma should be excluded.
Lung involvement	14	Small airway disease is the predominant pathology.
Vasculitis	11	The most common clinical manifestation is cutaneous palpable purpura.
Kidney involvement	9	Interstitial kidney disease is usually asymptomatic. Glomerulonephritis is associated with cryoglobulinemia.
Liver involvement	6	Primary biliary cirrhosis stage I.
Lymphoma	6	Glandular MALT ^a lymphoma is most common.
Peripheral neuropathy	2	Polyneuropathy, either sensory or sensorimotor.
Myositis	1	Sporadic causes of myositis and inclusion body myositis have been reported.

^aMucosa-associated lymphoid tissue.

arthritis. Patients with primary Sjögren's syndrome more often report easy fatigability, low-grade fever, Raynaud's phenomenon, myalgias, and arthralgias. Most patients with primary Sjögren's syndrome experience at least one episode of non-erosive arthritis during the course of their disease. Manifestations of pulmonary involvement are frequently evident histologically but are rarely important clinically. Dry cough is the major manifestation that is attributed to small airway disease. Renal involvement includes interstitial nephritis, clinically manifested by hyposthenuria and renal tubular dysfunction with or without acidosis. Untreated acidosis may lead to nephrocalcinosis. Glomerulonephritis is a rare finding that occurs in patients with mixed cryoglobulinemia or with systemic lupus erythematosus overlapping with Sjögren's syndrome. Vasculitis affects small and medium-sized vessels. The most common clinical features are purpura, recurrent urticaria, skin ulcerations, glomerulonephritis, and mononeuritis multiplex.

Different autoantibodies may determine the clinical expression of the disease. Patients positive for anticentromere autoantibody present with a clinical picture similar to that of limited scleroderma (Chap. 382). Antimitochondrial antibodies may connote liver involvement in the form of primary biliary cirrhosis (Chap. 369). Autoantibodies to 21-hydroxylase have recently been described in almost 20% of patients; their presence is associated with a blunted adrenal response.

Central nervous system involvement is rarely recognized. A few cases of myelitis associated with antibody to aquaporin 4 have been described.

Lymphoma is a well-known manifestation of Sjögren's syndrome that usually presents later in the illness. Persistent parotid gland enlargement, purpura, leukopenia, cryoglobulinemia, low C4 complement levels, and ectopic germinal centers in minor salivary gland biopsy samples are manifestations suggesting the development of lymphoma. It is interesting that the same risk factors account for glomerulonephritis and lymphoma and that these risk factors are the ones that confer increased mortality risk. Most lymphomas are extranodal, low-grade marginal-zone B cell lymphomas and are usually detected incidentally during evaluation of the labial biopsy. The affected lymph nodes are usually peripheral. Survival rates are decreased in patients with B symptoms, lymph node mass >7 cm in diameter, and high or intermediate histologic grade.

Routine laboratory tests in Sjögren's syndrome reveal mild normochromic, normocytic anemia. An elevated erythrocyte sedimentation rate is found in ~70% of patients.

TABLE 383-3 DIFFERENTIAL DIAGNOSIS OF SICCA SYMPTOMS

Xerostomia	Dry Eye	Bilateral Parotid Gland Enlargement
Viral infections	Inflammation	Viral infections
Drugs	Stevens-Johnson syndrome	Mumps
Psychotherapeutic	Pemphigoid	Influenza
Parasympatholytic	Chronic conjunctivitis	Epstein-Barr virus
Antihypertensive	Chronic blepharitis	Coxsackievirus A
Psychogenic origin	Sjögren's syndrome	Cytomegalovirus
Irradiation	Toxicity	HIV, HCV
Diabetes mellitus	Burns	Sarcoidosis
Trauma	Drugs	IgG4 syndrome
Sjögren's syndrome	Neurologic conditions	Sjögren's syndrome
Amyloidosis	Impaired lacrimal gland function	Metabolic disorders
	Impaired eyelid function	Diabetes mellitus
	Miscellaneous	Hyperlipoproteinemias
	Trauma	Chronic pancreatitis
	Hypovitaminosis A	Hepatic cirrhosis
	Blink abnormality	Endocrine
	Anesthetic cornea	Acromegaly
	Lid scarring	Gonadal hypofunction
	Epithelial irregularity	

DIAGNOSIS AND DIFFERENTIAL DIAGNOSIS

Primary Sjögren's syndrome is diagnosed if (1) the patient presents with eye and/or mouth dryness, (2) eye tests disclose keratoconjunctivitis sicca, (3) mouth evaluation reveals the classic manifestations of the syndrome, and/or (4) the patient's serum reacts with Ro/SS-A and/or La/SS-B autoantigens. Labial biopsy is needed when the diagnosis is uncertain or to rule out other conditions that may cause dry mouth or eyes or parotid gland enlargement (Tables 383-3 and 383-4). Validated diagnostic criteria have been established by a European study and have now been further improved by a European-American study group (Table 383-5). Hepatitis C virus infection should be ruled out since, apart from serologic tests, the clinicopathologic picture is almost identical to that of Sjögren's syndrome. Enlargement of major salivary glands, particularly in seronegative patients, should raise the suspicion of IgG4-related syndrome, which may present also as chronic pancreatitis, interstitial nephritis, retroperitoneal fibrosis, and aortitis.

TREATMENT SJÖGREN'S SYNDROME

Treatment of Sjögren's syndrome is aimed at symptom relief and limitation of the damaging local effects of chronic xerostomia and keratoconjunctivitis sicca through substitution for or stimulation of the missing secretions (Fig. 383-1).

TABLE 383-4 DIFFERENTIAL DIAGNOSIS OF SJÖGREN'S SYNDROME

HIV Infection and Sicca Syndrome	Sjögren's Syndrome	Sarcoidosis
Predominant in young males	Predominant in middle-aged women	No age or sex preference
Lack of autoantibodies to Ro/SS-A and/or La/SS-B	Presence of autoantibodies	Lack of autoantibodies to Ro/SS-A and/or La/SS-B
Lymphoid infiltrates of salivary glands by CD8+ T lymphocytes	Lymphoid infiltrates of salivary glands by CD4+ T lymphocytes	Granulomas in salivary glands
Association with HLA-DR5	Association with HLA-DR3 and DRw52	Unknown
Positive serologic tests for HIV	Negative serologic tests for HIV	Negative serologic tests for HIV

- I. Ocular symptoms: a positive response to at least one of three validated questions.
 1. Have you had daily, persistent, troublesome dry eyes for more than 3 months?
 2. Do you have a recurrent sensation of sand or gravel in the eyes?
 3. Do you use tear substitutes more than three times a day?
- II. Oral symptoms: a positive response to at least one of three validated questions.
 1. Have you had a daily feeling of dry mouth for more than 3 months?
 2. Have you had recurrent or persistently swollen salivary glands as an adult?
 3. Do you frequently drink liquids to aid in swallowing dry foods?
- III. Ocular signs: objective evidence of ocular involvement defined as a positive result to at least one of the following two tests:
 1. Shirmer's I test, performed without anesthesia (≤ 5 mm in 5 min)
 2. Rose Bengal score or other ocular dye score (≥ 4 according to van Bijsterveld's scoring system)
- IV. Histopathology: In minor salivary glands focal lymphocytic sialoadenitis, with a focus score ≥ 1 .
- V. Salivary gland involvement: objective evidence of salivary gland involvement defined by a positive result to at least one of the following diagnostic tests:
 1. Unstimulated whole salivary flow (≤ 1.5 mL in 15 min)
 2. Parotid sialography
 3. Salivary scintigraphy
- VI. Antibodies in the serum to Ro/SS-A or La/SS-B antigens, or both.

^aExclusion criteria: past head and neck radiation treatment, hepatitis C infection, AIDS, preexisting lymphoma, sarcoidosis, graft-versus-host disease, use of anticholinergic drugs. ^bPrimary Sjögren's syndrome: any four of the six items, as long as item IV (histopathology) or VI (serology) is positive; or any three of the four objective-criteria items (III, IV, V, VI). ^cIn patients with a potentially associated disease (e.g., another well-defined connective tissue disease), the presence of item I or item II plus any two from among items III, IV, and V may be considered indicative of secondary Sjögren's syndrome.

Source: From C Vitali et al: Ann Rheum Dis 61:554, 2002. ©2002 with permission from BMJ Publishing Group Ltd.

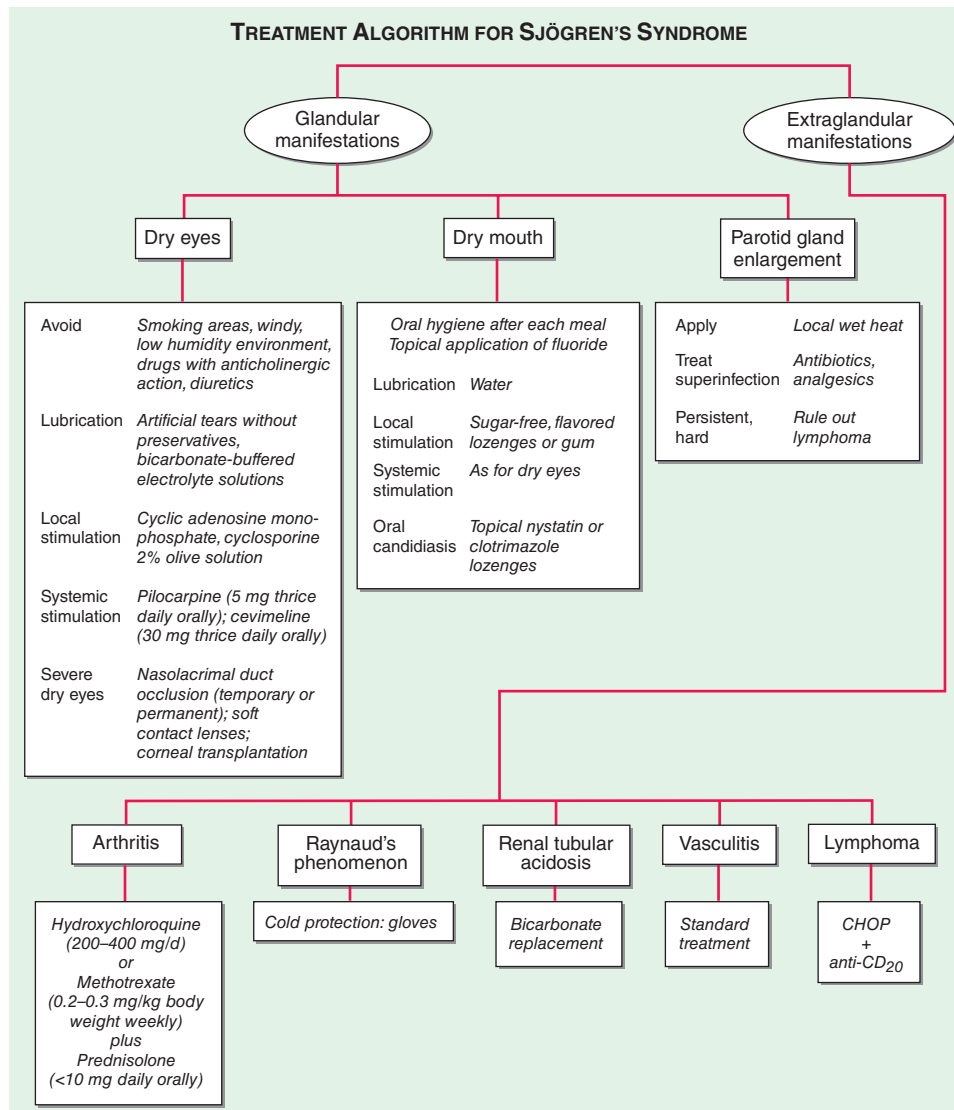


FIGURE 383-1 Treatment algorithm for Sjögren's syndrome. CHOP, cyclophosphamide, adriamycin (hydroxydaunorubicin), vincristine (oncovin), and prednisone.

To replace deficient tears, several ophthalmic preparations are readily available (hydroxypropyl methylcellulose; polyvinyl alcohol; 0.5% methylcellulose; Hypo Tears). If corneal ulcerations are present, eye patching and boric acid ointments are recommended. Certain drugs that may decrease lacrimal and salivary secretions, such as diuretics, antihypertensive drugs, anticholinergics, and antidepressants, should be avoided.

For xerostomia, the best replacement is water. Propionic acid gels may be used to treat vaginal dryness. To stimulate secretions, orally administered pilocarpine (5 mg thrice daily) or cevimeline (30 mg thrice daily) appears to improve sicca manifestations, and both are well tolerated. Hydroxychloroquine (200 mg) is helpful for arthralgias and mild arthritis.

Patients with renal tubular acidosis should receive sodium bicarbonate by mouth (0.5–2 mmol/kg in four divided doses). Glucocorticoids (1 mg/kg per day) and/or immunosuppressive agents (e.g., cyclophosphamide) are indicated only for the treatment of systemic vasculitis. Anti-tumor necrosis factor agents are ineffective. Monoclonal antibody to CD20 appears to be effective in patients with systemic disease, particularly in those with vasculitis, arthritis, and fatigue. Combination of anti-CD-20 with a classic CHOP regimen (cyclosporine, adriamycin [hydroxydaunorubicin], vincristine [oncovin], and prednisone) leads to increased survival rates among patients with high-grade lymphomas.

384 The Spondyloarthritis

Joel D. Taurog, John D. Carter

The spondyloarthritis are a group of overlapping disorders that share certain clinical features and genetic associations. These disorders include ankylosing spondylitis (AS), reactive arthritis, psoriatic arthritis and spondylitis, enteropathic arthritis and spondylitis, juvenile-onset spondyloarthritis (SpA), and undifferentiated SpA. The similarities in clinical manifestations and genetic predisposition suggest that these disorders share pathogenic mechanisms.

ANKYLOSING SPONDYLITIS

AS is an inflammatory disorder of unknown cause that primarily affects the axial skeleton; peripheral joints and extraarticular structures are also frequently involved. The disease usually begins in the second or third decade; male-to-female prevalence is between 2:1 and 3:1. The term *axial spondyloarthritis* is coming into common use, supported by criteria formulated in 2009 (Table 384-1). This classification includes both definite AS and early stages that do not yet meet classical criteria for AS, but it probably also includes other conditions with a different natural history.

EPIDEMIOLOGY

AS shows a striking correlation with the histocompatibility antigen HLA-B27 and occurs worldwide roughly in proportion to the prevalence of B27 (Chap. 373e). In North American whites, the prevalence of B27 is 7%, whereas it is 90% in patients with AS, independent of disease severity.

In population surveys, AS is present in 1–6% of adults inheriting B27, whereas the prevalence is 10–30% among B27+ adult first-degree relatives of AS probands. Concordance rate in identical twins is about 65%. Susceptibility to AS is determined largely by genetic factors, with B27 comprising less than one-half of the genetic component. Genome-wide single-nucleotide polymorphism (SNP) analysis has identified over 30 additional susceptibility alleles.

TABLE 384-1 ASAS CRITERIA FOR CLASSIFICATION OF AXIAL SPONDYLOARTHRITIS (TO BE APPLIED FOR PATIENTS WITH BACK PAIN ≥3 MONTHS AND AGE OF ONSET <45 YEARS)^a

Sacroiliitis on Imaging Plus ≥1 SpA Feature	or	HLA-B27 Plus ≥2 Other SpA Features
Sacroiliitis on imaging		SpA features
• Active (acute) inflammation on MRI highly suggestive of SpA-associated sacroiliitis ^b		• Inflammatory back pain ^d
and/or		• Arthritis ^e
• Definite radiographic sacroiliitis according to modified New York criteria ^f		• Enthesitis (heel) ^f
		• Anterior uveitis ^g
		• Dactylitis ^e
		• Psoriasis ^e
		• Crohn's disease or ulcerative colitis ^e
		• Good response to NSAIDs ^h
		• Family history of SpA ⁱ
		• HLA-B27
		• Elevated CRP ^j

^aSensitivity 83%, specificity 84%. The imaging arm (sacroiliitis) alone has a sensitivity of 66% and a specificity of 97%. ^bBone marrow edema and/or osteitis on short tau inversion recovery (STIR) or gadolinium-enhanced T1 image. ^cBilateral grade ≥2 or unilateral grade 3 or 4. ^dSee text for criteria. ^ePast or present, diagnosed by a physician. ^fPast or present pain or tenderness on examination at calcaneus insertion of Achilles tendon or plantar fascia. ^gPast or present, confirmed by an ophthalmologist. ^hSubstantial relief of back pain at 24–48 h after a full dose of NSAID. ⁱFirst- or second-degree relatives with ankylosing spondylitis (AS), psoriasis, uveitis, reactive arthritis (ReA), or inflammatory bowel disease (IBD). ^jAfter exclusion of other causes of elevated CRP.

Abbreviations: ASAS, Assessment of Spondyloarthritis International Society; CRP, C-reactive protein; MRI, magnetic resonance imaging; NSAIDs, nonsteroidal anti-inflammatory drugs; SpA, spondyloarthritis.

Source: From M Rudwaleit et al: Ann Rheum Dis 68:777, 2009. Copyright 2009, with permission from BMJ Publishing Group Ltd.

PATHOLOGY

Sacroiliitis is often the earliest manifestation of AS. Knowledge of its pathology comes from both biopsy and autopsy studies that cover a range of disease durations. Synovitis and myxoid marrow represent the earliest changes, followed by pannus and subchondral granulation tissue. Marrow edema, enthesitis, and chondroid differentiation are also found. Macrophages, T cells, plasma cells, and osteoclasts are prevalent. Eventually the eroded joint margins are gradually replaced by fibrocartilage regeneration and then by ossification. The joint may become totally obliterated.

In the spine, the specimens studied have either been surgically resected in advanced disease or taken from autopsies. There is inflammatory granulation tissue in the paravertebral connective tissue at the junction of annulus fibrosus and vertebral bone, and in some cases along the entire outer annulus. The outer annular fibers are eroded and eventually replaced by bone, forming the beginning of a syndesmophyte, which then grows by continued endochondral ossification, ultimately bridging the adjacent vertebral bodies. Ascending progression of this process leads to the “bamboo spine.” Other lesions in the spine include diffuse osteoporosis (loss of trabecular bone despite accretion of periosteal bone), erosion of vertebral bodies at the disk margin, “squaring” or “barreling” of vertebrae, and inflammation and destruction of the disk-bone border. Inflammatory arthritis of the apophyseal (facet) joints is common, with synovitis, inflammation at the bony attachment of the joint capsule, and subchondral bone marrow granulation tissue. Erosion of joint cartilage by pannus is often followed by bony ankylosis. This may precede formation of syndesmophytes bridging the adjacent disks. Bone mineral density is diminished in the spine and proximal femur early in the course of the disease.

Peripheral synovitis in AS shows marked vascularity, which is also evident as tortuous macrovasculature seen during arthroscopy. Lining layer hyperplasia, lymphoid infiltration, and pannus formation are also found. Central cartilaginous erosions caused by proliferation of subchondral granulation tissue are common. It should be

2170 emphasized that the characteristics of peripheral arthritis in AS and other forms of SpA are similar, and distinct from those of rheumatoid arthritis.

Inflammation in the fibrocartilaginous enthesis, the region where a tendon, ligament, or joint capsule attaches to bone, is a characteristic lesion in AS and other SpAs, both at axial and peripheral sites. Enthesitis is associated with prominent edema of the adjacent bone marrow and is often characterized by erosive lesions that eventually undergo ossification.

Subclinical intestinal inflammation has been found in the colon or distal ileum in a majority of patients with SpA. The histology is described below under “Enteropathic Arthritis.”

PATHOGENESIS

The pathogenesis of AS is immune-mediated, but there is little direct evidence for antigen-specific autoimmunity, and there is evidence to suggest more of an autoinflammatory pathogenesis. Uncertainty remains regarding the primary site of disease initiation. The dramatic response of the disease to therapeutic blockade of tumor necrosis factor α (TNF- α) indicates that this cytokine plays a central role in the immunopathogenesis of AS. Other genes related to TNF pathways show association with AS, including *TNFRSF1A*, *LTBR*, and *TBKBP1*. More recent evidence strongly implicates the interleukin (IL) 23/IL-17 cytokine pathway in AS pathogenesis. At least five genes in this pathway show association with AS, including *IL23R*, *PTER4*, *IL12B*, *CARD9*, and *TYK2*. All of these genes are also associated with inflammatory bowel disease (IBD), and three of them are associated with psoriasis. Serum levels of IL-23 and IL-17 are elevated in AS patients. Mice expressing high levels of IL-23 show spontaneous infiltration in the entheses of CD3+CD4-CD8- cells bearing IL-23 receptors and producing IL-17 and IL-22. This finding suggests the possibility that site-specific innate immune cells may play a critical role in the anatomic specificity of the lesions. Mast cells and, to a lesser extent, neutrophils appear to be the major IL-17-producing cells in peripheral arthritis, whereas neutrophils producing IL-17 are prominent in apophyseal joints. High levels of circulating $\gamma\delta$ T cells expressing IL-23 receptors and producing IL-17 have been found in AS patients.

Other associated genes encode other cytokines or cytokine receptors (*IL6R*, *IL1R1*, *IL1R2*, *IL7R*, *IL27*), transcription factors involved in the differentiation of immune cells (*RUNX3*, *EOMES*, *BACH2*, *NKX2-3*, *TBX21*), or other molecules involved in activation or regulation of immune or inflammatory responses (*FCGR2A*, *ZMIZ1*, *NOS2*, *ICOSLG*).

The inflamed sacroiliac joint is infiltrated with CD4+ and CD8+ T cells and macrophages and shows high levels of TNF- α , particularly early in the disease. Abundant transforming growth factor β (TGF- β) has been found in more advanced lesions. Peripheral synovitis in AS and the other spondyloarthritides is characterized by neutrophils, macrophages expressing CD68 and CD163, CD4+ and CD8+ T cells, and B cells. There is prominent staining for intercellular adhesion molecule 1 (ICAM-1), vascular cell adhesion molecule 1 (VCAM-1), matrix metalloproteinase 3 (MMP-3), and myeloid-related proteins 8 and 14 (MRP-8 and MRP-14). Unlike rheumatoid arthritis (RA) synovium, citrullinated proteins and cartilage gp39 peptide-major histocompatibility complexes (MHCs) are absent. However, citrullinated proteins can be seen in the circulation.

No specific event or exogenous agent that triggers the onset of disease has been identified, although overlapping features with reactive arthritis and IBD and the involvement of the IL-23/IL-17 pathway suggest that enteric bacteria may play a role, and microdamage from mechanical stress at enthesial sites has also been implicated.

It is firmly established that HLA-B27 plays a direct role in AS pathogenesis, but its precise role at the molecular level remains unresolved. Rats transgenic for HLA-B27 develop arthritis and spondylitis, and this is unaffected by the absence of CD8. It thus appears that classical peptide antigen presentation to CD8+ T cells may not be the primary disease mechanism. However, the association of AS with ERAP1, which strongly influences the MHC class I peptide repertoire, is only found in B27+ patients, and this suggests that peptide binding to B27 is nonetheless important. The pairs of ERAP1 alleles found in AS patients show diminished peptidase activity,

compared with those found in healthy controls. The B27 heavy chain has an unusual tendency to misfold, a process that may be proinflammatory. Genetic and functional studies in humans have suggested a role for natural killer (NK) cells in AS, possibly through interaction with B27 heavy chain homodimers. SpA-prone B27 rats show defective dendritic cell function and share with AS patients a characteristic “reverse interferon” gene expression signature in antigen-presenting cells.

New bone formation in AS appears to be largely based on enchondral bone formation and occurs only in the periosteal compartment. It correlates with lack of regulation of the Wnt signaling pathway, which controls the differentiation of mesenchymal cells into osteophytes, by the inhibitors DKK-1 and sclerostin. Indirect evidence and data from animal models also implicate bone morphogenic proteins, hedgehog proteins, and prostaglandin E₂. There is sharp controversy as to whether vertebral new bone formation in AS is a sequela of inflammation or whether it arises independently of inflammation. The second hypothesis is based on the observation that syndesmophyte formation is not suppressed by anti-TNF- α therapy that potently suppresses inflammation. TNF- α is also a known inducer of DKK-1, which inhibits bone formation. Recent magnetic resonance imaging (MRI) studies suggest that it is vertebral inflammatory lesions that undergo metaplasia to fat (increased T1-weighted signal) that are the predominant site of subsequent syndesmophytes despite anti-TNF- α therapy, whereas early acute inflammatory lesions resolve. A recent study suggested that the rate of syndesmophyte formation decreases after >4 years of anti-TNF- α therapy.

CLINICAL MANIFESTATIONS

The symptoms of the disease are usually first noticed in late adolescence or early adulthood; the median age in Western countries is approximately 23 years. In 5% of patients, symptoms begin after age 40. The initial symptom is usually dull pain, insidious in onset, felt deep in the lower lumbar or gluteal region, accompanied by low-back morning stiffness of up to a few hours' duration that improves with activity and returns following inactivity. Within a few months, the pain has usually become persistent and bilateral. Nocturnal exacerbation of pain often forces the patient to rise and move around.

In some patients, bony tenderness (presumably reflecting enthesitis or osteitis) may accompany back pain or stiffness, whereas in others it may be the predominant complaint. Common sites include the costosternal junctions, spinous processes, iliac crests, greater trochanters, ischial tuberosities, tibial tubercles, and heels. Hip and shoulder (“root” joint) arthritis is considered part of the axial disease. Hip arthritis occurs in 25–35% of patients. Shoulder arthritis is much less common. Severe isolated hip arthritis or bony chest pain may be the presenting complaint, and symptomatic hip disease can dominate the clinical picture. Arthritis of peripheral joints other than the hips and shoulders, usually asymmetric, occurs in up to 30% of patients. Neck pain and stiffness from involvement of the cervical spine are usually relatively late manifestations but are occasionally dominant symptoms. Rare patients, particularly in the older age group, present with predominantly constitutional symptoms.

AS often has a juvenile onset in developing countries. Peripheral arthritis and enthesitis usually predominate, with axial symptoms supervening in late adolescence.

Initially, physical findings mirror the inflammatory process. The most specific findings involve loss of spinal mobility, with limitation of anterior and lateral flexion and extension of the lumbar spine and of chest expansion. Limitation of motion is usually out of proportion to the degree of bony ankylosis and is thought to possibly reflect muscle spasm secondary to pain and inflammation. Pain in the sacroiliac joints may be elicited either with direct pressure or with stress on the joints. In addition, there is commonly tenderness upon palpation of the posterior spinous processes and other sites of symptomatic bony tenderness.

The modified Schober test is a useful measure of lumbar spine flexion. The patient stands erect, with heels together, and marks are made on the spine at the lumbosacral junction (identified by a horizontal line between the posterosuperior iliac spines) and 10 cm above. The patient then bends forward maximally with knees fully extended, and the distance between

the two marks is measured. This distance increases by ≥ 5 cm in the case of normal mobility and by < 4 cm in the case of decreased mobility. Chest expansion is measured as the difference between maximal inspiration and maximal forced expiration in the fourth intercostal space in males or just below the breasts in females, with the patient's hands resting on or just behind the head. Normal chest expansion is ≥ 5 cm. Lateral bending measures the distance the patient's middle finger travels down the leg with maximal lateral bending. Normal is > 10 cm.

Limitation or pain with motion of the hips or shoulders is usually present if these joints are involved. It should be emphasized that early in the course of mild cases, symptoms may be subtle and nonspecific, and the physical examination may be unrevealing.

The course of the disease is extremely variable, ranging from the individual with mild stiffness and normal radiographs to the patient with a totally fused spine and severe bilateral hip arthritis, accompanied by severe peripheral arthritis and extraarticular manifestations. Pain tends to be persistent early in the disease and intermittent later, with alternating exacerbations and quiescent periods. In a typical severe untreated case with progression of the spondylitis to syndesmophyte formation, the patient's posture undergoes characteristic changes, with obliterated lumbar lordosis, buttock atrophy, and accentuated thoracic kyphosis. There may be a forward stoop of the neck or flexion contractures at the hips, compensated by flexion at the knees. Disease progression can be estimated clinically from loss of height, limitation of chest expansion and spinal flexion, and occiput-to-wall distance. Occasional individuals are encountered with advanced deformities who report having never had significant symptoms.

The factors most predictive of radiographic progression (see below) are the presence of existing syndesmophytes, high inflammatory markers, and smoking. In some but not all studies, onset of AS in adolescence and early hip involvement correlate with a worse prognosis. In women, AS tends to progress less frequently to total spinal ankylosis, although there may be an increased prevalence of isolated cervical ankylosis and peripheral arthritis. In industrialized countries, peripheral arthritis (distal to hips and shoulders) occurs in less than one-half of patients with AS, usually as a late manifestation, whereas in developing countries, the prevalence is much higher, with onset typically early in the disease course. Pregnancy has no consistent effect on AS, with symptoms improving, remaining the same, or deteriorating in one-third of pregnant patients, respectively.

The most serious complication of the spinal disease is spinal fracture, which can occur with even minor trauma to the rigid, osteoporotic spine. The lower cervical spine is most commonly involved. These fractures are often displaced and cause spinal cord injury. A recent survey suggested a $> 10\%$ lifetime risk of fracture. Occasionally, fracture through a diskovertebral junction and adjacent neural arch, termed *pseudoarthrosis*, most common in the thoracolumbar spine, can be an unrecognized source of persistent localized pain and/or neurologic dysfunction. Wedging of thoracic vertebrae is common and correlates with accentuated kyphosis.

The most common extraarticular manifestation is acute anterior uveitis, which occurs in up to 40% of patients and can antedate the spondylitis. Attacks are typically unilateral, causing pain, photophobia, and increased lacrimation. These tend to recur, often in the opposite eye. Cataracts and secondary glaucoma are not uncommon sequelae. Up to 60% of patients with AS have inflammation in the colon or ileum. This is usually asymptomatic, but frank IBD occurs in 5–10% of patients with AS (see "Enteropathic Arthritis," below). About 10% of patients meeting criteria for AS have psoriasis (see "Psoriatic Arthritis," below). Aortic insufficiency, sometimes leading to congestive heart failure, occurs in a small percentage of patients, occasionally early. Third-degree heart block may occur alone or together with aortic insufficiency. Subclinical pulmonary lesions and cardiac dysfunction may be relatively common. Cauda equina syndrome and upper pulmonary lobe fibrosis are rare late complications. Retroperitoneal fibrosis is a rare associated condition. Prostatitis has been reported to have an increased prevalence. Amyloidosis is rare (Chap. 137).

Several validated measures of disease activity and functional outcome are in widespread use in the study and management of AS,

particularly the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI) and the Ankylosing Spondylitis Disease Activity Score (ASDAS), both measures of disease activity; the Bath Ankylosing Spondylitis Functional Index (BASFI), a measure of limitation in activities of daily living; and several measures of radiographic changes. The Harris hip score, although not specific for AS, can be helpful. Despite persistence of the disease, most patients remain gainfully employed. Some but not all studies of survival in AS have suggested that AS shortens life span, compared with the general population. Mortality attributable to AS is largely the result of spinal trauma, aortic insufficiency, respiratory failure, amyloid nephropathy, or complications of therapy such as upper gastrointestinal hemorrhage. The impact of anti-TNF therapy on outcome and mortality is not yet known, except for significantly improved work productivity.

LABORATORY FINDINGS

No laboratory test is diagnostic of AS. In most ethnic groups, HLA-B27 is present in 80–90% of patients. Erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP) are often, but not always, elevated. Mild anemia may be present. Patients with severe disease may show an elevated alkaline phosphatase level. Elevated serum IgA levels are common. Rheumatoid factor, anti-cyclic citrullinated peptide (CCP), and antinuclear antibodies (ANAs) are largely absent unless caused by a coexistent disease, although ANAs may appear with anti-TNF therapy. Circulating levels of CD8+ T cells tend to be low, and serum matrix metalloproteinase 3 levels correlate with disease activity. Synovial fluid from peripheral joints in AS is nonspecifically inflammatory. In cases with restriction of chest wall motion, decreased vital capacity and increased functional residual capacity are common, but airflow is normal and ventilatory function is usually well maintained.

RADIOGRAPHIC FINDINGS

Radiographically demonstrable sacroiliitis, usually symmetric, is eventually present in AS. The earliest changes by standard radiography are blurring of the cortical margins of the subchondral bone, followed by erosions and sclerosis. Progression of the erosions leads to "pseudowidening" of the joint space; as fibrous and then bony ankylosis supervenes, the joints may become obliterated.

In the lumbar spine, progression of the disease leads to straightening, caused by loss of lordosis, and reactive sclerosis, caused by osteitis of the anterior corners of the vertebral bodies with subsequent erosion, leading to "squaring" or even "barreling" of one or more vertebral bodies. Progressive ossification leads to eventual formation of marginal syndesmophytes, visible on plain films as bony bridges connecting successive vertebral bodies anteriorly and laterally.

Years may elapse before unequivocal sacroiliac abnormalities are evident on plain radiographs, and consequently, MRI is being increasingly used in diagnosing AS. Active sacroiliitis is best visualized by dynamic MRI with fat saturation, either T2-weighted turbo spin-echo sequence or short tau inversion recovery (STIR) with high resolution, or T1-weighted images with contrast enhancement. These techniques sensitively identify early intraarticular inflammation, cartilage changes, and underlying bone marrow edema in sacroiliitis (Fig. 384-1). They are also highly sensitive for evaluation of acute and chronic spinal changes (Fig. 384-2).

Reduced bone mineral density can be detected by dual-energy x-ray absorptiometry of the femoral neck and the lumbar spine. Use of a lateral projection of the L3 vertebral body can prevent falsely elevated readings related to spinal ossification.

DIAGNOSIS

It is important to establish the diagnosis of early AS before the development of irreversible deformity. This goal presents a challenge for several reasons: (1) Back pain is very common, but AS is much less common; (2) an early presumptive diagnosis often relies on clinical grounds requiring considerable expertise; and (3) young individuals with symptoms of AS often do not seek medical care. The widely used modified New York criteria (1984) are based on the presence of definite radiographic sacroiliitis and are too insensitive in early

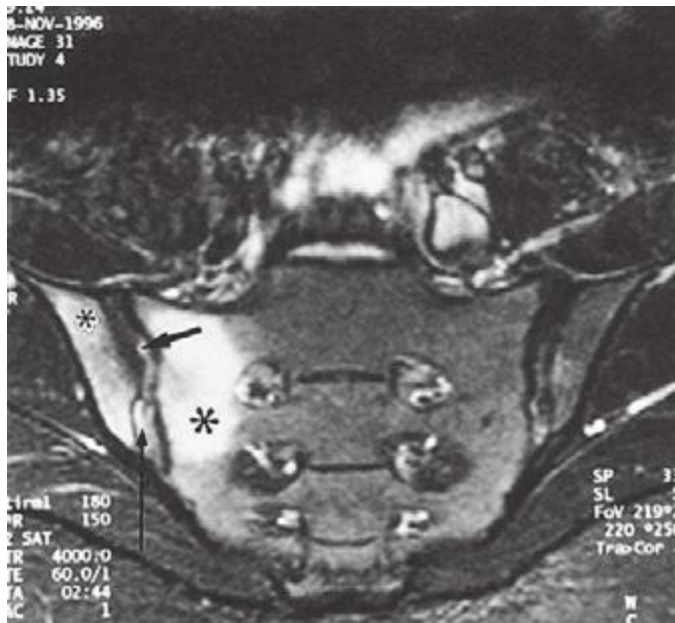


FIGURE 384-1 Early sacroiliitis in a patient with ankylosing spondylitis, indicated by prominent edema in the juxtaarticular bone marrow (asterisks), synovium and joint capsule (thin arrow), and interosseous ligaments (thick arrow) on a short tau inversion recovery (STIR) magnetic resonance image. (From M Bollow et al: *Zeitschrift für Rheumatologie* 58:61, 1999. Reproduced with permission.)

or mild cases. In 2009, new criteria for axial SpA were proposed by the Assessment of Spondyloarthritis International Society (ASAS) (Table 384-1). They are applicable to individuals with ≥ 3 months of back pain with age of onset < 45 years old. Active inflammation of the sacroiliac joints as determined by dynamic MRI is considered equivalent to definite radiographic sacroiliitis (see below).

AS must be differentiated from numerous other causes of low-back pain, some far more common than AS. To qualify as the criterion for inflammatory back pain of axial SpA (Table 384-1), the chronic (≥ 3 months) back pain should have four or more of the following characteristic features: (1) age of onset < 40 years old; (2) insidious onset; (3) improvement with exercise; (4) no improvement with rest; and (5) pain at night with improvement upon getting up. Other common

features of inflammatory back pain include morning stiffness > 30 min, awakening from back pain during only the second half of the night, and alternating buttock pain. In clinical decision-making, all of these features are additive. The most common causes of back pain other than AS are primarily mechanical or degenerative rather than primarily inflammatory and tend not to show clustering of these features.

Less-common metabolic, infectious, and malignant causes of back pain must also be differentiated from AS, including infectious spondylitis, spondylodiskitis, and sacroiliitis, and primary or metastatic tumor. Ochronosis can produce a phenotype that is clinically and radiographically similar to AS. Calcification and ossification of paraspinous ligaments occur in *diffuse idiopathic skeletal hyperostosis* (DISH), which occurs in the middle-aged and elderly and is usually not symptomatic. Ligamentous calcification gives the appearance of “flowing wax” on the anterior bodies of the vertebrae. Intervertebral disk spaces are preserved, and sacroiliac and apophyseal joints appear normal, helping to differentiate DISH from spondylosis and from AS, respectively.

TREATMENT ANKYLOSING SPONDYLITIS

All management of AS should include an exercise program designed to maintain posture and range of motion. Nonsteroidal anti-inflammatory drugs (NSAIDs) are the first line of pharmacologic therapy for AS. These agents reduce pain and tenderness and increase mobility in many patients with AS. There is mounting evidence that continuous high-dose NSAID therapy slows radiographic progression, particularly in patients who are at higher risk for progression. However, many patients with AS have continued symptoms despite NSAID therapy and are likely to benefit from anti-TNF- α therapy. Patients with AS treated with infliximab (chimeric human/mouse anti-TNF- α monoclonal antibody), etanercept (soluble p75 TNF- α receptor-IgG fusion protein), adalimumab, or golimumab (human anti-TNF- α monoclonal antibodies, or certolizumab pegol [humanized mouse anti-TNF- α monoclonal antibody]) have shown rapid, profound, and sustained reductions in all clinical and laboratory measures of disease activity. In a good response, there is significant improvement in both objective and subjective indicators of disease activity and function, including morning stiffness, pain, spinal mobility, peripheral joint swelling, CRP, and ESR. MRI studies indicate substantial resolution of bone marrow edema, enthesitis, and joint effusions in the sacroiliac joints, spine, and peripheral joints (Fig. 384-2). Similar results have been obtained in large randomized controlled trials of all four agents and

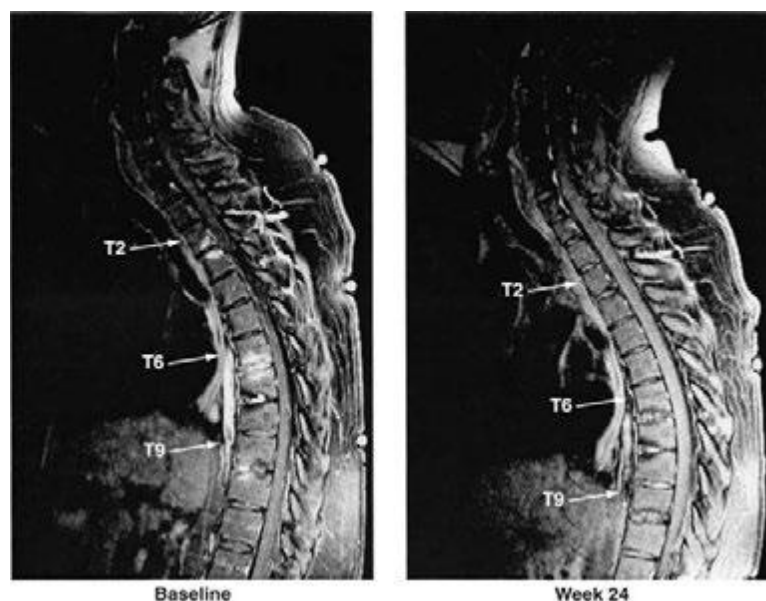


FIGURE 384-2 Spinal inflammation (spondylodiskitis) in a patient with ankylosing spondylitis and its dramatic response to treatment with infliximab. Gadolinium-enhanced T1-weighted magnetic resonance images, with fat saturation, at baseline and after 24 weeks of infliximab therapy. (From J Braun et al: *Arthritis Rheum* 54:1646, 2006.)

many open-label studies. About one-half of the patients achieve a $\geq 50\%$ reduction in the BASDAI. The response tends to be stable over time, and partial or full remissions are common. Predictors of the best responses include younger age, shorter disease duration, higher baseline inflammatory markers, and lower baseline functional disability. Nonetheless, some patients with long-standing disease and even spinal ankylosis can obtain significant benefit. Increased bone mineral density is found as early as 24 weeks after onset of therapy. There is evidence that anti-TNF therapy does not prevent syndesmophyte formation, although this may apply mainly during the early years of therapy. A mechanism for this has been proposed based on the observation that TNF- α inhibits new bone formation by upregulating DKK-1, a negative regulator of the wingless (Wnt) signaling pathway that promotes osteoblast activity. Serum DKK-1 levels are inappropriately low in AS patients and are also suppressed by anti-TNF therapy.

Infliximab is given intravenously, 3–5 mg/kg body weight, and then repeated 2 weeks later, again 6 weeks later, and then at 8-week intervals. Etanercept is given by subcutaneous injection, 50 mg once weekly. Adalimumab is given by subcutaneous injection, 40 mg biweekly. Golimumab is given by subcutaneous injection, 50 or 100 mg every 4 weeks. Certolizumab pegol is given by subcutaneous injection, 400 mg every 4 weeks.

Although these potent immunosuppressive agents have thus far been relatively safe, patients are at increased risk for serious infections, including disseminated tuberculosis. Hypersensitivity infusion or injection site reactions are not uncommon. Cases of anti-TNF-induced psoriasis have been increasingly recognized. Rare cases of systemic lupus erythematosus–related disease have been reported, as have hematologic disorders such as pancytopenia, demyelinating disorders, exacerbation of congestive heart failure, and severe liver disease. The overall incidence of malignancy does not appear to be increased in AS patients treated with anti-TNF therapy, but isolated cases of hematologic malignancy have occurred shortly after the start of treatment.

Because of the expense, potentially serious side effects, and unknown long-term effects of these agents, their use should be restricted to patients with a definite diagnosis and active disease (BASDAI ≥ 4 out of 10 and expert opinion) that is inadequately responsive to therapy with at least two different NSAIDs. Before initiation of anti-TNF therapy, all patients should be tested for tuberculin (TB) reactivity, and reactors (≥ 5 mm on PPD testing or a positive quantiferon test) should be treated with anti-TB agents. Contraindications include active infection or high risk of infection; malignancy or premalignancy; and history of systemic lupus erythematosus, multiple sclerosis, or related autoimmunity. Pregnancy and breast-feeding are relative contraindications. Continuation beyond 12 weeks of therapy requires either a 50% reduction in BASDAI or absolute reduction of ≥ 2 out of 10, and favorable expert opinion. Switching to a second anti-TNF agent may be effective, especially if there was a response to the first that was lost rather than primary failure. Sulfasalazine, in doses of 2–3 g/d, has been shown to be of modest benefit, primarily for peripheral arthritis. A therapeutic trial of this agent should precede any use of anti-TNF agents in patients with predominantly peripheral arthritis. Methotrexate, although widely used, has not been shown to be of benefit in AS, nor has any therapeutic role for gold or oral glucocorticoids been documented. Potential benefit in AS has been reported for thalidomide, 200 mg/d, perhaps acting through inhibition of TNF- α .

Ustekinumab (anti-IL-12/23) and secukinumab (anti-IL-17) monoclonal antibodies have shown promising efficacy in clinical trials, but have not yet been approved for use in AS.

The most common indication for surgery in patients with AS is severe hip joint arthritis, the pain and stiffness of which are usually dramatically relieved by total hip arthroplasty. Rare patients may benefit from surgical correction of extreme flexion deformities of the spine or of atlantoaxial subluxation.

Attacks of uveitis are usually managed effectively with local glucocorticoid administration in conjunction with mydriatic agents, although systemic glucocorticoids, immunosuppressive drugs,

or anti-TNF therapy may be required. TNF inhibitors reduce the frequency of attacks of uveitis in patients with AS, although cases of new or recurrent uveitis after use of a TNF inhibitor have been observed, especially with etanercept.

Coexistent cardiac disease may require pacemaker implantation and/or aortic valve replacement. Management of axial osteoporosis is at present similar to that used for primary osteoporosis, since data specific for AS are not available.

REACTIVE ARTHRITIS

Reactive arthritis (ReA) refers to acute nonpurulent arthritis complicating an infection elsewhere in the body. In recent years, the term has been used primarily to refer to SpA following enteric or urogenital infections.

Other forms of reactive and infection-related arthritis not associated with B27 and showing a spectrum of clinical features different from SpA, such as Lyme disease and rheumatic fever, are discussed in Chaps. 210 and 381.

HISTORIC BACKGROUND

The association of acute arthritis with episodes of diarrhea or urethritis has been recognized for centuries. A large number of cases during World Wars I and II focused attention on the triad of arthritis, urethritis, and conjunctivitis, often with additional mucocutaneous lesions, which became widely known by eponyms that are now of historic interest only.

The identification of bacterial species capable of triggering the clinical syndrome and the finding that many patients possess the B27 antigen led to the unifying concept of ReA as a clinical syndrome triggered by specific etiologic agents in a genetically susceptible host. A similar spectrum of clinical manifestations can be triggered by enteric infection with any of several *Shigella*, *Salmonella*, *Yersinia*, and *Campylobacter* species; by genital infection with *Chlamydia trachomatis*; and by other agents as well. The triad of arthritis, urethritis, and conjunctivitis represents a small part of the spectrum of the clinical manifestations of ReA and only a minority of patients present with this “classic triad” of symptoms. Although emerging data suggest that asymptomatic *Chlamydia trachomatis* infections might trigger ReA, for the purposes of this chapter, the use of the term *ReA* will be restricted to those cases of SpA in which there is at least presumptive evidence for a related symptomatic antecedent infection. Patients with clinical features of ReA who lack evidence of an antecedent infection will be considered to have *undifferentiated spondyloarthritis*, discussed below.

EPIDEMIOLOGY

Initial reports may have overestimated the association of ReA with HLA-B27, since 60–85% of patients who developed ReA triggered by *Shigella*, *Yersinia*, or *Chlamydia* were B27-positive. However, other studies demonstrated a lower prevalence of B27 in ReA triggered by *Salmonella*, with one study suggesting no association in *Campylobacter*-induced ReA. Several more recent community-based or common-source epidemic studies demonstrated that the prevalence of B27 in ReA was below 50%. The most common age range is 18–40 years, but ReA can occur rarely in children and occasionally in older adults.

The attack rate of postenteric ReA generally ranges from 1% to about 30% depending on the study and causative organism, whereas the attack rate of postchlamydial ReA is about 4–8%. The gender ratio in ReA following enteric infection is nearly 1:1, whereas venereally acquired ReA occurs mainly in men. The overall prevalence and incidence of ReA are difficult to assess because of the lack of validated diagnostic criteria, variable prevalence and arthritogenic potential of the triggering infectious agents, and inconstant genetic susceptibility factors in different populations. In Scandinavia, an annual incidence of 10–28:100,000 has been reported. The spondyloarthritides were formerly almost unknown in sub-Saharan Africa. However, ReA and other peripheral SpAs have now become the most common rheumatic diseases in Africans in the wake of the AIDS epidemic, without association to B27, which is very rare in these populations. ReA is often the first manifestation of HIV infection and often remits with disease progression. In contrast,

PATHOLOGY

Synovial histology is similar to that of other SpAs. Enthesitis shows increased vascularity and macrophage infiltration of fibrocartilage. Microscopic histopathologic evidence of inflammation mimicking IBD has routinely been demonstrated in the colon and ileum of patients with postenteric ReA and less commonly in postvenereal ReA. The skin lesions of keratoderma blennorrhagica, associated mainly with venereally acquired ReA, are histologically indistinguishable from pustular psoriatic lesions.

ETIOLOGY AND PATHOGENESIS

The bacteria identified as definitive triggers of ReA include several *Salmonella* spp., *Shigella* spp., *Yersinia enterocolitica*, *Yersinia pseudotuberculosis*, *Campylobacter jejuni*, and *Chlamydia trachomatis*. These triggering microbes are gram-negative bacteria with a lipopolysaccharide component to their cell walls. All four *Shigella* species (*Shigella sonnei*, *Shigella boydii*, *Shigella flexneri*, and *Shigella dysenteriae*) have been implicated in cases of ReA, with *S. flexneri* and *S. sonnei* being the most common. After *Salmonella* infection, individuals of Caucasian descent may be more likely than those of Asian descent to develop ReA. Children may be less susceptible to ReA caused by *Salmonella* and *Campylobacter*. *Yersinia* species in Europe and Scandinavia may have greater arthritogenic potential than in other parts of the world, and *C. trachomatis* appears to be a common cause worldwide. The ocular serovars of *C. trachomatis* appear to be particularly, perhaps uniquely, arthritogenic.

There is also evidence implicating *Clostridium difficile*, *Campylobacter coli*, certain toxigenic *Escherichia coli*, and possibly *Ureaplasma urealyticum* and *Mycoplasma genitalium* as potential triggers of ReA. *Chlamydia pneumoniae* is another trigger of ReA, albeit far less common than *C. trachomatis*. There have also been numerous isolated reports of acute arthritis preceded by other bacterial, viral, or parasitic infections, and even following intravesicular bacillus Calmette-Guérin (BCG) treatment for bladder cancer.

It has not been determined whether ReA occurs by the same pathogenic mechanism following infection with each of these microorganisms, nor has the mechanism been elucidated in the case of any one of the known bacterial triggers. Most, if not all, of the organisms well established to be triggers share a capacity to attack mucosal surfaces, to invade host cells, and to survive intracellularly. Antigens from *Chlamydia*, *Yersinia*, *Salmonella*, and *Shigella* have been shown to be present in the synovium and/or synovial fluid leukocytes of patients with ReA for long periods following the acute attack. In ReA triggered by *Y. enterocolitica*, bacterial lipopolysaccharide (LPS) and heat-shock protein antigens have been found in peripheral blood cells years after the triggering infection. *Yersinia* DNA and *C. trachomatis* DNA and RNA have been detected in synovial tissue from ReA patients, suggesting the presence of viable organisms despite uniform failure to culture the organism from these specimens. In *C. trachomatis*-induced ReA specifically, the bacterial load in synovial tissue of patients with remitting disease is lower than that of active disease, but mRNAs encoding proinflammatory proteins are equal to or higher than those of active disease. The specificity of these findings is unclear, however, since chromosomal bacterial DNA and 16S rRNA from a very wide variety of bacteria have also been found in synovium in other rheumatic diseases, albeit less frequently. In several older studies, synovial T cells that specifically responded to antigens of the inciting organism were reported and characterized as predominantly CD4+ with a T_H2 or T regulatory phenotype. More recent work has documented high levels of IL-17 in ReA synovial fluid, but the source has not been identified. HLA-B27 seems to be associated with more severe and chronic forms of the “classic triad” of ReA, but its pathogenic role remains to be determined. HLA-B27 significantly prolongs the intracellular survival of *Y. enterocolitica* and *Salmonella enteritidis* in human and mouse cell lines. Prolonged intracellular bacterial survival, promoted by B27, other factors, or both, may permit trafficking of infected leukocytes from the site

of primary infection to joints, where an innate and/or adaptive immune response to persistent bacterial antigens may then promote arthritis.

CLINICAL FEATURES

The clinical manifestations of ReA constitute a spectrum that ranges from an isolated, transient monoarthritis or enthesitis to severe multisystem disease. A careful history will usually elicit evidence of an antecedent infection 1–4 weeks before onset of symptoms of the reactive disease, particularly in postenteric ReA. However, in a sizable minority, no clinical or laboratory evidence of an antecedent infection can be found, particularly in the case of postchlamydial ReA. In cases of presumed venereally acquired reactive disease, there is often a history of a recent new sexual partner, even without laboratory evidence of infection.

Constitutional symptoms are common, including fatigue, malaise, fever, and weight loss. The musculoskeletal symptoms are usually acute in onset. Arthritis is usually asymmetric and additive, with involvement of new joints occurring over a few days to 1–2 weeks. The joints of the lower extremities, especially the knee, ankle, and subtalar, metatarsophalangeal, and toe interphalangeal joints, are most commonly involved, but the wrist and fingers can be involved as well. The arthritis is usually quite painful, and tense joint effusions are not uncommon, especially in the knee. Dactylitis, or “sausage digit,” a diffuse swelling of a solitary finger or toe, is a distinctive feature of ReA and other peripheral spondyloarthritides but can be seen in polyarticular gout and sarcoidosis. Tendinitis and fasciitis are particularly characteristic lesions, producing pain at multiple insertion sites (entheses), especially the Achilles insertion, the plantar fascia, and sites along the axial skeleton. Spinal, low-back, and buttock pain are quite common and may be caused by insertional inflammation, muscle spasm, acute sacroiliitis, or, presumably, arthritis in intervertebral joints.

Urogenital lesions may occur throughout the course of the disease. In males, urethritis may be marked or relatively asymptomatic and may be either an accompaniment of the triggering infection or a result of the reactive phase of the disease; interestingly, it occurs in both postvenereal and postenteric ReA. Prostatitis is also common. Similarly, in females, cervicitis or salpingitis may be caused either by the infectious trigger or by the sterile reactive process.

Ocular disease is common, ranging from transient, asymptomatic conjunctivitis to an aggressive anterior uveitis that occasionally proves refractory to treatment and may result in blindness.

Mucocutaneous lesions are frequent. Oral ulcers tend to be superficial, transient, and often asymptomatic. The characteristic skin lesions, *keratoderma blennorrhagica*, consist of vesicles and/or pustules that become hyperkeratotic, ultimately forming a crust before disappearing. They are most common on the palms and soles but may occur elsewhere as well. In patients with HIV infection, these lesions are often extremely severe and extensive, sometimes dominating the clinical picture (Chap. 226). Lesions may occur on the glans penis, termed *circinate balanitis*; these consist of vesicles that quickly rupture to form painless superficial erosions, which in circumcised individuals can form crusts similar to those of *keratoderma blennorrhagica*. Nail changes are common and consist of onycholysis, distal yellowish discoloration, and/or heaped-up hyperkeratosis.

Less-frequent or rare manifestations of ReA include cardiac conduction defects, aortic insufficiency, central or peripheral nervous system lesions, and pleuropulmonary infiltrates.

Arthritis typically persists for 3–5 months, but more chronic courses do occur. Chronic joint symptoms persist in about 15% of patients and in up to 60% of patients in hospital-based series, but these tend to be less severe than in the acute stage. Recurrences of the acute syndrome are also common. Work disability or forced change in occupation is common in those with persistent joint symptoms. Chronic heel pain is often particularly distressing. Low-back pain, sacroiliitis, and frank AS are also common sequelae. In most studies, HLA-B27-positive patients have shown a worse outcome than B27-negative patients. Patients with *Yersinia*- or *Salmonella*-induced arthritis have less chronic disease than those whose initial episode follows epidemic shigellosis.

LABORATORY AND RADIOGRAPHIC FINDINGS

The ESR and acute-phase reactants are usually elevated during the acute phase of the disease, often markedly so. Mild anemia may be present. Synovial fluid is nonspecifically inflammatory. In most ethnic groups, 30–50% of the patients are B27-positive. The triggering infection usually does not persist at the site of primary mucosal infection through the time of onset of the reactive disease, but it may be possible to culture the organism, e.g., in the case of *Yersinia*- or *Chlamydia*-induced disease. Serologic evidence of exposure to one of the causative organisms with elevation of antibodies is nonspecific and is of questionable utility. Polymerase chain reaction (PCR) for chlamydial DNA in first-voided urine specimens may have high sensitivity in the acute stage but is less useful with chronic disease.

In early or mild disease, radiographic changes may be absent or confined to juxtaarticular osteoporosis. With long-standing persistent disease, radiographic features share those of psoriatic arthritis; marginal erosions and loss of joint space can be seen in affected joints. Periostitis with reactive new bone formation is characteristic, as in all the SpAs. Spurs at the insertion of the plantar fascia are common.

Sacroiliitis and spondylitis may be seen as late sequelae. Sacroiliitis is more commonly asymmetric than in AS, and spondylitis, rather than ascending symmetrically, can begin anywhere along the lumbar spine. The syndesmophytes are described as nonmarginal; they are coarse, asymmetric, and “comma”-shaped, arising from the middle of a vertebral body, a pattern less commonly seen in primary AS. Progression to spinal fusion is uncommon.

DIAGNOSIS

ReA is a clinical diagnosis with no definitively diagnostic laboratory test or radiographic finding. The diagnosis should be entertained in any patient with an acute inflammatory, asymmetric, additive arthritis or tendinitis. The evaluation should include questioning regarding possible triggering events such as an episode of diarrhea or dysuria. On physical examination, attention must be paid to the distribution of the joint and tendon involvement and to possible sites of extraarticular involvement, such as the eyes, mucous membranes, skin, nails, and genitalia. Synovial fluid analysis may be helpful in excluding septic or crystal-induced arthritis. Culture, serology, or molecular methods may help to identify a triggering infection, but they cannot be relied upon.

Although typing for B27 has low negative predictive value in ReA, it may have prognostic significance in terms of severity, chronicity, and the propensity for spondylitis and uveitis. Furthermore, if positive, it can be helpful diagnostically in atypical cases. HIV testing is often indicated and may be necessary in order to select appropriate therapy.

It is important to differentiate ReA from disseminated gonococcal disease (Chap. 181), both of which can be venereally acquired and associated with urethritis. Unlike ReA, gonococcal arthritis and tenosynovitis tend to involve both upper and lower extremities equally, spare the axial skeleton, and be associated with characteristic vesicular skin lesions. A positive gonococcal culture from the urethra or cervix does not exclude a diagnosis of ReA; however, culturing gonococci from blood, skin lesion, or synovium establishes the diagnosis of disseminated gonococcal disease. PCR assay for *Neisseria gonorrhoeae* and *C. trachomatis* may be helpful. Occasionally, only a therapeutic trial of antibiotics can distinguish the two.

ReA shares many features in common with psoriatic arthropathy. However, psoriatic arthritis is usually gradual in onset; the arthritis tends to affect primarily the upper extremities; there is less associated periartthritis; and there are usually no associated mouth ulcers, urethritis, or bowel symptoms.

TREATMENT REACTIVE ARTHRITIS

Most patients with ReA benefit to some degree from high-dose NSAIDs, although acute symptoms are rarely completely ameliorated, and some patients fail to respond at all. Indomethacin, 75–150 mg/d in divided doses, is the initial treatment of choice, but other NSAIDs may be tried.

Prompt, appropriate antibiotic treatment of acute chlamydial urethritis or enteric infection may prevent the emergence of ReA, but is not universally successful. Data regarding the potential benefit of antibiotic therapy that is initiated after onset of arthritis are conflicting, but several trials suggest no benefit. One long-term follow-up study suggested that although antibiotic therapy had no effect on the acute episode of ReA, it helped prevent subsequent chronic SpA. Another such study failed to demonstrate any long-term benefit. A promising recent double-blind placebo-controlled study assessing combination antibiotics showed that a majority of patients with chronic ReA due to *Chlamydia* benefited significantly from a 6-month course of rifampin 300 mg daily plus azithromycin 500 mg daily for 5 days, then twice weekly, or 6 months of rifampin 300 mg daily plus doxycycline 100 mg twice daily. The possibility remains that acute *Chlamydia*-induced ReA might respond more favorably to antibiotic therapy than the postenteric variety.

Multicenter trials have suggested that sulfasalazine, up to 3 g/d in divided doses, may be beneficial to patients with persistent ReA.¹ Patients with persistent disease may respond to azathioprine, 1–2 mg/kg per day, or to methotrexate, up to 20 mg per week; however, these therapeutic regimens have never formally been studied. Although no controlled trials of anti-TNF- α in ReA have been reported, anecdotal evidence supports the use of these agents in severe chronic cases, although lack of response has also been observed.¹

Tendinitis and other enthesitic lesions may benefit from intral-lesional glucocorticoids. Uveitis may require aggressive treatment to prevent serious sequelae (see above). Skin lesions ordinarily require only symptomatic topical treatment. In patients with HIV infection and ReA, many of whom have severe skin lesions, the skin lesions in particular respond to antiretroviral therapy. Cardiac complications are managed conventionally; management of neurologic complications is symptomatic.

Comprehensive management includes counseling of patients in the avoidance of sexually transmitted disease and exposure to enteropathogens, as well as appropriate use of physical therapy, vocational counseling, and continued surveillance for long-term complications such as AS. Patients with a history of ReA are at increased risk for recurrent attacks following repeated exposures.

PSORIATIC ARTHRITIS

Psoriatic arthritis (PsA) refers to an inflammatory musculoskeletal disease that has both autoimmune and autoinflammatory features characteristically occurring in individuals with psoriasis.

HISTORIC BACKGROUND

The association between arthritis and psoriasis was noted in the nineteenth century. In the 1960s, on the basis of epidemiologic and clinical studies, it became clear that unlike RA, the arthritis associated with psoriasis was usually seronegative, often involved the distal interphalangeal (DIP) joints of the fingers and the spine and sacroiliac joints, had distinctive radiographic features, and showed considerable familial aggregation. In the 1970s, PsA was included in the broader category of the spondyloarthritides because of features similar to those of AS and ReA.

EPIDEMIOLOGY

Estimates of the prevalence of PsA among individuals with psoriasis range from 5 to 42%. The prevalence of PsA appears to be increasing in parallel with disease awareness; recent data using screening tools suggest that 20% or more of patients with psoriasis have undiagnosed PsA. The duration and severity of psoriasis increase one's likelihood of developing PsA. In white populations, psoriasis is estimated to have a prevalence of

¹Azathioprine, methotrexate, sulfasalazine, pamidronate, and thalidomide have not been approved for this purpose by the U.S. Food and Drug Administration at the time of publication.

2176 1–3%. Psoriasis and PsA are less common in other races in the absence of HIV infection, and the prevalence of PsA in individuals with psoriasis may be less common. First-degree relatives of PsA patients have an elevated risk for psoriasis, for PsA itself, and for other forms of SpA. Of patients with psoriasis, up to 30% have an affected first-degree relative. In monozygotic twins, the reported concordance for psoriasis varies from 35 to 72%, and for PsA from 10 to 30%. A variety of HLA associations have been found. The *HLA-Cw*0602* gene is directly associated with psoriasis, particularly familial juvenile-onset (type I) psoriasis. HLA-B27 is associated with psoriatic spondylitis (see below). HLA-DR7, -DQ3, and -B57 are associated with PsA because of linkage disequilibrium with Cw6. Other associations with PsA include HLA-B13, -B37, -B38, -B39, -C12, and -DR4. A recent genome-wide scan found association of both psoriasis and PsA with a polymorphism at the HCP5 locus closely linked to HLA-B, and also to IL-23R, IL-12B (chromosome 5q31), IL-13, and several other chromosomal regions. Certain genetic loci are associated with PsA but not psoriasis, e.g., RUNX3 and IL-13.

PATHOLOGY

The inflamed synovium in PsA resembles that of RA, although with somewhat less hyperplasia and cellularity than in RA. As noted with AS above, the synovial vascular pattern in PsA is generally greater and more tortuous than in RA, independent of disease duration. Some studies have indicated a higher tendency to synovial fibrosis in PsA. Unlike RA, PsA shows prominent enthesitis, with histology similar to that of the other spondyloarthritides.

PATHOGENESIS

PsA is almost certainly immune-mediated and probably shares pathogenic mechanisms with psoriasis. PsA synovium is characterized by lining layer hyperplasia; diffuse infiltration with T cells, B cells, macrophages, and NK receptor-expressing cells, with upregulation of leukocyte homing receptors; and neutrophil proliferation with angiogenesis. Clonally expanded T cell subpopulations are frequent and have been demonstrated both in the synovium and the skin. Plasmacytoid dendritic cells are thought to play a key role in psoriasis, and there is some evidence for their participation in PsA. There is abundant synovial overexpression of proinflammatory cytokines, and synovial tissue staining has identified an overexpression of monocyte-derived cytokines, such as myeloid-related protein (S100A8/A9). Interferon γ , TNF- α , and IL-1 β , -2, -6, -8, -10, -12, -13, and -15 are found in PsA synovium or synovial fluid. T_H17-derived cytokines are important in PsA, given the genetic association with genes in the IL-12/IL-23 axis and the therapeutic response to an antibody to the shared IL-12/23 p40 subunit (see below). T_H17 cells have been identified from the dermal extracts of psoriatic lesions and the synovial fluid of PsA patients. The majority of these CD4⁺ IL-17⁺ T cells are of memory phenotype (CD4RO[+]CD45RA[-]CD11a[+]). Consistent with the extensive bone remodeling in PsA, patients with PsA have been found to have a marked increase in osteoclastic precursors in peripheral blood and upregulation of receptor activator of nuclear factor κ B ligand (RANKL) in the synovial lining layer. Increased serum levels of TNF- α , RANKL, leptin, and omentin positively correlate with these osteoclastic precursors.

CLINICAL FEATURES

In 60–70% of cases, psoriasis precedes joint disease. In 15–20% of cases, the two manifestations appear within 1 year of each other. In about 15–20% of cases, the arthritis precedes the onset of psoriasis and can present a diagnostic challenge. The frequency in men and women is almost equal, although the frequency of disease patterns differs somewhat in the two sexes. The disease can begin in childhood or late in life but typically begins in the fourth or fifth decade, at an average age of 37 years.

The spectrum of arthropathy associated with psoriasis is quite broad. Many classification schemes have been proposed. In the original scheme of Wright and Moll, five patterns are described: (1) arthritis of the DIP joints; (2) asymmetric oligoarthritis; (3) symmetric polyarthritis similar to RA; (4) axial involvement (spine and sacroiliac joints); and (5) arthritis mutilans, a highly destructive form of



FIGURE 384-3 Characteristic lesions of psoriatic arthritis.

Inflammation is prominent in the distal interphalangeal joints (left 5th, 4th, 2nd; right 2nd, 3rd, and 5th) and proximal interphalangeal joints (left 2nd, right 2nd, 4th, and 5th). There is dactylitis in the left 2nd finger and thumb, with pronounced telescoping of the left 2nd finger. Nail dystrophy (hyperkeratosis and onycholysis) affects each of the fingers except the left 3rd finger, the only finger without arthritis. (Courtesy of Donald Raddatz, MD; with permission.)

disease. These patterns frequently coexist, and the pattern that persists chronically often differs from that of the initial presentation. A simpler scheme in recent use contains three patterns: oligoarthritis, polyarthritis, and axial arthritis.

Nail changes in the fingers or toes occur in up to 90% of patients with PsA, compared with 40% of psoriatic patients without arthritis, and pustular psoriasis is said to be associated with more severe arthritis. Several articular features distinguish PsA from other joint disorders; such hallmark features include dactylitis and enthesitis. Dactylitis occurs in >30%; enthesitis and tenosynovitis are also common and are probably present in most patients, although often not appreciated on physical examination. Shortening of digits because of underlying osteolysis is particularly characteristic of PsA (Fig. 384-3), and there is a much greater tendency than in RA for both fibrous and bony ankylosis of small joints. Rapid ankylosis of one or more proximal interphalangeal (PIP) joints early in the course of disease is not uncommon. Back and neck pain and stiffness are also common in PsA.

Arthropathy confined to the DIP joints occurs in about 5% of cases. Accompanying nail changes in the affected digits are almost always present. These joints are also often affected in the other patterns of PsA. Approximately 30% of patients have asymmetric oligoarthritis. This pattern commonly involves a knee or another large joint with a few small joints in the fingers or toes, often with dactylitis. Symmetric polyarthritis occurs in about 40% of PsA patients at presentation. It may be indistinguishable from RA in terms of the joints involved, but other features characteristic of PsA are usually also present. Almost any peripheral joint can be involved. Axial arthropathy without peripheral involvement is found in about 5% of PsA patients. It may be clinically indistinguishable from idiopathic AS, although more neck involvement and less thoracolumbar spinal involvement are characteristic, and nail changes are not found in idiopathic AS. A small percentage of PsA patients have arthritis mutilans, in which there can be widespread shortening of digits (“telescoping”), sometimes coexisting with ankylosis and contractures in other digits.

Six patterns of nail involvement are identified: pitting, horizontal ridging, onycholysis, yellowish discoloration of the nail margins, dystrophic hyperkeratosis, and combinations of these findings. Other extraarticular manifestations of the spondyloarthritides are common. Eye involvement, either conjunctivitis or uveitis, is reported in 7–33% of PsA patients. Unlike the uveitis associated with AS, the uveitis in PsA is more often bilateral, chronic, and/or posterior. Aortic valve insufficiency has been found in <4% of patients, usually after long-standing disease.

Widely varying estimates of clinical outcome have been reported in PsA. At its worst, severe PsA with arthritis mutilans is potentially at least as crippling and ultimately fatal as severe RA. Unlike RA, however, many patients with PsA experience temporary remissions. Overall, erosive disease develops in the majority of patients, progressive disease with deformity and disability is common, and in some large published series, mortality was found to be significantly increased compared with the general population. There appears to be a greater incidence of cardiovascular death in psoriatic disease.

The psoriasis and associated arthropathy seen with HIV infection both tend to be severe and can occur in populations with very little psoriasis in noninfected individuals. Severe enthesopathy, dactylitis, and rapidly progressive joint destruction are seen, but axial involvement is very rare. This condition is prevented by or responds well to antiretroviral therapy.

LABORATORY AND RADIOGRAPHIC FINDINGS

There are no laboratory tests diagnostic of PsA. ESR and CRP are often elevated. A small percentage of patients may have low titers of rheumatoid factor or antinuclear antibodies. About 10% of patients have anti-CCP antibodies. Uric acid may be elevated in the presence of extensive psoriasis. HLA-B27 is found in 50–70% of patients with axial disease, but in $\leq 20\%$ of patients with only peripheral joint involvement.

The peripheral and axial arthropathies in PsA show a number of radiographic features that distinguish them from RA and AS, respectively. Characteristics of peripheral PsA include DIP involvement, including the classic “pencil-in-cup” deformity; marginal erosions with adjacent bony proliferation (“whiskering”); small-joint ankylosis; osteolysis of phalangeal and metacarpal bone, with telescoping of digits; and periostitis and proliferative new bone at sites of enthesitis. Characteristics of axial PsA include asymmetric sacroiliitis. When compared with idiopathic AS, axial PsA manifests less zygapophyseal joint arthritis; nonmarginal, bulky, “comma”-shaped syndesmophytes that tend to be fewer and less symmetric and delicate than the marginal syndesmophytes of AS; fluffy hyperperiostosis on anterior vertebral bodies; severe cervical spine involvement, with a tendency to atlantoaxial subluxation but relative sparing of the thoracolumbar spine; and paravertebral ossification. Ultrasound and MRI both readily demonstrate enthesitis and tendon sheath effusions that can be difficult to assess on physical examination. A recent MRI study of 68 PsA patients found sacroiliitis in 35%, unrelated to B27 but correlated with restricted spinal movement.

DIAGNOSIS

Classification criteria for PsA were published in 2006 (Classification of Psoriatic Arthritis [CASPAR] criteria) that have been widely accepted (Table 384-2). The sensitivity and specificity of these criteria exceed 90%, and they are useful for early diagnosis. The

TABLE 384-2 THE CASPAR (CLASSIFICATION CRITERIA FOR PSORIATIC ARTHRITIS) CRITERIA^a

To meet the CASPAR criteria, a patient must have inflammatory articular disease (joint, spine, or enthesal) with ≥ 3 points from any of the following five categories:

1. Evidence of current psoriasis,^{b,c} a personal history of psoriasis, or a family history of psoriasis^d
2. Typical psoriatic nail dystrophy^e observed on current physical examination
3. A negative test result for rheumatoid factor
4. Either current dactylitis^f or a history of dactylitis recorded by a rheumatologist
5. Radiographic evidence of juxtaarticular new bone formation^g in the hand or foot

^aSpecificity of 99% and sensitivity of 91%. ^bCurrent psoriasis is assigned 2 points; all other features are assigned 1 point. ^cPsoriatic skin or scalp disease present at the time of examination, as judged by a rheumatologist or dermatologist. ^dHistory of psoriasis in a first- or second-degree relative. ^eOnycholysis, pitting, or hyperkeratosis. ^fSwelling of an entire digit. ^gIll-defined ossification near joint margins, excluding osteophyte formation.

Source: From W Taylor et al: Arthritis Rheum, 54:2665, 2006.

criteria are based on the history, presence of psoriasis, characteristic peripheral or spinal joint symptoms, signs, and imaging. Diagnosis can be challenging when the arthritis precedes psoriasis, the psoriasis is undiagnosed or obscure, or the joint involvement closely resembles another form of arthritis. A high index of suspicion is needed in any patient with an undiagnosed inflammatory arthropathy. The history should include inquiry about psoriasis in the patient and family members. Patients should be asked to disrobe for the physical examination, and psoriasiform lesions should be sought in the scalp, ears, umbilicus, and gluteal folds in addition to more accessible sites; the finger and toe nails should also be carefully examined. Axial symptoms or signs, dactylitis, enthesitis, ankylosis, the pattern of joint involvement, and characteristic radiographic changes can be helpful clues. The differential diagnosis includes all other forms of arthritis, which can occur coincidentally in individuals with psoriasis. The differential diagnosis of isolated DIP involvement is short. Osteoarthritis (Heberden’s nodes) is usually not inflammatory; gout involving more than one DIP joint often involves other sites and may be accompanied by tophi; the very rare entity multicentric reticulohistiocytosis involves other joints and has characteristic small pearly periungual skin nodules; and the uncommon entity inflammatory osteoarthritis, like the others, lacks the nail changes of PsA. Radiography can be helpful in all of these cases and in distinguishing between psoriatic spondylitis and idiopathic AS. A history of trauma to an affected joint preceding the onset of arthritis is said to occur more frequently in PsA than in other types of arthritis, perhaps reflecting the Koebner phenomenon in which psoriatic skin lesions can arise at sites of the skin trauma.

TREATMENT PSORIATIC ARTHRITIS

Ideally, coordinated therapy is directed at both the skin and joints in PsA. As described above for AS, use of the anti-TNF- α agents has revolutionized the treatment of PsA. Prompt and dramatic resolution of both arthritis and skin lesions has been observed in large, randomized controlled trials of etanercept, infliximab, adalimumab, and golimumab. Many of the responding patients had long-standing disease that was resistant to all previous therapy, as well as extensive skin disease. The clinical response is often more dramatic than in RA, and delay of disease progression has been demonstrated radiographically. The potential additive effect of methotrexate to anti-TNF- α agents in PsA remains uncertain. As noted above, anti-TNF therapy, paradoxically, has been reported to trigger exacerbation or de novo appearance of psoriasis, typically the palmoplantar pustular variety. In some cases, the therapy can nevertheless be continued.

Ustekinumab, a monoclonal antibody to the shared IL-23/IL-12p40 subunit, is an efficacious treatment for psoriasis and has shown promise in PsA in clinical trials. Other newer drugs that have shown efficacy for both psoriasis and PsA include the anti-IL-17 pathway agents, such as secukinumab and brodalumab, and an oral phosphodiesterase-4 inhibitor, apremilast. Data on the oral Jak inhibitor, tofacitinib, has been very limited but promising.

Other treatment for PsA has been based on drugs that have efficacy in RA and/or in psoriasis. Until recently, controlled clinical trial data on methotrexate in doses of 15–25 mg/week and sulfasalazine (usually given in doses of 2–3 g/d) suggesting clinical efficacy have been somewhat limited, but neither regimen effectively halts progression of erosive joint disease. A recent double-blind trial assessing methotrexate 15 mg weekly in PsA demonstrated no benefit to the joint-based inflammation, but improvement was seen in patient and assessor global scores and skin scores. Other agents with efficacy in psoriasis reported to benefit PsA are cyclosporine, retinoic acid derivatives, and psoralens plus ultraviolet A light (PUVA). There is controversy regarding the efficacy in PsA of gold and antimalarials, which have been widely used in RA. The pyrimidine synthetase inhibitor leflunomide has been shown in a randomized controlled trial to be beneficial in both psoriasis and PsA.

2178 All of these treatments require careful monitoring. Immunosuppressive therapy may be used cautiously in HIV-associated PsA if the HIV infection is well controlled.

UNDIFFERENTIATED AND JUVENILE-ONSET SPONDYLOARTHRITIS

Many patients, usually young adults, present with some features of one or more of the spondyloarthritides discussed above. Until recently, these patients were said to have *undifferentiated spondyloarthritis*, or simply *spondyloarthritis*, as defined by the 1991 European Spondyloarthropathy Study Group criteria. For example, a patient may present with inflammatory synovitis of one knee, Achilles tendinitis, and dactylitis of one digit. Some of these patients may have ReA in which the triggering infection remains clinically silent. In some other cases, the patient subsequently develops IBD or psoriasis, or the process eventually meets criteria for AS. This diagnosis of undifferentiated SpA was also commonly applied to patients with inflammatory back pain, who did meet modified New York criteria for AS. Most of these would now be classified under the new category of axial SpA (Table 384-1).

Comparable to the classification criteria for axial symptoms, the ASAS has recently formulated criteria for peripheral SpA. This is intended to exclude patients with axial symptoms and thus to divide the universe of patients with SpA into axial and exclusively peripheral subsets. These criteria are shown in Table 384-3.

Approximately one-half of the patients with undifferentiated SpA are HLA-B27-positive, and thus the absence of B27 is not useful in establishing or excluding the diagnosis. In familial cases, which are much more frequently B27-positive, there is often eventual progression to classical AS.

In juvenile-onset SpA, which begins between ages 7 and 16, most commonly in boys (60–80%), an asymmetric, predominantly lower-extremity oligoarthritis and enthesitis without extraarticular features is the typical mode of presentation. The prevalence of B27 in this condition, which has been termed the *seronegative enthesopathy and arthropathy (SEA) syndrome*, is approximately 80%. Many, but not all, of these patients go on to develop AS in late adolescence or adulthood.

Management of undifferentiated SpA is similar to that of the other spondyloarthritides. Response to anti-TNF- α therapy has been documented, and this therapy is indicated in severe, persistent cases not responsive to other treatment.

Current pediatric textbooks and journals should be consulted for information on management of juvenile-onset SpA.

ENTEROPATHIC ARTHRITIS

HISTORIC BACKGROUND

A relationship between arthritis and IBD was observed in the 1930s. The relationship was further defined by the epidemiologic studies

in the 1950s and 1960s and included in the concept of the spondyloarthritides in the 1970s.

EPIDEMIOLOGY

Both of the common forms of IBD, ulcerative colitis (UC) and Crohn's disease (CD) (Chap. 351), are associated with SpA. UC and CD both have an estimated prevalence of 0.05–0.1%, and the incidence of each is thought to have increased in recent decades. AS and peripheral arthritis are both associated with UC and CD. Wide variations have been reported in the estimated frequencies of these associations. In recent series, AS was diagnosed in 1–10%, and peripheral arthritis in 10–50% of patients with IBD. Inflammatory back pain and enthesopathy are common, and many patients have sacroiliitis on imaging studies.

The prevalence of UC or CD in patients with AS is thought to be 5–10%. However, investigation of unselected SpA patients by ileocolonoscopy has revealed that from one-third to two-thirds of patients with AS have subclinical intestinal inflammation that is evident either macroscopically or histologically. These lesions have also been found in patients with undifferentiated SpA or ReA (both enterically and urogenitally acquired).

Both UC and CD have a tendency to familial aggregation, more so for CD. HLA associations have been weak and inconsistent. HLA-B27 is found in up to 70% of patients with IBD and AS, but in $\leq 15\%$ of patients with IBD and peripheral arthritis or IBD alone. Three alleles of the *NOD2/CARD15* gene on chromosome 16 have been found in approximately one-half of patients with CD. These alleles are not associated with the spondyloarthritides per se. However, they are found significantly more often in (1) CD patients with sacroiliitis than in those without sacroiliitis, and (2) SpA patients with chronic inflammatory gut lesions than in those with normal gut histology. These associations are independent of HLA-B27. In addition to *NOD2*, over 100 other genes have been found to be associated with CD, UC, or both. Around 20 of these are also associated with AS.

PATHOLOGY

Available data for IBD-associated peripheral arthritis suggest a synovial histology similar to other spondyloarthritides. Association with arthropathy does not affect the gut histology of UC or CD (Chap. 351). The subclinical inflammatory lesions in the colon and distal ileum associated with SpA have been classified as either acute or chronic. The former resemble acute bacterial enteritis, with largely intact architecture and neutrophilic infiltration in the lamina propria. The latter resemble the lesions of CD, with distortion of villi and crypts, aphthoid ulceration, and mononuclear cell infiltration in the lamina propria.

PATHOGENESIS

Both IBD and SpA are immune-mediated, but the specific pathogenic mechanisms are poorly understood, and the connection between the two is obscure. The shared genetics evidently reflects shared pathogenic mechanisms. A number of rodent models showing various immune perturbations manifest both IBD and arthritis. Several lines of evidence indicate trafficking of leukocytes between the gut and the joint. Mucosal leukocytes from IBD patients have been shown to bind avidly to synovial vasculature through several different adhesion molecules. Macrophages expressing CD163 are prominent in the inflammatory lesions of both gut and synovium in the spondyloarthritides.

CLINICAL FEATURES

AS associated with IBD is clinically indistinguishable from idiopathic AS. It runs a course independent of the bowel disease, and in some patients, it precedes the onset of IBD, sometimes by many years. Peripheral arthritis may also begin before onset of overt bowel disease. The spectrum of peripheral arthritis includes acute self-limited attacks of oligoarthritis that often coincide with relapses of IBD, and more chronic and symmetric polyarticular arthritis that runs a course independent of IBD activity. The patterns of joint involvement are similar in UC and CD. In general, erosions and deformities are infrequent in IBD-associated peripheral arthritis, and joint surgery is infrequently required. Isolated destructive hip arthritis is a rare complication of CD,

TABLE 384-3 ASAS CRITERIA FOR PERIPHERAL SPONDYLOARTHRITIS^a

Arthritis ^b	or plus	Enthesitis
One or more of the following:		
• Uveitis		
• Psoriasis		
• Crohn's disease or ulcerative colitis		
OR two or more of the following:		
• Arthritis		
• Enthesitis		
• Dactylitis		
• Inflammatory back pain ever		
• Family history for SpA		

^aSensitivity 79.5%, specificity 83.3% ^bPeripheral arthritis, usually predominantly lower limb and/or asymmetric.

apparently distinct from osteonecrosis and septic arthritis. Dactylitis and enthesopathy are occasionally found. In addition to the ~20% of IBD patients with SpA, a comparable percentage have arthralgias or fibromyalgia symptoms.

Other extraintestinal manifestations of IBD are seen in addition to arthropathy, including uveitis, pyoderma gangrenosum, erythema nodosum, and finger clubbing, all somewhat more commonly in CD than UC. The uveitis shares the features described above for PsA-associated uveitis.

LABORATORY AND RADIOGRAPHIC FINDINGS

Laboratory findings reflect the inflammatory and metabolic manifestations of IBD. Joint fluid is usually at least mildly inflammatory. Of patients with AS and IBD, 30–70% carry the HLA-B27 gene, compared with >85% of patients with AS alone and 50–70% of those with AS and psoriasis. Hence, definite or probable AS in a B27-negative individual in the absence of psoriasis should prompt a search for occult IBD. Radiographic changes in the axial skeleton are the same as in uncomplicated AS. Erosions are uncommon in peripheral arthritis but may occur, particularly in the metatarsophalangeal joints. Isolated destructive hip disease has been described.

DIAGNOSIS

Diarrhea and arthritis are both common conditions that can coexist for a variety of reasons. When etiopathogenically related, ReA and IBD-associated arthritis are the most common causes. Rare causes include celiac disease, blind loop syndromes, and Whipple's disease. In most cases, diagnosis depends on investigation of the bowel disease.

TREATMENT

ENTEROPATHIC ARTHRITIS

Treatment of CD has been improved by therapy with anti-TNF agents. Infliximab, adalimumab, and certolizumab pegol are effective for induction and maintenance of clinical remission in CD, and infliximab has been shown to be effective in fistulizing CD. IBD-associated arthritis also responds to these agents. Other treatment for IBD, including sulfasalazine and related drugs, systemic glucocorticoids, and immunosuppressive drugs, is also usually of benefit for associated peripheral arthritis. NSAIDs are generally helpful and well tolerated, but they can precipitate flares of IBD. As noted above for psoriasis, rare cases of IBD, either CD or UC, have apparently been precipitated by anti-TNF therapy, usually etanercept, given for any of several rheumatic diseases.

SAPHO SYNDROME

The syndrome of synovitis, acne, pustulosis, hyperostosis, and osteitis (SAPHO) is characterized by a variety of skin and musculoskeletal manifestations. Dermatologic manifestations include palmo-plantar pustulosis, acne conglobata, acne fulminans, and hidradenitis suppurativa. The main musculoskeletal findings are sternoclavicular and spinal hyperostosis, chronic recurrent foci of sterile osteomyelitis, and axial or peripheral arthritis. Cases with one or a few manifestations are probably the rule. The ESR is usually elevated, sometimes dramatically. In some cases, bacteria, most often *Propionibacterium acnes*, have been cultured from bone biopsy specimens and occasionally other sites. IBD was coexistent in 8% of patients in one large series. B27 is not associated. Either bone scan or computed tomography scan is helpful diagnostically. An MRI report described characteristic vertebral body corner cortical erosions in 12 of 12 patients. High-dose NSAIDs may provide relief from bone pain. A number of uncontrolled series and case reports describe successful therapy with pamidronate or other bisphosphonates. Response to anti-TNF- α therapy has also been observed, although in a few cases this has been associated with a flare of skin manifestations. Successful prolonged antibiotic therapy has also been reported. Recent reports suggest a possible autoinflammatory pathogenesis and successful treatment with the IL-1 receptor antagonist anakinra.

385 The Vasculitis Syndromes

Carol A. Langford, Anthony S. Fauci

DEFINITION

Vasculitis is a clinicopathologic process characterized by inflammation of and damage to blood vessels. The vessel lumen is usually compromised, and this is associated with ischemia of the tissues supplied by the involved vessel. A broad and heterogeneous group of syndromes may result from this process, since any type, size, and location of blood vessel may be involved. Vasculitis and its consequences may be the primary or sole manifestation of a disease; alternatively, vasculitis may be a secondary component of another disease. Vasculitis may be confined to a single organ, such as the skin, or it may simultaneously involve several organ systems.

CLASSIFICATION

A major feature of the vasculitic syndromes as a group is the fact that there is a great deal of heterogeneity at the same time as there is considerable overlap among them. This heterogeneity and overlap in addition to a lack of understanding of the pathogenesis of these syndromes have been major impediments to the development of a coherent classification system for these diseases. [Table 385-1](#) lists the major vasculitis syndromes. The distinguishing and overlapping features of these syndromes are discussed below.

PATHOPHYSIOLOGY AND PATHOGENESIS

Generally, most of the vasculitic syndromes are assumed to be mediated at least in part by immunopathogenic mechanisms that occur in response to certain antigenic stimuli. However, evidence supporting this hypothesis is for the most part indirect and may reflect epiphenomena as opposed to true causality. Furthermore, it is unknown why some individuals might develop vasculitis in response to certain antigenic stimuli, whereas others do not. It is likely that a number of factors are involved in the ultimate expression of a vasculitic syndrome. These include the genetic predisposition, environmental exposures, and the regulatory mechanisms associated with immune response to certain antigens. Although immune complex formation, antineutrophil cytoplasmic antibodies (ANCA), and pathogenic T lymphocyte responses ([Table 385-2](#)) have been among the prominent hypothesized

TABLE 385-1 VASCULITIS SYNDROMES

Primary Vasculitis Syndromes	Secondary Vasculitis Syndromes
Granulomatosis with polyangiitis (Wegener's)	Vasculitis associated with probable etiology
Microscopic polyangiitis	Drug-induced vasculitis
Eosinophilic granulomatosis with polyangiitis (Churg-Strauss)	Hepatitis C virus-associated cryoglobulinemic vasculitis
IgA vasculitis (Henoch-Schönlein)	Hepatitis B virus-associated vasculitis
Cryoglobulinemic vasculitis	Cancer-associated vasculitis
Polyarteritis nodosa	Vasculitis associated with systemic disease
Kawasaki disease	Lupus vasculitis
Giant cell arteritis	Rheumatoid vasculitis
Takayasu arteritis	Sarcoid vasculitis
Behçet's disease	
Cogan's syndrome	
Single-organ vasculitis	
Cutaneous leukocytoclastic angiitis	
Cutaneous arteritis	
Primary central nervous system vasculitis	
Isolated aortitis	

Source: Adapted from JC Jennette et al: *Arthritis Rheum* 65:1, 2013.

TABLE 385-2 POTENTIAL MECHANISMS OF VESSEL DAMAGE IN VASCULITIS SYNDROMES

Pathogenic immune-complex formation and/or deposition
IgA vasculitis (Henoch-Schönlein)
Lupus vasculitis
Serum sickness and cutaneous vasculitis syndromes
Hepatitis C virus–associated cryoglobulinemic vasculitis
Hepatitis B virus–associated vasculitis
Production of antineutrophilic cytoplasmic antibodies
Granulomatosis with polyangiitis (Wegener’s)
Microscopic polyangiitis
Eosinophilic granulomatosis with polyangiitis (Churg-Strauss)
Pathogenic T lymphocyte responses and granuloma formation
Giant cell arteritis
Takayasu arteritis
Granulomatosis with polyangiitis (Wegener’s)
Eosinophilic granulomatosis with polyangiitis (Churg-Strauss)

Source: Adapted from MC Sneller, AS Fauci: Med Clin North Am 81:221, 1997.

mechanisms, it is likely that the pathogenesis of individual forms of vasculitis is complex and varied.

PATHOGENIC IMMUNE-COMPLEX FORMATION

Deposition of immune complexes was the first and most widely accepted pathogenic mechanism of vasculitis. However, the causal role of immune complexes has not been clearly established in most of the vasculitic syndromes. Circulating immune complexes need not result in deposition of the complexes in blood vessels with ensuing vasculitis, and many patients with active vasculitis do not have demonstrable circulating or deposited immune complexes. The actual antigen contained in the immune complex has only rarely been identified in vasculitic syndromes. In this regard, hepatitis B antigen has been identified in both the circulating and deposited immune complexes in a subset of patients who have features of a systemic vasculitis, most notably in polyarteritis nodosa (see “Polyarteritis Nodosa”). Cryoglobulinemic vasculitis is strongly associated with hepatitis C virus infection; hepatitis C virions and hepatitis C virus antigen-antibody complexes have been identified in the cryoprecipitates of these patients (see “Cryoglobulinemic Vasculitis”).

The mechanisms of tissue damage in immune complex-mediated vasculitis resemble those described for serum sickness. In this model, antigen-antibody complexes are formed in antigen excess and are deposited in vessel walls whose permeability has been increased by vasoactive amines such as histamine, bradykinin, and leukotrienes released from platelets or from mast cells as a result of IgE-triggered mechanisms. The deposition of complexes results in activation of complement components, particularly C5a, which is strongly chemotactic for neutrophils. These cells then infiltrate the vessel wall, phagocytose the immune complexes, and release their intracytoplasmic enzymes, which damage the vessel wall. As the process becomes subacute or chronic, mononuclear cells infiltrate the vessel wall. The common denominator of the resulting syndrome is compromise of the vessel lumen with ischemic changes in the tissues supplied by the involved vessel. Several variables may explain why only certain types of immune complexes cause vasculitis and why only certain vessels are affected in individual patients. These include the ability of the reticuloendothelial system to clear circulating complexes from the blood, the size and physicochemical properties of immune complexes, the relative degree of turbulence of blood flow, the intravascular hydrostatic pressure in different vessels, and the preexisting integrity of the vessel endothelium.

ANTINEUTROPHIL CYTOPLASMIC ANTIBODIES (ANCA)

ANCA are antibodies directed against certain proteins in the cytoplasmic granules of neutrophils and monocytes. These autoantibodies are present in a high percentage of patients with active granulomatosis with polyangiitis (Wegener’s) and microscopic polyangiitis, and in a

lower percentage of patients with eosinophilic granulomatosis with polyangiitis (Churg-Strauss). Because these diseases share the presence of ANCA and small-vessel vasculitis, some investigators have come to refer to them collectively as “ANCA-associated vasculitis.” However, as these diseases possess unique clinical phenotypes in which ANCA may be absent, it remains our opinion that granulomatosis with polyangiitis (Wegener’s), microscopic polyangiitis, and eosinophilic granulomatosis with polyangiitis (Churg-Strauss) should continue to be viewed as separate entities.

There are two major categories of ANCA based on different targets for the antibodies. The terminology of *cytoplasmic ANCA* (cANCA) refers to the diffuse, granular cytoplasmic staining pattern observed by immunofluorescence microscopy when serum antibodies bind to indicator neutrophils. Proteinase-3, a 29-kDa neutral serine proteinase present in neutrophil azurophilic granules, is the major cANCA antigen. More than 90% of patients with typical active granulomatosis with polyangiitis (Wegener’s) have detectable antibodies to proteinase-3 (see below). The terminology of *perinuclear ANCA* (pANCA) refers to the more localized perinuclear or nuclear staining pattern of the indicator neutrophils. The major target for pANCA is the enzyme myeloperoxidase; other targets that can produce a pANCA pattern of staining include elastase, cathepsin G, lactoferrin, lysozyme, and bactericidal/permeability-increasing protein. However, only antibodies to myeloperoxidase have been convincingly associated with vasculitis. Antimyeloperoxidase antibodies have been reported to occur in variable percentages of patients with microscopic polyangiitis, eosinophilic granulomatosis with polyangiitis (Churg-Strauss), isolated necrotizing crescentic glomerulonephritis, and granulomatosis with polyangiitis (Wegener’s) (see below). A pANCA pattern of staining that is not due to antimyeloperoxidase antibodies has been associated with nonvasculitic entities such as rheumatic and nonrheumatic autoimmune diseases, inflammatory bowel disease, certain drugs, and infections such as endocarditis and bacterial airway infections in patients with cystic fibrosis.

It is unclear why patients with these vasculitis syndromes develop antibodies to myeloperoxidase or proteinase-3 or what role these antibodies play in disease pathogenesis. There are a number of in vitro observations that suggest possible mechanisms whereby these antibodies can contribute to the pathogenesis of the vasculitis syndromes. Proteinase-3 and myeloperoxidase reside in the azurophilic granules and lysosomes of resting neutrophils and monocytes, where they are apparently inaccessible to serum antibodies. However, when neutrophils or monocytes are primed by tumor necrosis factor α (TNF- α) or interleukin 1 (IL-1), proteinase-3 and myeloperoxidase translocate to the cell membrane, where they can interact with extracellular ANCA. The neutrophils then degranulate and produce reactive oxygen species that can cause tissue damage. Furthermore, ANCA-activated neutrophils can adhere to and kill endothelial cells in vitro. Activation of neutrophils and monocytes by ANCA also induces the release of proinflammatory cytokines such as IL-1 and IL-8. Adoptive transfer experiments in genetically engineered mice provide further evidence for a direct pathogenic role of ANCA in vivo. In contradiction, however, a number of clinical and laboratory observations argue against a primary pathogenic role for ANCA. Patients may have active granulomatosis with polyangiitis (Wegener’s) in the absence of ANCA; the absolute height of the antibody titers does not correlate well with disease activity; and patients with granulomatosis with polyangiitis (Wegener’s) in remission may continue to have high antiproteinase-3 (cANCA) titers for years (see below).

PATHOGENIC T LYMPHOCYTE RESPONSES AND GRANULOMA FORMATION

The histopathologic feature of granulomatous vasculitis has provided evidence to support a role of pathogenic T lymphocyte responses and cell-mediated immune injury. Vascular endothelial cells can express HLA class II molecules following activation by cytokines such as interferon (IFN) γ . This allows these cells to participate in immunologic reactions such as interaction with CD4⁺ T lymphocytes in a manner similar to antigen-presenting macrophages. Endothelial cells can secrete IL-1, which may activate T lymphocytes and initiate or

propagate in situ immunologic processes within the blood vessel. In addition, IL-1 and TNF- α are potent inducers of endothelial-leukocyte adhesion molecule 1 (ELAM-1) and vascular cell adhesion molecule 1 (VCAM-1), which may enhance the adhesion of leukocytes to endothelial cells in the blood vessel wall.

APPROACH TO THE PATIENT: General Principles of Diagnosis

The diagnosis of vasculitis is often considered in any patient with an unexplained systemic illness. However, there are certain clinical abnormalities that when present alone or in combination should suggest a diagnosis of vasculitis. These include palpable purpura, pulmonary infiltrates and microscopic hematuria, chronic inflammatory sinusitis, mononeuritis multiplex, unexplained ischemic events, and glomerulonephritis with evidence of multisystem disease. A number of nonvasculitic diseases may also produce some or all of these abnormalities. Thus, the first step in the workup of a patient with suspected vasculitis is to exclude other diseases that produce clinical manifestations that can mimic vasculitis (Table 385-3). It is particularly important to exclude infectious diseases with features that overlap those of vasculitis, especially if the patient's clinical condition is deteriorating rapidly and empirical immunosuppressive treatment is being contemplated.

Once diseases that mimic vasculitis have been excluded, the workup should follow a series of progressive steps that establish the diagnosis of vasculitis and determine, where possible, the category of the vasculitis syndrome (Fig. 385-1). This approach is of considerable importance since several of the vasculitis syndromes require aggressive therapy with glucocorticoids and other immunosuppressive agents, whereas other syndromes usually resolve spontaneously and require symptomatic treatment only. The definitive

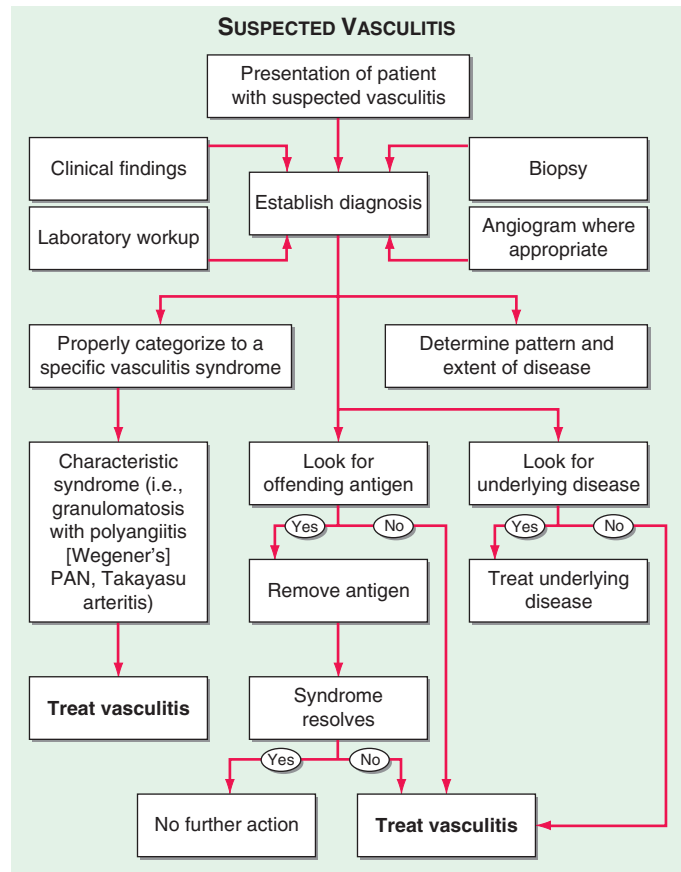


FIGURE 385-1 Algorithm for the approach to a patient with suspected diagnosis of vasculitis. PAN, polyarteritis nodosa.

TABLE 385-3 CONDITIONS THAT CAN MIMIC VASCULITIS

Infectious diseases
Bacterial endocarditis
Disseminated gonococcal infection
Pulmonary histoplasmosis
Coccidioidomycosis
Syphilis
Lyme disease
Rocky Mountain spotted fever
Whipple's disease
Coagulopathies/thrombotic microangiopathies
Antiphospholipid syndrome
Thrombotic thrombocytopenic purpura
Neoplasms
Atrial myxoma
Lymphoma
Carcinomatosis
Drug toxicity
Cocaine
Levamisole
Amphetamines
Ergot alkaloids
Methysergide
Arsenic
Sarcoidosis
Atheroembolic disease
Antiglomerular basement membrane disease (Goodpasture's syndrome)
Amyloidosis
Migraine
Reversible cerebral vasoconstrictive syndrome

diagnosis of vasculitis is usually made based on biopsy of involved tissue. The yield of "blind" biopsies of organs with no subjective or objective evidence of involvement is very low and should be avoided. When syndromes such as polyarteritis nodosa, Takayasu arteritis, or primary central nervous system (CNS) vasculitis are suspected, arteriogram of organs with suspected involvement should be performed.

GENERAL PRINCIPLES OF TREATMENT Once a diagnosis of vasculitis has been established, a decision regarding therapeutic strategy must be made (Fig. 385-1). If an offending antigen that precipitates the vasculitis is recognized, the antigen should be removed where possible. If the vasculitis is associated with an underlying disease such as an infection, neoplasm, or connective tissue disease, the underlying disease should be treated. If the syndrome represents a primary vasculitic disease, treatment should be initiated according to the category of the vasculitis syndrome. Specific therapeutic regimens are discussed below for the individual vasculitis syndromes; however, certain general principles regarding therapy should be considered. Decisions regarding treatment should be based on the use of regimens for which there has been published literature supporting efficacy for that particular vasculitic disease. Since the potential toxic side effects of certain therapeutic regimens may be substantial, the risk-versus-benefit ratio of any therapeutic approach should be weighed carefully. On the one hand, glucocorticoids and/or other immunosuppressive agents should be instituted immediately in diseases where irreversible organ system dysfunction and high morbidity and mortality rates have been clearly established. Granulomatosis with polyangiitis (Wegener's) is the prototype of a severe systemic vasculitis requiring such a therapeutic approach (see below). On the other hand, when feasible, aggressive therapy should be avoided for vasculitic manifestations that rarely result in irreversible organ system dysfunction and that usually do not respond to such therapy. For example, isolated

idiopathic cutaneous vasculitis usually resolves with symptomatic treatment, and prolonged courses of glucocorticoids uncommonly result in clinical benefit. Cytotoxic agents have not proved to be beneficial in idiopathic cutaneous vasculitis, and their toxic side effects generally outweigh any potential beneficial effects. Glucocorticoids should be initiated in those systemic vasculitides that cannot be specifically categorized or for which there is no established standard therapy; or other immunosuppressive therapy should be added in these diseases only if an adequate response does not result or if remission can only be achieved and maintained with an unacceptably toxic regimen of glucocorticoids. When remission is achieved, one should continually attempt to taper glucocorticoids and discontinue when possible. When using other immunosuppressive regimens, one should base the choice of agent upon the available therapeutic data supporting efficacy in that disease, the site and severity of organ involvement, and the toxicity profile of the drug.

Physicians should be thoroughly aware of the toxic side effects of therapeutic agents employed that can include both acute and long-term complications (Table 385-4). Morbidity and mortality can occur as a result of treatment, and strategies to monitor for and prevent toxicity represent an essential part of patient care. Glucocorticoids are an important part of treatment for most vasculitides but are associated with substantial toxicities. Monitoring and prevention of glucocorticoid-induced bone loss are important in all patients. With the use of daily cyclophosphamide, strategies are particularly important and are directed toward minimization

TABLE 385-4 MAJOR TOXIC SIDE EFFECTS OF DRUGS USED IN THE TREATMENT OF SYSTEMIC VASCULITIS

Glucocorticoids	
Osteoporosis	Growth suppression in children
Cataracts	Hypertension
Glaucoma	Avascular necrosis of bone
Diabetes mellitus	Myopathy
Electrolyte abnormalities	Alterations in mood
Metabolic abnormalities	Psychosis
Suppression of inflammatory and immune responses leading to opportunistic infections	Pseudotumor cerebri Peptic ulcer diathesis Pancreatitis
Cushingoid features	
Cyclophosphamide	
Bone marrow suppression	Hypogammaglobulinemia
Cystitis	Pulmonary fibrosis
Bladder carcinoma	Myelodysplasia
Gonadal suppression	Oncogenesis
Gastrointestinal intolerance	Teratogenicity Opportunistic infections
Methotrexate	
Gastrointestinal intolerance	Pneumonitis
Stomatitis	Teratogenicity
Bone marrow suppression	Opportunistic infections
Hepatotoxicity (may lead to fibrosis or cirrhosis)	
Azathioprine	
Gastrointestinal intolerance	Opportunistic infections
Bone marrow suppression	Hypersensitivity
Hepatotoxicity	
Rituximab	
Infusion reactions	Opportunistic infections
Progressive multifocal leukoencephalopathy	Hepatitis B reactivation
Mucocutaneous reactions	Tumor lysis syndrome

of bladder toxicity and prevention of leukopenia. Instructing the patient to take cyclophosphamide all at once in the morning with a large amount of fluid throughout the day in order to maintain a dilute urine can reduce the risk of bladder injury. Bladder cancer can occur several years after discontinuation of cyclophosphamide therapy; therefore, monitoring for bladder cancer should continue indefinitely in patients who have received cyclophosphamide. Bone marrow suppression is an important toxicity of cyclophosphamide and can be observed during glucocorticoid tapering or over time, even after periods of stable measurements. Monitoring of the complete blood count every 1–2 weeks for as long as the patient receives cyclophosphamide can effectively prevent cytopenias. Maintaining the white blood cell (WBC) count at $>3000/\mu\text{L}$ and the neutrophil count at $>1500/\mu\text{L}$ is essential to reduce the risk of life-threatening infections.

Methotrexate and azathioprine are also associated with bone marrow suppression, and complete blood counts should be obtained every 1–2 weeks for the first 1–2 months after their initiation and once a month thereafter. To lessen toxicity, methotrexate is often given together with folic acid, 1 mg daily, or folinic acid, 5–10 mg once a week 24 h following methotrexate. Prior to initiation of azathioprine, thiopurine methyltransferase (TPMT), an enzyme involved in the metabolism of azathioprine, should be assayed because inadequate levels may result in severe cytopenia.

Infection represents a significant toxicity for all vasculitis patients treated with immunosuppressive therapy. Infections with *Pneumocystis jiroveci* and certain fungi can be seen even in the face of WBCs that are within normal limits, particularly in patients receiving glucocorticoids. All vasculitis patients who are receiving daily glucocorticoids in combination with another immunosuppressive agent should receive trimethoprim-sulfamethoxazole (TMP-SMX) or another prophylactic therapy to prevent *P. jiroveci* infection.

Finally, it should be emphasized that each patient is unique and requires individual decision-making. The above outline should serve as a framework to guide therapeutic approaches; however, flexibility should be practiced in order to provide maximal therapeutic efficacy with minimal toxic side effects in each patient.

GRANULOMATOSIS WITH POLYANGIITIS (WEGENER'S)

DEFINITION

Granulomatosis with polyangiitis (Wegener's) is a distinct clinicopathologic entity characterized by granulomatous vasculitis of the upper and lower respiratory tracts together with glomerulonephritis. In addition, variable degrees of disseminated vasculitis involving both small arteries and veins may occur.

INCIDENCE AND PREVALENCE

Granulomatosis with polyangiitis (Wegener's) is an uncommon disease with an estimated prevalence of 3 per 100,000. It is extremely rare in blacks compared with whites; the male-to-female ratio is 1:1. The disease can be seen at any age; ~15% of patients are <19 years of age, but only rarely does the disease occur before adolescence; the mean age of onset is ~40 years.

PATHOLOGY AND PATHOGENESIS

The histopathologic hallmarks of granulomatosis with polyangiitis (Wegener's) are necrotizing vasculitis of small arteries and veins together with granuloma formation, which may be either intravascular or extravascular (Fig. 385-2). Lung involvement typically appears as multiple, bilateral, nodular cavitary infiltrates (Fig. 385-3), which on biopsy almost invariably reveal the typical necrotizing granulomatous vasculitis. Upper airway lesions, particularly those in the sinuses and nasopharynx, typically reveal inflammation, necrosis, and granuloma formation, with or without vasculitis.

In its earliest form, renal involvement is characterized by a focal and segmental glomerulitis that may evolve into a rapidly progressive

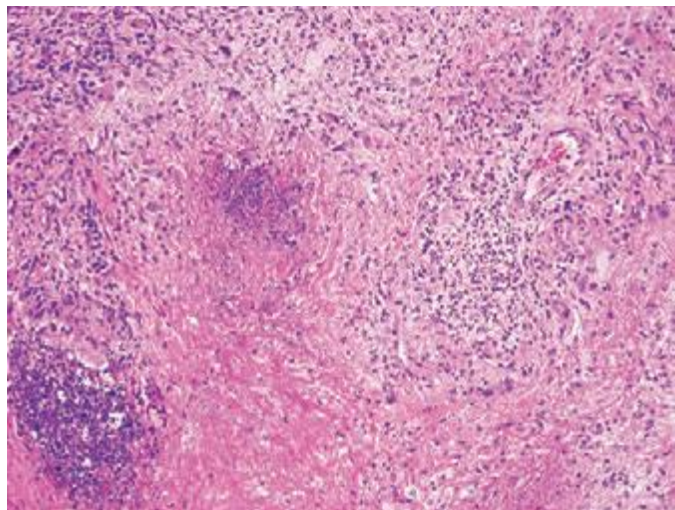


FIGURE 385-2 Lung histology in granulomatosis with polyangiitis (Wegener's). This area of geographic necrosis has a serpiginous border of histiocytes and giant cells surrounding a central necrotic zone. Vasculitis is also present with neutrophils and lymphocytes infiltrating the wall of a small arteriole (upper right). (Courtesy of William D. Travis, MD; with permission.)

crescentic glomerulonephritis. Granuloma formation is only rarely seen on renal biopsy. In contrast to other forms of glomerulonephritis, evidence of immune complex deposition is not found in the renal lesion of granulomatosis with polyangiitis (Wegener's). In addition to the classic triad of disease of the upper and lower respiratory tracts and kidney, virtually any organ can be involved with vasculitis, granuloma, or both.

The immunopathogenesis of this disease is unclear, although the involvement of upper airways and lungs with granulomatous vasculitis suggests an aberrant cell-mediated immune response to an exogenous or even endogenous antigen that enters through or resides in the upper airway. Chronic nasal carriage of *Staphylococcus aureus* has been reported to be associated with a higher relapse rate of granulomatosis with polyangiitis (Wegener's); however, there is no evidence for a role of this organism in the pathogenesis of the disease.

Peripheral blood mononuclear cells obtained from patients with granulomatosis with polyangiitis (Wegener's) manifest increased secretion of IFN- γ but not of IL-4, IL-5, or IL-10 compared to normal controls. In addition, TNF- α production from peripheral blood mononuclear cells and CD4+ T cells is elevated. Furthermore, monocytes from patients with granulomatosis with polyangiitis (Wegener's)



FIGURE 385-3 Computed tomography scan of a patient with granulomatosis with polyangiitis (Wegener's). The patient developed multiple, bilateral, and cavitary infiltrates.

produce increased amounts of IL-12. These findings indicate an unbalanced T_H1-type T cell cytokine pattern in this disease that may have pathogenic and perhaps ultimately therapeutic implications.

A high percentage of patients with granulomatosis with polyangiitis (Wegener's) develop ANCA, and these autoantibodies may play a role in the pathogenesis of this disease (see above).

CLINICAL AND LABORATORY MANIFESTATIONS

Involvement of the upper airways occurs in 95% of patients with granulomatosis with polyangiitis (Wegener's). Patients often present with severe upper respiratory tract findings such as paranasal sinus pain and drainage and purulent or bloody nasal discharge, with or without nasal mucosal ulceration (Table 385-5). Nasal septal perforation may follow, leading to saddle nose deformity. Serous otitis media may occur as a result of eustachian tube blockage. Subglottic tracheal stenosis resulting from active disease or scarring occurs in ~16% of patients and may result in severe airway obstruction.

TABLE 385-5 GRANULOMATOSIS WITH POLYANGIITIS (WEGENER'S): FREQUENCY OF CLINICAL MANIFESTATIONS IN 158 PATIENTS STUDIED AT THE NATIONAL INSTITUTES OF HEALTH

Manifestation	Percentage at Disease Onset	Percentage Throughout Course of Disease
Kidney		
Glomerulonephritis	18	77
Ear/Nose/Throat		
Sinusitis	51	85
Nasal disease	36	68
Otitis media	25	44
Hearing loss	14	42
Subglottic stenosis	1	16
Ear pain	9	14
Oral lesions	3	10
Lung		
Pulmonary infiltrates	25	66
Pulmonary nodules	24	58
Hemoptysis	12	30
Pleuritis	10	28
Eyes		
Conjunctivitis	5	18
Dacryocystitis	1	18
Scleritis	6	16
Proptosis	2	15
Eye pain	3	11
Visual loss	0	8
Retinal lesions	0	4
Corneal lesions	0	1
Iritis	0	2
Other^a		
Arthralgias/arthritis	32	67
Fever	23	50
Cough	19	46
Skin abnormalities	13	46
Weight loss (>10% body weight)	15	35
Peripheral neuropathy	1	15
Central nervous system disease	1	8
Pericarditis	2	6
Hyperthyroidism	1	3

^aFewer than 1% had parotid, pulmonary artery, breast, or lower genitourinary (urethra, cervix, vagina, testicular) involvement.

Source: GS Hoffman et al: Ann Intern Med 116:488, 1992.

Pulmonary involvement may be manifested as asymptomatic infiltrates or may be clinically expressed as cough, hemoptysis, dyspnea, and chest discomfort. It is present in 85–90% of patients. Endobronchial disease, either in its active form or as a result of fibrous scarring, may lead to obstruction with atelectasis.

Eye involvement (52% of patients) may range from a mild conjunctivitis to dacryocystitis, episcleritis, scleritis, granulomatous sclerouveitis, ciliary vessel vasculitis, and retroorbital mass lesions leading to proptosis.

Skin lesions (46% of patients) appear as papules, vesicles, palpable purpura, ulcers, or subcutaneous nodules; biopsy reveals vasculitis, granuloma, or both. Cardiac involvement (8% of patients) manifests as pericarditis, coronary vasculitis, or, rarely, cardiomyopathy. Nervous system manifestations (23% of patients) include cranial neuritis, mononeuritis multiplex, or, rarely, cerebral vasculitis and/or granuloma.

Renal disease (77% of patients) generally dominates the clinical picture and, if left untreated, accounts directly or indirectly for most of the mortality rate in this disease. Although it may smolder in some cases as a mild glomerulitis with proteinuria, hematuria, and red blood cell casts, it is clear that once clinically detectable renal functional impairment occurs, rapidly progressive renal failure usually ensues unless appropriate treatment is instituted.

While the disease is active, most patients have nonspecific symptoms and signs such as malaise, weakness, arthralgias, anorexia, and weight loss. Fever may indicate activity of the underlying disease but more often reflects secondary infection, usually of the upper airway.

Characteristic laboratory findings include a markedly elevated erythrocyte sedimentation rate (ESR), mild anemia and leukocytosis, mild hypergammaglobulinemia (particularly of the IgA class), and mildly elevated rheumatoid factor. Thrombocytosis may be seen as an acute-phase reactant. Approximately 90% of patients with active granulomatosis with polyangiitis (Wegener's) have a positive antiproteinase-3 ANCA. However, in the absence of active disease, the sensitivity drops to ~60–70%. A small percentage of patients with granulomatosis with polyangiitis (Wegener's) may have antityeloperoxidase rather than antiproteinase-3 antibodies, and up to 20% may lack ANCA.

Patients with granulomatosis with polyangiitis (Wegener's) have been found to have an increased incidence of venous thrombotic events. Although routine anticoagulation for all patients is not recommended, a heightened awareness for any clinical features suggestive of deep venous thrombosis or pulmonary emboli is warranted.

DIAGNOSIS

The diagnosis of granulomatosis with polyangiitis (Wegener's) is made by the demonstration of necrotizing granulomatous vasculitis on tissue biopsy in a patient with compatible clinical features. Pulmonary tissue offers the highest diagnostic yield, almost invariably revealing granulomatous vasculitis. Biopsy of upper airway tissue usually reveals granulomatous inflammation with necrosis but may not show vasculitis. Renal biopsy can confirm the presence of pauci-immune glomerulonephritis.

The specificity of a positive antiproteinase-3 ANCA for granulomatosis with polyangiitis (Wegener's) is very high, especially if active glomerulonephritis is present. However, the presence of ANCA should be adjunctive and, with rare exceptions, should not substitute for a tissue diagnosis. False-positive ANCA titers have been reported in certain infectious and neoplastic diseases.

In its typical presentation, the clinicopathologic complex of granulomatosis with polyangiitis (Wegener's) usually provides ready differentiation from other disorders. However, if all the typical features are not present at once, it needs to be differentiated from the other vasculitides, antiglomerular basement membrane disease (Goodpasture's syndrome) (Chap. 338), relapsing polychondritis (Chap. 389), tumors of the upper airway or lung, and infectious diseases such as histoplasmosis (Chap. 236), mucocutaneous leishmaniasis (Chap. 251), and rhinoscleroma (Chap. 44) as well as noninfectious granulomatous diseases.

Of particular note is the differentiation from other *midline destructive diseases*. These diseases lead to extreme tissue destruction and mutilation localized to the midline upper airway structures including the sinuses; erosion through the skin of the face commonly occurs,

a feature that is extremely rare in granulomatosis with polyangiitis (Wegener's). Although blood vessels may be involved in the intense inflammatory reaction and necrosis, primary vasculitis is not seen. *Upper airway neoplasms* and specifically *extranodal natural killer (NK)/T cell lymphoma (nasal type)* are important causes of midline destructive disease. These lesions are diagnosed based on histology, which reveals polymorphous atypical lymphoid cells with an NK cell immunophenotype, typically Epstein-Barr virus (Chap. 134). Such cases are treated based on their degree of dissemination, and localized lesions have responded to irradiation. Upper airway lesions should never be irradiated in granulomatosis with polyangiitis (Wegener's). Cocaine-induced tissue injury can be another important mimic of granulomatosis with polyangiitis (Wegener's) in patients who present with isolated midline destructive disease. ANCA that target human neutrophil elastase can be found in patients with cocaine-induced midline destructive lesions and can complicate the differentiation from granulomatosis with polyangiitis (Wegener's). This has been further confounded by the high frequency of levamisole adulteration of cocaine, which can result in cutaneous infarction and serologic changes that may mimic vasculitis. Granulocytopenia is a common finding in levamisole-induced disease that would not be associated with granulomatosis with polyangiitis (Wegener's).

Granulomatosis with polyangiitis (Wegener's) must also be differentiated from *lymphomatoid granulomatosis*, which is an Epstein-Barr virus–positive B cell proliferation that is associated with an exuberant T cell reaction. Lymphomatoid granulomatosis is characterized by lung, skin, CNS, and kidney involvement in which atypical lymphocytoid and plasmacytoid cells infiltrate nonlymphoid tissue in an angioinvasive manner. In this regard, it clearly differs from granulomatosis with polyangiitis (Wegener's) in that it is not an inflammatory vasculitis in the classic sense but an angiocentric perivascular infiltration of atypical mononuclear cells. Up to 50% of patients may develop a true malignant lymphoma.

TREATMENT

GRANULOMATOSIS WITH POLYANGIITIS (WEGENER'S)

Prior to the introduction of effective therapy, granulomatosis with polyangiitis (Wegener's) was universally fatal within a few months of diagnosis. Glucocorticoids alone led to some symptomatic improvement, with little effect on the ultimate course of the disease. The development of treatment with cyclophosphamide dramatically changed patient outcome such that marked improvement was seen in >90% of patients, complete remission in 75% of patients, and 5-year patient survival was seen in over 80%.

Despite the ability to successfully induce remission, 50–70% of remissions are later associated with one or more relapses. The determination of relapse should be based on objective evidence of disease activity, taking care to rule out other features that may have a similar appearance such as infection, medication toxicity, or chronic disease sequelae. The ANCA titer can be misleading and should not be used to assess disease activity. Many patients who achieve remission continue to have elevated titers for years. Results from a large prospective study found that increases in ANCA were not associated with relapse and that only 43% relapsed within 1 year of an increase in ANCA levels. Thus, a rise in ANCA by itself is not a harbinger of immediate disease relapse and should not lead to reinstitution or increase in immunosuppressive therapy. Reinduction of remission after relapse is almost always achieved; however, a high percentage of patients ultimately have some degree of damage from irreversible features of their disease, such as varying degrees of renal insufficiency, hearing loss, tracheal stenosis, saddle nose deformity, and chronically impaired sinus function. Patients who developed irreversible renal failure but who achieved subsequent remission have undergone successful renal transplantation.

Because long-term cyclophosphamide is associated with substantial toxicity, approaches have been developed that seek to minimize the duration of exposure to cyclophosphamide while still

taking advantage of its efficacy for severe disease. Treatment of granulomatosis with polyangiitis (Wegener's) is currently viewed as having two phases: *induction*, where active disease is put into remission, followed by *maintenance*. The decision regarding which agents to use for induction and maintenance is based on disease severity together with individual patient factors that include contraindication, relapse history, and comorbidities.

CYCLOPHOSPHAMIDE INDUCTION FOR SEVERE DISEASE

For patients with severe disease, daily cyclophosphamide combined with glucocorticoids has been repeatedly proved to effectively induce remission and prolong survival. At the initiation of therapy, glucocorticoids are usually given as prednisone, 1 mg/kg per day for the first month, followed by gradual tapering on an alternate-day or daily schedule with discontinuation after ~6–9 months.

Cyclophosphamide is given in doses of 2 mg/kg per day orally, but as it is renally eliminated, dosage reduction should be considered in patients with renal insufficiency. Some reports have indicated therapeutic success with less frequent and severe toxic side effects using IV cyclophosphamide. In a recent randomized trial, IV cyclophosphamide 15 mg/kg, three infusions given every 2 weeks, then every 3 weeks thereafter, was compared to cyclophosphamide 2 mg/kg daily given for 3 months followed by 1.5 mg/kg daily. Although IV cyclophosphamide was found to have a comparable rate of remission with a lower cumulative cyclophosphamide dose and occurrence of leukopenia, the use of a consolidation phase and an insufficient frequency of blood count monitoring may have negatively influenced the results in those who received daily cyclophosphamide. Of note in this study was that relapse occurred in 19% of those who received IV cyclophosphamide as compared to 9% who received daily oral administration. We continue to strongly favor daily cyclophosphamide with utilization of blood count monitoring every 1–2 weeks (as discussed above) and limiting the duration of induction exposure to 3–6 months.

In patients with imminently life-threatening disease, such as rapidly progressive glomerulonephritis with a creatinine greater than 4.0 mg/dL or pulmonary hemorrhage requiring mechanical ventilation, a regimen of daily cyclophosphamide and glucocorticoids is the treatment of choice to induce remission. Adjunctive plasmapheresis was found to further improve renal recovery in a study of patients with rapidly progressive glomerulonephritis who had a creatinine of greater than 5.8 mg/dL.

REMISSION MAINTENANCE AFTER CYCLOPHOSPHAMIDE

After 3–6 months of induction treatment, cyclophosphamide should be stopped and switched to another agent for remission maintenance. The agents with which there has been the greatest published experience are methotrexate and azathioprine. Methotrexate is administered orally or subcutaneously starting at a dosage of 0.3 mg/kg as a single weekly dose, not to exceed 15 mg/week. If the treatment is well tolerated after 1–2 weeks, the dosage should be increased by 2.5 mg weekly up to a dosage of 20–25 mg/week and maintained at that level. Azathioprine, 2 mg/kg per day, has also proved effective in maintaining remission following induction with daily cyclophosphamide. In a randomized trial comparing methotrexate to azathioprine for remission maintenance, comparable rates of toxicity and relapse were seen. Therefore, the choice of agent is often based on toxicity profile, because methotrexate cannot be given to patients with renal insufficiency or chronic liver disease, as well as on other individual patient factors. In patients who are unable to receive methotrexate or azathioprine or who have relapsed through such treatment, mycophenolate mofetil, 1000 mg twice a day, may also sustain remission following cyclophosphamide induction.

The optimal duration of maintenance therapy is uncertain. In the absence of toxicity, maintenance therapy is usually given for a minimum of 2 years past remission, after which time consideration can be given for tapering over a 6–12 month period until discontinuation. Patients with significant organ damage or a history of relapse may benefit from longer-term continuation of a maintenance agent.

RITUXIMAB INDUCTION FOR SEVERE DISEASE

Rituximab is a chimeric monoclonal antibody directed against CD20 present on normal and malignant B lymphocytes that is U.S. Food and Drug Administration (FDA) approved for the treatment of granulomatosis with polyangiitis (Wegener's) and microscopic polyangiitis. In two randomized trials that enrolled ANCA-positive patients with severe active granulomatosis with polyangiitis (Wegener's) or microscopic polyangiitis, rituximab 375 mg/m² once a week for 4 weeks in combination with glucocorticoids was found to be as effective as cyclophosphamide with glucocorticoids for inducing disease remission. In the trial, which also enrolled patients with relapsing disease, rituximab was found to be statistically superior to cyclophosphamide.

Although the data support that rituximab is effective for remission induction of severe active granulomatosis with polyangiitis (Wegener's) or microscopic polyangiitis, there remain a number of ongoing questions regarding rituximab that must be considered in weighing its use in the individual patient. The optimal approach to remission maintenance after treatment with rituximab remains unclear, as does whether this should include conventional maintenance agents such as methotrexate or azathioprine versus scheduled retreatment with rituximab. In addition, there are no long-term safety data with rituximab in granulomatosis with polyangiitis (Wegener's) or microscopic polyangiitis.

Although rituximab does not have the bladder toxicity or infertility concerns, as can occur with cyclophosphamide, in both of the randomized trials, the rate of adverse events was similar in the rituximab and cyclophosphamide arms. Serious side effects of rituximab include infusion reactions, severe mucocutaneous reactions, and rare reports of progressive multifocal leukoencephalopathy. Because rituximab can bring about reactivation of hepatitis B, all patients should undergo hepatitis screening prior to treatment with rituximab.

OTHER BIOLOGIC THERAPIES

Etanercept, a dimeric fusion protein containing the 75-kDa TNF receptor bound to human IgG1, was not found to sustain remission when used adjunctively to standard therapy and should not be used in the treatment of granulomatosis with polyangiitis (Wegener's).

METHOTREXATE INDUCTION FOR NONSEVERE DISEASE

For selected patients whose disease is not immediately life threatening, methotrexate together with glucocorticoids given at the dosages described above may be considered as an alternative for induction therapy, which is then continued for maintenance.

TRIMETHOPRIM-SULFAMETHOXAZOLE

Although certain reports have indicated that TMP-SMX may be of benefit in the treatment of granulomatosis with polyangiitis (Wegener's) isolated to the sinonasal tissues, it should never be used alone to treat active granulomatosis with polyangiitis (Wegener's) outside of the upper airway such as in patients with renal or pulmonary disease. In a study examining the effect of TMP-SMX on relapse, decreased relapses were shown only with regard to upper airway disease, and no differences in major organ relapses were observed.

ORGAN-SPECIFIC TREATMENT

Not all manifestations of granulomatosis with polyangiitis (Wegener's) require or respond to immunosuppressive therapy. In managing non-major organ disease, such as that isolated to the sinus, joints, or skin, the risks of treatment should be carefully weighed against the benefits. Treatment with cyclophosphamide is rarely if ever justified for the treatment of isolated sinus disease in granulomatosis with polyangiitis (Wegener's). Although patients with non-major organ disease may be effectively treated without immunosuppressive therapy, these individuals must be monitored closely for the development of disease activity affecting the lungs,

2186 kidneys, or other major organs. Subglottic stenosis and endobronchial stenosis are examples of disease manifestations that do not typically respond to systemic immunosuppressive treatment.

MICROSCOPIC POLYANGIITIS

DEFINITION

The term *microscopic polyarteritis* was introduced into the literature by Davson in 1948 in recognition of the presence of glomerulonephritis in patients with polyarteritis nodosa. In 1992, the Chapel Hill Consensus Conference on the Nomenclature of Systemic Vasculitis adopted the term *microscopic polyangiitis* to connote a necrotizing vasculitis with few or no immune complexes affecting small vessels (capillaries, venules, or arterioles). Glomerulonephritis is very common in microscopic polyangiitis, and pulmonary capillaritis often occurs. The absence of granulomatous inflammation in microscopic polyangiitis is said to differentiate it from granulomatosis with polyangiitis (Wegener's).

INCIDENCE AND PREVALENCE

The incidence of microscopic polyangiitis has not yet been reliably established due to its previous inclusion as part of polyarteritis nodosa. The mean age of onset is ~57 years of age, and males are slightly more frequently affected than females.

PATHOLOGY AND PATHOGENESIS

The vasculitis seen in microscopic polyangiitis has a predilection to involve capillaries and venules in addition to small and medium-sized arteries. Immunohistochemical staining reveals a paucity of immunoglobulin deposition in the vascular lesion of microscopic polyangiitis, suggesting that immune-complex formation does not play a role in the pathogenesis of this syndrome. The renal lesion seen in microscopic polyangiitis is identical to that of granulomatosis with polyangiitis (Wegener's). Like granulomatosis with polyangiitis (Wegener's), microscopic polyangiitis is highly associated with the presence of ANCA, which may play a role in pathogenesis of this syndrome (see above).

CLINICAL AND LABORATORY MANIFESTATIONS

Because of its predilection to involve the small vessels, microscopic polyangiitis and granulomatosis with polyangiitis (Wegener's) share similar clinical features. Disease onset may be gradual, with initial symptoms of fever, weight loss, and musculoskeletal pain; however, it is often acute. Glomerulonephritis occurs in at least 79% of patients and can be rapidly progressive, leading to renal failure. Hemoptysis may be the first symptom of alveolar hemorrhage, which occurs in 12% of patients. Other manifestations include mononeuritis multiplex and gastrointestinal tract and cutaneous vasculitis. Upper airway disease and pulmonary nodules are not typically found in microscopic polyangiitis and, if present, suggest granulomatosis with polyangiitis (Wegener's).

Features of inflammation may be seen, including an elevated ESR, anemia, leukocytosis, and thrombocytosis. ANCA are present in 75% of patients with microscopic polyangiitis, with antilysozyme antibodies being the predominant ANCA associated with this disease.

DIAGNOSIS

The diagnosis is based on histologic evidence of vasculitis or paucimmune glomerulonephritis in a patient with compatible clinical features of multisystem disease. Although microscopic polyangiitis is strongly ANCA-associated, no studies have as yet established the sensitivity and specificity of ANCA in this disease.

TREATMENT MICROSCOPIC POLYANGIITIS

The 5-year survival rate for patients with treated microscopic polyangiitis is 74%, with disease-related mortality occurring from alveolar hemorrhage or gastrointestinal, cardiac, or renal disease. Studies on treatment have come from trials that have included patients with granulomatosis with polyangiitis (Wegener's) or microscopic

polyangiitis. Currently, the treatment approach for microscopic polyangiitis is the same as is used for granulomatosis with polyangiitis (Wegener's) (see "Granulomatosis with Polyangiitis [Wegener's]" for a detailed description of this therapeutic regimen), and patients with immediately life-threatening disease should be treated with the combination of prednisone and daily cyclophosphamide or rituximab. Disease relapse has been observed in at least 34% of patients. Treatment for such relapses would be similar to that used at the time of initial presentation and based on site and severity of disease.

EOSINOPHILIC GRANULOMATOSIS WITH POLYANGIITIS (CHURG-STRAUSS)

DEFINITION

Eosinophilic granulomatosis with polyangiitis (Churg-Strauss) was described in 1951 by Churg and Strauss and is characterized by asthma, peripheral and tissue eosinophilia, extravascular granuloma formation, and vasculitis of multiple organ systems.

INCIDENCE AND PREVALENCE

Eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is an uncommon disease with an estimated annual incidence of 1–3 per million. The disease can occur at any age with the possible exception of infants. The mean age of onset is 48 years, with a female-to-male ratio of 1.2:1.

PATHOLOGY AND PATHOGENESIS

The necrotizing vasculitis of eosinophilic granulomatosis with polyangiitis (Churg-Strauss) involves small and medium-sized muscular arteries, capillaries, veins, and venules. A characteristic histopathologic feature of eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is granulomatous reactions that may be present in the tissues or even within the walls of the vessels themselves. These are usually associated with infiltration of the tissues with eosinophils. This process can occur in any organ in the body; lung involvement is predominant, with skin, cardiovascular system, kidney, peripheral nervous system, and gastrointestinal tract also commonly involved. Although the precise pathogenesis of this disease is uncertain, its strong association with asthma and its clinicopathologic manifestations, including eosinophilia, granuloma, and vasculitis, point to aberrant immunologic phenomena.

CLINICAL AND LABORATORY MANIFESTATIONS

Patients with eosinophilic granulomatosis with polyangiitis (Churg-Strauss) often exhibit nonspecific manifestations such as fever, malaise, anorexia, and weight loss, which are characteristic of a multisystem disease. The pulmonary findings in eosinophilic granulomatosis with polyangiitis (Churg-Strauss) clearly dominate the clinical picture with severe asthmatic attacks and the presence of pulmonary infiltrates. Mononeuritis multiplex is the second most common manifestation and occurs in up to 72% of patients. Allergic rhinitis and sinusitis develop in up to 61% of patients and are often observed early in the course of disease. Clinically recognizable heart disease occurs in ~14% of patients and is an important cause of mortality. Skin lesions occur in ~51% of patients and include purpura in addition to cutaneous and subcutaneous nodules. The renal disease in eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is less common and generally less severe than that of granulomatosis with polyangiitis and microscopic polyangiitis.

The characteristic laboratory finding in virtually all patients with eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is a striking eosinophilia, which reaches levels >1000 cells/ μ L in >80% of patients. Evidence of inflammation as evidenced by elevated ESR, fibrinogen, or α_2 -globulins can be found in 81% of patients. The other laboratory findings reflect the organ systems involved. Approximately 48% of patients with eosinophilic granulomatosis with polyangiitis (Churg-Strauss) have circulating ANCA that is usually antilysozyme.

DIAGNOSIS

Although the diagnosis of eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is optimally made by biopsy in a patient with the characteristic clinical manifestations (see above), histologic confirmation can be challenging because the pathognomonic features often do not occur simultaneously. In order to be diagnosed with eosinophilic granulomatosis with polyangiitis (Churg-Strauss), a patient should have evidence of asthma, peripheral blood eosinophilia, and clinical features consistent with vasculitis.

TREATMENT **EOSINOPHILIC GRANULOMATOSIS WITH POLYANGIITIS (CHURG-STRAUSS)**

The prognosis of untreated eosinophilic granulomatosis with polyangiitis (Churg-Strauss) is poor, with a reported 5-year survival of 25%. With treatment, prognosis is favorable, with one study finding a 78-month actuarial survival rate of 72%. Myocardial involvement is the most frequent cause of death and is responsible for 39% of patient mortality. Echocardiography should be performed in all newly diagnosed patients because this may influence therapeutic decisions.

Glucocorticoids alone appear to be effective in many patients. Dosage tapering is often limited by asthma, and many patients require low-dose prednisone for persistent asthma many years after clinical recovery from vasculitis. In glucocorticoid failure or in patients who present with fulminant multisystem disease, particularly cardiac involvement, the treatment of choice is a combined regimen of daily cyclophosphamide and prednisone (see “Granulomatosis with Polyangiitis [Wegener’s]” for a detailed description of this therapeutic regimen). Recent studies of mepolizumab (anti-IL-5 antibody) in eosinophilic granulomatosis with polyangiitis (Churg-Strauss) have been encouraging, but this treatment requires further investigation.

POLYARTERITIS NODOSA**DEFINITION**

Polyarteritis nodosa was described in 1866 by Kussmaul and Maier. It is a multisystem, necrotizing vasculitis of small and medium-sized muscular arteries in which involvement of the renal and visceral arteries is characteristic. Polyarteritis nodosa does not involve pulmonary arteries, although bronchial vessels may be involved; granulomas, significant eosinophilia, and an allergic diathesis are not observed.

INCIDENCE AND PREVALENCE

It is difficult to establish an accurate incidence of polyarteritis nodosa because previous reports have included polyarteritis nodosa and microscopic polyangiitis as well as other related vasculitides. Polyarteritis nodosa, as currently defined, is felt to be a very uncommon disease.

PATHOLOGY AND PATHOGENESIS

The vascular lesion in polyarteritis nodosa is a necrotizing inflammation of small and medium-sized muscular arteries. The lesions are segmental and tend to involve bifurcations and branchings of arteries. They may spread circumferentially to involve adjacent veins. However, involvement of venules is not seen in polyarteritis nodosa and, if present, suggests microscopic polyangiitis (see below). In the acute stages of disease, polymorphonuclear neutrophils infiltrate all layers of the vessel wall and perivascular areas, which results in intimal proliferation and degeneration of the vessel wall. Mononuclear cells infiltrate the area as the lesions progress to the subacute and chronic stages. Fibrinoid necrosis of the vessels ensues with compromise of the lumen, thrombosis, infarction of the tissues supplied by the involved vessel, and, in some cases, hemorrhage. As the lesions heal, there is collagen deposition, which may lead to further occlusion of the vessel lumen. Aneurysmal dilations up to 1 cm in size along the involved arteries are characteristic of polyarteritis nodosa. Granulomas and substantial eosinophilia with eosinophilic tissue infiltrations are not characteristically found and suggest eosinophilic granulomatosis with polyangiitis (Churg-Strauss) (see above).

Multiple organ systems are involved, and the clinicopathologic findings reflect the degree and location of vessel involvement and the resulting ischemic changes. As mentioned above, pulmonary arteries are not involved in polyarteritis nodosa, and bronchial artery involvement is uncommon. The pathology in the kidney in classic polyarteritis nodosa is that of arteritis without glomerulonephritis. In patients with significant hypertension, typical pathologic features of glomerulosclerosis may be seen. In addition, pathologic sequelae of hypertension may be found elsewhere in the body.

The presence of a polyarteritis nodosa–like vasculitis in patients with hepatitis B together with the isolation of circulating immune complexes composed of hepatitis B antigen and immunoglobulin, and the demonstration by immunofluorescence of hepatitis B antigen, IgM, and complement in the blood vessel walls, strongly suggest the role of immunologic phenomena in the pathogenesis of this disease. A polyarteritis nodosa–like vasculitis has also been reported in patients with hepatitis C. Hairy cell leukemia can be associated with polyarteritis nodosa; the pathogenic mechanisms of this association are unclear.

CLINICAL AND LABORATORY MANIFESTATIONS

Nonspecific signs and symptoms are the hallmarks of polyarteritis nodosa. Fever, weight loss, and malaise are present in over one-half of cases. Patients usually present with vague symptoms such as weakness, malaise, headache, abdominal pain, and myalgias that can rapidly progress to a fulminant illness. Specific complaints related to the vascular involvement within a particular organ system may also dominate the presenting clinical picture as well as the entire course of the illness (Table 385-6). In polyarteritis nodosa, renal involvement most commonly manifests as hypertension, renal insufficiency, or hemorrhage due to microaneurysms.

There are no diagnostic serologic tests for polyarteritis nodosa. In >75% of patients, the leukocyte count is elevated with a predominance of neutrophils. Eosinophilia is seen only rarely and, when present at high levels, suggests the diagnosis of eosinophilic granulomatosis with polyangiitis (Churg-Strauss). The anemia of chronic disease may be seen, and an elevated ESR is almost always present. Other common laboratory findings reflect the particular organ involved. Hypergammaglobulinemia may be present, and all patients should be screened for hepatitis B and C. Antibodies against myeloperoxidase or proteinase-3 (ANCA) are rarely found in patients with polyarteritis nodosa.

DIAGNOSIS

The diagnosis of polyarteritis nodosa is based on the demonstration of characteristic findings of vasculitis on biopsy material of involved

TABLE 385-6 CLINICAL MANIFESTATIONS RELATED TO ORGAN SYSTEM INVOLVEMENT IN POLYARTERITIS NODOSA

Organ System	Percent Incidence	Clinical Manifestations
Renal	60	Renal failure, hypertension
Musculoskeletal	64	Arthritis, arthralgia, myalgia
Peripheral nervous system	51	Peripheral neuropathy, mononeuritis multiplex
Gastrointestinal tract	44	Abdominal pain, nausea and vomiting, bleeding, bowel infarction and perforation, cholecystitis, hepatic infarction, pancreatic infarction
Skin	43	Rash, purpura, nodules, cutaneous infarcts, livedo reticularis, Raynaud’s phenomenon
Cardiac	36	Congestive heart failure, myocardial infarction, pericarditis
Genitourinary	25	Testicular, ovarian, or epididymal pain
Central nervous system	23	Cerebral vascular accident, altered mental status, seizure

Source: From TR Cupps, AS Fauci: *The Vasculitides*. Philadelphia, Saunders, 1981.

2188 organs. In the absence of easily accessible tissue for biopsy, the arteriographic demonstration of involved vessels, particularly in the form of aneurysms of small and medium-sized arteries in the renal, hepatic, and visceral vasculature, is sufficient to make the diagnosis. This should consist of a catheter-directed dye arteriogram because magnetic resonance and computed tomography arteriograms do not have sufficient resolution at the current time to visualize the vessels affected in polyarteritis nodosa. Aneurysms of vessels are not pathognomonic of polyarteritis nodosa; furthermore, aneurysms need not always be present, and arteriographic findings may be limited to stenotic segments and obliteration of vessels. Biopsy of symptomatic organs such as nodular skin lesions, painful testes, and nerve/muscle provides the highest diagnostic yields.

TREATMENT POLYARTERITIS NODOSA

The prognosis of untreated polyarteritis nodosa is extremely poor, with a reported 5-year survival rate between 10 and 20%. Death usually results from gastrointestinal complications, particularly bowel infarcts and perforation, and cardiovascular causes. Intractable hypertension often compounds dysfunction in other organ systems, such as the kidneys, heart, and CNS, leading to additional late morbidity and mortality in polyarteritis nodosa. With the introduction of treatment, survival rate has increased substantially. Favorable therapeutic results have been reported in polyarteritis nodosa with the combination of prednisone and cyclophosphamide (see “Granulomatosis with Polyangiitis [Wegener’s]” for a detailed description of this therapeutic regimen). In less severe cases of polyarteritis nodosa, glucocorticoids alone have resulted in disease remission. In patients with hepatitis B who have a polyarteritis nodosa-like vasculitis, antiviral therapy represents an important part of therapy and has been used in combination with glucocorticoids and plasma exchange. Careful attention to the treatment of hypertension can lessen the acute and late morbidity and mortality rates associated with renal, cardiac, and CNS complications of polyarteritis nodosa. Following successful treatment, relapse of polyarteritis nodosa has been estimated to occur in 10–20% of patients.

GIANT CELL ARTERITIS AND POLYMYALGIA RHEUMATICA

DEFINITION

Giant cell arteritis, historically referred to as *temporal arteritis*, is an inflammation of medium- and large-sized arteries. It characteristically involves one or more branches of the carotid artery, particularly the temporal artery. However, it is a systemic disease that can involve arteries in multiple locations, particularly the aorta and its main branches.

Giant cell arteritis is closely associated with *polymyalgia rheumatica*, which is characterized by stiffness, aching, and pain in the muscles of the neck, shoulders, lower back, hips, and thighs. Most commonly, polymyalgia rheumatica occurs in isolation, but it may be seen in 40–50% of patients with giant cell arteritis. In addition, ~10–20% of patients who initially present with features of isolated polymyalgia rheumatica later go on to develop giant cell arteritis. This strong clinical association together with data from pathophysiologic studies has increasingly supported that giant cell arteritis and polymyalgia rheumatica represent differing clinical spectrums of a single disease process.

INCIDENCE AND PREVALENCE

Giant cell arteritis occurs almost exclusively in individuals >50 years. It is more common in women than in men and is rare in blacks. The incidence of giant cell arteritis varies widely in different studies and in different geographic regions. A high incidence has been found in Scandinavia and in regions of the United States with large Scandinavian populations, compared to a lower incidence in southern Europe. The annual incidence rates in individuals ≥50 years range from 6.9 to 32.8 per 100,000 population. Familial aggregation has been reported, as has an association with HLA-DR4. In addition, genetic linkage studies have demonstrated an association of giant cell arteritis with alleles at the HLA-DRB1 locus, particularly

HLA-DRB1*04 variants. In Olmsted County, Minnesota, the annual incidence of polymyalgia rheumatica in individuals ≥50 years is 58.7 per 100,000 population.

PATHOLOGY AND PATHOGENESIS

Although the temporal artery is most frequently involved in giant cell arteritis, patients often have a systemic vasculitis of multiple medium- and large-sized arteries, which may go undetected. Histopathologically, the disease is a panarteritis with inflammatory mononuclear cell infiltrates within the vessel wall with frequent giant cell formation. There is proliferation of the intima and fragmentation of the internal elastic lamina. Pathophysiologic findings in organs result from the ischemia related to the involved vessels.

Experimental data support that giant cell arteritis is an antigen-driven disease in which activated T lymphocytes, macrophages, and dendritic cells play a critical role in the disease pathogenesis. Sequence analysis of the T cell receptor of tissue-infiltrating T cells in lesions of giant cell arteritis indicates restricted clonal expansion, suggesting the presence of an antigen residing in the arterial wall. Giant cell arteritis is believed to be initiated in the adventitia where CD4+ T cells enter through the vasa vasorum, become activated, and orchestrate macrophage differentiation. T cells recruited to vasculitic lesions in patients with giant cell arteritis produce predominantly IL-2 and IFN- γ , and the latter has been suggested to be involved in the progression to overt arteritis. Recent data demonstrate that at least two separate lineages of CD4 T cells—IFN- γ -producing T_H1 cells and IL-17-producing T_H17 cells—participate in vascular inflammation and may have differing levels of responsiveness to glucocorticoids.

CLINICAL AND LABORATORY MANIFESTATIONS

Giant cell arteritis is most commonly characterized clinically by the complex of fever, anemia, high ESR, and headaches in a patient over the age of 50 years. Other phenotypic manifestations include features of systemic inflammation including malaise, fatigue, anorexia, weight loss, sweats, arthralgias, polymyalgia rheumatica, or large-vessel disease.

In patients with involvement of the cranial arteries, headache is the predominant symptom and may be associated with a tender, thickened, or nodular artery, which may pulsate early in the disease but may become occluded later. Scalp pain and claudication of the jaw and tongue may occur. A well-recognized and dreaded complication of giant cell arteritis, particularly in untreated patients, is ischemic optic neuropathy, which may lead to serious visual symptoms, even sudden blindness in some patients. However, most patients have complaints relating to the head or eyes before visual loss. Attention to such symptoms with institution of appropriate therapy (see below) will usually avoid this complication. Other cranial ischemic complications include strokes and scalp or tongue infarction.

Up to one-third of patients can have large-vessel disease that can be the primary presentation of giant cell arteritis or can emerge at a later point in patients who have had previous cranial arteritis features or polymyalgia rheumatica. Manifestations of large-vessel disease can include subclavian artery stenosis that can present as arm claudication or aortic aneurysms involving the thoracic and to a lesser degree the abdominal aorta, which carry risks of rupture or dissection.

Characteristic laboratory findings in addition to the elevated ESR include a normochromic or slightly hypochromic anemia. Liver function abnormalities are common, particularly increased alkaline phosphatase levels. Increased levels of IgG and complement have been reported. Levels of enzymes indicative of muscle damage such as serum creatine kinase are not elevated.

DIAGNOSIS

The diagnosis of giant cell arteritis and its associated clinicopathologic syndrome can often be suggested clinically by the demonstration of the complex of fever, anemia, and high ESR with or without symptoms of polymyalgia rheumatica in a patient >50 years. The diagnosis is confirmed by biopsy of the temporal artery. Since involvement of the vessel may be segmental, positive yield is increased by obtaining a biopsy segment of 3–5 cm together with serial sectioning of biopsy specimens. Ultrasonography

of the temporal artery has been reported to be helpful in diagnosis. A temporal artery biopsy should be obtained as quickly as possible in the setting of ocular signs and symptoms, and under these circumstances, therapy should not be delayed pending a biopsy. In this regard, it has been reported that temporal artery biopsies may show vasculitis even after ~14 days of glucocorticoid therapy. A dramatic clinical response to a trial of glucocorticoid therapy can further support the diagnosis.

Large-vessel disease may be suggested by symptoms and findings on physical examination such as diminished pulses or bruits. It is confirmed by vascular imaging, most commonly through magnetic resonance or computed tomography.

Isolated polymyalgia rheumatica is a clinical diagnosis made by the presence of typical symptoms of stiffness, aching, and pain in the muscles of the hip and shoulder girdle, an increased ESR, the absence of clinical features suggestive of giant cell arteritis, and a prompt therapeutic response to low-dose prednisone.

TREATMENT GIANT CELL ARTERITIS AND POLYMYALGIA RHEUMATICA

Acute disease-related mortality directly from giant cell arteritis is very uncommon, with fatalities occurring from cerebrovascular events or myocardial infarction. However, patients are at risk of late mortality from aortic aneurysm rupture or dissection as patients with giant cell arteritis are 18 times more likely to develop thoracic aortic aneurysms than the general population.

The goals of treatment in giant cell arteritis are to reduce symptoms and, most importantly, to prevent visual loss. The treatment approach for cranial and large-vessel disease in giant cell arteritis is currently the same. Giant cell arteritis and its associated symptoms are exquisitely sensitive to glucocorticoid therapy. Treatment should begin with prednisone, 40–60 mg/d for ~1 month, followed by a gradual tapering. When ocular signs and symptoms occur, consideration should be given for the use of methylprednisolone 1000 mg daily for 3 days to protect remaining vision. Although the optimal duration of glucocorticoid therapy has not been established, most series have found that patients require treatment for ≥2 years. Symptom recurrence during prednisone tapering develops in 60–85% of patients with giant cell arteritis, requiring a dosage increase. The ESR can serve as a useful indicator of inflammatory disease activity in monitoring and tapering therapy and can be used to judge the pace of the tapering schedule. However, minor increases in the ESR can occur as glucocorticoids are being tapered and do not necessarily reflect an exacerbation of arteritis, particularly if the patient remains symptom-free. Under these circumstances, the tapering should continue with caution. Glucocorticoid toxicity occurs in 35–65% of patients and represents an important cause of patient morbidity. Aspirin 81 mg daily has been found to reduce the occurrence of cranial ischemic complications in giant cell arteritis and should be given in addition to glucocorticoids in patients who do not have contraindications. The use of weekly methotrexate as a glucocorticoid-sparing agent has been examined in two randomized placebo-controlled trials that reached conflicting conclusions. Infliximab, a monoclonal antibody to TNF, was studied in a randomized trial and was not found to provide benefit. Recent reports have shown favorable response of giant cell arteritis to tocilizumab (anti-IL-6 receptor), but this treatment requires further study before use in clinical practice.

Patients with isolated polymyalgia rheumatica respond promptly to prednisone, which can be started at a lower dose of 10–20 mg/d. Similar to giant cell arteritis, the ESR can serve as a useful indicator in monitoring and prednisone reduction. Recurrent polymyalgia symptoms develop in the majority of patients during prednisone tapering. One study of weekly methotrexate found that the use of this drug reduced the prednisone dose on average by only 1 mg and did not decrease prednisone-related side effects. A randomized trial in polymyalgia rheumatica did not find infliximab to lessen relapse or glucocorticoid requirements.

TAKAYASU ARTERITIS

DEFINITION

Takayasu arteritis is an inflammatory and stenotic disease of medium- and large-sized arteries characterized by a strong predilection for the aortic arch and its branches.

INCIDENCE AND PREVALENCE

Takayasu arteritis is an uncommon disease with an estimated annual incidence rate of 1.2–2.6 cases per million. It is most prevalent in adolescent girls and young women. Although it is more common in Asia, it is neither racially nor geographically restricted.

PATHOLOGY AND PATHOGENESIS

The disease involves medium- and large-sized arteries, with a strong predilection for the aortic arch and its branches; the pulmonary artery may also be involved. The most commonly affected arteries seen by arteriography are listed in [Table 385-7](#). The involvement of the major branches of the aorta is much more marked at their origin than distally. The disease is a panarteritis with inflammatory mononuclear cell infiltrates and occasionally giant cells. There are marked intimal proliferation and fibrosis, scarring and vascularization of the media, and disruption and degeneration of the elastic lamina. Narrowing of the lumen occurs with or without thrombosis. The vasa vasorum are frequently involved. Pathologic changes in various organs reflect the compromise of blood flow through the involved vessels.

Immunopathogenic mechanisms, the precise nature of which is uncertain, are suspected in this disease. As with several of the vasculitis syndromes, circulating immune complexes have been demonstrated, but their pathogenic significance is unclear.

CLINICAL AND LABORATORY MANIFESTATIONS

Takayasu arteritis is a systemic disease with generalized as well as vascular symptoms. The generalized symptoms include malaise, fever, night sweats, arthralgias, anorexia, and weight loss, which may occur months before vessel involvement is apparent. These symptoms may merge into those related to vascular compromise and organ ischemia. Pulses are commonly absent in the involved vessels, particularly the subclavian artery. The frequency of arteriographic abnormalities and the potentially associated clinical

TABLE 385-7 FREQUENCY OF ARTERIOGRAPHIC ABNORMALITIES AND POTENTIAL CLINICAL MANIFESTATIONS OF ARTERIAL INVOLVEMENT IN TAKAYASU ARTERITIS

Artery	Percentage of Arteriographic Abnormalities	Potential Clinical Manifestations
Subclavian	93	Arm claudication, Raynaud's phenomenon
Common carotid	58	Visual changes, syncope, transient ischemic attacks, stroke
Abdominal aorta ^a	47	Abdominal pain, nausea, vomiting
Renal	38	Hypertension, renal failure
Aortic arch or root	35	Aortic insufficiency, congestive heart failure
Vertebral	35	Visual changes, dizziness
Coeliac axis ^a	18	Abdominal pain, nausea, vomiting
Superior mesenteric ^a	18	Abdominal pain, nausea, vomiting
Iliac	17	Leg claudication
Pulmonary	10–40	Atypical chest pain, dyspnea
Coronary	<10	Chest pain, myocardial infarction

^aArteriographic lesions at these locations are usually asymptomatic but may potentially cause these symptoms.

Source: G Kerr et al: *Ann Intern Med* 120:919, 1994.

2190 manifestations are listed in Table 385-7. Hypertension occurs in 32–93% of patients and contributes to renal, cardiac, and cerebral injury.

Characteristic laboratory findings include an elevated ESR, mild anemia, and elevated immunoglobulin levels.

DIAGNOSIS

The diagnosis of Takayasu arteritis should be suspected strongly in a young woman who develops a decrease or absence of peripheral pulses, discrepancies in blood pressure, and arterial bruits. The diagnosis is confirmed by the characteristic pattern on arteriography, which includes irregular vessel walls, stenosis, poststenotic dilation, aneurysm formation, occlusion, and evidence of increased collateral circulation. Complete aortic arteriography by catheter-directed dye arteriography or magnetic resonance arteriography should be obtained in order to fully delineate the distribution and degree of arterial disease. Histopathologic demonstration of vessel wall inflammation that is predominantly lymphocytic with granuloma formation and giant cells involving the media and adventitia adds confirmatory data; however, tissue is rarely readily available for examination. IgG4-related disease is a potential cause of aortitis and periaortitis that is histologically differentiated from Takayasu arteritis by a dense lymphoplasmacytic infiltrate rich in IgG4-positive plasma cells, a storiform pattern of fibrosis, and obliterative phlebitis.

TREATMENT TAKAYASU ARTERITIS

The long-term outcome of patients with Takayasu arteritis has varied widely between studies. Although two North American reports found overall survival to be $\geq 94\%$, the 5-year mortality rate from other studies has ranged from 0 to 35%. Disease-related mortality most often occurs from congestive heart failure, cerebrovascular events, myocardial infarction, aneurysm rupture, or renal failure. Even in the absence of life-threatening disease, Takayasu arteritis can be associated with significant morbidity. The course of the disease is variable, and although spontaneous remissions may occur, Takayasu arteritis is most often chronic and relapsing. Although glucocorticoid therapy in doses of 40–60 mg prednisone per day alleviates symptoms, there are no convincing studies that indicate that it increases survival. The combination of glucocorticoid therapy for acute signs and symptoms and an aggressive surgical and/or arterioplasty approach to stenosed vessels has markedly improved outcome and decreased morbidity by lessening the risk of stroke, correcting hypertension due to renal artery stenosis, and improving blood flow to ischemic viscera and limbs. Unless it is urgently required, surgical correction of stenosed arteries should be undertaken only when the vascular inflammatory process is well controlled with medical therapy. In individuals who are refractory to or unable to taper glucocorticoids, methotrexate in doses up to 25 mg per week has yielded encouraging results. Preliminary results with anti-TNF therapies have been encouraging, but will require further study through randomized trials to determine efficacy.

IgA VASCULITIS (HENOCH-SCHÖNLEIN)

DEFINITION

IgA vasculitis (Henoch-Schönlein) is a small-vessel vasculitis characterized by palpable purpura (most commonly distributed over the buttocks and lower extremities), arthralgias, gastrointestinal signs and symptoms, and glomerulonephritis.

INCIDENCE AND PREVALENCE

IgA vasculitis (Henoch-Schönlein) is usually seen in children; most patients range in age from 4 to 7 years; however, the disease may also be seen in infants and adults. It is not a rare disease; in one series it accounted for between 5 and 24 admissions per year at a pediatric hospital. The male-to-female ratio is 1.5:1. A seasonal variation with a peak incidence in spring has been noted.

PATHOLOGY AND PATHOGENESIS

The presumptive pathogenic mechanism for IgA (Henoch-Schönlein) vasculitis is immune-complex deposition. A number of inciting antigens have been suggested including upper respiratory tract infections, various drugs, foods, insect bites, and immunizations. IgA is the antibody class most often seen in the immune complexes and has been demonstrated in the renal biopsies of these patients.

CLINICAL AND LABORATORY MANIFESTATIONS

In pediatric patients, palpable purpura is seen in virtually all patients; most patients develop polyarthralgias in the absence of frank arthritis. Gastrointestinal involvement, which is seen in almost 70% of pediatric patients, is characterized by colicky abdominal pain usually associated with nausea, vomiting, diarrhea, or constipation and is frequently accompanied by the passage of blood and mucus per rectum; bowel intussusception may occur. Renal involvement occurs in 10–50% of patients and is usually characterized by mild glomerulonephritis leading to proteinuria and microscopic hematuria, with red blood cell casts in the majority of patients (**Chap. 338**); it usually resolves spontaneously without therapy. Rarely, a progressive glomerulonephritis will develop. In adults, presenting symptoms are most frequently related to the skin and joints, while initial complaints related to the gut are less common. Although certain studies have found that renal disease is more frequent and more severe in adults, this has not been a consistent finding. However, the course of renal disease in adults may be more insidious and thus requires close follow-up. Myocardial involvement can occur in adults but is rare in children.

Laboratory studies generally show a mild leukocytosis, a normal platelet count, and occasionally eosinophilia. Serum complement components are normal, and IgA levels are elevated in about one-half of patients.

DIAGNOSIS

The diagnosis of IgA vasculitis (Henoch-Schönlein) is based on clinical signs and symptoms. Skin biopsy specimen can be useful in confirming leukocytoclastic vasculitis with IgA and C3 deposition by immunofluorescence. Renal biopsy is rarely needed for diagnosis but may provide prognostic information in some patients.

TREATMENT IgA VASCULITIS (HENOCH-SCHÖNLEIN)

The prognosis of IgA vasculitis (Henoch-Schönlein) is excellent. Mortality is exceedingly rare, and 1–5% of children progress to end-stage renal disease. Most patients recover completely, and some do not require therapy. Treatment is similar for adults and children. When glucocorticoid therapy is required, prednisone, in doses of 1 mg/kg per day and tapered according to clinical response, has been shown to be useful in decreasing tissue edema, arthralgias, and abdominal discomfort; however, it has not proved beneficial in the treatment of skin or renal disease and does not appear to shorten the duration of active disease or lessen the chance of recurrence. Patients with rapidly progressive glomerulonephritis have been anecdotally reported to benefit from intensive plasma exchange combined with cytotoxic drugs. Disease recurrences have been reported in 10–40% of patients.

CRYOGLOBULINEMIC VASCULITIS

DEFINITION

Cryoglobulins are cold-precipitable monoclonal or polyclonal immunoglobulins. Cryoglobulinemia may be associated with a systemic vasculitis characterized by palpable purpura, arthralgias, weakness, neuropathy, and glomerulonephritis. Although this can be observed in association with a variety of underlying disorders including multiple myeloma, lymphoproliferative disorders, connective tissue diseases, infection, and liver disease, in many instances it appears to be idiopathic. Because of the apparent absence of an underlying disease and the presence of cryoprecipitate containing oligoclonal/polyclonal immunoglobulins, this entity was referred to as *essential mixed cryoglobulinemia*. Since the discovery of hepatitis C, it has been established that the vast majority of patients

who were considered to have essential mixed cryoglobulinemia have cryoglobulinemic vasculitis related to hepatitis C infection.

INCIDENCE AND PREVALENCE

The incidence of cryoglobulinemic vasculitis has not been established. It has been estimated, however, that 5% of patients with chronic hepatitis C will develop cryoglobulinemic vasculitis.

PATHOLOGY AND PATHOGENESIS

Skin biopsies in cryoglobulinemic vasculitis reveal an inflammatory infiltrate surrounding and involving blood vessel walls, with fibrinoid necrosis, endothelial cell hyperplasia, and hemorrhage. Deposition of immunoglobulin and complement is common. Abnormalities of uninvolved skin including basement membrane alterations and deposits in vessel walls may be found. Membranoproliferative glomerulonephritis is responsible for 80% of all renal lesions in cryoglobulinemic vasculitis.

The association between hepatitis C and cryoglobulinemic vasculitis has been supported by the high frequency of documented hepatitis C infection, the presence of hepatitis C RNA and anti-hepatitis C antibodies in serum cryoprecipitates, evidence of hepatitis C antigens in vasculitic skin lesions, and the effectiveness of antiviral therapy (see below). Current evidence suggests that in the majority of cases, cryoglobulinemic vasculitis occurs when an aberrant immune response to hepatitis C infection leads to the formation of immune complexes consisting of hepatitis C antigens, polyclonal hepatitis C-specific IgG, and monoclonal IgM rheumatoid factor. The deposition of these immune complexes in blood vessel walls triggers an inflammatory cascade that results in cryoglobulinemic vasculitis.

CLINICAL AND LABORATORY MANIFESTATIONS

The most common clinical manifestations of cryoglobulinemic vasculitis are cutaneous vasculitis, arthritis, peripheral neuropathy, and glomerulonephritis. Renal disease develops in 10–30% of patients. Life-threatening rapidly progressive glomerulonephritis or vasculitis of the CNS, gastrointestinal tract, or heart occurs infrequently.

The presence of circulating cryoprecipitates is the fundamental finding in cryoglobulinemic vasculitis. Rheumatoid factor is almost always found and may be a useful clue to the disease when cryoglobulins are not detected. Hypocomplementemia occurs in 90% of patients. An elevated ESR and anemia occur frequently. Evidence for hepatitis C infection must be sought in all patients by testing for hepatitis C antibodies and hepatitis C RNA.

TREATMENT CRYOGLOBULINEMIC VASCULITIS

Acute mortality directly from cryoglobulinemic vasculitis is uncommon, but the presence of glomerulonephritis is a poor prognostic sign for overall outcome. In such patients, 15% progress to end-stage renal disease, with 40% later experiencing fatal cardiovascular disease, infection, or liver failure. As indicated above, the majority of cases are associated with hepatitis C infection. In such patients, treatment with antiviral therapy (Chap. 360) can prove beneficial and should be considered first-line therapy for hepatitis C-associated cryoglobulinemic vasculitis. Clinical improvement with antiviral therapy is dependent on the virologic response. Patients who clear hepatitis C from the blood have objective improvement in their vasculitis along with significant reductions in levels of circulating cryoglobulins, IgM, and rheumatoid factor. However, substantial portions of patients with hepatitis C do not have a sustained virologic response to such therapy, and the vasculitis typically relapses with the return of viremia. While transient improvement can be observed with glucocorticoids, a complete response is seen in only 7% of patients. Plasmapheresis and cytotoxic agents have been used in anecdotal reports. These observations have not been confirmed, and such therapies carry significant risks. Randomized trials with rituximab (anti-CD20) in hepatitis C-associated cryoglobulinemic vasculitis have provided evidence of benefit such that this agent should be considered in patients with active vasculitis either in combination with antiviral therapy or alone

in patients who have relapsed through, are intolerant to, or have contraindications to antiviral agents.

SINGLE-ORGAN VASCULITIS

The potential for vasculitis to affect single organs has become increasingly recognized. This has been defined as vasculitis in arteries or veins of any size in a single organ that has no features that indicate that it is a limited expression of a systemic vasculitis. Examples include isolated aortitis, testicular vasculitis, vasculitis of the breast, isolated cutaneous vasculitis, and primary CNS vasculitis. In some instances, this may be discovered at the time of surgery such as orchiectomy for a testicular mass where there is concern for neoplasm that is found instead to be vasculitis. Some patients originally diagnosed with single-organ vasculitis may later develop additional manifestations of a more systemic disease. In instances where there is no evidence of systemic vasculitis and the affected organ has been removed in its entirety, the patient may be followed closely without immunosuppressive therapy. In other instances, such as primary CNS vasculitis or some patients with isolated cutaneous vasculitis, medical intervention is warranted.

IDIOPATHIC CUTANEOUS VASCULITIS

DEFINITION

The term *cutaneous vasculitis* is defined broadly as inflammation of the blood vessels of the dermis. Due to its heterogeneity, cutaneous vasculitis has been described by a variety of terms including *hypersensitivity vasculitis* and *cutaneous leukocytoclastic angiitis*. However, cutaneous vasculitis is not one specific disease but a manifestation that can be seen in a variety of settings. In >70% of cases, cutaneous vasculitis occurs either as part of a primary systemic vasculitis or as a secondary vasculitis related to an inciting agent or an underlying disease (see “Secondary Vasculitis,” below). In the remaining 30% of cases, cutaneous vasculitis occurs idiopathically.

INCIDENCE AND PREVALENCE

Cutaneous vasculitis represents the most commonly encountered vasculitis in clinical practice. The exact incidence of idiopathic cutaneous vasculitis has not been determined due to the predilection for cutaneous vasculitis to be associated with an underlying process and the variability of its clinical course.

PATHOLOGY AND PATHOGENESIS

The typical histopathologic feature of cutaneous vasculitis is the presence of vasculitis of small vessels. Postcapillary venules are the most commonly involved vessels; capillaries and arterioles may be involved less frequently. This vasculitis is characterized by a *leukocytoclasia*, a term that refers to the nuclear debris remaining from the neutrophils that have infiltrated in and around the vessels during the acute stages. In the subacute or chronic stages, mononuclear cells predominate; in certain subgroups, eosinophilic infiltration is seen. Erythrocytes often extravasate from the involved vessels, leading to palpable purpura. *Cutaneous arteritis* can also occur, which involves slightly larger-sized vessels within the dermis.

CLINICAL AND LABORATORY MANIFESTATIONS

The hallmark of idiopathic cutaneous vasculitis is the predominance of skin involvement. Skin lesions may appear typically as palpable purpura; however, other cutaneous manifestations of the vasculitis may occur, including macules, papules, vesicles, bullae, subcutaneous nodules, ulcers, and recurrent or chronic urticaria. The skin lesions may be pruritic or even quite painful, with a burning or stinging sensation. Lesions most commonly occur in the lower extremities in ambulatory patients or in the sacral area in bedridden patients due to the effects of hydrostatic forces on the postcapillary venules. Edema may accompany certain lesions, and hyperpigmentation often occurs in areas of recurrent or chronic lesions.

There are no specific laboratory tests diagnostic of idiopathic cutaneous vasculitis. A mild leukocytosis with or without eosinophilia is characteristic, as is an elevated ESR. Laboratory studies should be

2192 aimed toward ruling out features to suggest an underlying disease or a systemic vasculitis.

DIAGNOSIS

The diagnosis of cutaneous vasculitis is made by the demonstration of vasculitis on biopsy. An important diagnostic principle in patients with cutaneous vasculitis is to search for an etiology of the vasculitis—be it an exogenous agent, such as a drug or an infection, or an endogenous condition, such as an underlying disease (Fig. 385-1). In addition, a careful physical and laboratory examination should be performed to rule out the possibility of systemic vasculitis. This should start with the least invasive diagnostic approach and proceed to the more invasive only if clinically indicated.

TREATMENT IDIOPATHIC CUTANEOUS VASCULITIS

When an antigenic stimulus is recognized as the precipitating factor in the cutaneous vasculitis, it should be removed; if this is a microbe, appropriate antimicrobial therapy should be instituted. If the vasculitis is associated with another underlying disease, treatment of the latter often results in resolution of the former. In situations where disease is apparently self-limited, no therapy, except possibly symptomatic therapy, is indicated. When cutaneous vasculitis persists and when there is no evidence of an inciting agent, an associated disease, or an underlying systemic vasculitis, the decision to treat should be based on weighing the balance between the degree of symptoms and the risk of treatment. Some cases of idiopathic cutaneous vasculitis resolve spontaneously, whereas others remit and relapse. In patients with persistent vasculitis, a variety of therapeutic regimens have been tried with variable results. In general, the treatment of idiopathic cutaneous vasculitis has not been satisfactory. Fortunately, since the disease is generally limited to the skin, this lack of consistent response to therapy usually does not lead to a life-threatening situation. Agents with which there have been anecdotal reports of success include dapsone, colchicine, hydroxychloroquine, and nonsteroidal anti-inflammatory agents. Glucocorticoids are often used in the treatment of idiopathic cutaneous vasculitis. Therapy is usually instituted as prednisone, 1 mg/kg per day, with rapid tapering where possible, either directly to discontinuation or by conversion to an alternate-day regimen followed by ultimate discontinuation. In cases that prove refractory to glucocorticoids, a trial of a cytotoxic agent may be indicated. Patients with chronic vasculitis isolated to cutaneous venules rarely respond dramatically to any therapeutic regimen, and cytotoxic agents should be used only as a last resort in these patients. Methotrexate and azathioprine have been used in such situations in anecdotal reports. Although cyclophosphamide is the most effective therapy for the systemic vasculitides, it should almost never be used for idiopathic cutaneous vasculitis because of the potential toxicity.

PRIMARY CENTRAL NERVOUS SYSTEM (CNS) VASCULITIS

Primary CNS vasculitis is an uncommon clinicopathologic entity characterized by vasculitis restricted to the vessels of the CNS without other apparent systemic vasculitis. The inflammatory process is usually composed of mononuclear cell infiltrates with or without granuloma formation.

Patients may present with headaches, altered mental function, and focal neurologic defects. Systemic symptoms are generally absent. Devastating neurologic abnormalities may occur depending on the extent of vessel involvement. The diagnosis can be suggested by abnormal magnetic resonance imaging of the brain, an abnormal lumbar puncture, and/or demonstration of characteristic vessel abnormalities on arteriography (Fig. 385-4), but it is confirmed by biopsy of the brain parenchyma and leptomeninges. In the absence of a brain biopsy, care should be taken not to misinterpret as true primary vasculitis arteriographic abnormalities that might actually be related to another

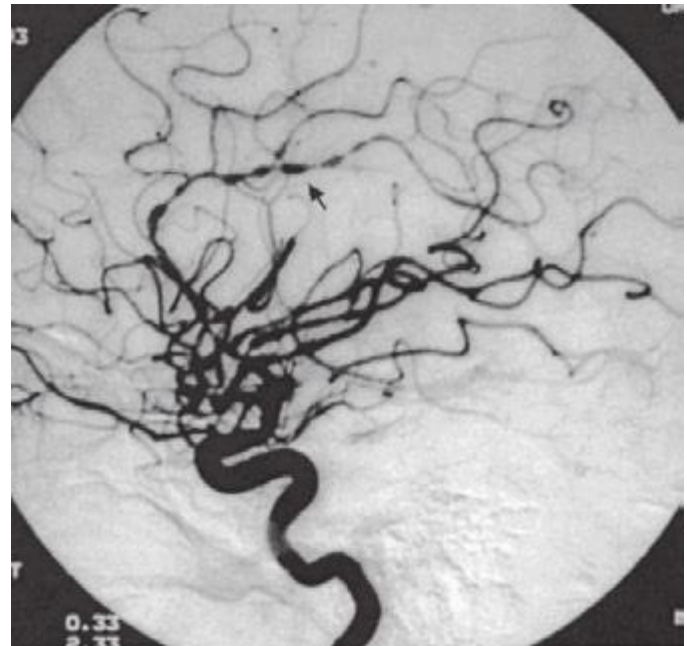


FIGURE 385-4 Cerebral arteriogram from a 32-year-old male with primary central nervous system vasculitis. Dramatic beading (arrow) typical of vasculitis is seen.

cause. An important entity in the differential diagnosis is reversible cerebral vasoconstrictive syndrome, which typically presents with “thunderclap” headache and is associated with arteriographic abnormalities that mimic primary CNS vasculitis that are reversible. Other diagnostic considerations include infection, atherosclerosis, emboli, connective tissue disease, sarcoidosis, malignancy, and drug-associated causes. The prognosis of granulomatous primary CNS vasculitis is poor; however, some reports indicate that glucocorticoid therapy, alone or together with cyclophosphamide administered as described above, has induced clinical remissions.

BEHÇET’S DISEASE

Behçet’s disease is a clinicopathologic entity characterized by recurrent episodes of oral and genital ulcers, iritis, and cutaneous lesions. The underlying pathologic process is a leukocytoclastic venulitis, although vessels of any size and in any organ can be involved. [This disorder is described in detail in Chap. 387.](#)

COGAN’S SYNDROME

Cogan’s syndrome is characterized by interstitial keratitis together with vestibuloauditory symptoms. It may be associated with a systemic vasculitis, particularly aortitis with involvement of the aortic valve. Glucocorticoids are the mainstay of treatment. Initiation of treatment as early as possible after the onset of hearing loss improves the likelihood of a favorable outcome.

KAWASAKI DISEASE

Kawasaki disease is an acute, febrile, multisystem disease of children. Some 80% of cases occur prior to the age of 5, with the peak incidence occurring at ≤ 2 years. It is characterized by nonsuppurative cervical adenitis and changes in the skin and mucous membranes such as edema; congested conjunctivae; erythema of the oral cavity, lips, and palms; and desquamation of the skin of the fingertips. Although the disease is generally benign and self-limited, it is associated with coronary artery aneurysms in $\sim 25\%$ of cases, with an overall case fatality rate of 0.5–2.8%. These complications usually occur between the third and fourth weeks of illness during the convalescent stage. Vasculitis of

the coronary arteries is seen in almost all the fatal cases that have been autopsied. There is typical intimal proliferation and infiltration of the vessel wall with mononuclear cells. Beadlike aneurysms and thromboses may be seen along the artery. Other manifestations include pericarditis, myocarditis, myocardial ischemia and infarction, and cardiomegaly.

Apart from the up to 2.8% of patients who develop fatal complications, the prognosis of this disease for uneventful recovery is excellent. High-dose IV γ -globulin (2 g/kg as a single infusion over 10 h) together with aspirin (100 mg/kg per day for 14 days followed by 3–5 mg/kg per day for several weeks) have been shown to be effective in reducing the prevalence of coronary artery abnormalities when administered early in the course of the disease. Surgery may be necessary for Kawasaki disease patients who have giant coronary artery aneurysms or other coronary complications. Surgical treatment most commonly includes thromboendarterectomy, thrombus clearing, aneurysmal reconstruction, and coronary artery bypass grafting.

POLYANGIITIS OVERLAP SYNDROMES

Some patients with systemic vasculitis manifest clinicopathologic characteristics that do not fit precisely into any specific disease but have overlapping features of different vasculitides. Active systemic vasculitis in such settings has the same potential for causing irreversible organ system damage as when it occurs in one of the defined syndromes listed in Table 385-1. The diagnostic and therapeutic considerations as well as the prognosis for these patients depend on the sites and severity of active vasculitis. Patients with vasculitis that could potentially cause irreversible damage to a major organ system should be treated as described under “Granulomatosis with Polyangiitis (Wegener’s).”

SECONDARY VASCULITIS

DRUG-INDUCED VASCULITIS

Vasculitis associated with drug reactions usually presents as palpable purpura that may be generalized or limited to the lower extremities or other dependent areas; however, urticarial lesions, ulcers, and hemorrhagic blisters may also occur (Chap. 74). Signs and symptoms may be limited to the skin, although systemic manifestations such as fever, malaise, and polyarthralgias may occur. Although the skin is the predominant organ involved, systemic vasculitis may result from drug reactions. Drugs that have been implicated in vasculitis include allopurinol, thiazides, gold, sulfonamides, phenytoin, and penicillin (Chap. 74).

An increasing number of drugs have been reported to cause vasculitis associated with antimyeloperoxidase ANCA. Of these, the best evidence of causality exists for hydralazine and propylthiouracil. The clinical manifestations in ANCA-positive drug-induced vasculitis can range from cutaneous lesions to glomerulonephritis and pulmonary hemorrhage. Outside of drug discontinuation, treatment should be based on the severity of the vasculitis. Patients with immediately life-threatening small-vessel vasculitis should initially be treated with glucocorticoids and cyclophosphamide as described for granulomatosis with polyangiitis (Wegener’s). Following clinical improvement, consideration may be given for tapering such agents along a more rapid schedule.

SERUM SICKNESS AND SERUM SICKNESS LIKE REACTIONS

These reactions are characterized by the occurrence of fever, urticaria, polyarthralgias, and lymphadenopathy 7–10 days after primary exposure and 2–4 days after secondary exposure to a heterologous protein (classic serum sickness) or a nonprotein drug such as penicillin or sulfa (serum sickness–like reaction). Most of the manifestations are not due to a vasculitis; however, occasional patients will have typical cutaneous venulitis that may progress rarely to a systemic vasculitis.

VASCULITIS ASSOCIATED WITH OTHER UNDERLYING DISEASES

Certain *infections* may directly trigger an inflammatory vasculitic process. For example, rickettsias can invade and proliferate in the endothelial cells of small blood vessels causing a vasculitis (Chap. 211). In

addition, the inflammatory response around blood vessels associated with certain systemic fungal diseases such as histoplasmosis (Chap. 236) may mimic a primary vasculitic process. A leukocytoclastic vasculitis predominantly involving the skin with occasional involvement of other organ systems may be a minor component of many other infections. These include *subacute bacterial endocarditis*, *Epstein-Barr virus infection*, *HIV infection*, and a number of other infections.

Vasculitis can be associated with certain *malignancies*, particularly lymphoid or reticuloendothelial neoplasms. Leukocytoclastic venulitis confined to the skin is the most common finding; however, widespread systemic vasculitis may occur. Of particular note is the association of *hairy cell leukemia* (Chap. 134) with polyarteritis nodosa.

A number of *connective tissue diseases* have vasculitis as a secondary manifestation of the underlying primary process. Foremost among these are *systemic lupus erythematosus* (Chap. 378), *rheumatoid arthritis* (Chap. 380), *inflammatory myositis* (Chap. 388), *relapsing polychondritis* (Chap. 389), and *Sjögren’s syndrome* (Chap. 383). The most common form of vasculitis in these conditions is the small-vessel venulitis isolated to the skin. However, certain patients may develop a fulminant systemic necrotizing vasculitis.

Secondary vasculitis has also been observed in association with *ulcerative colitis*, *congenital deficiencies of various complement components*, *sarcoidosis*, *primary biliary cirrhosis*, α_1 -*antitrypsin deficiency*, and *intestinal bypass surgery*.

Diagnosis of the vasculitic syndromes is usually based on characteristic histologic or arteriographic findings in a patient who has clinically compatible features. The images provided in this atlas highlight some of the characteristic histologic and radiographic findings that may be seen in the vasculitic diseases. These images demonstrate the importance that tissue histology may have in securing the diagnosis of vasculitis, the utility of diagnostic imaging in the vasculitic diseases, and the improvements in the care of vasculitis patients that have resulted from radiologic innovations.

Tissue biopsies represent vital information in many patients with a suspected vasculitic syndrome, not only in confirming the presence of vasculitis and other characteristic histologic features, but also in ruling out other diseases that can have similar clinical presentations. The determination of where biopsies should be performed is based on the presence of clinical disease in an affected organ, the likelihood of a positive diagnostic yield from data contained in the published literature, and the risk of performing a biopsy in an affected site. Common sites where biopsies may be performed include the lung, kidney, and skin. Other sites such as sural nerve, brain, testicle, and gastrointestinal tissues may also demonstrate features of vasculitis and be appropriate locations for biopsy when clinically affected.

Surgical biopsies of radiographically abnormal pulmonary parenchyma have a diagnostic yield of 90% in patients with granulomatosis with polyangiitis (Wegener's) and play an important role in ruling out infection or malignancy. The yield of lung biopsies is highly associated with amount of tissue that can be obtained, and transbronchial biopsies, while less invasive, have a yield of only 7%. Lung biopsies also play an important role in microscopic polyangiitis, eosinophilic granulomatosis with polyangiitis (Churg-Strauss), and any vasculitic disease where an immunosuppressed patient has pulmonary disease that is suspected to be an infection.

Kidney biopsy findings of a focal, segmental, crescentic, necrotizing glomerulonephritis with few to no immune complexes (pauci-immune glomerulonephritis) are characteristic in patients with granulomatosis with polyangiitis (Wegener's), microscopic polyangiitis, or eosinophilic granulomatosis with polyangiitis (Churg-Strauss), who have active renal disease. These findings not only distinguish these entities from other causes of glomerulonephritis, but can also confirm the presence of active glomerulonephritis that requires treatment. As a result, renal biopsies can also be helpful to guide management decisions in these diseases when an established patient has worsening renal function and an inactive or equivocal urine sediment. Cryoglobulinemic

vasculitis and IgA vasculitis (Henoch-Schönlein) are other vasculitides where renal involvement may occur and where biopsy may be important in diagnosis or prognosis.

Biopsies of the skin are commonly performed and are well tolerated. Because not all purpuric or ulcerative lesions are due to vasculitis, skin biopsy plays an important role to confirm the presence of vasculitis as the cause of the manifestation. Cutaneous vasculitis represents the most common vasculitic feature that affects people and can be seen in a broad spectrum of settings including infections, medications, malignancies, and connective tissue diseases. As a result, for systemic vasculitides that will require aggressive immunosuppressive treatment, a skin biopsy may not represent sufficient evidence to secure the diagnosis.

Diagnostic imaging represents a critical assessment tool in patients who are known or suspected to have a systemic vasculitic disease. Imaging contributes unique information about the patient that, when taken together with the history, physical examination, and laboratory determinations, can guide the differential diagnosis and the subsequent assessment or treatment plan. A diverse range of imaging techniques is used in the assessment of vasculitis including plain radiography, ultrasonography, computed tomography (CT), magnetic resonance imaging (MRI), positron emission tomography, and catheter-directed dye arteriography. These procedures have specific utilities that can allow differing perspectives on the spectrum and severity of vasculitis.

For vasculitic diseases that involve the large- or medium-sized blood vessels, arteriography provides information regarding blood vessel stenoses or aneurysms that can support the diagnosis. Catheter-directed dye arteriography provides information on central blood pressure and offers the most precise detail regarding vessel lumen dimensions but carries risks related to dye exposure and the invasive nature of the procedure. Advancements in magnetic resonance (MR) and CT arteriography have brought about noninvasive options to view the lumen and vessel wall, thus enhancing the ability to perform serial studies for patient monitoring in large-vessel vasculitis. However, in patients suspected to have a medium-vessel vasculitis such as polyarteritis nodosa, catheter-directed dye arteriography should still be performed because MR and CT arteriograms do not currently have sufficient resolution to visualize arteries of this size.

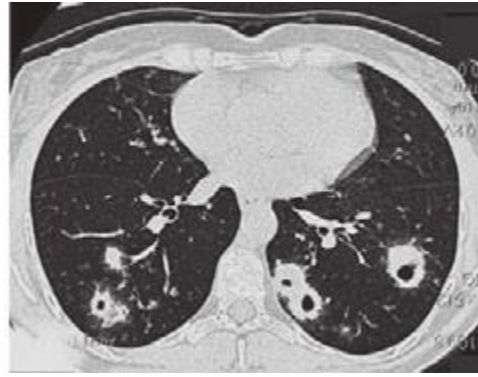
Although vasculitis involving the small blood vessels cannot be directly visualized, diagnostic imaging plays an essential role in detecting tissue injury that occurs as result of blood vessel and tissue inflammation. In granulomatosis with polyangiitis (Wegener's), 80% of patients may have pulmonary involvement during their disease course. Chest imaging should be obtained whenever active disease is suspected, because up to one-third of patients with radiographic abnormalities are asymptomatic. Pulmonary imaging is also important to detect complications of vasculitis therapy such as opportunistic pneumonias and medication-related pneumonitis.



FIGURE 386e-1 Bilateral nodular infiltrates seen on computed tomography of the chest in a 40-year-old woman with granulomatosis with polyangiitis (Wegener's).



A

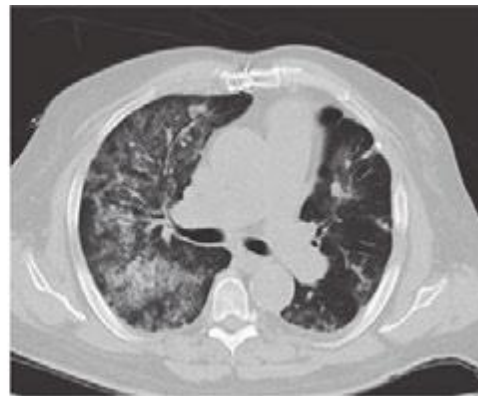


B

FIGURE 386e-2 Computed tomography of the chest in two patients with granulomatosis with polyangiitis (Wegener's) demonstrating (A) single and (B) multiple cavitary lung lesions.



A



B

FIGURE 386e-3 Bilateral ground-glass infiltrates due to alveolar hemorrhage from pulmonary capillaritis as seen in the same patient by (A) chest radiograph and (B) computed tomography. This manifestation can occur in granulomatosis with polyangiitis (Wegener's) or microscopic polyangiitis.



FIGURE 386e-4 Computed tomography of the chest demonstrating a dense infiltrate with air bronchograms involving a segment of the right upper lobe due to bacterial pneumonia in an immunosuppressed patient with granulomatosis with polyangiitis (Wegener's). Collapse of the left upper lobe secondary to endobronchial stenosis from granulomatosis with polyangiitis (Wegener's) also is seen on this image.

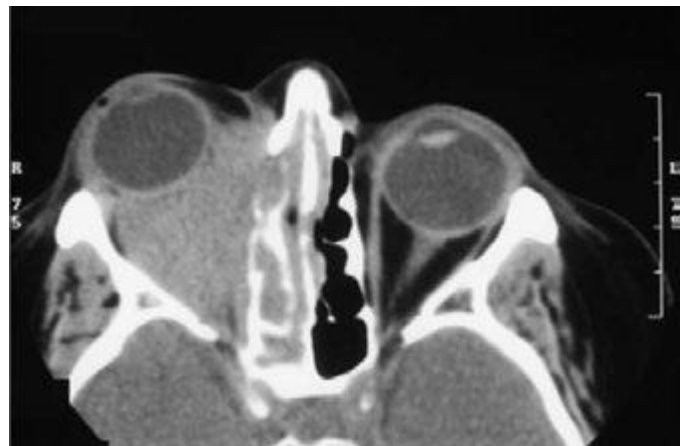
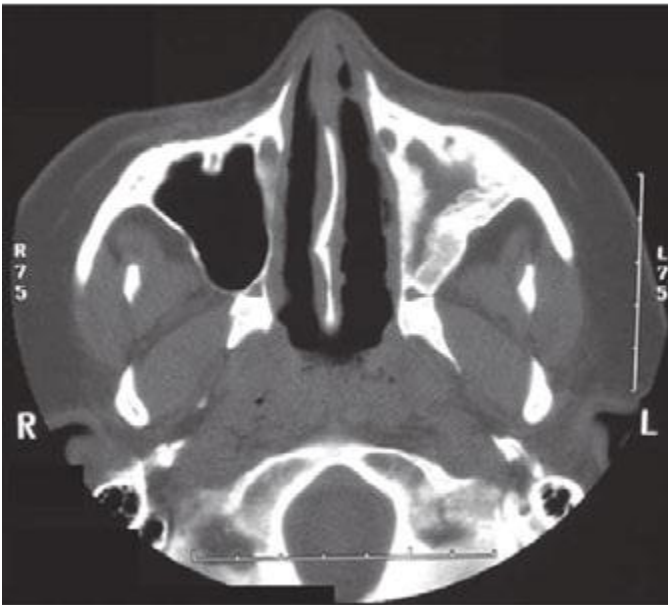


FIGURE 386e-5 Computed tomography of the orbits in a patient with granulomatosis with polyangiitis (Wegener's) who presented with right-eye proptosis. The image demonstrates inflammatory tissue extending from the ethmoid sinus through the lamina papyracea and filling the orbital space.



A



B

FIGURE 386e-6 Computed tomography of the sinuses in two patients with granulomatosis with polyangiitis (Wegener's). (**A**) Mucosal thickening of the bilateral maxillary sinuses and a perforation of the nasal septum. (**B**) Osteitis with obliteration of the left maxillary sinus in a patient with long-standing sinus disease.

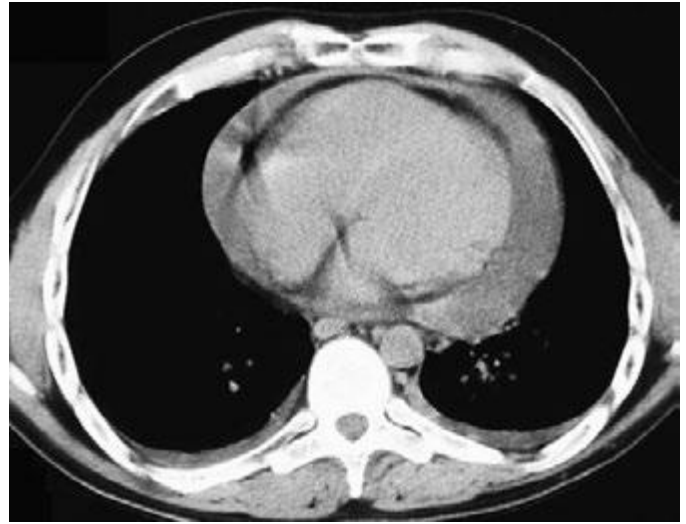


FIGURE 386e-7 Computed tomography of the chest demonstrating a large pericardial effusion in a patient with eosinophilic granulomatosis with polyangiitis (Churg-Strauss). Cardiac involvement is an important cause of morbidity and mortality in eosinophilic granulomatosis with polyangiitis and can include myocarditis, endocarditis, and pericarditis.

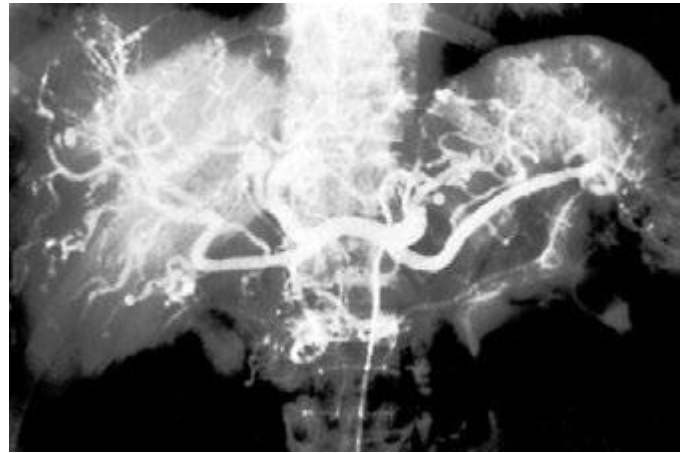


FIGURE 386e-8 Arteriogram of a 40-year-old man with polyarteritis nodosa demonstrating microaneurysms in the hepatic circulation.



FIGURE 386e-9 Arteriogram of a 19-year-old man with polyarteritis nodosa demonstrating multiple microaneurysms in the renal circulation. The patient presented with headache and severe hypertension that was due to medium-vessel vasculitis affecting the kidney.



FIGURE 386e-10 Cerebral arteriogram demonstrating beading along branches of the internal carotid artery in a patient with primary central nervous system vasculitis.

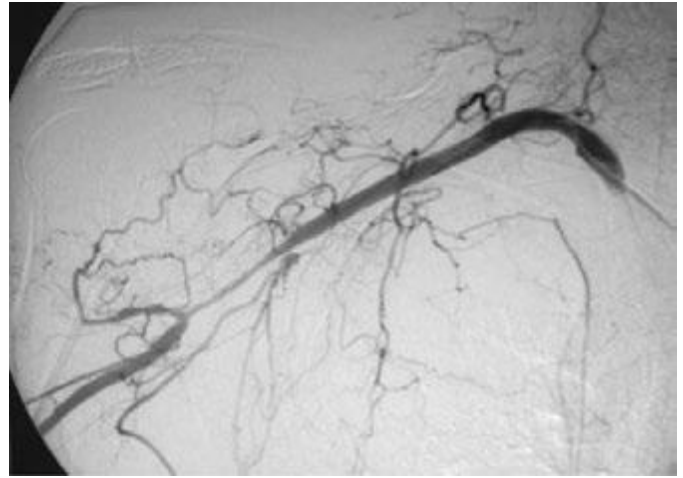


FIGURE 386e-11 Upper-extremity arteriogram demonstrating a long stenotic lesion of the axillary artery in a 75-year-old female with giant cell arteritis.



FIGURE 386e-12 Magnetic resonance imaging demonstrating extensive aneurysmal disease of the thoracic aorta in an 80-year-old female. The patient had been diagnosed with biopsy-proven giant cell arteritis 10 years prior to presenting with this aneurysm.



FIGURE 386e-13 Arteriogram of the aortic arch demonstrating complete occlusion of the left common carotid artery just after its origin from the aorta. This 20-year-old female presented with syncope and was subsequently diagnosed with Takayasu arteritis.



FIGURE 386e-14 Arteriogram demonstrating stenosis of the abdominal aorta in a 25-year-old female with Takayasu arteritis.

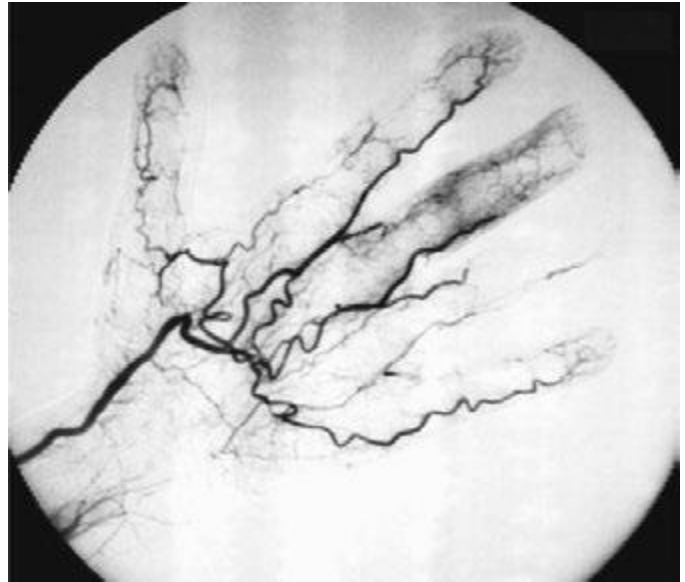


FIGURE 386e-15 Arteriogram of the hand demonstrating arterial skip lesions and vessel cutoffs in a patient with cryoglobulinemic vasculitis due to multiple myeloma.

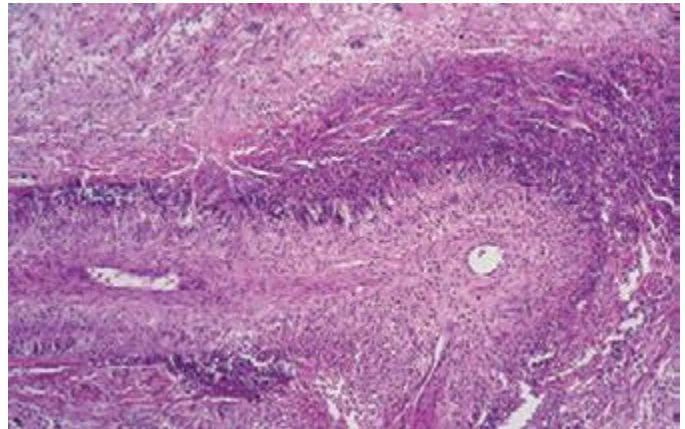


FIGURE 386e-16 Lung histology in granulomatosis with polyangiitis (Wegener's). This lung biopsy shows areas of geographic necrosis with a border of histiocytes and giant cells. There is also vasculitis with neutrophils, lymphocytes, and giant cells infiltrating the wall of an artery.

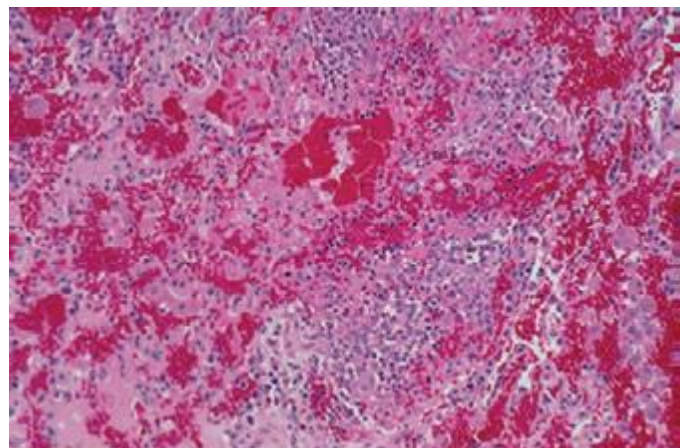


FIGURE 386e-17 Lung histology in microscopic polyangiitis. This lung biopsy demonstrates hemorrhage in the alveolar spaces due to capillaritis in a patient with microscopic polyangiitis. Similar findings can also be seen in granulomatosis with polyangiitis (Wegener's) and less commonly in eosinophilic granulomatosis with polyangiitis (Churg-Strauss).

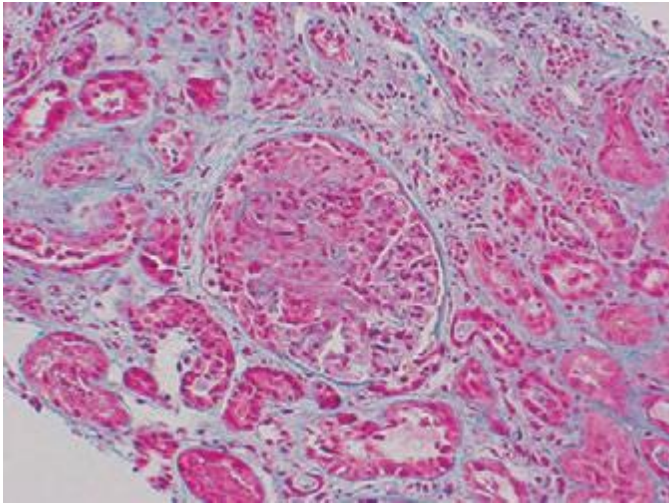


FIGURE 386e-18 Kidney biopsy in granulomatosis with polyangiitis (Wegener's). This renal biopsy shows a crescentic and necrotizing glomerulonephritis. These findings were focal and segmental with normal and scarred glomeruli also being found in the biopsy. By immunofluorescence and electron microscopy, no immune deposits were present, indicative of a pauci-immune glomerulonephritis. Similar findings can also be seen in microscopic polyangiitis and eosinophilic granulomatosis with polyangiitis (Churg-Strauss).

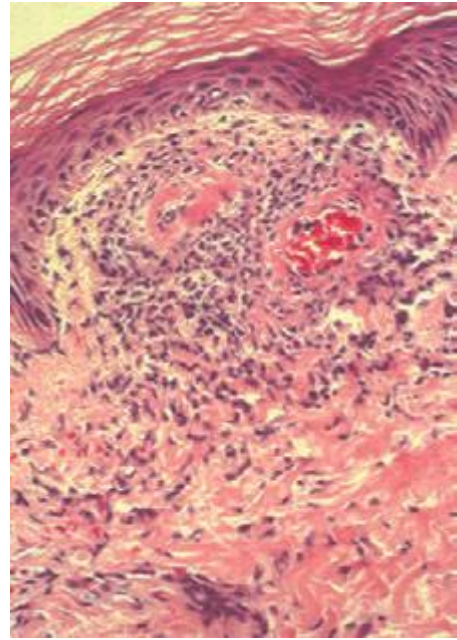


FIGURE 386e-21 Cutaneous vasculitis. This skin biopsy reveals two arterioles beneath the dermis with a neutrophilic inflammatory infiltrate in and around the vessel wall with leukocytoclasia (nuclear debris). While such features are diagnostic of vasculitis, they can be seen in a variety of settings and are not specific for any single disease.

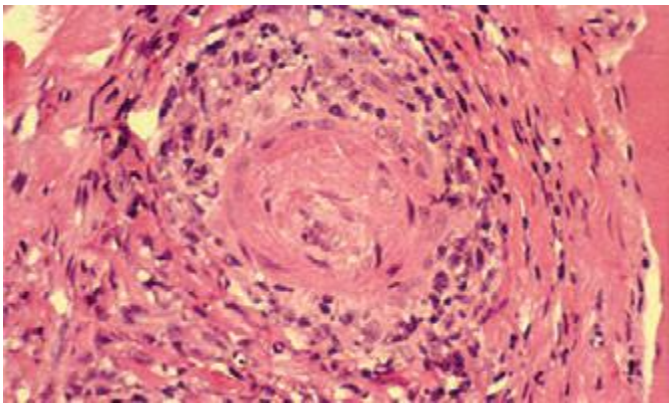


FIGURE 386e-19 Sural nerve biopsy in polyarteritis nodosa. This sural nerve biopsy was performed in a patient with polyarteritis nodosa, who had presented with a mononeuritis multiplex. Neutrophils are seen infiltrating all layers of this medium-sized vessel, which resulted in vessel occlusion and nerve infarction.

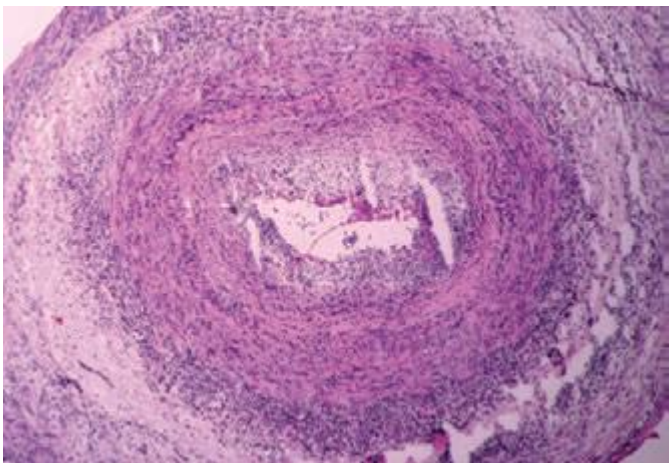


FIGURE 386e-20 Temporal artery biopsy in giant cell arteritis. This temporal artery biopsy demonstrates a panmural infiltration of mononuclear cells and lymphocytes that are particularly seen in the media and adventitia. Scattered giant cells are also present.

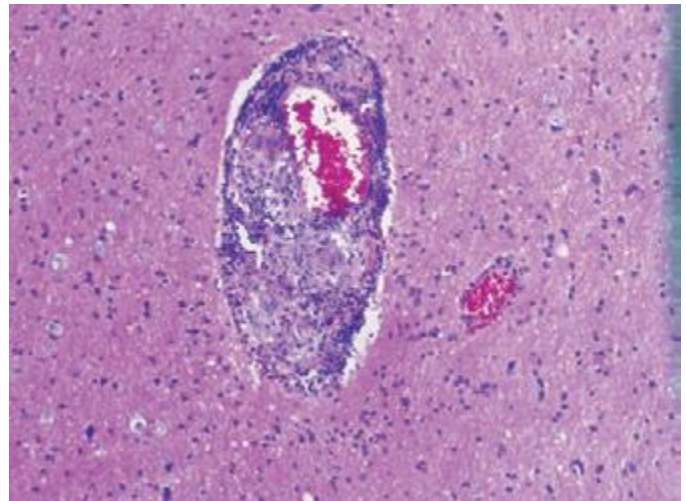


FIGURE 386e-22 Granulomatous primary central nervous system vasculitis. This brain biopsy demonstrates a medium-sized artery with granulomatous inflammation present within the vessel wall indicative of a granulomatous vasculitis. This patient presented with progressive headache, clinical and radiographic features of a stroke, and had arteriographic features consistent with vasculitis. Because no evidence of vasculitis could be found outside of the brain, this was consistent with granulomatous primary central nervous system vasculitis.

387 Behçet's Syndrome

Haralampos M. Moutsopoulos

DEFINITION, INCIDENCE, AND PREVALENCE

Behçet's syndrome is a multisystem disorder presenting with recurrent oral and genital ulcerations as well as ocular involvement. The diagnosis is clinical and based on internationally agreed diagnostic criteria (Table 387-1).

The syndrome affects young males and females from the Mediterranean region, the Middle East, and the Far East, suggesting a link with the ancient Silk Route. Males and females are affected equally, but males often have more severe disease. Blacks are very infrequently affected.

PATHOGENESIS

The etiology and pathogenesis of this syndrome remain obscure. The disease appears to be in the crossroads of autoinflammatory and autoimmune disorders. The main pathologic lesion is systemic perivasculitis with early neutrophil infiltration and endothelial swelling. In some patients, diffuse inflammatory disease, involving all layers of large vessels and resulting to formation of pseudoaneurysms, suggests vasculitis of vasa vasorum. Apart from activated neutrophils, increased numbers of infiltrating T_H1 , T_H17 , cytotoxic CD8+, and $\gamma\delta$ T cells are observed, suggesting a link between innate and adaptive autoreactive immune response. Circulating autoantibodies against α -enolase of endothelial cells, selenium binding protein, and anti-*Saccharomyces cerevisiae* antibodies have been observed, but their pathogenic role remains unclear. A recent genome-wide association study confirmed the known association of Behçet's syndrome with HLA-B*51 and identified a second, independent association within the major histocompatibility complex (MHC) class I region. In addition, an association with interleukin (IL) 10 and the IL-23R-IL-12RB2 locus was also observed. Interestingly, the disease-associated IL-10 variant was correlated with diminished mRNA expression and low protein production.

CLINICAL FEATURES

The recurrent aphthous ulcerations are a sine qua non for the diagnosis. The ulcers are usually painful, are shallow or deep with a central yellowish necrotic base, appear singly or in crops, and are located anywhere in the oral cavity. Small ulcers, less than 10 mm in diameter, are seen in 85% of patients, whereas large or herpetiform lesions are less frequent. The ulcers persist for 1–2 weeks and subside without leaving scars. The genital ulcers are less common but more specific, are painful, do not affect the glans penis or urethra, and produce scrotal scars.

Skin involvement is observed in 80% of patients and includes folliculitis, erythema nodosum, an acne-like exanthem, and, infrequently, vasculitis, Sweet syndrome, and pyoderma gangrenosum. Nonspecific skin inflammatory reactivity to any scratches or intradermal saline injection (pathergy test) is a common and specific manifestation.

Eye involvement with scarring and bilateral panuveitis is the most dreaded complication, since it occasionally progresses rapidly to blindness. The eye disease, occurring in 50% of patients, is usually present at the onset but may also develop within the first few years. In addition to iritis, posterior uveitis, retinal vessel occlusions, and optic neuritis can be seen in some patients with the syndrome.

Nondeforming arthritis or arthralgias are seen in 50% of patients and affect the knees and ankles.

Superficial or deep peripheral vein thrombosis is seen in 30% of patients. Pulmonary emboli are a rare complication. The superior vena cava is obstructed occasionally, producing a dramatic clinical picture. Arterial involvement occurs in less than 5% of patients and presents with aortitis or peripheral arterial aneurysm and arterial thrombosis. Pulmonary artery vasculitis presenting with dyspnea, cough, chest pain, hemoptysis, and infiltrates on chest roentgenograms has been reported in 5% of patients and should be differentiated from thromboembolic disease since it warrants anti-inflammatory and not thrombolytic therapy.

Neurologic involvement (5–10%) appears mainly in the parenchymal form (80%); it is associated with brainstem involvement and has a serious prognosis (*central nervous system [CNS]-Behçet's syndrome*). IL-6 is persistently raised in cerebrospinal fluid of these patients. Cerebral venous thrombosis is most frequently observed in the superior sagittal and transverse sinuses and is associated with headache and increased intracranial pressure. Magnetic resonance imaging (MRI) and/or proton magnetic resonance spectroscopy (MRS) are very sensitive and should be employed if CNS-Behçet's syndrome is suspected.

Gastrointestinal involvement is seen more frequently in patients from Japan and consists of mucosal ulcerations of the gut, resembling Crohn's disease.

Epididymitis is seen in 5% of patients, whereas amyloidosis of AA type and glomerulonephritis are uncommon.

Laboratory findings are mainly nonspecific indices of inflammation, such as leukocytosis and elevated erythrocyte sedimentation rate, as well as C-reactive protein levels.

TREATMENT BEHÇET'S SYNDROME

The severity of the syndrome usually abates with time. Apart from the patients with CNS-Behçet's syndrome and major vessel disease, the life expectancy seems to be normal and the only serious complication is blindness.

Mucous membrane involvement may respond to topical glucocorticoids in the form of mouthwash or paste. In more serious cases, thalidomide (100 mg/d) is effective. Thrombophlebitis is treated with aspirin, 325 mg/d. Colchicine can be beneficial for the mucocutaneous manifestations and arthritis. Uveitis and CNS-Behçet's syndrome require systemic glucocorticoid therapy (prednisone, 1 mg/kg per day) and azathioprine (2–3 mg/kg per day). Cyclosporine (5 mg/kg) has been used for sight-threatening uveitis, alone or in combination with azathioprine. Pulse doses of cyclophosphamide are useful early in the course of the disease for pulmonary or peripheral arterial aneurysms. Anti-tumor necrosis factor therapy is recommended in panuveitis refractory to immunosuppressives. Administration of this therapy improves visual acuity in more than two-thirds of patients.

388 Polymyositis, Dermatomyositis, and Inclusion Body Myositis

Marinos C. Dalakas

The inflammatory myopathies represent the largest group of acquired and potentially treatable causes of skeletal muscle weakness. They are classified into three major groups: polymyositis (PM), dermatomyositis (DM), and inclusion body myositis (IBM).

CLINICAL FEATURES

The prevalence of the inflammatory myopathies is estimated at 1 in 100,000. PM as a stand-alone entity is a rare disease. DM affects both

TABLE 387-1 DIAGNOSTIC CRITERIA OF BEHÇET'S SYNDROME

Recurrent oral ulceration plus two of the following:

- Recurrent genital ulceration
- Eye lesions
- Skin lesions
- Pathergy test

children and adults and women more often than men. IBM is three times more frequent in men than in women, more common in whites than blacks, and is most likely to affect persons age >50 years.

These disorders present as progressive and symmetric muscle weakness except for IBM, which can have an asymmetric pattern. Patients usually report increasing difficulty with everyday tasks requiring the use of proximal muscles, such as getting up from a chair, climbing steps, stepping onto a curb, lifting objects, or combing hair. Fine-motor movements that depend on the strength of distal muscles, such as buttoning a shirt, sewing, knitting, or writing, are affected only late in the course of PM and DM, but fairly early in IBM. Falling is common in IBM because of early involvement of the quadriceps muscle, with buckling of the knees. Ocular muscles are spared, even in advanced, untreated cases; if these muscles are affected, the diagnosis of inflammatory myopathy should be questioned. Facial muscles are unaffected in PM and DM, but mild facial muscle weakness is common in patients with IBM. In all forms of inflammatory myopathy, pharyngeal and neck-flexor muscles are often involved, causing dysphagia or difficulty in holding up the head (*head drop*). In advanced and rarely in acute cases, respiratory muscles may also be affected. Severe weakness, if untreated, is almost always associated with muscle wasting. Sensation remains normal. The tendon reflexes are preserved but may be absent in severely weakened or atrophied muscles, especially in IBM, where atrophy of the quadriceps and the distal muscles is common. Myalgia and muscle tenderness may occur in a small number of patients, usually early in the disease, and particularly in DM associated with connective tissue disorders. Weakness in PM and DM progresses subacutely over a period of weeks or months and rarely acutely; by contrast, IBM progresses very slowly, over years, simulating a late-life muscular dystrophy (Chap. 462e) or slowly progressive motor neuron disorder (Chap. 452).

SPECIFIC FEATURES

(Table 388-1)

Characteristic	Polymyositis	Dermatomyositis	Inclusion Body Myositis
Age at onset	>18 years	Adulthood and childhood	>50 years
Familial association	No	No	Yes, in some rare cases
Extramuscular manifestations	Yes	Yes	Yes
Associated conditions			
Connective tissue diseases	Yes ^a	Scleroderma and mixed connective tissue disease (overlap syndromes)	Yes, in up to 20% of cases ^a
Systemic autoimmune diseases ^b	Frequent	Infrequent	Infrequent
Malignancy	No	Yes, in up to 15% of cases	No
Viruses	Yes ^c	Unproven	Yes ^c
Drugs ^d	Yes	Yes, rarely	No
Parasites and bacteria ^e	Yes	No	No

^aSystemic lupus erythematosus, rheumatoid arthritis, Sjögren's syndrome, systemic sclerosis, mixed connective tissue disease. ^bCrohn's disease, vasculitis, sarcoidosis, primary biliary cirrhosis, adult celiac disease, chronic graft-versus-host disease, discoid lupus, ankylosing spondylitis, Behçet's syndrome, myasthenia gravis, acne fulminans, dermatitis herpetiformis, psoriasis, Hashimoto's disease, granulomatous diseases, agammaglobulinemia, monoclonal gammopathy, hyper eosinophilic syndrome, Lyme disease, Kawasaki disease, autoimmune thrombocytopenia, hypergammaglobulinemic purpura, hereditary complement deficiency, IgA deficiency. ^cHIV (human immunodeficiency virus) and HTLV-1 (human T cell lymphotropic virus type 1). ^dDrugs include penicillamine (dermatomyositis and polymyositis), zidovudine (polymyositis), statins (necrotizing, toxic, or autoimmune myositis), and contaminated tryptophan (dermatomyositis-like illness). Other myotoxic drugs may cause myopathy but not an inflammatory myopathy (see text for details). ^eParasites (protozoa, cestodes, nematodes), tropical and bacterial myositis (pyomyositis).

Polymyositis The actual onset of PM is often not easily determined, and patients typically delay seeking medical advice for several weeks or even months. This is in contrast to DM, in which the rash facilitates early recognition (see below). PM mimics many other myopathies and is a diagnosis of exclusion. It is a subacute inflammatory myopathy affecting adults, and rarely children, who *do not have* any of the following: rash, involvement of the extraocular and facial muscles, family history of a neuromuscular disease, history of exposure to myotoxic drugs or toxins, endocrinopathy, neurogenic disease, muscular dystrophy, biochemical muscle disorder (deficiency of a muscle enzyme), or IBM as excluded by muscle biopsy analysis (see below). As an isolated entity, PM is a rare (and overdiagnosed) disorder; more commonly, PM occurs in association with a systemic autoimmune or connective tissue disease or with a known viral or bacterial infection. Drugs, especially D-penicillamine, statins, or zidovudine (AZT), may also trigger an inflammatory myopathy similar to PM.

Dermatomyositis DM is a distinctive entity identified by a characteristic rash accompanying, or more often preceding, muscle weakness. The rash may consist of a blue-purple discoloration on the upper eyelids with edema (heliotrope rash; see Fig. 73-3), a flat red rash on the face and upper trunk, and erythema of the knuckles with a raised violaceous scaly eruption (*Gottron's sign*; see Fig. 73-4). The erythematous rash can also occur on other body surfaces, including the knees, elbows, malleoli, neck and anterior chest (often in a *V sign*), or back and shoulders (*shawl sign*), and may worsen after sun exposure. In some patients, the rash is pruritic, especially on the scalp, chest, and back. Dilated capillary loops at the base of the fingernails are also characteristic. The cuticles may be irregular, thickened, and distorted, and the lateral and palmar areas of the fingers may become rough and cracked, with irregular, "dirty" horizontal lines, resembling *mechanic's hands*. The weakness can be mild, moderate, or severe enough to lead to quadriparesis. At times, the muscle strength appears normal, hence the term *dermatomyositis sine myositis*. When muscle biopsy is performed in such cases, however, significant perivascular and perimysial inflammation is often seen.

DM usually occurs alone but may overlap with scleroderma and mixed connective tissue disease. Fasciitis and thickening of the skin, similar to that seen in chronic cases of DM, have occurred in patients with the *eosinophilia-myalgia syndrome* associated with the ingestion of contaminated L-tryptophan.

Inclusion Body Myositis In patients ≥ 50 years of age, IBM is the most common of the inflammatory myopathies. It is often misdiagnosed as PM and is suspected only later when a patient with presumed PM does not respond to therapy. Weakness and atrophy of the distal muscles, especially foot extensors and deep finger flexors, occur in almost all cases of IBM and may be a clue to early diagnosis. Some patients present with falls because their knees collapse due to early quadriceps weakness. Others present with weakness in the small muscles of the hands, especially finger flexors, and complain of inability to hold objects such as golf clubs or perform tasks such as turning keys or tying knots. On occasion, the weakness and accompanying atrophy can be asymmetric and selectively involve the quadriceps, iliopsoas, triceps, biceps, and finger flexors, resembling a lower motor neuron disease. Dysphagia is common, occurring in up to 60% of IBM patients, and may lead to episodes of choking. Sensory examination is generally normal; some patients have mildly diminished vibratory sensation at the ankles that presumably is age-related. The pattern of distal weakness, which superficially resembles motor neuron or peripheral nerve disease, results from the myopathic process affecting distal muscles selectively. Disease progression is slow but steady, and most patients require an assistive device such as cane, walker, or wheelchair within several years of onset.

In at least 20% of cases, IBM is associated with systemic autoimmune or connective tissue diseases. Familial aggregation of typical IBM may occur; such cases have been designated as *familial inflammatory IBM*. This disorder is distinct from *hereditary inclusion body myopathy* (h-IBM), which describes a heterogeneous group of recessive,

2196 and less frequently dominant, inherited syndromes; the h-IBMs are noninflammatory myopathies. A subset of h-IBM that spares the quadriceps muscles has emerged as a distinct entity. This disorder, originally described in Iranian Jews and now seen in many ethnic groups, is linked to chromosome 9p1 and results from mutations in the UDP-*N*-acetylglucosamine 2-epimerase/*N*-acetylmannosamine kinase (*GNE*) gene.

ASSOCIATED CLINICAL FINDINGS

Extramuscular Manifestations These may be present to a varying degree in patients with PM or DM, and include:

1. *Systemic symptoms*, such as fever, malaise, weight loss, arthralgia, and Raynaud's phenomenon, especially when inflammatory myopathy is associated with a connective tissue disorder.
2. *Joint contractures*, mostly in DM and especially in children.
3. *Dysphagia and gastrointestinal symptoms*, due to involvement of oropharyngeal striated muscles and upper esophagus, especially in DM and IBM.
4. *Cardiac disturbances*, including atrioventricular conduction defects, tachyarrhythmias, dilated cardiomyopathy, a low ejection fraction, and congestive heart failure, which may rarely occur either from the disease itself or from hypertension associated with long-term use of glucocorticoids.
5. *Pulmonary dysfunction*, due to weakness of the thoracic muscles, interstitial lung disease, or drug-induced pneumonitis (e.g., from methotrexate), which may cause dyspnea, nonproductive cough, and aspiration pneumonia. Interstitial lung disease may precede myopathy or occur early in the disease and develops in up to 10% of patients with PM or DM, most of whom have antibodies to t-RNA synthetases, as described below.
6. *Subcutaneous calcifications*, in DM, sometimes extruding on the skin and causing ulcerations and infections.
7. *Arthralgias, synovitis, or deforming arthropathy with subluxation* in the interphalangeal joints, which can occur in some patients with DM and PM who have Jo-1 antibodies (see below).

Association with Malignancies Although all the inflammatory myopathies can have a chance association with malignant lesions, especially in older age groups, the incidence of malignant conditions appears to be specifically increased only in patients with DM and not in those with PM or IBM. The most common tumors associated with DM are ovarian cancer, breast cancer, melanoma, colon cancer, and non-Hodgkin's lymphoma. The extent of the search that should be conducted for an occult neoplasm in adults with DM depends on the clinical circumstances. Tumors in these patients are usually uncovered by abnormal findings in the medical history and physical examination and not through an extensive blind search. The weight of evidence argues against performing expensive, invasive, and nondirected tumor searches. A complete annual physical examination with pelvic, breast (mammogram, if indicated), and rectal examinations (with colonoscopy according to age and family history); urinalysis; complete blood count; blood chemistry tests; and a chest film should suffice in most cases. In Asians, nasopharyngeal cancer is common, and a careful examination of ears, nose, and throat is indicated. If malignancy is clinically suspected, screening with a whole-body positron emission tomography (PET) scan should be considered.

Overlap Syndromes These describe the association of inflammatory myopathies with connective tissue diseases. A well-characterized overlap syndrome occurs in patients with DM who also have manifestations of systemic sclerosis or mixed connective tissue disease, such as sclerotic thickening of the dermis, contractures, esophageal hypomotility, microangiopathy, and calcium deposits (Table 388-1). By contrast, signs of rheumatoid arthritis, systemic lupus erythematosus, or Sjögren's syndrome are very rare in patients with DM. Patients with the overlap syndrome of DM and systemic sclerosis may have a specific antinuclear antibody, the anti-PM/Scl, directed against a nucleolar-protein complex.

PATHOGENESIS

An autoimmune etiology of the inflammatory myopathies is indirectly supported by an association with other autoimmune or connective tissue diseases; the presence of various autoantibodies; an association with specific major histocompatibility complex (MHC) genes; demonstration of T cell-mediated myocytotoxicity or complement-mediated microangiopathy; and a response to immunotherapy.

Autoantibodies and Immunogenetics Various autoantibodies against nuclear antigens (antinuclear antibodies) and cytoplasmic antigens are found in up to 30% of patients with inflammatory myopathies. The antibodies to cytoplasmic antigens are directed against ribonucleoproteins involved in protein synthesis (antisynthetases) or translational transport (anti-signal-recognition particles). The antibody directed against the histidyl-transfer RNA synthetase, called *anti-Jo-1*, accounts for 75% of all the antisynthetases and is clinically useful because up to 80% of patients with anti-Jo-1 antibodies have interstitial lung disease. Some patients with the anti-Jo-1 antibody also have Raynaud's phenomenon, nonerosive arthritis, and the MHC molecules DR3 and DRw52. DR3 haplotypes (molecular designation DRB1*0301, DQB1*0201) occur in up to 75% of patients with PM and IBM, whereas in juvenile DM, there is an increased frequency of DQA1*0501 (Chap. 373e). Antibodies against the cytosolic 5'-nucleotidase 1A, an enzyme abundantly expressed in muscle and thought to be involved in DNA degradation and repair, have been detected in one-third of IBM patients. Although the pathogenic significance of these antibodies is still unknown, they highlight the presence of an immune response, as discussed below.

Immunopathologic Mechanisms In DM, humoral immune mechanisms are implicated, resulting in a microangiopathy and muscle ischemia (Fig. 388-1). Endomysial inflammatory infiltrates are composed of B cells located in proximity to CD4 T cells, plasmacytoid dendritic cells, and macrophages; there is a relative absence of lymphocytic invasion of nonnecrotic muscle fibers. Activation of the complement C5b-9 membranolytic attack complex is thought to be a critical early event that triggers release of proinflammatory cytokines and chemokines, induces expression of vascular cell adhesion molecule (VCAM) 1 and intercellular adhesion molecule (ICAM) 1 on endothelial cells, and facilitates migration of activated lymphoid cells to the perimysial and endomysial spaces. Necrosis of the endothelial cells, reduced numbers of endomysial capillaries, ischemia, and muscle-fiber destruction resembling microinfarcts occur. The remaining capillaries often have dilated lumens in response to the ischemic process. Larger intramuscular blood vessels may also be affected in the same pattern. Residual perifascicular atrophy reflects the endofascicular hypoperfusion that is prominent in the periphery of muscle fascicles. Increased expression of type I interferon-inducible proteins is also noted in these regions.

By contrast, in PM and IBM, a mechanism of T cell-mediated cytotoxicity is likely. CD8 T cells, along with macrophages, initially surround and eventually invade and destroy healthy, nonnecrotic muscle fibers that aberrantly express class I MHC molecules. MHC-I expression, absent from the sarcolemma of normal muscle fibers, is probably induced by cytokines secreted by activated T cells and macrophages. The CD8/MHC-I complex is characteristic of PM and IBM; its detection can aid in confirming the histologic diagnosis of PM, as discussed below. The cytotoxic CD8 T cells contain perforin and granzyme granules directed toward the surface of the muscle fibers and capable of inducing myonecrosis. Analysis of T cell receptor molecules expressed by the infiltrating CD8 cells has revealed clonal expansion and conserved sequences in the antigen-binding region, both suggesting an antigen-driven T cell response. Whether the putative antigens are endogenous (e.g., muscle) or exogenous (e.g., viral) sequences is unknown. Viruses have not been identified within the muscle fibers. Co-stimulatory molecules and their counterreceptors, which are fundamental for T cell activation and antigen recognition, are strongly upregulated in PM and IBM. As noted above, the possibility that B cells and the humoral immune system might also play a role in IBM is suggested by the identification of anti-cytosolic 5'-nucleotidase

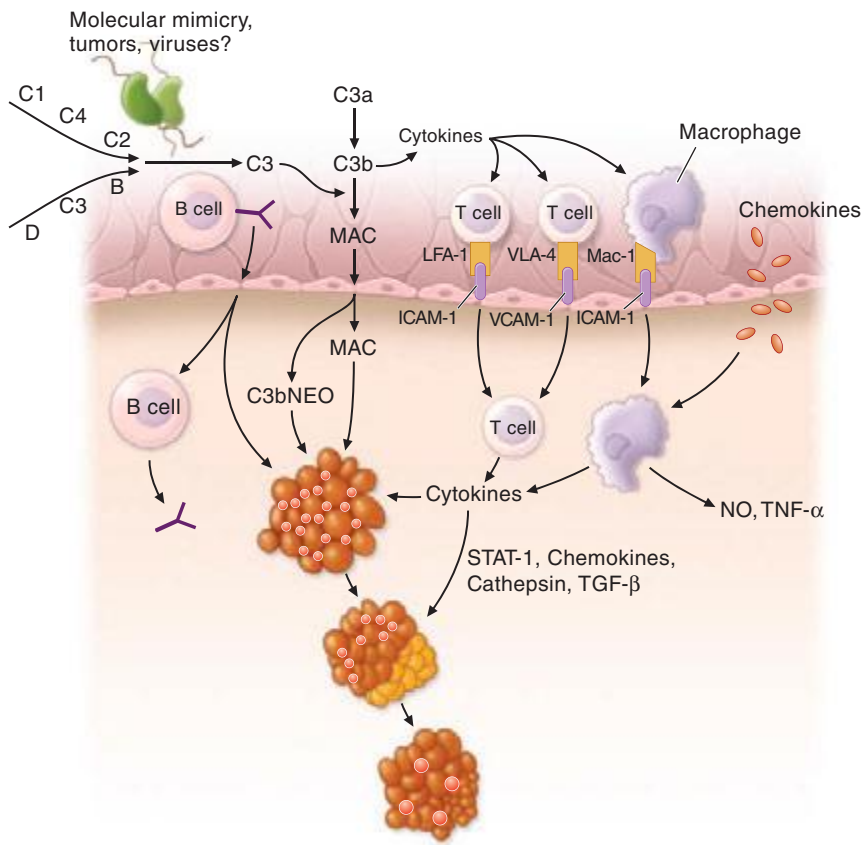


FIGURE 388-1 Immunopathogenesis of dermatomyositis. Activation of complement, possibly by autoantibodies (Y), against endothelial cells and formation of C3 via the classic or alternative pathway. Activated C3 leads to formation of C3b, C3bNEO, and membrane attack complexes (MAC), which are deposited in and around the endothelial cell wall of the endomysial capillaries. Deposition of MAC leads to destruction of capillaries, ischemia, or microinfarcts, most prominent in the periphery of the fascicles, and perifascicular atrophy. B cells, plasmacytoid dendritic cells, CD4 T cells, and macrophages traffic from the circulation to the muscle. Endothelial expression of vascular cell adhesion molecule (VCAM) and intercellular adhesion molecule (ICAM) is induced by cytokines released by the mononuclear cells. Integrins, specifically very late activation antigen (VLA)-4 and lymphocyte function-associated antigen (LFA)-1, bind VCAM and ICAM and promote T cell and macrophage infiltration of muscle through the endothelial cell wall.

1A antibodies in some patients. Key molecules involved in T cell-mediated cytotoxicity are depicted in [Fig. 388-2](#).

The Role of Nonimmune Factors in IBM In IBM, the presence of Congo red-positive amyloid deposits within some vacuolated muscle fibers and abnormal mitochondria with cytochrome oxidase-negative fibers suggest that, in addition to the autoimmune component, there is also a degenerative process. Similar to Alzheimer's disease, the intracellular amyloid deposits in IBM are immunoreactive against amyloid precursor protein (APP), β -amyloid, chymotrypsin, apolipoprotein E, presenilin, ubiquitin, and phosphorylated tau, but it is unclear whether these deposits, which are also seen in other vacuolar myopathies, are directly pathogenic or represent secondary phenomena. The same is true for the mitochondrial abnormalities, which may also be secondary to the effects of aging or a bystander effect of upregulated cytokines. Expression of cytokines and upregulation of MHC class I by the muscle fibers may cause an endoplasmic reticulum stress response resulting in intracellular accumulation of stressor molecules or misfolded glycoproteins and activation of nuclear factor κ B (NF- κ B), leading to further cytokine activation.

Association with Viral Infections and the Role of Retroviruses Several viruses, including coxsackieviruses, influenza, paramyxoviruses, mumps, cytomegalovirus, and Epstein-Barr virus, have been indirectly associated with myositis. For the coxsackieviruses, an autoimmune

myositis triggered by molecular mimicry has been proposed because of structural homology between histidyl-transfer RNA synthetase that is the target of the Jo-1 antibody (see above) and genomic RNA of an animal picornavirus, the encephalomyocarditis virus. Sensitive polymerase chain reaction (PCR) studies, however, have repeatedly failed to confirm the presence of such viruses in muscle biopsies.

The best evidence of a viral connection in PM and IBM is with the retroviruses. Some individuals infected with HIV or with human T cell lymphotropic virus 1 (HTLV-1) develop PM or IBM; a similar disorder has been described in nonhuman primates infected with the simian immunodeficiency virus. The inflammatory myopathy may occur as the initial manifestation of a retroviral infection, or myositis may develop later in the disease course. Retroviral antigens have been detected only in occasional endomysial macrophages and not within the muscle fibers themselves, suggesting that persistent infection and viral replication within the muscle does not occur. Histologic findings are identical to retroviral-negative PM or IBM. The infiltrating T cells in the muscle have a clonal bias and a number of them are retroviral-specific. This disorder should be distinguished from a toxic myopathy related to long-term therapy with AZT, characterized by fatigue, myalgia, mild muscle weakness, and mild elevation of creatine kinase (CK). AZT-induced myopathy, which generally improves when the drug is discontinued, is a mitochondrial disorder characterized histologically by "ragged-red" fibers. AZT inhibits γ -DNA polymerase, an enzyme found solely in the mitochondrial matrix.

GLOBAL ISSUES



Inadequate data exist with respect to possible differences in the prevalence of the inflammatory myopathies in various parts of the world. PM appears to be reported more often in Asia and southern Europe, whereas IBM seems to be more frequently recognized in North America, northern Europe, and Australia. Whether this represents differences in diagnostic methods and disease awareness or true disease prevalence remains unclear. Pyomyositis and parasitic myositis are clearly more common in the tropics, whereas HIV-associated PM and IBM are more commonly encountered in areas endemic for HIV. In patients from Asia, nasopharyngeal cancer appears to be a malignancy more commonly associated with DM, necessitating the need to specifically search for these tumors in this population.

DIFFERENTIAL DIAGNOSIS

The clinical picture of the typical skin rash and proximal or diffuse muscle weakness has few causes other than DM. However, proximal muscle weakness without skin involvement can be due to many conditions other than PM or IBM.

Subacute or Chronic Progressive Muscle Weakness This may be due to denervating conditions such as the spinal muscular atrophies or amyotrophic lateral sclerosis ([Chap. 452](#)). In addition to the muscle weakness, upper motor neuron signs in the latter and signs of denervation detected by electromyography (EMG) aid in the diagnosis. The muscular dystrophies ([Chap. 462e](#)) may be additional considerations; however, these disorders usually develop over years rather than weeks or months and rarely present after the age of 30 years. It may be difficult, even with a muscle biopsy, to distinguish chronic PM from a rapidly advancing muscular dystrophy. This is particularly true of facioscapulohumeral muscular dystrophy, dysferlin myopathy, and

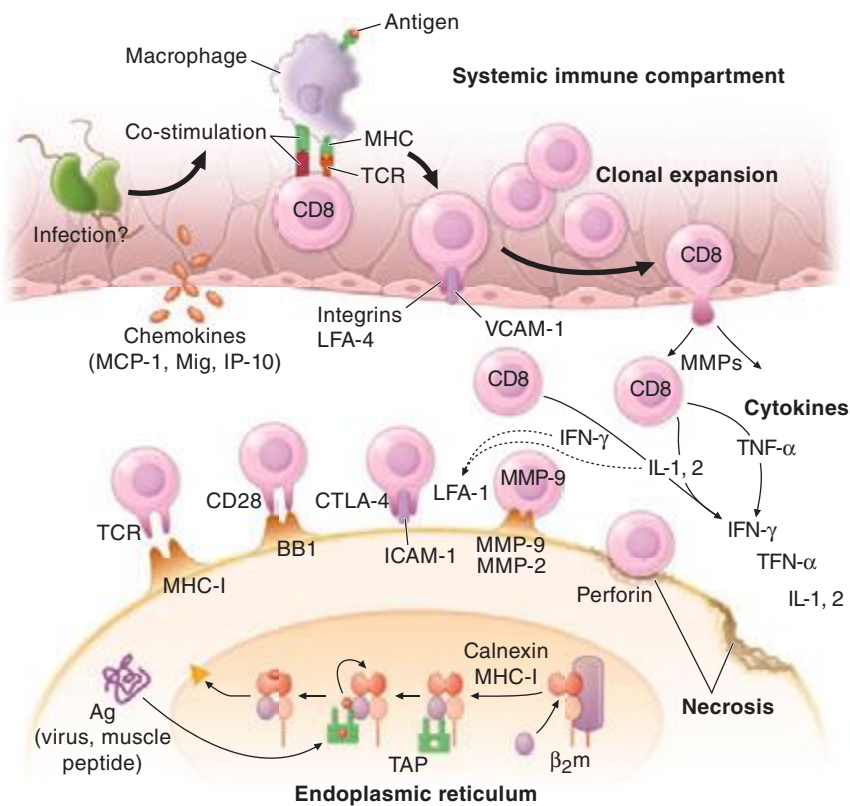


FIGURE 388-2 Cell-mediated mechanisms of muscle damage in polymyositis (PM) and inclusion body myositis (IBM). Antigen-specific CD8 cells are expanded in the periphery, cross the endothelial barrier, and bind directly to muscle fibers via T cell receptor (TCR) molecules that recognize aberrantly expressed major histocompatibility complex (MHC)-I. Engagement of co-stimulatory molecules (BB1 and ICOSL) with their ligands (CD28, CTLA-4, and ICOS), along with ICAM-1/LFA-1, stabilize the CD8–muscle fiber interaction. Metalloproteinases (MMPs) facilitate the migration of T cells and their attachment to the muscle surface. Muscle fiber necrosis occurs via perforin granules released by the autoaggressive T cells. A direct myocytotoxic effect exerted by the cytokines interferon (IFN) γ , interleukin (IL) 1, or tumor necrosis factor (TNF) α may also play a role. Death of the muscle fiber is mediated by necrosis. MHC class I molecules consist of a heavy chain and a light chain (β_2 microglobulin [β_2m]) complexed with an antigenic peptide that is transported into the endoplasmic reticulum by TAP proteins (**Chap. 373e**).

the dystrophinopathies where inflammatory cell infiltration is often found early in the disease. Such doubtful cases should always be given an adequate trial of glucocorticoid therapy and undergo genetic testing to exclude muscular dystrophy. Identification of the MHC/CD8 lesion by muscle biopsy is helpful to identify cases of PM. Some metabolic myopathies, including glycogen storage disease due to myophosphorylase or acid maltase deficiency, lipid storage myopathies due to carnitine deficiency, and mitochondrial diseases produce weakness that is often associated with other characteristic clinical signs; diagnosis rests upon histochemical and biochemical studies of the muscle biopsy. The endocrine myopathies such as those due to hypercortico-steroidism, hyper- and hypothyroidism, and hyper- and hypoparathyroidism require the appropriate laboratory investigations for diagnosis. Muscle wasting in patients with an underlying neoplasm may be due to disuse, cachexia, or rarely a paraneoplastic neuromyopathy (**Chap. 122**).

Diseases of the neuromuscular junction, including myasthenia gravis or the Lambert-Eaton myasthenic syndrome, cause fatiguing weakness that also affects ocular and other cranial muscles (**Chap. 461**). Repetitive nerve stimulation and single-fiber EMG studies aid in diagnosis.

Acute Muscle Weakness This may be caused by an acute neuropathy such as Guillain-Barré syndrome (**Chap. 460**), transverse myelitis (**Chap. 456**), a neurotoxin (**Chap. 462e**), or a neurotropic viral infection such as poliomyelitis or West Nile virus (**Chap. 164**). When acute weakness is associated

with very high levels of serum CK (often in the thousands), painful muscle cramps, rhabdomyolysis, and myoglobinuria, it may be due to a necrotizing autoimmune myositis, as discussed below, a viral infection or a metabolic disorder such as myophosphorylase deficiency, or carnitine palmitoyltransferase deficiency (**Chap. 462e**). Several animal parasites, including protozoa (*Toxoplasma*, *Trypanosoma*), cestodes (cysticerci), and nematodes (trichinae), may produce a focal or diffuse inflammatory myopathy known as *parasitic polymyositis*. *Staphylococcus aureus*, *Yersinia*, *Streptococcus*, or anaerobic bacteria may produce a suppurative myositis, known as *tropical polymyositis*, or *pyomyositis*. Pyomyositis, previously rare in the west, is now occasionally seen in AIDS patients. Other bacteria, such as *Borrelia burgdorferi* (Lyme disease) and *Legionella pneumophila* (Legionnaire's disease), may infrequently cause myositis.

Patients with periodic paralysis experience recurrent episodes of acute muscle weakness without pain, always beginning in childhood. Chronic alcoholics may develop painful myopathy with myoglobinuria after a bout of heavy drinking. Acute painless muscle weakness with myoglobinuria may occur with prolonged hypokalemia, or hypophosphatemia and hypomagnesemia, usually in chronic alcoholics or in patients on nasogastric suction receiving parenteral hyperalimentation.

Myofasciitis This distinctive inflammatory disorder affecting muscle and fascia presents as diffuse myalgias, skin induration, fatigue, and mild muscle weakness; mild elevations of serum CK are usually present. The most common form is eosinophilic myofasciitis characterized by peripheral blood eosinophilia and eosinophilic infiltrates in the endomysial tissue. In some patients, the eosinophilic myositis/fasciitis occurs in the context of parasitic infections, vasculitis, mixed connective tissue disease, hypereosinophilic syndrome, or toxic exposures (e.g., toxic oil syndrome, contaminated L-tryptophan) or with mutations in the calpain gene. A distinct subset of myofasciitis is characterized by pronounced infiltration of the connective tissue around the muscle by sheets of periodic acid–Schiff-positive macrophages and occasional CD8 T cells (macrophagic myofasciitis or inflammatory myositis with abundant macrophages [IMAM]). A focal form of this disorder, limited to sites of previous vaccinations, administered months or years earlier, has been linked to an aluminum-containing substrate in vaccines. This disorder, which to date has not been observed outside of France, responds to glucocorticoid therapy, and the overall prognosis seems favorable.

Necrotizing Autoimmune Myositis This is an increasingly recognized entity that has distinct features, even though it is often labeled as PM. It presents as an acute or subacute onset of symmetric muscle weakness; CK is typically extremely high. The weakness can be severe. Coexisting interstitial lung disease and cardiomyopathy may be present. The disorder may develop after a viral infection, in association with cancer, or in patients taking statins when the myopathy continues to worsen after statin withdrawal. Some patients have antibodies against signal recognition particle (SRP) or against the 3-hydroxy-3-methylglutaryl-coenzyme A reductase (HMGCR), a 100-kDa protein considered the pharmacologic target of statins. The muscle biopsy demonstrates necrotic fibers infiltrated by macrophages but only rare, if any, T cell infiltrates. Muscle MHC-I expression is only slightly and focally upregulated. The capillaries may be swollen with hyalinization, thickening of the capillary wall, and deposition of complement. Most patients respond to immunotherapy, but some are resistant.

Hyperacute Necrotizing Fasciitis/Myositis (Flesh-Eating Disease) This a fulminant infectious disease, seen most often in the tropics or in conditions with poor hygiene, characterized by widespread necrosis of the superficial fascia and muscle of a limb; if the scrotum, perineum, and abdominal wall are affected, the condition is referred to as Fournier's gangrene. It may be caused by group A β -hemolytic *Streptococcus*, methicillin-sensitive *S. aureus*, *Pseudomonas aeruginosa*, *Vibrio vulnificus*, clostridial species (gas gangrene; Chap. 179), or polymicrobial infection with anaerobes and facultative bacteria (Chap. 201); toxins from these bacteria may act as superantigens (Chap. 372e). The port of bacterial entry is usually a trivial cut or skin abrasion, and the source is contact with carriers of the organism. Individuals with diabetes mellitus, immunodeficiency states, or systemic illnesses such as liver failure are most susceptible. Systemic varicella is a predisposing factor in children.

The disease presents with swelling, pain, and redness in the involved area followed by a rapid tissue necrosis of fascia and muscle that progresses at an estimated rate of 3 cm/h. Emergency debridement, antibiotics, IV immunoglobulin (IVIg), and even hyperbaric oxygen have been recommended. In progressive or advanced cases, amputation of the affected limb may be necessary to avoid a fatal outcome.

Drug-Induced Myopathies D-Penicillamine, procainamide, and statins may produce a true myositis resembling PM or necrotizing myositis. A DM-like illness has been associated with the contaminated preparations of L-tryptophan. As noted above, AZT causes a mitochondrial myopathy. Other drugs may elicit a toxic noninflammatory myopathy that is histologically different from DM, PM, or IBM. These include cholesterol-lowering agents such as clofibrate, lovastatin, simvastatin, or pravastatin, especially when combined with cyclosporine, amiodarone, or gemfibrozil. Mild statin-induced myopathic symptoms (such as myalgia, fatigue, or asymptomatic elevations of CK) are self-limited and usually improve after discontinuation of the drug. In rare patients, however, muscle weakness continues to progress even after the statin is withdrawn; in these cases, a diagnostic muscle biopsy is indicated and search for antibodies to HMGCR is suggested; if histologic evidence of PM or necrotizing myositis is present, immunotherapy should be initiated. Rhabdomyolysis and myoglobinuria have been rarely associated with amphotericin B, ϵ -aminocaproic acid, fenfluramine, heroin, and phencyclidine. The use of amiodarone, chloroquine, colchicine, carbimazole, emetine, etretinate, and ipecac syrup; chronic laxative or licorice use resulting in hypokalemia; and glucocorticoid or growth hormone administration have also been associated with myopathic muscle weakness. Some neuromuscular blocking agents such as pancuronium, in combination with glucocorticoids, may cause an acute critical illness myopathy. A careful drug history is essential for diagnosis of these drug-induced myopathies, which do not require immunosuppressive

therapy except when an autoimmune myopathy has been triggered, as noted above.

"Weakness" due to Muscle Pain and Muscle Tenderness A number of conditions including *polymyalgia rheumatica* (Chap. 385) and arthritic disorders of adjacent joints may enter into the differential diagnosis of inflammatory myopathy, even though they do not cause myositis. The muscle biopsy is either normal or discloses type II muscle fiber atrophy. Patients with *fibrositis* and *fibromyalgia* (Chap. 396) complain of focal or diffuse muscle tenderness, fatigue, and aching, which is sometimes poorly differentiated from joint pain. Some patients, however, have muscle tenderness, painful muscles on movement, and signs suggestive of a collagen vascular disorder, such as an increased erythrocyte sedimentation rate, C-reactive protein, antinuclear antibody, or rheumatoid factor, along with modest elevation of the serum CK and aldolase. They demonstrate a "break-away" pattern of weakness with difficulty sustaining effort but not true muscle weakness. The muscle biopsy is usually normal or nonspecific. Many such patients show some response to nonsteroidal anti-inflammatory agents or glucocorticoids, although most continue to have indolent complaints. An indolent fasciitis in the setting of an ill-defined connective tissue disorder may be at times present, and these patients should not be labeled as having a psychosomatic disorder. *Chronic fatigue syndrome*, which may follow a viral infection, can present with debilitating fatigue, sore throat, painful lymphadenopathy, myalgia, arthralgia, sleep disorder, and headache (Chap. 464e). These patients do not have muscle weakness, and the muscle biopsy is normal.

DIAGNOSIS

The clinically suspected diagnosis of PM, DM, IBM, or necrotizing myositis is confirmed by analysis of serum muscle enzymes, EMG findings, and muscle biopsy (Table 388-2).

The most sensitive enzyme is CK, which in active disease can be elevated as much as 50-fold. Although the CK level usually parallels disease activity, it can be normal in some patients with active IBM or DM, especially when associated with a connective tissue disease. The CK is always elevated in patients with active PM. Along with the CK, the serum glutamic-oxaloacetic and glutamate pyruvate transaminases, lactate dehydrogenase, and aldolase may be elevated.

Needle EMG shows myopathic potentials characterized by short-duration, low-amplitude polyphasic units on voluntary activation and increased spontaneous activity with fibrillations, complex repetitive discharges, and positive sharp waves. Mixed potentials (polyphasic units of short and long duration) indicating a chronic process and muscle fiber regeneration are often present in IBM. These EMG findings are not diagnostic of an inflammatory myopathy but are useful

TABLE 388-2 CRITERIA FOR DIAGNOSIS OF INFLAMMATORY MYOPATHIES

Criterion	Polymyositis		Dermatomyositis	Inclusion Body Myositis
	Definite	Probable		
Myopathic muscle weakness ^a	Yes	Yes	Yes ^b	Yes; slow onset, early involvement of distal muscles, frequent falls
Electromyographic findings	Myopathic	Myopathic	Myopathic	Myopathic with mixed potentials
Muscle enzymes	Elevated (up to 50-fold)	Elevated (up to 50-fold)	Elevated (up to 50-fold) or normal	Elevated (up to 10-fold) or normal
Muscle biopsy findings ^c	"Primary" inflammation with the CD8/MHC-I complex and no vacuoles	Ubiquitous MHC-I expression but minimal inflammation and no vacuoles ^d	Perifascicular, perimysial, or perivascular infiltrates, perifascicular atrophy	Primary inflammation with CD8/MHC-I complex; vacuolated fibers with β -amyloid deposits; cytochrome oxygenase-negative fibers; signs of chronic myopathy ^e
Rash or calcinosis	Absent	Absent	Present ^f	Absent

^aMyopathic muscle weakness, affecting proximal muscles more than distal ones and sparing eye and facial muscles, is characterized by a subacute onset (weeks to months) and rapid progression in patients who have no family history of neuromuscular disease, no endocrinopathy, no exposure to myotoxic drugs or toxins, and no biochemical muscle disease (excluded on the basis of muscle biopsy findings). ^bIn some cases with the typical rash, the muscle strength is seemingly normal (dermatomyositis sine myositis); these patients often have new onset of easy fatigue and reduced endurance. Careful muscle testing may reveal mild muscle weakness. ^cSee text for details. ^dAn adequate trial of prednisone or other immunosuppressive drugs is warranted in probable cases. If, in retrospect, the disease is unresponsive to therapy, another muscle biopsy should be considered to exclude other diseases or possible evolution in inclusion body myositis. ^eIf the muscle biopsy does not contain vacuolated fibers but shows chronic myopathy with hypertrophic fibers, primary inflammation with the CD8/MHC-I complex, and cytochrome oxygenase-negative fibers, the diagnosis is probable inclusion body myositis. ^fIf rash is absent but muscle biopsy findings are characteristic of dermatomyositis, the diagnosis is probable dermatomyositis.

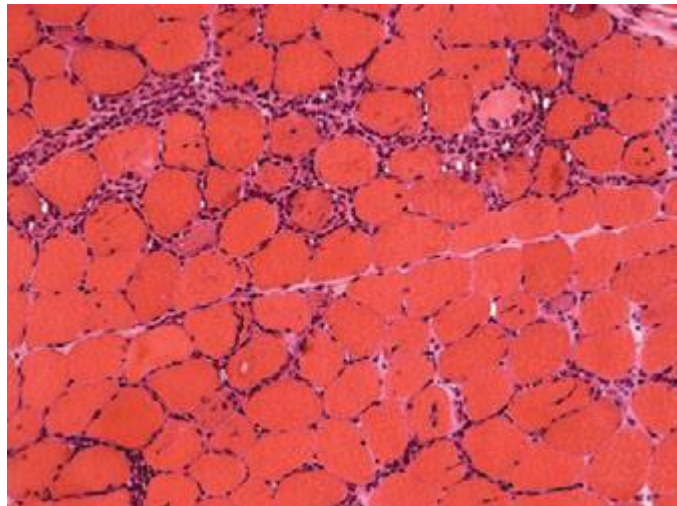


FIGURE 388-3 Cross-section of a muscle biopsy from a patient with polymyositis demonstrates scattered inflammatory foci with lymphocytes invading or surrounding muscle fibers. Note lack of chronic myopathic features (increased connective tissue, atrophic or hypertrophic fibers) as seen in inclusion body myositis.

to identify the presence of active or chronic myopathy and to exclude neurogenic disorders.

Magnetic resonance imaging (MRI) is not routinely used for the diagnosis of PM, DM, or IBM. However, it may provide information or guide the location of the muscle biopsy in certain clinical settings.

Muscle biopsy—despite occasional variability in demonstrating all of the typical pathologic findings—is the most sensitive and specific test for establishing the diagnosis of inflammatory myopathy and for excluding other neuromuscular diseases. Inflammation is the histologic hallmark for these diseases; however, additional features are characteristic of each subtype (Figs. 388-3, 388-4, and 388-5).

In PM the inflammation is *primary*, a term used to indicate that the inflammation is not reactive and the T cell infiltrates, located primarily within the muscle fascicles (endomysially), surround individual, healthy muscle fibers and result in phagocytosis and necrosis (Fig. 388-3). The MHC-I molecule is ubiquitously expressed on the sarcolemma, even in fibers not invaded by CD8+ cells. The CD8/MHC-I lesion is characteristic and useful to confirm or establish the diagnosis and to exclude disorders with secondary, nonspecific, inflammation, such as in some muscular dystrophies. When the disease is chronic, connective tissue is increased and may react positively with alkaline phosphatase. In necrotizing myositis, there are abundant necrotic fibers invaded or

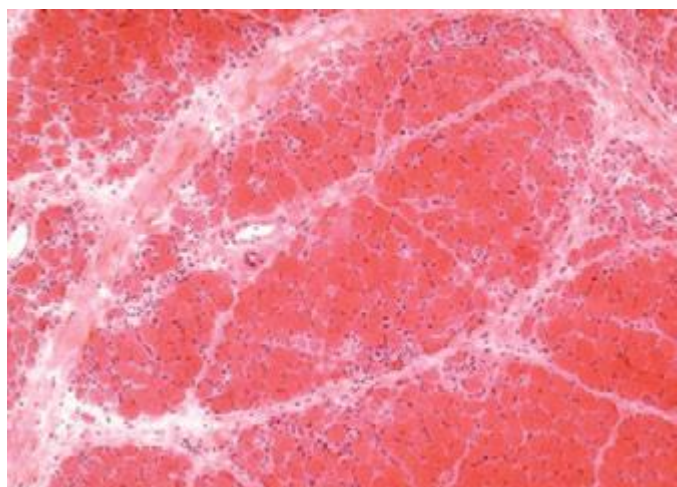


FIGURE 388-4 Cross-section of a muscle biopsy from a patient with dermatomyositis demonstrates atrophy of the fibers at the periphery of the fascicle (perifascicular atrophy).

surrounded by macrophages, but no lymphocytic infiltrates or MHC-I expression beyond the necrotic fibers.

In DM the endomysial inflammation is predominantly perivascular or in the interfascicular septae and around—rather than within—the muscle fascicles (Fig. 388-4). The intramuscular blood vessels show endothelial hyperplasia with tubuloreticular profiles, fibrin thrombi, and obliteration of capillaries. The muscle fibers undergo necrosis, degeneration, and phagocytosis, often in groups involving a portion of a muscle fasciculus in a wedge-like shape or at the periphery of the fascicle, due to microinfarcts within the muscle. This results in perifascicular atrophy, characterized by 2–10 layers of atrophic fibers at the periphery of the fascicles. The presence of perifascicular atrophy is diagnostic of DM, *even in the absence of inflammation*.

In IBM (Fig. 388-5), there is endomysial inflammation with T cells invading MHC-I-expressing nonvacuolated muscle fibers; basophilic granular deposits distributed around the edge of slit-like vacuoles (rimmed vacuoles); loss of fibers, replaced by fat and connective tissue, hypertrophic fibers, and angulated or round fibers; rare eosinophilic cytoplasmic inclusions; abnormal mitochondria characterized by the presence of ragged-red fibers or cytochrome oxidase–negative fibers; and amyloid deposits within or next to the vacuoles best visualized with crystal violet or Congo-red staining viewed with fluorescent optics. Electron microscopy demonstrates filamentous inclusions in the vicinity of the rimmed vacuoles. In at least 15% of patients with the typical clinical phenotype of IBM, there is brisk inflammation in the muscle biopsy but no vacuoles or amyloid deposits, leading to an erroneous diagnosis of PM. Such patients are often referred to as having “clinical IBM.” Close clinicopathologic correlations are therefore essential; if uncertain, a repeat muscle biopsy from another site is often helpful.

TREATMENT THERAPY OF INFLAMMATORY MYOPATHIES

The goal of therapy is to improve muscle strength, thereby improving function in activities of daily living, and ameliorate the extramuscular manifestations (rash, dysphagia, dyspnea, fever). When strength improves, the serum CK falls concurrently; however, the reverse is not always true. Unfortunately, there is a common tendency to “chase” or treat the CK level instead of the muscle weakness, a practice that has led to prolonged and unnecessary use of immunosuppressive drugs and erroneous assessment of their efficacy. It is prudent to discontinue these drugs if, after an adequate trial, there is no objective improvement in muscle strength whether or not CK levels are reduced. Agents used in the treatment of PM and DM include the following:

1. **Glucocorticoids.** Oral prednisone is the initial treatment of choice; the effectiveness and side effects of this therapy determine the future need for stronger immunosuppressive drugs. High-dose prednisone, at least 1 mg/kg per day, is initiated as early in the disease as possible. After 3–4 weeks, prednisone is tapered slowly over a period of 10 weeks to 1 mg/kg every other day. If there is evidence of efficacy and no serious side effects, the dosage is then further reduced by 5 or 10 mg every 3–4 weeks until the lowest possible dose that controls the disease is reached. The efficacy of prednisone is determined by an objective increase in muscle strength and activities of daily living, which almost always occurs by the third month of therapy. A feeling of increased energy or a reduction of the CK level without a concomitant increase in muscle strength is not a reliable sign of improvement. If prednisone provides no objective benefit after ~3 months of high-dose therapy, the disease is probably unresponsive to the drug and tapering should be accelerated while the next-in-line immunosuppressive drug is started. Although controlled trials have not been performed, almost all patients with true PM or DM respond to glucocorticoids to *some degree and for some period of time*; in general, DM responds better than PM.

The long-term use of prednisone may cause increased weakness associated with a normal or unchanged CK level; this effect is referred to as *steroid myopathy*. In a patient who previously

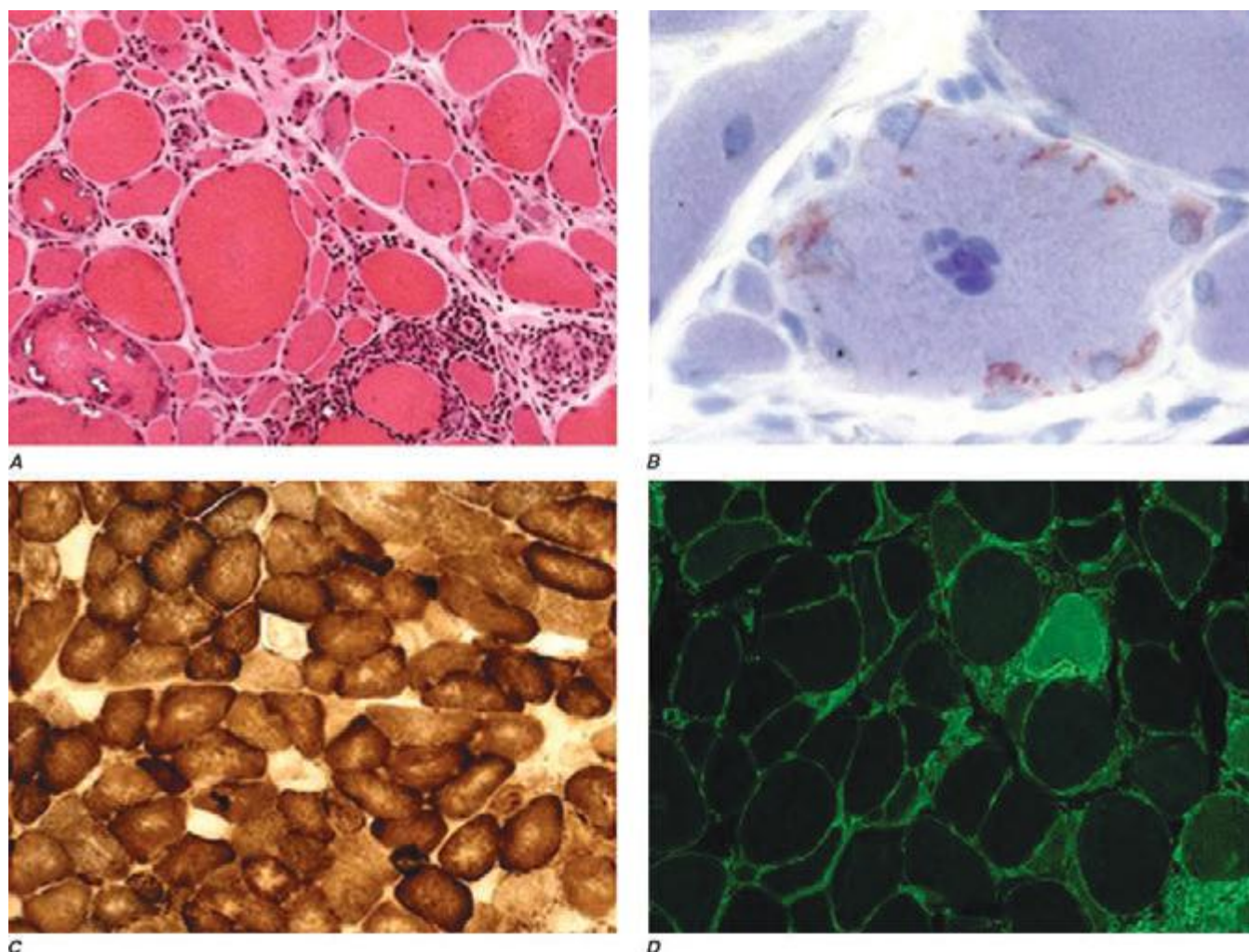


FIGURE 388-5 Cross-sections of a muscle biopsy from a patient with inclusion body myositis demonstrate the typical features of vacuoles with lymphocytic infiltrates surrounding nonvacuolated or necrotic fibers (A), tiny endomysial deposits of amyloid visualized with crystal violet (B), cytochrome oxidase–negative fibers, indicative of mitochondrial dysfunction (C), and ubiquitous major histocompatibility complex class I expression at the periphery of all fibers (D).

responded to high doses of prednisone, the development of new weakness may be related to steroid myopathy or to disease activity that either will respond to a higher dose of glucocorticoids or has become glucocorticoid-resistant. In uncertain cases, the prednisone dosage can be steadily increased or decreased as desired; the cause of the weakness is usually evident in 2–8 weeks.

2. *Other immunosuppressive drugs.* Approximately 75% of patients ultimately require additional treatment. This occurs when a patient fails to respond adequately to glucocorticoids after a 3-month trial, the patient becomes glucocorticoid-resistant, glucocorticoid-related side effects appear, attempts to lower the prednisone dose repeatedly result in a new relapse, or rapidly progressive disease with evolving severe weakness and respiratory failure develops.

The following drugs are commonly used but have never been tested in controlled studies: (1) *Azathioprine* is well tolerated, has few side effects, and appears to be as effective for long-term therapy as other drugs. The dose is up to 3 mg/kg daily. (2) *Methotrexate* has a faster onset of action than azathioprine. It is given orally starting at 7.5 mg weekly for the first 3 weeks (2.5 mg every 12 h for 3 doses), with gradual dose escalation by 2.5 mg per week to a total of 25 mg weekly. A rare side effect is methotrexate pneumonitis, which can be difficult to distinguish from the interstitial lung disease of the primary myopathy associated with Jo-1 antibodies (described above). (3) *Mycophenolate mofetil* also has a faster onset of action than azathioprine. At doses up to 2.5 or 3 g/d in two divided doses, it is well tolerated for long-term

use. (4) Monoclonal anti-CD20 antibody (rituximab) has been shown in a small uncontrolled series to benefit patients with DM and PM, but a controlled study did not show differences between patients randomized 8 weeks apart. (5) *Cyclosporine* has inconsistent and mild benefit. (6) *Cyclophosphamide* (0.5–1 g/m² IV monthly for 6 months) has limited success and significant toxicity. (7) Tacrolimus (formerly known as Fk506) has been effective in some difficult cases of PM especially with interstitial lung disease.

3. *Immunomodulation.* In a controlled trial of patients with refractory DM, IVIg improved not only strength and rash but also the underlying immunopathology. The benefit is often short-lived (≤ 8 weeks), and repeated infusions every 6–8 weeks are generally required to maintain improvement. A dose of 2 g/kg divided over 2–5 days per course is recommended. Uncontrolled observations suggest that IVIg may also be beneficial for patients with PM. Neither plasmapheresis nor leukapheresis appears to be effective in PM and DM.

The following sequential empirical approach to the treatment of PM and DM is suggested: *Step 1:* high-dose prednisone; *Step 2:* azathioprine, mycophenolate, or methotrexate for steroid-sparing effect; *Step 3:* IVIg; *Step 4:* a trial, with guarded optimism, of one of the following agents, chosen according to the patient's age, degree of disability, tolerance, experience with the drug, and general health: rituximab, cyclosporine, cyclophosphamide, or tacrolimus. Patients with interstitial lung disease may benefit from aggressive treatment with cyclophosphamide or tacrolimus.

A patient with presumed PM who has not responded to any form of immunotherapy most likely has IBM or another disease, usually a metabolic myopathy, a muscular dystrophy, a drug-induced myopathy, or an endocrinopathy. In these cases, a repeat muscle biopsy and a renewed search for another cause of the myopathy is indicated.

Calcinosis, a manifestation of DM, is difficult to treat; however, new calcium deposits may be prevented if the primary disease responds to the available therapies. Bisphosphonates, aluminum hydroxide, probenecid, colchicine, low doses of warfarin, calcium blockers, and surgical excision have all been tried without success.

IBM is generally resistant to immunosuppressive therapies. Prednisone together with azathioprine or methotrexate is often tried for a few months in newly diagnosed patients, although results are generally disappointing. Because occasional patients may feel subjectively weaker after these drugs are discontinued, some clinicians prefer to maintain these patients on low-dose, every-other-day prednisone along with mycophenolate in an effort to slow disease progression, even though there is no objective evidence or controlled study to support this practice. In two controlled studies of IVIg in IBM, minimal benefit in up to 30% of patients was found; the strength gains, however, were not of sufficient magnitude to justify its routine use. Another trial of IVIg combined with prednisone was ineffective. Nonetheless, some experts believe that a 2- to 3-month trial with IVIg may be reasonable for selected patients with IBM who experience rapid progression of muscle weakness or choking episodes due to worsening dysphagia.

PROGNOSIS

The 5-year survival rate for treated patients with PM and DM is ~95%, and the 10-year survival rate is 84%; death is usually due to pulmonary, cardiac, or other systemic complications. The prognosis is worse for patients who are severely affected at presentation, when initial treatment is delayed, and in cases with severe dysphagia or respiratory difficulties. Older patients and those with associated cancer also have a worse prognosis. DM responds more favorably to therapy than PM and thus has a better prognosis. Most patients improve with therapy, and many make a full functional recovery, which is often sustained with maintenance therapy. Up to 30% may be left with some residual muscle weakness. Relapses may occur at any time.

IBM has the least favorable prognosis of the inflammatory myopathies. Most patients will require the use of an assistive device such as a cane, walker, or wheelchair within 5–10 years of onset. In general, the older the age of onset in IBM, the more rapidly progressive is the course.

389 Relapsing Polychondritis

Carol A. Langford

Relapsing polychondritis is an uncommon disorder of unknown cause characterized by inflammation of cartilage predominantly affecting the ears, nose, and laryngotracheobronchial tree. Other manifestations include scleritis, neurosensory hearing loss, polyarthritis, cardiac abnormalities, skin lesions, and glomerulonephritis. Relapsing polychondritis has been estimated to have an incidence of 3.5 per million population per year. The peak age of onset is between the ages of 40 and 50 years, but relapsing polychondritis may affect children and the elderly. It is found in all races, and both sexes are equally affected. No familial tendency is apparent. A significantly higher frequency of HLA-DR4 has been found in patients with relapsing polychondritis than in healthy individuals. A predominant subtype allele(s) of HLA-DR4 was not found. Approximately 30% of patients with relapsing polychondritis will have another rheumatologic disorder, the most frequent being systemic vasculitis, followed by rheumatoid arthritis, and systemic lupus erythematosus (SLE). Nonrheumatic

TABLE 389-1 DISORDERS ASSOCIATED WITH RELAPSING POLYCHONDRITIS^a

Systemic vasculitis
Rheumatoid arthritis
Systemic lupus erythematosus
Overlapping connective tissue disease
Spondyloarthritis
Behçet's disease
Polymyalgia rheumatica
Primary biliary cirrhosis
Pulmonary fibrosis
Hashimoto's thyroiditis
Myelodysplastic syndrome

^aSystemic vasculitis is the most common association, followed by rheumatoid arthritis and systemic lupus erythematosus.

Source: Modified from CJ Michet et al: *Ann Intern Med* 104:74, 1986.

disorders associated with relapsing polychondritis include Hashimoto's thyroiditis, primary biliary cirrhosis, and myelodysplastic syndrome (Table 389-1). In most cases, these disorders antedate the appearance of relapsing polychondritis, usually by months or years; however, in other instances, the onset of relapsing polychondritis can accompany disease presentation.

PATHOLOGY AND PATHOPHYSIOLOGY

The earliest abnormality of hyaline and elastic cartilage noted histologically is a focal or diffuse loss of basophilic staining indicating depletion of proteoglycan from the cartilage matrix. Inflammatory infiltrates are found adjacent to involved cartilage and consist predominantly of mononuclear cells and occasional plasma cells. In acute disease, polymorphonuclear white cells may also be present. Destruction of cartilage begins at the outer edges and advances centrally. There is lacunar breakdown and loss of chondrocytes. Degenerating cartilage is replaced by granulation tissue and later by fibrosis and focal areas of calcification. Small loci of cartilage regeneration may be present. Immunofluorescence studies have shown immunoglobulins and complement at sites of involvement. Extracellular granular material observed in the degenerating cartilage matrix by electron microscopy has been interpreted to be enzymes, immunoglobulins, or proteoglycans.

Immunologic mechanisms play a role in the pathogenesis of relapsing polychondritis. The accumulating data strongly suggest that both humoral and cell-mediated immunity play an important role in the pathogenesis of relapsing polychondritis. Immunoglobulin and complement deposits are found at sites of inflammation. In addition, antibodies to type II collagen and to matrilin-1 and immune complexes are detected in the sera of some patients. The possibility that an immune response to type II collagen may be important in the pathogenesis is supported experimentally by the occurrence of auricular chondritis in rats immunized with type II collagen. Antibodies to type II collagen are found in the sera of these animals, and immune deposits are detected at sites of ear inflammation. Humoral immune responses to type IX and type XI collagen, matrilin-1, and cartilage oligomeric matrix protein have been demonstrated in some patients. In a study, rats immunized with matrilin-1 were found to develop severe inspiratory stridor and swelling of the nasal septum. The rats had severe inflammation with erosions of the involved cartilage, which was characterized by increased numbers of CD4+ and CD8+ T cells in the lesions. The cartilage of the joints and ear pinna was not involved. All had IgG antibodies to matrilin-1. Matrilin-1 is a noncollagenous protein present in the extracellular matrix in cartilage. It is present in high concentrations in the trachea and is also present in the nasal septum but not in articular cartilage. A subsequent study demonstrated serum anti-matrilin-1 antibodies in approximately 13% of patients with relapsing polychondritis; approximately 70% of these patients had respiratory symptoms. Cell-mediated immunity may also be operative in causing tissue injury, since lymphocyte transformation can be demonstrated when lymphocytes of patients are exposed to cartilage extracts. T cells specific for type II collagen have been found in some

TABLE 389-2 CLINICAL MANIFESTATIONS OF RELAPSING POLYCHONDritis

Clinical Feature	Presenting	Cumulative
		Frequency, %
Auricular chondritis	43	89
Arthritis	32	72
Nasal chondritis	21	61
Ocular inflammation	18	59
Laryngotracheal symptoms	23	55
Reduced hearing	7	40
Saddle nose deformity	11	25
Cutaneous	4	25
Laryngotracheal stricture	15	23
Vasculitis	2	14
Elevated creatinine	7	13
Aortic or mitral regurgitation	0	12

Source: Modified from PD Kent et al: *Curr Opin Rheumatol* 16:56, 2004.

patients, and CD4+ T cells have been observed at sites of cartilage inflammation.

CLINICAL MANIFESTATIONS

The onset of relapsing polychondritis is frequently abrupt, with the appearance of one or two sites of cartilaginous inflammation. The pattern of cartilaginous involvement and the frequency of episodes vary widely among patients. Noncartilaginous presentations may also occur. Systemic inflammatory features such as fever, fatigue, and weight loss occur and may precede the clinical signs of relapsing polychondritis by several weeks. Relapsing polychondritis may go unrecognized for several months or even years in patients who only initially manifest intermittent joint pain and/or swelling, or who have unexplained eye inflammation, hearing loss, valvular heart disease, or pulmonary symptoms.

Auricular chondritis is the most frequent presenting manifestation of relapsing polychondritis, occurring in 40% of patients and eventually affecting about 85% of patients (Table 389-2). One or both ears are involved, either sequentially or simultaneously. Patients experience the sudden onset of pain, tenderness, and swelling of the cartilaginous portion of the ear (Fig. 389-1). This typically involves the pinna of the ears, sparing the earlobes because they do not contain cartilage. The overlying skin has a beefy red or violaceous color. Prolonged or recurrent episodes lead to cartilage destruction and result in a flabby or



FIGURE 389-1 *Left.* The pinna is erythematous, swollen, and tender. Not shown is the ear lobule that is spared as there is no underlying cartilage. *Right.* The pinna is thickened and deformed. The destruction of the underlying cartilage results in a floppy ear. (Reprinted from the *Clinical Slide Collection on the Rheumatic Diseases*, ©1991, 1995, 1997, 1998, 1999. Used by permission of the American College of Rheumatology.)



FIGURE 389-2 Saddle nose results from destruction and collapse of the nasal cartilage. (Reprinted from the *Clinical Slide Collection on the Rheumatic Diseases*, ©1991, 1995, 1997, 1998, 1999. Used by permission of the American College of Rheumatology.)

droopy ear. Swelling may close off the eustachian tube or the external auditory meatus, either of which can impair hearing. Inflammation of the internal auditory artery or its cochlear branch produces hearing loss, vertigo, ataxia, nausea, and vomiting. Vertigo is almost always accompanied by hearing loss.

Approximately 61% of patients will develop nasal involvement, with 21% having this at the time of presentation. Patients may experience nasal stuffiness, rhinorrhea, and epistaxis. The bridge of the nose and surrounding tissue become red, swollen, and tender and may collapse, producing a saddle nose deformity (Fig. 389-2). In some patients, nasal deformity develops insidiously without overt inflammation. Saddle nose is observed more frequently in younger patients, especially in women.

Joint involvement is the presenting manifestation in relapsing polychondritis in approximately one-third of patients and may be present for several months before other features appear. Eventually, more than one-half of the patients will have arthralgias or arthritis. The arthritis is usually asymmetric and oligo- or polyarticular, and it involves both large and small peripheral joints. An episode of arthritis lasts from a few days to several weeks and resolves spontaneously without joint erosion or deformity. Attacks of arthritis may not be temporally related to other manifestations of relapsing polychondritis. Joint fluid has been reported to be noninflammatory. In addition to peripheral joints, inflammation may involve the costochondral, sternomanubrial, and sternoclavicular cartilages. Destruction of these cartilages may result in a pectus excavatum deformity or even a flail anterior chest wall.

Eye manifestations occur in more than one-half of patients and include conjunctivitis, episcleritis, scleritis, iritis, uveitis, and keratitis. Ocular inflammation can be severe and visually threatening. Other manifestations include eyelid and periorbital edema, proptosis, optic neuritis, extraocular muscle palsies, retinal vasculitis, and renal vein occlusion.

Laryngotracheobronchial involvement occurs in ~50% of patients and is among the most serious manifestations of relapsing polychondritis. Symptoms include hoarseness, a nonproductive cough, and tenderness over the larynx and proximal trachea. Mucosal edema, strictures, and/or collapse of laryngeal or tracheal cartilage may cause stridor and life-threatening airway obstruction necessitating tracheostomy. Involvement can extend into the lower airways resulting in tracheobronchomalacia. Collapse of cartilage in bronchi leads to pneumonia and, when extensive, to respiratory insufficiency.

Cardiac valvular regurgitation occurs in about 5–10% of patients and is due to progressive dilation of the valvular ring or to destruction

2204 of the valve cusps. Aortic regurgitation occurs in about 7% of patients, with the mitral and other heart valves being affected less often. Other cardiac manifestations include pericarditis, myocarditis, coronary vasculitis, and conduction abnormalities. Aneurysms of the proximal, thoracic, or abdominal aorta may occur even in the absence of active chondritis and occasionally rupture.

Renal disease occurs in about 10% of patients. The most common renal lesions include mesangial expansion or segmental necrotizing glomerulonephritis, which have been reported to have small amounts of electron-dense deposits in the mesangium where there is also faint deposition of C3 and/or IgG or IgM. Tubulointerstitial disease and IgA nephropathy have also been reported.

Approximately 25% of patients have skin lesions, which can include purpura, erythema nodosum, erythema multiforme, angioedema/urticaria, livedo reticularis, and panniculitis.

Features of vasculitis are seen in up to 25% of patients and can affect any size vessel. Large vessel vasculitis may present with aortic aneurysms, and medium vessel disease may affect the coronary, hepatic, mesenteric, or renal arteries or vessel supplying nerves. Skin vessel disease and involvement of the postcapillary venules can also occur. A variety of primary vasculitides have also been reported to occur in association with relapsing polychondritis (**Chap. 385**). One specific overlap is the “MAGIC” syndrome (mouth and genital ulcers with inflamed cartilage) in which patients present with features of both relapsing polychondritis and Behçet’s disease (**Chap. 387**).

LABORATORY FINDINGS AND DIAGNOSTIC IMAGING

There are no laboratory features that are diagnostic for relapsing polychondritis. Mild leukocytosis and normocytic, normochromic anemia are often present. Eosinophilia is observed in 10% of patients. The erythrocyte sedimentation rate and C-reactive protein are usually elevated. Rheumatoid factor and antinuclear antibody tests are occasionally positive in low titers, and complement levels are normal. Antibodies to type II collagen are present in fewer than one-half of the patients and are not specific. Circulating immune complexes may be detected, especially in patients with early active disease. Elevated levels of γ globulin may be present. Antineutrophil cytoplasmic antibodies (ANCA), either cytoplasmic (cANCA) or perinuclear (pANCA), are found in some patients with active disease. However, on target antigen-specific testing, there are only occasional reports of positive myeloperoxidase-ANCA, and proteinase 3-ANCA are very rarely found in relapsing polychondritis.

The upper and lower airways can be evaluated by imaging techniques such as computed tomography and magnetic resonance imaging (MRI). Bronchoscopy provides direct visualization of the airways but can be a high-risk procedure in patients with airway compromise. Pulmonary function testing with flow-volume loops can show inspiratory and/or expiratory obstruction. Imaging can also be useful to detect extracartilaginous disease. The chest film may show widening of the ascending or descending aorta due to an aneurysm, and cardiomegaly when aortic insufficiency is present. MRI can assess aortic aneurysmal dilatation. Electrocardiography and echocardiography can be useful in further evaluating for cardiac features of disease.

DIAGNOSIS

Diagnosis is based on recognition of the typical clinical features. Biopsies of the involved cartilage from the ear, nose, or respiratory tract will confirm the diagnosis but are only necessary when clinical features are not typical. Diagnostic criteria were suggested in 1976 by McAdam et al and modified by Damiani and Levine in 1979. These criteria continue to be generally used in clinical practice. McAdam et al proposed the following: (1) recurrent chondritis of both auricles; (2) nonerosive inflammatory arthritis; (3) chondritis of nasal cartilage; (4) inflammation of ocular structures, including conjunctivitis, keratitis, scleritis/episcleritis, and/or uveitis; (5) chondritis of the laryngeal and/or tracheal cartilages; and (6) cochlear and/or vestibular damage manifested by neurosensory hearing loss, tinnitus, and/or vertigo. The diagnosis is certain when three or more of these features are present along with a positive biopsy from the ear, nasal, or respiratory cartilage. Damiani and Levine later suggested that the diagnosis could be

made when one or more of the above features and a positive biopsy were present, when two or more separate sites of cartilage inflammation were present that responded to glucocorticoids or dapson, or when three or more of the above features were present.

The differential diagnosis of relapsing polychondritis is centered around its sites of clinical involvement. Patients with granulomatosis with polyangiitis (Wegener’s) may have a saddle nose and tracheal involvement but can be distinguished by the primary inflammation occurring in the mucosa at these sites, the absence of auricular involvement, and the presence of pulmonary parenchymal disease. Patients with Cogan’s syndrome have interstitial keratitis and vestibular and auditory abnormalities, but this syndrome does not involve the respiratory tract or ears. Reactive arthritis may initially resemble relapsing polychondritis because of oligoarticular arthritis and eye involvement, but it is distinguished in time by the appearance of urethritis and typical mucocutaneous lesions and the absence of nose or ear cartilage involvement. Rheumatoid arthritis may initially suggest relapsing polychondritis because of arthritis and eye inflammation. The arthritis in rheumatoid arthritis, however, is erosive and symmetric. In addition, rheumatoid factor titers are usually high compared with those in relapsing polychondritis, and anti-cyclic citrullinated peptide is usually not seen. Bacterial infection of the pinna may be mistaken for relapsing polychondritis but differs by usually involving only one ear, including the earlobe. Auricular cartilage may also be damaged by trauma or frostbite. Nasal destructive disease and auricular abnormalities can also be seen in patients using cocaine adulterated with levamisole. Ear involvement in this setting differs from relapsing polychondritis by typically manifesting as purpuric plaques with necrosis extending to the pinna, which does not contain cartilage.

TREATMENT RELAPSING POLYCHONDritis

In patients with active chondritis, prednisone, 40–60 mg/d, is often effective in suppressing disease activity; it is tapered gradually once disease is controlled. In some patients, prednisone can be stopped, whereas in others, low doses in the range of 5–10 mg/d are required for continued suppression of disease. Dapsone 50–100 mg/d has been effective for cartilage inflammation and joint features in some patients. Other immunosuppressive drugs such as cyclophosphamide, methotrexate, azathioprine, or cyclosporine should be reserved for patients who have severe organ-threatening disease, fail to respond to prednisone, or require high doses to control disease activity. Patients with significant ocular inflammation often require intraocular glucocorticoids as well as high doses of prednisone. There are a small number of reports on the use of tumor necrosis factor antagonists, rituximab (anti-CD20), and tocilizumab (anti-interleukin 6 receptor), which are too few in number to assess efficacy. Heart valve replacement or repair of an aortic aneurysm may be necessary. When airway obstruction is severe, tracheostomy is required. Stents may be necessary in patients with tracheobronchial collapse.

PATIENT OUTCOME, PROGNOSIS, AND SURVIVAL

The course of relapsing polychondritis is highly variable. Some patients experience inflammatory episodes lasting from a few days to several weeks that then subside spontaneously or with treatment. Attacks may recur at intervals varying from weeks to months. In other patients, the disease has a chronic, smoldering course that may be severe. In one study, the 5-year estimated survival rate was 74% and the 10-year survival rate was 55%. In contrast to earlier series, only about one-half of the deaths could be attributed to relapsing polychondritis or complications of treatment. Airway complications accounted for only 10% of all fatalities. In general, patients with more widespread disease have a worse prognosis.

ACKNOWLEDGMENT

This chapter represents a revised version of the text authored by Dr. Bruce C. Gilliland that appeared in previous editions of Harrison’s Principles of Internal Medicine. Dr. Gilliland passed away on February 17, 2007, and had been a contributor to Harrison’s since the 11th edition.

DEFINITION

Sarcoidosis is an inflammatory disease characterized by the presence of noncaseating granulomas. The disease is often multisystem and requires the presence of involvement in two or more organs for a specific diagnosis. The finding of granulomas is not specific for sarcoidosis, and other conditions known to cause granulomas must be ruled out. These conditions include mycobacterial and fungal infections, malignancy, and environmental agents such as beryllium. Although sarcoidosis can affect virtually every organ of the body, the lung is most commonly affected. Other organs commonly affected are the liver, skin, and eye. The clinical outcome of sarcoidosis varies, with remission occurring in over one-half of patients within a few years of diagnosis; however, the remaining patients may develop a chronic disease that lasts for decades.

ETIOLOGY

Despite multiple investigations, the cause of sarcoidosis remains unknown. Currently, the most likely etiology is an infectious or non-infectious environmental agent that triggers an inflammatory response in a genetically susceptible host. Among the possible infectious agents, careful studies have shown a much higher incidence of *Propionibacter acnes* in the lymph nodes of sarcoidosis patients compared to controls. An animal model has shown that *P. acnes* can induce a granulomatous response in mice similar to sarcoidosis. Others have demonstrated the presence of a mycobacterial protein (*Mycobacterium tuberculosis* catalase-peroxidase [mKatG]) in the granulomas of some sarcoidosis patients. This protein is very resistant to degradation and may represent the persistent antigen in sarcoidosis. Immune response to this and other mycobacterial proteins has been documented by another laboratory. These studies suggest that a mycobacterium similar to *M. tuberculosis* could be responsible for sarcoidosis. The mechanism exposure/infection with such agents has been the focus of other studies. Environmental exposures to insecticides and mold have been associated with an increased risk for disease. In addition, health care workers appear to have an increased risk. Also, sarcoidosis in a donor organ has occurred after transplantation into a sarcoidosis patient. Some authors have suggested that sarcoidosis is not due to a single agent but represents a particular host response to multiple agents. Some studies have been able to correlate the environmental exposures to genetic markers. These studies have supported the hypothesis that a genetically susceptible host is a key factor in the disease.

INCIDENCE, PREVALENCE, AND GLOBAL IMPACT

Sarcoidosis is seen worldwide, with the highest prevalence reported in the Nordic population. In the United States, the disease has been reported more commonly in African Americans than whites, with the ratio of African Americans to whites ranging from 3:1 to 17:0. Women appear to be slightly more susceptible than men. The higher incidence in African Americans may have been influenced by the fact that African Americans seem to develop more extensive and chronic pulmonary disease. Because most sarcoidosis clinics are run by pulmonologists, a selection bias may have occurred. Worldwide, the prevalence of the disease varies from 20–60 per 100,000 for many groups such as Japanese, Italians, and American whites. Higher rates occur in Ireland and Nordic countries. In one closely observed community in Sweden, the lifetime risk for developing sarcoidosis was 3%.

Sarcoidosis often occurs in young, otherwise healthy adults. It is uncommon to diagnose the disease in someone under age 18. However, it has become clear that a second peak in incidence develops around age 60. In a study of >700 newly diagnosed sarcoidosis patients in the United States, one-half of the patients were ≥40 years at the time of diagnosis.

Although most cases of sarcoidosis are sporadic, a familial form of the disease exists. At least 5% of patients with sarcoidosis will have a family member with sarcoidosis. Sarcoidosis patients who are Irish or African American seem to have a two to three times higher rate of familial disease.

PATHOPHYSIOLOGY AND IMMUNOPATHOGENESIS

The granuloma is the pathologic hallmark of sarcoidosis. A distinct feature of sarcoidosis is the local accumulation of inflammatory cells. Extensive studies in the lung using bronchoalveolar lavage (BAL) have demonstrated that the initial inflammatory response is an influx of T helper cells. In addition, there is an accumulation of activated monocytes. **Figure 390-1** is a proposed model for sarcoidosis. Using the HLA-CD4 complex, antigen-presenting cells present an unknown antigen to the helper T cell. Studies have clarified that specific HLA haplotypes such as HLA-DRB1*1101 are associated with an increased risk for developing sarcoidosis. In addition, different HLA haplotypes are associated with different clinical outcomes.

The macrophage/helper T cell cluster leads to activation with the increased release of several cytokines. These include interleukin (IL)-2 released from the T cell and interferon γ and tumor necrosis factor (TNF) released by the macrophage. The T cell is a necessary part of the initial inflammatory response. In advanced, untreated HIV infection, patients who lack helper T cells rarely develop sarcoidosis. In contrast, several reports confirm that sarcoidosis becomes unmasked as HIV-infected individuals receive antiretroviral therapy, with subsequent restoration of their immune system. In contrast, treatment of established pulmonary sarcoidosis with cyclosporine, a drug that downregulates helper T cell responses, seems to have little impact on sarcoidosis.

The granulomatous response of sarcoidosis can resolve with or without therapy. However, in at least 20% of patients with sarcoidosis, a chronic form of the disease develops. This persistent form of the disease is associated with the secretion of high levels of IL-8. Also, studies have reported that patients with this chronic form of disease release excessive amounts of TNF in areas of inflammation. Specific

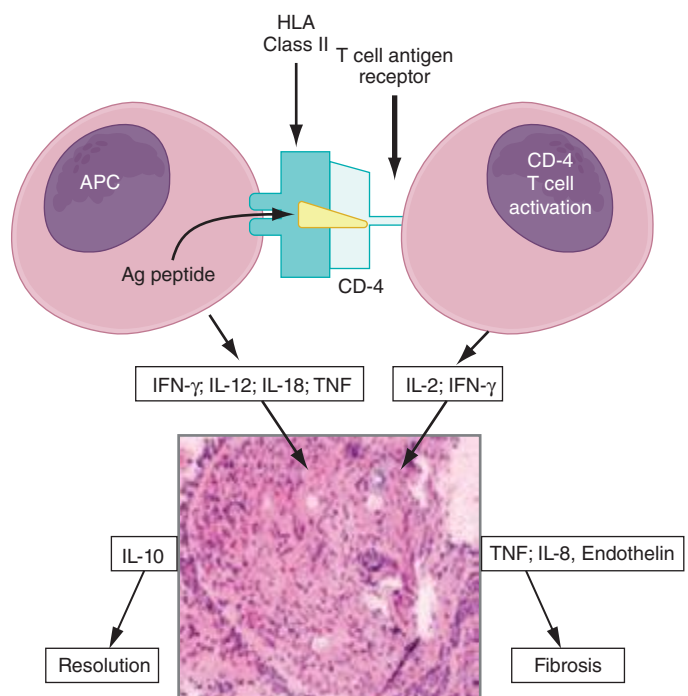


FIGURE 390-1 Schematic representation of initial events of sarcoidosis. The antigen-presenting cell and helper T cell complex leads to the release of multiple cytokines. This forms a granuloma. Over time, the granuloma may resolve or lead to chronic disease, including fibrosis. APC, antigen-presenting cell; HLA, human leukocyte antigen; IFN, interferon; IL, interleukin; TNF, tumor necrosis factor.

2206 gene signatures have been associated with more severe disease, such as cardiac, neurologic, and fibrotic pulmonary disease.

At diagnosis the natural history of the disease may be difficult to predict. One form of the disease, *Löfgren's syndrome*, consists of erythema nodosum and hilar adenopathy on chest roentgenogram. In some cases, periarticular arthritis may be identified without erythema nodosum. Löfgren's syndrome is associated with a good prognosis, with >90% of patients experiencing disease resolution within 2 years. Recent studies have demonstrated that the HLA-DRB1*03 was found in two-thirds of Scandinavian patients with Löfgren's syndrome. More than 95% of those patients who were HLA-DRB1*03 positive had resolution of their disease within 2 years, whereas nearly one-half of the remaining patients had disease for more than 2 years. It remains to be determined whether these observations can be applied to a non-Scandinavian population.

CLINICAL MANIFESTATIONS

The presentation of sarcoidosis ranges from patients who are asymptomatic to those with organ failure. It is unclear how often sarcoidosis is asymptomatic. In countries where routine chest roentgenogram screening is performed, 20–30% of pulmonary cases are detected in asymptomatic individuals. The inability to screen for other asymptomatic forms of the disease would suggest that as many as one-third of sarcoidosis patients are asymptomatic.

Respiratory complaints including cough and dyspnea are the most common presenting symptoms. In many cases, the patient presents with a 2- to 4-week history of these symptoms. Unfortunately, due to the nonspecific nature of pulmonary symptoms, the patient may see physicians for up to a year before a diagnosis is confirmed. For these patients, the diagnosis of sarcoidosis is usually only suggested when a chest roentgenogram is performed.

Symptoms related to cutaneous and ocular disease are the next two most common complaints. Skin lesions are often nonspecific. However, because these lesions are readily observed, the patient and treating physician are often led to a diagnosis. In contrast to patients with pulmonary disease, patients with cutaneous lesions are more likely to be diagnosed within 6 months of symptoms.

Nonspecific constitutional symptoms include fatigue, fever, night sweats, and weight loss. Fatigue is perhaps the most common constitutional symptom that affects these patients. Given its insidious nature, patients are usually not aware of the association with their sarcoidosis until their disease resolves.

The overall incidence of sarcoidosis at the time of diagnosis and eventual common organ involvement are summarized in **Table 390-1**. Over time, skin, eye, and neurologic involvement seem more apparent. In the United States, the frequency of specific organ involvement appears to be affected by age, race, and gender. For example, eye disease is more common among African Americans. Under the age of 40, it occurs more frequently in women. However, in those diagnosed over the age of 40, eye disease is more common in men.

TABLE 390-1 FREQUENCY OF COMMON ORGAN INVOLVEMENT AND LIFETIME RISK^a

	Presentation, % ^b	Follow-Up, % ^c
Lung	95	94
Skin	24	43
Eye	12	29
Extrathoracic lymph node	15	16
Liver	12	14
Spleen	7	8
Neurologic	5	16
Cardiac	2	3

^aPatients could have more than one organ involved. ^bFrom ACCESS study of 736 patients evaluated within 6 months of diagnosis. ^cFrom follow-up of 1024 sarcoidosis patients seen at the University of Cincinnati Interstitial Lung Disease and Sarcoidosis Clinic from 2002–2006.



FIGURE 390-2 Posterior-anterior chest roentgenogram demonstrating bilateral hilar adenopathy, stage 1 disease.

LUNG

Lung involvement occurs in >90% of sarcoidosis patients. The most commonly used method for detecting lung disease is still the chest roentgenogram. **Figure 390-2** illustrates the chest roentgenogram from a sarcoidosis patient with bilateral hilar adenopathy. Although the computed tomography (CT) scan has changed the diagnostic approach to interstitial lung disease, the CT scan is not usually considered a monitoring tool for patients with sarcoidosis. **Figure 390-3** demonstrates some of the characteristic CT features, including peribronchial thickening and reticular nodular changes, which are predominantly subpleural. The peribronchial thickening seen on CT scan seems to explain the high yield of granulomas from bronchial biopsies performed for diagnosis.

Although the CT scan is more sensitive, the standard scoring system described by Scadding in 1961 for chest roentgenograms remains the preferred method of characterizing the chest involvement. Stage 1 is hilar adenopathy alone (Fig. 390-2), often with right paratracheal involvement. Stage 2 is a combination of adenopathy plus infiltrates, whereas stage 3 reveals infiltrates alone. Stage 4 consists of fibrosis. Usually the infiltrates in sarcoidosis are predominantly an upper lobe process. Only in a few noninfectious diseases is an upper lobe

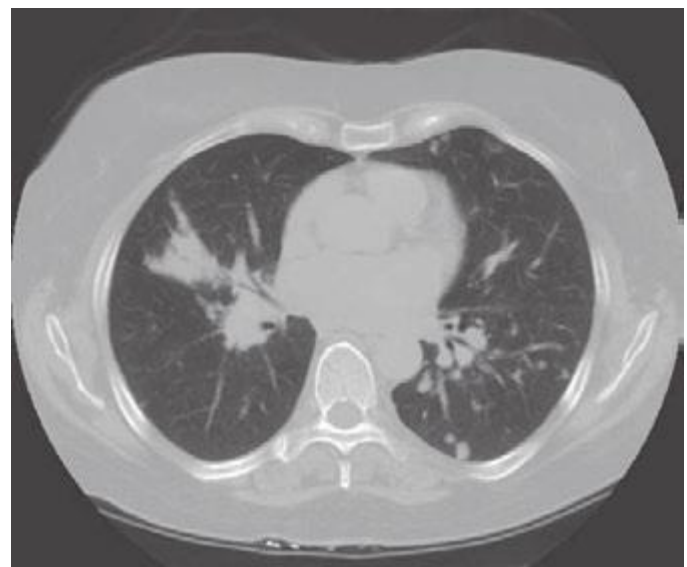


FIGURE 390-3 High-resolution computed tomography scan of chest demonstrating patchy reticular nodularity, including areas of confluence.

predominance noted. In addition to sarcoidosis, the differential diagnosis of upper lobe disease includes hypersensitivity pneumonitis, silicosis, and Langerhans cell histiocytosis. For infectious diseases, tuberculosis and *Pneumocystis pneumonia* can often present as upper lobe diseases.

Lung volumes, mechanics, and diffusion are all useful in evaluating interstitial lung diseases such as sarcoidosis. The diffusion of carbon monoxide (DL_{CO}) is the most sensitive test for an interstitial lung disease. Reduced lung volumes are a reflection of the restrictive lung disease seen in sarcoidosis. However, a third of the patients presenting with sarcoidosis still have lung volumes within the normal range, despite abnormal chest roentgenograms and dyspnea.

Approximately one-half of sarcoidosis patients present with obstructive disease, reflected by a reduced ratio of forced vital capacity expired in 1 second (FEV_1/FVC). Cough is a very common symptom. Airway involvement causing varying degrees of obstruction underlies the cough in most sarcoidosis patients. Airway hyperreactivity, as determined by methacholine challenge, will be positive in some of these patients. A few patients with cough will respond to traditional bronchodilators as the only form of treatment. In some cases, high-dose inhaled glucocorticoids alone are useful. Airway obstruction can be due to large airway stenosis, which can become fibrotic and unresponsive to anti-inflammatory therapy.

Pulmonary arterial hypertension is reported in at least 5% of sarcoidosis patients. Either direct vascular involvement or the consequence of fibrotic changes in the lung can lead to pulmonary arterial hypertension. In sarcoidosis patients with end-stage fibrosis awaiting lung transplant, 70% will have pulmonary arterial hypertension. This is a much higher incidence than that reported for other fibrotic lung diseases. In less advanced, but still symptomatic, patients, pulmonary arterial hypertension has been noted in up to 50% of the cases. Because sarcoidosis-associated pulmonary arterial hypertension may respond to therapy, evaluation for this should be considered in persistently dyspneic patients.

SKIN

Skin involvement is eventually identified in over a third of patients with sarcoidosis. The classic cutaneous lesions include erythema nodosum, maculopapular lesions, hyper- and hypopigmentation, keloid formation, and subcutaneous nodules. A specific complex of involvement of the bridge of the nose, the area beneath the eyes, and the cheeks is referred to as *lupus pernio* (Fig. 390-4) and is diagnostic for a chronic form of sarcoidosis.



FIGURE 390-4 Chronic inflammatory lesions around nose, eyes, and cheeks, referred to as lupus pernio.



FIGURE 390-5 Maculopapular lesions on the trunk of a sarcoidosis patient.

In contrast, erythema nodosum is a transient rash that can be seen in association with hilar adenopathy and uveitis (Löfgren's syndrome). Erythema nodosum is more common in women and in certain self-described demographic groups including whites and Puerto Ricans. In the United States, the other manifestations of skin sarcoidosis, especially lupus pernio, are more common in African Americans than whites.

The maculopapular lesions from sarcoidosis are the most common chronic form of the disease (Fig. 390-5). These are often overlooked by the patient and physician, because they are chronic and not painful. Initially, these lesions are usually purplish papules and are often indurated. They can become confluent and infiltrate large areas of the skin. With treatment, the color and induration may fade. Because these lesions are caused by noncaseating granulomas, the diagnosis of sarcoidosis can be readily made by a skin biopsy.

EYE

The frequency of ocular manifestations for sarcoidosis varies depending on race. In Japan, >70% of sarcoidosis patients develop ocular disease, whereas in the United States only 30% have eye disease, with problems more common in African Americans than whites. Although the most common manifestation is an anterior uveitis, over a quarter of patients will have inflammation at the posterior of the eye, including retinitis and pars planitis. Although symptoms such as photophobia, blurred vision, and increased tearing can occur, some asymptomatic patients still have active inflammation. Initially asymptomatic patients with ocular sarcoidosis can eventually develop blindness. Therefore, it is recommended that all patients with sarcoidosis receive a dedicated ophthalmologic examination. Sicca is seen in over one-half of the chronic sarcoidosis patients. Dry eyes appear to be a reflection of prior lacrimal gland disease. Although the patient may no longer have active inflammation, the dry eyes may require natural tears or other lubricants.

LIVER

Using biopsies to detect granulomatous disease, liver involvement can be identified in over one-half of sarcoidosis patients. However, using liver function studies, only 20–30% of patients will have evidence of liver involvement. The most common abnormality of liver function is an elevation of the alkaline phosphatase level, consistent with an obstructive pattern. In addition, elevated transaminase levels can occur. An elevated bilirubin level is a marker for more advanced liver disease. Overall, only 5% of sarcoidosis patients have sufficient symptoms from their liver disease to require specific therapy. Although symptoms can be due to hepatomegaly, more frequently symptoms

2208 result from extensive intrahepatic cholestasis leading to portal hypertension. In this case, ascites and esophageal varices can occur. It is rare that a sarcoidosis patient will require a liver transplant, because even the patient with cirrhosis due to sarcoidosis can respond to systemic therapy. On a cautionary note, patients with both sarcoidosis and hepatitis C should avoid therapy with interferon a because of its association with the development or worsening of granulomatous disease.

BONE MARROW AND SPLEEN

One or more bone marrow manifestations can be identified in many sarcoidosis patients. The most common hematologic problem is lymphopenia, which is a reflection of sequestration of the lymphocytes into the areas of inflammation. Anemia occurs in 20% of patients, and leukopenia is less common. Bone marrow examination will reveal granulomas in about a third of patients. Although splenomegaly can be detected in 5–10% of patients, splenic biopsy reveals granulomas in 60% of patients. The CT scan can be relatively specific for sarcoidosis involvement of the spleen (Fig. 390-6). Both bone marrow and spleen involvement are more common in African Americans than whites. Although these manifestations alone are rarely an indication for therapy, on rare occasion, splenectomy may be indicated for massive symptomatic splenomegaly or profound pancytopenia. Nonthoracic lymphadenopathy can occur in up to 20% of patients.

CALCIUM METABOLISM

Hypercalcemia and/or hypercalciuria occurs in about 10% of sarcoidosis patients. It is more common in whites than African Americans and in men. The mechanism of abnormal calcium metabolism is increased production of 1,25-dihydroxyvitamin D by the granuloma itself. The 1,25-dihydroxyvitamin D causes increased intestinal absorption of calcium, leading to hypercalcemia with a suppressed parathyroid hormone (PTH) level (Chap. 424). Increased exogenous vitamin D from diet or sunlight exposure may exacerbate this problem. Serum calcium should be determined as part of the initial evaluation of all sarcoidosis patients, and a repeat determination may be useful during the summer months with increased sun exposure. In patients with a history of renal calculi, a 24-h urine calcium measurement should be obtained. If a sarcoidosis patient with a history of renal calculi is to be placed on calcium supplements, a follow-up 24-h urine calcium level should be measured.

RENAL DISEASE

Direct kidney involvement occurs in <5% of sarcoidosis patients. It is associated with granulomas in the kidney itself and can lead to nephritis.

However, hypercalcemia is the most likely cause of sarcoidosis-associated renal disease. In 1–2% of sarcoidosis patients, acute renal failure may develop as a result of hypercalcemia. Successful treatment of hypercalcemia with glucocorticoids and other therapies often improves but usually does not totally resolve the renal dysfunction.

NERVOUS SYSTEM

Neurologic disease is reported in 5–10% of sarcoidosis patients and appears to be of equal frequency across all ethnic groups. Any part of the central or peripheral nervous system can be affected. The presence of granulomatous inflammation is often visible on magnetic resonance imaging (MRI) studies. The MRI with gadolinium enhancement may demonstrate space-occupying lesions, but the MRI can be negative due to small lesions or the effect of systemic therapy in reducing the inflammation. The cerebral spinal fluid (CSF) findings include lymphocytic meningitis with a mild increase in protein. The CSF glucose is usually normal but can be low. Certain areas of the nervous system are more commonly affected in neurosarcoidosis. These include cranial nerve involvement, basilar meningitis, myelopathy, and anterior hypothalamic disease with associated diabetes insipidus (Chap. 404). Seizures and cognitive changes also occur. Of the cranial nerves, seventh nerve paralysis can be transient and mistaken for Bell's palsy (idiopathic seventh nerve paralysis). Because this form of neurosarcoidosis often resolves within weeks and may not recur, it may have occurred prior to a definitive diagnosis of sarcoidosis. Optic neuritis is another cranial nerve manifestation of sarcoidosis. This manifestation is more chronic and usually requires long-term systemic therapy. It can be associated with both anterior and posterior uveitis. Differentiating between neurosarcoidosis and multiple sclerosis can be difficult at times. Optic neuritis can occur in both diseases. In some patients with sarcoidosis, multiple enhancing white matter abnormalities may be detected by MRI, suggesting multiple sclerosis. In such cases, the presence of meningeal enhancement or hypothalamic involvement suggests neurosarcoidosis, as does evidence of extraneurologic disease such as pulmonary or skin involvement, which also suggests sarcoidosis. Because the response of neurosarcoidosis to glucocorticoids and cytotoxic therapy is different from that of multiple sclerosis, differentiating between these disease entities is important.

CARDIAC

The presence of cardiac involvement is influenced by race. Although over a quarter of Japanese sarcoidosis patients develop cardiac disease, only 5% of sarcoidosis patients in the United States and Europe develop symptomatic cardiac disease. However, there is no apparent racial predilection between whites and African Americans. Cardiac disease, which usually presents as either congestive heart failure or cardiac arrhythmias, results from infiltration of the heart muscle by granulomas. Diffuse granulomatous involvement of the heart muscle can lead to profound dysfunction with left ventricular ejection fractions below 10%. Even in this situation, improvement in the ejection fraction can occur with systemic therapy. Arrhythmias can also occur with diffuse infiltration or with more patchy cardiac involvement. If the atrioventricular (AV) node is infiltrated, heart block can occur, which can be detected by routine electrocardiography. Ventricular arrhythmias and sudden death due to ventricular tachycardia are common causes of death. Arrhythmias are best detected using 24-h ambulatory monitoring, and electrophysiology studies may be negative. Other screening tests for cardiac disease include routine electrocardiography and echocardiography. The confirmation of cardiac sarcoidosis is usually performed with either MRI or positron emission tomography (PET) scanning. Because ventricular arrhythmias are usually multifocal due to patchy multiple granulomas in the heart, ablation therapy is not useful. Patients with significant ventricular arrhythmias should be considered for an implanted defibrillator, which appears to have reduced the rate of death in cardiac sarcoidosis. Although systemic therapy can be useful in treating the arrhythmias, patients may still have malignant arrhythmias up to 6 months after starting successful treatment, and the risk for recurrent arrhythmias occurs whenever medications are tapered.



FIGURE 390-6 Computed tomography scan of the abdomen after oral and intravenous contrast. The stomach is compressed by the enlarged spleen. Within the spleen, areas of hypo- and hyperdensity are identified.



FIGURE 390-7 Positron emission tomography and computed tomography scan merged demonstrating increased activity in spleen, ribs, and spine of patient with sarcoidosis.

MUSCULOSKELETAL SYSTEM

Direct granulomatous involvement of bone and muscle can be documented by radiography (x-ray, MRI, PET scan [Fig. 390-7], or gallium scan) or confirmed by biopsy in about 10% of sarcoidosis patients. However, a larger percentage of sarcoidosis patients complain of myalgias and arthralgias. These complaints are similar to those reported by patients with other inflammatory diseases, including chronic infections such as mononucleosis. Fatigue associated with sarcoidosis may be overwhelming for many patients. Recent studies have demonstrated a link between fatigue and small peripheral nerve fiber disease in sarcoidosis.

OTHER ORGAN INVOLVEMENT

Although sarcoidosis can affect any organ of the body, rarely does it involve the breast, testes, ovary, or stomach. Because of the rarity of involvement, a mass in one of these areas requires a biopsy to rule out other diseases including cancer. For example, in a study of breast problems in female sarcoidosis patients, a breast lesion was more likely to be a granuloma from sarcoidosis than from breast cancer. However, findings on the physical examination or mammogram cannot reliably differentiate between these lesions. More importantly, as women with sarcoidosis age, breast cancer becomes more common. Therefore, it is recommended that routine screening including mammography be performed along with other imaging studies (ultrasound, MRI) or biopsy as clinically indicated.

COMPLICATIONS

Sarcoidosis is usually a self-limited, non-life-threatening disease. However, organ-threatening disease can occur. These complications can include blindness, paraplegia, or renal failure. Death from sarcoidosis occurs in about 5% of patients seen in sarcoidosis referral clinics. The usual causes of death related to sarcoidosis are from lung, cardiac, neurologic, or liver involvement. In respiratory failure, an elevation of the right atrial pressure is a poor prognostic finding. Lung complications can also include infections such as mycetoma, which can

subsequently lead to massive bleeding. In addition, the use of immunosuppressive agents can increase the incidence of serious infections.

LABORATORY FINDINGS

The chest roentgenogram remains the most commonly used tool to assess lung involvement in sarcoidosis. As noted above, the chest roentgenogram classifies involvement into four stages, with stages 1 and 2 having hilar and paratracheal adenopathy. The CT scan has been used increasingly in evaluating interstitial lung disease. In sarcoidosis, the presence of adenopathy and a nodular infiltrate is not specific for sarcoidosis. Adenopathy up to 2 cm can be seen in other inflammatory lung diseases such as idiopathic pulmonary fibrosis. However, adenopathy >2 cm in the short axis supports the diagnosis of sarcoidosis over other interstitial lung diseases.

The PET scan has increasingly replaced gallium-67 scanning to identify areas of granulomatous disease in the chest and other parts of the body (Fig. 390-7). Both tests can be used to identify potential areas for biopsy. Cardiac PET scanning has also proved useful in assessing cardiac sarcoidosis. The identification of hypermetabolic activity may be due to the granulomas from sarcoidosis and not to disseminated malignancy.

MRI has also proved useful in the assessment of extrapulmonary sarcoidosis. Gadolinium enhancement has been demonstrated in areas of inflammation in the brain, heart, and bone. MRI scans may detect asymptomatic lesions. Like PET scan, MRI changes appear similar to those seen with malignancy and infection. In some cases, biopsy may be necessary to determine the cause of the radiologic abnormality.

Serum levels of angiotensin-converting enzyme (ACE) can be helpful in the diagnosis of sarcoidosis. However, the test has somewhat low sensitivity and specificity. Elevated levels of ACE are reported in 60% of patients with acute disease and only 20% of patients with chronic disease. Although there are several causes for mild elevation of ACE, including diabetes, elevations of >50% of the upper limit of normal are seen in only a few conditions including sarcoidosis, leprosy, Gaucher's disease, hyperthyroidism, and disseminated granulomatous infections such as miliary tuberculosis. Because the ACE level is determined by a biologic assay, the concurrent use of an ACE inhibitor such as lisinopril will lead to a very low ACE level.

DIAGNOSIS

The diagnosis of sarcoidosis requires both compatible clinical features and pathologic findings. Because the cause of sarcoidosis remains elusive, the diagnosis cannot be made with 100% certainty. Nevertheless, the diagnosis can be made with reasonable certainty based on history and physical features along with laboratory and pathologic findings.

Patients are usually evaluated for possible sarcoidosis based on two scenarios (Fig. 390-8). In the first scenario, a patient may undergo a biopsy revealing a noncaseating granuloma in either a pulmonary or an extrapulmonary organ. If the clinical presentation is consistent with sarcoidosis and there is no alternative cause for the granulomas identified, then the patient is felt to have sarcoidosis.

In the second scenario, signs or symptoms suggesting sarcoidosis such as the presence of bilateral adenopathy may be present in an otherwise asymptomatic patient or a patient with uveitis or a rash consistent with sarcoidosis. At this point, a diagnostic procedure should be performed. For the patient with a compatible skin lesion, a skin biopsy should be considered. Other biopsies to consider could include liver, extrathoracic lymph node, or muscle. In some cases, a biopsy of the affected organ may not be easy to perform (such as a brain or spinal cord lesion). In other cases, such as an endomyocardial biopsy, the likelihood of a positive biopsy is low. Because of the high rate of pulmonary involvement in these cases, the lung may be easier to approach by bronchoscopy. During the bronchoscopy, a transbronchial biopsy, bronchial biopsy, or transbronchial needle aspirate can be performed. The endobronchial ultrasonography-guided (EBUS) transbronchial needle aspirate can assist in diagnosing sarcoidosis in patients with mediastinal adenopathy (stage 1 or 2 radiographic pulmonary disease), whereas transbronchial biopsy has a higher diagnostic yield for those

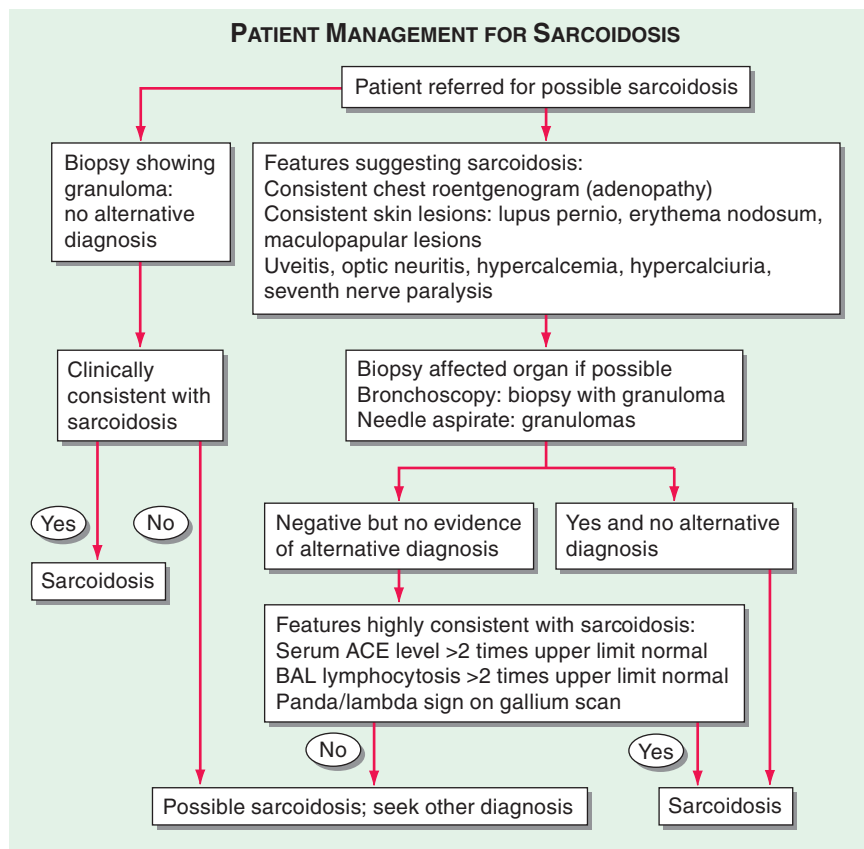


FIGURE 390-8 Proposed approach to management of patient with possible sarcoidosis. Presence of one or more of these features supports the diagnosis of sarcoidosis: uveitis, optic neuritis, hypercalcemia, hypercalciuria, seventh cranial nerve paralysis, diabetes insipidus. ACE, angiotensin-converting enzyme; BAL, bronchoalveolar lavage.

with only parenchymal lung disease (stage 3). These tests are complementary and may be performed together.

If the biopsy reveals granulomas, an alternative diagnosis such as infection or malignancy must be excluded. Bronchoscopic washings can be sent for cultures for fungi and tuberculosis. For the pathologist, the more tissue that is provided, the more comfortable is the diagnosis of sarcoidosis. A needle aspirate may be adequate in an otherwise classic case of sarcoidosis, but may be insufficient in a patient in whom lymphoma or fungal infection is a likely alternative diagnosis. Because granulomas can be seen on the edge of a lymphoma, the presence of a few granulomas from a needle aspirate may not be sufficient to clarify the diagnosis. Mediastinoscopy provides a larger sample to confirm the presence or absence of lymphoma in the mediastinum. Alternatively, for most patients, evidence of extrathoracic disease (e.g., eye involvement) may further support the diagnosis of sarcoidosis.

For patients with negative pathology, positive supportive tests may increase the likelihood of the diagnosis of sarcoidosis. These tests include an elevated ACE level, which can also be elevated in other granulomatous diseases but not in malignancy. A positive PET scan can support the diagnosis if multiple organs are affected. A BAL is often performed during the bronchoscopy. An increase in the percentage of lymphocytes supports the diagnosis of sarcoidosis. The use of the lymphocyte markers CD4 and CD8 can be used to determine the CD4/CD8 ratio of these increased lymphocytes in the BAL fluid. A ratio of >3.5 is strongly supportive of sarcoidosis but is less sensitive than an increase in lymphocytes alone. Although in general, an increase in BAL lymphocytes is supportive of the diagnosis, other conditions must be considered.

Supportive findings, when combined with commonly associated but nondiagnostic clinical features of the disease, improve the diagnostic probability of sarcoidosis. These clinical features include uveitis, renal stones, hypercalcemia, seventh cranial nerve paralysis, or erythema

nodosum. The presence of one or more of these features in a patient suspected of having sarcoidosis increases the probability of sarcoidosis.

The *Kviem-Siltzbach procedure* is a specific diagnostic test for sarcoidosis. An intradermal injection of specially prepared tissue derived from the spleen of a known sarcoidosis patient is biopsied 4–6 weeks after injection. If noncaseating granulomas are seen, this is highly specific for the diagnosis of sarcoidosis. Unfortunately, there is no commercially available Kviem-Siltzbach reagent, and some locally prepared batches have lower specificity. Thus, this test is of historic interest and is rarely used in current clinical practice.

Because the diagnosis of sarcoidosis can never be certain, over time other features may arise that lead to an alternative diagnosis. Conversely, evidence for new organ involvement may eventually confirm the diagnosis of sarcoidosis.

PROGNOSIS

The risk of death or loss of organ function remains low in sarcoidosis. Poor outcomes usually occur in patients who present with advanced disease in whom treatment seems to have little impact. In these cases, irreversible fibrotic changes have frequently occurred. Over the past 20 years, the reported mortality from sarcoidosis has increased in the United States and England. Whether this is due to heightened awareness of the chronic nature of this disease or to other factors such as more widespread immunosuppressive therapy usage remains unclear.

For the majority of patients, initial presentation occurs during the granulomatous phase of the disease as depicted in Fig. 390-1. It is clear that

many patients resolve their disease within 2–5 years. These patients are felt to have acute, self-limiting sarcoidosis. However, there is a form of the disease that does not resolve within the first 2–5 years. These chronic patients can be identified at presentation by certain risk factors at presentation such as fibrosis on chest roentgenogram, presence of lupus pernio, bone cysts, cardiac or neurologic disease (except isolated seventh nerve paralysis), and presence of renal calculi due to hypercalciuria. Recent studies also indicate that patients who require glucocorticoids for any manifestation of their disease in the first 6 months of presentation have a $>50\%$ chance of having chronic disease. In contrast, $<10\%$ of patients who require no systemic therapy in the first 6 months will require chronic therapy.

TREATMENT SARCOIDOSIS

Indications for therapy should be based on symptoms or presence of organ- or life-threatening disease, including disease involving the eye, heart, or nervous system. The patient with asymptomatic elevated liver function tests or an abnormal chest roentgenogram probably does not benefit from treatment. However, these patients should be monitored for evidence of progressive, symptomatic disease.

One approach to therapy is summarized in Figs. 390-9 and 390-10. We have divided the approach into treating acute versus chronic disease. For acute disease, no therapy remains a viable option for patients with no or mild symptoms. For symptoms confined to only one organ, topical therapy is preferable. For multi-organ disease or disease too extensive for topical therapy, an approach to systemic therapy is outlined. Glucocorticoids remain the drugs of choice for this disease. However, the decision to continue to treat with glucocorticoids or to add steroid-sparing agents depends on the tolerability, duration, and dosage of glucocorticoids.

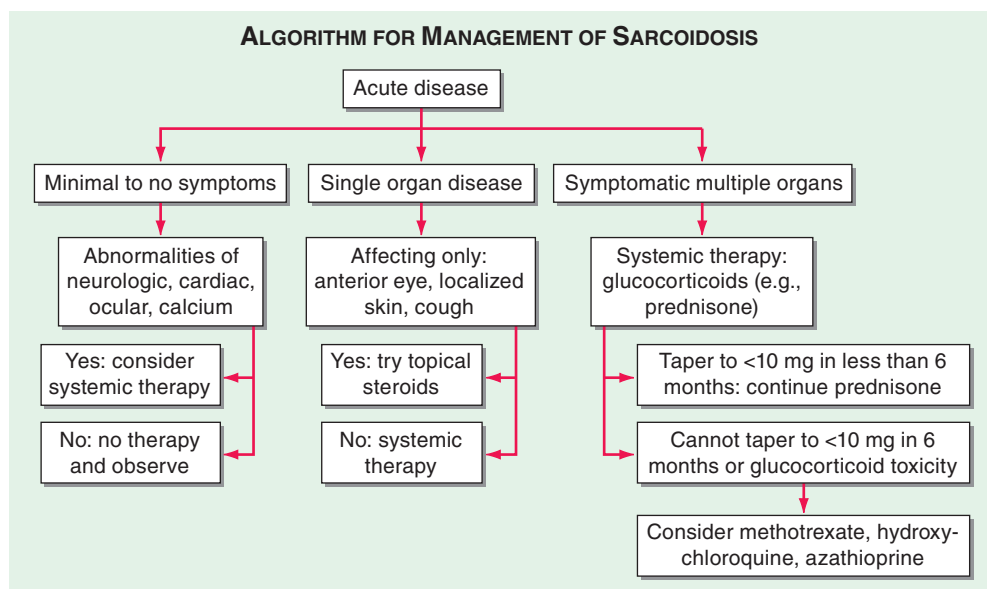


FIGURE 390-9 The management of acute sarcoidosis is based on level of symptoms and extent of organ involvement. In patients with mild symptoms, no therapy may be needed unless specified manifestations are noted.

Table 390-2 summarizes the dosage and monitoring of several commonly used drugs. According to the available trials, evidence-based recommendations are made. Most of these recommendations are for pulmonary disease because most of the trials were performed only in pulmonary disease. Treatment recommendations for extrapulmonary disease are usually similar with a few modifications. For example, the dosage of glucocorticoids is usually higher for neurosarcoidosis and lower for cutaneous disease. There was some suggestion that higher doses would be beneficial for cardiac sarcoidosis, but one study found that initial prednisone doses >40 mg/d were associated with a worse outcome because of toxicity.

Systemic therapies for sarcoidosis are usually immunosuppressive including glucocorticoids, cytotoxics, or biologics. Although most patients receive glucocorticoids as their initial systemic therapy, toxicity associated with prolonged therapy often leads to steroid-sparing alternatives. The antimalarial drugs such as hydroxychloroquine are more effective for skin than pulmonary disease. Minocycline may also be useful for cutaneous sarcoidosis.

For pulmonary and other extrapulmonary disease, cytotoxic agents are often used. These include methotrexate, azathioprine, leflunomide, mycophenolate, and cyclophosphamide. The most widely studied cytotoxic agent has been methotrexate. This agent works in approximately two-thirds of sarcoidosis patients, regardless of the disease manifestation. In one retrospective study comparing methotrexate to azathioprine, both drugs were equally effective. However, methotrexate was associated with significantly less toxicity. As noted in Table 390-2, specific guidelines for monitoring therapy have been recommended. Cytokine modulators such as thalidomide and pentoxifylline have also been used in a limited number of cases.

The biologic anti-TNF agents have recently been studied in sarcoidosis, with prospective randomized trials completed for both etanercept and infliximab. Etanercept has a limited role as a steroid-sparing agent. Conversely, infliximab significantly improved lung function when administered to glucocorticoid and cytotoxic pretreated patients with chronic disease. The difference in response between these two agents is similar to that observed in

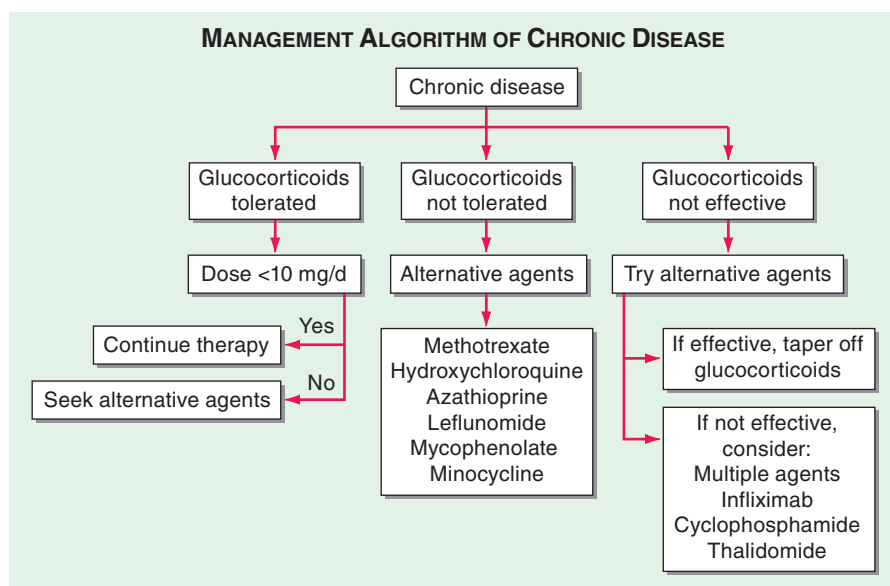


FIGURE 390-10 Approach to chronic disease is based on whether glucocorticoid therapy is tolerated or not.

TABLE 390-2 COMMONLY USED DRUGS TO TREAT SARCOIDOSIS

Drug	Initial Dose	Maintenance Dose	Monitoring	Toxicity	Support Therapy ^a	Support Monitoring ^a
Prednisone	20–40 mg qd	Taper to 5–10 mg	Glucose, blood pressure, bone density	Diabetes, osteoporosis	A: Acute pulmonary D: Extrapulmonary	
Hydroxychloroquine	200–400 mg qd	400 mg qd	Eye exam q6–12 mo	Ocular	B: Some forms of disease	D: Routine eye exam
Methotrexate	10 mg qwk	2.5–15 mg qwk	CBC, renal, hepatic q2mo	Hematologic, nausea, hepatic, pulmonary	B: Steroid sparing C: Some forms chronic disease	D: Routine hematologic, renal, and hepatic monitoring
Azathioprine	50–150 mg qd	50–200 mg qd	CBC, renal q2mo	Hematologic, nausea	C: Some forms chronic disease	D: Routine hematologic monitoring
Infliximab	3–5 mg/kg q2wk for 2 doses	3–10 mg/kg q4–8 wk	Initial PPD	Infections, allergic reaction, carcinogen	A: Chronic pulmonary disease	B: Caution in patients with latent tuberculosis or advanced congestive heart failure

^aGrade A: supported by at least two double-blind randomized control trials; grade B: supported by prospective cohort studies; grade C: supported primarily by two or more retrospective studies; grade D: only one retrospective study or based on experience in other diseases.

Abbreviations: CBC, complete blood count; PPD, purified protein derivative test for tuberculosis.

Source: Adapted from RP Baughman, O Selroos: Evidence-based approach to treatment of sarcoidosis, in PG Gibson et al (eds): *Evidence-Based Respiratory Medicine*. Oxford, BMJ Books Blackwell, 2005, pp 491–508.

Crohn's disease, where infliximab is effective and etanercept is not. However, there is a higher risk for reactivation of tuberculosis with infliximab compared to etanercept. The differential response rate could be explained by differences in mechanism of action because etanercept is a TNF receptor antagonist and infliximab is a monoclonal antibody against TNF. In contrast to etanercept, infliximab also binds to TNF on the surface of some cells that release TNF, which leads to cell lysis. This effect has been documented in Crohn's disease. Adalimumab is a humanized monoclonal anti-TNF antibody that also appears effective for sarcoidosis when dosed at higher strengths, as recommended for the treatment of Crohn's disease. The role of the newer therapeutic agents for sarcoidosis is still evolving. However, these targeted therapies confirm that TNF may be an important target, especially in the treatment of chronic disease. However, these agents are not a panacea, because sarcoidosis-like disease has occurred in patients treated with anti-TNF agents for nonsarcoidosis indications.

IgG4-related disease (IgG4-RD) is a fibroinflammatory condition characterized by a tendency to form tumefactive lesions. The clinical manifestations of this disease, however, are protean, and continue to be defined. IgG4-RD has now been described in virtually every organ system. Commonly affected organs are the biliary tree, salivary glands, periorbital tissues, kidneys, lungs, lymph nodes, and retroperitoneum. In addition, IgG4-RD involvement of the meninges, aorta, prostate, thyroid, pericardium, skin, and other organs is well described. The disease is believed to affect the brain parenchyma, the joints, the bone marrow, and the bowel mucosa only rarely (if ever).

The clinical features of IgG4-RD are numerous, but the pathologic findings are consistent across all affected organs. These findings include a lymphoplasmacytic infiltrate with a high percentage of IgG4-positive plasma cells; a characteristic pattern of fibrosis termed “storiform”; a tendency to target blood vessels, particularly veins, through an obliterative process (“obliterative phlebitis”); and a mild to moderate tissue eosinophilia.

IgG4-RD encompasses a number of conditions previously regarded as separate, organ-specific entities. A condition once known as “lymphoplasmacytic sclerosing pancreatitis” (among many other terms) became the paradigm of IgG4-RD in 2000, when Japanese investigators recognized that these patients had elevated serum concentrations of IgG4. This form of sclerosing pancreatitis is now termed type 1 (IgG4-related) autoimmune pancreatitis (AIP). By 2003, extrapancreatic

disease manifestations had been identified in patients with type 1 AIP, and since then, the manifestations of IgG4-RD in many organs have been catalogued. *Mikulicz’s disease*, once considered to be a subset of Sjögren’s syndrome that affected the lacrimal, parotid, and submandibular glands, is now considered part of the IgG4-RD spectrum. Similarly, a subset of patients previously diagnosed as having primary sclerosing cholangitis was known to respond well to glucocorticoids, in contrast to the majority of patients with that diagnosis. This steroid-responsive subset is now explained by the fact that such patients actually have a separate disease, i.e., IgG4-related sclerosing cholangitis. In this manner, the understanding of IgG4-RD has extended to include nearly every specialty of medicine.

CLINICAL FEATURES

The major organ lesions are summarized in [Table 391e-1](#). IgG4-RD usually presents subacutely, and most patients do not have severe constitutional symptoms. Fevers and dramatic elevations of C-reactive protein are unusual; however, some patients report substantial weight loss occurring over periods of months. Clinically apparent disease can evolve over months, years, or even decades before the manifestations within a given organ becomes sufficiently severe to bring the patient to medical attention. Some patients have disease that is marked by the appearance and then resolution or temporary improvement in symptoms within a particular organ. Other patients accumulate new organ involvement as their disease persists in previously affected organs. Many patients with IgG4-RD are misdiagnosed as having other conditions, particularly malignancies, or their findings are attributed initially to nonspecific inflammation. The disorder is often identified incidentally through radiologic findings or unexpectedly in pathology specimens.

TABLE 391e-1 ORGAN MANIFESTATIONS OF IgG4-RELATED DISEASE

Organ	Major Clinical Features
Orbits and periorbital tissues	Painless eyelid or periocular tissue swelling; orbital pseudotumor; dacryoadenitis; dacryocystitis; orbital myositis; and mass lesions extending into the pterygopalatine fossa and infiltrating along the trigeminal nerve
Ears, nose, and sinuses	Allergic phenomena (nasal polyps, asthma, allergic rhinitis, peripheral eosinophilia); nasal obstruction, rhinorrhea, anosmia, chronic sinusitis; occasional bone-destructive lesions
Salivary glands	Submandibular and/or parotid gland enlargement (isolated bilateral submandibular gland involvement more common); minor salivary glands sometimes involved
Meninges	Headache, radiculopathy, cranial nerve palsies, or other symptoms resulting from spinal cord compression; tendency to form mass lesions; magnetic resonance imaging shows marked thickening and enhancement of dura
Hypothalamus and pituitary	Clinical syndromes resulting from involvement of the hypothalamus and pituitary, e.g., anterior pituitary hormone deficiency, central diabetes insipidus, or both; imaging reveals thickened pituitary stalk or mass formation on the stalk, swelling of the pituitary gland, or mass formation within the pituitary
Lymph nodes	Generalized lymphadenopathy or localized disease adjacent to a specific affected organ; the lymph nodes involved are generally 1–2 cm in diameter and nontender
Thyroid gland	Riedel’s thyroiditis; fibrosing variant of Hashimoto’s thyroiditis
Lungs	Asymptomatic finding on lung imaging; cough, hemoptysis, dyspnea, pleural effusion, or chest discomfort; associated with parenchymal lung involvement, pleural disease, or both; four main clinical syndromes: inflammatory pseudotumor, central airway disease, localized or diffuse interstitial pneumonia, and pleuritis; pleural lesions have severe, nodular thickening of the visceral or parietal pleura with diffuse sclerosing inflammation, sometimes associated with pleural effusion
Aorta	Asymptomatic finding on radiologic studies; surprise finding at elective aortic surgery; aortic dissection; clinicopathologic syndromes described include lymphoplasmacytic aortitis of thoracic or abdominal aorta, aortic dissection, periaortitis and periarteritis, and inflammatory abdominal aneurysm
Retroperitoneum	Backache, lower abdominal pain, lower extremity edema, hydronephrosis from ureteral involvement, asymptomatic finding on radiologic studies
Kidneys	Tubulointerstitial nephritis; membranous glomerulonephritis in a small minority; asymptomatic tumoral lesions, typically multiple and bilateral, are sometimes detected on radiologic studies; renal involvement strongly associated with hypocomplementemia
Pancreas	Type 1 autoimmune pancreatitis, presenting as mild abdominal pain; weight loss; acute, obstructive jaundice, mimicking adenocarcinoma of the pancreas (including a pancreatic mass); between 20 and 50% of patients present with acute glucose intolerance; imaging shows diffuse (termed “sausage-shaped pancreas”) or segmental pancreatic enlargement, with loss of normal lobularity; a mass often raises the suspicion of malignancy
Biliary tree	Obstructive jaundice associated with autoimmunity in most cases; weight loss; steatorrhea; abdominal pain; and new-onset diabetes mellitus; mimicker of primary sclerosing cholangitis
Liver	Painless jaundice associated with mild to moderate abdominal discomfort, weight loss, steatorrhea; new-onset diabetes mellitus; mimicker of primary sclerosing cholangitis and cholangiocarcinoma
Other organs involved	Gallbladder, breast (pseudotumor), prostate (prostatitis), pericardium (constrictive pericarditis), mesentery (sclerosing mesenteritis), mediastinum (fibrosing mediastinitis), skin (erythematous or flesh-colored papules), peripheral nerve (perineural inflammation)

Multiorgan disease may be evident at diagnosis but can also evolve over months to years. Some patients have disease confined to a single organ for many years. Others have either known or subclinical organ involvement at the same time as the major clinical feature. Patients with type 1 AIP may have their major disease focus in the pancreas; however, thorough evaluations by history, physical examination, blood tests, urinalysis, and cross-sectional imaging may demonstrate lacrimal gland enlargement, sialoadenitis, lymphadenopathy, a variety of pulmonary findings, tubulointerstitial nephritis, hepatobiliary disease, aortitis, retroperitoneal fibrosis, or other organ involvement. Spontaneous improvement, sometimes leading to clinical resolution of certain organ system manifestations, is reported in a small percentage of patients.

Two common characteristics of IgG4-RD are allergic disease and the tendency to form tumefactive lesions that mimic malignancies (Fig. 391e-1). Many IgG4-RD patients have allergic features such as atopy, eczema, asthma, nasal polyps, sinusitis, and modest peripheral eosinophilia. IgG4-RD also appears to account for a significant proportion of tumorous swellings—pseudotumors—in many organ systems. Some patients undergo major surgeries (e.g., Whipple procedures or thyroidectomy) for the purpose of resecting malignancies before the correct diagnosis is identified. Frequent sites of pseudotumors are the major salivary glands, lacrimal glands, lungs, and kidneys; however, nearly all organs have been affected with this manifestation.



A



B

FIGURE 391e-1 A major clinical feature of IgG4-related disease is its tendency to form tumefactive lesions. Shown here are mass lesions of the lacrimal glands (A) and the submandibular glands (B).

IgG4-RD often causes major morbidity and can lead to organ failure; however, its general pattern is to cause damage in a subacute manner. Destructive bone lesions in the sinuses, head, and middle ear spaces that mimic granulomatous polyangiitis (formerly Wegener's granulomatosis) also occur in IgG4-RD; less aggressive lesions are the rule in most organs. In regions such as the retroperitoneum, substantial fibrosis often occurs before the diagnosis is established, leading to ureteral entrapment, hydronephrosis, postobstructive uropathy, renal atrophy, and chronic pain, possibly resulting from the encasement of peripheral nerves by the inflammatory process. Undiagnosed or undertreated IgG4-related cholangitis can lead to hepatic failure within months. Similarly, IgG4-related aortitis, believed to be associated with between 10 and 50% of cases of inflammatory aortitis, can cause aneurysms and dissections. Substantial renal dysfunction and even renal failure can ensue from IgG4-related tubulointerstitial nephritis, and renal atrophy is a frequent sequel to this disease complication.

SEROLOGIC FINDINGS

The majority of patients with IgG4-RD have elevated serum IgG4 concentrations; however, the range of elevation varies widely. Serum concentrations of IgG4 as high as 30 or 40 times the upper limit of normal sometimes occur, usually in patients with disease that affects multiple organ systems simultaneously. Approximately 30% of patients have normal serum IgG4 concentrations despite classic histopathologic and immunohistochemical findings. Such patients tend to have disease that affects fewer organs. Patients with IgG4-related retroperitoneal fibrosis have a high likelihood of normal serum IgG4 concentrations, perhaps because the process has advanced to a fibrotic stage by the time the diagnosis is considered.

The correlation between serum IgG4 concentrations and disease activity and the need for treatment is imperfect. Serum IgG4 concentrations typically decline swiftly with the institution of therapy but often do not normalize completely. Patients can achieve clinical remissions yet have persistently elevated serum IgG4 concentrations. Rapidly rising serum IgG4 concentrations may identify patients at greatest risk for clinical flares and monitoring of serial IgG4 concentrations identifies early relapse in some patients; however, the temporal relationship between modest IgG4 elevations and the need for clinical treatment is poor. Clinical relapses occur in some patients despite persistently normal IgG4 concentrations.

IgG4 concentrations in serum are usually measured by nephelometry assays. These assays can lead to reports of spuriously low IgG4 values because of the prozone effect. This effect can be corrected by dilution of the serum sample in the laboratory. The prozone effect should be considered when the results of serologic testing for IgG4 concentrations appear to be at odds with the patient's clinical features.

EPIDEMIOLOGY

The typical patient with IgG4-RD is a middle-aged to elderly man. This epidemiology stands in stark contrast to that of many classic autoimmune conditions, which tend to affect young women. Studies of AIP patients in Japan indicate that the male-to-female ratio in that disease subset is on the order of 3:1. Even more striking, male predominance has been reported in IgG4-related tubulointerstitial nephritis and IgG4-related retroperitoneal fibrosis. Among IgG4-RD manifestations that involve organs of the head and neck, the sex ratio may be closer to 1:1.

PATHOLOGY

The key histopathology characteristics of IgG4-RD are a dense lymphoplasmacytic infiltrate (Fig. 391e-2) that is organized in a storiform pattern (resembling a basket-weave), obliterative phlebitis, and a mild to moderate eosinophilic infiltrate. Lymphoid follicles and germinal centers are frequently observed. The infiltrate tends to aggregate around ductal structures when it affects glands such as the lacrimal, submandibular, and parotid glands or the pancreas. The inflammatory lesion often aggregates into tumefactive masses that destroy the involved tissue.

Obliterative arteritis is observed in some organs, particularly the lung; however, venous involvement is more common (and is indeed

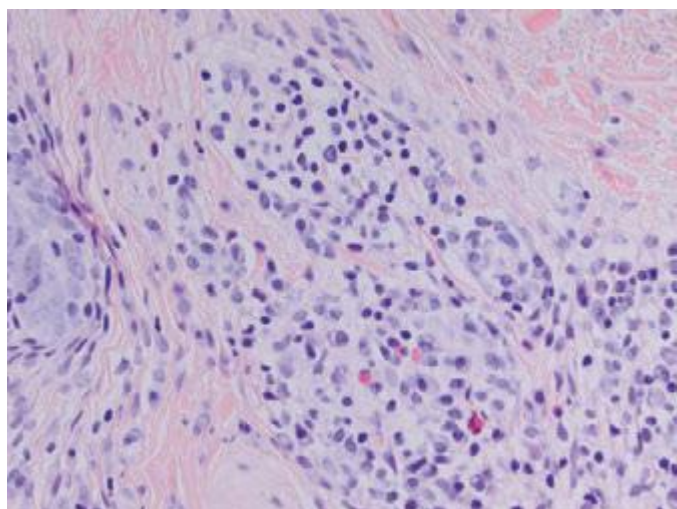


FIGURE 391e-2 Hallmark histopathology characteristics of IgG4-related disease (IgG4-RD) are a dense lymphoplasmacytic infiltrate and a mild to moderate eosinophilic infiltrate. The cellular inflammation is often encased in a distinctive type of fibrosis termed “storiform,” which often has a basket weave pattern. Abundant fibroblasts and strands of fibrosis accompany the lymphoplasmacytic infiltrate and eosinophils in this figure. This biopsy was taken from a nodular lesion on the cheek; however, the findings are identical to the pathology found in the pancreas, kidneys, lungs, salivary glands, and other organs affected by IgG4-RD.

a hallmark of IgG4-RD). Several histopathology features are uncommon in IgG4-RD and, when detected, mitigate against the diagnosis of IgG4-RD. These include intense neutrophilic infiltration, leukocytoclasia, granulomatous inflammation, multinucleated giant cells, and fibrinoid necrosis.

The inflammatory infiltrate is composed of an admixture of B and T lymphocytes. B cells are typically organized in germinal centers. Plasma cells staining for CD19, CD138, and IgG4 appear to radiate out from the germinal centers. In contrast, the T cells, usually CD4+, are distributed more diffusely throughout the lesion and generally represent the most abundant cell type. Fibroblasts, histiocytes, and eosinophils can all be observed in moderate numbers. Some biopsy samples are particularly enriched with eosinophils. In other samples, particularly from long-standing cases, fibrosis predominates.

The histologic appearance of IgG4-RD, although highly characteristic, requires immunohistochemical confirmation of the diagnosis with IgG4 immunostaining. IgG4-positive plasma cells predominate within the lesion, but plasma cells containing immunoglobulins from each subclass can be found. The number of IgG4-positive plasma cells can be quantified by either counting the number of cells per high-power field (HPF) or by calculating the ratio of IgG4- to IgG-bearing plasma cells. Tissue fibrosis predominates in the latter phases of organ involvement, and in this relatively acellular phase of inflammation, both the IgG4:total IgG ratio and the pattern of tissue fibrosis are more important than the number of IgG4-positive cells per HPF in establishing the diagnosis. In situ hybridization techniques are also now used to circumvent problems posed by increased background staining in conventional immunostaining techniques.

PATHOPHYSIOLOGY

The IgG4 molecule is believed to play an indirect role in the pathophysiology of disease in most organs. However, the molecule has properties that are unique among the immunoglobulin subclasses and that may contribute to tissue injury in some circumstances. As an example, IgG4 molecules have the ability to undergo Fab exchange, a phenomenon in which the two halves of the molecule dissociate from each other and reassociate with dissimilar hemi-molecules from other IgG4 molecules. This property is unique among the immunoglobulin subclasses. Partly as a result of Fab exchange, however, IgG4 antibodies bind antigen loosely. The molecules have low affinities for Fc receptors and C1q and are regarded generally as noninflammatory immunoglobulins. The low affinities for Fc receptors and C1q impair the ability of IgG4 antibodies to induce phagocyte activation, antibody-dependent cellular cytotoxicity, and complement-mediated damage. It is possible that the increased concentrations of IgG4 in serum and IgG4-bearing plasma cells in tissue are merely the result of other effector pathways, such as T_H2 /Treg cytokines, that are more central to the inflammation and tissue damage.

TREATMENT

Not every disease manifestation of IgG4-RD requires immediate treatment because the disease takes an indolent form in many patients. IgG4-related lymphadenopathy, for example, can be asymptomatic for years, without evolution to other disease manifestations. Thus, watchful waiting is prudent in some cases. Vital organ involvement must be treated aggressively, however, because IgG4-RD can lead to serious organ dysfunction and failure. Aggressive disease can lead quickly to end-stage liver disease, permanent impairment of pancreatic function, renal atrophy, aortic dissection or aneurysms, and destructive lesions in the sinuses and nasopharynx.

Glucocorticoids are the first line of therapy. Treatment regimens, extrapolated from experience with the management of type 1 AIP, generally begin with 40 mg/d of prednisone, with tapering to discontinuation or maintenance doses of 5 mg/d within 2 or 3 months. The clinical response to glucocorticoids is usually swift and striking; however, longitudinal data indicate that disease flares occur in more than 90% of patients within 3 years. Conventional steroid-sparing agents such as azathioprine and mycophenolate mofetil have been used in some patients; however, evidence for their efficacy is lacking.

For patients with relapsing or glucocorticoid-resistant disease, B cell depletion with rituximab is an excellent second-line therapy. Rituximab treatment (two doses of 1 g IV, separated by approximately 15 days) leads to a targeted, precipitous decline in serum IgG4 concentrations, suggesting that rituximab achieves its effects in part by preventing the repletion of short-lived plasma cells that produce IgG4. More important than its effects on IgG4 concentrations, however, may be the effect of B cell depletion on T cell function. Specific effects of rituximab on CD4+ effector T cells have been documented in IgG4-RD.

Rituximab may be an appropriate first-line therapy for some patients, particularly those at high risk for glucocorticoid toxicity and patients with immediately organ-threatening disease. The optimal approaches to remission maintenance, by either re-treatment with rituximab or continuous low-dose glucocorticoid therapy, require further study.

392 Familial Mediterranean Fever and Other Hereditary Autoinflammatory Diseases

Daniel L. Kastner

Familial Mediterranean fever (FMF) is the prototype of a group of inherited diseases (Table 392-1) that are characterized by recurrent episodes of fever with serosal, synovial, or cutaneous inflammation and, in some individuals, the eventual development of systemic AA amyloidosis (Chap. 137). Because of the relative infrequency of high-titer autoantibodies or antigen-specific T cells, the term *autoinflammatory* has been proposed to describe these disorders, rather than autoimmune. The innate immune system, with its myeloid effector cells and germline receptors for pathogen-associated molecular patterns and endogenous danger signals, plays a predominant role in the pathogenesis of the autoinflammatory diseases. Although the hereditary recurrent fevers comprise a major category of the autoinflammatory diseases, other inherited disorders of inflammation in which recurrent fever plays a less prominent role are now also considered to be autoinflammatory.

BACKGROUND AND PATHOPHYSIOLOGY

FMF was first recognized among Armenians, Arabs, Turks, and non-Ashkenazi (primarily North African and Iraqi) Jews. With the advent of genetic testing, FMF has been documented with increasing frequency among Ashkenazi Jews, Italians, and other Mediterranean populations, and occasional cases have been confirmed even in the absence of known Mediterranean ancestry. FMF is generally regarded as recessively inherited, but there is an increasing awareness of clear-cut clinical cases with only a single demonstrable genetic mutation, and, for certain relatively rare FMF mutations, there is strong evidence for dominant inheritance. Particularly in countries where families are small, a positive family history can only be elicited in ~50% of cases. DNA testing demonstrates carrier frequencies as high as 1:3 among affected populations, suggesting a heterozygote advantage.

The FMF gene encodes a 781-amino acid, ~95 kDa protein denoted *pyrin* (or *marenostrin*) that is expressed in granulocytes, eosinophils, monocytes, dendritic cells, and synovial and peritoneal fibroblasts. The N-terminal 92 amino acids of pyrin define a motif, the PYRIN domain, that is similar in structure to death domains, death effector

TABLE 392-1 THE HEREDITARY RECURRENT FEVER SYNDROMES

	FMF	TRAPS	HIDS	MWS	FCAS	NOMID
Ethnicity	Jewish, Arab, Turkish, Armenian, Italian	Any ethnic group	Predominantly Dutch, northern European	Any ethnic group	Any ethnic group	Any ethnic group
Inheritance	Recessive ^a	Dominant	Recessive	Dominant	Dominant	Most commonly de novo mutations; somatic mosaicism in a significant minority
Gene/chromosome	<i>MEFV</i> /16p13.3	<i>TNFRSF1A</i> /12p13	<i>MVK</i> /12q24	<i>NLRP3</i> /1q44	<i>NLRP3</i> /1q44	<i>NLRP3</i> /1q44
Protein	Pyrin	p55 TNF receptor	Mevalonate kinase	Cryopyrin (NLRP3)	Cryopyrin (NLRP3)	Cryopyrin (NLRP3)
Attack length	1–3 days	Often >7 days	3–7 days	1–2 days	Minutes–3 days	Continuous, with flares
Serosa	Pleurisy, peritonitis; asymptomatic pericardial effusions	Pleurisy, peritonitis, pericarditis	Abdominal pain, but seldom peritonitis; pleurisy, pericarditis uncommon	Abdominal pain; pleurisy, pericarditis rare	Rare	Rare
Skin	Erysipeloid erythema	Centrifugally migrating erythema	Diffuse maculopapular rash; oral ulcers	Diffuse urticaria-like rash	Cold-induced urticaria-like rash	Diffuse urticaria-like rash
Joints	Acute monoarthritis; chronic hip arthritis (rare)	Acute monoarthritis, arthralgia	Arthralgia, oligoarthritis	Arthralgia, large joint oligoarthritis	Polyarthralgia	Epiphyseal, patellar overgrowth, clubbing
Muscle	Exercise-induced myalgia common; protracted febrile myalgia rare	Migratory myalgia	Uncommon	Myalgia common	Sometimes myalgia	Sometimes myalgia
Eyes, ears	Uncommon	Periorbital edema, conjunctivitis, rarely uveitis	Uncommon	Conjunctivitis, episcleritis, optic disc edema; sensorineural hearing loss	Conjunctivitis	Conjunctivitis, uveitis, optic disc edema, blindness, sensorineural hearing loss
CNS	Aseptic meningitis rare	Headache	Headache	Headache	Headache	Aseptic meningitis, seizures
Amyloidosis	Most common in M694V homozygotes	~15% of cases, most often cysteine mutations, T50M	Uncommon	~25% of cases	Uncommon	Late complication
Treatment	Oral colchicine prophylaxis, IL-1 inhibitors for refractory cases	Glucocorticoids, etanercept, IL-1 inhibitors	NSAIDs for fever; IL-1 and TNF inhibitors	Anakinra, rilonacept, canakinumab	Anakinra, rilonacept, canakinumab	Anakinra

^aA substantial percentage of patients with clinical FMF have only a single demonstrable *MEFV* mutation on DNA sequencing.

Abbreviations: FCAS, familial cold autoinflammatory syndrome; FMF, familial Mediterranean fever; HIDS, hyperimmunoglobulinemia D with periodic fever syndrome; IL, interleukin; MWS, Muckle-Wells syndrome; NOMID, neonatal-onset multisystem inflammatory disease; NSAIDs, nonsteroidal anti-inflammatories; TNF, tumor necrosis factor; TRAPS, TNF receptor-associated periodic syndrome.

domains, and caspase recruitment domains. PYRIN domains mediate homotypic protein-protein interactions and have been found in several other proteins, including cryopyrin (NLRP3), which is mutated in three other recurrent fever syndromes. Through a number of mechanisms, including the interaction of the PYRIN domain with an intermediary adaptor protein, pyrin regulates caspase-1 (interleukin [IL] 1 β -converting enzyme), and thereby IL-1 β secretion. Mice bearing FMF-associated pyrin mutations exhibit inflammation and excessive IL-1 production.

ACUTE ATTACKS

Febrile episodes in FMF may begin even in early infancy; 90% of patients have had their first attack by age 20. Typical FMF episodes generally last 24–72 h, with arthritic attacks tending to last somewhat longer. In some patients, the episodes occur with great regularity, but more often, the frequency of attacks varies over time, ranging from as often as once every few days to remissions lasting several years. Attacks are often unpredictable, although some patients relate them to physical exertion, emotional stress, or menses; pregnancy may be associated with remission.

If measured, fever is nearly always present throughout FMF attacks. Severe hyperpyrexia and even febrile seizures may be seen in infants, and fever is sometimes the only manifestation of FMF in young children.

Over 90% of FMF patients experience abdominal attacks at some time. Episodes range in severity from dull, aching pain and distention with mild tenderness on direct palpation to severe generalized pain with absent bowel sounds, rigidity, rebound tenderness, and air-fluid levels on upright radiographs. Computed tomography (CT) scanning may demonstrate a small amount of fluid in the abdominal cavity. If such patients undergo exploratory laparotomy, a sterile, neutrophil-rich peritoneal exudate is present, sometimes with adhesions from previous episodes. Ascites is rare.

Pleural attacks are usually manifested by unilateral, sharp, stabbing chest pain. Radiographs may show atelectasis and sometimes an effusion. If performed, thoracentesis demonstrates an exudative fluid rich in neutrophils. After repeated attacks, pleural thickening may develop.

FMF arthritis is most frequent among individuals homozygous for the M694V mutation, which is especially common in the non-Ashkenazi Jewish population. Acute arthritis in FMF is usually monoarticular, affecting the knee, ankle, or hip, although other patterns can be seen, particularly in children. Large sterile effusions rich in neutrophils are frequent, without commensurate erythema or warmth. Even after repeated arthritic attacks, radiographic changes are rare. Before the advent of colchicine prophylaxis, chronic arthritis of the knee or hip was seen in ~5% of FMF patients with arthritis. Chronic sacroiliitis can occur in FMF irrespective of the HLA-B27 antigen, even in the face of

2214 colchicine therapy. In the United States, FMF patients are much more likely to have arthralgia than arthritis.

The most characteristic cutaneous manifestation of FMF is erysipelas-like erythema, a raised erythematous rash that most commonly occurs on the dorsum of the foot, ankle, or lower leg alone or in combination with abdominal pain, pleurisy, or arthritis. Biopsy demonstrates perivascular infiltrates of granulocytes and monocytes. This rash is seen most often in M694V homozygotes and is relatively rare in the United States.

Exercise-induced (nonfebrile) myalgia is common in FMF, and a small percentage of patients develop a protracted febrile myalgia that can last several weeks. Symptomatic pericardial disease is rare, although some patients have small pericardial effusions as an incidental echocardiographic finding. Unilateral acute scrotal inflammation may occur in prepubertal boys. Aseptic meningitis has been reported in FMF, but the causal connection is controversial. Vasculitis, including Henoch-Schönlein purpura and polyarteritis nodosa (**Chap. 385**), may be seen at increased frequency in FMF. The M694V FMF mutation has recently been shown to be a risk factor for Behçet's disease.

Laboratory features of FMF attacks are consistent with acute inflammation and include an elevated erythrocyte sedimentation rate, leukocytosis, thrombocytosis (in children), and elevations in C-reactive protein, fibrinogen, haptoglobin, and serum immunoglobulins. Transient albuminuria and hematuria may also be seen.

AMYLOIDOSIS

Before the advent of colchicine prophylaxis, systemic amyloidosis was a common complication of FMF. It is caused by deposition of a fragment of serum amyloid A, an acute-phase reactant, in the kidneys, adrenals, intestine, spleen, lung, and testes (**Chap. 137**). Amyloidosis should be suspected in patients who have proteinuria between attacks; renal or rectal biopsy is used most often to establish the diagnosis. Risk factors include the M694V homozygous genotype, positive family history (independent of FMF mutational status), the SAA 1 genotype, male gender, noncompliance with colchicine therapy, and having grown up in the Middle East.

DIAGNOSIS

For typical cases, physicians experienced with FMF can often make the diagnosis on clinical grounds alone. Clinical criteria sets for FMF have been shown to have high sensitivity and specificity in parts of the world where the pretest probability of FMF is high. Genetic testing can provide a useful adjunct in ambiguous cases or for physicians not experienced in FMF. Most of the more severe disease-associated FMF mutations are in exon 10 of the gene, with a smaller group of milder variants in exon 2. An updated list of mutations for FMF and other hereditary recurrent fevers can be found online at <http://fmf.igh.cnrs.fr/infevers/>.

Genetic testing has permitted a broadening of the clinical spectrum and geographic distribution of FMF and may be of prognostic value. Most studies indicate that M694V homozygotes have an earlier age of onset and a higher frequency of arthritis, rash, and amyloidosis. In contrast, the E148Q variant is quite common in certain populations and is more likely to affect overall levels of inflammation than to cause clinical FMF. E148Q is sometimes found in *cis* with exon 10 mutations, which may complicate the interpretation of genetic test results. Only ~70% of patients with clinically typical FMF have two identifiable mutations in *trans*. The inability to identify a second mutation even after intensive molecular analysis suggests that one FMF mutation may be sufficient to cause disease under some circumstances. In these cases clinical judgment is very important, and sometimes a therapeutic trial of colchicine may help to confirm the diagnosis. Genetic testing of unaffected individuals is usually inadvisable, because of the possibility of nonpenetrance and the potential impact of a positive test on future insurability.

If a patient is seen during his or her first attack, the differential diagnosis may be broad, although delimited by the specific organ involvement. After several attacks the differential diagnosis may include the other hereditary recurrent fever syndromes (Table 392-1); the syndrome of periodic fever with aphthous ulcers, pharyngitis, and cervical adenopathy (PFAPA); systemic-onset juvenile rheumatoid arthritis or adult Still's disease; porphyria; hereditary angioedema; inflammatory bowel disease; and, in women, gynecologic disorders.

TREATMENT FAMILIAL MEDITERRANEAN FEVER

The treatment of choice for FMF is daily oral colchicine, which decreases the frequency and intensity of attacks and prevents the development of amyloidosis in compliant patients. Intermittent dosing at the onset of attacks is not as effective as daily prophylaxis and is of unproven value in preventing amyloidosis. The usual adult dose of colchicine is 1.2–1.8 mg/d, which causes substantial reduction in symptoms in two-thirds of patients and some improvement in >90%. Children may require lower doses, although not proportionately to body weight.

Common side effects of colchicine include bloating, abdominal cramps, lactose intolerance, and diarrhea. They can be minimized by starting at a low dose and gradually advancing as tolerated, splitting the dose, use of simethicone for flatulence, and avoidance of dairy products. If taken by either parent at the time of conception, colchicine may cause a small increase in the risk of trisomy 21 (Down's syndrome). In elderly patients with renal insufficiency, colchicine can cause a myoneuropathy characterized by proximal muscle weakness and elevation of the creatine kinase. Cyclosporine inhibits hepatic excretion of colchicine by its effects on the MDR-1 transport system, sometimes leading to colchicine toxicity in patients who have undergone renal transplantation for amyloidosis. Intravenous colchicine should generally not be administered to patients already taking oral colchicine, because severe, sometimes fatal, toxicity can occur in this setting.

For FMF patients who do not respond to colchicine or cannot tolerate therapeutic doses, injectable IL-1 inhibitors appear to be effective in preventing the acute attacks. In a small randomized placebo-controlled trial, weekly subcutaneous riloncept, a recombinant IL-1 receptor fusion protein, significantly reduced the frequency of attacks. There is also substantial anecdotal experience with daily subcutaneous anakinra, a recombinant IL-1 receptor antagonist, in preventing the acute attacks of FMF, and in some cases reducing established amyloid deposits. Canakinumab, a monoclonal antibody to IL-1 β , and tumor necrosis factor (TNF) inhibitors may also have a role in the treatment of colchicine-unresponsive or intolerant patients. Bone marrow transplantation has been suggested for refractory FMF, but the risk-benefit ratio is currently regarded as unacceptable.

OTHER HEREDITARY RECURRENT FEVERS

Within 5 years of the discovery of the FMF gene, three additional genes causing five other hereditary recurrent fever syndromes were identified, catalyzing a paradigm shift in diagnosis and treatment of these disorders.

TNF RECEPTOR-ASSOCIATED PERIODIC SYNDROME (TRAPS)

TRAPS is caused by dominantly inherited mutations in the extracellular domains of the 55-kDa TNF receptor (TNFR1, p55). Although originally described in a large Irish family (and hence the name *familial Hibernian fever*), TRAPS has a broad ethnic distribution. TRAPS episodes often begin in childhood. The duration of attacks ranges from 1–2 days to as long as several weeks, and in severe cases symptoms may be nearly continuous. In addition to peritoneal, pleural, and synovial attacks similar to FMF, TRAPS patients frequently have ocular

inflammation (most often conjunctivitis and/or periorbital edema), and a distinctive migratory myalgia with overlying painful erythema may be present. TRAPS patients generally respond better to glucocorticoids than to prophylactic colchicine. Untreated, about 15% develop amyloidosis. The diagnosis of TRAPS is based on the demonstration of *TNFRSF1A* mutations in the presence of characteristic symptoms. Two particular variants, R92Q and P46L, are common in certain populations and may act more as functional polymorphisms than as disease-causing mutations. In contrast, pathogenic *TNFRSF1A* mutations, including a number of substitutions at highly conserved cysteine residues, are associated with intracellular TNFR1 misfolding, aggregation, and retention, with consequent ligand-independent kinase activation, mitochondrial reactive oxygen species production, and pro-inflammatory cytokine release. Etanercept, a TNF inhibitor, ameliorates TRAPS attacks, but the long-term experience with this agent has been less favorable. Perhaps because of the ligand-independent signaling abnormalities in TRAPS, IL-1 inhibition has been beneficial in a large percentage of the patients in whom it has been used. Monoclonal anti-TNF antibodies should be avoided, because they may exacerbate TRAPS attacks.

HYPERIMMUNOGLOBULINEMIA D WITH PERIODIC FEVER SYNDROME (HIDS)

HIDS is a recessively inherited recurrent fever syndrome found primarily in individuals of northern European ancestry. It is caused by mutations in mevalonate kinase (*MVK*), encoding an enzyme involved in the synthesis of cholesterol and nonsterol isoprenoids. Attacks usually begin in infancy and last 3–5 days. Clinically distinctive features include painful cervical adenopathy, a diffuse maculopapular rash sometimes affecting the palms and soles, and aphthous ulcers; pleurisy is rare, as is amyloidosis. Although originally defined by the persistent elevation of serum IgD, disease activity is not related to IgD levels, and some patients with FMF or TRAPS may have modestly increased serum IgD. Moreover, occasional patients with *MVK* mutations and recurrent fever have normal IgD levels. For these reasons, some have proposed renaming this disorder *mevalonate kinase deficiency (MKD)*. All patients with mutations have markedly elevated urinary mevalonate levels during their febrile attacks, although the inflammatory manifestations are likely to be due to a deficiency of isoprenoids rather than an excess of mevalonate. There is currently no established treatment for HIDS/MKD, although intermittent or continuous IL-1 inhibition and TNF inhibitors have been effective in small series.

THE CRYOPYRINOPATHIES, OR CRYOPYRIN-ASSOCIATED PERIODIC SYNDROMES (CAPS)

Three hereditary febrile syndromes, familial cold autoinflammatory syndrome (FCAS), Muckle-Wells syndrome (MWS), and neonatal-onset multisystem inflammatory disease (NOMID), are all caused by mutations in *NLRP3* (formerly known as *CIAS1*), the gene encoding cryopyrin (or NLRP3), and represent a clinical spectrum of disease. FCAS patients develop chills, fever, headache, arthralgia, conjunctivitis, and an urticaria-like rash in response to generalized cold exposure. In MWS, an urticarial rash is noted, but it is not usually induced by cold; MWS patients also develop fevers, abdominal pain, limb pain, arthritis, conjunctivitis, and, over time, sensorineural hearing loss. NOMID is the most severe of the three disorders, with chronic aseptic meningitis, a characteristic arthropathy, and rash. Like the FMF protein, pyrin, cryopyrin has an N-terminal PYRIN domain. Cryopyrin regulates

IL-1 β production through the formation of a macromolecular complex termed the *inflammasome*. Peripheral blood leukocytes from patients with FCAS, MWS, and NOMID release increased amounts of IL-1 β upon in vitro stimulation, relative to healthy controls. Macrophages from cryopyrin-deficient mice exhibit decreased IL-1 β production in response to certain gram-positive bacteria, bacterial RNA, and monosodium urate crystals. Patients with all three cryopyrinopathies show a dramatic response to injections of IL-1 inhibitors. Approximately one-third of patients with clinical manifestations of NOMID do not have germline mutations in *NLRP3*, but have been found to be mosaic for somatic *NLRP3* mutations. Such patients also respond dramatically to IL-1 inhibition. Similarly, somatic mosaicism in *NLRP3* has been found in Schnitzler's syndrome, which presents in middle age with recurrent fever, urticarial rash, elevated acute phase reactants, monoclonal IgM gammopathy, and abnormal bone remodeling. IL-1 inhibition is the treatment of choice for Schnitzler's syndrome.

OTHER INHERITED AUTOINFLAMMATORY DISEASES

There are a number of other Mendelian autoinflammatory diseases in which recurrent fevers are not a prominent clinical sign but that involve abnormalities of innate immunity. The syndrome of pyogenic arthritis with pyoderma gangrenosum and acne (PAPA) is a dominantly inherited disorder that presents with episodes of sterile pyogenic monoarthritis often induced by trauma, severe pyoderma gangrenosum, and severe cystic acne, usually beginning in puberty. It is caused by mutations in *PSTPIP1*, which encodes a pyrin-binding protein, and the arthritic manifestations often respond to IL-1 inhibition. Patients with the recessively inherited deficiency of the IL-1 receptor antagonist (DIRA) present with a generalized pustular rash and multifocal sterile osteomyelitis, and show dramatic clinical responses to anakinra, the recombinant form of the protein they lack. IL-36 is another member of the IL-1 family of cytokines that is regulated by an endogenous receptor antagonist. The recessively inherited deficiency of the IL-36 receptor antagonist (DITRA) presents with episodes of generalized pustular psoriasis and dramatic systemic inflammation.

Whereas PAPA, DIRA, and DITRA all involve mutations in IL-1-related molecules, other autoinflammatory diseases are caused by mutations in other components of innate immunity. *Blau's syndrome* is caused by mutations in *CARD15* (also known as *NOD2*), which regulates nuclear factor- κ B activation. *Blau's syndrome* is characterized by granulomatous dermatitis, uveitis, and arthritis; distinct *CARD15* variants predispose to Crohn's disease. Recessive mutations in one or more components of the proteasome lead to excessive interferon signaling and the syndrome of chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature (CANDLE), a severe form of generalized panniculitis. De novo gain-of-function mutations in *TMEM173*, encoding the stimulator of interferon genes (STING), cause severe vasculopathy and pulmonary fibrosis. Recessive loss-of-function mutations in *CERCR1*, encoding adenosine deaminase 2 (ADA2), cause a vasculopathy that can manifest as livedoid rash, early-onset lacunar strokes, or polyarteritis nodosa.

Finally, it should be noted that a number of common, genetically complex disorders are now sometimes considered autoinflammatory, because of evidence that components of the innate immune system, such as the inflammasome, may play a role in the pathogenesis. Two prominent examples are gout and atherosclerosis. Large clinical trials of IL-1 inhibitors have been initiated in both conditions.

393 Approach to Articular and Musculoskeletal Disorders

John J. Cush

Musculoskeletal complaints account for >315 million outpatient visits per year and over 20% of all outpatient visits in the United States. The Centers for Disease Control and Prevention estimate that 22.7% (52.5 million) of the U.S. population has physician-diagnosed arthritis and 22 million have significant functional limitation. While many patients will have self-limited conditions requiring minimal evaluation and only symptomatic therapy and reassurance, specific musculoskeletal presentations or their persistence may herald a more serious condition that requires further evaluation or laboratory testing to establish a diagnosis. The goal of the musculoskeletal evaluation is to formulate a differential diagnosis that leads to an accurate diagnosis and timely therapy, while avoiding excessive diagnostic testing and unnecessary treatment (Table 393-1). There are several urgent conditions that must be diagnosed promptly to avoid significant morbid or mortal sequelae. These “red flag” diagnoses include septic arthritis, acute crystal-induced arthritis (e.g., gout), and fracture. Each may be suspected by its acute onset and monarticular or focal musculoskeletal pain (see below).

Despite well-known links between certain disorders and laboratory testing, the majority of individuals with musculoskeletal complaints can be diagnosed with a thorough history and a comprehensive physical and musculoskeletal examination. The initial encounter should determine whether the musculoskeletal complaint signals a red flag condition (septic arthritis, gout, or fracture) or not. The evaluation should proceed to ascertain if the complaint is (1) *articular* or *non-articular* in origin, (2) *inflammatory* or *noninflammatory* in nature, (3) *acute* or *chronic* in duration, and (4) *localized (monarticular)* or *widespread (polyarticular)* in distribution.

With such an approach and an understanding of the pathophysiologic processes, the musculoskeletal complaint or presentation can be characterized (e.g., acute inflammatory monoarthritis or a chronic noninflammatory, nonarticular widespread pain) to narrow the diagnostic possibilities. A diagnosis can be made in the vast majority of individuals. However, some patients will not fit immediately into an established diagnostic category. Many musculoskeletal disorders resemble each other at the outset, and some may take weeks or months (but not years) to evolve into a recognizable diagnostic entity. This consideration should temper the desire to establish a definitive diagnosis at the first encounter.

TABLE 393-1 EVALUATION OF PATIENTS WITH MUSCULOSKELETAL COMPLAINTS

Goals
Accurate diagnosis
Timely provision of therapy
Avoidance of unnecessary diagnostic testing
Identification of acute, focal/monarticular “red flag” conditions
Approach
Determination of chronology (acute vs chronic)
Determination of the nature of the pathologic process (inflammatory vs noninflammatory)
Determination of the extent of involvement (monarticular, polyarticular, focal, widespread)
Anatomic localization of complaint (articular vs nonarticular)
Consider the most common disorders first
Formulate a differential diagnosis

ARTICULAR VERSUS NONARTICULAR

The musculoskeletal evaluation must discriminate the anatomic origin(s) of the patient’s complaint. For example, ankle pain can result from a variety of pathologic conditions involving disparate anatomic structures, including gonococcal arthritis, calcaneal fracture, Achilles tendinitis, plantar fasciitis, cellulitis, and peripheral or entrapment neuropathy. Distinguishing between articular and nonarticular conditions requires a careful and detailed examination. Articular structures include the synovium, synovial fluid, articular cartilage, intraarticular ligaments, joint capsule, and juxtaarticular bone. Nonarticular (or periarticular) structures, such as supportive extraarticular ligaments, tendons, bursae, muscle, fascia, bone, nerve, and overlying skin, may be involved in the pathologic process. Although musculoskeletal complaints are often ascribed to the joints, nonarticular disorders more frequently underlie such complaints. Distinguishing between these potential sources of pain may be challenging to the unskilled examiner. Articular disorders may be characterized by deep or diffuse pain, pain or limited range of motion on active and passive movement, and swelling (caused by synovial proliferation, effusion, or bony enlargement), crepitation, instability, “locking,” or deformity. By contrast, nonarticular disorders tend to be painful on active, but not passive (or assisted), range of motion. Periarticular conditions often demonstrate point or focal tenderness in regions adjacent to articular structures, are elicited with a specific movement or position, and have physical findings remote from the joint capsule. Moreover, nonarticular disorders seldom demonstrate swelling, crepitus, instability, or deformity of the joint itself.

INFLAMMATORY VERSUS NONINFLAMMATORY DISORDERS

In the course of a musculoskeletal evaluation, the examiner should determine the nature of the underlying pathologic process and whether inflammatory or noninflammatory findings exist. Inflammatory disorders may be infectious (*Neisseria gonorrhoeae* or *Mycobacterium tuberculosis*), crystal-induced (gout, pseudogout), immune-related (rheumatoid arthritis [RA], systemic lupus erythematosus [SLE]), reactive (rheumatic fever, reactive arthritis), or idiopathic. Inflammatory disorders may be identified by any of the four cardinal signs of inflammation (erythema, warmth, pain, or swelling), systemic symptoms (fatigue, fever, rash, weight loss), or laboratory evidence of inflammation (elevated erythrocyte sedimentation rate [ESR] or C-reactive protein [CRP], thrombocytosis, anemia of chronic disease, or hypoalbuminemia). Articular stiffness commonly accompanies chronic musculoskeletal disorders. The duration of stiffness may be prolonged (hours) with inflammatory disorders (such as RA or polymyalgia rheumatica) and may improve with activity. By contrast, intermittent stiffness (also known as gel phenomenon) is typical of noninflammatory conditions (such as osteoarthritis [OA]), shorter in duration (<60 min), and exacerbated by activity. Fatigue may accompany inflammation (as seen in RA and polymyalgia rheumatica) but may also be a consequence of fibromyalgia (a noninflammatory disorder), chronic pain, poor sleep, depression, anemia, cardiac failure, endocrinopathy, or malnutrition. Noninflammatory disorders may be related to trauma (rotator cuff tear), repetitive use (bursitis, tendinitis), degeneration or ineffective repair (OA), neoplasm (pigmented villonodular synovitis), or pain amplification (fibromyalgia). Noninflammatory disorders are often characterized by pain without synovial swelling or warmth, absence of inflammatory or systemic features, daytime gel phenomena rather than morning stiffness, and normal (for age) or negative laboratory investigations.

Identification of the nature of the underlying process and the site of the complaint will enable the examiner to characterize the musculoskeletal presentation (e.g., acute inflammatory monoarthritis, chronic noninflammatory, nonarticular widespread pain), narrow the diagnostic considerations, and assess the need for immediate diagnostic or

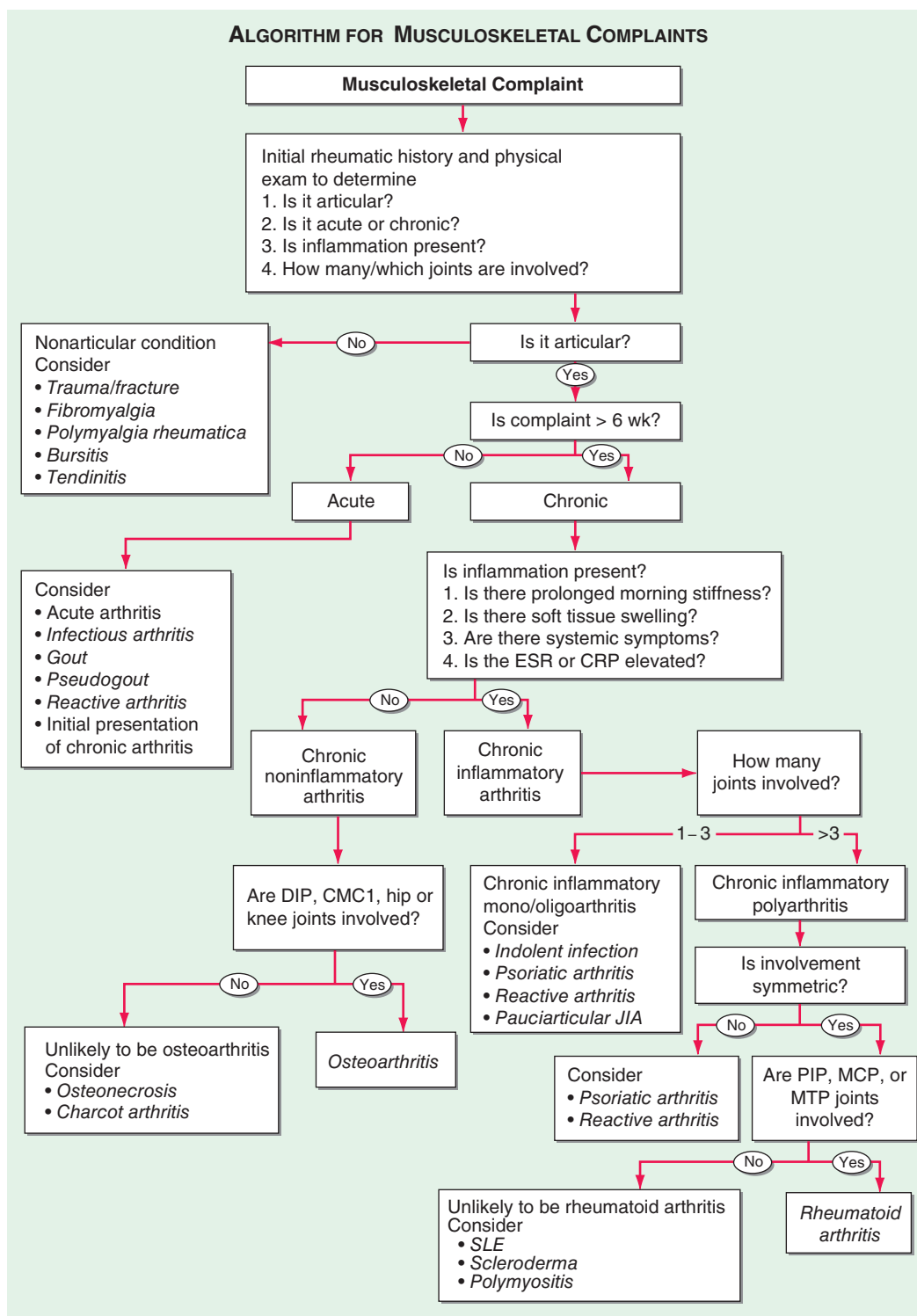


FIGURE 393-1 Algorithm for the diagnosis of musculoskeletal complaints. An approach to formulating a differential diagnosis (shown in italics). CMC, carpometacarpal; CRP, C-reactive protein; DIP, distal interphalangeal; ESR, erythrocyte sedimentation rate; JIA, juvenile idiopathic arthritis; MCP, metacarpophalangeal; MTP, metatarsophalangeal; PIP, proximal interphalangeal; PMR, polymyalgia rheumatica; SLE, systemic lupus erythematosus.

therapeutic intervention or for continued observation. **Figure 393-1** presents an algorithmic approach to the evaluation of patients with musculoskeletal complaints. This approach relies on clinical and historic features, rather than laboratory testing, to diagnose many common rheumatic disorders.

A simpler, alternative approach would consider the most commonly encountered complaints first, based on frequency in younger versus older populations. The most prevalent causes of musculoskeletal complaints are shown in **Fig. 393-2**. Because trauma, fracture, overuse

syndromes, and fibromyalgia are among the most common causes of joint pain, these should be considered during the initial encounter. If these possibilities are excluded, other frequently occurring disorders should be considered according to the patient's age. Hence, those younger than 60 years are commonly affected by repetitive use/strain disorders, gout (men only), RA, spondyloarthritis, and uncommonly, infectious arthritis. Patients over age 60 years are frequently affected by OA, crystal (gout and pseudogout) arthritis, polymyalgia rheumatica, osteoporotic fracture, and uncommonly, septic arthritis.

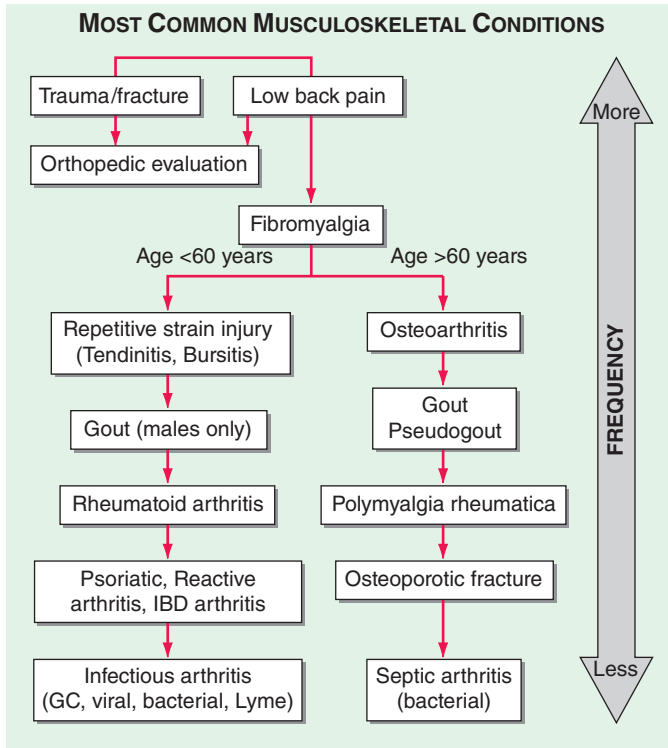


FIGURE 393-2 Algorithm for consideration of the most common musculoskeletal conditions. GC, gonococcal; IBD, inflammatory bowel disease.

These conditions are between 10 and 100 times more prevalent than other serious autoimmune conditions, such as SLE, scleroderma, polymyositis, and vasculitis.

CLINICAL HISTORY

Additional historic features may reveal important clues to the diagnosis. Aspects of the patient profile, complaint chronology, extent of joint involvement, and precipitating factors can provide important information. Certain diagnoses are more frequent in different age groups. SLE and reactive arthritis occur more frequently in the young, whereas fibromyalgia and RA are frequent in middle age, and OA and polymyalgia rheumatica are more prevalent among the elderly. Diagnostic clustering is also evident when *sex* and *race* are considered. Gout, spondyloarthritis, and ankylosing spondylitis are more common in men, whereas RA, fibromyalgia, and lupus are more frequent in women. *Racial predilections* may be evident. Thus, polymyalgia rheumatica, giant cell arteritis, and granulomatosis with polyangiitis (GPA; formerly called Wegener's granulomatosis) commonly affect whites, whereas sarcoidosis and SLE more commonly affect African Americans. *Familial aggregation* is most common with ankylosing spondylitis, gout, and Heberden's nodes of OA.

The chronology of the complaint is an important diagnostic feature and can be divided into the *onset*, *evolution*, and *duration*. The onset of disorders such as septic arthritis or gout tends to be abrupt, whereas OA, RA, and fibromyalgia may have more indolent presentations. The patients' complaints may evolve differently and be classified as chronic (OA), intermittent (crystal or Lyme arthritis), migratory (rheumatic fever, gonococcal or viral arthritis), or additive (RA, psoriatic arthritis). Musculoskeletal disorders are typically classified as acute or chronic based on a symptom duration that is either less than or greater than 6 weeks, respectively. Acute arthropathies tend to be infectious, crystal-induced, or reactive. Chronic conditions include noninflammatory or immunologic arthritides (e.g., OA, RA) and nonarticular disorders (e.g., fibromyalgia).

The *extent* or *distribution* of articular involvement is often informative. Articular disorders are classified based on the number of joints involved, as either *monarticular* (one joint), *oligoarticular* or

pauciarticular (two or three joints), or *polyarticular* (four or more joints). Although crystal and infectious arthritis are often mono- or oligoarticular, OA and RA are polyarticular disorders. Nonarticular disorders may be classified as either focal or widespread. Complaints secondary to tendinitis or carpal tunnel syndrome are typically focal, whereas weakness and myalgia, caused by polymyositis or fibromyalgia, are more widespread in their presentation. Joint involvement in RA tends to be symmetric and polyarticular. By contrast, spondyloarthritis, reactive arthritis, gout, and sarcoid are often asymmetric and oligo- or polyarticular. The upper extremities are frequently involved in RA and OA, whereas lower extremity arthritis is characteristic of reactive arthritis and gout at their onset. Involvement of the axial skeleton is common in OA and ankylosing spondylitis but is infrequent in RA, with the notable exception of the cervical spine.

The clinical history should also identify *precipitating events*, such as trauma (osteonecrosis, meniscal tear), drug administration (Table 393-2), antecedent or intercurrent infection (rheumatic fever, reactive arthritis, hepatitis), or illnesses that may have contributed to the patient's complaint. Certain comorbidities may have musculoskeletal consequences. This is especially so for diabetes mellitus (carpal tunnel syndrome), renal insufficiency (gout), depression or insomnia (fibromyalgia), myeloma (low back pain), cancer (myositis), and osteoporosis (fracture) or when using certain drugs such as glucocorticoids (osteonecrosis, septic arthritis) and diuretics or chemotherapy (gout) (Table 393-2).

Lastly, a thorough *rheumatic review of systems* may disclose useful diagnostic information. A variety of musculoskeletal disorders may be associated with systemic features such as fever (SLE, infection), rash (SLE, psoriatic arthritis), nail abnormalities (psoriatic or reactive arthritis), myalgias (fibromyalgia, statin- or drug-induced myopathy),

TABLE 393-2 DRUG-INDUCED MUSCULOSKELETAL CONDITIONS

Arthralgias

Quinidine, cimetidine, beta blockers, quinolones, chronic acyclovir, interferons, IL-2, nifedipine, vaccines, rifabutin, aromatase and HIV protease inhibitors

Myalgias/myopathy

Glucocorticoids, penicillamine, hydroxychloroquine, AZT, lovastatin, simvastatin, atorvastatin, pravastatin, clofibrate, amiodarone, interferon, IL-2, alcohol, cocaine, paclitaxel, docetaxel, imatinib mesylate, colchicine, quinolones, cyclosporine, tacrolimus, protease inhibitors

Tendon rupture/tendinitis

Quinolones, glucocorticoids, isotretinoin, statins, collagenase injections

Gout

Diuretics, aspirin, cytotoxics, cyclosporine, alcohol, moonshine, ethambutol, fructose-containing soft drinks

Drug-induced lupus

Hydralazine, procainamide, quinidine, phenytoin, carbamazepine, methyl-dopa, isoniazid, chlorpromazine, lithium, penicillamine, tetracyclines, TNF inhibitors, ACE inhibitors, ticlopidine

Drug-induced subacute lupus

Proton pump inhibitors, calcium channel blockers (diltiazem), ACE inhibitors, TNF inhibitors, terbinafine, interferons (α and β -1a), paclitaxel, docetaxel, HCTZ

Osteonecrosis

Glucocorticoids, alcohol, radiation, bisphosphonates

Osteopenia

Glucocorticoids, chronic heparin, phenytoin

Scleroderma

Vinyl chloride, bleomycin, baricitinib, pentazocine, organic solvents, carbido-pa, tryptophan, rapeseed oil

Vasculitis

Allopurinol, amphetamines, cocaine (often levamisole adulterated), thiazides, penicillamine, propylthiouracil, montelukast, TNF inhibitors, hepatitis B vaccine, trimethoprim/sulfamethoxazole, minocycline, hydralazine

Abbreviations: ACE, angiotensin-converting enzyme; AZT, zidovudine; HCTZ, hydrochlorothiazide; IL-2, interleukin 2; TNF, tumor necrosis factor.

or weakness (polymyositis, neuropathy). In addition, some conditions are associated with involvement of other organ systems including the eyes (Behçet's disease, sarcoidosis, spondyloarthritis), gastrointestinal tract (scleroderma, inflammatory bowel disease), genitourinary tract (reactive arthritis, gonococemia), or nervous system (Lyme disease, vasculitis).

RHEUMATOLOGIC EVALUATION OF THE ELDERLY

The incidence of rheumatic diseases rises with age, such that 58% of those >65 years will have joint complaints. Musculoskeletal disorders in elderly patients are often not diagnosed because the signs and symptoms may be insidious, overlooked, or overshadowed by comorbidities. These difficulties are compounded by the diminished reliability of laboratory testing in the elderly, who often manifest nonpathologic abnormal results. For example, the ESR may be misleadingly elevated, and low-titer positive tests for rheumatoid factor and antinuclear antibodies (ANAs) may be seen in up to 15% of elderly patients. Although nearly all rheumatic disorders afflict the elderly, geriatric patients are particularly prone to OA, osteoporosis, osteoporotic fractures, gout, pseudogout, polymyalgia rheumatica, vasculitis, and drug-induced disorders (Table 393-2). The elderly should be approached in the same manner as other patients with musculoskeletal complaints, but with an emphasis on identifying the potential rheumatic consequences of medical comorbidities and therapies. The physical examination should identify the nature of the musculoskeletal complaint as well as coexisting diseases that may influence diagnosis and choice of treatment.

RHEUMATOLOGIC EVALUATION OF THE HOSPITALIZED PATIENT

Evaluation of a hospitalized patient with rheumatic complaints differs from that of an outpatient, owing to greater symptom severity, more acute presentations, and greater interplay of comorbidities. Patients with rheumatic disorders tend to be admitted for one of several reasons: (1) acute onset of inflammatory arthritis; (2) undiagnosed systemic or febrile illness; (3) musculoskeletal trauma; (4) exacerbation or deterioration of an existing autoimmune disorder (e.g., SLE); or (5) new medical comorbidities (e.g., thrombotic event, lymphoma, infection) arising in patients with an established rheumatic disorder. Notably, rheumatic patients are seldom if ever admitted because of widespread pain or serologic abnormalities or for the initiation of new therapies.

Acute monoarticular inflammatory arthritis may be a “red flag” condition (e.g., septic arthritis, gout, pseudogout) that will require arthrocentesis and, on occasion, hospitalization if infection is suspected. However, new-onset inflammatory polyarthritis will have a wider differential diagnosis (e.g., RA, hepatitis-related arthritis, serum sickness, drug-induced lupus, polyarticular septic arthritis) and may require targeted laboratory investigations rather than synovial fluid analyses. Patients with febrile, multisystem disorders will require exclusion of crystal, infectious, or neoplastic etiologies and an evaluation driven by the dominant symptom/finding with the greatest specificity. Conditions worthy of consideration may include gout or pseudogout, vasculitis (giant cell arteritis in the elderly or polyarteritis nodosa in younger patients), adult-onset Still's disease, SLE, antiphospholipid antibody syndrome, and sarcoidosis. Because the misdiagnosis of connective tissue disorders is common, patients who present with a reported preexisting rheumatic condition (e.g., SLE, RA, ankylosing spondylitis) should have their diagnosis confirmed by careful history, physical and musculoskeletal examination, and review of their medical records. It is important to note that when established rheumatic disease patients are admitted to the hospital, it is usually not for a medical problem related to their autoimmune disease, but rather because of either a comorbid condition or complication of drug therapy. Patients with chronic inflammatory disorders (e.g., RA, SLE, psoriasis) have an augmented risk of infection, cardiovascular events, and neoplasia.

Certain conditions, such as acute gout, can be precipitated in hospitalized patients by surgery, dehydration, or other events and should be considered when hospitalized patients are evaluated for the acute onset

of a musculoskeletal condition. Lastly, overly aggressive and unfocused laboratory testing will often yield abnormal findings that are better explained by the patient's preexisting condition(s) rather than a new inflammatory or autoimmune disorder.

PHYSICAL EXAMINATION

The goal of the physical examination is to ascertain the structures involved, the nature of the underlying pathology, the functional consequences of the process, and the presence of systemic or extraarticular manifestations. A knowledge of topographic anatomy is necessary to identify the primary site(s) of involvement and differentiate articular from nonarticular disorders. The musculoskeletal examination depends largely on careful inspection, palpation, and a variety of specific physical maneuvers to elicit diagnostic signs (Table 393-3). Although most articulations of the appendicular skeleton can be examined in this manner, adequate inspection and palpation are not possible for many axial (e.g., zygapophyseal) and inaccessible (e.g., sacroiliac or hip) joints. For such joints, there is a greater reliance on specific maneuvers and imaging for assessment.

Examination of involved and uninvolved joints will determine whether *pain*, *warmth*, *erythema*, or *swelling* is present. The locale and level of pain elicited by palpation or movement should be quantified. One standard would be to count the number of tender joints on palpation of 28 easily examined joints (proximal interphalangeals, metacarpophalangeals, wrists, elbows, shoulders, and knees). Similarly, the number of swollen joints (0–28) can be counted and recorded. Careful examination should distinguish between true articular swelling (caused by bony hypertrophy, synovial effusion or proliferation) and nonarticular (or periarticular) involvement, which usually extends beyond the normal joint margins. Synovial effusion can be distinguished from synovial hypertrophy or bony hypertrophy by palpation or specific maneuvers. For example, small to moderate knee effusions may be identified by the “bulge sign” or “ballottement of the patellae.” Bursal effusions (e.g., effusions of the olecranon or prepatellar bursa) are often focal, periarticular, overlie bony prominences, and are fluctuant with sharply defined borders. Joint *stability* can be assessed by stabilizing the proximal joint, by palpation, and by the application of manual stress to the distal appendage. *Subluxation* or *dislocation*, which may be secondary to traumatic, mechanical, or inflammatory

TABLE 393-3 GLOSSARY OF MUSCULOSKELETAL TERMS

Crepitus

A palpable (less commonly audible) vibratory or crackling sensation elicited with joint motion; fine joint crepitus is common and often insignificant in large joints; coarse joint crepitus indicates advanced cartilaginous and degenerative changes (as in osteoarthritis)

Subluxation

Alteration of joint alignment such that articulating surfaces incompletely approximate each other

Dislocation

Abnormal displacement of articulating surfaces such that the surfaces are not in contact

Range of motion

For diarthrodial joints, the arc of measurable movement through which the joint moves in a single plane

Contracture

Loss of full movement resulting from a fixed resistance caused either by tonic spasm of muscle (reversible) or by fibrosis of periarticular structures (permanent)

Deformity

Abnormal shape or size resulting from bony hypertrophy, malalignment of articulating structures, or damage to periarticular supportive structures

Enthesitis

Inflammation of the entheses (tendinous or ligamentous insertions on bone)

Epicondylitis

Infection or inflammation involving an epicondyle

causes, can be assessed by inspection and palpation. Joint *swelling* or *volume* can be assessed by palpation. Distention of the articular capsule usually causes pain and evident enlargement or fluctuance. The patient will attempt to minimize the pain by maintaining the joint in the position of least intraarticular pressure and greatest volume, usually partial flexion. For this reason, inflammatory effusions may give rise to flexion contractures. Clinically, this may be detected as fluctuant or “squishy” swelling in larger joints and grape-like compressibility in smaller joints. Inflammation may result in fixed flexion deformities or diminished range of motion—especially on extension, when intraarticular pressure is increased. Active and passive *range of motion* should be assessed in all planes, with contralateral comparison. A goniometer may be used to quantify the arc of movement. Each joint should be passively manipulated through its full range of motion (including, as appropriate, flexion, extension, rotation, abduction, adduction, lateral bending, inversion, eversion, supination, pronation, medial/lateral deviation, and plantar- or dorsiflexion). Extreme range of motion may be seen with hypermobility syndrome, with joint pain and connective tissue laxity, often associated with Ehlers-Danlos or Marfan’s syndrome. Limitation of motion is frequently caused by inflammation, effusion, pain, deformity, contracture, or restriction from neuromyopathic causes. If passive motion exceeds active motion, a periarticular process (e.g., tendinitis, tendon rupture, or myopathy) should be considered. *Contractures* may reflect antecedent synovial inflammation or trauma. Minor joint *crepitus* is common during joint palpation and maneuvers but only indicates significant cartilage degeneration as it becomes coarser (e.g., OA). Joint *deformity* usually indicates a long-standing or aggressive pathologic process. Deformities may result from ligamentous destruction, soft tissue contracture, bony enlargement, ankylosis, erosive disease, subluxation, trauma, or loss of proprioception. Examination of the musculature will document strength, atrophy, pain, or spasm. Appendicular muscle weakness should be characterized as proximal or distal. Muscle strength should be assessed by observing the patient’s performance (e.g., walking, rising from a chair, grasping, writing). Strength may also be graded on a 5-point scale: 0 for no movement; 1 for trace movement or twitch; 2 for movement with gravity eliminated; 3 for movement against gravity only; 4 for movement against gravity and resistance; and 5 for normal strength. The examiner should assess for often-overlooked nonarticular or periarticular involvement, especially when articular complaints are not supported by objective findings referable to the joint capsule. The identification of soft tissue/nonarticular pain will prevent unwarranted and often expensive additional evaluations. Specific maneuvers may reveal common nonarticular abnormalities, such as a carpal tunnel syndrome (which can be identified by Tinel’s or Phalen’s sign). Other examples of soft tissue abnormalities include olecranon bursitis, epicondylitis (e.g., tennis elbow), enthesitis (e.g., Achilles tendinitis), and tender trigger points associated with fibromyalgia.

APPROACH TO REGIONAL RHEUMATIC COMPLAINTS

Although all patients should be evaluated in a logical and thorough manner, many cases with focal musculoskeletal complaints are caused by commonly encountered disorders that exhibit a predictable pattern of onset, evolution, and localization; they can often be diagnosed immediately on the basis of limited historic information and selected maneuvers or tests. Although nearly every joint could be approached in this manner, the evaluation of four common involved anatomic regions—the hand, shoulder, hip, and knee—are reviewed here.

HAND PAIN

Focal or unilateral hand pain may result from trauma, overuse, infection, or a reactive or crystal-induced arthritis. By contrast, bilateral hand complaints commonly suggest a degenerative (e.g., OA), systemic, or inflammatory/immune (e.g., RA) etiology. The distribution or pattern of joint involvement is highly suggestive of certain disorders (Fig. 393-3). Thus, OA (or degenerative arthritis) may manifest as distal interphalangeal (DIP) and proximal interphalangeal (PIP) joint pain with bony hypertrophy sufficient to produce Heberden’s and Bouchard’s nodes, respectively. Pain, with or without bony swelling,

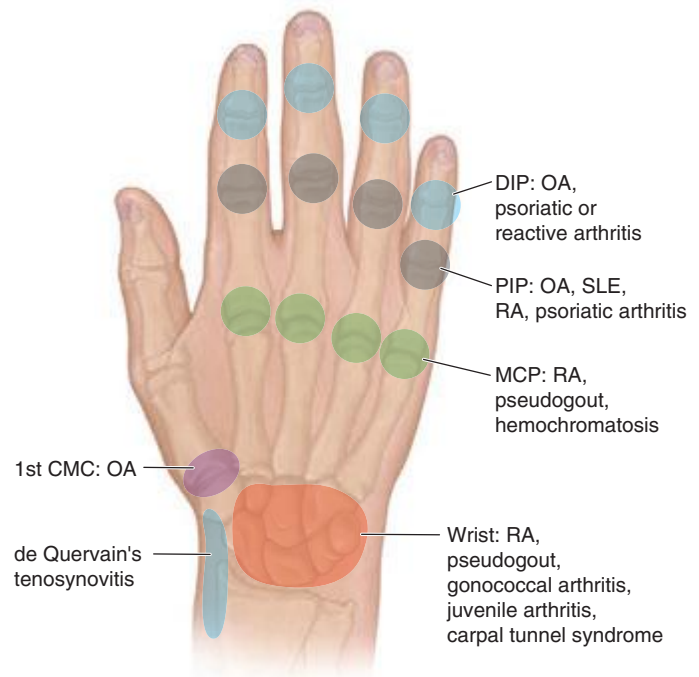


FIGURE 393-3 Sites of hand or wrist involvement and their potential disease associations. CMC, carpometacarpal; DIP, distal interphalangeal; MCP, metacarpophalangeal; OA, osteoarthritis; PIP, proximal interphalangeal; RA, rheumatoid arthritis; SLE, systemic lupus erythematosus. (From JJ Cush et al: *Evaluation of musculoskeletal complaints, in Rheumatology: Diagnosis and Therapeutics, 2nd ed, JJ Cush et al [eds]. Philadelphia, Lippincott Williams & Wilkins, 2005, pp 3–20. Used with permission from Dr. John J. Cush.*)

involving the base of the thumb (first carpometacarpal joint) is also highly suggestive of OA. By contrast, RA tends to cause symmetric, polyarticular involvement of the PIP, metacarpophalangeal (MCP), intercarpal, and carpometacarpal joints (wrist) with pain and palpable synovial tissue hypertrophy. Psoriatic arthritis may mimic the pattern of joint involvement seen in OA (DIP and PIP joints), but can be distinguished by the presence of inflammatory signs (erythema, warmth, synovial swelling), with or without carpal involvement, nail pitting, or onycholysis. Whereas lateral or medial subluxations at the PIP or DIP joints are most likely due to inflammatory OA or psoriatic arthritis, dorsal or ventral deformities (swan neck or boutonnière deformities) are typical of RA. Hemochromatosis should be considered when degenerative changes (bony hypertrophy) are seen at the second and third MCP joints with associated radiographic chondrocalcinosis or episodic, inflammatory wrist arthritis.

Dactylitis manifests as soft tissue swelling of the whole digit and may have a sausage-like appearance. Common causes of dactylitis include psoriatic arthritis, spondyloarthritis, juvenile spondylitis, mixed connective tissue disease, scleroderma, sarcoidosis, and sickle cell disease. Soft tissue swelling over the dorsum of the hand and wrist may suggest an inflammatory extensor tendon tenosynovitis possibly caused by gonococcal infection, gout, or inflammatory arthritis (e.g., RA). Tenosynovitis is suggested by localized warmth, swelling, or pitting edema and may be confirmed when the soft tissue swelling tracks with tendon movement, such as flexion and extension of fingers, or when pain is induced while stretching the extensor tendon sheaths (flexing the digits distal to the MCP joints and maintaining the wrist in a fixed, neutral position).

Focal wrist pain localized to the radial aspect may be caused by de Quervain’s tenosynovitis resulting from inflammation of the tendon sheath(s) involving the abductor pollicis longus or extensor pollicis brevis (Fig. 393-3). This commonly results from overuse or follows pregnancy and may be diagnosed with Finkelstein’s test. A positive

result is present when radial wrist pain is induced after the thumb is flexed and placed inside a clenched fist and the patient actively deviates the hand downward with ulnar deviation at the wrist. Carpal tunnel syndrome is another common disorder of the upper extremity and results from compression of the median nerve within the carpal tunnel. Manifestations include pain in the wrist that may radiate with paresthesia to the thumb, second and third fingers, and radial half of the fourth finger and, at times, atrophy of thenar musculature. Carpal tunnel syndrome is commonly associated with pregnancy, edema, trauma, OA, inflammatory arthritis, and infiltrative disorders (e.g., amyloidosis). The diagnosis may be suggested by a positive Tinel's or Phalen's sign. With each test, paresthesia in a median nerve distribution is induced or increased by either "thumping" the volar aspect of the wrist (Tinel's sign) or pressing the extensor surfaces of both flexed wrists against each other (Phalen's sign). The low sensitivity and moderate specificity of these tests may require nerve conduction velocity testing to confirm a suspected diagnosis.

SHOULDER PAIN

During the evaluation of shoulder disorders, the examiner should carefully note any history of trauma, fibromyalgia, infection, inflammatory disease, occupational hazards, or previous cervical disease. In addition, the patient should be questioned as to the activities or movement(s) that elicit shoulder pain. While arthritis is suggested by pain on movement in all planes, pain with specific active motion suggests a periarticular (nonarticular) process. Shoulder pain may originate in the glenohumeral or acromioclavicular joints, subacromial (subdeltoid) bursa, periarticular soft tissues (e.g., fibromyalgia, rotator cuff tear/tendinitis), or cervical spine (Fig. 393-4). Shoulder pain is referred frequently from the cervical spine but may also be referred from intrathoracic lesions (e.g., a Pancoast tumor) or from gallbladder, hepatic, or diaphragmatic disease. These same visceral causes may also manifest as focal scapular pain. Fibromyalgia should be suspected

when glenohumeral pain is accompanied by diffuse periarticular (i.e., subacromial, bicipital) pain and tender points (i.e., trapezius or supraspinatus). The shoulder should be put through its full range of motion both actively and passively (with examiner assistance): forward flexion, extension, abduction, adduction, and internal and external rotation. Manual inspection of the periarticular structures will often provide important diagnostic information. Glenohumeral involvement is best detected by placing the thumb over the glenohumeral joint just medial and inferior to the coracoid process and applying pressure anteriorly while internally and externally rotating the humeral head. Pain localized to this region is indicative of glenohumeral pathology. Synovial effusion or tissue is seldom palpable but, if present, may suggest infection, RA, amyloidosis, or an acute tear of the rotator cuff. The examiner should apply direct manual pressure over the subacromial bursa that lies lateral to and immediately beneath the acromion (Fig. 393-4). Subacromial bursitis is a frequent cause of shoulder pain. Anterior to the subacromial bursa, the bicipital tendon traverses the bicipital groove. This tendon is best identified by palpating it in its groove as the patient rotates the humerus internally and externally. Direct pressure over the tendon may reveal pain indicative of bicipital tendinitis. Palpation of the acromioclavicular joint may disclose local pain, bony hypertrophy, or, uncommonly, synovial swelling. Whereas OA and RA commonly affect the acromioclavicular joint, OA seldom involves the glenohumeral joint, unless there is a traumatic or occupational cause.

Rotator cuff tendinitis or tear is a very common cause of shoulder pain. Nearly 30 percent of the elderly will have shoulder pain, with rotator cuff tendinitis or tear as the primary cause. The rotator cuff is formed by four tendons that attach the scapula to the proximal humerus (supraspinatus, infraspinatus, teres minor, and subscapularis tendons). Of these, the supraspinatus muscle is the most commonly damaged. Rotator cuff tendinitis is suggested by pain on active abduction (but not passive abduction), pain over the lateral deltoid muscle, night pain, and evidence of the impingement signs (pain with overhead arm activities). The Neer test for impingement is performed by the examiner raising the patient's arm into forced flexion while stabilizing and preventing rotation of the scapula. A positive sign is present if pain develops before 180° of forward flexion. Tear of the rotator cuff is common in the elderly and often results from trauma; it may manifest in the same manner as tendinitis. The drop arm test is abnormal with supraspinatus pathology and is demonstrated by passive abduction of the arm to 90° by the examiner. If the patient is unable to hold the arm up actively or unable to lower the arm slowly without dropping, the test is positive. Tendinitis or tear of the rotator cuff is best confirmed by magnetic resonance imaging (MRI) or ultrasound.

KNEE PAIN

Knee pain may result from intraarticular (OA, RA) or periarticular (anserine bursitis, collateral ligament strain) processes or be referred from hip pathology. A careful history should delineate the chronology of the knee complaint and whether there are predisposing conditions, trauma, or medications that might underlie the complaint. For example, patellofemoral disease (e.g., OA) may cause anterior knee pain that worsens with climbing stairs. Observation of the patient's gait is also important. The knee should be carefully inspected in the upright (weight-bearing) and supine positions for swelling, erythema, malalignment, visible trauma, muscle wasting, and leg length discrepancy. The most common malalignment in the knee is *genu varum* (bowlegs) or *genu valgum* (knock-knees) resulting from asymmetric cartilage loss medially or laterally. Bony swelling of the knee joint commonly results from hypertrophic osseous changes seen with disorders such as OA and neuropathic arthropathy. Swelling caused by hypertrophy of the synovium or synovial effusion may manifest as a fluctuant, ballotable, or soft tissue enlargement in the suprapatellar pouch (suprapatellar reflection of the synovial cavity) or regions lateral and medial to the patella. Synovial effusions may also be detected by balloting the patella downward toward the femoral groove or by eliciting a "bulge sign." With the knee extended, the examiner should manually compress, or "milk," synovial fluid down from the suprapatellar pouch and lateral to the patellae. The

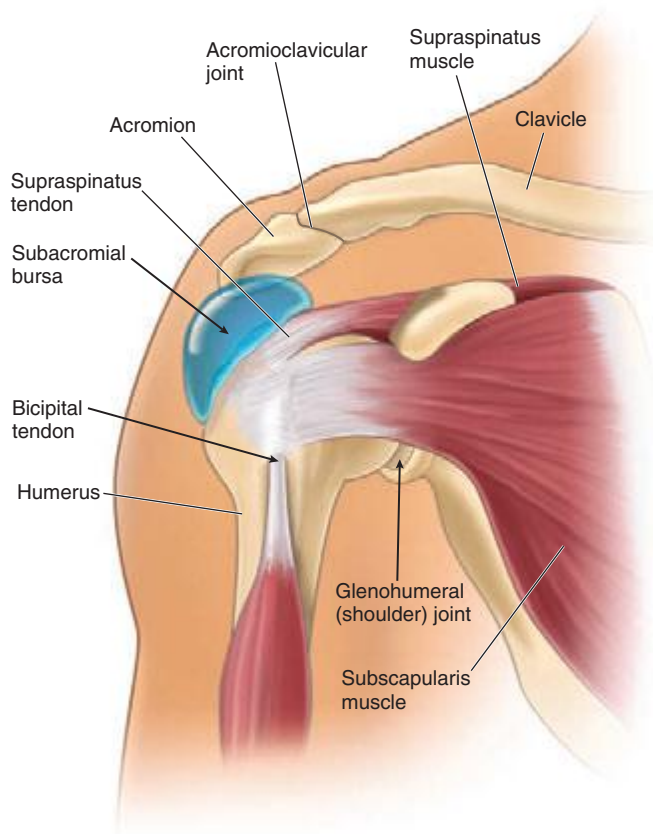


FIGURE 393-4 Origins of shoulder pain. The schematic diagram of the shoulder indicates with arrows the anatomic origins of shoulder pain.

2222 application of manual pressure lateral to the patella may cause an observable shift in synovial fluid (bulge) to the medial aspect. The examiner should note that this maneuver is only effective in detecting small to moderate effusions (<100 mL). Inflammatory disorders such as RA, gout, pseudogout, and psoriatic arthritis may involve the knee joint and produce significant pain, stiffness, swelling, or warmth. A popliteal or *Baker's cyst* may be palpated with the knee partially flexed and is best viewed posteriorly with the patient standing and knees fully extended to visualize isolated or unilateral popliteal swelling or fullness.

Anserine bursitis is an often missed periarticular cause of knee pain in adults. The pes anserine bursa underlies the insertion of the conjoined tendons (sartorius, gracilis, semitendinosus) on the anteromedial proximal tibia and may be painful following trauma, overuse, or inflammation. It is often tender in patients with fibromyalgia, obesity, and knee OA. Other forms of bursitis may also present as knee pain. The prepatellar bursa is superficial and is located over the inferior portion of the patella. The infrapatellar bursa is deeper and lies beneath the patellar ligament before its insertion on the tibial tubercle.

Internal derangement of the knee may result from trauma or degenerative processes. Damage to the meniscal cartilage (medial or lateral) frequently presents as chronic or intermittent knee pain. Such an injury should be suspected when there is a history of trauma, athletic activity, or chronic knee arthritis, and when the patient relates symptoms of "locking" or "giving way" of the knee. With the knee flexed 90° and the patient's foot on the table, pain elicited during palpation over the joint line or when the knee is stressed laterally or medially may suggest a meniscal tear. A positive McMurray test may also indicate a meniscal tear. To perform this test, the knee is first flexed at 90°, and the leg is then extended while the lower extremity is simultaneously torqued medially or laterally. A painful click during inward rotation may indicate a lateral meniscus tear, and pain during outward rotation may indicate a tear in the medial meniscus. Lastly, damage to the cruciate ligaments should be suspected with acute onset of pain, possibly with swelling, a history of trauma, or a synovial fluid aspirate that is grossly bloody. Examination of the cruciate ligaments is best accomplished by eliciting a drawer sign. With the patient recumbent, the knee should be partially flexed and the foot stabilized on the examining surface. The examiner should manually attempt to displace the tibia anteriorly or posteriorly with respect to the femur. If anterior movement is detected, then anterior cruciate ligament damage is likely. Conversely, significant posterior movement may indicate posterior cruciate damage. Contralateral comparison will assist the examiner in detecting significant anterior or posterior movement.

HIP PAIN

The hip is best evaluated by observing the patient's gait and assessing range of motion. The vast majority of patients reporting "hip pain" localize their pain unilaterally to the posterior gluteal musculature (Fig. 393-5). Such pain tends to radiate down the posterolateral aspect of the thigh and may or may not be associated with complaints of low back pain. This presentation frequently results from degenerative arthritis of the lumbosacral spine or disks and commonly follows a dermatomal distribution with involvement of nerve roots between L4 and S1. Sciatica is caused by impingement of the L4, L5, or S1 nerve (i.e., from a herniated disk) and manifests as unilateral neuropathic pain extending from the gluteal region down the posterolateral leg to the foot. Some individuals instead localize their "hip pain" laterally to the area overlying the trochanteric bursa. Because of the depth of this bursa, swelling and warmth are usually absent. Diagnosis of trochanteric bursitis or enthesitis can be confirmed by inducing point tenderness over the trochanteric bursa. Gluteal

and trochanteric pain are common findings in fibromyalgia. Range of movement may be limited by pain. Pain in the hip joint is less common and tends to be located anteriorly, over the inguinal ligament; it may radiate medially to the groin. Uncommonly, iliopsoas bursitis may mimic true hip joint pain. Diagnosis of iliopsoas bursitis may be suggested by a history of trauma or inflammatory arthritis. Pain associated with iliopsoas bursitis is localized to the groin or anterior thigh and tends to worsen with hyperextension of the hip; many patients prefer to flex and externally rotate the hip to reduce the pain from a distended bursa.

LABORATORY INVESTIGATIONS

The vast majority of musculoskeletal disorders can be easily diagnosed by a complete history and physical examination. An additional objective of the initial encounter is to determine whether additional investigations or immediate therapy is required. Additional evaluation is indicated with: (1) monarticular conditions; (2) traumatic or inflammatory conditions; (3) the presence of neurologic findings; (4) systemic manifestations; or (5) chronic symptoms (>6 weeks) and a lack of response to symptomatic measures. The extent and nature of the additional investigation should be dictated by the clinical features and suspected pathologic process. Laboratory tests should be used to confirm a specific clinical diagnosis and not be used to screen or evaluate patients with vague rheumatic complaints. Indiscriminate use of broad batteries of diagnostic tests and radiographic procedures is rarely a useful or cost-effective means to establish a diagnosis.

Besides a complete blood count, including a white blood cell (WBC) and differential count, the routine evaluation should include a determination of an acute-phase reactant such as the ESR or CRP, which can be useful in discriminating inflammatory from noninflammatory disorders. Both are inexpensive, easily obtained, and may be elevated with infection, inflammation, autoimmune disorders, neoplasia, pregnancy, renal insufficiency, advanced age, or hyperlipidemia. Extreme elevation of the acute-phase reactants (CRP, ESR) is seldom seen without evidence of serious illness (e.g., sepsis, pleuropericarditis, polymyalgia rheumatica, giant cell arteritis, adult Still's disease).

Serum uric acid determinations are useful in the diagnosis of gout and in monitoring the response to urate-lowering therapy. Uric acid, the end product of purine metabolism, is primarily excreted in the urine. Serum values range from 238 to 516 μmol/L (4.0–8.6 mg/dL) in men; the lower values (178–351 μmol/L [3.0–5.9 mg/dL]) seen in

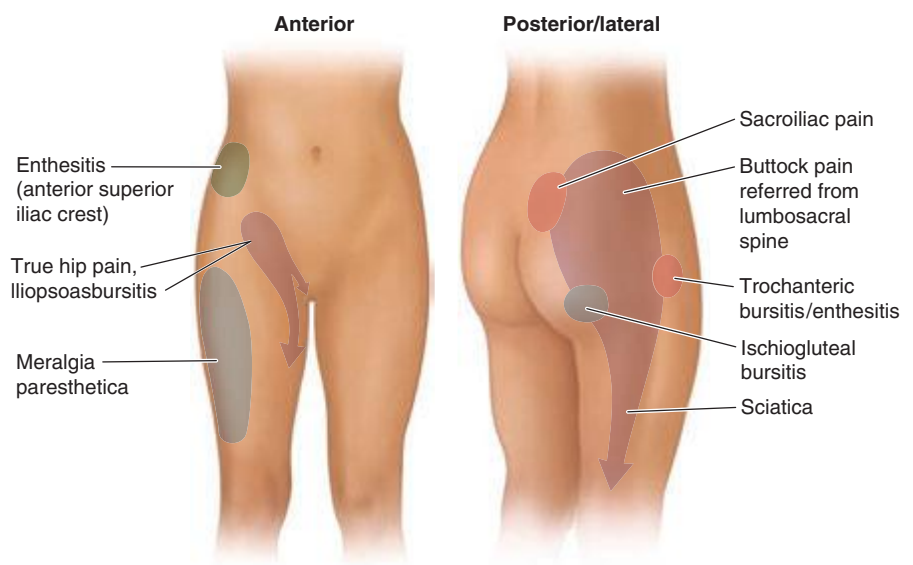


FIGURE 393-5 Origins of hip pain and dysesthesias. (From JJ Cush et al: *Evaluation of musculoskeletal complaints, in Rheumatology: Diagnosis and Therapeutics, 2nd ed, JJ Cush et al [eds]. Philadelphia, Lippincott Williams & Wilkins, 2005, pp 3–20. Used with permission from Dr. John J. Cush.*)

women are caused by the uricosuric effects of estrogen. Urinary uric acid levels are normally <750 mg per 24 h. Although hyperuricemia (especially levels >535 $\mu\text{mol/L}$ [9 mg/dL]) is associated with an increased incidence of gout and nephrolithiasis, levels may not correlate with the severity of articular disease. Uric acid levels (and the risk of gout) may be increased by inborn errors of metabolism (Lesch-Nyhan syndrome), disease states (renal insufficiency, myeloproliferative disease, psoriasis), or drugs (alcohol, cytotoxic therapy, thiazides). Although nearly all patients with gout will demonstrate hyperuricemia at some time during their illness, up to 50% of patients with an acute gouty attack will have normal serum uric acid levels. Monitoring serum uric acid may be useful in assessing the response to urate-lowering therapy or chemotherapy, with the target goal being a serum urate <6 mg/dL.

Serologic tests for rheumatoid factor (RF), cyclic citrullinated peptide (CCP or ACPA) antibodies, ANAs, complement levels, Lyme and antineutrophil cytoplasmic antibodies (ANCA), or antistreptolysin O (ASO) titer should be carried out only when there is clinical evidence to specifically suggest an associated diagnosis, because these have poor predictive value when used for screening, especially when the pretest probability is low. For most of these, there is no value to repeated or serial serologic testing. Although 4–5% of a healthy population will have positive tests for RF and ANAs, only 1% and <0.4% of the population will have RA or SLE, respectively. IgM RF (autoantibodies against the Fc portion of IgG) is found in 80% of patients with RA and may also be seen in low titers in patients with chronic infections (tuberculosis, leprosy, hepatitis); other autoimmune diseases (SLE, Sjögren's syndrome); and chronic pulmonary, hepatic, or renal diseases. When considering RA, both serum RF and anti-CCP antibodies should be obtained as these are complementary. Both are comparably sensitive, but CCP antibodies are more specific than RF. In RA, the presence of anti-CCP and RF antibodies may indicate a greater risk for more severe, erosive polyarthritis. ANAs are found in nearly all patients with SLE and may also be seen in patients with other autoimmune diseases (polymyositis, scleroderma, antiphospholipid syndrome, Sjögren's syndrome), drug-induced lupus (Table 393-2), chronic liver or renal disorders, and advanced age. Positive ANAs are found in 5% of adults and in up to 14% of elderly or chronically ill individuals. The ANA test is very sensitive but poorly specific for lupus, as only 1–2% of all positive results will be caused by lupus alone. The interpretation of a positive ANA test may depend on the magnitude of the titer and the pattern observed by immunofluorescence microscopy (Table 393-4). Diffuse and speckled patterns are least specific, whereas a peripheral, or rim, pattern (related to autoantibodies against double-strand [native] DNA) is highly specific and suggestive of lupus. Centromeric patterns are seen in patients with limited

scleroderma (calcinosis, Raynaud's phenomenon, esophageal involvement, sclerodactyly, telangiectasia [CREST] syndrome) or primary biliary sclerosis, and nucleolar patterns may be seen in patients with diffuse systemic sclerosis or inflammatory myositis.

Aspiration and analysis of synovial fluid are always indicated in acute monoarthritis or when an infectious or crystal-induced arthropathy is suspected. Synovial fluid may distinguish between noninflammatory and inflammatory processes by analysis of the appearance, viscosity, and cell count. Tests for synovial fluid glucose, protein, lactate dehydrogenase, lactic acid, or autoantibodies are not recommended because they have no diagnostic value. Normal synovial fluid is clear or a pale straw color and is viscous, primarily because of the high levels of hyaluronate. Noninflammatory synovial fluid is clear, viscous, and amber-colored, with a WBC count of <2000/ μL and a predominance of mononuclear cells. The viscosity of synovial fluid is assessed by expressing fluid from the syringe one drop at a time. Normally, there is a stringing effect, with a long tail behind each synovial drop. Effusions caused by OA or trauma will have normal viscosity. Inflammatory fluid is turbid and yellow, with an increased WBC count (2000–50,000/ μL) and a polymorphonuclear leukocyte predominance. Inflammatory fluid has reduced viscosity, diminished hyaluronate, and little or no tail following each drop of synovial fluid. Such effusions are found in RA, gout, and other inflammatory arthritides. Septic fluid is opaque and purulent, with a WBC count usually >50,000/ μL , a predominance of polymorphonuclear leukocytes (>75%), and low viscosity. Such effusions are typical of septic arthritis but may also occur with RA or gout. In addition, hemorrhagic synovial fluid may be seen with trauma, hemarthrosis, or neuropathic arthritis. An algorithm for synovial fluid aspiration and analysis is shown in Fig. 393-6. Synovial fluid should be analyzed immediately for appearance, viscosity, and cell count. Monosodium urate crystals (observed in gout) are seen by polarized microscopy and are long, needle-shaped, negatively birefringent, and usually intracellular. In chondrocalcinosis and pseudogout, calcium pyrophosphate dihydrate crystals are usually short, rhomboid-shaped, and positively birefringent. Whenever infection is suspected, synovial fluid should be Gram stained and cultured appropriately. If gonococcal arthritis is suspected, nucleic acid amplification tests should be used to detect either *Chlamydia trachomatis* or *N. gonorrhoeae* infection. Synovial fluid from patients with chronic monoarthritis should also be cultured for *M. tuberculosis* and fungi. Last, it should be noted that crystal-induced arthritis and septic arthritis occasionally occur together in the same joint.

DIAGNOSTIC IMAGING IN JOINT DISEASES

Conventional radiography has been a valuable tool in the diagnosis and staging of articular disorders. Plain x-rays are most appropriate and cost effective when there is a history of trauma, suspected chronic infection, progressive disability, or monoarticular involvement; when therapeutic alterations are considered; or when a baseline assessment is desired for what appears to be a chronic process. However, in acute inflammatory arthritis, early radiography is rarely helpful in establishing a diagnosis and may only reveal soft tissue swelling or juxtaarticular demineralization. As the disease progresses, calcification (of soft tissues, cartilage, or bone), joint space narrowing, erosions, bony ankylosis, new bone formation (sclerosis, osteophytes, or periostitis), or subchondral cysts may develop and suggest specific clinical entities. Consultation with a radiologist will help define the optimal imaging modality, technique, or positioning and prevent the need for further studies.

Additional imaging techniques may possess greater diagnostic sensitivity and facilitate early diagnosis in a limited number of articular disorders and in selected circumstances and are indicated when conventional radiography is inadequate or nondiagnostic (Table 393-5). *Ultrasonography* is useful in the detection of soft tissue abnormalities, such as tendinitis, tenosynovitis, enthesitis, bursitis, and entrapment neuropathies. Wider use, lower cost, better technology, and enhanced site-specific transducers now allow for routine use in outpatient care. Owing to low cost, portability, and wider use, ultrasound use has grown and is the preferred method for the evaluation of synovial (Baker's) cysts, rotator cuff tears, tendinitis and tendon injury, and

TABLE 393-4 ANTINUCLEAR ANTIBODY (ANA) PATTERNS AND CLINICAL ASSOCIATIONS

ANA Pattern	Antigen Identified	Clinical Correlate
Diffuse	Deoxyribonucleoprotein	Nonspecific
	Histones	Drug-induced lupus, lupus
Peripheral (rim)	ds-DNA	50% of SLE (specific)
Speckled	U1-RNP	>90% of MCTD
	Sm	30% of SLE (specific)
	Ro (SS-A)	Sjögren's 60%, SCLC, neonatal lupus, ANA(–) lupus
	La (SS-B)	50% of Sjögren's, 15% lupus
	Scl-70	40% of diffuse scleroderma
	PM-1	Polymyositis (PM), dermatomyositis
	Jo-1	PM w/pneumonitis + arthritis
Nucleolar	RNA polymerase I, others	40% of PSS
Centromere	Kinetochore	75% CREST (limited scleroderma)

Abbreviations: ANA, antinuclear antibody; CREST, calcinosis, Raynaud phenomenon, esophageal involvement, sclerodactyly, and telangiectasia; MCTD, mixed connective tissue disease; PSS, progressive systemic sclerosis; SCLC, subacute cutaneous lupus erythematosus; SLE, systemic lupus erythematosus.

INTERPRETATION OF SYNOVIAL FLUID ASPIRATION

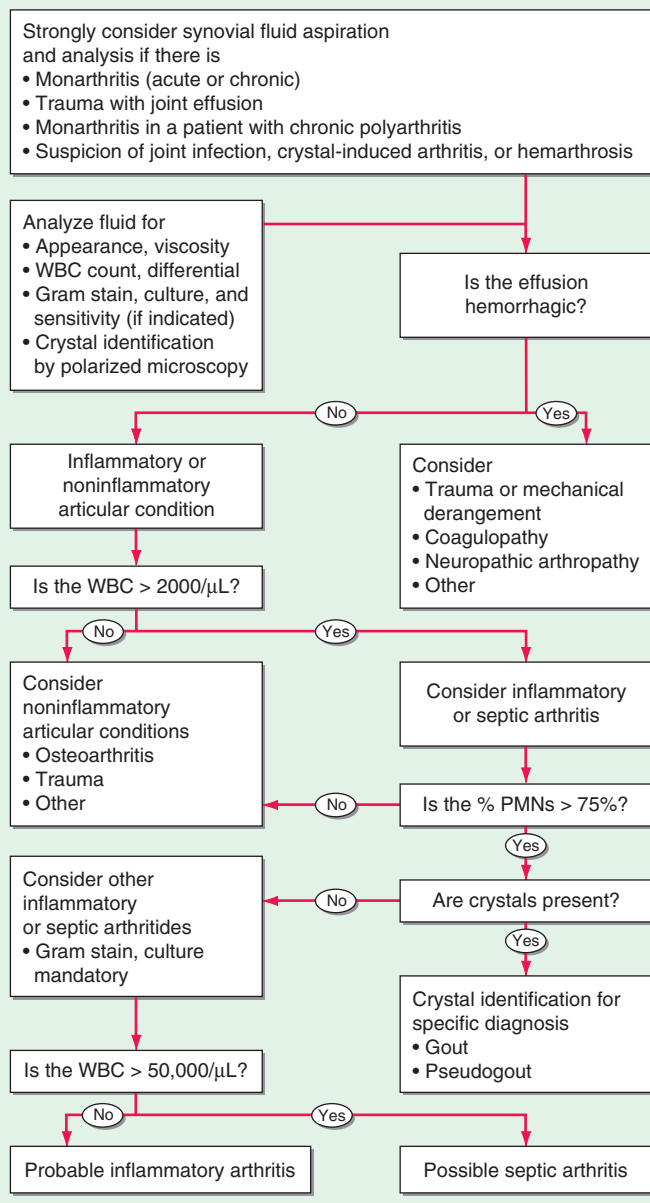


FIGURE 393-6 Algorithmic approach to the use and interpretation of synovial fluid aspiration and analysis. PMNs, polymorphonuclear (leukocytes); WBC, white blood cell (count).

crystal deposition on cartilage. Use of power Doppler allows for early detection of synovitis and bony erosions. *Radionuclide scintigraphy* is a very sensitive, but poorly specific, means of detecting inflammatory or metabolic alterations in bone or periarticular soft tissue structures. Scintigraphy is best suited for total-body assessment (extent and distribution) of skeletal involvement (neoplasia, Paget's disease) and the assessment of patients with undiagnosed polyarthralgias, looking for occult arthritis. The use of scintigraphy has declined with greater use and declining cost of ultrasound and MRI. The limited tissue contrast resolution of scintigraphy may obscure the distinction between a bony or periarticular process and may necessitate the additional use of MRI. Scintigraphy using ^{99m}Tc , ^{67}Ga , or ^{111}In -labeled WBCs has been applied to a variety of articular disorders with variable success (Table 393-5). Although [^{99m}Tc] diphosphate scintigraphy may be useful in identifying osseous infection, neoplasia, inflammation, increased blood flow, bone remodeling, heterotopic bone formation, or avascular necrosis, MRI is preferred in most instances. Gallium scanning uses ^{67}Ga , which binds serum and cellular transferrin and lactoferrin and is preferentially taken up by neutrophils, macrophages,

TABLE 393-5 DIAGNOSTIC IMAGING TECHNIQUES FOR MUSCULOSKELETAL DISORDERS

Method	Imaging Time, h	Cost ^a	Current Indications
Ultrasound	<1	++	Synovial (Baker's) cysts Rotator cuff tears Bursitis, tendinitis, tendon injury Enthesitis Carpal tunnel syndrome Urate or calcium pyrophosphate deposition on cartilage Early detection of synovial inflammation or erosions Ultrasound-guided injection/arthrocentesis
Radionuclide scintigraphy			
^{99m}Tc	1–4	++	Metastatic bone survey Evaluation of Paget's disease Identifying occult arthritis in patients with undiagnosed polyarthralgia
^{111}In -WBC	24	+++	Acute infection Prosthetic infection Acute osteomyelitis
^{67}Ga	24–48	++++	Acute and chronic infection Acute osteomyelitis
Computed tomography (CT)	<1	+++	Herniated intervertebral disk Sacroiliitis Spinal stenosis Spinal trauma Osteoid osteoma Stress fracture
Dual-energy CT	<1	NA	Uric acid deposition Tophus localization
Magnetic resonance imaging	1/2–2	++++	Avascular necrosis Osteomyelitis Septic arthritis, infected prosthetic joints Early sacroiliitis Intraarticular derangement and soft tissue injury Derangements of axial skeleton and spinal cord Herniated intervertebral disk Pigmented villonodular synovitis Inflammatory and metabolic muscle pathology

^aRelative cost for imaging study.

Abbreviations: NA, not commercially available; WBC, white blood cell.

bacteria, and tumor tissue (e.g., lymphoma). As such, it is primarily used in the identification of occult infection or malignancy. Scanning with ^{111}In -labeled WBCs has been used to detect osteomyelitis and infectious or inflammatory arthritis. Despite their utility, ^{111}In -labeled WBC or ^{67}Ga scanning has largely been replaced by MRI, except when there is a suspicion of septic joint or prosthetic joint infections.

Computed tomography (CT) provides detailed visualization of the axial skeleton. Articulations previously considered difficult to visualize by radiography (e.g., zygapophyseal, sacroiliac, sternoclavicular, hip joints) can be effectively evaluated using CT. CT has been demonstrated to be useful in the diagnosis of low back pain syndromes (e.g., spinal stenosis vs herniated disk), sacroiliitis, osteoid osteoma, and stress fractures. Helical or spiral CT (with or without contrast angiography) is a novel technique that is rapid, cost effective, and sensitive



FIGURE 393-7 Dual-energy computed tomography (DECT) scan from a 45-year-old woman with right ankle swelling around the lateral malleolus. Three-dimensional volume-rendered coronal reformatted DECT image shows that the mass is composed of monosodium urate (red) in keeping with tophus (arrow). (Used with permission from S Nicolaou et al: *AJR* 194:1072, 2010.)

in diagnosing pulmonary embolism or obscure fractures, often in the setting of initially equivocal findings. High-resolution CT can be advocated in the evaluation of suspected or established infiltrative lung disease (e.g., scleroderma or rheumatoid lung). The recent use of hybrid (positron emission tomography [PET] or single-photon emission CT [SPECT]) CT scans in metastatic evaluations has incorporated CT to provide better anatomic localization of scintigraphic abnormalities.

¹⁸F-Fluorodeoxyglucose (FDG) is the most commonly used radiopharmaceutical in PET scanning. FDG-PET/CT scans have been seldom used in the evaluation of septic or inflammatory arthritis. Dual-energy CT (DECT) scanning, developed in urology to identify urinary calculi, has been a highly sensitive and specific method used to identify and quantify uric acid deposition in tissues (Fig. 393-7).

MRI has significantly advanced the ability to image musculoskeletal structures. MRI has the advantages of providing multiplanar images with fine anatomic detail and contrast resolution (Fig. 393-8) that allows for the superior ability to visualize bone marrow and soft tissue periarticular structures. Although more costly with a longer procedural time than CT, the MRI has become the preferred technique when evaluating complex musculoskeletal disorders.

MRI can image fascia, vessels, nerve, muscle, cartilage, ligaments, tendons, pannus, synovial effusions, and bone marrow. Visualization of particular structures can be enhanced by altering the pulse sequence to produce either T1- or T2-weighted spin echo, gradient echo, or inversion recovery (including short tau inversion recovery [STIR]) images. Because of its sensitivity to changes in marrow fat, MRI is a sensitive but nonspecific means of detecting osteonecrosis, osteomyelitis, and marrow inflammation indicating overlying synovitis or osteitis (Fig. 393-8). Because of its enhanced soft tissue resolution, MRI is more sensitive than arthrography or CT in the diagnosis of soft tissue injuries (e.g., meniscal and rotator cuff tears); intraarticular derangements; marrow abnormalities (osteonecrosis, myeloma); and spinal cord or nerve root damage, synovitis, or cartilage damage or loss.

ACKNOWLEDGMENT

The author acknowledges the contributions of Dr. Peter E. Lipsky to this chapter in previous editions.



FIGURE 393-8 Superior sensitivity of magnetic resonance imaging (MRI) in the diagnosis of osteonecrosis of the femoral head.

A 45-year-old woman receiving high-dose glucocorticoids developed right hip pain. Conventional x-rays (top) demonstrated only mild sclerosis of the right femoral head. T1-weighted MRI (bottom) demonstrated low-density signal in the right femoral head, diagnostic of osteonecrosis.

394 Osteoarthritis

David T. Felson

Osteoarthritis (OA) is the most common type of arthritis. Its high prevalence, especially in the elderly, and the high rate of disability related to disease make it a leading cause of disability in the elderly. Because of the aging of Western populations and because obesity, a major risk factor, is increasing in prevalence, the occurrence of OA is on the rise. In the United States, OA prevalence will increase by 66–100% by 2020.

OA affects certain joints, yet spares others (Fig. 394-1). Commonly affected joints include the cervical and lumbosacral spine, hip, knee, and first metatarsal phalangeal joint (MTP). In the hands, the distal and proximal interphalangeal joints and the base of the thumb are often affected. Usually spared are the wrist, elbow, and ankle. Our joints were designed, in an evolutionary sense, for brachiating apes, animals that still walked on four limbs. We thus develop OA in joints that were ill designed for human tasks such as pincer grip (OA in the thumb base) and walking upright (OA in knees and hips). Some joints, like the ankles, may be spared because their articular cartilage may be uniquely resistant to loading stresses.

OA can be diagnosed based on structural abnormalities or on the symptoms these abnormalities evoke. According to cadaveric studies, by elderly years, structural changes of OA are nearly universal. These include cartilage loss (seen as joint space loss on x-rays) and osteophytes. Many persons with x-ray evidence of OA have no joint symptoms, and although the prevalence of structural abnormalities is of interest in understanding disease pathogenesis, what matters more from a clinical perspective is the prevalence of symptomatic OA. Symptoms, usually joint pain, determine disability, visits to clinicians, and disease costs.

Symptomatic OA of the knee (pain on most days of a recent month in a knee plus x-ray evidence of OA in that knee) occurs in ~12% of persons age ≥ 60 in the United States and 6% of all adults age ≥ 30 . Symptomatic hip OA is roughly one-third as common as disease in the knee. Although radiographically evident hand OA and the appearance of bony enlargement in affected hand joints (Fig. 394-2) are extremely common in older persons, most cases are often not symptomatic. Even so, symptomatic

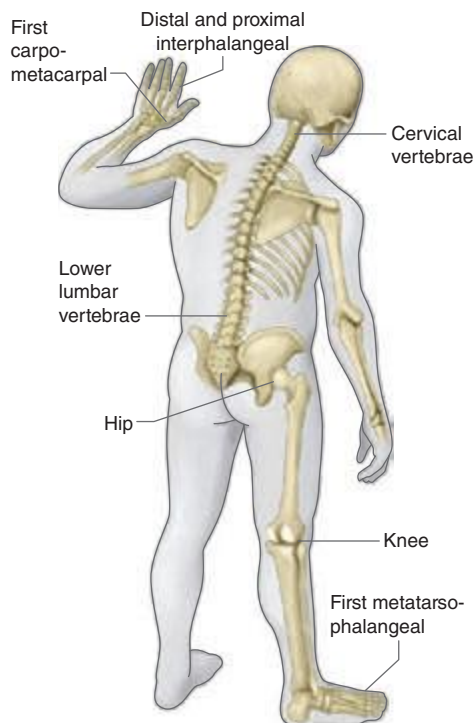


FIGURE 394-1 Joints commonly affected by osteoarthritis.



FIGURE 394-2 Severe osteoarthritis of the hands affecting the distal interphalangeal joints (Heberden's nodes) and the proximal interphalangeal joints (Bouchard's nodes). There is no clear bony enlargement of the other common site in the hands, the thumb base.

hand OA occurs in ~10% of elderly individuals and often produces measurable limitation in function.

The prevalence of OA rises strikingly with age. Regardless of how it is defined, OA is uncommon in adults under age 40 and highly prevalent in those over age 60. It is also a disease that, at least in middle-aged and elderly persons, is much more common in women than in men, and sex differences in prevalence increase with age.

X-ray evidence of OA is common in the lower back and neck, but back pain and neck pain have not been tied to findings of OA on x-ray. Thus, back pain and neck pain are treated separately (Chap. 22).

DEFINITION

OA is joint failure, a disease in which all structures of the joint have undergone pathologic change, often in concert. The pathologic sine qua non of disease is hyaline articular cartilage loss, present in a focal and, initially, nonuniform manner. This is accompanied by increasing thickness and sclerosis of the subchondral bony plate, by outgrowth of osteophytes at the joint margin, by stretching of the articular capsule, by mild synovitis in many affected joints, and by weakness of muscles bridging the joint. In knees, meniscal degeneration is part of the disease. There are numerous pathways that lead to joint failure, but the initial step is often joint injury in the setting of a failure of protective mechanisms.

JOINT PROTECTIVE MECHANISMS AND THEIR FAILURE

Joint protectors include joint capsule and ligaments, muscle, sensory afferents, and underlying bone. Joint capsule and ligaments serve as joint protectors by providing a limit to excursion, thereby fixing the range of joint motion.

Synovial fluid reduces friction between articulating cartilage surfaces, thereby serving as a protector against friction-induced cartilage wear. This lubrication function depends on *hyaluronic acid* and on *lubricin*, a mucinous glycoprotein secreted by synovial fibroblasts whose concentration diminishes after joint injury and in the face of synovial inflammation.

The ligaments, along with overlying skin and tendons, contain mechanoreceptor sensory afferent nerves. These mechanoreceptors fire at different frequencies throughout a joint's range of motion, providing feedback by way of the spinal cord to muscles and tendons. As a consequence, these muscles and tendons can assume the right tension at appropriate points in joint excursion to act as optimal joint protectors, anticipating joint loading.

Muscles and tendons that bridge the joint are key joint protectors. Their coordinated contractions at the appropriate time in joint movement provide the appropriate power and acceleration for the limb to accomplish its tasks. Focal stress across the joint is minimized by muscle contraction that decelerates the joint before impact and assures that when joint impact arrives, it is distributed broadly across the joint surface.

Failure of these joint protectors increases the risk of joint injury and OA. For example, in animals, OA develops rapidly when a sensory nerve to the joint is sectioned and joint injury induced. Similarly, in humans, Charcot's arthropathy, a severe and rapidly progressive OA, develops when minor joint injury occurs in the presence of posterior column peripheral neuropathy. Another example of joint protector failure is rupture of ligaments, a well-known cause of the early development of OA.

CARTILAGE AND ITS ROLE IN JOINT FAILURE

In addition to being a primary target tissue for disease, cartilage also functions as a joint protector. A thin rim of tissue at the ends of two opposing bones, cartilage is lubricated by synovial fluid to provide an almost frictionless surface across which these two bones move. The compressible stiffness of cartilage compared to bone provides the joint with impact-absorbing capacity.

The earliest changes of OA may occur in cartilage, and abnormalities there can accelerate disease development. The two major macromolecules in cartilage are type 2 collagen, which provides cartilage its tensile strength, and aggrecan, a proteoglycan macromolecule linked with hyaluronic acid, which consists of highly negatively charged glycosaminoglycans. In normal cartilage, type 2 collagen is woven tightly, constraining the aggrecan molecules in the interstices between collagen strands, forcing these highly negatively charged molecules into close proximity with one another. The aggrecan molecule, through electrostatic repulsion of its negative charges, gives cartilage its compressive stiffness. Chondrocytes, the cells within this avascular tissue, synthesize all elements of the matrix and produce enzymes that break down the matrix. Synovium and chondrocytes synthesize and release cytokines and growth factors, which provide feedback that modulates synthesis of matrix molecules (Fig. 394-3). Cartilage matrix synthesis and catabolism are in a dynamic equilibrium influenced by the cytokine and growth factor environment. Mechanical and osmotic stress on chondrocytes induces these cells to alter gene expression and increase production of inflammatory cytokines and matrix-degrading enzymes. While chondrocytes synthesize numerous enzymes, matrix metalloproteinases (MMP) (especially collagenases and ADAMTS-5) are critical enzymes in the breakdown of cartilage matrix. Both collagenase and aggrecanases act primarily in the territorial matrix surrounding chondrocytes; however, as the osteoarthritic process develops, their activities and effects spread throughout the matrix, especially in the superficial layers of cartilage.

The synovium, cartilage, and bone all influence disease development through cytokines, chemokines, and even complement activation (Fig. 394-3). These act on chondrocyte cell surface receptors and ultimately have transcriptional effects. Matrix fragments released from cartilage stimulate synovitis. Among the most important cytokines are interleukin (IL) 1 β , which exerts transcriptional effects on chondrocytes, stimulating production of proteinases and suppressing cartilage matrix synthesis. Tumor necrosis factor (TNF) α may play a similar role to that of IL-1. These cytokines also induce chondrocytes to synthesize prostaglandin E₂ and nitric oxide, which have complex effects on matrix synthesis and degradation. At early stages in the matrix response to injury and in the healthy response to loading, the net effect of cytokine stimulation may be matrix synthesis, but ultimately, the combination of effects on chondrocytes triggers matrix degradation. Enzymes in the matrix are held in check by activation inhibitors, including tissue inhibitor of metalloproteinase (TIMP). Growth factors are also part of this complex network, with BMP-2 and transforming growth factor β playing prominent roles in stimulating the development of osteophytes. Whereas healthy articular cartilage is avascular in part due to angiogenesis inhibitors present in cartilage, disease is characterized by the invasion of blood vessels into cartilage from underlying bone and proliferation of vessels within synovium. This is influenced by vascular endothelial growth factor (VEGF) synthesis in the cartilage and bone. With these blood vessels come nerves that may bring nociceptive innervation.

Probably as a result of chronic oxidative damage, articular chondrocytes exhibit an age-related decline in synthetic capacity while maintaining the ability to produce proinflammatory mediators and matrix-degrading enzymes, findings characteristic of a senescent secretory phenotype. These chondrocytes are unable to maintain tissue homeostasis (such as after insults of a mechanical or inflammatory nature). Thus, with age, cartilage is easily damaged by minor sometimes unnoticed injuries, including those that are part of daily activities.

OA cartilage is characterized by gradual depletion of aggrecan, an unfurling of the tightly woven collagen matrix, and loss of type 2 collagen. With these changes comes increasing vulnerability of cartilage, which loses its compressive stiffness.

RISK FACTORS

Joint vulnerability and joint loading are the two major factors contributing to the development of OA. On the one hand, a vulnerable joint whose protectors are dysfunctional can develop OA with minimal

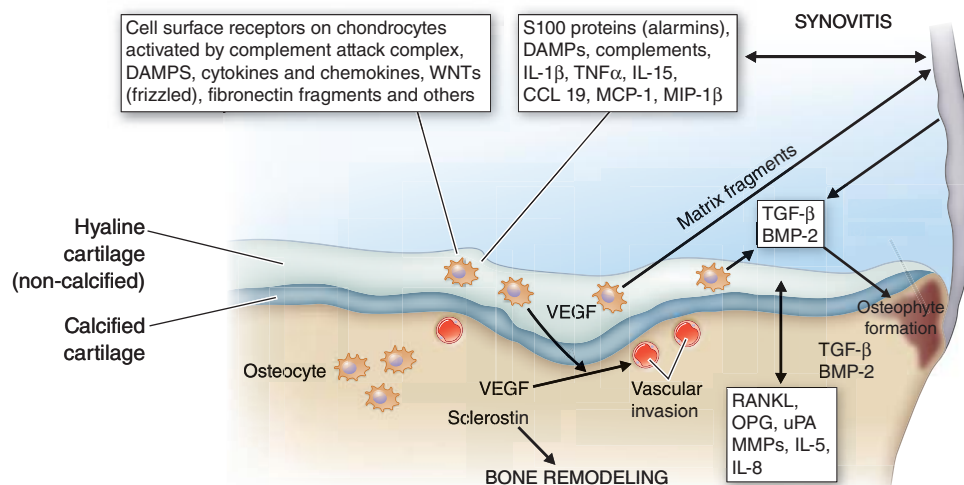


FIGURE 394-3 Selected factors involved in the osteoarthritic process including chondrocytes, bone, and synovium. Synovitis causes release of cytokines, alarmins, damage-associated molecular pattern (DAMP) molecules, and complement, which activate chondrocytes through cell surface receptors. Chondrocytes produce matrix molecules (collagen type 2, aggrecan) and the enzymes responsible for the degradation of the matrix (e.g., ADAMTS-5 and matrix metalloproteinases [MMPs]). Bone invasion occurs through the calcified cartilage, triggered by vascular endothelial growth factor (VEGF) and other molecules. IL, interleukin; TGF, transforming growth factor; TNF, tumor necrosis factor. (From RF Loeser et al: *Arthritis Rheum* 64:1697, 2012.)

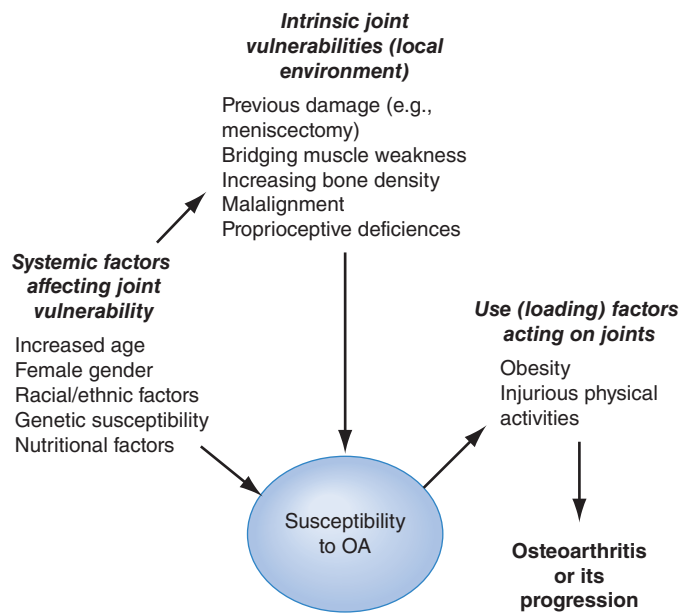


FIGURE 394-4 Risk factors for osteoarthritis (OA) either contribute to the susceptibility of the joint (systemic factors or factors in the local joint environment) or increase risk by the load they put on the joint. Usually a combination of loading and susceptibility factors is required to cause disease or its progression.

levels of loading, perhaps even levels encountered during everyday activities. On the other hand, in a young joint with competent protectors, a major acute injury or long-term overloading is necessary to precipitate disease. Risk factors for OA can be understood in terms of their effect either on joint vulnerability or on loading (Fig. 394-4).

SYSTEMIC RISK FACTORS

Age is the most potent risk factor for OA. Radiographic evidence of OA is rare in individuals under age 40; however, in some joints, such as the hands, OA occurs in >50% of persons over age 70. Aging increases joint vulnerability through several mechanisms. Whereas dynamic loading of joints stimulates cartilage matrix synthesis by chondrocytes in young cartilage, aged cartilage is less responsive to these stimuli. Partly because of this failure to synthesize matrix with loading, cartilage thins with age, and thinner cartilage experiences higher shear stress at basal layers and is at greater risk of cartilage damage. Also, joint protectors fail more often with age. Muscles that bridge the joint become weaker with age and also respond less quickly to oncoming impulses. Sensory nerve input slows with age, retarding the feedback loop of mechanoreceptors to muscles and tendons related to their tension and position. Ligaments stretch with age, making them less able to absorb impulses. These factors work in concert to increase the vulnerability of older joints to OA.

Older women are at high risk of OA in all joints, a risk that emerges as women reach their sixth decade. Although hormone loss with menopause may contribute to this risk, there is little understanding of the unique vulnerability of older women versus men to OA.

HERITABILITY AND GENETICS

OA is a highly heritable disease, but its heritability is joint specific. Fifty percent of the hand and hip OA in the community is attributable to inheritance, i.e., to disease present in other members of the family. However, the heritable proportion of knee OA is at most 30%, with some studies suggesting no heritability at all. Whereas many people with OA have disease in multiple joints, this “generalized OA” phenotype is rarely inherited and is more often a consequence of aging.

Emerging evidence has identified genetic mutations that confer a high risk of OA, the best replicated is a polymorphism within the growth differentiation factor 5 gene. This polymorphism diminishes the quantity of GDF5; GDF5 has its main influence on joint shape, and genes predisposing to OA are likely to increase risk of disease based on their effects on joint development and shape.

GLOBAL CONSIDERATIONS



Hip OA is rare in China and in immigrants from China to the United States. However, OA in the knees is at least as common, if not more so, in Chinese than in whites from the United States, and knee OA represents a major cause of disability in China, especially in rural areas. Anatomic differences between Chinese and white hips may account for much of the difference in hip OA prevalence, with white hips having a higher prevalence of anatomic predispositions to the development of OA. Persons from Africa, but not African Americans, may also have a very low rate of hip OA.

RISK FACTORS IN THE JOINT ENVIRONMENT

Some risk factors increase vulnerability of the joint through local effects on the joint environment. With changes in joint anatomy, for example, load across the joint is no longer distributed evenly across the joint surface, but rather shows an increase in focal stress. In the hip, three uncommon developmental abnormalities occurring in utero or in childhood, congenital dysplasia, Legg-Perthes disease, and slipped capital femoral epiphysis, leave a child with distortions of hip joint anatomy that often lead to OA later in life. Girls are predominantly affected by acetabular dysplasia, a mild form of congenital dislocation, whereas the other abnormalities more often affect boys. Depending on the severity of the anatomic abnormalities, hip OA occurs either in young adulthood (severe abnormalities) or middle age (mild abnormalities).

Major injuries to a joint also can produce anatomic abnormalities that leave the joint susceptible to OA. For example, a fracture through the joint surface often causes OA in joints in which the disease is otherwise rare such as the ankle and the wrist. Avascular necrosis can lead to collapse of dead bone at the articular surface, producing anatomic irregularities and subsequent OA.

Tears of ligamentous and fibrocartilaginous structures that protect the joints, such as the anterior cruciate ligament and the meniscus in the knee and the labrum in the hip, can lead to premature OA. Meniscal tears increase with age and when chronic are often asymptomatic but lead to adjacent cartilage damage and accelerated OA. Even injuries in which the affected person never received a diagnosis may increase risk of OA. For example, in the Framingham Study subjects, men with a history of major knee injury, but no surgery, had a 3.5-fold increased risk for subsequent knee OA.

Another source of anatomic abnormality is malalignment across the joint (Fig. 394-5). This factor has been best studied in the knee, which is the fulcrum of the longest lever arm in the body. Varus (bow-legged) knees with OA are at exceedingly high risk of cartilage loss in the medial or inner compartment of the knee, whereas valgus (knock-kneed) malalignment predisposes to rapid cartilage loss in the lateral compartment. Malalignment causes this effect by increasing stress on a focal area of cartilage, which then breaks down. There is evidence that malalignment in the knee not only causes cartilage loss but leads

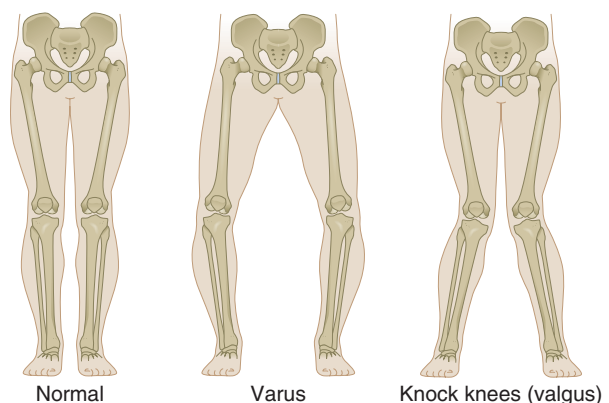


FIGURE 394-5 The two types of limb malalignment in the frontal plane: varus, in which the stress is placed across the medial compartment of the knee joint, and valgus, which places excess stress across the lateral compartment of the knee.

to underlying bone damage, producing bone marrow lesions seen on magnetic resonance imaging (MRI). Malalignment in the knee often produces such a substantial increase in focal stress within the knee (as evidenced by its destructive effects on subchondral bone) that severely malaligned knees may be destined to progress regardless of the status of other risk factors.

Weakness in the quadriceps muscles bridging the knee increases the risk of the development of painful OA in the knee.

Patients with knee OA have impaired proprioception across their knees, and this may predispose them to further disease progression. The role of bone in serving as a shock absorber for impact load is not well understood, but persons with increased bone density are at high risk of OA, suggesting that the resistance of bone to impact during joint use may play a role in disease development.

LOADING FACTORS

Obesity Three to six times body weight is transmitted across the knee during single-leg stance. Any increase in weight may be multiplied by this factor to reveal the excess force across the knee in overweight persons during walking. Obesity is a well-recognized and potent risk factor for the development of knee OA and, less so, for hip OA. Obesity precedes the development of disease and is not just a consequence of the inactivity present in those with disease. It is a stronger risk factor for disease in women than in men, and in women, the relationship of weight to the risk of disease is linear, so that with each increase in weight, there is a commensurate increase in risk. Weight loss in women lowers the risk of developing symptomatic disease. Not only is obesity a risk factor for OA in weight-bearing joints, but obese persons have more severe symptoms from the disease.

Obesity's effect on the development and progression of disease is mediated mostly through the increased loading in weight-bearing joints that occurs in overweight persons. However, a modest association of obesity with an increased risk of hand OA suggests that there may be a systemic metabolic factor circulating in obese persons that affects disease risk also.

Repeated Use of Joint and Exercise There are two categories of repetitive joint use, occupational use and leisure time physical activities. Workers performing repetitive tasks as part of their occupations for many years are at high risk of developing OA in the joints they use repeatedly. For example, farmers are at high risk for hip OA, and miners have high rates of OA in knees and spine. Workers whose jobs require regular knee bending or lifting or carrying heavy loads have a high rate of knee OA. One reason why workers may get disease is that during long days at work, their muscles may gradually become exhausted, no longer serving as effective joint protectors.

It is widely recommended for people to adopt an exercise-filled lifestyle, and long-term studies of exercise suggest no consistent association of exercise with OA risk in the majority of persons. However, persons who already have injured joints may put themselves at greater risk by engaging in certain types of exercise. For example, persons who have already sustained major knee injuries are at increased risk of progressive knee OA as a consequence of running. In addition, compared to nonrunners, elite runners (professional runners and those on Olympic teams) have high risks of both knee and hip OA. Lastly, although recreational runners are not at increased risk of knee OA, studies suggest that they have a modest increased risk of disease in the hip.

PATHOLOGY

The pathology of OA provides evidence of the involvement of many joint structures in disease. Cartilage initially shows surface fibrillation and irregularity. As disease progresses, focal erosions develop there, and these eventually extend down to the subjacent bone. With further progression, cartilage erosion down to bone expands to involve a larger proportion of the joint surface, even though OA remains a focal disease with nonuniform loss of cartilage (Fig. 394-6).

After an injury to cartilage, chondrocytes undergo mitosis and clustering. Although the metabolic activity of these chondrocyte clusters is

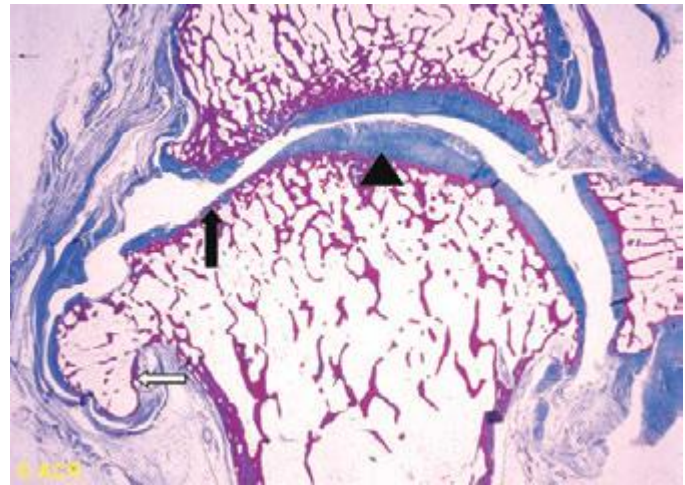


FIGURE 394-6 Pathologic changes of osteoarthritis in a toe joint.

Note the nonuniform loss of cartilage (*arrowhead vs solid arrow*), the increased thickness of the subchondral bone envelope (*solid arrow*), and the osteophyte (*open arrow*). (From the American College of Rheumatology slide collection.)

high, the net effect of this activity is to promote proteoglycan depletion in the matrix surrounding the chondrocytes. This is because the catabolic is greater than the synthetic activity. As disease develops, collagen matrix becomes damaged, the negative charges of proteoglycans get exposed, and cartilage swells from ionic attraction to water molecules. Because in damaged cartilage proteoglycans are no longer forced into close proximity, cartilage does not bounce back after loading as it did when healthy, and cartilage becomes vulnerable to further injury. Chondrocytes at the basal level of cartilage undergo apoptosis.

With loss of cartilage come alterations in subchondral bone. Stimulated by growth factors and cytokines, osteoclasts and osteoblasts in the subchondral bony plate, just underneath cartilage, become activated. Bone formation produces a thickening and stiffness of the subchondral plate that occurs even before cartilage ulcerates. Trauma to bone during joint loading may be the primary factor driving this bone response, with healing from injury (including microcracks) producing stiffness. Small areas of osteonecrosis usually exist in joints with advanced disease. Bone death may also be caused by bone trauma with shearing of microvasculature, leading to a cutoff of vascular supply to some bone areas.

At the margin of the joint, near areas of cartilage loss, osteophytes form. These start as outgrowths of new cartilage, and with neurovascular invasion from the bone, this cartilage ossifies. Osteophytes are an important radiographic hallmark of OA. In malaligned joints, osteophytes grow larger on the side of the joint subject to most loading stress (e.g., in varus knees, osteophytes grow larger on the medial side).

The synovium produces lubricating fluids that minimize shear stress during motion. In healthy joints, the synovium consists of a single discontinuous layer filled with fat and containing two types of cells, macrophages and fibroblasts, but in OA, it can sometimes become edematous and inflamed. There is a migration of macrophages from the periphery into the tissue, and cells lining the synovium proliferate. Enzymes secreted by the synovium digest cartilage matrix that has been released from the surface of the cartilage.

Additional pathologic changes occur in the capsule, which stretches, becomes edematous, and can become fibrotic.

The pathology of OA is not identical across joints. In hand joints with severe OA, for example, there are often cartilage erosions in the center of the joint probably produced by bony pressure from the opposite side of the joint.

Basic calcium phosphate and calcium pyrophosphate dihydrate crystals are present microscopically in most joints with end-stage OA. Their role in osteoarthritic cartilage is unclear, but their release from cartilage into the joint space and joint fluid likely triggers synovial

2230 inflammation, which can, in turn, produce release of enzymes and trigger nociceptive stimulation.

SOURCES OF PAIN

Because cartilage is aneural, cartilage loss in a joint is not accompanied by pain. Thus, pain in OA likely arises from structures outside the cartilage. Innervated structures in the joint include the synovium, ligaments, joint capsule, muscles, and subchondral bone. Most of these are not visualized by the x-ray, and the severity of x-ray changes in OA correlates poorly with pain severity.

Based on MRI studies in osteoarthritic knees comparing those with and without pain and on studies mapping tenderness in unanesthetized joints, likely sources of pain include synovial inflammation, joint effusions, and bone marrow edema. Modest synovitis develops in many but not all osteoarthritic joints. Some diseased joints have no synovitis, whereas others have synovial inflammation that approaches the severity of joints with rheumatoid arthritis (Chap. 380). The presence of synovitis on MRI is correlated with the presence and severity of knee pain. Capsular stretching from fluid in the joint stimulates nociceptive fibers there, inducing pain. Increased focal loading as part of the disease not only damages cartilage but probably also injures the underlying bone. As a consequence, bone marrow edema appears on the MRI; histologically, this edema signals the presence of microcracks and scar, which are the consequences of trauma. These lesions may stimulate bone nociceptive fibers. Also, hemostatic pressure within bone rises in OA, and the increased pressure itself may stimulate nociceptive fibers, causing pain.

Pain may arise from outside the joint also, including bursae near the joints. Common sources of pain near the knee are anserine bursitis and iliotibial band syndrome.

Persons with chronic OA pain may develop nervous system alterations as a consequence of disease, changes which decrease inhibitory controls on nociception and its distribution. This may produce allodynia and hyperalgesia in some patients with OA.

CLINICAL FEATURES

Joint pain from OA is activity-related. Pain comes on either during or just after joint use and then gradually resolves. Examples include knee or hip pain with going up or down stairs, pain in weight-bearing joints when walking, and, for hand OA, pain when cooking. Early in disease, pain is episodic, triggered often by a day or two of overactive use of a diseased joint, such as a person with knee OA taking a long run and noticing a few days of pain thereafter. As disease progresses, the pain becomes continuous and even begins to be bothersome at night. Stiffness of the affected joint may be prominent, but morning stiffness is usually brief (<30 min).

In knees, buckling may occur, in part, due to weakness of muscles crossing the joint. Mechanical symptoms, such as buckling, catching, or locking, could also signify internal derangement, such as meniscal tears, and need to be evaluated. In the knee, pain with activities requiring knee flexion, such as stair climbing and arising from a chair, often emanates from the patellofemoral compartment of the knee, which does not actively articulate until the knee is bent ~35°.

OA is the most common cause of chronic knee pain in persons over age 45, but the differential diagnosis is long. Inflammatory arthritis is likely if there is prolonged morning stiffness and many other joints are affected. Bursitis occurs commonly around knees and hips. A physical examination should focus on whether tenderness is over the joint line (at the junction of the two bones around which the joint is articulating) or is outside of it. Anserine bursitis, medial and distal to the knee, is an extremely common cause of chronic knee pain that may respond to a glucocorticoid injection. Prominent nocturnal pain in the absence of end-stage OA merits a distinct workup. For hip pain, OA can be detected by loss of internal rotation on passive movement, and pain isolated to an area lateral to the hip joint usually reflects the presence of trochanteric bursitis.

No blood tests are routinely indicated for workup of patients with OA unless symptoms and signs suggest inflammatory arthritis.



FIGURE 394-7 X-ray of knee with medial osteoarthritis. Note the narrowed joint space on medial side of the joint only (white arrow), the sclerosis of the bone in the medial compartment providing evidence of cortical thickening (black arrow), and the osteophytes in the medial femur (white wedge).

Examination of the synovial fluid is often more helpful diagnostically than an x-ray. If the synovial fluid white count is >1000/ μ L, inflammatory arthritis or gout or pseudogout is likely, the latter two being also identified by the presence of crystals.

X-rays are indicated to evaluate chronic hand pain and hip pain thought to be due to OA, as the diagnosis is often unclear without confirming radiographs. For knee pain, x-rays should be obtained if symptoms or signs are not typical of OA or if knee pain persists after inauguration of effective treatment. In OA, radiographic findings (Fig. 394-7) correlate poorly with the presence and severity of pain. Further, radiographs may be normal in early disease as they are insensitive to cartilage loss and other early findings.

Although MRI may reveal the extent of pathology in an osteoarthritic joint, it is not indicated as part of the diagnostic workup. Findings such as meniscal tears and cartilage and bone lesions occur in most patients with OA in the knee, but almost never warrant a change in therapy.

TREATMENT OSTEOARTHRITIS

The goals of the treatment of OA are to alleviate pain and minimize loss of physical function. To the extent that pain and loss of function are consequences of inflammation, of weakness across the joint, and of laxity and instability, the treatment of OA involves addressing each of these impairments. Comprehensive therapy consists of a multimodality approach including nonpharmacologic and pharmacologic elements.

Patients with mild and intermittent symptoms may need only reassurance or nonpharmacologic treatments. Patients with ongoing, disabling pain are likely to need both nonpharmacotherapy and pharmacotherapy.

Treatments for knee OA have been more completely evaluated than those for hip and hand OA or for disease in other joints. Thus, although the principles of treatment are identical for OA in all joints, we shall focus below on the treatment of knee OA, noting specific recommendations for disease in other joints, especially when they differ from those for the knee.

NONPHARMACOTHERAPY

Because OA is a mechanically driven disease, the mainstay of treatment involves altering loading across the painful joint and improving the function of joint protectors, so they can better distribute load across the joint. Ways of lessening focal load across the joint include:

1. avoiding activities that overload the joint, as evidenced by their causing pain;
2. improving the strength and conditioning of muscles that bridge the joint, so as to optimize their function; and
3. unloading the joint, either by redistributing load within the joint with a brace or a splint or by unloading the joint during weight bearing with a cane or a crutch.

The simplest effective treatment for many patients is to avoid activities that precipitate pain. For example, for the middle-aged patient whose long-distance running brings on symptoms of knee OA, a less demanding form of weight-bearing activity may alleviate all symptoms. For an older person whose daily constitutional up and down hills bring on knee pain, routing the constitutional away from hills might eliminate symptoms.

Each pound of weight increases the loading across the knee three- to sixfold. Weight loss may have a commensurate multiplier effect, unloading both knees and hips and probably relieving pain in those joints.

In hand joints affected by OA, splinting, by limiting motion, often minimizes pain for patients with involvement especially in the base of the thumb. Weight-bearing joints such as knees and hips can be unloaded by using a cane in the hand opposite to the affected joint for partial weight bearing. A physical therapist can help teach the patient how to use the cane optimally, including ensuring that its height is optimal for unloading. Crutches or walkers can serve a similar beneficial function.

Exercise Osteoarthritic pain in knees or hips during weight bearing results in lack of activity and poor mobility, and because OA is so common, the inactivity that results represents a public health concern, increasing the risk of cardiovascular disease and obesity. Aerobic capacity is poor in most elders with symptomatic knee OA, worse than others of the same age.

Weakness in muscles that bridge osteoarthritic joints is multifactorial in etiology. First, there is a decline in strength with age. Second, with limited mobility comes disuse muscle atrophy. Third, patients with painful knee or hip OA alter their gait so as to lessen loading across the affected joint, and this further diminishes muscle use. Fourth, “arthrogenous inhibition” may occur, whereby contraction of muscles bridging the joint is inhibited by a nerve afferent feedback loop emanating in a swollen and stretched joint capsule; this prevents maximal attainment of voluntary maximal strength. Because adequate muscle strength and conditioning are critical to joint protection, weakness in a muscle that bridges a diseased joint makes the joint more susceptible to further damage and pain. The degree of weakness correlates strongly with the severity of joint pain and the degree of physical limitation. One of the cardinal elements of the treatment of OA is to improve the functioning of muscles surrounding the joint.

For knee and hip OA, trials have shown that exercise lessens pain and improves physical function. Most effective exercise regimens consist of aerobic and/or resistance training, the latter of which focuses on strengthening muscles across the joint. Exercises are likely to be effective, especially if they train muscles for the activities a person performs daily. Activities that increase pain in the joint should be avoided, and the exercise regimen needs to be individualized to optimize effectiveness. Range-of-motion exercises, which do not strengthen muscles, and isometric exercises that strengthen muscles, but not through range of motion, are unlikely to be effective by themselves. Low-impact exercises, including water aerobics and water resistance training, are often better tolerated by patients than exercises involving impact loading, such as running or treadmill exercises. A patient should be referred to an exercise class or to a therapist who can create an individualized regimen, and then an individualized home-based regimen can be crafted. In addition to conventional exercise regimens, tai chi may be effective for knee OA. However, there is no strong evidence that patients with hand OA benefit from therapeutic exercise.

Adherence over the long term is the major challenge to an exercise prescription. In trials involving patients with knee OA, who are engaged in exercise treatment, from a third to over half of patients stopped exercising by 6 months. Less than 50% continued regular exercise at 1 year. The strongest predictor of a patient’s continued exercise is a previous personal history of successful exercise. Physicians should reinforce the exercise prescription at each clinic visit, help the patient recognize barriers to ongoing exercise, and identify convenient times for exercise to be done routinely. The combination of exercise with calorie restriction and weight loss is especially effective in lessening pain.

Correction of Malalignment Malalignment in the frontal plane (varus-valgus) markedly increases the stress across the joint, which can lead to progression of disease and to pain and disability (Fig. 394-5). Correcting malalignment, either surgically or with bracing, may relieve pain in persons whose knees are malaligned. Malalignment develops over years as a consequence of gradual anatomic alterations of the joint and bone, and correcting it is often very challenging. One way is with a fitted brace, which takes an often varus osteoarthritic knee and straightens it by putting valgus stress across the knee. Unfortunately, many patients are unwilling to wear a realigning knee brace; in addition, in patients with obese legs, braces may slip with usage and lose their realigning effect. They are indicated for willing patients who can learn to put them on correctly and on whom they do not slip.

Other ways of correcting malalignment across the knee include the use of orthotics in footwear. Unfortunately, although they may have modest effects on knee alignment, trials have heretofore not demonstrated efficacy of a lateral wedge orthotic versus placebo wedges.

Pain from the patellofemoral compartment of the knee can be caused by tilting of the patella or patellar malalignment with the patella riding laterally or medially in the femoral trochlear groove. Using a brace to realign the patella, or tape to pull the patella back into the trochlear sulcus or reduce its tilt, has been shown, when compared to placebo taping in clinical trials, to lessen patellofemoral pain. However, patients may find it difficult to apply tape, and skin irritation from the tape is common. Commercial patellar braces may be a solution, but there is insufficient evidence on their efficacy to recommend them.

Although their effect on malalignment is questionable, neoprene sleeves pulled to cover the knee lessen pain and are easy to use and popular among patients. The explanation for their therapeutic effect on pain is unclear.

In patients with knee OA, acupuncture produces modest pain relief compared to placebo needles and may be an adjunctive treatment.

PHARMACOTHERAPY

Although nonpharmacologic approaches to therapy constitute its mainstay, pharmacotherapy serves an important adjunctive role in OA treatment. Available drugs are administered using oral, topical, and intraarticular routes.

Acetaminophen, Nonsteroidal Anti-Inflammatory Drugs (NSAIDs), and Cyclooxygenase-2 (COX-2) Inhibitors Acetaminophen (paracetamol) is the initial analgesic of choice for patients with OA in knees, hips, or hands. For some patients, it is adequate to control symptoms, in which case more toxic drugs such as NSAIDs can be avoided. Doses up to 1 g three times daily can be used ([Table 394-1](#)).

NSAIDs are the most popular drugs to treat osteoarthritic pain. They can be administered either topically or orally. In clinical trials, oral NSAIDs produce ~30% greater improvement in pain than high-dose acetaminophen. Occasional patients treated with NSAIDs experience dramatic pain relief, whereas others experience little improvement. Initially, NSAIDs should be administered topically or taken orally on an “as needed” basis because side effects are less frequent with low intermittent doses. If occasional medication use is insufficiently effective, then daily treatment may be indicated, with

TABLE 394-1 PHARMACOLOGIC TREATMENT FOR OSTEOARTHRITIS

Treatment	Dosage	Comments
Acetaminophen	Up to 1 g tid	Prolongs half-life of warfarin. Make sure patient is not taking other treatments containing acetaminophen to avoid hepatic toxicity.
Oral NSAIDs and COX-2 inhibitors	375–500 mg bid 1500 mg bid	Take with food. Increased risk of myocardial infarction and stroke for some NSAIDs and especially COX-2 inhibitors. High rates of gastrointestinal side effects, including ulcers and bleeding, occur. Patients at high risk for gastrointestinal side effects should also take either a proton pump inhibitor or misoprostol. ^a There is an increase in gastrointestinal side effects or bleeding when taken with acetylsalicylic acid. Can also cause edema and renal insufficiency.
Naproxen	600–800 mg 3–4 times a day	
Salsalate		
Ibuprofen		
Topical NSAIDs		Rub onto joint. Few systemic side effects. Skin irritation common.
Diclofenac Na 1% gel	4 g qid (for knees, hands)	
Opiates	Various	Common side effects include dizziness, sedation, nausea or vomiting, dry mouth, constipation, urinary retention, and pruritus. Respiratory and central nervous system depression can occur.
Capsaicin	0.025–0.075% cream 3–4 times a day	Can irritate mucous membranes.
Intraarticular injections		
Steroids		
Hyaluronans	Varies from 3–5 weekly injections depending on preparation	Mild to moderate pain at injection site. Controversy exists regarding efficacy.

^aPatients at high risk include those with previous gastrointestinal events, persons ≥ 60 years, and persons taking glucocorticoids. Trials have shown the efficacy of proton pump inhibitors and misoprostol in the prevention of ulcers and bleeding. Misoprostol is associated with a high rate of diarrhea and cramping; therefore, proton pump inhibitors are more widely used to reduce NSAID-related gastrointestinal symptoms.

Abbreviations: COX-2, cyclooxygenase-2; NSAIDs, nonsteroidal anti-inflammatory drugs.

Source: Adapted from DT Felson: *N Engl J Med* 354:841, 2006.

an anti-inflammatory dose selected (Table 394-1). Patients should be reminded to take low-dose aspirin and ibuprofen at different times to eliminate a drug interaction.

NSAIDs taken orally have substantial and frequent side effects, the most common of which is upper gastrointestinal toxicity, including dyspepsia, nausea, bloating, gastrointestinal bleeding, and ulcer disease. Some 30–40% of patients experience upper gastrointestinal (GI) side effects so severe as to require discontinuation of medication. To minimize the risk of nonsteroidal-related GI side effects, patients should not take two NSAIDs and should take medications after food; if risk is high, patients should take a gastroprotective agent, such as a proton pump inhibitor. Certain oral agents are safer to the stomach than others, including nonacetylated salicylates and nabumetone. Major NSAID-related GI side effects can occur in patients who do not complain of upper GI symptoms. In one study of patients hospitalized for GI bleeding, 81% had no premonitory symptoms.

Because of the increased rates of cardiovascular events associated with COX-2 inhibitors and with some conventional NSAIDs such as diclofenac, many of these drugs are not appropriate long-term treatment choices for older persons with OA, especially those at high risk of heart disease or stroke. The American Heart Association has identified rofecoxib and all other COX-2 inhibitors as putting patients at high risk, although low doses of celecoxib (≤ 200 mg/d) may not be associated with an elevation of risk. The only conventional NSAID that appears safe from a cardiovascular perspective is naproxen, but it does have GI toxicity.

There are other common side effects of NSAIDs, including the tendency to develop edema because of prostaglandin inhibition of afferent blood supply to glomeruli in the kidneys and, for similar reasons, a predilection toward reversible renal insufficiency. Blood pressure may increase modestly in some NSAID-treated patients. Oral NSAIDs should not be used in patients with stage IV or V renal disease and should be used with caution in those with stage III disease.

NSAIDs can be placed into a gel or topical solution with another chemical modality that enhances penetration of the skin barrier creating a topical NSAID. When absorbed through the skin, plasma concentrations are an order of magnitude lower than with the same amount of drug administered orally or parenterally. However, when these drugs are administered topically in proximity to a superficial

joint (knees, hands, but not hips), the drug can be found in joint tissues such as the synovium and cartilage. Trial results have varied but generally have found that topical NSAIDs are slightly less efficacious than oral agents, but have far fewer GI and systemic side effects. Unfortunately, topical NSAIDs often cause local skin irritation where the medication is applied, inducing redness, burning, or itching in up to 40% of patients (see Table 394-1).

Intraarticular Injections: Glucocorticoids and Hyaluronic Acid Because synovial inflammation is likely to be a major cause of pain in patients with OA, local anti-inflammatory treatments administered intra-articularly may be effective in ameliorating pain, at least temporarily. Glucocorticoid injections provide such efficacy, but response is variable, with some patients having little relief of pain whereas others experience pain relief lasting several months. Glucocorticoid injections are useful to get patients over acute flares of pain and may be especially indicated if the patient has coexistent OA and crystal deposition disease, especially from calcium pyrophosphate dihydrate crystals (**Chap. 395**). There is no evidence that repeated glucocorticoid injections into the joint are dangerous.

Hyaluronic acid injections can be given for treatment of symptoms in knee and hip OA, but there is controversy as to whether they have efficacy versus placebo (Table 394-1).

Other Classes of Drugs and Nutraceuticals For patients with symptomatic knee or hip OA who have not had an adequate response to the treatments above and are either unwilling to undergo or are not candidates for total joint arthroplasty, opioid analgesics have shown modest efficacy and can be tried. Opioid management plans and patient selection are critical. Another option is the use of duloxetine, which has demonstrated modest efficacy in OA.

Recent guidelines recommend against the use of glucosamine or chondroitin for OA. Large publicly supported trials have failed to show that, compared with placebo, these compounds relieve pain in persons with disease.

Optimal nonsurgical therapy for OA is often achieved by trial and error, with each patient having idiosyncratic responses to specific treatments. When medical therapies have failed and the patient has an unacceptable reduction in their quality of life and ongoing pain

and disability, then at least for knee and hip OA, total joint arthroplasty is indicated.

SURGERY

For knee OA, several operations are available. Arthroscopic debridement and lavage have diminished in popularity after randomized trials evaluating this operation have showed that its efficacy is no greater than that of sham surgery or no treatment for relief of pain or disability. Even mechanical symptoms such as buckling, which are extremely common in patients with knee OA, do not respond to arthroscopic debridement. Although arthroscopic meniscectomy is indicated for acute meniscal tears in which symptoms such as locking and acute pain are clearly related temporally to a knee injury that produced the tear, recent trials show that doing a partial meniscectomy in persons with OA and a symptomatic meniscal tear does not relieve knee pain or improve function.

For patients with knee OA isolated to the medial compartment, operations to realign the knee to lessen medial loading can relieve pain. These include a high tibial osteotomy, in which the tibia is broken just below the tibial plateau and realigned so as to load the lateral, nondiseased compartment, or a unicompartmental replacement with realignment. Each surgery may provide the patient with years of pain relief before a total knee replacement is required.

Ultimately, when the patient with knee or hip OA has failed medical treatment modalities and remains in pain, with limitations of physical function that compromise the quality of life, the patient should be referred for total knee or hip arthroplasty. These are highly efficacious operations that relieve pain and improve function in the vast majority of patients, although rates of success are higher for hip than knee replacement. Currently failure rates for both are ~1% per year, although these rates are higher in obese patients. The chance of surgical success is greater in centers where at least 25 such operations are performed yearly or with surgeons who perform multiple operations annually. The timing of knee or hip replacement is critical. If the patient suffers for many years until their functional status has declined substantially, with considerable muscle weakness, postoperative functional status may not improve to a level achieved by others who underwent operation earlier in their disease course.

Cartilage Regeneration Chondrocyte transplantation has not been found to be efficacious in OA, perhaps because OA includes pathology of joint mechanics, which is not corrected by chondrocyte transplants. Similarly, abrasion arthroplasty (chondroplasty) has not been well studied for efficacy in OA, but it produces fibrocartilage in place of damaged hyaline cartilage. Both of these surgical attempts to regenerate and reconstitute articular cartilage may be more likely to be efficacious early in disease when joint malalignment and many of the other noncartilage abnormalities that characterize OA have not yet developed.

395 Gout and Other Crystal-Associated Arthropathies

H. Ralph Schumacher, Lan X. Chen

The use of polarizing light microscopy during synovial fluid analysis in 1961 by McCarty and Hollander and the subsequent application of other crystallographic techniques, such as electron microscopy, energy-dispersive elemental analysis, and x-ray diffraction, have allowed investigators to identify the roles of different microcrystals, including monosodium urate (MSU), calcium pyrophosphate (CPP), calcium apatite (apatite), and calcium oxalate (CaOx), in inducing

TABLE 395-1 MUSCULOSKELETAL MANIFESTATIONS OF CRYSTAL-INDUCED ARTHRITIS

Acute mono- or polyarthritis	Destructive arthropathies
Bursitis	Chronic inflammatory arthritis
Tendinitis	Spinal arthritis
Enthesitis	Peculiar type of osteoarthritis
Tophaceous deposits	Carpal tunnel syndrome

acute or chronic arthritis or peri-arthritis. The clinical events that result from deposition of MSU, CPP, apatite, and CaOx have many similarities but also have important differences. Because of often similar clinical presentations, the need to perform synovial fluid analysis to distinguish the type of crystal involved must be emphasized. Polarized light microscopy alone can identify most typical crystals; apatite, however, is an exception. Aspiration and analysis of effusions are also important to assess the possibility of infection. Apart from the identification of specific microcrystalline materials or organisms, synovial fluid characteristics in crystal-associated diseases are nonspecific, and synovial fluid can be inflammatory or noninflammatory. Without crystal identification, these diseases can be confused with rheumatoid or other types of arthritis. A list of possible musculoskeletal manifestations of crystal-associated arthritis is shown in [Table 395-1](#).

GOUT

Gout is a metabolic disease that most often affects middle-aged to elderly men and postmenopausal women. It results from an increased body pool of urate with hyperuricemia. It typically is characterized by episodic acute arthritis or chronic arthritis caused by deposition of MSU crystals in joints and connective tissue tophi and the risk for deposition in kidney interstitium or uric acid nephrolithiasis ([Chap. 431e](#)).

ACUTE AND CHRONIC ARTHRITIS

Acute arthritis is the most common early clinical manifestation of gout. Usually, only one joint is affected initially, but polyarticular acute gout can occur in subsequent episodes. The metatarsophalangeal joint of the first toe often is involved, but tarsal joints, ankles, and knees also are affected commonly. Especially in elderly patients or in advanced disease, finger joints may be involved. Inflamed Heberden's or Bouchard's nodes may be a first manifestation of gouty arthritis. The first episode of acute gouty arthritis frequently begins at night with dramatic joint pain and swelling. Joints rapidly become warm, red, and tender, with a clinical appearance that often mimics that of cellulitis. Early attacks tend to subside spontaneously within 3–10 days, and most patients have intervals of varying length with no residual symptoms until the next episode. Several events may precipitate acute gouty arthritis: dietary excess, trauma, surgery, excessive ethanol ingestion, hypouricemic therapy, and serious medical illnesses such as myocardial infarction and stroke.

After many acute mono- or oligoarticular attacks, a proportion of gouty patients may present with a chronic nonsymmetric synovitis, causing potential confusion with rheumatoid arthritis ([Chap. 380](#)). Less commonly, chronic gouty arthritis will be the only manifestation, and, more rarely, the disease will manifest only as periarticular tophaceous deposits in the absence of synovitis. Women represent only 5–20% of all patients with gout. Most women with gouty arthritis are postmenopausal and elderly, have osteoarthritis and arterial hypertension that causes mild renal insufficiency, and usually are receiving diuretics. Premenopausal gout is rare. Kindreds of precocious gout in young females caused by decreased renal urate clearance and renal insufficiency have been described.

Laboratory Diagnosis Even if the clinical appearance strongly suggests gout, the presumptive diagnosis ideally should be confirmed by needle aspiration of acutely or chronically involved joints or tophaceous deposits. Acute septic arthritis, several of the other crystalline-associated arthropathies, palindromic rheumatism, and psoriatic arthritis may present with similar clinical features. During acute gouty



FIGURE 395-1 Extracellular and intracellular monosodium urate crystals, as seen in a fresh preparation of synovial fluid, illustrate needle- and rod-shaped crystals. These crystals are strongly negative birefringent crystals under compensated polarized light microscopy; 400 \times .

attacks, needle-shaped MSU crystals typically are seen both intracellularly and extracellularly (Fig. 395-1). With compensated polarized light, these crystals are brightly birefringent with negative elongation. Synovial fluid leukocyte counts are elevated from 2000 to 60,000/ μ L. Effusions appear cloudy due to the increased numbers of leukocytes. Large amounts of crystals occasionally produce a thick pasty or chalky joint fluid. Bacterial infection can coexist with urate crystals in synovial fluid; if there is any suspicion of septic arthritis, joint fluid must be cultured.

MSU crystals also can often be demonstrated in the first metatarsophalangeal joint and in knees not acutely involved with gout. Arthrocentesis of these joints is a useful technique to establish the diagnosis of gout between attacks.

Serum uric acid levels can be normal or low at the time of an acute attack, as inflammatory cytokines can be uricosuric and effective initiation of hypouricemic therapy can precipitate attacks. This limits the value of serum uric acid determinations for the diagnosis of gout. Nevertheless, serum urate levels are almost always elevated at some time and are important to use to follow the course of hypouricemic therapy. A 24-h urine collection for uric acid can, in some cases, be useful in assessing the risk of stones, elucidating overproduction or underexcretion of uric acid, and deciding whether it may be appropriate to use a uricosuric therapy (Chap. 431e). Excretion of >800 mg of uric acid per 24 h on a regular diet suggests that causes of overproduction of purine should be considered. Urinalysis, serum creatinine, hemoglobin, white blood cell (WBC) count, liver function tests, and serum lipids should be obtained because of possible pathologic sequelae of gout and other associated diseases requiring treatment and as baselines because of possible adverse effects of gout treatment.

Radiographic Features Cystic changes, well-defined erosions with sclerotic margins (often with overhanging bony edges), and soft tissue masses are characteristic radiographic features of advanced chronic tophaceous gout. Ultrasound may aid earlier diagnosis by showing a double contour sign overlying the articular cartilage. Dual-energy computed tomography (CT) can show specific features establishing the presence of urate crystals.

TREATMENT GOUT

ACUTE GOUTY ARTHRITIS

The mainstay of treatment during an acute attack is the administration of anti-inflammatory drugs such as nonsteroidal anti-inflammatory drugs (NSAIDs), colchicine, or glucocorticoids. NSAIDs are used most often in individuals without complicating comorbid conditions. Both colchicine and NSAIDs may be poorly tolerated and dangerous in the elderly and in the presence of renal insufficiency and gastrointestinal

disorders. Ice pack applications and rest of the involved joints can be helpful. Colchicine given orally is a traditional and effective treatment if used early in an attack. Useful regimens are one 0.6-mg tablet given every 8 h with subsequent tapering or 1.2 mg followed by 0.6 mg in 1 h with subsequent day dosing depending on response. This is generally better tolerated than the formerly advised higher dose regimens. The drug must be at least temporarily discontinued promptly at the first sign of loose stools, and symptomatic treatment must be given for the diarrhea. Intravenous colchicine has been taken off the market. NSAIDs given in full anti-inflammatory doses are effective in ~90% of patients, and the resolution of signs and symptoms usually occurs in 5–8 days. The most effective drugs are any of those with a short half-life and include indomethacin, 25–50 mg tid; naproxen, 500 mg bid; ibuprofen, 800 mg tid; diclofenac, 50 mg tid; and celecoxib 800 mg followed by 400 mg 12 h later, then 400 mg bid.

Glucocorticoids given IM or orally, for example, prednisone, 30–50 mg/d as the initial dose and gradually tapered with the resolution of the attack, can be effective in polyarticular gout. For a single joint or a few involved joints, intraarticular triamcinolone acetonide, 20–40 mg, or methylprednisolone, 25–50 mg, have been effective and well tolerated. Based on recent evidence on the essential role of the inflammasome and interleukin 1 β (IL-1 β) in acute gout, anakinra has been used, and other inhibitors of IL-1 β , including canakinumab and rilonacept, are under investigation.

HYPOURICEMIC THERAPY

Ultimate control of gout requires correction of the basic underlying defect: the hyperuricemia. Attempts to normalize serum uric acid to <300–360 μ mol/L (5.0–6.0 mg/dL) to prevent recurrent gouty attacks and eliminate tophaceous deposits are critical and entail a commitment to hypouricemic regimens and medications that generally are required for life. Hypouricemic drug therapy should be considered when, as in most patients, the hyperuricemia cannot be corrected by simple means (control of body weight, low-purine diet, increase in liquid intake, limitation of ethanol use, decreased use of fructose-containing foods and beverages, and avoidance of diuretics). The decision to initiate hypouricemic therapy usually is made taking into consideration the number of acute attacks (urate lowering may be cost-effective after two attacks), serum uric acid levels (progression is more rapid in patients with serum uric acid >535 μ mol/L [>9.0 mg/dL]), the patient's willingness to commit to lifelong therapy, or the presence of uric acid stones. Urate-lowering therapy should be initiated in any patient who already has tophi or chronic gouty arthritis. Uricosuric agents such as probenecid can be used in patients with good renal function who underexcrete uric acid, with <600 mg in a 24-h urine sample. Urine volume should be maintained by ingestion of 1500 mL of water every day. Probenecid can be started at a dose of 250 mg twice daily and increased gradually as needed up to 3 g per day to achieve and maintain a serum uric acid level of less than 6 mg/dL. Probenecid is generally not effective in patients with serum creatinine levels >177 μ mol/L (2 mg/dL). These patients may require allopurinol or benzbromarone (not available in the United States). Benzbromarone is another uricosuric drug that is more effective in patients with chronic kidney disease. Some agents used to treat common comorbidities, including losartan, fenofibrate, and amlodipine, have some mild uricosuric effects.

The xanthine oxidase inhibitor allopurinol is by far the most commonly used hypouricemic agent and is the best drug to lower serum urate in overproducers, urate stone formers, and patients with renal disease. It can be given in a single morning dose, usually 100 mg initially and increasing up to 800 mg if needed. In patients with chronic renal disease, the initial allopurinol dose should be lower and adjusted depending on the serum creatinine concentration; for example, with a creatinine clearance of 10 mL/min, one generally would use 100 mg every other day. Doses can be increased gradually to reach the target urate level of less than 6 mg/dL. Toxicity of allopurinol has been recognized increasingly in patients who use thiazide diuretics, in patients allergic to penicillin and ampicillin,

and in Asians expressing HLA-B*5801. The most serious side effects include life-threatening toxic epidermal necrolysis, systemic vasculitis, bone marrow suppression, granulomatous hepatitis, and renal failure. Patients with mild cutaneous reactions to allopurinol can reconsider the use of a uricosuric agent, undergo an attempt at desensitization to allopurinol, or take febuxostat, a new, chemically unrelated specific xanthine oxidase inhibitor. Febuxostat is approved in the United States at 40 or 80 mg once a day and does not require dose adjustment in mild to moderate renal disease. Pegloticase is a pegylated uricase, now available for patients who do not tolerate or fail full doses of other treatments. It is given intravenously usually at 8 mg every 2 weeks and can dramatically lower serum uric acid in up to 50% of such patients. New uricosurics are also undergoing investigation.

Urate-lowering drugs are generally not initiated during acute attacks but after the patient is stable and low-dose colchicine has been initiated to decrease the risk of the flares that often occur with urate lowering. Colchicine anti-inflammatory prophylaxis in doses of 0.6 mg one to two times daily should be given along with the hypouricemic therapy until the patient is normouricemic and without gouty attacks for 6 months or as long as tophi are present. Colchicine should not be used in dialysis patients and is given in lower doses in patients with renal disease or with P glycoprotein or CYP3A4 inhibitors such as clarithromycin that can increase toxicity of colchicine.

CALCIUM PYROPHOSPHATE DEPOSITION (CPPD) DISEASE

PATHOGENESIS

The deposition of CPP crystals in articular tissues is most common in the elderly, occurring in 10–15% of persons age 65–75 years and 30–50% of those >85 years. In most cases, this process is asymptomatic and the cause of CPPD is uncertain. Because >80% of patients are >60 years and 70% have preexisting joint damage from other conditions, it is likely that biochemical changes in aging or diseased cartilage favor crystal nucleation. In patients with CPPD arthritis, there is increased production of inorganic pyrophosphate and decreased levels of pyrophosphatases in cartilage extracts. Mutations in the *ANKH* gene, as described in both familial and sporadic cases, can increase elaboration and extracellular transport of pyrophosphate. The increase in pyrophosphate production appears to be related to enhanced activity of ATP pyrophosphohydrolase and 5'-nucleotidase, which catalyze the reaction of ATP to adenosine and pyrophosphate. This pyrophosphate could combine with calcium to form CPP crystals in matrix vesicles or on collagen fibers. There are decreased levels of cartilage glycosaminoglycans that normally inhibit and regulate crystal nucleation. High activities of transglutaminase enzymes also may contribute to the deposition of CPP crystals.

Release of CPP crystals into the joint space is followed by the phagocytosis of those crystals by monocyte-macrophages and neutrophils, which respond by releasing chemotactic and inflammatory substances and, as with MSU crystals, activating the inflammasome.

A minority of patients with CPPD arthropathy have metabolic abnormalities or hereditary CPP disease (Table 395-2). These associations suggest that a variety of different metabolic products may enhance CPP crystal deposition either by directly altering cartilage or by inhibiting inorganic pyrophosphatases. Included among these conditions are hyperparathyroidism, hemochromatosis, hypophosphatasia, hypomagnesemia, and possibly myxedema. The presence of CPPD arthritis in individuals <50 years old should lead to consideration of these metabolic disorders (Table 395-2) and inherited forms of disease, including those identified in a variety of ethnic groups. Genomic DNA studies performed on different kindreds have shown a possible location of genetic defects on chromosome 8q or on chromosome 5p in a region that expresses the gene of the membrane pyrophosphate channel (*ANKH* gene). As noted above, mutations described in the *ANKH* gene in kindreds with CPPD arthritis can increase extracellular

TABLE 395-2 CONDITIONS ASSOCIATED WITH CALCIUM PYROPHOSPHATE CRYSTAL DEPOSITION DISEASE

Aging
Disease-associated
Primary hyperparathyroidism
Hemochromatosis
Hypophosphatasia
Hypomagnesemia
Chronic gout
Postmeniscectomy
Gitelman's syndrome
Epiphyseal dysplasias

pyrophosphate and induce CPP crystal formation. Investigation of younger patients with CPPD should include inquiry for evidence of familial aggregation and evaluation of serum calcium, phosphorus, alkaline phosphatase, magnesium, iron, and transferrin.

CLINICAL MANIFESTATIONS

CPPD arthropathy may be asymptomatic, acute, subacute, or chronic or may cause acute synovitis superimposed on chronically involved joints. Acute CPPD arthritis originally was termed *pseudogout* by McCarty and co-workers because of its striking similarity to gout. Other clinical manifestations of CPPD include (1) association with or enhancement of peculiar forms of osteoarthritis; (2) induction of severe destructive disease that may radiographically mimic neuropathic arthritis; (3) production of chronic symmetric synovitis that is clinically similar to rheumatoid arthritis; (4) intervertebral disk and ligament calcification with restriction of spine mobility, the crowned dens syndrome, or spinal stenosis (most commonly seen in the elderly); and (5) rarely periarticular tophus-like nodules.

The knee is the joint most frequently affected in CPPD arthropathy. Other sites include the wrist, shoulder, ankle, elbow, and hands. The temporomandibular joint may be involved. Clinical and radiographic evidence indicates that CPPD deposition is polyarticular in at least two-thirds of patients. When the clinical picture resembles that of slowly progressive osteoarthritis, diagnosis may be difficult. Joint distribution may provide important clues suggesting CPPD disease. For example, primary osteoarthritis less often involves metacarpophalangeal, wrist, elbow, shoulder, or ankle joints. If radiographs or ultrasound reveal punctate and/or linear radiodense deposits within fibrocartilaginous joint menisci or articular hyaline cartilage (*chondrocalcinosis*), the diagnostic likelihood of CPPD disease is further increased. *Definitive diagnosis* requires demonstration of typical rhomboid or rodlike crystals (generally weakly positively birefringent or nonbirefringent with polarized light) in synovial fluid or articular tissue (Fig. 395-2). In the absence of joint effusion or indications to obtain a synovial biopsy, chondrocalcinosis is presumptive of CPPD. One exception is chondrocalcinosis due to CaOx in some patients with chronic renal failure.

Acute attacks of CPPD arthritis may be precipitated by trauma. Rapid diminution of serum calcium concentration, as may occur in severe medical illness or after surgery (especially parathyroidectomy), can also lead to attacks.

In as many as 50% of cases, episodes of CPPD-induced inflammation are associated with low-grade fever and, on occasion, temperatures as high as 40°C (104°F). In such cases, synovial fluid analysis with microbial cultures is essential to rule out the possibility of infection. In fact, infection in a joint with any microcrystalline deposition process can lead to crystal shedding and subsequent synovitis from both crystals and microorganisms. The leukocyte count in synovial fluid in acute CPPD can range from several thousand cells to 100,000 cells/μL, with the mean being about 24,000 cells/μL and the predominant cell being the neutrophil. CPP crystals may be seen inside tissue fragments and fibrin clots and in neutrophils (Fig. 395-2). CPP crystals may coexist with MSU and apatite in some cases.

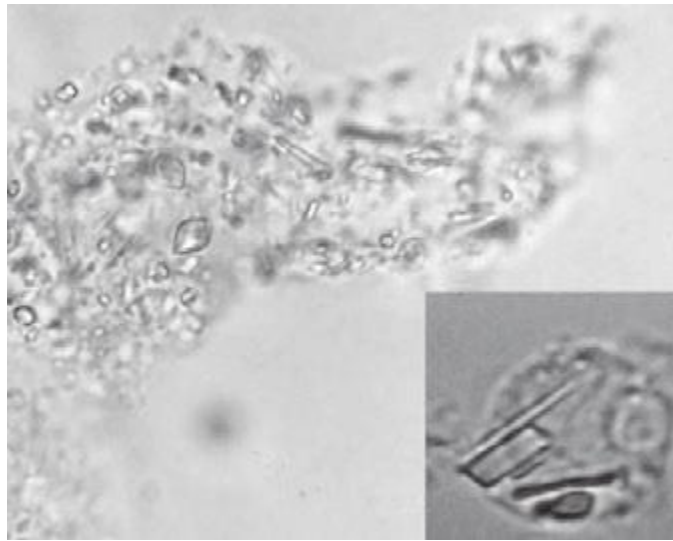


FIGURE 395-2 Intracellular and extracellular calcium pyrophosphate (CPP) crystals, as seen in a fresh preparation of synovial fluid, illustrate rectangular, rod-shaped, and rhomboid crystals that are weakly positively or nonbirefringent crystals (compensated polarized light microscopy; 400 \times).

TABLE 395-3 CONDITIONS ASSOCIATED WITH APATITE DEPOSITION DISEASE

Aging
Osteoarthritis
Hemorrhagic shoulder effusions in the elderly (Milwaukee shoulder)
Destructive arthropathy
Tendinitis, bursitis
Tumoral calcinosis (sporadic cases)
Disease-associated
Hyperparathyroidism
Milk-alkali syndrome
Renal failure/long-term dialysis
Connective tissue diseases (e.g., systemic sclerosis, dermatomyositis, SLE)
Heterotopic calcification after neurologic catastrophes (e.g., stroke, spinal cord injury)
Heredity
Bursitis, arthritis
Tumoral calcinosis
Fibrodysplasia ossificans progressiva

Abbreviation: SLE, systemic lupus erythematosus.

mitosis and markedly increase the release of prostaglandin E₂, various cytokines, and also collagenases and neutral proteases, underscoring the destructive potential of abnormally stimulated synovial lining cells.

CLINICAL MANIFESTATIONS

Periarticular or articular deposits may occur and may be associated with acute reversible inflammation and/or chronic damage to the joint capsule, tendons, bursa, or articular surfaces. The most common sites of apatite deposition include bursae and tendons in and/or around the knees, shoulders, hips, and fingers. Clinical manifestations include asymptomatic radiographic abnormalities, acute synovitis, bursitis, tendinitis, and chronic destructive arthropathy. Although the true incidence of apatite arthritis is not known, 30–50% of patients with osteoarthritis have apatite microcrystals in their synovial fluid. Such crystals frequently can be identified in clinically stable osteoarthritic joints, but they are more likely to come to attention in persons experiencing acute or subacute worsening of joint pain and swelling. The synovial fluid leukocyte count in apatite arthritis is usually low (<2000/ μ L) despite dramatic symptoms, with predominance of mononuclear cells.

DIAGNOSIS

Intra- and/or periarticular calcifications with or without erosive, destructive, or hypertrophic changes may be seen on radiographs (Fig. 395-3). They should be distinguished from the linear calcifications typical of CPPD.

Definitive diagnosis of apatite arthropathy, also called basic calcium phosphate disease, depends on identification of crystals from synovial fluid or tissue (Fig. 395-3). Individual crystals are very small and can be seen only by electron microscopy. Clumps of crystals may appear as 1- to 20- μ m shiny intra- or extracellular nonbirefringent globules or aggregates that stain purplish with Wright's stain and bright red with alizarin red S. Tetracycline binding and other investigative techniques are under consideration as labeling alternatives. Absolute identification depends on electron microscopy with energy-dispersive elemental analysis, x-ray diffraction, infrared spectroscopy, or Raman microspectroscopy, but these techniques usually are not required in clinical diagnosis.

TREATMENT CALCIUM APATITE DEPOSITION DISEASE

Treatment of apatite arthritis or periartthritis is nonspecific. Acute attacks of bursitis or synovitis may be self-limiting, resolving in days to several weeks. Aspiration of effusions and the use of either NSAIDs or oral colchicine for 2 weeks or intra- or periarticular injection of a depot glucocorticoid appear to shorten the duration and

TREATMENT CPPD DISEASE

Untreated acute attacks may last a few days to as long as a month. Treatment by rest, joint aspiration, and NSAIDs or by intraarticular glucocorticoid injection may result in more rapid return to prior status. For patients with frequent recurrent attacks, daily prophylactic treatment with low doses of colchicine may be helpful in decreasing the frequency of the attacks. Severe polyarticular attacks usually require short courses of glucocorticoids or, as recently reported, an IL-1 β antagonist, anakinra. Unfortunately, there is no effective way to remove CPP deposits from cartilage and synovium. Uncontrolled studies suggest that the administration of NSAIDs (with a gastric protective agent if required), hydroxychloroquine, or even methotrexate may be helpful in controlling persistent synovitis. Patients with progressive destructive large-joint arthropathy may require joint replacement.

CALCIUM APATITE DEPOSITION DISEASE

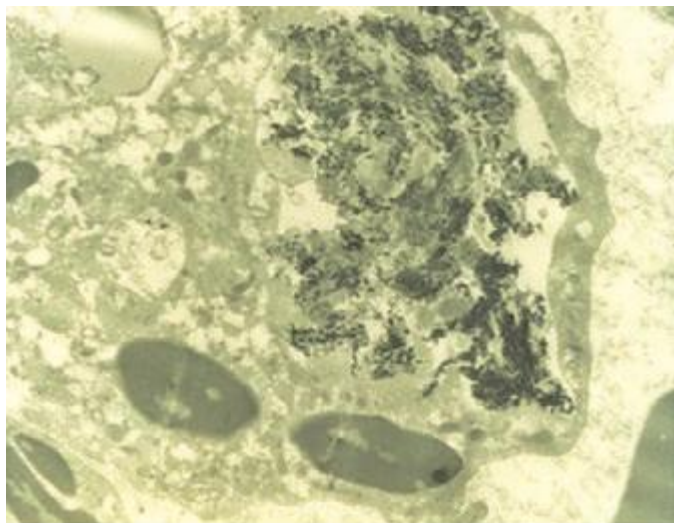
PATHOGENESIS

Apatite is the primary mineral of normal bone and teeth. Abnormal accumulation of basic calcium phosphates, largely carbonate substituted apatite, can occur in areas of tissue damage (dystrophic calcification), hypercalcemic or hyperparathyroid states (metastatic calcification), and certain conditions of unknown cause (Table 395-3). In chronic renal failure, hyperphosphatemia can contribute to extensive apatite deposition both in and around joints. Familial aggregation is rarely seen; no association with *ANKH* mutations has been described thus far. Apatite crystals are deposited primarily on matrix vessels. Incompletely understood alterations in matrix proteoglycans, phosphatases, hormones, and cytokines probably can influence crystal formation.

Apatite aggregates are commonly present in synovial fluid in an extremely destructive chronic arthropathy of the elderly that occurs most often in the shoulders (Milwaukee shoulder) and in a similar process in hips, knees, and erosive osteoarthritis of fingers. Joint destruction is associated with damage to cartilage and supporting structures, leading to instability and deformity. Progression tends to be indolent. Symptoms range from minimal to severe pain and disability that may lead to joint replacement surgery. Whether severely affected patients represent an extreme synovial tissue response to the apatite crystals that are so common in osteoarthritis is uncertain. Synovial lining cell or fibroblast cultures exposed to apatite (or CPP) crystals can undergo



A



B

FIGURE 395-3 **A.** Radiograph showing calcification due to apatite crystals surrounding an eroded joint. **B.** An electron micrograph demonstrates dark needle-shaped apatite crystals within a vacuole of a synovial fluid mononuclear cell (30,000 \times).

intensity of symptoms. Local injection of disodium ethylenediaminetetraacetic acid (EDTA) and SC anakinra have been suggested as effective in single studies of acute calcific tendinitis at the shoulder. Other reports have described that IV gamma globulin, rituximab, calcium channel blockers, or bisphosphonates may help diffuse calcinosis. Periarticular apatite deposits may be resorbed with resolution of attacks. Agents to lower serum phosphate levels may lead to resorption of deposits in renal failure patients receiving hemodialysis. In patients with underlying severe destructive articular changes, response to medical therapy is usually less rewarding.

CAOX DEPOSITION DISEASE

PATHOGENESIS

Primary oxalosis is a rare hereditary metabolic disorder (Chap. 434e). Enhanced production of oxalic acid may result from at least two

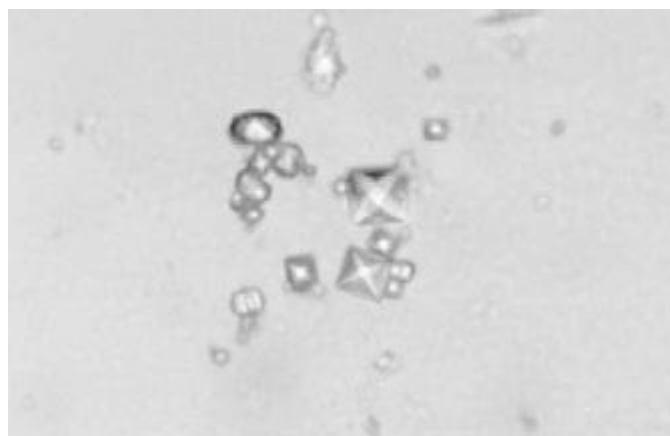


FIGURE 395-4 Bipyramidal and small polymorphic calcium oxalate crystals from synovial fluid are a classic finding in calcium oxalate arthropathy (ordinary light microscopy; 400 \times).

different enzyme defects, leading to hyperoxalemia and deposition of CaOx crystals in tissues. Nephrocalcinosis and renal failure are typical results. Acute and/or chronic CaOx arthritis, peri-arthritis, and bone disease may complicate primary oxalosis during later years of illness.

Secondary oxalosis is more common than the primary disorder. In chronic renal disease, calcium oxalate deposits have long been recognized in visceral organs, blood vessels, bones, and cartilage and are now known to be one of the causes of arthritis in chronic renal failure. Thus far, reported patients have been dependent on long-term hemodialysis or peritoneal dialysis (Chap. 336), and many had received ascorbic acid supplements. Ascorbic acid is metabolized to oxalate, which is inadequately cleared in uremia and by dialysis. Such supplements and foods high in oxalate content usually are avoided in dialysis programs because of the risk of enhancing hyperoxalosis and its sequelae.

CLINICAL MANIFESTATIONS AND DIAGNOSIS

CaOx aggregates can be found in bone, articular cartilage, synovium, and periarticular tissues. From these sites, crystals may be shed, causing acute synovitis. Persistent aggregates of CaOx can, like apatite and CPP, stimulate synovial cell proliferation and enzyme release, resulting in progressive articular destruction. Deposits have been documented in fingers, wrists, elbows, knees, ankles, and feet.

Clinical features of acute CaOx arthritis may not be distinguishable from those due to urate, CPP, or apatite. Radiographs may reveal chondrocalcinosis or soft tissue calcifications. CaOx-induced synovial effusions are usually noninflammatory, with <2000 leukocytes/ μ L, or mildly inflammatory. Neutrophils or mononuclear cells can predominate. CaOx crystals have a variable shape and variable birefringence to polarized light. The most easily recognized forms are bipyramidal, have strong birefringence (Fig. 395-4), and stain with alizarin red S.

TREATMENT CALCIUM OXALATE DEPOSITION DISEASE

Treatment of CaOx arthropathy with NSAIDs, colchicine, intra-articular glucocorticoids, and/or an increased frequency of dialysis has produced only slight improvement. In primary oxalosis, liver transplantation has induced a significant reduction in crystal deposits (Chap. 434e).

ACKNOWLEDGMENT

This chapter has been revised for this and the previous two editions from an original version written by Antonio Reginato, MD, in earlier editions of Harrison's Principles of Internal Medicine.

396 Fibromyalgia

Leslie J. Crofford

DEFINITION

Fibromyalgia (FM) is characterized by chronic widespread musculoskeletal pain and tenderness. Although FM is defined primarily as a pain syndrome, patients also commonly report associated neuropsychological symptoms of fatigue, unrefreshing sleep, cognitive dysfunction, anxiety, and depression. Patients with FM have an increased prevalence of other syndromes associated with pain and fatigue, including chronic fatigue syndrome ([Chap. 464e](#)), temporomandibular disorder, chronic headaches, irritable bowel syndrome, interstitial cystitis/painful bladder syndrome, and other pelvic pain syndromes. Available evidence implicates the central nervous system as key to maintaining pain and other core symptoms of FM and related conditions. The presence of FM is associated with substantial negative consequences for physical and social functioning.

EPIDEMIOLOGY



In clinical settings, a diagnosis of FM is made in ~2% of the population and is far more common in women than in men, with a ratio of ~9:1. However, in population-based survey studies worldwide, the prevalence rate is ~2–5%, with a female-to-male ratio of only 2–3:1 and with some variability depending on the method of ascertainment. The prevalence data are similar across socioeconomic classes. Cultural factors may play a role in determining whether patients with FM symptoms seek medical attention; however, even in cultures in which secondary gain is not expected to play a significant role, the prevalence of FM remains in this range.

CLINICAL MANIFESTATIONS

Pain and Tenderness At presentation, patients with FM most commonly report “pain all over.” These patients have pain that is typically both above and below the waist on both sides of the body and involves the axial skeleton (neck, back, or chest). The pain attributable to FM is poorly localized, difficult to ignore, severe in its intensity, and associated with a reduced functional capacity. For a diagnosis of FM, pain

should have been present most of the day on most days for at least 3 months.

The clinical pain of FM is associated with increased evoked pain sensitivity. In clinical practice, this elevated sensitivity may be determined by a tender-point examination in which the examiner uses the thumbnail to exert pressure of ~4 kg/m² (or the amount of pressure leading to blanching of the tip of the thumbnail) on well-defined musculotendinous sites ([Fig. 396-1](#)). Previously, the classification criteria of the American College of Rheumatology required that 11 of 18 sites be perceived as painful for a diagnosis of FM. In practice, tenderness is a continuous variable, and strict application of a categorical threshold for diagnostic specifics is not necessary. Newer criteria eliminate the need for tender points and focus instead on clinical symptoms of widespread pain and neuropsychological symptoms. The newer criteria perform well in a clinical setting in comparison to the older, tender-point criteria. However, it appears that when the new criteria are applied to populations, the result is an increase in prevalence of FM and a change in the sex ratio (see “Epidemiology,” earlier).

Patients with FM often have peripheral pain generators that are thought to serve as triggers for the more widespread pain attributed to central nervous system factors. Potential pain generators such as arthritis, bursitis, tendinitis, neuropathies, and other inflammatory or degenerative conditions should be identified by history and physical examination. More subtle pain generators may include joint hypermobility and scoliosis. In addition, patients may have chronic myalgias triggered by infectious, metabolic, or psychiatric conditions that can also serve as triggers for the development of FM. These conditions are often identified in the differential diagnosis of patients with FM, and a major challenge is to distinguish the ongoing activity of a triggering condition from FM that is occurring as a consequence of a comorbid condition and that should itself be treated.

Neuropsychological Symptoms In addition to widespread pain, FM patients typically report fatigue, stiffness, sleep disturbance, cognitive dysfunction, anxiety, and depression. These symptoms are present to varying degrees in most FM patients but are not present in every patient or at all times in a given patient. Relative to pain, such symptoms may, however, have an equal or even greater impact on function and quality of life. Fatigue is highly prevalent in patients under primary care who ultimately are diagnosed with FM. Pain, stiffness, and fatigue often

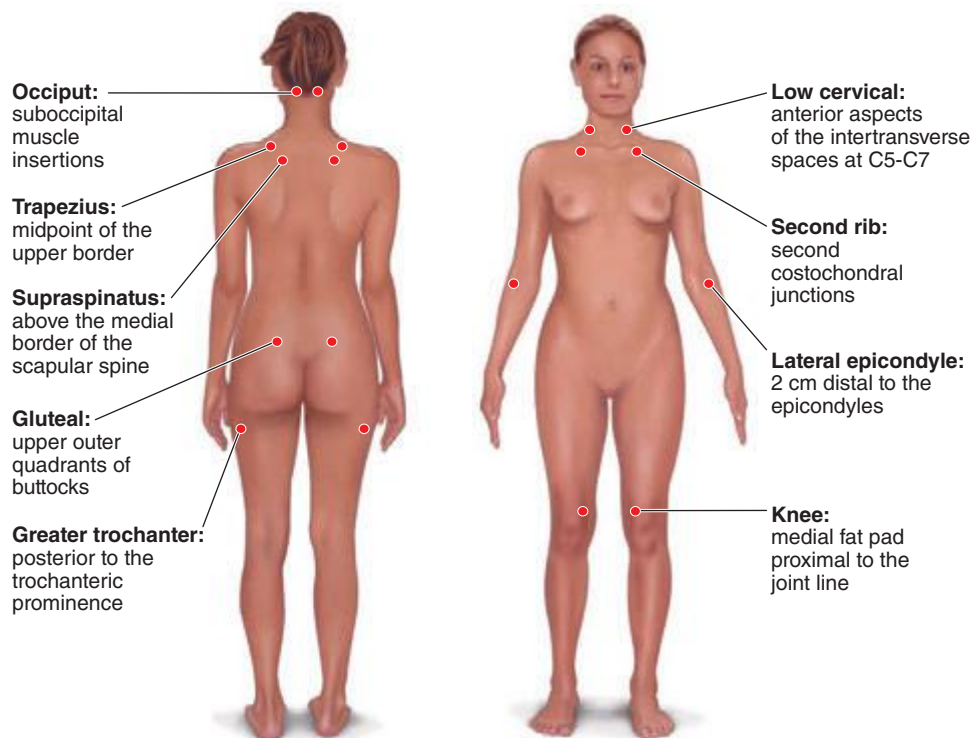


FIGURE 396-1 Tender-point assessment in patients with fibromyalgia. (Figure created using data from F Wolfe et al. *Arthritis Care Res* 62:600, 2010.)

are worsened by exercise or unaccustomed activity (postexertional malaise). The sleep complaints include difficulty falling asleep, difficulty staying asleep, and early-morning awakening. Regardless of the specific complaint, patients awake feeling unrefreshed. Patients with FM may meet criteria for restless legs syndrome and sleep-disordered breathing; frank sleep apnea can also be documented. Cognitive issues are characterized as slowness in processing, difficulties with attention or concentration, problems with word retrieval, and short-term memory loss. Studies have demonstrated altered cognitive function in these domains in patients with FM, though speed of processing is age-appropriate. Symptoms of anxiety and depression are common, and the lifetime prevalence of mood disorders in patients with FM approaches 80%. Although depression is neither necessary nor sufficient for the diagnosis of FM, it is important to screen for major depressive disorders by querying for depressed mood and anhedonia. Analysis of genetic factors that are likely to predispose to FM reveals shared neurobiologic pathways with mood disorders, providing the basis for comorbidity (see later in this chapter).

Overlapping Syndromes Because FM can overlap in presentation with other chronic pain conditions, review of systems often reveals headaches, facial/jaw pain, regional myofascial pain particularly involving the neck or back, and arthritis. Visceral pain involving the gastrointestinal tract, bladder, and pelvic or perineal region is often present as well. Patients may or may not meet defined criteria for specific syndromes. It is important for patients to understand that shared pathways may mediate symptoms and that treatment strategies effective for one condition may help with global symptom management.

Comorbid Conditions FM is often comorbid with chronic musculoskeletal, infectious, metabolic, or psychiatric conditions. Whereas FM affects only 2–5% of the general population, it occurs in 20% or more of patients with degenerative or inflammatory rheumatic disorders, likely because these conditions serve as peripheral pain generators to alter central pain-processing pathways. Similarly, chronic infectious, metabolic, or psychiatric diseases associated with musculoskeletal pain can mimic FM and/or serve as a trigger for the development of FM. It is particularly important for clinicians to be sensitive to pain management of these comorbid conditions so that when FM emerges—characterized by pain outside the boundaries of what could reasonably be explained by the triggering condition, development of neuropsychological symptoms, or tenderness on physical examination—treatment of central pain processes will be undertaken as opposed to a continued focus on treatment of peripheral or inflammatory causes of pain.

Psychosocial Considerations Symptoms of FM often have their onset and are exacerbated during periods of high-level real or perceived stress. This pattern may reflect an interaction among central stress physiology, vigilance or anxiety, and central pain-processing pathways. An understanding of current psychosocial stressors will aid in patient management, as many factors that exacerbate symptoms cannot be addressed by pharmacologic approaches. Furthermore, there is a high prevalence of exposure to previous interpersonal and other forms of violence in patients with FM and related conditions. If post-traumatic stress disorder is an issue, the clinician should be aware of it and consider treatment options.

Functional Impairment It is crucial to evaluate the impact of FM symptoms on function and role fulfillment. In defining the success of a management strategy, improved function is a key measure. Functional assessment should include physical, mental, and social domains. A recognition of the ways in which role functioning falls short will be helpful in the establishment of treatment goals.

DIFFERENTIAL DIAGNOSIS

Because musculoskeletal pain is such a common complaint, the differential diagnosis of FM is broad. **Table 396-1** lists some of the more common conditions that should be considered. Patients with inflammatory causes for widespread pain should be identifiable on the basis of specific history, physical findings, and laboratory or radiographic tests.

TABLE 396-1 COMMON CONDITIONS IN THE DIFFERENTIAL DIAGNOSIS OF FIBROMYALGIA

Inflammatory
Polymyalgia rheumatica
Inflammatory arthritis: rheumatoid arthritis, spondyloarthritis
Connective tissue diseases: systemic lupus erythematosus, Sjögren's syndrome
Infectious
Hepatitis C
HIV infection
Lyme disease
Parvovirus B19 infection
Epstein-Barr virus infection
Noninflammatory
Degenerative joint/spine/disk disease
Myofascial pain syndromes
Bursitis, tendinitis, repetitive strain injuries
Endocrine
Hypo- or hyperthyroidism
Hyperparathyroidism
Neurologic Diseases
Multiple sclerosis
Neuropathic pain syndromes
Psychiatric Disease
Major depressive disorder
Drugs
Statins
Aromatase inhibitors

LABORATORY OR RADIOGRAPHIC TESTING

Routine laboratory and radiographic tests yield normal results in FM. Thus diagnostic testing is focused on exclusion of other diagnoses and evaluation for pain generators or comorbid conditions (**Table 396-2**). Most patients with new chronic widespread pain should be assessed for the most common entities in the differential diagnosis. Radiographic testing should be used sparingly and only for diagnosis of inflammatory arthritis. After the patient has been evaluated thoroughly, repeat testing is discouraged unless the symptom complex changes. Particularly to be discouraged is advanced imaging (MRI) of the spine unless there are features suggesting inflammatory spine disease or neurologic symptoms.

GENETICS AND PHYSIOLOGY


 As in most complex diseases, it is likely that a number of genes contribute to vulnerability to the development of FM. To date, these genes appear to be in pathways controlling pain and stress

TABLE 396-2 LABORATORY AND RADIOGRAPHIC TESTING IN PATIENTS WITH FIBROMYALGIA SYMPTOMS

Routine
Erythrocyte sedimentation rate (ESR) or C-reactive protein (CRP)
Complete blood count (CBC)
Thyroid-stimulating hormone (TSH)
Guided by History and Physical Examination
Complete metabolic panel
Antinuclear antibody (ANA)
Anti-SSA (anti-Sjögren's syndrome A) and anti-SSB
Rheumatoid factor and anti-cyclic citrullinated peptide (anti-CCP)
Creatine phosphokinase (CPK)
Viral and bacterial serologies
Spine and joint radiographs

Source: LM Arnold et al: *J Women's Health* 21:231, 2012; MA Fitzcharles et al: *J Rheumatol* 40:1388, 2013.

2240 responses. Some of the genetic underpinnings of FM are shared across other chronic pain conditions. Genes associated with metabolism, transport, and receptors of serotonin and other monoamines have been implicated in FM and overlapping conditions. Genes associated with other pathways involved in pain transmission have also been described as vulnerability factors for FM. Taken together, the pathways in which polymorphisms have been identified in FM patients further implicate central factors in mediation of the physiology that leads to the clinical manifestations of FM.

Psychophysical testing of patients with FM has demonstrated altered sensory afferent pain processing and impaired descending noxious inhibitory control leading to hyperalgesia and allodynia. Functional MRI and other research imaging procedures clearly demonstrate activation of the brain regions involved in the experience of pain in response to stimuli that are innocuous in study participants without FM. Pain perception in FM patients is influenced by the emotional and cognitive dimensions, such as catastrophizing and perceptions of control, providing a solid basis for recommendations for cognitive and behavioral treatment strategies.

APPROACH TO THE PATIENT: Fibromyalgia

FM is common and has an extraordinary impact on the patient's function and health-related quality of life. However, its symptoms and impact can be managed effectively by physicians and other health professionals. Developing a partnership with patients is essential for improving the outcome of FM, with a goal of understanding the factors involved, implementing a treatment strategy, and choosing appropriate nonpharmacologic and pharmacologic treatments.

TREATMENT FIBROMYALGIA

NONPHARMACOLOGIC TREATMENT

Patients with chronic pain, fatigue, and other neuropsychological symptoms require a framework for understanding the symptoms that have such an important impact on their function and quality of life. Explaining the genetics, triggers, and physiology of FM can be an important adjunct in relieving associated anxiety and in reducing the overall cost of health care resources. In addition, patients must be educated regarding expectations for treatment. The physician should focus on improved function and quality of life rather than elimination of pain. Illness behaviors, such as frequent physician visits, should be discouraged and behaviors that focus on improved function strongly encouraged.

Treatment strategies should include physical conditioning, with encouragement to begin at low levels of aerobic exercise and to proceed with slow but consistent advancement. Patients who have been physically inactive or who report postexertional malaise may do best in supervised or water-based programs at the start. Activities that promote improved physical function with relaxation, such as yoga and Tai Chi, may also be helpful. Strength training may be recommended after patients reach their aerobic goals. Exercise programs are helpful in reducing tenderness and enhancing self-efficacy. Cognitive-behavioral strategies to improve sleep hygiene and reduce illness behaviors can also be helpful in management.

PHARMACOLOGIC APPROACHES

It is essential for the clinician to treat any comorbid triggering condition and to clearly delineate for the patient the treatment goals for each medication. For example, glucocorticoids or nonsteroidal anti-inflammatory drugs may be useful for management of inflammatory triggers but are not effective against FM-related symptoms. At present, the treatment approaches that have proved most successful in FM patients target afferent or descending pain pathways. [Table 396-3](#) lists the drugs with demonstrated effectiveness. It should be emphasized

TABLE 396-3 PHARMACOLOGIC AGENTS EFFECTIVE FOR TREATMENT OF FIBROMYALGIA

Antidepressants: balanced serotonin–norepinephrine reuptake inhibitors
Amitriptyline ^a
Duloxetine ^{b,c}
Milnacipran ^{b,c}
Anticonvulsants: ligands of the alpha-2-delta subunit of voltage-gated calcium channels
Gabapentin
Pregabalin ^b

^aRA Moore et al: *Cochrane Database Syst Rev* 12:CD008242, 2012. ^bApproved by the U.S. Food and Drug Administration. ^cW Hauser et al: *Cochrane Database Syst Rev* 1: CD010292, 2013.

Source: LM Arnold: *Arthritis Rheum* 56:1336, 2007.

strongly that opioid analgesics are to be avoided in patients with FM. These agents have no demonstrated efficacy in FM and are associated with opioid-induced hyperalgesia that can worsen both symptoms and function. Use of single agents to treat multiple symptom domains is strongly encouraged. For example, if a patient's symptom complex is dominated by pain and sleep disturbance, use of an agent that exerts both analgesic and sleep-promoting effects is desirable. These agents include sedating antidepressants such as amitriptyline and alpha-2-delta ligands such as gabapentin and pregabalin. For patients whose pain is associated with fatigue, anxiety, or depression, drugs that have both analgesic and antidepressant/anxiolytic effects, such as duloxetine or milnacipran, may be the best first choice.

397 Arthritis Associated with Systemic Disease, and Other Arthritides

Carol A. Langford, Brian F. Mandell

ARTHRITIS ASSOCIATED WITH SYSTEMIC DISEASE

ARTHROPATHY OF ACROMEGALY

Acromegaly is the result of excessive production of growth hormone by an adenoma in the anterior pituitary gland ([Chap. 403](#)). The excessive secretion of growth hormone along with insulin-like growth factor I stimulates proliferation of cartilage, periarticular connective tissue, and bone, resulting in several musculoskeletal problems, including osteoarthritis, back pain, muscle weakness, and carpal tunnel syndrome.

Osteoarthritis is a common feature, most often affecting the knees, shoulders, hips, and hands. Single or multiple joints may be affected. Hypertrophy of cartilage initially produces radiographic widening of the joint space. The newly synthesized cartilage is abnormally susceptible to fissuring, ulceration, and destruction. Ligament laxity of joints further contributes to the development of osteoarthritis. Cartilage degrades, the joint space narrows, and subchondral sclerosis and osteophytes develop. Joint examination reveals crepitus and laxity. Joint fluid is noninflammatory. Calcium pyrophosphate dihydrate crystals are found in the cartilage in some cases of acromegaly arthropathy and, when shed into the joint, can elicit attacks of pseudogout. Chondrocalcinosis may be observed on radiographs. Back pain is extremely common, perhaps as a result of spine hypermobility. Spine radiographs show normal or widened intervertebral disk spaces, hypertrophic anterior osteophytes, and ligament calcification. The latter changes are similar to those observed in patients with diffuse idiopathic skeletal hyperostosis. Dorsal kyphosis in conjunction with elongation of the ribs contributes to the development of the barrel chest seen in acromegalic patients. The hands and feet

become enlarged as a result of soft tissue proliferation. The fingers are thickened and have spadelike distal tufts. One-third of patients have a thickened heel pad. Approximately 25% of patients exhibit Raynaud's phenomenon. Carpal tunnel syndrome occurs in about half of patients. The median nerve is compressed by excess connective tissue in the carpal tunnel. Patients with acromegaly may develop proximal muscle weakness, which is thought to be caused by the effect of growth hormone on muscle. Serum muscle enzyme levels and electromyographic findings are normal. Muscle biopsy specimens contain muscle fibers of varying size without inflammation.

ARTHROPATHY OF HEMOCHROMATOSIS

Hemochromatosis is a disorder of iron storage. Absorption of excessive amounts of iron from the intestine leads to iron deposition in parenchymal cells, which results in impairment of organ function (Chap. 428). Symptoms of hemochromatosis usually begin between the ages of 40 and 60 but can appear earlier. Arthropathy, which occurs in 20–40% of patients, usually begins after the age of 50 and may be the first clinical feature of hemochromatosis. The arthropathy is an osteoarthritis-like disorder affecting the small joints of the hands and later the larger joints, such as knees, ankles, shoulders, and hips. The second and third metacarpophalangeal joints of both hands are often the first and most prominent joints affected; this clinical picture may provide an important clue to the possibility of hemochromatosis because these joints are not predominantly affected by “routine” osteoarthritis. Patients experience some morning stiffness and pain with use of involved joints. The affected joints are enlarged and mildly tender. Radiographs show narrowing of the joint space, subchondral sclerosis, subchondral cysts, and juxtaarticular proliferation of bone. Hooklike osteophytes are seen in up to 20% of patients; although they are regarded as a characteristic feature of hemochromatosis, they can also occur in osteoarthritis and are not disease specific. The synovial fluid is noninflammatory. The synovium shows mild to moderate proliferation of iron-containing lining cells, fibrosis, and some mononuclear cell infiltration. In approximately half of patients, there is evidence of calcium pyrophosphate deposition disease, and some patients late in the course of disease experience episodes of acute pseudogout (Chap. 395). An early diagnosis is suggested by high serum transferrin saturation, which is more sensitive than ferritin elevation.

Iron may damage the articular cartilage in several ways. Iron catalyzes superoxide-dependent lipid peroxidation, which may play a role in joint damage. In animal models, ferric iron has been shown to interfere with collagen formation and increase the release of lysosomal enzymes from cells in the synovial membrane. Iron inhibits synovial tissue pyrophosphatase in vitro and therefore may inhibit pyrophosphatase in vivo, resulting in chondrocalcinosis.

TREATMENT ARTHROPATHY OF HEMOCHROMATOSIS

The treatment of hemochromatosis is repeated phlebotomy. Unfortunately, this treatment has little effect on established arthritis, which, along with chondrocalcinosis, may progress. Symptom-based treatment of the arthritis consists of administration of acetaminophen and nonsteroidal anti-inflammatory drugs (NSAIDs), as tolerated. Acute pseudogout attacks are treated with high doses of an NSAID or a short course of glucocorticoids. Hip or knee total joint replacement has been successful in advanced disease.

HEMOPHILIC ARTHROPATHY

Hemophilia is a sex-linked recessive genetic disorder characterized by the absence or deficiency of factor VIII (hemophilia A, or classic hemophilia) or factor IX (hemophilia B, or Christmas disease) (Chap. 141). Hemophilia A constitutes 85% of cases. Spontaneous hemarthrosis is a common problem with both types of hemophilia and can lead to a deforming arthritis. The frequency and severity of hemarthrosis are related to the degree of clotting factor deficiency. Hemarthrosis is not common in other disorders of coagulation such as von Willebrand disease, factor V deficiency, warfarin therapy, or thrombocytopenia.

Hemarthrosis occurs after 1 year of age, when a child begins to walk and run. In order of frequency, the joints most commonly affected are the knees, ankles, elbows, shoulders, and hips. Small joints of the hands and feet are occasionally involved.

In the initial stage of arthropathy, hemarthrosis produces a warm, tensely swollen, and painful joint. The patient holds the affected joint in flexion and guards against any movement. Blood in the joint remains liquid because of the absence of intrinsic clotting factors and the absence of tissue thromboplastin in the synovium. The synovial blood is resorbed over a period of ≥ 1 week, with the precise interval depending on the size of the hemarthrosis. Joint function usually returns to normal or baseline in ~ 2 weeks. Low-grade temperature elevation may accompany hemarthrosis, but a fever $>101^\circ\text{F}$ (38.3°C) warrants concern about infection.

Recurrent hemarthrosis may result in chronic arthritis. The involved joints remain swollen, and flexion deformities develop. Joint motion may be restricted and function severely limited. Restricted joint motion or laxity with subluxation is a feature of end-stage disease.

Bleeding into muscle and soft tissue also causes musculoskeletal dysfunction. When bleeding into the iliopsoas muscle occurs, the hip is held in flexion because of the pain, resulting in a hip flexion contracture. Rotation of the hip is preserved, which distinguishes this problem from hemarthrosis or other causes of hip synovitis. Expansion of the hematoma may place pressure on the femoral nerve, resulting in femoral neuropathy. Hemorrhage into a closed compartment space, such as the calf or the volar compartment in the forearm, can result in muscle necrosis, neuropathy, and flexion deformities of the ankles, wrists, and fingers. When bleeding involves periosteum or bone, a painful pseudotumor forms. These pseudotumors occur distal to the elbows or knees in children and improve with treatment of hemophilia. Surgical removal is indicated if the pseudotumor continues to enlarge. In adults, pseudotumors develop in the femur and pelvis and are usually refractory to treatment. When bleeding occurs in muscle, cysts may develop within the muscle. Needle aspiration of a cyst is contraindicated because this procedure can induce further bleeding; however, if the cyst becomes secondarily infected, drainage may be necessary (after factor repletion).

Septic arthritis is rare in hemophilia and is difficult to distinguish from acute hemarthrosis on physical examination. If there is serious suspicion of an infected joint, the joint should be aspirated immediately, the fluid cultured, and treatment with broad-spectrum antibiotics administered, with coverage for microorganisms including *Staphylococcus*, until culture results become available. Clotting-factor deficiency should be corrected before arthrocentesis to minimize the risk of traumatic bleeding.

Radiographs of joints reflect the stage of disease. In early stages, there is only capsule distention; later, juxtaarticular osteopenia, marginal erosions, and subchondral cysts develop. Late in the disease, the joint space is narrowed and there is bony overgrowth similar to that in osteoarthritis.

TREATMENT HEMARTHROSIS

The treatment of musculoskeletal bleeding is initiated with the immediate infusion of factor VIII or IX at the first sign of joint or muscle hemorrhage. Patients who have developed factor inhibitors are at elevated risk for joint damage and may benefit from receiving recombinant activated factor VII or activated prothrombin complex concentrate. The joint should be rested in a position of forced extension, as tolerated, to avoid contracture. Analgesia should be provided; nonselective NSAIDs, which can diminish platelet function, should be avoided if possible. Selective cyclooxygenase-2 inhibitors do not interfere with platelet function, although cardiovascular and gastrointestinal risks must still be weighed. Synovectomy—open or arthroscopic—may be attempted in patients with chronic symptomatic synovial proliferation and recurrent hemarthrosis, although hypertrophied synovium is highly vascular and subject to bleeding. Both types of synovectomy reduce the number of hemarthroses. Open surgical synovectomy, however, is associated with some loss

2242 of range of motion. Both require aggressive prophylaxis against bleeding. Radiosynovectomy with either yttrium 90 silicate or phosphorus 31 colloid has been effective and may be attempted when surgical synovectomy is not practical. Total joint replacement is indicated for severe joint destruction and incapacitating pain.

ARTHROPATHIES ASSOCIATED WITH HEMOGLOBINOPATHIES

Sickle Cell Disease Sickle cell disease (Chap. 127) is associated with several musculoskeletal abnormalities (Table 397-1). Children under the age of 5 years may develop diffuse swelling, tenderness, and warmth of the hands and feet lasting 1–3 weeks. This condition, referred to as *sickle cell dactylitis* or *hand-foot syndrome*, has also been observed in sickle cell thalassemia. Dactylitis is believed to result from infarction of the bone marrow and cortical bone leading to periostitis and soft tissue swelling. Radiographs show periosteal elevation, subperiosteal new-bone formation, and areas of radiolucency and increased density involving the metacarpals, metatarsals, and proximal phalanges. These bone changes disappear after several months. The syndrome leaves little or no residual damage. Because hematopoiesis ceases in the small bones of the hands and feet with age, the syndrome is rarely seen after age 5.

Sickle cell crisis is associated with periarticular pain and occasionally with joint effusions. The joint and periarticular area are warm and tender. Knees and elbows are most often affected, but other joints can be involved. Joint effusions are usually noninflammatory. Acute synovial infarction can cause a sterile effusion with high neutrophil counts in synovial fluid. Synovial biopsies have shown mild lining-cell proliferation and microvascular thrombosis with infarctions. Scintigraphic studies have shown decreased marrow uptake adjacent to the involved joint. The treatment for sickle cell crisis is detailed in Chap. 127.

Patients with sickle cell disease seem predisposed to osteomyelitis, which commonly involves the long tubular bones (Chap. 158); *Salmonella* is a particularly common cause (Chap. 190). Radiographs of the involved site initially show periosteal elevation, with subsequent disruption of the cortex. Treatment of the infection results in healing of the bone lesion. In addition, sickle cell disease is associated with bone infarction resulting from vaso-occlusion secondary to the sickling of red cells. Bone infarction also occurs in hemoglobin sickle cell disease and sickle cell thalassemia (Chap. 127). The bone pain in sickle cell crisis is due to infarction of bone and bone marrow. In children, infarction of the epiphyseal growth plate interferes with normal growth of the affected extremity. Radiographically, infarction of the bone cortex results in periosteal elevation and irregular thickening of the bone cortex. Infarction in the bone marrow leads to lysis, fibrosis, and new bone formation. Clinical distinction between osteomyelitis and bone infarctions can be difficult; imaging can be helpful.

Avascular necrosis of the head of the femur occurs in ~5% of patients. It also occurs in the humeral head and less commonly in the distal femur, tibial condyles, distal radius, vertebral bodies, and other juxtaarticular sites. Irregularity of the femoral head and other articular surfaces often results in degenerative joint disease. Radiography of the affected joint may show patchy radiolucency and density followed by flattening of the bone. MRI is a sensitive technique for detecting early avascular necrosis as well as bone infarction elsewhere. Total hip replacement and placement of prostheses in other joints may improve function and relieve joint pain in these patients.

Septic arthritis is occasionally encountered in sickle cell disease (Chap. 157). Multiple joints may be infected. Joint infection may result from bacteremia due to splenic dysfunction or from contiguous

osteomyelitis. The more common microorganisms include *Staphylococcus aureus*, *Streptococcus*, and *Salmonella*. *Salmonella* does not cause septic arthritis as frequently as it causes osteomyelitis. Acute gouty arthritis is uncommon in sickle cell disease, even though 40% of patients are hyperuricemic. However, it may occur in patients generally not expected to get gout (young patients, female patients). Hyperuricemia is due to overproduction of uric acid secondary to increased red cell turnover as well as suboptimal renal excretion. Attacks may be polyarticular, and diagnostic arthrocentesis should be performed to distinguish infection from gout or synovial infarction.

The bone marrow hyperplasia in sickle cell disease results in widening of the medullary cavities, thinning of the cortices, and coarse trabeculations and central cupping of the vertebral bodies. These changes are also seen to a lesser degree in hemoglobin sickle cell disease and sickle cell thalassemia. In normal individuals red marrow is located mostly in the axial skeleton, but in sickle cell disease red marrow is found in the bones of the extremities and even in the tarsal and carpal bones. Vertebral compression may lead to dorsal kyphosis, and softening of the bone in the acetabulum may result in protrusio acetabuli.

Thalassemia A congenital disorder of hemoglobin synthesis, β thalassemia is characterized by impaired production of β chains (Chap. 127). Bone and joint abnormalities occur in β thalassemia, being most common in the major and intermedia groups. In one study, ~50% of patients with β thalassemia had evidence of symmetric ankle arthropathy characterized by a dull aching pain that was aggravated by weight bearing. The onset came most often in the second or third decade of life. The degree of ankle pain in these patients varied. Some patients experienced self-limited ankle pain that occurred only after strenuous physical activity and lasted several days or weeks. Other patients had chronic ankle pain that became worse with walking. Symptoms eventually abated in a few patients. Compression of the ankle, calcaneus, or forefoot was painful in some patients. Synovial fluid from two patients was noninflammatory. Radiographs of the ankle showed osteopenia, widened medullary spaces, thin cortices, and coarse trabeculations—findings that are largely the result of bone marrow expansion. The joint space was preserved. Specimens of bone from three patients revealed osteomalacia, osteopenia, and microfractures. Increased numbers of osteoblasts as well as increased foci of bone resorption were present on the bone surface. Iron staining was found in the bone trabeculae, in osteoid, and in the cement line. Synovium showed hyperplasia of lining cells, which contained deposits of hemosiderin. This arthropathy was considered to be related to the underlying bone pathology. The role of iron overload or abnormal bone metabolism in the pathogenesis of this arthropathy is not known. The arthropathy was treated with analgesics and splints. Patients also received transfusions to decrease hematopoiesis and bone marrow expansion.

In patients with β -thalassemia major and β -thalassemia intermedia, other joints are also involved, including the knees, hips, and shoulders. Acquired hemochromatosis with arthropathy has been described in a patient with thalassemia. Gouty arthritis and septic arthritis can occur. Avascular necrosis is not a feature of thalassemia because there is no sickling of red cells leading to thrombosis and infarction.

β -Thalassemia minor (also known as *β -thalassemia trait*) is likewise associated with joint manifestations. Chronic seronegative oligoarthritis affecting predominantly ankles, wrists, and elbows has been described; the affected patients had mild persistent synovitis without large effusions or joint erosions. Recurrent episodes of acute asymmetric arthritis have also been reported; episodes last <1 week and may affect the knees, ankles, shoulders, elbows, wrists, and metacarpal phalangeal joints. The mechanism underlying this arthropathy is unknown. Treatment with NSAIDs is not particularly effective.

MUSCULOSKELETAL DISORDERS ASSOCIATED WITH HYPERLIPIDEMIA

(See also Chap. 421) Musculoskeletal or cutaneous manifestations may be the first clinical indication of a specific hereditary disorder of lipoprotein metabolism. Patients with familial hypercholesterolemia (previously referred to as *type II hyperlipoproteinemia*) may have recurrent migratory polyarthritis involving the knees and other large

TABLE 397-1 MUSCULOSKELETAL ABNORMALITIES IN SICKLE CELL DISEASE

Sickle Cell Dactylitis	Avascular Necrosis
Joint effusions in sickle cell crises	Bone changes secondary to marrow hyperplasia
Osteomyelitis	Septic arthritis
Infarction of bone	Gouty arthritis
Infarction of bone marrow	

peripheral joints and, to a lesser degree, peripheral small joints. Pain ranges from moderate to incapacitating. The involved joints can be warm, erythematous, swollen, and tender. Arthritis usually has a sudden onset, lasts from a few days to 2 weeks, and does not cause joint damage. Episodes may suggest acute gout attacks. Several attacks occur per year. Synovial fluid from involved joints is not inflammatory and contains few white cells and no crystals. Joint involvement may actually represent inflammatory peri-arthritis or peritendinitis and not true arthritis. The recurrent, transient nature of the arthritis may suggest rheumatic fever, especially because patients with hyperlipoproteinemia may have an elevated erythrocyte sedimentation rate and elevated antistreptolysin O titers (the latter being quite common). Attacks of tendinitis, including the large Achilles and patellar tendons, may come on gradually and last only a few days or may be acute as described above. Patients may be asymptomatic between attacks. Achilles tendinitis and other joint manifestations often precede the appearance of xanthomas and may be the first clinical indication of hyperlipoproteinemia. Attacks of tendinitis may follow treatment with a lipid-lowering drug. Over time, patients may develop tendinous xanthomas in the Achilles, patellar, and extensor tendons of the hands and feet. Xanthomas have also been reported in the peroneal tendon, the plantar aponeurosis, and the periosteum overlying the distal tibia. These xanthomas are located within tendon fibers. Tuberos xanthomas are soft subcutaneous masses located over the extensor surfaces of the elbows, knees, and hands as well as on the buttocks. They appear during childhood in homozygous patients and after the age of 30 in heterozygous patients. Patients with elevated plasma levels of very-low-density lipoprotein (VLDL) and triglycerides (previously referred to as *type IV hyperlipoproteinemia*) may also have a mild inflammatory arthritis affecting large and small peripheral joints, usually in an asymmetric pattern, with only a few joints involved at a time. The onset of arthritis usually comes in middle age. Arthritis may be persistent or recurrent, with episodes lasting a few days or weeks. Some patients may experience severe joint pain or morning stiffness. Joint tenderness and periarticular hyperesthesia may also be present, as may synovial thickening. Joint fluid is usually noninflammatory and without crystals but may have increased white blood cell counts with predominantly mononuclear cells. Radiographs may show juxtaarticular osteopenia and cystic lesions. Large bone cysts have been noted in a few patients. Xanthoma and bone cysts are also observed in other lipoprotein disorders. The pathogenesis of arthritis in patients with familial hypercholesterolemia or with elevated levels of VLDL and triglycerides is not well understood. NSAIDs or analgesics usually provide adequate relief of symptoms when used on an as-needed basis.

Patients may improve clinically as they are treated with lipid-lowering agents; however, patients treated with an HMG-CoA reductase inhibitor may experience myalgias, and a few patients develop myopathy, myositis, or even rhabdomyolysis. Patients who develop myositis during statin therapy may be susceptible to this adverse effect because of an underlying muscle disorder and should be reevaluated after discontinuation of the drug. Myositis has also been reported with the use of niacin (Chap. 388) but is less common than myalgias.

Musculoskeletal syndromes have not clearly been associated with the more common mixed hyperlipidemias seen in general practice.

OTHER ARTHRITIDES

NEUROPATHIC JOINT DISEASE

Neuropathic joint disease (Charcot joint) is a progressive destructive arthritis associated with loss of pain sensation, proprioception, or both. Normal muscular reflexes that modulate joint movement are impaired. Without these protective mechanisms, joints are subjected to repeated trauma, resulting in progressive cartilage and bone damage. Today, diabetes mellitus is the most frequent cause of neuropathic joint disease (Fig. 397-1). A variety of other disorders are associated with neuropathic arthritis, including tabes dorsalis, leprosy, yaws, syringomyelia, meningomyelocele, congenital indifference to pain, peroneal muscular atrophy (Charcot-Marie-Tooth disease), and amyloidosis. An arthritis resembling neuropathic joint disease has been reported in



FIGURE 397-1 Charcot arthropathy associated with diabetes mellitus. Lateral foot radiograph demonstrating complete loss of the arch due to bony fragmentation and dislocation in the midfoot. (Courtesy of Andrew Neckers, MD, and Jean Schils, MD; with permission.)

patients who have received intraarticular glucocorticoid injections, but this is a rare complication and was not observed in one series of patients with knee osteoarthritis who received intraarticular glucocorticoid injections every 3 months for 2 years. The distribution of joint involvement depends on the underlying neurologic disorder (Table 397-2). In tabes dorsalis, the knees, hips, and ankles are most commonly affected; in syringomyelia, the glenohumeral joint, elbow, and wrist; and in diabetes mellitus, the tarsal and tarsometatarsal joints.

PATHOLOGY AND PATHOPHYSIOLOGY

The pathologic changes in the neuropathic joint are similar to those found in the severe osteoarthritic joint. There is fragmentation and eventual loss of articular cartilage with eburnation of the underlying bone. Osteophytes are found at the joint margins. With more advanced disease, erosions are present on the joint surface. Fractures, devitalized bone, intraarticular loose bodies, and microscopic fragments of cartilage and bone may be present.

At least two underlying mechanisms are believed to be involved in the pathogenesis of neuropathic arthritis. An abnormal autonomic nervous system is thought to be responsible for the dysregulated blood flow to the joint with subsequent resorption of bone. Loss of bone, particularly in the diabetic foot, may be the initial finding. With the loss of deep pain, proprioception, and protective neuromuscular reflexes, the joint is subjected to repeated microtrauma, resulting in ligament tears and bone fractures. The injury that follows frequent intraarticular glucocorticoid injections is thought to be due to the analgesic effect of glucocorticoids, leading to overuse of an already damaged joint; the result is accelerated cartilage damage, although steroid-induced cartilage damage is more common in some other animal species than in humans. It is not understood why only a few patients with neuropathy develop clinically evident neuropathic arthritis.

CLINICAL MANIFESTATIONS

Neuropathic joint disease usually begins in a single joint and then becomes apparent in other joints, depending on the underlying neurologic disorder. The involved joint becomes progressively enlarged as a result of bony overgrowth and synovial effusion. Loose bodies may be palpated in the joint cavity. Joint instability, subluxation, and crepitus occur as the disease progresses. Neuropathic joints may develop rapidly, and a totally disorganized joint with multiple bony fragments may evolve within weeks or months. The amount of pain experienced

TABLE 397-2 DISORDERS ASSOCIATED WITH NEUROPATHIC JOINT DISEASE

Diabetes mellitus	Amyloidosis
Tabes dorsalis	Leprosy
Meningomyelocele	Congenital indifference to pain
Syringomyelia	Peroneal muscular atrophy

2244 by the patient is less than would be anticipated from the degree of joint damage. Patients may experience sudden joint pain from intraarticular fractures of osteophytes or condyles.

Neuropathic arthritis is encountered most often in patients with diabetes mellitus, with an incidence of ~0.5%. The onset of disease usually comes at an age of ≥ 50 years in a patient who has had diabetes for several years, but exceptions occur. The tarsal and tarsometatarsal joints are most often affected, with the metatarsophalangeal and talotibial joints next most commonly involved. The knees and spine are occasionally involved. Patients often attribute the onset of foot pain to antecedent trauma such as twisting of the foot. Neuropathic changes may develop rapidly after a foot fracture or dislocation. The foot and ankle are often swollen. Downward collapse of the tarsal bones leads to convexity of the sole, referred to as a “rocker foot.” Large osteophytes may protrude from the top of the foot. Calluses frequently form over the metatarsal heads and may lead to infected ulcers and osteomyelitis. The value of protective inserts and orthotics, as well as regular foot examination, cannot be overstated. Radiographs may show resorption and tapering of the distal metatarsal bones. The term *Lisfranc fracture-dislocation* is sometimes used to describe the destructive changes at the tarsometatarsal joints.

DIAGNOSIS

The diagnosis of neuropathic arthritis is based on the clinical features and characteristic radiographic findings in a patient with underlying sensory neuropathy. The differential diagnosis of neuropathic arthritis depends upon the severity of the process and includes osteomyelitis, avascular necrosis, advanced osteoarthritis, stress fractures, and calcium pyrophosphate deposition disease. Radiographs in neuropathic arthritis initially show changes of osteoarthritis with joint space narrowing, subchondral bone sclerosis, osteophytes, and joint effusions; marked destructive and hypertrophic changes follow later. The radiographic findings of neuropathic arthritis may be difficult to differentiate from those of osteomyelitis, especially in the diabetic foot. The joint margins in a neuropathic joint tend to be distinct, while in osteomyelitis they are blurred. Imaging studies may be helpful, but cultures of tissue from the joint are often required to exclude osteomyelitis. MRI and bone scans using indium 111-labeled white blood cells or indium 111-labeled immunoglobulin G, which will show increased uptake in osteomyelitis but not in a neuropathic joint, may be useful. A technetium bone scan will not distinguish osteomyelitis from neuropathic arthritis, as increased uptake is observed in both. The joint fluid in neuropathic arthritis is noninflammatory; may be xanthochromic or even bloody; and may contain fragments of synovium, cartilage, and bone. The finding of calcium pyrophosphate dihydrate crystals supports the diagnosis of crystal-associated arthropathy. In the absence of such crystals, an increased number of leukocytes may indicate osteomyelitis.

TREATMENT NEUROPATHIC JOINT DISEASE

The primary focus of treatment is to stabilize the joint. Treatment of the underlying disorder, even if successful, does not usually affect established joint disease. Braces and splints are helpful. Their use requires close surveillance, because patients may be unable to appreciate pressure from a poorly adjusted brace. In the diabetic patient, early recognition of Charcot foot and its treatment—prohibition of weight bearing by the foot for at least 8 weeks—may possibly prevent severe disease from developing. Fusion of an unstable joint may improve function and reduce pain, but nonunion is frequent, especially when immobilization of the joint is inadequate.

HYPERTROPHIC OSTEOARTHROPATHY AND CLUBBING

Hypertrophic osteoarthropathy (HOA) is characterized by clubbing of digits and, in more advanced stages, by periosteal new-bone formation and synovial effusions. HOA may be primary or familial and may begin in childhood. Secondary HOA is associated with intrathoracic malignancies, suppurative and some hypoxemic lung diseases,



FIGURE 397-2 Clubbing of the fingers. (Reprinted from the *Clinical Slide Collection on the Rheumatic Diseases*, © 1991, 1995. Used by permission of the American College of Rheumatology.)

congenital heart disease, and a variety of other disorders. Clubbing is almost always a feature of HOA but can occur as an isolated manifestation (Fig. 397-2). The presence of clubbing in isolation may be congenital or represent either an early stage or one element in the spectrum of HOA. Isolated acquired clubbing has the same clinical significance as clubbing associated with periostitis.

Pathology and Pathophysiology of Acquired HOA In HOA, bone changes in the distal extremities begin as periostitis followed by new bone formation. At this stage, a radiolucent area may be observed between the new periosteal bone and the subjacent cortex. As the process progresses, multiple layers of new bone are deposited and become contiguous with the cortex, with consequent cortical thickening. The outer portion of the bone is laminated in appearance, with an irregular surface. Initially, the process of periosteal new-bone formation involves the proximal and distal diaphyses of the tibia, fibula, radius, and ulna and, less frequently, the femur, humerus, metacarpals, metatarsals, and phalanges. Occasionally, scapulae, clavicles, ribs, and pelvic bones are also affected. The adjacent interosseous membranes may become ossified. The distribution of bone manifestations is usually bilateral and symmetric. The soft tissue overlying the distal third of the arms and legs may be thickened. Proliferation of connective tissue occurs in the nail bed and volar pad of digits, giving the distal phalanges a clubbed appearance. Small blood vessels in the clubbed digits are dilated and have thickened walls. In addition, the number of arteriovenous anastomoses is increased.

Several theories have been suggested for the pathogenesis of HOA, but many have been disproved or have not explained the condition's development in all clinical disorders with which it is associated. Previously proposed neurogenic and humoral theories are no longer considered likely explanations for HOA. Studies have suggested a role for platelets in the development of HOA. It has been observed that megakaryocytes and large platelet particles present in the venous circulation are fragmented in their passage through normal lung. In patients with cyanotic congenital heart disease and in other disorders associated with right-to-left shunts, these large platelet particles bypass the lung and reach the distal extremities, where they can interact with endothelial cells. Platelet-endothelial cell activation in the distal portion of the extremities may result in the release of platelet-derived growth factor (PDGF) and other factors leading to the proliferation of connective tissue and periosteum. Stimulation of fibroblasts by PDGF and transforming growth factor β results in cell growth and collagen synthesis. Elevated plasma levels of von Willebrand factor antigen have been found in patients with both primary and secondary forms of HOA, indicating endothelial activation or damage. Abnormalities of collagen synthesis have been demonstrated in the involved skin of patients with primary HOA. Other factors are undoubtedly involved in the pathogenesis of HOA, and further studies are needed to elucidate this disorder.

Clinical Manifestations Primary or familial HOA, also referred to as *pachydermoperiostitis* or *Touraine-Solente-Golé syndrome*, usually

begins insidiously at puberty. In a smaller proportion of patients, the onset comes in the first year of life. The disorder is inherited as an autosomal dominant trait with variable expression and is nine times more common among boys than among girls. Approximately one-third of patients have a family history of primary HOA.

Primary HOA is characterized by clubbing, periostitis, and unusual skin features. A small number of patients with this syndrome do not express clubbing. The skin changes and periostitis are prominent features of this syndrome. The skin becomes thickened and coarse. Deep nasolabial folds develop, and the forehead may become furrowed. Patients may have heavy-appearing eyelids and ptosis. The skin is often greasy, and there may be excessive sweating of the hands and feet. Patients may also experience acne vulgaris, seborrhea, and folliculitis. In a few patients, the skin over the scalp becomes very thick and corrugated, a feature that has been descriptively termed *cutis verticis gyrata*. The distal extremities, particularly the legs, become thickened as a consequence of the proliferation of new bone and soft tissue; when the process is extensive, the distal lower extremities resemble those of an elephant. The periostitis usually is not painful, which it can be in secondary HOA. Clubbing of the fingers may be extensive, producing large, bulbous deformities and clumsiness. Clubbing also affects the toes. Patients may experience articular and periarticular pain, especially in the ankles and knees, and joint motion may be mildly restricted by periarticular bone overgrowth. Noninflammatory effusions occur in the wrists, knees, and ankles. Synovial hypertrophy is not found. Associated abnormalities observed in patients with primary HOA include hypertrophic gastropathy, bone marrow failure, female escutcheon, gynecomastia, and cranial suture defects. In patients with primary HOA, the symptoms disappear when adulthood is reached.

HOA secondary to an underlying disease occurs more frequently than primary HOA. It accompanies a variety of disorders and may precede clinical features of the associated disorder by months. Clubbing is more frequent than the full syndrome of HOA in patients with associated illnesses. Because clubbing evolves over months and is usually asymptomatic, it is often recognized first by the physician and not the patient. Patients may experience a burning sensation in their fingertips. Clubbing is characterized by widening of the fingertips, enlargement of the distal volar pad, convexity of the nail contour, and the loss of the normal 15° angle between the proximal nail and cuticle. The thickness of the digit at the base of the nail is greater than the thickness at the distal interphalangeal joint. An objective measurement of finger clubbing can be made by determining the diameter at the base of the nail and at the distal interphalangeal joint of all 10 digits. Clubbing is present when the sum of the individual digit ratios is >10. At the bedside, clubbing can be appreciated by having the patient place the dorsal surface of the distal phalanges of the fourth fingers together with the nails opposing each other. Normally, an open area is visible between the bases of the opposing fingernails; when clubbing is present, this open space is no longer visible. The base of the nail feels spongy when compressed, and the nail can be easily rocked on its bed. When clubbing is advanced, the finger may have a drumstick appearance, and the distal interphalangeal joint can be hyperextended. Periosteal involvement in the distal extremities may produce a burning or deep-seated aching pain. The pain, which can be quite incapacitating, is aggravated by dependency and relieved by elevation of the affected limbs. Pressure applied over the distal forearms and legs or gentle percussion of distal long bones like the tibia may be quite painful.

Patients may experience joint pain, most often in the ankles, wrists, and knees. Joint effusions may be present; usually, they are small and noninflammatory. The small joints of the hands are rarely affected. Severe joint or long bone pain may be the presenting symptom of an underlying lung malignancy and may precede the appearance of clubbing. In addition, the progression of HOA tends to be more rapid when associated with malignancies, most notably bronchogenic carcinoma. Noninflammatory but variably painful knee effusions may occur prior to the appearance of clubbing and symptoms of distal periostitis. Unlike primary HOA, secondary HOA does not commonly include excessive sweating and oiliness of the skin or thickening of the facial skin.

TABLE 397-3 DISORDERS ASSOCIATED WITH HYPERTROPHIC OSTEOARTHROPATHY

Pulmonary
Bronchogenic carcinoma and other neoplasms
Lung abscesses, empyema, bronchiectasis
Chronic interstitial pneumonitis
Cystic fibrosis
Sarcoidosis
Gastrointestinal
Inflammatory bowel disease
Sprue
Neoplasms: esophagus, liver, bowel
Cardiovascular
Cyanotic congenital heart disease
Subacute bacterial endocarditis
Infected arterial grafts ^a
Aortic aneurysm ^b
Aneurysm of major extremity artery ^a
Patent ductus arteriosus ^b
Arteriovenous fistula of major extremity vessel ^a
Thyroid (thyroid acropachy)
Hyperthyroidism (Graves' disease)

^aUnilateral involvement. ^bBilateral lower-extremity involvement.

HOA occurs in 5–10% of patients with intrathoracic malignancies, the most common being bronchogenic carcinoma and pleural tumors (Table 397-3). Lung metastases infrequently cause HOA. HOA is also seen in patients with intrathoracic infections, including lung abscesses, empyema, and bronchiectasis, but is uncommon in pulmonary tuberculosis. HOA may accompany chronic interstitial pneumonitis, sarcoidosis, and cystic fibrosis. In cystic fibrosis, clubbing is more common than the full syndrome of HOA. Other causes of clubbing include congenital heart disease with right-to-left shunts, bacterial endocarditis, Crohn's disease, ulcerative colitis, sprue, and neoplasms of the esophagus, liver, and small and large bowel. In patients who have congenital heart disease with right-to-left shunts, clubbing alone occurs more often than the full syndrome of HOA.

Unilateral clubbing has been found in association with aneurysms of major extremity arteries, with infected arterial grafts, and with arteriovenous fistulas of brachial vessels. Clubbing of the toes but not the fingers has been associated with an infected abdominal aortic aneurysm and patent ductus arteriosus. Clubbing of a single digit may follow trauma and has been reported in tophaceous gout and sarcoidosis. While clubbing occurs more commonly than the full syndrome in most diseases, periostitis in the absence of clubbing has been observed in the affected limb of patients with infected arterial grafts.

Hyperthyroidism (Graves' disease), treated or untreated, is occasionally associated with clubbing and periostitis of the bones of the hands and feet. This condition is referred to as *thyroid acropachy*. Periostitis may be asymptomatic and occurs in the midshaft and diaphyseal portion of the metacarpal and phalangeal bones. Significant hand-joint pain may occur; this pain may respond to successful therapy for thyroid dysfunction. The long bones of the extremities are seldom affected. Elevated levels of long-acting thyroid stimulator are found in the sera of these patients.

Laboratory Findings The laboratory abnormalities reflect the underlying disorder. The synovial fluid of involved joints has <500 white cells/ μ L, and the cells are predominantly mononuclear. Radiographs show a faint radiolucent line beneath the new periosteal bone along the shaft of long bones at their distal end. These changes are observed most frequently at the ankles, wrists, and knees. The ends of the distal phalanges may show osseous resorption. Radionuclide studies show pericortical linear uptake along the cortical margins of long bones that may precede any radiographic changes.

The treatment of HOA aims to identify the associated disorder and treat it appropriately. The symptoms and signs of HOA may disappear completely with removal of or effective chemotherapy for a tumor or with antibiotic therapy for a chronic pulmonary infection and drainage of the infected site. Vagotomy or percutaneous block of the vagus nerve leads to symptomatic relief in some patients. NSAIDs or analgesics may help control symptoms of HOA.

REFLEX SYMPATHETIC DYSTROPHY SYNDROME

The reflex sympathetic dystrophy syndrome is now referred to as *complex regional pain syndrome, type 1*, according to the new classification system of the International Association for the Study of Pain. This syndrome is characterized by pain and swelling, usually of a distal extremity, accompanied by vasomotor instability, trophic skin changes, and the rapid development of bony demineralization. Reflex sympathetic dystrophy syndrome, including its treatment, is covered in greater detail in [Chap. 454](#).

TIETZE SYNDROME AND COSTOCHONDRITIS

Tietze syndrome is manifested by painful swelling of one or more costochondral articulations. The age of onset is usually before 40, and both sexes are affected equally. In most patients, only one joint is involved, usually the second or third costochondral joint. The onset of anterior chest pain may be sudden or gradual. The pain may radiate to the arms or shoulders and is aggravated by sneezing, coughing, deep inspirations, or twisting motions of the chest. The term *costochondritis* is often used interchangeably with *Tietze syndrome*, but some workers restrict the former term to pain of the costochondral articulations without swelling. Costochondritis is observed in patients over age 40; tends to affect the third, fourth, and fifth costochondral joints; and occurs more often in women. Both syndromes may mimic cardiac or upper abdominal causes of pain. Rheumatoid arthritis, ankylosing spondylitis, and reactive arthritis may involve costochondral joints but are distinguished easily by their other clinical features. Other skeletal causes of anterior chest wall pain are xiphoidalgia and the slipping rib syndrome, which usually involves the tenth rib. Malignancies such as breast cancer, prostate cancer, plasma cell cytoma, and sarcoma can invade the ribs, thoracic spine, or chest wall and produce symptoms suggesting Tietze syndrome. Patients with osteomalacia may have significant rib pain, with or without documented microfractures. These conditions should be distinguishable by radiography, bone scanning, vitamin D measurement, or biopsy. Analgesics, anti-inflammatory drugs, and local glucocorticoid injections usually relieve symptoms of costochondritis/Tietze syndrome. Care should be taken to avoid overdiagnosing these syndromes in patients with acute chest pain syndromes; many patients will be tender to overly vigorous palpation of the costochondral joints.

MYOFASCIAL PAIN SYNDROME

Myofascial pain syndrome is characterized by multiple areas of localized musculoskeletal pain and tenderness in association with tender points. The pain is deep and aching and may be accompanied by a burning sensation. Myofascial pain may be regional and follow trauma, overuse, or prolonged static contraction of a muscle or muscle group, which may occur when an individual is reading or writing at a desk or working at a computer. In addition, this syndrome may be associated with underlying osteoarthritis of the neck or low back. Pain may be referred from tender points to defined areas distant from the area of original tenderness. Palpation of the tender point reproduces or accentuates the pain. The tender points are usually located in the center of a muscle belly, but they can occur at other sites such as costosternal junctions, the xiphoid process, ligamentous and tendinous insertions, fascia, and fatty areas. Tender point sites in muscle have been described as feeling indurated and taut, and palpation may cause the muscle to twitch. These findings, however, have been shown not to be unique to myofascial pain syndrome: in a controlled study, they were also present in some “normal” subjects.

Myofascial pain most often involves the posterior neck, low back, shoulders, and chest. Chronic pain in the muscles of the posterior neck may involve referral of pain from a tender point in the erector neck muscle or upper trapezius to the head, leading to persistent headaches that may last for days. Tender points in the paraspinal muscles of the low back may refer pain to the buttock. Pain may be referred down the leg from a tender point in the gluteus medius and can mimic sciatica. A tender point in the infraspinatus muscle may produce local and referred pain over the lateral deltoid and down the outside of the arm into the hand. Injection of a local anesthetic such as 1% lidocaine into the tender point site often results in at least transient pain relief. Another useful technique is first to spray an agent such as ethyl chloride from the tender point toward the area of referred pain and then to stretch the muscle. This maneuver may need to be repeated several times. Massage and application of ultrasound to the affected area also may be beneficial. Patients should be instructed in methods to prevent muscle stresses related to work and recreation. Posture and resting positions are important in preventing muscle tension. The prognosis in most patients is good. In some patients, regionally localized myofascial pain syndrome may seem to evolve into more generalized fibromyalgia ([Chap. 396](#)). Abnormal or nonrestorative sleep is a common accompaniment in these patients and may need to be specifically addressed.

NEOPLASIAS AND ARTHRITIS

Primary tumors and tumor-like disorders of synovium are uncommon but should be considered in the differential diagnosis of monarticular joint disease. In addition, metastases to bone and primary bone tumors adjacent to a joint may produce joint symptoms.

Pigmented villonodular synovitis (PVNS) is characterized by the slowly progressive, exuberant, benign proliferation of synovial tissue, usually involving a single joint. The most common age of onset is in the third decade, and women are affected slightly more often than men. The cause of this disorder is unknown.

The synovium has a brownish color and numerous large, finger-like villi that fuse to form pedunculated nodules. There is marked hyperplasia of synovial cells in the stroma of the villi. Hemosiderin granules and lipids are found in the cytoplasm of macrophages and in the interstitial tissue. Multinucleated giant cells may be present. The proliferative synovium grows into the subsynovial tissue and invades adjacent cartilage and bone.

The clinical picture of PVNS is characterized by the insidious onset of persistent swelling and pain in affected joints, most commonly the knee. Other joints affected include the hips, ankles, calcaneocuboid joints, elbows, and small joints of the fingers or toes. The disease may also involve the common flexor sheath of the hands or fingers. Less often, tendon sheaths in the wrist, ankle, or foot may be involved. Symptoms of pain, a catching sensation, or stiffness may initially be mild and intermittent and may be present for years before the patient seeks medical attention. Radiographs may show joint space narrowing, erosions, and subchondral cysts. The diagnosis of PVNS is strongly suggested by gradient echo MRI, which reveals a synovial mass lesion of low signal intensity typical of tissue containing hemosiderin ([Fig. 397-3](#)). The joint fluid contains blood and is dark red or almost black in color. Lipid-containing macrophages may be present in the fluid. The joint fluid may be clear if hemorrhage has not occurred. Some patients have polyarticular involvement.

The treatment for PVNS is complete synovectomy. With incomplete synovectomy, the villonodular synovitis recurs, and the rate of tissue growth may be faster than it was originally. Irradiation of the involved joint has been successful in some patients.

Synovial chondromatosis is a disorder characterized by multiple focal metaplastic growths of normal-appearing cartilage in the synovium or tendon sheath. Segments of cartilage break loose and continue to grow as loose bodies. When calcification and ossification of loose bodies occur, the disorder is referred to as *synovial osteochondromatosis*. The disorder is usually monarticular and affects young to middle-aged individuals. The knee is most often involved, followed by hip, elbow, and shoulder. Symptoms are pain, swelling, and decreased

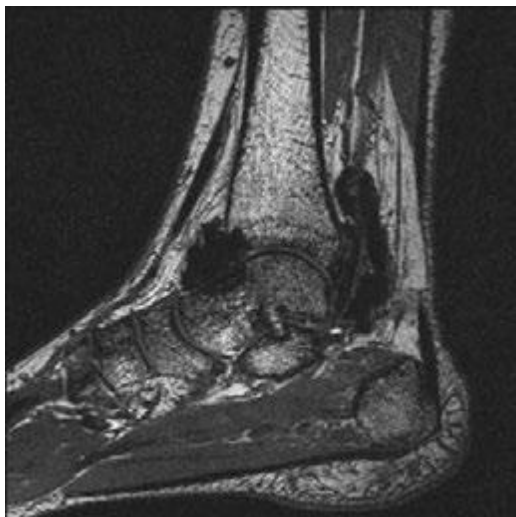


FIGURE 397-3 Pigmented villonodular synovitis. MRI gradient echo sagittal image showing a mass that abuts the neck of the talus with marked low signal typical of tissue containing hemosiderin. (Courtesy of Donald Flemming, MD; with permission.)

motion of the joint. Radiographs may show several rounded calcifications within the joint cavity. Treatment is synovectomy; however, as in PVNS, the tumor may recur.

Synovial sarcoma is a malignant neoplasm often found near a large joint of both upper and lower extremities, being more common in the lower extremity. It seldom arises within the joint itself. Synovial sarcomas constitute 10% of soft tissue sarcomas. The tumor is believed to arise from primitive mesenchymal tissue that differentiates into epithelial cells and/or spindle cells. Small foci of calcification may be present in the tumor mass. Synovial sarcoma occurs most often in young adults and is more common in men. The tumor presents as a slowly growing deep-seated mass near a joint, without much pain. The area of the knee is the most common site, followed by the foot, ankle, elbow, and shoulder. Other primary sites include the buttocks, abdominal wall, retroperitoneum, and mediastinum. The tumor spreads along tissue planes. The most common site of visceral metastasis is the lung. The diagnosis is made by biopsy. Treatment consists of wide resection of the tumor, including adjacent muscle and regional lymph nodes, followed by chemotherapy and radiation therapy. Amputation of the involved distal extremity may be required. Chemotherapy may be beneficial in some patients with metastatic disease. Isolated sites of pulmonary metastasis can be surgically removed. The 5-year survival rate with treatment is variable and depends on the staging of the tumor, ranging from ~25% to ≥60%. Synovial sarcomas tend to recur locally and metastasize to regional lymph nodes, lungs, and skeleton.

In addition to the rare direct metastases of solid cell tumors to the highly vascular synovium, neoplasia arising from nonarticular organ sites can affect joints in other ways. Acute leukemias in children can mimic juvenile inflammatory arthritis with severe joint pain and fever. In adults, chronic and acute myeloid leukemia can infiltrate the synovium in rare instances. The rarely occurring hairy cell leukemia has a peculiar tendency to cause episodic inflammatory oligoarthritis and tenosynovitis; these episodes are dramatic and mimic acute gout attacks. They respond to potent anti-inflammatory therapy with glucocorticoids; with remission of the leukemia, they may abate. Carcinomas can be associated with several paraneoplastic articular syndromes, including HOA (discussed above). Acute palmar fasciitis with polyarthritis is a well-described but rare condition associated with certain cancers, mainly adenocarcinomas. Clinically, this syndrome is fairly abrupt in onset, with pain in the metacarpophalangeal and proximal interphalangeal joints of the hands and rapidly evolving contractures of the fingers due to thickening of the palmar (flexor) tendons. A similar syndrome can be seen in diabetics. Paraneoplastic arthritis has been described and may occur in several patterns: asymmetric

disease predominantly affecting the lower extremity joints and symmetric polyarthritis with hand joint involvement. Tumors are often found after the onset of the arthritis, and many patients have a preceding period of malaise or weight loss. The onset is often acute, and patients tend to be older men. These features should raise the specter of an underlying malignancy (or a viral infection such as hepatitis C) as the cause of the arthritis. In one series, the symptoms resolved with successful therapy for the malignancy and did not recur with relapse of the malignancy. Dermatomyositis has a well-described association with neoplasms and may include joint pain and arthritis. Malignancy-associated arthritis may be responsive to NSAIDs and to treatment of the primary neoplasm.

ACKNOWLEDGMENT

This chapter represents a revised version of the chapter authored by Dr. Bruce C. Gilliland that appeared in previous editions of Harrison's. Dr. Gilliland passed away on February 17, 2007. He had been a contributor to Harrison's Principles of Internal Medicine since the 11th edition.

398 Periarticular Disorders of the Extremities

Carol A. Langford

A number of periarticular disorders have become increasingly common, due in part to greater participation in recreational sports by individuals of a wide range of ages. Periarticular disorders most commonly affect the knee or shoulder. With the exception of bursitis, hip pain is most often articular or is being referred from disease affecting another structure ([Chap 393](#)). This chapter discusses some of the more common periarticular disorders.

BURSITIS

Bursitis is inflammation of a bursa, which is a thin-walled sac lined with synovial tissue. The function of the bursa is to facilitate movement of tendons and muscles over bony prominences. Excessive frictional forces from overuse, trauma, systemic disease (e.g., rheumatoid arthritis, gout), or infection may cause bursitis. *Subacromial bursitis* (subdeltoid bursitis) is the most common form of bursitis. The subacromial bursa, which is contiguous with the subdeltoid bursa, is located between the undersurface of the acromion and the humeral head and is covered by the deltoid muscle. Bursitis is caused by repetitive overhead motion and often accompanies rotator cuff tendinitis. Another frequently encountered form is *trochanteric bursitis*, which involves the bursa around the insertion of the gluteus medius onto the greater trochanter of the femur. Patients experience pain over the lateral aspect of the hip and upper thigh and have tenderness over the posterior aspect of the greater trochanter. External rotation and resisted abduction of the hip elicit pain. *Olecranon bursitis* occurs over the posterior elbow, and when the area is acutely inflamed, infection or gout should be excluded by aspirating the bursa and performing a Gram stain and culture on the fluid as well as examining the fluid for urate crystals. *Achilles bursitis* involves the bursa located above the insertion of the tendon to the calcaneus and results from overuse and wearing tight shoes. *Retrocalcaneal bursitis* involves the bursa that is located between the calcaneus and posterior surface of the Achilles tendon. The pain is experienced at the back of the heel, and swelling appears on the medial and/or lateral side of the tendon. It occurs in association with spondyloarthritis, rheumatoid arthritis, gout, or trauma. *Ischial bursitis* affects the bursa separating the gluteus medius from the ischial tuberosity and develops from prolonged sitting and pivoting on hard surfaces. *Iliopsoas bursitis* affects the bursa that lies between the iliopsoas muscle and hip joint and is lateral to the femoral vessels. Pain is experienced over this area and is made worse

2248 by hip extension and flexion. *Anserine bursitis* is an inflammation of the sartorius bursa located over the medial side of the tibia just below the knee and under the conjoint tendon and is manifested by pain on climbing stairs. Tenderness is present over the insertion of the conjoint tendon of the sartorius, gracilis, and semitendinosus. *Prepatellar bursitis* occurs in the bursa situated between the patella and overlying skin and is caused by kneeling on hard surfaces. Gout or infection may also occur at this site. Bursitis is typically diagnosed by history and physical examination, but visualization by ultrasound may play a useful role in selected instances for diagnosis and directed guidance of glucocorticoid injection. Treatment of bursitis consists of prevention of the aggravating situation, rest of the involved part, administration of a nonsteroidal anti-inflammatory drug (NSAID) where appropriate for an individual patient, or local glucocorticoid injection.

ROTATOR CUFF TENDINITIS AND IMPINGEMENT SYNDROME

Tendinitis of the rotator cuff is the major cause of a painful shoulder and is currently thought to be caused by inflammation of the tendon(s). The rotator cuff consists of the tendons of the supraspinatus, infraspinatus, subscapularis, and teres minor muscles, and inserts on the humeral tuberosities. Of the tendons forming the rotator cuff, the supraspinatus tendon is the most often affected, probably because of its repeated impingement (*impingement syndrome*) between the humeral head and the undersurface of the anterior third of the acromion and coracoacromial ligament above as well as the reduction in its blood supply that occurs with abduction of the arm (Fig. 398-1). The tendon of the infraspinatus and that of the long head of the biceps are less commonly involved. The process begins with edema and hemorrhage of the rotator cuff, which evolves to fibrotic thickening and eventually to rotator cuff degeneration with tendon tears and bone spurs. Subacromial bursitis also accompanies this syndrome. Symptoms usually appear after injury or overuse, especially with activities involving elevation of the arm with some degree of forward flexion. Impingement syndrome occurs in persons participating in baseball, tennis, swimming, or occupations that require repeated elevation of the arm. Those over age 40 are particularly susceptible. Patients complain of a dull aching in the shoulder, which may interfere with sleep. Severe pain is experienced when the arm is actively abducted into an overhead position. The arc between 60° and 120° is especially painful. Tenderness is present over the lateral aspect of the humeral head just below the acromion. NSAIDs, local glucocorticoid injection, and physical therapy may relieve symptoms. Surgical decompression of the subacromial space may be necessary in patients refractory to conservative treatment.

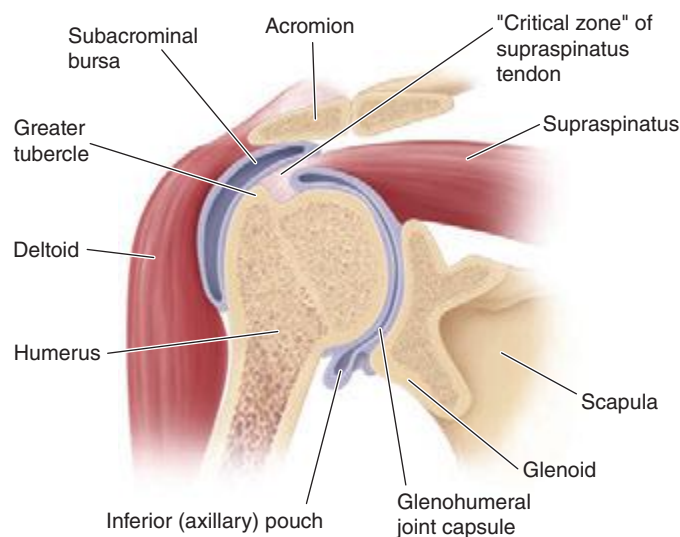


FIGURE 398-1 Coronal section of the shoulder illustrating the relationships of the glenohumeral joint, the joint capsule, the subacromial bursa, and the rotator cuff (supraspinatus tendon). (From F Kozin, in *Arthritis and Allied Conditions*, 13th ed, WJ Koopman [ed]. Baltimore, Williams & Wilkins, 1997, with permission.)

Patients may tear the supraspinatus tendon acutely by falling on an outstretched arm or lifting a heavy object. Symptoms are pain along with weakness of abduction and external rotation of the shoulder. Atrophy of the supraspinatus muscles develops. The diagnosis is established by arthrogram, ultrasound, or magnetic resonance imaging (MRI). Surgical repair may be necessary in patients who fail to respond to conservative measures. In patients with moderate-to-severe tears and functional loss, surgery is indicated.

CALCIFIC TENDINITIS

This condition is characterized by deposition of calcium salts, primarily hydroxyapatite, within a tendon. The exact mechanism of calcification is not known but may be initiated by ischemia or degeneration of the tendon. The supraspinatus tendon is most often affected because it is frequently impinged on and has a reduced blood supply when the arm is abducted. The condition usually develops after age 40. Calcification within the tendon may evoke acute inflammation, producing sudden and severe pain in the shoulder. However, it may be asymptomatic or not related to the patient's symptoms. Diagnosis of calcific tendonitis can be made by ultrasound or radiograph. Most cases are self-limited and respond to conservative therapy with physical therapy and/or NSAIDs. A subset of patients is refractory and requires ultrasound-guided percutaneous needle aspiration and lavage or surgery.

BICIPITAL TENDINITIS AND RUPTURE

Bicipital tendinitis, or tenosynovitis, is produced by friction on the tendon of the long head of the biceps as it passes through the bicipital groove. When the inflammation is acute, patients experience anterior shoulder pain that radiates down the biceps into the forearm. Abduction and external rotation of the arm are painful and limited. The bicipital groove is very tender to palpation. Pain may be elicited along the course of the tendon by resisting supination of the forearm with the elbow at 90° (Yergason's supination sign). Acute rupture of the tendon may occur with vigorous exercise of the arm and is often painful. In a young patient, it should be repaired surgically. Rupture of the tendon in an older person may be associated with little or no pain and is recognized by the presence of persistent swelling of the biceps produced by the retraction of the long head of the biceps. Surgery is usually not necessary in this setting.

DE QUERVAIN'S TENOSYNOVITIS

In this condition, inflammation involves the abductor pollicis longus and the extensor pollicis brevis as these tendons pass through a fibrous sheath at the radial styloid process. The usual cause is repetitive twisting of the wrist. It may occur in pregnancy, and it also occurs in mothers who hold their babies with the thumb outstretched. Patients experience pain on grasping with their thumb, such as with pinching. Swelling and tenderness are often present over the radial styloid process. The Finkelstein sign is positive, which is elicited by having the patient place the thumb in the palm and close the fingers over it. The wrist is then ulnarly deviated, resulting in pain over the involved tendon sheath in the area of the radial styloid. Treatment consists initially of splinting the wrist and an NSAID. When severe or refractory to conservative treatment, glucocorticoid injections can be very effective.

PATELLAR TENDINITIS (JUMPER'S KNEE)

Tendinitis involves the patellar tendon at its attachment to the lower pole of the patella. Patients may experience pain when jumping during basketball or volleyball, going up stairs, or doing deep knee squats. Tenderness is noted on examination over the lower pole of the patella. Treatment consists of rest, icing, and NSAIDs, followed by strengthening and increasing flexibility.

ILIOTIBIAL BAND SYNDROME

The iliotibial band is a thick connective tissue that runs from the ilium to the fibula. Patients with iliotibial band syndrome most commonly present with aching or burning pain at the site where the band courses over the lateral femoral condyle of the knee; pain may also radiate

up the thigh, toward the hip. Predisposing factors for iliotibial band syndrome include a varus alignment of the knee, excessive running distance, poorly fitted shoes, or continuous running on uneven terrain. Treatment consists of rest, NSAIDs, physical therapy, and addressing risk factors such as shoes and running surface. Glucocorticoid injection into the area of tenderness can provide relief, but running must be avoided for at least 2 weeks after the injection. Surgical release of the iliotibial band has been helpful in rare patients for whom conservative treatment has failed.

ADHESIVE CAPSULITIS

Often referred to as “frozen shoulder,” adhesive capsulitis is characterized by pain and restricted movement of the shoulder, usually in the absence of intrinsic shoulder disease. Adhesive capsulitis may follow bursitis or tendinitis of the shoulder or be associated with systemic disorders such as chronic pulmonary disease, myocardial infarction, and diabetes mellitus. Prolonged immobility of the arm contributes to the development of adhesive capsulitis. Pathologically, the capsule of the shoulder is thickened, and a mild chronic inflammatory infiltrate and fibrosis may be present.

Adhesive capsulitis occurs more commonly in women after age 50. Pain and stiffness usually develop gradually but progress rapidly in some patients. Night pain is often present in the affected shoulder, and pain may interfere with sleep. The shoulder is tender to palpation, and both active and passive movement are restricted. Radiographs of the shoulder show osteopenia. The diagnosis is typically made by physical examination but can be confirmed if necessary by arthrography, in that only a limited amount of contrast material, usually <15 mL, can be injected under pressure into the shoulder joint.

In most patients, the condition improves spontaneously 1–3 years after onset. While pain usually improves, many patients are left with some limitation of shoulder motion. Early mobilization of the arm following an injury to the shoulder may prevent the development of this disease. Physical therapy provides the foundation of treatment for adhesive capsulitis. Local injections of glucocorticoids and NSAIDs may also provide relief of symptoms. Slow but forceful injection of contrast material into the joint may lyse adhesions and stretch the capsule, resulting in improvement of shoulder motion. Manipulation under anesthesia may be helpful in some patients.

LATERAL EPICONDYLITIS (TENNIS ELBOW)

Lateral epicondylitis, or tennis elbow, is a painful condition involving the soft tissue over the lateral aspect of the elbow. The pain originates at or near the site of attachment of the common extensors to the lateral epicondyle and may radiate into the forearm and dorsum of the wrist. The pain usually appears after work or recreational activities involving repeated motions of wrist extension and supination against resistance. Most patients with this disorder injure themselves in activities other than tennis, such as pulling weeds, carrying suitcases or briefcases, or using a screwdriver. The injury in tennis usually occurs when hitting a backhand with the elbow flexed. Shaking hands and opening doors can reproduce the pain. Striking the lateral elbow against a solid object may also induce pain.

The treatment is usually rest along with administration of an NSAID. Ultrasound, icing, and friction massage may also help relieve pain. When pain is severe, the elbow is placed in a sling or splinted at 90° of flexion. When the pain is acute and well localized, injection of a glucocorticoid using a small-gauge needle may be effective. Following injection, the patient should be advised to rest the arm for at least 1 month and avoid activities that would aggravate the elbow. Once symptoms have subsided, the patient should begin rehabilitation to strengthen and increase flexibility of the extensor muscles before resuming physical activity involving the arm. A forearm band placed 2.5–5.0 cm (1–2 in.) below the elbow may help to reduce tension on the extensor muscles at their attachment to the lateral epicondyle. The patient should be advised to restrict activities requiring forcible extension and supination of the wrist. Improvement may take several months. The patient may continue to experience mild pain but, with

care, can usually avoid the return of debilitating pain. Occasionally, surgical release of the extensor aponeurosis may be necessary.

MEDIAL EPICONDYLITIS

Medial epicondylitis is an overuse syndrome resulting in pain over the medial side of the elbow with radiation into the forearm. The cause of this syndrome is considered to be repetitive resisted motions of wrist flexion and pronation, which lead to microtears and granulation tissue at the origin of the pronator teres and forearm flexors, particularly the flexor carpi radialis. This overuse syndrome is usually seen in patients >35 years and is much less common than lateral epicondylitis. It occurs most often in work-related repetitive activities but also occurs with recreational activities such as swinging a golf club or throwing a baseball. On physical examination, there is tenderness just distal to the medial epicondyle over the origin of the forearm flexors. Pain can be reproduced by resisting wrist flexion and pronation with the elbow extended. Radiographs are usually normal. The differential diagnosis of patients with medial elbow symptoms includes tears of the pronator teres, acute medial collateral ligament tear, and medial collateral ligament instability. Ulnar neuritis has been found in 25–50% of patients with medial epicondylitis and is associated with tenderness over the ulnar nerve at the elbow as well as hypesthesia and paresthesia on the ulnar side of the hand.

The initial treatment of medial epicondylitis is conservative, involving rest, NSAIDs, friction massage, ultrasound, and icing. Some patients may require splinting. Injections of glucocorticoids at the painful site may also be effective. Patients should be instructed to rest for at least 1 month. Also, patients should start physical therapy once the pain has subsided. In patients with chronic debilitating medial epicondylitis that remains unresponsive after at least a year of treatment, surgical release of the flexor muscle at its origin may be necessary and is often successful.

PLANTAR FASCIITIS

Plantar fasciitis is a common cause of foot pain in adults, with the peak incidence occurring in people between the ages of 40 and 60 years. The pain originates at or near the site of the plantar fascia attachment to the medial tuberosity of the calcaneus. Several factors that increase the risk of developing plantar fasciitis include obesity, pes planus (flat foot or absence of the foot arch when standing), pes cavus (high-arched foot), limited dorsiflexion of the ankle, prolonged standing, walking on hard surfaces, and faulty shoes. In runners, excessive running and a change to a harder running surface may precipitate plantar fasciitis.

The diagnosis of plantar fasciitis can usually be made on the basis of history and physical examination alone. Patients experience severe pain with the first steps on arising in the morning or following inactivity during the day. The pain usually lessens with weight-bearing activity during the day, only to worsen with continued activity. Pain is made worse on walking barefoot or up stairs. On examination, maximal tenderness is elicited on palpation over the inferior heel corresponding to the site of attachment of the plantar fascia.

Imaging studies may be indicated when the diagnosis is not clear. Plain radiographs may show heel spurs, which are of little diagnostic significance. Ultrasonography in plantar fasciitis can demonstrate thickening of the fascia and diffuse hypoechoogenicity, indicating edema at the attachment of the plantar fascia to the calcaneus. MRI is a sensitive method for detecting plantar fasciitis, but it is usually not required for establishing the diagnosis.

The differential diagnosis of inferior heel pain includes calcaneal stress fractures, the spondyloarthritides, rheumatoid arthritis, gout, neoplastic or infiltrative bone processes, and nerve compression/entrapment syndromes.

Resolution of symptoms occurs within 12 months in more than 80% of patients with plantar fasciitis. The patient is advised to reduce or discontinue activities that can exacerbate plantar fasciitis. Initial treatment consists of ice, heat, massage, and stretching. Orthotics provide medial arch support and can be effective. Foot strapping or taping are commonly performed, and some patients may benefit by wearing a

2250 night splint designed to keep the ankle in a neutral position. A short course of NSAIDs can be given to patients when the benefits outweigh the risks. Local glucocorticoid injections have also been shown to be efficacious but may carry an increased risk for plantar fascia rupture. Plantar fasciotomy is reserved for those patients who have failed to improve after at least 6–12 months of conservative treatment.

ACKNOWLEDGMENT

This chapter represents a revised version of the chapter authored by Dr. Bruce C. Gilliland that was in the previous editions of Harrison's. Dr. Gilliland passed away on February 17, 2007, and had been a contributor to Harrison's Principles of Internal Medicine since the 11th edition.

SECTION 1 ENDOCRINOLOGY

399 Approach to the Patient with Endocrine Disorders

J. Larry Jameson

The management of endocrine disorders requires a broad understanding of intermediary metabolism, reproductive physiology, bone metabolism, and growth. Accordingly, the practice of endocrinology is intimately linked to a conceptual framework for understanding hormone secretion, hormone action, and principles of feedback control (Chap. 400e). The endocrine system is evaluated primarily by measuring hormone concentrations, arming the clinician with valuable diagnostic information. Most disorders of the endocrine system are amenable to effective treatment once the correct diagnosis is determined. Endocrine deficiency disorders are treated with physiologic hormone replacement; hormone excess conditions, which usually are caused by benign glandular adenomas, are managed by removing tumors surgically or reducing hormone levels medically.

SCOPE OF ENDOCRINOLOGY

The specialty of endocrinology encompasses the study of glands and the hormones they produce. The term *endocrine* was coined by Starling to contrast the actions of hormones secreted internally (*endocrine*) with those secreted externally (*exocrine*) or into a lumen, such as the gastrointestinal tract. The term *hormone*, derived from a Greek phrase meaning “to set in motion,” aptly describes the dynamic actions of hormones as they elicit cellular responses and regulate physiologic processes through feedback mechanisms.

Unlike many other specialties in medicine, it is not possible to define endocrinology strictly along anatomic lines. The classic endocrine glands—pituitary, thyroid, parathyroid, pancreatic islets, adrenals, and gonads—communicate broadly with other organs through the nervous system, hormones, cytokines, and growth factors. In addition to its traditional synaptic functions, the brain produces a vast array of peptide hormones, and this has led to the discipline of neuroendocrinology. Through the production of hypothalamic releasing factors, the central nervous system (CNS) exerts a major regulatory influence over pituitary hormone secretion (Chap. 401e). The peripheral nervous system stimulates the adrenal medulla. The immune and endocrine systems are also intimately intertwined. The adrenal hormone cortisol is a powerful immunosuppressant. Cytokines and interleukins (ILs) have profound effects on the functions of the pituitary, adrenal, thyroid, and gonads. Common endocrine diseases such as autoimmune thyroid disease and type 1 diabetes mellitus are caused by dysregulation of immune surveillance and tolerance. Less common diseases such as polyglandular failure, Addison’s disease, and lymphocytic hypophysitis also have an immunologic basis.

The interdigitation of endocrinology with physiologic processes in other specialties sometimes blurs the role of hormones. For example, hormones play an important role in maintenance of blood pressure, intravascular volume, and peripheral resistance in the cardiovascular system. Vasoactive substances such as catecholamines, angiotensin II, endothelin, and nitric oxide are involved in dynamic changes of vascular tone in addition to their multiple roles in other tissues. The heart is

the principal source of atrial natriuretic peptide, which acts in classic endocrine fashion to induce natriuresis at a distant target organ (the kidney). Erythropoietin, a traditional circulating hormone, is made in the kidney and stimulates erythropoiesis in bone marrow (Chap. 77). The kidney is also integrally involved in the renin-angiotensin axis (Chap. 406) and is a primary target of several hormones, including parathyroid hormone (PTH), mineralocorticoids, and vasopressin. The gastrointestinal tract produces a surprising number of peptide hormones, such as cholecystokinin, ghrelin, gastrin, secretin, and vasoactive intestinal peptide, among many others. Carcinoid and islet tumors can secrete excessive amounts of these hormones, leading to specific clinical syndromes (Chap. 113). Many of these gastrointestinal hormones are also produced in the CNS, where their functions are poorly understood. Adipose tissue produces leptin, which acts centrally to control appetite, along with adiponectin, resistin, and other hormones that regulate metabolism. As hormones such as inhibin, ghrelin, and leptin are discovered, they become integrated into the science and practice of medicine on the basis of their functional roles rather than their tissues of origin.

Characterization of hormone receptors frequently reveals unexpected relationships to factors in nonendocrine disciplines. The growth hormone (GH) and leptin receptors, for example, are members of the cytokine receptor family. The G protein–coupled receptors (GPCRs), which mediate the actions of many peptide hormones, are used in numerous physiologic processes, including vision, smell, and neurotransmission.

PATHOLOGIC MECHANISMS OF ENDOCRINE DISEASE

Endocrine diseases can be divided into three major types of conditions: (1) hormone excess, (2) hormone deficiency, and (3) hormone resistance (Table 399-1).

CAUSES OF HORMONE EXCESS

Syndromes of hormone excess can be caused by neoplastic growth of endocrine cells, autoimmune disorders, and excess hormone administration. Benign endocrine tumors, including parathyroid, pituitary, and adrenal adenomas, often retain the capacity to produce hormones, perhaps reflecting the fact that these tumors are relatively well differentiated. Many endocrine tumors exhibit subtle defects in their “set points” for feedback regulation. For example, in Cushing’s disease, impaired feedback inhibition of adrenocorticotropic hormone (ACTH) secretion is associated with autonomous function. However, the tumor cells are not completely resistant to feedback, as evidenced by ACTH suppression by higher doses of dexamethasone (e.g., high-dose dexamethasone test) (Chap. 406). Similar set point defects are also typical of parathyroid adenomas and autonomously functioning thyroid nodules.

The molecular basis of some endocrine tumors, such as the multiple endocrine neoplasia (MEN) syndromes (MEN 1, 2A, 2B), have provided important insights into tumorigenesis (Chap. 408). MEN 1 is characterized primarily by the triad of parathyroid, pancreatic islet, and pituitary tumors. MEN 2 predisposes to medullary thyroid carcinoma, pheochromocytoma, and hyperparathyroidism. The *MEN1* gene, located on chromosome 11q13, encodes a putative tumor-suppressor gene, *menin*. Analogous to the paradigm first described for retinoblastoma, the affected individual inherits a mutant copy of the *MEN1* gene, and tumorigenesis ensues after a somatic “second hit”

Type of Endocrine Disorder	Examples
Hyperfunction	
Neoplastic	
Benign	Pituitary adenomas, hyperparathyroidism, autonomous thyroid or adrenal nodules, pheochromocytoma
Malignant	Adrenal cancer, medullary thyroid cancer, carcinoid
Ectopic	Ectopic ACTH, SIADH secretion
Multiple endocrine neoplasia (MEN)	MEN 1, MEN 2
Autoimmune	Graves' disease
Iatrogenic	Cushing's syndrome, hypoglycemia
Infectious/inflammatory	Subacute thyroiditis
Activating receptor mutations	LH, TSH, Ca ²⁺ , PTH receptors, G _s α
Hypofunction	
Autoimmune	Hashimoto's thyroiditis, type 1 diabetes mellitus, Addison's disease, polyglandular failure
Iatrogenic	Radiation-induced hypopituitarism, hypothyroidism, surgical
Infectious/inflammatory	Adrenal insufficiency, hypothalamic sarcoidosis
Hormone mutations	GH, LHβ, FSHβ, vasopressin
Enzyme defects	21-Hydroxylase deficiency
Developmental defects	Kallmann syndrome, Turner's syndrome, transcription factors
Nutritional/vitamin deficiency	Vitamin D deficiency, iodine deficiency
Hemorrhage/infarction	Sheehan's syndrome, adrenal insufficiency
Hormone Resistance	
Receptor mutations	
Membrane	GH, vasopressin, LH, FSH, ACTH, GnRH, GHRH, PTH, leptin, Ca ²⁺
Nuclear	AR, TR, VDR, ER, GR, PPARγ
Signaling pathway mutations	Albright's hereditary osteodystrophy
Postreceptor	Type 2 diabetes mellitus, leptin resistance

Abbreviations: ACTH, adrenocorticotropic hormone; AR, androgen receptor; ER, estrogen receptor; FSH, follicle-stimulating hormone; GHRH, growth hormone-releasing hormone; GnRH, gonadotropin-releasing hormone; GR, glucocorticoid receptor; LH, luteinizing hormone; PPAR, peroxisome proliferator activated receptor; PTH, parathyroid hormone; SIADH, syndrome of inappropriate antidiuretic hormone; TR, thyroid hormone receptor; TSH, thyroid-stimulating hormone; VDR, vitamin D receptor.

leads to loss of function of the normal *MEN1* gene (through deletion or point mutations).

In contrast to inactivation of a tumor-suppressor gene, as occurs in MEN 1 and most other inherited cancer syndromes, MEN 2 is caused by activating mutations in a single allele. In this case, activating mutations of the *RET* protooncogene, which encodes a receptor tyrosine kinase, leads to thyroid C cell hyperplasia in childhood before the development of medullary thyroid carcinoma. Elucidation of this pathogenic mechanism has allowed early genetic screening for *RET* mutations in individuals at risk for MEN 2, permitting identification of those who may benefit from prophylactic thyroidectomy and biochemical screening for pheochromocytoma and hyperparathyroidism.

Mutations that activate hormone receptor signaling have been identified in several GPCRs. For example, activating mutations of the luteinizing hormone (LH) receptor cause a dominantly transmitted form of male-limited precocious puberty, reflecting premature stimulation of testosterone synthesis in Leydig cells (Chap. 411). Activating mutations in these GPCRs are located predominantly in the transmembrane domains and induce receptor coupling to G_sα even in the absence of hormone. Consequently, adenylate cyclase is activated, and cyclic adenosine monophosphate (AMP) levels increase in a manner that mimics hormone action. A similar phenomenon results from activating mutations in G_sα. When these mutations occur early in development, they cause McCune-Albright syndrome. When they occur only in somatotropes, the activating G_sα mutations cause GH-secreting tumors and acromegaly (Chap. 403).

In autoimmune Graves' disease, antibody interactions with the thyroid-stimulating hormone (TSH) receptor mimic TSH action, leading to hormone overproduction (Chap. 405). Analogous to the effects of activating mutations of the TSH receptor, these stimulating autoantibodies induce conformational changes that release the

receptor from a constrained state, thereby triggering receptor coupling to G proteins.

CAUSES OF HORMONE DEFICIENCY

Most examples of hormone deficiency states can be attributed to glandular destruction caused by autoimmunity, surgery, infection, inflammation, infarction, hemorrhage, or tumor infiltration (Table 399-1). Autoimmune damage to the thyroid gland (Hashimoto's thyroiditis) and pancreatic islet β cells (type 1 diabetes mellitus) is a prevalent cause of endocrine disease. Mutations in a number of hormones, hormone receptors, transcription factors, enzymes, and channels can also lead to hormone deficiencies.

HORMONE RESISTANCE

Most severe hormone resistance syndromes are due to inherited defects in membrane receptors, nuclear receptors, or the pathways that transduce receptor signals. These disorders are characterized by defective hormone action despite the presence of increased hormone levels. In complete androgen resistance, for example, mutations in the androgen receptor result in a female phenotypic appearance in genetic (XY) males, even though LH and testosterone levels are increased (Chap. 408). In addition to these relatively rare genetic disorders, more common acquired forms of functional hormone resistance include insulin resistance in type 2 diabetes mellitus, leptin resistance in obesity, and GH resistance in catabolic states. The pathogenesis of functional resistance involves receptor downregulation and postreceptor desensitization of signaling pathways; functional forms of resistance are generally reversible.

CLINICAL EVALUATION OF ENDOCRINE DISORDERS

Because most glands are relatively inaccessible, the physical examination usually focuses on the manifestations of hormone excess or

deficiency as well as direct examination of palpable glands, such as the thyroid and gonads. For these reasons, it is important to evaluate patients in the context of their presenting symptoms, review of systems, family and social history, and exposure to medications that may affect the endocrine system. Astute clinical skills are required to detect subtle symptoms and signs suggestive of underlying endocrine disease. For example, a patient with Cushing's syndrome may manifest specific findings, such as central fat redistribution, striae, and proximal muscle weakness, in addition to features seen commonly in the general population, such as obesity, plethora, hypertension, and glucose intolerance. Similarly, the insidious onset of hypothyroidism—with mental slowing, fatigue, dry skin, and other features—can be difficult to distinguish from similar, nonspecific findings in the general population. Clinical judgment that is based on knowledge of disease prevalence and pathophysiology is required to decide when to embark on more extensive evaluation of these disorders. Laboratory testing plays an essential role in endocrinology by allowing quantitative assessment of hormone levels and dynamics. Radiologic imaging tests such as computed tomography (CT) scan, magnetic resonance imaging (MRI), thyroid scan, and ultrasound are also used for the diagnosis of endocrine disorders. However, these tests generally are employed only after a hormonal abnormality has been established by biochemical testing.

HORMONE MEASUREMENTS AND ENDOCRINE TESTING

Immunoassays are the most important diagnostic tool in endocrinology, as they allow sensitive, specific, and quantitative determination of steady-state and dynamic changes in hormone concentrations. Immunoassays use antibodies to detect specific hormones. For many peptide hormones, these measurements are now configured to use two different antibodies to increase binding affinity and specificity. There are many variations of these assays; a common format involves using one antibody to capture the antigen (hormone) onto an immobilized surface and a second antibody, coupled to a chemiluminescent (immunochemiluminescent assay [ICMA]) or radioactive (immunoradiometric assay [IRMA]) signal, to detect the antigen. These assays are sensitive enough to detect plasma hormone concentrations in the picomolar to nanomolar range, and they can readily distinguish structurally related proteins, such as PTH from PTH-related peptide (PTHrP). A variety of other techniques are used to measure specific hormones, including mass spectroscopy, various forms of chromatography, and enzymatic methods; bioassays are now rarely used.

Most hormone measurements are based on plasma or serum samples. However, urinary hormone determinations remain useful for the evaluation of some conditions. Urinary collections over 24 h provide an integrated assessment of the production of a hormone or metabolite, many of which vary during the day. It is important to assure complete collections of 24-h urine samples; simultaneous measurement of creatinine provides an internal control for the adequacy of collection and can be used to normalize some hormone measurements. A 24-h urine free cortisol measurement largely reflects the amount of unbound cortisol, thus providing a reasonable index of biologically available hormone. Other commonly used urine determinations include 17-hydroxycorticosteroids, 17-ketosteroids, vanillylmandelic acid, metanephrine, catecholamines, 5-hydroxyindoleacetic acid, and calcium.

The value of quantitative hormone measurements lies in their correct interpretation in a clinical context. The normal range for most hormones is relatively broad, often varying by a factor of two- to tenfold. The normal ranges for many hormones are sex- and age-specific. Thus, using the correct normative database is an essential part of interpreting hormone tests. The pulsatile nature of hormones and factors that can affect their secretion, such as sleep, meals, and medications, must also be considered. Cortisol values increase fivefold between midnight and dawn; reproductive hormone levels vary dramatically during the female menstrual cycle.

For many endocrine systems, much information can be gained from basal hormone testing, particularly when different components of an endocrine axis are assessed simultaneously. For example, low testosterone and elevated LH levels suggest a primary gonadal problem, whereas a hypothalamic-pituitary disorder is likely if both LH and testosterone are low. Because TSH is a sensitive indicator of thyroid function, it is generally recommended as a first-line test for thyroid disorders. An elevated TSH level is almost always the result of primary hypothyroidism, whereas a low TSH is most often caused by thyrotoxicosis. These predictions can be confirmed by determining the free thyroxine level. In the less common circumstance when free thyroxine and TSH are both low, it is important to consider secondary hypopituitarism caused by hypothalamic-pituitary disease. Elevated calcium and PTH levels suggest hyperparathyroidism, whereas PTH is suppressed in hypercalcemia caused by malignancy or granulomatous diseases. A suppressed ACTH in the setting of hypercortisolemia, or increased urine free cortisol, is seen with hyperfunctioning adrenal adenomas.

It is not uncommon, however, for baseline hormone levels associated with pathologic endocrine conditions to overlap with the normal range. In this circumstance, dynamic testing is useful to separate the two groups further. There are a multitude of dynamic endocrine tests, but all are based on principles of feedback regulation, and most responses can be rationalized based on principles that govern the regulation of endocrine axes. *Suppression tests* are used in the setting of suspected endocrine hyperfunction. An example is the dexamethasone suppression test used to evaluate Cushing's syndrome (Chaps. 403 and 406). *Stimulation tests* generally are used to assess endocrine hypofunction. The ACTH stimulation test, for example, is used to assess the adrenal gland response in patients with suspected adrenal insufficiency. Other stimulation tests use hypothalamic-releasing factors such as corticotropin-releasing hormone (CRH) and growth hormone-releasing hormone (GHRH) to evaluate pituitary hormone reserve (Chap. 403). Insulin-induced hypoglycemia also evokes pituitary ACTH and GH responses. Stimulation tests based on reduction or inhibition of endogenous hormones are now used infrequently. Examples include metyrapone inhibition of cortisol synthesis and clomiphene inhibition of estrogen feedback.

SCREENING AND ASSESSMENT OF COMMON ENDOCRINE DISORDERS

Many endocrine disorders are prevalent in the adult population (Table 399-2) and can be diagnosed and managed by general internists, family practitioners, or other primary health care providers. The high prevalence and clinical impact of certain endocrine diseases justifies vigilance for features of these disorders during routine physical examinations; laboratory screening is indicated in selected high-risk populations.

Disorder	Approx. Prevalence in Adults ^a	Screening/Testing Recommendations ^b	Chapter(s)
Obesity	34% BMI \geq 30 68% BMI \geq 25	Calculate BMI Measure waist circumference Exclude secondary causes Consider comorbid complications	416
Type 2 diabetes mellitus	>7%	Beginning at age 45, screen every 3 years, or earlier in high-risk groups: Fasting plasma glucose (FPG) >126 mg/dL Random plasma glucose >200 mg/dL An elevated HbA1c Consider comorbid complications	417
Hyperlipidemia	20–25%	Cholesterol screening at least every 5 years; more often in high-risk groups Lipoprotein analysis (LDL, HDL) for increased cholesterol, CAD, diabetes Consider secondary causes	421
Metabolic syndrome	35%	Measure waist circumference, FPG, BP, lipids	422
Hypothyroidism	5–10%, women 0.5–2%, men	TSH; confirm with free T ₄ Screen women after age 35 and every 5 years thereafter	405
Graves' disease	1–3%, women 0.1%, men	TSH, free T ₄	405
Thyroid nodules and neoplasia	2–5% palpable >25% by ultrasound	Physical examination of thyroid Fine-needle aspiration biopsy	405
Osteoporosis	5–10%, women 2–5%, men	Bone mineral density measurements in women >65 years or in postmenopausal women or men at risk Exclude secondary causes	425
Hyperparathyroidism	0.1–0.5%, women > men	Serum calcium PTH, if calcium is elevated Assess comorbid conditions	424
Infertility	10%, couples	Investigate both members of couple Semen analysis in male Assess ovulatory cycles in female Specific tests as indicated	411, 412
Polycystic ovarian syndrome	5–10%, women	Free testosterone, DHEAS Consider comorbid conditions	412
Hirsutism	5–10%	Free testosterone, DHEAS Exclude secondary causes Additional tests as indicated	68
Menopause	Median age, 51	FSH	413
Hyperprolactinemia	15% in women with amenorrhea or galactorrhea	PRL level MRI, if not medication-related	403
Erectile dysfunction	10–25%	Careful history, PRL, testosterone Consider secondary causes (e.g., diabetes)	67
Hypogonadism, male	1–2%	Testosterone, LH	411
Gynecomastia	15%	Often, no tests are indicated Consider Klinefelter's syndrome Consider medications, hypogonadism, liver disease	411
Klinefelter's syndrome	0.2%, men	Karyotype Testosterone	410
Vitamin D deficiency	10%	Measure serum 25-OH vitamin D Consider secondary causes	423
Turner's syndrome	0.03%, women	Karyotype Consider comorbid conditions	410

^aThe prevalence of most disorders varies among ethnic groups and with aging. Data based primarily on U.S. population. ^bSee individual chapters for additional information on evaluation and treatment. Early testing is indicated in patients with signs and symptoms of disease and in those at increased risk.

Abbreviations: BMI, body mass index; BP, blood pressure; CAD, coronary artery disease; DHEAS, dehydroepiandrosterone; FSH, follicle-stimulating hormone; HDL, high-density lipoprotein; LDL, low-density lipoprotein; LH, luteinizing hormone; MRI, magnetic resonance imaging; PRL, prolactin; PTH, parathyroid hormone; TSH, thyroid-stimulating hormone.

400e Mechanisms of Hormone Action

J. Larry Jameson

CLASSES OF HORMONES

Hormones can be divided into five major types: (1) *amino acid derivatives* such as dopamine, catecholamine, and thyroid hormone; (2) *small neuropeptides* such as gonadotropin-releasing hormone (GnRH), thyrotropin-releasing hormone (TRH), somatostatin, and vasopressin; (3) *large proteins* such as insulin, luteinizing hormone (LH), and parathyroid hormone (PTH); (4) *steroid hormones* such as cortisol and estrogen that are synthesized from cholesterol-based precursors; and (5) *vitamin derivatives* such as retinoids (vitamin A) and vitamin D. A variety of *peptide growth factors*, most of which act locally, share actions with hormones. As a rule, amino acid derivatives and peptide hormones interact with cell-surface membrane receptors. Steroids, thyroid hormones, vitamin D, and retinoids are lipid-soluble and interact with intracellular nuclear receptors, although many also interact with membrane receptors or intracellular signaling proteins as well.

HORMONE AND RECEPTOR FAMILIES

Hormones and receptors can be grouped into families, reflecting structural similarities and evolutionary origins (Table 400e-1). The evolution of these families generates diverse but highly selective pathways of hormone action. Recognition of these relationships has proven useful for extrapolating information gleaned from one hormone or receptor to other family members.

The glycoprotein hormone family, consisting of thyroid-stimulating hormone (TSH), follicle-stimulating hormone (FSH), LH, and human chorionic gonadotropin (hCG), illustrates many features of related hormones. The glycoprotein hormones are heterodimers that share the α subunit in common; the β subunits are distinct and confer

specific biologic actions. The overall three-dimensional architecture of the β subunits is similar, reflecting the locations of conserved disulfide bonds that restrain protein conformation. The cloning of the β -subunit genes from multiple species suggests that this family arose from a common ancestral gene, probably by gene duplication and subsequent divergence to evolve new biologic functions.

As hormone families enlarge and diverge, their receptors must co-evolve to derive new biologic functions. Related G protein-coupled receptors (GPCRs), for example, have evolved for each of the glycoprotein hormones. These receptors are structurally similar, and each is coupled predominantly to the $G_s\alpha$ signaling pathway. However, there is minimal overlap of hormone binding. For example, TSH binds with high specificity to the TSH receptor but interacts minimally with the LH or FSH receptors. Nonetheless, there can be subtle physiologic consequences of hormone cross-reactivity with other receptors. Very high levels of hCG during pregnancy stimulate the TSH receptor and increase thyroid hormone levels, resulting in a compensatory decrease in TSH.

Insulin and insulin-like growth factor I (IGF-I) and IGF-II have structural similarities that are most apparent when precursor forms of the proteins are compared. In contrast to the high degree of specificity seen with the glycoprotein hormones, there is moderate cross-talk among the members of the insulin/IGF family. High concentrations of an IGF-II precursor produced by certain tumors (e.g., sarcomas) can cause hypoglycemia, partly because of binding to insulin and IGF-I receptors (Chap. 424). High concentrations of insulin also bind to the IGF-I receptor, perhaps accounting for some of the clinical manifestations seen in conditions with chronic hyperinsulinemia.

Another important example of receptor cross-talk is seen with PTH and parathyroid hormone-related peptide (PTHrP) (Chap. 424). PTH is produced by the parathyroid glands, whereas PTHrP is expressed at high levels during development and by a variety of tumors (Chap. 121). These hormones have amino acid sequence similarity, particularly in their amino-terminal regions. Both hormones bind to a single PTH receptor that is expressed in bone and kidney. Hypercalcemia and hypophosphatemia therefore may result from excessive production of either hormone, making it difficult to distinguish hyperparathyroidism from hypercalcemia of malignancy solely on the basis of serum chemistries. However, sensitive and specific assays for PTH and PTHrP now allow these disorders to be distinguished more readily.

Based on their specificities for DNA binding sites, the nuclear receptor family can be subdivided into type 1 receptors (glucocorticoid receptor, mineralocorticoid receptor, androgen receptor, estrogen receptor, progesterone receptor) that bind steroids and type 2 receptors (thyroid hormone receptor, vitamin D receptor, retinoic acid receptor, peroxisome proliferator activated receptor) that bind thyroid hormone, vitamin D, retinoic acid, or lipid derivatives. Certain functional domains in nuclear receptors, such as the zinc finger DNA-binding domains, are highly conserved. However, selective amino acid differences within this domain confer DNA sequence specificity. The hormone-binding domains are more variable, providing great diversity in the array of small molecules that bind to different nuclear receptors. With few exceptions, hormone binding is highly specific for a single type of nuclear receptor. One exception involves the glucocorticoid and mineralocorticoid receptors. Because the mineralocorticoid receptor also binds glucocorticoids with high affinity, an enzyme (11 β -hydroxysteroid dehydrogenase) in renal tubular cells inactivates glucocorticoids, allowing selective responses to mineralocorticoids such as aldosterone. However, when very high glucocorticoid concentrations occur, as in Cushing's syndrome, the glucocorticoid degradation pathway becomes saturated, allowing excessive cortisol levels to exert mineralocorticoid effects (sodium retention, potassium wasting). This phenomenon is particularly pronounced in ectopic adrenocorticotrophic hormone (ACTH) syndromes (Chap. 406). Another example of relaxed nuclear receptor specificity involves the estrogen receptor, which can bind an array of compounds, some of which have little apparent structural similarity to the high-affinity ligand estradiol. This feature of the estrogen receptor makes it susceptible to activation by "environmental estrogens" such as resveratrol, octylphenol, and many other aromatic hydrocarbons. However, this lack of specificity

TABLE 400e-1 EXAMPLES OF MEMBRANE RECEPTOR FAMILIES AND SIGNALING PATHWAYS		
Receptors	Effectors	Signaling Pathways
G Protein-Coupled Seven-Transmembrane Receptor (GPCR)		
β -Adrenergic, LH, FSH, TSH	$G_s\alpha$, adenylate cyclase	Stimulation of cyclic AMP production, protein kinase A
Glucagon, PTH, PTHrP, ACTH, MSH, GHRH, CRH	Ca^{2+} channels	Calmodulin, Ca^{2+} -dependent kinases
α -Adrenergic, somatostatin	$G_i\alpha$	Inhibition of cyclic AMP production
TRH, GnRH	G_q, G_{11}	Activation of K^+ , Ca^{2+} channels
		Phospholipase C, diacylglycerol, IP_3 , protein kinase C, voltage-dependent Ca^{2+} channels
Receptor Tyrosine Kinase		
Insulin, IGF-I	Tyrosine kinases, IRS	MAP kinases, PI 3-kinase; AKT
EGF, NGF	Tyrosine kinases, ras	Raf, MAP kinases, RSK
Cytokine Receptor-Linked Kinase		
GH, PRL	JAK, tyrosine kinases	STAT, MAP kinase, PI 3-kinase, IRS-1
Serine Kinase		
Activin, TGF- β , MIS	Serine kinase	Smads

Abbreviations: IP_3 , inositol triphosphate; IRS, insulin receptor substrates; MAP, mitogen-activated protein; MSH, melanocyte-stimulating hormone; NGF, nerve growth factor; PI, phosphatidylinositol; RSK, ribosomal S6 kinase; TGF- β , transforming growth factor β . For all other abbreviations, see text. Note that most receptors interact with multiple effectors and activate networks of signaling pathways.

400e-2 provides an opportunity to synthesize a remarkable series of clinically useful antagonists (e.g., tamoxifen) and selective estrogen response modulators (SERMs) such as raloxifene. These compounds generate distinct conformations that alter receptor interactions with components of the transcription machinery (see below), thereby conferring their unique actions.

HORMONE SYNTHESIS AND PROCESSING

The synthesis of peptide hormones and their receptors occurs through a classic pathway of gene expression: transcription → mRNA → protein → posttranslational protein processing → intracellular sorting, followed by membrane integration or secretion (**Chap. 82**).

Many hormones are embedded within larger precursor polypeptides that are proteolytically processed to yield the biologically active hormone. Examples include proopiomelanocortin (POMC) → ACTH; proglucagon → glucagon; proinsulin → insulin; and pro-PTH → PTH, among others. In many cases, such as POMC and proglucagon, these precursors generate multiple biologically active peptides. It is provocative that hormone precursors are typically inactive, presumably adding an additional level of regulatory control. Prohormone conversion occurs not only for peptide hormones but also for certain steroids (testosterone → dihydrotestosterone) and thyroid hormone (T_4 → T_3).

Peptide precursor processing is intimately linked to intracellular sorting pathways that transport proteins to appropriate vesicles and enzymes, resulting in specific cleavage steps, followed by protein folding and translocation to secretory vesicles. Hormones destined for secretion are translocated across the endoplasmic reticulum under the guidance of an amino-terminal signal sequence that subsequently is cleaved. Cell-surface receptors are inserted into the membrane via short segments of hydrophobic amino acids that remain embedded within the lipid bilayer. During translocation through the Golgi and endoplasmic reticulum, hormones and receptors are subject to a variety of posttranslational modifications, such as glycosylation and phosphorylation, which can alter protein conformation, modify circulating half-life, and alter biologic activity.

Synthesis of most steroid hormones is based on modifications of the precursor, cholesterol. Multiple regulated enzymatic steps are required for the synthesis of testosterone (**Chap. 411**), estradiol (**Chap. 412**), cortisol (**Chap. 406**), and vitamin D (**Chap. 423**). This large number of synthetic steps predisposes to multiple genetic and acquired disorders of steroidogenesis.

Endocrine genes contain regulatory DNA elements similar to those found in many other genes, but their exquisite control by hormones reflects the presence of specific hormone response elements. For example, the TSH genes are repressed directly by thyroid hormones acting through the thyroid hormone receptor (TR), a member of the nuclear receptor family. Steroidogenic enzyme gene expression requires specific transcription factors, such as steroidogenic factor-1 (SF-1), acting in conjunction with signals transmitted by trophic hormones (e.g., ACTH or LH). For some hormones, substantial regulation occurs at the level of translational efficiency. Insulin biosynthesis, although it requires ongoing gene transcription, is regulated primarily at the translational and secretory levels in response to elevated levels of glucose or amino acids.

HORMONE SECRETION, TRANSPORT, AND DEGRADATION

The level of a hormone is determined by its rate of secretion and its circulating half-life. After protein processing, peptide hormones (e.g., GnRH, insulin, growth hormone [GH]) are stored in secretory granules. As these granules mature, they are poised beneath the plasma membrane for imminent release into the circulation. In most instances, the stimulus for hormone secretion is a releasing factor or neural signal that induces rapid changes in intracellular calcium concentrations, leading to secretory granule fusion with the plasma membrane and release of its contents into the extracellular environment and bloodstream. Steroid hormones, in contrast, diffuse into the circulation as they are synthesized. Thus, their secretory rates are closely aligned with rates of synthesis. For example, ACTH and LH induce steroidogenesis

by stimulating the activity of the steroidogenic acute regulatory (StAR) protein (transports cholesterol into the mitochondrion) along with other rate-limiting steps (e.g., cholesterol side-chain cleavage enzyme, CYP11A1) in the steroidogenic pathway.

Hormone transport and degradation dictate the rapidity with which a hormonal signal decays. Some hormone signals are evanescent (e.g., somatostatin), whereas others are longer-lived (e.g., TSH). Because somatostatin exerts effects in virtually every tissue, a short half-life allows its concentrations and actions to be controlled locally. Structural modifications that impair somatostatin degradation have been useful for generating long-acting therapeutic analogues such as octreotide (**Chap. 403**). In contrast, the actions of TSH are highly specific for the thyroid gland. Its prolonged half-life accounts for relatively constant serum levels even though TSH is secreted in discrete pulses.

An understanding of circulating hormone half-life is important for achieving physiologic hormone replacement, as the frequency of dosing and the time required to reach steady state are intimately linked to rates of hormone decay. T_4 , for example, has a circulating half-life of 7 days. Consequently, >1 month is required to reach a new steady state, and single daily doses are sufficient to achieve constant hormone levels. T_3 , in contrast, has a half-life of 1 day. Its administration is associated with more dynamic serum levels, and it must be administered two to three times per day. Similarly, synthetic glucocorticoids vary widely in their half-lives; those with longer half-lives (e.g., dexamethasone) are associated with greater suppression of the hypothalamic-pituitary-adrenal (HPA) axis. Most protein hormones (e.g., ACTH, GH, prolactin [PRL], PTH, LH) have relatively short half-lives (<20 min), leading to sharp peaks of secretion and decay. The only accurate way to profile the pulse frequency and amplitude of these hormones is to measure levels in frequently sampled blood (every 10 min or less) over long durations (8–24 h). Because this is not practical in a clinical setting, an alternative strategy is to pool three to four samples drawn at about 30-min intervals, or interpret the results in the context of a relatively wide normal range. Rapid hormone decay is useful in certain clinical settings. For example, the short half-life of PTH allows the use of intraoperative PTH determinations to confirm successful removal of an adenoma. This is particularly valuable diagnostically when there is a possibility of multicentric disease or parathyroid hyperplasia, as occurs with multiple endocrine neoplasia (MEN) or renal insufficiency.

Many hormones circulate in association with serum-binding proteins. Examples include (1) T_4 and T_3 binding to thyroxine-binding globulin (TBG), albumin, and thyroxine-binding prealbumin (TBPA); (2) cortisol binding to cortisol-binding globulin (CBG); (3) androgen and estrogen binding to sex hormone-binding globulin (SHBG); (4) IGF-I and -II binding to multiple IGF-binding proteins (IGFBPs); (5) GH interactions with GH-binding protein (GHBP), a circulating fragment of the GH receptor extracellular domain; and (6) activin binding to follistatin. These interactions provide a hormonal reservoir, prevent otherwise rapid degradation of unbound hormones, restrict hormone access to certain sites (e.g., IGFBPs), and modulate the unbound, or “free,” hormone concentrations. Although a variety of binding protein abnormalities have been identified, most have little clinical consequence aside from creating diagnostic problems. For example, TBG deficiency can reduce total thyroid hormone levels greatly but the free concentrations of T_4 and T_3 remain normal. Liver disease and certain medications can also influence binding protein levels (e.g., estrogen increases TBG) or cause displacement of hormones from binding proteins (e.g., salsalate displaces T_4 from TBG). In general, only unbound hormone is available to interact with receptors and thus elicit a biologic response. Short-term perturbations in binding proteins change the free hormone concentration, which in turn induces compensatory adaptations through feedback loops. SHBG changes in women are an exception to this self-correcting mechanism. When SHBG decreases because of insulin resistance or androgen excess, the unbound testosterone concentration is increased, potentially leading to hirsutism (**Chap. 68**). The increased unbound testosterone level does not result in an adequate compensatory feedback correction because estrogen, not testosterone, is the primary regulator of the reproductive axis.

An additional exception to the unbound hormone hypothesis involves megalin, a member of the low-density lipoprotein (LDL) receptor family that serves as an endocytotic receptor for carrier-bound vitamins A and D and SHBG-bound androgens and estrogens. After internalization, the carrier proteins are degraded in lysosomes and release their bound ligands within the cells. Membrane transporters have also been identified for thyroid hormones.

Hormone degradation can be an important mechanism for regulating concentrations locally. As noted above, 11β -hydroxysteroid dehydrogenase inactivates glucocorticoids in renal tubular cells, preventing actions through the mineralocorticoid receptor. Thyroid hormone deiodinases convert T_4 to T_3 and can inactivate T_3 . During development, degradation of retinoic acid by Cyp26b1 prevents primordial germ cells in the male from entering meiosis, as occurs in the female ovary.

HORMONE ACTION THROUGH RECEPTORS

Receptors for hormones are divided into two major classes: membrane and nuclear. *Membrane receptors* primarily bind peptide hormones and catecholamines. *Nuclear receptors* bind small molecules that can diffuse across the cell membrane, such as steroids and vitamin D. Certain general principles apply to hormone-receptor interactions regardless of the class of receptor. Hormones bind to receptors with specificity and an affinity that generally coincides with the dynamic range of circulating hormone concentrations. Low concentrations of free hormone (usually 10^{-12} to 10^{-9} M) rapidly associate and dissociate from receptors in a bimolecular reaction such that the occupancy of the receptor at any given moment is a function of hormone concentration and the receptor's affinity for the hormone. Receptor numbers vary greatly in different target tissues, providing one of the major determinants of specific tissue responses to circulating hormones. For example, ACTH receptors are located almost exclusively in the adrenal cortex, and FSH receptors are found predominantly in the gonads. In contrast, insulin and TRs are widely distributed, reflecting the need for metabolic responses in all tissues.

MEMBRANE RECEPTORS

Membrane receptors for hormones can be divided into several major groups: (1) seven transmembrane GPCRs, (2) tyrosine kinase receptors, (3) cytokine receptors, and (4) serine kinase receptors (Fig. 400e-1). The *seven transmembrane GPCR family* binds a remarkable array of hormones, including large proteins (e.g., LH, PTH), small peptides (e.g., TRH, somatostatin), catecholamines (epinephrine, dopamine), and even minerals (e.g., calcium). The extracellular domains of GPCRs

vary widely in size and are the major binding site for large hormones. The transmembrane-spanning regions are composed of hydrophobic α -helical domains that traverse the lipid bilayer. Like some channels, these domains are thought to circularize and form a hydrophobic pocket into which certain small ligands fit. Hormone binding induces conformational changes in these domains, transducing structural changes to the intracellular domain, which is a docking site for G proteins.

The large family of *G proteins*, so named because they bind guanine nucleotides (guanosine triphosphate [GTP], guanosine diphosphate [GDP]), provides great diversity for coupling receptors to different signaling pathways. G proteins form a heterotrimeric complex that is composed of various α and $\beta\gamma$ subunits. The α subunit contains the guanine nucleotide-binding site and hydrolyzes $GTP \rightarrow GDP$. The $\beta\gamma$ subunits are tightly associated and modulate the activity of the α subunit as well as mediating their own effector signaling pathways. G protein activity is regulated by a cycle that involves GTP hydrolysis and dynamic interactions between the α and $\beta\gamma$ subunits. Hormone binding to the receptor induces GDP dissociation, allowing $G\alpha$ to bind GTP and dissociate from the $\alpha\beta$ complex. Under these conditions, the $G\alpha$ subunit is activated and mediates signal transduction through various enzymes, such as adenylate cyclase and phospholipase C. GTP hydrolysis to GDP allows reassociation with the $\beta\gamma$ subunits and restores the inactive state. As described below, a variety of endocrinopathies result from G protein mutations or from mutations in receptors that modify their interactions with G proteins. G proteins interact with other cellular proteins, including kinases, channels, G protein-coupled receptor kinases (GRKs), and arrestins, that mediate signaling as well as receptor desensitization and recycling.

The *tyrosine kinase receptors* transduce signals for insulin and a variety of growth factors, such as IGF-I, epidermal growth factor (EGF), nerve growth factor, platelet-derived growth factor, and fibroblast growth factor. The cysteine-rich extracellular ligand-binding domains contain growth factor binding sites. After ligand binding, this class of receptors undergoes autophosphorylation, inducing interactions with intracellular adaptor proteins such as Shc and insulin receptor substrates (IRS). In the case of the insulin receptor, multiple kinases are activated, including the Raf-Ras-MAPK and the Akt/protein kinase B pathways. The tyrosine kinase receptors play a prominent role in cell growth and differentiation as well as in intermediary metabolism.

The GH and PRL receptors belong to the *cytokine receptor* family. Analogous to the tyrosine kinase receptors, ligand binding induces

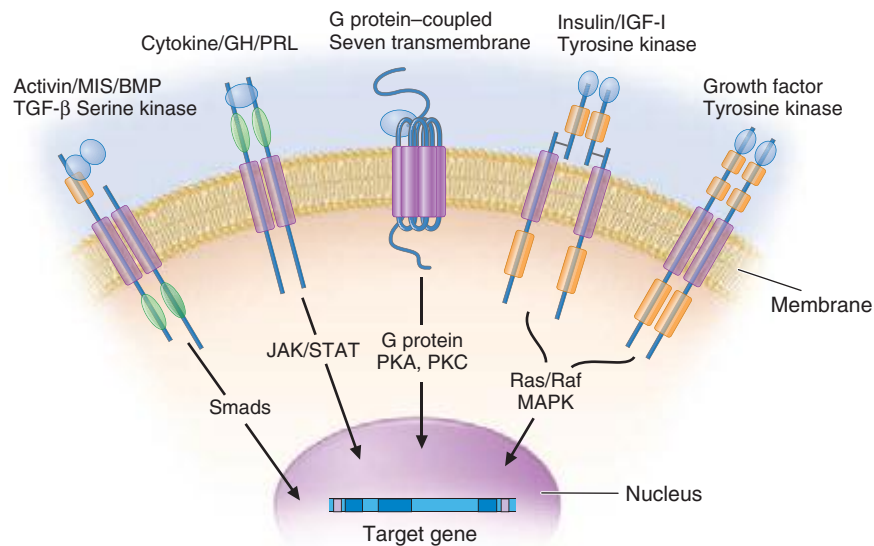


FIGURE 400e-1 Membrane receptor signaling. MAPK, mitogen-activated protein kinase; PKA, C, protein kinase A, C; TGF, transforming growth factor. For other abbreviations, see text.

400e-4 receptor interaction with intracellular kinases—the Janus kinases (JAKs), which phosphorylate members of the signal transduction and activators of transcription (STAT) family—as well as with other signaling pathways (Ras, PI3-K, MAPK). The activated STAT proteins translocate to the nucleus and stimulate expression of target genes.

The *serine kinase receptors* mediate the actions of activins, transforming growth factor β , müllerian-inhibiting substance (MIS, also known as anti-müllerian hormone, AMH), and bone morphogenic proteins (BMPs). This family of receptors (consisting of type I and II subunits) signals through proteins termed *smads* (fusion of terms for *Caenorhabditis elegans sma* + mammalian *mad*). Like the STAT proteins, the smads serve a dual role of transducing the receptor signal and acting as transcription factors. The pleomorphic actions of these growth factors dictate that they act primarily in a local (paracrine or autocrine) manner. Binding proteins such as follistatin (which binds activin and other members of this family) function to inactivate the growth factors and restrict their distribution.

NUCLEAR RECEPTORS

The family of nuclear receptors has grown to nearly 100 members, many of which are still classified as orphan receptors because their ligands, if they exist, have not been identified (Fig. 400e-2). Otherwise, most nuclear receptors are classified on the basis of their ligands. Although all nuclear receptors ultimately act to increase or decrease gene transcription, some (e.g., glucocorticoid receptor) reside primarily in the cytoplasm, whereas others (e.g., TR) are located in the nucleus. After ligand binding, the cytoplasmically localized receptors translocate to the nucleus. There is growing evidence that certain nuclear receptors (e.g., glucocorticoid, estrogen) can also act at the membrane or in the cytoplasm to activate or repress signal transduction pathways, providing a mechanism for cross-talk between membrane and nuclear receptors.

The structures of nuclear receptors have been studied extensively, including by x-ray crystallography. The DNA binding domain, consisting of two zinc fingers, contacts specific DNA recognition sequences in target genes. Most nuclear receptors bind to DNA as dimers. Consequently, each monomer recognizes an individual DNA motif, referred to as a “half-site.” The steroid receptors, including the glucocorticoid, estrogen, progesterone, and androgen receptors, bind to DNA as homodimers. Consistent with this twofold symmetry, their DNA recognition half-sites are palindromic. The thyroid, retinoid, peroxisome

proliferator activated, and vitamin D receptors bind to DNA preferentially as heterodimers in combination with retinoid X receptors (RXRs). Their DNA half-sites are typically arranged as direct repeats.

The carboxy-terminal hormone-binding domain mediates transcriptional control. For type II receptors such as TR and retinoic acid receptor (RAR), co-repressor proteins bind to the receptor in the absence of ligand and silence gene transcription. Hormone binding induces conformational changes, triggering the release of co-repressors and inducing the recruitment of coactivators that stimulate transcription. Thus, these receptors are capable of mediating dramatic changes in the level of gene activity. Certain disease states are associated with defective regulation of these events. For example, mutations in the TR prevent co-repressor dissociation, resulting in an autosomal dominant form of hormone resistance (Chap. 405). In promyelocytic leukemia, fusion of RAR α to other nuclear proteins causes aberrant gene silencing that prevents normal cellular differentiation. Treatment with retinoic acid reverses this repression and allows cellular differentiation and apoptosis to occur. Most type I steroid receptors interact weakly with co-repressors, but ligand binding still induces interactions with an array of coactivators. X-ray crystallography shows that various SERMs induce distinct estrogen receptor conformations. The tissue-specific responses caused by these agents in breast, bone, and uterus appear to reflect distinct interactions with coactivators. The receptor-coactivator complex stimulates gene transcription by several pathways, including (1) recruitment of enzymes (histone acetyl transferases) that modify chromatin structure, (2) interactions with additional transcription factors on the target gene, and (3) direct interactions with components of the general transcription apparatus to enhance the rate of RNA polymerase II-mediated transcription. Studies of nuclear receptor-mediated transcription show that these are dynamic events that involve relatively rapid (e.g., 30–60 min) cycling of transcription complexes on any specific target gene.

FUNCTIONS OF HORMONES

The functions of individual hormones are described in detail in subsequent chapters. Nevertheless, it is useful to illustrate how most biologic responses require integration of several different hormone pathways. The physiologic functions of hormones can be divided into three general areas: (1) growth and differentiation, (2) maintenance of homeostasis, and (3) reproduction.

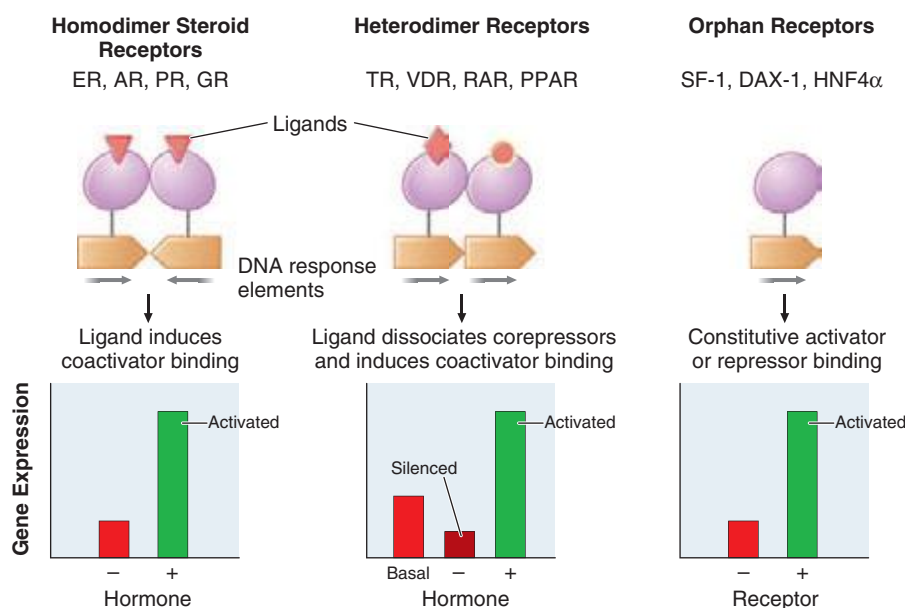


FIGURE 400e-2 Nuclear receptor signaling. AR, androgen receptor; DAX, dosage-sensitive sex-reversal, adrenal hypoplasia congenita, X-chromosome; ER, estrogen receptor; GR, glucocorticoid receptor; HNF4 α , hepatic nuclear factor 4 α ; PPAR, peroxisome proliferator activated receptor; PR, progesterone receptor; RAR, retinoic acid receptor; SF-1, steroidogenic factor-1; TR, thyroid hormone receptor; VDR, vitamin D receptor.

GROWTH

Multiple hormones and nutritional factors mediate the complex phenomenon of growth (Chap. 401e). Short stature may be caused by GH deficiency, hypothyroidism, Cushing's syndrome, precocious puberty, malnutrition, chronic illness, or genetic abnormalities that affect the epiphyseal growth plates (e.g., *FGFR3* and *SHOX* mutations). Many factors (GH, IGF-I, thyroid hormones) stimulate growth, whereas others (sex steroids) lead to epiphyseal closure. Understanding these hormonal interactions is important in the diagnosis and management of growth disorders. For example, delaying exposure to high levels of sex steroids may enhance the efficacy of GH treatment.

MAINTENANCE OF HOMEOSTASIS

Although virtually all hormones affect homeostasis, the most important among them are the following:

1. Thyroid hormone—controls about 25% of basal metabolism in most tissues
2. Cortisol—exerts a permissive action for many hormones in addition to its own direct effects
3. PTH—regulates calcium and phosphorus levels
4. Vasopressin—regulates serum osmolality by controlling renal free-water clearance
5. Mineralocorticoids—control vascular volume and serum electrolyte (Na^+ , K^+) concentrations
6. Insulin—maintains euglycemia in the fed and fasted states

The defense against hypoglycemia is an impressive example of integrated hormone action (Chap. 420). In response to the fasting state and falling blood glucose, insulin secretion is suppressed, resulting in decreased glucose uptake and enhanced glycogenolysis, lipolysis, proteolysis, and gluconeogenesis to mobilize fuel sources. If hypoglycemia develops (usually from insulin administration or sulfonylureas), an orchestrated counterregulatory response occurs—glucagon and epinephrine rapidly stimulate glycogenolysis and gluconeogenesis, whereas GH and cortisol act over several hours to raise glucose levels and antagonize insulin action.

Although free-water clearance is controlled primarily by vasopressin, cortisol and thyroid hormone are also important for facilitating renal tubular responses to vasopressin (Chap. 404). PTH and vitamin D function in an interdependent manner to control calcium metabolism (Chap. 423). PTH stimulates renal synthesis of 1,25-dihydroxyvitamin D, which increases calcium absorption in the gastrointestinal tract and enhances PTH action in bone. Increased calcium, along with vitamin D, feeds back to suppress PTH, thus maintaining calcium balance.

Depending on the severity of a specific stress and whether it is acute or chronic, multiple endocrine and cytokine pathways are activated to mount an appropriate physiologic response. In severe acute stress such as trauma or shock, the sympathetic nervous system is activated and catecholamines are released, leading to increased cardiac output and a primed musculoskeletal system. Catecholamines also increase mean blood pressure and stimulate glucose production. Multiple stress-induced pathways converge on the hypothalamus, stimulating several hormones, including vasopressin and corticotropin-releasing hormone (CRH). These hormones, in addition to cytokines (tumor necrosis factor α , interleukin [IL] 2, IL-6) increase ACTH and GH production. ACTH stimulates the adrenal gland, increasing cortisol, which in turn helps sustain blood pressure and dampen the inflammatory response. Increased vasopressin acts to conserve free water.

REPRODUCTION

The stages of reproduction include (1) sex determination during fetal development (Chap. 410); (2) sexual maturation during puberty (Chaps. 411 and 412); (3) conception, pregnancy, lactation, and child rearing (Chap. 412); and (4) cessation of reproductive capability at menopause (Chap. 413). Each of these stages involves an orchestrated interplay of multiple hormones, a phenomenon well illustrated by the dynamic hormonal changes that occur during each 28-day menstrual cycle. In the early follicular phase, pulsatile secretion of LH and FSH

stimulates the progressive maturation of the ovarian follicle. This results in gradually increasing estrogen and progesterone levels, leading to enhanced pituitary sensitivity to GnRH, which, when combined with accelerated GnRH secretion, triggers the LH surge and rupture of the mature follicle. Inhibin, a protein produced by the granulosa cells, enhances follicular growth and feeds back to the pituitary to selectively suppress FSH without affecting LH. Growth factors such as EGF and IGF-I modulate follicular responsiveness to gonadotropins. Vascular endothelial growth factor and prostaglandins play a role in follicle vascularization and rupture.

During pregnancy, the increased production of prolactin, in combination with placentally derived steroids (e.g., estrogen and progesterone), prepares the breast for lactation. Estrogens induce the production of progesterone receptors, allowing for increased responsiveness to progesterone. In addition to these and other hormones involved in lactation, the nervous system and oxytocin mediate the suckling response and milk release.

HORMONAL FEEDBACK REGULATORY SYSTEMS

Feedback control, both negative and positive, is a fundamental feature of endocrine systems. Each of the major hypothalamic-pituitary-hormone axes is governed by negative feedback, a process that maintains hormone levels within a relatively narrow range (Chap. 401e). Examples of hypothalamic-pituitary negative feedback include (1) thyroid hormones on the TRH-TSH axis, (2) cortisol on the CRH-ACTH axis, (3) gonadal steroids on the GnRH-LH/FSH axis, and (4) IGF-I on the growth hormone-releasing hormone (GHRH)-GH axis (Fig. 400e-3). These regulatory loops include both positive (e.g., TRH, TSH) and negative (e.g., T_4 , T_3) components, allowing for exquisite control of hormone levels. As an example, a small reduction of thyroid hormone triggers a rapid increase of TRH and TSH secretion, resulting in thyroid gland stimulation and increased thyroid hormone production. When thyroid hormone reaches a normal level, it feeds back to suppress TRH and TSH, and a new steady state is attained. Feedback regulation also occurs for endocrine systems that do not involve the pituitary gland, such as calcium feedback on PTH, glucose inhibition of insulin secretion, and leptin feedback on the hypothalamus. An

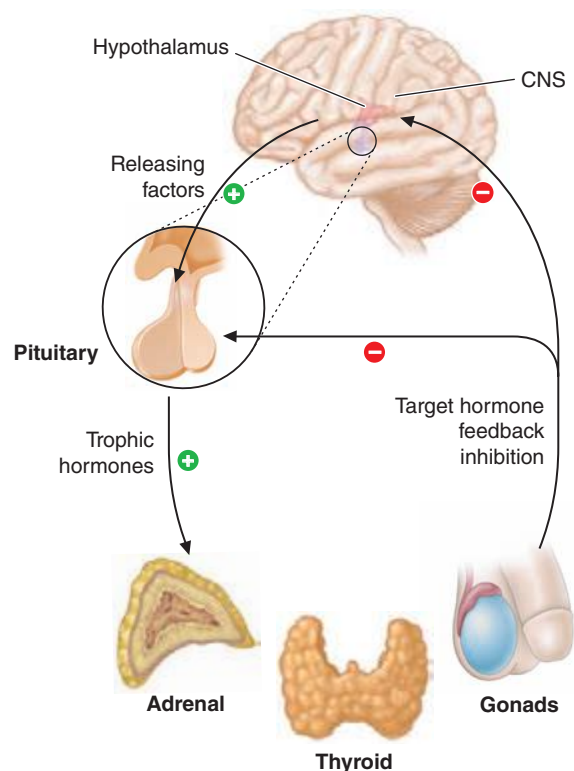


FIGURE 400e-3 Feedback regulation of endocrine axes. CNS, central nervous system.

400e-6 understanding of feedback regulation provides important insights into endocrine testing paradigms (see below).

Positive feedback control also occurs but is not well understood. The primary example is estrogen-mediated stimulation of the midcycle LH surge. Although chronic low levels of estrogen are inhibitory, gradually rising estrogen levels stimulate LH secretion. This effect, which is illustrative of an endocrine rhythm (see below), involves activation of the hypothalamic GnRH pulse generator. In addition, estrogen-primed gonadotropes are extraordinarily sensitive to GnRH, leading to amplification of LH release.

PARACRINE AND AUTOCRINE CONTROL

The previously mentioned examples of feedback control involve classic endocrine pathways in which hormones are released by one gland and act on a distant target gland. However, local regulatory systems, often involving growth factors, are increasingly recognized. *Paracrine regulation* refers to factors released by one cell that act on an adjacent cell in the same tissue. For example, somatostatin secretion by pancreatic islet δ cells inhibits insulin secretion from nearby β cells. *Autocrine regulation* describes the action of a factor on the same cell from which it is produced. IGF-I acts on many cells that produce it, including chondrocytes, breast epithelium, and gonadal cells. Unlike endocrine actions, paracrine and autocrine control are difficult to document because local growth factor concentrations cannot be measured readily.

Anatomic relationships of glandular systems also greatly influence hormonal exposure: the physical organization of islet cells enhances their intercellular communication; the portal vasculature of the hypothalamic-pituitary system exposes the pituitary to high concentrations of hypothalamic releasing factors; testicular seminiferous tubules gain exposure to high testosterone levels produced by the interdigitated Leydig cells; the pancreas receives nutrient information and local exposure to peptide hormones (incretins) from the gastrointestinal tract; and the liver is the proximal target of insulin action because of portal drainage from the pancreas.

HORMONAL RHYTHMS

The feedback regulatory systems described above are superimposed on hormonal rhythms that are used for adaptation to the environment. Seasonal changes, the daily occurrence of the light-dark cycle, sleep, meals, and stress are examples of the many environmental events that affect hormonal rhythms. The *menstrual cycle* is repeated on average every 28 days, reflecting the time required to follicular maturation and ovulation (**Chap. 412**). Essentially all pituitary hormone rhythms are entrained to sleep and to the *circadian cycle*, generating reproducible patterns that are repeated approximately every 24 h. The HPA axis, for

example, exhibits characteristic peaks of ACTH and cortisol production in the early morning, with a nadir during the night. Recognition of these rhythms is important for endocrine testing and treatment. Patients with Cushing's syndrome characteristically exhibit increased midnight cortisol levels compared with normal individuals (**Chap. 406**). In contrast, morning cortisol levels are similar in these groups, as cortisol is normally high at this time of day in normal individuals. The HPA axis is more susceptible to suppression by glucocorticoids administered at night as they blunt the early-morning rise of ACTH. Understanding these rhythms allows glucocorticoid replacement that mimics diurnal production by administering larger doses in the morning than in the afternoon. Disrupted sleep rhythms can alter hormonal regulation. For example, sleep deprivation causes mild insulin resistance, food craving, and hypertension, which are reversible, at least in the short term. Emerging evidence indicates that circadian clock pathways not only regulate sleep-wake cycles but also play important roles in virtually every cell type. For example, tissue-specific deletion of clock genes alters rhythms and levels of gene expression, as well as metabolic responses in liver, adipose, and other tissues.

Other endocrine rhythms occur on a more rapid time scale. Many peptide hormones are secreted in discrete bursts every few hours. LH and FSH secretion are exquisitely sensitive to GnRH pulse frequency. Intermittent pulses of GnRH are required to maintain pituitary sensitivity, whereas continuous exposure to GnRH causes pituitary gonadotropes desensitization. This feature of the hypothalamic-pituitary-gonadotrope axis forms the basis for using long-acting GnRH agonists to treat central precocious puberty or to decrease testosterone levels in the management of prostate cancer. It is important to be aware of the pulsatile nature of hormone secretion and the rhythmic patterns of hormone production in relating serum hormone measurements to normal values. For some hormones, integrated markers have been developed to circumvent hormonal fluctuations. Examples include 24-h urine collections for cortisol, IGF-I as a biologic marker of GH action, and HbA1c as an index of long-term (weeks to months) blood glucose control.

Often, one must interpret endocrine data only in the context of other hormones. For example, PTH levels typically are assessed in combination with serum calcium concentrations. A high serum calcium level in association with elevated PTH is suggestive of hyperparathyroidism, whereas a suppressed PTH in this situation is more likely to be caused by hypercalcemia of malignancy or other causes of hypercalcemia. Similarly, TSH should be elevated when T_4 and T_3 concentrations are low, reflecting reduced feedback inhibition. When this is not the case, it is important to consider secondary hypothyroidism, which is caused by a defect at the level of the pituitary.

401e Anterior Pituitary: Physiology of Pituitary Hormones

Shlomo Melmed, J. Larry Jameson

The anterior pituitary often is referred to as the “master gland” because, together with the hypothalamus, it orchestrates the complex regulatory functions of many other endocrine glands. The anterior pituitary gland produces six major hormones: (1) prolactin (PRL), (2) growth hormone (GH), (3) adrenocorticotrophic hormone (ACTH), (4) luteinizing hormone (LH), (5) follicle-stimulating hormone (FSH), and (6) thyroid-stimulating hormone (TSH) (Table 401e-1). Pituitary hormones are secreted in a pulsatile manner, reflecting stimulation by an array of specific hypothalamic releasing factors. Each of these pituitary hormones elicits specific responses in peripheral target tissues. The hormonal products of those peripheral glands, in turn, exert feedback control at the level of the hypothalamus and pituitary to modulate pituitary function (Fig. 401e-1). Pituitary tumors cause characteristic hormone excess syndromes. Hormone deficiency may be inherited or acquired. Fortunately, there are efficacious treatments for many pituitary hormone excess and deficiency syndromes. Nonetheless, these diagnoses are often elusive; this emphasizes the importance of recognizing subtle clinical manifestations and performing the correct laboratory diagnostic tests. **For discussion of disorders of the posterior pituitary, or neurohypophysitis, see Chap. 404.**

ANATOMY AND DEVELOPMENT

ANATOMY

The pituitary gland weighs ~600 mg and is located within the sella turcica ventral to the diaphragma sella; it consists of anatomically and functionally distinct anterior and posterior lobes. The bony sella is contiguous to vascular and neurologic structures, including the cavernous sinuses, cranial nerves, and optic chiasm. Thus, expanding intrasellar pathologic processes may have significant central mass effects in addition to their endocrinologic impact.

Hypothalamic neural cells synthesize specific releasing and inhibiting hormones that are secreted directly into the portal vessels of the pituitary stalk. Blood supply of the pituitary gland comes from the superior and inferior hypophyseal arteries (Fig. 401e-2). The hypothalamic-pituitary portal plexus provides the major blood source for the anterior pituitary, allowing reliable transmission of hypothalamic peptide pulses

without significant systemic dilution; consequently, pituitary cells are exposed to releasing or inhibiting factors and in turn release their hormones as discrete pulses into the systemic circulation (Fig. 401e-3).

The posterior pituitary is supplied by the inferior hypophyseal arteries. In contrast to the anterior pituitary, the posterior lobe is directly innervated by hypothalamic neurons (supraopticohypophyseal and tuberohypophyseal nerve tracts) via the pituitary stalk (Chap. 404). Thus, posterior pituitary production of vasopressin (antidiuretic hormone [ADH]) and oxytocin is particularly sensitive to neuronal damage by lesions that affect the pituitary stalk or hypothalamus.

PITUITARY DEVELOPMENT

The embryonic differentiation and maturation of anterior pituitary cells have been elucidated in considerable detail. Pituitary development from Rathke’s pouch involves a complex interplay of lineage-specific transcription factors expressed in pluripotent precursor cells and gradients of locally produced growth factors (Table 401e-1). The transcription factor Prop-1 induces pituitary development of Pit-1-specific lineages as well as gonadotropes. The transcription factor Pit-1 determines cell-specific expression of GH, PRL, and TSH in somatotropes, lactotropes, and thyrotropes. Expression of high levels of estrogen receptors in cells that contain Pit-1 favors PRL expression, whereas thyrotrope embryonic factor (TEF) induces TSH expression. Pit-1 binds to GH, PRL, and TSH gene regulatory elements as well as to recognition sites on its own promoter, providing a mechanism for maintaining specific pituitary hormone phenotypic stability. Gonadotrope cell development is further defined by the cell-specific expression of the nuclear receptors steroidogenic factor (SF-1) and *d* osage-sensitive sex reversal, *a* drenal hypoplasia critical region, on chromosome X, gene 1 (DAX-1). Development of corticotrope cells, which express the proopiomelanocortin (POMC) gene, requires the T-Pit transcription factor. Abnormalities of pituitary development caused by mutations of Pit-1, Prop-1, SF-1, DAX-1, and T-Pit result in a rare, selective or combined pituitary hormone deficit syndromes.

ANTERIOR PITUITARY HORMONES

Each anterior pituitary hormone is under unique control, and each exhibits highly specific normal and dysregulated secretory characteristics.

PROLACTIN

Synthesis PRL consists of 198 amino acids and has a molecular mass of 21,500 kDa; it is weakly homologous to GH and human placental lactogen (hPL), reflecting the duplication and divergence of a common GH-PRL-hPL precursor gene. PRL is synthesized in lactotropes,

TABLE 401e-1 ANTERIOR PITUITARY HORMONE EXPRESSION AND REGULATION

Cell	Corticotrope	Somatotrope	Lactotrope	Thyrotrope	Gonadotrope
Tissue-specific transcription factor	T-Pit	Prop-1, Pit-1	Prop-1, Pit-1	Prop-1, Pit-1, TEF	SF-1, DAX-1
Fetal appearance	6 weeks	8 weeks	12 weeks	12 weeks	12 weeks
Hormone	POMC	GH	PRL	TSH	FSH, LH
Protein	Polypeptide	Polypeptide	Polypeptide	Glycoprotein α , β subunits	Glycoprotein α , β subunits
Amino acids	266 (ACTH 1–39)	191	199	211	210, 204
Stimulators	CRH, AVP, gp-130 cytokines	GHRH, ghrelin	Estrogen, TRH, VIP	TRH	GnRH, activins, estrogen
Inhibitors	Glucocorticoids	Somatostatin, IGF-I	Dopamine	T ₃ , T ₄ , dopamine, somatostatin, glucocorticoids	Sex steroids, inhibin
Target gland	Adrenal	Liver, bone, other tissues	Breast, other tissues	Thyroid	Ovary, testis
Trophic effect	Steroid production	IGF-I production, growth induction, insulin antagonism	Milk production	T ₄ synthesis and secretion	Sex steroid production, follicle growth, germ cell maturation
Normal range	ACTH, 4–22 pg/L	<0.5 μ g/L ^a	M <15 μ g/L; F <20 μ g/L	0.1–5 mU/L	M, 5–20 IU/L; F (basal), 5–20 IU/L

^aHormone secretion integrated over 24 h.

Abbreviations: M, male; F, female. For other abbreviations, see text.

Source: Adapted from I Shimon, S Melmed, in S Melmed, P Conn (eds): *Endocrinology: Basic and Clinical Principles*. Totowa, NJ, Humana, 2005.

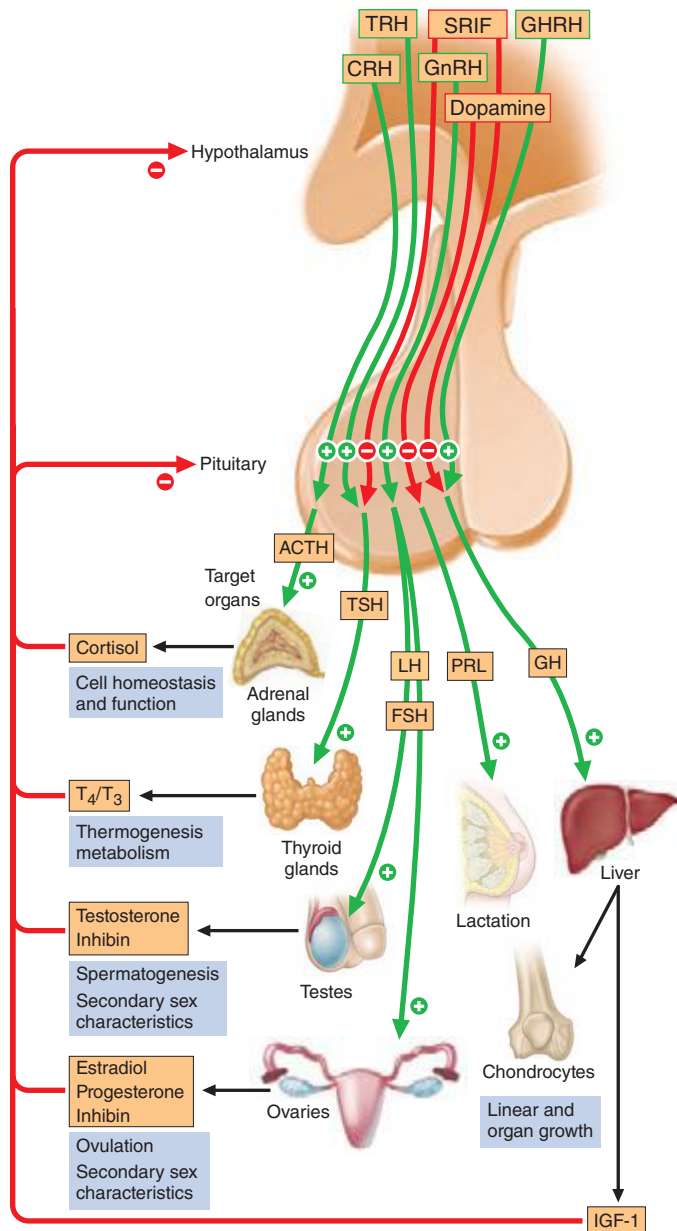


FIGURE 401e-1 Diagram of pituitary axes. Hypothalamic hormones regulate anterior pituitary trophic hormones that in turn determine target gland secretion. Peripheral hormones feed back to regulate hypothalamic and pituitary hormones. For abbreviations, see text.

which constitute about 20% of anterior pituitary cells. Lactotropes and somatotropes are derived from a common precursor cell that may give rise to a tumor that secretes both PRL and GH. Marked lactotrope cell hyperplasia develops during pregnancy and the first few months of lactation. These transient functional changes in the lactotrope population are induced by estrogen.

Secretion Normal adult serum PRL levels are about 10–25 $\mu\text{g/L}$ in women and 10–20 $\mu\text{g/L}$ in men. PRL secretion is pulsatile, with the highest secretory peaks occurring during rapid eye movement sleep. Peak serum PRL levels (up to 30 $\mu\text{g/L}$) occur between 4:00 and 6:00 A.M. The circulating half-life of PRL is about 50 min.

PRL is unique among the pituitary hormones in that the predominant central control mechanism is inhibitory, reflecting dopamine-mediated suppression of PRL release. This regulatory pathway accounts for the spontaneous PRL hypersecretion that occurs with pituitary stalk section, often a consequence of compressive mass lesions at the skull base. Pituitary dopamine type 2 (D_2) receptors mediate inhibition of

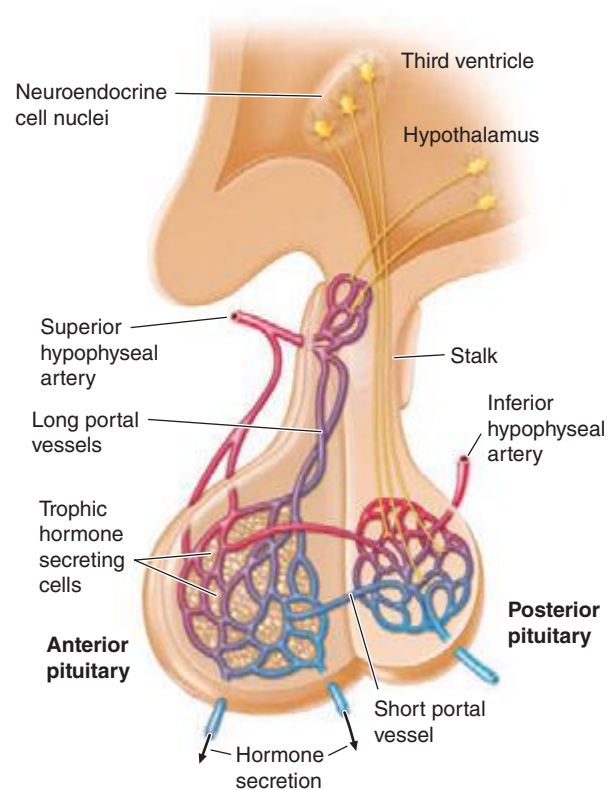


FIGURE 401e-2 Diagram of hypothalamic-pituitary vasculature. The hypothalamic nuclei produce hormones that traverse the portal system and impinge on anterior pituitary cells to regulate pituitary hormone secretion. Posterior pituitary hormones are derived from direct neural extensions.

PRL synthesis and secretion. Targeted disruption (gene knockout) of the murine D_2 receptor in mice results in hyperprolactinemia and lactotrope proliferation. As discussed below, dopamine agonists play a central role in the management of hyperprolactinemic disorders.

Thyrotropin-releasing hormone (TRH) (pyro Glu-His-Pro-NH₂) is a hypothalamic tripeptide that elicits PRL release within 15–30 min after intravenous injection. The physiologic relevance of TRH for PRL regulation is unclear, and it appears primarily to regulate TSH (**Chap. 405**). *Vasoactive intestinal peptide* (VIP) also induces PRL release, whereas glucocorticoids and thyroid hormone weakly suppress PRL secretion.

Serum PRL levels rise transiently after exercise, meals, sexual intercourse, minor surgical procedures, general anesthesia, chest wall injury, acute myocardial infarction, and other forms of acute stress. PRL levels increase markedly (about tenfold) during pregnancy and decline rapidly within 2 weeks of parturition. If breast-feeding is initiated, basal PRL levels remain elevated; suckling stimulates transient reflex increases in PRL levels that last for about 30–45 min. Breast suckling activates neural afferent pathways in the hypothalamus that

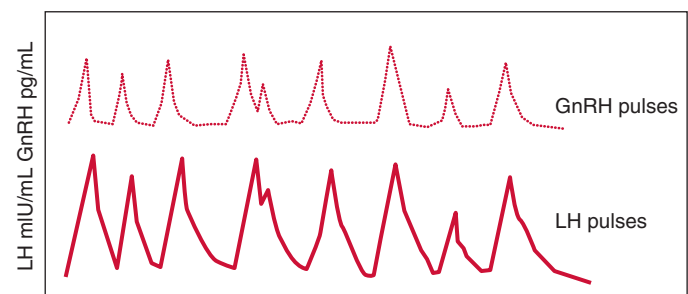


FIGURE 401e-3 Hypothalamic gonadotropin-releasing hormone (GnRH) pulses induce secretory pulses of luteinizing hormone (LH).

induce PRL release. With time, suckling-induced responses diminish and interfeeding PRL levels return to normal.

Action The PRL receptor is a member of the type I cytokine receptor family that also includes GH and interleukin (IL) 6 receptors. Ligand binding induces receptor dimerization and intracellular signaling by Janus kinase (JAK), which stimulates translocation of the signal transduction and activators of transcription (STAT) family to activate target genes. In the breast, the lobuloalveolar epithelium proliferates in response to PRL, placental lactogens, estrogen, progesterone, and local paracrine growth factors, including insulin-like growth factor I (IGF-I).

PRL acts to induce and maintain lactation, decrease reproductive function, and suppress sexual drive. These functions are geared toward ensuring that maternal lactation is sustained and not interrupted by pregnancy. PRL inhibits reproductive function by suppressing hypothalamic gonadotropin-releasing hormone (GnRH) and pituitary gonadotropin secretion and by impairing gonadal steroidogenesis in both women and men. In the ovary, PRL blocks folliculogenesis and inhibits granulosa cell aromatase activity, leading to hypoestrogenism and anovulation. PRL also has a luteolytic effect, generating a shortened, or inadequate, luteal phase of the menstrual cycle. In men, attenuated LH secretion leads to low testosterone levels and decreased spermatogenesis. These hormonal changes decrease libido and reduce fertility in patients with hyperprolactinemia.

GROWTH HORMONE

Synthesis GH is the most abundant anterior pituitary hormone, and GH-secreting somatotrope cells constitute up to 50% of the total anterior pituitary cell population. Mammosomatotrope cells, which coexpress PRL with GH, can be identified by using double immunostaining techniques. Somatotrope development and GH transcription are determined by expression of the cell-specific Pit-1 nuclear transcription factor. Five distinct genes encode GH and related proteins. The pituitary GH gene (*hGH-N*) produces two alternatively spliced products that give rise to 22-kDa GH (191 amino acids) and a less abundant 20-kDa GH molecule with similar biologic activity. Placental syncytiotrophoblast cells express a GH variant (*hGH-V*) gene; the related hormone human chorionic somatotropin (HCS) is expressed by distinct members of the gene cluster.

Secretion GH secretion is controlled by complex hypothalamic and peripheral factors. *GH-releasing hormone* (GHRH) is a 44-amino-acid hypothalamic peptide that stimulates GH synthesis and release. Ghrelin, an octanoylated gastric-derived peptide, and synthetic agonists of the *GHS-R* induce GHRH and also directly stimulate GH release. *Somatostatin* (somatotropin-release inhibiting factor [SRIF]) is synthesized in the medial preoptic area of the hypothalamus and inhibits GH secretion. GHRH is secreted in discrete spikes that elicit GH pulses, whereas SRIF sets basal GH secretory tone. SRIF also is expressed in many extrahypothalamic tissues, including the central nervous system (CNS), gastrointestinal tract, and pancreas, where it also acts to inhibit islet hormone secretion. *IGF-I*, the peripheral target hormone for GH, feeds back to inhibit GH; estrogen induces GH, whereas chronic glucocorticoid excess suppresses GH release.

Surface receptors on the somatotrope regulate GH synthesis and secretion. The GHRH receptor is a G protein-coupled receptor (GPCR) that signals through the intracellular cyclic AMP pathway to stimulate somatotrope cell proliferation as well as GH production. Inactivating mutations of the GHRH receptor cause profound dwarfism. A distinct surface receptor for ghrelin, the gastric-derived GH secretagogue, is expressed in both the hypothalamus and pituitary. Somatostatin binds to five distinct receptor subtypes (SSTR1 to SSTR5); SSTR2 and SSTR5 subtypes preferentially suppress GH (and TSH) secretion.

GH secretion is pulsatile, with highest peak levels occurring at night, generally correlating with sleep onset. GH secretory rates decline markedly with age so that hormone levels in middle age are about 15% of pubertal levels. These changes are paralleled by an age-related decline in lean muscle mass. GH secretion is also reduced in obese individuals, although IGF-I levels may not be suppressed, suggesting a change in

the setpoint for feedback control. Elevated GH levels occur within an hour of deep sleep onset as well as after exercise, physical stress, and trauma and during sepsis. Integrated 24-h GH secretion is higher in women and is also enhanced by estrogen replacement likely reflective of increased peripheral GH-resistance. Using standard assays, random GH measurements are undetectable in ~50% of daytime samples obtained from healthy subjects and are also undetectable in most obese and elderly subjects. Thus, single random GH measurements do not distinguish patients with adult GH deficiency from normal persons.

GH secretion is profoundly influenced by nutritional factors. Using newer ultrasensitive GH assays with a sensitivity of 0.002 µg/L, a glucose load suppresses GH to <0.7 µg/L in women and to <0.07 µg/L in men. Increased GH pulse frequency and peak amplitudes occur with chronic malnutrition or prolonged fasting. GH is stimulated by intravenous L-arginine, dopamine, and apomorphine (a dopamine receptor agonist), as well as by α-adrenergic pathways. β-Adrenergic blockade induces basal GH and enhances GHRH- and insulin-evoked GH release.

Action The pattern of GH secretion may affect tissue responses. The higher GH pulsatility observed in men compared with the relatively continuous basal GH secretion in women may be an important biologic determinant of linear growth patterns and liver enzyme induction.

The 70-kDa peripheral GH receptor protein has structural homology with the cytokine/hematopoietic superfamily. A fragment of the receptor extracellular domain generates a soluble GH binding protein (GHBP) that interacts with GH in the circulation. The liver and cartilage contain the greatest number of GH receptors. GH binding to preformed receptor dimers is followed by internal rotation and subsequent signaling through the JAK/STAT pathway. Activated STAT proteins translocate to the nucleus, where they modulate expression of GH-regulated target genes. GH analogues that bind to the receptor but are incapable of mediating receptor signaling are potent antagonists of GH action. A GH receptor antagonist (pegvisomant) is approved for treatment of acromegaly.

GH induces protein synthesis and nitrogen retention and impairs glucose tolerance by antagonizing insulin action. GH also stimulates lipolysis, leading to increased circulating fatty acid levels, reduced omental fat mass, and enhanced lean body mass. GH promotes sodium, potassium, and water retention and elevates serum levels of inorganic phosphate. Linear bone growth occurs as a result of complex hormonal and growth factor actions, including those of IGF-I. GH stimulates epiphyseal prechondrocyte differentiation. These precursor cells produce IGF-I locally, and their proliferation is also responsive to the growth factor.

Insulin-Like Growth Factors Although GH exerts direct effects in target tissues, many of its physiologic effects are mediated indirectly through IGF-I, a potent growth and differentiation factor. The liver is the major source of circulating IGF-I. In peripheral tissues, IGF-I also exerts local paracrine actions that appear to be both dependent on and independent of GH. Thus, GH administration induces circulating IGF-I as well as stimulating local IGF-I production in multiple tissues.

Both IGF-I and IGF-II are bound to high-affinity circulating IGF-binding proteins (IGFBPs) that regulate IGF bioactivity. Levels of IGFBP3 are GH-dependent, and it serves as the major carrier protein for circulating IGF-I. GH deficiency and malnutrition usually are associated with low IGFBP3 levels. IGFBP1 and IGFBP2 regulate local tissue IGF action but do not bind appreciable amounts of circulating IGF-I.

Serum IGF-I concentrations are profoundly affected by physiologic factors. Levels increase during puberty, peak at 16 years, and subsequently decline by >80% during the aging process. IGF-I concentrations are higher in women than in men. Because GH is the major determinant of hepatic IGF-I synthesis, abnormalities of GH synthesis or action (e.g., pituitary failure, GHRH receptor defect, GH receptor defect or pharmacologic GH receptor blockade) reduce IGF-I levels. Hypocaloric states are associated with GH resistance; IGF-I levels are therefore low with cachexia, malnutrition, and sepsis. In acromegaly, IGF-I levels are invariably high and reflect a log-linear relationship with circulating GH concentrations.

401e-4 IGF-I PHYSIOLOGY Injected IGF-I (100 µg/kg) induces hypoglycemia, and lower doses improve insulin sensitivity in patients with severe insulin resistance and diabetes. In cachectic subjects, IGF-I infusion (12 µg/kg per hour) enhances nitrogen retention and lowers cholesterol levels. Longer-term subcutaneous IGF-I injections enhance protein synthesis and are anabolic. Although bone formation markers are induced, bone turnover also may be stimulated by IGF-I. IGF-I has only been approved for use in patients with GH-resistance syndromes.

IGF-I side effects are dose-dependent, and overdose may result in hypoglycemia, hypotension, fluid retention, temporomandibular jaw pain, and increased intracranial pressure, all of which are reversible. Avascular femoral head necrosis has been reported. Chronic excess IGF-I administration presumably would result in features of acromegaly.

ADRENOCORTICOTROPIC HORMONE

(See also Chap. 406)

Synthesis ACTH-secreting corticotrope cells constitute about 20% of the pituitary cell population. ACTH (39 amino acids) is derived from the POMC precursor protein (266 amino acids) that also generates several other peptides, including β-lipotropin, β-endorphin, met-enkephalin, α-melanocyte-stimulating hormone (α-MSH), and corticotropin-like intermediate lobe protein (CLIP). The POMC gene is potently suppressed by glucocorticoids and induced by corticotropin-releasing hormone (CRH), arginine vasopressin (AVP), and proinflammatory cytokines, including IL-6, as well as leukemia inhibitory factor.

CRH, a 41-amino-acid hypothalamic peptide synthesized in the paraventricular nucleus as well as in higher brain centers, is the predominant stimulator of ACTH synthesis and release. The CRH receptor is a GPCR that is expressed on the corticotrope and signals to induce POMC transcription.

Secretion ACTH secretion is pulsatile and exhibits a characteristic circadian rhythm, peaking at about 6 A.M. and reaching a nadir about midnight. Adrenal glucocorticoid secretion, which is driven by ACTH, follows a parallel diurnal pattern. ACTH circadian rhythmicity is determined by variations in secretory pulse amplitude rather than changes in pulse frequency. Superimposed on this endogenous rhythm, ACTH levels are increased by physical and psychological stress, exercise, acute illness, and insulin-induced hypoglycemia.

Glucocorticoid-mediated negative regulation of the hypothalamic-pituitary-adrenal (HPA) axis occurs as a consequence of both hypothalamic CRH suppression and direct attenuation of pituitary POMC gene expression and ACTH release. In contrast, loss of cortisol feedback inhibition, as occurs in primary adrenal failure, results in extremely high ACTH levels.

Acute inflammatory or septic insults activate the HPA axis through the integrated actions of proinflammatory cytokines, bacterial toxins, and neural signals. The overlapping cascade of ACTH-inducing cytokines (tumor necrosis factor [TNF]; IL-1, -2, and -6; and leukemia inhibitory factor) activates hypothalamic CRH and AVP secretion, pituitary POMC gene expression, and local pituitary paracrine cytokine networks. The resulting cortisol elevation restrains the inflammatory response and enables host protection. Concomitantly, cytokine-mediated central glucocorticoid receptor resistance impairs glucocorticoid suppression of the HPA. Thus, the neuroendocrine stress response reflects the net result of highly integrated hypothalamic, intrapituitary, and peripheral hormone and cytokine signals acting to regulate cortisol secretion.

Action The major function of the HPA axis is to maintain metabolic homeostasis and mediate the neuroendocrine stress response. ACTH induces adrenocortical steroidogenesis by sustaining adrenal cell proliferation and function. The receptor for ACTH, designated *melanocortin-2 receptor*, is a GPCR that induces steroidogenesis by stimulating a cascade of steroidogenic enzymes (Chap. 406).

GONADOTROPINS: FSH AND LH

Synthesis and Secretion Gonadotrope cells constitute about 10% of anterior pituitary cells and produce two gonadotropin hormones—

LH and FSH. Like TSH and hCG, LH and FSH are glycoprotein hormones that comprise α and β subunits. The α subunit is common to these glycoprotein hormones; specificity of hormone function is conferred by the β subunits, which are expressed by separate genes.

Gonadotropin synthesis and release are dynamically regulated. This is particularly true in women, in whom rapidly fluctuating gonadal steroid levels vary throughout the menstrual cycle. Hypothalamic GnRH, a 10-amino-acid peptide, regulates the synthesis and secretion of both LH and FSH. Brain kisspeptin, a product of the *KISS1* gene regulates hypothalamic GnRH release. GnRH is secreted in discrete pulses every 60–120 min, and the pulses in turn elicit LH and FSH pulses (Fig. 401e-3). The pulsatile mode of GnRH input is essential to its action; pulses prime gonadotrope responsiveness, whereas continuous GnRH exposure induces desensitization. Based on this phenomenon, long-acting GnRH agonists are used to suppress gonadotropin levels in children with precocious puberty and in men with prostate cancer (Chap. 115) and are used in some ovulation-induction protocols to reduce levels of endogenous gonadotropins (Chap. 412). Estrogens act at both the hypothalamus and the pituitary to modulate gonadotropin secretion. Chronic estrogen exposure is inhibitory, whereas rising estrogen levels, as occur during the preovulatory surge, exert positive feedback to increase gonadotropin pulse frequency and amplitude. Progesterone slows GnRH pulse frequency but enhances gonadotropin responses to GnRH. Testosterone feedback in men also occurs at the hypothalamic and pituitary levels and is mediated in part by its conversion to estrogens.

Although GnRH is the main regulator of LH and FSH secretion, FSH synthesis is also under separate control by the gonadal peptides inhibin and activin, which are members of the transforming growth factor β (TGF-β) family. Inhibin selectively suppresses FSH, whereas activin stimulates FSH synthesis (Chap. 412).

Action The gonadotropin hormones interact with their respective GPCRs expressed in the ovary and testis, evoking germ cell development and maturation and steroid hormone biosynthesis. In women, FSH regulates ovarian follicle development and stimulates ovarian estrogen production. LH mediates ovulation and maintenance of the corpus luteum. In men, LH induces Leydig cell testosterone synthesis and secretion, and FSH stimulates seminiferous tubule development and regulates spermatogenesis.

THYROID-STIMULATING HORMONE

Synthesis and Secretion TSH-secreting thyrotrope cells constitute 5% of the anterior pituitary cell population. TSH shares a common α subunit with LH and FSH but contains a specific TSH β subunit. TRH is a hypothalamic tripeptide (pyroglutamyl histidylprolinamide) that acts through a pituitary GPCR to stimulate TSH synthesis and secretion; it also stimulates the lactotrope cell to secrete PRL. TSH secretion is stimulated by TRH, whereas thyroid hormones, dopamine, somatostatin, and glucocorticoids suppress TSH by overriding TRH induction.

Thyrotrope cell proliferation and TSH secretion are both induced when negative feedback inhibition by thyroid hormones is removed. Thus, thyroid damage (including surgical thyroidectomy), radiation-induced hypothyroidism, chronic thyroiditis, and prolonged goitrogen exposure are associated with increased TSH levels. Long-standing untreated hypothyroidism can lead to elevated TSH levels as well as thyrotrope hyperplasia and pituitary enlargement, which may be evident on magnetic resonance imaging.

Action TSH is secreted in pulses, although the excursions are modest in comparison to other pituitary hormones because of the low amplitude of the pulses and the relatively long half-life of TSH. Consequently, single determinations of TSH suffice to precisely assess its circulating levels. TSH binds to a GPCR on thyroid follicular cells to stimulate thyroid hormone synthesis and release (Chap. 405).

402 Hypopituitarism

Shlomo Melmed, J. Larry Jameson

Inadequate production of anterior pituitary hormones leads to features of hypopituitarism. Impaired production of one or more of the anterior pituitary trophic hormones can result from inherited disorders; more commonly, adult hypopituitarism is acquired and reflects the compressive mass effects of tumors or the consequences of local pituitary or hypothalamic traumatic, inflammatory, or vascular damage. These processes also may impair synthesis or secretion of hypothalamic hormones, with resultant pituitary failure (Table 402-1).

DEVELOPMENTAL AND GENETIC CAUSES OF HYPOPITUITARISM

Pituitary Dysplasia Pituitary dysplasia may result in aplastic, hypoplastic, or ectopic pituitary gland development. Because pituitary development follows midline cell migration from the nasopharyngeal Rathke's pouch, midline craniofacial disorders may be associated with

TABLE 402-1 ETIOLOGY OF HYPOPITUITARISM^a

Development/structural
Transcription factor defect
Pituitary dysplasia/aplasia
Congenital central nervous system mass, encephalocele
Primary empty sella
Congenital hypothalamic disorders (septo-optic dysplasia, Prader-Willi syndrome, Laurence-Moon-Biedl syndrome, Kallmann syndrome)
Traumatic
Surgical resection
Radiation damage
Head injuries
Neoplastic
Pituitary adenoma
Parasellar mass (germinoma, ependymoma, glioma)
Rathke's cyst
Craniopharyngioma
Hypothalamic hamartoma, gangliocytoma
Pituitary metastases (breast, lung, colon carcinoma)
Lymphoma and leukemia
Meningioma
Infiltrative/inflammatory
Lymphocytic hypophysitis
Hemochromatosis
Sarcoidosis
Histiocytosis X
Granulomatous hypophysitis
Transcription factor antibodies
Vascular
Pituitary apoplexy
Pregnancy-related (infarction with diabetes; postpartum necrosis)
Sickle cell disease
Arteritis
Infections
Fungal (histoplasmosis)
Parasitic (toxoplasmosis)
Tuberculosis
<i>Pneumocystis carinii</i>

^aTrophic hormone failure associated with pituitary compression or destruction usually occurs sequentially: growth hormone > follicle-stimulating hormone > luteinizing hormone > thyroid-stimulating hormone > adrenocorticotropic hormone. During childhood, growth retardation is often the presenting feature, and in adults, hypogonadism is the earliest symptom.

2256 pituitary dysplasia. Acquired pituitary failure in the newborn also can be caused by birth trauma, including cranial hemorrhage, asphyxia, and breech delivery.

SEPTO-OPTIC DYSPLASIA Hypothalamic dysfunction and hypopituitarism may result from dysgenesis of the septum pellucidum or corpus callosum. Affected children have mutations in the *HESX1* gene, which is involved in early development of the ventral prosencephalon. These children exhibit variable combinations of cleft palate, syndactyly, ear deformities, hypertelorism, optic nerve hypoplasia, micropenis, and anosmia. Pituitary dysfunction leads to diabetes insipidus, growth hormone (GH) deficiency and short stature, and, occasionally, thyroid-stimulating hormone (TSH) deficiency.

Tissue-Specific Factor Mutations Several pituitary cell-specific transcription factors, such as Pit-1 and Prop-1, are critical for determining the development and committed function of differentiated anterior pituitary cell lineages. Autosomal dominant or recessive Pit-1 mutations cause combined GH, prolactin (PRL), and TSH deficiencies. These patients usually present with growth failure and varying degrees of hypothyroidism. The pituitary may appear hypoplastic on magnetic resonance imaging (MRI).

Prop-1 is expressed early in pituitary development and appears to be required for Pit-1 function. Familial and sporadic *PROPI* mutations result in combined GH, PRL, TSH, and gonadotropin deficiency. Over 80% of these patients have growth retardation; by adulthood, all are deficient in TSH and gonadotropins, and a small minority later develop adrenocorticotrophic hormone (ACTH) deficiency. Because of gonadotropin deficiency, these individuals do not enter puberty spontaneously. In some cases, the pituitary gland appears enlarged on MRI. *TPIT* mutations result in ACTH deficiency associated with hypocortisolism.

Developmental Hypothalamic Dysfunction • KALLMANN SYNDROME Kallmann syndrome results from defective hypothalamic gonadotropin-releasing hormone (GnRH) synthesis and is associated with anosmia or hyposmia due to olfactory bulb agenesis or hypoplasia (**Chap. 411**). Classically, the syndrome may also be associated with color blindness, optic atrophy, nerve deafness, cleft palate, renal abnormalities, cryptorchidism, and neurologic abnormalities such as mirror movements. The initial genetic cause was identified in the X-linked *KAL* gene, mutations of which impair embryonic migration of GnRH neurons from the hypothalamic olfactory placode to the hypothalamus. Based on further studies, at least a dozen other genetic abnormalities, in addition to *KAL* mutations, have been found to cause isolated GnRH deficiency. Autosomal recessive (i.e., *GPR54*, *KISS1*) and dominant (i.e., *FGFR1*) modes of transmission have been described, and there is a growing list of genes associated with GnRH deficiency (*GNRH1*, *PROK2*, *PROKR2*, *CH7*, *PCSK1*, *FGF8*, *NELF*, *WDR11*, *TAC3*, *TACR3*). A fraction of patients have digenic mutations. Associated clinical features, in addition to GnRH deficiency, vary depending on the genetic cause. GnRH deficiency prevents progression through puberty. Males present with delayed puberty and pronounced hypogonadal features, including micropenis, probably the result of low testosterone levels during infancy. Females present with primary amenorrhea and failure of secondary sexual development.

Kallmann syndrome and other causes of congenital GnRH deficiency are characterized by low luteinizing hormone (LH) and follicle-stimulating hormone (FSH) levels and low concentrations of sex steroids (testosterone or estradiol). In sporadic cases of isolated gonadotropin deficiency, the diagnosis is often one of exclusion after other known causes of hypothalamic-pituitary dysfunction have been eliminated. Repetitive GnRH administration restores normal pituitary gonadotropin responses, pointing to a hypothalamic defect in these patients.

Long-term treatment of males with human chorionic gonadotropin (hCG) or testosterone restores pubertal development and secondary sex characteristics; women can be treated with cyclic estrogen and progestin. Fertility also may be restored by the administration of gonadotropins or by using a portable infusion pump to deliver subcutaneous, pulsatile GnRH.

BARDET-BIEDL SYNDROME This very rare genetically heterogeneous disorder is characterized by mental retardation, renal abnormalities, obesity, and hexadactyly, brachydactyly, or syndactyly. Central diabetes insipidus may or may not be associated. GnRH deficiency occurs in 75% of males and half of affected females. Retinal degeneration begins in early childhood, and most patients are blind by age 30. Numerous subtypes of Bardet-Biedl syndrome (BBS) have been identified, with genetic linkage to at least nine different loci. Several of the loci encode genes involved in basal body cilia function, and this may account for the diverse clinical manifestations.

LEPTIN AND LEPTIN RECEPTOR MUTATIONS Deficiencies of leptin or its receptor cause a broad spectrum of hypothalamic abnormalities, including hyperphagia, obesity, and central hypogonadism (**Chap. 415e**). Decreased GnRH production in these patients results in attenuated pituitary FSH and LH synthesis and release.

PRADER-WILLI SYNDROME This is a contiguous gene syndrome that results from deletion of the paternal copies of the imprinted *SNRPN* gene, the *NECDIN* gene, and possibly other genes on chromosome 15q. Prader-Willi syndrome is associated with hypogonadotropic hypogonadism, hyperphagia-obesity, chronic muscle hypotonia, mental retardation, and adult-onset diabetes mellitus (**Chap. 83e**). Multiple somatic defects also involve the skull, eyes, ears, hands, and feet. Diminished hypothalamic oxytocin- and vasopressin-producing nuclei have been reported. Deficient GnRH synthesis is suggested by the observation that chronic GnRH treatment restores pituitary LH and FSH release.

ACQUIRED HYPOPITUITARISM

Hypopituitarism may be caused by accidental or neurosurgical trauma; vascular events such as apoplexy; pituitary or hypothalamic neoplasms, craniopharyngioma, lymphoma, or metastatic tumors; inflammatory disease such as lymphocytic hypophysitis; infiltrative disorders such as sarcoidosis, hemochromatosis (**Chap. 428**), and tuberculosis; or irradiation.

Increasing evidence suggests that patients with brain injury, including contact sports trauma, subarachnoid hemorrhage, and irradiation, have transient hypopituitarism and require intermittent long-term endocrine follow-up, because permanent hypothalamic or pituitary dysfunction will develop in 25–40% of these patients.

Hypothalamic Infiltration Disorders These disorders—including sarcoidosis, histiocytosis X, amyloidosis, and hemochromatosis—frequently involve both hypothalamic and pituitary neuronal and neurochemical tracts. Consequently, diabetes insipidus occurs in half of patients with these disorders. Growth retardation is seen if attenuated GH secretion occurs before puberty. Hypogonadotropic hypogonadism and hyperprolactinemia are also common.

Inflammatory Lesions Pituitary damage and subsequent secretory dysfunction can be seen with chronic site infections such as tuberculosis, with opportunistic fungal infections associated with AIDS, and in tertiary syphilis. Other inflammatory processes, such as granulomas and sarcoidosis, may mimic the features of a pituitary adenoma. These lesions may cause extensive hypothalamic and pituitary damage, leading to trophic hormone deficiencies.

Cranial Irradiation Cranial irradiation may result in long-term hypothalamic and pituitary dysfunction, especially in children and adolescents, as they are more susceptible to damage after whole-brain or head and neck therapeutic irradiation. The development of hormonal abnormalities correlates strongly with irradiation dosage and the time interval after completion of radiotherapy. Up to two-thirds of patients ultimately develop hormone insufficiency after a median dose of 50 Gy (5000 rad) directed at the skull base. The development of hypopituitarism occurs over 5–15 years and usually reflects hypothalamic damage rather than primary destruction of pituitary cells. Although the pattern of hormone loss is variable, GH deficiency is most common, followed by gonadotropin and ACTH deficiency. When deficiency of one or more hormones is documented, the possibility of diminished reserve of other hormones is likely. Accordingly, anterior pituitary function should be continually

evaluated over the long term in previously irradiated patients, and replacement therapy instituted when appropriate (see below).

Lymphocytic Hypophysitis This occurs most often in postpartum women; it usually presents with hyperprolactinemia and MRI evidence of a prominent pituitary mass that often resembles an adenoma, with mildly elevated PRL levels. Pituitary failure caused by diffuse lymphocytic infiltration may be transient or permanent but requires immediate evaluation and treatment. Rarely, isolated pituitary hormone deficiencies have been described, suggesting a selective autoimmune process targeted to specific cell types. Most patients manifest symptoms of progressive mass effects with headache and visual disturbance. The erythrocyte sedimentation rate often is elevated. Because the MRI image may be indistinguishable from that of a pituitary adenoma, hypophysitis should be considered in a postpartum woman with a newly diagnosed pituitary mass before an unnecessary surgical intervention is undertaken. The inflammatory process often resolves after several months of glucocorticoid treatment, and pituitary function may be restored, depending on the extent of damage.

Pituitary Apoplexy Acute intrapituitary hemorrhagic vascular events can cause substantial damage to the pituitary and surrounding sellar structures. Pituitary apoplexy may occur spontaneously in a preexisting adenoma; postpartum (Sheehan's syndrome); or in association with diabetes, hypertension, sickle cell anemia, or acute shock. The hyperplastic enlargement of the pituitary, which occurs normally during pregnancy, increases the risk for hemorrhage and infarction. Apoplexy is an endocrine emergency that may result in severe hypoglycemia, hypotension and shock, central nervous system (CNS) hemorrhage, and death. Acute symptoms may include severe headache with signs of meningeal irritation, bilateral visual changes, ophthalmoplegia, and, in severe cases, cardiovascular collapse and loss of consciousness. Pituitary computed tomography (CT) or MRI may reveal signs of intratumoral or sellar hemorrhage, with pituitary stalk deviation and compression of pituitary tissue.

Patients with no evident visual loss or impaired consciousness can be observed and managed conservatively with high-dose glucocorticoids. Those with significant or progressive visual loss, cranial nerve palsy, or loss of consciousness require urgent surgical decompression. Visual recovery after sellar surgery is inversely correlated with the length of time after the acute event. Therefore, severe ophthalmoplegia or visual deficits are indications for early surgery. Hypopituitarism is common after apoplexy.

Empty Sella A partial or apparently totally empty sella is often an incidental MRI finding, and may be associated with intracranial hypertension. These patients usually have normal pituitary function, implying that the surrounding rim of pituitary tissue is fully functional. Hypopituitarism, however, may develop insidiously. Pituitary masses also may undergo clinically silent infarction and involution with development of a partial or totally empty sella by cerebrospinal fluid (CSF) filling the dural herniation. Rarely, small but functional pituitary adenomas may arise within the rim of normal pituitary tissue, and they are not always visible on MRI.

PRESENTATION AND DIAGNOSIS

The clinical manifestations of hypopituitarism depend on which hormones are lost and the extent of the hormone deficiency. GH deficiency causes growth disorders in children and leads to abnormal body composition in adults (see below). Gonadotropin deficiency causes menstrual disorders and infertility in women and decreased sexual function, infertility, and loss of secondary sexual characteristics in men. TSH and ACTH deficiency usually develop later in the course of pituitary failure. TSH deficiency causes growth retardation in children and features of hypothyroidism in children and adults. The secondary form of adrenal insufficiency caused by ACTH deficiency leads to hypocortisolism with relative preservation of mineralocorticoid production. PRL deficiency causes failure of lactation. When lesions involve the posterior pituitary, polyuria and polydipsia reflect loss of vasopressin secretion. In patients with long-standing pituitary damage, epidemiologic studies document

an increased mortality rate, primarily from increased cardiovascular and cerebrovascular disease. Previous head or neck irradiation is also a determinant of increased mortality rates in patients with hypopituitarism, especially from cerebrovascular disease.

LABORATORY INVESTIGATION

Biochemical diagnosis of pituitary insufficiency is made by demonstrating low levels of respective pituitary trophic hormones in the setting of low levels of target hormones. For example, low free thyroxine in the setting of a low or inappropriately normal TSH level suggests secondary hypothyroidism. Similarly, a low testosterone level without elevation of gonadotropins suggests hypogonadotropic hypogonadism. Provocative tests may be required to assess pituitary reserve (**Table 402-2**). GH responses to insulin-induced hypoglycemia, arginine, L-dopa, growth hormone-releasing hormone (GHRH), or growth hormone-releasing peptides (GHRPs) can be used to assess GH reserve. Corticotropin-releasing hormone (CRH) administration induces ACTH release, and administration of synthetic ACTH (cosyntropin) evokes adrenal cortisol release as an indirect indicator of pituitary ACTH reserve (**Chap. 406**). ACTH reserve is most reliably assessed by measuring ACTH and cortisol levels during insulin-induced hypoglycemia. However, this test should be performed cautiously in patients with suspected adrenal insufficiency because of enhanced susceptibility to hypoglycemia and hypotension. Administering insulin to induce hypoglycemia is contraindicated in patients with active coronary artery disease or known seizure disorders.

TREATMENT HYPOPITUITARISM

Hormone replacement therapy, including glucocorticoids, thyroid hormone, sex steroids, growth hormone, and vasopressin, is usually safe and free of complications. Treatment regimens that mimic physiologic hormone production allow for maintenance of satisfactory clinical homeostasis. Effective dosage schedules are outlined in **Table 402-3**. Patients in need of glucocorticoid replacement require careful dose adjustments during stressful events such as acute illness, dental procedures, trauma, and acute hospitalization.

DISORDERS OF GROWTH AND DEVELOPMENT

Skeletal Maturation and Somatic Growth The growth plate is dependent on a variety of hormonal stimuli, including GH, insulin-like growth factor (IGF) I, sex steroids, thyroid hormones, paracrine growth factors, and cytokines. The growth-promoting process also requires caloric energy, amino acids, vitamins, and trace metals and consumes about 10% of normal energy production. Malnutrition impairs chondrocyte activity, increases GH resistance, and reduces circulating IGF-I and IGFBP3 levels.

Linear bone growth rates are very high in infancy and are pituitary-dependent. Mean growth velocity is ~6 cm/year in later childhood and usually is maintained within a given range on a standardized percentile chart. Peak growth rates occur during midpuberty when bone age is 12 (girls) or 13 (boys). Secondary sexual development is associated with elevated sex steroids that cause progressive epiphyseal growth plate closure. *Bone age* is delayed in patients with all forms of true GH deficiency or GH receptor defects that result in attenuated GH action.

Short stature may occur as a result of constitutive intrinsic growth defects or because of acquired extrinsic factors that impair growth. In general, delayed bone age in a child with short stature is suggestive of a hormonal or systemic disorder, whereas normal bone age in a short child is more likely to be caused by a genetic cartilage dysplasia or growth plate disorder (**Chap. 427**).

GH Deficiency in Children • GH DEFICIENCY Isolated GH deficiency is characterized by short stature, micropenis, increased fat, high-pitched voice, and a propensity to hypoglycemia due to relatively unopposed insulin action. Familial modes of inheritance are seen in at least one-third of these individuals and may be autosomal dominant, recessive,

TABLE 402-2 TESTS OF PITUITARY SUFFICIENCY

Hormone	Test	Blood Samples	Interpretation
Growth hormone (GH)	Insulin tolerance test: Regular insulin (0.05–0.15 U/kg IV)	–30, 0, 30, 60, 120 min for glucose and GH	Glucose <40 mg/dL; GH should be >3 µg/L
	GHRH test: 1 µg/kg IV	0, 15, 30, 45, 60, 120 min for GH	Normal response is GH >3 µg/L
	L-Arginine test: 30 g IV over 30 min	0, 30, 60, 120 min for GH	Normal response is GH >3 µg/L
	L-Dopa test: 500 mg PO	0, 30, 60, 120 min for GH	Normal response is GH >3 µg/L
Prolactin	TRH test: 200–500 µg IV	0, 20, and 60 min for TSH and PRL	Normal prolactin is >2 µg/L and increase >200% of baseline
ACTH	Insulin tolerance test: regular insulin (0.05–0.15 U/kg IV)	–30, 0, 30, 60, 90 min for glucose and cortisol	Glucose <40 mg/dL Cortisol should increase by >7 µg/dL or to >20 µg/dL
	CRH test: 1 µg/kg ovine CRH IV at 8 A.M.	0, 15, 30, 60, 90, 120 min for ACTH and cortisol	Basal ACTH increases 2- to 4-fold and peaks at 20–100 pg/mL Cortisol levels >20–25 µg/dL
	Metypapone test: Metypapone (30 mg/kg) at midnight	Plasma 11-deoxycortisol and cortisol at 8 A.M.; ACTH can also be measured	Plasma cortisol should be <4 g/dL to assure an adequate response Normal response is 11-deoxycortisol >7.5 µg/dL or ACTH >75 pg/mL
	Standard ACTH stimulation test: ACTH 1-24 (cosyntropin), 0.25 mg IM or IV	0, 30, 60 min for cortisol and aldosterone	Normal response is cortisol >21 g/dL and aldosterone response of >4 ng/dL above baseline
	Low-dose ACTH test: ACTH 1-24 (cosyntropin), 1 µg IV	0, 30, 60 min for cortisol	Cortisol should be >21 g/dL
TSH	3-day ACTH stimulation test consists of 0.25 mg ACTH 1-24 given IV over 8 h each day		Cortisol >21 g/dL
	Basal thyroid function tests: T ₄ , T ₃ , TSH	Basal measurements	Low free thyroid hormone levels in the setting of TSH levels that are not appropriately increased indicate pituitary insufficiency
	TRH test: 200–500 µg IV	0, 20, 60 min for TSH and PRL ^a	TSH should increase by >5 mU/L unless thyroid hormone levels are increased
LH, FSH	LH, FSH, testosterone, estrogen	Basal measurements	Basal LH and FSH should be increased in postmenopausal women Low testosterone levels in the setting of low LH and FSH indicate pituitary insufficiency
	GnRH test: GnRH (100 µg) IV	0, 30, 60 min for LH and FSH	In most adults, LH should increase by 10 IU/L and FSH by 2 IU/L Normal responses are variable
Multiple hormones	Combined anterior pituitary test: GHRH (1 g/kg), CRH (1 µg/kg), GnRH (100 g), TRH (200 µg) are given IV	–30, 0, 15, 30, 60, 90, 120 min for GH, ACTH, cortisol, LH, FSH, and TSH	Combined or individual releasing hormone responses must be elevated in the context of basal target gland hormone values and may not be uniformly diagnostic (see text)

^aEvoked PRL response indicates lactotrope integrity.

Abbreviations: T₃, triiodothyronine; T₄, thyroxine; TRH, thyrotropin-releasing hormone. For other abbreviations, see text.

or X-linked. About 10% of children with GH deficiency have mutations in the *GH-N* gene, including gene deletions and a wide range of point mutations. Mutations in transcription factors Pit-1 and Prop-1, which control somatotrope development, result in GH deficiency in combination with other pituitary hormone deficiencies, which may become manifest only in adulthood. The diagnosis of *idiopathic GH deficiency* (IGHD) should be made only after known molecular defects have been rigorously excluded.

GHRH RECEPTOR MUTATIONS Recessive mutations of the GHRH receptor gene in subjects with severe proportionate dwarfism are associated with low basal GH levels that cannot be stimulated by exogenous GHRH, GHRP, or insulin-induced hypoglycemia, as well as anterior pituitary hypoplasia. The syndrome exemplifies the importance of the GHRH receptor for somatotrope cell proliferation and hormonal responsiveness.

GH INSENSITIVITY This is caused by defects of GH receptor structure or signaling. Homozygous or heterozygous mutations of the GH receptor are associated with partial or complete GH insensitivity and growth failure (*Laron's syndrome*). The diagnosis is based on normal or high GH levels, with decreased circulating GH-binding protein (GHBP), and low IGF-I levels. Very rarely, defective IGF-I, IGF-I receptor, or IGF-I signaling defects are also encountered. *STAT5B* mutations result in both immunodeficiency as well as abrogated GH signaling, leading to short stature with normal or elevated GH levels and low IGF-I levels.

Circulating GH receptor antibodies may rarely cause peripheral GH insensitivity.

NUTRITIONAL SHORT STATURE Caloric deprivation and malnutrition, uncontrolled diabetes, and chronic renal failure represent secondary causes of abrogated GH receptor function. These conditions also stimulate production of proinflammatory cytokines, which act to exacerbate the block of GH-mediated signal transduction. Children with these conditions typically exhibit features of acquired short stature with normal or elevated GH and low IGF-I levels.

PSYCHOSOCIAL SHORT STATURE Emotional and social deprivation lead to growth retardation accompanied by delayed speech, discordant hyperphagia, and an attenuated response to administered GH. A nurturing environment restores growth rates.

PRESENTATION AND DIAGNOSIS

Short stature is commonly encountered in clinical practice, and the decision to evaluate these children requires clinical judgment in association with auxologic data and family history. Short stature should be evaluated comprehensively if a patient's height is >3 standard deviations (SD) below the mean for age or if the growth rate has decelerated. Skeletal maturation is best evaluated by measuring a radiologic bone age, which is based mainly on the degree of wrist bone growth plate fusion. Final height can be predicted using standardized scales (Bayley-Pinneau or Tanner-Whitehouse) or

TABLE 402-3 HORMONE REPLACEMENT THERAPY FOR ADULT HYPOPITUITARISM^a

Trophic Hormone Deficit	Hormone Replacement
ACTH	Hydrocortisone (10–20 mg <i>A.M.</i> ; 5–10 mg <i>P.M.</i>) Cortisone acetate (25 mg <i>A.M.</i> ; 12.5 mg <i>P.M.</i>) Prednisone (5 mg <i>A.M.</i>)
TSH	L-Thyroxine (0.075–0.15 mg daily)
FSH/LH	Males Testosterone gel (5–10 g/d) Testosterone skin patch (5 mg/d) Testosterone enanthate (200 mg IM every 2 weeks) Females Conjugated estrogen (0.65–1.25 mg qd for 25 days) Progesterone (5–10 mg qd) on days 16–25 Estradiol skin patch (0.025–0.1 mg every week), adding progesterone on days 16–25 if uterus intact For fertility: menopausal gonadotropins, human chorionic gonadotropins
GH	Adults: Somatotropin (0.1–1.25 mg SC qd) Children: Somatotropin (0.02–0.05 mg/kg per day)
Vasopressin	Intranasal desmopressin (5–20 g twice daily) Oral 300–600 µg qd

^aAll doses shown should be individualized for specific patients and should be reassessed during stress, surgery, or pregnancy. Male and female fertility requirements should be managed as discussed in [Chaps. 411 and 412](#).

Note: For abbreviations, see text.

estimated by adding 6.5 cm (boys) or subtracting 6.5 cm (girls) from the midparental height.

LABORATORY INVESTIGATION

Because GH secretion is pulsatile, GH deficiency is best assessed by examining the response to provocative stimuli, including exercise, insulin-induced hypoglycemia, and other pharmacologic tests that normally increase GH to >7 µg/L in children. Random GH measurements do not distinguish normal children from those with true GH deficiency. Adequate adrenal and thyroid hormone replacement should be assured before testing. Age- and sex-matched IGF-I levels are not sufficiently sensitive or specific to make the diagnosis but can be useful to confirm GH deficiency. Pituitary MRI may reveal pituitary mass lesions or structural defects. Molecular analyses for known mutations should be undertaken when the cause of short stature remains cryptic, or when additional clinical features suggest a genetic cause.

TREATMENT DISORDERS OF GROWTH AND DEVELOPMENT

Replacement therapy with recombinant GH (0.02–0.05 mg/kg per day SC) restores growth velocity in GH-deficient children to ~10 cm/year. If pituitary insufficiency is documented, other associated hormone deficits should be corrected, especially adrenal steroids. GH treatment is also moderately effective for accelerating growth rates in children with Turner's syndrome and chronic renal failure.

In patients with GH insensitivity and growth retardation due to mutations of the GH receptor, treatment with IGF-I bypasses the dysfunctional GH receptor.

ADULT GH DEFICIENCY (AGHD)

This disorder usually is caused by acquired hypothalamic or pituitary somatotrope damage. Acquired pituitary hormone deficiency follows a typical pattern in which loss of adequate GH reserve foreshadows

TABLE 402-4 FEATURES OF ADULT GROWTH HORMONE DEFICIENCY

Clinical
Impaired quality of life Decreased energy and drive Poor concentration Low self-esteem Social isolation
Body composition changes Increased body fat mass Central fat deposition Increased waist-to-hip ratio Decreased lean body mass
Reduced exercise capacity Reduced maximum O ₂ uptake Impaired cardiac function Reduced muscle mass
Cardiovascular risk factors Impaired cardiac structure and function Abnormal lipid profile Decreased fibrinolytic activity Atherosclerosis Omental obesity
Imaging
Pituitary: mass or structural damage Bone: reduced bone mineral density Abdomen: excess omental adiposity
Laboratory
Evoked GH <3 ng/mL IGF-I and IGFBP3 low or normal Increased LDL cholesterol Concomitant gonadotropin, TSH, and/or ACTH reserve deficits may be present

Abbreviation: LDL, low-density lipoprotein. For other abbreviations, see text.

subsequent hormone deficits. The sequential order of hormone loss is usually GH → FSH/LH → TSH → ACTH. Patients previously diagnosed with childhood-onset GH deficiency should be retested as adults to affirm the diagnosis.

PRESENTATION AND DIAGNOSIS

The clinical features of AGHD include changes in body composition, lipid metabolism, and quality of life and cardiovascular dysfunction ([Table 402-4](#)). Body composition changes are common and include reduced lean body mass, increased fat mass with selective deposition of intraabdominal visceral fat, and increased waist-to-hip ratio. Hyperlipidemia, left ventricular dysfunction, hypertension, and increased plasma fibrinogen levels also may be present. Bone mineral content is reduced, with resultant increased fracture rates. Patients may experience social isolation, depression, and difficulty maintaining gainful employment. Adult hypopituitarism is associated with a threefold increase in cardiovascular mortality rates in comparison to age- and sex-matched controls, and this may be due to GH deficiency, as patients in these studies were replaced with other deficient pituitary hormones.

LABORATORY INVESTIGATION

AGHD is rare, and in light of the nonspecific nature of associated clinical symptoms, patients appropriate for testing should be selected carefully on the basis of well-defined criteria. With few exceptions, testing should be restricted to patients with the following predisposing factors: (1) pituitary surgery, (2) pituitary or hypothalamic tumor or granulomas, (3) history of cranial irradiation, (4) radiologic evidence of a pituitary lesion, (5) childhood requirement for GH replacement therapy, and rarely (6) unexplained low age- and sex-matched IGF-I levels. The transition of a GH-deficient adolescent to adulthood

2260 requires retesting to document subsequent adult GH deficiency. Up to 20% of patients previously treated for childhood-onset GH deficiency are found to be GH-sufficient on repeat testing as adults.

A significant proportion (~25%) of truly GH-deficient adults have low-normal IGF-I levels. Thus, as in the evaluation of GH deficiency in children, valid age- and sex-matched IGF-I measurements provide a useful index of therapeutic responses but are not sufficiently sensitive for diagnostic purposes. The most validated test to distinguish pituitary-sufficient patients from those with AGHD is insulin-induced (0.05–0.1 U/kg) hypoglycemia. After glucose reduction to ~40 mg/dL, most individuals experience neuroglycopenic symptoms (Chap. 420), and peak GH release occurs at 60 min and remains elevated for up to 2 h. About 90% of healthy adults exhibit GH responses >5 µg/L; AGHD is defined by a peak GH response to hypoglycemia of <3 µg/L. Although insulin-induced hypoglycemia is safe when performed under appropriate supervision, it is contraindicated in patients with diabetes, ischemic heart disease, cerebrovascular disease, or epilepsy and in elderly patients. Alternative stimulatory tests include intravenous arginine (30 g), GHRH (1 µg/kg), GHRP-6 (90 µg), and glucagon (1 mg). Combinations of these tests may evoke GH secretion in subjects who are not responsive to a single test.

TREATMENT ADULT GH DEFICIENCY

Once the diagnosis of AGHD is unequivocally established, replacement of GH may be indicated. Contraindications to therapy include the presence of an active neoplasm, intracranial hypertension, and uncontrolled diabetes and retinopathy. The starting dose of 0.1–0.2 mg/d should be titrated (up to a maximum of 1.25 mg/d) to maintain IGF-I levels in the mid-normal range for age- and sex-matched controls (Fig. 402-1). Women require higher doses than men, and elderly patients require less GH. Long-term GH maintenance sustains normal IGF-I levels and is associated with persistent body composition changes (e.g., enhanced lean body mass and lower body fat). High-density lipoprotein cholesterol increases, but total cholesterol and insulin levels may not change significantly. Lumbar spine bone mineral density increases, but this response is gradual (>1 year). Many patients note significant improvement in quality of life when evaluated by standardized questionnaires. The effect of GH replacement on mortality rates in GH-deficient patients is currently the subject of long-term prospective investigation.

About 30% of patients exhibit reversible dose-related fluid retention, joint pain, and carpal tunnel syndrome, and up to 40% exhibit myalgias and paresthesia. Patients receiving insulin require careful monitoring for dosing adjustments, as GH is a potent counterregulatory hormone for insulin action. Patients with type 2 diabetes mellitus initially develop further insulin resistance. However, glycemic control usually improves with the sustained loss of abdominal fat associated with long-term GH replacement. Headache, increased intracranial pressure, hypertension, and tinnitus occur rarely. Pituitary tumor regrowth and progression of skin lesions or other tumors are being assessed in long-term surveillance programs. To date, development of these potential side effects does not appear significant.

ACTH DEFICIENCY

PRESENTATION AND DIAGNOSIS

Secondary adrenal insufficiency occurs as a result of pituitary ACTH deficiency. It is characterized by fatigue, weakness, anorexia, nausea, vomiting, and, occasionally, hypoglycemia. In contrast to primary adrenal failure, hypocortisolism associated with pituitary failure usually is not accompanied by hyperpigmentation or mineralocorticoid deficiency.

ACTH deficiency is commonly due to glucocorticoid withdrawal after treatment-associated suppression of the hypothalamic-pituitary-adrenal (HPA) axis. Isolated ACTH deficiency may occur after surgical resection of an ACTH-secreting pituitary adenoma that has suppressed the HPA axis; this phenomenon is in fact suggestive of a surgical cure. The mass effects of other pituitary adenomas or sellar lesions may lead to ACTH deficiency, but usually in combination with other pituitary hormone deficiencies. Partial ACTH deficiency may be unmasked in the presence of an acute medical or surgical illness, when clinically significant hypocortisolism reflects diminished ACTH reserve. Rarely, *TP1T* or *POMC* mutations result in primary ACTH deficiency.

LABORATORY DIAGNOSIS

Inappropriately low ACTH levels in the setting of low cortisol levels are characteristic of diminished ACTH reserve. Low basal serum cortisol levels are associated with blunted cortisol responses to ACTH stimulation and impaired cortisol response to insulin-induced hypoglycemia, or testing with metyrapone or CRH. For a description of provocative ACTH tests, see Chap. 406.

TREATMENT ACTH DEFICIENCY

Glucocorticoid replacement therapy improves most features of ACTH deficiency. The total daily dose of hydrocortisone replacement preferably should not exceed 25 mg daily, divided into two or three doses. Prednisone (5 mg each morning) is longer acting and has fewer mineralocorticoid effects than hydrocortisone. Some authorities advocate lower maintenance doses in an effort to avoid cushingoid side effects. Doses should be increased severalfold during periods of acute illness or stress.

GONADOTROPIN DEFICIENCY

Hypogonadism is the most common presenting feature of adult hypopituitarism even when other pituitary hormones are also deficient. It is often a harbinger of hypothalamic or pituitary lesions that impair GnRH production or delivery through the pituitary stalk. As noted below, hypogonadotropic hypogonadism is a common presenting feature of hyperprolactinemia.

A variety of inherited and acquired disorders are associated with isolated hypogonadotropic hypogonadism (IHH) (Chap. 411). Hypothalamic defects associated with GnRH deficiency include Kallmann syndrome and mutations in more than a dozen genes that regulate GnRH neuron migration, development, and function (see above). Mutations in *GPR54*, *DAX1*, kisspeptin, the GnRH receptor, and the *LHβ* or *FSHβ* subunit genes also cause pituitary gonadotropin

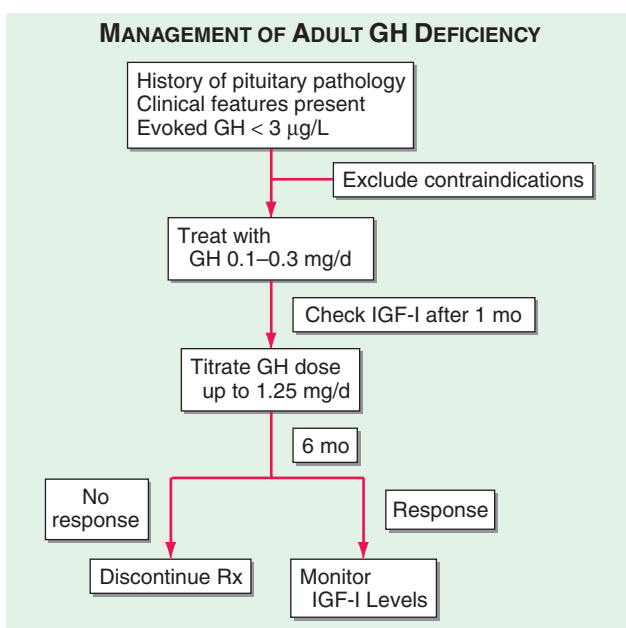


FIGURE 402-1 Management of adult growth hormone (GH) deficiency. IGF, insulin-like growth factor; Rx, Treatment.

deficiency. Acquired forms of GnRH deficiency leading to hypogonadotropism are seen in association with anorexia nervosa, stress, starvation, and extreme exercise but also may be idiopathic. Hypogonadotropic hypogonadism in these disorders is reversed by removal of the stressful stimulus or by caloric replenishment.

PRESENTATION AND DIAGNOSIS

In premenopausal women, hypogonadotropic hypogonadism presents as diminished ovarian function leading to oligomenorrhea or amenorrhea, infertility, decreased vaginal secretions, decreased libido, and breast atrophy. In hypogonadal adult men, secondary testicular failure is associated with decreased libido and potency, infertility, decreased muscle mass with weakness, reduced beard and body hair growth, soft testes, and characteristic fine facial wrinkles. Osteoporosis occurs in both untreated hypogonadal women and men.

LABORATORY INVESTIGATION

Central hypogonadism is associated with low or inappropriately normal serum gonadotropin levels in the setting of low sex hormone concentrations (testosterone in men, estradiol in women). Because gonadotropin secretion is pulsatile, valid assessments may require repeated measurements or the use of pooled serum samples. Men have reduced sperm counts.

Intravenous GnRH (100 µg) stimulates gonadotropes to secrete LH (which peaks within 30 min) and FSH (which plateaus during the ensuing 60 min). Normal responses vary according to menstrual cycle stage, age, and sex of the patient. Generally, LH levels increase about threefold, whereas FSH responses are less pronounced. In the setting of gonadotropin deficiency, a normal gonadotropin response to GnRH indicates intact pituitary gonadotrope function and suggests a hypothalamic abnormality. An absent response, however, does not reliably distinguish pituitary from hypothalamic causes of hypogonadism. For this reason, GnRH testing usually adds little to the information gained from baseline evaluation of the hypothalamic-pituitary-gonadotrope axis except in cases of isolated GnRH deficiency (e.g., Kallmann syndrome).

MRI examination of the sellar region and assessment of other pituitary functions usually are indicated in patients with documented central hypogonadism.

TREATMENT GONADOTROPIN DEFICIENCY

In males, testosterone replacement is necessary to achieve and maintain normal growth and development of the external genitalia, secondary sex characteristics, male sexual behavior, and androgenic anabolic effects, including maintenance of muscle function and bone mass. Testosterone may be administered by intramuscular injections every 1–4 weeks or by using skin patches that are replaced daily (Chap. 411). Testosterone gels are also available. Gonadotropin injections (hCG or human menopausal gonadotropin [hMG]) over 12–18 months are used to restore fertility. Pulsatile GnRH therapy (25–150 ng/kg every 2 h), administered by a subcutaneous infusion pump, is also effective for treatment of hypothalamic hypogonadism when fertility is desired.

In premenopausal women, cyclical replacement of estrogen and progesterone maintains secondary sexual characteristics and integrity of genitourinary tract mucosa and prevents premature osteoporosis (Chap. 412). Gonadotropin therapy is used for ovulation induction. Follicular growth and maturation are initiated using hMG or recombinant FSH; hCG or human luteinizing hormone (hLH) is subsequently injected to induce ovulation. As in men, pulsatile GnRH therapy can be used to treat hypothalamic causes of gonadotropin deficiency.

DIABETES INSIPIDUS

See Chap. 404 for diagnosis and treatment of diabetes insipidus.

403 Anterior Pituitary Tumor Syndromes

Shlomo Melmed, J. Larry Jameson

HYPOTHALAMIC, PITUITARY, AND OTHER SELLAR MASSES

EVALUATION OF SELLAR MASSES

Local Mass Effects Clinical manifestations of sellar lesions vary, depending on the anatomic location of the mass and the direction of its extension (Table 403-1). The dorsal sellar diaphragm presents the least resistance to soft tissue expansion from the sella; consequently, pituitary adenomas frequently extend in a suprasellar direction. Bony invasion may occur as well.

Headaches are common features of small intrasellar tumors, even with no demonstrable suprasellar extension. Because of the confined nature of the pituitary, small changes in intrasellar pressure stretch the dural plate; however, headache severity correlates poorly with adenoma size or extension.

Suprasellar extension can lead to visual loss by several mechanisms, the most common being compression of the optic chiasm, but rarely, direct invasion of the optic nerves or obstruction of cerebrospinal fluid (CSF) flow leading to secondary visual disturbances can occur. Pituitary stalk compression by a hormonally active or inactive intrasellar mass may compress the portal vessels, disrupting pituitary access to hypothalamic hormones and dopamine; this results in early hyperprolactinemia and later concurrent loss of other pituitary hormones. This “stalk section” phenomenon may also be caused by trauma, whiplash injury with posterior clinoid stalk compression, or skull base fractures. Lateral mass invasion may impinge on the cavernous sinus and compress its neural contents, leading to cranial nerve III,

TABLE 403-1 FEATURES OF SELLAR MASS LESIONS^a

Impacted Structure	Clinical Impact
Pituitary	Hypogonadism Hypothyroidism Growth failure and adult hyposomatotropism Hypoadrenalism
Optic chiasm	Loss of red perception Bitemporal hemianopia Superior or bitemporal field defect Scotoma Blindness
Hypothalamus	Temperature dysregulation Appetite and thirst disorders Obesity Diabetes insipidus Sleep disorders Behavioral dysfunction Autonomic dysfunction
Cavernous sinus	Ophthalmoplegia with or without ptosis or diplopia Facial numbness
Frontal lobe	Personality disorder Anosmia
Brain	Headache Hydrocephalus Psychosis Dementia Laughing seizures

^aAs the intrasellar mass expands, it first compresses intrasellar pituitary tissue, then usually invades dorsally through the dura to lift the optic chiasm or laterally to the cavernous sinuses. Bony erosion is rare, as is direct brain compression. Microadenomas may present with headache.

2262 IV, and VI palsies as well as effects on the ophthalmic and maxillary branches of the fifth cranial nerve (Chap. 455). Patients may present with diplopia, ptosis, ophthalmoplegia, and decreased facial sensation, depending on the extent of neural damage. Extension into the sphenoid sinus indicates that the pituitary mass has eroded through the sellar floor. Aggressive tumors rarely invade the palate roof and cause nasopharyngeal obstruction, infection, and CSF leakage. Temporal and frontal lobe involvement may rarely lead to uncinete seizures, personality disorders, and anosmia. Direct hypothalamic encroachment by an invasive pituitary mass may cause important metabolic sequelae, including precocious puberty or hypogonadism, diabetes insipidus, sleep disturbances, dysthermia, and appetite disorders.

Magnetic Resonance Imaging Sagittal and coronal T1-weighted magnetic resonance imaging (MRI) before and after administration of gadolinium allows precise visualization of the pituitary gland with clear delineation of the hypothalamus, pituitary stalk, pituitary tissue and surrounding suprasellar cisterns, cavernous sinuses, sphenoid sinus, and optic chiasm. Pituitary gland height ranges from 6 mm in children to 8 mm in adults; during pregnancy and puberty, the height may reach 10–12 mm. The upper aspect of the adult pituitary is flat or slightly concave, but in adolescent and pregnant individuals, this surface may be convex, reflecting physiologic pituitary enlargement. The stalk should be midline and vertical. Computed tomography (CT) scan is reserved to define the extent of bony erosion or the presence of calcification.

Anterior pituitary gland soft tissue consistency is slightly heterogeneous on MRI, and signal intensity resembles that of brain matter on T1-weighted imaging (Fig. 403-1). Adenoma density is usually lower than that of surrounding normal tissue on T1-weighted imaging, and the signal intensity increases with T2-weighted images. The high phospholipid content of the posterior pituitary results in a “pituitary bright spot.”

Sellar masses are encountered commonly as incidental findings on MRI, and most of them are pituitary adenomas (incidentalomas). In the absence of hormone hypersecretion, these small intrasellar lesions can be monitored safely with MRI, which is performed annually and then less often if there is no evidence of further growth. Resection should be considered for incidentally discovered larger macroadenomas, because about one-third become invasive or cause local pressure effects. If hormone hypersecretion is evident, specific therapies are indicated as described below. When larger masses (>1 cm) are encountered, they should also be distinguished from nonadenomatous lesions. Meningiomas often are associated with bony hyperostosis;



FIGURE 403-1 Pituitary adenoma. Coronal T1-weighted postcontrast magnetic resonance image shows a homogeneously enhancing mass (arrowheads) in the sella turcica and suprasellar region compatible with a pituitary adenoma; the small arrows outline the carotid arteries.

TABLE 403-2 SCREENING TESTS FOR FUNCTIONAL PITUITARY ADENOMAS

	Test	Comments
Acromegaly	Serum IGF-I	Interpret IGF-I relative to age- and sex-matched controls
	Oral glucose tolerance test with GH obtained at 0, 30, and 60 min	Normal subjects should suppress growth hormone to <1 g/L
Prolactinoma	Serum PRL	Exclude medications MRI of the sella should be ordered if PRL is elevated
Cushing's disease	24-h urinary free cortisol	Ensure urine collection is total and accurate
	Dexamethasone (1 mg) at 11 P.M. and fasting plasma cortisol measured at 8 A.M.	Normal subjects suppress to <5 g/dL
	ACTH assay	Distinguishes adrenal adenoma (ACTH suppressed) from ectopic ACTH or Cushing's disease (ACTH normal or elevated)

Abbreviations: ACTH, adrenocorticotropic hormone; GH, growth hormone; IGF-I, insulin-like growth factor I; MRI, magnetic resonance imaging; PRL, prolactin.

craniopharyngiomas may be calcified and are usually hypodense, whereas gliomas are hyperdense on T2-weighted images.

Ophthalmologic Evaluation Because optic tracts may be contiguous to an expanding pituitary mass, reproducible visual field assessment using perimetry techniques should be performed on all patients with sellar mass lesions that impinge the optic chiasm (Chap. 39). Bitemporal hemianopia, often more pronounced superiorly, is observed classically. It occurs because nasal ganglion cell fibers, which cross in the optic chiasm, are especially vulnerable to compression of the ventral optic chiasm. Occasionally, homonymous hemianopia occurs from postchiasmal compression or monocular temporal field loss from prechiasmal compression. Invasion of the cavernous sinus can produce diplopia from ocular motor nerve palsy. Early diagnosis reduces the risk of optic atrophy, vision loss, or eye misalignment.

Laboratory Investigation The presenting clinical features of functional pituitary adenomas (e.g., acromegaly, prolactinomas, or Cushing's syndrome) should guide the laboratory studies (Table 403-2). However, for a sellar mass with no obvious clinical features of hormone excess, laboratory studies are geared toward determining the nature of the tumor and assessing the possible presence of hypopituitarism. When a pituitary adenoma is suspected based on MRI, initial hormonal evaluation usually includes (1) basal prolactin (PRL); (2) insulin-like growth factor (IGF) I; (3) 24-h urinary free cortisol (UFC) and/or overnight oral dexamethasone (1 mg) suppression test; (4) α subunit, follicle-stimulating hormone (FSH), and luteinizing hormone (LH); and (5) thyroid function tests. Additional hormonal evaluation may be indicated based on the results of these tests. Pending more detailed assessment of hypopituitarism, a menstrual history, measurement of testosterone and 8 A.M. cortisol levels, and thyroid function tests usually identify patients with pituitary hormone deficiencies that require hormone replacement before further testing or surgery.

Histologic Evaluation Immunohistochemical staining of pituitary tumor specimens obtained at transsphenoidal surgery confirms clinical and laboratory studies and provides a histologic diagnosis when hormone studies are equivocal and in cases of clinically nonfunctioning tumors. Occasionally, ultrastructural assessment by electron microscopy is required for diagnosis.

TREATMENT HYPOTHALAMIC, PITUITARY, AND OTHER SELLAR MASSES

OVERVIEW Successful management of sellar masses requires accurate diagnosis as well as selection of optimal therapeutic modalities. Most pituitary tumors are benign and slow-growing. Clinical features

result from local mass effects and hormonal hyper- or hyposecretion syndromes caused directly by the adenoma or occurring as a consequence of treatment. Thus, lifelong management and follow-up are necessary for these patients.

MRI with gadolinium enhancement for pituitary visualization, new advances in transsphenoidal surgery and in stereotactic radiotherapy (including gamma-knife radiotherapy), and novel therapeutic agents have improved pituitary tumor management. The goals of pituitary tumor treatment include normalization of excess pituitary secretion, amelioration of symptoms and signs of hormonal hypersecretion syndromes, and shrinkage or ablation of large tumor masses with relief of adjacent structure compression. Residual anterior pituitary function should be preserved during treatment and sometimes can be restored by removing the tumor mass. Ideally, adenoma recurrence should be prevented.

TRANSSPHEOIDAL SURGERY Transsphenoidal rather than transfrontal resection is the desired surgical approach for pituitary tumors, except for the rare invasive suprasellar mass surrounding the frontal or middle fossa or the optic nerves or invading posteriorly behind the clivus. Intraoperative microscopy facilitates visual distinction between adenomatous and normal pituitary tissue as well as microdissection of small tumors that may not be visible by MRI (Fig. 403-2). Transsphenoidal surgery also avoids the cranial invasion and manipulation of brain tissue required by subfrontal surgical approaches. Endoscopic techniques with three-dimensional

intraoperative localization have also improved visualization and access to tumor tissue. Individual surgical experience is a major determinant of outcome efficacy with these techniques.

In addition to correction of hormonal hypersecretion, pituitary surgery is indicated for mass lesions that impinge on surrounding structures. Surgical decompression and resection are required for an expanding pituitary mass accompanied by persistent headache, progressive visual field defects, cranial nerve palsies, hydrocephalus, and, occasionally, intrapituitary hemorrhage and apoplexy. Transsphenoidal surgery sometimes is used for pituitary tissue biopsy to establish a histologic diagnosis. Whenever possible, the pituitary mass lesion should be selectively excised; normal pituitary tissue should be manipulated or resected only when critical for effective mass dissection. Nonselective hemihypophysectomy or total hypophysectomy may be indicated if no hypersecreting mass lesion is clearly discernible, multifocal lesions are present, or the remaining nontumorous pituitary tissue is obviously necrotic. This strategy, however, increases the likelihood of hypopituitarism and the need for lifelong hormone replacement.

Preoperative mass effects, including visual field defects and compromised pituitary function, may be reversed by surgery, particularly when the deficits are not long-standing. For large and invasive tumors, it is necessary to determine the optimal balance between maximal tumor resection and preservation of anterior pituitary function, especially for preserving growth and reproductive function in younger patients. Similarly, tumor invasion outside the sella is rarely amenable to surgical cure; the surgeon must judge the risk-versus-benefit ratio of extensive tumor resection.

Side Effects Tumor size, the degree of invasiveness, and experience of the surgeon largely determine the incidence of surgical complications. Operative mortality rate is about 1%. Transient diabetes insipidus and hypopituitarism occur in up to 20% of patients. Permanent diabetes insipidus, cranial nerve damage, nasal septal perforation, or visual disturbances may be encountered in up to 10% of patients. CSF leaks occur in 4% of patients. Less common complications include carotid artery injury, loss of vision, hypothalamic damage, and meningitis. Permanent side effects are rare after surgery for microadenomas.

RADIATION

Radiation is used either as a primary therapy for pituitary or parasellar masses or, more commonly, as an adjunct to surgery or medical therapy. Focused megavoltage irradiation is achieved by precise MRI localization, using a high-voltage linear accelerator and accurate isocentric rotational arcing. A major determinant of accurate irradiation is reproduction of the patient's head position during multiple visits and maintenance of absolute head immobility. A total of <50 Gy (5000 rad) is given as 180-cGy (180-rad) fractions divided over about 6 weeks. Stereotactic radiosurgery delivers a large single high-energy dose from a cobalt-60 source (gamma knife), linear accelerator, or cyclotron. Long-term effects of gamma-knife surgery are unclear but appear to be similar to those encountered with conventional radiation. Proton beam therapy is available in some centers and provides concentrated radiation doses within a localized region.

The role of radiation therapy in pituitary tumor management depends on multiple factors, including the nature of the tumor, the age of the patient, and the availability of surgical and radiation expertise. Because of its relatively slow onset of action, radiation therapy is usually reserved for postsurgical management. As an adjuvant to surgery, radiation is used to treat residual tumor and in an attempt to prevent regrowth. Irradiation offers the only means for potentially ablating significant postoperative residual nonfunctioning tumor tissue. In contrast, PRL- and growth hormone (GH)-secreting tumor tissues are amenable to medical therapy.

Side Effects In the short term, radiation may cause transient nausea and weakness. Alopecia and loss of taste and smell may be more long-lasting. Failure of pituitary hormone synthesis is common in patients who have undergone head and neck or pituitary-directed irradiation. More than 50% of patients develop loss of GH, adrenocorticotropin

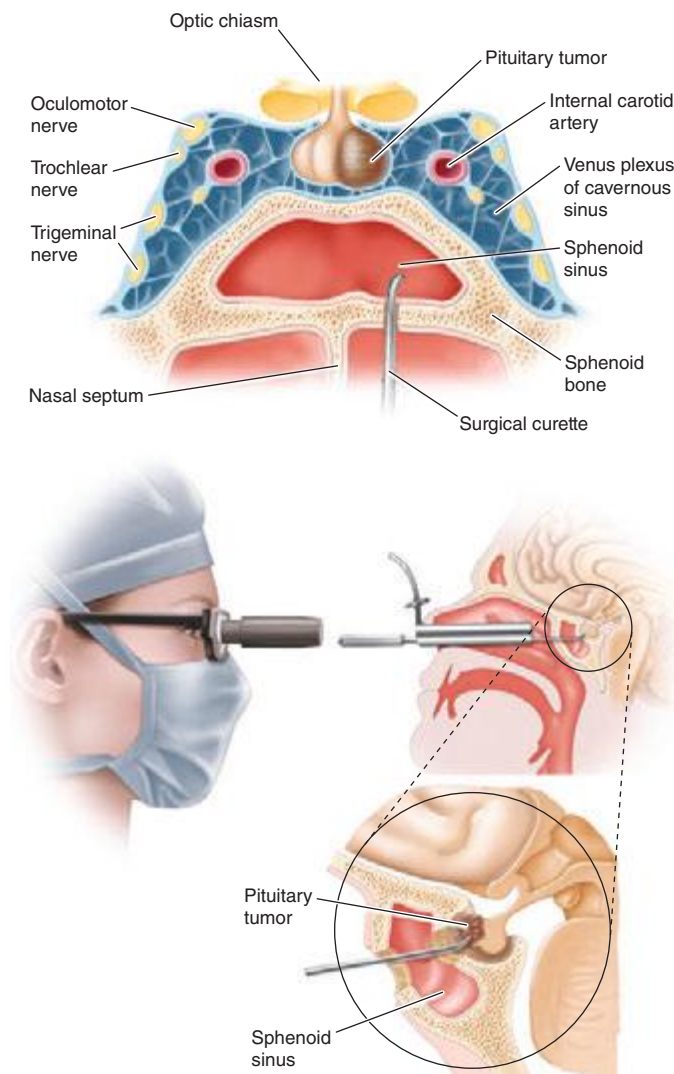


FIGURE 403-2 Transsphenoidal resection of pituitary mass via the endonasal approach. (Adapted from R Fahlbusch: *Endocrinol Metab Clin* 21:669, 1992.)

2264 hormone (ACTH), thyroid-stimulating hormone (TSH), and/or gonadotropin secretion within 10 years, usually due to hypothalamic damage. Lifelong follow-up with testing of anterior pituitary hormone reserve is therefore required after radiation treatment. Optic nerve damage with impaired vision due to optic neuritis is reported in about 2% of patients who undergo pituitary irradiation. Cranial nerve damage is uncommon now that radiation doses are ≤ 2 Gy (200 rad) at any one treatment session and the maximum dose is < 50 Gy (5000 rad). The use of stereotactic radiotherapy may reduce damage to adjacent structures. Radiotherapy for pituitary tumors has been associated with adverse mortality rates, mainly from cerebrovascular disease. The cumulative risk of developing a secondary tumor after conventional radiation is 1.3% after 10 years and 1.9% after 20 years.

MEDICAL

Medical therapy for pituitary tumors is highly specific and depends on tumor type. For prolactinomas, dopamine agonists are the treatment of choice. For acromegaly, somatostatin analogues and GH receptor antagonists are indicated. For TSH-secreting tumors, somatostatin analogues and occasionally dopamine agonists are indicated. ACTH-secreting tumors and nonfunctioning tumors are generally not responsive to medications and require surgery and/or irradiation.

SELLAR MASSES

Sellar masses other than pituitary adenomas may arise from brain, hypothalamic, or pituitary tissues. Each exhibit features related to the lesion location but also unique to the specific etiology.

Hypothalamic Lesions Lesions involving the anterior and preoptic hypothalamic regions cause paradoxical vasoconstriction, tachycardia, and hyperthermia. Acute hyperthermia usually is due to a hemorrhagic insult, but poikilothermia may also occur. Central disorders of thermoregulation result from posterior hypothalamic damage. The *periodic hypothermia syndrome* is characterized by episodic attacks of rectal temperatures $< 30^{\circ}\text{C}$ (86°F), sweating, vasodilation, vomiting, and bradycardia (**Chap. 478e**). Damage to the ventromedial hypothalamic nuclei by craniopharyngiomas, hypothalamic trauma, or inflammatory disorders may be associated with *hyperphagia* and *obesity*. This region appears to contain an energy-satiety center where melanocortin receptors are influenced by leptin, insulin, pro-opiomelanocortin (POMC) products, and gastrointestinal peptides (**Chap. 415e**). Polydipsia and hypodipsia are associated with damage to central osmoreceptors located in preoptic nuclei (**Chap. 404**). Slow-growing hypothalamic lesions can cause increased somnolence and disturbed sleep cycles as well as obesity, hyperthermia, and emotional outbursts. Lesions of the central hypothalamus may stimulate sympathetic neurons, leading to elevated serum catecholamine and cortisol levels. These patients are predisposed to cardiac arrhythmias, hypertension, and gastric erosions.

Craniopharyngiomas are benign, suprasellar cystic masses that present with headaches, visual field deficits, and variable degrees of hypopituitarism. They are derived from Rathke's pouch and arise near the pituitary stalk, commonly extending into the suprasellar cistern. Craniopharyngiomas are often large, cystic, and locally invasive. Many are partially calcified, exhibiting a characteristic appearance on skull x-ray and CT images. More than half of all patients present before age 20, usually with signs of increased intracranial pressure, including headache, vomiting, papilledema, and hydrocephalus. Associated symptoms include visual field abnormalities, personality changes and cognitive deterioration, cranial nerve damage, sleep difficulties, and weight gain. Hypopituitarism can be documented in about 90%, and diabetes insipidus occurs in about 10% of patients. About half of affected children present with growth retardation. MRI is generally superior to CT for evaluating cystic structure and tissue components of craniopharyngiomas. CT is useful to define calcifications and evaluate invasion into surrounding bony structures and sinuses.

Treatment usually involves transcranial or transsphenoidal surgical resection followed by postoperative radiation of residual tumor.

Surgery alone is curative in less than half of patients because of recurrences due to adherence to vital structures or because of small tumor deposits in the hypothalamus or brain parenchyma. The goal of surgery is to remove as much tumor as possible without risking complications associated with efforts to remove firmly adherent or inaccessible tissue. In the absence of radiotherapy, about 75% of craniopharyngiomas recur, and 10-year survival is less than 50%. In patients with incomplete resection, radiotherapy improves 10-year survival to 70–90% but is associated with increased risk of secondary malignancies. Most patients require lifelong pituitary hormone replacement.

Developmental failure of Rathke's pouch obliteration may lead to *Rathke's cysts*, which are small (< 5 mm) cysts entrapped by squamous epithelium and are found in about 20% of individuals at autopsy. Although Rathke's cleft cysts do not usually grow and are often diagnosed incidentally, about a third present in adulthood with compressive symptoms, diabetes insipidus, and hyperprolactinemia due to stalk compression. Rarely, hydrocephalus develops. The diagnosis is suggested preoperatively by visualizing the cyst wall on MRI, which distinguishes these lesions from craniopharyngiomas. Cyst contents range from CSF-like fluid to mucoid material. *Arachnoid cysts* are rare and generate an MRI image that is isointense with CSF.

Sella chordomas usually present with bony clival erosion, local invasiveness, and, on occasion, calcification. Normal pituitary tissue may be visible on MRI, distinguishing chordomas from aggressive pituitary adenomas. Mucinous material may be obtained by fine-needle aspiration.

Meningiomas arising in the sellar region may be difficult to distinguish from nonfunctioning pituitary adenomas. Meningiomas typically enhance on MRI and may show evidence of calcification or bony erosion. Meningiomas may cause compressive symptoms.

Histiocytosis X includes a variety of syndromes associated with foci of eosinophilic granulomas. Diabetes insipidus, exophthalmos, and punched-out lytic bone lesions (*Hand-Schüller-Christian disease*) are associated with granulomatous lesions visible on MRI, as well as a characteristic axillary skin rash. Rarely, the pituitary stalk may be involved.

Pituitary metastases occur in $\sim 3\%$ of cancer patients. Bloodborne metastatic deposits are found almost exclusively in the posterior pituitary. Accordingly, diabetes insipidus can be a presenting feature of lung, gastrointestinal, breast, and other pituitary metastases. About half of pituitary metastases originate from breast cancer; about 25% of patients with metastatic breast cancer have such deposits. Rarely, pituitary stalk involvement results in anterior pituitary insufficiency. The MRI diagnosis of a metastatic lesion may be difficult to distinguish from an aggressive pituitary adenoma; the diagnosis may require histologic examination of excised tumor tissue. Primary or metastatic lymphoma, leukemias, and plasmacytomas also occur within the sella.

Hypothalamic hamartomas and *gangliocytomas* may arise from astrocytes, oligodendrocytes, and neurons with varying degrees of differentiation. These tumors may overexpress hypothalamic neuropeptides, including gonadotropin-releasing hormone (GnRH), growth hormone-releasing hormone (GHRH), and corticotropin-releasing hormone (CRH). With GnRH-producing tumors, children present with precocious puberty, psychomotor delay, and laughing-associated seizures. Medical treatment of GnRH-producing hamartomas with long-acting GnRH analogues effectively suppresses gonadotropin secretion and controls premature pubertal development. Rarely, hamartomas also are associated with craniofacial abnormalities; imperforate anus; cardiac, renal, and lung disorders; and pituitary failure as features of *Pallister-Hall syndrome*, which is caused by mutations in the carboxy terminus of the *GLI3* gene. Hypothalamic hamartomas are often contiguous with the pituitary, and preoperative MRI diagnosis may not be possible. Histologic evidence of hypothalamic neurons in tissue resected at transsphenoidal surgery may be the first indication of a primary hypothalamic lesion.

Hypothalamic gliomas and *optic gliomas* occur mainly in childhood and usually present with visual loss. Adults have more aggressive tumors; about a third are associated with neurofibromatosis.

Brain germ cell tumors may arise within the sellar region. They include *dysgerminomas*, which frequently are associated with diabetes insipidus and visual loss. They rarely metastasize. *Germinomas*,

embryonal carcinomas, teratomas, and choriocarcinomas may arise in the parasellar region and produce hCG. These germ cell tumors present with precocious puberty, diabetes insipidus, visual field defects, and thirst disorders. Many patients are GH-deficient with short stature.

PITUITARY ADENOMAS AND HYPERSECRETION SYNDROMES

Pituitary adenomas are the most common cause of pituitary hormone hypersecretion and hyposecretion syndromes in adults. They account for ~15% of all intracranial neoplasms and have been identified with a population prevalence of ~80/100,000. At autopsy, up to one-quarter of all pituitary glands harbor an unsuspected microadenoma (<10 mm diameter). Similarly, pituitary imaging detects small clinically inapparent pituitary lesions in at least 10% of individuals.

Pathogenesis Pituitary adenomas are benign neoplasms that arise from one of the five anterior pituitary cell types. The clinical and biochemical phenotypes of pituitary adenomas depend on the cell type from which they are derived. Thus, tumors arising from lactotrope (PRL), somatotrope (GH), corticotrope (ACTH), thyrotrope (TSH), or gonadotrope (LH, FSH) cells hypersecrete their respective hormones (Table 403-3). Plurihormonal tumors express various combinations of GH, PRL, TSH, ACTH, or the glycoprotein hormone α or β subunits. They may be diagnosed by careful immunocytochemistry or may manifest as clinical syndromes that combine features of these hormonal hypersecretory syndromes. Morphologically, these tumors may arise from a single polysecreting cell type or include cells with mixed function within the same tumor.

Hormonally active tumors are characterized by autonomous hormone secretion with diminished feedback responsiveness to physiologic inhibitory pathways. Hormone production does not always correlate with tumor size. Small hormone-secreting adenomas may cause significant clinical perturbations, whereas larger adenomas that produce less hormone may be clinically silent and remain undiagnosed (if no central compressive effects occur). About one-third of all adenomas are clinically nonfunctioning and produce no distinct clinical hypersecretory syndrome. Most of them arise from gonadotrope cells and may secrete small amounts of α - and β -glycoprotein hormone subunits or, very rarely, intact circulating gonadotropins. True pituitary carcinomas with documented extracranial metastases are exceedingly rare.

Almost all pituitary adenomas are monoclonal in origin, implying the acquisition of one or more somatic mutations that confer a selective growth advantage. Consistent with their clonal origin, complete surgical resection of small pituitary adenomas usually cures hormone

hypersecretion. Nevertheless, hypothalamic hormones such as GHRH and CRH also enhance mitotic activity of their respective pituitary target cells in addition to their role in pituitary hormone regulation. Thus, patients who harbor rare abdominal or chest tumors that elaborate ectopic GHRH or CRH may present with somatotrope or corticotrope hyperplasia with GH or ACTH hypersecretion.

Several etiologic genetic events have been implicated in the development of pituitary tumors. The pathogenesis of sporadic forms of acromegaly has been particularly informative as a model of tumorigenesis. GHRH, after binding to its G protein-coupled somatotrope receptor, uses cyclic adenosine monophosphate (AMP) as a second messenger to stimulate GH secretion and somatotrope proliferation. A subset (~35%) of GH-secreting pituitary tumors contains sporadic mutations in Gsa (Arg 201 \rightarrow Cys or His; Gln 227 \rightarrow Arg). These mutations attenuate intrinsic GTPase activity, resulting in constitutive elevation of cyclic AMP, Pit-1 induction, and activation of cyclic AMP response element binding protein (CREB), thereby promoting somatotrope cell proliferation and GH secretion.

Characteristic loss of heterozygosity (LOH) in various chromosomes has been documented in large or invasive macroadenomas, suggesting the presence of putative tumor suppressor genes at these loci in up to 20% of sporadic pituitary tumors, including GH-, PRL-, and ACTH-producing adenomas and some nonfunctioning tumors. Lineage-specific cell cycle disruptions with elevated levels of CDK inhibitors are present in most of these adenomas.

Compelling evidence also favors growth factor promotion of pituitary tumor proliferation. Basic fibroblast growth factor (bFGF) is abundant in the pituitary and stimulates pituitary cell mitogenesis, whereas epithelial growth factor (EGF) receptor signaling induces both hormone synthesis and cell proliferation. Other factors involved in initiation and promotion of pituitary tumors include loss of negative-feedback inhibition (as seen with primary hypothyroidism or hypogonadism) and estrogen-mediated or paracrine angiogenesis. Growth characteristics and neoplastic behavior also may be influenced by several activated oncogenes, including RAS and pituitary tumor transforming gene (PTTG), or inactivation of growth suppressor genes, including MEG3.

Genetic Syndromes Associated with Pituitary Tumors Several familial syndromes are associated with pituitary tumors, and the genetic mechanisms for some of them have been unraveled (Table 403-4).

Multiple endocrine neoplasia (MEN) 1 is an autosomal dominant syndrome characterized primarily by a genetic predisposition to parathyroid, pancreatic islet, and pituitary adenomas (Chap. 408). MEN1 is caused by inactivating germline mutations in *MEN1*, a constitutively expressed tumor-suppressor gene located on chromosome 11q13.

TABLE 403-3 CLASSIFICATION OF PITUITARY ADENOMAS*

Adenoma Cell Origin	Hormone Product	Clinical Syndrome
Lactotrope	PRL	Hypogonadism, galactorrhea
Gonadotrope	FSH, LH, subunits	Silent or hypogonadism
Somatotrope	GH	Acromegaly/gigantism
Corticotrope	ACTH	Cushing's disease
Mixed growth hormone and prolactin cell	GH, PRL	Acromegaly, hypogonadism, galactorrhea
Other plurihormonal cell	Any	Mixed
Acidophil stem cell	PRL, GH	Hypogonadism, galactorrhea, acromegaly
Mammomatotrope	PRL, GH	Hypogonadism, galactorrhea, acromegaly
Thyrotrope	TSH	Thyrotoxicosis
Null cell	None	Pituitary failure
Oncocytoma	None	Pituitary failure

*Hormone-secreting tumors are listed in decreasing order of frequency. All tumors may cause local pressure effects, including visual disturbances, cranial nerve palsy, and headache.

Note: For abbreviations, see text.

Source: Adapted from S Melmed, in JL Jameson (ed): *Principles of Molecular Medicine*. Totowa, NJ, Humana Press, 1998.

TABLE 403-4 FAMILIAL PITUITARY TUMOR SYNDROMES

	Gene Mutated	Clinical Features
Multiple endocrine neoplasia 1 (MEN 1)	<i>MEN1</i> (11q13)	Hyperparathyroidism Pancreatic neuroendocrine tumors Foregut carcinoids Adrenal adenomas Skin lesions Pituitary adenomas (40%)
Multiple endocrine neoplasia 4 (MEN 4)	<i>CDKN1B</i> (12p13)	Hyperparathyroidism Pituitary adenomas Other tumors
Carney complex	<i>PRKAR1A</i> (17q23-24)	Pituitary hyperplasia and adenomas (10%) Atrial myxomas Schwannomas Adrenal hyperplasia Lentiginosities
Familial pituitary adenomas	<i>AIP</i> (11q13.3)	Acromegaly/gigantism (~15% of afflicted families)

2266 Loss of heterozygosity or a somatic mutation of the remaining normal *MENIN* allele leads to tumorigenesis. About half of affected patients develop prolactinomas; acromegaly and Cushing's syndrome are less commonly encountered.

Carney's syndrome is characterized by spotty skin pigmentation, myxomas, and endocrine tumors, including testicular, adrenal, and pituitary adenomas. Acromegaly occurs in about 20% of these patients. A subset of patients have mutations in the R1 α regulatory subunit of protein kinase A (*PRKARIA*).

McCune-Albright syndrome consists of polyostotic fibrous dysplasia, pigmented skin patches, and a variety of endocrine disorders, including acromegaly, adrenal adenomas, and autonomous ovarian function (**Chap. 426e**). Hormonal hypersecretion results from constitutive cyclic AMP production caused by inactivation of the GTPase activity of Gsa. The Gsa mutations occur postzygotically, leading to a mosaic pattern of mutant expression.

Familial acromegaly is a rare disorder in which family members may manifest either acromegaly or gigantism. A subset of families with a predisposition for familial pituitary tumors, especially acromegaly, have been found to harbor germline mutations in the *AIP* gene, which encodes the aryl hydrocarbon receptor interacting protein.

HYPERPROLACTINEMIA

Etiology Hyperprolactinemia is the most common pituitary hormone hypersecretion syndrome in both men and women. PRL-secreting pituitary adenomas (prolactinomas) are the most common cause of PRL levels >200 $\mu\text{g/L}$ (see below). Less pronounced PRL elevation can also be seen with microprolactinomas but is more commonly caused by drugs, pituitary stalk compression, hypothyroidism, or renal failure (**Table 403-5**).

Pregnancy and lactation are the important physiologic causes of hyperprolactinemia. Sleep-associated hyperprolactinemia reverts to normal within an hour of awakening. Nipple stimulation and sexual orgasm also may increase PRL. Chest wall stimulation or trauma (including chest surgery and herpes zoster) invoke the reflex suckling arc with resultant hyperprolactinemia. Chronic renal failure elevates PRL by decreasing peripheral clearance. Primary hypothyroidism is associated with mild hyperprolactinemia, probably because of compensatory TRH secretion.

Lesions of the hypothalamic-pituitary region that disrupt hypothalamic dopamine synthesis, portal vessel delivery, or lactotrope responses are associated with hyperprolactinemia. Thus, hypothalamic tumors, cysts, infiltrative disorders, and radiation-induced damage cause elevated PRL levels, usually in the range of 30–100 $\mu\text{g/L}$. Plurihormonal adenomas (including GH and ACTH tumors) may hypersecrete PRL directly. Pituitary masses, including clinically nonfunctioning pituitary tumors, may compress the pituitary stalk to cause hyperprolactinemia.

Drug-induced inhibition or disruption of dopaminergic receptor function is a common cause of hyperprolactinemia (**Table 403-5**). Thus, antipsychotics and antidepressants are a relatively common cause of mild hyperprolactinemia. Most patients receiving risperidone have elevated prolactin levels, sometimes exceeding 200 $\mu\text{g/L}$. Methyl dopa inhibits dopamine synthesis, and verapamil blocks dopamine release, also leading to hyperprolactinemia. Hormonal agents that induce PRL include estrogens and thyrotropin-releasing hormone (TRH).

Presentation and Diagnosis Amenorrhea, galactorrhea, and infertility are the hallmarks of hyperprolactinemia in women. If hyperprolactinemia develops before menarche, primary amenorrhea results. More commonly, hyperprolactinemia develops later in life and leads to oligomenorrhea and ultimately to amenorrhea. If hyperprolactinemia is sustained, vertebral bone mineral density can be reduced compared with age-matched controls, particularly when it is associated with pronounced hypoestrogenemia. Galactorrhea is present in up to 80% of hyperprolactinemic women. Although usually bilateral and spontaneous, it may be unilateral or expressed only manually. Patients also may complain of decreased libido, weight gain, and mild hirsutism.

In men with hyperprolactinemia, diminished libido, infertility, and visual loss (from optic nerve compression) are the usual presenting symptoms. Gonadotropin suppression leads to reduced testosterone,

TABLE 403-5 ETIOLOGY OF HYPERPROLACTINEMIA

I. Physiologic hypersecretion

- Pregnancy
- Lactation
- Chest wall stimulation
- Sleep
- Stress

II. Hypothalamic–pituitary stalk damage

- Tumors
 - Craniopharyngioma
 - Suprasellar pituitary mass
 - Meningioma
 - Dysgerminoma
 - Metastases
- Empty sella
- Lymphocytic hypophysitis
- Adenoma with stalk
- Compression
- Granulomas
- Rathke's cyst
- Irradiation
- Trauma
 - Pituitary stalk section
 - Suprasellar surgery

III. Pituitary hypersecretion

- Prolactinoma
- Acromegaly

IV. Systemic disorders

- Chronic renal failure
- Hypothyroidism
- Cirrhosis
- Pseudocyesis
- Epileptic seizures

V. Drug-induced hypersecretion

- Dopamine receptor blockers
 - Atypical antipsychotics: risperidone
 - Phenothiazines: chlorpromazine, perphenazine
 - Butyrophenones: haloperidol
 - Thioxanthenes
 - Metoclopramide
- Dopamine synthesis inhibitors
 - α -Methyl dopa
- Catecholamine depletors
 - Reserpine
- Opiates
- H₂ antagonists
 - Cimetidine, ranitidine
- Imipramines
 - Amitriptyline, amoxapine
- Serotonin reuptake inhibitors
 - Fluoxetine
- Calcium channel blockers
 - Verapamil
- Estrogens
- Thyrotropin-releasing hormone

Note: Hyperprolactinemia >200 $\mu\text{g/L}$ almost invariably is indicative of a prolactin-secreting pituitary adenoma. Physiologic causes, hypothyroidism, and drug-induced hyperprolactinemia should be excluded before extensive evaluation.

impotence, and oligospermia. True galactorrhea is uncommon in men with hyperprolactinemia. If the disorder is long-standing, secondary effects of hypogonadism are evident, including osteopenia, reduced muscle mass, and decreased beard growth.

The diagnosis of idiopathic hyperprolactinemia is made by exclusion of known causes of hyperprolactinemia in the setting of a normal pituitary MRI. Some of these patients may harbor small microadenomas below visible MRI sensitivity (~2 mm).

GALACTORRHEA

Galactorrhea, the inappropriate discharge of milk-containing fluid from the breast, is considered abnormal if it persists longer than 6 months after childbirth or discontinuation of breast-feeding. Postpartum galactorrhea associated with amenorrhea is a self-limiting disorder usually associated with moderately elevated PRL levels. Galactorrhea may occur spontaneously, or it may be elicited by nipple pressure. In both men and women, galactorrhea may vary in color and consistency (transparent, milky, or bloody) and arise either unilaterally or bilaterally. Mammography or ultrasound is indicated for bloody discharges (particularly from a single nipple), which may be caused by breast cancer. Galactorrhea is commonly associated with hyperprolactinemia caused by any of the conditions listed in Table 403-5. Acromegaly is associated with galactorrhea in about one-third of patients. Treatment of galactorrhea usually involves managing the underlying disorder (e.g., replacing T₄ for hypothyroidism, discontinuing a medication, treating prolactinoma).

Laboratory Investigation Basal, fasting morning PRL levels (normally <20 µg/L) should be measured to assess hypersecretion. Both false-positive and false-negative results may be encountered. In patients with markedly elevated PRL levels (>1000 µg/L), reported results may be falsely lowered because of assay artifacts; sample dilution is required to measure these high values accurately. Falsely elevated values may be caused by aggregated forms of circulating PRL, which are usually biologically inactive (macroprolactinemia). Hypothyroidism should be excluded by measuring TSH and T₄ levels.

TREATMENT HYPERPROLACTINEMIA

Treatment of hyperprolactinemia depends on the cause of elevated PRL levels. Regardless of the etiology, however, treatment should be aimed at normalizing PRL levels to alleviate suppressive effects on gonadal function, halt galactorrhea, and preserve bone mineral density. Dopamine agonists are effective for most causes of hyperprolactinemia (see the treatment section for prolactinoma, below) regardless of the underlying cause.

If the patient is taking a medication known to cause hyperprolactinemia, the drug should be withdrawn, if possible. For psychiatric patients who require neuroleptic agents, supervised dose titration or the addition of a dopamine agonist can help restore normoprolactinemia and alleviate reproductive symptoms. However, dopamine agonists may worsen the underlying psychiatric condition, especially at high doses. Hyperprolactinemia usually resolves after adequate thyroid hormone replacement in hypothyroid patients or after renal transplantation in patients undergoing dialysis. Resection of hypothalamic or sellar mass lesions can reverse hyperprolactinemia caused by stalk compression and reduced dopamine tone. Granulomatous infiltrates occasionally respond to glucocorticoid administration. In patients with irreversible hypothalamic damage, no treatment may be warranted. In up to 30% of patients with hyperprolactinemia—usually without a visible pituitary microadenoma—the condition may resolve spontaneously.

PROLACTINOMA

Etiology and Prevalence Tumors arising from lactotrope cells account for about half of all functioning pituitary tumors, with a population prevalence of ~10/100,000 in men and ~30/100,000 in women. Mixed tumors that secrete combinations of GH and PRL, ACTH and PRL, and rarely TSH and PRL are also seen. These plurihormonal tumors are usually recognized by immunohistochemistry, sometimes without

apparent clinical manifestations from the production of additional hormones. Microadenomas are classified as <1 cm in diameter and usually do not invade the parasellar region. Macroadenomas are >1 cm in diameter and may be locally invasive and impinge on adjacent structures. The female-to-male ratio for microprolactinomas is 20:1, whereas the sex ratio is near 1:1 for macroadenomas. Tumor size generally correlates directly with PRL concentrations; values >250 µg/L usually are associated with macroadenomas. Men tend to present with larger tumors than women, possibly because the features of male hypogonadism are less readily evident. PRL levels remain stable in most patients, reflecting the slow growth of these tumors. About 5% of microadenomas progress in the long term to macroadenomas.

Presentation and Diagnosis Women usually present with amenorrhea, infertility, and galactorrhea. If the tumor extends outside the sella, visual field defects or other mass effects may be seen. Men often present with impotence, loss of libido, infertility, or signs of central nervous system (CNS) compression, including headaches and visual defects. Assuming that physiologic and medication-induced causes of hyperprolactinemia are excluded (Table 403-5), the diagnosis of prolactinoma is likely with a PRL level >200 µg/L. PRL levels <100 µg/L may be caused by microadenomas, other sellar lesions that decrease dopamine inhibition, or nonneoplastic causes of hyperprolactinemia. For this reason, an MRI should be performed in all patients with hyperprolactinemia. It is important to remember that hyperprolactinemia caused secondarily by the mass effects of nonlactotrope lesions is also corrected by treatment with dopamine agonists despite failure to shrink the underlying mass. Consequently, PRL suppression by dopamine agonists does not necessarily indicate that the underlying lesion is a prolactinoma.

TREATMENT PROLACTINOMA

Because microadenomas rarely progress to become macroadenomas, no treatment may be needed if patients are asymptomatic and fertility is not desired; these patients should be monitored by regular serial PRL measurements and MRI scans. For symptomatic microadenomas, therapeutic goals include control of hyperprolactinemia, reduction of tumor size, restoration of menses and fertility, and resolution of galactorrhea. Dopamine agonist doses should be titrated to achieve maximal PRL suppression and restoration of reproductive function (Fig. 403-3). A normalized PRL level does not ensure reduced tumor size. However, tumor shrinkage usually is not seen in those who do not respond with lowered PRL levels. For macroadenomas, formal visual field testing should be performed before initiating dopamine agonists. MRI and visual fields should be assessed at 6- to 12-month intervals until the mass shrinks and annually thereafter until maximum size reduction has occurred.

MEDICAL

Oral dopamine agonists (cabergoline and bromocriptine) are the mainstay of therapy for patients with micro- or macroprolactinomas. Dopamine agonists suppress PRL secretion and synthesis as well as lactotrope cell proliferation. In patients with microadenomas who have achieved normoprolactinemia and significant reduction of tumor mass, the dopamine agonist may be withdrawn after 2 years. These patients should be monitored carefully for evidence of prolactinoma recurrence. About 20% of patients (especially males) are resistant to dopaminergic treatment; these adenomas may exhibit decreased D₂ dopamine receptor numbers or a postreceptor defect. D₂ receptor gene mutations in the pituitary have not been reported.

Cabergoline An ergoline derivative, cabergoline is a long-acting dopamine agonist with high D₂ receptor affinity. The drug effectively suppresses PRL for >14 days after a single oral dose and induces prolactinoma shrinkage in most patients. Cabergoline (0.5–1.0 mg twice weekly) achieves normoprolactinemia and resumption of normal gonadal function in ~80% of patients with microadenomas; galactorrhea improves or resolves in 90% of patients. Cabergoline normalizes PRL and shrinks ~70% of macroprolactinomas. Mass

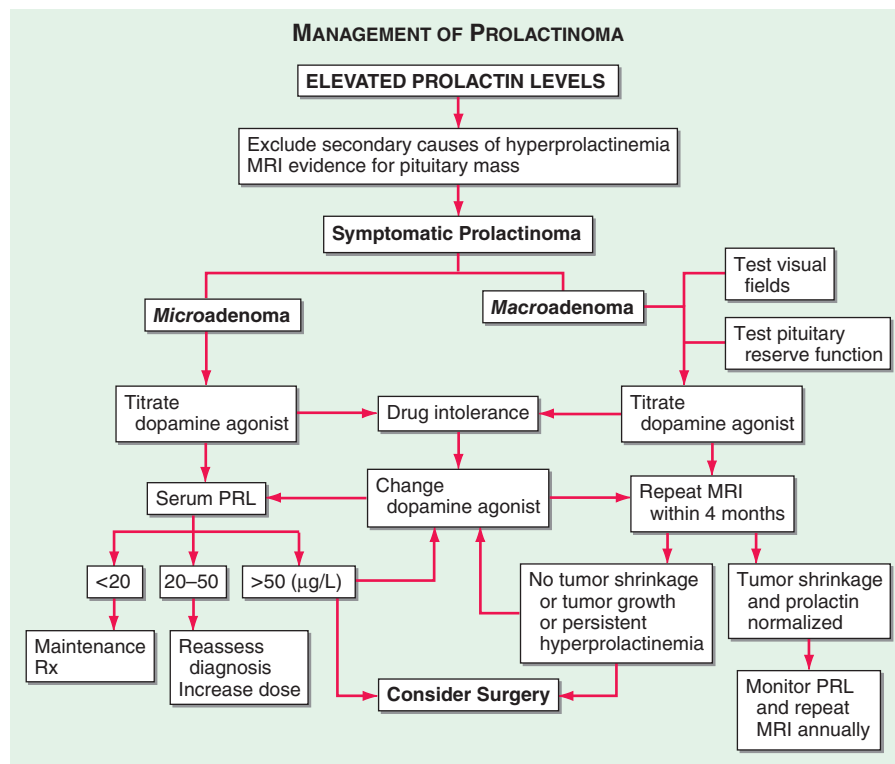


FIGURE 403-3 Management of prolactinoma. MRI, magnetic resonance imaging; PRL, prolactin.

effect symptoms, including headaches and visual disorders, usually improve dramatically within days after cabergoline initiation; improvement of sexual function requires several weeks of treatment but may occur before complete normalization of prolactin levels. After initial control of PRL levels has been achieved, cabergoline should be reduced to the lowest effective maintenance dose. In ~5% of treated patients harboring a microadenoma, hyperprolactinemia may resolve and not recur when dopamine agonists are discontinued after long-term treatment. Cabergoline also may be effective in patients resistant to bromocriptine. Adverse effects and drug intolerance are encountered less commonly than with bromocriptine.

BROMOCRRIPTINE

The ergot alkaloid bromocriptine mesylate is a dopamine receptor agonist that suppresses prolactin secretion. Because it is short-acting, the drug is preferred when pregnancy is desired. In microadenomas, bromocriptine rapidly lowers serum prolactin levels to normal in up to 70% of patients, decreases tumor size, and restores gonadal function. In patients with macroadenomas, prolactin levels are also normalized in 70% of patients, and tumor mass shrinkage ($\geq 50\%$) is achieved in most patients.

Therapy is initiated by administering a low bromocriptine dose (0.625–1.25 mg) at bedtime with a snack, followed by gradually increasing the dose. Most patients are controlled with a daily dose of ≤ 7.5 mg (2.5 mg tid).

SIDE EFFECTS

Side effects of dopamine agonists include constipation, nasal stuffiness, dry mouth, nightmares, insomnia, and vertigo; decreasing the dose usually alleviates these problems. Nausea, vomiting, and postural hypotension with faintness may occur in ~25% of patients after the initial dose. These symptoms may persist in some patients. In general, fewer side effects are reported with cabergoline. For the approximately 15% of patients who are intolerant of oral bromocriptine, cabergoline may be better tolerated. Intravaginal administration of bromocriptine is often efficacious in patients with intractable gastrointestinal side effects. Auditory hallucinations, delusions, and mood swings have been reported in up to 5% of patients and may

be due to the dopamine agonist properties or to the lysergic acid derivative of the compounds. Rare reports of leukopenia, thrombocytopenia, pleural fibrosis, cardiac arrhythmias, and hepatitis have been described. Patients with Parkinson's disease who receive at least 3 mg of cabergoline daily have been reported to be at risk for development of cardiac valve regurgitation. Studies analyzing over 500 prolactinoma patients receiving recommended doses of cabergoline (up to 2 mg weekly) have shown no evidence for an increased incidence of valvular disorders. Nevertheless, because no controlled prospective studies in pituitary tumor patients are available, it is prudent to perform echocardiograms before initiating standard-dose cabergoline therapy.

Surgery Indications for surgical adenoma debulking include dopamine resistance or intolerance and the presence of an invasive macroadenoma with compromised vision that fails to improve after drug treatment. Initial PRL normalization is achieved in about 70% of microprolactinomas after surgical resection, but only 30% of macroadenomas can be resected successfully. Follow-up studies have shown that hyperprolactinemia recurs in up to 20% of patients within the first year after surgery; long-term recurrence rates exceed 50% for macroadenomas. Radiotherapy for prolactinomas is reserved for patients with aggressive tumors that do not respond to maximally tolerated dopamine agonists and/or surgery.

PREGNANCY

The pituitary increases in size during pregnancy, reflecting the stimulatory effects of estrogen and perhaps other growth factors on pituitary vascularity and lactotrope cell hyperplasia. About 5% of microadenomas significantly increase in size, but 15–30% of macroadenomas grow during pregnancy. Bromocriptine has been used for more than 30 years to restore fertility in women with hyperprolactinemia, without evidence of teratogenic effects. Nonetheless, most authorities recommend strategies to minimize fetal exposure to the drug. For women taking bromocriptine who desire pregnancy, mechanical contraception should be used through three regular menstrual cycles to allow for conception timing. When pregnancy is confirmed, bromocriptine should

be discontinued and PRL levels followed serially, especially if headaches or visual symptoms occur. For women harboring macroadenomas, regular visual field testing is recommended, and the drug should be reinstated if tumor growth is apparent. Although pituitary MRI may be safe during pregnancy, this procedure should be reserved for symptomatic patients with severe headache and/or visual field defects. Surgical decompression may be indicated if vision is threatened. Although comprehensive data support the efficacy and relative safety of bromocriptine-facilitated fertility, patients should be advised of potential unknown deleterious effects and the risk of tumor growth during pregnancy. Because cabergoline is long-acting with a high D_2 -receptor affinity, it is not recommended for use in women when fertility is desired.

ACROMEGALY

Etiology GH hypersecretion is usually the result of a somatotrope adenoma but may rarely be caused by extrapituitary lesions (Table 403-6). In addition to the more common GH-secreting somatotrope adenomas, mixed mammosomatotrope tumors and acidophilic stem-cell adenomas secrete both GH and PRL. In patients with acidophilic stem-cell adenomas, features of hyperprolactinemia (hypogonadism and galactorrhea) predominate over the less clinically evident signs of acromegaly. Occasionally, mixed plurihormonal tumors are encountered that also secrete ACTH, the glycoprotein hormone α subunit, or TSH in addition to GH. Patients with partially empty sellas may present with GH hypersecretion due to a small GH-secreting adenoma within the compressed rim of pituitary tissue; some of these may reflect the spontaneous necrosis of tumors that were previously larger. GH-secreting tumors rarely arise from ectopic pituitary tissue remnants in the nasopharynx or midline sinuses.

There are case reports of ectopic GH secretion by tumors of pancreatic, ovarian, lung, or hematopoietic origin. Rarely, excess GHRH production may cause acromegaly because of chronic stimulation of somatotropes. These patients present with classic features of acromegaly, elevated GH levels, pituitary enlargement on MRI, and pathologic characteristics of pituitary hyperplasia. The most common cause of GHRH-mediated acromegaly is a chest or abdominal carcinoid tumor.

Although these tumors usually express positive GHRH immunoreactivity, clinical features of acromegaly are evident in only a minority of patients with carcinoid disease. Excessive GHRH also may be elaborated by hypothalamic tumors, usually choristomas or neuromas.

Presentation and Diagnosis Protean manifestations of GH and IGF-I hypersecretion are indolent and often are not clinically diagnosed for 10 years or more. Acral bony overgrowth results in frontal bossing, increased hand and foot size, mandibular enlargement with prognathism, and widened space between the lower incisor teeth. In children and adolescents, initiation of GH hypersecretion before epiphyseal long bone closure is associated with development of pituitary gigantism (Fig. 403-4). Soft tissue swelling results in increased heel pad thickness, increased shoe or glove size, ring tightening, characteristic coarse facial features, and a large fleshy nose. Other commonly encountered clinical features include hyperhidrosis, a deep and hollow-sounding voice, oily skin, arthropathy, kyphosis, carpal tunnel syndrome, proximal muscle weakness and fatigue, acanthosis nigricans, and skin tags. Generalized visceromegaly occurs, including cardiomegaly, macroglossia, and thyroid gland enlargement.

The most significant clinical impact of GH excess occurs with respect to the cardiovascular system. Coronary heart disease, cardiomyopathy with arrhythmias, left ventricular hypertrophy, decreased diastolic function, and hypertension ultimately occur in most patients if untreated. Upper airway obstruction with sleep apnea occurs in more than 60% of patients and is associated with both soft tissue laryngeal airway obstruction and central sleep dysfunction. Diabetes mellitus develops in 25% of patients with acromegaly, and most patients are intolerant of a glucose load (as GH counteracts the action of insulin). Acromegaly is associated with an increased risk of colon polyps and mortality from colonic malignancy; polyps are diagnosed in up to one-third of patients. Overall mortality is increased about threefold and is due primarily to cardiovascular and cerebrovascular disorders and respiratory disease. Unless GH levels are controlled, survival is reduced by an average of 10 years compared with an age-matched control population.

Laboratory Investigation Age-matched serum IGF-I levels are elevated in acromegaly. Consequently, an IGF-I level provides a useful laboratory screening measure when clinical features raise the possibility of acromegaly. Due to the pulsatility of GH secretion, measurement of a single random GH level is not useful for the diagnosis or exclusion of acromegaly and does not correlate with disease severity. The diagnosis of acromegaly is confirmed by demonstrating the failure of GH suppression to $<0.4 \mu\text{g/L}$ within 1–2 h of an oral glucose load (75 g). When newer ultrasensitive GH assays are used, normal nadir GH levels are even lower ($<0.05 \mu\text{g/L}$). About 20% of patients exhibit a paradoxical GH rise after glucose. PRL should be measured, as it is elevated in ~25% of patients with acromegaly. Thyroid function, gonadotropins, and sex steroids may be attenuated because of tumor mass effects. Because most patients will undergo surgery with glucocorticoid coverage, tests of ACTH reserve in asymptomatic patients are more efficiently deferred until after surgery.

TABLE 403-6 CAUSES OF ACROMEGALY

	Prevalence, %
Excess Growth Hormone Secretion	
Pituitary	98
Densely or sparsely granulated GH cell adenoma	60
Mixed GH cell and PRL cell adenoma	25
Mammosomatotrope cell adenoma	10
Plurihormonal adenoma	
GH cell carcinoma or metastases	
Multiple endocrine neoplasia 1 (GH cell adenoma)	
McCune-Albright syndrome	
Ectopic sphenoid or parapharyngeal sinus pituitary adenoma	
Extrapituitary tumor	
Pancreatic islet cell tumor	<1
Lymphoma	
Excess Growth Hormone–Releasing Hormone Secretion	
Central	<1
Hypothalamic hamartoma, choristoma, ganglioneuroma	<1
Peripheral	<1
Bronchial carcinoid, pancreatic islet cell tumor, small cell lung cancer, adrenal adenoma, medullary thyroid carcinoma, pheochromocytoma	

Abbreviations: GH, growth hormone; PRL, prolactin.

Source: Adapted from S Melmed: *N Engl J Med* 355:2558–2573, 2006.

TREATMENT ACROMEGALY

The goal of treatment is to control GH and IGF-I hypersecretion, ablate or arrest tumor growth, ameliorate comorbidities, restore mortality rates to normal, and preserve pituitary function.

Surgical resection of GH-secreting adenomas is the initial treatment for most patients (Fig. 403-5). Somatostatin analogues are used as adjuvant treatment for preoperative shrinkage of large invasive macroadenomas, immediate relief of debilitating symptoms, and reduction of GH hypersecretion; in frail patients experiencing morbidity; and in patients who decline surgery or, when surgery fails, to achieve biochemical control. Irradiation or repeat surgery may be required for patients who cannot tolerate or do not respond to adjunctive medical therapy. The high rate of late hypopituitarism and the slow rate (5–15 years) of biochemical response are the

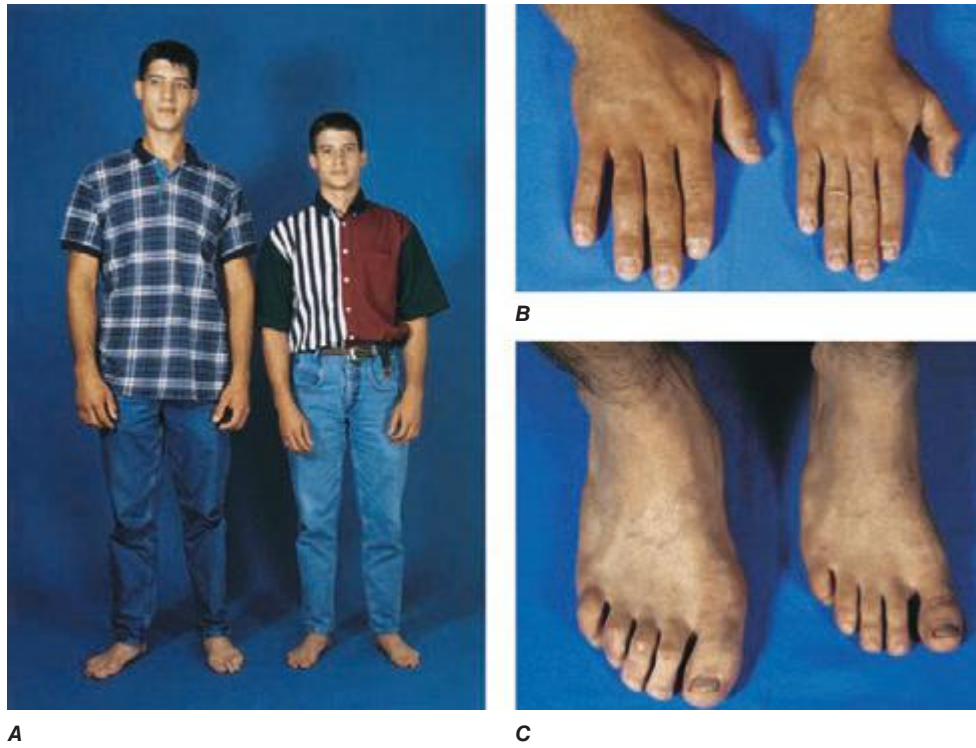


FIGURE 403-4 Features of acromegaly/gigantism. A 22-year-old man with gigantism due to excess growth hormone is shown to the left of his identical twin. The increased height and prognathism (A) and enlarged hand (B) and foot (C) of the affected twin are apparent. Their clinical features began to diverge at the age of approximately 13 years. (Reproduced from R Gagel, IE McCutcheon: *N Engl J Med* 324:524, 1999; with permission.)

main disadvantages of radiotherapy. Irradiation is also relatively ineffective in normalizing IGF-I levels. Stereotactic ablation of GH-secreting adenomas by gamma-knife radiotherapy is promising, but initial reports suggest that long-term results and side effects are similar to those observed with conventional radiation. Somatostatin analogues may be required while awaiting the full benefits of radiotherapy. Systemic co-morbid sequelae of acromegaly, including cardiovascular disease, diabetes, and arthritis, should be managed aggressively. Mandibular surgical repair may be indicated.

SURGERY

Transsphenoidal surgical resection by an experienced surgeon is the preferred primary treatment for both microadenomas (remission rate ~70%) and macroadenomas (<50% in remission). Soft tissue swelling improves immediately after tumor resection. GH levels return to normal within an hour, and IGF-I levels are normalized within 3–4 days. In ~10% of patients, acromegaly may recur several years after apparently successful surgery; hypopituitarism develops in up to 15% of patients after surgery.

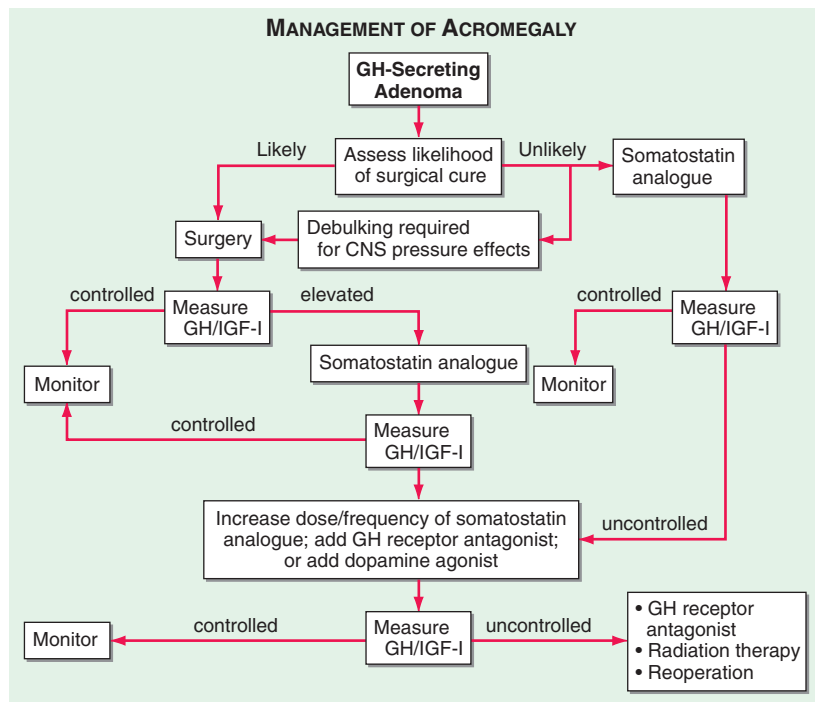


FIGURE 403-5 Management of acromegaly. GH, growth hormone; CNS, central nervous system; IGF, insulin-like growth factor. (Adapted from S Melmed et al: *J Clin Endocrinol Metab* 94:1509–1517, 2009; © The Endocrine Society.)

SOMATOSTATIN ANALOGUES

Somatostatin analogues exert their therapeutic effects through SSTR2 and SSTR5 receptors, both of which are expressed by GH-secreting tumors. Octreotide acetate is an eight-amino-acid synthetic somatostatin analogue. In contrast to native somatostatin, the analogue is relatively resistant to plasma degradation. It has a 2-h serum half-life and possesses fortyfold greater potency than native somatostatin to suppress GH. Octreotide is administered by subcutaneous injection, beginning with 50 µg tid; the dose can be increased gradually up to 1500 µg/d. Fewer than 10% of patients do not respond to the analogue. Octreotide suppresses integrated GH levels and normalizes IGF-I levels in ~60% of treated patients.

The long-acting somatostatin depot formulations, octreotide and lanreotide, are the preferred medical treatment for patients with acromegaly. *Sandostatin-LAR* is a sustained-release, long-acting formulation of octreotide incorporated into microspheres that sustain drug levels for several weeks after intramuscular injection. GH suppression occurs for as long as 6 weeks after a 30-mg intramuscular injection; long-term monthly treatment sustains GH and IGF-I suppression and also reduces pituitary tumor size in ~50% of patients. *Lanreotide* autogel, a slow-release depot somatostatin preparation, is a cyclic somatostatin octapeptide analogue that suppresses GH and IGF-I hypersecretion after a 60-mg subcutaneous injection. Long-term (4–6 weeks) administration controls GH hypersecretion in about two-thirds of treated patients and improves patient compliance because of the long interval required between drug injections. Rapid relief of headache and soft tissue swelling occurs in ~75% of patients within days to weeks of somatostatin analogue initiation. Most patients report symptomatic improvement, including amelioration of headache, perspiration, obstructive apnea, and cardiac failure.

Side Effects Somatostatin analogues are well tolerated in most patients. Adverse effects are short-lived and mostly relate to drug-induced suppression of gastrointestinal motility and secretion. Transient nausea, abdominal discomfort, fat malabsorption, diarrhea, and flatulence occur in one-third of patients, and these symptoms usually remit within 2 weeks. Octreotide suppresses postprandial gallbladder contractility and delays gallbladder emptying; up to 30% of patients develop long-term echogenic sludge or asymptomatic cholesterol gallstones. Other side effects include mild glucose intolerance due to transient insulin suppression, asymptomatic bradycardia, hypothyroxinemia, and local injection site discomfort.

GH RECEPTOR ANTAGONIST

Pegvisomant antagonizes endogenous GH action by blocking peripheral GH binding to its receptor. Consequently, serum IGF-I levels are suppressed, reducing the deleterious effects of excess endogenous GH. Pegvisomant is administered by daily subcutaneous injection (10–20 mg) and normalizes IGF-I in ~70% of patients. GH levels, however, remain elevated as the drug does not target the pituitary adenoma. Side effects include reversible liver enzyme elevation, lipodystrophy, and injection site pain. Tumor size should be monitored by MRI.

Combined treatment with monthly somatostatin analogues and weekly or biweekly pegvisomant injections has been used effectively in resistant patients.

DOPAMINE AGONISTS

Bromocriptine and cabergoline may modestly suppress GH secretion in some patients. Very high doses of bromocriptine (≥20 mg/d) or cabergoline (0.5 mg/d) are usually required to achieve modest GH therapeutic efficacy. Combined treatment with octreotide and cabergoline may induce additive biochemical control compared with either drug alone.

RADIATION

External radiation therapy or high-energy stereotactic techniques are used as adjuvant therapy for acromegaly. An advantage of radiation is that patient compliance with long-term treatment is not required. Tumor mass is reduced, and GH levels are attenuated over time.

However, 50% of patients require at least 8 years for GH levels to be suppressed to <5 µg/L; this level of GH reduction is achieved in about 90% of patients after 18 years but represents suboptimal GH suppression. Patients may require interim medical therapy for several years before attaining maximal radiation benefits. Most patients also experience hypothalamic-pituitary damage, leading to gonadotropin, ACTH, and/or TSH deficiency within 10 years of therapy.

In summary, surgery is the preferred primary treatment for GH-secreting microadenomas (Fig. 403-5). The high frequency of GH hypersecretion after macroadenoma resection usually necessitates adjuvant or primary medical therapy for these larger tumors. Patients unable to receive or respond to unimodal medical treatment may benefit from combined treatments, or can be offered radiation.

CUSHING'S SYNDROME (ACTH-PRODUCING ADENOMA)

(See also Chap. 406)

Etiology and Prevalence Pituitary corticotrope adenomas account for 70% of patients with endogenous causes of Cushing's syndrome. However, it should be emphasized that iatrogenic hypercortisolism is the most common cause of cushingoid features. Ectopic tumor ACTH production, cortisol-producing adrenal adenomas, adrenal carcinoma, and adrenal hyperplasia account for the other causes; rarely, ectopic tumor CRH production is encountered.

ACTH-producing adenomas account for about 10–15% of all pituitary tumors. Because the clinical features of Cushing's syndrome often lead to early diagnosis, most ACTH-producing pituitary tumors are relatively small microadenomas. However, macroadenomas also are seen and some ACTH-expressing adenomas are clinically silent. Cushing's disease is 5–10 times more common in women than in men. These pituitary adenomas exhibit unrestrained ACTH secretion, with resultant hypercortisolemia. However, they retain partial suppressibility in the presence of high doses of administered glucocorticoids, providing the basis for dynamic testing to distinguish pituitary from nonpituitary causes of Cushing's syndrome.

Presentation and Diagnosis The diagnosis of Cushing's syndrome presents two great challenges: (1) to distinguish patients with pathologic cortisol excess from those with physiologic or other disturbances of cortisol production and (2) to determine the etiology of pathologic cortisol excess.

Typical features of chronic cortisol excess include thin skin, central obesity, hypertension, plethoric moon facies, purple striae and easy bruising, glucose intolerance or diabetes mellitus, gonadal dysfunction, osteoporosis, proximal muscle weakness, signs of hyperandrogenism (acne, hirsutism), and psychological disturbances (depression, mania, and psychoses) (Table 403-7). Hematopoietic features of hypercortisolism include leukocytosis, lymphopenia, and eosinopenia. Immune suppression includes delayed hypersensitivity and infection propensity. These protean yet commonly encountered manifestations of hypercortisolism make it challenging to decide which patients mandate formal laboratory evaluation. Certain features make pathologic causes of hypercortisolism more likely; they include characteristic central redistribution of fat, thin skin with striae and bruising, and proximal muscle weakness. In children and young females, early osteoporosis may be particularly prominent. The primary cause of death is cardiovascular disease, but life-threatening infections and risk of suicide are also increased.

Rapid development of features of hypercortisolism associated with skin hyperpigmentation and severe myopathy suggests an ectopic tumor source of ACTH. Hypertension, hypokalemic alkalosis, glucose intolerance, and edema are also more pronounced in these patients. Serum potassium levels <3.3 mmol/L are evident in ~70% of patients with ectopic ACTH secretion but are seen in <10% of patients with pituitary-dependent Cushing's syndrome.

Laboratory Investigation The diagnosis of Cushing's syndrome is based on laboratory documentation of endogenous hypercortisolism. Measurement of 24-h urine free cortisol (UFC) is a precise and cost-effective screening test. Alternatively, the failure to suppress plasma cortisol after an overnight 1-mg dexamethasone suppression test can

TABLE 403-7 CLINICAL FEATURES OF CUSHING'S SYNDROME (ALL AGES)

Symptoms/Signs	Frequency, %
Obesity or weight gain (>115% ideal body weight)	80
Thin skin	80
Moon facies	75
Hypertension	75
Purple skin striae	65
Hirsutism	65
Menstrual disorders (usually amenorrhea)	60
Plethora	60
Abnormal glucose tolerance	55
Impotence	55
Proximal muscle weakness	50
Truncal obesity	50
Acne	45
Bruising	45
Mental changes	45
Osteoporosis	40
Edema of lower extremities	30
Hyperpigmentation	20
Hypokalemic alkalosis	15
Diabetes mellitus	15

Source: Adapted from MA Magiokou et al, in ME Wierman (ed): *Diseases of the Pituitary*. Totowa, NJ, Humana, 1997.

be used to identify patients with hypercortisolism. As nadir levels of cortisol occur at night, elevated midnight serum or salivary samples of cortisol are suggestive of Cushing's syndrome. Basal plasma ACTH levels often distinguish patients with ACTH-independent (adrenal or exogenous glucocorticoid) from those with ACTH-dependent (pituitary, ectopic ACTH) Cushing's syndrome. Mean basal ACTH levels are about eightfold higher in patients with ectopic ACTH secretion than in those with pituitary ACTH-secreting adenomas. However, extensive overlap of ACTH levels in these two disorders precludes using ACTH measurements to make the distinction. Preferably, dynamic testing based on differential sensitivity to glucocorticoid feedback or ACTH stimulation in response to CRH or cortisol reduction is used to distinguish ectopic from pituitary sources of excess ACTH (Table 403-8). Very rarely, circulating CRH levels are elevated, reflecting ectopic tumor-derived secretion of CRH and often ACTH. **For further discussion of dynamic testing for Cushing's syndrome, see Chap. 406.**

Most ACTH-secreting pituitary tumors are <5 mm in diameter, and about half are undetectable by sensitive MRI. The high prevalence of incidental pituitary microadenomas diminishes the ability to distinguish ACTH-secreting pituitary tumors accurately from nonsecreting incidentalomas.

Inferior Petrosal Venous Sampling Because pituitary MRI with gadolinium enhancement is insufficiently sensitive to detect small (<2 mm) pituitary ACTH-secreting adenomas, bilateral inferior petrosal sinus ACTH sampling before and after CRH administration may be required to distinguish these lesions from ectopic ACTH-secreting tumors that may have similar clinical and biochemical characteristics. Simultaneous assessment of ACTH in each inferior petrosal vein and in the diagnosis of peripheral circulation provides a strategy for confirming and localizing pituitary ACTH production. Sampling is performed at baseline and 2, 5, and 10 min after intravenous bovine CRH (1 µg/kg) injection. An increased ratio (>2) of inferior petrosal:peripheral vein ACTH confirms pituitary Cushing's syndrome. After CRH injection, peak petrosal:peripheral ACTH ratios ≥3 confirm the presence of a pituitary ACTH-secreting tumor. The sensitivity of this test is >95%, with very rare false-positive results. False-negative results may be encountered in patients with aberrant venous drainage. Petrosal sinus catheterizations are technically difficult, and about 0.05% of patients develop neurovascular complications. The procedure should not be performed in patients with hypertension, in patients with known cerebrovascular disease, or in the presence of a well-visualized pituitary adenoma on MRI.

TABLE 403-8 DIFFERENTIAL DIAGNOSIS OF ACTH-DEPENDENT CUSHING'S SYNDROME*

	ACTH-Secreting Pituitary Tumor	Ectopic ACTH Secretion
Etiology	Pituitary corticotrope adenoma Plurihormonal adenoma	Bronchial, abdominal carcinoid Small cell lung cancer Thymoma
Sex	F > M	M > F
Clinical features	Slow onset	Rapid onset Pigmentation Severe myopathy
Serum potassium <3.3µg/L	<10%	75%
24-h urinary free cortisol (UFC)	High	High
Basal ACTH level	Inappropriately high	Very high
Dexamethasone suppression		
1 mg overnight		
Low-dose (0.5 mg q6h)	Cortisol >5 µg/dL	Cortisol >5 µg/dL
High-dose (2 mg q6h)	Cortisol <5 µg/dL	Cortisol >5 µg/dL
UFC >80% suppressed	Microadenomas: 90% Macroadenomas: 50%	10%
Inferior petrosal sinus sampling (IPSS)		
Basal		
IPSS: peripheral	>2	<2
CRH-induced		
IPSS: peripheral	>3	<3

*ACTH-independent causes of Cushing's syndrome are diagnosed by suppressed ACTH levels and an adrenal mass in the setting of hypercortisolism. Iatrogenic Cushing's syndrome is excluded by history.

Abbreviations: ACTH, adrenocorticotropic hormone; CRH, corticotropin-releasing hormone; F, female; M, male.

TREATMENT CUSHING'S SYNDROME

Selective transsphenoidal resection is the treatment of choice for Cushing's disease (Fig. 403-6). The remission rate for this procedure is ~80% for microadenomas but <50% for macroadenomas. However, surgery is rarely successful when the adenoma is not visible on MRI. After successful tumor resection, most patients experience a postoperative period of symptomatic ACTH deficiency that may last up to 12 months. This usually requires low-dose cortisol replacement, as patients experience both steroid withdrawal symptoms and have a suppressed hypothalamic-pituitary-adrenal axis. Biochemical recurrence occurs in approximately 5% of patients in whom surgery was initially successful.

When initial surgery is unsuccessful, repeat surgery is sometimes indicated, particularly when a pituitary source for ACTH is well documented. In older patients, in whom issues of growth and fertility are less important, hemi- or total hypophysectomy may be necessary if a discrete pituitary adenoma is not recognized. Pituitary irradiation may be used after unsuccessful surgery, but it cures only about 15% of patients. Because the effects of radiation are slow and only partially effective in adults, steroidogenic inhibitors are used in combination with pituitary irradiation to block adrenal effects of persistently high ACTH levels.

Pasireotide (600 or 900 µg/day subcutaneously), a somatostatin analog with high affinity for SST5 > SST2 receptors, has been approved for treating patients with ACTH-secreting pituitary tumors when surgery is not an option or has been unsuccessful. In clinical trials, the drug lowered plasma ACTH levels, normalized 24-h urinary free cortisol levels in about 25% of patients, and resulted in up to 40% mean pituitary tumor shrinkage. Side effects include development of

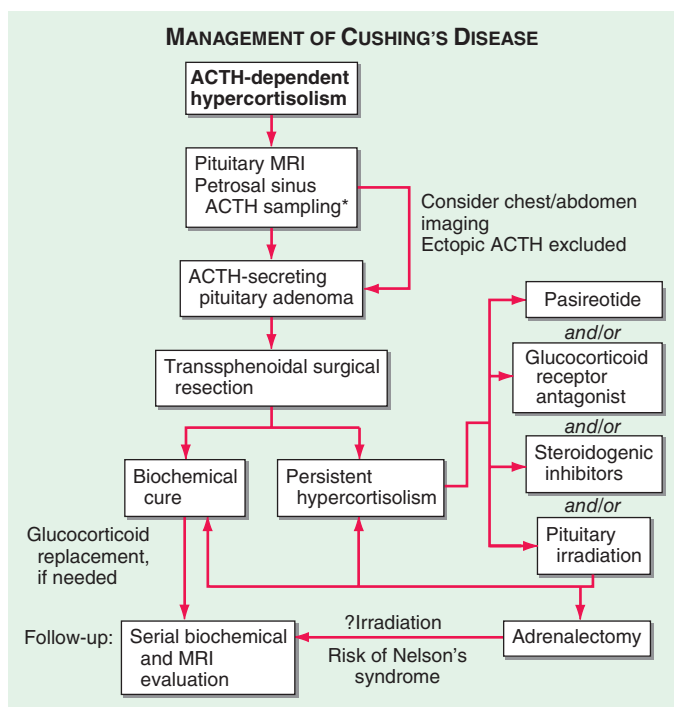


FIGURE 403-6 Management of Cushing's syndrome. ACTH, adrenocorticotropin hormone; MRI, magnetic resonance imaging. *, Not usually required.

hyperglycemia and diabetes in about 70% of patients, likely due to suppressed pancreatic secretion of insulin and incretins. Because patients with hypercortisolism are insulin-resistant, hyperglycemia should be rigorously managed. Other side effects are similar to those encountered for somatostatin analogs and include transient abdominal discomfort, diarrhea, nausea, and gallstones (20% of patients). The drug requires consistent long-term administration.

Ketoconazole, an imidazole derivative antimycotic agent, inhibits several P450 enzymes and effectively lowers cortisol in most patients with Cushing's disease when administered twice daily (600–1200 mg/d). Elevated hepatic transaminases, gynecomastia, impotence, gastrointestinal upset, and edema are common side effects.

Mifepristone (300–1200 mg/d), a glucocorticoid receptor antagonist, blocks peripheral cortisol action and is approved to treat hyperglycemia in Cushing's disease. Because the drug does not target the pituitary tumor, both ACTH and cortisol levels remain elevated, thus obviating a reliable circulating biomarker. Side effects are largely due to general antagonism of other steroid hormones and include hypokalemia, endometrial hyperplasia, hypoadrenalism, and hypertension.

Metyrapone (2–4 g/d) inhibits 11 β -hydroxylase activity and normalizes plasma cortisol in up to 75% of patients. Side effects include nausea and vomiting, rash, and exacerbation of acne or hirsutism. *Mitotane* (*o,p'*-DDD; 3–6 g/d orally in four divided doses) suppresses cortisol hypersecretion by inhibiting 11 β -hydroxylase and cholesterol side-chain cleavage enzymes and by destroying adrenocortical cells. Side effects of mitotane include gastrointestinal symptoms, dizziness, gynecomastia, hyperlipidemia, skin rash, and hepatic enzyme elevation. It also may lead to hypoadosteronism. Other agents include *aminoglutethimide* (250 mg tid), *trilostane* (200–1000 mg/d), *cyproheptadine* (24 mg/d), and IV *etomidate* (0.3 mg/kg per hour). Glucocorticoid insufficiency is a potential side effect of agents used to block steroidogenesis.

The use of steroidogenic inhibitors has decreased the need for bilateral adrenalectomy. Surgical removal of both adrenal glands corrects hypercortisolism but may be associated with significant morbidity rates and necessitates permanent glucocorticoid and mineralocorticoid replacement. Adrenalectomy in the setting of residual corticotrope adenoma tissue predisposes to the development of *Nelson's syndrome*, a disorder characterized by rapid pituitary tumor

enlargement and increased pigmentation secondary to high ACTH levels. Prophylactic radiation therapy may be indicated to prevent the development of Nelson's syndrome after adrenalectomy.

NONFUNCTIONING AND GONADOTROPIN-PRODUCING PITUITARY ADENOMAS

Etiology and Prevalence Nonfunctioning pituitary adenomas include those that secrete little or no pituitary hormones as well as tumors that produce too little hormone to result in recognizable clinical features. They are the most common type of pituitary adenoma and are usually macroadenomas at the time of diagnosis because clinical features are not apparent until tumor mass effects occur. Based on immunohistochemistry, most clinically nonfunctioning adenomas can be shown to originate from gonadotrope cells. These tumors typically produce small amounts of intact gonadotropins (usually FSH) as well as uncombined α , LH β , and FSH β subunits. Tumor secretion may lead to elevated α and FSH β subunits and, rarely, to increased LH β subunit levels. Some adenomas express α subunits without FSH or LH. TRH administration often induces an atypical increase of tumor-derived gonadotropins or subunits.

Presentation and Diagnosis Clinically nonfunctioning tumors often present with optic chiasm pressure and other symptoms of local expansion or may be incidentally discovered on an MRI performed for another indication (incidentaloma). Rarely, menstrual disturbances or ovarian hyperstimulation occur in women with large tumors that produce FSH and LH. More commonly, adenoma compression of the pituitary stalk or surrounding pituitary tissue leads to attenuated LH and features of hypogonadism. PRL levels are usually slightly increased, also because of stalk compression. It is important to distinguish this circumstance from true prolactinomas, as nonfunctioning tumors do not shrink in response to treatment with dopamine agonists.

Laboratory Investigation The goal of laboratory testing in clinically nonfunctioning tumors is to classify the type of the tumor, identify hormonal markers of tumor activity, and detect possible hypopituitarism. Free α subunit levels may be elevated in 10–15% of patients with nonfunctioning tumors. In female patients, peri- or postmenopausal basal FSH concentrations are difficult to distinguish from tumor-derived FSH elevation. Premenopausal women have cycling FSH levels, also preventing clear-cut diagnostic distinction from tumor-derived FSH. In men, gonadotropin-secreting tumors may be diagnosed because of slightly increased gonadotropins (FSH > LH) in the setting of a pituitary mass. Testosterone levels are usually low despite the normal or increased LH level, perhaps reflecting reduced LH bioactivity or the loss of normal LH pulsatility. Because this pattern of hormone test results is also seen in primary gonadal failure and, to some extent, with aging (Chap. 411), the finding of increased gonadotropins alone is insufficient for the diagnosis of a gonadotropin-secreting tumor. In the majority of patients with gonadotrope adenomas, TRH administration stimulates LH β subunit secretion; this response is not seen in normal individuals. GnRH testing, however, is not helpful for making the diagnosis. For nonfunctioning and gonadotropin-secreting tumors, the diagnosis usually rests on immunohistochemical analyses of surgically resected tumor tissue, as the mass effects of these tumors usually necessitate resection.

Although acromegaly or Cushing's syndrome usually presents with unique clinical features, clinically inapparent (silent) somatotrope or corticotrope adenomas may only be diagnosed by immunostaining of resected tumor tissue. If PRL levels are <100 μ g/L in a patient harboring a pituitary mass, a nonfunctioning adenoma causing pituitary stalk compression should be considered.

TREATMENT NONFUNCTIONING AND GONADOTROPIN-PRODUCING PITUITARY ADENOMAS

Asymptomatic small nonfunctioning microadenomas adenomas with no threat to vision may be followed with regular MRI and visual field testing without immediate intervention. However, for macroadenomas, transsphenoidal surgery is indicated to reduce tumor size and relieve mass effects (Fig. 403-7). Although it is not usually

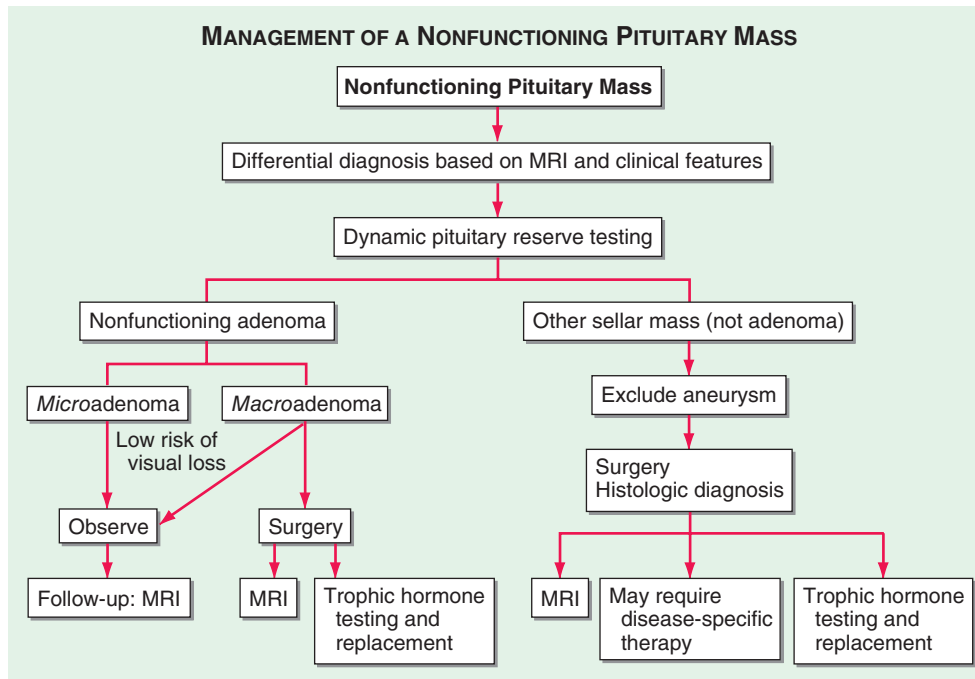


FIGURE 403-7 Management of a nonfunctioning pituitary mass. MRI, magnetic resonance imaging.

possible to remove all adenoma tissue surgically, vision improves in 70% of patients with preoperative visual field defects. Preexisting hypopituitarism that results from tumor mass effects may improve or resolve completely. Beginning about 6 months postoperatively, MRI scans should be performed yearly to detect tumor regrowth. Within 5–6 years after successful surgical resection, ~15% of nonfunctioning tumors recur. When substantial tumor remains after transsphenoidal surgery, adjuvant radiotherapy may be indicated to prevent tumor regrowth. Radiotherapy may be deferred if no postoperative residual mass is evident. Nonfunctioning pituitary tumors respond poorly to dopamine agonist treatment and somatostatin analogues are largely ineffective for shrinking these tumors. The selective GnRH antagonist Nal-Glu GnRH suppresses FSH hypersecretion but has no effect on adenoma size.

TSH-SECRETING ADENOMAS

TSH-producing macroadenomas are very rare but are often large and locally invasive when they occur. Patients usually present with thyroid goiter and hyperthyroidism, reflecting overproduction of TSH. Diagnosis is based on demonstrating elevated serum free T_4 levels, inappropriately normal or high TSH secretion, and MRI evidence of a pituitary adenoma. Elevated uncombined α subunits are seen in many patients.

It is important to exclude other causes of inappropriate TSH secretion, such as resistance to thyroid hormone, an autosomal dominant disorder caused by mutations in the thyroid hormone β receptor (Chap. 405). The presence of a pituitary mass and elevated β subunit levels are suggestive of a TSH-secreting tumor. Dysalbuminemic hyperthyroxinemia syndromes, caused by mutations in serum thyroid hormone binding proteins, are also characterized by elevated thyroid hormone levels, but with normal rather than suppressed TSH levels. Moreover, free thyroid hormone levels are normal in these disorders, most of which are familial.

TREATMENT TSH-SECRETING ADENOMAS

The initial therapeutic approach is to remove or debulk the tumor mass surgically, usually using a transsphenoidal approach. Total resection is not often achieved as most of these adenomas are large and locally invasive. Normal circulating thyroid hormone levels are achieved in about two-thirds of patients after surgery. Thyroid ablation or antithyroid drugs (methimazole and propylthiouracil) can be used to reduce thyroid hormone levels.

Somatostatin analogue treatment effectively normalizes TSH and α subunit hypersecretion, shrinks the tumor mass in 50% of patients, and improves visual fields in 75% of patients; euthyroidism is restored in most patients. Because somatostatin analogues markedly suppress TSH, biochemical hypothyroidism often requires concomitant thyroid hormone replacement, which may also further control tumor growth.

404 Disorders of the Neurohypophysis

Gary L. Robertson

The neurohypophysis, or posterior pituitary, is formed by axons that originate in large cell bodies in the supraoptic and paraventricular nuclei of the hypothalamus. It produces two hormones: (1) arginine vasopressin (AVP), also known as antidiuretic hormone, and (2) oxytocin. AVP acts on the renal tubules to reduce water loss by concentrating the urine. Oxytocin stimulates postpartum milk letdown in response to suckling. A deficiency of AVP secretion or action causes diabetes insipidus (DI), a syndrome characterized by the production of large amounts of dilute urine. Excessive or inappropriate AVP production impairs urinary water excretion and predisposes to hyponatremia if water intake is not reduced in parallel with urine output.

VASOPRESSIN

SYNTHESIS AND SECRETION

AVP is a nonapeptide composed of a six-member disulfide ring and a tripeptide tail (Fig. 404-1). It is synthesized via a polypeptide precursor that includes AVP, neurophysin, and copeptin, all encoded by a single gene on chromosome 20. After preliminary processing and folding, the precursor is packaged in neurosecretory vesicles, where it is transported down the axon; further processed to AVP, neurophysin, and copeptin; and stored in neurosecretory vesicles until released by exocytosis into peripheral blood.

AVP secretion is regulated primarily by the “effective” osmotic pressure of body fluids. This control is mediated by specialized hypothalamic cells known as *osmoreceptors*, which are extremely sensitive

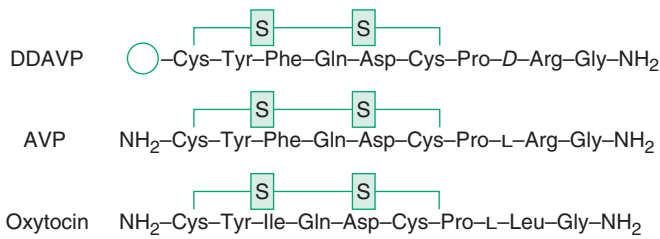


FIGURE 404-1 Primary structures of arginine vasopressin (AVP), oxytocin, and desmopressin (DDAVP).

to small changes in the plasma concentration of sodium and its anions but normally are insensitive to other solutes such as urea and glucose. The osmoreceptors appear to include inhibitory as well as stimulatory components that function in concert to form a threshold, or set point, control system. Below this threshold, plasma AVP is suppressed to levels that permit the development of a maximum water diuresis. Above it, plasma AVP rises steeply in direct proportion to plasma osmolarity, quickly reaching levels sufficient to effect a maximum antidiuresis. The absolute levels of plasma osmolarity/sodium at which minimally and maximally effective levels of plasma AVP occur, vary appreciably from person to person, apparently due to genetic influences on the set and sensitivity of the system. However, the average threshold, or set point, for AVP release corresponds to a plasma osmolarity or sodium of about 280 mosmol/L or 135 meq/L, respectively; levels only 2–4% higher normally result in maximum antidiuresis.

Although it is relatively stable in a healthy adult, the set point of the osmoregulatory system can be lowered by pregnancy, the menstrual cycle, estrogen, and relatively large, acute reductions in blood pressure or volume. Those reductions are mediated largely by neuronal afferents that originate in transmural pressure receptors of the heart and large arteries and project via the vagus and glossopharyngeal nerves to the brainstem, from which postsynaptic projections ascend to the hypothalamus. These pathways maintain a tonic inhibitory tone that decreases when blood volume or pressure falls by >10–20%. This baroregulatory system is probably of minor importance in the physiology of AVP secretion because the hemodynamic changes required to affect it usually do not occur during normal activities. However, the baroregulatory system undoubtedly plays an important role in AVP secretion in patients with disorders that produce large, acute disturbances of hemodynamic function. However, the baroregulatory system undoubtedly plays an important role in AVP secretion in patients with disorders that produce large, acute disturbances of hemodynamic function.

AVP secretion also can be stimulated by nausea, acute hypoglycemia, glucocorticoid deficiency, smoking, and, possibly, hyperangiotensinemia. The emetic stimuli are extremely potent since they typically elicit immediate, 50- to 100-fold increases in plasma AVP even when the nausea is transient and is not associated with vomiting or other symptoms. They appear to act via the emetic center in the medulla and can be blocked completely by treatment with antiemetics such as fluphenazine. There is no evidence that pain or other noxious stresses have any effect on AVP unless they elicit a vasovagal reaction with its associated nausea and hypotension.

ACTION

The most important, if not the only, physiologic action of AVP is to reduce water excretion by promoting concentration of urine. This antidiuretic effect is achieved by increasing the hydroosmotic permeability of cells that line the distal tubule and medullary collecting ducts of the kidney (Fig. 404-2). In the absence of AVP, these cells are impermeable to water and reabsorb little, if any, of the relatively large volume of dilute filtrate that enters from the proximal nephron. The lack of reabsorption results in the excretion of very large volumes (as much as 0.2 mL/kg per min) of maximally dilute urine (specific gravity and osmolarity ~1.000 and 50 mosmol/L, respectively), a condition known as *water diuresis*. In the presence of AVP, these cells become selectively permeable to water, allowing the water to diffuse back down the osmotic gradient created by the hypertonic renal medulla. As a result, the dilute fluid passing through the tubules is concentrated and the rate of urine flow decreases.

The magnitude of this effect varies in direct proportion to the plasma AVP concentration and the rate of solute excretion. At maximum levels of AVP and normal rates of solute excretion, it approximates a urine flow rate as low as 0.35 mL/min and a urine osmolarity as high as 1200 mosmol/L. This effect is reduced by a solute diuresis such as glucosuria in diabetes mellitus. Antidiuresis is mediated via binding to G protein-coupled V_2 receptors on the serosal surface of the cell, activation of adenylyl cyclase, and insertion into the luminal surface of water channels composed of a protein known as *aquaporin 2* (AQP2). The V_2 receptors and aquaporin 2 are encoded by genes on chromosomes Xq28 and 12q13, respectively.

At high concentrations, AVP also causes contraction of smooth muscle in blood vessels in the skin and gastrointestinal tract, induces glycogenolysis in the liver, and potentiates adrenocorticotrophic hormone (ACTH) release by corticotropin-releasing factor. These effects are mediated by V_{1a} or V_{1b} receptors that are coupled to phospholipase C. Their role, if any, in human physiology/pathophysiology is uncertain.

METABOLISM

AVP distributes rapidly into a space roughly equal to the extracellular fluid volume. It is cleared irreversibly with a half-life ($t_{1/2}$) of 10–30 min. Most AVP clearance is due to degradation in the liver and kidneys. During pregnancy, the metabolic clearance of AVP is increased three- to fourfold due to placental production of an N-terminal peptidase.

THIRST

Because AVP cannot reduce water loss below a certain minimum level obligated by urinary solute load and evaporation from skin and lungs, a mechanism for ensuring adequate intake is essential for preventing dehydration. This vital function is performed by the thirst mechanism. Like AVP, thirst is regulated primarily by an osmostat that is situated in the anteromedial hypothalamus and is able to detect very small changes in the plasma concentration of sodium and its anions. The thirst osmostat appears to be “set” about 3% higher than the AVP osmostat. This arrangement ensures that thirst, polydipsia, and dilution of body fluids do not occur until plasma osmolarity/sodium starts to exceed the defensive capacity of the antidiuretic mechanism.

OXYTOCIN

Oxytocin is also a nonapeptide that differs from AVP only at positions 3 and 8 (Fig. 404-1). However, it has relatively little antidiuretic effect and seems to act mainly on mammary ducts to facilitate milk letdown during nursing. It also may help initiate or facilitate labor by stimulating contraction of uterine smooth muscle, but it is not clear if this action is physiologic or necessary for normal delivery.

DEFICIENCIES OF AVP SECRETION AND ACTION

DIABETES INSIPIDUS

Clinical Characteristics A decrease of 75% or more in the secretion or action of AVP usually results in DI, a syndrome characterized by the production of abnormally large volumes of dilute urine. The 24-h urine volume exceeds 50 mL/kg body weight, and the osmolarity is less than 300 mosmol/L. The polyuria produces symptoms of urinary frequency, enuresis, and/or nocturia, which may disturb sleep and cause mild daytime fatigue or somnolence. It also results in a slight rise in plasma osmolarity that stimulates thirst and a commensurate increase in fluid intake (polydipsia). Overt clinical signs of dehydration are uncommon unless thirst and/or the compensatory increase of fluid intake are also impaired.



Etiology A primary deficiency of AVP secretion usually results from agenesis or irreversible destruction of the neurohypophysis. It is referred to variously as *neurohypophyseal DI*, *neurogenic DI*, *pituitary DI*, *cranial DI*, or *central DI*. It can be caused by a variety of congenital, acquired, or genetic disorders, but in about one-half of all adult patients, it is idiopathic (Table 404-1). Pituitary DI caused by surgery in or around the neurohypophysis usually appears within 24 h. After a few days, it may transition to a 2- to 3-week period of inappropriate antidiuresis, after which the DI may or may not recur permanently.

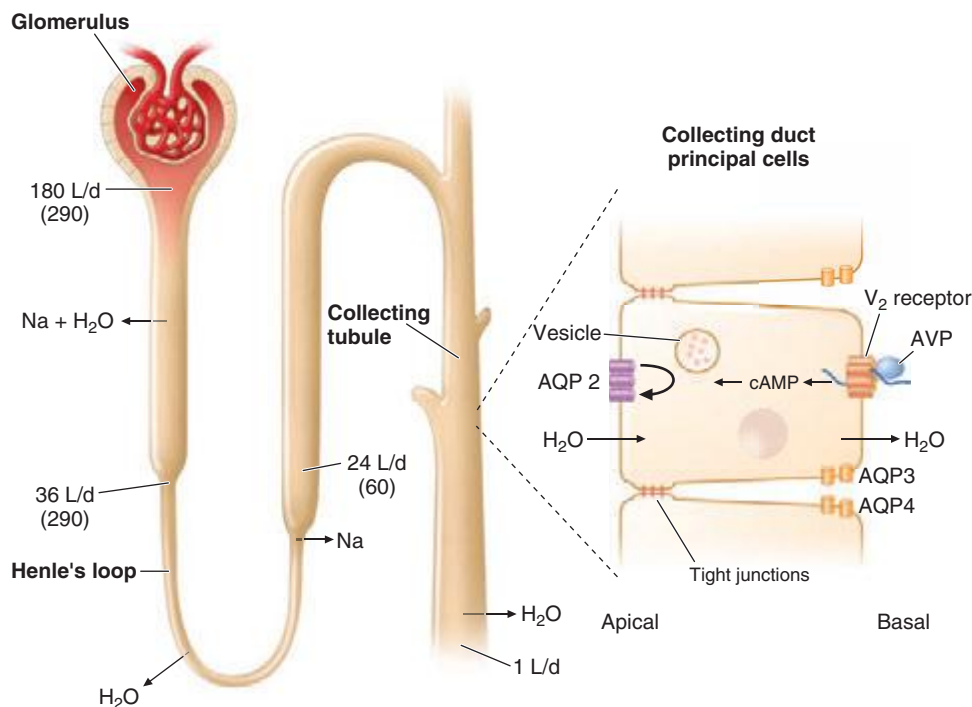


FIGURE 404-2 Antidiuretic effect of arginine vasopressin (AVP) in the regulation of urine volume. In a typical 70-kg adult, the kidney filters ~180 L/d of plasma. Of this, ~144 L (80%) is reabsorbed isosmotically in the proximal tubule and another 8 L (4–5%) is reabsorbed without solute in the descending limb of Henle's loop. The remainder is diluted to an osmolarity of ~60 mmol/kg by selective reabsorption of sodium and chloride in the ascending limb. In the absence of AVP, the urine issuing from the loop passes largely unmodified through the distal tubules and collecting ducts, resulting in a maximum water diuresis. In the presence of AVP, solute-free water is reabsorbed osmotically through the principal cells of the collecting ducts, resulting in the excretion of a much smaller volume of concentrated urine. This antidiuretic effect is mediated via a G protein–coupled V_2 receptor that increases intracellular cyclic AMP, thereby inducing translocation of aquaporin 2 (AQP 2) water channels into the apical membrane. The resultant increase in permeability permits an influx of water that diffuses out of the cell through AQP 3 and AQP 4 water channels in the basal-lateral surface. The net rate of flux across the cell is determined by the number of AQP 2 water channels in the apical membrane and the strength of the osmotic gradient between tubular fluid and the renal medulla. Tight junctions on the lateral surface of the cells serve to prevent unregulated water flow.

Five genetic forms of pituitary DI are now known. By far, the most common is transmitted in an autosomal dominant mode and is caused by diverse mutations in the coding region of one allele of the AVP–neurophysin II (or *AVP-NPII*) gene. All the mutations alter one or more amino acids known to be critical for correct processing and/or folding of the prohormone, thus interfering with its trafficking through the endoplasmic reticulum. The misfolded mutant precursor accumulates and interferes with production of AVP by the normal allele, eventually destroying the magnocellular neurons in which it is produced. The AVP deficiency and DI are usually not present at birth but develop gradually over a period of several months to years, progressing from partial to severe at different rates depending on the mutation. Once established, the deficiency of AVP is permanent, but for unknown reasons, the DI occasionally improves or remits spontaneously in late middle age. The parvocellular neurons that make AVP and the magnocellular neurons that make oxytocin appear to be unaffected. There are also rare autosomal recessive forms of pituitary DI. One is due to an inactivating mutation in the AVP portion of the gene; another is due to a large deletion involving the majority of the AVP gene and regulatory sequences in the intergenic region. A third form is caused by mutations of the *WFS 1* gene responsible for Wolfram's syndrome (DI, diabetes mellitus, optic atrophy, and neural deafness [DIDMOAD]). An X-linked recessive form linked to a region on Xq28 has also been described.

A primary deficiency of plasma AVP also can result from increased metabolism by an N-terminal aminopeptidase produced by the placenta. It is referred to as *gestational DI* because the signs and symptoms manifest during pregnancy and usually remit several weeks after delivery.

Secondary deficiencies of AVP secretion result from inhibition by excessive intake of fluids. They are referred to as *primary polydipsia* and can be divided into three subcategories. One of them, *dipsogenic DI*, is

characterized by inappropriate thirst caused by a reduction in the set of the osmoregulatory mechanism. It sometimes occurs in association with multifocal diseases of the brain such as neurosarcoid, tuberculous meningitis, and multiple sclerosis but is often idiopathic. The second subtype, *psychogenic polydipsia*, is not associated with thirst, and the polydipsia seems to be a feature of psychosis or obsessive compulsive disorder. The third subtype, *iatrogenic polydipsia*, results from recommendations to increase fluid intake for its presumed health benefits.

Primary deficiencies in the antidiuretic action of AVP result in *nephrogenic DI*. The causes can be genetic, acquired, or drug induced (Table 404-1). The most common genetic form is transmitted in a semirecessive X-linked manner. It is caused by mutations in the coding region of the V_2 receptor gene that impair trafficking and/or ligand binding of the mutant receptor. There are also autosomal recessive or dominant forms of nephrogenic DI. They are caused by *AQP2* gene mutations that result in complete or partial defects in trafficking and function of the water channels that mediate antidiuresis in the distal and collecting tubules of the kidney.

Secondary deficiencies in the antidiuretic response to AVP result from polyuria per se. They are caused by washout of the medullary concentration gradient and/or suppression of aquaporin function. They usually resolve 24–48 h after the polyuria is corrected but can complicate interpretation of some acute tests used for differential diagnosis.

Pathophysiology In pituitary, gestational, or nephrogenic DI, the polyuria results in a small (1–2%) decrease in body water and a commensurate increase in plasma osmolarity and sodium that stimulates thirst and a compensatory increase in water intake. As a result, *hypernatremia and other overt physical or laboratory signs of dehydration do not develop unless the patient also has a defect in thirst or fails to increase fluid intake for some other reason.*

TABLE 404-1 CAUSES OF DIABETES INSIPIDUS

Pituitary diabetes insipidus	Gestational diabetes insipidus
Acquired	Pregnancy (second and third trimesters)
Head trauma (closed and penetrating) including pituitary surgery	Nephrogenic diabetes insipidus
Neoplasms	Acquired
Primary	Drugs
Craniopharyngioma	Lithium
Pituitary adenoma (suprasellar)	Demeclocycline
Dysgerminoma	Methoxyflurane
Meningioma	Amphotericin B
Metastatic (lung, breast)	Aminoglycosides
Hematologic (lymphoma, leukemia)	Cisplatin
Granulomas	Rifampin
Sarcoidosis	Foscarnet
Histiocytosis	Metabolic
Xanthoma disseminatum	Hypercalcemia, hypercalciuria
Infectious	Hypokalemia
Chronic meningitis	Obstruction (ureter or urethra)
Viral encephalitis	Vascular
Toxoplasmosis	Sickle cell disease and trait
Inflammatory	Ischemia (acute tubular necrosis)
Lymphocytic infundibuloneurohypophysitis	Granulomas
Granulomatosis with polyangiitis (Wegener's)	Sarcoidosis
Lupus erythematosus	Neoplasms
Scleroderma	Sarcoma
Chemical toxins	Infiltration
Tetrodotoxin	Amyloidosis
Snake venom	Idiopathic
Vascular	Genetic
Sheehan's syndrome	X-linked recessive (<i>AVP receptor-2 gene</i>)
Aneurysm (internal carotid)	Autosomal recessive (<i>AQP2 gene</i>)
Aortocoronary bypass	Autosomal dominant (<i>AQP2 gene</i>)
Hypoxic encephalopathy	Primary polydipsia
Idiopathic	Acquired
Congenital malformations	Psychogenic
Septo-optic dysplasia	Schizophrenia
Midline craniofacial defects	Obsessive compulsive disorder
Holoprosencephaly	Dipsogenic (abnormal thirst)
Hypogenesis, ectopia of pituitary	Granulomas (sarcoidosis)
Genetic	Infectious (tuberculous meningitis)
Autosomal dominant	Head trauma (closed and penetrating)
(<i>AVP-neurophysin gene</i>)	Demyelination (multiple sclerosis)
Autosomal recessive	Drugs
Type A (<i>AVP-neurophysin gene</i>)	Idiopathic
Type B (<i>AVP-neurophysin gene</i>)	Iatrogenic
Type C (<i>Wolfram's [4p-WFS 1] gene</i>)	
X-linked recessive (Xq28)	

In pituitary and nephrogenic DI, the severity of the defect in AVP secretion or action varies significantly from patient to patient. In some, the defect is so severe that it cannot be overcome by even an intense stimulus such as nausea or severe dehydration. In others, the defect in AVP secretion or action is incomplete, and a modest stimulus such as a few hours of fluid deprivation, smoking, or a vasovagal reaction can raise urine osmolality as high as 800 mosmol/L. However, even when the defects are partial, the relation of urine osmolality to plasma AVP in patients with nephrogenic DI (Fig. 404-3A) or of plasma AVP to plasma osmolality and sodium in patients with pituitary DI (Fig. 404-3B) is subnormal.

In primary polydipsia, the pathogenesis of the polydipsia and polyuria is the reverse of that in pituitary, nephrogenic, and gestational DI. In primary polydipsia, an abnormality in cognition or thirst

causes excessive intake of fluids and an increase in body water that reduces plasma osmolality/sodium, AVP secretion, and urinary concentration. Dilution of the urine, in turn, results in a compensatory increase in urinary free-water excretion that usually offsets the increase in intake and stabilizes plasma osmolality/sodium at a level only 1–2% below basal. Thus, hyponatremia or clinically appreciable overhydration is uncommon unless the polydipsia is very severe or the compensatory water diuresis is impaired by a drug or disease that stimulates or mimics the antidiuretic effect of endogenous AVP. A rise in plasma osmolality and sodium produced by fluid deprivation or hypertonic saline infusion increases plasma AVP normally. However, the resultant increase in urine concentration is often subnormal because polyuria per se temporarily reduces the capacity of the kidney to concentrate the urine. Thus, the maximum level of urine osmolality achieved during fluid deprivation is often indistinguishable from that in patients with partial pituitary or partial nephrogenic DI.

Differential Diagnosis When symptoms of urinary frequency, enuresis, nocturia, and/or persistent thirst are present in the absence of glucosuria, the possibility of DI should be evaluated by collecting a 24-h urine on ad libitum fluid intake. If the volume exceeds 50 mL/kg per day (3500 mL in a 70-kg male) and the osmolality is below 300 mosmol/L, DI is confirmed and the patient should be evaluated further to determine the type in order to select the appropriate therapy.

The type of DI can sometimes be inferred from the clinical setting or medical history. Often, however, such information is lacking, ambiguous, or misleading, and other approaches to differential diagnosis are needed. If basal plasma osmolality and sodium are within normal limits, the traditional approach is to determine the effect of fluid deprivation and injection of antidiuretic hormone on urine osmolality. This approach suffices for differential diagnosis if fluid deprivation raises plasma osmolality and sodium above the normal range without inducing concentration of the urine. In that event, primary polydipsia and partial defects in AVP secretion and action are excluded, and the effect on urine osmolality of injecting 2 µg of the AVP analogue, desmopressin, indicates whether the patient has severe pituitary DI or severe nephrogenic DI. However, this approach is of little or no diagnostic value if fluid deprivation results in concentration of the urine because the increases in urine osmolality achieved both before and after the injection of desmopressin are similar in patients with partial pituitary DI,

partial nephrogenic DI, and primary polydipsia. These disorders can be differentiated by measuring plasma AVP during fluid deprivation and relating it to the concurrent level of plasma and urine osmolality (Fig. 404-3). However, this approach does not always differentiate clearly between partial pituitary DI and primary polydipsia unless the measurement is made when plasma osmolality and sodium are at or above the normal range. This level is difficult to achieve by fluid deprivation alone once urinary concentration occurs. Therefore it is usually necessary to give a short infusion of 3% saline condition (0.1 mL/kg body weight per minute for 60 to 90 minutes) and repeat the measurement of plasma AVP.

A simpler but equally reliable way to differentiate between pituitary DI, nephrogenic DI, and primary polydipsia is to measure basal plasma

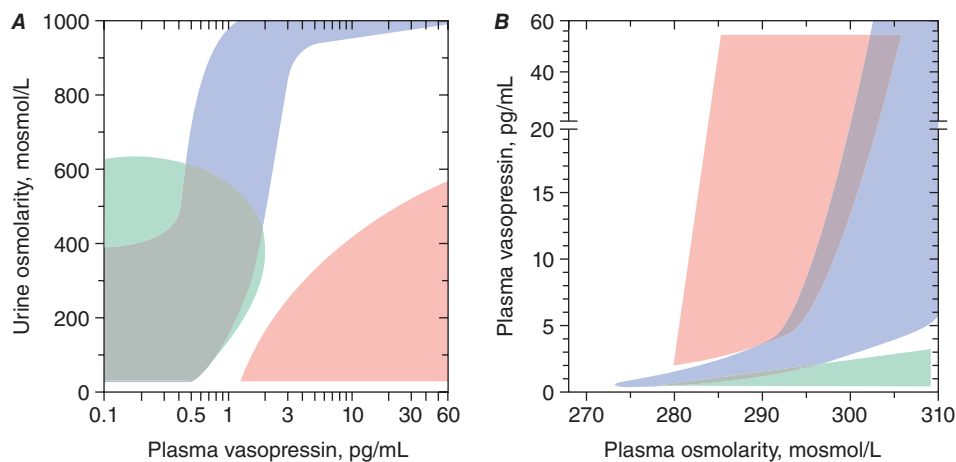


FIGURE 404-3 Relationship of plasma AVP to urine osmolarity (A) and plasma osmolarity (B) before and during fluid deprivation-hypertonic saline infusion test in patients who are normal or have primary polydipsia (blue zones), pituitary diabetes insipidus (green zones), or nephrogenic diabetes insipidus (pink zones).

AVP to determine if a brain magnetic resonance imaging (MRI) is needed and sufficient for diagnosis (Fig. 404-4). If plasma AVP on *ad libitum* fluid intake is normal or elevated (>1 pg/mL) when measured by a sensitive and specific assay, both primary polydipsia and pituitary DI are excluded and the diagnosis of nephrogenic DI can be confirmed, if desired, by a 1- to 2-day outpatient trial of desmopressin

therapy. If, however, basal plasma AVP is low or undetectable (<1 pg/mL), nephrogenic DI is very unlikely and MRI of the brain can be used to differentiate pituitary DI from primary polydipsia. In most healthy adults and children, the posterior pituitary emits a hyperintense signal visible in T1-weighted midsagittal images. This “bright spot” is almost always present in patients with primary polydipsia but is always absent or abnormally small in patients with pituitary DI, even if their AVP deficiency is partial. The MRI is also useful in searching for pathology responsible for pituitary DI or the dipsogenic form of primary polydipsia (Fig. 404-2). The principal caveat is that MRI is not reliable for differential diagnosis of DI in patients with empty sella because they typically lack a bright spot even when their AVP secretion and action are normal. MRI also cannot be used to differentiate pituitary from nephrogenic DI because many patients with nephrogenic DI also lack a posterior pituitary bright spot, probably because they have an abnormally high rate of AVP secretion and turnover.

If MRI and/or AVP assays with the requisite sensitivity and specificity are unavailable and a fluid deprivation test is impractical or undesirable, a third way to differentiate between pituitary DI, nephrogenic DI, and primary polydipsia is a trial of desmopressin therapy. Such a trial should be conducted with very close monitoring of serum sodium as well as urine output, preferably in hospital, because desmopressin will produce hyponatremia in 8–24 h if the patient has primary polydipsia.

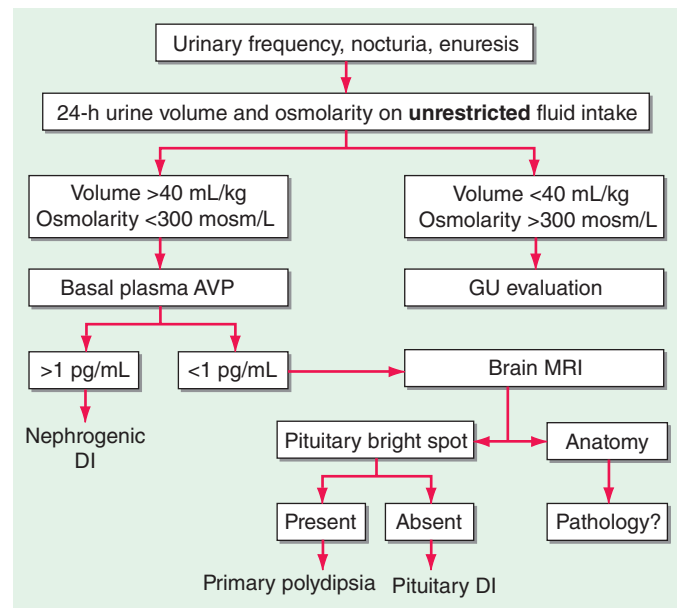


FIGURE 404-4 Simplified approach to the differential diagnosis of diabetes insipidus. When symptoms suggest diabetes insipidus (DI), the syndrome should be differentiated from a genitourinary (GU) abnormality by measuring the 24-h urine volume and osmolarity on unrestricted fluid intake. If DI is confirmed, basal plasma arginine vasopressin (AVP) should be measured on unrestricted fluid intake. If AVP is normal or elevated (>1 pg/mL), the patient probably has nephrogenic DI. However, if plasma AVP is low or undetectable, the patient has either pituitary DI or primary polydipsia. In that case, magnetic resonance imaging (MRI) of the brain can be performed to differentiate between these two conditions by determining whether or not the normal posterior pituitary bright spot is visible on T1-weighted midsagittal images. In addition, the MRI anatomy of the pituitary hypothalamic area can be examined to look for evidence of pathology that sometimes causes pituitary DI or the dipsogenic form of primary polydipsia. MRI is not reliable for differential diagnosis unless nephrogenic DI has been excluded because the bright spot is also absent, small, or faint in this condition.

TREATMENT DIABETES INSIPIDUS

The signs and symptoms of uncomplicated pituitary DI can be eliminated by treatment with desmopressin (DDAVP), a synthetic analogue of AVP (Fig. 404-1). DDAVP acts selectively at V₂ receptors to increase urine concentration and decrease urine flow in a dose-dependent manner. It is also more resistant to degradation than is AVP and has a three- to fourfold longer duration of action. DDAVP can be given by IV or SC injection, nasal inhalation, or orally by means of a tablet of melt. The doses required to control pituitary DI completely vary widely, depending on the patient and the route of administration. However, among adults, they usually range from 1–2 μg qd or bid by injection, 10–20 μg bid or tid by nasal spray, or 100–400 μg bid or tid orally. The onset of antidiuresis is rapid, ranging from as little as 15 min after injection to 60 min after oral administration. When given in a dose that normalizes 24-h urinary osmolarity (400–800 mosmol/L) and volume (15–30 mL/kg body weight), DDAVP produces a slight (1–3%) increase in total body water and a decrease in plasma osmolarity/sodium that rapidly eliminates thirst and polydipsia (Fig. 404-5). Consequently, water balance is maintained within the normal range. Hyponatremia does not develop unless urine volume is reduced too far (to less than 10 mL/kg per day) or fluid intake is excessive due to an associated

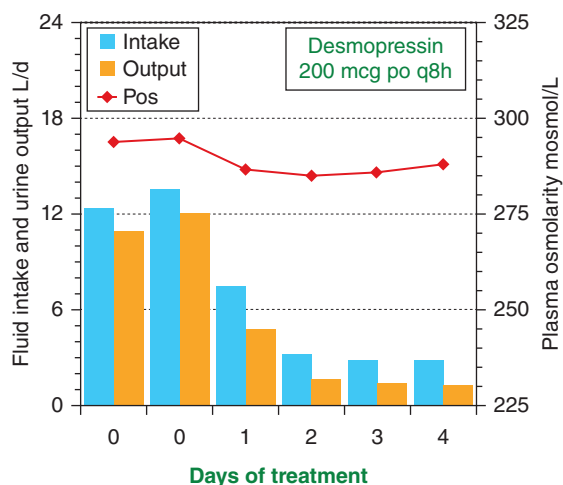


FIGURE 404-5 Effect of desmopressin therapy on fluid intake (blue bars), urine output (orange bars), and plasma osmolarity (red line) in a patient with uncomplicated pituitary diabetes insipidus. Note that treatment rapidly reduces fluid intake and urine output to normal, with only a slight increase in body water as evidenced by the slight decrease in plasma osmolarity.

abnormality in thirst or cognition. Fortunately, thirst abnormalities are rare, and if the patient is taught to drink only when truly thirsty, DDAVP can be given safely in doses sufficient to normalize urine output (~15–30 mL/kg per day) without the need for allowing intermittent escape to prevent water intoxication.

Primary polydipsia cannot be treated safely with DDAVP or any other antidiuretic drug because eliminating the polyuria does not eliminate the urge to drink. Therefore, it invariably produces hyponatremia and/or other signs of water intoxication, usually within 8–24 h if urine output is normalized completely. There is no consistently effective way to correct dipsogenic or psychogenic polydipsia, but the iatrogenic form may respond to patient education. To minimize the risk of water intoxication, all patients should be warned about the use of other drugs such as thiazide diuretics or carbamazepine (Tegretol) that can impair urinary free-water excretion directly or indirectly.

The polyuria and polydipsia of nephrogenic DI are not affected by treatment with standard doses of DDAVP. If resistance is partial, it may be overcome by tenfold higher doses, but this treatment is too expensive and inconvenient for long-term use. However, treatment with conventional doses of a thiazide diuretic and/or amiloride in conjunction with a low-sodium diet and a prostaglandin synthesis inhibitor (e.g., indomethacin) usually reduces the polyuria and polydipsia by 30–70% and may eliminate them completely in some patients. Side effects such as hypokalemia and gastric irritation can be minimized by the use of amiloride or potassium supplements and by taking medications with meals.

HYPODIPSIC HYPERNATREMIA

An increase in plasma osmolarity/sodium above the normal range (hypertonic hypernatremia) can be caused by either a decrease in total body water or an increase in total body sodium. The former results from a failure to drink enough to replace normal or increased urinary and insensible water loss. The deficient intake can be due either to water deprivation or a lack of thirst (hypodipsia). The most common cause of an increase in total body sodium is primary hyperaldosteronism (Chap. 406). Rarely, it can also result from ingestion of hypertonic saline in the form of sea water or incorrectly prepared infant formula. However, even in these forms of hypernatremia, inadequate intake of water also contributes. This chapter focuses on hypodipsic hypernatremia, the form of hypernatremia due to a primary defect in the thirst mechanism.

Clinical Characteristics Hypodipsic hypernatremia is a syndrome characterized by chronic or recurrent hypertonic dehydration. The hypernatremia varies widely in severity and usually is associated with signs

of hypovolemia such as tachycardia, postural hypotension, azotemia, hyperuricemia, and hypokalemia due to secondary hyperaldosteronism. Muscle weakness, pain, rhabdomyolysis, hyperglycemia, hyperlipidemia, and acute renal failure may also occur. Obtundation or coma may be present but are often absent. Despite inappropriately low levels of plasma AVP, DI usually is not evident at presentation but may develop during rehydration as blood volume, blood pressure, and plasma osmolarity/sodium return toward normal, further reducing plasma AVP.

Etiology Hypodipsia is usually due to hypogenesis or destruction of the osmoreceptors in the anterior hypothalamus that regulate thirst. These defects can result from various congenital malformations of midline brain structures or may be acquired due to diseases such as occlusions of the anterior communicating artery, primary or metastatic tumors in the hypothalamus, head trauma, surgery, granulomatous diseases such as sarcoidosis and histiocytosis, AIDS, and cytomegalovirus encephalitis. Because of their proximity, the osmoreceptors that regulate AVP secretion also are usually impaired. Thus, AVP secretion responds poorly or not at all to hyperosmotic stimulation (Fig. 404-6) but, in most cases, increases normally to nonosmotic

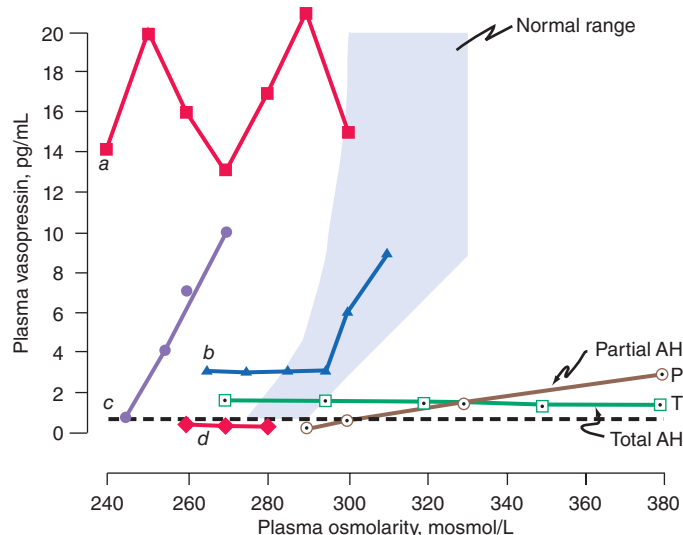


FIGURE 404-6 Heterogeneity of osmoregulatory dysfunction in adipsic hypernatremia (AH) and the syndrome of inappropriate antidiuresis (SIAD). Each line depicts schematically the relationship of plasma arginine vasopressin (AVP) to plasma osmolarity during water loading and/or infusion of 3% saline in a patient with either AH (open symbols) or SIAD (closed symbols). The shaded area indicates the normal range of the relationship. The horizontal broken line indicates the plasma AVP level below which the hormone is undetectable and urinary concentration usually does not occur. Lines P and T represent patients with a selective deficiency in the osmoregulation of thirst and AVP that is either partial (○) or total (□). In the latter, plasma AVP does not change in response to increases or decreases in plasma osmolarity but remains within a range sufficient to concentrate the urine even if overhydration produces hypotonic hyponatremia. In contrast, if the osmoregulatory deficiency is partial (○), rehydration of the patient suppresses plasma AVP to levels that result in urinary dilution and polyuria before plasma osmolarity and sodium are reduced to normal. Lines a–d represent different defects in the osmoregulation of plasma AVP observed in patients with SIADH or SIAD. In a (■), plasma AVP is markedly elevated and fluctuates widely without relation to changes in plasma osmolarity, indicating complete loss of osmoregulation. In b (▲), plasma AVP remains fixed at a slightly elevated level until plasma osmolarity reaches the normal range, at which point it begins to rise appropriately, indicating a selective SIAD defect in the inhibitory component of the osmoregulatory mechanism. In c (●), plasma AVP rises in close correlation with plasma osmolarity before the latter reaches the normal range, indicating downward resetting of the osmostat. In d (◆), plasma AVP appears to be osmoregulated normally, suggesting that the inappropriate antidiuresis is caused by some other abnormality.

2280 stimuli such as nausea or large reductions in blood volume or blood pressure, indicating that the neurohypophysis is intact.

Pathophysiology Hypodipsia results in a failure to drink enough water to replenish obligatory renal and extrarenal losses. Consequently, plasma osmolality and sodium rise often to extremely high levels before the disorder is recognized. In most cases, urinary loss of water contributes little, if any, to the dehydration because AVP continues to be secreted in the small amounts necessary to concentrate the urine. In some patients this appears to be due to hypovolemic stimulation and/or incomplete destruction of AVP osmoreceptors because plasma AVP declines and DI develops during rehydration (Fig. 404-6). In others, however, plasma AVP does not decline during rehydration even if they are overhydrated. Consequently, they develop a hyponatremic syndrome indistinguishable from inappropriate antidiuresis. This suggests that the AVP osmoreceptors normally provide inhibitory and stimulatory input to the neurohypophysis and the patients can no longer osmotically stimulate or suppress tonic secretion of the hormone because both inputs have been totally eliminated by the same pathology that destroyed the osmoregulation of thirst. In a few patients, the neurohypophysis is also destroyed, resulting in a combination of chronic pituitary DI and hypodipsia that is particularly difficult to manage.

Differential Diagnosis Hypodipsic hypernatremia usually can be distinguished from other causes of inadequate fluid intake (e.g., coma, paralysis, restraints, absence of fresh water) by the clinical history and setting. Previous episodes and/or denial of thirst and failure to drink spontaneously when the patient is conscious, unrestrained, and hypernatremic are virtually diagnostic. The hypernatremia caused by excessive retention or intake of sodium can be distinguished by the presence of thirst as well as the physical and laboratory signs of hypervolemia rather than hypovolemia.

TREATMENT HYPODIPSIC HYPERNATREMIA

Hypodipsic hypernatremia should be treated by administering water orally if the patient is alert and cooperative or by infusing hypotonic fluids (0.45% saline or 5% dextrose and water) if the patient is not. The amount of free water in liters required to correct the deficit (ΔFW) can be estimated from body weight in kg (BW) and the serum sodium concentration in mmol/L (S_{Na}) by the formula $\Delta FW = 0.5BW \times ((S_{Na} - 140)/140)$. If serum glucose (S_{Glu}) is elevated, the measured S_{Na} should be corrected (S_{Na}^*) by the formula $S_{Na}^* = S_{Na} + ((S_{Glu} - 90)/36)$. This amount plus an allowance for continuing insensible and urinary losses should be given over a 24- to 48-h period. Close monitoring of serum sodium as well as fluid intake and urinary output is essential because, depending on the extent of osmoreceptor deficiency, some patients will develop AVP-deficient DI, requiring DDAVP therapy to complete rehydration; others will develop hyponatremia and a syndrome of inappropriate antidiuresis (SIAD)-like picture if overhydrated. If hyperglycemia and/or hypokalemia are present, insulin and/or potassium supplements should be given with the expectation that both can be discontinued soon after rehydration is complete. Plasma urea/creatinine should be monitored closely for signs of acute renal failure caused by rhabdomyolysis, hypovolemia, and hypotension.

Once the patient has been rehydrated, an MRI of the brain and tests of anterior pituitary function should be performed to look for the cause and collateral defects in other hypothalamic functions. A long-term management plan to prevent or minimize recurrence of the fluid and electrolyte imbalance also should be developed. This should include a practical method to regulate fluid intake in accordance with variations in water balance as indicated by changes in body weight or serum sodium determined by home monitoring analyzers. Prescribing a constant fluid intake is ineffective and potentially dangerous because it does not take into account the large, uncontrolled variations in insensible loss that inevitably result from changes in ambient temperature and physical activity.

HYPONATREMIA DUE TO INAPPROPRIATE ANTIDIURESIS

A decrease in plasma osmolality/sodium below the normal range (hypotonic hyponatremia) can be due to any of three different types of salt and water imbalance: (1) an increase in total body water that exceeds the increase in total body sodium (hypervolemic hyponatremia); (2) a decrease in body sodium greater than the decrease in body water (hypovolemic hyponatremia); or (3) an increase in body water with little or no change in body sodium (euvolemic hyponatremia) (Chap. 63). All three forms are associated with a failure to fully dilute the urine and mount a water diuresis in the face of hypotonic hyponatremia. The hypervolemic form typically occurs in disorders like severe congestive heart failure or cirrhosis. The hypovolemic form typically occurs in disorders such as severe diarrhea, diuretic abuse, or mineralocorticoid deficiency. Euvolemic hyponatremia, however, is due mainly to expansion of total body water caused by excessive intake in the face of a defect in urinary dilution. The impaired dilution is usually caused by a defect in the osmotic suppression of AVP that can have either of two causes. One is a nonhemodynamic stimulus such as nausea or a cortisol deficiency, which can be corrected quickly by treatment with antiemetics or cortisol. The other is a primary defect in osmoregulation caused by another disorder such as malignancy, stroke, or pneumonia that cannot be easily or quickly corrected. The latter is commonly known as the syndrome of inappropriate antidiuretic hormone (SIADH). Much less often, euvolemic hyponatremia can also result from AVP-independent activation of renal V_2 receptors, a variant known as nephrogenic inappropriate antidiuresis or NSIAD. Both of the latter will be discussed in this chapter.

Clinical Characteristics Antidiuresis of any cause decreases the volume and increases the concentration of urine. If not accompanied by a commensurate reduction in fluid intake or an increase in insensible loss, the reduction in urine output results in excess water retention which expands and dilutes body fluids. If the hyponatremia develops gradually or has been present for more than a few days, it may be largely asymptomatic. However, if it develops acutely, it is usually accompanied by symptoms and signs of water intoxication that may include mild headache, confusion, anorexia, nausea, vomiting, coma, and convulsions. Severe acute hyponatremia may be lethal. Other clinical signs and symptoms vary greatly, depending on the type of hyponatremia. The hypervolemic form is characterized by generalized edema and other signs of marked volume expansion. The opposite is evident in the hypovolemic form. However, overt signs of volume expansion or contraction are absent in SIADH, SIAD, and other forms of euvolemic hyponatremia.

Etiology In SIADH, the inappropriate secretion of AVP can have many different causes. They include ectopic production of AVP by lung cancer or other neoplasms; eutopic release induced by various diseases or drugs; and exogenous administration of AVP, DDAVP, or large doses of oxytocin (Table 404-2). The ectopic forms result from abnormal expression of the *AVP-NP11* gene by primary or metastatic malignancies. The eutopic forms occur most often in patients with acute infections or strokes but have also been associated with many other neurologic diseases and injuries. The mechanisms by which these diseases interfere with osmotic suppression of AVP are not known. The defect in osmoregulation can take any of four distinct forms (Fig. 404-6). In one of the most common (reset osmostat), AVP secretion remains fully responsive to changes in plasma osmolality/sodium, but the threshold, or set point, of the osmoregulatory system is abnormally low. These patients differ from those with the other types of SIADH in that they are able to maximally suppress plasma AVP and dilute their urine if their fluid intake is high enough to reduce their plasma osmolality and/or sodium to the new set point. In most patients, SIADH is self-limited and remits spontaneously within 2–3 weeks, but about 10% of cases are chronic. Another, smaller subgroup (~10% of the total) has inappropriate antidiuresis without a demonstrable defect in the osmoregulation of plasma AVP (Fig. 404-6). In some of them, all young boys, the inappropriate antidiuresis has been traced to a constitutively activating mutation of the V_2 receptor gene. This unusual variant may be referred to as familial nephrogenic SIAD (NSIAD) to distinguish it from

TABLE 404-2 CAUSES OF SYNDROME OF INAPPROPRIATE ANTIURETIC HORMONE (SIADH)

Neoplasms	Neurologic
Carcinomas	Guillain-Barré syndrome
Lung	Multiple sclerosis
Duodenum	Delirium tremens
Pancreas	Amyotrophic lateral sclerosis
Ovary	Hydrocephalus
Bladder, ureter	Psychosis
Other neoplasms	Peripheral neuropathy
Thymoma	Congenital malformations
Mesothelioma	Agenesis corpus callosum
Bronchial adenoma	Cleft lip/palate
Carcinoid	Other midline defects
Gangliocytoma	Metabolic
Ewing's sarcoma	Acute intermittent porphyria
Head trauma (closed and penetrating)	Pulmonary
Infections	Asthma
Pneumonia, bacterial or viral	Pneumothorax
Abscess, lung or brain	Positive-pressure respiration
Cavitation (aspergillosis)	Drugs
Tuberculosis, lung or brain	Vasopressin or desmopressin
Meningitis, bacterial or viral	Serotonin reuptake inhibitors
Encephalitis	Oxytocin, high dose
AIDS	Vincristine
Vascular	Carbamazepine
Cerebrovascular occlusions, hemorrhage	Nicotine
Cavernous sinus thrombosis	Phenothiazines
	Cyclophosphamide
	Tricyclic antidepressants
	Monoamine oxidase inhibitors

other possible causes of the syndrome. The inappropriate antidiuresis in these patients appears to be permanent, although the hyponatremia is variable owing presumably to individual differences in fluid intake.

Pathophysiology Impaired osmotic suppression of antidiuresis results in excessive retention of water and dilution of body fluids only if water intake exceeds insensible and urinary losses. The excess intake is sometimes due to an associated defect in the osmoregulation of thirst (dipsogenic) but can also be psychogenic or iatrogenic, including excessive IV administration of hypotonic fluids. In SIADH and other forms of euvolemic hyponatremia, the decrease in plasma osmolarity/sodium and the increase in extracellular and intracellular volume are proportional to the amount of water retained. Thus, an increase in body water of 10% (~4 L in a 70-kg adult) reduces plasma osmolarity and sodium by approximately 10% (~28 mosmol/L or 14 meq/L). An increase in body water of this magnitude is rarely detectable on physical examination but will be reflected in a weight gain of about 4 kg. It also increases glomerular filtration and atrial natriuretic hormone and suppresses plasma renin activity, thereby increasing urinary sodium excretion. The resultant reduction in total body sodium decreases the expansion of extracellular volume but aggravates the hyponatremia and further expands intracellular volume. The latter further increases brain swelling and intracranial pressure, which probably produces most of the symptoms of acute water intoxication. Within a few days, this swelling may be counteracted by inactivation or elimination of intracellular solutes, resulting in the remission of symptoms even though the hyponatremia persists.

In type I (hypervolemic) or type II (hypovolemic) hyponatremia, osmotic suppression of AVP secretion appears to be counteracted by a hemodynamic stimulus resulting from a large reduction in cardiac output and/or effective blood volume. The resultant antidiuresis is enhanced by decreased distal delivery of glomerular filtrate that

results from increased reabsorption of sodium in proximal nephron. If the reduction in urine output is not associated with a commensurate reduction in water intake or an increase in insensible loss, body fluids are expanded and diluted, resulting in hyponatremia despite an increase in body sodium. Unlike SIADH and other forms of euvolemic hyponatremia, however, glomerular filtration is reduced and plasma renin activity and aldosterone are elevated. Thus, the rate of urinary sodium excretion is low (unless sodium reabsorption is impaired by a diuretic), and the hyponatremia is usually accompanied by edema, hypokalemia, azotemia, and hyperuricemia. In type II (hypovolemic) hyponatremia, sodium and water are also retained as an appropriate compensatory response to the severe depletion.

Differential Diagnosis SIADH is a diagnosis of exclusion that usually can be made from the history, physical examination, and basic laboratory data. If hyperglycemia is present, its contribution to the reduction in plasma sodium can be estimated either by measuring plasma osmolarity for a more accurate estimate of the true "effective" tonicity of body fluids or by correcting the measured plasma sodium for the reduction caused by the hyperglycemia using the simplified formula

$$\text{corrected } P_{\text{na}} = \text{measured } P_{\text{na}} + (P_{\text{glu}} - 90)/36$$

where P_{na} = plasma sodium in meq/L and P_{glu} = plasma glucose in mg/dL.

If the plasma osmolarity and/or corrected plasma sodium are below normal limits, hypotonic hyponatremia is present and further evaluation to determine the type should be undertaken in order to administer safe and effective treatment. This differentiation is usually possible by evaluating standard clinical indicators of the extracellular fluid volume (Table 404-3). If these findings are ambiguous or contradictory, measuring plasma renin activity or the rate of urinary sodium excretion may be helpful provided that the hyponatremia is not in the recovery phase or is due to a primary defect in renal conservation of sodium, diuretic abuse, or hyporeninemic hypoaldosteronism. The latter may be suspected if serum potassium is elevated instead of low, as it usually is in types I and II hyponatremia. Measurements of plasma AVP are currently of no value in differentiating SIADH from the other types of hyponatremia since the plasma levels are elevated similarly in all. In patients who fulfill the clinical criteria for type III (euvolemic) hyponatremia, morning plasma cortisol should also be measured to exclude secondary adrenal insufficiency. If it is normal and there is no history of nausea/vomiting, the diagnosis of SIADH is confirmed, and a careful search for occult lung cancer or other common causes of the syndrome (Table 404-2) should be undertaken.

SIAD due to an activating mutation of the V_2 receptor gene should be suspected if the hyponatremia occurs in a child or several members of the family or is refractory to treatment with a vaptan (see below). In that case, plasma AVP should be measured to confirm that it is appropriately suppressed while the hyponatremia and antidiuresis are present, and the V_2 receptor gene should be sequenced, if possible.

TREATMENT HYPONATREMIA

The management of hyponatremia differs depending on the type and the severity and duration of symptoms. In acute symptomatic SIADH, the aim should be to raise plasma osmolarity and/or plasma sodium at a rate approximating 1% an hour until they reach levels of about 270 mosmol/L or 130 meq/L, respectively. This can be accomplished in either of two ways. One is to infuse hypertonic (3%) saline at a rate of about 0.05 mL/kg body weight per minute. This treatment also has the advantage of correcting the sodium deficiency that is partly responsible for the hyponatremia and often produces a solute diuresis that serves to remove some of the excess water. The other treatment is to reduce body water by giving an AVP receptor-2 antagonist (vaptan) to block the antidiuretic effect of AVP and increase urine output (Fig. 404-7). One of the vaptans, a combined V_2/V_{1a} antagonist (Conivaptan), has been approved for short-term, in-hospital IV treatment of SIADH, and others are in various stages of development. With either approach, fluid intake should be restricted to less than urine

TABLE 404-3 DIFFERENTIAL DIAGNOSIS OF HYPONATREMIA BASED ON CLINICAL ASSESSMENT OF EXTRACELLULAR FLUID VOLUME (ECFV)

Clinical Findings	Type I, Hypervolemic	Type II, Hypovolemic	Type III, Euvolemic	SIADH and SIAD Euvolemic
History				
CHF, cirrhosis, or nephrosis	Yes	No	No	No
Salt and water loss	No	Yes	No	No
ACTH–cortisol deficiency and/or nausea and vomiting	No	No	Yes	No
Physical examination				
Generalized edema, ascites	Yes	No	No	No
Postural hypotension	Maybe	Maybe	Maybe ^a	No
Laboratory				
BUN, creatinine	High-normal	High-normal	Low-normal	Low-normal
Uric acid	High-normal	High-normal	Low-normal	Low-normal
Serum potassium	Low-normal	Low-normal ^b	Normal ^c	Normal
Serum urate	High	High	Low	Low
Serum albumin	Low-normal	High-normal	Normal	Normal
Serum cortisol	Normal-high	Normal-high ^d	Low ^e	Normal
Plasma renin activity	High	High	Low ^f	Low
Urinary sodium (meq per unit of time) ^g	Low	Low ^h	High ⁱ	High ⁱ

^aPostural hypotension may occur in secondary (ACTH-dependent) adrenal insufficiency even though extracellular fluid volume and aldosterone are usually normal. ^bSerum potassium may be high if hypovolemia is due to aldosterone deficiency. ^cSerum potassium may be low if vomiting causes alkalosis. ^dSerum cortisol is low if hypovolemia is due to primary adrenal insufficiency (Addison's disease). ^eSerum cortisol will be normal or high if the cause is nausea and vomiting rather than secondary (ACTH-dependent) adrenal insufficiency. ^fPlasma renin activity may be high if the cause is secondary (ACTH) adrenal insufficiency. ^gUrinary sodium should be expressed as the *rate of excretion* rather than the concentration. In a hyponatremic adult, an excretion rate >25 meq/d (or 25 μ eq/mg of creatinine) could be considered high. ^hThe rate of urinary sodium excretion may be high if the hypovolemia is due to diuretic abuse, primary adrenal insufficiency, or other causes of renal sodium wasting. ⁱThe rate of urinary sodium excretion may be low if intake is curtailed by symptoms or treatment.

Abbreviations: ACTH, adrenocorticotropic hormone; BUN, blood urea nitrogen; CHF, congestive heart failure; SIAD, syndrome of inappropriate antidiuresis.

output, and serum sodium should be checked at least once every 2h to ensure it is not raised too fast or too far. Doing so may result in central pontine myelinolysis, an acute, potentially fatal neurologic syndrome characterized by quadriplegia, ataxia, and abnormal extraocular movements.

In chronic and/or minimally symptomatic SIADH, the hyponatremia can and should be corrected more gradually. This can be achieved by restricting total fluid intake to less than the sum of urinary and insensible losses. Because the water derived from food

(300–700 mL/d) usually approximates basal insensible losses in adults, the aim should be to reduce total discretionary intake (all liquids) to approximately 500 mL less than urinary output. Adherence to this regimen is often problematic and, even if achieved, usually reduces body water and increases serum sodium by only about 1–2% per day. Hence, additional approaches are usually desirable if not necessary. The best approach for treatment of chronic SIADH is the administration of an oral vaptan, tolvaptan, a selective V_2 antagonist that also increases urinary water excretion by blocking the antidiuretic effect of AVP. Some restriction of fluid intake may also be necessary to achieve satisfactory control of the hyponatremia. It is approved for treatment of nonemergent SIADH with initial in-hospital dosing. Other approaches include demeclocycline, 150–300 mg PO tid or qid, or fludrocortisone, 0.05–0.2 mg PO bid. The effect of the demeclocycline manifests in 7–14 days and is due to induction of a reversible form of nephrogenic DI. Potential side effects include phototoxicity and azotemia. The effect of fludrocortisone also requires 1–2 weeks and is partly due to increased retention of sodium and possibly inhibition of thirst. It also increases urinary potassium excretion, which may require replacement through dietary adjustments or supplements and may induce hypertension, occasionally necessitating discontinuation of the treatment.

In euvolemic hyponatremia caused by protracted nausea and vomiting or isolated glucocorticoid deficiency (type III), all abnormalities can be corrected quickly and completely by giving an antiemetic or stress doses of hydrocortisone (for glucocorticoid deficiency). As with other treatments, care must be taken to ensure that serum sodium does not rise too quickly or too far.

In SIAD due to an activating mutation of the V_2 receptor, the V_2 antagonists usually do not block the antidiuresis or raise plasma osmolality/sodium. In that condition, use of an osmotic diuretic such as urea is reported to be effective in preventing or correcting hyponatremia. However, some vaptans may be effective in patients with a different type of activating mutation so the response to this therapy may be neither predictable nor diagnostic.

In hypervolemic hyponatremia, fluid restriction is also appropriate and somewhat effective if it can be maintained. However,

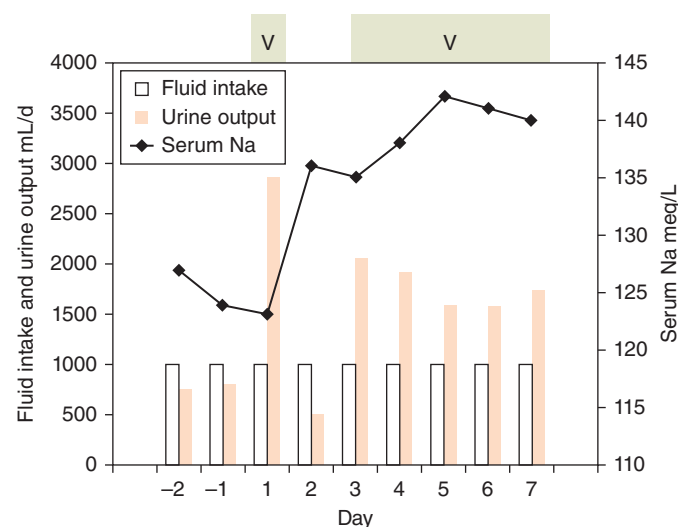


FIGURE 404-7 The effect of vaptan therapy on water balance in a patient with chronic syndrome of inappropriate antidiuretic hormone (SIADH). The periods of vaptan (V) therapy are indicated by the green shaded boxes at the top. Urine output is indicated by orange bars. Fluid intake is shown by the open bars. Intake was restricted to 1 L/d throughout. Serum sodium is indicated by the black line. Note that sodium increased progressively when vaptan increased urine output to levels that clearly exceeded fluid intake.

infusion of hypertonic saline is contraindicated because it further increases total body sodium and edema and may precipitate cardiovascular decompensation. However, as in SIADH, the V_2 receptor antagonists are also safe and effective in the treatment of hypervolemic hyponatremia caused by congestive heart failure. Tolvaptan is approved by the Food and Drug Administration for this indication with the caveat that treatment should be initiated or reinitiated in hospital. Its use should also be limited to 30 days at a time because of reports that longer periods may be associated with abnormal liver chemistries.

In hypovolemic hyponatremia, the defect in AVP secretion and water balance usually can be corrected easily and quickly by stopping the loss of sodium and water and/or replacing the deficits by mouth or IV infusion of normal or hypertonic saline. As with the treatment of other forms of hyponatremia, care must be taken to ensure that plasma sodium does not increase too rapidly or too far. Fluid restriction and administration of AVP antagonists are contraindicated in type II hyponatremia because they would only aggravate the underlying volume depletion and could result in hemodynamic collapse.

GLOBAL PERSPECTIVES



The incidence, clinical characteristics, etiology, pathophysiology, differential diagnosis, and treatments of fluid and electrolyte disorders in tropical and nonindustrialized countries differ in some respects from those in the United States and other industrialized parts of the world. Hyponatremia, for example, appears to be more common and is more likely to be due to infectious diseases such as cholera, shigellosis, and other diarrheal disorders. In these circumstances, hyponatremia is probably due to gastrointestinal losses of salt and water (hypovolemia type II), but other abnormalities, including undefined infectious toxins, also may contribute. The causes of DI are similar worldwide except that malaria and venoms from snake or insect bites are much more common.

405 Disorders of the Thyroid Gland

J. Larry Jameson, Susan J. Mandel, Anthony P. Weetman

The thyroid gland produces two related hormones, thyroxine (T_4) and triiodothyronine (T_3) (Fig. 405-1). Acting through thyroid hormone receptors α and β , these hormones play a critical role in cell differentiation during development and help maintain thermogenic and metabolic homeostasis in the adult. Autoimmune disorders of the thyroid gland can stimulate overproduction of thyroid hormones (*thyrotoxicosis*) or cause glandular destruction and hormone deficiency (*hypothyroidism*). In addition, benign nodules and various forms of thyroid cancer are relatively common and amenable to detection by physical examination.

ANATOMY AND DEVELOPMENT

The thyroid (Greek *thyreos*, shield, plus *eidosis*, form) consists of two lobes connected by an isthmus. It is located anterior to the trachea between the cricoid cartilage and the suprasternal notch. The normal thyroid is 12–20 g in size, highly vascular, and soft in consistency. Four parathyroid glands, which produce parathyroid hormone (Chap. 424), are located posterior to each pole of the thyroid. The recurrent laryngeal nerves traverse the lateral borders of the thyroid gland and must be identified during thyroid surgery to avoid injury and vocal cord paralysis.

The thyroid gland develops from the floor of the primitive pharynx during the third week of gestation. The developing gland migrates along the thyroglossal duct to reach its final location in the neck. This feature accounts for the rare ectopic location of thyroid tissue at the base of the tongue (lingual thyroid) as well as the occurrence of thyroglossal duct cysts along this developmental tract. Thyroid hormone synthesis normally begins at about 11 weeks' gestation.

Neural crest derivatives from the ultimobranchial body give rise to thyroid medullary C cells that produce calcitonin, a calcium-lowering hormone. The C cells are interspersed throughout the thyroid gland, although their density is greatest in the juncture of the upper one-third and lower two-thirds of the gland. Calcitonin plays a minimal role in calcium homeostasis in humans but the C-cells are important because of their involvement in medullary thyroid cancer.

Thyroid gland development is orchestrated by the coordinated expression of several developmental transcription factors. Thyroid transcription factor (TTF)-1, TTF-2, and paired homeobox-8 (PAX-8) are expressed selectively, but not exclusively, in the thyroid gland. In combination, they dictate thyroid cell development and the induction of thyroid-specific genes such as thyroglobulin (Tg), thyroid peroxidase (TPO), the sodium iodide symporter (Na^+/I^- , NIS), and the thyroid-stimulating hormone receptor (TSH-R). Mutations in these developmental transcription factors or their downstream target genes are rare causes of thyroid agenesis or dyshormonogenesis, although the causes of most forms of congenital hypothyroidism remain unknown (Table 405-1). Because congenital hypothyroidism occurs in approximately 1 in 4000 newborns, neonatal screening is now performed in most industrialized countries (see below). Transplacental passage of maternal thyroid hormone occurs before the fetal thyroid gland begins to function and provides partial hormone support to a fetus with congenital hypothyroidism. Early thyroid hormone replacement in newborns with congenital hypothyroidism prevents potentially severe developmental abnormalities.

The thyroid gland consists of numerous spherical follicles composed of thyroid follicular cells that surround secreted colloid, a proteinaceous fluid containing large amounts of thyroglobulin, the protein precursor of thyroid hormones (Fig. 405-2). The thyroid follicular cells are polarized—the basolateral surface is apposed to the bloodstream and an apical surface faces the follicular lumen. Increased demand for thyroid hormone is regulated by thyroid-stimulating hormone (TSH), which binds to its receptor on the basolateral surface of the follicular cells. This binding leads to Tg reabsorption from the follicular lumen and proteolysis within the cytoplasm, yielding thyroid hormones for secretion into the bloodstream.

REGULATION OF THE THYROID AXIS

TSH, secreted by the thyrotrope cells of the anterior pituitary, plays a pivotal role in control of the thyroid axis and serves as the most useful

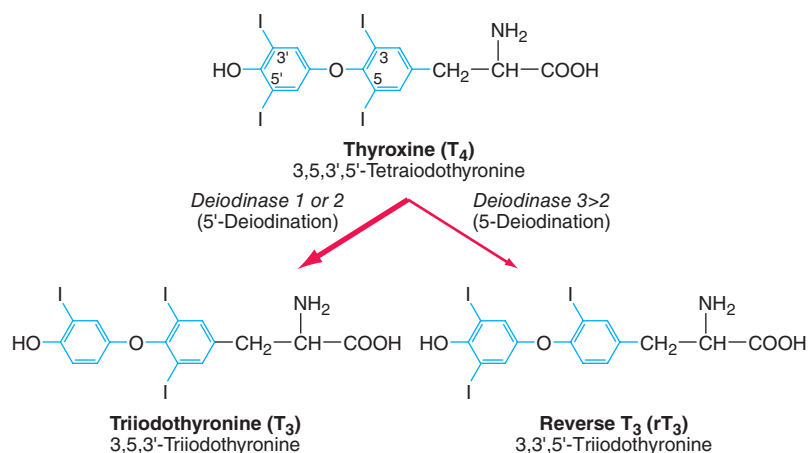


FIGURE 405-1 Structures of thyroid hormones. Thyroxine (T_4) contains four iodine atoms. Deiodination leads to production of the potent hormone triiodothyronine (T_3) or the inactive hormone reverse T_3 .

Defective Gene Protein	Inheritance	Consequences
PROP-1	Autosomal recessive	Combined pituitary hormone deficiencies with preservation of adrenocorticotropic hormone
PIT-1	Autosomal recessive Autosomal dominant	Combined deficiencies of growth hormone, prolactin, thyroid-stimulating hormone (TSH)
TSH β	Autosomal recessive	TSH deficiency
TTF-1 (TITF-1)	Autosomal dominant	Variable thyroid hypoplasia, cho- reoathetosis, pulmonary problems
TTF-2 (FOXE-1)	Autosomal recessive	Thyroid agenesis, choanal atresia, spiky hair
PAX-8	Autosomal dominant	Thyroid dysgenesis
TSH-receptor	Autosomal recessive	Resistance to TSH
G _{3α} (Albright hereditary osteodystrophy)	Autosomal dominant	Resistance to TSH
Na ⁺ /I ⁻ symporter	Autosomal recessive	Inability to transport iodide
DUOX2 (THOX2)	Autosomal dominant	Organification defect
DUOXA2	Autosomal recessive	Organification defect
Thyroid peroxidase	Autosomal recessive	Defective organification of iodide
Thyroglobulin	Autosomal recessive	Defective synthesis of thyroid hormone
Pendrin	Autosomal recessive	Pendred syndrome: sensorineural deafness and partial organification defect in thyroid
Dehalogenase 1	Autosomal recessive	Loss of iodide reutilization

physiologic marker of thyroid hormone action. TSH is a 31-kDa hormone composed of α and β subunits; the α subunit is common to the other glycoprotein hormones (luteinizing hormone, follicle-stimulating hormone, human chorionic gonadotropin [hCG]), whereas the TSH β subunit is unique to TSH. The extent and nature of carbohydrate modification are modulated by thyrotropin-releasing hormone (TRH) stimulation and influence the biologic activity of the hormone.

The thyroid axis is a classic example of an endocrine feedback loop. Hypothalamic TRH stimulates pituitary production of TSH, which, in turn, stimulates thyroid hormone synthesis and secretion. Thyroid hormones act via negative feedback predominantly through thyroid hormone receptor $\beta 2$ (TR $\beta 2$) to inhibit TRH and TSH production (Fig. 405-2). The “set-point” in this axis is established by TSH. TRH is the major positive regulator of TSH synthesis and secretion. Peak TSH secretion occurs ~15 min after administration of exogenous TRH. Dopamine, glucocorticoids, and somatostatin suppress TSH but are not of major physiologic importance except when these agents are administered in pharmacologic doses. Reduced levels of thyroid hormone increase basal TSH production and enhance TRH-mediated stimulation of TSH. High thyroid hormone levels rapidly and directly suppress TSH gene expression secretion and inhibit TRH stimulation of TSH, indicating that thyroid hormones are the dominant regulator of TSH production. Like other pituitary hormones, TSH is released in a pulsatile manner and exhibits a diurnal rhythm; its highest levels occur at night. However, these TSH excursions are modest in comparison to those of other pituitary hormones, in part, because TSH has a relatively long plasma half-life (50 min). Consequently, single measurements of TSH are adequate for assessing its circulating level. TSH is measured using immunoradiometric assays that are highly sensitive and specific. These assays readily distinguish between normal and suppressed TSH values; thus, TSH can be used for the diagnosis of hyperthyroidism (low TSH) as well as hypothyroidism (high TSH).

THYROID HORMONE SYNTHESIS, METABOLISM, AND ACTION

THYROID HORMONE SYNTHESIS

Thyroid hormones are derived from Tg, a large iodinated glycoprotein. After secretion into the thyroid follicle, Tg is iodinated on tyrosine

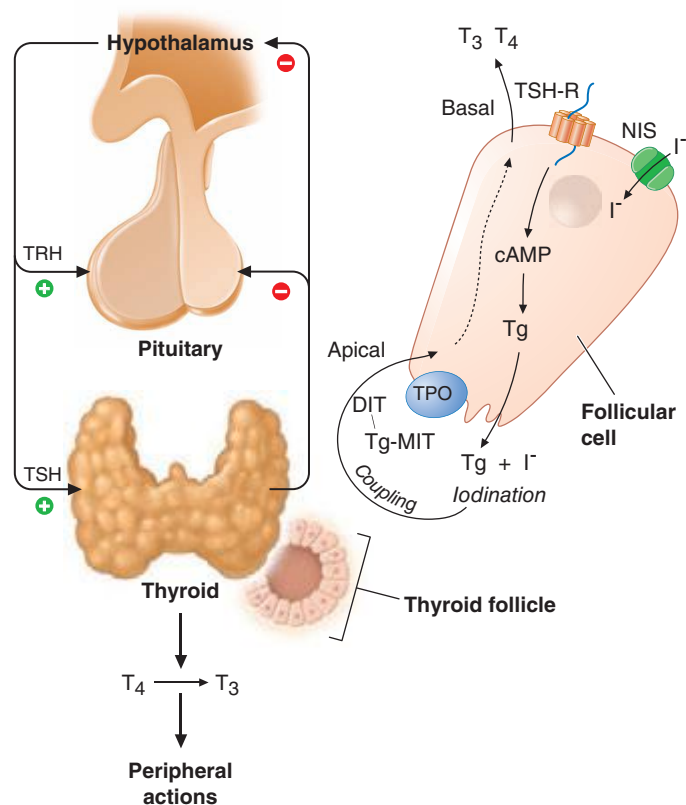



FIGURE 405-2 Regulation of thyroid hormone synthesis. Left.

Thyroid hormones T₄ and T₃ feed back to inhibit hypothalamic production of thyrotropin-releasing hormone (TRH) and pituitary production of thyroid-stimulating hormone (TSH). TSH stimulates thyroid gland production of T₄ and T₃. **Right.** Thyroid follicles are formed by thyroid epithelial cells surrounding proteinaceous colloid, which contains thyroglobulin. Follicular cells, which are polarized, synthesize thyroglobulin and carry out thyroid hormone biosynthesis (see text for details). DIT, diiodotyrosine; MIT, monoiodotyrosine; NIS, sodium iodide symporter; Tg, thyroglobulin; TPO, thyroid peroxidase; TSH-R, thyroid-stimulating hormone receptor.

residues that are subsequently coupled via an ether linkage. Reuptake of Tg into the thyroid follicular cell allows proteolysis and the release of newly synthesized T₄ and T₃.

Iodine Metabolism and Transport Iodide uptake is a critical first step in thyroid hormone synthesis. Ingested iodine is bound to serum proteins, particularly albumin. Unbound iodine is excreted in the urine. The thyroid gland extracts iodine from the circulation in a highly efficient manner. For example, 10–25% of radioactive tracer (e.g., ¹²³I) is taken up by the normal thyroid gland over 24 h; this value can rise to 70–90% in Graves’ disease. Iodide uptake is mediated by NIS, which is expressed at the basolateral membrane of thyroid follicular cells. NIS is most highly expressed in the thyroid gland, but low levels are present in the salivary glands, lactating breast, and placenta. The iodide transport mechanism is highly regulated, allowing adaptation to variations in dietary supply. Low iodine levels increase the amount of NIS and stimulate uptake, whereas high iodine levels suppress NIS expression and uptake. The selective expression of NIS in the thyroid allows isotopic scanning, treatment of hyperthyroidism, and ablation of thyroid cancer with radioisotopes of iodine, without significant effects on other organs. Mutation of the *NIS* gene is a rare cause of congenital hypothyroidism, underscoring its importance in thyroid hormone synthesis. Another iodine transporter, pendrin, is located on the apical surface of thyroid cells and mediates iodine efflux into the lumen. Mutation of the *pendrin* gene causes *Pendred syndrome*, a disorder characterized by defective organification of iodine, goiter, and sensorineural deafness.

 Iodine deficiency is prevalent in many mountainous regions and in central Africa, central South America, and northern Asia (Fig. 405-3). Europe remains mildly iodine-deficient, and health surveys indicate that iodine intake has been falling in the United States and Australia. The World Health Organization (WHO) estimates that about 2 billion people are iodine-deficient, based on urinary excretion data. In areas of relative iodine deficiency, there is an increased prevalence of goiter and, when deficiency is severe, hypothyroidism and cretinism. *Cretinism* is characterized by mental and growth retardation and occurs when children who live in iodine-deficient regions are not treated with iodine or thyroid hormone to restore normal thyroid hormone levels during early life. These children are often born to mothers with iodine deficiency, and it is likely that maternal thyroid hormone deficiency worsens the condition. Concomitant selenium deficiency may also contribute to the neurologic manifestations of cretinism. Iodine supplementation of salt, bread, and other food substances has markedly reduced the prevalence of cretinism. Unfortunately, however, iodine deficiency remains the most common cause of preventable mental deficiency, often because of societal resistance to food additives or the cost of supplementation. In addition to overt cretinism, mild iodine deficiency can lead to subtle reduction of IQ. Oversupply of iodine, through supplements or foods enriched in iodine (e.g., shellfish, kelp), is associated with an increased incidence of autoimmune thyroid disease. The recommended average daily intake of iodine is 150–250 µg/d for adults, 90–120 µg/d for children, and 250 µg/d for pregnant and lactating women. Urinary iodine is >10 µg/dL in iodine-sufficient populations.

Organification, Coupling, Storage, and Release After iodide enters the thyroid, it is trapped and transported to the apical membrane of thyroid follicular cells, where it is oxidized in an organification reaction that involves TPO and hydrogen peroxide produced by dual oxidase (DUOX) and DUOX maturation factor (DUOXA). The reactive iodine atom is added to selected tyrosyl residues within Tg, a large (660 kDa) dimeric protein that consists of 2769 amino acids. The iodotyrosines in Tg are then coupled via an ether linkage in a reaction that is also catalyzed by TPO. Either T₄ or T₃ can be produced by this reaction, depending on the number of iodine atoms present in the iodotyrosines. After coupling, Tg is taken back into the thyroid cell, where it is processed in lysosomes to release T₄ and T₃. Uncoupled mono- and diiodotyrosines (MIT, DIT) are deiodinated by the enzyme dehalogenase, thereby recycling any iodide that is not converted into thyroid hormones.

Disorders of thyroid hormone synthesis are rare causes of congenital hypothyroidism. The vast majority of these disorders are due to recessive mutations in TPO or Tg, but defects have also been identified in the TSH-R, NIS, pendrin, hydrogen peroxide generation, and dehalogenase. Because of the biosynthetic defect, the gland is incapable of synthesizing adequate amounts of hormone, leading to increased TSH and a large goiter.

TSH Action TSH regulates thyroid gland function through the TSH-R, a seven-transmembrane G protein-coupled receptor (GPCR). The TSH-R is coupled to the α subunit of stimulatory G protein (G_{sa}), which activates adenylyl cyclase, leading to increased production of cyclic adenosine monophosphate (AMP). TSH also stimulates phosphatidylinositol turnover by activating phospholipase C. The functional role of the TSH-R is exemplified by the consequences of naturally occurring mutations. Recessive loss-of-function mutations cause thyroid hypoplasia and congenital hypothyroidism. Dominant gain-of-function mutations cause sporadic or familial hyperthyroidism that is characterized by goiter, thyroid cell hyperplasia, and autonomous function. Most of these activating mutations occur in the transmembrane domain of the receptor. They mimic the conformational changes induced by TSH binding or the interactions of thyroid-stimulating immunoglobulins (TSI) in Graves' disease. Activating TSH-R mutations also occur as somatic events, leading to clonal selection and expansion of the affected thyroid follicular cell and autonomously functioning thyroid nodules (see below).

Other Factors That Influence Hormone Synthesis and Release Although TSH is the dominant hormonal regulator of thyroid gland growth and function, a variety of growth factors, most produced locally in the thyroid gland, also influence thyroid hormone synthesis. These include insulin-like growth factor I (IGF-I), epidermal growth factor, transforming growth factor β (TGF-β), endothelins, and various cytokines. The quantitative roles of these factors are not well understood, but they are important in selected disease states. In acromegaly, for example, increased levels of growth hormone and IGF-I are associated with goiter and predisposition to multinodular goiter (MNG). Certain cytokines and interleukins (ILs) produced in association with autoimmune thyroid disease induce thyroid growth, whereas others lead to apoptosis. Iodine deficiency increases thyroid blood flow and upregulates the NIS, stimulating more efficient iodine uptake. Excess iodide transiently inhibits thyroid iodide organification, a phenomenon known as the *Wolff-Chaikoff effect*. In individuals with a normal thyroid, the gland escapes from this inhibitory effect and iodide organification resumes; the suppressive action of high iodide may persist, however, in patients with underlying autoimmune thyroid disease.

THYROID HORMONE TRANSPORT AND METABOLISM

Serum Binding Proteins T₄ is secreted from the thyroid gland in about twentyfold excess over T₃ (Table 405-2). Both hormones are bound to plasma proteins, including thyroxine-binding globulin (TBG), transthyretin (TTR, formerly known as thyroxine-binding prealbumin, or TBPA), and albumin. The plasma-binding proteins increase the pool of circulating hormone, delay hormone clearance, and may modulate hormone delivery to selected tissue sites. The concentration of TBG is relatively low (1–2 mg/dL), but because of its high affinity for thyroid hormones (T₄ > T₃), it carries about 80% of the bound hormones. Albumin has relatively low affinity for thyroid hormones but has a high plasma concentration (~3.5 g/dL), and it binds up to 10% of T₄ and 30% of T₃. TTR carries about 10% of T₄ but little T₃.

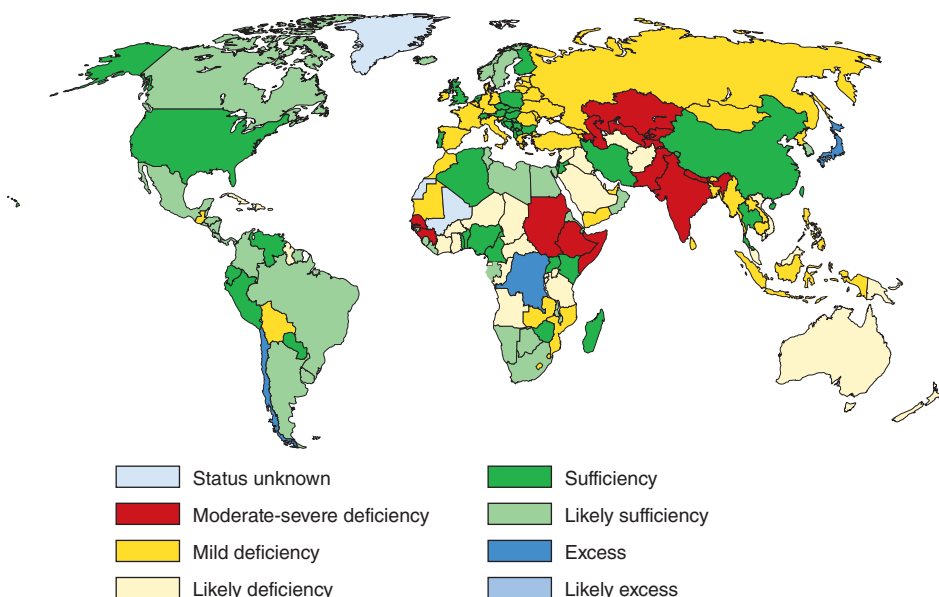


FIGURE 405-3 Worldwide iodine nutrition. Data are from the World Health Organization and the International Council for the Control of Iodine Deficiency Disorders (<http://indorgs.virginia.edu/iccidd/mi/cidds.html>).

TABLE 405-2 CHARACTERISTICS OF CIRCULATING T₄ AND T₃

Hormone Property	T ₄	T ₃
Serum concentrations		
Total hormone	8 µg/dL	0.14 µg/dL
Fraction of total hormone in the unbound form	0.02%	0.3%
Unbound (free) hormone	21 × 10 ⁻¹² M	6 × 10 ⁻¹² M
Serum half-life	7 d	2 d
Fraction directly from the thyroid	100%	20%
Production rate, including peripheral conversion	90 µg/d	32 µg/d
Intracellular hormone fraction	~20%	~70%
Relative metabolic potency	0.3	1
Receptor binding	10 ⁻¹⁰ M	10 ⁻¹¹ M

When the effects of the various binding proteins are combined, approximately 99.98% of T₄ and 99.7% of T₃ are protein-bound. Because T₃ is less tightly bound than T₄, the fraction of unbound T₃ is greater than unbound T₄, but there is less unbound T₃ in the circulation because it is produced in smaller amounts and cleared more rapidly than T₄. The unbound or “free” concentrations of the hormones are ~2 × 10⁻¹¹ M for T₄ and ~6 × 10⁻¹² M for T₃, which roughly correspond to the thyroid hormone receptor binding constants for these hormones (see below). The unbound hormone is thought to be biologically available to tissues. Nonetheless, the homeostatic mechanisms that regulate the thyroid axis are directed toward maintenance of normal concentrations of unbound hormones.

Abnormalities of Thyroid Hormone Binding Proteins A number of inherited and acquired abnormalities affect thyroid hormone binding proteins. X-linked TBG deficiency is associated with very low levels of total T₄ and T₃. However, because unbound hormone levels are normal, patients are euthyroid and TSH levels are normal. It is important to recognize this disorder to avoid efforts to normalize total T₄ levels, because this leads to thyrotoxicosis and is futile because of rapid hormone clearance in the absence of TBG. TBG levels are elevated by estrogen, which increases sialylation and delays TBG clearance. Consequently, in women who are pregnant or taking estrogen-containing contraceptives, elevated TBG increases total T₄ and T₃ levels; however, unbound T₄ and T₃ levels are normal. These features are part of the explanation for why women with hypothyroidism require increased amounts of L-thyroxine replacement as TBG levels are increased by pregnancy or estrogen treatment. Mutations in TBG, TTR, and albumin may increase the binding affinity for T₄ and/or T₃ and cause disorders known as *euthyroid hyperthyroxinemia* or *familial dysalbuminemic hyperthyroxinemia* (FDH) (Table 405-3). These disorders result in increased total T₄ and/or T₃, but unbound hormone levels are normal. The familial nature of the disorders, and the fact that TSH levels are normal rather than suppressed, should suggest this diagnosis. Unbound hormone levels (ideally measured by dialysis) are normal in FDH. The diagnosis can be confirmed by using tests that measure the affinities of radiolabeled hormone binding to specific transport proteins or by performing DNA sequence analyses of the abnormal transport protein genes.

Certain medications, such as salicylates and salsalate, can displace thyroid hormones from circulating binding proteins. Although these drugs transiently perturb the thyroid axis by increasing free thyroid hormone levels, TSH is suppressed

until a new steady state is reached, thereby restoring euthyroidism. Circulating factors associated with acute illness may also displace thyroid hormone from binding proteins (see “Sick Euthyroid Syndrome,” below).

Deiodinases T₄ may be thought of as a precursor for the more potent T₃. T₄ is converted to T₃ by the deiodinase enzymes (Fig. 405-1). Type I deiodinase, which is located primarily in thyroid, liver, and kidneys, has a relatively low affinity for T₄. Type II deiodinase has a higher affinity for T₄ and is found primarily in the pituitary gland, brain, brown fat, and thyroid gland. Expression of type II deiodinase allows it to regulate T₃ concentrations locally, a property that may be important in the context of levothyroxine (T₄) replacement. Type II deiodinase is also regulated by thyroid hormone; hypothyroidism induces the enzyme, resulting in enhanced T₄ → T₃ conversion in tissues such as brain and pituitary. T₄ → T₃ conversion is impaired by fasting, systemic illness or acute trauma, oral contrast agents, and a variety of medications (e.g., propylthiouracil, propranolol, amiodarone, glucocorticoids). Type III deiodinase inactivates T₄ and T₃ and is the most important source of reverse T₃ (rT₃), including in the sick euthyroid syndrome. This enzyme is expressed in the human placenta but is not active in healthy individuals. In the sick euthyroid syndrome, especially with hypoperfusion, the type III deiodinase is activated in muscle and liver. Massive hemangiomas that express type III deiodinase are a rare cause of hypothyroidism in infants.

THYROID HORMONE ACTION

Thyroid Hormone Transport Circulating thyroid hormones enter cells by passive diffusion and via specific transporters such as the monocarboxylate 8 transporter (MCT8), MCT10, and organic anion-transporting polypeptide 1C1. Mutations in the *MCT8* gene have been identified in patients with X-linked psychomotor retardation and thyroid function abnormalities (low T₄, high T₃, and high TSH). After entering cells, thyroid hormones act primarily through nuclear receptors, although they also have nongenomic actions through stimulating mitochondrial enzymatic responses and may act directly on blood vessels and the heart through integrin receptors.

Nuclear Thyroid Hormone Receptors Thyroid hormones bind with high affinity to nuclear *thyroid hormone receptors* (TRs) α and β. Both TRα and TRβ are expressed in most tissues, but their relative expression levels vary

TABLE 405-3 CONDITIONS ASSOCIATED WITH EUTHYROID HYPERTHYROXINEMIA

Disorder	Cause	Transmission	Characteristics
Familial dysalbuminemic hyperthyroxinemia (FDH)	Albumin mutations, usually R218H	AD	Increased T ₄ Normal unbound T ₄ Rarely increased T ₃
TBG			
Familial excess	Increased TBG production	XL	Increased total T ₄ , T ₃ Normal unbound T ₄ , T ₃
Acquired excess	Medications (estrogen), pregnancy, cirrhosis, hepatitis	Acquired	Increased total T ₄ , T ₃ Normal unbound T ₄ , T ₃
Transthyretin ^a			
Excess	Islet tumors	Acquired	Usually normal T ₄ , T ₃
Mutations	Increased affinity for T ₄ or T ₃	AD	Increased total T ₄ , T ₃ Normal unbound T ₄ , T ₃
Medications: propranolol, ipodate, iopanoic acid, amiodarone	Decreased T ₄ → T ₃ conversion	Acquired	Increased T ₄ Decreased T ₃ Normal or increased TSH
Resistance to thyroid hormone (RTH)	Thyroid hormone receptor β mutations	AD	Increased unbound T ₄ , T ₃ Normal or increased TSH Some patients clinically thyrotoxic

^aAlso known as thyroxine-binding prealbumin (TBPA).

Abbreviations: AD, autosomal dominant; TBG, thyroxine-binding globulin; TSH, thyroid-stimulating hormone; XL, X-linked.

among organs; TR α is particularly abundant in brain, kidneys, gonads, muscle, and heart, whereas TR β expression is relatively high in the pituitary and liver. Both receptors are variably spliced to form unique isoforms. The TR β 2 isoform, which has a unique amino terminus, is selectively expressed in the hypothalamus and pituitary, where it plays a role in feedback control of the thyroid axis (see above). The TR α 2 isoform contains a unique carboxy terminus that precludes thyroid hormone binding; it may function to block the action of other TR isoforms.

The TRs contain a central DNA-binding domain and a C-terminal ligand-binding domain. They bind to specific DNA sequences, termed *thyroid response elements* (TREs), in the promoter regions of target genes (Fig. 405-4). The receptors bind as homodimers or, more commonly, as heterodimers with retinoic acid X receptors (RXRs) (Chap. 400e). The activated receptor can either stimulate gene transcription (e.g., myosin heavy chain α) or inhibit transcription (e.g., TSH β -subunit gene), depending on the nature of the regulatory elements in the target gene.

Thyroid hormones (T_3 and T_4) bind with similar affinities to TR α and TR β . However, structural differences in the ligand binding domains provide the potential for developing receptor-selective agonists or antagonists, and these are under investigation. T_3 is bound with 10–15 times greater affinity than T_4 , which explains its increased hormonal potency. Although T_4 is produced in excess of T_3 , receptors are occupied mainly by T_3 , reflecting $T_4 \rightarrow T_3$ conversion by peripheral tissues, greater T_3 bioavailability in the plasma, and the greater affinity of receptors for T_3 . After binding to TRs, thyroid hormone induces conformational changes in the receptors that modify its interactions with accessory transcription factors. Importantly, in the absence of thyroid hormone binding, the aporeceptors bind to co-repressor proteins that inhibit gene transcription. Hormone binding dissociates the co-repressors and allows the recruitment of co-activators that enhance transcription. The discovery of TR interactions with co-repressors explains the fact that TR silences gene expression in the absence of hormone binding. Consequently, hormone deficiency has a profound effect on gene expression because it causes gene repression as well as loss of hormone-induced stimulation. This concept has been corroborated by the finding that targeted deletion of the TR genes in mice has a less pronounced phenotypic effect than hormone deficiency.

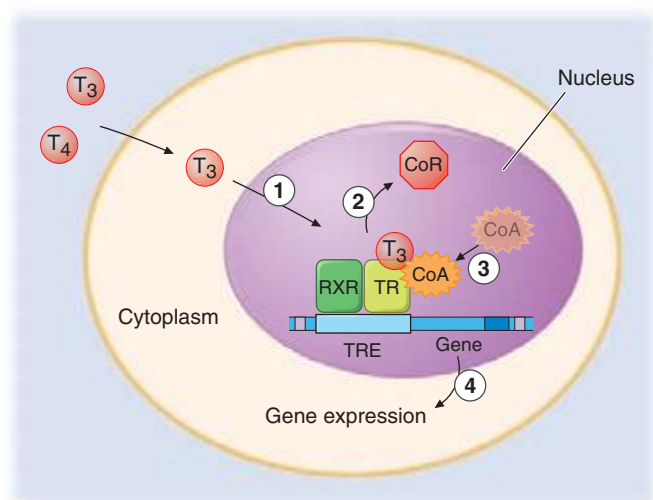


FIGURE 405-4 Mechanism of thyroid hormone receptor action.

The thyroid hormone receptor (TR) and retinoid X receptor (RXR) form heterodimers that bind specifically to thyroid hormone response elements (TRE) in the promoter regions of target genes. In the absence of hormone, TR binds co-repressor (CoR) proteins that silence gene expression. The numbers refer to a series of ordered reactions that occur in response to thyroid hormone: (1) T_4 or T_3 enters the nucleus; (2) T_3 binding dissociates CoR from TR; (3) co-activators (CoA) are recruited to the T_3 -bound receptor; and (4) gene expression is altered.

Thyroid Hormone Resistance Resistance to thyroid hormone (RTH) is an autosomal dominant disorder characterized by elevated thyroid hormone levels and inappropriately normal or elevated TSH. Individuals with RTH do not, in general, exhibit signs and symptoms that are typical of hypothyroidism because hormone resistance is partial and is compensated by increased levels of thyroid hormone. The clinical features of RTH can include goiter, attention deficit disorder, mild reduction in IQ, delayed skeletal maturation, tachycardia, and impaired metabolic responses to thyroid hormone.

Classical forms of RTH are caused by mutations in the TR β gene. These mutations, located in restricted regions of the ligand-binding domain, cause loss of receptor function. However, because the mutant receptors retain the capacity to dimerize with RXRs, bind to DNA, and recruit co-repressor proteins, they function as antagonists of the remaining normal TR β and TR α receptors. This property, referred to as “dominant negative” activity, explains the autosomal dominant mode of transmission. The diagnosis is suspected when unbound thyroid hormone levels are increased without suppression of TSH. Similar hormonal abnormalities are found in other affected family members, although the TR β mutation arises *de novo* in about 20% of patients. DNA sequence analysis of the TR β gene provides a definitive diagnosis. RTH must be distinguished from other causes of euthyroid hyperthyroxinemia (e.g., FDH) and inappropriate secretion of TSH by TSH-secreting pituitary adenomas (Chap. 403). In most patients, no treatment is indicated; the importance of making the diagnosis is to avoid inappropriate treatment of mistaken hyperthyroidism and to provide genetic counseling.

A distinct form of RTH is caused by mutations in the TR α gene. Affected patients have many clinical features of congenital hypothyroidism including growth retardation, skeletal dysplasia, and severe constipation. In contrast to RTH caused by mutations in TR β , thyroid function tests include normal TSH, low or normal T_4 , and normal or elevated T_3 levels. These distinct clinical and laboratory features underscore the different tissue distribution and functional roles of TR β and TR α . Optimal treatment of patients with RTH caused by TR α mutations has not been established.

PHYSICAL EXAMINATION

In addition to the examination of the thyroid itself, the physical examination should include a search for signs of abnormal thyroid function and the extrathyroidal features of ophthalmopathy and dermopathy (see below). Examination of the neck begins by inspecting the seated patient from the front and side and noting any surgical scars, obvious masses, or distended veins. The thyroid can be palpated with both hands from behind or while facing the patient, using the thumbs to palpate each lobe. It is best to use a combination of these methods, especially when nodules are small. The patient’s neck should be slightly flexed to relax the neck muscles. After locating the cricoid cartilage, the isthmus, which is attached to the lower one-third of the thyroid lobes, can be identified and then followed laterally to locate either lobe (normally, the right lobe is slightly larger than the left). By asking the patient to swallow sips of water, thyroid consistency can be better appreciated as the gland moves beneath the examiner’s fingers.

Features to be noted include thyroid size, consistency, nodularity, and any tenderness or fixation. An estimate of thyroid size (normally 12–20 g) should be made, and a drawing is often the best way to record findings. However, ultrasound is the method of choice when it is important to determine thyroid size accurately. The size, location, and consistency of any nodules should also be defined. A bruit or thrill over the gland, located over the insertion of the superior and inferior thyroid arteries (supero- or inferolaterally), indicates increased vascularity, as occurs in hyperthyroidism. If the lower borders of the thyroid lobes are not clearly felt, a goiter may be retrosternal. Large retrosternal goiters can cause venous distention over the neck and difficulty breathing, especially when the arms are raised (Pemberton’s sign). With any central mass above the thyroid, the tongue should be extended, as thyroglossal cysts then move upward. The thyroid examination is not complete without assessment for lymphadenopathy in the supraclavicular and cervical regions of the neck.

Measurement of Thyroid Hormones The enhanced sensitivity and specificity of *TSH assays* have greatly improved laboratory assessment of thyroid function. Because TSH levels change dynamically in response to alterations of T_4 and T_3 , a logical approach to thyroid testing is to first determine whether TSH is suppressed, normal, or elevated. With rare exceptions (see below), a normal TSH level excludes a primary abnormality of thyroid function. This strategy depends on the use of immunochemiluminometric assays (ICMAs) for TSH that are sensitive enough to discriminate between the lower limit of the reference range and the suppressed values that occur with thyrotoxicosis. Extremely sensitive (fourth-generation) assays can detect TSH levels ≤ 0.004 mIU/L, but, for practical purposes, assays sensitive to ≤ 0.1 mIU/L are sufficient. The widespread availability of the TSH ICMA has rendered the TRH stimulation test obsolete, because the failure of TSH to rise after an intravenous bolus of 200–400 μg TRH has the same implications as a suppressed basal TSH measured by ICMA.

The finding of an abnormal TSH level must be followed by measurements of circulating thyroid hormone levels to confirm the diagnosis of hyperthyroidism (suppressed TSH) or hypothyroidism (elevated TSH). Radioimmunoassays are widely available for serum *total* T_4 and *total* T_3 . T_4 and T_3 are highly protein-bound, and numerous factors (illness, medications, genetic factors) can influence protein binding. It is useful, therefore, to measure the free, or unbound, hormone levels, which correspond to the biologically available hormone pool. Two direct methods are used to measure *unbound thyroid hormones*: (1) unbound thyroid hormone competition with radiolabeled T_4 (or an analogue) for binding to a solid-phase antibody, and (2) physical separation of the unbound hormone fraction by ultracentrifugation or equilibrium dialysis. Although early unbound hormone immunoassays suffered from artifacts, newer assays correlate well with the results of the more technically demanding and expensive physical separation methods. An indirect method that is now less commonly used to estimate unbound thyroid hormone levels is to calculate the free T_3 or free T_4 index from the total T_4 or T_3 concentration and the *thyroid hormone binding ratio* (THBR). The latter is derived from the *T_3 -resin uptake test*, which determines the distribution of radiolabeled T_3 between an absorbent resin and the unoccupied thyroid hormone binding proteins in the sample. The binding of the labeled T_3 to the resin is increased when there is reduced unoccupied protein binding sites (e.g., TBG deficiency) or increased total thyroid hormone in the sample; it is decreased under the opposite circumstances. The product of THBR and total T_3 or T_4 provides the *free T_3 or T_4 index*. In effect, the index corrects for anomalous total hormone values caused by abnormalities in hormone-protein binding.

Total thyroid hormone levels are *elevated* when TBG is increased due to estrogens (pregnancy, oral contraceptives, hormone therapy, tamoxifen, selective estrogen receptor modulators, inflammatory liver disease) and *decreased* when TBG binding is reduced (androgens, nephrotic syndrome). Genetic disorders and acute illness can also cause abnormalities in thyroid hormone binding proteins, and various drugs (phenytoin, carbamazepine, salicylates, and nonsteroidal anti-inflammatory drugs [NSAIDs]) can interfere with thyroid hormone binding. Because unbound thyroid hormone levels are normal and the patient is euthyroid in all of these circumstances, assays that measure unbound hormone are preferable to those for total thyroid hormones.

For most purposes, the unbound T_4 level is sufficient to confirm thyrotoxicosis, but 2–5% of patients have only an elevated T_3 level (T_3 toxicosis). Thus, unbound T_3 levels should be measured in patients with a suppressed TSH but normal unbound T_4 levels.

There are several clinical conditions in which the use of TSH as a screening test may be misleading, particularly without simultaneous unbound T_4 determinations. Any severe nonthyroidal illness can cause abnormal TSH levels (see below). Although hypothyroidism is the most common cause of an elevated TSH level, rare causes include a TSH-secreting pituitary tumor (**Chap. 403**), thyroid hormone resistance, and assay artifact. Conversely, a suppressed TSH level, particularly < 0.01 mIU/L, usually indicates thyrotoxicosis. However, subnormal TSH levels between 0.01 and 0.1 mIU/L may be seen during the first trimester of pregnancy (due to hCG secretion), after treatment

of hyperthyroidism (because TSH can remain suppressed for several months), and in response to certain medications (e.g., high doses of glucocorticoids or dopamine). Importantly, secondary hypothyroidism, caused by hypothalamic-pituitary disease, is associated with a variable (low to high-normal) TSH level, which is inappropriate for the low T_4 level. Thus, *TSH should not be used as an isolated laboratory test to assess thyroid function in patients with suspected or known pituitary disease.*

Tests for the end-organ effects of thyroid hormone excess or depletion, such as estimation of basal metabolic rate, tendon reflex relaxation rates, or serum cholesterol, are not useful as clinical determinants of thyroid function.

Tests to Determine the Etiology of Thyroid Dysfunction Autoimmune thyroid disease is detected most easily by measuring circulating antibodies against TPO and Tg. Because antibodies to Tg alone are uncommon, it is reasonable to measure only TPO antibodies. About 5–15% of euthyroid women and up to 2% of euthyroid men have thyroid antibodies; such individuals are at increased risk of developing thyroid dysfunction. Almost all patients with autoimmune hypothyroidism, and up to 80% of those with Graves' disease, have TPO antibodies, usually at high levels.

TSIs are antibodies that stimulate the TSH-R in Graves' disease. They are most commonly measured by commercially available tracer displacement assays called TRAb (TSH receptor antibody) with the assumption that elevated levels in the setting of clinical hyperthyroidism reflect stimulatory effects on the TSH receptor. A bioassay is less commonly used. The main use of these assays is to predict neonatal thyrotoxicosis caused by high maternal levels of TRAb or TSI ($> 3\times$ upper limit of normal) in the last trimester of pregnancy.

Serum Tg levels are increased in all types of thyrotoxicosis except *thyrotoxicosis factitia* caused by self-administration of thyroid hormone. Tg levels are particularly increased in thyroiditis, reflecting thyroid tissue destruction and release of Tg. The main role for Tg measurement, however, is in the follow-up of thyroid cancer patients. After total thyroidectomy and radioablation, Tg levels should be undetectable; in the absence of anti-Tg antibodies, measurable levels indicate incomplete ablation or recurrent cancer.

Radioiodine Uptake and Thyroid Scanning The thyroid gland selectively transports radioisotopes of iodine (^{123}I , ^{125}I , ^{131}I) and $^{99\text{m}}\text{Tc}$ pertechnetate, allowing thyroid imaging and quantitation of radioactive tracer fractional uptake.

Nuclear imaging of Graves' disease is characterized by an enlarged gland and increased tracer uptake that is distributed homogeneously. Toxic adenomas appear as focal areas of increased uptake, with suppressed tracer uptake in the remainder of the gland. In toxic MNG, the gland is enlarged—often with distorted architecture—and there are multiple areas of relatively increased (functioning nodules) or decreased tracer uptake (suppressed thyroid parenchyma or nonfunctioning nodules). Subacute, viral, and postpartum thyroiditis are associated with very low uptake because of follicular cell damage and TSH suppression. Thyrotoxicosis factitia is also associated with low uptake. In addition, if there is excessive circulating exogenous iodine (e.g., from dietary sources of iodinated contrast dye), the radionuclide uptake is low even in the presence of increased thyroid hormone production.

Thyroid scintigraphy is not used in the routine evaluation of patients with thyroid nodules, but should be performed if the serum TSH level is subnormal to determine if functioning thyroid nodules are present. Functioning or “hot” nodules are almost never malignant, and fine-needle aspiration (FNA) biopsy is not indicated. The vast majority of thyroid nodules do not produce thyroid hormone (“cold” nodules), and these are more likely to be malignant (~5–10%). Whole-body and thyroid scanning is also used in the treatment and surveillance of thyroid cancer. After thyroidectomy for thyroid cancer, the TSH level is raised by either using a thyroid hormone withdrawal protocol or recombinant human TSH injection (see below). Administration of ^{131}I allows whole-body scanning (WBS) to confirm remnant ablation and to detect any functioning metastases. In addition, WBS may be helpful in surveillance of patients at risk for recurrence.

TABLE 405-4 GRAYSCALE SONOGRAPHIC FEATURES ASSOCIATED WITH THYROID CANCER

	Median Sensitivity [range]	Median Specificity [range]
Hypoechoic compared with surrounding thyroid	81% [48–90%]	53% [36–92%]
Marked hypoechogenicity	41% [27–59%]	94% [92–94%]
Microcalcifications	44% [26–73%]	89% [69–98%]
Irregular, microlobulated margins	55% [17–84%]	79% [62–85%]
Solid consistency	86% [78–91%]	48% [30–58%]
Taller than wide shape on transverse view	48% [33–84%]	92% [82–93%]

Thyroid Ultrasound Ultrasonography is valuable for the diagnosis and evaluation of patients with nodular thyroid disease (Table 405-4). Evidence-based guidelines recommend thyroid ultrasonography for all patients suspected of having thyroid nodules by either physical examination or another imaging study. Using 10- to 12-MHz linear transducers, resolution and image quality are excellent, allowing the characterization of nodules and cysts >3 mm. Certain sonographic patterns are highly suggestive of malignancy (e.g., hypoechoic solid nodules with infiltrative borders and microcalcifications), whereas other features correlate with benignity (e.g., spongiform nodules defined as those with multiple small internal cystic areas) (Fig. 405-5). In addition to evaluating thyroid nodules, ultrasound is useful for monitoring nodule size and for the aspiration of nodules or cystic lesions. Ultrasound-guided FNA biopsy of thyroid lesions lowers the rate of inadequate sampling and decreases sample error, thereby reducing the false-negative rate of FNA cytology. Ultrasonography of the central and lateral cervical lymph node compartments is indispensable in the evaluation thyroid cancer patients, preoperatively and during follow-up.

HYPOTHYROIDISM

Iodine deficiency remains a common cause of hypothyroidism worldwide. In areas of iodine sufficiency, autoimmune disease (Hashimoto's

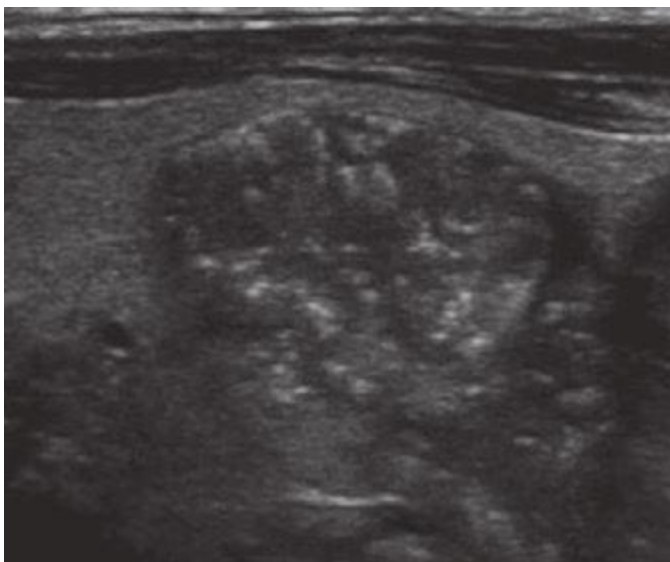


FIGURE 405-5 Sonographic patterns of thyroid nodules. **A.** High suspicion ultrasound pattern for thyroid malignancy (hypoechoic solid nodule with irregular borders and microcalcifications). **B.** Very low suspicion ultrasound pattern for thyroid malignancy (spongiform nodule with microcystic areas comprises over >50% of nodule volume).

TABLE 405-5 CAUSES OF HYPOTHYROIDISM

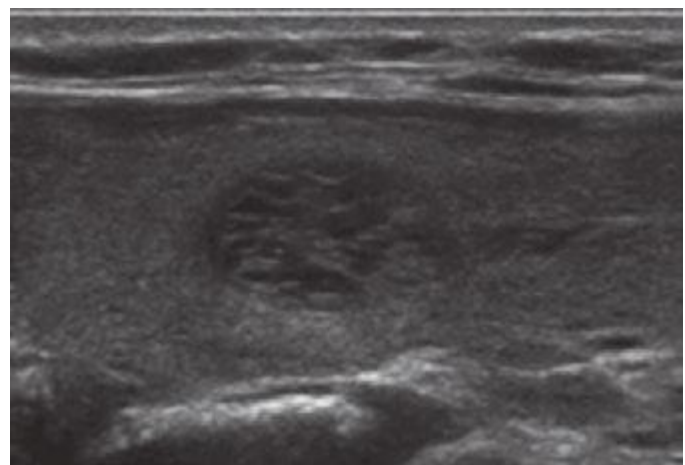
Primary
Autoimmune hypothyroidism: Hashimoto's thyroiditis, atrophic thyroiditis
Iatrogenic: ¹³¹ I treatment, subtotal or total thyroidectomy, external irradiation of neck for lymphoma or cancer
Drugs: iodine excess (including iodine-containing contrast media and amiodarone), lithium, antithyroid drugs, <i>p</i> -aminosalicylic acid, interferon α and other cytokines, aminoglutethimide, tyrosine kinase inhibitors (e.g., sunitinib)
Congenital hypothyroidism: absent or ectopic thyroid gland, dysmorphogenesis, TSH-R mutation
Iodine deficiency
Infiltrative disorders: amyloidosis, sarcoidosis, hemochromatosis, scleroderma, cystinosis, Riedel's thyroiditis
Overexpression of type 3 deiodinase in infantile hemangioma and other tumors
Transient
Silent thyroiditis, including postpartum thyroiditis
Subacute thyroiditis
Withdrawal of supraphysiologic thyroxine treatment in individuals with an intact thyroid
After ¹³¹ I treatment or subtotal thyroidectomy for Graves' disease
Secondary
Hypopituitarism: tumors, pituitary surgery or irradiation, infiltrative disorders, Sheehan's syndrome, trauma, genetic forms of combined pituitary hormone deficiencies
Isolated TSH deficiency or inactivity
Bexarotene treatment
Hypothalamic disease: tumors, trauma, infiltrative disorders, idiopathic

Abbreviations: TSH, thyroid-stimulating hormone; TSH-R, TSH receptor.

thyroiditis) and iatrogenic causes (treatment of hyperthyroidism) are most common (Table 405-5).

CONGENITAL HYPOTHYROIDISM

Prevalence Hypothyroidism occurs in about 1 in 4000 newborns. It may be transient, especially if the mother has TSH-R blocking antibodies or has received antithyroid drugs, but permanent hypothyroidism occurs in the majority. Neonatal hypothyroidism is due to thyroid gland dysgenesis in 80–85%, to inborn errors of thyroid hormone synthesis in 10–15%, and is TSH-R antibody-mediated in 5% of affected newborns. The developmental abnormalities are twice as common in girls. Mutations that cause congenital hypothyroidism are being increasingly identified, but most remain idiopathic (Table 405-1).



B

TABLE 405-6 SIGNS AND SYMPTOMS OF HYPOTHYROIDISM (DESCENDING ORDER OF FREQUENCY)

Symptoms	Signs
Tiredness, weakness	Dry coarse skin; cool peripheral extremities
Dry skin	Puffy face, hands, and feet (myxedema)
Feeling cold	Diffuse alopecia
Hair loss	Bradycardia
Difficulty concentrating and poor memory	Peripheral edema
Constipation	Delayed tendon reflex relaxation
Weight gain with poor appetite	Carpal tunnel syndrome
Dyspnea	Serous cavity effusions
Hoarse voice	
Menorrhagia (later oligomenorrhea or amenorrhea)	
Paresthesia	
Impaired hearing	

Clinical Manifestations The majority of infants appear normal at birth, and <10% are diagnosed based on clinical features, which include prolonged jaundice, feeding problems, hypotonia, enlarged tongue, delayed bone maturation, and umbilical hernia. Importantly, permanent neurologic damage results if treatment is delayed. Typical features of adult hypothyroidism may also be present (Table 405-6). Other congenital malformations, especially cardiac, are four times more common in congenital hypothyroidism.

Diagnosis and Treatment Because of the severe neurologic consequences of untreated congenital hypothyroidism, neonatal screening programs have been established. These are generally based on measurement of TSH or T_4 levels in heel-prick blood specimens. When the diagnosis is confirmed, T_4 is instituted at a dose of 10–15 $\mu\text{g}/\text{kg}$ per day, and the dose is adjusted by close monitoring of TSH levels. T_4 requirements are relatively great during the first year of life, and a high circulating T_4 level is usually needed to normalize TSH. Early treatment with T_4 results in normal IQ levels, but subtle neurodevelopmental abnormalities may occur in those with the most severe hypothyroidism at diagnosis or when treatment is delayed or suboptimal.

AUTOIMMUNE HYPOTHYROIDISM

Classification Autoimmune hypothyroidism may be associated with a goiter (Hashimoto's, or *goitrous thyroiditis*) or, at the later stages of the disease, minimal residual thyroid tissue (*atrophic thyroiditis*). Because the autoimmune process gradually reduces thyroid function, there is a phase of compensation when normal thyroid hormone levels are maintained by a rise in TSH. Although some patients may have minor symptoms, this state is called *subclinical hypothyroidism*. Later, unbound T_4 levels fall and TSH levels rise further; symptoms become more readily apparent at this stage (usually TSH >10 mIU/L), which is referred to as *clinical hypothyroidism* or *overt hypothyroidism*.

Prevalence The mean annual incidence rate of autoimmune hypothyroidism is up to 4 per 1000 women and 1 per 1000 men. It is more common in certain populations, such as the Japanese, probably because of genetic factors and chronic exposure to a high-iodine diet. The mean age at diagnosis is 60 years, and the prevalence of overt hypothyroidism increases with age. Subclinical hypothyroidism is found in 6–8% of women (10% over the age of 60) and 3% of men. The annual risk of developing clinical hypothyroidism is about 4% when subclinical hypothyroidism is associated with positive TPO antibodies.

Pathogenesis In Hashimoto's thyroiditis, there is a marked lymphocytic infiltration of the thyroid with germinal center formation, atrophy of the thyroid follicles accompanied by oxyphil metaplasia, absence of colloid, and mild to moderate fibrosis. In atrophic thyroiditis, the fibrosis is much more extensive, lymphocyte infiltration is less pronounced, and thyroid follicles are almost completely absent.

Atrophic thyroiditis likely represents the end stage of Hashimoto's thyroiditis rather than a distinct disorder.

As with most autoimmune disorders, susceptibility to autoimmune hypothyroidism is determined by a combination of genetic and environmental factors, and the risk of either autoimmune hypothyroidism or Graves' disease is increased among siblings. HLA-DR polymorphisms are the best documented genetic risk factors for autoimmune hypothyroidism, especially HLA-DR3, -DR4, and -DR5 in Caucasians. A weak association also exists between polymorphisms in *CTLA-4*, a T cell–regulatory gene, and autoimmune hypothyroidism. Both of these genetic associations are shared by other autoimmune diseases, which may explain the relationship between autoimmune hypothyroidism and other autoimmune diseases, especially type 1 diabetes mellitus, Addison's disease, pernicious anemia, and vitiligo. HLA-DR and *CTLA-4* polymorphisms account for approximately half of the genetic susceptibility to autoimmune hypothyroidism. Other contributory loci remain to be identified. A gene on chromosome 21 may be responsible for the association between autoimmune hypothyroidism and Down's syndrome. The female preponderance of thyroid autoimmunity is most likely due to sex steroid effects on the immune response, but an X chromosome–related genetic factor is also possible and may account for the high frequency of autoimmune hypothyroidism in Turner's syndrome. Environmental susceptibility factors are poorly defined at present. A high iodine intake and decreased exposure to microorganisms in childhood increase the risk of autoimmune hypothyroidism. These factors may account for the increase in prevalence over the last two to three decades.

The thyroid lymphocytic infiltrate in autoimmune hypothyroidism is composed of activated CD4+ and CD8+ T cells as well as B cells. Thyroid cell destruction is primarily mediated by the CD8+ cytotoxic T cells, which destroy their targets by either perforin-induced cell necrosis or granzyme B–induced apoptosis. In addition, local T cell production of cytokines, such as tumor necrosis factor (TNF), IL-1, and interferon γ (IFN- γ), may render thyroid cells more susceptible to apoptosis mediated by death receptors, such as Fas, which are activated by their respective ligands on T cells. These cytokines also impair thyroid cell function directly and induce the expression of other proinflammatory molecules by the thyroid cells themselves, such as cytokines, HLA class I and class II molecules, adhesion molecules, CD40, and nitric oxide. Administration of high concentrations of cytokines for therapeutic purposes (especially IFN- α) is associated with increased autoimmune thyroid disease, possibly through mechanisms similar to those in sporadic disease.

Antibodies to TPO and Tg are clinically useful markers of thyroid autoimmunity, but any pathogenic effect is restricted to a secondary role in amplifying an ongoing autoimmune response. TPO antibodies fix complement, and complement membrane-attack complexes are present in the thyroid in autoimmune hypothyroidism. However, transplacental passage of Tg or TPO antibodies has no effect on the fetal thyroid, which suggests that T cell–mediated injury is required to initiate autoimmune damage to the thyroid.

Up to 20% of patients with autoimmune hypothyroidism have antibodies against the TSH-R, which, in contrast to TSI, do not stimulate the receptor but prevent the binding of TSH. These TSH-R-blocking antibodies, therefore, cause hypothyroidism and, especially in Asian patients, thyroid atrophy. Their transplacental passage may induce transient neonatal hypothyroidism. Rarely, patients have a mixture of TSI and TSH-R-blocking antibodies, and thyroid function can oscillate between hyperthyroidism and hypothyroidism as one or the other antibody becomes dominant. Predicting the course of disease in such individuals is difficult, and they require close monitoring of thyroid function. Bioassays can be used to document that TSH-R-blocking antibodies reduce the cyclic AMP–inducing effect of TSH on cultured TSH-R-expressing cells, but these assays are difficult to perform. Thyrotropin-binding inhibitory immunoglobulin (TBII) assays that measure the binding of antibodies to the receptor by competition with radiolabeled TSH do not distinguish between TSI- and TSH-R-blocking antibodies, but a positive result in a patient with spontaneous hypothyroidism is strong evidence for the presence of blocking antibodies. The use of these assays does not generally alter clinical

management, although it may be useful to confirm the cause of transient neonatal hypothyroidism.

Clinical Manifestations The main clinical features of hypothyroidism are summarized in Table 405-6. The onset is usually insidious, and the patient may become aware of symptoms only when euthyroidism is restored. Patients with Hashimoto's thyroiditis may present because of goiter rather than symptoms of hypothyroidism. The goiter may not be large, but it is usually irregular and firm in consistency. It is often possible to palpate a pyramidal lobe, normally a vestigial remnant of the thyroglossal duct. Rarely is uncomplicated Hashimoto's thyroiditis associated with pain.

Patients with atrophic thyroiditis or the late stage of Hashimoto's thyroiditis present with symptoms and signs of hypothyroidism. The skin is dry, and there is decreased sweating, thinning of the epidermis, and hyperkeratosis of the stratum corneum. Increased dermal glycosaminoglycan content traps water, giving rise to skin thickening without pitting (*myxedema*). Typical features include a puffy face with edematous eyelids and nonpitting pretibial edema (Fig. 405-6). There is pallor, often with a yellow tinge to the skin due to carotene accumulation. Nail growth is retarded, and hair is dry, brittle, difficult to manage, and falls out easily. In addition to diffuse alopecia, there is thinning of the outer third of the eyebrows, although this is not a specific sign of hypothyroidism.

Other common features include constipation and weight gain (despite a poor appetite). In contrast to popular perception, the weight gain is usually modest and due mainly to fluid retention in the myxedematous tissues. Libido is decreased in both sexes, and there may be oligomenorrhea or amenorrhea in long-standing disease, but menorrhagia is also common. Fertility is reduced, and the incidence of miscarriage is increased. Prolactin levels are often modestly increased (Chap. 403) and may contribute to alterations in libido and fertility and cause galactorrhea.

Myocardial contractility and pulse rate are reduced, leading to a reduced stroke volume and bradycardia. Increased peripheral resistance may be accompanied by hypertension, particularly diastolic. Blood flow is diverted from the skin, producing cool extremities. Pericardial effusions occur in up to 30% of patients but rarely compromise cardiac



FIGURE 405-6 Facial appearance in hypothyroidism. Note puffy eyes and thickened skin.

function. Although alterations in myosin heavy chain isoform expression have been documented, cardiomyopathy is unusual. Fluid may also accumulate in other serous cavities and in the middle ear, giving rise to conductive deafness. Pulmonary function is generally normal, but dyspnea may be caused by pleural effusion, impaired respiratory muscle function, diminished ventilatory drive, or sleep apnea.

Carpal tunnel and other entrapment syndromes are common, as is impairment of muscle function with stiffness, cramps, and pain. On examination, there may be slow relaxation of tendon reflexes and pseudomyotonia. Memory and concentration are impaired. Experimentally, positron emission tomography (PET) scans examining glucose metabolism in hypothyroid subjects show lower regional activity in the amygdala, hippocampus, and perigenual anterior cingulate cortex, among other regions, and this activity corrects after thyroxine replacement. Rare neurologic problems include reversible cerebellar ataxia, dementia, psychosis, and myxedema coma. *Hashimoto's encephalopathy* has been defined as a steroid-responsive syndrome associated with TPO antibodies, myoclonus, and slow-wave activity on electroencephalography, but the relationship with thyroid autoimmunity or hypothyroidism is not established. The hoarse voice and occasionally clumsy speech of hypothyroidism reflect fluid accumulation in the vocal cords and tongue.

The features described above are the consequence of thyroid hormone deficiency. However, autoimmune hypothyroidism may be associated with signs or symptoms of other autoimmune diseases, particularly vitiligo, pernicious anemia, Addison's disease, alopecia areata, and type 1 diabetes mellitus. Less common associations include celiac disease, dermatitis herpetiformis, chronic active hepatitis, rheumatoid arthritis, systemic lupus erythematosus (SLE), myasthenia gravis, and Sjögren's syndrome. Thyroid-associated ophthalmopathy, which usually occurs in Graves' disease (see below), occurs in about 5% of patients with autoimmune hypothyroidism.

Autoimmune hypothyroidism is uncommon in children and usually presents with slow growth and delayed facial maturation. The appearance of permanent teeth is also delayed. Myopathy, with muscle swelling, is more common in children than in adults. In most cases, puberty is delayed, but precocious puberty sometimes occurs. There may be intellectual impairment if the onset is before 3 years and the hormone deficiency is severe.

Laboratory Evaluation A summary of the investigations used to determine the existence and cause of hypothyroidism is provided in Fig. 405-7. A normal TSH level excludes primary (but not secondary) hypothyroidism. If the TSH is elevated, an unbound T_4 level is needed to confirm the presence of clinical hypothyroidism, but T_4 is inferior to TSH when used as a screening test, because it will not detect subclinical hypothyroidism. Circulating unbound T_3 levels are normal in about 25% of patients, reflecting adaptive deiodinase responses to hypothyroidism. T_3 measurements are, therefore, not indicated.

Once clinical or subclinical hypothyroidism is confirmed, the etiology is usually easily established by demonstrating the presence of TPO antibodies, which are present in >90% of patients with autoimmune hypothyroidism. TBII can be found in 10–20% of patients, but measurement is not needed routinely. If there is any doubt about the cause of a goiter associated with hypothyroidism, FNA biopsy can be used to confirm the presence of autoimmune thyroiditis. Other abnormal laboratory findings in hypothyroidism may include increased creatine phosphokinase, elevated cholesterol and triglycerides, and anemia (usually normocytic or macrocytic). Except when accompanied by iron deficiency, the anemia and other abnormalities gradually resolve with thyroxine replacement.

Differential Diagnosis An asymmetric goiter in Hashimoto's thyroiditis may be confused with a MNG or thyroid carcinoma, in which thyroid antibodies may also be present. Ultrasound can be used to show the presence of a solitary lesion or an MNG rather than the heterogeneous thyroid enlargement typical of Hashimoto's thyroiditis. FNA biopsy is useful in the investigation of focal nodules. Other causes of hypothyroidism are discussed below and in Table 405-5 but rarely cause diagnostic confusion.

EVALUATION OF HYPOTHYROIDISM

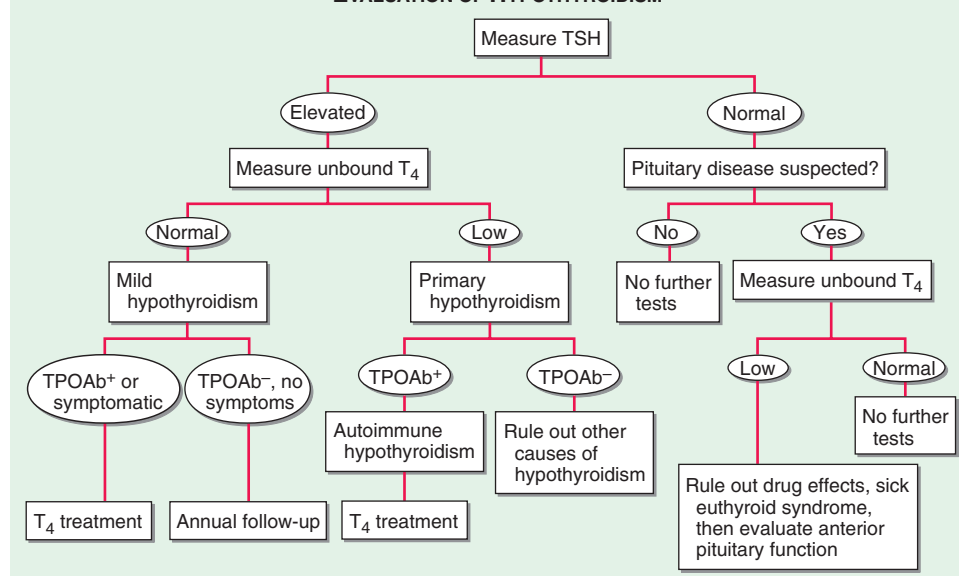


FIGURE 405-7 Evaluation of hypothyroidism. TPOAb⁺, thyroid peroxidase antibodies present; TPOAb⁻, thyroid peroxidase antibodies not present; TSH, thyroid-stimulating hormone.

OTHER CAUSES OF HYPOTHYROIDISM

Iatrogenic hypothyroidism is a common cause of hypothyroidism and can often be detected by screening before symptoms develop. In the first 3–4 months after radioiodine treatment, transient hypothyroidism may occur due to reversible radiation damage. Low-dose thyroxine treatment can be withdrawn if recovery occurs. Because TSH levels are suppressed by hyperthyroidism, unbound T₄ levels are a better measure of thyroid function than TSH in the months following radioiodine treatment. Mild hypothyroidism after subtotal thyroidectomy may also resolve after several months, as the gland remnant is stimulated by increased TSH levels.

Iodine deficiency is responsible for endemic goiter and cretinism but is an uncommon cause of adult hypothyroidism unless the iodine intake is very low or there are complicating factors, such as the consumption of thiocyanates in cassava or selenium deficiency. Although hypothyroidism due to iodine deficiency can be treated with thyroxine, public health measures to improve iodine intake should be advocated to eliminate this problem. Iodized salt or bread or a single bolus of oral or intramuscular iodized oil have all been used successfully.

Paradoxically, chronic iodine excess can also induce goiter and hypothyroidism. The intracellular events that account for this effect are unclear, but individuals with autoimmune thyroiditis are especially susceptible. Iodine excess is responsible for the hypothyroidism that occurs in up to 13% of patients treated with amiodarone (see below). Other drugs, particularly lithium, may also cause hypothyroidism. Transient hypothyroidism caused by thyroiditis is discussed below.

Secondary hypothyroidism is usually diagnosed in the context of other anterior pituitary hormone deficiencies; isolated TSH deficiency is very rare (Chap. 402). TSH levels may be low, normal, or even slightly increased in secondary hypothyroidism; the latter is due to secretion of immunoactive but bioinactive forms of TSH. The diagnosis is confirmed by detecting a low unbound T₄ level. The goal of treatment is to maintain T₄ levels in the upper half of the reference range, because TSH levels cannot be used to monitor therapy.

TREATMENT HYPOTHYROIDISM

CLINICAL HYPOTHYROIDISM

If there is no residual thyroid function, the daily replacement dose of levothyroxine is usually 1.6 µg/kg body weight (typically 100–150 µg), ideally taken at least 30 min before breakfast. In many patients,

however, lower doses suffice until residual thyroid tissue is destroyed. In patients who develop hypothyroidism after the treatment of Graves' disease, there is often underlying autonomous function, necessitating lower replacement doses (typically 75–125 µg/d).

Adult patients under 60 years old without evidence of heart disease may be started on 50–100 µg levothyroxine (T₄) daily. The dose is adjusted on the basis of TSH levels, with the goal of treatment being a normal TSH, ideally in the lower half of the reference range. TSH responses are gradual and should be measured about 2 months after instituting treatment or after any subsequent change in levothyroxine dosage. The clinical effects of levothyroxine replacement are slow to appear. Patients may not experience full relief from symptoms until 3–6 months after normal TSH levels are restored. Adjustment of levothyroxine dosage is made in 12.5- or 25-µg increments if the TSH is high; decrements of the same magnitude should be

made if the TSH is suppressed. Patients with a suppressed TSH of any cause, including T₄ overtreatment, have an increased risk of atrial fibrillation and reduced bone density.

Although desiccated animal thyroid preparations (thyroid extract USP) are available, they are not recommended because the ratio of T₃ to T₄ is nonphysiologic. The use of levothyroxine combined with liothyronine (triiodothyronine, T₃) has been investigated, but benefit has not been confirmed in prospective studies. There is no place for liothyronine alone as long-term replacement, because the short half-life necessitates three or four daily doses and is associated with fluctuating T₃ levels.

Once full replacement is achieved and TSH levels are stable, follow-up measurement of TSH is recommended at annual intervals and may be extended to every 2–3 years if a normal TSH is maintained over several years. It is important to ensure ongoing adherence, however, as patients do not feel any symptomatic difference after missing a few doses of levothyroxine, and this sometimes leads to self-discontinuation.

In patients of normal body weight who are taking ≥200 µg of levothyroxine per day, an elevated TSH level is often a sign of poor adherence to treatment. This is also the likely explanation for fluctuating TSH levels, despite a constant levothyroxine dosage. Such patients often have normal or high unbound T₄ levels, despite an elevated TSH, because they remember to take medication for a few days before testing; this is sufficient to normalize T₄, but not TSH levels. It is important to consider variable adherence, because this pattern of thyroid function tests is otherwise suggestive of disorders associated with inappropriate TSH secretion (Table 405-3). Because T₄ has a long half-life (7 days), patients who miss a dose can be advised to take two doses of the skipped tablets at once. Other causes of increased levothyroxine requirements must be excluded, particularly malabsorption (e.g., celiac disease, small-bowel surgery), estrogen or selective estrogen receptor modulator therapy, ingestion with a meal, and drugs that interfere with T₄ absorption or metabolism such as cholestyramine, ferrous sulfate, calcium supplements, proton pump inhibitors, lovastatin, aluminum hydroxide, rifampicin, amiodarone, carbamazepine, phenytoin, and tyrosine kinase inhibitors.

SUBCLINICAL HYPOTHYROIDISM

By definition, subclinical hypothyroidism refers to biochemical evidence of thyroid hormone deficiency in patients who have few or no apparent clinical features of hypothyroidism. There are no

universally accepted recommendations for the management of subclinical hypothyroidism, but levothyroxine is recommended if the patient is a woman who wishes to conceive or is pregnant, or when TSH levels are above 10 mIU/L. When TSH levels are below 10 mIU/L, treatment should be considered when patients have suggestive symptoms of hypothyroidism, positive TPO antibodies, or any evidence of heart disease. It is important to confirm that any elevation of TSH is sustained over a 3-month period before treatment is given. As long as excessive treatment is avoided, there is no risk in correcting a slightly increased TSH. Treatment is administered by starting with a low dose of levothyroxine (25–50 µg/d) with the goal of normalizing TSH. If levothyroxine is not given, thyroid function should be evaluated annually.

SPECIAL TREATMENT CONSIDERATIONS

Rarely, levothyroxine replacement is associated with pseudotumor cerebri in children. Presentation appears to be idiosyncratic and occurs months after treatment has begun.

Women with a history or high risk of hypothyroidism should ensure that they are euthyroid prior to conception and during early pregnancy because maternal hypothyroidism may adversely affect fetal neural development and cause preterm delivery. The presence of thyroid autoantibodies alone, in a euthyroid patient, is also associated with miscarriage and preterm delivery; it is unclear if levothyroxine therapy improves outcomes. Thyroid function should be evaluated immediately after pregnancy is confirmed and every 4 weeks during the first half of the pregnancy, with less frequent testing after 20 weeks' gestation (every 6–8 weeks depending on whether levothyroxine dose adjustment is ongoing). The levothyroxine dose may need to be increased by up to 50% during pregnancy, with a goal TSH of less than 2.5 mIU/L during the first trimester and less than 3.0 mIU/L during the second and third trimesters. After delivery, thyroxine doses typically return to prepregnancy levels. Pregnant women should be counseled to separate ingestion of prenatal vitamins and iron supplements from levothyroxine by at least 4 h.

Elderly patients may require 20% less thyroxine than younger patients. In the elderly, especially patients with known coronary artery disease, the starting dose of levothyroxine is 12.5–25 µg/d with similar increments every 2–3 months until TSH is normalized. In some patients, it may be impossible to achieve full replacement despite optimal antianginal treatment. *Emergency surgery* is generally safe in patients with untreated hypothyroidism, although routine surgery in a hypothyroid patient should be deferred until euthyroidism is achieved.

Myxedema coma still has a 20–40% mortality rate, despite intensive treatment, and outcomes are independent of the T₄ and TSH levels. Clinical manifestations include reduced level of consciousness, sometimes associated with seizures, as well as the other features of hypothyroidism (Table 405-6). Hypothermia can reach 23°C (74°F). There may be a history of treated hypothyroidism with poor compliance, or the patient may be previously undiagnosed. Myxedema coma almost always occurs in the elderly and is usually precipitated by factors that impair respiration, such as drugs (especially sedatives, anesthetics, and antidepressants), pneumonia, congestive heart failure, myocardial infarction, gastrointestinal bleeding, or cerebrovascular accidents. Sepsis should also be suspected. Exposure to cold may also be a risk factor. Hypoventilation, leading to hypoxia and hypercapnia, plays a major role in pathogenesis; hypoglycemia and dilutional hyponatremia also contribute to the development of myxedema coma.

Levothyroxine can initially be administered as a single IV bolus of 500 µg, which serves as a loading dose. Although further levothyroxine is not strictly necessary for several days, it is usually continued at a dose of 50–100 µg/d. If suitable IV preparation is not available, the same initial dose of levothyroxine can be given by nasogastric tube (although absorption may be impaired in myxedema). An alternative is to give liothyronine (T₃) intravenously or via nasogastric tube, in doses ranging from 10 to 25 µg every 8–12 h. This treatment has been advocated because T₄ → T₃ conversion is impaired in myxedema

coma. However, excess liothyronine has the potential to provoke arrhythmias. Another option is to combine levothyroxine (200 µg) and liothyronine (25 µg) as a single, initial IV bolus followed by daily treatment with levothyroxine (50–100 µg/d) and liothyronine (10 µg every 8 h).

Supportive therapy should be provided to correct any associated metabolic disturbances. External warming is indicated only if the temperature is <30°C, as it can result in cardiovascular collapse (Chap. 478e). Space blankets should be used to prevent further heat loss. Parenteral hydrocortisone (50 mg every 6 h) should be administered, because there is impaired adrenal reserve in profound hypothyroidism. Any precipitating factors should be treated, including the early use of broad-spectrum antibiotics, pending the exclusion of infection. Ventilatory support with regular blood gas analysis is usually needed during the first 48 h. Hypertonic saline or IV glucose may be needed if there is severe hyponatremia or hypoglycemia; hypotonic IV fluids should be avoided because they may exacerbate water retention secondary to reduced renal perfusion and inappropriate vasopressin secretion. The metabolism of most medications is impaired, and sedatives should be avoided if possible or used in reduced doses. Medication blood levels should be monitored, when available, to guide dosage.

THYROTOXICOSIS

Thyrotoxicosis is defined as the state of thyroid hormone excess and is not synonymous with *hyperthyroidism*, which is the result of excessive thyroid function. However, the major etiologies of thyrotoxicosis are hyperthyroidism caused by Graves' disease, toxic MNG, and toxic adenomas. Other causes are listed in Table 405-7.

GRAVES' DISEASE

Epidemiology Graves' disease accounts for 60–80% of thyrotoxicosis. The prevalence varies among populations, reflecting genetic factors and iodine intake (high iodine intake is associated with an increased prevalence of Graves' disease). Graves' disease occurs in up to 2% of women but is one-tenth as frequent in men. The disorder rarely begins before adolescence and typically occurs between 20 and 50 years of age; it also occurs in the elderly.

TABLE 405-7 CAUSES OF THYROTOXICOSIS

Primary Hyperthyroidism
Graves' disease
Toxic multinodular goiter
Toxic adenoma
Functioning thyroid carcinoma metastases
Activating mutation of the TSH receptor
Activating mutation of G _{sα} (McCune-Albright syndrome)
Struma ovarii
Drugs: iodine excess (Jod-Basedow phenomenon)
Thyrotoxicosis Without Hyperthyroidism
Subacute thyroiditis
Silent thyroiditis
Other causes of thyroid destruction: amiodarone, radiation, infarction of adenoma
Ingestion of excess thyroid hormone (thyrotoxicosis factitia) or thyroid tissue
Secondary Hyperthyroidism
TSH-secreting pituitary adenoma
Thyroid hormone resistance syndrome: occasional patients may have features of thyrotoxicosis
Chorionic gonadotropin-secreting tumors ^a
Gestational thyrotoxicosis ^a

^aCirculating TSH levels are low in these forms of secondary hyperthyroidism.

Abbreviations: TSH, thyroid-stimulating hormone.

2294 Pathogenesis As in autoimmune hypothyroidism, a combination of environmental and genetic factors, including polymorphisms in HLA-DR, the immunoregulatory genes *CTLA-4*, *CD25*, *PTPN22*, *FCRL3*, and *CD226*, as well as the TSH-R, contribute to Graves' disease susceptibility. The concordance for Graves' disease in monozygotic twins is 20–30%, compared to <5% in dizygotic twins. Indirect evidence suggests that stress is an important environmental factor, presumably operating through neuroendocrine effects on the immune system. Smoking is a minor risk factor for Graves' disease and a major risk factor for the development of ophthalmopathy. Sudden increases in iodine intake may precipitate Graves' disease, and there is a three-fold increase in the occurrence of Graves' disease in the postpartum period. Graves' disease may occur during the immune reconstitution phase after highly active antiretroviral therapy (HAART) or alemtuzumab treatment.

The hyperthyroidism of Graves' disease is caused by TSI that are synthesized in the thyroid gland as well as in bone marrow and lymph nodes. Such antibodies can be detected by bioassays or by using the more widely available TBII assays. The presence of TBII in a patient with thyrotoxicosis implies the existence of TSI, and these assays are useful in monitoring pregnant Graves' patients in whom high levels of TSI can cross the placenta and cause neonatal thyrotoxicosis. Other thyroid autoimmune responses, similar to those in autoimmune hypothyroidism (see above), occur concurrently in patients with Graves' disease. In particular, TPO antibodies occur in up to 80% of cases and serve as a readily measurable marker of autoimmunity. Because the coexisting thyroiditis can also affect thyroid function, there is no direct correlation between the level of TSI and thyroid hormone levels in Graves' disease. In the long term, spontaneous autoimmune hypothyroidism may develop in up to 15% of patients with Graves' disease.

Cytokines appear to play a major role in thyroid-associated ophthalmopathy. There is infiltration of the extraocular muscles by activated T cells; the release of cytokines such as IFN- γ , TNF, and IL-1 results in fibroblast activation and increased synthesis of glycosaminoglycans that trap water, thereby leading to characteristic muscle swelling. Late in the disease, there is irreversible fibrosis of the muscles. Orbital fibroblasts may be particularly sensitive to cytokines, perhaps explaining the anatomic localization of the immune response. Though the pathogenesis of thyroid-associated ophthalmopathy remains unclear, there is mounting evidence that the TSH-R may be a shared autoantigen that is expressed in the orbit; this would explain the close association with autoimmune thyroid disease. Increased fat is an additional cause of retrobulbar tissue expansion. The increase in intraorbital pressure can lead to proptosis, diplopia, and optic neuropathy.

Clinical Manifestations Signs and symptoms include features that are common to any cause of thyrotoxicosis (Table 405-8) as well as those specific for Graves' disease. The clinical presentation depends on the severity of thyrotoxicosis, the duration of disease, individual susceptibility to excess thyroid hormone, and the patient's age. In the elderly, features of thyrotoxicosis may be subtle or masked, and patients may present mainly with fatigue and weight loss, a condition known as *apathetic thyrotoxicosis*.

Thyrotoxicosis may cause unexplained weight loss, despite an enhanced appetite, due to the increased metabolic rate. Weight gain occurs in 5% of patients, however, because of increased food intake. Other prominent features include hyperactivity, nervousness, and irritability, ultimately leading to a sense of easy fatigability in some patients. Insomnia and impaired concentration are common; apathetic thyrotoxicosis may be mistaken for depression in the elderly. Fine tremor is a frequent finding, best elicited by having patients stretch out their fingers while feeling the fingertips with the palm. Common neurologic manifestations include hyperreflexia, muscle wasting, and proximal myopathy without fasciculation. Chorea is rare. Thyrotoxicosis is sometimes associated with a form of hypokalemic periodic paralysis; this disorder is particularly common in Asian males with thyrotoxicosis, but it occurs in other ethnic groups as well.

The most common cardiovascular manifestation is sinus tachycardia, often associated with palpitations, occasionally caused by supraventricular tachycardia. The high cardiac output produces a bounding pulse, widened pulse pressure, and an aortic systolic murmur and can lead to worsening of angina or heart failure in the elderly or those with preexisting heart disease. Atrial fibrillation is more common in patients >50 years of age. Treatment of the thyrotoxic state alone converts atrial fibrillation to normal sinus rhythm in about half of patients, suggesting the existence of an underlying cardiac problem in the remainder.

The skin is usually warm and moist, and the patient may complain of sweating and heat intolerance, particularly during warm weather. Palmar erythema, onycholysis, and, less commonly, pruritus, urticaria, and diffuse hyperpigmentation may be evident. Hair texture may become fine, and a diffuse alopecia occurs in up to 40% of patients, persisting for months after restoration of euthyroidism. Gastrointestinal transit time is decreased, leading to increased stool frequency, often with diarrhea and occasionally mild steatorrhea. Women frequently experience oligomenorrhea or amenorrhea; in men, there may be impaired sexual function and, rarely, gynecomastia. The direct effect of thyroid hormones on bone resorption leads to osteopenia in long-standing thyrotoxicosis; mild hypercalcemia occurs in up to 20% of patients, but hypercalciuria is more common. There is a small increase in fracture rate in patients with a previous history of thyrotoxicosis.

In Graves' disease, the thyroid is usually diffusely enlarged to two to three times its normal size. The consistency is firm, but not nodular. There may be a thrill or bruit, best detected at the inferolateral margins of the thyroid lobes, due to the increased vascularity of the gland and the hyperdynamic circulation.

Lid retraction, causing a staring appearance, can occur in any form of thyrotoxicosis and is the result of sympathetic overactivity. However, Graves' disease is associated with specific eye signs that comprise *Graves' ophthalmopathy* (Fig. 405-8A). This condition is also called *thyroid-associated ophthalmopathy*, because it occurs in the absence of hyperthyroidism in 10% of patients. Most of these individuals have autoimmune hypothyroidism or thyroid antibodies. The onset of Graves' ophthalmopathy occurs within the year before or after the diagnosis of thyrotoxicosis in 75% of patients but can sometimes precede or follow thyrotoxicosis by several years, accounting for some cases of euthyroid ophthalmopathy.

Some patients with Graves' disease have little clinical evidence of ophthalmopathy. However, the enlarged extraocular muscles typical of the disease, and other subtle features, can be detected in almost all patients when investigated by ultrasound or computed tomography (CT) imaging of the orbits. Unilateral signs are found in up to 10% of patients. The earliest manifestations of ophthalmopathy are usually a sensation of grittiness, eye discomfort, and excess tearing. About one-third of patients have proptosis, best detected by visualization of the sclera between the lower border of the iris and the lower eyelid, with the eyes in the primary position. Proptosis can be measured using an exophthalmometer. In severe cases, proptosis may cause corneal exposure and damage, especially if the lids fail to close during sleep. Periorbital edema, scleral injection, and chemosis are also frequent. In 5–10% of patients, the muscle swelling is so severe that diplopia results, typically, but not exclusively, when the patient looks up and laterally. The most serious manifestation is compression of the optic nerve at

TABLE 405-8 SIGNS AND SYMPTOMS OF THYROTOXICOSIS (DESCENDING ORDER OF FREQUENCY)

Symptoms	Signs ^a
Hyperactivity, irritability, dysphoria	Tachycardia; atrial fibrillation in the elderly
Heat intolerance and sweating	Tremor
Palpitations	Goiter
Fatigue and weakness	Warm, moist skin
Weight loss with increased appetite	Muscle weakness, proximal myopathy
Diarrhea	Lid retraction or lag
Polyuria	Gynecomastia
Oligomenorrhea, loss of libido	

^aExcludes the signs of ophthalmopathy and dermopathy specific for Graves' disease.



FIGURE 405-8 Features of Graves' disease. **A.** Ophthalmopathy in Graves' disease; lid retraction, periorbital edema, conjunctival injection, and proptosis are marked. **B.** Thyroid dermopathy over the lateral aspects of the shins. **C.** Thyroid acropachy.

the apex of the orbit, leading to papilledema; peripheral field defects; and, if left untreated, permanent loss of vision.

The "NO SPECS" scoring system to evaluate ophthalmopathy is an acronym derived from the following changes:

- 0 = No signs or symptoms
- 1 = Only signs (lid retraction or lag), no symptoms
- 2 = Soft tissue involvement (periorbital edema)
- 3 = Proptosis (>22 mm)
- 4 = Extraocular muscle involvement (diplopia)
- 5 = Corneal involvement
- 6 = Sight loss

Although useful as a mnemonic, the NO SPECS scheme is inadequate to describe the eye disease fully, and patients do not necessarily progress from one class to another; alternative scoring systems that assess disease activity are preferable for monitoring purposes. When Graves' eye disease is active and severe, referral to an ophthalmologist is indicated and objective measurements are needed, such as lid-fissure width; corneal staining with fluorescein; and evaluation of extraocular muscle function (e.g., Hess chart), intraocular pressure and visual fields, acuity, and color vision.

Thyroid dermopathy occurs in <5% of patients with Graves' disease (Fig. 405-8B), almost always in the presence of moderate or severe ophthalmopathy. Although most frequent over the anterior and lateral aspects of the lower leg (hence the term *pretibial myxedema*), skin changes can occur at other sites,

particularly after trauma. The typical lesion is a noninflamed, indurated plaque with a deep pink or purple color and an "orange skin" appearance. Nodular involvement can occur, and the condition can rarely extend over the whole lower leg and foot, mimicking elephantiasis. *Thyroid acropachy* refers to a form of clubbing found in <1% of patients with Graves' disease (Fig. 405-8C). It is so strongly associated with thyroid dermopathy that an alternative cause of clubbing should be sought in a Graves' patient without coincident skin and orbital involvement.

Laboratory Evaluation Investigations used to determine the existence and cause of thyrotoxicosis are summarized in Fig. 405-9. In Graves' disease, the TSH level is suppressed, and total and unbound thyroid hormone levels are increased. In 2–5% of patients (and more in areas of borderline iodine intake), only T_3 is increased (T_3 toxicosis). The converse state of T_4 toxicosis, with elevated total and unbound T_4 and normal T_3 levels, is occasionally seen when hyperthyroidism is induced by excess iodine, providing surplus substrate for thyroid hormone synthesis. Measurement of TPO antibodies or TRAb may be useful if the diagnosis is unclear clinically but is not needed routinely. Associated abnormalities that may cause diagnostic confusion in thyrotoxicosis include elevation of bilirubin, liver enzymes, and ferritin. Microcytic anemia and thrombocytopenia may occur.

Differential Diagnosis Diagnosis of Graves' disease is straightforward in a patient with biochemically confirmed thyrotoxicosis, diffuse goiter on palpation, ophthalmopathy, and often a personal or family history of autoimmune disorders. For patients with thyrotoxicosis who lack these features, the diagnosis is generally established by a radionuclide (^{99m}Tc , ^{123}I , or ^{131}I) scan and uptake of the thyroid, which will distinguish the diffuse, high uptake of Graves' disease from destructive thyroiditis, ectopic thyroid tissue, and factitious thyrotoxicosis. Scintigraphy is the preferred diagnostic test; however, TRAb measurement can be used to assess autoimmune activity. In secondary hyperthyroidism due to a

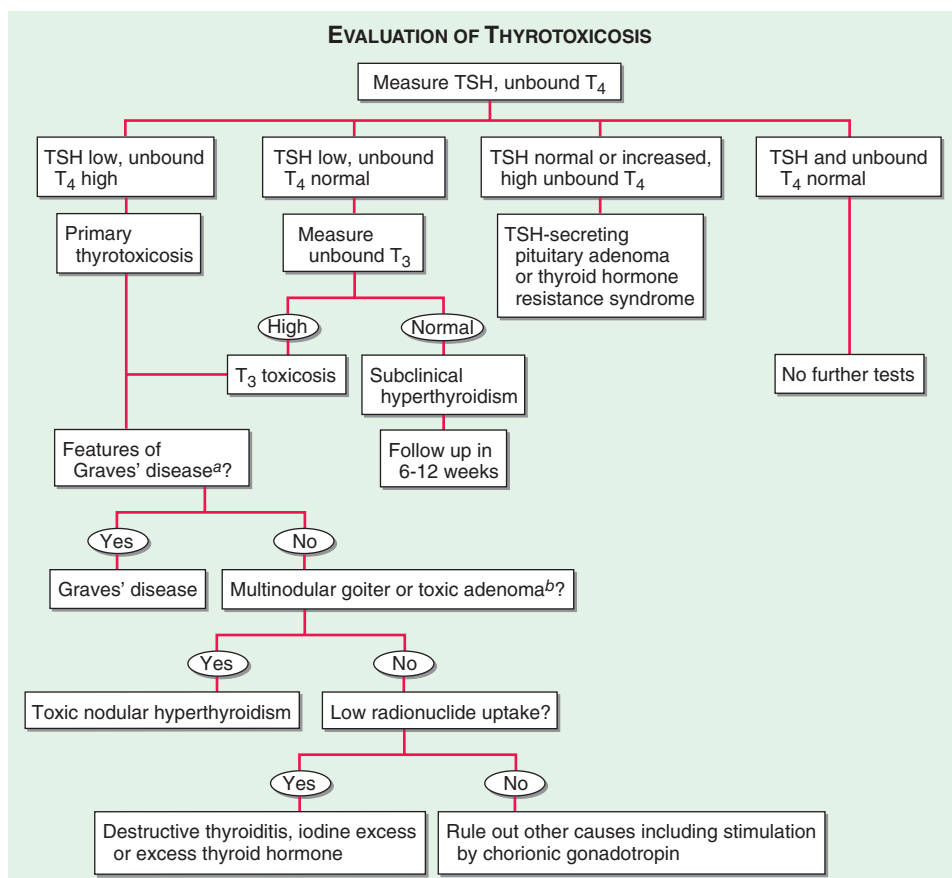


FIGURE 405-9 Evaluation of thyrotoxicosis. ^aDiffuse goiter, positive TPO antibodies or TRAb, ophthalmopathy, dermopathy. ^bCan be confirmed by radionuclide scan. TSH, thyroid-stimulating hormone.

2296 TSH-secreting pituitary tumor, there is also a diffuse goiter. The presence of a nonsuppressed TSH level and the finding of a pituitary tumor on CT or magnetic resonance scan (MRI) scan suggest this diagnosis.

Clinical features of thyrotoxicosis can mimic certain aspects of other disorders, including panic attacks, mania, pheochromocytoma, and weight loss associated with malignancy. The diagnosis of thyrotoxicosis can be easily excluded if the TSH and unbound T_4 and T_3 levels are normal. A normal TSH also excludes Graves' disease as a cause of diffuse goiter.

Clinical Course Clinical features generally worsen without treatment; mortality was 10–30% before the introduction of satisfactory therapy. Some patients with mild Graves' disease experience spontaneous relapses and remissions. Rarely, there may be fluctuation between hypo- and hyperthyroidism due to changes in the functional activity of TSH-R antibodies. About 15% of patients who enter remission after treatment develop hypothyroidism 10–15 years later as a result of the destructive autoimmune process.

The clinical course of ophthalmopathy does not follow that of the thyroid disease. Ophthalmopathy typically worsens over the initial 3–6 months, followed by a plateau phase over the next 12–18 months, with spontaneous improvement, particularly in the soft tissue changes. However, the course is more fulminant in up to 5% of patients, requiring intervention in the acute phase if there is optic nerve compression or corneal ulceration. Diplopia may appear late in the disease due to fibrosis of the extraocular muscles. Radioiodine treatment for hyperthyroidism worsens the eye disease in a small proportion of patients (especially smokers). Antithyroid drugs or surgery have no adverse effects on the clinical course of ophthalmopathy. Thyroid dermopathy, when it occurs, usually appears 1–2 years after the development of Graves' hyperthyroidism; it may improve spontaneously.

TREATMENT GRAVES' DISEASE

The *hyperthyroidism* of Graves' disease is treated by reducing thyroid hormone synthesis, using antithyroid drugs, or reducing the amount of thyroid tissue with radioiodine (^{131}I) treatment or by thyroidectomy. Antithyroid drugs are the predominant therapy in many centers in Europe and Japan, whereas radioiodine is more often the first line of treatment in North America. These differences reflect the fact that no single approach is optimal and that patients may require multiple treatments to achieve remission.

The main *antithyroid drugs* are the thionamides, such as propylthiouracil, carbimazole (not available in the United States), and the active metabolite of the latter, methimazole. All inhibit the function of TPO, reducing oxidation and organification of iodide. These drugs also reduce thyroid antibody levels by mechanisms that remain unclear, and they appear to enhance rates of remission. Propylthiouracil inhibits deiodination of $T_4 \rightarrow T_3$. However, this effect is of minor benefit, except in the most severe thyrotoxicosis, and is offset by the much shorter half-life of this drug (90 min) compared to methimazole (6 h). Due to the hepatotoxicity of propylthiouracil, the U.S. Food and Drug Administration (FDA) has limited indications for its use to the first trimester of pregnancy, the treatment of thyroid storm, and patients with minor adverse reactions to methimazole. If propylthiouracil is used, monitoring of liver function tests is recommended.

There are many variations of antithyroid drug regimens. The initial dose of carbimazole or methimazole is usually 10–20 mg every 8 or 12 h, but once-daily dosing is possible after euthyroidism is restored. Propylthiouracil is given at a dose of 100–200 mg every 6–8 h, and divided doses are usually given throughout the course. Lower doses of each drug may suffice in areas of low iodine intake. The starting dose of antithyroid drugs can be gradually reduced (titration regimen) as thyrotoxicosis improves. Alternatively, high doses may be given combined with levothyroxine supplementation (block-replace regimen) to avoid drug-induced hypothyroidism. The titration regimen is preferred to minimize the dose of antithyroid drug and provide an index of treatment response.

Thyroid function tests and clinical manifestations are reviewed 4–6 weeks after starting treatment, and the dose is titrated based on unbound T_4 levels. Most patients do not achieve euthyroidism until 6–8 weeks after treatment is initiated. TSH levels often remain suppressed for several months and therefore do not provide a sensitive index of treatment response. The usual daily maintenance doses of antithyroid drugs in the titration regimen are 2.5–10 mg of carbimazole or methimazole and 50–100 mg of propylthiouracil. In the block-replace regimen, the initial dose of antithyroid drug is held constant, and the dose of levothyroxine is adjusted to maintain normal unbound T_4 levels. When TSH suppression is alleviated, TSH levels can also be used to monitor therapy.

Maximum remission rates (up to 30–60% in some populations) are achieved by 12–18 months for the titration regimen and by 6 months for the block-replace regimen. For unclear reasons, remission rates appear to vary in different geographic regions. Younger patients, males, smokers, and patients with severe hyperthyroidism and large goiters are most likely to relapse when treatment stops, but outcomes are difficult to predict. All patients should be followed closely for relapse during the first year after treatment and at least annually thereafter.

The common minor side effects of antithyroid drugs are rash, urticaria, fever, and arthralgia (1–5% of patients). These may resolve spontaneously or after substituting an alternative antithyroid drug. Rare but major side effects include hepatitis (propylthiouracil; avoid use in children) and cholestasis (methimazole and carbimazole); an SLE-like syndrome; and, most important, agranulocytosis (<1%). It is essential that antithyroid drugs are stopped and not restarted if a patient develops major side effects. Written instructions should be provided regarding the symptoms of possible agranulocytosis (e.g., sore throat, fever, mouth ulcers) and the need to stop treatment pending an urgent complete blood count to confirm that agranulocytosis is not present. Management of agranulocytosis is described in [Chap. 130](#). It is not useful to monitor blood counts prospectively, because the onset of agranulocytosis is idiosyncratic and abrupt.

Propranolol (20–40 mg every 6 h) or longer-acting selective β_1 receptor blockers such as atenolol may be helpful to control adrenergic symptoms, especially in the early stages before antithyroid drugs take effect. Beta blockers are also useful in patients with thyrotoxic periodic paralysis, pending correction of thyrotoxicosis. In consultation with a cardiologist, anticoagulation with warfarin should be considered in all patients with atrial fibrillation who often spontaneously revert to sinus rhythm with control of hyperthyroidism. Decreased warfarin doses are required when patients are thyrotoxic. If digoxin is used, increased doses are often needed in the thyrotoxic state.

Radioiodine causes progressive destruction of thyroid cells and can be used as initial treatment or for relapses after a trial of antithyroid drugs. There is a small risk of thyrotoxic crisis (see below) after radioiodine, which can be minimized by pretreatment with antithyroid drugs for at least a month before treatment. Antecedent treatment with antithyroid drugs should be considered for all elderly patients or for those with cardiac problems to deplete thyroid hormone stores before administration of radioiodine. Carbimazole or methimazole must be stopped 3–5 days before radioiodine administration to achieve optimum iodine uptake. Propylthiouracil appears to have a prolonged radioprotective effect and should be stopped for a longer period before radioiodine is given, or a larger dose of radioiodine will be necessary.

Efforts to calculate an optimal dose of radioiodine that achieves euthyroidism without a high incidence of relapse or progression to hypothyroidism have not been successful. Some patients inevitably relapse after a single dose because the biologic effects of radiation vary between individuals, and hypothyroidism cannot be uniformly avoided even using accurate dosimetry. A practical strategy is to give a fixed dose based on clinical features, such as the severity of thyrotoxicosis, the size of the goiter (increases the dose needed), and the level of radioiodine uptake (decreases the dose needed). ^{131}I dosage generally ranges between 370 MBq (10 mCi) and 555 MBq

(15 mCi). Most authorities favor an approach aimed at thyroid ablation (as opposed to euthyroidism), given that levothyroxine replacement is straightforward and most patients ultimately progress to hypothyroidism over 5–10 years, frequently with some delay in the diagnosis of hypothyroidism.

Certain radiation safety precautions are necessary in the first few days after radioiodine treatment, but the exact guidelines vary depending on local protocols. In general, patients need to avoid close, prolonged contact with children and pregnant women for 5–7 days because of possible transmission of residual isotope and exposure to radiation emanating from the gland. Rarely, there may be mild pain due to radiation thyroiditis 1–2 weeks after treatment. Hyperthyroidism can persist for 2–3 months before radioiodine takes full effect. For this reason, β -adrenergic blockers or antithyroid drugs can be used to control symptoms during this interval. Persistent hyperthyroidism can be treated with a second dose of radioiodine, usually 6 months after the first dose. The risk of hypothyroidism after radioiodine depends on the dosage but is at least 10–20% in the first year and 5% per year thereafter. Patients should be informed of this possibility before treatment and require close follow-up during the first year followed by annual thyroid function testing.

Pregnancy and breast-feeding are absolute contraindications to radioiodine treatment, but patients can conceive safely 6 months after treatment. The presence of severe ophthalmopathy requires caution, and some authorities advocate the use of prednisone, 40 mg/d, at the time of radioiodine treatment, tapered over 6–12 weeks to prevent exacerbation of ophthalmopathy. The overall risk of cancer after radioiodine treatment in adults is not increased. Although many physicians avoid radioiodine in children and adolescents because of the theoretical risks of malignancy, emerging evidence suggests that radioiodine can be used safely in older children.

Subtotal or near-total thyroidectomy is an option for patients who relapse after antithyroid drugs and prefer this treatment to radioiodine. Some experts recommend surgery in young individuals, particularly when the goiter is very large. Careful control of thyrotoxicosis with antithyroid drugs, followed by potassium iodide (3 drops SSKI orally tid), is needed prior to surgery to avoid thyrotoxic crisis and to reduce the vascularity of the gland. The major complications of surgery—bleeding, laryngeal edema, hypoparathyroidism, and damage to the recurrent laryngeal nerves—are unusual when the procedure is performed by highly experienced surgeons. Recurrence rates in the best series are <2%, but the rate of hypothyroidism is only slightly less than that following radioiodine treatment.

The titration regimen of antithyroid drugs should be used to manage Graves' disease in pregnancy because transplacental passage of these drugs may produce fetal hypothyroidism and goiter if the maternal dose is excessive. If available, propylthiouracil should be used in early gestation because of the association of rare cases of fetal *aplasia cutis* and other defects, such as choanal atresia with carbimazole and methimazole. As noted above, because of its rare association with hepatotoxicity, propylthiouracil should be limited to the first trimester and then maternal therapy should be converted to methimazole (or carbimazole) at a ratio of 15–20 mg of propylthiouracil to 1 mg of methimazole. The lowest effective antithyroid drug dose should be used throughout gestation to maintain the maternal serum free T_4 level at the upper limit of the nonpregnant normal reference range. It is often possible to stop treatment in the last trimester because TSIs tend to decline in pregnancy. Nonetheless, the transplacental transfer of these antibodies rarely causes *fetal or neonatal thyrotoxicosis*. Poor intrauterine growth, a fetal heart rate of >160 beats/min, and high levels of maternal TSI in the last trimester may herald this complication. Antithyroid drugs given to the mother can be used to treat the fetus and may be needed for 1–3 months after delivery, until the maternal antibodies disappear from the baby's circulation. The postpartum period is a time of major risk for relapse of Graves' disease. Breast-feeding is safe with low doses of antithyroid drugs. Graves' disease in children is usually managed with methimazole or

carbimazole (avoid propylthiouracil), often given as a prolonged course of the titration regimen. Surgery or radioiodine may be indicated for severe disease.

Thyrotoxic crisis, or thyroid storm, is rare and presents as a life-threatening exacerbation of hyperthyroidism, accompanied by fever, delirium, seizures, coma, vomiting, diarrhea, and jaundice. The mortality rate due to cardiac failure, arrhythmia, or hyperthermia is as high as 30%, even with treatment. Thyrotoxic crisis is usually precipitated by acute illness (e.g., stroke, infection, trauma, diabetic ketoacidosis), surgery (especially on the thyroid), or radioiodine treatment of a patient with partially treated or untreated hyperthyroidism. Management requires intensive monitoring and supportive care, identification and treatment of the precipitating cause, and measures that reduce thyroid hormone synthesis. Large doses of propylthiouracil (500–1000 mg loading dose and 250 mg every 4 h) should be given orally or by nasogastric tube or per rectum; the drug's inhibitory action on $T_4 \rightarrow T_3$ conversion makes it the antithyroid drug of choice. If not available, methimazole can be used in doses up to 30 mg every 12 h. One hour after the first dose of propylthiouracil, stable iodide is given to block thyroid hormone synthesis via the Wolff-Chaikoff effect (the delay allows the antithyroid drug to prevent the excess iodine from being incorporated into new hormone). A saturated solution of potassium iodide (5 drops SSKI every 6 h) or, where available, ipodate or iopanoic acid (500 mg per 12 h) may be given orally. Sodium iodide, 0.25 g IV every 6 h, is an alternative but is not generally available. Propranolol should also be given to reduce tachycardia and other adrenergic manifestations (60–80 mg PO every 4 h; or 2 mg IV every 4 h). Although other β -adrenergic blockers can be used, high doses of propranolol decrease $T_4 \rightarrow T_3$ conversion, and the doses can be easily adjusted. Caution is needed to avoid acute negative inotropic effects, but controlling the heart rate is important, as some patients develop a form of high-output heart failure. Short-acting IV esmolol can be used to decrease heart rate while monitoring for signs of heart failure. Additional therapeutic measures include glucocorticoids (e.g., hydrocortisone 300 mg IV bolus, then 100 mg every 8 h), antibiotics if infection is present, cooling, oxygen, and IV fluids.

Ophthalmopathy requires no active treatment when it is mild or moderate, because there is usually spontaneous improvement. General measures include meticulous control of thyroid hormone levels, cessation of smoking, and an explanation of the natural history of ophthalmopathy. Discomfort can be relieved with artificial tears (e.g., 1% methylcellulose), eye ointment, and the use of dark glasses with side frames. Periorbital edema may respond to a more upright sleeping position or a diuretic. Corneal exposure during sleep can be avoided by using patches or taping the eyelids shut. Minor degrees of diplopia improve with prisms fitted to spectacles. Severe ophthalmopathy, with optic nerve involvement or chemosis resulting in corneal damage, is an emergency requiring joint management with an ophthalmologist. Pulse therapy with IV methylprednisolone (e.g., 500 mg of methylprednisolone once weekly for 6 weeks, then 250 mg once weekly for 6 weeks) is preferable to oral glucocorticoids, which are used for moderately active disease. When glucocorticoids are ineffective, orbital decompression can be achieved by removing bone from any wall of the orbit, thereby allowing displacement of fat and swollen extraocular muscles. The transantral route is used most often because it requires no external incision. Proptosis recedes an average of 5 mm, but there may be residual or even worsened diplopia. Once the eye disease has stabilized, surgery may be indicated for relief of diplopia and correction of the appearance. External beam radiotherapy of the orbits has been used for many years, but the efficacy of this therapy remains unclear, and it is best reserved for those with moderately active disease who have failed or are not candidates for glucocorticoid therapy. Other immunosuppressive agents such as rituximab have shown some benefit, but their role is yet to be established.

Thyroid dermatopathy does not usually require treatment, but it can cause cosmetic problems or interfere with the fit of shoes. Surgical

2298 removal is not indicated. If necessary, treatment consists of topical, high-potency glucocorticoid ointment under an occlusive dressing. Octreotide may be beneficial in some cases.

OTHER CAUSES OF THYROTOXICOSIS

Destructive thyroiditis (subacute or silent thyroiditis) typically presents with a short thyrotoxic phase due to the release of preformed thyroid hormones and catabolism of Tg (see “Subacute Thyroiditis,” below). True hyperthyroidism is absent, as demonstrated by a low radionuclide uptake. Circulating Tg levels are usually increased. Other causes of thyrotoxicosis with low or absent thyroid radionuclide uptake include *thyrotoxicosis factitia*, iodine excess, and, rarely, ectopic thyroid tissue, particularly teratomas of the ovary (*struma ovarii*) and functional metastatic follicular carcinoma. Whole-body radionuclide studies can demonstrate ectopic thyroid tissue, and thyrotoxicosis factitia can be distinguished from destructive thyroiditis by the clinical features and low levels of Tg. Amiodarone treatment is associated with thyrotoxicosis in up to 10% of patients, particularly in areas of low iodine intake (see below).

TSH-secreting pituitary adenoma is a rare cause of thyrotoxicosis. It is characterized by the presence of an inappropriately normal or increased TSH level in a patient with hyperthyroidism, diffuse goiter, and elevated T₄ and T₃ levels (Chap. 403). Elevated levels of the α -subunit of TSH, released by the TSH-secreting adenoma, support this diagnosis, which can be confirmed by demonstrating the pituitary tumor on MRI or CT scan. A combination of transphenoidal surgery, sella irradiation, and octreotide may be required to normalize TSH, because many of these tumors are large and locally invasive at the time of diagnosis. Radioiodine or antithyroid drugs can be used to control thyrotoxicosis.

Thyrotoxicosis caused by *toxic MNG* and *hyperfunctioning solitary nodules* is discussed below.

THYROIDITIS

A clinically useful classification of thyroiditis is based on the onset and duration of disease (Table 405-9).

ACUTE THYROIDITIS

Acute thyroiditis is rare and due to suppurative infection of the thyroid. In children and young adults, the most common cause is the presence of a piriform sinus, a remnant of the fourth branchial pouch that connects the oropharynx with the thyroid. Such sinuses are predominantly left-sided. A long-standing goiter and degeneration in a thyroid malignancy are risk factors in the elderly. The patient presents with thyroid pain, often referred to the throat or ears, and a small, tender goiter that may be asymmetric. Fever, dysphagia, and erythema over the thyroid are common, as are systemic symptoms of a febrile illness and lymphadenopathy.

TABLE 405-9 CAUSES OF THYROIDITIS

Acute	
Bacterial infection: especially <i>Staphylococcus</i> , <i>Streptococcus</i> , and <i>Enterobacter</i>	
Fungal infection: <i>Aspergillus</i> , <i>Candida</i> , <i>Coccidioides</i> , <i>Histoplasma</i> , and <i>Pneumocystis</i>	
Radiation thyroiditis after ¹³¹ I treatment	
Amiodarone (may also be subacute or chronic)	
Subacute	
Viral (or granulomatous) thyroiditis	
Silent thyroiditis (including postpartum thyroiditis)	
Mycobacterial infection	
Drug induced (interferon, amiodarone)	
Chronic	
Autoimmunity: focal thyroiditis, Hashimoto's thyroiditis, atrophic thyroiditis	
Riedel's thyroiditis	
Parasitic thyroiditis: echinococcosis, strongyloidiasis, cysticercosis	
Traumatic: after palpation	

The differential diagnosis of *thyroid pain* includes subacute or, rarely, chronic thyroiditis; hemorrhage into a cyst; malignancy including lymphoma; and, rarely, amiodarone-induced thyroiditis or amyloidosis. However, the abrupt presentation and clinical features of acute thyroiditis rarely cause confusion. The erythrocyte sedimentation rate (ESR) and white cell count are usually increased, but thyroid function is normal. FNA biopsy shows infiltration by polymorphonuclear leukocytes; culture of the sample can identify the organism. Caution is needed in immunocompromised patients as fungal, mycobacterial, or *Pneumocystis* thyroiditis can occur in this setting. Antibiotic treatment is guided initially by Gram stain and, subsequently, by cultures of the FNA biopsy. Surgery may be needed to drain an abscess, which can be localized by CT scan or ultrasound. Tracheal obstruction, septicemia, retropharyngeal abscess, mediastinitis, and jugular venous thrombosis may complicate acute thyroiditis but are uncommon with prompt use of antibiotics.

SUBACUTE THYROIDITIS

This is also termed *de Quervain's thyroiditis*, *granulomatous thyroiditis*, or *viral thyroiditis*. Many viruses have been implicated, including mumps, coxsackie, influenza, adenoviruses, and echoviruses, but attempts to identify the virus in an individual patient are often unsuccessful and do not influence management. The diagnosis of subacute thyroiditis is often overlooked because the symptoms can mimic pharyngitis. The peak incidence occurs at 30–50 years, and women are affected three times more frequently than men.

Pathophysiology The thyroid shows a characteristic patchy inflammatory infiltrate with disruption of the thyroid follicles and multinucleated giant cells within some follicles. The follicular changes progress to granulomas accompanied by fibrosis. Finally, the thyroid returns to normal, usually several months after onset. During the initial phase of follicular destruction, there is release of Tg and thyroid hormones, leading to increased circulating T₄ and T₃ and suppression of TSH (Fig. 405-10). During this destructive phase, radioactive iodine uptake is low or undetectable. After several weeks, the thyroid is depleted of stored thyroid hormone and a phase of hypothyroidism typically occurs, with low unbound T₄ (and sometimes T₃) and moderately increased TSH levels. Radioactive iodine uptake returns to normal or is even increased as a result of the rise in TSH. Finally, thyroid hormone and TSH levels return to normal as the disease subsides.

Clinical Manifestations The patient usually presents with a painful and enlarged thyroid, sometimes accompanied by fever. There may be

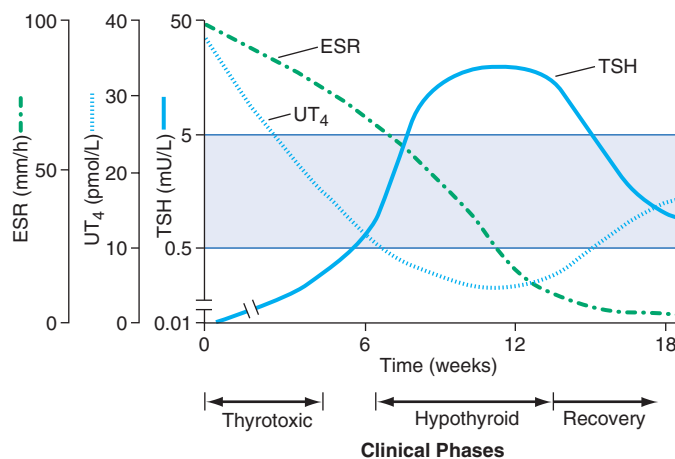


FIGURE 405-10 Clinical course of subacute thyroiditis. The release of thyroid hormones is initially associated with a thyrotoxic phase and suppressed thyroid-stimulating hormone (TSH). A hypothyroid phase then ensues, with low T₄ and TSH levels that are initially low but gradually increase. During the recovery phase, increased TSH levels combined with resolution of thyroid follicular injury lead to normalization of thyroid function, often several months after the beginning of the illness. ESR, erythrocyte sedimentation rate; UT₄, free or unbound T₄.

features of thyrotoxicosis or hypothyroidism, depending on the phase of the illness. Malaise and symptoms of an upper respiratory tract infection may precede the thyroid-related features by several weeks. In other patients, the onset is acute, severe, and without obvious antecedent. The patient typically complains of a sore throat, and examination reveals a small goiter that is exquisitely tender. Pain is often referred to the jaw or ear. Complete resolution is the usual outcome, but late-onset permanent hypothyroidism occurs in 15% of cases, particularly in those with coincidental thyroid autoimmunity. A prolonged course over many months, with one or more relapses, occurs in a small percentage of patients.

Laboratory Evaluation As depicted in Fig. 405-10, thyroid function tests characteristically evolve through three distinct phases over about 6 months: (1) thyrotoxic phase, (2) hypothyroid phase, and (3) recovery phase. In the thyrotoxic phase, T_4 and T_3 levels are increased, reflecting their discharge from the damaged thyroid cells, and TSH is suppressed. The T_4/T_3 ratio is greater than in Graves' disease or thyroid autonomy, in which T_3 is often disproportionately increased. The diagnosis is confirmed by a high ESR and low uptake of radioiodine (<5%) or ^{99m}Tc pertechnetate (as compared to salivary gland pertechnetate concentration). The white blood cell count may be increased, and thyroid antibodies are negative. If the diagnosis is in doubt, FNA biopsy may be useful, particularly to distinguish unilateral involvement from bleeding into a cyst or neoplasm.

TREATMENT SUBACUTE THYROIDITIS

Relatively large doses of aspirin (e.g., 600 mg every 4–6 h) or NSAIDs are sufficient to control symptoms in many cases. If this treatment is inadequate, or if the patient has marked local or systemic symptoms, glucocorticoids should be given. The usual starting dose is 40–60 mg of prednisone, depending on severity. The dose is gradually tapered over 6–8 weeks, in response to improvement in symptoms and the ESR. If a relapse occurs during glucocorticoid withdrawal, treatment should be started again and withdrawn more gradually. In these patients, it is useful to wait until the radioactive iodine uptake normalizes before stopping treatment. Thyroid function should be monitored every 2–4 weeks using TSH and unbound T_4 levels. Symptoms of thyrotoxicosis improve spontaneously but may be ameliorated by β -adrenergic blockers; antithyroid drugs play no role in treatment of the thyrotoxic phase. Levothyroxine replacement may be needed if the hypothyroid phase is prolonged, but doses should be low enough (50–100 μg daily) to allow TSH-mediated recovery.

SILENT THYROIDITIS

Painless thyroiditis, or “*silent*” *thyroiditis*, occurs in patients with underlying autoimmune thyroid disease and has a clinical course similar to that of subacute thyroiditis. The condition occurs in up to 5% of women 3–6 months after pregnancy and is then termed *postpartum thyroiditis*. Typically, patients have a brief phase of thyrotoxicosis lasting 2–4 weeks, followed by hypothyroidism for 4–12 weeks, and then resolution; often, however, only one phase is apparent. The condition is associated with the presence of TPO antibodies antepartum, and it is three times more common in women with type 1 diabetes mellitus. As in subacute thyroiditis, the uptake of ^{99m}Tc pertechnetate or radioactive iodine is initially suppressed. In addition to the painless goiter, silent thyroiditis can be distinguished from subacute thyroiditis by a normal ESR and the presence of TPO antibodies. Glucocorticoid treatment is not indicated for silent thyroiditis. Severe thyrotoxic symptoms can be managed with a brief course of propranolol, 20–40 mg three or four times daily. Thyroxine replacement may be needed for the hypothyroid phase but should be withdrawn after 6–9 months, as recovery is the rule. Annual follow-up thereafter is recommended, because a proportion of these individuals develop permanent hypothyroidism. The condition may recur in subsequent pregnancies.

DRUG-INDUCED THYROIDITIS

Patients receiving cytokines such as IFN- α or IL-2 may develop painless thyroiditis. IFN- α , which is used to treat chronic hepatitis B or C and hematologic and skin malignancies, causes thyroid dysfunction in up to 5% of treated patients. It has been associated with painless thyroiditis, hypothyroidism, and Graves' disease, and is most common in women with TPO antibodies prior to treatment. For discussion of amiodarone, see “Amiodarone Effects on Thyroid Function,” below.

CHRONIC THYROIDITIS

Focal thyroiditis is present in 20–40% of euthyroid autopsy cases and is associated with serologic evidence of autoimmunity, particularly the presence of TPO antibodies. The most common clinically apparent cause of chronic thyroiditis is *Hashimoto's thyroiditis*, an autoimmune disorder that often presents as a firm or hard goiter of variable size (see above). *Riedel's thyroiditis* is a rare disorder that typically occurs in middle-aged women. It presents with an insidious, painless goiter with local symptoms due to compression of the esophagus, trachea, neck veins, or recurrent laryngeal nerves. Dense fibrosis disrupts normal gland architecture and can extend outside the thyroid capsule. Despite these extensive histologic changes, thyroid dysfunction is uncommon. The goiter is hard, nontender, often asymmetric, and fixed, leading to suspicion of a malignancy. Diagnosis requires open biopsy as FNA biopsy is usually inadequate. Treatment is directed to surgical relief of compressive symptoms. Tamoxifen may also be beneficial. There is an association between Riedel's thyroiditis and IgG4-related systemic disease causing idiopathic fibrosis at other sites (retroperitoneum, mediastinum, biliary tree, lung, and orbit).

SICK EUTHYROID SYNDROME (NONTHYROIDAL ILLNESS)

Any acute, severe illness can cause abnormalities of circulating TSH or thyroid hormone levels in the absence of underlying thyroid disease, making these measurements potentially misleading. The major cause of these hormonal changes is the release of cytokines such as IL-6. Unless a thyroid disorder is strongly suspected, the routine testing of thyroid function should be avoided in acutely ill patients.

The most common hormone pattern in sick euthyroid syndrome (SES) is a decrease in total and unbound T_3 levels (low T_3 syndrome) with normal levels of T_4 and TSH. The magnitude of the fall in T_3 correlates with the severity of the illness. T_4 conversion to T_3 via peripheral 5' (outer ring) deiodination is impaired, leading to increased reverse T_3 (rT_3). Since rT_3 is metabolized by 5' deiodination, its clearance is also reduced. Thus, decreased clearance rather than increased production is the major basis for increased rT_3 . Also, T_4 is alternately metabolized to the hormonally inactive T_3 sulfate. It is generally assumed that this low T_3 state is adaptive, because it can be induced in normal individuals by fasting. Teleologically, the fall in T_3 may limit catabolism in starved or ill patients.

Very sick patients may exhibit a dramatic fall in total T_4 and T_3 levels (low T_4 syndrome). With decreased tissue perfusion, muscle and liver expression of the type 3 deiodinase leads to accelerated T_4 and T_3 metabolism. This state has a poor prognosis. Another key factor in the fall in T_4 levels is altered binding to TBG. The commonly used free T_4 assays are subject to artifact when serum binding proteins are low and underestimate the true free T_4 level. Fluctuation in TSH levels also creates challenges in the interpretation of thyroid function in sick patients. TSH levels may range from <0.1 mIU/L in very ill patients, especially with dopamine or glucocorticoid therapy, to >20 mIU/L during the recovery phase of SES. The exact mechanisms underlying the subnormal TSH seen in 10% of sick patients and the increased TSH seen in 5% remain unclear but may be mediated by cytokines including IL-12 and IL-18.

Any severe illness can induce changes in thyroid hormone levels, but certain disorders exhibit a distinctive pattern of abnormalities. Acute liver disease is associated with an initial rise in total (but not unbound) T_3 and T_4 levels due to TBG release; these levels become subnormal with progression to liver failure. A transient increase in total and unbound T_4 levels, usually with a normal T_3 level, is seen in 5–30% of acutely ill psychiatric patients. TSH values may be transiently

2300 low, normal, or high in these patients. In the early stage of HIV infection, T_3 and T_4 levels rise, even if there is weight loss. T_3 levels fall with progression to AIDS, but TSH usually remains normal. Renal disease is often accompanied by low T_3 concentrations, but with normal rather than increased rT_3 levels, due to an unknown factor that increases uptake of rT_3 into the liver.

The diagnosis of SES is challenging. Historic information may be limited, and patients often have multiple metabolic derangements. Useful features to consider include previous history of thyroid disease and thyroid function tests, evaluation of the severity and time course of the patient's acute illness, documentation of medications that may affect thyroid function or thyroid hormone levels, and measurements of rT_3 together with unbound thyroid hormones and TSH. The diagnosis of SES is frequently presumptive, given the clinical context and pattern of laboratory values; only resolution of the test results with clinical recovery can clearly establish this disorder. Treatment of SES with thyroid hormone (T_4 and/or T_3) is controversial, but most authorities recommend monitoring the patient's thyroid function tests during recovery, without administering thyroid hormone, unless there is historic or clinical evidence suggestive of hypothyroidism. Sufficiently large randomized controlled trials using thyroid hormone are unlikely to resolve this therapeutic controversy in the near future, because clinical presentations and outcomes are highly variable.

AMIODARONE EFFECTS ON THYROID FUNCTION

Amiodarone is a commonly used type III antiarrhythmic agent (Chap. 277). It is structurally related to thyroid hormone and contains 39% iodine by weight. Thus, typical doses of amiodarone (200 mg/d) are associated with very high iodine intake, leading to greater than forty-fold increases in plasma and urinary iodine levels. Moreover, because amiodarone is stored in adipose tissue, high iodine levels persist for >6 months after discontinuation of the drug. Amiodarone inhibits deiodinase activity, and its metabolites function as weak antagonists of thyroid hormone action. Amiodarone has the following effects on thyroid function: (1) acute, transient suppression of thyroid function; (2) hypothyroidism in patients susceptible to the inhibitory effects of a high iodine load; and (3) thyrotoxicosis that may be caused by either a Jod-Basedow effect from the iodine load, in the setting of MNG or incipient Graves' disease, or a thyroiditis-like condition.

The initiation of amiodarone treatment is associated with a transient decrease of T_4 levels, reflecting the inhibitory effect of iodine on T_4 release. Soon thereafter, most individuals escape from iodide-dependent suppression of the thyroid (Wolff-Chaikoff effect), and the inhibitory effects on deiodinase activity and thyroid hormone receptor action become predominant. These events lead to the following pattern of thyroid function tests: increased T_4 , decreased T_3 , increased rT_3 , and a transient TSH increase (up to 20 mIU/L). TSH levels normalize or are slightly suppressed within 1–3 months.

The incidence of hypothyroidism from amiodarone varies geographically, apparently correlating with iodine intake. Hypothyroidism occurs in up to 13% of amiodarone-treated patients in iodine-replete countries, such as the United States, but is less common (<6% incidence) in areas of lower iodine intake, such as Italy or Spain. The pathogenesis appears to involve an inability of the thyroid gland to escape from the Wolff-Chaikoff effect in autoimmune thyroiditis. Consequently, amiodarone-associated hypothyroidism is more common in women and individuals with positive TPO antibodies. It is usually unnecessary to discontinue amiodarone for this side effect, because levothyroxine can be used to normalize thyroid function. TSH levels should be monitored, because T_4 levels are often increased for the reasons described above.

The management of amiodarone-induced thyrotoxicosis (AIT) is complicated by the fact that there are different causes of thyrotoxicosis and because the increased thyroid hormone levels exacerbate underlying arrhythmias and coronary artery disease. Amiodarone treatment causes thyrotoxicosis in 10% of patients living in areas of low iodine intake and in 2% of patients in regions of high iodine intake. There are two major forms of AIT, although some patients have features of both. Type 1 AIT is associated with an underlying thyroid abnormality (preclinical Graves'

disease or nodular goiter). Thyroid hormone synthesis becomes excessive as a result of increased iodine exposure (Jod-Basedow phenomenon). Type 2 AIT occurs in individuals with no intrinsic thyroid abnormalities and is the result of drug-induced lysosomal activation leading to destructive thyroiditis with histiocyte accumulation in the thyroid; the incidence rises as cumulative amiodarone dosage increases. Mild forms of type 2 AIT can resolve spontaneously or can occasionally lead to hypothyroidism. Color-flow Doppler thyroid scanning shows increased vascularity in type 1 AIT but decreased vascularity in type 2 AIT. Thyroid scintiscans are difficult to interpret in this setting because the high endogenous iodine levels diminish tracer uptake. However, the presence of normal or rarely increased uptake favors type 1 AIT.

In AIT, the drug should be stopped, if possible, although this is often impractical because of the underlying cardiac disorder. Discontinuation of amiodarone will not have an acute effect because of its storage and prolonged half-life. High doses of antithyroid drugs can be used in type 1 AIT but are often ineffective. In type 2 AIT, oral contrast agents, such as sodium ipodate (500 mg/d) or sodium tyropanoate (500 mg, 1–2 doses/d), rapidly reduce T_4 and T_3 levels, decrease $T_4 \rightarrow T_3$ conversion, and may block tissue uptake of thyroid hormones. Potassium perchlorate, 200 mg every 6 h, has been used to reduce thyroidal iodide content. Perchlorate treatment has been associated with agranulocytosis, although the risk appears relatively low with short-term use. Glucocorticoids, as administered for subacute thyroiditis, have modest benefit in type 2 AIT. Lithium blocks thyroid hormone release and can also provide some benefit. Near-total thyroidectomy rapidly decreases thyroid hormone levels and may be the most effective long-term solution if the patient can undergo the procedure safely.

THYROID FUNCTION IN PREGNANCY

Five factors alter thyroid function in pregnancy: (1) the transient increase in hCG during the first trimester, which stimulates the TSH-R; (2) the estrogen-induced rise in TBG during the first trimester, which is sustained during pregnancy; (3) alterations in the immune system, leading to the onset, exacerbation, or amelioration of an underlying autoimmune thyroid disease (see above); (4) increased thyroid hormone metabolism by the placenta; and (5) increased urinary iodide excretion, which can cause impaired thyroid hormone production in areas of marginal iodine sufficiency. Women with a precarious iodine intake (<50 $\mu\text{g}/\text{d}$) are most at risk of developing a goiter during pregnancy or giving birth to an infant with a goiter and hypothyroidism. The World Health Organization recommends a daily iodine intake of 250 μg during pregnancy and prenatal vitamins should contain 150 μg per tablet.

The rise in circulating hCG levels during the first trimester is accompanied by a reciprocal fall in TSH that persists into the middle of pregnancy. This reflects the weak binding of hCG, which is present at very high levels, to the TSH-R. Rare individuals have been described with variant TSH-R sequences that enhance hCG binding and TSH-R activation. hCG-induced changes in thyroid function can result in transient gestational hyperthyroidism that may be associated with *hyperemesis gravidarum*, a condition characterized by severe nausea and vomiting and risk of volume depletion. However, since the hyperthyroidism is not causal, antithyroid drugs are not indicated unless concomitant Graves' disease is suspected. Parenteral fluid replacement usually suffices until the condition resolves.

During pregnancy, subclinical hypothyroidism occurs in 2% of women, but overt hypothyroidism is present in only 1 in 500. Prospective randomized controlled trials have not shown a benefit for universal thyroid disease screening in pregnancy. Targeted TSH testing for hypothyroidism is recommended for women planning a pregnancy if they have a strong family history of autoimmune thyroid disease, other autoimmune disorders (e.g., type 1 diabetes), prior preterm delivery or recurrent miscarriage, or signs or symptoms of thyroid disease. Thyroid hormone requirements are increased by up to 50% during pregnancy in levothyroxine-treated hypothyroid women (see above section on treatment of hypothyroidism).

GOITER AND NODULAR THYROID DISEASE

Goiter refers to an enlarged thyroid gland. Biosynthetic defects, iodine deficiency, autoimmune disease, and nodular diseases can each lead to goiter, although by different mechanisms. Biosynthetic defects and iodine deficiency are associated with reduced efficiency of thyroid hormone synthesis, leading to increased TSH, which stimulates thyroid growth as a compensatory mechanism to overcome the block in hormone synthesis. Graves' disease and Hashimoto's thyroiditis are also associated with goiter. In Graves' disease, the goiter results mainly from the TSH-R-mediated effects of TSI. The goitrous form of Hashimoto's thyroiditis occurs because of acquired defects in hormone synthesis, leading to elevated levels of TSH and its consequent growth effects. Lymphocytic infiltration and immune system-induced growth factors also contribute to thyroid enlargement in Hashimoto's thyroiditis. Nodular disease is characterized by the disordered growth of thyroid cells, often combined with the gradual development of fibrosis. Because the management of goiter depends on the etiology, the detection of thyroid enlargement on physical examination should prompt further evaluation to identify its cause.

Nodular thyroid disease is common, occurring in about 3–7% of adults when assessed by physical examination. Using ultrasound, nodules are present in up to 50% of adults, with the majority being <1 cm in diameter. Thyroid nodules may be solitary or multiple, and they may be functional or nonfunctional.

DIFFUSE NONTOXIC (SIMPLE) GOITER

Etiology and Pathogenesis When diffuse enlargement of the thyroid occurs in the absence of nodules and hyperthyroidism, it is referred to as a *diffuse nontoxic goiter*. This is sometimes called *simple goiter*, because of the absence of nodules, or *colloid goiter*, because of the presence of uniform follicles that are filled with colloid. Worldwide, diffuse goiter is most commonly caused by iodine deficiency and is termed *endemic goiter* when it affects >5% of the population. In nonendemic regions, *sporadic goiter* occurs, and the cause is usually unknown. Thyroid enlargement in teenagers is sometimes referred to as *juvenile goiter*. In general, goiter is more common in women than men, probably because of the greater prevalence of underlying autoimmune disease and the increased iodine demands associated with pregnancy.

In *iodine-deficient areas*, thyroid enlargement reflects a compensatory effort to trap iodide and produce sufficient hormone under conditions in which hormone synthesis is relatively inefficient. Somewhat surprisingly, TSH levels are usually normal or only slightly increased, suggesting increased sensitivity to TSH or activation of other pathways that lead to thyroid growth. Iodide appears to have direct actions on thyroid vasculature and may indirectly affect growth through vasoactive substances such as endothelins and nitric oxide. Endemic goiter is also caused by exposure to environmental *goitrogens* such as cassava root, which contains a thiocyanate; vegetables of the Cruciferae family (known as cruciferous vegetables) (e.g., Brussels sprouts, cabbage, and cauliflower); and milk from regions where goitrogens are present in grass. Although relatively rare, inherited defects in thyroid hormone synthesis lead to a diffuse nontoxic goiter. Abnormalities at each step in hormone synthesis, including iodide transport (NIS), Tg synthesis, organification and coupling (TPO), and the regeneration of iodide (dehalogenase), have been described.

CLINICAL MANIFESTATIONS AND DIAGNOSIS

If thyroid function is preserved, most goiters are asymptomatic. Examination of a diffuse goiter reveals a symmetrically enlarged, nontender, generally soft gland without palpable nodules. Goiter is defined, somewhat arbitrarily, as a lateral lobe with a volume greater than the thumb of the individual being examined. If the thyroid is markedly enlarged, it can cause tracheal or esophageal compression. These features are unusual, however, in the absence of nodular disease and fibrosis. *Substernal goiter* may obstruct the thoracic inlet. *Pemberton's sign* refers to symptoms of faintness with evidence of facial congestion and external jugular venous obstruction when the arms are raised above the head, a maneuver that draws the thyroid into the thoracic inlet. Respiratory flow measurements and CT or MRI

should be used to evaluate substernal goiter in patients with obstructive signs or symptoms.

Thyroid function tests should be performed in all patients with goiter to exclude thyrotoxicosis or hypothyroidism. It is not unusual, particularly in iodine deficiency, to find a low total T_4 , with normal T_3 and TSH, reflecting enhanced $T_4 \rightarrow T_3$ conversion. A low TSH with a normal free T_3 and free T_4 , particularly in older patients, suggests the possibility of thyroid autonomy or undiagnosed Graves' disease, and is termed *subclinical thyrotoxicosis*. The benefit of treatment (typically with radioiodine) in subclinical thyrotoxicosis, versus follow-up and implementing treatment if free T_3 or free T_4 levels become abnormal, is unclear, but treatment is increasingly recommended in the elderly to reduce the risk of atrial fibrillation and bone loss. TPO antibodies may be useful to identify patients at increased risk of autoimmune thyroid disease. Low urinary iodine levels (<50 $\mu\text{g/L}$) support a diagnosis of iodine deficiency. Thyroid scanning is not generally necessary but will reveal increased uptake in iodine deficiency and most cases of dysmorphogenesis. Ultrasound is not generally indicated in the evaluation of diffuse goiter unless a nodule is palpable on physical examination.

TREATMENT DIFFUSE NONTOXIC (SIMPLE) GOITER

Iodine replacement induces variable regression of goiter in iodine deficiency, depending on how long it has been present and the degree of fibrosis that has developed. Surgery is rarely indicated for diffuse goiter. Exceptions include documented evidence of tracheal compression or obstruction of the thoracic inlet, which are more likely to be associated with substernal MNGs (see below). Subtotal or near-total thyroidectomy for these or cosmetic reasons should be performed by an experienced surgeon to minimize complication rates. Surgery should be followed by replacement with levothyroxine, with the aim of keeping the TSH level at the lower end of the reference range to prevent regrowth of the goiter.

NONTOXIC MULTINODULAR GOITER

Etiology and Pathogenesis Depending on the population studied, MNG or nodular enlargement of the thyroid occurs in up to 12% of adults. MNG is more common in women than men and increases in prevalence with age. It is more common in iodine-deficient regions but also occurs in regions of iodine sufficiency, reflecting multiple genetic, autoimmune, and environmental influences on the pathogenesis.

There is typically wide variation in nodule size. Histology reveals a spectrum of morphologies ranging from hypercellular regions to cystic areas filled with colloid. Fibrosis is often extensive, and areas of hemorrhage or lymphocytic infiltration may be seen. Using molecular techniques, most nodules within an MNG are polyclonal in origin, suggesting a hyperplastic response to locally produced growth factors and cytokines. TSH, which is usually not elevated, may play a permissive or contributory role. Monoclonal lesions also occur within an MNG, reflecting mutations in genes that confer a selective growth advantage to the progenitor cell.

Clinical Manifestations Most patients with nontoxic MNG are asymptomatic and euthyroid. MNG typically develops over many years and is detected on routine physical examination, when an individual notices an enlargement in the neck, or as an incidental finding on imaging. If the goiter is large enough, it can ultimately lead to compressive symptoms including difficulty swallowing, respiratory distress (tracheal compression), or plethora (venous congestion), but these symptoms are uncommon. Symptomatic MNGs are usually extraordinarily large and/or develop fibrotic areas that cause compression. Sudden pain in an MNG is usually caused by hemorrhage into a nodule but should raise the possibility of invasive malignancy. Hoarseness, reflecting laryngeal nerve involvement, also suggests malignancy.

Diagnosis On examination, thyroid architecture is distorted, and multiple nodules of varying size can be appreciated. Because many nodules are deeply embedded in thyroid tissue or reside in posterior or

2302 substernal locations, it is not possible to palpate all nodules. Pemberton's sign, characterized by facial suffusion when the patient's arms are elevated above the head, suggests that the goiter has increased pressure in the thoracic inlet. A TSH level should be measured to exclude subclinical hyper- or hypothyroidism, but thyroid function is usually normal. Tracheal deviation is common, but compression must usually exceed 70% of the tracheal diameter before there is significant airway compromise. Pulmonary function testing can be used to assess the functional effects of compression, which characteristically causes inspiratory stridor. CT or MRI can be used to evaluate the anatomy of the goiter and the extent of substernal extension or tracheal narrowing. A barium swallow may reveal the extent of esophageal compression. The risk of malignancy in MNG is similar to that in solitary nodules. Ultrasonography can be used to identify which nodules should be biopsied based on sonographic features (see section above on ultrasound) and size. For nodules with more suspicious imaging characteristics (e.g., hypoechogenicity, microcalcifications, irregular margins), biopsy is recommended when ≥ 1 cm.

TREATMENT NONTOXIC MULTINODULAR GOITER

Most nontoxic MNGs can be managed conservatively. T_4 suppression is rarely effective for reducing goiter size and introduces the risk of subclinical or overt thyrotoxicosis, particularly if there is underlying autonomy or if it develops during treatment. If levothyroxine is used, it should be started at low doses (50 μg daily) and advanced gradually while monitoring the TSH level to avoid excessive suppression. Contrast agents and other iodine-containing substances should be avoided because of the risk of inducing the *Jod-Basedow effect*, characterized by enhanced thyroid hormone production by autonomous nodules. Radioiodine is used with increasing frequency in areas where large goiters are more prevalent because it can decrease goiter size and may selectively ablate regions of autonomy. Dosage of ^{131}I depends on the size of the goiter and radioiodine uptake but is usually about 3.7 MBq (0.1 mCi) per gram of tissue, corrected for uptake (typical dose 370–1070 MBq [10 to 29 mCi]). Repeat treatment may be needed and effectiveness may be increased by concurrent administration of low-dose recombinant TSH (0.1 mg IM). It is possible to achieve a 40–50% reduction in goiter size in most patients. Earlier concerns about radiation-induced thyroid swelling and tracheal compression have diminished, as studies have shown this complication to be rare. When acute compression occurs, glucocorticoid treatment or surgery may be needed. Radiation-induced hypothyroidism is less common than after treatment for Graves' disease. However, posttreatment autoimmune thyrotoxicosis may occur in up to 5% of patients treated for nontoxic MNG. Surgery remains highly effective but is not without risk, particularly in older patients with underlying cardiopulmonary disease.

TOXIC MULTINODULAR GOITER

The pathogenesis of toxic MNG appears to be similar to that of nontoxic MNG; the major difference is the presence of functional autonomy in toxic MNG. The molecular basis for autonomy in toxic MNG remains unknown. As in nontoxic goiters, many nodules are polyclonal, whereas others are monoclonal and vary in their clonal origins. Genetic abnormalities known to confer functional autonomy, such as activating TSH-R or G_{sa} mutations (see below), are not usually found in the autonomous regions of toxic MNG goiter.

In addition to features of goiter, the clinical presentation of toxic MNG includes subclinical hyperthyroidism or mild thyrotoxicosis. The patient is usually elderly and may present with atrial fibrillation or palpitations, tachycardia, nervousness, tremor, or weight loss. Recent exposure to iodine, from contrast dyes or other sources, may precipitate or exacerbate thyrotoxicosis. The TSH level is low. The uncombined T_4 level may be normal or minimally increased; T_3 is often elevated to a greater degree than T_4 . Thyroid scan shows heterogeneous uptake with multiple regions of increased and decreased

uptake; 24-h uptake of radioiodine may not be increased but is usually in the upper normal range.

Prior to definitive treatment of the hyperthyroidism, ultrasound imaging should be performed to assess the presence of discrete nodules corresponding to areas of decreased uptake ("cold" nodules). If present, FNA may be indicated based on sonographic features and size cutoffs. The cytology results, if indeterminate or suspicious, may direct the therapy to surgery.

TREATMENT TOXIC MULTINODULAR GOITER

Antithyroid drugs normalize thyroid function and are particularly useful in the elderly or ill patients with limited lifespan. In contrast to Graves' disease, spontaneous remission does not occur and so treatment is long-term. Radioiodine is generally the treatment of choice; it treats areas of autonomy as well as decreasing the mass of the goiter. Sometimes, however, a degree of autonomy remains, presumably because multiple autonomous regions emerge as soon as others are treated, and further radioiodine treatment may be necessary. Surgery provides definitive treatment of underlying thyrotoxicosis as well as goiter. Patients should be rendered euthyroid using an antithyroid drug before operation.

HYPERFUNCTIONING SOLITARY NODULE

A solitary, autonomously functioning thyroid nodule is referred to as *toxic adenoma*. The pathogenesis of this disorder has been unraveled by demonstrating the functional effects of mutations that stimulate the TSH-R signaling pathway. Most patients with solitary hyperfunctioning nodules have acquired somatic, activating mutations in the TSH-R (Fig. 405-11). These mutations, located primarily in the receptor transmembrane domain, induce constitutive receptor coupling to G_{sa} , increasing cyclic AMP levels and leading to enhanced thyroid follicular cell proliferation and function. Less commonly, somatic mutations are identified in G_{sa} . These mutations, which are similar to those seen in

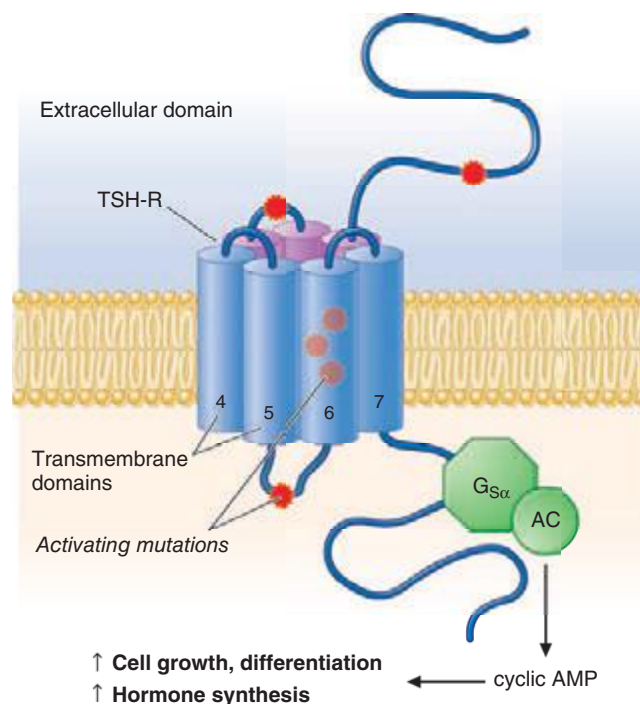


FIGURE 405-11 Activating mutations of the thyroid-stimulating hormone receptor (TSH-R). Mutations (*) that activate TSH-R reside mainly in transmembrane 5 and intracellular loop 3, although mutations have occurred in a variety of different locations. The effect of these mutations is to induce conformational changes that mimic TSH binding, thereby leading to coupling to stimulatory G protein (G_{sa}) and activation of adenylate cyclase (AC), an enzyme that generates cyclic AMP.

McCune-Albright syndrome (Chap. 412) or in a subset of somatotrope adenomas (Chap. 403), impair guanosine triphosphate (GTP) hydrolysis, causing constitutive activation of the cyclic AMP signaling pathway. In most series, activating mutations in either the TSH-R or the G_{sa} subunit genes are identified in >90% of patients with solitary hyperfunctioning nodules.

Thyrotoxicosis is usually mild. The disorder is suggested by a subnormal TSH level; the presence of the thyroid nodule, which is generally large enough to be palpable; and the absence of clinical features suggestive of Graves' disease or other causes of thyrotoxicosis. A thyroid scan provides a definitive diagnostic test, demonstrating focal uptake in the hyperfunctioning nodule and diminished uptake in the remainder of the gland, as activity of the normal thyroid is suppressed.

TREATMENT HYPERFUNCTIONING SOLITARY NODULE

Radioiodine ablation is usually the treatment of choice. Because normal thyroid function is suppressed, ^{131}I is concentrated in the hyperfunctioning nodule with minimal uptake and damage to normal thyroid tissue. Relatively large radioiodine doses (e.g., 370–1110 MBq [10–29.9 mCi] ^{131}I) have been shown to correct thyrotoxicosis in about 75% of patients within 3 months. Hypothyroidism occurs in <10% of those patients over the next 5 years. Surgical resection is also effective and is usually limited to enucleation of the adenoma or lobectomy, thereby preserving thyroid function and minimizing risk of hypoparathyroidism or damage to the recurrent laryngeal nerves. Medical therapy using antithyroid drugs and beta blockers can normalize thyroid function but is not an optimal long-term treatment. Using ultrasound guidance, repeated ethanol injections and percutaneous radiofrequency thermal ablation have been used successfully in some centers to ablate hyperfunctioning nodules, and these techniques have also been used to reduce the size of nonfunctioning thyroid nodules.

BENIGN NEOPLASMS

The various types of benign thyroid nodules are listed in Table 405-10. These lesions are common (5–10% adults), particularly when assessed by sensitive techniques such as ultrasound. The risk of malignancy is very low for *macrofollicular adenomas* and *normofollicular adenomas*. *Microfollicular*, *trabecular*, and *Hürthle cell variants* raise greater concern, and the histology is more difficult to interpret. Many are mixed cystic/solid lesions on ultrasound and may appear spongiform reflecting the pathology of macrofollicular structure. However, the majority of solid nodules (whether hypo-, iso-, or hyperechoic) are also benign. FNA, usually performed with ultrasound guidance, is the diagnostic procedure of choice to evaluate thyroid nodules (see the “Approach to the Patient” section on thyroid nodules). Pure thyroid cysts, <2% of all thyroid growths, consist of colloid and are benign as well. Cysts frequently recur, even after repeated aspiration, and may require surgical excision if they are large. Ethanol ablation to sclerose the cyst has been used successfully for patients who are symptomatic.

TSH suppression with levothyroxine therapy does not decrease thyroid nodule size in iodine-sufficient populations. However, if there is relative iodine deficiency, both iodine and levothyroxine therapy may decrease nodule volume. If levothyroxine is administered in this situation and the nodule has not decreased in size after 6–12 months of suppressive therapy, treatment should be discontinued because little benefit is likely to accrue from long-term treatment; the risk of iatrogenic subclinical thyrotoxicosis should also be considered.

THYROID CANCER

Thyroid carcinoma is the most common malignancy of the endocrine system. Malignant tumors derived from the follicular epithelium are classified according to histologic features. Differentiated tumors, such as papillary thyroid cancer (PTC) or follicular thyroid cancer (FTC), are often curable, and the prognosis is good for patients identified with early-stage disease. In contrast, anaplastic thyroid cancer (ATC)

TABLE 405-10 CLASSIFICATION OF THYROID NEOPLASMS

Benign	
Follicular epithelial cell adenomas	
Macrofollicular (colloid)	
Normofollicular (simple)	
Microfollicular (fetal)	
Trabecular (embryonal)	
Hürthle cell variant (oncocytic)	
Malignant	
	Approximate Prevalence, %
Follicular epithelial cell	
Well-differentiated carcinomas	
Papillary carcinomas	80–90
Pure papillary	
Follicular variant	
Diffuse sclerosing variant	
Tall cell, columnar cell variants	
Follicular carcinomas	5–10
Minimally invasive	
Widely invasive	
Hürthle cell carcinoma (oncocytic)	
Insular carcinoma	
Undifferentiated (anaplastic) carcinomas	
C cell (calcitonin-producing)	
Medullary thyroid cancer	<10
Sporadic	
Familial	
MEN 2	
Other malignancies	
Lymphomas	1–2
Sarcomas	
Metastases	
Others	

Abbreviation: MEN, multiple endocrine neoplasia.

is aggressive, responds poorly to treatment, and is associated with a bleak prognosis.

The incidence of thyroid cancer is ~12/100,000 per year in the United States and increases with age. Prognosis is worse in older persons (>65 years). Thyroid cancer is twice as common in women as men, but male gender is associated with a worse prognosis. Additional important risk factors include a history of childhood head or neck irradiation, large nodule size (≥ 4 cm), evidence for local tumor fixation or invasion into lymph nodes, and the presence of metastases (Table 405-11). Several unique features of thyroid cancer facilitate its management: (1) thyroid nodules are amenable to biopsy by FNA; (2) iodine radioisotopes can be used to diagnose (^{123}I) and treat (^{131}I) differentiated thyroid cancer, reflecting the unique uptake of this anion by the thyroid gland; and (3) serum markers allow the detection of

TABLE 405-11 RISK FACTORS FOR THYROID CARCINOMA IN PATIENTS WITH THYROID NODULE

History of head and neck irradiation, including total-body irradiation for bone marrow transplant and brain radiation for childhood leukemia	Family history of thyroid cancer, MEN 2, or other genetic syndromes associated with thyroid malignancy (e.g., Cowden's syndrome, familial polyposis, Carney complex)
Exposure to ionizing radiation from fallout in childhood or adolescence	Vocal cord paralysis, hoarse voice
Age <20 or >65 years	Nodule fixed to adjacent structures
Increased nodule size (>4 cm)	Extrathyroidal extension
New or enlarging neck mass	Lateral cervical lymphadenopathy
Male gender	

Abbreviation: MEN, multiple endocrine neoplasia.

2304 residual or recurrent disease, including the use of Tg levels for PTC and FTC, and calcitonin for medullary thyroid cancer (MTC).

CLASSIFICATION

Thyroid neoplasms can arise in each of the cell types that populate the gland, including thyroid follicular cells, calcitonin-producing C cells, lymphocytes, and stromal and vascular elements, as well as metastases from other sites (Table 405-10). The American Joint Committee on Cancer (AJCC) has designated a staging system using the tumor, node, metastasis (TNM) classification (Table 405-12). Several other classification and staging systems are also widely used, some of which place greater emphasis on histologic features or risk factors such as age or gender.

PATHOGENESIS AND GENETIC BASIS

Radiation Early studies of the pathogenesis of thyroid cancer focused on the role of external radiation, which predisposes to chromosomal breaks, leading to genetic rearrangements and loss of tumor-suppressor genes. External radiation of the mediastinum, face, head, and neck region was administered in the past to treat an array of conditions, including acne and enlargement of the thymus, tonsils, and adenoids. Radiation exposure increases the risk of benign and malignant thyroid nodules, is associated with multicentric cancers, and shifts the incidence of thyroid cancer to an earlier age group. Radiation from nuclear fallout also increases the risk of thyroid cancer. Children seem more predisposed to the effects of radiation than adults. Of note, radiation derived from ¹³¹I therapy appears to contribute minimal increased risk of thyroid cancer.

TSH and Growth Factors Many differentiated thyroid cancers express TSH receptors and, therefore, remain responsive to TSH. Higher serum TSH levels, even within normal range, are associated with increased thyroid cancer risk in patients with thyroid nodules. These

observations provide the rationale for T₄ suppression of TSH in patients with thyroid cancer. Residual expression of TSH receptors also allows TSH-stimulated uptake of ¹³¹I therapy (see below).

Oncogenes and Tumor-Suppressor Genes Thyroid cancers are monoclonal in origin, consistent with the idea that they originate as a consequence of mutations that confer a growth advantage to a single cell. In addition to increased rates of proliferation, some thyroid cancers exhibit impaired apoptosis and features that enhance invasion, angiogenesis, and metastasis. Thyroid neoplasms have been analyzed for a variety of genetic alterations, but without clear evidence of an ordered acquisition of somatic mutations as they progress from the benign to the malignant state. On the other hand, certain mutations are relatively specific for thyroid neoplasia, some of which correlate with histologic classification (Table 405-13).

As described above, activating mutations of the TSH-R and the G_{sα} subunit are associated with autonomously functioning nodules. Although these mutations induce thyroid cell growth, this type of nodule is almost always benign.

Activation of the RET-RAS-BRAF signaling pathway is seen in up to 70% of PTCs, although the types of mutations are heterogeneous. A variety of rearrangements involving the *RET* gene on chromosome 10 bring this receptor tyrosine kinase under the control of other promoters, leading to receptor overexpression. *RET* rearrangements occur in 20–40% of PTCs in different series and were observed with increased frequency in tumors developing after the Chernobyl radiation accident. Rearrangements in PTC have also been observed for another tyrosine kinase gene, *TRK1*, which is located on chromosome 1. To date, the identification of PTC with *RET* or *TRK1* rearrangements has not proven useful for predicting prognosis or treatment responses. *BRAF V600E* mutations appear to be the most common genetic alteration in PTC. These mutations activate the kinase, which stimulates the mitogen-activated protein MAP kinase (MAPK) cascade. *RAS* mutations, which also stimulate the MAPK cascade, are found in about 20–30% of thyroid neoplasms (*NRAS* > *HRAS* > *KRAS*), including both PTC and FTC. Of note, simultaneous *RET*, *BRAF*, and *RAS* mutations rarely occur in the same tumor, suggesting that activation of the MAPK cascade is critical for tumor development, independent of the step that initiates the cascade.

RAS mutations also occur in FTCs. In addition, a rearrangement of the thyroid developmental transcription factor *PAX8* with the nuclear receptor *PPARγ* is identified in a significant fraction of FTCs. Overall, about 70% of follicular cancers have mutations or genetic rearrangements. Loss of heterozygosity of 3p or 11q, consistent with deletions of tumor-suppressor genes, is also common in FTCs.

Most of the mutations seen in differentiated thyroid cancers have also been detected in ATCs. *BRAF* mutations are seen in up to 50% of ATCs. Mutations in *CTNNB1*, which encodes β-catenin, occur in about two-thirds of ATCs, but not in PTC or FTC. Mutations of the tumor-suppressor *P53* also play an important role in the development of ATC. Because *P53* plays a role in cell cycle surveillance, DNA repair, and apoptosis, its loss may contribute to the rapid acquisition of genetic instability as well as poor treatment responses (Chap. 102e) (Table 405-13).

The role of molecular diagnostics in the clinical management of thyroid cancer is under investigation. In principle, analyses of specific mutations might aid in classification, prognosis, or choice of treatment. Although *BRAF V600E* mutations are associated with loss of iodine uptake by tumor cells, there is no clear evidence to date that this information alters clinical decision making. Higher recurrence rates have been variably reported in patients with *BRAF*-positive PTC, but the impact on survival rates is unclear. Sequencing of thyroid cancers as part of the Cancer Genome Atlas (TCGA) is likely to lead to new classification schemes based on molecular abnormalities in tumors.

MTC, when associated with multiple endocrine neoplasia (MEN) type 2, harbors an inherited mutation of the *RET* gene. Unlike the rearrangements of *RET* seen in PTC, the mutations in MEN 2 are

TABLE 405-12 THYROID CANCER CLASSIFICATION^a

Papillary or Follicular Thyroid Cancers		
	<45 years	>45 years
Stage I	Any T, any N, M0	T1, N0, M0
Stage II	Any T, any N, M1	T2, N0, M0
Stage III	—	T3, N0, M0
Stage IVA	—	T1–T3, N1a, M0
Stage IVB	—	T4a, any N, M0
Stage IVC	—	T1–T3, N1b, M0
Stage IVB	—	T4b, any N, M0
Stage IVC	—	Any T, any N, M1
Anaplastic Thyroid Cancer		
Stage IV	All cases are stage IV	
Medullary Thyroid Cancer		
Stage I	T1, N0, M0	
Stage II	T2 or T3, N0, M0	
Stage III	T1–T3, N1a, M0	
Stage IVA	T4a, any N, M0	
Stage IVB	T1–T3, N1b, M0	
Stage IVB	T4b, any N, M0	
Stage IVC	Any T, any N, M1	

^aCriteria include: T, the size and extent of the primary tumor (T1a ≤1 cm; T1b >1 cm but ≤2 cm; T2 >2 cm but ≤4 cm; T3 >4 cm or any tumor with extension into perithyroidal soft tissue or sternothyroid muscle; T4a invasion into subcutaneous soft tissues, larynx, trachea, esophagus, or recurrent laryngeal nerve; T4b invasion into prevertebral fascia or encasement of carotid artery or mediastinal vessels); N, the absence (N0) or presence (N1a level IV central compartment; N1b levels II–V lateral compartment, upper mediastinal or retro-/parapharyngeal) of regional node involvement; M, the absence (M0) or presence (M1) of distant metastases.

Source: American Joint Committee on Cancer staging system for thyroid cancers using the TNM classification, 7th edition.

TABLE 405-13 GENETIC ALTERATIONS IN THYROID NEOPLASIA

Gene/Protein	Type of Gene	Chromosomal Location	Genetic Abnormality	Tumor
TSH receptor	GPCR receptor	14q31	Point mutations	Toxic adenoma, differentiated carcinomas
G _{sα}	G protein	20q13.2	Point mutations	Toxic adenoma, differentiated carcinomas
RET/PTC	Receptor tyrosine kinase	10q11.2	Rearrangements PTC1: inv(10)(q11.2q21) PTC2: t(10;17)(q11.2;q23) PTC3: ELE1/TK	PTC (more common in radiation-induced tumors)
RET	Receptor tyrosine kinase	10q11.2	Point mutations	MEN 2, medullary thyroid cancer
BRAF	MEK kinase	7q24	Point mutations, rearrangements	PTC, ATC
TRK	Receptor tyrosine kinase	1q23-24	Rearrangements	Multinodular goiter, papillary thyroid cancer
RAS	Signal transducing p21	NRAS 1p13.2 (most common); HRAS 11p15.5; KRAS 12p12.1	Point mutations	Follicular thyroid cancer, PTC follicular variant, adenomas
p53	Tumor suppressor, cell cycle control, apoptosis	17p13	Point mutations Deletion, insertion	Anaplastic cancer
APC	Tumor suppressor, adenomatous polyposis coli gene	5q21-q22	Point mutations	Anaplastic cancer, also associated with familial polyposis coli
p16 (MTS1, CDKN2A)	Tumor suppressor, cell cycle control	9p21	Deletions	Differentiated carcinomas
p21/WAF	Tumor suppressor, cell cycle control	6p21.2	Overexpression	Anaplastic cancer
MET	Receptor tyrosine kinase	7q31	Overexpression	Follicular thyroid cancer
c-MYC	Receptor tyrosine kinase	8q24.12-13	Overexpression	Differentiated carcinoma
PTEN	Phosphatase	10q23	Point mutations	PTC in Cowden's syndrome (multiple hamartomas, breast tumors, gastrointestinal polyps, thyroid tumors)
CTNNB1	β-Catenin	3p22	Point mutations	Anaplastic cancer
Loss of heterozygosity (LOH)	? Tumor suppressors	3p; 11q13, other loci	Deletions	Differentiated thyroid carcinomas, anaplastic cancer
PAX8-PPARγ1	Transcription factor-nuclear receptor fusion	t(2;3)(q13;p25)	Translocation	Follicular adenoma or carcinoma, rare PTC follicular variant

Abbreviations: APC, adenomatous polyposis coli; ATC, anaplastic thyroid cancer; BRAF, v-raf homologue, B1; CDKN2A, cyclin-dependent kinase inhibitor 2A; c-MYC, cellular homologue of myelocytomatosis virus protooncogene; ELE1/TK, RET-activating gene ele1/tyrosine kinase; GPCR, G protein-coupled receptor; G_{sα}, G-protein stimulating α-subunit; MEK, mitogen extracellular signal-regulated kinase; MEN 2, multiple endocrine neoplasia-2; MET, met protooncogene (hepatocyte growth factor receptor); MTS, multiple tumor suppressor; p53, p53 tumor suppressor gene; PTC, papillary thyroid cancer; PTEN, phosphatase and tensin homologue; RAS, rat sarcoma protooncogene; RET, rearranged during transfection protooncogene; p21, p21 tumor suppressor; PAX8, paired domain transcription factor; PPARγ1, peroxisome-proliferator activated receptor γ1; TRK, tyrosine kinase receptor; TSH, thyroid-stimulating hormone; WAF, wild-type p53 activated fragment.

Source: Adapted with permission from P Kopp, JL Jameson, in JL Jameson (ed): *Principles of Molecular Medicine*. Totowa, NJ, Humana Press, 1998.

point mutations that induce constitutive activity of the tyrosine kinase (**Chap. 408**). MTC is preceded by hyperplasia of the C cells, raising the likelihood that as-yet-unidentified “second hits” lead to cellular transformation. A subset of sporadic MTC contains somatic mutations that activate *RET*.

WELL-DIFFERENTIATED THYROID CANCER

Papillary PTC is the most common type of thyroid cancer, accounting for 70–90% of well-differentiated thyroid malignancies. Microscopic PTC is present in up to 25% of thyroid glands at autopsy, but most of these lesions are very small (several millimeters) and are not clinically significant. Characteristic cytologic features of PTC help make the diagnosis by FNA or after surgical resection; these include psammoma bodies, cleaved nuclei with an “orphan-Annie” appearance caused by large nucleoli, and the formation of papillary structures.

PTC tends to be multifocal and to invade locally within the thyroid gland as well as through the thyroid capsule and into adjacent structures in the neck. It has a propensity to spread via the lymphatic system but can metastasize hematogenously as well, particularly to bone and lung. Because of the relatively slow growth of the tumor, a significant burden of pulmonary metastases may accumulate, sometimes with remarkably few symptoms. The prognostic implication of lymph node

spread is debated. Lymph node involvement by thyroid cancer can be well tolerated but appears to increase the risk of recurrence and mortality, particularly in older patients. The staging of PTC by the TNM system is outlined in Table 405-12. Most papillary cancers are identified in the early stages (>80% stages I or II) and have an excellent prognosis, with survival curves similar to expected survival (**Fig. 405-12**). Mortality is markedly increased in stage IV disease, especially in the presence of distant metastases (stage IVC), but this group comprises only about 1% of patients. The treatment of PTC is described below.

Follicular The incidence of FTC varies widely in different parts of the world; it is more common in iodine-deficient regions. Currently, FTC accounts for only about 5% of all thyroid cancers diagnosed in the United States. FTC is difficult to diagnose by FNA because the distinction between benign and malignant follicular neoplasms rests largely on evidence of invasion into vessels, nerves, or adjacent structures. FTC tends to spread by hematogenous routes leading to bone, lung, and central nervous system metastases. Mortality rates associated with FTC are less favorable than for PTC, in part because a larger proportion of patients present with stage IV disease. Poor prognostic features include distant metastases, age >50 years, primary tumor size >4 cm, Hürthle cell histology, and the presence of marked vascular invasion.

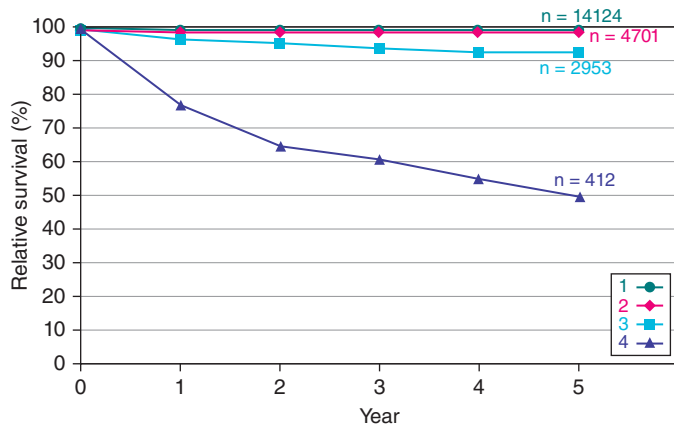


FIGURE 405-12 Survival rates of patients with different stages of papillary cancer. (Adapted with permission from Edge SB, Byrd DR: *Thyroid*, in Compton CC, Fritz AB, Greene FL, Trotti A [eds]: *AJCC Cancer Staging Manual*, 7th ed. New York, Springer, 2010, pp 87–92.)

TREATMENT WELL-DIFFERENTIATED THYROID CANCER

SURGERY

All well-differentiated thyroid cancers should be surgically excised. In addition to removing the primary lesion, surgery allows accurate histologic diagnosis and staging, and multicentric disease is commonly found in the contralateral thyroid lobe. Preoperative sonography should be performed in all patients to assess the central and lateral cervical lymph node compartments for suspicious adenopathy, which if present, can undergo FNA and then be removed at surgery. Bilateral, near-total thyroidectomy has been shown to reduce recurrence rates in all patients except those with T1a tumors (≤ 1 cm). If cytology is diagnostic for thyroid cancer, bilateral surgery should be done. If malignancy is identified pathologically after lobectomy, completion surgery is recommended unless the tumor is T1a or is a minimally invasive follicular cancer. Bilateral surgery for patients at higher risk allows monitoring of serum Tg levels and administration of radioiodine for remnant ablation and potential treatment of iodine-avid metastases, if indicated. Therefore, near-total thyroidectomy is preferable in almost all patients; complication rates are acceptably low if the surgeon is highly experienced in the procedure.

TSH SUPPRESSION THERAPY

Because most tumors are still TSH-responsive, levothyroxine suppression of TSH is a mainstay of thyroid cancer treatment. Although TSH suppression clearly provides therapeutic benefit, there are no prospective studies that define the optimal level of TSH suppression. The degree of TSH suppression should be individualized based on a patient's risk of recurrence. It should be adjusted over time as surveillance blood tests and imaging confirm absence of disease or, alternatively, indicate possible residual/recurrent cancer. For patients at low risk of recurrence, TSH should be suppressed into the low but detectable range (0.1–0.5 mIU/L). If subsequent surveillance testing indicates no evidence of disease, the TSH target may rise to the lower half of the normal range. For patients at high risk of recurrence or with known metastatic disease, TSH levels should be kept to <0.1 mIU/L if there are no strong contraindications to mild thyrotoxicosis. In this instance, unbound T_4 must also be monitored to avoid excessive treatment.

RADIOIODINE TREATMENT

After near-total thyroidectomy, substantial thyroid tissue often remains, particularly in the thyroid bed and surrounding the parathyroid glands. Postsurgical radioablation of the remnant thyroid eliminates residual normal thyroid, facilitating the use of Tg

determinations and radioiodine scanning for long-term follow-up. In addition, well-differentiated thyroid cancer often incorporates radioiodine, although less efficiently than normal thyroid follicular cells. Radioiodine uptake is determined primarily by expression of the NIS and is stimulated by TSH, requiring expression of the TSH-R. The retention time for radioactivity is influenced by the extent to which the tumor retains differentiated functions such as iodide trapping and organification. Consequently, for patients at risk of recurrence and for those with known distant metastatic disease, ^{131}I ablation may also potentially treat residual tumor cells.

Indications Not all patients benefit from radioiodine therapy. Neither recurrence nor survival rates are improved in stage I patients with T1 tumors (≤ 2 cm) confined to the thyroid. However, in higher risk patients (larger tumors, more aggressive variants of papillary cancer, tumor vascular invasion, presence of large-volume lymph node metastases), radioiodine reduces recurrence and may increase survival.

^{131}I Thyroid Ablation and Treatment As noted above, the decision to use ^{131}I for thyroid ablation should be coordinated with the surgical approach, because radioablation is much more effective when there is minimal remaining normal thyroid tissue. Radioiodine is administered after iodine depletion (patient follows a low-iodine diet for 1–2 weeks) and in the presence of elevated serum TSH levels to stimulate uptake of the isotope into both the remnant and potentially any residual tumor. To achieve high serum TSH levels, there are two approaches. A patient may be withdrawn from thyroid hormone so that endogenous TSH is secreted and, ideally, the serum TSH level is >25 mIU/L at the time of ^{131}I therapy. A typical strategy is to treat the patient for several weeks postoperatively with liothyronine (25 μg qd or bid), followed by thyroid hormone withdrawal for 2 weeks. Alternatively, recombinant human TSH (rhTSH) is administered as two daily consecutive injections (0.9 mg) with administration of ^{131}I 24 h after the second injection. The patient can continue to take levothyroxine and remains euthyroid. Both approaches have equal success in achieving remnant ablation.

A pretreatment scanning dose of ^{131}I (usually 111–185 MBq [3–5 mCi]) or ^{123}I (74 MBq [2 mCi]) can reveal the amount of residual tissue and provides guidance about the dose needed to accomplish ablation. However, because of concerns about radioactive “stunning” that impairs subsequent treatment, there is a trend to avoid pretreatment scanning with ^{131}I and use either ^{123}I or proceed directly to ablation, unless there is suspicion that the amount of residual tissue will alter therapy or that there is distant metastatic disease. In the United States, outpatient doses of up to 6475 MBq (175 mCi) can be given at most centers. The administered dose depends on the indication for therapy with lower doses of 1850–2775 MBq (50–75 mCi) given for remnant ablation but higher doses of 3700–5500 MBq (100–150 mCi) used as adjuvant therapy when residual disease may be present. A WBS following radioiodine treatment is used to confirm the ^{131}I uptake in the remnant and to identify possible metastatic disease.

Follow-Up Whole-Body Thyroid Scanning and Thyroglobulin Determinations Serum thyroglobulin is a sensitive marker of residual/recurrent thyroid cancer after ablation of the residual postsurgical thyroid tissue. However, newer Tg assays have functional sensitivities as low as 0.1 ng/mL, as opposed to older assays with functional sensitivities of 1 ng/mL, reducing the number of patients with truly undetectable serum Tg levels. Because the vast majority of papillary thyroid cancer recurrences are in cervical lymph nodes, a neck ultrasound should be performed about 6 months after thyroid ablation; ultrasound has been shown to be more sensitive than WBS in this scenario.

In low-risk patients who have no clinical evidence of residual disease after ablation and a basal Tg <1 ng/mL on levothyroxine, an rhTSH-stimulated Tg level should be obtained 6–12 months after ablation, without WBS. If stimulated Tg levels are low (<1 ng/mL)

and, ideally, undetectable, the risk of recurrence is <5% at 5 years. Newer data indicate that rhTSH stimulation may not be required for patients with undetectable basal Tg levels in sensitive assays, if there is documented absence of Tg antibodies. These patients can be followed with unstimulated Tg every 6–12 months and neck ultrasound as indicated. Levothyroxine dosing may then be titrated to a higher TSH level of 0.5–1.5 mIU/L.

The use of WBS is reserved for patients with known iodine-avid metastases or those with elevated serum thyroglobulin levels and negative imaging with ultrasound, chest CT, and neck cross-sectional imaging who may require additional ¹³¹I therapy.

In addition, most authorities advocate radioiodine treatment for scan-negative, Tg-positive (Tg >5–10 ng/mL) patients, as many derive therapeutic benefit from a large dose of ¹³¹I. For such patients, rhTSH preparation is not FDA approved for the treatment of metastatic disease, and the traditional approach of thyroid hormone withdrawal should be followed. This involves switching patients from levothyroxine (T₄) to the more rapidly cleared hormone liothyronine (T₃), thereby allowing TSH to increase more quickly. Whenever ¹³¹I is administered, posttherapy WBS is the gold standard to assess iodine-avid metastases.

In addition to radioiodine, external beam radiotherapy is also used to treat specific metastatic lesions, particularly when they cause bone pain or threaten neurologic injury (e.g., vertebral metastases).

New Potential Therapies Kinase inhibitors are being explored as a means to target pathways known to be active in thyroid cancer, including the RAS, BRAF, EGFR, VEGFR, and angiogenesis pathways. A multicenter randomized controlled trial of the multikinase inhibitor sorafenib in 417 patients with progressive metastatic thyroid cancer reported a doubling of progression-free survival to 10.8 months in the treatment group compared with the placebo group. Ongoing trials are exploring whether differentiation protocols with kinase inhibitors or other approaches might enhance radioiodine uptake and efficacy.

ANAPLASTIC AND OTHER FORMS OF THYROID CANCER

Anaplastic Thyroid Cancer As noted above, ATC is a poorly differentiated and aggressive cancer. The prognosis is poor, and most patients die within 6 months of diagnosis. Because of the undifferentiated state of these tumors, the uptake of radioiodine is usually negligible, but it can be used therapeutically if there is residual uptake. Chemotherapy has been attempted with multiple agents, including anthracyclines and paclitaxel, but it is usually ineffective. External beam radiation therapy can be attempted and continued if tumors are responsive.

Thyroid Lymphoma Lymphoma in the thyroid gland often arises in the background of Hashimoto's thyroiditis. A rapidly expanding thyroid mass suggests the possibility of this diagnosis. Diffuse large-cell lymphoma is the most common type in the thyroid. Biopsies reveal sheets of lymphoid cells that can be difficult to distinguish from small-cell lung cancer or ATC. These tumors are often highly sensitive to external radiation. Surgical resection should be avoided as initial therapy because it may spread disease that is otherwise localized to the thyroid. If staging indicates disease outside of the thyroid, treatment should follow guidelines used for other forms of lymphoma (**Chap. 134**).

MEDULLARY THYROID CARCINOMA

MTC can be sporadic or familial and accounts for about 5% of thyroid cancers. There are three familial forms of MTC: MEN 2A, MEN 2B, and familial MTC without other features of MEN (**Chap. 408**). In general, MTC is more aggressive in MEN 2B than in MEN 2A, and familial MTC is more aggressive than sporadic MTC. Elevated serum calcitonin provides a marker of residual or recurrent disease. All patients with MTC should be tested for *RET* mutations, because genetic counseling and testing of family members can be offered to those individuals who test positive for mutations.

The management of MTC is primarily surgical. Unlike tumors derived from thyroid follicular cells, these tumors do not take up radioiodine. External radiation treatment and chemotherapy may provide palliation in patients with advanced disease (**Chap. 408**).

APPROACH TO THE PATIENT:

Thyroid Nodules

Palpable thyroid nodules are found in about 5% of adults, but the prevalence varies considerably worldwide. Given this high prevalence rate, practitioners commonly identify thyroid nodules either on physical examination or as incidental findings on imaging performed for another indication (e.g., carotid ultrasound, cervical spine MRI). The main goal of this evaluation is to identify, in a cost-effective manner, the small subgroup of individuals with malignant lesions.

Nodules are more common in iodine-deficient areas, in women, and with aging. Most palpable nodules are >1 cm in diameter, but the ability to feel a nodule is influenced by its location within the gland (superficial versus deeply embedded), the anatomy of the patient's neck, and the experience of the examiner. More sensitive methods of detection, such as CT, thyroid ultrasound, and pathologic studies, reveal thyroid nodules in up to 50% of glands in individuals over the age of 50. The presence of these thyroid incidentalomas has led to much debate about how to detect nodules and which nodules to investigate further.

An approach to the evaluation of a solitary nodule is outlined in **Fig. 405-13**. Most patients with thyroid nodules have normal thyroid function tests. Nonetheless, thyroid function should be assessed by measuring a TSH level, which may be suppressed by one or more autonomously functioning nodules. If the TSH is suppressed, a radionuclide scan is indicated to determine if the identified nodule is "hot," as lesions with increased uptake are almost never malignant and FNA is unnecessary. Otherwise, the next step in evaluation is performance of a thyroid ultrasound for three reasons: (1) Ultrasound will confirm if the palpable nodule is indeed a nodule. About 15% of "palpable" nodules are not confirmed on imaging, and therefore, no further evaluation is required. (2) Ultrasound will assess if there are additional nonpalpable nodules for which FNA may be recommended based on imaging features and size. (3) Ultrasound will characterize the imaging features of the nodule, which, combined with the nodule's size, facilitate decision making about FNA.

Evidence-based guidelines from both the American Thyroid Association and the American Association of Clinical Endocrinologists provide recommendations for nodule FNA based on sonographic imaging features and size cut offs, with lower size cut offs for nodules with more suspicious ultrasound characteristics. FNA biopsy, ideally performed with ultrasound guidance, has good sensitivity and specificity when performed by physicians familiar with the procedure and when the results are interpreted by experienced cytopathologists. The technique is particularly useful for detecting PTC. However, the distinction between benign and malignant follicular lesions is often not possible using cytology alone.

In several large studies, FNA biopsies yielded the following findings: 65% benign, 5% malignant or suspicious for malignancy, 10% nondiagnostic or yielding insufficient material for diagnosis, and 20% indeterminate. The Bethesda System is now widely used to provide more uniform terminology for reporting thyroid nodule FNA cytology results. This six-tiered classification system with the respective estimated malignancy rates is shown in **Table 405-14**. Specifically, the Bethesda System subcategorized cytology specimens previously labeled as indeterminate into three categories: atypia or follicular lesion of undetermined significance (AUS/FLUS), follicular neoplasm, and suspicious for malignancy.

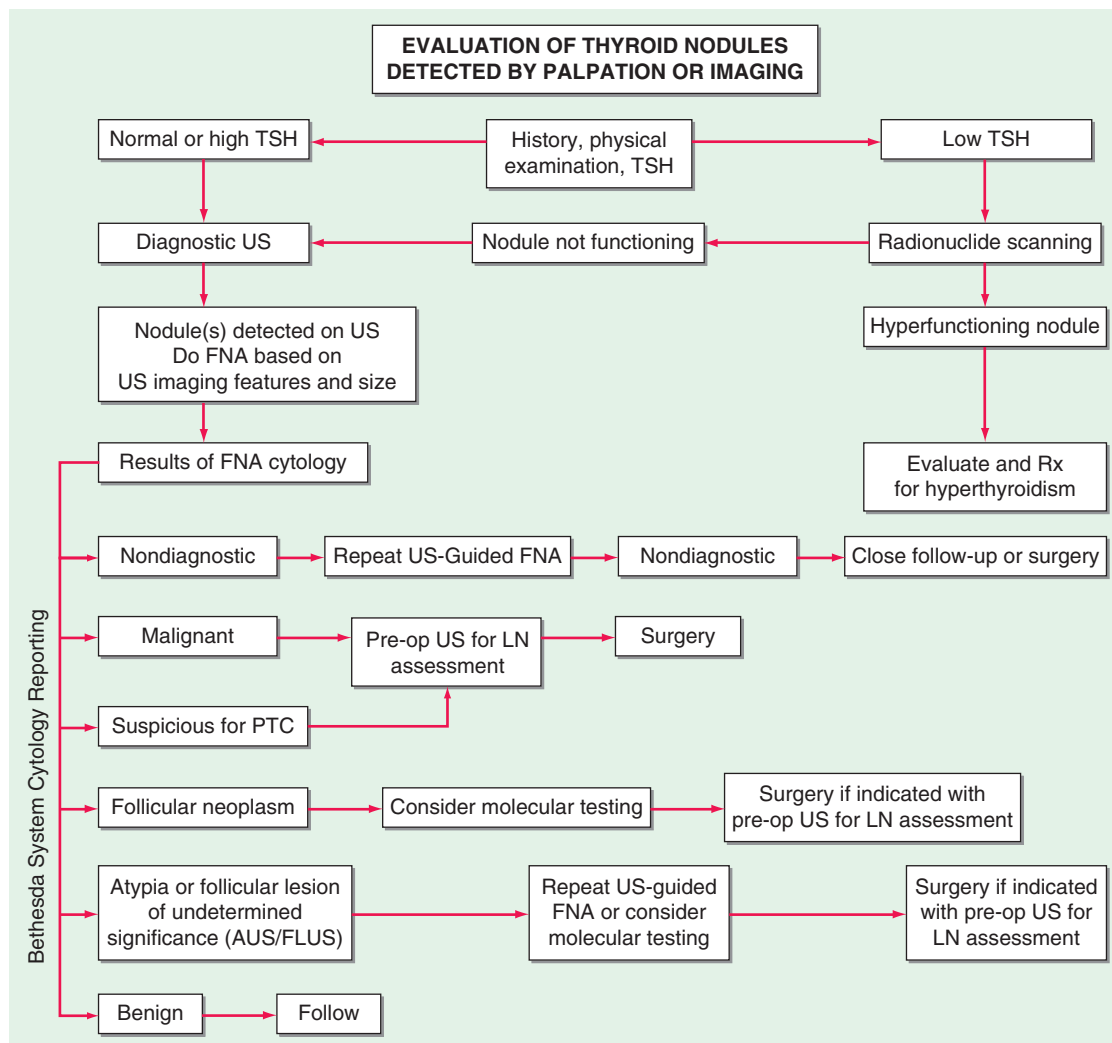


FIGURE 405-13 Approach to the patient with a thyroid nodule. See text and references for details. FNA, fine-needle aspiration; LN, lymph node; PTC, papillary thyroid cancer; TSH, thyroid-stimulating hormone; US, ultrasound.

Cytology results indicative of malignancy mandate surgery, after performing preoperative sonography to evaluate the cervical lymph nodes. Nondiagnostic cytology specimens generally result from cystic lesions but may also occur in fibrous long-standing nodules. Ultrasound-guided FNA is indicated when a repeat FNA is necessary. Repeat FNA will yield a diagnostic cytology in about 50% of cases. Benign nodules should be monitored by ultrasound for growth, and repeat FNA should be considered if the nodule enlarges. The use of levothyroxine to suppress serum TSH is not effective in shrinking nodules in iodine-replete populations, and therefore, levothyroxine should not be used.

The three cytology classifications introduced by the Bethesda System are associated with different risks of malignancy (Table 405-14).

TABLE 405-14 BETHESDA CLASSIFICATION FOR THYROID CYTOLOGY

Diagnostic Category	Risk of Malignancy
Nondiagnostic or unsatisfactory	1–5%
Benign	2–4%
Atypia or follicular lesion of unknown significance (AUS/FLUS)	15–20%
Follicular neoplasm	20–30%
Suspicious for malignancy	60–75%
Malignant	97–100%

For nodules with suspicious for malignancy cytology, surgery is recommended after ultrasound assessment of cervical lymph nodes. Options to be discussed with the patient include: (1) lobectomy with intraoperative frozen section; (2) near-total thyroidectomy; and (3) mutational analysis mainly for *BRAF V600E*, which is virtually diagnostic of PTC, and bilateral rather than unilateral thyroid surgery is required.

On the other hand, the majority of nodules with AUS/FLUS and follicular neoplasm cytology results are benign; only 10–30% are malignant. The traditional approach for these patients is diagnostic lobectomy for histopathologic diagnosis. Therefore, up to 85% of patients undergo surgery for benign nodules. A high-sensitivity (~90%) novel molecular test using gene expression profiling technology may reduce the need for unnecessary surgery in these two groups. In a multicenter trial of over 265 such nodules, a negative gene expression classifier test reduced the risk of malignancy to about 6%, leading to clinical recommendations for follow-up rather than surgery.

The evaluation of a thyroid nodule is stressful for most patients. They are concerned about the possibility of thyroid cancer, whether verbalized or not. It is constructive, therefore, to review the diagnostic approach and to reassure patients when no malignancy is found. When a suspicious lesion or thyroid cancer is identified, the generally favorable prognosis and available treatment options can be reassuring.

The adrenal cortex produces three classes of corticosteroid hormones: glucocorticoids (e.g., cortisol), mineralocorticoids (e.g., aldosterone), and adrenal androgen precursors (e.g., dehydroepiandrosterone [DHEA]) (Fig. 406-1). Glucocorticoids and mineralocorticoids act through specific nuclear receptors, regulating aspects of the physiologic stress response as well as blood pressure and electrolyte homeostasis. Adrenal androgen precursors are converted in the gonads and peripheral target cells to sex steroids that act via nuclear androgen and estrogen receptors.

Disorders of the adrenal cortex are characterized by deficiency or excess of one or several of the three major corticosteroid classes. Hormone deficiency can be caused by inherited glandular or enzymatic disorders or by destruction of the pituitary or adrenal gland by autoimmune disorders, infection, infarction, or iatrogenic events such as surgery or hormonal suppression. Hormone excess is usually

the result of neoplasia, leading to increased production of adrenocorticotropic hormone (ACTH) by the pituitary or neuroendocrine cells (ectopic ACTH) or increased production of glucocorticoids, mineralocorticoids, or adrenal androgen precursors by adrenal nodules. Adrenal nodules are increasingly identified incidentally during abdominal imaging performed for other reasons.

ADRENAL ANATOMY AND DEVELOPMENT

The normal adrenal glands weigh 6–11 g each. They are located above the kidneys and have their own blood supply. Arterial blood flows initially to the subcapsular region and then meanders from the outer cortical zona glomerulosa through the intermediate zona fasciculata to the inner zona reticularis and eventually to the adrenal medulla. The right suprarenal vein drains directly into the vena cava, while the left suprarenal vein drains into the left renal vein.

During early embryonic development, the adrenals originate from the urogenital ridge and then separate from gonads and kidneys at about the sixth week of gestation. Concordant with the time of sexual differentiation (seventh to ninth week of gestation, Chap. 410), the adrenal cortex starts to produce cortisol and the adrenal sex steroid

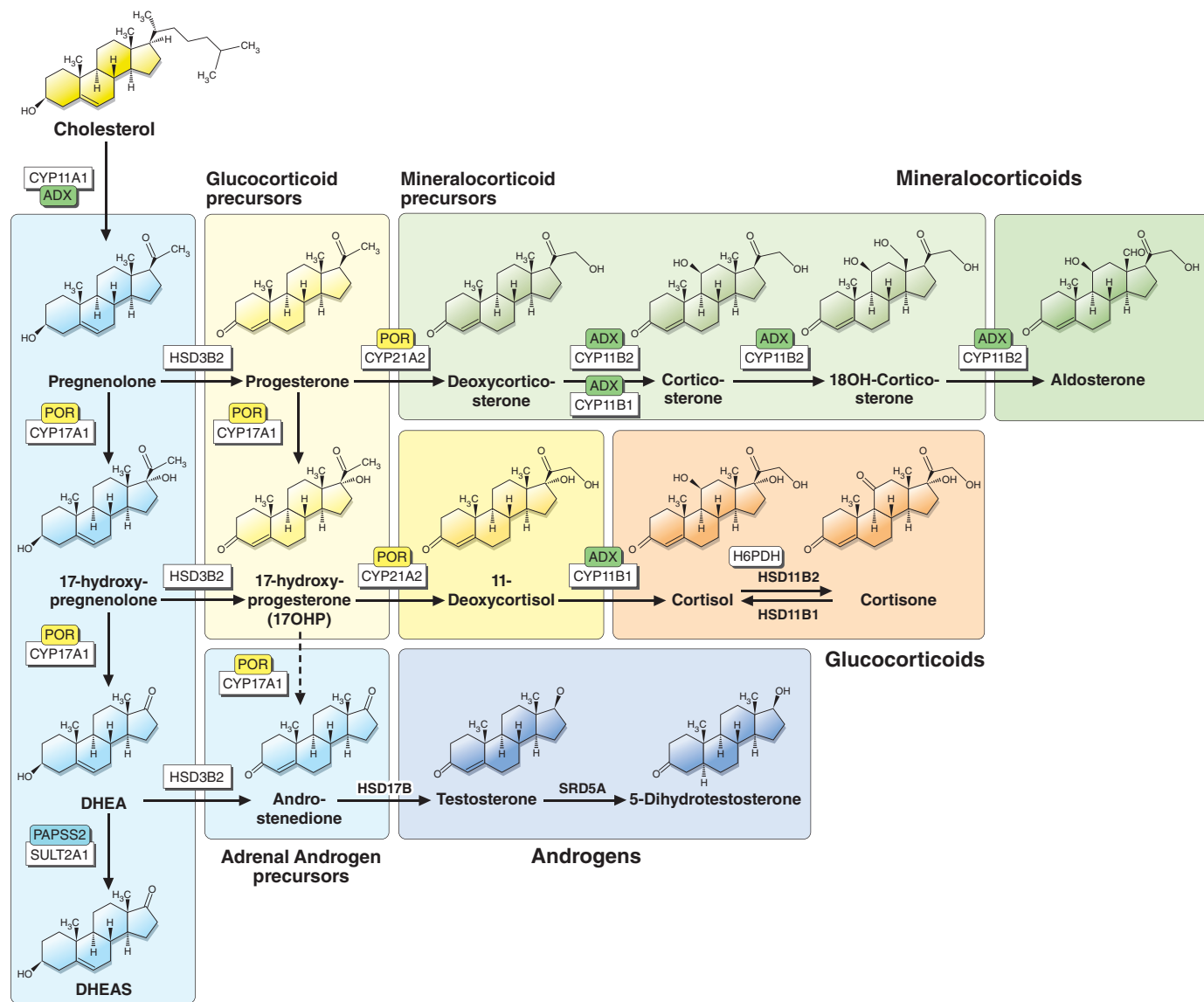


FIGURE 406-1 Adrenal steroidogenesis. ADX, adrenodoxin; CYP11A1, side chain cleavage enzyme; CYP11B1, 11 β -hydroxylase; CYP11B2, aldosterone synthase; CYP17A1, 17 α -hydroxylase/17,20 lyase; CYP21A2, 21-hydroxylase; DHEA, dehydroepiandrosterone; DHEAS, dehydroepiandrosterone sulfate; H6PDH, hexose-6-phosphate dehydrogenase; HSD11B1, 11 β -hydroxysteroid dehydrogenase type 1; HSD11B2, 11 β -hydroxysteroid dehydrogenase type 2; HSD17B, 17 β -hydroxysteroid dehydrogenase; HSD3B2, 3 β -hydroxysteroid dehydrogenase type 2; PAPSS2, PAPS synthase type 2; POR, P450 oxidoreductase; SRD5A, 5 α -reductase; SULT2A1, DHEA sulfotransferase.

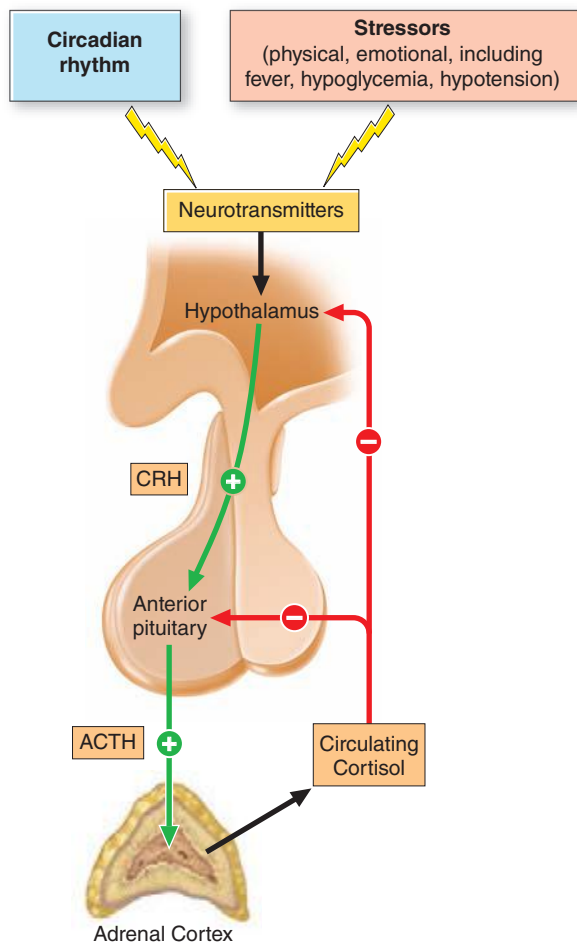


FIGURE 406-2 Regulation of the hypothalamic-pituitary-adrenal (HPA) axis. ACTH, adrenocorticotropic hormone; CRH, corticotropin-releasing hormone.

precursor DHEA. The orphan nuclear receptors SF1 (steroidogenic factor 1; encoded by the gene *NR5A1*) and DAX1 (dosage-sensitive sex reversal gene 1; encoded by the gene *NROB1*), among others, play a crucial role during this period of development, as they regulate a multitude of adrenal genes involved in steroidogenesis.

REGULATORY CONTROL OF STEROIDOGENESIS

Production of glucocorticoids and adrenal androgens is under the control of the hypothalamic-pituitary-adrenal (HPA) axis, whereas mineralocorticoids are regulated by the renin-angiotensin-aldosterone (RAA) system.

Glucocorticoid synthesis is under inhibitory feedback control by the hypothalamus and the pituitary (Fig. 406-2). Hypothalamic release of corticotropin-releasing hormone (CRH) occurs in response to endogenous or exogenous stress. CRH stimulates the cleavage of the 241-amino acid polypeptide proopiomelanocortin (POMC) by pituitary-specific prohormone convertase 1 (PC1), yielding the 39-amino acid peptide ACTH. ACTH is released by the corticotrope cells of the anterior pituitary and acts as the pivotal regulator of adrenal cortisol synthesis, with additional short-term effects on mineralocorticoid and adrenal androgen synthesis. The release of CRH, and subsequently ACTH, occurs in a pulsatile fashion that follows a circadian rhythm under the control of

the hypothalamus, specifically its suprachiasmatic nucleus (SCN), with additional regulation by a complex network of cell-specific clock genes. Reflecting the pattern of ACTH secretion, adrenal cortisol secretion exhibits a distinct circadian rhythm, starting to rise in the early morning hours prior to awakening, with peak levels in the morning and low levels in the evening (Fig. 406-3).

Diagnostic tests assessing the HPA axis make use of the fact that it is regulated by negative feedback. Glucocorticoid excess is diagnosed by employing a dexamethasone suppression test. Dexamethasone, a potent synthetic glucocorticoid, suppresses CRH/ACTH by binding hypothalamic-pituitary glucocorticoid receptors and, therefore, results in downregulation of endogenous cortisol synthesis. Various versions of the dexamethasone suppression test are described in detail in Chap. 403. If cortisol production is autonomous (e.g., adrenal nodule), ACTH is already suppressed and dexamethasone has little additional effect. If cortisol production is driven by an ACTH-producing pituitary adenoma, dexamethasone suppression is ineffective at low doses but usually induces suppression at high doses. If cortisol production is driven by an ectopic source of ACTH, the tumors are usually resistant to dexamethasone suppression. Thus, the dexamethasone suppression test is useful to establish the diagnosis of Cushing's syndrome and to assist with the differential diagnosis of cortisol excess.

Conversely, to assess glucocorticoid deficiency, ACTH stimulation of cortisol production is used. The ACTH peptide contains 39 amino acids but the first 24 are sufficient to elicit a physiologic response. The standard ACTH stimulation test involves administration of cosyntropin (ACTH 1-24), 0.25 mg IM or IV, and collection of blood samples at 0, 30, and 60 min for cortisol. A normal response is defined as a cortisol level $>20 \mu\text{g/dL}$ ($>550 \text{ nmol/L}$) 30–60 min after cosyntropin stimulation. A low-dose (1 μg cosyntropin IV) version of this test has been advocated; however, it has no superior diagnostic value and is more cumbersome to carry out. Alternatively, an insulin tolerance test (ITT) can be used to assess adrenal function. It involves injection of insulin to induce hypoglycemia, which represents a strong stress signal that triggers hypothalamic CRH release and activation of the entire HPA axis. The ITT involves administration of regular insulin 0.1 U/kg IV (dose should be lower if hypopituitarism is likely) and collection of blood samples at 0, 30, 60, and 120 min for glucose, cortisol, and growth hormone (GH), if also assessing the GH axis. Oral or IV glucose is administered after the patient has achieved symptomatic hypoglycemia (usually glucose $<40 \text{ mg/dL}$). A normal response is defined as a cortisol $>20 \mu\text{g/dL}$ and GH $>5.1 \mu\text{g/L}$. The ITT requires careful clinical

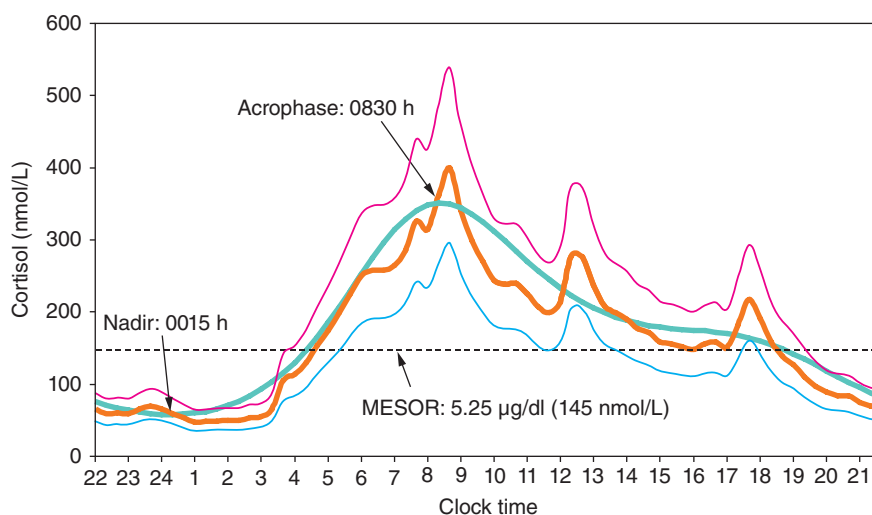


FIGURE 406-3 Physiologic cortisol circadian rhythm. Circulating cortisol concentrations (geometrical mean \pm standard deviation values and fitted cosinor) drop under the rhythm-adjusted mean (MESOR) in the early evening hours, with nadir levels around midnight and a rise in the early morning hours; peak levels are observed $\sim 8:30 \text{ AM}$ (acrophase). (Modified after M Debono et al: Modified-release hydrocortisone to provide circadian cortisol profiles. *J Clin Endocrinol Metab* 94:1548, 2009.)

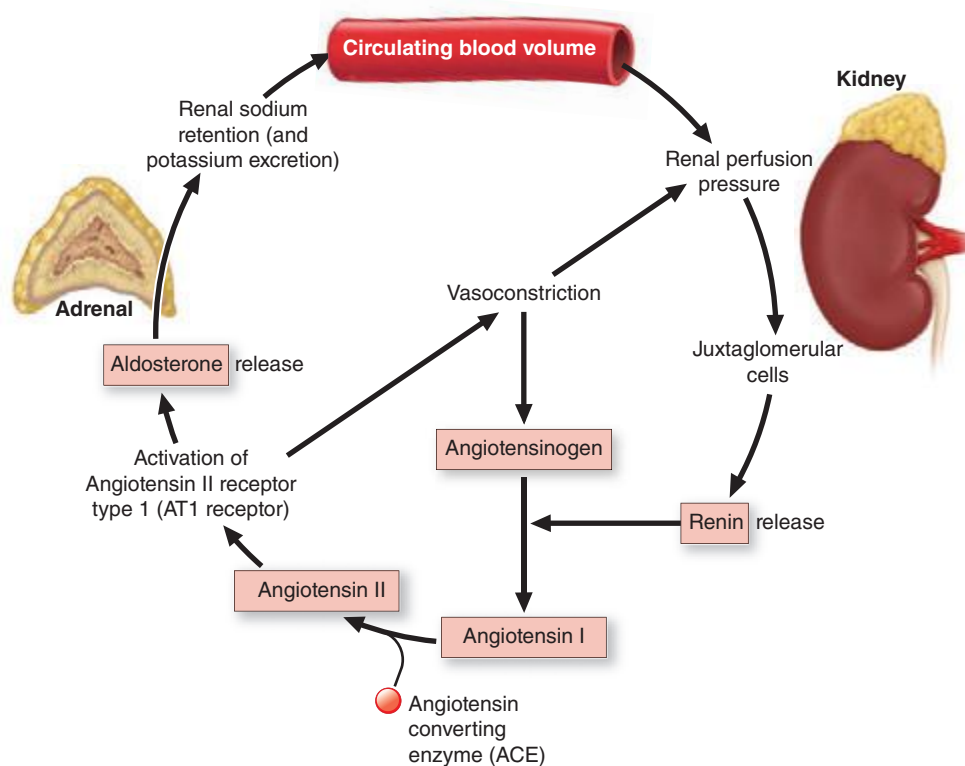


FIGURE 406-4 Regulation of the renin-angiotensin-aldosterone (RAA) system.

monitoring and sequential measurements of glucose. It is contraindicated in patients with coronary disease, cerebrovascular disease, or seizure disorders, which has made the short cosyntropin test the commonly accepted first-line test.

Mineralocorticoid production is controlled by the RAA regulatory cycle, which is initiated by the release of renin from the juxtaglomerular cells in the kidney, resulting in cleavage of angiotensinogen to angiotensin I in the liver (Fig. 406-4). Angiotensin-converting enzyme (ACE) cleaves angiotensin I to angiotensin II, which binds and activates the angiotensin II receptor type 1 (AT1 receptor [AT1R]), resulting in increased adrenal aldosterone production and vasoconstriction. Aldosterone enhances sodium retention and potassium excretion, and increases the arterial perfusion pressure, which in turn regulates renin release. Because mineralocorticoid synthesis is primarily under the control of the RAA system, hypothalamic-pituitary damage does not significantly impact the capacity of the adrenal to synthesize aldosterone.

Similar to the HPA axis, the assessment of the RAA system can be used for diagnostic purposes. If mineralocorticoid excess is present, there is a counter-regulatory downregulation of plasma renin (see below for testing). Conversely, in mineralocorticoid deficiency, plasma renin is markedly increased. Physiologically, oral or IV sodium loading results in suppression of aldosterone, a response that is attenuated or absent in patients with autonomous mineralocorticoid excess.

STEROID HORMONE SYNTHESIS, METABOLISM, AND ACTION

ACTH stimulation is required for the initiation of steroidogenesis. The ACTH receptor MC2R (melanocortin 2 receptor) interacts with the MC2R-accessory protein MRAP, and the complex is transported to the adrenocortical cell membrane, where it binds to ACTH (Fig. 406-5). ACTH stimulation generates cyclic AMP (cAMP), which upregulates the protein kinase A (PKA) signaling pathway. Inactive PKA is a tetramer of two regulatory and two catalytic subunits that is dissociated by cAMP into a dimer of two regulatory subunits bound to cAMP and two free and active catalytic subunits. PKA activation impacts steroidogenesis in three distinct ways: (1) increases the import of cholesterol esters; (2) increases the activity of hormone-sensitive lipase, which cleaves cholesterol esters to cholesterol for import into the mitochondrion; and (3) increases the availability and phosphorylation

of CREB (cAMP response element binding), a transcription factor that enhances transcription of CYP11A1 and other enzymes required for glucocorticoid synthesis.

Adrenal steroidogenesis occurs in a zone-specific fashion, with mineralocorticoid synthesis occurring in the outer zona glomerulosa, glucocorticoid synthesis in the zona fasciculata, and adrenal androgen synthesis in the inner zona reticularis (Fig. 406-1). All steroidogenic pathways require cholesterol import into the mitochondrion, a process initiated by the action of the steroidogenic acute regulatory (StAR) protein, which shuttles cholesterol from the outer to the inner mitochondrial membrane. The majority of steroidogenic enzymes are cytochrome P450 (CYP) enzymes, which are either located in the mitochondrion (side chain cleavage enzyme, CYP11A1; 11 β -hydroxylase, CYP11B1; aldosterone synthase, CYP11B2) or in the endoplasmic reticulum membrane (17 α -hydroxylase, CYP17A1; 21-hydroxylase, CYP21A2; aromatase, CYP19A1). These enzymes require electron donation via specific redox cofactor enzymes, P450 oxidoreductase (POR), and adrenodoxin/adrenodoxin reductase (ADX/ADR) for the microsomal and mitochondrial CYP enzymes, respectively. In addition, the short-chain dehydrogenase 3 β -hydroxysteroid dehydrogenase type 2 (3 β -HSD2), also termed Δ 4, Δ 5 isomerase, plays a major role in adrenal steroidogenesis.

The cholesterol side chain cleavage enzyme CYP11A1 generates pregnenolone. Glucocorticoid synthesis requires conversion of pregnenolone to progesterone by 3 β -HSD2, followed by conversion to 17-hydroxyprogesterone by CYP17A1, further hydroxylation at carbon 21 by CYP21A2, and eventually, 11 β -hydroxylation by CYP11B1 to generate active cortisol (Fig. 406-1). Mineralocorticoid synthesis also requires progesterone, which is first converted to deoxycorticosterone by CYP21A2 and then converted via corticosterone and 18-hydroxycorticosterone to aldosterone in three steps catalyzed by CYP11B2. For adrenal androgen synthesis, pregnenolone undergoes conversion by CYP17A1, which uniquely catalyzes two enzymatic reactions. Via its 17 α -hydroxylase activity, CYP17A1 converts pregnenolone to 17-hydroxypregnenolone, followed by generation of the universal sex steroid precursor DHEA via CYP17A1 17,20 lyase activity. The majority of DHEA is secreted by the adrenal in the form of its sulfate ester, DHEAS, generated by DHEA sulfotransferase (SULT2A1).

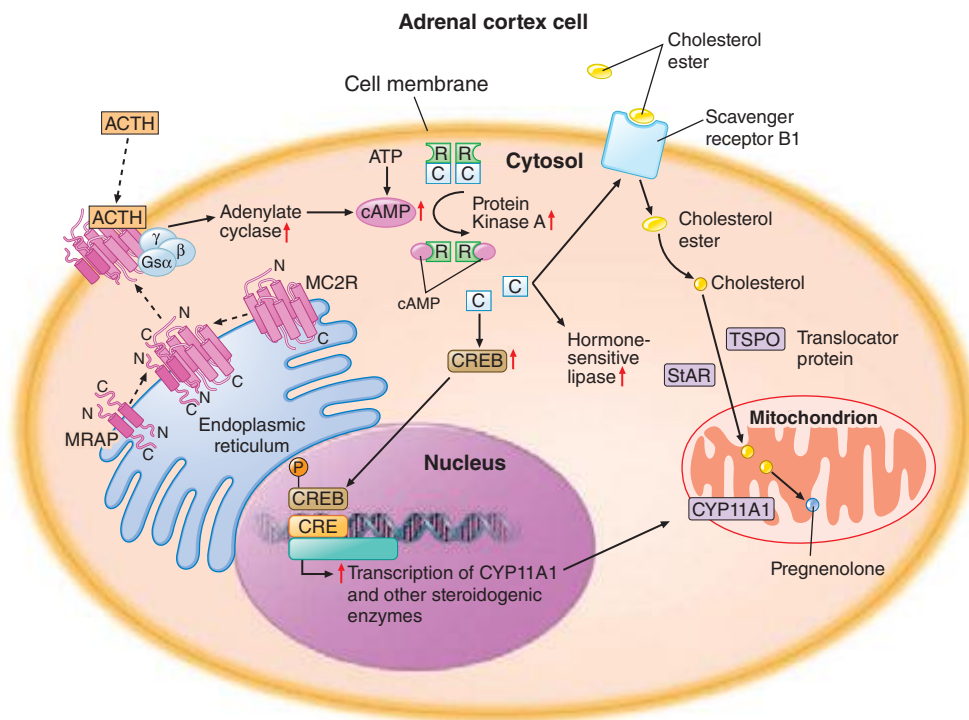


FIGURE 406-5 ACTH effects on adrenal steroidogenesis. ACTH, adrenocorticotropic hormone; binding protein; MRAP, MC2R-accessory protein; protein kinase A catalytic subunit (C; *PRKACA*), PKA regulatory subunit (R; *PRKAR1A*); StAR, steroidogenic acute regulatory (protein); TSPO, translocator protein.

Following its release from the adrenal, cortisol circulates in the bloodstream mainly bound to cortisol-binding globulin (CBG) and to a lesser extent to albumin, with only a minor fraction circulating as free, unbound hormone. Free cortisol is thought to enter cells directly, not requiring active transport. In addition, in a multitude of peripheral target tissues of glucocorticoid action, including adipose, liver, muscle, and brain, cortisol is generated from inactive cortisone within the cell by the enzyme 11 β -hydroxysteroid dehydrogenase type 1 (11 β -HSD1) (Fig. 406-6). Thereby, 11 β -HSD1 functions as a tissue-specific prereceptor regulator of glucocorticoid action. For the conversion

of inactive cortisone to active cortisol, 11 β -HSD1 requires nicotinamide adenine dinucleotide phosphate (NADPH [reduced form]), which is provided by the enzyme hexose-6-phosphate dehydrogenase (H6PDH). Like the catalytic domain of 11 β -HSD1, H6PDH is located in the lumen of the endoplasmic reticulum, and converts glucose-6-phosphate (G6P) to 6-phosphogluconate (6PGL), thereby regenerating NADP⁺ to NADPH, which drives the activation of cortisol from cortisone by 11 β -HSD1.

In the cytosol of target cells, cortisol binds and activates the glucocorticoid receptor (GR), which results in dissociation of heat shock

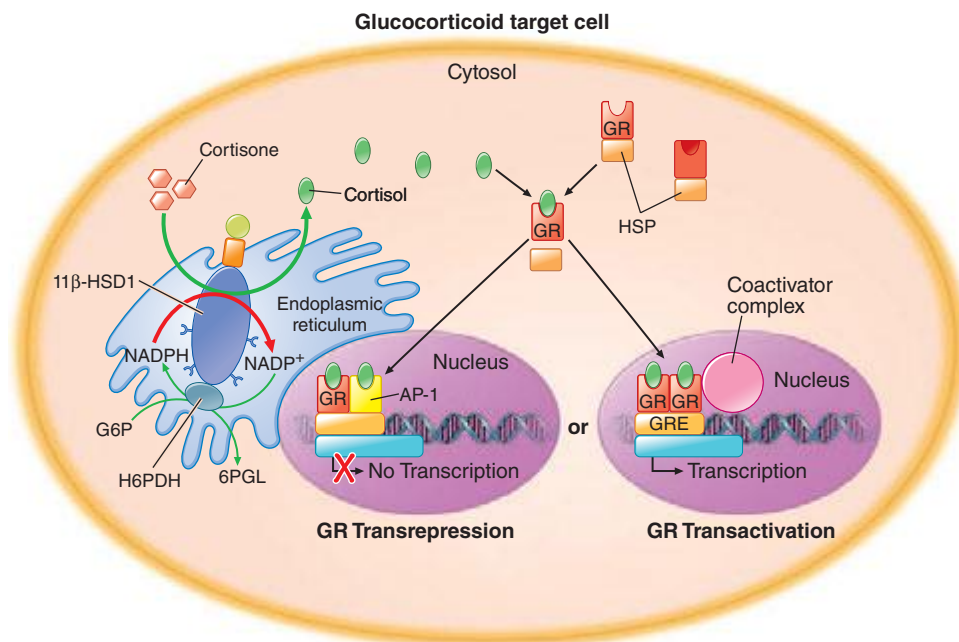


FIGURE 406-6 Prereceptor activation of cortisol and glucocorticoid receptor (GR) action. AP-1 activator protein-1; G6P, glucose-6-phosphate; GRE, glucocorticoid response elements; HSP, heat shock proteins; NADPH, nicotinamide adenine dinucleotide phosphate (reduced form); 6PGL, 6-phosphogluconate.

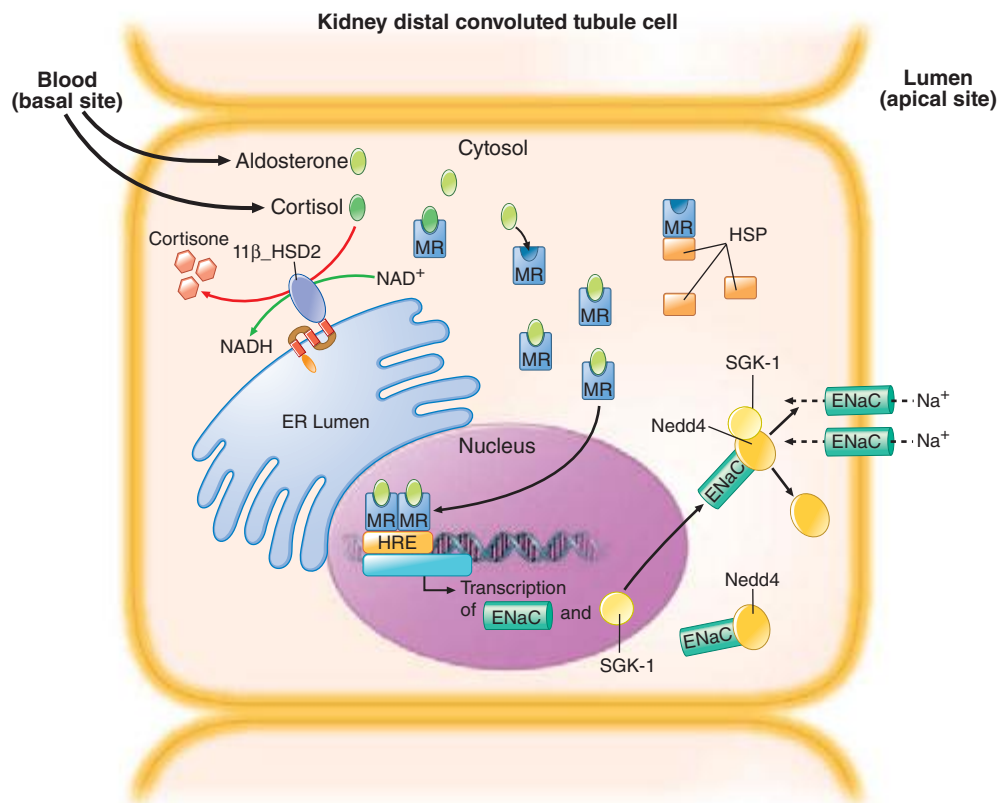


FIGURE 406-7 Prereceptor inactivation of cortisol and mineralocorticoid receptor action. ENaC, epithelial sodium channel; HRE, hormone response element; NADH, nicotinamide adenine dinucleotide; SGK-1, serum glucocorticoid-inducible kinase-1.

proteins (HSP) from the receptor and subsequent dimerization (Fig. 406-6). Cortisol-bound GR dimers translocate to the nucleus and activate glucocorticoid response elements (GRE) in the DNA sequence, thereby enhancing transcription of glucocorticoid-regulated genes (GR transactivation). However, cortisol-bound GR can also form heterodimers with transcription factors such as AP-1 or NF- κ B, resulting in transrepression of proinflammatory genes, a mechanism of major importance for the anti-inflammatory action of glucocorticoids. It is important to note that corticosterone also exerts glucocorticoid activity, albeit much weaker than cortisol itself. However, in rodents, corticosterone is the major glucocorticoid, and in patients with 17-hydroxylase deficiency, lack of cortisol can be compensated for by higher concentrations of corticosterone that accumulates as a consequence of the enzymatic block.

Cortisol is inactivated to cortisone by the microsomal enzyme 11 β -hydroxysteroid dehydrogenase type 2 (11 β -HSD2) (Fig. 406-7), mainly in the kidney, but also in the colon, salivary glands, and other target tissues. Cortisol and aldosterone bind the mineralocorticoid receptor (MR) with equal affinity; however, cortisol circulates in the bloodstream at about a thousandfold higher concentration. Thus, only rapid inactivation of cortisol to cortisone by 11 β -HSD2 prevents MR activation by excess cortisol, thereby acting as a tissue-specific modulator of the MR pathway. In addition to cortisol and aldosterone, deoxycorticosterone (DOC) (Fig. 406-1) also exerts mineralocorticoid activity. DOC accumulation due to 11 β -hydroxylase deficiency or due to tumor-related excess production can result in mineralocorticoid excess.

Aldosterone synthesis in the adrenal zona glomerulosa cells is driven by the enzyme aldosterone synthase (CYP11B2). The binding of angiotensin II to the AT1 receptor causes glomerulosa cell membrane depolarization by increasing intracellular sodium through inhibition of sodium potassium (Na⁺/K⁺) ATPase enzymes as well as potassium channels. This drives an increase in intracellular calcium by opening of voltage-dependent calcium channels or inhibition of calcium (Ca²⁺) ATPase enzymes. Consequently, the calcium signaling pathway is triggered, resulting in upregulation of CYP11B2 transcription (Fig. 406-8).

Analogous to cortisol action via the GR, aldosterone (or cortisol) binding to the MR in the kidney tubule cell dissociates the HSP–receptor complex, allowing homodimerization of the MR, and translocation of the hormone-bound MR dimer to the nucleus (Fig. 406-7). The activated MR enhances transcription of the epithelial sodium channel (ENaC) and serum glucocorticoid-inducible kinase 1 (SGK-1). In the cytosol, interaction of ENaC with Nedd4 prevents cell surface expression of ENaC. However, SGK-1 phosphorylates serine residues within the Nedd4 protein, reduces the interaction between Nedd4 and ENaC, and consequently, enhances the trafficking of ENaC to the cell surface, where it mediates sodium retention.

CUSHING'S SYNDROME

(See also Chap. 403) Cushing's syndrome reflects a constellation of clinical features that result from chronic exposure to excess glucocorticoids of any etiology. The disorder can be ACTH-dependent (e.g., pituitary corticotrope adenoma, ectopic secretion of ACTH by nonpituitary tumor) or ACTH-independent (e.g., adrenocortical adenoma, adrenocortical carcinoma, nodular adrenal hyperplasia), as well as iatrogenic (e.g., administration of exogenous glucocorticoids to treat various inflammatory conditions). The term *Cushing's disease* refers specifically to Cushing's syndrome caused by a pituitary corticotrope adenoma.

Epidemiology Cushing's syndrome is generally considered a rare disease. It occurs with an incidence of 1–2 per 100,000 population per year. However, it is debated whether mild cortisol excess may be more prevalent among patients with several features of Cushing's such as centripetal obesity, type 2 diabetes, and osteoporotic vertebral fractures, recognizing that these are relatively nonspecific and common in the population.

In the overwhelming majority of patients, Cushing's syndrome is caused by an ACTH-producing corticotrope adenoma of the pituitary (Table 406-1), as initially described by Harvey Cushing in 1912. Cushing's disease more frequently affects women, with the exception of prepubertal cases, where it is more common in boys. By contrast,

Adrenal zona glomerulosa cell

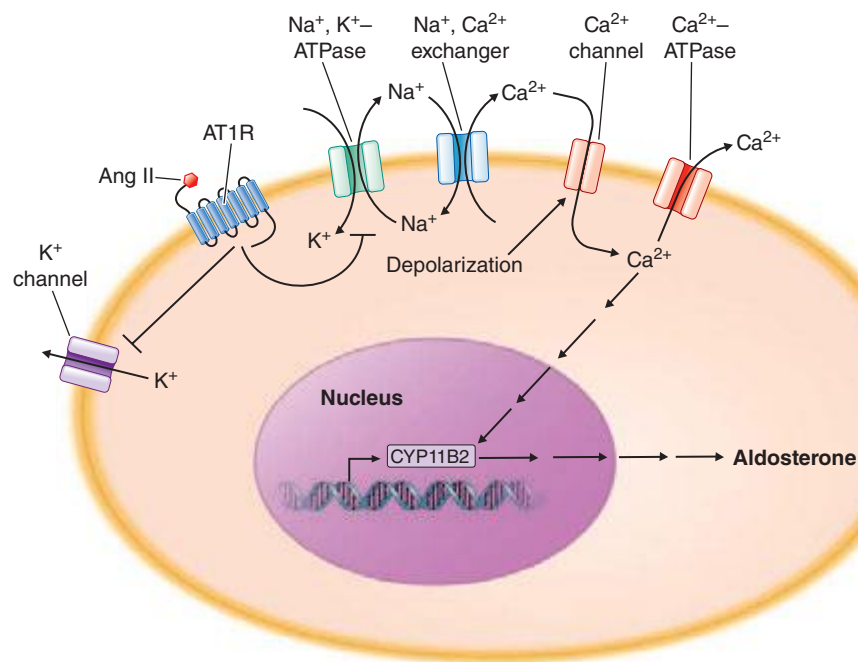


FIGURE 406-8 Regulation of adrenal aldosterone synthesis. AngII, angiotensin II; AT1R, angiotensin II receptor type 1; CYP11B2, aldosterone synthase. (Modified after F Beuschlein: Regulation of aldosterone secretion: from physiology to disease. *Eur J Endocrinol* 168:R85, 2013.)

ectopic ACTH syndrome is more frequently identified in men. Only 10% of patients with Cushing's syndrome have a primary, adrenal cause of their disease (e.g., autonomous cortisol excess independent of ACTH), and most of these patients are women. Overall, the medical use of glucocorticoids for immunosuppression, or for the treatment of inflammatory disorders, is the most common cause of Cushing's syndrome.

Etiology In at least 90% of patients with Cushing's disease, ACTH excess is caused by a corticotrope pituitary microadenoma, often only a few millimeters in diameter. Pituitary macroadenomas (i.e., tumors >1 cm in size) are found in only 5–10% of patients. Pituitary corticotrope adenomas usually occur sporadically but very rarely can be found in the context of multiple endocrine neoplasia type 1 (MEN 1) (Chap. 408).

Ectopic ACTH production is predominantly caused by occult carcinoid tumors, most frequently in the lung, but also in thymus or pancreas. Because of their small size, these tumors are often difficult to locate. Advanced small-cell lung cancer can cause ectopic ACTH

production. In rare cases, ectopic CRH and/or ACTH production has been found to originate from medullary thyroid carcinoma or pheochromocytoma, the latter co-secreting catecholamines and ACTH.

The majority of patients with ACTH-independent cortisol excess harbor a cortisol-producing adrenal adenoma; intratumor mutations, i.e., somatic mutations in the PKA catalytic subunit PRKACA, have been identified as cause of disease in 40% of these tumors. Adrenocortical carcinomas may also cause ACTH-independent disease and are often large, with excess production of several corticosteroid classes.

A rare but notable cause of adrenal cortisol excess is macronodular adrenal hyperplasia with low circulating ACTH, but with evidence for autocrine stimulation of cortisol production via intraadrenal ACTH production. These hyperplastic nodules are often also characterized by ectopic expression of G protein-coupled receptors not usually found in the adrenal, including receptors for luteinizing hormone, vasopressin, serotonin, interleukin 1, catecholamines, or gastric inhibitory peptide (GIP), the cause of food-dependent Cushing's. Activation of these receptors results in upregulation of PKA signaling, as physiologically occurs with ACTH, with a subsequent increase in cortisol production. A combination of germline and somatic mutations in the tumor-suppressor gene *ARMC5* have been identified as a prevalent cause of Cushing's due to macronodular adrenal hyperplasia. Germline mutations in the PKA catalytic subunit PRKACA can represent a rare cause of macronodular adrenal hyperplasia associated with cortisol excess.

Mutations in one of the regulatory subunits of PKA, PRKARIA, are found in patients with primary pigmented nodular adrenal disease (PPNAD) as part of *Carney's complex*, an autosomal dominant multiple neoplasia condition associated with cardiac myxomas, hyperlentiginosis, Sertoli cell tumors, and PPNAD. PPNAD can present as micronodular or macronodular hyperplasia, or both. Phosphodiesterases can influence intracellular cAMP and can thereby impact PKA activation. Mutations in PDE11A and PDE8B have been identified in patients with bilateral adrenal hyperplasia and Cushing's, with and without evidence of PPNAD.

Another rare cause of ACTH-independent Cushing's is *McCune-Albright syndrome*, also associated with polyostotic fibrous dysplasia, unilateral café-au-lait spots, and precocious puberty. McCune-Albright syndrome is caused by activating mutations in the stimulatory

TABLE 406-1 CAUSES OF CUSHING'S SYNDROME

Causes of Cushing's Syndrome	Female:Male Ratio	%
ACTH-Dependent Cushing's		90
Cushing's disease (= ACTH-producing pituitary adenoma)	4:1	75
Ectopic ACTH syndrome (due to ACTH secretion by bronchial or pancreatic carcinoid tumors, small-cell lung cancer, medullary thyroid carcinoma, pheochromocytoma and others)	1:1	15
ACTH-Independent Cushing's	4:1	10
Adrenocortical adenoma		5–10
Adrenocortical carcinoma		1
Rare causes: macronodular adrenal hyperplasia; primary pigmented nodular adrenal disease (micro- and/or macronodular); McCune-Albright syndrome		<1

Abbreviation: ACTH, adrenocorticotrophic hormone.

TABLE 406-2 SIGNS AND SYMPTOMS OF CUSHING'S SYNDROME

Body Compartment/System	Signs and Symptoms
Body fat	Weight gain, central obesity, rounded face, fat pad on back of neck ("buffalo hump")
Skin	Facial plethora, thin and brittle skin, easy bruising, broad and purple stretch marks, acne, hirsutism
Bone	Osteopenia, osteoporosis (vertebral fractures), decreased linear growth in children
Muscle	Weakness, proximal myopathy (prominent atrophy of gluteal and upper leg muscles with difficulty climbing stairs or getting up from a chair)
Cardiovascular system	Hypertension, hypokalemia, edema, atherosclerosis
Metabolism	Glucose intolerance/diabetes, dyslipidemia
Reproductive system	Decreased libido, in women amenorrhea (due to cortisol-mediated inhibition of gonadotropin release)
Central nervous system	Irritability, emotional lability, depression, sometimes cognitive defects; in severe cases, paranoid psychosis
Blood and immune system	Increased susceptibility to infections, increased white blood cell count, eosinopenia, hypercoagulation with increased risk of deep vein thrombosis and pulmonary embolism

G protein alpha subunit 1, GNAS-1 (guanine nucleotide binding protein alpha stimulating activity polypeptide 1), and such mutations have also been found in bilateral macronodular hyperplasia without other McCune-Albright features and, in rare instances, also in isolated cortisol-producing adrenal adenomas (Table 406-1; Chap. 426).

Clinical Manifestations Glucocorticoids affect almost all cells of the body, and thus signs of cortisol excess impact multiple physiologic systems (Table 406-2), with upregulation of gluconeogenesis, lipolysis, and

protein catabolism causing the most prominent features. In addition, excess glucocorticoid secretion overcomes the ability of 11β -HSD2 to rapidly inactivate cortisol to cortisone in the kidney, thereby exerting mineralocorticoid actions, manifest as diastolic hypertension, hypokalemia, and edema. Excess glucocorticoids also interfere with central regulatory systems, leading to suppression of gonadotropins with subsequent hypogonadism and amenorrhea, and suppression of the hypothalamic-pituitary-thyroid axis, resulting in decreased thyroid-stimulating hormone (TSH) secretion.

The majority of clinical signs and symptoms observed in Cushing's syndrome are relatively nonspecific and include features such as obesity, diabetes, diastolic hypertension, hirsutism, and depression that are commonly found in patients who do not have Cushing's. Therefore, careful clinical assessment is an important aspect of evaluating suspected cases. A diagnosis of Cushing's should be considered when several clinical features are found in the same patient, in particular when more specific features are found. These include fragility of the skin, with easy bruising and broad (>1 cm), purplish striae (Fig. 406-9), and signs of proximal myopathy, which becomes most obvious when trying to stand up from a chair without the use of hands or when climbing stairs. Clinical manifestations of Cushing's do not differ substantially among the different causes of Cushing's. In ectopic ACTH syndrome, hyperpigmentation of the knuckles, scars, or skin areas exposed to increased friction can be observed (Fig. 406-9) and is caused by stimulatory effects of excess ACTH and other POMC cleavage products on melanocyte pigment production. Furthermore, patients with ectopic ACTH syndrome, and some with adrenocortical carcinoma as the cause of Cushing's, may have a more brisk onset and rapid progression of clinical signs and symptoms.

Patients with Cushing's syndrome can be acutely endangered by deep vein thrombosis, with subsequent pulmonary embolism due to a hypercoagulable state associated with Cushing's. The majority of patients also experience psychiatric symptoms, mostly in the form of anxiety or depression, but acute paranoid or depressive psychosis



FIGURE 406-9 Clinical features of Cushing's syndrome. **A.** Note central obesity and broad, purple stretch marks (**B.** close-up). **C.** Note thin and brittle skin in an elderly patient with Cushing's syndrome. **D.** Hyperpigmentation of the knuckles in a patient with ectopic adrenocorticotrophic hormone (ACTH) excess.

2316 may also occur. Even after cure, long-term health may be affected by persistently impaired health-related quality of life and increased risk of cardiovascular disease and osteoporosis with vertebral fractures, depending on the duration and degree of exposure to significant cortisol excess.

Diagnosis The most important first step in the management of patients with suspected Cushing's syndrome is to establish the correct diagnosis. Most mistakes in clinical management, leading to unnecessary imaging or surgery, are made because the diagnostic protocol is not followed (Fig. 406-10). This protocol requires establishing the diagnosis of Cushing's beyond doubt prior to employing any tests used for the differential diagnosis of the condition. In principle, after excluding exogenous glucocorticoid use as the cause of clinical signs and symptoms, suspected cases should be tested if there are multiple and progressive features of Cushing's, particularly features with a potentially higher discriminatory value. Exclusion of Cushing's is also indicated in patients with incidentally discovered adrenal masses.

A diagnosis of Cushing's can be considered as established if the results of several tests are consistently suggestive of Cushing's. These tests may include increased 24-h urinary free cortisol excretion in three separate collections, failure to appropriately suppress morning cortisol after overnight exposure to dexamethasone, and evidence of loss of diurnal cortisol secretion with high levels at midnight, the time of the physiologically lowest secretion (Fig. 406-10). Factors potentially affecting the outcome of these diagnostic tests have to be excluded such as incomplete 24-h urine collection or rapid inactivation of dexamethasone due to concurrent intake of CYP3A4-inducing drugs (e.g., antiepileptics, rifampicin). Concurrent intake of oral contraceptives that raise CBG and thus total cortisol can cause failure to suppress after dexamethasone. If in doubt, testing should be repeated after 4–6 weeks off estrogens. Patients with pseudo-Cushing states, i.e., alcohol-related, and those with cyclic Cushing's may require further testing to safely confirm or exclude the diagnosis of Cushing's. In addition, the biochemical assays employed can affect the test results, with specificity representing a common problem with antibody-based

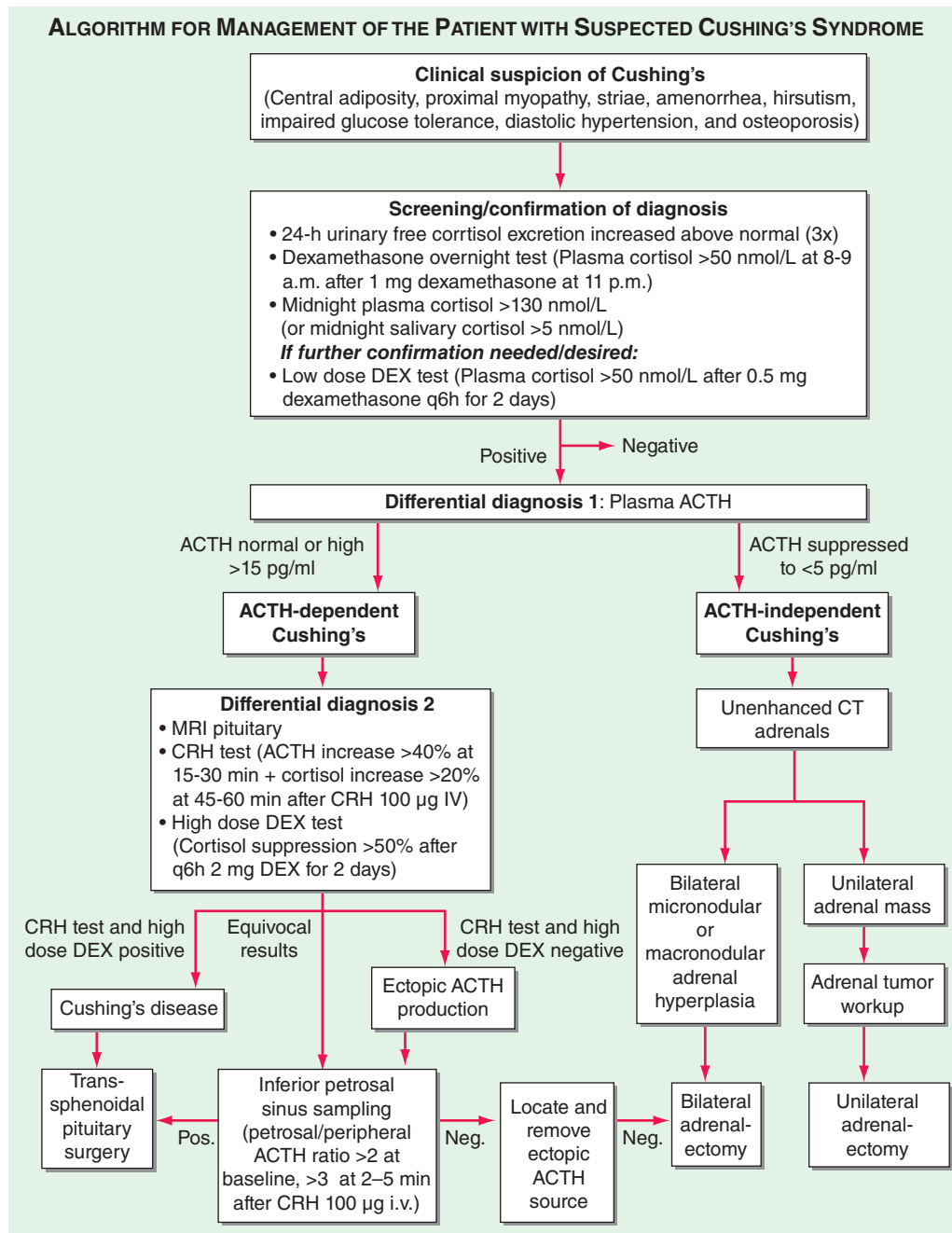


FIGURE 406-10 Management of the patient with suspected Cushing's syndrome. ACTH, adrenocorticotropic hormone; CRH, corticotropin-releasing hormone; CT, computed tomography; DEX, dexamethasone; MRI, magnetic resonance imaging.

assays for the measurement of urinary free cortisol. These assays have been greatly improved by the introduction of highly specific tandem mass spectrometry.

Differential Diagnosis The evaluation of patients with confirmed Cushing's should be carried out by an endocrinologist and begins with the differential diagnosis of ACTH-dependent and ACTH-independent cortisol excess (Fig. 406-10). Generally, plasma ACTH levels are suppressed in cases of autonomous adrenal cortisol excess, as a consequence of enhanced negative feedback to the hypothalamus and pituitary. By contrast, patients with ACTH-dependent Cushing's have normal or increased plasma ACTH, with very high levels being found in some patients with ectopic ACTH syndrome. Importantly, imaging should only be used after it is established whether the cortisol excess is ACTH-dependent or ACTH-independent, because nodules in the pituitary or the adrenal are a common finding in the general population. In patients with confirmed ACTH-independent excess, adrenal imaging is indicated (Fig. 406-11), preferably using an unenhanced computed tomography (CT) scan. This allows assessment of adrenal morphology and determination of precontrast tumor density in Hounsfield units (HU), which helps to distinguish between benign and malignant adrenal lesions.

For ACTH-dependent cortisol excess (Chap. 403), a magnetic resonance image (MRI) of the pituitary is the investigation of choice, but it may not show an abnormality in up to 40% of cases because of small tumors below the sensitivity of detection. Characteristically, pituitary corticotrope adenomas fail to enhance following gadolinium administration on T1-weighted MRI images. In all cases of confirmed ACTH-dependent Cushing's, further tests are required for the differential diagnosis of pituitary Cushing's disease and ectopic ACTH syndrome.

These tests exploit the fact that most pituitary corticotrope adenomas still display regulatory features, including residual ACTH suppression by high-dose glucocorticoids and CRH responsiveness. In contrast, ectopic sources of ACTH are typically resistant to dexamethasone suppression and unresponsive to CRH (Fig. 406-10). However, it should be noted that a small minority of ectopic ACTH-producing tumors exhibit dynamic responses similar to pituitary corticotrope tumors. If the two tests show discordant results, or if there is any other reason for doubt, the differential diagnosis can be further clarified by performing bilateral inferior petrosal sinus sampling (IPSS) with concurrent blood sampling for ACTH in the right and left inferior petrosal sinus and a peripheral vein. An increased central/peripheral plasma ACTH ratio >2 at baseline and >3 at 2–5 min after CRH injection is indicative of Cushing's disease (Fig. 406-10), with very high sensitivity and specificity. Of note, the results of the IPSS cannot be reliably used for lateralization (i.e., prediction of the location of the tumor within the pituitary), because there is broad interindividual variability in the venous drainage of the pituitary region. Importantly, no cortisol-lowering agents should be used prior to IPSS.

If the differential diagnostic testing indicates ectopic ACTH syndrome, then further imaging should include high-resolution, fine-cut CT scanning of the chest and abdomen for scrutiny of the lung, thymus, and pancreas. If no lesions are identified, an MRI of the chest can be considered because carcinoid tumors usually show high signal intensity on T2-weighted images. Furthermore, octreotide scintigraphy can be helpful in some cases because ectopic ACTH-producing tumors often express somatostatin receptors. Depending on the suspected cause, patients with ectopic ACTH syndrome should also undergo blood sampling for fasting gut hormones, chromogranin A, calcitonin, and biochemical exclusion of pheochromocytoma.

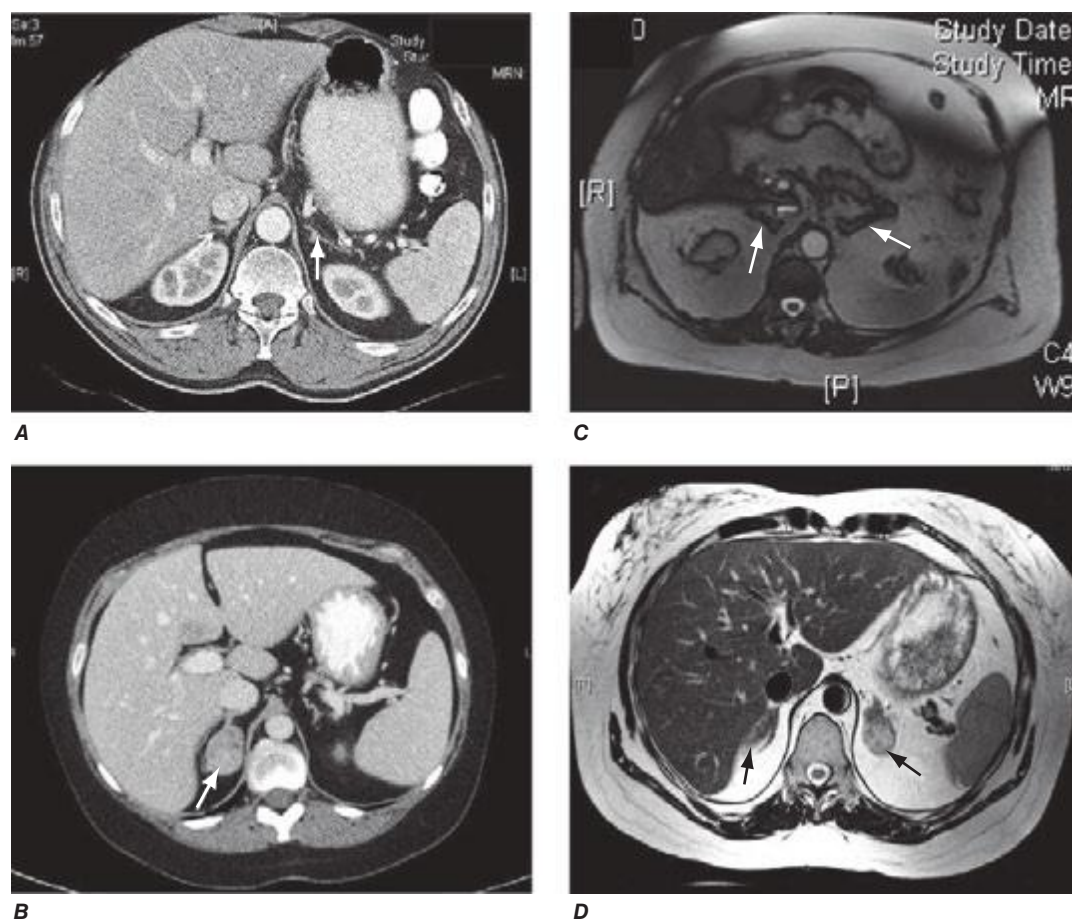


FIGURE 406-11 Adrenal imaging in Cushing's syndrome. **A.** Adrenal computed tomography (CT) showing normal bilateral adrenal morphology (arrows). **B.** CT scan depicting a right adrenocortical adenoma (arrow) causing Cushing's syndrome. **C.** Magnetic resonance imaging (MRI) showing bilateral adrenal hyperplasia due to excess adrenocorticotrophic hormone stimulation in Cushing's disease. **D.** MRI showing bilateral macronodular hyperplasia causing Cushing's syndrome.

Overt Cushing's is associated with a poor prognosis if left untreated. In ACTH-independent disease, treatment consists of surgical removal of the adrenal tumor. For smaller tumors, a minimally invasive approach can be used, whereas for larger tumors and those suspected of malignancy, an open approach is preferred.

In Cushing's disease, the treatment of choice is selective removal of the pituitary corticotrope tumor, usually via an endoscopic transphenoidal approach. This results in an initial cure rate of 70–80% when performed by a highly experienced surgeon. However, even after initial remission following surgery, long-term follow-up is important because late relapse occurs in a significant number of patients. If pituitary disease recurs, there are several options, including second surgery, radiotherapy, stereotactic radiosurgery, and bilateral adrenalectomy. These options need to be applied in a highly individualized fashion.

In some patients with very severe, overt Cushing's (e.g., difficult to control hypokalemic hypertension or acute psychosis), it may be necessary to introduce medical therapy to rapidly control the cortisol excess during the period leading up to surgery. Similarly, patients with metastasized, glucocorticoid-producing carcinomas may require long-term antiglucocorticoid drug treatment. In case of ectopic ACTH syndrome, in which the tumor cannot be located, one must carefully weigh whether drug treatment or bilateral adrenalectomy is the most appropriate choice, with the latter facilitating immediate cure but requiring life-long corticosteroid replacement. In this instance, it is paramount to ensure regular imaging follow-up for identification of the ectopic ACTH source.

Oral agents with established efficacy in Cushing's syndrome are metyrapone and ketoconazole. Metyrapone inhibits cortisol synthesis at the level of 11 β -hydroxylase (Fig. 406-1), whereas the antimycotic drug ketoconazole inhibits the early steps of steroidogenesis. Typical starting doses are 500 mg tid for metyrapone (maximum dose, 6 g) and 200 mg tid for ketoconazole (maximum dose, 1200 mg). Mitotane, a derivative of the insecticide o,p'DDD, is an adrenolytic agent that is also effective for reducing cortisol. Because of its side effect profile, it is most commonly used in the context of adrenocortical carcinoma, but low-dose treatment (500–1000 mg/d)

has also been used in benign Cushing's. In severe cases of cortisol excess, etomidate can be used to lower cortisol. It is administered by continuous IV infusion in low, nonanesthetic doses.

After the successful removal of an ACTH- or cortisol-producing tumor, the HPA axis will remain suppressed. Thus, hydrocortisone replacement needs to be initiated at the time of surgery and slowly tapered following recovery, to allow physiologic adaptation to normal cortisol levels. Depending on degree and duration of cortisol excess, the HPA axis may require many months or even years to resume normal function.

MINERALOCORTICOID EXCESS

Epidemiology Following the first description of a patient with an aldosterone-producing adrenal adenoma (*Conn's syndrome*), mineralocorticoid excess was thought to represent a rare cause of hypertension. However, in studies systematically screening all patients with hypertension, a much higher prevalence is now recognized, ranging from 5 to 12%. The prevalence is higher when patients are preselected for hypokalemic hypertension.

Etiology The most common cause of mineralocorticoid excess is primary aldosteronism, reflecting excess production of aldosterone by the adrenal zona glomerulosa. Bilateral micronodular hyperplasia is somewhat more common than unilateral adrenal adenomas (Table 406-3). Somatic mutations in channels and enzymes responsible for increasing sodium and calcium influx in adrenal zona glomerulosa cells have been identified as prevalent causes of aldosterone-producing adrenal adenomas (Table 406-3) and, in the case of germline mutations, also of primary aldosteronism due to bilateral macronodular adrenal hyperplasia. However, bilateral adrenal hyperplasia as a cause of mineralocorticoid excess is usually micronodular but can also contain larger nodules that might be mistaken for a unilateral adenoma. In rare instances, primary aldosteronism is caused by an adrenocortical carcinoma. Carcinomas should be considered in younger patients and in those with larger tumors, because benign aldosterone-producing adenomas usually measure <2 cm in diameter.

A rare cause of aldosterone excess is glucocorticoid-remediable aldosteronism (GRA), which is caused by a chimeric gene resulting

TABLE 406-3 CAUSES OF MINERALOCORTICOID EXCESS

Causes of Mineralocorticoid Excess	Mechanism	%
Primary Aldosteronism		
Adrenal (Conn's) adenoma	Autonomous aldosterone excess can be caused by somatic (intratumor) mutations in the potassium channel GIRK4 (encoded by <i>KCNJ5</i> ; identified as cause of disease in 40% of aldosterone-producing adenomas; rare germline mutations can cause bilateral macronodular adrenal hyperplasia). Further causes include somatic mutations affecting the α -subunit of the Na ⁺ /K ⁺ -ATPase (encoded by <i>ATP1A1</i>), the plasma membrane calcium-transporting ATPase 3 (encoded by <i>ATP2B3</i>), and somatic or germline mutations in <i>CACNA1D</i> encoding the voltage-gated calcium channel Cav1.3. All mutations result in upregulation of CYP11B2 and hence aldosterone synthesis.	60
Bilateral (micronodular) adrenal hyperplasia	Autonomous aldosterone excess	60
Glucocorticoid-remediable hyperaldosteronism (dexamethasone-suppressible hyperaldosteronism)	Crossover between the <i>CYP11B1</i> and <i>CYP11B2</i> genes results in ACTH-driven aldosterone production	<1
Other Causes (Rare)		
Syndrome of apparent mineralocorticoid excess (SAME)	Mutations in <i>HSD11B2</i> result in lack of renal inactivation of cortisol to cortisone, leading to excess activation of the MR by cortisol	
Cushing's syndrome	Cortisol excess overcomes the capacity of HSD11B2 to inactivate cortisol to cortisone, consequently flooding the MR	
Glucocorticoid resistance	Upregulation of cortisol production due to GR mutations results in flooding of the MR by cortisol	
Adrenocortical carcinoma	Autonomous aldosterone and/or DOC excess	
Congenital adrenal hyperplasia	Accumulation of DOC due to mutations in <i>CYP11B1</i> or <i>CYP17A1</i>	
Progesterone-induced hypertension	Progesterone acts as an abnormal ligand due to mutations in the MR gene	
Liddle's syndrome	Mutant ENaC β or γ subunits resulting in reduced degradation of ENaC keeping the membrane channel in open conformation for longer, enhancing mineralocorticoid action	

Abbreviations: ACTH, adrenocorticotropic hormone; DOC, deoxycorticosterone; ENaC, epithelial sodium channel; GR, glucocorticoid receptor; HSD11B2, 11 β -hydroxysteroid dehydrogenase type 2; MR, mineralocorticoid receptor.

from cross-over of promoter sequences between the *CYP11B1* and *CYP11B2* genes that are involved in glucocorticoid and mineralocorticoid synthesis, respectively (Fig. 406-1). This rearrangement brings *CYP11B2* transcription under the control of ACTH receptor signaling; consequently, aldosterone production is regulated by ACTH rather than by renin. The family history can be helpful because there may be evidence for dominant transmission of hypertension. Recognition of the disorder is important because it can be associated with early-onset hypertension and strokes. In addition, glucocorticoid suppression can reduce aldosterone production.

Other rare causes of mineralocorticoid excess are listed in Table 406-3. An important cause is excess binding and activation of the mineralocorticoid receptor by a steroid other than aldosterone. Cortisol acts as a potent mineralocorticoid if it escapes efficient inactivation to cortisone by 11 β -HSD2 in the kidney (Fig. 406-7). This can be caused by inactivating mutations in the *HSD11B2* gene resulting in the syndrome of apparent mineralocorticoid excess (SAME) that characteristically manifests with severe hypokalemic hypertension in childhood. However, milder mutations may cause normokalemic hypertension manifesting in adulthood (type II SAME). Inhibition of 11 β -HSD2 by excess licorice ingestion also results in hypokalemic hypertension, as does overwhelming of 11 β -HSD2 conversion capacity by cortisol excess in Cushing's syndrome. Deoxycorticosterone (DOC) also binds and activates the mineralocorticoid receptor and can cause hypertension if its circulating concentrations are increased. This can arise through autonomous DOC secretion by an adrenocortical carcinoma, but also when DOC accumulates as a consequence of an adrenal enzymatic block, as seen in congenital adrenal hyperplasia due to *CYP11B1* (11 β -hydroxylase) or *CYP17A1* (17 α -hydroxylase) deficiency (Fig. 406-1). Progesterone can cause hypokalemic hypertension in rare individuals who harbor a mineralocorticoid receptor mutation that enhances binding and activation by progesterone; physiologically, progesterone normally exerts antimineralocorticoid activity. Finally, excess mineralocorticoid activity can be caused by mutations in the β or γ subunits of the ENaC, disrupting its interaction with Nedd4 (Fig. 406-7), and thereby decreasing receptor internalization and degradation. The constitutively active ENaC drives hypokalemic hypertension, resulting in an autosomal dominant disorder termed *Liddle's syndrome*.

Clinical Manifestations Excess activation of the mineralocorticoid receptor leads to potassium depletion and increased sodium retention, with the latter causing an expansion of extracellular and plasma volume. Increased ENaC activity also results in hydrogen depletion that can cause metabolic alkalosis. Aldosterone also has direct effects on the vascular system, where it increases cardiac remodeling and decreases compliance. Aldosterone excess may cause direct damage to the myocardium and the kidney glomeruli, in addition to secondary damage due to systemic hypertension.

The clinical hallmark of mineralocorticoid excess is hypokalemic hypertension; serum sodium tends to be normal due to the concurrent fluid retention, which in some cases can lead to peripheral edema. Hypokalemia can be exacerbated by thiazide drug treatment, which leads to increased delivery of sodium to the distal renal tubule, thereby driving potassium excretion. Severe hypokalemia can be associated with muscle weakness, overt proximal myopathy, or even hypokalemic paralysis. Severe alkalosis contributes to muscle cramps and, in severe cases, can cause tetany.

Diagnosis Diagnostic screening for mineralocorticoid excess is not currently recommended for all patients with hypertension, but should be restricted to those who exhibit hypertension associated with drug resistance, hypokalemia, an adrenal mass, or onset of disease before the age of 40 years (Fig. 406-12). The accepted screening test is concurrent measurement of plasma renin and aldosterone with subsequent calculation of the aldosterone-renin ratio (ARR) (Fig. 406-12); serum potassium needs to be normalized prior to testing. Stopping antihypertensive medication can be cumbersome, particularly in patients with severe hypertension. Thus, for practical purposes, in the first instance the patient can remain on the usual antihypertensive medications, with the exception that mineralocorticoid receptor antagonists need to be ceased at least

4 weeks prior to ARR measurement. The remaining antihypertensive drugs usually do not affect the outcome of ARR testing, except that beta blocker treatment can cause false-positive results and ACE/AT1R inhibitors can cause false-negative results in milder cases (Table 406-4).

ARR screening is positive if the ratio is >750 pmol/L per ng/mL per hour, with a concurrently high normal or increased aldosterone (Fig. 406-12). If one relies on the ARR only, the likelihood of a false-positive ARR becomes greater when renin levels are very low. The characteristics of the biochemical assays are also important. Some labs measure plasma renin activity, whereas others measure plasma renin concentrations. Antibody-based assays for the measurement of serum aldosterone lack the reliability of tandem mass spectrometry assays, but these are not yet ubiquitously available.

Diagnostic confirmation of mineralocorticoid excess in a patient with positive ARR screening result should be undertaken by an endocrinologist as the tests lack optimized validation. The most straightforward is the saline infusion test, which involves the IV administration of 2 L of physiologic saline over a 4-h period. Failure of aldosterone to suppress below 140 pmol/L (5 ng/dL) is indicative of autonomous mineralocorticoid excess. Alternative tests are the oral sodium loading test (300 mmol NaCl/d for 3 days) or the fludrocortisone suppression test (0.1 mg q6h with 30 mmol NaCl q8h for 4 days); the latter can be difficult because of the risk of profound hypokalemia and increased hypertension. In patients with overt hypokalemic hypertension, strongly positive ARR, and concurrently increased aldosterone levels, confirmatory testing is usually not necessary.

Differential Diagnosis and Treatment After the diagnosis of hyperaldosteronism is established, the next step is to use adrenal imaging to further assess the cause. Fine-cut CT scanning of the adrenal region is the method of choice because it provides excellent visualization of adrenal morphology. CT will readily identify larger tumors suspicious of malignancy but may miss lesions smaller than 5 mm. The differentiation between bilateral micronodular hyperplasia and a unilateral adenoma is only required if a surgical approach is feasible and desired. Consequently, selective adrenal vein sampling (AVS) should only be carried out in surgical candidates with either no obvious lesion on CT or evidence of a unilateral lesion in patients older than 40 years, because the latter patients have a high likelihood of harboring a coincidental, endocrine-inactive adrenal adenoma (Fig. 406-12). AVS is used to compare aldosterone levels in the inferior vena cava and between the right and left adrenal veins. AVS requires concurrent measurement of cortisol to document correct placement of the catheter in the adrenal veins and should demonstrate a cortisol gradient >3 between the vena cava and each adrenal vein. Lateralization is confirmed by an aldosterone/cortisol ratio that is at least twofold higher on one side than the other. AVS is a complex procedure that requires a highly skilled interventional radiologist. Even then, the right adrenal vein can be difficult to cannulate correctly, which, if not achieved, invalidates the procedure. There is also no agreement as to whether the two adrenal veins should be cannulated simultaneously or successively and whether ACTH stimulation enhances the diagnostic value of AVS.

Patients younger than 40 years with confirmed mineralocorticoid excess and a unilateral lesion on CT can go straight to surgery, which is also indicated in patients with confirmed lateralization documented by a valid AVS procedure. Laparoscopic adrenalectomy is the preferred approach. Patients who are not surgical candidates, or with evidence of bilateral hyperplasia based on CT or AVS, should be treated medically (Fig. 406-12). Medical treatment, which can also be considered prior to surgery to avoid postsurgical hypoaldosteronism, consists primarily of the mineralocorticoid receptor antagonist spironolactone. It can be started at 12.5–50 mg bid and titrated up to a maximum of 400 mg/d to control blood pressure and normalize potassium. Side effects include menstrual irregularity, decreased libido, and gynecomastia. The more selective MR antagonist eplerenone can also be used. Doses start at 25 mg bid, and it can be titrated up to 200 mg/d. Another useful drug is the sodium channel blocker amiloride (5–10 mg bid).

In patients with normal adrenal morphology and family history of early-onset, severe hypertension, a diagnosis of GRA should be

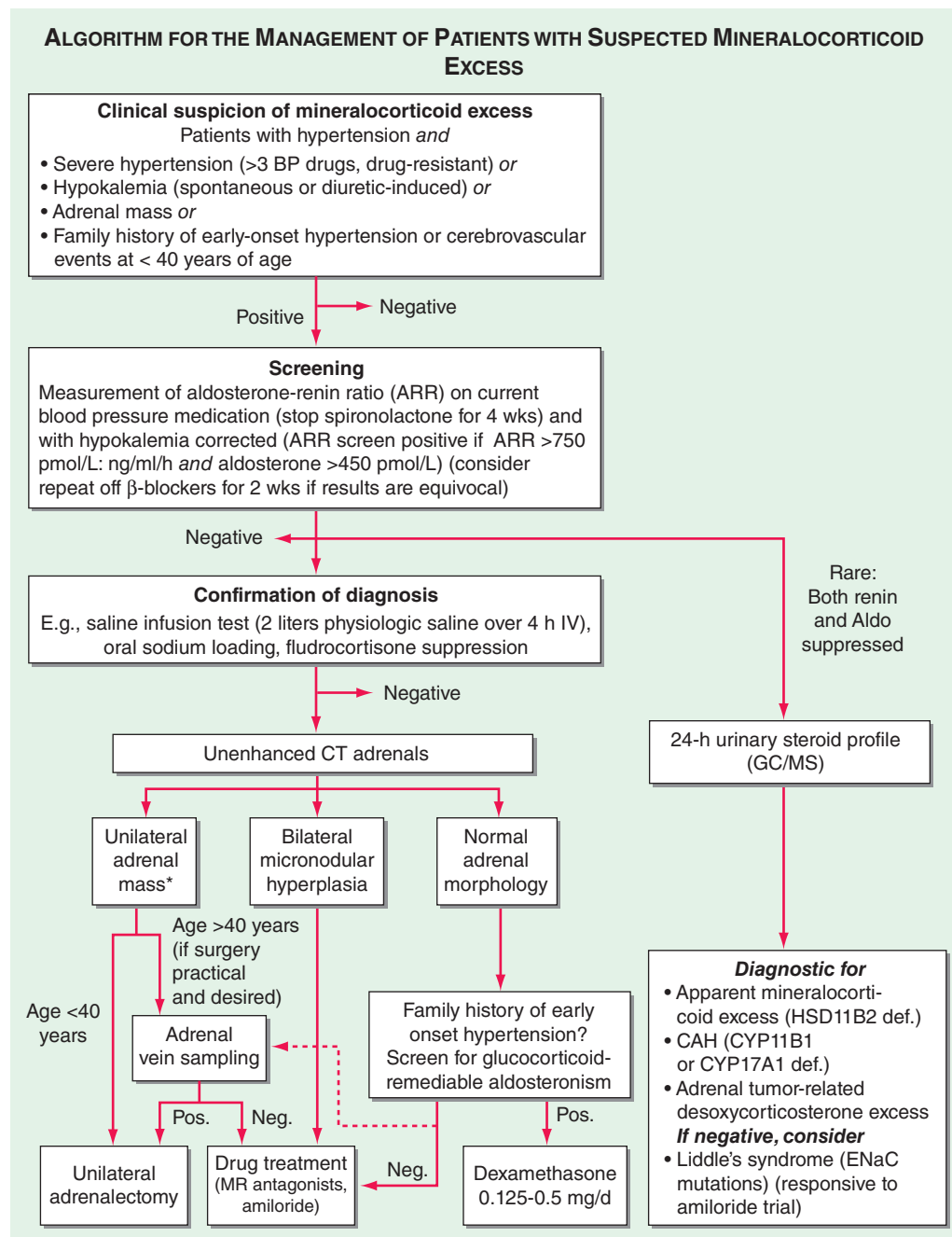


FIGURE 406-12 Management of patients with suspected mineralocorticoid excess. *Perform adrenal tumor workup (see Fig. 406-13). BP, blood pressure; CAH, congenital adrenal hyperplasia; CT, computed tomography; GC/MS, gas chromatography/mass spectrometry; PRA, plasma renin activity.

TABLE 406-4 EFFECTS OF ANTIHYPERTENSIVE DRUGS ON THE ALDOSTERONE-RENIN-RATIO (ARR)

Drug	Effect on Renin	Effect on Aldosterone	Net Effect on ARR
β Blockers	↓	↑	↑
α_1 Blockers	→	→	→
α_2 Sympathomimetics	→	→	→
ACE inhibitors	↑	↓	↓
AT1R blockers	↑	↓	↓
Calcium antagonists	→	→	→
Diuretics	(↑)	(↑)	→/(↓)

Abbreviations: ACE, angiotensin-converting enzyme; AT1R, angiotensin II receptor type 1.

considered and can be evaluated using genetic testing. Treatment of GRA consists of administering dexamethasone, using the lowest dose possible to control blood pressure. Some patients also require additional MR antagonist treatment.

The diagnosis of nonaldosterone-related mineralocorticoid excess is based on documentation of suppressed renin and suppressed aldosterone in the presence of hypokalemic hypertension. This testing is best carried out by employing urinary steroid metabolite profiling by gas chromatography/mass spectrometry (GC/MS). An increased free cortisol over free cortisone ratio is suggestive of SAME and can be treated with dexamethasone. Steroid profiling by GC/MS also detects the steroids associated with CYP11B1 and CYP17A1 deficiency or the irregular steroid secretion pattern in a DOC-producing adrenocortical carcinoma (Fig. 406-12). If the GC/MS profile is normal, then Liddle's syndrome should be considered. It is very sensitive to amiloride treatment but will not respond to MR antagonist treatment, because the defect is due to a constitutively active ENaC.

APPROACH TO THE PATIENT: INCIDENTALLY DISCOVERED ADRENAL MASS

Epidemiology Incidentally discovered adrenal masses, commonly termed adrenal “incidentalomas,” are common, with a prevalence of at least 2% in the general population as documented in CT and autopsy series. The prevalence increases with age, with 1% of 40-year-olds and 7% of 70-year-olds harboring an adrenal mass.

Etiology Most solitary adrenal tumors are monoclonal neoplasms. Several genetic syndromes, including MEN 1 (*MEN1*), MEN 2 (*RET*), Carney’s complex (*PRKARIA*), and McCune-Albright (*GNAS1*), can have adrenal tumors as one of their features. Somatic mutations in *MEN1*, *GNAS1*, and *PRKARIA* have been identified in a small proportion of sporadic adrenocortical adenomas. Aberrant expression of membrane receptors (gastric inhibitory peptide, α - and β -adrenergic, luteinizing hormone, vasopressin V1, and interleukin 1 receptors) have been identified in some sporadic cases of macronodular adrenocortical hyperplasia.

The majority of adrenal nodules are endocrine-inactive adrenocortical adenomas. However, larger series suggest that up to 25% of adrenal nodules are hormonally active, due to a cortisol- or aldosterone-producing adrenocortical adenoma or a pheochromocytoma associated with catecholamine excess (Table 406-5). Adrenocortical carcinoma is rare but is the cause of an adrenal mass in 5% of patients. However, the most common cause of a malignant adrenal mass is metastasis originating from another solid tissue tumor (Table 406-5).

Differential Diagnosis and Treatment Patients with an adrenal mass >1 cm require a diagnostic evaluation. Two key questions need to be addressed: (1) Does the tumor autonomously secrete hormones that could have a detrimental effect on health? (2) Is the adrenal mass benign or malignant?

Hormone secretion by an adrenal mass occurs along a continuum, with a gradual increase in clinical manifestations in parallel with hormone levels. Exclusion of catecholamine excess from a pheochromocytoma arising from the adrenal medulla is a mandatory part of the diagnostic workup (Fig. 406-13). Furthermore, autonomous cortisol and aldosterone secretion resulting in Cushing’s syndrome or primary

aldosteronism, respectively, require exclusion. Adrenal incidentalomas can be associated with lower levels of autonomous cortisol secretion, and patients may lack overt clinical features of Cushing’s syndrome. Nonetheless, they may exhibit one or more components of the metabolic syndrome (e.g., obesity, type 2 diabetes, or hypertension). There is ongoing debate about the optimal treatment for these patients with mild or subclinical Cushing’s syndrome. Overproduction of adrenal androgen precursors, DHEA and its sulfate, is rare and most frequently seen in the context of adrenocortical carcinoma, as are increased levels of steroid precursors such as 17-hydroxyprogesterone.

For the differentiation of benign from malignant adrenal masses, imaging is relatively sensitive, although specificity is suboptimal. CT is the procedure of choice for imaging the adrenal glands (Fig. 406-11). The risk of adrenocortical carcinoma, pheochromocytoma, and benign adrenal myelolipoma increases with the diameter of the adrenal mass. However, size alone is of poor predictive value, with only 80% sensitivity and 60% specificity for the differentiation of benign from malignant masses when using a 4-cm cut-off. Metastases are found with similar frequency in adrenal masses of all sizes. Tumor density on unenhanced CT is of additional diagnostic value, with most adrenocortical adenomas being lipid rich and thus presenting with low attenuation values (i.e., densities of <10 HU). By contrast, adrenocortical carcinomas, but also pheochromocytomas, usually have high attenuation values (i.e., densities >20 HU on precontrast scans). Generally, benign lesions are rounded and homogenous, whereas most malignant lesions appear lobulated and inhomogeneous. Pheochromocytoma and adrenomyelolipoma may also exhibit lobulated and inhomogeneous features. Additional information can be obtained from CT by assessment of contrast wash-out after 15 min, which is >50% in benign lesions but <40% in malignant lesions, which usually have a more extensive vascularization. MRI also allows for the visualization of the adrenal glands with somewhat lower resolution than CT. However, because it does not involve exposure to ionizing radiation, it is preferred in children, young adults, and during pregnancy. MRI has a valuable role in the characterization of indeterminate adrenal lesions using chemical shift analysis, with malignant tumors rarely showing loss of signal on opposed-phase MRI.

Fine-needle aspiration (FNA) or CT-guided biopsy of an adrenal mass is almost never indicated. FNA of a pheochromocytoma can cause a life-threatening hypertensive crisis. FNA of an adrenocortical carcinoma violates the tumor capsule and can cause needle track metastasis. FNA should only be considered in a patient with a history of nonadrenal malignancy and a newly detected adrenal mass, after careful exclusion of pheochromocytoma, and if the outcome will influence therapeutic management. It is important to recognize that in 25% of patients with a previous history of nonadrenal malignancy, a newly detected mass on CT is not a metastasis.

Adrenal masses associated with confirmed hormone excess or suspected malignancy are usually treated surgically (Fig. 406-13) or, if adrenalectomy is not feasible or desired, with medication. Preoperative exclusion of glucocorticoid excess is particularly important for the prediction of postoperative suppression of the contralateral adrenal gland, which requires glucocorticoid replacement peri- and postoperatively. If the initial decision is for observation, imaging and biochemical testing should be repeated about a year after the first assessment. However, this may be performed earlier in patients with borderline imaging or hormonal findings. There is no agreement with regard to the required long-term follow-up beyond 1 year in patients with normal biochemistry and no evidence of increased tumor size at follow-up.

ADRENOCORTICAL CARCINOMA

Adrenocortical carcinoma (ACC) is a rare malignancy with an annual incidence of 1–2 per million population. ACC is generally considered a highly malignant tumor; however, it presents with broad interindividual variability with regard to biologic characteristics and clinical behavior. Somatic mutations in the tumor-suppressor gene *TP53* are found in 25% of apparently sporadic ACC. Germline *TP53* mutations are the cause of the Li-Fraumeni syndrome associated with multiple solid organ cancers including ACC and are found in 25% of pediatric

TABLE 406-5 CLASSIFICATION OF UNILATERAL ADRENAL MASSES

Mass	Approximate Prevalence (%)
Benign	
Adrenocortical adenoma	
Endocrine-inactive	60–85
Cortisol-producing	5–10
Aldosterone-producing	2–5
Pheochromocytoma	5–10
Adrenal myelolipoma	<1
Adrenal ganglioneuroma	<0.1
Adrenal hemangioma	<0.1
Adrenal cyst	<1
Adrenal hematoma/hemorrhagic infarction	<1
Indeterminate	
Adrenocortical oncocytoma	<1
Malignant	
Adrenocortical carcinoma	2–5
Malignant pheochromocytoma	<1
Adrenal neuroblastoma	<0.1
Lymphomas (including primary adrenal lymphoma)	<1
Metastases (most frequent: breast, lung)	15

Note: Bilateral adrenal enlargement/masses may be caused by congenital adrenal hyperplasia, bilateral macronodular hyperplasia, bilateral hemorrhage (due to antiphospholipid syndrome or sepsis-associated Waterhouse-Friderichsen syndrome), granuloma, amyloidosis, or infiltrative disease including tuberculosis.

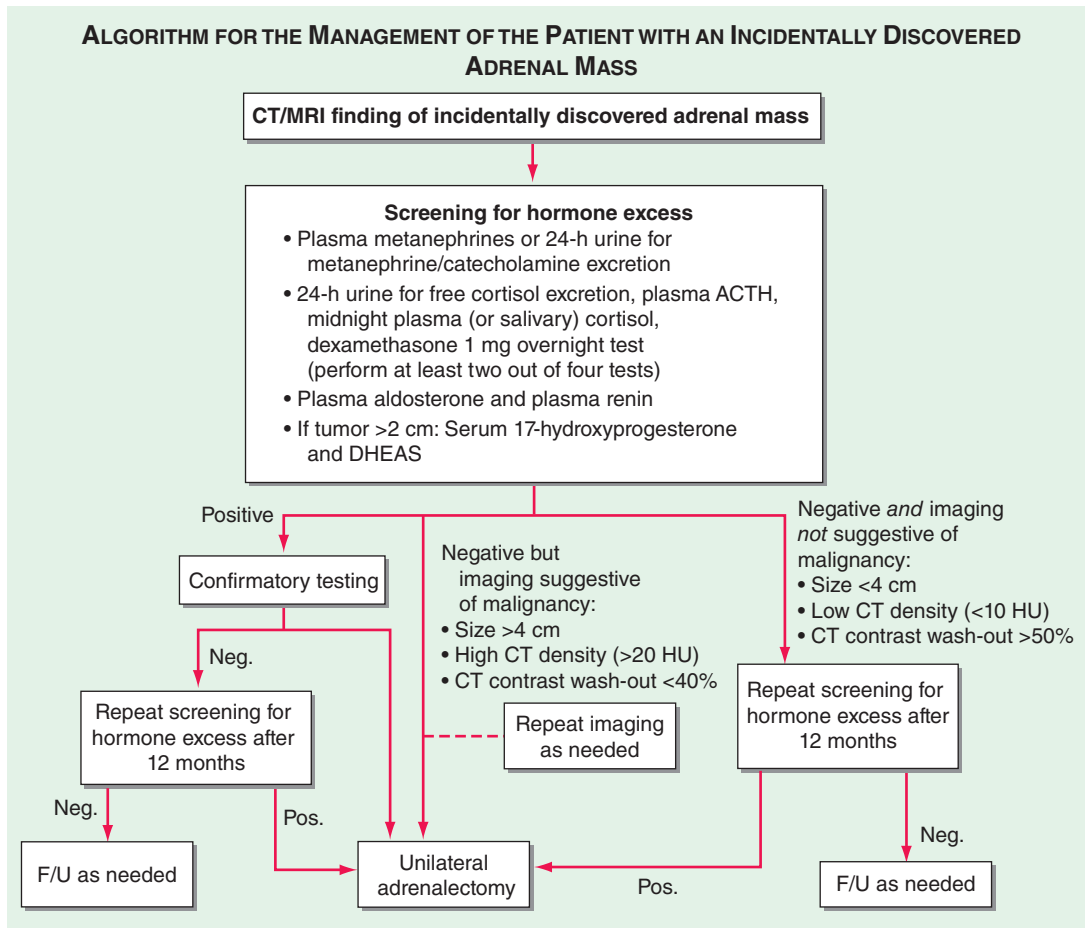


FIGURE 406-13 Management of the patient with an incidentally discovered adrenal mass. CT, computed tomography; F/U, follow-up; MRI, magnetic resonance imaging.

ACC cases; the *TP53* mutation R337H is found in almost all pediatric ACC in Brazil. Other genetic changes identified in ACC include alterations in the Wnt/ β -catenin pathway and in the insulin-like growth factor 2 (*IGF2*) cluster; *IGF2* overexpression is found in 90% of ACC.

Patients with large adrenal tumors suspicious of malignancy should be managed by a multidisciplinary specialist team, including an endocrinologist, an oncologist, a surgeon, a radiologist, and a histopathologist. FNA is not indicated in suspected ACC: first, cytology and also histopathology of a core biopsy cannot differentiate between benign and malignant primary adrenal masses; second, FNA violates the tumor capsule and may even cause needle canal metastasis. Even when the entire tumor specimen is available, the histopathologic differentiation between benign and malignant lesions is a diagnostic challenge. The most common histopathologic classification is the Weiss score, taking into account high nuclear grade; mitotic rate (>5/HPF); atypical mitosis; <25% clear cells; diffuse architecture; and presence of necrosis, venous invasion, and invasion of sinusoidal structures and tumor capsule. The presence of three or more elements suggests ACC.

Although 60–70% of ACCs show biochemical evidence of steroid overproduction, in many patients, this is not clinically apparent due to the relatively inefficient steroid production by the adrenocortical cancer cells. Excess production of glucocorticoids and adrenal androgen precursors are most common. Mixed excess production of several corticosteroid classes by an adrenal tumor is generally indicative of malignancy.

Tumor staging at diagnosis (Table 406-6) has important prognostic implications and requires scanning of the chest and abdomen for local organ invasion, lymphadenopathy, and metastases. Intravenous contrast medium is necessary for maximum sensitivity for hepatic metastases. An adrenal origin may be difficult to determine on standard axial CT imaging if the tumors are large and invasive, but CT reconstructions and MRI are more informative (Fig. 406-14) using multiple planes and different sequences. Vascular and adjacent organ

invasion is diagnostic of malignancy. 18-Fluoro-2-deoxy-D-glucose positron emission tomography (18-FDG PET) is highly sensitive for the detection of malignancy and can be used to detect small metastases or local recurrence that may not be obvious on CT (Fig. 406-14). However, FDG PET is not specific and therefore cannot be used for differentiating benign from malignant adrenal lesions. Metastasis in ACC most frequently occurs to liver and lung.

There is no established grading system for ACC, and the Weiss score carries no prognostic value; the most important prognostic histopathologic parameter is the Ki67 proliferation index, with Ki67 <10% indicative of slow to moderate growth velocity, whereas a Ki67 \geq 10% is associated with poor prognosis including high risk of recurrence and rapid progression.

Cure of ACC can only be achieved by early detection and complete surgical removal. Capsule violation during primary surgery, metastasis at diagnosis, and primary treatment in a nonspecialist center are major determinants of poor survival. If the primary tumor invades adjacent organs, en bloc removal of kidney and spleen should be considered to reduce the risk of recurrence. Surgery can also be considered in a patient with metastases if there is severe tumor-related hormone excess. This indication needs to be carefully weighed against surgical risk, including thromboembolic complications, and the resulting delay in the introduction of other therapeutic options. Patients with confirmed ACC and successful removal of the primary tumor should receive adjuvant treatment with mitotane (o,p'DDD), particularly in patients with a high risk of recurrence as determined by tumor size >8 cm, histopathologic signs of vascular invasion, capsule invasion or violation, and a Ki67 proliferation index \geq 10%. Adjuvant mitotane should be continued for at least 2 years, if the patient can tolerate side effects. Regular monitoring of plasma mitotane levels is mandatory (therapeutic range 14–20 mg/L; neurotoxic complications more frequent at >20 mg/L). Mitotane is usually started at 500 mg tid, with stepwise increases to a maximum dose of 2000 mg tid

TABLE 406-6 CLASSIFICATION SYSTEM FOR STAGING OF ADRENOCORTICAL CARCINOMA

ENSAT Stage	TNM Stage	TNM Definitions
I	T1,N0,M0	T1, tumor ≤5 cm N0, no positive lymph node M0, no distant metastases
II	T2,N0,M0	T2, tumor >5 cm N0, no positive lymph node M0, no distant metastases
III	T1–T2,N1,M0 T3–T4,N0–N1,M0	N1, positive lymph node(s) M0, no distant metastases T3, tumor infiltration into surrounding tissue T4, tumor invasion into adjacent organs or venous tumor thrombus in vena cava or renal vein
IV	T1–T4,N0–N1,M1	M1, presence of distant metastases

Abbreviations: ENSAT, European Network for the Study of Adrenal Tumors; TNM, tumor, node, metastasis.

in days (high-dose saturation) or weeks (low-dose saturation) as tolerated. Once therapeutic range plasma mitotane levels are achieved, the dose can be tapered to maintenance doses mostly ranging from 1000 to 1500 mg tid. Mitotane treatment results in disruption of cortisol synthesis and thus requires glucocorticoid replacement; glucocorticoid replacement dose should be at least double of that usually used in adrenal insufficiency (i.e., 20 mg tid) because mitotane induces hepatic CYP3A4 activity resulting in rapid inactivation of glucocorticoids. Mitotane also increases circulating CBG, thereby decreasing the available free cortisol fraction. Single metastases can be addressed surgically or with radiofrequency ablation as appropriate. If the tumor recurs or progresses during mitotane treatment, chemotherapy should be considered; the established first-line chemotherapy regimen is the combination of cisplatin, etoposide, and doxorubicin plus continuing mitotane. Painful bone metastasis responds to irradiation. Overall survival in ACC is still poor, with 5-year survival rates of 30–40% and a median survival of 15 months in metastatic ACC.

ADRENAL INSUFFICIENCY

Epidemiology The prevalence of well-documented, permanent adrenal insufficiency is 5 in 10,000 in the general population. Hypothalamic-pituitary origin of disease is most frequent, with a prevalence of 3 in

10,000, whereas primary adrenal insufficiency has a prevalence of 2 in 10,000. Approximately one-half of the latter cases are acquired, mostly caused by autoimmune destruction of the adrenal glands; the other one-half are genetic, most commonly caused by distinct enzymatic blocks in adrenal steroidogenesis affecting glucocorticoid synthesis (i.e., congenital adrenal hyperplasia).

Adrenal insufficiency arising from suppression of the HPA axis as a consequence of exogenous glucocorticoid treatment is much more common, occurring in 0.5–2% of the population in developed countries.

Etiology *Primary adrenal insufficiency* is most commonly caused by autoimmune adrenalitis. Isolated autoimmune adrenalitis accounts for 30–40%, whereas 60–70% develop adrenal insufficiency as part of autoimmune polyglandular syndromes (APS) (Chap. 408) (Table 406-7). APS1, also termed APECED (autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy), is the underlying cause in 10% of patients affected by APS. APS1 is transmitted in an autosomal recessive manner and is caused by mutations in the autoimmune regulator gene *AIRE*. Associated autoimmune conditions overlap with those seen in APS2, but may also include total alopecia, primary hypoparathyroidism, and, in rare cases, lymphoma. APS1 patients invariably develop chronic mucocutaneous candidiasis, usually manifest in childhood, and preceding adrenal insufficiency by years or decades. The much more prevalent APS2 is of polygenic inheritance, with confirmed associations with the *HLA-DR3* gene region in the major histocompatibility complex and distinct gene regions involved in immune regulation (*CTLA-4*, *PTPN22*, *CLEC16A*). Coincident autoimmune disease most frequently includes thyroid autoimmune disease, vitiligo, and premature ovarian failure. Less commonly, additional features may include type 1 diabetes and pernicious anemia caused by vitamin B₁₂ deficiency.

X-linked adrenoleukodystrophy has an incidence of 1:20,000 males and is caused by mutations in the *X-ALD* gene encoding the peroxisomal membrane transporter protein ABCD1; its disruption results in accumulation of very long chain (>24 carbon atoms) fatty acids. Approximately 50% of cases manifest in early childhood with rapidly progressive white matter disease (cerebral ALD); 35% present during adolescence or in early adulthood with neurologic features indicative of myelin and peripheral nervous system involvement (adrenomyeloneuropathy [AMN]). In the remaining 15%, adrenal insufficiency is the sole manifestation of disease. Of note, distinct mutations manifest with variable penetrance and phenotypes within affected families.

Rarer causes of adrenal insufficiency involve destruction of the adrenal glands as a consequence of infection, hemorrhage, or infiltration

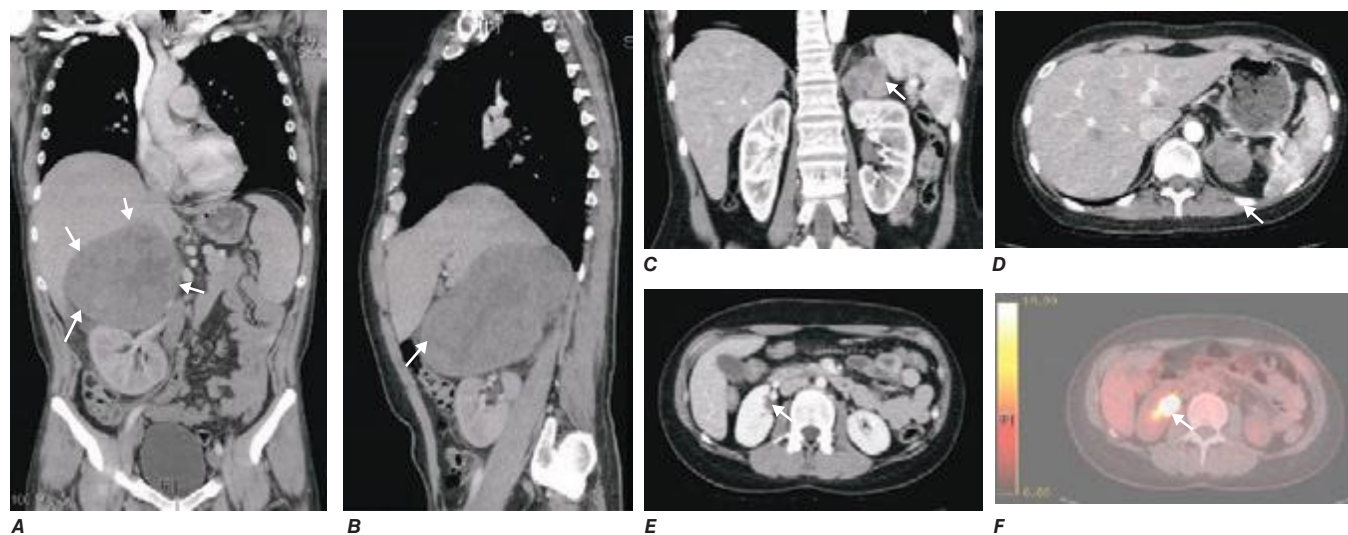


FIGURE 406-14 Imaging in adrenocortical carcinoma. Magnetic resonance imaging scan with (A) frontal and (B) lateral views of a right adrenocortical carcinoma that was detected incidentally. Computed tomography (CT) scan with (C) coronal and (D) transverse views depicting a right-sided adrenocortical carcinoma. Note the irregular border and inhomogeneous structure. CT scan (E) and positron emission tomography/CT (F) visualizing a peritoneal metastasis of an adrenocortical carcinoma in close proximity to the right kidney (arrow).

TABLE 406-7 CAUSES OF PRIMARY ADRENAL INSUFFICIENCY

Diagnosis	Gene	Associated Features
Autoimmune polyglandular syndrome 1 (APS1)	<i>AIRE</i>	Hypoparathyroidism, chronic mucocutaneous candidiasis, other autoimmune disorders, rarely lymphomas
Autoimmune polyglandular syndrome 2 (APS2)	Associations with HLA-DR3, CTLA-4	Hypothyroidism, hyperthyroidism, premature ovarian failure, vitiligo, type 1 diabetes mellitus, pernicious anemia
Isolated autoimmune adrenalitis	Associations with HLA-DR3, CTLA-4	
Congenital adrenal hyperplasia (CAH)	<i>CYP21A2</i> , <i>CYP11B1</i> , <i>CYP17A1</i> , <i>HSD3B2</i> , <i>POR</i>	See Table 406-10 (see also Chap. 410)
Congenital lipoid adrenal hyperplasia (CLAH)	<i>STAR</i> , <i>CYP11A1</i>	46,XY DSD, gonadal failure (see also Chap. 410)
Adrenal hypoplasia congenita (AHC)	<i>NROB1 (DAX-1)</i> , <i>NR5A1 (SF-1)</i>	46,XY DSD, gonadal failure (see also Chap. 410)
Adrenoleukodystrophy (ALD), adrenomyeloneuropathy (AMN)	<i>X-ALD</i>	Demyelination of central nervous system (ALD) or spinal cord and peripheral nerves (AMN)
Familial glucocorticoid deficiency	<i>MC2R</i>	Tall stature
	<i>MRAP</i>	None
	<i>STAR</i>	None
	<i>NNT</i>	None
Triple A syndrome	<i>MCM4</i>	Growth retardation, natural killer cell deficiency
	<i>AAAS</i>	Alacrima, achalasia, neurologic impairment
Smith-Lemli-Opitz syndrome	<i>SLOS</i>	Cholesterol synthesis disorder associated with mental retardation, craniofacial malformations, growth failure
Kearns-Sayre syndrome	Mitochondrial DNA deletions	Progressive external ophthalmoplegia, pigmentary retinal degeneration, cardiac conduction defects, gonadal failure, hypoparathyroidism, type 1 diabetes,
IMAge syndrome	<i>CDKN1C</i>	Intrauterine growth retardation, metaphyseal dysplasia, genital anomalies
Adrenal infections		Tuberculosis, HIV, CMV, cryptococcosis, histoplasmosis, coccidioidomycosis
Adrenal infiltration		Metastases, lymphomas, sarcoidosis, amyloidosis, hemochromatosis
Adrenal hemorrhage		Meningococcal sepsis (Waterhouse-Friderichsen syndrome), primary antiphospholipid syndrome
Drug-induced		Mitotane, aminoglutethimide, abiraterone, trilostane, etomidate, ketoconazole, suramin, RU486
Bilateral adrenalectomy		E.g., in the management of Cushing's or after bilateral nephrectomy

Abbreviations: AIRE, autoimmune regulator; CMV, cytomegalovirus; DSD, disordered sex development; MC2R, ACTH receptor; MCM4, mini chromosome maintenance-deficient 4 homologue; MRAP, MC2R-accessory protein; NNT, nicotinamide nucleotide transhydrogenase.

(Table 406-7); tuberculous adrenalitis is still a frequent cause of disease in developing countries. Adrenal metastases rarely cause adrenal insufficiency, and this occurs only with bilateral, bulky metastases.

Inborn causes of primary adrenal insufficiency other than congenital adrenal hyperplasia are rare, causing less than 1% of cases.

However, their elucidation provides important insights into adrenal gland development and physiology. Mutations causing primary adrenal insufficiency (Table 406-7) include factors regulating adrenal development and steroidogenesis (*DAX-1*, *SF-1*), cholesterol synthesis, import and cleavage (*DHCR7*, *StAR*, *CYP11A1*), and elements of the adrenal ACTH response pathway (*MC2R*, *MRAP*) (Fig. 406-5), and factors involved in redox regulation (*NNT*) and DNA repair (*MCM4*, *CDKN1C*).

Secondary adrenal insufficiency is the consequence of dysfunction of the hypothalamic-pituitary component of the HPA axis ([Table 406-8](#)). Excluding iatrogenic suppression, the overwhelming majority of cases are caused by pituitary or hypothalamic tumors or their treatment by surgery or irradiation ([Chap. 403](#)). Rarer causes include pituitary apoplexy, either as a consequence of an infarcted pituitary adenoma or transient reduction in the blood supply of the pituitary during surgery or after rapid blood loss associated with parturition, also termed Sheehan's syndrome. Isolated ACTH deficiency is rarely caused by autoimmune disease or pituitary infiltration (Table 406-8). Mutations in the ACTH precursor POMC or in factors regulating pituitary development are genetic causes of ACTH deficiency (Table 406-8).

Clinical Manifestations In principle, the clinical features of primary adrenal insufficiency (Addison's disease) are characterized by the loss of both glucocorticoid and mineralocorticoid secretion ([Table 406-9](#)). In secondary adrenal insufficiency, only glucocorticoid deficiency is present, as the adrenal itself is intact and thus still amenable to regulation by the RAA system. Adrenal androgen secretion is disrupted in both primary and secondary adrenal insufficiency (Table 406-9). Hypothalamic-pituitary disease can lead to additional clinical manifestations due to involvement of other endocrine axes (thyroid, gonads, growth hormone, prolactin) or visual impairment with bitemporal hemianopia caused by chiasmal compression. It is important to recognize that iatrogenic adrenal insufficiency caused by exogenous glucocorticoid suppression of the HPA axis may result in all symptoms associated with glucocorticoid deficiency (Table 406-9), if exogenous glucocorticoids are stopped abruptly. However, patients will appear clinically cushingoid as a result of the preceding overexposure to glucocorticoids.

Chronic adrenal insufficiency manifests with relatively nonspecific signs and symptoms such as fatigue and loss of energy, often resulting in delayed or missed diagnoses (e.g., as depression or anorexia). A distinguishing feature of primary adrenal insufficiency is hyperpigmentation, which is caused by excess ACTH stimulation of melanocytes. Hyperpigmentation is most pronounced in skin areas exposed to increased friction or shear stress and is increased by sunlight ([Fig. 406-15](#)). Conversely, in secondary adrenal insufficiency, the skin has an alabaster-like paleness due to lack of ACTH secretion.

Hyponatremia is a characteristic biochemical feature in primary adrenal insufficiency and is found in 80% of patients at presentation. Hyperkalemia is present in 40% of patients at initial diagnosis. Hyponatremia is primarily caused by mineralocorticoid deficiency but can also occur in secondary adrenal insufficiency due to diminished inhibition of antidiuretic hormone (ADH) release by cortisol, resulting in mild syndrome of inappropriate secretion of antidiuretic hormone (SIADH). Glucocorticoid deficiency also results in slightly increased TSH concentrations that normalize within days to weeks after initiation of glucocorticoid replacement.

Acute adrenal insufficiency usually occurs after a prolonged period of nonspecific complaints and is more frequently observed in patients with primary adrenal insufficiency, due to the loss of both glucocorticoid and mineralocorticoid secretion. Postural hypotension may progress to hypovolemic shock. Adrenal insufficiency may mimic features of acute abdomen with abdominal tenderness, nausea, vomiting, and fever. In some cases, the primary presentation may resemble neurologic disease, with decreased responsiveness, progressing to stupor and coma. An adrenal crisis can be triggered by an intercurrent illness, surgical or other stress, or increased glucocorticoid inactivation (e.g., hyperthyroidism).

TABLE 406-8 CAUSES OF SECONDARY ADRENAL INSUFFICIENCY

Diagnosis	Gene	Associated Features
Pituitary tumors (endocrine active and inactive adenomas, very rare: carcinoma)		Depending on tumor size and location: visual field impairment (bilateral hemianopia), hyperprolactinemia, secondary hypothyroidism, hypogonadism, growth hormone deficiency
Other mass lesions affecting the hypothalamic-pituitary region		Craniopharyngioma, meningioma, ependymoma, metastases
Pituitary irradiation		Radiotherapy administered for pituitary tumors, brain tumors, or craniospinal irradiation in leukemia
Autoimmune hypophysitis		Often associated with pregnancy; may present with panhypopituitarism or isolated ACTH deficiency; can be associated with autoimmune thyroid disease, more rarely with vitiligo, premature ovarian failure, type 1 diabetes, pernicious anemia
Pituitary apoplexy/hemorrhage		Hemorrhagic infarction of large pituitary adenomas or pituitary infarction consequent to traumatic major blood loss (e.g., surgery or pregnancy: Sheehan's syndrome)
Pituitary infiltration		Tuberculosis, actinomycosis, sarcoidosis, histiocytosis X, granulomatosis with polyangiitis (Wegener's), metastases
Drug-induced		Chronic glucocorticoid excess (endogenous or exogenous)
Congenital isolated ACTH deficiency	<i>TBX19</i> (Tpit)	
Combined pituitary hormone deficiency (CPHD)	<i>PROP-1</i>	Progressive development of CPHD in the order GH, PRL, TSH, LH/FSH, ACTH
	<i>HESX1</i>	CPHD and septo-optic dysplasia
	<i>LHX3</i>	CPHD and limited neck rotation, sensorineural deafness
	<i>LHX4</i>	CPHD and cerebellar abnormalities
Proopiomelanocortin (POMC) deficiency	<i>SOX3</i>	CPHD and variable mental retardation
	<i>POMC</i>	Early-onset obesity, red hair pigmentation

Abbreviations: ACTH, adrenocorticotropic hormone; GH, growth hormone; LH/FSH, luteinizing hormone/follicle-stimulating hormone; PRL, prolactin; TSH, thyroid-stimulating hormone.

Diagnosis The diagnosis of adrenal insufficiency is established by the short cosyntropin test, a safe and reliable tool with excellent predictive diagnostic value (Fig. 406-16). The cut-off for failure is usually defined at cortisol levels of <500–550 nmol/L (18–20 µg/dL) sampled 30–60 min after ACTH stimulation; the exact cut-off is dependent on the locally available assay. During the early phase of HPA disruption (e.g., within 4 weeks of pituitary insufficiency), patients may still respond to exogenous ACTH stimulation. In this circumstance, the ITT is an alternative choice but is more invasive and should be carried out only under a specialist's supervision (see above). Induction of hypoglycemia is contraindicated in individuals with diabetes mellitus, cardiovascular disease, or history of seizures. Random serum cortisol measurements are of limited diagnostic value, because baseline cortisol levels may be coincidentally low due to the physiologic diurnal rhythm of cortisol secretion (Fig. 406-3). Similarly, many patients with secondary adrenal insufficiency have relatively normal baseline cortisol levels but fail to mount an appropriate cortisol response to ACTH, which can only be revealed by stimulation testing. Importantly, tests to establish the diagnosis of adrenal insufficiency should never delay treatment. Thus, in a patient with suspected adrenal crisis, it is reasonable to draw baseline cortisol levels, provide replacement therapy, and defer formal stimulation testing until a later time.

TABLE 406-9 SIGNS AND SYMPTOMS OF ADRENAL INSUFFICIENCY

Signs and Symptoms Caused by Glucocorticoid Deficiency
TB>Fatigue, lack of energy
Weight loss, anorexia
Myalgia, joint pain
Fever
Normochromic anemia, lymphocytosis, eosinophilia
Slightly increased TSH (due to loss of feedback inhibition of TSH release)
Hypoglycemia (more frequent in children)
Low blood pressure, postural hypotension
Hyponatremia (due to loss of feedback inhibition of AVP release)
Signs and Symptoms Caused by Mineralocorticoid Deficiency (Primary Adrenal Insufficiency Only)
Abdominal pain, nausea, vomiting
Dizziness, postural hypotension
Salt craving
Low blood pressure, postural hypotension
Increased serum creatinine (due to volume depletion)
Hyponatremia
Hyperkalemia
Signs and Symptoms Caused by Adrenal Androgen Deficiency
Lack of energy
Dry and itchy skin (in women)
Loss of libido (in women)
Loss of axillary and pubic hair (in women)
Other Signs and Symptoms
Hyperpigmentation (primary adrenal insufficiency only) (due to excess of proopiomelanocortin [POMC]-derived peptides)
Alabaster-colored pale skin (secondary adrenal insufficiency only) (due to deficiency of POMC-derived peptides)

Abbreviations: AVP, arginine vasopressin; TSH, thyroid-stimulating hormone.

Once adrenal insufficiency is confirmed, measurement of plasma ACTH is the next step, with increased or inappropriately low levels defining primary and secondary origin of disease, respectively (Fig. 406-16). In primary adrenal insufficiency, increased plasma renin will confirm the presence of mineralocorticoid deficiency. At initial presentation, patients with primary adrenal insufficiency should undergo screening for steroid autoantibodies as a marker of autoimmune adrenalitis. If these tests are negative, adrenal imaging by CT is indicated to investigate possible hemorrhage, infiltration, or masses. In male patients with negative autoantibodies in the plasma, very-long-chain fatty acids should be measured to exclude X-ALD. Patients with inappropriately low ACTH, in the presence of confirmed cortisol deficiency, should undergo hypothalamic-pituitary imaging by MRI. Features suggestive of preceding pituitary apoplexy, such as sudden-onset severe headache or history of previous head trauma, should be carefully explored, particularly in patients with no obvious MRI lesion.

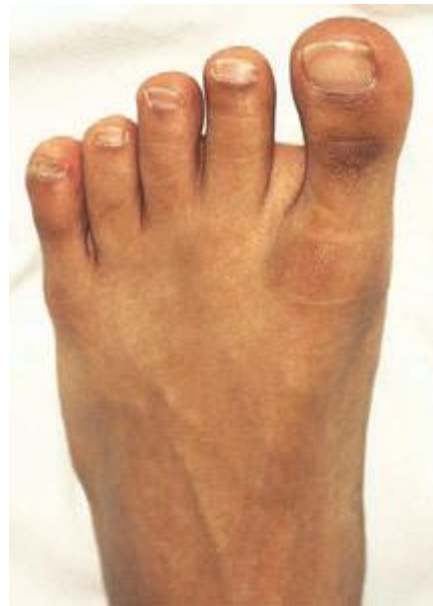
TREATMENT ACUTE ADRENAL INSUFFICIENCY

Acute adrenal insufficiency requires immediate initiation of rehydration, usually carried out by saline infusion at initial rates of 1 L/h with continuous cardiac monitoring. Glucocorticoid replacement should be initiated by bolus injection of 100 mg hydrocortisone, followed by the administration of 100–200 mg hydrocortisone over 24 h, either by continuous infusion or by bolus IV or IM injections. Mineralocorticoid replacement can be initiated once the daily hydrocortisone dose has been reduced to <50 mg because at higher doses hydrocortisone provides sufficient stimulation of mineralocorticoid receptors.

Glucocorticoid replacement for the treatment of chronic adrenal insufficiency should be administered at a dose that replaces the physiologic daily cortisol production, which is usually achieved by



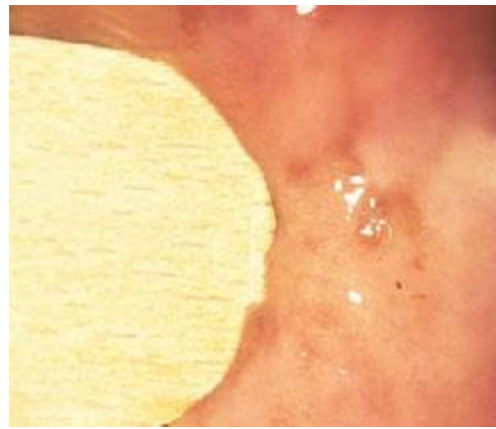
A



B



C



D

FIGURE 406-15 Clinical features of Addison's disease. Note the hyperpigmentation in areas of increased friction including (A) palmar creases, (B) dorsal foot, (C) nipples and axillary region, and (D) patchy hyperpigmentation of the oral mucosa.

the oral administration of 15–25 mg hydrocortisone in two to three divided doses. Pregnancy may require an increase in hydrocortisone dose by 50% during the last trimester. In all patients, at least one-half of the daily dose should be administered in the morning. Currently available glucocorticoid preparations fail to mimic the physiologic cortisol secretion rhythm (Fig. 406-3). Long-acting glucocorticoids such as prednisolone or dexamethasone are not preferred because they result in increased glucocorticoid exposure due to extended glucocorticoid receptor activation at times of physiologically low cortisol secretion. There are no well-established dose equivalencies, but as a guide, equipotency can be assumed for 1 mg hydrocortisone, 1.6 mg cortisone acetate, 0.2 mg prednisolone, 0.25 mg prednisone, and 0.025 mg dexamethasone.

Monitoring of glucocorticoid replacement is mainly based on the history and examination for signs and symptoms suggestive of glucocorticoid over- or underreplacement, including assessment of body weight and blood pressure. Plasma ACTH, 24-h urinary free cortisol, or serum cortisol day curves reflect whether hydrocortisone has been taken or not, but do not convey reliable information about replacement quality. In patients with isolated primary adrenal insufficiency, monitoring should include screening for autoimmune thyroid disease, and female patients should be made aware of the possibility of premature ovarian failure. Supraphysiologic glucocorticoid treatment with doses equivalent to 30 mg hydrocortisone or more will affect bone metabolism, and these patients should

undergo regular bone mineral density evaluation. All patients with adrenal insufficiency need to be instructed about the requirement for stress-related glucocorticoid dose adjustments. These generally consist of doubling the routine oral glucocorticoid dose in the case of intercurrent illness with fever and bed rest and the need for IV hydrocortisone injection at a daily dose of 100 mg in cases of prolonged vomiting, surgery, or trauma. Patients living or traveling in regions with delayed access to acute health care should carry a hydrocortisone self-injection emergency kit, in addition to their usual steroid emergency cards and bracelets.

Mineralocorticoid replacement in primary adrenal insufficiency should be initiated at a dose of 100–150 µg fludrocortisone. The adequacy of treatment can be evaluated by measuring blood pressure, sitting and standing, to detect a postural drop indicative of hypovolemia. In addition, serum sodium, potassium, and plasma renin should be measured regularly. Renin levels should be kept in the upper normal reference range. Changes in glucocorticoid dose may also impact on mineralocorticoid replacement as cortisol also binds the mineralocorticoid receptor; 40 mg hydrocortisone is equivalent to 100 µg fludrocortisone. In patients living or traveling in areas with hot or tropical weather conditions, the fludrocortisone dose should be increased by 50–100 µg during the summer. Mineralocorticoid dose may also need to be adjusted during pregnancy, due to the antiminerlocorticoid activity of progesterone, but this is less often required than hydrocortisone dose adjustment.

ALGORITHM FOR THE MANAGEMENT OF THE PATIENT WITH SUSPECTED ADRENAL INSUFFICIENCY

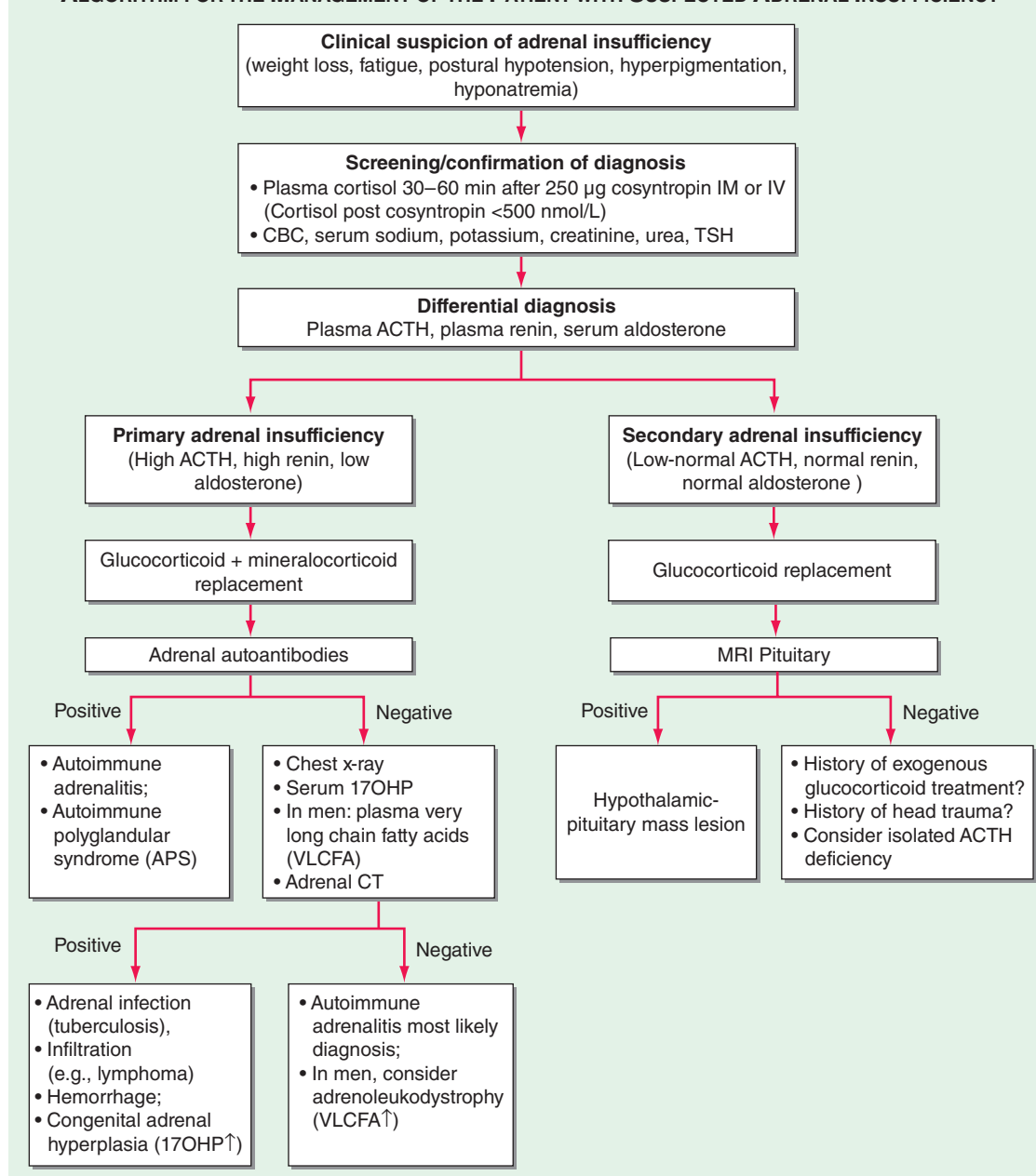


FIGURE 406-16 Management of the patient with suspected adrenal insufficiency. ACTH, adrenocorticotropic hormone; CBC, complete blood count; MRI, magnetic resonance imaging; PRA, plasma renin activity; TSH, thyroid-stimulating hormone.

Plasma renin cannot serve as a monitoring tool during pregnancy, because renin rises physiologically during gestation.

Adrenal androgen replacement is an option in patients with lack of energy, despite optimized glucocorticoid and mineralocorticoid replacement. It may also be indicated in women with features of androgen deficiency, including loss of libido. Adrenal androgen replacement can be achieved by once-daily administration of 25–50 mg DHEA. Treatment is monitored by measurement of DHEAS, androstenedione, testosterone, and sex hormone-binding globulin (SHBG) 24 h after the last DHEA dose.

CONGENITAL ADRENAL HYPERPLASIA

(See also Chap. 410) Congenital adrenal hyperplasia (CAH) is caused by mutations in genes encoding steroidogenic enzymes involved in glucocorticoid synthesis (*CYP21A2*, *CYP17A1*, *HSD3B2*, *CYP11B1*) or in the cofactor enzyme P450 oxidoreductase that serves as an electron donor to *CYP21A2* and *CYP17A1* (Fig. 406-1). Invariably, patients affected by CAH exhibit glucocorticoid deficiency. Depending on the

exact step of enzymatic block, they may also have excess production of mineralocorticoids or deficient production of sex steroids (Table 406-10). The diagnosis of CAH is readily established by measurement of the steroids accumulating before the distinct enzymatic block, either in serum or in urine, preferably by the use of mass spectrometry-based assays (Table 406-10).

Mutations in *CYP21A2* are the most prevalent cause of CAH, responsible for 90–95% of cases. 21-Hydroxylase deficiency disrupts glucocorticoid and mineralocorticoid synthesis (Fig. 406-1), resulting in diminished negative feedback via the HPA axis. This leads to increased pituitary ACTH release, which drives increased synthesis of adrenal androgen precursors and subsequent androgen excess. The degree of impairment of glucocorticoid and mineralocorticoid secretion depends on the severity of mutations. Major loss-of-function mutations result in combined glucocorticoid and mineralocorticoid deficiency (classic CAH, neonatal presentation), whereas less severe mutations affect glucocorticoid synthesis only (simple virilizing CAH, neonatal or early childhood presentation). The mildest mutations result in the least severe clinical phenotype, nonclassic CAH, usually

Variant	Gene	Impact on Steroid Synthesis	Diagnostic Marker Steroids in Serum (and Urine)
21-Hydroxylase deficiency (21OHD)	<i>CYP21A2</i>	Glucocorticoid deficiency, mineralocorticoid deficiency, adrenal androgen excess	17-Hydroxyprogesterone, 21-deoxycortisol (pregnanetriol, 17-hydroxypregnanolone, pregnanetriolone)
11 β -Hydroxylase deficiency (11OHD)	<i>CYP11B1</i>	Glucocorticoid deficiency, mineralocorticoid excess, adrenal androgen excess	11-Deoxycortisol, 11-deoxycorticosterone (tetrahydro-11-deoxycortisol, tetrahydro-11-deoxycorticosterone)
17 α -Hydroxylase deficiency (17OHD)	<i>CYP17A1</i>	(Glucocorticoid deficiency), mineralocorticoid excess, androgen deficiency	11-Deoxycorticosterone, corticosterone, pregnenolone, progesterone (tetrahydro-11-deoxycorticosterone, tetrahydrocorticosterone, pregnenediol, pregnanediol)
3 β -Hydroxysteroid dehydrogenase deficiency (3 β HSD)	<i>HSD3B2</i>	Glucocorticoid deficiency, (mineralocorticoid deficiency), adrenal androgen excess	17-Hydroxypregnanolone (pregnanetriol)
P450 oxidoreductase deficiency (ORD)	<i>POR</i>	Glucocorticoid deficiency, (mineralocorticoid excess), androgen deficiency, skeletal malformations	Pregnenolone, progesterone, 17-hydroxyprogesterone (pregnanediol, pregnanetriol)

presenting during adolescence and early adulthood and with preserved glucocorticoid production.

Androgen excess is present in all patients and manifests with broad phenotypic variability, ranging from severe virilization of the external genitalia in neonatal girls (e.g., 46,XX disordered sex development [DSD]) to hirsutism and oligomenorrhea resembling a polycystic ovary syndrome phenotype in young women with nonclassic CAH. In countries without neonatal screening for CAH, boys with classic CAH usually present with life-threatening adrenal crisis in the first few weeks of life (salt-wasting crisis); a simple-virilizing genotype manifests with precocious pseudo-puberty and advanced bone age in early childhood, whereas men with nonclassic CAH are usually detected only through family screening.

Glucocorticoid treatment is more complex than for other causes of primary adrenal insufficiency as it not only needed to replace missing glucocorticoids but also to control the increased ACTH drive and subsequent androgen excess. Current treatment is hampered by the lack of glucocorticoid preparations that mimic the diurnal cortisol secretion profile, resulting in a prolonged period of ACTH stimulation and subsequent androgen production during the early morning hours. In childhood,

optimization of growth and pubertal development are important goals of glucocorticoid treatment, in addition to prevention of adrenal crisis and treatment of 46,XX DSD. In adults, the focus shifts to preserving fertility and preventing side effects of glucocorticoid overtreatment, namely, the metabolic syndrome and osteoporosis. Fertility can be compromised in women due to oligomenorrhea/amenorrhea with chronic anovulation as a consequence of androgen excess. Men may develop so-called testicular adrenal rest tumors (Fig. 406-17). These consist of hyperplastic cells with adrenocortical characteristics located in the rete testis and should not be confused with testicular tumors. Testicular adrenal rest tissue can compromise sperm production and induce fibrosis that may be irreversible.

TREATMENT CONGENITAL ADRENAL HYPERPLASIA

Hydrocortisone is a good treatment option for the prevention of adrenal crisis, but longer acting prednisolone may be needed to control androgen excess. In children, hydrocortisone is given in divided doses at 1–1.5 times the normal cortisol production rate (about 10–13 mg/m² per day). In adults, if hydrocortisone does not suffice, intermediate-acting glucocorticoids (e.g., prednisone) may be

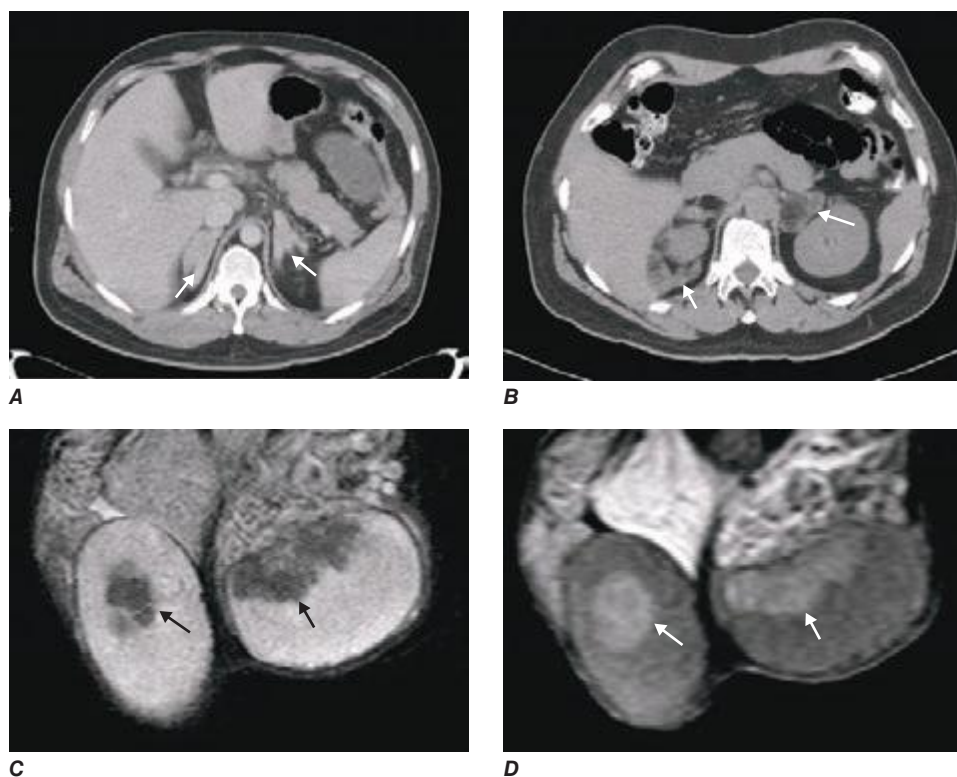


FIGURE 406-17 Imaging in congenital adrenal hyperplasia (CAH). Adrenal computed tomography scans showing homogenous bilateral hyperplasia in a young patient with classic CAH (A) and macronodular bilateral hyperplasia (B) in a middle-aged patient with classic CAH with longstanding poor disease control. Magnetic resonance imaging scan with T1-weighted (C) and T2-weighted (D) images showing bilateral testicular adrenal rest tumors (arrows) in a young patient with salt-wasting congenital adrenal hyperplasia. (Courtesy of N. Reisch.)

given, using the lowest dose necessary to suppress excess androgen production. For achieving fertility, dexamethasone treatment may be required, but should be only given for the shortest possible time period to limit adverse metabolic side effects. Biochemical monitoring should include androstenedione and testosterone, aiming for the normal sex-specific reference range. 17-Hydroxyprogesterone (17OHP) is a useful marker of overtreatment, indicated by 17OHP levels within the normal range of healthy controls. Glucocorticoid overtreatment may suppress the hypothalamic-pituitary-gonadal axis. Thus, treatment needs to be carefully titrated against clinical features of disease control. Stress dose glucocorticoids should be given at double or triple the daily dose for surgery, acute illness, or severe trauma. Poorly controlled CAH can result in adrenocortical hyperplasia, which gave the disease its name, and may present as macronodular hyperplasia subsequent to long-standing ACTH excess (Fig. 406-17). The nodular areas can develop autonomous adrenal androgen production and may be unresponsive to glucocorticoid treatment.

Mineralocorticoid requirements change during life and are higher in children, explained by relative mineralocorticoid resistance that diminishes with ongoing maturation of the kidney. Children with CAH usually receive mineralocorticoid and salt replacement. However, young adults with CAH should undergo reassessment of their mineralocorticoid reserve. Plasma renin should be regularly monitored and kept within the upper half of the normal reference range.

407 Pheochromocytoma

Hartmut P. H. Neumann

Pheochromocytomas and paragangliomas are catecholamine-producing tumors derived from the sympathetic or parasympathetic nervous system. These tumors may arise sporadically or be inherited as features of multiple endocrine neoplasia type 2, von Hippel–Lindau disease, or several other pheochromocytoma-associated syndromes. The diagnosis of pheochromocytomas identifies a potentially correctable cause of hypertension, and their removal can prevent hypertensive crises that can be lethal. The clinical presentation is variable, ranging from an adrenal incidentaloma to a hypertensive crisis with associated cerebrovascular or cardiac complications.

EPIDEMIOLOGY

Pheochromocytoma is estimated to occur in 2–8 of 1 million persons per year, and ~0.1% of hypertensive patients harbor a pheochromocytoma. The mean age at diagnosis is ~40 years, although the tumors can occur from early childhood until late in life. The classic “rule of tens” for pheochromocytomas states that ~10% are bilateral, 10% are extra-adrenal, and 10% are malignant.

ETIOLOGY AND PATHOGENESIS

Pheochromocytomas and paragangliomas are well-vascularized tumors that arise from cells derived from the sympathetic (e.g., adrenal medulla) or parasympathetic (e.g., carotid body, glomus vagale) paraganglia (Fig. 407-1). The name *pheochromocytoma* reflects the black-colored staining caused by chromaffin oxidation of catecholamines; although a variety of terms have been used to describe these tumors, most clinicians use this designation to describe symptomatic catecholamine-producing tumors, including those in extra-adrenal retroperitoneal, pelvic, and thoracic sites. The term *paraganglioma* is used to describe catecholamine-producing tumors in the skull base and neck; these tumors may secrete little or no catecholamine. In contrast to common clinical parlance, the World Health Organization (WHO) restricts the term *pheochromocytoma* to adrenal tumors and applies the term *paraganglioma* to tumors at all other sites.

The etiology of sporadic pheochromocytomas and paragangliomas is unknown. However, 25–33% of patients have an inherited condition, including germ-line mutations in the classically recognized *RET*, *VHL*, *NF1*, *SDHB*, *SDHC*, and *SDHD* genes or in the more recently recognized *SDHA*, *SDHAF2*, *TMEM127*, and *MAX* genes. Biallelic gene inactivation has been demonstrated for the *VHL*, *NF1*, and *SDH* genes, whereas *RET* mutations activate receptor tyrosine kinase activity. *SDH* is an enzyme of the Krebs cycle and the mitochondrial respiratory chain. The *VHL* protein is a component of a ubiquitin E3 ligase. *VHL* mutations reduce protein degradation, resulting in upregulation of components involved in cell cycle progression, glucose metabolism, and oxygen sensing.

CLINICAL FEATURES

Its clinical presentation is so variable that pheochromocytoma has been termed “the great masquerader” (Table 407-1). Among the presenting manifestations, episodes of palpitation, headache, and profuse sweating are typical, and these manifestations constitute a classic triad. The presence of all three manifestations in association with hypertension makes pheochromocytoma a likely diagnosis. However, a pheochromocytoma can be asymptomatic for years, and some tumors grow to a considerable size before patients note symptoms.

The dominant sign is hypertension. Classically, patients have episodic hypertension, but sustained hypertension is also common. Catecholamine crises can lead to heart failure, pulmonary edema, arrhythmias, and intracranial hemorrhage. During episodes of hormone release, which can occur at widely divergent intervals, patients are anxious and pale, and they experience tachycardia and palpitations. These paroxysms generally last <1 h and may be precipitated by surgery, positional changes, exercise, pregnancy, urination (particularly with bladder pheochromocytomas), and various medications (e.g., tricyclic antidepressants, opiates, metoclopramide).

DIAGNOSIS

The diagnosis is based on documentation of catecholamine excess by biochemical testing and localization of the tumor by imaging. These two criteria are of equal importance, although measurement of catecholamines or metanephrines (their methylated metabolites) is traditionally the first step in diagnosis.

Biochemical testing Pheochromocytomas and paragangliomas synthesize and store catecholamines, which include norepinephrine (noradrenaline), epinephrine (adrenaline), and dopamine. Elevated plasma and urinary levels of catecholamines and metanephrines form the cornerstone of diagnosis. The characteristic fluctuations in the hormonal activity of tumors results in considerable variation in serial catecholamine measurements. However, most tumors continuously leak O-methylated metabolites, which are detected by measurement of metanephrines.

Catecholamines and metanephrines can be measured by different methods, including high-performance liquid chromatography, enzyme-linked immunosorbent assay, and liquid chromatography/mass spectrometry. When pheochromocytoma is suspected on clinical grounds (i.e., when values are three times the upper limit of normal), this diagnosis is highly likely regardless of the assay used. However, as summarized in Table 407-2, the sensitivity and specificity of available biochemical tests vary greatly, and these differences are important in assessing patients with borderline elevations of different compounds. Urinary tests for metanephrines (total or fractionated) and catecholamines are widely available and are used commonly for initial evaluation. Among these tests, those for the fractionated metanephrines and catecholamines are the most sensitive. Plasma tests are more convenient and include measurements of catecholamines and metanephrines. Measurements of plasma metanephrine are the most sensitive and are less susceptible to false-positive elevations from stress, including venipuncture. Although the incidence of false-positive test results has been reduced by the introduction of newer assays, physiologic stress responses and medications that increase catecholamine levels still can confound testing. Because the tumors are relatively rare, borderline elevations are likely to represent false-positive results. In

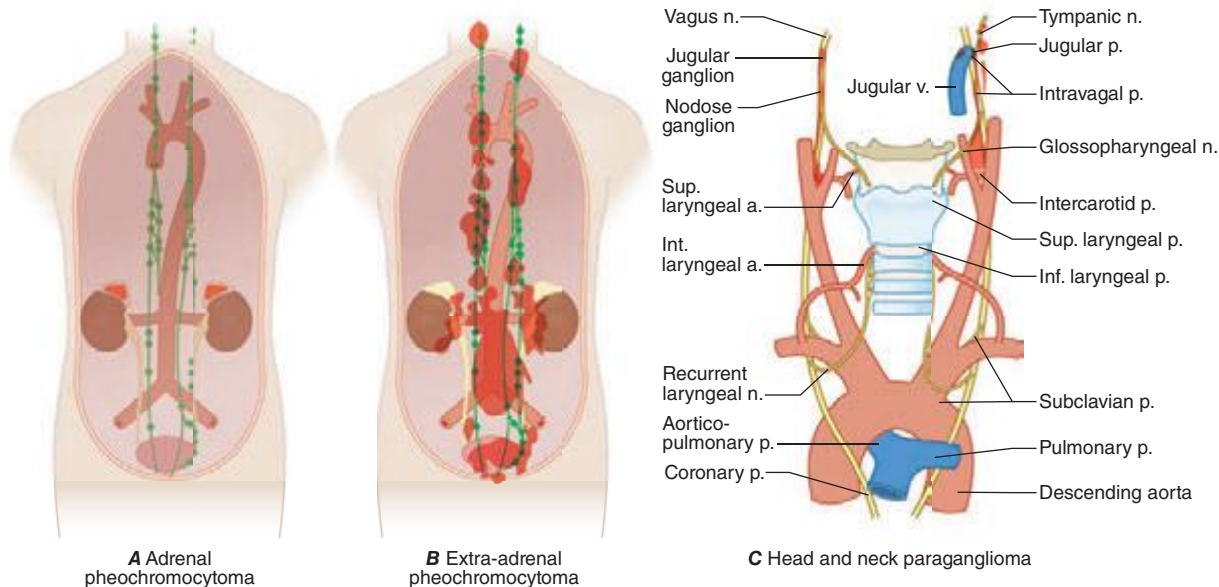


FIGURE 407-1 The paraganglial system and topographic sites (in red) of pheochromocytomas and paragangliomas. (Parts A and B from WM Manger, RW Gifford: *Clinical and experimental pheochromocytoma*. Cambridge, Blackwell Science, 1996; Part C from GG Glenner, PM Grimley: *Tumors of the Extra-adrenal Paraganglion System [Including Chemoreceptors]*, Atlas of Tumor Pathology, 2nd Series, Fascicle 9. Washington, DC, AFIP, 1974.)

TABLE 407-1 CLINICAL FEATURES ASSOCIATED WITH PHEOCHROMOCYTOMA, LISTED BY FREQUENCY OF OCCURRENCE

1. Headaches	10. Weight loss
2. Profuse sweating	11. Paradoxical response to antihypertensive drugs
3. Palpitations and tachycardia	12. Polyuria and polydipsia
4. Hypertension, sustained or paroxysmal	13. Constipation
5. Anxiety and panic attacks	14. Orthostatic hypotension
6. Pallor	15. Dilated cardiomyopathy
7. Nausea	16. Erythrocytosis
8. Abdominal pain	17. Elevated blood sugar
9. Weakness	18. Hypercalcemia

this circumstance, it is important to exclude dietary or drug-related factors (withdrawal of levodopa or use of sympathomimetics, diuretics, tricyclic antidepressants, alpha and beta blockers) that might cause false-positive results and then to repeat testing or perform a clonidine suppression test (i.e., the measurement of plasma normetanephrine 3 h after oral administration of 300 µg of clonidine). Other pharmacologic tests, such as the phentolamine test and the glucagon provocation test, are of relatively low sensitivity and are not recommended.

Diagnostic Imaging A variety of methods have been used to localize pheochromocytomas and paragangliomas (Table 407-2). CT and MRI are similar in sensitivity and should be performed with contrast. T2-weighted MRI with gadolinium contrast is optimal for detecting extraadrenal pheochromocytomas and paragangliomas. About 5% of adrenal incidentalomas, which usually are detected by CT or MRI, prove to be pheochromocytomas upon endocrinologic evaluation.

Tumors also can be localized by procedures using radioactive tracers, including ¹³¹I- or ¹²³I-metaiodobenzylguanidine (MIBG) scintigraphy, ¹¹¹In-somatostatin analogue scintigraphy, ¹⁸F-DOPA positron emission tomography (PET), or ¹⁸F-fluorodeoxyglucose (FDG) PET. Because these agents exhibit selective uptake in paragangliomas, nuclear imaging is particularly useful in the hereditary syndromes.

Differential Diagnosis When the possibility of a pheochromocytoma is being entertained, other disorders to consider include essential hypertension, anxiety attacks, use of cocaine or amphetamines, mastocytosis or carcinoid syndrome (usually without hypertension), intracranial lesions, clonidine withdrawal, autonomic epilepsy, and factitious crises (usually from use of sympathomimetic amines). When an asymptomatic adrenal mass is identified, likely diagnoses other than pheochromocytoma include a nonfunctioning adrenal adenoma, an aldosteronoma, and a cortisol-producing adenoma (Cushing's syndrome).

TABLE 407-2 BIOCHEMICAL AND IMAGING METHODS USED FOR DIAGNOSIS OF PHEOCHROMOCYTOMA AND PARAGANGLIOMA

Diagnostic Method	Sensitivity	Specificity
24-h urinary tests		
Catecholamines	+++	+++
Fractionated metanephrines	++++	++
Total metanephrines	+++	++++
Plasma tests		
Catecholamines	+++	++
Free metanephrines	++++	+++
Imaging		
CT	++++	+++
MRI	++++	+++
MIBG scintigraphy	+++	++++
Somatostatin receptor scintigraphy ^a	++	++
¹⁸ F-DOPA PET/CT	+++	++++

^aValues are particularly high in head and neck paragangliomas.

Abbreviations: MIBG, metaiodobenzylguanidine; PET/CT, positron emission tomography plus CT. For the biochemical tests, the ratings correspond globally to sensitivity and specificity rates as follows: ++, <85%; +++, 85–95%; and +, >95%.

TREATMENT PHEOCHROMOCYTOMA

Complete tumor removal, the ultimate therapeutic goal, can be achieved by partial or total adrenalectomy. It is important to preserve the normal adrenal cortex, particularly in hereditary disorders in which bilateral pheochromocytomas are most likely. Preoperative preparation of the patient is important. Before surgery, blood pressure should be consistently below 160/90 mmHg. Classically, blood pressure has been controlled by α-adrenergic blockers (oral

phenoxybenzamine, 0.5–4 mg/kg of body weight). Because patients are volume-constricted, liberal salt intake and hydration are necessary to avoid severe orthostasis. Oral prazosin or intravenous phentolamine can be used to manage paroxysms while adequate alpha blockade is awaited. Beta blockers (e.g., 10 mg of propranolol three or four times per day) can then be added. Other antihypertensives, such as calcium channel blockers or angiotensin-converting enzyme inhibitors, have also been used effectively.

Surgery should be performed by teams of surgeons and anesthesiologists with experience in the management of pheochromocytomas. Blood pressure can be labile during surgery, particularly at the outset of intubation or when the tumor is manipulated. Nitroprusside infusion is useful for intraoperative hypertensive crises, and hypotension usually responds to volume infusion.

Minimally invasive techniques (laparoscopy or retroperitoneoscopy) have become the standard approaches in pheochromocytoma surgery. They are associated with fewer complications, a faster recovery, and optimal cosmetic results. Extra-adrenal abdominal and most thoracic pheochromocytomas also can also be removed endoscopically. Postoperatively, catecholamine normalization should be documented. An adrenocorticotropic hormone test should be used to exclude cortisol deficiency when bilateral adrenal cortex-sparing surgery has been performed.

MALIGNANT PHEOCHROMOCYTOMA

About 5–10% of pheochromocytomas and paragangliomas are malignant. The diagnosis of malignant pheochromocytoma is problematic. The typical histologic criteria of cellular atypia, presence of mitoses, and invasion of vessels or adjacent tissues are insufficient for the diagnosis of malignancy in pheochromocytoma. Thus, the term *malignant pheochromocytoma* is restricted to tumors with distant metastases, most commonly found by nuclear medicine imaging in lungs, bone, or liver—locations suggesting a vascular pathway of spread. Because hereditary syndromes are associated with multifocal tumor sites, these

features should be anticipated in patients with germ-line mutations of *RET*, *VHL*, *SDHD*, or *SDHB*. However, distant metastases also occur in these syndromes, especially in carriers of *SDHB* mutations.

Treatment of malignant pheochromocytoma or paraganglioma is challenging. Options include tumor mass reduction, alpha blockers for symptoms, chemotherapy, and nuclear medicine radiotherapy. The first-line choice is nuclear medicine therapy for scintigraphically documented metastases, preferably with ^{131}I -MIBG in 200-mCi doses at monthly intervals over three to six cycles. Averbuch's chemotherapy protocol includes dacarbazine (600 mg/m² on days 1 and 2), cyclophosphamide (750 mg/m² on day 1), and vincristine (1.4 mg/m² on day 1), all repeated every 21 days for three to six cycles. Palliation (stable disease to shrinkage) is achieved in about one-half of patients. Other chemotherapeutic options are sunitinib and temozolomide/thalidomide. The prognosis of metastatic pheochromocytoma or paraganglioma is variable, with 5-year survival rates of 30–60%.

PHEOCHROMOCYTOMA IN PREGNANCY

Pheochromocytomas occasionally are diagnosed in pregnancy. Endoscopic removal, preferably in the fourth to sixth month of gestation, is possible and can be followed by uneventful childbirth. Regular screening in families with inherited pheochromocytomas provides an opportunity to identify and remove asymptomatic tumors in women of reproductive age.

PHEOCHROMOCYTOMA-ASSOCIATED SYNDROMES

About 25–33% of patients with a pheochromocytoma or paraganglioma have an inherited syndrome. At diagnosis, patients with inherited syndromes are a mean of ~15 years younger than patients with sporadic tumors.

Neurofibromatosis type 1 (NF1) was the first described pheochromocytoma-associated syndrome (**Chap. 118**). The *NF1* gene functions as a tumor suppressor by regulating the Ras signaling cascade. Classic features of neurofibromatosis include multiple neurofibromas, café au lait spots, axillary freckling, and Lisch nodules of the iris (**Fig. 407-2**). Pheochromocytomas occur in only ~1% of these patients

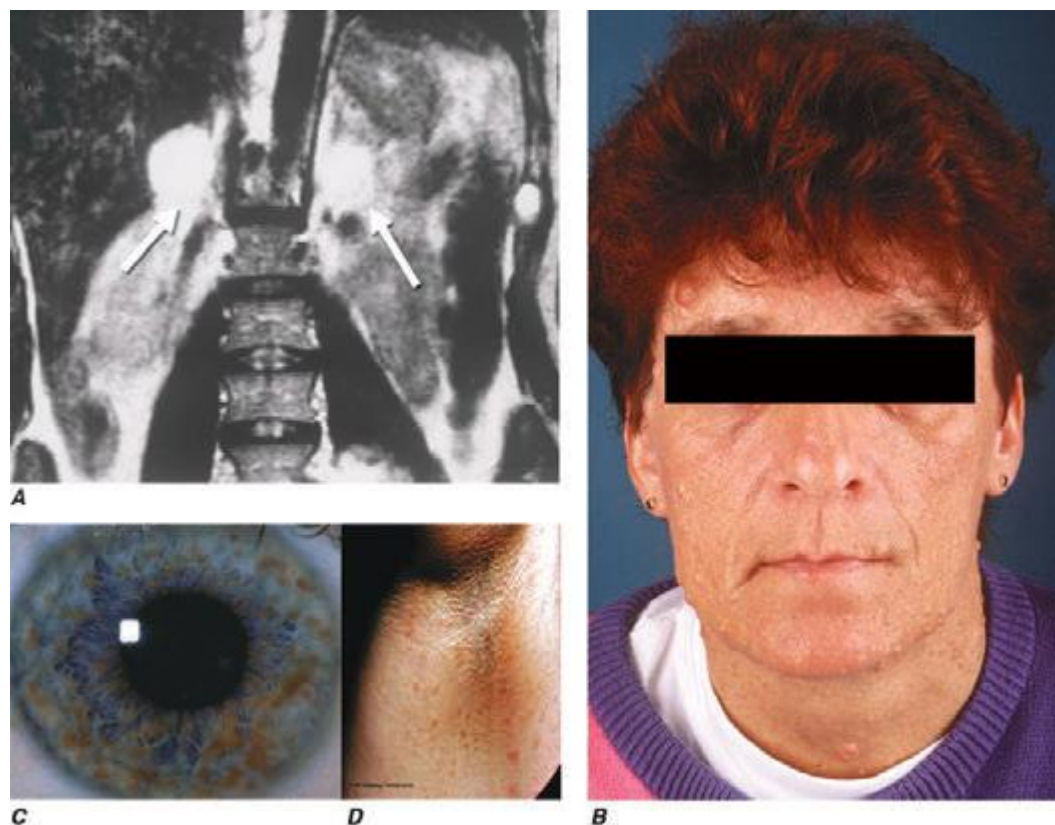


FIGURE 407-2 Neurofibromatosis. **A.** MRI of bilateral adrenal pheochromocytoma. **B.** Cutaneous neurofibromas. **C.** Lisch nodules of the iris. **D.** Axillary freckling. (Part A from HPH Neumann et al. *Keio J Med* 54:15, 2005; with permission.)

2332 and are located predominantly in the adrenals. Malignant pheochromocytoma is not uncommon.

The best-known pheochromocytoma-associated syndrome is the autosomal dominant disorder *multiple endocrine neoplasia type 2* (MEN2) (Chap. 408). Both types of MEN2 (2A and 2B) are caused by mutations in *RET* (rearranged during transfection), which encodes a tyrosine kinase. The locations of *RET* mutations correlate with the severity of disease and the type of MEN2 (Chap. 408). MEN2A is characterized by medullary thyroid carcinoma (MTC), pheochromocytoma, and hyperparathyroidism; MEN2B also includes MTC and pheochromocytoma as well as multiple mucosal neuromas, marfanoid habitus, and other developmental disorders, though it typically lacks hyperparathyroidism. MTC is found in virtually all patients with MEN2, but pheochromocytoma occurs in only ~50% of these patients. Nearly all pheochromocytomas in MEN2 are benign and located in the adrenals, often bilaterally (Fig. 407-3). Pheochromocytoma may be symptomatic before MTC. Prophylactic thyroidectomy is being performed in many carriers of *RET* mutations; pheochromocytomas should be excluded before any surgery in these patients.

Von Hippel-Lindau syndrome (VHL) is an autosomal dominant disorder that predisposes to retinal and cerebellar hemangioblastomas, which also occur in the brainstem and spinal cord (Fig. 407-4). Other important features of VHL are clear cell renal carcinomas, pancreatic neuroendocrine tumors, endolymphatic sac tumors of the inner ear, cystadenomas of the epididymis and broad ligament, and multiple pancreatic or renal cysts.

The *VHL* gene (among other genes) encodes an E3 ubiquitin ligase that regulates expression of hypoxia-inducible factor 1. Loss of *VHL* is associated with increased expression of vascular endothelial growth factor (VEGF), which induces angiogenesis. Although the *VHL* gene can be inactivated by all types of mutations, patients with pheochromocytoma predominantly have missense mutations. About 20–30% of patients with VHL have pheochromocytomas, but in some families the incidence can reach 90%. The recognition of pheochromocytoma as a VHL-associated feature provides an opportunity to diagnose retinal, central nervous system, renal, and pancreatic tumors at a stage when effective treatment may still be possible.

The *paraganglioma syndromes* (PGLs) have been classified by genetic analyses of families with head and neck paragangliomas. The susceptibility genes encode subunits of the enzyme succinate dehydrogenase (SDH), a component in the Krebs cycle and the mitochondrial electron transport chain. SDH is formed by four subunits (A–D). Mutations of *SDHB* (PGL4), *SDHC* (PGL3), *SDHD* (PGL1), and *SDHAF2* (PGL2) predispose to the PGLs. The transmission of the disease in carriers of *SDHB* and *SDHC* germ-line mutations is autosomal dominant. In contrast, in *SDHD* and *SDHAF2* families, only the progeny of affected fathers develop tumors if they inherit the mutation. PGL1 is most common, followed by PGL4; PGL2 and PGL3 are rare. Adrenal, extra-adrenal abdominal, and thoracic pheochromocytomas, which are components of PGL1 and PGL4, are rare in PGL3 and absent in PGL2 (Fig. 407-5). About one-third of patients with PGL4 develop metastases.

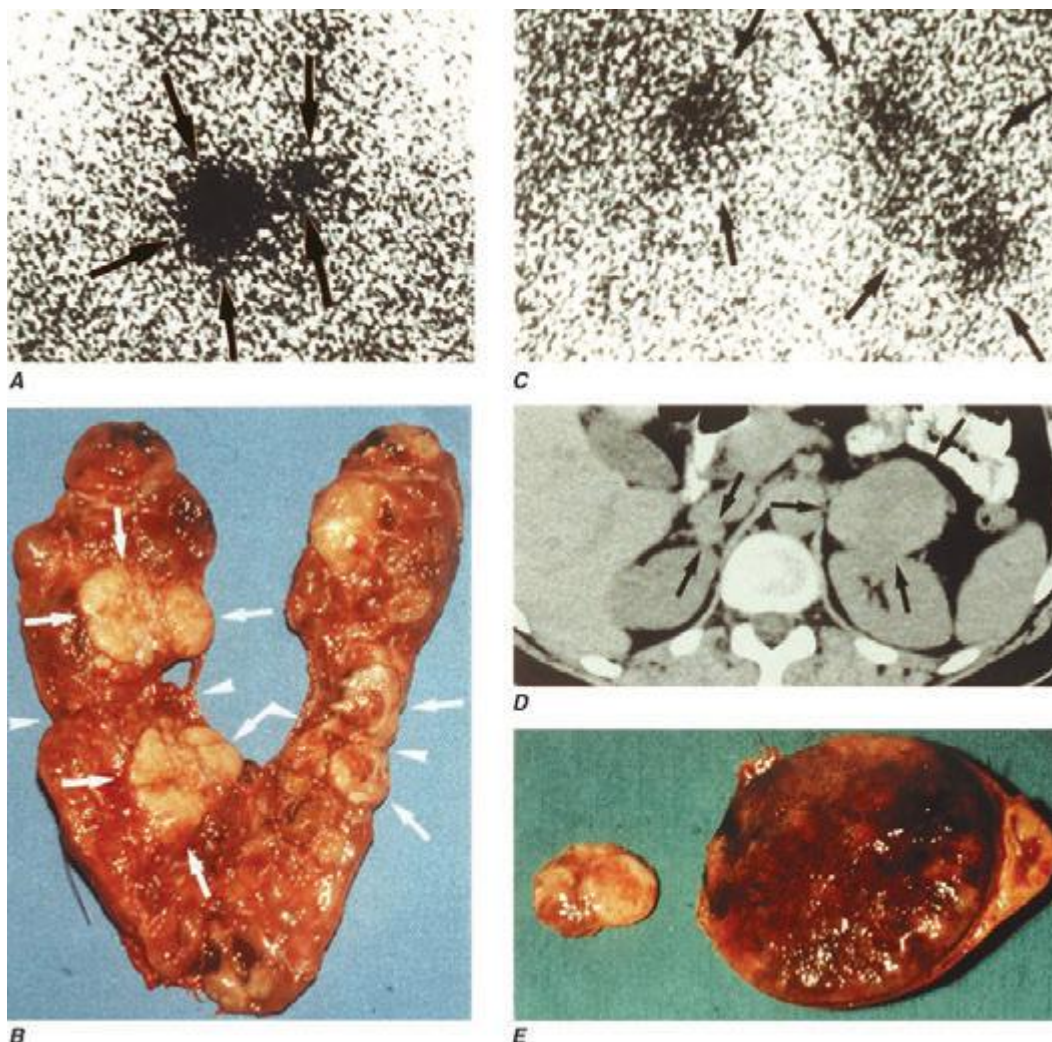


FIGURE 407-3 Multiple endocrine neoplasia type 2. **A, B.** Multifocal medullary thyroid carcinoma shown by MIBG scintigraphy (A) and operative specimen (B). Arrows demonstrate the tumors; arrowheads show the tissue bridge of the cut specimen. **C–E.** Bilateral adrenal pheochromocytoma shown by MIBG scintigraphy (C), CT imaging (D), and operative specimens (E). (From HPH Neumann et al: *Keio J Med* 54:15, 2005; with permission.)

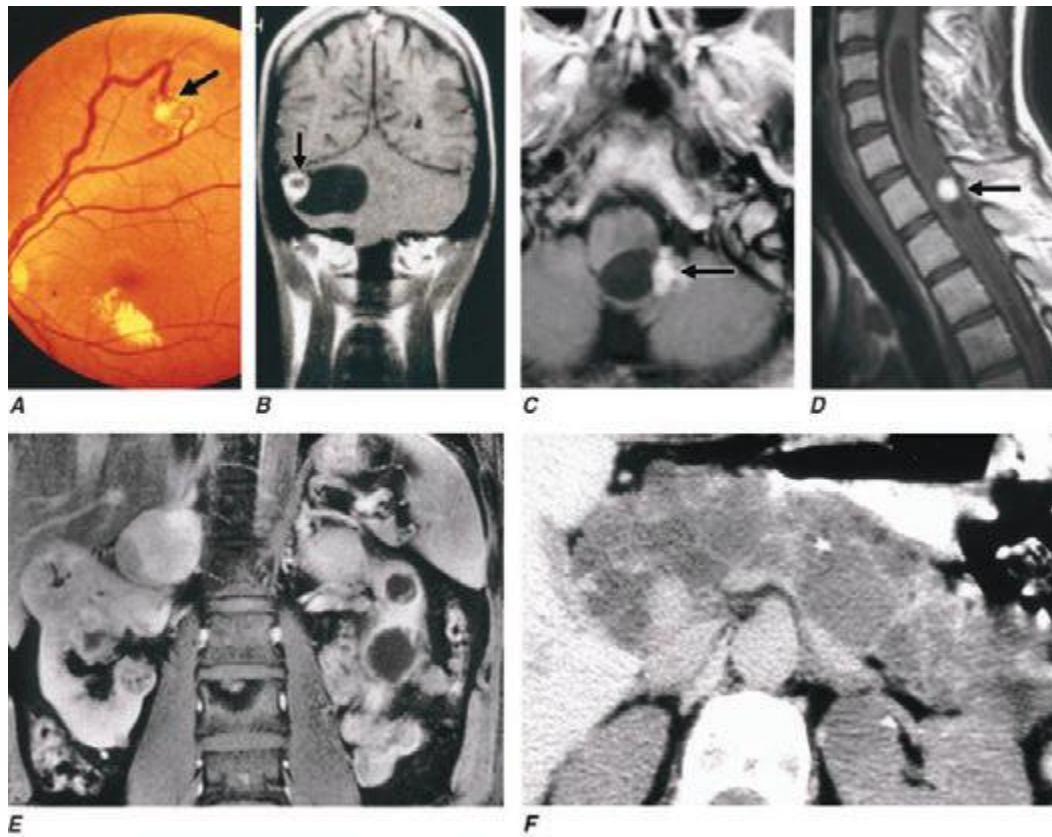


FIGURE 407-4 Von Hippel-Lindau disease. **A.** Retinal angioma. All subsequent panels show findings on MRI: **B–D.** Hemangioblastomas of the cerebellum (**B**) in brainstem (**C**) and spinal cord (**D**). **E.** Bilateral pheochromocytomas and bilateral renal clear cell carcinomas **F.** Multiple pancreatic cysts. (Parts A and D from HPH Neumann et al: *Adv Nephrol Necker Hosp* 27:361, 1997. © Elsevier. Part B from SH Morgan, J-P Grunfeld [eds]: *Inherited Disorders of the Kidney*. Oxford, UK, Oxford University Press, 1998. Part F from HPH Neumann et al: *Contrib Nephrol* 136:193, 2001. © S. Karger AG, Basel.)

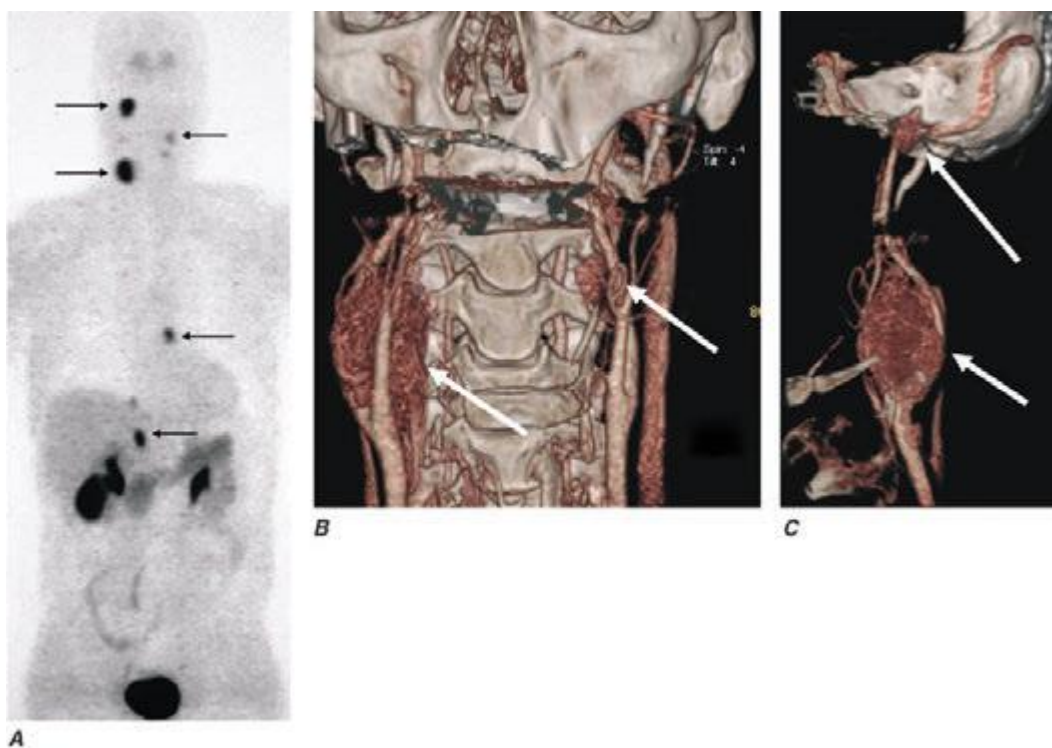


FIGURE 407-5 Paraganglioma syndrome. A patient with the SDHD W5X mutation and PGL1 underwent incomplete resection of a left carotid body tumor. **A.** ^{18}F -DOPA positron emission tomography demonstrating tumor uptake in the right jugular glomus, the right carotid body, the left carotid body, the left coronary glomus, and the right adrenal gland. Note the physiologic accumulation of the radiopharmaceutical agent in the kidneys, liver, gallbladder, renal pelvis, and urinary bladder. **B** and **C.** CT angiography with three-dimensional reconstruction. Arrows point to the paraganglial tumors. (From S Hoegerle et al: *Eur J Nucl Med Mol Imaging* 30:689, 2003; with permission.)

2334 *Familial pheochromocytoma (FP)* has been attributed to hereditary, mainly adrenal tumors in patients with germ-line mutations in the genes *TMEM127*, *MAX*, and *SDHA*. Transmission is also autosomal dominant, and mutations of *MAX*, like those of *SDHD*, cause tumors only if inherited from the father.

GUIDELINES FOR GENETIC SCREENING OF PATIENTS WITH PHEOCHROMOCYTOMA OR PARAGANGLIOMA

In addition to family history, general features suggesting an inherited syndrome include young age, multifocal tumors, extra-adrenal tumors, and malignant tumors (Fig. 407-6). Because of the relatively high prevalence of familial syndromes among patients who present with pheochromocytoma or paraganglioma, it is useful to identify

germ-line mutations even in patients without a known family history. A first step is to search for clinical features of inherited syndromes and to obtain an in-depth, multigenerational family history. Each of these syndromes exhibits autosomal dominant transmission with variable penetrance, but a proband with a mother affected by paraganglial tumors is not predisposed to *PLG1* (*SDHD* mutation carrier). Cutaneous neurofibromas, café au lait spots, and axillary freckling suggest neurofibromatosis. Germ-line mutations in *NF1* have not been reported in patients with sporadic pheochromocytomas. Thus, *NF1* testing need not be performed in the absence of other clinical features of neurofibromatosis. A personal or family history of MTC or an elevation of serum calcitonin strongly suggests MEN 2 and should prompt testing for *RET* mutations. A history of visual impairment or

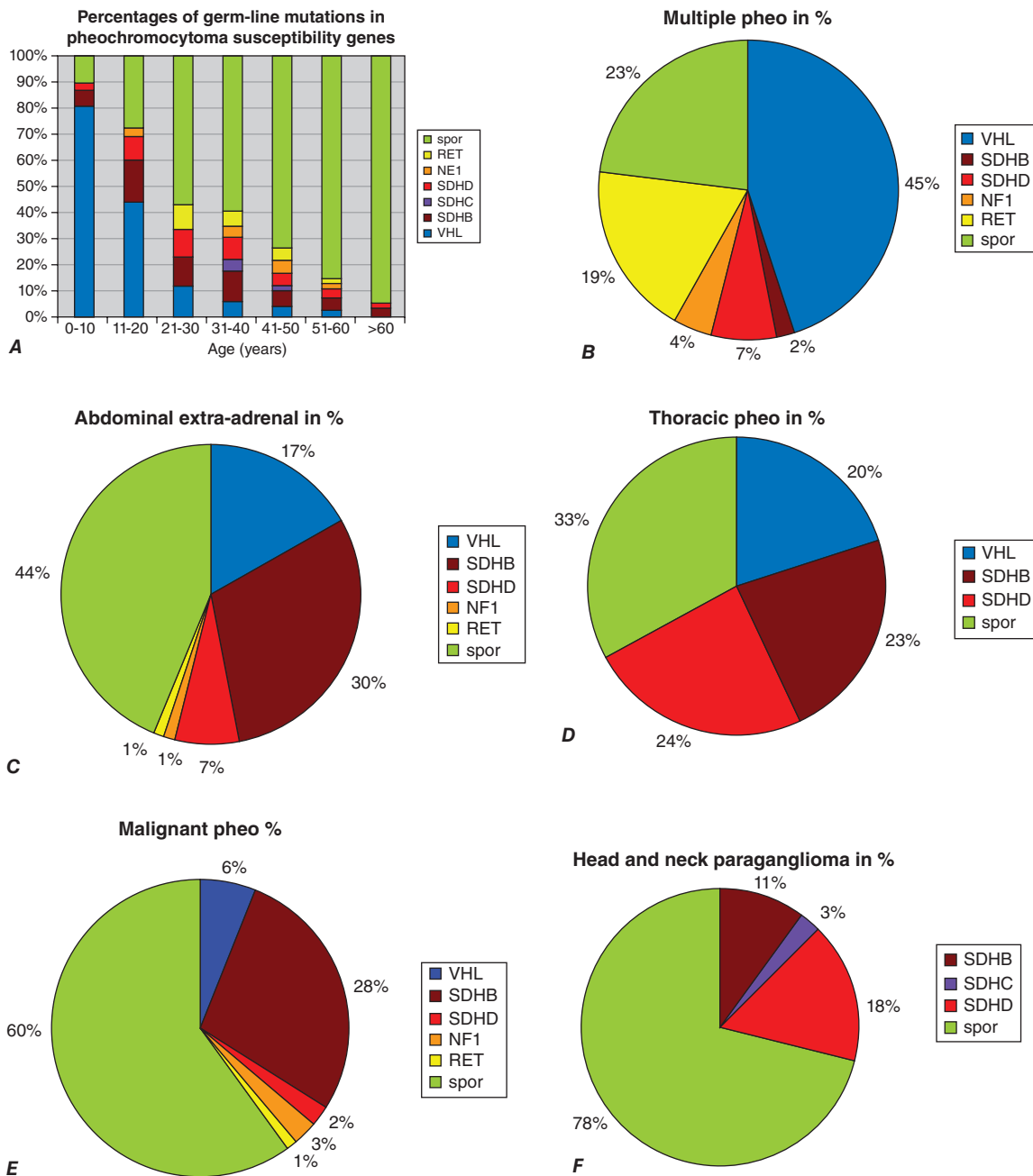


FIGURE 407-6 Mutation distribution in the *VHL*, *RET*, *SDHB*, *SDHC*, *SDHD*, and *NF1* genes in 2021 patients with pheochromocytomas and paragangliomas from the European-American Pheochromocytoma-Paraganglioma Registry based in Freiburg, Germany, as updated on March 1, 2014. **A.** Correlation with age. The bars depict the frequency of sporadic (spor) or various inherited forms of pheochromocytoma in different age groups. The inherited disorders are much more common among younger individuals presenting with pheochromocytoma. Patients with mutations in the *TMEM127*, *MAX*, and *SDHA* genes are not included, since they contribute <1% in decades 4–7 only. **B–F.** Germ-line mutations according to multiple (B), extra-adrenal retroperitoneal (C), thoracic (D), and malignant (E) pheochromocytomas and head and neck paragangliomas (F). (Data from the Freiburg International Pheochromocytoma and Paraganglioma Registry, 2014.)

tumors of the cerebellum, kidney, brainstem, or spinal cord suggests the possibility of VHL. A personal and/or family history of head and neck paraganglioma suggests PGL1 or PGL4.

A single adrenal pheochromocytoma in a patient with an otherwise unremarkable history may still be associated with mutations of *VHL*, *RET*, *SDHB*, or *SDHD* (in decreasing order of frequency). Two-thirds of extra-adrenal tumors are associated with one of these syndromes, and multifocal tumors occur with decreasing frequency in carriers of *RET*, *SDHD*, *VHL*, and *SDHB* mutations. About 30% of head and neck paragangliomas are associated with germ-line mutations of one of the SDH subunit genes (most often *SDHD*) and are rare in carriers of *VHL*, *RET*, and *TMEM127* mutations (Fig. 407-6F).

Immunohistochemistry is helpful in the preselection of hereditary pheochromocytoma. Negative immunostaining with antibodies to SDHB, TMEM127, and MAX may predict mutations of the *SDH*, *TMEM127*, and *MAX* genes, respectively.

Once the underlying syndrome is diagnosed, the benefit of genetic testing can be extended to relatives. For this purpose, it is necessary to identify the germ-line mutation in the proband and, after genetic counseling, to perform DNA sequence analyses of the responsible gene in relatives to determine whether they are affected (Chap. 84). Other family members may benefit when individuals who carry a germ-line mutation are biochemically screened for paraganglial tumors.

Asymptomatic paraganglial tumors, now often detected in patients with hereditary tumors and their relatives, are challenging to manage. Watchful waiting strategies have been introduced. Head and neck paragangliomas—mainly carotid body, jugular, and vagal tumors—are increasingly treated by radiation, since surgery is frequently associated with permanent palsy of cranial nerves II, VII, IX, X, XI, and XII. Nevertheless, tympanic paragangliomas are symptomatic early, and most of these tumors can easily be resected, with subsequent improvement of hearing and alleviation of tinnitus.

408 Multiple Endocrine Neoplasia

Rajesh V. Thakker

Multiple endocrine neoplasia (MEN) is characterized by a predilection for tumors involving two or more endocrine glands. Four major forms of MEN are recognized and referred to as MEN types 1–4 (MEN 1–4) (Table 408-1). Each type of MEN is inherited as an autosomal dominant syndrome or may occur sporadically; that is, without a family history. However, this distinction between familial and sporadic forms is often difficult because family members with the disease may have died before symptoms developed. In addition to MEN 1–4, at least six other syndromes are associated with multiple endocrine and other organ neoplasias (MEONs) (Table 408-2). These MEONs include the hyperparathyroidism–jaw tumor syndrome, Carney complex, von Hippel–Lindau disease (Chap. 407), neurofibromatosis type 1 (Chap. 118), Cowden's syndrome, and McCune–Albright syndrome (Chap. 426e); all of these are inherited as autosomal dominant disorders, except for McCune–Albright syndrome, which is caused by mosaic expression of a postzygotic somatic cell mutation (Table 408-2).

A diagnosis of a MEN or MEON syndrome may be established in an individual by one of three criteria: (1) clinical features (two or more of the associated tumors [or lesions] in an individual); (2) familial pattern (one of the associated tumors [or lesions] in a first-degree relative of a patient with a clinical diagnosis of the syndrome); and (3) genetic analysis (a germline mutation in the associated gene in an individual, who may be clinically affected or asymptomatic). Mutational analysis in MEN and MEON syndromes is helpful in clinical practice to: (1) confirm the clinical diagnosis; (2) identify family members who harbor the mutation and require screening for relevant tumor detection and early/appropriate treatment; and (3) identify the ~50% of family

members who do not harbor the germline mutation and can, therefore, be alleviated of the anxiety of developing associated tumors. This latter aspect also helps to reduce health care costs by reducing the need for unnecessary biochemical and radiologic investigations.

MULTIPLE ENDOCRINE NEOPLASIA TYPE 1

Clinical Manifestations MEN type 1 (MEN 1), which is also referred to as Wermer's syndrome, is characterized by the triad of tumors involving the parathyroids, pancreatic islets, and anterior pituitary. In addition, adrenal cortical tumors, carcinoid tumors usually of the foregut, meningiomas, facial angiofibromas, collagenomas, and lipomas may also occur in some patients with MEN 1. Combinations of the affected glands and their pathologic features (e.g., hyperplastic adenomas of the parathyroid glands) may differ in members of the same family and even between identical twins. In addition, a nonfamilial (e.g., sporadic) form occurs in 8–14% of patients with MEN 1, and molecular genetic studies have confirmed the occurrence of de novo mutations of the *MEN1* gene in approximately 10% of patients with MEN 1. The prevalence of MEN 1 is approximately 0.25% based on randomly chosen postmortem studies but is 1–18% among patients with primary hyperparathyroidism, 16–38% among patients with pancreatic islet tumors, and <3% among patients with pituitary tumors. The disorder affects all age groups, with a reported age range of 5 to 81 years, with clinical and biochemical manifestations developing in the vast majority by the fifth decade. The clinical manifestations of MEN 1 are related to the sites of tumors and their hormonal products. In the absence of treatment, endocrine tumors are associated with an earlier mortality in patients with MEN 1, with a 50% probability of death by the age of 50 years. The cause of death is usually a malignant tumor, often from a pancreatic neuroendocrine tumor (NET) or foregut carcinoid. In addition, the treatment outcomes of patients with MEN 1–associated tumors are not as successful as those in patients with non–MEN 1 tumors. This is because MEN 1–associated tumors, with the exception of pituitary NETs, are usually multiple, making it difficult to achieve a successful surgical cure. Occult metastatic disease is also more prevalent in MEN 1, and the tumors may be larger, more aggressive, and resistant to treatment.

Parathyroid Tumors (See also Chap. 424) Primary hyperparathyroidism occurs in approximately 90% of patients and is the most common feature of MEN 1. Patients may have asymptomatic hypercalcemia or vague symptoms associated with hypercalcemia (e.g., polyuria, polydipsia, constipation, malaise, or dyspepsia). Nephrolithiasis and osteitis fibrosa cystica (less commonly) may also occur. Biochemical investigations reveal hypercalcemia, usually in association with elevated circulating parathyroid hormone (PTH) (Table 408-3). The hypercalcemia is usually mild, and severe hypercalcemia or parathyroid cancer is a rare occurrence. Additional differences in the primary hyperparathyroidism of patients with MEN 1, as opposed to those without MEN 1, include an earlier age at onset (20–25 years vs 55 years) and an equal male-to-female ratio (1:1 vs 1:3). Preoperative imaging (e.g., neck ultrasound with ^{99m}Tc-sestamibi parathyroid scintigraphy) is of limited benefit because all parathyroid glands may be affected, and neck exploration may be required irrespective of preoperative localization studies.

TREATMENT PARATHYROID TUMORS

Surgical removal of the abnormally overactive parathyroids in patients with MEN 1 is the definitive treatment. However, it is controversial whether to perform subtotal (e.g., removal of 3.5 glands) or total parathyroidectomy with or without autotransplantation of parathyroid tissue in the forearm, and whether surgery should be performed at an early or late stage. Minimally invasive parathyroidectomy is not recommended because all four parathyroid glands are usually affected with multiple adenomas or hyperplasia. Surgical experience should be taken into account given the variability in pathology in MEN 1. Calcimimetics (e.g., cinacalcet), which act via

TABLE 408-1 MULTIPLE ENDOCRINE NEOPLASIA (MEN) SYNDROMES

Type (Chromosomal Location)	Tumors (Estimated Penetrance)	Gene and Most Frequently Mutated Codons
MEN 1 (11q13)	Parathyroid adenoma (90%) Enteropancreatic tumor (30–70%) <ul style="list-style-type: none"> • Gastrinoma (>50%) • Insulinoma (10–30%) • Nonfunctioning and PPoma (20–55%) • Glucagonoma (<3%) • VIPoma (<1%) Pituitary adenoma (15–50%) <ul style="list-style-type: none"> • Prolactinoma (60%) • Somatotrophinoma (25%) • Corticotropinoma (<5%) • Nonfunctioning (<5%) Associated tumors <ul style="list-style-type: none"> • Adrenal cortical tumor (20–70%) • Pheochromocytoma (<1%) • Bronchopulmonary NET (2%) • Thymic NET (2%) • Gastric NET (10%) • Lipomas (>33%) • Angiofibromas (85%) • Collagenomas (70%) • Meningiomas (8%) 	<i>MEN1</i> 83/84, 4-bp del (≈4%) 119, 3-bp del (≈3%) 209-211, 4-bp del (≈8%) 418, 3-bp del (≈4%) 514-516, del or ins (≈7%) Intron 4 ss (≈10%)
MEN 2 (10 cen-10q11.2)		
MEN 2A	MTC (90%) Pheochromocytoma (>50%) Parathyroid adenoma (10–25%)	<i>RET</i> 634, e.g., Cys → Arg (~85%)
MTC only	MTC (100%)	<i>RET</i> 618, missense (>50%)
MEN 2B (also known as MEN 3)	MTC (>90%) Pheochromocytoma (>50%) Associated abnormalities (40–50%) <ul style="list-style-type: none"> • Mucosal neuromas • Marfanoid habitus • Medullated corneal nerve fibers • Megacolon 	<i>RET</i> 918, Met → Thr (>95%)
MEN 4 (12p13)	Parathyroid adenoma ^a Pituitary adenoma ^a Reproductive organ tumors ^a (e.g., testicular cancer, neuroendocrine cervical carcinoma) ?Adrenal + renal tumors ^a	<i>CDKN1B</i> ; no common mutations identified to date

^aInsufficient numbers reported to provide prevalence information.

Note: Autosomal dominant inheritance of the MEN syndromes has been established.

Abbreviations: del, deletion; ins, insertion; MTC, medullary thyroid cancer; NET, neuroendocrine tumor; PPoma, pancreatic polypeptide–secreting tumor; VIPoma, vasoactive intestinal polypeptide–secreting tumor.

Source: Reproduced from RV Thakker et al: *J Clin Endocrinol Metab* 97:2990, 2012.

the calcium-sensing receptor, have been used to treat primary hyperparathyroidism in some patients when surgery is unsuccessful or contraindicated.

Pancreatic Tumors (See also Chap. 113) The incidence of pancreatic islet cell tumors, which are NETs, in patients with MEN 1 ranges from 30 to 80% in different series. Most of these tumors (Table 408-1) produce excessive amounts of hormone (e.g., gastrin, insulin, glucagon, vasoactive intestinal polypeptide [VIP]) and are associated with distinct clinical syndromes, although some are nonfunctioning or nonsecretory. These pancreatic islet cell tumors have an earlier age at onset in patients with MEN 1 than in patients without MEN 1.

Gastrinoma Gastrin-secreting tumors (gastrinomas) are associated with marked gastric acid production and recurrent peptic ulcerations, a combination referred to as the Zollinger-Ellison syndrome. Gastrinomas occur more often in patients with MEN 1 who are older than age 30 years. Recurrent severe multiple peptic ulcers, which may perforate, and cachexia are major contributors to the high mortality. Patients with Zollinger-Ellison syndrome may also suffer from diarrhea and steatorrhea. The diagnosis is established by demonstration of an elevated fasting serum gastrin concentration in association with increased basal gastric acid secretion (Table 408-3). However, the diagnosis of Zollinger-Ellison syndrome may be difficult in hypercalcemic MEN 1 patients, because hypercalcemia can also cause hypergastrinemia.

TABLE 408-2 MULTIPLE ENDOCRINE AND OTHER ORGAN NEOPLASIA SYNDROMES (MEONS)

Disease ^a	Gene Product	Chromosomal Location
Hyperparathyroidism-jaw tumor (HPT-JT)	Parafibromin	1q31.2
Carney complex		
CNC1	PPKAR1A	17q24.2
CNC2	? ^b	2p16
von Hippel-Lindau disease (VHL)	pVHL (elongin)	3p25
Neurofibromatosis type 1 (NF1)	Neurofibromin	17q11.2
Cowden's syndrome (CWD)		
CWD1	PTEN	10q23.31
CWD2	SDHB	1p36.13
CWD3	SDHD	11q23.1
CWD4	KLLN	10q23.31
CWD5	PIK3CA	3q26.32
CWD6	AKT1	14q32.33
McCune-Albright syndrome (MAS)	Gsa	20q13.32

^aThe inheritance for these disorders is autosomal dominant, except MAS, which is due to mosaicism that results from the postzygotic somatic cell mutation of the *GNAS1* gene, encoding Gsa. ^b?, unknown.

Ultrasonography, endoscopic ultrasonography, computed tomography (CT), nuclear magnetic resonance imaging (MRI), selective abdominal angiography, venous sampling, and somatostatin receptor scintigraphy are helpful in localizing the tumor prior to surgery. Gastrinomas represent more than 50% of all pancreatic NETs in patients with MEN 1, and approximately 20% of patients with gastrinomas will be found to have MEN 1. Gastrinomas, which may also occur in the duodenal mucosa, are the major cause of morbidity and mortality in patients with MEN 1. Most MEN 1 gastrinomas are malignant and metastasize before a diagnosis is established.

TREATMENT GASTRINOMA

Medical treatment of patients with MEN 1 and Zollinger-Ellison syndrome is directed toward reducing basal acid output to <10 mmol/L. Parietal cell H⁺-K⁺-adenosine triphosphatase (ATPase) inhibitors (e.g., omeprazole or lansoprazole) reduce acid output and are the drugs of choice for gastrinomas. Some patients may also require additional treatment with the histamine H₂ receptor antagonists, cimetidine or ranitidine. The role of surgery in the treatment of gastrinomas in patients with MEN 1 is controversial. The goal of surgery is to reduce the risk of distant metastatic disease and improve survival. For a nonmetastatic gastrinoma situated in the pancreas, surgical excision is often effective. However, the risk of hepatic metastases increases with tumor size, such that 25–40% of patients

with pancreatic NETs >4 cm develop hepatic metastases, and 50–70% of patients with tumors 2–3 cm in size have lymph node metastases. Survival in MEN 1 patients with gastrinomas <2.5 cm in size is 100% at 15 years, but 52% at 15 years, if metastatic disease is present. The presence of lymph node metastases does not appear to adversely affect survival. Surgery for gastrinomas that are >2–2.5 cm has been recommended, because the disease-related survival in these patients is improved following surgery. In addition, duodenal gastrinomas, which occur more frequently in patients with MEN 1, have been treated successfully with surgery. However, in most patients with MEN 1, gastrinomas are multiple or extrapancreatic, and with the exception of duodenal gastrinomas, surgery is rarely successful. For example, the results of one study revealed that only ~15% of patients with MEN 1 were free of disease immediately after surgery, and at 5 years, this number had decreased to ~5%; the respective outcomes in patients without MEN 1 were better, at 45% and 40%. Given these findings, most specialists recommend a nonsurgical management for gastrinomas in MEN 1, except as noted earlier for smaller, isolated lesions. Treatment of disseminated gastrinomas is difficult. Chemotherapy with streptozotocin and 5-fluorouracil; hormonal therapy with octreotide or lanreotide, which are human somatostatin analogues; hepatic artery embolization; administration of human leukocyte interferon; and removal of all resectable tumor have been successful in some patients.

Insulinoma These β islet cell insulin-secreting tumors represent 10–30% of all pancreatic tumors in patients with MEN 1. Patients with an insulinoma present with hypoglycemic symptoms (e.g., weakness, headaches, sweating, faintness, seizures, altered behavior, weight gain) that typically develop after fasting or exertion and improve after glucose intake. The most reliable test is a supervised 72-h fast. Biochemical investigations reveal increased plasma insulin concentrations in association with hypoglycemia (Table 408-3). Circulating concentrations of C peptide and proinsulin, which are also increased, are useful in establishing the diagnosis. It also is important to demonstrate the absence of sulfonylureas in plasma and urine samples obtained during the investigation of hypoglycemia (Table 408-3). Surgical success is greatly enhanced by preoperative localization by endoscopic ultrasonography, CT scanning, or celiac axis angiography. Additional localization methods may include preoperative and perioperative percutaneous transhepatic portal venous sampling, selective intraarterial stimulation with hepatic venous sampling, and intraoperative direct pancreatic ultrasonography. Insulinomas occur in association with gastrinomas in 10% of patients with MEN 1, and the two tumors may arise at different times. Insulinomas occur more often in patients with MEN 1 who are younger than 40 years, and some arise in individuals younger than 20 years. In contrast, in patients without MEN 1, insulinomas generally occur in those older than 40 years. Insulinomas may be the first manifestation of MEN 1 in 10% of patients, and approximately 4% of patients with insulinomas will have MEN 1.

TABLE 408-3 BIOCHEMICAL AND RADIOLOGICAL SCREENING IN MULTIPLE ENDOCRINE NEOPLASIA TYPE 1

Tumor	Age to Begin (Years)	Biochemical Test (Plasma or Serum) Annually	Imaging Test (Time Interval)
Parathyroid	8	Calcium, PTH	None
Pancreatic NETs			
Gastrinoma	20	Gastrin (\pm gastric pH)	None
Insulinoma	5	Fasting glucose, insulin	None
Other pancreatic NET	<10	Chromogranin A; pancreatic polypeptide, glucagon, vasoactive intestinal peptide	MRI, CT, or EUS (annually)
Anterior pituitary	5	Prolactin, IGF-I	MRI (every 3 years)
Adrenal	<10	None unless symptoms or signs of functioning tumor and/or tumor >1 cm identified on imaging	MRI or CT (annually with pancreatic imaging)
Thymic and bronchial carcinoid	15	None	CT or MRI (every 1–2 years)

Abbreviations: CT, computed tomography; EUS, endoscopic ultrasound; IGF-I, insulin-like growth factor I; MRI, magnetic resonance imaging; PTH, parathyroid hormone.

Source: Reproduced from RV Thakker et al. *J Clin Endocrinol Metab* 97:2990, 2012.

TREATMENT **INSULINOMA**

Medical treatment, which consists of frequent carbohydrate meals and diazoxide or octreotide, is not always successful, and surgery is the optimal treatment. Surgical treatment, which ranges from enucleation of a single tumor to a distal pancreatectomy or partial pancreatectomy, has been curative in many patients. Chemotherapy may include streptozotocin, 5-fluorouracil, and doxorubicin. Hepatic artery embolization has been used for metastatic disease.

Glucagonoma These glucagon-secreting pancreatic NETs occur in <3% of patients with MEN 1. The characteristic clinical manifestations of a skin rash (necrolytic migratory erythema), weight loss, anemia, and stomatitis may be absent. The tumor may have been detected in an asymptomatic patient with MEN 1 undergoing pancreatic imaging or by the finding of glucose intolerance and hyperglucagonemia.

TREATMENT **GLUCAGONOMA**

Surgical removal of the glucagonoma is the treatment of choice. However, treatment may be difficult because approximately 50–80% of patients have metastases at the time of diagnosis. Medical treatment with somatostatin analogues (e.g., octreotide or lanreotide) or chemotherapy with streptozotocin and 5-fluorouracil has been successful in some patients, and hepatic artery embolization has been used to treat metastatic disease.

Vasoactive Intestinal Peptide (VIP) Tumors (VIPomas) VIPomas have been reported in only a few patients with MEN 1. This clinical syndrome is characterized by watery diarrhea, hypokalemia, and achlorhydria and is also referred to as the Verner-Morrison syndrome, the WDHA (watery diarrhea, hypokalemia, and achlorhydria) syndrome, or the VIPoma syndrome. The diagnosis is established by excluding laxative and diuretic abuse, by confirming a stool volume in excess of 0.5–1.0 L/d during a fast, and by documenting a markedly increased plasma VIP concentration.

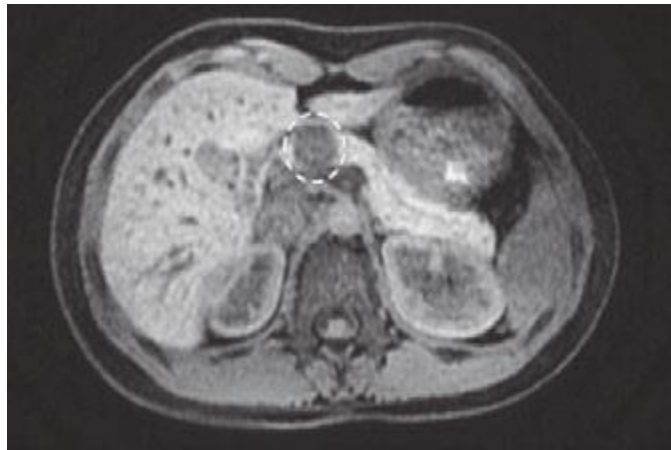
TREATMENT **VIPOMAS**

Surgical management of VIPomas, which are mostly located in the tail of the pancreas, can be curative. However, in patients with unresectable tumor, somatostatin analogues, such as octreotide and lanreotide, may be effective. Streptozotocin with 5-fluorouracil may be beneficial, along with hepatic artery embolization for the treatment of metastases.

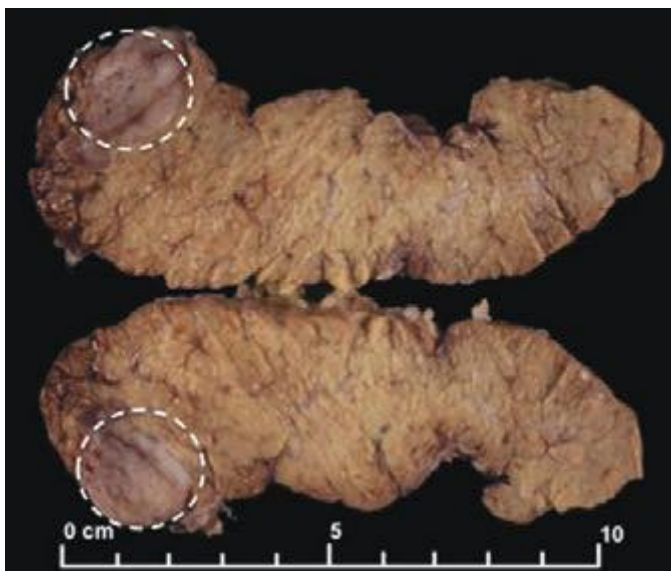
Pancreatic Polypeptide-Secreting Tumors (Ppomas) and Nonfunctioning Pancreatic NETs PPomas are found in a large number of patients with MEN 1. No pathologic sequelae of excessive polypeptide (PP) secretion are apparent, and the clinical significance of PP is unknown. Many PPomas may have been unrecognized or classified as nonfunctioning pancreatic NETs, which likely represent the most common enteropancreatic NET associated with MEN 1 (Fig. 408-1). The absence of both a clinical syndrome and specific biochemical abnormalities may result in a delayed diagnosis of nonfunctioning pancreatic NETs, which are associated with a worse prognosis than other functioning tumors, including insulinoma and gastrinoma. The optimum screening method and its timing interval for nonfunctioning pancreatic NETs remain to be established. At present, endoscopic ultrasound likely represents the most sensitive method of detecting small pancreatic tumors, but somatostatin receptor scintigraphy is the most reliable method for detecting metastatic disease (Table 408-3).

TREATMENT **PPOMAS AND NONFUNCTIONING PANCREATIC NETS**

The management of nonfunctioning pancreatic NETs in the asymptomatic patient is controversial. One recommendation is to undertake surgery irrespective of tumor size after biochemical



A



B

FIGURE 408-1 Pancreatic nonfunctioning neuroendocrine tumor (NET) in a 14-year-old patient with multiple endocrine neoplasia type 1 (MEN 1). **A.** An abdominal magnetic resonance imaging scan revealed a low-intensity >2.0 cm (anteroposterior maximal diameter) tumor within the neck of pancreas. There was no evidence of invasion of adjacent structures or metastases. The tumor is indicated by *white dashed circle*. **B.** The pancreatic NET was removed by surgery, and macroscopic examination confirmed the location of the tumor (*white dashed circles*) in the neck of the pancreas. Immunohistochemistry showed the tumor to immunostain for chromogranin A, but not gastrointestinal peptides or menin, thereby confirming that it was a non-secreting NET due to loss of menin expression. (Part A adapted with permission from PJ Newey et al: *J Clin Endocrinol Metab* 10:3640, 2009.)

assessment is complete. Alternatively, other experts recommend surgery based on tumor size, using either >1 cm or >3 cm at different centers. Pancreatoduodenal surgery is successful in removing the tumors in 80% of patients, but more than 40% of patients develop complications, including diabetes mellitus, frequent steatorrhea, early and late dumping syndromes, and other gastrointestinal symptoms. However, ~50–60% of patients treated surgically survive >5 years. When considering these recommendations, it is important to consider that occult metastatic disease (e.g., tumors not detected by imaging investigations) is likely to be present in a substantial proportion of these patients at the time of presentation. Inhibitors of tyrosine kinase receptors (TKRs) and of the mammalian

target of rapamycin (mTOR) signaling pathway have been reported to be effective in treating pancreatic NETs and in doubling the progression-free survival time.

Other Pancreatic NETs NETs secreting growth hormone-releasing hormone (GHRH), GHRHomas, have been reported rarely in patients with MEN 1. It is estimated that ~33% of patients with GHRHomas have other MEN 1-related tumors. GHRHomas may be diagnosed by demonstrating elevated serum concentrations of growth hormone and GHRH. More than 50% of GHRHomas occur in the lung, 30% occur in the pancreas, and 10% are found in the small intestine. Somatostatinomas secrete somatostatin, a peptide that inhibits the secretion of a variety of hormones, resulting in hyperglycemia, cholelithiasis, low acid output, steatorrhea, diarrhea, abdominal pain, anemia, and weight loss. Although 7% of pancreatic NETs secrete somatostatin, the clinical features of somatostatinoma syndrome are unusual in patients with MEN 1.

Pituitary Tumors (See also Chap. 403) Pituitary tumors occur in 15–50% of patients with MEN 1 (Table 408-1). These occur as early as 5 years of age or as late as the ninth decade. MEN 1 pituitary adenomas are more frequent in women than men and significantly are macroadenomas (i.e., diameter >1 cm). Moreover, about one-third of these pituitary tumors show invasive features such as infiltration of tumor cells into surrounding normal juxtatumoral pituitary tissue. However, no specific histologic parameters differentiate between MEN 1 and non-MEN 1 pituitary tumors. Approximately 60% of MEN 1-associated pituitary tumors secrete prolactin, <25% secrete growth hormone, 5% secrete adrenocorticotrophic hormone (ACTH), and the remainder appear to be nonfunctioning, with some secreting glycoprotein subunits (Table 408-1). However, pituitary tumors derived from MEN 1 patients may exhibit immunoreactivity to several hormones. In particular, there is a greater frequency of somatotrophic tumors. Prolactinomas are the first manifestation of MEN 1 in ~15% of patients, whereas somatotrophic tumors occur more often in patients older than 40 years of age. Fewer than 3% of patients with anterior pituitary tumors will have MEN 1. Clinical manifestations are similar to those in patients with sporadic pituitary tumors without MEN 1 and depend on the hormone secreted and the size of the pituitary tumor. Thus, patients may have symptoms of hyperprolactinemia (e.g., amenorrhea, infertility, and galactorrhea in women, or impotence and infertility in men) or have features of acromegaly or Cushing's disease. In addition, enlarging pituitary tumors may compress adjacent structures such as the optic chiasm or normal pituitary tissue, causing visual disturbances and/or hypopituitarism. In asymptomatic patients with MEN 1, periodic biochemical monitoring of serum prolactin and insulin-like growth factor I (IGF-I) levels, as well as MRI of the pituitary, can lead to early identification of pituitary tumors (Table 408-3). In patients with abnormal results, hypothalamic-pituitary testing should characterize the nature of the pituitary lesion and its effects on the secretion of other pituitary hormones.

TREATMENT PITUITARY TUMORS

Treatment of pituitary tumors in patients with MEN 1 consists of therapies similar to those used in patients without MEN 1 and includes appropriate medical therapy (e.g., bromocriptine or cabergoline for prolactinoma; or octreotide or lanreotide for somatotrophic tumors) or selective transsphenoidal adenectomy, if feasible, with radiotherapy reserved for residual unresectable tumor tissue. Pituitary tumors in MEN 1 patients may be more aggressive and less responsive to medical or surgical treatments.

Associated Tumors Patients with MEN 1 may also develop carcinoid tumors, adrenal cortical tumors, facial angiofibromas, collagenomas, thyroid tumors, and lipomatous tumors.

Carcinoid Tumors (See also Chap. 113) Carcinoid tumors occur in more than 3% of patients with MEN 1 (Table 408-1). The carcinoid tumor

may be located in the bronchi, gastrointestinal tract, pancreas, or thymus. At the time of diagnosis, most patients are asymptomatic and do not have clinical features of the carcinoid syndrome. Importantly, no hormonal or biochemical abnormality (e.g., plasma chromogranin A) is consistently observed in individuals with thymic or bronchial carcinoid tumors. Thus, screening for these tumors is dependent on radiologic imaging. The optimum method for screening has not been established. CT and MRI are sensitive for detecting thymic and bronchial tumors (Table 408-3), although repeated CT scanning raises concern about exposure to repeated doses of ionizing radiation. Octreotide scintigraphy may also reveal some thymic and bronchial carcinoids, although there is insufficient evidence to recommend its routine use. Gastric carcinoids, of which the type II gastric enterochromaffin-like (ECL) cell carcinoids (ECLomas) are associated with MEN 1 and Zollinger-Ellison syndrome, may be detected incidentally at the time of gastric endoscopy for dyspeptic symptoms in MEN 1 patients. These tumors, which may be found in >10% of MEN 1 patients, are usually multiple and smaller than 1.5 cm. Bronchial carcinoids in patients with MEN 1 occur predominantly in women (male-to-female ratio, 1:4). In contrast, thymic carcinoids in European patients with MEN 1 occur predominantly in men (male-to-female ratio, 20:1), with cigarette smokers having a higher risk for these tumors; thymic carcinoids in Japanese patients with MEN 1 have a less marked sex difference (male-to-female ratio 2:1). The course of thymic carcinoids in MEN 1 appears to be particularly aggressive. The presence of thymic tumors in patients with MEN 1 is associated with a median survival after diagnosis of approximately 9.5 years, with 70% of patients dying as a direct result of the tumor.

TREATMENT CARCINOID TUMORS

If resectable, surgical removal of carcinoid tumors is the treatment of choice. For unresectable tumors and those with metastatic disease, treatment with radiotherapy or chemotherapeutic agents (e.g., cisplatin, etoposide) may be used. In addition, somatostatin analogues, such as octreotide or lanreotide, have resulted in symptom improvement and regression of some tumors. Little is known about the malignant potential of gastric type II ECLomas, but treatment with somatostatin analogues, such as octreotide or lanreotide, has resulted in regression of these ECLomas.

Adrenocortical Tumors (See also Chap. 406) Asymptomatic adrenocortical tumors occur in 20–70% of patients with MEN 1 depending on the radiologic screening methods used (Table 408-1). Most of these tumors, which include cortical adenomas, hyperplasia, multiple adenomas, nodular hyperplasia, cysts, and carcinomas, are nonfunctioning. Indeed, <10% of patients with enlarged adrenal glands have hormonal hypersecretion, with primary hyperaldosteronism and ACTH-independent Cushing's syndrome being encountered most commonly. Occasionally, hyperandrogenemia may occur in association with adrenocortical carcinoma. Pheochromocytoma in association with MEN 1 is rare. Biochemical investigation (e.g., plasma renin and aldosterone concentrations, low-dose dexamethasone suppression test, urinary catecholamines, and/or metanephrines) should be undertaken in those with symptoms or signs suggestive of functioning adrenal tumors or in those with tumors >1 cm. Adrenocortical carcinoma occurs in approximately 1% of MEN 1 patients but increases to >10% for adrenal tumors larger than 1 cm.

TREATMENT ADRENOCORTICAL TUMORS

Consensus has not been reached about the management of MEN 1-associated nonfunctioning adrenal tumors, because the majority are benign. However, the risk of malignancy increases with size, particularly for tumors with a diameter >4 cm. Indications for surgery for adrenal tumors include: size >4 cm in diameter; atypical or suspicious radiologic features (e.g., increased Hounsfield unit on unenhanced CT scan) and size of 1–4 cm in diameter; or significant

2340 measurable growth over a 6-month period. The treatment of functioning (e.g., hormone-secreting) adrenal tumors is similar to that for tumors occurring in non-MEN 1 patients.

Meningioma Central nervous system (CNS) tumors, including ependymomas, schwannomas, and meningiomas, have been reported in MEN 1 patients (Table 408-1). Meningiomas are found in <10% of patients with other clinical manifestations of MEN 1 (e.g., primary hyperparathyroidism) for >15 years. The majority of meningiomas are not associated with symptoms, and 60% do not enlarge. The treatment of MEN 1-associated meningiomas is similar to that in non-MEN 1 patients.

Lipomas Subcutaneous lipomas occur in >33% of patients with MEN 1 (Table 408-1) and are frequently multiple. In addition, visceral, pleural, or retroperitoneal lipomas may occur in patients with MEN 1. Management is conservative. However, when surgically removed for cosmetic reasons, they typically do not recur.

Facial Angiofibromas and Collagenomas The occurrence of multiple facial angiofibromas in patients with MEN 1 may range from >20 to >90%, and occurrence of collagenomas may range from 0 to >70% (Table 408-1). These cutaneous findings may allow presymptomatic diagnosis of MEN 1 in the relatives of a patient with MEN 1. Treatment for these cutaneous lesions is usually not required.

Thyroid Tumors Thyroid tumors, including adenomas, colloid goiters, and carcinomas, have been reported to occur in >25% of patients with MEN 1. However, the prevalence of thyroid disorders in the general population is high, and it has been suggested that the association of thyroid abnormalities in patients with MEN 1 may be incidental. The treatment of thyroid tumors in MEN 1 patients is similar to that for non-MEN 1 patients.

Genetics and Screening The *MEN1* gene is located on chromosome 11q13 and consists of 10 exons, which encode a 610-amino acid protein, menin, that regulates transcription, genome stability, cell division, and proliferation. The pathophysiology of MEN 1 follows the Knudson two-hit hypothesis with a tumor-suppressor role for menin. Inheritance of a germline *MEN1* mutation predisposes an individual to developing a tumor that arises following a somatic mutation, which may be a point mutation or more commonly a deletion, leading to loss of heterozygosity (LOH) in the tumor DNA. The germline mutations of the *MEN1* gene are scattered throughout the entire 1830-bp coding region and splice sites, and there is no apparent correlation between the location of *MEN1* mutations and clinical manifestations of the disorder, in contrast with the situation in patients with MEN 2 (Table 408-1). More than 10% of *MEN1* germline mutations arise de novo and may be transmitted to subsequent generations. Some families with MEN 1 mutations develop parathyroid tumors as the sole endocrinopathy, and this condition is referred to as familial isolated hyperparathyroidism (FIHP). However, between 5 and 25% of patients with MEN 1 do not harbor germline mutations or deletions of the *MEN1* gene. Such patients with MEN 1-associated tumors but without *MEN1* mutations may represent phenocopies or have mutations involving other genes. Other genes associated with MEN 1-like features include: *CDC73*, which encodes parafibromin, whose mutations result in the hyperparathyroid-jaw tumor syndrome; the calcium-sensing receptor gene (*CaSR*), whose mutations result in familial benign hypocalcemic hypercalcemia (FBHH); and the aryl hydrocarbon receptor interacting protein gene (*AIP*), a tumor suppressor located on chromosome 11q13 whose mutations are associated with familial isolated pituitary adenomas (FIPA). Genetic testing to determine the *MEN1* mutation status in symptomatic family members within a MEN 1 kindred, as well as to all index cases (e.g., patients) with two or more endocrine tumors, is advisable. If an *MEN1* mutation is not identified in the index case with two or more endocrine tumors, then clinical and genetic tests for other disorders such as hyperparathyroid-jaw tumor syndrome, FBHH, FIPA, MEN 2, or MEN 4 should be considered, because these patients may represent phenocopies for MEN 1.

The current guidelines recommend that *MEN1* mutational analysis should be undertaken in: (1) an index case with two or more MEN 1-associated endocrine tumors (e.g., parathyroid, pancreatic, or pituitary tumors); (2) asymptomatic first-degree relatives of a known *MEN1* mutation carrier; and (3) first-degree relatives of a *MEN1* mutation carrier with symptoms, signs, or biochemical or radiologic evidence for one or more MEN 1-associated tumors. In addition, *MEN1* mutational analysis should be considered in patients with suspicious or atypical MEN 1. This would include individuals with parathyroid adenomas before the age of 30 years or multigland parathyroid disease; individuals with gastrinoma or multiple pancreatic NETs at any age; or individuals who have two or more MEN 1-associated tumors that are not part of the classical triad of parathyroid, pancreatic islet, and anterior pituitary tumors (e.g., parathyroid tumor plus adrenal tumor). Family members, including asymptomatic individuals who have been identified to harbor a *MEN1* mutation, will require biochemical and radiologic screening (Table 408-3). In contrast, relatives who do not harbor the *MEN1* mutation have a risk of developing MEN 1-associated endocrine tumors that is similar to that of the general population; thus, relatives without the *MEN1* mutation do not require repeated screening.

Mutational analysis in asymptomatic individuals should be undertaken at the earliest opportunity and, if possible, in the first decade of life because tumors have developed in some children by the age of 5 years. Appropriate biochemical and radiologic investigations (Table 408-3) aimed at detecting the development of tumors should then be undertaken in affected individuals. Mutant gene carriers should undergo biochemical screening at least once per annum and also have baseline pituitary and abdominal imaging (e.g., MRI or CT), which should then be repeated at 1- to 3-year intervals (Table 408-3). Screening should commence after 5 years of age and should continue for life because the disease may develop as late as the eighth decade. The screening history and physical examination elicit the symptoms and signs of hypercalcemia, nephrolithiasis, peptic ulcer disease, neuroglycopenia, hypopituitarism, galactorrhea and amenorrhea in women, acromegaly, Cushing's disease, and visual field loss and the presence of subcutaneous lipomas, angiofibromas, and collagenomas. Biochemical screening should include measurements of serum calcium, PTH, gastrointestinal hormones (e.g., gastrin, insulin with a fasting glucose, glucagon, VIP, PP), chromogranin A, prolactin, and IGF-I in all individuals. More specific endocrine function tests should be undertaken in individuals who have symptoms or signs suggestive of a specific clinical syndrome. Biochemical screening for the development of MEN 1 tumors in asymptomatic members of families with MEN 1 is of great importance to reduce morbidity and mortality from the associated tumors.

MULTIPLE ENDOCRINE NEOPLASIA TYPE 2 AND TYPE 3

Clinical Manifestations MEN type 2 (MEN 2), which is also called Sipple's syndrome, is characterized by the association of medullary thyroid carcinoma (MTC), pheochromocytomas, and parathyroid tumors (Table 408-1). Three clinical variants of MEN 2 are recognized: MEN 2A, MEN 2B, and MTC only. MEN 2A, which is often referred to as MEN 2, is the most common variant. In MEN 2A, MTC is associated with pheochromocytomas in 50% of patients (may be bilateral) and with parathyroid tumors in 20% of patients. MEN 2A may rarely occur in association with Hirschsprung's disease, caused by the absence of autonomic ganglion cells in the terminal hindgut, resulting in colonic dilatation, severe constipation, and obstruction. MEN 2A may also be associated with cutaneous lichen amyloidosis, which is a pruritic lichenoid lesion that is usually located on the upper back. MEN 2B, which is also referred to as MEN 3, represents 5% of all cases of MEN 2 and is characterized by the occurrence of MTC and pheochromocytoma in association with a Marfanoid habitus; mucosal neuromas of the lips, tongue, and eyelids; medullated corneal fibers; and intestinal autonomic ganglion dysfunction leading to multiple diverticulae and megacolon. Parathyroid tumors do not usually occur in MEN 2B. MTC only (FMTC) is a variant in which MTC is the sole manifestation of the syndrome. However, the distinction between

TABLE 408-4 RECOMMENDATIONS FOR TESTS AND SURGERY IN MEN 2 AND MEN 3^a

RET Mutation, Exon (Ex) Location, and Codon Involved	Risk ^b	Recommended Age (years) for Test/Intervention				
		RET Mutational Analysis	First Serum Calcitonin and Neck Ultrasound	Prophylactic Thyroidectomy	Screening for Pheochromocytoma	Screening for PHPT
Ex13 (768, 790) ^c ; Ex14 (804) ^c ; Ex15 (891) ^c	+	<3–5	<3–5	5 ^d	20	20
Ex10 (609, 611, 618, 620) ^c ; Ex11 (630) ^c	++	<3–5	<3–5	<5 ^e	20	20
Ex11 (634) ^c	+++	<3–5	<3–5	<5	8	20
Ex15 (883) ^c ; Ex16 (918) ^f	++++	ASAP and by <1	ASAP and by <0.5–1	ASAP and by <1	8	— ^g

^aAdapted from American Thyroid Association Guidelines, RT Kloos et al: *Thyroid* 6:565, 2009. ^bRisk for early development of metastasis and aggressive growth of medullary thyroid cancer: +++++, highest; +++, high; ++, intermediate; and +, lowest. ^cMutations associated with MEN 2A (or medullary thyroid carcinoma only). ^dConsider surgery at 5 years or later if serum calcitonin is normal, neck ultrasound is normal, and there is a less aggressive family history and family preference. ^eConsider surgery before 5 years or later if serum calcitonin is normal, neck ultrasound is normal, and there is a less aggressive family history and family preference. ^fMutations associated with MEN 2B (MEN 3). ^gNot required because PHPT is not a feature of MEN 2B (MEN 3).

Abbreviations: ASAP, as soon as possible; MEN, multiple endocrine neoplasia; PHPT, primary hyperparathyroidism.

FMTC and MEN 2A is difficult and should only be considered if there are at least four family members above the age of 50 years who are affected by MTC but not pheochromocytomas or primary hyperparathyroidism. All of the MEN 2 variants are due to mutations of the rearranged during transfection (*RET*) protooncogene, which encodes a TKR. Moreover, there is a correlation between the locations of *RET* mutations and MEN 2 variants. Thus, ~95% of MEN 2A patients have mutations involving the cysteine-rich extracellular domain, with mutations of codon 634 accounting for ~85% of MEN 2A mutations; FMTC patients also have mutations of the cysteine-rich extracellular domain, with most mutations occurring in codon 618. In contrast, ~95% of MEN 2B/MEN 3 patients have mutations of codon 918 of the intracellular tyrosine kinase domain (Table 408-1 and Table 408-4).

Medullary Thyroid Carcinoma MTC is the most common feature of MEN 2A and MEN 2B and occurs in almost all affected individuals. MTC represents 5–10% of all thyroid gland carcinomas, and 20% of MTC patients have a family history of the disorder. The use of *RET* mutational analysis to identify family members at risk for hereditary forms of MTC has altered the presentation of MTC from that of symptomatic tumors to a preclinical disease for which prophylactic thyroidectomy (Table 408-4) is undertaken to improve the prognosis and ideally result in cure. However, in patients who do not have a known family history of MEN 2A, FMTC, or MEN 2B, and therefore have not had *RET* mutational analysis, MTC may present as a palpable mass in the neck, which may be asymptomatic or associated with symptoms of pressure or dysphagia in >15% of patients. Diarrhea occurs in 30% of patients and is associated either with elevated circulating concentrations of calcitonin or tumor-related secretion of serotonin and prostaglandins. Some patients may also experience flushing. In addition, ectopic ACTH production by MTC may cause Cushing's syndrome. The diagnosis of MTC relies on the demonstration of hypercalcitoninemia (>90 pg/mL in the basal state); stimulation tests using IV pentagastrin (0.5 mg/kg) and or calcium infusion (2 mg/kg) are rarely used now, reflecting improvements in the assay for calcitonin. Neck ultrasonography with fine-needle aspiration of the nodules can confirm the diagnosis. Radionuclide thyroid scans may reveal MTC tumors as “cold” nodules. Radiography may reveal dense irregular calcification within the involved portions of the thyroid gland and in lymph nodes involved with metastases. Positron emission tomography (PET) may help to identify the MTC and metastases (Fig. 408-2). Metastases of MTC usually occur to the cervical lymph nodes in the early stages and to the mediastinal nodes, lung, liver, trachea, adrenal, esophagus, and bone in later stages. Elevations in serum calcitonin concentrations are often the first sign of recurrence or persistent disease, and the serum calcitonin doubling time is useful for determining prognosis. MTC can have an aggressive clinical course, with early metastases and death in approximately 10% of patients. A family history of aggressive MTC or MEN 2B may be elicited.

TREATMENT MEDULLARY THYROID CARCINOMA

Individuals with *RET* mutations who do not have clinical manifestations of MTC should be offered prophylactic surgery between the ages of <1 and 5 years. The timing of surgery will depend on the type of *RET* mutation and its associated risk for early development, metastasis, and aggressive growth of MTC (Table 408-4). Such patients should have a total thyroidectomy with a systematic central neck dissection to remove occult nodal metastasis, although

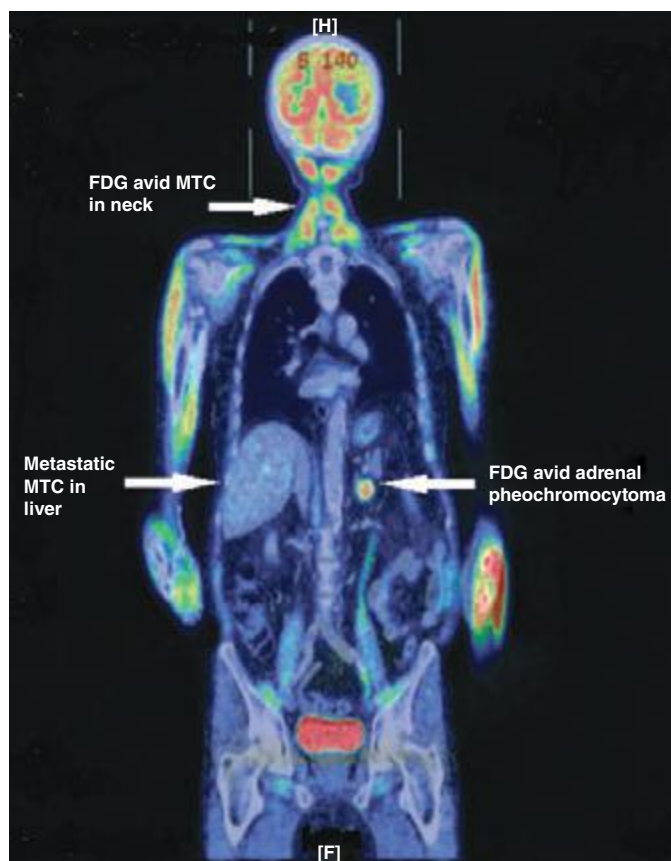


FIGURE 408-2 Fluorodeoxyglucose (FDG) positron emission tomography scan in a patient with multiple endocrine neoplasia type 2A, showing medullary thyroid cancer (MTC) with hepatic and skeletal (left arm) metastasis and a left adrenal pheochromocytoma. Note the presence of excreted FDG compound in the bladder. (Reproduced with permission from A Naziat et al: *Clin Endocrinol [Oxf]* 78:966, 2013.)


2342 the value of undertaking a central neck dissection has been subject to debate. Prophylactic thyroidectomy, with life-long thyroxine replacement, has dramatically improved outcomes in patients with MEN 2 and MEN 3, such that ~90% of young patients with *RET* mutations who had a prophylactic thyroidectomy have no evidence of persistent or recurrent MTC at 7 years after surgery. In patients with clinically evident MTC, a total thyroidectomy with bilateral central resection is recommended, and an ipsilateral lateral neck dissection should be undertaken if the primary tumor is >1 cm in size or there is evidence of nodal metastasis in the central neck. Surgery is the only curative therapy for MTC. The 10-year survival in patients with metastatic MTC is ~20%. For inoperable MTC or metastatic disease, the tyrosine kinase inhibitors, vandetanib and cabozantinib, have improved the progression-free survival times. Other types of chemotherapy are of limited efficacy, but radiotherapy may help to palliate local disease.

Pheochromocytoma (See also Chap. 407) These noradrenaline- and adrenaline-secreting tumors occur in >50% of patients with MEN 2A and MEN 2B and are a major cause of morbidity and mortality. Patients may have symptoms and signs of catecholamine secretion (e.g., headaches, palpitations, sweating, poorly controlled hypertension), or they may be asymptomatic with detection through biochemical screening based on a history of familial MEN 2A, MEN 2B, or MTC. Pheochromocytomas in patients with MEN 2A and MEN 2B differ significantly in distribution when compared with patients without MEN 2A and MEN 2B. Extra-adrenal pheochromocytomas, which occur in 10% of patients without MEN 2A and MEN 2B, are observed rarely in patients with MEN 2A and MEN 2B. Malignant pheochromocytomas are much less common in patients with MEN 2A and MEN 2B. The biochemical and radiologic investigation of pheochromocytoma in patients with MEN 2A and MEN 2B is similar to that in non-MEN 2 patients and includes the measurement of plasma (obtained from supine patients) and urinary free fractionated metanephrines (e.g., normetanephrine and metanephrines measured separately), CT or MRI scanning, radionuclide scanning with meta-iodo-(¹²³I or ¹³¹I)-benzyl guanidine (MIBG), and PET using (¹⁸F)-fluorodopamine or (¹⁸F)-fluoro-2-dexoxy-D-glucose (Fig. 408-2).

TREATMENT PHEOCHROMOCYTOMA

Surgical removal of pheochromocytoma, using α and β adreno-receptor blockade before and during the operation, is the recommended treatment. Endoscopic adrenal-sparing surgery, which decreases postoperative morbidity, hospital stay, and expense, as opposed to open surgery, has become the method of choice.

Parathyroid Tumors (See also Chap. 424) Parathyroid tumors occur in 10–25% of patients with MEN 2A. However, >50% of these patients do not have hypercalcemia. The presence of abnormally enlarged parathyroids, which are unusually hyperplastic, is often seen in the normocalcemic patient undergoing thyroidectomy for MTC. The biochemical investigation and treatment of hypercalcemic patients with MEN 2A is similar to that of patients with MEN 1.


 **Genetics and Screening** To date, approximately 50 different *RET* mutations have been reported, and these are located in exons 5, 8, 10, 11, 13, 14, 15, and 16. *RET* germline mutations are detected in >95% of MEN 2A, FMTC, and MEN 2B families, with Cys634Arg being most common in MEN 2A, Cys618Arg being most common in FMTC, and Met918Thr being most common in MEN 2B (Tables 408-1 and 408-4). Between 5 and 10% of patients with MTC or MEN 2A-associated tumors have de novo *RET* germline mutations, and ~50% of patients with MEN 2B have de novo *RET* germline mutations. These de novo *RET* germline mutations always occur on the paternal allele. Approximately 5% of patients with sporadic pheochromocytoma have a germline *RET* mutation, but such germline *RET* mutations do not appear to be associated with sporadic primary hyperparathyroidism.

Thus, *RET* mutational analysis should be performed in: (1) all patients with MTC who have a family history of tumors associated with MEN 2, FMTC, or MEN 3, such that the diagnosis can be confirmed and genetic testing offered to asymptomatic relatives; (2) all patients with MTC and pheochromocytoma without a known family history of MEN 2 or MEN 3; (3) all patients with MTC, but without a family history of MEN 2, FMTC, or MEN 3, because these patients may have a de novo germline *RET* mutations; (4) all patients with bilateral pheochromocytoma; and (5) patients with unilateral pheochromocytoma, particularly if this occurs with increased calcitonin levels.

Screening for MEN 2/MEN 3-associated tumors in patients with *RET* germline mutations should be undertaken annually and include serum calcitonin measurements, a neck ultrasound for MTC, plasma and 24-h urinary fractionated metanephrines for pheochromocytoma, and albumin-corrected serum calcium or ionized calcium with PTH for primary hyperparathyroidism. In patients with MEN 2-associated *RET* mutations, screening for MTC should begin by 3 to 5 years; for pheochromocytoma by 20 years; and for primary hyperparathyroidism by 20 years of age (Table 408-4).


MULTIPLE ENDOCRINE NEOPLASIA TYPE 4

Clinical Manifestations Patients with MEN 1-associated tumors, such as parathyroid adenomas, pituitary adenomas, and pancreatic NETs, occurring in association with gonadal, adrenal, renal, and thyroid tumors have been reported to have mutations of the gene encoding the 196-amino acid cyclin-dependent kinase inhibitor (CK1) p27 kip1 (*CDNK1B*). Such families with MEN 1-associated tumors and *CDNK1B* mutations are designated to have MEN 4 (Table 408-1). The investigations and treatments for the MEN 4-associated tumors are similar to those for MEN 1 and non-MEN 1 tumors.

 **Genetics and Screening** To date, eight different MEN 4-associated mutations of *CDNK1B*, which is located on chromosome 12p13, have been reported, and all of these are associated with a loss of function. These MEN 4 patients may represent ~3% of the 5–10% of patients with MEN 1 who do not have mutations of the *MEN1* gene. Germline *CDNK1B* mutations may rarely be found in patients with sporadic (i.e., nonfamilial) forms of primary hyperparathyroidism.

HYPERPARATHYROIDISM-JAW TUMOR SYNDROME (SEE ALSO CHAP. 424)

Clinical Manifestations Hyperparathyroidism-jaw tumor (HPT-JT) syndrome is an autosomal dominant disorder characterized by the development of parathyroid tumors (15% are carcinomas) and fibro-osseous jaw tumors. In addition, some patients may also develop Wilms' tumors, renal cysts, renal hematomas, renal cortical adenomas, papillary renal cell carcinomas, pancreatic adenocarcinomas, uterine tumors, testicular mixed germ cell tumors with a major seminoma component, and Hürthle cell thyroid adenomas. The parathyroid tumors may occur in isolation and without any evidence of jaw tumors, and this may cause confusion with other hereditary hypercalcemic disorders, such as MEN 1. However, genetic testing to identify the causative mutation will help to establish the correct diagnosis. The investigation and treatment for HPT-JT-associated tumors are similar to those in non-HPT-JT patients, except that early parathyroidectomy is advisable because of the increased frequency of parathyroid carcinoma.

 **Genetics and Screening** The gene that causes HPT-JT is located on chromosome 1q31.2 and encodes a 531-amino acid protein, parafibromin (Table 408-2). Parafibromin is also referred to as cell division cycle protein 73 (CDC73) and has a role in transcription. Genetic testing in families helps to identify mutation carriers who should be periodically screened for the development of tumors (Table 408-5).

VON HIPPEL-LINDAU DISEASE (SEE ALSO CHAP. 407)

Clinical Manifestations von Hippel-Lindau (VHL) disease is an autosomal dominant disorder characterized by hemangioblastomas of the retina and CNS; cysts involving the kidneys, pancreas, and epididymis; renal cell carcinomas; pheochromocytomas; and pancreatic islet cell tumors. The retinal and CNS hemangioblastomas are benign vascular tumors that may be multiple; those in the CNS may cause symptoms

TABLE 408-5 HPT-JT SCREENING GUIDELINES


Tumor ^a	Test	Frequency ^b
Parathyroid	Serum Ca, PTH	6–12 months
Ossifying jaw fibroma	Panoramic jaw x-ray with neck shielding ^c	5 years
Renal	Abdominal MRI ^{c,d}	5 years
Uterine	Ultrasound (transvaginal or transabdominal) and additional imaging ± D&C if indicated ^e	Annual

^aScreening for most common HPT-JT-associated tumors is considered. Assessment for other reported tumor types may be indicated (e.g., pancreatic, thyroid, testicular tumors). ^bFrequency of repeating test after baseline tests performed. ^cX-rays and imaging involving ionizing radiation should ideally be avoided to minimize risk of generating subsequent mutations. ^dUltrasound scan recommended if MRI unavailable. ^eSuch selective pelvic imaging should be considered after obtaining a detailed menstrual history.

Abbreviations: Ca, calcium; D&C, dilatation and curettage; HPT-JT, hyperparathyroidism-jaw tumor syndrome; MRI, magnetic resonance imaging; PTH, parathyroid hormone.

Source: Reproduced from PJ Newey et al: *Hum Mutat* 31:295, 2010.


by compressing adjacent structures and/or increasing intracranial pressure. In the CNS, the cerebellum and spinal cord are the most frequently involved sites. The renal abnormalities consist of cysts and carcinomas, and the lifetime risk of a renal cell carcinoma (RCC) in VHL is 70%. The endocrine tumors in VHL consist of pheochromocytomas and pancreatic islet cell tumors. The clinical presentation of pheochromocytoma in VHL disease is similar to that in sporadic cases, except there is a higher frequency of bilateral or multiple tumors, which may involve extra-adrenal sites in VHL disease. The most frequent pancreatic lesions in VHL are multiple cyst-adenomas, which rarely cause clinical disease. However, nonsecreting pancreatic islet cell tumors occur in <10% of VHL patients, who are usually asymptomatic. The pancreatic tumors in these patients are often detected by regular screening using abdominal imaging. Pheochromocytomas should be investigated and treated as described earlier for MEN 2. The pancreatic islet cell tumors frequently become malignant, and early surgery is recommended.

 **Genetics and Screening** The *VHL* gene, which is located on chromosome 3p26-p25, is widely expressed in human tissues and encodes a 213–amino acid protein (pVHL) (Table 408-2). A wide variety of germline *VHL* mutations have been identified. *VHL* acts as a tumor-suppressor gene. A correlation appears to exist between the type of mutation and the clinical phenotype; large deletions and protein-truncating mutations are associated with a low incidence of pheochromocytomas, whereas some missense mutations in VHL patients are associated with pheochromocytoma (referred to as VHL type 2C). Other missense mutations may be associated with hemangioblastomas and RCC but not pheochromocytoma (referred to as VHL type 1), whereas distinct missense mutations are associated with hemangioblastomas, RCC, and pheochromocytoma (VHL type 2B). VHL type 2A, which refers to the occurrence of hemangioblastomas and pheochromocytoma without RCC, is associated with rare missense mutations. The basis for these complex genotype-phenotype relationships remains to be elucidated. One major function of pVHL, which is also referred to as elongin, is to downregulate the expression of vascular endothelial growth factor (VEGF) and other hypoxia-inducible mRNAs. Thus, pVHL, in complex with other proteins, regulates the expression of hypoxia-inducible factors (HIF-1 and HIF-2) such that loss of functional pVHL leads to a stabilization of the HIF protein complexes, resulting in VEGF overexpression and tumor angiogenesis. Screening for the development of pheochromocytomas and pancreatic islet cell tumors is as described earlier for MEN 2 and MEN 1, respectively (Tables 408-3 and 408-4).

NEUROFIBROMATOSIS

Clinical Manifestations Neurofibromatosis type 1 (NF1), which is also referred to as von Recklinghausen's disease, is an autosomal dominant disorder characterized by the following manifestations: neurologic (e.g., peripheral and spinal neurofibromas); ophthalmologic (e.g., optic

gliomas and iris hamartomas such as Lisch nodules); dermatologic (e.g., café au lait macules); skeletal (e.g., scoliosis, macrocephaly, short stature, and pseudoarthrosis); vascular (e.g., stenoses of renal and intracranial arteries); and endocrine (e.g., pheochromocytoma, carcinoid tumors, and precocious puberty). Neurofibromatosis type 2 (NF2) is also an autosomal dominant disorder but is characterized by the development of bilateral vestibular schwannomas (acoustic neuromas) that lead to deafness, tinnitus, or vertigo. Some patients with NF2 also develop meningiomas, spinal schwannomas, peripheral nerve neurofibromas, and café au lait macules. Endocrine abnormalities are not found in NF2 and are associated solely with NF1. Pheochromocytomas, carcinoid tumors, and precocious puberty occur in about 1% of patients with NF1, and growth hormone deficiency has been also reported. The features of pheochromocytomas in NF1 are similar to those in non-NF1 patients, with 90% of tumors being located within the adrenal medulla and the remaining 10% at an extra-adrenal location, which often involves the para-aortic region. Primary carcinoid tumors are often periampullary and may also occur in the ileum but rarely in the pancreas, thyroid, or lungs. Hepatic metastases are associated with symptoms of the carcinoid syndrome, which include flushing, diarrhea, bronchoconstriction, and tricuspid valve disease. Precocious puberty is usually associated with the extension of an optic glioma into the hypothalamus with resultant early activation of gonadotropin-releasing hormone secretion. Growth hormone deficiency has also been observed in some NF1 patients, who may or may not have optic chiasmal gliomas, but it is important to note that short stature is frequent in the absence of growth hormone deficiency in patients with NF1. The investigation and treatment for tumors are similar to those undertaken for each respective tumor type in non-NF1 patients.

 **Genetics and Screening** The *NF1* gene, which is located on chromosome 17q11.2 and acts as a tumor suppressor, consists of 60 exons that span more than 350 kb of genomic DNA (Table 408-2). Mutations in *NF1* are of diverse types and are scattered throughout the exons. The NF1 gene product is the protein neurofibromin, which has homologies to the p120GAP (GTPase activating protein) and acts on p21ras by converting the active GTP bound form to its inactive GDP form. Mutations of *NF1* impair this downregulation of the p21ras signaling pathways, which in turn results in abnormal cell proliferation. Screening for the development of pheochromocytomas and carcinoid tumors is as described earlier for MEN 2 and MEN 1, respectively (Tables 408-3 and 408-4).

CARNEY COMPLEX

Clinical Manifestations Carney complex (CNC) is an autosomal dominant disorder characterized by spotty skin pigmentation (usually of the face, labia, and conjunctiva), myxomas (usually of the eyelids and heart, but also the tongue, palate, breast, and skin), psammomatous melanotic schwannomas (usually of the sympathetic nerve chain and upper gastrointestinal tract), and endocrine tumors that involve the adrenals, Sertoli cells, somatotropes, thyroid, and ovary. Cushing's syndrome, the result of primary pigmented nodular adrenal disease (PPNAD), is the most common endocrine manifestation of CNC and may occur in one-third of patients. Patients with CNC and Cushing's syndrome often have an atypical appearance by being thin (as opposed to having truncal obesity). In addition, they may have short stature, muscle and skin wasting, and osteoporosis. These patients often have levels of urinary free cortisol that are normal or increased only marginally. Cortisol production may fluctuate periodically with days or weeks of hypercortisolism; this pattern is referred to as "periodic Cushing's syndrome." Patients with Cushing's syndrome usually have loss of the circadian rhythm of cortisol production. Acromegaly, the result of a somatotrope tumor, affects ~10% of patients with CNC. Testicular tumors may also occur in one-third of patients with CNC. These may either be large-cell calcifying Sertoli cell tumors, adrenocortical rests, or Leydig cell tumors. The Sertoli cell tumors occasionally may be estrogen-secreting and lead to precocious puberty or gynecomastia. Some patients with CNC have been reported to develop thyroid follicular tumors, ovarian cysts, or breast duct adenomas.

Genetics and Screening CNC type 1 (CNC1) is due to mutations of the protein kinase A (PKA) regulatory subunit 1 α (R1 α) (*PPKARIA*), a tumor suppressor, whose gene is located on chromosome 17q.24.2 (Table 408-2). The gene causing CNC type 2 (CNC2) is located on chromosome 2p16 and has not yet been identified. It is interesting to note, however, that some tumors do not show LOH of 2p16 but instead show genomic instability, suggesting that this CNC gene may not be a tumor suppressor. Screening and treatment of these endocrine tumors are similar to those described earlier for patients with MEN 1 and MEN 2 (Tables 408-3 and 408-4).

COWDEN'S SYNDROME

Clinical Manifestations Multiple hamartomatous lesions, especially of the skin, mucous membranes (e.g., buccal, intestinal, and colonic), breast, and thyroid are characteristic of Cowden's (CWD) syndrome, which is an autosomal dominant disorder. Thyroid abnormalities occur in two-thirds of patients with CWD syndrome, and these usually consist of multinodular goiters or benign adenomas, although <10% of patients may have a follicular thyroid carcinoma. Breast abnormalities occur in >75% of patients and consist of either fibrocystic disease or adenocarcinomas. The investigation and treatment for CWD tumors are similar to those undertaken for non-CWD patients.

Genetics and Screening CWD syndrome is genetically heterogeneous, and six types (CWD1-6) are recognized (Table 408-2). CWD is due to mutations of the phosphate and tensin homologue deleted on chromosome 10 (*PTEN*) gene, located on chromosome 10q23.31. CWD2 is caused by mutations of the succinate dehydrogenase subunit B (*SDHB*) gene, located on chromosome 1p36.13; and CWD3 is caused by mutations of the *SDHD* gene, located on chromosome 11q13.1. *SDHB* and *SDHD* mutations are also associated with pheochromocytoma. CWD4 is caused by hypermethylation of the Killin (*KLLN*) gene, the promoter of which shares the same transcription site as *PTEN* on chromosome 10q23.31. CWD5 is caused by mutations of the phosphatidylinositol 3-kinase catalytic alpha (*PIK3CA*) gene on chromosome 3q26.32, and CWD6 is caused by mutations of the V-Akt murine thymoma viral oncogene homolog 1 (*AKT1*) gene on chromosome 14q32.33. Screening for thyroid abnormalities entails neck ultrasonography and fine-needle aspiration with analysis of cell cytology.

MCCUNE-ALBRIGHT SYNDROME (SEE ALSO CHAP. 426e)

Clinical Manifestations McCune-Albright syndrome (MAS) is characterized by the triad of polyostotic fibrous dysplasia, which may be associated with hypophosphatemic rickets; café au lait skin pigmentation; and peripheral precocious puberty; other endocrine abnormalities include thyrotoxicosis, which may be associated with a multinodular goiter, somatotrope tumors, and Cushing's syndrome (due to adrenal tumors). Investigation and treatment for each endocrinopathy are similar to those used in patients without MAS.

Genetics and Screening MAS is a disorder of mosaicism that results from postzygotic somatic cell mutations of the G protein a stimulating subunit (Gsa), encoded by the *GNAS1* gene, located on chromosome 20q13.32 (Table 408-2). The Gsa mutations, which include Arg201Cys, Arg201His, Glu227Arg, or Glu227His, are activating and are found only in cells of the abnormal tissues. Screening for hyperfunction of relevant endocrine glands and development of hypophosphatemia, which may be associated with elevated serum fibroblast growth factor 23 (FGF23) concentrations, is undertaken in MAS patients.

ACKNOWLEDGMENTS

The author is grateful to the Medical Research Council (UK) for support and to Mrs. Tracey Walker for typing the manuscript.

409 Autoimmune Polyendocrine Syndromes

Peter A. Gottlieb

Polyglandular deficiency syndromes have been given many different names, reflecting the wide spectrum of disorders that have been associated with these syndromes and the heterogeneity of their clinical presentations. The name used in this chapter for this group of disorders is *autoimmune polyendocrine syndrome* (APS). In general, these disorders are divided into two major categories, APS type 1 (APS-1) and APS type 2 (APS-2). Some groups have further subdivided APS-2 into APS type 3 (APS-3) and APS type 4 (APS-4) depending on the type of autoimmunity involved. For the most part, this additional classification does not clarify our understanding of disease pathogenesis or prevention of complications in individual patients. Importantly, there are many nonendocrine disease associations included in these syndromes, suggesting that although the underlying autoimmune disorder predominantly involves endocrine targets, it does not exclude other tissues. The disease associations found in APS-1 and APS-2 are summarized in Table 409-1. Understanding these syndromes and their disease manifestations can lead to early diagnosis and treatment of additional disorders in patients and their family members.

TABLE 409-1 DISEASE ASSOCIATIONS WITH AUTOIMMUNE POLYENDOCRINE SYNDROMES

Autoimmune Polyendocrine Syndrome Type 1	Autoimmune Polyendocrine Syndrome Type 2	Other Autoimmune Polyendocrine Disorders
Endocrine	Endocrine	IPEX (immune dysfunction polyendocrinopathy X-linked)
Addison's disease	Addison's disease	Thymic tumors
Hypoparathyroidism	Type 1 diabetes	Anti-insulin receptor antibodies
Hypogonadism	<i>Graves' disease or autoimmune thyroiditis</i>	POEMS syndrome
<i>Graves' disease or autoimmune thyroiditis</i>	<i>Hypogonadism</i>	Insulin autoimmune syndrome (Hirata's syndrome)
<i>Type 1 diabetes</i>		Adult combined pituitary hormone deficiency (CPHD) with anti-Pit1 autoantibodies
		Kearns-Sayre syndrome
		DIDMOAD syndrome
Nonendocrine	Nonendocrine	Congenital rubella associated with thyroiditis and/or diabetes
Mucocutaneous candidiasis	Celiac disease, dermatitis herpetiformis	
Chronic active hepatitis	Pernicious anemia	
Pernicious anemia	Vitiligo	
Vitiligo	<i>Alopecia</i>	
Asplenism	<i>Myasthenia gravis</i>	
Ectodermal dysplasia	<i>IgA deficiency</i>	
<i>Alopecia</i>	<i>Parkinson's disease</i>	
<i>Malabsorption syndromes</i>	<i>Idiopathic thrombocytopenia</i>	
<i>IgA deficiency</i>		

Abbreviations: DIDMOAD, diabetes insipidus, diabetes mellitus, progressive bilateral optic atrophy, and sensorineural deafness; POEMS, polyneuropathy, organomegaly, endocrinopathy, M-protein, and skin changes.

Note: Italics denote less common disorders.

APS-1

APS-1 (Online Mendelian Inheritance in Man [OMIM] 240300) has also been called autoimmune polyendocrinopathy–candidiasis–ectodermal dystrophy (APECED). Mucocutaneous candidiasis, hypoparathyroidism, and Addison’s disease form the three major components of this disorder. However, as summarized in Table 409-1, many other organ systems can be involved over time. APS-1 is rare, with fewer than 500 cases reported in the literature. It is an autosomal recessive disorder caused by mutations in the *AIRE* gene (autoimmune regulator gene) found on chromosome 21. This gene is most highly expressed in thymic medullary epithelial cells (mTECs) where it appears to control the expression of tissue-specific self-antigens (e.g., insulin). Deletion of this regulator leads to decreased expression of tissue-specific self-antigens and is hypothesized to allow autoreactive T cells to avoid clonal deletion, which normally occurs during T cell maturation in the thymus. The *AIRE* gene is also expressed in epithelial cells found in peripheral lymphoid organs, but its role in these extrathymic cells remains controversial. A number of mutations have been described in this gene, and there is a higher frequency within certain ethnic groups including Iranian Jews, Sardinians, Finns, Norwegians, and Irish.

Clinical Manifestations APS-1 develops very early in life, often in infancy (Table 409-2). Chronic mucocutaneous candidiasis without signs of systemic disease is often the first manifestation. It affects the mouth and nails more frequently than the skin and esophagus. Chronic oral candidiasis can result in atrophic disease with areas suggestive of leukoplakia, which can pose a risk for future carcinoma. The etiology is associated with anticytokine autoantibodies (anti-IL-17A, -IL-17F, and -IL-22) related to T helper (T_H) 17 T cells and depressed production of these cytokines by peripheral blood mononuclear cells. Hypoparathyroidism usually develops next, followed by adrenal insufficiency. The time from development of one component of the disorder to the next can be many years, and the order of disease appearance is variable.

Chronic candidiasis is nearly always present and is not very responsive to treatment. Hypoparathyroidism is found in >85% of cases, and Addison’s disease is found in nearly 80%. Gonadal failure appears to affect women more than men (70% vs 25%, respectively), and hypoplasia of the dental enamel also occurs frequently (77% of patients). Other endocrine disorders that occur less frequently include type 1 diabetes (23%) and autoimmune thyroid disease (18%). Nonendocrine manifestations that present less frequently include alopecia (40%), vitiligo (26%), intestinal malabsorption (18%), pernicious anemia (31%), chronic active hepatitis (17%), and nail dystrophy. An unusual and debilitating manifestation of the disorder is the development of refractory diarrhea/obstipation that may be related to autoantibody-mediated destruction of enterochromaffin or enterochromaffin-like cells.

TABLE 409-2 COMPARISON OF APS-1 AND APS-2

APS-1	APS-2
Early onset: infancy	Later onset
Siblings often affected and at risk	Multigenerational
Equivalent sex distribution	Females > males affected
Monogenic: <i>AIRE</i> gene, chromosome 21, autosomal recessive	Polygenic: <i>HLA</i> , <i>MICA</i> , <i>PTNP22</i> , <i>CTLA4</i>
Not HLA associated for entire syndrome, some specific component risk	DR3/DR4 associated; other HLA class III gene associations noted
Autoantibodies to type 1 interferons and IL-17 and IL-22	No autoantibodies to cytokines
Autoantibodies to specific target organs	Autoantibodies to specific target organs
Asplenia	No defined immunodeficiency
Mucocutaneous candidiasis	Association with other nonendocrine immunologic disorders like myasthenia gravis and idiopathic thrombocytopenic purpura

Abbreviations: APS, autoimmune polyendocrine syndrome; IL, interleukin.

The incidence rates for many of these disorders peak in the first or second decade of life, but the individual disease components continue to emerge over time. Therefore, prevalence rates may be higher than originally reported.

Diagnosis The diagnosis of APS-1 is usually made clinically when two of the three major component disorders are found in an individual patient. Siblings of individuals with APS-1 should be considered affected even if only one component disorder has been detected due to the known inheritance of the syndrome. Genetic analysis of the *AIRE* gene should be undertaken to identify mutations. Initial sequencing may detect the common mutations, but rare mutations are continually being noted, and an initial negative genetic analysis should not dissuade one from the clinical diagnosis until more extensive DNA sequencing can be performed. Detection of anti-interferon α and anti-interferon γ antibodies can identify nearly 100% of cases with APS-1. The autoantibody arises independent of the type of *AIRE* gene mutation and is not found in other autoimmune disorders.

Diagnosis of each underlying disorder should be done based on their typical clinical presentations (Table 409-3). Mucocutaneous candidiasis may present throughout the gastrointestinal tract, and it may be detected in the oral mucosa or from stool samples. Evaluation by a gastroenterologist to examine the esophagus for candidiasis or secondary stricture may be merited based on symptoms. Other gastrointestinal manifestations of APS-1, including malabsorption and obstipation, may also bring these young patients to the attention of gastroenterologists for first evaluation. Specific physical examination findings of hyperpigmentation, vitiligo, alopecia, tetany, and signs of hyper- or hypothyroidism should be considered as signs of development of component disorders.

The development of disease-specific autoantibody assays can help confirm disease and also detect risk for future disease. For example, where possible, detection of anticytokine antibodies to interleukin (IL) 17 and IL-22 would confirm the diagnosis of mucocutaneous candidiasis due to APS-1. The presence of anti-21-hydroxylase antibody or anti-17-hydroxylase antibody (which may be found more commonly in adrenal insufficiency associated with APS-1) would confirm the presence or risk for Addison’s disease. Other autoantibodies found in type 1 diabetes (e.g., anti-GAD65), pernicious anemia, and other component conditions should be screened for on a regular basis (6- to 12-month intervals depending on the age of the subject).

Laboratory tests, including a complete metabolic panel, phosphorous and magnesium, thyroid-stimulating hormone (TSH), adrenocorticotropic hormone (ACTH; morning), hemoglobin A_{1c}, plasma vitamin B₁₂ level, and complete blood count with peripheral smear looking for Howell-Jolly bodies (asplenia), should also be performed at these time points. Detection of abnormal physical findings or test results should prompt subsequent examinations of the relevant organ system (e.g., presence of Howell-Jolly bodies indicates need for ultrasound of spleen).

TREATMENT APS-1

Therapy of individual disease components is carried out as outlined in other relevant chapters. Replacement of deficient hormones (e.g., adrenal, pancreas, ovaries/testes) will treat most of the endocrinopathies noted. Several unique issues merit special emphasis. Adrenal insufficiency can be masked by primary hypothyroidism by prolonging the half-life of cortisol. The caveat therefore is that replacement therapy with thyroid hormone can precipitate an adrenal crisis in an undiagnosed individual. Hence, all patients with hypothyroidism and the possibility of APS should be screened for adrenal insufficiency to allow treatment with glucocorticoids prior to the initiation of thyroid hormone replacement. Treatment of mucocutaneous candidiasis with ketoconazole in an individual with subclinical adrenal insufficiency may also precipitate adrenal crisis. Furthermore, mucocutaneous candidiasis may be difficult to eradicate entirely. Severe cases of disease involvement may require systemic immunomodulatory therapy, but this is not commonly needed.

TABLE 409-3 CLINICAL FEATURES AND RECOMMENDED FOLLOW-UP FOR APS-1 AND APS-2

Component Disease	Recommended Evaluation
APS-1	
Addison's disease	Sodium, potassium, ACTH, cortisol, 21- and 17-hydroxylase autoantibodies
Diarrhea	History
Ectodermal dysplasia	Physical examination
Hypoparathyroidism	Serum calcium, phosphate, PTH
Hepatitis	Liver function tests
Hypothyroidism/ Graves' disease	TSH; thyroid peroxidase and/or thyroglobulin autoantibodies and anti-TSH receptor Ab
Male hypogonadism	FSH/LH, testosterone
Malabsorption	Physical examination, anti-IL-17 and anti-IL-22 autoantibodies
Mucocutaneous candidiasis	Physical examination, mucosal swab, stool samples
Obstipation	History
Ovarian failure	FSH/LH, estradiol
Pernicious anemia	CBC, vitamin B ₁₂ levels
Splenic atrophy	Blood smear for Howell-Jolly bodies; platelet count; ultrasound if positive
Type 1 diabetes	Glucose, hemoglobin A _{1c} , diabetes-associated autoantibodies (insulin, GAD65, IA-2, ZnT8)
APS-2	
Addison's disease	21-Hydroxylase autoantibodies, ACTH stimulation testing if positive
Alopecia	Physical examination
Autoimmune hyper- or hypothyroidism	TSH; thyroid peroxidase and/or thyroglobulin autoantibodies, anti-TSH receptor Ab
Celiac disease	Transglutaminase autoantibodies; small intestine biopsy if positive
Cerebellar ataxia	Dictated by signs and symptoms of disease
Chronic inflamma- tory demyelinating polyneuropathy	Dictated by signs and symptoms of disease
Hypophysitis	Dictated by signs and symptoms of disease, anti-Pit1 autoantibody
Idiopathic heart block	Dictated by signs and symptoms of disease
IgA deficiency	IgA level
Myasthenia gravis	Dictated by signs and symptoms of disease, antiacetylcholinesterase Ab
Myocarditis	Dictated by signs and symptoms of disease
Pernicious anemia	Anti-parietal cell autoantibodies CBC, vitamin B ₁₂ levels if positive
Serositis	Dictated by signs and symptoms of disease
Stiff man syndrome	Dictated by signs and symptoms of disease
Vitiligo	Physical examination, NALP-1 polymorphism

Abbreviations: Ab, antibody; ACTH, adrenocorticotropic hormone; APS, autoimmune polyendocrine syndrome; CBC, complete blood count; FSH, follicle-stimulating hormone; IL, interleukin; LH, luteinizing hormone; PTH, parathyroid hormone; TSH, thyroid-stimulating hormone.

APS-2

APS-2 (OMIM 269200) is more common than APS-1 with a prevalence of 1 in 100,000. It has a gender bias and occurs more often in female patients with a ratio of at least 3:1 compared to male patients. In contrast to APS-1, APS-2 often has its onset in adulthood with a peak incidence between 20 and 60 years of age. It shows a familial, multigenerational heritage (Table 409-2). The presence of two or more of the following endocrine deficiencies in the same patient defines the presence of APS-2: primary adrenal insufficiency (Addison's disease; 50–70%), Graves' disease or autoimmune thyroiditis (15–69%), type 1 diabetes mellitus (T1D; 40–50%), and primary hypogonadism.

Frequently associated autoimmune conditions include celiac disease (3–15%), myasthenia gravis, vitiligo, alopecia, serositis, and pernicious anemia. These conditions occur with increased frequency in affected patients but are also found in their family members (Table 409-3).

Genetic Considerations The overwhelming risk factor for APS-2 has been localized to the genes in the human lymphocyte antigen complex on chromosome 6. Primary adrenal insufficiency in APS-2, but not APS-1, is strongly associated with both HLA-DR3 and HLA-DR4. Other class I and class II genes and alleles, such as HLA-B8, HLA-DQ2 and HLA-DQ8, and HLA-DR subtype such as DRB1*0404, appear to contribute to organ-specific disease susceptibility (Table 409-4). HLA-B8- and HLA-DR3-associated illnesses include selective IgA deficiency, juvenile dermatomyositis, dermatitis herpetiformis, alopecia, scleroderma, autoimmune thrombocytopenia purpura, hypophysitis, metaphyseal osteopenia, and serositis.

Several other immune genes have been proposed to be associated with Addison's disease and therefore with APS-2 (Table 409-3). The "5.1" allele of a major histocompatibility complex (MHC) gene is an atypical class I HLA molecule MIC-A. The MIC-A5.1 allele has a very strong association with Addison's disease that is not accounted for by linkage disequilibrium with DR3 or DR4. Its role is complicated because certain HLA class I genes can offset this effect. PTPN22 codes for a polymorphism in a protein tyrosine phosphatase, which acts on intracellular signaling pathways in both T and B lymphocytes. It has been implicated in T1D, Addison's disease, and other autoimmune conditions. CTLA4 is a receptor on the T cell surface that modulates the activation state of the cell as part of the signal 2 pathway. Polymorphisms of this gene appear to cause downregulation of the cell surface expression of the receptor, leading to decreased T cell activation and proliferation. This appears to contribute to disease in Addison's disease and potentially other components of APS-2. Allelic variants of the IL-2Ra are linked to development of T1D and autoimmune thyroid disease and could contribute to the phenotype of APS-2 in certain individuals.

Diagnosis When one of the component disorders is present, a second associated disorder occurs more commonly than in the general population (Table 409-3). There is controversy as to which tests to use and how often to screen individuals for disease. A strong family history of autoimmunity should raise suspicion in an individual with an initial component diagnosis. The development of a rarer form of autoimmunity, such as Addison's disease, should prompt more extensive screening for other linked disorders compared to the diagnosis of autoimmune thyroid disease, which is relatively common.

Circulating autoantibodies, as previously discussed, can precede the development of disease by many years but would allow the clinician to follow the patient and identify the disease onset at its earliest time point (Tables 409-3 and 409-4). For each of the endocrine components of the disorder, appropriate autoantibody assays are listed and, if positive, should prompt physiologic testing to diagnose clinical or subclinical disease. For Addison's disease, antibodies to 21-hydroxylase antibodies are highly diagnostic for risk of adrenal insufficiency. However, individuals may take many years to develop overt hypoadrenalism. Screening of 21-hydroxylase antibody-positive patients can be performed measuring morning ACTH and cortisol on a yearly basis. Rising ACTH values over time or low morning cortisol in association with signs or symptoms of adrenal insufficiency should prompt testing via the cosyntropin stimulation test (Chap. 406). T1D can be screened for by measuring autoantibodies including anti-insulin, anti-GAD65, anti-IA-2, and anti-ZnT8. Risk for progression to disease can be based on the number of antibodies, and in some cases the titer (insulin autoantibody), as well as other metabolic factors (impaired oral glucose tolerance test). National Institutes of Health-sponsored trial groups such as Type 1 Diabetes TrialNet are screening first- and second-degree family members for these autoantibodies and identifying prediabetic individuals who may qualify for intervention trials to change the course of the disease prior to onset.

TABLE 409-4 APS-2 AND OTHER POLYENDOCRINE DISORDER ASSOCIATIONS

Disease	HLA Association	Initiating Factor	Mechanism	Autoantigen
Graves' Disease	DR3	Iodine Anti-CD52	Antibody	TSH receptor
Myasthenia gravis	DR3, DR7	Thymoma Penicillamine	Antibody	Acetylcholine receptor
Anti-insulin receptor	?	SLE or other autoimmune disease	Antibody	Insulin receptor
Hypoparathyroidism	?	?	Antibody	Cell surface inhibitor
Insulin autoimmune syndrome	DR4, DRB1*0406	Methimazole Sulfhydryl-containing drugs	Antibody	Insulin
Celiac disease	DQ2/DQ8	Gluten diet	T cell	Transglutaminase
Type 1 diabetes	DR3/DR4 DQ2/DQ8	? Congenital rubella	T cell	Insulin, GAD65, IA-2, ZnT8, IGRP
Addison's disease	DR3/DR4 DRB1*0404	Unknown	T cell	21-Hydroxylase P450-5cc
Thyroiditis	DR3/DQB1*0201 DQA1*0301	Iodine Interferon α	T cell	Thyroglobulin Thyroid peroxidase
Pernicious anemia	?	?	T cell	Intrinsic factor H+/K+ ATPase
Vitiligo	?	Melanoma Antigen Immunization	?	Melanocyte
Chromosome dysgenesis–trisomy 21 and Turner's syndrome	DQA1*0301	?	?	Thyroid, islet, transglutaminase
Hypophysitis	?	Pit-1, TDRD6	?	Pituitary, Pit-1

Abbreviations: APS, autoimmune polyendocrine syndrome; SLE, systemic lupus erythematosus; TSH, thyroid-stimulating hormone.

Screening tests for thyroid disease can include anti-thyroid peroxidase (TPO) or anti-thyroglobulin autoantibodies or anti-TSH receptor antibodies for Graves' disease. Yearly measurements of TSH can then be used to follow these individuals. Celiac disease can be screened for using the anti-tissue transglutaminase (tTg) antibody test. For those <20 years of age, testing every 1–2 years should be performed, whereas less frequent testing is indicated after the age of 20 because the majority of individuals who develop celiac disease have the antibody earlier in life. Positive tTg antibody test results should be confirmed on repeat testing, followed by small-bowel biopsy to document pathologic changes of celiac disease. Many patients have asymptomatic celiac disease that is nevertheless associated with osteopenia and impaired growth. If left untreated, symptomatic celiac disease has been reported to be associated with an increased risk of gastrointestinal malignancy, especially lymphoma.

The knowledge of the particular disease associations should guide other autoantibody or laboratory testing. A complete history and physical examination should be performed every 1–3 years including CBC, metabolic panel, TSH, and vitamin B₁₂ levels to screen for most of the possible abnormalities. More specific tests should be based on specific findings from the history and physical.

TREATMENT APS-2

With the exception of Graves' disease, the management of each of the endocrine components of APS-2 involves hormone replacement and is covered in detail in the chapters on adrenal (**Chap. 406**), thyroid (**Chap. 405**), gonadal (**Chaps. 411 and 412**), and parathyroid disease (**Chap. 424**). As noted for APS-1, adrenal insufficiency can be masked by primary hypothyroidism and should be considered and treated as discussed above. In patients with T1D, decreasing insulin requirements or hypoglycemia, without obvious secondary causes,

may indicate the emergence of adrenal insufficiency. Hypocalcemia in APS-2 patients is more likely due to malabsorption than hypoparathyroidism.

Immunotherapy for autoimmune endocrine disease has been reserved for T1D, for the most part, reflecting the lifetime burden of the disease for the individual patient and society. Although several immunotherapies (e.g., modified anti-CD3, rituximab, abatacept) can prolong the honeymoon phase of T1D, none has achieved long-term success. Active research using new approaches and combination therapy may change the treatment of this disease or other autoimmune conditions that share similar pathways. Furthermore, treatment of subclinical disease diagnosed by the presence of autoantibodies may provide a mechanism to preempt the development of overt disease and is the subject of active basic and clinical research.

IPEX

Immune dysregulation, polyendocrinopathy, enteropathy, and X-linked disease (IPEX; OMIM 304790) is a rare X-linked recessive disorder. The disease onset is in infancy and is characterized by severe enteropathy, T1D, and skin disease, as well as variable association with several other autoimmune disorders. Many infants die within the first days of life, but the course is variable, with some children surviving for 12–15 years. Early onset of T1D, often at birth, is highly suggestive of the diagnosis because nearly 80% of IPEX patients develop T1D. Although treatment of the individual disorders can temporarily improve the situation, treatment of the underlying immune deficiency is required and includes immunosuppressive therapy generally followed by hematopoietic stem cell transplantation. Transplantation is the only life-saving form of therapy and can be fully curative by normalizing the imbalanced immune system found in this disorder.

IPEX is caused by mutations in the *FOXP3* gene, which is also mutated in the Scurfy mouse, an animal model that shares much of the phenotype of IPEX patients. The *FOXP3* transcription factor is expressed in regulatory T cells designated CD4+CD25+FOXP3+ (Treg). Lack of this factor causes a profound deficiency of this Treg population and results in rampant autoimmunity due to the lack of peripheral tolerance normally provided by these cells. Certain mutations may lead to varying forms of expression of the full syndrome, and there are rare cases where the *FOXP3* gene is intact but other genes involved in this pathway (e.g., CD25, IL-2R α) may be causative.

THYMIC TUMORS

Thymomas and thymic hyperplasia are associated with several autoimmune diseases, with the most common being myasthenia gravis (44%) and red cell aplasia (20%). Graves' disease, T1D, and Addison's disease may also be associated with thymic tumors. Patients with myasthenia gravis and thymoma may have unique anti-acetylcholine receptor autoantibodies. Many thymomas lack AIRE expression within the thymoma, and this could be a potential factor in the development of autoimmunity. In support of this concept, thymoma is the one other disease with "frequent" development of anticytokine antibodies and mucocutaneous candidiasis in adults. The majority of tumors are malignant, and temporary remissions of the autoimmune condition can occur with resection of the tumor.

ANTI-INSULIN RECEPTOR ANTIBODIES

This is a very rare disorder where severe insulin resistance (type B) is caused by the presence of anti-insulin receptor antibodies. It is associated with acanthosis nigricans, which can also be associated with other forms of less severe insulin resistance. About one-third of patients have an associated autoimmune illness such as systemic lupus erythematosus or Sjögren's syndrome. Therefore, the presence of antinuclear antibodies, elevated erythrocyte sedimentation rate, hyperglobulinemia, leukopenia, and hypocomplementemia may accompany the presentation. The presence of anti-insulin receptor autoantibodies leads to marked insulin resistance, requiring more than 100,000 units of insulin to be given daily with only partial control of hyperglycemia. Patients can also have severe hypoglycemia due to partial activation of the insulin receptor by the antibody. The course of the disease is variable, and several patients have had spontaneous remissions. Therapy targeting B lymphocytes including rituximab, cyclophosphamide, and pulse steroids can induce remission of the disease.

INSULIN AUTOIMMUNE SYNDROME (HIRATA'S SYNDROME)

The insulin autoimmune syndrome, associated with Graves' disease and methimazole therapy (or other sulfhydryl-containing medications), is of particular interest due to a remarkably strong association with a specific HLA haplotype. Such patients with elevated titers of anti-insulin autoantibodies frequently present with hypoglycemia. In Japan, the disease is restricted to HLA-DR4-positive individuals with DRB1*0406. Curiously, a recent report demonstrated that five out of six Caucasian patients taking lipoic acid (sulfhydryl group) who developed insulin autoimmune syndrome were primarily DRB1*0403 (which is related to DRB1*0406); the sixth was DRB1*0406. In Hirata's syndrome the anti-insulin autoantibodies are often polyclonal. Discontinuation of the medication generally leads to resolution of the syndrome over time.

POEMS SYNDROME

POEMS (polyneuropathy, organomegaly, endocrinopathy, M-protein, and skin changes; also known as Crow-Fukase syndrome; OMIM

192240) patients usually present with a progressive sensorimotor polyneuropathy, diabetes mellitus (50%), primary gonadal failure (70%), and a plasma cell dyscrasia with sclerotic bony lesions. Associated findings can be hepatosplenomegaly, lymphadenopathy, and hyperpigmentation. Patients often present in the fifth to sixth decade of life and have a median survival after diagnosis of less than 3 years. The syndrome is assumed to be secondary to circulating immunoglobulins, but patients have excess vascular endothelial growth factor as well as elevated levels of other inflammatory cytokines such as IL1- β , IL-6, and tumor necrosis factor α . A small series of patients have been treated with thalidomide, leading to a decrease in vascular endothelial growth factor. Hyperglycemia responds to small, subcutaneous doses of insulin. The hypogonadism is due to primary gonadal disease with elevated plasma levels of follicle-stimulating hormone and luteinizing hormone. Temporary resolution of the features of POEMS, including normalization of blood glucose, may occur after radiotherapy for localized plasma cell lesions of bone or after chemotherapy, thalidomide, plasmapheresis, autologous stem cell transplantation, or treatment with all-*trans*-retinoic acid.

OTHER DISORDERS

Other diseases can exhibit polyendocrine deficiencies, including Kearns-Sayre syndrome, DIDMOAD syndrome (*d*iabetes *i*nsipidus, *d*iabetes mellitus, progressive bilateral *o*ptic atrophy, and sensorineural *d*eafness; also termed Wolfram's syndrome), Down's syndrome or trisomy 21 (OMIM 190685), Turner's syndrome (monosomy X, 45,X), and congenital rubella.

Kearns-Sayre syndrome (OMIM 530000) is a rare mitochondrial DNA disorder characterized by myopathic abnormalities leading to ophthalmoplegia and progressive weakness in association with several endocrine abnormalities, including hypoparathyroidism, primary gonadal failure, diabetes mellitus, and hypopituitarism. Crystalline mitochondrial inclusions are found in muscle biopsy specimens, and such inclusions have also been observed in the cerebellum. Antiparathyroid antibodies have not been described; however, antibodies to the anterior pituitary gland and striated muscle have been identified, and the disease may have autoimmune components. These mitochondrial DNA mutations occur sporadically and do not appear to be associated with a familial syndrome.

Wolfram's syndrome (OMIM 222300, chromosome 4; OMIM 598500, mitochondrial) is a rare autosomal recessive disease that is also called DIDMOAD. Neurologic and psychiatric disturbances are prominent in most patients and can cause severe disability. The disease is caused by defects in wolframin, a 100-kDa transmembrane protein that has been localized to the endoplasmic reticulum and is found in neuronal and neuroendocrine tissue. Its expression induces ion channel activity with a resultant increase in intracellular calcium and may play an important role in intracellular calcium homeostasis. Wolfram's syndrome appears to be a slowly progressive neurodegenerative process, and there is nonautoimmune selective destruction of the pancreatic beta cells. Diabetes mellitus with an onset in childhood is usually the first manifestation. Diabetes mellitus and optic atrophy are present in all reported cases, but expression of the other features is variable.

Down's syndrome, or trisomy 21 (OMIM 190685), is associated with the development of T1D, thyroiditis, and celiac disease. Patients with Turner's syndrome also appear to be at increased risk for the development of thyroid disease and celiac disease. It is recommended to screen patients with trisomy 21 and Turner's syndrome for associated autoimmune diseases on a regular basis.

SECTION 2 REPRODUCTIVE ENDOCRINOLOGY

410 Disorders of Sex Development

John C. Achermann, J. Larry Jameson

Sex development begins in utero but continues into young adulthood with the achievement of sexual maturity and reproductive capability. The major determinants of sex development can be divided into three components: chromosomal sex, gonadal sex (sex determination), and phenotypic sex (sex differentiation) (Fig. 410-1). Variations at each of these stages can result in disorders (or differences) of sex development (DSDs) (Table 410-1). In the newborn period, approximately 1 in 4000 babies require investigation because of ambiguous (atypical) genitalia. Urgent assessment is required, because some causes such as congenital adrenal hyperplasia (CAH) can be associated with life-threatening adrenal crises. Support for the parents and clear communication about the diagnosis and management options are essential. The involvement of an experienced multidisciplinary team is important for counseling, planning appropriate investigations, and discussing long-term well-being. DSDs can also present at other ages and to a range of health professionals. Subtler forms of gonadal dysfunction (e.g., Klinefelter's syndrome [KS], Turner's syndrome [TS]) often are diagnosed later in life by internists. Because these conditions are associated with a variety of psychological, reproductive, and potential medical consequences, an open dialogue must be established between the patient and health care providers to ensure continuity and attention to these issues.

SEX DEVELOPMENT

Chromosomal sex, defined by a karyotype, describes the X and/or Y chromosome complement (46,XY; 46,XX) that is established at the time of fertilization. The presence of a normal Y chromosome determines that testis development will occur even in the presence of multiple X chromosomes (e.g., 47,XXY or 48,XXXY). The loss of an X chromosome impairs gonad development (45,X or 45,X/46,XY mosaicism). Fetuses with no X chromosome (45,Y) are not viable.

Gonadal sex refers to the histologic and functional characteristics of gonadal tissue as testis or ovary. The embryonic gonad is bipotential and can develop (from ~42 days after conception) into either a testis or an ovary, depending on which genes are expressed (Fig. 410-2). Testis development is initiated by expression of the Y chromosome gene *SRY*

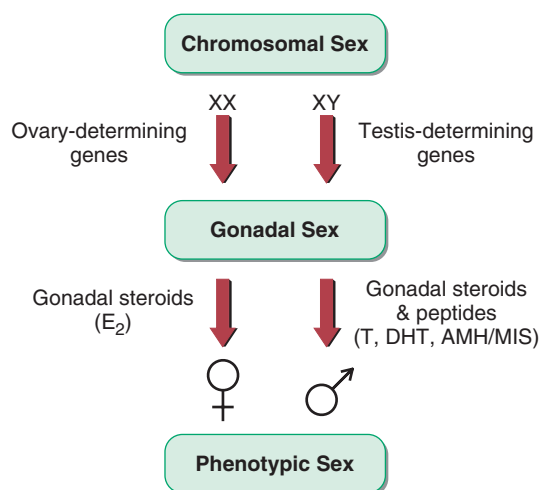


FIGURE 410-1 Sex development can be divided into three major components: chromosomal sex, gonadal sex, and phenotypic sex. DHT, dihydrotestosterone; MIS, müllerian-inhibiting substance also known as anti-müllerian hormone, AMH; T, testosterone.

(sex-determining region on the Y chromosome) that encodes an HMG box transcription factor. *SRY* is expressed transiently in cells destined to become Sertoli cells and serves as a pivotal switch to establish the testis lineage. Mutation of *SRY* prevents testis development in 46,XY individuals, whereas translocation of *SRY* in 46,XX individuals is sufficient to induce testis development and a male phenotype. Other genes are necessary to continue testis development. *SOX9* (*SRY*-related HMG-box gene 9) is upregulated by *SRY* in the developing testis but is suppressed in the ovary. *WT1* (Wilms' tumor-related gene 1) acts early in the genetic pathway and regulates the transcription of several genes, including *SFI* (*NR5A1*), *DAX1* (*NR0B1*), and *AMH* (encoding müllerian-inhibiting substance [MIS]). *SFI* encodes steroidogenic factor 1, a nuclear receptor that functions in cooperation with other transcription factors to regulate a large array of adrenal and gonadal genes, including *SOX9* and many genes involved in steroidogenesis. *SFI* mutations causing loss of function are found in ~10% of XY patients with gonadal dysgenesis and impaired androgenization. In contrast, duplication of a related gene *DAX1* also impairs testis development, revealing the exquisite sensitivity of the testis-determining pathway to gene dosage effects. *DAX1* loss-of-function mutations cause adrenal hypoplasia, hypogonadotropic hypogonadism, and testicular dysgenesis. In addition to the genes mentioned above, studies of humans and mice indicate that at least 30 other genes are also involved in gonad development (Fig. 410-2). These genes encode an array of signaling molecules and paracrine growth factors in addition to transcription factors.

Although ovarian development once was considered a "default" process, it is now clear that specific genes are expressed during the earliest stages of ovary development. Some of these factors may repress testis development (e.g., *WNT4*, R-spondin-1) (Fig. 410-2). Once the ovary has formed, additional factors are required for normal follicular development (e.g., follicle-stimulating hormone [FSH] receptor, *GDF9*). Steroidogenesis in the ovary requires the development of follicles that contain granulosa cells and theca cells surrounding the oocytes (Chap. 412). Thus, there is relatively limited ovarian steroidogenesis until puberty.

Germ cells also develop in a sex dimorphic manner. In the developing ovary, primordial germ cells (PGCs) proliferate and enter meiosis, whereas they proliferate and then undergo mitotic arrest in the developing testis. PGC entry into meiosis is initiated by retinoic acid that activates *STRA8* (stimulated by retinoic acid 8) and other genes involved in meiosis. The developing testis produces high levels of *CYP26B1*, an enzyme that degrades retinoic acid, preventing PGC entry into meiosis. Approximately 7 million germ cells are present in the fetal ovary in the second trimester, and 1 million remain at birth. Only 400 are ovulated during a woman's reproductive life span (Chap. 412).

Phenotypic sex refers to the structures of the external and internal genitalia and secondary sex characteristics. The developing testis releases anti-müllerian hormone (AMH; also known as müllerian-inhibiting substance [MIS]) from Sertoli cells and testosterone from Leydig cells. AMH is a member of the transforming growth factor (TGF) β family and acts through specific receptors to cause regression of the müllerian structures from 60–80 days after conception. At ~60–140 days after conception, testosterone supports the development of wolffian structures, including the epididymides, vasa deferentia, and seminal vesicles. Testosterone is the precursor for dihydrotestosterone (DHT), a potent androgen that promotes development of the external genitalia, including the penis and scrotum (65–100 days, and thereafter) (Fig. 410-3). The urogenital sinus develops into the prostate and prostatic urethra in the male and into the urethra and lower portion of the vagina in the female. The genital tubercle becomes the glans penis in the male and the clitoris in the female. The urogenital swellings form the scrotum or the labia majora, and the urethral folds fuse to form the shaft of the penis and the male urethra or the labia minora. In the female, wolffian ducts regress and the müllerian ducts form the

TABLE 410-1 CLASSIFICATION OF DISORDERS OF SEX DEVELOPMENT (DSDS)

Sex Chromosome DSD	46,XY DSD (see Table 410-3)	46,XX DSD (see Table 410-4)
47,XXY (Klinefelter's syndrome and variants)	Disorders of gonadal (testis) development Complete or partial gonadal dysgenesis (e.g., SRY, SOX9, SF1, WT1, DHH, MAP3K1)	Disorders of gonadal (ovary) development Gonadal dysgenesis Ovotesticular DSD Testicular DSD (e.g., SRY+, dup SOX9, RSPO1)
45,X (Turner's syndrome and variants)	Impaired fetal Leydig cell function (e.g., SF1/NR5A1, Cxorf6/MAMLD1)	Androgen excess
45,X/46,XY mosaicism (mixed gonadal dysgenesis)	Ovotesticular DSD	Fetal
46,XX/46,XY (chimerism/mosaicism)	Testis regression	3β-Hydroxysteroid dehydrogenase II (HSD3β2) 21-Hydroxylase (CYP21A2) P450 oxidoreductase (POR) 11β-Hydroxylase (CYP11B1) Glucocorticoid receptor mutations
	Disorders in androgen synthesis or action	Fetoplacental
	Disorders of androgen biosynthesis	Aromatase deficiency (CYP19) Oxidoreductase deficiency (POR)
	LH receptor (LHCGR)	Maternal
	Smith-Lemli-Opitz syndrome	Maternal virilizing tumors (e.g., luteomas)
	Steroidogenic acute regulatory (StAR) protein	Androgenic drugs
	Cholesterol side-chain cleavage (CYP11A1)	Other
	3β-Hydroxysteroid dehydrogenase II (HSD3B2)	Syndromic associations (e.g., cloacal anomalies)
	17α-Hydroxylase/17,20-lyase (CYP17A1)	Müllerian agenesis/hypoplasia (e.g., MRKH)
	P450 oxidoreductase (POR)	Uterine abnormalities (e.g., MODY5)
	Cytochrome b5 (CYB5A)	Vaginal atresia (e.g., McKusick-Kaufman)
	17β-Hydroxysteroid dehydrogenase III (HSD17B3)	Labial adhesions
	5α-Reductase II (SRD5A2)	
	Aldo-keto reductase 1C2 (AKR1C2)	
	Disorders of androgen action	
	Androgen insensitivity syndrome	
	Drugs and environmental modulators	
	Other	
	Syndromic associations of male genital development	
	Persistent müllerian duct syndrome	
	Vanishing testis syndrome	
	Isolated hypospadias	
	Congenital hypogonadotropic hypogonadism	
	Cryptorchidism	
	Environmental influences	

Source: Modified from IA Hughes: Arch Dis Child 91:554, 2006.

fallopian tubes, uterus, and upper segment of the vagina. A female phenotype will develop in the absence of the gonad, but estrogen is needed for maturation of the uterus and breast at puberty.

DISORDERS OF CHROMOSOMAL SEX

Variations in sex chromosome number and structure can present as DSDs (e.g., 45,X/46,XY). KS (47,XXY) and TS (45,X) do not usually present with genital ambiguity but are associated with gonadal dysfunction (Table 410-2).

KLINFELTER'S SYNDROME (47,XXY)

Pathophysiology The classic form of KS (47,XXY) occurs after meiotic nondisjunction of the sex chromosomes during gametogenesis (40% during spermatogenesis, 60% during oogenesis) (Chap. 83e). Mosaic forms of KS (46,XY/47,XXY) are thought to result from chromosomal mitotic nondisjunction within the zygote and occur in at least 10% of individuals with this condition. Other chromosomal variants of KS (e.g., 48,XXYY, 48,XXXY) have been reported but are less common.

Clinical Features KS is characterized by small testes, infertility, gynecomastia, tall stature/increased leg length, and hypogonadism in phenotypic males. It has an incidence of at least 1 in 1000 men, but approximately 75% of cases are not diagnosed. Of those who are diagnosed, only 10% are identified prepubertally, usually because of small genitalia or cryptorchidism. Others are diagnosed after puberty, usually based on impaired androgenization and/or gynecomastia. Developmental delay, speech difficulties, and poor motor skills may be features but are variable, especially in adolescence. Later in life, body habitus or infertility leads to the diagnosis. Testes are small and

firm (median length 2.5 cm [4 mL volume]; almost always <3.5 cm [12 mL]) and typically seem inappropriately small for the degree of androgenization. Biopsies are not usually necessary but typically reveal seminiferous tubule hyalinization and azoospermia. Other clinical features of KS are listed in Table 410-2. Plasma concentrations of FSH and luteinizing hormone (LH) are increased in most adults with 47,XXY, and plasma testosterone is decreased (50–75%), reflecting primary gonadal failure. Estradiol is often increased, likely because of chronic Leydig cell stimulation by LH and aromatization of androstenedione by adipose tissue; the increased ratio of estradiol-to-testosterone results in gynecomastia (Chap. 411). Patients with mosaic forms of KS have less severe clinical features, have larger testes, and sometimes achieve spontaneous fertility.

TREATMENT KLINEFELTER'S SYNDROME

Growth, endocrine function, and bone mineralization should be monitored, especially from adolescence. Educational and psychological support is important for many individuals with KS. Androgen supplementation improves virilization, libido, energy, hypofibrinolysis, and bone mineralization in men with low testosterone levels but may occasionally worsen gynecomastia (Chap. 411). Gynecomastia can be treated by surgical reduction if it causes concern (Chap. 411). Fertility has been achieved by using in vitro fertilization in men with oligospermia or with intracytoplasmic sperm injection (ICSI) after retrieval of spermatozoa by testicular sperm extraction techniques. In specialized centers, successful spermatozoa retrieval using this technique is possible in >50% of men with nonmosaic

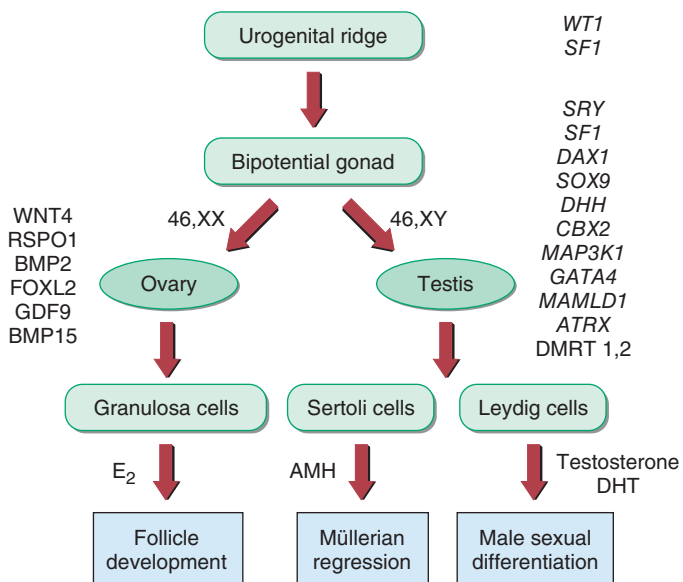


FIGURE 410-2 The genetic regulation of gonadal development.

AMH, anti-Müllerian hormone (Müllerian-inhibiting substance); *ATRX*, α -thalassemia, mental retardation on the X; *BMP2* and *15*, bone morphogenetic factors 2 and 15; *CBX2*, chromobox homologue 2; *DAX1*, dosage sensitive sex-reversal, adrenal hypoplasia congenita on the X chromosome, gene 1; *DHH*, desert hedgehog; *DHT*, dihydrotestosterone; *DMRT 1,2*, doublesex MAB3-related transcription factor 1,2; *FOXL2*, forkhead transcription factor L2; *GATA4*, GATA binding protein 4; *GDF9*, growth differentiation factor 9; *MAMLD1*, mastermind-like domain containing 1; *MAP3K1*, mitogen-activated protein kinase kinase kinase 1; *RSPO1*, R-spondin 1; *SF1*, steroidogenic factor 1 (also known as *NR5A1*); *SOX9*, *SRY*-related HMG-box gene 9; *SRY*, sex-determining region on the Y chromosome; *WNT4*, wingless-type MMTV integration site 4; *WT1*, Wilms' tumor-related gene 1.

KS. Results may be better in younger men. After ICSI and embryo transfer, successful pregnancies can be achieved in ~50% of these cases. The risk of transmission of this chromosomal abnormality needs to be considered, and preimplantation screening may be

desired, although this outcome is much less common than originally predicted. Long-term monitoring of men with KS is important given the increased risk of breast cancer, cardiovascular disease, metabolic syndrome, and autoimmune disorders. Because most men with KS are never diagnosed, it is important that all internists consider this diagnosis in men with these features who might be seeking medical advice for other conditions.

TURNER'S SYNDROME (GONADAL DYSGENESIS; 45,X)

Pathophysiology Approximately one-half of women with TS have a 45,X karyotype, about 20% have 45,X/46,XX mosaicism, and the remainder have structural abnormalities of the X chromosome such as X fragments, isochromosomes, or rings. The clinical features of TS result from haploinsufficiency of multiple X chromosomal genes (e.g., short stature homeobox, *SHOX*). However, imprinted genes also may be affected when the inherited X has different parental origins.

Clinical Features TS is characterized by bilateral streak gonads, primary amenorrhea, short stature, and multiple congenital anomalies in phenotypic females. It affects ~1 in 2500 women and is diagnosed at different ages depending on the dominant clinical features (Table 410-2). Prenatally, a diagnosis of TS usually is made incidentally after chorionic villus sampling or amniocentesis for unrelated reasons such as advanced maternal age. Prenatal ultrasound findings include increased nuchal translucency. The postnatal diagnosis of TS should be considered in female neonates or infants with lymphedema, nuchal folds, low hairline, or left-sided cardiac defects and in girls with unexplained growth failure or pubertal delay. Although limited spontaneous pubertal development occurs in up to 30% of girls with TS (10%, 45,X; 30–40%, 45,X/46,XX) and ~2% reach menarche, the vast majority of women with TS develop complete ovarian insufficiency. Therefore, this diagnosis should be considered in all women who present with primary or secondary amenorrhea and elevated gonadotropin levels.

TREATMENT TURNER'S SYNDROME

The management of girls and women with TS requires a multidisciplinary approach because of the number of potentially involved organ systems. Detailed cardiac and renal evaluation should be performed at the time of diagnosis. Individuals with congenital heart

TABLE 410-2 CLINICAL FEATURES OF CHROMOSOMAL DISORDERS OF SEX DEVELOPMENT (DSD)

Disorder	Common Chromosomal Complement	Gonad	Genitalia		
			External	Internal	Breast Development
Klinefelter's syndrome	47,XXY or 46,XY/47,XXY	Hyalinized testes	Male	Male	Gynecomastia
Clinical Features					
Small testes, azoospermia, decreased facial and axillary hair, decreased libido, tall stature and increased leg length, decreased penile length, increased risk of breast tumors, thromboembolic disease, learning difficulties, speech delay and decreased verbal IQ, obesity, diabetes mellitus, metabolic syndrome, varicose veins, hypothyroidism, systemic lupus erythematosus, epilepsy					
Turner's syndrome	45,X or 45,X/46,XX	Streak gonad or immature ovary	Female	Hypoplastic female	Immature female
Clinical Features					
Infancy: lymphedema, web neck, shield chest, low-set hairline, cardiac defects and coarctation of the aorta, urinary tract malformations, and horseshoe kidney					
Childhood: short stature, cubitus valgus, short neck, short fourth metacarpals, hypoplastic nails, micrognathia, scoliosis, otitis media and sensorineural hearing loss, ptosis and amblyopia, multiple nevi and keloid formation, autoimmune thyroid disease, visuospatial learning difficulties					
Adulthood: pubertal failure and primary amenorrhea, hypertension, obesity, dyslipidemia, impaired glucose tolerance and insulin resistance, autoimmune thyroid disease, cardiovascular disease, aortic root dilation, osteoporosis, inflammatory bowel disease, chronic hepatic dysfunction, increased risk of colon cancer, hearing loss					
45,X/46,XY mosaicism	45,X/46,XY	Testis or streak gonad	Variable	Variable	Usually male
Clinical Features					
Short stature, increased risk of gonadal tumors, some Turner's syndrome features					
Ovotesticular DSD (true hermaphroditism)	46,XX/46,XY	Testis and ovary or ovo-testis	Variable	Variable	Gynecomastia
Clinical Features					
Possible increased risk of gonadal tumors					

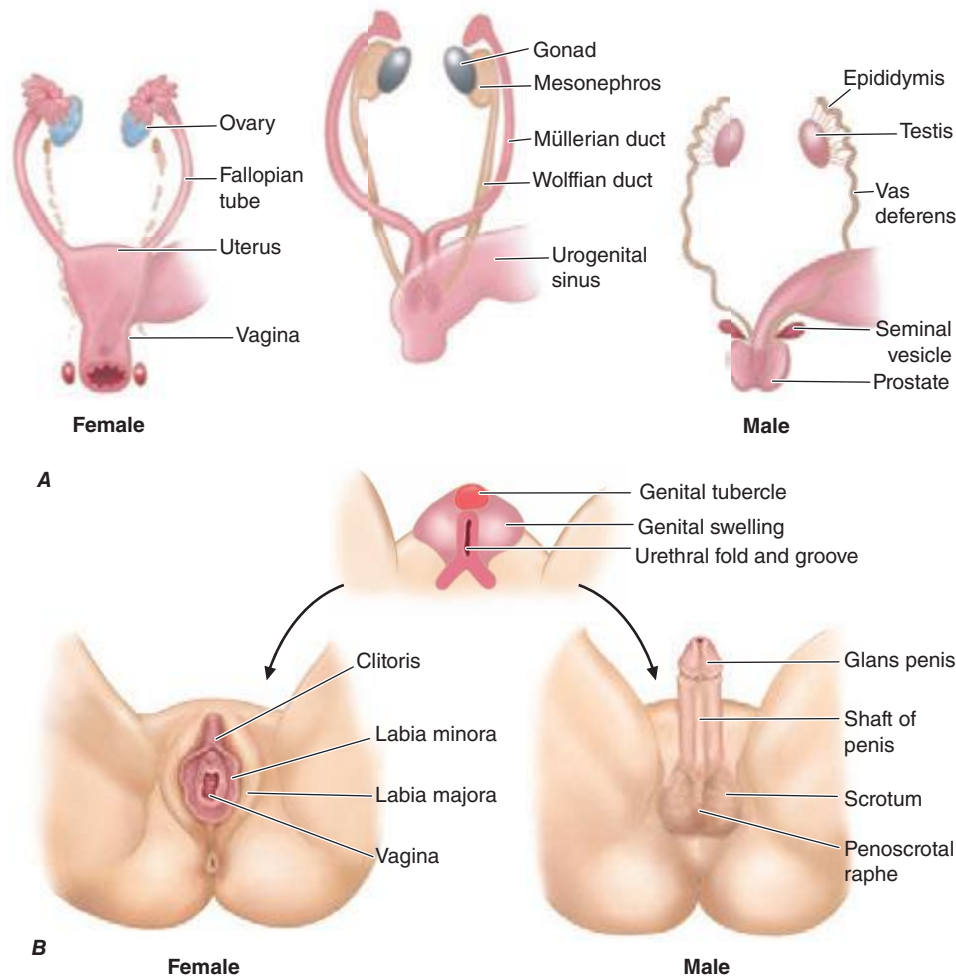


FIGURE 410-3 Sex development. A. Internal urogenital tract. **B.** External genitalia. (After E Braunwald et al [eds]: *Harrison's Principles of Internal Medicine*, 15th ed. New York, McGraw-Hill, 2001.)

defects (CHDs) (30%) (bicuspid aortic valve, 30–50%; coarctation of the aorta, 30%; aortic root dilation, 5%) require long-term follow-up by an experienced cardiologist, antibiotic prophylaxis for dental or surgical procedures, and serial magnetic resonance imaging (MRI) of aortic root dimensions, because progressive aortic root dilation is associated with increased risk of aortic dissection. Individuals found to have congenital renal and urinary tract malformations (30%) are at risk for urinary tract infections, hypertension, and nephrocalcinosis. Hypertension can occur independently of cardiac and renal malformations and should be monitored and treated as in other patients with essential hypertension. Clitoral enlargement or other evidence of virilization suggests the presence of covert, translocated Y chromosomal material and is associated with increased risk of gonadoblastoma. Regular assessment of thyroid function, weight, dentition, hearing, speech, vision, and educational issues should be performed during childhood. Otitis media and middle-ear disease are prevalent in childhood (50–85%), and sensorineural hearing loss becomes progressively common with age (70–90%). Autoimmune hypothyroidism (15–30%) can occur in childhood but has a mean age of onset in the third decade. Counseling about long-term growth and fertility issues should be provided. Patient support groups are active throughout the world and can play an invaluable role.

Short stature can be an issue for some girls because untreated final height rarely exceeds 150 cm in nonmosaic 45,X TS. High-dose recombinant growth hormone stimulates growth rate in children with TS and is occasionally combined with low doses of the nonaromatizable anabolic steroid oxandrolone (up to 0.05 mg/kg per day) in an older child (>9 years). However, final height increments are often about 5–10 cm, and individualization of treatment response

to regimens may be beneficial. Girls with evidence of ovarian insufficiency require estrogen replacement to induce breast and uterine development, support growth, and maintain bone mineralization. Most physicians now initiate low-dose estrogen therapy (one-tenth to one-eighth of the adult replacement dose) to induce puberty at an age-appropriate time (~12 years). Doses of estrogen are increased gradually to allow development over a 2- to 4-year period. Progestins are added later to regulate withdrawal bleeds. Some women with TS have achieved successful pregnancy after ovum donation and in vitro fertilization but are high risk, and cardiac assessment is required. Long-term follow-up of women with TS involves careful surveillance of sex hormone replacement and reproductive function, bone mineralization, cardiac function and aortic root dimensions, blood pressure, weight and glucose tolerance, hepatic and lipid profiles, thyroid function, and hearing. This service is provided by a dedicated TS clinic in some centers.

45,X/46,XY MOSAICISM (MIXED GONADAL DYSGENESIS)

The phenotype of individuals with 45,X/46,XY mosaicism (sometimes called *mixed gonadal dysgenesis*) can vary considerably. Some have a predominantly female phenotype with somatic features of TS, streak gonads, and müllerian structures, and are managed as TS with a Y chromosome. Most 45,X/46,XY individuals have a male phenotype and testes, and the diagnosis is made incidentally after amniocentesis or during investigation of infertility. In practice, most newborns referred for assessment have atypical genitalia and variable somatic features. Management is complex and needs to be individualized. A female sex-of-rearing is often assigned if uterine structures are present, gonads are intraabdominal, and phallic development is incomplete. In

such situations, gonadectomy usually is considered to prevent further androgen secretion at puberty and prevent risk of gonadoblastoma (up to 25%). Individuals raised as males usually require reconstructive surgery for hypospadias and removal of dysgenetic or streak gonads if the gonads cannot be brought down into the scrotum. Scrotal testes can be preserved but require regular examination for tumor development and sonography at the time of puberty. Biopsy for carcinoma *in situ* is recommended in adolescence, and testosterone supplementation may be required to support androgenization in puberty or if low testosterone is detected in adulthood. Height potential is usually attenuated; some children receive recombinant growth hormone using TS protocols. Screening for cardiac, renal, and other TS features should be considered, and psychological support offered for the family and young person.

OVOTESTICULAR DSD

Ovotesticular DSD (formerly called *true hermaphroditism*) occurs when both an ovary and a testis—or when an ovotestis—are found in one individual. Most individuals with this diagnosis have a 46,XX karyotype, especially in sub-Saharan Africa, and present with ambiguous genitalia at birth or with breast development and phallic development at puberty. A 46,XX/46,XY chimeric karyotype is less common and has a variable phenotype.

DISORDERS OF GONADAL AND PHENOTYPIC SEX

Disorders of gonadal and phenotypic sex can result in underandrogenization of individuals with a 46,XY karyotype (46,XY DSD) and the excess androgenization of individuals with a 46,XX karyotype (46,XX DSD) (Table 410-1). These disorders cover a spectrum of phenotypes ranging from “46,XY phenotypic females” or “46,XX phenotypic males” to individuals with atypical genitalia.

46,XY DSD

Underandrogenization of the 46,XY fetus (formerly called *male pseudohermaphroditism*) reflects defects in androgen production or action. It can result from disorders of testis development, defects of androgen synthesis, or resistance to testosterone and DHT (Table 410-1).

Disorders of Testis Development • TESTICULAR DYSGENESIS *Pure* (or *complete*) *gonadal dysgenesis* (*Swyer’s syndrome*) is associated with streak gonads, müllerian structures (due to insufficient AMH/MIS secretion), and a complete absence of androgenization. Phenotypic females with this condition often present because of absent pubertal development and are found to have a 46,XY karyotype. Serum sex steroids, AMH/MIS, and inhibin B are low, and LH and FSH are elevated. Patients with *partial gonadal dysgenesis* (*dysgenetic testes*) may produce enough MIS to regress the uterus and sufficient testosterone for partial androgenization, and therefore usually present in the newborn period with atypical genitalia. Gonadal dysgenesis can result from mutations or deletions of testis-promoting genes (*WT1*, *CBX2*, *SF1*, *SRY*, *SOX9*, *MAP3K1*, *DHH*, *GATA4*, *ATRX*, *ARX*, *DMRT*) or duplication of chromosomal loci containing “antitestis” genes (e.g., *WNT4/RSPO1*, *DAX1*) (Table 410-3). Among these, deletions or mutations of *SRY* and heterozygous mutations of *SF1* (*NR5A1*) appear to be most common but still account collectively for <25% of cases. Associated clinical features may be present, reflecting additional functional roles for these genes. For example, renal dysfunction occurs in patients with specific *WT1* mutations (Denys-Drash and Frasier’s syndromes), primary adrenal failure occurs in some patients with *SF1* mutations, and severe cartilage abnormalities (campomelic dysplasia) are the predominant clinical feature of *SOX9* mutations. A family history of DSD, infertility, or early menopause is important because mutations in *SF1/NR5A1* can be inherited from a mother in a sex-limited dominant manner (which can mimic X-linked inheritance). In some cases, a woman may later develop primary ovarian insufficiency because of the effect of *SF1* on the ovary. Intraabdominal dysgenetic testes should be removed to prevent malignancy, and estrogens can be used to induce secondary sex characteristics and uterine development in 46,XY individuals raised as females, if it is felt that a female gender identity is established. *Absent*

(*vanishing*) *testis syndrome* (*bilateral anorchia*) reflects regression of the testis during development. The etiology is unknown, but the absence of müllerian structures indicates adequate secretion of AMH early in utero. In most cases, androgenization of the external genitalia is either normal or slightly impaired (e.g., small penis, hypospadias). These individuals can be offered testicular prostheses and should receive androgen replacement in adolescence.

Disorders of Androgen Synthesis Defects in the pathway that regulates androgen synthesis (Fig. 410-4) cause underandrogenization of the 46,XY fetus (Table 410-1). Müllerian regression is unaffected because Sertoli cell function is preserved. Most of these conditions can present with a spectrum of genital phenotypes, ranging from female-typical external genitalia or clitoromegaly in the more severe situations to penoscrotal hypospadias or a small phallus in others.

LH RECEPTOR Mutations in the LH receptor (LHCGR) cause Leydig cell hypoplasia and androgen deficiency, due to impaired actions of human chorionic gonadotropin in utero and LH late in gestation and during the neonatal period. As a result, testosterone and DHT synthesis are insufficient for complete androgenization.

STEROIDOGENIC ENZYME PATHWAYS Mutations in *steroidogenic acute regulatory protein* (*StAR*) and *CYP11A1* affect both adrenal and gonadal steroidogenesis (Fig. 410-4) (Chap. 406). Affected individuals (46,XY) usually have severe early-onset salt-losing adrenal failure and a female phenotype, although later-onset milder variants have been reported. Defects in *3 β -hydroxysteroid dehydrogenase type 2* (*HSD3 β 2*) also cause adrenal insufficiency in severe cases, but the accumulation of dehydroepiandrosterone (DHEA) has a mild androgenizing effect, resulting in ambiguous genitalia or hypospadias. Salt loss occurs in many but not all cases. Patients with CAH due to *17 α -hydroxylase* (*CYP17*) *deficiency* have variable underandrogenization and develop hypertension and hypokalemia due to the potent salt-retaining effects of corticosterone and 11-deoxycorticosterone. Patients with complete loss of *17 α -hydroxylase* function often present as phenotypic females who fail to enter puberty and are found to have inguinal testes and hypertension in adolescence. Some mutations in *CYP17* selectively impair 17,20-lyase activity without altering *17 α -hydroxylase* activity, leading to underandrogenization without mineralocorticoid excess and hypertension. Disruption of the coenzyme, *cytochrome b5* (*CYB5A*), can present similarly, and methemoglobinemia is usually present. Mutations in *P450 oxidoreductase* (*POR*) affect multiple steroidogenic enzymes, leading to impaired androgenization and a biochemical pattern of apparent combined 21-hydroxylase and *17 α -hydroxylase* deficiency, sometimes with skeletal abnormalities (Antley-Bixler craniosynostosis). Defects in *17 β -hydroxysteroid dehydrogenase type 3* (*HSD17 β 3*) and *5 α -reductase type 2* (*SRD5A2*) interfere with the synthesis of testosterone and DHT, respectively. These conditions are characterized by minimal or absent androgenization in utero, but some phallic development can occur during adolescence due to the action of other enzyme isoforms. Individuals with *5 α -reductase type 2* deficiency have normal wolffian structures and usually do not develop breast tissue. At puberty, the increase in testosterone induces muscle mass and other virilizing features despite DHT deficiency. Some individuals change gender from female to male at puberty. Thus, the management of this disorder is challenging. DHT cream can improve prepubertal phallic growth in patients raised as male. Gonadectomy before adolescence and estrogen replacement at puberty can be considered in individuals raised as females who have a female gender identity. Disruption of alternative pathways to fetal DHT production might also present with 46,XY DSD (*AKR1C2/AKR1C4*).

Disorders of Androgen Action • ANDROGEN INSENSITIVITY SYNDROME Mutations in the androgen receptor cause resistance to androgen (testosterone, DHT) action or the *androgen insensitivity syndrome* (*AIS*). *AIS* is a spectrum of disorders that affects at least 1 in 100,000 46,XY individuals. Because the androgen receptor is X-linked, only 46,XY offspring are affected if the mother is a carrier of a mutation. XY individuals with *complete AIS* (formerly called *testicular feminization syndrome*) have a female phenotype, normal breast development

Gene	Inheritance	Gonad	Uterus	External Genitalia	Associated Features
Disorders of Testis Development					
<i>WT1</i>	AD	Dysgenetic testis	+/-	Female or ambiguous	Wilms' tumor, renal abnormalities, gonadal tumors (WAGR, Denys-Drash and Frasier's syndromes)
<i>CBX2</i>	AD	Ovary	+	Female	
<i>SF1</i>	AR/AD (SL)	Dysgenetic testis/Leydig dysfunction	+/-	Female or ambiguous	Primary adrenal failure; primary ovarian insufficiency in female (46,XX) relatives
<i>SRY</i>	Y	Dysgenetic testis or ovotestis	+/-	Female or ambiguous	
<i>SOX9</i>	AD	Dysgenetic testis or ovotestis	+/-	Female or ambiguous	Campomelic dysplasia
<i>MAP3K1</i>	AD (SL)	Dysgenetic testis	+/-	Female or ambiguous	
<i>DHH</i>	AR	Dysgenetic testis	+	Female	Minifascicular neuropathy
<i>GATA4</i>	AD	Dysgenetic testis	-	Ambiguous or male	Congenital heart disease
<i>ATRX</i>	X	Dysgenetic testis	-	Female or ambiguous	α Thalassemia, developmental delay
<i>ARX</i>	X	Dysgenetic testis	-	Male or ambiguous	Developmental delay; X-linked lissencephaly
<i>MAMLD1</i>	X	Dysgenetic testis/Leydig dysfunction	-	Hypospadias	
<i>DAX1</i>	dupXp21	Dysgenetic testis	+/-	Female or ambiguous	
<i>WNT4/RSP01</i>	dup1p35	Dysgenetic testis	+	Ambiguous	
Disorders of Androgen Synthesis					
<i>LHR</i>	AR	Testis	-	Female, ambiguous or micropenis	Leydig cell hypoplasia
<i>DHCR7</i>	AR	Testis	-	Variable	Smith-Lemli-Opitz syndrome: coarse facies, second-third toe syndactyly, failure to thrive, developmental delay, cardiac and visceral abnormalities
<i>StAR</i>	AR	Testis	-	Female or ambiguous	Congenital lipoid adrenal hyperplasia (primary adrenal failure)
<i>CYP11A1</i>	AR	Testis	-	Ambiguous	Primary adrenal failure
<i>HSD3B2</i>	AR	Testis	-	Ambiguous	CAH, primary adrenal failure \pm salt loss, partial androgenization due to \uparrow DHEA
<i>CYP17</i>	AR	Testis	-	Female or ambiguous	CAH, hypertension due to \uparrow corticosterone and 11-deoxycorticosterone, except in isolated 17,20-lyase deficiency
<i>CYB5A</i>	AR	Testis	-	Ambiguous	Apparent isolated 17,20-lyase deficiency; methemoglobinemia
<i>POR</i>	AR	Testis	-	Ambiguous or male	Mixed features of 21-hydroxylase deficiency and 17 α -hydroxylase/17,20-lyase deficiency, sometimes associated with Antley-Bixler craniosynostosis
<i>HSD17B3</i>	AR	Testis	-	Female or ambiguous	Partial androgenization at puberty, \uparrow androstenedione-to-testosterone ratio
<i>SRD5A2</i>	AR	Testis	-	Ambiguous or micropenis	Partial androgenization at puberty, \uparrow testosterone-to-dihydrotestosterone ratio
<i>AKR1C2</i> (<i>AKR1C4</i>)	AR	Testis	-	Female or ambiguous	Decreased fetal DHT production
Disorders of Androgen Action					
Androgen receptor	X	Testis	-	Female, ambiguous, micropenis or normal male	Phenotypic spectrum from complete androgen insensitivity syndrome (female external genitalia) and partial androgen insensitivity (ambiguous) to normal male genitalia and infertility

Abbreviations: AD, autosomal dominant; *AKR1C2*, aldo-keto reductase family 1 member 2; AR, autosomal recessive; *ARX*, aristaless related homeobox, X-linked; *ATRX*, α -thalassemia, mental retardation on the X; CAH, congenital adrenal hyperplasia; *CBX2*, chromobox homologue 2; *CYB5A*, cytochrome b5 POR, P450 oxidoreductase; *CYP11A1*, P450 cholesterol side-chain cleavage; *CYP17*, 17 α -hydroxylase and 17,20-lyase; *DAX1*, dosage sensitive sex-reversal, adrenal hypoplasia congenita on the X chromosome, gene 1; DHEA, dehydroepiandrosterone; *DHCR7*, sterol 7 δ reductase; *DHH*, desert hedgehog; *GATA4*, GATA binding protein 4; *HSD17B3*, 17 β -hydroxysteroid dehydrogenase type 3; *HSD3B2*, 3 β -hydroxysteroid dehydrogenase type 2; *LHR*, LH receptor; *MAP3K1*, mitogen-activated protein kinase kinase kinase 1; *SF1*, steroidogenic factor 1; SL, sex-limited; *SOX9*, *SRY*-related HMG-box gene 9; *SRD5A2*, 5 α -reductase type 2; *SRY*, sex-related gene on the Y chromosome; *StAR*, steroidogenic acute regulatory protein; WAGR, Wilms' tumor, aniridia, genitourinary anomalies, and mental retardation; *WNT4*, wingless-type mouse mammary tumor virus integration site, 4; *WT1*, Wilms' tumor-related gene 1.

(due to aromatization of testosterone), a short vagina but no uterus (because MIS production is normal), scanty pubic and axillary hair, and a female gender identity and sex role behavior. Gonadotropins and testosterone levels can be low, normal, or elevated, depending on the degree of androgen resistance and the contribution of estradiol to feedback inhibition of the hypothalamic-pituitary-gonadal axis. AMH/MIS levels in childhood are normal or high. Most patients present with inguinal hernias (containing testes) in childhood or with primary amenorrhea in late adolescence. Gonadectomy sometimes is

offered for girls diagnosed in childhood, because there is a low risk of malignancy, and estrogen replacement is prescribed. Alternatively, the gonads can be left in situ until breast development is complete and removed because of tumor risk. Some adults with complete AIS decline gonadectomy, but should be counseled about the risk of malignancy, especially because early detection of premalignant changes by imaging or biomarkers is currently not possible. The use of graded dilators in adolescence is usually sufficient to dilate the vagina for sexual intercourse.

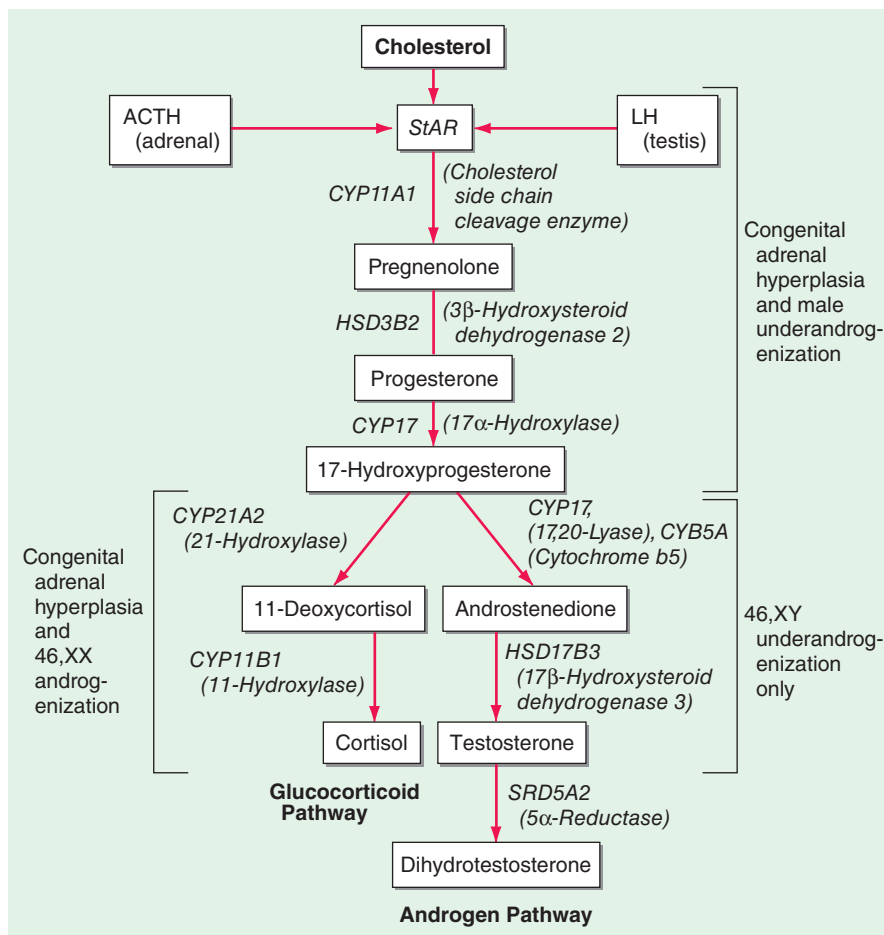


FIGURE 410-4 Simplified overview of glucocorticoid and androgen synthesis

pathways. Defects in *CYP21A2* and *CYP11B1* shunt steroid precursors into the androgen pathway and cause androgenization of the 46,XX fetus. Testosterone is synthesized in the testicular Leydig cells and converted to dihydrotestosterone peripherally. Defects in enzymes involved in androgen synthesis result in underandrogenization of the 46,XY fetus. *StAR*, steroidogenic acute regulatory protein. (After E Braunwald et al [eds]: *Harrison's Principles of Internal Medicine*, 15th ed. New York, McGraw-Hill, 2001.)

Partial AIS (Reifenstein's syndrome) results from androgen receptor mutations that maintain residual function. Patients often present in infancy with penoscrotal hypospadias and small undescended testes and with gynecomastia at the time of puberty. Those individuals raised as males usually require hypospadias repair in childhood and may need breast reduction in adolescence. Some boys enter puberty spontaneously. High-dose testosterone has been given to support development if puberty does not progress, but long-term data are limited. More severely underandrogenized patients present with clitoral enlargement and labial fusion and may be raised as females. The surgical and psychosexual management of these patients is complex and requires active involvement of the parents and the patient during the appropriate stages of development. *Azoospermia* and male-factor infertility also have been described in association with mild loss-of-function mutations in the androgen receptor.

OTHER DISORDERS AFFECTING 46,XY MALES

Persistent müllerian duct syndrome is the presence of a uterus in an otherwise phenotypic male. This condition can result from mutations in AMH or its receptor (*AMHR2*). The uterus may be removed, but only if damage to the vasa deferentia and blood supply can be avoided. *Isolated hypospadias* occurs in ~1 in 250 males and is usually repaired surgically. Most cases are idiopathic, although evidence of penoscrotal hypospadias, poor phallic development, and/or bilateral cryptorchidism requires investigation for an underlying DSD (e.g., partial gonadal dysgenesis, mild defect in testosterone action, or even severe

forms of 46,XX CAH). Unilateral undescended testes (cryptorchidism) affect more than 3% of boys at birth. Orchidopexy should be considered if the testis has not descended by 6–9 months of age. Bilateral cryptorchidism occurs less frequently and should raise suspicion of gonadotropin deficiency or DSD. A small subset of patients with cryptorchidism may have mutations in the insulin-like 3 (*INSL3*) gene or its receptor *LGR8* (also known as *GREAT*), which mediates normal testicular descent. *Syndromic associations* and *intrauterine growth retardation* also occur relatively frequently in association with impaired testicular function or target tissue responsiveness, but the underlying etiology of many of these conditions is unknown.

46,XX DSD

Inappropriate androgenization of the 46,XX fetus (formerly called *female pseudohermaphroditism*) occurs when the gonad (ovary) contains androgen-secreting testicular material or after increased androgen exposure, which is usually adrenal in origin (Table 410-1).

46,XX Testicular/Ovotesticular DSD Testicular tissue can develop in 46,XX testicular DSD (46,XX males) after translocation of *SRY*, duplication of *SOX9*, or defects in *RSPO1* (Table 410-4).

Increased Androgen Exposure • 21-HYDROXYLASE DEFICIENCY (CONGENITAL ADRENAL HYPERPLASIA)

The classic form of 21-hydroxylase deficiency (21-OHD) is the most common cause of CAH (Chap. 406). It has an incidence between 1 in 10,000 and 1 in 15,000 and is the most common cause of androgenization in chromosomal 46,XX females (Table 410-4). Affected individuals are homozygous or compound heterozygous for severe mutations in the enzyme 21-hydroxylase (*CYP21A2*). This mutation causes a block in adrenal glucocorticoid and mineralocorticoid synthesis, increasing 17-hydroxyprogesterone and shunting steroid precursors into the androgen synthesis pathway (Fig. 410-4). Glucocorticoid insufficiency causes a compensatory elevation of adrenocorticotropic (ACTH), resulting in adrenal hyperplasia and additional synthesis of steroid precursors proximal to the enzymatic block. Increased androgen synthesis in utero causes androgenization of the 46,XX fetus in the first trimester. Ambiguous genitalia are seen at birth, with varying degrees of clitoral enlargement and labial fusion. Excess androgen production causes gonadotropin-independent precocious puberty in males with 21-OHD.

The *salt-wasting* form of 21-OHD results from severe combined glucocorticoid and mineralocorticoid deficiency. A salt-wasting crisis usually manifests between 5 and 21 days of life and is a potentially life-threatening event that requires urgent fluid resuscitation and steroid treatment. Thus, a diagnosis of 21-OHD should be considered in any baby with atypical genitalia with bilateral nonpalpable gonads. Males (46,XY) with 21-OHD have no genital abnormalities at birth but are equally susceptible to adrenal insufficiency and salt-losing crises.

Females with the *classic simple virilizing* form of 21-OHD also present with genital ambiguity. They have impaired cortisol biosynthesis but do not develop salt loss. Patients with *nonclassic 21-OHD* produce normal amounts of cortisol and aldosterone but at the expense of producing excess androgens. Hirsutism (60%), oligomenorrhea (50%), and acne (30%) are the most common presenting features. This is one of the most common recessive disorders in humans, with an incidence as high as 1 in 100 to 500 in many populations and 1 in 27 in Ashkenazi Jews of Eastern European origin.

TABLE 410-4 SELECTED GENETIC CAUSES OF 46,XX DISORDERS OF SEX DEVELOPMENT (DSDs)

Gene	Inheritance	Gonad	Uterus	External Genitalia	Associated Features
Testicular/Ovotesticular DSD					
<i>SRY</i>	Translocation	Testis or ovotestis	–	Male or ambiguous	
<i>SOX9</i>	dup17q24	Unknown	–	Male or ambiguous	
<i>RSP01</i>	AR	Testis or ovotestis	±	Male or ambiguous	Palmar plantar hyperkeratosis, squamous cell skin carcinoma
<i>WNT4</i>	AR	Testis or ovotestis	–	Male or ambiguous	SERKAL syndrome (renal dysgenesis, adrenal and lung hypoplasia)
Increased Androgen Synthesis					
<i>HSD3B2</i>	AR	Ovary	+	Clitoromegaly	CAH, primary adrenal failure, mild androgenization due to ↑ DHEA
<i>CYP21A2</i>	AR	Ovary	+	Ambiguous	CAH, phenotypic spectrum from severe salt-losing forms associated with adrenal failure to simple virilizing forms with compensated adrenal function, ↑ 17-hydroxyprogesterone
<i>POR</i>	AR	Ovary	+	Ambiguous or female	Mixed features of 21-hydroxylase deficiency and 17 α -hydroxylase/17,20-lyase deficiency, sometimes associated with Antley-Bixler craniosynostosis
<i>CYP11B1</i>	AR	Ovary	+	Ambiguous	CAH, hypertension due to ↑ 11-deoxycortisol and 11-deoxycorticosterone
<i>CYP19</i>	AR	Ovary	+	Ambiguous	Maternal virilization during pregnancy, absent breast development at puberty
Glucocorticoid receptor	AR	Ovary	+	Ambiguous	↑ ACTH, 17-hydroxyprogesterone and cortisol; failure of dexamethasone suppression

Abbreviations: ACTH, adrenocorticotropic; AR, autosomal recessive; CAH, congenital adrenal hyperplasia; *CYP11B1*, 11 β -hydroxylase; *CYP19*, aromatase; *CYP21A2*, 21-hydroxylase; DHEA, dehydroepiandrosterone; *HSD3B2*, 3 β -hydroxysteroid dehydrogenase type 2; *POR*, P450 oxidoreductase; *RSP01*, R-spondin 1; *SOX9*, *SRY*-related HMG-box gene 9; *SRY*, sex-related gene on the Y chromosome.

Biochemical features of acute salt-wasting 21-OHD are hyponatremia, hyperkalemia, hypoglycemia, inappropriately low cortisol and aldosterone, and elevated 17-hydroxyprogesterone, ACTH, and plasma renin activity. Presymptomatic diagnosis of classic 21-OHD is now made by neonatal screening tests for increased 17-hydroxyprogesterone in many centers. In most cases, 17-hydroxyprogesterone is markedly increased. In adults, ACTH stimulation (0.25 mg of cosyntropin IV) with assays for 17-hydroxyprogesterone at 0 and 30 min can be useful for detecting nonclassic 21-OHD and heterozygotes (Chap. 406).

TREATMENT CONGENITAL ADRENAL HYPERPLASIA

Acute salt-wasting crises require fluid resuscitation, IV hydrocortisone, and correction of hypoglycemia. Once the patient is stabilized, glucocorticoids must be given to correct the cortisol insufficiency and suppress ACTH stimulation, thereby preventing further virilization, rapid skeletal maturation, and the development of polycystic ovaries. Typically, hydrocortisone (10–15 mg/m² per day in three divided doses) is used in childhood with a goal of partially suppressing 17-hydroxyprogesterone (100 to <1000 ng/dL). The aim of treatment is to use the lowest glucocorticoid dose that adequately suppresses adrenal androgen production without causing signs of glucocorticoid excess such as impaired growth and obesity. Salt-wasting conditions are treated with mineralocorticoid replacement. Infants usually need salt supplements up to the first year of life. Plasma renin activity and electrolytes are used to monitor mineralocorticoid replacement. Some patients with simple virilizing 21-OHD also benefit from mineralocorticoid supplements. Parents and patients should be educated about the need for increased doses of steroids during sickness, and patients should carry medic alert systems.

Steroid treatment for older adolescents and adults varies depending on lifestyle, age, and factors such as a desire to optimize fertility. Hydrocortisone remains a useful approach, but treatment with prednisolone at night may provide more complete ACTH suppression. Steroid doses should be adjusted to individual requirements because overtreatment can result in iatrogenic Cushing's-like features, including weight gain, insulin resistance, hypertension,

and osteopenia. Because it is long acting, dexamethasone given at night is useful for ACTH suppression but is often associated with more side effects, making hydrocortisone or prednisolone preferable for most patients. Androstenedione and testosterone may be useful measurements of long-term control, with less fluctuation than 17-hydroxyprogesterone. Mineralocorticoid requirements often decrease in adulthood, and doses should be reassessed and reduced to avoid hypertension in adults. In very severe cases, adrenalectomy has been advocated but incurs the risks of surgery and total adrenal insufficiency.

Girls with significant genital androgenization due to classic 21-OHD usually undergo vaginal reconstruction and sometimes clitoral reduction (maintaining the glans and nerve supply), but the optimal timing of these procedures is debated, as is the need for the individual to be able to consent. There is a higher threshold for undertaking clitoral surgery in some centers because long-term sensation and ability to achieve orgasm can be affected, but the long-term results of newer techniques are not yet known. Full information about all options should be provided. If surgery is performed in infancy, surgical revision or regular vaginal dilatation may be needed in adolescence or adulthood, and long-term psychological support and psychosexual counseling may be appropriate. Women with 21-OHD frequently develop polycystic ovaries and have reduced fertility, especially when control is poor. Fecundity is achieved in 60–90% of women with good metabolic control, but ovulation induction (or even adrenalectomy) may be required. Dexamethasone should be avoided in pregnancy. Men with poorly controlled 21-OHD may develop testicular adrenal rests and are at risk for reduced fertility. Prenatal treatment of 21-OHD by the administration of dexamethasone to mothers is still under evaluation. However, pending methods to diagnose the disorder early in pregnancy, both affected and nonaffected fetuses will be exposed because treatment is started ideally before 6 to 7 weeks. The long-term effects of prenatal dexamethasone exposure on fetal development are still under evaluation, and current guidelines recommend full informed consent before treatment, ideally in a protocol that allows long-term follow-up of all children treated. Newer techniques such as cell-free fetal DNA testing may potentially reduce treatment of nonaffected fetuses.

The treatment of other forms of CAH includes mineralocorticoid and glucocorticoid replacement for salt-losing conditions (e.g., *StAR*, *CYP11A1*, *HSD3 β 2*), suppression of ACTH drive with glucocorticoids in disorders associated with hypertension (e.g., *CYP17*, *CYP11B1*), and appropriate sex hormone replacement in adolescence and adulthood, when necessary.

OTHER CAUSES Increased androgen synthesis can also occur in CAH due to defects in *POR*, *11 β -hydroxylase* (*CYP11B1*), and *3 β -hydroxysteroid dehydrogenase type 2* (*HSD3B2*) and with mutations in the genes encoding *aromatase* (*CYP19*) and the glucocorticoid receptor. Increased androgen exposure in utero can occur with maternal virilizing tumors and with ingestion of androgenic compounds.

OTHER DISORDERS AFFECTING 46,XX FEMALES

Congenital absence of the vagina occurs in association with *müllerian agenesis* or *hypoplasia* as part of the Mayer-Rokitansky-Kuster-Hauser (MRKH) syndrome (rarely caused by *WNT4* mutations). This diagnosis should be considered in otherwise phenotypically normal females with primary amenorrhea. Associated features include renal (agenesis) and cervical spinal abnormalities.

GLOBAL CONSIDERATIONS



The approach to a child or adolescent with ambiguous genitalia or another DSD requires cultural sensitivity, as the concepts of sex and gender vary widely. Rare genetic DSDs can occur more frequently in specific populations (e.g., *5 α -reductase type 2* in the Dominican Republic). Different forms of CAH also show ethnic and geographic variability. In many countries, appropriate biochemical tests may not be readily available, and access to appropriate forms of treatment and support may be limited.

411 Disorders of the Testes and Male Reproductive System

Shalender Bhasin, J. Larry Jameson

The male reproductive system regulates sex differentiation, virilization, and the hormonal changes that accompany puberty, ultimately leading to spermatogenesis and fertility. Under the control of the pituitary hormones—luteinizing hormone (LH) and follicle-stimulating hormone (FSH)—the Leydig cells of the testes produce testosterone and germ cells are nurtured by Sertoli cells to divide, differentiate, and mature into sperm. During embryonic development, testosterone and dihydrotestosterone (DHT) induce the wolffian duct and virilization of the external genitalia. During puberty, testosterone promotes somatic growth and the development of secondary sex characteristics. In the adult, testosterone is necessary for spermatogenesis, stimulation of libido and normal sexual function, and maintenance of muscle and bone mass. This chapter focuses on the physiology of the testes and disorders associated with decreased androgen production, which may be caused by gonadotropin deficiency or by primary testis dysfunction. A variety of testosterone formulations now allow more physiologic androgen replacement. Infertility occurs in ~5% of men and is increasingly amenable to treatment by hormone replacement or by using sperm transfer techniques. **For further discussion of sexual dysfunction, disorders of the prostate, and testicular cancer, see Chaps. 67, 115, and 116, respectively.**

DEVELOPMENT AND STRUCTURE OF THE TESTIS

The fetal testis develops from the undifferentiated gonad after expression of a genetic cascade that is initiated by the *SRY* (sex-related gene on the Y chromosome) (**Chap. 410**). *SRY* induces differentiation of

Sertoli cells, which surround germ cells and, together with peritubular myoid cells, form testis cords that will later develop into seminiferous tubules. Fetal Leydig cells and endothelial cells migrate into the gonad from the adjacent mesonephros but may also arise from interstitial cells that reside between testis cords. Leydig cells produce testosterone, which supports the growth and differentiation of wolffian duct structures that develop into the epididymis, vas deferens, and seminal vesicles. Testosterone is also converted to DHT (see below), which induces formation of the prostate and the external male genitalia, including the penis, urethra, and scrotum. Testicular descent through the inguinal canal is controlled in part by Leydig cell production of insulin-like factor 3 (*INSL3*), which acts via a receptor termed *Great* (*G* protein-coupled receptor affecting testis descent). Sertoli cells produce müllerian-inhibiting substance (*MIS*), which causes regression of the müllerian structures, including the fallopian tube, uterus, and upper segment of the vagina.

NORMAL MALE PUBERTAL DEVELOPMENT

Although *puberty* commonly refers to the maturation of the reproductive axis and the development of secondary sex characteristics, it involves a coordinated response of multiple hormonal systems including the adrenal gland and the growth hormone (GH) axis (**Fig. 411-1**). The development of secondary sex characteristics is initiated by *adrenarche*, which usually occurs between 6 and 8 years of age when the adrenal gland begins to produce greater amounts of androgens from the zona reticularis, the principal site of dehydroepiandrosterone (DHEA) production. The sex maturation process is greatly accelerated by the activation of the hypothalamic-pituitary axis and the production of gonadotropin-releasing hormone (GnRH). The GnRH pulse generator in the hypothalamus is active during fetal life and early infancy but is restrained until the early stages of puberty by a neuroendocrine brake imposed by the inhibitory actions of glutamate, γ -aminobutyric acid (GABA), and neuropeptide Y. Although the pathways that initiate reactivation of the GnRH pulse generator at the onset of puberty have been elusive, mounting evidence supports involvement of GPR54, a G protein-coupled receptor that binds an endogenous ligand, kisspeptin. Individuals with mutations of GPR54 fail to enter puberty, and experiments in primates demonstrate that infusion of the ligand is sufficient to induce premature puberty. Kisspeptin signaling plays an important role in mediating the feedback action of sex steroids on gonadotropin secretion and in regulating the tempo of sexual maturation at puberty. Leptin, a hormone produced by adipose cells, plays a permissive role in the resurgence of GnRH secretion at the onset of puberty, as leptin-deficient individuals also fail to enter puberty (**Chap. 415e**). The adipocyte hormone leptin, gut hormone ghrelin, neuropeptide Y, and kisspeptin integrate the signals

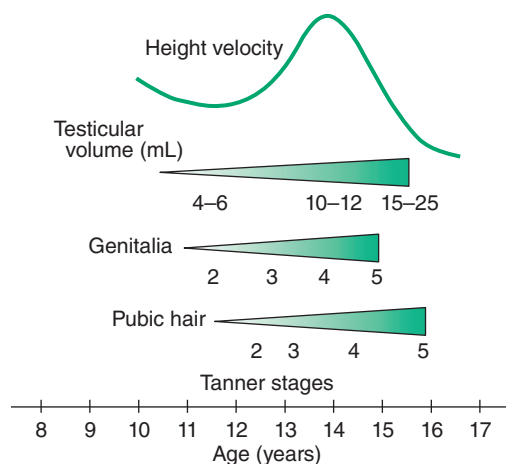


FIGURE 411-1 Pubertal events in males. Sexual maturity ratings for genitalia and pubic hair and divided into five stages. (From WA Marshall, JM Tanner: *Variations in the pattern of pubertal changes in boys*. *Arch Dis Child* 45:13, 1970.)

2358 originating in energy stores and metabolic tissues with mechanisms that control onset of puberty through regulation of GnRH secretion. Energy deficit and excess and metabolic stress are associated with disturbed reproductive maturation and timing of pubertal onset.

The early stages of puberty are characterized by nocturnal surges of LH and FSH. Growth of the testes is usually the first sign of puberty, reflecting an increase in seminiferous tubule volume. Increasing levels of testosterone deepen the voice and increase muscle growth. Conversion of testosterone to DHT leads to growth of the external genitalia and pubic hair. DHT also stimulates prostate and facial hair growth and initiates recession of the temporal hairline. The growth spurt occurs at a testicular volume of about 10–12 mL. GH increases early in puberty and is stimulated in part by the rise in gonadal steroids. GH increases the level of insulin-like growth factor I (IGF-I), which enhances linear bone growth. The prolonged pubertal exposure to gonadal steroids (mainly estradiol) ultimately causes epiphyseal closure and limits further bone growth.

REGULATION OF TESTICULAR FUNCTION

REGULATION OF THE HYPOTHALAMIC-PITUITARY-TESTIS AXIS IN ADULT MAN

Hypothalamic GnRH regulates the production of the pituitary gonadotropins LH and FSH (Fig. 411-2). GnRH is released in discrete pulses approximately every 2 h, resulting in corresponding pulses of LH and FSH. These dynamic hormone pulses account in part for the wide variations in LH and testosterone, even within the same individual. LH acts primarily on the Leydig cell to stimulate testosterone synthesis. The regulatory control of androgen synthesis is mediated by testosterone and estrogen feedback on both the hypothalamus and the pituitary. FSH acts on the Sertoli cell to regulate spermatogenesis and the production of Sertoli products such as inhibin B, which acts to selectively suppress pituitary FSH. Despite these somewhat distinct Leydig and Sertoli cell-regulated pathways, testis function is integrated at several levels: GnRH regulates both gonadotropins; spermatogenesis requires high levels of testosterone; and numerous paracrine interactions between Leydig and Sertoli cells are necessary for normal testis function.

THE LEYDIG CELL: ANDROGEN SYNTHESIS

LH binds to its seven-transmembrane, G protein-coupled receptor to activate the cyclic AMP pathway. Stimulation of the LH receptor induces steroid acute regulatory (StAR) protein, along with several steroidogenic enzymes involved in androgen synthesis. LH receptor mutations cause Leydig cell hypoplasia or agenesis, underscoring the importance of this pathway for Leydig cell development and function. The rate-limiting process in testosterone synthesis is the delivery of cholesterol by the StAR protein to the inner mitochondrial membrane. Peripheral benzodiazepine receptor, a mitochondrial cholesterol-binding protein, is also an acute regulator of Leydig cell steroidogenesis. The five major enzymatic steps involved in testosterone synthesis are summarized in Fig. 411-3. After cholesterol transport into the mitochondrion, the formation of pregnenolone by CYP11A1 (side chain cleavage enzyme) is a limiting enzymatic step. The 17 α -hydroxylase and the 17,20-lyase reactions are catalyzed by a single enzyme, CYP17; posttranslational modification (phosphorylation) of this enzyme and the presence of specific enzyme cofactors confer 17,20-lyase activity selectively in the testis and zona reticularis of the adrenal gland. Testosterone can be converted to the more potent DHT by 5 α -reductase, or it can be aromatized to estradiol by CYP19 (aromatase). Two isoforms of steroid 5 α -reductase, SRD5A1 and SRD5A2, have been described; all known kindreds with 5 α -reductase deficiency have had mutations in SRD5A2, the predominant form in the prostate and the skin.

Testosterone Transport and Metabolism In males, 95% of circulating testosterone is derived from testicular production (3–10 mg/d). Direct secretion of testosterone by the adrenal and the peripheral conversion of androstenedione to testosterone collectively account for another 0.5 mg/d of testosterone. Only a small amount of DHT (70 μ g/d) is secreted directly by the testis; most circulating DHT is derived from

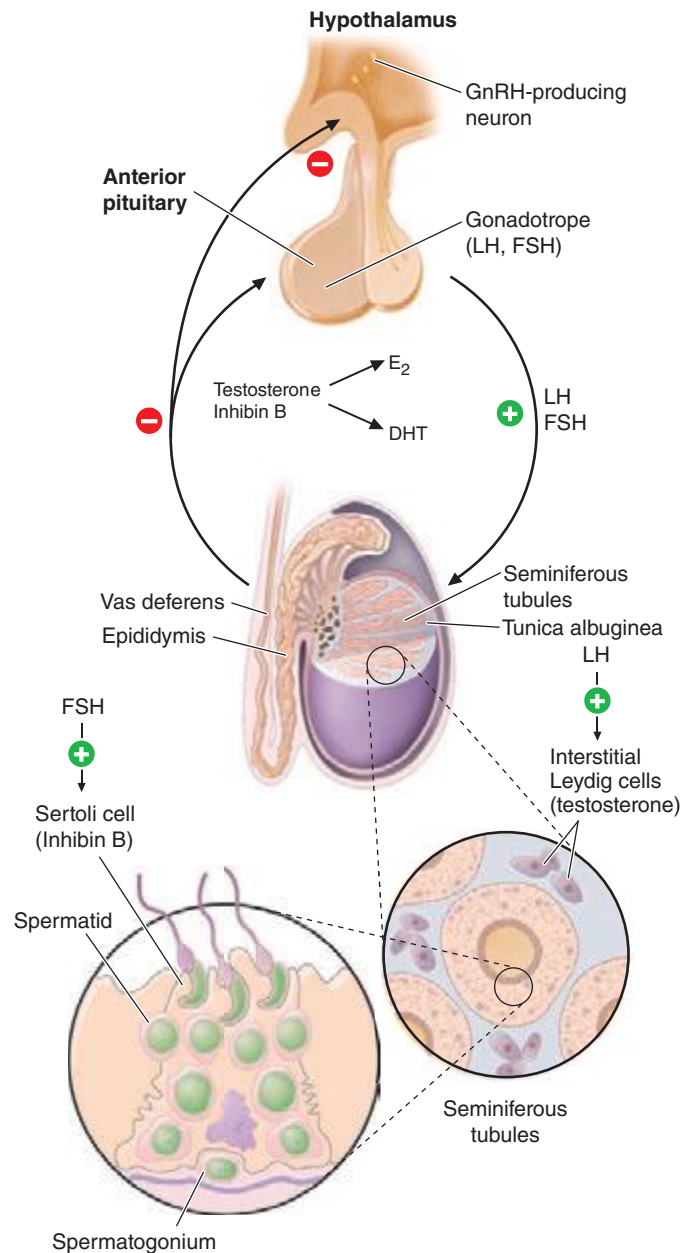


FIGURE 411-2 Human pituitary gonadotropin axis, structure of testis, and seminiferous tubule. E₂, 17 β -estradiol; DHT, dihydrotestosterone; FSH, follicle-stimulating hormones; GnRH, gonadotropin-releasing; LH, luteinizing hormone.

peripheral conversion of testosterone. Most of the daily production of estradiol (~45 μ g/d) in men is derived from aromatase-mediated peripheral conversion of testosterone and androstenedione.

Circulating testosterone is bound to two plasma proteins: sex hormone-binding globulin (SHBG) and albumin (Fig. 411-4). SHBG binds testosterone with much greater affinity than albumin. Only 0.5–3% of testosterone is unbound. According to the “free hormone” hypothesis, only the unbound fraction is biologically active; however, albumin-bound hormone dissociates readily in the capillaries and may be bioavailable. SHBG-bound testosterone also may be internalized through endocytic pits by binding to a protein called megalin. SHBG concentrations are decreased by androgens, obesity, diabetes mellitus, insulin, and nephrotic syndrome. Conversely, estrogen administration, hyperthyroidism, many chronic inflammatory illnesses, infections such as HIV or hepatitis B and C, and aging are associated with high SHBG concentrations.

Testosterone is metabolized predominantly in the liver, although some degradation occurs in peripheral tissues, particularly the

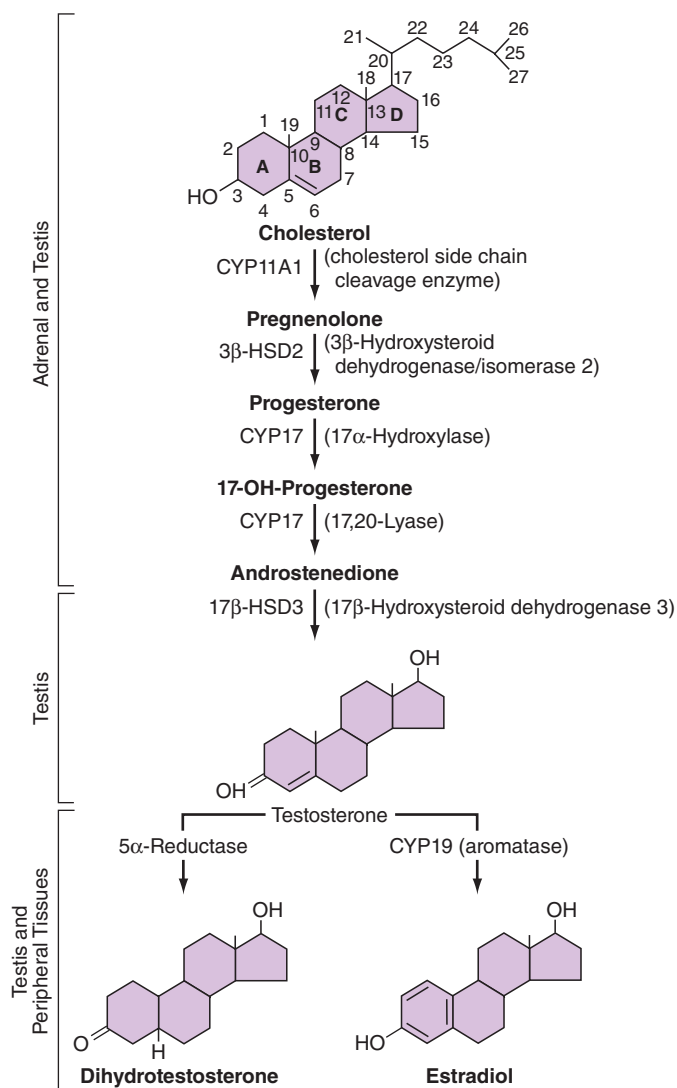


FIGURE 411-3 The biochemical pathway in the conversion of 27-carbon sterol cholesterol to androgens and estrogens.

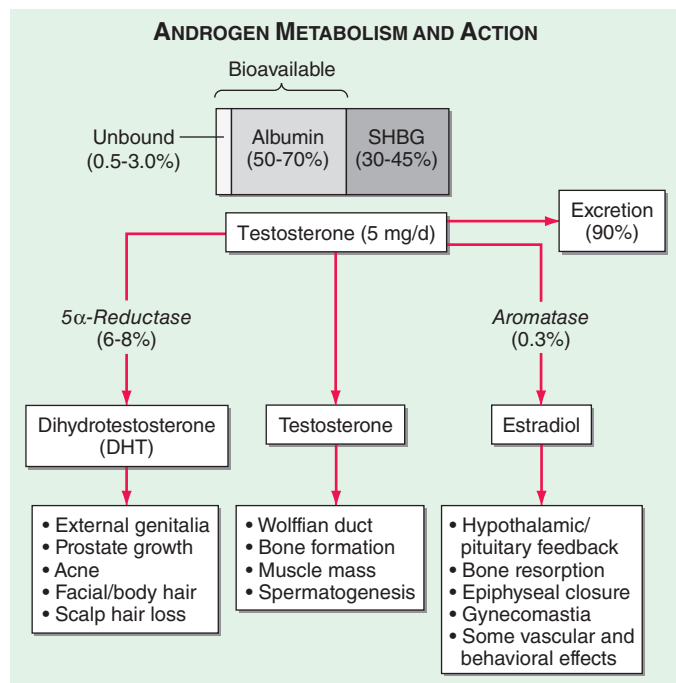


FIGURE 411-4 Androgen metabolism and actions. SHBG, sex hormone-binding globulin.

prostate and the skin. In the liver, testosterone is converted by a series of enzymatic steps that involve 5α- and 5β-reductases, 3α- and 3β-hydroxysteroid dehydrogenases, and 17β-hydroxysteroid dehydrogenase into androsterone, etiocholanolone, DHT, and 3-α-androstenediol. These compounds undergo glucuronidation or sulfation before being excreted by the kidneys.

Mechanism of Androgen Action Testosterone exerts some of its biologic effects by binding to androgen receptor, either directly or after its conversion to DHT by the steroid 5-α reductase. Testosterone's effects on the skeletal muscle, erythropoiesis, and bone in men do not require its obligatory conversion to DHT. However, the conversion of testosterone to DHT is necessary for the masculinization of the urogenital sinus and genital tubercle. Aromatization of testosterone to estradiol mediates additional effects of testosterone on the bone resorption, epiphyseal closure, sexual desire, vascular endothelium, and fat. DHT can also be converted in some tissues by 3-keto reductase/3β-hydroxysteroid dehydrogenase enzymes to 5α-androstane-3β,17β-diol, which is a high-affinity ligand and agonist of estrogen receptor β.

The androgen receptor (AR) is structurally related to the nuclear receptors for estrogen, glucocorticoids, and progesterone (Chap. 400e). The AR is encoded by a gene on the long arm of the X chromosome and has a molecular mass of about 110 kDa. A polymorphic region in the amino terminus of the receptor, which contains a variable number of glutamine repeats, modifies the transcriptional activity of the receptor. The AR protein is distributed in both the cytoplasm and the nucleus. The ligand binding to the AR induces conformational changes that allow the recruitment and assembly of tissue-specific cofactors and causes it to translocate into the nucleus, where it binds to DNA or other transcription factors already bound to DNA. Thus, the AR is a ligand-regulated transcription factor that regulates the expression of androgen-dependent genes in a tissue-specific manner. Some androgen effects may be mediated by nongenomic AR signal transduction pathways. Testosterone binds to AR with half the affinity of DHT. The DHT-AR complex also has greater thermostability and a slower dissociation rate than the testosterone-AR complex. However, the molecular basis for selective testosterone versus DHT actions remains incompletely explained.

THE SEMINIFEROUS TUBULES: SPERMATOGENESIS

The seminiferous tubules are convoluted, closed loops with both ends emptying into the rete testis, a network of progressively larger efferent ducts that ultimately form the epididymis (Fig. 411-2). The seminiferous tubules total about 600 m in length and comprise about two-thirds of testis volume. The walls of the tubules are formed by polarized Sertoli cells that are apposed to peritubular myoid cells. Tight junctions between Sertoli cells create a blood-testis barrier. Germ cells compose the majority of the seminiferous epithelium (~60%) and are intimately embedded within the cytoplasmic extensions of the Sertoli cells, which function as “nurse cells.” Germ cells progress through characteristic stages of mitotic and meiotic divisions. A pool of type A spermatogonia serve as stem cells capable of self-renewal. Primary spermatocytes are derived from type B spermatogonia and undergo meiosis before progressing to spermatids that undergo spermiogenesis (a differentiation process involving chromatin condensation, acquisition of an acrosome, elongation of cytoplasm, and formation of a tail) and are released from Sertoli cells as mature spermatozoa. The complete differentiation process into mature sperm requires 74 days. Peristaltic-type action by peritubular myoid cells transports sperm into the efferent ducts. The spermatozoa spend an additional 21 days in the epididymis, where they undergo further maturation and capacitation. The normal adult testes produce >100 million sperm per day.

Naturally occurring mutations in the *FSHβ* gene and in the FSH receptor confirm an important, but not essential, role for this pathway in spermatogenesis. Females with these mutations are hypogonadal and infertile because ovarian follicles do not mature; males exhibit variable degrees of reduced spermatogenesis, presumably because of impaired Sertoli cell function. Because Sertoli cells produce inhibin B, an inhibitor of FSH, seminiferous tubule damage (e.g., by radiation)

2360 causes a selective increase of FSH. Testosterone reaches very high concentrations locally in the testis and is essential for spermatogenesis. The cooperative actions of FSH and testosterone are important in the progression of meiosis and spermiation. FSH and testosterone regulate germ cell survival via the intrinsic and the extrinsic apoptotic mechanisms. FSH may also play an important role in supporting spermatogonia. Gonadotropin-regulated testicular RNA helicase (GRTH/DDX25), a testis-specific gonadotropin/androgen-regulated RNA helicase, is present in germ cells and Leydig cells and may be an important factor in the paracrine regulation of germ cell development. Several cytokines and growth factors are also involved in the regulation of spermatogenesis by paracrine and autocrine mechanisms. A number of knockout mouse models exhibit impaired germ cell development or spermatogenesis, presaging possible mutations associated with male infertility. The human Y chromosome contains a small pseudoautosomal region that can recombine with homologous regions of the X chromosome. Most of the Y chromosome does not recombine with the X chromosome and is referred to as the male-specific region of the Y (MSY). The MSY contains 156 transcription units that encode for 26 proteins, including nine families of Y-specific multicopy genes; many of these Y-specific genes are also testis-specific and necessary for spermatogenesis. Microdeletions of several Y chromosome azoospermia factor (AZF) genes (e.g., RNA-binding motif, *RBM*; deleted in azoospermia, *DAZ*) are associated with oligospermia or azoospermia.

TREATMENT MALE FACTOR INFERTILITY

Treatment options for male factor infertility have expanded greatly in recent years. Secondary hypogonadism is highly amenable to treatment with pulsatile GnRH or gonadotropins (see below). Assisted reproductive technologies such as the in vitro fertilization (IVF) and intracytoplasmic sperm injection (ICSI) have provided new opportunities for patients with primary testicular failure and disorders of sperm transport. Choice of initial treatment options depends on sperm concentration and motility. Expectant management should be attempted initially in men with mild male factor infertility (sperm count of $15\text{--}20 \times 10^6/\text{mL}$ and normal motility). Moderate male factor infertility ($10\text{--}15 \times 10^6/\text{mL}$ and $20\text{--}40\%$ motility) should begin with intrauterine insemination alone or in combination with treatment of the female partner with clomiphene or gonadotropins, but it may require IVF with or without ICSI. For men with a severe defect (sperm count of $<10 \times 10^6/\text{mL}$, 10% motility), IVF with ICSI or donor sperm should be used.

CLINICAL AND LABORATORY EVALUATION OF MALE REPRODUCTIVE FUNCTION

HISTORY AND PHYSICAL EXAMINATION

The history should focus on developmental stages such as puberty and growth spurts, as well as androgen-dependent events such as early morning erections, frequency and intensity of sexual thoughts, and frequency of masturbation or intercourse. Although libido and the overall frequency of sexual acts are decreased in androgen-deficient men, young hypogonadal men may achieve erections in response to visual erotic stimuli. Men with acquired androgen deficiency often report decreased energy and increased irritability.

The physical examination should focus on secondary sex characteristics such as hair growth, gynecomastia, testicular volume, prostate, and height and body proportions. *Eunuchoid proportions* are defined as an arm span >2 cm greater than height and suggest that androgen deficiency occurred before epiphyseal fusion. Hair growth in the face, axilla, chest, and pubic regions is androgen-dependent; however, changes may not be noticeable unless androgen deficiency is severe and prolonged. Ethnicity also influences the intensity of hair growth (**Chap. 68**). Testicular volume is best assessed by using a Prader orchidometer. Testes range from 3.5 to 5.5 cm in length, which corresponds to a volume of 12–25 mL. Advanced age does not influence testicular size, although the consistency becomes less firm. Asian men generally

have smaller testes than Western Europeans, independent of differences in body size. Because of its possible role in infertility, the presence of varicocele should be sought by palpation while the patient is standing; it is more common on the left side. Patients with Klinefelter's syndrome have markedly reduced testicular volumes (1–2 mL). In congenital hypogonadotropic hypogonadism, testicular volumes provide a good index for the degree of gonadotropin deficiency and the likelihood of response to therapy.

GONADOTROPIN AND INHIBIN MEASUREMENTS

LH and FSH are measured using two-site immunoradiometric, immunofluorometric, or chemiluminescent assays, which have very low cross-reactivity with other pituitary glycoprotein hormones and human chorionic gonadotropin (hCG) and have sufficient sensitivity to measure the low levels present in patients with hypogonadotropic hypogonadism. In men with a low testosterone level, an LH level can distinguish primary (high LH) versus secondary (low or inappropriately normal LH) hypogonadism. An elevated LH level indicates a primary defect at the testicular level, whereas a low or inappropriately normal LH level suggests a defect at the hypothalamic-pituitary level. LH pulses occur about every 1–3 h in normal men. Thus, gonadotropin levels fluctuate, and samples should be pooled or repeated when results are equivocal. FSH is less pulsatile than LH because it has a longer half-life. Selective increase in FSH suggests damage to the seminiferous tubules. Inhibin B, a Sertoli cell product that suppresses FSH, is reduced with seminiferous tubule damage. Inhibin B is a dimer with $\alpha\text{-}\beta_{\text{B}}$ subunits and is measured by two-site immunoassays.

GnRH Stimulation Testing The GnRH test is performed by measuring LH and FSH concentrations at baseline and at 30 and 60 min after intravenous administration of 100 μg of GnRH. A minimally acceptable response is a twofold LH increase and a 50% FSH increase. In the prepubertal period or with severe GnRH deficiency, the gonadotrope may not respond to a single bolus of GnRH because it has not been primed by endogenous hypothalamic GnRH; in these patients, GnRH responsiveness may be restored by chronic, pulsatile GnRH administration. With the availability of sensitive and specific LH assays, GnRH stimulation testing is used rarely except to evaluate gonadotrope function in patients who have undergone pituitary surgery or have a space-occupying lesion in the hypothalamic-pituitary region.

TESTOSTERONE ASSAYS

Total Testosterone Total testosterone includes both unbound and protein-bound testosterone and is measured by radioimmunoassays, immunometric assays, or liquid chromatography tandem mass spectrometry (LC-MS/MS). LC-MS/MS involves extraction of serum by organic solvents, separation of testosterone from other steroids by high-performance liquid chromatography and mass spectrometry, and quantitation of unique testosterone fragments by mass spectrometry. LC-MS/MS provides accurate and sensitive measurements of testosterone levels even in the low range and is emerging as the method of choice for testosterone measurement. Laboratories that have been certified by the Centers for Disease Control and Prevention (CDC) Hormone Standardization Program for Testosterone (HoST) can ensure that testosterone measurements are accurate and calibrated to an international standard. A single fasting morning sample provides a good approximation of the average testosterone concentration with the realization that testosterone levels fluctuate in response to pulsatile LH. Testosterone is generally lower in the late afternoon and is reduced by acute illness. The testosterone concentration in healthy young men ranges from 300 to 1000 ng/dL in most laboratories, and efforts are under way to generate harmonized population-based reference ranges that can be applied to all CDC-certified laboratories. Alterations in SHBG levels due to aging, obesity, diabetes mellitus, hyperthyroidism, some types of medications, or chronic illness or on a congenital basis can affect total testosterone levels. Heritable factors contribute substantially to the population-level variation in testosterone levels, and genome-wide association studies have revealed polymorphisms in the *SHBG* gene as important contributors to variation in testosterone levels.

Measurement of Unbound Testosterone Levels Most circulating testosterone is bound to SHBG and to albumin; only 0.5–3% of circulating testosterone is unbound, or “free.” The unbound testosterone concentration can be measured by equilibrium dialysis or calculated from total testosterone, SHBG, and albumin concentrations. Recent research has shown that testosterone binding to SHBG is a multistep process that involves complex homoallostery within the SHBG dimer; a novel allosteric model of testosterone binding to SHBG dimers provides good estimates of free testosterone concentrations. The previous law of mass action equations based on linear models of testosterone binding to SHBG have been shown to be erroneous. Tracer analogue methods are relatively inexpensive and convenient, but they are inaccurate. *Bioavailable testosterone* refers to unbound testosterone plus testosterone that is loosely bound to albumin; it can be determined by the ammonium sulfate precipitation method.

hCG Stimulation Test The hCG stimulation test is performed by administering a single injection of 1500–4000 IU of hCG intramuscularly and measuring testosterone levels at baseline and 24, 48, 72, and 120 h after hCG injection. An alternative regimen involves three injections of 1500 units of hCG on successive days and measuring testosterone levels 24 h after the last dose. An acceptable response to hCG is a doubling of the testosterone concentration in adult men. In prepubertal boys, an increase in testosterone to >150 ng/dL indicates the presence of testicular tissue. No response may indicate an absence of testicular tissue or marked impairment of Leydig cell function. Measurement of MIS, a Sertoli cell product, is also used to detect the presence of testes in prepubertal boys with cryptorchidism.

SEMEN ANALYSIS

Semen analysis is the most important step in the evaluation of male infertility. Samples are collected by masturbation following a period of abstinence for 2–3 days. Semen volumes and sperm concentrations vary considerably among fertile men, and several samples may be needed before concluding that the results are abnormal. Analysis should be performed within an hour of collection. Using semen samples from over 4500 men in 14 countries, whose partners had a time-to-pregnancy of less than 12 months, the World Health Organization (WHO) has generated the following one-sided reference limits for semen parameters: semen volume, 1.5 mL; total sperm number, 39 million per ejaculate; sperm concentration, 15 million/mL; vitality, 58% live; progressive motility, 32%; total (progressive + nonprogressive) motility, 40%; morphologically normal forms, 4.0%. Some men with low sperm counts are nevertheless fertile. A variety of tests for sperm function can be performed in specialized laboratories, but these add relatively little to the treatment options.

TESTICULAR BIOPSY

Testicular biopsy is useful in some patients with oligospermia or azospermia as an aid in diagnosis and indication for the feasibility of treatment. Using local anesthesia, fine-needle aspiration biopsy is performed to aspirate tissue for histology. Alternatively, open biopsies can be performed under local or general anesthesia when more tissue is required. A normal biopsy in an azospermic man with a normal FSH level suggests obstruction of the vas deferens, which may be correctable surgically. Biopsies are also used to harvest sperm for ICSI and to classify disorders such as hypospermatogenesis (all stages present but in reduced numbers), germ cell arrest (usually at primary spermatocyte stage), and Sertoli cell–only syndrome (absent germ cells) or hyalinization (sclerosis with absent cellular elements).

DISORDERS OF SEXUAL DIFFERENTIATION

See Chap. 410.

DISORDERS OF PUBERTY

The onset and tempo of puberty varies greatly in the general population and is affected by genetic and environmental factors. Although some of the variance in the timing of puberty is explained by heritable factors, the genes involved remain unknown.

PRECOCIOUS PUBERTY

Puberty in boys before age 9 is considered precocious. *Isosexual precocity* refers to premature sexual development consistent with phenotypic sex and includes features such as the development of facial hair and phallic growth. Isosexual precocity is divided into gonadotropin-dependent and gonadotropin-independent causes of androgen excess (**Table 411-1**). *Heterosexual precocity* refers to the premature development of estrogenic features in boys, such as breast development.

Gonadotropin-Dependent Precocious Puberty This disorder, called *central precocious puberty* (CPP), is less common in boys than in girls. It is caused by premature activation of the GnRH pulse generator, sometimes because of central nervous system (CNS) lesions such as hypothalamic hamartomas, but it is often idiopathic. CPP is characterized by gonadotropin levels that are inappropriately elevated for age. Because pituitary priming has occurred, GnRH elicits LH and FSH responses typical of those seen in puberty or in adults. Magnetic resonance imaging (MRI) should be performed to exclude a mass, structural defect, infection, or inflammatory process. Mutations in *MKRN3*, an imprinted gene encoding makorin ring-finger protein 3, which is expressed only from the paternally inherited allele, have been associated with CPP.

Gonadotropin-Independent Precocious Puberty In gonadotropin-independent precocious puberty, androgens from the testis or the adrenal are increased, but gonadotropins are low. This group of disorders includes hCG-secreting tumors; congenital adrenal hyperplasia; sex steroid-producing tumors of the testis, adrenal, and ovary; accidental or deliberate exogenous sex steroid administration; hypothyroidism; and activating mutations of the LH receptor or $G_s\alpha$ subunit.

TABLE 411-1 CAUSES OF PRECOCIOUS OR DELAYED PUBERTY IN BOYS

- I. Precocious puberty
 - A. Gonadotropin-dependent
 1. Idiopathic
 2. Hypothalamic hamartoma or other lesions
 3. CNS tumor or inflammatory state
 - B. Gonadotropin-independent
 1. Congenital adrenal hyperplasia
 2. hCG-secreting tumor
 3. McCune-Albright syndrome
 4. Activating LH receptor mutation
 5. Exogenous androgens
- II. Delayed puberty
 - A. Constitutional delay of growth and puberty
 - B. Systemic disorders
 1. Chronic disease
 2. Malnutrition
 3. Anorexia nervosa
 - C. CNS tumors and their treatment (radiotherapy and surgery)
 - D. Hypothalamic-pituitary causes of pubertal failure (low gonadotropins)
 1. Congenital disorders (see Table 411-2)
 2. Acquired disorders
 - a. Pituitary tumors
 - b. Hyperprolactinemia
 - E. Gonadal causes of pubertal failure (elevated gonadotropins)
 1. Klinefelter’s syndrome
 2. Bilateral undescended testes
 3. Orchitis
 4. Chemotherapy or radiotherapy
 5. Anorchia
 - F. Androgen insensitivity

Abbreviations: CNS, central nervous system; GnRH, gonadotropin-releasing hormone; hCG, human chorionic gonadotropin; LH, luteinizing hormone.

2362 FAMILIAL MALE-LIMITED PRECOCIOUS PUBERTY Also called *testotoxicosis*, familial male-limited precocious puberty is an autosomal dominant disorder caused by activating mutations in the LH receptor, leading to constitutive stimulation of the cyclic AMP pathway and testosterone production. Clinical features include premature androgenization in boys, growth acceleration in early childhood, and advanced bone age followed by premature epiphyseal fusion. Testosterone is elevated, and LH is suppressed. Treatment options include inhibitors of testosterone synthesis (e.g., ketoconazole), AR antagonists (e.g., flutamide and bicalutamide), and aromatase inhibitors (e.g., anastrozole).

MCCUNE-ALBRIGHT SYNDROME This is a sporadic disorder caused by somatic (postzygotic) activating mutations in the G_{α} subunit that links G protein-coupled receptors to intracellular signaling pathways (**Chap. 426e**). The mutations impair the guanosine triphosphatase activity of the G_{α} protein, leading to constitutive activation of adenylyl cyclase. Like activating LH receptor mutations, this stimulates testosterone production and causes gonadotropin-independent precocious puberty. In addition to sexual precocity, affected individuals may have autonomy in the adrenals, pituitary, and thyroid glands. Café au lait spots are characteristic skin lesions that reflect the onset of the somatic mutations in melanocytes during embryonic development. Polyostotic fibrous dysplasia is caused by activation of the parathyroid hormone receptor pathway in bone. Treatment is similar to that in patients with activating LH receptor mutations. Bisphosphonates have been used to treat bone lesions.

CONGENITAL ADRENAL HYPERPLASIA Boys with congenital adrenal hyperplasia (CAH) who are not well controlled with glucocorticoid suppression of adrenocorticotrophic hormone (ACTH) can develop premature virilization because of excessive androgen production by the adrenal gland (**Chaps. 406 and 410**). LH is low, and the testes are small. Adrenal rests may develop within the testis of poorly controlled patients with CAH because of chronic ACTH stimulation; adrenal rests do not require surgical removal and regress with effective glucocorticoid therapy. Some children with CAH may develop gonadotropin-dependent precocious puberty with early maturation of the hypothalamic-pituitary-gonadal axis, elevated gonadotropins, and testicular growth.

Heterosexual Sexual Precocity Breast enlargement in prepubertal boys can result from familial aromatase excess, estrogen-producing tumors in the adrenal gland, Sertoli cell tumors in the testis, marijuana smoking, or exogenous estrogens or androgens. Occasionally, germ cell tumors that secrete hCG can be associated with breast enlargement due to excessive stimulation of estrogen production (see “Gynecomastia,” below).

APPROACH TO THE PATIENT: Precocious Puberty

After verification of precocious development, serum LH and FSH levels should be measured to determine whether gonadotropins are increased in relation to chronologic age (gonadotropin-dependent) or whether sex steroid secretion is occurring independent of LH and FSH (gonadotropin-independent). In children with gonadotropin-dependent precocious puberty, CNS lesions should be excluded by history, neurologic examination, and MRI scan of the head. If organic causes are not found, one is left with the diagnosis of idiopathic central precocity. Patients with high testosterone but suppressed LH concentrations have gonadotropin-independent sexual precocity; in these patients, DHEA sulfate (DHEAS) and 17α -hydroxyprogesterone should be measured. High levels of testosterone and 17α -hydroxyprogesterone suggest the possibility of CAH due to 21α -hydroxylase or 11β -hydroxylase deficiency. If testosterone and DHEAS are elevated, adrenal tumors should be excluded by obtaining a computed tomography (CT) scan of the adrenal glands. Patients with elevated testosterone but without increased 17α -hydroxyprogesterone or DHEAS should undergo careful evaluation of the testis by palpation and ultrasound to

exclude a Leydig cell neoplasm. Activating mutations of the LH receptor should be considered in children with gonadotropin-independent precocious puberty in whom CAH, androgen abuse, and adrenal and testicular neoplasms have been excluded.

TREATMENT PRECOCIOUS PUBERTY

In patients with a known cause (e.g., a CNS lesion or a testicular tumor), therapy should be directed toward the underlying disorder. In patients with idiopathic CPP, long-acting GnRH analogues can be used to suppress gonadotropins and decrease testosterone, halt early pubertal development, delay accelerated bone maturation, prevent early epiphyseal closure, promote final height gain, and mitigate the psychosocial consequences of early pubertal development without causing osteoporosis. The treatment is most effective for increasing final adult height if it is initiated before age 6. Puberty resumes after discontinuation of the GnRH analogue. Counseling is an important aspect of the overall treatment strategy.

In children with gonadotropin-independent precocious puberty, inhibitors of steroidogenesis, such as ketoconazole, and AR antagonists have been used empirically. Long-term treatment with spironolactone (a weak androgen antagonist) and ketoconazole has been reported to normalize growth rate and bone maturation and to improve predicted height in small, nonrandomized trials in boys with familial male-limited precocious puberty. Aromatase inhibitors, such as testolactone and letrozole, have been used as an adjunct to antiandrogen and GnRH analogue therapy for children with familial male-limited precocious puberty, CAH, and McCune-Albright syndrome.

DELAYED PUBERTY

Puberty is delayed in boys if it has not ensued by age 14, an age that is 2–2.5 standard deviations above the mean for healthy children. Delayed puberty is more common in boys than in girls. There are four main categories of delayed puberty: (1) constitutional delay of growth and puberty (~60% of cases); (2) functional hypogonadotropic hypogonadism caused by systemic illness or malnutrition (~20% of cases); (3) hypogonadotropic hypogonadism caused by genetic or acquired defects in the hypothalamic-pituitary region (~10% of cases); and (4) hypergonadotropic hypogonadism secondary to primary gonadal failure (~15% of cases) (Table 411-1). Functional hypogonadotropic hypogonadism is more common in girls than in boys. Permanent causes of hypogonadotropic or hypergonadotropic hypogonadism are identified in >25% of boys with delayed puberty.

APPROACH TO THE PATIENT: Delayed Puberty

Any history of systemic illness, eating disorders, excessive exercise, social and psychological problems, and abnormal patterns of linear growth during childhood should be verified. Boys with pubertal delay may have accompanying emotional and physical immaturity relative to their peers, which can be a source of anxiety. Physical examination should focus on height; arm span; weight; visual fields; and secondary sex characteristics, including hair growth, testicular volume, phallic size, and scrotal reddening and thinning. Testicular size >2.5 cm generally indicates that the child has entered puberty.

The main diagnostic challenge is to distinguish those with constitutional delay, who will progress through puberty at a later age, from those with an underlying pathologic process. Constitutional delay should be suspected when there is a family history and when there are delayed bone age and short stature. Pituitary priming by pulsatile GnRH is required before LH and FSH are synthesized and secreted normally. Thus, blunted responses to exogenous GnRH can be seen in patients with constitutional delay, GnRH deficiency, or pituitary disorders (see “GnRH Stimulation Testing,” above). On

the other hand, low-normal basal gonadotropin levels or a normal response to exogenous GnRH is consistent with an early stage of puberty, which is often heralded by nocturnal GnRH secretion. Thus, constitutional delay is a diagnosis of exclusion that requires ongoing evaluation until the onset of puberty and the growth spurt.

TREATMENT DELAYED PUBERTY

If therapy is considered appropriate, it can begin with 25–50 mg testosterone enanthate or testosterone cypionate every 2 weeks, or by using a 2.5-mg testosterone patch or 25-mg testosterone gel. Because aromatization of testosterone to estrogen is obligatory for mediating androgen effects on epiphyseal fusion, concomitant treatment with aromatase inhibitors may allow attainment of greater final adult height. Testosterone treatment should be interrupted after 6 months to determine if endogenous LH and FSH secretion have ensued. Other causes of delayed puberty should be considered when there are associated clinical features or when boys do not enter puberty spontaneously after a year of observation or treatment.

Reassurance without hormonal treatment is appropriate for many individuals with presumed constitutional delay of puberty. However, the impact of delayed growth and pubertal progression on a child's social relationships and school performance should be weighed. Also, boys with constitutional delay of puberty are less likely to achieve their full genetic height potential and have reduced total-body bone mass as adults, mainly due to narrow limb bones and vertebrae as a result of impaired periosteal expansion during puberty. Administration of androgen therapy to boys with constitutional delay does not affect final height, and when administered with an aromatase inhibitor, it may improve final height.

DISORDERS OF THE MALE REPRODUCTIVE AXIS DURING ADULTHOOD

HYPOGONADOTROPIC HYPOGONADISM

Because LH and FSH are trophic hormones for the testes, impaired secretion of these pituitary gonadotropins results in secondary hypogonadism, which is characterized by low testosterone in the setting of low LH and FSH. Those with the most severe deficiency have complete absence of pubertal development, sexual infantilism, and, in some cases, hypospadias and undescended testes. Patients with partial gonadotropin deficiency have delayed or arrested sex development. The 24-h LH secretory profiles are heterogeneous in patients with hypogonadotropic hypogonadism, reflecting variable abnormalities of LH pulse frequency or amplitude. In severe cases, basal LH is low and there are no LH pulses. A smaller subset of patients has low-amplitude LH pulses or markedly reduced pulse frequency. Occasionally, only sleep-entrained LH pulses occur, reminiscent of the pattern seen in the early stages of puberty. Hypogonadotropic hypogonadism can be classified into congenital and acquired disorders. Congenital disorders most commonly involve GnRH deficiency, which leads to gonadotropin deficiency. Acquired disorders are much more common than congenital disorders and may result from a variety of sellar mass lesions or infiltrative diseases of the hypothalamus or pituitary.

Congenital Disorders Associated with Gonadotropin Deficiency Congenital hypogonadotropic hypogonadism is a heterogeneous group of disorders characterized by decreased gonadotropin secretion and testicular dysfunction either due to impaired function of the GnRH pulse generator or the gonadotrope. The disorders characterized by GnRH deficiency represent a family of oligogenic disorders whose phenotype spans a wide spectrum. Some individuals with GnRH deficiency may suffer from complete absence of pubertal development, while others may manifest varying degrees of gonadotropin deficiency and pubertal delay, and a subset that carries the same mutations as their affected family members may even have normal reproductive function. In approximately 10% of men with idiopathic hypogonadotropic hypogonadism,

reversal of gonadotropin deficiency may occur in adult life after sex steroid therapy. Also, a small fraction of men with idiopathic hypogonadotropic hypogonadism may present with androgen deficiency and infertility in adult life after having gone through apparently normal pubertal development. Nutritional, emotional, or metabolic stress may unmask gonadotropin deficiency and reproductive dysfunction (analogous to hypothalamic amenorrhea) in some patients who harbor mutations in the candidate genes but who previously had normal reproductive function. The clinical phenotype may include isolated anosmia or hyposmia. These striking variations in phenotypic presentation of GnRH deficiency have highlighted the important role of oligogenicity and gene-gene and gene-environment interactions in shaping the clinical phenotype.

Mutations in a number of genes involved in the development and migration of GnRH neurons or in the regulation of GnRH secretion have been linked to GnRH deficiency, although the genetic defect remains elusive in nearly two-thirds of cases. Familial hypogonadotropic hypogonadism can be transmitted as an X-linked (20%), autosomal recessive (30%), or autosomal dominant (50%) trait. Some individuals with idiopathic hypogonadotropic hypogonadism (IHH) have sporadic mutations in the same genes that cause inherited forms of the disorder. The genetic defects associated with GnRH deficiency can be conveniently classified as anosmic (Kallmann's syndrome) or normosmic (Table 411-2), although the occurrence of both anosmic and normosmic forms of GnRH deficiency in the same families suggests commonality of pathophysiologic mechanisms. *Kallmann's syndrome*, the anosmic form of GnRH deficiency, can result from mutations in one or more genes associated with olfactory bulb morphogenesis and the migration of GnRH neurons from their origin in the region of the olfactory placode, along the scaffold established by the olfactory nerves, through the cribriform plate into their final location into the preoptic region of the hypothalamus. Thus, mutations in *KAL1*, *FGF8*, *FGFR1*, *NELF*, *PROK2*, *PROK2R*, and *CHD7* have been described in patients with Kallmann's syndrome. An X-linked form of IHH is caused by mutations in the *KAL1* gene, which encodes anosmin, a protein that mediates the migration of neural progenitors of the olfactory bulb and GnRH-producing neurons. These individuals have GnRH deficiency and variable combinations of anosmia or hyposmia, renal defects, and neurologic abnormalities including mirror movements. Mutations in the *FGFR1* gene cause an autosomal dominant form of hypogonadotropic hypogonadism that clinically resembles Kallmann's syndrome; mutations in its putative ligand, *FGF8* gene product, have also been associated with IHH. Prokineticin 2 (*PROK2*) also encodes a protein involved in migration and development of olfactory and GnRH neurons. Recessive mutations in *PROK2* or in its receptor, *PROKR2*, have been associated with both anosmic and normosmic forms of hypogonadotropic hypogonadism.

Normosmic GnRH deficiency results from defects in pulsatile GnRH secretion, its regulation, or its action on the gonadotrope and has been associated with mutations in *GnRHR*, *GNRH1*, *KISS1R*, *TAC3*, *TACR3*, and *NROB1* (*DAX1*). Some mutations, such as those in *PROK2*, *PROKR2*, and *CHD7*, have been associated with both the anosmic and normosmic forms of IHH. *GnRHR* mutations, the most frequent identifiable cause of normosmic IHH, account for ~40% of autosomal recessive and 10% of sporadic cases of hypogonadotropic hypogonadism. These patients have decreased LH response to exogenous GnRH. Some receptor mutations alter GnRH binding affinity, allowing apparently normal responses to pharmacologic doses of exogenous GnRH, whereas other mutations may alter signal transduction downstream of hormone binding. Mutations of the *GnRH1* gene have also been reported in patients with hypogonadotropic hypogonadism, although they are rare. G protein-coupled receptor *KISS1R* (*GPR54*) and its cognate ligand, kisspeptin (*KISS1*), are important regulators of sexual maturation in primates. Recessive mutations in *GPR54* cause gonadotropin deficiency without anosmia. Patients retain responsiveness to exogenous GnRH, suggesting an abnormality in the neural pathways controlling GnRH release. The genes encoding neurokinin B (*TAC3*), which is involved in preferential activation of GnRH release in early development, and its receptor (*TAC3R*) have been implicated

TABLE 411-2 CAUSES OF CONGENITAL HYPOGONADOTROPIC HYPOGONADISM

Gene	Locus	Inheritance	Associated Features
A. Hypogonadotropic Hypogonadism due to GnRH Deficiency			
A1. GnRH Deficiency Associated with Hyposmia or Anosmia			
<i>KAL1</i>	Xp22	X-linked	Anosmia, renal agenesis, synkinesia, cleft lip/palate, oculomotor/visuospatial defects, gut malformations
<i>NELF</i>	9q34.3	AR	Anosmia, hypogonadotropic hypogonadism
<i>FGFR1</i>	8p11-p12	AD	Anosmia, cleft lip/palate, synkinesia, syndactyly
<i>PROK2</i>	3p21	AR	Anosmia/sleep dysregulation
<i>PROK2R</i>	20p12.3	AR	Variable
<i>CHD7</i>	8q12.1		Anosmia, other features of CHARGE syndrome
A2. GnRH Deficiency with Normal Sense of Smell			
<i>GNRHR</i>	4q21	AR	None
<i>GnRH1</i>	8p21	AR	None
<i>KISS1R</i>	19p13	AR	None
<i>TAC3</i>	12q13	AR	Microphallus, cryptorchidism, reversal of GnRH deficiency
<i>TAC3R</i>	4q25	AR	Microphallus, cryptorchidism, reversal of GnRH deficiency
<i>LEPR</i>	1p31	AR	Obesity
<i>LEP</i>	7q31	AR	Obesity
<i>FGF8</i>	10q24	AR	Skeletal abnormalities
B. Hypogonadotropic Hypogonadism not due to GnRH Deficiency			
<i>PC1</i>	5q15-21	AR	Obesity, diabetes mellitus, ACTH deficiency
<i>HESX1</i>	3p21	AR	Septo-optic dysplasia, CPHD
		AD	Isolated GH insufficiency
<i>LHX3</i>	9q34	AR	CPHD (ACTH spared), cervical spine rigidity
<i>PROP1</i>	5q35	AR	CPHD (ACTH usually spared)
<i>FSHβ</i>	11p13	AR	↑ LH
<i>LHβ</i>	19q13	AR	↑ FSH
<i>SF1</i>	9p33	AD/AR	Primary adrenal failure, XY sex reversal (NR5A1)

Abbreviations: ACTH, adrenocorticotropic hormone; AD, autosomal dominant; AR, autosomal recessive; CHARGE, eye coloboma, choanal atresia, growth and developmental retardation, genitourinary anomalies, ear anomalies; CPHD, combined pituitary hormone deficiency; *DAX1*, dosage-sensitive sex-reversal, adrenal hypoplasia congenita, X-chromosome; *FGFR1*, fibroblast growth factor receptor 1; FSH, follicle-stimulating hormone; *FSHβ*, follicle-stimulating hormone β-subunit; GH, growth hormone; GnRH, gonadotropin-releasing hormone; *GNRHR*, gonadotropin-releasing hormone receptor; *GPR54*, G protein-coupled receptor 54; *HESX1*, homeo box gene expressed in embryonic stem cells 1; *KAL1*, interval-1 gene; *LEP*, leptin; *LEPR*, leptin receptor; LH, luteinizing hormone; *LHβ*, luteinizing hormone β-subunit; *LHX3*, LIM homeobox gene 3; *NELF*, nasal embryonic LHRH factor; *PC1*, prohormone convertase 1; *PROK2*, prokineticin 2; *PROP1*, Prophet of Pit 1; *SF1*, steroidogenic factor 1; *TAC3*, tachykinin 3; *TAC3R*, tachykinin 3 receptor.

in some families with normosmic IHH. Mutations in more than one gene (digenicity or oligogenicity) may contribute to clinical heterogeneity in IHH patients. X-linked hypogonadotropic hypogonadism also occurs in *adrenal hypoplasia congenita*, a disorder caused by mutations in the *DAX1* gene, which encodes a nuclear receptor in the adrenal gland and reproductive axis. Adrenal hypoplasia congenita is characterized by absent development of the adult zone of the adrenal cortex, leading to neonatal adrenal insufficiency. Puberty usually does not occur or is arrested, reflecting variable degrees of gonadotropin deficiency. Although sexual differentiation is normal, **most** patients have testicular dysgenesis and impaired spermatogenesis despite gonadotropin replacement. Less commonly, adrenal hypoplasia congenita, sex reversal, and hypogonadotropic hypogonadism can be caused by mutations of steroidogenic factor 1 (SF1). Rarely, recessive mutations in the *LHβ* or *FSHβ* gene have been described in patients with selective deficiencies of these gonadotropins. In approximately 10% of men with IHH, reversal of gonadotropin deficiency may occur in adult life.

Also, a small fraction of men with IHH may present with androgen deficiency and infertility in adult life after having gone through apparently normal pubertal development.

A number of homeodomain transcription factors are involved in the development and differentiation of the specialized hormone-producing cells within the pituitary gland (Table 411-2). Patients with mutations of *PROPI* have combined pituitary hormone deficiency that includes GH, prolactin (PRL), thyroid-stimulating hormone (TSH), LH, and FSH, but not ACTH. *LHX3* mutations cause combined pituitary hormone deficiency in association with cervical spine rigidity. *HESX1* mutations cause septo-optic dysplasia and combined pituitary hormone deficiency.

Prader-Willi syndrome is characterized by obesity, hypotonic musculature, mental retardation, hypogonadism, short stature, and small hands and feet. Prader-Willi syndrome is a genomic imprinting disorder caused by deletions of the proximal portion of the paternally derived chromosome 15q11-15q13 region, which contains a bipartite imprinting center, uniparental disomy of the maternal alleles, or mutations of the genes/loci involved in imprinting (Chap. 83e). *Laurence-Moon syndrome* is an autosomal recessive disorder characterized by obesity, hypogonadism, mental retardation, polydactyly, and retinitis pigmentosa. Recessive mutations of leptin, or its receptor, cause severe obesity and pubertal arrest, apparently because of hypothalamic GnRH deficiency (Chap. 415e).

Acquired Hypogonadotropic Disorders • SEVERE ILLNESS, STRESS, MALNUTRITION, AND EXERCISE These factors may cause reversible gonadotropin deficiency. Although gonadotropin deficiency and reproductive dysfunction are well documented in these conditions in women, men exhibit similar but less pronounced responses. Unlike women, most male runners and other endurance athletes have normal gonadotropin and sex steroid levels, despite low body fat and frequent intensive exercise. Testosterone levels fall at the onset of illness and recover during recuperation. The magnitude of gonadotropin suppression generally correlates with the severity of illness. Although hypogonadotropic hypogonadism is the most common cause of androgen deficiency in patients with acute illness, some have elevated levels of LH and FSH, which suggest primary gonadal dysfunction. The pathophysiology of reproductive dysfunction during acute illness is unknown but likely involves a combination of cytokine and/or glucocorticoid effects. There is a high frequency of low testosterone levels in patients with chronic illnesses such as HIV infection, end-stage renal disease, chronic obstructive lung disease, and many types of cancer and in patients receiving glucocorticoids. About 20% of HIV-infected men with low testosterone levels have elevated LH and FSH levels; these patients presumably have primary testicular dysfunction. The remaining 80% have either normal or low LH and FSH levels; these men have a central hypothalamic-pituitary defect or a dual defect involving both the testis and the hypothalamic-pituitary centers. Muscle wasting is common in chronic diseases associated with hypogonadism, which also leads to debility, poor quality of life, and adverse outcome of disease. There is great interest in exploring strategies that can reverse androgen deficiency or attenuate the sarcopenia associated with chronic illness.

Men using opioids for relief of cancer or noncancerous pain or because of addiction often have suppressed testosterone and LH levels and high prevalence of sexual dysfunction and osteoporosis; the degree of suppression is dose-related and particularly severe with long-acting opioids such as methadone. Opioids suppress GnRH secretion and alter the sensitivity to feedback inhibition by gonadal steroids. Men who are heavy users of marijuana have decreased testosterone secretion and sperm production. The mechanism of marijuana-induced hypogonadism is decreased GnRH secretion. Gynecomastia observed in marijuana users can also be caused by plant estrogens in crude preparations. Androgen deprivation therapy in men with prostate cancer has been associated with increased risk of bone fractures, diabetes mellitus, cardiovascular events, fatigue, sexual dysfunction, and poor quality of life.

OBESITY In men with mild to moderate obesity, SHBG levels decrease in proportion to the degree of obesity, resulting in lower

total testosterone levels. However, free testosterone levels usually remain within the normal range. The decrease in SHBG levels is caused by increased circulating insulin, which inhibits SHBG production. Estradiol levels are higher in obese men compared to healthy, nonobese controls, because of aromatization of testosterone to estradiol in adipose tissue. Weight loss is associated with reversal of these abnormalities including an increase in total and free testosterone levels and a decrease in estradiol levels. A subset of obese men with moderate to severe obesity may have a defect in the hypothalamic-pituitary axis as suggested by low free testosterone in the absence of elevated gonadotropins. Weight gain in adult men can accelerate the rate of age-related decline in testosterone levels.

HYPERPROLACTINEMIA (See also Chap. 403) Elevated PRL levels are associated with hypogonadotropic hypogonadism. PRL inhibits hypothalamic GnRH secretion either directly or through modulation of tuberoinfundibular dopaminergic pathways. A PRL-secreting tumor may also destroy the surrounding gonadotropes by invasion or compression of the pituitary stalk. Treatment with dopamine agonists reverses gonadotropin deficiency, although there may be a delay relative to PRL suppression.

SELLAR MASS LESIONS Neoplastic and nonneoplastic lesions in the hypothalamus or pituitary can directly or indirectly affect gonadotrope function. In adults, pituitary adenomas constitute the largest category of space-occupying lesions affecting gonadotropin and other pituitary hormone production. Pituitary adenomas that extend into the suprasellar region can impair GnRH secretion and mildly increase PRL secretion (usually <50 µg/L) because of impaired tonic inhibition by dopaminergic pathways. These tumors should be distinguished from prolactinomas, which typically secrete higher PRL levels. The presence of diabetes insipidus suggests the possibility of a craniopharyngioma, infiltrative disorder, or other hypothalamic lesions (Chap. 404).

HEMOCHROMATOSIS (See also Chap. 428) Both the pituitary and testis can be affected by excessive iron deposition. However, the pituitary defect is the predominant lesion in most patients with hemochromatosis and hypogonadism. The diagnosis of hemochromatosis is suggested by the association of characteristic skin discoloration, hepatic enlargement or dysfunction, diabetes mellitus, arthritis, cardiac conduction defects, and hypogonadism.

PRIMARY TESTICULAR CAUSES OF HYPOGONADISM

Common causes of primary testicular dysfunction include Klinefelter's syndrome, uncorrected cryptorchidism, cancer chemotherapy, radiation to the testes, trauma, torsion, infectious orchitis, HIV infection, anorchia syndrome, and myotonic dystrophy. Primary testicular disorders may be associated with impaired spermatogenesis, decreased androgen production, or both. See Chap. 410 for disorders of testis development, androgen synthesis, and androgen action.

Klinefelter's Syndrome (See also Chap. 410) Klinefelter's syndrome is the most common chromosomal disorder associated with testicular dysfunction and male infertility. It occurs in about 1 in 600 live-born males. Azoospermia is the rule in men with Klinefelter's syndrome who have the 47,XXY karyotype; however, men with mosaicism may have germ cells, especially at a younger age. The clinical phenotype of Klinefelter's syndrome can be heterogeneous possibly because of mosaicism, polymorphisms in AR gene, variable testosterone levels, or other genetic factors. Testicular histology shows hyalinization of seminiferous tubules and absence of spermatogenesis. Although their function is impaired, the number of Leydig cells appears to increase. Testosterone is decreased and estradiol is increased, leading to clinical features of undervirilization and gynecomastia. Men with Klinefelter's syndrome are at increased risk of systemic lupus erythematosus, Sjögren's syndrome, breast cancer, diabetes mellitus, osteoporosis, non-Hodgkin's lymphoma, and lung cancer, and reduced risk of prostate cancer. Periodic mammography for breast cancer surveillance is recommended for men with Klinefelter's syndrome. Fertility has been achieved by intracytoplasmic injection of sperm retrieved surgically from testicular biopsies of men with Klinefelter's syndrome, including

some men with the nonmosaic form of Klinefelter's syndrome. The karyotypes 48,XXXXY and 49,XXXXXY are associated with a more severe phenotype, increased risk of congenital malformations, and lower intelligence than 47,XXY individuals.

Cryptorchidism Cryptorchidism occurs when there is incomplete descent of the testis from the abdominal cavity into the scrotum. About 3% of full-term and 30% of premature male infants have at least one undescended testis at birth, but descent is usually complete by the first few weeks of life. The incidence of cryptorchidism is <1% by 9 months of age. Androgens regulate predominantly the inguinoscrotal descent of the testes through degeneration of the craniosuspensory ligament and a shortening of the gubernaculum, respectively. Mutations in *INSL3* and leucine-rich repeat family of G protein-coupled receptor 8 (*LGR8*), which regulate the transabdominal portion of testicular descent, have been found in some patients with cryptorchidism.

Cryptorchidism is associated with increased risk of malignancy, infertility, inguinal hernia, and torsion. Unilateral cryptorchidism, even when corrected before puberty, is associated with decreased sperm count, possibly reflecting unrecognized damage to the fully descended testis or other genetic factors. Epidemiologic, clinical, and molecular evidence supports the idea that cryptorchidism, hypospadias, impaired spermatogenesis, and testicular cancer may be causally related to common genetic and environment perturbations and are components of the testicular dysgenesis syndrome.

Acquired Testicular Defects *Viral orchitis* may be caused by the mumps virus, echovirus, lymphocytic choriomeningitis virus, and group B arboviruses. Orchitis occurs in as many as one-fourth of adult men with mumps; the orchitis is unilateral in about two-thirds and bilateral in the remainder. Orchitis usually develops a few days after the onset of parotitis but may precede it. The testis may return to normal size and function or undergo atrophy. Semen analysis returns to normal for three-fourths of men with unilateral involvement but for only one-third of men with bilateral orchitis. *Trauma*, including testicular torsion, can also cause secondary atrophy of the testes. The exposed position of the testes in the scrotum renders them susceptible to both thermal and physical trauma, particularly in men with hazardous occupations.

The testes are sensitive to *radiation damage*. Doses >200 mGy (20 rad) are associated with increased FSH and LH levels and damage to the spermatogonia. After ~800 mGy (80 rad), oligospermia or azoospermia develops, and higher doses may obliterate the germinal epithelium. Permanent androgen deficiency in adult men is uncommon after therapeutic radiation; however, most boys given direct testicular radiation therapy for acute lymphoblastic leukemia have permanently low testosterone levels. Sperm banking should be considered before patients undergo radiation treatment or chemotherapy.

Drugs interfere with testicular function by several mechanisms, including inhibition of testosterone synthesis (e.g., ketoconazole), blockade of androgen action (e.g., spironolactone), increased estrogen (e.g., marijuana), or direct inhibition of spermatogenesis (e.g., chemotherapy).

Combination chemotherapy for acute leukemia, Hodgkin's disease, and testicular and other cancers may impair Leydig cell function and cause infertility. The degree of gonadal dysfunction depends on the type of chemotherapeutic agent and the dose and duration of therapy. Because of high response rates and the young age of these men, infertility and androgen deficiency have emerged as important long-term complications of cancer chemotherapy. Cyclophosphamide and combination regimens containing procarbazine are particularly toxic to germ cells. Thus, 90% of men with Hodgkin's lymphoma receiving MOPP (mechlorethamine, vincristine, procarbazine, prednisone) therapy develop azoospermia or extreme oligozoospermia; newer regimens that do not include procarbazine, such as ABVD (doxorubicin, bleomycin, vinblastine, dacarbazine), are less toxic to germ cells.

Alcohol, when consumed in excess for prolonged periods, decreases testosterone, independent of liver disease or malnutrition. Elevated estradiol and decreased testosterone levels may occur in men taking digitalis.

The occupational and recreational history should be carefully evaluated in all men with infertility because of the toxic effects of many *chemical agents* on spermatogenesis. Known environmental hazards include pesticides (e.g., vinclozolin, dicofol, atrazine), sewage contaminants (e.g., ethinyl estradiol in birth control pills, surfactants such as octylphenol, nonylphenol), plasticizers (e.g., phthalates), flame retardants (e.g., polychlorinated biphenyls, polybrominated diphenol ethers), industrial pollutants (e.g., heavy metals cadmium and lead, dioxins, polycyclic aromatic hydrocarbons), microwaves, and ultrasound. In some populations, sperm density is said to have declined by as much as 40% in the past 50 years. Environmental estrogens or antiandrogens may be partly responsible.

Testicular failure also occurs as a part of *polyglandular autoimmune insufficiency* (Chap. 408). Sperm antibodies can cause isolated male infertility. In some instances, these antibodies are secondary phenomena resulting from duct obstruction or vasectomy. Granulomatous diseases can affect the testes, and testicular atrophy occurs in 10–20% of men with lepromatous leprosy because of direct tissue invasion by the mycobacteria. The tubules are involved initially, followed by endarteritis and destruction of Leydig cells.

Systemic disease can cause primary testis dysfunction in addition to suppressing gonadotropin production. In cirrhosis, a combined testicular and pituitary abnormality leads to decreased testosterone production independent of the direct toxic effects of ethanol. Impaired hepatic extraction of adrenal androstenedione leads to extraglandular conversion to estrone and estradiol, which partially suppresses LH. Testicular atrophy and gynecomastia are present in approximately one-half of men with cirrhosis. In chronic renal failure, androgen synthesis and sperm production decrease despite elevated gonadotropins. The elevated LH level is due to reduced clearance, but it does not restore normal testosterone production. About one-fourth of men with renal failure have hyperprolactinemia. Improvement in testosterone production with hemodialysis is incomplete, but successful renal transplantation may return testicular function to normal. Testicular atrophy is present in one-third of men with sickle cell anemia. The defect may be at either the testicular or the hypothalamic-pituitary level. Sperm density can decrease temporarily after acute febrile illness in the absence of a change in testosterone production. Infertility in men with celiac disease is associated with a hormonal pattern typical of androgen resistance, namely elevated testosterone and LH levels.

Neurologic diseases associated with altered testicular function include myotonic dystrophy, spinobulbar muscular atrophy, and paraplegia. In myotonic dystrophy, small testes may be associated with impairment of both spermatogenesis and Leydig cell function. Spinobulbar muscular atrophy is caused by an expansion of the glutamine repeat sequences in the amino-terminal region of the AR; this expansion impairs function of the AR, but it is unclear how the alteration is related to the neurologic manifestations. Men with spinobulbar muscular atrophy often have undervirilization and infertility as a late manifestation. Spinal cord lesions that cause paraplegia can lead to a temporary decrease in testosterone levels and may cause persistent defects in spermatogenesis; some patients retain the capacity for penile erection and ejaculation.

ANDROGEN INSENSITIVITY SYNDROMES

Mutations in the AR cause resistance to the action of testosterone and DHT. These X-linked mutations are associated with variable degrees of defective male phenotypic development and undervirilization (Chap. 410). Although not technically hormone-insensitivity syndromes, two genetic disorders impair testosterone conversion to active sex steroids. Mutations in the *SRD5A2* gene, which encodes 5 α -reductase type 2, prevent the conversion of testosterone to DHT, which is necessary for the normal development of the male external genitalia. Mutations in the *CYP19* gene, which encodes aromatase, prevent testosterone conversion to estradiol. Males with *CYP19* mutations have delayed epiphyseal fusion, tall stature, eunuchoid proportions, and osteoporosis, consistent with evidence from an estrogen receptor-deficient individual that these testosterone actions are mediated indirectly via estrogen.

GYNecomastia

Gynecomastia refers to enlargement of the male breast. It is caused by excess estrogen action and is usually the result of an increased estrogen-to-androgen ratio. True gynecomastia is associated with glandular breast tissue that is >4 cm in diameter and often tender. Glandular tissue enlargement should be distinguished from excess adipose tissue: glandular tissue is firmer and contains fibrous-like cords. Gynecomastia occurs as a normal physiologic phenomenon in the newborn (due to transplacental transfer of maternal and placental estrogens), during puberty (high estrogen-to-androgen ratio in early stages of puberty), and with aging (increased fat tissue and increased aromatase activity), but it can also result from pathologic conditions associated with androgen deficiency or estrogen excess. The prevalence of gynecomastia increases with age and body mass index (BMI), likely because of increased aromatase activity in adipose tissue. Medications that alter androgen metabolism or action may also cause gynecomastia. The relative risk of breast cancer is increased in men with gynecomastia, although the absolute risk is relatively small.

PATHOLOGIC GYNecomastia

Any cause of *androgen deficiency* can lead to gynecomastia, reflecting an increased estrogen-to-androgen ratio, because estrogen synthesis still occurs by aromatization of residual adrenal and gonadal androgens. Gynecomastia is a characteristic feature of Klinefelter's syndrome (Chap. 410). *Androgen insensitivity* disorders also cause gynecomastia. *Excess estrogen production* may be caused by tumors, including Sertoli cell tumors in isolation or in association with Peutz-Jeghers syndrome or Carney complex. Tumors that produce hCG, including some testicular tumors, stimulate Leydig cell estrogen synthesis. *Increased conversion of androgens to estrogens* can be a result of increased availability of substrate (androstenedione) for extraglandular estrogen formation (CAH, hyperthyroidism, and most feminizing adrenal tumors) or of diminished catabolism of androstenedione (liver disease) so that estrogen precursors are shunted to aromatase in peripheral sites. Obesity is associated with increased aromatization of androgen precursors to estrogens. Extraglandular aromatase activity can also be increased in tumors of the liver or adrenal gland or rarely as an inherited disorder. Several families with *increased peripheral aromatase activity* inherited as an autosomal dominant or as an X-linked disorder have been described. In some families with this disorder, an inversion in chromosome 15q21.2-3 causes the *CYP19* gene to be activated by the regulatory elements of contiguous genes, resulting in excessive estrogen production in the fat and other extragonadal tissues. *Drugs* can cause gynecomastia by acting directly as estrogenic substances (e.g., oral contraceptives, phytoestrogens, digitalis) or by inhibiting androgen synthesis (e.g., ketoconazole) or action (e.g., spironolactone).

Because up to two-thirds of pubertal boys and half of hospitalized men have palpable glandular tissue that is benign, detailed investigation or intervention is not indicated in all men presenting with gynecomastia (Fig. 411-5). In addition to the extent of gynecomastia, recent onset, rapid growth, tender tissue, and occurrence in a lean subject should prompt more extensive evaluation. This should include a careful drug history, measurement and examination of the testes, assessment of virilization, evaluation of liver function, and hormonal measurements including testosterone, estradiol, and androstenedione, LH, and hCG. A karyotype should be obtained in men with very small testes to exclude Klinefelter's syndrome. Despite extensive evaluation, the etiology is established in fewer than one-half of patients.

TREATMENT GYNecomastia

When the primary cause can be identified and corrected, breast enlargement usually subsides over several months. However, if gynecomastia is of long duration, surgery is the most effective therapy. Indications for surgery include severe psychological and/or cosmetic problems, continued growth or tenderness, or suspected malignancy. In patients who have painful gynecomastia and in

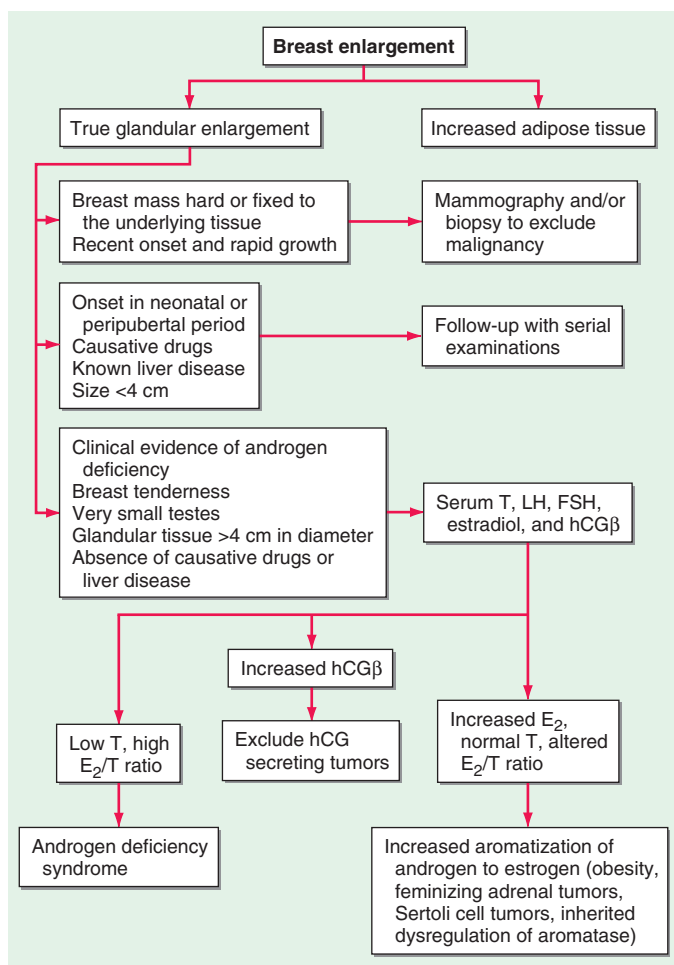


FIGURE 411-5 Evaluation of gynecomastia. E₂, 17β-estradiol; hCGβ, human chorionic gonadotropin β; T, testosterone.

whom surgery cannot be performed, treatment with antiestrogens such as tamoxifen (20 mg/d) can reduce pain and breast tissue size in over half the patients. Estrogen receptor antagonists, tamoxifen and raloxifene, have been reported in small trials to reduce breast size in men with pubertal gynecomastia, although complete regression of breast enlargement is unusual with the use of estrogen receptor antagonists. Aromatase inhibitors can be effective in the early proliferative phase of the disorder. However, in a randomized trial in men with established gynecomastia, anastrozole proved no more effective than placebo in reducing breast size. Tamoxifen is effective in the prevention and treatment of breast enlargement and breast pain in men with prostate cancer who are receiving antiandrogen therapy.

AGING-RELATED CHANGES IN MALE REPRODUCTIVE FUNCTION

A number of cross-sectional and longitudinal studies (e.g., The Baltimore Longitudinal Study of Aging, the Framingham Heart Study, the Massachusetts Male Aging Study, and the European Male Aging Study) have established that testosterone concentrations decrease with advancing age. This age-related decline starts in the third decade of life and progresses slowly; the rate of decline in testosterone concentrations is greater in obese men, men with chronic illness, and those taking medications than in healthy older men. Because SHBG concentrations are higher in older men than in younger men, free or bioavailable testosterone concentrations decline with aging to a greater extent than total testosterone concentrations. The age-related decline in testosterone is due to defects at all levels of the hypothalamic-pituitary-testicular axis: pulsatile GnRH secretion is attenuated, LH response to GnRH is reduced, and testicular response to LH is impaired. However, the gradual rise of LH with aging suggests that testis dysfunction is

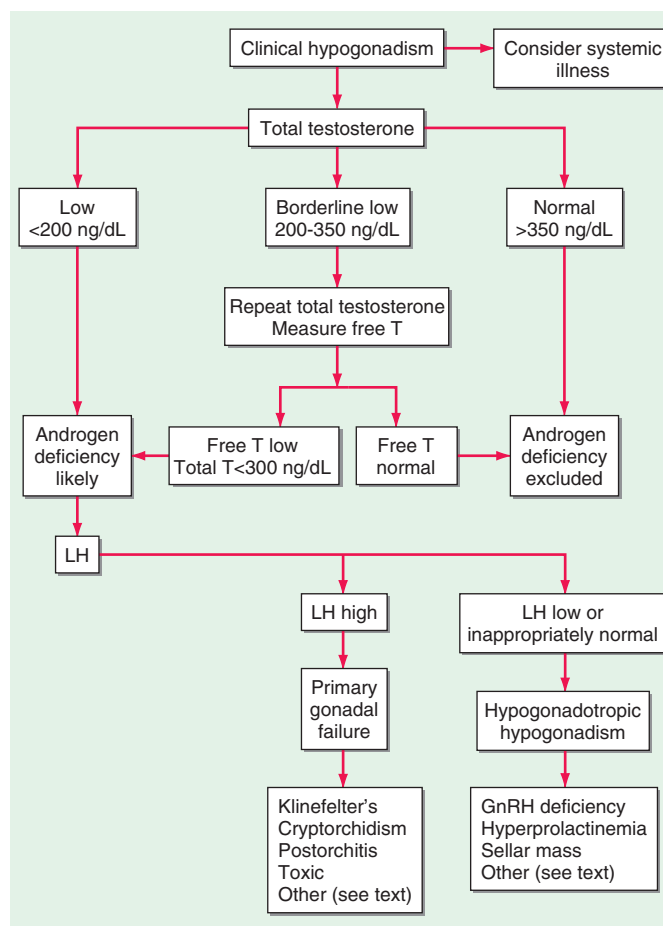


FIGURE 411-6 Evaluation of hypogonadism. GnRH, gonadotropin-releasing hormone; LH, luteinizing hormone; T, testosterone.

the main cause of declining androgen levels. The term *andropause* has been used to denote age-related decline in testosterone concentrations; this term is a misnomer because there is no discrete time when testosterone concentrations decline abruptly. The approach to evaluating hypogonadism is summarized in Fig. 411-6.

In epidemiologic surveys, low total and bioavailable testosterone concentrations have been associated with decreased appendicular skeletal muscle mass and strength, decreased self-reported physical function, higher visceral fat mass, insulin resistance, and increased risk of coronary artery disease and mortality, although the associations are weak. An analysis of signs and symptoms in older men in the European Male Aging Study revealed a syndromic association of sexual symptoms with total testosterone levels below 320 ng/dL and free testosterone levels below 64 pg/mL in community-dwelling older men. In systematic reviews of randomized controlled trials, testosterone therapy of healthy older men with low or low-normal testosterone levels was associated with greater increments in lean body mass, grip strength, and self-reported physical function compared with placebo. Testosterone therapy also induced greater improvement in vertebral but not femoral bone mineral density. Testosterone therapy of older men with sexual dysfunction and unequivocally low testosterone levels improves libido, but testosterone effects on erectile function and response to selective phosphodiesterase inhibitors have been inconsistent. Testosterone therapy has not been shown to improve depression scores, fracture risk, cognitive function, response to phosphodiesterase inhibitors, or clinical outcomes in older men. Furthermore, neither the long-term risks nor clinical benefits of testosterone therapy in older men have been demonstrated in adequately powered trials. Although there is no evidence that testosterone causes prostate cancer, there is concern that testosterone therapy might cause subclinical prostate cancers to grow. Testosterone therapy is associated with increased risk of detection of prostate events (Fig. 411-7).

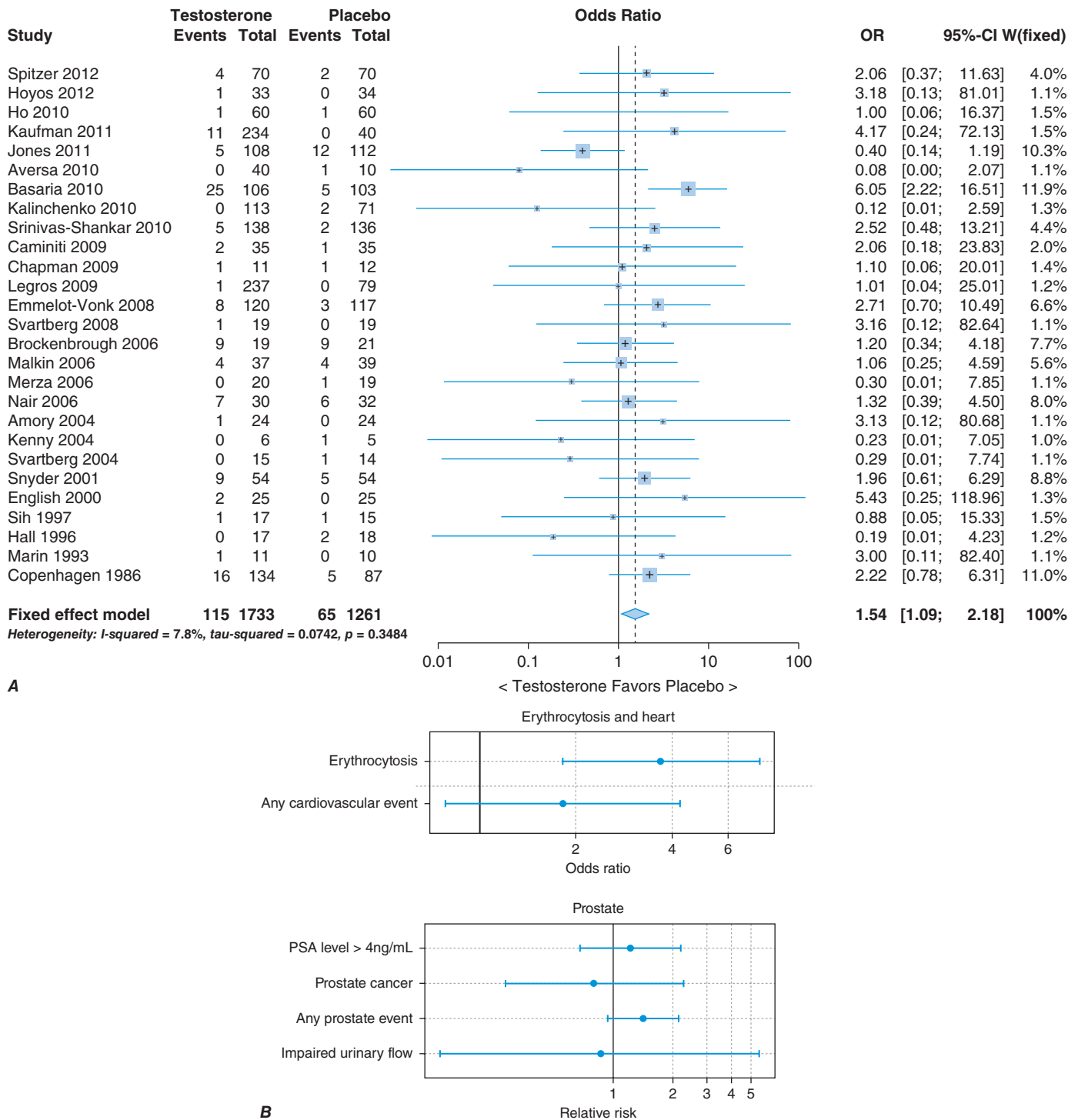


FIGURE 411-7 Meta-analyses of cardiovascular and prostate adverse events associated with testosterone therapy. **A.** A meta-analysis of cardiovascular-related events in randomized testosterone trials of 12 weeks or longer in duration. Randomization to testosterone was associated with a significantly increased risk of cardiovascular-related event (odds ratio [OR] 1.54). (Modified with permission from L Xu et al: Testosterone therapy and cardiovascular events among men: a systematic review and meta-analysis of placebo-controlled randomized trials *BMC Med* 11:108, 2013.) **B.** The relative risk of prostate events and the associated 95% confidence intervals (CIs) in a meta-analysis of randomized testosterone trials. PSA, prostate-specific antigen. (Data were derived from a meta-analysis by MM Fernández-Balsells et al: *J Clin Endocrinol Metab* 95:2560, 2010, and the figure was reproduced with permission from M Spitzer et al: *Nat Rev Endocrinol* 9:414, 2013.)

One randomized testosterone trial in older men with mobility limitation and high burden of chronic conditions, such as diabetes, heart disease, hypertension, and hyperlipidemia, reported a greater number of cardiovascular events in men randomized to the testosterone arm of the study than in those randomized to the placebo arm. Since then, two large retrospective analyses of patient databases have reported higher frequency of cardiovascular events, including myocardial infarction, in older men with preexisting heart disease (Fig. 411-7).

Population screening of all older men for low testosterone levels is not recommended, and testing should be restricted to men who have symptoms or physical features attributable to androgen deficiency. Testosterone therapy is not recommended for all older men with low testosterone levels. In older men with significant symptoms of androgen deficiency who have testosterone levels below 200 ng/dL, testosterone therapy may be considered on an individualized basis and should be instituted after careful

discussion of the risks and benefits (see “Testosterone Replacement,” below).

Testicular morphology, semen production, and fertility are maintained up to a very old age in men. Although concern has been expressed about age-related increases in germ cell mutations and impairment of DNA repair mechanisms, there is no clear evidence that the frequency of chromosomal aneuploidy is increased in the sperm of older men. However, the incidence of autosomal dominant diseases, such as achondroplasia, polyposis coli, Marfan’s syndrome, and Apert’s syndrome, increases in the offspring of men who are advanced in age, consistent with transmission of sporadic missense mutations. Advanced paternal age may be associated with increased rates of de novo mutations, which may contribute to an increased risk of neurodevelopmental diseases such as schizophrenia and autism. The somatic mutations in male germ cells that enhance the proliferation of germ cells could lead to within-testis expansion of mutant clonal lines, thus favoring the propagation of germ cells carrying these pathogenic mutations and increasing the risk of mutations in the offspring of older fathers (the “selfish spermatogonial selection” hypothesis).

APPROACH TO THE PATIENT: Androgen Deficiency

Hypogonadism is often characterized by decreased sex drive, reduced frequency of sexual activity, inability to maintain erections, reduced beard growth, loss of muscle mass, decreased testicular size, and gynecomastia. Erectile dysfunction and androgen deficiency are two distinct clinical disorders that can coexist in middle-aged and older men. Less than 10% of patients with erectile dysfunction have testosterone deficiency. Thus, it is useful to evaluate men presenting with erectile dysfunction for androgen deficiency. Except when extreme, these clinical features of androgen deficiency may be difficult to distinguish from changes that occur with normal aging. Moreover, androgen deficiency may develop gradually. Several epidemiologic studies, such as the Framingham Heart Study, the Massachusetts Male Aging Study, the Baltimore Longitudinal Study of Aging, and the Study of Osteoporotic Fractures in Men, have reported a high prevalence of low testosterone levels in middle-aged and older men. The age-related decline in testosterone should be distinguished from classical hypogonadism due to diseases of the testes, the pituitary, and the hypothalamus.

When symptoms or clinical features suggest possible androgen deficiency, the laboratory evaluation is initiated by the measurement of total testosterone, preferably in the morning using a reliable assay, such as LC-MS/MS that has been calibrated to an international testosterone standard (Fig. 411-6). A consistently low total testosterone level <300 ng/dL measured by a reliable assay, in association with symptoms, is evidence of testosterone deficiency. An early-morning testosterone level >400 ng/dL makes the diagnosis of androgen deficiency unlikely. In men with testosterone levels between 200 and 400 ng/dL, the total testosterone level should be repeated and a free testosterone level should be measured. In older men and in patients with other clinical states that are associated with alterations in SHBG levels, a direct measurement of free testosterone level by equilibrium dialysis can be useful in unmasking testosterone deficiency.

When androgen deficiency has been confirmed by the consistently low testosterone concentrations, LH should be measured to classify the patient as having primary (high LH) or secondary (low or inappropriately normal LH) hypogonadism. An elevated LH level indicates that the defect is at the testicular level. Common causes of primary testicular failure include Klinefelter’s syndrome, HIV infection, uncorrected cryptorchidism, cancer chemotherapeutic agents, radiation, surgical orchiectomy, or prior infectious orchitis. Unless causes of primary testicular failure are known, a karyotype should be performed in men with low testosterone and elevated LH to exclude Klinefelter’s syndrome. Men who have low testosterone levels but “inappropriately normal” or low LH

levels have secondary hypogonadism; their defect resides at the hypothalamic-pituitary level. Common causes of acquired secondary hypogonadism include space-occupying lesions of the sella, hyperprolactinemia, chronic illness, hemochromatosis, excessive exercise, and the use of anabolic-androgenic steroids, opiates, marijuana, glucocorticoids, and alcohol. Measurement of PRL and MRI scan of the hypothalamic-pituitary region can help exclude the presence of a space-occupying lesion. Patients in whom known causes of hypogonadotropic hypogonadism have been excluded are classified as having IHH. It is not unusual for congenital causes of hypogonadotropic hypogonadism, such as Kallmann’s syndrome, to be diagnosed in young adults.

TREATMENT ANDROGEN DEFICIENCY

GONADOTROPINS

Gonadotropin therapy is used to establish or restore fertility in patients with gonadotropin deficiency of any cause. Several gonadotropin preparations are available. Human menopausal gonadotropin (hMG; purified from the urine of postmenopausal women) contains 75 IU FSH and 75 IU LH per vial. hCG (purified from the urine of pregnant women) has little FSH activity and resembles LH in its ability to stimulate testosterone production by Leydig cells. Recombinant LH is now available. Because of the expense of hMG, treatment is usually begun with hCG alone, and hMG is added later to promote the FSH-dependent stages of spermatid development. Recombinant human FSH (hFSH) is now available and is indistinguishable from purified urinary hFSH in its biologic activity and pharmacokinetics in vitro and in vivo, although the mature β subunit of recombinant hFSH has seven fewer amino acids. Recombinant hFSH is available in ampoules containing 75 IU (~7.5 μ g FSH), which accounts for >99% of protein content. Once spermatogenesis is restored using combined FSH and LH therapy, hCG alone is often sufficient to maintain spermatogenesis.

Although a variety of treatment regimens are used, 1000–2000 IU of hCG or recombinant human LH (rhLH) administered intramuscularly three times weekly is a reasonable starting dose. Testosterone levels should be measured 6–8 weeks later and 48–72 h after the hCG or rhLH injection; the hCG/rhLH dose should be adjusted to achieve testosterone levels in the mid-normal range. Sperm counts should be monitored on a monthly basis. It may take several months for spermatogenesis to be restored; therefore, it is important to forewarn patients about the potential length and expense of the treatment and to provide conservative estimates of success rates. If testosterone levels are in the mid-normal range but the sperm concentrations are low after 6 months of therapy with hCG alone, FSH should be added. This can be done by using hMG, highly purified urinary hFSH, or recombinant hFSH. The selection of FSH dose is empirical. A common practice is to start with the addition of 75 IU FSH three times each week in conjunction with the hCG/rhLH injections. If sperm densities are still low after 3 months of combined treatment, the FSH dose should be increased to 150 IU. Occasionally, it may take \geq 18–24 months for spermatogenesis to be restored.

The two best predictors of success using gonadotropin therapy in hypogonadotropic men are testicular volume at presentation and time of onset. In general, men with testicular volumes >8 mL have better response rates than those who have testicular volumes >4 mL. Patients who became hypogonadotropic after puberty experience higher success rates than those who have never undergone pubertal changes. Spermatogenesis can usually be reinitiated by hCG alone, with high rates of success for men with postpubertal onset of hypogonadotropism. The presence of a primary testicular abnormality, such as cryptorchidism, will attenuate testicular response to gonadotropin therapy. Prior androgen therapy does not preclude subsequent response to gonadotropin therapy, although some studies suggest that it may attenuate response to subsequent gonadotropin therapy.

2370 TESTOSTERONE REPLACEMENT

Androgen therapy is indicated to restore testosterone levels to normal to correct features of androgen deficiency. Testosterone replacement improves libido and overall sexual activity; increases energy, lean muscle mass, and bone density; and decreases fat mass. The benefits of testosterone replacement therapy have only been proven in men who have documented androgen deficiency, as demonstrated by testosterone levels that are well below the lower limit of normal.

Testosterone is available in a variety of formulations with distinct pharmacokinetics (Table 411-3). Testosterone serves as a prohormone and is converted to 17β-estradiol by aromatase and to 5α-dihydrotestosterone by steroid 5α-reductase. Therefore, when evaluating testosterone formulations, it is important to consider whether the formulation being used can achieve physiologic estradiol and DHT concentrations, in addition to normal testosterone concentrations. Although testosterone concentrations at the lower end of the normal male range can restore sexual function, it is not

TABLE 411-3 CLINICAL PHARMACOLOGY OF SOME TESTOSTERONE FORMULATIONS

Formulation	Regimen	Pharmacokinetic Profile	DHT and E ₂	Advantages	Disadvantages
Testosterone enanthate or cypionate	150–200 mg IM q2wk or 75–100 mg/wk	After a single IM injection, serum T levels rise into the supra-physiologic range, then decline gradually into the hypogonadal range by the end of the dosing interval	DHT and E ₂ levels rise in proportion to the increase in T levels; T:DHT and T:E ₂ ratios do not change	Corrects symptoms of androgen deficiency; relatively inexpensive, if self-administered; flexibility of dosing	Requires IM injection; peaks and valleys in serum T levels
Topical testosterone gels and axillary testosterone solution	Available in sachets, tubes, and pumps	When used in appropriate doses, these topical formulations restore serum T and E ₂ levels to the physiologic male range	Serum DHT levels are higher and T:DHT ratios are lower in hypogonadal men treated with the transdermal gels than in healthy eugonadal men	Corrects symptoms of androgen deficiency, provides flexibility of dosing, ease of application, good skin tolerability	Potential of transfer to a female partner or child by direct skin-to-skin contact; skin irritation in a small proportion of treated men; moderately high DHT levels; considerable interindividual and intraindividual variation in on-treatment T levels
Transdermal testosterone patch	1 or 2 patches, designed to nominally deliver 5–10 mg T over 24 h applied every day on nonpressure areas	Restores serum T, DHT, and E ₂ levels to the physiologic male range	T:DHT and T:E ₂ levels are in the physiologic male range	Ease of application, corrects symptoms of androgen deficiency	Serum T levels in some androgen-deficient men may be in the low-normal range; these men may need application of 2 patches daily; skin irritation at the application site occurs frequently in many patients
Buccal, bioadhesive, testosterone tablets	30-mg controlled-release, bioadhesive tablets bid	Absorbed from the buccal mucosa	Normalizes serum T and DHT levels in hypogonadal men	Corrects symptoms of androgen deficiency in healthy, hypogonadal men	Gum-related adverse events in 16% of treated men
Testosterone pellets	2–6 pellets implanted SC; dose and regimen vary with formulation	Serum T peaks at 1 month and then is sustained in normal range for 3–6 months, depending on formulation	T:DHT and T:E ₂ ratios do not change	Corrects symptoms of androgen deficiency	Requires surgical incision for insertions; pellets may extrude spontaneously
17-α-Methyl testosterone	This 17-α-alkylated compound should not be used because of potential for liver toxicity	Orally active			Clinical responses are variable; potential for liver toxicity; should not be used for treatment of androgen deficiency
Oral testosterone undecanoate ^a	40–80 mg PO bid or tid with meals	When administered in oleic acid, T undecanoate is absorbed through the lymphatics, bypassing the portal system; considerable variability in the same individual on different days and among individuals	High DHT:T ratio	Convenience of oral administration	Not approved in the United States; variable clinical responses, variable serum T levels, high DHT:T ratio
Injectable long-acting testosterone undecanoate in oil ^a	European regimen 1000 mg IM, followed by 1000 mg at 6 weeks, and 1000 mg every 10–14 weeks	When administered at a dose of 750–1000 mg IM, serum T levels are maintained in the normal range in a majority of treated men	DHT and E ₂ levels rise in proportion to the increase in T levels; T:DHT and T:E ₂ ratios do not change	Corrects symptoms of androgen deficiency; requires infrequent administration	Requires IM injection of a large volume (4 mL); cough reported immediately after injection in a very small number of men
Testosterone-in-adhesive matrix patch ^a	2 × 60 cm ² patches delivering approximately 4.8 mg of T/d	Restores serum T, DHT, and E ₂ to the physiologic range	T:DHT and T:E ₂ are in the physiologic range	Lasts 2 d	Some skin irritation

^aThese formulations are not approved for clinical use in the United States, but are available outside the United States in many countries. Physicians in those countries where these formulations are available should follow the approved drug regimens.

Abbreviations: DHT, dihydrotestosterone; E₂, estradiol; T, testosterone.

clear whether low-normal testosterone levels can maintain bone mineral density and muscle mass. The current recommendation is to restore testosterone levels to the mid-normal range.

Oral Derivatives of Testosterone Testosterone is well-absorbed after oral administration but is quickly degraded during the first pass through the liver. Therefore, it is difficult to achieve sustained blood levels of testosterone after oral administration of crystalline testosterone. 17 α -Alkylated derivatives of testosterone (e.g., 17 α -methyl testosterone, oxandrolone, fluoxymesterone) are relatively resistant to hepatic degradation and can be administered orally; however, because of the potential for hepatotoxicity, including cholestatic jaundice, peliosis, and hepatoma, these formulations should not be used for testosterone replacement. Hereditary angioedema due to C1 esterase deficiency is the only exception to this general recommendation; in this condition, oral 17 α -alkylated androgens are useful because they stimulate hepatic synthesis of the C1 esterase inhibitor.

Injectable Forms of Testosterone The esterification of testosterone at the 17 β -hydroxy position makes the molecule hydrophobic and extends its duration of action. The slow release of testosterone ester from an oily depot in the muscle accounts for its extended duration of action. The longer the side chain, the greater is the hydrophobicity of the ester and the longer is the duration of action. Thus, testosterone enanthate, cypionate, and undecanoate with longer side chains have longer duration of action than testosterone propionate. Within 24 h after intramuscular administration of 200 mg testosterone enanthate or cypionate, testosterone levels rise into the high-normal or supraphysiologic range and then gradually decline into the hypogonadal range over the next 2 weeks. A bimonthly regimen of testosterone enanthate or cypionate therefore results in peaks and troughs in testosterone levels that are accompanied by changes in a patient's mood, sexual desire, and energy level. The kinetics of testosterone enanthate and cypionate are similar. Estradiol and DHT levels are normal if testosterone replacement is physiologic.

Transdermal Testosterone Patch The nongenital testosterone patch, when applied in an appropriate dose, can normalize testosterone, DHT, and estradiol levels 4–12 h after application. Sexual function and well-being are restored in androgen-deficient men treated with the nongenital patch. One 5-mg patch may not be sufficient to increase testosterone into the mid-normal male range in all hypogonadal men; some patients may need two 5-mg patches daily to achieve the targeted testosterone concentrations. The use of testosterone patches may be associated with skin irritation in some individuals.

Testosterone Gel Several transdermal testosterone gels (e.g., Androgel, Testim, Fortesta, and Axiron), when applied topically to the skin in appropriate doses (Table 411-3), can maintain total and free testosterone concentrations in the normal range in hypogonadal men. The current recommendations are to begin with an initial U.S. Food and Drug Administration–approved dose and adjust the dose based on testosterone levels. The advantages of the testosterone gel include the ease of application and its flexibility of dosing. A major concern is the potential for inadvertent transfer of the gel to a sexual partner or to children who may come in close contact with the patient. The ratio of DHT to testosterone concentrations is higher in men treated with the testosterone gel than in healthy men. Also, there is considerable intra- and interindividual variation in serum testosterone levels in men treated with the transdermal gel due to variations in transdermal absorption and plasma clearance of testosterone. Therefore, monitoring of serum testosterone levels and multiple dose adjustments may be required to achieve and maintain testosterone levels in the target range.

Buccal Adhesive Testosterone A buccal testosterone tablet, which adheres to the buccal mucosa and releases testosterone as it is slowly dissolved, has been approved. After twice-daily application of 30-mg tablets, serum testosterone levels are maintained within

the normal male range in a majority of treated hypogonadal men. The adverse effects include buccal ulceration and gum problems in a few subjects. The effects of food and brushing on absorption have not been studied in detail.

Implants of crystalline testosterone can be inserted in the subcutaneous tissue by means of a trocar through a small skin incision. Testosterone is released by surface erosion of the implant and absorbed into the systemic circulation. Two to six 200-mg implants can maintain testosterone in the mid- to high-normal range for up to 6 months. Potential drawbacks include incising the skin for insertion and removal and spontaneous extrusions and fibrosis at the site of the implant.

Testosterone Formulations Not Available in the United States

Testosterone undecanoate, when administered orally in oleic acid, is absorbed preferentially through the lymphatics into the systemic circulation and is spared the first-pass degradation in the liver. Doses of 40–80 mg orally, two or three times daily, are typically used. However, the clinical responses are variable and suboptimal. DHT-to-testosterone ratios are higher in hypogonadal men treated with oral testosterone undecanoate, as compared to eugonadal men.

After initial priming, long-acting testosterone undecanoate in oil, when administered intramuscularly every 12 weeks, maintains serum testosterone, estradiol, and DHT in the normal male range and corrects symptoms of androgen deficiency in a majority of treated men. However, large injection volume (4 mL) is its relative drawback.

Novel Androgen Formulations A number of androgen formulations with better pharmacokinetics or more selective activity profiles are under development. A long-acting ester, testosterone undecanoate, when injected intramuscularly, can maintain circulating testosterone concentrations in the male range for 7–12 weeks. Initial clinical trials have demonstrated the feasibility of administering testosterone by the sublingual or buccal routes. 7 α -Methyl-19-nortestosterone is an androgen that cannot be 5 α -reduced; therefore, compared to testosterone, it has relatively greater agonist activity in muscle and gonadotropin suppression but lesser activity on the prostate.

Selective AR modulators (SARMs) are a class of AR ligands that bind the AR and display tissue-selective actions. A number of nonsteroidal SARMs that act as full agonists on the muscle and bone and that spare the prostate to varying degrees have advanced to phase 3 human trials. Nonsteroidal SARMs do not serve as substrates for either the steroid 5 α -reductase or the CYP19 aromatase. SARM binding to AR induces specific conformational changes in the AR protein, which then modulates protein-protein interactions between AR and its coregulators, resulting in tissue-specific regulation of gene expression.

Pharmacologic Uses of Androgens Androgens and SARMs are being evaluated as anabolic therapies for functional limitations associated with aging and chronic illness. Testosterone supplementation increases skeletal muscle mass, maximal voluntary strength, and muscle power in healthy men, hypogonadal men, older men with low testosterone levels, HIV-infected men with weight loss, and men receiving glucocorticoids. These anabolic effects of testosterone are related to testosterone dose and circulating concentrations. Systematic reviews have confirmed that testosterone therapy of HIV-infected men with weight loss promotes improvements in body weight, lean body mass, muscle strength, and depression indices, leading to the recommendation that testosterone be considered as an adjunctive therapy in HIV-infected men who are experiencing unexplained weight loss and who have low testosterone levels. Similarly, in glucocorticoid-treated men, testosterone therapy should be considered to maintain muscle mass and strength and vertebral bone mineral density. It is unknown whether testosterone therapy of older men with functional limitations is safe and effective in improving physical function, vitality, and health-related quality of life and reducing disability. Concerns about potential adverse effects of testosterone on prostate and cardiovascular event rates

2372 have encouraged the development of SARMs that are preferentially anabolic and spare the prostate.

Testosterone administration induces hypertrophy of both type 1 and 2 fibers and increases satellite cell (muscle progenitor cells) and myonuclear number. Androgens promote the differentiation of mesenchymal, multipotent progenitor cells into the myogenic lineage and inhibit their differentiation into the adipogenic lineage. Testosterone may have additional effects on satellite cell replication and muscle protein synthesis, which may contribute to an increase in skeletal muscle mass.

Other indications for androgen therapy are in selected patients with anemia due to bone marrow failure (an indication largely supplanted by erythropoietin) or for hereditary angioedema.

Male Hormonal Contraception Based on Combined Administration of Testosterone and Gonadotropin Inhibitors Supraphysiologic doses of testosterone (200 mg testosterone enanthate weekly) suppress LH and FSH secretion and induce azoospermia in 50% of Caucasian men and >95% of Chinese men. The WHO-supported multicenter efficacy trials have demonstrated that suppression of spermatogenesis to azoospermia or severe oligozoospermia (<3 million/mL) by administration of testosterone enanthate to men results in highly effective contraception. Because of concern about long-term adverse effects of supraphysiologic testosterone doses, regimens that combine other gonadotropin inhibitors, such as GnRH antagonists and progestins with replacement doses of testosterone, are being investigated. Oral etonogestrel daily in combination with intramuscular testosterone decanoate every 4–6 weeks induced azoospermia or severe oligozoospermia (sperm density <1 million/mL) in 99% of treated men over a 1-year period. This regimen was associated with weight gain, decreased testicular volume, and decreased plasma high-density lipoprotein (HDL) cholesterol, and its long-term safety has not been demonstrated. SARMs that are more potent inhibitors of gonadotropins than testosterone and spare the prostate hold promise for their contraceptive potential.

Recommended Regimens for Androgen Replacement Testosterone esters are administered typically at doses of 75–100 mg intramuscularly every week, or 150–200 mg every 2 weeks. One or two 5-mg nongenital testosterone patches can be applied daily over the skin of the back, thigh, or upper arm away from pressure areas. Testosterone gels are typically applied over a covered area of skin at initial doses that vary with the formulation; patients should wash their hands after gel application. Bioadhesive buccal testosterone tablets at a dose of 30 mg are typically applied twice daily on the buccal mucosa.

Establishing Efficacy of Testosterone Replacement Therapy Because a clinically useful marker of androgen action is not available, restoration of testosterone levels to the mid-normal range remains the goal of therapy. Measurements of LH and FSH are not useful in assessing the adequacy of testosterone replacement. Testosterone should be measured 3 months after initiating therapy to assess adequacy of therapy. There is substantial interindividual variability in serum testosterone levels, especially with transdermal gels, presumably due to genetic differences in testosterone clearance and transdermal absorption. In patients who are treated with testosterone enanthate or cypionate, testosterone levels should be 350–600 ng/dL 1 week after the injection. If testosterone levels are outside this range, adjustments should be made either in the dose or in the interval between injections. In men on transdermal patch, gel, or buccal testosterone therapy, testosterone levels should be in the mid-normal range (500–700 ng/dL) 4–12 h after application. If testosterone levels are outside this range, the dose should be adjusted. Multiple dose adjustments are often necessary to achieve testosterone levels in the desired therapeutic range.

Restoration of sexual function, secondary sex characteristics, energy, and well-being and maintenance of muscle and bone health are important objectives of testosterone replacement therapy. The patient should be asked about sexual desire and activity,

the presence of early morning erections, and the ability to achieve and maintain erections adequate for sexual intercourse. Some hypogonadal men continue to complain about sexual dysfunction even after testosterone replacement has been instituted; these patients may benefit from counseling. The hair growth in response to androgen replacement is variable and depends on ethnicity. Hypogonadal men with prepubertal onset of androgen deficiency who begin testosterone therapy in their late twenties or thirties may find it difficult to adjust to their newly found sexuality and may benefit from counseling. If the patient has a sexual partner, the partner should be included in counseling because of the dramatic physical and sexual changes that occur with androgen treatment.

Contraindications for Androgen Administration Testosterone administration is contraindicated in men with a history of prostate or breast cancer (Table 411-4). Testosterone therapy should not be administered without further urologic evaluation to men with a palpable prostate nodule or induration; to men with prostate-specific antigen levels >4 ng/mL or >3 ng/mL in men at high risk for prostate cancer such as African Americans or men with first-degree relatives with prostate cancer; or to men with severe lower urinary tract symptoms (American Urological Association lower urinary tract symptom score >19). Testosterone replacement should not be administered to men with baseline hematocrit \geq 50%, severe untreated obstructive sleep apnea, uncontrolled or poorly controlled congestive heart failure, or myocardial infarction, stroke, or acute coronary syndrome in the preceding 6 months.

Monitoring Potential Adverse Experiences The clinical effectiveness and safety of testosterone replacement therapy should be assessed 3 to 6 months after initiating testosterone therapy and annually thereafter (Table 411-5). Potential adverse effects include acne, oiliness of skin, erythrocytosis, breast tenderness and enlargement, leg edema, induction and exacerbation of obstructive sleep apnea, and increased risk of detection of prostate events. In addition, there may be formulation-specific adverse effects such as skin irritation with transdermal patch, risk of gel transfer to a sexual partner with testosterone gels, buccal ulceration and gum problems with buccal testosterone, and pain and mood fluctuation with injectable testosterone esters. Older men with preexisting heart disease may be at increased risk of cardiovascular events after initiation of testosterone therapy.

HEMOGLOBIN LEVELS Administration of testosterone to androgen-deficient men is typically associated with a ~3% increase in hemoglobin

TABLE 411-4 CONDITIONS IN WHICH TESTOSTERONE ADMINISTRATION IS ASSOCIATED WITH A RISK OF ADVERSE OUTCOME

Conditions in which testosterone administration is associated with very high risk of serious adverse outcomes:

- Metastatic prostate cancer
- Breast cancer

Conditions in which testosterone administration is associated with moderate to high risk of adverse outcomes:

- Undiagnosed prostate nodule or induration
- PSA >4 ng/mL (>3 ng/mL in individuals at high risk for prostate cancer, such as African Americans or men with first-degree relatives who have prostate cancer)
- Erythrocytosis (hematocrit >50%)
- Severe lower urinary tract symptoms associated with benign prostatic hypertrophy as indicated by American Urological Association/International Prostate Symptom Score >19
- Uncontrolled or poorly controlled congestive heart failure
- Myocardial infarction, stroke, or acute coronary syndrome in the preceding 6 months

Abbreviation: PSA, prostate-specific antigen.

Source: Reproduced from the Endocrine Society Guideline for Testosterone Therapy of Androgen Deficiency Syndromes in Men (S Bhasin et al: J Clin Endocrinol Metab 95:2536, 2010).

TABLE 411-5 MONITORING MEN RECEIVING TESTOSTERONE THERAPY

- Evaluate the patient 3–6 months after treatment initiation and then annually to assess whether symptoms have responded to treatment and whether the patient is suffering from any adverse effects.
- Monitor testosterone level 3–6 months after initiation of testosterone therapy:
 - Therapy should aim to raise serum testosterone level into the mid-normal range.
 - Injectable testosterone enanthate or cypionate: Measure serum testosterone level midway between injections. If testosterone is >700 ng/dL (24.5 nmol/L) or >400 ng/dL (14.1 nmol/L), adjust dose or frequency.
 - Transdermal patches: Assess testosterone level 3–12 h after application of the patch; adjust dose to achieve testosterone level in the mid-normal range.
 - Buccal testosterone bioadhesive tablet: Assess level immediately before or after application of fresh system.
 - Transdermal gels and solution: Assess testosterone level 2–8 h after patient has been on treatment for at least 2 weeks; adjust dose to achieve serum testosterone level in the mid-normal range.
 - Testosterone pellets: Measure testosterone levels at the end of the dosing interval. Adjust the number of pellets and/or the dosing interval to achieve serum testosterone levels in the normal range.
 - Oral testosterone undecanoate^a: Monitor serum testosterone level 3–5 h after ingestion.
 - Injectable testosterone undecanoate: Measure serum testosterone level just prior to each subsequent injection and adjust the dosing interval to maintain serum testosterone in mid-normal range.
- Check hematocrit at baseline, at 3–6 months, and then annually. If hematocrit is >54%, stop therapy until hematocrit decreases to a safe level; evaluate the patient for hypoxia and sleep apnea; reinstate therapy with a reduced dose.
- Measure bone mineral density of lumbar spine and/or femoral neck after 1–2 years of testosterone therapy in hypogonadal men with osteoporosis or low trauma fracture, consistent with regional standard of care.
- In men 40 years of age or older with baseline PSA >0.6 ng/mL, perform digital rectal examination and check PSA level before initiating treatment, at 3–6 months, and then in accordance with guidelines for prostate cancer screening depending on the age and race of the patient.
- Obtain urologic consultation if there is:
 - An increase in serum PSA concentration >1.4 ng/mL within any 12-month period of testosterone treatment.
 - A PSA velocity of >0.4 ng/mL per year using the PSA level after 6 months of testosterone administration as the reference (only applicable if PSA data are available for a period exceeding 2 years).
 - Detection of a prostatic abnormality on digital rectal examination.
 - An AUA/IPSS prostate symptom score of >19.
- Evaluate formulation-specific adverse effects at each visit:
 - Buccal testosterone tablets: Inquire about alterations in taste and examine the gums and oral mucosa for irritation.
 - Injectable testosterone esters (enanthate, cypionate, and undecanoate): Ask about fluctuations in mood or libido and, rarely, cough after injections.
 - Testosterone patches: Look for skin reaction at the application site.
 - Testosterone gels: Advise patients to cover the application sites with a shirt and to wash the skin with soap and water before having skin-to-skin contact, because testosterone gels leave a testosterone residue on the skin that can be transferred to a woman or child who might come in close contact. Serum testosterone levels are maintained when the application site is washed 4–6 h after application of the testosterone gel.
 - Testosterone pellets: Look for signs of infection, fibrosis, or pellet extrusion.

^aNot approved for clinical use in the United States.

Abbreviations: AUA/IPSS, American Urological Association/International Prostate Symptom Score; PSA, prostate-specific antigen.

Source: Reproduced with permission from the Endocrine Society Guideline for Testosterone Therapy of Androgen Deficiency Syndromes in Adult Men (S Bhasin et al: J Clin Endocrinol Metab 95:2536, 2010).

levels, due to increased erythropoiesis, suppression of hepcidin, and increased iron availability for erythropoiesis. The magnitude of hemoglobin increase during testosterone therapy is greater in older men than younger men and in men who have sleep apnea, a significant smoking history, or chronic obstructive lung disease. The frequency of erythrocytosis is higher in hypogonadal men treated with injectable testosterone esters than in those treated with transdermal formulations, presumably due to the higher testosterone dose delivered by the typical regimens of testosterone esters. Erythrocytosis is the most frequent adverse event reported in testosterone trials in middle-aged and older men and is also the most frequent cause of treatment discontinuation in these trials. If hematocrit rises above 54%, testosterone therapy should be stopped until hematocrit has fallen to <50%. After evaluation of the patient for hypoxia and sleep apnea, testosterone therapy may be reinstated at a lower dose.

PROSTATE AND SERUM PROSTATE-SPECIFIC ANTIGEN LEVELS Testosterone replacement therapy increases prostate volume to the size seen in age-matched controls but does not increase prostate volume beyond that expected for age. There is no evidence that testosterone therapy causes prostate cancer. However, androgen administration can exacerbate preexisting metastatic prostate cancer. Many older men harbor microscopic foci of cancer in their prostates. It

is not known whether long-term testosterone administration will induce these microscopic foci to grow into clinically significant cancers.

Prostate-specific antigen (PSA) levels are lower in testosterone-deficient men and are restored to normal after testosterone replacement. There is considerable test-retest variability in PSA measurements. Increments in PSA levels after testosterone supplementation in androgen-deficient men are generally <0.5 ng/mL, and increments >1.0 ng/mL over a 3- to 6-month period are unusual. The 90% confidence interval for the change in PSA values in men with benign prostatic hypertrophy, measured 3–6 months apart, is 1.4 ng/mL. Therefore, the Endocrine Society expert panel suggested that an increase in PSA >1.4 ng/mL in any 1 year after starting testosterone therapy, if confirmed, should lead to urologic evaluation. PSA velocity criterion can be used for patients who have sequential PSA measurements for >2 years; a change of >0.40 ng/mL per year merits closer urologic follow-up.

CARDIOVASCULAR RISK In epidemiologic studies, testosterone concentrations are negatively related to the risk of diabetes mellitus, heart disease, and all-cause and cardiovascular mortality. A recent testosterone trial in older men with mobility limitation was stopped early because of the higher rates of cardiovascular events in the testosterone arm than in the placebo arm of this trial. Meta-analyses

2374 of testosterone trials have found a significant increase in cardiovascular event rates in older men receiving testosterone therapy. Inferences about adverse events from previous trials included in these meta-analyses are limited by poor ascertainment, small numbers of events, heterogeneity of study populations, and small numbers of participants. Two retrospective analyses also found a higher frequency of cardiovascular events in association with testosterone therapy in older men with preexisting heart disease. Retrospective database analyses are limited by their inherent inability to verify the indication for treatment, diagnoses, or other relevant quantitative information and are susceptible to confounding by many other factors. Adequately powered prospective studies are needed to determine the effect on testosterone replacement on cardiovascular risk.

Androgen Abuse by Athletes and Recreational Bodybuilders The illicit use of androgenic-anabolic steroids (AAS) to enhance athletic performance first surfaced in the 1950s among power lifters and spread rapidly to other sports, professional as well as high school athletes, and recreational bodybuilders. In the early 1980s, the use of AAS spread beyond the athletic community into the general population, and now as many as 3 million Americans, most of them men, have likely used these compounds. Most AAS users are not athletes, but rather recreational weightlifters, who use these drugs to look lean and more muscular. The most commonly used AAS include testosterone esters, nandrolone, stanozolol, methandienone, and methenolol. AAS users generally use increasing doses of multiple steroids in a practice known as stacking.

The adverse effects of long-term AAS abuse remain poorly understood. Most of the information about the adverse effects of AAS has emerged from case reports, uncontrolled studies, or clinical trials that used replacement doses of testosterone. The adverse event data from clinical trials using physiologic replacement doses of testosterone have been extrapolated unjustifiably to AAS users who may administer 10–100 times the replacement doses of testosterone over many years and to support the claim that AAS use is safe. A substantial fraction of androgenic steroid users also use other drugs that are perceived to be muscle building or performance enhancing, such as GH; erythropoiesis-stimulating agents; insulin; and stimulants such as amphetamine, clenbuterol, cocaine, ephedrine, and thyroxine; and drugs perceived to reduce adverse effects such as hCG, aromatase inhibitors, or estrogen antagonists. The men who abuse androgenic steroids are more likely to engage in other high-risk behaviors than nonusers. The adverse events associated with AAS use may be due to AAS themselves, concomitant use of other drugs, high-risk behaviors, and host characteristics that may render these individuals more susceptible to AAS use or to other high-risk behaviors.

The high rates of mortality and morbidities observed in AAS users are alarming. One Finnish study reported 4.6 times the risk of death among elite power lifters than in age-matched men from the general population. The causes of death among power lifters included suicides, myocardial infarction, hepatic coma, and non-Hodgkin's lymphoma. A retrospective review of patient records in Sweden also reported higher standardized mortality ratios for AAS users than for nonusers. Thiblin and colleagues found that 32% of deaths among AAS users were suicidal, 26% homicidal, and 35% accidental. The median age of death among AAS users (24 years) is even lower than that for heroin or amphetamine users.

Numerous reports of cardiac death among young AAS users raise concerns about the adverse cardiovascular effects of AAS. High doses of AAS may induce proatherogenic dyslipidemia, increase thrombosis risk via effects on clotting factors and platelets, and induce vasospasm through their effects on vascular nitric oxide.

Replacement doses of testosterone, when administered parenterally, are associated with only a small decrease in HDL cholesterol and little or no effect on total cholesterol, low-density lipoprotein (LDL) cholesterol, and triglyceride levels. In contrast, supraphysiologic doses of testosterone and orally administered, 17 α -alkylated, nonaromatizable AAS are associated with marked reductions in HDL cholesterol and increases in LDL cholesterol.

Recent studies of AAS users using tissue Doppler and strain imaging and MRI have reported diastolic and systolic dysfunction, including significantly lower early and late diastolic tissue velocities, reduced E/A ratio, and reduced peak systolic strain in AAS users than in nonusers. Power athletes using AAS often have short QT intervals but increased QT dispersion, which may predispose them to ventricular arrhythmias. Long-term AAS use may be associated with myocardial hypertrophy and fibrosis. Myocardial tissue of power lifters using AAS has been shown to be infiltrated with fibrous tissue and fat droplets. The finding of ARs on myocardial cells suggests that AAS might be directly toxic to myocardial cells.

Long-term AAS use suppresses LH and FSH secretion and inhibits endogenous testosterone production and spermatogenesis. Men who have used AAS for more than a few months experience marked suppression of the hypothalamic-pituitary-testicular (HPT) axis after stopping AAS that may be associated with sexual dysfunction, fatigue, infertility, and depression; in some AAS users, HPT suppression may last more than a year, and in a few individuals, complete recovery may never occur. The symptoms of androgen deficiency caused by androgen withdrawal may cause some men to revert back to using AAS, leading to continued use and AAS dependence. As many as 30% of AAS users develop a syndrome of AAS dependence, characterized by long-term AAS use despite adverse medical and psychiatric effects.

Supraphysiologic doses of testosterone may also impair insulin sensitivity. Orally administered androgens also have been associated with insulin resistance and diabetes.

Unsafe injection practices, high-risk behaviors, and increased rates of incarceration render AAS users at increased risk of HIV and hepatitis B and C. In one survey, nearly 1 in 10 gay men had injected AAS or other substances, and AAS users were more likely to report high-risk unprotected anal sex than other men.

Some AAS users develop hypomanic or manic symptoms during AAS exposure (irritability, aggressiveness, reckless behavior, and occasional psychotic symptoms, sometimes associated with violence) and major depression (sometimes associated with suicidality) during AAS withdrawal. Users may also develop other forms of illicit drug use, which may be potentiated or exacerbated by AAS.

Elevated liver enzymes, cholestatic jaundice, hepatic neoplasms, and peliosis hepatis have been reported with oral, 17 α -alkylated AAS. AAS use may cause muscle hypertrophy without compensatory adaptations in tendons, ligaments, and joints, thus increasing the risk of tendon and joint injuries. AAS use is associated with acne, baldness, and increased body hair.

The suspicion of AAS use may be raised by the increased hemoglobin and hematocrit, suppressed LH and FSH and testosterone levels, low high-density lipoproteins cholesterol, and low testicular volume and sperm density in a person who looks highly muscular. Accredited laboratories use gas chromatography–mass spectrometry or liquid chromatography–mass spectrometry to detect anabolic steroid abuse. In recent years, the availability of high-resolution mass spectrometry and tandem mass spectrometry has further improved the sensitivity of detecting androgen abuse. Illicit testosterone use is detected generally by the application of the measurement of the urinary testosterone-to-epitestosterone ratio and further confirmed by the use of the $^{13}\text{C}:^{12}\text{C}$ ratio in testosterone by the use of isotope ratio combustion mass spectrometry. Exogenous testosterone administration increases urinary testosterone glucuronide excretion and consequently the testosterone-to-epitestosterone ratio. Ratios >4 suggest exogenous testosterone use but can also reflect genetic variation. Genetic variations in the uridine diphosphoglucuronyl transferase 2B17 (*UGT2B17*), the major enzyme for testosterone glucuronidation, affect the testosterone-to-epitestosterone ratio. Synthetic testosterone has a lower $^{13}\text{C}:^{12}\text{C}$ ratio than endogenously produced testosterone, and these differences in $^{13}\text{C}:^{12}\text{C}$ ratio can be detected by isotope ratio combustion mass spectrometry, which is used to confirm exogenous testosterone use in individuals with a high testosterone-to-epitestosterone ratio.

412 Disorders of the Female Reproductive System

Janet E. Hall

The female reproductive system regulates the hormonal changes responsible for puberty and adult reproductive function. Normal reproductive function in women requires the dynamic integration of hormonal signals from the hypothalamus, pituitary, and ovary, resulting in repetitive cycles of follicle development, ovulation, and preparation of the endometrial lining of the uterus for implantation should conception occur. It is critical to understand pubertal development in normal girls (and boys) as a yardstick for identifying precocious and delayed puberty.

For further discussion of related topics, see the following chapters: amenorrhea and pelvic pain (Chap. 69), infertility and contraception (Chap. 414), menopause (Chap. 413), disorders of sex development (Chap. 410), and disorders of the male reproductive system (Chap. 411).

DEVELOPMENT OF THE OVARY AND EARLY FOLLICULAR GROWTH

The ovary orchestrates the development and release of a mature oocyte and also elaborates hormones (e.g., estrogen, progesterone, inhibin, relaxin) that are critical for pubertal development and preparation of the uterus for conception, implantation, and the early stages of pregnancy. To achieve these functions in repeated monthly cycles, the ovary undergoes some of the most dynamic changes of any organ in the body. Primordial germ cells can be identified by the third week of gestation, and their migration to the genital ridge is complete by 6 weeks of gestation. Germ cells persist within the genital ridge, are then referred to as *oogonia*, and are essential for induction of ovarian development. Although one X chromosome undergoes X inactivation in somatic cells, it is reactivated in oogonia and genes on both

X chromosomes are required for normal ovarian development. A streak ovary containing only stromal cells is found in patients with 45,X Turner's syndrome (Chap. 410).

The germ cell population expands, and starting at ~8 weeks of gestation, oogonia begin to enter prophase of the first meiotic division and become primary oocytes. This allows the oocyte to be surrounded by a single layer of flattened granulosa cells to form a primordial follicle (Fig. 412-1). Granulosa cells are derived from mesonephric cells that invade the ovary early in its development, pushing the germ cells to the periphery. The weight of evidence supports the concept that for the most part, the ovary contains a nonrenewable pool of germ cells. Through the combined processes of mitosis, meiosis, and atresia, the population of oogonia reaches its maximum of 6–7 million by 20 weeks of gestation, after which there is a progressive loss of both oogonia and primordial follicles through the process of atresia. At birth, oogonia are no longer present in the ovary, and only 1–2 million germ cells remain in the form of primordial follicles (Fig. 412-2). The oocyte persists in prophase of the first meiotic division until just before ovulation, when meiosis resumes.

The quiescent primordial follicles are recruited to further growth and differentiation through a highly regulated process that limits the size of the developing cohort to ensure that folliculogenesis can continue throughout the reproductive life span. This initial recruitment of primordial follicles to form primary follicles (Fig. 412-1) is characterized by growth of the oocyte and the transition from squamous to cuboidal granulosa cells. The theca interna cells that surround the developing follicle begin to form as the primary follicle grows. Acquisition of a zona pellucida by the oocyte and the presence of several layers of surrounding cuboidal granulosa cells mark the development of secondary follicles. It is at this stage that granulosa cells develop follicle-stimulating hormone (FSH), estradiol, and androgen receptors and communicate with one another through the development of gap junctions.

Bidirectional signaling between the germ cells and the somatic cells in the ovary is a necessary component underlying the maturation

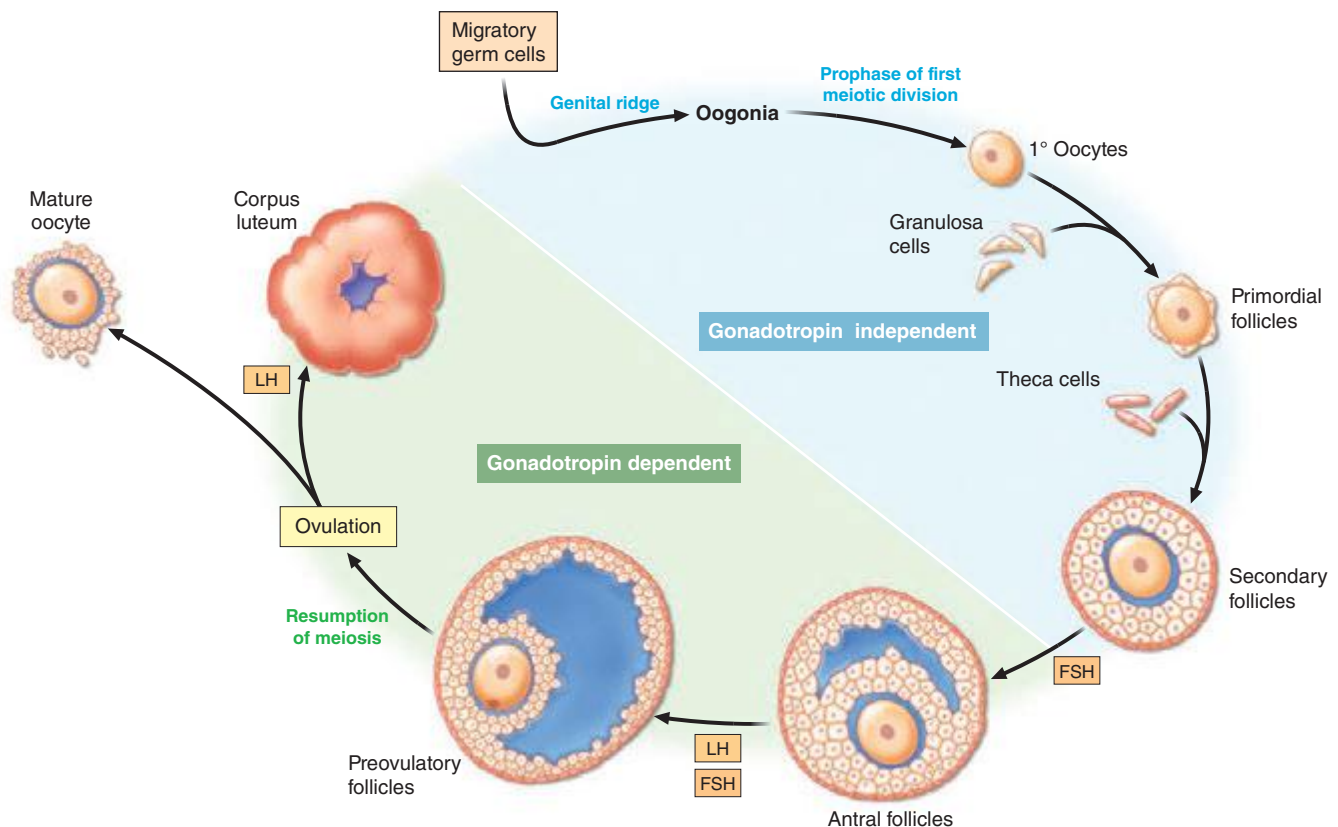


FIGURE 412-1 Stages of ovarian development from the arrival of the migratory germ cells at the genital ridge through gonadotropin-independent and gonadotropin-dependent phases that ultimately result in ovulation of a mature oocyte. FSH, follicle-stimulating hormone; LH, luteinizing hormone.

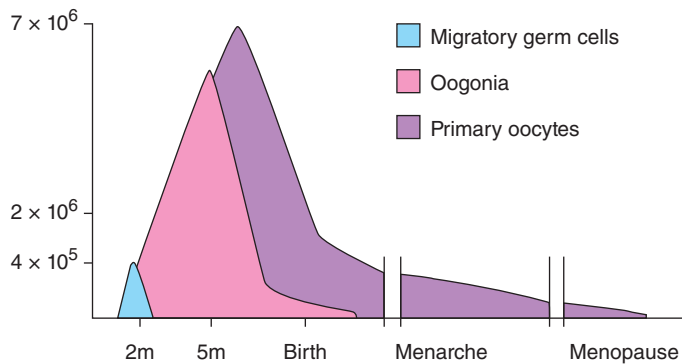


FIGURE 412-2 Ovarian germ cell number is maximal at mid-gestation and then decreases precipitously.

of the oocyte and the capacity for hormone secretion. For example, oocyte-derived growth differentiation factor 9 (GDF-9) and bone morphogenic protein-15 (BMP-15), also known as GDF-9b, are required for migration of pregranulosa and pretheca cells to the outer surface of the developing follicle and, hence, initial follicle formation. GDF-9 is also required for formation of secondary follicles, as are granulosa cell–derived KIT ligand (KITL) and the forkhead transcription factor (FOXL2). All of these genes are potential candidates for premature ovarian failure in women, and mutations in the human *FOXL2* gene have been shown to cause the syndrome of blepharophimosis/ptosis/epicanthus inversus, which is associated with ovarian failure.

DEVELOPMENT OF A MATURE FOLLICLE

The early stages of follicle growth are primarily driven by intraovarian factors and may take up to a year from development of the primary follicle to the dominant follicle stage. Further maturation to the preovulatory state, including the resumption of meiosis in the oocyte, requires the combined stimulus of FSH and luteinizing hormone (LH) (Fig. 412-1) and can be accomplished within weeks. Recruitment of secondary follicles from the resting follicle pool requires the direct action of FSH, whereas anti-müllerian hormone (AMH) produced from small growing follicles, restrains this effect of FSH. Accumulation of follicular fluid between the layers of granulosa cells creates an antrum that divides the granulosa cells into two functionally distinct groups: mural cells that line the follicle wall and cumulus cells that surround the oocyte (Fig. 412-3). Recent evidence suggests that, in addition to its role in normal development of the müllerian system, the WNT

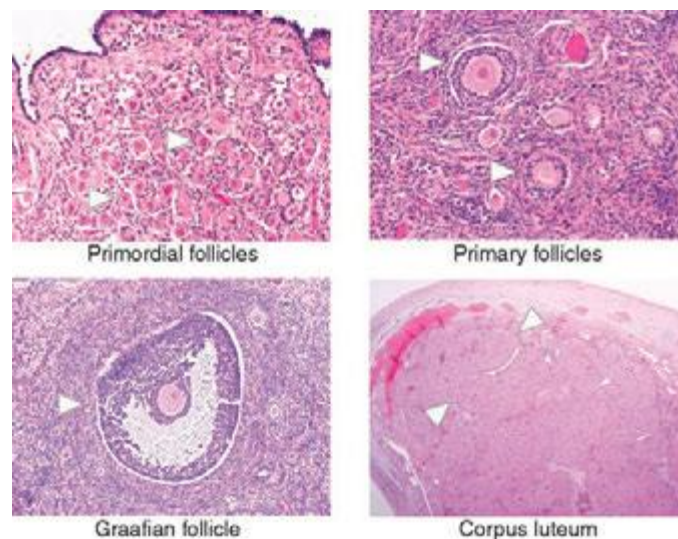


FIGURE 412-3 Development of ovarian follicles. The Graafian follicle is also known as a tertiary or preovulatory follicle. (Courtesy of JH Eichhorn and D. Roberts, Massachusetts General Hospital; with permission.)

signaling pathway is required for normal antral follicle development and may also play a role in ovarian steroidogenesis. A single dominant follicle emerges from the growing follicle pool within the first 5–7 days after the onset of menses, and the majority of follicles fall off their growth trajectory and become atretic. Autocrine actions of activin and BMP-6, derived from the granulosa cells, and paracrine actions of GDF-9, BMP-15, BMP-6, and Gpr149, derived from the oocyte, are involved in granulosa cell proliferation and modulation of FSH responsiveness. Differential exposure to these factors may explain the mechanism whereby a given follicle is selected for continued growth to the preovulatory stage. The dominant follicle can be distinguished by its size, evidence of granulosa cell proliferation, large number of FSH receptors, high aromatase activity, and elevated concentrations of estradiol and inhibin A in follicular fluid.

The dominant follicle undergoes rapid expansion during the 5–6 days prior to ovulation, reflecting granulosa cell proliferation and accumulation of follicular fluid. FSH induces LH receptors on the granulosa cells, and the preovulatory, or Graafian, follicle moves to the outer ovarian surface in preparation for ovulation. The LH surge triggers the resumption of meiosis, the suppression of granulosa cell proliferation, and the induction of cyclooxygenase 2 (COX-2), prostaglandins, the progesterone receptor, and the epidermal growth factor (EGF)-like growth factors amphiregulin, epiregulin, betacellulin, and neuroregulin 1, all of which are required for ovulation. EGF-like factors are thought to mediate these follicular responses to LH. Ovulation also involves production of extracellular matrix leading to expansion of the cumulus cell population that surrounds the oocyte and the controlled expulsion of the egg and follicular fluid. Both progesterone and prostaglandins (induced by the ovulatory stimulus) are essential for this process. After ovulation, luteinization is induced by LH in conjunction with the acquisition of a rich vascular network in response to vascular endothelial growth factor (VEGF) and basic fibroblast growth factor (FGF). Traditional regulators of central reproductive control, gonadotropin-releasing hormone (GnRH) and its receptor (GnRH-R), are also produced in the ovary and may be involved in corpus luteum function.

REGULATION OF OVARIAN FUNCTION

HYPOTHALAMIC AND PITUITARY SECRETION

GnRH neurons develop from epithelial cells outside the central nervous system and migrate, initially alongside the olfactory neurons, to the medial basal hypothalamus. Studies in GnRH-deficient patients who fail to undergo puberty have provided insights into genes that control the ontogeny and function of GnRH neurons (Fig. 412-4). *KAL1*, *FGF8/FGFR1*, *PROK2/PROKR2*, *NSMF*, and *CDH7*, among others (Chap. 411), have been implicated in the migration of GnRH neurons to the hypothalamus. Approximately 7000 GnRH neurons, scattered throughout the medial basal hypothalamus, establish contacts with capillaries of the pituitary portal system in the median eminence. GnRH is secreted into the pituitary portal system in discrete pulses to stimulate synthesis and secretion of LH and FSH from pituitary gonadotropes, which comprise ~10% of cells in the pituitary (Chap. 401e). Functional connections of GnRH neurons with the portal system are established by the end of the first trimester, coinciding with the production of pituitary gonadotropins. Thus, like the ovary, the hypothalamic and pituitary components of the reproductive system are present before birth. However, the high levels of estradiol and progesterone produced by the placenta suppress hypothalamic-pituitary stimulation of ovarian hormonal secretion in the fetus.

After birth and the loss of placenta-derived steroids, gonadotropin levels rise. FSH levels are much higher in girls than in boys. This rise in FSH results in ovarian activation (evident on ultrasound) and increased inhibin B and estradiol levels. Studies that have identified mutations in *TAC3*, which encodes neurokinin B, and its receptor, *TAC3R*, in patients with GnRH deficiency indicate that both are involved in control of GnRH secretion and may be particularly important at this early stage of development. By 12–20 months of age, the reproductive axis is again suppressed, and a period of relative quiescence persists until puberty (Fig. 412-5). At the onset of puberty,

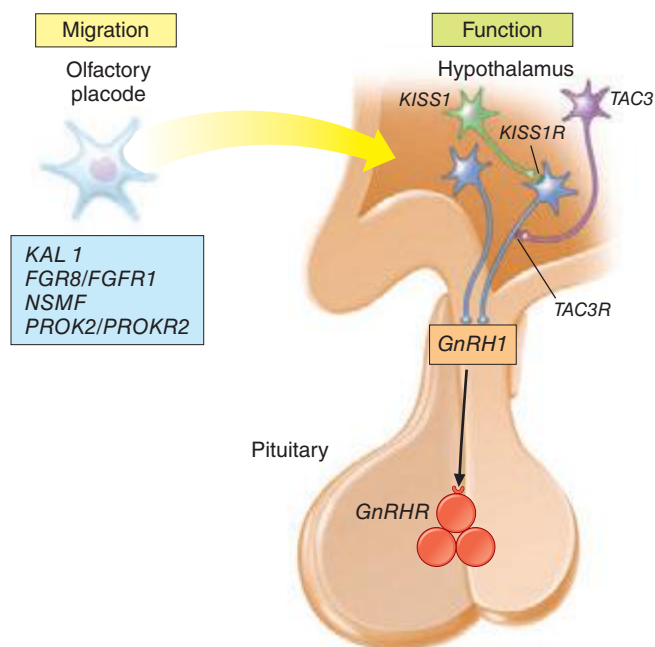


FIGURE 412-4 Establishment of a functional gonadotropin-releasing hormone (GnRH) system requires the participation of a number of genes that are essential for development and migration of GnRH neurons from the olfactory placode to the hypothalamus in addition to genes involved in the functional control of GnRH secretion and action.

pulsatile GnRH secretion induces pituitary gonadotropin production. In the early stages of puberty, LH and FSH secretion are apparent only during sleep, but as puberty develops, pulsatile gonadotropin secretion occurs throughout the day and night.

The mechanisms responsible for the childhood quiescence and pubertal reactivation of the reproductive axis remain incompletely understood. GnRH neurons in the hypothalamus respond to both excitatory and inhibitory factors. Increased sensitivity to the inhibitory influence of gonadal steroids has long been implicated in the inhibition of GnRH secretion during childhood but has not been definitively established in the human. Metabolic signals, such as adipocyte-derived leptin, play a permissive role in reproductive function (Chap. 415e). Studies of patients with isolated GnRH deficiency reveal that mutations in the G protein-coupled receptor 54 (*GPR54*) gene (now known as *KISS1R*) preclude the onset of puberty. The ligand for this receptor, metastin, is derived from the parent peptide, kisspeptin-1 (*KISS1*), and is a powerful stimulant for GnRH release. A potential role for kisspeptin in the onset of puberty has been suggested by upregulation of *KISS1* and *KISS1R* transcripts in the hypothalamus at the time of puberty. *TAC3* and dynorphin (*Dyn*), which appear to play an inhibitory rather than stimulatory role in GnRH control, are co-expressed

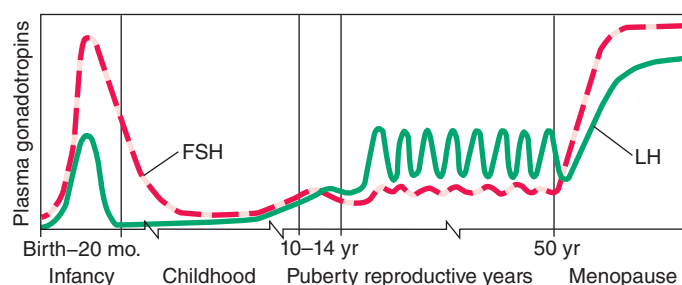


FIGURE 412-5 Follicle-stimulating hormone (FSH) and luteinizing hormone (LH) are increased during the neonatal years but go through a period of childhood quiescence before increasing again during puberty. Gonadotropin levels are cyclic during the reproductive years and increase dramatically with the loss of negative feedback that accompanies menopause.

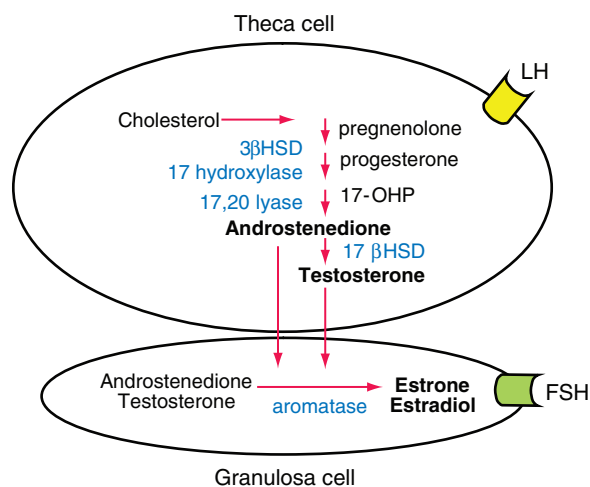


FIGURE 412-6 Estrogen production in the ovary requires the cooperative function of the theca and granulosa cells under the control of luteinizing hormone (LH) and follicle-stimulating hormone (FSH). HSD, hydroxysteroid dehydrogenase; OHP, hydroxyprogesterone.

with *KISS1* in KNDy neurons that project to GnRH neurons. This system is intimately involved with estrogen negative feedback regulation of GnRH secretion.

OVARIAN STEROIDS

Ovarian steroid-producing cells do not store hormones but produce them in response to LH and FSH during the normal menstrual cycle. The sequence of steps and the enzymes involved in the synthesis of steroid hormones are similar in the ovary, adrenal, and testis. However, the enzymes required to catalyze specific steps are compartmentalized and may not be abundant or even present in all cell types. Within the developing ovarian follicle, estrogen synthesis from cholesterol requires close integration between theca and granulosa cells—sometimes called the *two-cell model for steroidogenesis* (Fig. 412-6). FSH receptors are confined to the granulosa cells, whereas LH receptors are restricted to the theca cells until the late stages of follicular development, when they are also found on granulosa cells. The theca cells surrounding the follicle are highly vascularized and use cholesterol, derived primarily from circulating lipoproteins, as the starting point for the synthesis of androstenedione and testosterone under the control of LH. Androstenedione and testosterone are transferred across the basal lamina to the granulosa cells, which receive no direct blood supply. The mural granulosa cells are particularly rich in aromatase and, under the control of FSH, produce estradiol, the primary steroid secreted from the follicular phase ovary and the most potent estrogen. Theca cell-produced androstenedione and, to a lesser extent, testosterone are also secreted into peripheral blood, where they can be converted to dihydrotestosterone in skin and to estrogens in adipose tissue. The hilar interstitial cells of the ovary are functionally similar to Leydig cells and are also capable of secreting androgens. Although stromal cells proliferate in response to androgens (as in polycystic ovarian syndrome [PCOS]), they do not secrete androgens.

Development of the rich capillary network following rupture of the follicle at the time of ovulation makes it possible for large molecules such as low-density lipoprotein (LDL) to reach the luteinized granulosa and theca lutein cells. As in the follicle, both cell types are required for steroidogenesis in the corpus luteum. The large luteinized granulosa cells are the main source of progesterone production, whereas the smaller theca lutein cells produce 17-hydroxyprogesterone, a substrate for aromatization to estradiol by the luteinized granulosa cells. LH is critical for normal structure and function of the corpus luteum. Because LH and human chorionic gonadotropin (hCG) bind to a common receptor, the role of LH in support of the corpus luteum can be replaced by hCG in the first 10 weeks after conception, and hCG is commonly used for luteal phase support in the treatment of infertility.

2378 Steroid Hormone Actions Both estrogen and progesterone play critical roles in the expression of secondary sexual characteristics in women (**Chap. 400e**). Estrogen promotes development of the ductule system in the breast, whereas progesterone is responsible for glandular development. In the reproductive tract, estrogens create a receptive environment for fertilization and support pregnancy and parturition through carefully coordinated changes in the endometrium, thickening of the vaginal mucosa, thinning of the cervical mucus, and uterine growth and contractions. Progesterone induces secretory activity in the estrogen-primed endometrium, increases the viscosity of cervical mucus, and inhibits uterine contractions. Both gonadal steroids play critical roles in the negative and positive feedback controls of gonadotropin secretion. Progesterone also increases basal body temperature and has therefore been used clinically as a marker of ovulation.

The vast majority of circulating estrogens and androgens are carried in the blood bound to carrier proteins, which restrain their free diffusion into cells and prolong their clearance, serving as a reservoir. High-affinity binding proteins include sex hormone-binding globulin (SHBG), which binds androgens with somewhat greater affinity than estrogens, and corticosteroid-binding globulin (CBG), which also binds progesterone. Modulations in binding protein levels by insulin, androgens, and estrogens contribute to high bioavailable testosterone levels in PCOS and to high circulating estrogen and progesterone levels during pregnancy.

Estrogens act primarily through binding to the nuclear receptors, estrogen receptor (ER) α and β . Transcriptional coactivators and corepressors modulate ER action (**Chap. 400e**). Both ER subtypes are present in the hypothalamus, pituitary, ovary, and reproductive tract. Although ER α and β exhibit some functional redundancy, there is also a high degree of specificity, particularly in expression within cell types. For example, ER α functions in the ovarian theca cells, whereas ER β is critical for granulosa cell function. There is also evidence for membrane-initiated signaling by estrogen. Similar signaling mechanisms pertain for progesterone with evidence of transcriptional regulation through progesterone receptor (PR) A and B protein isoforms, as well as rapid membrane signaling.

OVARIAN PEPTIDES

Inhibin was initially isolated from gonadal fluids based on its ability to selectively inhibit FSH secretion from pituitary cells. Inhibin is a heterodimer composed of an α subunit and a β A or β B subunit to form inhibin A or inhibin B, both of which are secreted from the ovary. Activin is a homodimer of inhibin β subunits with the capacity to stimulate the synthesis and secretion of FSH. Inhibins and activins are members of the transforming growth factor β (TGF- β) superfamily of growth and differentiation factors. During the purification of inhibin, follistatin, an unrelated monomeric protein that inhibits FSH secretion, was discovered. Within the pituitary, follistatin inhibits FSH secretion indirectly through binding and neutralizing activin.

Inhibin B is secreted from the granulosa cells of small antral follicles, whereas inhibin A is present in both granulosa and theca cells and is secreted by dominant follicles. Inhibin A is also present in luteinized granulosa cells and is a major secretory product of the corpus luteum. Inhibin B is constitutively secreted by granulosa cells and increases in serum in conjunction with recruitment of secondary follicles to the pool of actively growing follicles under the control of FSH. Inhibin B has been used clinically as a marker of ovarian reserve. Inhibin B is an important inhibitor of FSH, independent of estradiol, during the menstrual cycle. Although activin is also secreted from the ovary, the excess of follistatin in serum, combined with its nearly irreversible binding of activin, make it unlikely that ovarian activin plays an endocrine role in FSH regulation. However, there is evidence that activin plays an autocrine/paracrine role in the ovary, in addition to its intrapituitary role in modulation of FSH production.

AMH (also known as müllerian-inhibiting substance) is important in ovarian biology in addition to the function from which it derived its name (i.e., promotion of the degeneration of the müllerian system during embryogenesis in the male). AMH is produced by granulosa cells from small follicles and, like inhibin B, is a marker of ovarian reserve.

AMH inhibits the recruitment of primordial follicles into the follicle pool and counters FSH stimulation of aromatase expression.

Relaxin, which is produced by the theca lutein cells of the corpus luteum, is thought to play a role in decidualization of the endometrium and suppression of myometrial contractile activity, both of which are essential for the early establishment of pregnancy.

HORMONAL INTEGRATION OF THE NORMAL MENSTRUAL CYCLE

The sequence of changes responsible for mature reproductive function is coordinated through a series of negative and positive feedback loops that alter pulsatile GnRH secretion, the pituitary response to GnRH, and the relative secretion of LH and FSH from the gonadotrope. The frequency and amplitude of pulsatile GnRH secretion differentially modulate the synthesis and secretion of LH and FSH, with slow frequencies favoring FSH synthesis and increased amplitudes favoring LH synthesis. Activin is produced in both pituitary gonadotropes and folliculostellate cells and stimulates the synthesis and secretion of FSH. Inhibins function as potent antagonists of activins through sequestration of the activin receptors. Although inhibin is expressed in the pituitary, gonadal inhibin is the principal source of feedback inhibition of FSH.

For the majority of the cycle, the reproductive system functions in a classic endocrine negative feedback mode. Estradiol and progesterone inhibit GnRH secretion, and the inhibins act at the pituitary to selectively inhibit FSH synthesis and secretion (**Fig. 412-7**). This negative feedback control of FSH is critical for development of the single mature oocyte that characterizes normal reproductive function in women. In addition to these negative feedback controls, the menstrual cycle is uniquely dependent on estrogen-induced positive feedback to produce an LH surge that is essential for ovulation of a mature follicle. Estrogen negative feedback in women occurs primarily at the hypothalamus with a small pituitary contribution, whereas estrogen positive feedback occurs at the pituitary with hypothalamic GnRH secretion playing a permissive role.

THE FOLLICULAR PHASE

The follicular phase is characterized by recruitment of a cohort of secondary follicles and the ultimate selection of a dominant preovulatory follicle (**Fig. 412-8**). The follicular phase begins, by convention, on the first day of menses. However, follicle recruitment is initiated by the rise in FSH that begins in the late luteal phase of the previous cycle in conjunction with the loss of negative feedback of gonadal steroids and likely inhibin A. The fact that a 20–30% increase in FSH is adequate for follicular recruitment speaks to the marked sensitivity of the resting follicle pool to FSH. The resultant granulosa cell proliferation is responsible for increasing early follicular phase levels of inhibin B. Inhibin B in conjunction with rising levels of estradiol, and

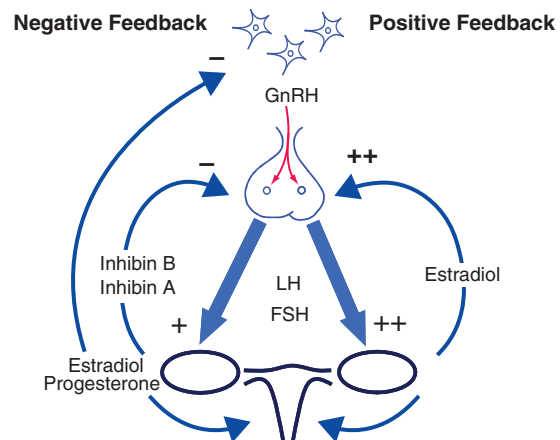


FIGURE 412-7 The reproductive system in women is critically dependent on both negative feedback of gonadal steroids and inhibin to modulate follicle-stimulating hormone (FSH) secretion and on estrogen positive feedback to generate the preovulatory luteinizing hormone (LH) surge. GnRH, gonadotropin-releasing hormone.

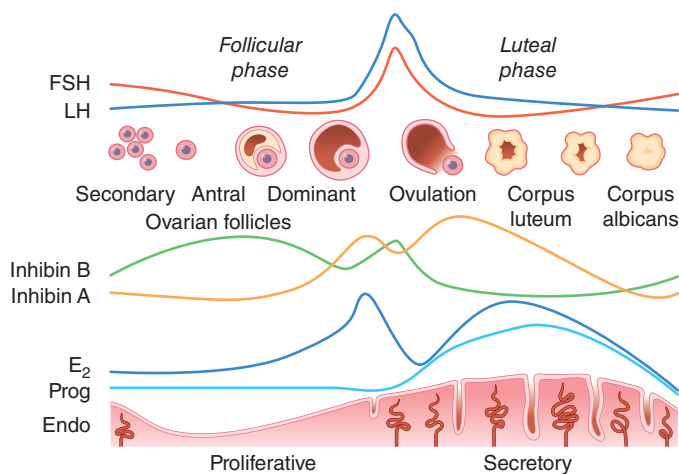


FIGURE 412-8 Relationship between gonadotropins, follicle development, gonadal secretion, and endometrial changes during the normal menstrual cycle. E₂, estradiol; Endo, endometrium; FSH, follicle-stimulating hormone; LH, luteinizing hormone; Prog, progesterone.

probably inhibin A, restrain FSH secretion during this critical period such that only a single follicle matures in the vast majority of cycles. The increased risk of multiple gestation associated with the increased levels of FSH characteristic of advanced maternal age, or with exogenous gonadotropin administration in the treatment of infertility, attests to the importance of negative feedback regulation of FSH. With further growth of the dominant follicle, estradiol and inhibin A increase exponentially and the follicle acquires LH receptors. Increasing levels of estradiol are responsible for proliferative changes in the endometrium. The exponential rise in estradiol results in positive feedback on the pituitary, leading to the generation of an LH surge (and a smaller FSH surge), thereby triggering ovulation and luteinization of the granulosa cells.

THE LUTEAL PHASE

The luteal phase begins with the formation of the corpus luteum from the ruptured follicle (Fig. 412-8). Progesterone and inhibin A are produced from the luteinized granulosa cells, which continue to aromatize theca-derived androgen precursors, producing estradiol. The combined actions of estrogen and progesterone are responsible for the secretory changes in the endometrium that are necessary for implantation. The corpus luteum is supported by LH but has a finite life span because of diminished sensitivity to LH. The demise of the corpus luteum results in a progressive decline in hormonal support of the endometrium. Inflammation or local hypoxia and ischemia result in vascular changes in the endometrium, leading to the release of cytokines, cell death, and shedding of the endometrium.

If conception occurs, hCG produced by the trophoblast binds to LH receptors on the corpus luteum, maintaining steroid hormone production and preventing involution of the corpus luteum. The corpus luteum is essential for the hormonal maintenance of the endometrium during the first 6–10 weeks of pregnancy until this function is taken over by the placenta.

CLINICAL ASSESSMENT OF OVARIAN FUNCTION

Menstrual bleeding should become regular within 2–4 years of menarche, although anovulatory and irregular cycles are common before that. For the remainder of adult reproductive life, the cycle length counted from the first day of menses to the first day of subsequent menses is ~28 days, with a range of 25–35 days. However, cycle-to-cycle variability for an individual woman is ± 2 days. Luteal phase length is relatively constant between 12 and 14 days in normal cycles; thus, the major variability in cycle length is due to variations in the follicular phase. The duration of menstrual bleeding in ovulatory cycles varies between 4 and 6 days. There is a gradual

shortening of cycle length with age such that women over the age of 35 have cycles that are shorter than during their younger reproductive years. Anovulatory cycles increase as women approach menopause, and bleeding patterns may be erratic.

Women who report regular monthly bleeding with cycles that do not vary by >4 days generally have ovulatory cycles, but several other clinical signs can be used to assess the likelihood of ovulation. Some women experience *mittelschmerz*, described as midcycle pelvic discomfort that is thought to be caused by the rapid expansion of the dominant follicle at the time of ovulation. A constellation of premenstrual moliminal symptoms such as bloating, breast tenderness, and food cravings often occur several days before menses in ovulatory cycles, but their absence cannot be used as evidence of anovulation. Methods that can be used to determine whether ovulation is likely include a serum progesterone level >5 ng/mL ~7 days before expected menses, an increase in basal body temperature of 0.24°C (>0.5°F) in the second half of the cycle due to the thermoregulatory effect of progesterone, or the detection of the urinary LH surge using ovulation predictor kits. Because ovulation occurs ~36 h after the LH surge, urinary LH can be helpful in timing intercourse to coincide with ovulation.

Ultrasound can be used to detect the growth of the fluid-filled antrum of the developing follicle and to assess endometrial proliferation in response to increasing estradiol levels in the follicular phase, as well as the characteristic echogenicity of the secretory endometrium of the luteal phase.

PUBERTY

NORMAL PUBERTAL DEVELOPMENT IN GIRLS

The first menstrual period (*menarche*) occurs relatively late in the series of developmental milestones that characterize normal pubertal development (Table 412-1). Menarche is preceded by the appearance of pubic and then axillary hair (*adrenarche*) as a result of maturation of the zona reticularis in the adrenal gland and increased adrenal androgen secretion, particularly dehydroepiandrosterone (DHEA). The triggers for adrenarche remain unknown but may involve increases in body mass index, as well as in utero and neonatal factors. Menarche is also preceded by breast development (*thelarche*). The breast is exquisitely sensitive to the very low levels of estrogen that result from peripheral conversion of adrenal androgens and the low levels of estrogen secreted from the ovary early in pubertal maturation. Breast development precedes the appearance of pubic and axillary hair in ~60% of girls. The interval between the onset of breast development and menarche is ~2 years. There has been a gradual decline in the age of menarche over the past century, attributed in large part to improvement in nutrition, and there is a relationship between adiposity and earlier sexual maturation in girls. In the United States, menarche occurs at an average age of 12.5 years (Table 412-1). Much of the variation in the timing of puberty is due to genetic factors, with heritability estimates of 50–80%. Both adrenarche and thelarche occur ~1 year earlier in black compared with white girls, although the timing of menarche differs by only 6 months between these ethnic groups.

Other important hormonal changes also occur in conjunction with puberty. Growth hormone (GH) levels increase early in puberty, stimulated in part by the pubertal increases in estrogen secretion. GH increases insulin-like growth factor-I (IGF-I), which enhances linear growth. The growth spurt is generally less pronounced in girls than in boys, with a peak growth velocity of ~7 cm/year. Linear growth is ultimately limited by closure of epiphyses in the long bones as a result of prolonged exposure to estrogen. Puberty is also associated with mild insulin resistance.

TABLE 412-1 MEAN AGE (YEARS) OF PUBERTAL MILESTONES IN GIRLS

	Onset of Breast/ Pubic Hair Development	Age of Peak Height Velocity	Menarche	Final Breast/ Pubic Hair Development	Adult Height
White	10.2	11.9	12.6	14.3	17.1
Black	9.6	11.5	12	13.6	16.5

Source: From FM Biro et al: J Pediatr 148:234, 2006.

TABLE 412-2 DIFFERENTIAL DIAGNOSIS OF PRECOCIOUS PUBERTY

Central (GnRH Dependent)	Peripheral (GnRH Independent)
Idiopathic	Congenital adrenal hyperplasia
CNS tumors	Estrogen-producing tumors
Hamartomas	Adrenal tumors
Astrocytomas	Ovarian tumors
Adenomyomas	Gonadotropin/hCG-producing tumors
Gliomas	Exogenous exposure to estrogen or androgen
Germinomas	McCune-Albright syndrome
CNS infection	Aromatase excess syndrome
Head trauma	
Iatrogenic	
Radiation	
Chemotherapy	
Surgical	
CNS malformation	
Arachnoid or suprasellar cysts	
Septo-optic dysplasia	
Hydrocephalus	

Abbreviations: CNS, central nervous system; GnRH, gonadotropin-releasing hormone; hCG, human chorionic gonadotropin.

DISORDERS OF PUBERTY

The differential diagnosis of precocious and delayed puberty is similar in boys (Chap. 411) and girls. However, there are differences in the timing of normal puberty and differences in the relative frequency of specific disorders in girls compared with boys.

Precocious Puberty Traditionally, precocious puberty has been defined as the development of secondary sexual characteristics before the age of 8 in girls based on data from Marshall and Tanner in British girls studied in the 1960s. More recent studies led to recommendations that girls be evaluated for precocious puberty if breast development or pubic hair is present at <7 years of age for white girls or <6 years for black girls.

Precocious puberty in girls is most often centrally mediated (Table 412-2), resulting from early activation of the hypothalamic-pituitary-ovarian axis. It is characterized by pulsatile LH secretion (which is initially associated with deep sleep) and an enhanced LH and FSH response to exogenous GnRH (two- to threefold stimulation) (Table 412-3). True precocity is marked by advancement in bone age of >2 standard deviations, a recent history of growth acceleration, and progression of secondary sexual characteristics. In girls, centrally mediated precocious puberty is idiopathic in ~85% of cases; however, neurogenic causes must be considered. Mutations in genes associated with GnRH deficiency have been reported in small numbers of patients with idiopathic precocious puberty (*KISS*, *KISS1R*, *TAC3*, *TAC3R*, and *DAX-1*), but their frequency is insufficient to warrant their use in clinical testing. GnRH agonists that induce pituitary desensitization are the mainstay of treatment to prevent premature epiphyseal closure and preserve adult height, as well as to manage psychosocial repercussions of precocious puberty.

Peripherally mediated precocious puberty does not involve activation of the hypothalamic-pituitary-ovarian axis and is characterized by suppressed gonadotropins in the presence of elevated estradiol. Management of peripheral precocious puberty involves treating the underlying disorder (Table 412-2) and limiting the effects of gonadal steroids using aromatase inhibitors, inhibitors of steroidogenesis, and ER blockers. It is important to be aware that central precocious puberty can also develop in girls whose precocity was initially peripherally mediated, as in McCune-Albright syndrome and congenital adrenal hyperplasia.

Incomplete and intermittent forms of precocious puberty may also occur. For example, premature breast development may occur in girls before the age of 2 years, with no further progression and without significant advancement in bone age, estrogen production, or compromised

TABLE 412-3 EVALUATION OF PRECOCIOUS AND DELAYED PUBERTY

	Precocious	Delayed
Initial Screening Tests		
History and physical	×	×
Assessment of growth velocity	×	×
Bone age	×	×
LH, FSH	×	×
Estradiol, testosterone	×	×
DHEAS	×	×
17-Hydroxyprogesterone	×	×
TSH, T ₄	×	×
Complete blood count		×
Sedimentation rate, C-reactive protein		×
Electrolytes, renal function		×
Liver enzymes		×
IGF-I, IGFBP-3		×
Urinalysis		×
Secondary Tests		
Pelvic ultrasound	×	×
Cranial MRI	×	×
β-hCG	×	
GnRH/agonist stimulation test	×	×
ACTH stimulation test	×	
Inflammatory bowel disease panel	×	×
Celiac disease panel		×
Prolactin		×
Karyotype		×

Abbreviations: ACTH, adrenocorticotropic hormone; DHEAS, dehydroepiandrosterone sulfate; FSH, follicle-stimulating hormone; hCG, human chorionic gonadotropin; IGF-I, insulin-like growth factor-I; IGFBP-3, IGF-binding protein 3; LH, luteinizing hormone; MRI, magnetic resonance imaging; TSH, thyroid-stimulating hormone; T₄, thyroxine.

height. Premature adrenarche can also occur in the absence of progressive pubertal development, but it must be distinguished from late-onset congenital adrenal hyperplasia and androgen-secreting tumors, in which case it may be termed *heterosexual precocity*. Premature adrenarche may be associated with obesity, hyperinsulinemia, and the subsequent predisposition to PCOS.

Delayed Puberty Delayed puberty (Table 412-4) is defined as the absence of secondary sexual characteristics by age 13 in girls. The diagnostic considerations are very similar to those for primary amenorrhea (Chap. 69). Between 25 and 40% of delayed puberty in girls is of ovarian origin, with Turner's syndrome accounting for the majority of such patients. Functional hypogonadotropic hypogonadism encompasses diverse etiologies such as systemic illnesses, including celiac disease and chronic renal disease, and endocrinopathies such as diabetes and hypothyroidism. In addition, girls appear to be particularly susceptible to the adverse effects of decreased energy balance resulting from exercise, dieting, and/or eating disorders. Together these reversible conditions account for ~25% of delayed puberty in girls. Congenital hypogonadotropic hypogonadism in girls or boys can be caused by mutations in several different genes or combinations of genes (Fig. 412-4, Chap. 411, Table 411-2). Approximately 50% of girls with congenital hypogonadotropic hypogonadism, with or without anosmia, have a history of some degree of breast development, and 10% report one to two episodes of vaginal bleeding. Family studies suggest that genes identified in association with absent puberty may also cause delayed puberty, and recent reports have further suggested that a genetic susceptibility to environmental stresses such as diet and exercise may account for at least some cases of functional hypothalamic amenorrhea. Although neuroanatomic causes of delayed puberty are considerably less common in girls than in boys, it is always important to rule these out in the setting of hypogonadotropic hypogonadism.

TABLE 412-4 DIFFERENTIAL DIAGNOSIS OF DELAYED PUBERTY**Hypergonadotropic**

Ovarian

Turner's syndrome
Gonadal dysgenesis
Chemotherapy/radiation therapy
Galactosemia
Autoimmune oophoritis
Congenital lipoid hyperplasia

Steroidogenic enzyme abnormalities

17 α -Hydroxylase deficiency
Aromatase deficiency

Gonadotropin/receptor mutations

FSH β , *LHR*, *FSHR*

Androgen resistance syndrome

Hypogonadotropic

Genetic

Hypothalamic syndromes

Leptin/leptin receptor
HESX1 (septo-optic dysplasia)
PC1 (prohormone convertase)

IHH and Kallmann's syndrome

KAL1, *FGF8*, *FGFR1*, *NSMF*, *PROK2*, *PROKR2*,
KISS1, *KISS1R*, *TAC3*, *TAC3R*, *GnRH1*, *GnRHR*, *SEM3A*, *HS6ST1*, *WDR11*, *CHD7*

Abnormalities of pituitary development/function

PROP1

CNS tumors/infiltrative disorders

Craniopharyngioma
Astrocytoma, germinoma, glioma
Prolactinomas, other pituitary tumors
Histiocytosis X

Chemotherapy/radiation

Functional

Chronic diseases
Malnutrition
Excessive exercise
Eating disorders

Abbreviations: *CHD7*, chromodomain-helicase-DNA-binding protein 7; CNS, central nervous system; *FGF8*, fibroblast growth factor 8; *FGFR1*, fibroblast growth factor 1 receptor; *FSH β* , follicle-stimulating hormone β chain; *FSHR*, FSH receptor; *GnRHR*, gonadotropin-releasing hormone receptor; *HESX1*, homeobox, embryonic stem cell expressed 1; *HS6ST1*, heparin sulfate 6-O sulfotransferase 1; IHH, idiopathic hypogonadotropic hypogonadism; *KAL1*, Kallmann; *KISS1*, kisspeptin 1; *KISS1R*, *KISS1* receptor; *LHR*, luteinizing hormone receptor; *NSMF*, NMDA receptor synaptonuclear signaling and neuronal migration factor; *PROK2*, prokineticin 2; *PROKR2*, prokineticin receptor 2; *PROP1*, prophet of Pit1, paired-like homeodomain transcription factor *SEMA3A*, semaphorin-3A; *WDR11*, WD repeat-containing protein 11.

413 Menopause and Postmenopausal Hormone Therapy

JoAnn E. Manson, Shari S. Bassuk

Menopause is the permanent cessation of menstruation due to loss of ovarian follicular function. It is diagnosed retrospectively after 12 months of amenorrhea. The average age at menopause is 51 years among U.S. women. *Perimenopause* refers to the time period preceding menopause, when fertility wanes and menstrual cycle irregularity increases, until the first year after cessation of menses. The onset of

perimenopause precedes the final menses by 2–8 years, with a mean duration of 4 years. Smoking accelerates the menopausal transition by 2 years.

Although the peri- and postmenopausal transitions share many symptoms, the physiology and clinical management of the two differ. Low-dose oral contraceptives have become a therapeutic mainstay in perimenopause, whereas postmenopausal hormone therapy (HT) has been a common method of symptom alleviation after menstruation ceases.

PERIMENOPAUSE**PHYSIOLOGY**

Ovarian mass and fertility decline sharply after age 35 and even more precipitously during perimenopause; depletion of primary follicles, a process that begins before birth, occurs steadily until menopause (Chap. 412). In perimenopause, intermenstrual intervals shorten significantly (typically by 3 days) as a result of an accelerated follicular phase. Follicle-stimulating hormone (FSH) levels rise because of altered folliculogenesis and reduced inhibin secretion. In contrast to the consistently high FSH and low estradiol levels seen in menopause, perimenopause is characterized by “irregularly irregular” hormone levels. The propensity for anovulatory cycles can produce a hyperestrogenic, hypoprogesteragenic environment that may account for the increased incidence of endometrial hyperplasia or carcinoma, uterine polyps, and leiomyoma observed among women of perimenopausal age. Mean serum levels of selected ovarian and pituitary hormones during the menopausal transition are shown in Fig. 413-1. With transition into menopause, estradiol levels fall markedly, whereas estrone levels are relatively preserved, a pattern reflecting peripheral aromatization of adrenal and ovarian androgens. Levels of FSH increase more than those of luteinizing hormone, presumably because of the loss of inhibin as well as estrogen feedback.

DIAGNOSTIC TESTS

The Stages of Reproductive Aging Workshop +10 (STRAW+10) classification provides a comprehensive framework for the clinical assessment of ovarian aging. As shown in Fig. 413-2, menstrual cycle characteristics are the principal criteria for characterizing the menopausal transition, with biomarker measures as supportive criteria. Because of their extreme intraindividual variability, FSH and estradiol levels are imperfect diagnostic indicators of perimenopause in menstruating women. However, a consistently low FSH level in the early follicular phase (days 2–5) of the menstrual cycle does not support a diagnosis of perimenopause, while levels >25 IU/L in a random blood sample are characteristic of the late menopause transition. FSH measurement can also aid in assessing fertility; levels of <20 IU/L, 20 to <30 IU/L, and \geq 30 IU/L measured on day 3 of the cycle indicate a good, fair, and poor likelihood of achieving pregnancy, respectively. Antimüllerian hormone and inhibin B may also be useful for assessing reproductive aging.

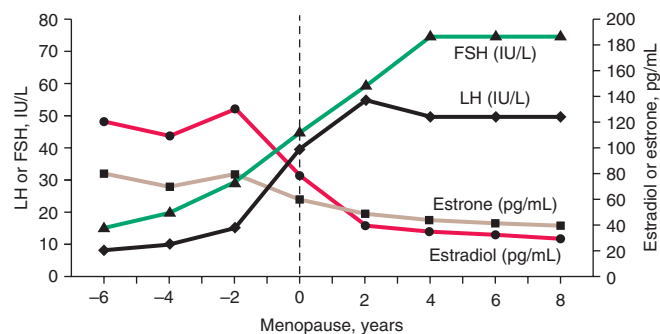


FIGURE 413-1 Mean serum levels of ovarian and pituitary hormones during the menopausal transition. FSH, follicle-stimulating hormone; LH, luteinizing hormone. (From JL Shifren, I Schiff: *J Womens Health Gen Based Med* 9 Suppl 1:S3, 2000.)

Stage	-5	-4	-3b	-3a	-2	-1	+1a	+1b	+1c	+2
Terminology	Reproductive				Menopausal transition		Postmenopause			
	Early	Peak	Late		Early	Late	Early			Late
					Perimenopause					
Duration	Variable				Variable	1–3 years	2 years (1+1)	3–6 years	Remaining lifespan	
Principal criteria										
Menstrual cycle	Variable to regular	Regular	Regular	Subtle changes in flow/length	Variable Length Persistent ≥7-day difference in length of consecutive cycles	Interval of amenorrhea of ≥60 days				
Supportive criteria										
Endocrine FSH AMH Inhibin B			Low Low	Variable* Low Low	↑ Variable* Low Low	↑ >25 IU/L** Low Low	↑ Variable Low Low	Stabilizes Very low Very low		
Antral follicle count			Low	Low	Low	Low	Very low	Very low		
Descriptive characteristics										
Symptoms							Vasomotor symptoms <i>Likely</i>	Vasomotor symptoms <i>Most likely</i>		Increasing symptoms of urogenital atrophy

*Blood draw on cycle days 2–5 ↑ = elevated.

**Approximate expected level based on assays using current international pituitary standard.

FIGURE 413-2 The Stages of Reproductive Aging Workshop +10 (STRAW +10) staging system for reproductive aging in women. AMH, antimüllerian hormone; FSH, follicle-stimulating hormone. (From SD Harlow et al: *Menopause* 14:387, 2012. Reproduced with permission.)

SYMPTOMS

Determining whether symptoms that develop in midlife are due to ovarian senescence or to other age-related changes is difficult. There is strong evidence that the menopausal transition can cause hot flashes, night sweats, irregular bleeding, and vaginal dryness, and there is moderate evidence that it can cause sleep disturbances in some women. There is inconclusive or insufficient evidence that ovarian aging is a major cause of mood swings, depression, impaired memory or concentration, somatic symptoms, urinary incontinence, or sexual dysfunction. In one U.S. study, nearly 60% of women reported hot flashes in the 2 years before their final menses. Symptom intensity, duration, frequency, and effects on quality of life are highly variable.

TREATMENT PERIMENOPAUSE

PERIMENOPAUSAL THERAPY

For women with irregular or heavy menses or hormone-related symptoms that impair quality of life, low-dose combined oral contraceptives are a staple of therapy. Static doses of estrogen and progestin (e.g., 20 µg of ethinyl estradiol and 1 mg of norethindrone acetate daily for 21 days each month) can eliminate vasomotor symptoms and restore regular cyclicality. Oral contraceptives provide other benefits, including protection against ovarian and endometrial cancers and increased bone density, although it is not clear whether use during perimenopause decreases fracture risk later in life. Moreover, the contraceptive benefit is important, given that the unintentional pregnancy rate among women in their forties rivals that of adolescents. Contraindications to oral contraceptive use include cigarette smoking, liver disease, a history of thromboembolism or cardiovascular disease, breast cancer, or unexplained vaginal bleeding. Progestin-only formulations (e.g., 0.35 mg of norethindrone daily) or medroxyprogesterone (Depo-Provera) injections (e.g., 150 mg IM every 3 months) may provide an alternative for the

treatment of perimenopausal menorrhagia in women who smoke or have cardiovascular risk factors. Although progestins neither regularize cycles nor reduce the number of bleeding days, they reduce the volume of menstrual flow.

Nonhormonal strategies to reduce menstrual flow include the use of nonsteroidal anti-inflammatory agents such as mefenamic acid (an initial dose of 500 mg at the start of menses, then 250 mg qid for 2–3 days) or, when medical approaches fail, endometrial ablation. It should be noted that menorrhagia requires an evaluation to rule out uterine disorders. Transvaginal ultrasound with saline enhancement is useful for detecting leiomyomata or polyps, and endometrial aspiration can identify hyperplastic changes.

TRANSITION TO MENOPAUSE

For sexually active women using contraceptive hormones to alleviate perimenopausal symptoms, the question of when and if to switch to HT must be individualized. Doses of estrogen and progestogen (either synthetic progestins or natural forms of progesterone) in HT are lower than those in oral contraceptives and have not been documented to prevent pregnancy. Although a 1-year absence of spontaneous menses reliably indicates ovulation cessation, it is not possible to assess the natural menstrual pattern while a woman is taking an oral contraceptive. Women willing to switch to a barrier method of contraception should do so; if menses occur spontaneously, oral contraceptive use can be resumed. The average age of final menses among relatives can serve as a guide for when to initiate this process, which can be repeated yearly until menopause has occurred.

MENOPAUSE AND POSTMENOPAUSAL HORMONE THERAPY

One of the most complex health care decisions facing women is whether to use postmenopausal HT. Once prescribed primarily to relieve vasomotor symptoms, HT has been promoted as a strategy to

forestall various disorders that accelerate after menopause, including osteoporosis and cardiovascular disease. In 2000, nearly 40% of postmenopausal women age 50–74 in the United States had used HT. This widespread use occurred despite the paucity of conclusive data, until recently, on the health consequences of such therapy. Although many women rely on their health care providers for a definitive answer to the question of whether to use postmenopausal hormones, balancing the benefits and risks for an individual patient is challenging.

Although observational studies suggest that HT prevents cardiovascular and other chronic diseases, the apparent benefits may result at least in part from differences between women who opt to take postmenopausal hormones and women who do not. Those choosing HT tend to be healthier, have greater access to medical care, are more compliant with prescribed treatments, and maintain a more health-promoting lifestyle. Randomized trials, which eliminate these confounding factors, have not consistently confirmed the benefits found in observational studies. Indeed, the largest HT trial to date, the Women's Health Initiative (WHI), which examined more than 27,000 postmenopausal women age 50–79 (mean age, 63) for an average of 5–7 years, was stopped early because of an overall unfavorable benefit-risk ratio in the estrogen-progestin arm and an excess risk of stroke that was not offset by a reduced risk of coronary heart disease (CHD) in the estrogen-only arm.

The following summary offers a decision-making guide based on a synthesis of currently available evidence. Prevention of cardiovascular disease is eliminated from the equation due to lack of evidence for such benefits in recent randomized clinical trials.

BENEFITS AND RISKS OF POSTMENOPAUSAL HORMONE THERAPY

See [Table 413-1](#).

Definite Benefits • SYMPTOMS OF MENOPAUSE Compelling evidence, including data from randomized clinical trials, indicates that estrogen therapy is highly effective for controlling vasomotor and genitourinary symptoms. Alternative approaches, including the use of antidepressants (such as paroxetine, 7.5 mg/d; or venlafaxine, 75–150 mg/d), gabapentin (300–900 mg/d), clonidine (0.1–0.2 mg/d), or vitamin E (400–800 IU/d), or the consumption of soy-based products or other phytoestrogens, may also alleviate vasomotor symptoms, although they are less effective than HT. Paroxetine is the only nonhormonal drug approved by the U.S. Food and Drug Administration for treatment of vasomotor symptoms. Bazedoxifene, an estrogen agonist/antagonist, in combination with conjugated estrogens has also received approval for vasomotor symptom management. For genitourinary symptoms, the efficacy of vaginal estrogen is similar to that of oral or transdermal estrogen; oral ospemifene is an additional option.

OSTEOPOROSIS (See also [Chap. 425](#))

Bone density By reducing bone turnover and resorption rates, estrogen slows the aging-related bone loss experienced by most postmenopausal women. More than 50 randomized trials have demonstrated that postmenopausal estrogen therapy, with or without a progestogen, rapidly increases bone mineral density at the spine by 4–6% and at the hip by 2–3% and that those increases are maintained during treatment.

Fractures Data from observational studies indicate a 50–80% lower risk of vertebral fracture and a 25–30% lower risk of hip, wrist, and other peripheral fractures among current estrogen users; addition of a progestogen does not appear to modify this benefit. In the WHI, 5–7 years of either combined estrogen-progestin or estrogen-only therapy was associated with a 33% reduction in hip fractures and 25–30% fewer total fractures among a population unselected for osteoporosis. Bisphosphonates (such as alendronate, 10 mg/d or 70 mg once per week; risedronate, 5 mg/d or 35 mg once per week; or ibandronate, 2.5 mg/d or 150 mg once per month or 3 mg every 3 months IV) and raloxifene (60 mg/d), a selective estrogen receptor modulator (SERM), have been shown in randomized trials to increase bone mass density and decrease fracture rates. Other options for treatment of osteoporosis are bazedoxifene in combination with conjugated estrogens and

parathyroid hormone (teriparatide, 20 µg/d SC). These agents, unlike estrogen, do not appear to have adverse effects on the endometrium or breast. Increased physical activity, adequate calcium intake (1000–1200 mg/d through diet or supplements in two or three divided doses), and adequate vitamin D intake (600–1000 IU/d) may also reduce the risk of osteoporosis-related fractures. According to the Institute of Medicine's 2011 report, 25-hydroxyvitamin D blood levels of ≥ 50 nmol/L are sufficient for bone-density maintenance and fracture prevention. The Fracture Risk Assessment (FRAX[®]) score, an algorithm that combines an individual's bone-density score with age and other risk factors to predict her 10-year risk of hip and major osteoporotic fracture, may be of use in guiding decisions about pharmacologic treatment (see [www.shef.ac.uk/FRAX/](#)).

Definite Risks • ENDOMETRIAL CANCER (WITH ESTROGEN ALONE) A combined analysis of 30 observational studies found a tripling of endometrial cancer risk among short-term users (1–5 years) of unopposed estrogen and a nearly tenfold increased risk among long-term users (≥ 10 years). These findings are supported by results from the randomized Postmenopausal Estrogen/Progestin Interventions (PEPI) trial, in which 24% of women assigned to unopposed estrogen for 3 years developed atypical endometrial hyperplasia—a premalignant lesion—as opposed to only 1% of women assigned to placebo. Use of a progestogen, which opposes the effects of estrogen on the endometrium, eliminates these risks and may even reduce risk (see later).

VENOUS THROMBOEMBOLISM A meta-analysis of observational studies found that current oral estrogen use was associated with a 2.5-fold increase in risk of venous thromboembolism in postmenopausal women. A meta-analysis of randomized trials, including the WHI, found a 2.1-fold increase in risk. Results from the WHI indicate a nearly twofold increase in risk of pulmonary embolism and deep vein thrombosis with estrogen-progestin and a 35–50% increase in these risks with estrogen-only therapy. Transdermal estrogen, taken alone or with certain progestogens (micronized progesterone or pregnane derivatives), appears to be a safer alternative with respect to thrombotic risk.

BREAST CANCER (WITH ESTROGEN-PROGESTIN) An increased risk of breast cancer has been found among current or recent estrogen users in observational studies; this risk is directly related to duration of use. In a meta-analysis of 51 case-control and cohort studies, short-term use (<5 years) of postmenopausal HT did not appreciably elevate breast cancer incidence, whereas long-term use (≥ 5 years) was associated with a 35% increase in risk. In contrast to findings for endometrial cancer, combined estrogen-progestin regimens appear to increase breast cancer risk more than estrogen alone. Data from randomized trials also indicate that estrogen-progestin raises breast cancer risk. In the WHI, women assigned to receive combination hormones for an average of 5.6 years were 24% more likely to develop breast cancer than women assigned to placebo, but 7.1 years of estrogen-only therapy did not increase risk. Indeed, the WHI showed a trend toward a reduction in breast cancer risk with estrogen alone, although it is unclear whether this finding would pertain to formulations of estrogen other than conjugated equine estrogens or to treatment durations of >7 years. In the Heart and Estrogen/Progestin Replacement Study (HERS), 4 years of combination therapy was associated with a 27% increase in breast cancer risk. Although the latter finding was not statistically significant, the totality of evidence strongly implicates estrogen-progestin therapy in breast carcinogenesis.

Some observational data suggest that the length of the interval between menopause onset and HT initiation may influence the association between such therapy and breast cancer risk, with a “gap time” of <3–5 years conferring a higher HT-associated breast cancer risk. (This pattern of findings contrasts with that for CHD, as discussed later in this Chapter.) However, this association remains inconclusive and may be a spurious finding attributable to higher rates of screening mammography and thus earlier cancer detection in HT users than in nonusers, especially in early menopause. Indeed, in the WHI trial, hazard ratios for HT and breast cancer risk did not differ among women

TABLE 413-1 BENEFITS AND RISKS OF POSTMENOPAUSAL HORMONE THERAPY IN THE OVERALL STUDY POPULATION OF WOMEN 50–79 YEARS OF AGE IN THE INTERVENTION PHASE OF THE WOMEN'S HEALTH INITIATIVE (WHI) ESTROGEN-PROGESTIN AND ESTROGEN-ALONE TRIALS^a

Outcome	Effect	Estrogen-Progestin		Estrogen Alone	
		Relative Benefit or Risk	Absolute Benefit or Risk ^b	Relative Benefit or Risk	Absolute Benefit or Risk ^b
Definite Benefits					
Symptoms of menopause	Definite improvement	↓65–90% decreased risk ^c		↓65–90% decreased risk ^c	
Osteoporosis	Definite increase in bone mineral density and decrease in fracture risk	↓33% decreased risk for hip fracture	6 fewer cases (11 vs. 17) of hip fracture	↓33% decreased risk for hip fracture	6 fewer cases (13 vs. 19) of hip fracture
Definite Risks^b					
Endometrial cancer	Definite increase in risk with estrogen alone (see below for estrogen-progestin)	See below	See below		4.6 excess cases (observational studies)
Pulmonary embolism	Definite increase in risk	↑98% increased risk	9 excess cases (18 vs. 9)	↑35% increased risk (n.s.)	4 excess cases (14 vs. 10)
Deep vein thrombosis	Definite increase in risk	↑87% increased risk	11.5 excess cases (25 vs. 14)	↑48% increased risk	7.5 excess cases (23 vs. 15)
Breast cancer	Definite increase in risk with long-term use (≥5 years) of estrogen-progestin	↑24% increased risk	8.5 excess cases (43 vs. 35)	↓21% decreased risk (n.s.)	7 fewer cases (28 vs. 35)
Gallbladder disease	Definite increase in risk	↑57% increased risk	47 excess cases (131 vs. 84)	↑55% increased risk	58 excess cases (164 vs. 106)
Probable or Uncertain Risks and Benefits^b					
Coronary heart disease ^d	Probable increase in risk among older women and women many years past menopause; possible decrease in risk or no effect in younger or recently menopausal women ^e	↑18% increased risk (n.s.)	6 excess cases (41 vs. 35)	No increase in risk	No difference in risk
Myocardial infarction	Significant interaction by age group for estrogen alone, with reduced risk in younger—but not older—women (<i>p</i> for trend by age = .02)	↑24% increased risk (n.s.)	6 excess cases (35 vs. 29)	No increase in risk ^e	No difference in risk ^e
Stroke	Probable increase in risk	↑37% increased risk	9 excess cases (33 vs. 24)	↑35% increased risk	11 excess cases (45 vs. 34)
Ovarian cancer	Probable increase in risk with long-term use (≥5 years)	↑41% increased risk (n.s.)	1 excess cases (5 vs. 4)	Not available	Not available
Endometrial cancer	Probable decrease in risk with estrogen-progestin during long-term follow-up (see above for estrogen alone)	↓33% decreased risk ^f	3 fewer cases (7 vs. 10)	See above	See above
Urinary incontinence	Probable increase in risk	↑49% increased risk	549 excess cases (1661 vs. 1112)	↑61% increased risk	852 excess cases (2255 vs. 1403)
Colorectal cancer	Probable decrease in risk with estrogen-progestin; possible increase in risk in older women with estrogen alone (<i>p</i> for trend by age = .02 for estrogen alone)	↓38% decreased risk	6.5 fewer cases (10 vs. 17)	No increase or decrease in risk ^e	No difference in risk ^e
Type 2 diabetes	Probable decrease in risk	↓19% decreased risk	16 fewer cases (72 vs. 88)	↓14% decreased risk	21 fewer cases (134 vs. 155)
Dementia (age ≥65)	Increase in risk in older women (but inconsistent data from observational studies and randomized trials)	↑101% increased risk	23 excess cases (46 vs. 23)	↑47% increased risk (n.s.)	15 excess cases (44 vs. 29)
Total mortality	Possible increase in risk among older women and women many years past menopause; possible decrease in risk or no effect in younger or recently menopausal women (<i>p</i> for trend by age <.05 for both trials combined)	No increase in risk	No difference in risk	No increase in risk ^e	No difference in risk ^e
Global index ^g	Probable increase in risk or no effect among older women and women many years past menopause; possible decrease in risk or no effect in younger or recently menopausal women (<i>p</i> for trend by age = 0.02 for estrogen alone)	↑12% increased risk	20.5 excess cases (189 vs. 168)	No increase in risk ^e	No difference in risk ^e

^aThe estrogen-progestin arm of the WHI assessed 5.6 years of conjugated equine estrogen (0.625 mg/d) plus medroxyprogesterone acetate (2.5 mg/d) versus placebo. The estrogen-alone arm of the WHI assessed 7.1 years of conjugated equine estrogen (0.625 mg/d) versus placebo. ^bNumber of cases per 10,000 women per year. ^cThe WHI was not designed to assess the effect of HT on menopausal symptoms. Data from other randomized trials suggest that HT reduces risk for menopausal symptoms by 65–90%. ^dCoronary heart disease is defined as nonfatal myocardial infarction or coronary death. ^eThere was a significant interaction by age; that is, the association between HT and the specified outcome was different in younger women and older women. ^fThis is the risk reduction that was observed during a cumulative 12-year follow-up period (5.6 years of treatment plus 6.8 years of postintervention observation). ^gThe global index is a composite outcome representing the first event for each participant from among the following: coronary heart disease, stroke, pulmonary embolism, breast cancer, colorectal cancer, endometrial cancer (estrogen-progestin arm only), hip fracture, and death. Because participants can experience more than one type of event, the global index cannot be derived by a simple summing of the component events. ^hIncludes some outcomes where results were divergent between the estrogen-progestin arm and the estrogen-alone arm.

Abbreviation: n.s., not statistically significant.

Source: Data from JE Manson et al: JAMA 310:1353, 2013.

50–59, those 60–69, and those 70–79 years of age at trial entry. (There was insufficient power to examine finer age categories.) Additional research is needed to clarify the issue.

GALLBLADDER DISEASE Large observational studies report a two- to threefold increased risk of gallstones or cholecystectomy among postmenopausal women taking oral estrogen. In the WHI, women randomized to estrogen-progestin or estrogen alone were ~55% more likely to develop gallbladder disease than those assigned to placebo. Risks were also increased in HERS. Transdermal HT might be a safer alternative, but further research is needed.

Probable or Uncertain Risks and Benefits • CORONARY HEART DISEASE/STROKE Until recently, HT had been enthusiastically recommended as a possible cardioprotective agent. In the past three decades, multiple observational studies suggested, in the aggregate, that estrogen use leads to a 35–50% reduction in CHD incidence among postmenopausal women. The biologic plausibility of such an association is supported by data from randomized trials demonstrating that exogenous estrogen lowers plasma low-density lipoprotein (LDL) cholesterol levels and raises high-density lipoprotein (HDL) cholesterol levels by 10–15%. Administration of estrogen also favorably affects lipoprotein(a) levels, LDL oxidation, endothelial vascular function, fibrinogen, and plasminogen activator inhibitor 1. However, estrogen therapy has unfavorable effects on other biomarkers of cardiovascular risk: it boosts triglyceride levels; promotes coagulation via factor VII, prothrombin fragments 1 and 2, and fibrinopeptide A elevations; and raises levels of the inflammatory marker C-reactive protein.

Randomized trials of estrogen or combined estrogen-progestin in women with preexisting cardiovascular disease have not confirmed the benefits reported in observational studies. In HERS (a secondary-prevention trial designed to test the efficacy and safety of estrogen-progestin therapy with regard to clinical cardiovascular outcomes), the 4-year incidence of coronary death and nonfatal myocardial infarction was similar in the active-treatment and placebo groups, and a 50% increase in risk of coronary events was noted during the first year among participants assigned to the active-treatment group. Although it is possible that progestin may mitigate estrogen's benefits, the Estrogen Replacement and Atherosclerosis (ERA) trial indicated that angiographically determined progression of coronary atherosclerosis was unaffected by either opposed or unopposed estrogen treatment. Moreover, no cardiovascular benefit was found in the Papworth Hormone Replacement Therapy Atherosclerosis Study, a trial of transdermal estradiol with and without norethindrone; the Women's Estrogen for Stroke Trial (WEST), a trial of oral 17 β -estradiol; or the Estrogen in the Prevention of Reinfarction Trial (ESPRIT), a trial of oral estradiol valerate. Thus, in clinical trials, HT has not proved effective for the secondary prevention of cardiovascular disease in postmenopausal women.

Primary-prevention trials also suggest an early increase in cardiovascular risk and an absence of cardioprotection with postmenopausal HT. In the WHI, women assigned to 5.6 years of estrogen-progestin therapy were 18% more likely to develop CHD (defined in primary analyses as nonfatal myocardial infarction or coronary death) than those assigned to placebo, although this risk elevation was not statistically significant. However, during the trial's first year, there was a significant 80% increase in risk, which diminished in subsequent years (p for trend by time = .03). In the estrogen-only arm of the WHI, no overall effect on CHD was observed during the 7.1 years of the trial or in any specific year of follow-up. This pattern of results was similar to that for the outcome of total myocardial infarction.

However, a closer look at available data suggests that timing of initiation of HT may critically influence the association between such therapy and CHD. Estrogen may slow early stages of atherosclerosis but have adverse effects on advanced atherosclerotic lesions. It has been hypothesized that the prothrombotic and proinflammatory effects of estrogen manifest themselves predominantly among women with subclinical lesions who initiate HT well after the menopausal transition, whereas women with less arterial damage who start HT early in menopause may derive cardiovascular benefit because they have not yet developed advanced lesions. Nonhuman primate data

support this concept. Conjugated estrogens had no effect on the extent of coronary artery plaque in cynomolgus monkeys assigned to receive estrogen alone or combined with progestin starting 2 years (~6 years in human terms) after oophorectomy and well after the establishment of atherosclerosis. However, administration of exogenous hormones immediately after oophorectomy, during the early stages of atherosclerosis, reduced the extent of plaque by 70%.

Lending further credence to this hypothesis are results of subgroup analyses of observational and clinical trial data. For example, among women who entered the WHI trial with a relatively favorable cholesterol profile, estrogen with or without progestin led to a 40% lower risk of incident CHD. Among women who entered with a worse cholesterol profile, therapy resulted in a 73% higher risk (p for interaction = .02). The presence or absence of the metabolic syndrome (Chap. 422) also strongly influenced the relation between HT and incident CHD. Among women with the metabolic syndrome, HT more than doubled CHD risk, whereas no association was observed among women without the syndrome. Moreover, although there was no association between estrogen-only therapy and CHD in the WHI trial cohort as a whole, such therapy was associated with a CHD risk reduction of 40% among participants age 50–59; in contrast, a risk reduction of only 5% was observed among those age 60–69, and a risk increase of 9% was found among those age 70–79 (p for trend by age = .08). For the outcome of total myocardial infarction, estrogen alone was associated with a borderline-significant 45% reduction and a nonsignificant 24% increase in risk among the youngest and oldest women, respectively (p for trend by age = .02). Estrogen was also associated with lower levels of coronary artery calcified plaque in the younger age group. Although age did not have a similar effect in the estrogen-progestin arm of the WHI, CHD risks increased with years since menopause (p for trend = .08), with a significantly elevated risk among women who were ≥ 20 years past menopause. For the outcome of total myocardial infarction, estrogen-progestin was associated with a 9% risk reduction among women <10 years past menopause as opposed to a 16% increase in risk among women 10–19 years past menopause and a twofold increase in risk among women >20 years past menopause (p for trend = .01). In the large observational Nurses' Health Study, women who chose to start HT within 4 years of menopause experienced a lower risk of CHD than did nonusers, whereas those who began therapy ≥ 10 years after menopause appeared to receive little coronary benefit. Observational studies include a high proportion of women who begin HT within 3–4 years of menopause, whereas clinical trials include a high proportion of women ≥ 12 years past menopause; this difference helps to reconcile some of the apparent discrepancies between the two types of studies.

For the outcome of stroke, WHI participants assigned to estrogen-progestin or estrogen alone were ~35% more likely to suffer a stroke than those assigned to placebo. Whether or not age at initiation of HT influences stroke risk is not well understood. In the WHI and the Nurses' Health Study, HT was associated with an excess risk of stroke in all age groups. Further research is needed on age, time since menopause, and other individual characteristics (including biomarkers) that predict increases or decreases in cardiovascular risk associated with exogenous HT. Furthermore, it remains uncertain whether different doses, formulations, or routes of administration of HT will produce different cardiovascular effects.

COLORECTAL CANCER Observational studies have suggested that HT reduces risks of colon and rectal cancer, although the estimated magnitudes of the relative benefits have ranged from 8% to 34% in various meta-analyses. In the WHI (the sole trial to examine the issue), estrogen-progestin was associated with a significant 38% reduction in colorectal cancer over a 5.6-year period, although no benefit was seen with 7 years of estrogen-only therapy. However, a modifying effect of age was observed, with a doubling of risk with HT in women age 70–79 but no risk elevation in younger women (p for trend by age = .02).

COGNITIVE DECLINE AND DEMENTIA A meta-analysis of 10 case-control and two cohort studies suggested that postmenopausal HT is associated with a 34% decreased risk of dementia. Subsequent randomized trials (including the WHI), however, have failed to demonstrate any benefit

2386 of estrogen or estrogen-progestin therapy on the progression of mild to moderate Alzheimer's disease and/or have indicated a potential adverse effect of HT on the incidence of dementia, at least in women ≥ 65 years of age. Among women randomized to HT (as opposed to placebo) at age 50–55 in the WHI, no effect on cognition was observed during the postintervention phase. Determining whether timing of initiation of HT influences cognitive outcomes will require further study.

OVARIAN CANCER AND OTHER DISORDERS On the basis of limited observational and randomized data, it has been hypothesized that HT increases the risk of ovarian cancer and reduces the risk of type 2 diabetes mellitus. Results from the WHI support these hypotheses. The WHI also found that HT use was associated with an increased risk of urinary incontinence and that estrogen-progestin was associated with increased rates of lung cancer mortality.

ENDOMETRIAL CANCER (WITH ESTROGEN-PROGESTIN) In the WHI, use of estrogen-progestin was associated with a nonsignificant 17% reduction in risk of endometrial cancer. A significant reduction in risk emerged during the postintervention period (see later).

ALL-CAUSE MORTALITY In the overall WHI cohort, estrogen with or without progestin was not associated with all-cause mortality. However, there was a trend toward reduced mortality in younger women, particularly with estrogen alone. For women 50–59, 60–69, and 70–79 years of age, relative risks (RRs) associated with estrogen-only therapy were 0.70, 1.01, and 1.21, respectively (p for trend = .04).

OVERALL BENEFIT-RISK PROFILE Estrogen-progestin was associated with an unfavorable benefit-risk profile (excluding relief from menopausal symptoms) as measured by a “global index”—a composite outcome including CHD, stroke, pulmonary embolism, breast cancer, colorectal cancer, endometrial cancer, hip fracture, and death (Table 413-1)—in the WHI cohort as a whole, and this association did not vary by 10-year age group. Estrogen-only therapy was associated with a neutral benefit-risk profile in the WHI cohort as a whole. However, there was a significant trend toward a more favorable benefit-risk profile among younger women and a less favorable profile among older women, with RRs of 0.84, 0.99, and 1.17 for women 50–59, 60–69, and 70–79 years of age, respectively (p for trend by age = .02).

CHANGES IN HEALTH STATUS AFTER DISCONTINUATION OF HORMONE THERAPY In the WHI, many but not all risks and benefits associated with active use of HT dissipated within 5–7 years after discontinuation of therapy. For estrogen-progestin, an elevated risk of breast cancer persisted (RR = 1.28 [95% confidence interval, 1.11–1.48]) during a cumulative 12-year follow-up period (5.6 years of treatment plus 6.8 years of postintervention observation), but most cardiovascular disease risks became neutral. A reduction in hip fracture risk persisted (RR = 0.81 [0.68–0.97]), and a significant reduction in endometrial cancer risk emerged (RR = 0.67 [0.49–0.91]). For estrogen alone, the reduction in breast cancer risk became statistically significant (RR = 0.79 [0.65–0.97]) during a cumulative 12-year follow-up period (6.8 years of treatment plus 5.1 years of postintervention observation), and significant differences by age group persisted for total myocardial infarction and the global index, with more favorable results for younger women.

APPROACH TO THE PATIENT: Postmenopausal Hormone Therapy

The rational use of postmenopausal HT requires balancing the potential benefits and risks. **Figure 413-3** provides one approach to decision making. The clinician should first determine whether the patient has moderate to severe menopausal symptoms—the primary indication for initiation of systemic HT. Systemic HT may also be used to prevent osteoporosis in women at high risk of fracture who cannot tolerate alternative osteoporosis therapies. (Vaginal estrogen or other medications may be used to treat urogenital symptoms in the absence of vasomotor symptoms.) The benefits and risks of such therapy should be reviewed with the patient, giving more emphasis to absolute than to relative measures

of effect and pointing out uncertainties in clinical knowledge where relevant. Because chronic disease rates generally increase with age, absolute risks tend to be greater in older women, even when relative risks remain similar. Potential side effects—especially vaginal bleeding that may result from use of the combined estrogen-progestogen formulations recommended for women with an intact uterus—should be noted. The patient's own preference regarding therapy should be elicited and factored into the decision. Contraindications to HT should be assessed routinely and include unexplained vaginal bleeding, active liver disease, venous thromboembolism, history of endometrial cancer (except stage 1 without deep invasion) or breast cancer, and history of CHD, stroke, transient ischemic attack, or diabetes. Relative contraindications include hypertriglyceridemia (>400 mg/dL) and active gallbladder disease; in such cases, transdermal estrogen may be an option. Primary prevention of heart disease should not be viewed as an expected benefit of HT, and an increase in the risk of stroke as well as a small early increase in the risk of coronary artery disease should be considered. Nevertheless, such therapy may be appropriate if the noncoronary benefits of treatment clearly outweigh the risks. A woman who suffers an acute coronary event or stroke while taking HT should discontinue therapy immediately.

Short-term use (<5 years for estrogen-progestogen and <7 years for estrogen alone) is appropriate for relief of menopausal symptoms among women without contraindications to such use. However, such therapy should be avoided by women with an elevated baseline risk of future cardiovascular events. Women who have contraindications for or are opposed to HT may derive benefit from the use of certain antidepressants (including venlafaxine, fluoxetine, or paroxetine), gabapentin, clonidine, soy, or black cohosh and, for genitourinary symptoms, intravaginal estrogen creams or devices, or ospemifene.

Long-term use (≥ 5 years for estrogen-progestogen and ≥ 7 years for estrogen alone) is more problematic because a heightened risk of breast cancer must be factored into the decision, especially for estrogen-progestogen. Reasonable candidates for such use include the small percentage of postmenopausal women who have persistent severe vasomotor symptoms along with an increased risk of osteoporosis (e.g., those with osteopenia, a personal or family history of nontraumatic fracture, or a weight below 125 lbs), who also have no personal or family history of breast cancer in a first-degree relative or other contraindications, and who have a strong personal preference for therapy. Poor candidates are women with elevated cardiovascular risk, those at increased risk of breast cancer (e.g., women who have a first-degree relative with breast cancer, susceptibility genes such as *BRCA1* or *BRCA2*, or a personal history of cellular atypia detected by breast biopsy), and those at low risk of osteoporosis. Even for reasonable candidates, strategies to minimize dose and duration of use should be employed. For example, women using HT to relieve intense vasomotor symptoms in early postmenopause should consider discontinuing therapy within 5 years, resuming it only if such symptoms persist. Because of the role of progestogens in increasing breast cancer risk, regimens that employ cyclic rather than continuous progestogen exposure as well as formulations other than medroxyprogesterone acetate should be considered if treatment is extended. For prevention of osteoporosis, alternative therapies such as bisphosphonates or SERMs should be considered. Research on alternative progestogens and androgen-containing preparations has been limited, particularly with respect to long-term safety. Additional research on the effects of these agents on cardiovascular disease, glucose tolerance, and breast cancer will be of particular interest.

In addition to HT, lifestyle choices such as smoking abstinence, adequate physical activity, and a healthy diet can play a role in controlling symptoms and preventing chronic disease. An expanding array of pharmacologic options (e.g., bisphosphonates, SERMs, and other agents for osteoporosis; cholesterol-lowering or antihypertensive agents for cardiovascular disease) should also reduce the widespread reliance on hormone use. However, short-term HT may still benefit some women.

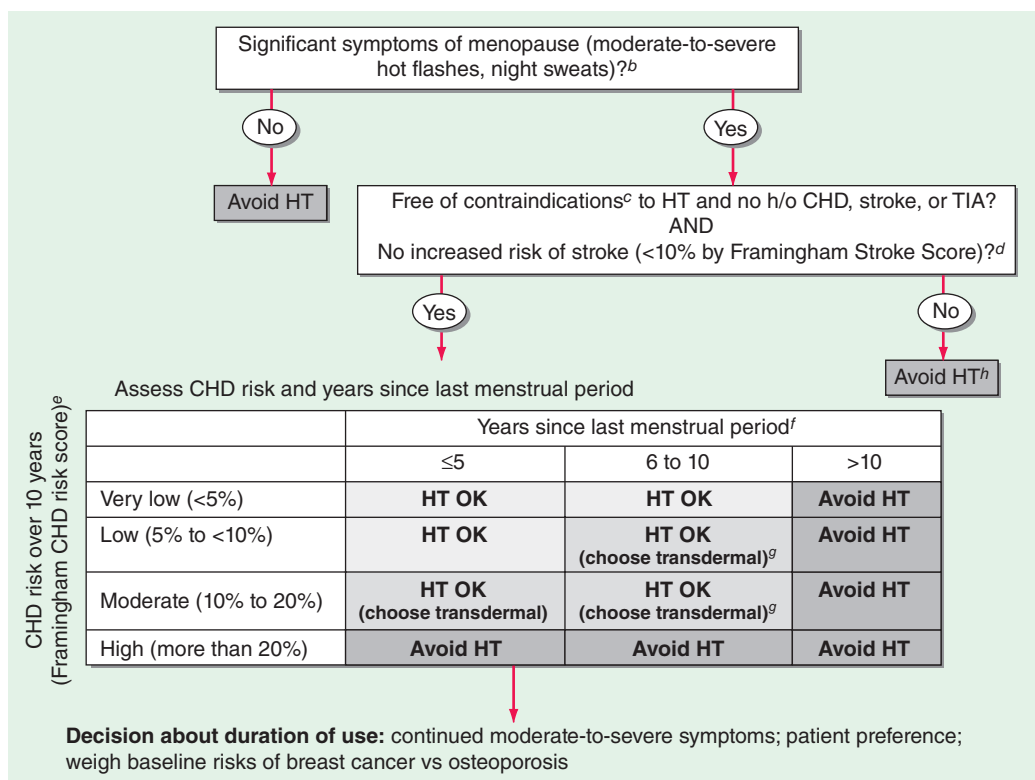


FIGURE 413-3 Chart for identifying appropriate candidates for postmenopausal hormone therapy (HT).^a

^aReassess each step at least once every 6–12 months (assuming the patient's continued preference for HT). ^bWomen who are at high risk of osteoporotic fracture but are unable to tolerate alternative preventive medications may also be reasonable candidates for systemic HT even if they do not have moderate to severe vasomotor symptoms. Women who have vaginal dryness without moderate to severe vasomotor symptoms may be candidates for vaginal estrogen. ^cTraditional contraindications are unexplained vaginal bleeding; active liver disease; history of venous thromboembolism due to pregnancy, oral contraceptive use, or an unknown etiology; blood-clotting disorder; history of breast or endometrial cancer; and diabetes. Oral HT should be avoided but transdermal HT may be an option (see *g* below) for other contraindications, including high triglyceride levels (>400 mg/dL); active gallbladder disease; and history of venous thromboembolism due to past immobility, surgery, or bone fracture. ^dTen-year risk of stroke, based on Framingham Stroke Risk Score (RB D'Agostino et al: Stroke risk profile: Adjustment for antihypertensive medication. The Framingham Study. Stroke 25:40, 1994), as modified by JE Manson, SS Bassuk: *Hot Flashes, Hormones & Your Health*. New York, McGraw-Hill, 2007. ^eTen-year risk of CHD, based on Framingham Coronary Heart Disease Risk Score (Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults: JAMA 285:2486, 2001), as modified by JE Manson, SS Bassuk: *Hot Flashes, Hormones & Your Health*. New York, McGraw-Hill, 2007. ^fWomen >10 years past menopause are not good candidates for initiation (first use) of HT. ^gAvoid oral HT. Transdermal HT may be an option because it has a less adverse effect on clotting factors, triglyceride levels, and inflammation factors than oral HT. ^hConsider selective serotonin or serotonin–norepinephrine reuptake inhibitor, gabapentin, clonidine, soy, or another alternative.

Abbreviations: CHD, coronary heart disease; h/o, history of; TIA, transient ischemic attack. (Adapted from JE Manson, SS Bassuk: *Hot Flashes, Hormones & Your Health*. New York, McGraw-Hill, 2007. Copyright © 2007 by the President and Fellows of Harvard College. All rights reserved.)

414 Infertility and Contraception

Janet E. Hall

INFERTILITY

DEFINITION AND PREVALENCE

Infertility has traditionally been defined as the inability to conceive after 12 months of unprotected sexual intercourse. In women who ultimately conceived, pregnancy occurred in ~50% within 3 months, 75–82% within 6 months, and 85–92% within 12 months. The World Health Organization (WHO) considers infertility as a disability (an impairment of function) and thus access to health care falls under the Convention on the Rights of Persons with Disability. Thirty-four million women, predominantly from developing countries, have infertility resulting from maternal sepsis and unsafe abortion. In populations <60 years old, infertility is ranked the fifth highest serious global disability. In the United States, the rate of infertility in married women age 15–44 is 6% based on the National Survey of Family Growth,

although prospective studies suggest that it may be as high as 12–15%. The infertility rate has remained relatively stable over the past 30 years in most countries. However, the proportion of couples without children has risen, reflecting both higher numbers of couples in childbearing years and a trend to delay childbearing. This trend has important implications because of an age-related decrease in fecundability: the incidence of primary infertility increases from ~8% between the ages of 18 and 38 to 25% and 30% between the ages of 35 and 39 and 40 and 44, respectively. It is estimated that 14% of couples in the United States have received medical assistance for infertility; of these, two-thirds received counseling, ~12% underwent infertility testing of the female and/or male partner, and 17% received drugs to induce ovulation.

CAUSES OF INFERTILITY

The spectrum of infertility ranges from reduced conception rates or the need for medical intervention to irreversible causes of infertility. Infertility can be attributed primarily to male factors in 25% of couples and female factors in 58% of couples and is unexplained in about 17% of couples (Fig. 414-1). Not uncommonly, both male and female factors contribute to infertility. Decreases in the ability to conceive as a

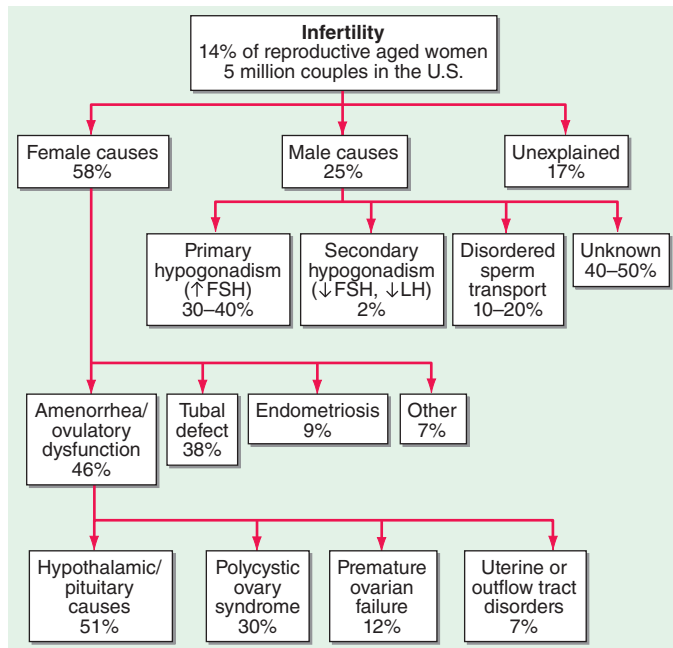


FIGURE 414-1 Causes of infertility. FSH, follicle-stimulating hormone; LH, luteinizing hormone.

function of age in women has led to recommendations that women >34 years old who are not at increased risk of infertility seek attention after 6 months, rather than 12 months as suggested for younger women, and receive an expedited work-up and approach to treatment.

APPROACH TO THE PATIENT: Infertility

INITIAL EVALUATION

In all couples presenting with infertility, the initial evaluation includes discussion of the appropriate timing of intercourse and discussion of modifiable risk factors such as smoking, alcohol, caffeine, and obesity. The range of required investigations should be reviewed as well as a brief description of infertility treatment options, including adoption. Initial investigations are focused on determining whether the primary cause of the infertility is male, female, or both. These investigations include a semen analysis in the male, confirmation of ovulation in the female, and, in the majority of situations, documentation of tubal patency in the female. In some cases, after an extensive workup excluding all male and female factors, a specific cause cannot be identified, and infertility may ultimately be classified as unexplained.

PSYCHOLOGICAL ASPECTS OF INFERTILITY

Infertility is invariably associated with psychological stress related not only to the diagnostic and therapeutic procedures themselves but also to repeated cycles of hope and loss associated with each new procedure or cycle of treatment that does not result in the birth of a child. These feelings are often combined with a sense of isolation from friends and family. Counseling and stress-management techniques should be introduced early in the evaluation of infertility. Importantly, infertility and its treatment do not appear to be associated with long-term psychological sequelae.

FEMALE CAUSES

Abnormalities in menstrual function constitute the most common cause of female infertility. These disorders, which include ovulatory dysfunction and abnormalities of the uterus or outflow tract, may present as amenorrhea or as irregular or short menstrual cycles. A careful history and physical examination and a limited number of laboratory tests will help to determine whether the abnormality is

(1) hypothalamic or pituitary (low follicle-stimulating hormone [FSH], luteinizing hormone [LH], and estradiol with or without an increase in prolactin), (2) polycystic ovary syndrome (PCOS; irregular cycles and hyperandrogenism in the absence of other causes of androgen excess), (3) ovarian (low estradiol with increased FSH), or (4) a uterine or outflow tract abnormality. The frequency of these diagnoses depends on whether the amenorrhea is primary or occurs after normal puberty and menarche (see Fig. 69-2).

The approach to further evaluation of these disorders is described in detail in Chap. 69.

Ovulatory Dysfunction In women with a history of regular menstrual cycles, evidence of ovulation should be sought (Chap. 412). Even in the presence of ovulatory cycles, evaluation of ovarian reserve is recommended for women age >35 years if they are interested in fertility. Measurement of FSH on day 3 of the cycle (an FSH level <10 IU/mL on cycle day 3 predicts adequate ovarian oocyte reserve) is the most cost-effective test. Other tests include measurement of FSH in response to clomiphene citrate (blocks estrogen negative feedback on FSH), antral follicle count on ultrasound, and anti-müllerian hormone (AMH); <0.5 ng/mL predicts reduced ovarian reserve although there is variability between labs).

Tubal Disease Tubal dysfunction may result from pelvic inflammatory disease (PID), appendicitis, endometriosis, pelvic adhesions, tubal surgery, previous use of an intrauterine device (IUD), and a previous ectopic pregnancy. However, a cause is not identified in up to 50% of patients with documented tubal factor infertility. Because of the high prevalence of tubal disease, evaluation of tubal patency by hysterosalpingogram (HSG) or laparoscopy should occur early in the majority of couples with infertility. Subclinical infections with *Chlamydia trachomatis* may be an underdiagnosed cause of tubal infertility and requires the treatment of both partners.

Endometriosis Endometriosis is defined as the presence of endometrial glands or stroma outside the endometrial cavity and uterine musculature and accounts for 40% of infertility not due to ovulatory disorders, tubal obstruction, or male factor. Its presence is suggested by a history of dyspareunia (painful intercourse), worsening dysmenorrhea that often begins before menses, or a thickened rectovaginal septum or deviation of the cervix on pelvic examination. Mild endometriosis does not appear to impair fertility; the pathogenesis of the infertility associated with moderate and severe endometriosis may be multifactorial with impairments of folliculogenesis, fertilization, and implantation, as well as adhesions. Endometriosis is often clinically silent, however, and can only be excluded definitively by laparoscopy.

MALE CAUSES (SEE ALSO CHAP. 411)

Known causes of male infertility include primary testicular disease, genetic disorders (particularly Y chromosome microdeletions), disorders of sperm transport, and hypothalamic-pituitary disease resulting in secondary hypogonadism. However, the etiology is not ascertained in up to one-half of men with suspected male factor infertility. The key initial diagnostic test is a semen analysis. Testosterone levels should be measured if the sperm count is low on repeated examination or if there is clinical evidence of hypogonadism. Gonadotropin levels will help to determine a gonadal versus a central cause of hypogonadism.

TREATMENT INFERTILITY

In addition to addressing the negative impact of smoking on fertility and pregnancy outcome, counseling about nutrition and weight is a fundamental component of infertility and pregnancy management. Both low and increased body mass index (BMI) are associated with infertility in women and with increased morbidity during pregnancy. Obesity has also been associated with infertility in men. The treatment of infertility should be tailored to the problems unique to each couple. In many situations, including unexplained infertility,

mild-to-moderate endometriosis, and/or borderline semen parameters, a stepwise approach to infertility is optimal, beginning with low-risk interventions and moving to more invasive, higher risk interventions only if necessary. After determination of all infertility factors and their correction, if possible, this approach might include, in increasing order of complexity: (1) expectant management, (2) clomiphene citrate or an aromatase inhibitor (see below) with or without intrauterine insemination (IUI), (3) gonadotropins with or without IUI, and (4) in vitro fertilization (IVF). The time used for evaluation, correction of problems identified, and expectant management can be longer in women age <30 years, but this process should be advanced rapidly in women age >35 years. In some situations, expectant management will not be appropriate.

OVULATORY DYSFUNCTION

Treatment of ovulatory dysfunction should first be directed at identification of the etiology of the disorder to allow specific management when possible. Dopamine agonists, for example, may be indicated in patients with hyperprolactinemia (Chap. 403); lifestyle modification may be successful in women with obesity, low body weight, or a history of intensive exercise.

Medications used for ovulation induction include agents that increase FSH through alteration of negative feedback, gonadotropins, and pulsatile GnRH. *Clomiphene citrate* is a nonsteroidal estrogen antagonist that increases FSH and LH levels by blocking estrogen negative feedback at the hypothalamus. The efficacy of clomiphene for ovulation induction is highly dependent on patient selection. In appropriate patients, it induces ovulation in ~60% of women with PCOS and has traditionally been the initial treatment of choice. Combination with agents that modify insulin levels such as metformin does not appear to improve outcome. Clomiphene citrate is less successful in patients with hypogonadotropic hypogonadism. *Aromatase inhibitors* have also been investigated for the treatment of infertility. Studies suggest they may have advantages over clomiphene, but these medications have not been approved for this indication.

Gonadotropins are highly effective for ovulation induction in women with hypogonadotropic hypogonadism and PCOS and are used to induce the development of multiple follicles in unexplained infertility and in older reproductive-age women. Disadvantages include a significant risk of multiple gestation and the risk of ovarian hyperstimulation, particularly in women with polycystic ovaries, with or without other features of PCOS. Careful monitoring and a conservative approach to ovarian stimulation reduce these risks. Currently available gonadotropins include urinary preparations of LH and FSH, highly purified FSH, and recombinant FSH. Although FSH is the key component, LH is essential for steroidogenesis in hypogonadotropic patients, and LH or human chorionic gonadotropin (hCG) may improve results through effects on terminal differentiation of the oocyte. These methods are commonly combined with IUI.

None of these methods are effective in women with premature ovarian failure, in whom donor oocyte or adoption is the method of choice.

TUBAL DISEASE

If hysterosalpingography suggests a tubal or uterine cavity abnormality or if a patient is age ≥ 35 at the time of initial evaluation, laparoscopy with tubal lavage is recommended, often with a hysteroscopy. Although tubal reconstruction may be attempted if tubal disease is identified, it is generally being replaced by the use of IVF. These patients are at increased risk of developing an ectopic pregnancy.

ENDOMETRIOSIS

Although 60% of women with minimal or mild endometriosis may conceive within 1 year without treatment, laparoscopic resection or ablation appears to improve conception rates. Medical management of advanced stages of endometriosis is widely used for symptom control but has not been shown to enhance fertility. In moderate and severe endometriosis, conservative surgery is associated with

pregnancy rates of 50 and 39%, respectively, compared with rates of 25 and 5% with expectant management alone. In some patients, IVF may be the treatment of choice.

MALE FACTOR INFERTILITY

The treatment options for male factor infertility have expanded greatly in recent years (Chap. 411). Secondary hypogonadism is highly amenable to treatment with gonadotropins or pulsatile gonadotropin-releasing hormone (GnRH) where available. In vitro techniques have provided new opportunities for patients with primary testicular failure and disorders of sperm transport. Choice of initial treatment options depends on sperm concentration and motility. Expectant management should be attempted initially in men with mild male factor infertility (sperm count of 15 to 20×10^6 /mL and normal motility). Moderate male factor infertility (10 to 15×10^6 /mL and 20–40% motility) should begin with IUI alone or in combination with treatment of the female partner with ovulation induction, but it may require IVF with or without intracytoplasmic sperm injection (ICSI). For men with a severe defect (sperm count of $<10 \times 10^6$ /mL, 10% motility), IVF with ICSI or donor sperm should be used. If ICSI is performed because of azoospermia due to congenital bilateral absence of the vas deferens, genetic testing and counseling should be provided because of the risk of cystic fibrosis.

ASSISTED REPRODUCTIVE TECHNOLOGIES

The development of assisted reproductive technologies (ARTs) has dramatically altered the treatment of male and female infertility. IVF is indicated for patients with many causes of infertility that have not been successfully managed with more conservative approaches. IVF or ICSI is often the treatment of choice in couples with a significant male factor or tubal disease, whereas IVF using donor oocytes is used in patients with premature ovarian failure and in women of advanced reproductive age. Success rates are influenced by cause of infertility and age, varying between 15 and 40%. Success rates are highest in anovulatory women and lowest in women with decreased ovarian reserve. In the United States, success rates are higher in white than in black, Asian, or Hispanic women. Although often effective, IVF is expensive and requires careful monitoring of ovulation induction and invasive techniques, including the aspiration of multiple follicles. IVF is associated with a significant risk of multiple gestation, particularly in women age <35, in whom the rate can be as high as 30%, which has led to specific recommendations for numbers of embryos or blastocysts to transfer based on age and specific prognostic factors.

CONTRACEPTION

Although use of contraception worldwide has increased in the last two decades, as of 2010, 146 million women worldwide age 15–49 years who were married or in a union had an unmet need for family planning. The absolute number of married women who use contraception or have an unmet need for family planning is projected to grow from 900 million (876–922 million) in 2010 to 962 million (927–992 million) in 2015.

Only 15% of couples in the United States report having unprotected sexual intercourse in the past 3 months. However, despite the wide availability and widespread use of a variety of effective methods of contraception, approximately one-half of all births in the United States are the result of unintended pregnancy. Teenage pregnancies continue to represent a serious public health problem in the United States, with >1 million unintended pregnancies each year—a significantly greater incidence than in other industrialized nations.

Of the contraceptive methods available (Table 414-1), a reversible form of contraception is used by >50% of couples, whereas sterilization (male or female) has been used as a permanent form of contraception by over one-third of couples. Pregnancy termination is relatively safe when directed by health care professionals but is rarely the option of choice.

TABLE 414-1 EFFECTIVENESS OF DIFFERENT FORMS OF CONTRACEPTION

Method of Contraception	Theoretical ^a Effectiveness, %	Actual ^a Effectiveness, %	Percent Continuing Use at 1 Year ^b	Contraceptive Methods Used by U.S. Women ^c
Barrier methods				
Condoms	98	88	63	18
Diaphragm	94	82	58	2
Cervical cap	94	82	50	<1
Spermicides	97	79	43	1
Sterilization				
Male	99.9	99.9	100	9
Female	99.8	99.6	100	27
Intrauterine device				
Copper T380	99	97	78	1
Progestasert	98	97	81	
Mirena	99.9	99.8		
Hormonal contraceptives				
Combination pill	99.7	92	72	31
Progestin only pill				
Transdermal patch				
Vaginal ring				
Monthly injection				
Long-acting progestins				

^aAdapted from J Trussel et al: *Obstet Gynecol* 76:558, 1990. ^bAdapted from *Contraceptive Technology Update*. *Contraceptive Technology*, Feb. 1996, Vol 17, No 1, pp 13–24. ^cAdapted from LJ Piccinino, WD Mosher: *Fam Plan Perspective* 30:4, 1998.

No single contraceptive method is ideal, although all are safer than carrying a pregnancy to term. The effectiveness of a given method of contraception does not just depend on the efficacy of the method itself. Discrepancies between theoretical and actual effectiveness emphasize the importance of patient education and compliance when considering various forms of contraception (Table 414-1). Knowledge of the advantages and disadvantages of each contraceptive is essential for counseling an individual about the methods that are safest and most consistent with his or her lifestyle. The WHO has extensive family planning resources for the physician and patient that can be accessed online. Similar resources for determining medical eligibility are available through the Centers for Disease Control and Prevention (CDC). Considerations for contraceptive use in obese patients and after bariatric surgery are discussed below.

BARRIER METHODS

Barrier contraceptives (such as condoms, diaphragms, and cervical caps) and spermicides are easily available, reversible, and have fewer side effects than hormonal methods. However, their effectiveness is highly dependent on adherence and proper use (Table 414-1). A major advantage of barrier contraceptives is the protection provided against sexually transmitted infections (STIs) (Chap. 163). Consistent use is associated with a decreased risk of HIV, gonorrhea, nongonococcal urethritis, and genital herpes, probably due in part to the concomitant use of spermicides. Natural membrane condoms may be less effective than latex condoms, and petroleum-based lubricants can degrade condoms and decrease their efficacy for preventing HIV infection. Barrier methods used by women include the diaphragm, cervical cap, and contraceptive sponge. The cervical cap and sponge are less effective than the diaphragm, and there have been rare reports of toxic shock syndrome with the diaphragm and contraceptive sponge.

STERILIZATION

Sterilization is the method of birth control most frequently chosen by fertile men and multiparous women >30 years old (Table 414-1). Sterilization refers to a procedure that prevents fertilization by

surgical interruption of the fallopian tubes in women or the vas deferens in men. Although tubal ligation and vasectomy are potentially reversible, these procedures should be considered permanent and should not be undertaken without patient counseling.

Several methods of *tubal ligation* have been developed, all of which are highly effective with a 10-year cumulative pregnancy rate of 1.85 per 100 women. However, when pregnancy does occur, the risk of ectopic pregnancy may be as high as 30%. The success rate of tubal reanastomosis depends on the method of ligation used, but even after successful reversal, the risk of ectopic pregnancy remains high. In addition to prevention of pregnancy, tubal ligation reduces the risk of ovarian cancer, possibly by limiting the upward migration of potential carcinogens.

Vasectomy is a highly effective outpatient surgical procedure that has little risk. The development of azoospermia may be delayed for 2–6 months, and other forms of contraception must be used until two sperm-free ejaculations provide proof of sterility. Reanastomosis may restore fertility in 30–50% of men, but the success rate declines with time after vasectomy and may be influenced by factors such as the development of antisperm antibodies.

INTRAUTERINE DEVICES

IUDs inhibit pregnancy through several mechanisms, primarily via a spermicidal effect caused by a sterile inflammatory reaction induced by the presence of a foreign body in the uterine cavity (copper IUDs) or by the release of progestins (Progestasert, Mirena). IUDs provide a high level of efficacy in the absence of systemic metabolic effects, and ongoing motivation is not required to ensure efficacy once the device has been placed. However, only 1% of women in the United States use this method compared to a utilization rate of 15–30% in much of Europe and Canada, despite evidence that the newer devices are not associated with increased rates of pelvic infection and infertility, as occurred with earlier devices. An IUD should not be used in women at high risk for development of STI or in women at high risk for bacterial endocarditis. The IUD may not be effective in women with uterine leiomyomas because they alter the size or shape of the uterine cavity. IUD use is associated with increased menstrual blood flow, although this is less pronounced with the progestin-releasing IUD, which is associated with a more frequent occurrence of spotting or amenorrhea.

HORMONAL METHODS

Oral Contraceptive Pills Because of their ease of use and efficacy, oral contraceptive pills are the most widely used form of hormonal contraception. They act by suppressing ovulation, changing cervical mucus, and altering the endometrium. The current formulations are made from synthetic estrogens and progestins. The estrogen component of the pill consists of ethinyl estradiol or mestranol, which is metabolized to ethinyl estradiol. Multiple synthetic progestins are used. Norethindrone and its derivatives are used in many formulations. Low-dose norgestimate and the more recently developed (third-generation) progestins (desogestrel, gestodene, drospirenone) have a less androgenic profile; levonorgestrel appears to be the most androgenic of the progestins and should be avoided in patients with hyperandrogenism. The three major formulations of oral contraceptives are (1) fixed-dose estrogen-progestin combination, (2) phasic estrogen-progestin combination, and (3) progestin only. Each of these formulations is administered daily for 3 weeks followed by a week of

TABLE 414-2 ORAL CONTRACEPTIVES: CONTRAINDICATIONS AND DISEASE RISK

Contraindications	
Absolute	
	Previous thromboembolic event or stroke
	History of an estrogen-dependent tumor
	Active liver disease
	Pregnancy
	Undiagnosed abnormal uterine bleeding
	Hypertriglyceridemia
	Women age >35 years who smoke heavily
Relative	
	Hypertension
	Women receiving anticonvulsant drug therapy
	Women following bariatric surgery (malabsorptive procedure)
Disease Risks	
Increased	
	Coronary heart disease—increased in smokers >35; no relation to progestin type
	Hypertension—relative risk 1.8 (current users) and 1.2 (previous users)
	Venous thrombosis—relative risk ~4; may be higher with third-generation progestin, drospirenone, and patch; compounded by obesity (tenfold increased risk compared with nonobese, no OCP); markedly increased with factor V Leiden or prothrombin gene mutations
	Stroke—slight increase; unclear relation to migraine headache
	Cerebral vein thrombosis—relative risk ~13–15; synergistic with prothrombin gene mutation
	Cervical cancer—relative risk 2–4
	Breast cancer—may increase risk in carriers of <i>BRCA1</i> and possibly <i>BRCA2</i>
Decreased	
	Ovarian cancer—50% reduction in risk
	Endometrial cancer—40% reduction in risk

Abbreviation: OCP, oral contraceptive pill.

no medication during which menstrual bleeding generally occurs. Two extended oral contraceptives are approved for use in the United States; Seasonale is a 3-month preparation with 84 days of active drug and 7 days of placebo, whereas Lybrel is a continuous preparation. Current doses of ethinyl estradiol range from 10 to 50 µg. However, indications for the 50-µg dose are rare, and the majority of formulations contain 30–35 µg of ethinyl estradiol. The reduced estrogen and progestin content in the second- and third-generation pills has decreased both side effects and risks associated with oral contraceptive use (Table 414-2). At the currently used doses, patients must be cautioned not to miss pills due to the potential for ovulation. Side effects, including breakthrough bleeding, amenorrhea, breast tenderness, and weight gain, often respond to a change in formulation. Even the lower dose oral contraceptives have been associated with an increased risk of cardiovascular disease (myocardial infarction, stroke, venous thromboembolism [VTE]), but the absolute excess risk is extremely low. VTE risk is higher with the third-generation than the second-generation progestins, and the risk of stroke and VTE is also higher with drospirenone (although not cyproterone), but the absolute excess risk is small and may be outweighed by contraceptive benefits and reduction in ovarian and endometrial cancer risk.

The microdose progestin-only minipill is less effective as a contraceptive, having a pregnancy rate of 2–7 per 100 women-years. However, it may be appropriate for women at increased risk for cardiovascular disease or for women who cannot tolerate synthetic estrogens.

Alternative Methods A weekly contraceptive patch (Ortho Evra) is available and has similar efficacy to oral contraceptives. Approximately 2% of patches fail to adhere, and a similar percentage of women have skin reactions. Efficacy is lower in women weighing >90 kg. The amount of estrogen delivered may be comparable to that of a 40-µg ethinyl

estradiol oral contraceptive, raising the possibility of increased risk of VTE, which must be balanced against potential benefits for women not able to successfully use other methods. A monthly contraceptive estrogen/progestin injection (Lunelle) is highly effective, with a first-year failure rate of <0.2%, but it may be less effective in obese women. Its use is associated with bleeding irregularities that diminish over time. Fertility returns rapidly after discontinuation. A monthly vaginal ring (NuvaRing) that is intended to be left in place during intercourse is also available for contraceptive use. It is highly effective, with a 12-month failure rate of 0.7%. Ovulation returns within the first recovery cycle after discontinuation.

Long-Term Contraceptives Long-term progestin administration acts primarily by inhibiting ovulation and causing changes in the endometrium and cervical mucus that result in decreased implantation and sperm transport. Depot medroxyprogesterone acetate (Depo-Provera, DMPA), the only injectable form available in the United States, is effective for 3 months, but return of fertility after discontinuation may be delayed for up to 12–18 months. DMPA is now available for both SC and IM injection. Irregular bleeding, amenorrhea, and weight gain are the most common side effects. This form of contraception may be particularly good for women in whom an estrogen-containing contraceptive is contraindicated (e.g., migraine exacerbation, sickle cell anemia, fibroids).

POSTCOITAL CONTRACEPTION

The probability of pregnancy without relation to time of the month is 8%, but the probability varies significantly in relation to proximity to ovulation and may be as high as 30%. In order of efficacy, methods of postcoital contraception include the following:

1. Copper IUD insertion within a maximum of 5 days has a reported efficacy of 99–100% and prevents pregnancy by its spermicidal effect; insertion is frequently available through family planning clinics.
2. Oral antiprogesterins (ulipristal acetate, 30 mg single dose, available worldwide, or mifepristone, 600 mg single dose, not available for this indication in the United States) prevent pregnancy by delaying or preventing ovulation; when administered, ideally within 72 h but up to 120 h after intercourse, they have an efficacy of 98–99%; require a prescription.
3. Levonorgestrel (1.5 mg as a single dose) delays or prevents ovulation and is not effective after ovulation; should be taken within 72 h of unprotected intercourse, and has an efficacy that varies between 60 and 94%; it is available over the counter.

Combined estrogen and progestin regimens have lower efficacy and are no longer recommended. A pregnancy test is not necessary before the use of oral methods, but pregnancy should be excluded before IUD insertion. Risk factors for failure of oral regimens include close proximity to ovulation and unprotected intercourse after use. In addition, there is an increased risk of pregnancy in obese and overweight women using levonorgestrel for postcoital contraception and an increased risk in obese women using an antiprogesterin.

IMPACT OF OBESITY ON CONTRACEPTIVE CHOICE

Approximately one-third of adults in the United States are obese. Although obesity is associated with some reduction in fertility, the vast majority of obese women can conceive. The risk of pregnancy-associated complications is higher in obese women. Intrauterine contraception may be more effective than oral or transdermal methods for obese women. The WHO guidelines provide no restrictions (class 1) for the use of intrauterine contraception, DMPA, and progestin-only pills for obese women (BMI ≥30) in the absence of coexistent medical problems, whereas methods that include estrogen (pill, patch, ring) are considered class 2 (advantages generally outweigh theoretical or proven risks) due to the increased risk of thromboembolic disease. There are no restrictions to the use of any contraceptive methods following restrictive bariatric surgery procedures, but both combined and progestin-only pills are relatively less effective following procedures associated with malabsorption.

SECTION 3 OBESITY, DIABETES MELLITUS, AND METABOLIC SYNDROME

415e Biology of Obesity

Jeffrey S. Flier, Eleftheria Maratos-Flier

In a world where food supplies are intermittent, the ability to store energy in excess of what is required for immediate use is essential for survival. Fat cells, residing within widely distributed adipose tissue depots, are adapted to store excess energy efficiently as triglyceride and, when needed, to release stored energy as free fatty acids for use at other sites. This physiologic system, orchestrated through endocrine and neural pathways, permits humans to survive starvation for as long as several months. However, in the presence of nutritional abundance and a sedentary lifestyle, and influenced importantly by genetic endowment, this system increases adipose energy stores and produces adverse health consequences.

DEFINITION AND MEASUREMENT

Obesity is a state of excess adipose tissue mass. Although often viewed as equivalent to increased body weight, this need not be the case—lean but very muscular individuals may be overweight by numerical standards without having increased adiposity. Body weights are distributed continuously in populations, so that choice of a medically meaningful distinction between lean and obese is somewhat arbitrary. Obesity is therefore defined by assessing its linkage to morbidity or mortality.

Although not a direct measure of adiposity, the most widely used method to gauge obesity is the *body mass index* (BMI), which is equal to weight/height² (in kg/m²) (Fig. 415e-1). Other approaches to quantifying obesity include anthropometry (skin-fold thickness), densitometry (underwater weighing), computed tomography (CT) or magnetic resonance imaging (MRI), and electrical impedance. Using data from the Metropolitan Life Tables, BMIs for the mid-point of all heights and frames among both men and women range from 19 to 26 kg/m²; at a similar BMI, women have more body fat than men. Based on data of substantial morbidity, a BMI of 30 is most commonly used as a threshold for obesity in both men and women. Most but not all large-scale epidemiologic studies suggest that all-cause, metabolic, cancer, and cardiovascular morbidity begin to rise (albeit at a slow rate) when BMIs are ≥ 25 . Most authorities use the term *overweight* (rather than obese) to describe individuals with BMIs between 25 and 30. A BMI between 25 and 30 should be viewed as medically significant and worthy of therapeutic intervention in the presence of risk factors that are influenced by adiposity, such as hypertension and glucose intolerance.

The distribution of adipose tissue in different anatomic depots also has substantial implications for morbidity. Specifically,

intraabdominal and abdominal subcutaneous fat have more significance than subcutaneous fat present in the buttocks and lower extremities. This distinction is most easily made clinically by determining the waist-to-hip ratio, with a ratio >0.9 in women and >1.0 in men being abnormal. Many of the most important complications of obesity, such as insulin resistance, diabetes, hypertension, hyperlipidemia, and hyperandrogenism in women, are linked more strongly to intraabdominal and/or upper body fat than to overall adiposity (Chap. 422). The mechanism underlying this association is unknown but may relate to the fact that intraabdominal adipocytes are more lipolytically active than those from other depots. Release of free fatty acids into the portal circulation has adverse metabolic actions, especially on the liver. Adipokines and cytokines that are differentially secreted by adipocyte depots may play a role in the systemic complications of obesity.

PREVALENCE

Data from the National Health and Nutrition Examination Surveys (NHANES) show that the percentage of the American adult population

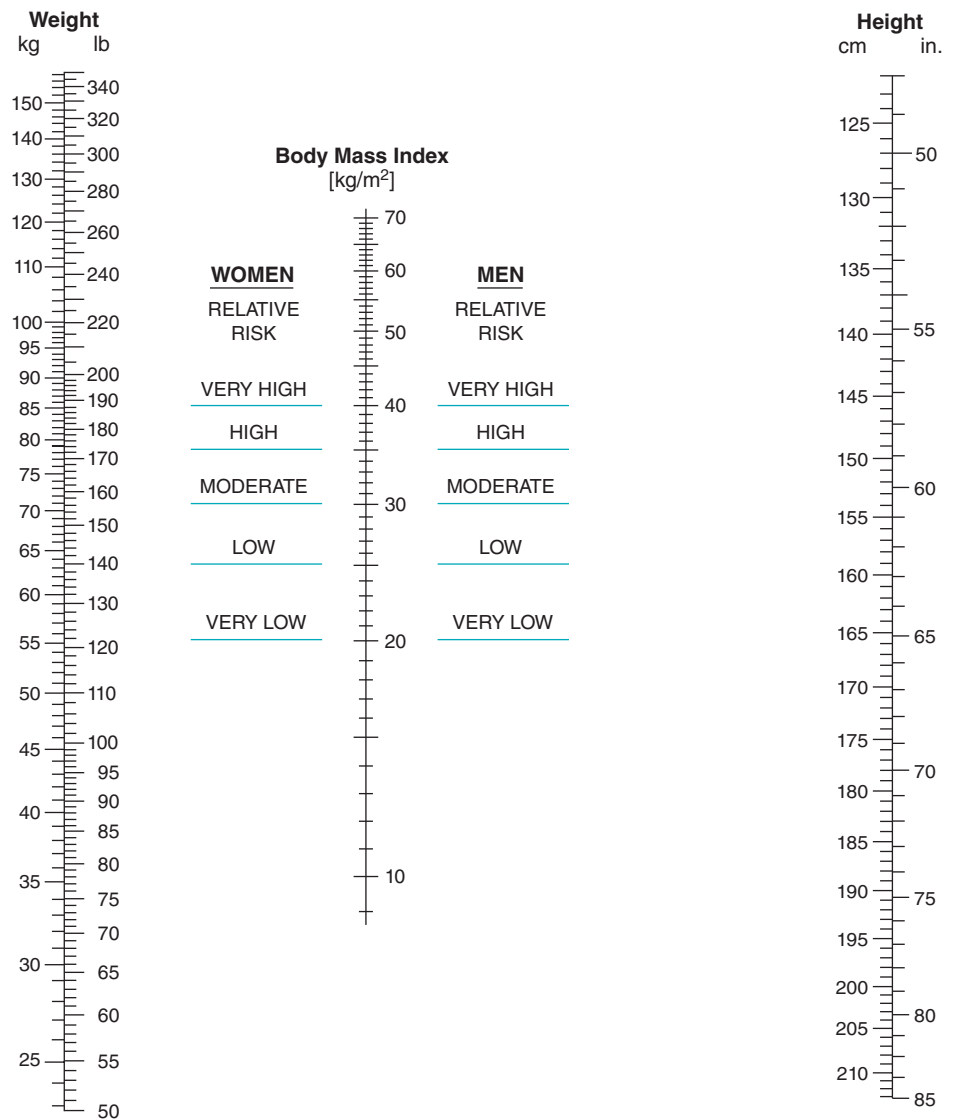


FIGURE 415e-1 Nomogram for determining body mass index. To use this nomogram, place a ruler or other straight edge between the body weight (without clothes) in kilograms or pounds located on the left-hand line and the height (without shoes) in centimeters or inches located on the right-hand line. The body mass index is read from the middle of the scale and is in metric units. (Copyright 1979, George A. Bray, MD; used with permission.)

415e-2 with obesity (BMI >30) has increased from 14.5% (between 1976 and 1980) to 35.7% (between 2009 and 2010). As many as 68% of U.S. adults aged ≥20 years were overweight (defined as BMI >25) between the years of 2007 and 2008. Extreme obesity (BMI ≥40) has also increased and affects 5.7% of the population. The increasing prevalence of medically significant obesity raises great concern. Overall, the prevalence of obesity is comparable in men and women. In women, poverty is associated with increased prevalence. Obesity is more common among blacks and Hispanics. The prevalence in children and adolescents has been rising at a worrisome rate, reaching 15.9% in 2009/2010, but may be leveling off.

PHYSIOLOGIC REGULATION OF ENERGY BALANCE

Substantial evidence suggests that body weight is regulated by both endocrine and neural components that ultimately influence the effector arms of energy intake and expenditure. This complex regulatory system is necessary because even small imbalances between energy intake and expenditure will ultimately have large effects on body weight. For example, a 0.3% positive imbalance over 30 years would result in a 9-kg (20-lb) weight gain. This exquisite regulation of energy balance cannot be monitored easily by calorie-counting in relation to physical activity. Rather, body weight regulation or dysregulation depends on a complex interplay of hormonal and neural signals. Alterations in stable weight by forced overfeeding or food deprivation induce physiologic changes that resist these perturbations: with weight loss, appetite increases and energy expenditure falls; with overfeeding, appetite falls and energy expenditure increases. This latter compensatory mechanism frequently fails, however, permitting obesity to develop when food is abundant and physical activity is limited. A major regulator of these adaptive responses is the adipocyte-derived hormone leptin, which acts through brain circuits (predominantly in the hypothalamus) to influence appetite, energy expenditure, and neuroendocrine function (see below).

Appetite is influenced by many factors that are integrated by the brain, most importantly within the hypothalamus (Fig. 415e-2). Signals that impinge on the hypothalamic center include neural afferents, hormones, and metabolites. Vagal inputs are particularly important, bringing information from viscera, such as gut distention. Hormonal signals include leptin, insulin, cortisol, and gut peptides. Among the latter is ghrelin, which is made in the stomach and stimulates feeding, and peptide YY (PYY) and cholecystokinin, which is made in the small intestine and signals to the brain through direct action on hypothalamic control centers and/or via the vagus nerve. Metabolites, including glucose, can influence appetite, as seen by the effect of hypoglycemia to induce hunger; however, glucose is not

normally a major regulator of appetite. These diverse hormonal, metabolic, and neural signals act by influencing the expression and release of various hypothalamic peptides (e.g., neuropeptide Y [NPY], Agouti-related peptide [AgRP], α-melanocyte-stimulating hormone [α-MSH], and melanin-concentrating hormone [MCH]) that are integrated with serotonergic, catecholaminergic, endocannabinoid, and opioid signaling pathways (see below). Psychological and cultural factors also play a role in the final expression of appetite. Apart from rare genetic syndromes involving leptin, its receptor, and the melanocortin system, specific defects in this complex appetite control network that influence common cases of obesity are not well defined.

Energy expenditure includes the following components: (1) resting or basal metabolic rate; (2) the energy cost of metabolizing and storing food; (3) the thermic effect of exercise; and (4) adaptive thermogenesis, which varies in response to long-term caloric intake (rising with increased intake). Basal metabolic rate accounts for ~70% of daily energy expenditure, whereas active physical activity contributes 5–10%. Thus, a significant component of daily energy consumption is fixed.

Genetic models in mice indicate that mutations in certain genes (e.g., targeted deletion of the insulin receptor in adipose tissue) protect against obesity, apparently by increasing energy expenditure. Adaptive thermogenesis occurs in *brown adipose tissue* (BAT), which plays an important role in energy metabolism in many mammals. In contrast to white adipose tissue, which is used to store energy in the form of lipids, BAT expends stored energy as heat. A mitochondrial *uncoupling protein* (UCP-1) in BAT dissipates the hydrogen ion gradient in the oxidative respiration chain and releases energy as heat. The metabolic activity of BAT is increased by a central action of leptin, acting through the sympathetic nervous system that heavily innervates this tissue. In rodents, BAT deficiency causes obesity and diabetes; stimulation of BAT with a specific adrenergic agonist (β₃ agonist) protects against diabetes and obesity. BAT exists in humans (especially neonates), and although its physiologic role is not yet established, identification of functional BAT in many adults using positron emission tomography (PET) imaging has increased interest in the implications of the tissue for pathogenesis and therapy of obesity. Beige fat cells, recently described, resemble BAT cells in expressing UCP-1. They are scattered through white adipose tissue, and their thermogenic potential is uncertain.

THE ADIPOCYTE AND ADIPOSE TISSUE

Adipose tissue is composed of the lipid-storing adipose cell and a stromal/vascular compartment in which cells including preadipocytes and macrophages reside. Adipose mass increases by enlargement of adipose cells through lipid deposition, as well as by an increase in the number of adipocytes. Obese adipose tissue is also characterized by increased numbers of infiltrating macrophages. The process by which adipose cells are derived from a mesenchymal preadipocyte involves an orchestrated series of differentiation steps mediated by a cascade of specific transcription factors. One of the key transcription factors is *peroxisome proliferator-activated receptor γ* (PPARγ), a nuclear receptor that binds the thiazolidinedione class of insulin-sensitizing drugs used in the treatment of type 2 diabetes (Chap. 418).

Although the adipocyte has generally been regarded as a storage depot for fat, it is also an endocrine cell that releases numerous molecules in a regulated fashion (Fig. 415e-3). These include the energy balance-regulating hormone leptin, cytokines such as tumor necrosis factor (TNF)-α and interleukin (IL)-6, complement factors such as factor D (also known as *adipsin*), prothrombotic agents such as plasminogen activator inhibitor I, and a component of the blood pressure-regulating system, angiotensinogen. Adiponectin, an abundant adipose-derived protein whose levels are reduced in obesity, enhances insulin sensitivity and lipid oxidation and has vascular-protective effects, whereas resistin and RBP4, whose levels are increased in obesity, may induce insulin resistance. These factors, and others not yet identified, play a role in the physiology of lipid homeostasis, insulin sensitivity, blood pressure control, coagulation, and vascular health, and are likely to contribute to obesity-related pathologies.

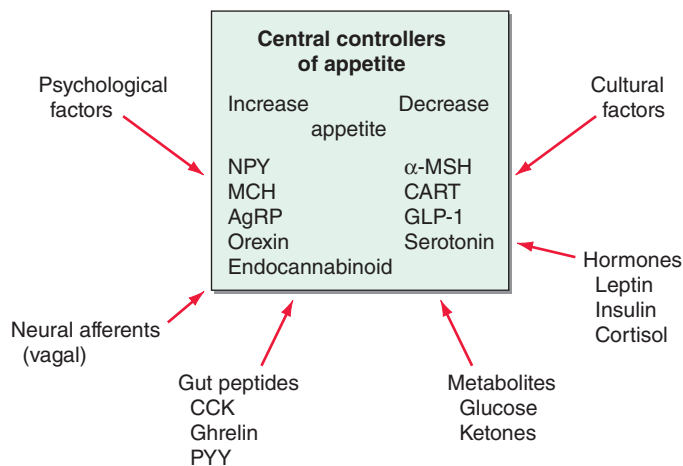


FIGURE 415e-2 The factors that regulate appetite through effects on central neural circuits. Some factors that increase or decrease appetite are listed. AgRP, Agouti-related peptide; CART, cocaine- and amphetamine-related transcript; CCK, cholecystokinin; GLP-1, glucagon-related peptide-1; MCH, melanin-concentrating hormone; α-MSH, α-melanocyte-stimulating hormone; NPY, neuropeptide Y.

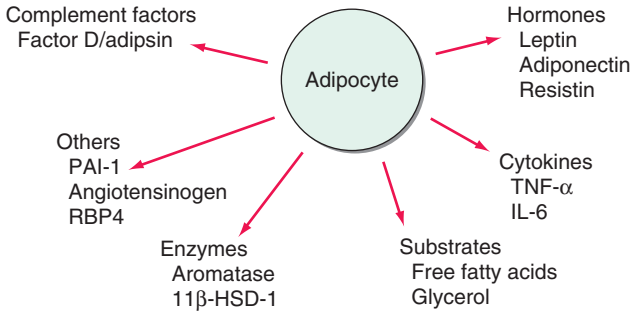


FIGURE 415e-3 Factors released by the adipocyte that can affect peripheral tissues. IL-6, interleukin 6; PAI, plasminogen activator inhibitor; RBP4, retinal binding protein 4; TNF, tumor necrosis factor.

ETIOLOGY OF OBESITY

Although the molecular pathways regulating energy balance are beginning to be illuminated, the causes of obesity remain elusive. In part, this reflects the fact that obesity is a heterogeneous group of disorders. At one level, the pathophysiology of obesity seems simple: a chronic excess of nutrient intake relative to the level of energy expenditure. However, due to the complexity of the neuroendocrine and metabolic systems that regulate energy intake, storage, and expenditure, it has been difficult to quantitate all the relevant parameters (e.g., food intake and energy expenditure) over time in human subjects.

Role of Genes Versus Environment Obesity is commonly seen in families, and the heritability of body weight is similar to that for height. Inheritance is usually not Mendelian, however, and it is difficult to distinguish the role of genes and environmental factors. Adoptees more closely resemble their biologic than adoptive parents with respect to obesity, providing strong support for genetic influences. Likewise, identical twins have very similar BMIs whether reared together or apart, and their BMIs are much more strongly correlated than those of dizygotic twins. These genetic effects appear to relate to both energy intake and expenditure. Currently, identified genetic variants, both common and rare, account for less than 5% of the variance of body weight.

Whatever the role of genes, it is clear that the environment plays a key role in obesity, as evidenced by the fact that famine prevents obesity in even the most obesity-prone individual. In addition, the recent increase in the prevalence of obesity in the United States is far too rapid to be due to changes in the gene pool. Undoubtedly, genes influence the susceptibility to obesity in response to specific diets and availability of nutrition. Cultural factors are also important—these relate to both availability and composition of the diet and to changes in the level of physical activity. In industrial societies, obesity is more common among poor women, whereas in underdeveloped countries, wealthier women are more often obese. In children, obesity correlates to some degree with time spent watching television. Although the role of diet composition in obesity continues to generate controversy, it appears that high-fat diets may, when combined with simple, rapidly absorbed carbohydrates, promote obesity. Specific genes are likely to influence the response to specific diets, but these genes are largely unidentified.

Additional environmental factors may contribute to the increasing obesity prevalence. Both epidemiologic correlations and experimental data suggest that sleep deprivation leads to increased obesity. Changes in gut microbiome with capacity to alter energy balance are receiving experimental support from animal studies, and a possible role for obesogenic viral infections continues to receive sporadic attention.

Specific Genetic Syndromes For many years, obesity in rodents has been known to be caused by a number of distinct mutations distributed through the genome. Most of these single-gene mutations cause both hyperphagia and diminished energy expenditure, suggesting a physiologic link between these two parameters of energy homeostasis. Identification of the *ob* gene mutation in genetically obese (*ob/ob*) mice represented a major breakthrough in the field. The *ob/ob* mouse

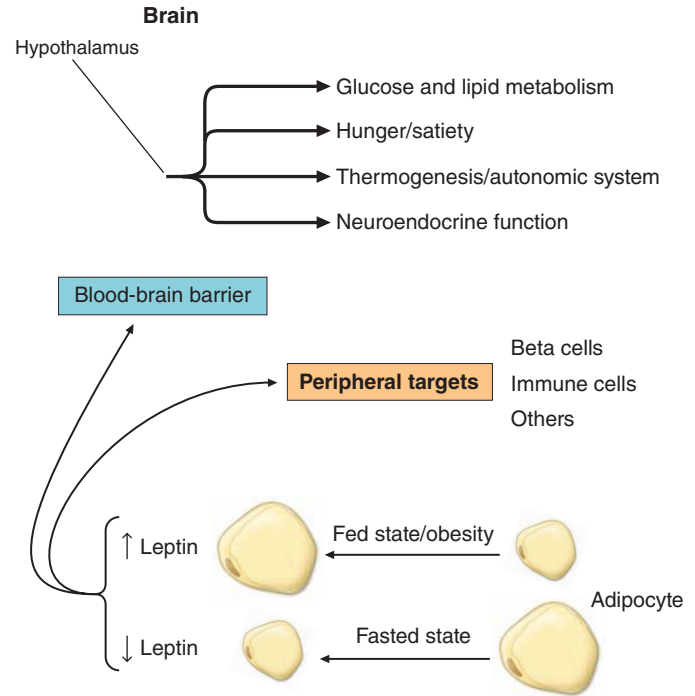


FIGURE 415e-4 The physiologic system regulated by leptin. Rising or falling leptin levels act through the hypothalamus to influence appetite, energy expenditure, and neuroendocrine function and through peripheral sites to influence systems such as the immune system.

develops severe obesity, insulin resistance, and hyperphagia, as well as efficient metabolism (e.g., it gets fat even when ingesting the same number of calories as lean litter mates). The product of the *ob* gene is the peptide leptin, a name derived from the Greek root *leptos*, meaning thin. Leptin is secreted by adipose cells and acts primarily through the hypothalamus. Its level of production provides an index of adipose energy stores (Fig. 415e-4). High leptin levels decrease food intake and increase energy expenditure. Another mouse mutant, *db/db*, which is resistant to leptin, has a mutation in the leptin receptor and develops a similar syndrome. The *ob* gene is present in humans where it is also expressed in fat. Several families with morbid, early-onset obesity caused by inactivating mutations in either leptin or the leptin receptor have been described, thus demonstrating the biologic relevance of the leptin pathway in humans. Obesity in these individuals begins shortly after birth, is severe, and is accompanied by neuroendocrine abnormalities. The most prominent of these is hypogonadotropic hypogonadism, which is reversed by leptin replacement in the leptin-deficient subset. Central hypothyroidism and growth retardation are seen in the mouse model, but their occurrence in leptin-deficient humans is less clear. Mutations in the leptin or leptin receptor genes do not play a prominent role in common forms of obesity.

Mutations in several other genes cause severe obesity in humans (Table 415e-1); each of these syndromes is rare. Mutations in the gene encoding proopiomelanocortin (POMC) cause severe obesity through failure to synthesize α -MSH, a key neuropeptide that inhibits appetite in the hypothalamus. The absence of POMC also causes secondary adrenal insufficiency due to absence of adrenocorticotropic hormone (ACTH), as well as pale skin and red hair due to absence of α -MSH. Proenzyme convertase 1 (PC-1) mutations are thought to cause obesity by preventing synthesis of α -MSH from its precursor peptide, POMC. α -MSH binds to the type 4 melanocortin receptor (MC4R), a key hypothalamic receptor that inhibits eating. Heterozygous loss-of-function mutations of this receptor account for as much as 5% of severe obesity. Loss of function of MRAP2, a protein required for normal MC4R signaling, has been found in rare cases of severe obesity. These six genetic defects define a pathway through which leptin (by stimulating POMC and increasing α -MSH) restricts food intake and limits weight

TABLE 415e-1 SELECTED OBESITY GENES IN HUMANS AND MICE

Gene	Gene Product	Mechanism of Obesity	In Human	In Rodent
<i>Lep (ob)</i>	Leptin, a fat-derived hormone	Mutation prevents leptin from delivering satiety signal; brain perceives starvation	Yes	Yes
<i>LepR (db)</i>	Leptin receptor	Same as above	Yes	Yes
<i>POMC</i>	Proopiomelanocortin, a precursor of several hormones and neuropeptides	Mutation prevents synthesis of melanocyte-stimulating hormone (MSH), a satiety signal	Yes	Yes
<i>MC4R</i>	Type 4 receptor for MSH	Mutation prevents reception of satiety signal from MSH	Yes	Yes
<i>AgRP</i>	Agouti-related peptide, a neuropeptide expressed in the hypothalamus	Overexpression inhibits signal through <i>MC4R</i>	No	Yes
<i>PC-1</i>	Prohormone convertase 1, a processing enzyme	Mutation prevents synthesis of neuropeptide, probably MSH	Yes	No
<i>Fat</i>	Carboxypeptidase E, a processing enzyme	Same as above	No	Yes
<i>Tub</i>	Tub, a hypothalamic protein of unknown function	Hypothalamic dysfunction	No	Yes
<i>TrkB</i>	TrkB, a neurotrophin receptor	Hyperphagia due to uncharacterized hypothalamic defect	Yes	Yes

(Fig. 415e-5). The results of genomewide association studies to identify genetic loci responsible for obesity in the general population have so far been disappointing. More than 40 replicated loci linked to obesity have been identified, but together they account for less than 3% of interindividual variation in BMI. The most replicated of these is a gene named *FTO*, which is of unknown function, but like many of the other recently described candidates, is expressed in the brain. Because the heritability of obesity is estimated to be 40–70%, it is likely that many more loci remain to be identified. It is possible that epistatic interactions between causative loci or unknown gene-environment interactions explain the poor success at identifying causal loci.

In addition to these human obesity genes, studies in rodents reveal several other molecular candidates for hypothalamic mediators of human obesity or leanness. The *tub* gene encodes a hypothalamic peptide of unknown function; mutation of this gene causes late-onset obesity. The *fat* gene encodes carboxypeptidase E, a peptide-processing enzyme; mutation of this gene is thought to cause obesity by disrupting production of one or more neuropeptides. *AgRP* is coexpressed with NPY in arcuate nucleus neurons. *AgRP* antagonizes α -MSH action at *MC4* receptors, and its overexpression induces obesity. In contrast, a

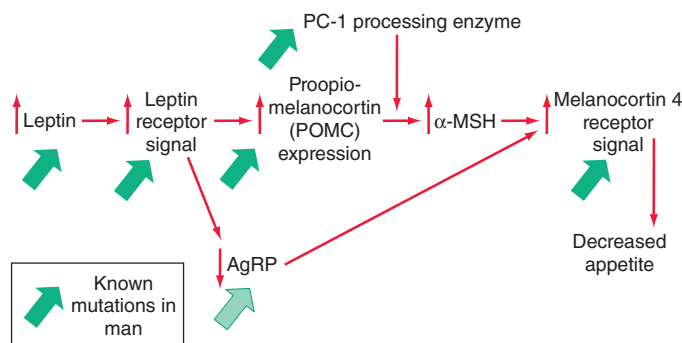


FIGURE 415e-5 A central pathway through which leptin acts to regulate appetite and body weight. Leptin signals through proopiomelanocortin (POMC) neurons in the hypothalamus to induce increased production of α -melanocyte-stimulating hormone (α -MSH), requiring the processing enzyme PC-1 (proenzyme convertase 1). α -MSH acts as an agonist on melanocortin-4 receptors to inhibit appetite, and the neuropeptide *AgRP* (Agouti-related peptide) acts as an antagonist of this receptor. Mutations that cause obesity in humans are indicated by the solid green arrows.

mouse deficient in the peptide MCH, whose administration causes feeding, is lean.

A number of complex human syndromes with defined inheritance are associated with obesity (Table 415e-2). Although specific genes have limited definition at present, their identification will likely enhance our understanding of more common forms of human obesity. In the Prader-Willi syndrome, a multigenic neurodevelopmental disorder, obesity coexists with short stature, mental retardation, hypogonadotropic hypogonadism, hypotonia, small hands and feet, fish-shaped mouth, and hyperphagia. Most patients have reduced expression of imprinted paternally inherited genes encoded in the 15q11-13 chromosomal region. Reduced expression of *Snord116*, a small nucleolar RNA highly expressed in hypothalamus, may be an important cause of defective hypothalamic function in this disorder. Bardet-Biedl syndrome (BBS) is a genetically heterogeneous disorder characterized by obesity, mental retardation, retinitis pigmentosa, diabetes, renal and cardiac malformations, polydactyly, and hypogonadotropic hypogonadism.

At least 12 genetic loci have been identified, and most of the encoded proteins form two multiprotein complexes that are involved in ciliary function and microtubule-based intracellular transport. Some evidence suggests that mutations might disrupt leptin receptor trafficking in key hypothalamic neurons, causing leptin resistance.

Other Specific Syndromes Associated with Obesity • CUSHING'S SYNDROME

Although obese patients commonly have central obesity, hypertension, and glucose intolerance, they lack other specific stigmata of Cushing's syndrome (Chap. 406). Nonetheless, a potential diagnosis of Cushing's syndrome is often entertained. Cortisol production and urinary metabolites (17OH steroids) may be increased in simple obesity. Unlike in Cushing's syndrome, however, cortisol levels in blood and urine in the basal state and in response to corticotropin-releasing hormone (CRH) or ACTH are normal; the overnight 1-mg dexamethasone suppression test is normal in 90%, with the remainder being normal on a standard 2-day low-dose dexamethasone suppression test. Obesity may be associated with excessive local reactivation of cortisol in fat by 11β -hydroxysteroid dehydrogenase 1, an enzyme that converts inactive cortisone to cortisol.

HYPOTHYROIDISM The possibility of hypothyroidism should be considered, but it is an uncommon cause of obesity; hypothyroidism is easily ruled out by measuring thyroid-stimulating hormone (TSH). Much of the weight gain that occurs in hypothyroidism is due to myxedema (Chap. 405).

INSULINOMA Patients with insulinoma often gain weight as a result of overeating to avoid hypoglycemic symptoms (Chap. 420). The increased substrate plus high insulin levels promote energy storage in fat. This can be marked in some individuals but is modest in most.

CRANIOPHARYNGIOMA AND OTHER DISORDERS INVOLVING THE HYPOTHALAMUS Whether through tumors, trauma, or inflammation, hypothalamic dysfunction of systems controlling satiety, hunger, and energy expenditure can cause varying degrees of obesity (Chap. 402). It is uncommon to identify a discrete anatomic basis for these disorders. Subtle hypothalamic dysfunction is probably a more common cause of obesity than can be documented using currently available imaging techniques. Growth hormone (GH), which exerts lipolytic activity, is diminished in obesity and is increased with weight loss. Despite low GH levels, insulin-like growth factor (IGF) I (somatomedin) production is normal, suggesting that GH suppression may be a compensatory response to increased nutritional supply.

TABLE 415e-2 A COMPARISON OF SYNDROMES OF OBESITY—HYPOGONADISM AND MENTAL RETARDATION

Feature	Syndrome				
	Prader-Willi	Laurence-Moon-Biedl	Ahlstrom's	Cohen's	Carpenter's
Inheritance	Sporadic; two-thirds have defect	Autosomal recessive	Autosomal recessive	Probably autosomal recessive	Autosomal recessive
Stature	Short	Normal; infrequently short	Normal; infrequently short	Short or tall	Normal
Obesity	Generalized Moderate to severe Onset 1–3 years	Generalized Early onset, 1–2 years	Truncal Early onset, 2–5 years	Truncal Mid-childhood, age 5	Truncal, gluteal
Craniofacies	Narrow bifrontal diameter Almond-shaped eyes Strabismus V-shaped mouth High-arched palate	Not distinctive	Not distinctive	High nasal bridge Arched palate Open mouth Short philtrum	Acrocephaly Flat nasal bridge High-arched palate
Limbs	Small hands and feet Hypotonia	Polydactyly	No abnormalities	Hypotonia Narrow hands and feet	Polydactyly Syndactyly Genu valgum
Reproductive status	1° Hypogonadism	1° Hypogonadism	Hypogonadism in males but not in females	Normal gonadal function or hypogonadotropic hypogonadism	2° Hypogonadism
Other features	Enamel hypoplasia Hyperphagia Temper tantrums Nasal speech			Dysplastic ears Delayed puberty	
Mental retardation	Mild to moderate		Normal intelligence	Mild	Slight

Pathogenesis of Common Obesity Obesity can result from increased energy intake, decreased energy expenditure, or a combination of the two. Thus, identifying the etiology of obesity should involve measurements of both parameters. However, it is difficult to perform direct and accurate measurements of energy intake in free-living individuals; and the obese, in particular, often underreport intake. Measurements of chronic energy expenditure are possible using doubly labeled water or metabolic chamber/rooms. In subjects at stable weight and body composition, energy intake equals expenditure. Consequently, these techniques allow assessment of energy intake in free-living individuals. The level of energy expenditure differs in established obesity, during periods of weight gain or loss, and in the pre- or postobese state. Studies that fail to take note of this phenomenon are not easily interpreted.

There is continued interest in the concept of a body weight “set point.” This idea is supported by physiologic mechanisms centered around a sensing system in adipose tissue that reflects fat stores and a receptor, or “adipostat,” that is in the hypothalamic centers. When fat stores are depleted, the adipostat signal is low, and the hypothalamus responds by stimulating hunger and decreasing energy expenditure to conserve energy. Conversely, when fat stores are abundant, the signal is increased, and the hypothalamus responds by decreasing hunger and increasing energy expenditure. The recent discovery of the *ob* gene, and its product leptin, and the *db* gene, whose product is the leptin receptor, provides important elements of a molecular basis for this physiologic concept (see above).

What Is the Status of Food Intake in Obesity? (Do the Obese Eat More Than the Lean?) This question has stimulated much debate, due in part to the methodologic difficulties inherent in determining food intake. Many obese individuals believe that they eat small quantities of food, and this claim has often been supported by the results of food intake questionnaires. However, it is now established that average energy expenditure increases as individuals get more obese, due primarily to the fact that metabolically active lean tissue mass increases with obesity. Given the laws of thermodynamics, the obese person must therefore eat more than the average lean person to maintain their increased weight. It may be the case, however, that a subset of individuals who are predisposed to obesity have the capacity to become obese initially without an absolute increase in caloric consumption.

What Is the State of Energy Expenditure in Obesity? The average total daily energy expenditure is higher in obese than lean individuals when measured at stable weight. However, energy expenditure falls as weight is lost, due in part to loss of lean body mass and to decreased sympathetic nerve activity. When reduced to near-normal weight and maintained there for a while, (some) obese individuals have lower energy expenditure than (some) lean individuals. There is also a tendency for those who will develop obesity as infants or children to have lower resting energy expenditure rates than those who remain lean. The physiologic basis for variable rates of energy expenditure (at a given body weight and level of energy intake) is essentially unknown.

Another component of thermogenesis, called *nonexercise activity thermogenesis* (NEAT), has been linked to obesity. It is the thermogenesis that accompanies physical activities other than volitional exercise such as the activities of daily living, fidgeting, spontaneous muscle contraction, and maintaining posture. NEAT accounts for about two-thirds of the increased daily energy expenditure induced by overfeeding. The wide variation in fat storage seen in overfed individuals is predicted by the degree to which NEAT is induced. The molecular basis for NEAT and its regulation is unknown.

Leptin in Typical Obesity The vast majority of obese persons have increased leptin levels but do not have mutations of either leptin or its receptor. They appear, therefore, to have a form of functional “leptin resistance.” Data suggesting that some individuals produce less leptin per unit fat mass than others or have a form of relative leptin deficiency that predisposes to obesity are at present contradictory and unsettled. The mechanism for leptin resistance, and whether it can be overcome by raising leptin levels or combining leptin with other treatments in a subset of obese individuals, is not yet established. Some data suggest that leptin may not effectively cross the blood-brain barrier as levels rise. It is also apparent from animal studies that leptin-signaling inhibitors, such as SOCS3 and PTP1b, are involved in the leptin-resistant state.

PATHOLOGIC CONSEQUENCES OF OBESITY

(See also Chap. 416) Obesity has major adverse effects on health. Obesity is associated with an increase in mortality, with a 50–100% increased risk of death from all causes compared to normal-weight

individuals, mostly due to cardiovascular causes. Obesity and overweight together are the second leading cause of preventable death in the United States, accounting for 300,000 deaths per year. Mortality rates rise as obesity increases, particularly when obesity is associated with increased intraabdominal fat (see above). Life expectancy of a moderately obese individual could be shortened by 2–5 years, and a 20- to 30-year-old male with a BMI >45 may lose 13 years of life. It is likely that the degree to which obesity affects particular organ systems is influenced by susceptibility genes that vary in the population.

Insulin Resistance and Type 2 Diabetes Mellitus Hyperinsulinemia and insulin resistance are pervasive features of obesity, increasing with weight gain and diminishing with weight loss (Chap. 422). Insulin resistance is more strongly linked to intraabdominal fat than to fat in other depots. Molecular links between obesity and insulin resistance in fat, muscle, and liver have been sought for many years. Major factors include: (1) insulin itself, by inducing receptor downregulation; (2) free fatty acids that are increased and capable of impairing insulin action; (3) intracellular lipid accumulation; and (4) several circulating peptides produced by adipocytes, including the cytokines TNF- α and IL-6, RBP4, and the “adipokines” adiponectin and resistin, which have altered expression in obese adipocytes and can modify insulin action. Additional mechanisms are obesity-linked inflammation, including infiltration of macrophages into tissues including fat, and induction of the endoplasmic reticulum stress response, which can bring about resistance to insulin action in cells. Despite the prevalence of insulin resistance, most obese individuals do not develop diabetes, suggesting that diabetes requires an interaction between obesity-induced insulin resistance and other factors such as impaired insulin secretion (Chap. 417). Obesity, however, is a major risk factor for diabetes, and as many as 80% of patients with type 2 diabetes mellitus are obese. Weight loss and exercise, even of modest degree, increase insulin sensitivity and often improve glucose control in diabetes.

Reproductive Disorders Disorders that affect the reproductive axis are associated with obesity in both men and women. Male hypogonadism is associated with increased adipose tissue, often distributed in a pattern more typical of females. In men whose weight is >160% ideal body weight (IBW), plasma testosterone and sex hormone-binding globulin (SHBG) are often reduced, and estrogen levels (derived from conversion of adrenal androgens in adipose tissue) are increased (Chap. 411). Gynecomastia may be seen. However, masculinization, libido, potency, and spermatogenesis are preserved in most of these individuals. Free testosterone may be decreased in morbidly obese men whose weight is >200% IBW.

Obesity has long been associated with menstrual abnormalities in women, particularly in women with upper body obesity (Chap. 412). Common findings are increased androgen production, decreased SHBG, and increased peripheral conversion of androgen to estrogen. Most obese women with oligomenorrhea have polycystic ovarian syndrome (PCOS), with its associated anovulation and ovarian hyperandrogenism; 40% of women with PCOS are obese. Most nonobese women with PCOS are also insulin-resistant, suggesting that insulin resistance, hyperinsulinemia, or the combination of the two are causative or contribute to the ovarian pathophysiology in PCOS in both obese and lean individuals. Increasing evidence supports a role for adipokines in mediating a link between obesity and the reproductive dysfunction of PCOS. In obese women with PCOS, weight loss or treatment with insulin-sensitizing drugs often restores normal menses. The increased conversion of androstenedione to estrogen, which occurs to a greater degree in women with lower body obesity, may contribute to the increased incidence of uterine cancer in postmenopausal women with obesity.

Cardiovascular Disease The Framingham Study revealed that obesity was an independent risk factor for the 26-year incidence of cardiovascular disease in men and women (including coronary disease, stroke, and congestive heart failure). The waist-to-hip ratio may be the best predictor of these risks. When the additional effects of hypertension

and glucose intolerance associated with obesity are included, the adverse impact of obesity is even more evident. The effect of obesity on cardiovascular mortality in women may be seen at BMIs as low as 25. Obesity, especially abdominal obesity, is associated with an atherogenic lipid profile; with increased low-density lipoprotein cholesterol, very-low-density lipoprotein, and triglyceride; and with decreased high-density lipoprotein cholesterol and decreased levels of the vascular protective adipokine adiponectin (Chap. 421). Obesity is also associated with hypertension. Measurement of blood pressure in the obese requires use of a larger cuff size to avoid artifactual increases. Obesity-induced hypertension is associated with increased peripheral resistance and cardiac output, increased sympathetic nervous system tone, increased salt sensitivity, and insulin-mediated salt retention; it is often responsive to modest weight loss.

Pulmonary Disease Obesity may be associated with a number of pulmonary abnormalities. These include reduced chest wall compliance, increased work of breathing, increased minute ventilation due to increased metabolic rate, and decreased functional residual capacity and expiratory reserve volume. Severe obesity may be associated with obstructive sleep apnea and the “obesity hypoventilation syndrome” with attenuated hypoxic and hypercapnic ventilatory responses. Sleep apnea can be obstructive (most common), central, or mixed and is associated with hypertension. Weight loss (10–20 kg) can bring substantial improvement, as can major weight loss following gastric bypass or restrictive surgery. Continuous positive airway pressure has been used with some success.

Hepatobiliary Disease Obesity is frequently associated with nonalcoholic fatty liver disease (NAFLD), and this association represents one of the most common causes of liver disease in industrialized countries. The hepatic fatty infiltration of NAFLD progresses in a subset to inflammatory nonalcoholic steatohepatitis (NASH) and more rarely to cirrhosis and hepatocellular carcinoma. Steatosis typically improves following weight loss, secondary to diet or bariatric surgery. The mechanism for the association remains unclear. Obesity is associated with enhanced biliary secretion of cholesterol, supersaturation of bile, and a higher incidence of gallstones, particularly cholesterol gallstones (Chap. 369). A person 50% above IBW has about a sixfold increased incidence of symptomatic gallstones. Paradoxically, fasting increases supersaturation of bile by decreasing the phospholipid component. Fasting-induced cholecystitis is a complication of extreme diets.

Cancer Obesity is associated with increased risk of several cancer types, and in addition can lead to poorer treatment outcomes and increased cancer mortality. Obesity in males is associated with higher mortality from cancer of the esophagus, colon, rectum, pancreas, liver, and prostate; obesity in females is associated with higher mortality from cancer of the gallbladder, bile ducts, breasts, endometrium, cervix, and ovaries. Some of the latter may be due to increased rates of conversion of androstenedione to estrone in adipose tissue of obese individuals. Other possible mechanistic links may involve hormones, growth factors, and cytokines whose levels are linked to nutritional state, including insulin, leptin, adiponectin, and IGF-I, as well as activation of signaling pathways linked to both obesity and cancer. It has been estimated that obesity accounts for 14% of cancer deaths in men and 20% in women in the United States.

Bone, Joint, and Cutaneous Disease Obesity is associated with an increased risk of osteoarthritis, no doubt partly due to the trauma of added weight bearing, but potentially linked as well to activation of inflammatory pathways that could promote synovial pathology. The prevalence of gout may also be increased (Chap. 395). One of the skin problems associated with obesity is acanthosis nigricans, manifested by darkening and thickening of the skinfolds on the neck, elbows, and dorsal interphalangeal spaces. Acanthosis reflects the severity of underlying insulin resistance and diminishes with weight loss. Friability of skin may be increased, especially in skinfolds, enhancing the risk of fungal and yeast infections. Finally, venous stasis is increased in the obese.

416 Evaluation and Management of Obesity

Robert F. Kushner

More than 66% of U.S. adults are categorized as overweight or obese, and the prevalence of obesity is increasing rapidly in most of the industrialized world. Children and adolescents also are becoming more obese, indicating that the current trends will accelerate over time. Obesity is associated with an increased risk of multiple health problems, including hypertension, type 2 diabetes, dyslipidemia, obstructive sleep apnea, nonalcoholic fatty liver disease, degenerative joint disease, and some malignancies. Thus, it is important for physicians to identify, evaluate, and treat patients for obesity and associated comorbid conditions.

EVALUATION

Physicians should screen all adult patients for obesity and offer intensive counseling and behavioral interventions to promote sustained weight loss. The five main steps in the evaluation of obesity, as described below, are (1) a focused obesity-related history, (2) a physical examination to determine the degree and type of obesity, (3) assessment of comorbid conditions, (4) determination of fitness level, and (5) assessment of the patient's readiness to adopt lifestyle changes.

The Obesity-Focused History Information from the history should address the following seven questions:

- What factors contribute to the patient's obesity?
- How is the obesity affecting the patient's health?

- What is the patient's level of risk from obesity?
- What does the patient find difficult about managing weight?
- What are the patient's goals and expectations?
- Is the patient motivated to begin a weight management program?
- What kind of help does the patient need?

Although the vast majority of cases of obesity can be attributed to behavioral factors that affect diet and physical activity patterns, the history may suggest secondary causes that merit further evaluation. Disorders to consider include polycystic ovarian syndrome, hypothyroidism, Cushing's syndrome, and hypothalamic disease. Drug-induced weight gain also should be considered. Common causes include medications for diabetes (insulin, sulfonylureas, thiazolidinediones); steroid hormones; psychotropic agents; mood stabilizers (lithium); antidepressants (tricyclics, monoamine oxidase inhibitors, paroxetine, mirtazapine); and antiepileptic drugs (valproate, gabapentin, carbamazepine). Other medications, such as nonsteroidal anti-inflammatory drugs and calcium channel blockers, may cause peripheral edema but do not increase body fat.

The patient's current diet and physical activity patterns may reveal factors that contribute to the development of obesity and may identify behaviors to target for treatment. This type of historic information is best obtained by the combination of a questionnaire and an interview.

Body Mass Index (BMI) and Waist Circumference Three key anthropometric measurements are important in evaluating the degree of obesity: weight, height, and waist circumference. The BMI, calculated as weight (kg)/height (m)² or as weight (lbs)/height (inches)² × 703, is used to classify weight status and risk of disease (Tables 416-1 and 416-2). BMI provides an estimate of body fat and is related to disease risk. Lower BMI thresholds for overweight and obesity have been proposed for the Asia-Pacific region since this population appears to be at risk for glucose and lipid abnormalities at lower body weights.

Excess abdominal fat, assessed by measurement of waist circumference or waist-to-hip ratio, is independently associated with a higher risk for diabetes mellitus and cardiovascular disease. Measurement of the waist circumference is a surrogate for visceral adipose tissue and should be performed in the horizontal plane above the iliac crest (Table 416-3).

Physical Fitness Several prospective studies have demonstrated that physical fitness, reported by questionnaire or measured by a maximal treadmill exercise test, is an important predictor of all-cause mortality rate independent of BMI and body composition. These observations highlight the importance of taking a physical activity and exercise history during examination as well as emphasizing physical activity as a treatment approach.

Obesity-Associated Comorbid Conditions The evaluation of comorbid conditions should be based on presentation of symptoms, risk factors, and index of suspicion. For all patients, a fasting lipid panel should be performed (total, low-density lipoprotein, and high-density lipoprotein cholesterol and triglyceride levels) and a fasting blood glucose level and blood pressure determined. Symptoms and diseases that are directly or indirectly related to obesity are listed in Table 416-4. Although individuals vary, the number and severity of organ-specific comorbid conditions usually rise with increasing levels of obesity. Patients at very high absolute risk include those with the following: established coronary heart disease; presence of other atherosclerotic diseases, such as peripheral arterial disease, abdominal aortic aneurysm, and symptomatic carotid artery disease; type 2 diabetes; and sleep apnea.

Assessing the Patient's Readiness to Change An attempt to initiate lifestyle changes when the patient is not ready usually leads to frustration and may hamper future weight-loss efforts. Assessment includes

TABLE 416-1 BODY MASS INDEX (BMI)

BMI	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35		
Height (Inches)	Body Weight (Pounds)																		
58	91	96	100	105	110	115	119	124	129	134	138	143	148	153	158	162	167		
59	94	99	104	109	114	119	124	128	133	138	143	148	153	158	163	168	173		
60	97	102	107	112	118	123	128	133	138	143	148	153	158	163	168	174	179		
61	100	106	111	116	122	127	132	137	143	148	153	158	164	169	174	180	185		
62	104	109	115	120	126	131	136	142	147	153	158	164	169	175	180	186	191		
63	107	113	118	124	130	135	141	146	152	158	163	169	175	180	186	191	197		
64	110	116	122	128	134	140	145	151	157	163	169	174	180	186	192	197	204		
65	114	120	126	132	138	144	150	156	162	168	174	180	186	192	198	204	210		
66	118	124	130	136	142	148	155	161	167	173	179	186	192	198	204	210	216		
67	121	127	134	140	146	153	159	166	172	178	185	191	198	204	211	217	223		
68	125	131	138	144	151	158	164	171	177	184	190	197	203	210	216	223	230		
69	128	135	142	149	155	162	169	176	182	189	196	203	209	216	223	230	236		
70	132	139	146	153	160	167	174	181	188	195	202	209	216	222	229	236	243		
71	136	143	150	157	165	172	179	186	193	200	208	215	222	229	236	243	250		
72	140	147	154	162	169	177	184	191	199	206	213	221	228	235	242	250	258		
73	144	151	159	166	174	182	189	197	204	212	219	227	235	242	250	257	265		
74	148	155	163	171	179	186	194	202	210	218	225	233	241	249	256	264	272		
75	152	160	168	176	184	192	200	208	216	224	232	240	248	256	264	272	279		
76	156	164	172	180	189	197	205	213	221	230	238	246	254	263	271	279	287		
BMI	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
58	172	177	181	186	191	196	201	205	210	215	220	224	229	234	239	244	248	253	258
59	178	183	188	193	198	203	208	212	217	222	227	232	237	242	247	252	257	262	267
60	184	189	194	199	204	209	215	220	225	230	235	240	245	250	255	261	266	271	276
61	190	195	201	206	211	217	222	227	232	238	243	248	254	259	264	269	275	280	285
62	196	202	207	213	218	224	229	235	240	246	251	256	262	267	273	278	284	289	295
63	203	208	214	220	225	231	237	242	248	254	259	265	270	278	282	287	293	299	304
64	209	215	221	227	232	238	244	250	256	262	267	273	279	285	291	296	302	308	314
65	216	222	228	234	240	246	252	258	264	270	276	282	288	294	300	306	312	318	324
66	223	229	235	241	247	253	260	266	272	278	284	291	297	303	309	315	322	328	334
67	230	236	242	249	255	261	268	274	280	287	293	299	306	312	319	325	331	338	344
68	236	243	249	256	262	269	276	282	289	295	302	308	315	322	328	335	341	348	354
69	243	250	257	263	270	277	284	291	297	304	311	318	324	331	338	345	351	358	365
70	250	257	264	271	278	285	292	299	306	313	320	327	334	341	348	355	362	369	376
71	257	265	272	279	286	293	301	308	315	322	329	338	343	351	358	365	372	379	386
72	265	272	279	287	294	302	309	316	324	331	338	346	353	361	368	375	383	390	397
73	272	280	288	295	302	310	318	325	333	340	348	355	363	371	378	386	393	401	408
74	280	287	295	303	311	319	326	334	342	350	358	365	373	381	389	396	404	412	420
75	287	295	303	311	319	327	335	343	351	359	367	375	383	391	399	407	415	423	431
76	295	304	312	320	328	336	344	353	361	369	377	385	394	402	410	418	426	435	443

patient motivation and support, stressful life events, psychiatric status, time availability and constraints, and appropriateness of goals and expectations. Readiness can be viewed as the balance of two opposing forces: (1) motivation, or the patient's desire to change; and (2) resistance, or the patient's resistance to change.

TABLE 416-2 CLASSIFICATION OF WEIGHT STATUS AND DISEASE RISK

Classification	Body Mass Index (kg/m ²)	Obesity Class	Disease Risk
Underweight	<18.5	—	—
Healthy weight	18.5–24.9	—	—
Overweight	25.0–29.9	—	Increased
Obesity	30.0–34.9	I	High
Obesity	35.0–39.9	II	Very high
Extreme obesity	≥40	III	Extremely high

Source: Adapted from the National Institutes of Health, National Heart, Lung, and Blood Institute: *Clinical Guidelines on the Identification, Evaluation, and Treatment of Overweight and Obesity in Adults*. U.S. Department of Health and Human Services, U.S. Public Health Service, 1998.

A helpful method to begin a readiness assessment is to use the motivational interviewing technique of “anchoring” the patient's interest and confidence to change on a numerical scale. With this technique, the patient is asked to rate—on a scale from 0 to 10, with 0 being not so important (or confident) and 10 being very important (or confident)—his or her level of interest in and confidence about losing weight at this time. This exercise helps establish readiness to change and also serves as a basis for further dialogue.

TREATMENT OBESITY

THE GOAL OF THERAPY

The primary goals of treatment are to improve obesity-related comorbid conditions and to reduce the risk of developing future comorbidities. Information obtained from the history, physical examination, and diagnostic tests is used to determine risk and develop a treatment plan (Fig. 416-1). The decision of how aggressively to treat the patient and which modalities to use is determined by the patient's risk status, expectations, and available resources.

TABLE 416-3 ETHNIC-SPECIFIC CUTPOINT VALUES FOR WAIST CIRCUMFERENCE

Ethnic Group	Waist Circumference
Europeans	
Men	>94 cm (>37 in)
Women	>80 cm (>31.5 in)
South Asians and Chinese	
Men	>90 cm (>35 in)
Women	>80 cm (>31.5 in)
Japanese	
Men	>85 cm (>33.5 in)
Women	>90 cm (>35 in)
Ethnic South and Central Americans	Use South Asian recommendations until more specific data are available.
Sub-Saharan Africans	Use European data until more specific data are available.
Eastern Mediterranean and Middle Eastern (Arab) populations	Use European data until more specific data are available.

Source: From KGMM Alberti et al for the IDF Epidemiology Task Force Consensus Group: *Lancet* 366:1059, 2005.

Not all patients who are deemed obese by BMI alone need to be treated, as exemplified by the concepts of obesity paradox or the metabolically healthy obese. However, patients who present with obesity-related comorbidities and who would benefit from weight loss intervention should be managed proactively. Therapy for obesity always begins with lifestyle management and may include pharmacotherapy or surgery, depending on BMI risk category (Table 416-5). Setting an initial weight-loss goal of 8–10% over 6 months is a realistic target.

LIFESTYLE MANAGEMENT

Obesity care involves attention to three essential elements of lifestyle: dietary habits, physical activity, and behavior modification. Because obesity is fundamentally a disease of energy imbalance, all patients must learn how and when energy is consumed (diet), how and when energy is expended (physical activity), and how to incorporate this information into their daily lives (behavioral therapy). Lifestyle management has been shown to result in a modest (typically 3–5 kg) weight loss when compared with no treatment or usual care.

Diet Therapy The primary focus of diet therapy is to reduce overall calorie consumption. Guidelines from the National Heart, Lung, and Blood Institute recommend initiating treatment with a calorie deficit of 500–1000 kcal/d compared with the patient's habitual diet. This reduction is consistent with a goal of losing ~1–2 lbs per week. The calorie deficit can be instituted through dietary substitutions or alternatives. Examples include choosing smaller portion sizes, eating more fruits and vegetables, consuming more whole-grain cereals, selecting leaner cuts of meat and skimmed dairy products, reducing consumption of fried foods and other foods with added fats and oils, and drinking water instead of sugar-sweetened beverages. It is important that dietary counseling remain patient centered and that the goals set be practical, realistic, and achievable.

The macronutrient composition of the diet will vary with the patient's preference and medical condition. The 2010 U.S. Department of Agriculture Dietary Guidelines for Americans (Chap. 95e), which focus on health promotion and risk reduction, can be applied to treatment of overweight or obese patients. The recommendations include maintaining a diet rich in whole grains, fruits, vegetables, and dietary fiber; consuming two servings (8 oz) of fish high in omega 3 fatty acids per week; decreasing sodium intake to <2300 mg/d; consuming 3 cups of milk (or equivalent low-fat or fat-free dairy products) per day; limiting cholesterol intake to <300 mg/d; and keeping total fat intake at 20–35% of daily calories and saturated fat intake at <10% of daily calories. Application of these guidelines to specific calorie goals can be found on the website www.choosemyplate.gov. The revised Dietary Reference Intakes for Macronutrients released by the Institute of Medicine recommends that 45–65% of calories come from carbohydrates, 20–35% from fat, and 10–35% from protein. The guidelines also recommend daily fiber intake of 38 g (men) and 25 g (women) for persons over 50 years of age and 30 g (men) and 21 g (women) for those under age 50.

Since portion control is one of the most difficult strategies for patients to manage, the use of pre-prepared products such as meal replacements is a simple and convenient suggestion. Examples include frozen entrees, canned beverages, and bars. Use of meal replacements in the diet has been shown to result in a 7–8% weight loss.

TABLE 416-4 OBESITY-RELATED ORGAN SYSTEMS REVIEW

Cardiovascular	Respiratory
Hypertension	Dyspnea
Congestive heart failure	Obstructive sleep apnea
Cor pulmonale	Hypoventilation syndrome
Varicose veins	Pickwickian syndrome
Pulmonary embolism	Asthma
Coronary artery disease	Gastrointestinal
Endocrine	Gastroesophageal reflux disease
Metabolic syndrome	Nonalcoholic fatty-liver disease
Type 2 diabetes	Cholelithiasis
Dyslipidemia	Hernias
Polycystic ovarian syndrome	Colon cancer
Musculoskeletal	Genitourinary
Hyperuricemia and gout	Urinary stress incontinence
Immobility	Obesity-related glomerulopathy
Osteoarthritis (knees and hips)	Hypogonadism (male)
Low back pain	Breast and uterine cancer
Carpal tunnel syndrome	Pregnancy complications
Psychological	Neurologic
Depression/low self-esteem	Stroke
Body image disturbance	Idiopathic intracranial hypertension
Social stigmatization	Meralgia paresthetica
Integument	Dementia
Striae distensae	
Stasis pigmentation of legs	
Lymphedema	
Cellulitis	
Intertrigo, carbuncles	
Acanthosis nigricans	
Acrochordons (skin tags)	
Hidradenitis suppurativa	

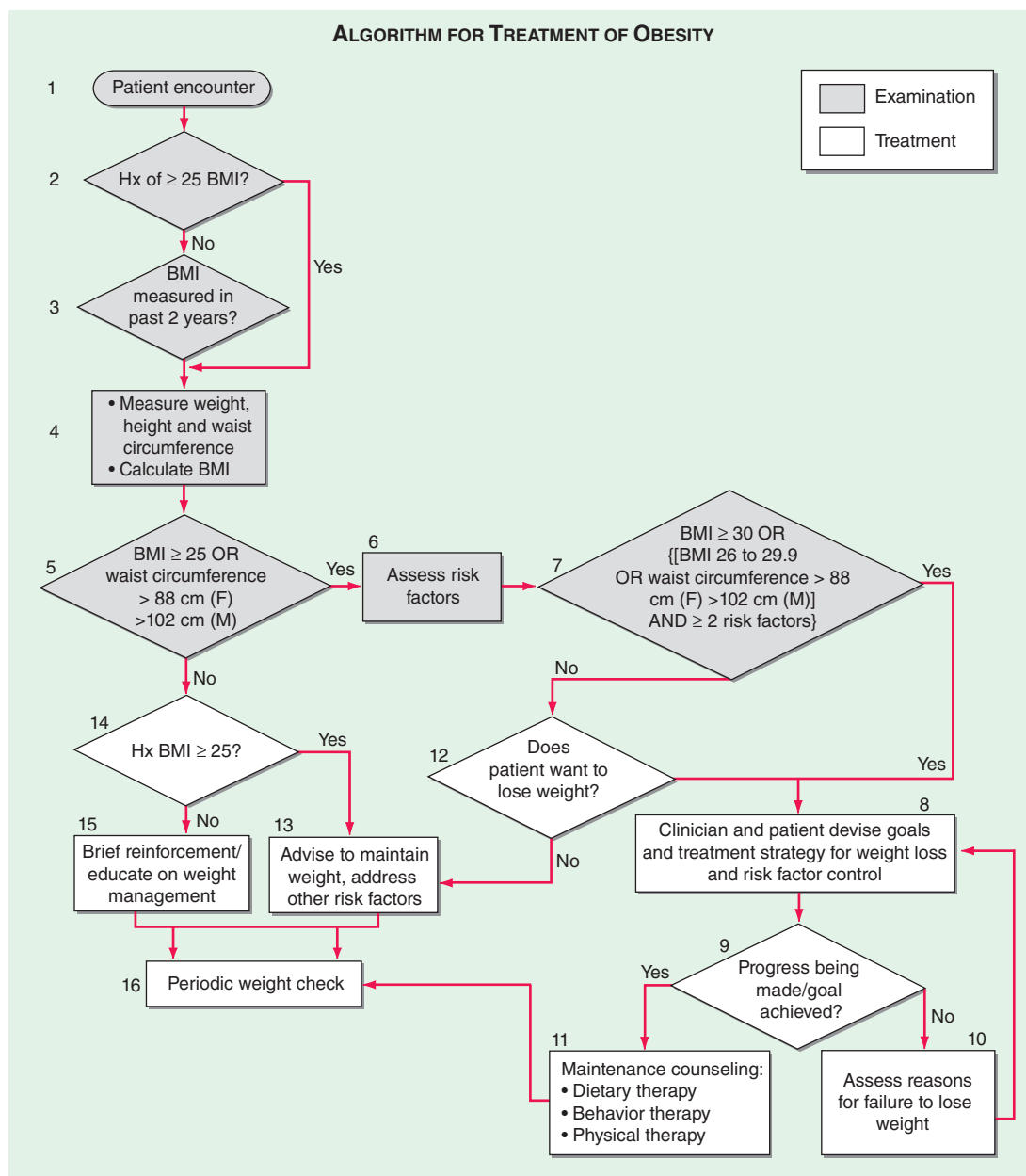


FIGURE 416-1 Algorithm for the treatment of obesity. This algorithm applies only to assessment for overweight and obesity and subsequent decisions based on that assessment. It does not reflect initial overall assessment for other conditions that the physician may wish to perform. BMI, body mass index; Hx, history. (From the National, Heart, Lung, and Blood Institute: *Clinical guidelines on the identification, evaluation, and treatment of overweight and obesity in adults: The evidence report*. Washington, DC, US Department of Health and Human Services, 1998.)

Numerous randomized trials comparing diets of different macronutrient composition (e.g., low-carbohydrate, low-fat, Mediterranean) have shown that weight loss depends primarily on reduction of total caloric intake and adherence to the prescribed diet, not the specific proportions of carbohydrate, fat, and protein in the diet. The macronutrient composition will ultimately be determined by the

patient's taste preferences, cooking style, and culture. However, the patient's underlying medical problems are also important in guiding the recommended dietary composition. The dietary prescription will vary according to the patient's metabolic profile and risk factors. A consultation with a registered dietitian for medical nutrition therapy is particularly useful in considering patient preference and treatment of comorbid diseases.

Another dietary approach to consider is based on the concept of *energy density*, which refers to the number of calories (i.e., amount of energy) a food contains per unit of weight. People tend to ingest a constant volume of food regardless of caloric or macronutrient content. Adding water or fiber to a food decreases its energy density by increasing weight without affecting caloric content. Examples of foods with low-energy density include soups, fruits, vegetables, oatmeal, and lean meats. Dry foods and high-fat foods such as pretzels, cheese, egg yolks, potato chips, and red meat have a high-energy

TABLE 416-5 A GUIDE TO OPTING FOR TREATMENT FOR OBESITY

Treatment	BMI Category (kg/m ²)				
	25–26.9	27–29.9	30–34.9	35–39.9	≥40
Diet, exercise, behavioral therapy	With comorbidities	With comorbidities	+	+	+
Pharmacotherapy	—	With comorbidities	+	+	+
Surgery	—	—	—	With comorbidities	+

Source: From the National Heart, Lung, and Blood Institute, North American Association for the Study of Obesity (2000).

2396 density. Diets containing low-energy-dense foods have been shown to control hunger and thus to result in decreased caloric intake and weight loss.

Occasionally, very low-calorie diets (VLCDs) are prescribed as a form of aggressive dietary therapy. The primary purpose of a VLCD is to promote a rapid and significant (13- to 23-kg) short-term weight loss over a 3- to 6-month period. The proprietary formulas designed for this purpose typically supply ≤ 800 kcal, 50–80 g of protein, and 100% of the recommended daily intake for vitamins and minerals. According to a review by the National Task Force on the Prevention and Treatment of Obesity, indications for initiating a VLCD include the involvement of well-motivated individuals who are moderately to severely obese (BMI, >30 kg/m²), have failed at more conservative approaches to weight loss, and have a medical condition that would be immediately improved with rapid weight loss. These conditions include poorly controlled type 2 diabetes, hypertriglyceridemia, obstructive sleep apnea, and symptomatic peripheral edema. The risk for gallstone formation increases exponentially at rates of weight loss >1.5 kg/week (3.3 lb/week). Prophylaxis against gallstone formation with ursodeoxycholic acid (600 mg/d) is effective in reducing this risk. Because of the need for close metabolic monitoring, VLCDs usually are prescribed by physicians specializing in obesity care.

Physical Activity Therapy Although exercise alone is only moderately effective for weight loss, the combination of dietary modification and exercise is the most effective behavioral approach for the treatment of obesity. The most important role of exercise appears to be in the maintenance of the weight loss. The 2008 Physical Activity Guidelines for Americans (www.health.gov/paguidelines) recommend that adults should engage in 150 min of moderate-intensity or 75 min a week of vigorous-intensity aerobic physical activity per week, performed in episodes of at least 10 min and preferably spread throughout the week. Focusing on simple ways to add physical activity into the normal daily routine through leisure activities, travel, and domestic work should be suggested. Examples include walking, using the stairs, doing housework and yard work, and engaging in sports. Asking the patient to wear a pedometer or accelerometer to monitor total accumulation of steps or kcal expended as part of the activities of daily living is a useful strategy. Step counts are highly correlated with activity level. Studies have demonstrated that lifestyle activities are as effective as structured exercise programs for improving cardiorespiratory fitness and weight loss. A high level of physical activity (>300 min of moderate-intensity activity per week) is often needed to lose weight and sustain weight loss. These exercise recommendations are daunting to most patients and need to be implemented gradually. Consultation with an exercise physiologist or personal trainer may be helpful.

Behavioral Therapy Cognitive behavioral therapy is used to help change and reinforce new dietary and physical activity behaviors. Strategies include self-monitoring techniques (e.g., journaling, weighing, and measuring food and activity); stress management; stimulus control (e.g., using smaller plates, not eating in front of the television or in the car); social support; problem solving; and cognitive restructuring to help patients develop more positive and realistic thoughts about themselves. When recommending any behavioral lifestyle change, the patient should be asked to identify what, when, where, and how the behavioral change will be performed. The patient should keep a record of the anticipated behavioral change so that progress can be reviewed at the next office visit. Because these techniques are time-consuming to implement, their supervision is often undertaken by ancillary office staff, such as a nurse-clinician or registered dietitian.

PHARMACOTHERAPY

Adjuvant pharmacologic treatments should be considered for patients with a BMI ≥ 30 kg/m² or—for patients who have concomitant obesity-related diseases and for whom dietary and physical activity therapy has not been successful—a BMI ≥ 27 kg/m². When an antiobesity medication is prescribed, patients should be actively

engaged in a lifestyle program that provides the strategies and skills needed to use the drug effectively, since such support increases total weight loss.

Medications for obesity have traditionally fallen into two major categories: appetite suppressants (*anorexiant*s) and gastrointestinal fat blockers. Appetite-suppressing medications have primarily targeted three monoamine receptor systems in the hypothalamus: noradrenergic, dopaminergic, and serotonergic receptors. Two new appetite suppressants were approved by the U.S. Food and Drug Administration (FDA) in 2012: lorcaserin and phentermine/topiramate (PHEN/TPM) extended release. Gastrointestinal fat blockers reduce the absorption of selective macronutrients, such as fat, from the gastrointestinal tract.

Centrally Acting Anorexiant Medications Anorexiant affect *satiety* (the absence of hunger after eating) and hunger (the biologic sensation that prompts eating). By increasing satiety and decreasing hunger, these agents help patients reduce caloric intake without a sense of deprivation. The target site for the actions of anorexiant is the ventromedial and lateral hypothalamic regions in the central nervous system (**Chap. 415e**). The biologic effect of these agents on appetite regulation is produced by augmentation of the neurotransmission of three monoamines: norepinephrine; serotonin (5-hydroxytryptamine, or 5-HT); and, to a lesser degree, dopamine. The classic sympathomimetic adrenergic agents (benzphetamine, phendimetrazine, diethylpropion, mazindol, and phentermine) function by stimulating norepinephrine release or by blocking its reuptake. Among the anorexiant, phentermine has been the most commonly prescribed; there is limited long-term data on its effectiveness. A 2002 review of six randomized, placebo-controlled trials of phentermine for weight control found that patients lost 0.6–6.0 additional kilograms of weight over 2–24 weeks of treatment. The most common side effects of the amphetamine-derived anorexiant are restlessness, insomnia, dry mouth, constipation, and increased blood pressure and heart rate.

PHEN/TPM is a combination drug that contains a catecholamine releaser (phentermine) and an anticonvulsant (topiramate). Topiramate is approved by the FDA as an anticonvulsant for the treatment of epilepsy and for the prophylaxis of migraine headaches. Weight loss was identified as an unintended side effect of topiramate during clinical trials for epilepsy. The mechanism responsible for weight loss is uncertain but is thought to be mediated through the drug's modulation of γ -aminobutyric acid receptors, inhibition of carbonic anhydrase, and antagonism of glutamate. PHEN/TPM has undergone two 1-year pivotal randomized, placebo-controlled, double-blind trials of efficacy and safety: EQUIP and CONQUER. In a third study, SEQUEL, 78% of CONQUER participants continued to receive their blinded treatment for an additional year. All participants received diet and exercise counseling. Participant numbers, eligibility, characteristics, and weight loss outcomes are displayed in **Table 416-6**. Intention-to-treat 1-year placebo-subtracted weight loss for the PHEN/TPM 15-mg/92-mg dose was 9.3% and 8.6%, respectively, in the EQUIP and CONQUER trials. Clinical and statistical dose-dependent improvements were seen in selected cardiovascular and metabolic outcome measurements that were related to the weight loss. The most common adverse events experienced by the drug-randomized group were paresthesias, dry mouth, constipation, dysgeusia, and insomnia. Because of an increased risk of congenital fetal oral-cleft formation from topiramate, the FDA approval of PHEN/TPM stipulated a Risk Evaluation and Mitigation Strategies requirement to educate prescribers about the need for active birth control among women of childbearing age and a contraindication for use during pregnancy.

Lorcaserin is a selective 5-HT_{2C} receptor agonist with a functional selectivity ~ 15 times that of 5-HT_{2A} receptors and 100 times that of 5-HT_{2B} receptors. This selectivity is important, since the drug-induced valvulopathy documented with two other serotonergic agents that were removed from the market—fenfluramine

TABLE 416-6 CLINICAL TRIALS FOR WEIGHT LOSS MEDICATIONS^a

	Lorcaserin		PHEN/TPM ^d	
	BLOOM ^b	BLOSSOM ^c	EQUIP	CONQUER
No. of participants (ITT-LOCF)	3182	4008	1230	2448
Age (years)	18–65	18–65	≥35	27–45
BMI (kg/m ²)	27–45	27–45	18–70	18–70
Comorbid conditions (cardiovascular and metabolic)	≥1	≥1	≥1	≥2
Mean weight loss (%) with treatment vs. placebo	5.8 vs. 2.2	4.8 vs. 2.8	11 vs. 1.6	10.4 vs. 1.8
Placebo-subtracted weight loss (%)	3.6	3.0	9.3	8.6
Categorical change in 5% weight loss with treatment vs. placebo	47.5 vs. 20.3	47.2 vs. 25	67 vs. 17	70 vs. 21
Completion rate (%)	Lorcaserin, 55.4; placebo, 45.1	55.5	59.9	62

^aTable shows a comparison of two 1-year prospective, randomized, double-blind trials of lorcaserin (BLOOM and BLOSSOM) and phentermine-topiramate extended release (EQUIP and CONQUER). ^bLorcaserin dose: 10 mg bid. ^cLorcaserin dose: 10 mg bid or qd. ^dPhentermine-topiramate extended release dose: 15 mg/92 mg.

Abbreviations: BMI, body mass index (see Table 416-1); ITT-LOCF, intention to treat, last observation carried forward; PHEN/TPM, phentermine-topiramate extended release.

and dexfenfluramine—was due to activation of the 5-HT_{2B} receptors expressed on cardiac valvular interstitial cells. By activating the 5-HT_{2C} receptor, lorcaserin is thought to decrease food intake through the pro-opiomelanocortin system of neurons.

Lorcaserin has undergone two randomized, placebo-controlled, double-blind trials for efficacy and safety. Participants were randomized to receive lorcaserin (10 mg bid) or placebo in the BLOOM study and to receive lorcaserin (10 mg bid or qd) or placebo in the BLOSSOM study. All participants received diet and exercise counseling. Participant numbers, eligibility, characteristics, and weight loss outcomes are displayed in Table 416-6. Overweight or obese subjects had at least one coexisting condition (hypertension, dyslipidemia, cardiovascular disease, impaired glucose tolerance, or sleep apnea)—medical conditions that are commonly seen in the office setting. Intention-to-treat 1-year placebo-subtracted weight loss was 3.6% and 3.0%, respectively, in the BLOOM and BLOSSOM trials. Echocardiography was performed at the screening visit and at scheduled time points over the course of the studies. There was no difference in the development of FDA-defined valvulopathy between drug-treated and placebo-treated participants at 1 year or 2 years. Modest statistical improvements consistent with the weight loss were seen in selected cardiovascular and metabolic outcome measurements. The most common adverse events experienced by the drug group were headache, dizziness, and nausea.

In approving both PHEN/TPM and lorcaserin, the FDA introduced a new provision with important clinical relevance: a prescription trial period to assess effectiveness. Response to both medications should be assessed after 3 months of treatment. For lorcaserin, the medication should be discontinued if the patient has not lost at least 5% of body weight by that point. For PHEN/TPM, if the patient has not lost at least 3% of body weight at 3 months, the clinician can either escalate the dose and reassess progress at 6 months or discontinue treatment entirely.

Peripherally Acting Medications Orlistat (XenicalTM) is a synthetic hydrogenated derivative of a naturally occurring lipase inhibitor, lipostatin, that is produced by the mold *Streptomyces toxytricini*. This drug is a potent, slowly reversible inhibitor of pancreatic, gastric, and carboxylester lipases and phospholipase A₂, which are required for the hydrolysis of dietary fat into fatty acids and monoacylglycerols. Orlistat acts in the lumen of the stomach and small intestine by forming a covalent bond with the active site of these lipases. Taken at a therapeutic dose of 120 mg tid, orlistat blocks the digestion and absorption of ~30% of dietary fat. After discontinuation of the drug, fecal fat content usually returns to normal within 48–72 h.

Multiple randomized, double-blind, placebo-controlled studies have shown that, after 1 year, orlistat produces a weight loss of ~9–10%, whereas placebo recipients have a 4–6% weight loss. Because orlistat is minimally (<1%) absorbed from the gastrointestinal tract, it has no systemic side effects. The drug's tolerability is related to the malabsorption of dietary fat and the subsequent passage of fat in the feces. Adverse gastrointestinal effects, including flatus with discharge, fecal urgency, fatty/oily stool, and increased defecation, are reported in at least 10% of orlistat-treated patients. These side effects generally are experienced early, diminish as patients control their dietary fat intake, and only infrequently cause patients to withdraw from clinical trials. When taken concomitantly, psyllium mucilloid is helpful in controlling orlistat-induced gastrointestinal side effects. Because serum concentrations of the fat-soluble vitamins D and E and β-carotene may be reduced by orlistat treatment, vitamin supplements are recommended to prevent potential deficiencies. Orlistat was approved for over-the-counter use in 2007.

Antiobesity Drugs in Development Two additional medications are currently in development. Bupropion and naltrexone (ContraveTM)—a dopamine and norepinephrine reuptake inhibitor and an opioid receptor antagonist, respectively—are theoretically combined to dampen the motivation/reinforcement that food brings (dopamine effect) and the pleasure/palatability of eating (opioid effect). In the COR-1 randomized, double-blind, placebo-controlled trial, 1742 enrolled participants, who were 18–65 years of age and had BMIs of 30–45 kg/m², were randomized to receive naltrexone (16 mg/d) plus bupropion (360 mg/d), naltrexone (32 mg/d) plus bupropion (360 mg/d), or placebo. Mean change in body weight for the three groups was 5.0%, 6.1%, and 1.3%, respectively. The most common adverse events were nausea, headache, constipation, dizziness, vomiting, and dry mouth. However, the FDA rejected the drug in 2011 because of cardiovascular concerns and concluded that a large-scale study of the long-term cardiovascular effects of naltrexone would be needed before approval could be considered.

Liraglutide, a glucagon-like peptide 1 receptor agonist currently approved for the treatment of type 2 diabetes, has independent weight loss effects via hypothalamic neural activation causing appetite suppression. In a double-blind, placebo-controlled trial, 564 adults with BMIs of 30–40 kg/m² were randomized to receive once-daily SC liraglutide (1.2, 1.8, 2.4, or 3.0 mg), placebo, or open-label orlistat (120 mg tid) for 1 year. The liraglutide and placebo recipients were switched to 2.4 mg of liraglutide during the second year and then to 3.0 mg for an additional year. One-year placebo-subtracted mean weight loss was 5.8 kg for liraglutide and 3.8 kg more than those on orlistat. The most common side effects were nausea, vomiting, and change in bowel habits.

Bariatric surgery (Fig. 416-2) can be considered for patients with severe obesity (BMI, ≥ 40 kg/m²) or for those with moderate obesity (BMI, ≥ 35 kg/m²) associated with a serious medical condition. Weight loss surgeries have traditionally been classified into three categories on the basis of anatomic changes: restrictive, restrictive malabsorptive, and malabsorptive. More recently, however, the clinical benefits of bariatric surgery in achieving weight loss and alleviating metabolic comorbidities have been attributed largely to changes in the physiologic responses of gut hormones and in adipose tissue metabolism. Metabolic effects resulting from bypassing the foregut include altered responses of ghrelin, glucagon-like peptide 1, peptide YY3-36, and oxyntomodulin. Additional effects on food intake and body weight control may be attributed to changes in vagal signaling. The loss of fat mass, particularly visceral fat, is associated with multiple metabolic, adipokine, and inflammatory changes that include improved insulin sensitivity and glucose disposal; reduced free fatty acid flux; increased adiponectin levels; and decreased interleukin 6, tumor necrosis factor α , and high-sensitivity C-reactive protein levels.

Restrictive surgeries limit the amount of food the stomach can hold and slow the rate of gastric emptying. *Laparoscopic adjustable gastric banding* is the prototype of this category. The first banding device, the LAP-BAND, was approved for use in the United States in 2001 and the second, the REALIZE band, in 2007. In contrast to previous devices, these bands have diameters that are adjustable by way of their connection to a reservoir that is implanted under the skin. Injection of saline into the reservoir and removal of saline from the reservoir tighten and loosen the band's internal diameter, respectively, thus changing the size of the gastric opening. The

mean percentage of total body weight lost at 5 years is estimated at 20–25%. In *laparoscopic sleeve gastrectomy*, the stomach is restricted by stapling and dividing it vertically, removing ~80% of the greater curvature, and leaving a slim banana-shaped remnant stomach along the lesser curvature. Weight loss after this procedure is superior to that after laparoscopic adjustable gastric banding.

The three restrictive-malabsorptive bypass procedures combine the elements of gastric restriction and selective malabsorption. These procedures are Roux-en-Y gastric bypass, biliopancreatic diversion, and biliopancreatic diversion with duodenal switch (Fig. 416-2). Roux-en-Y is the most commonly undertaken and most accepted bypass procedure. It may be performed with an open incision or by laparoscopy.

These procedures generally produce a 30–35% average total body weight loss that is maintained in nearly 60% of patients at 5 years. In general, mean weight loss is greater after the combined restrictive-malabsorptive procedures than after the restrictive procedures. Significant improvement in multiple obesity-related comorbid conditions, including type 2 diabetes, hypertension, dyslipidemia, obstructive sleep apnea, quality of life, and long-term cardiovascular events, has been reported. A meta-analysis of controlled clinical trials comparing bariatric surgery versus no surgery showed that surgery was associated with a reduced odds ratio (OR) risk of global mortality (OR = 0.55), cardiovascular death (OR = 0.58), and all-cause mortality (OR = 0.70).

Among the observed improvements in comorbidities, the prevention and treatment of type 2 diabetes resulting from bariatric surgery has garnered the most attention. Fifteen-year data from the Swedish Obese Subjects study demonstrated a marked reduction (i.e., by 78%) in the incidence of type 2 diabetes development among

obese patients who underwent bariatric surgery. Several randomized controlled studies have shown greater weight loss and more improved glycemic control at 1 and 2 years among surgical patients than among patients receiving conventional medical therapy. A retrospective cohort study of more than 4000 adults with diabetes found that, overall, 68.2% of patients experienced an initial complete type 2 diabetes remission within 5 years after surgery. However, among these patients, one-third redeveloped type 2 diabetes within 5 years. The rapid improvement seen in diabetes after restrictive-malabsorptive procedures is thought to be due to surgery-specific, weight-independent effects on glucose homeostasis brought about by alteration of gut hormones.

The mortality rate from bariatric surgery is generally <1% but varies with the procedure, the patient's age and comorbid conditions, and the experience of the surgical team. The most common surgical complications include stomal stenosis or marginal ulcers (occurring in 5–15% of patients) that present as prolonged nausea and vomiting after eating or inability to advance the diet to solid foods. These complications typically are treated by endoscopic balloon dilation and acid suppression therapy, respectively. For patients who undergo laparoscopic adjustable gastric banding, there are no intestinal absorptive abnormalities other than mechanical reduction in gastric size and outflow. Therefore, selective deficiencies are uncommon unless eating habits become unbalanced. In contrast, the restrictive-malabsorptive procedures carry an increased risk for micronutrient deficiencies of vitamin B₁₂, iron, folate, calcium, and vitamin D. Patients with restrictive-malabsorptive procedures require lifelong supplementation with these micronutrients.

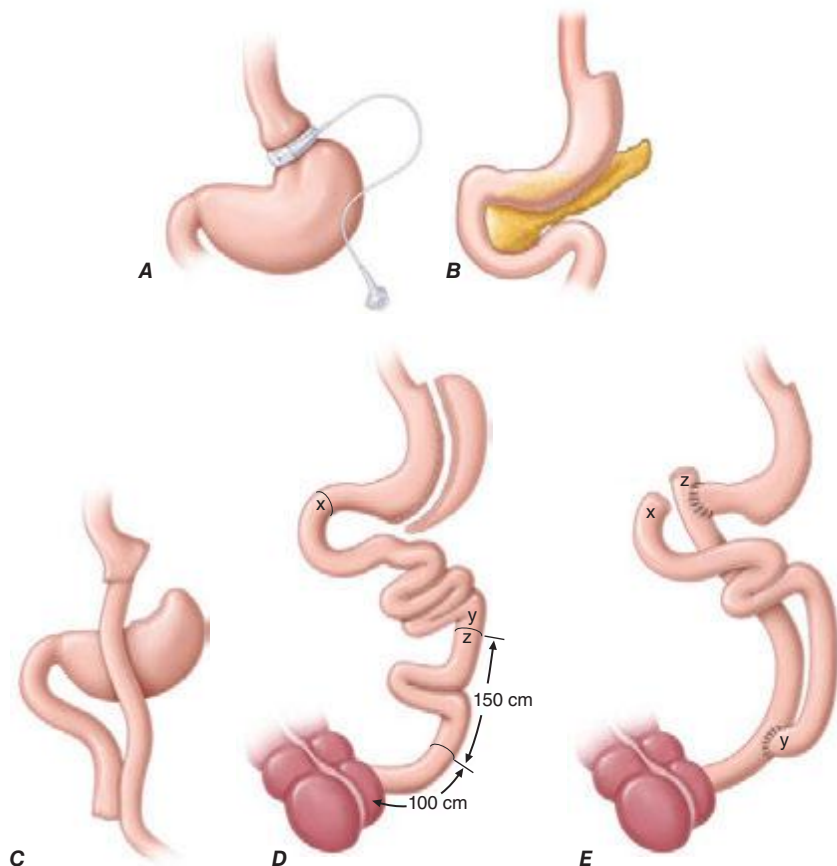


FIGURE 416-2 Bariatric surgical procedures. Examples of operative interventions used for surgical manipulation of the gastrointestinal tract. **A.** Laparoscopic adjustable gastric banding. **B.** Laparoscopic sleeve gastrectomy. **C.** The Roux-en-Y gastric bypass. **D.** Biliopancreatic diversion with duodenal switch. **E.** Biliopancreatic diversion. (From ML Kendrick, GF Dakin: *Mayo Clin Proc* 815:518, 2006; with permission.)

417 Diabetes Mellitus: Diagnosis, Classification, and Pathophysiology

Alvin C. Powers

Diabetes mellitus (DM) refers to a group of common metabolic disorders that share the phenotype of hyperglycemia. Several distinct types of DM are caused by a complex interaction of genetics and environmental factors. Depending on the etiology of the DM, factors contributing to hyperglycemia include reduced insulin secretion, decreased glucose utilization, and increased glucose production. The metabolic dysregulation associated with DM causes secondary pathophysiologic changes in multiple organ systems that impose a tremendous burden on the individual with diabetes and on the health care system. In the United States, DM is the leading cause of end-stage renal disease (ESRD), nontraumatic lower extremity amputations, and adult blindness. It also predisposes to cardiovascular diseases. With an increasing incidence worldwide, DM will be likely a leading cause of morbidity and mortality in the future.

CLASSIFICATION

DM is classified on the basis of the pathogenic process that leads to hyperglycemia, as opposed to earlier criteria such as age of onset or type of therapy (Fig. 417-1). There are two broad categories of DM, designated type 1 and type 2 (Table 417-1). However, there is increasing recognition of other forms of diabetes in which the pathogenesis is better understood. These other forms of diabetes may share features of

Type of Diabetes	Normal glucose tolerance	Hyperglycemia			
		Pre-diabetes*	Diabetes Mellitus		
			Impaired fasting glucose or impaired glucose tolerance	Not insulin requiring	Insulin required for control
Type 1			→		
Type 2		←			→
Other specific types		←			→
Gestational Diabetes		←			→
Time (years)		→			
FPG	<5.6 mmol/L (100 mg/dL)	5.6–6.9 mmol/L (100–125 mg/dL)	≥7.0 mmol/L (126 mg/dL)		
2-h PG	<7.8 mmol/L (140 mg/dL)	7.8–11.0 mmol/L (140–199 mg/dL)	≥11.1 mmol/L (200 mg/dL)		
HbA1C	<5.6%	5.7–6.4%	≥6.5%		

FIGURE 417-1 Spectrum of glucose homeostasis and diabetes mellitus (DM). The spectrum from normal glucose tolerance to diabetes in type 1 DM, type 2 DM, other specific types of diabetes, and gestational DM is shown from left to right. In most types of DM, the individual traverses from normal glucose tolerance to impaired glucose tolerance to overt diabetes (these should be viewed not as abrupt categories but as a spectrum). Arrows indicate that changes in glucose tolerance may be bidirectional in some types of diabetes. For example, individuals with type 2 DM may return to the impaired glucose tolerance category with weight loss; in gestational DM, diabetes may revert to impaired glucose tolerance or even normal glucose tolerance after delivery. The fasting plasma glucose (FPG), the 2-h plasma glucose (PG) after a glucose challenge, and the hemoglobin A_{1c} (HbA_{1c}) for the different categories of glucose tolerance are shown at the lower part of the figure. These values do not apply to the diagnosis of gestational DM. Some types of DM may or may not require insulin for survival. *Some use the term *increased risk for diabetes* or *intermediate hyperglycemia* (World Health Organization) rather than *prediabetes*. (Adapted from the American Diabetes Association, 2014.)

TABLE 417-1 ETIOLOGIC CLASSIFICATION OF DIABETES MELLITUS

1. Type 1 diabetes (beta cell destruction, usually leading to absolute insulin deficiency)
 - A. Immune-mediated
 - B. Idiopathic
- II. Type 2 diabetes (may range from predominantly insulin resistance with relative insulin deficiency to a predominantly insulin secretory defect with insulin resistance)
- III. Other specific types of diabetes
 - A. Genetic defects of beta cell development or function characterized by mutations in:
 1. Hepatocyte nuclear transcription factor (HNF) 4a (MODY 1)
 2. Glucokinase (MODY 2)
 3. HNF-1α (MODY 3)
 4. Insulin promoter factor-1 (IPF-1; MODY 4)
 5. HNF-1β (MODY 5)
 6. NeuroD1 (MODY 6)
 7. Mitochondrial DNA
 8. Subunits of ATP-sensitive potassium channel
 9. Proinsulin or insulin
 10. Other pancreatic islet regulators/proteins such as *KLF11*, *PAX4*, *BLK*, *GATA4*, *GATA6*, *SLC2A2* (GLUT2), *RFX6*, *GLIS3*
 - B. Genetic defects in insulin action
 1. Type A insulin resistance
 2. Leprechaunism
 3. Rabson-Mendenhall syndrome
 4. Lipodystrophy syndromes
 - C. Diseases of the exocrine pancreas—pancreatitis, pancreatectomy, neoplasia, cystic fibrosis, hemochromatosis, fibrocalculus pancreatopathy, mutations in carboxyl ester lipase
 - D. Endocrinopathies—acromegaly, Cushing's syndrome, glucagonoma, pheochromocytoma, hyperthyroidism, somatostatinoma, aldosteronoma
 - E. Drug- or chemical-induced—glucocorticoids, vacor (a rodenticide), pentamidine, nicotinic acid, diazoxide, β-adrenergic agonists, thiazides, calcineurin and mTOR inhibitors, hydantoin, asparaginase, α-interferon, protease inhibitors, antipsychotics (atypicals and others), epinephrine
 - F. Infections—congenital rubella, cytomegalovirus, coxsackievirus
 - G. Uncommon forms of immune-mediated diabetes—"stiff-person" syndrome, anti-insulin receptor antibodies
 - H. Other genetic syndromes sometimes associated with diabetes—Wolfram's syndrome, Down's syndrome, Klinefelter's syndrome, Turner's syndrome, Friedreich's ataxia, Huntington's chorea, Laurence-Moon-Biedl syndrome, myotonic dystrophy, porphyria, Prader-Willi syndrome
- IV. Gestational diabetes mellitus (GDM)

Abbreviation: MODY, maturity-onset diabetes of the young.

Source: Adapted from American Diabetes Association: Diabetes Care 37(Suppl 1):S14, 2014.

type 1 and/or type 2 DM. Both type 1 and type 2 DM are preceded by a phase of abnormal glucose homeostasis as the pathogenic processes progress. Type 1 DM is the result of complete or near-total insulin deficiency. Type 2 DM is a heterogeneous group of disorders characterized by variable degrees of insulin resistance, impaired insulin secretion, and increased glucose production. Distinct genetic and metabolic defects in insulin action and/or secretion give rise to the common phenotype of hyperglycemia in type 2 DM and have important potential therapeutic implications now that pharmacologic agents are available to target specific metabolic derangements. Type 2 DM is preceded by a period of abnormal glucose homeostasis classified as impaired fasting glucose (IFG) or impaired glucose tolerance (IGT).

Two features of the current classification of DM merit emphasis from previous classifications. First, the terms *insulin-dependent diabetes mellitus* (IDDM) and *non-insulin-dependent diabetes mellitus* (NIDDM) are obsolete. Because many individuals with type 2 DM eventually require insulin treatment for control of

2400 glycemia, the use of the term NIDDM generated considerable confusion. A second difference is that age or treatment modality is not a criterion. Although type 1 DM most commonly develops before the age of 30, an autoimmune beta cell destructive process can develop at any age. It is estimated that between 5 and 10% of individuals who develop DM after age 30 years have type 1 DM. Although type 2 DM more typically develops with increasing age, it is now being diagnosed more frequently in children and young adults, particularly in obese adolescents.

OTHER TYPES OF DM

Other etiologies for DM include specific genetic defects in insulin secretion or action, metabolic abnormalities that impair insulin secretion, mitochondrial abnormalities, and a host of conditions that impair glucose tolerance (Table 417-1). *Maturity-onset diabetes of the young* (MODY) and *monogenic diabetes* are subtypes of DM characterized by autosomal dominant inheritance, early onset of hyperglycemia (usually <25 years; sometimes in neonatal period), and impaired insulin secretion (discussed below). Mutations in the insulin receptor cause a group of rare disorders characterized by severe insulin resistance.

DM can result from pancreatic exocrine disease when the majority of pancreatic islets are destroyed. Cystic fibrosis-related DM is an important consideration in that patient population. Hormones that antagonize insulin action can also lead to DM. Thus, DM is often a feature of endocrinopathies such as acromegaly and Cushing's disease. Viral infections have been implicated in pancreatic islet destruction but are an extremely rare cause of DM. A form of acute onset of type 1 diabetes, termed *fulminant diabetes*, has been noted in Japan and may be related to viral infection of islets.

GESTATIONAL DIABETES MELLITUS

Glucose intolerance developing during pregnancy is classified as gestational diabetes mellitus (GDM). Insulin resistance is related to the metabolic changes of late pregnancy, and the increased insulin requirements may lead to IGT or diabetes. GDM occurs in ~7% (range 1–14%) of pregnancies in the United States; most women revert to normal glucose tolerance postpartum but have a substantial risk (35–60%) of developing DM in the next 10–20 years. The International Association of the Diabetes and Pregnancy Study Groups and the American Diabetes Association (ADA) recommend that diabetes diagnosed at the initial prenatal visit should be classified as “overt” diabetes rather than GDM. With the rising rates of obesity, the number of women being diagnosed with GDM or overt diabetes is rising worldwide.

EPIDEMIOLOGY AND GLOBAL CONSIDERATIONS



The worldwide prevalence of DM has risen dramatically over the past two decades, from an estimated 30 million cases in 1985 to 382 million in 2013 (Fig. 417-2). Based on current trends, the International Diabetes Federation projects that 592 million individuals will have diabetes by the year 2035 (see <http://www.idf.org/>). Although the prevalence of both type 1 and type 2 DM is increasing worldwide, the prevalence of type 2 DM is rising much more rapidly, presumably because of increasing obesity, reduced activity levels as countries become more industrialized, and the aging of the population. In 2013, the prevalence of diabetes in individuals from age 20–79 ranged from 23 to 37% in the 10 countries with the highest prevalence (Tuvalu, Federated States of Micronesia, Marshall Islands, Kiribati, Vanuatu, Cook Islands, Saudi Arabia, Nauru, Kuwait, and Qatar, in descending order of prevalence). The countries with the greatest number of individuals with diabetes in 2013 are China (98.4

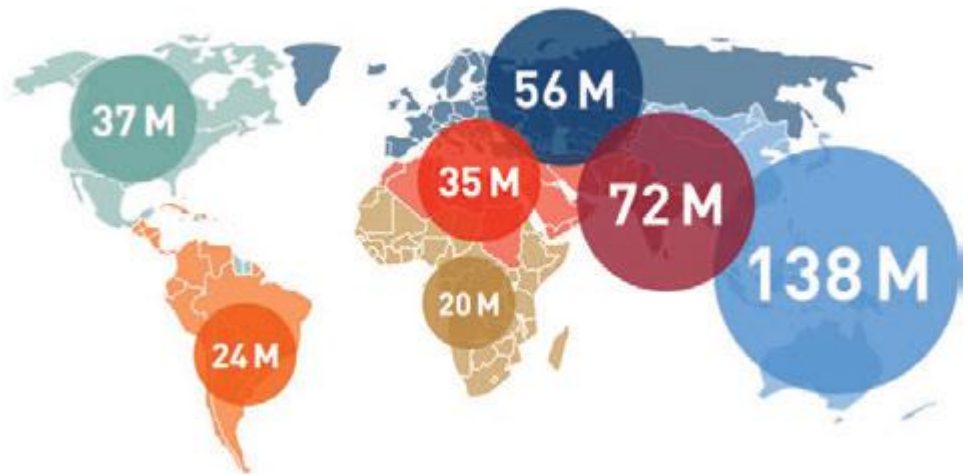


FIGURE 417-2 Worldwide prevalence of diabetes mellitus. Global estimate is 382 million individuals with diabetes. Regional estimates of the number of individuals with diabetes (20–79 years of age) are shown (2013). (Used with permission from the IDF Diabetes Atlas, the International Diabetes Federation, 2013.)

million), India (65.1 million), United States (24.4 million), Brazil (11.9 million), and the Russian Federation (10.9 million). Up to 80% of individuals with diabetes live in low-income or medium-income countries. In the most recent estimate for the United States (2012), the Centers for Disease Control and Prevention (CDC) estimated that 9.3% of the population had diabetes (~28% of the individuals with diabetes were undiagnosed); globally, it is estimated that 50% of individuals may be undiagnosed). The CDC estimated that the incidence and prevalence of diabetes doubled from 1990–2008, but appears to have plateaued from 2008–2012. DM increases with age. In 2012, the prevalence of DM in the United States was estimated to be 0.2% in individuals age <20 years and 12% in individuals age >20 years. In individuals age >65 years, the prevalence of DM was 26.9%. The prevalence is similar in men and women throughout most age ranges (14% and 11%, respectively, in individuals age >20 years). Worldwide, most individuals with diabetes are between the ages of 40 and 59 years.

There is considerable geographic variation in the incidence of both type 1 and type 2 DM. Scandinavia has the highest incidence of type 1 DM; the lowest incidence is in the Pacific Rim where it is 20- to 30-fold lower. Northern Europe and the United States have an intermediate rate. Much of the increased risk of type 1 DM is believed to reflect the frequency of high-risk human leukocyte antigen (HLA) alleles among ethnic groups in different geographic locations. The prevalence of type 2 DM and its harbinger, IGT, is highest in certain Pacific islands and the Middle East and intermediate in countries such as India and the United States. This variability is likely due to genetic, behavioral, and environmental factors. DM prevalence also varies among different ethnic populations within a given country, with indigenous populations usually having a greater incidence of diabetes than the general population of the country. For example, the CDC estimated that the age-adjusted prevalence of DM in the United States (age >20 years; 2010–2012) was 8% in non-Hispanic whites, 9% in Asian Americans, 13% in Hispanics, 13% in non-Hispanic blacks, and 16% in American-Indian and Alaskan native populations. The onset of type 2 DM occurs, on average, at an earlier age in ethnic groups other than non-Hispanic whites. In Asia, the prevalence of diabetes is increasing rapidly, and the diabetes phenotype appears to be somewhat different from that in the United States and Europe, with an onset at a lower body mass index (BMI) and younger age, greater visceral adiposity, and reduced insulin secretory capacity.

Diabetes is a major cause of mortality, but several studies indicate that diabetes is likely underreported as a cause of death. In the United States, diabetes was listed as the seventh leading cause of death in 2010. A recent estimate suggested that diabetes was responsible for almost 5.1 million deaths or 8% of deaths worldwide in 2013. In 2013, it was estimated that \$548 billion or 11% of health care expenditures worldwide were spent on individuals with diabetes.

TABLE 417-2 CRITERIA FOR THE DIAGNOSIS OF DIABETES MELLITUS

- Symptoms of diabetes plus random blood glucose concentration ≥ 11.1 mmol/L (200 mg/dL)^a or
- Fasting plasma glucose ≥ 7.0 mmol/L (126 mg/dL)^b or
- Hemoglobin A_{1c} $\geq 6.5\%$ ^c or
- 2-h plasma glucose ≥ 11.1 mmol/L (200 mg/dL) during an oral glucose tolerance test^d

^aRandom is defined as without regard to time since the last meal. ^bFasting is defined as no caloric intake for at least 8 h. ^cHemoglobin A_{1c} test should be performed in a laboratory using a method approved by the National Glycohemoglobin Standardization Program and correlated to the reference assay of the Diabetes Control and Complications Trial. Point-of-care hemoglobin A_{1c} should not be used for diagnostic purposes. ^dThe test should be performed using a glucose load containing the equivalent of 75 g anhydrous glucose dissolved in water, not recommended for routine clinical use.

Note: In the absence of unequivocal hyperglycemia and acute metabolic decompensation, these criteria should be confirmed by repeat testing on a different day.

Source: Adapted from American Diabetes Association: Diabetes Care 37(Suppl 1):S14, 2014.

DIAGNOSIS

Glucose tolerance is classified into three broad categories: normal glucose homeostasis, DM, or impaired glucose homeostasis. Glucose tolerance can be assessed using the fasting plasma glucose (FPG), the response to oral glucose challenge, or the hemoglobin A_{1c} (HbA_{1c}). An FPG < 5.6 mmol/L (100 mg/dL), a plasma glucose < 140 mg/dL (11.1 mmol/L) following an oral glucose challenge, and an HbA_{1c} $< 5.7\%$ are considered to define normal glucose tolerance. The International Expert Committee with members appointed by the ADA, the European Association for the Study of Diabetes, and the International Diabetes Federation have issued diagnostic criteria for DM (Table 417-2) based on the following premises: (1) the FPG, the response to an oral glucose challenge (oral glucose tolerance test [OGTT]), and HbA_{1c} differ among individuals, and (2) DM is defined as the level of glycemia at which diabetes-specific complications occur rather than on deviations from a population-based mean. For example, the prevalence of retinopathy in Native Americans (Pima Indian population) begins to increase at an FPG > 6.4 mmol/L (116 mg/dL) (Fig. 417-3).

An FPG ≥ 7.0 mmol/L (126 mg/dL), a glucose ≥ 11.1 mmol/L (200 mg/dL) 2 h after an oral glucose challenge, or an HbA_{1c} $\geq 6.5\%$ warrants the diagnosis of DM (Table 417-2). A random plasma glucose

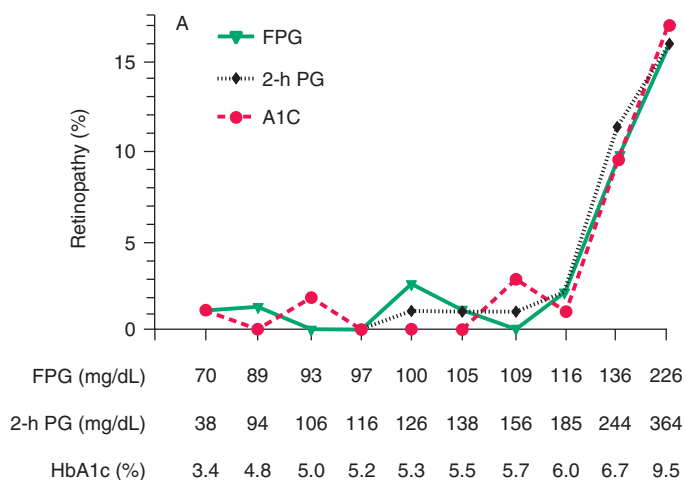


FIGURE 417-3 Relationship of diabetes-specific complication and glucose tolerance. This figure shows the incidence of retinopathy in Pima Indians as a function of the fasting plasma glucose (FPG), the 2-h plasma glucose after a 75-g oral glucose challenge (2-h PG), or the hemoglobin A_{1c} (HbA_{1c}). Note that the incidence of retinopathy greatly increases at a fasting plasma glucose > 116 mg/dL, a 2-h plasma glucose of 185 mg/dL, or an HbA_{1c} $> 6.5\%$. (Blood glucose values are shown in mg/dL; to convert to mmol/L, divide value by 18.) (Copyright 2002, American Diabetes Association. From Diabetes Care 25(Suppl 1): S5–S20, 2002.)

concentration ≥ 11.1 mmol/L (200 mg/dL) accompanied by classic symptoms of DM (polyuria, polydipsia, weight loss) is also sufficient for the diagnosis of DM (Table 417-2).

Abnormal glucose homeostasis (Fig. 417-1) is defined as (1) FPG = 5.6–6.9 mmol/L (100–125 mg/dL), which is defined as *impaired fasting glucose* (IFG); (2) plasma glucose levels between 7.8 and 11 mmol/L (140 and 199 mg/dL) following an oral glucose challenge, which is termed *impaired glucose tolerance* (IGT); or (3) HbA_{1c} of 5.7–6.4%. An HbA_{1c} of 5.7–6.4%, IFG, and IGT do not identify the same individuals, but individuals in all three groups are at greater risk of progressing to type 2 DM, have an increased risk of cardiovascular disease, and should be counseled about ways to decrease these risks (see below). Some use the terms *prediabetes*, *increased risk of diabetes*, or *intermediate hyperglycemia* (World Health Organization) for this category. These values for the fasting plasma glucose, the glucose following an oral glucose challenge, and HbA_{1c} are continuous variables and not discrete categories. The current criteria for the diagnosis of DM emphasize the HbA_{1c} or the FPG as the most reliable and convenient tests for identifying DM in asymptomatic individuals (however, some individuals may meet criteria for one test but not the other). OGTT, although still a valid means for diagnosing DM, is not often used in routine clinical care.

The diagnosis of DM has profound implications for an individual from both a medical and a financial standpoint. Thus, abnormalities on screening tests for diabetes should be repeated before making a definitive diagnosis of DM, unless acute metabolic derangements or a markedly elevated plasma glucose are present (Table 417-2). These criteria also allow for the diagnosis of DM to be withdrawn in situations when the glucose intolerance reverts to normal.

SCREENING

Widespread use of the FPG or the HbA_{1c} as a screening test for type 2 DM is recommended because (1) a large number of individuals who meet the current criteria for DM are asymptomatic and unaware that they have the disorder, (2) epidemiologic studies suggest that type 2 DM may be present for up to a decade before diagnosis, (3) some individuals with type 2 DM have one or more diabetes-specific complications at the time of their diagnosis, (4) treatment of type 2 DM may favorably alter the natural history of DM, diagnosis of prediabetes should spur efforts for diabetes prevention. The ADA recommends screening all individuals > 45 years every 3 years and screening individuals at an earlier age if they are overweight (BMI > 25 kg/m² or ethnically relevant definition for overweight) and have one additional risk factor for diabetes (Table 417-3). In contrast to type 2 DM, a long asymptomatic period of hyperglycemia is rare prior to the diagnosis of type 1 DM. A number of immunologic markers for type 1 DM are becoming available (discussed below), but their routine use outside a clinical trial is discouraged, pending the identification of clinically beneficial interventions for individuals at high risk for developing type 1 DM.

TABLE 417-3 RISK FACTORS FOR TYPE 2 DIABETES MELLITUS

- Family history of diabetes (i.e., parent or sibling with type 2 diabetes)
- Obesity (BMI ≥ 25 kg/m² or ethnically relevant definition for overweight)
- Physical inactivity
- Race/ethnicity (e.g., African American, Latino, Native American, Asian American, Pacific Islander)
- Previously identified with IFG, IGT, or an hemoglobin A_{1c} of 5.7–6.4%
- History of GDM or delivery of baby > 4 kg (9 lb)
- Hypertension (blood pressure $\geq 140/90$ mmHg)
- HDL cholesterol level < 35 mg/dL (0.90 mmol/L) and/or a triglyceride level > 250 mg/dL (2.82 mmol/L)
- Polycystic ovary syndrome or acanthosis nigricans
- History of cardiovascular disease

Abbreviations: BMI, body mass index; GDM, gestational diabetes mellitus; HDL, high-density lipoprotein; IFG, impaired fasting glucose; IGT, impaired glucose tolerance.

Source: Adapted from American Diabetes Association: Diabetes Care 37(Suppl 1):S14, 2014.

OVERALL REGULATION OF GLUCOSE HOMEOSTASIS

Glucose homeostasis reflects a balance between hepatic glucose production and peripheral glucose uptake and utilization. Insulin is the most important regulator of this metabolic equilibrium, but neural input, metabolic signals, and other hormones (e.g., glucagon) result in integrated control of glucose supply and utilization (Fig. 417-4). The organs that regulate glucose and lipids communicate by neural and humoral mechanisms with fat and muscle producing adipokines, myokines, and metabolites that influence liver function. In the fasting state, low insulin levels increase glucose production by promoting hepatic gluconeogenesis and glycogenolysis and reduce glucose uptake in insulin-sensitive tissues (skeletal muscle and fat), thereby promoting mobilization of stored precursors such as amino acids and free fatty acids (lipolysis). Glucagon, secreted by pancreatic alpha cells when blood glucose or insulin levels are low, stimulates glycogenolysis and gluconeogenesis by the liver and renal medulla (Chap. 420). Postprandially, the glucose load elicits a rise in insulin and fall in glucagon, leading to a reversal of these processes. Insulin, an anabolic hormone, promotes the storage of carbohydrate and fat and protein synthesis. The major portion of postprandial glucose is used by skeletal muscle, an effect of insulin-stimulated glucose uptake. Other tissues, most notably the brain, use glucose in an insulin-independent fashion. Factors secreted by skeletal myocytes (irisin), adipocytes (leptin, resistin, adiponectin, etc.), and bone also influence glucose homeostasis.

INSULIN BIOSYNTHESIS

Insulin is produced in the beta cells of the pancreatic islets. It is initially synthesized as a single-chain 86-amino-acid precursor polypeptide, proinsulin. Subsequent proteolytic processing removes the amino-terminal signal peptide, giving rise to proinsulin. Proinsulin is structurally related to insulin-like growth factors I and II, which bind weakly to the insulin receptor. Cleavage of an internal 31-residue fragment from proinsulin generates the C peptide and the A (21 amino acids) and B (30 amino acids) chains of insulin, which are connected by disulfide bonds. The mature insulin molecule and C peptide are stored together and co-secreted from secretory granules in the beta cells. Because C peptide is cleared more slowly than insulin, it is a useful marker of insulin secretion and allows discrimination of endogenous and exogenous sources of insulin in the evaluation of hypoglycemia (Chaps. 420 and 113). Pancreatic beta cells co-secrete islet amyloid polypeptide (IAPP) or amylin, a 37-amino-acid peptide, along with insulin. The role of IAPP in normal physiology is incompletely defined, but it is the major component of the amyloid fibrils found in the islets of patients with type 2 diabetes, and an analogue

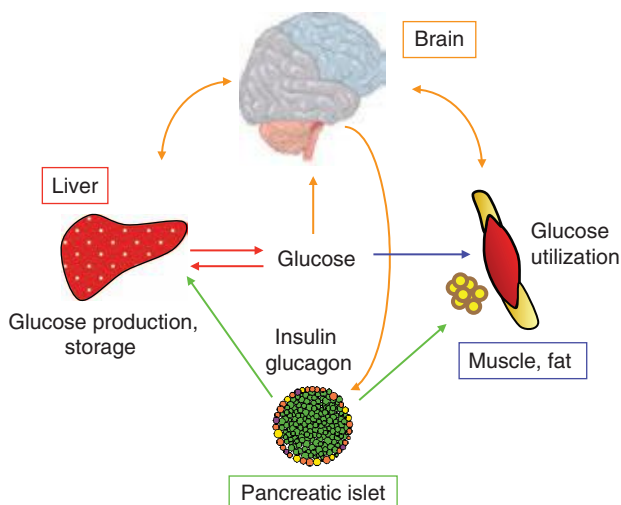


FIGURE 417-4 Regulation of glucose homeostasis. The organs shown contribute to glucose utilization, production, or storage. See text for a description of the communications (arrows), which can be neural or humoral.

is sometimes used in treating type 1 and type 2 DM. Human insulin is produced by recombinant DNA technology; structural alterations at one or more amino acid residues modify its physical and pharmacologic characteristics (Chap. 418).

INSULIN SECRETION

Glucose is the key regulator of insulin secretion by the pancreatic beta cell, although amino acids, ketones, various nutrients, gastrointestinal peptides, and neurotransmitters also influence insulin secretion. Glucose levels >3.9 mmol/L (70 mg/dL) stimulate insulin synthesis, primarily by enhancing protein translation and processing. Glucose stimulation of insulin secretion begins with its transport into the beta cell by a facilitative glucose transporter (Fig. 417-5). Glucose phosphorylation by glucokinase is the rate-limiting step that controls glucose-regulated insulin secretion. Further metabolism of glucose-6-phosphate via glycolysis generates ATP, which inhibits the activity of an ATP-sensitive K^+ channel. This channel consists of two separate proteins: one is the binding site for certain oral hypoglycemics (e.g., sulfonylureas, meglitinides); the other is an inwardly rectifying K^+ channel protein (Kir6.2). Inhibition of this K^+ channel induces beta cell membrane depolarization, which opens voltage-dependent calcium channels (leading to an influx of calcium) and stimulates insulin secretion. Insulin secretory profiles reveal a pulsatile pattern of hormone release, with small secretory bursts occurring about every 10 min, superimposed upon greater amplitude oscillations of about 80–150 min. Incretins are released from neuroendocrine cells of the gastrointestinal tract following food ingestion and amplify glucose-stimulated insulin secretion and suppress glucagon secretion. Glucagon-like peptide 1 (GLP-1), the most potent incretin, is released from L cells in the small intestine and stimulates insulin secretion only when the blood glucose is above the fasting level. Incretin analogues or pharmacologic agents that prolong the activity of endogenous GLP-1 enhance insulin secretion.

INSULIN ACTION

Once insulin is secreted into the portal venous system, ~50% is removed and degraded by the liver. Unextracted insulin enters the

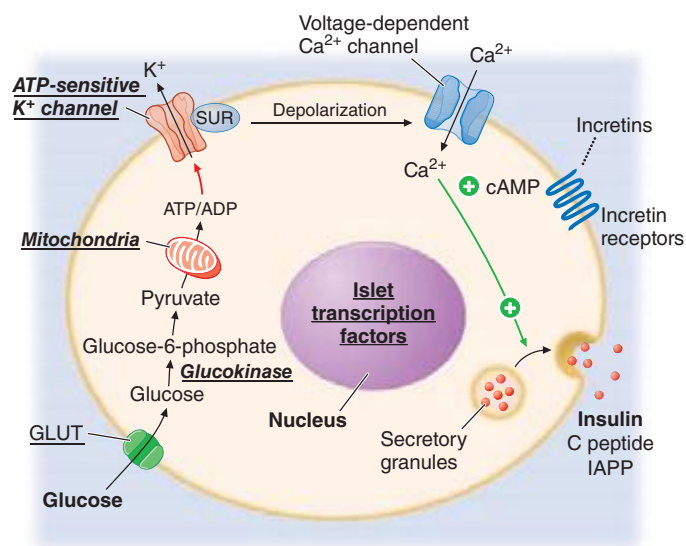


FIGURE 417-5 Mechanisms of glucose-stimulated insulin secretion and abnormalities in diabetes.

Glucose and other nutrients regulate insulin secretion by the pancreatic beta cell. Glucose is transported by a glucose transporter (GLUT1 and/or GLUT2 in humans, GLUT2 in rodents); subsequent glucose metabolism by the beta cell alters ion channel activity, leading to insulin secretion. The SUR receptor is the binding site for some drugs that act as insulin secretagogues. Mutations in the events or proteins underlined are a cause of monogenic forms of diabetes. ADP, adenosine diphosphate; ATP, adenosine triphosphate; cAMP, cyclic adenosine monophosphate; IAPP, islet amyloid polypeptide or amylin; SUR, sulfonylurea receptor.

systemic circulation where it binds to receptors in target sites. Insulin binding to its receptor stimulates intrinsic tyrosine kinase activity, leading to receptor autophosphorylation and the recruitment of intracellular signaling molecules, such as insulin receptor substrates (IRS). IRS and other adaptor proteins initiate a complex cascade of phosphorylation and dephosphorylation reactions, resulting in the widespread metabolic and mitogenic effects of insulin. As an example, activation of the phosphatidylinositol-3'-kinase (PI-3-kinase) pathway stimulates translocation of a facilitative glucose transporter (e.g., GLUT4) to the cell surface, an event that is crucial for glucose uptake by skeletal muscle and fat. Activation of other insulin receptor signaling pathways induces glycogen synthesis, protein synthesis, lipogenesis, and regulation of various genes in insulin-responsive cells.

PATHOGENESIS

TYPE 1 DM

Type 1 DM is the result of interactions of genetic, environmental, and immunologic factors that ultimately lead to the destruction of the pancreatic beta cells and insulin deficiency. Type 1 DM, which can develop at any age, develops most commonly before 20 years of age. Worldwide, the incidence of type 1 DM is increasing at the rate of 3–4% per year for uncertain reasons. Type 1 DM results from autoimmune beta cell destruction, and most, but not all, individuals have evidence of islet-directed autoimmunity. Some individuals who have the clinical phenotype of type 1 DM lack immunologic markers indicative of an autoimmune process involving the beta cells and the genetic markers of type 1 DM. These individuals are thought to develop insulin deficiency by unknown, nonimmune mechanisms and may be ketosis prone; many are African American or Asian in heritage. The temporal development of type 1 DM is shown schematically as a function of beta cell mass in Fig. 417-6. Individuals with a genetic susceptibility are thought to have normal beta cell mass at birth but begin to lose beta cells secondary to autoimmune destruction that occurs over months to years. This autoimmune process is thought to be triggered by an infectious or environmental stimulus and to be sustained by a beta cell-specific molecule. In the majority of patients, immunologic

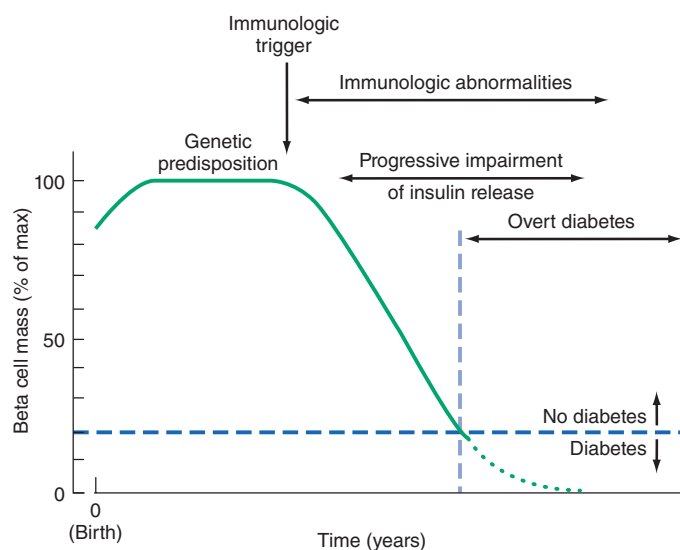


FIGURE 417-6 Temporal model for development of type 1 diabetes. Individuals with a genetic predisposition are exposed to a trigger that initiates an autoimmune process, resulting in a gradual decline in beta cell mass. The downward slope of the beta cell mass varies among individuals and may not be continuous. This progressive impairment in insulin release results in diabetes when ~80% of the beta cell mass is destroyed. A “honeymoon” phase may be seen in the first 1 or 2 years after the onset of diabetes and is associated with reduced insulin requirements. (Adapted from ER Kaufman: *Medical Management of Type 1 Diabetes*, 6th ed. American Diabetes Association, Alexandria, VA, 2012.)

markers appear after the triggering event but before diabetes becomes clinically overt. Beta cell mass then begins to decrease, and insulin secretion progressively declines, although normal glucose tolerance is maintained. The rate of decline in beta cell mass varies widely among individuals, with some patients progressing rapidly to clinical diabetes and others evolving more slowly. Features of diabetes do not become evident until a majority of beta cells are destroyed (70–80%). At this point, residual functional beta cells exist but are insufficient in number to maintain glucose tolerance. The events that trigger the transition from glucose intolerance to frank diabetes are often associated with increased insulin requirements, as might occur during infections or puberty. After the initial clinical presentation of type 1 DM, a “honeymoon” phase may ensue during which time glycemic control is achieved with modest doses of insulin or, rarely, insulin is not needed. However, this fleeting phase of endogenous insulin production from residual beta cells disappears and the individual becomes insulin deficient. Many individuals with long-standing type 1 DM produce a small amount of insulin (as reflected by C-peptide production), and some individuals with more than 50 years of type 1 DM have insulin-positive cells in the pancreas at autopsy.

GENETIC CONSIDERATIONS

Susceptibility to type 1 DM involves multiple genes. The concordance of type 1 DM in identical twins ranges between 40 and 60%, indicating that additional modifying factors are likely involved in determining whether diabetes develops. The major susceptibility gene for type 1 DM is located in the HLA region on chromosome 6. Polymorphisms in the HLA complex account for 40–50% of the genetic risk of developing type 1 DM. This region contains genes that encode the class II major histocompatibility complex (MHC) molecules, which present antigen to helper T cells and thus are involved in initiating the immune response (Chap. 373e). The ability of class II MHC molecules to present antigen is dependent on the amino acid composition of their antigen-binding sites. Amino acid substitutions may influence the specificity of the immune response by altering the binding affinity of different antigens for class II molecules.

Most individuals with type 1 DM have the HLA DR3 and/or DR4 haplotype. Refinements in genotyping of HLA loci have shown that the haplotypes DQA1*0301, DQB1*0302, and DQB1*0201 are most strongly associated with type 1 DM. These haplotypes are present in 40% of children with type 1 DM as compared to 2% of the normal U.S. population. However, most individuals with predisposing haplotypes do not develop diabetes.

In addition to MHC class II associations, genome association studies have identified at least 20 different genetic loci that contribute susceptibility to type 1 DM (polymorphisms in the promoter region of the insulin gene, the CTLA-4 gene, interleukin 2 receptor, *CTLA4*, and *PTPN22*, etc.). Genes that confer protection against the development of the disease also exist. The haplotype DQA1*0102, DQB1*0602 is extremely rare in individuals with type 1 DM (<1%) and appears to provide protection from type 1 DM.

Although the risk of developing type 1 DM is increased tenfold in relatives of individuals with the disease, the risk is relatively low: 3–4% if the parent has type 1 DM and 5–15% in a sibling (depending on which HLA haplotypes are shared). Hence, most individuals with type 1 DM do not have a first-degree relative with this disorder.

Pathophysiology Although other islet cell types (alpha cells [glucagon-producing], delta cells [somatostatin-producing], or PP cells [pancreatic polypeptide-producing]) are functionally and embryologically similar to beta cells and express most of the same proteins as beta cells, they are spared from the autoimmune destruction. Pathologically, the pancreatic islets have a modest infiltration of lymphocytes (a process termed *insulinitis*). After beta cells are destroyed, it is thought that the inflammatory process abates and the islets become atrophic. Studies of the autoimmune process in humans and in animal models of type 1 DM (NOD mouse and BB rat) have identified the following abnormalities in the humoral and cellular arms of the immune system: (1) islet cell autoantibodies; (2) activated lymphocytes in the islets,

2404 peripancreatic lymph nodes, and systemic circulation; (3) T lymphocytes that proliferate when stimulated with islet proteins; and (4) release of cytokines within the insulinitis. Beta cells seem to be particularly susceptible to the toxic effect of some cytokines (tumor necrosis factor α [TNF- α], interferon γ , and interleukin 1 [IL-1]). The precise mechanisms of beta cell death are not known but may involve formation of nitric oxide metabolites, apoptosis, and direct CD8+ T cell cytotoxicity. The islet destruction is mediated by T lymphocytes rather than islet autoantibodies, as these antibodies do not generally react with the cell surface of islet cells and are not capable of transferring DM to animals. Efforts to suppress the autoimmune process at the time of diagnosis of diabetes have largely been ineffective or only temporarily effective in slowing beta cell destruction.

Pancreatic islet molecules targeted by the autoimmune process include insulin, glutamic acid decarboxylase (GAD; the biosynthetic enzyme for the neurotransmitter GABA), ICA-512/IA-2 (homology with tyrosine phosphatases), and a beta cell-specific zinc transporter (ZnT-8). Most of the autoantigens are not beta cell-specific, which raises the question of how the beta cells are selectively destroyed. Current theories favor initiation of an autoimmune process directed at one beta cell molecule, which then spreads to other islet molecules as the immune process destroys beta cells and creates a series of secondary autoantigens. The beta cells of individuals who develop type 1 DM do not differ from beta cells of normal individuals because islets transplanted from a genetically identical twin are destroyed by a recurrence of the autoimmune process of type 1 DM.

Immunologic Markers Islet cell autoantibodies (ICAs) are a composite of several different antibodies directed at pancreatic islet molecules such as GAD, insulin, IA-2/ICA-512, and ZnT-8, and serve as a marker of the autoimmune process of type 1 DM. Assays for autoantibodies to GAD-65 are commercially available. Testing for ICAs can be useful in classifying the type of DM as type 1 and in identifying nondiabetic individuals at risk for developing type 1 DM. ICAs are present in the majority of individuals (>85%) diagnosed with new-onset type 1 DM, in a significant minority of individuals with newly diagnosed type 2 DM (5–10%), and occasionally in individuals with GDM (<5%). ICAs are present in 3–4% of first-degree relatives of individuals with type 1 DM. In combination with impaired insulin secretion after IV glucose tolerance testing, they predict a >50% risk of developing type 1 DM within 5 years. At present, the measurement of ICAs in nondiabetic individuals is a research tool because no treatments have been demonstrated to prevent the occurrence or progression to type 1 DM.

Environmental Factors Numerous environmental events have been proposed to trigger the autoimmune process in genetically susceptible individuals; however, none have been conclusively linked to diabetes. Identification of an environmental trigger has been difficult because the event may precede the onset of DM by several years (Fig. 417-6). Putative environmental triggers include viruses (coxsackie, rubella, enteroviruses most prominently), bovine milk proteins, and nitrosourea compounds. There is increasing interest in the microbiome and type 1 diabetes (Chap. 86e).


Prevention of Type 1 DM A number of interventions have prevented diabetes in animal models. None of these interventions have been successful in preventing type 1 DM in humans. For example, the Diabetes Prevention Trial-Type 1 concluded that administering insulin (IV or PO) to individuals at high risk for developing type 1 DM did not prevent type 1 DM. This is an area of active clinical investigation.

TYPE 2 DM

Insulin resistance and abnormal insulin secretion are central to the development of type 2 DM. Although the primary defect is controversial, most studies support the view that insulin resistance precedes an insulin secretory defect but that diabetes develops only when insulin secretion becomes inadequate. Type 2 DM likely encompasses a range of disorders with common phenotype of hyperglycemia. Most of our current understanding (and the discussion below) of the pathophysiology and genetics is based on studies of individuals of European

descent. It is becoming increasingly apparent that DM in other ethnic groups (Asian, African, and Latin American) has a somewhat different, but yet undefined, pathophysiology. In general, Latinos have greater insulin resistance and East Asians and South Asians have more beta cell dysfunction, but both defects are present in both populations. East and South Asians appear to develop type 2 DM at a younger age and a lower BMI. In some groups, DM that is ketosis prone (often obese) or ketosis-resistant (often lean) is seen.

GENETIC CONSIDERATIONS

 Type 2 DM has a strong genetic component. The concordance of type 2 DM in identical twins is between 70 and 90%. Individuals with a parent with type 2 DM have an increased risk of diabetes; if both parents have type 2 DM, the risk approaches 40%. Insulin resistance, as demonstrated by reduced glucose utilization in skeletal muscle, is present in many nondiabetic, first-degree relatives of individuals with type 2 DM. The disease is polygenic and multifactorial, because in addition to genetic susceptibility, environmental factors (such as obesity, nutrition, and physical activity) modulate the phenotype. The in utero environment also contributes, and either increased or reduced birth weight increases the risk of type 2 DM in adult life. The genes that predispose to type 2 DM are incompletely identified, but recent genome-wide association studies have identified a large number of genes that convey a relatively small risk for type 2 DM (>70 genes, each with a relative risk of 1.06–1.5). Most prominent is a variant of the transcription factor 7-like 2 gene that has been associated with type 2 DM in several populations and with IGT in one population at high risk for diabetes. Genetic polymorphisms associated with type 2 DM have also been found in the genes encoding the peroxisome proliferator-activated receptor γ , inward rectifying potassium channel, zinc transporter, IRS, and calpain 10. The mechanisms by which these genetic loci increase the susceptibility to type 2 DM are not clear, but most are predicted to alter islet function or development or insulin secretion. Although the genetic susceptibility to type 2 DM is under active investigation (it is estimated that <10% of genetic risk is determined by loci identified thus far), it is currently not possible to use a combination of known genetic loci to predict type 2 DM.

Pathophysiology Type 2 DM is characterized by impaired insulin secretion, insulin resistance, excessive hepatic glucose production, and abnormal fat metabolism. Obesity, particularly visceral or central (as evidenced by the hip-waist ratio), is very common in type 2 DM ($\geq 80\%$ of patients are obese). In the early stages of the disorder, glucose tolerance remains near-normal, despite insulin resistance, because the pancreatic beta cells compensate by increasing insulin output (Fig. 417-7). As insulin resistance and compensatory hyperinsulinemia progress, the pancreatic islets in certain individuals are unable to sustain the hyperinsulinemic state. IGT, characterized by elevations in postprandial glucose, then develops. A further decline in insulin secretion and an increase in hepatic glucose production lead to overt diabetes with fasting hyperglycemia. Ultimately, beta cell failure ensues. Although both insulin resistance and impaired insulin secretion contribute to the pathogenesis of type 2 DM, the relative contribution of each varies from individual to individual.

Metabolic Abnormalities • ABNORMAL MUSCLE AND FAT METABOLISM Insulin resistance, the decreased ability of insulin to act effectively on target tissues (especially muscle, liver, and fat), is a prominent feature of type 2 DM and results from a combination of genetic susceptibility and obesity. Insulin resistance is relative, however, because supranormal levels of circulating insulin will normalize the plasma glucose. Insulin dose-response curves exhibit a rightward shift, indicating reduced sensitivity, and a reduced maximal response, indicating an overall decrease in maximum glucose utilization (30–60% lower than in normal individuals). Insulin resistance impairs glucose utilization by insulin-sensitive tissues and increases hepatic glucose output; both effects contribute to the hyperglycemia. Increased hepatic glucose output predominantly accounts for increased FPG levels, whereas decreased peripheral glucose usage results in postprandial hyperglycemia. In skeletal muscle, there is a greater impairment in nonoxidative

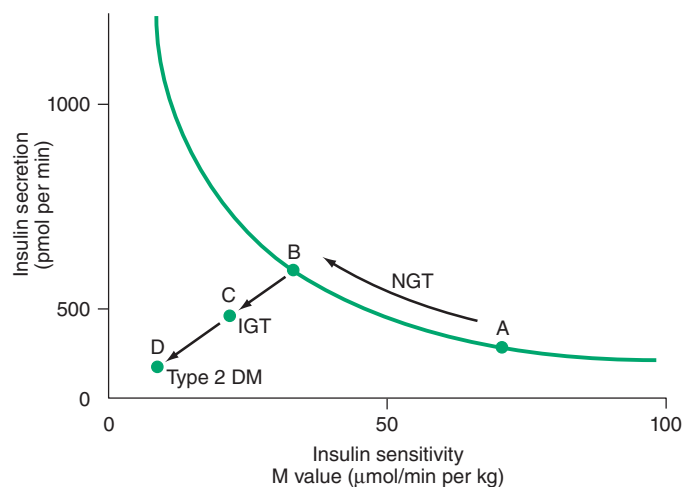


FIGURE 417-7 Metabolic changes during the development of type 2 diabetes mellitus (DM). Insulin secretion and insulin sensitivity are related, and as an individual becomes more insulin resistant (by moving from point A to point B), insulin secretion increases. A failure to compensate by increasing the insulin secretion results initially in impaired glucose tolerance (IGT; point C) and ultimately in type 2 DM (point D). NGT, normal glucose tolerance. (Adapted from SE Kahn: *J Clin Endocrinol Metab* 86:4047, 2001; RN Bergman, M Ader: *Trends Endocrinol Metab* 11:351, 2000.)

glucose usage (glycogen formation) than in oxidative glucose metabolism through glycolysis. Glucose metabolism in insulin-independent tissues is not altered in type 2 DM.

The precise molecular mechanism leading to insulin resistance in type 2 DM has not been elucidated. Insulin receptor levels and tyrosine kinase activity in skeletal muscle are reduced, but these alterations are most likely secondary to hyperinsulinemia and are not a primary defect. Therefore, “postreceptor” defects in insulin-regulated phosphorylation/dephosphorylation appear to play the predominant role in insulin resistance. Abnormalities include the accumulation of lipid within skeletal myocytes, which may impair mitochondrial oxidative phosphorylation and reduce insulin-stimulated mitochondrial ATP production. Impaired fatty acid oxidation and lipid accumulation within skeletal myocytes also may generate reactive oxygen species such as lipid peroxides. Of note, not all insulin signal transduction pathways are resistant to the effects of insulin (e.g., those controlling cell growth and differentiation using the mitogenic-activated protein kinase pathway). Consequently, hyperinsulinemia may increase the insulin action through these pathways, potentially accelerating diabetes-related conditions such as atherosclerosis.

The obesity accompanying type 2 DM, particularly in a central or visceral location, is thought to be part of the pathogenic process (Chap. 415e). In addition to these white fat depots, humans now are recognized to have brown fat, which has much greater thermogenic capacity. Efforts are under way to increase the activity or quantity of brown fat (e.g., a myokine, irisin, may convert white to brown fat). The increased adipocyte mass leads to increased levels of circulating free fatty acids and other fat cell products. For example, adipocytes secrete a number of biologic products (nonesterified free fatty acids, retinol-binding protein 4, leptin, TNF- α , resistin, IL-6, and adiponectin). In addition to regulating body weight, appetite, and energy expenditure, adipokines also modulate insulin sensitivity. The increased production of free fatty acids and some adipokines may cause insulin resistance in skeletal muscle and liver. For example, free fatty acids impair glucose utilization in skeletal muscle, promote glucose production by the liver, and impair beta cell function. In contrast, the production by adipocytes of adiponectin, an insulin-sensitizing peptide, is reduced in obesity, and this may contribute to hepatic insulin resistance. Adipocyte products and adipokines also produce an inflammatory state and may explain why markers of inflammation such as IL-6 and C-reactive protein are often elevated in type 2 DM. In addition, inflammatory cells

have been found infiltrating adipose tissue. Inhibition of inflammatory signaling pathways such as the nuclear factor- κ B (NF- κ B) pathway appears to reduce insulin resistance and improve hyperglycemia in animal models and is being tested in humans.

IMPAIRED INSULIN SECRETION Insulin secretion and sensitivity are interrelated (Fig. 417-7). In type 2 DM, insulin secretion initially increases in response to insulin resistance to maintain normal glucose tolerance. Initially, the insulin secretory defect is mild and selectively involves glucose-stimulated insulin secretion, including a greatly reduced first secretory phase. The response to other nonglucose secretagogues, such as arginine, is preserved, but overall beta function is reduced by as much as 50% at the onset of type 2 DM. Abnormalities in proinsulin processing are reflected by increased secretion of proinsulin in type 2 DM. Eventually, the insulin secretory defect is progressive.

The reason(s) for the decline in insulin secretory capacity in type 2 DM is unclear. The assumption is that a second genetic defect—superimposed upon insulin resistance—leads to beta cell failure. Beta cell mass is decreased by approximately 50% in individuals with long-standing type 2 DM. Islet amyloid polypeptide or amylin, co-secreted by the beta cell, forms the amyloid fibrillar deposit found in the islets of individuals with long-standing type 2 DM. Whether such islet amyloid deposits are a primary or secondary event is not known. The metabolic environment of diabetes may also negatively impact islet function. For example, chronic hyperglycemia paradoxically impairs islet function (“glucose toxicity”) and leads to a worsening of hyperglycemia. Improvement in glycemic control is often associated with improved islet function. In addition, elevation of free fatty acid levels (“lipotoxicity”) and dietary fat may also worsen islet function. Reduced GLP-1 action may contribute to the reduced insulin secretion.

INCREASED HEPATIC GLUCOSE AND LIPID PRODUCTION In type 2 DM, insulin resistance in the liver reflects the failure of hyperinsulinemia to suppress gluconeogenesis, which results in fasting hyperglycemia and decreased glycogen storage by the liver in the postprandial state. Increased hepatic glucose production occurs early in the course of diabetes, although likely after the onset of insulin secretory abnormalities and insulin resistance in skeletal muscle. As a result of insulin resistance in adipose tissue, lipolysis and free fatty acid flux from adipocytes are increased, leading to increased lipid (very-low-density lipoprotein [VLDL] and triglyceride) synthesis in hepatocytes. This lipid storage or steatosis in the liver may lead to nonalcoholic fatty liver disease (Chap. 367e) and abnormal liver function tests. This is also responsible for the dyslipidemia found in type 2 DM (elevated triglycerides, reduced high-density lipoprotein [HDL], and increased small dense low-density lipoprotein [LDL] particles).

Insulin Resistance Syndromes The insulin resistance condition comprises a spectrum of disorders, with hyperglycemia representing one of the most readily diagnosed features. The *metabolic syndrome*, the *insulin resistance syndrome*, and *syndrome X* are terms used to describe a constellation of metabolic derangements that includes insulin resistance, hypertension, dyslipidemia (decreased HDL and elevated triglycerides), central or visceral obesity, type 2 DM or IGT/IFG, and accelerated cardiovascular disease. This syndrome is discussed in Chap. 422.

A number of relatively rare forms of severe insulin resistance include features of type 2 DM or IGT (Table 417-1). Mutations in the insulin receptor that interfere with binding or signal transduction are a rare cause of insulin resistance. Acanthosis nigricans and signs of hyperandrogenism (hirsutism, acne, and oligomenorrhea in women) are also common physical features. Two distinct syndromes of severe insulin resistance have been described in adults: (1) type A, which affects young women and is characterized by severe hyperinsulinemia, obesity, and features of hyperandrogenism; and (2) type B, which affects middle-aged women and is characterized by severe hyperinsulinemia, features of hyperandrogenism, and autoimmune disorders. Individuals with the type A insulin resistance syndrome have an undefined defect in the insulin-signaling pathway; individuals with the type B insulin resistance syndrome have autoantibodies directed at

2406 the insulin receptor. These receptor autoantibodies may block insulin binding or may stimulate the insulin receptor, leading to intermittent hypoglycemia.

Polycystic ovary syndrome (PCOS) is a common disorder that affects premenopausal women and is characterized by chronic anovulation and hyperandrogenism (**Chap. 412**). Insulin resistance is seen in a significant subset of women with PCOS, and the disorder substantially increases the risk for type 2 DM, independent of the effects of obesity.

Prevention Type 2 DM is preceded by a period of IGT or IFG, and a number of lifestyle modifications and pharmacologic agents prevent or delay the onset of DM. Individuals with prediabetes or increased risk of diabetes should be referred to a structured program to reduce body weight and increase physical activity as well as being screened for cardiovascular disease. The Diabetes Prevention Program (DPP) demonstrated that intensive changes in lifestyle (diet and exercise for 30 min/d five times/week) in individuals with IGT prevented or delayed the development of type 2 DM by 58% compared to placebo. This effect was seen in individuals regardless of age, sex, or ethnic group. In the same study, metformin prevented or delayed diabetes by 31% compared to placebo. The lifestyle intervention group lost 5–7% of their body weight during the 3 years of the study. Studies in Finnish and Chinese populations noted similar efficacy of diet and exercise in preventing or delaying type 2 DM. A number of agents, including α -glucosidase inhibitors, metformin, thiazolidinediones, GLP-1 receptor pathway modifiers, and orlistat, prevent or delay type 2 DM but are not approved for this purpose. Individuals with a strong family history of type 2 DM and individuals with IFG or IGT should be strongly encouraged to maintain a normal BMI and engage in regular physical activity. Pharmacologic therapy for individuals with prediabetes is currently controversial because its cost-effectiveness and safety profile are not known. The ADA has suggested that metformin be considered in individuals with both IFG and IGT who are at very high risk for progression to diabetes (age <60 years, BMI \geq 35 kg/m², family history of diabetes in first-degree relative, and women with a history of GDM). Individuals with IFG, IGT, or an HbA_{1c} of 5.7–6.4% should be monitored annually to determine if diagnostic criteria for diabetes are present.

GENETICALLY DEFINED, MONOGENIC FORMS OF DIABETES MELLITUS RELATED TO REDUCED INSULIN SECRETION

Several monogenic forms of DM have been identified. More than 10 different variants of MODY, caused by mutations in genes encoding islet-enriched transcription factors or glucokinase (Fig. 417-5; Table 417-1), are transmitted as autosomal dominant disorders. MODY 1, MODY 3, and MODY 5 are caused by mutations in hepatocyte nuclear transcription factor (HNF) 4 α , HNF-1 α , and HNF-1 β , respectively. As their names imply, these transcription factors are expressed in the liver but also in other tissues, including the pancreatic islets and kidney. These factors most likely affect islet development or the expression of genes important in glucose-stimulated insulin secretion or the maintenance of beta cell mass. For example, individuals with an HNF-1 α mutation (MODY 3) have a progressive decline in glycemic control but may respond to sulfonylureas. In fact, some of these patients were initially thought to have type 1 DM but were later shown to respond to a sulfonylurea, and insulin was discontinued. Individuals with a HNF-1 β mutation have progressive impairment of insulin secretion and hepatic insulin resistance, and require insulin treatment (minimal response to sulfonylureas). These individuals often have other abnormalities such as renal cysts, mild pancreatic exocrine insufficiency, and abnormal liver function tests. Individuals with MODY 2, the result of mutations in the glucokinase gene, have mild-to-moderate, stable hyperglycemia that does not respond to oral hypoglycemic agents. Glucokinase catalyzes the formation of glucose-6-phosphate from glucose, a reaction that is important for glucose sensing by the beta cells (Fig. 417-5) and for glucose utilization by the liver. As a result of glucokinase mutations, higher glucose levels are required to elicit insulin secretory responses, thus altering the set point for insulin secretion. Studies of populations with type 2 DM suggest that mutations in MODY-associated genes are an uncommon (<5%)

cause of type 2 DM. Mutations in mitochondrial DNA are associated with diabetes and deafness.

Transient or permanent neonatal diabetes (onset <12 months of age) occurs. Permanent neonatal diabetes may be caused by several genetic mutations, usually requires treatment with insulin, and phenotypically is similar to type 1 DM. Mutations in the ATP-sensitive potassium channel subunits (Kir6.2 and ABCC8) and the insulin gene (interfere with proinsulin folding and processing) (Fig. 417-5) are the major causes of permanent neonatal diabetes. Although these activating mutations in the ATP-sensitive potassium channel subunits impair glucose-stimulated insulin secretion, these individuals may respond to sulfonylureas and can be treated with these agents. These mutations are often associated with a spectrum of neurologic dysfunction. MODY 4 is a rare variant caused by mutations in the insulin promoter factor (IPF) 1, a transcription factor that regulates pancreatic development and insulin gene transcription. Homozygous inactivating mutations cause pancreatic agenesis, whereas heterozygous mutations may result in DM. Mutations in the transcription factor of GATA6 are the most common cause of pancreatic agenesis. Homozygous glucokinase mutations cause a severe form of neonatal diabetes.

APPROACH TO THE PATIENT: Diabetes Mellitus

Once the diagnosis of DM is made, attention should be directed to symptoms related to diabetes (acute and chronic) and classifying the type of diabetes. DM and its complications produce a wide range of symptoms and signs; those secondary to acute hyperglycemia may occur at any stage of the disease, whereas those related to chronic hyperglycemia begin to appear during the second decade of hyperglycemia (**Chap. 419**). Individuals with previously undetected type 2 DM may present with chronic complications of DM at the time of diagnosis. The history and physical examination should assess for symptoms or signs of acute hyperglycemia and should screen for the chronic complications and conditions associated with DM.

HISTORY

A complete medical history should be obtained with special emphasis on DM-relevant aspects such as weight, family history of DM and its complications, risk factors for cardiovascular disease, exercise, smoking, and ethanol use. Symptoms of hyperglycemia include polyuria, polydipsia, weight loss, fatigue, weakness, blurry vision, frequent superficial infections (vaginitis, fungal skin infections), and slow healing of skin lesions after minor trauma. Metabolic derangements relate mostly to hyperglycemia (osmotic diuresis) and to the catabolic state of the patient (urinary loss of glucose and calories, muscle breakdown due to protein degradation and decreased protein synthesis). Blurred vision results from changes in the water content of the lens and resolves as the hyperglycemia is controlled.

In a patient with established DM, the initial assessment should also include special emphasis on prior diabetes care, including the type of therapy, prior HbA_{1c} levels, self-monitoring blood glucose results, frequency of hypoglycemia, presence of DM-specific complications, and assessment of the patient's knowledge about diabetes, exercise, and nutrition. Diabetes-related complications may afflict several organ systems, and an individual patient may exhibit some, all, or none of the symptoms related to the complications of DM (**Chap. 419**). In addition, the presence of DM-related comorbidities should be sought (cardiovascular disease, hypertension, dyslipidemia). Pregnancy plans should be ascertained in women of childbearing age.

PHYSICAL EXAMINATION

In addition to a complete physical examination, special attention should be given to DM-relevant aspects such as weight or BMI,

retinal examination, orthostatic blood pressure, foot examination, peripheral pulses, and insulin injection sites. Blood pressure >140/80 mmHg is considered hypertension in individuals with diabetes. Because periodontal disease is more frequent in DM, the teeth and gums should also be examined.

An annual foot examination should (1) assess blood flow, sensation (vibratory sensation [128-MHz tuning fork at the base of the great toe], the ability to sense touch with a monofilament [5.07, 10-g monofilament], pinprick sensation, testing for ankle reflexes, and vibration perception threshold using a biothesiometer), ankle reflexes, and nail care; (2) look for the presence of foot deformities such as hammer or claw toes and Charcot foot; and (3) identify sites of potential ulceration. The ADA recommends annual screening for distal symmetric neuropathy beginning with the initial diagnosis of diabetes and annual screening for autonomic neuropathy 5 years after diagnosis of type 1 DM and at the time of diagnosis of type 2 DM. This includes testing for loss of protective sensation (LOPS) using monofilament testing plus one of the following tests: vibration, pinprick, ankle reflexes, or vibration perception threshold (using a biothesiometer). If the monofilament test or one of the other tests is abnormal, the patient is diagnosed with LOPS and counseled accordingly (Chap. 419).

CLASSIFICATION OF DM IN AN INDIVIDUAL PATIENT

The etiology of diabetes in an individual with new-onset disease can usually be assigned on the basis of clinical criteria. Individuals with type 1 DM tend to have the following characteristics: (1) onset of disease prior to age 30 years; (2) lean body habitus; (3) requirement of insulin as the initial therapy; (4) propensity to develop ketoacidosis; and (5) an increased risk of other autoimmune disorders such as autoimmune thyroid disease, adrenal insufficiency, pernicious anemia, celiac disease, and vitiligo. In contrast, individuals with type 2 DM often exhibit the following features: (1) develop diabetes after the age of 30 years; (2) are usually obese (80% are obese, but elderly individuals may be lean); (3) may not require insulin therapy initially; and (4) may have associated conditions such as insulin resistance, hypertension, cardiovascular disease, dyslipidemia, or PCOS. In type 2 DM, insulin resistance is often associated with abdominal obesity (as opposed to hip and thigh obesity) and hypertriglyceridemia. Although most individuals diagnosed with type 2 DM are older, the age of diagnosis is declining, and there is a marked increase among overweight children and adolescents. Some individuals with phenotypic type 2 DM present with diabetic ketoacidosis but lack autoimmune markers and may be later treated with oral glucose-lowering agents rather than insulin (this clinical picture is sometimes referred to as *ketosis-prone type 2 DM*). On the other hand, some individuals (5–10%) with the phenotypic appearance of type 2 DM do not have absolute insulin deficiency but have autoimmune markers (GAD and other ICA autoantibodies) suggestive of type 1 DM (termed *latent autoimmune diabetes of the adult*). Such individuals are more likely to be <50 years of age, thinner, and have a personal or family history of other autoimmune disease than individuals with type 2 DM. They are much more likely to require insulin treatment within 5 years. Monogenic forms of diabetes (discussed above) should be considered in those with diabetes onset at <30 years of age, an autosomal pattern of diabetes inheritance, and the lack of nearly complete insulin deficiency. Despite recent advances in the understanding of the pathogenesis of diabetes, it remains difficult to categorize some patients unequivocally. Individuals who deviate from the clinical profile of type 1 and type 2 DM, or who have other associated defects such as deafness, pancreatic exocrine disease, and other endocrine disorders, should be classified accordingly (Table 417-1).

LABORATORY ASSESSMENT

The laboratory assessment should first determine whether the patient meets the diagnostic criteria for DM (Table 417-2) and then assess the degree of glycemic control (Chap. 418). In addition to the

standard laboratory evaluation, the patient should be screened for DM-associated conditions (e.g., albuminuria, dyslipidemia, thyroid dysfunction).

The classification of the type of DM may be facilitated by laboratory assessments. Serum insulin or C-peptide measurements often do not distinguish type 1 from type 2 DM, but a low C-peptide level confirms a patient's need for insulin. Many individuals with new-onset type 1 DM retain some C-peptide production. Measurement of islet cell antibodies at the time of diabetes onset may be useful if the type of DM is not clear based on the characteristics described above.

418 Diabetes Mellitus: Management and Therapies

Alvin C. Powers

OVERALL GOALS

The goals of therapy for type 1 or type 2 diabetes mellitus (DM) are to (1) eliminate symptoms related to hyperglycemia, (2) reduce or eliminate the long-term microvascular and macrovascular complications of DM (Chap. 419), and (3) allow the patient to achieve as normal a lifestyle as possible. To reach these goals, the physician should identify a target level of glycemic control for each patient, provide the patient with the educational and pharmacologic resources necessary to reach this level, and monitor/treat DM-related complications. Symptoms of diabetes usually resolve when the plasma glucose is <11.1 mmol/L (200 mg/dL), and thus most DM treatment focuses on achieving the second and third goals. This chapter first reviews the ongoing treatment of diabetes in the outpatient setting and then discusses the treatment of severe hyperglycemia, as well as the treatment of diabetes in hospitalized patients.

The care of an individual with either type 1 or type 2 DM requires a multidisciplinary team. Central to the success of this team are the patient's participation, input, and enthusiasm, all of which are essential for optimal diabetes management. Members of the health care team include the primary care provider and/or the endocrinologist or diabetologist, a certified diabetes educator, a nutritionist, and a psychologist. In addition, when the complications of DM arise, subspecialists (including neurologists, nephrologists, vascular surgeons, cardiologists, ophthalmologists, and podiatrists) with experience in DM-related complications are essential.

ONGOING ASPECTS OF COMPREHENSIVE DIABETES CARE

A number of names are sometimes applied to different approaches to diabetes care, such as intensive insulin therapy, intensive glycemic control, and "tight control." The current chapter, and other sources, uses the term *comprehensive diabetes care* to emphasize the fact that optimal diabetes therapy involves more than plasma glucose management and medications. Although glycemic control is central to optimal diabetes therapy, comprehensive diabetes care of both type 1 and type 2 DM should also detect and manage DM-specific complications (Chap. 419) and modify risk factors for DM-associated diseases. The key elements of comprehensive diabetes care are summarized in Table 418-1. In addition to the physical aspects of DM, social, family, financial, cultural, and employment-related issues may impact diabetes care. The International Diabetes Federation (IDF), recognizing that resources available for diabetes care vary widely throughout the world, has issued guidelines for "recommended care" (a well-developed service base and with health care funding systems consuming a significant part of their national wealth), "limited care" (health care settings with very limited resources), and "comprehensive care" (health care settings with considerable resources). This chapter provides guidance

TABLE 418-1 GUIDELINES FOR ONGOING, COMPREHENSIVE MEDICAL CARE FOR PATIENTS WITH DIABETES

- Optimal and individualized glycemic control
- Self-monitoring of blood glucose (individualized frequency)
- HbA_{1c} testing (2–4 times/year)
- Patient education in diabetes management (annual); diabetes-self management education and support
- Medical nutrition therapy and education (annual)
- Eye examination (annual or biannual; **Chap. 419**)
- Foot examination (1–2 times/year by physician; daily by patient; **Chap. 419**)
- Screening for diabetic nephropathy (annual; **Chap. 419**)
- Blood pressure measurement (quarterly)
- Lipid profile and serum creatinine (estimate GFR) (annual; **Chap. 419**)
- Influenza/pneumococcal/hepatitis B immunizations
- Consider antiplatelet therapy (**Chap. 419**)

Abbreviations: GFR, glomerular filtration rate; HbA_{1c}, hemoglobin A_{1c}.

for this comprehensive level of diabetes care. The treatment goals for patients with diabetes are summarized in **Table 418-2** and should be individualized.

DETECTION AND PREVENTION OF COMPLICATIONS RELATED TO DIABETES

The morbidity and mortality rates of DM-related complications (**Chap. 419**) can be greatly reduced by timely and consistent surveillance procedures (**Table 418-1**). These screening procedures are indicated for all individuals with DM, but many individuals with diabetes do not receive comprehensive diabetes care. A comprehensive eye examination should be performed by a qualified optometrist or ophthalmologist. Because many individuals with type 2 DM have had asymptomatic diabetes for several years before diagnosis, the American Diabetes Association (ADA) recommends the following ophthalmologic examination schedule: (1) individuals with type 1 DM should have an initial eye examination within 5 years of diagnosis, (2) individuals with type 2 DM should have an initial eye examination at the time of diabetes diagnosis, (3) women with DM who are pregnant or contemplating pregnancy should have an eye examination prior to conception and during the first trimester, and (4) if eye exam is normal, repeat examination in 2–3 years is appropriate.

TABLE 418-2 TREATMENT GOALS FOR ADULTS WITH DIABETES^a

Index	Goal
Glycemic control ^b	
HbA _{1c}	<7.0% ^c
Preprandial capillary plasma glucose	4.4–7.2 mmol/L (80–130 mg/dL)
Peak postprandial capillary plasma glucose ^d	<10.0 mmol/L (<180 mg/dL)
Blood pressure	<140/90 mmHg ^e
Lipids ^f	
Low-density lipoprotein	<2.6 mmol/L (100 mg/dL) ^g
High-density lipoprotein	>1 mmol/L (40 mg/dL) in men >1.3 mmol/L (50 mg/dL) in women
Triglycerides	<1.7 mmol/L (150 mg/dL)

^aAs recommended by the American Diabetes Association; goals should be individualized for each patient (see text). Goals may be different for certain patient populations. ^bHbA_{1c} is primary goal. ^cDiabetes Control and Complications Trial–based assay. ^d1–2 h after beginning of a meal. ^eGoal of <130/80 mmHg may be appropriate for younger individuals. ^fIn decreasing order of priority. Recent guidelines from the American College of Cardiology and American Heart Association no longer advocate specific LDL and HDL goals (see **Chaps. 291e and 419**). ^gGoal of <1.8 mmol/L (70 mg/dL) may be appropriate for individuals with cardiovascular disease.

Abbreviation: HbA_{1c}, hemoglobin A_{1c}.

Source: Adapted from American Diabetes Association: *Diabetes Care* 38(Suppl 1):S1, 2015.

PATIENT EDUCATION ABOUT DM, NUTRITION, AND EXERCISE

The patient with type 1 or type 2 DM should receive education about nutrition, exercise, care of diabetes during illness, and medications to lower the plasma glucose. Along with improved compliance, patient education allows individuals with DM to assume greater responsibility for their care. Patient education should be viewed as a continuing process with regular visits for reinforcement; it should not be a process that is completed after one or two visits to a nurse educator or nutritionist. The ADA refers to education about the individualized management plan for the patient as diabetes self-management education (DSME) and diabetes self-management support (DSMS). DSME and DSMS are ways to improve the patient's knowledge, skills, and abilities necessary for diabetes self-care and should also emphasize psychosocial issues and emotional well-being. More frequent contact between the patient and the diabetes management team (e.g., electronic, telephone) improves glycemic control.

Diabetes Education The diabetes educator is a health care professional (nurse, dietician, or pharmacist) with specialized patient education skills who is certified in diabetes education (e.g., American Association of Diabetes Educators). Education topics important for optimal diabetes care include self-monitoring of blood glucose; urine ketone monitoring (type 1 DM); insulin administration; guidelines for diabetes management during illnesses; prevention and management of hypoglycemia (**Chap. 420**); foot and skin care; diabetes management before, during, and after exercise; and risk factor–modifying activities.

Psychosocial Aspects Because the individual with DM can face challenges that affect many aspects of daily life, psychosocial assessment and treatment are a critical part of providing comprehensive diabetes care. The individual with DM must accept that he or she may develop complications related to DM. Even with considerable effort, normoglycemia can be an elusive goal, and solutions to worsening glycemic control may not be easily identifiable. The patient should view him- or herself as an essential member of the diabetes care team and not as someone who is cared for by the diabetes management team. Emotional stress may provoke a change in behavior so that individuals no longer adhere to a dietary, exercise, or therapeutic regimen. This can lead to the appearance of either hyper- or hypoglycemia. Eating disorders, including binge eating disorders, bulimia, and anorexia nervosa, appear to occur more frequently in individuals with type 1 or type 2 DM.

Nutrition *Medical nutrition therapy* (MNT) is a term used by the ADA to describe the optimal coordination of caloric intake with other aspects of diabetes therapy (insulin, exercise, weight loss). Primary prevention measures of MNT are directed at preventing or delaying the onset of type 2 DM in high-risk individuals (obese or with prediabetes) by promoting weight reduction. Medical treatment of obesity is a rapidly evolving area and is discussed in **Chap. 416**. Secondary prevention measures of MNT are directed at preventing or delaying diabetes-related complications in diabetic individuals by improving glycemic control. Tertiary prevention measures of MNT are directed at managing diabetes-related complications (cardiovascular disease, nephropathy) in diabetic individuals. MNT in patients with diabetes and cardiovascular disease should incorporate dietary principles used in nondiabetic patients with cardiovascular disease. Although the recommendations for all three types of MNT overlap, this chapter emphasizes secondary prevention measures of MNT. Pharmacologic approaches that facilitate weight loss and bariatric surgery should be considered in selected patients (**Chaps. 415e and 416**).

In general, the components of optimal MNT are similar for individuals with type 1 or type 2 DM and similar to those for the general population (fruits, vegetables, fiber-containing foods, and low fat; **Table 418-3**). MNT education is an important component of comprehensive diabetes care and should be reinforced by regular patient education. Historically, nutrition education imposed restrictive, complicated regimens on the patient. Current practices have greatly changed, although many patients and health care providers still view the diabetic diet as monolithic and static. For example, MNT now includes foods

TABLE 418-3 NUTRITIONAL RECOMMENDATIONS FOR ADULTS WITH DIABETES OR PREDIABETES^a

Weight loss diet (in prediabetes and type 2 DM)
• Hypocaloric diet that is low-carbohydrate
Fat in diet (optimal % of diet is not known; should be individualized)
• Minimal <i>trans</i> fat consumption
• Mediterranean-style diet rich in monounsaturated fatty acids may be better
Carbohydrate in diet (optimal % of diet is not known; should be individualized)
• Monitor carbohydrate intake in regard to calories
• Sucrose-containing foods may be consumed with adjustments in insulin dose, but minimize intake
• Amount of carbohydrate determined by estimating grams of carbohydrate in diet (type 1 DM)
• Use glycemic index to predict how consumption of a particular food may affect blood glucose
• Fructose preferred over sucrose or starch
Protein in diet (optimal % of diet is not known; should be individualized)
Other components
• Dietary fiber, vegetable, fruits, whole grains, dairy products, and sodium intake as advised for general population
• Nonnutrient sweeteners
• Routine supplements of vitamins, antioxidants, or trace elements not advised

^aSee text for differences for patients with type 1 or type 2 diabetes.

Source: Adapted from American Diabetes Association: *Diabetes Care* 37(Suppl 1):S14, 2014.

with sucrose and seeks to modify other risk factors such as hyperlipidemia and hypertension rather than focusing exclusively on weight loss in individuals with type 2 DM. The *glycemic index* is an estimate of the postprandial rise in the blood glucose when a certain amount of that food is consumed. Consumption of foods with a low glycemic index appears to reduce postprandial glucose excursions and improve glycemic control. Reduced-calorie and nonnutritive sweeteners are useful. Currently, evidence does not support supplementation of the diet with vitamins, antioxidants (vitamin C and E), or micronutrients (chromium) in patients with diabetes.

The goal of MNT in the individual with type 1 DM is to coordinate and match the caloric intake, both temporally and quantitatively, with the appropriate amount of insulin. MNT in type 1 DM and self-monitoring of blood glucose must be integrated to define the optimal insulin regimen. The ADA encourages patients and providers to use carbohydrate counting or exchange systems to estimate the nutrient content of a meal or snack. Based on the patient's estimate of the carbohydrate content of a meal, an insulin-to-carbohydrate ratio determines the bolus insulin dose for a meal or snack. MNT must be flexible enough to allow for exercise, and the insulin regimen must allow for deviations in caloric intake. An important component of MNT in type 1 DM is to minimize the weight gain often associated with intensive diabetes management.

The goals of MNT in type 2 DM should focus on weight loss and address the greatly increased prevalence of cardiovascular risk factors (hypertension, dyslipidemia, obesity) and disease in this population. The majority of these individuals are obese, and weight loss is strongly encouraged and should remain an important goal. Hypocaloric diets and modest weight loss (5–7%) often result in rapid and dramatic glucose lowering in individuals with new-onset type 2 DM. Nevertheless, numerous studies document that long-term weight loss is uncommon. MNT for type 2 DM should emphasize modest caloric reduction (low-carbohydrate) and increased physical activity. Increased consumption of soluble, dietary fiber may improve glycemic control in individuals with type 2 DM. Weight loss and exercise improve insulin resistance.

Exercise Exercise has multiple positive benefits including cardiovascular risk reduction, reduced blood pressure, maintenance of muscle

mass, reduction in body fat, and weight loss. For individuals with type 1 or type 2 DM, exercise is also useful for lowering plasma glucose (during and following exercise) and increasing insulin sensitivity. In patients with diabetes, the ADA recommends 150 min/week (distributed over at least 3 days) of moderate aerobic physical activity with no gaps longer than 2 days. The exercise regimen should also include resistance training.

Despite its benefits, exercise presents challenges for individuals with DM because they lack the normal glucoregulatory mechanisms (normally, insulin falls and glucagon rises during exercise). Skeletal muscle is a major site for metabolic fuel consumption in the resting state, and the increased muscle activity during vigorous, aerobic exercise greatly increases fuel requirements. Individuals with type 1 DM are prone to either hyperglycemia or hypoglycemia during exercise, depending on the preexercise plasma glucose, the circulating insulin level, and the level of exercise-induced catecholamines. If the insulin level is too low, the rise in catecholamines may increase the plasma glucose excessively, promote ketone body formation, and possibly lead to ketoacidosis. Conversely, if the circulating insulin level is excessive, this relative hyperinsulinemia may reduce hepatic glucose production (decreased glycogenolysis, decreased gluconeogenesis) and increase glucose entry into muscle, leading to hypoglycemia.

To avoid exercise-related hyper- or hypoglycemia, individuals with type 1 DM should (1) monitor blood glucose before, during, and after exercise; (2) delay exercise if blood glucose is >14 mmol/L (250 mg/dL) and ketones are present; (3) if the blood glucose is <5.6 mmol/L (100 mg/dL), ingest carbohydrate before exercising; (4) monitor glucose during exercise and ingest carbohydrate to prevent hypoglycemia; (5) decrease insulin doses (based on previous experience) before exercise and inject insulin into a nonexercising area; and (6) learn individual glucose responses to different types of exercise and increase food intake for up to 24 h after exercise, depending on intensity and duration of exercise. In individuals with type 2 DM, exercise-related hypoglycemia is less common but can occur in individuals taking either insulin or insulin secretagogues.

Despite asymptomatic cardiovascular disease appearing at a younger age in both type 1 and type 2 DM, routine screening for coronary artery disease has not been shown to be effective and is not recommended (Chap. 419). Untreated proliferative retinopathy is a relative contraindication to vigorous exercise, because this may lead to vitreous hemorrhage or retinal detachment.

MONITORING THE LEVEL OF GLYCEMIC CONTROL

Optimal monitoring of glycemic control involves plasma glucose measurements by the patient and an assessment of long-term control by the physician (measurement of hemoglobin A_{1c} [HbA_{1c}] and review of the patient's self-measurements of plasma glucose). These measurements are complementary: the patient's measurements provide a picture of short-term glycemic control, whereas the HbA_{1c} reflects average glycemic control over the previous 2–3 months.

Self-Monitoring of Blood Glucose Self-monitoring of blood glucose (SMBG) is the standard of care in diabetes management and allows the patient to monitor his or her blood glucose at any time. In SMBG, a small drop of blood and an easily detectable enzymatic reaction allow measurement of the capillary plasma glucose. Many glucose monitors can rapidly and accurately measure glucose (calibrated to provide plasma glucose value even though blood glucose is measured) in small amounts of blood (3–10 μL) obtained from the fingertip; alternative testing sites (e.g., forearm) are less reliable, especially when the blood glucose is changing rapidly (postprandially). A large number of blood glucose monitors are available, and the certified diabetes educator is critical in helping the patient select the optimal device and learn to use it properly. By combining glucose measurements with diet history, medication changes, and exercise history, the diabetes management team and patient can improve the treatment program.

The frequency of SMBG measurements must be individualized and adapted to address the goals of diabetes care. Individuals with type 1

2410 DM or individuals with type 2 DM taking multiple insulin injections each day should routinely measure their plasma glucose three or more times per day to estimate and select mealtime boluses of short-acting insulin and to modify long-acting insulin doses. Most individuals with type 2 DM require less frequent monitoring, although the optimal frequency of SMBG has not been clearly defined. Individuals with type 2 DM who are taking insulin should use SMBG more frequently than those on oral agents. Individuals with type 2 DM who are on oral medications should use SMBG as a means of assessing the efficacy of their medication and the impact of diet. Because plasma glucose levels fluctuate less in these individuals, one to two SMBG measurements per day (or fewer in patients who are on oral agents or are diet-controlled) may be sufficient. Most measurements in individuals with type 1 or type 2 DM should be performed prior to a meal and supplemented with postprandial measurements to assist in reaching postprandial glucose targets (Table 418-2).

Devices for continuous glucose monitoring (CGM) have been approved by the U.S. Food and Drug Administration (FDA), and others are in various stages of development. These devices do not replace the need for traditional glucose measurements and require calibration with SMBG. This rapidly evolving technology requires substantial expertise on the part of the diabetes management team and the patient. Current CGM systems measure the glucose in interstitial fluid, which is in equilibrium with the blood glucose. These devices provide useful short-term information about the patterns of glucose changes as well as an enhanced ability to detect hypoglycemic episodes. Alarms notify the patient if the blood glucose falls into the hypoglycemic range. Clinical experience with these devices is rapidly growing, and they are most useful in individuals with hypoglycemia unawareness, individuals with frequent hypoglycemia, or those who have not achieved glycemic targets despite major efforts. The utility of CGM in the intensive care unit (ICU) setting remains to be determined.

Assessment of Long-Term Glycemic Control Measurement of glycated hemoglobin (HbA_{1c}) is the standard method for assessing long-term glycemic control. When plasma glucose is consistently elevated, there is an increase in nonenzymatic glycation of hemoglobin; this alteration reflects the glycemic history over the previous 2–3 months, because erythrocytes have an average life span of 120 days (glycemic level in the preceding month contributes about 50% to the HbA_{1c} value). Measurement of HbA_{1c} at the “point of care” allows for more rapid feedback and may therefore assist in adjustment of therapy.

HbA_{1c} should be measured in all individuals with DM during their initial evaluation and as part of their comprehensive diabetes care. As the primary predictor of long-term complications of DM, the HbA_{1c} should mirror, to a certain extent, the short-term measurements of SMBG. These two measurements are complementary in that recent intercurrent illnesses may impact the SMBG measurements but not the HbA_{1c}. Likewise, postprandial and nocturnal hyperglycemia may not be detected by the SMBG of fasting and preprandial capillary plasma glucose but will be reflected in the HbA_{1c}. In standardized assays, the HbA_{1c} approximates the following mean plasma glucose values: an HbA_{1c} of 6% = 7.0 mmol/L (126 mg/dL), 7% = 8.6 mmol/L (154 mg/dL), 8% = 10.2 mmol/L (183 mg/dL), 9% = 11.8 mmol/L (212 mg/dL), 10% = 13.4 mmol/L (240 mg/dL), 11% = 14.9 mmol/L (269 mg/dL), and 12% = 16.5 mmol/L (298 mg/dL). In patients achieving their glycemic goal, the ADA recommends measurement of the HbA_{1c} at least twice per year. More frequent testing (every 3 months) is warranted when glycemic control is inadequate or when therapy has changed. Laboratory standards for the HbA_{1c} test have been established and should be correlated to the reference assay of the Diabetes Control and Complications Trial (DCCT). Clinical conditions such as hemoglobinopathies, anemias, reticulocytosis, transfusions, and uremia may interfere with the HbA_{1c} result. The degree of glycation of other proteins, such as albumin, can be used as an alternative indicator of glycemic control when the HbA_{1c} is inaccurate. The fructosamine assay (measuring glycated albumin) reflects the glycemic status over the prior 2 weeks.

PHARMACOLOGIC TREATMENT OF DIABETES

Comprehensive care of type 1 and type 2 DM requires an emphasis on nutrition, exercise, and monitoring of glycemic control but also usually involves glucose-lowering medication(s). This chapter discusses classes of such medications but does not describe every glucose-lowering agent available worldwide. The initial step is to select an individualized, glycemic goal for the patient.

ESTABLISHMENT OF TARGET LEVEL OF GLYCEMIC CONTROL

Because the complications of DM are related to glycemic control, normoglycemia or near-normoglycemia is the desired, but often elusive, goal for most patients. Normalization or near-normalization of the plasma glucose for long periods of time is extremely difficult, as demonstrated by the DCCT and United Kingdom Prospective Diabetes Study (UKPDS). Regardless of the level of hyperglycemia, improvement in glycemic control will lower the risk of diabetes-specific complications (**Chap. 419**).

The target for glycemic control (as reflected by the HbA_{1c}) must be individualized, and the goals of therapy should be developed in consultation with the patient after considering a number of medical, social, and lifestyle issues. The ADA calls this a *patient-centered approach*, and other organizations such as the IDF and American Association of Clinical Endocrinologists (AACE) also suggest an individualized glycemic goal. Important factors to consider include the patient's age and ability to understand and implement a complex treatment regimen, presence and severity of complications of diabetes, known cardiovascular disease (CVD), ability to recognize hypoglycemic symptoms, presence of other medical conditions or treatments that might affect survival or the response to therapy, lifestyle and occupation (e.g., possible consequences of experiencing hypoglycemia on the job), and level of support available from family and friends.

In general, the ADA suggests that the goal is to achieve an HbA_{1c} as close to normal as possible without significant hypoglycemia. In most individuals, the target HbA_{1c} should be <7% (Table 418-2) with a more stringent target for some patients. For instance, the HbA_{1c} goal in a young adult with type 1 DM may be 6.5%. A higher HbA_{1c} goal may be appropriate for the very young or old or in individuals with limited life span or comorbid conditions. For example, an appropriate HbA_{1c} goal in elderly individuals with multiple, chronic illnesses and impaired activities of daily living might be 8.0 or 8.5%. A major consideration is the frequency and severity of hypoglycemia, because this becomes more common with a more stringent HbA_{1c} goal.

More stringent glycemic control (HbA_{1c} of ≤6%) is not beneficial, and may be detrimental, in patients with type 2 DM and a high risk of CVD. Large clinical trials (UKPDS, Action to Control Cardiovascular Risk in Diabetes [ACCORD], Action in Diabetes and Vascular Disease: Preterax and Diamicon MR Controlled Evaluation [ADVANCE], Veterans Affairs Diabetes Trial [VADT]; **Chap. 419**) have examined glycemic control in type 2 DM in individuals with low risk of CVD, with high risk of CVD, or with established CVD and have found that more intense glycemic control is not beneficial and, in some patient populations, may have a negative impact on some outcomes. These divergent outcomes stress the need for individualized glycemic goals based on the following general guidelines: (1) early in the course of type 2 diabetes when the CVD risk is lower, improved glycemic control likely leads to improved cardiovascular outcome, but this benefit occurs more than a decade after the period of improved glycemic control; (2) intense glycemic control in individuals with established CVD or at high risk for CVD is not advantageous, and may be deleterious, over a follow-up of 3–5 years; an HbA_{1c} goal <7.0% is not appropriate in this population; (3) hypoglycemia in such high-risk populations (elderly, CVD) should be avoided; and (4) improved glycemic control reduces microvascular complications of diabetes (**Chap. 419**) even if it does not improve macrovascular complications like CVD.

TYPE 1 DIABETES MELLITUS

General Aspects The ADA recommendations for fasting and bedtime glycemic goals and HbA_{1c} targets are summarized in Table 418-2.

The goal is to design and implement insulin regimens that mimic physiologic insulin secretion. Because individuals with type 1 DM partially or completely lack endogenous insulin production, administration of basal insulin is essential for regulating glycogen breakdown, gluconeogenesis, lipolysis, and ketogenesis. Likewise, insulin replacement for meals should be appropriate for the carbohydrate intake and promote normal glucose utilization and storage.

Intensive Management Intensive diabetes management has the goal of achieving euglycemia or near-normal glycemia. This approach requires multiple resources, including thorough and continuing patient education, comprehensive recording of plasma glucose measurements and nutrition intake by the patient, and a variable insulin regimen that matches glucose intake and insulin dose. Insulin regimens usually include multiple-component insulin regimens, multiple daily injections (MDIs), or insulin infusion devices (each discussed below).

The benefits of intensive diabetes management and improved glycemic control include a reduction in the microvascular complications of DM and a reduction in diabetes-related complications. From a psychological standpoint, the patient experiences greater control over his or her diabetes and often notes an improved sense of well-being, greater flexibility in the timing and content of meals, and the capability to alter insulin dosing with exercise. In addition, intensive diabetes management prior to and during pregnancy reduces the risk of fetal malformations and morbidity. Intensive diabetes management is encouraged in newly diagnosed patients with type 1 DM because it may prolong the period of C-peptide production, which may result in better glycemic control and a reduced risk of serious hypoglycemia. Although intensive management confers impressive benefits, it is also accompanied by significant personal and financial costs and is therefore not appropriate for all individuals.

Insulin Preparations Current insulin preparations are generated by recombinant DNA technology and consist of the amino acid sequence of human insulin or variations thereof. In the United States, most insulin is formulated as U-100 (100 units/mL). Regular insulin formulated as U-500 (500 units/mL) is available and sometimes useful in patients with severe insulin resistance. Human insulin has been formulated with distinctive pharmacokinetics or genetically modified to more closely mimic physiologic insulin secretion. Insulins can be classified as short-acting or long-acting (Table 418-4). For example, one short-acting insulin formulation, insulin lispro, is an insulin analogue in which the 28th and 29th amino acids (lysine and proline) on the insulin B chain have been reversed by recombinant DNA technology. Insulin aspart and insulin glulisine are genetically modified insulin analogues with properties similar to lispro. All three of the insulin analogues have full biologic activity but less tendency for self-aggregation, resulting in more rapid absorption and onset of action and a shorter duration of action. These characteristics are particularly advantageous for allowing entrainment of insulin injection and action to rising plasma glucose levels following meals. The shorter duration of action also appears to be associated with a decreased number of hypoglycemic episodes, primarily because the decay of insulin action corresponds to the decline in plasma glucose after a meal. Thus, insulin aspart, lispro, or glulisine is preferred over regular insulin for prandial coverage. Insulin glargine is a long-acting biosynthetic human insulin that differs from normal insulin in that asparagine is replaced by glycine at amino acid 21, and two arginine residues are added to the C terminus of the B chain. Compared to neutral protamine Hagedorn (NPH) insulin, the onset of insulin glargine action is later, the duration of action is longer (~24 h), and there is a less pronounced peak. A lower incidence of hypoglycemia, especially at night, has been reported with insulin glargine when compared to NPH insulin. The most recent evidence does not support an association between glargine and increased cancer risk. Insulin detemir has a fatty acid side chain that prolongs its action by slowing absorption and catabolism. Twice-daily injections of glargine or detemir are sometimes required to provide 24-h coverage. Regular and NPH insulin have the native insulin amino acid sequence.

Basal insulin requirements are provided by long-acting (NPH insulin, insulin glargine, or insulin detemir) insulin formulations. These

TABLE 418-4 PROPERTIES OF INSULIN PREPARATIONS^a

Preparation	Time of Action		
	Onset, h	Peak, h	Effective Duration, h
Short-acting			
Aspart	<0.25	0.5–1.5	2–4
Glulisine	<0.25	0.5–1.5	2–4
Lispro	<0.25	0.5–1.5	2–4
Regular	0.5–1.0	2–3	3–6
Long-acting			
Detemir	1–4	— ^b	12–24 ^c
Glargine	2–4	— ^b	20–24
NPH	2–4	4–10	10–16
Insulin combinations ^d			
75/25–75% protamine lispro, 25% lispro	<0.25	Dual ^e	10–16
70/30–70% protamine aspart, 30% aspart	<0.25	Dual ^e	15–18
50/50–50% protamine lispro, 50% lispro	<0.25	Dual ^e	10–16
70/30–70% NPH, 30% regular	0.5–1	Dual ^e	10–16

^aInsulin preparations available in the United States; others are available in the United Kingdom and Europe. ^bGlargine and detemir have minimal peak activity. ^cDuration is dose-dependent (shorter at lower doses). ^dOther insulin combinations are available. ^eDual: two peaks—one at 2–3 h and the second one several hours later.

Source: Adapted from FR Kaufman: *Medical Management of Type 1 Diabetes*, 6th edition. Alexandria, VA: American Diabetes Association, 2012.

are usually prescribed with short-acting insulin in an attempt to mimic physiologic insulin release with meals. Although mixing of NPH and short-acting insulin formulations is common practice, this mixing may alter the insulin absorption profile (especially the short-acting insulins). For example, lispro absorption is delayed by mixing with NPH. The alteration in insulin absorption when the patient mixes different insulin formulations should not prevent mixing insulins. However, the following guidelines should be followed: (1) mix the different insulin formulations in the syringe immediately before injection (inject within 2 min after mixing); (2) do not store insulin as a mixture; (3) follow the same routine in terms of insulin mixing and administration to standardize the physiologic response to injected insulin; and (4) do not mix insulin glargine or detemir with other insulins. The miscibility of some insulins allows for the production of combination insulins that contain 70% NPH and 30% regular (70/30), or equal mixtures of NPH and regular (50/50). By including the insulin analogue mixed with protamine, several combinations have a short-acting and long-acting profile (Table 418-4). Although more convenient for the patient (only two injections/day), combination insulin formulations do not allow independent adjustment of short-acting and long-acting activity. Several insulin formulations are available as insulin “pens,” which may be more convenient for some patients. Insulin delivery by inhalation has recently been approved but is not yet available. Other insulins, such as one with a duration of action of several days, are under development but are not currently available in the United States.

Insulin Regimens Representations of the various insulin regimens that may be used in type 1 DM are illustrated in Fig. 418-1. Although the insulin profiles are depicted as “smooth,” symmetric curves, there is considerable patient-to-patient variation in the peak and duration. In all regimens, long-acting insulins (NPH, glargine, or detemir) supply basal insulin, whereas regular, insulin aspart, glulisine, or lispro insulin provides prandial insulin. Short-acting insulin analogues should be injected just before (<10 min) or just after a meal; regular insulin is given 30–45 min prior to a meal. Sometimes short-acting insulin analogues are injected just after a meal (gastroparesis, unpredictable food intake).

A shortcoming of current insulin regimens is that injected insulin immediately enters the systemic circulation, whereas endogenous

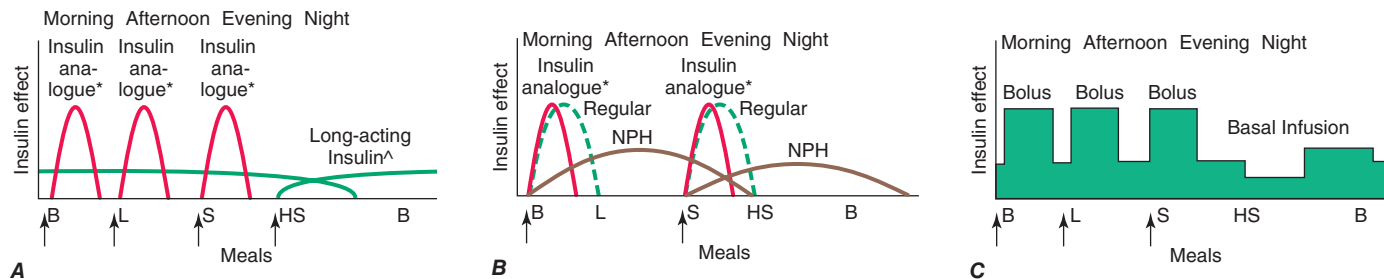


FIGURE 418-1 Representative insulin regimens for the treatment of diabetes. For each panel, the y-axis shows the amount of insulin effect and the x-axis shows the time of day. B, breakfast; HS, bedtime; L, lunch; S, supper. *Lispro, glulisine, or insulin aspart can be used. The time of insulin injection is shown with a vertical arrow. The type of insulin is noted above each insulin curve. **A.** Multiple-component insulin regimen consisting of long-acting insulin (e.g. glargine or detemir) to provide basal insulin coverage and three shots of glulisine, lispro, or insulin aspart to provide glycemic coverage for each meal. **B.** Injection of two shots of long-acting insulin (NPH) and short-acting insulin analogue (glulisine, lispro, insulin aspart [solid red line], or regular insulin [green dashed line]). Only one formulation of short-acting insulin is used. **C.** Insulin administration by insulin infusion device is shown with the basal insulin and a bolus injection at each meal. The basal insulin rate is decreased during the evening and increased slightly prior to the patient awakening in the morning. Glulisine, lispro, or insulin aspart is used in the insulin pump. (Adapted from H Lebovitz [ed]: *Therapy for Diabetes Mellitus*. American Diabetes Association, Alexandria, VA, 2004.)

insulin is secreted into the portal venous system. Thus, exogenous insulin administration exposes the liver to subphysiologic insulin levels. No insulin regimen reproduces the precise insulin secretory pattern of the pancreatic islet. However, the most physiologic regimens entail more frequent insulin injections, greater reliance on short-acting insulin, and more frequent capillary plasma glucose measurements. In general, individuals with type 1 DM require 0.5–1 U/kg per day of insulin divided into multiple doses, with ~50% of the insulin given as basal insulin.

Multiple-component insulin regimens refer to the combination of basal insulin and bolus insulin (preprandial short-acting insulin). The timing and dose of short-acting, preprandial insulin are altered to accommodate the SMBG results, anticipated food intake, and physical activity. Such regimens offer the patient with type 1 diabetes more flexibility in terms of lifestyle and the best chance for achieving near normoglycemia. One such regimen, shown in Fig. 418-1B, consists of basal insulin with glargine or detemir and preprandial lispro, glulisine, or insulin aspart. The insulin aspart, glulisine, or lispro dose is based on individualized algorithms that integrate the preprandial glucose and the anticipated carbohydrate intake. To determine the meal component of the preprandial insulin dose, the patient uses an insulin-to-carbohydrate ratio (a common ratio for type 1 DM is 1–1.5 units/10 g of carbohydrate, but this must be determined for each individual). To this insulin dose is added the supplemental or correcting insulin based on the preprandial blood glucose (one formula uses 1 unit of insulin for every 2.7 mmol/L [50 mg/dL] over the preprandial glucose target; another formula uses $[\text{body weight in kg}] \times [\text{blood glucose} - \text{desired glucose in mg/dL}] / 1500$). An alternative multiple-component insulin regimen consists of bedtime NPH insulin, a small dose of NPH insulin at breakfast (20–30% of bedtime dose), and preprandial short-acting insulin. Other variations of this regimen are in use but have the disadvantage that NPH has a significant peak, making hypoglycemia more common. Frequent SMBG (more than three times per day) is absolutely essential for these types of insulin regimens.

In the past, one commonly used regimen consisted of twice-daily injections of NPH mixed with a short-acting insulin before the morning and evening meals (Fig. 418-1B). Such regimens usually prescribe two-thirds of the total daily insulin dose in the morning (with about two-thirds given as long-acting insulin and one-third as short-acting) and one-third before the evening meal (with approximately one-half given as long-acting insulin and one-half as short-acting). The drawback to such a regimen is that it forces a rigid schedule on the patient, in terms of daily activity and the content and timing of meals. Although it is simple and effective at avoiding severe hyperglycemia, it does not generate near-normal glycemic control in individuals with type 1 DM. Moreover, if the patient's meal pattern or content varies or if physical activity is increased, hyperglycemia or hypoglycemia may result. Moving the long-acting insulin from before the evening

meal to bedtime may avoid nocturnal hypoglycemia and provide more insulin as glucose levels rise in the early morning (so-called dawn phenomenon). The insulin dose in such regimens should be adjusted based on SMBG results with the following general assumptions: (1) the fasting glucose is primarily determined by the prior evening long-acting insulin; (2) the pre-lunch glucose is a function of the morning short-acting insulin; (3) the pre-supper glucose is a function of the morning long-acting insulin; and (4) the bedtime glucose is a function of the pre-supper, short-acting insulin. This is not an optimal regimen for the patient with type 1 DM, but is sometimes used for patients with type 2 DM.

Continuous SC insulin infusion (CSII) is a very effective insulin regimen for the patient with type 1 DM (Fig. 418-1C). To the basal insulin infusion, a preprandial insulin (“bolus”) is delivered by the insulin infusion device based on instructions from the patient, who uses an individualized algorithm incorporating the preprandial plasma glucose and anticipated carbohydrate intake. These sophisticated insulin infusion devices can accurately deliver small doses of insulin (microliters per hour) and have several advantages: (1) multiple basal infusion rates can be programmed to accommodate nocturnal versus daytime basal insulin requirement; (2) basal infusion rates can be altered during periods of exercise; (3) different waveforms of insulin infusion with meal-related bolus allow better matching of insulin depending on meal composition; and (4) programmed algorithms consider prior insulin administration and blood glucose values in calculating the insulin dose. These devices require instruction by a health professional with considerable experience with insulin-infusion devices and very frequent patient interactions with the diabetes management team. Insulin-infusion devices present unique challenges, such as infection at the infusion site, unexplained hyperglycemia because the infusion set becomes obstructed, or diabetic ketoacidosis if the pump becomes disconnected. Because most physicians use lispro, glulisine, or insulin aspart in CSII, the extremely short half-life of these insulins quickly leads to insulin deficiency if the delivery system is interrupted. Essential to the safe use of infusion devices is thorough patient education about pump function and frequent SMBG. Efforts to create a closed-loop system in which data from continuous glucose measurement regulate the insulin infusion rate are under way.

Other Agents That Improve Glucose Control The role of amylin, a 37-amino-acid peptide co-secreted with insulin from pancreatic beta cells, in normal glucose homeostasis is uncertain. However, based on the rationale that patients who are insulin deficient are also amylin deficient, an analogue of amylin (pramlintide) was created and found to reduce postprandial glycemic excursions in type 1 and type 2 diabetic patients taking insulin. Pramlintide injected just before a meal slows gastric emptying and suppresses glucagon but does not alter insulin levels. Pramlintide is approved for insulin-treated patients

with type 1 and type 2 DM. Addition of pramlintide produces a modest reduction in the HbA_{1c} and seems to dampen meal-related glucose excursions. In type 1 DM, pramlintide is started as a 15- μ g SC injection before each meal and titrated up to a maximum of 30–60 μ g as tolerated. In type 2 DM, pramlintide is started as a 60- μ g SC injection before each meal and may be titrated up to a maximum of 120 μ g. The major side effects are nausea and vomiting, and dose escalations should be slow to limit these side effects. Because pramlintide slows gastric emptying, it may influence absorption of other medications and should not be used in combination with other drugs that slow GI motility. The short-acting insulin given before the meal should initially be reduced to avoid hypoglycemia and then titrated as the effects of the pramlintide become evident. α -Glucosidase inhibitors are sometimes used with insulin in type 1 DM.

TYPE 2 DIABETES MELLITUS

General Aspects The goals of glycemia-controlling therapy for type 2 DM are similar to those in type 1 DM. Whereas glycemic control tends to dominate the management of type 1 DM, the care of individuals with type 2 DM must also include attention to the treatment of conditions associated with type 2 DM (e.g., obesity, hypertension, dyslipidemia, CVD) and detection/management of DM-related complications (Fig. 418-2). Reduction in cardiovascular risk is of paramount importance because this is the leading cause of mortality in these individuals.

Type 2 DM management should begin with MNT (discussed above). An exercise regimen to increase insulin sensitivity and promote weight loss should also be instituted. Pharmacologic approaches to the management of type 2 DM include oral glucose-lowering agents, insulin, and other agents that improve glucose control; most physicians and patients prefer oral glucose-lowering agents as the initial choice. Any therapy that improves glycemic control reduces “glucose toxicity” to beta cells and improves endogenous insulin secretion. However, type 2 DM is a progressive disorder and ultimately requires multiple therapeutic agents and often insulin in most patients.

Glucose-Lowering Agents Advances in the therapy of type 2 DM have generated oral glucose-lowering agents that target different pathophysiologic processes in type 2 DM. Based on their mechanisms of action, glucose-lowering agents are subdivided into agents that increase insulin secretion, reduce glucose production, increase insulin sensitivity, enhance GLP-1 action, or promote urinary excretion of glucose (Table 418-5). Glucose-lowering agents other than insulin (with the exception of amylin analogue and α -glucosidase inhibitors) are ineffective in type 1 DM and should not be used for glucose management of severely ill individuals with type 2 DM. Insulin is sometimes the initial glucose-lowering agent in type 2 DM.

BIGUANIDES Metformin, representative of this class of agents, reduces hepatic glucose production and improves peripheral glucose utilization slightly (Table 418-5). Metformin activates AMP-dependent protein kinase and enters cells through organic cation transporters

(polymorphisms of these may influence the response to metformin). Recent evidence indicates that metformin’s mechanism for reducing hepatic glucose production is to antagonize glucagon’s ability to generate cAMP in hepatocytes. Metformin reduces fasting plasma glucose (FPG) and insulin levels, improves the lipid profile, and promotes modest weight loss. An extended-release form is available and may have fewer gastrointestinal side effects (diarrhea, anorexia, nausea, metallic taste). Because of its relatively slow onset of action and gastrointestinal symptoms with higher doses, the initial dose should be low and then escalated every 2–3 weeks based on SMBG measurements. Metformin is effective as monotherapy and can be used in combination with other oral agents or with insulin. The major toxicity of metformin, lactic acidosis, is very rare and can be prevented by careful patient selection. Vitamin B₁₂ levels are ~30% lower during metformin treatment. Metformin should not be used in patients with renal insufficiency (glomerular filtration rate [GFR] <60 mL/min), any form of acidosis, unstable congestive heart failure (CHF), liver disease, or severe hypoxemia. Some feel that these guidelines are too restrictive and prevent individuals with mild to moderate renal impairment from being safely treated with metformin. The National Institute for Health and Clinical Excellence in the United Kingdom suggests that metformin be used at a GFR >30 mL/min, with a reduced dose when the GFR is <45 mL/min. Metformin should be discontinued in hospitalized patients, in patients who can take nothing orally, and in those receiving radiographic contrast material. Insulin should be used until metformin can be restarted.

INSULIN SECRETAGOGUES—AGENTS THAT AFFECT THE ATP-SENSITIVE K⁺ CHANNEL Insulin secretagogues stimulate insulin secretion by interacting with the ATP-sensitive potassium channel on the beta cell (Chap. 417). These drugs are most effective in individuals with type 2 DM of relatively recent onset (<5 years) who have residual endogenous insulin production. First-generation sulfonylureas (chlorpropamide, tolazamide, tolbutamide) have a longer half-life, a greater incidence of hypoglycemia, and more frequent drug interactions, and are no longer used. Second-generation sulfonylureas have a more rapid onset of action and better coverage of the postprandial glucose rise, but the shorter half-life of some agents may require more than once-a-day dosing. Sulfonylureas reduce both fasting and postprandial glucose and should be initiated at low doses and increased at 1- to 2-week intervals based on SMBG. In general, sulfonylureas increase insulin acutely and thus should be taken shortly before a meal; with chronic therapy, though, the insulin release is more sustained. Glimepiride and glipizide can be given in a single daily dose and are preferred over glyburide, especially in the elderly. Repaglinide, nateglinide, and mitiglinide are not sulfonylureas but also interact with the ATP-sensitive potassium channel. Because of their short half-life, these agents are given with each meal or immediately before to reduce meal-related glucose excursions.

Insulin secretagogues, especially the longer acting ones, have the potential to cause hypoglycemia, especially in elderly individuals. Hypoglycemia is usually related to delayed meals, increased physical activity, alcohol intake, or renal insufficiency. Individuals who ingest an overdose of some agents develop prolonged and serious hypoglycemia and should be monitored closely in the hospital (Chap. 420). Most sulfonylureas are metabolized in the liver to compounds (some of which are active) that are cleared by the kidney. Thus, their use in individuals with significant hepatic or renal dysfunction is not advisable. Weight gain, a common side effect of sulfonylurea therapy, results from the increased insulin levels and improvement in glycemic control. Some sulfonylureas have significant drug interactions with alcohol and some medications including warfarin, aspirin, ketoconazole, α -glucosidase inhibitors, and fluconazole. A related isoform of ATP-sensitive potassium channels is present in the myocardium and the brain. All of these agents except glyburide have a low affinity for this isoform. Despite concerns that this agent might affect the myocardial response to ischemia and observational studies suggesting that sulfonylureas increase cardiovascular risk, studies have not shown an increased cardiac mortality with glyburide or other agents in this class.

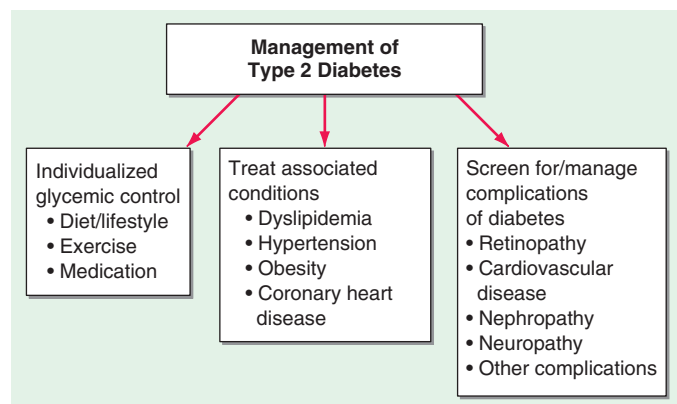


FIGURE 418-2 Essential elements in comprehensive care of type 2 diabetes.

	Mechanism of Action	Examples ^a	HbA _{1c} Reduction (%) ^b	Agent-Specific Advantages	Agent-Specific Disadvantages	Contraindications
Oral						
Biguanides ^{c*}	↓ Hepatic glucose production	Metformin	1–2	Weight neutral, do not cause hypoglycemia, inexpensive, extensive experience, ↓ CV events	Diarrhea, nausea, lactic acidosis	Serum creatinine >1.5 mg/dL (men) >1.4 mg/dL (women) (see text), CHF, radiographic contrast studies, hospitalized patients, acidosis
α-Glucosidase inhibitors ^{c**}	↓ GI glucose absorption	Acarbose, miglitol, voglibose	0.5–0.8	Reduce postprandial glycemia	GI flatulence, liver function tests	Renal/liver disease
Dipeptidyl peptidase IV inhibitors ^{c***}	Prolong endogenous GLP-1 action	Alogliptin, Anagliptin, Gemigliptin, linaagliptin, saxagliptin, sitagliptin, teneligliptin, vildagliptin	0.5–0.8	Well tolerated, do not cause hypoglycemia		Reduced dose with renal disease; one associated with increase heart failure risk; possible association with ACE inhibitor-induced angioedema
Insulin secretagogues: Sulfonylureas ^{c*}	↑ Insulin secretion	Glibornuride, gliclazide, glimepiride, glipizide, gliquidone, glyburide, glycopyramide	1–2	Short onset of action, lower postprandial glucose, inexpensive	Hypoglycemia, weight gain	Renal/liver disease
Insulin secretagogues: Nonsulfonylureas ^{c***}	↑ Insulin secretion	Nateglinide, repaglinide, mitiglinide	0.5–1.0	Short onset of action, lower postprandial glucose	Hypoglycemia	Renal/liver disease
Sodium-glucose cotransporter 2 inhibitors ^{c***}	↑ Urinary glucose excretion	Canagliflozin, dapagliflozin, empagliflozin	0.5–1.0	Insulin secretion and action independent	Urinary and vaginal infections, dehydration, exacerbate tendency to hyperkalemia	Limited clinical experience; moderate renal insufficiency
Thiazolidinediones ^{c***}	↓ Insulin resistance, ↑ glucose utilization	Rosiglitazone, pioglitazone	0.5–1.4	Lower insulin requirements	Peripheral edema, CHF, weight gain, fractures, macular edema	CHF, liver disease
Parenteral						
Amylin agonists ^{c,d***}	Slow gastric emptying, ↓ glucagon	Pramlintide	0.25–0.5	Reduce postprandial glycemia, weight loss	Injection, nausea, ↑ risk of hypoglycemia with insulin	Agents that also slow GI motility
GLP-1 receptor agonists ^{c***}	↑ Insulin, ↓ glucagon, slow gastric emptying, satiety	Exenatide, liraglutide, dulaglutide	0.5–1.0	Weight loss, do not cause hypoglycemia	Injection, nausea, ↑ risk of hypoglycemia with insulin secretagogues	Renal disease, agents that also slow GI motility; medullary carcinoma of thyroid
Insulin ^{c,d****}	↑ Glucose utilization, ↓ hepatic glucose production, and other anabolic actions	See text and Table 418-4	Not limited	Known safety profile	Injection, weight gain, hypoglycemia	
Medical nutrition therapy and physical activity^{c*}	↓ Insulin resistance, ↑ insulin secretion	Low-calorie, low-fat diet, exercise	1–3	Other health benefits	Compliance difficult, long-term success low	

^aExamples are approved for use in at least one country, but may not be available in the United States or all countries. Examples may not include all agents in the class. ^bHbA_{1c} reduction (absolute) depends partly on starting HbA_{1c}. ^cUsed for treatment of type 2 diabetes. ^dUsed in conjunction with insulin for treatment of type 1 diabetes. Cost of agent: 'low, **moderate, ***high, ****variable.

Note: Some agents used to treat type 2 DM are not included in table (see text).

Abbreviations: ACE, angiotensin-converting enzyme; CHF, congestive heart failure; CV, cardiovascular; GI, gastrointestinal; HbA_{1c}, hemoglobin A_{1c}.

INSULIN SECRETAGOGUES—AGENTS THAT ENHANCE GLP-1 RECEPTOR SIGNALING “Incretins” amplify glucose-stimulated insulin secretion (**Chap. 417**). Agents that either act as a GLP-1 receptor agonist or enhance endogenous GLP-1 activity are approved for the treatment of type 2 DM (Table 418-5). Agents in this class do not cause hypoglycemia because of the glucose-dependent nature of incretin-stimulated insulin secretion (unless there is concomitant use of an agent that can lead to hypoglycemia—sulfonylureas, etc.). Exenatide, a synthetic version of a peptide initially identified in the saliva of the Gila monster (exenatide-4), is an analogue of GLP-1. Unlike native GLP-1, which has a half-life of >5 min, differences in the exenatide amino acid sequence render it resistant to the enzyme that degrades GLP-1 (dipeptidyl

peptidase IV [DPP-IV]). Thus, exenatide has prolonged GLP-1-like action and binds to GLP-1 receptors found in islets, the gastrointestinal tract, and the brain. Liraglutide, another GLP-1 receptor agonist, is almost identical to native GLP-1 except for an amino acid substitution and addition of a fatty acyl group (coupled with a γ-glutamic acid spacer) that promote binding to albumin and plasma proteins and prolong its half-life. GLP-1 receptor agonists increase glucose-stimulated insulin secretion, suppress glucagon, and slow gastric emptying. These agents do not promote weight gain; in fact, most patients experience modest weight loss and appetite suppression. Treatment with these agents should start at a low dose to minimize initial side effects (nausea being the limiting one). GLP-1 receptor agonists, available in twice

daily, daily, and weekly injectable formulations, can be used as combination therapy with metformin, sulfonylureas, and thiazolidinediones. Some patients taking insulin secretagogues may require a reduction in those agents to prevent hypoglycemia. The major side effects are nausea, vomiting, and diarrhea. Some formulations carry a black box warning from the FDA because of an increased risk of thyroid C-cell tumors in rodents and are contraindicated in individuals with medullary carcinoma of the thyroid or multiple endocrine neoplasia. Because GLP-1 receptor agonists slow gastric emptying, they may influence the absorption of other drugs. Whether GLP-1 receptor agonists enhance beta cell survival, promote beta cell proliferation, or alter the natural history of type 2 DM is not known. Other GLP-1 receptor agonists and formulations are under development.

DPP-IV inhibitors inhibit degradation of native GLP-1 and thus enhance the incretin effect. DPP-IV, which is widely expressed on the cell surface of endothelial cells and some lymphocytes, degrades a wide range of peptides (not GLP-1 specific). DPP-IV inhibitors promote insulin secretion in the absence of hypoglycemia or weight gain and appear to have a preferential effect on postprandial blood glucose. The levels of GLP-1 action in the patient are greater with the GLP-1 receptor agonists than with DPP-IV inhibitors. DPP-IV inhibitors are used either alone or in combination with other oral agents in type 2 DM. Reduced doses should be given to patients with renal insufficiency. Initial concerns about the pancreatic side effects of GLP-1 receptor agonists and DPP-IV inhibitors (pancreatitis, possible premalignant lesions) appear to be unfounded.

α -GLUCOSIDASE INHIBITORS α -Glucosidase inhibitors reduce postprandial hyperglycemia by delaying glucose absorption; they do not affect glucose utilization or insulin secretion (Table 418-5). Postprandial hyperglycemia, secondary to impaired hepatic and peripheral glucose disposal, contributes significantly to the hyperglycemic state in type 2 DM. These drugs, taken just before each meal, reduce glucose absorption by inhibiting the enzyme that cleaves oligosaccharides into simple sugars in the intestinal lumen. Therapy should be initiated at a low dose with the evening meal and increased to a maximal dose over weeks to months. The major side effects (diarrhea, flatulence, abdominal distention) are related to increased delivery of oligosaccharides to the large bowel and can be reduced somewhat by gradual upward dose titration. α -Glucosidase inhibitors may increase levels of sulfonylureas and increase the incidence of hypoglycemia. Simultaneous treatment with bile acid resins and antacids should be avoided. These agents should not be used in individuals with inflammatory bowel disease, gastroparesis, or a serum creatinine $>177 \mu\text{mol/L}$ (2 mg/dL). This class of agents is not as potent as other oral agents in lowering the HbA_{1c} but is unique because it reduces the postprandial glucose rise even in individuals with type 1 DM. If hypoglycemia from other diabetes treatments occurs while taking these agents, the patient should consume glucose because the degradation and absorption of complex carbohydrates will be retarded.

THIAZOLIDINEDIONES Thiazolidinediones (Table 418-5) reduce insulin resistance by binding to the PPAR- γ (peroxisome proliferator-activated receptor γ) nuclear receptor (which forms a heterodimer with the retinoid X receptor). The PPAR- γ receptor is found at highest levels in adipocytes but is expressed at lower levels in many other tissues. Agonists of this receptor regulate a large number of genes, promote adipocyte differentiation, reduce hepatic fat accumulation, and promote fatty acid storage. Thiazolidinediones promote a redistribution of fat from central to peripheral locations. Circulating insulin levels decrease with use of the thiazolidinediones, indicating a reduction in insulin resistance. Although direct comparisons are not available, the two currently available thiazolidinediones appear to have similar efficacy. The prototype of this class of drugs, troglitazone, was withdrawn from the U.S. market after reports of hepatotoxicity and an association with an idiosyncratic liver reaction that sometimes led to hepatic failure. Although rosiglitazone and pioglitazone do not appear to induce the liver abnormalities seen with troglitazone, the FDA recommends measurement of liver function tests prior to initiating therapy.

Rosiglitazone raises low-density lipoprotein (LDL), high-density lipoprotein (HDL), and triglycerides slightly. Pioglitazone raises HDL to a greater degree and LDL a lesser degree but lowers triglycerides. The clinical significance of the lipid changes with these agents is not known and may be difficult to ascertain because most patients with type 2 DM are also treated with a statin.

Thiazolidinediones are associated with weight gain (2–3 kg), a small reduction in the hematocrit, and a mild increase in plasma volume. Peripheral edema and CHF are more common in individuals treated with these agents. These agents are contraindicated in patients with liver disease or CHF (class III or IV). The FDA has issued an alert that rare patients taking these agents may experience a worsening of diabetic macular edema. An increased risk of fractures has been noted in women taking these agents. Thiazolidinediones have been shown to induce ovulation in premenopausal women with polycystic ovary syndrome. Women should be warned about the risk of pregnancy because the safety of thiazolidinediones in pregnancy is not established.

Concerns about increased cardiovascular risk associated with rosiglitazone led to considerable restrictions on its use and to the FDA issuing a “black box” warning in 2007. However, based on new information, the FDA has revised its guidelines and categorizes rosiglitazone similar to other drugs for type 2 DM. Because of a possible increased risk of bladder cancer, pioglitazone is part of an ongoing FDA safety review.

Sodium-Glucose Co-Transporter 2 Inhibitors (SLGT2) These agents (Table 418-5) lower the blood glucose by selectively inhibiting this co-transporter, which is expressed almost exclusively in the proximal, convoluted tubule in the kidney. This inhibits glucose reabsorption, lowers the renal threshold for glucose, and leads to increased urinary glucose excretion. Thus, the glucose-lowering effect is insulin independent and not related to changes in insulin sensitivity or secretion. Because these agents are the newest class to treat type 2 DM (Table 418-5), clinical experience is limited. Due to the increased urinary glucose, urinary or vaginal infections are more common, and the diuretic effect can lead to reduced intravascular volume. As part of the FDA approval of canagliflozin in 2013, postmarketing studies for cardiovascular outcomes and for monitoring bladder and urinary cancer risk are under way.

OTHER THERAPIES FOR TYPE 2 DM

Bile acid-binding resins Evidence indicates that bile acids, by signaling through nuclear receptors, may have a role in metabolism. Bile acid metabolism is abnormal in type 2 DM. The bile acid-binding resin colesevelam has been approved for the treatment of type 2 DM (already approved for treatment of hypercholesterolemia). Because bile acid-binding resins are minimally absorbed into the systemic circulation, how bile acid-binding resins lower blood glucose is not known. The most common side effects are gastrointestinal (constipation, abdominal pain, and nausea). Bile acid-binding resins can increase plasma triglycerides and should be used cautiously in patients with a tendency for hypertriglyceridemia. The role of this class of drugs in the treatment of type 2 DM is not yet defined.

Bromocriptine A formulation of the dopamine receptor agonist bromocriptine (Cycloset) has been approved by the FDA for the treatment of type 2 DM. However, its role in the treatment of type 2 DM is uncertain.

INSULIN THERAPY IN TYPE 2 DM Insulin should be considered as the initial therapy in type 2 DM, particularly in lean individuals or those with severe weight loss, in individuals with underlying renal or hepatic disease that precludes oral glucose-lowering agents, or in individuals who are hospitalized or acutely ill. Insulin therapy is ultimately required by a substantial number of individuals with type 2 DM because of the progressive nature of the disorder and the relative insulin deficiency that develops in patients with long-standing diabetes. Both physician and patient reluctance often delay the initiation of insulin therapy, but glucose control and patient well-being are improved by insulin therapy in patients who have not reached the glycemic target.

Because endogenous insulin secretion continues and is capable of providing some coverage of mealtime caloric intake, insulin is usually initiated in a single dose of long-acting insulin (0.3–0.4 U/kg per day), given in the evening (NPH) or just before bedtime (NPH, glargine, detemir). Because fasting hyperglycemia and increased hepatic glucose production are prominent features of type 2 DM, bedtime insulin is more effective in clinical trials than a single dose of morning insulin. Glargine given at bedtime has less nocturnal hypoglycemia than NPH insulin. Some physicians prefer a relatively low, fixed starting dose of long-acting insulin (5–15 units) or a weight-based dose (0.2 units/kg). The insulin dose may then be adjusted in 10% increments as dictated by SMBG results. Both morning and bedtime long-acting insulin may be used in combination with oral glucose-lowering agents. Initially, basal insulin may be sufficient, but often prandial insulin coverage with multiple insulin injections is needed as diabetes progresses (see insulin regimens used for type 1 DM). Other insulin formulations that have a combination of short-acting and long-acting insulin (Table 418-4) are sometimes used in patients with type 2 DM because of convenience but do not allow independent adjustment of short-acting and long-acting insulin dose and often do not achieve the same degree of glycemic control as basal/bolus regimens. In selected patients with type 2 DM, insulin-infusion devices may be considered.

CHOICE OF INITIAL GLUCOSE-LOWERING AGENT The level of hyperglycemia and the patient's individualized goal (see "Establishment of Target Level of Glycemic Control") should influence the initial choice of therapy. Assuming that maximal benefit of MNT and increased physical activity has been realized, patients with mild to moderate hyperglycemia (FPG <11.1–13.9 mmol/L [200–250 mg/dL]) often respond well to a single, oral glucose-lowering agent. Patients with more severe hyperglycemia (FPG >13.9 mmol/L [250 mg/dL]) may respond partially but are unlikely to achieve normoglycemia with oral monotherapy. A stepwise approach that starts with a single agent and adds a second agent to achieve the glycemic target can be used (see "Combination therapy with glucose-lowering agents," below). Insulin can be used as initial therapy in individuals with severe hyperglycemia (FPG <13.9–16.7 mmol/L [250–300 mg/dL]) or in those who are symptomatic from the hyperglycemia. This approach is based on the rationale that more rapid glycemic control will reduce "glucose toxicity" to the islet cells, improve endogenous insulin secretion, and possibly allow oral glucose-lowering agents to be more effective. If this occurs, the insulin may be discontinued.

Insulin secretagogues, biguanides, α -glucosidase inhibitors, thiazolidinediones, GLP-1 receptor agonists, DPP-IV inhibitors, SGLT2 inhibitors, and insulin are approved for monotherapy of type 2 DM. Although each class of oral glucose-lowering agents has advantages and disadvantages (Table 418-5), certain generalizations apply: (1) insulin secretagogues, biguanides, GLP-1 receptor agonists, and thiazolidinediones improve glycemic control to a similar degree (1–2% reduction in HbA_{1c}) and are more effective than α -glucosidase inhibitors, DPP-IV inhibitors, and SGLT2 inhibitors; (2) assuming a similar degree of glycemic improvement, no clinical advantage to one class of drugs has been demonstrated; any therapy that improves glycemic control is likely beneficial; (3) insulin secretagogues, GLP-1 receptor agonists, DPP-IV inhibitors, α -glucosidase inhibitors, and SGLT2 inhibitors begin to lower the plasma glucose immediately, whereas the glucose-lowering effects of the biguanides and thiazolidinediones are delayed by weeks; (4) not all agents are effective in all individuals with type 2 DM; (5) biguanides, α -glucosidase inhibitors, GLP-1 receptor agonists, DPP-IV inhibitors, thiazolidinediones, and SGLT2 inhibitors do not directly cause hypoglycemia; (6) most individuals will eventually require treatment with more than one class of oral glucose-lowering agents or insulin, reflecting the progressive nature of type 2 DM; and (7) durability of glycemic control is slightly less for glyburide compared to metformin or rosiglitazone.

Considerable clinical experience exists with metformin and sulfonylureas because they have been available for several decades. It is assumed that the α -glucosidase inhibitors, GLP-1 agonists, DPP-IV inhibitors, thiazolidinediones, and SGLT2 inhibitors will reduce

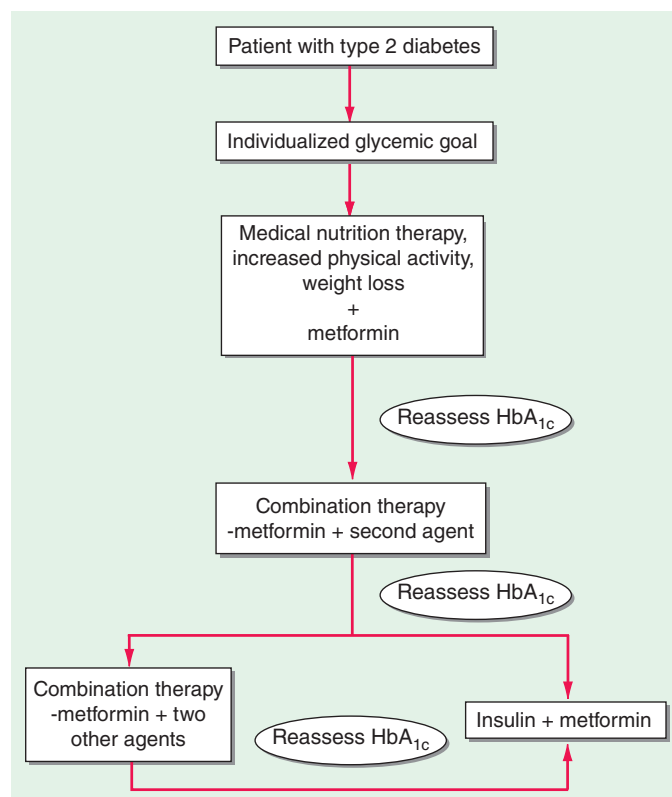


FIGURE 418-3 Glycemic management of type 2 diabetes. See text for discussion of treatment of severe hyperglycemia or symptomatic hyperglycemia. Agents that can be combined with metformin include insulin secretagogues, thiazolidinediones, α -glucosidase inhibitors, DPP-IV inhibitors, GLP-1 receptor agonists, SGLT2 inhibitors, and insulin. HbA_{1c}, hemoglobin HbA_{1c}.

DM-related complications by improving glycemic control, but long-term data are not yet available. The thiazolidinediones are theoretically attractive because they target a fundamental abnormality in type 2 DM, namely insulin resistance. However, all of these agents are currently more costly than metformin and sulfonylureas.

Treatment algorithms by several professional societies (ADA/European Association for the Study of Diabetes [EASD], IDF, AACE) suggest metformin as initial therapy because of its efficacy, known side effect profile, and low cost (Fig. 418-3). Metformin's advantages are that it promotes mild weight loss, lowers insulin levels, and improves the lipid profile slightly. Based on SMBG results and the HbA_{1c}, the dose of metformin should be increased until the glycemic target is achieved or maximum dose is reached. If metformin is not tolerated, then initial therapy with an insulin secretagogue or DPP-IV inhibitor is reasonable.

COMBINATION THERAPY WITH GLUCOSE-LOWERING AGENTS A number of combinations of therapeutic agents are successful in type 2 DM (metformin + second oral agent, metformin + GLP-1 receptor agonist, or metformin + insulin), and the dosing of agents in combination is the same as when the agents are used alone. Because mechanisms of action of the first and second agents should be different, the effect on glycemic control is usually additive. There are little data to support the choice of one combination over another combination. Medication costs vary considerably (Table 418-5), and this often factors into medication choice. Several fixed-dose combinations of oral agents are available, but evidence that they are superior to titration of single agent to a maximum dose and then addition of a second agent is lacking. If adequate control is not achieved with the combination of two agents (based on reassessment of the HbA_{1c} every 3 months), a third oral agent or basal insulin should be added (Fig. 418-3). Treatment approaches vary considerably from country to country. For example, α -glucosidase inhibitors are used commonly in South Asian patients

(Indian), but infrequently in the United States or Europe. Whether this reflects an underlying difference in the disease or physician preference is not clear.

Treatment with insulin becomes necessary as type 2 DM enters the phase of relative insulin deficiency (as seen in long-standing DM) and is signaled by inadequate glycemic control with one or two oral glucose-lowering agents. Insulin alone or in combination should be used in patients who fail to reach the glycemic target. For example, a single dose of long-acting insulin at bedtime is often effective in combination with metformin. In contrast, insulin secretagogues have little utility once insulin therapy is started. Experience using incretin therapies and insulin is limited. As endogenous insulin production falls further, multiple injections of long-acting and short-acting insulin regimens are necessary to control postprandial glucose excursions. These insulin regimens are identical to the long-acting and short-acting combination regimens discussed above for type 1 DM. Because the hyperglycemia of type 2 DM tends to be more “stable,” these regimens can be increased in 10% increments every 2–3 days using the fasting blood glucose results. Weight gain and hypoglycemia are the major adverse effects of insulin therapy. The daily insulin dose required can become quite large (1–2 units/kg per day) as endogenous insulin production falls and insulin resistance persists. Individuals who require >1 unit/kg per day of long-acting insulin should be considered for combination therapy with metformin or a thiazolidinedione. The addition of metformin or a thiazolidinedione can reduce insulin requirements in some individuals with type 2 DM, while maintaining or even improving glycemic control. Insulin plus a thiazolidinedione promotes weight gain and is associated with peripheral edema. Addition of a thiazolidinedione to a patient’s insulin regimen may necessitate a reduction in the insulin dose to avoid hypoglycemia. Patients requiring large doses of insulin (>200 units/day) can be treated with a more concentrated form of insulin, U-500.

EMERGING THERAPIES

Whole pancreas transplantation (performed concomitantly with a renal transplant) may normalize glucose tolerance and is an important therapeutic option in type 1 DM with end-stage renal disease, although it requires substantial expertise and is associated with the side effects of immunosuppression. Pancreatic islet transplantation has been plagued by limitations in pancreatic islet supply and graft survival and remains an area of clinical investigation. Many individuals with long-standing type 1 DM still produce very small amounts of insulin or have insulin-positive cells within the pancreas. This suggests that beta cells may slowly regenerate but are quickly destroyed by the autoimmune process. Thus, efforts to suppress the autoimmune process and to stimulate beta cell regeneration are being tested both at the time of diagnosis and in years after the diagnosis of type 1 DM. Closed-loop pumps that infuse the appropriate amount of insulin in response to changing glucose levels are potentially feasible now that CGM technology has been developed. Bi-hormonal pumps that deliver both insulin and glucagon are under development. New therapies under development for type 2 DM include activators of glucokinase, inhibitors of 11 β -hydroxysteroid dehydrogenase-1, GPR40 agonists, monoclonal antibodies to reduce inflammation, and salsalate.

Bariatric surgery for obese individuals with type 2 DM has shown considerable promise, sometimes with dramatic resolution of the diabetes or major reductions in the needed dose of glucose-lowering therapies (Chap. 416). Several large, unblinded clinical trials have demonstrated a much greater efficacy of bariatric surgery compared to medical management in the treatment of type 2 DM; the durability of the diabetes reversal or improvement is uncertain. The ADA clinical guidelines state that bariatric surgery should be considered in individuals with DM and a body mass index >35 kg/m².

ADVERSE EFFECTS OF THERAPY FOR DIABETES MELLITUS

As with any therapy, the benefits of efforts directed toward glycemic control must be balanced against the risks of treatment (Table 418-5). Side effects of intensive treatment include an increased frequency of serious hypoglycemia, weight gain, increased economic costs, and greater

demands on the patient. In the DCCT, quality of life was very similar in the intensive and standard therapy groups. The most serious complication of therapy for DM is hypoglycemia, and its treatment with oral glucose or glucagon injection is discussed in Chap. 420. Severe, recurrent hypoglycemia warrants examination of treatment regimen and glycemic goal for the individual patient. Weight gain occurs with most (insulin, insulin secretagogues, thiazolidinediones) but not all (metformin, α -glucosidase inhibitors, GLP-1 receptor agonists, DPP-IV inhibitors) therapies. The weight gain is partially due to the anabolic effects of insulin and the reduction in glucosuria. As a result of recent controversies about the optimal glycemic goal and concerns about safety, the FDA now requires information about the cardiovascular safety profile as part of its evaluation of new treatments for type 2 DM.

ACUTE DISORDERS RELATED TO SEVERE HYPERGLYCEMIA

Individuals with type 1 or type 2 DM and severe hyperglycemia (>16.7 mmol/L [300 mg/dL]) should be assessed for clinical stability, including mentation and hydration. Depending on the patient and the rapidity and duration of the severe hyperglycemia, an individual may require more intense and rapid therapy to lower the blood glucose. However, many patients with poorly controlled diabetes and hyperglycemia have few symptoms. The physician should assess if the patient is stable or if diabetic ketoacidosis or a hyperglycemic hyperosmolar state should be considered. Ketones, an indicator of diabetic ketoacidosis, should be measured in individuals with type 1 DM when the plasma glucose is >16.7 mmol/L (300 mg/dL), during a concurrent illness, or with symptoms such as nausea, vomiting, or abdominal pain. Blood measurement of β -hydroxybutyrate is preferred over urine testing with nitroprusside-based assays that measure only acetoacetate and acetone.

Diabetic ketoacidosis (DKA) and hyperglycemic hyperosmolar state (HHS) are acute, severe disorders directly related to diabetes. DKA was formerly considered a hallmark of type 1 DM, but also occurs in individuals who lack immunologic features of type 1 DM and who can sometimes subsequently be treated with oral glucose-lowering agents (these obese individuals with type 2 DM are often of Hispanic or African-American descent). HHS is primarily seen in individuals with type 2 DM. Both disorders are associated with absolute or relative insulin deficiency, volume depletion, and acid-base abnormalities. DKA and HHS exist along a continuum of hyperglycemia, with or without ketosis. The metabolic similarities and differences in DKA and HHS are highlighted in Table 418-6. Both

TABLE 418-6 LABORATORY VALUES IN DIABETIC KETOACIDOSIS (DKA) AND HYPERGLYCEMIC HYPEROSMOLAR STATE (HHS) (REPRESENTATIVE RANGES AT PRESENTATION)

	DKA	HHS
Glucose, ^a mmol/L (mg/dL)	13.9–33.3 (250–600)	33.3–66.6 (600–1200)
Sodium, meq/L	125–135	135–145
Potassium ^{a,b}	Normal to \uparrow	Normal
Magnesium ^a	Normal	Normal
Chloride ^a	Normal	Normal
Phosphate ^{a,b}	Normal	Normal
Creatinine	Slightly \uparrow	Moderately \uparrow
Osmolality (mOsm/mL)	300–320	330–380
Plasma ketones ^a	++++	+/-
Serum bicarbonate, ^a meq/L	<15	Normal to slightly \downarrow
Arterial pH	6.8–7.3	>7.3
Arterial Pco ₂ , ^a mmHg	20–30	Normal
Anion gap ^a (Na – [Cl + HCO ₃])	\uparrow	Normal to slightly \uparrow

^aLarge changes occur during treatment of DKA. ^bAlthough plasma levels may be normal or high at presentation, total-body stores are usually depleted.

2418 disorders are associated with potentially serious complications if not promptly diagnosed and treated.

DIABETIC KETOACIDOSIS

Clinical Features The symptoms and physical signs of DKA are listed in [Table 418-7](#) and usually develop over 24 h. DKA may be the initial symptom complex that leads to a diagnosis of type 1 DM, but more frequently, it occurs in individuals with established diabetes. Nausea and vomiting are often prominent, and their presence in an individual with diabetes warrants laboratory evaluation for DKA. Abdominal pain may be severe and can resemble acute pancreatitis or ruptured viscus. Hyperglycemia leads to glucosuria, volume depletion, and tachycardia. Hypotension can occur because of volume depletion in combination with peripheral vasodilatation. Kussmaul respirations and a fruity odor on the patient's breath (secondary to metabolic acidosis and increased acetone) are classic signs of the disorder. Lethargy and central nervous system depression may evolve into coma with severe DKA but should also prompt evaluation for other reasons for altered mental status (e.g., infection, hypoxemia). Cerebral edema, an extremely serious complication of DKA, is seen most frequently in children. Signs of infection, which may precipitate DKA, should be sought on physical examination, even in the absence of fever. Tissue ischemia (heart, brain) can also be a precipitating factor. Omission of insulin because of an eating disorder, mental health disorders, or an unstable psychosocial environment may sometimes be a factor precipitating DKA.

Pathophysiology DKA results from relative or absolute insulin deficiency combined with counterregulatory hormone excess (glucagon, catecholamines, cortisol, and growth hormone). Both insulin deficiency and glucagon excess, in particular, are necessary for DKA to develop. The decreased ratio of insulin to glucagon promotes gluconeogenesis, glycogenolysis, and ketone body formation in the liver, as well as increases in substrate delivery from fat and muscle (free fatty acids, amino acids) to the liver. Markers of inflammation (cytokines, C-reactive protein) are elevated in both DKA and HHS

The combination of insulin deficiency and hyperglycemia reduces the hepatic level of fructose-2,6-bisphosphate, which alters the activity of phosphofructokinase and fructose-1,6-bisphosphatase. Glucagon excess decreases the activity of pyruvate kinase, whereas insulin deficiency increases the activity of phosphoenolpyruvate carboxykinase. These changes shift the handling of pyruvate toward glucose synthesis and away from glycolysis. The increased levels of glucagon and catecholamines in the face of low insulin levels promote glycogenolysis. Insulin deficiency also reduces levels of the GLUT4 glucose transporter, which impairs glucose uptake into skeletal muscle and fat and reduces intracellular glucose metabolism.

Ketosis results from a marked increase in free fatty acid release from adipocytes, with a resulting shift toward ketone body synthesis in the liver. Reduced insulin levels, in combination with elevations in catecholamines and growth hormone, increase lipolysis and the

release of free fatty acids. Normally, these free fatty acids are converted to triglycerides or very-low-density lipoprotein (VLDL) in the liver. However, in DKA, hyperglucagonemia alters hepatic metabolism to favor ketone body formation, through activation of the enzyme carnitine palmitoyltransferase I. This enzyme is crucial for regulating fatty acid transport into the mitochondria, where beta oxidation and conversion to ketone bodies occur. At physiologic pH, ketone bodies exist as ketoacids, which are neutralized by bicarbonate. As bicarbonate stores are depleted, metabolic acidosis ensues. Increased lactic acid production also contributes to the acidosis. The increased free fatty acids increase triglyceride and VLDL production. VLDL clearance is also reduced because the activity of insulin-sensitive lipoprotein lipase in muscle and fat is decreased. Hypertriglyceridemia may be severe enough to cause pancreatitis.

DKA is often precipitated by increased insulin requirements, as occurs during a concurrent illness ([Table 418-7](#)). Failure to augment insulin therapy often compounds the problem. Complete omission or inadequate administration of insulin by the patient or health care team (in a hospitalized patient with type 1 DM) may precipitate DKA. Patients using insulin-infusion devices with short-acting insulin may develop DKA, because even a brief interruption in insulin delivery (e.g., mechanical malfunction) quickly leads to insulin deficiency.

Laboratory Abnormalities and Diagnosis The timely diagnosis of DKA is crucial and allows for prompt initiation of therapy. DKA is characterized by hyperglycemia, ketosis, and metabolic acidosis (increased anion gap) along with a number of secondary metabolic derangements ([Table 418-6](#)). Occasionally, the serum glucose is only minimally elevated. Serum bicarbonate is frequently <10 mmol/L, and arterial pH ranges between 6.8 and 7.3, depending on the severity of the acidosis. Despite a total-body potassium deficit, the serum potassium at presentation may be mildly elevated, secondary to the acidosis. Total-body stores of sodium, chloride, phosphorus, and magnesium are reduced in DKA but are not accurately reflected by their levels in the serum because of hypovolemia and hyperglycemia. Elevated blood urea nitrogen (BUN) and serum creatinine levels reflect intravascular volume depletion. Interference from acetoacetate may falsely elevate the serum creatinine measurement. Leukocytosis, hypertriglyceridemia, and hyperlipoproteinemia are commonly found as well. Hyperamylasemia may suggest a diagnosis of pancreatitis, especially when accompanied by abdominal pain. However, in DKA the amylase is usually of salivary origin and thus is not diagnostic of pancreatitis. Serum lipase should be obtained if pancreatitis is suspected.

The measured serum sodium is reduced as a consequence of the hyperglycemia (1.6-mmol/L [1.6-meq] reduction in serum sodium for each 5.6-mmol/L [100-mg/dL] rise in the serum glucose). A normal serum sodium in the setting of DKA indicates a more profound water deficit. In “conventional” units, the calculated serum osmolality ($2 \times [\text{serum sodium} + \text{serum potassium}] + \text{plasma glucose} [\text{mg/dL}] / 18 + \text{BUN} / 2.8$) is mildly to moderately elevated, although to a lesser degree than that found in HHS (see below).

In DKA, the ketone body, β -hydroxybutyrate, is synthesized at a threefold greater rate than acetoacetate; however, acetoacetate is preferentially detected by a commonly used ketosis detection reagent (nitroprusside). Serum ketones are present at significant levels (usually positive at serum dilution of $\geq 1:8$). The nitroprusside tablet, or stick, is often used to detect urine ketones; certain medications such as captopril or penicillamine may cause false-positive reactions. Serum or plasma assays for β -hydroxybutyrate are preferred because they more accurately reflect the true ketone body level.

The metabolic derangements of DKA exist along a spectrum, beginning with mild acidosis with moderate hyperglycemia evolving into more severe findings. The degree of acidosis and hyperglycemia do not necessarily correlate closely because a variety of factors determine the level of hyperglycemia (oral intake, urinary glucose loss). Ketonemia is a consistent finding in DKA and distinguishes it from simple hyperglycemia. The differential diagnosis of DKA includes starvation ketosis, alcoholic ketoacidosis (bicarbonate usually >15 meq/L), and other forms of increased anion-gap acidosis ([Chap. 66](#)).

TABLE 418-7 MANIFESTATIONS OF DIABETIC KETOACIDOSIS

Symptoms	Physical Findings
Nausea/vomiting	Tachycardia
Thirst/polyuria	Dehydration/hypotension
Abdominal pain	Tachypnea/Kussmaul respirations/respiratory distress
Shortness of breath	Abdominal tenderness (may resemble acute pancreatitis or surgical abdomen)
Precipitating events	Lethargy/obtundation/cerebral edema/possibly coma
Inadequate insulin administration	
Infection (pneumonia/UTI/gastroenteritis/sepsis)	
Infarction (cerebral, coronary, mesenteric, peripheral)	
Drugs (cocaine)	
Pregnancy	

Abbreviation: UTI, urinary tract infection.

The management of DKA is outlined in **Table 418-8**. After initiating IV fluid replacement and insulin therapy, the agent or event that precipitated the episode of DKA should be sought and aggressively treated. If the patient is vomiting or has altered mental status, a nasogastric tube should be inserted to prevent aspiration of gastric contents. Central to successful treatment of DKA is careful monitoring and frequent reassessment to ensure that the patient and the metabolic derangements are improving. A comprehensive flow sheet should record chronologic changes in vital signs, fluid intake and output, and laboratory values as a function of insulin administered.

After the initial bolus of normal saline, replacement of the sodium and free water deficit is carried out over the next 24 h (fluid deficit is often 3–5 L). When hemodynamic stability and adequate urine output are achieved, IV fluids should be switched to 0.45% saline depending on the calculated volume deficit. The change to 0.45% saline helps to reduce the trend toward hyperchloremia later in the course of DKA. Alternatively, initial use of lactated Ringer's IV solution may reduce the hyperchloremia that commonly occurs with normal saline.

A bolus of IV (0.1 units/kg) short-acting insulin should be administered immediately (Table 418-8), and subsequent treatment should provide continuous and adequate levels of circulating insulin. IV administration is preferred (0.1 units/kg of regular insulin per hour) because it ensures rapid distribution and allows adjustment of the infusion rate as the patient responds to therapy. In mild episodes of DKA, short-acting insulin can be used SC. IV insulin

should be continued until the acidosis resolves and the patient is metabolically stable. As the acidosis and insulin resistance associated with DKA resolve, the insulin infusion rate can be decreased (to 0.05–0.1 units/kg per hour). Long-acting insulin, in combination with SC short-acting insulin, should be administered as soon as the patient resumes eating, because this facilitates transition to an outpatient insulin regimen and reduces length of hospital stay. It is crucial to continue the insulin infusion until adequate insulin levels are achieved by administering long-acting insulin by the SC route. Even relatively brief periods of inadequate insulin administration in this transition phase may result in DKA relapse.

Hyperglycemia usually improves at a rate of 4.2–5.6 mmol/L (75–100 mg/dL) per hour as a result of insulin-mediated glucose disposal, reduced hepatic glucose release, and rehydration. The latter reduces catecholamines, increases urinary glucose loss, and expands the intravascular volume. The decline in the plasma glucose within the first 1–2 h may be more rapid and is mostly related to volume expansion. When the plasma glucose reaches 13.9 mmol/L (250 mg/dL), glucose should be added to the 0.45% saline infusion to maintain the plasma glucose in the 8.3–13.9 mmol/L (150–250 mg/dL) range, and the insulin infusion should be continued. Ketoacidosis begins to resolve as insulin reduces lipolysis, increases peripheral ketone body use, suppresses hepatic ketone body formation, and promotes bicarbonate regeneration. However, the acidosis and ketosis resolve more slowly than hyperglycemia. As ketoacidosis improves, β -hydroxybutyrate is converted to acetoacetate. Ketone body levels may appear to increase if measured by laboratory assays that use the nitroprusside reaction, which only detects acetoacetate and acetone. The improvement in acidosis and anion gap, a result of bicarbonate regeneration and decline in ketone bodies, is reflected by a rise in the serum bicarbonate level and the arterial pH. Depending on the rise of serum chloride, the anion gap (but not bicarbonate) will normalize. A hyperchloremic acidosis (serum bicarbonate of 15–18 mmol/L [15–18 meq/L]) often follows successful treatment and gradually resolves as the kidneys regenerate bicarbonate and excrete chloride.

Potassium stores are depleted in DKA (estimated deficit 3–5 mmol/kg [3–5 meq/kg]). During treatment with insulin and fluids, various factors contribute to the development of hypokalemia. These include insulin-mediated potassium transport into cells, resolution of the acidosis (which also promotes potassium entry into cells), and urinary loss of potassium salts of organic acids. Thus, potassium repletion should commence as soon as adequate urine output and a normal serum potassium are documented. If the initial serum potassium level is elevated, then potassium repletion should be delayed until the potassium falls into the normal range. Inclusion of 20–40 meq of potassium in each liter of IV fluid is reasonable, but additional potassium supplements may also be required. To reduce the amount of chloride administered, potassium phosphate or acetate can be substituted for the chloride salt. The goal is to maintain the serum potassium at >3.5 mmol/L (3.5 meq/L).

Despite a bicarbonate deficit, bicarbonate replacement is not usually necessary. In fact, theoretical arguments suggest that bicarbonate administration and rapid reversal of acidosis may impair cardiac function, reduce tissue oxygenation, and promote hypokalemia. The results of most clinical trials do not support the routine use of bicarbonate replacement, and one study in children found that bicarbonate use was associated with an increased risk of cerebral edema. However, in the presence of severe acidosis (arterial pH <7.0), the ADA advises bicarbonate (50 mmol/L [meq/L] of sodium bicarbonate in 200 mL of sterile water with 10 meq/L KCl per hour for 2 h until the pH is >7.0). Hypophosphatemia may result from increased glucose usage, but randomized clinical trials have not demonstrated that phosphate replacement is beneficial in DKA. If the serum phosphate is <0.32 mmol/L (1 mg/dL), then phosphate supplement should be considered and the serum calcium monitored. Hypomagnesemia may develop during DKA therapy and may also require supplementation.

TABLE 418-8 MANAGEMENT OF DIABETIC KETOACIDOSIS

1. Confirm diagnosis (\uparrow plasma glucose, positive serum ketones, metabolic acidosis).
2. Admit to hospital; intensive care setting may be necessary for frequent monitoring or if pH <7.00 or unconscious.
3. Assess:
 - Serum electrolytes (K^+ , Na^+ , Mg^{2+} , Cl^- , bicarbonate, phosphate)
 - Acid-base status—pH, HCO_3^- , Pco_2 , β -hydroxybutyrate
 - Renal function (creatinine, urine output)
4. Replace fluids: 2–3 L of 0.9% saline over first 1–3 h (10–20 mL/kg per hour); subsequently, 0.45% saline at 250–500 mL/h; change to 5% glucose and 0.45% saline at 150–250 mL/h when plasma glucose reaches 250 mg/dL (13.9 mmol/L).
5. Administer short-acting insulin: IV (0.1 units/kg), then 0.1 units/kg per hour by continuous IV infusion; increase two- to threefold if no response by 2–4 h. If the initial serum potassium is <3.3 mmol/L (3.3 meq/L), do not administer insulin until the potassium is corrected.
6. Assess patient: What precipitated the episode (noncompliance, infection, trauma, pregnancy, infarction, cocaine)? Initiate appropriate workup for precipitating event (cultures, CXR, ECG).
7. Measure capillary glucose every 1–2 h; measure electrolytes (especially K^+ , bicarbonate, phosphate) and anion gap every 4 h for first 24 h.
8. Monitor blood pressure, pulse, respirations, mental status, fluid intake and output every 1–4 h.
9. Replace K^+ : 10 meq/h when plasma K^+ <5.0–5.2 meq/L (or 20–30 meq/L of infusion fluid), ECG normal, urine flow and normal creatinine documented; administer 40–80 meq/h when plasma K^+ <3.5 meq/L or if bicarbonate is given. If initial serum potassium is >5.2 mmol/L (5.2 meq/L), do not supplement K^+ until the potassium is corrected.
10. See text about bicarbonate or phosphate supplementation.
11. Continue above until patient is stable, glucose goal is 8.3–13.9 mmol/L (150–250 mg/dL), and acidosis is resolved. Insulin infusion may be decreased to 0.05–0.1 units/kg per hour.
12. Administer long-acting insulin as soon as patient is eating. Allow for a 2–4 hour overlap in insulin infusion and SC insulin injection.

Abbreviations: CXR, chest x-ray; ECG, electrocardiogram.

Source: Adapted from M Sperling, in *Therapy for Diabetes Mellitus and Related Disorders*, American Diabetes Association, Alexandria, VA, 1998; and AE Kitabchi et al: *Diabetes Care* 32:1335, 2009.

With appropriate therapy, the mortality rate of DKA is low (<1%) and is related more to the underlying or precipitating event, such as infection or myocardial infarction. Venous thrombosis, upper gastrointestinal bleeding, and acute respiratory distress syndrome occasionally complicate DKA. The major nonmetabolic complication of DKA therapy is cerebral edema, which most often develops in children as DKA is resolving. The etiology of and optimal therapy for cerebral edema are not well established, but overreplacement of free water should be avoided.

Following treatment, the physician and patient should review the sequence of events that led to DKA to prevent future recurrences. Foremost is patient education about the symptoms of DKA, its precipitating factors, and the management of diabetes during a concurrent illness. During illness or when oral intake is compromised, patients should (1) frequently measure the capillary blood glucose; (2) measure urinary ketones when the serum glucose is >16.5 mmol/L (300 mg/dL); (3) drink fluids to maintain hydration; (4) continue or increase insulin; and (5) seek medical attention if dehydration, persistent vomiting, or uncontrolled hyperglycemia develop. Using these strategies, early DKA can be prevented or detected and treated appropriately on an outpatient basis.

HYPERGLYCEMIC HYPEROSMOLAR STATE

Clinical Features The prototypical patient with HHS is an elderly individual with type 2 DM, with a several-week history of polyuria, weight loss, and diminished oral intake that culminates in mental confusion, lethargy, or coma. The physical examination reflects profound dehydration and hyperosmolality and reveals hypotension, tachycardia, and altered mental status. Notably absent are symptoms of nausea, vomiting, and abdominal pain and the Kussmaul respirations characteristic of DKA. HHS is often precipitated by a serious, concurrent illness such as myocardial infarction or stroke. Sepsis, pneumonia, and other serious infections are frequent precipitants and should be sought. In addition, a debilitating condition (prior stroke or dementia) or social situation that compromises water intake usually contributes to the development of the disorder.

Pathophysiology Relative insulin deficiency and inadequate fluid intake are the underlying causes of HHS. Insulin deficiency increases hepatic glucose production (through glycogenolysis and gluconeogenesis) and impairs glucose utilization in skeletal muscle (see above discussion of DKA). Hyperglycemia induces an osmotic diuresis that leads to intravascular volume depletion, which is exacerbated by inadequate fluid replacement. The absence of ketosis in HHS is not understood. Presumably, the insulin deficiency is only relative and less severe than in DKA. Lower levels of counterregulatory hormones and free fatty acids have been found in HHS than in DKA in some studies. It is also possible that the liver is less capable of ketone body synthesis or that the insulin/glucagon ratio does not favor ketogenesis.

Laboratory Abnormalities and Diagnosis The laboratory features in HHS are summarized in Table 418-6. Most notable are the marked hyperglycemia (plasma glucose may be >55.5 mmol/L [1000 mg/dL]), hyperosmolality (>350 mosmol/L), and prerenal azotemia. The measured serum sodium may be normal or slightly low despite the marked hyperglycemia. The corrected serum sodium is usually increased (add 1.6 meq to measured sodium for each 5.6-mmol/L [100-mg/dL] rise in the serum glucose). In contrast to DKA, acidosis and ketonemia are absent or mild. A small anion-gap metabolic acidosis may be present secondary to increased lactic acid. Moderate ketonuria, if present, is secondary to starvation.

TREATMENT HYPERGLYCEMIC HYPEROSMOLAR STATE

Volume depletion and hyperglycemia are prominent features of both HHS and DKA. Consequently, therapy of these disorders shares several elements (Table 418-8). In both disorders, careful

monitoring of the patient's fluid status, laboratory values, and insulin infusion rate is crucial. Underlying or precipitating problems should be aggressively sought and treated. In HHS, fluid losses and dehydration are usually more pronounced than in DKA due to the longer duration of the illness. The patient with HHS is usually older, more likely to have mental status changes, and more likely to have a life-threatening precipitating event with accompanying comorbidities. Even with proper treatment, HHS has a substantially higher mortality rate than DKA (up to 15% in some clinical series).

Fluid replacement should initially stabilize the hemodynamic status of the patient (1–3 L of 0.9% normal saline over the first 2–3 h). Because the fluid deficit in HHS is accumulated over a period of days to weeks, the rapidity of reversal of the hyperosmolar state must balance the need for free water repletion with the risk that too rapid a reversal may worsen neurologic function. If the serum sodium is >150 mmol/L (150 meq/L), 0.45% saline should be used. After hemodynamic stability is achieved, the IV fluid administration is directed at reversing the free water deficit using hypotonic fluids (0.45% saline initially, then 5% dextrose in water [D₅W]). The calculated free water deficit (which averages 9–10 L) should be reversed over the next 1–2 days (infusion rates of 200–300 mL/h of hypotonic solution). Potassium repletion is usually necessary and should be dictated by repeated measurements of the serum potassium. In patients taking diuretics, the potassium deficit can be quite large and may be accompanied by magnesium deficiency. Hypophosphatemia may occur during therapy and can be improved by using KPO₄ and beginning nutrition.

As in DKA, rehydration and volume expansion lower the plasma glucose initially, but insulin is also required. A reasonable regimen for HHS begins with an IV insulin bolus of 0.1 unit/kg followed by IV insulin at a constant infusion rate of 0.1 unit/kg per hour. If the serum glucose does not fall, increase the insulin infusion rate by twofold. As in DKA, glucose should be added to IV fluid when the plasma glucose falls to 13.9 mmol/L (250 mg/dL), and the insulin infusion rate should be decreased to 0.05–0.1 unit/kg per hour. The insulin infusion should be continued until the patient has resumed eating and can be transferred to a SC insulin regimen. The patient should be discharged from the hospital on insulin, although some patients can later switch to oral glucose-lowering agents.

MANAGEMENT OF DIABETES IN A HOSPITALIZED PATIENT

Virtually all medical and surgical subspecialties are involved in the care of hospitalized patients with diabetes. Hyperglycemia, whether in a patient with known diabetes or in someone without known diabetes, appears to be a predictor of poor outcome in hospitalized patients. General anesthesia, surgery, infection, or concurrent illness raises the levels of counterregulatory hormones (cortisol, growth hormone, catecholamines, and glucagon) and cytokines that may lead to transient insulin resistance and hyperglycemia. These factors increase insulin requirements by increasing glucose production and impairing glucose utilization and thus may worsen glycemic control. The concurrent illness or surgical procedure may lead to variable insulin absorption and also prevent the patient with DM from eating normally and, thus, may promote hypoglycemia. Glycemic control should be assessed on admission using the HbA_{1c}. Electrolytes, renal function, and intravascular volume status should be assessed as well. The high prevalence of CVD in individuals with DM (especially in type 2 DM) may necessitate preoperative cardiovascular evaluation (Chap. 419).

The goals of diabetes management during hospitalization are normoglycemia, avoidance of hypoglycemia, and transition back to the outpatient diabetes treatment regimen. Upon hospital admission, frequent glycemic monitoring should begin, as should planning for diabetes management after discharge. Glycemic control appears to improve the clinical outcomes in a variety of settings, but optimal glycemic goals for the hospitalized patient are incompletely defined.

In a number of cross-sectional studies of patients with diabetes, a greater degree of hyperglycemia was associated with worse cardiac, neurologic, and infectious outcomes. In some studies, patients who do not have preexisting diabetes but who develop modest blood glucose elevations during their hospitalization appear to benefit from achieving near-normoglycemia using insulin treatment. However, a large randomized clinical trial (Normoglycemia in Intensive Care Evaluation Survival Using Glucose Algorithm Regulation [NICE-SUGAR]) of individuals in the ICU (most of whom were receiving mechanical ventilation) found an increased mortality rate and a greater number of episodes of severe hypoglycemia with very strict glycemic control (target blood glucose of 4.5–6 mmol/L or 81–108 mg/dL) compared to individuals with a more moderate glycemic goal (mean blood glucose of 8 mmol/L or 144 mg/dL). Currently, most data suggest that very strict blood glucose control in acutely ill patients likely worsens outcomes and increases the frequency of hypoglycemia. The ADA suggests the following glycemic goals for hospitalized patients: (1) in critically ill patients: glucose of 7.8–10.0 mmol/L or 140–180 mg/dL; (2) in non-critically ill patients: premeal glucose <7.8 mmol/L (140 mg/dL) and at other times blood glucose <10 mmol/L (180 mg/dL).

Critical aspects for optimal diabetes care in the hospital include the following. (1) A hospital system approach to treatment of hyperglycemia and prevention of hypoglycemia is needed. Inpatient diabetes management teams consisting of nurse practitioners and physicians are increasingly common. (2) Diabetes treatment plans should focus on the transition from the ICU and the transition from the inpatient to outpatient setting. (3) Adjustment of the discharge treatment regimen of patients whose diabetes was poorly controlled on admission (as reflected by the HbA_{1c}) is necessary.

The physician caring for an individual with diabetes in the perioperative period, during times of infection or serious physical illness, or simply when the patient is fasting for a diagnostic procedure must monitor the plasma glucose vigilantly, adjust the diabetes treatment regimen, and provide glucose infusion as needed. Hypoglycemia is frequent in hospitalized patients, and many of these episodes are avoidable. Hospital systems should have a diabetes management protocol to avoid inpatient hypoglycemia. Measures to reduce or prevent hypoglycemia include frequent glucose monitoring and anticipating potential modifications of insulin/glucose administration because of changes in the clinical situation or treatment (e.g., tapering of glucocorticoids) or interruption of enteral or parenteral infusions or PO intake.

Depending on the severity of the patient's illness and the hospital setting, the physician can use either an insulin infusion or SC insulin. Insulin infusions are preferred in the ICU or in a clinically unstable setting. The absorption of SC insulin may be variable in such situations. Insulin infusions can also effectively control plasma glucose in the perioperative period and when the patient is unable to take anything by mouth. Regular insulin is used rather than insulin analogues for IV insulin infusion because it is less expensive and equally effective. The physician must consider carefully the clinical setting in which an insulin infusion will be used, including whether adequate ancillary personnel are available to monitor the plasma glucose frequently and whether they can adjust the insulin infusion rate to maintain the plasma glucose within the optimal range. Insulin-infusion algorithms should integrate the insulin sensitivity of the patient, frequent blood glucose monitoring, and the trend of changes in the blood glucose to determine the insulin-infusion rate. Insulin-infusion algorithms jointly developed and implemented by nursing and physician staff are advised. Because of the short half-life of IV regular insulin, it is necessary to administer long-acting insulin prior to discontinuation of the insulin infusion (2–4 h before the infusion is stopped) to avoid a period of insulin deficiency.

In patients who are not critically ill or not in the ICU, basal or “scheduled” insulin is provided by SC, long-acting insulin supplemented by prandial and/or “corrective” insulin using a short-acting insulin (insulin analogues preferred). The use of “sliding scale,” short-acting insulin alone, where no insulin is given unless the blood

glucose is elevated, is inadequate for inpatient glucose management and should not be used. The short-acting, preprandial insulin dose should include coverage for food consumption (based on anticipated carbohydrate intake) plus a corrective or supplemental insulin based on the patient's insulin sensitivity and the blood glucose. For example, if the patient is thin (and likely insulin-sensitive), a corrective insulin supplement might be 1 unit for each 2.7 mmol/L (50 mg/dL) over the glucose target. If the patient is obese and insulin-resistant, then the insulin supplement might be 2 units for each 2.7 mmol/L (50 mg/dL) over the glucose target. It is critical to individualize the regimen and adjust the basal or “scheduled” insulin dose frequently, based on the corrective insulin required. A consistent carbohydrate diabetes meal plan for hospitalized patients provides a predictable amount of carbohydrate for a particular meal each day (but not necessarily the same amount for breakfast, lunch, and supper). The hospital diet should be determined by a nutritionist; terms such as *ADA diet* or *low-sugar diet* are no longer used.

Individuals with type 1 DM who are undergoing general anesthesia and surgery or who are seriously ill should receive continuous insulin, either through an IV insulin infusion or by SC administration of a reduced dose of long-acting insulin. Short-acting insulin alone is insufficient. Prolongation of a surgical procedure or delay in the recovery room is not uncommon and may result in periods of insulin deficiency leading to DKA. Insulin infusion is the preferred method for managing patients with type 1 DM in the perioperative period or when serious concurrent illness is present (0.5–1.0 units/h of regular insulin). If the diagnostic or surgical procedure is brief and performed under local or regional anesthesia, a reduced dose of SC, long-acting insulin may suffice (30–50% reduction, with short-acting insulin withheld or reduced). This approach facilitates the transition back to long-acting insulin after the procedure. Glucose may be infused to prevent hypoglycemia. The blood glucose should be monitored frequently during the illness or in the perioperative period.

Individuals with type 2 DM can be managed with either an insulin infusion or SC long-acting insulin (25–50% reduction depending on clinical setting) plus preprandial, short-acting insulin. Oral glucose-lowering agents should be discontinued upon admission and are not useful in regulating the plasma glucose in clinical situations where the insulin requirements and glucose intake are changing rapidly. Moreover, these oral agents may be dangerous if the patient is fasting (e.g., hypoglycemia with sulfonylureas). Metformin should be withheld when radiographic contrast media will be given or if unstable CHF, acidosis, or declining renal function is present.

SPECIAL CONSIDERATIONS IN DIABETES MELLITUS

TOTAL PARENTERAL NUTRITION

(See also Chap. 98e) Total parenteral nutrition (TPN) greatly increases insulin requirements. In addition, individuals not previously known to have DM may become hyperglycemic during TPN and require insulin treatment. IV insulin infusion is the preferred treatment for hyperglycemia, and rapid titration to the required insulin dose is done most efficiently using a separate insulin infusion. After the total insulin dose has been determined, insulin may be added directly to the TPN solution or, preferably, given as a separate infusion or subcutaneously. Often, individuals receiving either TPN or enteral nutrition receive their caloric loads continuously and not at “meal times”; consequently, SC insulin regimens must be adjusted.

GLUCOCORTICOIDS

Glucocorticoids increase insulin resistance, decrease glucose utilization, increase hepatic glucose production, and impair insulin secretion. These changes lead to a worsening of glycemic control in individuals with DM and may precipitate diabetes in other individuals (“steroid-induced diabetes”). The effects of glucocorticoids on glucose homeostasis are dose-related, usually reversible, and most pronounced in the postprandial period. If the FPG is near the normal range, oral diabetes agents (e.g., sulfonylureas, metformin) may be sufficient to reduce

2422 hyperglycemia. If the FPG is >11.1 mmol/L (200 mg/dL), oral agents are usually not efficacious and insulin therapy is required. Short-acting insulin may be required to supplement long-acting insulin in order to control postprandial glucose excursions.

REPRODUCTIVE ISSUES

Reproductive capacity in either men or women with DM appears to be normal. Menstrual cycles may be associated with alterations in glycaemic control in women with DM. Pregnancy is associated with marked insulin resistance; the increased insulin requirements often precipitate DM and lead to the diagnosis of gestational diabetes mellitus (GDM). Glucose, which at high levels is a teratogen to the developing fetus, readily crosses the placenta, but insulin does not. Thus, hyperglycemia from the maternal circulation may stimulate insulin secretion in the fetus. The anabolic and growth effects of insulin may result in macrosomia. GDM complicates ~7% (range 1–14%) of pregnancies. The incidence of GDM is greatly increased in certain ethnic groups, including African Americans and Latinas, consistent with a similar increased risk of type 2 DM. Current recommendations advise screening for glucose intolerance between weeks 24 and 28 of pregnancy in women with increased risk for GDM (≥ 25 years; obesity; family history of DM; member of an ethnic group such as Latina, Native American, Asian American, African American, or Pacific Islander). Therapy for GDM is similar to that for individuals with pregnancy-associated diabetes and involves MNT and insulin, if hyperglycemia persists. Oral glucose-lowering agents are not approved for use during pregnancy, but studies using metformin or glyburide have shown efficacy and have not found toxicity. However, many physicians use insulin to treat GDM. With current practices, the morbidity and mortality rates of the mother with GDM and the fetus are not different from those in the nondiabetic population. Individuals who develop GDM are at marked increased risk for developing type 2 DM in the future and should be screened periodically for DM. Most individuals with GDM revert to normal glucose tolerance after delivery, but some will continue to have overt diabetes or impairment of glucose tolerance after delivery. In addition, children of women with GDM appear to be at risk for obesity and glucose intolerance and have an increased risk of diabetes beginning in the later stages of adolescence.

Pregnancy in individuals with known DM requires meticulous planning and adherence to strict treatment regimens. Intensive diabetes management and normalization of the HbA_{1c} are essential for individuals with existing DM who are planning pregnancy. The most crucial period of glycaemic control is soon after fertilization. The risk of fetal malformations is increased 4–10 times in individuals with uncontrolled DM at the time of conception, and normal plasma glucose during the preconception period and throughout the periods of organ development in the fetus should be the goal.

LIPODYSTROPHIC DM

Lipodystrophy, or the loss of subcutaneous fat tissue, may be generalized in certain genetic conditions such as leprechaunism. Generalized lipodystrophy is associated with severe insulin resistance and is often accompanied by acanthosis nigricans and dyslipidemia. Localized lipodystrophy associated with insulin injections has been reduced considerably by the use of human insulin.

Protease Inhibitors and Lipodystrophy Protease inhibitors used in the treatment of HIV disease (Chap. 226) have been associated with a centripetal accumulation of fat (visceral and abdominal area), accumulation of fat in the dorsocervical region, loss of extremity fat, decreased insulin sensitivity (elevations of the fasting insulin level and reduced glucose tolerance on IV glucose tolerance testing), and dyslipidemia. Although many aspects of the physical appearance of these individuals resemble Cushing's syndrome, increased cortisol levels do not account for this appearance. The possibility remains that this is related to HIV infection by some undefined mechanism, because some features of the syndrome were observed before the introduction of protease inhibitors. Therapy for HIV-related lipodystrophy is not well established.

419 Diabetes Mellitus: Complications

Alvin C. Powers

Diabetes-related complications affect many organ systems and are responsible for the majority of morbidity and mortality associated with the disease. Strikingly, in the United States, diabetes is the leading cause of new blindness in adults, renal failure, and nontraumatic lower extremity amputation. Diabetes-related complications usually do not appear until the second decade of hyperglycemia. Because type 2 diabetes mellitus (DM) often has a long asymptomatic period of hyperglycemia before diagnosis, many individuals with type 2 DM have complications at the time of diagnosis. Fortunately, many of the diabetes-related complications can be prevented or delayed with early detection, aggressive glycaemic control, and efforts to minimize the risks of complications.

Diabetes-related complications can be divided into vascular and nonvascular complications and are similar for type 1 and type 2 DM (Table 419-1). The vascular complications of DM are further subdivided into microvascular (retinopathy, neuropathy, nephropathy) and macrovascular complications (coronary heart disease [CHD], peripheral arterial disease [PAD], cerebrovascular disease). Microvascular complications are diabetes-specific, whereas macrovascular complications are similar to those in nondiabetics but occur at greater frequency in individuals with diabetes. Nonvascular complications include gastroparesis, infections, skin changes, and hearing loss. Whether type 2 DM increases the risk of dementia or impaired cognitive function is not clear.

GLYCEMIC CONTROL AND COMPLICATIONS

The microvascular complications of both type 1 and type 2 DM result from chronic hyperglycemia (Fig. 419-1). Evidence implicating a causative role for chronic hyperglycemia in the development of macrovascular complications is less conclusive. CHD events and mortality rate are two to four times greater in patients with type 2 DM and correlate with fasting and postprandial plasma glucose levels as well the

TABLE 419-1 DIABETES-RELATED COMPLICATIONS

Microvascular
Eye disease
Retinopathy (nonproliferative/proliferative)
Macular edema
Neuropathy
Sensory and motor (mono- and polyneuropathy)
Autonomic
Nephropathy (albuminuria and declining renal function)
Macrovascular
Coronary heart disease
Peripheral arterial disease
Cerebrovascular disease
Other
Gastrointestinal (gastroparesis, diarrhea)
Genitourinary (uropathy/sexual dysfunction)
Dermatologic
Infectious
Cataracts
Glaucoma
Cheiroarthropathy ^a
Periodontal disease
Hearing loss
Other comorbid conditions associated with diabetes (relationship to hyperglycemia is uncertain): depression, obstructive sleep apnea, fatty liver disease, hip fracture, osteoporosis (in type 1 diabetes), cognitive impairment or dementia, low testosterone in men

^aThickened skin and reduced joint mobility.

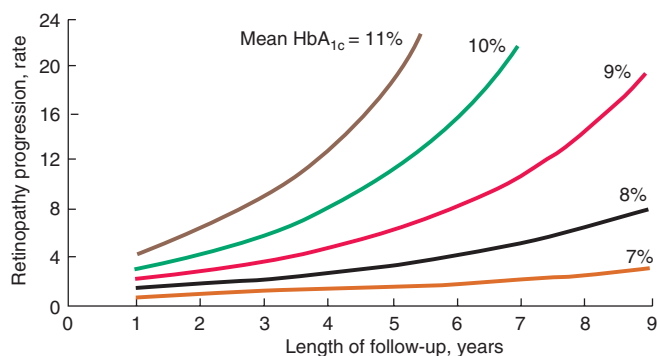


FIGURE 419-1 Relationship of glycemic control and diabetes duration to diabetic retinopathy. The progression of retinopathy in individuals in the Diabetes Control and Complications Trial is graphed as a function of the length of follow-up with different curves for different hemoglobin A_{1c} (HbA_{1c}) values. (Adapted from *The Diabetes Control and Complications Trial Research Group: Diabetes 44:968, 1995.*)

hemoglobin A_{1c} (HbA_{1c}). Other factors such as dyslipidemia and hypertension also play important roles in macrovascular complications.

The Diabetes Control and Complications Trial (DCCT) provided definitive proof that reduction in chronic hyperglycemia can prevent many complications of type 1 DM (Fig. 419-1). This large multicenter clinical trial randomized more than 1400 individuals with type 1 DM to either intensive or conventional diabetes management and prospectively evaluated the development of diabetes-related complications during a mean follow-up of 6.5 years. Individuals in the intensive diabetes management group received multiple administrations of insulin each day (injection or pump) along with extensive educational, psychological, and medical support. Individuals in the conventional diabetes management group received twice-daily insulin injections and quarterly nutritional, educational, and clinical evaluation. The goal in the former group was normoglycemia; the goal in the latter group was prevention of symptoms of diabetes. Individuals in the intensive diabetes management group achieved a substantially lower HbA_{1c} (7.3%) than individuals in the conventional diabetes management group (9.1%). After the DCCT results were reported in 1993, study participants continue to be followed in the Epidemiology of Diabetes Intervention and Complications (EDIC) trial, which recently completed 30 years of follow-up (DCCT + EDIC). At the end of the DCCT phase, study participants in both intensive and conventional arms were offered intensive therapy. However, during the subsequent follow-up of more than 18 years, the initial separation in glycemic control disappeared with both arms maintaining a mean HbA_{1c} of 8.0%.

The DCCT phase demonstrated that improvement of glycemic control reduced nonproliferative and proliferative retinopathy (47% reduction), microalbuminuria (39% reduction), clinical nephropathy (54% reduction), and neuropathy (60% reduction). Improved glycemic control also slowed the progression of early diabetic complications. During the DCCT phase, weight gain (4.6 kg) and severe hypoglycemia (requiring assistance of another person to treat) were more common in the intensive therapy group. The benefits of an improvement in glycemic control occurred over the entire range of HbA_{1c} values (Fig. 419-1), indicating that at any HbA_{1c} level, an improvement in glycemic control is beneficial. The results of the DCCT predicted that individuals in the intensive diabetes management group would gain 7.7 additional years of vision, 5.8 additional years free from end-stage renal disease (ESRD), and 5.6 years free from lower extremity amputations. If all complications of DM were combined, individuals in the intensive diabetes management group would experience 15.3 more years of life without significant microvascular or neurologic complications of DM, compared to individuals who received standard therapy. This translates into an additional 5.1 years of life expectancy for individuals in the intensive diabetes management group. The 30-year follow-up data in the intensively treated group show a continued reduction in retinopathy, nephropathy, and cardiovascular disease. For example, individuals in the intensive therapy group had a 42–57% reduction in

cardiovascular events (nonfatal myocardial infarction [MI], stroke, or death from a cardiovascular event) at a mean follow-up of 17 years, even though their subsequent glycemic control was the same as those in the conventional diabetes management group from years 6.5–17. During the EDIC phase, less than 1% of the cohort had become blind, lost a limb to amputation, or required dialysis.

The United Kingdom Prospective Diabetes Study (UKPDS) studied the course of >5000 individuals with type 2 DM for >10 years. This study used multiple treatment regimens and monitored the effect of intensive glycemic control and risk factor treatment on the development of diabetic complications. Newly diagnosed individuals with type 2 DM were randomized to (1) intensive management using various combinations of insulin, a sulfonylurea, or metformin or (2) conventional therapy using dietary modification and pharmacotherapy with the goal of symptom prevention. In addition, individuals were randomly assigned to different antihypertensive regimens. Individuals in the intensive treatment arm achieved an HbA_{1c} of 7%, compared to a 7.9% HbA_{1c} in the standard treatment group. The UKPDS demonstrated that each percentage point reduction in HbA_{1c} was associated with a 35% reduction in microvascular complications. As in the DCCT, there was a continuous relationship between glycemic control and development of complications. Improved glycemic control also reduced the cardiovascular event rate in the follow-up period of >10 years.

One of the major findings of the UKPDS was that strict blood pressure control significantly reduced both macro- and microvascular complications. In fact, the beneficial effects of blood pressure control were greater than the beneficial effects of glycemic control. Lowering blood pressure to moderate goals (144/82 mmHg) reduced the risk of DM-related death, stroke, microvascular endpoints, retinopathy, and heart failure (risk reductions between 32 and 56%).

Similar reductions in the risks of retinopathy and nephropathy were also seen in a small trial of lean Japanese individuals with type 2 DM randomized to either intensive glycemic control or standard therapy with insulin (Kumamoto study). These results demonstrate the effectiveness of improved glycemic control in individuals of different ethnicity and, presumably, a different etiology of DM (i.e., phenotypically different from those in the DCCT and UKPDS). The Action to Control Cardiovascular Risk in Diabetes (ACCORD) and Action in Diabetes and Vascular Disease: Preterax and Diamicon MR Controlled Evaluation (ADVANCE) trials also found that improved glycemic control reduced microvascular complications.

Thus, these large clinical trials in type 1 and type 2 DM indicate that chronic hyperglycemia plays a causative role in the pathogenesis of diabetic microvascular complications. In both the DCCT and the UKPDS, cardiovascular events were reduced at follow-up of >10 years, even though the improved glycemic control was not maintained. The positive impact of a period of improved glycemic control on later disease has been termed a *legacy effect* or *metabolic memory*.

A summary of the features of diabetes-related complications includes the following. (1) Duration and degree of hyperglycemia correlate with complications. (2) Intensive glycemic control is beneficial in all forms of DM. (3) Blood pressure control is critical, especially in type 2 DM. (4) Survival in patients with type 1 DM is improving, and diabetes-related complications are declining. (5) Not all individuals with diabetes develop diabetes-related complications. Other incompletely defined factors appear to modulate the development of complications. For example, despite long-standing DM, some individuals never develop nephropathy or retinopathy. Many of these patients have glycemic control that is indistinguishable from those who develop microvascular complications, suggesting a genetic susceptibility for developing particular complications.

MECHANISMS OF COMPLICATIONS

Although chronic hyperglycemia is an important etiologic factor leading to complications of DM, the mechanism(s) by which it leads to such diverse cellular and organ dysfunction is unknown. An emerging hypothesis is that hyperglycemia leads to epigenetic changes (Chap. 82) that influence gene expression in affected cells. For example, this may explain the legacy effect or metabolic memory mentioned above.

Four theories, which are not mutually exclusive, on how hyperglycemia might lead to the chronic complications of DM include the following pathways. (1) Increased intracellular glucose leads to the formation of advanced glycosylation end products, which bind to a cell surface receptor, via the nonenzymatic glycosylation of intra- and extracellular proteins, leading to cross-linking of proteins, accelerated atherosclerosis, glomerular dysfunction, endothelial dysfunction, and altered extracellular matrix composition. (2) Hyperglycemia increases glucose metabolism via the sorbitol pathway related to the enzyme aldose reductase. However, testing of this theory in humans, using aldose reductase inhibitors, has not demonstrated beneficial effects. (3) Hyperglycemia increases the formation of diacylglycerol, leading to activation of protein kinase C, which alters the transcription of genes for fibronectin, type IV collagen, contractile proteins, and extracellular matrix proteins in endothelial cells and neurons. (4) Hyperglycemia increases the flux through the hexosamine pathway, which generates fructose-6-phosphate, a substrate for O-linked glycosylation and proteoglycan production, leading to altered function by glycosylation of proteins such as endothelial nitric oxide synthase or by changes in gene expression of transforming growth factor β (TGF- β) or plasminogen activator inhibitor-1.

Growth factors may play an important role in some diabetes-related complications, and their production is increased by most of these proposed pathways. Vascular endothelial growth factor A (VEGF-A) is increased locally in diabetic proliferative retinopathy and decreases after laser photocoagulation. TGF- β is increased in diabetic nephropathy and stimulates basement membrane production of collagen and fibronectin by mesangial cells. A possible unifying mechanism is that hyperglycemia leads to increased production of reactive oxygen species or superoxide in the mitochondria; these compounds may activate all four of the pathways described above. Although hyperglycemia serves as the initial trigger for complications of diabetes, it is still unknown whether the same pathophysiologic processes are operative in all complications or whether some pathways predominate in certain organs.

OPHTHALMOLOGIC COMPLICATIONS OF DIABETES MELLITUS

DM is the leading cause of blindness between the ages of 20 and 74 in the United States. The gravity of this problem is highlighted by the finding that individuals with DM are 25 times more likely to become legally blind than individuals without DM. Severe vision loss is primarily the result of progressive diabetic retinopathy and clinically significant macular edema. Diabetic retinopathy is classified into two stages: nonproliferative and proliferative. Nonproliferative diabetic retinopathy usually appears late in the first decade or early in the second decade of the disease and is marked by retinal vascular microaneurysms, blot hemorrhages, and cotton-wool spots (Fig. 419-2). Mild nonproliferative retinopathy may progress to more extensive disease, characterized by changes in venous vessel caliber, intraretinal microvascular

abnormalities, and more numerous microaneurysms and hemorrhages. The pathophysiologic mechanisms invoked in nonproliferative retinopathy include loss of retinal pericytes, increased retinal vascular permeability, alterations in retinal blood flow, and abnormal retinal microvasculature, all of which can lead to retinal ischemia. A new concept is that the pathology involves inflammatory processes in the retinal neurovascular unit, which consists of neurons, glia, astrocytes, Müller cells, and specialized vasculature.

The appearance of neovascularization in response to retinal hypoxemia is the hallmark of proliferative diabetic retinopathy (Fig. 419-2). These newly formed vessels appear near the optic nerve and/or macula and rupture easily, leading to vitreous hemorrhage, fibrosis, and ultimately retinal detachment. Not all individuals with nonproliferative retinopathy go on to develop proliferative retinopathy, but the more severe the nonproliferative disease, the greater the chance of evolution to proliferative retinopathy within 5 years. This creates an important opportunity for early detection and treatment of diabetic retinopathy. Clinically significant macular edema can occur in the context of nonproliferative or proliferative retinopathy. Fluorescein angiography and optical coherence tomography are useful to detect macular edema, which is associated with a 25% chance of moderate visual loss over the next 3 years. Duration of DM and degree of glycemic control are the best predictors of the development of retinopathy; hypertension and nephropathy are also risk factors. Nonproliferative retinopathy is found in many individuals who have had DM for >20 years. Although there is genetic susceptibility for retinopathy, it confers less influence than either the duration of DM or the degree of glycemic control.

TREATMENT DIABETIC RETINOPATHY

The most effective therapy for diabetic retinopathy is prevention. Intensive glycemic and blood pressure control will delay the development or slow the progression of retinopathy in individuals with either type 1 or type 2 DM. Paradoxically, during the first 6–12 months of improved glycemic control, established diabetic retinopathy may transiently worsen. Fortunately, this progression is temporary, and in the long term, improved glycemic control is associated with less diabetic retinopathy. Individuals with known retinopathy may be candidates for prophylactic laser photocoagulation when initiating intensive therapy. Once advanced retinopathy is present, improved glycemic control imparts less benefit, although adequate ophthalmologic care can prevent most blindness.

Regular, comprehensive eye examinations are essential for all individuals with DM (see Table 418-1). Most diabetic eye disease can be successfully treated if detected early. Routine, nondilated eye examinations by the primary care provider or diabetes specialist are inadequate to detect diabetic eye disease, which requires an ophthalmologist for optimal care of these disorders. Laser photocoagulation is very successful in preserving vision. Proliferative retinopathy is usually treated with panretinal laser photocoagulation, whereas macular edema is treated with focal laser photocoagulation and anti-vascular endothelial growth factor therapy (ocular injection). Aspirin therapy (650 mg/d) does not appear to influence the natural history of diabetic retinopathy.

RENAL COMPLICATIONS OF DIABETES MELLITUS

Diabetic nephropathy is the leading cause of chronic kidney disease (CKD), ESRD, and CKD requiring renal replacement therapy. Furthermore, the prognosis of diabetic patients on dialysis is poor, with survival comparable to many forms of cancer. Albuminuria in individuals with DM is associated with an increased risk of cardiovascular disease. Individuals with diabetic nephropathy commonly have diabetic retinopathy.

Like other microvascular complications, the pathogenesis of diabetic nephropathy is related to chronic hyperglycemia. The mechanisms by which chronic hyperglycemia leads to diabetic nephropathy, although incompletely defined, involve the effects of soluble factors (growth factors, angiotensin II, endothelin, advanced glycation end



FIGURE 419-2 Diabetic retinopathy results in scattered hemorrhages, yellow exudates, and neovascularization. This patient has neovascular vessels proliferating from the optic disc, requiring urgent panretinal laser photocoagulation.

products [AGEs]), hemodynamic alterations in the renal microcirculation (glomerular hyperfiltration or hyperperfusion, increased glomerular capillary pressure), and structural changes in the glomerulus (increased extracellular matrix, basement membrane thickening, mesangial expansion, fibrosis). Some of these effects may be mediated through angiotensin II receptors. Smoking accelerates the decline in renal function. Because only 20–40% of patients with diabetes develop diabetic nephropathy, additional genetic or environmental susceptibility factors remain unidentified. Known risk factors include race and a family history of diabetic nephropathy. Diabetic nephropathy and ESRD secondary to DM develop more commonly in African Americans, Native Americans, and Hispanic individuals with diabetes.

The natural history of diabetic nephropathy is characterized by a fairly predictable sequence of events that was initially defined for individuals with type 1 DM but appears to be similar in type 2 DM (Fig. 419-3). Glomerular hyperperfusion and renal hypertrophy occur in the first years after the onset of DM and are associated with an increase of the glomerular filtration rate (GFR). During the first 5 years of DM, thickening of the glomerular basement membrane, glomerular hypertrophy, and mesangial volume expansion occur as the GFR returns to normal. After 5–10 years of type 1 DM, many individuals begin to excrete small amounts of albumin in the urine. The American Diabetes Association (ADA) recently suggested that the terms previously used to refer to increased urinary protein (microalbuminuria as defined as 30–299 mg/d in a 24-h collection or 30–299 μ g/mg creatinine in a spot collection or macroalbuminuria as defined as >300 mg/24 h) be replaced by the phrases “persistent albuminuria (30–299 mg/24 h)” and “persistent albuminuria (\geq 300 mg/24 h)” to better reflect the continuous nature of albumin excretion in the urine as risk factor for nephropathy and cardiovascular disease (CVD). This chapter uses the terms *microalbuminuria* and *macroalbuminuria*. Although the appearance of microalbuminuria in type 1 DM is an important risk factor for progression to macroalbuminuria, only ~50% of individuals progress to macroalbuminuria over the next 10 years. In some individuals with type 1 diabetes and microalbuminuria of short duration, the microalbuminuria regresses. Microalbuminuria is also a risk factor for CVD. Once macroalbuminuria is present, there is a steady decline in GFR, and ~50% of individuals reach ESRD in 7–10 years. Once macroalbuminuria develops, blood pressure rises slightly and the pathologic changes are likely irreversible.

The nephropathy that develops in type 2 DM differs from that of type 1 DM in the following respects: (1) microalbuminuria or macroalbuminuria may be present when type 2 DM is diagnosed, reflecting its long asymptomatic period; (2) hypertension more commonly accompanies microalbuminuria or macroalbuminuria in type 2 DM; and (3) microalbuminuria may be less predictive of diabetic nephropathy and likelihood of progression to macroalbuminuria in type 2 DM, in large part due to increased CV mortality in this population. Finally, it should be noted that albuminuria in type 2 DM may be secondary to factors unrelated to DM, such as hypertension, congestive heart failure (CHF), prostate disease, or infection.

As part of comprehensive diabetes care (Chap. 418), albuminuria should be detected at an early stage when effective therapies can be instituted. Because some individuals with type 1 or type 2 DM have a decline in GFR in the absence of albuminuria, annual measurement of the serum creatinine to estimate GFR should also be performed.

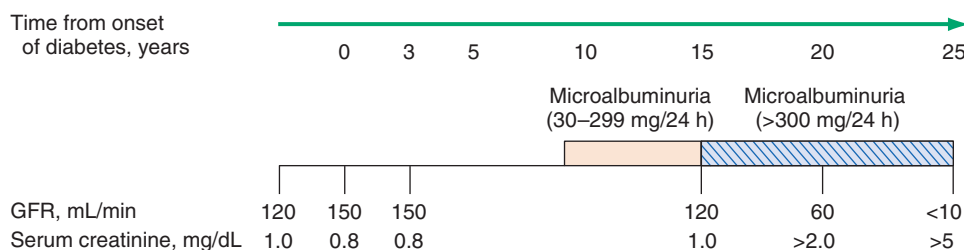


FIGURE 419-3 Time course of development of diabetic nephropathy. The relationship of time from onset of diabetes, the glomerular filtration rate (GFR), and the serum creatinine are shown. (Adapted from RA DeFranzo, in *Therapy for Diabetes Mellitus and Related Disorders*, 3rd ed. American Diabetes Association, Alexandria, VA, 1998.)

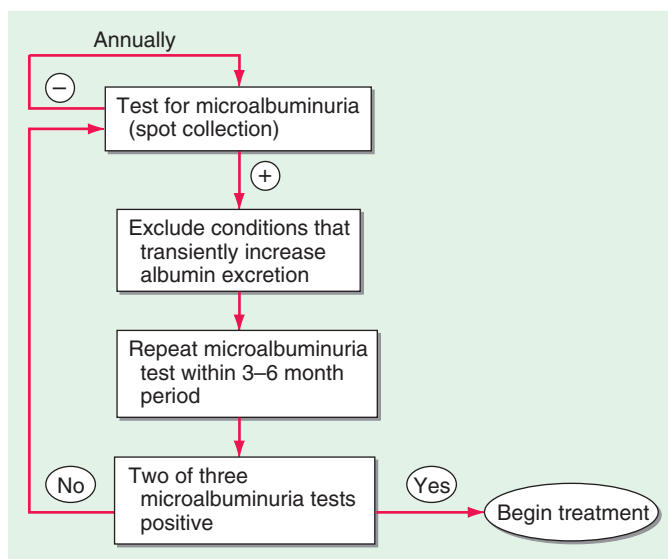


FIGURE 419-4 Screening for microalbuminuria should be performed in patients with type 1 diabetes for \geq 5 years, in patients with type 2 diabetes, and during pregnancy. Non-diabetes-related conditions that might increase microalbuminuria are urinary tract infection, hematuria, heart failure, febrile illness, severe hyperglycemia, severe hypertension, and vigorous exercise. (Adapted from RA DeFranzo, in *Therapy for Diabetes Mellitus and Related Disorders*, 3rd ed. American Diabetes Association, Alexandria, VA, 1998.)

An annual microalbuminuria measurement (albumin-to-creatinine ratio in spot urine) is advised in individuals with type 1 or type 2 DM (Fig. 419-4). The urine protein measurement in a routine urinalysis does not detect these low levels of albumin excretion. Screening for albuminuria should commence 5 years after the onset of type 1 DM and at the time of diagnosis of type 2 DM.

Type IV renal tubular acidosis (hyporeninemic hypoaldosteronism) may occur in type 1 or 2 DM. These individuals develop a propensity to hyperkalemia and acidemia, which may be exacerbated by medications (especially angiotensin-converting enzyme [ACE] inhibitors, angiotensin receptor blockers [ARBs], and spironolactone). Patients with DM are predisposed to radiocontrast-induced nephrotoxicity. Risk factors for radiocontrast-induced nephrotoxicity are preexisting nephropathy and volume depletion. Individuals with DM undergoing radiographic procedures with contrast dye should be well hydrated before and after dye exposure, and the serum creatinine should be monitored for 24–48 h following the procedure. Metformin should be held if indicated.

TREATMENT DIABETIC NEPHROPATHY

The optimal therapy for diabetic nephropathy is prevention by control of glycemia (Chap. 418 outlines **glycemic goals and approaches**). Interventions effective in slowing progression of albuminuria include (1) improved glycemic control, (2) strict blood pressure control, and (3) administration of an ACE inhibitor or ARB. Dyslipidemia should also be treated.

Improved glycemic control reduces the rate at which microalbuminuria appears and progresses in type 1 and type 2 DM. However, once macroalbuminuria is present, it is unclear whether improved glycemic control will slow progression of renal disease. During the later phase of declining renal function, insulin requirements may fall as the kidney is a site of insulin degradation. As the GFR decreases with progressive nephropathy, the use and dose of glucose-lowering agents should be reevaluated (see Table 418-5). Some glucose-lowering medications (sulfonylureas and metformin) are contraindicated in advanced renal insufficiency.

Many individuals with type 1 or type 2 DM develop hypertension. Numerous studies in both type 1 and type 2 DM demonstrate the effectiveness of strict blood pressure control in reducing albumin excretion and slowing the decline in renal function. Blood pressure should be maintained at <140/90 mmHg in diabetic individuals.

Either ACE inhibitors or ARBs should be used to reduce the albuminuria and the associated decline in GFR that accompanies it in individuals with type 1 or type 2 DM (see "Hypertension," below). Although direct comparisons of ACE inhibitors and ARBs are lacking, most experts believe that the two classes of drugs are equivalent in patient with diabetes. ARBs can be used as an alternative in patients who develop ACE inhibitor-associated cough or angioedema. After 2–3 months of therapy in patients with microalbuminuria, the drug dose is increased until the maximum tolerated dose is reached. Recent studies do not show benefit of intervention prior to onset of microalbuminuria. The combination of an ACE inhibitor and an ARB is not recommended and appears to be detrimental. If use of either ACE inhibitors or ARBs is not possible or the blood pressure is not controlled, then, diuretics, calcium channel blockers (nondihydropyridine class), or beta blockers should be used. These salutary effects are mediated by reducing intraglomerular pressure and inhibition of angiotensin-driven sclerosing pathways, in part through inhibition of TGF- β -mediated pathways.

The ADA does not suggest restriction of protein intake in diabetic individuals with albuminuria because studies have failed to show benefit.

Nephrology consultation should be considered when albuminuria appears and again when the estimated GFR is <60 mL/min per 1.743 m². As compared with nondiabetic individuals, hemodialysis in patients with DM is associated with more frequent complications, such as hypotension (due to autonomic neuropathy or loss of reflex tachycardia), more difficult vascular access, and accelerated progression of retinopathy. Complications of atherosclerosis are the leading cause of death in diabetic individuals with nephropathy and hyperlipidemia should be treated aggressively. Renal transplantation from a living related donor is the preferred therapy but requires chronic immunosuppression. Combined pancreas-kidney transplant offers the promise of normoglycemia and freedom from dialysis.

NEUROPATHY AND DIABETES MELLITUS

Diabetic neuropathy occurs in ~50% of individuals with long-standing type 1 and type 2 DM. It may manifest as polyneuropathy, mononeuropathy, and/or autonomic neuropathy. As with other complications of DM, the development of neuropathy correlates with the duration of diabetes and glycemic control. Additional risk factors are body mass index (BMI) (the greater the BMI, the greater the risk of neuropathy) and smoking. The presence of CVD, elevated triglycerides, and hypertension is also associated with diabetic peripheral neuropathy. Both myelinated and unmyelinated nerve fibers are lost. Because the clinical features of diabetic neuropathy are similar to those of other neuropathies, the diagnosis of diabetic neuropathy should be made only after other possible etiologies are excluded (Chap. 459).

Polyneuropathy/Mononeuropathy The most common form of diabetic neuropathy is distal symmetric polyneuropathy. It most frequently presents with distal sensory loss and pain, but up to 50% of patients

do not have symptoms of neuropathy. Hyperesthesia, paresthesia, and dysesthesia also may occur. Any combination of these symptoms may develop as neuropathy progresses. Symptoms may include a sensation of numbness, tingling, sharpness, or burning that begins in the feet and spreads proximally. Neuropathic pain develops in some of these individuals, occasionally preceded by improvement in their glycemic control. Pain typically involves the lower extremities, is usually present at rest, and worsens at night. Both an acute (lasting <12 months) and a chronic form of painful diabetic neuropathy have been described. The acute form is sometimes treatment-related, occurring in the context of improved glycemic control. As diabetic neuropathy progresses, the pain subsides and eventually disappears, but a sensory deficit in the lower extremities persists. Physical examination reveals sensory loss, loss of ankle deep-tendon reflexes, and abnormal position sense.

Diabetic polyradiculopathy is a syndrome characterized by severe disabling pain in the distribution of one or more nerve roots. It may be accompanied by motor weakness. Intercostal or truncal radiculopathy causes pain over the thorax or abdomen. Involvement of the lumbar plexus or femoral nerve may cause severe pain in the thigh or hip and may be associated with muscle weakness in the hip flexors or extensors (diabetic amyotrophy). Fortunately, diabetic polyradiculopathies are usually self-limited and resolve over 6–12 months.

Mononeuropathy (dysfunction of isolated cranial or peripheral nerves) is less common than polyneuropathy in DM and presents with pain and motor weakness in the distribution of a single nerve. Mononeuropathies can occur at entrapment sites such as carpal tunnel or be noncompressive. A vascular etiology for noncompressive mononeuropathies has been suggested, but the pathogenesis is unknown. Involvement of the third cranial nerve is most common and is heralded by diplopia. Physical examination reveals ptosis and ophthalmoplegia with normal pupillary constriction to light. Sometimes other cranial nerves, such as IV, VI, or VII (Bell's palsy), are affected. Peripheral mononeuropathies or simultaneous involvement of more than one nerve (mononeuropathy multiplex) may also occur.

Autonomic Neuropathy Individuals with long-standing type 1 or 2 DM may develop signs of autonomic dysfunction involving the cholinergic, noradrenergic, and peptidergic (peptides such as pancreatic polypeptide, substance P, etc.) systems. DM-related autonomic neuropathy can involve multiple systems, including the cardiovascular, gastrointestinal, genitourinary, sudomotor, and metabolic systems. Autonomic neuropathies affecting the cardiovascular system cause a resting tachycardia and orthostatic hypotension. Reports of sudden death have also been attributed to autonomic neuropathy. Gastroparesis and bladder-emptying abnormalities are often caused by the autonomic neuropathy seen in DM (discussed below). Hyperhidrosis of the upper extremities and anhidrosis of the lower extremities result from sympathetic nervous system dysfunction. Anhidrosis of the feet can promote dry skin with cracking, which increases the risk of foot ulcers. Autonomic neuropathy may reduce counterregulatory hormone release (especially catecholamines), leading to an inability to sense hypoglycemia appropriately (hypoglycemia unawareness; Chap. 420), thereby subjecting the patient to the risk of severe hypoglycemia and complicating efforts to improve glycemic control.

TREATMENT DIABETIC NEUROPATHY

Treatment of diabetic neuropathy is less than satisfactory. Improved glycemic control should be aggressively pursued and will improve nerve conduction velocity, but symptoms of diabetic neuropathy may not necessarily improve. Efforts to improve glycemic control in long-standing diabetes may be confounded by autonomic neuropathy and hypoglycemia unawareness. Risk factors for neuropathy such as hypertension and hypertriglyceridemia should be treated. Avoidance of neurotoxins (alcohol) and smoking, supplementation with vitamins for possible deficiencies (B₁₂, folate; Chap. 96e), and symptomatic treatment are the mainstays of therapy. Loss of sensation in the foot places the patient at risk for ulceration and its sequelae; consequently, prevention of

such problems is of paramount importance. Patients with symptoms or signs of neuropathy should check their feet daily and take precautions (footwear) aimed at preventing calluses or ulcerations. If foot deformities are present, a podiatrist should be involved.

Chronic, painful diabetic neuropathy is difficult to treat but may respond to duloxetine, amitriptyline, gabapentin, valproate, pregabalin, or opioids. Two agents, duloxetine and pregabalin, have been approved by the U.S. Food and Drug Administration (FDA) for pain associated with diabetic neuropathy, but no treatments are satisfactory. No direct comparisons of agents are available, and it is reasonable to switch agents if there is no response or if side effects develop. Referral to a pain management center may be necessary. Because the pain of acute diabetic neuropathy may resolve over time, medications may be discontinued as progressive neuronal damage from DM occurs.

Therapy of orthostatic hypotension secondary to autonomic neuropathy is also challenging. A variety of agents have limited success (fludrocortisone, midodrine, clonidine, octreotide, and yohimbine), but each has significant side effects. Nonpharmacologic maneuvers (adequate salt intake, avoidance of dehydration and diuretics, and lower extremity support hose) may offer some benefit.

GASTROINTESTINAL/GENITOURINARY DYSFUNCTION

Long-standing type 1 and 2 DM may affect the motility and function of the gastrointestinal (GI) and genitourinary systems. The most prominent GI symptoms are delayed gastric emptying (gastroparesis) and altered small- and large-bowel motility (constipation or diarrhea). Gastroparesis may present with symptoms of anorexia, nausea, vomiting, early satiety, and abdominal bloating. Microvascular complications (retinopathy and neuropathy) are usually present. Nuclear medicine scintigraphy after ingestion of a radiolabeled meal may document delayed gastric emptying, but may not correlate well with the patient's symptoms. Noninvasive "breath tests" following ingestion of a radiolabeled meal have been developed, but are not yet validated. Although parasympathetic dysfunction secondary to chronic hyperglycemia is important in the development of gastroparesis, hyperglycemia itself also impairs gastric emptying. Nocturnal diarrhea, alternating with constipation, is a feature of DM-related GI autonomic neuropathy. In type 1 DM, these symptoms should also prompt evaluation for celiac sprue because of its increased frequency. Esophageal dysfunction in long-standing DM may occur but is usually asymptomatic.

Diabetic autonomic neuropathy may lead to genitourinary dysfunction including cystopathy and female sexual dysfunction (reduced sexual desire, dyspareunia, reduced vaginal lubrication). Symptoms of diabetic cystopathy begin with an inability to sense a full bladder and a failure to void completely. As bladder contractility worsens, bladder capacity and the postvoid residual increase, leading to symptoms of urinary hesitancy, decreased voiding frequency, incontinence, and recurrent urinary tract infections. Diagnostic evaluation includes cystometry and urodynamic studies.

Erectile dysfunction and retrograde ejaculation are very common in DM and may be one of the earliest signs of diabetic neuropathy (Chap. 67). Erectile dysfunction, which increases in frequency with the age of the patient and the duration of diabetes, may occur in the absence of other signs of diabetic autonomic neuropathy.

TREATMENT GASTROINTESTINAL/GENITOURINARY DYSFUNCTION

Current treatments for these complications of DM are inadequate. Improved glycemic control should be a primary goal, because some aspects (neuropathy, gastric function) may improve. Smaller, more frequent meals that are easier to digest (liquid) and low in fat and fiber may minimize symptoms of gastroparesis. Metoclopramide has been used but is now restricted in both the United States and Europe and not advised for long-term use. Gastric electrical stimulatory devices are available but not approved. Diabetic diarrhea in the absence of bacterial overgrowth is treated symptomatically (Chap. 349).

Diabetic cystopathy should be treated with scheduled voiding or self-catheterization. Drugs that inhibit type 5 phosphodiesterase are effective for erectile dysfunction, but their efficacy in individuals with DM is slightly lower than in the nondiabetic population (Chap. 67). Sexual dysfunction in women may be improved with use of vaginal lubricants, treatment of vaginal infections, and systemic or local estrogen replacement.

CARDIOVASCULAR MORBIDITY AND MORTALITY

CVD is increased in individuals with type 1 or type 2 DM. The Framingham Heart Study revealed a marked increase in PAD, coronary artery disease, MI, and CHF (risk increase from one- to fivefold) in DM. In addition, the prognosis for individuals with diabetes who have coronary artery disease or MI is worse than for nondiabetics. CHD is more likely to involve multiple vessels in individuals with DM. In addition to CHD, cerebrovascular disease is increased in individuals with DM (threefold increase in stroke). Thus, after controlling for all known cardiovascular risk factors, type 2 DM increases the cardiovascular death rate twofold in men and fourfold in women.

The American Heart Association has designated DM as a "CHD risk equivalent," and type 2 DM patients without a prior MI have a similar risk for coronary artery-related events as nondiabetic individuals who have had a prior MI. However, the cardiovascular risk assessment in type 2 DM should encompass a more nuanced approach. Cardiovascular risk is lower and not equivalent in a younger individual with a brief duration of type 2 DM compared to an older individual with long-standing type 2DM. Because of the extremely high prevalence of underlying CVD in individuals with diabetes (especially in type 2 DM), evidence of atherosclerotic vascular disease (e.g., cardiac stress test) should be sought in an individual with diabetes who has symptoms suggestive of cardiac ischemia or peripheral or carotid arterial disease. The screening of asymptomatic individuals with diabetes for CHD, even with a risk-factor scale, is not recommended because recent studies have not shown a clinical benefit. The absence of chest pain ("silent ischemia") is common in individuals with diabetes, and a thorough cardiac evaluation should be considered prior to major surgical procedures.

The increase in cardiovascular morbidity and mortality rates in diabetes appears to relate to the synergism of hyperglycemia with other cardiovascular risk factors. Risk factors for macrovascular disease in diabetic individuals include dyslipidemia, hypertension, obesity, reduced physical activity, and cigarette smoking. Additional risk factors more prevalent in the diabetic population include microalbuminuria, macroalbuminuria, an elevation of serum creatinine, abnormal platelet function and endothelial dysfunction. The possibility of atherogenic potential of insulin is suggested by the data in nondiabetic individuals showing higher serum insulin levels (indicative of insulin resistance) in association with greater risk of cardiovascular morbidity and mortality. However, treatment with insulin and the sulfonylureas did not increase the risk of CVD in individuals with type 2 DM.

TREATMENT CARDIOVASCULAR DISEASE

In general, the treatment of coronary disease is not different in the diabetic individual (Chap. 293). Revascularization procedures for CHD, including percutaneous coronary interventions (PCI) and coronary artery bypass grafting (CABG), may be less efficacious in the diabetic individual. Initial success rates of PCI in diabetic individuals are similar to those in the nondiabetic population, but diabetic patients have higher rates of restenosis and lower long-term patency and survival rates in older studies.

Aggressive cardiovascular risk modification in all individuals with DM and glycemic control should be individualized, as discussed in Chap. 418. In patients with known CHD and type 2 DM, an ACE inhibitor (or ARB), a statin, and acetylsalicylic acid (ASA; aspirin) should be considered. Past trepidation about using beta blockers in individuals who have diabetes should not prevent use of these agents because they clearly benefit diabetic patients after MI. In

2428 patients with CHF, thiazolidinediones should not be used (**Chap. 418**). However, metformin can be used in patients with stable CHF if the renal function is normal.

Antiplatelet therapy reduces cardiovascular events in individuals with DM who have CHD and is recommended. Current recommendations by the ADA include the use of aspirin for primary prevention of coronary events in diabetic individuals with an increased 10-year cardiovascular risk >10% (at least one risk factor such as hypertension, smoking, family history, albuminuria, or dyslipidemia in men >50 years or women >60 years of age). ASA is not recommended for primary prevention in those with a 10-year cardiovascular risk <10%. The aspirin dose is the same as in nondiabetic individuals.

Cardiovascular Risk Factors • DYSLIPIDEMIA Individuals with DM may have several forms of dyslipidemia (**Chap. 421**). Because of the additive cardiovascular risk of hyperglycemia and hyperlipidemia, lipid abnormalities should be assessed aggressively and treated as part of comprehensive diabetes care (**Chap. 418**). The most common pattern of dyslipidemia is hypertriglyceridemia and reduced high-density lipoprotein (HDL) cholesterol levels. DM itself does not increase levels of low-density lipoprotein (LDL), but the small dense LDL particles found in type 2 DM are more atherogenic because they are more easily glycosylated and susceptible to oxidation.

Almost all treatment studies of diabetic dyslipidemia have been performed in individuals with type 2 DM because of the greater frequency of dyslipidemia in this form of diabetes. Interventional studies have shown that the beneficial effects of LDL reduction with statins are similar in the diabetic and nondiabetic populations. Large prospective trials of primary and secondary intervention for CHD have included some individuals with type 2 DM, and subset analyses have consistently found that reductions in LDL reduce cardiovascular events and morbidity in individuals with DM. No prospective studies have addressed similar questions in individuals with type 1 DM. Because the frequency of CVD is low in children and young adults with diabetes, assessment of cardiovascular risk should be incorporated into the guidelines discussed below.

Based on the guidelines provided by the ADA, priorities in the treatment of dyslipidemia are as follows: (1) lower the LDL cholesterol, (2) raise the HDL cholesterol, and (3) decrease the triglycerides. A treatment strategy depends on the pattern of lipoprotein abnormalities. Initial therapy for all forms of dyslipidemia should include dietary changes, as well as the same lifestyle modifications recommended in the nondiabetic population (smoking cessation, blood pressure control, weight loss, increased physical activity). The dietary recommendations for individuals with DM include increased monounsaturated fat and carbohydrates and reduced saturated fats and cholesterol (**Chap. 421**). According to guidelines of the ADA, the target lipid values in diabetic individuals (age >40 years) without CVD should be as follows: LDL <2.6 mmol/L (100 mg/dL); HDL >1 mmol/L (40 mg/dL) in men and >13 mmol/L (50 mg/dL) in women; and triglycerides <1.7 mmol/L (150 mg/dL). In patients >40 years, the ADA recommends addition of a statin, regardless of the LDL level, in patients with CHD and those without CHD who have CHD risk factors. Recently released guidelines by the American College of Cardiology (ACC) and American Heart Association (AHA) differ slightly and recommend that diabetic individuals aged 40–75 without CHD and a LDL of 70–189 mg/dl receive “moderate” intensity statin therapy (**Chap. 291e**). Improvement in glycemic control will lower triglycerides and have a modest beneficial effect by raising HDL.

If the patient is known to have CHD, the ADA recommends an LDL goal of <18 mmol/L (70 mg/dL) as an “option” (in keeping with evidence that such a goal is beneficial in nondiabetic individuals with CHD [**Chap. 421**]). The ACC/AHA guidelines do not advocate a specific LDL for statin therapy. HMG-CoA reductase inhibitors are the agents of choice for lowering LDL. Combination therapy with an HMG-CoA reductase inhibitor and a fibrate or another lipid-lowering agent (ezetimibe, niacin) may be considered but increases the possibility of side effects such as myositis and has not been shown to be

beneficial. Nicotinic acid effectively raises HDL and can be used in patients with diabetes, but may worsen glycemic control and increase insulin resistance and has not been shown to provide additional benefit beyond statin therapy alone. Bile acid-binding resins should not be used if hypertriglyceridemia is present. In large clinical trials, statin usage is associated with a mild increase in the risk of developing type 2 DM. This risk is greatest in individuals with other risk factors for type 2 DM (**Chap. 417**). However, the cardiovascular benefits of statin use outweigh the mildly increased risk of diabetes.

HYPERTENSION Hypertension can accelerate other complications of DM, particularly CVD, nephropathy, and retinopathy. In targeting a goal of blood pressure of <140/80 mmHg, therapy should first emphasize lifestyle modifications such as weight loss, exercise, stress management, and sodium restriction. The BP goal should be individualized. In some younger individuals, the provider may target a blood pressure of <130/80 mmHg. Realizing that more than one agent is usually required to reach the blood pressure goal, the ADA recommends that all patients with diabetes and hypertension be treated with an ACE inhibitor or an ARB. Subsequently, agents that reduce cardiovascular risk (beta blockers, thiazide diuretics, and calcium channel blockers) should be incorporated into the regimen. ACE inhibitors and ARBs are likely equivalent in most patients with diabetes and renal disease. Serum potassium and renal function should be monitored.

Because of the high prevalence of atherosclerotic disease in individuals with type 2 DM, the possibility of renovascular hypertension should be considered when the blood pressure is not readily controlled.

LOWER EXTREMITY COMPLICATIONS

DM is the leading cause of nontraumatic lower extremity amputation in the United States. Foot ulcers and infections are also a major source of morbidity in individuals with DM. The reasons for the increased incidence of these disorders in DM involve the interaction of several pathogenic factors: neuropathy, abnormal foot biomechanics, PAD, and poor wound healing. The peripheral sensory neuropathy interferes with normal protective mechanisms and allows the patient to sustain major or repeated minor trauma to the foot, often without knowledge of the injury. Disordered proprioception causes abnormal weight bearing while walking and subsequent formation of callus or ulceration. Motor and sensory neuropathy lead to abnormal foot muscle mechanics and to structural changes in the foot (hammer toe, claw toe deformity, prominent metatarsal heads, Charcot joint). Autonomic neuropathy results in anhidrosis and altered superficial blood flow in the foot, which promote drying of the skin and fissure formation. PAD and poor wound healing impede resolution of minor breaks in the skin, allowing them to enlarge and to become infected.

Many individuals with type 2 DM develop a foot ulcer (great toe or metatarsophalangeal areas are most common), and a significant subset who develop an ulceration will ultimately undergo amputation (14–24% risk with that ulcer or subsequent ulceration). Risk factors for foot ulcers or amputation include male sex, diabetes for >10 years, peripheral neuropathy, abnormal structure of foot (bony abnormalities, callus, thickened nails), PAD, smoking, history of previous ulcer or amputation, visual impairment, and poor glycemic control. Large calluses are often precursors to or overlies ulcerations.

TREATMENT LOWER EXTREMITY COMPLICATIONS

The optimal therapy for foot ulcers and amputations is prevention through identification of high-risk patients, education of the patient, and institution of measures to prevent ulceration. High-risk patients should be identified during the routine, annual foot examination performed on all patients with DM (see “Ongoing Aspects of Comprehensive Diabetes Care” in **Chap. 418**). If the monofilament test or one of the other tests is abnormal, the patient is diagnosed with loss of protective sensation (LOPS; **Chap. 417**). Providers should consider screening for asymptomatic PAD in individuals >50 years of age who have diabetes and other risk factors using ankle-brachial index testing in high-risk individuals (**Chap. 302**). Patient

education should emphasize (1) careful selection of footwear, (2) daily inspection of the feet to detect early signs of poor-fitting footwear or minor trauma, (3) daily foot hygiene to keep the skin clean and moist, (4) avoidance of self-treatment of foot abnormalities and high-risk behavior (e.g., walking barefoot), and (5) prompt consultation with a health care provider if an abnormality arises. Patients at high risk for ulceration or amputation may benefit from evaluation by a foot care specialist. Calluses and nail deformities should be treated by a podiatrist. Interventions directed at risk factor modification include orthotic shoes and devices, callus management, nail care, and prophylactic measures to reduce increased skin pressure from abnormal bony architecture. Attention to other risk factors for vascular disease (smoking, dyslipidemia, hypertension) and improved glycemic control are also important.

Despite preventive measures, foot ulceration and infection are common and represent a serious problem. Due to the multifactorial pathogenesis of lower extremity ulcers, management of these lesions is multidisciplinary and often demands expertise in orthopedics, vascular surgery, endocrinology, podiatry, and infectious diseases. The plantar surface of the foot is the most common site of ulceration. Ulcers may be primarily neuropathic (no accompanying infection) or may have surrounding cellulitis or osteomyelitis. Cellulitis without ulceration is also frequent and should be treated with antibiotics that provide broad-spectrum coverage, including anaerobes (see below).

An infected ulcer is a clinical diagnosis, because superficial culture of any ulceration will likely find multiple possible bacterial species. The infection surrounding the foot ulcer is often the result of multiple organisms, with aerobic gram-positive cocci (staphylococci including MRSA, Group A and B streptococci) being most common and with aerobic gram-negative bacilli and/or obligate anaerobes as co-pathogens.

Gas gangrene may develop in the absence of clostridial infection. Cultures taken from the surface of the ulcer are not helpful; a culture from the debrided ulcer base or from purulent drainage or aspiration of the wound is the most helpful. Wound depth should be determined by inspection and probing with a blunt-tipped sterile instrument. Plain radiographs of the foot should be performed to assess the possibility of osteomyelitis in chronic ulcers that have not responded to therapy. Magnetic resonance imaging (MRI) is the most specific modality, with nuclear medicine scans and labeled white cell studies as alternatives. Surgical debridement is often necessary.

Osteomyelitis is best treated by a combination of prolonged antibiotics (IV, then oral) and/or possibly debridement of infected bone. The possible contribution of vascular insufficiency should be considered in all patients. Peripheral arterial bypass procedures are often effective in promoting wound healing and in decreasing the need for amputation of the ischemic limb ([Chap. 302](#)).

A consensus statement from the ADA identified six interventions with demonstrated efficacy in diabetic foot wounds: (1) off-loading, (2) debridement, (3) wound dressings, (4) appropriate use of antibiotics, (5) revascularization, and (6) limited amputation. Off-loading is the complete avoidance of weight bearing on the ulcer, which removes the mechanical trauma that retards wound healing. Bed rest and a variety of orthotic devices or contact casting limit weight bearing on wounds or pressure points. Surgical debridement is important and effective, but clear efficacy of other modalities for wound cleaning (enzymes, soaking, whirlpools) is lacking. Dressings such as hydrocolloid dressings promote wound healing by creating a moist environment and protecting the wound. Antiseptic agents should be avoided. Topical antibiotics are of limited value. Referral for physical therapy, orthotic evaluation, and rehabilitation should occur once the infection is controlled.

Mild or non-limb-threatening infections can be treated with oral antibiotics directed predominantly at methicillin-susceptible staphylococci and streptococci (e.g., dicloxacillin, cephalosporin, amoxicillin/clavulanate). However the increasing prevalence of MRSA often requires the use of clindamycin, doxycycline, or trimethoprim-sulfamethoxazole. Trimethoprim-sulfamethoxazole exhibits

less reliable coverage of streptococci than the β -lactams, and diabetic patients may develop adverse effects including acute kidney injury and hyperkalemia. Surgical debridement of necrotic tissue, local wound care (avoidance of weight bearing over the ulcer), and close surveillance for progression of infection are crucial. More severe infections require IV antibiotics as well as bed rest and local wound care. Urgent surgical debridement may be required. Optimization of glycemic control should be a goal. IV antibiotics should provide broad-spectrum coverage directed toward *Staphylococcus aureus*, including MRSA, streptococci, gram-negative aerobes, and anaerobic bacteria. Initial antimicrobial regimens include vancomycin plus a β -lactam/ β -lactamase inhibitor or carbapenem or vancomycin plus a combination of quinolone plus metronidazole. Daptomycin, ceftaroline, or linezolid may be substituted for vancomycin. If the infection surrounding the ulcer is not improving with IV antibiotics, reassessment of antibiotic coverage and reconsideration of the need for surgical debridement or revascularization are indicated. With clinical improvement, oral antibiotics and local wound care can be continued on an outpatient basis with close follow-up.

INFECTIONS

Individuals with DM have a greater frequency and severity of infection. The reasons for this include incompletely defined abnormalities in cell-mediated immunity and phagocyte function associated with hyperglycemia, as well as diminished vascularization. Hyperglycemia aids the colonization and growth of a variety of organisms (*Candida* and other fungal species). Many common infections are more frequent and severe in the diabetic population, whereas several rare infections are seen almost exclusively in the diabetic population. Examples of this latter category include rhinocerebral mucormycosis, emphysematous infections of the gallbladder and urinary tract, and “malignant” or invasive otitis externa. Invasive otitis externa is usually secondary to *P. aeruginosa* infection in the soft tissue surrounding the external auditory canal, usually begins with pain and discharge, and may rapidly progress to osteomyelitis and meningitis. These infections should be sought, in particular, in patients presenting with severe hyperglycemia ([Chap. 418](#)).

Pneumonia, urinary tract infections, and skin and soft tissue infections are all more common in the diabetic population. In general, the organisms that cause pulmonary infections are similar to those found in the nondiabetic population; however, gram-negative organisms, *S. aureus*, and *Mycobacterium tuberculosis* are more frequent pathogens. Urinary tract infections (either lower tract or pyelonephritis) are the result of common bacterial agents such as *Escherichia coli*, although several yeast species (*Candida* and *Torulopsis glabrata*) are commonly observed. Complications of urinary tract infections include emphysematous pyelonephritis and emphysematous cystitis. Bacteriuria occurs frequently in individuals with diabetic cystopathy. Susceptibility to furunculosis, superficial candidal infections, and vulvovaginitis are increased. Poor glycemic control is a common denominator in individuals with these infections. Diabetic individuals have an increased rate of colonization of *S. aureus* in the skinfolds and nares. Diabetic patients also have a greater risk of postoperative wound infections.

DERMATOLOGIC MANIFESTATIONS

The most common skin manifestations of DM are xerosis and pruritus and are usually relieved by skin moisturizers. Protracted wound healing and skin ulcerations are also frequent complications. Diabetic dermopathy, sometimes termed *pigmented pretibial papules*, or “diabetic skin spots,” begins as an erythematous macule or papule that evolves into an area of circular hyperpigmentation. These lesions result from minor mechanical trauma in the pretibial region and are more common in elderly men with DM. Bullous diseases, such as bullosa diabeticorum (shallow ulcerations or erosions in the pretibial region), are also seen. *Necrobiosis lipoidica diabeticorum* is an uncommon disorder, accompanying diabetes in predominantly young women. This usually begins in the pretibial region as an erythematous plaque or papules that gradually enlarge, darken, and develop irregular margins,

2430 with atrophic centers and central ulceration. They are often painful. Vitiligo occurs at increased frequency in individuals with type 1 DM. *Acanthosis nigricans* (hyperpigmented velvety plaques seen on the neck, axilla, or extensor surfaces) is sometimes a feature of severe insulin resistance and accompanying diabetes. Generalized or localized *granuloma annulare* (erythematous plaques on the extremities or trunk) and *scleredema* (areas of skin thickening on the back or neck at the site of previous superficial infections) are more common in the diabetic population. *Lipoatrophy* and *lipohypertrophy* can occur at insulin injection sites but are now unusual with the use of human insulin.

420 Hypoglycemia

Philip E. Cryer, Stephen N. Davis

Hypoglycemia is most commonly caused by drugs used to treat diabetes mellitus or by exposure to other drugs, including alcohol. However, a number of other disorders, including critical organ failure, sepsis and inanition, hormone deficiencies, non- β -cell tumors, insulinoma, and prior gastric surgery, can cause hypoglycemia (Table 420-1). Hypoglycemia is most convincingly documented by *Whipple's triad*: (1) symptoms consistent with hypoglycemia, (2) a low plasma glucose concentration measured with a precise method (not a glucose monitor), and (3) relief of symptoms after the plasma glucose level is raised. The lower limit of the fasting plasma glucose concentration is normally ~ 70 mg/dL (~ 3.9 mmol/L), but lower venous glucose levels occur normally, late after a meal, during pregnancy, and during prolonged fasting (>24 h). Hypoglycemia can cause serious morbidity; if severe and prolonged, it can be fatal. It should be considered in any patient with episodes of confusion, an altered level of consciousness, or a seizure.

SYSTEMIC GLUCOSE BALANCE AND GLUCOSE COUNTERREGULATION

Glucose is an obligate metabolic fuel for the brain under physiologic conditions. The brain cannot synthesize glucose or store more than a few minutes' supply as glycogen and therefore requires a continuous supply of glucose from the arterial circulation. As the arterial plasma glucose concentration falls below the physiologic range, blood-to-brain glucose transport becomes insufficient to support brain energy metabolism and function. However, redundant glucose counterregulatory mechanisms normally prevent or rapidly correct hypoglycemia.

Plasma glucose concentrations are normally maintained within a relatively narrow range—roughly 70–110 mg/dL (3.9–6.1 mmol/L) in the fasting state, with transient higher excursions after a meal—despite wide variations in exogenous glucose delivery from meals and in endogenous glucose utilization by, for example, exercising muscle. Between meals and during fasting, plasma glucose levels are maintained by endogenous glucose production, hepatic glycogenolysis, and hepatic (and renal) gluconeogenesis (Fig. 420-1). Although hepatic glycogen stores are usually sufficient to maintain plasma glucose levels for ~ 8 h, this period can be shorter if glucose demand is increased by exercise or if glycogen stores are depleted by illness or starvation.

Gluconeogenesis normally requires low insulin levels and the presence of anti-insulin (counterregulatory) hormones together with a coordinated supply of precursors from muscle and adipose tissue to the liver (and kidneys). Muscle provides lactate, pyruvate, alanine, glutamine, and other amino acids. Triglycerides in adipose tissue are broken down into fatty acids and glycerol, which is a gluconeogenic precursor. Fatty acids provide an alternative oxidative fuel to tissues other than the brain (which requires glucose).

Systemic glucose balance—maintenance of the normal plasma glucose concentration—is accomplished by a network of hormones, neural signals, and substrate effects that regulate endogenous glucose production and glucose utilization by tissues other than the brain (Chap. 417). Among the regulatory factors, insulin plays a dominant role (Table 420-2; Fig. 420-1). As plasma glucose levels decline within

TABLE 420-1 CAUSES OF HYPOGLYCEMIA IN ADULTS

Ill or medicated individual

1. Drugs
 - Insulin or insulin secretagogue
 - Alcohol
 - Others
2. Critical illness
 - Hepatic, renal or cardiac failure
 - Sepsis
 - Inanition
3. Hormone deficiency
 - Cortisol
 - Glucagon and epinephrine (in insulin-deficient diabetes)
4. Non-islet cell tumor

Seemingly well individual

5. Endogenous hyperinsulinism
 - Insulinoma
 - Functional β -cell disorders (nesidioblastosis)
 - Noninsulinoma pancreatogenous hypoglycemia
 - Post-gastric bypass hypoglycemia
 - Insulin autoimmune hypoglycemia
 - Antibody to insulin
 - Antibody to insulin receptor
 - Insulin secretagogue
 - Other
6. Accidental, surreptitious, or malicious hypoglycemia

Source: From PE Cryer et al: J Clin Endocrinol Metab 94:709, 2009. ©The Endocrine Society, 2009.

the physiologic range in the fasting state, pancreatic β -cell insulin secretion decreases, thereby increasing hepatic glycogenolysis and hepatic (and renal) gluconeogenesis. Low insulin levels also reduce glucose utilization in peripheral tissues, inducing lipolysis and proteolysis and consequently releasing gluconeogenic precursors. Thus, a decrease in insulin secretion is the first defense against hypoglycemia.

As plasma glucose levels decline just below the physiologic range, glucose counterregulatory (plasma glucose-raising) hormones are released (Table 420-2; Fig. 420-1). Among these, pancreatic α -cell glucagon, which stimulates hepatic glycogenolysis, plays a primary role. Glucagon is the second defense against hypoglycemia. Adrenomedullary epinephrine, which stimulates hepatic glycogenolysis and gluconeogenesis (and renal gluconeogenesis), is not normally critical. However, it becomes critical when glucagon is deficient. Epinephrine is the third defense against hypoglycemia. When hypoglycemia is prolonged beyond ~ 4 h, cortisol and growth hormone also support glucose production and restrict glucose utilization to a limited amount ($\sim 20\%$ compared to epinephrine). Thus cortisol and growth hormone play no role in defense against acute hypoglycemia.

As plasma glucose levels fall further, symptoms prompt behavioral defense against hypoglycemia, including the ingestion of food (Table 420-2; Fig. 420-1). The normal glycemic thresholds for these responses to decreasing plasma glucose concentrations are shown in Table 420-2. However, these thresholds are dynamic. They shift to higher-than-normal glucose levels in people with poorly controlled diabetes, who can experience symptoms of hypoglycemia when their glucose levels decline toward the normal range (*pseudohypoglycemia*). On the other hand, thresholds shift to lower-than-normal glucose levels in people with recurrent hypoglycemia; e.g., patients with aggressively treated diabetes or an insulinoma have symptoms at glucose levels lower than those that cause symptoms in healthy individuals.

Clinical Manifestations Neuroglycopenic manifestations of hypoglycemia are the direct result of central nervous system glucose deprivation. These features include behavioral changes, confusion, fatigue, seizure, loss of consciousness, and, if hypoglycemia is severe and prolonged,

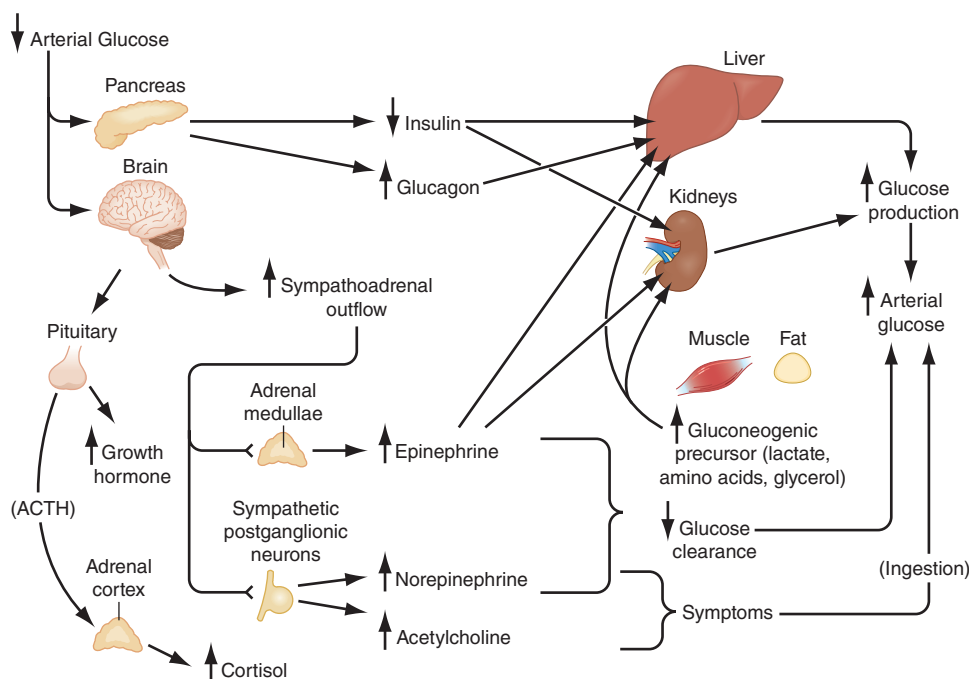


FIGURE 420-1 Physiology of glucose counterregulation: mechanisms that normally prevent or rapidly correct hypoglycemia. In insulin-deficient diabetes, the key counterregulatory responses—suppression of insulin and increases in glucagon—are lost, and stimulation of sympathoadrenal outflow is attenuated. ACTH, adrenocorticotropic hormone.

death. Neurogenic (or autonomic) manifestations of hypoglycemia result from the perception of physiologic changes caused by the central nervous system–mediated sympathoadrenal discharge that is triggered by hypoglycemia. They include *adrenergic* symptoms (mediated largely by norepinephrine released from sympathetic postganglionic neurons but perhaps also by epinephrine released from the adrenal medullae), such as palpitations, tremor, and anxiety, as well as *cholinergic* symptoms (mediated by acetylcholine released from sympathetic postganglionic neurons), such as sweating, hunger, and paresthesias. Clearly, these are nonspecific symptoms. Their attribution to hypoglycemia requires that the corresponding plasma glucose concentration be low and that the symptoms resolve after the glucose level is raised (as delineated by Whipple’s triad).

Common signs of hypoglycemia include diaphoresis and pallor. Heart rate and systolic blood pressure are typically increased but may not be raised in an individual who has experienced repeated, recent episodes of hypoglycemia. Neuroglycopenic manifestations are often observable. Transient focal neurologic deficits occur occasionally. Permanent neurologic deficits are rare.

Etiology and Pathophysiology Hypoglycemia is most commonly a result of the treatment of diabetes. This topic is therefore addressed before other causes of hypoglycemia are considered.

HYPOGLYCEMIA IN DIABETES

Impact and Frequency Hypoglycemia is the limiting factor in the glycemic management of diabetes mellitus. First, it causes recurrent morbidity in most people with type 1 diabetes (T1DM) and in many with advanced type 2 diabetes (T2DM), and it is sometimes fatal. Second, it precludes maintenance of euglycemia over a lifetime of diabetes and thus full realization of the well-established microvascular benefits of glycemic control. Third, it causes a vicious cycle of recurrent hypoglycemia by producing hypoglycemia-associated autonomic failure—i.e., the clinical syndromes of defective glucose counterregulation and of hypoglycemia unawareness (see later).

Hypoglycemia is a fact of life for people with T1DM. They suffer an average of two episodes of symptomatic hypoglycemia per week and at least one episode of severe, at least temporarily disabling hypoglycemia each year. An estimated 6–10% of people with T1DM die as a result

TABLE 420-2 PHYSIOLOGIC RESPONSES TO DECREASING PLASMA GLUCOSE CONCENTRATIONS

Response	Glycemic Threshold, mmol/L (mg/dL)	Physiologic Effects	Role in Prevention or Correction of Hypoglycemia (Glucose Counterregulation)
↓ Insulin	4.4–4.7 (80–85)	↑ R_a (↓ R_d)	Primary glucose regulatory factor/first defense against hypoglycemia
↑ Glucagon	3.6–3.9 (65–70)	↑ R_a	Primary glucose counterregulatory factor/second defense against hypoglycemia
↑ Epinephrine	3.6–3.9 (65–70)	↑ R_a , ↓ R_c	Third defense against hypoglycemia, critical when glucagon is deficient
↑ Cortisol and growth hormone	3.6–3.9 (65–70)	↑ R_a , ↓ R_c	Involved in defense against prolonged hypoglycemia; not critical
Symptoms	2.8–3.1 (50–55)	Recognition of hypoglycemia	Prompt behavioral defense against hypoglycemia (food ingestion)
↓ Cognition	<2.8 (<50)	—	Compromises behavioral defense against hypoglycemia

Note: R_a , rate of glucose appearance, glucose production by the liver and kidneys; R_d , rate of glucose disappearance, glucose utilization by insulin-sensitive tissues; R_c , rate of glucose clearance, glucose utilization by insulin-sensitive tissues such as skeletal muscle. R_d by the brain is not altered by insulin, glucagon, epinephrine, cortisol, or growth hormone.

Source: From PE Cryer, in S Melmed et al (eds): *Williams Textbook of Endocrinology*, 12th ed. New York, Elsevier, 2012.

2432 of hypoglycemia. The incidence of hypoglycemia is lower in T2DM than in T1DM. However, its prevalence in insulin-requiring T2DM is surprisingly high. Recent studies investigating insulin-pump or multiple-injection therapies have revealed a hypoglycemia prevalence approaching 70%. In fact, as patients with T2DM outnumber those with T1DM by ten- to twentyfold, the prevalence of hypoglycemia is now greater in T2DM. Insulin, a sulfonylurea, or a glinide can cause hypoglycemia in T2DM. Metformin, thiazolidinediones, α -glucosidase inhibitors, glucagon-like peptide 1 (GLP-1) receptor agonists, and dipeptidyl peptidase IV (DPP-IV) inhibitors should not cause hypoglycemia. However, they increase the risk when combined with one of the sulfonylureas or glinides, or with insulin. Notably, the frequency of hypoglycemia approaches that in T1DM as persons with T2DM develop absolute insulin deficiency and require more complex treatment with insulin.

Conventional Risk Factors The conventional risk factors for hypoglycemia in diabetes are identified on the basis of the premise that relative or absolute insulin excess is the sole determinant of risk. Relative or absolute insulin excess occurs when (1) insulin (or insulin secretagogue) doses are excessive, ill-timed, or of the wrong type; (2) the influx of exogenous glucose is reduced (e.g., during an overnight fast or after missed meals or snacks); (3) insulin-independent glucose utilization is increased (e.g., during exercise); (4) sensitivity to insulin is increased (e.g., with improved glycemic control, in the middle of the night, late after exercise, or with increased fitness or weight loss); (5) endogenous glucose production is reduced (e.g., after alcohol ingestion); and (6) insulin clearance is reduced (e.g., in renal failure). However, these conventional risk factors alone explain a minority of episodes; other factors are typically involved.

Hypoglycemia-Associated Autonomic Failure (HAAF) While marked insulin excess alone can cause hypoglycemia, iatrogenic hypoglycemia in diabetes is typically the result of the interplay of relative or absolute therapeutic insulin excess and compromised physiologic and behavioral defenses against falling plasma glucose concentrations (Table 420-2; Fig. 420-2). Defective glucose counterregulation compromises physiologic defense (particularly decrements in insulin and increments in glucagon and epinephrine), and hypoglycemia unawareness compromises behavioral defense (ingestion of carbohydrate).

DEFECTIVE GLUCOSE COUNTERREGULATION In the setting of absolute endogenous insulin deficiency, insulin levels do not decrease as plasma glucose levels fall; the first defense against hypoglycemia is lost. Furthermore, probably because the decrement in intraislet insulin is normally a signal to stimulate glucagon secretion, glucagon levels do not increase as plasma glucose levels fall further; a second defense against hypoglycemia is lost. Finally, the increase in epinephrine levels, a third defense against hypoglycemia, in response to a given level of hypoglycemia is typically attenuated. The glycemic threshold for the sympathoadrenal (adrenomedullary epinephrine and sympathetic neural norepinephrine) response is shifted to lower plasma glucose concentrations. That shift is typically the result of recent antecedent iatrogenic hypoglycemia. In the setting of absent decrements in insulin and of absent increments in glucagon, the attenuated increment in epinephrine causes the clinical syndrome of defective glucose counterregulation. Affected patients are at ≥ 25 -fold greater risk of severe iatrogenic hypoglycemia during aggressive glycemic therapy for their diabetes than are patients with normal epinephrine responses. This functional—and potentially reversible—disorder is distinct from classic diabetic autonomic neuropathy—a structural and irreversible disorder.

HYPOGLYCEMIA UNAWARENESS The attenuated sympathoadrenal response (largely the reduced sympathetic neural response) to hypoglycemia causes the clinical syndrome of *hypoglycemia unawareness*—i.e., loss of the warning adrenergic and cholinergic symptoms that previously allowed the patient to recognize developing hypoglycemia and therefore to abort the episode by ingesting carbohydrates. Affected patients are at a sixfold increased risk of severe iatrogenic hypoglycemia during aggressive glycemic therapy of their diabetes.

HAAF IN DIABETES The concept of HAAF in diabetes posits that recent antecedent iatrogenic hypoglycemia (or sleep or prior exercise) causes both defective glucose counterregulation (by reducing the epinephrine response to a given level of subsequent hypoglycemia in the setting of absent insulin and glucagon responses) and hypoglycemia unawareness (by reducing the sympathoadrenal response to a given level of subsequent hypoglycemia). These impaired responses create a vicious cycle of recurrent iatrogenic hypoglycemia (Fig. 420-2). Hypoglycemia unawareness and, to some extent, the reduced epinephrine component of defective glucose counterregulation are reversible by as little as 2–3 weeks of scrupulous avoidance of hypoglycemia in most affected patients.

On the basis of this pathophysiology, additional risk factors for hypoglycemia in diabetes include (1) absolute insulin deficiency, indicating that insulin levels will not decrease and glucagon levels will not increase as plasma glucose levels fall; (2) a history of severe hypoglycemia or of hypoglycemia unawareness, implying recent antecedent hypoglycemia, as well as prior exercise or sleep, indicating that the sympathoadrenal response will be attenuated; and (3) lower hemoglobin A_{1c} (HbA_{1c}) levels or lower glycemic goals that, all other factors being equal, increase the probability of recent antecedent hypoglycemia.

Hypoglycemia Risk Factor Reduction Several recent multicenter, randomized, controlled trials investigating the potential benefits of tight

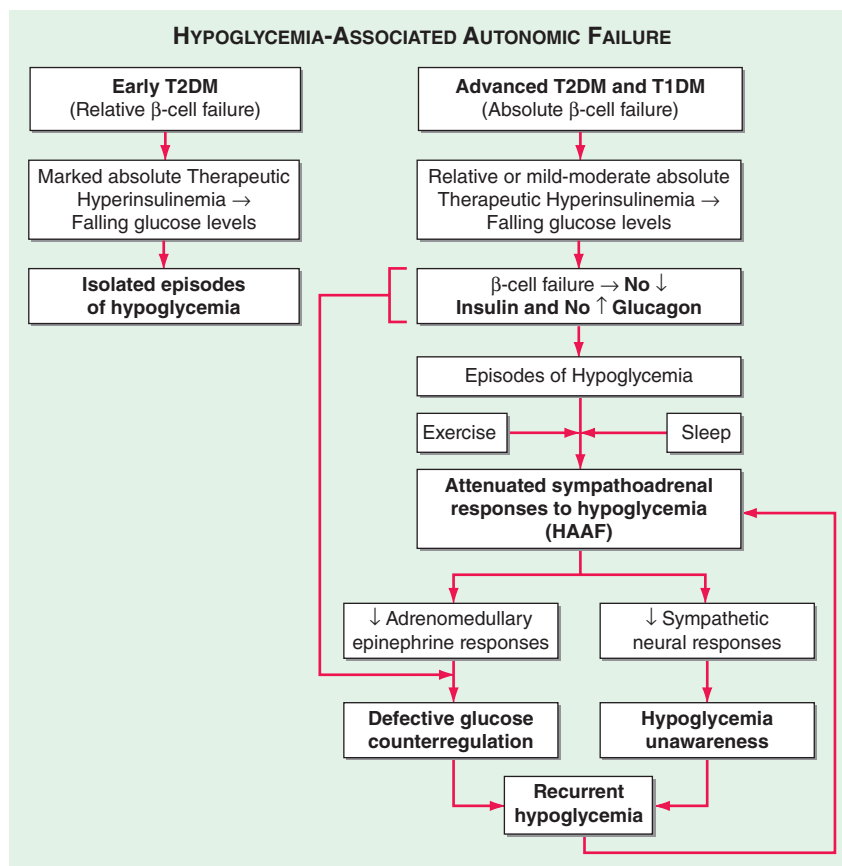


FIGURE 420-2 Hypoglycemia-associated autonomic failure (HAAF) in insulin-deficient diabetes. T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus. (Modified from PE Cryer: *Hypoglycemia in Diabetes. Pathophysiology, Prevalence, and Prevention*, 2nd ed. © American Diabetes Association, 2012.)

glucose control in either inpatient or outpatient settings have reported a high prevalence of severe hypoglycemia. In the NICE-SUGAR study, attempts to control in-hospital plasma glucose values towards physiologic levels resulted in increased mortality risk. The ADVANCE and ACCORD studies and the Veterans Affairs Diabetes Trial (VADT) also found a significant incidence of severe hypoglycemia among T2DM patients. Severe hypoglycemia with accompanying serious cardiovascular morbidity and mortality also occurred in the standard (e.g., not receiving intensified treatment) control group in both the ACCORD study and the VADT. Thus, severe hypoglycemia can and does occur at HbA_{1c} values of 8–9% in both T1DM and T2DM. Somewhat surprisingly, all three studies found little or no benefit of intensive glucose control to reduce macrovascular events in T2DM. In fact, the ACCORD study was ended early because of the increased mortality rate in the intensive glucose control arm. Whether iatrogenic hypoglycemia was the cause of the increased mortality risk is not known. In light of these findings, some new recommendations and paradigms have been formulated. Whereas there is little debate regarding the need to reduce hyperglycemia in the hospital, the glycemic maintenance goals have been modified to lie between 140 and 180 mg/dL. Accordingly, the benefits of insulin therapy and reduced hyperglycemia can be obtained while the prevalence of hypoglycemia is reduced.

Similarly, evidence exists that intensive glucose control can reduce the prevalence of microvascular disease in both T1DM and T2DM. These benefits need to be weighed against the increased prevalence of hypoglycemia. Certainly, the level of glucose control (i.e., the HbA_{1c} level) should be evaluated for each patient. Multicenter trials have demonstrated that individuals with recently diagnosed T1DM or T2DM can have better glycemic control with less hypoglycemia. In addition, there is still long-term benefit in reducing HbA_{1c} values from higher to lower, albeit still above recommended levels. Perhaps a reasonable therapeutic goal is the lowest HbA_{1c} level that does not cause severe hypoglycemia and that preserves awareness of hypoglycemia.

Pancreatic transplantation (both whole-organ and islet-cell) has been used in part as a treatment for severe hypoglycemia. Generally, rates of hypoglycemia are reduced after transplantation. This decrease appears to be due to increased physiologic insulin and glucagon responses during hypoglycemia.

The use of continuous glucose monitors offers some promise as a method of reducing hypoglycemia while improving HbA_{1c}. Other interventions to stimulate counterregulatory responses, such as selective serotonin-reuptake inhibitors, β -adrenergic receptor antagonists, opiate receptor antagonists, and fructose, remain experimental and have not been assessed in large-scale clinical trials.

Thus, intensive glycemic therapy (Chap. 418) needs to be applied along with the patient's education and empowerment, frequent self-monitoring of blood glucose, flexible insulin (and other drug) regimens (including the use of insulin analogues, both short- and longer-acting), individualized glycemic goals, and ongoing professional guidance, support, and consideration of both the conventional risk factors and those indicative of compromised glucose counterregulation. Given a history of hypoglycemia unawareness, a 2- to 3-week period of scrupulous avoidance of hypoglycemia is indicated.

HYPOGLYCEMIA WITHOUT DIABETES

There are many causes of hypoglycemia (Table 420-1). Because hypoglycemia is common in insulin- or insulin secretagogue-treated diabetes, it is often reasonable to assume that a clinically suspicious episode is the result of hypoglycemia. On the other hand, because hypoglycemia is rare in the absence of relevant drug-treated diabetes, it is reasonable to conclude that a hypoglycemic disorder is present only in patients in whom Whipple's triad can be demonstrated.

Particularly when patients are ill or medicated, the initial diagnostic focus should be on the possibility of drug involvement and then on critical illnesses, hormone deficiency, or non-islet cell tumor hypoglycemia. In the absence of any of these etiologic factors and in a seemingly well individual, the focus should shift to possible endogenous hyperinsulinism or accidental, surreptitious, or even malicious hypoglycemia.

Drugs Insulin and insulin secretagogues suppress glucose production and stimulate glucose utilization. Ethanol blocks gluconeogenesis but not glycogenolysis. Thus, alcohol-induced hypoglycemia typically occurs after a several-day ethanol binge during which the person eats little food, with consequent glycogen depletion. Ethanol is usually measurable in blood at the time of presentation, but its levels correlate poorly with plasma glucose concentrations. Because gluconeogenesis becomes the predominant route of glucose production during prolonged hypoglycemia, alcohol can contribute to the progression of hypoglycemia in patients with insulin-treated diabetes.

Many other drugs have been associated with hypoglycemia. These include commonly used drugs such as angiotensin-converting enzyme inhibitors and angiotensin receptor antagonists, β -adrenergic receptor antagonists, quinolone antibiotics, indomethacin, quinine, and sulfonamides.

Critical Illness Among hospitalized patients, serious illnesses such as renal, hepatic, or cardiac failure; sepsis; and inanition are second only to drugs as causes of hypoglycemia.

Rapid and extensive hepatic destruction (e.g., toxic hepatitis) causes fasting hypoglycemia because the liver is the major site of endogenous glucose production. The mechanism of hypoglycemia in patients with cardiac failure is unknown. Hepatic congestion and hypoxia may be involved. Although the kidneys are a source of glucose production, hypoglycemia in patients with renal failure is also caused by the reduced clearance of insulin and the reduced mobilization of gluconeogenic precursors in renal failure.

Sepsis is a relatively common cause of hypoglycemia. Increased glucose utilization is induced by cytokine production in macrophage-rich tissues such as the liver, spleen, and lung. Hypoglycemia develops if glucose production fails to keep pace. Cytokine-induced inhibition of gluconeogenesis in the setting of nutritional glycogen depletion, in combination with hepatic and renal hypoperfusion, may also contribute to hypoglycemia.

Hypoglycemia can be seen with starvation, perhaps because of loss of whole-body fat stores and subsequent depletion of gluconeogenic precursors (e.g., amino acids), necessitating increased glucose utilization.

Hormone Deficiencies Neither cortisol nor growth hormone is critical to the prevention of hypoglycemia, at least in adults. Nonetheless, hypoglycemia can occur with prolonged fasting in patients with primary adrenocortical failure (Addison's disease) or hypopituitarism. Anorexia and weight loss are typical features of chronic cortisol deficiency and likely result in glycogen depletion. Cortisol deficiency is associated with impaired gluconeogenesis and low levels of gluconeogenic precursors; these associations suggest that substrate-limited gluconeogenesis, in the setting of glycogen depletion, is the cause of hypoglycemia. Growth hormone deficiency can cause hypoglycemia in young children. In addition to extended fasting, high rates of glucose utilization (e.g., during exercise or in pregnancy) or low rates of glucose production (e.g., after alcohol ingestion) can precipitate hypoglycemia in adults with previously unrecognized hypopituitarism.

Hypoglycemia is not a feature of the epinephrine-deficient state that results from bilateral adrenalectomy when glucocorticoid replacement is adequate, nor does it occur during pharmacologic adrenergic blockade when other glucoregulatory systems are intact. Combined deficiencies of glucagon and epinephrine play a key role in the pathogenesis of iatrogenic hypoglycemia in people with insulin-deficient diabetes, as discussed earlier. Otherwise, deficiencies of these hormones are not usually considered in the differential diagnosis of a hypoglycemic disorder.

Non- β -Cell Tumors Fasting hypoglycemia, often termed *non-islet cell tumor hypoglycemia*, occurs occasionally in patients with large mesenchymal or epithelial tumors (e.g., hepatomas, adrenocortical carcinomas, carcinoids). The glucose kinetic patterns resemble those of hyperinsulinism (see next), but insulin secretion is suppressed appropriately during hypoglycemia. In most instances, hypoglycemia is due to overproduction of an incompletely processed form of insulin-like growth factor II ("big IGF-II") that does not complex normally with circulating binding proteins and thus more readily gains access to

2434 target tissues. The tumors are usually apparent clinically, plasma ratios of IGF-II to IGF-I are high, and free IGF-II levels (and levels of pro-IGF-II [1–21]) are elevated. Curative surgery is seldom possible, but reduction of tumor bulk may ameliorate hypoglycemia. Therapy with a glucocorticoid, a growth hormone, or both has also been reported to alleviate hypoglycemia. Hypoglycemia attributed to ectopic IGF-I production has been reported but is rare.

Endogenous Hyperinsulinism Hypoglycemia due to endogenous hyperinsulinism can be caused by (1) a primary β -cell disorder—typically a β -cell tumor (*insulinoma*), sometimes multiple insulinomas, or a functional β -cell disorder with β -cell hypertrophy or hyperplasia; (2) an antibody to insulin or to the insulin receptor; (3) a β -cell secretagogue such as a sulfonylurea; or perhaps (4) ectopic insulin secretion, among other very rare mechanisms. None of these causes is common.

The fundamental pathophysiologic feature of endogenous hyperinsulinism caused by a primary β -cell disorder or an insulin secretagogue is the failure of insulin secretion to fall to very low levels during hypoglycemia. This feature is assessed by measurement of plasma insulin, C-peptide (the connecting peptide that is cleaved from proinsulin to produce insulin), proinsulin, and glucose concentrations during hypoglycemia. Insulin, C-peptide, and proinsulin levels need not be high relative to normal, euglycemic values; rather, they are inappropriately high in the setting of a low plasma glucose concentration. Critical diagnostic findings are a plasma insulin concentration $\geq 3 \mu\text{U/mL}$ ($\geq 18 \text{ pmol/L}$), a plasma C-peptide concentration $\geq 0.6 \text{ ng/mL}$ ($\geq 0.2 \text{ nmol/L}$), and a plasma proinsulin concentration $\geq 5.0 \text{ pmol/L}$ when the plasma glucose concentration is $< 55 \text{ mg/dL}$ ($< 3.0 \text{ mmol/L}$) with symptoms of hypoglycemia. A low plasma β -hydroxybutyrate concentration ($\leq 2.7 \text{ mmol/L}$) and an increment in plasma glucose level of $> 25 \text{ mg/dL}$ ($> 1.4 \text{ mmol/L}$) after IV administration of glucagon (1.0 mg) indicate increased insulin (or IGF) actions.

The diagnostic strategy is (1) to measure plasma glucose, insulin, C-peptide, proinsulin, and β -hydroxybutyrate concentrations and to screen for circulating oral hypoglycemic agents during an episode of hypoglycemia and (2) to assess symptoms during the episode and seek their resolution following correction of hypoglycemia by IV injection of glucagon (i.e., to document Whipple's triad). This is straightforward if the patient is hypoglycemic when seen. Since endogenous hyperinsulinemic disorders usually, but not invariably, cause fasting hypoglycemia, a diagnostic episode may develop after a relatively short outpatient fast. Serial sampling during an inpatient diagnostic fast of up to 72 h or after a mixed meal is more problematic. An alternative is to give patients a detailed list of the required measurements and ask them to present to an emergency room, with the list, during a symptomatic episode. Obviously, a normal plasma glucose concentration during a symptomatic episode indicates that the symptoms are not the result of hypoglycemia.

An *insulinoma*—an insulin-secreting pancreatic islet β -cell tumor—is the prototypical cause of endogenous hyperinsulinism and therefore should be sought in patients with a compatible clinical syndrome. However, insulinoma is not the only cause of endogenous hyperinsulinism. Some patients with fasting endogenous hyperinsulinemic hypoglycemia have diffuse islet involvement with β -cell hypertrophy and sometimes hyperplasia. This pattern is commonly referred to as *nesidioblastosis*, although β -cells budding from ducts are not invariably found. Other patients have a similar islet pattern but with postprandial hypoglycemia, a disorder termed *noninsulinoma pancreatogenous hypoglycemia*. Postgastric bypass postprandial hypoglycemia, which most often follows Roux-en-Y gastric bypass, is also characterized by diffuse islet involvement and endogenous hyperinsulinism. Some have suggested that exaggerated GLP-1 responses to meals cause hyperinsulinemia and hypoglycemia, but the relevant pathogenesis has not been clearly established. If medical treatments with agents such as an α -glucosidase inhibitor, diazoxide, or octreotide fail, partial pancreatectomy may be required. Autoimmune hypoglycemias include those caused by an antibody to insulin that binds post-meal insulin and then gradually disassociates, with consequent late postprandial hypoglycemia. Alternatively, an insulin receptor antibody can function

as an agonist. The presence of an insulin secretagogue, such as a sulfonylurea or a glinide, results in a clinical and biochemical pattern similar to that of an insulinoma but can be distinguished by the presence of the circulating secretagogue. Finally, there are reports of very rare phenomena such as ectopic insulin secretion, a gain-of-function insulin receptor mutation, and exercise-induced hyperinsulinemia.

Insulinomas are uncommon, with an estimated yearly incidence of 1 in 250,000. Because more than 90% of insulinomas are benign, they are a treatable cause of potentially fatal hypoglycemia. The median age at presentation is 50 years in sporadic cases, but the tumor usually presents in the third decade when it is a component of multiple endocrine neoplasia type 1 (**Chap. 408**). More than 99% of insulinomas are within the substance of the pancreas, and the tumors are usually small ($< 2.0 \text{ cm}$ in diameter in 90% of cases). Therefore, they come to clinical attention because of hypoglycemia rather than mass effects. CT or MRI detects ~ 70 – 80% of insulinomas. These methods detect metastases in the roughly 10% of patients with a malignant insulinoma. Transabdominal ultrasound often identifies insulinomas, and endoscopic ultrasound has a sensitivity of $\sim 90\%$. Somatostatin receptor scintigraphy is thought to detect insulinomas in about half of patients. Selective pancreatic arterial calcium injections, with the endpoint of a sharp increase in hepatic venous insulin levels, regionalize insulinomas with high sensitivity, but this invasive procedure is seldom necessary except to confirm endogenous hyperinsulinism in the diffuse islet disorders. Intraoperative pancreatic ultrasonography almost invariably localizes insulinomas that are not readily palpable by the surgeon. Surgical resection of a solitary insulinoma is generally curative. Diazoxide, which inhibits insulin secretion, or the somatostatin analogue octreotide can be used to treat hypoglycemia in patients with unresectable tumors; everolimus, an mTOR (mammalian target of rapamycin) inhibitor, is promising.

ACCIDENTAL, SURREPTITIOUS, OR MALICIOUS HYPOGLYCEMIA

Accidental ingestion of an insulin secretagogue (e.g., as the result of a pharmacy or other medical error) or even accidental administration of insulin can occur. Factitious hypoglycemia, caused by surreptitious or even malicious administration of insulin or an insulin secretagogue, shares many clinical and laboratory features with insulinoma. It is most common among health care workers, patients with diabetes or their relatives, and people with a history of other factitious illnesses. However, it should be considered in all patients being evaluated for hypoglycemia of obscure cause. Ingestion of an insulin secretagogue causes hypoglycemia with increased C-peptide levels, whereas exogenous insulin causes hypoglycemia with low C-peptide levels reflecting suppression of insulin secretion.

Analytical error in the measurement of plasma glucose concentrations is rare. On the other hand, glucose monitors used to guide treatment of diabetes are not quantitative instruments, particularly at low glucose levels, and should not be used for the definitive diagnosis of hypoglycemia. Even with a quantitative method, low measured glucose concentrations can be artifactual—e.g., the result of continued glucose metabolism by the formed elements of the blood *ex vivo*, particularly in the presence of leukocytosis, erythrocytosis, or thrombocytosis or with delayed separation of the serum from the formed elements (pseudohypoglycemia).

INBORN ERRORS OF METABOLISM CAUSING HYPOGLYCEMIA

Nondiabetic hypoglycemia also results from inborn errors of metabolism. Such hypoglycemia most commonly occurs in infancy but can also occur in adulthood. Cases in adults can be classified into those resulting in fasting hypoglycemia, postprandial hypoglycemia, and exercise-induced hypoglycemia.

Fasting Hypoglycemia Although rare, disorders of glycogenolysis can result in fasting hypoglycemia. These disorders include glycogen storage disease (GSD) of types 0, I, III, and IV and Fanconi-Bickel syndrome (**Chap. 433e**). Patients with GSD types I and III characteristically have high blood lactate levels before and after meals, respectively. Both groups have hypertriglyceridemia, but ketones are high in GSD type III. Defects in fatty acid oxidation also result in fasting hypoglycemia. These defects can include (1) defects in the carnitine

cycle; (2) fatty-acid β -oxidation disorders; (3) electron transfer disturbances; and (4) ketogenesis disorders. Finally, defects in gluconeogenesis (fructose-1, 6-biphosphatase) have been reported to result in recurrent hypoglycemia and lactic acidosis.

Postprandial Hypoglycemia Inborn errors of metabolism resulting in postprandial hypoglycemia are also rare. These errors include (1) glucokinase, SUR1, and Kir6.2 potassium channel mutations; (2) congenital disorders of glycosylation; and (3) inherited fructose intolerance.

Exercise-Induced Hypoglycemia Exercise-induced hypoglycemia, by definition, follows exercise. It results in hyperinsulinemia caused by increased activity of monocarboxylate transporter 1 in β cells.

APPROACH TO THE PATIENT: Hypoglycemia

In addition to the recognition and documentation of hypoglycemia as well as its treatment (often on an urgent basis), diagnosis of the hypoglycemic mechanism is critical for the selection of therapy that prevents, or at least minimizes, recurrent hypoglycemia.

RECOGNITION AND DOCUMENTATION

Hypoglycemia is suspected in patients with typical symptoms; in the presence of confusion, an altered level of consciousness, or a seizure; or in a clinical setting in which hypoglycemia is known to occur. Blood should be drawn, whenever possible, before the administration of glucose to allow documentation of a low plasma glucose concentration. Convincing documentation of hypoglycemia requires the fulfillment of Whipple's triad. Thus, the ideal time to measure the plasma glucose level is during a symptomatic episode. A normal glucose level excludes hypoglycemia as the cause of the symptoms. A low glucose level confirms that hypoglycemia is the cause of the symptoms, provided the latter resolve after the glucose level is raised. When the cause of the hypoglycemic episode is obscure, additional measurements—made while the glucose level is low and before treatment—should include plasma insulin, C-peptide, proinsulin, and β -hydroxybutyrate levels; also critical are screening for circulating oral hypoglycemic agents and assessment of symptoms before and after the plasma glucose concentration is raised.

When the history suggests prior hypoglycemia and no potential mechanism is apparent, the diagnostic strategy is to evaluate the patient as just described and assess for Whipple's triad during and after an episode of hypoglycemia. On the other hand, while it cannot be ignored, a distinctly low plasma glucose concentration measured in a patient without corresponding symptoms raises the possibility of an artifact (pseudohypoglycemia).

DIAGNOSIS OF THE HYPOGLYCEMIC MECHANISM

In a patient with documented hypoglycemia, a plausible hypoglycemic mechanism can often be deduced from the history, physical examination, and available laboratory data (Table 420-1). Drugs, particularly alcohol or agents used to treat diabetes, should be the first consideration—even in the absence of known use of a relevant drug—given the possibility of surreptitious, accidental, or malicious drug administration. Other considerations include evidence of a relevant critical illness, hormone deficiencies (less commonly), and a non- β -cell tumor that can be pursued diagnostically (rarely). Absent one of these mechanisms in an otherwise seemingly well individual, the physician should consider endogenous hyperinsulinism and proceed with measurements and assessment of symptoms during spontaneous hypoglycemia or under conditions that might elicit hypoglycemia.

URGENT TREATMENT

If the patient is able and willing, oral treatment with glucose tablets or glucose-containing fluids, candy, or food is appropriate. A

reasonable initial dose is 20 g of glucose. If the patient is unable or unwilling (because of neuroglycopenia) to take carbohydrates orally, parenteral therapy is necessary. IV administration of glucose (25 g) should be followed by a glucose infusion guided by serial plasma glucose measurements. If IV therapy is not practical, SC or IM glucagon (1.0 mg in adults) can be used, particularly in patients with T1DM. Because it acts by stimulating glycogenolysis, glucagon is ineffective in glycogen-depleted individuals (e.g., those with alcohol-induced hypoglycemia). Glucagon also stimulates insulin secretion and is therefore less useful in T2DM. The somatostatin analogue octreotide can be used to suppress insulin secretion in sulfonyleurea-induced hypoglycemia. These treatments raise plasma glucose concentrations only transiently, and patients should therefore be urged to eat as soon as is practical to replete glycogen stores.

PREVENTION OF RECURRENT HYPOGLYCEMIA

Prevention of recurrent hypoglycemia requires an understanding of the hypoglycemic mechanism. Offending drugs can be discontinued or their doses reduced. Hypoglycemia caused by a sulfonyleurea can persist for hours or even days. Underlying critical illnesses can often be treated. Cortisol and growth hormone can be replaced if levels are deficient. Surgical, radiotherapeutic, or chemotherapeutic reduction of a non-islet cell tumor can alleviate hypoglycemia even if the tumor cannot be cured; glucocorticoid or growth hormone administration also may reduce hypoglycemic episodes in such patients. Surgical resection of an insulinoma is curative; medical therapy with diazoxide or octreotide can be used if resection is not possible and in patients with a nontumor β -cell disorder. Partial pancreatectomy may be necessary in the latter patients. The treatment of autoimmune hypoglycemia (e.g., with glucocorticoid or immunosuppressive drugs) is problematic, but these disorders are sometimes self-limited. Failing these treatments, frequent feedings and avoidance of fasting may be required. Administration of uncooked cornstarch at bedtime or even an overnight intragastric infusion of glucose may be necessary for some patients.

421 Disorders of Lipoprotein Metabolism

Daniel J. Rader, Helen H. Hobbs

Lipoproteins are complexes of lipids and proteins that are essential for transport of cholesterol, triglycerides, and fat-soluble vitamins. Previously, lipoprotein disorders were the purview of specialized lipidologists, but the demonstration that lipid-lowering therapy significantly reduces the clinical complications of atherosclerotic cardiovascular disease (ASCVD) has brought the diagnosis and treatment of these disorders into the domain of the internist. The number of individuals who are candidates for lipid-lowering therapy continues to increase. Therefore, the appropriate diagnosis and management of lipoprotein disorders is of critical importance in the practice of medicine. This chapter reviews normal lipoprotein physiology, the pathophysiology of disorders of lipoprotein metabolism, the effects of diet and other environmental factors that influence lipoprotein metabolism, and the practical approaches to the diagnosis and management of lipoprotein disorders.

LIPOPROTEIN METABOLISM

LIPOPROTEIN CLASSIFICATION AND COMPOSITION

Lipoproteins are large macromolecular complexes composed of lipids and proteins that transport poorly soluble lipids (primarily triglycerides, cholesterol, and fat-soluble vitamins) through body fluids

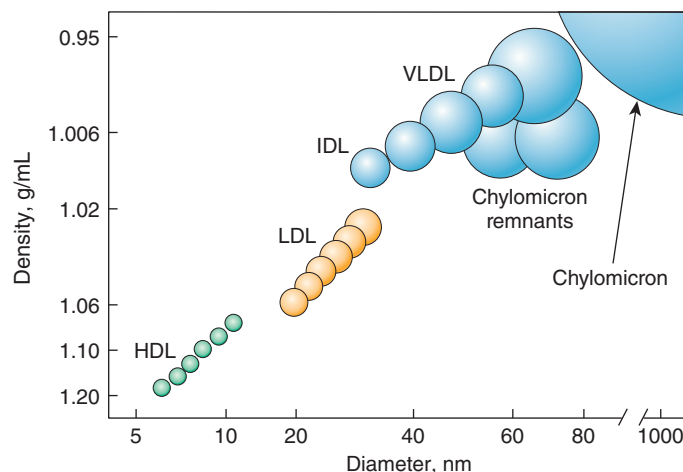


FIGURE 421-1 The density and size distribution of the major classes of lipoprotein particles. Lipoproteins are classified by density and size, which are inversely related. HDL, high-density lipoprotein; IDL, intermediate-density lipoprotein; LDL, low-density lipoprotein; VLDL, very-low-density lipoprotein.

(plasma, interstitial fluid, and lymph) to and from tissues. Lipoproteins play an essential role in the absorption of dietary cholesterol, long-chain fatty acids, and fat-soluble vitamins; the transport of triglycerides, cholesterol, and fat-soluble vitamins from the liver to peripheral tissues; and the transport of cholesterol from peripheral tissues to the liver and intestine.

Lipoproteins contain a core of hydrophobic lipids (triglycerides and cholesteryl esters) surrounded by a shell of hydrophilic lipids (phospholipids, unesterified cholesterol) and proteins (called apolipoproteins) that interact with body fluids. The plasma lipoproteins are divided into five major classes based on their relative density (Fig. 421-1 and Table 421-1): chylomicrons, very-low-density lipoproteins (VLDLs), intermediate-density lipoproteins (IDLs), low-density lipoproteins (LDLs), and high-density lipoproteins (HDLs). Each lipoprotein class comprises a family of particles that vary in density, size, and protein composition. Because lipid is less dense than water, the density of a lipoprotein particle is primarily determined by the amount of lipid per particle. Chylomicrons are the most lipid-rich and therefore least dense lipoprotein particles, whereas HDLs have the least lipid and are therefore the most dense lipoproteins. In addition to their density, lipoprotein particles can be classified according to their size, determined either by non-denaturing gel electrophoresis

TABLE 421-2 MAJOR APOLIPOPROTEINS

Apolipoprotein	Primary Source	Lipoprotein Association	Function
ApoA-I	Intestine, liver	HDL, chylomicrons	Structural protein for HDL Activates LCAT
ApoA-II	Liver	HDL, chylomicrons	Structural protein for HDL
ApoA-IV	Intestine, liver	HDL, chylomicrons	Unknown
ApoA-V	Liver	VLDL, chylomicrons	Promotes LPL-mediated triglyceride lipolysis
Apo(a)	Liver	Lp(a)	Unknown
ApoB-48	Intestine	Chylomicrons, chylomicron remnants	Structural protein for chylomicrons
ApoB-100	Liver	VLDL, IDL, LDL, Lp(a)	Structural protein for VLDL, LDL, IDL, Lp(a) Ligand for binding to LDL receptor
ApoC-I	Liver	Chylomicrons, VLDL, HDL	Unknown
ApoC-II	Liver	Chylomicrons, VLDL, HDL	Cofactor for LPL
ApoC-III	Liver, intestine	Chylomicrons, VLDL, HDL	Inhibits LPL activity and lipoprotein binding to receptors
ApoE	Liver	Chylomicron remnants, IDL, HDL	Ligand for binding to LDL receptor and other receptors

Abbreviations: HDL, high-density lipoprotein; IDL, intermediate-density lipoprotein; LCAT, lecithin-cholesterol acyltransferase; LDL, low-density lipoprotein; Lp(a), lipoprotein A; LPL, lipoprotein lipase; VLDL, very-low-density lipoprotein.

or by nuclear magnetic resonance profiling. There is a strong inverse relationship between density and size, with the largest particles being the most buoyant (chylomicrons) and the smallest particles being the most dense (HDL).

The proteins associated with lipoproteins, called *apolipoproteins* (Table 421-2), are required for the assembly, structure, function, and metabolism of lipoproteins. Apolipoproteins activate enzymes important in lipoprotein metabolism and act as ligands for cell surface receptors. ApoB is a very large protein and is the major structural protein of chylomicrons, VLDLs, IDLs, and LDLs; one molecule

TABLE 421-1 MAJOR LIPOPROTEIN CLASSES

Lipoprotein	Density, g/mL ^a	Size, nm ^b	Electrophoretic Mobility ^c	Apolipoproteins		
				Major	Other	Other Constituents
Chylomicrons	0.930	75–1200	Origin	ApoB-48	A-I, A-V, C-I, C-II, C-III, E	Retinyl esters
Chylomicron remnants	0.930–1.006	30–80	Slow pre-β	ApoB-48	A-I, A-V, C-I, C-II, C-III, E	Retinyl esters
VLDL	0.930–1.006	30–80	Pre-β	ApoB-100	A-I, A-II, A-V, C-I, C-II, C-III, E	Vitamin E
IDL	1.006–1.019	25–35	Slow pre-β	ApoB-100	C-I, C-II, C-III, E	Vitamin E
LDL	1.019–1.063	18–25	β	ApoB-100		Vitamin E
HDL	1.063–1.210	5–12	α	ApoA-I	A-II, A-IV, A-V, C-III, E	LCAT, CETP, paroxonase
Lp(a)	1.050–1.120	25	Pre-β	ApoB-100	Apo(a)	Oxidized phospholipids

^aThe density of the particle is determined by ultracentrifugation. ^bThe size of the particle is measured using gel electrophoresis. ^cThe electrophoretic mobility of the particle on agarose gel electrophoresis reflects the size and surface charge of the particle, with β being the position of LDL and α being the position of HDL.

Note: All of the lipoprotein classes contain phospholipids, esterified and unesterified cholesterol, and triglycerides to varying degrees.

Abbreviations: CETP, cholesteryl ester transfer protein; HDL, high-density lipoprotein; IDL, intermediate-density lipoprotein; LCAT, lecithin-cholesterol acyltransferase; LDL, low-density lipoprotein; Lp(a), lipoprotein A; VLDL, very-low-density lipoprotein.

of apoB, either apoB-48 (chylomicron) or apoB-100 (VLDL, IDL, or LDL), is present on each lipoprotein particle. The human liver synthesizes apoB-100, and the intestine makes apoB-48, which is derived from the same gene by mRNA editing. HDLs have different apolipoproteins that define this lipoprotein class, most importantly apoA-I, which is synthesized in the liver and intestine and is found on virtually all HDL particles. ApoA-II is the second most abundant HDL apolipoprotein and is on approximately two-thirds of the HDL particles. ApoC-I, apoC-II, and apoC-III participate in the metabolism of triglyceride-rich lipoproteins. ApoE also plays a critical role in the metabolism and clearance of triglyceride-rich particles. Most apolipoproteins, other than apoB, exchange actively among lipoprotein particles in the blood. Apolipoprotein(a) [apo(a)] is a distinctive apolipoprotein and is discussed more below.

TRANSPORT OF INTESTINALLY DERIVED DIETARY LIPIDS BY CHYLOMICRONS

One critical role of lipoproteins is the efficient transport of dietary lipids from the intestine to tissues that require fatty acids for energy or store and metabolize lipids (Fig. 421-2). Dietary triglycerides are hydrolyzed by lipases within the intestinal lumen and emulsified with bile acids to form micelles. Dietary cholesterol, fatty acids, and fat-soluble vitamins are absorbed in the proximal small intestine. Cholesterol and retinol are esterified (by the addition of a fatty acid) in the enterocyte to form cholesteryl esters and retinyl esters, respectively. Longer-chain fatty acids (>12 carbons) are incorporated into triglycerides and packaged with apoB-48, cholesteryl esters, retinyl esters, phospholipids, and cholesterol to form chylomicrons. Nascent chylomicrons are secreted into the intestinal lymph and delivered via the thoracic duct directly to the systemic circulation, where they are extensively processed by peripheral tissues before reaching the liver. The particles encounter lipoprotein lipase (LPL), which is anchored

to a glycosylphosphatidylinositol-anchored protein, GPIHBP1, that is attached to the endothelial surfaces of capillaries in adipose tissue, heart, and skeletal muscle (Fig. 421-2). The triglycerides of chylomicrons are hydrolyzed by LPL, and free fatty acids are released. ApoC-II, which is transferred to circulating chylomicrons from HDL, acts as a required cofactor for LPL in this reaction. The released free fatty acids are taken up by adjacent myocytes or adipocytes and either oxidized to generate energy or reesterified and stored as triglyceride. Some of the released free fatty acids bind albumin before entering cells and are transported to other tissues, especially the liver. The chylomicron particle progressively shrinks in size as the hydrophobic core is hydrolyzed and the hydrophilic lipids (cholesterol and phospholipids) and apolipoproteins on the particle surface are transferred to HDL, creating chylomicron remnants.

Chylomicron remnants are rapidly removed from the circulation by the liver through a process that requires apoE as a ligand for receptors in the liver. Consequently, few, if any, chylomicrons or chylomicron remnants are generally present in the blood after a 12-h fast, except in patients with certain disorders of lipoprotein metabolism.

TRANSPORT OF HEPATICALLY DERIVED LIPIDS BY VLDL AND LDL

Another key role of lipoproteins is the transport of hepatic lipids from the liver to the periphery (Fig. 421-2). VLDL particles resemble chylomicrons in protein composition but contain apoB-100 rather than apoB-48 and have a higher ratio of cholesterol to triglyceride (~1 mg of cholesterol for every 5 mg of triglyceride). The triglycerides of VLDL are derived predominantly from the esterification of long-chain fatty acids in the liver. The packaging of hepatic triglycerides with the other major components of the nascent VLDL particle (apoB-100, cholesteryl esters, phospholipids, and vitamin E) requires the action of the enzyme microsomal triglyceride transfer protein (MTP). After secretion into the plasma, VLDL acquires multiple copies of apoE and apolipoproteins of the C series by transfer from HDL. As with chylomicrons, the triglycerides of VLDL are hydrolyzed by LPL, especially in muscle, heart, and adipose tissue. After the VLDL remnants dissociate from LPL, they are referred to as IDLs, which contain roughly similar amounts of cholesterol and triglyceride. The liver removes approximately 40–60% of IDL by LDL receptor-mediated endocytosis via binding to apoE. The remainder of IDL is remodeled by hepatic lipase (HL) to form LDL. During this process, phospholipids and triglyceride in the particle are hydrolyzed, and all apolipoproteins except apoB-100 are transferred to other lipoproteins. Approximately 70% of LDL is removed from the circulation by the liver in a similar manner as IDL; however, in this case, apoB, rather than apoE, binds the LDL receptor.

Lp(a) is a lipoprotein similar to LDL in lipid and protein composition, but it contains an additional protein called apolipoprotein(a) [apo(a)]. Apo(a) is synthesized in the liver and attached to apoB-100 by a disulfide linkage. The major site of clearance of Lp(a) is the liver, but the uptake pathway is not known.

HDL METABOLISM AND REVERSE CHOLESTEROL TRANSPORT

All nucleated cells synthesize cholesterol, but only hepatocytes and enterocytes can effectively excrete cholesterol from the body, into either the bile or the gut lumen. In the liver, cholesterol is secreted into the bile, either directly or after conversion to bile acids. Cholesterol in peripheral cells is transported from the plasma membranes of peripheral cells to the liver and intestine by a process termed “reverse cholesterol transport” that is facilitated by HDL (Fig. 421-3).

Nascent HDL particles are synthesized by the intestine and the liver. Newly secreted apoA-I rapidly acquires phospholipids and unesterified cholesterol from its site

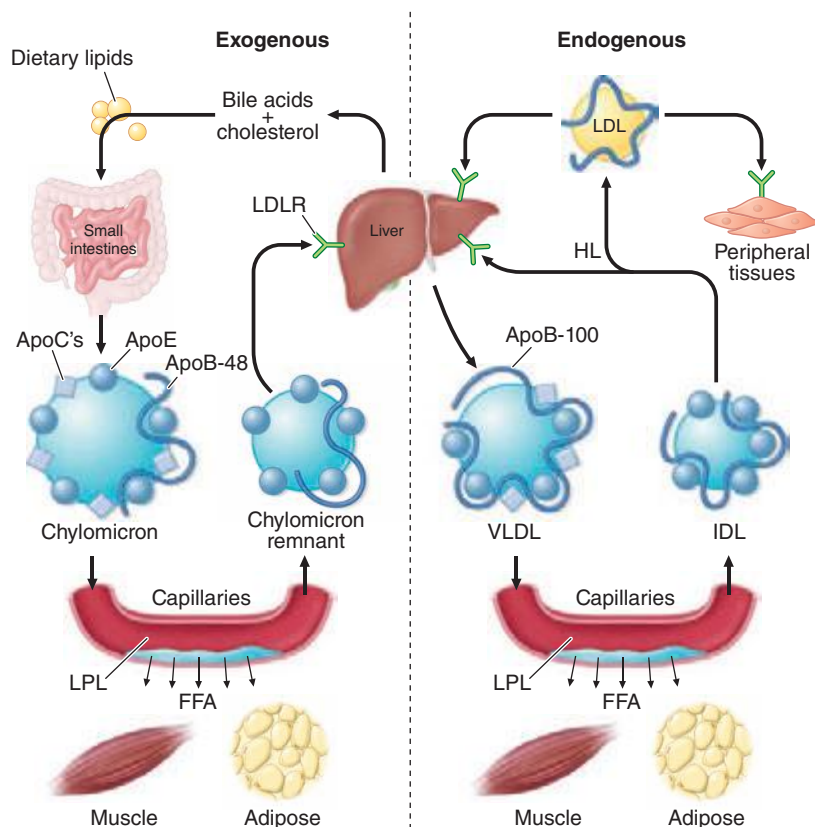


FIGURE 421-2 The exogenous and endogenous lipoprotein metabolic pathways. The exogenous pathway transports dietary lipids to the periphery and the liver. The endogenous pathway transports hepatic lipids to the periphery. FFA, free fatty acid; HL, hepatic lipase; IDL, intermediate-density lipoprotein; LDL, low-density lipoprotein; LDLR, low-density lipoprotein receptor; LPL, lipoprotein lipase; VLDL, very-low-density lipoprotein.

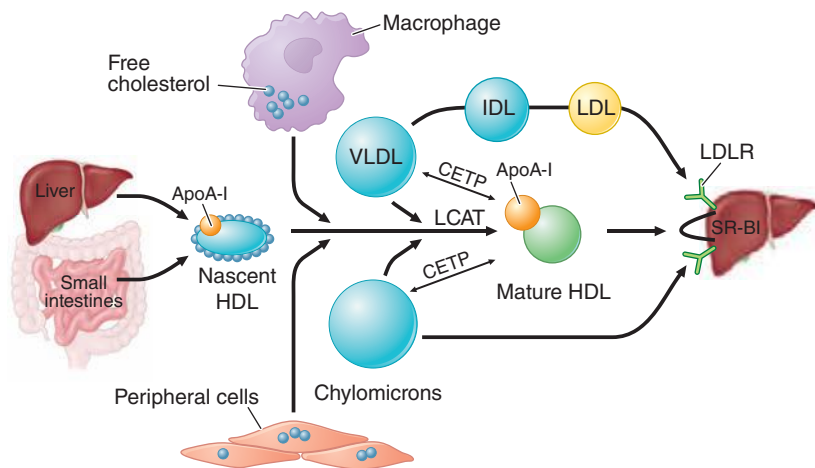


FIGURE 421-3 High-density lipoprotein (HDL) metabolism and reverse cholesterol transport. This pathway transports excess cholesterol from the periphery back to the liver for excretion in the bile. The liver and the intestine produce nascent HDLs. Free cholesterol is acquired from macrophages and other peripheral cells and esterified by lecithin-cholesterol acyltransferase (LCAT), forming mature HDLs. HDL cholesterol can be selectively taken up by the liver via SR-B1 (scavenger receptor class B1). Alternatively, HDL cholesterol ester can be transferred by cholesteryl ester transfer protein (CETP) from HDLs to very-low-density lipoproteins (VLDLs) and chylomicrons, which can then be taken up by the liver. IDL, intermediate-density lipoprotein; LDL, low-density lipoprotein; LDLR, low-density lipoprotein receptor.

of synthesis (intestine or liver) via efflux promoted by the membrane protein ATP-binding cassette protein A1 (ABCA1). This process results in the formation of discoidal HDL particles, which then recruit additional unesterified cholesterol from cells or circulating lipoproteins. Within the HDL particle, the cholesterol is esterified by lecithin-cholesterol acyltransferase (LCAT), a plasma enzyme associated with HDL, and the more hydrophobic cholesteryl ester moves to the core of the HDL particle. As HDL acquires more cholesteryl ester, it becomes spherical, and additional apolipoproteins and lipids are transferred to the particles from the surfaces of chylomicrons and VLDLs during lipolysis.

HDL cholesterol is transported to hepatocytes by both an indirect and a direct pathway. HDL cholesteryl esters can be transferred to apoB-containing lipoproteins in exchange for triglyceride by the cholesteryl ester transfer protein (CETP). The cholesteryl esters are then removed from the circulation by LDL receptor-mediated endocytosis. HDL cholesterol can also be taken up directly by hepatocytes via the scavenger receptor class B1 (SR-B1), a cell surface receptor that mediates the selective transfer of lipids to cells.

HDL particles undergo extensive remodeling within the plasma compartment by a variety of lipid transfer proteins and lipases. The phospholipid transfer protein (PLTP) transfers phospholipids from other lipoproteins to HDL or among different classes of HDL particles. After CETP- and PLTP-mediated lipid exchange, the triglyceride-enriched HDL becomes a much better substrate for HL, which hydrolyzes the triglycerides and phospholipids to generate smaller HDL particles. A related enzyme called *endothelial lipase* hydrolyzes HDL phospholipids, generating smaller HDL particles that are catabolized faster. Remodeling of HDL influences the metabolism, function, and plasma concentrations of HDL.

DISORDERS OF ELEVATED CHOLESTEROL AND TRIGLYCERIDES

Disorders of lipoprotein metabolism are collectively referred to as “dyslipidemias.” Dyslipidemias are generally characterized clinically by increased plasma levels of cholesterol, triglycerides, or both, variably accompanied by reduced levels of HDL cholesterol. Because plasma lipids are commonly screened (see below), dyslipidemia is frequently seen in clinical practice. The majority of patients with dyslipidemia have some combination of genetic predisposition (often polygenic) and environmental contribution (lifestyle, medical condition,

or drug). Many, but not all, patients with dyslipidemia are at increased risk for ASCVD, the primary reason for making the diagnosis, as intervention may reduce this risk. In addition, patients with substantially elevated levels of triglycerides may be at risk for acute pancreatitis and require intervention to reduce this risk.

Although literally hundreds of proteins influence lipoprotein metabolism and may interact to produce dyslipidemia in an individual patient, there are a limited number of discrete “nodes” that regulate lipoprotein metabolism. These include: (1) assembly and secretion of triglyceride-rich VLDLs by the liver; (2) lipolysis of triglyceride-rich lipoproteins by LPL; (3) receptor-mediated uptake of apoB-containing lipoproteins by the liver; (4) cellular cholesterol metabolism in the hepatocyte and the enterocyte; and (5) neutral lipid transfer and phospholipid hydrolysis in the plasma. The following discussion will focus on these regulatory nodes, recognizing that in many cases these nodes interact with and influence each other.

DYSLIPIDEMIA CAUSED BY EXCESSIVE HEPATIC SECRETION OF VLDL

Excessive production of VLDL by the liver is one of the most common causes of dyslipidemia. Individuals with excessive hepatic VLDL production usually have elevated fasting triglycerides and low levels of HDL cholesterol (HDL-C), with variable elevations in LDL cholesterol (LDL-C) but usually elevated plasma levels

of apoB. A cluster of other metabolic risk factors are often found in association with VLDL overproduction, including obesity, glucose intolerance, insulin resistance, and hypertension (the so-called metabolic syndrome, [Chap. 422](#)). Some of the major factors that drive hepatic VLDL secretion include obesity, insulin resistance, a high-carbohydrate diet, alcohol use, exogenous estrogens, and genetic predisposition.

Secondary Causes of VLDL Overproduction • HIGH-CARBOHYDRATE DIET Dietary carbohydrates are converted to fatty acids in the liver. Some of the newly synthesized fatty acids are esterified forming triglycerides (TGs) and secreted as constituents of VLDL. Thus, excessive intake of calories as carbohydrates, which is frequent in Western societies, leads to increased hepatic VLDL-TG secretion.

ALCOHOL Regular alcohol consumption inhibits hepatic oxidation of free fatty acids, thus promoting hepatic TG synthesis and VLDL secretion. Regular alcohol use also raises plasma levels of HDL-C and should be considered in patients with the unusual combination of elevated TGs and elevated HDL-C.

OBESITY AND INSULIN RESISTANCE (See also [Chaps. 416 and 417](#)) Obesity and insulin resistance are frequently accompanied by dyslipidemia characterized by elevated plasma levels of TG, low HDL-C, variable levels of LDL-C, and increased levels of small dense LDL. The increase in adipocyte mass and accompanying decreased insulin sensitivity associated with obesity have multiple effects on lipid metabolism, with one of the major effects being excessive hepatic VLDL production. More free fatty acids are delivered from the expanded and insulin-resistant adipose tissue to the liver, where they are reesterified in hepatocytes to form TGs, which are packaged into VLDLs for secretion into the circulation. In addition, the increased insulin levels promote increased fatty acid synthesis in the liver. In insulin-resistant patients who progress to type 2 diabetes mellitus, dyslipidemia remains common, even when the patient is under relatively good glycemic control. In addition to increased VLDL production, insulin resistance can also result in decreased LPL activity, resulting in reduced catabolism of chylomicrons and VLDLs and more severe hypertriglyceridemia (see below).

NEPHROTIC SYNDROME (See also [Chap. 335](#)) Nephrotic syndrome is a classic cause of excessive VLDL production. The molecular

mechanism of VLDL overproduction remains poorly understood but has been attributed to the effects of hypoalbuminemia leading to increased hepatic protein synthesis. Effective treatment of the underlying renal disease often normalizes the lipid profile, but most patients with chronic nephrotic syndrome require lipid-lowering drug therapy.

CUSHING'S SYNDROME (See also Chap. 406) Endogenous or exogenous glucocorticoid excess is associated with increased VLDL synthesis and secretion and hypertriglyceridemia. Patients with Cushing's syndrome frequently have dyslipidemia especially characterized by hypertriglyceridemia and low HDL-C, although elevations in plasma levels of LDL-C can also be seen.

Primary (Genetic) Causes of VLDL Overproduction Genetic variation influences hepatic VLDL production. A number of genes have been identified in which common and low-frequency variants likely contribute to increased VLDL production, likely involving interactions with diet and other environmental factors. The best recognized inherited condition associated with VLDL overproduction is familial combined hyperlipidemia.

FAMILIAL COMBINED HYPERLIPIDEMIA (FCHL) FCHL is generally characterized by elevations in plasma levels of TGs (VLDL) and LDL-C (including small dense LDL) and reduced plasma levels of HDL-C. It is estimated to occur in approximately 1 in 100–200 individuals and is an important cause of premature coronary heart disease (CHD); approximately 20% of patients who develop CHD under age 60 have FCHL. FCHL can manifest in childhood but is usually not fully expressed until adulthood. The disease clusters in families, with affected family members typically have one of three possible phenotypes: (1) elevated plasma levels of LDL-C, (2) elevated plasma levels of TGs due to elevation in VLDL, or (3) elevated plasma levels of both LDL-C and TG. The lipoprotein profile can switch among these three phenotypes in the same individual over time and may depend on factors such as diet, exercise, weight, and insulin sensitivity. Patients with FCHL almost always have significantly elevated plasma levels of apoB. The levels of apoB are disproportionately high relative to the plasma LDL-C concentration, indicating the presence of small, dense LDL particles, which are characteristic of this syndrome.

Individuals with this phenotype generally share the same metabolic defect, namely overproduction of VLDL by the liver. The molecular etiology of this condition remains poorly understood, and no single gene has been identified in which mutations cause this disorder. It is likely that defects in a combination of genes can cause the condition, suggesting that a more appropriate term for the disorder might be *polygenic combined hyperlipidemia*.

The presence of a mixed dyslipidemia (plasma TG levels between 200 and 600 mg/dL and total cholesterol levels between 200 and 400 mg/dL, usually with HDL-C levels <40 mg/dL in men and <50 mg/dL in women) and a family history of mixed dyslipidemia and/or premature CHD strongly suggests the diagnosis. Individuals with this phenotype should be treated aggressively due to significantly increased risk of premature CHD. Decreased dietary intake of simple carbohydrates, aerobic exercise, and weight loss can all have beneficial effects on the lipid profile. Patients with diabetes should be aggressively treated to maintain good glucose control. Most patients with FCHL require lipid-lowering drug therapy, starting with statins, to reduce lipoprotein levels and lower the risk of cardiovascular disease.

LIPODYSTROPHY Lipodystrophy is a condition in which the generation of adipose tissue generally or in certain fat depots is impaired. Lipodystrophies are often associated with insulin resistance and elevated plasma levels of VLDL and chylomicrons due to increased fatty acid synthesis and VLDL production, as well as reduced clearance of TG-rich particles. This disorder can be especially difficult to control. Patients with congenital generalized lipodystrophy are very rare and have nearly complete absence of subcutaneous fat, accompanied by profound insulin resistance and leptin deficiency, and accumulation of TGs in multiple tissues including the liver. Some patients with generalized lipodystrophy have been treated successfully with leptin

administration. Partial lipodystrophy is somewhat more common and can be caused by mutations in several different genes, most notably lamin A. Partial lipodystrophy is usually characterized by increased truncal fat accompanied by markedly reduced or absent subcutaneous fat in the extremities and buttocks. These patients generally have insulin resistance, often quite severe, accompanied by type 2 diabetes, hepatosteatosis, and dyslipidemia. The dyslipidemia is usually characterized by elevated TGs and cholesterol and can be difficult to manage clinically. Patients with partial lipodystrophy are at substantially increased risk of atherosclerotic vascular disease and should therefore be treated aggressively for their dyslipidemia with statins and, if necessary, additional lipid-lowering therapies.

DYSLIPIDEMIA CAUSED BY IMPAIRED LIPOLYSIS OF TRIGLYCERIDE-RICH LIPOPROTEINS

Impaired lipolysis of the TGs in TG-rich lipoproteins (TRLs) also commonly contributes to dyslipidemia. As noted above, LPL is the key enzyme responsible for hydrolyzing the TGs in chylomicrons and VLDL. LPL is synthesized and secreted into the extracellular space from adipocytes, myocytes, and cardiomyocytes. It is then transported from the subendothelial to the vascular endothelial surfaces by GPIHBP1. LPL is also synthesized in macrophages. Individuals with impaired LPL activity, whether secondary or due to a primary genetic disorder, have elevated fasting TGs and low levels of HDL-C, usually without elevation in LDL-C or apoB. Insulin resistance, in addition to causing excessive VLDL production, can also cause impaired LPL activity and lipolysis. A number of common and low-frequency genetic variants have been described that influence LPL activity, and single-gene Mendelian disorders that reduce LPL activity have also been described (Table 421-3).

Secondary Causes of Impaired Lipolysis of TRLs • OBESITY AND INSULIN RESISTANCE (See also Chaps. 415e, 416, and 417) In addition to hepatic overproduction of VLDL, as discussed above, obesity, insulin resistance, and type 2 diabetes have been reported to be associated with variably reduced LPL activity. This may be due in part to the effects of tissue insulin resistance leading to reduced transcription of LPL in skeletal muscle and adipose, as well as to increased production of the LPL inhibitor apoC-III by the liver. This reduction in LPL activity often contributes to the dyslipidemia seen in these patients.

Primary (Genetic) Causes and Genetic Predisposition to Impaired Lipolysis of TRLs • FAMILIAL CHYLOMICRONEMIA SYNDROME As noted above, LPL is required for the hydrolysis of TGs in chylomicrons and VLDLs, and apoC-II is a cofactor for LPL. Genetic deficiency or inactivity of either protein results in impaired lipolysis and profound elevations in plasma chylomicrons. These patients can also have elevated plasma levels of VLDL, but chylomicronemia predominates. The fasting plasma is turbid, and if left at 4°C (39.2°F) for a few hours, the chylomicrons float to the top and form a creamy supernatant. In these disorders, collectively called the *familial chylomicronemia syndrome*, fasting TG levels are almost invariably >1000 mg/dL. Fasting cholesterol levels are also elevated but to a lesser degree.

LPL deficiency has autosomal recessive inheritance and has a frequency of approximately 1 in 1 million in the population. *ApoC-II deficiency* is also recessive in inheritance pattern and is even less common than LPL deficiency. Multiple different mutations in the LPL and *APOC2* genes cause these diseases. Obligate LPL heterozygotes often have mild-to-moderate elevations in plasma TG levels, whereas individuals heterozygous for mutation in apoC-II do not have hypertriglyceridemia.

Both LPL and apoC-II deficiency usually present in childhood with recurrent episodes of severe abdominal pain due to acute pancreatitis. On fundoscopic examination, the retinal blood vessels are opalescent (lipemia retinalis). Eruptive xanthomas, which are small, yellowish-white papules, often appear in clusters on the back, buttocks, and extensor surfaces of the arms and legs. These typically painless skin lesions may become pruritic. Hepatosplenomegaly results from the uptake of circulating chylomicrons by reticuloendothelial cells in the liver and spleen. For unknown reasons, some patients with persistent

Genetic Disorder	Protein (Gene) Defect	Lipoproteins Elevated	Clinical Findings	Genetic Transmission	Estimated Incidence
Hypertriglyceridemia					
Lipoprotein lipase deficiency	LPL (<i>LPL</i>)	Chylomicrons, VLDL	Eruptive xanthomas, hepatosplenomegaly, pancreatitis	AR	~1/1,000,000
Familial apoC-II deficiency	ApoC-II (<i>APOC2</i>)	Chylomicrons, VLDL	Eruptive xanthomas, hepatosplenomegaly, pancreatitis	AR	<1/1,000,000
ApoA-V deficiency	ApoA-V (<i>APOA5</i>)	Chylomicrons, VLDL	Eruptive xanthomas, hepatosplenomegaly, pancreatitis	AR	<1/1,000,000
GPIIIBP1 deficiency	<i>GPIIIBP1</i>	Chylomicrons	Eruptive xanthomas, pancreatitis	AR	<1/1,000,000
Combined Hyperlipidemia					
Familial hepatic lipase deficiency	Hepatic lipase (<i>LIPC</i>)	VLDL remnants, HDL	Pancreatitis, CHD	AR	<1/1,000,000
Familial dysbetalipoproteinemia	ApoE (<i>APOE</i>)	Chylomicron remnants, VLDL remnants	Palmar and tuberoeruptive xanthomas, CHD, PVD	AR	~1/10,000
Hypercholesterolemia					
Familial hypercholesterolemia	LDL receptor (<i>LDLR</i>)	LDL	Tendon xanthomas, CHD	AD	~1/250 to 1/500
Familial defective apoB-100	ApoB-100 (<i>APOB</i>)	LDL	Tendon xanthomas, CHD	AD	<~1/1500
Autosomal dominant hypercholesterolemia, type 3	<i>PCSK9</i> (<i>PCSK9</i>)	LDL	Tendon xanthomas, CHD	AD	<1/1,000,000
Autosomal recessive hypercholesterolemia	ARH (<i>LDLRAP</i>)	LDL	Tendon xanthomas, CHD	AR	<1/1,000,000
Sitosterolemia	<i>ABCG5</i> or <i>ABCG8</i>	LDL	Tendon xanthomas, CHD	AR	<1/1,000,000

Abbreviations: AD, autosomal dominant; apo, apolipoprotein; AR, autosomal recessive; ARH, autosomal recessive hypercholesterolemia; CHD, coronary heart disease; LDL, low-density lipoprotein; LPL, lipoprotein lipase; PVD, peripheral vascular disease; VLDL, very-low density lipoprotein.

and pronounced chylomicronemia never develop pancreatitis, eruptive xanthomas, or hepatosplenomegaly. Premature CHD is not generally a feature of familial chylomicronemia syndromes.

The diagnoses of LPL and apoC-II deficiency are established enzymatically in specialized laboratories by assaying TG lipolytic activity in postheparin plasma. Blood is sampled after an IV heparin injection to release the endothelial-bound LPL. LPL activity is profoundly reduced in both LPL and apoC-II deficiency; in patients with apoC-II deficiency, it normalizes after the addition of normal plasma (providing a source of apoC-II). Molecular sequencing of the genes can be used to confirm the diagnosis.

The major therapeutic intervention in familial chylomicronemia syndrome is dietary fat restriction (to as little as 15 g/d) with fat-soluble vitamin supplementation. Consultation with a registered dietician familiar with this disorder is essential. Caloric supplementation with medium-chain TGs, which are absorbed directly into the portal circulation, can be useful, but there is uncertainty about their hepatic safety with prolonged use. If dietary fat restriction alone is not successful in resolving the chylomicronemia, fish oils have been effective in some patients. In patients with apoC-II deficiency, apoC-II can be provided by infusing fresh-frozen plasma to resolve the chylomicronemia in the acute setting. Management of patients with familial chylomicronemia syndrome is particularly challenging during pregnancy when VLDL production is increased. A gene therapy approach, called alipogene tiparvovec, is approved for LPL deficiency in Europe; it involves multiple intramuscular injections of an adeno-associated viral vector encoding a gain-of-function LPL variant, leading to skeletal myocyte expression of LPL.

APOA-V DEFICIENCY Another apolipoprotein, ApoA-V, facilitates the association of VLDL and chylomicrons with LPL and promotes their hydrolysis. Individuals harboring loss-of-function mutations in both *APOA5* alleles develop hyperchylomicronemia. Heterozygosity for variants in *APOA5* that reduce its function contributes to the polygenic basis of hypertriglyceridemia.

GPIIIBP1 DEFICIENCY Homozygosity for mutations that interfere with GPIIIBP1 synthesis or folding cause severe hypertriglyceridemia by compromising the transport of LPL to the vascular endothelium. The frequency of chylomicronemia due to mutations in *GPIIIBP1* has not been established but appears to be very rare.

FAMILIAL HYPERTRIGLYCERIDEMIA (FHTG) FHTG is characterized by elevated fasting TGs without a clear secondary cause, average to below average LDL-C levels, low HDL-C levels, and a family history of hypertriglyceridemia. Plasma LDL-C levels are often reduced due to defective conversion of TG-rich particles to LDL. In contrast to FCHL, apoB levels are not elevated. The identification of other first-degree relatives with hypertriglyceridemia is useful in making the diagnosis. Unlike in FCHL, this condition is not generally associated with a significantly increased risk of CHD. However, if the hypertriglyceridemia is exacerbated by environmental factors, medical conditions, or drugs, the TGs can rise to a level at which acute pancreatitis is a risk. Indeed, management of patients with this condition is mostly geared toward reduction of TGs to prevent pancreatitis.

Individuals with this phenotype generally have reduced lipolysis of TRLs, although overproduction of VLDL by the liver can also contribute. No single gene has been identified in which mutations cause this disorder, whereas combinations of gene variants have been shown to cause this phenotype. A more appropriate term for this condition might be *polygenic hypertriglyceridemia*.

It is important to consider and rule out secondary causes of the hypertriglyceridemia as discussed above. Increased intake of simple carbohydrates, obesity, insulin resistance, alcohol use, estrogen treatment, and certain medications can exacerbate this phenotype. Patients who are at high risk for CHD due to other risk factors should be treated with statin therapy. In patients who are otherwise not at high risk for CHD, lipid-lowering drug therapy can frequently be avoided with appropriate dietary and lifestyle changes. Patients with plasma TG levels >500 mg/dL after a trial of diet and exercise should be considered for drug therapy with a fibrate or fish oil to reduce TGs in order to prevent pancreatitis.

DYSLIPIDEMIA CAUSED BY IMPAIRED HEPATIC UPTAKE OF APOB-CONTAINING LIPOPROTEINS

Impaired uptake of LDL and remnant lipoproteins by the liver is another common cause of dyslipidemia. As discussed above, the LDL receptor is the major receptor responsible for uptake of LDL and remnant particles by the liver. Downregulation of LDL receptor activity or genetic variation that reduces the activity of the LDL receptor pathway leads to elevations in LDL-C. One major factor that reduces LDL receptor activity is a diet high in saturated and *trans* fats. Other medical conditions that reduce LDL receptor activity include hypothyroidism and estrogen

deficiency. In addition, genetic variation in a number of genes influences LDL clearance, and mutations in some of these genes cause several discrete Mendelian disorders of elevated LDL-C (Table 421-3).

Secondary Causes of Impaired Hepatic Uptake of Lipoproteins • HYPOTHYROIDISM (See also Chap. 405) Hypothyroidism is associated with elevated plasma LDL-C levels due primarily to a reduction in hepatic LDL receptor function and delayed clearance of LDL. Thyroid hormone increases hepatic expression of the LDL receptor. Hypothyroid patients also frequently have increased levels of circulating IDL, and some patients with hypothyroidism also have mild hypertriglyceridemia. Because hypothyroidism is often subtle and therefore easily overlooked, all patients presenting with elevated plasma levels of LDL-C, especially if there has been an unexplained increase in LDL-C, should be screened for hypothyroidism. Thyroid replacement therapy usually ameliorates the hypercholesterolemia; if not, the patient probably has a primary lipoprotein disorder and may require lipid-lowering drug therapy with a statin.

CHRONIC KIDNEY DISEASE (See also Chap. 335) Chronic kidney disease (CKD) is often associated with mild hypertriglyceridemia (<300 mg/dL) due to the accumulation of VLDLs and remnant lipoproteins in the circulation. TG lipolysis and remnant clearance are both reduced in patients with renal failure. Because the risk of ASCVD is increased in end-stage renal disease, subjects with hyperlipidemia, they should usually be aggressively treated with lipid-lowering agents, even though there is inadequate data at present to indicate that this population benefits from LDL-lowering therapy.

Patients with solid organ transplants often have increased lipid levels due to the effect of the drugs required for immunosuppression. These patients can present a difficult clinical management problem, since statins should be used cautiously in these patients due to untoward muscle-related side effects.

Primary (Genetic) Causes of Impaired Hepatic Uptake of Lipoproteins Genetic variation contributes substantially to elevated LDL-C levels in the general population. It has been estimated that at least 50% of variation in LDL-C is genetically determined. Many patients with elevated LDL-C have *polygenic hypercholesterolemia* characterized by hypercholesterolemia in the absence of secondary causes of hypercholesterolemia (other than dietary factors) or a primary Mendelian disorder. In patients who are genetically predisposed to higher LDL-C levels, diet plays a key role; indeed increased saturated and *trans* fats in the diet shifts the entire distribution of LDL levels in the population to the right. Inheritance of several variants that together elevate LDL-C, coupled with diet, is generally the cause of this condition; <10% of first-degree relatives themselves have hypercholesterolemia. However, single-gene (Mendelian) causes of elevated LDL-C are relatively common and should be considered in the differential diagnosis of elevated LDL-C.

FAMILIAL HYPERCHOLESTEROLEMIA (FH) FH, also known as autosomal dominant hypercholesterolemia (ADH) type 1, is an autosomal codominant disorder characterized by elevated plasma levels of LDL-C in the absence of hypertriglyceridemia. FH is caused by loss-of-function mutations in the gene encoding the LDL receptor. The reduction in LDL receptor activity in the liver results in a reduced rate of clearance of LDL from the circulation. The plasma level of LDL increases to a level such that the rate of LDL production equals the rate of LDL clearance by residual LDL receptor as well as non-LDL receptor mechanisms. More than 1600 different mutations have been reported in association with FH. The elevated levels of LDL-C in FH are primarily due to delayed removal of LDL from the blood; in addition, because the removal of IDL is also delayed, the production of LDL from IDL is also increased. Individuals with two mutated LDL receptor alleles (FH homozygotes, or compound heterozygotes) have much higher LDL-C levels than those with one mutant allele (FH heterozygotes).



Heterozygous FH is caused by the inheritance of one mutant LDL receptor allele. The population frequency of heterozygous FH due to LDL receptor mutations was originally

estimated to be 1 in 500 individuals, but recent data suggest it may be as high as approximately 1 in 250 individuals, making it one of the most common single-gene disorders in humans. FH has a higher prevalence in certain founder populations, such as South African Afrikaners, Christian Lebanese, and French Canadians. Heterozygous FH is characterized by elevated plasma levels of LDL-C (usually 200–400 mg/dL) and normal levels of TGs. Patients with heterozygous FH have hypercholesterolemia from birth, and disease recognition is usually based on detection of hypercholesterolemia on routine screening, the appearance of tendon xanthomas, or the development of symptomatic cardiovascular disease. Inheritance is dominant, meaning that the condition was inherited from one parent and ~50% of the patient's siblings can be expected to have hypercholesterolemia. The family history is frequently positive for premature CHD on the side of the family from which the mutation was inherited. Physical findings in many, but not all, patients with heterozygous FH include corneal arcus and tendon xanthomas particularly involving the dorsum of the hands and the Achilles tendons. Untreated heterozygous FH is associated with a markedly increased risk of cardiovascular disease. Untreated men with heterozygous FH have an ~50% chance of having a myocardial infarction before age 60 years, and women with heterozygous FH are at substantially increased risk as well. The age of onset of cardiovascular disease is highly variable and depends on the specific molecular defect, the level of LDL-C, and coexisting cardiovascular risk factors. FH heterozygotes with elevated plasma levels of Lp(a) (see below) appear to be at greater risk for cardiovascular disease.

No definitive diagnostic test for heterozygous FH is available, except in certain founder populations where selected mutations predominate. Most LDL receptor mutations are private and require sequencing of the LDL receptor gene for identification. Sequencing for clinical diagnosis is available but not standard of care and is rarely performed in the United States, because the clinical utility of identifying the specific mutation has not been demonstrated. A family history of hypercholesterolemia and/or premature coronary disease is supportive of the diagnosis. Secondary causes of significant hypercholesterolemia such as hypothyroidism, nephrotic syndrome, and obstructive liver disease should be excluded.

Heterozygous FH patients should be aggressively treated to lower plasma levels of LDL-C, starting in childhood. Initiation of a diet low in saturated and *trans* fats is recommended, but heterozygous FH patients virtually always require lipid-lowering drug therapy for effective control of their LDL-C levels. Statins are effective in heterozygous FH and are clearly the drug class of choice, and usually a more potent member of the class. However, some heterozygous FH patients cannot achieve adequate control of their LDL-C levels even with high-dose statin therapy and require additional drugs; a cholesterol absorption inhibitor and/or a bile acid sequestrant are the next-line classes of drugs. Currently, heterozygous FH patients whose LDL-C levels remain markedly elevated (>200 mg/dL with cardiovascular disease [CVD] or >300 mg/dL without CVD) on maximally tolerated drug therapy are candidates for LDL apheresis, a physical method of purging the blood of LDL in which the LDL particles are selectively removed from the circulation; LDL apheresis is usually performed every 2 weeks. A new class of drugs known as PCSK9 inhibitors is under clinical development and has the potential to effectively control LDL-C levels in the vast majority of patients with heterozygous FH who are inadequately controlled on a statin alone or who are statin intolerant.

Homozygous FH is caused by mutations in both alleles of the LDL receptor and therefore much rarer than heterozygous FH. Patients with homozygous FH have been classified into those patients with virtually no detectable LDL receptor activity (*receptor negative*) and those patients with markedly reduced but detectable LDL receptor activity (*receptor defective*). LDL-C levels in patients with homozygous FH range from about 400 to >1000 mg/dL, with receptor-defective patients at the lower end and receptor-negative patients at the higher end of the range. TGs are usually normal. Many patients with homozygous FH, particularly receptor-negative patients, present in childhood with cutaneous xanthomas on the hands, wrists, elbows, knees, heels, or

2442 buttocks. The devastating consequence of homozygous FH is accelerated ASCVD, which often presents in childhood or early adulthood. Atherosclerosis often develops first in the aortic root, where it can cause aortic valvular or supra-aortic stenosis, and typically extends into the coronary ostia, which become stenotic. Symptoms can be atypical, and sudden death is not uncommon. Untreated, receptor-negative patients with homozygous FH rarely survive beyond the second decade; patients with receptor-defective LDL receptor defects have a better prognosis but almost invariably develop clinically apparent atherosclerotic vascular disease by age 30, and often much sooner. Carotid and femoral disease develops later in life and is usually not clinically significant.

Homozygous FH should be suspected in a child or young adult with LDL >400 mg/dL without secondary cause. Cutaneous xanthomas, evidence of CVD, and hypercholesterolemia in both parents all are supportive of the diagnosis. Although the specific mutations in the LDL receptor can usually be identified by DNA sequencing, this is not generally performed, and the diagnosis is usually made on clinical grounds.

Patients with homozygous FH must be treated aggressively to delay the onset and progression of CVD. Receptor-defective patients sometimes respond to statins and other LDL-lowering drug classes such as a cholesterol absorption inhibitor or a bile acid sequestrant, which upregulate the LDL receptor activity. Two drugs that reduce the hepatic production of VLDL and thus LDL, a small-molecule inhibitor of the microsomal TG transfer protein (MTP) and an antisense oligonucleotide to apoB, are approved in the United States for the treatment of adults with homozygous FH and can be considered. PCSK9 inhibitors, which work through increasing LDL receptor availability, appear to have some benefit in receptor-defective patients and are under clinical development. LDL apheresis is used to lower plasma LDL levels in these patients and can promote regression of xanthomas as well as slow the progression of atherosclerosis. Because the liver is quantitatively the most important tissue for removing circulating LDLs via the LDL receptor, liver transplantation is effective in decreasing plasma LDL-C levels in this disorder but is infrequently used because of the associated problems with immunosuppression.

FAMILIAL DEFECTIVE APOB-100 (FDB) FDB, also known as autosomal dominant hypercholesterolemia (ADH) type 2, is a dominantly inherited disorder that clinically resembles heterozygous FH with elevated LDL-C levels and normal TGs. FDB is caused by mutations in the gene encoding apoB-100, specifically in the LDL receptor-binding domain of apoB-100. Several different mutations have been identified, but a single mutation predominates: substitution of glutamine for arginine at position 3500. The mutation results in a reduction in the affinity of LDL binding to the LDL receptor, so LDL is removed from the circulation at a reduced rate. FDB is less common than FH but is more prevalent in individuals of central European descent; the Lancaster County (United States) Amish are a founder population in which the prevalence of FDB is as high as 1 in 10 individuals. FDB is characterized by elevated plasma LDL-C levels with normal TGs; tendon xanthomas can be seen, although not as frequently as in FH, and there is an associated increase in risk of CHD. Patients with FDB cannot be clinically distinguished from patients with heterozygous FH, although patients with FDB tend to have somewhat lower plasma levels of LDL-C than FH heterozygotes, presumably due to the fact that IDL clearance is not impaired in this disorder. Homozygotes for FDB mutations have higher LDL-C levels than FDB heterozygotes but are not as severely affected as homozygous FH patients. The apoB-100 gene mutations can be detected directly through sequencing of the receptor-binding region of the apoB gene or genotyping for the most common mutation, but genetic diagnosis is not generally performed because there is no direct implication for clinical management. As with FH, patients are treated with statins first and, if necessary, with additional classes of LDL-lowering drugs.

AUTOSOMAL DOMINANT HYPERCHOLESTEROLEMIA DUE TO MUTATIONS IN PCSK9 (ADH-PCSK9 OR ADH3) ADH-PCSK9, also known as autosomal dominant

hypercholesterolemia (ADH) type 3, is a very rare autosomal dominant disorder caused by gain-of-function mutations in proprotein convertase subtilisin/kexin type 9 (PCSK9). PCSK9 is a secreted protein that binds to the LDL receptor, targeting it for degradation. Normally, after LDL binds to the LDL receptor, it is internalized along with the receptor, and in the low pH of the endosome, the LDL receptor dissociates from the LDL and recycles to the cell surface. When PCSK9 binds to the receptor, the complex is internalized and the receptor is directed to the lysosome, rather than to the cell surface. The missense mutations in PCSK9 that cause hypercholesterolemia enhance the activity of PCSK9. As a consequence, the number of hepatic LDL receptors is reduced. Patients with ADH-PCSK9 are similar clinically to patients with FH. They may be particularly responsive to PCSK9 inhibitors in clinical development. Loss-of-function mutations in PCSK9 cause low LDL-C levels (see below).

AUTOSOMAL RECESSIVE HYPERCHOLESTEROLEMIA (ARH) ARH is a very rare disorder that is mostly seen in individuals of Sardinian descent. The disease is caused by mutations in a protein, ARH (also called LDLR adaptor protein, LDLRAP), which is required for LDL receptor-mediated endocytosis in the liver. ARH binds to the cytoplasmic domain of the LDL receptor and links the receptor to the endocytic machinery. In the absence of LDLRAP, LDL binds to the extracellular domain of the LDL receptor, but the lipoprotein-receptor complex fails to be internalized. ARH, like homozygous FH, is characterized by hypercholesterolemia, tendon xanthomas, and premature coronary artery disease (CAD). The levels of plasma LDL-C tend to be intermediate between the levels present in FH homozygotes and FH heterozygotes, and CAD is not usually symptomatic until the third decade. LDL receptor function in cultured fibroblasts is normal or only modestly reduced in ARH, whereas LDL receptor function in lymphocytes and the liver is negligible. Unlike FH homozygotes, the hyperlipidemia responds to treatment with statins, but these patients usually require additional therapy to lower plasma LDL-C to acceptable levels.

SITOSTEROLEMIA Sitosterolemia is a rare autosomal recessive disease that can result in severe hypercholesterolemia, tendon xanthomas, and premature ASCVD. Sitosterolemia is caused by loss-of-function mutations in either of two members of the ATP-binding cassette (ABC) half transporter family, *ABCG5* and *ABCG8*. These genes are expressed in enterocytes and hepatocytes. The proteins heterodimerize to form a functional complex that transports plant sterols such as sitosterol and campesterol, and animal sterols, predominantly cholesterol, across the biliary membrane of hepatocytes into the bile and across the intestinal luminal surface of enterocytes into the gut lumen. In normal individuals, <5% of dietary plant sterols are absorbed by the proximal small intestine. The small amounts of plant sterols that enter the circulation are preferentially excreted into the bile. Thus, levels of plant sterols are kept very low in tissues. In sitosterolemia, the intestinal absorption of sterols is increased and biliary and fecal excretion of the sterols is reduced, resulting in increased plasma and tissue levels of both plant sterols and cholesterol. The increase in hepatic sterol levels results in transcriptional suppression of the expression of the LDL receptor, resulting in reduced uptake of LDL and substantially increased LDL-C levels. In addition to the usual clinical picture of hypercholesterolemia (i.e., tendon xanthomas and premature ASCVD), these patients also have anisocytosis and poikilocytosis of erythrocytes and megathrombocytes due to the incorporation of plant sterols into cell membranes. Episodes of hemolysis and splenomegaly are a distinctive clinical feature of this disease compared to other genetic forms of hypercholesterolemia and can be a clue to the diagnosis.

Sitosterolemia should be suspected in a patient with severe hypercholesterolemia without a family history of such or who responds dramatically to dietary therapy and/or ezetimibe but not statins. Sitosterolemia can be diagnosed by a laboratory finding of a substantial increase in the plasma level of sitosterol and/or other plant sterols. It is important to make the diagnosis, because bile acid sequestrants and cholesterol-absorption inhibitors are the most effective agents to reduce LDL-C and plasma plant sterol levels in these patients.

CHOLESTERYL ESTER STORAGE DISEASE (CESD) CESD, also known as *lysosomal acid lipase deficiency*, is an autosomal recessive disorder characterized by elevated LDL-C, usually in association with low HDL-C, together with progressive fatty liver ultimately leading to hepatic fibrosis. Plasma TG levels can also be mild to moderately increased in this disorder. The most severe form of this disorder, Wolman's disease, presents in infancy and is rapidly fatal. Both Wolman's disease and CESD are caused by loss-of-function variants in both alleles of the gene encoding lysosomal acid lipase (LAL; gene name *LIPA*). LAL is responsible for hydrolyzing neutral lipids, particularly TGs and cholesteryl esters, after delivery to the lysosome by cell-surface receptors such as the LDL receptor. It is particularly important in the liver, which clears large amounts of lipoproteins from the circulation. Genetic deficiency of LAL results in accumulation of neutral lipid in the hepatocytes, leading to hepatosplenomegaly, microvesicular steatosis, and ultimately fibrosis and end-stage liver disease. The etiology of the elevated LDL-C levels is uncertain; one study suggested that VLDL production is increased, but impaired LDL receptor-mediated clearance of LDL is also likely.

CESD should be particularly suspected in nonobese patients with elevated LDL-C, low HDL-C, and evidence of fatty liver in the absence of overt insulin resistance. The diagnosis can be made with a dried blood spot assay of LAL activity and confirmed by DNA genotyping for the most common mutation, followed if necessary by sequencing of the gene to find the second mutation. Liver biopsy is required to assess the degree of inflammation and fibrosis. It is important to make the diagnosis because it has implications for liver monitoring and potentially for therapeutic approaches under development.

FAMILIAL DYSBETALIPOPROTEINEMIA (FDBL) FDBL (also known as *type III hyperlipoproteinemia*) is usually a recessive disorder characterized by a mixed hyperlipidemia (elevated cholesterol and TGs) due to the accumulation of remnant lipoprotein particles (chylomicron remnants and VLDL remnants, or IDL). ApoE is present in multiple copies on chylomicron remnants and IDL, and mediates their removal via hepatic lipoprotein receptors (Fig. 421-2). FDBL is due to genetic variants of apoE, most commonly apoE2, that result in an apoE protein with reduced ability to bind lipoprotein receptors. The *APOE* gene is polymorphic in sequence, resulting in the expression of three common isoforms: apoE3, which is the most common; and apoE2 and apoE4, which both differ from apoE3 by a single amino acid. Although associated with slightly higher LDL-C levels and increased CHD risk, the apoE4 allele is not associated with FDBL. Individuals who carry one or two apoE4 alleles have an increased risk of Alzheimer's disease. ApoE2 has a lower affinity for the LDL receptor; therefore, chylomicron remnants and IDL containing apoE2 are removed from plasma at a slower rate. Individuals who are homozygous for the E2 allele (the E2/E2 genotype) comprise the most common subset of patients with FDBL.

Approximately 0.5% of the general population are apoE2/E2 homozygotes, but only a small minority of these individuals actually develop hyperlipidemia characteristic of FDBL. In most cases, an additional, sometimes identifiable, factor precipitates the development of hyperlipoproteinemia. The most common precipitating factors are a high-fat diet, diabetes mellitus, obesity, hypothyroidism, renal disease, HIV infection, estrogen deficiency, alcohol use, or certain drugs. The disease seldom presents in women before menopause. Other mutations in apoE can cause a dominant form of FDBL where the hyperlipidemia is fully manifest in the heterozygous state, but these mutations are very rare.

Patients with FDBL usually present in adulthood with hyperlipidemia, xanthomas, or premature coronary or peripheral vascular disease. In FDBL, in contrast to other disorders of elevated TGs, the plasma levels of cholesterol and TG are often elevated to a similar degree, and the level of HDL-C is usually normal or reduced. Two distinctive types of xanthomas, tuberoeruptive and palmar, are seen in FDBL patients. Tuberoeruptive xanthomas begin as clusters of small papules on the elbows, knees, or buttocks and can grow to the size of small grapes. Palmar xanthomas (alternatively called *xanthomata striata palmaris*) are orange-yellow discolorations of the creases in the palms and wrists.

Both of these xanthoma types are virtually pathognomonic for FDBL. Subjects with FDBL have premature ASCVD and tend to have more peripheral vascular disease than is typically seen in FH.

The definitive diagnosis of FDBL can be made either by the documentation of very high levels of remnant lipoproteins or by identification of the apoE2/E2 genotype. A variety of methods are used to identify remnant lipoproteins in the plasma, including "β-quantification" by ultracentrifugation (ratio of directly measured VLDL-C to total plasma TG >0.30), lipoprotein electrophoresis (broad β band), or nuclear magnetic resonance lipoprotein profiling. The Friedewald formula for calculation of LDL-C is not valid in FDBL because the VLDL particles are depleted in TG and enriched in cholesterol. The plasma levels of LDL-C are actually low in this disorder due to defective metabolism of VLDL to LDL. DNA-based methods (apoE genotyping) can be performed to confirm homozygosity for apoE2. However, absence of the apoE2/E2 genotype does not strictly rule out the diagnosis of FDBL, because other mutations in apoE can (rarely) cause this condition.

Because FDBL is associated with increased risk of premature ASCVD, it should be treated aggressively. Other metabolic conditions that can worsen the hyperlipidemia (see above) should be managed. Patients with FDBL are typically diet-responsive and can respond favorably to weight reduction and to low-cholesterol, low-fat diets. Alcohol intake should be curtailed. Pharmacologic therapy is often required, and statins are the first line in management. In the event of statin intolerance or insufficient control of hyperlipidemia, cholesterol absorption inhibitors, fibrates, and niacin are also effective in the treatment of FDBL.

HEPATIC LIPASE DEFICIENCY Hepatic lipase (HL; gene name *LIPC*) is a member of the same gene family as LPL and hydrolyzes TGs and phospholipids in remnant lipoproteins and HDL. Hydrolysis of lipids in remnant particles by HL contributes to their hepatic uptake via an apoE-mediated process. HL deficiency is a very rare autosomal recessive disorder characterized by elevated plasma levels of cholesterol and TGs (mixed hyperlipidemia) due to the accumulation of lipoprotein remnants, accompanied by elevated plasma level of HDL-C. The diagnosis is confirmed by measuring HL activity in postheparin plasma and/or confirmation of loss-of-function mutations in both alleles of *HL/LIPC*. Due to the small number of patients with HL deficiency, the association of this genetic defect with ASCVD is not entirely clear, although anecdotally patients with HL deficiency who have premature CVD have been described. As with FDBL, statin therapy is recommended to reduce remnant lipoproteins and cardiovascular risk.

Additional Secondary Causes of Dyslipidemia Many of the secondary causes of dyslipidemia (Table 421-4) have been described above. Additional considerations are discussed here.

LIVER DISORDERS (See also Chap. 357) Because the liver is the principal site of formation and clearance of lipoproteins, liver disorders can affect plasma lipid levels in a variety of ways. Hepatitis due to infection, drugs, or alcohol is often associated with increased VLDL synthesis and mild to moderate hypertriglyceridemia. Severe hepatitis and liver failure are associated with dramatic reductions in plasma cholesterol and TGs due to reduced lipoprotein biosynthetic capacity.

Cholestasis is associated with hypercholesterolemia, which can be very severe. A major pathway by which cholesterol is excreted from the body is via secretion into bile, either directly or after conversion to bile acids, and cholestasis blocks this critical excretory pathway. In cholestasis, free cholesterol, coupled with phospholipids, is secreted into the plasma as a constituent of a lamellar particle called *LP-X*. The particles can deposit in skinfolds, producing lesions resembling those seen in patients with FDBL (xanthomata strata palmaris). Planar and eruptive xanthomas can also be seen in patients with cholestasis.

DRUGS Many drugs have an impact on lipid metabolism and can result in significant alterations in the lipoprotein profile (Table 421-4). Estrogen administration is associated with increased VLDL and HDL synthesis, resulting in elevated plasma levels of both TGs and HDL-C. This lipoprotein pattern is distinctive because the levels of plasma TG and HDL-C are typically inversely related. Plasma TG levels should be monitored when birth control pills or postmenopausal estrogen

LDL		HDL		VLDL Elevated	IDL Elevated	Chylomicrons Elevated	Lp(a) Elevated
Elevated	Reduced	Elevated	Reduced				
Hypothyroidism	Severe liver disease	Alcohol	Smoking	Obesity	Multiple myeloma	Autoimmune disease	Chronic kidney disease Nephrotic syndrome
Nephrotic syndrome	Malabsorption	Exercise	DM type 2	DM type 2	Monoclonal gammopathy	DM type 2	Inflammation Menopause
Cholestasis	Malnutrition	Exposure to chlorinated hydrocarbons	Obesity	Glycogen storage disease			
Acute intermittent porphyria	Gaucher's disease	Drugs: estrogen	Gaucher's disease	Malnutrition	Autoimmune disease		Orchidectomy
	Chronic infectious disease			Nephrotic syndrome			
Anorexia nervosa	Hyperthyroidism		Cholesteryl ester storage disease	Alcohol	Hypothyroidism		Hypothyroidism
Hepatoma	Drugs: niacin toxicity			Renal failure			
Drugs: thiazides, cyclosporin, carbamazepine			Drugs: anabolic steroids, beta blockers	Stress			Drugs: growth hormone, isotretinoin
				Cushing's syndrome			
				Acromegaly			
				Lipodystrophy			
				Drugs: estrogen, beta blockers, glucocorticoids, bile acid binding resins, retinoic acid			

Abbreviations: DM, diabetes mellitus; HDL, high-density lipoprotein; IDL, intermediate-density lipoprotein; LDL, low-density lipoprotein; Lp(a), lipoprotein A; VLDL, very-low-density lipoprotein.

therapy is initiated to ensure that the increase in VLDL production does not lead to severe hypertriglyceridemia. Use of low-dose preparations of estrogen or the estrogen patch can minimize the effect of exogenous estrogen on lipids.

INHERITED CAUSES OF LOW LEVELS OF ApoB-CONTAINING LIPOPROTEINS

Plasma concentrations of LDL-C <60 mg/dL are unusual. Although in some cases LDL-C levels in this range may be reflective of malnutrition or serious chronic illness, LDL-C <60 mg/dL in an otherwise healthy individual suggests an inherited condition. The major inherited causes of low LDL-C are reviewed here.

Abetalipoproteinemia The synthesis and secretion of apoB-containing lipoproteins in the enterocytes of the proximal small bowel and in the hepatocytes of the liver involve a complex series of events that coordinate the coupling of various lipids with apoB-48 and apoB-100, respectively. Abetalipoproteinemia is a rare autosomal recessive disease caused by loss-of-function mutations in the gene encoding microsomal TG transfer protein (MTP; gene name *MTTP*), a protein that transfers lipids to nascent chylomicrons and VLDLs in the intestine and liver, respectively. Plasma levels of cholesterol and TG are extremely low in this disorder, and chylomicrons, VLDLs, LDLs, and apoB are undetectable in plasma. The parents of patients with abetalipoproteinemia (obligate heterozygotes) have normal plasma lipid and apoB levels. Abetalipoproteinemia usually presents in early childhood with diarrhea and failure to thrive due to fat malabsorption. The initial neurologic manifestations are loss of deep tendon reflexes, followed by decreased distal lower extremity vibratory and proprioceptive sense, dysmetria, ataxia, and the development of a spastic gait, often by the third or fourth decade. Patients with abetalipoproteinemia also develop a progressive pigmented retinopathy presenting with decreased night and color vision, followed by reductions in daytime visual acuity and ultimately progressing to near-blindness. The presence of spinocerebellar degeneration and pigmented retinopathy in this disease has resulted in some patients with abetalipoproteinemia being misdiagnosed as having Friedreich's ataxia.

Most of the clinical manifestations of abetalipoproteinemia result from defects in the absorption and transport of fat-soluble vitamins.

Vitamin E and retinyl esters are normally transported from enterocytes to the liver by chylomicrons, and vitamin E is dependent on VLDL for transport out of the liver and into the circulation. As a consequence of the inability of these patients to secrete apoB-containing particles, patients with abetalipoproteinemia are markedly deficient in vitamin E and are also mildly to moderately deficient in vitamins A and K. Patients with abetalipoproteinemia should be referred to specialized centers for confirmation of the diagnosis and appropriate therapy. Treatment consists of a low-fat, high-caloric, vitamin-enriched diet accompanied by large supplemental doses of vitamin E. It is imperative that treatment be initiated as soon as possible to prevent development of neurologic sequelae, which can progress even with appropriate therapy. New therapies for this serious disease are needed.

Familial Hypobetalipoproteinemia (FHBL) FHBL generally refers to a condition of low total cholesterol, LDL-C, and apoB due to mutations in apoB. Most of the mutations causing FHBL result in a truncated apoB protein, resulting in impaired assembly and secretion of chylomicrons from enterocytes and VLDL from the liver. Mutations that result in VLDL particles containing a truncated apoB protein are cleared from the circulation at an accelerated rate, which also contributes to patients with this disorder having low levels of LDL-C and apoB. Individuals heterozygous for these mutations usually have LDL-C levels <60–80 mg/dL and also tend to have lower levels of plasma TG. Many FHBL patients have elevated levels of hepatic fat (due to reduced VLDL export) and sometimes have increased levels of liver transaminases, although it appears that these patients infrequently develop associated inflammation and fibrosis.

Mutations in both apoB alleles cause homozygous FHBL, an extremely rare disorder resembling abetalipoproteinemia with nearly undetectable LDL-C and apoB. The neurologic defects in this form of hypobetalipoproteinemia tend to be less severe than is typically seen in abetalipoproteinemia. Homozygous hypobetalipoproteinemia can be distinguished from abetalipoproteinemia by examining the inheritance pattern of the plasma LDL-C level. The levels of LDL-C and apoB are normal in the parents of patients with abetalipoproteinemia and low in those of patients with homozygous hypobetalipoproteinemia.

PCSK9 Deficiency Another inherited cause of low LDL-C results from loss-of-function mutations in PCSK9. PCSK9 is a secreted protein that binds to the extracellular domain of the LDL receptor in the liver and promotes the degradation of the receptor. Heterozygosity for nonsense mutations in PCSK9 that interfere with the synthesis of the protein are associated with increased hepatic LDL receptor activity and reduced plasma levels of LDL-C. Such mutations are particularly frequent in individuals of African descent. Individuals who are heterozygous for a loss-of-function mutation in PCSK9 have an ~30–40% reduction in plasma levels of LDL-C and have a substantial protection from CHD relative to those without a PCSK9 mutation, presumably due to having lower plasma cholesterol levels since birth. This observation led to the development of PCSK9 inhibitors as a new strategy for reducing LDL-C levels and cardiovascular risk. Homozygotes for these nonsense mutations have been reported and have extremely low LDL-C levels (<20 mg/dL) but appear otherwise healthy. A sequence variation of somewhat higher frequency (R46L) is found predominantly in individuals of European descent. This mutation impairs, but does not completely destroy, PCSK9 function. As a consequence, the plasma levels of LDL-C in individuals carrying this mutation are more modestly reduced (~15–20%); individuals with these mutations have a 45% reduction in ASCVD risk.

DISORDERS OF REDUCED HDL CHOLESTEROL

Low levels of HDL-C are very commonly encountered in clinical practice. Low HDL-C is an important independent predictor of increased cardiovascular risk and has been used regularly in standardized risk calculators, including the most recent one from the American Heart Association (AHA)/American College of Cardiology (ACC). However, it remains very uncertain whether low HDL-C is directly causal for the development of ASCVD. HDL metabolism is strongly influenced by TRLs, insulin resistance, and inflammation, among other environmental and medical factors. Thus the HDL-C measurement integrates a number of cardiovascular risk factors, potentially explaining its strong inverse association with ASCVD.

The majority of patients with low HDL-C have some combination of genetic predisposition and secondary factors. Variants in dozens of genes have been shown to influence HDL-C levels. Even more important quantitatively, obesity and insulin resistance have strong suppressive effects on HDL-C, and low HDL-C in these conditions is widely observed. Furthermore, the vast majority of patients with elevated TGs have reduced levels of HDL-C. Most patients with low HDL-C who have been studied in detail have accelerated catabolism of HDL and its associated apoA-I as the physiologic basis for the low HDL-C. Importantly, although HDL-C remains an important biomarker for assessing cardiovascular risk, it is not currently a direct target of intervention for raising the level in order to reduce cardiovascular risk. Certain therapeutic approaches in clinical development, such as inhibitors of CETP (see below), have the potential to change this paradigm.

INHERITED CAUSES OF VERY LOW LEVELS OF HDL-C

Mutations in genes encoding proteins that play critical roles in HDL synthesis and catabolism can result in reductions in plasma levels of HDL-C. Unlike the genetic forms of hypercholesterolemia, which are invariably associated with premature coronary atherosclerosis, genetic forms of hypoalphalipoproteinemia (low HDL-C) are often not associated with clearly increased risk of ASCVD.

Gene Deletions in the APOA5-A1-C3-A4 Locus and Coding Mutations in APOA1 Complete genetic deficiency of apoA-I due to a complete deletion of the *APOA1* gene results in the virtual absence of circulating HDL and appears to increase the risk of premature ASCVD. The genes encoding *APOA5*, *APOA1*, *APOC3*, and *APOA4* are clustered together on chromosome 11. Some patients with no apoA-I have genomic deletions that include other genes in the cluster. ApoA-I is required for LCAT activity. In the absence of LCAT, free cholesterol levels increase in both plasma (not HDL) and in tissues. The free cholesterol

can form deposits in the cornea and in the skin, resulting in corneal opacities and planar xanthomas. Premature CHD is associated with apoA-I deficiency.

Missense and nonsense mutations in the apoA-I gene are present in some patients with low plasma levels of HDL-C (usually 15–30 mg/dL), but are a rare cause of low plasma HDL-C levels. Most individuals with low plasma HDL-C levels due to missense mutations in apoA-I do not appear to have premature CHD. Patients who are heterozygous for an Arg173Cys substitution in apoA-I (so-called apoA-I_{Milano}) have very low plasma levels of HDL-C due to impaired LCAT activation and accelerated clearance of the HDL particles containing the abnormal apoA-I. Despite having very low plasma levels of HDL-C, these individuals do not have an increased risk of premature CHD.

A few selected missense mutations in apoA-I and apoA-II promote the formation of amyloid fibrils, which can cause systemic amyloidosis.

Tangier Disease (ABCA1 Deficiency) Tangier disease is a rare autosomal co-dominant form of extremely low plasma HDL-C levels that is caused by mutations in the gene encoding ABCA1, a cellular transporter that facilitates efflux of unesterified cholesterol and phospholipids from cells to apoA-I (Fig. 421-3). ABCA1 in the liver and intestine rapidly lipidates the apoA-I secreted from the basolateral membranes of these tissues. In the absence of ABCA1, the nascent, poorly lipidated apoA-I is immediately cleared from the circulation. Thus, patients with Tangier disease have extremely low circulating plasma levels of HDL-C (<5 mg/dL) and apoA-I (<5 mg/dL). Cholesterol accumulates in the reticuloendothelial system of these patients, resulting in hepatosplenomegaly and pathognomonic enlarged, grayish yellow or orange tonsils. An intermittent peripheral neuropathy (mononeuritis multiplex) or a sphingomyelia-like neurologic disorder can also be seen in this disorder. Tangier disease is probably associated with some increased risk of premature atherosclerotic disease, although the association is not as robust as might be anticipated, given the very low levels of HDL-C and apoA-I in these patients. Patients with Tangier disease also have low plasma levels of LDL-C, which may attenuate the atherosclerotic risk. Obligate heterozygotes for ABCA1 mutations have moderately reduced plasma HDL-C levels (15–30 mg/dL), and their risk of premature CHD remains uncertain.

Familial LCAT Deficiency This rare autosomal recessive disorder is caused by mutations in LCAT, an enzyme synthesized in the liver and secreted into the plasma, where it circulates associated with lipoproteins (Fig. 421-3). As reviewed above, the enzyme is activated by apoA-I and mediates the esterification of cholesterol to form cholesteryl esters. Consequently, in familial LCAT deficiency, the proportion of free cholesterol in circulating lipoproteins is greatly increased (from ~25% to >70% of total plasma cholesterol). Deficiency in this enzyme interferes with the maturation of HDL particles and results in rapid catabolism of circulating apoA-I.

Two genetic forms of familial LCAT deficiency have been described in humans: complete deficiency (also called *classic LCAT deficiency*) and partial deficiency (also called *fish-eye disease*). Progressive corneal opacification due to the deposition of free cholesterol in the cornea, very low plasma levels of HDL-C (usually <10 mg/dL), and variable hypertriglyceridemia are characteristic of both disorders. In partial LCAT deficiency, there are no other known clinical sequelae. In contrast, patients with complete LCAT deficiency have hemolytic anemia and progressive renal insufficiency that eventually leads to end-stage renal disease. Remarkably, despite the extremely low plasma levels of HDL-C and apoA-I, premature ASCVD is not a consistent feature of either LCAT deficiency or fish eye disease. The diagnosis can be confirmed in a specialized laboratory by assaying plasma LCAT activity or by sequencing the *LCAT* gene.

Primary Hypoalphalipoproteinemia The condition of low plasma levels of HDL-C (the “alpha lipoprotein”) is referred to as *hypoalphalipoproteinemia*. Primary hypoalphalipoproteinemia is defined as a plasma HDL-C level below the tenth percentile in the setting of relatively normal cholesterol and TG levels, no apparent secondary causes of low plasma HDL-C, and no clinical signs of LCAT deficiency or Tangier

2446 disease. This syndrome is often referred to as *isolated low HDL*. A family history of low HDL-C facilitates the diagnosis of an inherited condition, which may follow an autosomal dominant pattern. The metabolic etiology of this disease appears to be primarily accelerated catabolism of HDL and its apolipoproteins. Some of these patients may have ABCA1 mutations and therefore technically have heterozygous Tangier disease. Several kindreds with primary hypoalphalipoproteinemia and an increased incidence of premature CHD have been described, although it is not clear if the low HDL-C level is the cause of the accelerated atherosclerosis in these families. Association of hypoalphalipoproteinemia with premature CHD may depend on the specific nature of the gene defect or the underlying metabolic defect that either directly or indirectly causes the low plasma HDL-C level.

INHERITED CAUSES OF VERY HIGH LEVELS OF HDL-C

CETP Deficiency Loss-of-function mutations in both alleles of the gene encoding CETP cause substantially elevated HDL-C levels (usually >150 mg/dL). As noted above, CETP transfers cholesteryl esters from HDL to apoB-containing lipoproteins (Fig. 421-3). Absence of this transfer activity results in an increase in the cholesteryl ester content of HDL and a reduction in plasma levels of LDL-C. The large, cholesterol-rich HDL particles circulating in these patients are cleared at a reduced rate. CETP deficiency was first diagnosed in Japanese persons and is rare outside of Japan. The relationship of CETP deficiency to ASCVD remains unresolved. Heterozygotes for CETP deficiency have only modestly elevated HDL-C levels. Based on the phenotype of high HDL-C in CETP deficiency, pharmacologic inhibition of CETP is under development as a new therapeutic approach to both raise HDL-C levels and lower LDL-C levels, but whether it will reduce risk of ASCVD remains to be determined.

SCREENING, DIAGNOSIS, AND MANAGEMENT OF DISORDERS OF LIPOPROTEIN METABOLISM

SCREENING

Plasma lipid and lipoprotein levels should be measured in all adults, preferably after a 12-h overnight fast. In most clinical laboratories, the total cholesterol and TGs in the plasma are measured enzymatically, and then the cholesterol in the supernatant is measured after precipitation of apoB-containing lipoproteins to determine the HDL-C. The LDL-C is then estimated using the following equation:

$$\text{LDL-C} = \text{total cholesterol} - (\text{TG}/5) - \text{HDL-C}$$

(The VLDL cholesterol content is estimated by dividing the plasma TG by 5, reflecting the ratio of TG to cholesterol in VLDL particles.) This formula (the Friedewald formula) is reasonably accurate if test results are obtained on fasting plasma and if the TG level does not exceed ~200 mg/dL; by convention it cannot be used if the TG level is >400 mg/dL. LDL-C can be directly measured by a number of methods. Further evaluation and treatment are based primarily on the clinical assessment of absolute cardiovascular risk using risk calculators such as the AHA/ACC risk calculator based on a large amount of observational data.

DIAGNOSIS

A critical first step in managing a lipoprotein disorder is to attempt to determine the class or classes of lipoproteins that are increased or decreased in the patient. Once the hyperlipidemia is accurately classified, efforts should be directed to rule out any possible secondary causes of the hyperlipidemia (Table 421-4). Although many patients with hyperlipidemia have a primary (i.e., genetic) cause of their lipid disorder, secondary factors frequently contribute to the hyperlipidemia. A careful social, medical, and family history should be obtained. A fasting glucose should be obtained in the initial workup of all subjects with an elevated TG level. Nephrotic syndrome and chronic renal insufficiency should be excluded by obtaining urine protein and serum creatinine. Liver function tests should be performed to rule out hepatitis and cholestasis. Hypothyroidism should be ruled out by measuring serum thyroid-stimulating hormone.

Once secondary causes have been ruled out, attempts should be made to diagnose the primary lipid disorder because the underlying genetic defect can provide important prognostic information regarding the risk of developing CHD, the response to drug therapy, and the management of other family members. Obtaining the correct diagnosis often requires a detailed family medical history, lipid analyses in family members, and sometimes specialized testing.

Severe Hypertriglyceridemia If the fasting plasma TG level is >1000 mg/dL, the patient has chylomicronemia. If the cholesterol-to-TG ratio is >10, familial chylomicronemia syndrome must be considered, and LPL activity measured in postheparin plasma can help with making that diagnosis. Most adults with chylomicronemia also have elevated VLDL levels. These individuals usually do not have a Mendelian disorder but instead are genetically predisposed and have secondary factors (diet, obesity, glucose intolerance, alcohol ingestion, estrogen therapy) that contribute to the hyperlipidemia. Such patients are a risk of acute pancreatitis and should be treated to reduce their TG levels and thus their risk of pancreatitis.

Severe Hypercholesterolemia If the levels of LDL-C are very high (greater than a ninety-fifth percentile for age and sex), it is likely that the patient has a genetic cause of hypercholesterolemia. At present, there is no compelling reason to perform molecular studies to further refine the molecular diagnosis because the clinical management is not affected. Recessive forms of severe hypercholesterolemia are rare, but if a patient with severe hypercholesterolemia has parents with normal cholesterol levels, ARH, sitosterolemia, and CESD should be considered. Patients with more moderate hypercholesterolemia that does not segregate in families as a monogenic trait are likely to have polygenic hypercholesterolemia.

Combined Hyperlipidemia The most common errors in the diagnosis of lipid disorders involve patients with combined hyperlipidemia. Elevations in the plasma levels of both cholesterol and TGs are seen in patients with increased plasma levels of VLDL and LDL or of remnant lipoproteins. A β -quantification to determine the VLDL cholesterol/TG ratio in plasma (see discussion of FDBL) or a direct measurement of the plasma LDL-C should be performed at least once prior to initiation of lipid-lowering therapy to determine if the hyperlipidemia is due to the accumulation of remnants or to an increase in both LDL and VLDL. Measurement of plasma apoB levels can help identify patients with FCHL who may require more aggressive treatment.

APPROACH TO THE PATIENT: Lipoprotein Disorders

The major goals in the clinical management of lipoprotein disorders are: (1) prevention of acute pancreatitis in patients with severe hypertriglyceridemia; and (2) prevention of CVD and related cardiovascular events.

MANAGEMENT OF SEVERE HYPERTRIGLYCERIDEMIA TO PREVENT PANCREATITIS

Although the observational relationship between severe hypertriglyceridemia, particularly chylomicronemia, and acute pancreatitis is well-established, there has never been a clinical trial designed or powered to prove that intervention to reduce TGs reduces the risk of pancreatitis. Nevertheless, it is generally considered appropriate medical practice to intervene in patients with TGs >500 mg/dL in order to reduce the risk of pancreatitis. It remains controversial whether individuals with severe hypertriglyceridemia are at increased risk for ASCVD.

Lifestyle Modifying the lifestyle of the patient with severe hypertriglyceridemia often is associated with a significant reduction in plasma TG level. Patients who drink alcohol should be encouraged to decrease or preferably eliminate their intake. Patients with severe

hypertriglyceridemia often benefit from a formal dietary consultation with a dietician intimately familiar with counseling patients on the dietary management of high TGs. Dietary fat intake should be restricted to reduce the formation of chylomicrons in the intestine. The excessive intake of simple carbohydrates should be discouraged because insulin drives TG production in the liver. Aerobic exercise and even increase in regular physical activity can have a positive effect in reducing TG levels and should be strongly encouraged. For patients who are overweight, weight loss can help to reduce TG levels. In extreme cases, bariatric surgery has been shown to not only produce effective weight loss but also substantially reduce plasma TG levels.

Pharmacologic Therapy for Severe Hypertriglyceridemia Despite the above interventions, however, many patients with severe hypertriglyceridemia require pharmacologic therapy (Table 421-5). Patients who persist in having fasting TG >500 mg/dL despite active lifestyle management are candidates for pharmacologic therapy. There are three classes of drugs that are used for management of these patients: fibrates, omega-3 fatty acids (fish oils), and niacin. In addition, statins can reduce plasma TG levels and also reduce ASCVD risk.

FIBRATES Fibrates, or fibric acid derivatives, are agonists of PPAR α , a nuclear receptor involved in the regulation of lipid metabolism. Fibrates stimulate LPL activity (enhancing TG hydrolysis), reduce apoC-III synthesis (enhancing lipoprotein remnant clearance), promote β -oxidation of fatty acids, and may reduce VLDL TG production. Fibrates are a first-line therapy for severe hypertriglyceridemia (>500 mg/dL). This class of therapeutic agents sometimes lowers but more often raises the plasma level of LDL-C in individuals with severe hypertriglyceridemia. Fibrates are generally well tolerated, but are associated with an increase in the incidence of gallstones. Fibrates can cause myopathy, especially when combined with other lipid-lowering therapy (statins, niacin), and can raise creatinine. Fibrates should be used with caution in patients with CKD. Importantly, fibrates can potentiate the effect of warfarin and certain oral hypoglycemic agents, so the anticoagulation status and plasma glucose levels should be closely monitored in patients on these agents.

OMEGA 3 FATTY ACIDS (FISH OILS) Omega-3 fatty acids, or omega-3 polyunsaturated fatty acids (n-3 PUFAs), commonly known as fish oils, are present in high concentration in fish and in flaxseed. The most widely used n-3 PUFAs for the treatment of hyperlipidemias are the two active molecules in fish oil: eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA). n-3 PUFAs have been concentrated into tablets and in doses of 3–4 g/d are effective at lowering fasting TG levels. Fish oils are a reasonable consideration for first-line therapy in patients with severe hypertriglyceridemia (>500 mg/dL) to prevent pancreatitis. Fish oils can cause an increase in plasma LDL-C levels in some patients. In general, fish oils are well tolerated, with the major side effect being dyspepsia. They appear to be safe, at least at doses up to 3–4 g, but can be associated with a prolongation in the bleeding time.

NICOTINIC ACID Nicotinic acid, or niacin, is a B-complex vitamin that has been used as a lipid-modifying agent for more than five decades. Niacin suppresses lipolysis in the adipocyte through its effect on the niacin receptor GPR109A and has other effects on hepatic lipid metabolism that are poorly understood. Niacin reduces plasma TG and LDL-C levels and also raises the plasma concentration of HDL-C. Because it has a number of side effects and can be difficult to use, it is at best a third-line agent for the management of severe hypertriglyceridemia. Niacin therapy is generally started at lower doses and gradually titrated up to higher doses. The most frequent side effect of niacin is cutaneous flushing, which is mediated by activating GPR109A in the skin. Niacin can cause dyspepsia and can exacerbate esophageal reflux and peptic ulcer disease. Mild elevations in transaminases occur in up to 15% of patients treated

with any form of niacin. Niacin can raise plasma levels of uric acid and precipitate gouty attacks in susceptible patients. Acanthosis nigricans, a dark-colored coarse skin lesion, and maculopathy are infrequent side effects of niacin.

MANAGEMENT OF CHOLESTEROL TO PREVENT CARDIOVASCULAR DISEASE

In contrast to hypertriglyceridemia and pancreatitis, there are abundant and compelling data that intervention to reduce LDL-C substantially reduces the risk of CVD, including myocardial infarction and stroke, as well as total mortality. Thus, it is imperative that patients with hypercholesterolemia be assessed for cardiovascular risk and for the need for intervention. It is also worth noting that patients at high risk for CVD who have plasma LDL-C levels in the “normal” or average range also benefit from intervention to reduce LDL-C levels.

Lifestyle The first approach to a patient with hypercholesterolemia and high cardiovascular risk is to make any necessary lifestyle changes. In obese patients, efforts should be made to reduce body weight to the ideal level. Patients should receive dietary counseling to reduce the content of saturated fats, *trans* fats, and cholesterol in the diet. Regular aerobic exercise has relatively little impact on reducing plasma LDL-C levels, although it has cardiovascular benefits independent of LDL lowering.

Pharmacologic Therapy for Hypercholesterolemia The decision to use LDL-lowering drug therapy (Table 421-5)—with a statin being first-line therapy—depends on the level of LDL-C as well as the level of cardiovascular risk. In general, patients with a Mendelian disorder of elevated LDL-C such as FH must be treated to reduce the very high lifetime risk of CVD, and treatment should be initiated as early as possible in adulthood or, in some cases, during childhood.

Otherwise, the decision to initiate LDL-lowering drug therapy is generally determined by the level of cardiovascular risk. In patients with established CVD, statin therapy is well supported by clinical trial data and should be used regardless of the LDL-C level. For patients >40 years old without clinical CVD, the AHA/ACC risk calculator (http://my.americanheart.org/professional/StatementsGuidelines/PreventionGuidelines/Prevention-Guidelines_UCM_457698_SubHomePage.jsp) can be used to determine the 10-year absolute risk for CVD, and current guidelines suggest that a 10-year risk >7.5% merits consideration of statin therapy regardless of plasma LDL-C level. For younger patients, the assessment of lifetime risk of CVD may help inform the decision to start a statin.

HMG-CoA REDUCTASE INHIBITORS (STATINS) Statins inhibit HMG-CoA reductase, a key enzyme in cholesterol biosynthesis. By inhibiting cholesterol biosynthesis, statins lead to increased hepatic LDL receptor activity and accelerated clearance of circulating LDL, resulting in a dose-dependent reduction in plasma levels of LDL-C. The magnitude of LDL lowering associated with statin treatment varies widely among individuals, but once a patient is on a statin, the doubling of the statin dose produces an ~6% further reduction in the level of plasma LDL-C. The statins currently available differ in their LDL-C-reducing potency (Table 421-5). Currently, there is no convincing evidence that any of the different statins confer an advantage that is independent of the effect on LDL-C. Statins also reduce plasma TGs in a dose-dependent fashion, which is roughly proportional to their LDL-C-lowering effects (if the TGs are <400 mg/dL). Statins have a modest HDL-raising effect (5–10%) that is not generally dose-dependent.

Statins are well tolerated and can be taken in tablet form once a day. Potential side effects include dyspepsia, headaches, fatigue, and muscle or joint pains. Severe myopathy and even rhabdomyolysis occur rarely with statin treatment. The risk of statin-associated myopathy is increased by the presence of older age, frailty, renal insufficiency, and coadministration of drugs that interfere with the metabolism of statins, such as erythromycin and related antibiotics, antifungal agents, immunosuppressive drugs, and fibric acid deriva-

TABLE 421-5 SUMMARY OF THE MAJOR APPROVED DRUGS USED FOR THE TREATMENT OF DYSLIPIDEMIA

Drug	Major Indications	Starting Dose	Maximal Dose	Mechanism	Common Side Effects
HMG-CoA reductase inhibitors (statins)	Elevated LDL-C; increased CV risk			↓ Cholesterol synthesis, ↑ Hepatic LDL receptors, ↓ VLDL production	Myalgias, arthralgias, elevated transaminases, dyspepsia
Lovastatin		20–40 mg daily	80 mg daily		
Pravastatin		40–80 mg daily	80 mg daily		
Simvastatin		20–40 mg daily	80 mg daily		
Fluvastatin		20–40 mg daily	80 mg daily		
Atorvastatin		20–40 mg daily	80 mg daily		
Rosuvastatin		5–20 mg daily	40 mg daily		
Pitavastatin		1–2 mg daily	4 mg daily		
Cholesterol absorption inhibitor	Elevated LDL-C			↓ Cholesterol absorption, ↑ LDL receptors	Elevated transaminases
Ezetimibe		10 mg daily	10 mg daily		
Bile acid sequestrants	Elevated LDL-C			↑ Bile acid excretion and ↑ LDL receptors	Bloating, constipation, elevated triglycerides
Cholestyramine		4 g daily	32 g daily		
Colestipol		5 g daily	40 g daily		
Colesevelam		3750 mg daily	4375 mg daily		
MTP inhibitor	HoFH			↓ VLDL production	Nausea, diarrhea, increased hepatic fat
Lomitapide		5 mg daily	60 mg daily		
ApoB inhibitor	HoFH			↓ VLDL production	Injection site reactions, flu-like symptoms, increased hepatic fat
Mipomersen		200 mg SC weekly	200 mg SC weekly		
Nicotinic acid	Elevated LDL-C, elevated TG			↓ VLDL production	Cutaneous flushing, GI upset, elevated glucose, uric acid, and elevated liver function tests
Immediate-release		100 mg tid	1 g tid		
Sustained-release		250 mg bid	1.5 g bid		
Extended-release		500 mg qhs	2 g qhs		
Fibric acid derivatives	Elevated TG			↑ LPL, ↓ VLDL synthesis	Dyspepsia, myalgia, gallstones, elevated transaminases
Gemfibrozil		600 mg bid	600 mg bid		
Fenofibrate		145 mg qd	145 mg qd		
Omega-3 fatty acids	Elevated TG			↑ TG catabolism	Dyspepsia, fishy odor to breath
Omega-3 acid ethyl esters		4 g daily	4 g daily		
Icosapent ethyl		4 g daily	4 g daily		

Abbreviations: GI, gastrointestinal; HDL-C, high-density lipoprotein cholesterol; HoFH, homozygous familial hypercholesterolemia; LDL, low-density lipoprotein; LDL-C, LDL-cholesterol; LPL, lipoprotein lipase; TG, triglyceride; VLDL, very-low-density lipoprotein.

tives (particularly gemfibrozil). Severe myopathy can usually be avoided by careful patient selection, avoidance of interacting drugs, and instructing the patient to contact the physician immediately in the event of unexplained muscle pain. In the event of muscle symptoms, the plasma creatine kinase (CK) level should be obtained to differentiate myopathy from myalgia. Serum CK levels need not be monitored on a routine basis in patients taking statins, because an elevated CK in the absence of symptoms does not predict the development of myopathy and does not necessarily suggest the need for discontinuing the drug.

Another consequence of statin therapy can be elevation in liver transaminases (alanine aminotransferase [ALT] and aspartate aminotransferase [AST]). They should be checked before starting therapy, at 2–3 months, and then annually. Substantial (greater than three times the upper limit of normal) elevation in transaminases is relatively rare, and mild-to-moderate (one to three times normal) elevation in transaminases in the absence of symptoms need not mandate discontinuing the medication. Severe clinical hepatitis associated with statins is exceedingly rare, and the trend is toward

less frequent monitoring of transaminases in patients taking statins. The statin-associated elevation in liver enzymes resolves upon discontinuation of the medication.

Statins appear to be remarkably safe. Meta-analyses of large randomized controlled clinical trials with statins do not suggest an increase in any major noncardiac diseases except type 2 diabetes. A small excess percentage of those taking statins will develop diabetes but the benefits associated with the reduction in cardiovascular events outweigh the increase in incidence of diabetes. Statins are the drug class of choice for LDL-C reduction and are by far the most widely used class of lipid-lowering drugs.

CHOLESTEROL ABSORPTION INHIBITORS Cholesterol within the lumen of the small intestine is derived from the diet (about one-third) and the bile (about two-thirds) and is actively absorbed by the enterocyte through a process that involves the protein NPC1L1. Ezetimibe (Table 421-5) is a cholesterol absorption inhibitor that binds directly to and inhibits NPC1L1 and blocks the intestinal absorption of cholesterol. Ezetimibe (10 mg) inhibits cholesterol

absorption by almost 60%, resulting in a reduction in delivery of dietary sterols in the liver and an increase in hepatic LDL receptor expression. The mean reduction in plasma LDL-C on ezetimibe (10 mg) is 18%, and the effect is additive when used in combination with a statin. Effects on TG and HDL-C levels are negligible. When used in combination with a statin, monitoring of liver transaminases is recommended. The only roles for ezetimibe in monotherapy are in patients who do not tolerate statins and in sitosterolemia.

BILE ACID SEQUESTRANTS (RESINS) Bile acid sequestrants bind bile acids in the intestine and promote their excretion rather than reabsorption in the ileum. To maintain the bile acid pool size, the liver diverts cholesterol to bile acid synthesis. The decreased hepatic intracellular cholesterol content results in upregulation of the LDL receptor and enhanced LDL clearance from the plasma. Bile acid sequestrants, including cholestyramine, colestipol, and colesevelam (Table 421-5), primarily reduce plasma LDL-C levels but can cause an increase in plasma TGs. Therefore, patients with hypertriglyceridemia generally should not be treated with bile acid-binding resins. Cholestyramine and colestipol are insoluble resins that must be suspended in liquids. Colesevelam is available as tablets but generally requires up to six to seven tablets per day for effective LDL-C lowering. Most side effects of resins are limited to the gastrointestinal tract and include bloating and constipation. Because bile acid sequestrants are not systemically absorbed, they are very safe and the cholesterol-lowering drug of choice in children and in women of childbearing age who are lactating, pregnant, or could become pregnant. They are effective in combination with statins and in combination with ezetimibe and are particularly useful with one or both of these drugs for patients with severe hypercholesterolemia or those with statin intolerance.

SPECIALIZED DRUGS FOR HOMOZYGOUS FH Two “orphan” drugs are approved specifically for the management of homozygous FH. They include a small-molecule inhibitor of MTP, called lomitapide, and an antisense oligonucleotide against apoB, called mipomersen. These drugs reduce VLDL production and LDL-C levels in homozygous FH patients. Due to their mechanism of action, each drug causes an increase in hepatic fat, the long-term consequences of which are unknown. In addition, lomitapide is associated with gastrointestinal-related side effects, and mipomersen is associated with skin reactions and flu-like symptoms.

LDL APHERESIS Patients who remain severely hypercholesterolemic despite optimally tolerated drug therapy are candidates for LDL apheresis. In this process, the patient’s plasma is passed over a column that selectively removes the LDL, and the LDL-depleted plasma is returned to the patient. Patients on maximally tolerated combination drug therapy who have CHD and a plasma LDL-C level >200 mg/dL or no CHD and a plasma LDL-C level >300 mg/dL are candidates for every-other-week LDL apheresis and should be referred to a specialized lipid center.

of high-density lipoprotein (HDL) cholesterol, hyperglycemia, and hypertension (Table 422-1).

EPIDEMIOLOGY



The most challenging feature of the metabolic syndrome to define is waist circumference. Intraabdominal circumference (visceral adipose tissue) is considered most strongly related to insulin resistance and risk of diabetes and CVD, and for any given waist circumference the distribution of adipose tissue between SC and visceral depots varies substantially. Thus, within and between populations, there is a lesser vs. greater risk at the same waist circumference. These differences in populations are reflected in the range of waist circumferences considered to confer risk in different geographic locations (Table 422-1).

The prevalence of the metabolic syndrome varies around the world, in part reflecting the age and ethnicity of the populations studied and the diagnostic criteria applied. In general, the prevalence of the metabolic syndrome increases with age. The highest recorded prevalence worldwide is among Native Americans, with nearly 60% of women ages 45–49 and 45% of men ages 45–49 meeting the criteria of the National Cholesterol Education Program and Adult Treatment Panel III (NCEP:ATPIII). In the United States, the metabolic syndrome is less common among African-American men and more common among Mexican-American women. Based on data from the National Health and Nutrition Examination Survey (NHANES) 2003–2006, the age-adjusted prevalence of the metabolic syndrome in U.S. adults without diabetes is 28% for men and 30% for women. In France, studies of a cohort of 30- to 60-year-olds have shown a <10% prevalence for each sex, although 17.5% of people 60–64 years of age are affected. Greater global industrialization is associated with rising rates of obesity, which are expected to increase the prevalence of the metabolic syndrome dramatically, especially as the population ages. Moreover, the rising prevalence and severity of obesity among children is reflected in features of the metabolic syndrome in a younger population.

The frequency distribution of the five components of the syndrome for the U.S. population (NHANES III) is summarized in Fig. 422-1. Increases in waist circumference predominate among women, whereas increases in fasting plasma triglyceride levels (i.e., to >150 mg/dL), reductions in HDL cholesterol levels, and hyperglycemia are more likely in men.

RISK FACTORS

Overweight/Obesity Although the metabolic syndrome was first described in the early twentieth century, the worldwide overweight/obesity epidemic has recently been the force driving its increasing recognition. Central adiposity is a key feature of the syndrome, and the syndrome’s prevalence reflects the strong relationship between waist circumference and increasing adiposity. However, despite the importance of obesity, patients who are of normal weight may also be insulin resistant and may have the metabolic syndrome.

Sedentary Lifestyle Physical inactivity is a predictor of CVD events and the related risk of death. Many components of the metabolic syndrome are associated with a sedentary lifestyle, including increased adipose tissue (predominantly central), reduced HDL cholesterol, and increased triglycerides, blood pressure, and glucose in genetically susceptible persons. Compared with individuals who watch television or videos or use the computer <1 h daily, those who do so for >4 h daily have a twofold increased risk of the metabolic syndrome.

Aging The metabolic syndrome affects nearly 50% of the U.S. population older than age 50, and at >60 years of age women are more often affected than men. The age dependency of the syndrome’s prevalence is seen in most populations around the world.

Diabetes Mellitus Diabetes mellitus is included in both the NCEP and the harmonizing definitions of the metabolic syndrome. It is estimated that the great majority (~75%) of patients with type 2 diabetes or impaired glucose tolerance have the metabolic syndrome. The presence of the metabolic syndrome in these populations relates to a

422 The Metabolic Syndrome

Robert H. Eckel

The metabolic syndrome (syndrome X, insulin resistance syndrome) consists of a constellation of metabolic abnormalities that confer increased risk of cardiovascular disease (CVD) and diabetes mellitus. Evolution of the criteria for the metabolic syndrome since the original definition by the World Health Organization in 1998 reflects growing clinical evidence and analysis by a variety of consensus conferences and professional organizations. The major features of the metabolic syndrome include central obesity, hypertriglyceridemia, low levels

NCEP:ATPIII 2001	Harmonizing Definition ^b		
Three or more of the following:	Three of the following:		
<ul style="list-style-type: none"> Central obesity: waist circumference >102 cm (M), >88 cm (F) Hypertriglyceridemia: triglyceride level \geq150 mg/dL or specific medication Low HDL^c cholesterol: <40 mg/dL and <50 mg/dL for men and women, respectively, or specific medication Hypertension: blood pressure \geq130 mmHg systolic or \geq85 mmHg diastolic or specific medication Fasting plasma glucose level \geq100 mg/dL or specific medication or previously diagnosed type 2 diabetes 	<ul style="list-style-type: none"> Waist circumference (cm) 		
	Men	Women	Ethnicity
	\geq 94	\geq 80	Europid, sub-Saharan African, Eastern and Middle Eastern
	\geq 90	\geq 80	South Asian, Chinese, and ethnic South and Central American
	\geq 85	\geq 90	Japanese
	<ul style="list-style-type: none"> Fasting triglyceride level >150 mg/dL or specific medication HDL cholesterol level <40 mg/dL and <50 mg/dL for men and women, respectively, or specific medication Blood pressure >130 mm systolic or >85 mm diastolic or previous diagnosis or specific medication Fasting plasma glucose level \geq100 mg/dL (alternative indication: drug treatment of elevated glucose levels) 		

^aNational Cholesterol Education Program and Adult Treatment Panel III. ^bIn this analysis, the following thresholds for waist circumference were used: white men, \geq 94 cm; African-American men, \geq 94 cm; Mexican-American men, \geq 90 cm; white women, \geq 80 cm; African-American women, \geq 80 cm; Mexican-American women, \geq 80 cm. For participants whose designation was "other race—including multiracial," thresholds that were once based on Europid cutoffs (\geq 94 cm for men and \geq 80 cm for women) and on South Asian cutoffs (\geq 90 cm for men and \geq 80 cm for women) were used. For participants who were considered "other Hispanic," the International Diabetes Federation thresholds for ethnic South and Central Americans were used. ^cHigh-density lipoprotein.

higher prevalence of CVD than in patients who have type 2 diabetes or impaired glucose tolerance but do not have this syndrome.

Cardiovascular Disease Individuals with the metabolic syndrome are twice as likely to die of cardiovascular disease as those who do not, and their risk of an acute myocardial infarction or stroke is three-fold higher. The approximate prevalence of the metabolic syndrome among patients with coronary heart disease (CHD) is 50%, with a prevalence of ~35% among patients with premature coronary artery disease (before or at age 45) and a particularly high prevalence among women. With appropriate cardiac rehabilitation and changes in lifestyle (e.g., nutrition, physical activity, weight reduction, and—in some cases—pharmacologic therapy), the prevalence of the syndrome can be reduced.

Lipodystrophy Lipodystrophic disorders in general are associated with the metabolic syndrome. Both genetic lipodystrophy (e.g., Berardinelli-Seip congenital lipodystrophy, Dunnigan familial partial lipodystrophy) and acquired lipodystrophy (e.g., HIV-related lipodystrophy in

patients receiving antiretroviral therapy) may give rise to severe insulin resistance and many of the components of the metabolic syndrome.

ETIOLOGY

Insulin Resistance The most accepted and unifying hypothesis to describe the pathophysiology of the metabolic syndrome is insulin resistance, which is caused by an incompletely understood defect in insulin action (Chap. 417). The onset of insulin resistance is heralded by postprandial hyperinsulinemia, which is followed by fasting hyperinsulinemia and ultimately by hyperglycemia.

An early major contributor to the development of insulin resistance is an overabundance of circulating fatty acids (Fig. 422-2). Plasma albumin-bound free fatty acids are derived predominantly from adipose-tissue triglyceride stores released by intracellular lipolytic enzymes. Fatty acids are also derived from the lipolysis of triglyceride-rich lipoproteins in tissues by lipoprotein lipase. Insulin mediates both antilipolysis and the stimulation of lipoprotein lipase in adipose tissue. Of note, the inhibition of lipolysis in adipose tissue is the most sensitive pathway of insulin action. Thus, when insulin resistance develops, increased lipolysis produces more fatty acids, which further decrease the antilipolytic effect of insulin. Excessive fatty acids enhance substrate availability and create insulin resistance by modifying downstream signaling. Fatty acids impair insulin-mediated glucose uptake and accumulate as triglycerides in both skeletal and cardiac muscle, whereas increased glucose production and triglyceride accumulation take place in the liver.

Leptin resistance has also been raised as a possible pathophysiologic mechanism to explain the metabolic syndrome. Physiologically, leptin reduces appetite, promotes energy expenditure, and enhances insulin sensitivity. In addition, leptin may regulate cardiac and vascular function through a nitric oxide-dependent mechanism. However, when obesity develops, hyperleptinemia ensues, with evidence of leptin resistance in the brain and other tissues resulting in inflammation, insulin resistance, hyperlipidemia, and a plethora of cardiovascular disorders, such as hypertension, atherosclerosis, CHD, and heart failure.

The oxidative stress hypothesis provides a unifying theory for aging and the predisposition to the metabolic syndrome. In studies of insulin-resistant individuals with obesity or type 2 diabetes, the offspring of patients with type 2 diabetes, and the elderly, a defect in mitochondrial oxidative phosphorylation that leads to the accumulation of triglycerides and related lipid molecules in muscle has been identified.

Recently, the gut microbiome has emerged as an important contributor to the development of obesity and related metabolic disorders, including the metabolic syndrome. Although the mechanism remains

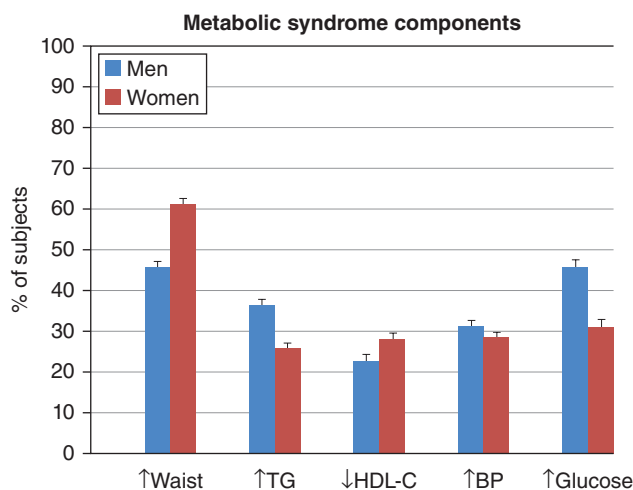


FIGURE 422-1 Prevalence of the metabolic syndrome components, from NHANES 2003–2006. NHANES, National Health and Nutrition Examination Survey; TG, triglyceride; HDL-C, high-density lipoprotein cholesterol; BP, blood pressure. The prevalence of elevated glucose includes individuals with known diabetes mellitus. (Created from data in ES Ford et al: *J Diabetes* 2:1753, 2010.)

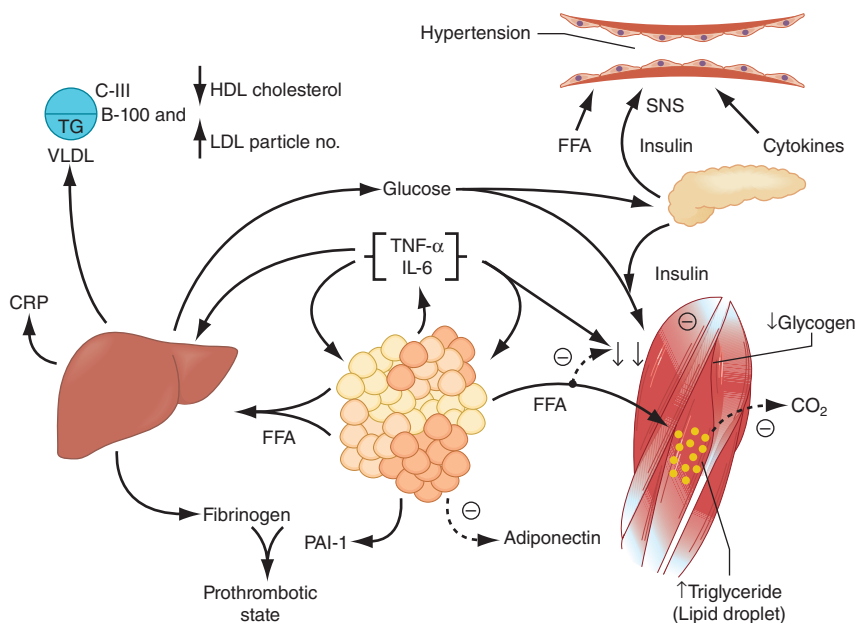


FIGURE 422-2 Pathophysiology of the metabolic syndrome. Free fatty acids (FFAs) are released in abundance from an expanded adipose tissue mass. In the liver, FFAs result in increased production of glucose and triglycerides and secretion of very low density lipoproteins (VLDLs). Associated lipid/lipoprotein abnormalities include reductions in high-density lipoprotein (HDL) cholesterol and an increased low-density lipoprotein (LDL) particle number (no.). FFAs also reduce insulin sensitivity in muscle by inhibiting insulin-mediated glucose uptake. Associated defects include a reduction in glucose partitioning to glycogen and increased lipid accumulation in triglyceride (TG). The increase in circulating glucose, and to some extent FFAs, increases pancreatic insulin secretion, resulting in hyperinsulinemia. Hyperinsulinemia may result in enhanced sodium reabsorption and increased sympathetic nervous system (SNS) activity and contribute to hypertension, as might higher levels of circulating FFAs. The proinflammatory state is superimposed and contributory to the insulin resistance produced by excessive FFAs. The enhanced secretion of interleukin 6 (IL-6) and tumor necrosis factor α (TNF- α) produced by adipocytes and monocyte-derived macrophages results in more insulin resistance and lipolysis of adipose tissue triglyceride stores to circulating FFAs. IL-6 and other cytokines also enhance hepatic glucose production, VLDL production by the liver, hypertension and insulin resistance in muscle. Cytokines and FFAs also increase hepatic production of fibrinogen and adipocyte production of plasminogen activator inhibitor 1 (PAI-1), resulting in a prothrombotic state. Higher levels of circulating cytokines stimulate hepatic production of C-reactive protein (CRP). Reduced production of the anti-inflammatory and insulin-sensitizing cytokine adiponectin is also associated with the metabolic syndrome. (Modified from RH Eckel et al: *Lancet* 365:1415, 2005.)

uncertain, interaction among genetic predisposition, diet, and the intestinal flora is important.

Increased Waist Circumference Waist circumference is an important component of the most recent and frequently applied diagnostic criteria for the metabolic syndrome. However, measuring waist circumference does not reliably distinguish increases in SC adipose tissue from those in visceral fat; this distinction requires CT or MRI. With increases in visceral adipose tissue, adipose tissue-derived free fatty acids are directed to the liver. In contrast, increases in abdominal SC fat release lipolysis products into the systemic circulation and avert more direct effects on hepatic metabolism. Relative increases in visceral versus SC adipose tissue with increasing waist circumference in Asians and Asian Indians may explain the greater prevalence of the syndrome in those populations than in African-American men, in whom SC fat predominates. It is also possible that visceral fat is a marker for—but not the source of—excess postprandial free fatty acids in obesity.

Dyslipidemia (See also Chap. 421) In general, free fatty acid flux to the liver is associated with increased production of ApoB-containing, triglyceride-rich, very low-density lipoproteins (VLDLs). The effect of insulin on this process is complex, but *hypertriglyceridemia* is an

excellent marker of the insulin-resistant condition. Not only is hypertriglyceridemia a feature of the metabolic syndrome, but patients with the metabolic syndrome have elevated levels of ApoCIII carried on VLDLs and other lipoproteins. This increase in ApoCIII is inhibitory to lipoprotein lipase, further contributing to hypertriglyceridemia and also associated with more atherosclerotic cardiovascular disease.

The other major lipoprotein disturbance in the metabolic syndrome is a *reduction in HDL cholesterol*. This reduction is a consequence of changes in HDL composition and metabolism. In the presence of hypertriglyceridemia, a decrease in the cholesterol content of HDL is a consequence of reduced cholesteryl ester content of the lipoprotein core in combination with cholesteryl ester transfer protein-mediated alterations in triglyceride that make the particle small and dense. This change in lipoprotein composition also results in increased clearance of HDL from the circulation. These changes in HDL have a relationship to insulin resistance that is probably indirect, occurring in concert with the changes in triglyceride-rich lipoprotein metabolism.

In addition to HDLs, low-density lipoproteins (LDLs) are modified in composition in the metabolic syndrome. With fasting serum triglycerides at >2.0 mM (~ 180 mg/dL), there is almost always a predominance of small, dense LDLs, which are thought to be more atherogenic although their association with hypertriglyceridemia and low HDLs make their independent contribution to CVD events difficult to assess. Individuals with hypertriglyceridemia often have increases in cholesterol content of both VLDL1 and VLDL2 subfractions and in LDL particle number. Both of these lipoprotein changes may contribute to atherogenic risk in patients with the metabolic syndrome.

Glucose Intolerance (See also Chap. 417) Defects in insulin action in the metabolic syndrome lead to impaired suppression of glucose production by the liver and kidney and reduced glucose uptake and metabolism in insulin-sensitive tissues—i.e., muscle and adipose tissue. The relationship between impaired fasting glucose or impaired glucose tolerance and insulin resistance is well supported by studies of humans, nonhuman primates, and rodents. To

compensate for defects in insulin action, insulin secretion and/or clearance must be modified so that euglycemia is sustained. Ultimately, this compensatory mechanism fails, usually because of defects in insulin secretion, resulting in progression from impaired fasting glucose and/or impaired glucose tolerance to diabetes mellitus.

Hypertension The relationship between insulin resistance and hypertension is well established. Paradoxically, under normal physiologic conditions, insulin is a vasodilator with secondary effects on sodium reabsorption in the kidney. However, in the setting of insulin resistance, the vasodilatory effect of insulin is lost but the renal effect on sodium reabsorption is preserved. Sodium reabsorption is increased in whites with the metabolic syndrome but not in Africans or Asians. Insulin also increases the activity of the sympathetic nervous system, an effect that may be preserved in the setting of insulin resistance. Insulin resistance is characterized by pathway-specific impairment in phosphatidylinositol-3-kinase signaling. In the endothelium, this impairment may cause an imbalance between the production of nitric oxide and the secretion of endothelin 1, with a consequent decrease in blood flow. Although these mechanisms are provocative, evaluation of insulin action by measurement of fasting insulin levels or by homeostasis model assessment shows that insulin

2452 resistance contributes only partially to the increased prevalence of hypertension in the metabolic syndrome.

Another possible mechanism underlying hypertension in the metabolic syndrome is the vasoactive role of perivascular adipose tissue. Reactive oxygen species released by NADPH oxidase impair endothelial function and result in local vasoconstriction. Other paracrine effects could be mediated by leptin or other proinflammatory cytokines released from adipose tissue, such as tumor necrosis factor α .

Hyperuricemia is another consequence of insulin resistance and is commonly observed in the metabolic syndrome. There is growing evidence not only that uric acid is associated with hypertension but also that reduction of uric acid normalizes blood pressure in hyperuricemic adolescents with hypertension. The mechanism appears to be related to an adverse effect of uric acid on nitric acid synthase in the macula densa of the kidney and stimulation of the renin-angiotensin aldosterone system.

Proinflammatory Cytokines The increases in proinflammatory cytokines—including interleukins 1, 6, and 18; resistin; tumor necrosis factor α ; and the systemic biomarker C-reactive protein—reflect overproduction by the expanded adipose tissue mass (Fig. 422-2). Adipose tissue–derived macrophages may be the primary source of proinflammatory cytokines locally and in the systemic circulation. It remains unclear, however, how much of the insulin resistance is caused by the paracrine effects of these cytokines and how much by the endocrine effects.

Adiponectin Adiponectin is an anti-inflammatory cytokine produced exclusively by adipocytes. Adiponectin enhances insulin sensitivity and inhibits many steps in the inflammatory process. In the liver, adiponectin inhibits the expression of gluconeogenic enzymes and the rate of glucose production. In muscle, adiponectin increases glucose transport and enhances fatty acid oxidation, partially through the activation of AMP kinase. Adiponectin levels are reduced in the metabolic syndrome. The relative contributions of adiponectin deficiency and overabundance of the proinflammatory cytokines are unclear.

CLINICAL FEATURES

Symptoms and Signs The metabolic syndrome typically is not associated with symptoms. On physical examination, waist circumference may be expanded and blood pressure elevated. The presence of either or both of these signs should prompt the clinician to search for other biochemical abnormalities that may be associated with the metabolic syndrome. Less frequently, lipoatrophy or acanthosis nigricans is found on examination. Because these physical findings characteristically are associated with severe insulin resistance, other components of the metabolic syndrome should be expected.

Associated Diseases • CARDIOVASCULAR DISEASE The relative risk for new-onset CVD in patients with the metabolic syndrome who do not have diabetes averages 1.5–3 fold. However, an 8-year follow-up of middle-aged participants in the Framingham Offspring Study documented that the population-attributable CVD risk in the metabolic syndrome was 34% among men and only 16% among women. In the same study, both the metabolic syndrome and diabetes predicted ischemic stroke, with greater risk among patients with the metabolic syndrome than among those with diabetes alone (19% vs. 7%) and a particularly large difference among women (27% vs. 5%). Patients with the metabolic syndrome are also at increased risk for peripheral vascular disease.

TYPE 2 DIABETES Overall, the risk for type 2 diabetes among patients with the metabolic syndrome is increased three- to fivefold. In the Framingham Offspring Study's 8-year follow-up of middle-aged participants, the population-attributable risk for developing type 2 diabetes was 62% among men and 47% among women.

Other Associated Conditions In addition to the features specifically associated with the metabolic syndrome, other metabolic alterations accompany insulin resistance. Those alterations include increases in ApoB and ApoCIII, uric acid, prothrombotic factors (fibrinogen, plasminogen activator inhibitor 1), serum viscosity, asymmetric

dimethylarginine, homocysteine, white blood cell count, proinflammatory cytokines, C-reactive protein, microalbuminuria, nonalcoholic fatty liver disease and/or nonalcoholic steatohepatitis, polycystic ovary syndrome, and obstructive sleep apnea.

NONALCOHOLIC FATTY LIVER DISEASE (SEE ALSO CHAP. 367e) Fatty liver is a relatively common condition, affecting 25–45% of the U.S. population. However, in nonalcoholic steatohepatitis, triglyceride accumulation and inflammation coexist. Nonalcoholic steatohepatitis is now present in 3–12% of the population of the United States and other Western countries. Of patients with the metabolic syndrome, ~25–60% have nonalcoholic fatty liver disease and up to 35% have nonalcoholic steatohepatitis. As the prevalence of overweight/obesity and the metabolic syndrome increases, nonalcoholic steatohepatitis may become one of the more common causes of end-stage liver disease and hepatocellular carcinoma.

HYPERURICEMIA (SEE ALSO CHAP. 431e) Hyperuricemia reflects defects in insulin action on the renal tubular reabsorption of uric acid and may contribute to hypertension through its effect on the endothelium. An increase in asymmetric dimethylarginine, an endogenous inhibitor of nitric oxide synthase, also relates to endothelial dysfunction. In addition, microalbuminuria may be caused by altered endothelial pathophysiology in the insulin-resistant state.

POLYCYSTIC OVARY SYNDROME (SEE ALSO CHAP. 412) Polycystic ovary syndrome is highly associated with insulin resistance (50–80%) and the metabolic syndrome, with a prevalence of the syndrome between 40% and 50%. Women with polycystic ovary syndrome are two to four times more likely to have the metabolic syndrome than are women without polycystic ovary syndrome.

OBSTRUCTIVE SLEEP APNEA (SEE ALSO CHAP. 38) Obstructive sleep apnea is commonly associated with obesity, hypertension, increased circulating cytokines, impaired glucose tolerance, and insulin resistance. With these associations, it is not surprising that individuals with obstructive sleep apnea frequently have the metabolic syndrome. Moreover, when biomarkers of insulin resistance are compared between patients with obstructive sleep apnea and weight-matched controls, insulin resistance is found to be more severe in those with apnea. Continuous positive airway pressure treatment improves insulin sensitivity in patients with obstructive sleep apnea.

DIAGNOSIS

The diagnosis of the metabolic syndrome relies on fulfillment of the criteria listed in Table 422-1, as assessed using tools at the bedside and in the laboratory. The medical history should include evaluation of symptoms for obstructive sleep apnea in all patients and polycystic ovary syndrome in premenopausal women. Family history will help determine risk for CVD and diabetes mellitus. Blood pressure and waist circumference measurements provide information necessary for the diagnosis.

Laboratory Tests Measurement of fasting lipids and glucose is needed in determining whether the metabolic syndrome is present. The measurement of additional biomarkers associated with insulin resistance can be individualized. Such tests might include those for ApoB, high-sensitivity C-reactive protein, fibrinogen, uric acid, urinary microalbumin, and liver function. A sleep study should be performed if symptoms of obstructive sleep apnea are present. If polycystic ovary syndrome is suspected on the basis of clinical features and anovulation, testosterone, luteinizing hormone, and follicle-stimulating hormone should be measured.

TREATMENT THE METABOLIC SYNDROME

LIFESTYLE (SEE ALSO CHAP. 416)

Obesity is the driving force behind the metabolic syndrome. Thus, weight reduction is the primary approach to the disorder. With weight reduction, improvement in insulin sensitivity is often accompanied by favorable modifications in many components of the

metabolic syndrome. In general, recommendations for weight loss include a combination of caloric restriction, increased physical activity, and behavior modification. Caloric restriction is the most important component, whereas increases in physical activity are important for maintenance of weight loss. Some but not all evidence suggests that the addition of exercise to caloric restriction may promote greater weight loss from the visceral depot. The tendency for weight regain after successful weight reduction underscores the need for long-lasting behavioral changes.

Diet Before prescribing a weight-loss diet, it is important to emphasize that it has taken the patient a long time to develop an expanded fat mass; thus, the correction need not occur quickly. Given that ~ 3500 kcal = 1 lb of fat, ~ 500 -kcal restriction daily equates to weight reduction of 1 lb per week. Diets restricted in carbohydrate typically provide a rapid initial weight loss. However, after 1 year, the amount of weight reduction is minimally reduced or no different from that with caloric restriction alone. Thus, adherence to the diet is more important than which diet is chosen. Moreover, there is concern about low-carbohydrate diets enriched in saturated fat, particularly for patients at risk for CVD. Therefore, a high-quality dietary pattern—i.e., a diet enriched in fruits, vegetables, whole grains, lean poultry, and fish—should be encouraged to maximize overall health benefit.

Physical Activity Before a physical activity recommendation is provided to patients with the metabolic syndrome, it is important to ensure that the increased activity does not incur risk. Some high-risk patients should undergo formal cardiovascular evaluation before initiating an exercise program. For an inactive participant, gradual increases in physical activity should be encouraged to enhance adherence and avoid injury. Although increases in physical activity can lead to modest weight reduction, 60–90 min of daily activity is required to achieve this goal. Even if an overweight or obese adult is unable to undertake this level of activity, a significant health benefit will follow from at least 30 min of moderate-intensity activity daily. The caloric value of 30 min of a variety of activities can be found at www.heart.org/HEARTORG/GettingHealthy/WeightManagement/LosingWeight/Losing-Weight_UCM_307904_Article.jsp. Of note, a variety of routine activities, such as gardening, walking, and housecleaning, require moderate caloric expenditure. Thus, physical activity need not be defined solely in terms of formal exercise such as jogging, swimming, or tennis.

Behavior Modification Behavioral treatment typically includes recommendations for dietary restriction and more physical activity, resulting in weight loss that benefits metabolic health. The subsequent challenge is the duration of the program because weight regain so often follows successful weight reduction. Long-term outcomes may be enhanced by a variety of methods, such as the Internet, social media, and telephone follow-up to maintain contact between providers and patients.

Obesity (See also Chap. 416) In some patients with the metabolic syndrome, treatment options need to extend beyond lifestyle intervention. Weight-loss drugs come in two major classes: appetite suppressants and absorption inhibitors. Appetite suppressants approved by the U.S. Food and Drug Administration include phentermine (for short-term use [3 months] only) as well as the more recent additions phentermine/topiramate and lorcaserin, which are approved without restrictions on the duration of therapy. In clinical trials, the phentermine/topiramate combination has resulted in $\sim 10\%$ weight loss in 50% of patients. Side effects include palpitations, headache, paresthesias, constipation, and insomnia. Lorcaserin results in less weight loss—typically $\sim 5\%$ beyond placebo—but can cause headache and nasopharyngitis. Orlistat inhibits fat absorption by $\sim 30\%$ and is moderately effective compared with placebo ($\sim 5\%$ more weight loss). Orlistat has been shown to reduce the incidence of type 2 diabetes, an effect that was especially evident among patients with impaired glucose tolerance at baseline. This drug is often difficult of take because of oily leakage per rectum.

Metabolic or bariatric surgery is an option for patients with the metabolic syndrome who have a body mass index >40 kg/m², or >35 kg/m² with comorbidities. An evolving application for metabolic surgery includes patients with a body mass index as low as 30 kg/m² and type 2 diabetes. Gastric bypass or vertical sleeve gastrectomy results in dramatic weight reduction and improvement in the features of the metabolic syndrome. A survival benefit with gastric bypass has also been realized.

LDL CHOLESTEROL (SEE ALSO CHAP. 421)

The rationale for the NCEP:ATPIII's development of criteria for the metabolic syndrome was to go beyond LDL cholesterol in identifying and reducing the risk of CVD. The working assumption by the panel was that LDL cholesterol goals had already been achieved and that increasing evidence supports a linear reduction in CVD events as a result of progressive lowering of LDL cholesterol with statins. For patients with the metabolic syndrome and diabetes, a statin should be prescribed. For those patients with diabetes and known CVD, the current evidence supports a maximum of penultimate dose of a potent statin (e.g., atorvastatin or rosuvastatin). For those patients with the metabolic syndrome but without diabetes, a score that predicts a 10-year CVD risk exceeding 7.5% should also take a statin. With a 10-year risk of $<7.5\%$, use of statin therapy is not evidence based.

Diets restricted in saturated fats ($<7\%$ of calories) and *trans*-fats (as few as possible) should be applied aggressively. Although less evidence exists, dietary cholesterol should also be restricted. If LDL cholesterol remains elevated, pharmacologic intervention is needed. Treatment with statins, which lower LDL cholesterol by 15–60%, is evidence based and is the first-choice medication intervention. Of note, for each doubling of the statin dose, LDL cholesterol is further lowered by only $\sim 6\%$. Hepatotoxicity (more than a threefold increase in hepatic aminotransferases) is rare, and myopathy is seen in $\sim 10\%$ of patients. The cholesterol absorption inhibitor ezetimibe is well tolerated and should be the second-choice medication intervention. Ezetimibe typically reduces LDL cholesterol by 15–20%. The bile acid sequestrants cholestyramine, colestipol, and colesevalam may be more effective than ezetimibe but, because they can increase triglyceride levels, must be used with caution in patients with the metabolic syndrome. In general, bile sequestrants should not be administered when fasting triglyceride levels are >250 mg/dL. Side effects include gastrointestinal symptoms (palatability, bloating, belching, constipation, anal irritation). Nicotinic acid has modest LDL cholesterol-lowering capabilities ($<20\%$). Fibrates are best employed to lower LDL cholesterol when both LDL cholesterol and triglycerides are elevated. Fenofibrate may be more effective than gemfibrozil in this setting.

TRIGLYCERIDES (SEE ALSO CHAP. 421)

The NCEP:ATPIII has focused on non-HDL cholesterol rather than on triglycerides. However, a fasting triglyceride value of <150 mg/dL is recommended. In general, the response of fasting triglycerides relates to the amount of weight reduction achieved: a weight reduction of $>10\%$ is necessary to lower fasting triglyceride levels.

A fibrate (gemfibrozil or fenofibrate) is the drug of choice to lower fasting triglyceride levels, which are typically reduced by 30–45%. Concomitant administration with drugs metabolized by the 3A4 cytochrome P450 system (including some statins) increases the risk of myopathy. In these cases, fenofibrate may be preferable to gemfibrozil. In the Veterans Affairs HDL Intervention Trial, gemfibrozil was administered to men with known CHD and levels of HDL cholesterol <40 mg/dL. A coronary disease event and mortality rate benefit was experienced predominantly among men with hyperinsulinemia and/or diabetes, many of whom were identified retrospectively as having the metabolic syndrome. Of note, the degree of triglyceride lowering in this trial did not predict benefit. Although levels of LDL cholesterol did not change, a decrease in LDL particle number correlated with benefit. Several additional clinical trials have not shown

2454 clear evidence that fibrates reduce CVD risk; however, post hoc analyses of several studies demonstrated that patients with baseline triglyceride levels >200 mg/dL and HDL cholesterol levels <35 mg/dL did benefit.

Other drugs that lower triglyceride levels include statins, nicotinic acid, and—in high doses—omega-3 fatty acids. For this purpose, an intermediate or high dose of the “more potent” statins (atorvastatin, rosuvastatin) is needed. The effect of nicotinic acid on fasting triglycerides is dose related and ~20–35%, an effect that is less pronounced than that of fibrates. In patients with the metabolic syndrome and diabetes, nicotinic acid may increase fasting glucose levels. Omega-3 fatty acid preparations that include high doses of docosahexaenoic acid plus eicosapentaenoic acid (~1.5–4.5 g/d) or eicosapentaenoic acid alone lower fasting triglyceride levels by ~30–40%. No drug interactions with fibrates or statins occur, and the main side effect of their use is eructation with a fishy taste. This taste can be partially blocked by ingestion of the nutraceutical after freezing. Clinical trials of nicotinic acid or high-dose omega-3 fatty acids in patients with the metabolic syndrome have not been reported.

HDL CHOLESTEROL (SEE ALSO CHAP. 421)

Very few lipid-modifying compounds increase HDL cholesterol levels. Statins, fibrates, and bile acid sequestrants have modest effects (5–10%), whereas ezetimibe and omega-3 fatty acids have no effect. Nicotinic acid is the only currently available drug with predictable HDL cholesterol-raising properties. The response is dose related, and nicotinic acid can increase HDL cholesterol by ~30% above baseline. After several trials of nicotinic acid versus placebo in statin-treated patients, there is still no evidence that raising HDL with nicotinic acid beneficially affects CVD events in patients with or without the metabolic syndrome.

BLOOD PRESSURE (SEE ALSO CHAP. 298)

The direct relationship between blood pressure and all-cause mortality rate has been well established in studies comparing patients

with hypertension (>140/90 mmHg), patients with pre-hypertension (>120/80 mmHg but <140/90 mmHg), and individuals with normal blood pressure (<120/80 mmHg). In patients who have the metabolic syndrome without diabetes, the best choice for the initial antihypertensive medication is an angiotensin-converting enzyme (ACE) inhibitor or an angiotensin II receptor blocker, as these two classes of drugs appear to reduce the incidence of new-onset type 2 diabetes. In all patients with hypertension, a sodium-restricted dietary pattern enriched in fruits and vegetables, whole grains, and low-fat dairy products should be advocated. Home monitoring of blood pressure may assist in maintaining good blood-pressure control.

IMPAIRED FASTING GLUCOSE (SEE ALSO CHAP. 417)

In patients with the metabolic syndrome and type 2 diabetes, aggressive glycemic control may favorably modify fasting levels of triglycerides and/or HDL cholesterol. In patients with impaired fasting glucose who do not have diabetes, a lifestyle intervention that includes weight reduction, dietary fat restriction, and increased physical activity has been shown to reduce the incidence of type 2 diabetes. Metformin also reduces the incidence of diabetes, although the effect is less pronounced than that of lifestyle intervention.

INSULIN RESISTANCE (SEE ALSO CHAP. 418)

Several drug classes (biguanides, thiazolidinediones [TZDs]) increase insulin sensitivity. Because insulin resistance is the primary pathophysiologic mechanism for the metabolic syndrome, representative drugs in these classes reduce its prevalence. Both metformin and TZDs enhance insulin action in the liver and suppress endogenous glucose production. TZDs, but not metformin, also improve insulin-mediated glucose uptake in muscle and adipose tissue. Benefits of both drugs have been seen in patients with nonalcoholic fatty liver disease and polycystic ovary syndrome, and the drugs have been shown to reduce markers of inflammation.

SECTION 4 DISORDERS OF BONE AND MINERAL METABOLISM

423 Bone and Mineral Metabolism in Health and Disease

F. Richard Bringham, Marie B. Demay,
Stephen M. Krane, Henry M. Kronenberg

BONE STRUCTURE AND METABOLISM

Bone is a dynamic tissue that is remodeled constantly throughout life. The arrangement of compact and cancellous bone provides strength and density suitable for both mobility and protection. In addition, bone provides a reservoir for calcium, magnesium, phosphorus, sodium, and other ions necessary for homeostatic functions. Bone also hosts and regulates hematopoiesis by providing niches for hematopoietic cell proliferation and differentiation. The skeleton is highly vascular and receives about 10% of the cardiac output. Remodeling of bone is accomplished by two distinct cell types: osteoblasts produce bone matrix, and osteoclasts resorb the matrix.

The extracellular components of bone consist of a solid mineral phase in close association with an organic matrix, of which 90–95% is type I collagen (Chap. 427). The noncollagenous portion of the organic matrix is heterogeneous and contains serum proteins such as albumin as well as many locally produced proteins, whose functions are incompletely understood. Those proteins include cell attachment/

signaling proteins such as thrombospondin, osteopontin, and fibronectin; calcium-binding proteins such as matrix gla protein and osteocalcin; and proteoglycans such as biglycan and decorin. Some of the proteins organize collagen fibrils; others influence mineralization and binding of the mineral phase to the matrix.

The mineral phase is made up of calcium and phosphate and is best characterized as a poorly crystalline hydroxyapatite. The mineral phase of bone is deposited initially in intimate relation to the collagen fibrils and is found in specific locations in the “holes” between the collagen fibrils. This architectural arrangement of mineral and matrix results in a two-phase material well suited to withstand mechanical stresses. The organization of collagen influences the amount and type of mineral phase formed in bone. Although the primary structures of type I collagen in skin and bone tissues are similar, there are differences in posttranslational modifications and distribution of intermolecular cross-links. The holes in the packing structure of the collagen are larger in mineralized collagen of bone and dentin than in unmineralized collagens such as those in tendon. Single amino acid substitutions in the helical portion of either the $\alpha 1$ (COL1A1) or $\alpha 2$ (COL1A2) chains of type I collagen disrupt the organization of bone in osteogenesis imperfecta. The severe skeletal fragility associated with this group of disorders highlights the importance of the fibrillar matrix in the structure of bone (Chap. 427).

Osteoblasts synthesize and secrete the organic matrix and regulate its mineralization. They are derived from cells of mesenchymal origin (Fig. 423-1A). Active osteoblasts are found on the surface of newly

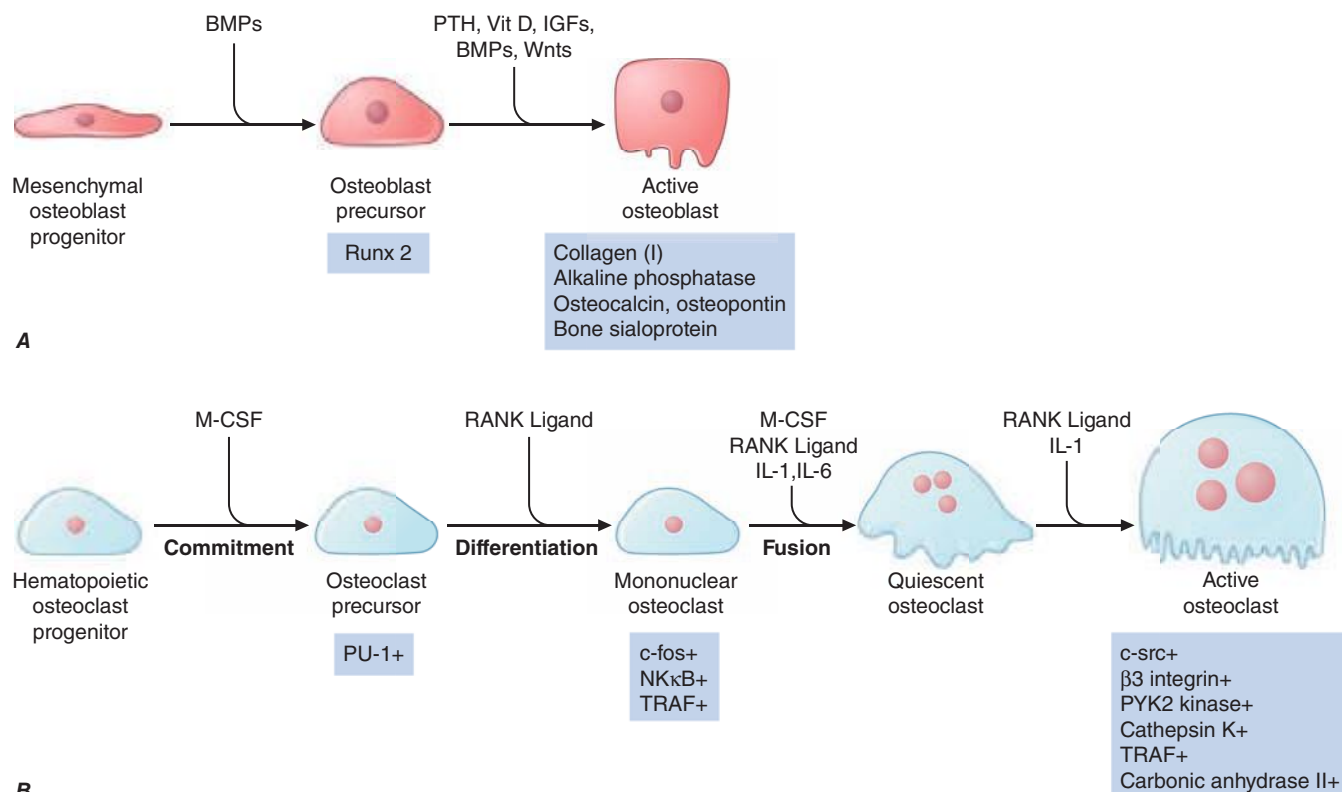


FIGURE 423-1 Pathways regulating development of (A) osteoblasts and (B) osteoclasts. Hormones, cytokines, and growth factors that control cell proliferation and differentiation are shown above the arrows. Transcription factors and other markers specific for various stages of development are depicted below the arrows. BMPs, bone morphogenic proteins; IGFs, insulin-like growth factors; IL-1, interleukin 1; IL-6, interleukin 6; M-CSF, macrophage colony-stimulating factor; NFκB, nuclear factor κB; PTH, parathyroid hormone; PU-1, a monocyte- and B lymphocyte-specific ets family transcription factor; RANK ligand, receptor activator of NFκB ligand; Runx2, Runt-related transcription factor 2; TRAF, tumor necrosis factor receptor-associated factors; Vit D, vitamin D; wnts, wingless-type mouse mammary tumor virus integration site. (Modified from T Suda et al: *Endocr Rev* 20:345, 1999, with permission.)

forming bone. As an osteoblast secretes matrix, which then is mineralized, the cell becomes an *osteocyte*, still connected with its blood supply through a series of canaliculi. Osteocytes account for the vast majority of the cells in bone. They are thought to be the mechanosensors in bone that communicate signals to surface osteoblasts and their progenitors through the canalicular network and thereby serve as master regulators of bone formation and resorption. Remarkably, osteocytes also secrete fibroblast growth factor 23 (FGF23), a major regulator of phosphate metabolism (see below). Mineralization of the matrix, both in trabecular bone and in osteons of compact cortical bone (*Haversian systems*), begins soon after the matrix is secreted (primary mineralization) but is not completed for several weeks or even longer (secondary mineralization). Although this mineralization takes advantage of the high concentrations of calcium and phosphate, already near saturation in serum, mineralization is a carefully regulated process that is dependent on the activity of osteoblast-derived alkaline phosphatase, which probably works by hydrolyzing inhibitors of mineralization.

Genetic studies in humans and mice have identified several key genes that control osteoblast development. *Runx2* is a transcription factor expressed specifically in chondrocyte (cartilage cells) and osteoblast progenitors as well as in hypertrophic chondrocytes and mature osteoblasts. *Runx2* regulates the expression of several important osteoblast proteins, including osterix (another transcription factor needed for osteoblast maturation), osteopontin, bone sialoprotein, type I collagen, osteocalcin, and receptor-activator of NFκB (RANK) ligand. *Runx2* expression is regulated in part by bone morphogenic proteins (BMPs). *Runx2*-deficient mice are devoid of osteoblasts, whereas mice with a deletion of only one allele (*Runx2* +/-) exhibit a delay in formation of the clavicles and some cranial bones. The latter abnormalities are similar to those in the human disorder *cleidocranial dysplasia*, which is also caused by heterozygous inactivating mutations in *Runx2*.

The paracrine signaling molecule, Indian hedgehog (Ihh), also plays a critical role in osteoblast development, as evidenced by *Ihh*-deficient mice that lack osteoblasts in the type of bone formed on a cartilage mold (endochondral ossification). Signals originating from members of the wnt (wingless-type mouse mammary tumor virus integration site) family of paracrine factors are also important for osteoblast proliferation and differentiation. Numerous other growth-regulatory factors affect osteoblast function, including the three closely related transforming growth factor βs, fibroblast growth factors (FGFs) 2 and 18, platelet-derived growth factor, and insulin-like growth factors (IGFs) I and II. Hormones such as parathyroid hormone (PTH) and 1,25-dihydroxyvitamin D (1,25[OH]₂D) activate receptors expressed by osteoblasts to assure mineral homeostasis and influence a variety of bone cell functions.

Resorption of bone is carried out mainly by *osteoclasts*, multinucleated cells that are formed by fusion of cells derived from the common precursor of macrophages and osteoclasts. Thus, these cells derive from the hematopoietic lineage, quite different from the mesenchymal cells that become osteoblasts. Multiple factors that regulate osteoclast development have been identified (Fig. 423-1B). Factors produced by osteoblasts or marrow stromal cells allow osteoblasts to control osteoclast development and activity. Macrophage colony-stimulating factor (M-CSF) plays a critical role during several steps in the pathway and ultimately leads to fusion of osteoclast progenitor cells to form multinucleated, active osteoclasts. RANK ligand, a member of the tumor necrosis factor (TNF) family, is expressed on the surface of osteoblast progenitors and stromal fibroblasts. In a process involving cell-cell interactions, RANK ligand binds to the RANK receptor on osteoclast progenitors, stimulating osteoclast differentiation and activation. Alternatively, a soluble decoy receptor, referred to as osteoprotegerin, can bind RANK ligand and inhibit osteoclast

2456 differentiation. Several growth factors and cytokines (including interleukins 1, 6, and 11; TNF; and interferon γ) modulate osteoclast differentiation and function. Most hormones that influence osteoclast function do not target these cells directly but instead act on cells of the osteoblast lineage to increase production of M-CSF and RANK. Both PTH and $1,25(\text{OH})_2\text{D}$ increase osteoclast number and activity by this indirect mechanism. Calcitonin, in contrast, binds to its receptor on the basal surface of osteoclasts and directly inhibits osteoclast function. Estradiol has multiple cellular targets in bone, including osteoclasts, immune cells, and osteoblasts; actions on all these cells serve to decrease osteoclast number and decrease bone resorption.

Osteoclast-mediated resorption of bone takes place in scalloped spaces (*Howship's lacunae*) where the osteoclasts are attached through a specific $\alpha_v\beta_3$ integrin to components of the bone matrix such as osteopontin. The osteoclast forms a tight seal to the underlying matrix and secretes protons, chloride, and proteinases into a confined space that has been likened to an extracellular lysosome. The active osteoclast surface forms a ruffled border that contains a specialized proton pump ATPase that secretes acid and solubilizes the mineral phase. Carbonic anhydrase (type II isoenzyme) within the osteoclast generates the needed protons. The bone matrix is resorbed in the acid environment adjacent to the ruffled border by proteases, such as cathepsin K, that act at low pH.

In the embryo and the growing child, bone develops mostly by remodeling and replacing previously calcified cartilage (endochondral bone formation) or, in a few bones, is formed without a cartilage matrix (intramembranous bone formation). During endochondral bone formation, chondrocytes proliferate, secrete and mineralize a matrix, enlarge (hypertrophy), and then die, enlarging bone and providing the matrix and factors that stimulate endochondral bone formation. This program is regulated by both local factors, such as IGF-I and -II, Ihh, PTH-related peptide (PTHrP), and FGFs, and by systemic hormones, such as growth hormone, glucocorticoids, and estrogen.

New bone, whether formed in infants or in adults during repair, has a relatively high ratio of cells to matrix and is characterized by coarse fiber bundles of collagen that are interlaced and randomly dispersed (woven bone). In adults, the more mature bone is organized with fiber bundles regularly arranged in parallel or concentric sheets (lamellar bone). In long bones, deposition of lamellar bone in a concentric arrangement around blood vessels forms the Haversian systems. Growth in length of bones is dependent on proliferation of cartilage cells and the endochondral sequence at the growth plate. Growth in width and thickness is accomplished by formation of bone at the periosteal surface and by resorption at the endosteal surface, with the rate of formation exceeding that of resorption. In adults, after the growth plates of cartilage close, growth in length and endochondral bone formation cease except for some activity in the cartilage cells beneath the articular surface. Even in adults, however, remodeling of bone (within Haversian systems as well as along the surfaces of trabecular bone) continues throughout life. In adults, ~4% of the surface of trabecular bone (such as iliac crest) is involved in active resorption, whereas 10–15% of trabecular surfaces are covered with osteoid, unmineralized new bone formed by osteoblasts. Radioisotope studies indicate that as much as 18% of the total skeletal calcium is deposited and removed each year. Thus, bone is an active metabolizing tissue that requires an intact blood supply. The cycle of bone resorption and formation is a highly orchestrated process carried out by the basic multicellular unit, which is composed of a group of osteoclasts and osteoblasts (Fig. 423-2).

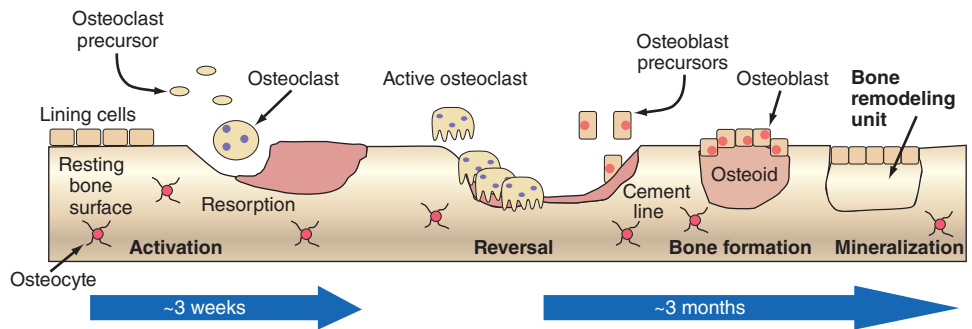


FIGURE 423-2 Schematic representation of bone remodeling. The cycle of bone remodeling is carried out by the basic multicellular unit (BMU), which consists of a group of osteoclasts and osteoblasts. In cortical bone, the BMUs tunnel through the tissue, whereas in cancellous bone, they move across the trabecular surface. The process of bone remodeling is initiated by contraction of the lining cells and the recruitment of osteoclast precursors. These precursors fuse to form multinucleated, active osteoclasts that mediate bone resorption. Osteoclasts adhere to bone and subsequently remove it by acidification and proteolytic digestion. As the BMU advances, osteoclasts leave the resorption site and osteoblasts move in to cover the excavated area and begin the process of new bone formation by secreting osteoid, which eventually is mineralized into new bone. After osteoid mineralization, osteoblasts flatten and form a layer of lining cells over new bone.

The response of bone to fractures, infection, and interruption of blood supply and to expanding lesions is relatively limited. Dead bone must be resorbed, and new bone must be formed, a process carried out in association with growth of new blood vessels into the involved area. In injuries that disrupt the organization of the tissue such as a fracture in which apposition of fragments is poor or when motion exists at the fracture site, progenitor stromal cells recapitulate the endochondral bone formation of early development and form cartilage that is replaced by bone and, variably, fibrous tissue. When there is good apposition with fixation and little motion at the fracture site, repair occurs predominantly by formation of new bone without other mediating tissue.

Remodeling of bone occurs along lines of force generated by mechanical stress. The signals from these mechanical stresses are sensed by osteocytes, which transmit signals to osteoclasts and osteoblasts or their precursors. One such signal made by osteocytes is sclerostin, an inhibitor of wnt signaling. Mechanical forces suppress sclerostin production and thus increase bone formation by osteoblasts. Expanding lesions in bone such as tumors induce resorption at the surface in contact with the tumor by producing ligands such as PTHrP that stimulate osteoclast differentiation and function. Even in a disorder as architecturally disruptive as Paget's disease, remodeling is dictated by mechanical forces. Thus, bone plasticity reflects the interaction of cells with each other and with the environment.

Measurement of the products of osteoblast and osteoclast activity can assist in the diagnosis and management of bone diseases. Osteoblast activity can be assessed by measuring serum bone-specific alkaline phosphatase. Similarly, osteocalcin, a protein secreted from osteoblasts, is made virtually only by osteoblasts. Osteoclast activity can be assessed by measurement of products of collagen degradation. Collagen molecules are covalently linked to each other in the extracellular matrix through the formation of hydroxyypyridinium cross-links (Chap. 427). After digestion by osteoclasts, these cross-linked peptides can be measured both in urine and in blood.

CALCIUM METABOLISM

Over 99% of the 1–2 kg of calcium present normally in the adult human body resides in the skeleton, where it provides mechanical stability and serves as a reservoir sometimes needed to maintain extracellular fluid (ECF) calcium concentration (Fig. 423-3). Skeletal calcium accretion first becomes significant during the third trimester of fetal life, accelerates throughout childhood and adolescence, reaches a peak in early adulthood, and gradually declines thereafter at rates that rarely

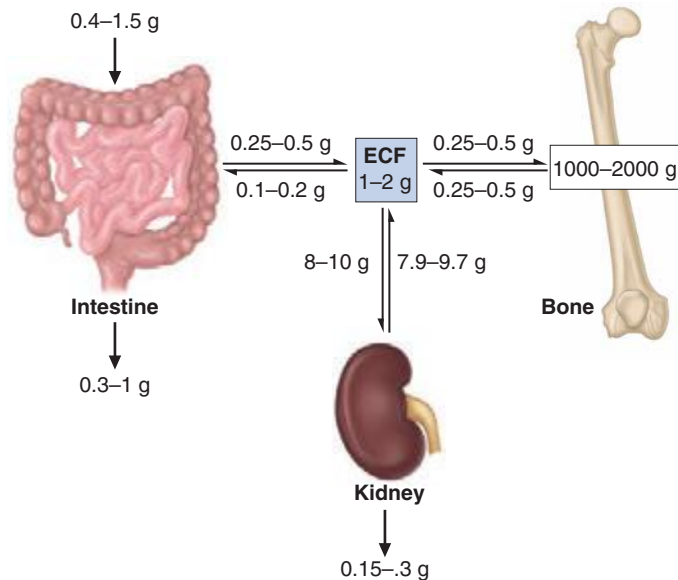


FIGURE 423-3 Calcium homeostasis. Schematic illustration of calcium content of extracellular fluid (ECF) and bone as well as of diet and feces; magnitude of calcium flux per day as calculated by various methods is shown at sites of transport in intestine, kidney, and bone. Ranges of values shown are approximate and were chosen to illustrate certain points discussed in the text. In conditions of calcium balance, rates of calcium release from and uptake into bone are equal.

exceed 1–2% per year. These slow changes in total skeletal calcium content contrast with relatively high daily rates of closely matched fluxes of calcium into and out of bone (~250–500 mg each), a process mediated by coupled osteoblastic and osteoclastic activity. Another 0.5–1% of skeletal calcium is freely exchangeable (e.g., in chemical equilibrium) with that in the ECF.

The concentration of ionized calcium in the ECF must be maintained within a narrow range because of the critical role calcium plays in a wide array of cellular functions, especially those involved in neuromuscular activity, secretion, and signal transduction. Intracellular cytosolic free calcium levels are ~100 nmol/L and are 10,000-fold lower than ionized calcium concentrations in the blood and ECF (1.1–1.3 mmol/L). Cytosolic calcium does not play the structural role played by extracellular calcium; instead, it serves a signaling function. The steep chemical gradient of calcium from outside to inside the cell promotes rapid calcium influx through various membrane calcium channels that can be activated by hormones, metabolites, or neurotransmitters, swiftly changing cellular function. In blood, total calcium concentration is normally 2.2–2.6 mM (8.5–10.5 mg/dL), of which ~50% is ionized. The remainder is bound ionically to negatively charged proteins (predominantly albumin and immunoglobulins) or loosely complexed with phosphate, citrate, sulfate, or other anions. Alterations in serum protein concentrations directly affect the total blood calcium concentration even if the ionized calcium concentration remains normal. An algorithm to correct for protein changes adjusts the total serum calcium (in mg/dL) upward by 0.8 times the deficit in serum albumin (g/dL) or by 0.5 times the deficit in serum immunoglobulin (in g/dL). Such corrections provide only rough approximations of actual free calcium concentrations, however, and may be misleading, particularly during acute illness. Acidosis also alters ionized calcium by reducing its association with proteins. The best practice is to measure blood ionized calcium directly by a method that employs calcium-selective electrodes in acute settings during which calcium abnormalities might occur.

Control of the ionized calcium concentration in the ECF ordinarily is accomplished by adjusting the rates of calcium movement across intestinal and renal epithelia. These adjustments are mediated mainly via changes in blood levels of the hormones, PTH and 1,25(OH)₂D. Blood ionized calcium directly suppresses PTH secretion by activating

calcium-sensing receptors (CaSRs) in parathyroid cells. Also, ionized calcium indirectly affects PTH secretion by lowering 1,25(OH)₂D production. This active vitamin D metabolite inhibits PTH production by an incompletely understood mechanism of negative feedback (Chap. 424).

Normal dietary calcium intake in the United States varies widely, ranging from 10–37 mmol/d (400–1500 mg/d). An Institute of Medicine report recommends a daily allowance of 25–30 mmol (1000–1200 mg) for most adults. Intestinal absorption of ingested calcium involves both active (transcellular) and passive (paracellular) mechanisms. Passive calcium absorption is nonsaturable and approximates 5% of daily calcium intake, whereas active absorption involves apical calcium entry via specific ion channels (TRPV5 and TRPV6), whose expression is controlled principally by 1,25(OH)₂D, and normally ranges from 20 to 70%. Active calcium transport occurs mainly in the proximal small bowel (duodenum and proximal jejunum), although some active calcium absorption occurs in most segments of the small intestine. Optimal rates of calcium absorption require gastric acid. This is especially true for weakly dissociable calcium supplements such as calcium carbonate. In fact, large boluses of calcium carbonate are poorly absorbed because of their neutralizing effect on gastric acid. In achlorhydric subjects and for those taking drugs that inhibit gastric acid secretion, supplements should be taken with meals to optimize their absorption. Use of calcium citrate may be preferable in these circumstances. Calcium absorption may also be blunted in disease states such as pancreatic or biliary insufficiency, in which ingested calcium remains bound to unabsorbed fatty acids or other food constituents. At high levels of calcium intake, synthesis of 1,25(OH)₂D is reduced; this decreases the rate of active intestinal calcium absorption. The opposite occurs with dietary calcium restriction. Some calcium, ~2.5–5 mmol/d (100–200 mg/d), is excreted as an obligate component of intestinal secretions and is not regulated by calciotropic hormones.

The feedback-controlled hormonal regulation of intestinal absorptive efficiency results in a relatively constant daily net calcium absorption of ~5–7.5 mmol/d (200–400 mg/d) despite large changes in daily dietary calcium intake. This daily load of absorbed calcium is excreted by the kidneys in a manner that is also tightly regulated by the concentration of ionized calcium in the blood. Approximately 8–10 g/d of calcium is filtered by the glomeruli, of which only 2–3% appears in the urine. Most filtered calcium (65%) is reabsorbed in the proximal tubules via a passive, paracellular route that is coupled to concomitant NaCl reabsorption and not specifically regulated. The cortical thick ascending limb of Henle's loop (cTAL) reabsorbs roughly another 20% of filtered calcium, also via a paracellular mechanism. Calcium reabsorption in the cTAL requires a tight-junctional protein called paracellin-1 and is inhibited by increased blood concentrations of calcium or magnesium, acting via the CaSR, which is highly expressed on basolateral membranes in this nephron segment. Operation of the renal CaSR provides a mechanism, independent of those engaged directly by PTH or 1,25(OH)₂D, by which serum ionized calcium can control renal calcium reabsorption. Finally, ~10% of filtered calcium is reabsorbed in the distal convoluted tubules (DCTs) by a transcellular mechanism. Calcium enters the luminal surface of the cell through specific apical calcium channels (TRPV5), whose number is regulated. It then moves across the cell in association with a specific calcium-binding protein (calbindin-D28k) that buffers cytosolic calcium concentrations from the large mass of transported calcium. Ca²⁺-ATPases and Na⁺/Ca²⁺ exchangers actively extrude calcium across the basolateral surface and thereby maintain the transcellular calcium gradient. All these processes are stimulated directly or indirectly by PTH. The DCT is also the site of action of thiazide diuretics, which lower urinary calcium excretion by inducing sodium depletion and thereby augmenting proximal calcium reabsorption. Conversely, dietary sodium loads, or increased distal sodium delivery caused by loop diuretics or saline infusion, induce calciuresis.

The homeostatic mechanisms that normally maintain a constant serum ionized calcium concentration may fail at extremes of calcium intake or when the hormonal systems or organs involved are compromised. Thus, even with maximal activity of the vitamin D-dependent

2458 intestinal active transport system, sustained calcium intakes <5 mmol/d (<200 mg/d) cannot provide enough net calcium absorption to replace obligate losses via the intestine, the kidney, sweat, and other secretions. In this case, increased blood levels of PTH and $1,25(\text{OH})_2\text{D}$ activate osteoclastic bone resorption to obtain needed calcium from bone, which leads to progressive bone loss and negative calcium balance. Increased PTH and $1,25(\text{OH})_2\text{D}$ also enhance renal calcium reabsorption, and $1,25(\text{OH})_2\text{D}$ enhances calcium absorption in the gut. At very high calcium intakes (>100 mmol/d [>4 g/d]), passive intestinal absorption continues to deliver calcium into the ECF despite maximally downregulated intestinal active transport and renal tubular calcium reabsorption. This can cause severe hypercalciuria, nephrocalcinosis, progressive renal failure, and hypercalcemia (e.g., “milk-alkali syndrome”). Deficiency or excess of PTH or vitamin D, intestinal disease, and renal failure represent other commonly encountered challenges to normal calcium homeostasis (**Chap. 424**).

PHOSPHORUS METABOLISM

Although 85% of the ~ 600 g of body phosphorus is present in bone mineral, phosphorus is also a major intracellular constituent both as the free anion(s) and as a component of numerous organophosphate compounds, including structural proteins, enzymes, transcription factors, carbohydrate and lipid intermediates, high-energy stores (adenosine triphosphate [ATP], creatine phosphate), and nucleic acids. Unlike calcium, phosphorus exists intracellularly at concentrations close to those present in ECF (e.g., 1–2 mmol/L). In cells and in the ECF, phosphorus exists in several forms, predominantly as H_2PO_4^- or NaHPO_4^- , with perhaps 10% as HPO_4^{2-} . This mixture of anions will be referred to here as “phosphate.” In serum, about 12% of phosphorus is bound to proteins. Concentrations of phosphates in blood and ECF generally are expressed in terms of elemental phosphorus, with the normal range in adults being 0.75–1.45 mmol/L (2.5–4.5 mg/dL). Because the volume of the intracellular fluid compartment is twice that of the ECF, measurements of ECF phosphate may not accurately reflect phosphate availability within cells that follows even modest shifts of phosphate from one compartment to the other.

Phosphate is widely available in foods and is absorbed efficiently (65%) by the small intestine even in the absence of vitamin D. However, phosphate absorptive efficiency may be enhanced (to 85–90%) via active transport mechanisms that are stimulated by $1,25(\text{OH})_2\text{D}$. These mechanisms involve activation of $\text{Na}^+/\text{PO}_4^{2-}$ co-transporters that move phosphate into intestinal cells against an unfavorable electrochemical gradient. Daily net intestinal phosphate absorption varies widely with the composition of the diet but is generally in the range of 500–1000 mg/d. Phosphate absorption can be inhibited by large doses of calcium salts or by sevelamer hydrochloride (Renagel), strategies commonly used to control levels of serum phosphate in renal failure. Aluminum hydroxide antacids also reduce phosphate absorption but are used less commonly because of the potential for aluminum toxicity. Low serum phosphate stimulates renal proximal tubular synthesis of $1,25(\text{OH})_2\text{D}$, perhaps by suppressing blood levels of FGF23 (see below).

Serum phosphate levels vary by as much as 50% on a normal day. This reflects the effect of food intake but also an underlying circadian rhythm that produces a nadir between 7:00 and 10:00 A.M. Carbohydrate administration, especially as IV dextrose solutions in fasting subjects, can decrease serum phosphate by >0.7 mmol/L (2 mg/dL) due to rapid uptake into and utilization by cells. A similar response is observed in the treatment of diabetic ketoacidosis and during metabolic or respiratory alkalosis. Because of this wide variation in serum phosphate, it is best to perform measurements in the basal, fasting state.

Control of serum phosphate is determined mainly by the rate of renal tubular reabsorption of the filtered load, which is ~ 4 –6 g/d. Because intestinal phosphate absorption is highly efficient, urinary excretion is not constant but varies directly with dietary intake. The fractional excretion of phosphate (ratio of phosphate to creatinine clearance) is generally in the range of 10–15%. The proximal tubule is the principal site at which renal phosphate reabsorption is regulated. This is accomplished by changes in the levels of apical expression and

activity of specific $\text{Na}^+/\text{PO}_4^{2-}$ co-transporters (NaPi-2a and NaPi-2c) in the proximal tubule. Levels of these transporters at the apical surface of these cells are reduced rapidly by PTH, a major hormonal regulator of renal phosphate excretion. FGF23 can impair phosphate reabsorption dramatically by a similar mechanism. Activating FGF23 mutations cause the rare disorder autosomal dominant hypophosphatemic rickets. In contrast to PTH, FGF23 also leads to reduced synthesis of $1,25(\text{OH})_2\text{D}$, which may worsen the resulting hypophosphatemia by lowering intestinal phosphate absorption. Renal reabsorption of phosphate is responsive to changes in dietary intake such that experimental dietary phosphate restriction leads to a dramatic lowering of urinary phosphate within hours, preceding any decline in serum phosphate (e.g., filtered load). This physiologic renal adaptation to changes in dietary phosphate availability occurs independently of PTH and may be mediated in part by changes in levels of serum FGF23. Findings in FGF23-knockout mice suggest that FGF23 normally acts to lower blood phosphate and $1,25(\text{OH})_2\text{D}$ levels. In turn, elevation of blood phosphate increases blood levels of FGF23.

Renal phosphate reabsorption is impaired by hypocalcemia, hypomagnesemia, and severe hypophosphatemia. Phosphate clearance is enhanced by ECF volume expansion and impaired by dehydration. Phosphate retention is an important pathophysiologic feature of renal insufficiency (**Chap. 335**).

HYPOPHOSPHATEMIA

Causes Hypophosphatemia can occur by one or more of three primary mechanisms: (1) inadequate intestinal phosphate absorption, (2) excessive renal phosphate excretion, and (3) rapid redistribution of phosphate from the ECF into bone or soft tissue (**Table 423-1**). Because phosphate is so abundant in foods, inadequate intestinal absorption is almost never observed now that aluminum hydroxide antacids, which bind phosphate in the gut, are no longer widely used. Fasting or starvation, however, may result in depletion of body phosphate and predispose to subsequent hypophosphatemia during refeeding, especially if this is accomplished with IV glucose alone.

Chronic hypophosphatemia usually signifies a persistent renal tubular phosphate-wasting disorder. Excessive activation of PTH/PTHrP receptors in the proximal tubule as a result of primary or secondary hyperparathyroidism or because of the PTHrP-mediated hypercalcemia syndrome in malignancy (**Chap. 424**) is among the more common causes of renal hypophosphatemia, especially because of the high prevalence of vitamin D deficiency in older Americans. Familial hypocalciuric hypercalcemia and Jansen’s chondrodystrophy are rare examples of genetic disorders in this category (**Chap. 424**).

Several genetic and acquired diseases cause PTH/PTHrP-independent tubular phosphate wasting with associated rickets and osteomalacia. All these diseases manifest severe hypophosphatemia; renal phosphate wasting, sometimes accompanied by aminoaciduria; inappropriately low blood levels of $1,25(\text{OH})_2\text{D}$; low-normal serum levels of calcium; and evidence of impaired cartilage or bone mineralization. Analysis of these diseases led to the discovery of the hormone FGF23, which is an important physiologic regulator of phosphate metabolism. FGF23 decreases phosphate reabsorption in the proximal tubule and also suppresses the 1α -hydroxylase responsible for synthesis of $1,25(\text{OH})_2\text{D}$. FGF23 is synthesized by cells of the osteoblast lineage, primarily osteocytes. High-phosphate diets increase FGF23 levels, and low-phosphate diets decrease them. Autosomal dominant hypophosphatemic rickets (ADHR) was the first disease linked to abnormalities in FGF23. ADHR results from activating mutations in the gene that encodes FGF23. These mutations alter a cleavage site that ordinarily allows for inactivation of intact FGF23. Several other genetic disorders exhibit elevated FGF23 and hypophosphatemia. The most common of these is X-linked hypophosphatemic rickets (XLH), which results from inactivating mutations in an endopeptidase termed *PHEX* (phosphate-regulating gene with homologies to endopeptidases on the X chromosome) that is expressed most abundantly on the surface of osteocytes and mature osteoblasts. Patients with XLH usually have high FGF23 levels, and ablation of the FGF23 gene reverses the hypophosphatemia found in the mouse version of XLH. How inactivation

TABLE 423-1 CAUSES OF HYPOPHOSPHATEMIA

- I. Reduced renal tubular phosphate reabsorption
 - A. PTH/PTHrP-dependent
 1. Primary hyperparathyroidism
 2. Secondary hyperparathyroidism
 - a. Vitamin D deficiency/resistance
 - b. Calcium starvation/malabsorption
 - c. Bartter's syndrome
 - d. Autosomal recessive renal hypercalciuria with hypomagnesemia
 3. PTHrP-dependent hypercalcemia of malignancy
 4. Familial hypocalciuric hypercalcemia
 - B. PTH/PTHrP-independent
 1. Excess FGF23 or other "phosphatonins"
 - a. X-linked hypophosphatemic rickets (XLH)
 - b. Autosomal recessive hypophosphatemia (ARHP)
 - c. Autosomal dominant hypophosphatemic rickets (ADHR) (DMP1, ENPP1 deficiency)
 - d. Tumor-induced osteomalacia syndrome (TIO)
 - e. McCune-Albright syndrome (fibrous dysplasia)
 - f. Epidermal nevus syndrome
 2. Intrinsic renal disease
 - a. Fanconi's syndrome(s)
 - b. Cystinosis
 - c. Wilson's disease
 - d. NaPi-2a or NaPi-2c mutations
 3. Other systemic disorders
 - a. Poorly controlled diabetes mellitus
 - b. Alcoholism
 - c. Hyperaldosteronism
 - d. Hypomagnesemia
 - e. Amyloidosis
 - f. Hemolytic-uremic syndrome
 - g. Renal transplantation or partial liver resection
 - h. Rewarming or induced hyperthermia
 4. Drugs or toxins
 - a. Ethanol
 - b. Acetazolamide, other diuretics
 - c. High-dose estrogens or glucocorticoids
 - d. Heavy metals (lead, cadmium, saccharated ferric oxide)
 - e. Toluene, *N*-methyl formamide
 - f. Cisplatin, ifosfamide, foscarnet, rapamycin
- II. Impaired intestinal phosphate absorption
 - A. Aluminum-containing antacids
 - B. Sevelamer
- III. Shifts of extracellular phosphate into cells
 - A. Intravenous glucose
 - B. Insulin therapy for prolonged hyperglycemia or diabetic ketoacidosis
 - C. Catecholamines (epinephrine, dopamine, albuterol)
 - D. Acute respiratory alkalosis
 - E. Gram-negative sepsis, toxic shock syndrome
 - F. Recovery from starvation or acidosis
 - G. Rapid cellular proliferation
 1. Leukemic blast crisis
 2. Intensive erythropoietin, other growth factor therapy
- IV. Accelerated net bone formation
 - A. After parathyroidectomy
 - B. Treatment of vitamin D deficiency, Paget's disease
 - C. Osteoblastic metastases

Abbreviations: PTH, parathyroid hormone; PTHrP, parathyroid hormone–related peptide.

of *PHEX* leads to increased levels of FGF23 has not been determined. Two rare autosomal recessive hypophosphatemic syndromes associated with elevated FGF23 are due to inactivating mutations of dentin matrix protein-1 (*DMP1*) and ectonucleotide pyrophosphatase/phosphodiesterase 1 (*ENPP1*), both of which normally are highly expressed in bone and regulate FGF23 production. An unusual hypophosphatemic disorder, tumor-induced osteomalacia (TIO), is an acquired disorder in which tumors, usually of mesenchymal origin and generally histologically benign, secrete FGF23 and/or other molecules that induce renal phosphate wasting. The hypophosphatemic syndrome resolves completely within hours to days after successful resection of the responsible tumor. Such tumors typically express large amounts of FGF23 mRNA, and patients with TIO usually exhibit elevations of FGF23 in their blood.

Dent's disease is an X-linked recessive disorder caused by inactivating mutations in *CLCN5*, a chloride transporter expressed in endosomes of the proximal tubule; features include hypercalciuria, hypophosphatemia, and recurrent kidney stones. Renal phosphate wasting is common among poorly controlled diabetic patients and alcoholics, who therefore are at risk for iatrogenic hypophosphatemia when treated with insulin or IV glucose, respectively. Diuretics and certain other drugs and toxins can cause defective renal tubular phosphate reabsorption (Table 423-1).

In hospitalized patients, hypophosphatemia is often attributable to massive redistribution of phosphate from the ECF into cells. Insulin therapy for diabetic ketoacidosis is a paradigm for this phenomenon, in which the severity of the hypophosphatemia is related to the extent of antecedent depletion of phosphate and other electrolytes (Chap. 417). The hypophosphatemia is usually greatest at a point many hours after initiation of insulin therapy and is difficult to predict from baseline measurements of serum phosphate at the time of presentation, when prerenal azotemia can obscure significant phosphate depletion. Other factors that may contribute to such acute redistributive hypophosphatemia include antecedent starvation or malnutrition, administration of IV glucose without other nutrients, elevated blood catecholamines (endogenous or exogenous), respiratory alkalosis, and recovery from metabolic acidosis.

Hypophosphatemia also can occur transiently (over weeks to months) during the phase of accelerated net bone formation that follows parathyroidectomy for severe primary hyperparathyroidism or during treatment of vitamin D deficiency or lytic Paget's disease. This is usually most prominent in patients who preoperatively have evidence of high bone turnover (e.g., high serum levels of alkaline phosphatase). Osteoblastic metastases can also lead to this syndrome.

Clinical and Laboratory Findings The clinical manifestations of severe hypophosphatemia reflect a generalized defect in cellular energy metabolism because of ATP depletion, a shift from oxidative phosphorylation toward glycolysis, and associated tissue or organ dysfunction. Acute, severe hypophosphatemia occurs mainly or exclusively in hospitalized patients with underlying serious medical or surgical illness and preexisting phosphate depletion due to excessive urinary losses, severe malabsorption, or malnutrition. Chronic hypophosphatemia tends to be less severe, with a clinical presentation dominated by musculoskeletal complaints such as bone pain, osteomalacia, pseudo-fractures, and proximal muscle weakness or, in children, rickets and short stature.

Neuromuscular manifestations of severe hypophosphatemia are variable but may include muscle weakness, lethargy, confusion, disorientation, hallucinations, dysarthria, dysphagia, oculomotor palsies, anisocoria, nystagmus, ataxia, cerebellar tremor, ballismus, hyporeflexia, impaired sphincter control, distal sensory deficits, paresthesia, hyperesthesia, generalized or Guillain-Barré–like ascending paralysis, seizures, coma, and even death. Serious sequelae such as paralysis, confusion, and seizures are likely only at phosphate concentrations <0.25 mmol/L (<0.8 mg/dL). Rhabdomyolysis may develop during rapidly progressive hypophosphatemia. The diagnosis of hypophosphatemia-induced rhabdomyolysis may be overlooked, as up to 30% of patients with acute hypophosphatemia (<0.7 mM) have creatine phosphokinase

2460 elevations that peak 1–2 days after the nadir in serum phosphate, when the release of phosphate from injured myocytes may have led to a near normalization of circulating levels of phosphate.

Respiratory failure and cardiac dysfunction, which are reversible with phosphate treatment, may occur at serum phosphate levels of 0.5–0.8 mmol/L (1.5–2.5 mg/dL). Renal tubular defects, including tubular acidosis, glycosuria, and impaired reabsorption of sodium and calcium, may occur. Hematologic abnormalities correlate with reductions in intracellular ATP and 2,3-diphosphoglycerate and may include erythrocyte microspherocytosis and hemolysis; impaired oxyhemoglobin dissociation; defective leukocyte chemotaxis, phagocytosis, and bacterial killing; and platelet dysfunction with spontaneous gastrointestinal hemorrhage.

TREATMENT HYPOPHOSPHATEMIA

Severe hypophosphatemia (<0.75 mmol/L [<2 mg/dL]), particularly in the setting of underlying phosphate depletion, constitutes a dangerous electrolyte abnormality that should be corrected promptly. Unfortunately, the cumulative deficit in body phosphate cannot be predicted easily from knowledge of the circulating level of phosphate, and therapy must be approached empirically. The threshold for IV phosphate therapy and the dose administered should reflect consideration of renal function, the likely severity and duration of the underlying phosphate depletion, and the presence and severity of symptoms consistent with those of hypophosphatemia. In adults, phosphate may be safely administered IV as neutral mixtures of sodium or potassium phosphate salts at initial doses of 0.2–0.8 mmol/kg of elemental phosphorus over 6 h (e.g., 10–50 mmol over 6 h), with doses >20 mmol/6 h reserved for those who have serum levels <0.5 mmol/L (1.5 mg/dL) and normal renal function. A suggested approach is presented in [Table 423-2](#). Serum levels of phosphate and calcium must be monitored closely (every 6–12 h) throughout treatment. It is necessary to avoid a serum calcium-phosphorus product >50 to reduce the risk of heterotopic calcification. Hypocalcemia, if present, should be corrected before administering IV phosphate. Less severe hypophosphatemia, in the range of 0.5–0.8 mmol/L (1.5–2.5 mg/dL), usually can be treated with oral phosphate in divided doses of 750–2000 mg/d as elemental phosphorus; higher doses can cause bloating and diarrhea.

Management of chronic hypophosphatemia requires knowledge of the cause(s) of the disorder. Hypophosphatemia related to the secondary hyperparathyroidism of vitamin D deficiency usually responds to treatment with vitamin D and calcium alone. XLH, ADHR, TIO, and related renal tubular disorders usually are managed with divided oral doses of phosphate, often with calcium and 1,25(OH)₂D supplements to bypass the block in renal 1,25(OH)₂D synthesis and prevent secondary hyperparathyroidism caused by suppression of ECF calcium levels. Thiazide diuretics may be used to prevent nephrocalcinosis in patients who are managed this way. Complete normalization of hypophosphatemia is generally not

possible in these conditions. Optimal therapy for TIO is extirpation of the responsible tumor, which may be localized by radiographic skeletal survey or bone scan (many are located in bone) or by radionuclide scanning using sestamibi or labeled octreotide. Successful treatment of TIO-induced hypophosphatemia with octreotide has been reported in a small number of patients.

HYPERPHOSPHATEMIA

Causes When the filtered load of phosphate and glomerular filtration rate (GFR) are normal, control of serum phosphate levels is achieved by adjusting the rate at which phosphate is reabsorbed by the proximal tubular NaPi-2 co-transporters. The principal hormonal regulators of NaPi-2 activity are PTH and FGF23. Hyperphosphatemia, defined in adults as a fasting serum phosphate concentration >1.8 mmol/L (5.5 mg/dL), usually results from impaired glomerular filtration, hypoparathyroidism, excessive delivery of phosphate into the ECF (from bone, gut, or parenteral phosphate therapy), or a combination of these factors ([Table 423-3](#)). The upper limit of normal serum phosphate concentrations is higher in children and neonates (2.4 mmol/L [7 mg/dL]). It is useful to distinguish hyperphosphatemia caused by impaired renal phosphate excretion from that which results from excessive delivery of phosphate into the ECF ([Table 423-3](#)).

In chronic renal insufficiency, reduced GFR leads to phosphate retention. Hyperphosphatemia in turn further impairs renal synthesis of 1,25(OH)₂D, increases FGF23 levels, and stimulates PTH secretion and hypertrophy both directly and indirectly (by lowering blood ionized calcium levels). Thus, hyperphosphatemia is a major cause of the secondary hyperparathyroidism of renal failure and must be addressed early in the course of the disease ([Chaps. 335 and 424](#)).

Hypoparathyroidism leads to hyperphosphatemia via increased expression of NaPi-2 co-transporters in the proximal tubule. Hypoparathyroidism, or parathyroid suppression, has multiple potential causes, including autoimmune disease; developmental, surgical, or radiation-induced absence of functional parathyroid tissue; vitamin D intoxication or other causes of PTH-independent hypercalcemia; cellular PTH resistance (pseudohypoparathyroidism or hypomagnesemia); infiltrative disorders such as Wilson's disease and hemochromatosis; and impaired PTH secretion caused by hypermagnesemia, severe hypomagnesemia, or activating mutations in the CaSR. Hypocalcemia may also contribute directly to impaired phosphate clearance, as calcium infusion can induce phosphaturia in hypoparathyroid subjects. Increased tubular phosphate reabsorption also occurs in acromegaly, during heparin administration, and in tumoral calcinosis. Tumoral calcinosis is caused by a rare group of genetic disorders in which FGF23 is processed in a way that leads to low levels of active FGF23 in the bloodstream. This may result from mutations in the FGF23 sequence or via inactivating mutations in the *GALNT3* gene, which encodes a galactosaminyl transferase that normally adds sugar residues to FGF23 that slow its proteolysis. A similar syndrome results from FGF23 resistance due to inactivating mutations of the FGF23 co-receptor Klotho. These abnormalities cause elevated serum 1,25(OH)₂D, parathyroid

TABLE 423-2 INTRAVENOUS THERAPY FOR HYPOPHOSPHATEMIA

Consider			
Likely severity of underlying phosphate depletion			
Concurrent parenteral glucose administration			
Presence of neuromuscular, cardiopulmonary, or hematologic complications of hypophosphatemia			
Renal function (reduce dose by 50% if serum creatinine >220 μ mol/L [>2.5 mg/dL])			
Serum calcium level (correct hypocalcemia first; reduce dose by 50% in hypercalcemia)			
Guidelines			
Serum Phosphorus, mM (mg/dL)	Rate of Infusion, mmol/h	Duration, h	Total Administered, mmol
<0.8 (<2.5)	2	6	12
<0.5 (<1.5)	4	6	24
<0.3 (<1)	8	6	48

Note: Rates shown are calculated for a 70-kg person; levels of serum calcium and phosphorus must be measured every 6–12 h during therapy; infusions can be repeated to achieve stable serum phosphorus levels >0.8 mmol/L (>2.5 mg/dL); most formulations available in the United States provide 3 mmol/mL of sodium or potassium phosphate.

TABLE 423-3 CAUSES OF HYPERPHOSPHATEMIA

- I. Impaired renal phosphate excretion
 - A. Renal insufficiency
 - B. Hypoparathyroidism
 1. Developmental
 2. Autoimmune
 3. After neck surgery or radiation
 4. Activating mutations of the calcium-sensing receptor
 - C. Parathyroid suppression
 1. Parathyroid-independent hypercalcemia
 - a. Vitamin D or vitamin A intoxication
 - b. Sarcoidosis, other granulomatous diseases
 - c. Immobilization, osteolytic metastases
 - d. Milk-alkali syndrome
 2. Severe hypermagnesemia or hypomagnesemia
 - D. Pseudohypoparathyroidism
 - E. Acromegaly
 - F. Tumoral calcinosis
 - G. Heparin therapy
- II. Massive extracellular fluid phosphate loads
 - A. Rapid administration of exogenous phosphate (intravenous, oral, rectal)
 - B. Extensive cellular injury or necrosis
 1. Crush injuries
 2. Rhabdomyolysis
 3. Hyperthermia
 4. Fulminant hepatitis
 5. Cytotoxic therapy
 6. Severe hemolytic anemia
 - C. Transcellular phosphate shifts
 1. Metabolic acidosis
 2. Respiratory acidosis

suppression, increased intestinal calcium absorption, and focal hyperostosis with large, lobulated periarticular heterotopic ossifications (especially at shoulders or hips) and are accompanied by hyperphosphatemia. In some forms of tumoral calcinosis, serum phosphorus levels are normal.

When large amounts of phosphate are delivered rapidly into the ECF, hyperphosphatemia can occur despite normal renal function. Examples include overzealous IV phosphate therapy, oral or rectal administration of large amounts of phosphate-containing laxatives or enemas (especially in children), extensive soft tissue injury or necrosis (crush injuries, rhabdomyolysis, hyperthermia, fulminant hepatitis, cytotoxic chemotherapy), extensive hemolytic anemia, and transcellular phosphate shifts induced by severe metabolic or respiratory acidosis.

Clinical Findings The clinical consequences of acute, severe hyperphosphatemia are due mainly to the formation of widespread calcium phosphate precipitates and resulting hypocalcemia. Thus, tetany, seizures, accelerated nephrocalcinosis (with renal failure, hyperkalemia, hyperuricemia, and metabolic acidosis), and pulmonary or cardiac calcifications (including development of acute heart block) may occur. The severity of these complications relates to the elevation of serum phosphate levels, which can reach concentrations as high as 7 mmol/L (20 mg/dL) in instances of massive soft tissue injury or tumor lysis syndrome.

TREATMENT HYPERPHOSPHATEMIA

Therapeutic options for management of severe hyperphosphatemia are limited. Volume expansion may enhance renal phosphate clearance. Aluminum hydroxide antacids or sevelamer may be helpful in chelating and limiting absorption of offending phosphate salts

present in the intestine. Hemodialysis is the most effective therapeutic strategy and should be considered early in the course of severe hyperphosphatemia, especially in the setting of renal failure and symptomatic hypocalcemia.

MAGNESIUM METABOLISM

Magnesium is the major intracellular divalent cation. Normal concentrations of extracellular magnesium and calcium are crucial for normal neuromuscular activity. Intracellular magnesium forms a key complex with ATP and is an important cofactor for a wide range of enzymes, transporters, and nucleic acids required for normal cellular function, replication, and energy metabolism. The concentration of magnesium in serum is closely regulated within the range of 0.7–1 mmol/L (1.5–2 meq/L; 1.7–2.4 mg/dL), of which 30% is protein-bound and another 15% is loosely complexed to phosphate and other anions. One-half of the 25 g (1000 mmol) of total body magnesium is located in bone, only one-half of which is insoluble in the mineral phase. Almost all extracellular magnesium is present within cells, where the total concentration is 5 mM, 95% of which is bound to proteins and other macromolecules. Because only 1% of body magnesium resides in the ECF, measurements of serum magnesium levels may not accurately reflect the level of total body magnesium stores.

Dietary magnesium content normally ranges from 6 to 15 mmol/d (140–360 mg/d), of which 30–40% is absorbed, mainly in the jejunum and ileum. Intestinal magnesium absorptive efficiency is stimulated by 1,25(OH)₂D and can reach 70% during magnesium deprivation. Urinary magnesium excretion normally matches net intestinal absorption and is ~4 mmol/d (100 mg/d). Regulation of serum magnesium concentrations is achieved mainly by control of renal magnesium reabsorption. Only 20% of filtered magnesium is reabsorbed in the proximal tubule, whereas 60% is reclaimed in the cTAL and another 5–10% in the DCT. Magnesium reabsorption in the cTAL occurs via a paracellular route that requires both a lumen-positive potential, created by NaCl reabsorption, and tight-junction proteins encoded by members of the Claudin gene family. Magnesium reabsorption in the cTAL is increased by PTH but inhibited by hypercalcemia or hypermagnesemia, both of which activate the CaSR in this nephron segment.

HYPOMAGNESEMIA

Causes Hypomagnesemia usually signifies substantial depletion of body magnesium stores (0.5–1 mmol/kg). Hypomagnesemia can result from intestinal malabsorption; protracted vomiting, diarrhea, or intestinal drainage; defective renal tubular magnesium reabsorption; or rapid shifts of magnesium from the ECF into cells, bone, or third spaces (Table 423-4). Dietary magnesium deficiency is unlikely except possibly in the setting of alcoholism. A rare genetic disorder that causes selective intestinal magnesium malabsorption has been described (primary infantile hypomagnesemia). Another rare inherited disorder (hypomagnesemia with secondary hypocalcemia) is caused by mutations in the gene encoding TRPM6, a protein that, along with TRPM7, forms a channel important for both intestinal and distal-tubular renal transcellular magnesium transport. Malabsorptive states, often compounded by vitamin D deficiency, can critically limit magnesium absorption and produce hypomagnesemia despite the compensatory effects of secondary hyperparathyroidism and of hypocalcemia and hypomagnesemia to enhance cTAL magnesium reabsorption. Diarrhea or surgical drainage fluid may contain ≥5 mmol/L of magnesium. Proton pump inhibitors (omeprazole and others) may produce hypomagnesemia by an unknown mechanism that does not involve renal wasting of magnesium.

Several genetic magnesium-wasting syndromes have been described, including inactivating mutations of genes encoding the DCT NaCl co-transporter (Gitelman's syndrome), proteins required for cTAL Na-K-2Cl transport (Bartter's syndrome), claudin 16 or claudin 19 (autosomal recessive renal hypomagnesemia with hypercalciuria), a DCT Na⁺,K⁺-ATPase γ -subunit (autosomal dominant renal hypomagnesemia with hypercalciuria), DCT K⁺ channels (Kv1.1, Kir4.1), and a mitochondrial gene encoding a tRNA. Activating mutations

- I. Impaired intestinal absorption
 - A. Hypomagnesemia with secondary hypocalcemia (TRPM6 mutations)
 - B. Malabsorption syndromes
 - C. Vitamin D deficiency
 - D. Proton pump inhibitors
- II. Increased intestinal losses
 - A. Protracted vomiting/diarrhea
 - B. Intestinal drainage, fistulas
- III. Impaired renal tubular reabsorption
 - A. Genetic magnesium-wasting syndromes
 1. Gitelman's syndrome
 2. Bartter's syndrome
 3. Claudin 16 or 19 mutations
 4. Potassium channel mutations (Kv1.1, Kir4.1)
 5. Na⁺,K⁺-ATPase γ -subunit mutations (FXD2)
 - B. Acquired renal disease
 1. Tubulointerstitial disease
 2. Postobstruction, ATN (diuretic phase)
 3. Renal transplantation
 - C. Drugs and toxins
 1. Ethanol
 2. Diuretics (loop, thiazide, osmotic)
 3. Cisplatin
 4. Pentamidine, foscarnet
 5. Cyclosporine
 6. Aminoglycosides, amphotericin B
 7. Cetuximab
 - D. Other
 1. Extracellular fluid volume expansion
 2. Hyperaldosteronism
 3. SIADH
 4. Diabetes mellitus
 5. Hypercalcemia
 6. Phosphate depletion
 7. Metabolic acidosis
 8. Hyperthyroidism
- IV. Rapid shifts from extracellular fluid
 - A. Intracellular redistribution
 1. Recovery from diabetic ketoacidosis
 2. Refeeding syndrome
 3. Correction of respiratory acidosis
 4. Catecholamines
 - B. Accelerated bone formation
 1. Postparathyroidectomy
 2. Treatment of vitamin D deficiency
 3. Osteoblastic metastases
 - C. Other
 1. Pancreatitis, burns, excessive sweating
 2. Pregnancy (third trimester) and lactation

Abbreviations: ATN, acute tubular necrosis; SIADH, syndrome of inappropriate antidiuretic hormone.

of the CaSR can cause hypomagnesemia as well as hypocalcemia. ECF expansion, hypercalcemia, and severe phosphate depletion may impair magnesium reabsorption, as can various forms of renal injury, including those caused by drugs such as cisplatin, cyclosporine, aminoglycosides, and pentamidine as well as the epidermal growth factor (EGF) receptor inhibitory antibody, cetuximab (EGF action is required for normal DCT apical expression of TRPM6) (Table 423-4). A rising blood concentration of ethanol directly impairs tubular magnesium

reabsorption, and persistent glycosuria with osmotic diuresis leads to magnesium wasting and probably contributes to the high frequency of hypomagnesemia in poorly controlled diabetic patients. Magnesium depletion is aggravated by metabolic acidosis, which causes intracellular losses as well.

Hypomagnesemia due to rapid shifts of magnesium from ECF into the intracellular compartment can occur during recovery from diabetic ketoacidosis, starvation, or respiratory acidosis. Less acute shifts may be seen during rapid bone formation after parathyroidectomy, with treatment of vitamin D deficiency, or with osteoblastic metastases. Large amounts of magnesium may be lost with acute pancreatitis, extensive burns, or protracted and severe sweating and during pregnancy and lactation.

Clinical and Laboratory Findings Hypomagnesemia may cause generalized alterations in neuromuscular function, including tetany, tremor, seizures, muscle weakness, ataxia, nystagmus, vertigo, apathy, depression, irritability, delirium, and psychosis. Patients are usually asymptomatic when serum magnesium concentrations are >0.5 mmol/L (1 meq/L; 1.2 mg/dL), although the severity of symptoms may not correlate with serum magnesium levels. Cardiac arrhythmias may occur, including sinus tachycardia, other supraventricular tachycardias, and ventricular arrhythmias. Electrocardiographic abnormalities may include prolonged PR or QT intervals, T-wave flattening or inversion, and ST straightening. Sensitivity to digitalis toxicity may be enhanced.

Other electrolyte abnormalities often seen with hypomagnesemia, including hypocalcemia (with hypocalciuria) and hypokalemia, may not be easily corrected unless magnesium is administered as well. The hypocalcemia may be a result of concurrent vitamin D deficiency, although hypomagnesemia can cause impaired synthesis of 1,25(OH)₂D, cellular resistance to PTH, and, at very low serum magnesium (<0.4 mmol/L [0.8 meq/L; <1 mg/dL]), a defect in PTH secretion; these abnormalities are reversible with therapy.

TREATMENT HYPOMAGNESEMIA

Mild, asymptomatic hypomagnesemia may be treated with oral magnesium salts (MgCl₂, MgO, Mg(OH)₂) in divided doses totaling 20–30 mmol/d (40–60 meq/d). Diarrhea may occur with larger doses. More severe hypomagnesemia should be treated parenterally, preferably with IV MgCl₂, which can be administered safely as a continuous infusion of 50 mmol/d (100 meq Mg²⁺/d) if renal function is normal. If GFR is reduced, the infusion rate should be lowered by 50–75%. Use of IM MgSO₄ is discouraged; the injections are painful and provide relatively little magnesium (2 mL of 50% MgSO₄ supplies only 4 mmol). MgSO₄ may be given IV instead of MgCl₂, although the sulfate anions may bind calcium in serum and urine and aggravate hypocalcemia. Serum magnesium should be monitored at intervals of 12–24 h during therapy, which may continue for several days because of impaired renal conservation of magnesium (only 50–70% of the daily IV magnesium dose is retained) and delayed repletion of intracellular deficits, which may be as high as 1–1.5 mmol/kg (2–3 meq/kg).

It is important to consider the need for calcium, potassium, and phosphate supplementation in patients with hypomagnesemia. Vitamin D deficiency frequently coexists and should be treated with oral or parenteral vitamin D or 25(OH)D (but not with 1,25(OH)₂D, which may impair tubular magnesium reabsorption, possibly via PTH suppression). In severely hypomagnesemic patients with concomitant hypocalcemia and hypophosphatemia, administration of IV magnesium alone may worsen hypophosphatemia, provoking neuromuscular symptoms or rhabdomyolysis, due to rapid stimulation of PTH secretion. This is avoided by administering both calcium and magnesium.

HYPERMAGNESEMIA

Causes Hypermagnesemia is rarely seen in the absence of renal insufficiency, as normal kidneys can excrete large amounts (250 mmol/d) of

TABLE 423-5 CAUSES OF HYPERMAGNESEMIA

- I. Excessive magnesium intake
 - A. Cathartics, urologic irrigants
 - B. Parenteral magnesium administration
- II. Rapid mobilization from soft tissues
 - A. Trauma, shock, sepsis
 - B. Cardiac arrest
 - C. Burns
- III. Impaired magnesium excretion
 - A. Renal failure
 - B. Familial hypocalciuric hypercalcemia
- IV. Other
 - A. Adrenal insufficiency
 - B. Hypothyroidism
 - C. Hypothermia

magnesium. Mild hypermagnesemia due to excessive reabsorption in the cTAL occurs with CaSR mutations in familial hypocalciuric hypercalcemia and has been described in some patients with adrenal insufficiency, hypothyroidism, or hypothermia. Massive exogenous magnesium exposures, usually via the gastrointestinal tract, can overwhelm renal excretory capacity and cause life-threatening hypermagnesemia (Table 423-5). A notable example of this is prolonged retention of even normal amounts of magnesium-containing cathartics in patients with intestinal ileus, obstruction, or perforation. Extensive soft tissue injury or necrosis can also deliver large amounts of magnesium into the ECF in patients who have suffered trauma, shock, sepsis, cardiac arrest, or severe burns.

Clinical and Laboratory Findings The most prominent clinical manifestations of hypermagnesemia are vasodilation and neuromuscular blockade, which may appear at serum magnesium concentrations >2 mmol/L (>4.8 meq/L; >4.8 mg/dL). Hypotension that is refractory to vasopressors or volume expansion may be an early sign. Nausea, lethargy, and weakness may progress to respiratory failure, paralysis, and coma, with hypoactive tendon reflexes, at serum magnesium levels >4 mmol/L. Other findings may include gastrointestinal hypomotility or ileus; facial flushing; pupillary dilation; paradoxical bradycardia; prolongation of PR, QRS, and QT intervals; heart block; and, at serum magnesium levels approaching 10 mmol/L, asystole.

Hypermagnesemia, acting via the CaSR, causes hypocalcemia and hypercalciuria due to both parathyroid suppression and impaired cTAL calcium reabsorption.

TREATMENT HYPERMAGNESEMIA

Successful treatment of hypermagnesemia generally involves identifying and interrupting the source of magnesium and employing measures to increase magnesium clearance from the ECF. Use of magnesium-free cathartics or enemas may be helpful in clearing ingested magnesium from the gastrointestinal tract. Vigorous IV hydration should be attempted, if appropriate. Hemodialysis is effective and may be required in patients with significant renal insufficiency. Calcium, administered IV in doses of 100–200 mg over 1–2 h, has been reported to provide temporary improvement in signs and symptoms of hypermagnesemia.

VITAMIN D

SYNTHESIS AND METABOLISM

1,25-Dihydroxyvitamin D ($1,25[\text{OH}]_2\text{D}$) is the major steroid hormone involved in mineral ion homeostasis regulation. Vitamin D and its metabolites are hormones and hormone precursors rather than vitamins, since in the proper biologic setting, they can be synthesized endogenously (Fig. 423-4). In response to ultraviolet radiation of the

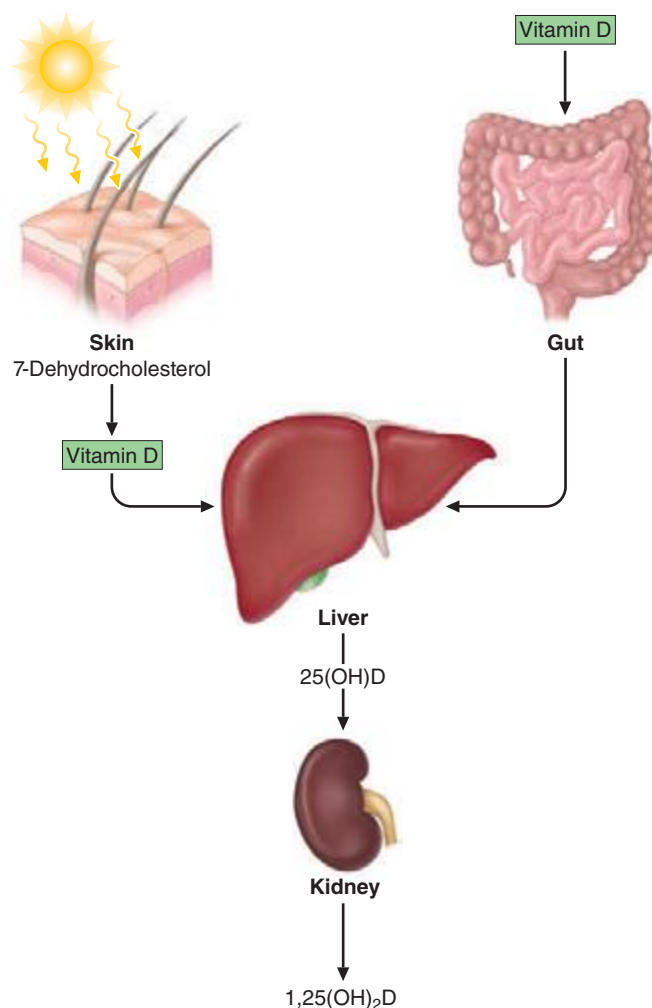


FIGURE 423-4 Vitamin D synthesis and activation. Vitamin D is synthesized in the skin in response to ultraviolet radiation and also is absorbed from the diet. It is then transported to the liver, where it undergoes 25-hydroxylation. This metabolite is the major circulating form of vitamin D. The final step in hormone activation, 1α -hydroxylation, occurs in the kidney.

skin, a photochemical cleavage results in the formation of vitamin D from 7-dehydrocholesterol. Cutaneous production of vitamin D is decreased by melanin and high solar protection factor sunblocks, which effectively impair skin penetration by ultraviolet light. The increased use of sunblocks in North America and Western Europe and a reduction in the magnitude of solar exposure of the general population over the last several decades has led to an increased reliance on dietary sources of vitamin D. In the United States and Canada, these sources largely consist of fortified cereals and dairy products, in addition to fish oils and egg yolks. Vitamin D from plant sources is in the form of vitamin D_2 , whereas that from animal sources is vitamin D_3 . These two forms have equivalent biologic activity and are activated equally well by the vitamin D hydroxylases in humans. Vitamin D enters the circulation, whether absorbed from the intestine or synthesized cutaneously, bound to vitamin D-binding protein, an α -globulin synthesized in the liver. Vitamin D is subsequently 25-hydroxylated in the liver by cytochrome P450-like enzymes in the mitochondria and microsomes. The activity of this hydroxylase is not tightly regulated, and the resultant metabolite, 25-hydroxyvitamin D ($25[\text{OH}]\text{D}$), is the major circulating and storage form of vitamin D. Approximately 88% of $25(\text{OH})\text{D}$ circulates bound to the vitamin D-binding protein, 0.03% is free, and the rest circulates bound to albumin. The half-life of $25(\text{OH})\text{D}$ is approximately 2–3 weeks; however, it is shortened dramatically when vitamin D-binding protein levels are reduced, as can occur with increased urinary losses in the nephrotic syndrome.

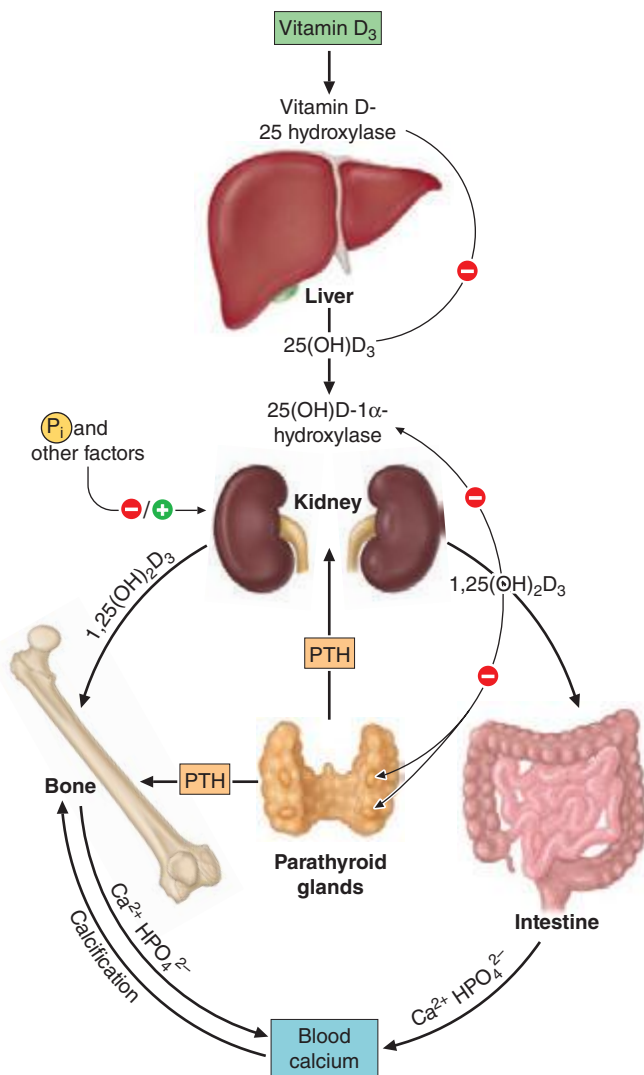


FIGURE 423-5 Schematic representation of the hormonal control loop for vitamin D metabolism and function.

A reduction in the serum calcium below ~ 2.2 mmol/L (8.8 mg/dL) prompts a proportional increase in the secretion of parathyroid hormone (PTH) and so mobilizes additional calcium from the bone. PTH promotes the synthesis of $1,25(\text{OH})_2\text{D}_3$ in the kidney, which in turn stimulates the mobilization of calcium from bone and intestine and regulates the synthesis of PTH by negative feedback.

The second hydroxylation, required for the formation of the mature hormone, occurs in the kidney (Fig. 423-5). The 25-hydroxyvitamin D-1 α -hydroxylase is a tightly regulated cytochrome P450-like mixed-function oxidase expressed in the proximal convoluted tubule cells of the kidney. PTH and hypophosphatemia are the major inducers of this microsomal enzyme, whereas calcium, FGF23, and the enzyme's product, $1,25(\text{OH})_2\text{D}_3$, repress it. The 25-hydroxyvitamin D-1 α -hydroxylase is also present in epidermal keratinocytes, but keratinocyte production of $1,25(\text{OH})_2\text{D}_3$ is not thought to contribute to circulating levels of this hormone. In addition to being present in the trophoblastic layer of the placenta, the 1 α -hydroxylase is produced by macrophages associated with granulomas and lymphomas. In these latter pathologic states, the activity of the enzyme is induced by interferon γ and TNF- α but is not regulated by calcium or $1,25(\text{OH})_2\text{D}_3$; therefore, hypercalcemia, associated with elevated levels of $1,25(\text{OH})_2\text{D}_3$, may be observed. Treatment of sarcoidosis-associated hypercalcemia with glucocorticoids, ketoconazole, or chloroquine reduces $1,25(\text{OH})_2\text{D}_3$ production and effectively lowers serum calcium. In contrast, chloroquine has not been shown to lower the elevated serum $1,25(\text{OH})_2\text{D}_3$ levels in patients with lymphoma.

The major pathway for inactivation of vitamin D metabolites is an additional hydroxylation step by the vitamin D 24-hydroxylase, an enzyme that is expressed in most tissues. $1,25(\text{OH})_2\text{D}_3$ is the major inducer of this enzyme; therefore, this hormone promotes its own inactivation, thereby limiting its biologic effects. Mutations of the gene encoding this enzyme (CYP24A1) can lead to infantile hypercalcemia and, in those less severely affected, long-standing hypercalciuria, nephrocalcinosis, and nephrolithiasis.

Polar metabolites of $1,25(\text{OH})_2\text{D}_3$ are secreted into the bile and reabsorbed via the enterohepatic circulation. Impairment of this recirculation, which is seen with diseases of the terminal ileum, leads to accelerated losses of vitamin D metabolites.

ACTIONS OF $1,25(\text{OH})_2\text{D}_3$

$1,25(\text{OH})_2\text{D}_3$ mediates its biologic effects by binding to a member of the nuclear receptor superfamily, the vitamin D receptor (VDR). This receptor belongs to the subfamily that includes the thyroid hormone receptors, the retinoid receptors, and the peroxisome proliferator-activated receptors; however, in contrast to the other members of this subfamily, only one VDR isoform has been isolated. The VDR binds to target DNA sequences as a heterodimer with the retinoid X receptor, recruiting a series of coactivators that modify chromatin and approximate the VDR to the basal transcriptional apparatus, resulting in the induction of target gene expression. The mechanism of transcriptional repression by the VDR varies with different target genes but has been shown to involve either interference with the action of activating transcription factors or the recruitment of novel proteins to the VDR complex, resulting in transcriptional repression.

The affinity of the VDR for $1,25(\text{OH})_2\text{D}_3$ is approximately three orders of magnitude higher than that for other vitamin D metabolites. In normal physiologic circumstances, these other metabolites are not thought to stimulate receptor-dependent actions. However, in states of vitamin D toxicity, the markedly elevated levels of $25(\text{OH})\text{D}_3$ may lead to hypercalcemia by interacting directly with the VDR and by displacing $1,25(\text{OH})_2\text{D}_3$ from vitamin D-binding protein, resulting in increased bioavailability of the active hormone.

The VDR is expressed in a wide range of cells and tissues. The molecular actions of $1,25(\text{OH})_2\text{D}_3$ have been studied most extensively in tissues involved in the regulation of mineral ion homeostasis. This hormone is a major inducer of calbindin 9K, a calcium-binding protein expressed in the intestine, which is thought to play an important role in the active transport of calcium across the enterocyte. The two major calcium transporters expressed by intestinal epithelia, TRPV5 and TRPV6 (transient receptor potential vanilloid), are also vitamin D responsive. By inducing the expression of these and other genes in the small intestine, $1,25(\text{OH})_2\text{D}_3$ increases the efficiency of intestinal calcium absorption, and it also has been shown to have several important actions in the skeleton. The VDR is expressed in osteoblasts and regulates the expression of several genes in this cell. These genes include the bone matrix proteins osteocalcin and osteopontin, which are upregulated by $1,25(\text{OH})_2\text{D}_3$, in addition to type I collagen, which is transcriptionally repressed by $1,25(\text{OH})_2\text{D}_3$. Both $1,25(\text{OH})_2\text{D}_3$ and PTH induce the expression of RANK ligand, which promotes osteoclast differentiation and increases osteoclast activity, by binding to RANK on osteoclast progenitors and mature osteoclasts. This is the mechanism by which $1,25(\text{OH})_2\text{D}_3$ induces bone resorption. However, the skeletal features associated with VDR-knockout mice (rickets, osteomalacia) are largely corrected by increasing calcium and phosphorus intake, underscoring the importance of vitamin D action in the gut.

The VDR is expressed in the parathyroid gland, and $1,25(\text{OH})_2\text{D}_3$ has been shown to have antiproliferative effects on parathyroid cells and to suppress the transcription of the PTH gene. These effects of $1,25(\text{OH})_2\text{D}_3$ on the parathyroid gland are an important part of the rationale for current therapies directed at preventing and treating hyperparathyroidism associated with renal insufficiency.

The VDR is also expressed in tissues and organs that do not play a role in mineral ion homeostasis. Notable in this respect is the observation that $1,25(\text{OH})_2\text{D}_3$ has an antiproliferative effect on several cell types, including keratinocytes, breast cancer cells, and prostate cancer

cells. The effects of $1,25(\text{OH})_2\text{D}$ and the VDR on keratinocytes are particularly intriguing. Alopecia is seen in humans and mice with mutant VDRs but is not a feature of vitamin D deficiency; thus, the effects of the VDR on the hair follicle are ligand-independent.

VITAMIN D DEFICIENCY

The mounting concern about the relationship between solar exposure and the development of skin cancer has led to increased reliance on dietary sources of vitamin D. Although the prevalence of vitamin D deficiency varies, the third National Health and Nutrition Examination Survey (NHANES III) revealed that vitamin D deficiency is prevalent throughout the United States. The clinical syndrome of vitamin D deficiency can be a result of deficient production of vitamin D in the skin, lack of dietary intake, accelerated losses of vitamin D, impaired vitamin D activation, or resistance to the biologic effects of $1,25(\text{OH})_2\text{D}$ (Table 423-6). The elderly and nursing home residents are particularly at risk for vitamin D deficiency, since both the efficiency of vitamin D synthesis in the skin and the absorption of vitamin D from the intestine decline with age. Similarly, intestinal malabsorption of dietary fats and short bowel syndrome, including that associated with intestinal bypass surgery, can lead to vitamin D deficiency. This is further exacerbated in the presence of terminal ileal disease, which results in impaired enterohepatic circulation of vitamin D metabolites. In addition to intestinal diseases, accelerated inactivation of vitamin D metabolites can be seen with drugs that induce hepatic cytochrome P450 mixed-function oxidases such as barbiturates, phenytoin, and rifampin. Impaired 25-hydroxylation, associated with severe liver disease or isoniazid, is an uncommon cause of vitamin D deficiency. A mutation in the gene responsible for 25-hydroxylation has been identified in one kindred. Impaired 1α -hydroxylation is prevalent in the population with profound renal dysfunction due to an increase in circulating FGF23 levels and a decrease in functional renal mass. Thus, therapeutic interventions should be considered in patients whose creatinine clearance is <0.5 mL/s (30 mL/min). Mutations in the renal 1α -hydroxylase are the basis for the genetic disorder, pseudovitamin D–deficiency rickets. This autosomal recessive disorder presents with the syndrome of vitamin D deficiency in the first year of life. Patients present with growth retardation, rickets, and hypocalcemic seizures. Serum $1,25(\text{OH})_2\text{D}$ levels are low despite normal 25(OH)D levels and elevated PTH levels. Treatment with vitamin D metabolites that do not require 1α -hydroxylation results in disease remission, although lifelong therapy is required. A second autosomal recessive disorder, hereditary vitamin D–resistant rickets, a consequence of vitamin D receptor mutations, is a greater therapeutic challenge. These patients present in a similar fashion during the first year of life, but alopecia often accompanies the disorder, demonstrating a functional role of the VDR in postnatal hair regeneration. Serum levels of $1,25(\text{OH})_2\text{D}$ are dramatically elevated in these individuals both because of increased production due to stimulation of 1α -hydroxylase activity as a consequence of secondary hyperparathyroidism and because of impaired inactivation, since induction of the 24-hydroxylase by $1,25(\text{OH})_2\text{D}$ requires an intact VDR. Because the receptor mutation results in hormone resistance, daily calcium and phosphorus infusions may be required to bypass the defect in intestinal mineral ion absorption.

TABLE 423-6 CAUSES OF IMPAIRED VITAMIN D ACTION

Vitamin D deficiency	Impaired 1α -hydroxylation
Impaired cutaneous production	Hypoparathyroidism
Dietary absence	Renal failure
Malabsorption	Ketoconazole
Accelerated loss of vitamin D	1α -hydroxylase mutation
Increased metabolism (barbiturates, phenytoin, rifampin)	Oncogenic osteomalacia
Impaired enterohepatic circulation	X-linked hypophosphatemic rickets
Nephrotic syndrome	Vitamin D receptor mutation
Impaired 25-hydroxylation	Phenytoin
Liver disease, isoniazid	

Regardless of the cause, the clinical manifestations of vitamin D deficiency are largely a consequence of impaired intestinal calcium absorption. Mild to moderate vitamin D deficiency is asymptomatic, whereas long-standing vitamin D deficiency results in hypocalcemia accompanied by secondary hyperparathyroidism, impaired mineralization of the skeleton (osteopenia on x-ray or decreased bone mineral density), and proximal myopathy. Vitamin D deficiency also has been shown to be associated with an increase in overall mortality, including cardiovascular causes. In the absence of an intercurrent illness, the hypocalcemia associated with long-standing vitamin D deficiency rarely presents with acute symptoms of hypocalcemia such as numbness, tingling, and seizures. However, the concurrent development of hypomagnesemia, which impairs parathyroid function, or the administration of potent bisphosphonates, which impair bone resorption, can lead to acute symptomatic hypocalcemia in vitamin D–deficient individuals.

Rickets and Osteomalacia In children, before epiphyseal fusion, vitamin D deficiency results in growth retardation associated with an expansion of the growth plate known as *rickets*. Three layers of chondrocytes are present in the normal growth plate: the reserve zone, the proliferating zone, and the hypertrophic zone. Rickets associated with impaired vitamin D action is characterized by expansion of the hypertrophic chondrocyte layer. The proliferation and differentiation of the chondrocytes in the rachitic growth plate are normal, and the expansion of the growth plate is a consequence of impaired apoptosis of the late hypertrophic chondrocytes, an event that precedes replacement of these cells by osteoblasts during endochondral bone formation. Investigations in murine models demonstrate that hypophosphatemia, which in vitamin D deficiency is a consequence of secondary hyperparathyroidism, is a key etiologic factor in the development of the rachitic growth plate.

The hypocalcemia and hypophosphatemia that accompany vitamin D deficiency result in impaired mineralization of bone matrix proteins, a condition known as *osteomalacia*. Osteomalacia is also a feature of long-standing hypophosphatemia, which may be a consequence of renal phosphate wasting or chronic use of etidronate or phosphate-binding antacids. This hypomineralized matrix is biomechanically inferior to normal bone; as a result, patients with vitamin D deficiency are prone to bowing of weight-bearing extremities and skeletal fractures. Vitamin D and calcium supplementation have been shown to decrease the incidence of hip fracture among ambulatory nursing home residents in France, suggesting that undermineralization of bone contributes significantly to morbidity in the elderly. Proximal myopathy is a striking feature of severe vitamin D deficiency both in children and in adults. Rapid resolution of the myopathy is observed upon vitamin D treatment.

Although vitamin D deficiency is the most common cause of rickets and osteomalacia, many disorders lead to inadequate mineralization of the growth plate and bone. Calcium deficiency without vitamin D deficiency, the disorders of vitamin D metabolism previously discussed, and hypophosphatemia can all lead to inefficient mineralization. Even in the presence of normal calcium and phosphate levels, chronic acidosis and drugs such as bisphosphonates can lead to osteomalacia. The inorganic calcium/phosphate mineral phase of bone cannot form at low pH, and bisphosphonates bind to and prevent mineral crystal growth. Because alkaline phosphatase is necessary for normal mineral deposition, probably because the enzyme can hydrolyze inhibitors of mineralization such as inorganic pyrophosphate, genetic inactivation of the alkaline phosphatase gene (hereditary hypophosphatasia) also can lead to osteomalacia in the setting of normal calcium and phosphate levels.

Diagnosis of Vitamin D Deficiency, Rickets, and Osteomalacia The most specific screening test for vitamin D deficiency in otherwise healthy individuals is a serum 25(OH)D level. Although the normal ranges vary, levels of 25(OH)D <37 nmol/L (<15 ng/mL) are associated with increasing PTH levels and lower bone density. The Institute of Medicine has defined vitamin D sufficiency as a vitamin D level >50 nmol/L (>20 ng/mL), although higher levels may be required to

2466 optimize intestinal calcium absorption in the elderly and those with underlying disease states. Vitamin D deficiency leads to impaired intestinal absorption of calcium, resulting in decreased serum total and ionized calcium values. This hypocalcemia results in secondary hyperparathyroidism, a homeostatic response that initially maintains serum calcium levels at the expense of the skeleton. Due to the PTH-induced increase in bone turnover, alkaline phosphatase levels are often increased. In addition to increasing bone resorption, PTH decreases urinary calcium excretion while promoting phosphaturia. This results in hypophosphatemia, which exacerbates the mineralization defect in the skeleton. With prolonged vitamin D deficiency resulting in osteomalacia, calcium stores in the skeleton become relatively inaccessible, since osteoclasts cannot resorb unmineralized osteoid, and frank hypocalcemia ensues. Because PTH is a major stimulus for the renal 25(OH)D 1 α -hydroxylase, there is increased synthesis of the active hormone, 1,25(OH)₂D. Paradoxically, levels of this hormone are often normal in severe vitamin D deficiency. Therefore, measurements of 1,25(OH)₂D are not accurate reflections of vitamin D stores and should not be used to diagnose vitamin D deficiency in patients with normal renal function.

Radiologic features of vitamin D deficiency in children include a widened, expanded growth plate that is characteristic of rickets. These findings not only are apparent in the long bones but also are present at the costochondral junction, where the expansion of the growth plate leads to swellings known as the “rachitic rosary.” Impairment of intramembranous bone mineralization leads to delayed fusion of the calvarial sutures and a decrease in the radiopacity of cortical bone in the long bones. If vitamin D deficiency occurs after epiphyseal fusion, the main radiologic finding is a decrease in cortical thickness and relative radiolucency of the skeleton. A specific radiologic feature of osteomalacia, whether associated with phosphate wasting or vitamin D deficiency, is pseudofractures, or Looser’s zones. These are radiolucent lines that occur where large arteries are in contact with the underlying skeletal elements; it is thought that the arterial pulsations lead to the radiolucencies. As a result, these pseudofractures are usually a few millimeters wide, are several centimeters long, and are seen particularly in the scapula, the pelvis, and the femoral neck.

TREATMENT VITAMIN D DEFICIENCY

Based on the Institute of Medicine 2010 report, the recommended daily intake of vitamin D is 600 IU from 1 to 70 years of age, and 800 IU for those over 70. Based on the observation that 800 IU of vitamin D, with calcium supplementation, decreases the risk of hip fractures in elderly women, this higher dose is thought to be an appropriate daily intake for prevention of vitamin D deficiency in adults. The safety margin for vitamin D is large, and vitamin D toxicity usually is observed only in patients taking doses in the range of 40,000 IU daily. Treatment of vitamin D deficiency should be directed at the underlying disorder, if possible, and also should be tailored to the severity of the condition. Vitamin D should always be repleted in conjunction with calcium supplementation because most of the consequences of vitamin D deficiency are a result of impaired mineral ion homeostasis. In patients in whom 1 α -hydroxylation is impaired, metabolites that do not require this activation step are the treatment of choice. They include 1,25(OH)₂D₃ (calcitriol [Rocaltrol], 0.25–0.5 μ g/d) and 1 α -hydroxyvitamin D₂ (Hectorol, 2.5–5 μ g/d). If the pathway required for activation of vitamin D is intact, severe vitamin D deficiency can be treated with pharmacologic repletion initially (50,000 IU weekly for 3–12 weeks), followed by maintenance therapy (800 IU daily). Pharmacologic doses may be required for maintenance therapy in patients who are taking medications, such as barbiturates or phenytoin, that accelerate metabolism of or cause resistance to 1,25(OH)₂D. Calcium supplementation should include 1.5–2 g/d of elemental calcium. Normocalcemia is usually observed within 1 week of the institution of therapy, although increases in PTH and alkaline phosphatase levels may persist for 3–6 months. The most efficacious methods to monitor treatment and resolution of vitamin D deficiency are serum and urinary calcium

measurements. In patients who are vitamin D replete and are taking adequate calcium supplementation, the 24-h urinary calcium excretion should be in the range of 100–250 mg/24 h. Lower levels suggest problems with adherence to the treatment regimen or with absorption of calcium or vitamin D supplements. Levels >250 mg/24 h predispose to nephrolithiasis and should lead to a reduction in vitamin D dosage and/or calcium supplementation.

424 Disorders of the Parathyroid Gland and Calcium Homeostasis

John T. Potts, Jr., Harald Jüppner

The four parathyroid glands are located posterior to the thyroid gland. They produce parathyroid hormone (PTH), which is the primary regulator of calcium physiology. PTH acts directly on bone, where it induces calcium release; on the kidney, where it enhances calcium reabsorption in the distal tubules; and in the proximal renal tubules, where it synthesizes 1,25-dihydroxyvitamin D (1,25[OH]₂D), a hormone that increases gastrointestinal calcium absorption. Serum PTH levels are tightly regulated by a negative feedback loop. Calcium, acting through the calcium-sensing receptor, and vitamin D, acting through its nuclear receptor, reduce PTH release and synthesis. Additional evidence indicates that fibroblast growth factor 23 (FGF23), a phosphaturic hormone, can suppress PTH secretion. Understanding the hormonal pathways that regulate calcium levels and bone metabolism is essential for effective diagnosis and management of a wide array of hyper- and hypocalcemic disorders.

Hyperparathyroidism, characterized by excess production of PTH, is a common cause of hypercalcemia and is usually the result of autonomously functioning adenomas or hyperplasia. Surgery for this disorder is highly effective and has been shown to reverse some of the deleterious effects of long-standing PTH excess on bone density. Humoral hypercalcemia of malignancy is also common and is usually due to the overproduction of parathyroid hormone-related peptide (PTHrP) by cancer cells. The similarities in the biochemical characteristics of hyperparathyroidism and humoral hypercalcemia of malignancy, first noted by Albright in 1941, are now known to reflect the actions of PTH and PTHrP through the same G protein-coupled PTH/PTHrP receptor.

The genetic basis of multiple endocrine neoplasia (MEN) types 1 and 2, familial hypocalciuric hypercalcemia (FHH), different forms of pseudohypoparathyroidism, Jansen’s syndrome, disorders of vitamin D synthesis and action, and the molecular events associated with parathyroid gland neoplasia have provided new insights into the regulation of calcium homeostasis. PTH and possibly some of its analogues are promising therapeutic agents for the treatment of postmenopausal or senile osteoporosis, and calcimimetic agents, which activate the calcium-sensing receptor, have provided new approaches for PTH suppression.

PARATHYROID HORMONE

PHYSIOLOGY

The primary function of PTH is to maintain the extracellular fluid (ECF) calcium concentration within a narrow normal range. The hormone acts directly on bone and kidney and indirectly on the intestine through its effects on synthesis of 1,25(OH)₂D to increase serum calcium concentrations; in turn, PTH production is closely regulated by the concentration of serum ionized calcium. This feedback system is the critical homeostatic mechanism for maintenance of ECF calcium. Any tendency toward hypocalcemia, as might be induced by calcium- or vitamin D-deficient diets, is counteracted by an increased secretion of PTH. This in turn (1) increases the rate of dissolution

of bone mineral, thereby increasing the flow of calcium from bone into blood; (2) reduces the renal clearance of calcium, returning more of the calcium and phosphate filtered at the glomerulus into ECF; and (3) increases the efficiency of calcium absorption in the intestine by stimulating the production of $1,25(\text{OH})_2\text{D}$. Immediate control of blood calcium is due to PTH effects on bone and, to a lesser extent, on renal calcium clearance. Maintenance of steady-state calcium balance, on the other hand, probably results from the effects of $1,25(\text{OH})_2\text{D}$ on calcium absorption (Chap. 423). The renal actions of the hormone are exerted at multiple sites and include inhibition of phosphate transport (proximal tubule), augmentation of calcium reabsorption (distal tubule), and stimulation of the renal $25(\text{OH})\text{D}-1\alpha$ -hydroxylase. As much as 12 mmol (500 mg) of calcium is transferred between the ECF and bone each day (a large amount in relation to the total ECF calcium pool), and PTH has a major effect on this transfer. The homeostatic role of the hormone can preserve calcium concentration in blood at the cost of bone demineralization.

PTH has multiple actions on bone, some direct and some indirect. PTH-mediated changes in bone calcium release can be seen within minutes. The chronic effects of PTH are to increase the number of bone cells, both osteoblasts and osteoclasts, and to increase the remodeling of bone; these effects are apparent within hours after the hormone is given and persist for hours after PTH is withdrawn. Continuous exposure to elevated PTH (as in hyperparathyroidism or long-term infusions in animals) leads to increased osteoclast-mediated bone resorption. However, the intermittent administration of PTH, elevating hormone levels for 1–2 h each day, leads to a net stimulation of bone formation rather than bone breakdown. Striking increases, especially in trabecular bone in the spine and hip, have been reported with the use of PTH in combination with estrogen. PTH(1–34) as monotherapy caused a highly significant reduction in fracture incidence in a worldwide placebo-controlled trial.

Osteoblasts (or stromal cell precursors), which have PTH/PTHrP receptors, are crucial to this bone-forming effect of PTH; osteoclasts, which mediate bone breakdown, lack such receptors. PTH-mediated stimulation of osteoclasts is indirect, acting in part through cytokines released from osteoblasts to activate osteoclasts; in experimental studies of bone resorption *in vitro*, osteoblasts must be present for PTH to activate osteoclasts to resorb bone (Chap. 423).

STRUCTURE

PTH is an 84-amino-acid single-chain peptide. The amino-terminal portion, PTH(1–34), is highly conserved and is critical for the biologic actions of the molecule. Modified synthetic fragments of the amino-terminal sequence as small as PTH(1–11) are sufficient to activate the PTH/PTHrP receptor (see below). The carboxyl-terminal region of the full-length PTH(1–84) molecule also can bind to a separate binding protein/receptor (cPTH-R), but this receptor has been incompletely characterized. Fragments shortened at the amino-terminus possibly by binding to cPTH-R can reduce, directly or indirectly, some of the biologic actions of full-length PTH(1–84) and of PTH(1–34).

BIOSYNTHESIS, SECRETION, AND METABOLISM

Synthesis Parathyroid cells have multiple methods of adapting to increased needs for PTH production. Most rapid (within minutes) is secretion of preformed hormone in response to hypocalcemia. Second, within hours, PTH mRNA expression is induced by sustained hypocalcemia. Finally, protracted challenge leads within days to cellular replication to increase parathyroid gland mass.

PTH is initially synthesized as a larger molecule (preparathyroid hormone, consisting of 115 amino acids). After a first cleavage step to remove the “pre” sequence of 25 amino acid residues, a second cleavage step removes the “pro” sequence of 6 amino acid residues before secretion of the mature peptide comprising 84 residues. Mutations in the preprotein region of the gene can cause hypoparathyroidism by interfering with hormone synthesis, transport, or secretion.

Transcriptional suppression of the PTH gene by calcium is nearly maximal at physiologic calcium concentrations. Hypocalcemia increases

transcriptional activity within hours. $1,25(\text{OH})_2\text{D}$ strongly suppresses PTH gene transcription. In patients with renal failure, IV administration of supraphysiologic levels of $1,25(\text{OH})_2\text{D}$ or analogues of this active metabolite can dramatically suppress PTH overproduction, which is sometimes difficult to control due to severe secondary hyperparathyroidism. Regulation of proteolytic destruction of preformed hormone (posttranslational regulation of hormone production) is an important mechanism for mediating rapid (within minutes) changes in hormone availability. High calcium increases and low calcium inhibit the proteolytic destruction of stored hormone.

Regulation of PTH Secretion PTH secretion increases steeply to a maximum value of about five times the basal rate of secretion as the calcium concentration falls from normal to the range of 1.9–2.0 mmol/L (7.6–8.0 mg/dL; measured as total calcium). However, the ionized fraction of blood calcium is the important determinant of hormone secretion. Severe intracellular magnesium deficiency impairs PTH secretion (see below).

ECF calcium controls PTH secretion by interaction with a calcium-sensing receptor (CaSR), a G protein-coupled receptor (GPCR) for which Ca^{2+} ions act as the primary ligand (see below). This receptor is a member of a distinctive subgroup of the GPCR superfamily that mediates its actions through the alpha-subunits of two related signaling G proteins, namely Gq and G11, and is characterized by a large extracellular domain suitable for “clamping” the small-molecule ligand. Stimulation of the CaSR by high calcium levels suppresses PTH secretion. The CaSR is present in parathyroid glands and the calcitonin-secreting cells of the thyroid (C cells), as well as in multiple other sites, including brain and kidney. Genetic evidence has revealed a key biologic role for the CaSR in parathyroid gland responsiveness to calcium and in renal calcium clearance. Heterozygous loss-of-function mutations in CaSR cause the syndrome of FHH, in which the blood calcium abnormality resembles that observed in hyperparathyroidism but with hypocalciuria; two more recently defined variants of FHH, FHH2 and FHH3, are caused either by heterozygous mutations in G11, one of the signaling proteins downstream of the CaSR, or by heterozygous mutations in *AP2S1*. Homozygous loss-of-function mutations in the CaSR are the cause of severe neonatal hyperparathyroidism, a disorder that can be lethal if not treated within the first days of life. On the other hand, heterozygous gain-of-function mutations cause a form of hypocalcemia resembling hypoparathyroidism (see below).

Metabolism The secreted form of PTH is indistinguishable by immunologic criteria and by molecular size from the 84-amino-acid peptide (PTH[1–84]) extracted from glands. However, much of the immunoreactive material found in the circulation is smaller than the extracted or secreted hormone. The principal circulating fragments of immunoreactive hormone lack a portion of the critical amino-terminal sequence required for biologic activity and, hence, are biologically inactive fragments (so-called middle and carboxyl-terminal fragments). Much of the proteolysis of the hormone occurs in the liver and kidney. Peripheral metabolism of PTH does not appear to be regulated by physiologic states (high versus low calcium, etc.); hence, peripheral metabolism of hormone, although responsible for rapid clearance of secreted hormone, appears to be a high-capacity, metabolically invariant catabolic process.

The rate of clearance of the secreted 84-amino-acid peptide from blood is more rapid than the rate of clearance of the biologically inactive fragment(s) corresponding to the middle and carboxyl-terminal regions of PTH. Consequently, the interpretation of results obtained with earlier PTH radioimmunoassays was influenced by the nature of the peptide fragments detected by the antibodies.

Although the problems inherent in PTH measurements have been largely circumvented by use of double-antibody immunometric assays, it is now known that some of these assays detect, besides the intact molecule, large amino-terminally truncated forms of PTH, which are present in normal and uremic individuals in addition to PTH(1–84). The concentration of these fragments relative to that of intact PTH(1–84) is higher with induced hypercalcemia than in eucalcemic

2468 or hypocalcemic conditions and is higher in patients with impaired renal function. PTH(7–84) has been identified as a major component of these amino-terminally truncated fragments. Growing evidence suggests that the PTH(7–84) (and probably related amino-terminally truncated fragments) can act, through yet undefined mechanisms, as an inhibitor of PTH action and may be of clinical significance, particularly in patients with chronic kidney disease. In this group of patients, efforts to prevent secondary hyperparathyroidism by a variety of measures (vitamin D analogues, higher calcium intake, higher dialysate calcium, phosphate-lowering strategies, and calcimetic drugs) can lead to oversuppression of the parathyroid glands since some amino-terminally truncated PTH fragments, such as PTH(7–84), react in many immunometric PTH assays (now termed second-generation assays; see below under “Diagnosis”), thus overestimating the levels of biologically active, intact PTH. Such excessive parathyroid gland suppression in chronic kidney disease can lead to adynamic bone disease (see below), which has been associated with further impaired growth in children and increased bone fracture rates in adults, and can furthermore lead to significant hypercalcemia. The measurement of PTH with newer third-generation immunoassays, which use detection antibodies directed against extreme amino-terminal PTH epitopes and thus detect only full-length PTH(1–84), may provide some advantage to prevent bone disease in chronic kidney disease.

PARATHYROID HORMONE–RELATED PROTEIN (PTHrP)

PTHrP is responsible for most instances of humoral hypercalcemia of malignancy (Chap. 121), a syndrome that resembles primary hyperparathyroidism but without elevated PTH levels. Most cell types normally produce PTHrP, including brain, pancreas, heart, lung, mammary tissue, placenta, endothelial cells, and smooth muscle. In fetal animals, PTHrP directs transplacental calcium transfer, and high concentrations of PTHrP are produced in mammary tissue and secreted into milk, but the biologic significance of the very high concentrations of this hormone in breast milk is unknown. PTHrP also plays an essential role in endochondral bone formation and in branching morphogenesis of the breast, and possibly in uterine contraction and other biologic functions.

PTH and PTHrP, although products of different genes, exhibit considerable functional and structural homology (Fig. 424-1) and have evolved from a shared ancestral gene. The structure of the gene encoding human PTHrP, however, is more complex than that of PTH, containing multiple additional exons, which can undergo alternate splicing patterns during formation of the mature mRNA. Protein products of 139, 141, and 173 amino acids are produced, and other molecular forms may result from tissue-specific degradation at accessible internal cleavage sites. The biologic roles of these various molecular species and the nature of the circulating forms of PTHrP are unclear. In fact, it is uncertain whether PTHrP circulates at any significant level in adults. As a paracrine factor, PTHrP may be produced, act, and be destroyed locally within tissues. In adults, PTHrP appears to have little influence on calcium homeostasis, except in disease states, when large tumors,

especially of the squamous cell type as well as renal cell carcinomas, lead to massive overproduction of the hormone and hypercalcemia.

PTH AND PTHrP HORMONE ACTION

Both PTH and PTHrP bind to and activate the PTH/PTHrP receptor. The PTH/PTHrP receptor (also known as the PTH-1 receptor, PTH1R) belongs to a subfamily of GPCRs that includes the receptors for calcitonin, glucagon, secretin, vasoactive intestinal peptide, and other peptides. Although both ligands activate the PTH1R, the two peptides induce distinct responses in the receptor, which explains how a single receptor without isoforms can serve two biologic roles. The extracellular regions of the receptor are involved in hormone binding, and the intracellular domains, after hormone activation, bind G protein subunits to transduce hormone signaling into cellular responses through the stimulation of second messenger formation. A second receptor that binds PTH, originally termed the *PTH-2 receptor* (PTH2R), is primarily expressed in brain, pancreas, and testis. Different mammalian PTH1Rs respond equivalently to PTH and PTHrP, at least when tested with traditional assays, whereas only the human PTH2R responds efficiently to PTH (but not to PTHrP). PTH2Rs from other species show little or no stimulation of second-messenger formation in response to PTH or PTHrP. The endogenous ligand of the PTH2R was shown to be a hypothalamic peptide referred to as tubular infundibular peptide of 39 residues, TIP39, that is distantly related to PTH and PTHrP. The PTH1R and the PTH2R can be traced backward in evolutionary time to fish; in fact, the zebrafish genome contains, in addition to the PTH1R and the PTH2R orthologs, a third receptor, the PTH3R, that is more closely related to the fish PTH1R than to the fish PTH2R. The evolutionary conservation of structure and function suggests important biologic roles for these receptors, even in fish, which lack discrete parathyroid glands but produce two molecules that are closely related to mammalian PTH.

Studies using the cloned PTH1R confirm that it can be coupled to more than one G protein and second-messenger pathway, apparently explaining the multiplicity of pathways stimulated by PTH. Activation of protein kinases (A and C) and calcium transport channels is associated with a variety of hormone-specific tissue responses. These responses include inhibition of phosphate and bicarbonate transport, stimulation of calcium transport, and activation of renal 1α -hydroxylase in the kidney. The responses in bone include effects on collagen synthesis, alkaline phosphatase, ornithine decarboxylase, citrate decarboxylase, and glucose-6-phosphate dehydrogenase activities; phospholipid synthesis; and calcium and phosphate transport. Ultimately, these biochemical events lead to an integrated hormonal response in bone turnover and calcium homeostasis. PTH also activates $\text{Na}^+/\text{Ca}^{2+}$ exchangers at renal distal tubular sites and stimulates translocation of preformed calcium transport channels, moving them from the interior to the apical surface to increase tubular uptake of calcium. PTH-dependent stimulation of phosphate excretion (reducing reabsorption—the opposite effect from actions on calcium in the kidney) involves the downregulation of two sodium-dependent phosphate co-transporters, NPT2a and NPT2c, and their expression at the apical membrane, thereby reducing phosphate reabsorption in

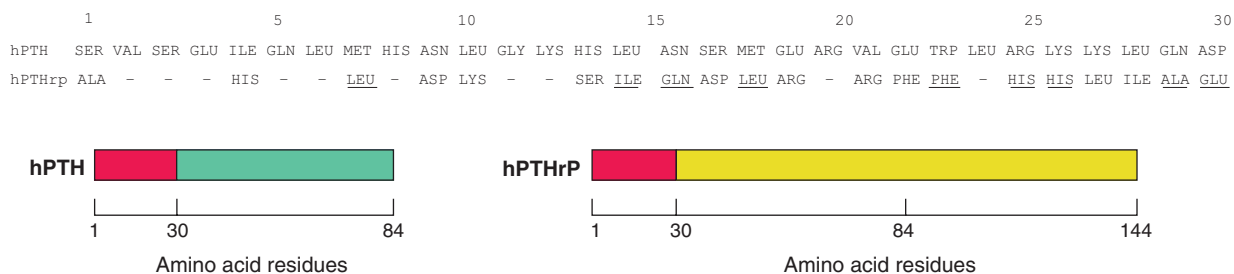


FIGURE 424-1 Schematic diagram to illustrate similarities and differences in structure of human parathyroid hormone (PTH) and human PTH-related peptide (PTHrP). Close structural (and functional) homology exists between the first 30 amino acids of hPTH and hPTHrP. The PTHrP sequence may be ≥ 144 amino acid residues in length. PTH is only 84 residues long; after residue 30, there is little structural homology between the two. *Dashed lines* in the PTHrP sequence indicate identity; *underlined residues*, although different from those of PTH, still represent conservative changes (charge or polarity preserved). Ten amino acids are identical, and a total of 20 of 30 are homologues.

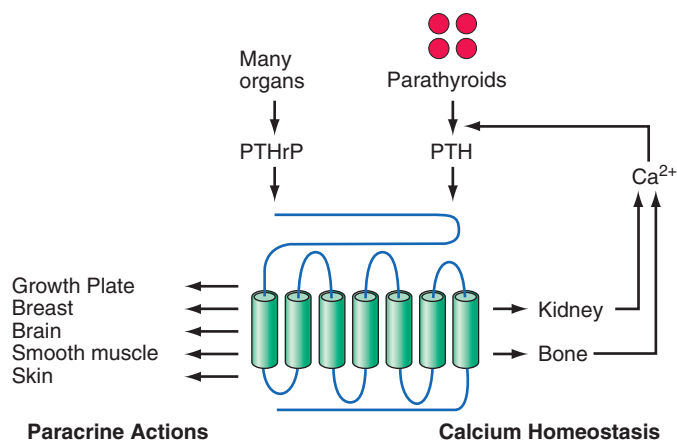


FIGURE 424-2 Dual role for the actions of the PTH/PTHrP receptor (PTH1R). Parathyroid hormone (PTH; endocrine–calcium homeostasis) and PTH-related peptide (PTHrP; paracrine–multiple tissue actions including growth plate cartilage in developing bone) use the single receptor for their disparate functions mediated by the amino-terminal 34 residues of either peptide. Other regions of both ligands interact with other receptors (not shown).

the proximal renal tubules. Similar mechanisms may be involved in other renal tubular transporters that are influenced by PTH. Recent studies reaffirm the critical linkage of blood phosphate lowering to net calcium entry into blood by PTH action and emphasize the participation of bone cells other than osteoclasts in the rapid calcium-elevating actions of PTH.

PTHrP exerts important developmental influences on fetal bone development and in adult physiology. A homozygous ablation of the gene encoding PTHrP (or disruption of the PTH1R gene) in mice causes a lethal phenotype in which animals are born with pronounced acceleration of chondrocyte maturation that resembles a lethal form of chondrodysplasia in humans that is caused by homozygous or compound heterozygous, inactivating PTH1R mutations (Fig. 424-2). Heterozygous PTH1R mutations in humans furthermore can be a cause of delayed tooth eruption, and mice that are heterozygous for ablation of the PTHrP gene display reduced mineral density consistent with osteoporosis. Experiments with these mouse models point to a hitherto unappreciated role of PTHrP as a paracrine/autocrine factor that modulates bone metabolism in adults as well as during bone development.

CALCITONIN

(See also Chap. 408) Calcitonin is a hypocalcemic peptide hormone that in several mammalian species acts as an indirect antagonist to the calcemic actions of PTH. Calcitonin seems to be of limited physiologic significance in humans, at least with regard to calcium homeostasis. It is of medical significance because of its role as a tumor marker in sporadic and hereditary cases of medullary carcinoma and its medical use as an adjunctive treatment in severe hypercalcemia and in Paget's disease of bone.

The hypocalcemic activity of calcitonin is accounted for primarily by inhibition of osteoclast-mediated bone resorption and secondarily by stimulation of renal calcium clearance. These effects are mediated by receptors on osteoclasts and renal tubular cells. Calcitonin exerts additional effects through receptors present in the brain, the gastrointestinal tract, and the immune system. The hormone, for example, exerts analgesic effects directly on cells in the hypothalamus and related structures, possibly by interacting with receptors for related peptide hormones such as calcitonin gene-related peptide (CGRP) or amylin. Both of these ligands have specific high-affinity receptors that share considerable structural similarity with the PTH1R and can also bind to and activate calcitonin receptors. The calcitonin receptor shares considerable structural similarity with the PTH1R.

The thyroid is the major source of the hormone, and the cells involved in calcitonin synthesis arise from neural crest tissue. During embryogenesis, these cells migrate into the ultimobranchial body, derived from the last branchial pouch. In submammalian vertebrates, the ultimobranchial body constitutes a discrete organ, anatomically separate from the thyroid gland; in mammals, the ultimobranchial gland fuses with and is incorporated into the thyroid gland.

The naturally occurring calcitonins consist of a peptide chain of 32 amino acids. There is considerable sequence variability among species. Calcitonin from salmon, which is used therapeutically, is 10–100 times more potent than mammalian forms in lowering serum calcium.

There are two calcitonin genes, α and β ; the transcriptional control of these genes is complex. Two different mRNA molecules are transcribed from the α gene; one is translated into the precursor for calcitonin, and the other message is translated into an alternative product, CGRP. CGRP is synthesized wherever the calcitonin mRNA is expressed (e.g., in medullary carcinoma of the thyroid). The β , or CGRP-2, gene is transcribed into the mRNA for CGRP in the central nervous system (CNS); this gene does not produce calcitonin, however. CGRP has cardiovascular actions and may serve as a neurotransmitter or play a developmental role in the CNS.

The circulating level of calcitonin in humans is lower than that in many other species. In humans, even extreme variations in calcitonin production do not change calcium and phosphate metabolism; no definite effects are attributable to calcitonin deficiency (totally thyroidectomized patients receiving only replacement thyroxine) or excess (patients with medullary carcinoma of the thyroid, a calcitonin-secreting tumor) (Chap. 408). Calcitonin has been a useful pharmacologic agent to suppress bone resorption in Paget's disease (Chap. 426e) and osteoporosis (Chap. 425) and in the treatment of hypercalcemia of malignancy (see below). However, bisphosphonates are usually more effective, and the physiologic role, if any, of calcitonin in humans is uncertain. On the other hand, ablation of the calcitonin gene (combined because of the close proximity with ablation of the CGRP gene) in mice leads to reduced bone mineral density, suggesting that its biologic role in mammals is still not fully understood.

HYPERCALCEMIA

(See also Chap. 65) Hypercalcemia can be a manifestation of a serious illness such as malignancy or can be detected coincidentally by laboratory testing in a patient with no obvious illness. The number of patients recognized with asymptomatic hypercalcemia, usually hyperparathyroidism, increased in the late twentieth century.

Whenever hypercalcemia is confirmed, a definitive diagnosis must be established. Although hyperparathyroidism, a frequent cause of asymptomatic hypercalcemia, is a chronic disorder in which manifestations, if any, may be expressed only after months or years, hypercalcemia can also be the earliest manifestation of malignancy, the second most common cause of hypercalcemia in the adult. The causes of hypercalcemia are numerous (Table 424-1), but hyperparathyroidism and cancer account for 90% of all cases.

Before undertaking a diagnostic workup, it is essential to be sure that true hypercalcemia, not a false-positive laboratory test, is present. A false-positive diagnosis of hypercalcemia is usually the result of inadvertent hemoconcentration during blood collection or elevation in serum proteins such as albumin. Hypercalcemia is a chronic problem, and it is cost-effective to obtain several serum calcium measurements; these tests need not be in the fasting state.

Clinical features are helpful in differential diagnosis. Hypercalcemia in an adult who is asymptomatic is usually due to primary hyperparathyroidism. In malignancy-associated hypercalcemia, the disease is usually not occult; rather, symptoms of malignancy bring the patient to the physician, and hypercalcemia is discovered during the evaluation. In such patients, the interval between detection of hypercalcemia and death, especially without vigorous treatment, is often <6 months. Accordingly, if an asymptomatic individual has had hypercalcemia or some manifestation of hypercalcemia such as kidney stones for more than 1 or 2 years, it is unlikely that malignancy

I. Parathyroid-Related

- A. Primary hyperparathyroidism
 - 1. Adenoma(s)
 - 2. Multiple endocrine neoplasia
 - 3. Carcinoma
- B. Lithium therapy
- C. Familial hypocalciuric hypercalcemia

II. Malignancy-Related

- A. Solid tumor with metastases (breast)
- B. Solid tumor with humoral mediation of hypercalcemia (lung, kidney)
- C. Hematologic malignancies (multiple myeloma, lymphoma, leukemia)

III. Vitamin D–Related

- A. Vitamin D intoxication
- B. ↑ 1,25(OH)₂D; sarcoidosis and other granulomatous diseases
- C. ↑ 1,25(OH)₂D; impaired 1,25(OH)₂D metabolism due to 24-hydroxylase deficiency

IV. Associated with High Bone Turnover

- A. Hyperthyroidism
- B. Immobilization
- C. Thiazides
- D. Vitamin A intoxication
- E. Fat necrosis

V. Associated with Renal Failure

- A. Severe secondary hyperparathyroidism
- B. Aluminum intoxication
- C. Milk-alkali syndrome

is the cause. Nevertheless, differentiating primary hyperparathyroidism from occult malignancy can occasionally be difficult, and careful evaluation is required, particularly when the duration of the hypercalcemia is unknown. Hypercalcemia not due to hyperparathyroidism or malignancy can result from excessive vitamin D action, impaired metabolism of 1,25(OH)₂D, high bone turnover from any of several causes, or renal failure (Table 424-1). Dietary history and a history of ingestion of vitamins or drugs are often helpful in diagnosing some of the less frequent causes. Immunometric PTH assays serve as the principal laboratory test in establishing the diagnosis.

Hypercalcemia from any cause can result in fatigue, depression, mental confusion, anorexia, nausea, vomiting, constipation, reversible renal tubular defects, increased urine output, a short QT interval in the electrocardiogram, and, in some patients, cardiac arrhythmias. There is a variable relation from one patient to the next between the severity of hypercalcemia and the symptoms. Generally, symptoms are more common at calcium levels >2.9–3.0 mmol/L (11.6–12.0 mg/dL), but some patients, even at this level, are asymptomatic. When the calcium level is >3.2 mmol/L (12.8 mg/dL), calcification in kidneys, skin, vessels, lungs, heart, and stomach occurs and renal insufficiency may develop, particularly if blood phosphate levels are normal or elevated due to impaired renal excretion. Severe hypercalcemia, usually defined as ≥3.7–4.5 mmol/L (14.8–18.0 mg/dL), can be a medical emergency; coma and cardiac arrest can occur.

Acute management of the hypercalcemia is usually successful. The type of treatment is based on the severity of the hypercalcemia and the nature of associated symptoms, as outlined below.

PRIMARY HYPERPARATHYROIDISM

Natural History and Incidence Primary hyperparathyroidism is a generalized disorder of calcium, phosphate, and bone metabolism due to an increased secretion of PTH. The elevation of circulating hormone usually leads to hypercalcemia and hypophosphatemia. There is great variation in the manifestations. Patients may present with multiple signs and symptoms, including recurrent nephrolithiasis, peptic ulcers, mental changes, and, less frequently, extensive bone

resorption. However, with greater awareness of the disease and wider use of multiphasic screening tests, including measurements of blood calcium, the diagnosis is frequently made in patients who have no symptoms and minimal, if any, signs of the disease other than hypercalcemia and elevated levels of PTH. The manifestations may be subtle, and the disease may have a benign course for many years or a lifetime. This milder form of the disease is usually termed *asymptomatic hyperparathyroidism*. Rarely, hyperparathyroidism develops or worsens abruptly and causes severe complications such as marked dehydration and coma, so-called hypercalcemic parathyroid crisis.

The annual incidence of the disease is calculated to be as high as 0.2% in patients >60, with an estimated prevalence, including undiscovered asymptomatic patients, of ≥1%; some reports suggest the incidence may be declining. If confirmed, these changing estimates may reflect less frequent routine testing of serum calcium in recent years, earlier overestimates in incidence, or unknown factors. The disease has a peak incidence between the third and fifth decades but occurs in young children and in the elderly.

Etiology Parathyroid tumors are most often encountered as isolated adenomas without other endocrinopathy. They may also arise in hereditary syndromes such as MEN syndromes. Parathyroid tumors may also arise as secondary to underlying disease (excessive stimulation in secondary hyperparathyroidism, especially chronic renal failure) or after other forms of excessive stimulation such as lithium therapy. These etiologies are discussed below.

SOLITARY ADENOMAS A single abnormal gland is the cause in ~80% of patients; the abnormality in the gland is usually a benign neoplasm or adenoma and rarely a parathyroid carcinoma. Some surgeons and pathologists report that the enlargement of multiple glands is common; double adenomas are reported. In ~15% of patients, all glands are hyperfunctioning; *chief cell parathyroid hyperplasia* is usually hereditary and frequently associated with other endocrine abnormalities.

HEREDITARY SYNDROMES AND MULTIPLE PARATHYROID TUMORS Hereditary hyperparathyroidism can occur without other endocrine abnormalities but is usually part of a *multiple endocrine neoplasia* (MEN) syndrome (**Chap. 408**). MEN 1 (Wermer's syndrome) consists of hyperparathyroidism and tumors of the pituitary and pancreas, often associated with gastric hypersecretion and peptic ulcer disease (Zollinger-Ellison syndrome). MEN 2A is characterized by pheochromocytoma and medullary carcinoma of the thyroid, as well as hyperparathyroidism; MEN 2B has additional associated features such as multiple neuromas but usually lacks hyperparathyroidism. Each of these MEN syndromes is transmitted in an apparent autosomal dominant manner, although, as noted below, the genetic basis of MEN 1 involves biallelic loss of a tumor suppressor.

The *hyperparathyroidism jaw tumor* (HPT-JT) syndrome occurs in families with parathyroid tumors (sometimes carcinomas) in association with benign jaw tumors. This disorder is caused by mutations in *CDC73* (*HRPT2*), and mutations in this gene are also observed in parathyroid cancers. Some kindreds exhibit hereditary hyperparathyroidism without other endocrinopathies. This disorder is often termed *nonsyndromic familial isolated hyperparathyroidism* (FIHP). There is speculation that these families may be examples of variable expression of the other syndromes such as MEN 1, MEN 2, or the HPT-JT syndrome, but they may also have distinctive, still unidentified genetic causes.

Pathology Adenomas are most often located in the inferior parathyroid glands, but in 6–10% of patients, parathyroid adenomas may be located in the thymus, the thyroid, the pericardium, or behind the esophagus. Adenomas are usually 0.5–5 g in size but may be as large as 10–20 g (normal glands weigh 25 mg on average). Chief cells are predominant in both hyperplasia and adenoma. With chief cell hyperplasia, the enlargement may be so asymmetric that some involved glands appear grossly normal. If generalized hyperplasia is present, however, histologic examination reveals a uniform pattern of chief cells and disappearance of fat even in the absence of an increase in gland weight. Thus, microscopic examination of biopsy specimens of several glands is essential to interpret findings at surgery.

Parathyroid carcinoma is often not aggressive. Long-term survival without recurrence is common if at initial surgery the entire gland is removed without rupture of the capsule. Recurrent parathyroid carcinoma is usually slow-growing with local spread in the neck, and surgical correction of recurrent disease may be feasible. Occasionally, however, parathyroid carcinoma is more aggressive, with distant metastases (lung, liver, and bone) found at the time of initial operation. It may be difficult to appreciate initially that a primary tumor is carcinoma; increased numbers of mitotic figures and increased fibrosis of the gland stroma may precede invasion. The diagnosis of carcinoma is often made in retrospect. Hyperparathyroidism from a parathyroid carcinoma may be indistinguishable from other forms of primary hyperparathyroidism but is usually more severe clinically. A potential clue to the diagnosis is offered by the degree of calcium elevation. Calcium values of 3.5–3.7 mmol/L (14–15 mg/dL) are frequent with carcinoma and may alert the surgeon to remove the abnormal gland with care to avoid capsular rupture. Recent findings concerning the genetic basis of parathyroid carcinoma (distinct from that of benign adenomas) indicate the need, in these kindreds, for family screening (see below).

GENETIC DEFECTS ASSOCIATED WITH HYPERPARATHYROIDISM

As in many other types of neoplasia, two fundamental types of genetic defects have been identified in parathyroid gland tumors: (1) overactivity of protooncogenes and (2) loss of function of tumor-suppressor genes. The former, by definition, can lead to uncontrolled cellular growth and function by activation (gain-of-function mutation) of a single allele of the responsible gene, whereas the latter requires loss of function of both allelic copies. Biallelic loss of function of a tumor-suppressor gene is usually characterized by a germline defect (all cells) and an additional somatic deletion/mutation in the tumor (Fig. 424-3).

Mutations in the *MEN1* gene locus, encoding the protein MENIN, on chromosome 11q13 are responsible for causing MEN 1; the normal allele of this gene fits the definition of a tumor-suppressor gene. Inheritance of one mutated allele in this hereditary syndrome, followed by loss of the other allele via somatic cell mutation, leads to monoclonal expansion and tumor development. Also, in ~15–20% of sporadic parathyroid adenomas, both alleles of the *MEN1* locus on chromosome 11 are somatically deleted, implying that the same defect responsible for MEN 1 can also cause the sporadic disease (Fig. 424-3A). Consistent with the Knudson hypothesis for two-step neoplasia in certain inherited cancer syndromes (Chap. 101e), the earlier onset of hyperparathyroidism in the hereditary syndromes reflects the need for only one mutational event to trigger the monoclonal outgrowth. In sporadic adenomas, typically occurring later in life, two different somatic events must occur before the *MEN1* gene is silenced.

Other presumptive anti-oncogenes involved in hyperparathyroidism include a still unidentified gene mapped to chromosome 1p seen in 40% of sporadic parathyroid adenomas and a gene mapped to chromosome Xp11 in patients with secondary hyperparathyroidism and renal failure, who progressed to “tertiary” hyperparathyroidism, now known to reflect monoclonal outgrowths within previously hyperplastic glands.

A more complex pattern, still incompletely resolved, arises with genetic defects and carcinoma of the parathyroids. This appears to be due to biallelic loss of a functioning copy of a gene, *HRPT2* (or *CDC73*), originally identified as the cause of the HPT-JT syndrome. Several inactivating mutations have been identified in *HRPT2* (located on chromosome 1q21-31), which encodes a 531-amino-acid protein called parafibromin. The responsible genetic mutations in *HRPT2* appear to be necessary, but not sufficient, for parathyroid cancer.

In general, the detection of additional genetic defects in these parathyroid tumor-related syndromes and the variations seen in

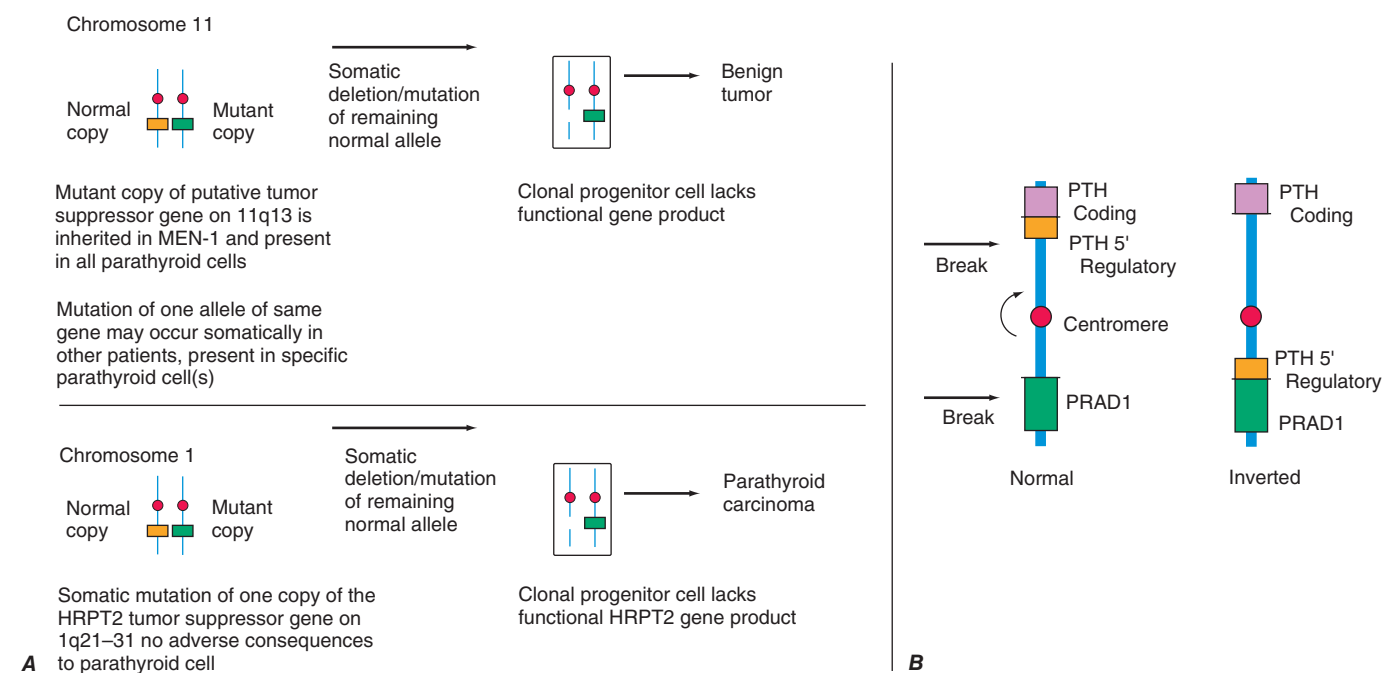


FIGURE 424-3 **A.** Schematic diagram indicating molecular events in tumor susceptibility. The patient with the hereditary abnormality (multiple endocrine neoplasia [MEN]) is envisioned as having one defective gene inherited from the affected parent on chromosome 11, but one copy of the normal gene is present from the other parent. In the monoclonal tumor (benign tumor), a somatic event, here partial chromosomal deletion, removes the remaining normal gene from a cell. In nonhereditary tumors, two successive somatic mutations must occur, a process that takes a longer time. By either pathway, the cell, deprived of growth-regulating influence from this gene, has unregulated growth and becomes a tumor. A different genetic locus also involving loss of a tumor-suppressor gene termed *HRPT2* is involved in the pathogenesis of parathyroid carcinoma. (From A Arnold: *J Clin Endocrine Metab* 77:1108, 1993. Copyright 1993, The Endocrine Society.) **B.** Schematic illustration of the mechanism and consequences of gene rearrangement and overexpression of the *PRAD1* protooncogene (pericentromeric inversion of chromosome 11) in parathyroid adenomas. The excessive expression of *PRAD1* (a cell cycle control protein, cyclin D₁) by the highly active parathyroid hormone (PTH) gene promoter in the parathyroid cell contributes to excess cellular proliferation. (From J Habener et al, in L DeGroot, JL Jameson [eds]: *Endocrinology*, 4th ed. Philadelphia, Saunders, 2001; with permission.)

2472 phenotypic expression/penetrance indicate the multiplicity of the genetic factors responsible. Nonetheless, the ability to detect the presence of the major genetic contributors has greatly aided a more informed management of family members of patients identified in the hereditary syndromes such as MEN 1, MEN 2, and HPT-JT.

An important contribution from studies on the genetic origin of parathyroid carcinoma has been the realization that the mutations involve a different pathway than that involved with the benign gland enlargements. Unlike the pathogenesis of genetic alterations seen in colon cancer, where lesions evolve from benign adenomas to malignant disease by progressive genetic changes, the alterations commonly seen in most parathyroid cancers (*HRPT2* mutations) are infrequently seen in sporadic parathyroid adenomas.

Abnormalities at the *Rb* gene were the first to be noted in parathyroid cancer. The *Rb* gene, a tumor-suppressor gene located on chromosome 13q14, was initially associated with retinoblastoma but has since been implicated in other neoplasias, including parathyroid carcinoma. Early studies implicated allelic deletions of the *Rb* gene in many parathyroid carcinomas and decreased or absent expression of the *Rb* protein. However, because there are often large deletions on chromosome 13 that include many genes in addition to the *Rb* locus (with similar findings in some pituitary carcinomas), it remains possible that other tumor-suppressor genes on chromosome 13 may be playing a role in parathyroid carcinoma.

Study of the parathyroid cancers found in some patients with the HPT-JT syndrome has led to identification of a much larger role for mutations in the *HRPT2* gene in most parathyroid carcinomas, including those that arise sporadically, without apparent association with the HPT-JT syndrome. Mutations in the coding region have been identified in 75–80% of all parathyroid cancers analyzed, leading to the conclusion that, with addition of presumed mutations in the noncoding regions, this genetic defect may be seen in essentially all parathyroid carcinomas. Of special importance was the discovery that, in some sporadic parathyroid cancers, germline mutations have been found; this, in turn, has led to careful investigation of the families of these patients and a new clinical indication for genetic testing in this setting.

Hypercalcemia occurring in family members (who are also found to have the germline mutations) can lead to the finding, at parathyroid surgery, of premalignant parathyroid tumors.

Overall, it seems there are multiple factors in parathyroid cancer, in addition to the *HRPT2* and *Rb* gene, although the *HRPT2* gene mutation is the most invariant abnormality. *RET* encodes a tyrosine kinase type receptor; specific inherited germline mutations lead to a constitutive activation of the receptor, thereby explaining the autosomal dominant mode of transmission and the relatively early onset of neoplasia. In the MEN 2 syndrome, the *RET* protooncogene may be responsible for the earliest disorder detected, the polyclonal disorder (C cell hyperplasia, which then is transformed into a clonal outgrowth—a medullary carcinoma with the participation of other, still uncharacterized genetic defects).

In some parathyroid adenomas, activation of a protooncogene has been identified (Fig. 424-3B). A reciprocal translocation involving chromosome 11 has been identified that juxtaposes the *PTH* gene promoter upstream of a gene product termed *PRAD-1*, encoding a cyclin D protein that plays a key role in normal cell division. This translocation plus other mechanisms that cause an equivalent overexpression of cyclin D1 are found in 20–40% of parathyroid adenomas.

Mouse models have confirmed the role of several of the major identified genetic defects in parathyroid disease and the MEN syndromes. Loss of the *MEN1* gene locus or overexpression of the *PRAD-1* protooncogene or the mutated *RET* protooncogene have been analyzed by genetic manipulation in mice, with the expected onset of parathyroid tumors or medullary carcinoma, respectively.

Signs and Symptoms Many patients with hyperparathyroidism are asymptomatic. Manifestations of hyperparathyroidism involve primarily the kidneys and the skeletal system. Kidney involvement, due either to deposition of calcium in the renal parenchyma or to recurrent nephrolithiasis, was present in 60–70% of patients prior to 1970. With earlier

detection, renal complications occur in <20% of patients in many large series. Renal stones are usually composed of either calcium oxalate or calcium phosphate. In occasional patients, repeated episodes of nephrolithiasis or the formation of large calculi may lead to urinary tract obstruction, infection, and loss of renal function. Nephrocalcinosis may also cause decreased renal function and phosphate retention.

The distinctive bone manifestation of hyperparathyroidism is *osteitis fibrosa cystica*, which occurred in 10–25% of patients in series reported 50 years ago. Histologically, the pathognomonic features are an increase in the giant multinucleated osteoclasts in scalloped areas on the surface of the bone (Howship's lacunae) and a replacement of the normal cellular and marrow elements by fibrous tissue. X-ray changes include resorption of the phalangeal tufts and replacement of the usually sharp cortical outline of the bone in the digits by an irregular outline (subperiosteal resorption). In recent years, osteitis fibrosa cystica is very rare in primary hyperparathyroidism, probably due to the earlier detection of the disease.

Dual-energy x-ray absorptiometry (DEXA) of the spine provides reproducible quantitative estimates (within a few percent) of spinal bone density. Similarly, bone density in the extremities can be quantified by densitometry of the hip or of the distal radius at a site chosen to be primarily cortical. Computed tomography (CT) is a very sensitive technique for estimating spinal bone density, but reproducibility of standard CT is no better than 5%. Newer CT techniques (spiral, "extreme" CT) are more reproducible but are currently available in a limited number of medical centers. Cortical bone density is reduced while cancellous bone density, especially in the spine, is relatively preserved. In symptomatic patients, dysfunctions of the CNS, peripheral nerve and muscle, gastrointestinal tract, and joints also occur. It has been reported that severe neuropsychiatric manifestations may be reversed by parathyroidectomy. When present in symptomatic patients, neuromuscular manifestations may include proximal muscle weakness, easy fatigability, and atrophy of muscles and may be so striking as to suggest a primary neuromuscular disorder. The distinguishing feature is the complete regression of neuromuscular disease after surgical correction of the hyperparathyroidism.

Gastrointestinal manifestations are sometimes subtle and include vague abdominal complaints and disorders of the stomach and pancreas. Again, cause and effect are unclear. In MEN 1 patients with hyperparathyroidism, duodenal ulcer may be the result of associated pancreatic tumors that secrete excessive quantities of gastrin (Zollinger-Ellison syndrome). Pancreatitis has been reported in association with hyperparathyroidism, but the incidence and the mechanism are not established.

Much attention has been paid in recent years to the manifestations of and optimum management strategies for asymptomatic hyperparathyroidism. This is now the most prevalent form of the disease. *Asymptomatic primary hyperparathyroidism* is defined as biochemically confirmed hyperparathyroidism (elevated or inappropriately normal PTH levels despite hypercalcemia) with the absence of signs and symptoms typically associated with more severe hyperparathyroidism such as features of renal or bone disease.

Three conferences on the topic have been held in the United States over the past two decades, with the most recent in 2008. The published proceedings include discussion of more subtle manifestations of disease, its natural history (without parathyroidectomy), and guidelines both for indications for surgery and medical monitoring in nonoperated patients.

Issues of concern include the potential for cardiovascular deterioration, the presence of subtle neuropsychiatric symptoms, and the longer-term status of skeletal integrity in patients not treated surgically. The current consensus is that medical monitoring rather than surgical correction of hyperparathyroidism may be justified in certain patients. The current recommendation is that patients who show mild disease, as defined by specific criteria (Table 424-2), can be safely followed under management guidelines (Table 424-3). There is, however, growing uncertainty about subtle disease manifestations and whether surgery is therefore indicated in most patients. Among the issues is the evidence of eventual (>8 years) deterioration in bone mineral density after a decade

TABLE 424-2 GUIDELINES FOR SURGERY IN ASYMPTOMATIC PRIMARY HYPERPARATHYROIDISM^a

Parameter	Guideline
Serum calcium (above normal)	>1 mg/dL
24-h urinary calcium	No indication
Creatinine clearance (calculated) ^b	If <60 mL/min
Bone density	T score <-2.5 at any of 3 sites ^c
Age	<50

^aJP Bilezikian et al: Guidelines for the management of asymptomatic primary hyperparathyroidism: Summary statement from the third international workshop. *J Clin Endocrinol Metab* 94:335, 2009. ^bCreatinine clearance calculated by Cockcroft-Gault equation or Modification of Diet in Renal Disease (MDRD) equation. ^cSpine, distal radius, hip.

of relative stability. There is concern that this late-onset deterioration in bone density in nonoperated patients could contribute significantly to the well-known age-dependent fracture risk (osteoporosis). One study reported significant and sustained improvements in bone mineral density after successful parathyroidectomy, again raising the issue regarding benefits of surgery. Other randomized studies, however, did not report major gains after surgery.

Cardiovascular disease including left ventricular hypertrophy, cardiac functional defects, and endothelial dysfunction have been reported as reversible in European patients with more severe symptomatic disease after surgery, leading to numerous studies of these cardiovascular features in those with milder disease. There are reports of endothelial dysfunction in patients with mild asymptomatic hyperparathyroidism, but the expert panels concluded that more observation is needed, especially regarding whether there is reversibility with surgery.

A topic of considerable interest and some debate is assessment of neuropsychiatric status and health-related quality of life (QOL) status in hyperparathyroid patients both before surgery and in response to parathyroidectomy. Several observational studies suggest considerable improvements in symptom score after surgery. Randomized studies of surgery versus observation, however, have yielded inconclusive results, especially regarding benefits of surgery. Most studies report that hyperparathyroidism is associated with increased neuropsychiatric symptoms, so the issue remains a significant factor in decisions regarding the impact of surgery in this disease.

DIAGNOSIS

The diagnosis is typically made by detecting an elevated immunoreactive PTH level in a patient with asymptomatic hypercalcemia (see “Differential Diagnosis: Special Tests,” below). Serum phosphate is usually low but may be normal, especially if renal failure has developed.

Several modifications in PTH assays have been introduced in efforts to improve their utility in light of information about metabolism of PTH (as discussed above). First-generation assays were based on displacement of radiolabeled PTH from antibodies that reacted with PTH (often also PTH fragments). Double-antibody or immunometric assays (one antibody that is usually directed against the carboxyl-terminal portion of intact PTH to capture the hormone and a second radio- or enzyme-labeled antibody that is usually directed against the amino-terminal portion of intact PTH) greatly improved the diagnostic discrimination of the tests by eliminating interference from

TABLE 424-3 GUIDELINES FOR MONITORING IN ASYMPTOMATIC PRIMARY HYPERPARATHYROIDISM^a

Parameter	Guideline
Serum calcium	Annually
24-h urinary calcium	Recommended
Creatinine clearance	Recommended
Serum creatinine ^b	Annually
Bone density	Annually (3 sites) ^c

^aUpdates guidelines (JP Bilezikian et al: *J Clin Endocrinol Metab* 2014; epub ahead of print). ^bCreatinine clearance calculated by Cockcroft-Gault equation or Modification of Diet in Renal Disease (MDRD) equation.

circulating biologically inactive fragments, detected by the original first-generation assays. Double-antibody assays are now referred to as second-generation. Such PTH assays have in some centers and testing laboratories been replaced by third-generation assays after it was discovered that large PTH fragments, devoid of only the extreme amino-terminal portion of the PTH molecule, are also present in blood and are detected, incorrectly, as intact PTH. These amino-terminally truncated PTH fragments were prevented from registering in the newer third-generation assays by use of a detection antibody directed against the extreme amino-terminal epitope. These assays may be useful for clinical research studies as in management of chronic renal disease, but the consensus is that either second- or third-generation assays are useful in the diagnosis of primary hyperparathyroidism and for the diagnosis of high-turnover bone disease in chronic kidney disease.

Many tests based on renal responses to excess PTH (renal calcium and phosphate clearance; blood phosphate, chloride, magnesium; urinary or nephrogenous cyclic AMP [cAMP]) were used in earlier decades. These tests have low specificity for hyperparathyroidism and are therefore not cost-effective; they have been replaced by PTH immunometric assays combined with simultaneous blood calcium measurements (Fig. 424-4).

TREATMENT HYPERPARATHYROIDISM

Surgical excision of the abnormal parathyroid tissue is the definitive therapy for this disease. As noted above, medical surveillance without operation for patients with mild, asymptomatic disease is, however, still preferred by some physicians and patients, particularly when the patients are more elderly. Evidence favoring surgery, if medically feasible, is growing because of concerns about skeletal,

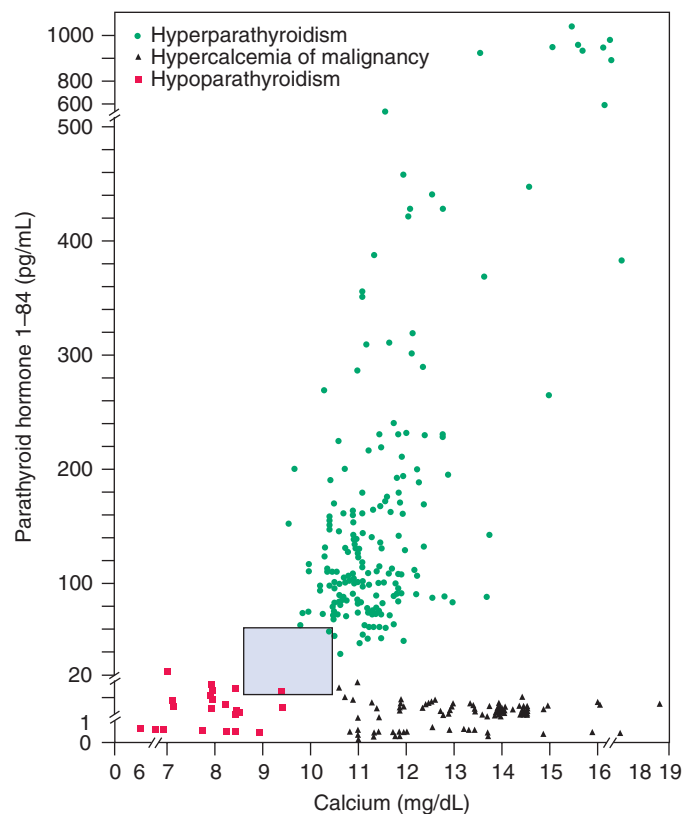


FIGURE 424-4 Levels of immunoreactive parathyroid hormone (PTH) detected in patients with primary hyperparathyroidism, hypercalcemia of malignancy, and hypoparathyroidism. Boxed area represents the upper and normal limits of blood calcium and/or immunoreactive PTH. (From SR Nussbaum, JT Potts, Jr, in L DeGroot, JL Jameson [eds]: *Endocrinology*, 4th ed. Philadelphia, Saunders, 2001; with permission.)

2474 cardiovascular, and neuropsychiatric disease, even in mild hyperparathyroidism.

Two surgical approaches are generally practiced. The conventional parathyroidectomy procedure was neck exploration with general anesthesia; this procedure is being replaced in many centers, whenever feasible, by an outpatient procedure with local anesthesia, termed *minimally invasive parathyroidectomy*.

Parathyroid exploration is challenging and should be undertaken by an experienced surgeon. Certain features help in predicting the pathology (e.g., multiple abnormal glands in familial cases). However, some critical decisions regarding management can be made only during the operation.

With conventional surgery, one approach is still based on the view that typically only one gland (the adenoma) is abnormal. If an enlarged gland is found, a normal gland should be sought. In this view, if a biopsy of a normal-sized second gland confirms its histologic (and presumed functional) normality, no further exploration, biopsy, or excision is needed. At the other extreme is the minority viewpoint that all four glands be sought and that most of the total parathyroid tissue mass be removed. The concern with the former approach is that the recurrence rate of hyperparathyroidism may be high if a second abnormal gland is missed; the latter approach could involve unnecessary surgery and an unacceptable rate of hypoparathyroidism. When normal glands are found in association with one enlarged gland, excision of the single adenoma usually leads to cure or at least years free of symptoms. Long-term follow-up studies to establish true rates of recurrence are limited.

Recently, there has been growing experience with new surgical strategies that feature a minimally invasive approach guided by improved preoperative localization and intraoperative monitoring by PTH assays. Preoperative ^{99m}Tc sestamibi scans with single-photon emission CT (SPECT) are used to predict the location of an abnormal gland and intraoperative sampling of PTH before and at 5-min intervals after removal of a suspected adenoma to confirm a rapid fall (>50%) to normal levels of PTH. In several centers, a combination of preoperative sestamibi imaging, cervical block anesthesia, minimal surgical incision, and intraoperative PTH measurements has allowed successful outpatient surgical management with a clear-cut cost benefit compared to general anesthesia and more extensive neck surgery. The use of these minimally invasive approaches requires clinical judgment to select patients unlikely to have multiple gland disease (e.g., MEN or secondary hyperparathyroidism). The growing acceptance of the technique and its relative ease for the patient has lowered the threshold for surgery.

Severe hypercalcemia may provide a preoperative clue to the presence of parathyroid carcinoma. In such cases, when neck exploration is undertaken, the tissue should be widely excised; care is taken to avoid rupture of the capsule to prevent local seeding of tumor cells.

Multiple-gland hyperplasia, as predicted in familial cases, poses more difficult questions of surgical management. Once a diagnosis of hyperplasia is established, all the glands must be identified. Two schemes have been proposed for surgical management. One is to totally remove three glands with partial excision of the fourth gland; care is taken to leave a good blood supply for the remaining gland. Other surgeons advocate total parathyroidectomy with immediate transplantation of a portion of a removed, minced parathyroid gland into the muscles of the forearm, with the view that surgical excision is easier from the ectopic site in the arm if there is recurrent hyperfunction.

In a minority of cases, if no abnormal parathyroid glands are found in the neck, the issue of further exploration must be decided. There are documented cases of five or six parathyroid glands and of unusual locations for adenomas such as in the mediastinum.

When a second parathyroid exploration is indicated, the minimally invasive techniques for preoperative localization such as ultrasound, CT scan, and isotope scanning are combined with venous sampling and/or selective digital arteriography in one of the centers specializing in these procedures. Intraoperative monitoring of PTH

levels by rapid PTH immunoassays may be useful in guiding the surgery. At one center, long-term cures have been achieved with selective embolization or injection of large amounts of contrast material into the end-arterial circulation feeding the parathyroid tumor.

A decline in serum calcium occurs within 24 h after successful surgery; usually blood calcium falls to low-normal values for 3–5 days until the remaining parathyroid tissue resumes full hormone secretion. Acute postoperative hypocalcemia is likely only if severe bone mineral deficits are present or if injury to all the normal parathyroid glands occurs during surgery. In general, there are few problems encountered in patients with uncomplicated disease such as a single adenoma (the clear majority), who do not have symptomatic bone disease or a large deficit in bone mineral, who are vitamin D and magnesium sufficient, and who have good renal and gastrointestinal function. The extent of postoperative hypocalcemia varies with the surgical approach. If all glands are biopsied, hypocalcemia may be transiently symptomatic and more prolonged. Hypocalcemia is more likely to be symptomatic after second parathyroid explorations, particularly when normal parathyroid tissue was removed at the initial operation and when the manipulation and/or biopsy of the remaining normal glands are more extensive in the search for the missing adenoma.

Patients with hyperparathyroidism have efficient intestinal calcium absorption due to the increased levels of 1,25(OH)₂D stimulated by PTH excess. Once hypocalcemia signifies successful surgery, patients can be put on a high-calcium intake or be given oral calcium supplements. Despite mild hypocalcemia, most patients do not require parenteral therapy. If the serum calcium falls to <2 mmol/L (8 mg/dL), and if the phosphate level rises simultaneously, the possibility that surgery has caused hypoparathyroidism must be considered. With unexpected hypocalcemia, coexistent hypomagnesemia should be considered, because it interferes with PTH secretion and causes functional hypoparathyroidism ([Chap. 423](#)).

Signs of hypocalcemia include symptoms such as muscle twitching, a general sense of anxiety, and positive Chvostek's and Trousseau's signs coupled with serum calcium consistently <2 mmol/L (8 mg/dL). Parenteral calcium replacement at a low level should be instituted when hypocalcemia is symptomatic. The rate and duration of IV therapy are determined by the severity of the symptoms and the response of the serum calcium to treatment. An infusion of 0.5–2 mg/kg per hour or 30–100 mL/h of a 1-mg/mL solution usually suffices to relieve symptoms. Usually, parenteral therapy is required for only a few days. If symptoms worsen or if parenteral calcium is needed for >2–3 days, therapy with a vitamin D analogue and/or oral calcium (2–4 g/d) should be started (see below). It is cost-effective to use calcitriol (doses of 0.5–1 µg/d) because of the rapidity of onset of effect and prompt cessation of action when stopped, in comparison to other forms of vitamin D. A rise in blood calcium after several months of vitamin D replacement may indicate restoration of parathyroid function to normal. It is also appropriate to monitor serum PTH serially to estimate gland function in such patients.

If magnesium deficiency was present, it can complicate the postoperative course since magnesium deficiency impairs the secretion of PTH. Hypomagnesemia should be corrected whenever detected. Magnesium replacement can be effective orally (e.g., MgCl₂, MgOH₂), but parenteral repletion is usual to ensure postoperative recovery, if magnesium deficiency is suspected due to low blood magnesium levels. Because the depressant effect of magnesium on central and peripheral nerve functions does not occur at levels <2 mmol/L (normal range 0.8–1.2 mmol/L), parenteral replacement can be given rapidly. A cumulative dose as great as 0.5–1 mmol/kg of body weight can be administered if severe hypomagnesemia is present; often, however, total doses of 20–40 mmol are sufficient.

MEDICAL MANAGEMENT

The guidelines for recommending surgical intervention, if feasible (Table 424-2), as well as for monitoring patients with asymptomatic hyperparathyroidism who elect not to undergo parathyroidectomy

(Table 424-3), reflect the changes over time since the first conference on the topic in 1990. Medical monitoring rather than corrective surgery is still acceptable, but it is clear that surgical intervention is the more frequently recommended option for the reasons noted above. Tightened guidelines favoring surgery include lowering the recommended level of serum calcium elevation, more careful attention to skeletal integrity through reference to peak skeletal mass at baseline (T scores) rather than age-adjusted bone density (Z scores), as well as the presence of any fragility fracture. The other changes noted in the two guidelines (Tables 424-2 and 424-3) reflect accumulated experience and practical consideration, such as a difficulty in quantity of urine collections. Despite the usefulness of the guidelines, the importance of individual patient and physician judgment and preference is clear in all recommendations.

When surgery is not selected, or not medically feasible, there is interest in the potential value of specific medical therapies. There is no long-term experience regarding specific clinical outcomes such as fracture prevention, but it has been established that bisphosphonates increase bone mineral density significantly without changing serum calcium (as does estrogen, but the latter is not favored because of reported adverse effects in other organ systems). Calcimimetics that lower PTH secretion lower calcium but do not affect bone mineral density.

OTHER PARATHYROID-RELATED CAUSES OF HYPERCALCEMIA

Lithium Therapy Lithium, used in the management of bipolar depression and other psychiatric disorders, causes hypercalcemia in ~10% of treated patients. The hypercalcemia is dependent on continued lithium treatment, remitting and recurring when lithium is stopped and restarted. The parathyroid adenomas reported in some hypercalcemic patients with lithium therapy may reflect the presence of an independently occurring parathyroid tumor; a permanent effect of lithium on parathyroid gland growth need not be implicated as most patients have complete reversal of hypercalcemia when lithium is stopped. However, long-standing stimulation of parathyroid cell replication by lithium may predispose to development of adenomas (as is documented in secondary hyperparathyroidism and renal failure).

At the levels achieved in blood in treated patients, lithium can be shown *in vitro* to shift the PTH secretion curve to the right in response to calcium; i.e., higher calcium levels are required to lower PTH secretion, probably acting at the calcium sensor (see below). This effect can cause elevated PTH levels and consequent hypercalcemia in otherwise normal individuals. Fortunately, there are usually alternative medications for the underlying psychiatric illness. Parathyroid surgery should not be recommended unless hypercalcemia and elevated PTH levels persist after lithium is discontinued.

GENETIC DISORDERS CAUSING HYPERPARATHYROID-LIKE SYNDROMES

Familial Hypocalciuric Hypercalcemia FHH (also called *familial benign hypercalcemia*) is inherited as an autosomal dominant trait. Affected individuals are discovered because of asymptomatic hypercalcemia. Most cases of FHH (FHH1) are caused by an inactivating mutation in a single allele of the CaSR (see below), leading to inappropriately normal or even increased secretion of PTH, whereas another hypercalcemic disorder, namely the exceedingly rare Jansen's disease, is caused by a constitutively active PTH/PTHrP receptor in target tissues. Neither FHH1 nor Jansen's disease, however, is a growth disorder of the parathyroids. Other forms of FHH are caused either by heterozygous mutations in *GNA11* (encoding G11), one of the signaling proteins downstream of the CaSR (FHH2), or by mutations in *AP2S1* (FHH3).

The pathophysiology of FHH1 is now understood. The primary defect is abnormal sensing of the blood calcium by the parathyroid gland and renal tubule, causing inappropriate secretion of PTH and excessive reabsorption of calcium in the distal renal tubules. The CaSR is a member of the third family of GPCRs (type C or type III). The receptor responds to increased ECF calcium concentration by suppressing PTH secretion through second-messenger signaling involving the G protein alpha-subunits G11 and Gq, thereby providing

negative-feedback regulation of PTH secretion. Many different inactivating CaSR mutations have been identified in patients with FHH1. These mutations lower the capacity of the sensor to bind calcium, and the mutant receptors function as though blood calcium levels were low; excessive secretion of PTH occurs from an otherwise normal gland. Approximately two-thirds of patients with FHH have mutations within the protein-coding region of the CaSR gene. The remaining one-third of kindreds may have mutations in the promoter of the CaSR gene or are caused by mutations in other genes.

Even before elucidation of the pathophysiology of FHH, abundant clinical evidence served to separate the disorder from primary hyperparathyroidism; these clinical features are still useful in differential diagnosis. Patients with primary hyperparathyroidism have <99% renal calcium reabsorption, whereas most patients with FHH have >99% reabsorption. The hypercalcemia in FHH is often detectable in affected members of the kindreds in the first decade of life, whereas hypercalcemia rarely occurs in patients with primary hyperparathyroidism or the MEN syndromes who are age <10 years. PTH may be elevated in the different forms of FHH, but the values are usually normal or lower for the same degree of calcium elevation than is observed in patients with primary hyperparathyroidism. Parathyroid surgery performed in a few patients with FHH before the nature of the syndrome was understood led to permanent hypoparathyroidism; nevertheless, hypocalciuria persisted, establishing that hypocalciuria is not PTH-dependent (now known to be due to the abnormal CaSR in the kidney).

Few clinical signs or symptoms are present in patients with FHH, whereas other endocrine abnormalities are not. Most patients are detected as a result of family screening after hypercalcemia is detected in a proband. In those patients inadvertently operated upon for primary hyperparathyroidism, the parathyroids appeared normal or moderately hyperplastic. Parathyroid surgery is not appropriate, nor, in view of the lack of symptoms, does medical treatment seem needed to lower the calcium. One striking exception to the rule against parathyroid surgery in this syndrome is the occurrence, usually in consanguineous marriages (due to the rarity of the gene mutation), of a homozygous or compound heterozygote state, resulting in severe impairment of CaSR function. In this condition, neonatal severe hypercalcemia, total parathyroidectomy is mandatory, but calcimimetics have been used as a temporary measure. Rare but well-documented cases of acquired hypocalciuric hypercalcemia are reported due to antibodies against the CaSR. They appear to be a complication of an underlying autoimmune disorder and respond to therapies directed against the underlying disorder.

Jansen's Disease Activating mutations in the PTH/PTHrP receptor (PTH1R) have been identified as the cause of this rare autosomal dominant syndrome. Because the mutations lead to constitutive activation of receptor function, one abnormal copy of the mutant receptor is sufficient to cause the disease, thereby accounting for its dominant mode of transmission. The disorder leads to short-limbed dwarfism due to abnormal regulation of chondrocyte maturation in the growth plates of the bone that are formed through the endochondral process. In adult life, there are numerous abnormalities in bone, including multiple cystic resorptive areas resembling those seen in severe hyperparathyroidism. Hypercalcemia and hypophosphatemia with undetectable or low PTH levels are typically observed. The pathogenesis of the growth plate abnormalities in Jansen's disease has been confirmed by transgenic experiments in which targeted expression of the mutant PTH/PTHrP receptor to the proliferating chondrocyte layer of growth plate emulated several features of the human disorder. Some of these genetic mutations in the parathyroid gland or PTH target cells that affect Ca²⁺ metabolism are illustrated in Fig. 424-5.

MALIGNANCY-RELATED HYPERCALCEMIA

Clinical Syndromes and Mechanisms of Hypercalcemia Hypercalcemia due to malignancy is common (occurring in as many as 20% of cancer patients, especially with certain types of tumor such as lung carcinoma), often severe and difficult to manage, and, on rare occasions,

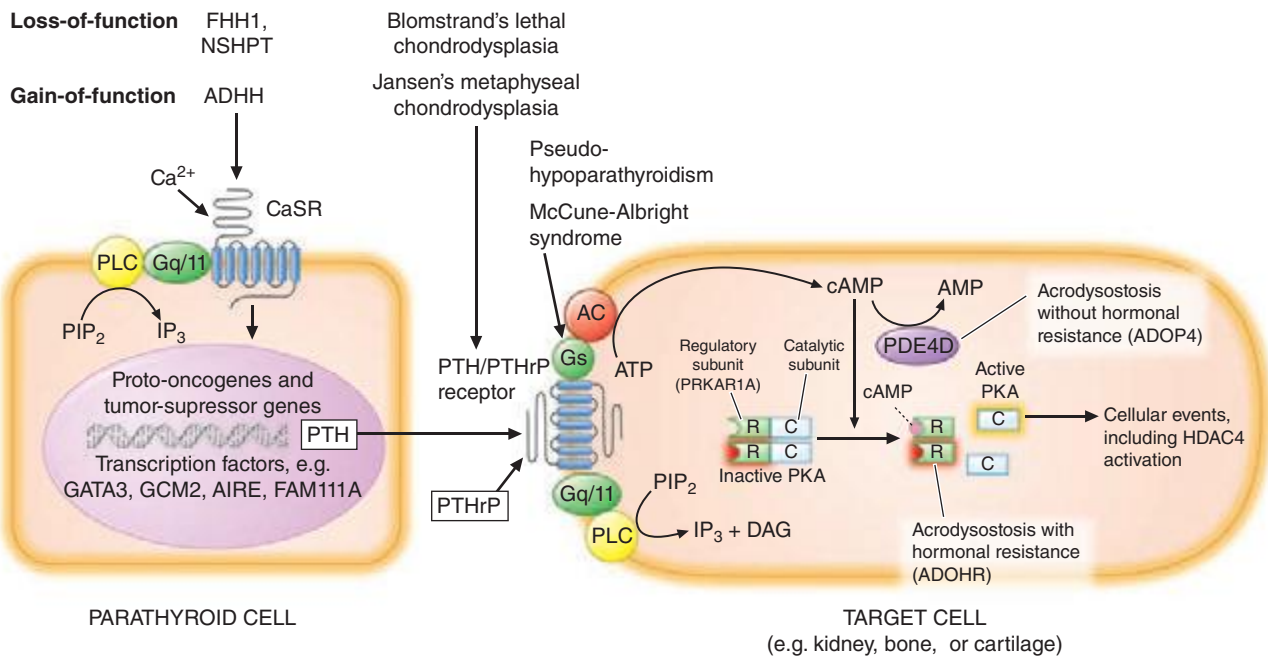


FIGURE 424-5 Illustration of some genetic mutations that alter calcium metabolism by effects on the parathyroid cell or target cells of parathyroid hormone (PTH) action. Alterations in PTH production by the parathyroid cell can be caused by changes in the response to extracellular fluid calcium (Ca^{2+}) that are detected by the calcium-sensing receptor (CaSR). Furthermore, PTH (or PTH-related peptide [PTHrP]) can show altered efficacy in target cells such as in proximal tubular cells, by altered function of its receptor (PTH/PTHrP receptor) or the signal transduction proteins, G proteins such as $\text{G}_s\alpha$, which is linked to adenylate cyclase (AC), the enzyme responsible for producing cyclic AMP (cAMP) (also illustrated are Gq/11, which activate an alternate pathway of receptor signal transmission involving the generation of inositol triphosphate [IP_3] or diacylglycerol [DAG]). Heterozygous loss-of-function mutations in the CaSR cause familial benign hypocalciuric hypercalcemia (FBHH), homozygous mutations (both alleles mutated), and severe neonatal hyperparathyroidism (NSHPT); heterozygous gain-of-function causes autosomal dominant hypercalciuric hypocalcemia (ADHH). Other defects in parathyroid cell function that occur at the level of gene regulation (oncogenes or tumor-suppressor genes) or transcription factors are discussed in the text. Blomstrand's lethal chondrodysplasia is due to homozygous or compound heterozygous loss-of-function mutations in the PTH/PTHrP receptor, a neonatally lethal disorder, while pseudohypoparathyroidism involves inactivation at the level of the G proteins, specifically mutations that eliminate or reduce $\text{G}_s\alpha$ activity in the kidney (see text for details). Acrodysostosis can occur with (acrodysostosis with hormonal resistance [ADOHR]; mutant regulatory subunit of PKA) or without hormonal resistance (ADOP4; mutant PDE4D). Jansen's metaphyseal chondrodysplasia and McCune-Albright syndrome represent gain-of-function mutations in the PTH/PTHrP receptor and $\text{G}_s\alpha$ protein, respectively.

difficult to distinguish from primary hyperparathyroidism. Although malignancy is often clinically obvious or readily detectable by medical history, hypercalcemia can occasionally be due to an occult tumor. Previously, hypercalcemia associated with malignancy was thought to be due to local invasion and destruction of bone by tumor cells; many cases are now known to result from the elaboration by the malignant cells of humoral mediators of hypercalcemia. PTHrP is the responsible humoral agent in most solid tumors that cause hypercalcemia.

The histologic character of the tumor is more important than the extent of skeletal metastases in predicting hypercalcemia. Small-cell carcinoma (oat cell) and adenocarcinoma of the lung, although the most common lung tumors associated with skeletal metastases, rarely cause hypercalcemia. By contrast, many patients with squamous cell carcinoma of the lung develop hypercalcemia. Histologic studies of bone in patients with squamous cell or epidermoid carcinoma of the lung, in sites invaded by tumor as well as areas remote from tumor invasion, reveal increased bone resorption.

Two main mechanisms of hypercalcemia are operative in cancer hypercalcemia. Many solid tumors associated with hypercalcemia, particularly squamous cell and renal tumors, produce and secrete PTHrP that causes increased bone resorption and mediate the hypercalcemia through systemic actions on the skeleton. Alternatively, direct bone marrow invasion occurs with hematologic malignancies such as leukemia, lymphoma, and multiple myeloma. Lymphokines and cytokines (including PTHrP) produced by cells involved in the marrow response to the tumors promote resorption of bone through local destruction. Several hormones, hormone analogues, cytokines, and growth factors have been implicated as the result of clinical assays, in vitro tests, or chemical isolation. The etiologic factor

produced by activated normal lymphocytes and by myeloma and lymphoma cells, originally termed *osteoclast activation factor*, now appears to represent the biologic action of several different cytokines, probably interleukin 1 and lymphotoxin or tumor necrosis factor (TNF). In some lymphomas, there is a third mechanism, caused by an increased blood level of $1,25(\text{OH})_2\text{D}$, produced by the abnormal lymphocytes.

In the more common mechanism, usually termed *humoral hypercalcemia of malignancy*, solid tumors (cancers of the lung and kidney, in particular), in which bone metastases are absent, minimal, or not detectable clinically, secrete PTHrP measurable by immunoassay. Secretion by the tumors of the PTH-like factor, PTHrP, activates the PTH1R, resulting in a pathophysiology closely resembling hyperparathyroidism, but with normal or suppressed PTH levels. The clinical picture resembles primary hyperparathyroidism (hypophosphatemia accompanies hypercalcemia), and elimination or regression of the primary tumor leads to disappearance of the hypercalcemia.

As in hyperparathyroidism, patients with the humoral hypercalcemia of malignancy have elevated urinary nephrogenous cAMP excretion, hypophosphatemia, and increased urinary phosphate clearance. However, in humoral hypercalcemia of malignancy, immunoreactive PTH is undetectable or suppressed, making the differential diagnosis easier. Other features of the disorder differ from those of true hyperparathyroidism. Although the biologic actions of PTH and PTHrP are exerted through the same receptor, subtle differences in receptor activation by the two ligands must account for some of the discordance in pathophysiology, when an excess of one or the other peptide occurs. Other cytokines elaborated by the malignancy may contribute to the variations from hyperparathyroidism in these patients as well. Patients

with humoral hypercalcemia of malignancy may have low to normal levels of $1,25(\text{OH})_2\text{D}$ instead of elevated levels as in true hyperparathyroidism. In some patients with the humoral hypercalcemia of malignancy, osteoclastic resorption is unaccompanied by an osteoblastic or bone-forming response, implying inhibition of the normal coupling of bone formation and resorption.

Several different assays (single- or double-antibody, different epitopes) have been developed to detect PTHrP. Most data indicate that circulating PTHrP levels are undetectable (or low) in normal individuals except perhaps in pregnancy (high in human milk) and elevated in most cancer patients with the humoral syndrome. The etiologic mechanisms in cancer hypercalcemia may be multiple in the same patient. For example, in breast carcinoma (metastatic to bone) and in a distinctive type of T cell lymphoma/leukemia initiated by human T cell lymphotropic virus I, hypercalcemia is caused by direct local lysis of bone as well as by a humoral mechanism involving excess production of PTHrP. Hyperparathyroidism has been reported to coexist with the humoral cancer syndrome, and rarely, ectopic hyperparathyroidism due to tumor elaboration of true PTH is reported.

Diagnostic Issues Levels of PTH measured by the double-antibody technique are undetectable or extremely low in tumor hypercalcemia, as would be expected with the mediation of the hypercalcemia by a factor other than PTH (the hypercalcemia suppresses the normal parathyroid glands). In a patient with minimal symptoms referred for hypercalcemia, low or undetectable PTH levels would focus attention on a possible occult malignancy (except for very rare cases of ectopic hyperparathyroidism).

Ordinarily, the diagnosis of cancer hypercalcemia is not difficult because tumor symptoms are prominent when hypercalcemia is detected. Indeed, hypercalcemia may be noted incidentally during the workup of a patient with known or suspected malignancy. Clinical suspicion that malignancy is the cause of the hypercalcemia is heightened when there are other signs or symptoms of a paraneoplastic process such as weight loss, fatigue, muscle weakness, or unexplained skin rash, or when symptoms specific for a particular tumor are present. Squamous cell tumors are most frequently associated with hypercalcemia, particularly tumors of the lung, kidney, head and neck, and urogenital tract. Radiologic examinations can focus on these areas when clinical evidence is unclear. Bone scans with technetium-labeled bisphosphonate are useful for detection of osteolytic metastases; the sensitivity is high, but specificity is low; results must be confirmed by conventional x-rays to be certain that areas of increased uptake are due to osteolytic metastases per se. Bone marrow biopsies are helpful in patients with anemia or abnormal peripheral blood smears.

TREATMENT MALIGNANCY-RELATED HYPERCALCEMIA

Treatment of the hypercalcemia of malignancy is first directed to control of tumor; reduction of tumor mass usually corrects hypercalcemia. If a patient has severe hypercalcemia yet has a good chance for effective tumor therapy, treatment of the hypercalcemia should be vigorous while awaiting the results of definitive therapy. If hypercalcemia occurs in the late stages of a tumor that is resistant to antitumor therapy, the treatment of the hypercalcemia should be judicious as high calcium levels can have a mild sedating effect. Standard therapies for hypercalcemia (discussed below) are applicable to patients with malignancy.

VITAMIN D-RELATED HYPERCALCEMIA

Hypercalcemia caused by vitamin D can be due to excessive ingestion or abnormal metabolism of the vitamin. Abnormal metabolism of the vitamin is usually acquired in association with a widespread granulomatous disorder. Vitamin D metabolism is carefully regulated, particularly the activity of renal 1α -hydroxylase, the enzyme responsible for the production of $1,25(\text{OH})_2\text{D}$ (Chap. 423). The regulation of

1α -hydroxylase and the normal feedback suppression by $1,25(\text{OH})_2\text{D}$ seem to work less well in infants than in adults and to operate poorly, if at all, in sites other than the renal tubule; these phenomena may explain the occurrence of hypercalcemia secondary to excessive $1,25(\text{OH})_2\text{D}$ production in infants with Williams' syndrome (see below) and in adults with sarcoidosis or lymphoma.

Vitamin D Intoxication Chronic ingestion of 40–100 times the normal physiologic requirement of vitamin D (amounts >40,000–100,000 U/d) is usually required to produce significant hypercalcemia in otherwise healthy individuals. The stated upper limit of safe dietary intake is 2000 U/d (50 $\mu\text{g}/\text{d}$) in adults because of concerns about potential toxic effects of cumulative supraphysiologic doses. These recommendations are now regarded as too restrictive, because some estimates are that in elderly individuals in northern latitudes, 2000 U/d or more may be necessary to avoid vitamin D insufficiency.

Hypercalcemia in vitamin D intoxication is due to an excessive biologic action of the vitamin, perhaps the consequence of increased levels of $25(\text{OH})\text{D}$ rather than merely increased levels of the active metabolite $1,25(\text{OH})_2\text{D}$ (the latter may not be elevated in vitamin D intoxication). $25(\text{OH})\text{D}$ has definite, if low, biologic activity in the intestine and bone. The production of $25(\text{OH})\text{D}$ is less tightly regulated than is the production of $1,25(\text{OH})_2\text{D}$. Hence concentrations of $25(\text{OH})\text{D}$ are elevated several-fold in patients with excess vitamin D intake.

The diagnosis is substantiated by documenting elevated levels of $25(\text{OH})\text{D}$ >100 mg/mL. Hypercalcemia is usually controlled by restriction of dietary calcium intake and appropriate attention to hydration. These measures, plus discontinuation of vitamin D, usually lead to resolution of hypercalcemia. However, vitamin D stores in fat may be substantial, and vitamin D intoxication may persist for weeks after vitamin D ingestion is terminated. Such patients are responsive to glucocorticoids, which in doses of 100 mg/d of hydrocortisone or its equivalent usually return serum calcium levels to normal over several days; severe intoxication may require intensive therapy.

Sarcoidosis and Other Granulomatous Diseases In patients with sarcoidosis and other granulomatous diseases, such as tuberculosis and fungal infections, excess $1,25(\text{OH})_2\text{D}$ is synthesized in macrophages or other cells in the granulomas. Indeed, increased $1,25(\text{OH})_2\text{D}$ levels have been reported in anephric patients with sarcoidosis and hypercalcemia. Macrophages obtained from granulomatous tissue convert $25(\text{OH})\text{D}$ to $1,25(\text{OH})_2\text{D}$ at an increased rate. There is a positive correlation in patients with sarcoidosis between $25(\text{OH})\text{D}$ levels (reflecting vitamin D intake) and the circulating concentrations of $1,25(\text{OH})_2\text{D}$, whereas normally there is no increase in $1,25(\text{OH})_2\text{D}$ with increasing $25(\text{OH})\text{D}$ levels due to multiple feedback controls on renal 1α -hydroxylase (Chap. 423). The usual regulation of active metabolite production by calcium and phosphate or by PTH does not operate in these patients. Clearance of $1,25(\text{OH})_2\text{D}$ from blood may be decreased in sarcoidosis as well. PTH levels are usually low and $1,25(\text{OH})_2\text{D}$ levels are elevated, but primary hyperparathyroidism and sarcoidosis may coexist in some patients.

Management of the hypercalcemia can often be accomplished by avoiding excessive sunlight exposure and limiting vitamin D and calcium intake. Presumably, however, the abnormal sensitivity to vitamin D and abnormal regulation of $1,25(\text{OH})_2\text{D}$ synthesis will persist as long as the disease is active. Alternatively, glucocorticoids in the equivalent of 100 mg/d of hydrocortisone or equivalent doses of glucocorticoids may help control hypercalcemia. Glucocorticoids appear to act by blocking excessive production of $1,25(\text{OH})_2\text{D}$, as well as the response to it in target organs.

Idiopathic Hypercalcemia of Infancy This rare disorder, usually referred to as *Williams' syndrome*, is an autosomal dominant disorder characterized by multiple congenital development defects, including supravalvular aortic stenosis, mental retardation, and an elfin facies, in association with hypercalcemia due to abnormal sensitivity to vitamin D. The hypercalcemia associated with the syndrome was first recognized in England after fortification of milk with vitamin D. The cardiac and developmental abnormalities were independently described,

2478 but the connection between these defects and hypercalcemia were not described until later. Levels of $1,25(\text{OH})_2\text{D}$ can be elevated, ranging from 46 to 120 nmol/L (150–500 pg/mL). The mechanism of the abnormal sensitivity to vitamin D and of the increased circulating levels of $1,25(\text{OH})_2\text{D}$ is still unclear. Studies suggest that genetic mutations involving microdeletions at the elastin locus and perhaps other genes on chromosome 7 may play a role in the pathogenesis. Another cause of hypercalcemia in infants and young children is a 24-hydroxylase deficiency that impairs metabolism of $1,25(\text{OH})_2\text{D}$.

HYPERCALCEMIA ASSOCIATED WITH HIGH BONE TURNOVER

Hyperthyroidism As many as 20% of hyperthyroid patients have high-normal or mildly elevated serum calcium concentrations; hypercalciuria is even more common. The hypercalcemia is due to increased bone turnover, with bone resorption exceeding bone formation. Severe calcium elevations are not typical, and the presence of such suggests a concomitant disease such as hyperparathyroidism. Usually, the diagnosis is obvious, but signs of hyperthyroidism may occasionally be occult, particularly in the elderly (**Chap. 405**). Hypercalcemia is managed by treatment of the hyperthyroidism. Reports that thyroid-stimulating hormone (TSH) itself normally has a bone-protective effect suggest that suppressed TSH levels also play a role in hypercalcemia.

Immobilization Immobilization is a rare cause of hypercalcemia in adults in the absence of an associated disease but may cause hypercalcemia in children and adolescents, particularly after spinal cord injury and paraplegia or quadriplegia. With resumption of ambulation, the hypercalcemia in children usually returns to normal.

The mechanism appears to involve a disproportion between bone formation and bone resorption; the former decreased and the latter increased. Hypercalciuria and increased mobilization of skeletal calcium can develop in normal volunteers subjected to extensive bed rest, although hypercalcemia is unusual. Immobilization of an adult with a disease associated with high bone turnover, however, such as Paget's disease, may cause hypercalcemia.

Thiazides Administration of benzothiadiazines (thiazides) can cause hypercalcemia in patients with high rates of bone turnover. Traditionally, thiazides are associated with aggravation of hypercalcemia in primary hyperparathyroidism, but this effect can be seen in other high-bone-turnover states as well. The mechanism of thiazide action is complex. Chronic thiazide administration leads to reduction in urinary calcium; the hypocalciuric effect appears to reflect the enhancement of proximal tubular resorption of sodium and calcium in response to sodium depletion. Some of this renal effect is due to augmentation of PTH action and is more pronounced in individuals with intact PTH secretion. However, thiazides cause hypocalciuria in hypoparathyroid patients on high-dose vitamin D and oral calcium replacement if sodium intake is restricted. This finding is the rationale for the use of thiazides as an adjunct to therapy in hypoparathyroid patients, as discussed below. Thiazide administration to normal individuals causes a transient increase in blood calcium (usually within the high-normal range) that reverts to preexisting levels after a week or more of continued administration. If hormonal function and calcium and bone metabolism are normal, homeostatic controls are reset to counteract the calcium-elevating effect of the thiazides. In the presence of hyperparathyroidism or increased bone turnover from another cause, homeostatic mechanisms are ineffective. The abnormal effects of the thiazide on calcium metabolism disappear within days of cessation of the drug.

Vitamin A Intoxication Vitamin A intoxication is a rare cause of hypercalcemia and is most commonly a side effect of dietary faddism (**Chap. 96e**). Calcium levels can be elevated into the 3–3.5-mmol/L (12–14 mg/dL) range after the ingestion of 50,000–100,000 units of vitamin A daily (10–20 times the minimum daily requirement). Typical features of severe hypercalcemia include fatigue, anorexia, and, in some, severe muscle and bone pain. Excess vitamin A intake is presumed to increase bone resorption.

The diagnosis can be established by history and by measurement of vitamin A levels in serum. Occasionally, skeletal x-rays reveal periosteal

calcifications, particularly in the hands. Withdrawal of the vitamin is usually associated with prompt disappearance of the hypercalcemia and reversal of the skeletal changes. As in vitamin D intoxication, administration of 100 mg/d of hydrocortisone or its equivalent leads to a rapid return of the serum calcium to normal.

HYPERCALCEMIA ASSOCIATED WITH RENAL FAILURE

Severe Secondary Hyperparathyroidism The pathogenesis of secondary hyperparathyroidism in chronic kidney disease is incompletely understood. Resistance to the normal level of PTH is a major factor contributing to the development of hypocalcemia, which, in turn, is a stimulus to parathyroid gland enlargement. However, recent findings have indicated that an increase of FGF23 production by osteocytes (and possibly osteoblasts) in bone occurs well before an elevation in PTH is detected. FGF23 is a potent inhibitor of the renal 1- α -hydroxylase, and the FGF23-dependent reduction in $1,25(\text{OH})_2$ vitamin D seems to be an important stimulus for the development of secondary hyperparathyroidism.

Secondary hyperparathyroidism occurs not only in patients with renal failure but also in those with osteomalacia due to multiple causes (**Chap. 423**), including deficiency of vitamin D action and pseudohypoparathyroidism (deficient response to PTH downstream of PTHR1). For both disorders, hypocalcemia seems to be the common denominator in initiating the development of secondary hyperparathyroidism. Primary (1°) and secondary (2°) hyperparathyroidism can be distinguished conceptually by the autonomous growth of the parathyroid glands in primary hyperparathyroidism (presumably irreversible) and the adaptive response of the parathyroids in secondary hyperparathyroidism (typically reversible). In fact, reversal over weeks from an abnormal pattern of secretion, presumably accompanied by involution of parathyroid gland mass to normal, occurs in patients with osteomalacia who have been treated effectively with calcium and vitamin D. However, it is now recognized that a true clonal outgrowth (irreversible) can arise in long-standing, inadequately treated chronic kidney disease (e.g., tertiary [3°] hyperparathyroidism; see below).

Patients with secondary hyperparathyroidism may develop bone pain, ectopic calcification, and pruritus. The bone disease seen in patients with secondary hyperparathyroidism and chronic kidney disease is termed *renal osteodystrophy* and affects primarily bone turnover. However, osteomalacia is frequently encountered as well and may be related to the circulating levels of FGF23.

Two other skeletal disorders have been frequently associated in the past with chronic kidney disease (CKD) patients treated by long-term dialysis, who received aluminum-containing phosphate binders. Aluminum deposition in bone (see below) leads to an osteomalacia-like picture. The other entity is a low-turnover bone disease termed “aplastic” or “adynamic” bone disease; PTH levels are lower than typically observed in CKD patients with secondary hyperparathyroidism. It is believed that the condition is caused, at least in part, by excessive PTH suppression, which may be even greater than previously appreciated in light of evidence that some of the immunoreactive PTH detected by most commercially available PTH assays is not the full-length biologically active molecule (as discussed above) but may consist of amino-terminally truncated fragments that do not activate the PTH1R.

TREATMENT SECONDARY HYPERPARATHYROIDISM

Medical therapy to reverse secondary hyperparathyroidism in CKD includes reduction of excessive blood phosphate by restriction of dietary phosphate, the use of nonabsorbable phosphate binders, and careful, selective addition of calcitriol (0.25–2 $\mu\text{g}/\text{d}$) or related analogues. Calcium carbonate became preferred over aluminum-containing antacids to prevent aluminum-induced bone disease. However, synthetic gels that also bind phosphate (such as sevelamer; **Chap. 335**) are now widely used, with the advantage of avoiding not only aluminum retention, but also excess calcium

loading, which may contribute to cardiovascular calcifications. Intravenous calcitriol (or related analogues), administered as several pulses each week, helps control secondary hyperparathyroidism. Aggressive but carefully administered medical therapy can often, but not always, reverse hyperparathyroidism and its symptoms and manifestations.

Occasional patients develop severe manifestations of secondary hyperparathyroidism, including hypercalcemia, pruritus, extraskel-etal calcifications, and painful bones, despite aggressive medical efforts to suppress the hyperparathyroidism. PTH hypersecretion no longer responsive to medical therapy, a state of severe hyperparathyroidism in patients with CKD that requires surgery, has been referred to as *tertiary hyperparathyroidism*. Parathyroid surgery is necessary to control this condition. Based on genetic evidence from examination of tumor samples in these patients, the emergence of autonomous parathyroid function is due to a monoclonal outgrowth of one or more previously hyperplastic parathyroid glands. The adaptive response has become an independent contributor to disease; this finding seems to emphasize the importance of optimal medical management to reduce the proliferative response of the parathyroid cells that enables the irreversible genetic change.

Aluminum Intoxication Aluminum intoxication (and often hypercalcemia as a complication of medical treatment) in the past occurred in patients on chronic dialysis; manifestations included acute dementia and unresponsive and severe osteomalacia. Bone pain, multiple non-healing fractures, particularly of the ribs and pelvis, and a proximal myopathy occur. Hypercalcemia develops when these patients are treated with vitamin D or calcitriol because of impaired skeletal responsiveness. Aluminum is present at the site of osteoid mineralization, osteoblastic activity is minimal, and calcium incorporation into the skeleton is impaired. The disorder is now rare because of the avoidance of aluminum-containing antacids or aluminum excess in the dialysis regimen (Chap. 429).

Milk-Alkali Syndrome The milk-alkali syndrome is due to excessive ingestion of calcium and absorbable antacids such as milk or calcium carbonate. It is much less frequent since proton pump inhibitors and other treatments became available for peptic ulcer disease. For a time, the increased use of calcium carbonate in the management of secondary hyperparathyroidism led to reappearance of the syndrome.

Several clinical presentations—acute, subacute, and chronic—have been described, all of which feature hypercalcemia, alkalosis, and renal failure. The chronic form of the disease, termed *Burnett's syndrome*, is associated with irreversible renal damage. The acute syndromes reverse if the excess calcium and absorbable alkali are stopped.

Individual susceptibility is important in the pathogenesis, because some patients are treated with calcium carbonate and alkali regimens without developing the syndrome. One variable is the fractional calcium absorption as a function of calcium intake. Some individuals absorb a high fraction of calcium, even with intakes ≥ 2 g of elemental calcium per day, instead of reducing calcium absorption with high intake, as occurs in most normal individuals. Resultant mild hypercalcemia after meals in such patients is postulated to contribute to the generation of alkalosis. Development of hypercalcemia causes increased sodium excretion and some depletion of total-body water. These phenomena and perhaps some suppression of endogenous PTH secretion due to mild hypercalcemia lead to increased bicarbonate resorption and to alkalosis in the face of continued calcium carbonate ingestion. Alkalosis per se selectively enhances calcium resorption in the distal nephron, thus aggravating the hypercalcemia. The cycle of mild hypercalcemia \rightarrow bicarbonate retention \rightarrow alkalosis \rightarrow renal calcium retention \rightarrow severe hypercalcemia perpetuates and aggravates hypercalcemia and alkalosis as long as calcium and absorbable alkali are ingested.

DIFFERENTIAL DIAGNOSIS: SPECIAL TESTS

Differential diagnosis of hypercalcemia is best achieved by using clinical criteria, but immunometric assays to measure PTH are especially useful in distinguishing among major causes (Fig. 424-6). The clinical features that deserve emphasis are the presence or absence of symptoms or signs of disease and evidence of chronicity. If one discounts fatigue or depression, $>90\%$ of patients with primary hyperparathyroidism have *asymptomatic hypercalcemia*; symptoms of malignancy are usually present in cancer-associated hypercalcemia. Disorders other than hyperparathyroidism and malignancy cause $<10\%$ of cases of hypercalcemia, and some of the nonparathyroid causes are associated with clear-cut manifestations such as renal failure.

Hyperparathyroidism is the likely diagnosis in patients with *chronic hypercalcemia*. If hypercalcemia has been manifest for >1 year, malignancy can usually be excluded as the cause. A striking feature of malignancy-associated hypercalcemia is the rapidity of the course,

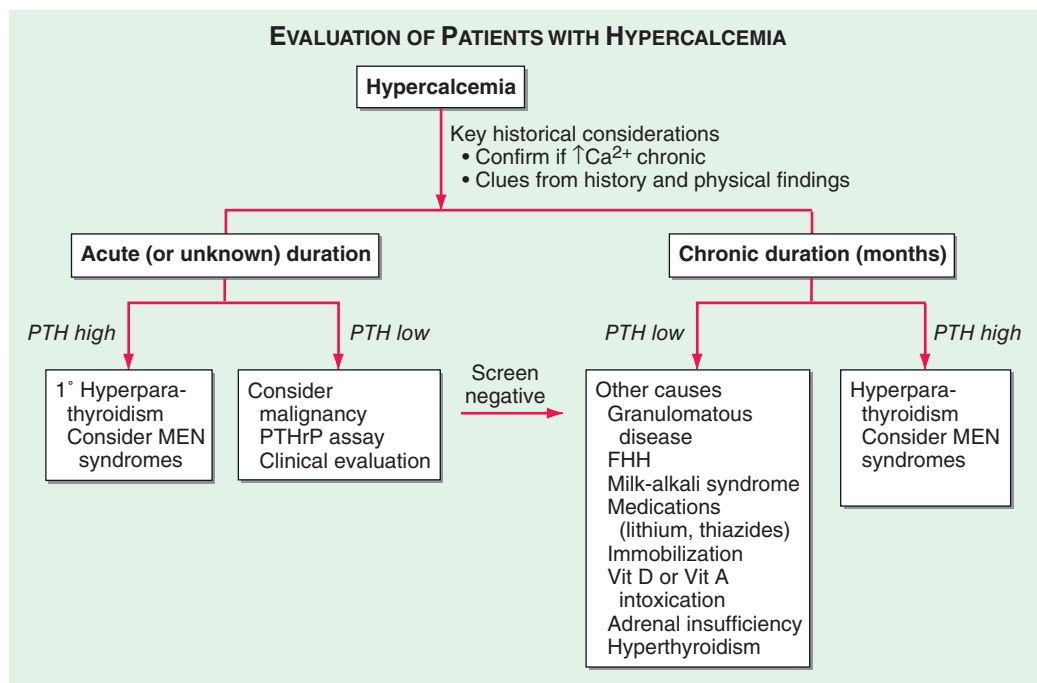


FIGURE 424-6 Algorithm for the evaluation of patients with hypercalcemia. See text for details. FHH, familial hypocalciuric hypercalcemia; MEN, multiple endocrine neoplasia; PTH, parathyroid hormone; PThrP, parathyroid hormone–related peptide.

2480 whereby signs and symptoms of the underlying malignancy are evident within months of the detection of hypercalcemia. Although clinical considerations are helpful in arriving at the correct diagnosis of the cause of hypercalcemia, appropriate laboratory testing is essential for definitive diagnosis. The immunoassay for PTH usually separates hyperparathyroidism from all other causes of hypercalcemia (exceptions are very rare reports of ectopic production of excess PTH by nonparathyroid tumors). Patients with hyperparathyroidism have elevated PTH levels despite hypercalcemia, whereas patients with malignancy and the other causes of hypercalcemia (except for disorders mediated by PTH such as lithium-induced hypercalcemia) have levels of hormone below normal or undetectable levels. Assays based on the double-antibody method for PTH exhibit very high sensitivity (especially if serum calcium is simultaneously evaluated) and specificity for the diagnosis of primary hyperparathyroidism (Fig. 424-4).

In summary, PTH values are elevated in >90% of parathyroid-related causes of hypercalcemia, undetectable or low in malignancy-related hypercalcemia, and undetectable or normal in vitamin D-related and high-bone-turnover causes of hypercalcemia. In view of the specificity of the PTH immunoassay and the high frequency of hyperparathyroidism in hypercalcemic patients, it is cost-effective to measure the PTH level in all hypercalcemic patients unless malignancy or a specific nonparathyroid disease is obvious. False-positive PTH assay results are rare. Immunoassays for PTHrP are helpful in diagnosing certain types of malignancy-associated hypercalcemia. Although FHH is parathyroid-related, the disease should be managed distinctively from hyperparathyroidism. Clinical features and the low urinary calcium excretion can help make the distinction. Because the incidence of malignancy and hyperparathyroidism both increase with age, they can coexist as two independent causes of hypercalcemia.

1,25(OH)₂D levels are elevated in many (but not all) patients with primary hyperparathyroidism. In other disorders associated with hypercalcemia, concentrations of 1,25(OH)₂D are low or, at the most, normal. However, this test is of low specificity and is not cost-effective, as not all patients with hyperparathyroidism have elevated 1,25(OH)₂D levels and not all nonparathyroid hypercalcemic patients have suppressed 1,25(OH)₂D. Measurement of 1,25(OH)₂D is, however, critically valuable in establishing the cause of hypercalcemia in sarcoidosis and certain lymphomas.

A useful general approach is outlined in Fig. 424-6. If the patient is *asymptomatic* and there is evidence of *chronicity* to the hypercalcemia,

hyperparathyroidism is almost certainly the cause. If PTH levels (usually measured at least twice) are elevated, the clinical impression is confirmed and little additional evaluation is necessary. If there is only a short history or no data as to the duration of the hypercalcemia, *occult malignancy* must be considered; if the PTH levels are not elevated, then a thorough workup must be undertaken for malignancy, including chest x-ray, CT of chest and abdomen, and bone scan. Immunoassays for PTHrP may be especially useful in such situations. Attention should also be paid to clues for underlying hematologic disorders such as anemia, increased plasma globulin, and abnormal serum immunoelectrophoresis; bone scans can be negative in some patients with metastases such as in multiple myeloma. Finally, if a patient with chronic hypercalcemia is asymptomatic and malignancy therefore seems unlikely on clinical grounds, but PTH values are not elevated, it is useful to search for other chronic causes of hypercalcemia such as occult sarcoidosis. A careful history of dietary supplements and drug use may suggest intoxication with vitamin D or vitamin A or the use of thiazides.

TREATMENT HYPERCALCEMIC STATES

The approach to medical treatment of hypercalcemia varies with its severity (Table 424-4). Mild hypercalcemia, <3 mmol/L (12 mg/dL), can be managed by hydration. More severe hypercalcemia (levels of 3.2–3.7 mmol/L [13–15 mg/dL]) must be managed aggressively; above that level, hypercalcemia can be life-threatening and requires emergency measures. By using a combination of approaches in severe hypercalcemia, the serum calcium concentration can be decreased by 0.7–2.2 mmol/L (3–9 mg/dL) within 24–48 h in most patients, enough to relieve acute symptoms, prevent death from hypercalcemic crisis, and permit diagnostic evaluation. Therapy can then be directed at the underlying disorder—the second priority.

Hypercalcemia develops because of excessive skeletal calcium release, increased intestinal calcium absorption, or inadequate renal calcium excretion. Understanding the particular pathogenesis helps guide therapy. For example, hypercalcemia in patients with malignancy is primarily due to excessive skeletal calcium release and is, therefore, minimally improved by restriction of dietary calcium. On the other hand, patients with vitamin D hypersensitivity or vitamin D intoxication have excessive intestinal calcium absorption, and

TABLE 424-4 THERAPIES FOR SEVERE HYPERCALCEMIA

Treatment	Onset of Action	Duration of Action	Advantages	Disadvantages
Most Useful Therapies				
Hydration with saline	Hours	During infusion	Rehydration invariably needed	Volume overload
Forced diuresis; saline plus loop diuretic	Hours	During treatment	Rapid action	Volume overload, cardiac decompensation, intensive monitoring, electrolyte disturbance, inconvenience
Pamidronate	1–2 days	10–14 days to weeks	High potency; intermediate onset of action	Fever in 20%, hypophosphatemia, hypocalcemia, hypomagnesemia, rarely jaw necrosis
Zoledronate	1–2 days	>3 weeks	Same as for pamidronate (may last longer)	Same as pamidronate above
Calcitonin	Hours	1–2 days	Rapid onset of action; useful as adjunct in severe hypercalcemia	Rapid tachyphylaxis
Special Use Therapies				
Phosphate oral	24 h	During use	Chronic management (with hypophosphatemia); low toxicity if phosphate <4 mg/dL	Limited use except as adjuvant or chronic therapy
Glucocorticoids	Days	Days, weeks	Oral therapy, antitumor agent	Active only in certain malignancies, vitamin D excess and sarcoidosis; glucocorticoid side effects
Dialysis	Hours	During use and 24–48 h afterward	Useful in renal failure; onset of effect in hours; can immediately reverse life-threatening hypercalcemia	Complex procedure, reserved for extreme or special circumstances

restriction of dietary calcium is beneficial. Decreased renal function or ECF depletion decreases urinary calcium excretion. In such situations, rehydration may rapidly reduce or reverse the hypercalcemia, even though increased bone resorption persists. As outlined below, the more severe the hypercalcemia, the greater the number of combined therapies that should be used. Rapid-acting (hours) approaches—rehydration, forced diuresis, and calcitonin—can be used with the most effective antiresorptive agents such as bisphosphonates (since severe hypercalcemia usually involves excessive bone resorption).

HYDRATION, INCREASED SALT INTAKE, MILD AND FORCED DIURESIS

The first principle of treatment is to restore normal hydration. Many hypercalcemic patients are dehydrated because of vomiting, inanition, and/or hypercalcemia-induced defects in urinary concentrating ability. The resultant drop in glomerular filtration rate is accompanied by an additional decrease in renal tubular sodium and calcium clearance. Restoring a normal ECF volume corrects these abnormalities and increases urine calcium excretion by 2.5–7.5 mmol/d (100–300 mg/d). Increasing urinary sodium excretion to 400–500 mmol/d increases urinary calcium excretion even further than simple rehydration. After rehydration has been achieved, saline can be administered, or furosemide or ethacrynic acid can be given twice daily to depress the tubular reabsorptive mechanism for calcium (care must be taken to prevent dehydration). The combined use of these therapies can increase urinary calcium excretion to ≥ 12.5 mmol/d (500 mg/d) in most hypercalcemic patients. Because this is a substantial percentage of the exchangeable calcium pool, the serum calcium concentration usually falls 0.25–0.75 mmol/L (1–3 mg/dL) within 24 h. Precautions should be taken to prevent potassium and magnesium depletion; calcium-containing renal calculi are a potential complication.

Under life-threatening circumstances, the preceding approach can be pursued more aggressively, but the availability of effective agents to block bone resorption (such as bisphosphonates) has reduced the need for extreme diuresis regimens (Table 424-4). Depletion of potassium and magnesium is inevitable unless replacements are given; pulmonary edema can be precipitated. The potential complications can be reduced by careful monitoring of central venous pressure and plasma or urine electrolytes; catheterization of the bladder may be necessary. Dialysis treatment may be needed when renal function is compromised.

BISPHOSPHONATES

The bisphosphonates are analogues of pyrophosphate, with high affinity for bone, especially in areas of increased bone turnover, where they are powerful inhibitors of bone resorption. These bone-seeking compounds are stable in vivo because phosphatase enzymes cannot hydrolyze the central carbon-phosphorus-carbon bond. The bisphosphonates are concentrated in areas of high bone turnover and are taken up by and inhibit osteoclast action; the mechanism of action is complex. The bisphosphonate molecules that contain amino groups in the side chain structure (see below) interfere with prenylation of proteins and can lead to cellular apoptosis. The highly active nonamino group-containing bisphosphonates are also metabolized to cytotoxic products.

The initial bisphosphonate widely used in clinical practice, etidronate, was effective but had several disadvantages, including the capacity to inhibit bone formation as well as blocking resorption. Subsequently, a number of second- or third-generation compounds have become the mainstays of antiresorptive therapy for treatment of hypercalcemia and osteoporosis. The newer bisphosphonates have a highly favorable ratio of blocking resorption versus inhibiting bone formation; they inhibit osteoclast-mediated skeletal resorption yet do not cause mineralization defects at ordinary doses. Although the bisphosphonates have similar structures, the routes of administration, efficacy, toxicity, and side effects vary. The potency of the compounds for inhibition of bone resorption varies more than 10,000-fold,

increasing in the order of etidronate, tiludronate, pamidronate, alendronate, risedronate, and zoledronate. The IV use of pamidronate and zoledronate is approved for the treatment of hypercalcemia; between 30 and 90 mg pamidronate, given as a single IV dose over a few hours, returns serum calcium to normal within 24–48 h with an effect that lasts for weeks in 80–100% of patients. Zoledronate given in doses of 4 or 8 mg/5-min infusion has a more rapid and more sustained effect than pamidronate in direct comparison.

These drugs are used extensively in cancer patients. Absolute survival improvements are noted with pamidronate and zoledronate in multiple myeloma, for example. However, although still rare, there are increasing reports of jaw necrosis, especially after dental surgery, mainly in cancer patients treated with multiple doses of the more potent bisphosphonates.

CALCITONIN

Calcitonin acts within a few hours of its administration, principally through receptors on osteoclasts, to block bone resorption. Calcitonin, after 24 h of use, is no longer effective in lowering calcium. Tachyphylaxis, a known phenomenon with this drug, seems to explain the results, since the drug is often effective in the first 24 h of use. Therefore, in life-threatening hypercalcemia, calcitonin can be used effectively within the first 24 h in combination with rehydration and saline diuresis while waiting for more sustained effects from a simultaneously administered bisphosphonate such as pamidronate. Usual doses of calcitonin are 2–8 U/kg of body weight IV, SC, or IM every 6–12 h.

OTHER THERAPIES

Denosumab, an antibody that blocks the RANK ligand (RANKL) and dramatically reduces osteoclast number and function, is approved for therapy of osteoporosis. It also appears to be an effective treatment to reverse hypercalcemia of malignancy, but is not yet approved for this indication. *Plicamycin* (formerly mithramycin), which inhibits bone resorption, and *gallium nitrate*, which exerts a hypocalcemic action also by inhibiting bone resorption, are no longer used because of superior alternatives such as bisphosphonates.

Glucocorticoids have utility, especially in hypercalcemia complicating certain malignancies. They increase urinary calcium excretion and decrease intestinal calcium absorption when given in pharmacologic doses, but they also cause negative skeletal calcium balance. In normal individuals and in patients with primary hyperparathyroidism, glucocorticoids neither increase nor decrease the serum calcium concentration. In patients with hypercalcemia due to certain osteolytic malignancies, however, glucocorticoids may be effective as a result of antitumor effects. The malignancies in which hypercalcemia responds to glucocorticoids include multiple myeloma, leukemia, Hodgkin's disease, other lymphomas, and carcinoma of the breast, at least early in the course of the disease. Glucocorticoids are also effective in treating hypercalcemia due to vitamin D intoxication and sarcoidosis. Glucocorticoids are also useful in the rare form of hypercalcemia, now recognized in certain autoimmune disorders in which inactivating antibodies against the receptor imitate FHH. Elevated PTH and calcium levels are effectively lowered by the glucocorticoids. In all the preceding situations, the hypocalcemic effect develops over several days, and the usual glucocorticoid dosage is 40–100 mg prednisone (or its equivalent) daily in four divided doses. The side effects of chronic glucocorticoid therapy may be acceptable in some circumstances.

Dialysis is often the treatment of choice for severe hypercalcemia complicated by renal failure, which is difficult to manage medically. Peritoneal dialysis with calcium-free dialysis fluid can remove 5–12.5 mmol (200–500 mg) of calcium in 24–48 h and lower the serum calcium concentration by 0.7–3 mmol/L (3–12 mg/dL). Large quantities of phosphate are lost during dialysis, and serum inorganic phosphate concentration usually falls, potentially aggravating hypercalcemia. Therefore, the serum inorganic phosphate concentration

2482 should be measured after dialysis, and phosphate supplements should be added to the diet or to dialysis fluids if necessary.

Phosphate therapy, PO or IV, has a limited role in certain circumstances (Chap. 423). Correcting hypophosphatemia lowers the serum calcium concentration by several mechanisms, including bone/calcium exchange. The usual oral treatment is 1–1.5 g of phosphorus per day for several days, given in divided doses. It is generally believed, but not established, that toxicity does not occur if therapy is limited to restoring serum inorganic phosphate concentrations to normal.

Raising the serum inorganic phosphate concentration above normal decreases serum calcium levels, sometimes strikingly. Intravenous phosphate is one of the most dramatically effective treatments available for severe hypercalcemia but is toxic and even dangerous (fatal hypocalcemia). For these reasons, it is used rarely and only in severely hypercalcemic patients with cardiac or renal failure where dialysis, the preferable alternative, is not feasible or is unavailable.

SUMMARY

The various therapies for hypercalcemia are listed in Table 424-4. The choice depends on the underlying disease, the severity of the hypercalcemia, the serum inorganic phosphate level, and the renal, hepatic, and bone marrow function. Mild hypercalcemia (≤ 3 mmol/L [12 mg/dL]) can usually be managed by hydration. Severe hypercalcemia (≥ 3.7 mmol/L [15 mg/dL]) requires rapid correction. Calcitonin should be given for its rapid, albeit short-lived, blockade of bone resorption, and IV pamidronate or zoledronate should be administered, although its onset of action is delayed for 1–2 days. In addition, for the first 24–48 h, aggressive sodium-calcium diuresis with IV saline should be given and, following rehydration, large doses of furosemide or ethacrynic acid, but only if appropriate monitoring is available and cardiac and renal function are adequate. Intermediate degrees of hypercalcemia between 3 and 3.7 mmol/L (12 and 15 mg/dL) should be approached with vigorous hydration and then the most appropriate selection for the patient of the combinations used with severe hypercalcemia.

HYPOCALCEMIA

(See also Chap. 65)

PATHOPHYSIOLOGY OF HYPOCALCEMIA: CLASSIFICATION BASED ON MECHANISM

Chronic hypocalcemia is less common than hypercalcemia; causes include chronic renal failure, hereditary and acquired hypoparathyroidism, vitamin D deficiency, pseudohypoparathyroidism, and hypomagnesemia (Table 424-5).

Acute rather than chronic hypocalcemia is seen in critically ill patients or as a consequence of certain medications and often does not require specific treatment. Transient hypocalcemia is seen with severe sepsis, burns, acute kidney injury, and extensive transfusions with citrated blood. Although as many as one-half of patients in an intensive care setting are reported to have calcium concentrations of < 2.1 mmol/L (8.5 mg/dL), most do not have a reduction in ionized calcium. Patients with severe sepsis may have a decrease in ionized calcium (true hypocalcemia), but in other severely ill individuals, hypoalbuminemia is the primary cause of the reduced total calcium concentration. Alkalosis increases calcium binding to proteins, and in this setting, direct measurements of ionized calcium should be made.

Medications such as protamine, heparin, and glucagon may cause transient hypocalcemia. These forms of hypocalcemia are usually not associated with tetany and resolve with improvement in the overall medical condition. The hypocalcemia after repeated transfusions of citrated blood usually resolves quickly.

Patients with acute pancreatitis have hypocalcemia that persists during the acute inflammation and varies in degree with disease severity. The cause of hypocalcemia remains unclear. PTH values are

TABLE 424-5 FUNCTIONAL CLASSIFICATION OF HYPOCALCEMIA (EXCLUDING NEONATAL CONDITIONS)

PTH Absent	
Hereditary hypoparathyroidism	Hypomagnesemia
Acquired hypoparathyroidism	
PTH Ineffective	
Chronic kidney disease	Active vitamin D ineffective
Active vitamin D lacking	Intestinal malabsorption
↓ Dietary intake or sunlight	Vitamin D–dependent rickets type II
Defective metabolism:	Pseudohypoparathyroidism
Anticonvulsant therapy	
Vitamin D–dependent rickets type I	
PTH Overwhelmed	
Severe, acute hyperphosphatemia	Osteitis fibrosa after parathyroidectomy
Tumor lysis	
Acute kidney injury	
Rhabdomyolysis	

Abbreviation: PTH, parathyroid hormone.

reported to be low, normal, or elevated, and both resistance to PTH and impaired PTH secretion have been postulated. Occasionally, a chronic low total calcium and low ionized calcium concentration are detected in an elderly patient without obvious cause and with a paucity of symptoms; the pathogenesis is unclear.

Chronic hypocalcemia, however, is usually symptomatic and requires treatment. Neuromuscular and neurologic manifestations of chronic hypocalcemia include muscle spasms, carpopedal spasm, facial grimacing, and, in extreme cases, laryngeal spasm and convulsions. Respiratory arrest may occur. Increased intracranial pressure occurs in some patients with long-standing hypocalcemia, often in association with papilledema. Mental changes include irritability, depression, and psychosis. The QT interval on the electrocardiogram is prolonged, in contrast to its shortening with hypercalcemia. Arrhythmias occur, and digitalis effectiveness may be reduced. Intestinal cramps and chronic malabsorption may occur. Chvostek's or Trousseau's sign can be used to confirm latent tetany.

The classification of hypocalcemia shown in Table 424-5 is based on an organizationally useful premise that PTH is responsible for minute-to-minute regulation of plasma calcium concentration and, therefore, that the occurrence of hypocalcemia must mean a failure of the homeostatic action of PTH. Failure of the PTH response can occur if there is hereditary or acquired parathyroid gland failure, if PTH is ineffective in target organs, or if the action of the hormone is overwhelmed by the loss of calcium from the ECF at a rate faster than it can be replaced.

PTH ABSENT

Whether hereditary or acquired, hypoparathyroidism has a number of common components. Symptoms of untreated hypocalcemia are shared by both types of hypoparathyroidism, although the onset of hereditary hypoparathyroidism can be more gradual and associated with other developmental defects. Basal ganglia calcification and extrapyramidal syndromes are more common and earlier in onset in hereditary hypoparathyroidism. In previous decades, acquired hypoparathyroidism secondary to surgery in the neck was more common than hereditary hypoparathyroidism, but the frequency of surgically induced parathyroid failure has diminished as a result of improved surgical techniques that spare the parathyroid glands and increased use of nonsurgical therapy for hyperthyroidism. Pseudohypoparathyroidism, an example of ineffective PTH action rather than a failure of parathyroid gland production, may share several features with hypoparathyroidism, including extrasosseous calcification and extrapyramidal manifestations such as choreoathetotic movements and dystonia.

Papilledema and raised intracranial pressure may occur in both hereditary and acquired hypoparathyroidism, as do chronic changes in fingernails and hair and lenticular cataracts, the latter usually reversible with treatment of hypocalcemia. Certain skin manifestations, including alopecia and candidiasis, are characteristic of hereditary hypoparathyroidism associated with autoimmune polyglandular failure (Chap. 408).

Hypocalcemia associated with hypomagnesemia is associated with both deficient PTH release and impaired responsiveness to the hormone. Patients with hypocalcemia secondary to hypomagnesemia have absent or low levels of circulating PTH, indicative of diminished hormone release despite a maximum physiologic stimulus by hypocalcemia. Plasma PTH levels return to normal with correction of the hypomagnesemia. Thus hypoparathyroidism with low levels of PTH in blood can be due to hereditary gland failure, acquired gland failure, or acute but reversible gland dysfunction (hypomagnesemia).

Genetic Abnormalities and Hereditary Hypoparathyroidism Hereditary hypoparathyroidism can occur as an isolated entity without other endocrine or dermatologic manifestations. More typically, it occurs in association with other abnormalities such as defective development of the thymus or failure of other endocrine organs such as the adrenal, thyroid, or ovary (Chap. 408). Hereditary hypoparathyroidism is often manifest within the first decade but may appear later.

Genetic defects associated with hypoparathyroidism serve to illuminate the complexity of organ development, hormonal biosynthesis and secretion, and tissue-specific patterns of endocrine effector function (Fig. 424-5). Often, hypoparathyroidism is isolated, signifying a highly specific functional disturbance. When hypoparathyroidism is associated with other developmental or organ defects, treatment of the hypocalcemia can still be effective.

A form of hypoparathyroidism associated with defective development of both the thymus and the parathyroid glands is termed the *DiGeorge syndrome*, or the *velocardiofacial syndrome*. Congenital cardiovascular, facial, and other developmental defects are present, and patients may die in early childhood with severe infections, hypocalcemia and seizures, or cardiovascular complications. Patients can survive into adulthood, and milder, incomplete forms occur. Most cases are sporadic, but an autosomal dominant form involving microdeletions of chromosome 22q11.2 has been described. Smaller deletions in chromosome 22 are seen in incomplete forms of the *DiGeorge syndrome*, appearing in childhood or adolescence, that are manifest primarily by parathyroid gland failure. The chromosome 22 defect is now termed *DSG1*; more recently, a defect in chromosome 10p is also recognized—now called *DSG2*. The phenotypes seem similar. Studies on the chromosome 22 defect have pinpointed a transcription factor, *TBX1*. Deletions of the orthologous mouse gene show a phenotype similar to the human syndrome.

Another autosomal dominant developmental defect, featuring hypoparathyroidism, deafness, and renal dysplasia (HDR), has been studied at the genetic level. Cytogenetic abnormalities in some, but not all kindreds, point to translocation defects on chromosome 10, as in *DiGeorge syndrome*. However, the lack of immunodeficiency and heart defects distinguishes the two syndromes. Mouse models, as well as deletion analysis in some HDR patients, has identified the transcription factor *GATA3*, which is important in embryonic development and is expressed in developing kidney, ear structures, and the parathyroids.

Another pair of linked developmental disorders involving the parathyroids is recognized. *Kenney-Caffey syndrome type 1* features hypoparathyroidism, short stature, osteosclerosis, and thick cortical bones. A defect seen in Middle Eastern patients, particularly in Saudi Arabia, termed *Sanjad-Sakati syndrome*, also exhibits growth failure and other dysmorphic features. This syndrome, which is clearly autosomal recessive, involves a gene on chromosome 1q42-q43. Both syndromes apparently involve a chaperone protein, called *TBCE*, relevant to tubulin function. Recently, a defect in *FAM111A* was identified as the cause of *Kenney-Caffey syndrome type 2*.

Hypoparathyroidism can occur in association with a complex hereditary autoimmune syndrome involving failure of the adrenals,

the ovaries, the immune system, and the parathyroids in association with recurrent mucocutaneous candidiasis, alopecia, vitiligo, and pernicious anemia (Chap. 408). The responsible gene on chromosome 21q22.3 has been identified. The protein product, which resembles a transcription factor, has been termed the *autoimmune regulator*, or *AIRE*. A stop codon mutation occurs in many Finnish families with the disorder, commonly referred to as *polyglandular autoimmune type 1 deficiency*, whereas another *AIRE* mutation (Y85C) is typically observed in Jews of Iraqi and Iranian descent.

Hypoparathyroidism is seen in two disorders associated with mitochondrial dysfunction and myopathy, one termed the *Kearns-Sayre syndrome* (KSS), with ophthalmoplegia and pigmentary retinopathy, and the other termed the *MELAS syndrome* (mitochondrial encephalopathy, lactic acidosis, and stroke-like episodes). Mutations or deletions in mitochondrial genes have been identified.

Several forms of hypoparathyroidism, each rare in frequency, are seen as isolated defects; the genetic mechanisms are varied. The inheritance includes autosomal dominant, autosomal recessive, and X-linked modes. Three separate autosomal defects involving the parathyroid gene have been recognized: one is dominant and the other two are recessive. The dominant form has a point mutation in the signal sequence, a critical region involved in intracellular transport of the hormone precursor. An Arg for Cys mutation interferes with processing of the precursor and is believed to trigger an apoptotic cellular response, hence acting as a dominant negative. The other two forms are recessive. One point mutation also blocks cleavage of the PTH precursor but requires both alleles to cause hypoparathyroidism. The third involves a single-nucleotide base change that results in an exon splicing defect; the lost exon contains the promoter—hence, the gene is silenced. An X-linked recessive form of hypoparathyroidism has been described in males, and the defect has been localized to chromosome Xq26-q27, perhaps involving the *SOX3* gene.

Abnormalities in the CaSR are detected in three distinctive hypocalcemic disorders. All are rare, but more than 10 different gain-of-function mutations have been found in one form of hypocalcemia termed *autosomal dominant hypocalcemic hypercalciuria* (ADHH). The receptor senses the ambient calcium level as excessive and suppresses PTH secretion, leading to hypocalcemia. The hypocalcemia is aggravated by constitutive receptor activity in the renal tubule causing excretion of inappropriate amounts of calcium. Recognition of the syndrome is important because efforts to treat the hypocalcemia with vitamin D analogues and increased oral calcium exacerbate the already excessive urinary calcium excretion (several grams or more per 24 h), leading to irreversible renal damage from stones and ectopic calcification.

Other causes of isolated hypoparathyroidism include homozygous, inactivating mutations in the parathyroid-specific transcription factor *GCM2*, which lead to an autosomal recessive form of the disease, or heterozygous point mutations in *GCM2*, which have a dominant negative effect on the wild-type protein and thus lead to an autosomal dominant form of hypoparathyroidism. Furthermore, heterozygous mutations in *G11*, one of the two signaling proteins downstream of the CaSR, have been identified as a cause of autosomal dominant hypoparathyroidism.

Bartter's syndrome is a group of disorders associated with disturbances in electrolyte and acid/base balance, sometimes with nephrocalcinosis and other features. Several types of ion channels or transporters are involved. Curiously, *Bartter's syndrome type V* has the electrolyte and pH disturbances seen in the other syndromes but appears to be due to a gain of function in the CaSR. The defect may be more severe than in ADHH and explains the additional features seen beyond hypocalcemia and hypercalciuria.

As with autoimmune disorders that block the CaSR (discussed above under hypercalcemic conditions), there are autoantibodies that at least transiently activate the CaSR, leading to suppressed PTH secretion and hypocalcemia. This disorder may wax and wane.

Acquired Hypoparathyroidism Acquired chronic hypoparathyroidism is usually the result of inadvertent surgical removal of all the parathyroid

2484 glands; in some instances, not all the tissue is removed, but the remainder undergoes vascular supply compromise secondary to fibrotic changes in the neck after surgery. In the past, the most frequent cause of acquired hypoparathyroidism was surgery for hyperthyroidism. Hypoparathyroidism now usually occurs after surgery for hyperparathyroidism when the surgeon, facing the dilemma of removing too little tissue and thus not curing the hyperparathyroidism, removes too much. Parathyroid function may not be totally absent in all patients with postoperative hypoparathyroidism.

Rare causes of acquired chronic hypoparathyroidism include radiation-induced damage subsequent to radioiodine therapy of hyperthyroidism and glandular damage in patients with hemochromatosis or hemosiderosis after repeated blood transfusions. Infection may involve one or more of the parathyroids but usually does not cause hypoparathyroidism because all four glands are rarely involved.

Transient hypoparathyroidism is frequent following surgery for hyperparathyroidism. After a variable period of hypoparathyroidism, normal parathyroid function may return due to hyperplasia or recovery of remaining tissue. Occasionally, recovery occurs months after surgery.

TREATMENT ACQUIRED AND HEREDITARY HYPOPARATHYROIDISM

Treatment involves replacement with vitamin D or 1,25(OH)₂D (calcitriol) combined with a high oral calcium intake. In most patients, blood calcium and phosphate levels are satisfactorily regulated, but some patients show resistance and a brittleness, with a tendency to alternate between hypocalcemia and hypercalcemia. For many patients, vitamin D in doses of 40,000–120,000 U/d (1–3 mg/d) combined with ≥1 g elemental calcium is satisfactory. The wide dosage range reflects the variation encountered from patient to patient; precise regulation of each patient is required. Compared to typical daily requirements in euparathyroid patients of 200 U/d (or in older patients as high as 800 U/d), the high dose of vitamin D (as much as 100-fold higher) reflects the reduced conversion of vitamin D to 1,25(OH)₂D. Many physicians now use 0.5–1 µg of calcitriol in management of such patients, especially if they are difficult to control. Because of its storage in fat, when vitamin D is withdrawn, weeks are required for the disappearance of the biologic effects, compared with a few days for calcitriol, which has a rapid turnover.

Oral calcium and vitamin D restore the overall calcium-phosphate balance but do not reverse the lowered urinary calcium reabsorption typical of hypoparathyroidism. Therefore, care must be taken to avoid excessive urinary calcium excretion after vitamin D and calcium replacement therapy; otherwise, nephrocalcinosis and kidney stones can develop, and the risk of CKD is increased. Thiazide diuretics lower urine calcium by as much as 100 mg/d in hypoparathyroid patients on vitamin D, provided they are maintained on a low-sodium diet. Use of thiazides seems to be of benefit in mitigating hypercalciuria and easing the daily management of these patients.

There are now trials of parenterally administered PTH (either PTH[1–34] or PTH[1–84]) in patients with hypoparathyroidism providing greater ease of maintaining serum calcium and reducing urinary calcium excretion (desirable to protect any renal damage). However, PTH therapy for the treatment of hypoparathyroidism is not approved as of yet.

Hypomagnesemia Severe hypomagnesemia (<0.4 mmol/L; <0.8 meq/L) is associated with hypocalcemia ([Chap. 423](#)). Restoration of the total-body magnesium deficit leads to rapid reversal of hypocalcemia. There are at least two causes of the hypocalcemia—impaired PTH secretion and reduced responsiveness to PTH. **For further discussion of causes and treatment of hypomagnesemia, see Chap. 423.**

The effects of magnesium on PTH secretion are similar to those of calcium; hypermagnesemia suppresses and hypomagnesemia stimulates PTH secretion. The effects of magnesium on PTH secretion are

normally of little significance, however, because the calcium effects dominate. Greater change in magnesium than in calcium is needed to influence hormone secretion. Nonetheless, hypomagnesemia might be expected to increase hormone secretion. It is therefore surprising to find that severe hypomagnesemia is associated with blunted secretion of PTH. The explanation for the paradox is that severe, chronic hypomagnesemia leads to intracellular magnesium deficiency, which interferes with secretion and peripheral responses to PTH. The mechanism of the cellular abnormalities caused by hypomagnesemia is unknown, although effects on adenylate cyclase (for which magnesium is a cofactor) have been proposed.

PTH levels are undetectable or inappropriately low in severe hypomagnesemia despite the stimulus of severe hypocalcemia, and acute repletion of magnesium leads to a rapid increase in PTH level. Serum phosphate levels are often not elevated, in contrast to the situation with acquired or idiopathic hypoparathyroidism, probably because phosphate deficiency is often seen in hypomagnesemia ([Chap. 393](#)).

Diminished peripheral responsiveness to PTH also occurs in some patients, as documented by subnormal response in urinary phosphorus and urinary cAMP excretion after administration of exogenous PTH to patients who are hypocalcemic and hypomagnesemic. Both blunted PTH secretion and lack of renal response to administered PTH can occur in the same patient. When acute magnesium repletion is undertaken, the restoration of PTH levels to normal or supra-normal may precede restoration of normal serum calcium by several days.

TREATMENT HYPOMAGNESEMIA

Repletion of magnesium cures the condition. Repletion should be parenteral. Attention must be given to restoring the intracellular deficit, which may be considerable. After IV magnesium administration, serum magnesium may return transiently to the normal range, but unless replacement therapy is adequate, serum magnesium will again fall. If the cause of the hypomagnesemia is renal magnesium wasting, magnesium may have to be given long-term to prevent recurrence ([Chap. 423](#)).

PTH INEFFECTIVE

PTH is not sufficiently active to fully prevent hypocalcemia (although retaining phosphaturic activity, for example). This problem occurs when the PTH1R–signaling protein complex is defective (as in the different forms of pseudohypoparathyroidism [PHP], discussed below); when PTH action to promote calcium absorption from the diet via the synthesis of 1,25(OH)₂D is insufficient because of vitamin D deficiency or because vitamin D is ineffective (defects in vitamin D receptor or vitamin D synthesis); or in CKD in which the calcium-elevating action of PTH is impaired.

Typically, hypophosphatemia is more severe than hypocalcemia in vitamin D deficiency states because of the increased secretion of PTH, which, although only partly effective in elevating blood calcium, is readily capable of promoting urinary phosphate excretion.

PHP, on the other hand, has a pathophysiology that is different from the other disorders of ineffective PTH action. PHP resembles hypoparathyroidism (in which PTH synthesis is deficient) and is manifested by hypocalcemia and hyperphosphatemia, yet elevated PTH levels. The cause of the disorder is defective PTH-dependent activation of the stimulatory G protein complex or the downstream effector protein kinase A, resulting in failure of PTH to increase intracellular cAMP or to respond to elevated cAMP levels (see below).

Chronic Kidney Disease Improved medical management of CKD now allows many patients to survive for decades and hence allows time enough to develop features of renal osteodystrophy, which must be controlled to avoid additional morbidity. Impaired production of 1,25(OH)₂D is now thought to be the principal factor that causes calcium deficiency, secondary hyperparathyroidism, and bone disease;

hyperphosphatemia typically occurs only in the later stages of the disease. Low levels of $1,25(\text{OH})_2\text{D}$ due to increased FGF23 production in bone are critical in the development of hypocalcemia. The uremic state also causes impairment of intestinal absorption by mechanisms other than defects in vitamin D metabolism. Nonetheless, treatment with supraphysiologic amounts of vitamin D or calcitriol can correct the impaired calcium absorption. Because increased FGF23 levels are seen even in early stages of CKD and have been reported to correlate with increased mortality and left ventricular hypertrophy, there is current interest in approaches to lower intestinal phosphate absorption early during the course of kidney disease and to thereby lower FGF23 levels. However, there is concern as to whether vitamin D supplementation increases the circulating FGF23 levels in CKD patients. Although vitamin D analogs improve survival in this patient population, it is notable that there are often dramatic elevations of FGF23.

Hyperphosphatemia in CKD lowers blood calcium levels by several mechanisms, including extrasosseous deposition of calcium and phosphate, impairment of the bone-resorbing action of PTH, and reduction in $1,25(\text{OH})_2\text{D}$ production by remaining renal tissue.

TREATMENT CHRONIC KIDNEY DISEASE

Therapy of CKD (**Chap. 335**) involves appropriate management of patients prior to dialysis and adjustment of regimens once dialysis is initiated. Attention should be paid to restriction of phosphate in the diet; avoidance of aluminum-containing phosphate-binding antacids to prevent the problem of aluminum intoxication; provision of an adequate calcium intake by mouth, usually 1–2 g/d; and supplementation with 0.25–1 $\mu\text{g}/\text{d}$ calcitriol or other activated forms of vitamin D. Each patient must be monitored closely. The aim of therapy is to restore normal calcium balance to prevent osteomalacia and severe secondary hyperparathyroidism (it is usually recommended to maintain PTH levels between 100 and 300 pg/mL) and, in light of evidence of genetic changes and monoclonal outgrowths of parathyroid glands in CKD patients, to prevent secondary hyperparathyroidism from becoming autonomous hyperparathyroidism. Reduction of hyperphosphatemia and restoration of normal intestinal calcium absorption by calcitriol can improve blood calcium levels and reduce the manifestations of secondary hyperparathyroidism. Because adynamic bone disease can occur in association with low PTH levels, it is important to avoid excessive suppression of the parathyroid glands while recognizing the beneficial effects of controlling the secondary hyperparathyroidism. These patients should probably be closely monitored with PTH assays that detect only the full-length PTH(1–84) to ensure that biologically active PTH and not inactive, inhibitory PTH fragments are measured. Use of phosphate-binding agents such as sevelamer is approved only in end-stage renal disease, but it may be necessary to initiate such treatment much earlier during the course of kidney disease to prevent the increase in FGF23 and its “off-target” effects.

Vitamin D Deficiency due to Inadequate Diet and/or Sunlight Vitamin D deficiency due to inadequate intake of dairy products enriched with vitamin D, lack of vitamin supplementation, and reduced sunlight exposure in the elderly, particularly during winter in northern latitudes, is more common in the United States than previously recognized. Biopsies of bone in elderly patients with hip fracture (documenting osteomalacia) and abnormal levels of vitamin D metabolites, PTH, calcium, and phosphate indicate that vitamin D deficiency may occur in as many as 25% of elderly patients, particularly in northern latitudes in the United States. Concentrations of $25(\text{OH})\text{D}$ are low or low-normal in these patients. Quantitative histomorphometric analysis of bone biopsy specimens from such individuals reveals widened osteoid seams consistent with osteomalacia (**Chap. 423**). PTH hypersecretion compensates for the tendency for the blood calcium to fall but also increases renal phosphate excretion and thus causes osteomalacia.

Treatment involves adequate replacement with vitamin D and calcium until the deficiencies are corrected. Severe hypocalcemia rarely occurs in moderately severe vitamin D deficiency of the elderly, but vitamin D deficiency must be considered in the differential diagnosis of mild hypocalcemia.

Mild hypocalcemia, secondary hyperparathyroidism, severe hypophosphatemia, and a variety of nutritional deficiencies occur with gastrointestinal diseases. Hepatocellular dysfunction can lead to reduction in $25(\text{OH})\text{D}$ levels, as in portal or biliary cirrhosis of the liver, and malabsorption of vitamin D and its metabolites, including $1,25(\text{OH})_2\text{D}$, may occur in a variety of bowel diseases, hereditary or acquired. Hypocalcemia itself can lead to steatorrhea, due to deficient production of pancreatic enzymes and bile salts. Depending on the disorder, vitamin D or its metabolites can be given parenterally, guaranteeing adequate blood levels of active metabolites.

Defective Vitamin D Metabolism • ANTICONVULSANT THERAPY Anticonvulsant therapy with any of several agents induces acquired vitamin D deficiency by increasing the conversion of vitamin D to inactive compounds and/or causing resistance to its action. The more marginal the vitamin D intake in the diet, the more likely that anticonvulsant therapy will lead to abnormal mineral and bone metabolism.

VITAMIN D–DEPENDENT RICKETS TYPE I Vitamin D–dependent rickets type I, previously termed *pseudo-vitamin D-resistant rickets*, differs from true vitamin D–resistant rickets (vitamin D–dependent rickets type II, see below) in that it is typically less severe and the biochemical and radiographic abnormalities can be reversed with appropriate doses of the vitamin’s active metabolite, $1,25(\text{OH})_2\text{D}$. Physiologic amounts of calcitriol cure the disease (**Chap. 423**). This finding fits with the pathophysiology of the disorder, which is autosomal recessive, and is now known to be caused by mutations in the gene encoding $25(\text{OH})\text{D}$ -1 α -hydroxylase. Both alleles are inactivated in affected patients, and compound heterozygotes, harboring distinct mutations, are common.

Clinical features include hypocalcemia, often with tetany or convulsions, hypophosphatemia, secondary hyperparathyroidism, and osteomalacia, often associated with skeletal deformities and increased alkaline phosphatase. Treatment involves physiologic replacement doses of $1,25(\text{OH})_2\text{D}$ (**Chap. 423**).

VITAMIN D–DEPENDENT RICKETS TYPE II Vitamin D–dependent rickets type II results from end-organ resistance to the active metabolite $1,25(\text{OH})_2\text{D}$. The clinical features resemble those of the type I disorder and include hypocalcemia, hypophosphatemia, secondary hyperparathyroidism, and rickets but also partial or total alopecia. Plasma levels of $1,25(\text{OH})_2\text{D}$ are elevated, in keeping with the refractoriness of the end organs. This disorder is caused by mutations in the gene encoding the vitamin D receptor; treatment is difficult and requires regular, usually nocturnal calcium infusions, which dramatically improve growth but do not restore hair growth (**Chap. 423**).

Pseudohypoparathyroidism PHP refers to a group of distinct inherited disorders. Patients affected by PHP type Ia (PHP-Ia) are characterized by symptoms and signs of hypocalcemia in association with distinctive skeletal and developmental defects. The hypocalcemia is due to a deficient response to PTH, which is probably restricted to the proximal renal tubules. Hyperplasia of the parathyroids, a response to hormone-resistant hypocalcemia, causes elevation of PTH levels. Studies, both clinical and basic, have clarified some aspects of these disorders, including the variable clinical spectrum, the pathophysiology, the genetic defects, and their mode of inheritance.

A working classification of the various forms of PHP is given in **Table 424-6**. The classification scheme is based on the signs of ineffective PTH action (low calcium and high phosphate), low or normal urinary cAMP response to exogenous PTH, the presence or absence of *Albright’s hereditary osteodystrophy* (AHO), and assays to measure the concentration of the G_α subunit of the adenylate cyclase enzyme. Using these criteria, there are four types: PHP types Ia and Ib; pseudopseudohypoparathyroidism (PPHP), and PHP-II.

TABLE 424-6 CLASSIFICATION OF PSEUDOHYPOPARATHYROIDISM (PHP) AND PSEUDOPSEUDOHYPOPARATHYROIDISM (PPHP)

Type	Hypocalcemia, Hyperphosphatemia	Response of Urinary cAMP to PTH	Serum PTH	G _s α Subunit Deficiency	AHO	Resistance to hormones other than PTH
PHP-Ia	Yes	↓	↑	Yes	Yes	Yes
PPHP	No	Normal	Normal	Yes	Yes	No
PHP-Ib	Yes	↓	↑	No	No	Yes (in some patients)
PHP-II	Yes	Normal	↑	No	No	No
Acrodysostosis with hormonal resistance	Yes	Normal (but ↓ phosphaturic response)	↑	No	Yes	Yes

Abbreviations: ↓, decreased; ↑, increased; AHO, Albright's hereditary osteodystrophy; PTH, parathyroid hormone.

PHP-IA AND PHP-IB Individuals with PHP-I, the most common of the disorders, show a deficient urinary cAMP response to administration of exogenous PTH. Patients with PHP-I are divided into type Ia and type Ib. Patients with PHP-Ia show evidence for AHO and reduced amounts of G_s α protein/activity, as determined in readily accessible tissues such as erythrocytes, lymphocytes, and fibroblasts. Patients with PHP-Ib typically lack evidence for AHO and they have normal G_s α activity. PHP-Ic, sometimes listed as a third form of PHP-I, is really a variant of PHP-Ia, although the mutant G_s α shows normal activity in certain in vitro assays.

Most patients who have PHP-Ia reveal characteristic features of AHO, which consist of short stature, round face, obesity, skeletal anomalies (brachydactyly), intellectual impairment, and/or heterotopic calcifications. Patients have low calcium and high phosphate levels, as with true hypoparathyroidism. PTH levels, however, are elevated, reflecting resistance to hormone action.

Amorphous deposits of calcium and phosphate are found in the basal ganglia in about one-half of patients. The defects in metacarpal and metatarsal bones are sometimes accompanied by short phalanges as well, possibly reflecting premature closing of the epiphyses. The typical findings are short fourth and fifth metacarpals and metatarsals. The defects are usually bilateral. Exostoses and radius curvus are frequent.

Inheritance and Genetic Defects Multiple defects at the *GNAS* locus have now been identified in PHP-Ia, PHP-Ib, and PPHP patients. This gene, which is located on chromosome 20q13.3, encodes the α-subunit of the stimulatory G protein (G_s α), among other products (see below). Mutations include abnormalities in splice junctions associated with deficient mRNA production, point mutations, insertions, and/or deletion that all result in a protein with defective function resulting in a 50% reduction of G_s α activity in erythrocytes or other cells.

Detailed analyses of disease transmission in affected kindreds have clarified many features of PHP-Ia, PPHP, and PHP-Ib (Fig. 424-7). The former two entities, often traced through multiple generations, have an inheritance pattern consistent with genetic imprinting. The phenomenon of gene imprinting, involving methylation of genetic loci, independent of any mutation, impairs transcription from either the maternal or the paternal allele (Chap. 82). The G_s α transcript is biallelically expressed in most tissues; expression from paternal allele is silenced through as-of-yet unknown mechanisms in some tissues including the proximal renal tubules and the thyroid; consequently, inheritance of a defective paternal allele has no implications with regard to hormonal function. Thus, females affected by either PHP-Ia or PPHP will have offspring with PHP-Ia, if these children inherit the allele carrying the *GNAS* mutation; in contrast, if the mutant allele is inherited from a male affected by either disorder, the offspring will exhibit PPHP. Consistent with these data in humans, gene-ablation studies in mice have shown that inheritance of the mutant G_s α allele from the female causes much reduced G_s α protein in renal cortex, hypocalcemia, and resistance to PTH. Offspring inheriting the mutant allele from the male showed no evidence of PTH resistance or hypocalcemia.

Imprinting is tissue selective. Paternal G_s α expression is not silenced in most tissues. It seems likely, therefore, that the AHO phenotype recognized in PPHP as well as PHP-Ia reflects G_s α haploinsufficiency during embryonic or postnatal development.

The complex mechanisms that control the *GNAS* gene contribute to challenges involved in unraveling the pathogenesis of these disorders, especially that of PHP-Ib. Much intensive work with families in which multiple members are affected by PHP-Ib, as well as studies of the complex regulation of the *GNAS* gene locus, have now shown that PHP-Ib is caused by microdeletions within or upstream of the *GNAS* locus, which are associated with a loss of DNA methylation at one or several loci of the maternal allele (Table 424-6). These abnormalities in methylation silence the expression of the gene. This leads in the proximal renal tubules—where G_s α appears to be expressed exclusively from the maternal allele—to PTH resistance.

PHP-Ib, lacking the AHO phenotype in most instances, shares with PHP-Ia the hypocalcemia and hyperphosphatemia caused by PTH resistance, and thus the blunted urinary cAMP response to administered PTH, a standard test to assess the presence or absence of hormone resistance (Table 424-6). Furthermore, these endocrine abnormalities become apparent only if the disease-causing mutation is inherited maternally. Bone responsiveness may be excessive rather than blunted in PHP-Ib (and in PHP-Ia) patients, based on case reports that have emphasized an osteitis fibrosa-like pattern in several PHP-Ib patients.

PHP-II refers to patients with hypocalcemia and hyperphosphatemia, who have a normal urinary cAMP but an impaired urinary phosphaturic

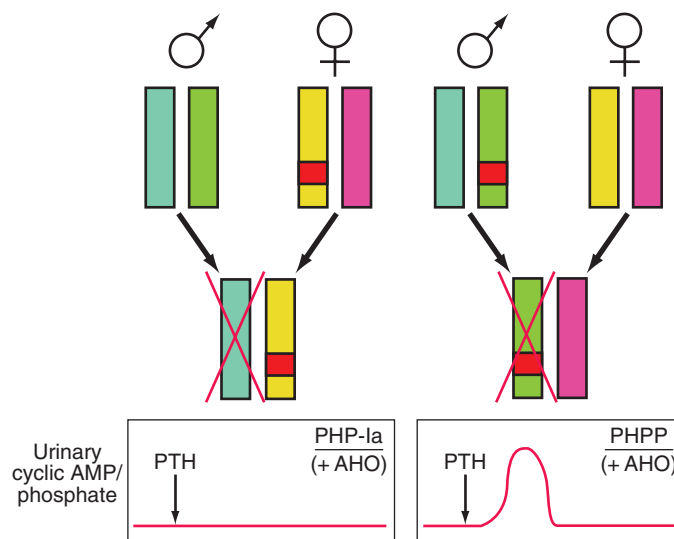


FIGURE 424-7 Paternal imprinting of renal parathyroid hormone (PTH) resistance. An impaired excretion of urinary cyclic AMP and phosphate is observed in patients with pseudohypoparathyroidism type Ia (PHP-Ia). In the renal cortex, there is selective silencing of paternal G_s α expression. The disease becomes manifest only in patients who inherit the defective gene from an obligate female carrier (left). If the genetic defect is inherited from an obligate male carrier, there is no biochemical abnormality; administration of PTH causes an appropriate increase in the urinary cyclic AMP and phosphate concentration (pseudo-PHP [PPHP]; right). Both patterns of inheritance lead to Albright's hereditary osteodystrophy (AHO), perhaps because of haplotype insufficiency—i.e., both copies of G_s α must be active for normal bone development.

response to PTH. In a PHP-II variant, referred to as acrodysostosis with hormonal resistance (ADOHR), patients have a defect in the regulatory subunit of PKA (PRKAR1A) that mediates the response to PTH distal to cAMP production. Acrodysostosis without hormonal resistance is caused by mutations in the cAMP-selective phosphodiesterase 4 (ADOP4). It remains unclear why the PTH-resistance in some patients, labeled as PHP-II without bony abnormalities, resolves upon treatment with vitamin D supplements.

The diagnosis of these hormone-resistant states can usually be made without difficulty when there is a positive family history for features of AHO, in association with the signs and symptoms of hypocalcemia. In both categories—PHP-Ia and PHP-Ib—serum PTH levels are elevated, particularly when patients are hypocalcemic. However, patients with PHP-Ib or PHP-II without acrodysostosis present only with hypocalcemia and high PTH levels, as evidence for hormone resistance. In PHP-Ia and PHP-Ib, the response of urinary cAMP to the administration of exogenous PTH is blunted. The diagnosis of PHP-II, in the absence of acrodysostosis, is more complex, and vitamin D deficiency must be excluded before such a diagnosis can be entertained.

TREATMENT PSEUDOHYPOPARATHYROIDISM

Treatment of PHP is similar to that of hypoparathyroidism, except that calcium and vitamin D doses are usually higher. Patients with PHP show no PTH-resistance in the distal tubules—hence, urinary calcium clearance is typically reduced, and they are not at risk of developing nephrocalcinosis as are patients with true hypoparathyroidism, unless overtreatment occurs, for example, after the completion of pubertal development and skeletal maturation, when calcium and 1,25(OH)₂D treatment should be reduced. Variability in response makes it necessary to establish the optimal regimen for each patient, based on maintaining appropriate blood calcium level and urinary calcium excretion and keeping the PTH level within or slightly above the normal range.

PTH OVERWHELMED

Occasionally, loss of calcium from the ECF is so severe that PTH cannot compensate. Such situations include acute pancreatitis and severe, acute hyperphosphatemia, often in association with renal failure, conditions in which there is rapid efflux of calcium from the ECF. Severe hypocalcemia can occur quickly; PTH rises in response to hypocalcemia but does not return blood calcium to normal.

Severe, Acute Hyperphosphatemia Severe hyperphosphatemia is associated with extensive tissue damage or cell destruction (Chap. 423). The combination of increased release of phosphate from muscle and impaired ability to excrete phosphorus because of renal failure causes moderate to severe hyperphosphatemia, the latter causing calcium loss from the blood and mild to moderate hypocalcemia. Hypocalcemia is usually reversed with tissue repair and restoration of renal function as phosphorus and creatinine values return to normal. There may even be a mild hypercalcemic period in the oliguric phase of renal function recovery. This sequence, severe hypocalcemia followed by mild hypercalcemia, reflects widespread deposition of calcium in muscle and subsequent redistribution of some of the calcium to the ECF after phosphate levels return to normal.

Other causes of hyperphosphatemia include hypothermia, massive hepatic failure, and hematologic malignancies, either because of high cell turnover of malignancy or because of cell destruction by chemotherapy.

TREATMENT SEVERE, ACUTE HYPERPHOSPHATEMIA

Treatment is directed toward lowering of blood phosphate by the administration of phosphate-binding antacids or dialysis, often needed for the management of CKD. Although calcium replacement may be necessary if hypocalcemia is severe and symptomatic, calcium administration during the hyperphosphatemic period tends

to increase extraosseous calcium deposition and aggravate tissue damage. The levels of 1,25(OH)₂D may be low during the hyperphosphatemic phase and return to normal during the oliguric phase of recovery.

Osteitis Fibrosa after Parathyroidectomy Severe hypocalcemia after parathyroid surgery is rare now that osteitis fibrosa cystica is an infrequent manifestation of hyperparathyroidism. When osteitis fibrosa cystica is severe, however, bone mineral deficits can be large. After parathyroidectomy, hypocalcemia can persist for days if calcium replacement is inadequate. Treatment may require parenteral administration of calcium; addition of calcitriol and oral calcium supplementation is sometimes needed for weeks to a month or two until bone defects are filled (which, of course, is of therapeutic benefit in the skeleton), making it possible to discontinue parenteral calcium and/or reduce the amount.

DIFFERENTIAL DIAGNOSIS OF HYPOCALCEMIA

Care must be taken to ensure that true hypocalcemia is present; in addition, acute transient hypocalcemia can be a manifestation of a variety of severe, acute illnesses, as discussed above. *Chronic hypocalcemia*, however, can usually be ascribed to a few disorders associated with absent or ineffective PTH. Important clinical criteria include the duration of the illness, signs or symptoms of associated disorders, and the presence of features that suggest a hereditary abnormality. A nutritional history can be helpful in recognizing a low intake of vitamin D and calcium in the elderly, and a history of excessive alcohol intake may suggest magnesium deficiency.

Hypoparathyroidism and PHP are typically lifelong illnesses, usually (but not always) appearing by adolescence; hence, a recent onset of hypocalcemia in an adult is more likely due to nutritional deficiencies, renal failure, or intestinal disorders that result in deficient or ineffective vitamin D. Neck surgery, even long past, however, can be associated with a delayed onset of postoperative hypoparathyroidism. A history of seizure disorder raises the issue of anticonvulsive medication. Developmental defects may point to the diagnosis of PHP. Rickets and a variety of neuromuscular syndromes and deformities may indicate ineffective vitamin D action, either due to defects in vitamin D metabolism or to vitamin D deficiency.

A pattern of *low calcium with high phosphorus* in the absence of renal failure or massive tissue destruction almost invariably means hypoparathyroidism or PHP. A *low calcium and low phosphorus* pattern points to absent or ineffective vitamin D, thereby impairing the action of PTH on calcium metabolism (but not phosphate clearance). The relative ineffectiveness of PTH in calcium homeostasis in vitamin D deficiency, anticonvulsant therapy, gastrointestinal disorders, and hereditary defects in vitamin D metabolism leads to secondary hyperparathyroidism as a compensation. The excess PTH on renal tubule phosphate transport accounts for renal phosphate wasting and hypophosphatemia.

Exceptions to these patterns may occur. Most forms of hypomagnesemia are due to long-standing nutritional deficiency as seen in chronic alcoholics. Despite the fact that the hypocalcemia is principally due to an acute absence of PTH, phosphate levels are usually low, rather than elevated, as in hypoparathyroidism. Chronic renal failure is often associated with hypocalcemia and hyperphosphatemia, despite secondary hyperparathyroidism.

Diagnosis is usually established by application of the PTH immunoassay, tests for vitamin D metabolites, and measurements of the urinary cAMP response to exogenous PTH. In hereditary and acquired hypoparathyroidism and in severe hypomagnesemia, PTH is either undetectable or inappropriately in the normal range (Fig. 424-4). This finding in a hypocalcemic patient is supportive of hypoparathyroidism, as distinct from ineffective PTH action, in which even mild hypocalcemia is associated with elevated PTH levels. Hence a failure to detect elevated PTH levels establishes the diagnosis of hypoparathyroidism; elevated levels suggest the presence of secondary hyperparathyroidism, as found in many of the situations in which the hormone is

2488 ineffective due to associated abnormalities in vitamin D action. Assays for 25(OH)D can be helpful. Low or low-normal 25(OH)D indicates vitamin D deficiency due to lack of sunlight, inadequate vitamin D intake, or intestinal malabsorption. Recognition that mild hypocalcemia, rickets, and hypophosphatemia are due to anticonvulsant therapy is made by history.

TREATMENT HYPOCALCEMIC STATES

The management of hypoparathyroidism, PHP, chronic renal failure, and hereditary defects in vitamin D metabolism involves the use of vitamin D or vitamin D metabolites and calcium supplementation. Vitamin D itself is the least expensive form of vitamin D replacement and is frequently used in the management of uncomplicated hypoparathyroidism and some disorders associated with ineffective vitamin D action. When vitamin D is used prophylactically, as in the elderly or in those with chronic anticonvulsant therapy, there is a wider margin of safety than with the more potent metabolites. However, most of the conditions in which vitamin D is administered chronically for hypocalcemia require amounts 50–100 times the daily replacement dose because the formation of 1,25(OH)₂D is deficient. In such situations, vitamin D is no safer than the active metabolite because intoxication can occur with high-dose therapy (because of storage in fat). Calcitriol is more rapid in onset of action and also has a short biologic half-life.

Vitamin D (at least 1000 U/d [2–3 µg/d] [higher levels required in older persons]) or calcitriol (0.25–1 µg/d) is required to prevent rickets in normal individuals. In contrast, 40,000–120,000 U (1–3 mg) of vitamin D₂ or D₃ is typically required in hypoparathyroidism. The dose of calcitriol is unchanged in hypoparathyroidism, because the defect is in hydroxylation by the 25(OH)D-1α-hydroxylase. Calcitriol is also used in disorders of 25(OH)D-1α-hydroxylase; vitamin D receptor defects are much more difficult to treat.

Patients with hypoparathyroidism should be given 2–3 g of elemental calcium PO each day. The two agents, vitamin D or calcitriol and oral calcium, can be varied independently. Urinary calcium excretion needs to be monitored carefully. If hypocalcemia alternates with episodes of hypercalcemia in high-brittleness patients with hypoparathyroidism, administration of calcitriol and use of thiazides, as discussed above, may make management easier. Clinical trials with PTH(1–34) or PTH(1–84) are promising, but these alternative treatments have not yet been approved.

–2.5. Postmenopausal women who fall at the lower end of the young normal range (a T-score <–1.0) are defined as having low bone density and are also at increased risk of osteoporosis. Although risk is lower in this group, more than 50% of fractures among postmenopausal women, including hip fractures, occur in this group with low bone density, because the number of individuals in this category is so much larger than that in the osteoporosis range. As a result, there are ongoing attempts to identify individuals within the low bone density range who are at high risk of fracture and might benefit from pharmacologic intervention. Furthermore, some have advocated using fracture risk as the “diagnostic” criterion for osteoporosis.

EPIDEMIOLOGY

In the United States, as many as 9 million adults have osteoporosis (T-score <–2.5 in either spine or hip), and an additional 48 million individuals have bone mass levels that put them at increased risk of developing osteoporosis (e.g., bone mass T-score <–1.0). Osteoporosis occurs more frequently with increasing age as bone tissue is lost progressively. In women, the loss of ovarian function at menopause (typically about age 50) precipitates rapid bone loss so that most women meet the diagnostic criterion for osteoporosis by age 70–80. As the population continues to age, the number of individuals with osteoporosis and fractures will also continue to increase, despite a recognized reduction in age-specific risk. It is estimated that about 2 million fractures occur each year in the United States as a consequence of osteoporosis, and that number is expected to increase as the population continues to age.

The epidemiology of fractures follows the trend for loss of bone density, with exponential increases in both hip and vertebral fractures with age. Fractures of the distal radius have a somewhat different epidemiology, increasing in frequency before age 50 and plateauing by age 60, with only a modest age-related increase thereafter. In contrast, incidence rates for hip fractures double every 5 years after age 70 (Fig. 425-1). This distinct epidemiology may be related to the way the elderly fall as they age, with fewer falls on an outstretched hand and more falls directly on the hip. About 300,000 hip fractures occur each year in the United States, most of which require hospital admission and surgical intervention. The probability that a 50-year-old white individual will have a hip fracture during his or her lifetime is 14% for women and 5% for men; the risk for African Americans is lower (about one-half those rates), and the risk for Asians is roughly equal to that for whites. Hip fractures are associated with a high incidence of deep vein thrombosis and pulmonary embolism (20–50%) and a mortality rate between 5 and 20% during the year after surgery.

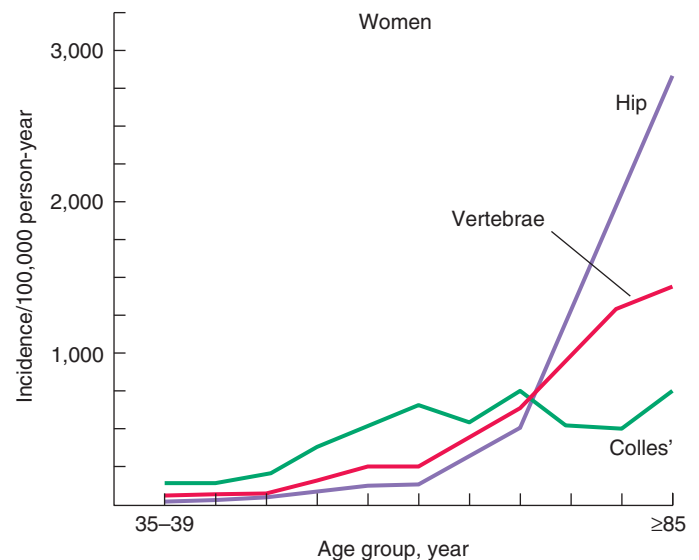


FIGURE 425-1 Epidemiology of vertebral, hip, and Colles' fractures with age. (Adapted from C Cooper, LJ Melton III: Trends Endocrinol Metab 3:224, 1992; with permission.)

425 Osteoporosis

Robert Lindsay, Felicia Cosman

Osteoporosis, a condition characterized by decreased bone strength, is prevalent among postmenopausal women but also occurs in men and women with underlying conditions or major risk factors associated with bone demineralization. Its chief clinical manifestations are vertebral and hip fractures, although fractures can occur at almost any skeletal site. Osteoporosis affects almost 10 million individuals in the United States, but only a small proportion are diagnosed and treated.

DEFINITION

Osteoporosis is defined as a reduction in the strength of bone that leads to an increased risk of fractures. Loss of bone tissue is associated with deterioration in skeletal microarchitecture. The World Health Organization (WHO) operationally defines osteoporosis as a bone density that falls 2.5 standard deviations (SD) below the mean for young healthy adults of the same sex—also referred to as a *T-score* of



FIGURE 425-2 Lateral spine x-ray showing severe osteopenia and a severe wedge-type deformity (severe anterior compression).

There is also significant morbidity, with about 20–40% of survivors requiring long-term care, and many who are unable to function as they did before the fracture.

There are about 550,000 vertebral crush fractures per year in the United States. Only a fraction (estimated to be one-third) of them are recognized clinically, because many are relatively asymptomatic and are identified incidentally during radiography for other purposes (Fig. 425-2). Vertebral fractures rarely require hospitalization but are associated with long-term morbidity and a slight increase in mortality rates, primarily related to pulmonary disease. Multiple vertebral fractures lead to height loss (often of several inches), kyphosis, and secondary pain and discomfort related to altered biomechanics of the back. Thoracic fractures can be associated with restrictive lung disease, whereas lumbar fractures are associated with abdominal symptoms that include distention, early satiety, and constipation.

Approximately 400,000 wrist fractures and 135,000 pelvic fractures occur in the United States each year. Fractures of the humerus and other bones (estimated to be about 675,000 per year) also occur with osteoporosis; this is not surprising in light of the fact that bone loss is a systemic phenomenon. Although some fractures result from major trauma, the threshold for fracture is reduced for an osteoporotic bone (Fig. 425-3). In addition to bone density, there are a number of risk factors for fracture; the common ones are summarized in Table 425-1. Age, prior fractures (especially recent fractures), a family history of osteoporosis-related fractures, low body weight, smoking, and excessive alcohol use are all independent predictors of fracture. Chronic diseases with inflammatory components that increase skeletal remodeling such as rheumatoid arthritis, increase the risk of osteoporosis, as do

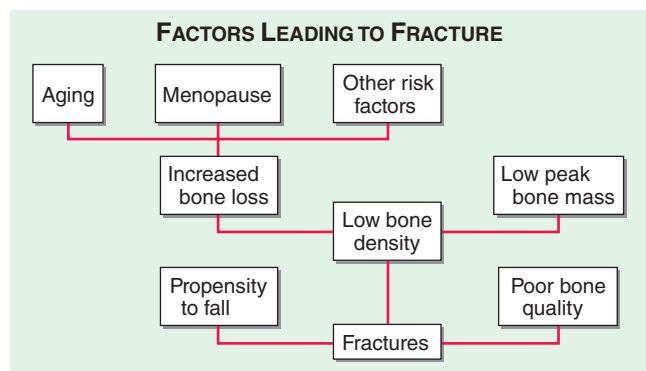


FIGURE 425-3 Factors leading to osteoporotic fractures.

diseases associated with malabsorption. Chronic diseases that increase the risk of falling or frailty, including dementia, Parkinson's disease, and multiple sclerosis, also increase fracture risk.

In the United States and Europe, osteoporosis-related fractures are more common among women than men, presumably due to a lower peak bone mass as well as postmenopausal bone loss in women. However, this sex difference in bone density and age-related increase in hip fractures is not as apparent in some other cultures, possibly due to genetics, physical activity level, or diet.

Fractures are themselves risk factors for future fractures (Table 425-1). Vertebral fractures increase the risk of other vertebral fractures as well as fractures of the peripheral skeleton such as the hip and wrist. Wrist fractures also increase the risk of vertebral and hip fractures. The risk for subsequent fractures is particularly high in the first several years after the first fracture, and the risk wanes considerably thereafter. Consequently, among individuals over age 50, any fracture should be considered as potentially related to osteoporosis regardless of the circumstances of the fracture. Osteoporotic bone is more likely to fracture than normal bone at any level of trauma, and a fracture in a person over 50 should trigger evaluation for osteoporosis. This often does not occur because postfracture care is not always well coordinated.

PATHOPHYSIOLOGY

BONE REMODELING

Osteoporosis results from bone loss due to age-related changes in bone remodeling as well as extrinsic and intrinsic factors that exaggerate this process. These changes may be superimposed on a low peak bone mass. Consequently, understanding the bone remodeling process is fundamental to understanding the pathophysiology of osteoporosis (Chap. 423). During growth, the skeleton increases in size by linear growth and by apposition of new bone tissue on the outer surfaces of the cortex (Fig. 425-4). The latter process is called *modeling*, a process that also allows the long bones to adapt in shape to the stresses placed on them. Increased sex hormone production at puberty is required for skeletal maturation, which reaches maximum mass and density in early adulthood. It is around puberty that the sexual dimorphism in skeletal size becomes obvious, although true bone density remains similar between the sexes. Nutrition and lifestyle also play an important role in growth, although genetic factors primarily determine peak skeletal mass and density. Numerous genes control skeletal growth, peak bone mass, and body size, as well as skeletal structure and density. Heritability estimates of 50–80% for bone density and size have been derived on the basis of twin studies. Although peak bone mass is often lower among individuals with a family history of osteoporosis, association studies of candidate genes (vitamin D receptor; type I collagen, the estrogen receptor [ER], and interleukin 6 [IL-6]; and insulin-like growth factor I [IGF-I]) and bone mass, bone turnover, and fracture prevalence have been inconsistent. Linkage studies suggest that a genetic locus on chromosome 11 is associated with high bone mass. Families with high bone mass and without much apparent age-related bone loss have been shown to have a point mutation in *LRP5*, a low-density lipoprotein receptor-related protein. The role of this gene in the general population is not clear, although a nonfunctional mutation results in osteoporosis-pseudoglioma syndrome, and *LRP5* signaling appears to be important in controlling bone formation. *LRP5* acts through the Wnt signaling pathway. With *LRP5* and Wnt activation, beta-catenin is translocated to the nucleus, allowing stimulation of osteoblast formation, activation, and life span as well as suppression of osteoclast activity, thereby increasing bone formation. The osteocyte product, sclerostin, is a negative inhibitor of Wnt signaling.

Genome-wide scans for low bone mass suggest multiple genes are involved, many of which are also implicated in control of body size.

In adults, bone remodeling, not modeling, is the principal metabolic skeletal process. Bone remodeling has two primary functions: (1) to repair microdamage within the skeleton to maintain skeletal strength and ensure the relative youth of the skeleton and (2) to supply calcium from the skeleton to maintain serum calcium. Remodeling may be

TABLE 425-1 CONDITIONS, DISEASES, AND MEDICATIONS THAT CONTRIBUTE TO OSTEOPOROSIS AND FRACTURES

Lifestyle factors		
Alcohol abuse	High salt intake	Falling
Low calcium intake	Inadequate physical activity	Excessive thinness
Vitamin D insufficiency	Immobilization	Prior fractures
Excess vitamin A	Smoking (active or passive)	
Genetic factors		
Cystic fibrosis	Homocystinuria	Osteogenesis imperfecta
Ehlers-Danlos syndrome	Hypophosphatasia	Parental history of hip fracture
Gaucher's disease	Idiopathic hypercalciuria	Porphyria
Glycogen storage diseases	Marfan's syndrome	Riley-Day syndrome
Hemochromatosis	Menkes' steely hair syndrome	
Hypogonadal states		
Androgen insensitivity	Hyperprolactinemia	Athletic amenorrhea
Anorexia nervosa and bulimia	Premature menopause	Panhypopituitarism
Turner's & Klinefelter's syndromes	Premature ovarian failure	
Endocrine disorders		
Adrenal insufficiency	Cushing's syndrome	Central adiposity
Diabetes mellitus (types 1 and 2)	Hyperparathyroidism	Thyrotoxicosis
Gastrointestinal disorders		
Celiac disease	Inflammatory bowel disease	Primary biliary cirrhosis
Gastric bypass	Malabsorption	
Gastrointestinal surgery	Pancreatic disease	
Hematologic disorders		
Multiple myeloma	Monoclonal gammopathies	Sickle cell disease
Hemophilia	Leukemia and lymphomas	Systemic mastocytosis
Thalassemia		
Rheumatologic and autoimmune diseases		
Ankylosing spondylitis	Lupus	Rheumatoid arthritis
Other rheumatic and autoimmune diseases		
Central nervous system disorders		
Epilepsy	Parkinson's disease	Stroke
Multiple sclerosis	Spinal cord injury	
Miscellaneous conditions and diseases		
AIDS/HIV	Congestive heart failure	Posttransplant bone disease
Alcoholism	Depression	Sarcoidosis
Amyloidosis	End-stage renal disease	Weight loss
Chronic metabolic acidosis	Hypercalciuria	
Chronic obstructive lung disease	Idiopathic scoliosis	
	Muscular dystrophy	
Medications		
Aluminum (in antacids)	Glucocorticoids (≥ 5 mg/d prednisone or equivalent for ≥ 3 months)	Tamoxifen (premenopausal use)
Anticoagulants (heparin)		Thiazolidinediones (such as pioglitazone and rosiglitazone)
Anticonvulsants	Gonadotropin-releasing hormone antagonists and agonists	Thyroid hormones (in excess)
Aromatase inhibitors	Lithium	Parenteral nutrition
Barbiturates	Methotrexate	
Cancer chemotherapeutic drugs	Proton pump inhibitors	
Cyclosporine A and tacrolimus	Selective serotonin reuptake inhibitors	
Depo-medroxyprogesterone (premenopausal contraception)		

Source: From the 2014 National Osteoporosis Foundation Clinician's Guide to the Prevention and Treatment of Osteoporosis. © National Osteoporosis Foundation.

activated by microdamage to bone as a result of excessive or accumulated stress. Acute demands for calcium involve osteoclast-mediated resorption as well as calcium transport by osteocytes. Chronic demands for calcium result in secondary hyperparathyroidism, increased bone remodeling, and overall loss of bone tissue.

Bone remodeling also is regulated by several circulating hormones, including estrogens, androgens, vitamin D, and parathyroid hormone (PTH), as well as locally produced growth factors such as IGF-I and immunoreactive growth hormone II (IGH-II), transforming growth factor β (TGF- β), parathyroid hormone-related peptide (PTHrP), interleukins (ILs), prostaglandins, and members of the tumor necrosis factor (TNF) superfamily. These factors primarily modulate the rate at which new remodeling sites are activated, a process that results initially in bone resorption by osteoclasts, followed by a period of repair during which new bone tissue is synthesized by osteoblasts. The cytokine responsible for communication between the osteoblasts, other marrow cells, and osteoclasts is RANK ligand (RANKL; receptor activator of nuclear factor- κ B [NF- κ B]). RANKL, a member of the TNF family, is secreted by osteoblasts and certain cells of the immune system (Chap. 423). The osteoclast receptor for this protein is referred to as RANK. Activation of RANK by RANKL is a final common path in osteoclast development, activation, and life span. A humoral decoy for RANKL, also secreted by osteoblasts, is *osteoprotegerin* (Fig. 425-5). Modulation of osteoclast recruitment and activity appears to be related to the interplay among these three factors. It appears that estrogens are pivotal in modulating secretion of osteoprotegerin (OPG) and perhaps also RANKL. Additional influences include nutrition (particularly calcium intake) and physical activity level.

In young adults, resorbed bone is replaced by an equal amount of new bone tissue. Thus, the mass of the skeleton remains constant after peak bone mass is achieved in adulthood. After age 30–45, however, the resorption and formation processes become imbalanced, and resorption exceeds formation. This imbalance may begin at different ages and varies at different skeletal sites; it becomes exaggerated in women after menopause. Excessive bone loss can be due to an increase in osteoclastic activity and/or a decrease in osteoblastic activity. In addition, an increase in remodeling activation frequency, and thus the number of remodeling sites, can magnify the small imbalance seen at each remodeling unit. Increased recruitment of bone remodeling sites produces a reversible reduction in bone tissue but also can result in permanent loss of tissue and disrupted skeletal architecture. In trabecular bone, if the osteoclasts penetrate trabeculae, they leave no template for new bone formation to occur, and, consequently, rapid bone loss ensues and cancellous connectivity becomes impaired. A higher number of remodeling sites increases the likelihood of this event. In cortical bone, increased activation of remodeling creates more porous bone. The effect of this increased porosity on cortical bone strength may be modest if the overall diameter of the bone is not changed. However, decreased apposition of new bone on the periosteal surface coupled with increased endocortical resorption of bone decreases the biomechanical strength of long bones. Even a slight exaggeration in normal bone loss increases the risk of osteoporosis-related fractures because of the architectural changes that occur, and osteoporosis is primarily a disease of disordered skeletal architecture. The main clinically available tool (dual-energy x-ray absorptiometry) measures mass not architecture. Emerging data from high-resolution peripheral quantitative computed tomography (CT) scans suggest that aging is associated with changes in microstructure of bone tissue, including increased cortical porosity and reduced cortical thickness.

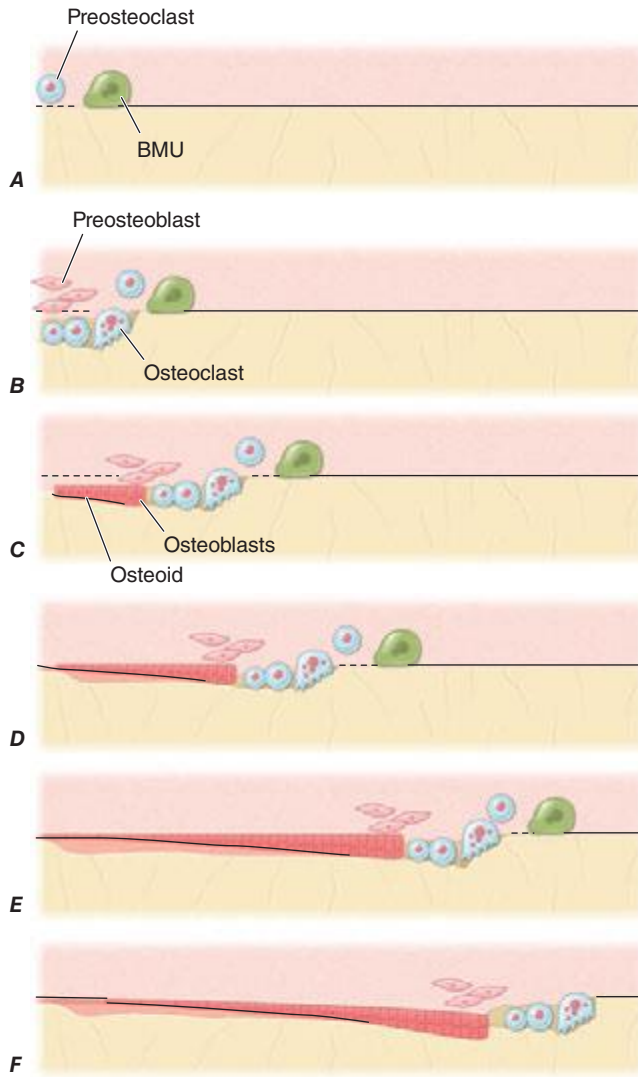


FIGURE 425-4 Mechanism of bone remodeling. The basic molecular unit (BMU) moves along the trabecular surface at a rate of about 10 $\mu\text{m}/\text{d}$. The figure depicts remodeling over ~ 120 days. **A.** Origination of BMU-lining cells contracts to expose collagen and attract preosteoclasts. **B.** Osteoclasts fuse into multinucleated cells that resorb a cavity. Mononuclear cells continue resorption, and preosteoblasts are stimulated to proliferate. **C.** Osteoblasts align at bottom of cavity and start forming osteoid (black). **D.** Osteoblasts continue formation and mineralization. Previous osteoid starts to mineralize (horizontal lines). **E.** Osteoblasts begin to flatten. **F.** Osteoblasts turn into lining cells; bone remodeling at initial surface (left of drawing) is now complete, but BMU is still advancing (to the right). (Adapted from SM Ott, in JP Bilezikian et al [eds]: *Principles of Bone Biology*, vol. 18. San Diego, Academic Press, 1996, pp 231–241.)

CALCIUM NUTRITION

Peak bone mass may be impaired by inadequate calcium intake during growth among other nutritional factors (calories, protein, and other minerals), leading to increased risk of osteoporosis later in life. During the adult phase of life, insufficient calcium intake contributes to relative secondary hyperparathyroidism and an increase in the rate of bone remodeling to maintain normal serum calcium levels. PTH stimulates the hydroxylation of vitamin D in the kidney, leading to increased levels of 1,25-dihydroxyvitamin D [$1,25(\text{OH})_2\text{D}$] and enhanced gastrointestinal calcium absorption. PTH also reduces renal calcium loss. Although these are all appropriate compensatory homeostatic responses for adjusting calcium economy, the long-term effects are detrimental to the skeleton because the increased remodeling rates and

the ongoing imbalance between resorption and formation at remodeling sites combine to accelerate loss of bone tissue.

Total daily calcium intakes < 400 mg are detrimental to the skeleton, and intakes in the range of 600–800 mg, which is about the average intake among adults in the United States, are also probably suboptimal. The recommended daily required intake of 1000–1200 mg for adults accommodates population heterogeneity in controlling calcium balance (Chap. 95e). Such intakes should preferentially come from dietary sources, and supplements should be used only when dietary intakes fall short. The supplement should contain enough calcium to bring total intake to about 1200 mg/d.

VITAMIN D

(See also Chap. 423) Severe vitamin D deficiency causes rickets in children and osteomalacia in adults. However, there is accumulating evidence that vitamin D insufficiency may be more prevalent than previously thought, particularly among individuals at increased risk such as the elderly; those living in northern latitudes; and individuals with poor nutrition, malabsorption, or chronic liver or renal disease. Dark-skinned individuals are also at high risk of vitamin D deficiency. There is controversy regarding optimal levels of serum 25-hydroxyvitamin D [$25(\text{OH})\text{D}$], with some advocating levels > 20 ng/mL and others advocating optimal targets > 75 nmol/L (30 ng/mL). To achieve this level for most adults requires an intake of 800–1000 units/d, particularly in individuals who avoid sunlight or routinely use ultraviolet-blocking lotions. Vitamin D insufficiency leads to compensatory secondary hyperparathyroidism and is an important risk factor for osteoporosis and fractures. Some studies have shown that $> 50\%$ of inpatients on a general medical service exhibit biochemical features of vitamin D deficiency, including increased levels of PTH and alkaline phosphatase and lower levels of ionized calcium. In women living in northern latitudes, vitamin D levels decline during the winter months. This is associated with seasonal bone loss, reflecting increased bone turnover. Even among healthy ambulatory individuals, mild vitamin D deficiency is increasing in prevalence, in part due to decreased exposure to sunlight coupled with increased use of potent sunscreens. Treatment with vitamin D can return levels to normal and prevent the associated increase in bone remodeling, bone loss, and fractures. Improved muscle function and gait associated with reduced falls and fracture rates also have been documented among individuals in northern latitudes who have greater vitamin D intake and higher $25(\text{OH})\text{D}$ levels (see below). Vitamin D adequacy also may affect risk and/or severity of other diseases, including cancers (colorectal, prostate, and breast), autoimmune diseases, and diabetes; however, many observational studies suggesting these potential extraskeletal benefits have not been confirmed with randomized controlled trials.

ESTROGEN STATUS

Estrogen deficiency probably causes bone loss by two distinct but interrelated mechanisms: (1) activation of new bone remodeling sites and (2) exaggeration of the imbalance between bone formation and resorption. The change in activation frequency causes a transient bone loss until a new steady state between resorption and formation is achieved. The remodeling imbalance, however, results in a permanent decrement in mass. In addition, the very presence of more remodeling sites in the skeleton increases the probability that trabeculae will be penetrated, eliminating the template on which new bone can be formed and accelerating the loss of bony tissue.

The most common estrogen-deficient state is the cessation of ovarian function at the time of menopause, which occurs on average at age 51 (Chap. 413). Thus, with current life expectancy, an average woman will spend about 30 years without an ovarian supply of estrogen. The mechanism by which estrogen deficiency causes bone loss is summarized in Fig. 425-5. Marrow cells (macrophages, monocytes, osteoclast precursors, mast cells) as well as bone cells (osteoblasts, osteocytes, osteoclasts) express ERs α and β . Loss of estrogen increases production of RANKL and may reduce production of OPG, increasing osteoclast recruitment. Estrogen also may play an important role in determining the life span of bone cells by controlling the rate of apoptosis. Thus,

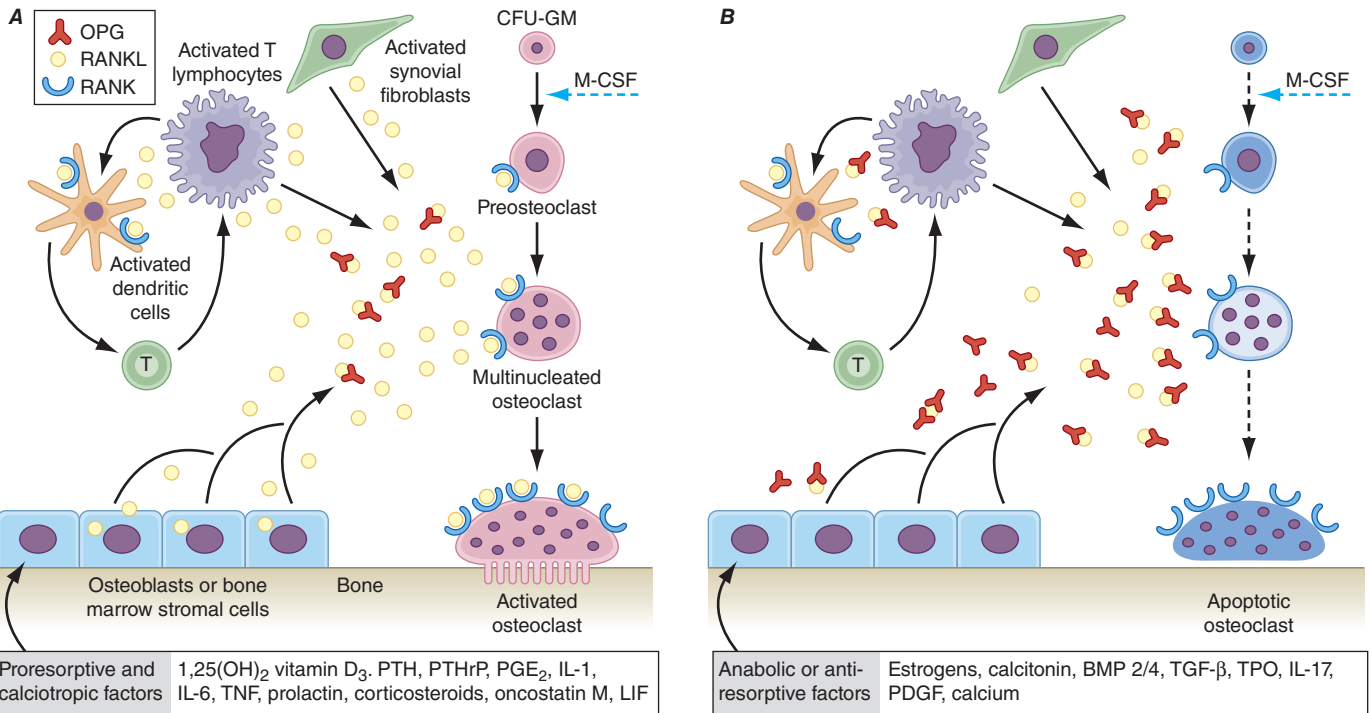


FIGURE 425-5 Hormonal control of bone resorption. A. Proresorptive and calciotropic factors. **B.** Anabolic and antiosteoclastic factors. RANK ligand (RANKL) expression is induced in osteoblasts, activated T cells, synovial fibroblasts, and bone marrow stromal cells. It binds to membrane-bound receptor RANK to promote osteoclast differentiation, activation, and survival. Conversely, osteoprotegerin (OPG) expression is induced by factors that block bone catabolism and promote anabolic effects. OPG binds and neutralizes RANKL, leading to a block in osteoclastogenesis and decreased survival of preexisting osteoclasts. CFU-GM, colony-forming units, granulocyte macrophage; IL, interleukin; LIF, leukemia inhibitory factor; M-CSF, macrophage colony-stimulating factor; OPG-L, osteoprotegerin-ligand; PDGF, platelet-derived growth factor; PGE₂, prostaglandin E₂; PTH, parathyroid hormone; RANKL, receptor activator of nuclear factor nuclear factor-κB; TGF-β, transforming growth factor β; TNF, tumor necrosis factor; TPO, thrombospondin. (From WJ Boyle et al: *Nature* 423: 337, 2003.)

in situations of estrogen deprivation, the life span of osteoblasts may be decreased, whereas the longevity and activity of osteoclasts are increased. The rate and duration of bone loss after menopause are heterogeneous and unpredictable. Once surfaces are lost in cancellous bone, the rate of bone loss must decline. In cortical bone, loss is slower but continues for a longer time period.

Because remodeling is initiated at the surface of bone, it follows that trabecular bone—which has a considerably larger surface area (80% of the total) than cortical bone—will be affected preferentially by estrogen deficiency. Fractures occur earliest at sites where trabecular bone contributes most to bone strength; consequently, vertebral fractures are the most common early consequence of estrogen deficiency.

PHYSICAL ACTIVITY

Inactivity, such as prolonged bed rest or paralysis, results in significant bone loss. Concordantly, athletes have higher bone mass than does the general population. These changes in skeletal mass are most marked when the stimulus begins during growth and before the age of puberty. Adults are less capable than children of increasing bone mass after restoration of physical activity. Epidemiologic data support the beneficial effects on the skeleton of chronic high levels of physical activity. Fracture risk is lower in rural communities and in countries where physical activity is maintained into old age. However, when exercise is initiated during adult life, the effects of moderate exercise on the skeleton are modest, with a bone mass increase of 1–2% in short-term studies of <2 years in duration. It is argued that more active individuals are less likely to fall and are more capable of protecting themselves upon falling, thereby reducing fracture risk.

CHRONIC DISEASE

Various genetic and acquired diseases are associated with an increase in the risk of osteoporosis (Table 425-1). Mechanisms that contribute to bone loss are unique for each disease and typically result from multiple factors, including nutrition, reduced physical activity levels,

and factors that affect rates of bone remodeling. In most, but not all, circumstances the primary diagnosis is made before osteoporosis presents clinically.

MEDICATIONS

A large number of medications used in clinical practice have potentially detrimental effects on the skeleton (Table 425-1). *Glucocorticoids* are the most common cause of medication-induced osteoporosis. It is often not possible to determine the extent to which osteoporosis is related to glucocorticoids or to other factors, because treatment is superimposed on the effects of the primary disease, which in itself may be associated with bone loss (e.g., rheumatoid arthritis). Excessive doses of thyroid hormone can accelerate bone remodeling and result in bone loss.

Other medications have less detrimental effects on the skeleton than pharmacologic doses of glucocorticoids. *Anticonvulsants* are thought to increase the risk of osteoporosis, although many affected individuals have concomitant insufficiency of 1,25(OH)₂D, as some anticonvulsants induce the cytochrome P450 system and vitamin D metabolism. Patients undergoing transplantation are at high risk for rapid bone loss and fracture not only from glucocorticoids but also from treatment with other *immunosuppressants* such as cyclosporine and tacrolimus (FK506). In addition, these patients often have underlying metabolic abnormalities, such as hepatic or renal failure, that predispose to bone loss.

Aromatase inhibitors, which potently block the aromatase enzyme that converts androgens and other adrenal precursors to estrogen, reduce circulating postmenopausal estrogen levels dramatically. These agents, which are used in various stages for breast cancer treatment, also have been shown to have a detrimental effect on bone density and risk of fracture. More recently a variety of agents have been implicated in increased bone loss and fractures. These include selective serotonin reuptake inhibitors, proton pump inhibitors, and thiazolidinediones. It is difficult in some cases to separate the risk

accrued by the underlying disease from that attributable to the medication. For example, both depression and diabetes are risk factors for fracture by themselves.

CIGARETTE CONSUMPTION

The use of cigarettes over a long period has detrimental effects on bone mass. These effects may be mediated directly by toxic effects on osteoblasts or indirectly by modifying estrogen metabolism. On average, cigarette smokers reach menopause 1–2 years earlier than the general population. Cigarette smoking also produces secondary effects that can modulate skeletal status, including intercurrent respiratory and other illnesses, frailty, decreased exercise, poor nutrition, and the need for additional medications (e.g., glucocorticoids for lung disease).

MEASUREMENT OF BONE MASS

Several noninvasive techniques are available for estimating skeletal mass or density. They include dual-energy x-ray absorptiometry (DXA), single-energy x-ray absorptiometry (SXA), quantitative CT, and ultrasound (US). DXA is a highly accurate x-ray technique that has become the standard for measuring bone density. Although it can be used for measurement in any skeletal site, clinical determinations usually are made of the lumbar spine and hip. DXA also can be used to measure body composition. In the DXA technique, two x-ray energies are used to estimate the area of mineralized tissue, and the mineral content is divided by the area, which partially corrects for body size. However, this correction is only partial because DXA is a two-dimensional scanning technique and cannot estimate the depth or posteroanterior length of the bone. Thus, small slim people tend to have lower than average bone mineral density (BMD), a feature that is important in interpreting BMD measurements when performed in young adults, and something that must be taken into account at any age. Bone spurs, which are common in osteoarthritis, tend to falsely increase bone density of the spine and are a particular problem in measuring the spine in older individuals. Because DXA instrumentation is provided by several different manufacturers, the output varies in absolute terms. Consequently, it has become standard practice to relate the results to “normal” values by using T-scores (a T-score of 1 equals 1 SD), which compare individual results to those in a young population that is matched for race and sex. Z-scores (also measured in SD) compare individual results to those of an age-matched population that also is matched for race and sex. Thus, a 60-year-old woman with a Z-score of -1 (1 SD below mean for age) has a T-score of -2.5 (2.5 SD below mean for a young control group) (Fig. 425-6). A T-score below -2.5 in the lumbar spine, femoral neck, or total hip has been defined as a diagnosis of osteoporosis. As noted above, because more than 50% of fractures occur in individuals with low bone mass rather than BMD osteoporosis, attempts are ongoing to redefine the disease as a fracture risk rather than a specific BMD. Consistent with this concept, fractures of the spine and hip that occur in the absence of major trauma would be considered to be sufficient to diagnose osteoporosis, regardless of BMD. Fractures of other sites, such as pelvis, proximal humerus, and wrist, would be tantamount to an osteoporosis diagnosis in the presence of low BMD. CT can also be used to measure the spine and the

hip, but is rarely used clinically, in part because of higher radiation exposure and cost, in addition to a lesser body of data confirming its ability to predict fracture risk, compared with BMD by DXA. High-resolution peripheral CT is used to measure bone in the forearm or tibia as a research tool to noninvasively provide some measure of skeletal architecture. Magnetic resonance imaging (MRI) can also be used in research settings to obtain some architectural information on the forearm and perhaps the hip.

DXA equipment can also be used to obtain lateral images of the spine, from T4 through L4, a technique called vertebral fracture assessment (VFA). Although not as definitive as radiography, it is a useful screening tool when height loss, back pain, or postural change suggests the presence of an undiagnosed vertebral fracture. Furthermore, because vertebral fractures are so prevalent with advancing age, screening vertebral imaging is recommended in women and men with low bone mass (T-score <1) by age 70 and 80, respectively.

US is used to measure bone mass by calculating the attenuation of the signal as it passes through bone or the speed with which it traverses the bone. It is unclear whether US assesses properties of bone other than mass (e.g., quality), but this is a potential advantage of the technique. Because of its relatively low cost and mobility, US is amenable for use as a screening procedure in stores or at health fairs.

All of these techniques for measuring BMD have been approved by the U.S. Food and Drug Administration (FDA) on the basis of their capacity to predict fracture risk. The hip is the preferred site of measurement in most individuals, because it predicts the risk of hip fracture, the most important consequence of osteoporosis, better than any other bone density measurement site. When hip measurements are performed by DXA, the spine can be measured at the same time. In younger individuals such as perimenopausal or early postmenopausal women, spine measurements may be the most sensitive indicator of bone loss. A risk assessment tool (FRAX) incorporates femoral neck BMD to assess 10-year fracture risk (see below).

WHEN TO MEASURE BONE MASS

Clinical guidelines have been developed for the use of bone densitometry in clinical practice. The original National Osteoporosis Foundation guidelines recommend bone mass measurements in postmenopausal women, assuming they have one or more risk factors for osteoporosis in addition to age, sex, and estrogen deficiency. The guidelines further recommend that bone mass measurement be considered in *all* women by age 65, a position ratified by the U.S. Preventive Health Services Task Force. Criteria approved for Medicare reimbursement of BMD are summarized in Table 425-2.

WHEN TO TREAT BASED ON BONE MASS RESULTS

Most guidelines suggest that patients be considered for treatment when BMD is >2.5 SD below the mean value for young adults (T-score ≤ -2.5), in either spine, total hip, or femoral neck. Treatment also should also be considered in postmenopausal women with fracture risk factors even if BMD is not in the osteoporosis range. Risk factors (age, prior fracture, family history of hip fracture, low body weight, cigarette consumption, excessive alcohol use, steroid use, and rheumatoid arthritis) can be combined with BMD to assess the likelihood of a

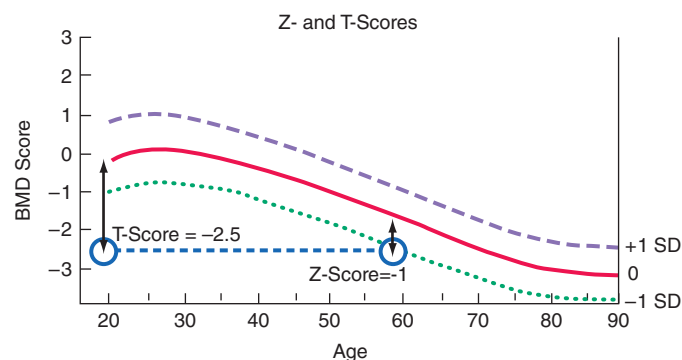


FIGURE 425-6 Relationship between Z-scores and T-scores in a 60-year-old woman. BMD, bone mineral density; SD, standard deviation.

TABLE 425-2 INDICATIONS FOR BONE DENSITY TESTING

Consider BMD testing in the following individuals:

- Women age 65 and older and men age 70 and older, regardless of clinical risk factors
- Younger postmenopausal women, women in the menopausal transition and men age 50–69 with clinical risk factors for fracture
- Adults who have a fracture after age 50
- Adults with a condition (e.g., rheumatoid arthritis) or taking a medication (e.g., glucocorticoids in a daily dose ≥ 5 mg prednisone or equivalent for ≥ 3 months) associated with low bone mass or bone loss

Source: From the 2014 National Osteoporosis Foundation Clinician's Guide to the Prevention and Treatment of Osteoporosis. © National Osteoporosis Foundation.

2494 fracture over a 5- or 10-year period. Treatment threshold depends on cost-effectiveness analyses but probably is ~1% per year of risk in the United States.

APPROACH TO THE PATIENT: Osteoporosis

The perimenopausal transition is a good opportunity to initiate a discussion about risk factors for osteoporosis and consideration of indications for a BMD test. A careful history and physical examination should be performed to identify risk factors for osteoporosis. A low Z-score increases the suspicion of a secondary disease. Height loss >2.5–3.8 cm (>1–1.5 in.) is an indication for VFA by DXA or radiography to rule out asymptomatic vertebral fractures, as is the presence of significant kyphosis or back pain, particularly if it began after menopause. In appropriate individuals, screening BMD and screening vertebral imaging should be recommended as above, even in the absence of any specific risk factors (Table 425-3). For patients who present with fractures, it is important to ensure that the fractures are not caused by an underlying malignancy. Usually this is clear on routine radiography, but on occasion, CT, MRI, or radionuclide scans may be necessary.

ROUTINE LABORATORY EVALUATION

There is no established algorithm for the evaluation of women who present with osteoporosis. A general evaluation that includes complete blood count, serum and 24-h urine calcium, renal and hepatic function tests, and a 25(OH)D level is useful for identifying selected secondary causes of low bone mass, particularly for women with fractures or very low Z-scores. An elevated serum calcium level suggests hyperparathyroidism or malignancy, whereas a reduced serum calcium level may reflect malnutrition and osteomalacia. In the presence of hypercalcemia, a serum PTH level differentiates between hyperparathyroidism (PTH↑) and malignancy (PTH↓), and a high PTHrP level can help document the presence of humoral hypercalcemia of malignancy (Chap. 424). A low urine calcium (<50 mg/24 h) suggests osteomalacia, malnutrition, or malabsorption; a high urine calcium (>300 mg/24 h) is indicative of hypercalciuria and must be investigated further. Hypercalciuria occurs primarily in three situations: (1) a renal calcium leak, which is more common in males with osteoporosis; (2) absorptive hypercalciuria, which can be idiopathic or associated with increased 1,25(OH)₂D in granulomatous disease; or (3) hematologic malignancies or conditions associated with excessive bone turnover such as Paget's disease, hyperparathyroidism, and hyperthyroidism. Renal hypercalciuria is treated with thiazide diuretics, which lower urine calcium and help improve calcium economy.

Individuals who have osteoporosis-related fractures or bone density in the osteoporotic range should have a measurement of serum 25(OH)D level, because the intake of vitamin D required to achieve a target level >20–30 ng/mL is highly variable. Vitamin D levels should be optimized in all individuals being treated for

osteoporosis. Hyperthyroidism should be evaluated by measuring thyroid-stimulating hormone (TSH).

When there is clinical suspicion of Cushing's syndrome, urinary free cortisol levels or a fasting serum cortisol should be measured after overnight dexamethasone. When bowel disease, malabsorption, or malnutrition is suspected, serum albumin, cholesterol, and a complete blood count should be checked. Asymptomatic malabsorption may be heralded by anemia (macrocytic—vitamin B₁₂ or folate deficiency; microcytic—iron deficiency) or low serum cholesterol or urinary calcium levels. If these or other features suggest malabsorption, further evaluation is required. Asymptomatic celiac disease with selective malabsorption is being found with increasing frequency; the diagnosis can be made by testing for antigliadin, antiendomysial, or transglutaminase antibodies but may require endoscopic biopsy. A trial of a gluten-free diet can be confirmatory (Chap. 349). When osteoporosis is found associated with symptoms of rash, multiple allergies, diarrhea, or flushing, mastocytosis should be excluded by using 24-h urine histamine collection or serum tryptase.

Myeloma can masquerade as generalized osteoporosis, although it more commonly presents with bone pain and characteristic “punched-out” lesions on radiography. Serum and urine electrophoresis and or serum free light chains are required to exclude this diagnosis. More commonly, a monoclonal gammopathy of unclear significance (MGUS) is found, and the patient is subsequently monitored to ensure that this is not an incipient myeloma. Approximately 1% of patients with MGUS progress to myeloma each year. A bone marrow biopsy may be required to rule out myeloma (in patients with equivocal electrophoretic results) and also can be used to exclude mastocytosis, leukemia, and other marrow infiltrative disorders such as Gaucher's disease. MGUS syndromes, although benign, may also be associated with reduced bone mass and elevated bone turnover.

BONE BIOPSY

Tetracycline labeling of the skeleton allows determination of the rate of remodeling as well as evaluation for other metabolic bone diseases. The current use of BMD tests, in combination with hormonal evaluation and biochemical markers of bone remodeling, has largely replaced the clinical use of bone biopsy, although it remains an important tool in clinical research and assessment of mechanism of action of medication for osteoporosis.

BIOCHEMICAL MARKERS

Several biochemical tests are available that provide an index of the overall rate of bone remodeling (Table 425-4). Biochemical markers usually are characterized as those related primarily to *bone formation* or *bone resorption*. These tests measure the overall state of bone remodeling at a single point in time. Clinical use of these tests has been hampered by biologic variability (in part related to circadian rhythm) as well as analytic variability, although the latter is improving.

TABLE 425-3 INDICATIONS FOR VERTEBRAL IMAGING

Consider vertebral imaging tests in the following individuals:

- In all women age 70 and older and all men age 80 and older if bone mineral density (BMD) T-score is –1.0 or below
- In women age 65–69 and men age 75–79 if BMD T-score is –1.5 or below
- In postmenopausal women age 50–64 and men age 50–69 with specific risk factors:
 - Low-trauma fracture
 - Historical height loss of 1.5 in. or more (4 cm)
 - Prospective height loss of 0.8 in. or more (2 cm)
 - Recent or ongoing long-term glucocorticoid treatment

Source: From the 2014 National Osteoporosis Foundation Clinician's Guide to the Prevention and Treatment of Osteoporosis. © National Osteoporosis Foundation.

TABLE 425-4 INDICATIONS FOR BIOCHEMICAL MARKERS

Biochemical markers of bone turnover may:

- Predict risk of fracture independently of bone density.
- Predict extent of fracture risk reduction when repeated after 3–6 months of treatment with FDA-approved therapies.
- Predict magnitude of BMD increases with FDA-approved therapies.
- Predict rapidity of bone loss.
- Help determine adequacy of patient compliance and persistence with osteoporosis therapy.
- Help determine duration of “drug holiday” (data are quite limited to support this use, but studies are under way).

Abbreviations: BMD, bone mineral density; FDA, U.S. Food and Drug Administration.

Source: Adapted from the 2014 National Osteoporosis Foundation Clinician's Guide to the Prevention and Treatment of Osteoporosis. © National Osteoporosis Foundation.

Biochemical markers of bone resorption may help in the prediction of fracture risk, independently of bone density, particularly in older individuals. In women ≥ 65 years, when bone density results are greater than the usual treatment thresholds noted above, a high level of bone resorption should prompt consideration of treatment. The primary use of biochemical markers is for monitoring the response to treatment. With the introduction of antiresorptive therapeutic agents, bone remodeling declines rapidly, with the fall in resorption occurring earlier than the fall in formation. Inhibition of bone resorption is maximal within 3 months or so. Thus, measurement of bone resorption (C-telopeptide [CTX] is the preferred marker) before initiating therapy and 3–6 months after starting therapy provides an earlier estimate of patient response than does bone densitometry. A decline in resorptive markers can be ascertained after treatment with potent antiresorptive agents such as bisphosphonates, denosumab, or standard-dose estrogen; this effect is less marked after treatment with weaker agents such as raloxifene or intranasal calcitonin. A biochemical marker response to therapy is particularly useful for asymptomatic patients and may help ensure long-term adherence to treatment. Bone turnover markers are also useful in monitoring the effects of osteoanabolic agents such as 1-34hPTH, or teriparatide, which rapidly increases bone formation (PINP is preferred, but osteocalcin is a reasonable alternative) and later bone resorption. The recent suggestion of “drug holidays” (see below) has created another use for biochemical markers, allowing evaluation of the off effect of drugs such as bisphosphonates.

TREATMENT OSTEOPOROSIS

MANAGEMENT OF PATIENTS WITH FRACTURES

Treatment of a patient with osteoporosis frequently involves management of acute fractures as well as treatment of the underlying disease. Hip fractures almost always require surgical repair if the patient is to become ambulatory again. Depending on the location and severity of the fracture, condition of the neighboring joint, and general status of the patient, procedures may include open reduction and internal fixation with pins and plates, hemiarthroplasties, and total arthroplasties. These surgical procedures are followed by intense rehabilitation in an attempt to return patients to their prefracture functional level. Long bone fractures (e.g., wrist) often require either external or internal fixation. Other fractures (e.g., vertebral, rib, and pelvic fractures) usually are managed with supportive care, requiring no specific orthopedic treatment.

Only ~25–30% of vertebral compression fractures present with sudden-onset back pain. For acutely symptomatic fractures, treatment with analgesics is required, including nonsteroidal anti-inflammatory agents and/or acetaminophen, sometimes with the addition of a narcotic agent (codeine or oxycodone). A few small, randomized clinical trials suggest that calcitonin may reduce pain related to acute vertebral compression fracture. Percutaneous injection of artificial cement (polymethylmethacrylate) into the vertebral body (vertebroplasty or kyphoplasty) may offer significant immediate pain relief in patients with severe pain from acute or subacute vertebral fractures. Safety concerns include extravasation of cement with neurologic sequelae and increased risk of fracture in neighboring vertebrae due to mechanical rigidity of the treated bone. Exactly which patients are the optimal candidates for this procedure remains unknown. Short periods of bed rest may be helpful for pain management, but in general, early mobilization is recommended because it helps prevent further bone loss associated with immobilization. Occasionally, use of a soft elastic-style brace may facilitate earlier mobilization. Muscle spasms often occur with acute compression fractures and can be treated with muscle relaxants and heat treatments.

Severe pain usually resolves within 6–10 weeks. More chronic severe pain might suggest the possibility of multiple myeloma or underlying metastatic disease. Chronic pain following vertebral

fracture is probably not bony in origin; instead, it is related to abnormal strain on muscles, ligaments, and tendons and to secondary facet-joint arthritis associated with alterations in thoracic and/or abdominal shape. Chronic pain is difficult to treat effectively and may require analgesics, sometimes including narcotic analgesics. Frequent intermittent rest in a supine or semireclining position is often required to allow the soft tissues, which are under tension, to relax. Back-strengthening exercises (paraspinal) may be beneficial. Heat treatments help relax muscles and reduce the muscular component of discomfort. Various physical modalities, such as US and transcutaneous nerve stimulation, may be beneficial in some patients. Pain also occurs in the neck region, not as a result of compression fractures (which almost never occur in the cervical spine as a result of osteoporosis) but because of chronic strain associated with trying to elevate the head in a person with a significant thoracic kyphosis.

Multiple vertebral fractures often are associated with psychological symptoms; this is not always appreciated. The changes in body configuration and back pain can lead to marked loss of self-image and a secondary depression. Altered balance, precipitated by the kyphosis and the anterior movement of the body's center of gravity, leads to a fear of falling, a consequent tendency to remain indoors, and the onset of social isolation. These symptoms sometimes can be alleviated by family support and/or psychotherapy. Medication may be necessary when depressive features are present. Multiple thoracic vertebral fractures may be associated with restrictive lung disease symptoms and increased pulmonary infections. Multiple lumbar vertebral fractures are often associated with abdominal pain, constipation, protuberance, and early satiety. Multiple vertebral fractures are associated with greater age-specific mortality.

Multiple studies show that the majority of patients presenting in adulthood with fractures are not evaluated or treated for osteoporosis. Estimates suggest only about 20% of fracture patients receive follow-up care. Patients who sustain acute fractures are at dramatically elevated risk for more fractures, particularly within the first several years, and pharmacologic intervention can reduce that risk substantially. Recently, several studies have demonstrated the effectiveness of a relatively simple and inexpensive program that reduces the risk of subsequent fractures. In the Kaiser system, it is estimated that a 20% decline in hip fracture occurrence was seen with the introduction of what is called a fracture liaison service. This typically involves a health care professional (usually a nurse) whose job is to coordinate follow-up care and education of fracture patients. If the Kaiser experience can be repeated, there would be significant savings of health care dollars, as well as a dramatic drop in hip fracture incidence and a marked improvement in morbidity and mortality among the aging population.

MANAGEMENT OF THE UNDERLYING DISEASE

Patients presenting with typical osteoporosis-related fractures (certainly hip and spine) can be assumed to have osteoporosis and can be treated appropriately. Patients with osteoporosis by BMD are handled in a similar fashion. Other fracture patients and those with reduced bone mass can be classified according to their future risk of fracture and treated if that risk is sufficiently high. It must be emphasized, however, that risk assessment is an inexact science when applied to individual patients. Fractures are chance occurrences that can happen to anyone. Patients often do not understand the relative benefits of medications, compared to the perceived risks of the medications themselves.

Risk Factor Reduction Several tools exist for risk assessment. The most commonly available is the FRAX tool, developed by a working party for the WHO, and available as part of the report from many DXA machines. It is also available online (<http://www.shef.ac.uk/FRAX/tool.jsp?locationValue=9>) (Fig. 425-7). In the United States, it has been estimated that it is cost-effective to treat a patient if the 10-year major fracture risk (including hip, clinical spine, proximal humerus, and tibia) from FRAX is $\geq 20\%$ and/or the 10-year risk of hip fracture is $\geq 3\%$. FRAX is an imperfect tool because it does not include

FRAX - WHO Fracture Risk Assessment Tool - Windows Internet Explorer

http://www.shef.ac.uk/FRAX/tool.jsp?locationValue=9

File Edit View Favorites Tools Help

Google frax fracture risk

Search

207 New

FRAX - WHO Fracture Risk Assessment Tool

Calculation Tool

Please answer the questions below to calculate the ten year probability of fracture with BMD.

Country: US(Caucasian) Name / ID: About the risk factors ⓘ

Questionnaire:

1. Age (between 40-90 years) or Date of birth
Age: Y: M: D:

2. Sex Male Female

3. Weight (kg)

4. Height (cm)

5. Previous fracture No Yes

6. Parent fractured hip No Yes

7. Current smoking No Yes

8. Glucocorticoids No Yes

9. Rheumatoid arthritis No Yes

10. Secondary osteoporosis No Yes

11. Alcohol 3 more units per day No Yes

12. Femoral neck BMD
T-score

Clear Calculate

BMI 22.6
The ten year probability of fracture (%)
with BMD

Major osteoporotic	13
Hip fracture	1.7

Weight Conversion: pound: convert 140 pound = 63.50 kg

Height Conversion: inch: convert 66 inch = 167.64 cm

FIGURE 425-7 FRAX calculation tool. When the answers to the indicated questions are filled in, the calculator can be used to assess the 10-year probability of fracture. The calculator (available online at <http://www.shef.ac.uk/FRAX/tool.jsp?locationValue=9>) also can risk adjust for various ethnic groups.

any assessment of fall risk and secondary causes are excluded when BMD is entered. Moreover, it does not include any term for multiple fractures or recent versus remote fracture. Nonetheless, it is useful as an educational tool for patients.

After risk assessment, patients should be thoroughly educated to reduce the impact of modifiable risk factors associated with bone loss and falling. All medications that increase risk of falls, bone loss, or fractures should be reviewed to ensure that they are necessary and being used at the lowest required dose. For those on thyroid hormone replacement, TSH testing should be performed to confirm that an excessive dose is not being used, because biochemical and symptomatic thyrotoxicosis can be associated with increased bone loss. In patients who smoke, efforts should be made to facilitate smoking cessation. Reducing risk factors for falling also include alcohol abuse treatment and a review of the medical regimen for any drugs that might be associated with orthostatic hypotension and/or sedation, including hypnotics and anxiolytics. If nocturia occurs, the frequency should be reduced, if possible (e.g., by decreasing or modifying diuretic use), because arising in the middle of sleep is a common precipitant of a fall. Patients should be instructed about environmental safety with regard to eliminating exposed wires, curtain strings, slippery rugs, and mobile tables. Avoiding stocking feet on wood floors, checking carpet condition (particularly on stairs), and providing good light in paths to bathrooms and outside the home are important preventive measures. Treatment for impaired vision is recommended, particularly a problem with depth perception, which is specifically associated with increased

falling risk. Elderly patients with neurologic impairment (e.g., stroke, Parkinson's disease, Alzheimer's disease) are particularly at risk of falling and require specialized supervision and care.

Nutritional Recommendations • CALCIUM A large body of data indicates that optimal calcium intake reduces bone loss and suppresses bone turnover. Recommended intakes from an Institute of Medicine report are shown in [Table 425-5](#).

The National Health and Nutrition Examination Surveys (NHANES) have consistently documented that average calcium intakes fall considerably short of these recommendations. Food sources of calcium are dairy products (milk, yogurt, and cheese) and fortified

TABLE 425-5 ADEQUATE CALCIUM INTAKE

Life Stage Group	Estimated Adequate Daily Calcium Intake, mg/d
Young children (1–3 years)	500
Older children (4–8 years)	800
Adolescents and young adults (9–18 years)	1300
Men and women (19–50 years)	1000
Men and women (51 and older)	1200

Note: Pregnancy and lactation needs are the same as for nonpregnant women (e.g., 1300 mg/d for adolescents/young adults and 1000 mg/d for ≥19 years).

Source: Adapted from the Standing Committee on the Scientific Evaluation of Dietary Reference Intakes. Food and Nutrition Board. Institute of Medicine. Washington, DC, 1997. National Academy Press.

TABLE 425-6 SIMPLE METHOD FOR CALCULATING DIETARY CALCIUM INTAKE**STEP 1:** Estimate calcium intake from calcium-rich foods

Product	# of Servings/d	Estimated calcium/serving, in mg	Calcium in mg
Milk (8 oz.)	_____	× 300	= _____
Yogurt (6 oz.)	_____	× 300	= _____
Cheese (1 oz. or 1 cubic in.)	_____	× 200	= _____
Fortified foods or juices	_____	× 80 to 1000	= _____
Subtotal = _____			
STEP 2: Total from above + 250 mg for nondairy sources			
= total dietary calcium		TOTAL Calcium, in mg = _____	

Source: Adapted from SM Krane, MF Holick, **Chap. 355**, in *Harrison's Principles of Internal Medicine*, 14th ed. New York, McGraw-Hill, 1998.

foods such as certain cereals, waffles, snacks, juices, and crackers. Some of these fortified foods contain as much calcium per serving as milk. Green leafy vegetables and nuts, particularly almonds, are also sources of calcium, although their bioavailability may be lower than with dairy products. Calcium intake from the diet can also be assessed (**Table 425-6**) and calculators are available at *NOF.org* or *NYSOPEP.org*.

If a calcium supplement is required, it should be taken in doses sufficient to supplement dietary intake to bring total intake to the required level (1000–1200 mg/d). Doses of supplements should be ≤600 mg at a time, because the calcium absorption fraction decreases at higher doses. Calcium supplements should be calculated on the basis of the elemental calcium content of the supplement, not the weight of the calcium salt. Calcium supplements containing carbonate are best taken with food because they require acid for solubility. Calcium citrate supplements can be taken at any time. To confirm bioavailability, calcium supplements can be placed in distilled vinegar. They should dissolve within 30 min.

Several controlled clinical trials of calcium, mostly plus vitamin D, have confirmed reductions in clinical fractures, including fractures of the hip (~20–30% risk reduction). All recent studies of pharmacologic agents have been conducted in the context of calcium replacement (± vitamin D). Thus, it is standard practice to ensure an adequate calcium and vitamin D intake in patients with osteoporosis whether they are receiving additional pharmacologic therapy or not. A systematic review confirmed a greater BMD response to antiresorptive therapy when calcium intake was adequate.

Although side effects from supplemental calcium are minimal (eructation and constipation mostly with carbonate salts), individuals with a history of kidney stones should have a 24-h urine calcium determination before starting increased calcium to avoid significant hypercalciuria. Many studies confirm a small but significant increase in the risk of renal stones with calcium supplements, but not dietary calcium. A recent analysis of published data has suggested that high intakes of calcium from supplements are associated with an increase in the risk of heart disease. This is an evolving story with additional studies that confirm or refute this finding. Because high calcium supplement intakes increase the risk of renal stones and confer no extra benefit to the skeleton, the recommendation that total intakes should be between 1000 and 1200 mg/d is reasonable.

VITAMIN D Vitamin D is synthesized in skin under the influence of heat and ultraviolet light (**Chap. 423**). However, large segments of the population do not obtain sufficient vitamin D to maintain what is now considered an adequate supply [serum 25(OH)D consistently >75 μmol/L (30 ng/mL)]. Because vitamin D supplementation at doses that would achieve these serum levels is safe and inexpensive, the Institute of Medicine (based on obtaining a serum level of 20 ng/mL) recommends daily intakes of 200 IU for adults <50 years of age, 400 IU for those 50–70 years, and 600 IU for those >70 years. Multivitamin tablets usually contain 400 IU, and many

calcium supplements also contain vitamin D. Some data suggest that higher doses (≥1000 IU) may be required in the elderly and chronically ill. The Institute of Medicine report suggests that it is safe to take up to 4000 IU/d. For those with osteoporosis or those at risk of osteoporosis, 1000–2000 IU/d can usually maintain serum 25(OH)D above 30 ng/mL.

OTHER NUTRIENTS Other nutrients such as salt, high animal protein intakes, and caffeine may have modest effects on calcium excretion or absorption. Adequate vitamin K status is required for optimal carboxylation of osteocalcin. States in which vitamin K nutrition or metabolism is impaired, such as with long-term warfarin therapy, have been associated with reduced bone mass. Research concerning cola intake is controversial but suggests a possible link to reduced bone mass through factors that are independent of caffeine. Although dark green leafy vegetables such as spinach and kale contain a fair amount of calcium, the high oxalate content reduces absorption of this calcium (but does not inhibit absorption of calcium from other food eaten simultaneously).

Magnesium is abundant in foods, and magnesium deficiency is quite rare in the absence of a serious chronic disease. Magnesium supplementation may be warranted in patients with inflammatory bowel disease, celiac disease, chemotherapy, severe diarrhea, malnutrition, or alcoholism. Dietary phytoestrogens, which are derived primarily from soy products and legumes (e.g., garbanzo beans [chickpeas] and lentils), exert some estrogenic activity but are insufficiently potent to justify their use in place of a pharmacologic agent in the treatment of osteoporosis.

Patients with hip fractures are often frail and relatively malnourished. Some data suggest an improved outcome in such patients when they are provided calorie and protein supplementation. Excessive protein intake can increase renal calcium excretion, but this can be corrected by an adequate calcium intake.

Exercise Exercise in young individuals increases the likelihood that they will attain the maximal genetically determined peak bone mass. Meta-analyses of studies performed in postmenopausal women indicate that weight-bearing exercise helps prevent bone loss but does not appear to result in substantial gain of bone mass. This beneficial effect wanes if exercise is discontinued. Most of the studies are short term, and a more substantial effect on bone mass is likely if exercise is continued over a long period. Exercise also has beneficial effects on neuromuscular function, and it improves coordination, balance, and strength, thereby reducing the risk of falling. A walking program is a practical way to start. Other activities, such as dancing, racquet sports, cross-country skiing, and use of gym equipment, are also recommended, depending on the patient's personal preference and general condition. Even women who cannot walk benefit from swimming or water exercises, not so much for the effects on bone, which are quite minimal, but because of effects on muscle. Exercise habits should be consistent, optimally at least three times a week.

PHARMACOLOGIC THERAPIES

Before the mid-1990s, estrogen treatment, either by itself or in concert with a progestin, was the primary therapeutic agent for prevention or treatment of osteoporosis. There are now a number of new medications approved for osteoporosis and more under development. Some are agents that specifically treat osteoporosis (bisphosphonates, calcitonin, denosumab, and teriparatide [1-34hPTH]); others, such as selective estrogen response modulators (SERMs) and, most recently, an estrogen/SERM combination medication, have broader effects. The availability of these drugs allows therapy to be tailored to the needs of an individual patient.

Estrogens A large body of clinical trial data indicates that various types of estrogens (conjugated equine estrogens, estradiol, estrone, esterified estrogens, ethinyl estradiol, and mestranol) reduce bone turnover, prevent bone loss, and induce small increases in bone mass of the spine, hip, and total body. The effects of estrogen are

2498 seen in women with natural or surgical menopause and in late postmenopausal women with or without established osteoporosis. Estrogens are efficacious when administered orally or transdermally. For both oral and transdermal routes of administration, combined estrogen/progestin preparations are now available in many countries, obviating the problem of taking two tablets or using a patch and oral progestin.

Dose of Estrogen For oral estrogens, the standard recommended doses have been 0.3 mg/d for esterified estrogens, 0.625 mg/d for conjugated equine estrogens, and 5 µg/d for ethinyl estradiol. For transdermal estrogen, the commonly used dose supplies 50 µg estradiol per day, but a lower dose may be appropriate for some individuals. Dose-response data for conjugated equine estrogens indicate that lower doses (0.3 and 0.45 mg/d) are effective. Doses even lower have been associated with bone mass protection.

FRACTURE DATA Epidemiologic databases indicate that women who take estrogen replacement have a 50% reduction, on average, of osteoporotic fractures, including hip fractures. The beneficial effect of estrogen is greatest among those who start replacement early and continue the treatment; the benefit declines after discontinuation to the extent that there is no residual protective effect against fracture by 10 years after discontinuation. The first clinical trial evaluating fractures as secondary outcomes, the Heart and Estrogen-Progestin Replacement Study (HERS) trial, showed no effect of hormone therapy on hip or other clinical fractures in women with established coronary artery disease. These data made the results of the Women's Health Initiative (WHI) exceedingly important (Chap. 413). The estrogen-progestin arm of the WHI in >16,000 postmenopausal healthy women indicated that hormone therapy reduces the risk of hip and clinical spine fracture by 34% and reduces the risk of all clinical fractures by 24%. There was similar antifracture efficacy seen with estrogen alone in women who had had a hysterectomy.

A few smaller clinical trials have evaluated spine fracture occurrence as an outcome with estrogen therapy. They have consistently shown that estrogen treatment reduces the incidence of vertebral compression fracture.

The WHI has provided a vast amount of data on the multisystemic effects of hormone therapy. Although earlier observational studies suggested that estrogen replacement might reduce heart disease, the WHI showed that combined estrogen-progestin treatment increased risk of fatal and nonfatal myocardial infarction by ~29%, confirming data from the HERS study. Other important relative risks included a 40% increase in stroke, a 100% increase in venous thromboembolic disease, and a 26% increase in risk of breast cancer. Subsequent analyses have confirmed the increased risk of stroke and, in a substudy, showed a twofold increase in dementia. Benefits other than the fracture reductions noted above included a 37% reduction in the risk of colon cancer. These relative risks have to be interpreted in light of absolute risk (Fig. 425-8). For example, out of 10,000 women treated with estrogen-progestin for 1 year, there will be 8 excess heart attacks, 8 excess breast cancers, 18 excess venous thromboembolic events, 5 fewer hip fractures, 44 fewer clinical fractures, and 6 fewer colorectal cancers. These numbers must be multiplied by years of hormone treatment. There was no effect of hormone treatment on the risk of uterine cancer or total mortality.

It is important to note that these WHI findings apply specifically to hormone treatment in the form of conjugated equine estrogen plus medroxyprogesterone acetate. The relative benefits and risks of unopposed estrogen in women who had hysterectomies vary somewhat. They still show benefits against fracture occurrence and increased risk of venous thrombosis and stroke, similar in magnitude to the risks for combined hormone therapy. In contrast, though, the estrogen-only arm of WHI indicated no increased risk of heart attack or breast cancer. The data suggest that at least some of the detrimental effects of combined therapy are related to the progestin component. In addition, there is the possibility, suggested by primate data, that the risk accrues mainly to women who have

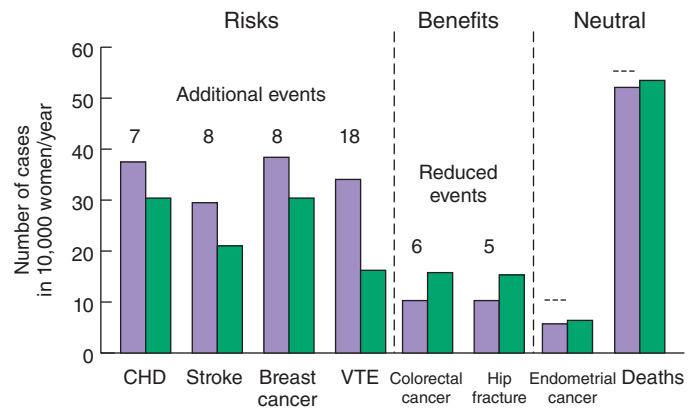


FIGURE 425-8 Effects of hormone therapy on event rates: green, placebo; purple, estrogen and progestin. CHD, coronary heart disease; VTE, venous thromboembolic events. (Adapted from *Women's Health Initiative. WHI HRT Update*. Available at <http://www.nhlbi.nih.gov/health/women/upd2002.htm>.)

some years of estrogen deficiency before initiating treatment. (The average woman in the WHI was more than 10 years from the last menstrual period). Nonetheless, there is reluctance among women to use estrogen/hormone therapy, and the U.S. Preventive Services Task Force has specifically suggested that estrogen/hormone therapy not be used for disease prevention.

MODE OF ACTION Two subtypes of ERs, α and β , have been identified in bone and other tissues. Cells of monocyte lineage express both ER α and ER β , as do osteoblasts. Estrogen-mediated effects vary with the receptor type. Using ER knockout mouse models, elimination of ER α produces a modest reduction in bone mass, whereas mutation of ER β has less of an effect on bone. A male patient with a homozygous mutation of ER α had markedly decreased bone density as well as abnormalities in epiphyseal closure, confirming the important role of ER α in bone biology. The mechanism of estrogen action in bone is an area of active investigation (Fig. 425-5). Although data are conflicting, estrogens may inhibit osteoclasts directly. However, the majority of estrogen (and androgen) effects on bone resorption are mediated through paracrine factors produced by osteoblasts and osteocytes. These actions include decreasing RANKL production and increasing OPG production by osteoblasts.

Progestins In women with a uterus, daily progestin or cyclical progestins at least 12 days per month are prescribed in combination with estrogens to reduce the risk of uterine cancer. Medroxyprogesterone acetate and norethindrone acetate blunt the high-density lipoprotein response to estrogen, but micronized progesterone does not. Neither medroxyprogesterone acetate nor micronized progesterone appears to have an independent effect on bone; at lower doses of estrogen, norethindrone acetate may have an additive benefit. On breast tissue, progestins may increase the risk of breast cancer.

SERMs

Two SERMs are used currently in postmenopausal women: raloxifene, which is approved for the prevention and treatment of osteoporosis as well as the prevention of breast cancer, and tamoxifen, which is approved for the prevention and treatment of breast cancer. A third SERM, bazedoxifene, has been complexed with conjugated estrogen, creating a tissue selective estrogen complex (TSEC). This agent has been approved for prevention of osteoporosis.

Tamoxifen reduces bone turnover and bone loss in postmenopausal women compared with placebo groups. These findings support the concept that tamoxifen acts as an estrogenic agent in bone. There are limited data on the effect of tamoxifen on fracture risk, but the Breast Cancer Prevention Study indicated a possible reduction in clinical vertebral, hip, and Colles' fractures. The major benefit of tamoxifen is on breast cancer occurrence. The breast

cancer prevention trial indicated that tamoxifen administration over 4–5 years reduced the incidence of new invasive and noninvasive breast cancer by ~45% in women at increased risk of breast cancer. The incidence of ER-positive breast cancers was reduced by 65%. Tamoxifen increases the risk of uterine cancer and increases risk of venous thrombosis, cataracts, and possibly stroke in postmenopausal women, limiting its use for breast cancer prevention in women at low or moderate risk.

Raloxifene (60 mg/d) has effects on bone turnover and bone mass that are very similar to those of tamoxifen, indicating that this agent is also estrogenic on the skeleton. The effect of raloxifene on bone density (+1.4–2.8% vs placebo in the spine, hip, and total body) is somewhat less than that seen with standard doses of estrogens. Raloxifene reduces the occurrence of vertebral fracture by 30–50%, depending on the population; however, there are no data confirming that raloxifene can reduce the risk of nonvertebral fractures over 8 years of observation.

Raloxifene, like tamoxifen and estrogen, has effects in other organ systems. The most beneficial effect appears to be a reduction in invasive breast cancer (mainly decreased ER-positive) occurrence of ~65% in women who take raloxifene compared to placebo. In a head-to-head study, raloxifene was as effective as tamoxifen in preventing breast cancer in high-risk women, and raloxifene is now FDA approved for this indication. In a further study, raloxifene had no effect on heart disease in women with increased risk for this outcome. In contrast to tamoxifen, raloxifene is not associated with an increase in the risk of uterine cancer or benign uterine disease. Raloxifene increases the occurrence of hot flashes but reduces serum total and low-density lipoprotein cholesterol, lipoprotein(a), and fibrinogen. Raloxifene, with positive effects on breast cancer and vertebral fractures, has become a useful agent for the treatment of the younger asymptomatic postmenopausal woman. In some women, a recurrence of menopausal hot flashes may occur. Usually this is evanescent, but occasionally, it is sufficiently impactful on daily life and sleep that the drug must be withdrawn. Raloxifene increases the risk of deep vein thrombosis and may increase the risk of death from stroke among older women. Consequently, it is not usually recommended for women over 70 years of age.

The main advantage of the *bazedoxifene/conjugated estrogen* compound is that the bazedoxifene protects uterine tissue from the effects of estrogen and makes it possible to avoid taking a progestin, while using an estrogen primarily for control of menopausal symptoms. The TSEC prevents bone loss somewhat more potently than raloxifene alone and appears safe for the breast.

MODE OF ACTION OF SERMS All SERMs bind to the ER, but each agent produces a unique receptor-drug conformation. As a result, specific co-activator or co-repressor proteins are bound to the receptor (Chap. 400e), resulting in differential effects on gene transcription that vary depending on other transcription factors present in the cell. Another aspect of selectivity is the affinity of each SERM for the different ER α and ER β subtypes, which are expressed differentially in various tissues. These tissue-selective effects of SERMs offer the possibility of tailoring estrogen therapy to best meet the needs and risk factor profile of an individual patient.

Bisphosphonates Alendronate, risedronate, ibandronate, and zoledronic acid are approved for the prevention and treatment of postmenopausal osteoporosis. Alendronate, risedronate, and zoledronic acid are also approved for the treatment of steroid-induced osteoporosis, and risedronate and zoledronic acid are approved for prevention of steroid-induced osteoporosis. Alendronate, risedronate, and zoledronic acid are approved for treatment of osteoporosis in men.

Alendronate has been shown to decrease bone turnover and increase bone mass in the spine by up to 8% versus placebo and by 6% versus placebo in the hip. Multiple trials have evaluated its effect on fracture occurrence. The Fracture Intervention Trial provided evidence in >2000 women with prevalent vertebral fractures that daily alendronate treatment (5 mg/d for 2 years and 10 mg/d for 9 months afterward) reduces vertebral fracture risk by about

50%, multiple vertebral fractures by up to 90%, and hip fractures by up to 50%. Several subsequent trials have confirmed these findings (Fig. 425-9). For example, in a study of >1900 women with low bone mass treated with alendronate (10 mg/d) versus placebo, the incidence of all nonvertebral fractures was reduced by ~47% after only 1 year. In the United States, the 10-mg dose is approved for treatment of osteoporosis and 5 mg/d is used for prevention.

Trials comparing once-weekly alendronate, 70 mg, with daily 10-mg dosing have shown equivalence with regard to bone mass and bone turnover responses. Consequently, once-weekly therapy generally is preferred because of the low incidence of gastrointestinal side effects and ease of administration. Alendronate should be given with a full glass of water before breakfast, because bisphosphonates are poorly absorbed. Because of the potential for esophageal irritation, alendronate is contraindicated in patients who have stricture or inadequate emptying of the esophagus. It is recommended that patients remain upright for at least 30 min after taking the medication to avoid esophageal irritation. Cases of esophagitis, esophageal ulcer, and esophageal stricture have been described, but the incidence appears to be low. In clinical trials, overall gastrointestinal symptomatology was no different with alendronate than with placebo. Alendronate is also available in a preparation that contains vitamin D.

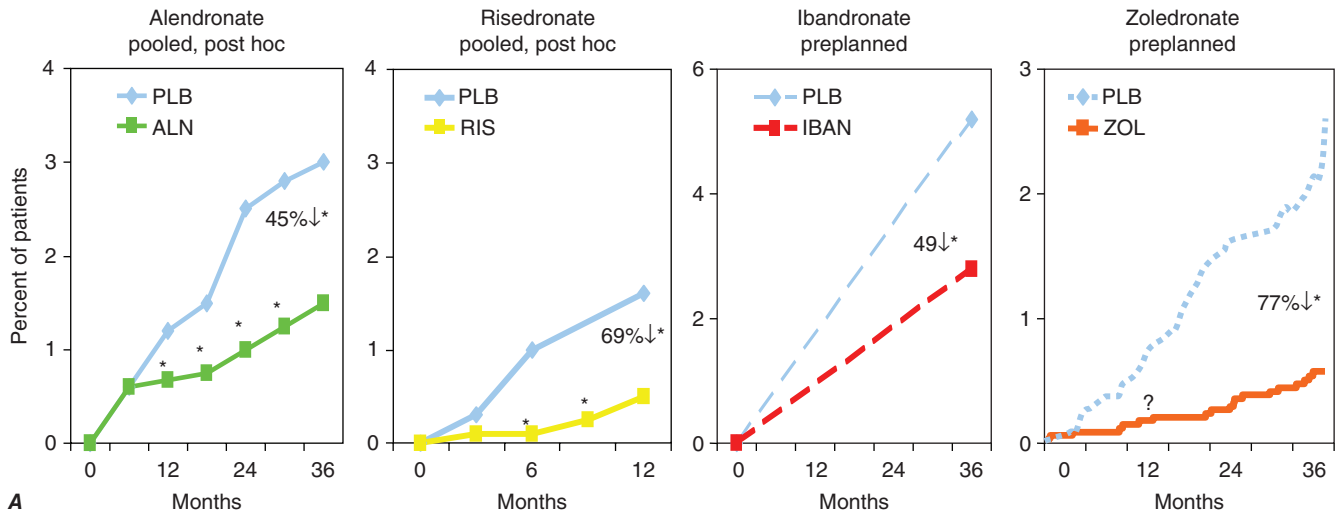
Risedronate also reduces bone turnover and increases bone mass. Controlled clinical trials have demonstrated 40–50% reduction in vertebral fracture risk over 3 years, accompanied by a 40% reduction in clinical nonspine fractures. The only clinical trial specifically designed to evaluate hip fracture outcome (HIP) indicated that risedronate reduced hip fracture risk in women in their seventies with confirmed osteoporosis by 40%. In contrast, risedronate was not effective at reducing hip fracture occurrence in older women (80+ years) without proven osteoporosis. Studies have shown that 35 mg of risedronate administered once weekly is therapeutically equivalent to 5 mg/d and that 150 mg once monthly is therapeutically equivalent to 35 mg once weekly. Patients should take risedronate with a full glass of plain water to facilitate delivery to the stomach and should not lie down for 30 min after taking the drug. The incidence of gastrointestinal side effects in trials with risedronate was similar to that of placebo. A new preparation, which allows risedronate to be taken with food, was recently approved.

Etidronate was the first bisphosphonate to be approved, initially for use in Paget's disease and hypercalcemia. This agent has also been used in osteoporosis trials of smaller magnitude than those performed for alendronate and risedronate but is not approved by the FDA for treatment of osteoporosis. Etidronate probably has some efficacy against vertebral fracture when given as an intermittent cyclical regimen (2 weeks on, 2.5 months off). Its effectiveness against nonvertebral fractures has not been studied.

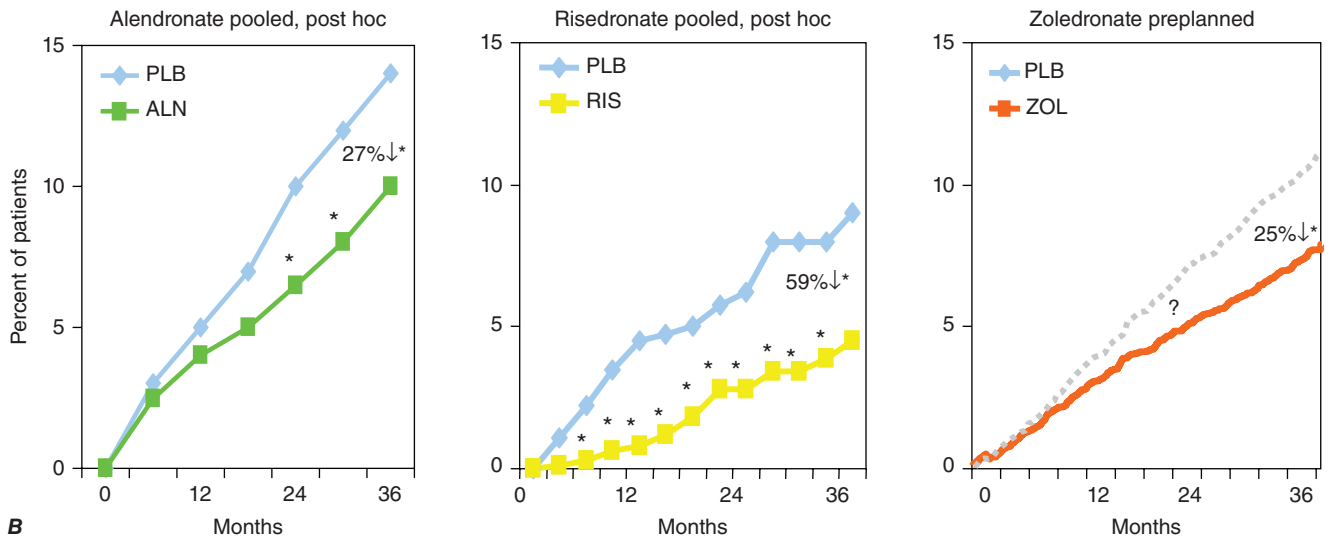
Ibandronate is the third amino-bisphosphonate approved in the United States. Ibandronate (2.5 mg/d) has been shown in clinical trials to reduce vertebral fracture risk by ~40% but with no overall effect on nonvertebral fractures. In a post hoc analysis of subjects with a femoral neck T-score of -3 or below, ibandronate reduced the risk of nonvertebral fractures by ~60%. In clinical trials, ibandronate doses of 150 mg/month PO or 3 mg every 3 months IV had greater effects on turnover and bone mass than did 2.5 mg/d. Patients should take oral ibandronate in the same way as other bisphosphonates, but with 1 h elapsing before other food or drink (other than plain water).

Zoledronic acid is a potent bisphosphonate with a unique administration regimen (5 mg by slow IV infusion annually). The data confirm that it is highly effective in fracture risk reduction. In a study of >7000 women followed for 3 years, zoledronic acid (three annual infusions) reduced the risk of vertebral fractures by 70%, nonvertebral fractures by 25%, and hip fractures by 40%. These results were associated with less height loss and disability. In the treated population, there was an increased risk of transient postdose symptoms (acute-phase reaction) manifested by fever, arthralgia, myalgias, and headache. The symptoms usually last less than 48 h. An increased risk of atrial fibrillation and transient but

Vertebral fractures



Nonvertebral fractures



Hip fractures

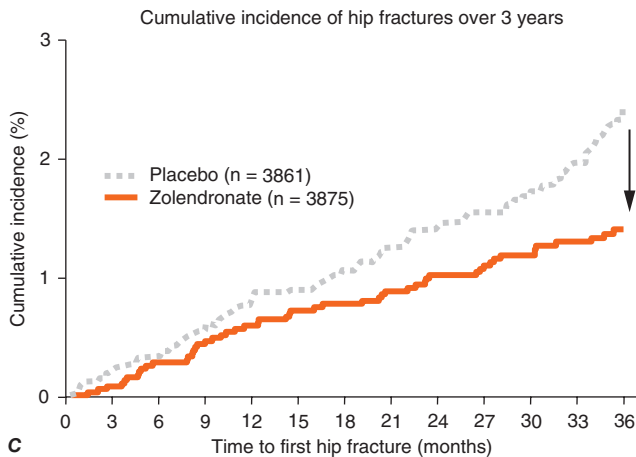


FIGURE 425-9 Effects of various bisphosphonates on clinical vertebral fractures (A), nonvertebral fractures (B), and hip fractures (C). PLB, placebo; RRR, relative risk reduction. (After DM Black et al: *J Clin Endocrinol Metab* 85:4118, 2000; C Roux et al: *Curr Med Res Opin* 4:433, 2004; CH Chesnut et al: *J Bone Miner Res* 19:1241, 2004; DM Black et al: *N Engl J Med* 356:1809, 2007; JT Harrington et al: *Calcif Tissue Int* 74:129, 2003.)

not permanent reduction in renal function was seen in comparison to placebo. Detailed evaluation of all bisphosphonates failed to confirm that these agents increased the risk of atrial fibrillation. Zoledronic acid is the only osteoporosis agent that has been studied in the elderly with a prior hip fracture. The risk of all clinical

fractures was reduced significantly by about 35%, and there was a trend toward reduced risk of a second hip fracture (effect size similar to that seen above). There was also a reduction in mortality of about 30% that was not completely accounted for the reduced hip fracture risk.

Recently there has been concern about two potential side effects associated with bisphosphonate use. The first is osteonecrosis of the jaw (ONJ). ONJ usually follows a dental procedure in which bone is exposed (extractions or dental implants). It is presumed that the exposed bone becomes infected and dies. It is not uncommon among cancer victims with multiple myeloma or patients receiving high doses of bisphosphonates for skeletal metastases, but is rare among persons with osteoporosis on usual doses of bisphosphonates. The second side effect is called atypical femur fracture. These are unusual fractures that occur distal to the lesser trochanter and anywhere along the femoral shaft. They are often preceded by pain in the lateral thigh or groin that can be present for weeks or months before the fracture. The fractures occur with trivial trauma, sometimes completely spontaneously, and are primarily transverse, with a medial break when complete and minimally comminuted. A localized periosteal reaction, consistent with a stress fracture, is often seen in the lateral cortex (Fig. 425-10). The overall risk is low (suggested to be about one-one hundredth to one-tenth that of hip fracture) but appears to increase in incidence with long-term use of bisphosphonates. Although the fractures may be bisphosphonate related in many individuals, they clearly occur in patients with no prior bisphosphonate exposure. When complete, they require surgical fixation and may be difficult to heal. Anabolic medication may accelerate healing of these fractures in some patients, and surgery can sometimes be avoided. Patients initiating bisphosphonates need to be warned that if they develop thigh or groin pain they must notify their physician. Routine x-rays will sometimes pick up cortical thickening or even a stress fracture, but more commonly MRI or technetium bone scan is required. The presence of an abnormality requires at minimum a period of modified weight bearing and may need prophylactic rodding of the femur. It is important to realize that these fractures may be bilateral, and when an abnormality is found, the other femur should be investigated.

MODE OF ACTION Bisphosphonates are structurally related to pyrophosphates, compounds that are incorporated into bone matrix. Bisphosphonates specifically impair osteoclast function and reduce osteoclast number, in part by inducing apoptosis. Recent evidence suggests that the nitrogen-containing bisphosphonates also inhibit protein prenylation, one of the end products in the mevalonic

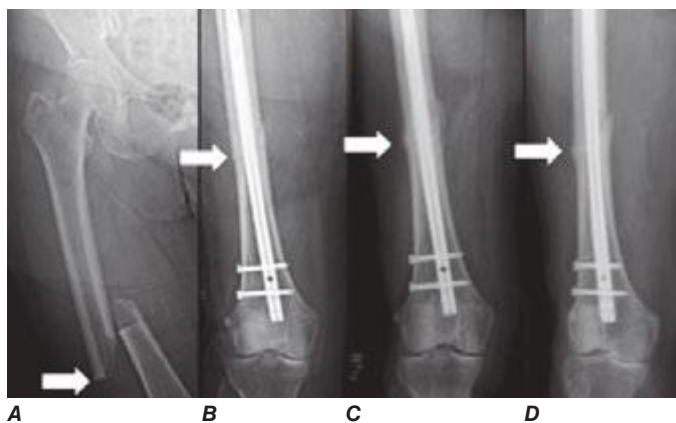


FIGURE 425-10 An atypical femur fracture (AFF) of the femoral diaphysis. **A.** Note the transverse fracture line in the lateral cortex that becomes oblique as it progresses medially across the femur (white arrow). **B.** On radiograph obtained immediately after intramedullary rod placement, a small area of periosteal thickening of the lateral cortex is visible (white arrow). **C.** On radiograph obtained at 6 weeks, note callus formation of the fracture site (white arrow). **D.** On radiograph obtained at 3 months, there is a mature callus that has failed to bridge the cortical gap (white arrow). Note the localized periosteal and/or endosteal thickening of the lateral cortex at the fracture site (white arrow). (From E Shane et al: *J Bone Min Res* 29:1-23, 2014. Courtesy of Fergus McKiernan.)

acid pathway, by inhibiting the enzyme farnesyl pyrophosphate synthase. This effect disrupts intracellular protein trafficking and ultimately may lead to apoptosis. Some bisphosphonates have very long retention in the skeleton and may exert long-term effects. The consequences of this, if any, are unknown.

Calcitonin Calcitonin is a polypeptide hormone produced by the thyroid gland (Chap. 424). Its physiologic role is unclear because no skeletal disease has been described in association with calcitonin deficiency or excess. Calcitonin preparations are approved by the FDA for Paget's disease, hypercalcemia, and osteoporosis in women >5 years past menopause. Concerns have been raised about an increase in the incidence of cancer associated with calcitonin use. Initially, the cancer noted was of the prostate, but an analysis of all data suggested a more general increase in cancer risk. In Europe, the European Medicines Agency (EMA) has removed the osteoporosis indication, and an FDA Advisory Committee has voted for a similar change in the United States.

Injectable calcitonin produces small increments in bone mass of the lumbar spine. However, difficulty of administration and frequent reactions, including nausea and facial flushing, make general use limited. A nasal spray containing calcitonin (200 IU/d) is available for treatment of osteoporosis in postmenopausal women. One study suggests that nasal calcitonin produces small increments in bone mass and a small reduction in new vertebral fractures in calcitonin-treated patients versus those on calcium alone. There has been no proven effectiveness against nonvertebral fractures.

Calcitonin is not indicated for prevention of osteoporosis and is not sufficiently potent to prevent bone loss in early postmenopausal women. Calcitonin might have an analgesic effect on bone pain, both in the subcutaneous and possibly the nasal form.

MODE OF ACTION Calcitonin suppresses osteoclast activity by direct action on the osteoclast calcitonin receptor. Osteoclasts exposed to calcitonin cannot maintain their active ruffled border, which normally maintains close contact with underlying bone.

Denosumab A novel agent that was given twice yearly by SC administration in a randomized controlled trial in postmenopausal women with osteoporosis has been shown to increase BMD in the spine, hip, and forearm and reduce vertebral, hip, and nonvertebral fractures over a 3-year period by 70, 40, and 20%, respectively (Fig. 425-11). Other clinical trials indicate ability to increase bone mass in postmenopausal women with low bone mass (above osteoporosis range) and in postmenopausal women with breast cancer treated with hormonal agents. Furthermore, a study of men with prostate cancer treated with gonadotropin-releasing hormone (GnRH) agonist therapy indicated the ability of denosumab to improve bone mass and reduce vertebral fracture occurrence. Denosumab was approved by the FDA in 2010 for the treatment of postmenopausal women who have a high risk for osteoporotic fractures, including those with a history of fracture or multiple risk factors for fracture, and those who have failed or are intolerant to other osteoporosis therapy. Denosumab is also approved for the treatment of osteoporosis in men at high risk, men with prostate cancer on GnRH agonist therapy, and women with breast cancer on aromatase inhibitor therapy.

MODE OF ACTION Denosumab is a fully human monoclonal antibody to RANKL, the final common effector of osteoclast formation, activity, and survival. Denosumab binds to RANKL, inhibiting its ability to initiate formation of mature osteoclasts from osteoclast precursors and to bring mature osteoclasts to the bone surface and initiate bone resorption. Denosumab also plays a role in reducing the survival of the osteoclast. Through these actions on the osteoclast, denosumab induces potent antiresorptive action, as assessed biochemically and histomorphometrically, and may contribute to the occurrence of ONJ. Atypical femur fractures have also been noted. Serious adverse reactions include hypocalcemia, skin infections (usually cellulitis of the lower extremity), and dermatologic reactions such as dermatitis, rashes, and eczema. The effects of denosumab are rapidly reversible.

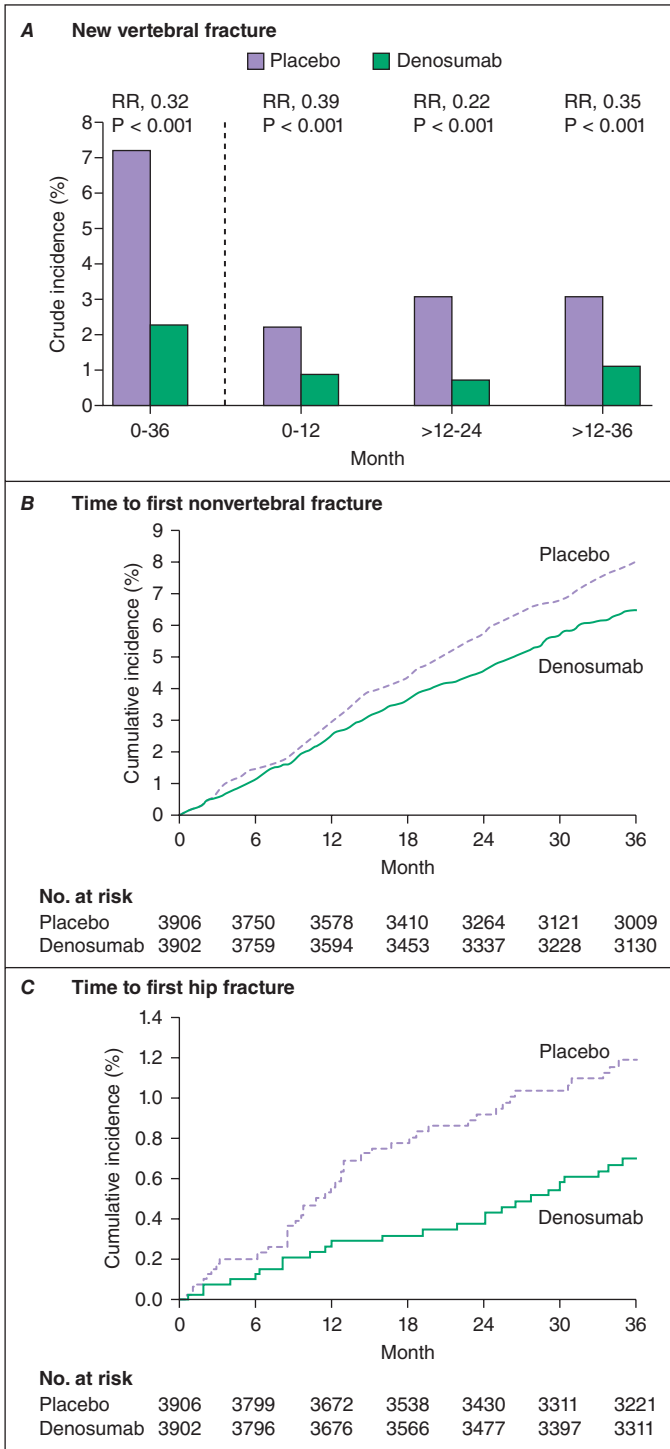


FIGURE 425-11 Effects of denosumab on new vertebral fractures (A) and times to nonvertebral and hip fracture (B and C). RR, relative risk. (After SR Cummings et al: *N Engl J Med* 361:756, 2009.)

If denosumab is stopped, bone will be lost rapidly if another agent is not used.

Parathyroid Hormone Endogenous PTH is an 84-amino-acid peptide that is largely responsible for calcium homeostasis (Chap. 424). Although chronic elevation of PTH, as occurs in hyperparathyroidism, is associated with bone loss (particularly cortical bone), PTH when given exogenously as a daily injection exerts anabolic effects on bone. Teriparatide (1-34hPTH) is approved for the treatment of osteoporosis in both men and women at high risk for fracture. In a pivotal study (median time of treatment, 19 months' duration), 20 µg of teriparatide daily by SC injection reduced vertebral fractures by 65% and nonvertebral fractures by 45% (Fig. 425-12). Treatment is administered as

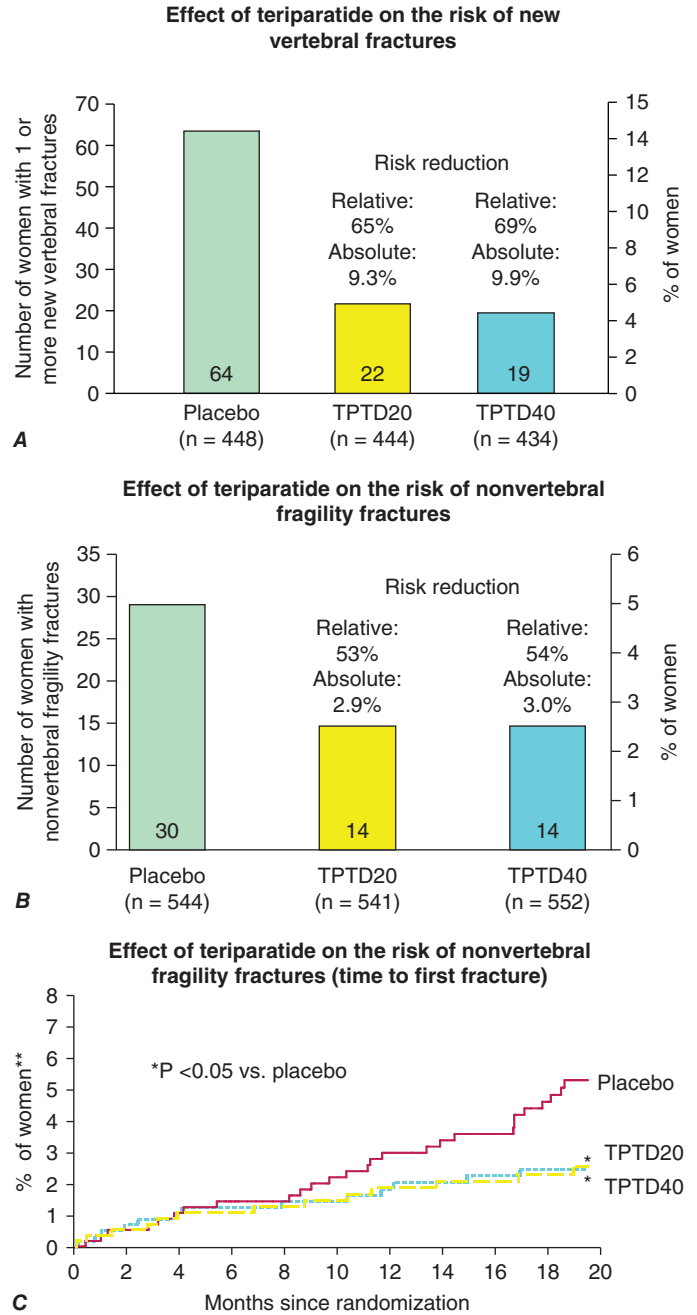


FIGURE 425-12 Effects of teriparatide (TPTD) on new vertebral fractures (A) and nonvertebral fragility fractures (B and C). (After RM Neer et al: *N Engl J Med* 344:1434, 2001.)

a single daily injection given for a maximum of 2 years. Teriparatide produces increases in bone mass and mediates architectural improvements in skeletal structure. These effects are lower when patients have been exposed previously to bisphosphonates, possibly in proportion to the potency of the antiresorptive effect. When teriparatide is being considered for treatment-naïve patients, it is best administered as monotherapy and followed by an antiresorptive agent such as a bisphosphonate. If teriparatide treatment is not followed by an antiresorptive agent, the bone gained is rapidly lost.

Side effects of teriparatide are generally mild and can include leg cramps, muscle pain, weakness, dizziness, headache, and nausea. Rodents given prolonged treatment with PTH in relatively high doses developed osteogenic sarcomas. Long-term surveillance studies suggest no association between 2 years of teriparatide administration and osteosarcoma risk in humans.

PTH use may be limited by its mode of administration; alternative modes of delivery are being investigated. The optimal frequency of administration also remains to be established, and it is possible that

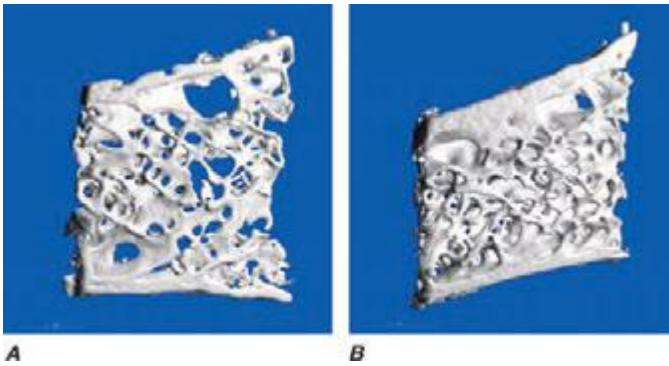


FIGURE 425-13 Effect of parathyroid hormone (PTH) treatment on bone microarchitecture. Paired biopsy specimens from a 64-year-old woman before (A) and after (B) treatment with PTH. (From DW Dempster et al: *J Bone Miner Res* 16:1846, 2001.)

PTH might be effective when used intermittently. Cost also may be a limiting factor. In some settings, the effect of PTH might be enhanced by combination with an antiresorptive agent. This might be particularly important in patients who have been treated previously with bisphosphonate medications.

MODE OF ACTION Exogenously administered PTH appears to have direct actions on osteoblast activity, with biochemical and histomorphometric evidence of *de novo* bone formation early in response to PTH, before activation of bone resorption. Subsequently, PTH activates bone remodeling but still appears to favor bone formation over bone resorption. PTH stimulates Wnt signaling, IGF-I, and collagen production and appears to increase osteoblast number by stimulating replication, enhancing osteoblast recruitment, and inhibiting apoptosis. Unlike all other treatments, PTH produces a true increase in bone tissue and an apparent restoration of bone microarchitecture (Fig. 425-13).

Fluoride Fluoride has been available for many years and is a potent stimulator of osteoprogenitor cells when studied *in vitro*. It has been used in multiple osteoporosis studies with conflicting results, in part because of the use of varying doses and preparations. Despite increments in bone mass of up to 10%, there are no consistent effects of fluoride on vertebral or nonvertebral fracture; the latter may actually increase when high doses of fluoride are used. Fluoride remains an experimental agent despite its long history and multiple studies.

Strontium Ranelate Strontium ranelate is approved in several European countries for the treatment of osteoporosis. It increases bone mass throughout the skeleton; in clinical trials, the drug reduced the risk of vertebral fractures by 37% and that of nonvertebral fractures by 14%. It appears to be modestly antiresorptive while at the same time not causing as much of a decrease in bone formation (measured biochemically). Strontium is incorporated into hydroxyapatite, replacing calcium, a feature that might explain some of its fracture benefits. Small increased risks of venous thrombosis, sometimes severe dermatologic reactions, seizures, and abnormal cognition have been seen and require further study. An increase in risk of cardiovascular disease has also been associated with use of strontium, such that the EMA has restricted its use at present.

Other Potential Anabolic Agents Several small studies of growth hormone (GH), alone or in combination with other agents, have not shown consistent or substantial positive effects on skeletal mass. Many of these studies have been relatively short term, and the effects of GH, growth hormone-releasing hormone, and the IGFs are still under investigation. Anabolic steroids, mostly derivatives of testosterone, act primarily as antiresorptive agents to reduce bone turnover but also may stimulate osteoblastic activity. Effects on bone mass remain unclear but appear weak in general, and use is limited by masculinizing side effects. Several observational studies

suggested that the statin drugs, used to treat hypercholesterolemia, may be associated with increased bone mass and reduced fractures, but conclusions from clinical trials have been largely negative. Early studies with sclerostin antibodies, which inhibit sclerostin, activate Wnt, and might be highly anabolic to bone, are under development. Odanacatib is a mixed antiresorptive, partial bone formation stimulator that is currently in the late stages of development.

NONPHARMACOLOGIC APPROACHES

In some early studies, protective pads worn around the outer thigh, which cover the trochanteric region of the hip, were able to prevent hip fractures in elderly residents in nursing homes. Randomized controlled trials of hip protectors have been unable to confirm these early findings. Therefore, the efficacy of hip protectors remains controversial at this time.

Kyphoplasty and *vertebroplasty* are also useful nonpharmacologic approaches for the treatment of painful vertebral fractures. However, no long-term data are available.

TREATMENT MONITORING

There are currently no well-accepted guidelines for monitoring treatment of osteoporosis. Because most osteoporosis treatments produce small or moderate bone mass increments on average, it is reasonable to consider BMD as a monitoring tool. Changes must exceed ~4% in the spine and 6% in the hip to be considered significant in any individual. The hip is the preferred site due to larger surface area and greater reproducibility. Medication-induced increments may require several years to produce changes of this magnitude (if they do at all). Consequently, it can be argued that BMD should be repeated at intervals >2 years. Only significant BMD reductions should prompt a change in medical regimen, because it is expected that many individuals will not show responses greater than the detection limits of the current measurement techniques.

Biochemical markers of bone turnover may prove useful for treatment monitoring, but little hard evidence currently supports this concept; it remains unclear which endpoint is most useful. If bone turnover markers are used, a determination should be made before therapy is started and repeated ≥ 4 months after therapy is initiated. In general, a change in bone turnover markers must be 30–40% lower than the baseline to be significant because of the biologic and technical variability in these tests. A positive change in biochemical markers and/or bone density can be useful to help patients adhere to treatment regimens.

GLUCOCORTICOID-INDUCED OSTEOPOROSIS

Osteoporotic fractures are a well-characterized consequence of the hypercortisolism associated with Cushing's syndrome. However, the therapeutic use of glucocorticoids is by far the most common form of glucocorticoid-induced osteoporosis. Glucocorticoids are used widely in the treatment of a variety of disorders, including chronic lung disorders, rheumatoid arthritis and other connective tissue diseases, inflammatory bowel disease, and after transplantation. Osteoporosis and related fractures are serious side effects of chronic glucocorticoid therapy. Because the effects of glucocorticoids on the skeleton are often superimposed on the consequences of aging and menopause, it is not surprising that women and the elderly are most frequently affected. The skeletal response to steroids is remarkably heterogeneous, however, and even young, growing individuals treated with glucocorticoids can present with fractures.

The risk of fractures depends on the dose and duration of glucocorticoid therapy, although recent data suggest that there may be no completely safe dose. Bone loss is more rapid during the early months of treatment, and trabecular bone is affected more severely than cortical bone. As a result, fractures have been shown to increase within 3 months of steroid treatment. There is an increase in fracture risk in both the axial skeleton and the appendicular skeleton, including risk of hip fracture. Bone loss can occur with any route of steroid administration, including high-dose inhaled glucocorticoids and intraarticular

2504 injections. Alternate-day delivery does not appear to ameliorate the skeletal effects of glucocorticoids.

PATHOPHYSIOLOGY

Glucocorticoids increase bone loss by multiple mechanisms, including (1) inhibition of osteoblast function and an increase in osteoblast apoptosis, resulting in impaired synthesis of new bone; (2) stimulation of bone resorption, probably as a secondary effect; (3) impairment of the absorption of calcium across the intestine, probably by a vitamin D-independent effect; (4) increase of urinary calcium loss and perhaps induction of some degree of secondary hyperparathyroidism; (5) reduction of adrenal androgens and suppression of ovarian and testicular secretion of estrogens and androgens; and (6) induction of glucocorticoid myopathy, which may exacerbate effects on skeletal and calcium homeostasis as well as increase the risk of falls.

EVALUATION OF THE PATIENT

Because of the prevalence of glucocorticoid-induced bone loss, it is important to evaluate the status of the skeleton in all patients starting or already receiving long-term glucocorticoid therapy. Modifiable risk factors should be identified, including those for falls. Examination should include testing of height and muscle strength. Laboratory evaluation should include an assessment of 24-h urinary calcium. All patients on long-term (>3 months) glucocorticoids should have measurement of bone mass at both the spine and the hip using DXA. If only one skeletal site can be measured, it is best to assess the spine in individuals <60 years and the hip in those >60 years.

PREVENTION

Bone loss caused by glucocorticoids can be prevented and the risk of fractures significantly reduced. Strategies must include using the lowest dose of glucocorticoid for disease management. Topical and inhaled routes of administration are preferred, where appropriate. Risk factor reduction is important, including smoking cessation, limitation of alcohol consumption, and participation in weight-bearing exercise, when appropriate. All patients should receive an adequate calcium and vitamin D intake from the diet or from supplements.

TREATMENT **GLUCOCORTICOID-INDUCED OSTEOPOROSIS**

Several bisphosphonates (alendronate, risedronate, and zoledronic acid) have been demonstrated in large clinical trials to reduce the risk of vertebral fractures in patients being treated with glucocorticoids, as well as improve bone mass in spine and hip. Teriparatide also improves bone mass and reduces fracture risk in glucocorticoid-treated osteoporosis compared to an active comparator (alendronate).

426e Paget's Disease and Other Dysplasias of Bone

Murray J. Favus, Tamara J. Vokes

PAGET'S DISEASE OF BONE

Paget's disease is a localized bone-remodeling disorder that affects widespread, noncontiguous areas of the skeleton. The pathologic process is initiated by overactive osteoclastic bone resorption followed by a compensatory increase in osteoblastic new bone formation, resulting in a structurally disorganized mosaic of woven and lamellar bone. Pagetic bone is expanded, less compact, and more vascular; thus, it is more susceptible to deformities and fractures. Although most patients are asymptomatic, symptoms resulting directly from bony involvement (bone pain, secondary arthritis, fractures) or secondarily from the expansion of bone causing compression of surrounding neural tissue are not uncommon.

Epidemiology There is a marked geographic variation in the frequency of Paget's disease, with high prevalence in Western Europe (Great Britain, France, and Germany, but not Switzerland or Scandinavia) and among those who have immigrated to Australia, New Zealand, South Africa, and North and South America. The disease is rare in native populations of the Americas, Africa, Asia, and the Middle East; when it does occur, the affected subjects usually have evidence of European ancestry, supporting the migration theory. For unclear reasons, the prevalence and severity of Paget's disease are decreasing, and the age of diagnosis is increasing.

The prevalence is greater in males and increases with age. Autopsy series reveal Paget's disease in about 3% of those over age 40. Prevalence of positive skeletal radiographs in patients over age 55 is 2.5% for men and 1.6% for women. Elevated alkaline phosphatase (ALP) levels in asymptomatic patients have an age-adjusted incidence of 12.7 and 7 per 100,000 person-years in men and women, respectively.

Etiology The etiology of Paget's disease of bone remains unknown, but evidence supports both genetic and viral etiologies. A positive family history is found in 15–25% of patients and, when present, raises the prevalence of the disease seven- to tenfold among first-degree relatives.

A clear genetic basis has been established for several rare familial bone disorders that clinically and radiographically resemble Paget's disease but have more severe presentation and earlier onset. A homozygous deletion of the *TNFRSF11B* gene, which encodes osteoprotegerin (Fig. 426e-1), causes *juvenile Paget's disease*, also known as *familial idiopathic hyperphosphatasia*, a disorder characterized by uncontrolled osteoclastic differentiation and resorption. Familial patterns of disease in several large kindred are consistent with an autosomal dominant pattern of inheritance with variable penetrance. *Familial expansile osteolysis*, *expansile skeletal hyperphosphatasia*, and *early-onset Paget's disease* are associated with mutations in *TNFRSF11A* gene, which encodes RANK (receptor activator of nuclear factor- κ B), a member of the tumor necrosis factor superfamily critical for osteoclast differentiation (Fig. 426e-1). Finally, mutations in the gene for valosin-containing protein cause a rare syndrome with autosomal dominant inheritance and variable penetrance known as *inclusion body myopathy with Paget's disease and frontotemporal dementia (IBMPFD)*. The role of genetic factors is less clear in the more common form of late-onset Paget's disease. Although a few families with mutations in the gene encoding RANK have been reported, the most common mutations identified in familial and sporadic cases of Paget's disease have been in the *SQSTM1* gene (sequestasome-1 or p62 protein) in the C-terminal ubiquitin-binding domain. The p62 protein is involved in nuclear factor κ B (NF- κ B) signaling and regulates osteoclastic differentiation. The phenotypic variability in patients with *SQSTM1* mutations suggests that additional factors, such as other genetic influences or viral infection, may influence clinical expression of the disease.

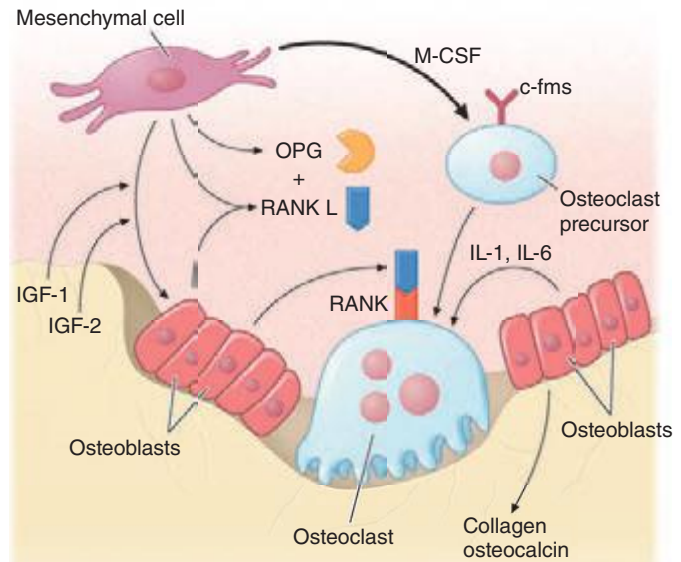


FIGURE 426e-1 Diagram illustrating factors that promote differentiation and function of osteoclasts and osteoblasts and the role of the RANK pathway. Stromal bone marrow (mesenchymal) cells and differentiated osteoblasts produce multiple growth factors and cytokines, including macrophage colony-stimulating factor (M-CSF), to modulate osteoclastogenesis. RANKL (receptor activator of nuclear factor- κ B ligand) is produced by osteoblast progenitors and mature osteoblasts and can bind to a soluble decoy receptor known as OPG (osteoprotegerin) to inhibit RANKL action. Alternatively, a cell-cell interaction between osteoblast and osteoclast progenitors allows RANKL to bind to its membrane-bound receptor, RANK, thereby stimulating osteoclast differentiation and function. RANK binds intracellular proteins called TRAFs (tumor necrosis factor receptor-associated factors) that mediate receptor signaling through transcription factors such as NF- κ B. M-CSF binds to its receptor, c-fms, which is the cellular homologue of the *fms* oncogene. See text for the potential role of these pathways in disorders of osteoclast function such as Paget's disease and osteopetrosis. IL, interleukin; IGF, insulin-like growth factor.

Several lines of evidence suggest that a viral infection may contribute to the clinical manifestations of Paget's disease, including (1) the presence of cytoplasmic and nuclear inclusions resembling paramyxoviruses (measles and respiratory syncytial virus) in pagetic osteoclasts and (2) viral mRNA in precursor and mature osteoclasts. The viral etiology is further supported by conversion of osteoclast precursors to pagetic-like osteoclasts by vectors containing the measles virus nucleocapsid or matrix genes. However, the viral etiology has been questioned by the inability to culture a live virus from pagetic bone and by failure to clone the full-length viral genes from material obtained from patients with Paget's disease.

Pathophysiology The principal abnormality in Paget's disease is the increased number and activity of osteoclasts. Pagetic osteoclasts are large, increased 10- to 100-fold in number, and have a greater number of nuclei (as many as 100 compared to 3–5 nuclei in the normal osteoclast). The overactive osteoclasts may create a sevenfold increase in resorptive surfaces and an erosion rate of 9 μ g/d (normal is 1 μ g/d). Several causes for the increased number and activity of pagetic osteoclasts have been identified: (1) osteoclastic precursors are hypersensitive to $1,25(\text{OH})_2\text{D}_3$; (2) osteoclasts are hyperresponsive to RANK ligand (RANKL), the osteoclast stimulatory factor that mediates the effects of most osteotropic factors on osteoclast formation; (3) marrow stromal cells from pagetic lesions have increased RANKL expression; (4) osteoclast precursor recruitment is increased by interleukin (IL) 6, which is increased in the blood of patients with active Paget's disease and is overexpressed in pagetic osteoclasts; (5) expression of the proto-oncogene *c-fos*, which increases osteoclastic activity, is increased; and

426e-2 (6) the antiapoptotic oncogene *Bcl-2* in pagetic bone is overexpressed. Numerous osteoblasts are recruited to active resorption sites and produce large amounts of new bone matrix. As a result, bone turnover is high, and bone mass is normal or increased, not reduced, unless there is concomitant deficiency of calcium and/or vitamin D.

The characteristic feature of Paget's disease is increased bone resorption accompanied by accelerated bone formation. An initial osteolytic phase involves prominent bone resorption and marked hypervascularization. Radiographically, this manifests as an advancing lytic wedge, or "blade of grass" lesion. The second phase is a period of very active bone formation and resorption that replaces normal lamellar bone with haphazard (woven) bone. Fibrous connective tissue may replace normal bone marrow. In the final sclerotic phase, bone resorption declines progressively and leads to a hard, dense, less vascular pagetic or mosaic bone, which represents the so-called burned-out phase of Paget's disease. All three phases may be present at the same time at different skeletal sites.

Clinical Manifestations Diagnosis is often made in asymptomatic patients because they have elevated ALP levels on routine blood chemistry testing or an abnormality on a skeletal radiograph obtained for another indication. The skeletal sites most commonly involved are the pelvis, vertebral bodies, skull, femur, and tibia. Familial cases with an early presentation often have numerous active sites of skeletal involvement.

The most common presenting symptom is pain, which may result from increased bony vascularity, expanding lytic lesions, fractures, bowing, or other deformities. Bowing of the femur or tibia causes gait abnormalities and abnormal mechanical stresses with secondary osteoarthritis of the hip or knee joints. Long bone bowing also causes extremity pain by stretching the muscles attached to the bone softened by the pagetic process. Back pain results from enlarged pagetic vertebrae, vertebral compression fractures, spinal stenosis, degenerative changes of the joints, and altered body mechanics with kyphosis and forward tilt of the upper back. Rarely, spinal cord compression may result from bone enlargement or from the vascular steal syndrome. Skull involvement may cause headaches, symmetric or asymmetric enlargement of the parietal or frontal bones (frontal bossing), and increased head size. Cranial expansion may narrow cranial foramina and cause neurologic complications including hearing loss from cochlear nerve damage from temporal bone involvement, cranial nerve palsies, and softening of the base of the skull (*platybasia*) with the risk of brainstem compression. Pagetic involvement of the facial bones may cause facial deformity; loss of teeth and other dental conditions; and, rarely, airway compression.

Fractures are serious complications of Paget's disease and usually occur in long bones at areas of active or advancing lytic lesions.

Common fracture sites are the femoral shaft and subtrochanteric regions. Neoplasms arising from pagetic bone are rare (<0.5%). The incidence of sarcoma appears to be decreasing, possibly because of earlier, more effective treatment with potent antiresorptive agents. The majority of tumors are osteosarcomas, which usually present with new pain in a long-standing pagetic lesion. Osteoclast-rich benign giant cell tumors may arise in areas adjacent to pagetic bone, and they respond to glucocorticoid therapy.

Cardiovascular complications may occur in patients with involvement of large (15–35%) portions of the skeleton and a high degree of disease activity (ALP four times above normal). The extensive arteriovenous shunting and marked increases in blood flow through the vascular pagetic bone lead to a high-output state and cardiac enlargement. However, high-output heart failure is relatively rare and usually develops in patients with concomitant cardiac pathology. In addition, calcific aortic stenosis and diffuse vascular calcifications have been associated with Paget's disease.

Diagnosis The diagnosis may be suggested on clinical examination by the presence of an enlarged skull with frontal bossing, bowing of an extremity, or short stature with simian posturing. An extremity with an area of warmth and tenderness to palpation may suggest an underlying pagetic lesion. Other findings include bony deformity of the pelvis, skull, spine, and extremities; arthritic involvement of the joints adjacent to lesions; and leg-length discrepancy resulting from deformities of the long bones.

Paget's disease is usually diagnosed from radiologic and biochemical abnormalities. Radiographic findings typical of Paget's disease include enlargement or expansion of an entire bone or area of a long bone, cortical thickening, coarsening of trabecular markings, and typical lytic and sclerotic changes. Skull radiographs (Fig. 426e-2) reveal regions of "cotton wool," or osteoporosis circumscripta, thickening of diploic areas, and enlargement and sclerosis of a portion or all of one or more skull bones. Vertebral cortical thickening of the superior and inferior end plates creates a "picture frame" vertebra. Diffuse radiodense enlargement of a vertebra is referred to as "ivory vertebra." Pelvic radiographs may demonstrate disruption or fusion of the sacroiliac joints; porotic and radiodense lesions of the ilium with whorls of coarse trabeculation; thickened and sclerotic iliopectineal line (brim sign); and softening with protrusio acetabuli, with axial migration of the hips and functional flexion contracture. Radiographs of long bones reveal bowing deformity and typical pagetic changes of cortical thickening and expansion and areas of lucency and sclerosis (Fig. 426e-3). Radionuclide ^{99m}Tc bone scans are less specific but are more sensitive than standard radiographs for identifying sites of active skeletal lesions. Although computed tomography (CT) and magnetic resonance imaging (MRI) studies are not necessary in most cases,

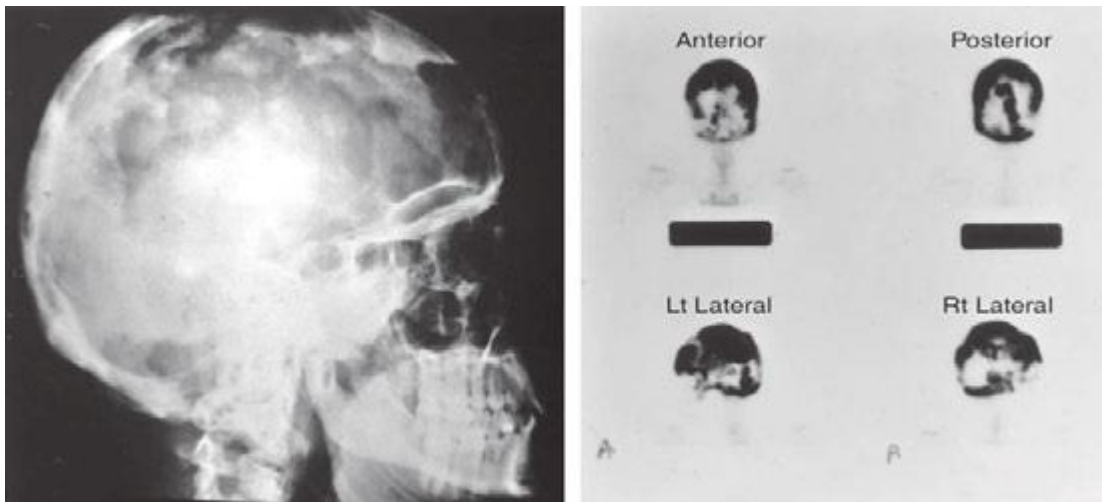


FIGURE 426e-2 A 48-year-old woman with Paget's disease of the skull. **Left.** Lateral radiograph showing areas of both bone resorption and sclerosis. **Right.** ^{99m}Tc HDP bone scan with anterior, posterior, and lateral views of the skull showing diffuse isotope uptake by the frontal, parietal, occipital, and petrous bones.



FIGURE 426e-3 Radiograph of a 73-year-old man with Paget's disease of the right proximal femur. Note the coarsening of the trabecular pattern with marked cortical thickening and narrowing of the joint space consistent with osteoarthritis secondary to pagetic deformity of the right femur.

CT may be useful for the assessment of possible fracture, and MRI is necessary to assess the possibility of sarcoma, giant cell tumor, or metastatic disease in pagetic bone. Definitive diagnosis of malignancy often requires bone biopsy.

Biochemical evaluation is useful in the diagnosis and management of Paget's disease. The marked increase in bone turnover can be monitored using biochemical markers of bone formation and resorption. The parallel rise in markers of bone formation and resorption confirms the coupling of bone formation and resorption in Paget's disease. The degree of bone marker elevation reflects the extent and severity of the disease. Patients with the highest elevation of ALP (10 × the upper limit of normal) typically have involvement of the skull and at least one other skeletal site. Lower values suggest less extensive involvement or a quiescent phase of the disease. For most patients, serum total ALP remains the test of choice both for diagnosis and assessing response to therapy. Occasionally, a symptomatic patient with evidence of progression at a single site may have a normal total ALP level but increased bone-specific ALP. For unclear reasons, serum osteocalcin, another marker of bone formation, is not always elevated and is not recommended for use in diagnosis or management of Paget's disease. Bone resorption markers (serum or urine N-telopeptide or C-telopeptide measured in the blood or urine) are also elevated in active Paget's disease and decrease more rapidly in response to therapy than does ALP.

Serum calcium and phosphate levels are normal in Paget's disease. Immobilization of a patient with active Paget's disease may rarely cause hypercalcemia and hypercalciuria and increase the risk for nephrolithiasis. However, the discovery of hypercalcemia, even in the presence of immobilization, should prompt a search for another cause of hypercalcemia. In contrast, hypocalcemia or mild secondary hyperparathyroidism may develop in Paget's patients with very active bone formation and insufficient calcium and vitamin D intake, particularly during bisphosphonate therapy when bone resorption is rapidly suppressed and active bone formation continues. Therefore, adequate calcium and vitamin D intake should be instituted prior to administration of bisphosphonates.

TREATMENT PAGET'S DISEASE OF BONE

The development of effective and potent pharmacologic agents (Table 426e-1) has changed the treatment philosophy from treating only symptomatic patients to treating asymptomatic patients who are at risk for complications. Pharmacologic therapy is indicated in

TABLE 426e-1 PHARMACOLOGIC AGENTS APPROVED FOR TREATMENT OF PAGET'S DISEASE

Name	Dose and Mode of Delivery	Normalization of ALP
Zoledronic acid	5 mg IV over 15 min	90% of patients at 6 mo
Pamidronate	30 mg IV/d over 4 h on 3 days	~50% of patients
Risedronate	30 mg PO/d for 2 mo	73% of patients
Alendronate	40 mg PO/d for 6 mo	63% of patients
Tiludronate	800 mg PO daily for 3 mo	35% of patients
Etidronate	200–400 mg PO/d × 6 mo	15% of patients
Calcitonin (Miacalcin)	100 U SC daily for 6–18 mo (may reduce to 50 U 3 × per wk)	(Reduction of ALP by up to 50%)

the following circumstances: to control symptoms caused by metabolically active Paget's disease such as bone pain, fracture, headache, pain from pagetic radiculopathy or arthropathy, or neurologic complications; to decrease local blood flow and minimize operative blood loss in patients who need surgery at an active pagetic site; to reduce hypercalciuria that may occur during immobilization; and to decrease the risk of complications when disease activity is high (elevated ALP) and when the site of involvement involves weight-bearing bones, areas adjacent to major joints, vertebral bodies, and the skull. Whether or not early therapy prevents late complications remains to be determined. A randomized study of over 1200 patients from the United Kingdom showed no difference in bone pain, fracture rates, quality of life, and hearing loss between patients who received pharmacologic therapy to control symptoms (bone pain) and those receiving bisphosphonates to normalize serum ALP. However, the most potent agent (zoledronic acid) was not used, and the duration of observation (mean of 3 years with a range of 2 to 5 years) may not be long enough to assess the impact of treatment on long-term outcomes. It seems likely that the restoration of normal bone architecture following suppression of pagetic activity will prevent further deformities and complications.

Agents approved for treatment of Paget's disease suppress the very high rates of bone resorption and secondarily decrease the high rates of bone formation (Table 426e-1). As a result of decreasing bone turnover, pagetic structural patterns, including areas of poorly mineralized woven bone, are replaced by more normal cancellous or lamellar bone. Reduced bone turnover can be documented by a decline in serum ALP and urine or serum resorption markers (N-telopeptide, C-telopeptide).

The first clinically useful agent, etidronate, is now rarely used because the doses required to suppress bone resorption may impair mineralization, necessitating that the drug be given for a maximum of 6 months followed by a 6-month drug-free period. The second-generation oral bisphosphonates—tiludronate, alendronate, and risedronate—are more potent than etidronate in controlling bone turnover and, thus, induce a longer remission at a lower dose. The lower doses reduce the risks of impaired mineralization and osteomalacia. Oral bisphosphonates should be taken first thing in the morning on an empty stomach, followed by maintenance of upright posture with no food, drink, or other medications for 30–60 min. The efficacy of different agents, based on their ability to normalize or decrease ALP levels, is summarized in Table 426e-1, although the response rates are not comparable because they are obtained from different studies.

Intravenous bisphosphonates approved for Paget's disease include pamidronate and zoledronic acid. Although the recommended dose for pamidronate is 30 mg dissolved in 500 mL of normal saline or dextrose IV over 4 h on 3 consecutive days, a more commonly used simpler regimen is a single infusion of 60–90 mg in patients with mild elevation of serum ALP and multiple 90-mg

426e-4 infusions in those with higher levels of ALP. In many patients, particularly those who have severe disease or need rapid normalization of bone turnover (neurologic symptoms, severe bone pain due to a lytic lesion, risk of an impending fracture, or pretreatment prior to elective surgery in an area of active disease), treatment with zoledronic acid is the first choice. It normalizes ALP in about 90% of patients by 6 months, and the therapeutic effect persists for at least 6 more months in most patients. About 10–20% of patients experience a flulike syndrome after the first infusion, which can be partly ameliorated by pretreatment with acetaminophen or nonsteroidal anti-inflammatory drugs (NSAIDs). In patients with high bone turnover, vitamin D and calcium should be provided to prevent hypocalcemia and secondary hyperparathyroidism. Remission following treatment with IV bisphosphonates, particularly zoledronic acid, may persist for well over 1 year. Bisphosphonates should not be used in patients with renal insufficiency (glomerular filtration rate <35 mL/min).

The subcutaneous injectable form of salmon calcitonin is approved for the treatment of Paget's disease. The common side effects of calcitonin therapy are nausea and facial flushing. Secondary resistance after prolonged use may be due to either the formation of anticalcitonin antibodies or downregulation of osteoclastic cell-surface calcitonin receptors. The lower potency and injectable mode of delivery make this agent a less attractive treatment option that should be reserved for patients who either do not tolerate bisphosphonates or have a contraindication to their use. In early reports, denosumab, an antibody to RANKL, has shown promise but has not been approved for this indication.

SCLEROSING BONE DISORDERS

OSTEOPETROSIS

Osteopetrosis refers to a group of disorders caused by severe impairment of osteoclast-mediated bone resorption. Other terms that are often used include marble bone disease, which captures the solid x-ray appearance of the involved skeleton, and Albers-Schonberg disease, which refers to the milder, adult form of osteopetrosis also known as autosomal dominant osteopetrosis type II. The major types of osteopetrosis include malignant (severe, infantile, autosomal recessive) osteopetrosis and benign (adult, autosomal dominant) osteopetrosis types I and II. A rare autosomal recessive intermediate form has a more benign prognosis. Autosomal recessive carbonic anhydrase (CA) II deficiency produces osteopetrosis of intermediate severity associated with renal tubular acidosis and cerebral calcification.

Etiology and Genetics Naturally occurring and gene-knockout animal models with phenotypes similar to those of the human disorders have been used to explore the genetic basis of osteopetrosis. The primary defect in osteopetrosis is the loss of osteoclastic bone resorption and preservation of normal osteoblastic bone formation. Osteoprotegerin (OPG) is a soluble decoy receptor that binds osteoblast-derived RANK ligand, which mediates osteoclast differentiation and activation (Fig. 426e-1). Transgenic mice that overexpress OPG develop osteopetrosis, presumably by blocking RANK ligand. Mice deficient in RANK lack osteoclasts and develop severe osteopetrosis.

Recessive mutations of CA II prevent osteoclasts from generating an acid environment in the clear zone between its ruffled border and the adjacent mineral surface. Absence of CA II, therefore, impairs osteoclastic bone resorption. Other forms of human disease have less clear genetic defects. About one-half of the patients with malignant infantile osteopetrosis have a mutation in the *TCIRG1* gene encoding the osteoclast-specific subunit of the vacuolar proton pump, which mediates the acidification of the interface between bone mineral and the osteoclast ruffled border. Mutations in the *CICN7* chloride channel gene cause autosomal dominant osteopetrosis type II.

Clinical Presentation The incidence of autosomal recessive severe (malignant) osteopetrosis ranges from 1 in 200,000 to 1 in 500,000 live births. As bone and cartilage fail to undergo modeling, paralysis of

one or more cranial nerves may occur due to narrowing of the cranial foramina. Failure of skeletal modeling also results in inadequate marrow space, leading to extramedullary hematopoiesis with hypersplenism and pancytopenia. Hypocalcemia due to lack of osteoclastic bone resorption may occur in infants and young children. The untreated infantile disease is fatal, often before age 5.

Adult (benign) osteopetrosis is an autosomal dominant disease that is usually diagnosed by the discovery of typical skeletal changes in young adults who undergo radiologic evaluation of a fracture. The prevalence is 1 in 100,000 to 1 in 500,000 adults. The course is not always benign, because fractures may be accompanied by loss of vision, deafness, psychomotor delay, mandibular osteomyelitis, and other complications usually associated with the juvenile form. In some kindred, nonpenetrance results in skip generations, while in other families, severely affected children are born into families with benign disease. The milder form of the disease does not usually require treatment.

Radiography Typically, there are generalized symmetric increases in bone mass with thickening of both cortical and trabecular bone. Diaphyses and metaphyses are broadened, and alternating sclerotic and lucent bands may be seen in the iliac crests, at the ends of long bones, and in vertebral bodies. The cranium is usually thickened, particularly at the base of the skull, and the paranasal and mastoid sinuses are underpneumatized.

Laboratory Findings The only significant laboratory findings are elevated serum levels of osteoclast-derived tartrate-resistant acid phosphatase (TRAP) and the brain isoenzyme of creatine kinase. Serum calcium may be low in severe disease, and parathyroid hormone and 1,25-dihydroxyvitamin D levels may be elevated in response to hypocalcemia.

TREATMENT OSTEOPETROSIS

Allogeneic HLA-identical bone marrow transplantation has been successful in some children. Following transplantation, the marrow contains progenitor cells and normally functioning osteoclasts. A cure is most likely when children are transplanted before age 4. Marrow transplantation from nonidentical HLA-matched donors has a much higher failure rate. Limited studies in small numbers of patients have suggested variable benefits following treatment with interferon γ -1 β , 1,25-dihydroxyvitamin D (which stimulates osteoclasts directly), methylprednisolone, and a low-calcium/high-phosphate diet.

Surgical intervention is indicated to decompress optic or auditory nerve compression. Orthopedic management is required for the surgical treatment of fractures and their complications including malunion and postfracture deformity.

PYKNODYSTOSIS

This is an autosomal recessive form of osteosclerosis that is believed to have affected the French impressionist painter Henri de Toulouse-Lautrec. The molecular basis involves mutations in the gene that encodes cathepsin K, a lysosomal metalloproteinase highly expressed in osteoclasts and important for bone-matrix degradation. Osteoclasts are present but do not function normally. Pyknodysostosis is a form of short-limb dwarfism that presents with frequent fractures but usually a normal life span. Clinical features include short stature; kyphoscoliosis and deformities of the chest; high arched palate; proptosis; blue sclerae; dysmorphic features including small face and chin, frontooccipital prominence, pointed beaked nose, large cranium, and obtuse mandibular angle; and small, square hands with hypoplastic nails. Radiographs demonstrate a generalized increase in bone density, but in contrast to osteopetrosis, the long bones are normally shaped. Separated cranial sutures, including the persistent patency of the anterior fontanel, are characteristic of the disorder. There may also be hypoplasia of the sinuses, mandible, distal clavicles, and terminal phalanges. Persistence of deciduous teeth and sclerosis of the calvarium and base of the skull

are also common. Histologic evaluation shows normal cortical bone architecture with decreased osteoblastic and osteoclastic activities. Serum chemistries are normal, and unlike osteopetrosis, there is no anemia. There is no known treatment for this condition, and there are no reports of attempted bone marrow transplant.

PROGRESSIVE DIAPHYSEAL DYSPLASIA

Also known as *Camurati-Engelmann disease*, progressive diaphyseal dysplasia is an autosomal dominant disorder that is characterized radiographically by diaphyseal hyperostosis and a symmetric thickening and increased diameter of the endosteal and periosteal surfaces of the diaphyses of the long bones, particularly the femur and tibia, and, less often, the fibula, radius, and ulna. The genetic defect responsible for the disease has been localized to the area of chromosome 19q13.2 encoding tumor growth factor (TGF) β 1. The mutation promotes activation of TGF- β 1. The clinical severity is variable. The most common presenting symptoms are pain and tenderness of the involved areas, fatigue, muscle wasting, and gait disturbance. The weakness may be mistaken for muscular dystrophy. Characteristic body habitus includes thin limbs with little muscle mass yet prominent and palpable bones and, when the skull is involved, large head with prominent forehead and proptosis. Patients may also display signs of cranial nerve palsies, hydrocephalus, central hypogonadism, and Raynaud's phenomenon. Radiographically, patchy progressive endosteal and periosteal new bone formation is observed along the diaphyses of the long bones. Bone scintigraphy shows increased radiotracer uptake in involved areas.

Treatment with low-dose glucocorticoids relieves bone pain and may reverse the abnormal bone formation. Intermittent bisphosphonate therapy has produced clinical improvement in a limited number of patients.

HYPEROSTOSIS CORTICALIS GENERALISATA

This is also known as *van Buchem's disease*; it is an autosomal recessive disorder characterized by endosteal hyperostosis in which osteosclerosis involves the skull, mandible, clavicles, and ribs. The major manifestations are due to narrowed cranial foramina with neural compressions that may result in optic atrophy, facial paralysis, and deafness. Adults may have an enlarged mandible. Serum ALP levels may be elevated, which reflect the uncoupled bone remodeling with high osteoblastic formation rates and low osteoclastic resorption. As a result, there is increased accumulation of normal bone. Endosteal hyperostosis with syndactyly, known as *sclerosteosis*, is a more severe form. The genetic defects for both sclerosteosis and van Buchem's disease have been assigned to the same region of the chromosome 17q12-q21. It is possible that both conditions may have deactivating mutations in the *BEER* (bone-expressed equilibrium regulator) gene.

MELORHEOSTOSIS

Melorheostosis (Greek, "flowing hyperostosis") may occur sporadically or follow a pattern consistent with an autosomal recessive disorder. The major manifestation is progressive linear hyperostosis in one or more bones of one limb, usually a lower extremity. The name comes from the radiographic appearance of the involved bone, which resembles melted wax that has dripped down a candle. Symptoms appear during childhood as pain or stiffness in the area of sclerotic bone. There may be associated ectopic soft tissue masses, composed of cartilage or osseous tissue, and skin changes overlying the involved bone, consisting of scleroderma-like areas and hypertrichosis. The disease does not progress in adults, but pain and stiffness may persist. Laboratory tests are unremarkable. No specific etiology is known. There is no specific treatment. Surgical interventions to correct contractures are often unsuccessful.

OSTEOPOIKILOSIS

The literal translation of osteopoikilosis is "spotted bones"; it is a benign autosomal dominant condition in which numerous small, variably shaped (usually round or oval) foci of bony sclerosis are seen in the epiphyses and adjacent metaphyses. The lesions may involve any

bone except the skull, ribs, and vertebrae. They may be misidentified as metastatic lesions. The main differentiating points are that bony lesions of osteopoikilosis are stable over time and do not accumulate radionuclide on bone scanning. In some kindred, osteopoikilosis is associated with connective tissue nevi known as *dermatofibrosis lenticularis disseminata*, also known as *Buschke-Ollendorff syndrome*. Histologic inspection reveals thickened but otherwise normal trabeculae and islands of normal cortical bone. No treatment is indicated.

HEPATITIS C-ASSOCIATED OSTEOSCLEROSIS

Hepatitis C-associated osteosclerosis (HCAO) is a rare acquired diffuse osteosclerosis in adults with prior hepatitis C infection. After a latent period of several years, patients develop diffuse appendicular bone pain and a generalized increase in bone mass with elevated serum ALP. Bone biopsy and histomorphometry reveal increased rates of bone formation, decreased bone resorption with a marked decrease in osteoclasts, and dense lamellar bone. One patient had increased serum OPG levels, and bone biopsy showed large numbers of osteoblasts positive for OPG and reduced osteoclast number. Empirical therapy includes pain control, and there may be beneficial response to bisphosphonate. Long-term antiviral therapy may reverse the bone disease.

DISORDERS ASSOCIATED WITH DEFECTIVE MINERALIZATION

HYPHOPHOSPHATASIA

This is a rare inherited disorder that presents as rickets in infants and children or osteomalacia in adults with paradoxically low serum levels of ALP. The frequency of the severe neonatal and infantile forms is about 1 in 100,000 live births in Canada, where the disease is most common because of its high prevalence among Mennonites and Hutterites. It is rare in African Americans. The severity of the disease is remarkably variable, ranging from intrauterine death associated with profound skeletal hypomineralization at one extreme to premature tooth loss as the only manifestation in some adults. Severe cases are inherited in an autosomal recessive manner, but the genetic patterns are less clear for the milder forms. The disease is caused by a deficiency of tissue nonspecific (bone/liver/kidney) ALP (TNSALP), which, although ubiquitous, results only in bone abnormalities. Protein levels and functions of the other ALP isozymes (germ cell, intestinal, placental) are normal. Defective ALP permits accumulation of its major naturally occurring substrates including phosphoethanolamine (PEA), inorganic pyrophosphate (PPi), and pyridoxal 5'-phosphate (PLP). The accumulation of PPi interferes with mineralization through its action as a potent inhibitor of hydroxyapatite crystal growth.

Perinatal hypophosphatasia becomes manifest during pregnancy and is often complicated by polyhydramnios and intrauterine death. The infantile form becomes clinically apparent before the age of 6 months with failure to thrive, rachitic deformities, functional craniosynostosis despite widely open fontanels (which are actually hypomineralized areas of the calvarium), raised intracranial pressure, and flail chest with predisposition to pneumonia. Hypercalcemia and hypercalciuria are common. This form has a mortality rate of about 50%. Prognosis seems to improve for the children who survive infancy. Childhood hypophosphatasia has variable clinical presentation. Premature loss of deciduous teeth (before age 5) is the hallmark of the disease. Rickets causes delayed walking with waddling gait, short stature, and dolichocephalic skull with frontal bossing. The disease often improves during puberty but may recur in adult life. Adult hypophosphatasia presents during middle age with painful, poorly healing metatarsal stress fractures or thigh pain due to femoral pseudofractures.

Laboratory investigation reveals low ALP levels and normal or elevated levels of serum calcium and phosphorus despite clinical and radiologic evidence of rickets or osteomalacia. Serum parathyroid hormone, 25-hydroxyvitamin D, and 1,25-dihydroxyvitamin D levels are normal. The elevation of PLP is specific for the disease and may even be present in asymptomatic parents of severely affected children. Because vitamin B₆ increases PLP levels, vitamin B₆ supplements should be discontinued 1 week before testing. Clinical testing is available to

426e-6 detect loss-of-function mutation(s) within the *ALPL* gene that encodes TNSALP.

There is no established medical therapy. In contrast to other forms of rickets and osteomalacia, calcium and vitamin D supplementation should be avoided because they may aggravate hypercalcemia and hypercalciuria. A low-calcium diet, glucocorticoids, and calcitonin have been used in a small number of patients with variable responses. Because fracture healing is poor, placement of intramedullary rods is best for acute fracture repair and for prophylactic prevention of fractures.

AXIAL OSTEOMALACIA

This is a rare disorder characterized by defective skeletal mineralization despite normal serum calcium and phosphate levels. Clinically, the disorder presents in middle-aged or elderly men with chronic axial skeletal discomfort. Cervical spine pain may also be present. Radiographic findings are mainly osteosclerosis due to coarsened trabecular patterns typical of osteomalacia. Spine, pelvis, and ribs are most commonly affected. Histologic changes show defective mineralization and flat, inactive osteoblasts. The primary defect appears to be an acquired defect in osteoblast function. The course is benign, and there is no established treatment. Calcium and vitamin D therapies are not effective.

FIBROGENESIS IMPERFECTA OSSIUM

This is a rare condition of unknown etiology. It presents in both sexes; in middle age or later; and with progressive, intractable skeletal pain and fractures; worsening immobilization; and a debilitating course. Radiographic evaluation reveals generalized osteomalacia, osteopenia, and occasional pseudofractures. Histologic features include a tangled pattern of collagen fibrils with abundant osteoblasts and osteoclasts. There is no effective treatment. Spontaneous remission has been reported in a small number of patients. Calcium and vitamin D have not been beneficial.

FIBROUS DYSPLASIA AND MCCUNE-ALBRIGHT SYNDROME

Fibrous dysplasia is a sporadic disorder characterized by the presence of one (monostotic) or more (polyostotic) expanding fibrous skeletal lesions composed of bone-forming mesenchyme. The association of the polyostotic form with café au lait spots and hyperfunction of an endocrine system such as pseudoprecocious puberty of ovarian origin is known as *McCune-Albright syndrome* (MAS). A spectrum of the phenotypes is caused by activating mutations in the *GNAS1* gene, which encodes the α subunit of the stimulatory G protein ($G_s\alpha$). As the postzygotic mutations occur at different stages of early development, the extent and type of tissue affected are variable and explain the mosaic pattern of skin and bone changes. GTP binding activates the $G_s\alpha$ regulatory protein and mutations in regions of $G_s\alpha$ that selectively inhibit GTPase activity, which results in constitutive stimulation of the cyclic AMP–protein kinase A signal transduction pathway. Such mutations of the $G_s\alpha$ protein–coupled receptor may cause autonomous function in bone (parathyroid hormone receptor); skin (melanocyte-stimulating hormone receptor); and various endocrine glands including ovary (follicle-stimulating hormone receptor), thyroid (thyroid-stimulating hormone receptor), adrenal (adrenocorticotropic hormone receptor), and pituitary (growth hormone-releasing hormone receptor). The skeletal lesions are composed largely of mesenchymal cells that do not differentiate into osteoblasts, resulting in the formation of imperfect bone. In some areas of bone, fibroblast-like cells develop features of osteoblasts in that they produce extracellular matrix that organizes into woven bone. Calcification may occur in some areas. In other areas, cells have features of chondrocytes and produce cartilage-like extracellular matrix.

Clinical Presentation Fibrous dysplasia occurs with equal frequency in both sexes, whereas MAS with precocious puberty is more common (10:1) in girls. The monostotic form is the most common and is usually diagnosed in patients between 20 and 30 years of age without

associated skin lesions. The polyostotic form typically manifests in children <10 years old and may progress with age. Early-onset disease is generally more severe. Lesions may become quiescent in puberty and progress during pregnancy or with estrogen therapy. In polyostotic fibrous dysplasia, the lesions most commonly involve the maxilla and other craniofacial bones, ribs, and metaphyseal or diaphyseal portions of the proximal femur or tibia. Expanding bone lesions may cause pain, deformity, fractures, and nerve entrapment. Sarcomatous degeneration involving the facial bones or femur is infrequent (<1%). The risk of malignant transformation is increased by radiation, which has proven to be ineffective treatment. In rare patients with widespread lesions, renal phosphate wasting and hypophosphatemia may cause rickets or osteomalacia. Hypophosphatemia may be due to production of a phosphaturic factor by the abnormal fibrous tissue.

MAS patients may have café au lait spots, which are flat, hyperpigmented skin lesions that have rough borders (“coast of Maine”) in contrast to the café au lait lesions of neurofibromatosis that have smooth borders (“coast of California”). The most common endocrinopathy is isosexual pseudoprecocious puberty in girls. Other less common endocrine disorders include thyrotoxicosis, Cushing’s syndrome, acromegaly, hyperparathyroidism, hyperprolactinemia, and pseudoprecocious puberty in boys.

Radiographic Findings In long bones, the fibrous dysplastic lesions are typically well-defined, radiolucent areas with thin cortices and a ground-glass appearance. Lesions may be lobulated with trabeculated areas of radiolucency (Fig. 426e-4). Involvement of facial bones usually presents as radiodense lesions, which may create a leonine appearance (leontiasis osea). Expansile cranial lesions may narrow foramina and cause optic lesions, reduce hearing, and create other manifestations of cranial nerve compression.

Laboratory Results Serum ALP is occasionally elevated but calcium, parathyroid hormone, 25-hydroxyvitamin D, and 1,25-dihydroxyvitamin D levels are normal. Patients with extensive polyostotic lesions may have hypophosphatemia, hyperphosphaturia, and osteomalacia. The hypophosphatemia and phosphaturia are directly related to the levels of fibroblast growth factor 23 (FGF23). Biochemical markers of bone turnover may be elevated.



FIGURE 426e-4 Radiograph of a 16-year-old male with fibrous dysplasia of the right proximal femur. Note the multiple cystic lesions, including the large lucent lesion in the proximal midshaft with scalloping of the interior surface. The femoral neck contains two lucent cystic lesions.

TREATMENT FIBROUS DYSPLASIA AND McCUNE-ALBRIGHT SYNDROME

Spontaneous healing of the lesions does not occur, and there is no established effective treatment. Improvement in bone pain and partial or complete resolution of radiographic lesions have been reported after IV bisphosphonate therapy. Surgical stabilization is used to prevent pathologic fracture or destruction of a major joint space and to relieve nerve root or cranial nerve compression or sinus obstruction.

OTHER DYSPLASIAS OF BONE AND CARTILAGE

PACHYDERMOPERIOSTOSIS

Pachydermoperiostosis, or hypertrophic osteoarthropathy (primary or idiopathic), is an autosomal dominant disorder characterized by periosteal new bone formation that involves the distal extremities. The lesions present as clubbing of the digits and hyperhidrosis and thickening of the skin, primarily of the face and forehead. The changes usually appear during adolescence, progress over the next decade, and then become quiescent. During the active phase, progressive enlargement of the hands and feet produces a paw-like appearance, which may be mistaken for acromegaly. Arthralgias, pseudogout, and limited mobility may also occur. The disorder must be differentiated from secondary hypertrophic osteopathy that develops during the course of serious pulmonary disorders. The two conditions can be differentiated by standard radiography of the digits in which secondary pachydermoperiostosis has exuberant periosteal new bone formation and a smooth and undulating surface. In contrast, primary hypertrophic osteopathy has an irregular periosteal surface.

There are no diagnostic blood or urine tests. Synovial fluid does not have an inflammatory profile. There is no specific therapy, although a limited experience with colchicine suggests some benefit in controlling the arthralgias.

OSTEOCHONDRODYSPLASIAS

These include several hundred heritable disorders of connective tissue. These primary abnormalities of cartilage manifest as disturbances in cartilage and bone growth. Selected growth-plate chondrodysplasias are described here. [For discussion of chondrodysplasias, see Chap. 427.](#)

Achondrodysplasia This is a relatively common form of short-limb dwarfism that occurs in 1 in 15,000 to 1 in 40,000 live births. The disease is caused by a mutation of the fibroblast growth factor receptor 3 (*FGFR3*) gene that results in a gain-of-function state. Most cases are sporadic mutations. However, when the disorder appears in families, the inheritance pattern is consistent with an autosomal dominant disorder. The primary defect is abnormal chondrocyte proliferation at the growth plate that causes development of short, but proportionately thick, long bones. Other regions of the long bones may be relatively unaffected. The disorder is manifest by the presence of short limbs (particularly the proximal portions), normal trunk, large head, saddle nose, and an exaggerated lumbar lordosis. Severe spinal deformity may lead to cord compression. The homozygous disorder is more serious than the sporadic form and may cause neonatal death. Pseudoachondroplasia clinically resembles achondrodysplasia but has no skull abnormalities.

Enchondromatosis This is also called *dyschondroplasia* or *Ollier's disease*; it is also a disorder of the growth plate in which the primary cartilage is not resorbed. Cartilage ossification proceeds normally, but it is not resorbed normally, leading to cartilage accumulation. The changes are most marked at the ends of long bones, where the highest growth rates occur. Chondrosarcoma develops infrequently. The association of enchondromatosis and cavernous hemangiomas of the skin and soft tissues is known as *Maffucci's syndrome*. Both Ollier's disease and Maffucci's syndrome are associated with various malignancies, including granulosa cell tumor of the ovary and cerebral glioma.

Multiple Exostoses This is also called *diaphyseal aclasis* or *osteochondromatosis*; it is a genetic disorder that follows an autosomal dominant pattern of inheritance. In this condition, areas of growth plates become displaced, presumably by growing through a defect in the perichondrium. The lesion begins with vascular invasion of the growth-plate cartilage, resulting in a characteristic radiographic finding of a mass that is in direct communication with the marrow cavity of the parent bone. The underlying cortex is resorbed. The disease is caused by inactivating mutations of the *EXT1* and *EXT2* genes, whose products normally regulate processing of chondrocyte cytoskeletal proteins. The products of the *EXT* gene likely function as tumor suppressors, with the loss-of-function mutation resulting in abnormal proliferation of growth-plate cartilage. Solitary or multiple lesions are located in the metaphyses of long bones. Although usually asymptomatic, the lesions may interfere with joint or tendon function or compress peripheral nerves. The lesions stop growing when growth ceases but may recur during pregnancy. There is a small risk for malignant transformation into chondrosarcoma.

EXTRASKELETAL (ECTOPIC) CALCIFICATION AND OSSIFICATION

Deposition of calcium phosphate crystals (*calcification*) or formation of true bone (*ossification*) in nonosseous soft tissue may occur by one of three mechanisms: (1) metastatic calcification due to a supranormal calcium \times phosphate concentration product in extracellular fluid; (2) dystrophic calcification due to mineral deposition into metabolically impaired or dead tissue despite normal serum levels of calcium and phosphate; and (3) ectopic ossification, or true bone formation. Disorders that may cause extraskeletal calcification or ossification are listed in [Table 426e-2](#).

METASTATIC CALCIFICATION

Soft tissue calcification may complicate diseases associated with significant hypercalcemia, hyperphosphatemia, or both. In addition, vitamin D and phosphate treatments or calcium administration in the presence of mild hyperphosphatemia, such as during hemodialysis, may induce ectopic calcification. Calcium phosphate precipitation may complicate any disorder when the serum calcium \times phosphate concentration product is >75 . The initial calcium phosphate deposition is in the form of small, poorly organized crystals, which subsequently organize into hydroxyapatite crystals. Calcifications that occur in hypercalcemic states with normal or low phosphate have a predilection for kidney, lungs, and gastric mucosa. Hyperphosphatemia with normal or low serum calcium may promote soft tissue calcification with predilection for the kidney and arteries. The disturbances of calcium and phosphate in renal failure and hemodialysis are common causes of soft tissue (metastatic) calcification.

TABLE 426e-2 DISEASES AND CONDITIONS ASSOCIATED WITH ECTOPIC CALCIFICATION AND OSSIFICATION

Metastatic calcification	Dystrophic calcification
Hypercalcemic states	Inflammatory disorders
Primary hyperparathyroidism	Scleroderma
Sarcoidosis	Dermatomyositis
Vitamin D intoxication	Systemic lupus erythematosus
Milk-alkali syndrome	Trauma-induced
Renal failure	Ectopic ossification
Hyperphosphatemia	Myositis ossificans
Tumoral calcinosis	Postsurgery
Secondary hyperparathyroidism	Burns
Pseudohypoparathyroidism	Neurologic injury
Renal failure	Other trauma
Hemodialysis	Fibrodysplasia ossificans
Cell lysis following chemotherapy	progressiva
Therapy with vitamin D and phosphate	

This is a rare genetic disorder characterized by masses of metastatic calcifications in soft tissues around major joints, most often shoulders, hips, and ankles. Tumoral calcinosis differs from other disorders in that the periarticular masses contain hydroxyapatite crystals or amorphous calcium phosphate complexes, while in fibrodysplasia ossificans progressiva (below), true bone is formed in soft tissues. About one-third of tumoral calcinosis cases are familial, with both autosomal recessive and autosomal dominant modes of inheritance reported. The disease is also associated with a variably expressed abnormality of dentition marked by short bulbous roots, pulp calcification, and radicular dentin deposited in swirls. The primary defect responsible for the metastatic calcification appears to be hyperphosphatemia resulting from the increased capacity of the renal tubule to reabsorb filtered phosphate. Spontaneous soft tissue calcification is related to the elevated serum phosphate, which, along with normal serum calcium, exceeds the concentration product of 75.

All of the North American patients reported have been African American. The disease usually presents in childhood and continues throughout the patient's life. The calcific masses are typically painless and grow at variable rates, sometimes becoming large and bulky. The masses are often located near major joints but remain extracapsular. Joint range of motion is not usually restricted unless the tumors are very large. Complications include compression of neural structures and ulceration of the overlying skin with drainage of chalky fluid and risk of secondary infection. Small deposits not detected by standard radiographs may be detected by ^{99m}Tc bone scanning. The most common laboratory findings are hyperphosphatemia and elevated serum 1,25-dihydroxyvitamin D levels. Serum calcium, parathyroid hormone, and ALP levels are usually normal. Renal function is also usually normal. Urine calcium and phosphate excretions are low, and calcium and phosphate balances are positive.

An acquired form of the disease may occur with other causes of hyperphosphatemia, such as secondary hyperparathyroidism associated with hemodialysis, hypoparathyroidism, pseudohypoparathyroidism, and massive cell lysis following chemotherapy for leukemia. Tissue trauma from joint movement may contribute to the periarticular calcifications. Metastatic calcifications are also seen in conditions associated with hypercalcemia, such as in sarcoidosis, vitamin D intoxication, milk-alkali syndrome, and primary hyperparathyroidism. In these conditions, however, mineral deposits are more likely to occur in proton-transporting organs such as kidney, lungs, and gastric mucosa in which an alkaline milieu is generated by the proton pumps.

TREATMENT TUMORAL CALCINOSIS

Therapeutic successes have been achieved with surgical removal of subcutaneous calcified masses, which tend not to recur if all calcification is removed from the site. Reduction of serum phosphate

by chronic phosphorus restriction may be accomplished using low dietary phosphorus intake alone or in combination with oral phosphate binders. The addition of the phosphaturic agent acetazolamide may be useful. Limited experience using the phosphaturic action of calcitonin deserves further testing.

DYSTROPHIC CALCIFICATION

Posttraumatic calcification may occur with normal serum calcium and phosphate levels and normal ion-solubility product. The deposited mineral is either in the form of amorphous calcium phosphate or hydroxyapatite crystals. Soft tissue calcification complicating connective tissue disorders such as scleroderma, dermatomyositis, and systemic lupus erythematosus may involve localized areas of the skin or deeper subcutaneous tissue and is referred to as *calcinosis circumscripta*. Mineral deposition at sites of deeper tissue injury including periarticular sites is called *calcinosis universalis*.

ECTOPIC OSSIFICATION

True extraskeletal bone formation that begins in areas of fasciitis following surgery, trauma, burns, or neurologic injury is referred to as *myositis ossificans*. The bone formed is organized as lamellar or trabecular, with normal osteoblasts and osteoclasts conducting active remodeling. Well-developed haversian systems and marrow elements may be present. A second cause of ectopic bone formation occurs in an inherited disorder, *fibrodysplasia ossificans progressiva*.

FIBRODYSPLASIA OSSIFICANS PROGRESSIVA

This is also called *myositis ossificans progressiva*; it is a rare autosomal dominant disorder characterized by congenital deformities of the hands and feet and episodic soft tissue swellings that ossify. Ectopic bone formation occurs in fascia, tendons, ligaments, and connective tissue within voluntary muscles. Tender, rubbery induration, sometimes precipitated by trauma, develops in the soft tissue and gradually calcifies. Eventually, heterotopic bone forms at these sites of soft tissue trauma. Morbidity results from heterotopic bone interfering with normal movement and function of muscle and other soft tissues. Mortality is usually related to restrictive lung disease caused by an inability of the chest to expand. Laboratory tests are unremarkable.

There is no effective medical therapy. Bisphosphonates, glucocorticoids, and a low-calcium diet have largely been ineffective in halting progression of the ossification. Surgical removal of ectopic bone is not recommended, because the trauma of surgery may precipitate formation of new areas of heterotopic bone. Dental complications including frozen jaw may occur following injection of local anesthetics. Thus, CT imaging of the mandible should be undertaken to detect early sites of soft tissue ossification before they are appreciated by standard radiography.

SECTION 5 DISORDERS OF INTERMEDIARY METABOLISM

427 Heritable Disorders of Connective Tissue

Darwin J. Prockop, John F. Bateman

CLASSIFICATION OF CONNECTIVE TISSUE DISORDERS

Some of the most common conditions that are transmitted genetically in families are disorders that produce clinically obvious changes in the skeleton, skin, or other relatively acellular tissues that have been loosely defined as connective tissues. Because of their heritability, the disorders were recognized as potentially traceable to mutated genes soon after the principles of genetics were introduced into medicine. In the last several decades, many of these disorders have been linked to mutations in several hundred different genes. However, classifying the disorders on the basis of either their clinical presentations or the mutations causing them presents a challenge for both the clinician and the geneticist.

A major development in the field was made by McKusick, who suggested that a group of disorders that included brittle bones in children (osteogenesis imperfecta), hyperextensible skin (Ehlers-Danlos syndrome), and characteristic distortions of skeleton (Marfan's

syndrome) be considered as "heritable disorders of connective tissue" and that mutations causing the disorders would be found in the genes coding for proteins of the tissues.

The information on the disorders has continued to develop on two levels. The initial clinical classifications suggested by McKusick, and others, had to be refined as additional patients were examined. For example, some patients had skin changes similar to those commonly seen in Ehlers-Danlos syndrome, but this feature was overshadowed by other features such as extreme hypotonia or sudden rupture of large blood vessels. To account for the full spectrum of presentations in patients and families, many of the disorders have been reclassified several times, and each has been divided into a series of subtypes. For example, a recent effort to classify all the heritable disorders that alter the skeleton defined 456 distinctive conditions that were divided into 40 major groups.

The identification of mutations causing the diseases has developed on a parallel track. The first genes cloned for connective tissues were the two genes coding for type I collagen, the most abundant protein in bones, skin, tendons, and several other tissues. Some of the first assays in patients with osteogenesis imperfecta (OI) revealed mutations in type I collagen genes. Biochemical data developed using cultures of skin fibroblasts from affected patients demonstrated that the mutations dramatically altered the synthesis or structure of collagen fibers. The results stimulated efforts to identify additional mutations in genes

coding for structural proteins. Genes for collagens provided an attractive paradigm to search for mutations, since a series of different types of collagens were found in different connective tissues and the collagen genes were readily isolated by their unique signature sequences. Also, the collagen genes are particularly vulnerable to a large number of different mutations because of unusual structural requirements of the protein. The search for mutations in collagen genes proved fruitful in that mutations were found in most patients with OI, in many patients with hyperextensible skin, in some patients with dwarfism, and in patients with other disorders, including Alport's syndrome, that were not initially classified as disorders of connective tissue. Also, mutations in collagen genes were found in subset of patients with a diagnosis of osteoarthritis and a subset of patients with the diagnosis of osteoporosis. However, the search for mutations quickly expanded to hundreds of other genes that included those for other structural proteins, for the posttranslational processing of structural proteins, and for growth factors and their receptors, and other genes whose functions are still not fully understood.

In many instances, the mutations helped to define the clinical subtype of the disorder. In others, however, they did not. Some patients with the same clinical presentations were found to have mutations in different genes. Also, some patients with different manifestations were found to have mutations in the same genes. In addition, it was difficult to establish whether a change in the structure of a gene caused the phenotypic changes in patients and was not simply a neutral polymorphism. Therefore, there has been a continuing debate as to whether the disorders should be classified by their clinical presentations or by the genetic abnormalities. As an illustration of the problem, mutations in 226 genes have been found to be associated with the 456 defined disorders of the skeleton, but the latest nosology remains a "hybrid" between a list of clinically defined disorders, waiting for molecular clarification,

and an annotated database documenting the phenotypic spectrum produced by mutations in a given gene. A simpler system of classification proved feasible for one rare heritable disorder of skin, epidermolysis bullosa. The disorder was first defined by the presence of friction-induced blister. It was then divided into subtypes that were defined by the ultrastructural layers of the skin that cleaved and blistered. Most patients in each subtype were subsequently shown to have mutations in genes expressed in the corresponding layer of skin. Even with these patients, however, the strength of the genotype-phenotype correlation varies, and mutations have not yet been found in every patient.

In the end, consensus reports by experts in the field and sources such as the Online Mendelian Inheritance in Man database provide valuable resources for physicians searching for diagnoses of patients with unusual clinical features. However, patients with the most common forms of the disorders have mutations in a limited number of genes. This chapter will focus primarily on these more common disorders.

COMPOSITION OF CONNECTIVE TISSUES

Connective tissues such as skin, bone, cartilage, ligaments, and tendons are the critical structural frameworks of the body important for development and function. They consist of a complex interacting extracellular matrix network of collagens, proteoglycans, and a large number of noncollagenous glycoproteins and proteins. Although these precise combinations of up to ~500 potential extracellular matrix building blocks provide tissue-specific function, there are many overarching similarities in composition, such as the role of composite collagen fibrils in providing strength and form, elastin fibrils and proteoglycans and other interacting proteins, and glycoproteins that fine-tune function (Table 427-1). The most abundant components are three similar fibrillar collagens (types I, II, and III). They have a similar tensile

TABLE 427-1 CONSTITUENTS OF CONNECTIVE TISSUES

Connective Tissue	Major Constituents	Approximate Amounts, % dry wt	Characteristics or Functions
Dermis, ligaments, tendons	Type I collagen	80	Large bundles of fibrils
	Type III collagen	5–15	Thin fibrils
	Type IV collagen, laminins, and nidogen	<5	Form basal laminae under epithelium
	Types V, VI, and VII collagens	<5	V modifies type I fibrils; VI forms beaded microfibrils; VII forms anchoring fibrils for epidermis
	Fibrillin aggregates/elastin	<5	Provide elasticity
	Fibronectin	<5	Associated with collagen fibers and cell surfaces
	Proteoglycans ^a /hyaluronan	<0.5	Provide resiliency
Bone (demineralized)	Type I collagen	90	Complex fibril network
	Type VI collagen	1–2	Beaded microfibrils
	Proteoglycans ^a /hyaluronan	1	Function unclear
	Osteonectin, osteopontin, osteocalcin, α2-glycoprotein, and sialoproteins	1–5	May regulate mineralization
Aorta	Type I collagen	20–40	Fibril network
	Type III collagen	20–40	Thin fibrils
	Fibrillin aggregates/elastin	20–40	Provide elasticity
	Type IV collagen, laminins, and nidogen	<5	Form basal lamina under endothelial cells
	Types V and VI collagens	<2	V modifies type I fibrils; VI forms beaded microfibrils
	Proteoglycans ^a /hyaluronan	<3	Provide resiliency
Cartilage	Type II collagen	40–50	Arcades of thin fibrils
	Type IX collagen	5–10	Links type II fibrils and other components
	Type VI collagen	<1	Beaded microfibrils, largely pericellular
	Type X collagen	5–10	Forms pericellular network in hypertrophic growth plate cartilage
	Type XI collagen	<10	Incorporated into some type II fibrils
	Proteoglycans ^a /hyaluronan	15–50	Provide resiliency
	Small leucine-rich repeat proteins (SLRPs; >6 kinds)	<5	Multiple functions in assembly and function of the tissue

^aOver 30 proteoglycans have been identified. They differ in the structures of their core proteins and their contents of glycosaminoglycan side chains of chondroitin-4-sulfate, chondroitin-6-sulfate, dermatan sulfate, and keratan sulfate. Basal lamina contains a proteoglycan with a side chain of heparan sulfate that resembles heparin.

2506 strength as steel wires. The three fibrillar collagens are distributed in a tissue-specific manner: type I collagen accounts for most of the protein of dermis, ligaments, tendons, and demineralized bone; type I and type III are the most abundant proteins of large blood vessels; and type II is the most abundant protein of cartilage.

BIOSYNTHESIS AND TURNOVER OF CONNECTIVE TISSUES

Connective tissues are among the most stable components in living organisms, but they are not inert. During embryonic development, connective tissue membranes appear as early as the four-cell blastocyst to provide strength and a structural scaffold for the developing embryo. With the development of blood vessels and skeleton, there is a rapid increase in the synthesis, degradation, and resynthesis of connective tissues. The turnover continues at a slower, but still rapid pace throughout postnatal development and then spikes during the growth spurt of puberty. During adulthood, the metabolic turnover of most connective tissues is slow, but it continues at a moderate pace in bone. With age, malnutrition, physical inactivity, and low gravitational stress, the rate of degradation of most connective tissues, especially in bone and skin, begins to exceed the rate of synthesis and the tissues shrink. In starvation, a large fraction of the collagen in skin and other connective tissues is degraded and provides amino acids for gluconeogenesis (Chap. 97). In both osteoarthritis and rheumatoid arthritis, there is extensive degradation of articular cartilage collagen. Glucocorticoids weaken most tissues by decreasing collagen synthesis. In many pathologic states, however, collagen is deposited in excess. With most injuries to tissues, inflammatory and immune responses stimulate the deposition of collagen fibrils in the form of fibrotic scars. The deposition of the fibrils is largely irreversible and prevents regeneration of normal tissues in hepatic cirrhosis, pulmonary fibrosis, atherosclerosis, and nephrosclerosis.

Structure and Biosynthesis of Fibrillar Collagens The tensile strength of collagen fibers derives primarily from the self-assembly of protein monomers into large fibril structures in a process that resembles crystallization. The self-assembly requires monomers of highly uniform and relatively rigid structure. It also requires a complex series of post-translational processing steps that maintain the solubility of the monomers until they are transported to the appropriate extracellular sites for fibril assembly. Because of the stringent requirements for correct self-assembly, it is not surprising that mutations in genes for fibrillar collagens cause many of the diseases of connective tissues.

The monomers of the three fibrillar collagens are formed from three polypeptide chains, called α chains, that are wrapped around each other into a rope-like triple-helical conformation. The triple helix is a unique structure among proteins, and it provides rigidity to the molecule. It also orients the side chains of amino acids in an “inside out” manner relative to most other proteins so that the charged and hydrophobic residues on the surface can direct self-assembly of the monomers into fibrils. The triple-helical conformation of the monomer is generated because each of the α chains has a repetitive amino acid sequence in which glycine (Gly) appears as every third amino acid. Each α chain contains about 1000 amino acids. Therefore, the sequence of each α chain can be designated as $(-\text{Gly-X-Y})_n$, where X and Y represent amino acids other than glycine and n is >338 . The presence of glycine, the smallest amino acid, in every third position in the sequence is critical because this residue must fit in a sterically restricted space in the middle of the helix where the three chains come together. The requirement for a glycine residue at every third position explains the severe effects of mutations that convert any of the glycine residues to an amino acid with a bulkier side chain (see below). Many of the X- and Y-position amino acids are proline and hydroxyproline, which, because of their ring structures, provide additional rigidity to the triple helix. Other X- and Y-positions are occupied by charged or hydrophobic amino acids that precisely direct lateral and longitudinal assembly of the monomers into highly ordered fibrils. Mutations that substitute amino acids in some X- and Y-positions can, in rare instances, also produce genetic diseases.

The fibers formed by the three fibrillar collagens differ in thickness and length, but they have a similar fine structure. As viewed by electron microscopy, they all have a characteristic pattern of cross-striations that are about one-quarter the length of the monomers and reflect the precise packing into fibrils. The three fibrillar collagens, however, differ in sequences found in the X- and Y-positions of the α chains and therefore in some of their physical properties. Type I collagen is composed of two identical $\alpha 1(\text{I})$ chains and a third $\alpha 2(\text{I})$ chain that differs slightly in its amino acid sequence. Type II collagen is composed of three identical $\alpha(\text{II})$ chains. Type III collagen is composed of three identical $\alpha 1(\text{III})$ chains.

To deliver a monomer of the correct structure to the appropriate site of fibril assembly, the biosynthesis of fibrillar collagens involves a large number of unique processing steps (Fig. 427-1). The monomer, first synthesized as a soluble precursor called *procollagen*, contains an additional globular domain at each end. As the pro α chains of procollagen are synthesized on ribosomes, the free N-terminal ends move into the cisternae of the rough endoplasmic reticulum. Signal peptides at the N-termini are cleaved, and additional posttranslational reactions begin. Proline residues in the Y-position of the repeating -Gly-X-Y- sequences are converted to hydroxyproline by the enzyme prolyl hydroxylase. The hydroxylation of prolyl residues is essential for the three α chains of the monomer to fold into a triple helix at body temperature. The enzyme requires ascorbic acid as one of its essential cofactors, an observation that explains why wounds fail to heal in scurvy (Chap. 96e). In scurvy, some of the underhydroxylated and unfolded protein accumulates in the cisternae of the rough endoplasmic reticulum and is degraded. Lysine residues in the Y-position are also hydroxylated to hydroxylysine by a separate lysyl hydroxylase. Many of the hydroxylysine residues are glycosylated with galactose or with galactose and glucose. A large mannose-rich oligosaccharide is assembled on the C-terminal propeptide of each chain. The pro α chains are assembled by interactions among these C-terminal propeptides that control the selection of the appropriate partner chains to form hetero- or homotrimers and provide the correct chain registration required for subsequent formation of the collagen triple helix. After the C-terminal propeptides assemble the three pro α chains, a nucleus of triple helix is formed near the C terminus, and the helical conformation is propagated toward the N terminus in a zipper-like manner that resembles crystallization. The folding into the triple helix is spontaneous in solution, but as discussed below, identification of rare mutations causing OI demonstrated that the folding *in cellulo* is assisted by ancillary proteins. The fully folded protein is then secreted. After secretion, procollagen is processed to collagen by cleavage of the N-propeptides and C-propeptides by two specific proteinases. The release of the propeptides decreases the solubility of the protein about 1000-fold. The entropic energy that is released drives the self-assembly of the collagen into fibrils. Self-assembled collagen fibers have considerable tensile strength, but their strength is increased further by cross-linking reactions that form covalent bonds between α chains in one molecule and α chains in adjacent molecules.

Although the assembly of collagen monomers into fibers is a spontaneous reaction, the process in tissues is modulated by the presence of less abundant collagens (type V with type I, and type XI with type II) and by other components such as a series of small leucine-rich proteins (SLRPs). Some of the less abundant components alter the rate of fibril assembly, whereas others change the morphology of the fibers or their interactions with cells and other molecules.

Collagen fibers are resistant to most proteases, but during degradation of connective tissues, they are cleaved by specific matrix metalloproteinases (collagenases) that cause partial unfolding of the triple helices into gelatin-like structures that are further degraded by less specific proteinases.

OTHER COLLAGENS AND RELATED MOLECULES

The unique properties of the triple helix are used to define a family of at least 28 collagens that contain repetitive -Gly-X-Y- sequences and form triple helices of varying length and complexity. The proteins are

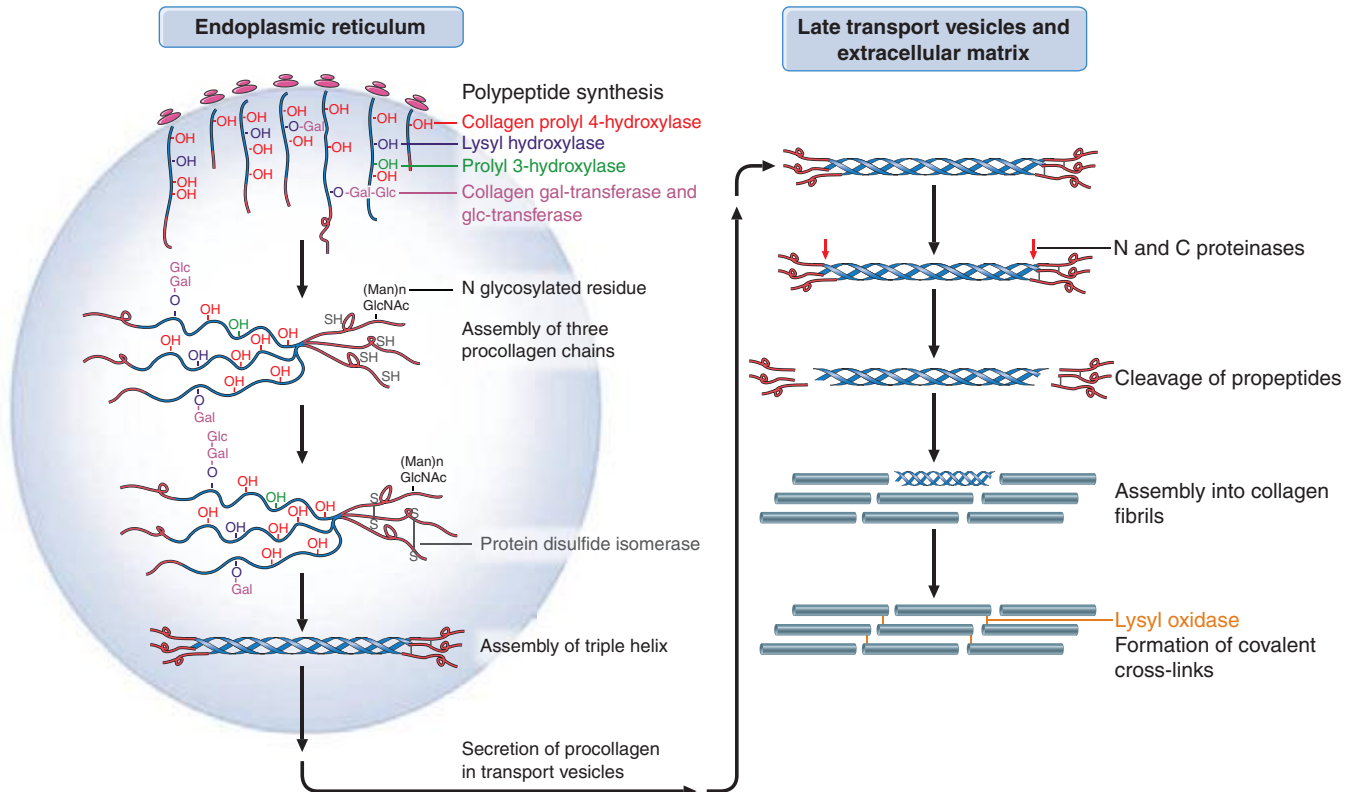


FIGURE 427-1 Schematic summary of biosynthesis of fibrillar collagens. (Modified and reproduced with permission from J Myllyharju, KI Kivirikko: *Trends in Genetics* 20:33, 2004.)

heterogeneous both in structure and function, and many are the sites of mutations causing genetic diseases. For example, the type IV collagen found in basement membranes is composed of three α chains synthesized from any of six different genes. Mutations in any of the six genes can cause Alport's syndrome.

Fibrillin Aggregates and Elastin In addition to tensile strength, many tissues such as the lung, large blood vessels, and ligaments require elasticity. The elasticity was originally ascribed to an amorphous rubber-like protein named *elastin*. Subsequent analyses, largely sparked by discoveries of mutations causing the Marfan's syndrome (MFS), demonstrated that the elasticity resided in thin fibrils composed primarily of large glycoproteins named *fibrillins*. The fibrillins contain large numbers of epidermal growth factor-like domains interspersed with characteristic cysteine-rich domains that are also found in latent transforming growth factor β (TGF- β) binding proteins. The fibrillins assemble into long beadlike strands that also contain numerous other components including small and variable amounts of elastin, bone morphogenic proteins (BMPs), and microfibril-associated glycoproteins (MAGPs). The principles whereby the fibrils provide elasticity to tissue and their biosynthetic assembly are still under investigation. As well as contributing to extracellular matrix structure, the fibrillins play a major role in TGF- β signaling.

Proteoglycans The resiliency to compression of connective tissues such as cartilage or the aorta is largely explained by the presence of proteoglycans. Proteoglycans are composed of a core protein to which are attached a large series of negatively charged polymers of disaccharides (largely chondroitin sulfates). At least 30 proteoglycans have been identified. They vary in their binding to collagens and other components of matrix, but specific functions have not been assigned to most. The major proteoglycan of cartilage, called *aggrecan*, has a core protein of 2000 amino acids that is decorated with about 100 side chains of chondroitin sulfate and keratin sulfate. The core protein, in turn, binds to long chains of the polymeric disaccharide hyaluronan to form proteoglycan aggregates, one of the largest soluble macromolecular structures

in nature. Because of its highly negative charge and extended structure, the proteoglycan aggregate binds large amounts of water and small ions to distend the three-dimensional arcade of collagen fibers found in the same tissues. It thereby makes the cartilage resilient to pressure.

SPECIFIC DISORDERS

OSTEOGENESIS IMPERFECTA (OI)

The central feature of OI is a severe decrease in bone mass that makes bones brittle. The disorder is frequently associated with blue sclerae, dental abnormalities (dentinogenesis imperfecta), progressive hearing loss, and a positive family history. Most patients have mutations in one of the two genes coding for type I collagen.

Classification OI was originally classified into two subtypes of *congenita* and *tarda* depending on the age of onset of the symptoms. Sillence suggested a series of subtypes based on clinical and radiologic findings and mode of inheritance. As with the other disorders discussed here, the description of rare recessive forms of OI and discovery of mutations in new genes have opened a debate as to whether the disorders should be classified by the clinical phenotypes or by the genes at fault. For the near term, the classification based on the clinical presentations seems the most useful (Table 427-2).

Type I is the mildest subtype and can produce either mild or no apparent deformities of the skeleton. Most patients have distinctly blue sclerae. Type II produces bone so brittle that it is lethal in utero or shortly after birth; it can be subclassified into types IIA, IIB, and IIC, depending on radiologic findings. Of the nonlethal forms, type III is progressively deforming with moderate to severe bone deformity, and type IV (common variable OI with normal sclerae) has mild to moderate bone fragility.

The classifications of patients by types of OI do not consistently predict the clinical course of the disease. Some patients appear normal at birth and become progressively worse; others have multiple fractures in infancy and childhood, improve after puberty, and fracture more frequently later in life. Women are particularly prone to fracture during

TABLE 427-2 CLASSIFICATION OF OSTEOGENESIS IMPERFECTA (OI)

Phenotype	Type	Typical Features	Inheritance	Gene Defect	Protein Defect
Nondeforming OI with blue sclerae	I	Mild to moderate bone fragility, normal or near-normal stature, blue sclerae, normal dentition in most, hearing loss in ~50%	AD	COL1A1 COL1A2	Collagen I haploinsufficiency
Perinatally lethal OI	II	Extreme bone fragility, short stature, long bone bowing, blue sclerae Normal/pale blue sclerae, normal dentition and hearing	AD	COL1A1 COL1A2	Collagen I structural mutations
			AR	<i>CRTAP</i> <i>LEPRE1</i> <i>PPIB</i>	Collagen posttranslational modification and folding machinery
Progressively deforming OI	III	Moderate to severe bone deformity, blue sclerae at birth, hearing loss and abnormal dentition common	AD	COL1A1 COL1A2	Collagen I structural mutations
			AR	<i>CRTAP</i> <i>LEPRE1</i> <i>PPIB</i> <i>FKBP10</i> <i>SERPINH1</i> <i>BMP1</i>	Collagen posttranslational modification and folding machinery
				<i>WNT1</i>	Proteolytic removal of procollagen N-propeptide
				<i>SERPINF1</i>	Wnt cell signaling pathway
				<i>TMEM38B</i>	PEDF - growth factor signaling?
					Cation channel, Ca ²⁺ release
Common variable OI with normal sclerae	IV	Mild to moderate, bone fragility, normal sclerae, variable dentition, hearing loss in <10%	AD	COL1A1 COL1A2	Collagen I structural mutations
			AR	<i>WNT1</i> <i>CRTAP</i> <i>FKBP10</i> <i>SP7/OSX</i>	Wnt cell signaling pathway Collagen posttranslational modification and folding machinery Transcription factor, bone formation defect
OI with calcification of the interosseous membranes	V	Calcification of the interosseous membranes in forearm and legs and/or hypertrophic callus; variable bone deformity, normal sclerae and dentition	AD	<i>IFITM5</i>	Transcription factor, bone formation defect
Bruck syndrome type 1		Contractures with pterygia, fractures in infancy or early childhood, postnatal short stature, severe limb deformity, and progressive scoliosis	AR	<i>FKBP10</i>	Collagen folding machinery
Bruck syndrome type 2		As for Bruck syndrome type 1	AR	<i>PLOD2</i>	Collagen posttranslational modification of lysine

Abbreviations: AD, autosomal dominant; AR, autosomal recessive.

Note: Predominant OI gene mutations (>90%) are in *COL1A1* and *COL1A2* (in bold typeface).

pregnancy and after menopause. A few women from families with mild variants of OI do not develop fractures until after menopause, and their disease may be difficult to distinguish from postmenopausal osteoporosis.

Incidence Type I OI has a frequency of about 1 in 15,000–20,000 births. Type II OI has a reported incidence of about 1 in 60,000. Only a limited number of patients with the severe forms of OI have been reported, and the combined incidence of the severe forms that are recognizable at birth (types II, III, and IV) may be higher than 1 in 60,000.

Skeletal Effects In type I OI, the fragility of bones may be severe enough to limit physical activity or be so mild that individuals are unaware of any disability. Radiographs of the skull in patients with mild disease may show a mottled appearance because of small islands of irregular ossification. In type II OI, ossification of many bones is frequently incomplete. Continuously beaded or broken ribs and crumpled long bones (accordina femora) may be present. For reasons that are not apparent, the long bones may be either thick or thin. In types III and IV, multiple fractures from minor physical stress can produce severe deformities. Kyphoscoliosis can impair respiration, cause cor pulmonale, and predispose to pulmonary infections. The appearance of “popcorn-like” deposits of mineral in x-rays of the ends of long bones is an ominous sign. Progressive neurologic symptoms may

result from basilar compression and communicating hydrocephalus. Type V OI is recognized by the presence of dislocated radial heads and hyperplastic callus formation.

In all forms of OI, bone mineral density is decreased. However, the degree of osteopenia may be difficult to evaluate because recurrent fractures limit exercise and thereby diminish bone mass. Surprisingly, fractures appear to heal normally.

Ocular Features The sclerae can be normal, gray, slightly bluish, or bright blue. The color is probably caused by a thinness of the collagen layers of the sclerae that allows the choroid layers to be seen. Blue sclerae, however, are an inherited trait in some families who do not have increased bone fragility.

Dentinogenesis The teeth may be normal, moderately discolored, or grossly abnormal. The enamel generally appears normal, but the teeth may have a characteristic amber, yellowish brown, or translucent bluish gray color because of a deficiency of dentin that is rich in type I collagen. The deciduous teeth are usually smaller than normal, whereas permanent teeth are frequently bell-shaped and restricted at the base. In some patients, the teeth readily fracture and need to be extracted. Similar tooth defects, however, can be inherited without any evidence of OI.

Hearing Loss Hearing loss usually begins during the second decade of life and occurs in more than 50% of individuals over age 30. The loss can be conductive, sensorineural, or mixed, and it varies in severity. The middle ear usually exhibits maldevelopment, deficient ossification, persistence of cartilage in areas that are normally ossified, and abnormal calcium deposits.

Other Features Changes in other connective tissues can include thin skin that scars extensively, joint laxity with permanent dislocations indistinguishable from those of Ehlers-Danlos syndrome (EDS), and occasionally, cardiovascular manifestations such as aortic regurgitation, floppy mitral valves, mitral incompetence, and fragility of large blood vessels. For unknown reasons, some patients develop bouts of a hypermetabolic state with elevated serum thyroxine levels, hyperthermia, and excessive sweating.



Molecular Defects Of the ~1360 unique gene mutations now described in OI, more than 90% are heterozygous mutations in either *COL1A1* or *COL1A2*, the genes coding for the pro α 1 or pro α 2 chain of type I procollagen (Table 427-2).

Most patients with type I OI and blue sclerae have mutations that reduce the synthesis of pro α 1 chains to about one-half. Mutations that reduce the synthesis of pro α 2 chains produce slightly more severe phenotypes and skin defects similar to EDS.

In contrast to the null mutations found in type I OI, most of the severe variants (types II, III, and IV) are caused by mutations that produce structurally abnormal pro α chains that have compromised assembly or abnormal folding of the triple helix. As with collagen mutations in other connective tissue diseases, these structural mutations generally fall into two functional categories. First, the relatively rare mutations in the C-propeptide domain can prevent or seriously impair initial assembly of the procollagen trimers. These misfolded chains are retained in the endoplasmic reticulum (ER) and targeted for degradation by the ER-associated proteasomal pathway. Because these mutations induce an ER stress response, the unfolded protein response (UPR) may have many downstream effects on cells. ER stress is a new concept in the pathophysiology of connective tissue disease and has been best characterized for chondrodysplasias (see below).

The most common type I collagen mutations, however, are single base substitutions that introduce an amino acid with a bulky side chain for one of the glycine residues that appear as every third amino acid in the triple helix. In effect, any of the 338 glycine residues in the helical domain of either the pro α 1 or pro α 2 chain of type I procollagen is a potential site for a disease-producing mutation. These mutations compromise the structural integrity of the triple helix, causing disruption to helix folding, retention of the mutant trimers in the ER, and increased posttranslational hydroxylation and glycosylation of lysines. Collagen-containing helix mutations can form insoluble aggregates in the ER that are degraded by the autophagosome-endosome system, rather than the proteasomes.

A similar sequence of events occurs with less common mutations that produce partial gene deletions, partial gene duplications, and splicing mutations. In addition to their intracellular effects, the structurally abnormal mutant-containing collagen that is secreted by the cell can also have important extracellular effects. For example, the presence of one abnormal pro α chain in a procollagen molecule can interfere with cleavage of the N-propeptide from the protein. The persistence of the N-propeptide on a fraction of the molecules interferes with the self-assembly of normal collagen so that thin and irregular collagen fibrils are formed. Furthermore, if structurally abnormal collagens are incorporated into fibrils, they may have a destabilizing effect and be selectively degraded, or they may alter the interactions of collagen with other connective tissue components, disturbing architecture and stability.

Several generalizations can be made about mutations in type I collagen genes. One is that unrelated patients rarely have the same mutation in the same gene. Glycine substitutions in the N-terminal region of the triple helix tend to produce milder phenotypes, apparently because they have less effect on the zipper-like propagation of the

triple-helical conformation from the C terminus. Rare substitutions of charged amino acids (Asp, Arg) or a branched amino acid (Val) in X- or Y-positions produce lethal phenotypes, apparently because they are located at sites for lateral assembly of the monomers or binding of other components of the matrix.

The search for mutations causing the less common and autosomal recessive forms of OI identified mutations in genes for a series of proteins that are essential for the timely folding of the procollagen monomer: cartilage-associated protein (*CRTAP*), prolyl-3-hydroxylase (*LEPRE1/P3H1*), cyclophilin B (*PPIB*), collagen chaperone-like protein HSP47 (*SERPINH1*), and the procollagen chaperone protein FKBP65 (*FKBP10*). Recently, mutations have been characterized in additional downstream components of the collagen fibrillogenesis pathway: *BMP1*, the gene coding for a metalloproteinase that cleaves the C-propeptide of type I procollagen, and *PLOD2* (LH2, lysyl oxidase 2), which is involved in establishing collagen cross-links. In addition to these mutations that affect the collagen assembly pathway, mutations have been characterized in genes involved in the regulation of bone formation and mineralization such as *SP7* (osterix), *IFITM5*, *WNT1*, and *TMEM38B* (Table 427-2).

Inheritance and Mosaicism in Germline Cells and in Somatic Cells Type I OI is inherited as an autosomal dominant trait. However, some patients with type I OI appear to represent sporadic new mutations or a diagnosis that was missed in earlier generations. Most lethal OI is the result of sporadic mutations that occur in the germ line in one of the parents. Because of the possibility for germline mosaicism for newly generated mutations, there is about a 7% probability that a second child could inherit a severe variant of OI.

Diagnosis OI is usually diagnosed on the basis of clinical criteria. The presence of fractures together with blue sclerae, dentinogenesis imperfecta, or family history of the disease is usually sufficient to make the diagnosis. Other causes of pathologic fractures must be excluded, including battered child syndrome, nutritional deficiencies, malignancies, and other inherited disorders such as chondrodysplasias and hypophosphatasia that can have overlapping presentations. The absence of superficial bruises can be helpful in distinguishing OI from battered child syndrome. X-rays usually reveal a decrease in bone density that can be verified by photon or x-ray absorptiometry. Bone microscopy can be helpful in the diagnosis. The diagnosis, as in other genetic disorders, is now routinely determined using targeted candidate gene sequencing and exome sequencing, but whole-genome sequencing may be used in the future.

TREATMENT OSTEOPENIA AND OSTEODYSPLASIA

Many patients with OI lead productive and successful lives despite severe deformities. Those with mild forms of the disease may need little treatment when fractures decrease after puberty, but women require special attention during pregnancy and after menopause, when fractures again increase. More severely affected children require a comprehensive program of physical therapy and surgical management of fractures and skeletal deformities.

Many fractures are only slightly displaced and have little soft tissue swelling and, therefore, can be treated with minimal support or traction for a week or two followed by a light cast. If fractures are relatively painless, physical therapy can be initiated early. A judicious amount of exercise prevents loss of bone mass secondary to physical inactivity. Some physicians advocate insertion of steel rods into long bones to correct limb deformities; the risk/benefits and cost/benefits of such procedures are difficult to evaluate. Aggressive conventional intervention is usually warranted for pneumonia and cor pulmonale. For severe hearing loss, stapedectomy or replacement of the stapes with a prosthesis may be successful. Moderately to severely affected patients should be evaluated periodically to anticipate possible neurologic problems. About half of children have a substantial increase in growth when given growth hormone. Treatment with bisphosphonates to decrease bone loss has been introduced for moderate to severe forms of OI. Improvements in bone mineral

2510 density are consistently seen in patients. Some clinical trials observed improvements in bone pain and fracture incidence; however, there are still unresolved questions about the best delivery protocols and the risks associated with long-term use in OI patients. For these reasons, the current consensus is that bisphosphonate therapy should be restricted to moderate to severe OI, where the possible benefits outweigh risks. Also, a clinical trial was performed in which patients were treated by intravenous infusion of cells from bone marrow referred to as mesenchymal stem cells, or multipotential stromal cells (MSCs; **see Chap. 90e**). Promising results were obtained, but the trial required a prior bone marrow transplant with marrow from a normal donor who subsequently was used as a source of normal MSCs. As a result, the procedure has not been widely adopted. However, the results raise the possibility that it may be possible in the future to develop effective stem cell therapies for OI.

Counseling and emotional support are important for patients and their parents; lay organizations in some countries provide help in these areas. Prenatal ultrasonography will detect severely affected fetuses at about 16 weeks of pregnancy. Diagnosis is routinely performed on DNA from blood.

EHLERS-DANLOS SYNDROME

EDS is characterized by hyperextensible skin and hypermobile joints, but the category includes rare patients with other distinctive features. Mutations in different types of collagen are found in many patients, but other genes are at fault in rare forms. Contrary to initial expectations, no patients have been found with mutations in the gene for elastin in EDS.

Classification Several types of EDS have been defined, based on the extent to which the skin, joints, and other tissues are involved, mode

of inheritance, and molecular and biochemical analysis (**Table 427-3**). Classical EDS includes a severe form of the disease (type I) and a milder form (type II), both characterized by joint hypermobility and skin that is velvety in texture, hyperextensible, and easily scarred. In hypermobile EDS (type III), joint hypermobility is more prominent than skin changes. In vascular-type EDS (type IV), the skin changes are more prominent than joint changes, and the patients are predisposed to sudden death from rupture of large blood vessels or other hollow organs. EDS type V is similar to EDS type II but is inherited as an X-linked trait. The ocular-scoliotic type of EDS (type VI) is characterized by scoliosis, ocular fragility, and a cone-shaped deformity of the cornea (keratoconus). The arthrochalasic type of EDS (type VIIA and VIIB) is characterized by marked joint hypermobility that is difficult to distinguish from EDS III except by the specific molecular defects in the processing of type I procollagen to collagen. The periodontic-type EDS (type VIII) is distinguished by prominent periodontal changes. EDS types IX, X, and XI were defined on the basis of preliminary biochemical and clinical data. EDS due to tenascin X deficiency has not been assigned a type; it is an autosomal recessive form of the syndrome similar to EDS II. The cardiac valvular form of EDS has similar features to EDS II, but also involves severe changes to the aorta. The progeroid form of EDS displays features of both EDS and progeria. Because of overlapping signs and symptoms, many patients and families with some of the features of EDS cannot be assigned to any of the defined types.

Incidence The overall incidence of EDS is about 1 in 5000 births, with a higher rate for blacks. Classical and hypermobile types of EDS are the most common. Patients with milder forms frequently do not seek medical attention.

Skin Skin changes vary from thin and velvety to skin that is either dramatically hyperextensible (“rubber person” syndrome) or easily

TABLE 427-3 DIFFERENT FORMS OF EHLERS-DANLOS SYNDROME

Type	Typical Features	Inheritance	Gene Defect	Protein Defect
Classic (EDS I—severe and EDS II—mild)	Skin hyperextensibility and fragility, joint hypermobility, tissue fragility manifested by widened atrophic scarring	AD	<i>COL5A1</i> <i>COL5A2</i>	Collagen V
		AD	<i>COL1A1</i>	Proa1 (I) and proa2 (I) chains of procollagen I
		AD, AR	<i>COL1A2</i>	
Hypermobile (EDS III)	Joint hypermobility, moderate skin involvement, absence of tissue fragility	AD	<i>TNXB</i>	Tenascin X
Vascular (EDS IV)	Markedly reduced life span due to spontaneous rupture of internal organs such as arteries and intestines; skin is thin, translucent, and fragile, with extensive bruising; hypermobile minor joints; characteristic facial appearance	AD	<i>COL3A1</i>	Collagen III
X-linked EDS (EDS V)	Similar to classic type	X-linked recessive	Unknown	Unknown
Ocular-scoliotic EDS VI (EDS VIA and EDS VIIB)	Features of classic EDS as well as severe muscular hypotonia after birth, progressive kyphoscoliosis, a Marfanoid habitus, osteopenia, occasionally rupture of the eye globe and great arteries	AR	<i>PLOD1</i>	Deficiency of procollagen-lysine 5-dioxygenase activity (EDS VIA)
			Unknown for EDS VIIB	Unknown for EDS VIIB
Arthrochalasic EDS VII (EDS VIIA and EDS VIIB)	Congenital bilateral hip dislocation, hypermobile joints, moderate skin involvement, osteopenia	AD	<i>COL1A1</i> <i>COL1A2</i>	Mutations that prevent cleavage of the N propeptides
		AR	<i>ADAMTS2</i>	Deficiency of procollagen I N-terminal proteinase
Periodontic EDS VIII	Absorptive periodontitis with premature loss of permanent teeth, fragility of the skin, skin lesions	AD	Unknown	Unknown
EDS due to tenascin X deficiency	Similar to EDS II	AR	<i>TNXB</i>	Tenascin X
EDS, progeroid form		AR?	<i>B4GALT7</i>	Deficiency of galactosyltransferase 7 (defective synthesis of dermatan sulfate proteoglycans)

Abbreviations: AD, autosomal dominant; AR, autosomal recessive.

torn or scarred. Patients with classical EDS develop characteristic “cigarette-paper” scars. In vascular-type EDS, extensive scars and hyperpigmentation develop over bony prominences, and the skin may be so thin that subcutaneous blood vessels are visible. In the periodontotic type of EDS, the skin is more fragile than hyperextensible, and it heals with atrophic, pigmented scars. Easy bruisability occurs in several types of EDS.

Ligament and Joint Changes Laxity and hypermobility of joints vary from mild to unreducible dislocations of hips and other large joints. In mild forms, patients learn to avoid dislocations by limiting physical activity. In more severe forms, surgical repair may be required. Some patients have progressive difficulty with age.

Other Features Mitral valve prolapse and hernias occur, particularly with type I. Pes planus and mild to moderate scoliosis are common. Extreme joint laxity and repeated dislocations may lead to degenerative arthritis. In the ocular-scoliotic type of EDS, the eye may rupture with minimal trauma, and kyphoscoliosis can cause respiratory impairment. Also, sclerae may be blue.



Molecular Defects Subsets of patients with different types of EDS have mutations in the structural genes for collagens (Table 427-3). These include mutations in the *COL1A1* gene in a few patients with moderately severe classical EDS (type I); mutations in *COL1A2* in rare patients with an aortic valvular form of EDS; mutations in two of the three genes (*COL5A1* and *COL5A2*) for type V collagen, a minor collagen found in association with type I collagen, in about half the patients with classical EDS (types I and II); and mutations in the *COL3A1* gene for type III collagen, which is abundant in the aorta in patients with the frequently lethal vascular EDS (type IV).

Some of the type I collagen-related mutations alter processing of the protein or genes for the processing enzymes. Arthrochalcic EDS (type VII) is caused by mutations in the amino acid sequence that make type I procollagen resistant to cleavage by procollagen N-proteinase or by mutations that decrease the activity of the enzyme. The persistence of the N propeptide causes the formation of collagen fibrils that are thin and irregular. Some of the patients have fragile bones and therefore a phenotype that overlaps with OI. The ocular-scoliotic type of EDS (type VI) is caused by homozygous or compound heterozygous mutations in the *PLOD1* gene, which encodes procollagen-lysine 5-dioxygenase (lysyl hydroxylase 1), an enzyme required for formation of stable cross-links in collagen fibers.

Some patients with the hypermobile EDS (type III) and a few with mild EDS (type II) have mutations in the *TNXB* gene, which encodes tenascin X, another minor component of connective tissue that appears to regulate the assembly of collagen fibers. Mutations in proteoglycans have been found in a few patients. The progeroid form of EDS results from autosomal recessive mutations in *B4GALT7*, the gene for β -1,4-galactosyltransferase 7, a key enzyme in the addition of glycosaminoglycan chains to proteoglycans.

Diagnosis The diagnosis is based on clinical criteria and increasingly on DNA sequencing. Correlations between genotype and phenotype can be challenging, but gene or biochemical tests are particularly useful for the diagnosis of vascular type IV EDS with its dire prognosis.

As with other heritable diseases of connective tissue, there is a large degree of variability among members of the same family carrying the same mutation. Some patients have increased fractures and are difficult to distinguish from OI. A few families with heritable aortic aneurysms have mutations in the gene for type III collagen without any evidence of EDS or OI.

TREATMENT EHLERS-DANLOS SYNDROME

Surgical repair and tightening of joint ligaments require careful evaluation of individual patients, as the ligaments frequently do not hold sutures. Patients with easy bruisability should be evaluated for bleeding disorders. Patients with type IV EDS and members of their families should be evaluated at regular intervals for early detection

of aneurysms, but surgical repair may be difficult because of friable tissues. Also, women with type IV EDS should be counseled about the increased risk of uterine rupture, bleeding, and other complications of pregnancy.

CHONDRODYSPLASIAS

(See also Chap. 426e) Chondrodysplasias (CDs), also referred to as skeletal dysplasias, are heritable skeletal disorders that are characterized by dwarfism and abnormal body proportions. The category also includes some individuals with normal stature and body proportions who have features such as ocular changes or cleft palate, which are common in more severe CDs. Many patients develop degenerative joint changes, and mild CD in adults may be difficult to differentiate from primary generalized osteoarthritis. An undefined number of patients have mutations in either the most abundant collagen in cartilage (type II) or the less abundant collagens (types X or XI). Other patients have mutations in genes that code for other components of cartilage or for proteins required for the embryonic development of cartilage, including a common mutation in a gene for a fibroblast growth factor receptor.

Classification Over 200 distinct types and subtypes have been defined based on criteria such as “bringing death” (thanatophoric), causing “twisted” bones (diastrophic), affecting metaphyses (metaphyseal), affecting epiphyses (epiphyseal), affecting spine (spondylo-), and producing histologic changes such as an apparent increase in the fibrous material in the epiphyses (fibrochondrogenesis). Also, a number of eponyms are based on the first or most comprehensive case reports. Severe forms of the diseases produce dwarfism with gross distortions of most cartilaginous structures and of other structures including the eye. Mild forms are more difficult to classify. Among the features are cataracts, degeneration of the vitreous, retinal detachment, high forehead hypoplastic facies, cleft palate, short extremities, and gross distortions of the epiphyses, metaphyses, and joint surfaces. Patients with Stickler’s syndrome (hereditary arthro-ophthalmopathy) have been classified into three types based on a combination of the ocular phenotype and mutated genes.

INCIDENCE

The overall incidence of all forms of CD ranges from 1 per 2500 to 1 per 4000 births. Data on the frequency of individual CDs are incomplete, but the incidence of Stickler’s syndrome is 1 in 10,000. Therefore, the disease is probably among the more common heritable disorders of connective tissue.



Molecular Defects Mutations in the *COL2A1* gene for the type II collagen of cartilage are found in a fraction of patients with both mild and severe CDs. For example, a mutation in the gene substituting a cysteine residue for an arginine was found in three unrelated families with spondyloepiphyseal dysplasia (SED) and precocious generalized osteoarthritis (OA). Mutations in the gene, often glycine substitution mutations with the collagen II triple helix, were also found in some lethal CDs characterized by gross deformities of bones and cartilage, such as those found in spondyloepiphyseal dysplasia congenita, spondyloepimetaphyseal dysplasia congenita, hypochondrogenesis/achondrogenesis type II, and Kniest’s syndrome. The highest incidence of *COL2A1* mutations, however, occurs in patients with the distinctive features of Stickler’s syndrome, which is characterized by skeletal changes, orofacial abnormalities, and auditory abnormalities. Most of the mutations in *COL2A1* are premature stop codons that produce haploinsufficiency. In addition, some patients with Stickler’s syndrome or a closely related syndrome have mutations in two genes specific for type XI collagen, which is an unusual heterotrimer formed from a chains encoded by the gene for type II collagen (*COL2A1*) and two distinctive genes for type XI collagen (*COL11A1* and *COL11A2*). Mutations in the *COL11A1* gene are also found in patients with Marshall’s syndrome, which is similar to classic Stickler’s syndrome, but with more severe hearing loss and dysmorphic features, such as a flat or retracted midface with a flat nasal bridge, short nose, anteverted nostrils, long philtrum, and large-appearing eyes.

CDs are also caused by mutations in the less abundant collagens found in cartilage. For example, patients with Schmid metaphyseal CD have mutations in the gene for type X collagen, a short, network-forming collagen found in the hypertrophic zone of endochondral cartilage. The syndrome is characterized by short stature, coxa vara, flaring metaphyses, and waddling gait. As with other collagen genes, the most common mutations are of two types: nonsense mutations that lead to haploinsufficiency and structural mutations that compromise collagen assembly. In type X collagen, all the structural mutations detected occur in the C-terminal NC1 domain that coordinates the formation of the trimers. This NC1 domain is functionally equivalent to the C-propeptide of the fibrillar collagens. These mutations disturb the structure of the NC1 domain, leading to misfolding and initiation of cellular ER stress via the unfolded protein response (UPR). While the UPR evolved to allow cells to adjust their ER folding capacity to differing protein folding loads, it is deployed by cells when mutant misfolded proteins accumulate in the ER. Activation of the UPR attenuates protein translation and activates mutant protein degradation pathways such as ER-associated degradation. If these strategies do not sufficiently reduce the stress response, cell death may occur. In Schmid metaphyseal CD, mutant misfolded type X collagen induces the UPR, resulting in downstream consequences that contribute to the pathophysiology. This general mechanism may also contribute to pathology in other CDs (and in other connective tissues disorders) where gene mutations lead to protein structural abnormalities.

Some patients have mutations in genes for proteins that interact with collagens. Patients with pseudoachondroplasia or autosomal dominant multiple epiphyseal dysplasia have mutations in the gene for the cartilage oligomeric matrix protein (*COMP*), a protein that interacts with both collagens and proteoglycans in cartilage. However, some families with multiple epiphyseal dysplasia have a defect in one of the three genes for type IX collagen (*COL9A1*, *COL9A2*, and *COL9A3*) or in matrilin-3, another extracellular protein found in cartilage. With misfolding mutations in *COMP* and matrilin-3, the activation of the UPR has been described, providing further evidence that the UPR is a component of pathology of these conditions.

Some CDs are caused by mutations in genes that affect early development of cartilage and related structures. The most common form of short-limbed dwarfism, achondroplasia, is caused by mutations in the gene for a receptor for a fibroblastic growth factor (*FGFR3*). The mutations in the *FGFR3* gene causing achondroplasias are unusual in several respects. The same single-base mutation in the gene that converts glycine to arginine at position 380 in the *FGFR3* gene is present in over 90% of patients. Most patients harbor sporadic new mutations, and therefore, this nucleotide change must be one of the most common recurring mutations in the human genome. The mutation causes unregulated signal transduction through the receptor and inappropriate development of cartilage. Mutations that alter other domains of *FGFR3* have been found in patients with the more severe disorders of hypochondroplasia and thanatophoric dysplasia and in a few families with a variant of craniosynostosis. However, most patients with craniosynostosis appear to have mutations in the related *FGFR2* gene. The similarities between the phenotypes produced by mutations in genes for FGF receptors and mutations in structural proteins of cartilage are probably explained by the observation that the activity of FGFs is regulated in part by binding of FGFs to proteins sequestered in the extracellular matrix. Therefore, the situation parallels the interactions between transforming growth factors (TGFs) and fibrillin in MFS (see below).

Other mutations involve the proteoglycans of cartilage, aggrecan (*AGC1*), and perlecan (*HSPG2*) and the proteoglycan posttranslational sulphation pathway (*DTDST*, *PAPSS2*, and *CHST3*). Mutations in more than 45 other genes have been defined in chondrodysplasias.

Diagnosis The diagnosis of CDs is made on the basis of the physical appearance, slit-lamp eye examinations, x-ray findings, histologic changes, and clinical course. Evaluation of patients by specialists in the field is usually required for a diagnosis. Targeted gene and exome sequencing or more global sequencing strategies are used for molecular diagnosis. Given the wide spectrum of CD phenotypes, these gene

tests are becoming critical diagnostic tools. For Stickler's syndrome, more precise diagnostic criteria have made it possible to identify type I variants with mutations in the *COL2A1* gene with a high degree of accuracy. It has been suggested that the type II variant with mutations in the *COL11A1* gene can be identified on the basis of a "beaded" vitreous phenotype, and the type III variant with mutations in the *COL11A2* gene can be identified on the basis of the characteristic systemic features without the ocular involvement. Prenatal diagnosis based on analysis of DNA obtained from chorionic villus or amniotic fluid is possible.

TREATMENT CHONDRODYSPLASIAS

The treatment is symptomatic and is directed to secondary features such as degenerative arthritis. Many patients require joint replacement surgery and corrective surgery for cleft palate. The eyes should be monitored carefully for the development of cataracts and the need for laser therapy to prevent retinal detachment. In general, patients should be advised to avoid obesity and contact sports. Counseling for the psychological problems of short stature is critical, and support groups have formed in many countries.

MARFAN'S SYNDROME (MFS)

MFS includes features that primarily affect the skeleton, the cardiovascular system, and the eyes. Most patients have mutations in the gene for fibrillin-1.

Classification MFS was initially characterized by a triad of features: (1) skeletal changes that include long, thin extremities, frequently associated with loose joints; (2) reduced vision as the result of dislocations of the lenses (ectopia lentis); and (3) aortic aneurysms. An international panel has developed a series of revised "Ghent criteria" that are useful in classifying patients.

Incidence and Inheritance The incidence of MFS is among the highest of any heritable disorder: about 1 in 3000/5000 births in most racial and ethnic groups. The related syndromes are less common. Mutations are generally inherited as autosomal dominant traits, but about one-fourth of patients have sporadic new mutations.

Skeletal Effects Patients have long limbs and are usually tall compared to other members of the same family. The ratio of the upper segment (top of the head to the top of the pubic ramus) to the lower segment (top of the pubic ramus to the floor) is usually two standard deviations below mean for age, race, and sex. The fingers and hands are long and slender and have a spider-like appearance (arachnodactyly). Many patients have severe chest deformities, including depression (pectus excavatum), protrusion (pectus carinatum), or asymmetry. Scoliosis is frequent and usually accompanied by kyphosis. High-arched palate and high pedal arches or pes planus are common. A few patients have severe joint hypermobility similar to EDS. Computed tomography or magnetic resonance imaging examinations of the lumbar sacral region frequently reveal enlargement of the neural canal, thinning of the pedicles and laminae, widening of the foramina, or anterior meningocele (dural ectasia).

Cardiovascular Features Cardiovascular abnormalities are the major source of morbidity and mortality (**Chap. 301**). Mitral valve prolapse develops early in life and progresses to mitral valve regurgitation of increasing severity in about one-quarter of patients. Dilation of the root of the aorta and the sinuses of Valsalva are characteristic and ominous features of the disease that can develop at any age. The rate of dilation is unpredictable, but it can lead to aortic regurgitation, dissection of the aorta, and rupture. Dilation is probably accelerated by physical and emotional stress, as well as by pregnancy. Patients usually differ from those with familial aortic aneurysms who tend to develop aneurysms in the abdominal aorta. The location of the aneurysms, however, is somewhat variable, and the high incidence of aortic aneurysms in the general population (1 in 100) makes the differential diagnosis difficult unless other features of MFS are clearly present.

Ocular Features Upward displacement of the lens is common. It is usually not progressive but may contribute to the formation of cataracts. The ocular globe is frequently elongated, and most patients are myopic, but with adequate vision. Retinal detachment can occur.

Other Features Striae may occur over the shoulders and buttocks. A number of patients develop spontaneous pneumothorax. Inguinal and incisional hernias are common. Patients are typically thin with little subcutaneous fat, but adults may develop centripetal obesity.



Molecular Defects More than 90% of patients clinically classified as having MFS by the “Ghent criteria” have a mutation in the gene for fibrillin-1 (*FBN1*). Mutations in the same gene are found in a few patients who do not meet the Ghent criteria. Also, a few MFS patients without mutations in the *FBN1* gene have mutations in the gene for TGF- β receptor 2 (*TGFBR2*). In addition, mutations in either *TGFBR2* or *TGFBR1* are found in the related Loeys-Dietz syndrome, which is characterized by aortic aneurysms, cleft palate, and hypertelorism. Mutations in the *FBN2* gene, which is structurally similar to the *FBN1* gene, are found in patients with MFS-like syndrome of congenital contractural arachnodactyly.

FBN1 gene mutations are scattered throughout its 65 coding exons. Most are private mutations, but about 10% are recurrent new mutations that are largely located in CpG sequences known to be “hot spots.” Most severe mutations are located in the central codons (24–32). About one-third of the mutations introduce premature termination codons, and about two-thirds are missense mutations that alter calcium-binding domains in the repetitive epidermal growth factor-like domains of the protein. Rarer mutations alter the processing of the protein. As in many genetic diseases, the severity of the phenotype cannot be predicted from the nature of the mutation.

The discovery that syndromes similar to MFS are caused by mutations in *TGFBR1* and *TGFBR2* refocused attention on structural similarity between fibrillin-1 and TGF- β binding proteins that sequester TGF- β in the extracellular matrix. As a result, some of the manifestations of MFS have been shown to arise from alterations in binding sites that modulate TGF- β bioavailability during development of the skeleton and other tissues. Likewise, *TGFBR1* and *TGFBR2* mutations in Loeys-Dietz syndrome alter TGF- β signaling. In both MFS and Loeys-Dietz syndrome, the pathogenic mechanisms involve increased TGF- β signaling, which contributes to aneurysm formation.

Diagnosis All patients with a suspected diagnosis of MFS should have a slit-lamp examination and an echocardiogram. Also, homocystinuria should be ruled out by amino acid analysis of plasma (Chap. 434e). The diagnosis of MFS according to the international Ghent standards places emphasis on major criteria that include presence of at least four skeletal abnormalities: ectopia lentis; dilation of the ascending aorta with or without dissection; dural ectasia; and a blood relative who meets the same criteria, with or without a DNA diagnosis. A final diagnosis is based on a balanced assessment of the major criteria together with several minor criteria. The absence of ocular changes suggests the Loeys-Dietz syndrome, and the presence of contractures with some of the signs of OI suggests congenital contractural arachnodactyly.

Diagnostic tests based on gene sequencing or detection of protein defects are available. These results are unlikely to alter the treatment or prognosis but are helpful to inform the patients and families and to rapidly exclude the diagnosis in unaffected family members.

TREATMENT MARFAN'S SYNDROME

Propranolol or other β -adrenergic blocking agents are used to lower blood pressure and thereby delay or prevent aortic dilation. Surgical correction of the aorta, aortic valve, and mitral valve has been successful in many patients, but tissues are frequently friable. Patients should be advised that the risks are increased by severe physical exertion, emotional stress, and pregnancy.

The scoliosis tends to be progressive and should be treated by mechanical bracing and physical therapy if $>20^\circ$ or by surgery if it progresses to $>45^\circ$. Dislocated lenses rarely require surgical

removal, but patients should be followed closely for retinal detachment. The finding that MFS pathophysiology involves alterations in TGF- β signaling has raised the possibility of new therapeutic strategies. Attenuation of TGF- β signaling with agents such as angiotensin II receptor blockers (e.g., losartan) was effective in animal studies and has been very promising in small observational studies on MFS patients, significantly reducing progressive aortic enlargement. Based on these results, large randomized clinical trials of angiotensin receptor blockers in MFS are under way.

ELASTIN-RELATED DISEASES

Mutations in the elastin gene (*ELN*) have been found in patients with supravalvular aortic stenosis and skin that hangs in loose and redundant folds (cutis laxa). As indicated in Table 427-3, patients with several forms of EDS have similar changes in skin that were initially thought to reflect changes in elastin.

EPIDERMOLYSIS BULLOSA (EB)

EB has been defined as the category of heritable disorders involving skin that is specifically characterized by blistering as a result of friction. Using this criterion, it was possible to define subtypes by the ultrastructural layer of skin in which the cleavage and blistering occurred. These functional and anatomical criteria made it possible to establish that most patients with a specific subtype have mutations in genes coding for a structural protein, or a cell adherence protein, expressed in the corresponding layer of skin.

Classification and Incidence The four major types of EB are: (1) EB simplex in which cleavage occurs within the epidermis, (2) junctional EB in which cleavage occurs within the lamina lucida, (3) dystrophic EB in which cleavage occurs within the sublamina densa, and (4) Kindler's syndrome with a mixed level of cleavage in different layers. Patients are then separated into major and minor subtypes based on clinical features and analysis of mutations.

The incidence of EB in the United States is about 1 in 50,000.



Molecular Defects The distinctive anatomic locations in skin have made it possible to relate the clinical subtypes of EB to mutations for specific components. In EB simplex, mutations are found primarily in the genes for the major keratins of basal epithelial cells (keratins 5 and 14) and the cell adhesion proteins plectin, $\alpha 6\beta 4$ integrin, plakophilin-1, and desmoplakin. Patients with the related syndrome, epidermolytic ichthyosis, have mutations in keratin 1 and keratin 10. In junctional EB, mutations occur in type XVII collagen, a laminin (laminin-332), and $\alpha 6\beta 4$ integrin. In the severe syndrome of dystrophic EB, mutations are found in the gene that codes for type VII collagen, which forms long loops anchoring the epidermis to the dermis. Patients with more complex features of what is classified as Kindler's syndrome have mutations in kindlin-1, a focal adhesion protein involved in integrin activation.

Diagnosis and Treatment The diagnosis is based on skin that readily breaks and forms blisters from minor trauma. EB simplex is generally milder than junctional EB or dystrophic EB. Dystrophic EB variants usually have large and prominent scars. Precise classification within subtypes usually requires immunofluorescent mapping. DNA diagnostic tests have been developed as research tools but are not widely available. The treatment is symptomatic. Novel therapeutic approaches such as gene therapy, protein replacement therapy, and cell therapy are being explored.

ALPORT'S SYNDROME (AS)

AS is an inherited disorder characterized by hematuria and several associated features. It was not initially considered as a disorder of connective tissue. However, the search for mutations in the genes coding for the collagen found that most patients had mutations in collagen found in basement membranes (type IV). Four forms of AS are now recognized: (1) classic AS, which is inherited as an X-linked disorder with hematuria, sensorineural deafness, and conical deformation of

2514 the anterior surface of the lens (lenticonus); (2) an X-linked form associated with diffuse leiomyomatosis; (3) an autosomal recessive form; and (4) an autosomal dominant form. Both autosomal recessive and dominant forms can cause renal disease without deafness or lenticonus.

Incidence The incidence of AS is about 1 in 10,000 births in the general population and as high as 1 in 5000 in some ethnic groups. About 80% of AS patients have the classical X-linked variant.

Molecular Defects Most patients have mutations in four of the six genes for the chains of type IV collagen (*COL4A3*, *COL4A4*, *COL4A5*, and *COL4A6*). The genes for the proteins are arranged in tandem pairs on different chromosomes in an unusual head-to-head orientation and with overlapping promoters; i.e., the *COL4A1* and *COL4A2* genes are head-to-head on chromosome 13q34, the *COL4A3* and *COL4A4* genes are on chromosome 2q35–37, and the *COL4A5* and *COL4A6* genes are on chromosome Xq22. The X-linked variants are caused by either mutations in the *COL4A5* gene or by partial deletions of both of the adjacent *COL4A4* and *COL4A5* genes. The autosomal recessive variants are caused by mutations in either the *COL4A3* or *COL4A4* gene. The mutations responsible for the autosomal dominant variants are still unknown, but they have been mapped to the same locus as the *COL4A3* and *COL4A4* genes.

Diagnosis The diagnosis of classic AS is based on X-linked inheritance of hematuria, sensorineural deafness, and lenticonus. The lenticonus together with hematuria is pathognomonic of classic AS. The sensorineural deafness is primarily in the high-tone range. It can frequently be detected only by an audiogram and is usually not progressive. Because of the X-linked transmission, women are generally underdiagnosed and are usually less severely affected than men. The hematuria usually progresses to nephritis and may cause renal failure in late adolescence in affected males and at older ages in some women. Renal transplantation is usually successful.

ACKNOWLEDGMENTS

The authors acknowledge the contributions of Helena Kuivaniemi, Gerard Tromp, Leena Ala-Kokko, and Malwina Czarny-Ratajczak to this chapter in previous editions of *Harrison's*. The authors also wish to thank David Silence for his expert advice on the classifications of types of *OI*.

428 Hemochromatosis

Lawrie W. Powell

DEFINITION

Hemochromatosis is a common inherited disorder of iron metabolism in which dysregulation of intestinal iron absorption results in deposition of excessive amounts of iron in parenchymal cells with eventual tissue damage and impaired function in a wide range of organs. The iron-storage pigment in tissues is called *hemosiderin* because it was believed to be derived from the blood. The term *hemosiderosis* is used to describe the presence of stainable iron in tissues, but tissue iron must be quantified to assess body-iron status accurately (see below and [Chap. 126](#)). *Hemochromatosis* refers to a group of genetic diseases that predispose to iron overload, potentially leading to fibrosis and organ failure. Cirrhosis of the liver, diabetes mellitus, arthritis, cardiomyopathy, and hypogonadotropic hypogonadism are the major clinical manifestations.

Although there is debate about definitions, the following terminology is widely accepted.

1. *Hereditary hemochromatosis* is most often caused by a mutant gene, termed *HFE* , which is tightly linked to the HLA-A locus on

TABLE 428-1 CLASSIFICATION OF IRON OVERLOAD STATES

Hereditary Hemochromatosis	
Hemochromatosis, <i>HFE</i> -related (type 1)	
C282Y homozygosity	
C282Y/H63D compound heterozygosity	
Hemochromatosis, non- <i>HFE</i> -related	
Juvenile hemochromatosis (type 2A) (hemojuvelin mutations)	
Juvenile hemochromatosis (type 2B) (hepcidin mutation)	
Mutated transferrin receptor 2, <i>TFR2</i> (type 3)	
Mutated ferroportin 1 gene, <i>SLC11A3</i> (type 4)	
Acquired Iron Overload	
Iron-loading anemias	Chronic liver disease
Thalassemia major	Hepatitis C
Sideroblastic anemia	Alcoholic cirrhosis, especially when advanced
Chronic hemolytic anemias	Nonalcoholic steatohepatitis
Transfusional and parenteral iron overload	Porphyria cutanea tarda
Dietary iron overload	Dysmetabolic iron overload syndrome
	Post-portacaval shunting
Miscellaneous	
Iron overload in sub-Saharan Africa	
Neonatal iron overload	
Aceruloplasminemia	
Congenital atransferrinemia	

chromosome 6p (see “Genetic Basis,” below). Persons who are homozygous for the mutation are at increased risk of iron overload and account for 80–90% of clinical hereditary hemochromatosis in persons of northern European descent. In such subjects, the presence of hepatic fibrosis, cirrhosis, arthropathy, or hepatocellular carcinoma constitutes iron overload–related disease. Rarer forms of non-*HFE* hemochromatosis are caused by mutations in other genes involved in iron metabolism ([Table 428-1](#)). The disease can be recognized during its early stages when iron overload and organ damage are minimal. At this stage, the disease is best referred to as *early hemochromatosis* or *precirrhotic hemochromatosis* .

2. *Secondary iron overload* occurs as a result of an iron-loading anemia, such as thalassemia or sideroblastic anemia, in which erythropoiesis is increased but ineffective. In the acquired iron-loading disorders, massive iron deposits in parenchymal tissues can lead to the same clinical and pathologic features as in hemochromatosis.

PREVALENCE

HFE-associated hemochromatosis mutations are among the most common inherited disease alleles, although the prevalence varies in different ethnic groups. It is most common in populations of northern European extraction in whom approximately 1 in 10 persons are heterozygous carriers and 0.3–0.5% are homozygotes. However, expression of the disease is variable and modified by several factors, especially alcohol consumption and dietary iron intake, blood loss associated with menstruation and pregnancy, and blood donation. Recent population studies indicate that approximately 30% of homozygous men develop iron overload–related disease and about 6% develop hepatic cirrhosis; for women, the figure is closer to 1%. Presumably there are as yet unidentified modifying genes responsible for expression and there is some early evidence to support this. Nearly 70% of untreated patients develop the first symptoms between ages 40 and 60. The disease is rarely evident before age 20, although with family screening (see “Screening for Hemochromatosis,” below) and periodic health examinations, asymptomatic subjects with iron overload can be identified, including young menstruating women.

In contrast to *HFE*-associated hemochromatosis, the non-*HFE*-associated forms of hemochromatosis ([Table 428-1](#)) are rare, but they affect all races and young people (juvenile hemochromatosis).

GENETIC BASIS

A homozygous G to A mutation in the *HFE* gene resulting in a cysteine to tyrosine substitution at position 282 (C282Y) is the most common mutation. It is identified in 85–90% of patients with hereditary hemochromatosis in populations of northern European descent but is found in only 60% of cases from Mediterranean populations (e.g., southern Italy). A second, relatively common *HFE* mutation (H63D) results in a substitution of histidine to aspartic acid at codon 63. Homozygosity for H63D is not associated with clinically significant iron overload. Some compound heterozygotes (e.g., one copy each of C282Y and H63D) have mild to moderately increased body-iron stores but develop clinical disease only in association with cofactors such as heavy alcohol intake or hepatic steatosis. Thus, *HFE*-associated hemochromatosis is inherited as an autosomal recessive trait; heterozygotes have no, or minimal, increase in iron stores. However, this slight increase in hepatic iron can act as a cofactor that may modify the expression of other diseases such as porphyria cutanea tarda (PCT) or nonalcoholic steatohepatitis.

Mutations in other genes involved in iron metabolism are responsible for non-*HFE*-associated hemochromatosis, including juvenile hemochromatosis, which affects persons in the second and third decades of life (Table 428-1). Mutations in the genes encoding hepcidin, transferrin receptor 2 (TfR2), and hemojuvelin (Fig. 428-1) result in clinicopathologic features that are indistinguishable from *HFE*-associated hemochromatosis. However, mutations in ferroportin, responsible for the efflux of iron from enterocytes and most other cell types, result in iron loading of reticuloendothelial cells and macrophages as well as parenchymal cells.

PATHOPHYSIOLOGY

Normally, the body-iron content of 3–4 g is maintained such that intestinal mucosal absorption of iron is equal to iron loss. This amount is approximately 1 mg/d in men and 1.5 mg/d in menstruating women. In hemochromatosis, mucosal absorption is greater than body requirements and amounts to 4 mg/d or more. The progressive accumulation of iron increases plasma iron and saturation of transferrin and results in a progressive increase of plasma ferritin (Fig. 428-2). A liver-derived peptide, hepcidin, represses basolateral iron transport in the intestine and iron release from macrophages and other cells by binding to ferroportin. Hepcidin, in turn, responds to signals in the liver mediated by *HFE*, TfR2, and hemojuvelin (Fig. 428-1). Thus, hepcidin is a crucial molecule in iron metabolism, linking body stores with intestinal iron absorption.

The *HFE* gene encodes a 343-amino-acid protein that is structurally related to MHC class I proteins (*HFE*). The basic defect in *HFE*-associated hemochromatosis is a lack of cell surface expression of *HFE* (due to the C282Y mutation). The normal (wild-type) *HFE* protein forms a complex with β_2 -microglobulin and transferrin receptor 1 (TfR1). The C282Y mutation completely abrogates this interaction. As a result, the mutant *HFE* protein remains trapped intracellularly, reducing TfR1-mediated iron uptake by the intestinal crypt cell. This impaired TfR1-mediated iron uptake leads to upregulation of the divalent metal transporter (DMT1) on the brush border of the villus cells, causing inappropriately increased intestinal iron absorption (Fig. 428-1). In advanced disease, the body may contain 20 g or more of iron that is deposited mainly in parenchymal cells of the liver, pancreas, and heart. Iron may be increased 50- to 100-fold in the liver and pancreas

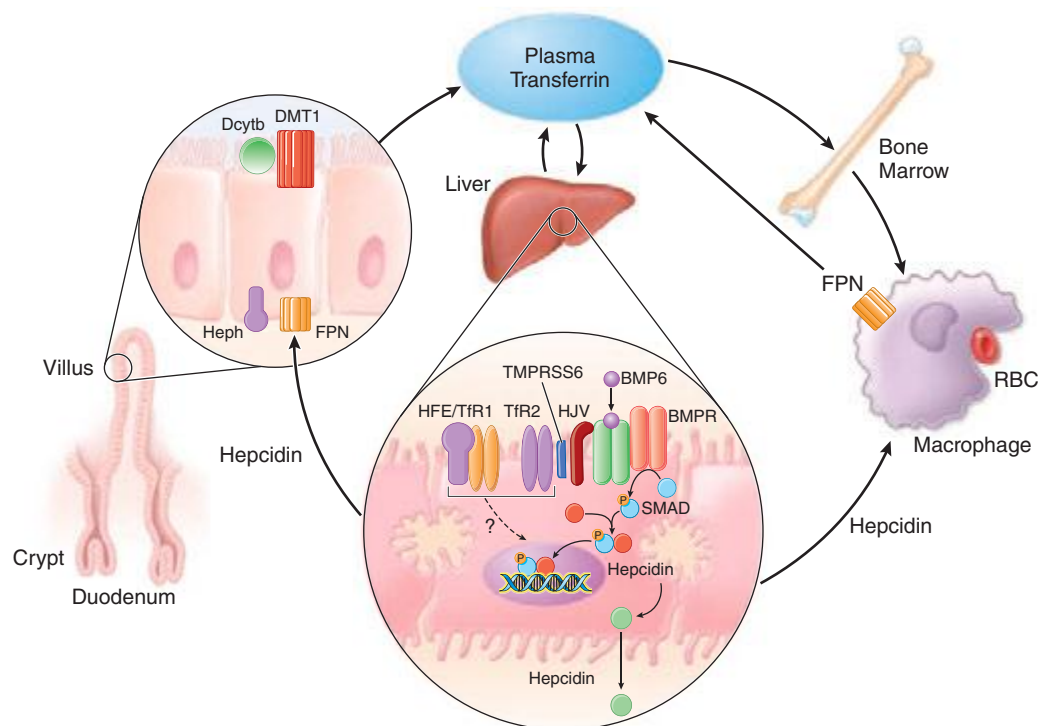


FIGURE 428-1 Pathways of normal iron homeostasis. Dietary inorganic iron traverses the brush border membrane of duodenal enterocytes via the divalent metal-ion transporter 1 (DMT1) after reduction of ferric (Fe^{3+}) iron to the ferrous (Fe^{2+}) state by duodenal cytochrome B (DcytB). Iron then moves from the enterocyte to the circulation via a process requiring the basolateral iron exporter ferroportin (FPN) and the iron oxidase hephaestin (Heph). In the circulation, iron binds to plasma transferrin and is thereby distributed to sites of iron utilization and storage. Much of the diferric transferrin supplies iron to immature erythrocyte cells in the bone marrow for hemoglobin synthesis. At the end of their life, senescent red blood cells (RBCs) are phagocytosed by macrophages, and iron is returned to the circulation after export through ferroportin. The liver-derived peptide hepcidin represses basolateral iron transport in the gut as well as iron released from macrophages and other cells and serves as a central regulator of body-iron traffic. Hepcidin responds to changes in body-iron requirements by signals mediated by diferric transferrin through two mechanisms. One involves *HFE* and TfR2, whereas the other involves hemojuvelin (HJV) and the bone morphogenetic protein (BMP)/SMAD pathway. *TMPRSS6* is a protease that modulates HJV activity. Heme is metabolized by heme oxygenase within the enterocytes, and the released iron then follows the same pathway. Mutations in the genes encoding *HFE*, TfR2, hemojuvelin, and hepcidin all lead to decreased hepcidin release and increased iron absorption, resulting in hemochromatosis (Table 428-1).

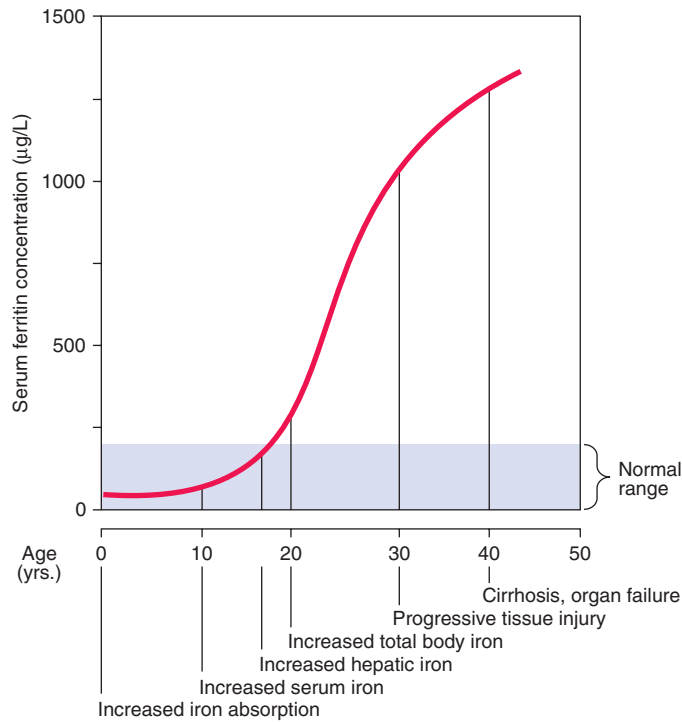


FIGURE 428-2 Sequence of events in genetic hemochromatosis and their correlation with the serum ferritin concentration.

Increased iron absorption is present throughout life. Overt, symptomatic disease usually develops between ages 40 and 60, but latent disease can be detected long before this.

and 5- to 25-fold in the heart. Iron deposition in the pituitary causes hypogonadotropic hypogonadism in both men and women. Tissue injury may result from disruption of iron-laden lysosomes, from lipid peroxidation of subcellular organelles by excess iron, or from stimulation of collagen synthesis by activated stellate cells.

Secondary iron overload with deposition in parenchymal cells occurs in chronic disorders of erythropoiesis, particularly in those due to defects in hemoglobin synthesis or ineffective erythropoiesis such as sideroblastic anemia and thalassemia (Chap. 127). In these disorders, iron absorption is increased. Moreover, these patients require blood transfusions and are frequently treated inappropriately with iron. PCT, a disorder characterized by a defect in porphyrin biosynthesis (Chap. 430), can also be associated with excessive parenchymal iron deposits. The magnitude of the iron load in PCT is usually insufficient to produce tissue damage. However, some patients with PCT also have mutations in the *HFE* gene, and some have associated hepatitis C virus (HCV) infection. Although the relationship between these disorders remains to be clarified, iron overload accentuates the inherited enzyme deficiency in PCT and should be avoided along with other agents (alcohol, estrogens, haloaromatic compounds) that may exacerbate PCT. Another cause of hepatic parenchymal iron overload is hereditary aceruloplasminemia. In this disorder, impairment of iron mobilization due to deficiency of ceruloplasmin (a ferroxidase) causes iron overload in hepatocytes.

Excessive iron ingestion over many years rarely results in hemochromatosis. An important exception has been reported in South Africa among groups who brew fermented beverages in vessels made of iron. Hemochromatosis has been described in apparently normal persons who have taken medicinal iron over many years, but such individuals probably had genetic disorders.

The common denominator in all patients with hemochromatosis is *excessive amounts of iron in parenchymal tissues*. Parenteral administration of iron in the form of blood transfusions or iron preparations results predominantly in reticuloendothelial cell iron overload. This appears to lead to less tissue damage than iron loading of parenchymal cells.

In the liver, parenchymal iron is in the form of ferritin and hemosiderin. In the early stages, these deposits are seen in the periportal

parenchymal cells, especially within lysosomes in the pericanalicular cytoplasm of the hepatocytes. This stage progresses to peribulbar fibrosis and eventually to deposition of iron in bile-duct epithelium, Kupffer cells, and fibrous septa due to activation of stellate cells. In the advanced stage, a macronodular or mixed macro- and micronodular cirrhosis develops. Hepatic fibrosis and cirrhosis correlate significantly with hepatic iron concentration.

At autopsy, the enlarged nodular liver and pancreas are rusty in color. Histologically, iron is increased in many organs, particularly in the liver, heart, and pancreas, and, to a lesser extent, in the endocrine glands. The epidermis of the skin is thin, and melanin is increased in the cells of the basal layer and dermis. Deposits of iron are present around the synovial lining cells of the joints.

CLINICAL MANIFESTATIONS

C282Y homozygotes can be characterized by the stage of progression as follows: (1) a genetic predisposition without abnormalities; (2) iron overload without symptoms; (3) iron overload with symptoms (e.g., arthritis and fatigue); and (4) iron overload with organ damage—in particular, cirrhosis. Thus, many subjects with significant iron overload are asymptomatic. For example, in a study of 672 asymptomatic C282Y homozygous subjects—identified by either family screening or routine health examinations—there was hepatic iron overload (grades 2–4) in 56% and 34.5% of male and female subjects, respectively; hepatic fibrosis (stages 2–4) in 18.4% and 5.4%, respectively; and cirrhosis in 5.6% and 1.9%, respectively.

Initial symptoms are often nonspecific and include lethargy, arthralgia, change in skin color, loss of libido, and features of diabetes mellitus. Hepatomegaly, increased pigmentation, spider angiomas, splenomegaly, arthropathy, ascites, cardiac arrhythmias, congestive heart failure, loss of body hair, testicular atrophy, and jaundice are prominent in advanced disease.

The *liver* is usually the first organ to be affected, and hepatomegaly is present in more than 95% of symptomatic patients. Hepatic enlargement may exist in the absence of symptoms or of abnormal liver-function tests. Manifestations of portal hypertension and esophageal varices occur less commonly than in cirrhosis from other causes. Hepatocellular carcinoma develops in about 30% of patients with cirrhosis, and it is the most common cause of death in treated patients—hence the importance of early diagnosis and therapy. The incidence increases with age, it is more common in men, and it occurs almost exclusively in cirrhotic patients.

Excessive skin pigmentation is present in patients with advanced disease. The characteristic metallic or slate-gray hue is sometimes referred to as *bronzing* and results from increased melanin and iron in the dermis. Pigmentation usually is diffuse and generalized, but it may be more pronounced on the face, neck, extensor aspects of the lower forearms, dorsa of the hands, lower legs, and genital regions, as well as in scars.

Diabetes mellitus occurs in about 65% of patients with advanced disease and is more likely to develop in those with a family history of diabetes, suggesting that direct damage to the pancreatic islets by iron deposition occurs in combination with other risk factors. The management is similar to that of other forms of diabetes, although insulin resistance is more common in association with hemochromatosis. Late complications are the same as seen in other causes of diabetes mellitus.

Arthropathy develops in 25–50% of symptomatic patients. It usually occurs after age 50 but may occur as a first manifestation or long after therapy. The joints of the hands, especially the second and third metacarpophalangeal joints, are usually the first joints involved, a feature that helps to distinguish the chondrocalcinosis associated with hemochromatosis from the idiopathic form (Chap. 395). A progressive polyarthritides involving wrists, hips, ankles, and knees may also ensue. Acute brief attacks of synovitis may be associated with deposition of calcium pyrophosphate (chondrocalcinosis or pseudogout), mainly in the knees. Radiologic manifestations include cystic changes of the subchondral bones, loss of articular cartilage with narrowing of the joint space, diffuse demineralization, hypertrophic bone proliferation, and calcification of the synovium. The arthropathy tends to progress

despite removal of iron by phlebotomy. Although the relation of these abnormalities to iron metabolism is not known, the fact that similar changes occur in other forms of iron overload suggests that iron is directly involved.

Cardiac involvement is the presenting manifestation in about 15% of symptomatic patients. The most common manifestation is congestive heart failure, which occurs in about 10% of young adults with the disease, especially those with juvenile hemochromatosis. Symptoms of congestive heart failure may develop suddenly, with rapid progression to death if untreated. The heart is diffusely enlarged; this may be misdiagnosed as idiopathic cardiomyopathy if other overt manifestations are absent. Cardiac arrhythmias include premature supraventricular beats, paroxysmal tachyarrhythmias, atrial flutter, atrial fibrillation, and varying degrees of atrioventricular block.

Hypogonadism occurs in both sexes and may antedate other clinical features. Manifestations include loss of libido, impotence, amenorrhea, testicular atrophy, gynecomastia, and sparse body hair. These changes are primarily the result of decreased production of gonadotropins due to impairment of hypothalamic-pituitary function by iron deposition. Adrenal insufficiency, hypothyroidism, and hypoparathyroidism are rare manifestations.

DIAGNOSIS

The association of (1) hepatomegaly, (2) skin pigmentation, (3) diabetes mellitus, (4) heart disease, (5) arthritis, and (6) hypogonadism should suggest the diagnosis. However, as stated above, significant iron overload may exist with none or only some of these manifestations. Therefore, a high index of suspicion is needed to make the diagnosis early. Treatment before permanent organ damage occurs can reverse the iron toxicity and restore life expectancy to normal.

The history should be particularly detailed in regard to disease in other family members; alcohol ingestion; iron intake; and ingestion of large doses of ascorbic acid, which promotes iron absorption (Chap. 96e). Appropriate tests should be performed to exclude iron deposition due to hematologic disease. The presence of liver, pancreatic, cardiac, and joint disease should be confirmed by physical examination, radiography, and standard function tests of these organs.

The degree of increase in total body iron stores can be assessed by (1) measurement of serum iron and the percent saturation of transferrin (or the unsaturated iron-binding capacity), (2) measurement of serum ferritin concentration, (3) liver biopsy with measurement of the iron concentration and calculation of the hepatic iron index (Table 428-2), and (4) magnetic resonance imaging (MRI) of the liver. In addition, a retrospective assessment of body-iron storage is also provided by performing weekly phlebotomy and calculating the amount of iron removed before iron stores are exhausted (1 mL blood = approximately 0.5 mg iron).

Each of these methods for assessing iron stores has advantages and limitations. The serum iron level and percent saturation of transferrin are elevated early in the course, but their specificity is reduced by significant false-positive and false-negative rates. For example, serum iron concentration may be increased in patients with alcoholic liver

disease without iron overload; in this situation, however, the hepatic iron index is usually not increased as in hemochromatosis (Table 428-1). In otherwise healthy persons, a fasting serum transferrin saturation greater than 45% is abnormal and suggests homozygosity for hemochromatosis.

The serum ferritin concentration is usually a good index of body-iron stores, whether decreased or increased. In fact, an increase of 1 µg/L in serum ferritin level reflects an increase of about 5 mg in body stores. In most untreated patients with hemochromatosis, the serum ferritin level is significantly increased (Fig. 428-2 and Table 428-1), and a serum ferritin level >1000 µg/L is the strongest predictor of disease expression among individuals homozygous for the C282Y mutation. However, in patients with inflammation and hepatocellular necrosis, serum ferritin levels may be elevated out of proportion to body-iron stores due to increased release from tissues. Therefore, a repeat determination of serum ferritin should be carried out after acute hepatocellular damage has subsided (e.g., in alcoholic liver disease). Ordinarily, the combined measurements of the percent transferrin saturation and serum ferritin level provide a simple and reliable screening test for hemochromatosis, including the precirrhotic phase of the disease. If either of these tests is abnormal, genetic testing for hemochromatosis should be performed (Fig. 428-3).

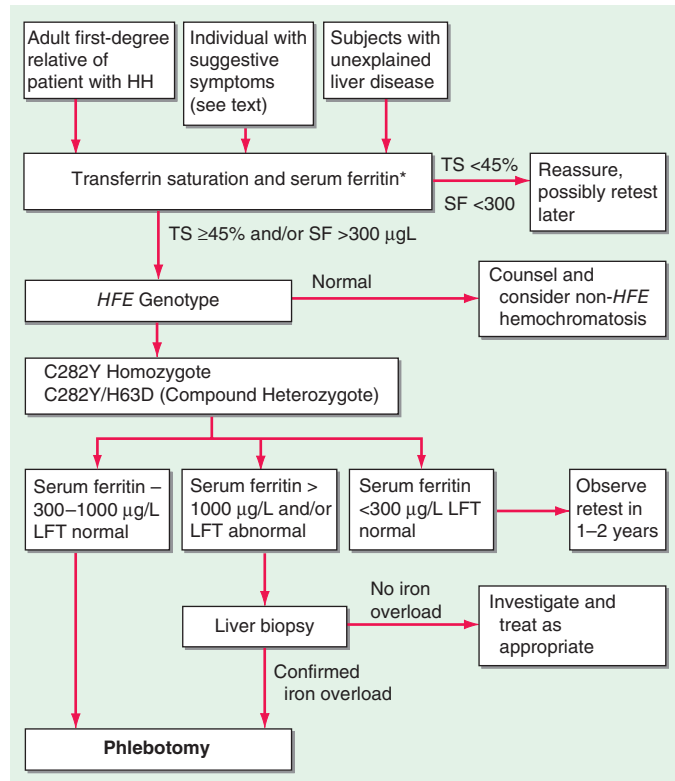
The role of liver biopsy in the diagnosis and management of hemochromatosis has been reassessed as a result of the widespread availability of genetic testing for the C282Y mutation. The absence of severe fibrosis can be accurately predicted in most patients using clinical and biochemical variables. Thus, there is virtually no risk of severe fibrosis in a C282Y homozygous subject with (1) serum ferritin level less than 1000 µg/L, (2) normal serum alanine aminotransferase values, (3) no hepatomegaly, and (4) no excess alcohol intake. However, it should be emphasized that liver biopsy is the only reliable method for establishing or excluding the presence of hepatic cirrhosis, which is the critical factor determining prognosis and the risk of developing hepatocellular carcinoma. Biopsy also permits histochemical estimation of tissue iron and measurement of hepatic iron concentration. Increased density of the liver due to iron deposition can be demonstrated by computed tomography (CT) or MRI, and with improved technology, MRI has become more accurate in determining hepatic iron concentration.

SCREENING FOR HEMOCHROMATOSIS

When the diagnosis of hemochromatosis is established, it is important to counsel and screen other family members (Chap. 84). Asymptomatic and symptomatic family members with the disease usually have an increased saturation of transferrin and an increased serum ferritin concentration. These changes occur even before the iron stores are greatly increased (Fig. 428-2). All adult first-degree relatives of patients with hemochromatosis should be tested for the C282Y and H63D mutations and counseled appropriately (Fig. 428-3). In affected individuals, it is important to confirm or exclude the presence of cirrhosis and begin therapy as early as possible. For children of an identified proband, testing for *HFE* of the other parent is helpful because if

TABLE 428-2 REPRESENTATIVE IRON VALUES IN NORMAL SUBJECTS, PATIENTS WITH HEMOCHROMATOSIS, AND PATIENTS WITH ALCOHOLIC LIVER DISEASE

Determination	Normal	Symptomatic Hemochromatosis	Homozygotes with Early, Asymptomatic Hemochromatosis	Heterozygotes	Alcoholic Liver Disease
Plasma iron, µmol/L (µg/dL)	9–27 (50–150)	32–54 (180–300)	Usually elevated	Elevated or normal	Often elevated
Total iron-binding capacity, µmol/L (µg/dL)	45–66 (250–370)	36–54 (200–300)	36–54 (200–300)	Elevated or normal	45–66 (250–370)
Transferrin saturation, %	22–45	50–100	50–100	Normal or elevated	27–60
Serum ferritin, µg/L		1000–6000	200–500	Usually <500	10–500
Men	20–250				
Women	15–150				
Liver iron, µg/g dry wt	300–1400	6000–18,000	2000–4000	300–3000	300–2000
Hepatic iron index	<1.0	>2	1.5–2	<2	<2



*For convenience both genotype and phenotype (iron tests) can be performed together at a single visit in first-degree relatives.

FIGURE 428-3 Algorithm for screening for *HFE*-associated hemochromatosis. HH, hereditary hemochromatosis, homozygous subject (C282Y +/+); LFT, liver function tests; SF, serum ferritin concentration; TS, transferrin saturation.

normal, the child is merely an obligate heterozygote and at no risk. Otherwise, for practical purposes, children need not be checked before they are 18 years old.

The role of population screening for hemochromatosis is controversial. Recent studies indicate that it is highly effective for primary care physicians to screen subjects using transferrin saturation and serum ferritin levels. Such screening also detects iron deficiency. Genetic screening of the normal population is feasible but is probably not cost effective.

TREATMENT HEMOCHROMATOSIS

The therapy of hemochromatosis involves removal of the excess body iron and supportive treatment of damaged organs. Iron removal is best accomplished by weekly or twice-weekly phlebotomy of 500 mL. Although there is an initial modest decline in the volume of packed red blood cells to about 35 mL/dL, the level stabilizes after several weeks. The plasma transferrin saturation remains increased until the available iron stores are depleted. In contrast, the plasma ferritin concentration falls progressively, reflecting the gradual decrease in body-iron stores. One 500-mL unit of blood contains 200–250 mg of iron, and up to 25 g of iron or more may have to be removed. Therefore, in patients with advanced disease, weekly phlebotomy may be required for 1–2 years, and it should be continued until the serum ferritin level is <50 µg/L. Thereafter, phlebotomies are performed at appropriate intervals to maintain ferritin levels between 50 and 100 µg/L. Usually one phlebotomy every 3 months will suffice.

Chelating agents such as deferoxamine, when given parenterally, remove 10–20 mg of iron per day, which is much less than that mobilized by once-weekly phlebotomy. Phlebotomy is also less expensive, more convenient, and safer for most patients. However,

chelating agents are indicated when anemia or hypoproteinemia is severe enough to preclude phlebotomy. Subcutaneous infusion of deferoxamine using a portable pump is the most effective means of its administration.

An effective oral iron chelating agent, deferasirox (Exjade), has recently become available but is still in clinical trials. This agent is effective in thalassemia and secondary iron overload, but its role in primary iron overload has yet to be established.

Alcohol consumption should be severely curtailed or eliminated because it increases the risk of cirrhosis in hereditary hemochromatosis nearly tenfold. Dietary adjustments are unnecessary, although vitamin C and iron supplements should be avoided. The management of hepatic failure, cardiac failure, and diabetes mellitus is similar to conventional therapy for these conditions. Loss of libido and change in secondary sex characteristics are managed with testosterone replacement or gonadotropin therapy (Chap. 411).

End-stage liver disease may be an indication for liver transplantation, although results are improved if the excess iron can be removed beforehand. The available evidence indicates that the fundamental metabolic abnormality in hemochromatosis is reversed by successful liver transplantation.

PROGNOSIS

The principal causes of death are cardiac failure, hepatocellular failure or portal hypertension, and hepatocellular carcinoma.

Life expectancy is improved by removal of the excessive stores of iron and maintenance of these stores at near-normal levels. The 5-year survival rate with therapy increases from 33 to 89%. With repeated phlebotomy, the liver decreases in size, liver function improves, pigmentation of skin decreases, and cardiac failure may be reversed. Diabetes improves in about 40% of patients, but removal of excess iron has little effect on hypogonadism or arthropathy. Hepatic fibrosis may decrease, but established cirrhosis is irreversible. Hepatocellular carcinoma occurs as a late sequela in patients who are cirrhotic at presentation. The apparent increase in its incidence in treated patients is probably related to their increased life span. Hepatocellular carcinoma rarely develops if the disease is treated in the precirrhotic stage. Indeed, the life expectancy of homozygotes treated before the development of cirrhosis is normal.

The importance of family screening and early diagnosis and treatment cannot be overemphasized. Asymptomatic individuals detected by family studies should have phlebotomy therapy if iron stores are moderately to severely increased. Assessment of iron stores at appropriate intervals is also important. With this management approach, most manifestations of the disease can be prevented.

ROLE OF *HFE* MUTATIONS IN OTHER LIVER DISEASES

There is considerable interest in the role of *HFE* mutations and hepatic iron in several other liver diseases. Several studies have shown an increased prevalence of *HFE* mutations in PCT patients. Iron accentuates the inherited enzyme deficiency in PCT and clinical manifestations of PCT. The situation in nonalcoholic steatohepatitis (NASH) is less clear, but some studies have shown an increased prevalence of *HFE* mutations in NASH patients. The role of phlebotomy therapy, however, is unproven. In chronic HCV infection, *HFE* mutations are not more common, but some subjects have increased hepatic iron. Before initiating antiviral therapy in these patients, it is reasonable to perform phlebotomy therapy to remove excess iron stores, because this reduces liver enzyme levels.

HFE mutations are not increased in frequency in alcoholic liver disease. Hemochromatosis in a heavy drinker can be distinguished from alcoholic liver disease by the presence of the C282Y mutation.

End-stage liver disease may also be associated with iron overload of the degree seen in hemochromatosis. The mechanism is uncertain, although studies have shown that alcohol suppresses hepatic hepcidin secretion. Hemolysis also plays a role. *HFE* mutations are uncommon.

A recent large population study has suggested that subjects homozygous for C282Y are at increased risk of breast and colorectal cancer.

GLOBAL CONSIDERATIONS



The *HFE* mutation is of northern European origin (Celtic or Nordic) with a heterozygous carrier rate of approximately 1 in 10 (1 in 8 in Ireland). Thus, *HFE*-associated hemochromatosis is quite rare in non-European populations, e.g., Asia. However, non-*HFE*-associated hemochromatosis resulting from mutations in other genes involved in iron metabolism (Fig. 428-1) is ubiquitous and should be considered when one encounters iron overload.

429 Wilson's Disease

George J. Brewer

Wilson's disease is an autosomal recessive disorder caused by mutations in the *ATP7B* gene, which encodes a membrane-bound, copper-transporting ATPase. Clinical manifestations are caused by copper toxicity and primarily involve the liver and the brain. Because effective treatment is available, it is important to make this diagnosis early.

The frequency of Wilson's disease in most populations is about 1 in 30,000–40,000, and the frequency of carriers of *ATP7B* mutations is ~1%. Siblings of a diagnosed patient have a 1 in 4 risk of Wilson's disease, whereas children of an affected patient have about a 1 in 200 risk. Because a large number of inactivating mutations have been reported in the *ATP7B* gene, mutation screening for diagnosis is not routine, although this approach may be practical in the future. DNA haplotype analysis can be used to genotype siblings of an affected patient. A rare multisystem disorder of copper metabolism with features of both Menkes and Wilson's diseases has been reported. It is termed the MEDNIK syndrome (mental retardation, enteropathy, deafness, neuropathy, ichthyosis, keratoderma) and is caused by mutations in the *APIS1* gene, which encodes an adaptor protein necessary for intracellular trafficking of copper pump proteins ATP7A (Menkes disease) and ATP7B (Wilson's disease).

PATHOGENESIS

ATP7B protein deficiency impairs biliary copper excretion, resulting in positive copper balance, hepatic copper accumulation, and copper toxicity from oxidant damage. Excess hepatic copper is initially bound to metallothionein; liver damage begins as this storage capacity is exceeded, sometimes by 3 years of age. Defective copper incorporation into apoceruloplasmin leads to excess catabolism and low blood levels of ceruloplasmin. Serum copper levels are usually lower than normal because of low blood levels of ceruloplasmin, which normally binds >90% of serum copper. As the disease progresses, nonceruloplasmin serum copper ("free" copper) levels increase, resulting in copper buildup in other parts of the body (e.g., in the brain, with consequent neurologic and psychiatric disease).

CLINICAL PRESENTATION

Hepatic Features Wilson's disease may present as hepatitis, cirrhosis, or hepatic decompensation. Patients typically present in the mid- to late teenage years in Western countries, although the age of presentation is quite broad and extends into the fifth decade of life. An episode of hepatitis may occur—with elevated serum aminotransferase levels, with or without jaundice—and then spontaneously regress. Hepatitis often recurs, and most of these patients eventually develop cirrhosis. Hepatic decompensation is associated with elevated serum bilirubin, reduced serum albumin and coagulation factors, ascites, peripheral edema, and hepatic encephalopathy. In severe hepatic failure, hemolytic anemia may develop because large amounts of copper derived from hepatocellular necrosis are released into the bloodstream. The association of hemolysis and liver disease makes Wilson's disease a likely diagnosis.

Neurologic Features The neurologic manifestations of Wilson's disease typically occur in patients in their early twenties, although the age of onset extends into the sixth decade of life. MRI and CT scans reveal damage in the basal ganglia and occasionally in the pons, medulla, thalamus, cerebellum, and subcortical areas. The three main movement disorders include dystonia, incoordination, and tremor. Dysarthria and dysphagia are common. In some patients, the clinical picture closely resembles that of Parkinson's disease. Dystonia can involve any part of the body and eventually leads to grotesque positions of the limbs, neck, and trunk. Autonomic disturbances may include orthostatic hypotension and sweating abnormalities as well as bowel, bladder, and sexual dysfunction. Memory loss, migraine-type headaches, and seizures may occur. Patients have difficulty focusing on tasks, but cognition usually is not grossly impaired. Sensory abnormalities and muscular weakness are not features of the disease.

Psychiatric Features Half of patients with neurologic disease have a history of behavioral disturbances with onset in the 5 years before diagnosis. The features are diverse and may include loss of emotional control (temper tantrums, crying bouts), depression, hyperactivity, or loss of sexual inhibition.

Other Manifestations Some female patients have repeated spontaneous abortions, and most become amenorrheic prior to diagnosis. Cholelithiasis and nephrolithiasis occur with increased frequency. Some patients have osteoarthritis, particularly of the knee. Microscopic hematuria is common, and levels of urinary excretion of phosphates, amino acids, glucose, or urates may increase; however, a full-blown Fanconi syndrome is rare. Sunflower cataracts and Kayser-Fleischer rings (copper deposits in the outer rim of the cornea) may be seen. Electrocardiographic and other cardiac abnormalities have been reported but are not common.

DIAGNOSIS

Diagnostic tests for Wilson's disease are listed in [Table 429-1](#). Serum ceruloplasmin levels should not be used for definitive diagnosis, because they are normal in up to 10% of affected patients and are reduced in 20% of carriers. Kayser-Fleischer rings ([Fig. 429-1](#)) can be definitively diagnosed only by an ophthalmologist using a slit lamp. They are present in >99% of patients with neurologic/psychiatric forms of the disease and have been described very rarely in the absence of Wilson's disease. Kayser-Fleischer rings are present in only ~30–50% of patients diagnosed in the hepatic or presymptomatic state; thus, the absence of rings does not exclude the diagnosis.

Urine copper measurement is an important diagnostic tool, but urine must be collected carefully to avoid contamination. Symptomatic patients invariably have urine copper levels >1.6 μmol (>100 μg) per 24 h. Heterozygotes have values <1.3 μmol (<80 μg) per 24 h. About half of presymptomatic patients who are ultimately affected have diagnostically elevated urine copper values, but the other half have levels that are in an intermediate range between 0.9 and 1.6 μmol (60–100 μg) per 24 h. Because heterozygotes may have values up to 1.3 μmol (80 μg) per 24 h, patients in this range may require a liver biopsy for definitive diagnosis.

The gold standard for diagnosis remains liver biopsy with quantitative copper assays. Affected patients have values >3.1 $\mu\text{mol/g}$ (>200 $\mu\text{g/g}$ [dry weight] of liver). Copper stains are not reliable. False-positive results can occur with long-standing obstructive liver disease, which can elevate hepatic and urine copper concentrations and rarely causes Kayser-Fleischer rings.

TREATMENT WILSON'S DISEASE

Recommended anticopper treatments are listed in [Table 429-2](#). Penicillamine was previously the primary anticopper treatment but now plays only a minor role because of its toxicity and because it often worsens existing neurologic disease if used as initial therapy. If penicillamine is given, it should always be accompanied by pyridoxine (25 mg/d). Trientine is a less toxic chelator and is supplanting penicillamine when a chelator is indicated.

TABLE 429-1 USEFUL TESTS FOR WILSON'S DISEASE

Test	Usefulness ^a	Normal Value	Heterozygous Carriers	Wilson's Disease
Serum ceruloplasmin	+	180–350 mg/L (18–35 mg/dL)	Low in 20%	Low in 90%
Kayser- Fleischer rings	++	Absent	Absent	Present in >99% if neurologic or psychiatric symptoms are present Present in 30–50% in hepatic presentation and presymptomatic state
Urine copper (24-h)	+++	0.3–0.8 μmol (20–50 μg)	Normal to 1.3 μmol (80 μg)	>1.6 μmol (>100 μg) in symptomatic patients; 0.9 to >1.6 μmol (60 to >100 μg) in presymptomatic patients
Liver copper	++++	0.3–0.8 μmol/g (20–50 μg/g of tissue)	Normal to 2.0 μmol (125 μg)	>3.1 μmol (>200 μg) (Obstructive liver disease can cause false-positive results.)
Haplotype analysis	++++ (siblings only)	0 matches	1 match	2 matches

^aUsefulness range: + (somewhat useful) to ++++ (very useful).

For patients with hepatitis or cirrhosis but without evidence of hepatic decompensation or neurologic/psychiatric symptoms, zinc is the therapy of choice although some experts advocate therapy with trientine. Zinc has proven efficacy in Wilson's disease and is essentially nontoxic. It produces a negative copper balance by blocking intestinal absorption of copper, and it induces hepatic metallothionein synthesis, thereby sequestering additional toxic copper. All presymptomatic patients should be treated prophylactically because the disease is close to 100% penetrant.

The first step in evaluating patients presenting with hepatic decompensation is to establish disease severity, which can be estimated with the Nazer prognostic index (Table 429-3). Patients with scores <7 can usually be managed with medical therapy. Patients with scores >9 should be considered immediately for liver transplantation. For patients with scores between 7 and 9, clinical judgment is required in deciding whether to recommend transplantation or medical therapy. A combination of trientine and zinc has been used to treat patients with Nazer scores as high as 9, but such patients should be watched carefully for indications of hepatic deterioration, which mandates transplantation.

For initial medical treatment of patients with hepatic decompensation, the recommended regimen is a chelator (preferably trientine) plus zinc (Table 429-2). Zinc should not, however, be ingested simultaneously with trientine, which chelates zinc and forms therapeutically ineffective complexes. Administration of the two drugs should be separated by at least 1 h.

For initial neurologic therapy, tetrathiomolybdate is emerging as the drug of choice because of its rapid control of free copper, preservation of neurologic function, and low toxicity. Penicillamine and trientine should be avoided because both have a high risk of worsening the neurologic condition. Until tetrathiomolybdate is commercially available, zinc therapy is recommended. Although

it is relatively slow-acting, zinc itself does not exacerbate neurologic abnormalities. Although hepatic transplantation may alleviate neurologic symptoms, it does so only by copper removal, which can be done more safely and inexpensively with anticopper drugs. Pregnant patients should be treated with zinc or trientine throughout pregnancy but without tight copper control because copper deficiency can be teratogenic.

Anticopper therapy must be lifelong. With treatment, liver function usually recovers after about a year although residual liver damage is usually present. Neurologic and psychiatric symptoms usually improve after 6–24 months of treatment.

MONITORING ANTICOPPER THERAPY

When trientine or penicillamine is first used, it is necessary to monitor for drug toxicity, particularly bone marrow suppression and proteinuria. Complete blood counts, standard biochemical profiles, and a urinalysis should be performed at weekly intervals for 1 month, then at twice-weekly intervals for 2 or 3 months, then at monthly intervals for 3 or 4 months, and at 4- to 6-month intervals thereafter.

The anticopper effects of trientine and penicillamine can be monitored by following 24-h “free” serum copper levels. Changes in urine copper levels are more difficult to interpret because excretion reflects the effect of the drug as well as body loading with copper. Free serum copper is calculated by subtracting the ceruloplasmin copper from

TABLE 429-2 RECOMMENDED ANTICOPPER DRUGS FOR WILSON'S DISEASE

Disease Status	First Choice	Second Choice
Initial hepatic		
Hepatitis or cirrhosis without decompensation	Zinc ^a	Trientine
Hepatic decompensation		
Mild	Trientine ^b and zinc	Penicillamine ^b and zinc
Moderate	Trientine and zinc	Hepatic transplantation
Severe	Hepatic transplantation	Trientine and zinc
Initial neurologic/psychiatric	Tetrathiomolybdate ^c and zinc	Zinc
Maintenance	Zinc	Trientine
Presymptomatic	Zinc	Trientine
Pediatric	Zinc	Trientine
Pregnant	Zinc	Trientine

^aZinc acetate is supplied as Galzin, manufactured by Gate Pharmaceutical. The recommended adult dose for all the above indications is 50 mg of elemental zinc three times daily, with each dose separated by at least 1 h from consumption of food and beverages other than water as well as from trientine or penicillamine doses. ^bTrientine is supplied as Syprine and penicillamine as Cuprimine, both manufactured by Merck. The recommended adult dosage for both drugs is 500 mg twice daily, with each dose at least 0.5 h before or 2 h after meals and separated by at least 1 h from zinc administration. ^cTetrathiomolybdate is being studied in clinical trials.



FIGURE 429-1 A Kayser-Fleischer ring. Although in this case, the brownish ring rimming the cornea is clearly visible to the naked eye, confirmation is usually made by slit-lamp examination.

TABLE 429-3 PROGNOSTIC INDEX OF NAZER

Laboratory Measurement	Normal Value	Score (in Points)				
		0	1	2	3	4
Serum bilirubin ^a	0.2–1.2 mg/dL	<5.8	5.8–8.8	8.8–11.7	11.7–17.5	>17.5
Serum aspartate aminotransferase	10–35 IU/L	<100	100–150	151–200	201–300	>300
Prolongation of prothrombin time (sec)	—	<4	4–8	9–12	13–20	>20

^aIf hemolysis is present, serum bilirubin cannot be used as a measure of liver function until the hemolysis subsides.

Source: Modified from H Nazer et al: Gut 27:1377, 1986; with permission from BMJ Publishing Group.

the total serum copper. Each 10 mg/L (1 mg/dL) of ceruloplasmin contributes 0.5 $\mu\text{mol/L}$ (3 $\mu\text{g/dL}$) of serum copper. The normal serum free copper value is 1.6–2.4 $\mu\text{mol/L}$ (10–15 $\mu\text{g/dL}$); the level is often as high as 7.9 $\mu\text{mol/L}$ (50 $\mu\text{g/dL}$) in untreated Wilson's disease. With treatment, the serum free copper should be <3.9 $\mu\text{mol/L}$ (<25 $\mu\text{g/dL}$).

Zinc treatment does not require monitoring of blood or urine for toxicity. Its only significant side effect is gastric burning or nausea in ~10% of patients, usually with the first morning dose. This effect can be mitigated if the first dose is taken an hour after breakfast or if zinc is taken with a small amount of protein. Because zinc mainly affects stool copper, 24-h urine copper can be used to reflect body loading. The typical value in untreated symptomatic patients is >3.1 μmol (>200 μg) per 24 h. This level should decrease during the first 1–2 years of therapy to <2.0 μmol (<125 μg) per 24 h. A normal value (0.3–0.8 μmol [20–50 μg]) is rarely reached during the first decade of therapy and should raise concern about overtreatment (copper deficiency), the first sign of which is anemia and/or leukopenia.

GLOBAL CONSIDERATIONS



The age of onset of clinical disease may be considerably younger in India and the Far East; in these regions, onset often occurs in children at only 5 or 6 years of age. The incidence of the disease may be increased in certain populations as a result of founder effects. For example, in Sardinia, the incidence may be 1 in 3000. In countries where penicillamine, trientine, and zinc acetate (as Galzin) are not available or are unaffordable, zinc salts such as gluconate or sulfate provide an alternative treatment option.

430 The Porphyrrias

Robert J. Desnick, Manisha Balwani

The porphyrias are metabolic disorders, each resulting from the deficiency of a specific enzyme in the heme biosynthetic pathway (Fig. 430-1 and Table 430-1). These enzyme deficiencies are inherited as autosomal dominant, autosomal recessive, or X-linked traits, with the exception of porphyria cutanea tarda (PCT), which usually is sporadic (Table 430-1). The porphyrias are classified as either *hepatic* or *erythropoietic*, depending on the primary site of overproduction and accumulation of their respective porphyrin precursors or porphyrins (Tables 430-1 and 430-2), although some have overlapping features. For example, PCT, the most common porphyria, is hepatic and presents with blistering cutaneous photosensitivity, which is typically characteristic of the erythropoietic porphyrias. The major manifestations of the acute hepatic porphyrias are neurologic, including neuropathic abdominal pain, peripheral motor neuropathy, and mental disturbances, with attacks often precipitated by dieting, certain drugs, and hormonal changes. While hepatic porphyrias are symptomatic primarily in adults, rare homozygous variants of the autosomal dominant hepatic porphyrias usually manifest clinically prior to puberty.

In contrast, the erythropoietic porphyrias usually present at birth or in early childhood with cutaneous photosensitivity, or in the case

of congenital erythropoietic porphyria (CEP), even in utero as nonimmune hydrops fetalis. Cutaneous sensitivity to sunlight results from excitation of excess porphyrins in the skin by long-wave ultraviolet light, leading to cell damage, scarring, and disfigurement. Thus, the porphyrias are metabolic disorders in which environmental, physiologic, and genetic factors interact to cause disease.

Because many symptoms of the porphyrias are nonspecific, diagnosis is often delayed. Laboratory measurement of porphyrin precursors (5'-aminolevulinic acid [ALA] and porphobilinogen [PBG]) or porphyrins in urine, plasma, erythrocytes, or feces is required to confirm or exclude the various types of porphyria (see below). However, a definite diagnosis requires demonstration of the specific gene defect (Table 430-3). The genes encoding all the heme biosynthetic enzymes have been characterized, permitting identification of the mutations causing each porphyria (Table 430-2). Molecular genetic analyses now make it possible to provide precise heterozygote or homozygote identification and prenatal diagnoses in families with known mutations.

In addition to recent reviews of the porphyrias, informative and up-to-date websites are sponsored by the American Porphyria Foundation (www.porphyrifoundation.com) and the European Porphyria Initiative (www.porphyrria-europe.org). An extensive list of unsafe and safe drugs for individuals with acute porphyrias is provided at the Drug Database for Acute Porphyrias (www.drugs-porphyrria.com).

GLOBAL CONSIDERATIONS



The porphyrias are panethnic metabolic diseases that affect individuals around the globe. The acute hepatic porphyrias—acute intermittent porphyria (AIP), hereditary coproporphyrin (HCP), and variegate porphyria (VP)—are autosomal dominant disorders. The frequency of AIP, the most common acute hepatic porphyria, is ~1 in 20,000 among Caucasian individuals of Western European ancestry, and it is particularly frequent in Scandinavians, with a frequency of ~1 in 10,000 in Sweden. VP is particularly frequent in South Africa, where its high prevalence (>10,000 affected patients) is in part due to a genetic “founder effect.” The autosomal recessive acute hepatic porphyria, ALA dehydratase-deficient porphyria (ADP), is very rare, and less than 20 patients have been identified worldwide.

The erythropoietic protoporphyrias—CEP, erythropoietic protoporphyria (EPP), and X-linked protoporphyria (XLP)—also are panethnic. EPP is the most common porphyria in children, whereas CEP is very rare, with about 200 reported cases worldwide. The frequency of EPP varies globally because most patients have the common low expression *FECH* mutation that varies in frequency in different populations. It rarely occurs in Africans, is present in about 10% of whites, and is frequent (~30%) in the Japanese.

The autosomal recessive porphyrias—ADP, CEP, EPP, and hepatoerythropoietic porphyria (HEP)—are more frequent in regions with high rates of consanguineous unions. PCT, which is typically sporadic, occurs more frequently in countries in which its predisposing risk factors such as hepatitis C and HIV are more prevalent.

HEME BIOSYNTHESIS

Heme biosynthesis involves eight enzymatic steps in the conversion of glycine and succinyl-CoA to heme (Fig. 430-2 and Table 430-2).

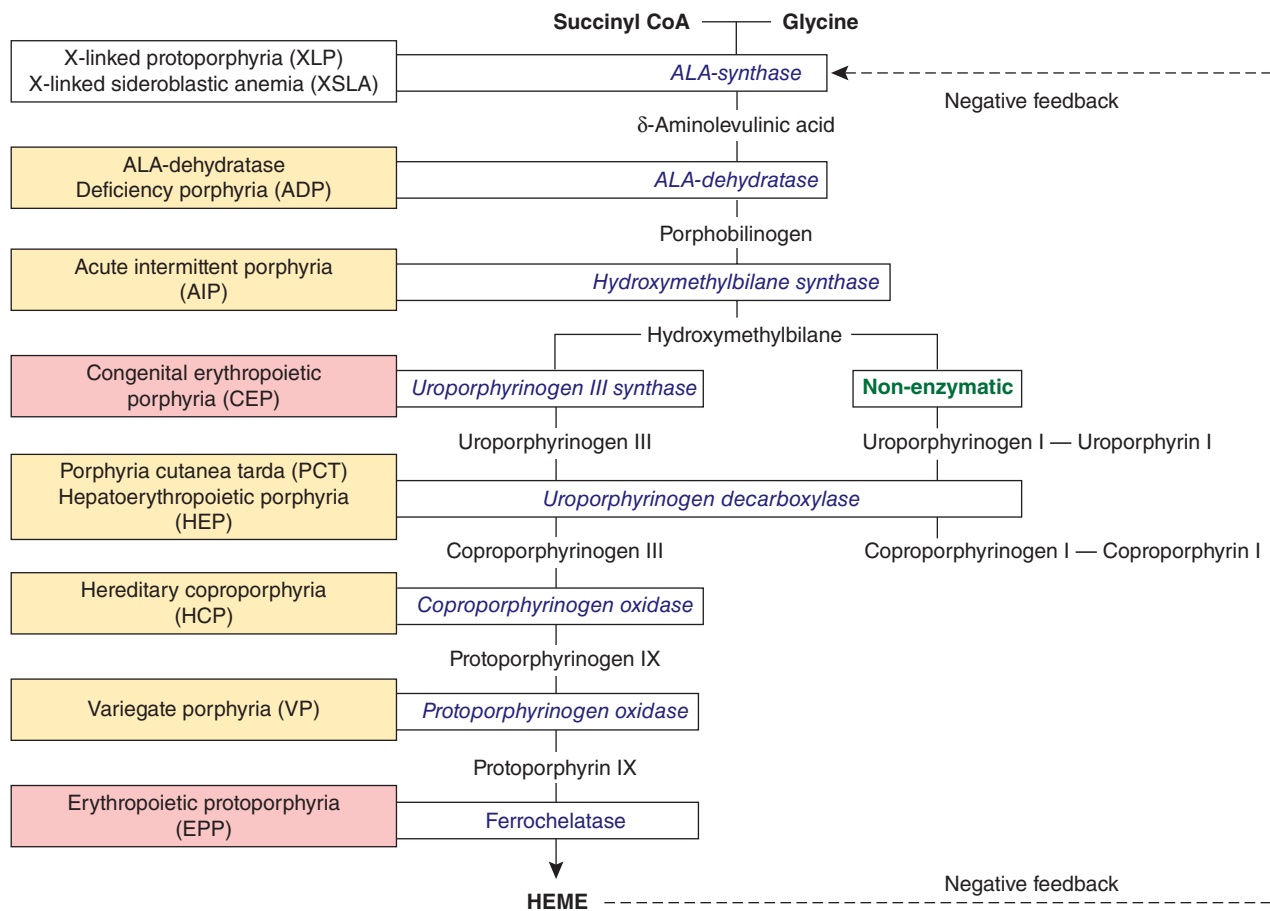


FIGURE 430-1 The human heme biosynthetic pathway indicating in *linked boxes* the enzyme that, when deficient, causes the respective porphyria. Hepatic porphyrias are shown in *yellow boxes* and erythropoietic porphyrias in *pink boxes*.

These eight enzymes are encoded by nine genes, as the first enzyme in the pathway, 5'-aminolevulinic acid synthase (ALA synthase), has two genes that encode unique housekeeping (*ALAS1*) and erythroid-specific (*ALAS2*) isoforms. The first and last three enzymes in the pathway are located in the mitochondrion, whereas the other four are

in the cytosol. Heme is required for a variety of hemoproteins such as hemoglobin, myoglobin, respiratory cytochromes, and the cytochrome P450 enzymes (CYPs). Hemoglobin synthesis in erythroid precursor cells accounts for approximately 85% of daily heme synthesis in humans. Hepatocytes account for most of the rest, primarily for

TABLE 430-1 HUMAN PORPHYRIAS: MAJOR CLINICAL AND LABORATORY FEATURES

Porphyria	Deficient Enzyme	Inheritance	Principal Symptoms: NV or CP+	Enzyme Activity % of Normal	Increased Porphyrin Precursors and/or Porphyrins		
					Erythrocytes	Urine	Stool
Hepatic Porphyrias							
5-ALA dehydratase-deficient porphyria (ADP)	ALA dehydratase	AR	NV	~5	Zn-Protoporphyrin	ALA, coproporphyrin III	—
Acute intermittent porphyria (AIP)	HMB synthase	AD	NV	~50	—	ALA ^a , PBG, uroporphyrin	—
Porphyria cutanea tarda (PCT)	URO decarboxylase	AD	CP	~20	—	Uroporphyrin, 7-carboxylate porphyrin	Isocoproporphyrin
Hereditary coproporphyria (HCP)	COPRO oxidase	AD	NV and CP	~50	—	ALA, PBG, coproporphyrin III	Coproporphyrin III
Variegate porphyria (VP)	PROTO oxidase	AD	NV and CP	~50	—	ALA, PBG, coproporphyrin III	Coproporphyrin III, protoporphyrin
Erythropoietic Porphyrias							
Congenital erythropoietic porphyria (CEP)	URO-synthase	AR	CP	1–5	Uroporphyrin I Coproporphyrin I	Uroporphyrin I ^b Coproporphyrin I ^b	Coproporphyrin I
Erythropoietic protoporphyria (EPP)	Ferrochelatase	AR ^c	CP	~20–30	Protoporphyrin	—	Protoporphyrin
X-linked protoporphyria (XLP)	ALA synthase 2	XL	CP	>100 ^c	Protoporphyrin	—	Protoporphyrin

^aPolymorphism in intron 3 of wild-type allele affects level of enzyme activity and clinical expression. ^bType I isomers. ^cIncreased activity due to "gain-of-function" mutations in *ALAS2* exon 11.

Abbreviations: AD, autosomal dominant; ALA, 5-aminolevulinic acid; AR, autosomal recessive; COPRO I, coproporphyrin I; COPRO III, coproporphyrin III; CP, cutaneous photosensitivity; ISOCOPRO, isocoproporphyrin; + NV, neurovisceral; PBG, porphobilinogen; PROTO, protoporphyrin IX; URO I, uroporphyrin I; URO III, uroporphyrin III; XL, X-linked.

TABLE 430-2 HUMAN HEME BIOSYNTHETIC ENZYMES AND GENES

Enzyme	Gene Symbol	Chromosomal Location	cDNA (bp)	Gene		Protein (aa)	Subcellular Location	Known Mutations ^b	Three-Dimensional Structure ^c
				Size (kb)	Exons ^a				
ALA synthase									
Housekeeping	ALAS1	3p21.1	2199	17	11	640	M	—	—
Erythroid-specific	ALAS2	Xp11.2	1937	22	11	587	M	>30	—
ALA dehydratase									
Housekeeping	ALAD	9q32	1149	15.9	12 (1A + 2 – 12)	330	C	12	Y
Erythroid-specific	ALAD	9q32	1154	15.9	12 (1B + 2 – 12)	330	C	—	—
HMB synthase									
Housekeeping	HMBS	11q23.3	1086	11	15 (1 + 3 – 15)	361	C	>315	E
Erythroid-specific	HMBS	11q23.3	1035	11	15 (2 – 15)	344	C	10	—
URO synthase									
Housekeeping	UROS	10q26.2	1296	34	10 (1 + 2B – 10)	265	C	39	H
Erythroid-specific	UROS	10q26.2	1216	34	10 (2A + 2B – 10)	265	C	4	—
URO decarboxylase	UROD	1p34.1	1104	3	10	367	C	108	H
COPRO oxidase	CPOX	3q12.1	1062	14	7	354	M	51	H
PROTO oxidase	PPOX	1q23.3	1431	5.5	13	477	M	129	—
Ferrochelatase	FECH	18q21.31	1269	45	11	423	M	125	B

^aNumber of exons and those encoding separate housekeeping and erythroid-specific forms indicated in parentheses. ^bNumber of known mutations from the Human Gene Mutation Database (www.hgmd.org). ^cCrystallized from human (H), murine (M), *Escherichia coli* (E), *Bacillus subtilis* (B), or yeast (Y) purified enzyme; references in Protein Data Bank (www.rcsb.org).

Abbreviations: C, cytoplasm; M, mitochondria.

Source: From KE Anderson et al: Disorders of heme biosynthesis: X-linked sideroblastic anemia and the porphyrias, in *The Metabolic and Molecular Bases of Inherited Diseases*, CR Scriver et al (eds). New York, McGraw-Hill, 2001, pp 2991–3062.

TABLE 430-3 DIAGNOSIS OF ACUTE AND CUTANEOUS PORPHYRIAS

Symptoms	First-Line Test: Abnormality	Possible Porphyria	Second-Line Testing if First-Line Testing Is Positive: To include: urine (U), plasma (P), and fecal (F) porphyrins; for acute porphyrias, add red blood cell (RBC) HMB synthase; for blistering skin lesions, add P and RBC porphyrins	Confirmatory Test: Enzyme Assay and/or Mutation Analysis
Neurovisceral	Spot U: ↑↑ALA and normal PBG	ADP	U porphyrins: ↑↑, mostly COPRO III P & F porphyrins: normal or slightly ↑ RBC HMB synthase: normal	Rule out other causes of elevated ALA; ↓↓RBC ALA dehydratase activity (<10%); ALA dehydratase mutation analysis
	Spot U: ↑↑PBG	AIP	U porphyrins: ↑↑, mostly URO and COPRO P & F porphyrins: normal or slightly ↑ RBC HMB synthase: usually ↓	HMB synthase mutation analysis
	"	HCP	U porphyrins: ↑↑, mostly COPRO III P porphyrins: normal or slightly ↑ (↑ if skin lesions present) F porphyrins: ↑↑, mostly COPRO III	Measure RBC HMB synthase: normal activity COPRO oxidase mutation analysis
	"	VP	U porphyrins: ↑↑, mostly COPRO III P porphyrins: ↑↑ (characteristic fluorescence peak at neutral pH) F porphyrins: ↑↑, mostly COPRO and PROTO	Measure RBC HMB synthase: normal activity PROTO oxidase mutation analysis
Blistering skin lesions	P: ↑ porphyrins	PCT and HEP	U porphyrins: ↑↑, mostly URO and heptacarboxylate porphyrin P porphyrins: ↑↑ F porphyrins: ↑↑, including increased isocoproporphyrin RBC porphyrins: ↑↑ zinc PROTO in HEP ^a	RBC URO decarboxylase activity: half-normal in familial PCT (~20% of all PCT cases); substantially deficient in HEP URO decarboxylase mutation analysis: mutation(s) present in familial PCT (heterozygous) and HEP (homozygous)
	"	HCP and VP CEP	See HCP and VP above. Also, U ALA and PBG: may be ↑ RBC and U porphyrins: ↑↑, mostly URO I and COPRO I F porphyrins: ↑↑; mostly COPRO I	↓↓ RBC URO synthase activity (<15%) URO synthase mutation analysis
Nonblistering photosensitivity	P: porphyrins usually ↑	EPP	RBC porphyrins: ↓↓, mostly free PROTO U porphyrins: normal F porphyrins: normal or ↓, mostly PROTO	FECH mutation analysis
	P: porphyrins usually ↑	XLP	RBC porphyrins: ↑↑, approximately equal free and zinc PROTO U porphyrins: normal F porphyrins: normal or ↑, mostly PROTO	ALAS2 mutation analysis

^aNonspecific increases in zinc protoporphyrins are common in other porphyrias.

Abbreviations: ADP, 5-ALA dehydratase-deficient porphyria; AIP, acute intermittent porphyria; ALA, 5-aminolevulinic acid; CEP, congenital erythropoietic porphyria; COPRO I, coproporphyrin I; COPRO III, coproporphyrin III; EPP, erythropoietic protoporphyria; F, fecal; HCP, hereditary coporphyria; HEP; ISOCOPRO, isocoproporphyrin; P, plasma; PBG, porphobilinogen; PCT, porphyria cutanea tarda; PROTO, protoporphyrin IX; RBC, erythrocytes; U, urine; URO I, uroporphyrin I; URO III, uroporphyrin III; VP, variegate porphyria; XLP, X-linked protoporphyria.

Source: Based on KE Anderson et al: *Ann Intern Med* 142:439, 2005.

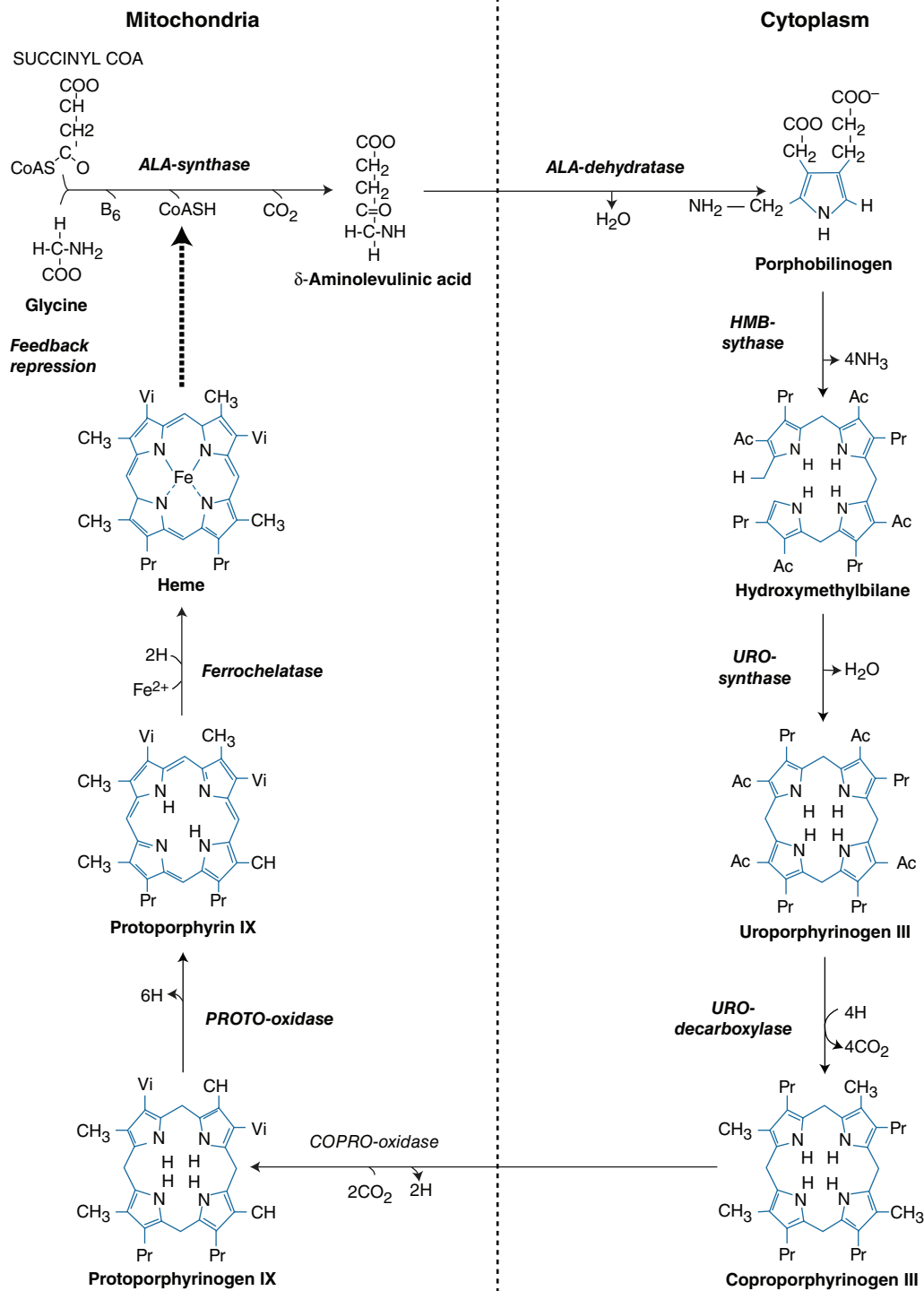


FIGURE 430-2 The heme biosynthetic pathway showing the eight enzymes and their substrates and products. Four of the enzymes are localized in the mitochondria and four in the cytosol.

the synthesis of CYPs, which are especially abundant in the liver endoplasmic reticulum, and turn over more rapidly than many other hemoproteins, such as the mitochondrial respiratory cytochromes. As shown in Fig. 430-2, pathway intermediates are the porphyrin precursors, ALA and PBG, and porphyrins (mostly in their reduced forms, known as *porphyrinogens*). At least in humans, these intermediates do not accumulate in significant amounts under normal conditions or have important physiologic functions.

The first enzyme, ALA synthase, catalyzes the condensation of glycine, activated by pyridoxal phosphate and succinyl coenzyme A, to form ALA. In the liver, this rate-limiting enzyme can be induced by a variety of drugs, steroids, and other chemicals. Distinct nonerythroid

(e.g., housekeeping) and erythroid-specific forms of ALA synthase are encoded by separate genes located on chromosome 3p21.1 (*ALAS1*) and Xp11.2 (*ALAS2*), respectively. Defects in the erythroid gene *ALAS2* that decrease its activity cause X-linked sideroblastic anemia (XLSA). Recently, gain-of-function mutations in the last exon (11) of *ALAS2* that increase its activity have been shown to cause an X-linked form of EPP, known as *X-linked protoporphyria* (XLP).

The second enzyme, ALA dehydratase, catalyzes the condensation of two molecules of ALA to form PBG. Hydroxymethylbilane synthase (HMB synthase; also known as PBG deaminase) catalyzes the head-to-tail condensation of four PBG molecules by a series of deaminations to form the linear tetrapyrrole, HMB. Uroporphyrinogen III synthase

(URO synthase) catalyzes the rearrangement and rapid cyclization of HMB to form the asymmetric, physiologic, octacarboxylate porphyrinogen, uroporphyrinogen (URO'gen) III.

The fifth enzyme in the pathway, uroporphyrinogen decarboxylase (URO decarboxylase), catalyzes the sequential removal of the four carboxyl groups from the acetic acid side chains of URO'gen III to form coproporphyrinogen (COPRO'gen) III, a tetracarboxylate porphyrinogen. This compound then enters the mitochondrion via a specific transporter, ABCB6, where COPRO oxidase, the sixth enzyme, catalyzes the decarboxylation of two of the four propionic acid groups to form the two vinyl groups of protoporphyrinogen (PROTO'gen) IX, a decarboxylate porphyrinogen. Next, PROTO oxidase oxidizes PROTO gen to protoporphyrin IX by the removal of six hydrogen atoms. The product of the reaction is a porphyrin (oxidized form), in contrast to the preceding tetrapyrrole intermediates, which are porphyrinogens (reduced forms). Finally, ferrous iron is inserted into protoporphyrin to form heme, a reaction catalyzed by the eighth enzyme in the pathway, ferrochelatase (also known as heme synthetase or protoheme ferolyase).

REGULATION OF HEME BIOSYNTHESIS

Regulation of heme synthesis differs in the two major heme-forming tissues, the liver and erythron. In the liver, the concentration of “free” heme regulates the synthesis and mitochondrial translocation of the housekeeping form of ALA synthase 1. Heme represses the synthesis of the ALA synthase 1 mRNA and interferes with the transport of the enzyme from the cytosol into mitochondria. Hepatic ALA synthase 1 is increased by many of the same chemicals that induce the cytochrome P450 enzymes in the endoplasmic reticulum of the liver. Because most of the heme in the liver is used for the synthesis of cytochrome P450 enzymes, hepatic ALA synthase 1 and the cytochrome P450s are regulated in a coordinated fashion, and many drugs that induce hepatic ALA synthase 1 also induce the CYP genes. The other hepatic heme biosynthetic enzymes are presumably expressed at constant levels, although their relative activities and kinetic properties differ. For example, normal individuals have high activities of ALA dehydratase, but low activities of HMB synthase, the latter being the second rate-limiting step in the pathway.

In the erythron, novel regulatory mechanisms allow for the production of the very large amounts of heme needed for hemoglobin synthesis. The response to stimuli for hemoglobin synthesis occurs during cell differentiation, leading to an increase in cell number. In contrast, the erythroid-specific ALA synthase 2 is expressed at higher levels than the housekeeping enzyme, and erythroid-specific control mechanisms regulate other pathway enzymes as well as iron transport into erythroid cells. Separate erythroid-specific and nonerythroid or “housekeeping” transcripts are known for the first four enzymes in the pathway. As noted above, housekeeping and erythroid-specific ALA synthases are encoded by genes on different chromosomes, but for each of the next three genes in the pathway, both erythroid and nonerythroid transcripts are transcribed by alternative promoters from their single respective genes (Table 430-2).

CLASSIFICATION OF THE PORPHYRIAS

As mentioned above, the porphyrias can be classified as either *hepatic* or *erythropoietic*, depending on whether the heme biosynthetic intermediates that accumulate arise initially from the liver or developing erythrocytes, or as *acute* or *cutaneous*, based on their clinical manifestations. Table 430-1 lists the porphyrias, their principal symptoms, and major biochemical abnormalities. Four of the five hepatic porphyrias—AIP, HCP, VP, and ADP—present during adult life with acute attacks of neurologic manifestations and elevated levels of one or both of the porphyrin precursors, ALA and PBG, and are thus classified as *acute porphyrias*. Patients with ADP have presented in infancy and adolescence. The fifth hepatic disorder, PCT, presents with blistering skin lesions. HCP and VP also may have cutaneous manifestations similar to PCT.

The erythropoietic porphyrias—CEP, EPP, and the recently described XLP—are characterized by elevations of porphyrins in bone

marrow and erythrocytes and present with cutaneous photosensitivity. The skin lesions in CEP resemble PCT but are usually much more severe, whereas EPP and XLP cause a more immediate, painful, and nonblistering type of photosensitivity. EPP is the most common porphyria to cause symptoms before puberty. Around 20% of EPP patients develop minor abnormalities of liver function, with up to about 5% developing hepatic complications that can become life-threatening. XLP has a clinical presentation similar to EPP causing photosensitivity and liver disease.

DIAGNOSIS OF PORPHYRIA

A few specific and sensitive first-line laboratory tests should be used whenever symptoms or signs suggest the diagnosis of porphyria (Table 430-3). If a first-line test is significantly abnormal, more comprehensive testing should follow to establish the type of porphyria, including the specific causative gene mutation.

Acute Porphyrrias An acute porphyria should be suspected in patients with neurovisceral symptoms after puberty, such as abdominal pain, and when the initial clinical evaluation does not suggest another cause. The urinary porphyrin precursors (ALA and PBG) should be measured (Fig. 430-2). Urinary PBG is virtually always increased during acute attacks of AIP, HCP, and VP and is not substantially increased in any other medical condition. Therefore, this measurement is both sensitive and specific. A method for rapid, in-house testing for urinary PBG, such as the Trace PBG kit (Thermo Scientific), can be used. Results from spot (single-void) urine specimens are highly informative because very substantial increases in PBG are expected during acute attacks of porphyria. A 24-h collection can unnecessarily delay diagnosis. The same spot urine specimen should be saved for quantitative determination of ALA, PBG, and creatinine, in order to confirm the qualitative PBG result and also to detect patients with ADP. Urinary porphyrins may remain increased longer than porphyrin precursors in HCP and VP. Therefore, it is useful to measure total urinary porphyrins in the same sample, keeping in mind that urinary porphyrin increases are often nonspecific. Measurement of urinary porphyrins alone should be avoided for screening, because these may be increased in disorders other than porphyrias, such as chronic liver disease, and misdiagnoses of porphyria can result from minimal increases in urinary porphyrins that have no diagnostic significance. Measurement of erythrocyte HMB synthase is not useful as a first-line test. Moreover, the enzyme activity is not decreased in all AIP patients, a borderline low normal value is not diagnostic, and the enzyme is not deficient in other acute porphyrias.

Cutaneous Porphyrrias Blistering skin lesions due to porphyria are virtually always accompanied by increases in total plasma porphyrins. A fluorometric method is preferred, because the porphyrins in plasma in VP are mostly covalently linked to plasma proteins and may be less readily detected by high-performance liquid chromatography (HPLC). The normal range for plasma porphyrins is somewhat increased in patients with end-stage renal disease.

Although a total plasma porphyrin determination will usually detect EPP and XLP, an erythrocyte protoporphyrin determination is more sensitive. Increases in erythrocyte protoporphyrin occur in many other conditions. Therefore, the diagnosis of EPP must be confirmed by showing a predominant increase in free protoporphyrin rather than zinc protoporphyrin. In XLP, both free and zinc protoporphyrin are markedly increased in approximately equal proportions. Interpretation of laboratory reports can be difficult, because the term *free erythrocyte protoporphyrin* sometimes actually represents zinc protoporphyrin.

More extensive testing is justified when an initial test is positive. A substantial increase in PBG may be due to AIP, HCP, or VP. These acute porphyrias can be distinguished by measuring urinary porphyrins (using the same spot urine sample), fecal porphyrins, and plasma porphyrins. Assays for COPRO oxidase or PROTO oxidase are not widely available. More specifically, mutation analysis by sequencing the genes encoding HMB synthase, COPRO oxidase, and PROTO oxidase will detect almost all disease-causing mutations, and will be

2526 diagnostic even when the levels of urinary ALA and PBG have returned to normal or near normal. The various porphyrias that cause blistering skin lesions are differentiated by measuring porphyrins in urine, feces, and plasma. These porphyrias also should be confirmed at the DNA level by the demonstration of the causative gene mutation(s). It is often difficult to diagnose or “rule out” porphyria in patients who have had suggestive symptoms months or years in the past, and in relatives of patients with acute porphyrias, because porphyrin precursors and porphyrins may be normal. In those situations, detection of the specific gene mutation in the index case can make the diagnosis and facilitate the diagnosis and genetic counseling of at-risk relatives. Consultation with a specialist laboratory and physician will assist in selecting the heme biosynthetic gene or genes to be sequenced.

THE HEPATIC PORPHYRIAS

Markedly elevated plasma and urinary concentrations of the porphyrin precursors, ALA and/or PBG, which originate from the liver, are especially evident during attacks of neurologic manifestations of the four acute porphyrias—ADP, AIP, HCP, and VP. In PCT, excess porphyrins also accumulate initially in the liver and cause chronic blistering of sun-exposed areas of the skin.

ALA DEHYDRATASE-DEFICIENT PORPHYRIA (ADP)

ADP is a rare autosomal recessive acute hepatic porphyria caused by a severe deficiency of ALA dehydratase activity. To date, there are only a few documented cases, some in children or young adults, in which specific gene mutations have been identified. These affected homozygotes had <10% of normal ALA dehydratase activity in erythrocytes, but their clinically asymptomatic parents and heterozygous relatives had about half-normal levels of activity and did not excrete increased levels of ALA. The frequency of ADP is unknown, but the frequency of heterozygous individuals with <50% normal ALA dehydratase activity was ~2% in a screening study in Sweden. Because there are multiple causes for deficient ALA dehydratase activity, it is important to confirm the diagnosis of ADP by mutation analysis.

Clinical Features The clinical presentation depends on the amount of residual ALA dehydratase activity. Four of the documented patients were male adolescents with symptoms resembling those of AIP, including abdominal pain and neuropathy. One patient was an infant with more severe disease, including failure to thrive beginning at birth. The earlier age of onset and more severe manifestations in this patient reflect a more significant deficiency of ALA dehydratase activity. Another patient developed an acute motor polyneuropathy at age 63 that was associated with a myeloproliferative disorder. He was heterozygous for an *ALAD* mutation that presumably was present in erythroblasts that underwent clonal expansion due to the bone marrow malignancy.

Diagnosis All patients had significantly elevated levels of plasma and urinary ALA and urinary coproporphyrin (COPRO) III; ALAD activities in erythrocytes were <10% of normal. Hereditary tyrosinemia type 1 (fumarylacetoacetase deficiency) and lead intoxication should be considered in the differential diagnosis because either succinylacetone (which accumulates in hereditary tyrosinemia and is structurally similar to ALA) or lead can inhibit ALA dehydratase, increase urinary excretion of ALA and COPRO III, and cause manifestations that resemble those of the acute porphyrias. Heterozygotes are clinically asymptomatic and do not excrete increased levels of ALA but can be detected by demonstration of intermediate levels of erythrocyte ALA dehydratase activity or a specific mutation in the *ALAD* gene. To date, molecular studies of ADP patients have identified nine point mutations, two splice-site mutations, and a two-base deletion in the *ALAD* gene (Human Gene Mutation Database; www.hgmd.org). The parents in each case were not consanguineous, and the index cases had inherited a different *ALAD* mutation from each parent. Prenatal diagnosis of this disorder is possible by determination of ALA dehydratase activity and/or gene mutations in cultured chorionic villi or amniocytes.

TREATMENT ALA-DEHYDRATASE DEFICIENT PORPHYRIA

The treatment of ADP acute attacks is similar to that of AIP (see below). The severely affected infant referred to above was supported by hyperalimentation and periodic blood transfusions but did not respond to intravenous hemin and died after liver transplantation.

ACUTE INTERMITTENT PORPHYRIA (AIP)

This hepatic porphyria is an autosomal dominant condition resulting from the half-normal level of HMB synthase activity. The disease is widespread but is especially common in Scandinavia and Great Britain. Clinical expression is highly variable, and activation of the disease is often related to environmental or hormonal factors, such as drugs, diet, and steroid hormones. Attacks can be prevented by avoiding known precipitating factors. Rare homozygous dominant AIP also has been described in children (see below).

Clinical Features Induction of the rate-limiting hepatic enzyme ALA synthase in heterozygotes who have half-normal HMB synthase activity is thought to underlie the acute attacks in AIP. The disorder remains latent (or asymptomatic) in the great majority of those who are heterozygous for *HMB*S mutations, and this is almost always the case prior to puberty. In patients with no history of acute symptoms, porphyrin precursor excretion is usually normal, suggesting that half-normal hepatic HMB synthase activity is sufficient and that hepatic ALA synthase activity is not increased. However, under conditions where heme synthesis is increased in the liver, half-normal HMB synthase activity may become limiting, and ALA, PBG, and other heme pathway intermediates may accumulate and be excreted in the urine. Common precipitating factors include endogenous and exogenous steroids, porphyrinogenic drugs, alcohol ingestion, and low-calorie diets, usually instituted for weight loss.

The fact that AIP is almost always latent before puberty suggests that adult levels of steroid hormones are important for clinical expression. Symptoms are more common in women, suggesting a role for estrogens or progestins. Premenstrual attacks are probably due to endogenous progesterone. Acute porphyrias are sometimes exacerbated by exogenous steroids, including oral contraceptive preparations containing progestins. Surprisingly, pregnancy is usually well tolerated, suggesting that beneficial metabolic changes may ameliorate the effects of high levels of progesterone. **Table 430-4** provides a partial list of the major drugs that are harmful in AIP (and also in HCP and VP). Extensive lists of unsafe and safe drugs are available on websites sponsored by the American Porphyria Foundation (www.porphyrifoundation.com) and the European Porphyria Initiative (www.porphyrria-europe.org), and at the Drug Database for Acute Porphyrias website (www.drugs-porphyrria.com). Reduced intake of calories and carbohydrate, as may occur with illness or attempts to lose weight, can also increase porphyrin precursor excretion and induce attacks of porphyria. Increased carbohydrate intake may ameliorate attacks. Studies in a knockout AIP mouse model indicate that the hepatic *ALAS1* gene is regulated by the peroxisome proliferator-activated receptor γ coactivator 1 α (PGC-1 α). Hepatic PGC-1 α is induced by fasting, which in turn activates *ALAS1* transcription, resulting in increased heme biosynthesis. This finding suggests an important link between nutritional status and the attacks in acute porphyrias. Attacks also can be provoked by infections, surgery, and ethanol.

Because the neurovisceral symptoms rarely occur before puberty and are often nonspecific, a high index of suspicion is required to make the diagnosis. The disease can be disabling but is rarely fatal. Abdominal pain, the most common symptom, is usually steady and poorly localized but may be cramping. Ileus, abdominal distention, and decreased bowel sounds are common. However, increased bowel sounds and diarrhea may occur. Abdominal tenderness, fever, and leukocytosis are usually absent or mild because the symptoms are neurologic rather than inflammatory. Nausea; vomiting; constipation; tachycardia; hypertension; mental symptoms; pain in the limbs, head, neck, or chest; muscle weakness; sensory loss; dysuria; and urinary

TABLE 430-4 UNSAFE DRUGS IN PORPHYRIA

Documented Porphyrinogenic	Probably Porphyrinogenic	Possibly Porphyrinogenic	
Carbamazepine	Altretamine	Aceclofenac	Parecoxib
Carisoprodol	Aminophylline	Acitretin	Pentifylline
Chloramphenicol	Amiodarone	Acrivastine	Pentoxifyverine
Clindamycin	Amitriptyline	Alfuzosin	Phenylpropanolamine + cinnarizine
Dextropropoxyphene	Amlodipine	Anastrozole	Pizotifen
Dihydralazine	Amprenavir	Auranofin	Polidocanol
Dihydroergotamine	Aprepitant	Azelastine	Polyestradiol
Drospirenone + estrogen	Atorvastatin	Benzotropine	Phosphate
Dydrogesterone	Azathioprine	Benzydamine	Potassium canrenoate
Etonogestrel	Bosentan	Betaxolol	Pravastatin
Fosphenytoin sodium	Bromocriptine	Bicalutamide	Prednisolone
Hydralazine	Buspirone	Biperiden	Prilocaine
Hydroxyzine	Busulfan	Bupropion	Progauil
Indinavir	Butylscopolamine	Carvedilol	Propafenone
Ketamine	Cabergoline	Chlorambucil	Pseudoephedrine + dexbrompheniramine
Ketoconazole	Ceftriaxone + lidocaine	Chlorcyclizine + guaifenesin	Quillaia extract
Lidocaine	Cerivastatin	Chloroquine	Quinagolide
Lynestrenol	Cetirizine	Chlorprothixene	Quinine
Lynestrenol + estrogen	Cholinetheophyllinate	Chlorzoxazone	Quinupristin + Dalfopristin
Mecillinam	Clarithromycin	Chorionic	Reboxetine
Medroxyprogesterone	Clemastine	Gonadotropin	Repaglinide
Megestrol	Clonidine	Ciclosporin	Rizatriptan
Methylergometrine	Cyclizine	Cisapride	Rofecoxib
Methyldopa	Cyproterone	Citalopram	Ropinirole
Mifepristone	Danazol	Clomethiazole	Ropivacaine
Nicotinic acid/meclozine/ hydroxyzine	Delavirdine	Clomiphene	Roxithromycin
Nitrofurantoin	Desogestrel + estrogen	Clomipramine	Sertraline
Norethisterone	Diazepam	Clopidogrel	Sevoflurane
Norgestimate + estrogen	Dienogest + estrogen	Clotrimazole	Sibutramine
Orphenadrine	Diclofenac	Cortisone	Sildenafil
Phenobarbital	Diltiazem	Cyclandelate	Sirolimus
Phenytoin	Diphenhydramine	Cyclophosphamide	Sodium aurothiomalate
Pivampicillin	Disopyramide	Cyproheptadine	Sodium oleate + chlorocymol
Pivmecillinam	Disulfiram	Dacarbazine	Stavudine
Primidone	Drospirenone + estrogen	Daunorubicin	Sulindac
Rifampicin	Dydrogesterone	Desogestrel	Sumatriptan
Ritonavir	Ergoloid mesylate	Dichlorobenzyl alcohol	Tacrolimus
Spirolactone	Erythromycin	Dithranol	Tadalafil
Sulfadiazine + trimethoprim	Estramustine	Docetaxel	Tegafur + uracil
Tamoxifen	Ethosuximide	Donepezil	Telmisartan
Testosterone, injection	Etoposide	Doxycycline	Thioridazine
Thiopental	Exemestane	Ebastine	Thioguanine
Trimethoprim	Felbamate	Econazole	Tolfenamic acid
Valproic acid	Felodipine	Efavirenz	Tolterodine
Venlafaxine	Fluconazole	Escitalopram	Torsemede
Vinblastine	Flunitrazepam	Esomeprazole	Triamcinolone
Vincristine	Fluvastatin	Estradiol/tablets	Trihexyphenidyl
Vindesine	Glibenclamide	Estriol/tablets	Trimipramine
Vinorelbine	Halothane	Estrio/vaginal crème, tablet	Valerian
Xylometazoline	Hyoscyamine	Estrogen, conjugate	Venlafaxine
Zaleplon	Ifosfamide	Finasteride	Vinblastine
Ziprasidone	Imipramine	Flecainide	Vincristine
Zolmitriptan	Irinotecan	Flucloxacillin	Vindesine
Zolpidem	Isoniazid	Fluoxetine	Vinorelbine
	Isradipine		Xylometazoline

(Continued)

TABLE 430-4 UNSAFE DRUGS IN PORPHYRIA (CONTINUED)

Documented Porphyrinogenic	Probably Porphyrinogenic	Possibly Porphyrinogenic	
Zuclopenthixol	Itraconazole	Flupentixol	Zaleplon
	Lamivudine + zidovudine	Flutamide	Ziprasidone
	Lansoprazole	Fluvoxamine	Zolmitriptan
	Lercanidipine	Follitropin alfa and beta	Zolpidem
	Levonorgestrel	Galantamine	Zuclopenthixol
	Lidocaine	Glimepiride	
	Lopinavir	Glipizide	
	Lutropin alfa	Gonadorelin	
	Lymecycline	Gramicidin	
	Meclozine	Guaifenesin	
	Medroxyprogesterone + estrogen	Hydrocortisone	
	Metoclopramide	Hydroxycarbamide	
	Metronidazole	Hydroxychloroquine	
	Metyrapone	Ibutilide	
	Moxonidine	Imatinib	
	Nandrolone	Indomethacin	
	Nefazodone	Ketobemidone + DDBA	
	Nelfinavir	Ketoconazole	
	Nevirapine	Ketorolac	
	Nifedipine	Lamotrigine	
	Nimodipine	Letrozole	
	Nitrazepam	Levodopa + benserazide	
	Norethisterone	Levonorgestrel intra-uterine	
	Nortriptyline	Levosimendan	
	Oxcarbazepine	Lidocaine	
	Oxytetracycline	Linezolid	
	Paclitaxel	Lofepramine	
	Paroxetine	Lomustine	
	Phenazone + caffeine	Malathion	
	Pioglitazone	Maprotiline	
	Probenecid	Mebendazole	
	Progesterone, vaginal gel	Mefloquine	
	Quinidine	Melperone	
	Rabeprazole	Melphalan	
	Raloxifene	Mepenzolate	
	Rifabutin	Mepivacaine	
	Riluzole	Mercaptopurine	
	Risperidone	Methadone	
	Rosiglitazone	Methylprednisolone	
	Saquinavir	Methixene	
	Selegiline	Metolazone	
	Simvastatin	Metronidazole	
	Sulfasalazine	Mexiletine	
	Telithromycin	Mianserin	
	Terbinafine	Midazolam	
	Terfenadine	Minoxidil	
	Testosterone, transdermal patch	Mirtazapine	
	Tetracycline	Mitomycin	
	Theophylline	Mitoxantrone	
	Thiamazole	Moclobemide	
	Tibolone	Montelukast	
	Ticlopidine	Morphine + scopolamine	
	Tinidazole	Multivitamins	
	Thiotepa	Mupirocin	
	Topiramate	Nabumetone	
	Topotecan	Nafarelin	
	Toremifene		

(Continued)

TABLE 430-4 UNSAFE DRUGS IN PORPHYRIA (CONTINUED)

Documented Porphyrinogenic	Probably Porphyrinogenic	Possibly Porphyrinogenic
	Tramadol	Naltrexone
	Trimegestone + estrogen	Nateglinide
	Verapamil	Nilutamide
	Voriconazole	Noscapine
	Zidovudine/AZT	Omeprazole
		Oxybutynin
		Oxycodone
		Pantoprazole
		Papaverine

Note: Based on list in "Patient's and Doctor's Guide to Medication in Acute Porphyria," Swedish Porphyria Association and Porphyria Centre Sweden. Also see the website Drug Database for Acute Porphyrias (www.drugs-porphyria.com) for a searchable list of safe and unsafe drugs.

retention are characteristic. Tachycardia, hypertension, restlessness, tremors, and excess sweating are due to sympathetic overactivity.

The peripheral neuropathy is due to axonal degeneration (rather than demyelination) and primarily affects motor neurons. Significant neuropathy does not occur with all acute attacks; abdominal symptoms are usually more prominent. Motor neuropathy affects the proximal muscles initially, more often in the shoulders and arms. The course and degree of involvement are variable and sometimes may be focal and involve cranial nerves. Deep tendon reflexes initially may be normal or hyperactive but become decreased or absent as the neuropathy advances. Sensory changes such as paresthesia and loss of sensation are less prominent. Progression to respiratory and bulbar paralysis and death occurs especially when the diagnosis and treatment are delayed. Sudden death may result from sympathetic overactivity and cardiac arrhythmia.

Mental symptoms such as anxiety, insomnia, depression, disorientation, hallucinations, and paranoia can occur in acute attacks. Seizures can be due to neurologic effects or to hyponatremia. Treatment of seizures is difficult because most antiseizure drugs can exacerbate AIP (clonazepam may be safer than phenytoin or barbiturates). Hyponatremia results from hypothalamic involvement and inappropriate vasopressin secretion or from electrolyte depletion due to vomiting, diarrhea, poor intake, or excess renal sodium loss. Persistent hypertension and impaired renal function may occur. When an attack resolves, abdominal pain may disappear within hours, and paresis begins to improve within days and may continue to improve over several years.

Homozygous dominant AIP is a rare form of AIP in which patients inherit *HMBS* mutations from each of their heterozygous parents and, therefore, have very low (<2%) enzyme activity. The disease has been described in a Dutch girl, two young British siblings, and a Spanish boy. In these homozygous affected patients, the disease presented in infancy with failure to thrive, developmental delay, bilateral cataracts, and/or hepatosplenomegaly. Urinary ALA and PBG concentrations were markedly elevated. All of these patients' *HMBS* mutations (R167W, R167Q, and R172Q) were in exon 10 within five bases of each other. Studies of the brain magnetic resonance images (MRIs) of children with homozygous AIP have suggested damage primarily in white matter that was myelinated postnatally, while tracks that myelinated prenatally were normal. Most children with homozygous AIP die at an early age.

Diagnosis ALA and PBG levels are substantially increased in plasma and urine, especially during acute attacks, and become normal only after prolonged latency. For example, urinary PBG excretion during an attack is usually 50–200 mg/24 h (220–880 μ mol/24 h) (normal, 0–4 mg/24 h, [0–18 μ mol/24 h]), and urinary ALA excretion is 20–100 mg/24 h (150–760 μ mol/24 h) (normal, 1–7 mg/24 h [8–53 μ mol/24 h]). Because levels often remain high after symptoms resolve, the diagnosis of an acute attack in a patient with biochemically proven AIP is based primarily on clinical features. Excretion of ALA and PBG decreases over a few days after intravenous heme administration. A normal urinary PBG level before heme effectively excludes AIP as a cause for current symptoms. Fecal porphyrins are usually normal or minimally increased in AIP, in contrast to HCP and VP. Most AIP heterozygotes

with no history of symptoms have normal urinary excretion of ALA and PBG. Therefore, the detection of the family's *HMBS* mutation will diagnose asymptomatic family members.

Patients with *HMBS* mutations in the initiation of translation codon in exon 1 and in the intron 15'-splice donor site have normal enzyme levels in erythrocytes and deficient activity only in nonerythroid tissues. This occurs because the erythroid and housekeeping forms of HMB synthase are encoded by a single gene, which has two promoters. Thus, the enzyme assay may not be diagnostic, and genetic testing should be used to confirm the diagnosis.

More than 390 *HMBS* mutations have been identified in AIP, including missense, nonsense, and splicing mutations and insertions and deletions, with most mutations found in only one or a few families (Human Gene Mutation Database, www.hgmd.org). The prenatal diagnosis of a fetus at risk can be made with cultured amniotic cells or chorionic villi. However, this is seldom done, because the prognosis of individuals with *HMBS* mutations is generally favorable.

TREATMENT ACUTE INTERMITTENT PORPHYRIA

During acute attacks, narcotic analgesics may be required for abdominal pain, and phenothiazines are useful for nausea, vomiting, anxiety, and restlessness. Chloral hydrate can be given for insomnia, and benzodiazepines are probably safe in low doses if a minor tranquilizer is required. Carbohydrate loading, usually with intravenous glucose (at least 300 g daily), may be effective in milder acute attacks of porphyria (without paresis, hyponatremia, etc.) if heme is not available. Intravenous heme is more effective and should be used as first-line therapy for all acute attacks. The standard regimen is 3–4 mg/kg of heme, in the form of lyophilized heme (Recordati Pharmaceuticals), heme albumin (heme albumin reconstituted with human albumin), or heme arginate (Orphan Europe), infused daily for 4 days. Heme arginate and heme albumin are chemically stable and are less likely than heme to produce phlebitis or an anticoagulant effect. Recovery depends on the degree of neuronal damage and usually is rapid if therapy is started early. Recovery from severe motor neuropathy may require months or years. Identification and avoidance of inciting factors can hasten recovery from an attack and prevent future attacks. Inciting factors are usually multiple, and removal of one or more hastens recovery and helps prevent future attacks. Frequent attacks that occur during the luteal phase of the menstrual cycle may be prevented with a gonadotropin-releasing hormone analogue, which prevents ovulation and progesterone production, or by prophylactic heme administration.

The long-term risk of hypertension and chronic renal disease is increased in AIP; a number of patients have undergone successful renal transplantation. Chronic, low-grade abnormalities in liver function tests are common, and the risk of hepatocellular carcinoma is increased. Hepatic imaging is recommended at least yearly for early detection of these tumors.

An allogeneic liver transplant was performed on a 19-year-old female AIP heterozygote who had 37 acute attacks in the 29 months

2530 prior to transplantation. After transplantation, her elevated urinary ALA and PBG levels returned to normal in 24 h, and she did not experience acute neurologic attacks for more than 3 years after transplant. Two AIP patients had combined liver and kidney transplants secondary to uncontrolled acute porphyria attacks, chronic peripheral neuropathy, and renal failure requiring dialysis. Both patients had a marked improvement with no attacks and normal urinary PBG levels after transplantation, as well as improvement of their neuropathic manifestations. More recently, a group from the United Kingdom reported their experience with liver transplantation in 10 AIP patients with recurrent attacks that were refractory to medical management and impaired quality of life. Patients had a complete biochemical and symptomatic resolution after transplant. The investigators reported a high rate of hepatic artery thrombosis in their series. Clearly, liver transplantation is a high-risk procedure and should be considered as a last resort in patients with severe recurrent attacks. Recently, liver-directed gene therapy has proven successful in the prevention of drug-induced biochemical attacks in a murine model of human AIP, and clinical trials of AAV-HMBS gene transfer have been initiated. In addition, preclinical studies of a hepatic-targeted RNA interference (RNAi) therapy directed to inhibit the markedly elevated hepatic *ALAS1* mRNA in the AIP mouse model prevented induced biochemical attacks and rapidly reduced the *ALAS1* mRNA during an ongoing attack.

PORPHYRIA CUTANEA TARDA (PCT)

PCT, the most common of the porphyrias, can be either sporadic (type 1) or familial (type 2) and can also develop after exposure to halogenated aromatic hydrocarbons. Hepatic URO decarboxylase is deficient in all types of PCT, and for clinical symptoms to manifest, this enzyme deficiency must be substantial (~20% of normal activity or less); it is currently attributed to generation of an URO decarboxylase inhibitor in the liver, which forms uroporphomethene in the presence of iron and under conditions of oxidative stress. The majority of PCT patients (~80%) have no *UROD* mutations and are said to have sporadic (type 1) disease. PCT patients heterozygous for *UROD* mutations have familial (type 2) PCT. In these patients, inheritance of a *UROD* mutation from one parent results in half-normal enzyme activity in liver and all other tissues, which is a significant predisposing factor, but is insufficient by itself to cause symptomatic PCT. As discussed below, other genetic and environmental factors contribute to susceptibility for both types of PCT. Because penetrance of the genetic trait is low, many patients with familial (type 2) PCT have no family history of the disease. HEP is an autosomal recessive form of porphyria that results from the marked systemic deficiency of URO decarboxylase activity with clinical symptoms in childhood.

Clinical Features Blistering skin lesions that appear most commonly on the backs of the hands are the major clinical feature (Fig. 430-3). These rupture and crust over, leaving areas of atrophy and scarring. Lesions may also occur on the forearms, face, legs, and feet. Skin friability and small white papules termed milia are common, especially on the backs of the hands and fingers. Hypertrichosis and hyperpigmentation, especially of the face, are especially troublesome in women. Occasionally, the skin over sun-exposed areas becomes severely thickened, with scarring and calcification that resembles systemic sclerosis. Neurologic features are absent.

A number of susceptibility factors, in addition to inherited *UROD* mutations in type 2 PCT, can be recognized clinically and can affect management. These include hepatitis C, HIV, excess alcohol, elevated iron levels, and estrogens. The importance of excess hepatic iron as a precipitating factor is underscored by the finding that the incidence of the common hemochromatosis-causing mutations, hemochromatosis gene (*HFE*) mutations C282Y and H63D, are increased in patients with types 1 and 2 PCT (Chap. 428). Excess alcohol is a long-recognized contributor, as is estrogen use in women. HIV is probably an independent but less common risk factor that, like hepatitis C, does not cause PCT in isolation. Multiple susceptibility factors that appear to act synergistically can be identified in the individual PCT patient. Patients



FIGURE 430-3 Typical cutaneous lesions in a patient with porphyria cutanea tarda. Chronic, crusted lesions resulting from blistering due to photosensitivity on the dorsum of the hand of a patient with porphyria cutanea tarda. (Courtesy of Dr. Karl E. Anderson; with permission.)

with PCT characteristically have chronic liver disease and sometimes cirrhosis and are at risk for hepatocellular carcinoma. Various chemicals can also induce PCT; an epidemic of PCT occurred in eastern Turkey in the 1950s as a consequence of wheat contaminated with the fungicide hexachlorobenzene. PCT also occurs after exposure to other chemicals, including di- and trichlorophenols and 2,3,7,8-tetrachlorodibenzo-(*p*)-dioxin (TCDD, dioxin).

Diagnosis Porphyrins are increased in the liver, plasma, urine, and stool. The urinary ALA level may be slightly increased, but the PBG level is normal. Urinary porphyrins consist mostly of uroporphyrins and heptacarboxylate porphyrin, with lesser amounts of coproporphyrin and hexa- and pentacarboxylate porphyrins. Plasma porphyrins are also increased, and fluorometric scanning of diluted plasma at neutral pH can rapidly distinguish VP and PCT (Table 430-3). Isocoproprophyrins, which are increased in feces and sometimes in plasma and urine, are diagnostic for hepatic URO decarboxylase deficiency.

Type 2 PCT and HEP can be distinguished from type 1 by finding decreased URO decarboxylase in erythrocytes. URO decarboxylase activity in liver, erythrocytes, and cultured skin fibroblasts in type 2 PCT is approximately 50% of normal in affected individuals and in family members with latent disease. In HEP, the URO decarboxylase activity is markedly deficient, with typical levels of 3–10% of normal. Over 121 mutations have been identified in the *UROD* gene (Human Gene Mutation Database; www.hgmd.org). Of the mutations listed in the database, ~65% are missense or nonsense and ~10% are splice-site mutations. Most *UROD* mutations have been identified in only one or two families.

TREATMENT PORPHYRIA CUTANEA TARDA

Alcohol, estrogens, iron supplements, and, if possible, any drugs that may exacerbate the disease should be discontinued, but this step does not always lead to improvement. A complete response can almost always be achieved by the standard therapy, repeated phlebotomy, to reduce hepatic iron. A unit (450 mL) of blood can be removed every 1–2 weeks. The aim is to gradually reduce excess hepatic iron until the serum ferritin level reaches the lower limits of normal. Because iron overload is not marked in most cases, remission may occur after only five or six phlebotomies; however, PCT patients with hemochromatosis may require more treatments to bring their iron levels down to the normal range. To document improvement in PCT, it is most convenient to follow the total plasma porphyrin concentration, which becomes normal some time after the target ferritin level is reached. Hemoglobin levels or hematocrits and serum ferritin should be followed closely to prevent development of iron deficiency and anemia. After remission, continued phlebotomy may not be needed. Plasma porphyrin levels are

followed at 6- to 12-month intervals for early detection of recurrences, which are treated by additional phlebotomy.

An alternative when phlebotomy is contraindicated or poorly tolerated is a low-dose regimen of chloroquine or hydroxychloroquine, both of which complex with the excess porphyrins and promote their excretion. Small doses (e.g., 125 mg chloroquine phosphate twice weekly) should be given, because standard doses can induce transient, sometimes marked increases in photosensitivity and hepatocellular damage. Recent studies indicate that low-dose hydroxychloroquine is as safe and effective as phlebotomy in PCT. Hepatic imaging can diagnose or exclude complicating hepatocellular carcinoma. Treatment of PCT in patients with end-stage renal disease is facilitated by administration of erythropoietin.

HEREDITARY COPROPORPHYRIA (HCP)

HCP is an autosomal dominant hepatic porphyria that results from the half-normal activity of COPRO oxidase. The disease presents with acute attacks, as in AIP. Cutaneous photosensitivity also may occur, but much less commonly than in VP. HCP patients may have acute attacks and cutaneous photosensitivity together or separately. HCP is less common than AIP and VP. Homozygous dominant HCP and harderoporphyria, a biochemically distinguishable variant of HCP, present with clinical symptoms in children (see below).

Clinical Features HCP is influenced by the same factors that cause attacks in AIP. The disease is latent before puberty, and symptoms, which are virtually identical to those of AIP, are more common in women. HCP is generally less severe than AIP. Blistering skin lesions are identical to PCT and VP and begin in childhood in rare homozygous cases.

Diagnosis COPRO III is markedly increased in the urine and feces in symptomatic patients, and often persists, especially in feces, when there are no symptoms. Urinary ALA and PBG levels are increased (but less than in AIP) during acute attacks, but may revert to normal more quickly than in AIP when symptoms resolve. Plasma porphyrins are usually normal or only slightly increased, but they may be higher in cases with skin lesions. The diagnosis of HCP is readily confirmed by increased fecal porphyrins consisting almost entirely of COPRO III, which distinguishes it from other porphyrias.

Although the diagnosis can be confirmed by measuring COPRO oxidase activity, the assays for this mitochondrial enzyme are not widely available and require cells other than erythrocytes. To date, over 64 mutations have been identified in the *CPOX* gene, 67% of which are missense or nonsense (Human Gene Mutation Database; www.hgmd.org). Detection of a *CPOX* mutation in a symptomatic individual permits the identification of asymptomatic family members.

TREATMENT HEREDITARY COPROPORPHYRIA

Neurologic symptoms are treated as in AIP (see above). Phlebotomy and chloroquine are not effective for the cutaneous lesions.

VARIEGATE PORPHYRIA (VP)



VP is an autosomal dominant hepatic porphyria that results from the deficient activity of PROTO oxidase, the seventh enzyme in the heme biosynthetic pathway, and can present with neurologic symptoms, photosensitivity, or both. VP is particularly common in South Africa, where 3 of every 1000 whites have the disorder. Most are descendants of a couple who emigrated from Holland to South Africa in 1688. In other countries, VP is less common than AIP. Rare cases of homozygous dominant VP, presenting in childhood with cutaneous symptoms, also have been reported.

Clinical Features VP can present with skin photosensitivity, acute neurovisceral crises, or both. In two large studies of VP patients, 59% had only skin lesions, 20% had only acute attacks, and 22% had both. Acute attacks are identical to those in AIP and are precipitated by the same factors as AIP (see above). Blistering skin manifestations are

similar to those in PCT, but are more difficult to treat and usually are of longer duration. Homozygous VP is associated with photosensitivity, neurologic symptoms, and developmental disturbances, including growth retardation, in infancy or childhood; all cases had increased erythrocyte levels of zinc protoporphyrin, a characteristic finding in all homozygous porphyrias so far described.

Diagnosis Urinary ALA and PBG levels are increased during acute attacks, but may return to normal more quickly than in AIP. Increases in fecal protoporphyrin and COPRO III and in urinary COPRO III are more persistent. Plasma porphyrin levels also are increased, particularly when there are cutaneous lesions. VP can be distinguished rapidly from all other porphyrias by examining the fluorescence emission spectrum of porphyrins in plasma since VP has a unique fluorescence peak at neutral pH.

Assays of PROTO oxidase activity in cultured fibroblasts or lymphocytes are not widely available. Over 174 mutations have been identified in the *PPOX* gene from unrelated VP patients (Human Gene Mutation Database; www.hgmd.org). The missense mutation R59W is the common mutation in most South Africans with VP of Dutch descent. Five missense mutations were common in English and French VP patients; however, most mutations have been found in only one or two families.

TREATMENT VARIEGATE PORPHYRIA

Acute attacks are treated as in AIP, and hemin should be started early in most cases. Other than avoiding sun exposure, there are few effective measures for treating the skin lesions. β -Carotene, phlebotomy, and chloroquine are not helpful.

THE ERYTHROPOIETIC PORPHYRIAS

In the erythropoietic porphyrias, excess porphyrins from bone marrow erythrocyte precursors are transported via the plasma to the skin and lead to cutaneous photosensitivity.

X-LINKED SIDEROBLASTIC ANEMIA (XLSA)

XLSA results from the deficient activity of the erythroid form of ALA synthase (ALA synthase 2) and is associated with ineffective erythropoiesis, weakness, and pallor.

Clinical Features Typically, males with XLSA develop refractory hemolytic anemia, pallor, and weakness during infancy. They have secondary hypersplenism, become iron overloaded, and can develop hemosiderosis. The severity depends on the level of residual erythroid ALA synthase activity and on the responsiveness of the specific mutation to pyridoxal 5'-phosphate supplementation (see below). Peripheral blood smears reveal a hypochromic, microcytic anemia with striking anisocytosis, poikilocytosis, and polychromasia; the leukocytes and platelets appear normal. Hemoglobin content is reduced, and the mean corpuscular volume and mean corpuscular hemoglobin concentration are decreased. Patients with milder, late-onset disease have been reported recently.

Diagnosis Bone marrow examination reveals hypercellularity with a left shift and megaloblastic erythropoiesis with an abnormal maturation. A variety of Prussian blue-staining sideroblasts are observed. Levels of urinary porphyrin precursors and of both urinary and fecal porphyrins are normal. The activity of erythroid ALA synthase 2 is decreased in bone marrow, but this enzyme is difficult to measure in the presence of the normal ALA synthase 1 housekeeping enzyme. Definitive diagnosis requires the demonstration of mutations in the erythroid *ALAS2* gene.

TREATMENT X-LINKED SIDEROBLASTIC ANEMIA

The severe anemia may respond to pyridoxine supplementation. This cofactor is essential for ALA synthase activity, and mutations in the pyridoxine binding site of the enzyme have been found in

2532 several responsive patients. Cofactor supplementation may make it possible to eliminate or reduce the frequency of transfusion. Unresponsive patients may be transfusion-dependent and require chelation therapy.

CONGENITAL ERYTHROPOIETIC PORPHYRIA (CEP)

CEP, also known as Günther's disease, is an autosomal recessive disorder. It is due to the markedly deficient, but not absent, activity of URO synthase and the resultant accumulation of URO I and COPRO I isomers. CEP is associated with hemolytic anemia and cutaneous lesions.

Clinical Features Severe cutaneous photosensitivity typically begins in early infancy. The skin over light-exposed areas is friable, and bullae and vesicles are prone to rupture and infection. Skin thickening, focal hypo- and hyperpigmentation, and hypertrichosis of the face and extremities are characteristic. Secondary infection of the cutaneous lesions can lead to disfigurement of the face and hands. Porphyrins are deposited in teeth and in bones. As a result, the teeth are brownish and fluoresce on exposure to long-wave ultraviolet light. Hemolysis is probably due to the marked increase in erythrocyte porphyrins and leads to splenomegaly. Adults with a milder later-onset form of the disease also have been described.

Diagnosis URO and COPRO (mostly type I isomers) accumulate in the bone marrow, erythrocytes, plasma, urine, and feces. The predominant porphyrin in feces is COPRO I. The diagnosis of CEP can be confirmed by demonstration of markedly deficient URO synthase activity and/or by the identification of specific mutations in the *UROS* gene. The disease can be detected in utero by measuring porphyrins in amniotic fluid and URO synthase activity in cultured amniotic cells or chorionic villi, or by the detection of the family's specific gene mutations. Molecular analyses of the mutant alleles from unrelated patients have revealed the presence of over 48 mutations in the *UROS* gene, including four in the erythroid-specific promoter of the *UROS* gene. Genotype/phenotype correlations can predict the severity of the disease. The CEP phenotype may be modulated by sequence variations in the erythroid specific ALA synthase 2, mutation of which typically causes XLP. One mutation (p.ArgR216WTrp) in *GATA1*, encoding the X-linked erythroid-specific transcription factor GATA binding protein 1 (*GATA1*), has been identified in an individual with CEP, thrombocytopenia, and β thalassemia.

TREATMENT CONGENITAL ERYTHROPOIETIC PORPHYRIA

Severe cases often require transfusions for anemia. Chronic transfusions of sufficient blood to suppress erythropoiesis are effective in reducing porphyrin production but result in iron overload. Splenectomy may reduce hemolysis and decrease transfusion requirements. Protection from sunlight and from minor skin trauma is important. β -Carotene may be of some value. Complicating bacterial infections should be treated promptly. Recently, bone marrow and cord blood transplantation has proven curative in several transfusion-dependent children, providing the rationale for stem cell gene therapy.

ERYTHROPOIETIC PROTOPORPHYRIA (EPP)

EPP is an inherited disorder resulting from the deficient activity of ferrochelatase (FECH), the last enzyme in the heme biosynthetic pathway. EPP is the most common erythropoietic porphyria in children and, after PCT, the second most common porphyria in adults. EPP patients have FECH activities as low as 15–25% of normal in lymphocytes and cultured fibroblasts. Protoporphyrin accumulates in bone marrow reticulocytes and then appears in plasma, is taken up in the liver, and is excreted in bile and feces. Protoporphyrin transported to the vessels in the skin causes the nonblistering photosensitivity. In most symptomatic patients (~90%) with this autosomal recessive disorder, a mutation in one *FECH* allele is inherited with a relatively

common (~10% of normal whites) intronic 3 (IVS3) alteration (IVS3-48T>C) that results in the low expression of the normal enzyme. In about 10% of EPP families, two *FECH* mutations have been found. Recently, deletion mutations in exon 11 of the *ALAS2* gene have been described, causing XLP that is clinically indistinguishable from EPP. The deletion of the C-terminal amino acids of *ALAS2* results in its increased activity and the accumulation of protoporphyrin. XLP accounts for approximately 2–10% of cases with the EPP phenotype in Europe and North America.

Clinical Features Skin photosensitivity, which differs from that in other porphyrias, usually begins in childhood and consists of pain, redness, and itching occurring within minutes of sunlight exposure (Fig. 430-4). Photosensitivity is associated with substantial elevations in erythrocyte protoporphyrin and occurs only in patients with genotypes that result in ferrochelatase activities below ~35% of normal. Vesicular lesions are uncommon. Redness, swelling, burning, and itching can develop shortly after sun exposure and resemble angioedema. Pain symptoms may seem out of proportion to the visible skin involvement. Sparse vesicles and bullae occur in ~10% of cases. Chronic skin changes may include lichenification, leathery pseudovesicles, labial grooving, and nail changes. Severe scarring is rare, as are pigment changes, friability, and hirsutism. Unless hepatic or other complications develop, protoporphyrin levels and symptoms of photosensitivity remain remarkably stable over many years in most patients. Factors that exacerbate the hepatic porphyrias play little or no role in EPP.

The primary source of excess protoporphyrin is the bone marrow reticulocytes. Erythrocyte protoporphyrin is free (not complexed with zinc) and is mostly bound to hemoglobin. In plasma, protoporphyrin is bound to albumin. Hemolysis and anemia are usually absent or mild.

Although EPP is an erythropoietic porphyria, up to 20% of EPP patients may have minor abnormalities of liver function, and in about 5% of these patients the accumulation of protoporphyrins causes chronic liver disease that can progress to liver failure and death. Protoporphyrin is insoluble, and excess amounts form crystalline structures in liver cells (Fig. 430-4) and can decrease hepatic bile flow. Studies in the mouse model of EPP have shown that the bile duct epithelium may be damaged by toxic bile, leading to biliary fibrosis. Thus, rapidly progressive liver disease appears to be related to the cholestatic effects of protoporphyrins and is associated with increasing hepatic protoporphyrin levels due to impaired hepatobiliary excretion and increased photosensitivity. The hepatic complications also are often characterized by increasing levels of protoporphyrins in erythrocytes and plasma as well as severe abdominal and back pains, especially in the right upper quadrant. Gallstones



FIGURE 430-4 Erythema and edema of the hands due to acute photosensitivity in a 10-year-old boy with erythropoietic protoporphyria. (From P Poblette-Gutierrez et al: *Eur J Dermatol* 16:230, 2006.)

composed at least in part of protoporphyrin occur in some patients. Hepatic complications appear to be higher in autosomal recessive EPP due to two *FECH* mutations and in XLP.

Diagnosis A substantial increase in erythrocyte protoporphyrin, which is predominantly free and not complexed with zinc, is the hallmark of EPP. Protoporphyrin levels are also variably increased in bone marrow, plasma, bile, and feces. Erythrocyte protoporphyrin concentrations are increased in other conditions such as lead poisoning, iron deficiency, various hemolytic disorders, all homozygous forms of other porphyrias, and sometimes even in acute porphyrias. In all these conditions, however, in contrast to EPP, protoporphyrin is complexed with zinc. Therefore, after an increase in erythrocyte protoporphyrin is found in a suspected EPP patient, it is important to confirm the diagnosis by an assay that distinguishes free and zinc-complexed protoporphyrin. Erythrocytes in EPP also exhibit red fluorescence under a fluorescence microscopy at 620 nm. Urinary levels of porphyrins and porphyrin precursors are normal. Ferrochelatase activity in cultured lymphocytes or fibroblasts is decreased. DNA diagnosis by mutation analysis is recommended to detect the causative *FECH* mutation(s) and/or the presence of the IVS3–48T>C low expression allele. To date, over 190 mutations have been identified in the *FECH* gene, many of which result in an unstable or absent enzyme protein (null alleles) (Human Gene Mutation Database; www.hgmd.org). Studies suggest that EPP patients with a null allele (and the IVS3–48T>C low expression allele) have a greater risk for developing severe liver complications.

In XLP, the erythrocyte protoporphyrin levels appear to be higher than other forms of EPP and the proportions of free and zinc protoporphyrins may reach 50%. To date, four *ALAS2* mutations, three deletions of one to four bases, and one novel nonsense mutation have been described, which markedly increase ALA synthase 2 activity and cause XLP. XLP accounts for about 2% of patients with the EPP phenotype in Western Europe. Recent studies show that about 10% of North American patients with the EPP phenotype have XLP.

TREATMENT ERYTHROPOIETIC PROTOPORPHYRIA

Avoiding sunlight exposure and wearing clothing designed to provide protection for conditions with chronic photosensitivity are essential. Oral β -carotene (120–180 mg/dL) may improve tolerance to sunlight in some patients. The beneficial effects of β -carotene may involve quenching of singlet oxygen or free radicals. The dosage may need to be adjusted to maintain serum carotene levels in the recommended range of 10–15 mmol/L (600–800 mg/dL). Mild skin discoloration due to carotenemia is the only significant side effect. Afamelanotide, an α -melanocyte-stimulating hormone (MSH) analogue has completed phase III clinical trials in the United States for patients with EPP and XLP.

Treatment of hepatic complications, which may be accompanied by motor neuropathy, is difficult. Cholestyramine and other porphyrin absorbents such as activated charcoal may interrupt the enterohepatic circulation of protoporphyrin and promote its fecal excretion, leading to some improvement. Splenectomy may be helpful when the disease is accompanied by hemolysis and significant splenomegaly. Plasmapheresis and intravenous hemein are sometimes beneficial.

Liver transplantation has been carried out in some EPP and XLP patients with severe liver complications and is often successful in the short term. However, the disease often recurs in the transplanted liver due to continued bone marrow production of excess protoporphyrin. In a retrospective study of 17 liver-transplanted EPP patients, 11 (65%) had recurrent EPP liver disease. Posttransplantation treatment with hemein and plasmapheresis should be considered to prevent the recurrence of liver disease. However, bone marrow transplantation, which has been successful in human EPP and which prevented liver disease in a mouse model, should be considered after liver transplantation, if a suitable donor can be found.

ACKNOWLEDGMENT

The authors thank Dr. Karl E. Anderson for his review of the manuscript and helpful comments and suggestions. This work is supported in part by the Porphyrrias Consortium (U54 DK083909), a part of the National Institutes of Health (NIH) Rare Disease Clinical Research Network (RDCRN), supported through collaboration between the NIH Office of Rare Diseases Research (ORDR) at the National Center for Advancing Translational Science (NCATS), and the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. MB is supported by a career development award K23-DK-095946

Purines (adenine and guanine) and pyrimidines (cytosine, thymine, uracil) serve fundamental roles in the replication of genetic material, gene transcription, protein synthesis, and cellular metabolism. Disorders that involve abnormalities of nucleotide metabolism range from relatively common diseases such as hyperuricemia and gout, in which there is increased production or impaired excretion of a metabolic end product of purine metabolism (uric acid), to rare enzyme deficiencies that affect purine and pyrimidine synthesis or degradation. Understanding these biochemical pathways has led, in some instances, to the development of specific forms of treatment, such as the use of allopurinol and febuxostat to reduce uric acid production.

URIC ACID METABOLISM

Uric acid is the final breakdown product of purine degradation in humans. It is a weak diprotic acid with pK_a values of 5.75 and 10.3. Urates, the ionized forms of uric acid, predominate in plasma, extracellular fluid, and synovial fluid, with ~98% existing as monosodium urate at pH 7.4.

Plasma is saturated with monosodium urate at a concentration of 405 $\mu\text{mol/L}$ (6.8 mg/dL) at 37°C. At higher concentrations, plasma is therefore supersaturated—a situation that creates the potential for urate crystal precipitation. However, plasma urate concentrations can reach 4800 $\mu\text{mol/L}$ (80 mg/dL) without precipitation, perhaps because of the presence of solubilizing substances.

The pH of urine greatly influences the solubility of uric acid. At pH 5.0, urine is saturated with uric acid at concentrations ranging from 360 to 900 $\mu\text{mol/L}$ (6–15 mg/dL). At pH 7, saturation is reached at concentrations from 9840 to 12,000 $\mu\text{mol/L}$ (158–200 mg/dL). Ionized forms of uric acid in urine include monosodium, disodium, potassium, ammonium, and calcium urates.

Although purine nucleotides are synthesized and degraded in all tissues, urate is produced only in tissues that contain xanthine oxidase, primarily the liver and small intestine. Urate production varies with the purine content of the diet and with rates of purine biosynthesis, degradation, and salvage (Fig. 431e-1). Normally, two-thirds to three-fourths of

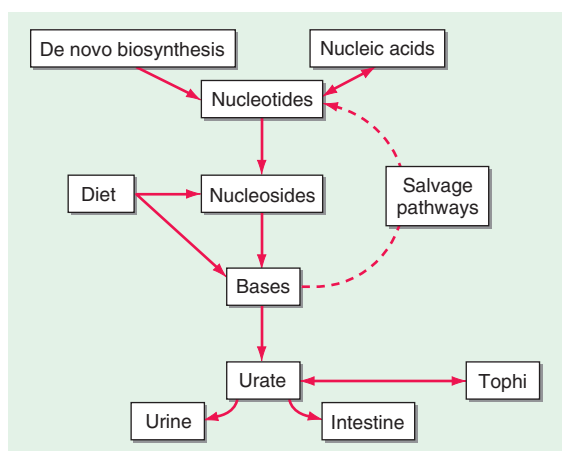


FIGURE 431e-1 The total-body urate pool is the net result between urate production and excretion. Urate production is influenced by dietary intake of purines and the rates of de novo biosynthesis of purines from nonpurine precursors, nucleic acid turnover, and salvage by phosphoribosyltransferase activities. The formed urate is normally excreted by urinary and intestinal routes. Hyperuricemia can result from increased production, decreased excretion, or a combination of both mechanisms. When hyperuricemia exists, urate can precipitate and deposit in tissues as tophi.

urate is excreted by the kidneys, and most of the remainder is eliminated through the intestines.

The kidneys clear urate from the plasma and maintain physiologic balance by utilizing specific organic anion transporters (OATs), including urate transporter 1 (URAT1, SLC22A12) (Fig. 431e-2). In humans, OAT1 (SLC22A6), OAT2 (SLC22A7), and OAT3 (SLC22A8) are located on the basolateral membrane of renal proximal tubule cells. OAT4 (SLC22A11), OAT10 (SLC22A13), and URAT1 are located on the apical brush-border membrane of these cells. The latter transporters carry urate and other organic anions into the tubular cells from the lumen in exchange for intracellular organic anions. Once inside the cell, urate must pass to the basolateral side of the lumen in a process controlled by voltage-dependent carriers, including glucose transporter 9 (GLUT9, SLC2A9). *Uricosuric* compounds (Table 431e-1) directly inhibit URAT1 on the apical side of the tubular cell (so-called *cis*-inhibition). In contrast, *antiuricosuric* compounds (those that promote hyperuricemia), such as nicotinate, pyrazinoate, lactate, and other aromatic organic acids, serve as the exchange anion inside the cell, thereby stimulating anion exchange and urate reabsorption (*trans*-stimulation). The activities of URAT1, other OATs, and sodium anion transporters result in excretion of 8–12% of the filtered urate as uric acid.

Most children have serum urate concentrations of 180–240 $\mu\text{mol/L}$ (3–4 mg/dL). Levels begin to rise in males during puberty but remain low in females until menopause. The most recent mean serum urate values for men and premenopausal women in the United States are 415 and 360 $\mu\text{mol/L}$ (6.14 and 4.87 mg/dL), respectively, according to National Health and Nutrition Evaluation Survey (NHANES) data for 2007–2008. After menopause, values for women increase to approximately those for men. In adulthood, concentrations rise steadily over time and vary with height, body weight, blood pressure, renal function, and alcohol intake.

HYPERURICEMIA

Hyperuricemia can result from increased production or decreased excretion of uric acid or from a combination of the two processes. Sustained hyperuricemia predisposes some individuals to develop clinical manifestations including gouty arthritis (Chap. 395), urolithiasis, and renal dysfunction (see below).

In general, hyperuricemia is defined as a plasma (or serum) urate concentration >405 $\mu\text{mol/L}$ (>6.8 mg/dL). The risk of developing gouty arthritis or urolithiasis increases with higher urate levels and escalates in proportion to the degree of elevation. The prevalence of hyperuricemia is increasing among ambulatory adults and even more markedly among hospitalized patients. The prevalence of gout in the United States more than doubled between the 1960s and the 1990s. Based on NHANES data from 2007–2008, these trends continue, with an approximate prevalence of gout among men of 5.9% (6.1 million) and among women of 2.0% (2.2 million). Mean serum urate levels rose to 6.14 mg/dL among men and 4.87 mg/dL among women, with consequent hyperuricemia prevalences of 21.2% and 21.6%, respectively (with *hyperuricemia* defined as a serum urate level of >7.0 mg/dL [415 $\mu\text{mol/L}$] for men and >5.7 mg/dL [340 $\mu\text{mol/L}$] for women). These numbers represent a 1.2% increase in the prevalence of gout, a 0.15-mg/dL increase in the serum urate level, and a 3.2% increase in the prevalence of hyperuricemia over figures reported in NHANES-III (1988–1994). These rises are thought to be driven by increased obesity and hypertension and perhaps also by better medical care and increased longevity.

CAUSES OF HYPERURICEMIA

Hyperuricemia may be classified as primary or secondary, depending on whether the cause is innate or an acquired disorder. However, it is more useful to classify hyperuricemia in relation to the underlying pathophysiology—i.e., whether it results from increased production, decreased excretion, or a combination of the two (Fig. 431e-1, Table 431e-2).

Increased Urate Production Diet contributes to the serum urate concentration in proportion to its purine content. Strict restriction of purine intake reduces the mean serum urate level by ~60 $\mu\text{mol/L}$

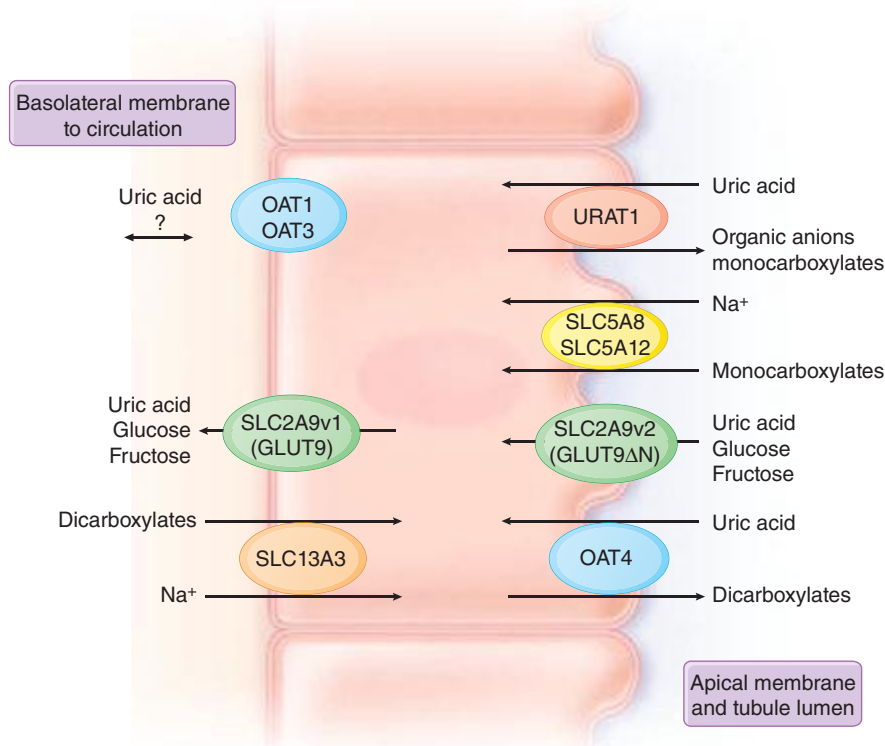


FIGURE 431e-2 Schematic for handling of uric acid by the kidney. A complex interplay of transporters on both the apical and basolateral aspects of the renal tubule epithelial cell is involved in the reabsorption of uric acid. See text for details. Most uricosuric compounds inhibit URAT1 on the apical side, as well as OAT1, OAT3, and GLUT9 on the basolateral side.

(~1 mg/dL) and urinary uric acid excretion by ~1.2 mmol/d (~200 mg/d). Foods high in nucleic acid content include liver, “sweetbreads” (i.e., thymus and pancreas), kidney, and anchovy.

Endogenous sources of purine production also influence the serum urate level (Fig. 431e-3). De novo purine biosynthesis is a multistep process that forms inosine monophosphate (IMP). The rates of purine biosynthesis and urate production are predominantly determined by amidophosphoribosyltransferase (amidoPRT), which combines phosphoribosylpyrophosphate (PRPP) and glutamine. A secondary regulatory pathway is the salvage of purine bases by hypoxanthine phosphoribosyltransferase (HPRT). HPRT catalyzes the combination of the purine bases hypoxanthine and guanine with PRPP to form the respective ribonucleotides IMP and guanosine monophosphate (GMP).

TABLE 431e-1 MEDICATIONS WITH URICOSURIC ACTIVITY

Acetohexamide	Glyceryl guaiacolate
Adrenocorticotrophic hormone	Glycopyrrolate
Ascorbic acid	Halofenate
Azauridine	Losartan
Benzbromarone	Meclofenamate
Calcitonin	Phenolsulfonphthalein
Chlorprothixene	Phenylbutazone
Citrate	Probenecid
Dicumarol	Radiographic contrast agents
Diflunisal	Salicylates (>2 g/d)
Estrogens	Sulfinpyrazone
Fenofibrate	Tetracycline that is outdated
Glucocorticoids	Zoxazolamine

Serum urate levels are closely coupled to the rates of de novo purine biosynthesis, which is driven in part by the level of PRPP, as evidenced by two X-linked inborn errors of purine metabolism (Table 431e-3). Both increased PRPP synthetase activity and HPRT deficiency are associated with overproduction of purines, hyperuricemia, and hyperuricaciduria (see below for clinical descriptions).

Accelerated purine nucleotide degradation can also cause hyperuricemia—i.e., with conditions of rapid cell turnover, proliferation, or cell death, as in leukemic blast crises, cytotoxic therapy for malignancy, hemolysis, or rhabdomyolysis. Hyperuricemia can result from excessive degradation of skeletal muscle ATP after strenuous physical exercise or status epilepticus and in glycogen storage disease types III, V, and VII (Chap. 433e). The hyperuricemia of myocardial infarction, smoke inhalation, and acute respiratory failure may also be related to accelerated breakdown of ATP.

Decreased Uric Acid Excretion More than 90% of individuals with sustained hyperuricemia have a defect in the renal handling of uric acid. For any given plasma urate concentration, patients who have gout excrete ~40% less uric acid than those who do not. When plasma urate levels are raised by purine ingestion or infusion, uric acid excretion increases in patients with and without gout; however, in those with gout, plasma urate concentrations must be 60–120 $\mu\text{mol/L}$ (1–2 mg/dL) higher than normal to achieve equivalent uric acid excretion rates.

Diminished uric acid excretion could theoretically result from decreased glomerular filtration, decreased tubular secretion, or enhanced tubular reabsorption. Decreased urate filtration does not appear to cause primary hyperuricemia but does contribute to the hyperuricemia of renal insufficiency. Although hyperuricemia is invariably present in chronic renal disease, the correlation among

TABLE 431e-2 CLASSIFICATION OF HYPERURICEMIA BY PATHOPHYSIOLOGY

Urate Overproduction		
Primary idiopathic	Myeloproliferative diseases	Rhabdomyolysis
HPRT deficiency	Polycythemia vera	Exercise
PRPP synthetase overactivity	Psoriasis	Alcohol
Hemolytic processes	Paget’s disease	Obesity
Lymphoproliferative diseases	Glycogenosis III, V, and VII	Purine-rich diet
Decreased Uric Acid Excretion		
Primary idiopathic	Starvation ketosis	Drug ingestion
Renal insufficiency	Berylliosis	Salicylates (<2 g/d)
Polycystic kidney disease	Sarcoidosis	Diuretics
Diabetes insipidus	Lead intoxication	Alcohol
Hypertension	Hyperparathyroidism	Levodopa
Acidosis	Hypothyroidism	Ethambutol
Lactic acidosis	Toxemia of pregnancy	Pyrazinamide
Diabetic ketoacidosis	Bartter’s syndrome	Nicotinic acid
	Down syndrome	Cyclosporine
Combined Mechanism		
Glucose-6-phosphatase deficiency	Fructose-1-phosphate aldolase deficiency	Alcohol
		Shock

Abbreviations: HPRT, hypoxanthine phosphoribosyltransferase; PRPP, phosphoribosylpyrophosphate.

Hyperuricemia does not necessarily represent a disease, nor is it a specific indication for therapy. The decision to treat depends on the cause and the potential consequences of hyperuricemia in each individual.

Quantification of uric acid excretion can be used to determine whether hyperuricemia is caused by overproduction or decreased excretion. On a purine-free diet, men with normal renal function excrete <3.6 mmol/d (600 mg/d). Thus, the hyperuricemia of individuals who excrete uric acid above this level while on a purine-free diet is due to purine overproduction; for those who excrete lower amounts on the purine-free diet, it is due to decreased excretion. If the assessment is performed while the patient is on a regular diet, the level of 4.2 mmol/d (800 mg/d) can be used as the discriminating value.

COMPLICATIONS

The most recognized complication of hyperuricemia is *gouty arthritis*. NHANES 2007–2008 found a prevalence of gout among U.S. adults of 3.9%, with figures of ~6% for men and ~2% for women. The higher the serum urate level, the more likely an individual is to develop gout. In one study, the incidence of gout was 4.9% among individuals with serum urate concentrations >540 $\mu\text{mol/L}$ (>9.0 mg/dL) as opposed to only 0.5% among those with values between 415 and 535 $\mu\text{mol/L}$ (7.0 and 8.9 mg/dL). The complications of gout correlate with both the duration and the severity of hyperuricemia. **For further discussion of gout, see Chap. 395.**

Hyperuricemia also causes several renal problems: (1) nephrolithiasis; (2) urate nephropathy, a rare cause of renal insufficiency attributed to monosodium urate crystal deposition in the renal interstitium; and (3) uric acid nephropathy, a reversible cause of acute renal failure resulting from deposition of large amounts of uric acid crystals in the renal collecting ducts, pelvis, and ureters.

Nephrolithiasis Uric acid nephrolithiasis occurs most commonly, but not exclusively, in individuals with gout. In gout, the prevalence of nephrolithiasis correlates with the serum and urinary uric acid levels, reaching ~50% with serum urate levels of 770 $\mu\text{mol/L}$ (13 mg/dL) or urinary uric acid excretion >6.5 mmol/d (1100 mg/d).

Uric acid stones can develop in individuals with no evidence of arthritis, only 20% of whom are hyperuricemic. Uric acid can also play a role in other types of kidney stones. Some individuals who do not have gout but have calcium oxalate or calcium phosphate stones have hyperuricemia or hyperuricaciduria. Uric acid may act as a nidus on which calcium oxalate can precipitate or lower the formation product for calcium oxalate crystallization.

Urate Nephropathy Urate nephropathy, sometimes referred to as *urate nephrosis*, is a late manifestation of severe gout and is characterized histologically by deposits of monosodium urate crystals surrounded by a giant-cell inflammatory reaction in the medullary interstitium and pyramids. The disorder is now rare and cannot be diagnosed in the absence of gouty arthritis. The lesions may be clinically silent or cause proteinuria, hypertension, and renal insufficiency.

Uric Acid Nephropathy This reversible cause of acute renal failure is due to precipitation of uric acid in renal tubules and collecting ducts that obstructs urine flow. Uric acid nephropathy develops following sudden urate overproduction and marked hyperuricaciduria. Factors that favor uric acid crystal formation include dehydration and acidosis. This form of acute renal failure occurs most often during an aggressive “blastic” phase of leukemia or lymphoma prior to or coincident with cytolytic therapy but has also been observed in individuals with other neoplasms, following epileptic seizures, and after vigorous exercise with heat stress. Autopsy studies have demonstrated intraluminal precipitates of uric acid, dilated proximal tubules, and normal glomeruli. The initial pathogenic events are believed to include obstruction of collecting ducts with uric acid and obstruction of the distal renal vasculature.

If recognized, uric acid nephropathy is potentially reversible. Appropriate therapy has reduced the mortality rate from ~50% to

practically nil. Serum levels cannot be relied on for diagnosis because this condition has developed in the presence of urate concentrations varying from 720 to 4800 $\mu\text{mol/L}$ (12–80 mg/dL). The distinctive feature is the urinary uric acid concentration. In most forms of acute renal failure with decreased urine output, urinary uric acid content is either normal or reduced, and the ratio of uric acid to creatinine is <1. In acute uric acid nephropathy, the ratio of uric acid to creatinine in a random urine sample or a 24-h specimen is >1, and a value that high is essentially diagnostic.

HYPERURICEMIA AND METABOLIC SYNDROME

Metabolic syndrome (**Chap. 422**) is characterized by abdominal obesity with visceral adiposity, impaired glucose tolerance due to insulin resistance with hyperinsulinemia, hypertriglyceridemia, increased low-density lipoprotein cholesterol, decreased high-density lipoprotein cholesterol, and hyperuricemia. Hyperinsulinemia reduces the renal excretion of uric acid and sodium. Not surprisingly, hyperuricemia resulting from euglycemic hyperinsulinemia may precede the onset of type 2 diabetes, hypertension, coronary artery disease, and gout in individuals with metabolic syndrome.

TREATMENT HYPERURICEMIA

ASYMPTOMATIC HYPERURICEMIA

Hyperuricemia is present in ~21% of the population and in at least 25% of hospitalized individuals. The vast majority of hyperuricemic persons are at no clinical risk. In the past, the association of hyperuricemia with cardiovascular disease and renal failure led to the use of urate-lowering agents for patients with asymptomatic hyperuricemia. This practice is no longer recommended except for individuals receiving cytolytic therapy for neoplastic disease, who are treated with urate-lowering agents in an effort to prevent uric acid nephropathy. Because hyperuricemia can be a component of the metabolic syndrome, its presence is an indication to screen for and aggressively treat any accompanying obesity, hyperlipidemia, diabetes mellitus, or hypertension.

Hyperuricemic individuals, especially those with higher serum urate levels, are at risk for the development of gouty arthritis. However, most hyperuricemic persons never develop gout, and prophylactic treatment is not indicated. Furthermore, neither structural kidney damage nor tophi are identifiable before the first attack. Reduced renal function cannot be attributed to asymptomatic hyperuricemia, and treatment of asymptomatic hyperuricemia does not alter the progression of renal dysfunction in patients with renal disease. An increased risk of stone formation in those with asymptomatic hyperuricemia has not been established.

Thus, because treatment with specific antihyperuricemic agents entails inconvenience, cost, and potential toxicity, routine treatment of asymptomatic hyperuricemia cannot be justified other than for prevention of acute uric acid nephropathy. In addition, routine screening for asymptomatic hyperuricemia is not recommended. If hyperuricemia is diagnosed, however, the cause should be determined. Causal factors should be corrected if the condition is secondary, and associated problems such as hypertension, hypercholesterolemia, diabetes mellitus, and obesity should be treated.

SYMPTOMATIC HYPERURICEMIA

See Chap. 395 for treatment of gout, including urate nephrosis.

Nephrolithiasis Antihyperuricemic therapy is recommended for the individual who has both gouty arthritis and either uric acid- or calcium-containing stones, both of which may occur in association with hyperuricaciduria. Regardless of the nature of the calculi, fluid ingestion should be sufficient to produce a daily urine volume >2 L. Alkalinization of the urine with sodium bicarbonate or acetazolamide may be justified to increase the solubility of uric acid. Specific treatment of uric acid calculi requires reducing the urine uric acid concentration with a xanthine oxidase inhibitor, such as allopurinol or febuxostat. These agents decrease the serum urate

concentration and the urinary excretion of uric acid in the first 24 h, with a maximal reduction within 2 weeks. Allopurinol can be given once a day because of the long half-life (18 h) of its active metabolite, oxypurinol. In the febuxostat trials, the generally recommended dose of allopurinol (300 mg/d) was effective at achieving a target serum urate concentration below 6.0 mg/dL (357 $\mu\text{mol/L}$) in <50% of patients; this result suggested that higher doses should be considered. The drug is effective in patients with renal insufficiency, but the dose should be reduced. Allopurinol is also useful in reducing the recurrence of calcium oxalate stones in patients with gout and in individuals with hyperuricemia or hyperuricaciduria who do not have gout. Febuxostat (40–80 mg/d) is also taken once daily, and doses do not need to be adjusted in the presence of mild to moderate renal dysfunction. Potassium citrate (30–80 mmol/d orally in divided doses) is an alternative therapy for patients with uric acid stones alone or mixed calcium/uric acid stones. A xanthine oxidase inhibitor is also indicated for the treatment of 2,8-dihydroxyadenine kidney stones.

Uric Acid Nephropathy Uric acid nephropathy is often preventable, and immediate appropriate therapy has greatly reduced the mortality rate. Vigorous IV hydration and diuresis with furosemide dilute the uric acid in the tubules and promote urine flow to ≥ 100 mL/h. The administration of acetazolamide (240–500 mg every 6–8 h) and sodium bicarbonate (89 mmol/L) IV enhances urine alkalinity and thereby solubilizes more uric acid. It is important to ensure that the urine pH remains >7.0 and to watch for circulatory overload. In addition, antihyperuricemic therapy in the form of allopurinol in a single dose of 8 mg/kg is administered to reduce the amount of urate that reaches the kidney. If renal insufficiency persists, subsequent daily doses should be reduced to 100–200 mg because oxypurinol, the active metabolite of allopurinol, accumulates in renal failure. Despite these measures, hemodialysis may be required. Urate oxidase (rasburicase) can also be administered IV to prevent or to treat tumor lysis syndrome.

HYPOURICEMIA

Hypouricemia, defined as a serum urate concentration <120 $\mu\text{mol/L}$ (<2.0 mg/dL), can result from decreased production of urate, increased excretion of uric acid, or a combination of both mechanisms. This condition occurs in $<0.2\%$ of the general population and $<0.8\%$ of hospitalized individuals. Hypouricemia causes no symptoms or pathology and therefore requires no therapy.

Most hypouricemia results from increased renal uric acid excretion. The finding of normal amounts of uric acid in a 24-h urine collection from an individual with hypouricemia is evidence for a renal cause. Medications with uricosuric properties (Table 431e-1) include aspirin

(at doses >2.0 g/d), losartan, fenofibrate, x-ray contrast materials, and glyceryl guaiacolate. Total parenteral hyperalimentation can also cause hypouricemia, possibly a result of the high glycine content of the infusion formula. Other causes of increased urate clearance include conditions such as neoplastic disease, hepatic cirrhosis, diabetes mellitus, and inappropriate secretion of vasopressin; defects in renal tubular transport such as primary Fanconi syndrome and Fanconi syndromes caused by Wilson's disease, cystinosis, multiple myeloma, and heavy metal toxicity; and isolated congenital defects in the bidirectional transport of uric acid. Hypouricemia can be a familial disorder that is generally inherited in an autosomal recessive manner. Most cases are caused by a loss of function mutation in *SLC22A12*, the gene that encodes URAT-1, resulting in increased renal urate clearance. Individuals with normal *SLC22A12* most likely have a defect in other urate transporters. Although hypouricemia is usually asymptomatic, some patients suffer from urate nephrolithiasis or exercise-induced renal failure.

SELECTED INBORN ERRORS OF PURINE AND PYRIMIDINE METABOLISM

(See also Table 431e-3, Table 431e-4, Fig. 431e-3, and Fig. 431e-4) More than 30 defects in human purine and pyrimidine metabolic pathways have been identified thus far. Many are benign, but about half are associated with clinical manifestations, some causing major morbidity and mortality. Advances in genetics, along with high-performance liquid chromatography and tandem mass spectrometry, have facilitated diagnosis.

PURINE DISORDERS

HPRT Deficiency The HPRT gene is located on the X chromosome. Affected males are hemizygous for the mutant gene; carrier females are asymptomatic. A complete deficiency of HPRT, the Lesch-Nyhan syndrome, is characterized by hyperuricemia, self-mutilative behavior, choreoathetosis, spasticity, and mental retardation. A partial deficiency of HPRT, the Kelley-Seegmiller syndrome, is associated with hyperuricemia but no central nervous system manifestations. In both disorders, the hyperuricemia results from urate overproduction and can cause uric acid crystalluria, nephrolithiasis, obstructive uropathy, and gouty arthritis. Early diagnosis and appropriate therapy with allopurinol can prevent or eliminate all the problems attributable to hyperuricemia without affecting behavioral or neurologic abnormalities.

Increased PRPP Synthetase Activity Like the HPRT deficiency states, PRPP synthetase overactivity is X-linked and results in gouty arthritis and uric acid nephrolithiasis. Nerve deafness occurs in some families.

Adenine Phosphoribosyltransferase (APRT) Deficiency APRT deficiency is inherited as an autosomal recessive trait. Affected individuals develop kidney stones composed of 2,8-dihydroxyadenine. Caucasians with the disorder have a complete deficiency (type I), whereas Japanese

TABLE 431e-4 INBORN ERRORS OF PYRIMIDINE METABOLISM

Enzyme	Activity	Inheritance	Clinical Features	Laboratory Features
Uridine-5'-monophosphate synthetase	Deficiency	Autosomal recessive	Orotic acid crystalluria; obstructive uropathy, hypochromic megaloblastic anemia	Orotic aciduria
Pyrimidine 5'-nucleotidase	Deficiency	Autosomal recessive	Hemolytic anemia	Basophilic stippling of erythrocytes; high levels of cytidine and uridine ribonucleotides
Pyrimidine 5'-nucleotidase	Superactivity	Uncertain	Developmental delay, seizures, ataxia, language deficit	Hypouricosuria
Thymidine phosphorylase	Deficiency	Autosomal recessive	Mitochondrial neurogastrointestinal encephalopathy	Hypouricosuria
Dihydropyrimidine dehydrogenase	Deficiency	Autosomal recessive	Seizures, motor and mental retardation	High levels of uracil, thymine, and 5-hydroxymethyluracil and low levels of dihydropyrimidines in urine
Dihydropyrimidinase	Deficiency	Uncertain	Seizures, mental retardation	Dihydropyrimidinuria
Ureidopropionase	Deficiency	Uncertain	Hypotonia, dystonia, developmental delay	High urinary excretion of <i>N</i> -carbamyl- β -alanine and <i>N</i> -carbamyl β -aminoisobutyric acid

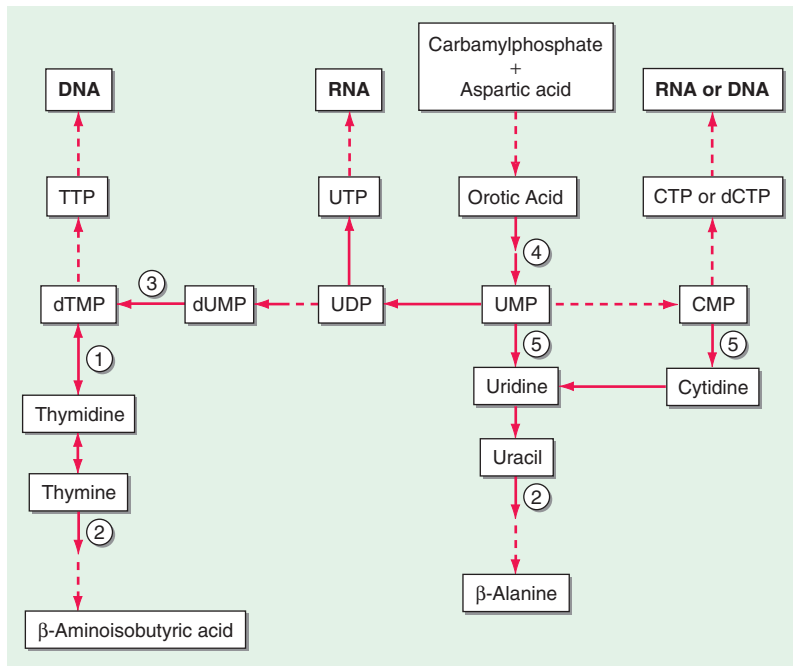


FIGURE 431e-4 Abbreviated scheme of pyrimidine metabolism. (1) Thymidine kinase, (2) dihydropyrimidine dehydrogenase, (3) thymidylate synthase, (4) UMP synthase, (5) 5'-nucleotidase. CMP, cytidine-5'-monophosphate; UMP, uridine-5'-monophosphate; UDP, uridine-5'-diphosphate; dUMP, deoxyuridine-5'-monophosphate; dTMP, deoxythymidine-5'-monophosphate; TTP, thymidine triphosphate; UTP, uridine triphosphate.

individuals have some measurable enzyme activity (type II). Expression of the defect is similar in the two populations, as is the frequency of the heterozygous state (0.4–1.1 per 100). Allopurinol treatment prevents stone formation.

Hereditary Xanthinuria A deficiency of xanthine oxidase causes all purine in the urine to occur in the form of hypoxanthine and xanthine. About two-thirds of deficient individuals are asymptomatic. The remainder develop kidney stones composed of xanthine.

Myoadenylate Deaminase Deficiency Primary (inherited) and secondary (acquired) forms of myoadenylate deaminase deficiency have been described. The primary form is inherited as an autosomal recessive trait. Clinically, some persons may have relatively mild myopathic symptoms with exercise or other triggers, but most individuals with this defect are asymptomatic. Therefore, another explanation for the myopathy should be sought in symptomatic patients with this deficiency. The acquired deficiency occurs in association with a wide variety of neuromuscular diseases, including muscular dystrophies, neuropathies, inflammatory myopathies, and collagen vascular diseases.

Adenylosuccinate Lyase Deficiency Deficiency of this enzyme is due to an autosomal recessive trait and causes profound psychomotor retardation, seizures, and other movement disorders. All individuals with this deficiency are mentally retarded, and most are autistic.

Adenosine Deaminase Deficiency and Purine Nucleoside Phosphorylase Deficiency See Chap. 374.

PYRIMIDINE DISORDERS

The pyrimidine cytosine is found in both DNA and RNA; it is a complementary base pair for guanine. Thymidine is found only in DNA,

where it is paired with adenine. Uridine is found only in RNA and can pair with either adenine or guanine in RNA secondary structures. Pyrimidines can be synthesized by a de novo pathway (Fig. 431e-4) or reused in a salvage pathway. Although more than 25 different enzymes are involved in pyrimidine metabolism, disorders of these pathways are rare. Seven disorders of pyrimidine metabolism have been discovered (Table 431e-4), three of which are discussed below.

Orotic Aciduria Hereditary orotic aciduria is caused by mutations in a bifunctional enzyme, uridine-5'-monophosphate (UMP) synthase, which converts orotic acid to UMP in the de novo synthesis pathway (Fig. 431e-4). The disorder is characterized by hypochromic megaloblastic anemia that is unresponsive to vitamin B₁₂ and folic acid, growth retardation, and neurologic abnormalities. Increased excretion of orotic acid causes crystalluria and obstructive uropathy. Replacement of uridine (100–200 mg/kg per day) corrects anemia, reduces orotic acid excretion, and improves the other sequelae of the disorder.

Pyrimidine 5'-nucleotidase Deficiency Pyrimidine 5'-nucleotidase catalyzes the removal of the phosphate group from pyrimidine ribonucleoside monophosphates (cytidine-5'-monophosphate or UMP) (Fig. 431e-4). An inherited deficiency of this enzyme causes hemolytic anemia with prominent basophilic stippling of erythrocytes. The accumulation of pyrimidines or cytidine diphosphate choline is thought to induce hemolysis. There is no specific treatment. Acquired pyrimidine 5'-nucleotidase deficiency has been reported in lead poisoning and in thalassemia.

Dihydropyrimidine Dehydrogenase Deficiency Dihydropyrimidine dehydrogenase is the rate-limiting enzyme in the pathway of uracil and thymine degradation (Fig. 431e-4). Deficiency of this enzyme causes excessive urinary excretion of uracil and thymine. In addition, this deficiency causes nonspecific cerebral dysfunction with convulsive disorders, motor retardation, and mental retardation. No specific treatment is available.

Medication Effect on Pyrimidine Metabolism A variety of medications can influence pyrimidine metabolism. The anticancer agents fluorodeoxyuridine and 5-fluorouracil and the antimicrobial agent fluorocytosine cause cytotoxicity when converted to fluorodeoxyuridylate, a specific suicide inhibitor of thymidylate synthase. Fluorocytosine must be converted to 5-fluorouracil to be effective. This conversion is catalyzed by cytosine deaminase activity. Fluorocytosine's action is selective because cytosine deaminase is present in bacteria and fungi but not in human cells. Dihydropyrimidine dehydrogenase is involved in the degradation of 5-fluorouracil. Consequently, deficiency of this enzyme is associated with 5-fluorouracil neurotoxicity.

Leflunomide, which is used to treat rheumatoid arthritis, inhibits de novo pyrimidine synthesis by inhibiting dihydroorotate dehydrogenase, resulting in an antiproliferative effect on T cells. Allopurinol, which inhibits xanthine oxidase in the purine metabolic pathway, also inhibits the activity of orotidine-5'-phosphate decarboxylase, a step in UMP synthesis. Consequently, allopurinol use is associated with increased excretion of orotidine and orotic acid. There are no known clinical effects of this inhibition.

Lysosomes are heterogeneous subcellular organelles containing specific hydrolases that allow selective processing or degradation of proteins, nucleic acids, carbohydrates, and lipids. There are more than 40 different lysosomal storage diseases (LSDs), classified according to the nature of the stored material (Table 432e-1). Several of the most prevalent disorders are reviewed here: Tay-Sachs disease, Fabry disease, Gaucher disease, Niemann-Pick disease, lysosomal acid lipase deficiencies, the mucopolysaccharidoses, and Pompe disease. LSDs should be considered in the differential diagnosis of patients with neurologic, renal, or muscular degeneration and/or unexplained hepatomegaly, splenomegaly, cardiomyopathy, or skeletal dysplasias and deformations. Physical findings are disease specific, and enzyme assays or genetic testing can be used to make a definitive diagnosis. Although the nosology of LSDs segregates the variants into distinct phenotypes, these are heuristic; in the clinic, each disease exhibits—to some degree—a continuous spectrum of manifestations, from severe to attenuated variants.

PATHOGENESIS

Lysosomal biogenesis involves ongoing synthesis of lysosomal hydrolases, membrane constitutive proteins, and new membranes. Lysosomes originate from the fusion of trans-Golgi network vesicles with late endosomes. Progressive vesicular acidification accompanies the maturation of these vesicles; this gradient facilitates the pH-dependent dissociation of receptors and ligands and also activates lysosomal hydrolases. Lysosomes are components of the lysosome/autophagy/mitophagy system, which can be disrupted in the LSDs.

Abnormalities at any biosynthetic step can impair enzyme activation and lead to a lysosomal storage disorder. After leader sequence clipping, remodeling of complex oligosaccharides (including the lysosomal targeting ligand mannose-6-phosphate as well as high-mannose oligosaccharide chains of many soluble lysosomal hydrolases) occurs during transit through the Golgi. Lysosomal integral or associated membrane proteins are sorted to the membrane or interior of the lysosome by several different peptide signals. Phosphorylation, sulfation, additional proteolytic processing, and macromolecular assembly of heteromers occur concurrently. Such posttranslational modifications are critical to enzyme function, and defects can result in multiple enzyme/protein deficiencies.

The common pathway for LSDs is the accumulation of specific macromolecules within tissues and cells that normally have a high flux of these substrates. The majority of lysosomal enzyme deficiencies result from point mutations or genetic rearrangements at a locus that encodes a single lysosomal hydrolase. However, some mutations cause deficiencies of several different lysosomal hydrolases by alteration of the enzymes/proteins involved in targeting, active site modifications, or macromolecular association or trafficking. All LSDs are inherited as autosomal recessive disorders except for Hunter (mucopolysaccharidosis type II), Danon, and Fabry diseases, which are X-linked. Substrate accumulation leads to lysosomal distortion, which has significant pathologic consequences. In addition, abnormal amounts of metabolites may also have pharmacologic effects important to disease pathophysiology and propagation.

For many LSDs, the accumulated substrates are endogenously synthesized within particular tissue sites of pathology. Other diseases have greater exogenous substrate supplies. For example, they are delivered by low-density lipoprotein receptor-mediated uptake in Fabry and cholesteryl ester storage diseases (CESDs) or by phagocytosis in Gaucher disease type 1. The *threshold hypothesis* refers to a level of enzyme activity below which disease develops; small changes in enzyme activity near the threshold can lead to or prevent disease. A critical element of this model is that enzymatic activity can be challenged by changes in substrate flux based on genetic background, cell

turnover, recycling, or metabolic demands. Thus, a set level of residual enzyme may be adequate for substrate in some tissues or cells but not in others. In addition, several variants of each LSD exist at a clinical level. These disorders therefore represent a continuum of manifestations that are not easily dissociated into discrete entities. The bases for such variations have not been elucidated in any detail.

SELECTED DISORDERS

TAY-SACHS DISEASE

About 1 in 30 Ashkenazi Jews is a carrier for Tay-Sachs disease, which is caused by total hexosaminidase A (Hex A) deficiency resulting from defective α -chains. The infantile form is a fatal neurodegenerative disease with macrocephaly, loss of motor skills, increased startle reaction, and a macular cherry red spot. The juvenile-onset form presents as ataxia and dementia, with death by age 10–15 years. The adult-onset disorder is characterized by clumsiness in childhood; progressive motor weakness in adolescence; and additional spinocerebellar and lower-motor-neuron symptoms and dysarthria in adulthood. Intelligence declines slowly, and psychosis is common. Screening for Tay-Sachs disease carriers is recommended in the Ashkenazi Jewish population. Sandhoff disease, due to a deficiency in both Hex A and Hex B resulting from defective β -chains, is phenotypically similar to Tay-Sachs disease but also includes hepatosplenomegaly and bony dysplasias.

FABRY DISEASE

Fabry disease is an X-linked disorder that results from mutations in the α -galactosidase A gene. The estimated prevalence of hemizygous males ranges from 1 in 40,000 to 1 in 3500 in selected populations. Clinically, the disease manifests with angiokeratomas (telangiectatic skin lesions), hypohidrosis, corneal and lenticular opacities, acroparesthesia; and progressive small-vessel disease of the kidney, heart, and brain.

The angiokeratomas and acroparesthesias may appear in childhood. Angiokeratomas are punctate, dark red to blue-black, flat or slightly raised, and usually symmetric; they do not blanch with pressure. They range from barely visible to several millimeters in diameter and have a tendency to increase in size and number with age. They usually are most dense between the umbilicus and the knees—the “bathing suit area”—but may occur anywhere, including the mucosal surfaces. Angiokeratomas also occur in several other very rare LSDs. Corneal and lenticular lesions, detectable on slit-lamp examination, may help in establishing a diagnosis of Fabry disease. Debilitating episodic burning pain of the hands, feet, and proximal extremities (acroparesthesia) can last from minutes to days and can be precipitated by changes in temperature, exercise, fatigue, or fever. Abdominal pain can resemble that from appendicitis or renal colic. Proteinuria, isosthenuria, and progressive renal dysfunction occur in the second to fourth decades; ~5% of male patients with idiopathic renal failure have α -galactosidase A mutations. Hypertension, left ventricular hypertrophy, anginal chest pain, and congestive heart failure can occur in the third to fourth decades. About 1–3% of patients with idiopathic hypertrophic cardiomyopathy have Fabry disease. Similarly, ~3–5% of male patients with idiopathic stroke at 35–50 years of age have α -galactosidase A mutations. Leg lymphedema without hypoproteinemia and episodic diarrhea also occur. Death is due to renal failure or cardiovascular or cerebrovascular disease in untreated male patients. Variants with residual α -galactosidase A activity may have late-onset manifestations that are limited to the cardiovascular system and resemble hypertrophic cardiomyopathy. Variants with predominant cardiac, renal, or central nervous system (CNS) manifestations are becoming better defined. Up to 70% of heterozygous females may exhibit clinical manifestations. However, in females, heart disease is the most common life-threatening manifestation, followed in frequency by stroke and then renal disease.

Phenytoin and carbamazepine diminish chronic and episodic acroparesthesia. Chronic hemodialysis or kidney transplantation can be lifesaving in patients with renal failure. Enzyme therapy clears stored lipids from a variety of cells, particularly those of the renal, cardiac, and skin vascular endothelium. Renal insufficiency appears

TABLE 432e-1 SELECTED LYSOSOMAL STORAGE DISEASES

Disorder ^a	Enzyme Deficiency [Specific Therapy]	Stored Material	Clinical Types (Onset)	Inheritance	Clinical Features					
					Neurologic	Liver, Spleen Enlargement	Skeletal Dysplasia	Ophthalmologic	Hematologic	Unique Features
Mucopolysaccharidoses CTMucopolysaccharidoses (MPS)										
MPS I H, Hurler (136)	α -L-Iduronidase [ET, BMT]	Dermatan sulfate Heparan sulfate	Infantile Intermediate	AR	Mental retardation	+++	++++	Corneal clouding	Vacuolated lymphocytes	Coarse facies; cardiovascular involvement; joint stiffness
MPS I H/S, Hurler/ Scheie			Adult		Mental retardation					
MPS I S, Scheie					None					
MPS II, Hunter (136)	Iduronate sulfatase [ET]	Dermatan sulfate Heparan sulfate	Severe infantile Mild juvenile	X-linked	Mental retardation, less in mild form	+++	++++	Retinal degeneration, no corneal clouding	Granulated lymphocytes	Coarse facies; cardiovascular involvement; joint stiffness; distinctive pebbly skin lesions
MPS III A, Sanfilippo A (136)	Heparan-N-sulfatase	Heparan sulfate	Late infantile	AR	Severe mental retardation	+	+	None	Granulated lymphocytes	Mild coarse facies
MPS III B, Sanfilippo B (136)	N-Acetyl- α -glucosaminidase	Heparan sulfate	Late infantile	AR	Severe mental retardation	+	+	None	Granulated lymphocytes	Mild coarse facies
MPS III C, Sanfilippo C (136)	Acetyl-CoA: α -glucosaminide N-acetyltransferase	Heparan sulfate	Late infantile	AR	Severe mental retardation	+	+	None	Granulated lymphocytes	Mild coarse facies
MPS III D, Sanfilippo D (136)	N-Acetylglucosamine- 6-sulfate sulfatase	Heparan sulfate	Late infantile	AR	Severe mental retardation	+	+	None	Granulated lymphocytes	Mild coarse facies
MPS IV A, Morquio A (136)	N-Acetylgalactosamine- 6-sulfate sulfatase [ET—trials]	Keratan sulfate Chondroitin-6 sulfate	Childhood	AR	None	+	++++	Corneal clouding	Granulated neutrophils	Distinctive skeletal deformity; odontoid hypoplasia; aortic valve disease
MPS IV B, Morquio (136)	β -Galactosidase		Childhood	AR	None	\pm	++++			
MPS VI, Maroteaux- Lamy (136)	Arylsulfatase B [ET, BMT]	Dermatan sulfate	Late infantile	AR	None	++	++++	Corneal clouding	Granulated neutrophils and lymphocytes	Coarse facies; valvular heart disease
MPS VII (136)	β -Glucuronidase	Dermatan sulfate Heparan sulfate	Neonatal Infantile Adult	AR	Mental retardation, absent in some adults	+++	+++	Corneal clouding	Granulated neutrophils	Coarse facies; vascular involvement; hydrops fetalis in neonatal form

GM₂ Gangliosidoses										
Tay-Sachs disease (153)	β-Hexosaminidase A	GM ₂ gangliosides	Infantile Juvenile	AR	Mental retardation; seizures; later juvenile form	None	None	Cherry red spot in infantile form	None	Macrocephaly; hyperacusis in infantile form
Sandhoff disease (153)	β-Hexosaminidases A and B	GM ₂ gangliosides	Infantile	AR	Mental retardation; seizures	++	±	Cherry red spot	None	Macrocephaly; hyperacusis
Neutral Glycosphingolipidoses										
Fabry disease (150)	α-Galactosidase A [ET]	Globotriaosylceramide	Childhood	X-linked	Painful acroparesthesias	None	None	Corneal dystrophy, vascular lesions	None	Cutaneous angiokeratomas; hypo-hydrosis
Gaucher disease (146)	Acid β-glucosidase [ET, SRT]	Glucosylceramide	Type 1 Type 2 Type 3	AR	None	++++ ++++ ++	++++ + ++++	None Eye movements Eye movements	Gaucher cells in bone marrow; cytopenias	Adult form highly variable
Niemann-Pick disease (144) A and B	Sphingomyelinase [ET—trials]	Sphingomyelin	Neuronopathic, type A Nonneuronopathic, type B	AR	Mental retardation; seizures	++++	None Osteoporosis	Macular degeneration	Foam cells in bone marrow	Pulmonary infiltrates Lung failure
Glycoproteinoses										
Fucosidosis (140)	α-Fucosidase	Glycopeptides; oligosaccharides	Infantile Juvenile	AR	Mental retardation	++	++	None	Vacuolated lymphocytes; foam cells	Coarse facies; angiokeratomas in juvenile form
α-Mannosidosis (140)	α-Mannosidase	Oligosaccharides	Infantile Milder variant	AR	Mental retardation	+++	+++	Cataracts, corneal clouding	Vacuolated lymphocytes, granulated neutrophils	Coarse facies; enlarged tongue
β-Mannosidosis (140)	β-Mannosidase	Oligosaccharides		AR	Seizures; mental retardation		++	None	Vacuolated lymphocytes, foam cells	Angiokeratomas
Aspartylglucosaminuria (141)	Aspartylglucosaminidase	Aspartylglucosamine; glycopeptides	Young adult	AR	Mental retardation	±	++	None	Vacuolated lymphocytes, foam cells	Coarse facies
Sialidosis (140)	Neuraminidase	Sialyloligosaccharides	Type I, congenital Type II, infantile and juvenile	AR	Myoclonus; mental retardation	++, less in type I	++, less in type I	Cherry red spot	Vacuolated lymphocytes	MPS phenotype in type II
Mucopolysaccharidoses (ML)										
ML-II, I-cell disease (138)	UDP-N- Acetylglucosamine- 1-phosphotransferase	Glycoprotein; glycolipids	Infantile	AR	Mental retardation	+	++++	Corneal clouding	Vacuolated and granulated neutrophils	Coarse facies; absence of mucopolysac- chariduria; gingival hypoplasia
ML-III, pseudo-Hurler polydystrophy (138)	UDP-N- Acetylglucosamine- 1-phosphotransferase	Glycoprotein; glycolipids	Late infantile	AR	Mild mental retardation	None	+++	Corneal clouding, mild retinopathy, hyperopic astigmatism		Coarse facies; stiffness of hands and shoulders

(Continued)

TABLE 432e-1 SELECTED LYSOSOMAL STORAGE DISEASES (CONTINUED)

Disorder ^a	Enzyme Deficiency [Specific Therapy]	Stored Material	Clinical Types (Onset)	Inheritance	Clinical Features					
					Neurologic	Liver, Spleen Enlargement	Skeletal Dysplasia	Ophthalmologic	Hematologic	Unique Features
Leukodystrophies										
Krabbe disease (147)	Galactosylceramidase [BMT/HSCT]	Galactosylceramide Galactosylsphingosine	Infantile	AR	Mental retardation	None	None	None	None	White matter globoid cells
Metachromatic leukodystrophy (148)	Arylsulfatase A	Cerebroside sulfate	Infantile Juvenile Adult	AR	Mental retardation; dementia; psychosis in adult	None	None	Optic atrophy	None	Gait abnormalities in late infantile form
Multiple sulfatase deficiency (149)	Active site cysteine to C ₆ -formylglycine- converting enzyme	Sulfatides; mucopoly- saccharides	Late infantile	AR	Mental retardation	+	++	Retinal degeneration	Vacuolated and granu- lated cells	Absent activity of all known cellular sulfatases
Disorders of Neutral Lipids										
Wolman disease (142)	Acid lysosomal lipase [ET—trials]	Cholesteryl esters; triglycerides	Infantile	AR	Mild mental retardation	+++	None	None	None	Adrenal calcification
Cholesteryl ester storage disease (142)	Acid lysosomal lipase [ET—trials]	Cholesteryl esters	Childhood	AR	None	Hepatomegaly	None	None	None	Fatty liver disease; cirrhosis
Farber disease (143)	Acid ceramidase	Ceramide	Infantile Juvenile	AR	Occasional mental retardation	±	None	Macular degeneration	None	Arthropathy, subcutaneous nodules
Disorders of Glycogen										
Pompe disease (135)	Acid α -glucosidase [ET]	Glycogen	Infantile, late onset	AR	Neuromuscular	±	None	None	None	Myocardiopathy
Late -onset GAA deficiency (135)	Acid α -glucosidase [ET]	Glycogen	Variable: juvenile to adulthood	AR	Neuromuscular	None	None	None	None	Respiratory insufficiency; neuromuscular disease
Danon disease (154)	LAMP-2 (lysosomal associated membrane protein-2)	Glycogen	Variable: childhood to adulthood	X-linked (?Dominant)	Cardiomyopathy Neuromuscular Inconsistent mental retardation	None	None	None	None	Myocardial vacuolar degeneration

^aNumbers in parentheses refer to the chapters in CR Scriver et al: *The Metabolic and Molecular Bases of Inherited Disease*, 9th ed. New York, McGraw-Hill, www.ommbid.com, which provide comprehensive reviews.

Abbreviations: AR, autosomal recessive; BMT/HSCT, bone marrow or stem cell transplantation; ET, enzyme therapy; SRT, substrate reduction therapy.

to be irreversible. Early institution of enzyme therapy may prevent or slow the progression of life-threatening complications.

GAUCHER DISEASE

Gaucher disease is an autosomal recessive disorder that results from defective activity of acid β -glucosidase; ~400 mutations have been described at the *GBA1* locus of such patients. Disease variants are classified by the absence or presence and progression of neuronopathic involvement.

Gaucher disease type 1 is a nonneuronopathic disease that can present in childhood to adulthood as slowly to rapidly progressive visceral disease. About 55–60% of patients are diagnosed at <20 years of age in white populations and at even younger ages in other groups. This pattern of presentation is distinctly bimodal, with peaks at <10–15 years and at ~25 years. Younger patients tend to have a greater degree of hepatosplenomegaly and accompanying blood cytopenias. In contrast, the older group has a greater tendency for chronic bone disease. Hepatosplenomegaly occurs in virtually all symptomatic patients and can be minor or massive. Accompanying anemia and thrombocytopenia are variable and are not directly related to liver or spleen volumes. Severe liver dysfunction is unusual. Splenic infarctions can resemble an acute abdomen. Pulmonary hypertension and alveolar Gaucher cell accumulation are uncommon but life-threatening and can occur at any age. *GBA1* mutations in the hetero- or homozygous state are a significant risk factor for early-onset or more rapidly progressive Parkinson disease.

All patients with Gaucher disease have nonuniform infiltration of bone marrow by lipid-laden macrophages termed *Gaucher cells*. This phenomenon can lead to marrow packing with subsequent infarction, ischemia, necrosis, and cortical bone destruction. Bone marrow involvement spreads from proximal to distal in the limbs and can involve the axial skeleton extensively, causing vertebral collapse. In addition to bone marrow involvement, bone remodeling is defective, with loss of total bone calcium leading to osteopenia, osteonecrosis, avascular infarction, and vertebral compression fractures and spinal cord involvement. Aseptic necrosis of the femoral head is common, as is fracture of the femoral neck. The mechanism by which diseased bone marrow macrophages interact with osteoclasts and/or osteoblasts to cause bone disease is not well understood. Chronic, ill-defined bone pain can be debilitating and poorly correlated with radiographic findings. “Bone crises” are associated with localized excruciating pain and, on occasion, local erythema, fever, and leukocytosis. These crises represent acute infarctions of bone, as evidenced in nuclear scans by localized absent uptake of pyrophosphate agents. Decreased acid β -glucosidase activity (0–20% of normal) in nucleated cells establishes the diagnosis. The enzyme is not present in bodily fluids. The sensitivity of enzyme testing is poor for heterozygote detection; molecular testing by *GBA1* sequencing is preferred. The disease frequency varies from about 1 in 1000 among Ashkenazi Jews to <1 in 100,000 in other populations; ~1 in 12–15 Ashkenazi Jews carries a Gaucher disease allele. Four common mutations account for ~85% of the mutations in that population of affected patients: N370S (1226G), 84GG (a G insertion at cDNA position 84), L444P (1448C), and IVS-2 (an intron 2 splice junction mutation).

Genotype/phenotype studies indicate a significant, though not absolute, correlation between disease type and severity and the *GBA1* genotype. The most common mutation in the Ashkenazi Jewish population (N370S) shares a 100% association with nonneuronopathic or type 1 Gaucher disease. The N370S/N370S and N370S/other mutant allele genotypes are associated with later-onset/less severe disease and with earlier-onset/severe disease, respectively. As many as 50–60% of individuals with the N370S/N370S genotype are asymptomatic. Other alleles include L444P (very low activity), 84GG (null), or IVS-2 (null) and rare/private or uncharacterized alleles. The L444P/L444P patients almost always have life-threatening to very severe/early-onset disease, and many, though not all, develop CNS involvement in the first two decades of life.

Symptom-based treatment of blood cytopenias and joint replacement surgeries continue to have important roles in management.

However, regular intravenous enzyme therapy is currently the treatment of choice in significantly affected patients and is highly efficacious and safe in diminishing hepatosplenomegaly and improving hematologic values. Bone disease is decreased but not eliminated by enzyme therapy. Adult patients may benefit from adjunctive treatment with bisphosphonates to improve bone density. Patients who cannot be treated with enzyme, either because it is not effective or because they have an allergy or other hypersensitivities, may receive *substrate reduction therapy* with medications that decrease the production of the complex lipid molecules that are broken down by acid β -glucosidase.

Gaucher disease type 2 is a rare, severe, progressive CNS disease that leads to death by 2 years of age. Gaucher disease type 3 has highly variable manifestations in the CNS and viscera. It can present in early childhood with rapidly progressive, massive visceral disease and slowly progress to static CNS involvement; in adolescence with dementia; or in early adulthood with rapidly progressive, uncontrollable myoclonic seizures and mild visceral disease. Visceral disease in type 3 is nearly identical to that in type 1 but is generally more severe. Early CNS findings may be limited to defects in lateral gaze tracking, which may remain static for decades. Mental retardation can be slowly progressive or static. This variant is most frequent among individuals of Swedish descent. Visceral—but not CNS—involvement responds to enzyme therapy.

NIEMANN-PICK DISEASE

Niemann-Pick diseases are autosomal recessive disorders that result from defects in acid sphingomyelinase. Types A and B are distinguished by the early age of onset and progressive CNS disease in type A. Type A typically has its onset in the first 6 months of life, with rapidly progressive CNS deterioration, spasticity, failure to thrive, and massive hepatosplenomegaly. Type B has a later, more variable onset and is characterized by a progression of hepatosplenomegaly, with eventual development of cirrhosis and hepatic replacement by foam cells. Affected patients develop progressive pulmonary disease with dyspnea, hypoxemia, and a reticular infiltrative pattern on chest x-ray. Foam cells are present in alveoli, lymphatic vessels, and pulmonary arteries. Progressive hepatic or lung disease leads to death in adolescence or early adulthood.

The diagnosis is established by markedly decreased (1–10% of normal) sphingomyelinase activity in nucleated cells. There is no specific treatment for Niemann-Pick disease. The efficacy of hepatic or bone marrow transplantation has not been clearly established. Clinical trials of enzyme therapy are in phases 2 and 3.

Niemann-Pick C diseases are progressive CNS diseases due to mutations in either NPC1 or NPC2. They present with liver or splenic disease, but their major manifestations are progressive CNS disease over one to two decades. Treatment with substrate inhibition agents (e.g., Miglustat) and substrate depletion with cyclodextrin have shown promise.

LYSOSOMAL ACID LIPASE

Lysosomal acid lipase deficiency may result in Wolman disease (severe deficiency) or CESD, which presents later and has some (3–10%) residual enzyme activity. Wolman disease presents in early infancy with hepatosplenomegaly, diarrhea, vomiting, and abdominal distention, sometimes accompanied by adrenal calcification, anemia, and mixed hyperlipidemia. Death occurs before the age of 1 year and is often due to severe intestinal malabsorption. CESD is heterogeneous and presents with hepatomegaly and hepatosteatosis at any age in childhood or adulthood. CESD should be included in the differential diagnosis for all patients with isolated hypercholesterolemia. The disease may progress to hepatic fibrosis, cirrhosis, and liver failure. In addition, patients often develop very early-onset atherosclerotic vascular disease, which may be life-threatening in childhood. Preliminary results of treatment with enzyme replacement therapy are promising for Wolman disease and CESD.

MUCOPOLYSACCHARIDOSES

Mucopolysaccharidosis type I (MPS I) is an autosomal recessive disorder caused by deficiency of α -L-iduronidase. The continuum of

432e-6 involvement traditionally has been divided into three categories: (1) Hurler disease (MPS I H) for severe deficiency with neurodegeneration, (2) Scheie disease (MPS I S) for later-onset disease without neurologic involvement and with relatively less severe disease in other organ systems, and (3) Hurler-Scheie syndrome (MPS I H/S) for patients intermediate between these extremes. MPS I H/S is characterized by severe somatic disease, usually without overt neurologic deterioration.

MPS I often presents in infancy or early childhood as chronic rhinitis, clouding of the corneas, and hepatosplenomegaly. As the disease progresses, nearly every organ system can be affected. In the more severe forms, cardiac and respiratory diseases become life-threatening in childhood. Skeletal disease can be quite severe, resulting in very limited mobility.

There are two current treatments for the MPS I diseases. Hematopoietic stem cell transplantation (HSCT) is the standard treatment for patients presenting at <2 years of age who appear to have or are at risk for neurologic degeneration. HSCT results in stabilization of CNS disease and reverses hepatosplenomegaly. It also beneficially affects cardiac and respiratory disease. HSCT does not eliminate corneal disease or result in the resolution of progressive skeletal disease. Enzyme therapy effectively addresses hepatosplenomegaly and alleviates cardiac and respiratory disease. The enzyme does not effectively penetrate the CNS and does not directly affect CNS disease. Enzyme therapy and HSCT appear to have similar effects on visceral signs and symptoms. Enzyme therapy poses a lower risk of life-threatening complications and may therefore be advantageous for patients who have attenuated manifestations without CNS disease. A combination of enzyme therapy and HSCT has been used, with enzyme therapy initiated prior to transplantation in an attempt to reduce the disease burden. The experience with this approach is not well documented, but it appears to have advantages over HSCT alone.

Enzyme therapy for Maroteaux-Lamy disease (MPS VI) has received U.S. Food and Drug Administration (FDA) approval. This very rare autosomal recessive disorder is characterized by hepatosplenomegaly, bone disease, heart disease, and respiratory compromise similar to

those seen in MPS I; however, MPS VI is due to deficiency of arylsulfatase B and is not associated with neurologic degeneration.

Hunter disease (MPS II) is an X-linked disorder due to deficiency in iduronate sulfate sulfatase and has manifestations similar to those of MPS I, including neurologic degeneration. There is no corneal clouding or other eye disease. Like MPS I, MPS II is clinically variable, with CNS and non-CNS variants. HSCT has not been successful in treating CNS disease associated with MPS II. The FDA and the European Medicines Agency (EMA) have approved enzyme therapy for the visceral manifestations of MPS II.

POMPE DISEASE

Acid maltase (acid α -glucosidase, GAA) deficiency, also called Pompe disease, is the only glycogen LSD. The classic severe infantile form presents with hypotonia, cardiomyopathy, and hepatosplenomegaly. This variant is rapidly progressive and generally results in death in the first year of life. However, as with other LSDs, there are early- and late-onset forms of this disorder. The late-onset variants may be as common as 1 in 40,000; patients typically present with a slowly progressive myopathy that may resemble limb-girdle muscular dystrophy. Respiratory insufficiency may be the presenting sign or may develop with advancing disease. In late stages of the disease, patients may require mechanical ventilation, report swallowing difficulties, and experience loss of bowel and bladder control. Myocardopathy is not usually seen in late-onset variants of Pompe disease.

The FDA and EMA have approved enzyme therapy for Pompe disease. This treatment clearly prolongs life in the infantile form, consistently resulting in improved cardiac function. Respiratory function is also improved in most treated infants. Some infants demonstrate marked improvement in motor functions, while others have minor changes in muscle tone or strength. Prevention of deterioration has been shown with GAA enzyme therapy in the late-onset forms. Early intervention with GAA enzyme therapy in such patients may limit or prevent deterioration, but very advanced disease will have significant irreversible components.

Priya S. Kishnani, Yuan-Tsong Chen

Carbohydrate metabolism plays a vital role in cellular function by providing the energy required for most metabolic processes. The relevant biochemical pathways involved in the metabolism of these carbohydrates are shown in **Fig. 433e-1**. Glucose is the principal substrate of energy metabolism in humans. Metabolism of glucose generates ATP through glycolysis and mitochondrial oxidative phosphorylation. The body obtains glucose through the ingestion of polysaccharides (primarily starch) and disaccharides (e.g., lactose, maltose, and sucrose). Galactose and fructose are two other monosaccharides that serve as sources of fuel for cellular metabolism; however, their role as fuel sources is much less significant than that of glucose. Galactose is derived from lactose (galactose + glucose), which is found in milk

products, and is an important component of certain glycolipids, glycoproteins, and glycosaminoglycans. Fructose is found in fruits, vegetables, and honey. Sucrose (fructose + glucose) is another dietary source of fructose and is a commonly used sweetener.

Glycogen, the storage form of glucose in animal cells, is composed of glucose residues joined in straight chains by α 1-4 linkages and branched at intervals of 4–10 residues by α 1-6 linkages. Glycogen forms a treelike molecule and can have a molecular weight of many millions. Glycogen may aggregate to form structures recognizable by electron microscopy. With the exception of type 0 disease, defects in glycogen metabolism typically cause an accumulation of glycogen in the tissues—hence the designation *glycogen storage diseases* (GSDs). The structure of stored glycogen can be normal or abnormal in the various disorders. Defects in gluconeogenesis or glycolytic pathways, including galactose and fructose metabolism, usually do not result in glycogen accumulation.

Clinical manifestations of the various disorders of carbohydrate metabolism differ markedly. The symptoms range from harmless to lethal. Unlike disorders of lipid metabolism, mucopolysaccharidoses, or other storage diseases, many carbohydrate disorders have been effectively managed with dietary therapy. All of the genes responsible for inherited defects of carbohydrate metabolism have been cloned, and mutations have been identified. Advances in our understanding of the molecular basis of these diseases are being used to improve diagnosis and management. Some of these disorders are candidates for enzyme replacement therapy, substrate reduction therapy, and early trials of gene therapy.

Historically, the GSDs were categorized numerically in the order in which the enzymatic defects were identified. They are also classified by the organs involved (liver, muscle, and/or heart) and clinical manifestations. The latter is the system followed in this chapter (**Table 433e-1**). The overall frequency of all forms of GSD is ~1 in 20,000 live births. Most are inherited as autosomal recessive traits; however, phosphoglycerate kinase deficiency—one form of liver phosphorylase kinase (PhK) deficiency—and lysosomal-associated membrane protein 2 (LAMP2) deficiency are X-linked disorders. The most common childhood disorders are glucose-6-phosphatase deficiency (type I), lysosomal acid α -glucosidase deficiency (type II), debrancher deficiency (type III), and liver PhK deficiency (type IX). The most common adult disorder is myophosphorylase deficiency (type V, or McArdle disease).

The overall frequency of all forms of GSD is ~1 in 20,000 live births. Most are inherited as autosomal recessive traits; however, phosphoglycerate kinase deficiency—one form of liver phosphorylase kinase (PhK) deficiency—and lysosomal-associated membrane protein 2 (LAMP2) deficiency are X-linked disorders. The most common childhood disorders are glucose-6-phosphatase deficiency (type I), lysosomal acid α -glucosidase deficiency (type II), debrancher deficiency (type III), and liver PhK deficiency (type IX). The most common adult disorder is myophosphorylase deficiency (type V, or McArdle disease).

SELECTED LIVER GLYCOGENOSES

DISORDERS WITH HEPATOMEGALY AND HYPOGLYCEMIA

Type I GSD (Glucose-6-Phosphatase or Translocase Deficiency, Von Gierke's Disease)

Type I GSD is an autosomal recessive disorder caused by glucose-6-phosphatase deficiency in liver, kidney, and intestinal mucosa. There are two subtypes of GSD I: type Ia, in which the glucose-6-phosphatase enzyme is defective, and

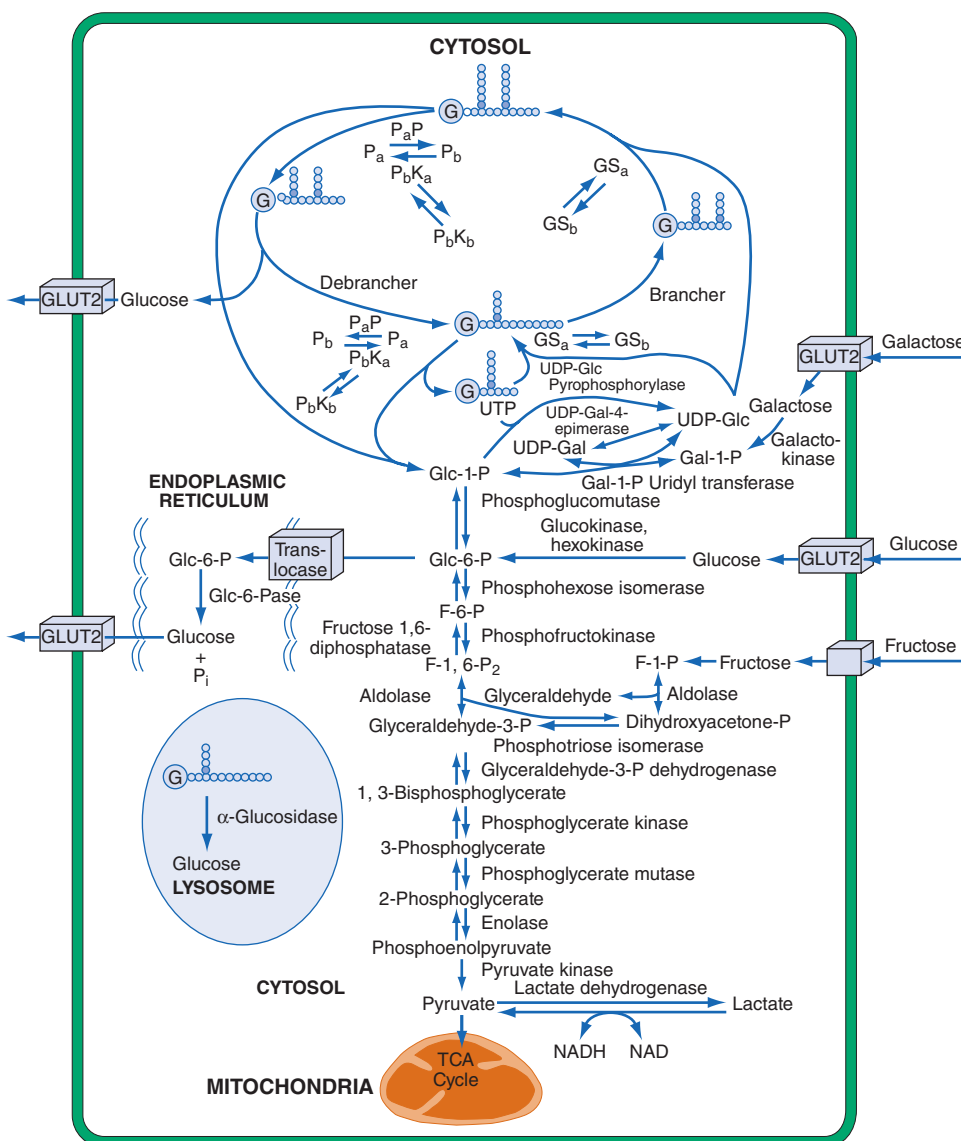


FIGURE 433e-1 Metabolic pathways related to glycogen storage diseases and galactose and fructose disorders. Nonstandard abbreviations are as follows: GS_a , active glycogen synthase; GS_b , inactive glycogen synthase; P_a , active phosphorylase; P_b , inactive phosphorylase; P_aP , phosphorylase a phosphatase; P_bK_a , active phosphorylase β kinase; P_bK_b , inactive phosphorylase β kinase; G, glycogenin, the primer protein for glycogen synthesis. (Modified from AR Beaudet, in KJ Isselbacher et al [eds]: *Harrison's Principles of Internal Medicine*, 13th ed., New York, McGraw-Hill, 1994, p 1855.)

TABLE 433e-1 FEATURES OF GLYCOGEN STORAGE DISEASES AND GALACTOSE AND FRUCTOSE DISORDERS

Type/Common Name	Basic Defect	Clinical Features	Comments
Liver Glycogenoses			
Disorders with hepatomegaly and hypoglycemia			
Ia/von Gierke	Glucose-6-phosphatase	Growth retardation, enlarged liver and kidney, hypoglycemia, elevated blood lactate, cholesterol, triglycerides, and uric acid	Common, severe hypoglycemia. Complications in adulthood include hepatic adenomas, hepatic carcinoma, and renal failure.
Ib	Glucose-6-phosphate translocase	As for Ia, with additional findings of neutropenia and neutrophil dysfunction	~10% of type I
IIla/Cori or Forbes	Liver and muscle debranching enzyme	<i>Childhood:</i> hepatomegaly, growth retardation, muscle weakness, hypoglycemia, hyperlipidemia, elevated liver aminotransferases. Liver symptoms improve with age. <i>Adulthood:</i> muscle atrophy and weakness; onset in third or fourth decades; variable cardiomyopathy, liver cirrhosis, progressive liver failure	Common, intermediate severity of hypoglycemia. Hepatic adenomas, liver cirrhosis, and hepatic carcinoma can occur.
IIlb	Liver debranching enzyme (normal muscle debrancher activity)	Liver symptoms same as in type IIIa; no muscle symptoms	~15% of type III
VI/Hers	Liver phosphorylase	Hepatomegaly, variable hypoglycemia, hyperlipidemia, and ketosis. Symptoms may improve with age.	Rare, often a “benign” glycogenosis, severe cases being recognized
IX/phosphorylase kinase deficiency	Liver phosphorylase kinase α subunit	As for VI	Common, X-linked, typically less severe than autosomal forms; clinical variability within and between subtypes; severe cases being recognized
0/glycogen synthase deficiency	Glycogen synthase	Fasting hypoglycemia and ketosis, elevated lactic acid and hyperglycemia after glucose load, no hepatomegaly	Decreased glycogen stores
XI/Fanconi-Bickel	Glucose transporter 2	Failure to thrive, short stature, hypophosphatemic rickets, metabolic acidosis, hepatomegaly, proximal renal tubular dysfunction, impaired glucose and galactose utilization	Rare, consanguinity in 70%
Disorders with liver cirrhosis			
IV/Andersen	Branching enzyme	Failure to thrive, hypotonia, hepatomegaly, splenomegaly, progressive liver cirrhosis and failure (death usually before fifth year); some without progression	One of the rarer glycogenoses. Other neuromuscular variants exist.
Muscle Glycogenoses			
Disorders with muscle-energy impairment			
V/McArdle	Muscle phosphorylase	Exercise intolerance, muscle cramps, myoglobinuria on strenuous exercise, increased CK, “second-wind” phenomenon	Common, male predominance
VII/Tarui	Phosphofructokinase—M subunit	As for type V, with additional findings of compensated hemolysis, myalgia	Prevalent in Ashkenazi Jews and Japanese
Phosphoglycerate kinase deficiency	Phosphoglycerate kinase	As for type V, with additional findings of hemolytic anemia and CNS dysfunction	Rare, X-linked
Phosphoglycerate mutase deficiency	Phosphoglycerate mutase—M subunit	As for type V	Rare, most patients African American
Lactate dehydrogenase deficiency	Lactic acid dehydrogenase—M subunit	As for type V, with additional findings of erythematous skin eruption and uterine stiffness resulting in childbirth difficulty in females	Rare
Fructose 1,6-bisphosphate aldolase A deficiency	Fructose 1,6-bisphosphate aldolase A	As for type V, with additional finding of hemolytic anemia	Rare
Pyruvate kinase deficiency	Pyruvate kinase—muscle isozyme	Muscle cramps and/or fixed muscle weakness	Rare
Muscle phosphorylase kinase deficiency	Muscle-specific phosphorylase kinase	As for type V. Some patients may have muscle weakness and atrophy.	Rare, autosomal recessive
β -Enolase deficiency	Muscle β -enolase	Exercise intolerance	Rare
Disorders with progressive skeletal muscle myopathy and/or cardiomyopathy			
II/Pompe	Lysosomal acid α -glucosidase	<i>Infantile:</i> hypotonia, muscle weakness, cardiac enlargement and failure, fatal early <i>late onset (juvenile and adult):</i> progressive skeletal muscle weakness and atrophy, proximal muscle and respiratory muscle seriously affected	Common, undetectable or very low level of enzyme activity in infantile form; variable residual enzyme activity in late-onset form
PRKAG2 deficiency	AMP-activated gamma 2 protein kinase	Severe cardiomyopathy and early heart failure (9–55 years). Congenital fetal form is rapidly fatal with hypertrophic cardiomyopathy and Wolff-Parkinson-White syndrome.	Very rare, autosomal dominant
Danon’s disease	Lysosomal-associated membrane protein 2 (LAMP2)	Severe cardiomyopathy and heart failure (8–15 years)	Very rare, X-linked

(Continued)

TABLE 433e-1 FEATURES OF GLYCOGEN STORAGE DISEASES AND GALACTOSE AND FRUCTOSE DISORDERS (CONTINUED)

Type/Common Name	Basic Defect	Clinical Features	Comments
Galactose Disorders			
Galactosemia with uridylyltransferase deficiency	Galactose 1-phosphate uridylyltransferase	Vomiting, hepatomegaly, jaundice, cataracts, amino aciduria, failure to thrive	Long-term complications exist despite early diagnosis and treatment.
Galactokinase deficiency	Galactokinase	Cataracts	Benign
Uridine diphosphate galactose 4-epimerase deficiency	Uridine diphosphate galactose 4-epimerase	Similar to transferase deficiency with additional findings of hypotonia and nerve deafness	Benign variant exists.
Fructose Disorders			
Essential fructosuria	Fructokinase	Asymptomatic, positive urine reducing substance	Benign, autosomal recessive
Hereditary fructose intolerance	Fructose 1,6-bisphosphate aldolase B	Vomiting, lethargy, failure to thrive, hepatic failure, aversion to sweets, severity of symptoms depending on age/quantity of sugar ingested	Prognosis good with early diagnosis and fructose restriction, autosomal recessive.
Fructose 1,6-diphosphatase deficiency	Fructose 1,6-diphosphatase	Episodic hypoglycemia, hyperlactic acidemia, and ketoacidosis usually following illness, hepatomegaly	Avoid fasting, good prognosis

Abbreviations: CK, creatine kinase; CNS, central nervous system; M, muscle.

type Ib, in which the translocase that transports glucose-6-phosphate across the microsomal membrane is defective. The defects in both subtypes lead to inadequate conversion of glucose-6-phosphate to glucose in the liver and thus make affected individuals susceptible to fasting hypoglycemia.

CLINICAL AND LABORATORY FINDINGS Persons with type I GSD may develop hypoglycemia and lactic acidosis during the neonatal period; however, more commonly, they exhibit hepatomegaly at 3–4 months of age. Hypoglycemia, hypoglycemic seizures, and lactic acidosis can develop after a short fast. These children usually have doll-like faces with fat cheeks, relatively thin extremities, short stature, and a protuberant abdomen that is due to massive hepatomegaly. The kidneys are enlarged, but the spleen and heart are of normal size. The hepatocytes are distended by glycogen and fat, with large and prominent lipid vacuoles. Despite hepatomegaly, liver enzyme levels are usually normal or near normal. Easy bruising and epistaxis are associated with a prolonged bleeding time as a result of impaired platelet aggregation/adhesion. Hyperuricemia is present. Hyperlipidemia includes elevation of triglycerides, low-density lipoproteins, and phospholipids. Type Ib patients have additional findings of neutropenia and impaired neutrophil function, which result in recurrent bacterial infections and oral and intestinal mucosal ulceration. GSD I patients may experience intermittent diarrhea, which can worsen with age. In GSD Ib, diarrhea is largely due to loss of mucosal barrier function caused by inflammation.

LONG-TERM COMPLICATIONS Gout usually becomes symptomatic at puberty as a result of long-term hyperuricemia. Puberty is often delayed. Nearly all female patients have ultrasound findings consistent with polycystic ovaries; however, the other clinical features of polycystic ovary syndrome, such as acne and hirsutism, are not seen. Several reports of successful pregnancy in women with GSD I suggest that fertility is not affected. Increased bleeding during menstrual cycles, including life-threatening menorrhagia, has been reported. Secondary to lipid abnormalities, there is an increased risk of pancreatitis. Patients with GSD I may be at increased risk for cardiovascular disease. In adult patients, frequent fractures can occur and radiographic evidence of osteopenia/osteoporosis can be found; in prepubertal patients, radial bone mineral content is significantly reduced. Pulmonary hypertension—although rare—has been reported. By the second or third decade of life, many patients with type I GSD develop hepatic adenomas that can hemorrhage and, in some cases, become malignant. Renal disease is a serious late complication. Almost all patients older than 20 years have proteinuria, and many have hypertension, kidney stones, nephrocalcinosis, and altered creatinine clearance. Laboratory findings include abnormally high levels of blood lactate, triglycerides, cholesterol, and uric acid. In some patients, renal function deteriorates and progresses to complete failure, requiring dialysis or transplantation.

DIAGNOSIS Clinical presentation and abnormal plasma lactate and lipid values suggest that a patient may have GSD I, but gene-based mutation analysis provides a noninvasive means of reaching a definitive diagnosis for most patients with types Ia and Ib disease. Before the glucose-6-phosphatase and glucose-6-phosphate translocase genes were cloned, a definitive diagnosis required a liver biopsy to demonstrate a deficiency.

Type III GSD (Debrancher Deficiency, Limit Dextrinosis) Type III GSD is an autosomal recessive disorder caused by a deficiency of glycogen debranching enzyme. Debranching and phosphorylase enzymes are responsible for complete degradation of glycogen. When the debranching enzyme is defective, glycogen breakdown is incomplete. Abnormal glycogen accumulates with short outer chains and resembles dextrin.

CLINICAL AND LABORATORY FINDINGS Deficiency of glycogen debranching enzyme causes hepatomegaly, hypoglycemia, short stature, variable skeletal myopathy, and cardiomyopathy. The disorder usually involves both liver and muscle and, in such cases, is termed *type IIIa* GSD. However, in ~15% of patients, the disease appears to involve only the liver and is classified as *type IIIb*. Hypoglycemia and hyperlipidemia occur in children. In type III disease (as opposed to type I disease), fasting ketosis can be prominent, aminotransferase levels are elevated, and blood lactate and uric acid concentrations are usually normal. Serum creatine kinase (CK) levels can sometimes be used to identify patients with muscle involvement, but normal levels do not rule out muscle enzyme deficiency. In most patients with type III disease, hepatomegaly improves with age; however, liver cirrhosis and hepatocellular carcinoma may occur in late adulthood. Hepatic adenomas may occur, although less commonly than in GSD I. Left ventricular hypertrophy and life-threatening arrhythmias have been reported. Patients with type IIIa disease may experience muscle weakness in childhood that can become severe after the third or fourth decade of life. Polycystic ovaries are common in GSD III, and some patients develop features of polycystic ovarian syndrome, such as hirsutism and irregular menstrual cycles. Reports of successful pregnancy in women with GSD III suggest that fertility is normal.

DIAGNOSIS In type IIIa GSD, deficient debranching enzyme activity can be demonstrated in liver, skeletal muscle, and heart. In contrast, patients with type IIIb have debranching enzyme deficiency in the liver but not in muscle. The liver has distended hepatocytes due to glycogen buildup; areas of fibrosis are also noted very early in the disease course. In the past, definitive assignment of subtype required enzyme assays in both liver and muscle. DNA-based analyses now provide a noninvasive way of subtyping these disorders in most patients. However, the large size of the gene and the distribution of private mutations across it pose challenges in DNA-based analysis.

433e-4 Type IX GSD (Liver Phosphorylase Kinase Deficiency) Defects of PhK cause a heterogeneous group of glycogenoses. The PhK enzyme complex consists of four subunits (α , β , γ , and δ). Each subunit is encoded by different genes (X chromosome as well as autosomes) that are differentially expressed in various tissues. PhK deficiency can be divided into several subtypes on the basis of the gene/subunit involved, the tissues primarily affected, and the mode of inheritance. The most common subtype is X-linked liver PhK deficiency, which is also one of the most common liver glycogenoses. PhK activity may also be deficient in erythrocytes and leukocytes but is normal in muscle. Typically, a child between the ages of 1 year and 5 years presents with growth retardation and hepatomegaly. Children tend eventually to exhibit normal growth patterns initiated by a delayed growth spurt during puberty. Liver fibrosis has been identified in some patients, including children. Levels of cholesterol, triglycerides, and liver enzymes are mildly elevated. Fasting ketosis is another feature of the disease. Lactic and uric acid levels are usually normal. Hypoglycemia is typically mild; however, phenotypic variability is being increasingly recognized, with significant involvement in some cases of the X-linked form. The accumulated glycogen in liver (β particles, rosette form) has a frayed or burst appearance and is less compact than the glycogen seen in type I or type III GSD. Hepatomegaly and abnormal blood chemistries gradually return to normal with age. Most adults reach a normal final height and are practically asymptomatic, despite persistent PhK deficiency. The prognosis is usually good, and adult patients have minimal hepatomegaly. Some patients have significant ketosis and progressive liver disease that can advance to fibrosis and liver failure. It is recommended that adult patients be monitored for hepatic complications with regular CT or MRI scans.

Treatment is symptom based. A high-carbohydrate diet and frequent feedings are effective in preventing hypoglycemia. Some patients require no specific treatment. Recent studies have shown that instituting a treatment regimen of cornstarch and protein feedings early, even in seemingly stable patients, may prevent long-term complications. Blood ketones and glucose should be evaluated during times of stress.

Other subtypes of type IX GSD include an autosomal recessive form of liver and muscle PhK deficiency, an autosomal recessive form of liver PhK deficiency that often develops into liver cirrhosis, a muscle-specific PhK deficiency that causes cramps and myoglobinuria with exercise, and a cardiac-specific PhK deficiency that is lethal during infancy because of massive glycogen deposition in the myocardium. The finding of PhK deficiency in the cardiac-specific PhK deficiency may be a secondary phenomenon, as a subset of these patients have mutations in PRKAG2.

Other Liver Glycogenoses with Hepatomegaly and Hypoglycemia These disorders include hepatic phosphorylase deficiency (Hers disease, type VI) and hepatic glycogenosis with renal Fanconi syndrome (type XI). Patients with GSD type VI can have growth retardation, hyperlipidemia, and hyperketosis in addition to hepatomegaly and hypoglycemia. Some patients have a benign clinical course. GSD XI is caused by defects in the facilitative glucose transporter 2 (GLUT-2), which transports glucose and galactose in and out of hepatocytes, pancreatic cells, and the basolateral membranes of intestinal and renal epithelial cells. The disease is characterized by proximal renal tubular dysfunction, impaired glucose and galactose utilization, and accumulation of glycogen in liver and kidney.

SELECTED MUSCLE GLYCOGENOSES

DISORDERS WITH MUSCLE-ENERGY IMPAIRMENT

Type V GSD (Muscle Phosphorylase Deficiency, McArdle Disease) Type V GSD is an autosomal recessive disorder caused by deficiency of muscle phosphorylase. McArdle disease is a prototypical muscle-energy disorder as the enzyme deficiency limits ATP generation by glycolysis and results in glycogen accumulation.

CLINICAL AND LABORATORY FINDINGS Usually, symptoms first develop in adulthood and involve exercise intolerance with muscle cramps. Two types of activity tend to cause symptoms: (1) brief exercise of great intensity, such as sprinting or carrying heavy loads; and (2) less intense but sustained activity, such as climbing stairs or walking uphill. Most

patients can engage in moderate exercise, such as walking on level ground, for long periods. Patients often exhibit the “second-wind” phenomenon, in which, after a short break from the initiation of strenuous physical effort, they are able to continue the activity without pain. Although most patients experience episodic muscle pain and cramping as a result of exercise, 35% report permanent pain that seriously affects sleep and other activities. About half of patients report burgundy-colored urine after exercise; this coloration results from myoglobinuria secondary to rhabdomyolysis. Intense myoglobinuria after vigorous exercise can lead to renal failure. Clinical heterogeneity is uncommon; however, there are cases with symptom onset as late as the eighth decade and cases that present early with hypotonia, generalized muscle weakness, and progressive respiratory insufficiency, which is often fatal.

Although cardiac involvement is not usually associated with muscle phosphorylase deficiency, hypertrophic cardiomyopathy has been observed in an adult patient with GSD V. In rare cases, electromyographic findings may suggest inflammatory myopathy, a diagnosis that may be confused with polymyositis. These patients may be at risk for statin-induced myopathy and rhabdomyolysis.

During rest, the serum CK level is usually elevated; after exercise, the CK level increases even more. Exercise also increases levels of blood ammonia, inosine, hypoxanthine, and uric acid; these abnormalities reflect residues of accelerated muscle purine nucleotide recycling as a result of insufficient ATP production. NADH is underproduced during physical exertion.

DIAGNOSIS Lack of an increase in blood lactate and exaggerated blood ammonia elevations after an ischemic exercise test are indicative of a muscle glycogenosis and suggest a defect in the conversion of glycogen or glucose to lactate. This abnormal exercise response, however, can also occur with other defects in glycogenolysis or glycolysis, such as deficiencies of muscle phosphofructokinase or debranching enzyme (when the test is done after fasting). The cycle test detects the hallmark heart rate observed during the second-wind phenomenon. A definitive diagnosis is made by enzymatic assay in muscle tissue or by mutation analysis of the myophosphorylase gene.

DISORDERS WITH PROGRESSIVE SKELETAL MUSCLE MYOPATHY AND/OR CARDIOMYOPATHY

Pompe Disease, Type II GSD (Acid α -1,4 Glucosidase Deficiency) Pompe disease is an autosomal recessive disorder caused by a deficiency of lysosomal acid α -1,4 glucosidase, an enzyme responsible for the degradation of glycogen in the lysosomes. This disease is characterized by the accumulation of glycogen in the lysosomes as opposed to accumulation in cytoplasm (as in the other glycogenoses).

CLINICAL AND LABORATORY FINDINGS The disorder encompasses a range of phenotypes. Each includes myopathy but differs in the age of onset, extent of organ involvement, and clinical severity. The most severe is the infantile form, with cardiomegaly, hypotonia, and death before the age of 1 year. Infants may appear normal at birth but soon develop generalized muscle weakness with feeding difficulties, macroglossia, hepatomegaly, and congestive heart failure due to hypertrophic cardiomyopathy.

The late-onset form (juvenile/late-childhood or adult form) is characterized by skeletal muscle manifestations, usually with minimal or no cardiac involvement, and a more slowly progressive course. The juvenile form typically presents as delayed motor milestones (if age of onset is early enough) and difficulty in walking. With disease progression, patients often develop swallowing difficulties, proximal muscle weakness, and respiratory muscle involvement. Death may occur before the end of the second decade.

Adults typically present between the second and seventh decades with slowly progressive myopathy without overt cardiac involvement. The clinical picture is dominated by slowly progressive proximal limb girdle muscle weakness. The pelvic girdle, paraspinal muscles, and diaphragm are most seriously affected. Respiratory symptoms include somnolence, morning headache, orthopnea, and exertional dyspnea. In rare instances, patients present with respiratory insufficiency as the initial symptom. Basilar artery aneurysms and dilation of the ascending aorta have been observed in patients with Pompe disease. Ptosis,

lingual weakness, gastrointestinal dysmotility, and incontinence due to poor sphincter tone are now being recognized as part of the clinical spectrum. Individuals with advanced disease often require some form of ventilation and are dependent on a walking aid or wheelchair.

Laboratory findings include elevated levels of serum CK, aspartate aminotransferase, and lactate dehydrogenase. Levels of urine glucose tetrasaccharide (Hex4), a breakdown product of glycogen, are elevated, especially on the severe end of the disease spectrum. In infants, chest x-ray shows massive cardiomegaly, and electrocardiographic findings include a high-voltage QRS complex and a shortened PR interval. Muscle biopsy shows vacuoles that stain positive for glycogen; the muscle acid phosphatase level is increased, presumably from a compensatory increase of lysosomal enzymes. Electromyography reveals myopathic features, with irritability of muscle fibers and pseudomyotonic discharges. Serum CK is not always elevated in adults, and, depending on the muscle biopsied or tested, muscle histology or electromyography may not be abnormal. The affected muscle should be examined.

DIAGNOSIS The confirmatory step for a diagnosis of Pompe disease is enzyme assay demonstrating deficient acid α -glucosidase or a gene sequence with two pathogenic mutations in the GAA gene. Enzyme activity can be measured in muscle, cultured skin fibroblasts, or blood. Deficiency is usually more severe in the infantile form. Early diagnosis is the key to treatment efficacy.

GSD Mimicking Hypertrophic Cardiomyopathy Deficiency of LAMP2—also called Danon's disease—of the protein kinase, AMP-activated gamma 2 noncatalytic subunit (PRKAG2), results in the accumulation of glycogen in the heart and skeletal muscle. LAMP2 deficiency is X-linked, whereas PRKAG2 deficiency is autosomal dominant. Clinically, both subsets of patients present primarily with hypertrophic cardiomyopathy. Their electrophysiologic abnormalities, particularly ventricular preexcitation and conduction defects, can distinguish them from patients with hypertrophic cardiomyopathy resulting from defects in sarcomere-protein genes. In patients with LAMP2 deficiency, mental delays are common and the onset of cardiac symptoms, including chest pain, palpitation, syncope, and cardiac arrest, can occur between the ages of 8 and 15 years—i.e., earlier than the average age of 33 years for patients with PRKAG2 deficiency. Patients as young as 9 years old have presented with PRKAG2 deficiency. A rapidly fatal congenital form of PRKAG2 presents in early infancy with severe hypertrophic cardiomyopathy and Wolff-Parkinson-White syndrome. In these patients, levels of PhK have been found to be low. The prognosis for LAMP2 deficiency is poor, with progressive end-stage heart failure early in adulthood. By contrast, except in the fatal congenital form, long-term survival is possible for patients with cardiomyopathy due to PRKAG2 mutations. Some patients may require the implantation of a pacemaker and aggressive control of arrhythmias. Congestive heart failure has been documented in patients with PRKAG2 deficiency.

TREATMENT GLYCOGEN STORAGE DISEASE MIMICKING HYPERTROPHIC CARDIOMYOPATHY

Heart transplantation has been suggested as a preventive measure for LAMP2 deficiency and noncongenital PRKAG2 deficiency.

SELECTED DISORDERS OF GALACTOSE METABOLISM

“Classic” galactosemia is caused by galactose 1-phosphate uridylyltransferase (GALT) deficiency. It is a serious disease with an incidence of 1 in 60,000 and an early onset of symptoms. The newborn infant normally receives up to 40% of caloric intake as lactose (glucose + galactose). Without the transferase, the infant is unable to metabolize galactose 1-phosphate (Fig. 433e-1), which consequently accumulates, resulting in injury to parenchymal cells of the kidney, liver, and brain. After the first feeding, infants can present with vomiting, diarrhea, hypotonia, jaundice, and hepatomegaly. Patients with galactosemia are at increased risk for *Escherichia coli* neonatal sepsis; the onset of sepsis often precedes the diagnosis of galactosemia.

Widespread newborn screening for galactosemia has identified these infants early and allowed them to be placed on dietary restriction. Elimination of galactose from the diet reverses growth failure as well as renal and hepatic dysfunction, improving the prognosis. However, on long-term follow-up, some patients still have ovarian failure manifesting as primary or secondary amenorrhea as well as developmental delays and learning disabilities that increase in severity with age. Of women with classic galactosemia, 80–90% or more report hypergonadotropic hypogonadism. While most female patients are infertile when they reach childbearing age, a few have given birth. Several mutations appear to be protective, particularly the p.Ser135Leu mutation, which is more common in the African-American population. Methods for fertility preservation, such as cryopreservation, are still in the experimental stages. In addition, most patients have speech disorders, and a smaller proportion demonstrate poor growth and impaired motor function and balance (with or without overt ataxia). Adults on dairy-free diets have developed cataracts, tremors, and low bone density. The treatment of galactosemia to prevent long-term complications remains a challenge.

Deficiency of galactokinase (Fig. 433e-1) causes cataracts. Deficiency of uridine diphosphate galactose 4-epimerase can be benign when the enzyme deficiency is limited to blood cells but can be as severe as classic galactosemia when the enzyme deficiency is generalized.

SELECTED DISORDERS OF FRUCTOSE METABOLISM

Fructokinase deficiency, or essential fructosemia (Fig. 433e-1), causes a benign condition that is usually an incidental finding made through the detection of fructose as a reducing substance in the urine.

Deficiency of fructose 1,6-bisphosphate aldolase (aldolase B; hereditary fructose intolerance) is a serious disease in infants. These patients are healthy and asymptomatic until fructose or sucrose (table sugar) is ingested (usually from fruit, sweetened cereal, or sucrose-containing formula). Clinical manifestations may include jaundice, hepatomegaly, vomiting, lethargy, irritability, and convulsions. The incidence of celiac disease is higher among patients with hereditary fructose intolerance (>10%) than in the general population (1–3%). Laboratory findings show prolonged clotting time, hypoalbuminemia, elevation of bilirubin and aminotransferase levels, and proximal renal tubular dysfunction. If the disease is not diagnosed and intake of the noxious sugar continues, hypoglycemic episodes recur, and liver and kidney failure progresses, eventually leading to death. Treatment requires the elimination of all sources of sucrose, fructose, and sorbitol from the diet. Through this treatment, liver and kidney dysfunction improve, and catch-up growth is common; intellectual development is usually unimpaired. Over time, the patient's symptoms become milder, even after fructose ingestion, and the long-term prognosis is good.

Fructose 1,6-diphosphatase deficiency is characterized by childhood life-threatening episodes of acidosis, hypoglycemia, hyperventilation, convulsions, and coma. These episodes are triggered by febrile infections and gastroenteritis when oral food intake decreases. Laboratory findings show low blood glucose levels, high lactate and uric acid levels, and metabolic acidosis. Unlike hereditary fructose intolerance, this deficiency usually is not associated with an aversion to sweets, and renal tubular and liver functions are normal. Treatment of acute attacks requires the correction of hypoglycemia and acidosis by IV infusion. Later, avoidance of fasting and elimination of fructose and sucrose from the diet prevent further episodes. A slowly released carbohydrate such as cornstarch is useful for the long-term prevention of hypoglycemia. The prognosis is good, as patients who survive childhood develop normally.

GLOBAL CONSIDERATIONS

The GSDs and other inherited disorders of carbohydrate metabolism, although rare, have been reported in most ethnic populations. The prevalent genetic mutations for each disease may vary in different ethnic populations, but clinical symptoms are remarkably similar and treatment guidelines apply to all populations. The practice of newborn screening should be considered worldwide to intercept the rapid progression of many of these disorders.

434e Inherited Disorders of Amino Acid Metabolism in Adults

Nicola Longo

Amino acids are not only the building blocks of proteins but also serve as neurotransmitters (glycine, glutamate, γ -aminobutyric acid) or as precursors of hormones, coenzymes, pigments, purines, or pyrimidines. Eight amino acids, referred to as *essential*, cannot be synthesized by humans and must be obtained from dietary sources. The others are formed endogenously. Each amino acid has a unique degradative pathway by which its nitrogen and carbon components are used for the synthesis of other amino acids, carbohydrates, and lipids. Disorders of amino acid metabolism and transport (Chap. 435e) are individually rare—the incidences range from 1 in 10,000 for cystinuria or phenylketonuria to 1 in 200,000 for homocystinuria or alkaptonuria—but collectively, they affect perhaps 1 in 1000 newborns. Almost all are transmitted as autosomal recessive traits.

The features of inherited disorders of amino acid catabolism are summarized in Table 434e-1. In general, these disorders are named for the compound that accumulates to highest concentration in blood (*-emias*) or urine (*-urias*). In the aminoacidopathies, the parent amino acid is found in excess, whereas products in the catabolic pathway accumulate in organic acidemias. Which compound(s) accumulates depends on the site of the enzymatic block, the reversibility of the reactions proximal to the lesion, and the availability of alternative pathways of metabolic “runoff.” Biochemical and genetic heterogeneity are common. Five distinct forms of hyperphenylalaninemia, nine forms of homocystinuria, and methylmalonic acidemia are recognized. Such heterogeneity reflects the presence of a large array of molecular defects.

The manifestations of these conditions differ widely (Table 434e-1). Some, such as sarcosinemia, produce no clinical consequences. At the other extreme, complete deficiency of ornithine transcarbamylase is lethal in the untreated neonate. Central nervous system (CNS) dysfunction, in the form of developmental retardation, seizures, alterations in sensorium, or behavioral disturbances, is present in more than half the disorders. Protein-induced vomiting, neurologic dysfunction, and hyperammonemia occur in many disorders of urea cycle intermediates. Metabolic ketoacidosis, often accompanied by hyperammonemia, is a frequent presenting finding in disorders of branched-chain amino acid metabolism. Some disorders produce focal tissue or organ involvement such as liver disease, renal failure, cutaneous abnormalities, or ocular lesions.

The analysis of plasma amino acids (by ion-exchange chromatography), urine organic acids (by gas chromatography/mass spectrometry), and plasma acylcarnitine profile (by tandem mass spectrometry) is commonly used to diagnose and monitor most of these disorders. The diagnosis is confirmed by enzyme assay on cells or tissues from the patients or by DNA testing. The clinical manifestations in many of these conditions can be prevented or mitigated if a diagnosis is achieved early and appropriate treatment (e.g., dietary protein or amino acid restriction or vitamin supplementation) is instituted promptly. For this reason, newborn screening programs seek to identify several of these disorders. Infants with a positive screening test need additional metabolic testing (usually suggested by the newborn screening program) to confirm or exclude the diagnosis. Confirmed cases should be referred to a metabolic center for initiation of therapy. The parents need to be counseled about the recurrence risk of the disease in future pregnancies. In some cases, parents need testing to exclude metabolic alterations seen in carriers for some of these disorders (such as some forms of homocystinuria) or because they might have a disorder themselves (such as glutaric acidemia type 1, methylcrotonyl coenzyme A carboxylase deficiency, or fatty acid oxidation defects). Some metabolic disorders can remain asymptomatic until adult age, presenting only when fasting or severe stress require full activity of affected metabolic pathways to provide energy.

Selected disorders that illustrate the principles, properties, and problems presented by the disorders of amino acid metabolism are discussed in this chapter.

434e-1

THE HYPERPHENYLALANINEMIAS

The hyperphenylalaninemias (Table 434e-1) result from impaired conversion of phenylalanine to tyrosine. The most common and clinically important is *phenylketonuria* (frequency 1:10,000), which is an autosomal recessive disorder characterized by an increased concentration of phenylalanine and its by-products in body fluids and by severe mental retardation if untreated in infancy. It results from reduced activity of phenylalanine hydroxylase. The accumulation of phenylalanine inhibits the transport of other amino acids required for protein or neurotransmitter synthesis, reduces synthesis and increases degradation of myelin, and leads to inadequate formation of norepinephrine and serotonin. Phenylalanine is a competitive inhibitor of tyrosinase, a key enzyme in the pathway of melanin synthesis, and accounts for the hypopigmentation of hair and skin. Untreated children with classic phenylketonuria are normal at birth but fail to attain early developmental milestones, develop microcephaly, and demonstrate progressive impairment of cerebral function. Hyperactivity, seizures, and severe intellectual disability are major clinical problems later in life. Electroencephalographic abnormalities; “mousy” odor of skin, hair, and urine (due to phenylacetate accumulation); and a tendency to develop hypopigmentation and eczema complete the devastating clinical picture. In contrast, affected children who are detected and treated at birth show none of these abnormalities.

TREATMENT PHENYLKETONURIA

To prevent intellectual disability, diagnosis and initiation of dietary treatment of classic phenylketonuria must occur before the child is 2 weeks of age. For this reason, most newborns in North America, Australia, and Europe are screened by determinations of blood phenylalanine levels. Abnormal values are confirmed using quantitative analysis of plasma amino acids. Dietary phenylalanine restriction is usually instituted if blood phenylalanine levels are $>360 \mu\text{mol/L}$ (6 mg/dL). Treatment consists of a special diet low in phenylalanine and supplemented with tyrosine, since tyrosine becomes an essential amino acid in phenylalanine hydroxylase deficiency. With therapy, plasma phenylalanine concentrations should be maintained between 120 and 360 $\mu\text{mol/L}$ (2 and 6 mg/dL). Dietary restriction should be continued and monitored indefinitely. Some patients with milder forms of phenylketonuria (phenylalanine $<1200 \mu\text{mol/L}$ at presentation) show increased tolerance to dietary proteins and improved metabolic control when treated with tetrahydrobiopterin (5–20 mg/kg per day), an essential cofactor of phenylalanine hydroxylase.

A number of women with phenylketonuria who have been treated since infancy will reach adulthood and become pregnant. If maternal phenylalanine levels are not strictly controlled before and during pregnancy, their offspring are at increased risk for congenital defects and microcephaly (*maternal phenylketonuria*). After birth, these children have severe intellectual disability and growth retardation. Pregnancy risks can be minimized by continuing lifelong phenylalanine-restricted diets and assuring strict phenylalanine restriction 2 months prior to conception and throughout gestation.

THE HOMOCYSTINURIAS (HYPERHOMOCYSTEINEMIAS)

The homocystinurias are nine biochemically and clinically distinct disorders (Table 434e-1) characterized by increased concentration of the sulfur-containing amino acid homocystine in blood and urine.

Classic homocystinuria, the most common (frequency 1:200,000), results from reduced activity of cystathionine β -synthase (Fig. 434e-1), the pyridoxal phosphate–dependent enzyme that condenses homocysteine with serine to form cystathionine. Most patients present between

TABLE 434e-1 INHERITED DISORDERS OF AMINO ACID METABOLISM

Amino Acid(s)	Condition	Enzyme Defect	Clinical Findings	Inheritance
Phenylalanine	Phenylketonuria	Phenylalanine hydroxylase	Intellectual disability, microcephaly, hypopigmented skin and hairs, eczema, "mousy" odor	AR
	DHPR deficiency	Dihydropteridine reductase	Intellectual disability, hypotonia, spasticity, myoclonus	AR
	PTPS deficiency	6-Pyruvoyl-tetrahydropterin synthase	Dystonia, neurologic deterioration, seizures, intellectual disability	AR
	GTP cyclohydrolase I deficiency	GTP cyclohydrolase I	Intellectual disability, seizures, dystonia, temperature instability	AR
Tyrosine	Carbinolamine dehydratase deficiency	Pterin-4 α -carbinolamine dehydratase	Transient hyperphenylalaninemia (benign)	AR
	Tyrosinemia type I (hepatorenal)	Fumarylacetoacetate hydrolase	Liver failure, cirrhosis, rickets, failure to thrive, peripheral neuropathy, "boiled cabbage" odor	AR
	Tyrosinemia type II (oculocutaneous)	Tyrosine transaminase	Palmoplantar keratosis, painful corneal erosions with photophobia, intellectual disability (?)	AR
	Tyrosinemia type III	4-Hydroxyphenylpyruvate dioxygenase	Hypertyrosinemia with normal liver function, occasional mental delay	AR
	Hawkinsinuria	4-Hydroxyphenylpyruvate dioxygenase	Transient failure to thrive, metabolic acidosis in infancy	AD
	Alkaptonuria	Homogentisic acid oxidase	Ochronosis, arthritis, cardiac valve involvement, coronary artery calcification	AR
	Albinism (oculocutaneous)	Tyrosinase	Hypopigmentation of hair, skin, and optic fundus; visual loss; photophobia	AR
GABA	Albinism (ocular)	Different enzymes or transporters	Hypopigmentation of optic fundus, visual loss	AR, XL
	DOPA-responsive dystonia	Tyrosine hydroxylase	Rigidity, truncal hypotonia, tremor, intellectual disability	AR
Tryptophan	4-Hydroxybutyric aciduria	Succinic semialdehyde dehydrogenase	Seizures, intellectual disability, ataxia	AR
	Tryptophanuria	Unknown	Intellectual disability, ataxia, skin photosensitivity	AR
Histidine	Hydroxykynureninuria	Kynureninase	Intellectual disability, spasticity	AR
	Histidinemia	Histidine-ammonia lyase	Benign	AR
	Urocanic aciduria	Urocanase	Benign	AR
Glycine	Formiminoglutamic aciduria	Formiminotransferase	Occasional intellectual disability	AR
	Glycine encephalopathy	Glycine cleavage (4 enzymes)	Infantile seizures, lethargy, apnea, profound intellectual disability	AR
	Sarcosinemia	Sarcosine dehydrogenase	Benign	AR
	Hyperoxaluria type I	Alanine:glyoxylate aminotransferase	Calcium oxalate nephrolithiasis, renal failure	AR
Serine	Hyperoxaluria type II	D-Glyceric acid dehydrogenase/glyoxylate reductase	Calcium oxalate nephrolithiasis, renal failure	AR
	Phosphoglycerate dehydrogenase deficiency	Phosphoglycerate dehydrogenase	Seizures, microcephaly, intellectual disability	AR
Proline	Hyperprolinemia type I	Proline oxidase	Benign	AR
	Hyperprolinemia type II	Δ^1 -Pyrroline-5-carboxylate dehydrogenase	Febrile seizures, intellectual disability	AR
	Hyperhydroxyprolinemia	Hydroxyproline oxidase	Benign	AR
Methionine	Prolidase deficiency	Prolidase	Mild intellectual disability, chronic dermatitis	AR
	Hypermethioninemia	Methionine adenosyltransferase	Usually benign	AR
	S-Adenosylhomocysteine hydrolase deficiency	S-Adenosylhomocysteine hydrolase	Hypotonia, intellectual disability, absent tendon reflexes, delayed myelination	AR
	Glycine N-methyltransferase deficiency	Glycine N-methyltransferase	Elevated liver transaminases	AR
Homocystine	Adenosine kinase deficiency	Adenosine kinase	Intellectual disability, seizures, liver dysfunction	AR
	Homocystinuria	Cystathionine β -synthase	Lens dislocation, thrombotic vascular disease, intellectual disability, osteoporosis	AR
	Homocystinuria	5,10-Methylenetetrahydrofolate reductase	Intellectual disability, gait and psychiatric abnormalities, recurrent strokes	AR
	Homocystinuria	Methionine synthase (cblE, -G)	Intellectual disability, hypotonia, seizures, megaloblastic anemia	AR
Cystathionine	Homocystinuria and methylmalonic acidemia	Vitamin B ₁₂ lysosomal efflux and metabolism (cblC, -D, -F, -J, -X)	Intellectual disability, lethargy, failure to thrive, hypotonia, seizures, megaloblastic anemia	AR
	Cystathioninuria	β -Cystathioninase	Benign	AR
Cystine	Cystinosis	Cystinosin CTNS (lysosomal efflux)	Renal Fanconi's syndrome, rickets, photophobia, hypotonia, renal failure	AR
S-Sulfo-L-cysteine	Sulfocysteinuria	Sulfate oxidase or molybdenum cofactor deficiency	Seizures, intellectual disability, dislocated lenses	AR

(Continued)

TABLE 434e-1 INHERITED DISORDERS OF AMINO ACID METABOLISM (CONTINUED)

Amino Acid(s)	Condition	Enzyme Defect	Clinical Findings	Inheritance
Lysine	Hyperlysinemia, saccharopinuria	α -Amino adipic semialdehyde synthase	Benign	AR
	Pyridoxine-dependent seizures	L- Δ^1 -Piperidine-6-carboxylate dehydrogenase	Seizures	AR
Lysine, tryptophan	α -Ketoadipic acidemia	α -Ketoadipic acid dehydrogenase	Benign	?
	Glutaric acidemia type I	Glutaryl-CoA dehydrogenase	Progressive severe dystonia and athetosis, motor delays	AR
	Glutaric acidemia type II	Electron transfer flavoproteins (ETF) or ETF:ubiquinone oxidoreductase	Hypoglycemia, metabolic acidosis, "sweaty feet" odor, hypotonia, cardiomyopathy	AR
Ornithine	Gyrate atrophy of the choroid and retina	Ornithine- δ -aminotransferase	Myopia, night blindness, loss of peripheral vision, cataracts, chorioretinal degeneration	AR
Urea cycle	Carbamoylphosphate synthase-1 deficiency	Carbamoylphosphate synthase-1	Lethargy progressing to coma, protein aversion, intellectual disability, hyperammonemia	AR
	N-Acetylglutamate synthase deficiency	N-Acetylglutamate synthase	Lethargy progressing to coma, protein aversion, intellectual disability, hyperammonemia	AR
	Ornithine transcarbamylase deficiency	Ornithine transcarbamylase	Lethargy progressing to coma, protein aversion, intellectual disability, hyperammonemia	XL
	Citrullinemia type I	Argininosuccinate synthase	Lethargy progressing to coma, protein aversion, intellectual disability, hyperammonemia	AR
	Argininosuccinic acidemia	Argininosuccinate lyase	Lethargy progressing to coma, protein aversion, intellectual disability, hyperammonemia, trichorhexis nodosa	AR
	Arginase deficiency	Arginase	Spastic tetraparesis, intellectual disability, mild hyperammonemia	AR
	Hyperornithinemia, hyperammonemia, homocitrullinuria	Mitochondrial ornithine carrier ORNT1	Vomiting, lethargy, failure to thrive, intellectual disability, episodic confusion, hyperammonemia, protein intolerance	AR
	Citrullinemia type 2	Mitochondrial aspartate/glutamate carrier CTLN2	Neonatal intrahepatic cholestasis, adult presentation with sudden behavioral changes and stupor, coma, hyperammonemia	AR
Proline, ornithine, arginine	Δ^1 -Pyrroline-5-carboxylate synthase deficiency	Δ^1 -Pyrroline-5-carboxylate synthase	Hypotonia, seizures, neurodegeneration, peripheral neuropathy, joint laxity, skin hyperelasticity, subcapsular cataracts, hyperammonemia	AR
Glutamine	Glutamine synthase deficiency	Glutamine synthase	Brain malformations, pachygyria, seizures, hypotonia, dysmorphic features	AR
Valine	Hypervalinemia	Valine aminotransferase	Vomiting, fever, failure to thrive, hypotonia	AR
	Isobutyryl-CoA dehydrogenase deficiency	Isobutyryl-CoA dehydrogenase	Failure to thrive, anemia, and dilated cardiomyopathy(?)	AR
Leucine, isoleucine	Hyperleucine-isoleucinemia	Leucine-isoleucine aminotransferase	Seizures, failure to thrive, intellectual disability	?
Valine, leucine, isoleucine	Maple syrup urine disease	Branched chain ketoacid dehydrogenase (E1 α , E1 β , E2, E3 deficiency)	Lethargy, vomiting, encephalopathy, seizures, intellectual disability, "maple syrup" odor, protein intolerance	AR
Leucine	Isovaleric acidemia	Isovaleryl-CoA dehydrogenase	Acidosis, ketosis, vomiting, coma, hyperammonemia, "sweaty feet" odor, protein intolerance	AR
	3-Methylcrotonyl glycinuria	3-Methylcrotonyl-CoA carboxylase	Stress-induced metabolic acidosis, hypotonia, hypoglycemia, "cat's urine" odor	AR
	3-Methylglutaconic aciduria type I	3-Methylglutaconyl-CoA hydratase deficiency	Stress-induced acidosis, leukodystrophy, hypotonia, hepatomegaly	AR
	3-Hydroxy-3-methylglutaric aciduria	3-Hydroxy-3-methylglutaryl-CoA lyase	Stress-induced hypoketotic hypoglycemia and acidosis, encephalopathy, hyperammonemia	AR
Isoleucine	2-Methylbutyryl-glycinuria	2-Methylbutyryl-CoA dehydrogenase	Fasting-induced metabolic acidosis/hypoglycemia	AR
	2-Methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency	2-Methyl-3-hydroxybutyryl-CoA dehydrogenase	Developmental regression, seizures, and rigidity sometimes triggered by illnesses	XL
	3-Oxothiolase deficiency	3-Oxothiolase	Fasting-induced acidosis and ketosis, vomiting, lethargy	AR
Valine, isoleucine, methionine, threonine	Propionic acidemia (pccA, -B, -C)	Propionyl-CoA carboxylase	Metabolic ketoacidosis, hyperammonemia, hypotonia, lethargy, coma, protein intolerance, intellectual disability, hyperglycinemia	AR
	Multiple carboxylase/biotinidase deficiency	Holocarboxylase synthase or biotinidase	Metabolic ketoacidosis, diffuse rash, alopecia, seizures, intellectual disability	AR
	Methylmalonic acidemia (mutase, cblA, -B, racemase)	Methylmalonyl-CoA mutase/racemase or cobalamin reductase/adenosyltransferase	Metabolic ketoacidosis, hyperammonemia, hypertonia, lethargy, coma, protein intolerance, intellectual disability, hyperglycinemia	AR

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; Cbl, cobalamin; DOPA, dihydroxyphenylalanine; GABA, γ -aminobutyric acid; GTP, guanosine 5'-triphosphate; XL, X-linked.

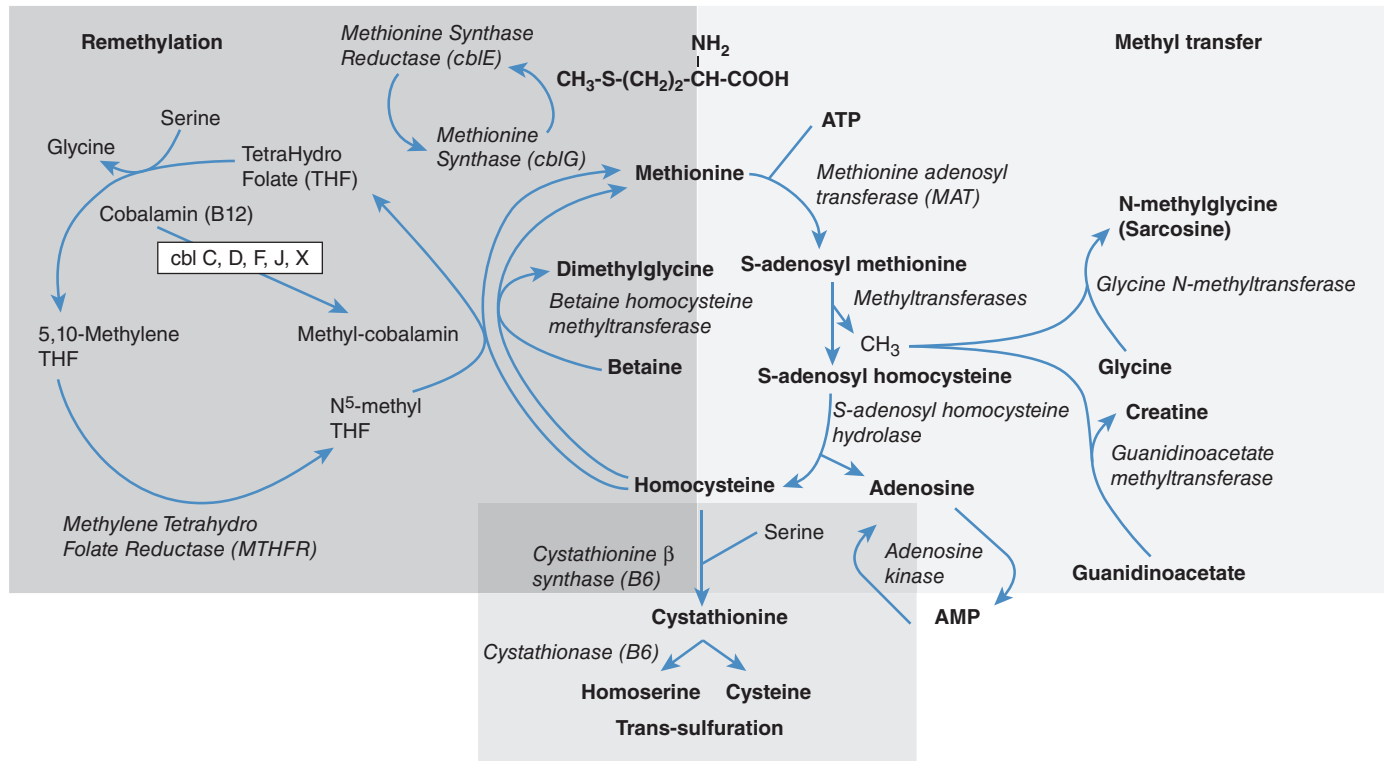


FIGURE 434e-1 Pathways, enzymes, and coenzymes involved in the homocystinurias. Methionine transfers a methyl group during its conversion to homocysteine. Defects in methyl transfer or in the subsequent metabolism of homocysteine by the pyridoxal phosphate (vitamin B₆)-dependent cystathionine β-synthase increase plasma methionine levels. Homocysteine is transformed into methionine via remethylation. This occurs through methionine synthase, a reaction requiring methylcobalamin and folic acid. Deficiencies in these enzymes or lack of cofactors is associated with decreased or normal methionine levels. In an alternative pathway, homocysteine can be remethylated by betaine:homocysteine methyl transferase.

3 and 5 years of age with dislocated optic lenses and intellectual disability (in about half of cases). Some patients develop a marfanoid habitus and radiologic evidence of osteoporosis.

Life-threatening vascular complications (affecting coronary, renal, and cerebral arteries) can occur during the first decade of life and are the major cause of morbidity and mortality. Classic homocystinuria can be diagnosed with analysis of plasma amino acids, showing elevated methionine and presence of free homocysteine. Total plasma homocysteine is also extremely elevated (usually >100 μM). Treatment consists of a special diet restricted in protein and methionine and supplemented with cystine. In approximately half of patients, oral pyridoxine (25–500 mg/d) produces a fall in plasma methionine and homocysteine concentration in body fluids. Folate and vitamin B₁₂ deficiency should be prevented by adequate supplementation. Betaine is also effective in reducing homocysteine levels in pyridoxine-unresponsive patients.

The other forms of homocystinuria are the result of impaired remethylation of homocysteine to methionine. This can be caused by defective methionine synthase or reduced availability of two essential cofactors, 5-methyltetrahydrofolate and methylcobalamin (methyl-vitamin B₁₂).

Hyperhomocysteinemia refers to increased total plasma concentration of homocysteine with or without an increase in free homocysteine (disulfide form). Hyperhomocysteinemia, in the absence of significant homocystinuria, is found in some heterozygotes for the genetic defects noted above or in homozygotes for milder variants. Changes of homocysteine levels are also observed with increasing age; with smoking; in postmenopausal women; in patients with renal failure, hypothyroidism, leukemias, inflammatory bowel disease, or psoriasis; and during therapy with drugs such as methotrexate, nitrous oxide, isoniazid, and some antiepileptic agents. Homocysteine acts as an atherogenic and thrombophilic agent. An increase in total plasma homocysteine is an independent risk factor for coronary, cerebrovascular, and peripheral

arterial disease as well as for deep vein thrombosis (Chap. 291e). Homocysteine is synergistic with hypertension and smoking, and it is additive with other risk factors that predispose to peripheral arterial disease. In addition, hyperhomocysteinemia and folate and vitamin B₁₂ deficiency have been associated with an increased risk of neural tube defects in pregnant women. Vitamin supplements are effective in reducing plasma homocysteine levels in these cases, although there are limited effects on cardiovascular disease.

ALKAPTONURIA

Alkaptonuria is a rare (frequency 1:200,000) disorder of tyrosine catabolism in which deficiency of homogentisate 1,2-dioxygenase (also known as *homogentisic acid oxidase*) leads to excretion of large amounts of homogentisic acid in urine and accumulation of oxidized homogentisic acid pigment in connective tissues (*ochronosis*). Alkaptonuria may go unrecognized until middle life, when degenerative joint disease develops. Prior to this time, about half of patients might be diagnosed for the presence of dark urine. Foci of gray-brown scleral pigment and generalized darkening of the concha, anthelix, and, finally, helix of the ear usually develop after age 30. Low back pain usually starts between 30 and 40 years of age. *Ochronotic arthritis* is heralded by pain, stiffness, and some limitation of motion of the hips, knees, and shoulders. Acute arthritis may resemble rheumatoid arthritis, but small joints are usually spared. Pigmentation of heart valves, larynx, tympanic membranes, and skin occurs, and occasional patients develop pigmented renal or prostatic calculi. Pigment deposition in the heart and blood vessels leads to aortic stenosis necessitating valve replacement, especially after 60 years of age. The diagnosis should be suspected in a patient whose urine darkens to blackness. Homogentisic acid in urine is identified by urine organic acid analysis. Ochronotic arthritis is treated symptomatically with pain medications, spinal surgery, and arthroplasty (Chap. 394). Ascorbic acid and protein

restriction are not effective in reducing homogentisic acid production. By contrast, nitisone (2-[2-nitro-4-trifluoromethylbenzoyl]-1,3-cyclohexanedione), a drug used in tyrosinemia type I, reduces urinary excretion of homogentisic acid and, in conjunction with a low-protein diet, might prevent the long-term complications of alkaptonuria.

UREA CYCLE DEFECTS

Excess ammonia generated from protein nitrogen is removed by the urea cycle, a process mediated by several enzymes and transporters (Table 434e-1). Complete absence of any of these enzymes usually causes severe hyperammonemia in newborns, while milder variants can be seen in adults. The accumulation of ammonia and glutamine leads to brain edema and direct neuronal toxicity. Deficiencies in urea cycle enzymes are individually rare, but as a group, they affect about 1:25,000 individuals. They are all transmitted as autosomal recessive traits, with the exception of ornithine transcarbamylase deficiency, which is X-linked. Hepatocytes of females with ornithine transcarbamylase deficiency express either the normal or the mutant allele due to random X-inactivation and may be unable to remove excess ammonia if mutant cells are predominant.

Infants with classic urea cycle defects present at 1–4 days of life with refusal to eat and lethargy progressing to coma and death. Milder enzyme deficiencies present with protein avoidance, recurrent vomiting, migraine, mood swings, chronic fatigue, irritability, and disorientation that can progress to coma. Females with ornithine transcarbamylase deficiency can present at time of childbirth due to the combination of involuntary fasting and stress that favors catabolism. The diagnosis requires measurement of plasma ammonia, plasma amino acids, and urine orotic acid, useful for differentiating ornithine transcarbamylase deficiency from carbamyl phosphate synthase-1 and

N-acetylglutamate synthase deficiency. Hyperammonemia can also be caused by liver disease from any cause and several organic acidemias and fatty acid oxidation defects (the latter two excluded by the analysis of urine organic acids and plasma acylcarnitine profile).

TREATMENT UREA CYCLE DEFECTS

Therapy is aimed at stopping catabolism and ammonia production by providing adequate calories (as IV glucose and lipids in the comatose patient) and, if needed, insulin. Excess nitrogen is removed by IV phenylacetate and benzoate (0.25 g/kg for the priming dose and subsequently as an infusion over 24 h) that conjugate with glutamine and glycine, respectively, to form phenylacetylglutamine and hippuric acid, water-soluble molecules efficiently excreted in urine. Arginine (200 mg/kg per day) becomes an essential amino acid (except in arginase deficiency) and should be provided intravenously to resume protein synthesis. If these measures fail to reduce ammonia, hemodialysis should be initiated promptly. Chronic therapy consists of a protein-restricted diet, phenylbutyrate, glycerol phenylbutyrate (a liquid drug better tolerated by most patients), arginine, or citrulline supplements, depending on the specific diagnosis. Liver transplantation should be considered in patients with severe urea cycle defects that are difficult to control medically.

Hyperammonemia due to a functional deficiency of glutamine synthase can occur in patients receiving chemotherapy for different malignancies or undergoing solid organ transplants. It can also be seen with hepatic cirrhosis. Several of these patients have been successfully rescued from hyperammonemia using the protocol described above for urea cycle defects.

435e Inherited Defects of Membrane Transport

Nicola Longo

Specific membrane transporters mediate the passage of a wide variety of substances across cellular membranes. Classes of substrates include amino acids, sugars, cations, anions, vitamins, and water. The number of inherited disorders of membrane transport continues to increase with the identification of new transporters on the plasma membrane or intracellular organelles and the clarification of the molecular basis of diseases with previously unknown pathophysiology. The first transport disorders identified affected the gut or the kidney, but transport processes are now proving essential for the normal function of every organ. Mutations in transporter molecules cause disorders of the heart, muscle, brain, and endocrine and sensory organs (Table 435e-1). Inherited defects impairing the transport of selected amino acids that can present in adults are discussed here as examples of the abnormalities encountered; others are considered elsewhere in this text.

CYSTINURIA

Cystinuria (frequency of 1 in 10,000 to 1 in 15,000) is an autosomal recessive disorder caused by defective transporters in the apical brush border of proximal renal tubule and small intestinal cells. It is characterized by impaired reabsorption and excessive urinary excretion of the dibasic amino acids lysine, arginine, ornithine, and cystine. Because cystine is poorly soluble, its excess excretion predisposes to the formation of renal, ureteral, and bladder stones. Such stones are responsible for the signs and symptoms of the disorder.

There are two variants of cystinuria. Homozygotes for both variants have high urinary excretion of cystine, lysine, arginine, and ornithine. Type I heterozygotes usually have normal urinary amino acid excretion, whereas most non-type I (formerly type II and type III) heterozygotes have moderately increased urinary excretion of each of the four amino acids. The gene for type I cystinuria (*SLC3A1*, chromosome 2p16.3) encodes a membrane glycoprotein. Non-type I cystinuria is caused by mutations in *SLC7A9* (chromosome 19q13) that encodes the b^{0,+} amino acid transporter. The glycoprotein encoded by *SLC3A1* favors the correct processing of the b^{0,+} membrane transporter and explains why mutations in two different genes cause a similar disease.

TABLE 435e-1 GENETIC DISORDERS OF MEMBRANE TRANSPORT (SELECTED EXAMPLES)

Class of Substance and Disorder	Individual Substrates	Tissues Manifesting Transport Defect	Molecular Defect	Major Clinical Manifestations	Inheritance
Amino Acids					
Cystinuria	Cystine, lysine, arginine, ornithine	Proximal renal tubule, jejunal mucosa	Shared dibasic-cystine transporter <i>SLC3A1</i> , <i>SLC7A9</i>	Cystine nephrolithiasis	AR
Dibasic aminoaciduria	Lysine, arginine, ornithine	Proximal renal tubule, jejunal mucosa	Dibasic transporter <i>SLC7A7</i>	Protein intolerance, hyperammonemia, intellectual disability	AR
Hartnup disease	Neutral amino acids	Proximal renal tubule, jejunal mucosa	Neutral amino acid transporter <i>SLC6A19</i>	Constant neutral aminoaciduria, intermittent symptoms of pellagra	AR
Citrullinemia type 2	Aspartate, glutamate, malate	Inner mitochondrial membrane	Mitochondrial aspartate/glutamate carrier 2 <i>SLC25A13</i>	Sudden behavioral changes with stupor, coma, hyperammonemia	AR
Hyperornithinemia, hyperammonemia, homocitrullinuria	Ornithine, citrulline	Inner mitochondrial membrane	Mitochondrial ornithine carrier <i>SLC25A15</i>	Vomiting, lethargy, failure to thrive, intellectual disability, episodic confusion, hyperammonemia, protein intolerance	AR
Histidinuria	Histidine	Proximal renal tubule, jejunal mucosa	Histidine transporter	Intellectual disability	AR
Iminoglycinuria	Glycine, proline, hydroxyproline	Proximal renal tubule, jejunal mucosa	Shared glycine-imino acid transporter <i>SLC36A2</i> , <i>SLC6A19</i> , <i>SLC6A20</i>	None	AR
Dicarboxylic aminoaciduria	Glutamic acid, aspartic acid	Proximal renal tubule, jejunal mucosa	Shared dicarboxylic amino acid transporter	None	Probable AR
Cystinosis	Cystine	Lysosomal membranes	Lysosomal cystine transporter	Renal failure, hypothyroidism, blindness	AR
Hexoses					
Glucose-galactose malabsorption	D-Glucose D-Galactose	Proximal renal tubule, jejunal mucosa	Sodium-dependent glucose/galactose transporter SGLT1	Watery diarrhea on feeding glucose, lactose, sucrose, or galactose	AR
Glucose-transport defect	D-Glucose	Ubiquitous blood-brain barrier	Facilitative glucose transporter GLUT1	Seizures, intellectual disability	AD
Fanconi-Bickel syndrome	D-Glucose	Liver, kidney, pancreas, intestine	Facilitative glucose transporter GLUT2	Growth retardation, rickets, hepatorenal glycogenosis, hypo- and hyperglycemia	AR
Urate					
Hypouricemia	Uric acid	Proximal renal tubule	Urate transporter <i>SLC22A12</i>	Hypouricemia, uric acid urolithiasis	AR
Vitamins					
Thiamine-responsive megaloblastic anemia	Thiamine	Ubiquitous	Thiamine transporter <i>SLC19A2</i>	Megaloblastic anemia, deafness, diabetes mellitus	AR
Other					
Carnitine deficiency	Carnitine	Kidney, muscle, heart	Carnitine transporter OCTN2	Hypoketotic hypoglycemia, cardiomyopathy, sudden death	AR
Creatine deficiency	Creatine	Brain	Creatine transporter <i>SLC6A8</i>	Intellectual disability, seizures, hypotonia	XL

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; XL, X-linked recessive.

Cystine stones account for 1–2% of all urinary tract calculi but are the most common cause of stones in children. Cystinuria homozygotes regularly excrete 2400–7200 μmol (600–1800 mg) of cystine daily. Since the maximum solubility of cystine in the physiologic urinary pH range of 4.5–7.0 is about 1200 $\mu\text{mol/L}$ (300 mg/L), cystine needs to be diluted to 2.5–7 L of water to prevent crystalluria. Stone formation usually manifests in the second or third decade but may occur in the first year of life. Symptoms and signs are those typical of urolithiasis: hematuria, flank pain, renal colic, obstructive uropathy, and infection (**Chap. 342**). Recurrent urolithiasis may lead to progressive renal insufficiency.

Cystinuria is suspected after observing typical hexagonal crystals in the sediment of acidified, concentrated, chilled urine or after performing a urinary nitroprusside test. Quantitative urine amino acid analysis confirms the diagnosis of cystinuria by showing selective overexcretion of cystine, lysine, arginine, and ornithine. Quantitative measurements are important for differentiating heterozygotes from homozygotes and for following free cystine excretion during therapy.

Management is aimed at preventing cystine crystal formation by increasing urinary volume and by maintaining an alkaline urine pH. Fluid ingestion in excess of 4 L/d is essential, and 5–7 L/d is optimal. Urinary cystine concentration should be <1000 $\mu\text{mol/L}$ (250 mg/L). The daily fluid ingestion necessary to maintain this dilution of excreted cystine should be spaced over 24 h, with one-third of the total volume ingested between bedtime and 3 A.M. Cystine solubility rises sharply above pH 7.5, and urinary alkalization (with bicarbonate or potassium citrate) can be therapeutic. Penicillamine (1–3 g/d) and tiopronin (α -mercaptopyrionylglycine, 800–1200 mg/d in four divided doses) undergo sulphydryl-disulfide exchange with cystine to form mixed disulfides. Because these disulfides are much more soluble than cystine, pharmacologic therapy can prevent and promote dissolution of calculi. Penicillamine can have significant side effects and should be reserved for patients who fail to respond to hydration alone or who are in a high-risk category (e.g., one remaining kidney, renal insufficiency). When medical management fails, shock wave lithotripsy, ureteroscopy, and percutaneous nephrolithotomy are effective for most stones. Open urologic surgery is considered for complex staghorn stones or when the patient has concomitant renal or ureteral abnormalities. Occasional patients progress to renal failure and require kidney transplantation.

DIBASIC AMINOACIDURIA (LYSINURIC PROTEIN INTOLERANCE)

This disorder is characterized by a defect in renal tubular reabsorption of the three dibasic amino acids lysine, arginine, and ornithine but not cystine (*lysinuric protein intolerance*). Homozygotes show defective intestinal transport of dibasic amino acids as well as exaggerated renal losses. Lysinuric protein intolerance is most common in Finland (1 in 60,000), southern Italy, and Japan, but is rare elsewhere. The transport defect affects basolateral rather than luminal membrane transport and is associated with impairment of the urea cycle. The defective gene (*SLC7A7*, chromosome 14q11.2) encodes the γ^+ LAT membrane transporter, which associates with the cell-surface glycoprotein 4F2 heavy chain to form the complete sodium-independent transporter γ^+ L.

Manifestations are related to impairment of the urea cycle and to immune dysfunction potentially attributable to nitric oxide overproduction secondary to arginine intracellular trapping. Affected patients present in childhood with hepatosplenomegaly, protein intolerance, and episodic ammonia intoxication. Older patients may present with severe osteoporosis, impairment of kidney function, pulmonary alveolar proteinosis, various autoimmune disorders, and an incompletely characterized immune deficiency. Plasma concentrations of lysine, arginine, and ornithine are reduced, whereas urinary excretion of lysine and orotic acid are increased. Hyperammonemia may develop after the ingestion of protein loads or with infections, probably because of insufficient amounts of arginine and ornithine to maintain proper function of the urea cycle. Therapy consists of dietary protein restriction and supplementation of citrulline (2–8 g/d), a neutral amino acid that fuels the urea cycle when metabolized to arginine and ornithine. Pulmonary disease responds to glucocorticoids or bronchoalveolar

lavage in some patients. Women with lysinuric protein intolerance who become pregnant have an increased risk of anemia, toxemia, and bleeding complications during delivery. These can be minimized by aggressive nutritional therapy and control of blood pressure. Their infants can have intrauterine growth restriction but have normal neurologic function

CITRULLINEMIA TYPE 2 (CITRIN DEFICIENCY)

Citrullinemia type 2 is a recessive condition caused by deficiency of the mitochondrial aspartate-glutamate carrier AGC2 (citrin). A defect in this transporter reduces the availability of cytoplasmic aspartate to combine with citrulline, impairing the urea cycle and decreasing the transfer of reducing equivalents from the cytosol to the mitochondria through the malate–aspartate NADH shuttle. Mutations in the *SLC25A13* gene on chromosome 7q21.3 that encodes for this transporter are rare in Caucasians, but affect about 1:20,000 people with ancestry from Japan, China, and Southeast Asia with variable penetrance.

The disease usually presents with sudden onset between 20 and 50 years of age with recurring episodes of hyperammonemia with associated neuropsychiatric symptoms such as altered mental status, irritability, seizures, or coma resembling hepatic encephalopathy. Some patients might come to medical attention for hypertriglyceridemia, pancreatitis, hepatoma, or fatty liver histologically similar to nonalcoholic steatohepatitis. Without therapy, most patients die with cerebral edema within a few years of onset. Episodes are usually triggered by medications (such as acetaminophen), surgery, alcohol consumption or high sugar intake, the latter conditions causing excess NADH production. NADH is not generated by the metabolism of proteins or fats, and many individuals with citrullinemia type 2 spontaneously prefer foods such as meat, eggs, and fish, and avoid carbohydrates.

Laboratory studies during an acute attack include elevated ammonia, citrulline, and arginine with low or normal levels of glutamine (the latter is usually increased in classic urea cycle defects). The diagnosis is confirmed by demonstrating mutations in the *SLC25A13* gene. Liver transplantation prevents progression of the disease and normalizes biochemical parameters. A diet high in fats and proteins and low in carbohydrates with supplements of arginine and pyruvate is also effective in preventing further episodes, at least in the short term.

HARTNUP DISEASE

Hartnup disease (frequency 1 in 24,000) is an autosomal recessive disorder characterized by pellagra-like skin lesions, variable neurologic manifestations, and neutral and aromatic aminoaciduria. Alanine, serine, threonine, valine, leucine, isoleucine, phenylalanine, tyrosine, tryptophan, glutamine, asparagine, and histidine are excreted in urine in quantities 5–10 times greater than normal, and intestinal transport of these same amino acids is defective. The defective neutral amino acid transporter, B^0 AT1 encoded by the *SLC6A19* gene on chromosome 5p15, requires either collectrin or angiotensin-converting enzyme 2 for surface expression in the kidney and intestine, respectively.

The clinical manifestations result from nutritional deficiency of the essential amino acid tryptophan, caused by its intestinal and renal malabsorption, and of niacin, which derives in part from tryptophan metabolism. Only a small fraction of patients with the chemical findings of this disorder develop a pellagra-like syndrome, implying that manifestations depend on other factors in addition to the transport defect. The diagnosis of Hartnup disease should be suspected in any patient with clinical features of pellagra who does not have a history of dietary niacin deficiency (**Chap. 96e**). The neurologic and psychiatric manifestations range from attacks of cerebellar ataxia to mild emotional lability to frank delirium, and they are usually accompanied by exacerbations of the erythematous, eczematoid skin rash. Fever, sunlight, stress, and sulfonamide therapy provoke clinical relapses. Diagnosis is made by detection of the neutral aminoaciduria, which does not occur in dietary niacin deficiency. Treatment is directed at niacin repletion and includes a high-protein diet and daily nicotinamide supplementation (50–250 mg).

The term *metabolism* is derived from the Greek *metabol*, meaning “to change.” This term encompasses the broad array of chemical pathways that are necessary for normal development and homeostasis. In practice, clinicians generally use the term *metabolism* in reference to energy utilization for anabolism or catabolism. Alternatively, intermediary metabolism describes the myriad cellular pathways that convert energy sources from one form to another (e.g., the citric acid cycle). The emerging field of *metabolomics* is based on the premise that the identification and measurement of metabolic products will enhance our understanding of physiology and disease.

Over the years, the classification of metabolic diseases has extended beyond traditional pathways involved in fuel metabolism to include disorders such as lysosomal storage diseases and connective tissue diseases. Thus, metabolic diseases really reflect disorders of cell biology, and many have a well-defined genetic basis. For example, lysosomal storage diseases (Chap. 432e) result from a variety of genetic defects, usually in a lysosomal enzyme, causing accumulation of a substrate within the lysosome. Certain lipodystrophies and cardiomyopathies can be caused by mutations in lamin A, a structural protein in the nuclear envelope. Membrane defects (Chap. 435e), usually involving transporters of amino acids, sugars, or ions, cause disorders such as cystinuria, Hartnup’s disease, or Wilson’s disease (Chap. 429). Connective tissue diseases (Chap. 427) frequently involve defects in collagen synthesis or structure (osteogenesis imperfecta, Ehlers-Danlos syndrome, Alport’s syndrome) or in other extracellular matrix structural proteins such as fibrillin (Marfan syndrome). Many metabolic disorders originate from defects in enzymes involved in the synthesis or degradation of amino acids, carbohydrates, lipids, purines, or pyrimidines (Chaps. 431e, 433e, and 434e). Lipoprotein disorders (Chap. 421) are caused by defects in a wide array of cellular pathways including membrane receptors (the low-density lipoprotein receptor), enzyme defects (lipoprotein lipase), carrier proteins (apolipoprotein B100), or transporters (ATP-binding cassette transporter ABCA1). In some instances, metabolic abnormalities induce compensatory physiologic responses that reflect the interactions of multiple metabolic pathways. For example, the metabolic syndrome (Chap. 422), which includes a constellation of clinical features (central obesity, hypertriglyceridemia, low high-density lipoprotein cholesterol, hyperglycemia, and hypertension), likely has multiple genetic and environmental origins. Cushing’s syndrome reflects the metabolic effects of excess cortisol on multiple tissues (Chap. 406).

This broader definition results in a plethora of metabolic diseases, numbering in the thousands. Fortunately, comprehensive reference sources exist, such as the Online Metabolic and Molecular Bases of Inherited Disease (OMMBID) (<http://www.ommbid.com/>) and the Online Mendelian Inheritance in Man (OMIM) (<http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=OMIM>). The study of metabolic diseases has been invaluable for advancing our understanding of human genetics by providing insight into principles such as patterns of inheritance, variable expressivity, phenotypic variation, and novel approaches to therapy, including screening programs, blood and organ transplantation, gene therapy, and enzyme replacement (Chap. 82).

This atlas provides a visual survey of selected metabolic disorders with references to the topics elsewhere in the text. The author encourages submission of additional illustrations that might facilitate learning among our peers and thereby enhance the recognition and care of patients with these disorders.



FIGURE 436e-1 “Gauntlet” of pellagra (niacin deficiency). Note indurated, lichenified, pigmented, and scaly skin on the dorsa of the hands. (Source: K Wolff et al: *Fitzpatrick’s Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 96e.



FIGURE 436e-2 Scurvy (vitamin C deficiency). Note perifollicular hemorrhage on the leg. The follicles are often plugged by keratin (perifollicular hyperkeratosis). This eruption occurred in a 46-year-old alcoholic, homeless man who also had bleeding gums and loose teeth. (Source: K Wolff et al: *Fitzpatrick’s Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 96e.



FIGURE 436e-3 Podagra with gouty inflammation of the left first metatarsophalangeal joint. Note swelling and erythema. (From KJ Knoop et al: *The Atlas of Emergency Medicine*, 2nd ed. New York, McGraw-Hill, 2002. Courtesy of Kevin J. Knoop, MD, MS; with permission.) See Chaps. 395 and 431e.

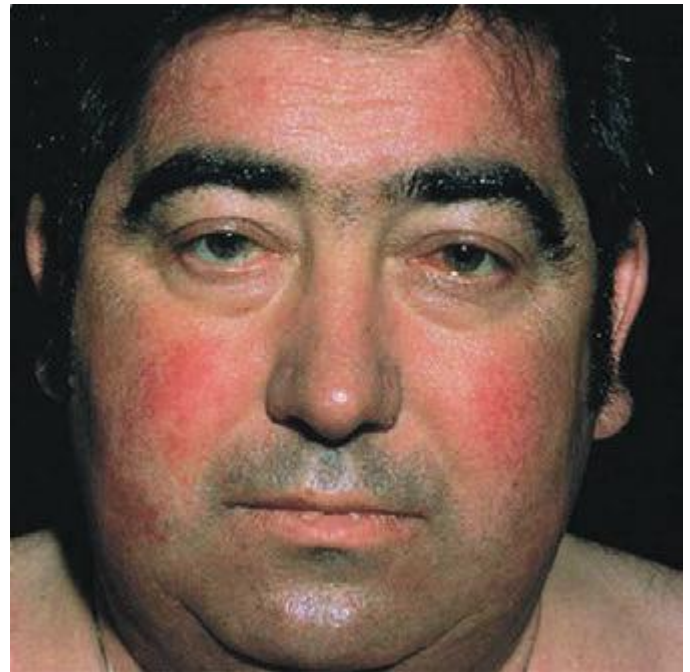


FIGURE 436e-6 Cushing's syndrome. Note plethoric moon facies with erythema and telangiectases of cheek and forehead. The face and neck show increased deposition of fat, which was also seen in the supraclavicular areas (not depicted here). (Source: K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 406.



FIGURE 436e-4 Large tophi of gout located in and around the right knee. (Courtesy of Daniel L. Savitt, MD; with permission.) See Chaps. 395 and 431e.



FIGURE 436e-5 Gouty arthritis of the finger. The finger is an unusual site for gouty arthritis. Examination of the synovial fluid confirmed the diagnosis. (Courtesy of Alan B. Storrow, MD; with permission.) See Chaps. 395 and 431e.



FIGURE 436e-7 Necrobiosis lipoidica diabeticorum. A large symmetric plaque with active tan-pink, well-demarcated, raised, firm borders and a yellow center in the pretibial regions of a 28-year-old diabetic woman is shown. The central parts of the lesion are depressed with atrophic changes of epidermal thinning and telangiectasis against a yellow background. (Source: K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 417.



FIGURE 436e-8 Patient with multiple endocrine neoplasia 2B syndrome. Note the multiple neuromas on the lips and tongue and the marfanoid facies. (Source: DG Gardner, D Shoback, eds: *Greenspan's Basic & Clinical Endocrinology*, 8th ed. New York, McGraw-Hill, 2006, www.accessmedicine.com.) See Chap. 408.



FIGURE 436e-10 Bone scan of a patient with severe Paget's disease of the skull, ribs, spine, pelvis, right femur, and acetabulum. Note localization of bone-seeking isotope (^{99m}Tc -labeled bisphosphonate) in these areas. (Source: DG Gardner, D Shoback, eds: *Greenspan's Basic & Clinical Endocrinology*, 8th ed. New York, McGraw-Hill, 2006, www.accessmedicine.com.) See Chap. 426e.



FIGURE 436e-9 Early and late radiographs of Paget's disease of the tibia of a male patient, taken at 45 (A) and 65 years of age (B). (Source: HB Skinner: *Current Diagnosis & Treatment in Orthopedics*, 4th ed. New York, McGraw-Hill, 2007, www.accessmedicine.com.) See Chap. 426e.



FIGURE 436e-11 Tendinous xanthomas. Large subcutaneous tumors adherent to the Achilles tendons. (Source: K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 421.



A



B

FIGURE 436e-12 Papular eruptive xanthomas. **A.** Multiple, discrete, red-to-yellow papules becoming confluent on the elbow of a white individual with uncontrolled diabetes mellitus; lesions were present on both elbows and buttocks. **B.** Papular eruptive xanthomas on the elbows and lower arms of an African-American patient. (Source: K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 421.

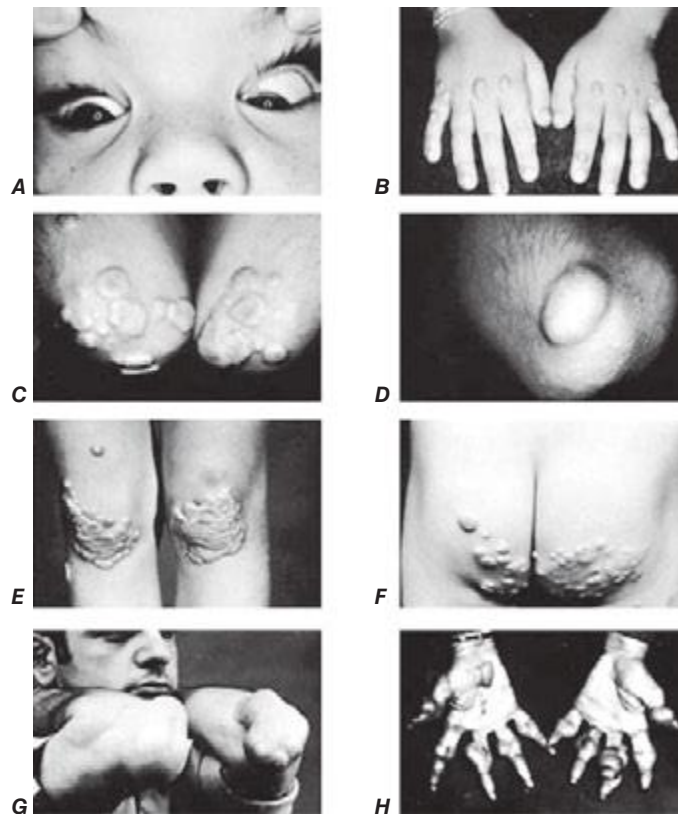


FIGURE 436e-13 Forms of xanthomas and other lipid deposits frequently seen in familial hypercholesterolemia homozygotes. **A.** Arcus corneae. **B, E,** and **F.** Cutaneous planar xanthomas, which usually have a bright orange hue. **C, D,** and **G.** Tuberous xanthomas on the elbows. (Source for panels C and D: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease* online, 8th ed. New York, McGraw-Hill, www.ommbid.com.) **H.** Tendon and tuberous xanthomas. (Panel H reproduced through the courtesy of Dr. A. Khachadurian; with permission.) See Chap. 421.



A

B

FIGURE 436e-14 Examples of xanthomas in type III hyperlipoproteinemic patients. **A.** Tuberous xanthomas of the elbows. **B.** Tuberous xanthomas of the digits and xanthomas of the palmar creases (xanthoma striata palmaris) (arrows). (Courtesy of Dr. Thomas P. Bersot; with permission.) See Chap. 421.



FIGURE 436e-15 A 17-year-old patient with abetalipoproteinemia, with generalized weakness, kyphoscoliosis, and lordosis. (Courtesy of Drs. Peter Herbert, Gerd Assmann, Antonio M. Gotto, Jr., and Donald Fredrickson; with permission.) See Chap. 421.



FIGURE 436e-16 Porphyria cutanea tarda. Periorbital and malar violaceous coloration, hyperpigmentation, and hypertrichosis on the face; bullae, crusts, and scars on the dorsa of the hands. (Source: K Wolff et al: *Fitzpatrick's Color Atlas & Synopsis of Clinical Dermatology*, 5th ed. New York, McGraw-Hill, 2005.) See Chap. 430.



FIGURE 436e-17 Mucopolysaccharidosis type IH (Hurler's syndrome) in a 4-year-old boy. The diagnosis was made at the age of 15 months, at which time he had developmental delay, hepatomegaly, and skeletal involvement. At the time of the picture, the patient had short stature, an enlarged tongue, persistent nasal discharge, stiff joints, and hydrocephalus. Verbal language skills consisted of four or five words. The patient had a severe hearing loss and wore hearing aids. (Source: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease* online, 8th ed. New York, McGraw-Hill, www.ommbid.com.) See Chap. 432e.

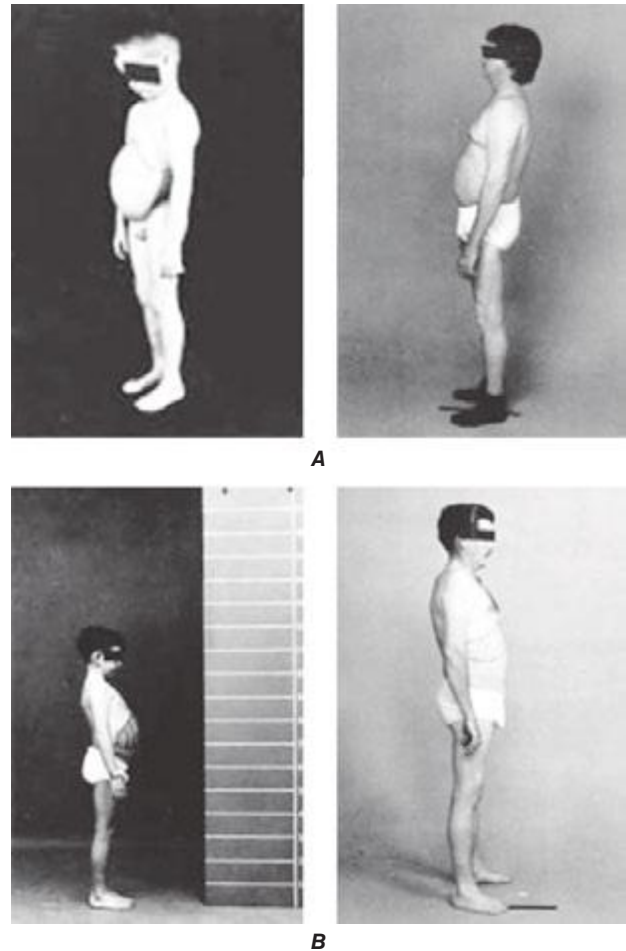


FIGURE 436e-18 Growth and development of two patients with type Ia glycogen storage disease. **A.** Patient at age 7 years and at age 39 years. **B.** Another patient at age 10 years and at age 33 years. Both patients survive despite inadequate treatment of their disease. Note that the abdomen is less protuberant with age. Hypoglycemia also improves with age. In adulthood, however, both patients continue to be short, and both have gout, multiple liver adenomas, and progressive renal disease. (Source: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease* online, 8th ed. New York, McGraw-Hill, www.ommbid.com.) See Chap. 433e.



FIGURE 436e-19 Progressive myopathy in a patient with type IIIa glycogen storage disease. The patient has a debrancher deficiency in both liver and muscle (subtype IIIa). As a child, he had hepatomegaly, hypoglycemia, and growth retardation. After puberty, he no longer had hepatomegaly, and his final height is normal. Note the muscle wasting in the lower legs and both hands at age 44 years of age (*left panel*); this condition progressed to pronounced muscle atrophy at age 53 years (*two right panels*). (Source: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease online*, 8th ed. New York, McGraw-Hill, www.ommbid.com.) See Chap. 433e.

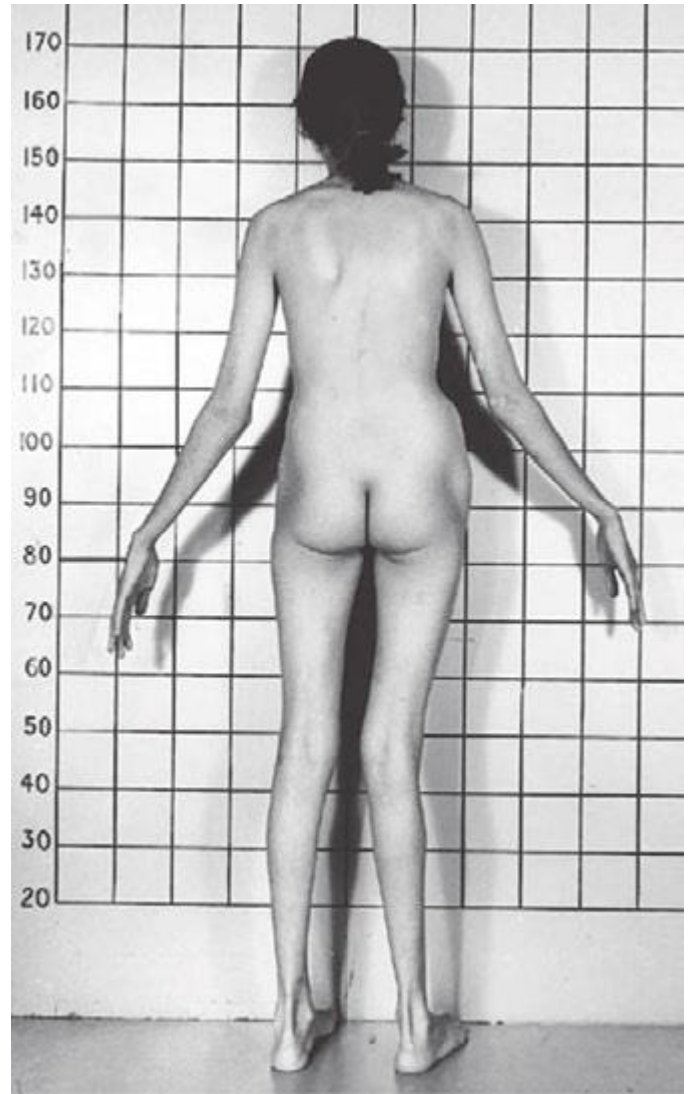


FIGURE 436e-20 Skeletal features of Marfan's syndrome in a 16-year-old girl. Note the long limbs (associated with disproportionately tall stature), long fingers, scoliosis, and genu valgum. (Source: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease online*, 8th ed. New York, McGraw-Hill, www.ommbid.com.) See Chap. 427.



A



B



C

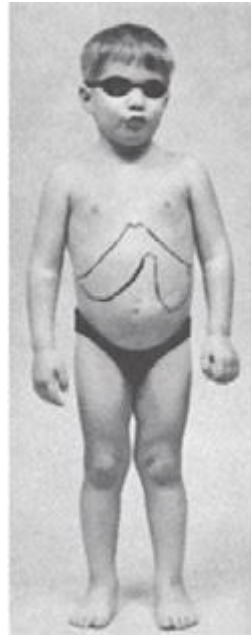


D

FIGURE 436e-21 Marfan's syndrome. **A.** Long, narrow face. **B.** Arachnodactyly and positive wrist sign. **C.** High-arched palate. **D.** Ectopia lentis associated with aortic aneurysm and severe aortic regurgitation in a teenage girl. (Source: V Fuster et al [eds]: *Hurst's The Heart*, 11th ed. New York, McGraw-Hill, 2004, www.accessmedicine.com.) See Chap. 427.



FIGURE 436e-22 Ochronotic pigmentation of the femur of a 56-year-old alkaptonuric patient. (Courtesy of Dr. H. W. Edmonds of the Washington Hospital Center, Washington, DC; with permission.) See Chap. 434e.

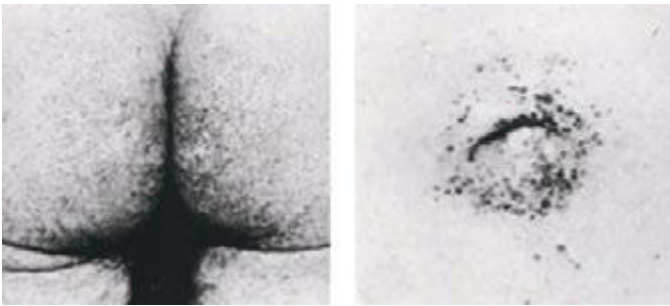


A



B

FIGURE 436e-24 Two patients with type B Niemann-Pick disease (NPD). **A.** A 4.7-year-old patient. (From DS Fredrickson, HR Sloan, in JB Stanbury et al: *The Metabolic Basis of Inherited Disease*, 3rd ed. New York, McGraw-Hill, 1972. Used by permission.) **B.** A 44-year-old patient. See Chap. 432e.



A

B

FIGURE 436e-23 Clusters of angiokeratomas (telangiectases) on the buttocks (**A**) and in the umbilical area (**B**) of a hemizygote with Fabry disease. (Source: CR Scriver et al [eds]: *The Metabolic and Molecular Bases of Inherited Disease* online, 8th ed. New York, McGraw-Hill, www.ommbid.com.) See Chap. 432e.

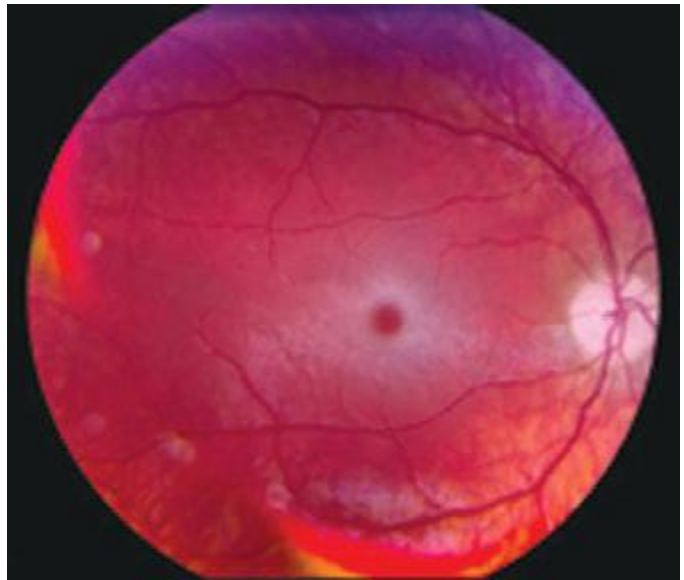


FIGURE 436e-25 "Cherry red" spot in the eye of a Tay-Sachs patient. (From <http://www.nei.nih.gov/resources/eyegene.asp>.) See Chap. 432e.

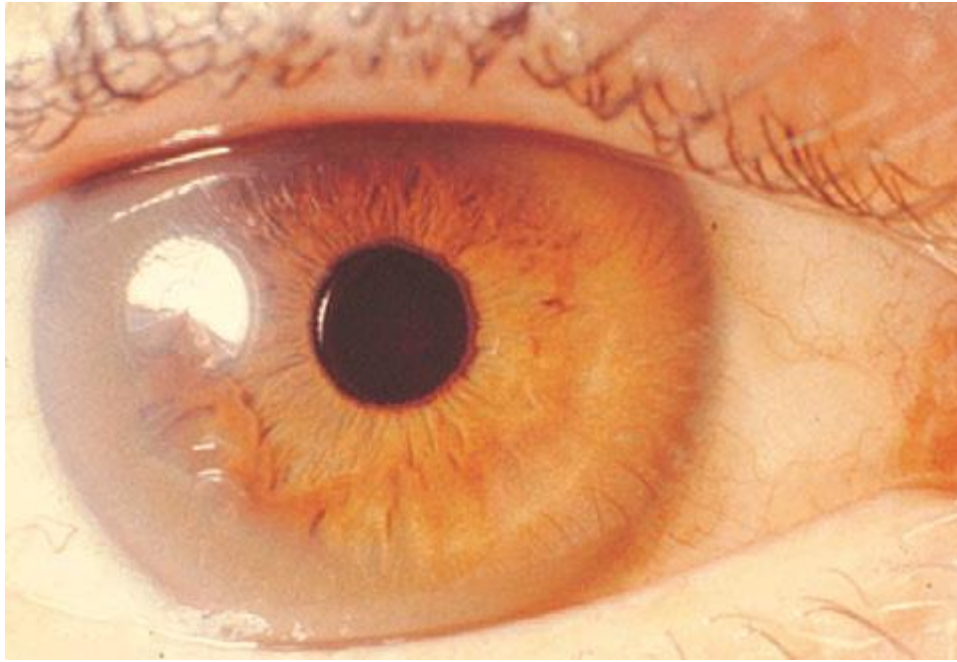


FIGURE 436e-26 Kayser-Fleischer ring. This manifestation develops in Wilson's disease from copper deposition in Descemet's membrane, which produces brownish discoloration of the peripheral cornea. It should not be confused with the yellow-white lipid ring of arcus senilis, which is common in the elderly and occasionally signifies hyperlipidemia, especially when it appears at a young age. (Courtesy of Jonathan C. Horton, MD, PhD; with permission.) See Chap. 429.

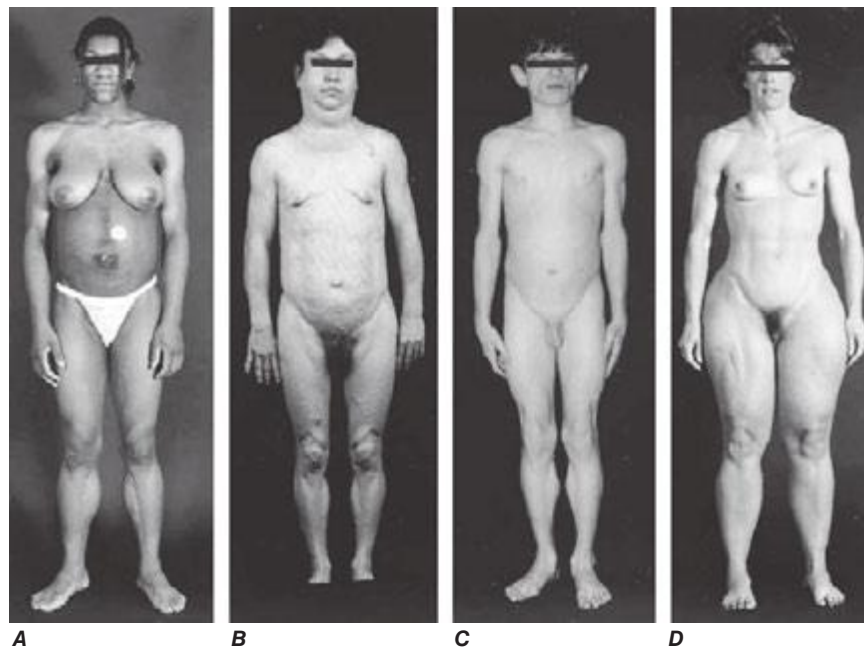


FIGURE 436e-27 Anterior view of patients with different forms of lipodystrophy. **A.** Congenital generalized lipodystrophy: a 16-year-old girl with generalized loss of fat, acromegaloid features, severe acanthosis nigricans affecting the axillae and abdomen, and umbilical hernia. (From A Garg et al: *J Clin Endocrinol Metab* 84:3390, 1999; with permission.) **B.** Familial partial lipodystrophy, Dunnigan variety: a 43-year-old woman with marked loss of subcutaneous fat from both the limbs and the trunk and excess fat deposition in the face, chin, supraclavicular area, and labia majora. (From JM Peters et al: *Nat Genet* 18:292, 1998; with permission.) **C.** Acquired generalized lipodystrophy: a 10-year-old boy who developed generalized loss of fat that also affected the palms and soles after panniculitis at the age of 3 months. **D.** Acquired partial lipodystrophy: a 30-year-old woman with onset of lipodystrophy at age 14 years. Note loss of fat from the face, neck, upper limbs, trunk, and anterior thighs. There is accumulation of excess fat in the hips and other regions of the lower limbs.

This page intentionally left blank

SECTION 1 DIAGNOSIS OF NEUROLOGIC DISORDERS

437 Approach to the Patient with Neurologic Disease

Daniel H. Lowenstein, Joseph B. Martin, Stephen L. Hauser

Neurologic diseases are common and costly. According to estimates by the World Health Organization, neurologic disorders affect over 1 billion people worldwide, constitute 12% of the global burden of disease, and cause 14% of global deaths (Table 437-1). These numbers are only expected to increase as the world's population ages. Most patients with neurologic symptoms seek care from internists and other generalists rather than from neurologists. Because therapies now exist for many neurologic disorders, a skillful approach to diagnosis is essential. Errors commonly result from an overreliance on costly neuroimaging procedures and laboratory tests, which, while useful, do not substitute for an adequate history and examination. The proper approach to the patient with a neurologic illness begins with the patient and focuses the clinical problem first in anatomic and then in pathophysiologic terms; only then should a specific diagnosis be entertained. This method ensures that technology is judiciously applied, a correct diagnosis is established in an efficient manner, and treatment is promptly initiated.

THE NEUROLOGIC METHOD

DEFINE THE ANATOMY

The first priority is to identify the region of the nervous system that is likely to be responsible for the symptoms. Can the disorder be mapped to one specific location, is it multifocal, or is a diffuse process present? Are the symptoms restricted to the nervous system, or do they arise in the context of a systemic illness? Is the problem in the central nervous system (CNS), the peripheral nervous system (PNS), or both? If in the CNS, is the cerebral cortex, basal ganglia, brainstem, cerebellum, or spinal cord responsible? Are the pain-sensitive meninges involved? If in the PNS, could the disorder be located in peripheral nerves and, if so, are motor or sensory nerves primarily affected, or is a lesion in the neuromuscular junction or muscle more likely?

The first clues to defining the anatomic area of involvement appear in the history, and the examination is then directed to confirm or rule out these impressions and to clarify uncertainties. A more detailed examination of a particular region of the CNS or PNS is often indicated.

TABLE 437-1 GLOBAL DISABILITY-ADJUSTED LIFE-YEARS (DALYs) AND NUMBER OF ANNUAL DEATHS FOR SELECTED NEUROLOGIC DISORDERS IN 2010

Disorder	DALYs	Deaths
Low back and neck pain	116,704,000	—
Cerebrovascular diseases	102,232,000	5,874,000
Meningitis and encephalitis	26,540,000	541,000
Migraine	22,362,000	—
Epilepsy	17,429,000	177,000
Dementia	11,349,000	485,000
Parkinson's disease	1,918,000	111,000
% of total DALYs or deaths for all causes that are neurologic	12.0%	13.6%
% change of DALYs for neurologic disorders between 2000 and 2010	51.6%	114.3%

Source: R Lozano et al: Lancet 380: 2095, 2012.

For example, the examination of a patient who presents with a history of ascending paresthesias and weakness should be directed toward deciding, among other things, if the location of the lesion is in the spinal cord or peripheral nerves. Focal back pain, a spinal cord sensory level, and incontinence suggest a spinal cord origin, whereas a stocking-glove pattern of sensory loss suggests peripheral nerve disease; areflexia usually indicates peripheral neuropathy but may also be present with spinal shock in acute spinal cord disorders.

Deciding “where the lesion is” accomplishes the task of limiting the possible etiologies to a manageable, finite number. In addition, this strategy safeguards against making serious errors. Symptoms of recurrent vertigo, diplopia, and nystagmus should not trigger “multiple sclerosis” as an answer (etiology) but “brainstem” or “pons” (location); then a diagnosis of brainstem arteriovenous malformation will not be missed for lack of consideration. Similarly, the combination of optic neuritis and spastic ataxic paraparesis suggests optic nerve and spinal cord disease; multiple sclerosis (MS), CNS syphilis, and vitamin B₁₂ deficiency are treatable disorders that can produce this syndrome. Once the question, “Where is the lesion?” is answered, then the question, “What is the lesion?” can be addressed.

IDENTIFY THE PATHOPHYSIOLOGY

Clues to the pathophysiology of the disease process may also be present in the history. Primary neuronal (gray matter) disorders may present as early cognitive disturbances, movement disorders, or seizures, whereas white matter involvement produces predominantly “long tract” disorders of motor, sensory, visual, and cerebellar pathways. Progressive and symmetric symptoms often have a metabolic or degenerative origin; in such cases lesions are usually not sharply circumscribed. Thus, a patient with paraparesis and a clear spinal cord sensory level is unlikely to have vitamin B₁₂ deficiency as the explanation. A Lhermitte symptom (electric shock-like sensations evoked by neck flexion) is due to ectopic impulse generation in white matter pathways and occurs with demyelination in the cervical spinal cord; among many possible causes, this symptom may indicate MS in a young adult or compressive cervical spondylosis in an older person. Symptoms that worsen after exposure to heat or exercise may indicate conduction block in demyelinated axons, as occurs in MS. A patient with recurrent episodes of diplopia and dysarthria associated with exercise or fatigue may have a disorder of neuromuscular transmission such as myasthenia gravis. Slowly advancing visual scotoma with luminous edges, termed *fortification spectra*, indicates spreading cortical depression, typically with migraine.

THE NEUROLOGIC HISTORY

Attention to the description of the symptoms experienced by the patient and substantiated by family members and others often permits an accurate localization and determination of the probable cause of the complaints, even before the neurologic examination is performed. The history also helps to bring a focus to the neurologic examination that follows. Each complaint should be pursued as far as possible to elucidate the location of the lesion, the likely underlying pathophysiology, and potential etiologies. For example, a patient complains of weakness of the right arm. What are the associated features? Does the patient have difficulty with brushing hair or reaching upward (proximal) or buttoning buttons or opening a twist-top bottle (distal)? Negative associations may also be crucial. A patient with a right hemiparesis without a language deficit likely has a lesion (internal capsule, brainstem, or spinal cord) different from that of a patient with a right hemiparesis and aphasia (left hemisphere). Other pertinent features of the history include the following:

1. *Temporal course of the illness.* It is important to determine the precise time of appearance and rate of progression of the symptoms experienced by the patient. The rapid onset of a neurologic complaint, occurring within seconds or minutes, usually indicates a vascular event, a seizure, or migraine. The onset of sensory symptoms located in one extremity that spread over a few seconds to adjacent portions of that extremity and then to the other regions of the body suggests a seizure. A more gradual onset and less well-localized symptoms point to the possibility of a transient ischemic attack (TIA). A similar but slower temporal march of symptoms accompanied by headache, nausea, or visual disturbance suggests migraine. The presence of “positive” sensory symptoms (e.g., tingling or sensations that are difficult to describe) or involuntary motor movements suggests a seizure; in contrast, transient loss of function (negative symptoms) suggests a TIA. A stuttering onset where symptoms appear, stabilize, and then progress over hours or days also suggests cerebrovascular disease; an additional history of transient remission or regression indicates that the process is more likely due to ischemia rather than hemorrhage. A gradual evolution of symptoms over hours or days suggests a toxic, metabolic, infectious, or inflammatory process. Progressing symptoms associated with the systemic manifestations of fever, stiff neck, and altered level of consciousness imply an infectious process. Relapsing and remitting symptoms involving different levels of the nervous system suggest MS or other inflammatory processes. Slowly progressive symptoms without remissions are characteristic of neurodegenerative disorders, chronic infections, gradual intoxications, and neoplasms.
2. *Patients’ descriptions of the complaint.* The same words often mean different things to different patients. “Dizziness” may imply impending syncope, a sense of disequilibrium, or true spinning vertigo. “Numbness” may mean a complete loss of feeling, a positive sensation such as tingling, or even weakness. “Blurred vision” may be used to describe unilateral visual loss, as in transient monocular blindness, or diplopia. The interpretation of the true meaning of the words used by patients to describe symptoms obviously becomes even more complex when there are differences in primary languages and cultures.
3. *Corroboration of the history by others.* It is almost always helpful to obtain additional information from family, friends, or other observers to corroborate or expand the patient’s description. Memory loss, aphasia, loss of insight, intoxication, and other factors may impair the patient’s capacity to communicate normally with the examiner or prevent openness about factors that have contributed to the illness. Episodes of loss of consciousness necessitate that details be sought from observers to ascertain precisely what has happened during the event.
4. *Family history.* Many neurologic disorders have an underlying genetic component. The presence of a Mendelian disorder, such as Huntington’s disease or Charcot-Marie-Tooth neuropathy, is often obvious if family data are available. More detailed questions about family history are often necessary in polygenic disorders such as MS, migraine, and many types of epilepsy. It is important to elicit family history about all illnesses, in addition to neurologic and psychiatric disorders. A familial propensity to hypertension or heart disease is relevant in a patient who presents with a stroke. There are numerous inherited neurologic diseases that are associated with multisystem manifestations that may provide clues to the correct diagnosis (e.g., neurofibromatosis, Wilson’s disease, mitochondrial disorders).
5. *Medical illnesses.* Many neurologic diseases occur in the context of systemic disorders. Diabetes mellitus, hypertension, and abnormalities of blood lipids predispose to cerebrovascular disease. A solitary mass lesion in the brain may be an abscess in a patient with valvular heart disease, a primary hemorrhage in a patient with a coagulopathy, a lymphoma or toxoplasmosis in a patient with AIDS, or a metastasis in a patient with underlying cancer. Patients with malignancy may also present with a neurologic paraneoplastic syndrome ([Chap. 122](#)) or complications from chemotherapy or radiotherapy. Marfan’s syndrome and related collagen disorders predispose to dissection of the cranial arteries and aneurysmal subarachnoid hemorrhage; the latter may also occur with polycystic kidney disease. Various neurologic disorders occur with dysthyroid states or other endocrinopathies. It is especially important to look for the presence of systemic diseases in patients with peripheral neuropathy. Most patients with coma in a hospital setting have a metabolic, toxic, or infectious cause.
6. *Drug use and abuse and toxin exposure.* It is essential to inquire about the history of drug use, both prescribed and illicit. Sedatives, antidepressants, and other psychoactive medications are frequently associated with acute confusional states, especially in the elderly. Aminoglycoside antibiotics may exacerbate symptoms of weakness in patients with disorders of neuromuscular transmission, such as myasthenia gravis, and may cause dizziness secondary to ototoxicity. Vincristine and other antineoplastic drugs can cause peripheral neuropathy, and immunosuppressive agents such as cyclosporine can produce encephalopathy. Excessive vitamin ingestion can lead to disease; examples include vitamin A and pseudotumor cerebri or pyridoxine and peripheral neuropathy. Many patients are unaware that over-the-counter sleeping pills, cold preparations, and diet pills are actually drugs. Alcohol, the most prevalent neurotoxin, is often not recognized as such by patients, and other drugs of abuse such as cocaine and heroin can cause a wide range of neurologic abnormalities. A history of environmental or industrial exposure to neurotoxins may provide an essential clue; consultation with the patient’s coworkers or employer may be required.
7. *Formulating an impression of the patient.* Use the opportunity while taking the history to form an impression of the patient. Is the information forthcoming, or does it take a circuitous course? Is there evidence of anxiety, depression, or hypochondriasis? Are there any clues to problems with language, memory, insight, comportment, or behavior? The neurologic assessment begins as soon as the patient comes into the room and the first introduction is made.

THE NEUROLOGIC EXAMINATION

The neurologic examination is challenging and complex; it has many components and includes a number of skills that can be mastered only through repeated use of the same techniques on a large number of individuals with and without neurologic disease. Mastery of the complete neurologic examination is usually important only for physicians in neurology and associated specialties. However, knowledge of the basics of the examination, especially those components that are effective in screening for neurologic dysfunction, is essential for all clinicians, especially generalists.

There is no single, universally accepted sequence of the examination that must be followed, but most clinicians begin with assessment of mental status followed by the cranial nerves, motor system, reflexes, sensory system, coordination, and gait. Whether the examination is basic or comprehensive, it is essential that it be performed in an orderly and systematic fashion to avoid errors and serious omissions. Thus, the best way to learn and gain expertise in the examination is to choose one’s own approach and practice it frequently and do it in the same exact sequence each time.

The detailed description that follows describes the more commonly used parts of the neurologic examination, with a particular emphasis on the components that are considered most helpful for the assessment of common neurologic problems. Each section also includes a brief description of the minimal examination necessary to adequately screen for abnormalities in a patient who has no symptoms suggesting neurologic dysfunction. A screening examination done in this way can be completed in 3–5 min.

Several additional points about the examination are worth noting. First, in recording observations, it is important to describe what is found rather than to apply a poorly defined medical term (e.g., “patient groans to sternal rub” rather than “obtunded”). Second, subtle CNS abnormalities are best detected by carefully comparing a patient’s performance on tasks that require simultaneous activation of both cerebral hemispheres (e.g., eliciting a pronator drift of an outstretched arm with the eyes closed; extinction on one side of bilaterally applied light touch, also with eyes closed; or decreased arm swing or a slight

asymmetry when walking). Third, if the patient's complaint is brought on by some activity, reproduce the activity in the office. If the complaint is of dizziness when the head is turned in one direction, have the patient do this and also look for associated signs on examination (e.g., nystagmus or dysmetria). If pain occurs after walking two blocks, have the patient leave the office and walk this distance and immediately return, and repeat the relevant parts of the examination. Finally, the use of tests that are individually tailored to the patient's problem can be of value in assessing changes over time. Tests of walking a 7.5-m (25-ft) distance (normal, 5–6 s; note assistance, if any), repetitive finger or toe tapping (normal, 20–25 taps in 5 s), or handwriting are examples.

MENTAL STATUS EXAMINATION

- *The bare minimum: During the interview, look for difficulties with communication and determine whether the patient has recall and insight into recent and past events.*

The mental status examination is under way as soon as the physician begins observing and speaking with the patient. If the history raises any concern for abnormalities of higher cortical function or if cognitive problems are observed during the interview, then detailed testing of the mental status is indicated. The patient's ability to understand the language used for the examination, cultural background, educational experience, sensory or motor problems, or comorbid conditions need to be factored into the applicability of the tests and interpretation of results.

The Folstein mini-mental status examination (MMSE) is a standardized screening examination of cognitive function that is extremely easy to administer and takes <10 min to complete. Using age-adjusted values for defining normal performance, the test is ~85% sensitive and 85% specific for making the diagnosis of dementia that is moderate or severe, especially in educated patients. When there is sufficient time available, the MMSE is one of the best methods for documenting the current mental status of the patient, and this is especially useful as a baseline assessment to which future scores of the MMSE can be compared.

Individual elements of the mental status examination can be subdivided into level of consciousness, orientation, speech and language, memory, fund of information, insight and judgment, abstract thought, and calculations.

Level of consciousness is the patient's relative state of awareness of the self and the environment, and ranges from fully awake to comatose. When the patient is not fully awake, the examiner should describe the responses to the minimum stimulus necessary to elicit a reaction, ranging from verbal commands to a brief, painful stimulus such as a squeeze of the trapezius muscle. Responses that are directed toward the stimulus and signify some degree of intact cerebral function (e.g., opening the eyes and looking at the examiner or reaching to push away a painful stimulus) must be distinguished from reflex responses of a spinal origin (e.g., triple flexion response—flexion at the ankle, knee, and hip in response to a painful stimulus to the foot).

Orientation is tested by asking the person to state his or her name, location, and time (day of the week and date); time is usually the first to be affected in a variety of conditions.

Speech is assessed by observing articulation, rate, rhythm, and prosody (i.e., the changes in pitch and accentuation of syllables and words).

Language is assessed by observing the content of the patient's verbal and written output, response to spoken commands, and ability to read. A typical testing sequence is to ask the patient to name successively more detailed components of clothing, a watch, or a pen; repeat the phrase "No ifs, ands, or buts"; follow a three-step, verbal command; write a sentence; and read and respond to a written command.

Memory should be analyzed according to three main time scales: (1) immediate memory is assessed by saying a list of three items and having the patient repeat the list immediately; (2) short-term memory is tested by asking the patient to recall the same three items 5 and 15 min later; and (3) long-term memory is evaluated by determining how well the patient is able to provide a coherent chronologic history of his or her illness or personal events.

Fund of information is assessed by asking questions about major historic or current events, with special attention to educational level and life experiences.

Abnormalities of *insight and judgment* are usually detected during the patient interview; a more detailed assessment can be elicited by asking the patient to describe how he or she would respond to situations having a variety of potential outcomes (e.g., "What would you do if you found a wallet on the sidewalk?").

Abstract thought can be tested by asking the patient to describe similarities between various objects or concepts (e.g., apple and orange, desk and chair, poetry and sculpture) or to list items having the same attributes (e.g., a list of four-legged animals).

Calculation ability is assessed by having the patient carry out a computation that is appropriate to the patient's age and education (e.g., serial subtraction of 7 from 100 or 3 from 20; or word problems involving simple arithmetic).

CRANIAL NERVE EXAMINATION

- *The bare minimum: Check the fundi, visual fields, pupil size and reactivity, extraocular movements, and facial movements.*

The cranial nerves (CN) are best examined in numerical order, except for grouping together CN III, IV, and VI because of their similar function.

CN I (Olfactory) Testing is often omitted unless there is suspicion for inferior frontal lobe disease (e.g., meningioma). With eyes closed, ask the patient to sniff a mild stimulus such as toothpaste or coffee and identify the odorant.

CN II (Optic) Check visual acuity (with eyeglasses or contact lens correction) using a Snellen chart or similar tool. Test the visual fields by confrontation, i.e., by comparing the patient's visual fields to your own. As a screening test, it is usually sufficient to examine the visual fields of both eyes simultaneously; individual eye fields should be tested if there is any reason to suspect a problem of vision by the history or other elements of the examination, or if the screening test reveals an abnormality. Face the patient at a distance of approximately 0.6–1.0 m (2–3 ft) and place your hands at the periphery of your visual fields in the plane that is equidistant between you and the patient. Instruct the patient to look directly at the center of your face and to indicate when and where he or she sees one of your fingers moving. Beginning with the two inferior quadrants and then the two superior quadrants, move your index finger of the right hand, left hand, or both hands simultaneously and observe whether the patient detects the movements. A single small-amplitude movement of the finger is sufficient for a normal response. Focal perimetry and tangent screen examinations should be used to map out visual field defects fully or to search for subtle abnormalities. Optic fundi should be examined with an ophthalmoscope, and the color, size, and degree of swelling or elevation of the optic disc noted, as well as the color and texture of the retina. The retinal vessels should be checked for size, regularity, arteriovenous nicking at crossing points, hemorrhage, exudates, etc.

CN III, IV, VI (Oculomotor, Trochlear, Abducens) Describe the size and shape of pupils and reaction to light and accommodation (i.e., as the eyes converge while following your finger as it moves toward the bridge of the nose). To check extraocular movements, ask the patient to keep his or her head still while tracking the movement of the tip of your finger. Move the target slowly in the horizontal and vertical planes; observe any paresis, nystagmus, or abnormalities of smooth pursuit (saccades, oculomotor ataxia, etc.). If necessary, the relative position of the two eyes, both in primary and multidirectional gaze, can be assessed by comparing the reflections of a bright light off both pupils. However, in practice it is typically more useful to determine whether the patient describes diplopia in any direction of gaze; true diplopia should almost always resolve with one eye closed. Horizontal nystagmus is best assessed at 45° and not at extreme lateral gaze (which is uncomfortable for the patient); the target must often be held at the lateral position for at least a few seconds to detect an abnormality.

CN V (Trigeminal) Examine sensation within the three territories of the branches of the trigeminal nerve (ophthalmic, maxillary, and mandibular) on each side of the face. As with other parts of the sensory examination, testing of two sensory modalities derived from different anatomic pathways (e.g., light touch and temperature) is sufficient

for a screening examination. Testing of other modalities, the corneal reflex, and the motor component of CN V (jaw clench—masseter muscle) is indicated when suggested by the history.

CN VII (Facial) Look for facial asymmetry at rest and with spontaneous movements. Test eyebrow elevation, forehead wrinkling, eye closure, smiling, and cheek puff. Look in particular for differences in the lower versus upper facial muscles; weakness of the lower two-thirds of the face with preservation of the upper third suggests an upper motor neuron lesion, whereas weakness of an entire side suggests a lower motor neuron lesion.

CN VIII (Vestibulocochlear) Check the patient's ability to hear a finger rub or whispered voice with each ear. Further testing for air versus mastoid bone conduction (Rinne) and lateralization of a 512-Hz tuning fork placed at the center of the forehead (Weber) should be done if an abnormality is detected by history or examination. Any suspected problem should be followed up with formal audiometry. **For further discussion of assessing vestibular nerve function in the setting of dizziness, hearing loss, or coma, see Chaps. 28, 43, and 328, respectively.**

CN IX, X (Glossopharyngeal, Vagus) Observe the position and symmetry of the palate and uvula at rest and with phonation ("aah"). The pharyngeal ("gag") reflex is evaluated by stimulating the posterior pharyngeal wall on each side with a sterile, blunt object (e.g., tongue blade), but the reflex is often absent in normal individuals.

CN XI (Spinal Accessory) Check shoulder shrug (trapezius muscle) and head rotation to each side (sternocleidomastoid) against resistance.

CN XII (Hypoglossal) Inspect the tongue for atrophy or fasciculations, position with protrusion, and strength when extended against the inner surface of the cheeks on each side.

MOTOR EXAMINATION

- *The bare minimum: Look for muscle atrophy and check extremity tone. Assess upper extremity strength by checking for pronator drift and strength of wrist or finger extensors. Assess lower extremity strength by checking strength of the toe extensors and having the patient walk normally and on heels and toes.*

The motor examination includes observations of muscle appearance, tone, and strength. Although gait is in part a test of motor function, it is usually evaluated separately at the end of the examination.

Appearance Inspect and palpate muscle groups under good light and with the patient in a comfortable and symmetric position. Check for muscle fasciculations, tenderness, and atrophy or hypertrophy. Involuntary movements may be present at rest (e.g., tics, myoclonus, choreoathetosis), during maintained posture (pill-rolling tremor of Parkinson's disease), or with voluntary movements (intention tremor of cerebellar disease or familial tremor).

Tone Muscle tone is tested by measuring the resistance to passive movement of a relaxed limb. Patients often have difficulty relaxing during this procedure, so it is useful to distract the patient to minimize active movements. In the upper limbs, tone is assessed by rapid pronation and supination of the forearm and flexion and extension at the wrist. In the lower limbs, while the patient is supine the examiner's hands are placed behind the knees and rapidly raised; with normal tone, the ankles drag along the table surface for a variable distance before rising, whereas increased tone results in an immediate lift of the heel off the surface. Decreased tone is most commonly due to lower motor neuron or peripheral nerve disorders. Increased tone may be evident as spasticity (resistance determined by the angle and velocity of motion; corticospinal tract disease), rigidity (similar resistance in all angles of motion; extrapyramidal disease), or paratonia (fluctuating changes in resistance; frontal lobe pathways or normal difficulty in relaxing). Cogwheel rigidity, in which passive motion elicits jerky interruptions in resistance, is seen in parkinsonism.

Strength Testing for pronator drift is an extremely useful method for screening upper limb weakness. The patient is asked to hold both arms fully extended and parallel to the ground with eyes closed. This position should be maintained for ~10 s; any flexion at the elbow or fingers or

pronation of the forearm, especially if asymmetric, is a sign of potential weakness. Muscle strength is further assessed by having the patient exert maximal effort for the particular muscle or muscle group being tested. It is important to isolate the muscles as much as possible, i.e., hold the limb so that only the muscles of interest are active. It is also helpful to palpate accessible muscles as they contract. Grading muscle strength and evaluating the patient's effort is an art that takes time and practice. Muscle strength is traditionally graded using the following scale:

- 0 = no movement
- 1 = flicker or trace of contraction but no associated movement at a joint
- 2 = movement with gravity eliminated
- 3 = movement against gravity but not against resistance
- 4- = movement against a mild degree of resistance
- 4 = movement against moderate resistance
- 4+ = movement against strong resistance
- 5 = full power

However, in many cases, it is more practical to use the following terms:

- Paralysis = no movement
- Severe weakness = movement with gravity eliminated
- Moderate weakness = movement against gravity but not against mild resistance
- Mild weakness = movement against moderate resistance
- Full strength

Noting the pattern of weakness is as important as assessing the magnitude of weakness. Unilateral or bilateral weakness of the upper limb extensors and lower limb flexors ("pyramidal weakness") suggests a lesion of the pyramidal tract, bilateral proximal weakness suggests myopathy, and bilateral distal weakness suggests peripheral neuropathy.

REFLEX EXAMINATION

- *The bare minimum: Check the biceps, patellar, and Achilles reflexes.*

Muscle Stretch Reflexes Those that are typically assessed include the biceps (C5, C6), brachioradialis (C5, C6), and triceps (C7, C8) reflexes in the upper limbs and the patellar or quadriceps (L3, L4) and Achilles (S1, S2) reflexes in the lower limbs. The patient should be relaxed and the muscle positioned midway between full contraction and extension. Reflexes may be enhanced by asking the patient to voluntarily contract other, distant muscle groups (Jendrassik maneuver). For example, upper limb reflexes may be reinforced by voluntary teeth-clenching, and the Achilles reflex by hooking the flexed fingers of the two hands together and attempting to pull them apart. For each reflex tested, the two sides should be tested sequentially, and it is important to determine the smallest stimulus required to elicit a reflex rather than the maximum response. Reflexes are graded according to the following scale:

- 0 = absent
- 1 = present but diminished
- 2 = normoactive
- 3 = exaggerated
- 4 = clonus

Cutaneous Reflexes The plantar reflex is elicited by stroking, with a noxious stimulus such as a tongue blade, the lateral surface of the sole of the foot beginning near the heel and moving across the ball of the foot to the great toe. The normal reflex consists of plantar flexion of the toes. With upper motor neuron lesions above the S1 level of the spinal cord, a paradoxical extension of the toe is observed, associated with fanning and extension of the other toes (termed an *extensor plantar response*, or *Babinski sign*). However, despite its popularity, the reliability and validity of the Babinski sign for identifying upper motor neuron weakness is limited—it is far more useful to rely on tests of tone, strength, stretch reflexes, and coordination. Superficial abdominal reflexes are elicited by gently stroking the abdominal surface near the umbilicus in a diagonal fashion with a sharp object (e.g., the wooden end of a cotton-tipped swab) and observing the movement of

the umbilicus. Normally, the umbilicus will pull toward the stimulated quadrant. With upper motor neuron lesions, these reflexes are absent. They are most helpful when there is preservation of the upper (spinal cord level T9) but not lower (T12) abdominal reflexes, indicating a spinal lesion between T9 and T12, or when the response is asymmetric. Other useful cutaneous reflexes include the cremasteric (ipsilateral elevation of the testicle following stroking of the medial thigh; mediated by L1 and L2) and anal (contraction of the anal sphincter when the perianal skin is scratched; mediated by S2, S3, S4) reflexes. It is particularly important to test for these reflexes in any patient with suspected injury to the spinal cord or lumbosacral roots.

Primitive Reflexes With disease of the frontal lobe pathways, several primitive reflexes not normally present in the adult may appear. The suck response is elicited by lightly touching with a tongue blade the center of the lips, and the root response the corner of the lips; the patient will move the lips to suck or root in the direction of the stimulus. The grasp reflex is elicited by touching the palm between the thumb and index finger with the examiner's fingers; a positive response is a forced grasp of the examiner's hand. In many instances, stroking the back of the hand will lead to its release. The palmomental response is contraction of the mentalis muscle (chin) ipsilateral to a scratch stimulus diagonally applied to the palm.

SENSORY EXAMINATION

- *The bare minimum: Ask whether the patient can feel light touch and the temperature of a cool object in each distal extremity. Check double simultaneous stimulation using light touch on the hands. Perform the Romberg maneuver.*

Evaluating sensation is usually the most unreliable part of the examination because it is subjective and is difficult to quantify. In the compliant and discerning patient, the sensory examination can be extremely helpful for the precise localization of a lesion. With patients who are uncooperative or lack an understanding of the tests, it may be useless. The examination should be focused on the suspected lesion. For example, in spinal cord, spinal root, or peripheral nerve abnormalities, all major sensory modalities should be tested while looking for a pattern consistent with a spinal level and dermatomal or nerve distribution. In patients with lesions at or above the brainstem, screening the primary sensory modalities in the distal extremities along with tests of "cortical" sensation is usually sufficient.

The five primary sensory modalities—light touch, pain, temperature, vibration, and joint position—are tested in each limb. Light touch is assessed by stimulating the skin with single, very gentle touches of the examiner's finger or a wisp of cotton. Pain is tested using a new pin, and temperature is assessed using a metal object (e.g., tuning fork) that has been immersed in cold and warm water. Vibration is tested using a 128-Hz tuning fork applied to the distal phalanx of the great toe or index finger just below the nail bed. By placing a finger on the opposite side of the joint being tested, the examiner compares the patient's threshold of vibration perception with his or her own. For joint position testing, the examiner grasps the digit or limb laterally and distal to the joint being assessed; small 1- to 2-mm excursions can usually be sensed. The Romberg maneuver is primarily a test of proprioception. The patient is asked to stand with the feet as close together as necessary to maintain balance while the eyes are open, and the eyes are then closed. A loss of balance with the eyes closed is an abnormal response.

"Cortical" sensation is mediated by the parietal lobes and represents an integration of the primary sensory modalities; testing cortical sensation is only meaningful when primary sensation is intact. Double simultaneous stimulation is especially useful as a screening test for cortical function; with the patient's eyes closed, the examiner lightly touches one or both hands and asks the patient to identify the stimuli. With a parietal lobe lesion, the patient may be unable to identify the stimulus on the contralateral side when both hands are touched. Other modalities relying on the parietal cortex include the discrimination of two closely placed stimuli as separate (two-point discrimination), identification of an object by touch and manipulation alone (stereognosis), and the identification of numbers or letters written on the skin surface (graphesthesia).

COORDINATION EXAMINATION

- *The bare minimum: Observe the patient at rest and during spontaneous movements. Test rapid alternating movements of the hands and feet and finger to nose.*

Coordination refers to the orchestration and fluidity of movements. Even simple acts require cooperation of agonist and antagonist muscles, maintenance of posture, and complex servomechanisms to control the rate and range of movements. Part of this integration relies on normal function of the cerebellar and basal ganglia systems. However, coordination also requires intact muscle strength and kinesthetic and proprioceptive information. Thus, if the examination has disclosed abnormalities of the motor or sensory systems, the patient's coordination should be assessed with these limitations in mind.

Rapid alternating movements in the upper limbs are tested separately on each side by having the patient make a fist, partially extend the index finger, and then tap the index finger on the distal thumb as quickly as possible. In the lower limb, the patient rapidly taps the foot against the floor or the examiner's hand. Finger-to-nose testing is primarily a test of cerebellar function; the patient is asked to touch his or her index finger repetitively to the nose and then to the examiner's outstretched finger, which moves with each repetition. A similar test in the lower extremity is to have the patient raise the leg and touch the examiner's finger with the great toe. Another cerebellar test in the lower limbs is the heel-knee-shin maneuver; in the supine position the patient is asked to slide the heel of each foot from the knee down the shin of the other leg. For all these movements, the accuracy, speed, and rhythm are noted.

GAIT EXAMINATION

- *The bare minimum: Observe the patient while walking normally, on the heels and toes, and along a straight line.*

Watching the patient walk is the most important part of the neurologic examination. Normal gait requires that multiple systems—including strength, sensation, and coordination—function in a highly integrated fashion. Unexpected abnormalities may be detected that prompt the examiner to return in more detail to other aspects of the examination. The patient should be observed while walking and turning normally, walking on the heels, walking on the toes, and walking heel-to-toe along a straight line. The examination may reveal decreased arm swing on one side (corticospinal tract disease), a stooped posture and short-stepped gait (parkinsonism), a broad-based unstable gait (ataxia), scissoring (spasticity), or a high-stepped, slapping gait (posterior column or peripheral nerve disease), or the patient may appear to be stuck in place (apraxia with frontal lobe disease).

NEUROLOGIC DIAGNOSIS

The clinical data obtained from the history and examination are interpreted to arrive at an anatomic localization that best explains the clinical findings (Table 437-2), to narrow the list of diagnostic possibilities, and to select the laboratory tests most likely to be informative. The laboratory assessment may include (1) serum electrolytes; complete blood count; and renal, hepatic, endocrine, and immune studies; (2) cerebrospinal fluid examination; (3) focused neuroimaging studies (Chap. 440e); or (4) electrophysiologic studies (Chap. 442e). The anatomic localization, mode of onset and course of illness, other medical data, and laboratory findings are then integrated to establish an etiologic diagnosis.

The neurologic examination may be normal even in patients with a serious neurologic disease, such as seizures, chronic meningitis, or a TIA. A comatose patient may arrive with no available history, and in such cases, the approach is as described in Chap. 328. In other patients, an inadequate history may be overcome by a succession of examinations from which the course of the illness can be inferred. In perplexing cases it is useful to remember that uncommon presentations of common diseases are more likely than rare etiologies. Thus, even in tertiary care settings, multiple strokes are usually due to emboli and not vasculitis, and dementia with myoclonus is usually Alzheimer's disease and not a prion disorder or a paraneoplastic illness. Finally, the most important task of a primary care physician

TABLE 437-2 FINDINGS HELPFUL FOR LOCALIZATIONS WITHIN THE NERVOUS SYSTEM

	Signs
Cerebrum	Abnormal mental status or cognitive impairment Seizures Unilateral weakness ^a and sensory abnormalities including head and limbs Visual field abnormalities Movement abnormalities (e.g., diffuse incoordination, tremor, chorea)
Brainstem	Isolated cranial nerve abnormalities (single or multiple) "Crossed" weakness ^a and sensory abnormalities of head and limbs, e.g., weakness of right face and left arm and leg
Spinal cord	Back pain or tenderness Weakness ^a and sensory abnormalities sparing the head Mixed upper and lower motor neuron findings Sensory level Sphincter dysfunction
Spinal roots	Radiating limb pain Weakness ^b or sensory abnormalities following root distribution (see Figs. 31-2 and 31-3) Loss of reflexes
Peripheral nerve	Mid or distal limb pain Weakness ^b or sensory abnormalities following nerve distribution (see Figs. 31-2 and 31-3) "Stocking or glove" distribution of sensory loss Loss of reflexes
Neuromuscular junction	Bilateral weakness including face (ptosis, diplopia, dysphagia) and proximal limbs Increasing weakness with exertion Sparing of sensation
Muscle	Bilateral proximal or distal weakness Sparing of sensation

^aWeakness along with other abnormalities having an "upper motor neuron" pattern, i.e., spasticity, weakness of extensors > flexors in the upper extremity and flexors > extensors in the lower extremity, and hyperreflexia. ^bWeakness along with other abnormalities having a "lower motor neuron" pattern, i.e., flaccidity and hyporeflexia.

faced with a patient who has a new neurologic complaint is to assess the urgency of referral to a specialist. Here, the imperative is to rapidly identify patients likely to have nervous system infections, acute strokes, and spinal cord compression or other treatable mass lesions and arrange for immediate care.

438e The Neurologic Screening Exam

Daniel H. Lowenstein

Knowledge of the basic neurologic examination is an essential clinical skill. A simple neurologic screening examination—assessment of mental status, cranial nerves, motor system, sensory system, coordination, and gait—can be reliably performed in 3–5 min. Although the components of the examination may appear daunting at first, skills usually improve rapidly with repetition and practice. In this video, the technique of performing a simple and efficient screening examination is presented.

439e Video Atlas of the Detailed Neurologic Examination

Martin A. Samuels

The comprehensive neurologic examination is an irreplaceable tool for the efficient diagnosis of neurologic disorders. Mastery of its details requires knowledge of normal nervous system anatomy and physiology combined with personal experience performing orderly and systematic examinations on large numbers of patients and healthy individuals. In the hands of a great clinician, the neurologic examination also becomes a thing of beauty—the pinnacle of the art of medicine. In this video, the most commonly used components of the examination are presented in detail, with a particular emphasis on those elements that are most helpful for assessment of common neurologic problems.

440e Neuroimaging in Neurologic Disorders

William P. Dillon

The clinician caring for patients with neurologic symptoms is faced with myriad imaging options, including computed tomography (CT), CT angiography (CTA), perfusion CT (pCT), magnetic resonance (MR) imaging (MRI), MR angiography (MRA), functional MRI (fMRI), MR spectroscopy (MRS), MR neurography (MRN), diffusion and diffusion tensor imaging, susceptibility-weighted MR imaging (SWI), arterial spin label MRI (ASL) and perfusion MRI (pMRI). In addition, an increasing number of interventional neuroradiologic techniques are available, including angiography catheter embolization, coiling, and stenting of vascular structures, and spine diagnostic and interventional techniques, such as diskography, transforaminal and translaminar epidural and nerve root injections, and blood patches. Multidetector CTA (MDCTA) and gadolinium-enhanced MRA have narrowed the indications for conventional angiography, which is now reserved for patients in whom small-vessel detail is essential for diagnosis or for whom concurrent interventional therapy is planned (Table 440e-1).

In general, MRI is more sensitive than CT for the detection of lesions affecting the central nervous system (CNS), particularly those of the spinal cord, cranial nerves, and posterior fossa structures. Diffusion MR, a sequence sensitive to the microscopic motion of water, is the most sensitive technique for detecting acute ischemic stroke of the brain or spinal cord, and it is also useful in the detection of encephalitis, abscesses, and prion diseases. CT, however, is quickly acquired and is widely available, making it a pragmatic choice for the initial evaluation of patients with acute changes in mental status, suspected acute stroke, hemorrhage, and intracranial or spinal trauma. CT is also more sensitive than MRI for visualizing fine osseous detail and is indicated in the initial imaging evaluation of conductive hearing loss as well as lesions affecting the skull base and calvarium. MR may, however, add important diagnostic information regarding bone marrow infiltrative processes that are difficult to detect on CT.

COMPUTED TOMOGRAPHY

TECHNIQUE

The CT image is a cross-sectional representation of anatomy created by a computer-generated analysis of the attenuation of x-ray beams passed through a section of the body. As the x-ray beam, collimated to the desired slice width, rotates around the patient, it passes through selected regions in the body. X-rays that are not attenuated by body structures are detected by sensitive x-ray detectors aligned 180° from the x-ray tube. A computer calculates a “back projection” image from the 360° x-ray attenuation profile. Greater x-ray attenuation (e.g., as caused by bone), results in areas of high “density” (whiter) on the scan, whereas soft tissue structures that have poor attenuation of x-rays, such as organs and air-filled cavities, are lower (black) in density. The resolution of an image depends on the radiation dose, the detector size, collimation (slice thickness), the field of view, and the matrix size of the display. A modern CT scanner is capable of obtaining sections as thin as 0.5–1 mm with 0.4-mm in-plane resolution at a speed of 0.3 s per rotation; complete studies of the brain can be completed in 1–10 s.

Multidetector CT (MDCT) is now standard in most radiology departments. Single or multiple (from 4 to 320) solid-state detectors positioned opposite to the x-ray source result in multiple slices per revolution of the beam around the patient. The table moves continuously through the rotating x-ray beam, generating a continuous “helix” of information that can be reformatted into various slice thicknesses and planes. Advantages of MDCT include shorter scan times, reduced patient and organ motion, and the ability to acquire images dynamically during the infusion of intravenous contrast, which can be used to

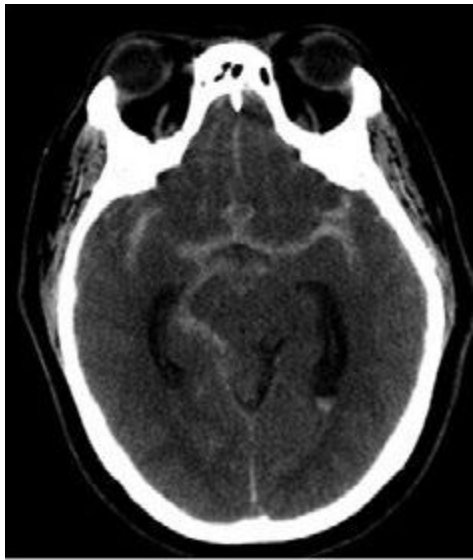
TABLE 440e-1 GUIDELINES FOR THE USE OF CT, ULTRASOUND, AND MRI

Condition	Recommended Technique
Hemorrhage	
Acute parenchymal	CT, MR
Subacute/chronic	MRI
Subarachnoid hemorrhage	CT, CTA, lumbar puncture → angiography
Aneurysm	Angiography > CTA, MRA
Ischemic infarction	
Hemorrhagic infarction	CT or MRI
Bland infarction	MRI with diffusion > CT, CTA, angiography
Carotid or vertebral dissection	MRI/MRA
Vertebral basilar insufficiency	CTA, MRI/MRA
Carotid stenosis	CTA, MRA > US
Suspected mass lesion	
Neoplasm, primary or metastatic	MRI + contrast
Infection/abscess	MRI + contrast
Immunosuppressed with focal findings	MRI + contrast
Vascular malformation	MRI ± angiography
White matter disorders	MRI
Demyelinating disease	MRI ± contrast
Dementia	MRI > CT
Trauma	
Acute trauma	CT
Shear injury/chronic hemorrhage	MRI + susceptibility-weighted imaging
Headache/migraine	CT/MRI
Seizure	
First time, no focal neurologic deficits	MRI > CT
Partial complex/refractory	MRI
Cranial neuropathy	MRI with contrast
Meningeal disease	MRI with contrast
Spine	
Low back pain	
No neurologic deficits	MRI or CT after >6 weeks
With focal deficits	MRI > CT
Spinal stenosis	MRI or CT
Cervical spondylosis	MRI, CT, CT myelography
Infection	MRI + contrast, CT
Myelopathy	MRI + contrast
Arteriovenous malformation	MRI + contrast, angiography

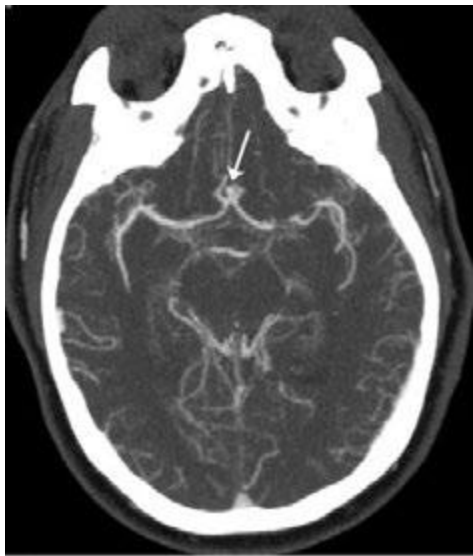
Abbreviations: CT, computed tomography; CTA, CT angiography; MRA, magnetic resonance angiography; MRI, magnetic resonance imaging.

construct CT angiograms of vascular structures and perfusion images (Figs. 440e-1B and C). CTA can be displayed in three dimensions to yield angiogram-like images (Figs. 440e-1C, 440e-2E and F, and see Fig. 446-4). CTA has proved useful in assessing the cervical and intracranial arterial and venous anatomy.

Intravenous iodinated contrast is often administered to identify both vascular structures and to detect defects in the blood-brain barrier (BBB) that are caused by tumors, infarcts, and infections. In the normal CNS, only vessels and structures lacking a BBB (e.g., the pituitary gland, choroid plexus, and dura) enhance after contrast administration. The use of iodinated contrast agents carries a small risk of allergic reaction and adds additional expense. While helpful in characterizing mass lesions as well as essential for the acquisition of CTA studies, the decision to use contrast material should always be considered carefully.



A



B



C

FIGURE 440e-1 Computed tomography (CT) angiography (CTA) of ruptured anterior cerebral artery aneurysm in a patient presenting with acute headache. **A.** Noncontrast CT demonstrates subarachnoid hemorrhage and mild obstructive hydrocephalus. **B.** Axial maximum-intensity projection from CTA demonstrates enlargement of the anterior cerebral artery (arrow). **C.** Three-dimensional surface reconstruction using a workstation confirms the anterior cerebral aneurysm and demonstrates its orientation and relationship to nearby vessels (arrow). CTA image is produced by 0.5- to 1-mm helical CT scans performed during a rapid bolus infusion of intravenous contrast medium.

INDICATIONS

CT is the primary study of choice in the evaluation of an acute change in mental status, focal neurologic findings, acute trauma to the brain and spine, suspected subarachnoid hemorrhage, and conductive hearing loss (Table 440e-1). CT is complementary to MR in the evaluation of the skull base, orbit, and osseous structures of the spine. In the spine, CT is useful in evaluating patients with osseous spinal stenosis and spondylosis, but MRI is often preferred in those with neurologic deficits. CT can also be obtained following intrathecal contrast injection to evaluate the intracranial cisterns (*CT cisternography*) for cerebrospinal fluid (CSF) fistula, as well as the spinal subarachnoid space (*CT myelography*), although intrathecal administration of gadolinium combined with MR may also be complementary.

COMPLICATIONS

CT is safe, fast, and reliable. Radiation exposure depends on the dose used but is normally between 2 and 5 mSv (millisievert) for a routine brain CT study. Care must be taken to reduce exposure when imaging children. With the advent of MDCT, CTA, and CT perfusion, the benefit must be weighed against the increased radiation doses associated with these techniques. Advanced noise reduction software now permits acceptable diagnostic CT scans at 30–40% lower radiation doses.

The most frequent complications are those associated with use of intravenous contrast agents. While two broad categories of contrast media, ionic and nonionic, are in use, ionic agents have been largely replaced by safer nonionic compounds.

Contrast nephropathy may result from hemodynamic changes, renal tubular obstruction and cell damage, or immunologic reactions to contrast agents. A rise in serum creatinine of at least 85 $\mu\text{mol/L}$ (1 mg/dL) within 48 h of contrast administration is often used as a definition of contrast nephropathy, although other causes of acute renal failure must be excluded. The prognosis is usually favorable, with serum creatinine levels returning to baseline within 1–2 weeks. Risk factors for contrast nephropathy include advanced age (>80 years), preexisting renal disease (serum creatinine exceeding 2 mg/dL), solitary kidney, diabetes mellitus, dehydration, paraproteinemia, concurrent use of nephrotoxic medication or chemotherapeutic agents, and high contrast dose. Patients with diabetes and those with mild renal failure should be well hydrated prior to the administration of contrast agents, although careful consideration should be given to alternative imaging techniques such as MRI, noncontrast CT, or ultrasound (US). Nonionic, low-osmolar media produce fewer abnormalities in renal blood flow and less endothelial cell damage but should still be used carefully in patients at risk for allergic reaction. Estimated glomerular filtration rate (eGFR) is a more reliable indicator of renal function compared to creatinine alone because it takes into account age, race, and sex. In one study, 15% of outpatients with a normal serum creatinine had an estimated creatinine clearance of 50 mL/min/1.73 m² or less (normal is ≥ 90 mL/min/1.73 m²). The exact eGFR threshold, below which withholding intravenous contrast should be considered, is controversial. The risk of contrast nephropathy increases in patients with an eGFR <60 mL/min/1.73 m²; however, the majority of these patients will only have a temporary rise in creatinine. The risk of dialysis after receiving contrast significantly increases in patients with eGFR <30 mL/min/1.73 m². Thus, an eGFR threshold between 60 and 30 mL/min/1.73 m² is appropriate; however, the exact number is somewhat arbitrary. A creatinine of 1.6 in a 70-year-old, non-African-American male corresponds to an eGFR of approximately 45 mL/min/1.73 m². The American College of Radiology suggests using an eGFR of 45 mL/min/1.73 m² as a threshold below which iodinated contrast should not be given without serious consideration of the potential for contrast nephropathy. If contrast must be administered to a patient with an eGFR below 45 mL/min/1.73 m², the patient should be well hydrated, and a reduction in the dose of contrast should be considered. Use of other agents such as bicarbonate and acetylcysteine may reduce the incidence of contrast nephropathy.

Allergy Immediate reactions following intravenous contrast media can occur through several mechanisms. The most severe reactions

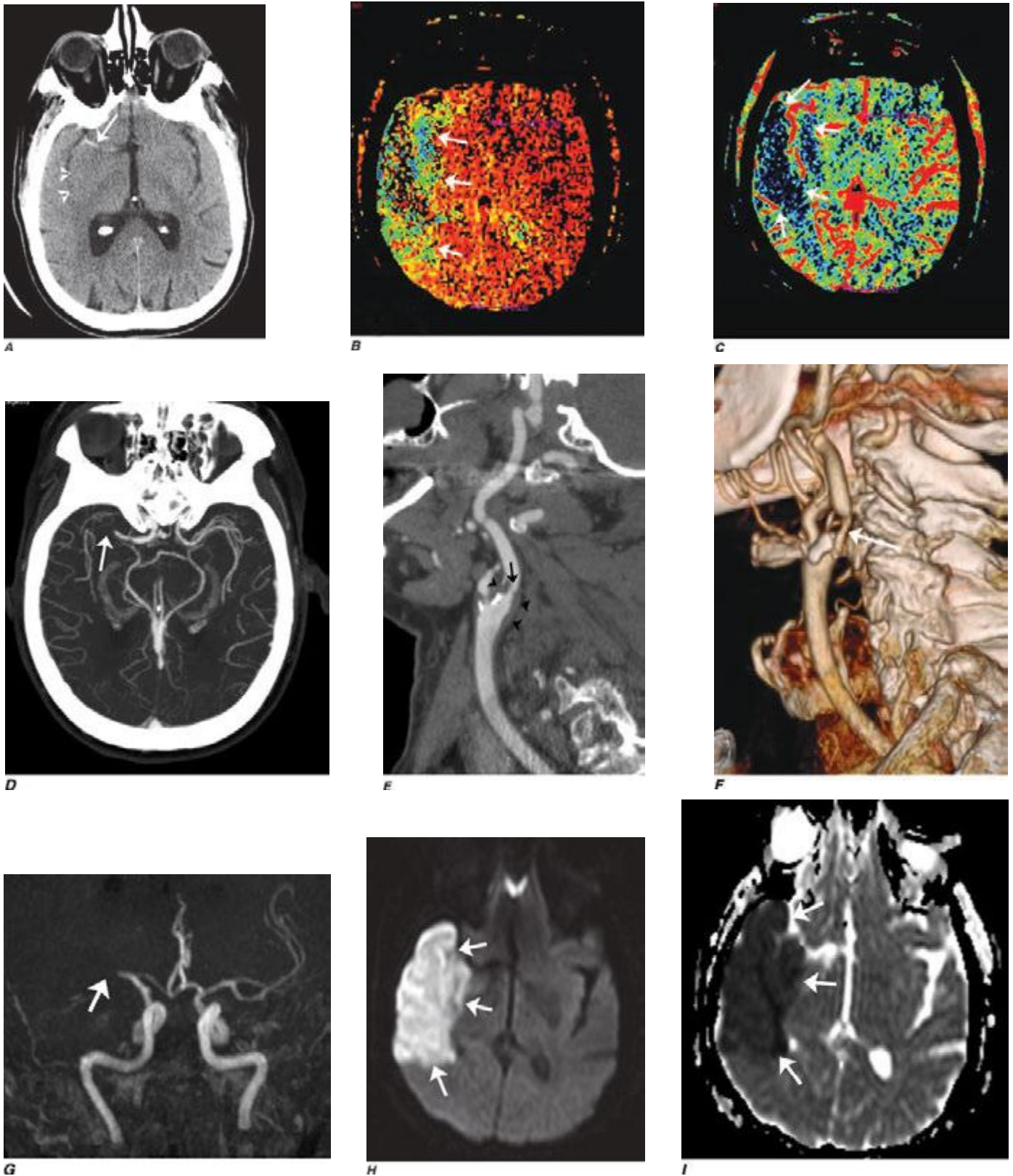


FIGURE 440e-2 Acute left hemiparesis due to middle cerebral artery occlusion. **A.** Axial noncontrast computed tomography (CT) scan demonstrates high density within the right middle cerebral artery (*arrow*) associated with subtle low density involving the right putamen (*arrowheads*). **B.** Mean transit time CT perfusion parametric map indicating prolonged mean transit time involving the right middle cerebral territory (*arrows*). **C.** Cerebral blood volume (CBV) map shows reduced CBV involving an area within the defect shown in **B**, indicating a high likelihood of infarction (*arrows*). **D.** Axial maximum-intensity projection from a CT angiography (CTA) study through the circle of Willis demonstrates an abrupt occlusion of the proximal right middle cerebral artery (*arrow*). **E.** Sagittal reformation through the right internal carotid artery demonstrates a low-density lipid-laden plaque (*arrowheads*) narrowing the lumen (*black arrow*). **F.** Three-dimensional surface-rendered CTA image demonstrates calcification and narrowing of the right internal carotid artery (*arrow*), consistent with atherosclerotic disease. **G.** Coronal maximum-intensity projection from magnetic resonance angiography shows right middle cerebral artery (MCA) occlusion (*arrow*). **H.** and **I.** Axial diffusion-weighted image (**H**) and apparent diffusion coefficient image (**I**) documents the presence of a right middle cerebral artery infarction.

TABLE 440e-2 GUIDELINES FOR PREMEDICATION OF PATIENTS WITH PRIOR CONTRAST ALLERGY**12 h prior to examination:**

Prednisone, 50 mg PO or methylprednisolone, 32 mg PO

2 h prior to examination:

Prednisone, 50 mg PO or methylprednisolone, 32 mg PO and cimetidine, 300 mg PO or ranitidine, 150 mg PO

Immediately prior to examination:

Benadryl, 50 mg IV (alternatively, can be given PO 2 h prior to exam)

are related to allergic hypersensitivity (anaphylaxis) and range from mild hives to bronchospasm and death. The pathogenesis of allergic hypersensitivity reactions is thought to include the release of mediators such as histamine, antibody-antigen reactions, and complement activation. Severe allergic reactions occur in ~0.04% of patients receiving nonionic media, sixfold lower than with ionic media. Risk factors include a history of prior contrast reaction (fivefold increased likelihood), food and/or drug allergies, and atopy (asthma and hay fever). The predictive value of specific allergies, such as those to shellfish, once thought important, actually is now recognized to be unreliable. Nonetheless, in patients with a history worrisome for potential allergic reaction, a noncontrast CT or MRI procedure should be considered as an alternative to contrast administration. If iodinated contrast is absolutely required, a nonionic agent should be used in conjunction with pretreatment with glucocorticoids and antihistamines (**Table 440e-2**); however, pretreatment does not guarantee safety. Patients with allergic reactions to iodinated contrast material do not usually react to gadolinium-based MR contrast material, although such reactions can occur. It would be wise to pretreat patients with a prior allergic history to MR contrast administration in a similar fashion. Nonimmediate (>1 h after injection) reactions are frequent and probably related to T cell-mediated immune reactions. These are typically urticarial but can occasionally be more severe. Drug provocation and skin testing may be required to determine the culprit agent involved as well as determine a safe alternative.

Other side effects of CT scanning are rare but include a sensation of warmth throughout the body and a metallic taste during intravenous administration of iodinated contrast media. Extravasation of contrast media, although rare, can be painful and lead to compartment syndrome. When this occurs, consultation with plastic surgery is indicated. Patients with significant cardiac disease may be at increased risk for contrast reactions, and in these patients, limits to the volume and osmolality of the contrast media should be considered. Patients who may undergo systemic radioactive iodine therapy for thyroid disease or cancer should not receive iodinated contrast media if possible, because this will decrease the uptake of the radioisotope into the tumor or thyroid (see the *American College of Radiology Manual on Contrast Media*, Version 9, 2013; http://www.acr.org/~media/ACR/Documents/PDF/QualitySafety/Resources/Contrast%20Manual/2013_Contrast_Media.pdf).

MAGNETIC RESONANCE IMAGING**TECHNIQUE**

MRI is a complex interaction between hydrogen protons in biologic tissues, a static magnetic field (the magnet), and energy in the form of radiofrequency (Rf) waves of a specific frequency introduced by coils placed next to the body part of interest. Images are made by computerized processing of resonance information received from protons in the body. Field strength of the magnet is directly related to signal-to-noise ratio. While 1.5-T magnets have become the standard high-field MRI units, 3-T magnets are now widely available and have distinct advantages in the brain and musculoskeletal systems. Even higher field magnets (7-T) and positron emission tomography (PET) MR machines promise increased resolution and anatomic-functional information on a variety of disorders. Spatial localization is achieved by magnetic gradients surrounding the main magnet, which impart slight changes

in magnetic field throughout the imaging volume. Rf pulses transiently excite the energy state of the hydrogen protons in the body. Rf is administered at a frequency specific for the field strength of the magnet. The subsequent return to equilibrium energy state (*relaxation*) of the hydrogen protons results in a release of Rf energy (the *echo*), which is detected by the coils that delivered the Rf pulses. Fourier analysis is used to transform the echo into the information used to form an MR image. The MR image thus consists of a map of the distribution of hydrogen protons, with signal intensity imparted by both density of hydrogen protons and differences in the relaxation times (see below) of hydrogen protons on different molecules. Although clinical MRI currently makes use of the ubiquitous hydrogen proton, research into sodium and carbon imaging and spectroscopy appears promising.

T1 and T2 Relaxation Times The rate of return to equilibrium of perturbed protons is called the *relaxation rate*. The relaxation rate varies among normal and pathologic tissues. The relaxation rate of a hydrogen proton in a tissue is influenced by local interactions with surrounding molecules and atomic neighbors. Two relaxation rates, T1 and T2, influence the signal intensity of the image. The T1 relaxation time is the time, measured in milliseconds, for 63% of the hydrogen protons to return to their normal equilibrium state, whereas the T2 relaxation is the time for 63% of the protons to become dephased owing to interactions among nearby protons. The intensity and image contrast of the signal within various tissues can be modulated by altering acquisition parameters such as the interval between Rf pulses (TR) and the time between the Rf pulse and the signal reception (TE). T1-weighted (T1W) images are produced by keeping the TR and TE relatively short, whereas using longer TR and TE times produces T2-weighted (T2W) images. Fat and subacute hemorrhage have relatively shorter T1 relaxation rates and thus higher signal intensity than brain on T1W images. Structures containing more water, such as CSF and edema, have long T1 and T2 relaxation rates, resulting in relatively lower signal intensity on T1W images and higher signal intensity on T2W images (**Table 440e-3**). Gray matter contains 10–15% more water than white matter, which accounts for much of the intrinsic contrast between the two on MRI (Fig. 440e-6B). T2W images are more sensitive than T1W images to edema, demyelination, infarction, and chronic hemorrhage, whereas T1W imaging is more sensitive to subacute hemorrhage and fat-containing structures.

Many different MR pulse sequences exist, and each can be obtained in various planes (**Figs. 440e-2, 440e-3, and 440e-4**). The selection of a proper protocol that will best answer a clinical question depends on an accurate clinical history and indication for the examination. Fluid-attenuated inversion recovery (FLAIR) is a useful pulse sequence that produces T2W images in which the normally high signal intensity of CSF is suppressed (Fig. 440e-6B). FLAIR images are more sensitive than standard spin echo images for any water-containing lesions or edema. Susceptibility-weighted imaging, such as gradient echo imaging, is very sensitive to magnetic susceptibility generated by blood, calcium, and air and routinely obtained in patients suspected of pathology that might result in microhemorrhages, such as amyloid, hemorrhagic metastases, and thrombotic states (**Fig. 440e-5C**). MR images can be generated in any plane without changing the patient's position. Each sequence, however, must be obtained separately and takes 1–10 min on average to complete. Three-dimensional volumetric imaging is also possible with MRI, resulting in a three-dimensional

TABLE 440e-3 SOME COMMON INTENSITIES ON T1- AND T2-WEIGHTED MRI SEQUENCES

Image	TR	TE	Signal Intensity			
			CSF	Fat	Brain	Edema
T1W	Short	Short	Low	High	Low	Low
T2W	Long	Long	High	Low	High	High
FLAIR (T2)	Long	Long	Low	Medium	High	High

Abbreviations: CSF, cerebrospinal fluid; FLAIR, fluid-attenuated inversion recovery; TE, interval between radiofrequency pulse and signal reception; TR, interval between radiofrequency pulses; T1W and T2W, T1- and T2-weighted.

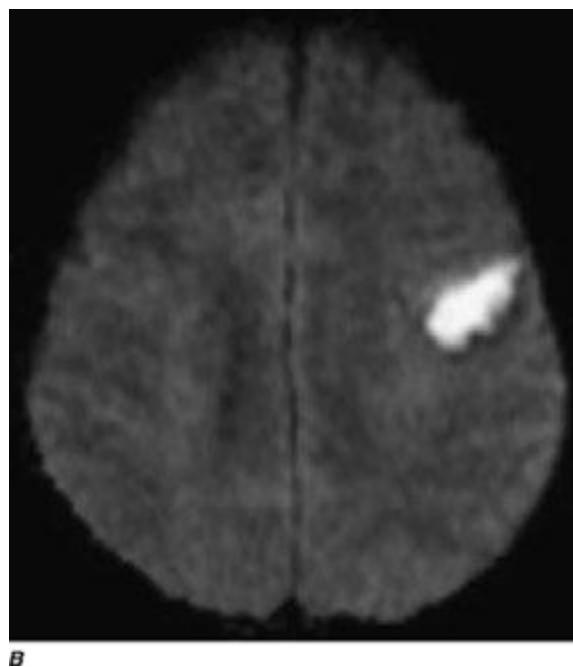
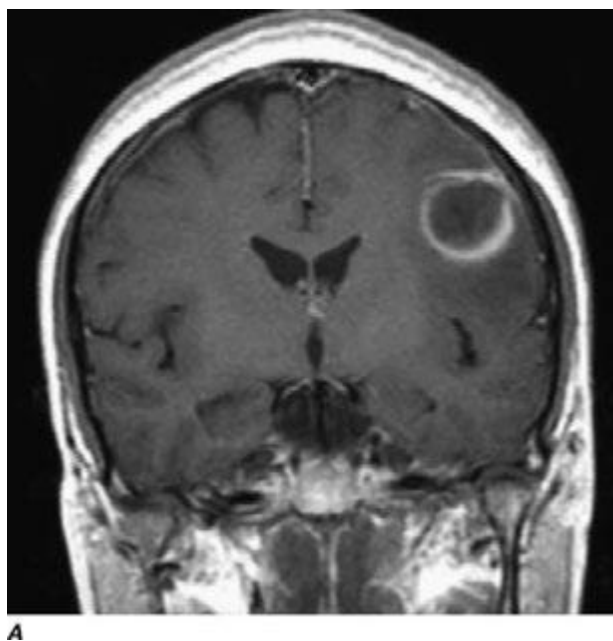


FIGURE 440e-3 Cerebral abscess in a patient with fever and a right hemiparesis. **A.** Coronal postcontrast T1-weighted image demonstrates a ring-enhancing mass in the left frontal lobe. **B.** Axial diffusion-weighted image demonstrates restricted diffusion (high signal intensity) within the lesion, which in this setting is highly suggestive of cerebral abscess.

volume of data that can be reformatted in any orientation to highlight certain disease processes.

MR Contrast Material The heavy-metal element gadolinium forms the basis of all currently approved intravenous MR contrast agents. Gadolinium is a paramagnetic substance, which means that it reduces the T1 and T2 relaxation times of nearby water protons, resulting in a high signal on T1W images and a low signal on T2W images (the latter requires a sufficient local concentration, usually in the form of an intravenous bolus). Unlike iodinated contrast agents, the effect of MR contrast agents depends on the presence of local hydrogen protons on which it must act to achieve the desired effect. There are nine different gadolinium agents approved in the United States for use with MRI. These differ according to the attached chelated moiety, which also affects the strength of chelation of the otherwise toxic gadolinium element. The chelating carrier molecule for gadolinium can be classified by whether it is macrocyclic or has linear geometry and whether it is ionic or nonionic. Most of these are excreted by the renal system. Cyclical agents are less likely to release the gadolinium element, and thus are considered the safest category.

ALLERGIC HYPERSENSITIVITY Gadolinium-DTPA (diethylenetriamine-pentaacetic acid) does not normally cross the intact BBB (Fig. 440e-3A) as well as areas of the brain that normally are devoid of the BBB (pituitary, dura, choroid plexus). However, gadolinium contrast has been noted to slowly cross an intact BBB over time and especially in the setting of reduced renal clearance or inflamed meninges. The agents are generally well tolerated; overall adverse events after injection range from 0.07–2.4%. True allergic reactions are rare (0.004–0.7%) but have been reported. Severe life-threatening reactions are exceedingly rare; in one report, only 55 reactions out of 20 million doses occurred. However, the adverse reaction rate in patients with a prior history of reaction to gadolinium is eight times higher than normal. Other risk factors include atopy or asthma (3.7%); although there is no cross-reactivity to iodinated contrast material, those with a prior allergic response to iodine should be considered at higher risk. Gadolinium contrast material can be administered safely to children as well as adults, although these agents are generally avoided in those under 6 months of age.

NEPHROTOXICITY Contrast-induced renal failure does not occur with gadolinium agents. A rare complication, nephrogenic systemic fibrosis

(NSF), has occurred in patients with severe renal insufficiency who have been exposed to gadolinium contrast agents. The onset of NSF has been reported between 5 and 75 days following exposure; histologic features include thickened collagen bundles with surrounding clefts, mucin deposition, and increased numbers of fibrocytes and elastic fibers in skin. In addition to dermatologic symptoms, other manifestations include widespread fibrosis of the skeletal muscle, bone, lungs, pleura, pericardium, myocardium, kidney, muscle, bone, testes, and dura. The American College of Radiology recommends that a glomerular filtration rate (GFR) assessment be obtained within 6 weeks prior to elective gadolinium-based MR contrast agent administration in patients with:

1. A history of renal disease (including solitary kidney, renal transplant, renal tumor)
2. Age >60 years
3. History of hypertension
4. History of diabetes
5. History of severe hepatic disease, liver transplant, or pending liver transplant; for these patients, it is recommended that the patient's GFR assessment be nearly contemporaneous with the MR examination

The incidence of NSF in patients with severe renal dysfunction (GFR <30) varies from 0.19 to 4%. Other risk factors for NSF include acute kidney injury, the use of nonmacrocyclic agents, and repeated or high-dose exposure to gadolinium. The American College of Radiology Committee on Drugs and Contrast Media states that patients receiving any gadolinium-containing agent should be considered at risk of NSF if they are on dialysis (of any form); have severe or end-stage chronic renal disease (eGFR <30 mL/min/1.73 m²) without dialysis; eGFR of 30–40 mL/min/1.73 m² without dialysis (as the GFR may fluctuate); or have acute renal insufficiency.

COMPLICATIONS AND CONTRAINDICATIONS

From the patient's perspective, an MRI examination can be intimidating, and a higher level of cooperation is required than with CT. The patient lies on a table that is moved into a long, narrow gap within the magnet. Approximately 5% of the population experiences severe claustrophobia in the MR environment. This can be reduced by mild sedation but remains a problem for some. Because it takes between 3 and

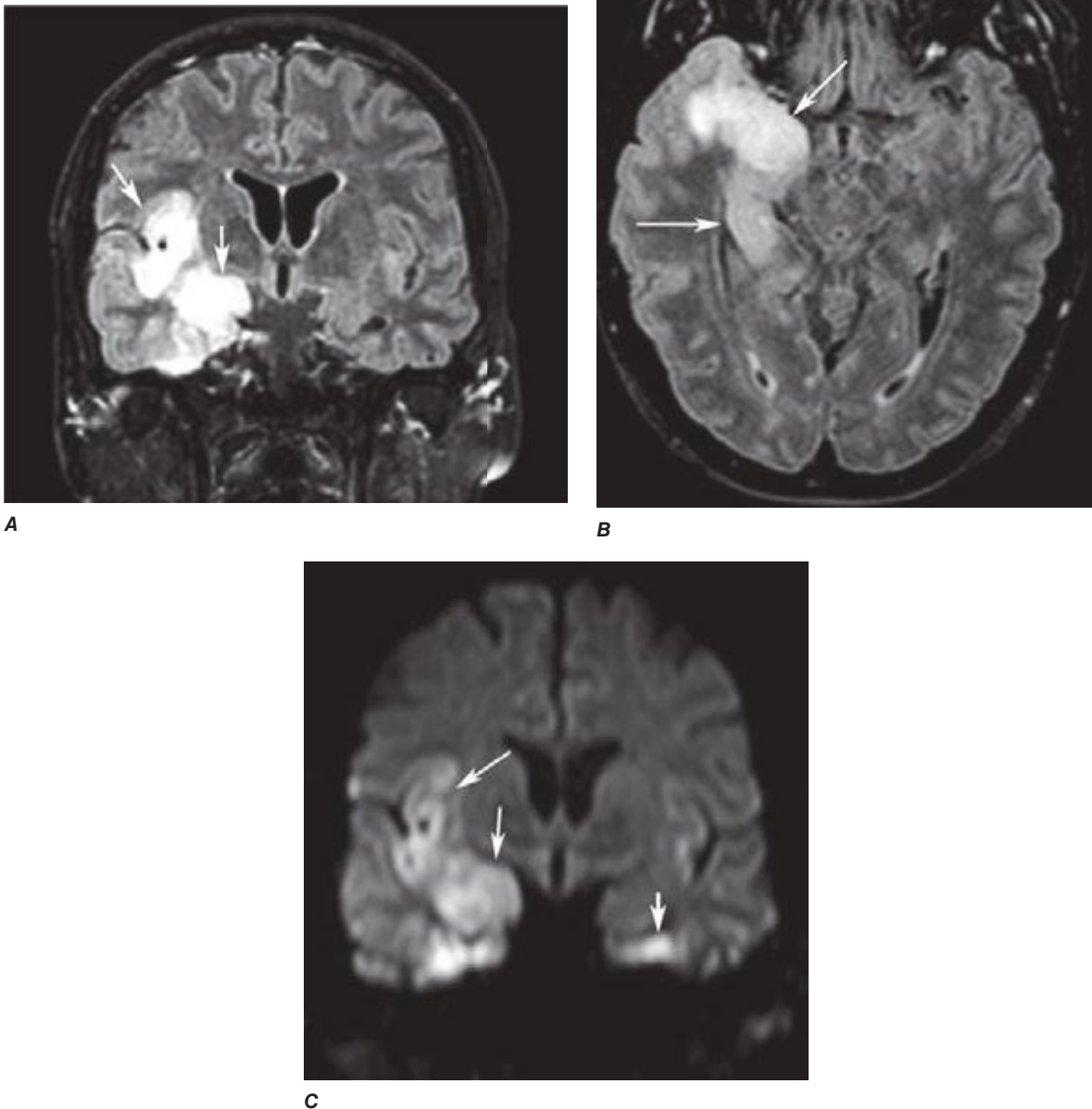


FIGURE 440e-4 Herpes simplex encephalitis in a patient presenting with altered mental status and fever. **A.** and **B.** Coronal (**A**) and axial (**B**) T2-weighted fluid-attenuated inversion recovery images demonstrate expansion and high signal intensity involving the right medial temporal lobe and insular cortex (*arrows*). **C.** Coronal diffusion-weighted image demonstrates high signal intensity indicating restricted diffusion involving the right medial temporal lobe and hippocampus (*arrows*) as well as subtle involvement of the left inferior temporal lobe (*arrowhead*). This is most consistent with neuronal death and can be seen in acute infarction as well as encephalitis and other inflammatory conditions. The suspected diagnosis of herpes simplex encephalitis was confirmed by cerebrospinal fluid polymerase chain reaction analysis.

10 min per sequence, movement of the patient during an MR exam distorts all of the images; therefore, uncooperative patients should either be sedated for the MR study or scanned with CT. Generally, children under the age of 8 years usually require conscious sedation in order to complete the MR examination without motion degradation.

MRI is considered safe for patients, even at very high field strengths. Serious injuries have been caused, however, by attraction of ferromagnetic objects into the magnet, which act as missiles if brought too close to the magnet. Likewise, ferromagnetic implants, such as aneurysm clips, may torque within the magnet, causing damage to vessels and even death. Metallic foreign bodies in the eye have moved and caused intraocular hemorrhage; screening for ocular metallic fragments is indicated in those with a history of metal work or ocular metallic foreign bodies. Implanted cardiac pacemakers are generally a contraindication

to MRI owing to the risk of induced arrhythmias; however, some newer pacemakers have been shown to be safe. All health care personnel and patients must be screened and educated thoroughly to prevent such disasters because the magnet is always “on.” **Table 440e-4** lists common contraindications for MRI.

MAGNETIC RESONANCE ANGIOGRAPHY

MR angiography is a general term describing several MR techniques that result in vascular-weighted images. These provide a vascular flow map rather than the anatomic map shown by conventional angiography. On routine spin echo MR sequences, moving protons (e.g., flowing blood, CSF) exhibit complex MR signals that range from high- to low-signal intensity relative to background stationary tissue.

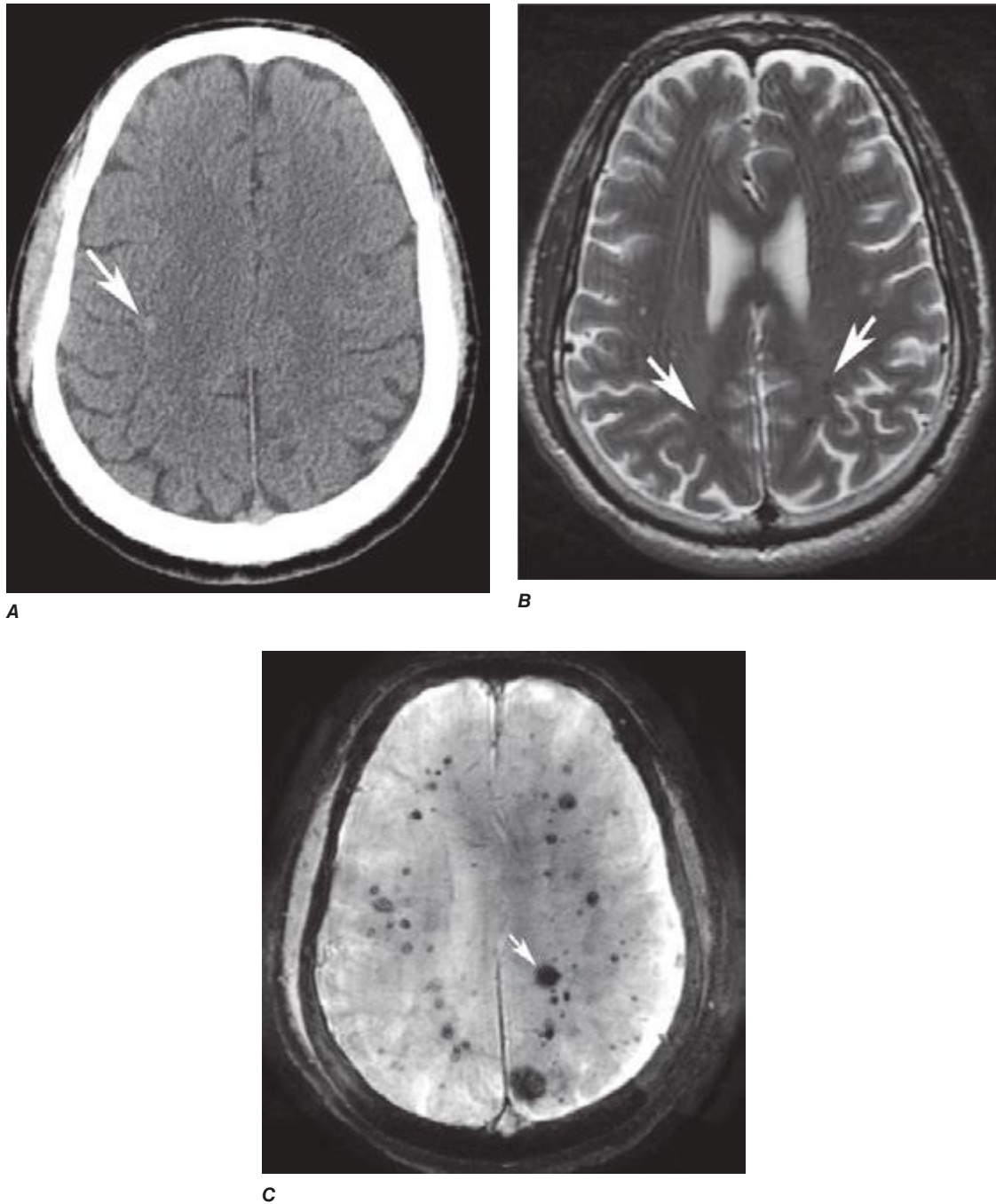


FIGURE 440e-5 Susceptibility-weighted imaging in a patient with familial cavernous malformations. **A.** Noncontrast computed tomography scan shows one hyperdense lesion in the right hemisphere (*arrow*). **B.** T2-weighted fast spin echo image shows subtle low-intensity lesions (*arrows*). **C.** Susceptibility-weighted image shows numerous low-intensity lesions consistent with hemosiderin-laden cavernous malformations (*arrow*).

Fast-flowing blood returns no signal (flow void) on routine T1W or T2W spin echo MR images. Slower-flowing blood, as occurs in veins or distal to arterial stenosis, may appear high in signal. However, using special pulse sequences called *gradient echo sequences*, it is possible to increase the signal intensity of moving protons in contrast to the low signal background intensity of stationary tissue. This creates angiography-like images, which can be manipulated in three dimensions to highlight vascular anatomy and relationships.

So called time-of-flight (TOF) MRA relies on the suppression of nonmoving tissue to provide a low-intensity background for the high signal intensity of flowing blood entering the section; arterial or venous structures may be highlighted. A typical TOF MRA sequence results in a series of contiguous, thin MR sections (0.6–0.9 mm thick), which can be viewed as a stack and manipulated to create an

angiographic image data set that can be reformatted and viewed in various planes and angles, much like that seen with conventional angiography (**Fig. 440e-2G**).

Phase-contrast MRA has a longer acquisition time than TOF MRA, but in addition to providing anatomic information similar to that of TOF imaging, it can be used to reveal the velocity and direction of blood flow in a given vessel. Through the selection of different imaging parameters, differing blood velocities can be highlighted; selective venous and arterial MRA images can thus be obtained. One advantage of phase-contrast MRA is the excellent suppression of high-signal-intensity background structures.

MRA can also be acquired during infusion of contrast material. Advantages include faster imaging times (1–2 min vs 10 min), fewer flow-related artifacts, and higher resolution images. Recently,

TABLE 440e-4 COMMON CONTRAINDICATIONS TO MAGNETIC RESONANCE IMAGING

Cardiac pacemaker or permanent pacemaker leads
Internal defibrillatory device
Cochlear prostheses
Bone growth stimulators
Spinal cord stimulators
Electronic infusion devices
Intracranial aneurysm clips (some but not all)
Ocular implants (some) or ocular metallic foreign body
McGee stapedectomy piston prosthesis
Duraphase penile implant
Swan-Ganz catheter
Magnetic stoma plugs
Magnetic dental implants
Magnetic sphincters
Ferromagnetic inferior vena cava filters, coils, stents—safe 6 weeks after implantation
Tattooed eyeliner (contains ferromagnetic material and may irritate eyes)

Note: See also <http://www.mrisafety.com>.

contrast-enhanced MRA has become the standard for extracranial vascular MRA. This technique entails rapid imaging using coronal three-dimensional TOF sequences during a bolus infusion of gadolinium contrast agent. Proper technique and timing of acquisition relative to bolus arrival are critical for success.

MRA has lower spatial resolution compared with conventional film-based angiography, and therefore the detection of small-vessel abnormalities, such as vasculitis and distal vasospasm, is problematic. MRA is also less sensitive to slowly flowing blood and thus may not reliably differentiate complete from near-complete occlusions. Motion, either by the patient or by anatomic structures, may distort the MRA images, creating artifacts. These limitations notwithstanding, MRA has proved useful in evaluation of the extracranial carotid and vertebral circulation as well as of larger-caliber intracranial arteries and dural sinuses. It has also proved useful in the noninvasive detection of intracranial aneurysms and vascular malformations.

ECHO-PLANAR MRI

Recent improvements in gradients, software, and high-speed computer processors now permit extremely rapid MRI of the brain. With echo-planar MRI (EPI), fast gradients are switched on and off at high speeds to create the information used to form an image. In routine spin echo imaging, images of the brain can be obtained in 5–10 min. With EPI, all of the information required for processing an image is accumulated in milliseconds, and the information for the entire brain can be obtained in less than 1–2 min, depending on the degree of resolution required or desired. Fast MRI reduces patient and organ motion and is the basis of perfusion imaging during contrast infusion and kinematic motion studies. EPI is also the sequence used to obtain diffusion imaging and tractography, as well as fMRI and arterial spin-labeled studies (Figs. 440e-2H, 440e-3, 440e-4C, and 440e-6; and see Fig. 446-16).

Perfusion and diffusion imaging are EPI techniques that are useful in early detection of ischemic injury of the brain and may be useful together to demonstrate infarcted tissue as well as ischemic but potentially viable tissue at risk of infarction (e.g., the ischemic penumbra). Diffusion-weighted imaging (DWI) assesses microscopic motion of water; abnormal restriction of motion appears as relative high-signal intensity on diffusion-weighted images. Infarcted tissue reduces the water motion within cells and in the interstitial tissues, resulting in high signal on DWI. DWI is the most sensitive technique for detection of acute cerebral infarction of <7 days in duration (Fig. 440e-2H). It is also quite sensitive for detecting dying or dead brain tissue secondary to encephalitis, as well as abscess formation (Fig. 440e-3B).

Perfusion MRI involves the acquisition of fast echo planar gradient images during a rapid intravenous bolus of gadolinium contrast material. Relative cerebral blood volume, mean transit time, and cerebral blood flow maps are then derived. Delay in mean transit time and reduction in cerebral blood volume and cerebral blood flow are typical of infarction. In the setting of reduced blood flow, a prolonged mean transit time of contrast but normal or elevated cerebral blood volume may indicate tissue supplied by collateral flow that is at risk of infarction. Perfusion MRI imaging can also be used in the assessment of brain tumors to differentiate intraaxial primary tumors, whose BBB is relatively intact, from extraaxial tumors or metastases, which demonstrate a relatively more permeable BBB.

Diffusion tensor imaging is derived from diffusion MRI imaging sequences, which assesses the direction of microscopic motion of water along white matter tracts. This technique has great potential in the assessment of brain maturation as well as disease entities that undermine the integrity of the white matter architecture. It has proven valuable in preoperative assessment of subcortical white matter tract anatomy prior to brain tumor surgery (Fig. 440e-6).

fMRI of the brain is an EPI technique that localizes regions of activity in the brain following task activation. Neuronal activity elicits a slight increase in the delivery of oxygenated blood flow to a specific region of activated brain. This results in an alteration in the balance of oxyhemoglobin and deoxyhemoglobin, which yields a 2–3% increase in signal intensity within veins and local capillaries. Further studies will determine whether these techniques are cost effective or clinically useful, but currently, preoperative somatosensory and auditory cortex localization is possible. This technique has proved useful to neuroscientists interested in interrogating the localization of certain brain functions.

ARTERIAL SPIN LABELING

ASL is a quantitative noninvasive MR technique that measures cerebral blood flow. Blood traversing in the neck is labeled by an MR pulse and then imaged in the brain after a short delay. The signal in the brain is reflective of blood flow. ASL is an especially important technique for patients with kidney failure and for pediatric patients in whom the use of radioactive tracers or exogenous contrast agents is contraindicated. Increased cerebral flow is more easily identified than slow flow, which can be sometimes difficult to quantify. This technique has also been shown useful in detecting arterial venous shunting in arteriovenous malformations and arteriovenous fistulas.

MAGNETIC RESONANCE NEUROGRAPHY

MRN is a T2W MR technique that shows promise in detecting increased signal in irritated, inflamed, or infiltrated peripheral nerves. Images are obtained with fat-suppressed fast spin echo imaging or short inversion recovery sequences. Irritated or infiltrated nerves will demonstrate high signal on T2W imaging. This is indicated in patients with radiculopathy whose conventional MR studies of the spine are normal, or in those suspected of peripheral nerve entrapment or trauma.

POSITRON EMISSION TOMOGRAPHY

PET relies on the detection of positrons emitted during the decay of a radionuclide that has been injected into a patient. The most frequently used moiety is 2-[¹⁸F]fluoro-2-deoxy-D-glucose (FDG), which is an analogue of glucose and is taken up by cells competitively with 2-deoxyglucose. Multiple images of glucose uptake activity are formed after 45–60 min. Images reveal differences in regional glucose activity among normal and pathologic brain structures. FDG-PET is used primarily for the detection of extracranial metastatic disease; however, a lower activity of FDG in the parietal lobes is associated with Alzheimer's disease, a finding that may simply reflect atrophy that occurs in the later stages of the disease. Combination PET-CT scanners, in which both CT and PET are obtained at one sitting, have largely replaced PET scans alone for most clinical indications. MR-PET scanners have also

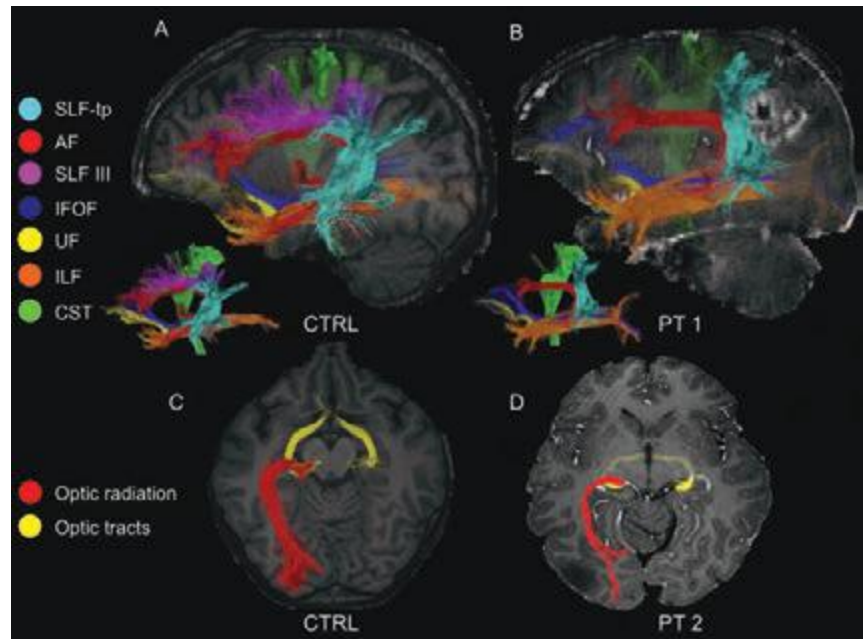
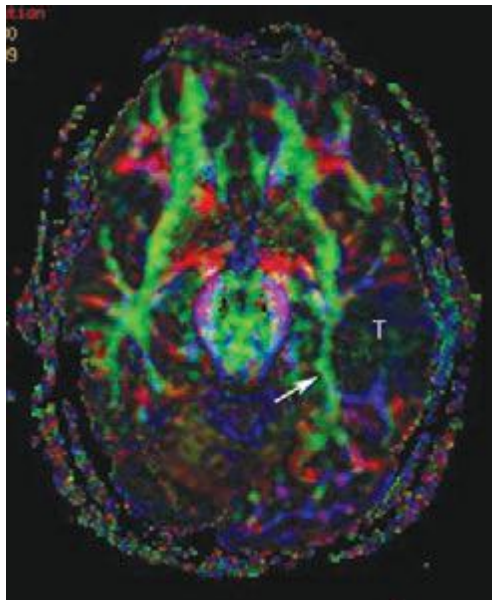
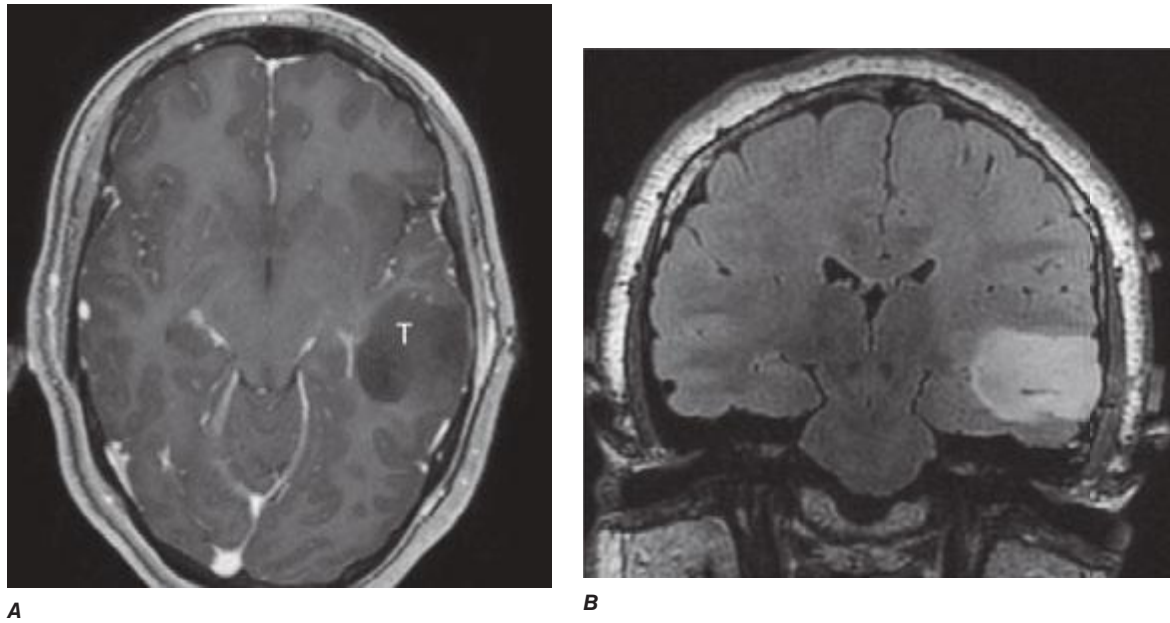


FIGURE 440e-6 Diffusion tractography in cerebral glioma. Associative and descending pathways in a healthy subject (**A**) and in a patient with parietal lobe glioblastoma (**B**) presenting with a language deficit: the mass causes a disruption of the arcuate-SLF complex, in particular of its anterior portion (SLF III). Also shown are bilateral optic tract and left optic radiation pathways in a healthy subject (**C**) and in a patient with left occipital grade II oligoastrocytoma (**D**): the mass causes a disruption of the left optic radiation. Shown in neurologic orientation, i.e., the left brain appears on the left side of the image. AF, long segment of the arcuate fascicle; CST, corticospinal tract; IFOF, inferior fronto-occipital tract; ILF, inferior longitudinal fascicle; SLF III, superior longitudinal fascicle III or anterior segment of the arcuate fascicle; SLF-tp, temporo-parietal portion of the superior longitudinal fascicle or posterior segment of the arcuate fascicle; T, tumor; UF, uncinated fascicle. (Part D courtesy of Eduardo Caverzasi and Roland Henry.)

been developed and may prove useful for imaging the brain and other organs without the radiation exposure of CT. More recent PET ligand developments include amyloid tracers, such as Pittsburgh compound B (PIB) and 18-F AV-45 (florbetapir), and tau PET tracers, such as 18F-T807 and T808. Studies have shown an increased percentage of amyloid deposition in patients with Alzheimer's disease compared with mild cognitive impairment and healthy controls; however, up to 25% of cognitively "normal" patients show abnormalities on amyloid PET imaging. This may either reflect subclinical disease processes or variation of normal. Tau imaging may be more specific for Alzheimer's disease, and clinical studies are under way.

MYELOGRAPHY

TECHNIQUE

Myelography involves the intrathecal instillation of specially formulated water-soluble iodinated contrast medium into the lumbar or cervical subarachnoid space. CT scanning is typically performed after myelography (*CT myelography*) to better demonstrate the spinal cord and roots, which appear as filling defects in the opacified subarachnoid space. *Low-dose CT myelography*, in which CT is performed after the subarachnoid injection of a small amount of relatively dilute contrast material, has replaced conventional myelography for many indications, thereby

440e-10 reducing exposure to radiation and contrast media. Newer multidetector scanners now obtain CT studies quickly so that reformations in sagittal and coronal planes, equivalent to traditional myelography projections, are now routine.

INDICATIONS

Myelography has been largely replaced by CT myelography and MRI for diagnosis of diseases of the spinal canal and cord (Table 440e-1). Remaining indications for conventional plain-film myelography include the evaluation of suspected meningeal or arachnoid cysts and the localization of CSF fistulas. Conventional myelography and CT myelography provide the most precise information in patients with prior spinal fusion and spinal fixation hardware.

CONTRAINDICATIONS

Myelography is relatively safe; however, it should be performed with caution in any patient with elevated intracranial pressure, evidence of a spinal block, or a history of allergic reaction to intrathecal contrast media. In patients with a suspected spinal block, MR is the preferred technique. If myelography is necessary, only a small amount of contrast medium should be instilled below the lesion in order to minimize the risk of neurologic deterioration. Lumbar puncture is to be avoided in patients with bleeding disorders, including patients receiving anticoagulant therapy, as well as in those with infections of the overlying soft tissues (Chap. 443e).

COMPLICATIONS

Headache is the most frequent complication of myelography and is reported to occur in 5–30% of patients. Nausea and vomiting may also occur rarely. Postural headache (post-lumbar puncture headache) is generally due to leakage of CSF from the puncture site, resulting in CSF hypotension. A higher incidence is noted among younger women and with the use of larger gauge cutting-type spinal needles. If significant headache persists for longer than 48 h, placement of an epidural blood patch should be considered. **Management of lumbar puncture headache is discussed in Chap. 21.** Vasovagal syncope may occur during lumbar puncture; it is accentuated by the upright position used during lumbar myelography. Adequate hydration before and after myelography will reduce the incidence of this complication.

Hearing loss is a rare complication of myelography. It may result from a direct toxic effect of the contrast medium or from an alteration of the pressure equilibrium between CSF and perilymph in the inner ear. Puncture of the spinal cord is a rare but serious complication of cervical (C1–2) or high lumbar puncture. The risk of cord puncture is greatest in patients with spinal stenosis, Chiari malformations, or conditions that reduce CSF volume. In these settings, a low-dose lumbar injection followed by thin-section CT or MRI is a safer alternative to cervical puncture. Intrathecal contrast reactions are rare, but aseptic meningitis and encephalopathy are reported complications. The latter is usually dose related and associated with contrast entering the intracranial subarachnoid space. Seizures occur following myelography in 0.1–0.3% of patients. Risk factors include a preexisting seizure disorder and the use of a total iodine dose of >4500 mg. Other reported complications include hyperthermia, hallucinations, depression, and anxiety states. These side effects have been reduced by the development of nonionic, water-soluble contrast agents as well as by head elevation and generous hydration following myelography.

SPINE INTERVENTIONS

DISKOGRAPHY

The evaluation of back pain and radiculopathy may require diagnostic procedures that attempt either to reproduce the patient's pain or relieve it, indicating its correct source prior to lumbar fusion. Diskography is performed by fluoroscopic placement of a 22- to 25-gauge needle into the intervertebral disk and subsequent injection of 1–3 mL of contrast media. The intradiskal pressure is recorded, as is an assessment of the patient's response to the injection of contrast material. Typically little or no pain is felt during injection of a normal disk, which does not

accept much more than 1 mL of contrast material, even at pressures as high as 415–690 kPa (60–100 lb/in²). CT and plain films are obtained following the procedure. Concerns have been raised that diskography may contribute to an accelerated rate of disk degeneration.

SELECTIVE NERVE ROOT AND EPIDURAL SPINAL INJECTIONS

Percutaneous selective nerve root and epidural blocks with glucocorticoid and anesthetic mixtures may be both therapeutic and diagnostic, especially if a patient's pain is relieved. Typically, 1–2 mL of an equal mixture of a long-acting glucocorticoid such as betamethasone and a long-acting anesthetic such as bupivacaine 0.75% is instilled under CT or fluoroscopic guidance in the intraspinal epidural space or adjacent to an existing nerve root.

ANGIOGRAPHY

Catheter angiography is indicated for evaluating intracranial small-vessel pathology (such as vasculitis), for assessing vascular malformations and aneurysms, and in endovascular therapeutic procedures (Table 440e-1). Angiography has been replaced for many indications by CT/CTA or MRI/MRA.

Angiography carries the greatest risk of morbidity of all diagnostic imaging procedures, owing to the necessity of inserting a catheter into a blood vessel, directing the catheter to the required location, injecting contrast material to visualize the vessel, and removing the catheter while maintaining hemostasis. Therapeutic transcatheter procedures (see below) have become important options for the treatment of some cerebrovascular diseases. The decision to undertake a diagnostic or therapeutic angiographic procedure requires careful assessment of the goals of the investigation and its attendant risks.

To improve tolerance to contrast agents, patients undergoing angiography should be well hydrated before and after the procedure. Because the femoral route is used most commonly, the femoral artery must be compressed after the procedure to prevent a hematoma from developing. The puncture site and distal pulses should be evaluated carefully after the procedure; complications can include thigh hematoma or lower extremity emboli.

COMPLICATIONS

A common femoral arterial puncture provides retrograde access via the aorta to the aortic arch and great vessels. The most feared complication of cerebral angiography is stroke. Thrombus can form on or inside the tip of the catheter, and atherosclerotic thrombus or plaque can be dislodged by the catheter or guide wire or by the force of injection and can embolize distally in the cerebral circulation. Risk factors for ischemic complications include limited experience on the part of the angiographer, atherosclerosis, vasospasm, low cardiac output, decreased oxygen-carrying capacity, advanced age, and prior history of migraine. The risk of a neurologic complication varies but is ~4% for transient ischemic attack and stroke, 1% for permanent deficit, and <0.1% for death.

Ionic contrast material injected into the cerebral vasculature can be neurotoxic if the BBB is breached, either by an underlying disease or by the injection of hyperosmolar contrast agent. Ionic contrast media are less well tolerated than nonionic media, probably because they can induce changes in cell membrane electrical potentials. Patients with dolichoectasia of the basilar artery can suffer reversible brainstem dysfunction and acute short-term memory loss during angiography, owing to the slow percolation of the contrast material and the consequent prolonged exposure of the brain. Rarely, an intracranial aneurysm ruptures during an angiographic contrast injection, causing subarachnoid hemorrhage, perhaps as a result of injection under high pressure.

SPINAL ANGIOGRAPHY

Spinal angiography may be indicated to evaluate vascular malformations and tumors and to identify the artery of Adamkiewicz (Chap. 456) prior to aortic aneurysm repair. The procedure is lengthy and requires the use of relatively large volumes of contrast; the incidence of serious complications, including paraparesis, subjective visual

blurring, and altered speech, is ~2%. Gadolinium-enhanced MRA has been used successfully in this setting, as has iodinated contrast CTA, which has promise for replacing diagnostic spinal angiography for some indications.

INTERVENTIONAL NEURORADIOLOGY

This rapidly developing field is providing new therapeutic options for patients with challenging neurovascular problems. Available procedures include detachable coil therapy for aneurysms, particulate or liquid adhesive embolization of arteriovenous malformations, stent retrieval systems for embolectomy, balloon angioplasty and stenting of arterial stenosis or vasospasm, transarterial or transvenous embolization of dural arteriovenous fistulas, balloon occlusion of carotid-cavernous and

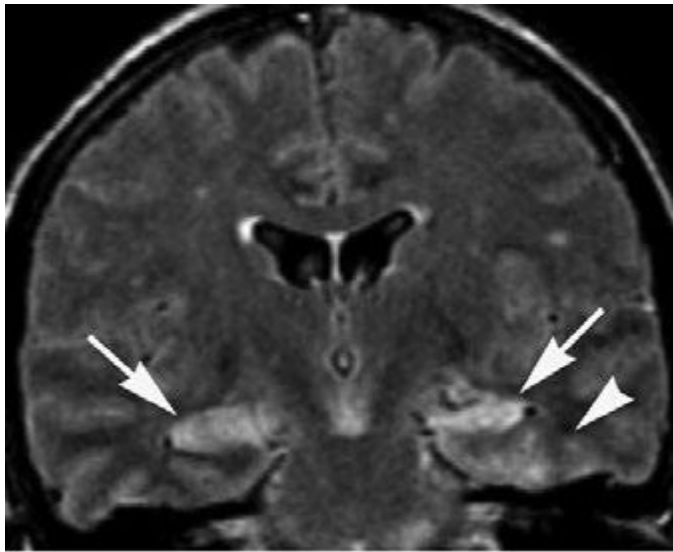
vertebral fistulas, endovascular treatment of vein-of-Galen malformations, preoperative embolization of tumors, and thrombolysis of acute arterial or venous thrombosis. Many of these disorders place the patient at high risk of cerebral hemorrhage, stroke, or death.

The highest complication rates are found with the therapies designed to treat the highest risk diseases. The advent of electrolytically detachable coils has ushered in a new era in the treatment of cerebral aneurysms. Two randomized trials found reductions of morbidity and mortality at 1 year among those treated for aneurysm with detachable coils compared with neurosurgical clipping. It remains to be determined what the role of coils will be relative to surgical options, but in many centers, coiling has become standard therapy for many aneurysms.

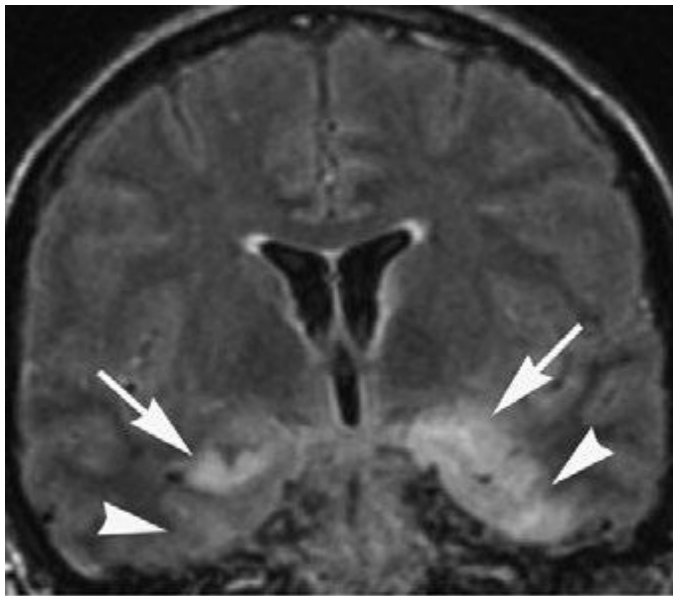
441e Atlas of Neuroimaging

Andre D. Furtado, William P. Dillon

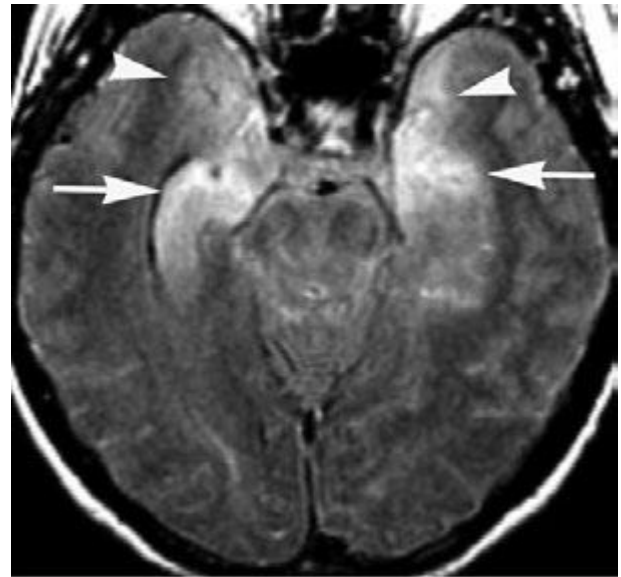
This atlas comprises 48 cases to assist the clinician caring for patients with neurologic symptoms. The majority of the images shown are magnetic resonance imaging (MRI) scans; other techniques illustrated include magnetic resonance (MR) and conventional angiography and computed tomography (CT) scans. Many different categories of neurologic disease are illustrated, including numerous examples of ischemic, inflammatory, inherited, vascular, and neoplastic etiologies.



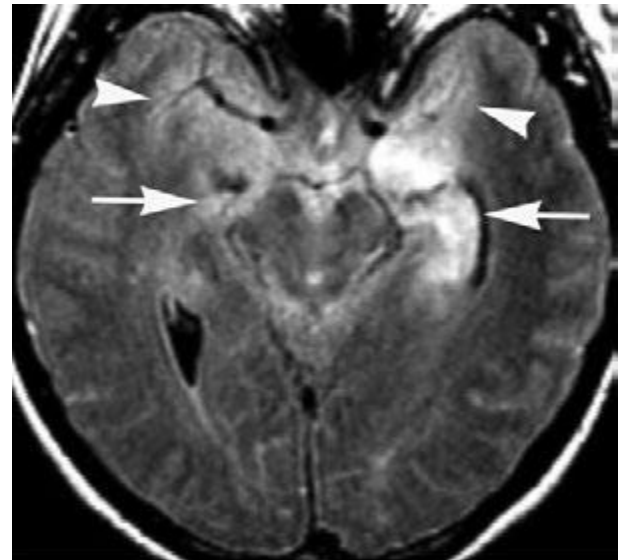
A



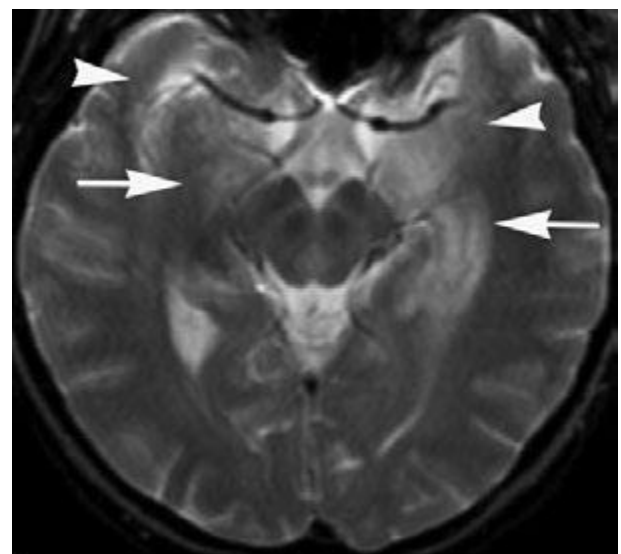
B



C



D



E

FIGURE 441e-1 Limbic encephalitis (Chap. 122). Coronal (A, B), axial fluid-attenuated inversion recovery (FLAIR) (C, D), and axial T2-weighted (E) MR images demonstrate abnormal high signal involving the bilateral mesial temporal lobes (arrowheads) including the hippocampi (left greater than right) without significant mass effect (arrows). There was no enhancement on postgadolinium images (not shown).

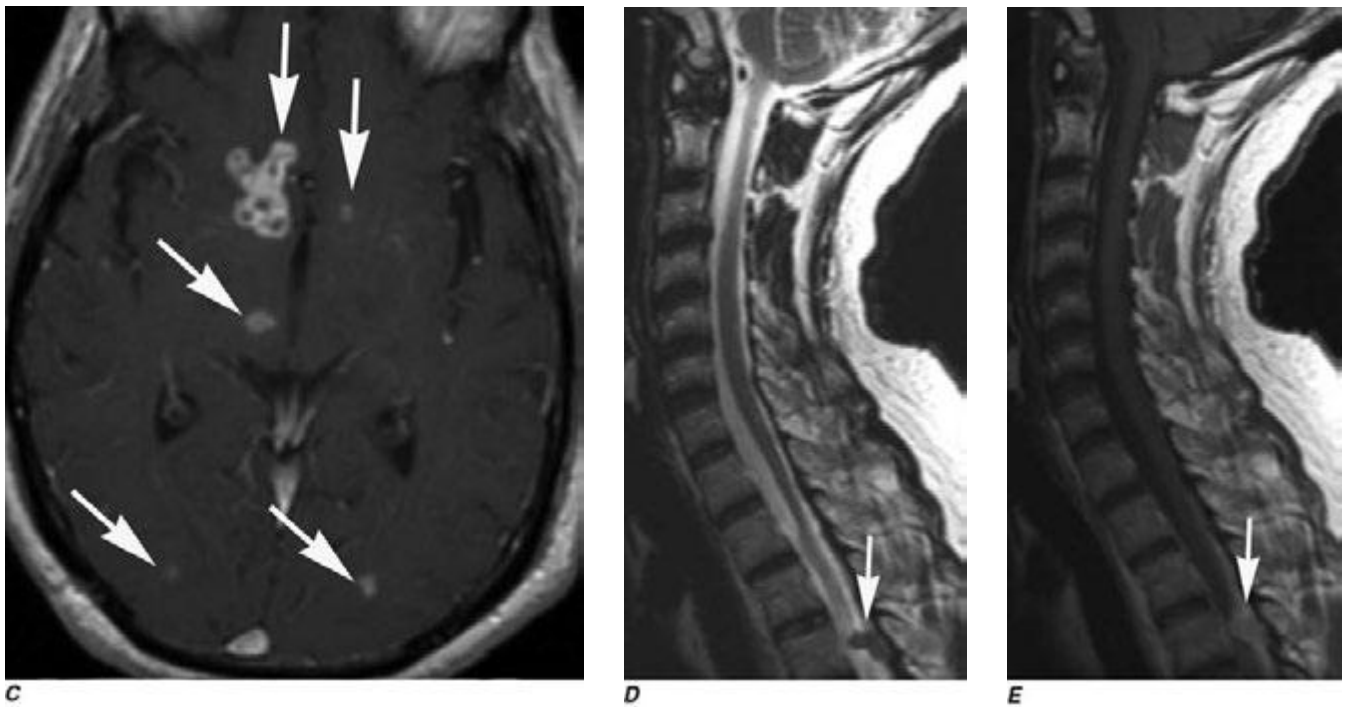
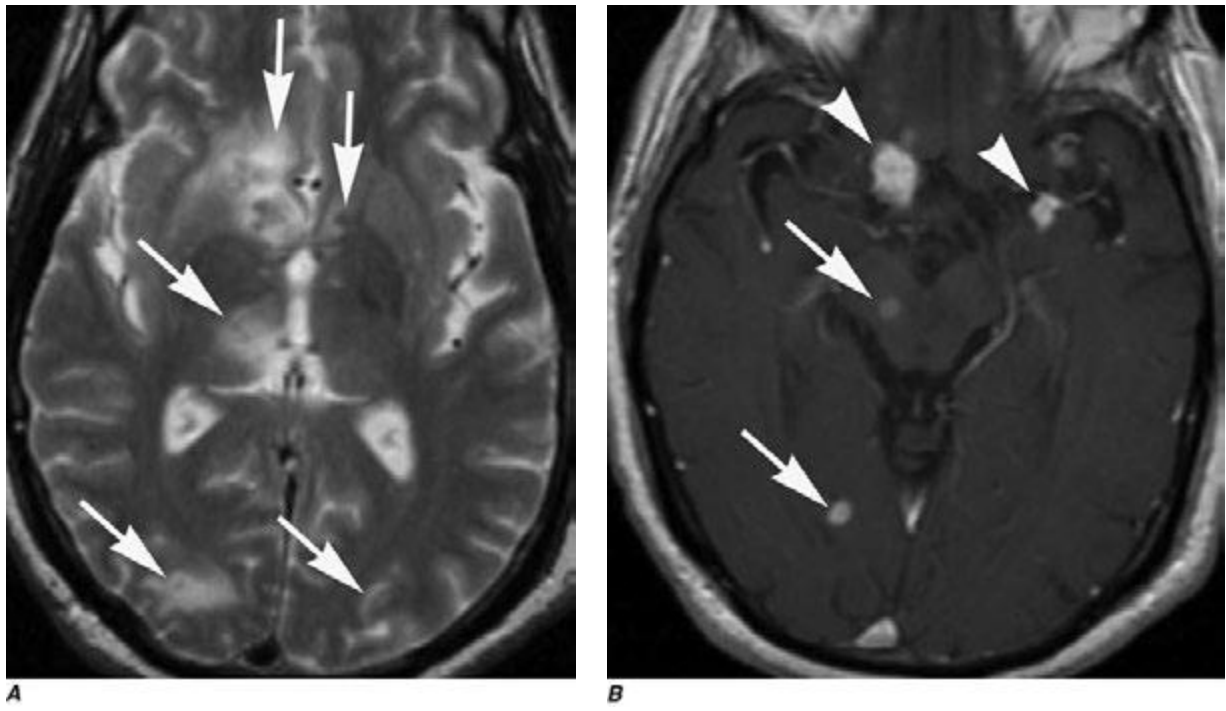


FIGURE 441e-2 Central nervous system tuberculosis (Chap. 202). Axial T2-weighted MRI (**A**) demonstrates multiple lesions (*arrows*) with peripheral high signal and central low signal, located predominantly in the cortex and subcortical white matter, as well as in the basal ganglia. Axial T1-weighted MR images postgadolinium (**B**, **C**) demonstrate ring enhancement of the lesions (*arrows*) and additional lesions in the subarachnoid space (*arrowheads*). Sagittal T2-weighted MR image of the cervical spine (**D**) demonstrates a hypointense lesion in the subarachnoid space at the level of T5 (*arrow*). Sagittal T1-weighted postgadolinium MRI of the cervical spine (**E**) demonstrates enhancement of the lesion in the subarachnoid space at the level of T5 (*arrow*).

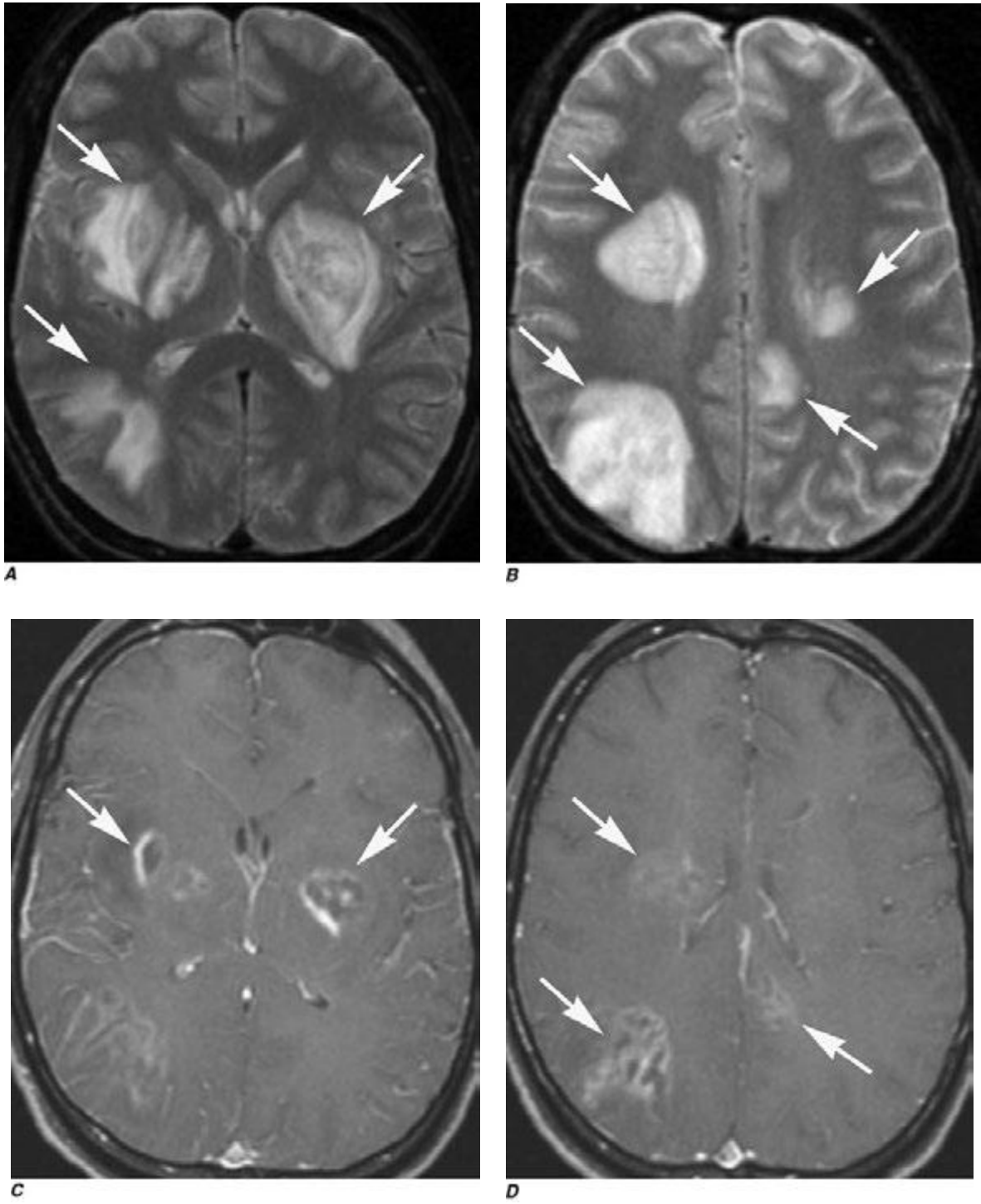
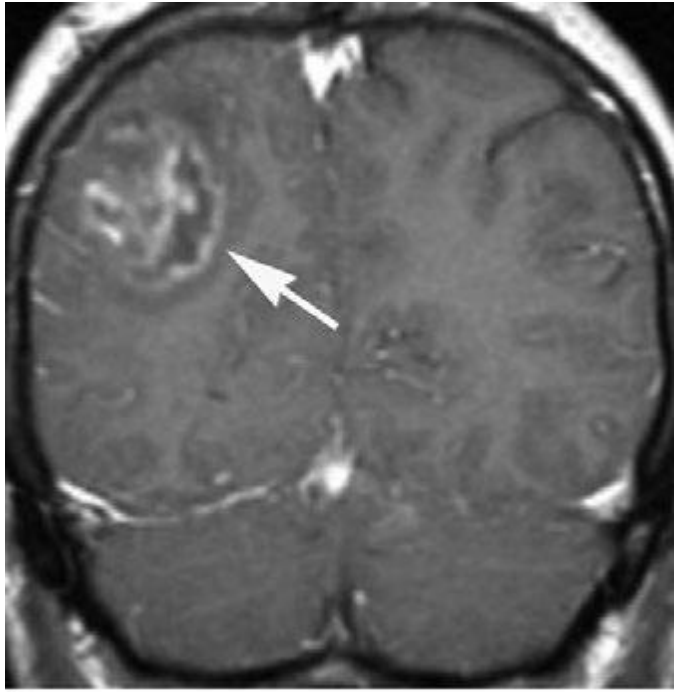
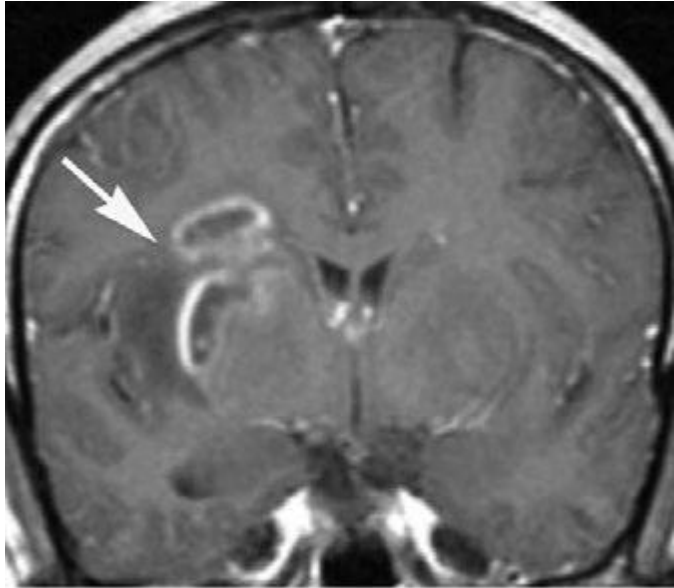


FIGURE 441e-3 Neurosyphilis (Chap. 206): Case I. Axial T2-weighted MRIs (**A, B**) demonstrate well-defined areas of abnormal high signal in the basal ganglia bilaterally and in a wedge-shaped distribution in the right parietal lobe (*arrows*). Axial (**C, D**) T1-weighted images postgadolinium. Coronal (**E, F**) T1-weighted images postgadolinium demonstrate irregular ring enhancement of the lesions (*arrows*).



E



F

FIGURE 441e-3 (Continued)

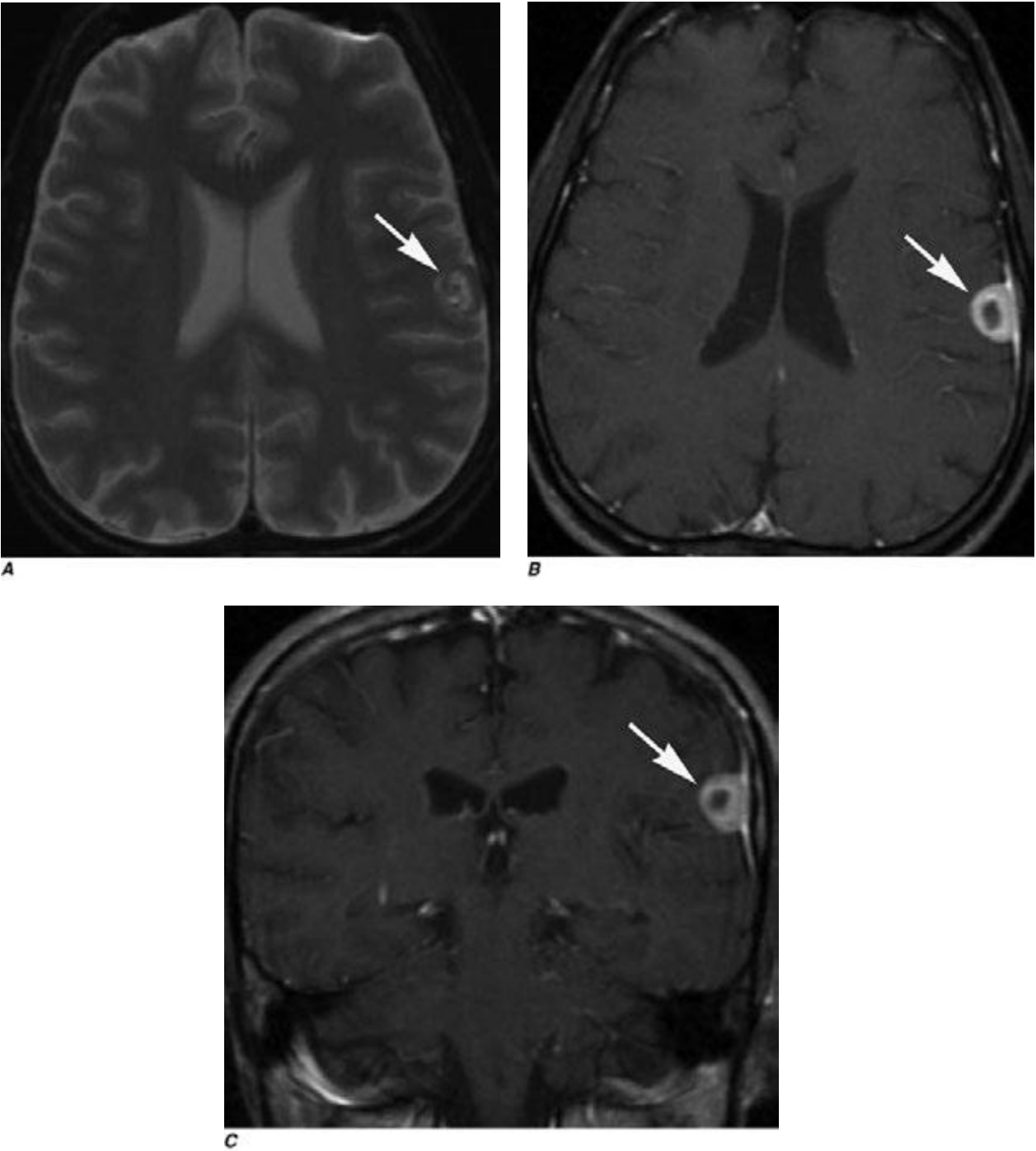
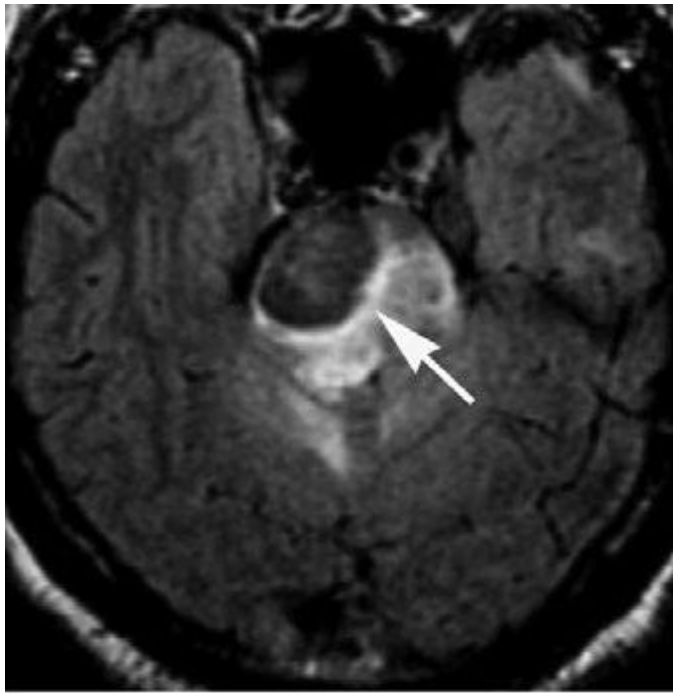
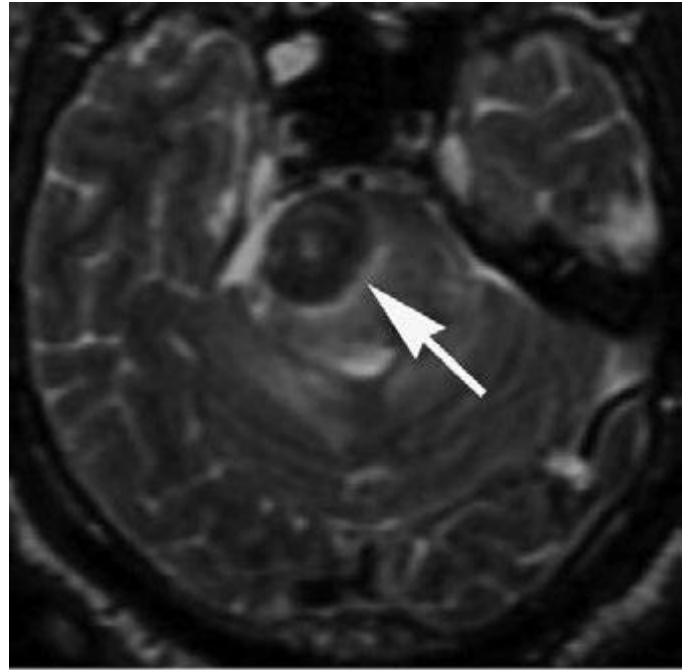


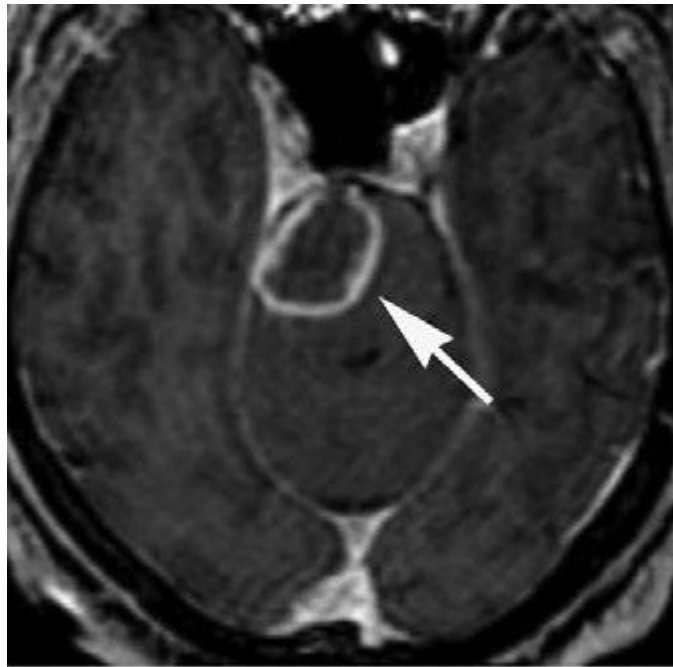
FIGURE 441e-4 Neurosyphilis (Chap. 206): Case II. Axial T2-weighted MRI (A) demonstrates a dural-based, peripherally hyperintense and centrally hypointense lesion located lateral to the left frontal lobe (*arrow*). Axial (B) and coronal (C) T1-weighted MRIs postgadolinium demonstrate peripheral enhancement of the lesion (*arrows*).



A

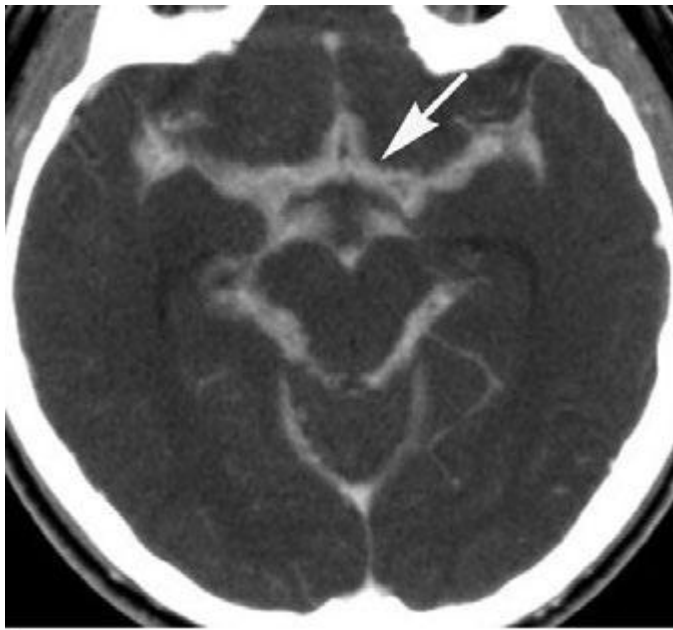


B

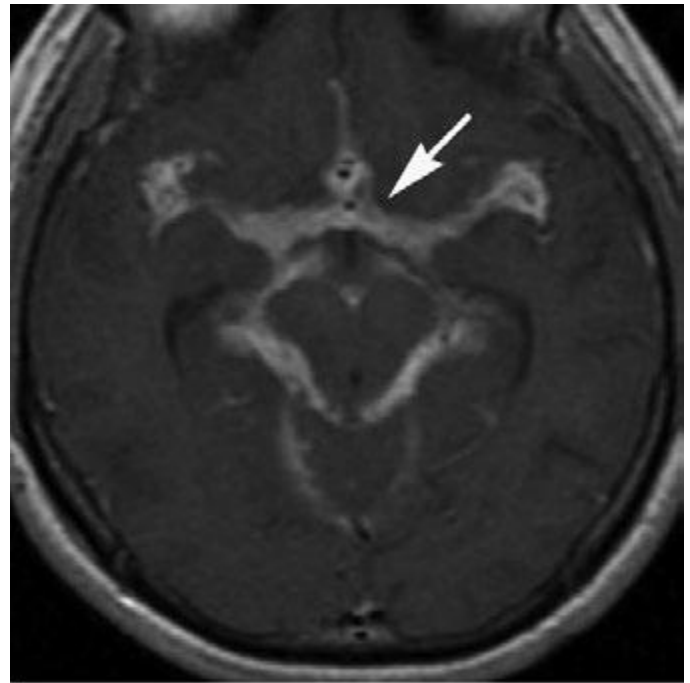


C

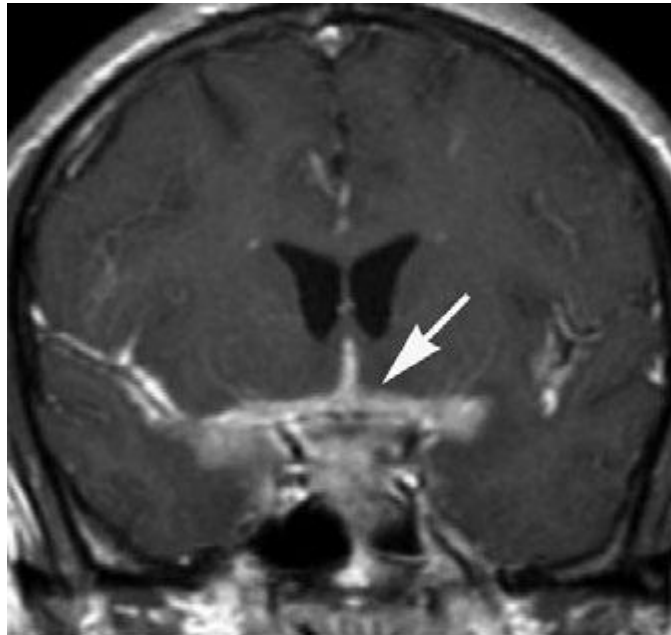
FIGURE 441e-5 Histoplasmosis of the pons (Chap. 236). Axial FLAIR (A) and T2-weighted (B) MRIs demonstrate a low signal mass in the right pons (arrows) with surrounding vasogenic edema. Axial T1-weighted MRI postgadolinium (C) demonstrates ring enhancement of the lesion in the right pons (arrow). Of note, there was no evidence of restricted diffusion (not shown).



A



B



C

FIGURE 441e-6 *Coccidiomycosis meningitis* (Chap. 237). Axial postcontrast CT (A) and axial (B) and coronal (C) T1-weighted MRIs postgadolinium demonstrate enhancement of the perimesencephalic cisterns (arrows), as well as the sylvian and interhemispheric fissures.

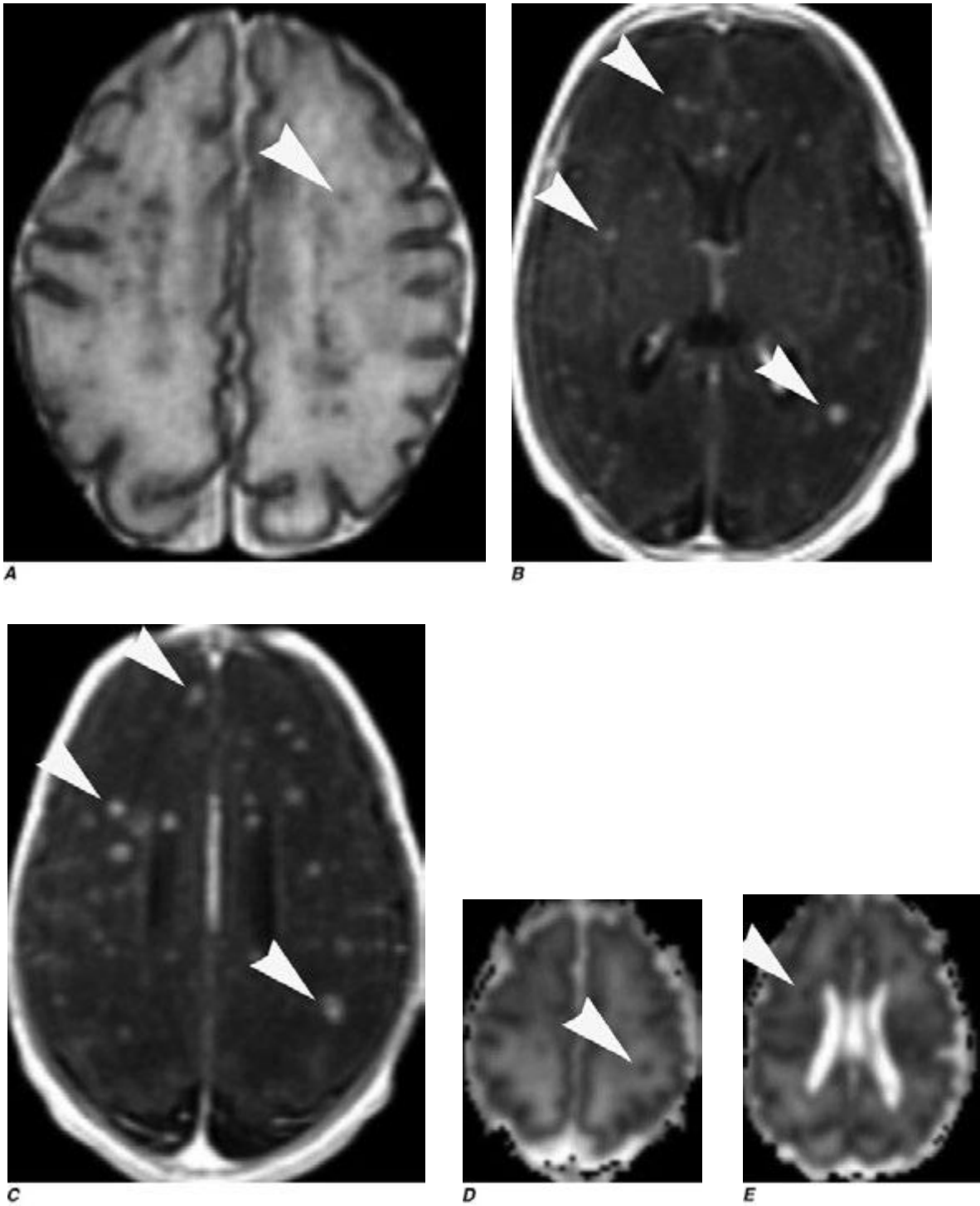


FIGURE 441e-7 Candidiasis in a newborn (Chap. 240). Axial T2-weighted MRI (A) demonstrates multiple punctate foci of low signal diffusely distributed in the brain parenchyma (arrowhead). Axial T1-weighted MRIs postgadolinium (B, C) demonstrate marked enhancement of the lesions (arrowheads). Apparent diffusion coefficient (ADC) map (D, E) demonstrates restricted diffusion of water molecules in the lesions (arrowheads).

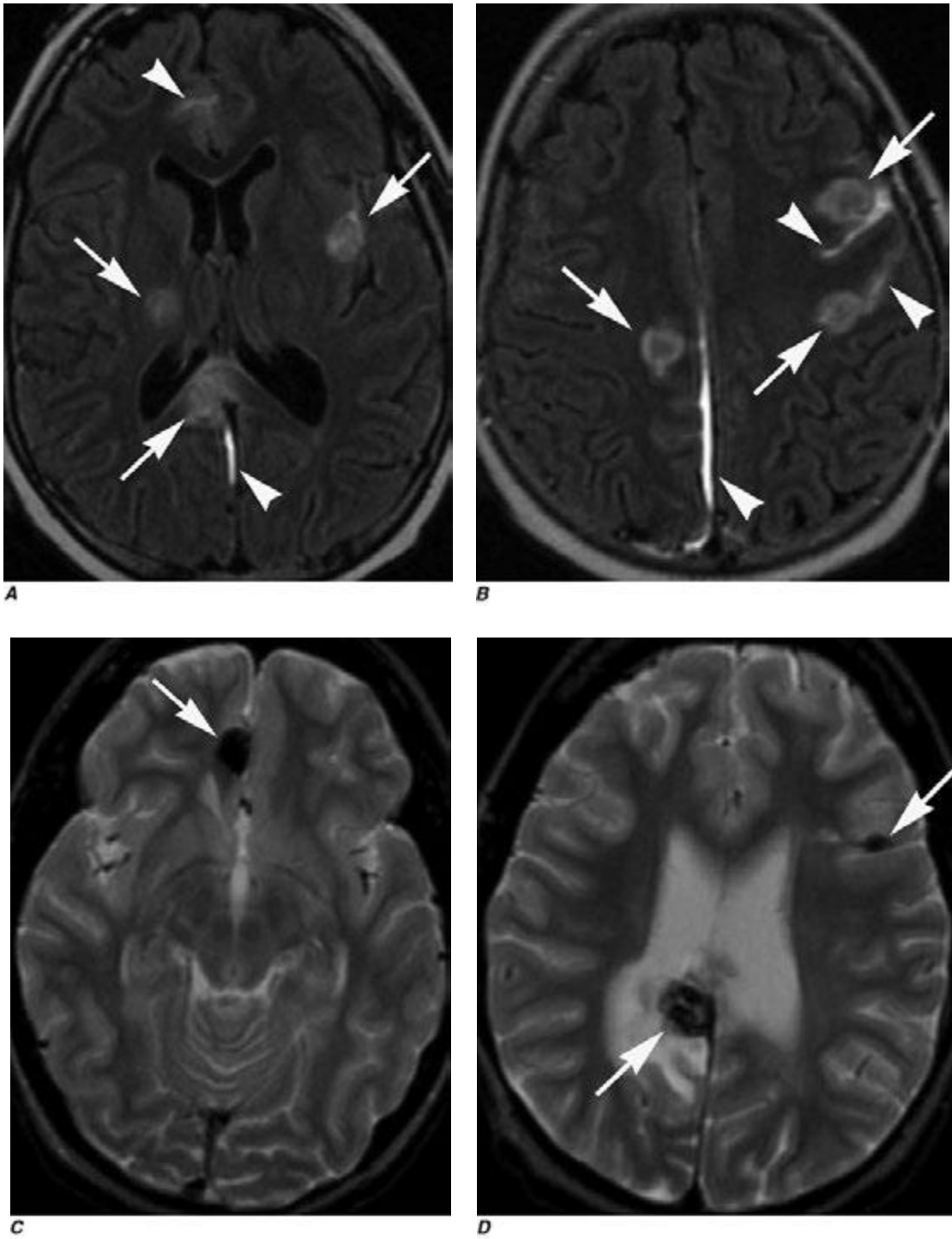
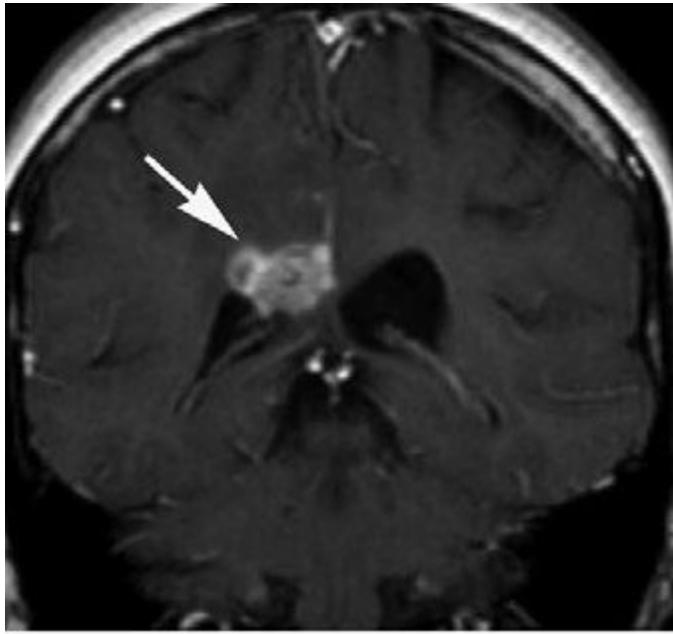
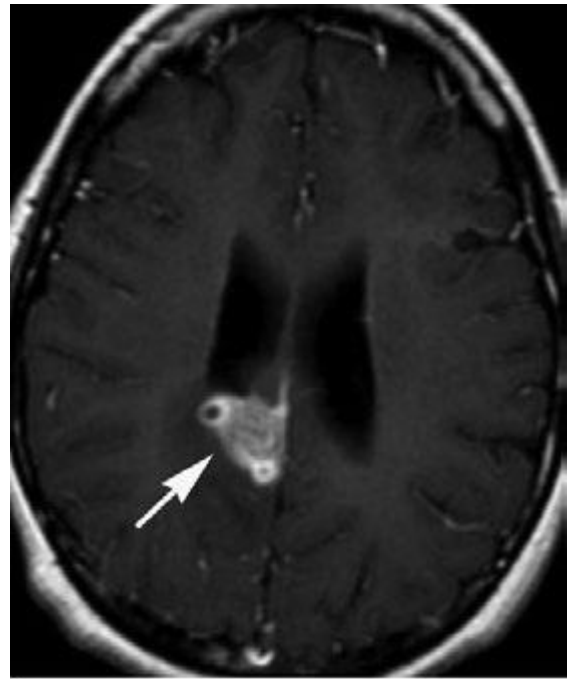


FIGURE 441e-8 Central nervous system (CNS) aspergillosis (Chap. 241). Axial FLAIR MRIs (**A**, **B**) demonstrate multiple areas of abnormal high signal in the basal ganglia as well as cortex and subcortical white matter (*arrows*). There is also abnormal high signal in the subarachnoid space adjacent to the lesions (*arrowheads*) that can correspond to blood or high protein content. Axial T2-weighted MRIs (**C**, **D**) demonstrate intrinsic low signal in the lesions (*arrows*), suggesting the presence of blood products. Some of the lesions also show vasogenic edema. Coronal (**E**) and axial (**F**) T1-weighted MRIs postgadolinium demonstrate peripheral enhancement of the lesions (*arrows*).

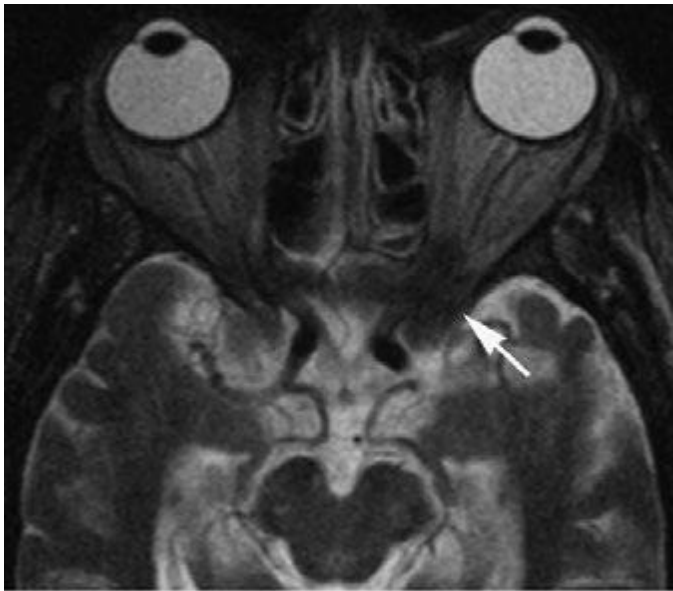


E

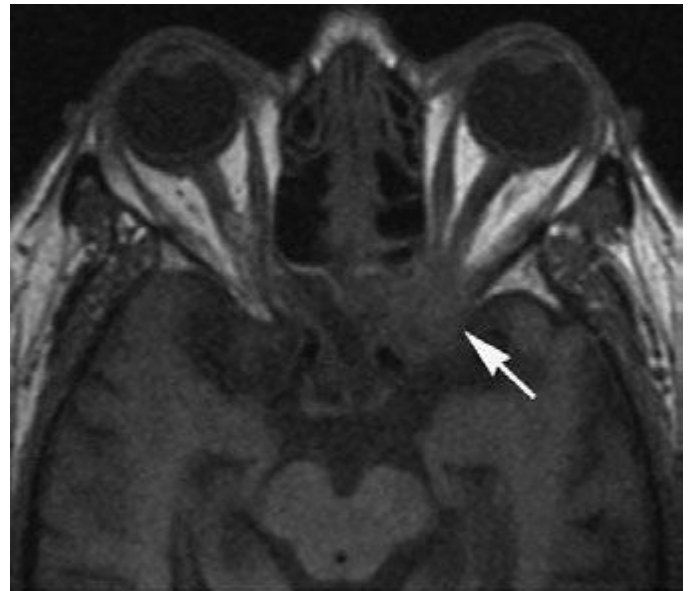


F

FIGURE 441e-8 (Continued)

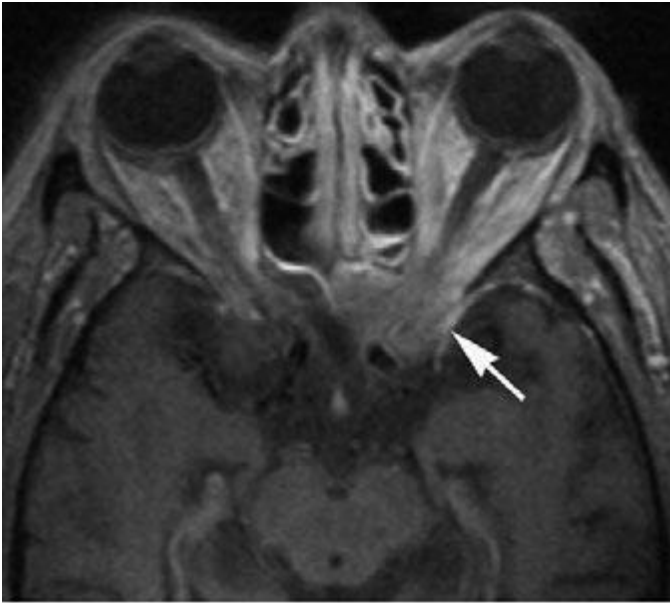


A



B

FIGURE 441e-9 Invasive sinonasal aspergillosis (Chap. 241). Axial T2-weighted MRI (A) demonstrates an irregularly shaped low signal lesion involving the left orbital apex (arrow). B. T1-weighted image pregadolinium demonstrates low signal in left anterior clinoid process (arrow). C. T1-weighted image postgadolinium demonstrates enhancement of lesion (arrow).



c

FIGURE 441e-9 (Continued)

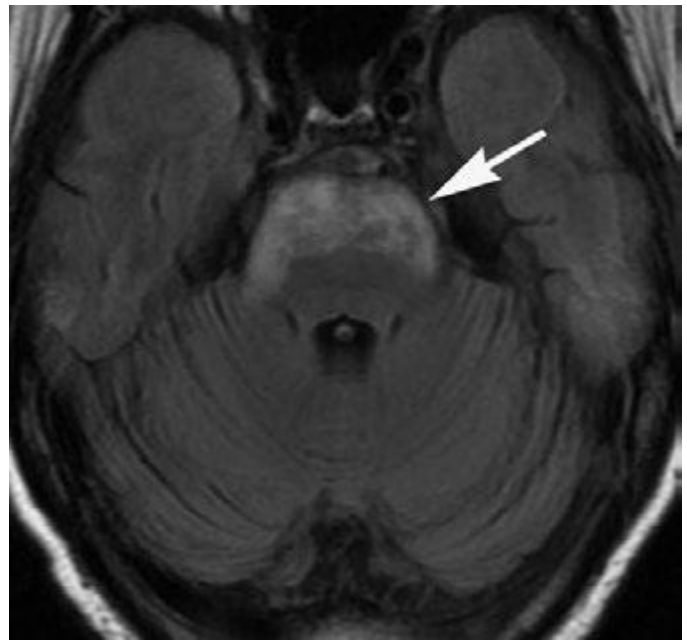
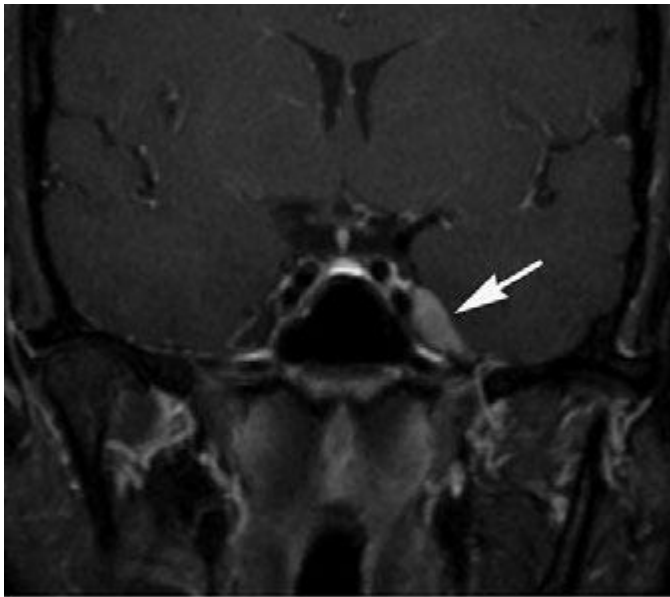
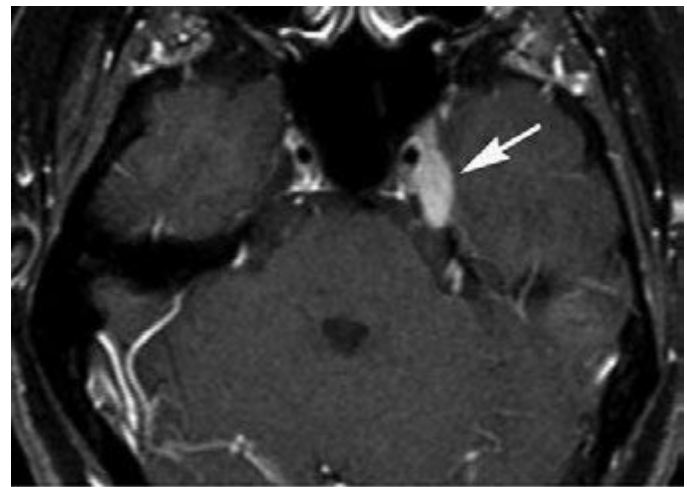


FIGURE 441e-10 Behçet's disease (Chap. 387). Axial FLAIR MRI demonstrates abnormal high signal involving the anterior pons (*arrow*); following gadolinium administration, the lesion was nonenhancing (not shown). Brainstem lesions are typical of Behçet's disease, caused primarily by vasculitis and in some cases demyelinating lesions.



A



B

FIGURE 441e-11 Neurosarcoid (Chap. 390): Case I. Coronal (A) and axial (B) T1-weighted images postgadolinium with fat suppression demonstrate a homogeneously enhancing well-circumscribed mass centered in the left Meckel's cave (*arrows*).

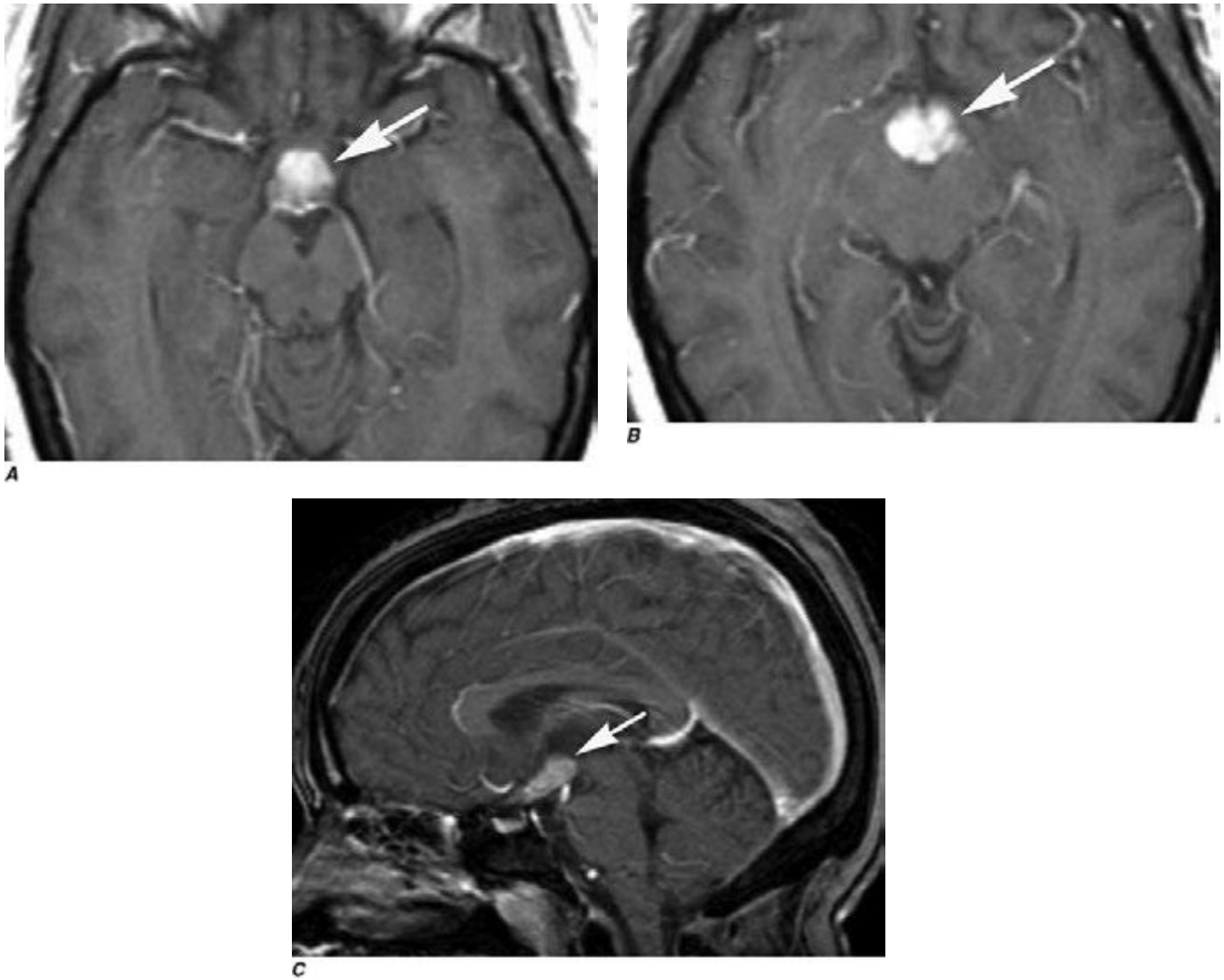


FIGURE 441e-12 Neurosarcoid (Chap. 390): Case II. Axial (*A, B*) and sagittal (*C*) T1-weighted images postgadolinium with fat suppression demonstrate a homogeneously enhancing mass involving the hypothalamus and the pituitary stalk (*arrows*).

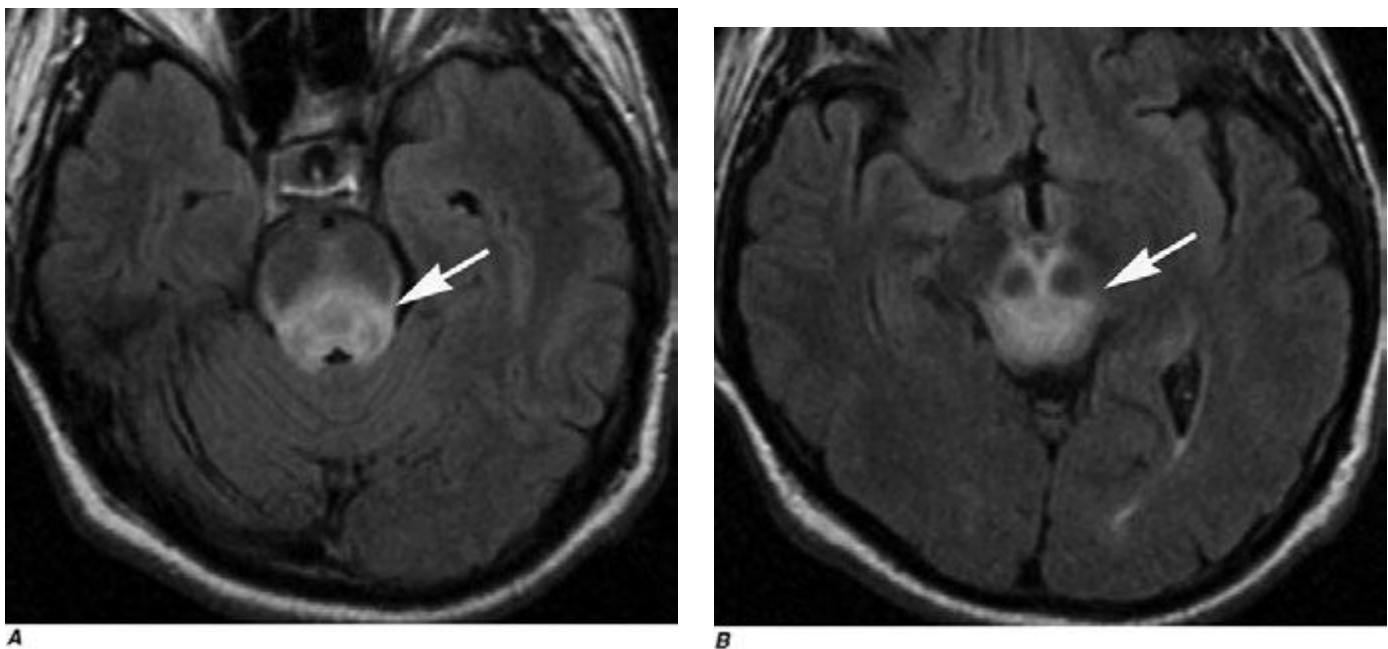
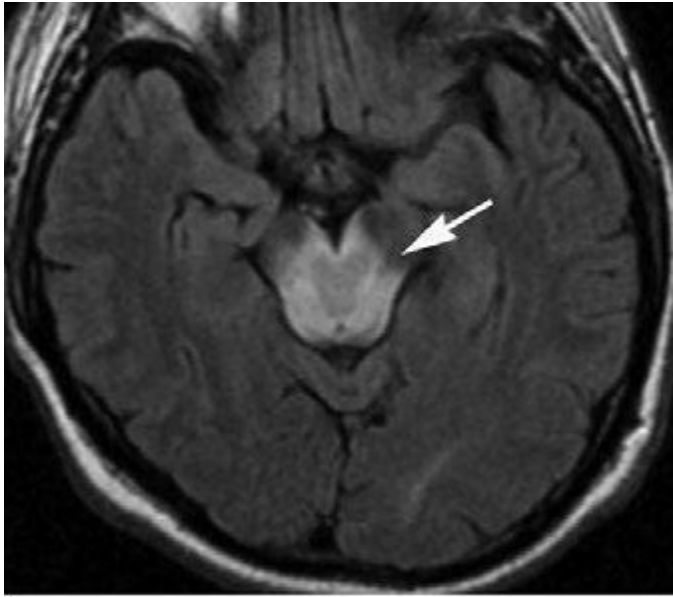
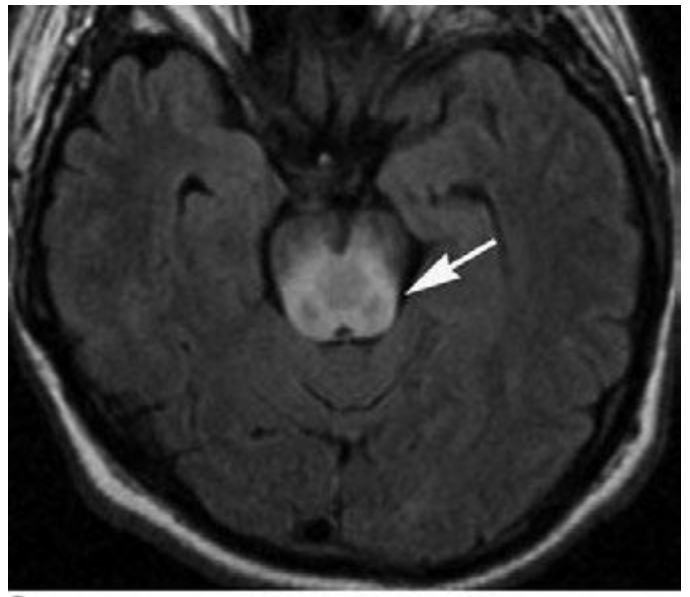


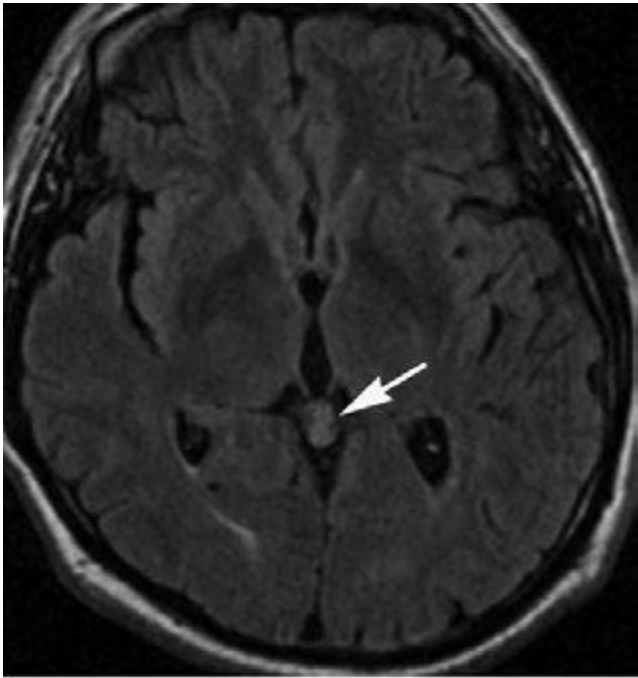
FIGURE 441e-13 Neurosarcoid (Chap. 390): Case III. Axial FLAIR images (*A–E*) demonstrate abnormal high signal and slight expansion in the midbrain, dorsal pons, and pineal region (*arrows*) without significant mass effect. Sagittal T1-weighted images postgadolinium (*F*) with fat suppression demonstrate abnormal enhancement in the midbrain, dorsal pons, and pineal region (*arrows*).



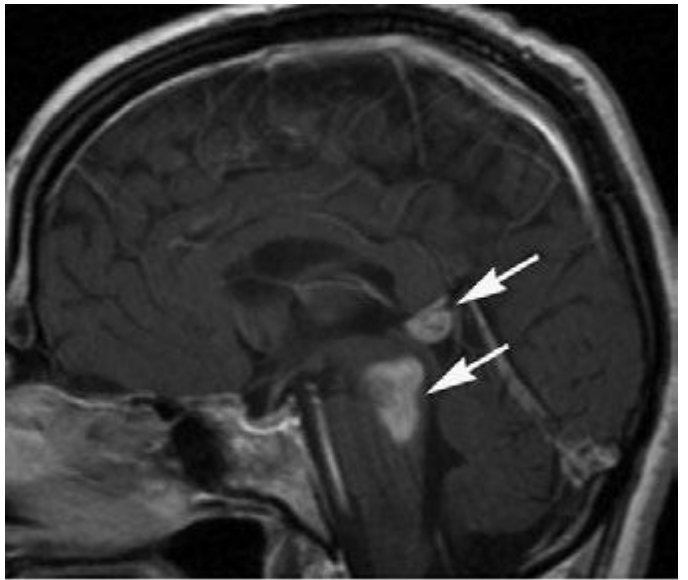
C



D

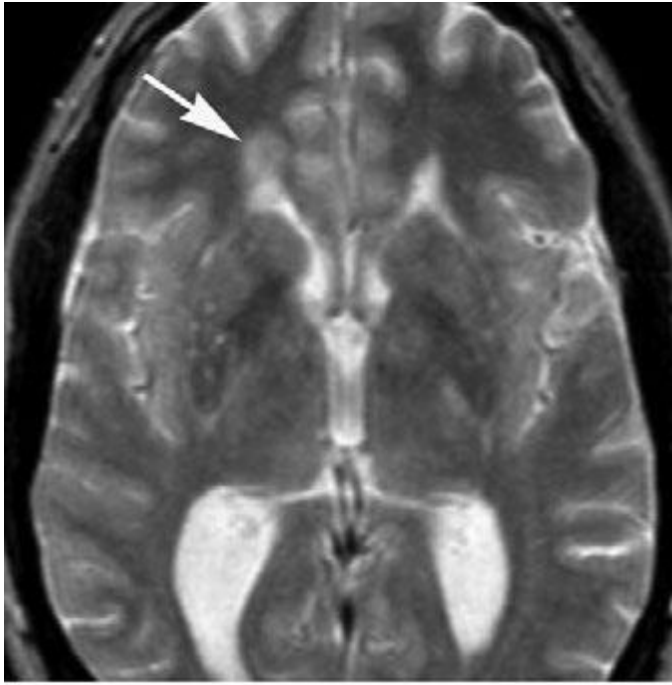


E

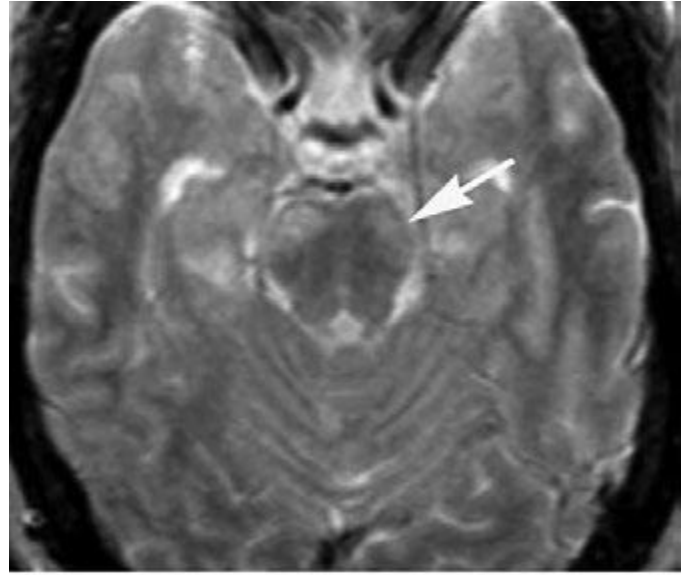


F

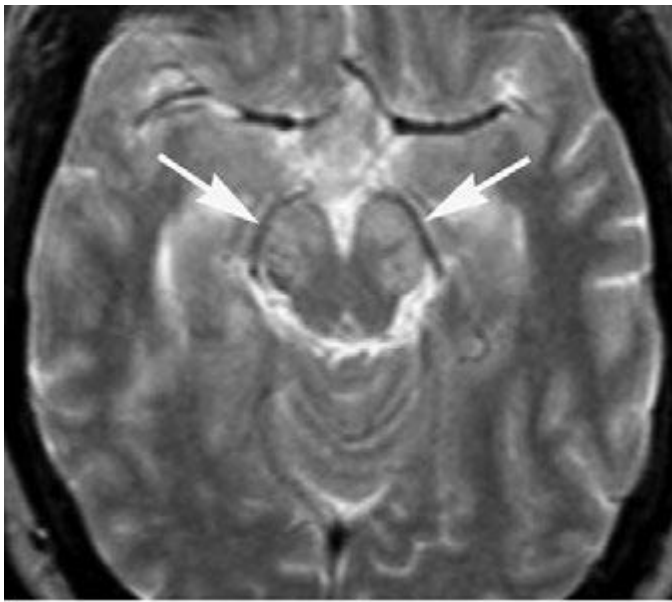
FIGURE 441e-13 (Continued)



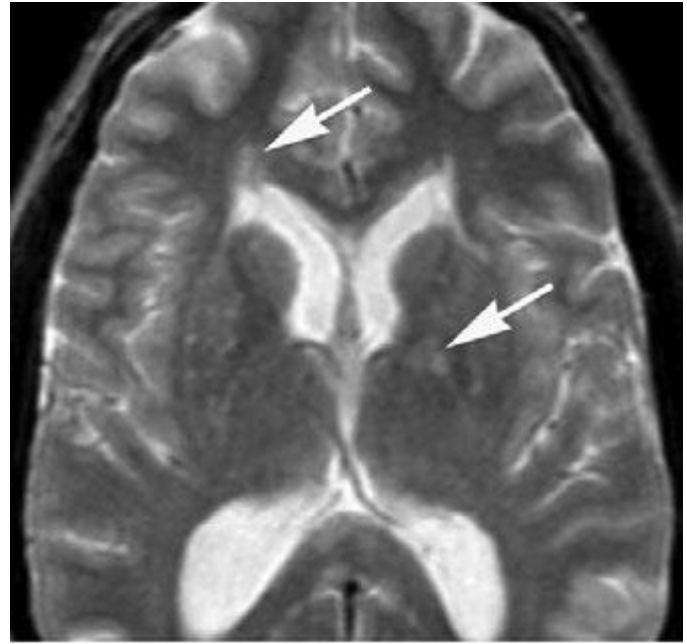
A



B

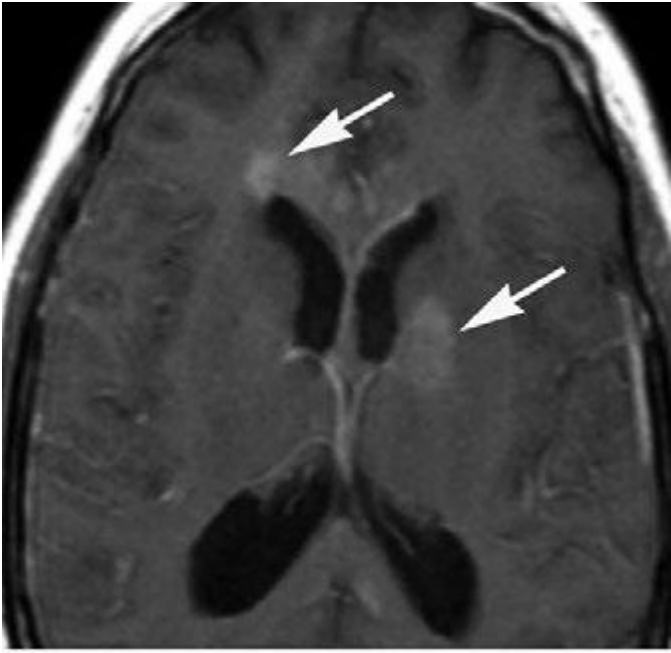


C

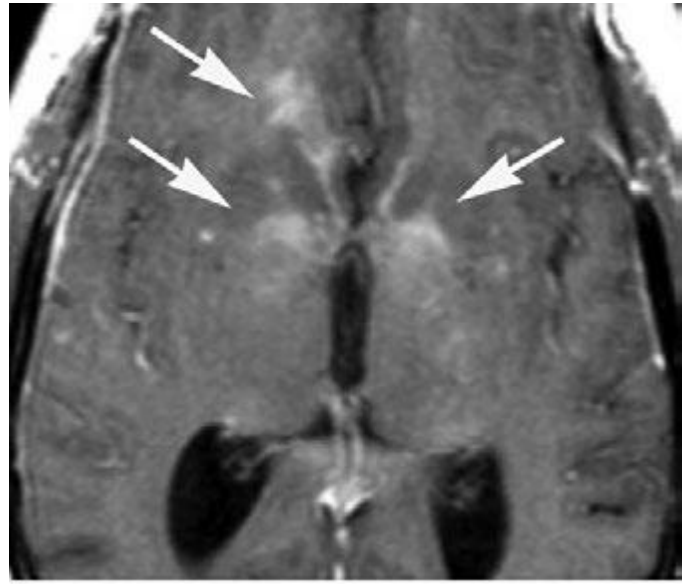


D

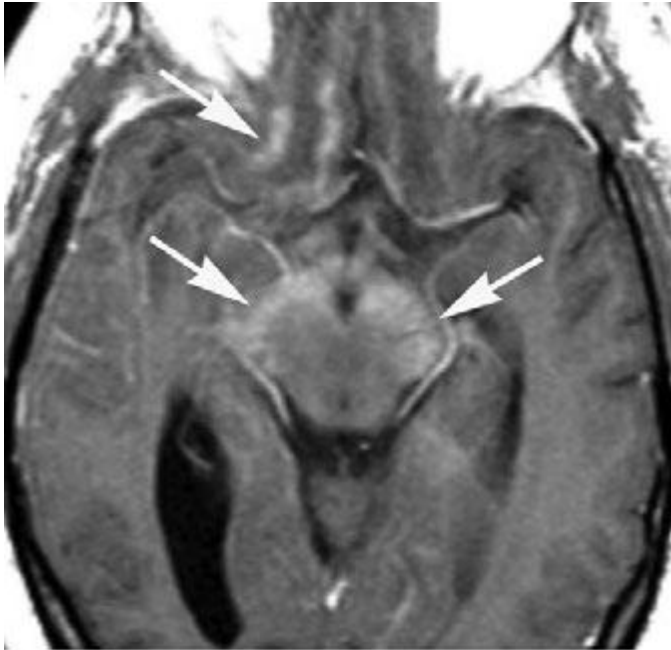
FIGURE 441e-14 Neurosarcoid (Chap. 390): Case IV. Axial T2-weighted images (A–D) demonstrate numerous areas of abnormal hyperintensity involving the corpus callosum, left internal capsule and globus pallidus, bilateral cerebral peduncles, bilateral gyrus rectus, right frontal lobe periventricular white matter, and patchy areas in bilateral temporal lobes. T1-weighted images postgadolinium (E–H) demonstrate abnormal enhancement of those areas with high T2 signal.



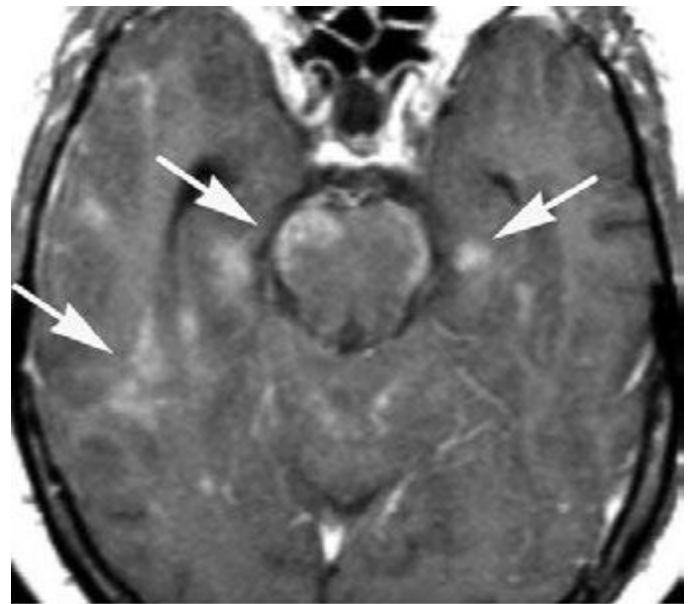
E



F



G



H

FIGURE 441e-14 (Continued)

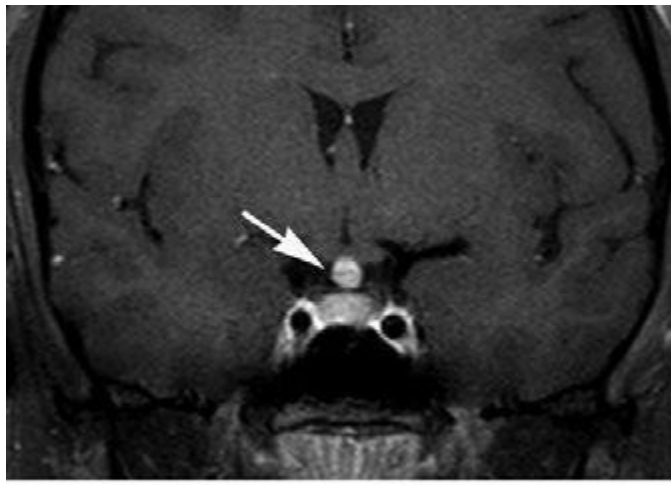
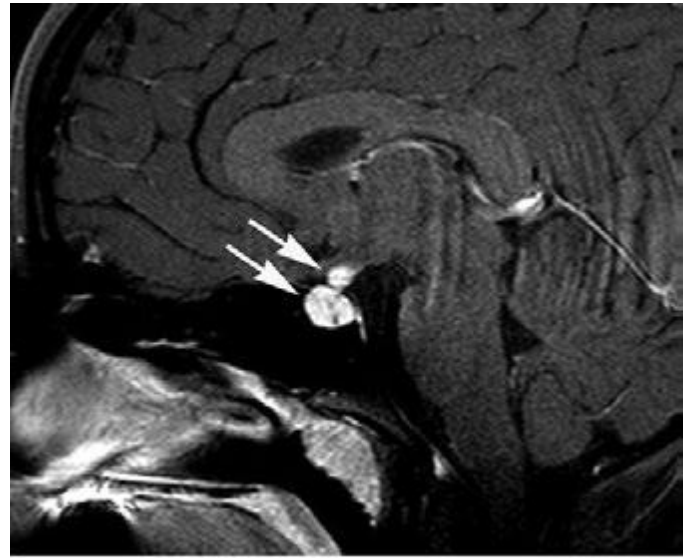
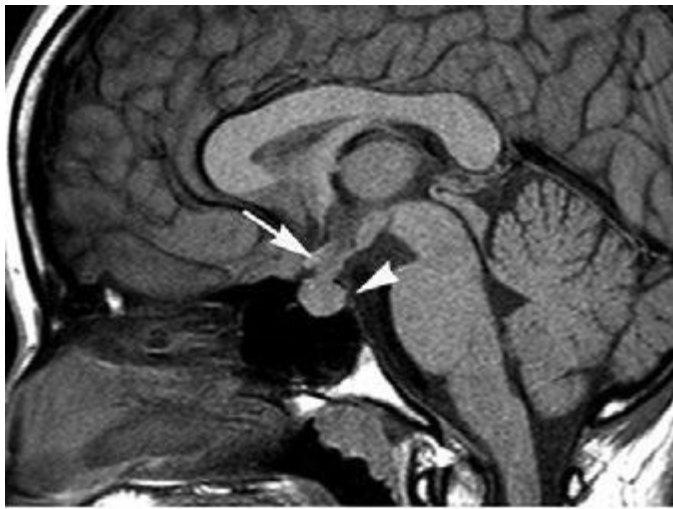


FIGURE 441e-15 Histiocytosis (Chap. 404). Sagittal T1-weighted image (A) demonstrates enlargement of the pituitary stalk (arrow) and absence of the posterior pituitary intrinsic T1 hyperintensity (arrowhead). Sagittal and coronal T1-weighted images postgadolinium (B, C) demonstrate enhancement of the pituitary stalk and infundibulum (arrows).



FIGURE 441e-16 Middle cerebral artery stenosis (Chap. 446). Time-of-flight (TOF) MR angiography (MRA) (A, B) reveals narrowing within the left M1 segment that is likely secondary to atherosclerosis (arrows).

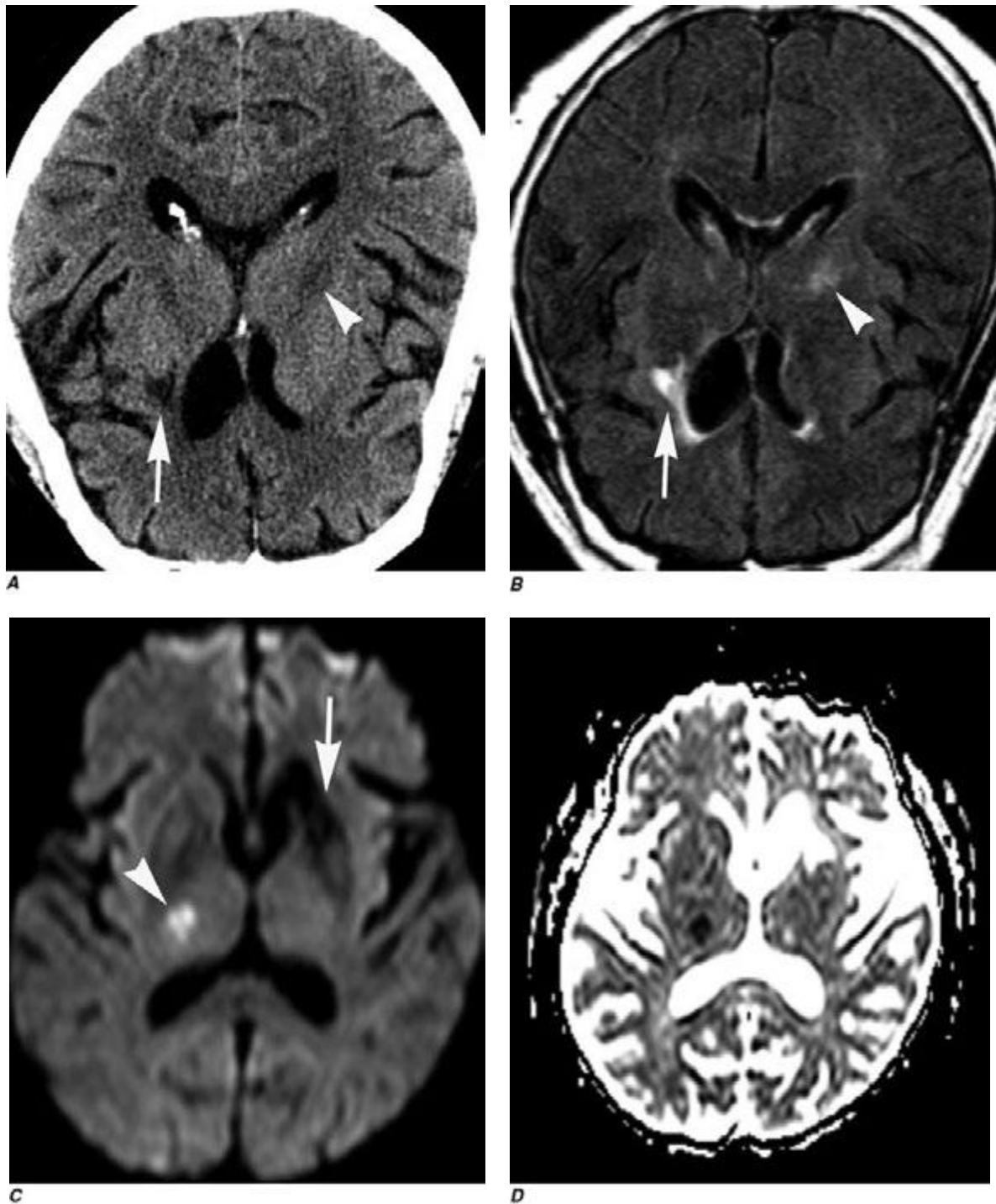
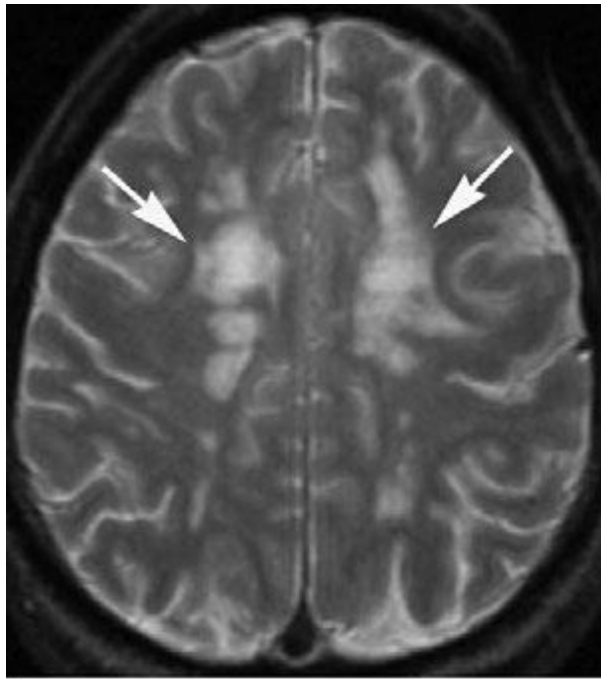
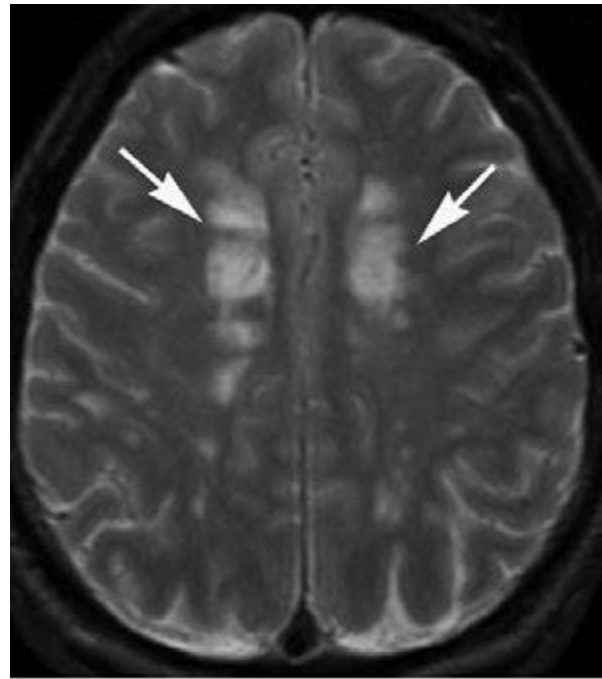


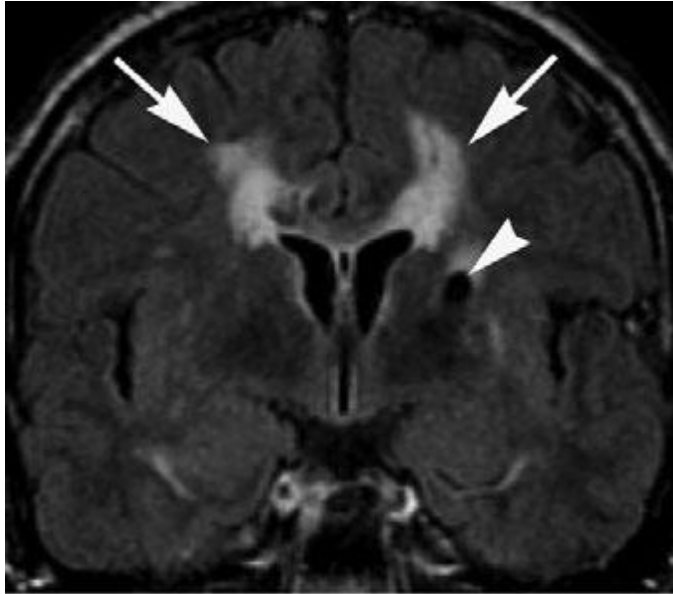
FIGURE 441e-17 Lacunar infarction (Chap. 446). Axial noncontrast CT (A) demonstrates abnormal hypodensity involving the left anterior putamen and anterior limb of internal capsule with ex-vacuo dilatation of the adjacent frontal horn of the left lateral ventricle, suggestive of an old infarction (*arrow*). A small area of slight hypodensity is also seen in the posterior limb of the right internal capsule that can correspond to an acute infarct (*arrowhead*). Axial FLAIR MRI (B) demonstrates abnormal high signal involving the left anterior putamen and anterior limb of internal capsule with ex-vacuo dilatation of the adjacent frontal horn of the left lateral ventricle, suggestive of an old infarction (*arrow*). A small area of slight hyperintensity is also seen in the posterior limb of the right internal capsule that can correspond to an acute lacunar infarct (*arrowhead*). Diffusion-weighted image (C) and apparent diffusion coefficient (ADC) map (D) demonstrate restricted water motion in the lesion of the posterior limb of the right internal capsule, strongly suggestive for an acute lacunar infarct (*arrowhead*). There is no evidence of restricted diffusion in the old infarct (*arrow*).



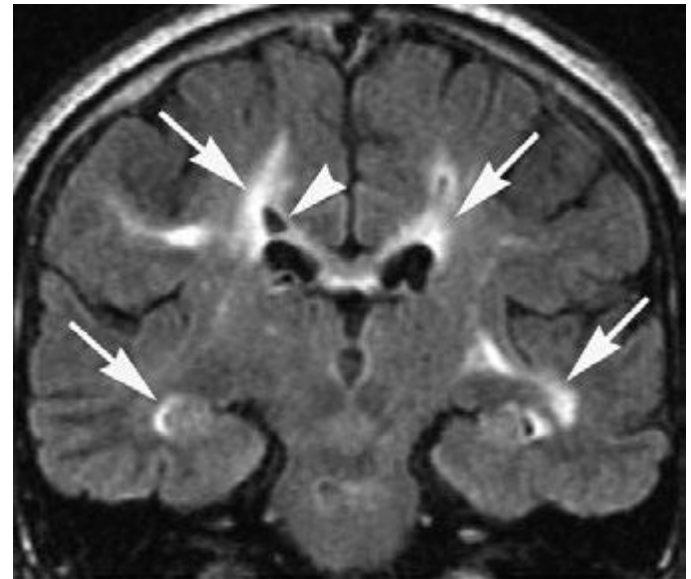
A



B



C



D

FIGURE 441e-18 Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL) (Chap. 446). Axial T2-weighted MRIs (A, B) demonstrate multiple patchy areas of abnormal high signal in the periventricular white matter (arrows). Coronal FLAIR MRI (C, D) demonstrates multiple patchy areas of abnormal high signal in the periventricular white matter bilaterally, including the temporal lobes (arrows). In some of these areas, there are small areas of tissue loss (encephalomalacia) (arrowheads).

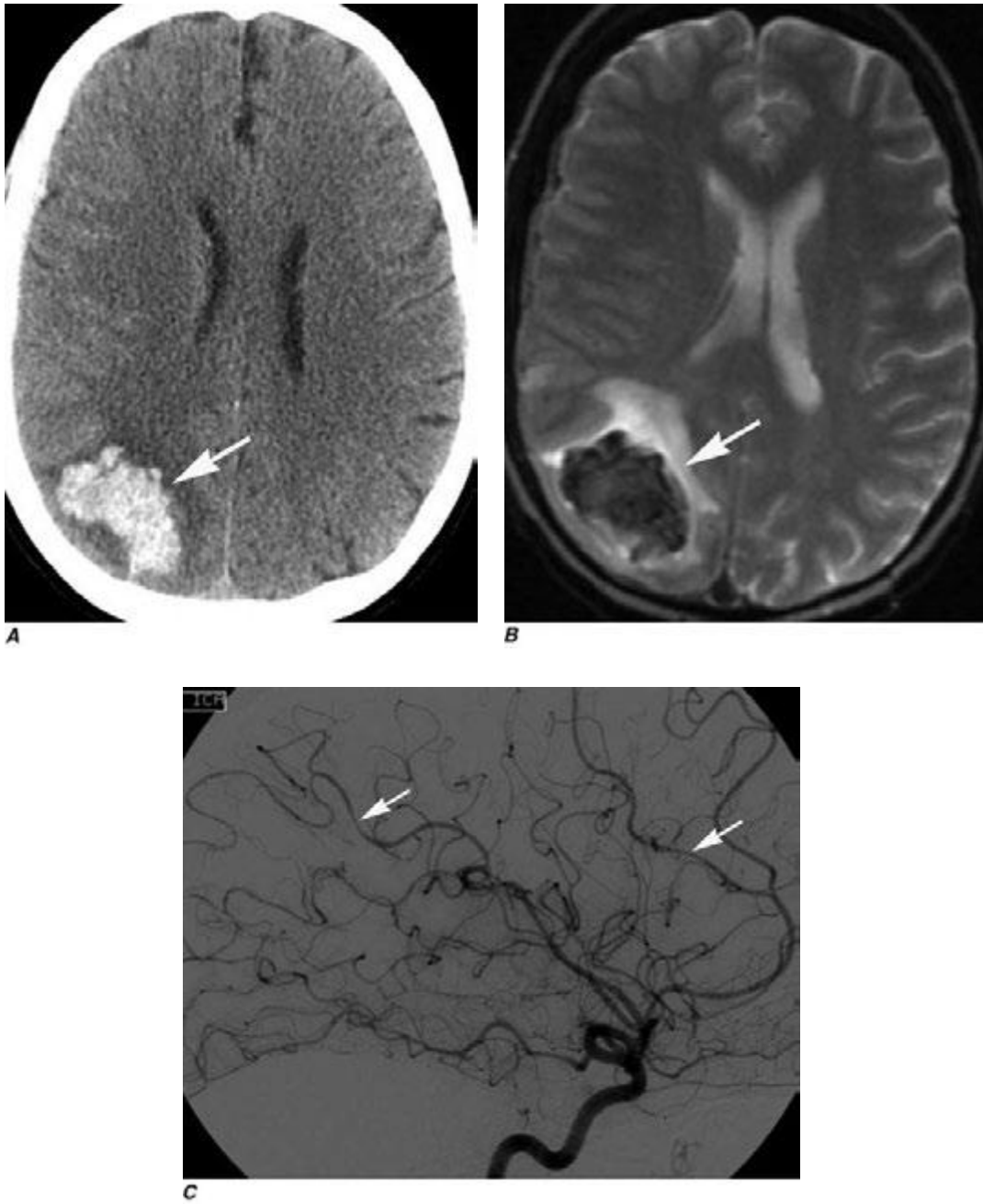


FIGURE 441e-19 CNS vasculitis (Chap. 446). Axial noncontrast CT (**A**) demonstrates a large hyperdense intraparenchymal hematoma surrounded by hypodense vasogenic edema in the right parietal lobe. Axial T2-weighted MRI (**B**) demonstrates a large hypointense intraparenchymal hematoma surrounded by hyperintense vasogenic edema in the right parietal lobe. Conventional angiography (**C**) demonstrates multiple segments of intracranial arterial narrowing, some of which have associated adjacent areas of focal arterial dilatation. These abnormalities are suggestive of vasculitis.

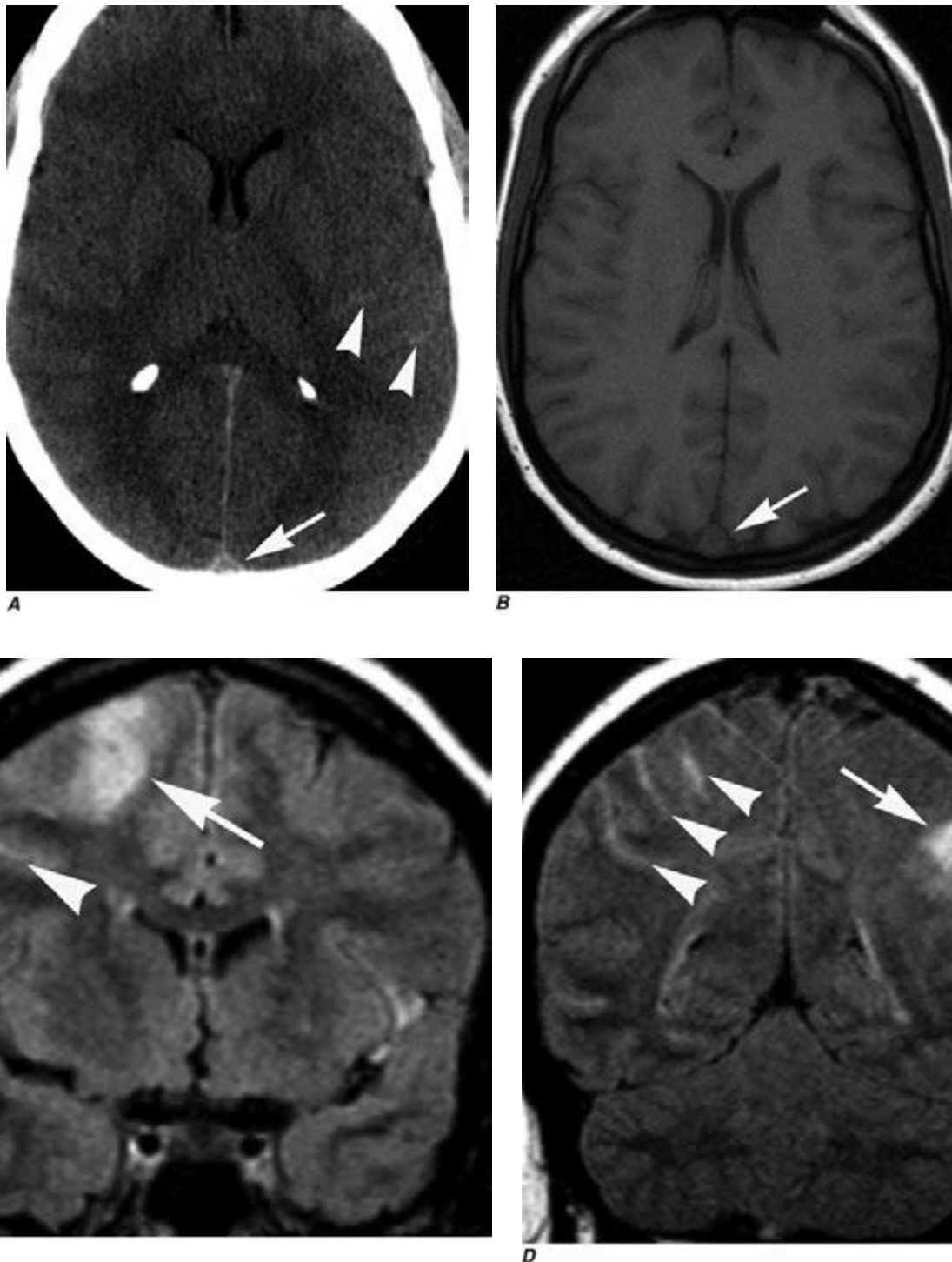
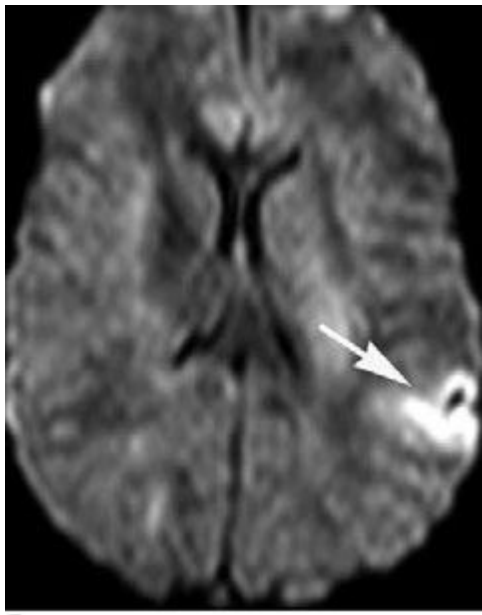
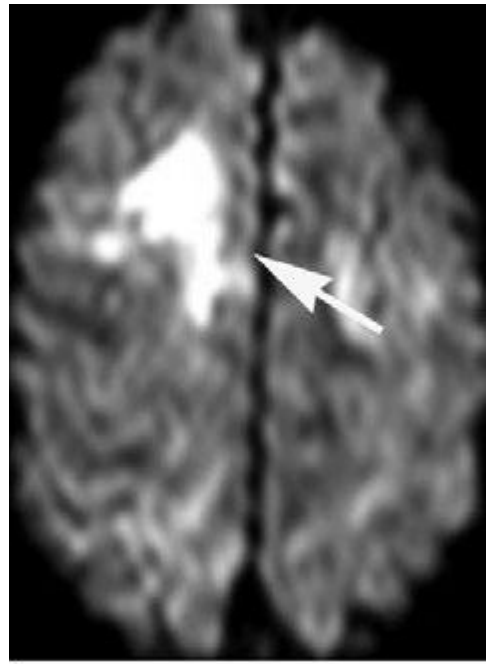


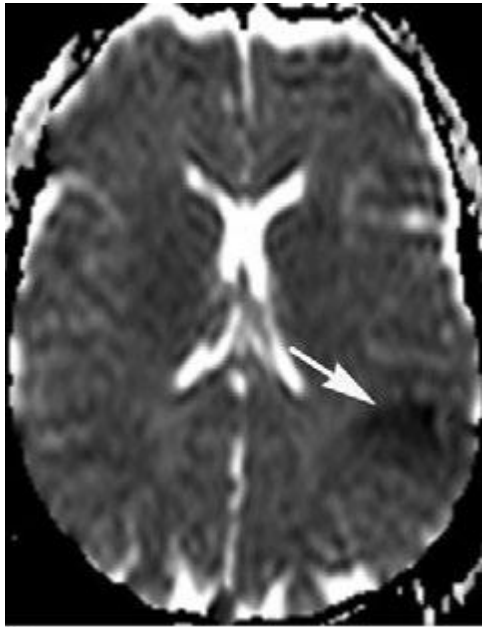
FIGURE 441e-20 Superior sagittal sinus thrombosis (Chap. 446). Noncontrast CT of the head (**A**) demonstrates increased density in the superior sagittal sinus, suggestive of thrombosis (*arrow*), and small linear hyperdensities in some temporal lobe sulci, suggestive of subarachnoid hemorrhage (*arrowheads*). Axial T1-weighted MRI (**B**) demonstrates absence of flow void in the superior sagittal sinus, suggestive of thrombosis. Coronal FLAIR images (**C**, **D**) demonstrate areas of abnormal high signal involving the gray and the subcortical white matter of the right frontal and left parietal lobes, as well as the adjacent sulci. These findings are suggestive of vasogenic edema with subarachnoid hemorrhage (*arrowheads*). Diffusion-weighted images (**E**, **F**) and ADC maps (**G**, **H**) demonstrate restricted diffusion of the abnormal areas on FLAIR, suggestive of infarct. Phase-contrast venography of the brain (**I**) demonstrates absence of signal in the superior sagittal sinus down to the torcular herophili, and left transverse sinus and jugular vein. Axial (**J**) and coronal (**K**) T1-weighted images postgadolinium demonstrate a filling defect in the superior sagittal sinus, suggestive of thrombosis.



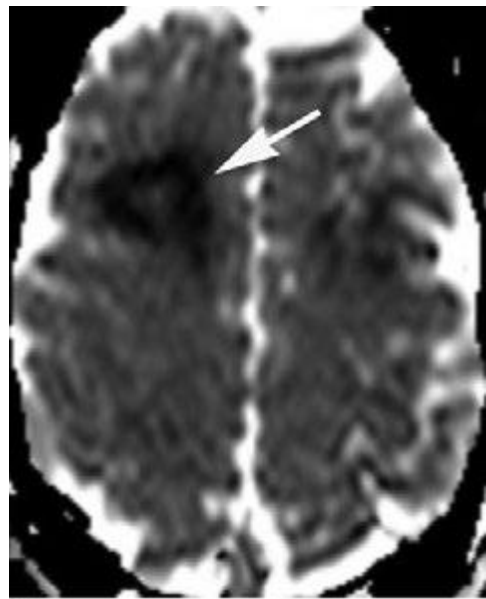
E



F



G



H

FIGURE 441e-20 (Continued)

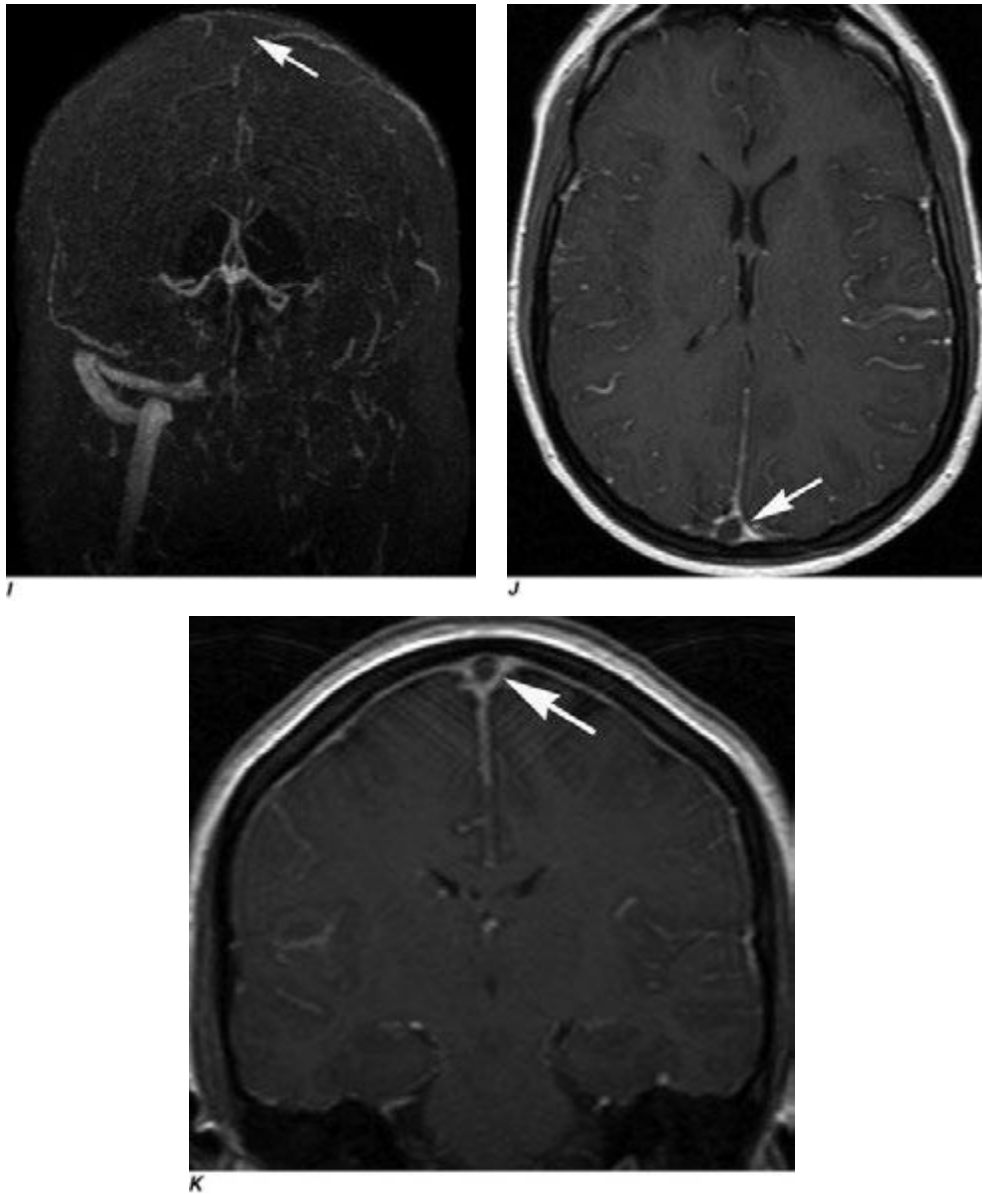


FIGURE 441e-20 (Continued)

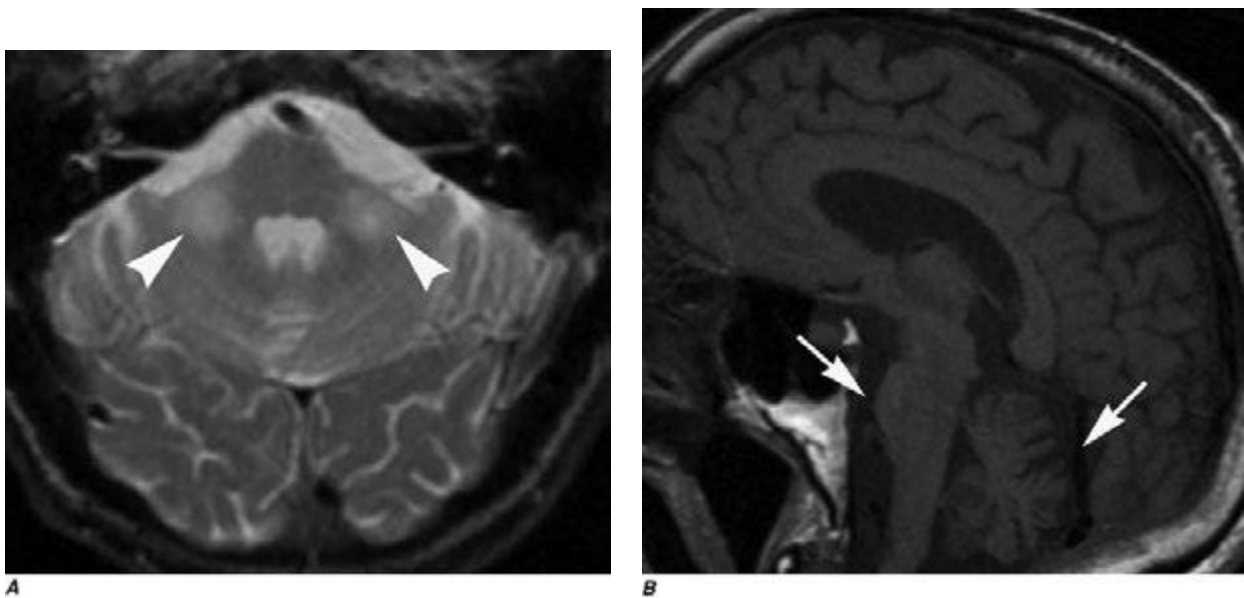


FIGURE 441e-21 **Multiple system atrophy** (Chap. 449). Axial T2-weighted MRI (**A**) reveals symmetric poorly circumscribed abnormal high signal in the middle cerebellar peduncles bilaterally (*arrowheads*). Sagittal T1-weighted MRI (**B**) demonstrates pontine atrophy and enlarged cerebellar fissures as a result of cerebellar atrophy (*arrows*).

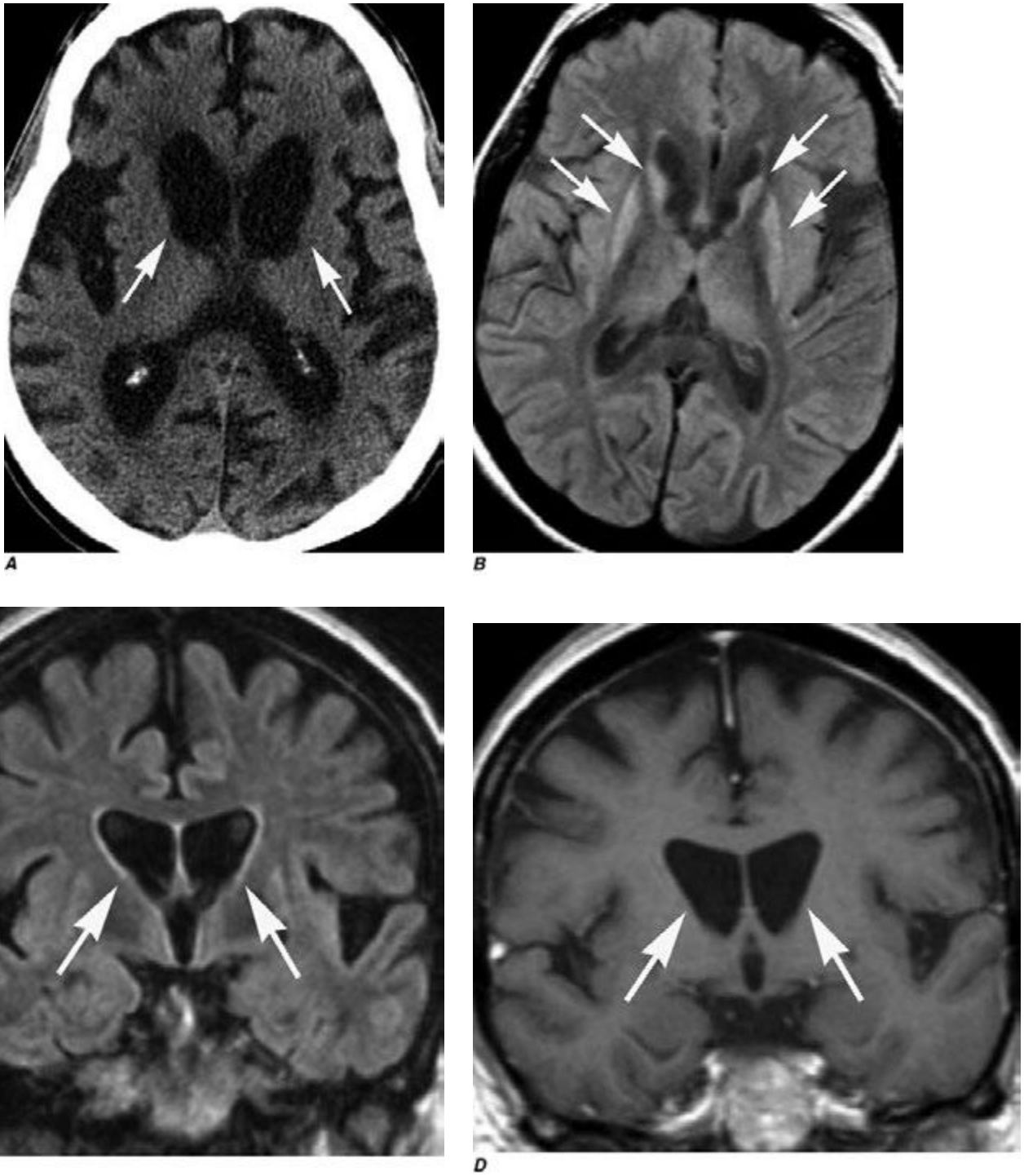
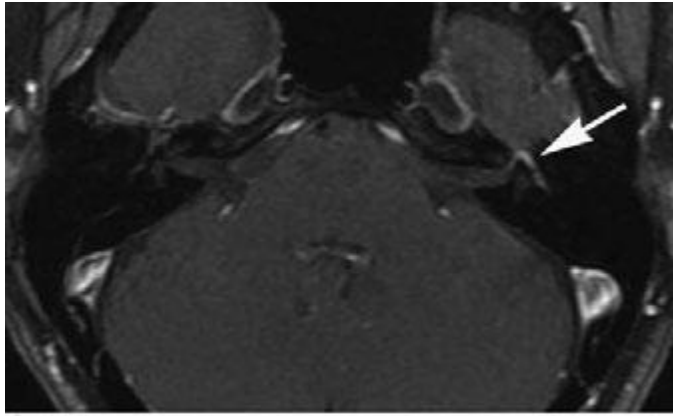
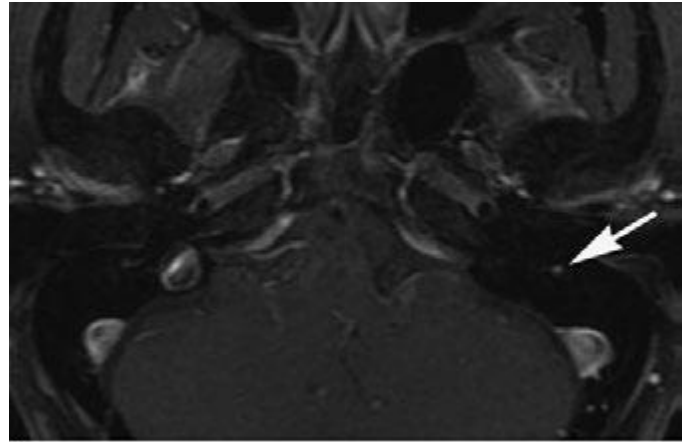


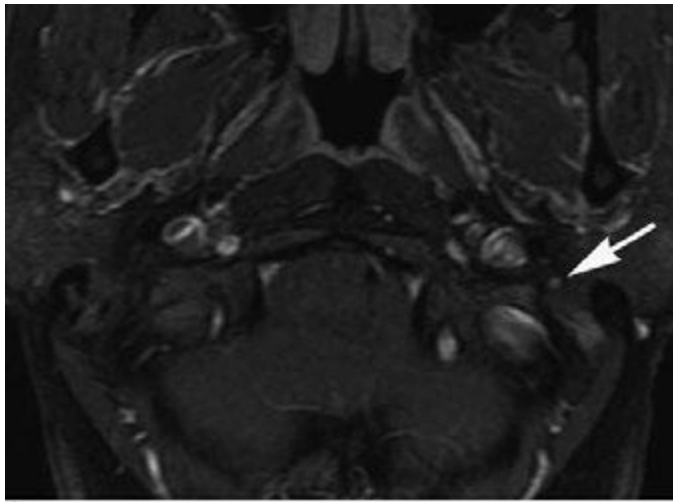
FIGURE 441e-22 Huntington's disease (Chap. 449). Axial noncontrast CT (A) demonstrates symmetric bilateral severe atrophy involving the caudate nuclei, putamen, and globus pallidi bilaterally with consequent enlargement of the frontal horns of the lateral ventricles (arrows). There is also diffuse prominence of the sulci indicating generalized cortical atrophy. Axial (B) and coronal (C) FLAIR images demonstrate bilateral symmetric abnormal high signal in the caudate and putamen. Coronal T1-weighted image (D) demonstrates enlarged frontal horns with abnormal configuration. Also note diffusely decreased marrow signal, which could represent anemia or myeloproliferative disease.



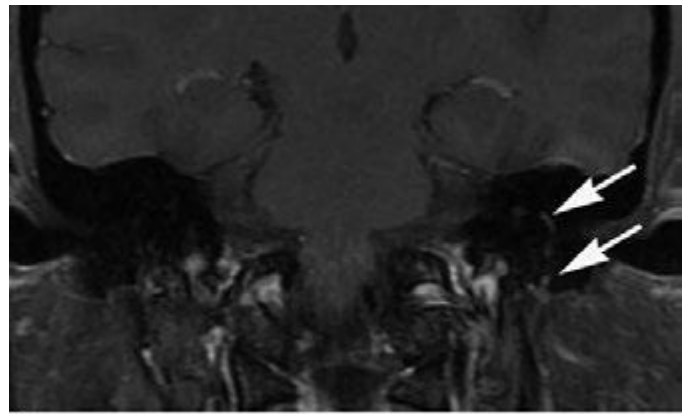
A



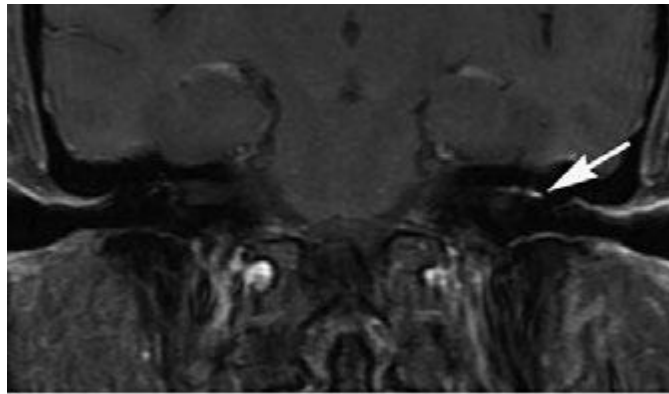
B



C



D



E

FIGURE 441e-23 Bell's palsy (Chap. 455). Axial T1-weighted images postgadolinium with fat suppression (**A–C**) demonstrate diffuse smooth linear enhancement along the left facial nerve, involving the second and third segments (genu, tympanic, and mastoid) within the temporal bone (*arrows*). Note that there is no evidence of a mass lesion. A potential pitfall for facial nerve enhancement in the stylomastoid foramen is the enhancement of the stylomastoid artery that enters the foramen and supplies the tympanic cavity, the tympanic antrum, mastoid cells, and the semicircular canals. Coronal T1-weighted images postgadolinium with fat suppression (**D, E**) demonstrate the course of the enhancing facial nerve (*arrows*). Although these findings are highly suggestive of Bell's palsy, the diagnosis is established on clinical grounds.



FIGURE 441e-24 Spinal cord infarction (Chap. 456). Sagittal T2-weighted MRI of the lumbar spine (**A**) demonstrates poorly defined areas of abnormal high signal in the conus medullaris and mild cord expansion (*arrow*). T1-weighted MRI of the lumbar spine postgadolinium (**B**) demonstrates mild enhancement (*arrow*). Sagittal diffusion-weighted MRI of the lumbar spine (**C**) demonstrates restricted diffusion (*arrow*) in the areas of abnormal high signal on the T2-weighted image (**A**).

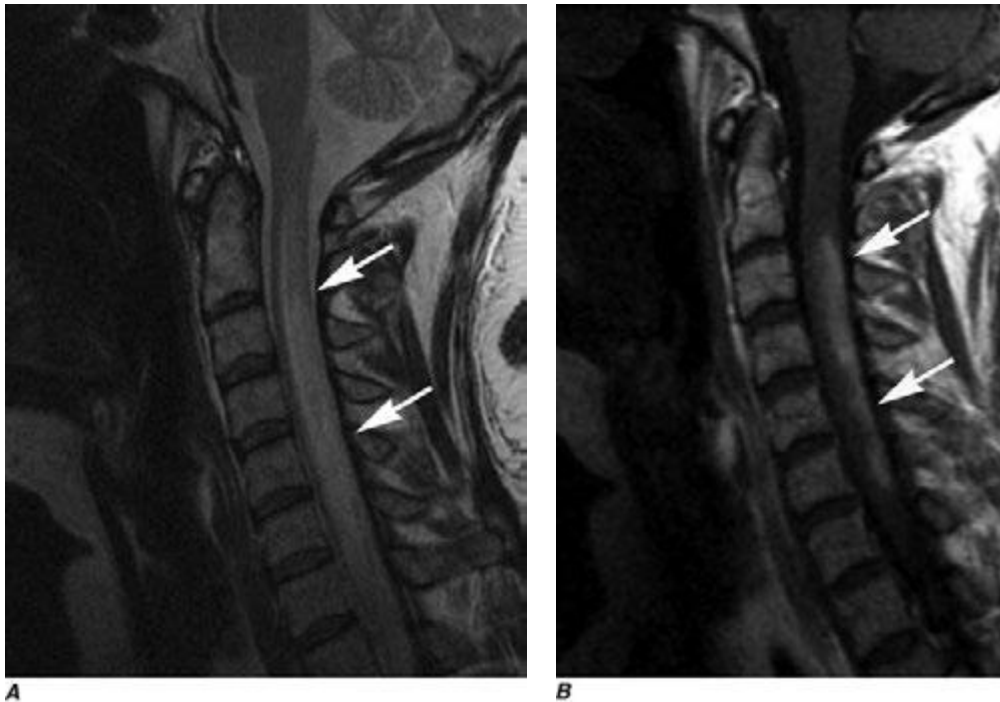
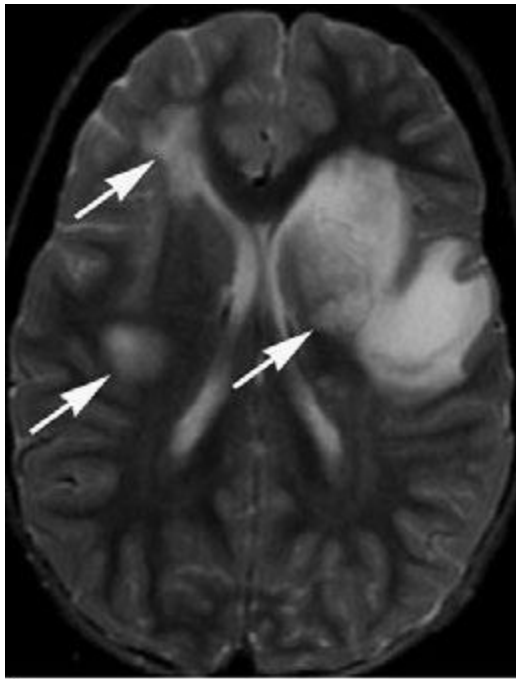
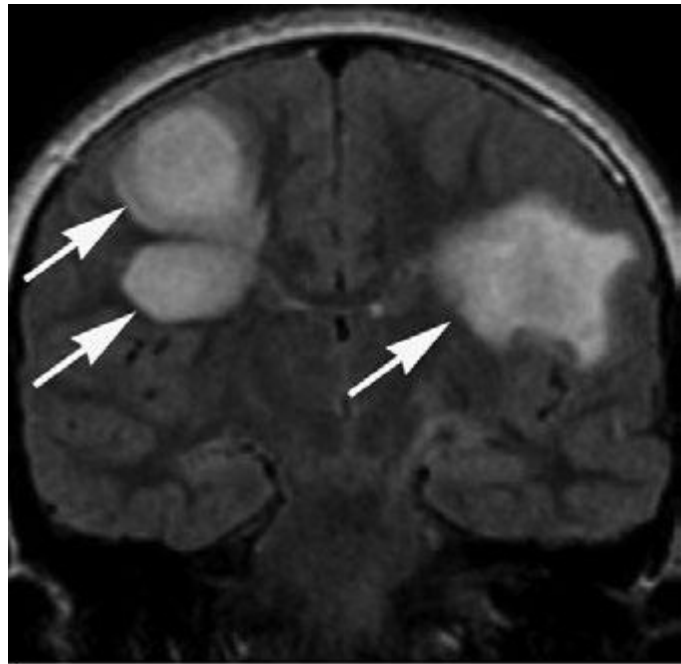


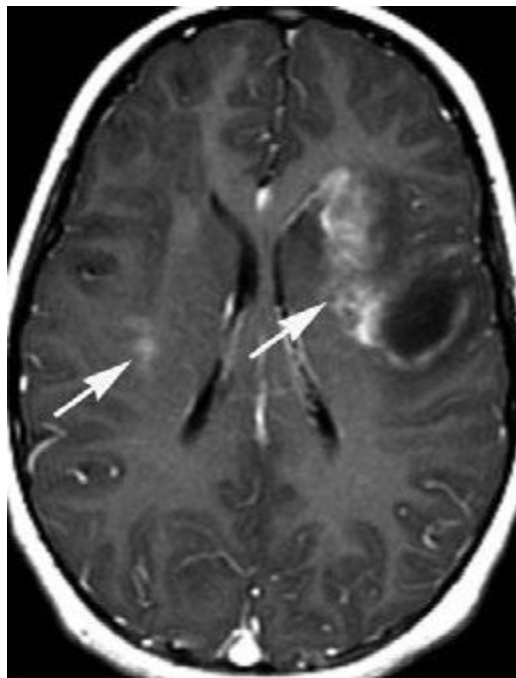
FIGURE 441e-25 Acute transverse myelitis (Chap. 456). Sagittal T2-weighted MRI (**A**) demonstrates abnormal high signal in the cervical cord extending from C1 to T1 with associated cord expansion (*arrows*). Sagittal T1-weighted MRI postgadolinium (**B**) demonstrates abnormal enhancement in the posterior half of the cord from C2 to T1 (*arrows*).



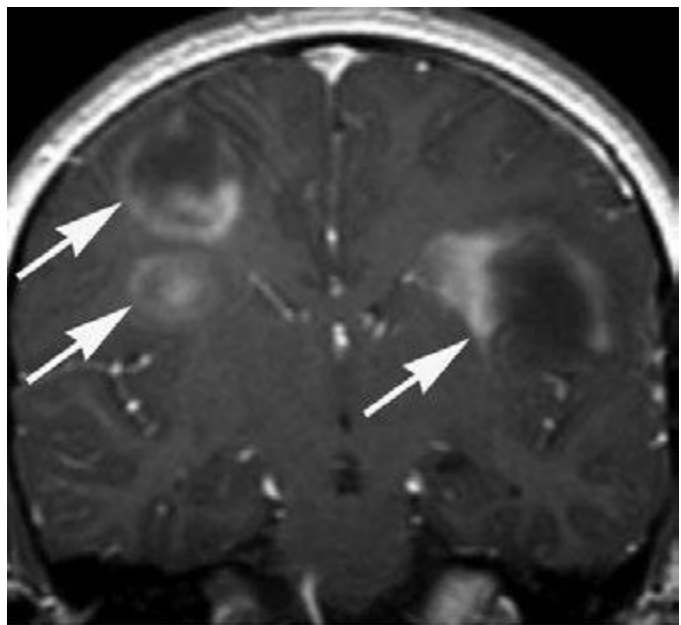
A



B

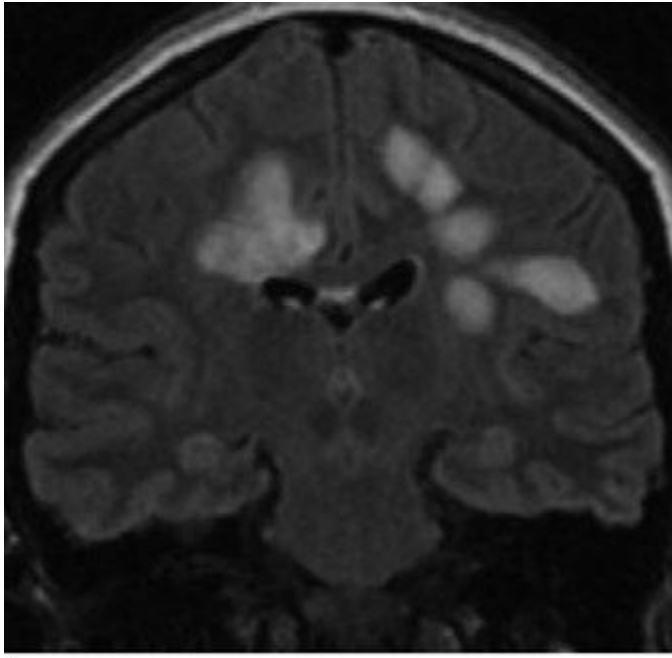


C

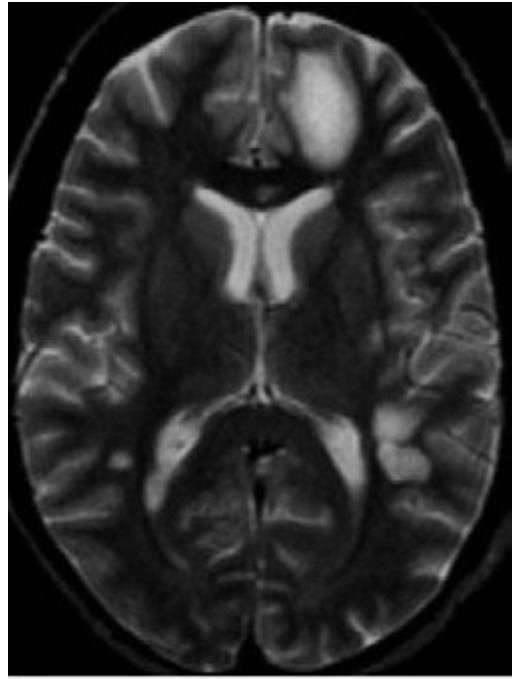


D

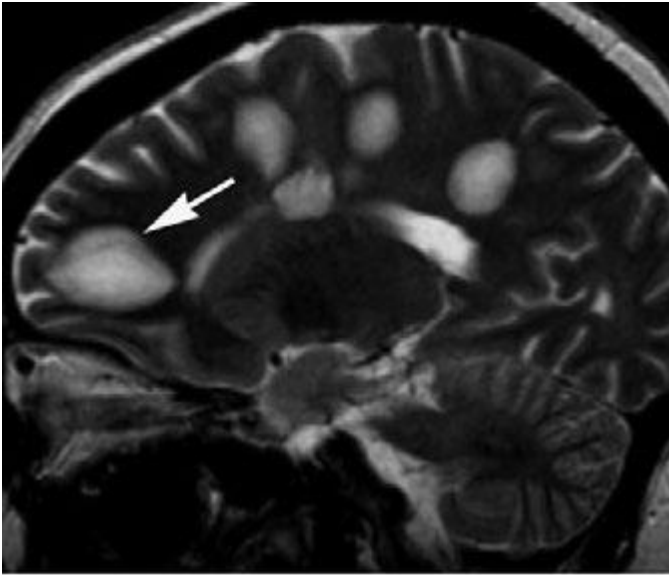
FIGURE 441e-26 Acute disseminated encephalomyelitis (ADEM) (Chap. 458). Axial T2-weighted (A) and coronal FLAIR (B) images demonstrate abnormal areas of high signal involving predominantly the subcortical white matter of the frontal lobe bilaterally and left caudate head. Following administration of gadolinium, corresponding axial (C) and coronal (D) T1-weighted images demonstrate irregular enhancement consistent with blood-brain barrier breakdown and inflammation; some lesions show incomplete rim enhancement, typical for demyelination.



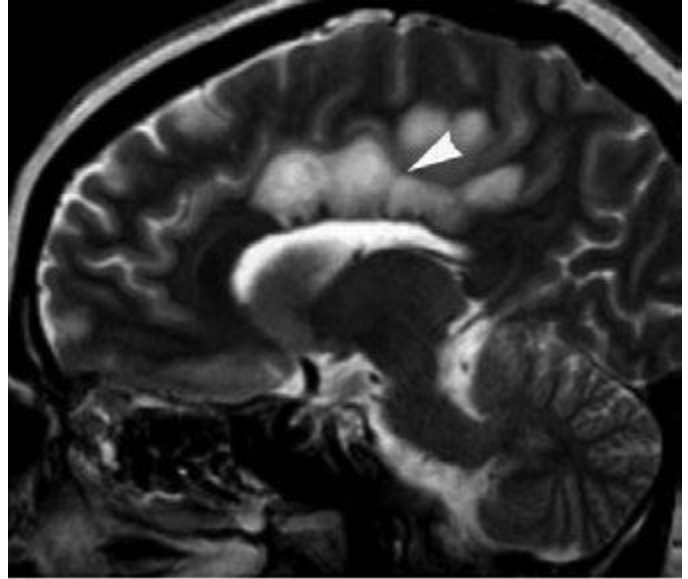
A



B

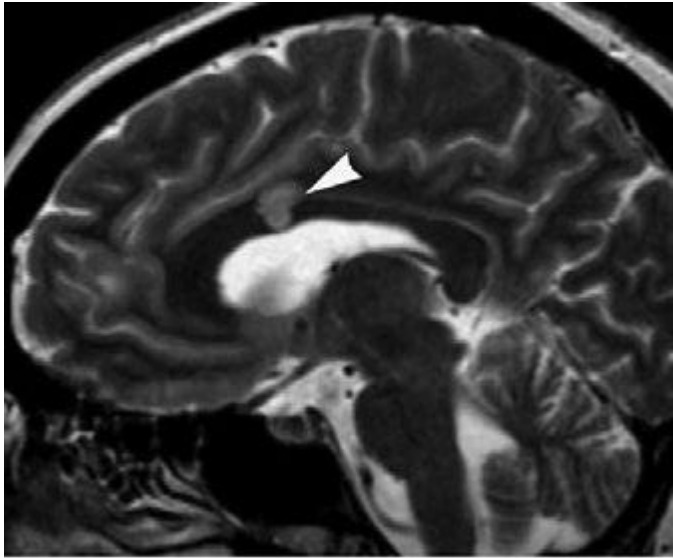


C

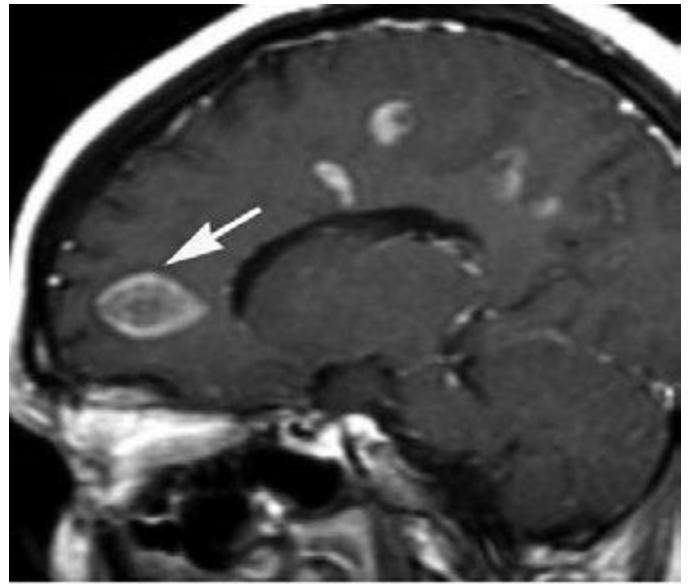


D

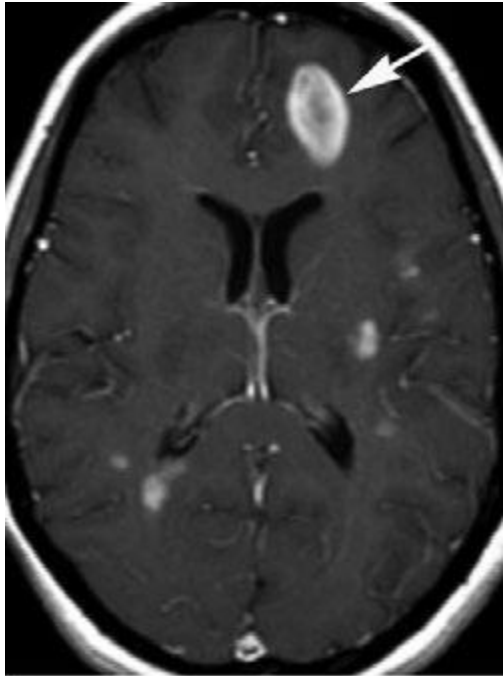
FIGURE 441e-27 Baló's concentric sclerosis (a variant of multiple sclerosis) (Chap. 458). Coronal FLAIR MRI (A) demonstrates multiple areas of abnormal high signal in the supratentorial white matter bilaterally. The lesions are ovoid in shape, perpendicular to the orientation of the lateral ventricles, and with little mass effect. Axial (B) and sagittal (C–E) T2-weighted MRIs demonstrate multiple areas of abnormal high signal in the supratentorial white matter bilaterally, as well as the involvement of the body and splenium of the corpus callosum and the callosal-septal interface (*arrowhead*). Some of the lesions reveal concentric layers, typical of Baló's concentric sclerosis (*arrows*). Sagittal (F) and axial (G, H) T1-weighted MRIs postgadolinium demonstrate abnormal enhancement of all lesions with some of the lesions demonstrating concentric ring enhancement (*arrows*).



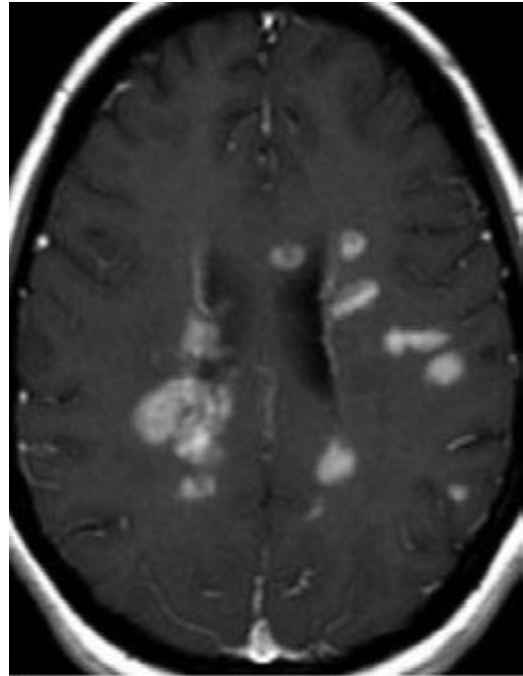
E



F



G



H

FIGURE 441e-27 (Continued)

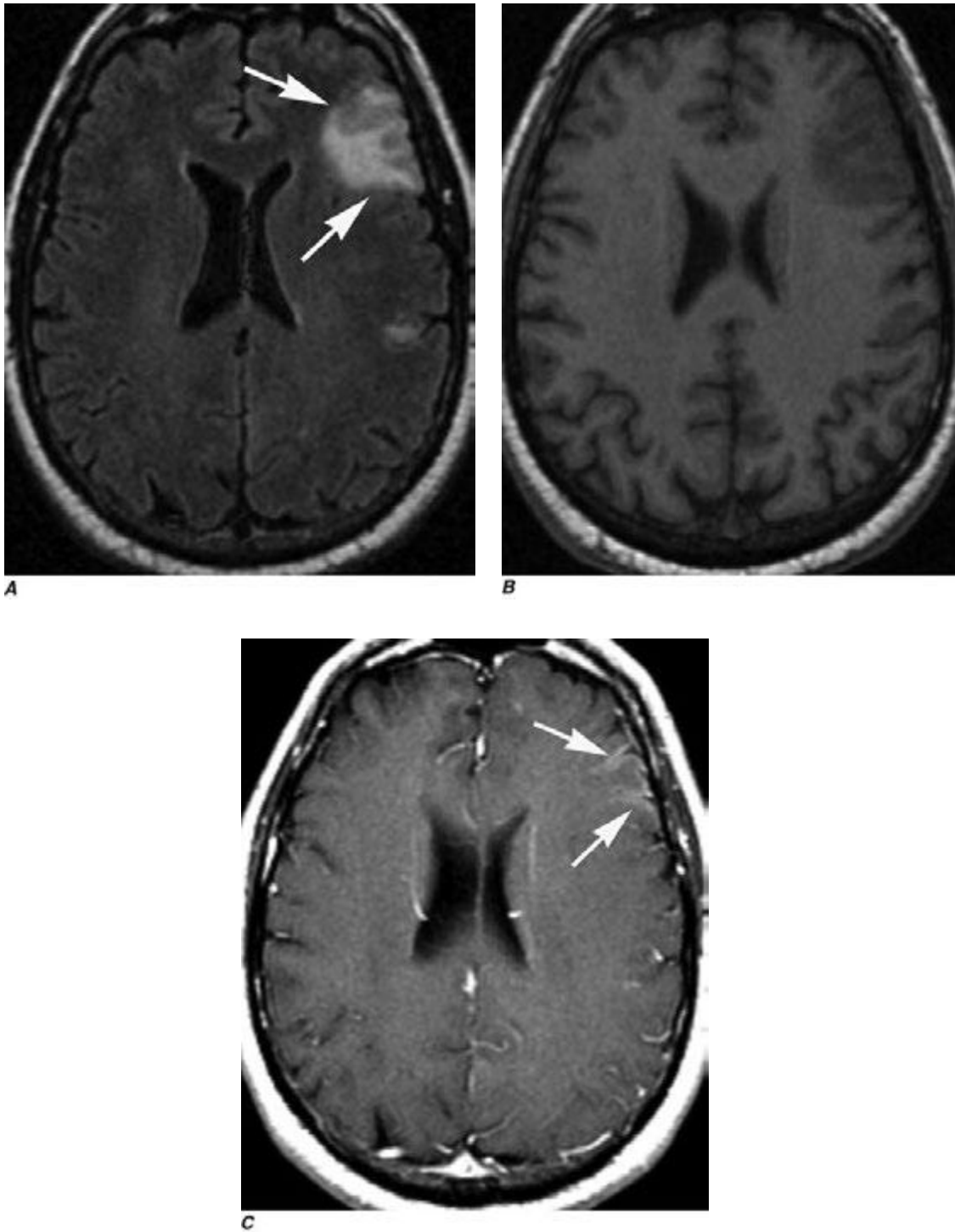
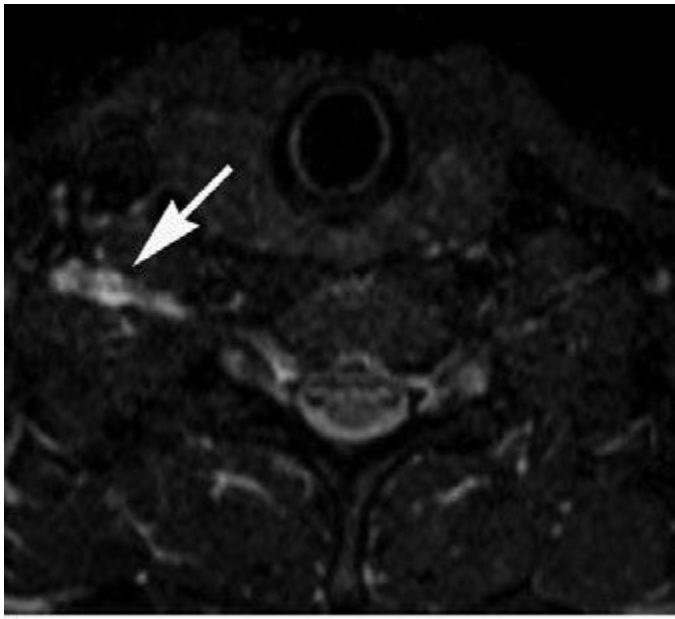
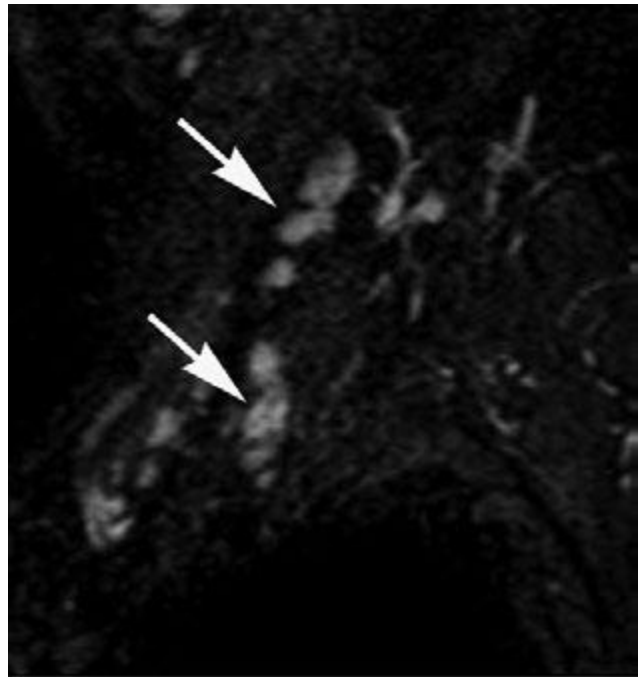


FIGURE 441e-28 Hashimoto's encephalopathy (Chap. 164). Axial FLAIR (A) demonstrates focal area of abnormal high signal involving the gray and white matter in the left frontal lobe. There is also a small area of abnormal high signal in the precentral gyrus. Axial T1-weighted images (B, C) pre and postgadolinium demonstrate cortical/pial enhancement in the region of high signal on FLAIR.



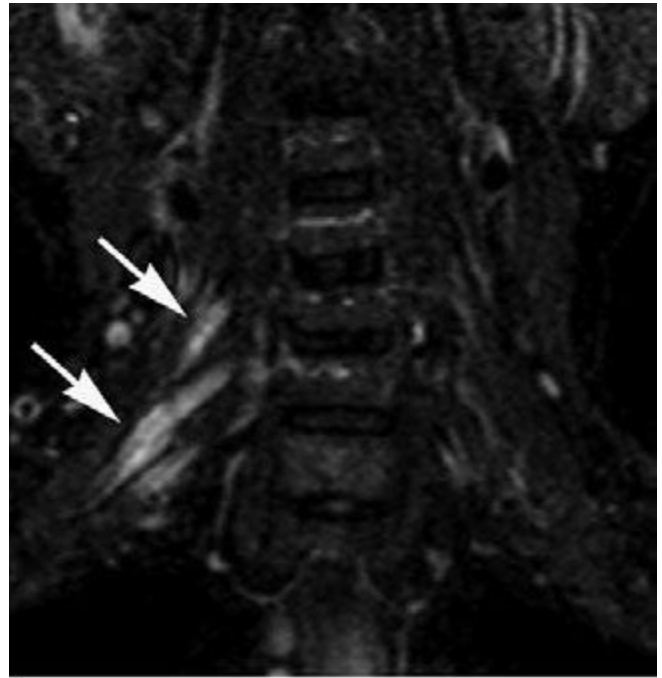
A



B



C



D

FIGURE 441e-29 Brachial plexopathy (Chap. 459). Axial (A), sagittal (B), and coronal (C, D) short tau inversion recovery (STIR) MRIs demonstrate abnormal enlargement and abnormal high signal involving the right C6, C7, and C8 nerve roots, and the trunks and divisions that originate from these roots (arrows). Diffusion-weighted MRI (E) demonstrates abnormal reduced diffusion within the right C6, C7, and C8 nerve roots and their corresponding trunks and divisions (arrow). These findings are compatible with radiation-induced brachial plexopathy.



E
FIGURE 441e-29 (Continued)

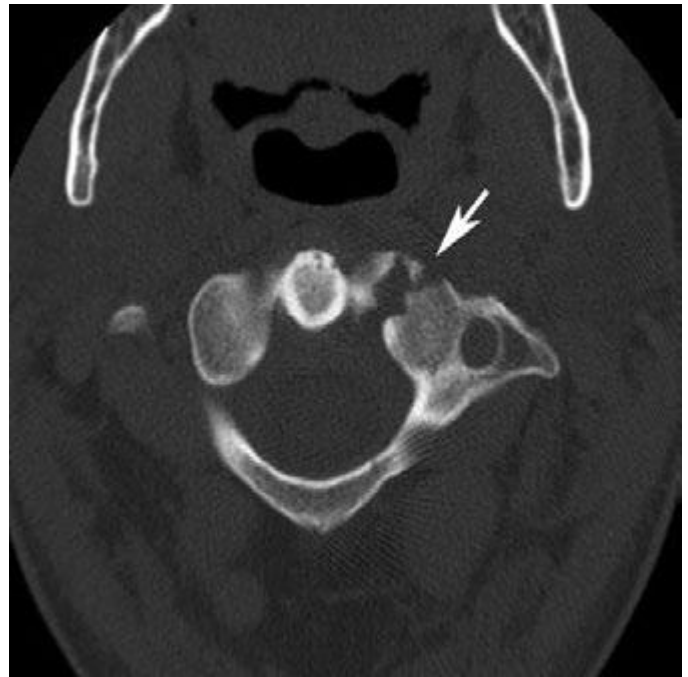


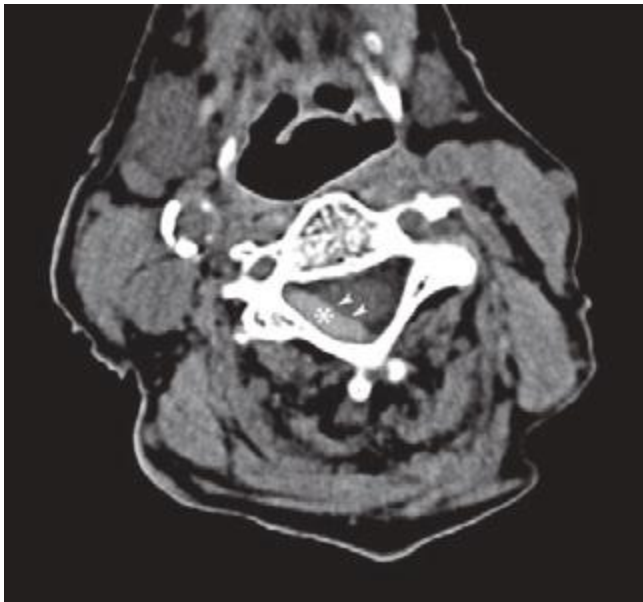
FIGURE 441e-31 **CT facet fracture.** Axial CT demonstrates fracture line along the C2 facet (*arrow*).



FIGURE 441e-30 **Anterior dens dislocation.** Sagittal CT demonstrates the tip of the dens below the anterior arch of C2 (*arrow*), indicating anterior dislocation.



FIGURE 441e-32 **Compression fracture.** Sagittal T2-weighted MRI demonstrates compression fracture of C7 (*) and high signal within the spinous processes of C6-C7 (*arrows*) and to lesser degree C5-C6. This is suggestive of interspinous ligament injury. Note the pad under the patient's neck to maintain neck alignment during the scanning time.



A



B

FIGURE 441e-33 Epidural hematoma. Axial noncontrast CT (**A**) demonstrates a high-density epidural collection in the cervical spine (*), which is consistent with acute hemorrhage. Also noted is mass effect on the spinal cord (*arrowheads*). Sagittal reformatted CT image (**B**) demonstrates the extension of the acute epidural hematoma (*) and a disk bulge (*arrowhead*), which further contributes to spinal canal narrowing. CT is the imaging procedure of choice to detect acute hematoma.



FIGURE 441e-34 Retropharyngeal soft tissue mass. Sagittal T1-weighted MRI demonstrates a hyperflexion fracture with retropulsion of the posterior wall in the canal at C5 and C6 (*arrow*). There is also a large retropharyngeal hematoma (*). The distance from the posterior wall of the airway to the anterior wall of the vertebral body should not measure more than 6 mm at C2 or more than 20 mm at C6 (mnemonic "6 at 2 and 20 at 6").

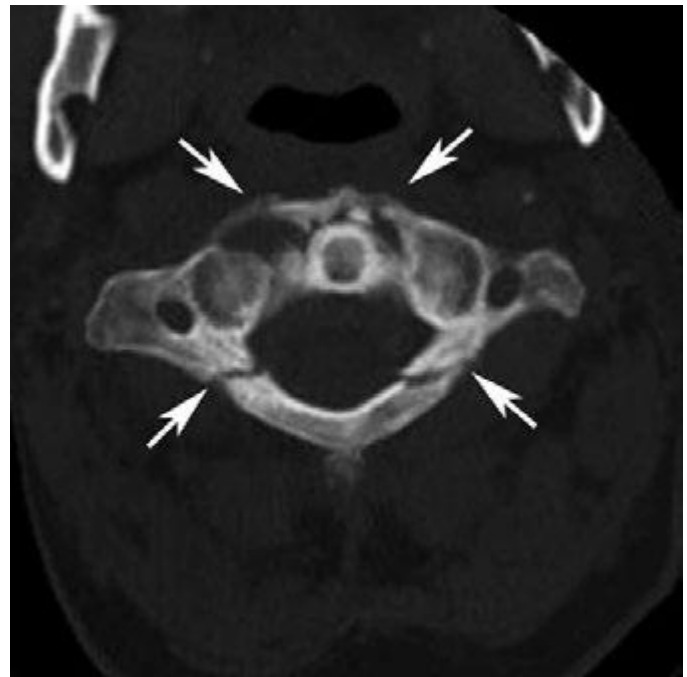


FIGURE 441e-35 Jefferson fracture. Axial CT demonstrates four fracture lines (*arrows*) separating C1 in four parts. Jefferson fracture is usually caused by axial impact to the head such as diving in shallow water.



FIGURE 441e-36 Ligament injury after trauma. Coronal CT reconstruction demonstrates abnormal asymmetry between the dens and the lateral masses of C1 indicating transverse ligament rupture.



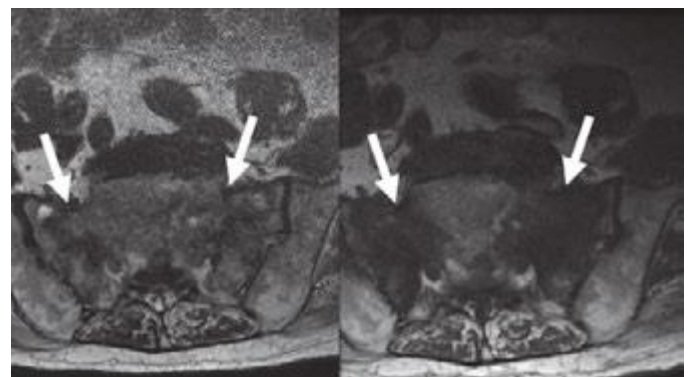
A

B

FIGURE 441e-38 Pathologic fracture. Sagittal T1-weighted MRI (A) demonstrates wedge-shaped T6 vertebral body (arrow). Sagittal postcontrast T1-weighted MRI (B) depicts tumor extension into the epidural space and the involvement of the posterior arch (*), which are highly suggestive of metastatic or primary bone tumor.



FIGURE 441e-37 Odontoid fracture. Sagittal CT demonstrates disruption of the main reference cervical lines. 1: Anterior vertebral body line; 2: Posterior vertebral body line; 3: Spinolaminar line.



A

B

FIGURE 441e-39 Sacral insufficiency fracture. Axial T2-weighted MRI (A) and T1-weighted MRI (B) demonstrate symmetric high T2 and low T1 signal involving the sacral alae longitudinally (arrows).

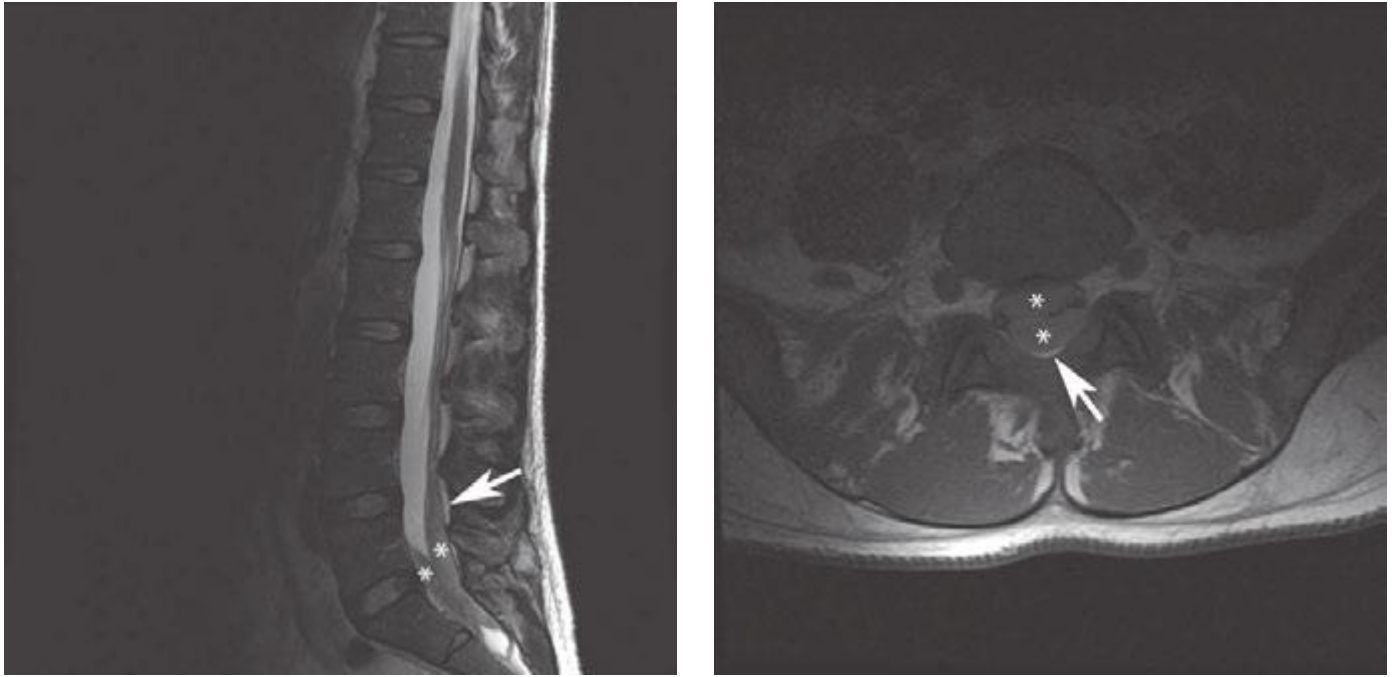


FIGURE 441e-40 Subdural hematoma. Sagittal T2-weighted MRI (**A**) and axial noncontrast T1-weighted MRI (**B**) demonstrate subdural collection in the lumbosacral region (**). Note that the epidural fat is compressed but not involved (*arrow*).

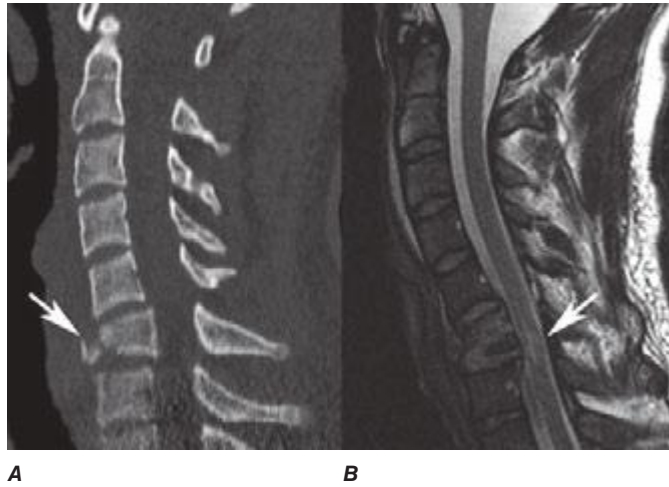


FIGURE 441e-41 Teardrop fracture. Sagittal CT (**A**) demonstrates fracture line separating the anteroinferior corner of C6 (*arrow*). Sagittal T2-weighted MRI (**B**) displays cord injury (*arrow*).

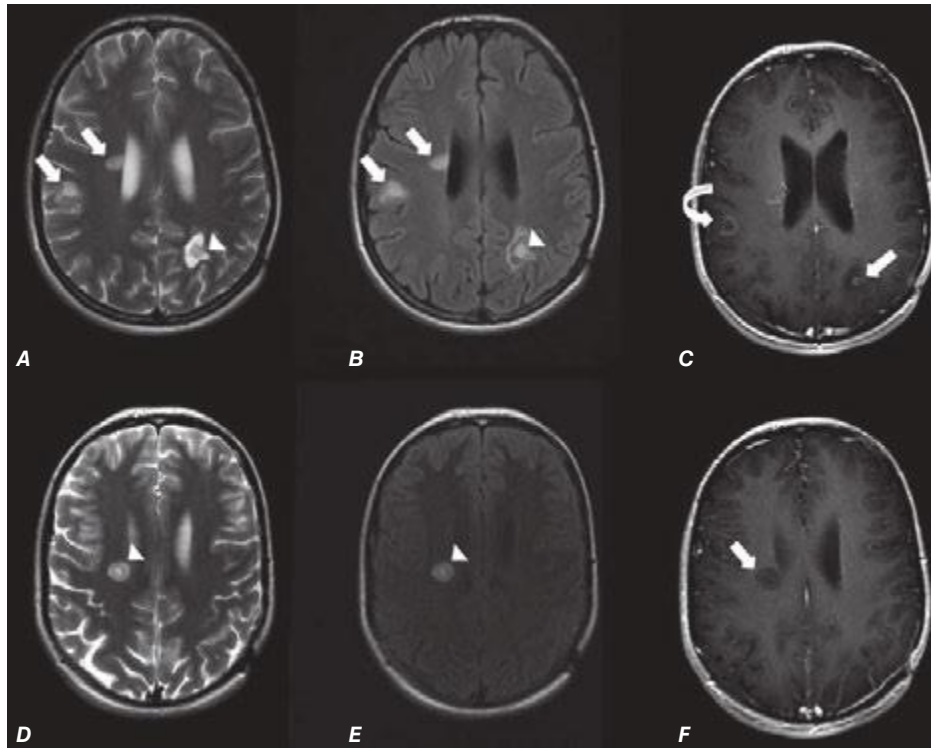


FIGURE 441e-42 Demyelinating disease (multiple sclerosis, Chap. 458). Axial T2-weighted MRI (**A, D**) and axial T2 FLAIR MRI (**B, E**) demonstrate multiple hyperintense lesions involving the periventricular and subcortical white matter (*arrows*). Although not always present, the appearance of a “lesion within and lesion” (*arrowheads*) is typical of demyelinating disease. Axial T1-weighted postcontrast MRI (**C, F**) shows partial enhancement of the lesion (*arrows*), which is often peripheral, incomplete, and “C-shaped” (*curved arrow*).

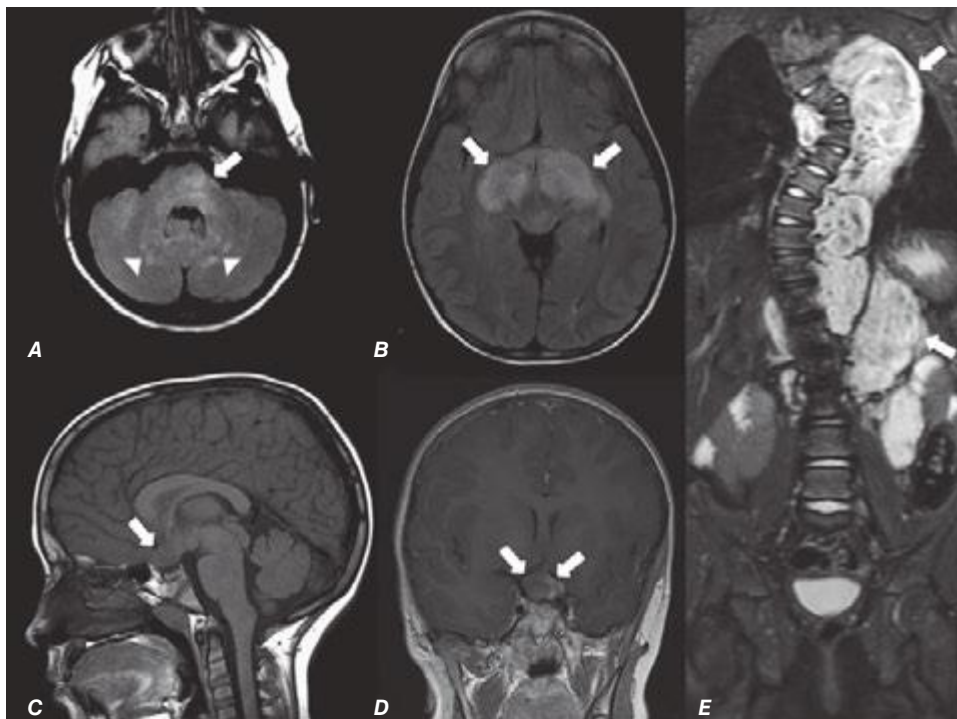


FIGURE 441e-43 Neurofibromatosis type 1 (Chap. 118). Axial T2 FLAIR MRI (**A, B**) demonstrates multiple hyperintense lesions involving the brainstem and basal ganglia (*arrows*) as well as deep cerebellar hemispheres (*arrowheads*). Sagittal and coronal T1-weighted postcontrast MRI (**C, D**) shows enlargement of the optic chiasm with an area of enhancement on the left, representing an optic pathway glioma (*arrows*). Coronal STIR MRI (**E**) shows thoracolumbar scoliosis and a large paravertebral plexiform neurofibroma (*arrows*).

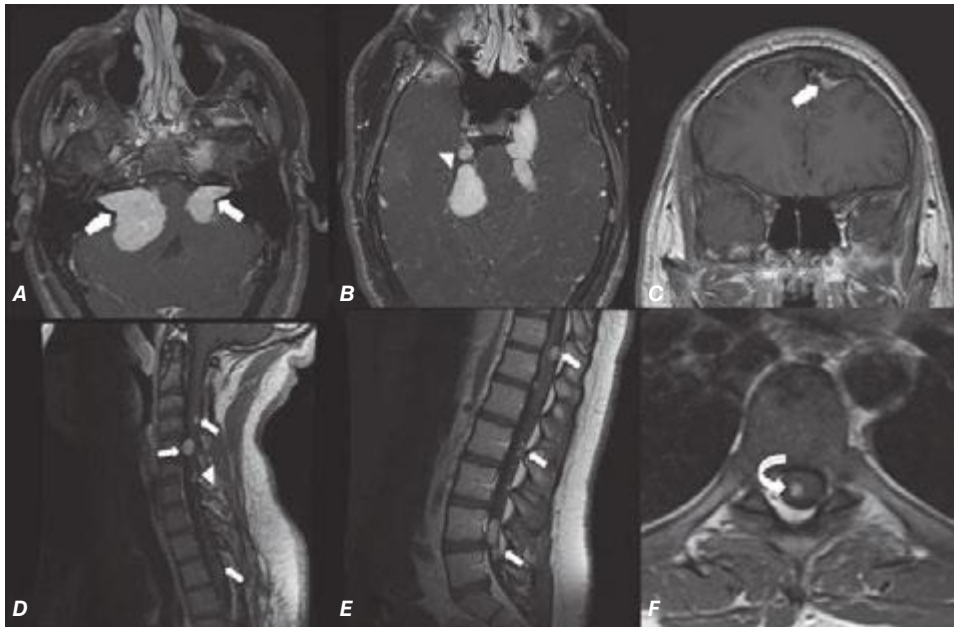


FIGURE 441e-44 Neurofibromatosis type 2 (Chap. 118). Axial T1-weighted postcontrast MRI (**A, B**) shows enhancing expansible lesions in the bilateral cerebellopontine cisterns extending in the internal auditory canals, consistent with vestibular schwannomas (*arrows*), as well as in the bilateral prepontine cistern, consistent with trigeminal schwannomas (*arrowheads*). Coronal axial T1-weighted image postgadolinium (**C**) demonstrates an intensely enhancing dural-based lesion typical for a small meningioma (*arrows*). Sagittal (**D, E**) T1-weighted images postgadolinium show intradural, extramedullary lesions, suggestive of multiple spinal schwannomas (*small arrows*). The flat dural-based lesion may represent a spinal meningioma (*arrowhead*). Axial T1-weighted image postgadolinium (**F**) shows an enhancing intramedullary lesion, most consistent with an ependymoma. (*curved arrow*).

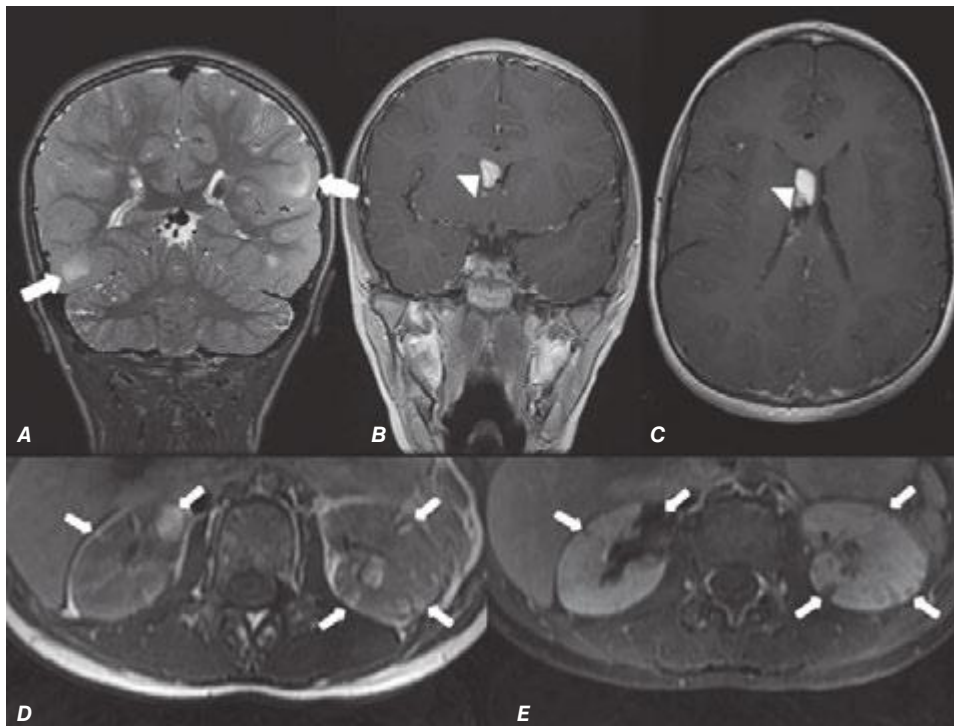


FIGURE 441e-45 Tuberosclerosis (Chap. 118). Coronal T2-weighted MRI (**A**) shows multiple T2 hyperintense lesions in a cortical and subcortical distribution (*arrows*). Coronal and axial postcontrast T1-weighted image (**B, C**) demonstrates an expanding nodule with intense enhancement in the proximity of the right foramen of Monro, consistent with a subependymal giant-cell astrocytoma (SEGA) (*arrowheads*). Surveillance T1-weighted image (**D**) and postcontrast T1-weighted image with fat saturation (**E**) show multiple bilateral renal lesions with signal intensity of fat, consistent with angiomyolipomas (*small arrows*).

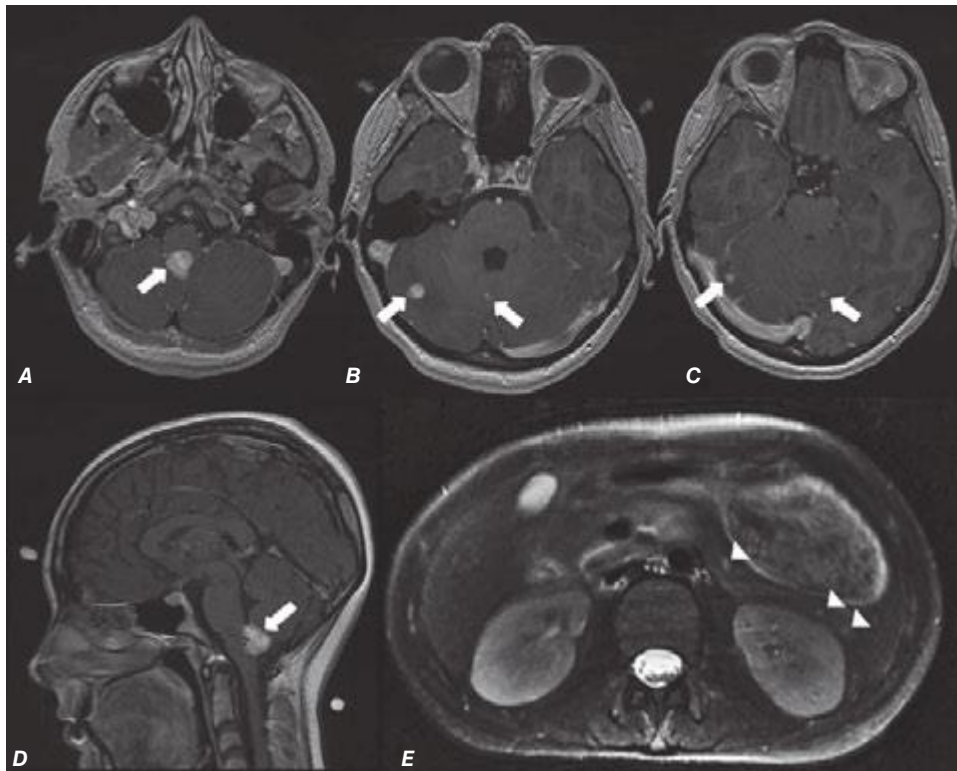


FIGURE 441e-46 Von Hippel-Lindau (VHL) (Chap. 408). Axial postcontrast T1 weighted images (A–C) demonstrates multiple enhancing nodules in the posterior fossa (arrows). Sagittal postcontrast T1-weighted image (D) shows vascular flow voids within the enhancing nodule in the region of the foramen of Magendie (arrow), indicating increased vascularity. Surveillance axial T2-weighted MRI of the abdomen (E) shows multiple small pancreatic cysts (arrowheads). This patient did not have an endolymphatic sac tumor, renal cell carcinoma, neuroendocrine pancreatic tumor, or pheochromocytoma, all of which may also occur in von Hippel-Lindau disease.

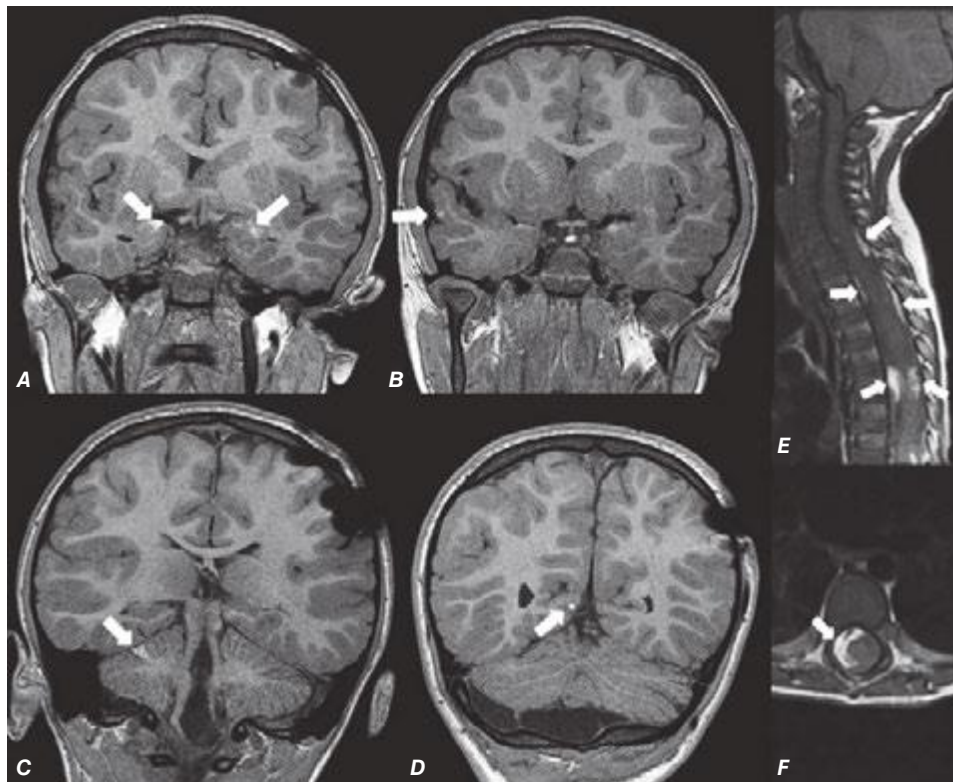


FIGURE 441e-47 Neurocutaneous melanosis. Coronal T1 weighted MRI (A–D) shows multiple lesions with intrinsic increased T1 signal in the bilateral amygdalae, right superior temporal gyrus, right cerebellar hemisphere, and right medial occipital cortex (arrows). Sagittal and axial T1-weighted images of the spine (E, F) show intradural, extramedullary lesions, also with intrinsic increased T1 signal, due to malignant melanoma (arrows).

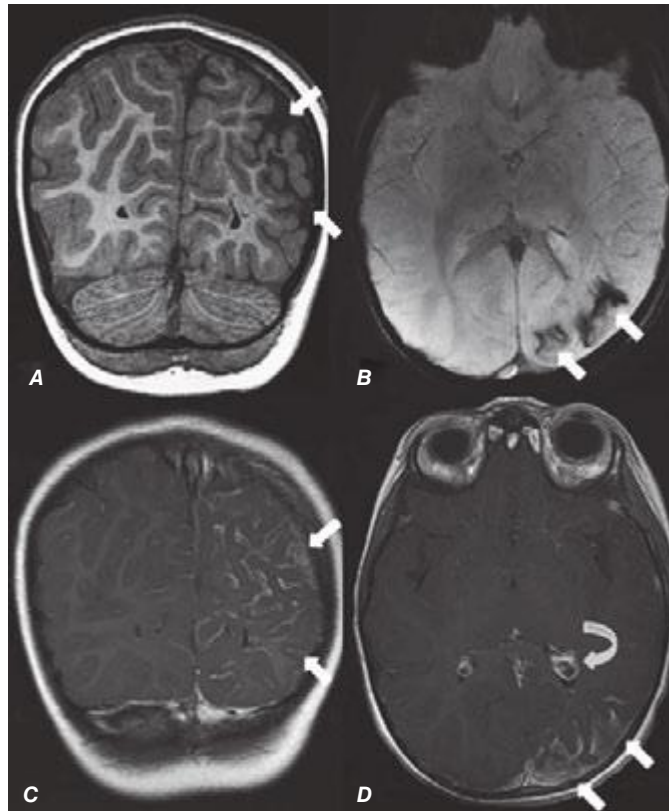


FIGURE 441e-48 Sturge-Weber syndrome. Coronal T1-weighted MRI (A) shows enlargement of the sulci in the left parietal lobe, consistent with brain parenchymal volume loss (arrows). Axial susceptibility weighted imaging shows susceptibility effect in this region, consistent with calcifications (arrows). Coronal and axial T1-weighted images postgadolinium (C, D) show increased leptomeningeal enhancement (arrows) and enlargement of the left choroid plexus (curved arrow).

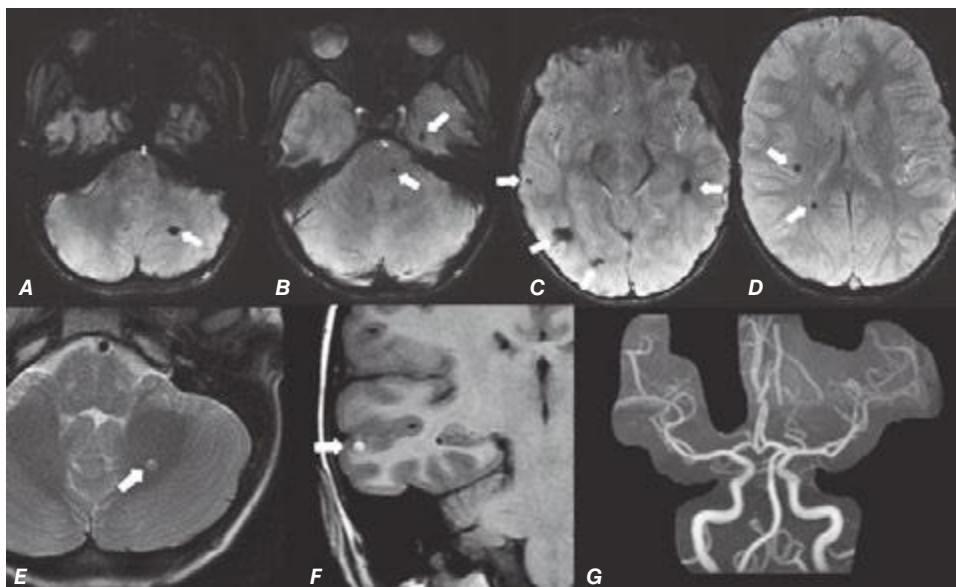
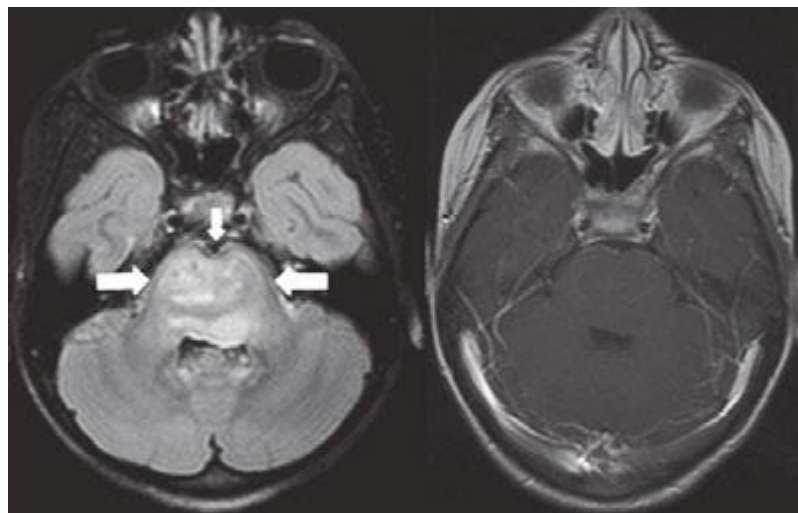
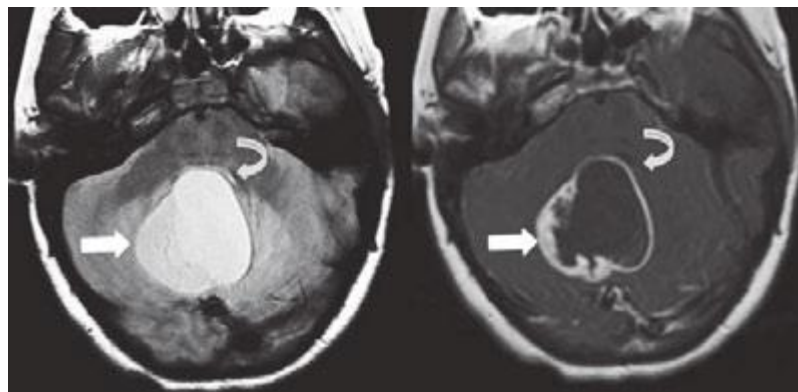


FIGURE 441e-49 Multiple cavernomas (Chap. 446). Axial susceptibility weighted images (A–D) show multiple foci of susceptibility involving the bilateral cerebral hemispheres, pons, and left cerebellum, which in a young patient most likely represents multiple cavernomas (arrows). These lesions have variable signal intensity on T2-weighted (E) and T1-weighted MRI (F), related to the different stages of hemoglobin degradation (arrows). Cavernomas are not seen on the time-of-flight MR angiography (G); thus these are designated as angiographically occult vascular malformations. Of note, in elderly patients, amyloid angiopathy may have a similar appearance.



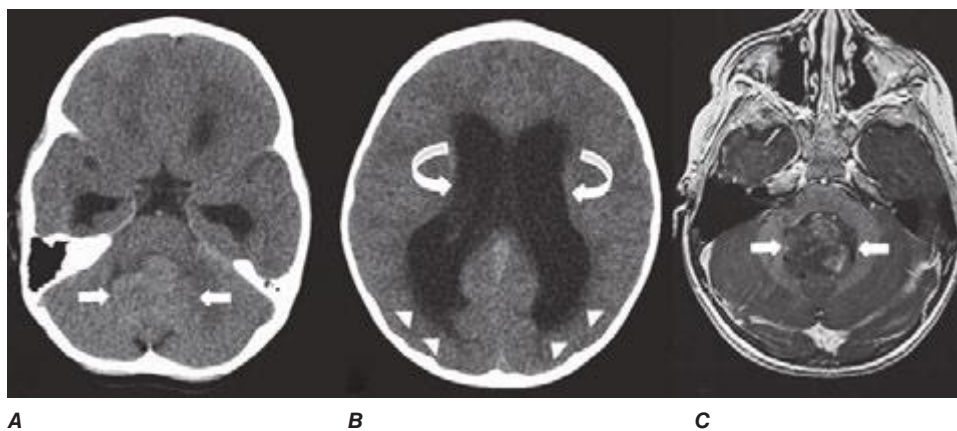
A **B**

FIGURE 441e-50 Brainstem glioma (Chap. 118). Axial T2 FLAIR MRI (**A**) demonstrates increased T2 signal and marked enlargement of the pons (*large arrows*), which engulfs the basilar artery (*small arrow*). These findings are characteristic of brainstem glioma. At the time of diagnosis, these lesions are typically low grade without abnormal enhancement, as shown by axial postcontrast T1-weighted image (**B**).



A **B**

FIGURE 441e-51 Pilocytic astrocytoma (Chap. 118). Axial T2-weighted and T1-weighted postgadolinium images (**A, B**) show a cystic lesion with peripheral enhancement and an enhancing solid component located in the posterior fossa (*arrows*). These findings are suggestive of a pilocytic astrocytoma. Note that the lesion exerts mass effect on the fourth ventricle, which is compressed (*curved arrows*).



A **B** **C**

FIGURE 441e-52 Fourth ventricle ependymoma causing hydrocephalus (Chap. 118). Unenhanced axial CT images (**A, B**) demonstrate an expansile mass lesion, isodense with brain parenchyma, filling the fourth ventricle (*arrows*) and causing obstruction of cerebrospinal fluid flow, dilation of the lateral ventricles, and hydrocephalus (*curved arrows*). The hypoattenuation within periventricular white matter bilaterally represents transependymal flow (*arrowheads*). Axial T1-weighted postgadolinium MRI confirms the presence of a heterogeneously enhancing mass filling the fourth ventricle (*arrow*), which is suggestive of ependymoma.

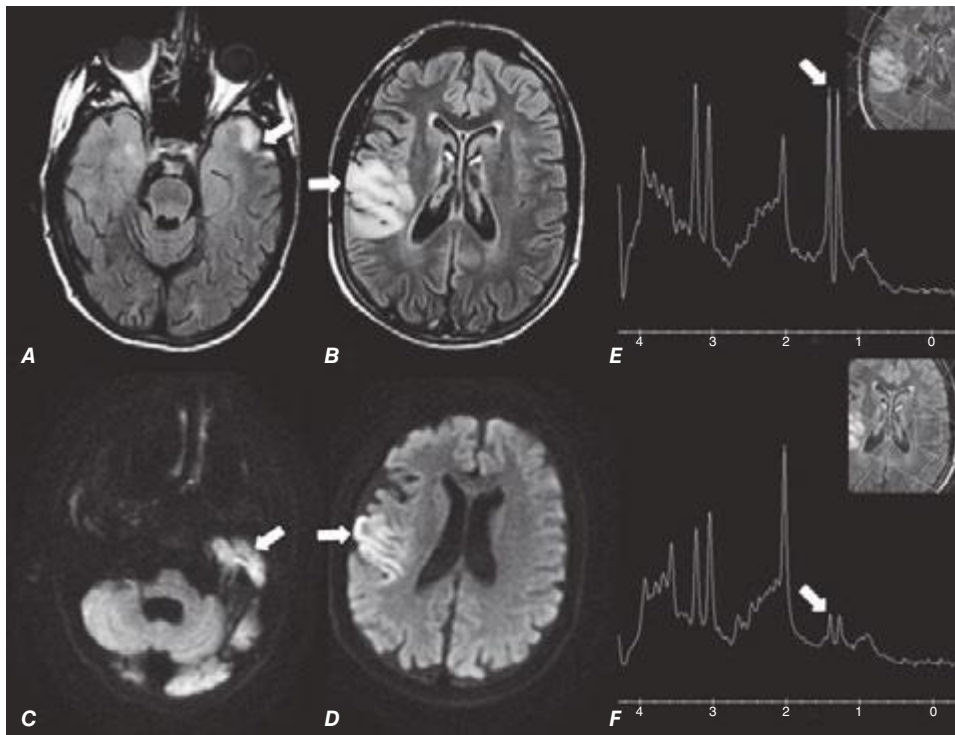


FIGURE 441e-53 Mitochondrial encephalopathy, lactic acidosis, and stroke-like episodes (MELAS) (Chap. 462e). Axial T2 FLAIR MRI (A, B) shows areas of increased T2 signal involving the cortex and subcortical white matter in the posterior right frontal lobe and anterior left temporal lobe, consistent with edema (arrows). Axial diffusion-weighted images (C, D) show reduced diffusion from cytotoxic edema, consistent with infarcts (arrows). MR spectroscopy of the right frontal lesion (E) demonstrates markedly increased lactate (arrow), an expected finding in infarction regardless of etiology. However, MR spectroscopy of the normal-appearing contralateral brain parenchyma (F) shows mildly elevated lactate (arrow), which is suggestive of a mitochondrial disorder.

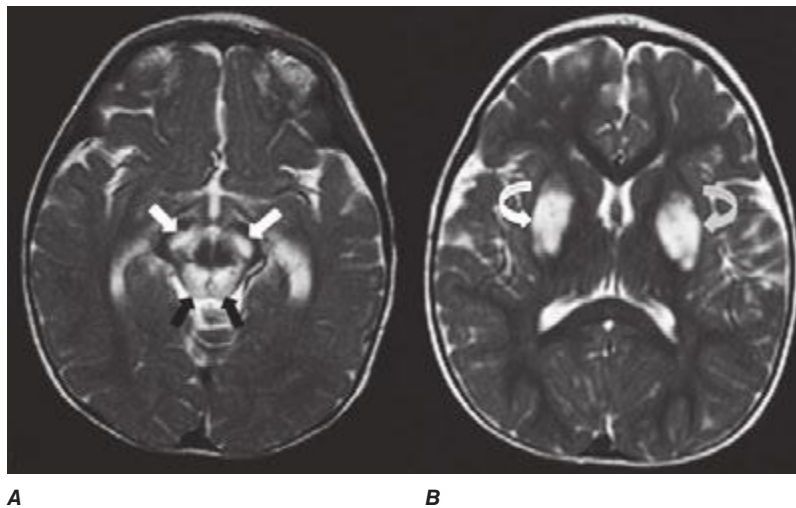


FIGURE 441e-54 Leigh's disease (subacute necrotizing encephalomyelopathy) (Chap. 85e). Axial T2-weighted MRI (A, B) demonstrates increased T2 signal involving the substantia nigra (white arrows) and dorsal midbrain (black arrows), as well as the putamen bilaterally (curved arrows). This is a common pattern for the mitochondrial disorder Leigh's disease secondary to cytochrome oxidase (CO IV) deficiency.

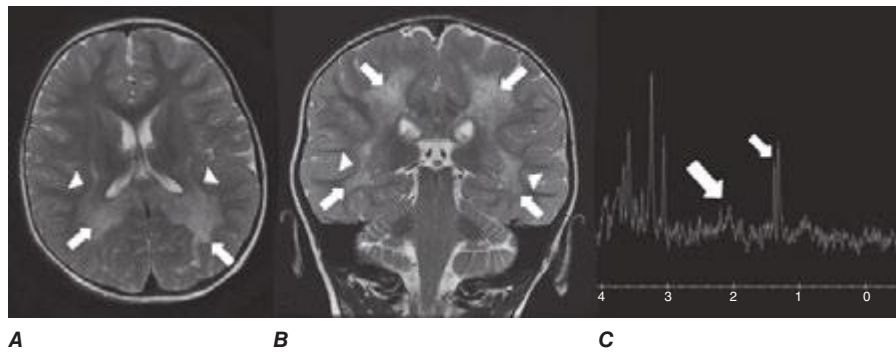


FIGURE 441e-55 Krabbe's disease (Chap. 432e). Axial and coronal T2-weighted MRIs (A, B) show increased T2 signal involving predominantly the posterior white matter bilaterally (arrows) with sparing of the subcortical U-fibers (arrowheads). MR spectroscopy of the left parietal white matter (C) shows markedly decreased N-acetylaspartate (large arrow) and increased lactate/lipids (small arrow), consistent with severe neuronal injury.

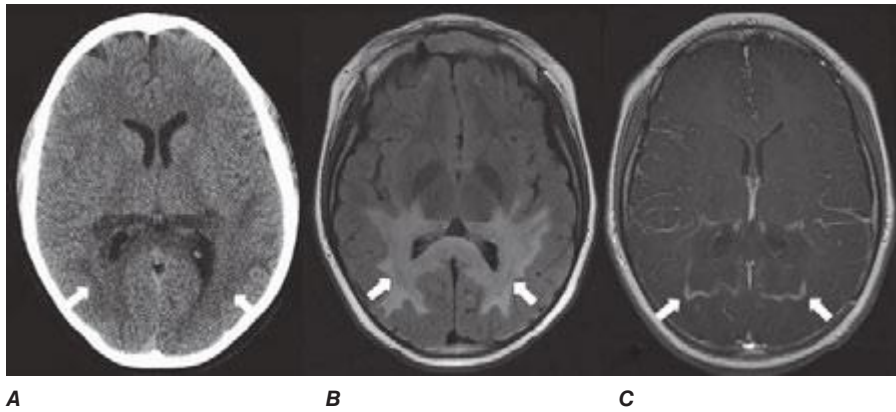


FIGURE 441e-56 X-linked adrenoleukodystrophy (Chap. 459). Axial unenhanced CT (A) demonstrates areas of attenuation involving the posterior white matter bilaterally (arrows). Axial T2 FLAIR MRI (B) displays increased T2 signal consistent with edema (arrows). Axial T1-weighted image postgadolinium (C) shows peripheral enhancement of the parietal lesions bilaterally (arrows). These findings are typical of adrenoleukodystrophy.

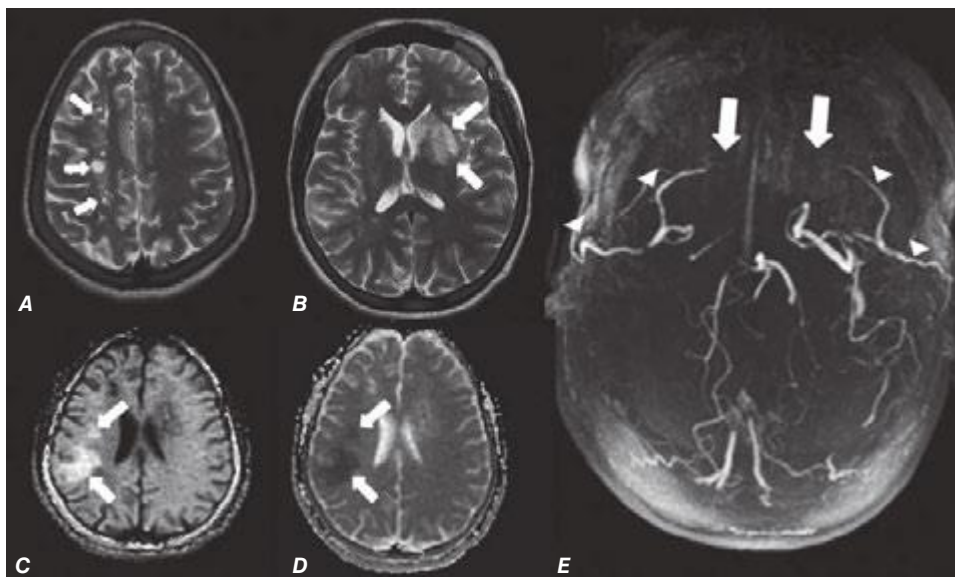


FIGURE 441e-57 Sickle cell disease and moyamoya disease (Chap. 446). Axial T2-weighted MRI (A, B) shows multiple small areas of encephalomalacia from prior infarcts in the watershed zones between the anterior and middle cerebral artery territories (small arrows). There is also an area of edema involving the left basal ganglia from an evolving subacute infarct (arrow). An axial diffusion-weighted image (C) with corresponding ADC map (D) shows an area of restricted diffusion in the right frontoparietal region, consistent with an acute infarct (arrows). Time-of-flight MR angiography (E) shows absence of flow in the distal internal carotid arteries and proximal middle cerebral arteries (arrows) due to moyamoya disease. Also note that this patient is status post-bilateral encephalo-duro-arterio-synangiosis (EDAS) (arrowheads), a surgical procedure to create indirect anastomosis between branches of the external carotid artery with distal branches of the middle cerebral artery.

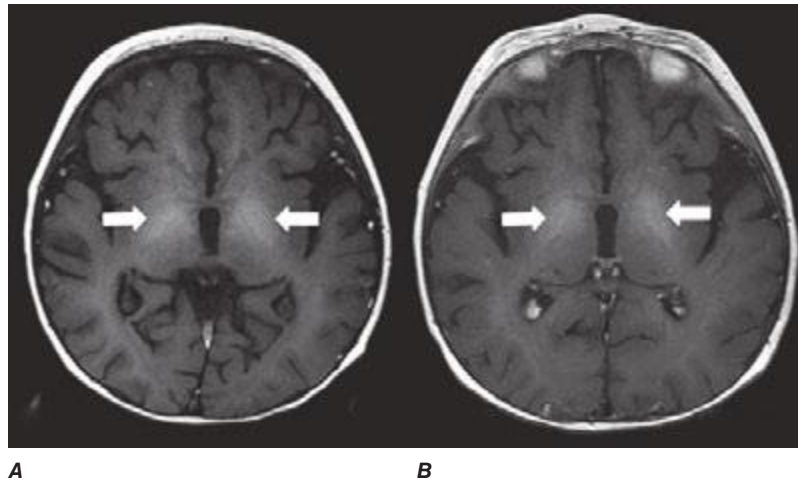


FIGURE 441e-58 Hepatic encephalopathy (Chap. 330). Axial T1-weighted MRI (A, B) shows increased intrinsic T1 signal involving the basal ganglia bilaterally, particularly the globus pallidus (arrows).



FIGURE 441e-59 Guillain-Barré syndrome (Chap. 460). Axial precontrast T1-weighted MRI (A) and axial and sagittal T1-weighted postgadolinium (B, C) images show thickening and increased enhancement of the anterior roots of the cauda equina (arrows).

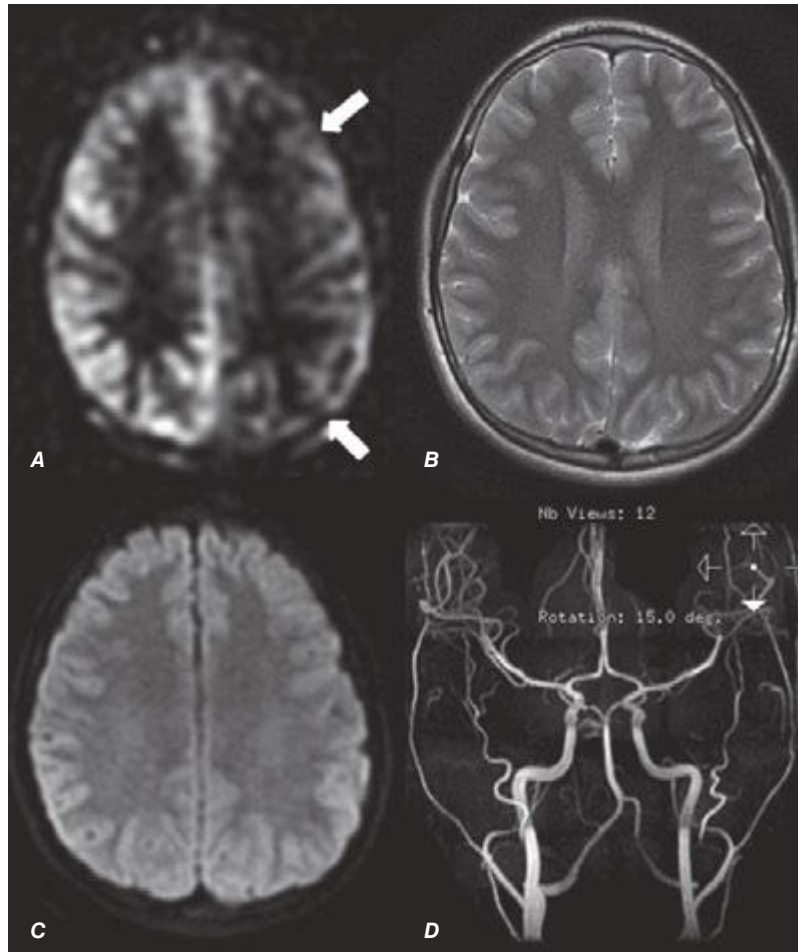


FIGURE 441e-60 Hemiplegic migraine (Chap. 447). Axial noncontrast perfusion MRI using an arterial spin labeling technique (A) demonstrates decreased cerebral blood flow to the left hemisphere (arrows) in a patient with right hemiparesis associated with migraine symptoms. No abnormality was seen on T2-weighted images, diffusion-weighted images, or time-of-flight MR angiography (B–D) to suggest stroke.

442e Electrodiagnostic Studies of Nervous System Disorders: EEG, Evoked Potentials, and EMG

Michael J. Aminoff

ELECTROENCEPHALOGRAPHY

The electrical activity of the brain (the electroencephalogram [EEG]) is easily recorded from electrodes placed on the scalp. The potential difference between pairs of electrodes on the scalp (bipolar derivation) or between individual scalp electrodes and a relatively inactive common reference point (referential derivation) is amplified and displayed on a computer monitor, oscilloscope, or paper. Digital systems allow the EEG to be reconstructed and displayed with any desired format and to be manipulated for more detailed analysis and also permit computerized techniques to be used to detect certain abnormalities. The characteristics of the normal EEG depend on the patient's age and level of arousal. The rhythmic activity normally recorded represents the postsynaptic potentials of vertically oriented pyramidal cells of the cerebral cortex and is characterized by its frequency. In normal awake adults lying quietly with the eyes closed, an 8- to 13-Hz alpha rhythm is seen posteriorly in the EEG, intermixed with a variable amount of generalized faster (beta) activity (>13 Hz); the alpha rhythm is attenuated when the eyes are opened (Fig. 442e-1). During drowsiness, the alpha rhythm is also attenuated; with light sleep, slower activity in the theta (4–7 Hz) and delta (<4 Hz) ranges becomes more conspicuous.

Activating procedures are generally undertaken while the EEG is recorded in an attempt to provoke abnormalities. Such procedures commonly include hyperventilation (for 3 or 4 min), photic stimulation, sleep, and sleep deprivation on the night prior to the recording.

Electroencephalography is relatively inexpensive and may aid clinical management in several different contexts.

THE EEG AND EPILEPSY

The EEG is most useful in evaluating patients with suspected epilepsy. The presence of electrographic seizure activity—i.e., of abnormal, repetitive, rhythmic activity having an abrupt onset and termination and a characteristic evolution—clearly establishes the diagnosis. The absence of such electrocerebral accompaniment to an episodic behavioral disturbance does not exclude a seizure disorder, however, because there may be no changes in the scalp-recorded EEG during certain focal seizures. With generalized tonic-clonic seizures, the EEG is always abnormal during the episode. It is often not possible to obtain an EEG during clinical events that may represent seizures, especially when such events occur unpredictably or infrequently. Continuous monitoring for prolonged periods in video-EEG telemetry units has made it easier to capture the electrocerebral accompaniments of such clinical episodes. Monitoring by these means is sometimes helpful in confirming that seizures are occurring, characterizing the nature of clinically equivocal episodes, and determining the frequency of epileptic events.

The EEG findings in the interictal period may show certain abnormalities that are strongly supportive of a diagnosis of epilepsy. Such *epileptiform activity* consists of bursts of abnormal discharges containing spikes or sharp waves. The presence of epileptiform activity is not specific for epilepsy, but it has a much greater prevalence in epileptic patients than in normal individuals. However, even in individuals with epilepsy, the initial routine interictal EEG may be normal up to 60% of the time. Thus, the EEG cannot establish the diagnosis of epilepsy in many cases.

The EEG findings have been used in classifying seizure disorders and selecting appropriate anticonvulsant medication for individual patients (Fig. 442e-2). The episodic generalized spike-wave activity that occurs during and between seizures in patients with typical absence epilepsy contrasts with focal interictal epileptiform discharges or ictal patterns found in patients with focal seizures. These latter seizures may have no correlates in the scalp-recorded EEG or may be associated with abnormal rhythmic activity of variable frequency, a localized or generalized distribution, and a stereotyped pattern that varies with the patient. Focal or lateralized epileptogenic lesions are important to recognize, especially if surgical treatment is contemplated. Intensive long-term monitoring of clinical behavior and the EEG is required for operative candidates, however, and this generally also involves recording from intracranial (subdural, extradural, or intracerebral) electrodes.

The EEG findings may indicate the prognosis of seizure disorders: In general, a normal EEG implies a better prognosis than otherwise, whereas an abnormal background or profuse epileptiform activity suggests a poor outlook. The EEG findings are not helpful in determining which patients with head injuries, stroke, or brain tumors will go on to develop seizures, because in such circumstances epileptiform activity is

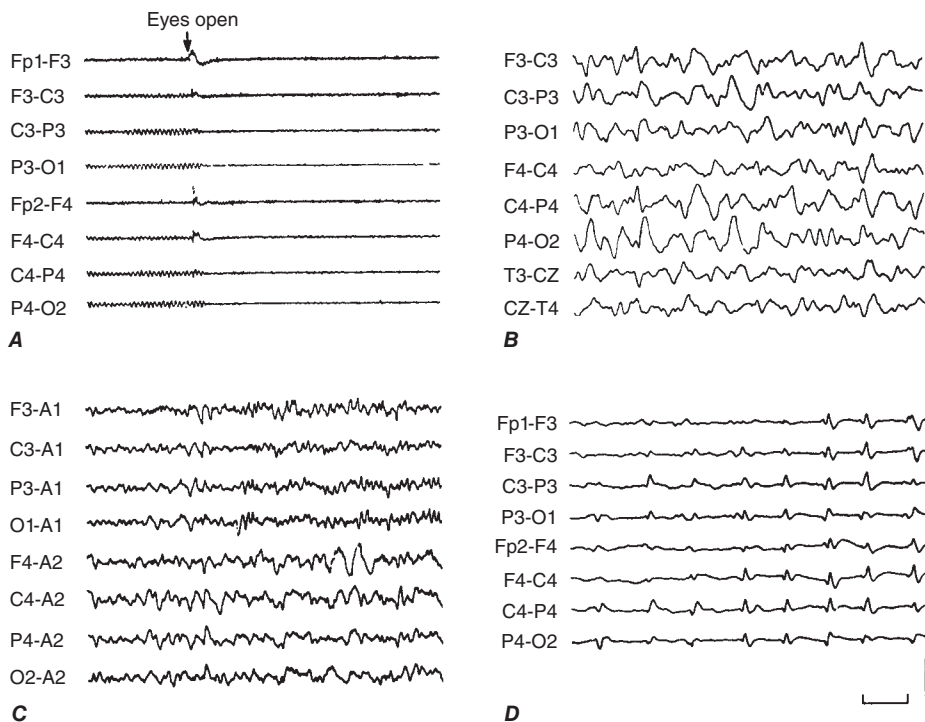


FIGURE 442e-1 **A.** Normal electroencephalogram (EEG) showing a posteriorly situated 9-Hz alpha rhythm that attenuates with eye opening. **B.** Abnormal EEG showing irregular diffuse slow activity in an obtunded patient with encephalitis. **C.** Irregular slow activity in the right central region, on a diffusely slowed background, in a patient with a right parietal glioma. **D.** Periodic complexes occurring once every second in a patient with Creutzfeldt-Jakob disease. Horizontal calibration: 1 s; vertical calibration: 200 μ V in A, 300 μ V in other panels. In this and the following figure, electrode placements are indicated at the left of each panel and accord with the international 10:20 system. A, earlobe; C, central; F, frontal; Fp, frontal polar; P, parietal; T, temporal; O, occipital. Right-sided placements are indicated by even numbers, left-sided placements by odd numbers, and midline placements by Z. (From MJ Aminoff [ed]: *Aminoff's Electrodiagnosis in Clinical Neurology*, 6th ed. Oxford, Elsevier Saunders, 2012.)

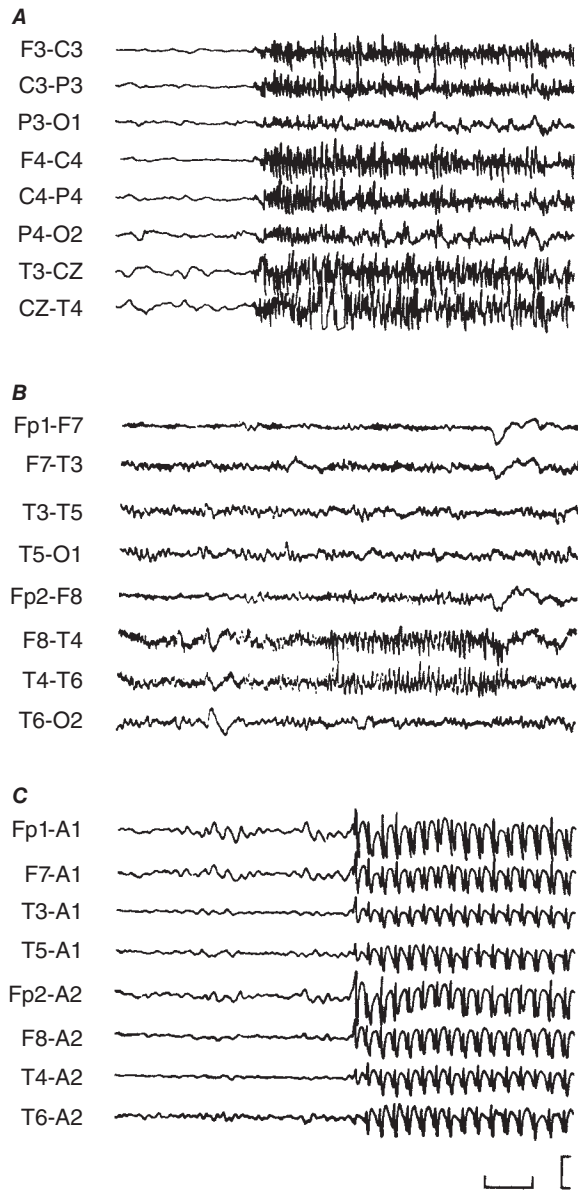


FIGURE 442e-2 Electrographic seizures. **A.** Onset of a tonic seizure showing generalized repetitive sharp activity with synchronous onset over both hemispheres. **B.** Burst of repetitive spikes occurring with sudden onset in the right temporal region during a clinical spell characterized by transient impairment of external awareness. **C.** Generalized 3-Hz spike-wave activity occurring synchronously over both hemispheres during an absence (petit mal) attack. Horizontal calibration: 1 s; vertical calibration: 400 μ V in A, 200 μ V in B, and 750 μ V in C. (From MJ Aminoff [ed]: *Aminoff's Electrodiagnosis in Clinical Neurology*, 6th ed. Oxford, Elsevier Saunders, 2012.)

commonly encountered regardless of whether seizures occur. The EEG findings are of limited utility in determining whether anticonvulsant medication can be discontinued after several seizure-free years. Further seizures may occur after withdrawal of anticonvulsant medication despite a normal EEG or, conversely, may not occur despite a continuing EEG abnormality. The decision to discontinue anticonvulsant medication is made on clinical grounds, and the EEG is helpful only for providing guidance when there is clinical ambiguity or the patient requires reassurance about a particular course of action.

The EEG has no role in the management of tonic-clonic status epilepticus except when there is clinical uncertainty about whether seizures are continuing in a comatose patient. In patients treated by drug-induced coma for refractory status epilepticus, the EEG findings indicate the level of anesthesia and whether seizures are occurring.

During status epilepticus, the EEG shows repeated electrographic seizures or continuous spike-wave discharges. In nonconvulsive status epilepticus, a disorder that may not be recognized unless an EEG is performed, the EEG may also show continuous spike-wave activity ("spike-wave stupor") or, less commonly, repetitive electrographic seizures (focal status epilepticus).

THE EEG AND COMA

The EEG tends to become slower as consciousness is depressed, regardless of the underlying cause (Fig. 442e-1). Other findings may suggest diagnostic possibilities, as when electrographic seizures are found or a focal abnormality indicates a structural lesion. The EEG generally slows in metabolic encephalopathies, and triphasic waves may be present. The findings do not permit differentiation of the underlying metabolic disturbance but help to exclude other encephalopathic processes by indicating the diffuse extent of cerebral dysfunction. An EEG responsive to external stimulation is helpful prognostically because electrocerebral responsiveness implies a lighter level of coma than a nonreactive EEG, and thus a better prognosis. Serial records provide a better guide to prognosis than a single record and supplement the clinical examination in following the course of events. As the depth of coma increases, the EEG becomes nonreactive and may show a burst-suppression pattern, with bursts of mixed-frequency activity separated by intervals of relative cerebral inactivity. In other instances there is a reduction in amplitude of the EEG until eventually activity cannot be detected. Such electrocerebral silence does not necessarily reflect irreversible brain damage, because it may occur reversibly in hypothermic patients or with drug overdose. The prognosis of electrocerebral silence, when recorded using an adequate technique, therefore depends on the clinical context in which it is found. In patients with severe cerebral anoxia, for example, electrocerebral silence in a technically satisfactory record implies that useful cognitive recovery will not occur.

In patients with clinically suspected brain death, an EEG recorded using appropriate technical standards may be confirmatory by showing electrocerebral silence, but disorders that may produce a similar but reversible EEG appearance must be excluded. The presence of residual EEG activity in suspected brain death fails to confirm the diagnosis but does not exclude it. The EEG is usually normal in patients with locked-in syndrome (Chap. 446), and helps in distinguishing this disorder from the comatose state with which it is sometimes confused clinically.

THE EEG IN OTHER NEUROLOGIC DISORDERS

In developed countries, computed tomography (CT) scanning and magnetic resonance imaging (MRI) are used as a noninvasive means of screening for focal structural abnormalities of the brain, such as tumors, infarcts, or hematomas (Fig. 442e-1). The EEG is still used for this purpose in many parts of the world, however, although infratentorial or slowly expanding lesions may not be recognized. Focal slow-wave disturbances, a localized loss of electrocerebral activity, or more generalized electrocerebral disturbances are common findings but do not indicate the nature of the underlying pathology.

In patients with an acute encephalopathy, focal or lateralized periodic slow-wave complexes, sometimes with a sharpened outline, suggest a diagnosis of herpes simplex encephalitis, and periodic lateralizing epileptiform discharges (PLEDs) are commonly found with acute hemispheric pathology such as a hematoma, abscess, or rapidly expanding tumor. The EEG findings in dementia are usually nonspecific and do not distinguish reliably between different underlying causes except in rare instances when the presence of complexes occurring with a regular repetition rate ("periodic complexes") supports a diagnosis of Creutzfeldt-Jakob disease (Fig. 442e-1) or subacute sclerosing panencephalitis. In most patients with dementia, the EEG is normal or diffusely slowed, and the findings alone cannot indicate whether a patient is demented or distinguish between dementia and pseudodementia.

CONTINUOUS EEG MONITORING

The brief EEG obtained routinely in the laboratory often fails to reveal abnormalities that are transient and infrequent. Continuous monitoring over 12 or 24 hours or longer may detect abnormalities or capture

clinical events that otherwise would be missed. The EEG is often recorded continuously in critically ill patients to detect early changes in neurologic status. Continuous EEG recording in this context has been used to detect acute events such as from nonconvulsive seizures or developing cerebral ischemia, to monitor cerebral function in patients with metabolic disorders such as liver failure, and to manage the level of anesthesia in pharmacologically induced coma.

MAGNETOENCEPHALOGRAPHY AND MAGNETIC SOURCE IMAGING

Recording the magnetic field of the electrical activity of the brain (magnetoencephalography [MEG]) provides a means of examining cerebral activity that is less subject to distortion by other biologic tissues than the EEG. MEG is used in only a few specialized centers because of the complexity and expense of the necessary equipment. It permits the source of activity to be localized and coregistered with the MRI in a technique that is known as *magnetic source imaging*. In patients with focal epilepsy, MEG is useful in localizing epileptogenic foci for surgery and for guiding the placement of intracranial electrodes for electrophysiologic monitoring. MEG has also been used for mapping brain tumors, identifying the central fissure preoperatively, and localizing functionally eloquent cortical areas such as those concerned with language.

EVOKED POTENTIALS

SENSORY EVOKED POTENTIALS

The noninvasive recording of spinal or cerebral potentials elicited by stimulation of specific afferent pathways allows the functional integrity of these pathways to be monitored but does not indicate the pathologic basis of lesions involving them. Such evoked potentials (EPs) are small compared to the background EEG activity, and the responses to a number of stimuli are therefore recorded and averaged with a computer to permit their recognition and definition. The background EEG activity, which has no fixed temporal relationship to the stimulus, is averaged out by this procedure.

Visual evoked potentials (VEPs) are elicited by monocular stimulation with a reversing checkerboard pattern and are recorded from the occipital region in the midline and on either side of the scalp. The component of major clinical importance is the so-called P100 response, a positive peak having a latency of approximately 100 ms. Its presence, latency, and symmetry over the two sides of the scalp are noted. Amplitude changes are less helpful for the recognition of pathology. VEPs are most useful in detecting dysfunction of the visual pathways anterior to the optic chiasm. In acute severe optic neuritis, the P100 is frequently lost or grossly attenuated; as clinical recovery occurs, it is restored but with an increased latency that generally remains abnormal indefinitely. The VEP findings are therefore helpful in indicating previous or subclinical optic neuritis. They may also be abnormal with ocular abnormalities and with other causes of optic nerve disease, such as ischemia or compression by a tumor. Flash-elicited VEPs may be normal in patients with cortical blindness.

Routine VEPs record a mass response over a relatively large cortical area and thus may be insensitive to localized waveform abnormalities. A newer technique, *multifocal VEP*, measures responses from 120 individual sectors within each affected eye and is more sensitive than routine VEP.

Brainstem auditory evoked potentials (BAEPs) are elicited by monaural stimulation with repetitive clicks and are recorded between the vertex of the scalp and the mastoid process or earlobe. A series of potentials, designated by roman numerals, occurs in the first 10 ms after the stimulus and represents in part the sequential activation of different structures in the pathway between the auditory nerve (wave I) and the inferior colliculus (wave V) in the midbrain. The presence, latency, and interpeak latency of the first five positive potentials recorded at the vertex are evaluated. The findings are helpful in screening for acoustic neuromas, detecting brainstem pathology, and evaluating comatose patients. The BAEPs are often normal in coma due to metabolic/toxic disorders or bihemispheric disease but are typically abnormal in the presence of brainstem pathology.

Somatosensory evoked potentials (SSEPs) are recorded over the scalp and spine in response to electrical stimulation of a peripheral (mixed or cutaneous) nerve. The configuration, polarity, and latency of the responses depend on the nerve that is stimulated and on the recording arrangements. SSEPs are used to evaluate proximal (otherwise inaccessible) portions of the peripheral nervous system and the integrity of the central somatosensory pathways, especially in patients who are comatose or suspected to be brain dead.

Clinical Utility of EPs EP studies may detect and localize lesions in afferent pathways in the central nervous system (CNS). They have been used particularly to investigate patients with suspected multiple sclerosis (MS), the diagnosis of which requires the recognition of multifocal white-matter lesions. In patients with clinical evidence of a single lesion, the electrophysiologic recognition of abnormalities in other sites helps to support the diagnosis but does not establish it unequivocally. Multimodality EP abnormalities are not specific for MS; they may occur in AIDS, Lyme disease, systemic lupus erythematosus, neurosyphilis, spinocerebellar degenerations, familial spastic paraplegia, and deficiency of vitamin E or B₁₂, among other disorders. The diagnostic utility of the EP findings therefore depends on the circumstances in which they are found. Abnormalities may aid in the localization of lesions to broad areas of the CNS, but attempts at precise localization may be misleading because the generators of many components are unknown.

The EP findings are sometimes of prognostic relevance. Bilateral loss of cortically generated SSEP components implies that cognition may not be regained in posttraumatic or postanoxic coma, and EP studies may also be useful in evaluating patients with suspected brain death. In patients who are comatose for uncertain reasons, preserved BAEPs suggest either a metabolic-toxic etiology or bihemispheric disease. In patients with spinal cord injuries, SSEPs have been used to indicate the completeness of the lesion. The presence or early return of a cortically generated response to stimulation of a nerve below the injured segment of the cord indicates an incomplete lesion and thus a better prognosis for functional recovery than otherwise. In surgery, intraoperative EP monitoring of neural structures placed at risk by the procedure may permit the early recognition of dysfunction and thereby permit a neurologic complication to be averted or minimized.

Visual and auditory acuity may be determined using EP techniques in patients whose age or mental state precludes traditional ophthalmologic or audiologic examinations.

COGNITIVE EVOKED POTENTIALS

Certain EP components depend on the mental attention of the subject and the setting in which the stimulus occurs, rather than simply on the physical characteristics of the stimulus. Such “event-related” potentials (ERPs) or “endogenous” potentials are related in some manner to the cognitive aspects of distinguishing an infrequently occurring target stimulus from other stimuli occurring more frequently. For clinical purposes, attention has been directed particularly at the so-called P3 component of the ERP, which is also designated the P300 component because of its positive polarity and latency of approximately 300–400 ms after onset of an auditory target stimulus. The P3 component is prolonged in latency in many patients with dementia, whereas it is generally normal in patients with depression or other disorders simulating dementia. ERPs are, therefore, sometimes helpful in making this distinction when there is clinical uncertainty, although a response of normal latency does not exclude dementia.

MOTOR EVOKED POTENTIALS

The electrical potentials recorded from muscle or the spinal cord following stimulation of the motor cortex or central motor pathways are referred to as *motor evoked potentials*. For clinical purposes, such responses are recorded most often as the compound muscle action potentials elicited by transcutaneous magnetic stimulation of the motor cortex. A strong but brief magnetic field is produced by passing a current through a coil, and this induces stimulating currents in the subjacent neural tissue. The procedure is painless and apparently safe.

Abnormalities have been described in several neurologic disorders with clinical or subclinical involvement of central motor pathways, including MS and motor neuron disease. In addition to a possible role in diagnosis or evaluating the extent of pathologic involvement, the technique provides information of prognostic relevance (e.g., in suggesting the likelihood of recovery of motor function after stroke) and provides a means of monitoring intraoperatively the functional integrity of central motor tracts. Nevertheless, it is not used widely for clinical purposes.

ELECTROPHYSIOLOGIC STUDIES OF MUSCLE AND NERVE

The motor unit is the basic element subserving motor function. It is defined as an anterior horn cell, its axon and neuromuscular junctions, and all the muscle fibers innervated by the axon. The number of motor units in a muscle ranges from approximately 10 in the extraocular muscles to several thousand in the large muscles of the legs. There is considerable variation in the average number of muscle fibers within the motor units of an individual muscle, i.e., in the innervation ratio of different muscles. Thus the innervation ratio is <25 in the human external rectus or platysma muscle and between 1600 and 1700 in the medial head of the gastrocnemius muscle. The muscle fibers of individual motor units are divided into two general types by distinctive contractile properties, histochemical stains, and characteristic responses to fatigue. Within each motor unit, all of the muscle fibers are of the same type.

ELECTROMYOGRAPHY

The pattern of electrical activity in muscle, i.e., the electromyogram (EMG), may be recorded both at rest and during activity from a needle electrode inserted into the muscle. The nature and pattern of abnormalities relate to disorders at different levels of the motor unit.

Relaxed muscle normally is electrically silent except in the end-plate region, but abnormal spontaneous activity (Fig. 442e-3) occurs in various neuromuscular disorders, especially those associated with denervation or inflammatory changes in affected muscle. *Fibrillation potentials* and *positive sharp waves* (which reflect muscle fiber irritability) and complex repetitive discharges are most often—but not always—found in denervated muscle and may also occur after muscle injury and in certain myopathic disorders, especially inflammatory disorders such as polymyositis. After an acute neuropathic lesion, they occur earlier in proximal than distal muscles and sometimes do not develop distally in the extremities for 4–6 weeks; once present, they may persist indefinitely unless reinnervation occurs or the muscle degenerates so completely that no viable tissue remains. *Fasciculation potentials*

(which reflect the spontaneous activity of individual motor units) are characteristic of slowly progressive neuropathic disorders, especially those with degeneration of anterior horn cells (such as amyotrophic lateral sclerosis). *Myotonic discharges*—high-frequency discharges of potentials derived from single muscle fibers that wax and wane in amplitude and frequency—are the signature of myotonic disorders such as myotonic dystrophy or myotonia congenita but occur occasionally in polymyositis or other, rarer, disorders.

Slight voluntary contraction of a muscle leads to activation of a small number of motor units. The potentials generated by muscle fibers of these units that are within the pickup range of the needle electrode will be recorded (Fig. 442e-3). The parameters of normal motor unit action potentials depend on the muscle under study and age of the patient, but their duration is normally between 5 and 15 ms, amplitude is between 200 μ V and 2 mV, and most are bi- or triphasic. The number of units activated depends on the degree of voluntary activity. An increase in muscle contraction is associated with an increase in the number of motor units that are activated (recruited) and in the frequency of discharge. With a full contraction, so many motor units are normally activated that individual motor unit action potentials can no longer be distinguished, and a complete interference pattern is said to have been produced.

The incidence of small, short-duration, polyphasic motor unit action potentials (i.e., having more than four phases) is usually increased in myopathic muscle, and an excessive number of units is activated for a specified degree of voluntary activity. By contrast, the loss of motor units that occurs in neuropathic disorders leads to a reduction in number of units activated during a maximal contraction and an increase in their firing rate, i.e., there is an incomplete or reduced interference pattern. The configuration and dimensions of the potentials may also be abnormal, depending on the duration of the neuropathic process. The surviving motor units are initially normal in configuration but, as reinnervation occurs, they increase in amplitude and duration and become polyphasic (Fig. 442e-3).

Action potentials from the same motor unit sometimes fire with a consistent temporal relationship to each other, so that double, triple, or multiple discharges are recorded, especially in tetany, hemifacial spasm, or myokymia.

Electrical silence characterizes the involuntary, sustained muscle contraction that occurs in phosphorylase deficiency, which is designated a contracture.

EMG enables disorders of the motor units to be detected and characterized as either neurogenic or myopathic. In neurogenic disorders, the pattern of affected muscles may localize the lesion to the anterior horn cells or to a specific site as the axons traverse a nerve root, limb plexus, and peripheral nerve to their terminal arborizations. The findings do not enable a specific etiologic diagnosis to be made, however, except in conjunction with the clinical findings and results of other laboratory studies.

The findings may provide a guide to the severity of an acute disorder of a peripheral or cranial nerve (by indicating whether denervation has occurred and the completeness of the lesion) and whether the pathologic process is active or progressive in chronic or degenerative disorders such as amyotrophic lateral sclerosis. Such information is important for prognostic purposes.

Various quantitative EMG approaches have been developed. The most common is to determine the mean duration and amplitude of 20 motor unit action potentials using a standardized technique. The technique of macro-EMG provides information about the number and size of muscle fibers in a larger volume of the motor unit territory and has also been used to estimate the number of motor units in a muscle. Scanning EMG is a computer-based technique that has been used to study the topography of motor unit action potentials and, in particular, the spatial and temporal distribution of activity in individual units. The technique of single-fiber EMG is discussed separately below.

NERVE CONDUCTION STUDIES

Recording of the electrical response of a muscle to stimulation of its motor nerve at two or more points along its course (Fig. 442e-4)

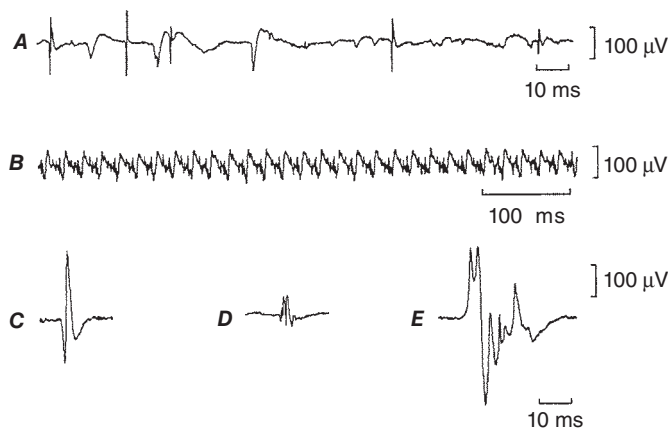


FIGURE 442e-3 Activity recorded during electromyography (EMG). **A.** Spontaneous fibrillation potentials and positive sharp waves. **B.** Complex repetitive discharges recorded in partially denervated muscle at rest. **C.** Normal triphasic motor unit action potential. **D.** Small, short-duration, polyphasic motor unit action potential such as is commonly encountered in myopathic disorders. **E.** Long-duration polyphasic motor unit action potential such as may be seen in chronic neuropathic disorders.

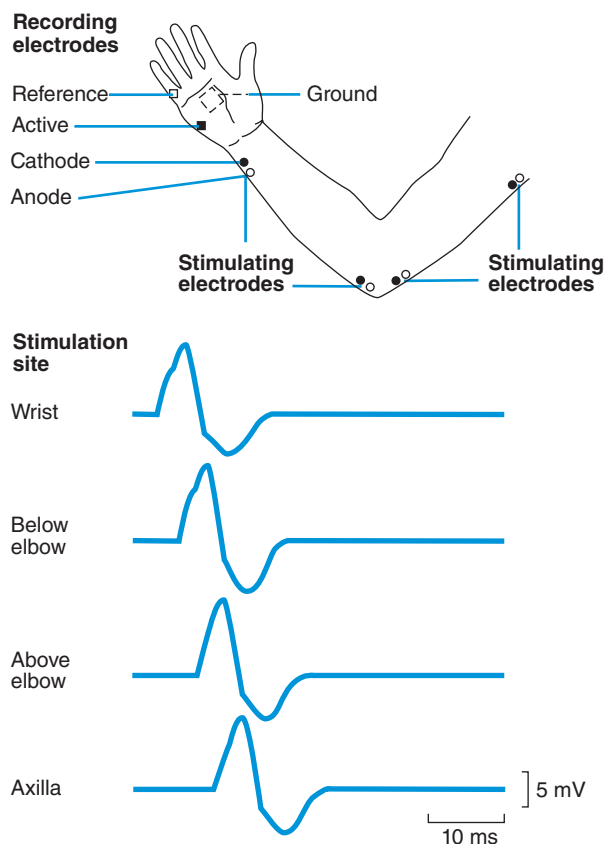


FIGURE 442e-4 Arrangement for motor conduction studies of the ulnar nerve. Responses are recorded with a surface electrode from the abductor digiti minimi muscle to supramaximal stimulation of the nerve at different sites, and are shown in the lower panel. (From MJ Aminoff: *Electromyography in Clinical Practice: Electrodiagnostic Aspects of Neuromuscular Disease*, 3rd ed. New York, Churchill Livingstone, 1998.)

permits conduction velocity to be determined in the fastest conducting motor fibers between the points of stimulation. The latency and amplitude of the electrical response of muscle (i.e., of the compound muscle action potential) to stimulation of its motor nerve at a distal site are also compared with values defined in normal subjects. Sensory nerve conduction studies are performed by determining the conduction velocity and amplitude of action potentials in sensory fibers when these fibers are stimulated at one point and the responses are recorded at another point along the course of the nerve. In adults, conduction velocity in the arms is normally between 50 and 70 m/s, and in the legs is between 40 and 60 m/s.

Nerve conduction studies complement the EMG examination, enabling the presence and extent of peripheral nerve pathology to be determined. They are particularly helpful in determining whether sensory symptoms are arising from pathology proximal or distal to the dorsal root ganglia (in the former instance, peripheral sensory conduction studies is normal) and whether neuromuscular dysfunction relates to peripheral nerve disease. In patients with a mononeuropathy, they are invaluable as a means of localizing a focal nerve lesion, determining the extent and severity of the underlying pathology, providing a guide to prognosis, and detecting subclinical involvement of other nerves. They enable a polyneuropathy to be distinguished from a mononeuropathy multiplex, which has important etiologic implications. Nerve conduction studies provide a means of following the progression and therapeutic response of peripheral nerve disorders and are used widely for this purpose in clinical trials. They may suggest the underlying pathologic basis in individual cases. Conduction velocity is often markedly slowed, terminal motor latencies are prolonged, and compound motor and sensory nerve action potentials may be dispersed in the demyelinating neuropathies (such as in Guillain-Barré syndrome, chronic inflammatory polyneuropathy, metachromatic leukodystrophy, or certain hereditary

neuropathies); conduction block is frequent in acquired varieties of these neuropathies. By contrast, conduction velocity is normal or slowed only mildly, sensory nerve action potentials are small or absent, and there is EMG evidence of denervation in axonal neuropathies such as occur in association with metabolic or toxic disorders.

The utility and complementary role of EMG and nerve conduction studies are best illustrated by reference to a common clinical problem. Numbness and paresthesias of the little finger and associated wasting of the intrinsic muscles of the hand may result from a spinal cord lesion, C8/T1 radiculopathy, brachial plexopathy (lower trunk or medial cord), or a lesion of the ulnar nerve. If sensory nerve action potentials can be recorded normally at the wrist following stimulation of the digital fibers in the affected finger, the pathology is probably proximal to the dorsal root ganglia (i.e., there is a radiculopathy or more central lesion); absence of the sensory potentials, by contrast, suggests distal pathology. EMG examination will indicate whether the pattern of affected muscles conforms to radicular or ulnar nerve territory or is more extensive (thereby favoring a plexopathy). Ulnar motor conduction studies will generally also distinguish between a radiculopathy (normal findings) and ulnar neuropathy (abnormal findings) and will often identify the site of an ulnar nerve lesion. The nerve is stimulated at several points along its course to determine whether the compound action potential recorded from a distal muscle that it supplies shows a marked alteration in size or area or a disproportionate change in latency, with stimulation at a particular site. The electrophysiologic findings thus permit a definitive diagnosis to be made and specific treatment instituted in circumstances where there is clinical ambiguity.

F-WAVE STUDIES

Stimulation of a motor nerve causes impulses to travel antidromically (i.e., toward the spinal cord) as well as orthodromically (to the nerve terminals). Such antidromic impulses cause a few of the anterior horn cells to discharge, producing a small motor response that occurs considerably later than the direct response elicited by nerve stimulation. The F wave so elicited is sometimes abnormal (absent or delayed) with proximal pathology of the peripheral nervous system, such as a radiculopathy, and may therefore be helpful in detecting abnormalities when conventional nerve conduction studies are normal. In general, however, the clinical utility of F-wave studies has been disappointing, except perhaps in Guillain-Barré syndrome, where they are often absent or delayed.

H-REFLEX STUDIES

The H reflex is easily recorded only from the soleus muscle (S1) in normal adults. It is elicited by low-intensity stimulation of the tibial nerve and represents a monosynaptic reflex in which spindle (Ia) afferent fibers constitute the afferent arc and alpha motor axons the efferent pathway. The H reflexes are often absent bilaterally in elderly patients or with polyneuropathies and may be lost unilaterally in S1 radiculopathies.

MUSCLE RESPONSE TO REPETITIVE NERVE STIMULATION

The size of the electrical response of a muscle to supramaximal electrical stimulation of its motor nerve relates to the number of muscle fibers that are activated. Neuromuscular transmission can be tested by several different protocols, but the most helpful is to record with surface electrodes the electrical response of a muscle to supramaximal stimulation of its motor nerve by repetitive (2–3 Hz) shocks delivered before and at selected intervals after a maximal voluntary contraction.

There is normally little or no change in size of the muscle response to repetitive stimulation of a motor nerve at 2–3 Hz with stimuli delivered at intervals after voluntary contraction of the muscle for about 20–30 s, even though preceding activity in the junctional region influences the release of acetylcholine and thus the size of the end-plate potentials elicited by a test stimulus. This is because more acetylcholine is normally released than is required to bring the motor end-plate potentials to the threshold for generating muscle fiber action potentials. In disorders of neuromuscular transmission this safety

factor is reduced. Thus in myasthenia gravis, repetitive stimulation, particularly at a rate of between 2 and 5 Hz, may lead to a depression of neuromuscular transmission, with a decrement in size of the response recorded from affected muscles. Similarly, immediately after a period of maximal voluntary activity, single or repetitive stimuli of the motor nerve may elicit larger muscle responses than before, indicating that more muscle fibers are responding. This postactivation facilitation of neuromuscular transmission is followed by a longer-lasting period of depression, maximal between 2 and 4 min after the conditioning period and lasting for as long as 10 min or so, during which responses are reduced in size.

Decrementing responses to repetitive stimulation at 2–5 Hz are common in myasthenia gravis but may also occur in the congenital myasthenic syndromes (Chap. 461). In Lambert-Eaton myasthenic syndrome, in which there is defective release of acetylcholine at the neuromuscular junction, the compound muscle action potential elicited by a single stimulus is generally very small. With repetitive stimulation at rates of up to 10 Hz, the first few responses may decline in size, but subsequent responses increase. If faster rates of stimulation are used (20–50 Hz), the increment may be dramatic so that the amplitude of compound muscle action potentials eventually reaches a size that is several times larger than the initial response. In patients with botulism, the response to repetitive stimulation is similar to that in Lambert-Eaton myasthenic syndrome, although the findings are somewhat more variable and not all muscles are affected.

SINGLE-FIBER ELECTROMYOGRAPHY

This technique is particularly helpful in detecting disorders of neuromuscular transmission. A special needle electrode is placed within

a muscle and positioned to record action potentials from two muscle fibers belonging to the same motor unit. The time interval between the two potentials will vary in consecutive discharges; this is called the *neuromuscular jitter*. The jitter can be quantified as the mean difference between consecutive interpotential intervals and is normally between 10 and 50 μ s. This value is increased when neuromuscular transmission is disturbed for any reason, and in some instances impulses in individual muscle fibers may fail to occur because of impulse blocking at the neuromuscular junction. Single-fiber EMG is more sensitive than repetitive nerve stimulation or determination of acetylcholine receptor antibody levels in diagnosing myasthenia gravis.

Single-fiber EMG can also be used to determine the mean fiber density of motor units (i.e., mean number of muscle fibers per motor unit within the recording area) and to estimate the number of motor units in a muscle, but this is of less immediate clinical relevance.

BLINK REFLEXES

Electrical or mechanical stimulation of the supraorbital nerve on one side leads to two separate reflex responses of the orbicularis oculi—an ipsilateral R1 response having a latency of approximately 10 ms and a bilateral R2 response with a latency in the order of 30 ms. The trigeminal and facial nerves constitute the afferent and efferent arcs of the reflex, respectively. Abnormalities of either nerve or intrinsic lesions of the medulla or pons may lead to uni- or bilateral loss of the response, and the findings may therefore be helpful in identifying or localizing such pathology.

443e Technique of Lumbar Puncture

Elizabeth Robbins, Stephen L. Hauser

In experienced hands, lumbar puncture (LP) is usually a safe procedure. Major complications are extremely uncommon but can include cerebral herniation, injury to the spinal cord or nerve roots, hemorrhage (spinal hematoma), or infection. Minor complications occur with greater frequency and can include backache, post-LP headache, and radicular pain or numbness.

IMAGING AND LABORATORY STUDIES PRIOR TO LP

Patients with an altered level of consciousness, a focal neurologic deficit, new-onset seizure, papilledema, or an immunocompromised state are at increased risk for potentially fatal cerebellar or tentorial herniation following LP. Neuroimaging should be obtained in these patients prior to LP to exclude a focal mass lesion or diffuse swelling. Imaging studies should include the spine in patients with symptoms suggesting spinal cord compression, such as back pain, leg weakness, urinary retention, or incontinence. In patients with suspected meningitis who require neuroimaging prior to diagnostic LP, administration of antibiotics, preferably following blood culture, should precede the neuroimaging study.

LP should not be performed through infected skin, as organisms can be introduced into the subarachnoid space (SAS).

Patients with coagulation defects including thrombocytopenia are at increased risk of post-LP spinal subdural or epidural hematomas, either of which can produce permanent nerve injury and/or paralysis. If a bleeding disorder is suspected, the platelet count, international normalized ratio (INR), and partial thromboplastin time should be checked prior to LP. There are no data available to assess the safety of LP in patients with low platelet counts; a count of $<20,000/\mu\text{L}$ is considered to be a contraindication to LP. Bleeding complications rarely occur in patients with platelet counts $\geq 50,000/\mu\text{L}$ and an INR ≤ 1.5 . Some institutions recommend that the platelet count be $>40,000$ prior to LP.

GUIDELINES FOR PATIENTS RECEIVING ANTICOAGULANT OR ANTIPLATELET MEDICATIONS

There is an increased risk of bleeding complications if an LP is performed in a patient receiving antiplatelet or anticoagulant medications. The risk is further increased when multiple anticoagulant medications are used or when the level of anticoagulation is high. The most common site of bleeding is the epidural space. Symptoms of bleeding following an LP can include a sensory or motor deficit and/or bowel/bladder dysfunction; back pain occurs less commonly. For serious deficits such as paraparesis, immediate surgical intervention, ideally within 8 h of onset of weakness, is important to minimize permanent disability; surgical intervention after 24 h is associated with a poor outcome.

Only limited data are available to guide decisions about performing LPs in patients receiving anticoagulant drugs. Information about managing antiplatelet and anticoagulation drugs during invasive surgical procedures is often available from the prescribing information provided by the drug manufacturer. Evidence-based guidelines for management of regional anesthetic procedures including spinal and epidural blocks in patients receiving anticoagulation have been developed by the American Society of Regional Anesthesia and Pain (ASRA); these guidelines can help guide decisions by physicians considering LP in patients receiving anticoagulation. Management of these patients can be complex and needs to consider both the risk of LP-related hemorrhage as well as the risk of reversing therapeutic anticoagulation prior to LP. Guidelines for some commonly used anticoagulants are summarized below.

Unfractionated Heparin, Therapeutic Dosing The ASRA 2010 Practice Advisory recommends discontinuing unfractionated heparin (UFH)

2–4 h prior to removal of spinal or epidural catheters to minimize risk of hematoma. Similar guidelines are reasonable for patients undergoing LP: discontinue UFH 2–4 h prior to LP; document normal partial thromboplastin time (PTT) prior to the procedure; and document a normal platelet count in patients who have received heparin for 4 days or longer because of the risk of heparin-induced thrombocytopenia (HIT). The half-life of heparin is 60–90 min.

Unfractionated Heparin, Prophylactic Dosing There are only a few case reports of spinal hematoma resulting from spinal or epidural anesthetic procedures in patients receiving low-dose subcutaneous UFH; ASRA guidelines state that there is no contraindication to the use of these techniques for anesthesia in patients receiving prophylactic UFH at a dose of 5000 U subcutaneously twice daily. Similarly, LP in patients receiving 5000 U of UFH subcutaneously twice daily is unlikely to cause spinal hematoma. Precautions to minimize risk include the following: document a normal PTT prior to the LP; document a normal platelet count in patients who have received heparin for 4 days or longer; and perform the LP 1–2 h prior to the next heparin dose, when the heparin effect should be minimal.

Low-Molecular-Weight Heparin, Therapeutic Dose (e.g., Enoxaparin 1 mg/kg Subcutaneously q12h) Patients receiving low-molecular-weight heparin (LMWH) are at increased risk of post-LP spinal or epidural hematoma. LMWH dose should be held for at least 24 h before the procedure.

Low-Molecular-Weight Heparin, Prophylactic Dose (e.g., Enoxaparin 0.5 mg/kg Subcutaneously q12h) Patients receiving prophylactic-dose LMWH have altered coagulation. ASRA guidelines recommend waiting at least 10–12 h after a prophylactic dose of LMWH before inserting a spinal or epidural catheter to minimize the risk of spinal or epidural hematoma. Similar guidelines are reasonable for patients undergoing LP.

Warfarin Spinal puncture is contraindicated during warfarin therapy.

Aspirin and Nonsteroidal Anti-inflammatory Drugs (NSAIDs) ASRA guidelines conclude that use of these drugs does not appear to be associated with an added significant risk of spinal bleeding in patients having spinal or epidural anesthesia. Similarly, LP in patients receiving one of these drugs is unlikely to cause bleeding. Reversal of drug effect on platelet function requires stopping the drug for approximately 10 days for aspirin and for 48 h for NSAIDs.

Ticlopidine/Clopidogrel The actual risk of spinal hematoma with these drugs is unknown. Based on drug prescribing information and surgical reviews, ASRA guidelines suggest discontinuing ticlopidine 14 days prior to a spinal or epidural procedure and discontinuing clopidogrel 7 days prior to the procedure. Similar guidelines are reasonable for performing LP.

Abciximab, Eptifibatide, and Other Platelet Glycoprotein IIb/IIIa Inhibitors The actual risk of spinal hematoma with these drugs is unknown. Platelet aggregation remains abnormal for 24–48 h following discontinuation of abciximab and 4–8 h following discontinuation of eptifibatide. ASRA guidelines recommend avoiding spinal or epidural procedures until platelet function is normal. Similar guidelines are reasonable for performing LP.

Direct Thrombin Inhibitors (e.g., Argatroban, Bivalirudin) ASRA guidelines recommend against performing spinal or epidural anesthesia in patients receiving thrombin inhibitors.

Oral Factor Xa Inhibitor (e.g., Rivaroxaban) Rivaroxaban prescribing information includes a black box warning that epidural or spinal hematomas have occurred in patients treated with rivaroxaban who are receiving spinal or epidural anesthesia or undergoing LP. LP should be avoided in patients receiving this drug.

ANALGESIA

Anxiety and pain can be minimized prior to beginning the procedure. Anxiety can be allayed by the use of lorazepam, 1–2 mg given PO 30 min prior to the procedure or IV 5 min prior to the procedure.

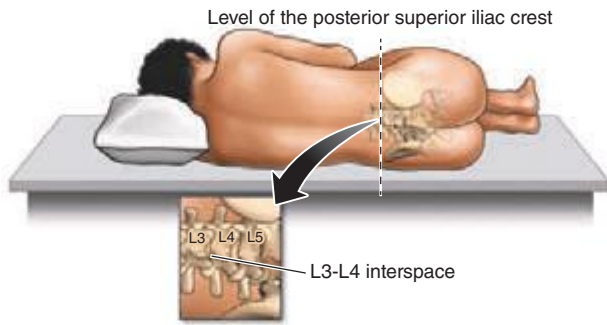


FIGURE 443e-1 Proper positioning of a patient in the lateral decubitus position. Note that the shoulders and hips are in a vertical plane; the torso is perpendicular to the bed. (From RP Simon et al [eds]: *Clinical Neurology*, 7th ed. New York, McGraw-Hill, 2009.)

Topical anesthesia can be achieved by the application of a lidocaine-based cream. Lidocaine 4% is effective when applied 30 min prior to the procedure; lidocaine/prilocaine requires 60–120 min. The cream should be applied in a thick layer so that it completely covers the skin; an occlusive dressing is used to keep the cream in place.

POSITIONING

Proper positioning of the patient is essential. The procedure should be performed on a firm surface; if the procedure is to be performed at the bedside, the patient should be positioned at the edge of the bed and not in the middle. The patient is asked to lie on his or her side, facing away from the examiner, and to “roll up into a ball.” The neck is gently ante-flexed and the thighs pulled up toward the abdomen; the shoulders and pelvis should be vertically aligned without forward or backward tilt (Fig. 443e-1). The spinal cord terminates at approximately the L1 vertebral level in 94% of individuals. In the remaining 6%, the conus extends to the L2–L3 interspace. LP is therefore performed at or below the L3–L4 interspace. A useful anatomic guide is a line drawn between the posterior superior iliac crests, which corresponds closely to the level of the L3–L4 interspace. The interspace is chosen following gentle palpation to identify the spinous processes at each lumbar level.

An alternative to the lateral recumbent position is the seated position. The patient sits at the side of the bed, with feet supported on a chair. The patient is instructed to curl forward, trying to touch the nose to the umbilicus. It is important that the patient not simply lean forward onto a bedside tabletop, as this is not an optimal position for opening up the spinous processes. LP is sometimes more easily performed in obese patients if they are sitting. A disadvantage of the seated position is that measurement of opening pressure is not accurate. In situations in which LP is difficult using palpable spinal landmarks, bedside ultrasound to guide needle placement may be used. In some particularly difficult situations, a computed tomography (CT)-guided needle placement may be necessary.

TECHNIQUE

Once the desired site for needle insertion has been identified, the examiner should put on sterile gloves. A mask is worn if the clinician will be injecting material into the spinal or epidural space to prevent droplet spread of oral flora during the procedure. After cleansing the skin with povidone-iodine or similar disinfectant, the area is draped with a sterile cloth; the needle insertion site is blotted dry using a sterile gauze pad. Proper local disinfection reduces the risk of introducing skin bacteria into the SAS or other sites. Local anesthetic, typically 1% lidocaine, 3–5 mL total, is injected into the subcutaneous tissue; in nonemergency situations, a topical anesthetic cream can be applied (see above). When time permits, pain associated with the injection of lidocaine can be minimized by slow, serial injections, each one progressively deeper than the last, over a period of ~5 min. Approximately 0.5–1 mL of lidocaine is injected at a time; the needle

is not usually withdrawn between injections. A pause of ~15 s between injections helps to minimize the pain of the subsequent injection. The goal is to inject each mini-bolus of anesthetic into an area of skin that has become numb from the preceding injection. Approximately 5–10 mini-boluses are injected, using a total of ~5 mL of lidocaine.

If possible, the LP should be delayed for 10–15 min following the completion of the injection of anesthetic; this significantly decreases and can even eliminate pain from the procedure. Even a delay of 5 min will help to reduce pain.

The LP needle (typically 20- to 22-gauge) is inserted in the midline, midway between two spinous processes, and slowly advanced. The bevel of the needle should be maintained in a horizontal position, parallel to the direction of the dural fibers and with the flat portion of the bevel pointed upward; this minimizes injury to the fibers as the dura is penetrated. When LP is performed in patients who are sitting, the bevel should be maintained in the vertical position. In most adults, the needle is advanced 4–5 cm (1–2 in.) before the SAS is reached; the examiner usually recognizes entry as a sudden release of resistance, a “pop.” If no fluid appears despite apparently correct needle placement, then the needle may be rotated 90°–180°. If there is still no fluid, the stylet is reinserted and the needle is advanced slightly. Some examiners halt needle advancement periodically to remove the stylet and check for flow of cerebrospinal fluid (CSF). If the needle cannot be advanced because it hits bone, if the patient experiences sharp radiating pain down one leg, or if no fluid appears (“dry tap”), the needle is partially withdrawn and reinserted at a different angle. If on the second attempt the needle still hits bone (indicating lack of success in introducing it between the spinous processes), then the needle should be completely withdrawn and the patient should be repositioned. The second attempt is sometimes more successful if the patient straightens the spine completely prior to repositioning. The needle can then be reinserted at the same level or at an adjacent one.

Once the SAS is reached, a manometer is attached to the needle and the opening pressure measured. The examiner should look for normal oscillations in CSF pressure associated with pulse and respirations. The upper limit of normal opening pressure with the patient supine is 180 mmH₂O in adults but may be as high as 200–250 mmH₂O in obese adults.

CSF is allowed to drip into collection tubes; it should not be withdrawn with a syringe. Depending on the clinical indication, fluid is obtained for studies including: (1) cell count with differential; (2) protein and glucose concentrations; (3) culture (bacterial, fungal, mycobacterial, viral); (4) smears (e.g., Gram’s and acid-fast stained smears); (5) antigen tests (e.g., latex agglutination); (6) polymerase chain reaction (PCR) amplification of DNA or RNA of microorganisms (e.g., herpes simplex virus, enteroviruses); (7) antibody levels against microorganisms; (8) immunoelectrophoresis for determination of γ -globulin level and oligoclonal banding; and (9) cytology. Although 15 mL of CSF is sufficient to obtain all of the listed studies, the yield of fungal and mycobacterial cultures and cytology increases when larger volumes are sampled. In general 20–30 mL may be safely removed from adults.

A bloody tap due to penetration of a meningeal vessel (a “traumatic tap”) may result in confusion with subarachnoid hemorrhage (SAH). In these situations a specimen of CSF should be centrifuged immediately after it is obtained; clear supernatant following CSF centrifugation supports the diagnosis of a bloody tap, whereas xanthochromic supernatant suggests SAH. In general, bloody CSF due to the penetration of a meningeal vessel clears gradually in successive tubes, whereas blood due to SAH does not. In addition to SAH, xanthochromic CSF may also be present in patients with liver disease and when the CSF protein concentration is markedly elevated (>1.5–2 g/L [150–200 mg/dL]).

Prior to removing the LP needle, the stylet is reinserted to avoid the possibility of entrapment of a nerve root in the dura as the needle is being withdrawn; entrapment could result in a dural CSF leak, causing headache. Some practitioners question the safety of this maneuver, with its potential risk of causing a needle-stick injury to the examiner. Injury is unlikely, however, given the flexibility of the small-diameter stylet, which tends to bend, rather than penetrate, on contact. Following LP,

the patient is customarily positioned in a comfortable, recumbent position for 30–60 min before rising, although this may be unnecessary because it does not appear to affect the development of headache (see below).

POST-LP HEADACHE

The principal complication of LP is headache, occurring in 10–30% of patients. Younger age and female gender are associated with an increased risk of post-LP headache. Headache usually begins within 48 h but may be delayed for up to 12 days. Head pain is dramatically positional; it begins when the patient sits or stands upright; there is relief upon reclining or with abdominal compression. The longer the patient is upright, the longer the latency before head pain subsides. The pain is usually a dull ache but may be throbbing; its location is occipitofrontal. Nausea and stiff neck often accompany headache, and occasionally, patients report blurred vision, photophobia, tinnitus, and vertigo. In more than three-quarters of patients, symptoms completely resolve within a week, but in a minority they can persist for weeks or even months.

Post-LP headache is caused by a drop in CSF pressure related to persistent leakage of CSF at the site where the needle entered the SAS. Loss of CSF volume decreases the brain's supportive cushion, so that when a patient is upright there is probably dilation and tension placed on the brain's anchoring structures, the pain-sensitive dural sinuses, resulting in pain. Although intracranial hypotension is the usual explanation for severe LP headache, the syndrome can occur in patients with normal CSF pressure.

Because post-LP headache usually resolves without specific treatment, care is largely supportive with oral analgesics (acetaminophen, NSAIDs, opioids [Chap. 18]) and antiemetics. Patients may obtain relief by lying in a comfortable (especially a recumbent or head-down Trendelenburg) position. For some patients, beverages with caffeine can provide temporary pain relief.

For patients with persistent pain, treatment with IV caffeine (500 mg in 500 mL saline administered over 2 h) may be effective; atrial fibrillation is a rare side effect. Alternatively, an epidural blood patch accomplished by injection of 15 mL of autologous whole blood is usually effective; the injection is directed at the epidural space at the level of the initial LP. This procedure is most often performed by a pain specialist or anesthesiologist. The blood patch has an immediate effect, making it unlikely that sealing off a dural hole with blood clot is its sole mechanism of action. The acute benefit may be due to compression of the CSF space by the clot, increasing CSF pressure. Some clinicians reserve epidural blood patch for patients who do not respond to caffeine, while others prefer to use blood patch as initial management for unremitting post-LP symptoms.

Strategies to decrease the incidence of post-LP headache are listed in Table 443e-1. Use of a smaller caliber needle is associated with a lower risk: in one study, the risk of headache following use of a 24- to 27-gauge standard (Quincke) needle was 5–12%, compared to 20–40% when a 20- or 22-gauge needle was used. The smallest gauge needles usually require the use of an introducer needle and are associated with a slower CSF flow rate. Use of an “atraumatic” (Sprotte, “pencil

TABLE 443e-1 REDUCING THE INCIDENCE OF POST-LP HEADACHE

Effective Strategies

Use of small-diameter needle (22-gauge or smaller)
Use of atraumatic needle (Sprotte and others)
Replacement of stylet prior to removal of needle
Insertion of needle with bevel oriented in a cephalad to caudad direction (when using standard needle)

Ineffective Strategies

Bed rest (up to 4 h) following LP
Supplemental fluids
Minimizing the volume of spinal fluid removed
Immediate mobilization following LP

Abbreviation: LP, lumbar puncture.

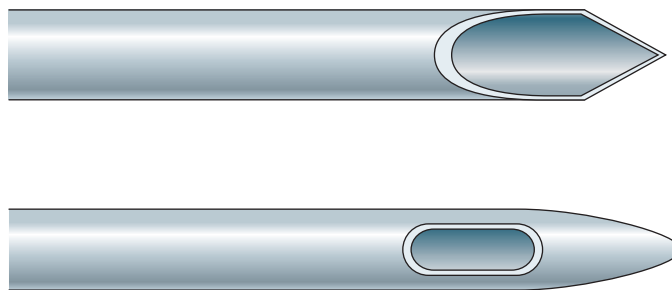


FIGURE 443e-2 Comparison of the standard (“cutting” or Quincke) lumbar puncture (LP) needle with the “atraumatic” (Sprotte). The “atraumatic” needle has its opening on the top surface of the needle, a design intended to reduce the chance of cutting dural fibers that, by protruding through the dura, could be responsible for subsequent cerebrospinal fluid leak and post-LP headache. (From SR Thomas et al: *BMJ* 321:986, 2000.)

point,” or “noncutting”) needle also reduces the incidence of moderate to severe headache compared with standard LP (Quincke, or “traumatic”) needles (Fig. 443e-2). However, because atraumatic needles are more difficult to use, more attempts may be required to perform the LP, particularly in overweight patients. It may also be necessary to use an introducer with the atraumatic needle, which does not have the customary cutting, beveled tip. There is a low risk of needle damage, e.g., breakage, with the Sprotte atraumatic needle. Another strategy to decrease the incidence of headache is to replace the stylet before removing the LP needle.

Patients are often advised to remain in a recumbent position for up to an hour following LP. However, studies comparing mobilization immediately following LP with bed rest for periods up to 4 h show no significant differences in the incidence of headache, suggesting that the customary practice of remaining in a recumbent position post-LP may be unnecessary.

TABLE 443e-2 CEREBROSPINAL FLUID (CSF)^a

Constituent	SI Units	Conventional Units
Glucose	2.22–3.89 mmol/L	40–70 mg/dL
Lactate	1–2 mmol/L	10–20 mg/dL
Total protein		
Lumbar	0.15–0.5 g/L	15–50 mg/dL
Cisternal	0.15–0.25 g/L	15–25 mg/dL
Ventricular	0.06–0.15 g/L	6–15 mg/dL
Albumin	0.066–0.442 g/L	6.6–44.2 mg/dL
IgG	0.009–0.057 g/L	0.9–5.7 mg/dL
IgG index ^b	0.29–0.59	
Oligoclonal bands (OGB)	<2 bands not present in matched serum sample	
Ammonia	15–47 μmol/L	25–80 μg/dL
CSF pressure		50–180 mmH ₂ O
CSF volume (adult)	~150 mL	
Red blood cells	0	0
Leukocytes		
Total	0–5 mononuclear cells per mm ³	
Differential		
Lymphocytes	60–70%	
Monocytes	30–50%	
Neutrophils	None	

^aBecause CSF concentrations are equilibrium values, measurements of the same parameters in blood plasma obtained at the same time are recommended. However, there is a time lag in attainment of equilibrium, and cerebrospinal levels of plasma constituents that can fluctuate rapidly (such as plasma glucose) may not achieve stable values until after a significant lag phase. ^bIgG index = CSF IgG (mg/dL) × Serum albumin (g/dL) / Serum IgG (g/dL) × CSF albumin (mg/dL).

443e-4 NORMAL VALUES

(See Table 443e-2) In uninfected CSF, the normal white blood cell count is fewer than five mononuclear cells (lymphocytes and monocytes) per μL . Polymorphonuclear leukocytes (PMNs) are not found in normal unconcentrated CSF; however, rare PMNs can be found

in centrifuged or concentrated CSF specimens such as those used for cytologic examination. Red blood cells (RBCs) are not normally present in CSF; if RBCs are present from a traumatic tap, their number decreases as additional CSF is collected. CSF glucose concentrations $<2.2 \text{ mmol/L}$ ($<40 \text{ mg/dL}$) are abnormal.

444e **Biology of Neurologic Diseases**

Stephen L. Hauser, Stanley B. Prusiner, M. Flint Beal

The human nervous system is the organ of consciousness, cognition, ethics, and behavior; as such, it is the most intricate structure known to exist. More than one-third of the 23,000 genes encoded in the human genome are expressed in the nervous system. Each mature brain is composed of 100 billion neurons, several million miles of axons and dendrites, and $>10^{15}$ synapses. Neurons exist within a dense parenchyma of multifunctional glial cells that synthesize myelin, preserve homeostasis, and regulate immune responses. Measured against this background of complexity, the achievements of molecular neuroscience have been extraordinary. This chapter reviews selected themes in neuroscience that provide a context for understanding fundamental mechanisms underlying neurologic disorders.

NEUROGENETICS

The landscape of neurology has been transformed by modern molecular genetics. Several hundred neurologic and psychiatric disorders can now be diagnosed through genetic testing (<http://www.ncbi.nlm.nih.gov/sites/GeneTests?db=GeneTests>). The vast majority of these represent highly penetrant mutations that cause rare neurologic disorders; alternatively, they represent rare monogenic causes of common phenotypes. Examples of the latter include mutations of the amyloid precursor protein in familial Alzheimer's disease, the microtubule-associated protein tau (MAPT) in frontotemporal dementia, and α -synuclein in Parkinson's disease. These discoveries have been profoundly important because the mutated gene in the familial disorder often encodes a protein that is also pathogenetically involved (although not mutated) in the typical, sporadic form. The common mechanism involves disordered processing and, ultimately, aggregation of the protein leading to cell death (see "Protein Aggregation and Neurodegeneration," below).

There is optimism that complex genetic disorders, caused by combinations of both genetic and environmental factors, have now become tractable problems. Genome-wide association studies (GWAS) have been carried out in many complex neurologic disorders, with many hundreds of variants identified, nearly all of which confer only a small increment in disease risk (1.15- to 1.5-fold). GWAS studies are rooted in the "common disease, common variant" hypothesis, as they examine potential risk alleles that are relatively frequent (e.g. $>5\%$) in the general population. More than 1500 GWAS studies have been carried out, with notable successes such as the identification of 110 risk alleles for multiple sclerosis (**Chap. 458**). Furthermore, using bioinformatics tools, risk variants can be aligned in functional biologic pathways to identify novel pathogenic mechanisms as well as to reveal heterogeneity (e.g., different pathways in different individuals). Despite these successes, many experienced geneticists question the real value of common disease-associated variants, particularly whether they are actually causative or merely mark the approximate locations of more important—truly causative—rare mutations.

This debate has set the stage for the next revolution in human genetics, made possible by the development of increasingly efficient and cost-effective high-throughput sequencing methodologies. It is already possible to sequence an entire human genome in approximately an hour, at a cost of only \$1300 for the entire coding sequence ("whole-exome") or \$3000 for the entire genome; it is certain that these costs will continue to decline. This makes it feasible to look for disease-causing sequence variations in individual patients with the possibility of identifying rare variants that cause disease. The utility of this approach was demonstrated by whole-genome sequencing in a patient with Charcot-Marie-Tooth neuropathy, in which compound heterozygous mutations were identified in the *SH3TC2* gene, which then were shown to co-segregate with the disease in other members of the family.

It is increasingly recognized that not all genetic diseases or predispositions are caused by simple changes in the linear nucleotide sequence of genes. Disease-causative mutations also occur commonly in

noncoding sequences of DNA. For example, large intronic GGGGCC repeat expansions in a gene of unknown function *C9orf72* (chromosome 9 open reading frame 72) were recently identified as a common cause of both frontotemporal dementia and amyotrophic lateral sclerosis (ALS). This mutation is the most common cause of both familial and sporadic ALS identified thus far. It was shown to be associated with TDP-43 (tar DNA binding protein-43) inclusions in both hippocampal and cerebral neurons. Interestingly, despite the absence of a start codon, the three alternate dipeptide sequences consisting of two amino acids are translated and found in postmortem brain tissue of affected patients. Three potential pathogenic mechanisms have been proposed, including (1) haploinsufficiency, (2) repeat RNA-mediated toxicity, and (3) dipeptide protein toxicity. The possibility of RNA toxicity is supported by the finding of intranuclear RNA foci containing *C9orf72* hexanucleotide repeats and specific RNA-binding proteins. The *C9orf72* mRNA forms quadruplexes of DNA and RNA, which then can halt transcription and also bind transcription factors. Mutations in both *TARDP* (transactive region DNA binding protein) and *FUS* (fused in sarcoma) also encode DNA/RNA-processing polypeptides and are a cause of familial and sporadic ALS.

As the complex architecture of the human genome becomes better defined, many disorders that result from alterations in copy numbers of genes ("gene-dosage" effects) resulting from unequal crossing-over are also likely to be identified. As much as 5–10% of the human genome consists of nonhomologous duplications and deletions, and these appear to occur with a much higher mutational rate than is the case for single base pair mutations. The first copy-number disorders to be recognized were Charcot-Marie-Tooth disease type 1A (CMT1A), caused by a duplication in the gene encoding the myelin protein PMP22, and the reciprocal deletion of the gene causing hereditary liability to pressure palsies (**Chap. 459**). Gene-dosage effects are causative in some cases of Parkinson's disease (α -synuclein), Alzheimer's disease (amyloid precursor protein), spinal muscular atrophy (survival motor neuron 2), the dysmyelinating disorder Pelizaeus-Merzbacher syndrome (proteolipid protein 1), late-onset leukodystrophy (lamin B1), and a variety of developmental neurologic disorders. It is likely that copy-number variations contribute substantially to normal human genomic variation for numerous genes involved in neurologic function, regulation of cell growth, and regulation of metabolism. It is also already clear that gene-dosage effects will influence many behavioral phenotypes, learning disorders, and autism spectrum disorders. Deletions at ch444eq and ch15q have been associated with schizophrenia, and deletions at 15q and 16p with autism. Interestingly, the 16p deletion is also associated with epilepsy. Duplications of the X-linked *MeCP2* gene cause autism in males and psychiatric disorders with anxiety in females, whereas point mutations in this gene produce the neurodevelopmental disorder Rett's syndrome. The understanding of the role of copy number variation in human disease is still in its infancy.

The role of splicing variation as a contributor to neurologic disease is another area of active investigation. *Alternative splicing* refers to the inclusion of different combinations of exons in mature mRNA, resulting in the potential for many different protein products encoded by a single gene. Alternative splicing represents a powerful mechanism for generation of complexity and variation, and this mechanism appears to be highly prevalent in the nervous system, affecting key processes such as neurotransmitter receptors and ion channels. Numerous diseases are already known to result from abnormalities in alternative splicing. Increased inclusion of exon 10-containing transcripts of *MAPT* can cause frontotemporal dementia. Aberrant splicing also contributes to the pathogenesis of Duchenne's, myotonic, and facioscapulohumeral muscular dystrophies; ataxia telangiectasia; neurofibromatosis; some inherited ataxias; and fragile X syndrome, among other disorders. It is also likely that subtle variations of splicing will influence many genetically complex disorders. A splicing variant of the interleukin 7 receptor α chain, resulting in production of more soluble and less membrane-bound receptor, is associated with susceptibility to multiple sclerosis (MS) in multiple different populations.

Epigenetics refers to the mechanisms by which levels of gene expression can be exquisitely modulated, not by variations in the primary

genetic sequence of DNA but rather by postgenomic alterations in DNA and chromatin structure, which influence how, when, and where genes are expressed. DNA methylation and methylation and acetylation of histone proteins that interact with nuclear DNA to form chromatin are key mediators of these events. Epigenetic processes appear to be dynamically active even in post-mitotic neurons. *Imprinting* refers to an epigenetic feature, present for a subset of genes, in which the predominant expression of one allele is determined by its parent of origin. The distinctive neurodevelopmental disorders Prader-Willi syndrome (mild mental retardation and endocrine abnormalities) and Angelman's syndrome (cortical atrophy, cerebellar dysmyelination, Purkinje cell loss) are classic examples of imprinting disorders whose distinctive features are determined by whether the paternal or maternal copy of the chromosome of the critical genetic region 15q11-13 is responsible. In a study of discordant monozygotic twins for MS in which the entire DNA sequence, transcriptome (e.g., mRNA levels), and methylome were assessed genome-wide, tantalizing allelic differences in the use of the paternal, compared to maternal, copy for a group of genes were identified. Preferential allelic expression, whether due to imprinting, resistance to X-inactivation, or other mechanisms, is likely to play a major role in determining complex behaviors and susceptibility to many neurologic and psychiatric disorders.

Another advance is the development of transgenic mouse models of neurologic diseases, which has been particularly fruitful in producing models relevant to Alzheimer's disease, Parkinson's disease, Huntington's disease, and ALS. These models are useful in both studying disease pathogenesis and developing and testing new therapies. New transgenic mouse models with conditional expression have fostered investigations in which late gene expression avoids developmental compensation or in which the reversibility of a disease phenotype can be examined by turning a gene off after the disease phenotype has manifested. One can also examine the effects of gene expression in specific subsets of neurons, such as entorhinal cortex, or selectively in neurons, astrocytes, or microglia. Models in both *Caenorhabditis elegans* and *Drosophila* have also been extremely useful, particularly in studying genetic modifiers and therapeutic interventions.

ION CHANNELS AND CHANNELOPATHIES

The resting potential of neurons and the action potentials responsible for impulse conduction are generated by ion currents and ion channels. Most ion channels are gated, meaning that they can transition between conformations that are open or closed to ion conductance. Individual ion channels are distinguished by the specific ions they conduct; by their kinetics; and by whether they directly sense voltage, are linked to receptors for neurotransmitters or other ligands such as neurotrophins, or are activated by second messengers. The diverse characteristics of different ion channels provide a means by which neuronal excitability can be exquisitely modulated at both the cellular and the subcellular levels. Disorders of ion channels—channelopathies—are responsible for a growing list of human neurologic diseases (Table 444e-1). Most are caused by mutations in ion channel genes or by autoantibodies against ion channel proteins. One example is epilepsy, a syndrome of diverse causes characterized by repetitive, synchronous firing of neuronal action potentials. Action potentials are normally generated by the opening of sodium channels and the inward movement of sodium ions down the intracellular concentration gradient. Depolarization of the neuronal membrane opens potassium channels, resulting in outward movement of potassium ions, repolarization, closure of the sodium channel, and hyperpolarization. Sodium or potassium channel subunit genes have long been considered candidate disease genes in inherited

TABLE 444e-1 EXAMPLES OF NEUROLOGIC CHANNELOPATHIES

Category	Disorder	Channel Type	Mutated Gene	Chap. Ref.
Genetic				
Ataxias	Episodic ataxia-1	K	<i>KCNA1</i>	450
	Episodic ataxia-2	Ca	<i>CACNL1A</i>	
	Spinocerebellar ataxia-6	Ca	<i>CACNL1A</i>	
Migraine	Familial hemiplegic migraine 1	Ca	<i>CACNL1A</i>	447
	Familial hemiplegic migraine 2	Na	<i>SCN1A</i>	
Epilepsy	Benign neonatal familial convulsions	K	<i>KCNQ2, KCNQ3</i>	445
	Generalized epilepsy with febrile convulsions plus	Na	<i>SCN1B</i>	
Periodic paralysis	Hyperkalemic periodic paralysis	Na	<i>SCN4A</i>	462e
	Hypokalemic periodic paralysis	Ca	<i>CACNL1A3</i>	
Myotonia	Myotonia congenita	Cl	<i>CLCN1</i>	462e
	Paramyotonia congenita	Na	<i>SCN4A</i>	
Deafness	Jervell and Lange-Nielsen syndrome (deafness, prolonged QT interval, and arrhythmia)	K	<i>KCNQ1, KCNE1</i>	43
	Autosomal dominant progressive deafness	K	<i>KCNQ4</i>	
Autoimmune				
Paraneoplastic	Limbic encephalitis	Kv1	—	122
	Acquired neuromyotonia	Kv1	—	
	Cerebellar ataxia	Ca (P/Q type)	—	
	Lambert-Eaton syndrome	Ca (P/Q type)	—	

epilepsy syndromes, and recently such mutations have been identified. These mutations appear to alter the normal gating function of these channels, increasing the inherent excitability of neuronal membranes in regions where the abnormal channels are expressed.

Whereas the specific clinical manifestations of channelopathies are quite variable, one common feature is that manifestations tend to be intermittent or paroxysmal, such as occurs in epilepsy, migraine, ataxia, myotonia, or periodic paralysis. Exceptions are clinically progressive channel disorders such as autosomal dominant hearing impairment. The genetic channelopathies identified to date are all uncommon disorders caused by obvious mutations in channel genes. As the full repertoire of human ion channels and related proteins is identified, it is likely that additional channelopathies will be discovered. In addition to rare disorders that result from obvious mutations, it is also likely that less penetrant allelic variations in channel genes or in their pattern of expression might underlie susceptibility to some apparently sporadic forms of epilepsy, migraine, or other disorders. For example, mutations in the potassium channel gene *Kir2.6* have been found in many individuals with thyrotoxic hypokalemic periodic paralysis, a disorder similar to hypokalemic periodic paralysis but precipitated by stress from thyrotoxicosis or carbohydrate loading.

NEUROTRANSMITTERS AND NEUROTRANSMITTER RECEPTORS

Synaptic neurotransmission is the predominant means by which neurons communicate with each other. Classic neurotransmitters are synthesized in the presynaptic region of the nerve terminal; stored in vesicles; and released into the synaptic cleft, where they bind to receptors on the postsynaptic cell. Secreted neurotransmitters are eliminated by reuptake into the presynaptic neuron (or glia), by diffusion away from the synaptic cleft, and/or by specific inactivation. In addition to the classic neurotransmitters, many neuropeptides have been identified as definite or probable neurotransmitters; these include substance P, neurotensin, enkephalins, β -endorphin, histamine, vasoactive intestinal polypeptide, cholecystokinin, neuropeptide Y, and somatostatin.

Peptide neurotransmitters are synthesized in the cell body rather than the nerve terminal and may colocalize with classic neurotransmitters in single neurons. A number of neuropeptides are important in pain modulation including substance P and calcitonin gene-related peptide (CGRP), which causes migraine-like headaches in patients. As a consequence, CGRP receptor antagonists have been developed and shown to be effective in treating migraine headaches. Nitric oxide and carbon monoxide are gases that appear also to function as neurotransmitters, in part by signaling in a retrograde fashion from the postsynaptic to the presynaptic cell.

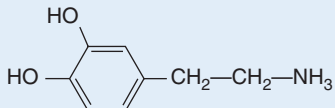
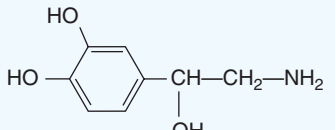
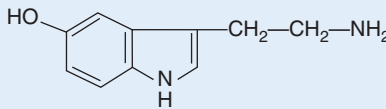
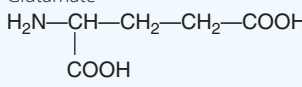
Neurotransmitters modulate the function of postsynaptic cells by binding to specific neurotransmitter receptors, of which there are two major types. *Ionotropic receptors* are direct ion channels that open after engagement by the neurotransmitter. *Metabotropic receptors* interact with G proteins, stimulating production of second messengers and activating protein kinases, which modulate a variety of cellular events. Ionotropic receptors are multiple subunit structures, whereas metabotropic receptors are composed of single subunits only. One important difference between ionotropic and metabotropic receptors is that the

kinetics of ionotropic receptor effects are fast (generally <1 ms) because neurotransmitter binding directly alters the electrical properties of the postsynaptic cell, whereas metabotropic receptors function over longer time periods. These different properties contribute to the potential for selective and finely modulated signaling by neurotransmitters.

Neurotransmitter systems are perturbed in a large number of clinical disorders, several of which are highlighted in [Table 444e-2](#). One example is the involvement of dopaminergic neurons originating in the substantia nigra of the midbrain and projecting to the striatum (nigrostriatal pathway) in Parkinson's disease and in heroin addicts after the ingestion of the toxin MPTP (1-methyl-4-phenyl-1,2,5,6-tetrahydropyridine).

A second important dopaminergic system arising in the midbrain is the mesocorticolimbic pathway, which is implicated in the pathogenesis of addictive behaviors including drug reward. Its key components include the midbrain ventral tegmental area (VTA), median forebrain bundle, and nucleus accumbens (see [Fig. 465e-2](#)). The cholinergic pathway originating in the nucleus basalis of Meynert plays a role in memory function in Alzheimer's disease.

TABLE 444e-2 PRINCIPAL CLASSIC NEUROTRANSMITTERS

Neurotransmitter	Anatomy	Clinical Aspects
Acetylcholine (ACh) $\text{CH}_3-\overset{\text{O}}{\parallel}{\text{C}}-\text{O}-\text{CH}_2-\text{N}-(\text{CH}_3)_3$	Motor neurons in spinal cord → neuromuscular junction Basal forebrain → widespread cortex Interneurons in striatum Autonomic nervous system (preganglionic and postganglionic parasympathetic; preganglionic sympathetic)	Acetylcholinesterases (nerve gases) Myasthenia gravis (antibodies to ACh receptor) Congenital myasthenic syndromes (mutations in ACh receptor subunits) Lambert-Eaton syndrome (antibodies to Ca channels impair ACh release) Botulism (toxin disrupts ACh release by exocytosis) Alzheimer disease (selective cell death) Autosomal dominant frontal lobe epilepsy (mutations in CNS ACh receptor) Parkinson's disease (tremor)
Dopamine 	Substantia nigra → striatum (nigrostriatal pathway) Substantia nigra → limbic system and widespread cortex Arcuate nucleus of hypothalamus → anterior pituitary (via portal veins)	Parkinson's disease (selective cell death) MPTP parkinsonism (toxin transported into neurons) Addiction, behavioral disorders Inhibits prolactin secretion
Norepinephrine (NE) 	Locus coeruleus (pons) → limbic system, hypothalamus, cortex Medulla → locus coeruleus, spinal cord Postganglionic neurons of sympathetic nervous system	Mood disorders (MAOA inhibitors and tricyclics increase NE and improve depression) Anxiety Orthostatic tachycardia syndrome (mutations in NE transporter)
Serotonin 	Pontine raphe nuclei → widespread projections Medulla/pons → dorsal horn of spinal cord	Mood disorders (SSRIs improve depression) Migraine pain pathway Pain pathway
γ-Aminobutyric acid (GABA) $\text{H}_2\text{N}-\text{CH}_2-\text{CH}_2-\text{CH}_2-\text{COOH}$	Major inhibitory neurotransmitter in brain; widespread cortical interneurons and long projection pathways	Stiff person syndrome (antibodies to glutamic acid decarboxylase, the biosynthetic enzyme for GABA) Epilepsy (gabapentin and valproic acid increase GABA)
Glycine $\text{H}_2\text{N}-\text{CH}_2-\text{COOH}$	Major inhibitory neurotransmitter in spinal cord	Spasticity Hyperekplexia (myoclonic startle syndrome) due to mutations in glycine receptor
Glutamate 	Major excitatory neurotransmitter; located throughout CNS, including cortical pyramidal cells	Seizures due to ingestion of domoic acid (a glutamate analogue) Rasmussen's encephalitis (antibody against glutamate receptor 3) Excitotoxic cell death

Abbreviations: CNS, central nervous system; MAOA, monoamine oxidase A; MPTP, 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine; SSRI, selective serotonin reuptake inhibitor.

Addictive drugs share the property of increasing dopamine release in the nucleus accumbens (Chap. 465e). Amphetamine increases intracellular release of dopamine from vesicles and reverses transport of dopamine through the dopamine transporters. Patients prone to addiction show increased activation of the nucleus accumbens following administration of amphetamine. Cocaine binds to dopamine transporters and inhibits dopamine reuptake. Ethanol inhibits inhibitory neurons in the VTA, leading to increased dopamine release in the nucleus accumbens. Opioids also disinhibit these dopaminergic neurons by binding to μ receptors expressed by γ -aminobutyric acid (GABA)-containing interneurons in the VTA. Nicotine increases dopamine release by activating nicotinic acetylcholine receptors on cell bodies and nerve terminals of dopaminergic VTA neurons. Tetrahydrocannabinol, the active ingredient of cannabis, also increases dopamine levels in the nucleus accumbens. Blockade of dopamine in the nucleus accumbens can terminate the rewarding effects of addictive drugs.

Not all cell-to-cell communication in the nervous system occurs via neurotransmission. Gap junctions provide for direct neuron-neuron electrical conduction and also create openings for the diffusion of ions and metabolites between cells. In addition to neurons, gap junctions are also widespread in glia, creating a syncytium that protects neurons by removing glutamate and potassium from the extracellular environment. Gap junctions consist of membrane-spanning proteins, termed *connexins*, that pair across adjacent cells. Mechanisms that involve gap junctions have been related to a variety of neurologic disorders. Mutations in connexin 32, a gap junction protein expressed by Schwann cells, are responsible for the X-linked form of Charcot-Marie-Tooth (CMT) disease (Chap. 459). Mutations in either of two gap junction proteins expressed in the inner ear—connexin 26 and connexin 31—result in autosomal dominant progressive hearing loss (Chap. 43). Glial calcium waves mediated through gap junctions also appear to explain the phenomenon of spreading depression associated with migraine auras and the march of epileptic discharges. Spreading depression is a neural response that follows a variety of different stimuli and is characterized by a circumferentially expanding negative potential that propagates at a characteristic speed of 20 m/s and is associated with an increase in extracellular potassium.

SIGNALING PATHWAYS AND GENE TRANSCRIPTION

The fundamental issue of how memory, learning, and thinking are encoded in the nervous system is likely to be clarified by identifying the signaling pathways involved in neuronal differentiation, axon guidance, and synapse formation, and by understanding how these pathways are modulated by experience. Many families of transcription factors, each comprising multiple individual components, are expressed in the nervous system. Elucidation of these signaling pathways has already begun to provide insights into the cause of a variety of neurologic disorders, including inherited disorders of cognition such as X-linked mental retardation. This problem affects ~1 in 500 males, and linkage studies in different families suggest that as many as 60 different X-chromosome-encoded genes may be responsible. The formation of RNA-DNA duplexes that block transcription has also been observed with the CGG repeat expansions that occur in fragile X gene-associated mental retardation. Rett's syndrome, a common cause of (dominant) X-linked progressive mental retardation in females, is due to a mutation in a gene (*MECP2*) encoding a DNA-binding protein involved in transcriptional repression. Because the X chromosome comprises only ~3% of germline DNA, then by extrapolation, the number of genes that potentially contribute to clinical disorders affecting intelligence in humans must be potentially very large. As discussed below, there is increasing evidence that abnormal gene transcription may play a role in neurodegenerative diseases, such as Huntington's disease, in which proteins with polyglutamine expansions bind to and sequester transcription factors. A critical transcription factor for neuronal survival is CREB (cyclic adenosine monophosphate responsive element-binding) protein, which also plays an important role in memory in the hippocampus. The regulatory gene repressor element 1-silencing transcription factor (REST)

coordinates the expression of neuroprotective stress genes during normal aging. It turns off genes involved in cell death and pathology and boosts protective factors. High levels of REST are associated with normal cognition even in the presence of both amyloid plaques and neurofibrillary tangles. Although REST increases with normal aging, it fails to increase in the nucleus in patients with Alzheimer's disease and is found clumped with amyloid in autophagosomes.

MYELIN AND NEUROINFLAMMATION

Myelin is the multilayered insulating substance that surrounds axons and speeds impulse conduction by permitting action potentials to jump between naked regions of axons (nodes of Ranvier) and across myelinated segments. Molecular interactions between the myelin membrane and axon are required to maintain the stability, function, and normal life span of both structures. A single oligodendrocyte usually ensheathes multiple axons in the central nervous system (CNS), whereas in the peripheral nervous system (PNS), each Schwann cell typically myelinates a single axon. Myelin is a lipid-rich material formed by a spiraling process of the membrane of the myelinating cell around the axon, creating multiple membrane bilayers that are tightly apposed (compact myelin) by charged protein interactions. Several inhibitors of axon growth are expressed on the innermost (periaxonal) lamellae of the myelin membrane (see "Stem Cells and Transplantation," below). A number of clinically important neurologic disorders are caused by inherited mutations in myelin proteins of the CNS or PNS (Fig. 444e-1). Constituents of myelin also have a

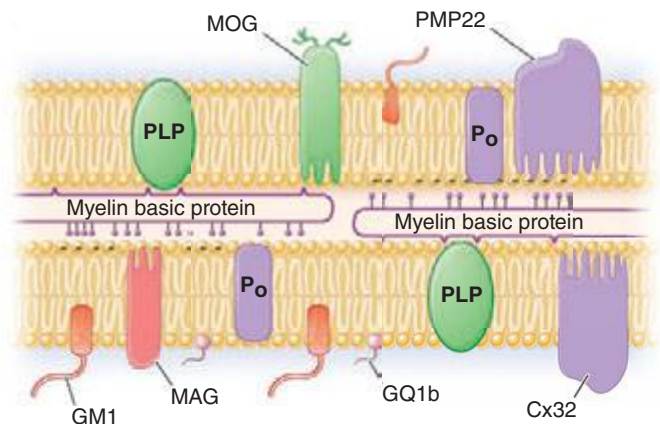


FIGURE 444e-1 The molecular architecture of the myelin sheath illustrating the most important disease-related proteins. The illustration represents a composite of central nervous system (CNS) and peripheral nervous system (PNS) myelin. Proteins restricted to CNS myelin are shown in green, proteins of PNS myelin are lavender, and proteins present in both CNS and PNS are red. In the CNS, the X-linked allelic disorders, Pelizaeus-Merzbacher disease and one variant of familial spastic paraplegia, are caused by mutations in the gene for proteolipid protein (PLP) that normally promotes extracellular compaction between adjacent myelin lamellae. The homologue of PLP in the PNS is the P_0 protein, mutations in which cause the neuropathy Charcot-Marie-Tooth disease (CMT) type 1B. The most common form of CMT is the 1A subtype caused by a duplication of the *PMP22* gene; deletions in *PMP22* are responsible for another inherited neuropathy termed *hereditary liability to pressure palsies* (Chap. 459).

In multiple sclerosis (MS), myelin basic protein (MBP) and the quantitatively minor CNS protein, myelin oligodendrocyte glycoprotein (MOG), are likely T cell and B cell antigens, respectively (Chap. 458). The location of MOG at the outermost lamella of the CNS myelin membrane may facilitate its targeting by autoantibodies. In the PNS, autoantibodies against myelin gangliosides are implicated in a variety of disorders, including GQ1b in the Fisher variant of Guillain-Barré syndrome, GM1 in multifocal motor neuropathy, and sulfatide constituents of myelin-associated glycoprotein (MAG) in peripheral neuropathies associated with monoclonal gammopathies (Chap. 460).

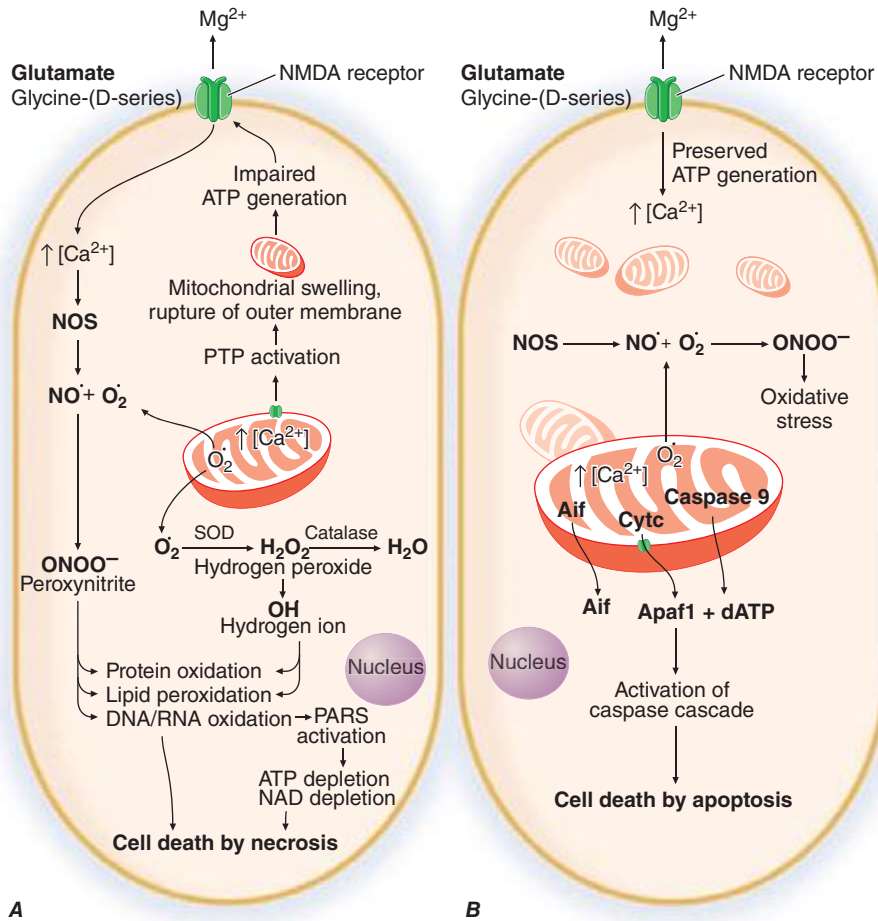


FIGURE 444e-2 Involvement of mitochondria in cell death. A severe excitotoxic insult (**A**) results in cell death by necrosis, whereas a mild excitotoxic insult (**B**) results in apoptosis. After a severe insult (such as ischemia), there is a large increase in glutamate activation of *N*-methyl-D-aspartate (NMDA) receptors, an increase in intracellular Ca^{2+} concentrations, activation of nitric oxide synthase (NOS), and increased mitochondrial Ca^{2+} and superoxide generation followed by the formation of $ONOO^-$. This sequence results in damage to cellular macromolecules including DNA, leading to activation of poly-ADP-ribose polymerase (PARS). Both mitochondrial accumulation of Ca^{2+} and oxidative damage lead to activation of the permeability transition pore (PTP) that is linked to excitotoxic cell death. A mild excitotoxic insult can occur due either to an abnormality in an excitotoxicity amino acid receptor, allowing more Ca^{2+} flux, or to impaired functioning of other ionic channels or of energy production, which may allow the voltage-dependent NMDA receptor to be activated by ambient concentrations of glutamate. This event can then lead to increased mitochondrial Ca^{2+} and free radical production, yet relatively preserved ATP generation. The mitochondria may then release cytochrome c (Cyt c), caspase 9, apoptosis-inducing factor (Aif), and perhaps other mediators that lead to apoptosis. The precise role of the PTP in this mode of cell death is still being clarified, but there does appear to be involvement of the adenine nucleotide transporter that is a key component of the PTP.

propensity to be targeted as autoantigens in autoimmune demyelinating disorders (Fig. 444e-2).

Specification to oligodendrocyte precursor cells (OPCs) is transcriptionally regulated by the *Olig 2* and *Yin Yang 1* genes, whereas myelination mediated by postmitotic oligodendrocytes depends on a different transcription factor, *myelin gene regulatory factor (MRF)*. It is noteworthy that in the normal adult brain, large numbers of OPCs (expressing PDGFR- α and NG2) are widely distributed but do not myelinate axons, even in demyelinating environments such as in lesions of MS. Several families of molecules have now been identified that regulate oligodendrocyte differentiation and myelination, including LINGO-1, PSA-NCAM, hyaluronan, Nogo-A, the Wnt pathway, notch signaling (and its receptor Jagged), and the retinoic acid receptor RXR γ ; all are inhibitory, with the exception of RXR γ , which is excitatory. All are also potential targets for myelin repair therapies, and a monoclonal antibody against LINGO-1 is in clinical testing for remyelination in MS. Very recently, a series of observations has called into question the traditional concept that axon-derived cues are always required for myelination to occur. Fixed (i.e., dead) axons could be efficiently myelinated by oligodendrocytes in vitro, as could artificial polystyrene nanowires of a similar diameter. This led to development of new

high-throughput screening assays based on myelination of polystyrene nanowires to identify compounds that could promote myelination.

Macrophages and microglia represent the major cell types in the nervous system responsible for antigen presentation and innate immunity. Brain macrophages are derived from either hematopoietic stem cell-derived bone marrow monocytes or from brain microglia that migrate from the yolk sac early in embryogenesis before the blood-brain barrier is formed. In a murine model of autoimmune demyelination, experimental allergic encephalomyelitis (EAE) (Fig. 444e-3), macrophages derived from bone marrow monocytes, but not microglia, were found to represent the critical population that initiated inflammatory demyelination at paraxonal regions near nodes of Ranvier. An additional, unexpected role for brain microglia was also identified in the regulation of neural circuits through pruning of excitatory synapses and control of dendritic spine densities; mice depleted of microglia during development exhibited a variety of cognitive learning and behavioral deficits, including abnormal social behaviors. Remarkably, depletion of microglia in adult mice by administration of a selective inhibitor of colony-stimulating factor receptor 1 (CSFR1) was followed by their rapid repopulation, suggesting that a pool of resident microglial precursor cells may exist throughout the CNS.

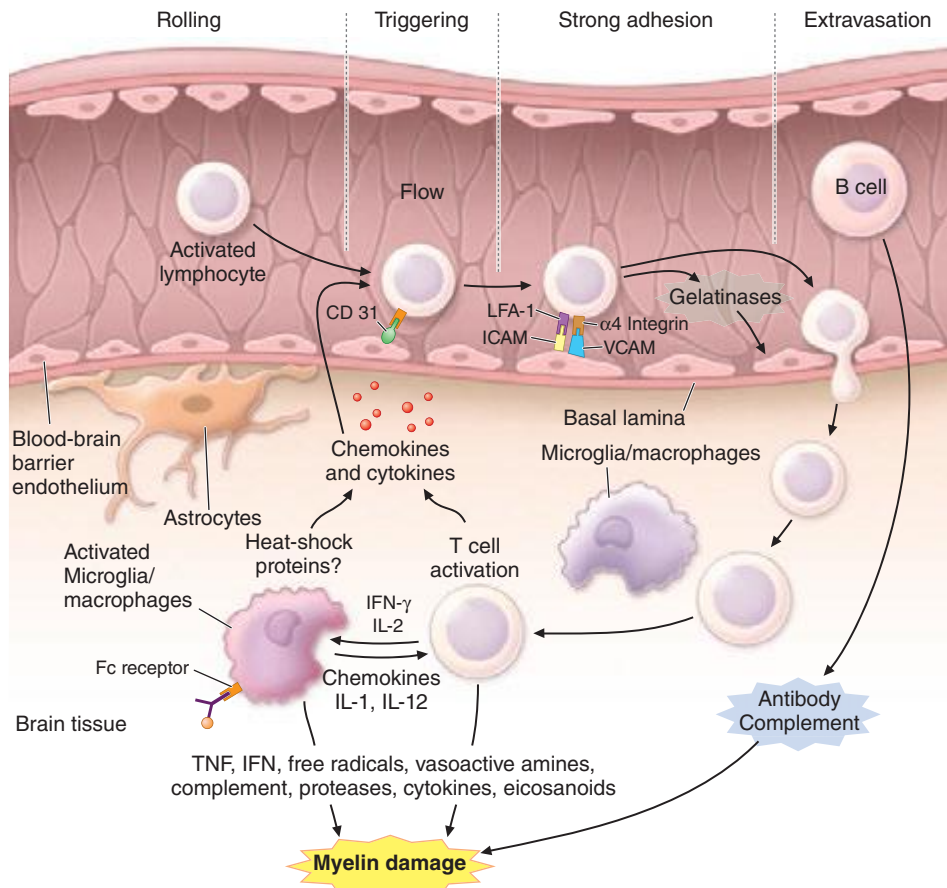


FIGURE 444e-3 A model for experimental allergic encephalomyelitis (EAE). Crucial steps for disease initiation and progression include peripheral activation of preexisting autoreactive T cells; homing to the central nervous system (CNS) and extravasation across the blood-brain barrier; reactivation of T cells by exposed autoantigens; secretion of cytokines; activation of microglia and astrocytes and recruitment of a secondary inflammatory wave; and immune-mediated myelin destruction. ICAM, intercellular adhesion molecule; IFN, interferon; IL, interleukin; LFA-1, leukocyte function-associated antigen-1; TNF, tumor necrosis factor; VCAM, vascular cell adhesion molecule.

MICROBIOTA AND NEUROLOGIC DISEASE

The human microbiome (Chap. 86e) represents the collective set of genes from the 10^{14} organisms living in our gut, skin, mucosa, and other sites. Different microbial communities are associated with different ethnicities, diets, and environments. In any individual, the predominant gut microbiota can be remarkably stable over decades, but also can be altered by exposure to certain microbial species, for example by ingestion of probiotics.

There is compelling evidence that gut microbes can shape immune responses through the interaction of their metabolism with that of humans. These gut-brain interactions are likely to be important in understanding the pathogenesis of many autoimmune neurologic diseases. For example, mice treated with broad-spectrum antibiotics are resistant to EAE, an effect associated with decreases in production of proinflammatory cytokines, and conversely more production of the immunosuppressive cytokines interleukin (IL) 10 and IL-13 and an increase in regulatory T and B lymphocytes. Oral administration of polysaccharide A (PSA) from *Bacillus fragilis* also protects mice from EAE, via increases in IL-10.

In addition to nonspecific effects on immune homeostasis mediated by cytokines and regulatory cells, some microbial proteins can trigger, in susceptible individuals, a cross-reactive immune response against a homologous protein in the nervous system, a mechanism termed *molecular mimicry*. Examples include cross-reactivity between the astrocyte water channel aquaporin-4 and an ABC transporter permease from *Clostridia perfringens* in neuromyelitis optica (Chap. 458); the neural ganglioside Gm1 and similar sialic acid-containing structures from *Campylobacter jejuni* in Guillain-Barré syndrome (Chap. 460); and the sleep-promoting protein hypocretin and hemagglutinin from H1N1 influenza virus in narcolepsy (Chap. 38), among others.

Recently, a number of tantalizing observations have incriminated the microbial environment in the pathogenesis of a much wider spectrum of neurologic conditions and behaviors, extending well beyond the traditional boundaries of immune-mediated pathologies. This is perhaps not surprising, as it has long been known in neurology that gut bacteria can influence brain function, based mostly on classic studies demonstrating that products of gut microbes can worsen hepatic encephalopathy, forming the basis of treatment with antibiotics for this condition.

Mice that developed in a completely germ-free environment displayed less anxiety, lower responses to stressful situations, more exploratory locomotive behaviors, and impaired memory formation compared with non-germ-free counterparts. These behaviors were related to changes in gene expression in pathways related to neural signaling, synaptic function, and modulation of neurotransmitters. Moreover, this behavior could be reversed when the germ-free mice were co-housed with non-germ-free mice.

The enteric autonomic nervous system in humans provides a bidirectional neural connection between the brain and gut. The vagus nerve, which innervates the upper gut and proximal colon, has been implicated in anxiety- and depression-like behaviors in mice. Ingestion of *Lactobacillus rhamnosus* induced changes in expression of the inhibitory neurotransmitter GABA1b in neurons of the limbic cortex, hippocampus, and amygdala, associated with reduced levels of corticosteroids and reduced anxiety- and depression-like behaviors. Remarkably, these changes could be blocked by vagotomy.

Another area of emerging interest is in a possible contribution of the gut microbiome to autism and related disorders. Children with autistic spectrum disorders have long been known to have gastrointestinal disturbances, and it has been claimed that the severity of dysbiosis correlates

TABLE 444e-3 NEUROTROPHIC FACTORS

Neurotrophin family	Transforming growth factor β family
Nerve growth factor	Glial-derived neurotrophic family
Brain-derived neurotrophic factor	Neurturin
Neurotrophin-3	Persephin
Neurotrophin-4	Fibroblast growth factor family
Neurotrophin-6	Hepatocyte growth factor
Cytokine family	Insulin-like growth factor (IGF) family
Ciliary neurotrophic factor	IGF-1
Leukemia inhibitory factor	IGF-2
Interleukin 6	
Cardiotrophin-1	

with the severity of autism. A murine model of autism was recently induced in offspring after injecting the pregnant mother with the viral RNA mimic polyinosinic:polycytidylic acid (poly I:C). Remarkably, oral treatment of offspring with *B. fragilis* corrected a range of autistic behaviors in these mice and also improved gut permeability.

NEUROTROPHIC FACTORS

Neurotrophic factors (Table 444e-3) are secreted proteins that modulate neuronal growth, differentiation, repair, and survival; some have additional functions, including roles in neurotransmission and in the synaptic reorganization involved in learning and memory. The neurotrophin (NT) family contains nerve growth factor (NGF), brain-derived neurotrophic factor (BDNF), NT3, and NT4/5. The neurotrophins act at TrK and p75 receptors to promote survival of neurons. BDNF is linked to synaptogenesis. Certain polymorphisms are linked to increased risk for Alzheimer's disease (AD), and BDNF is depleted in Huntington's disease. Because of their survival-promoting and antiapoptotic effects, neurotrophic factors are in theory outstanding candidates for therapy of disorders characterized by premature death of neurons such as occurs in ALS and other degenerative motor neuron disorders. Knockout mice lacking receptors for ciliary neurotrophic factor (CNTF) or BDNF show loss of motor neurons, and experimental motor neuron death can be rescued by treatment with various neurotrophic factors including CNTF, BDNF, and vascular endothelial growth factor (VEGF). However, in phase 3 clinical trials, growth factors were ineffective in human ALS. The growth factor glial-derived neurotrophic factor (GDNF) is important for survival of dopaminergic neurons. Direct infusions of GDNF showed initial promise in Parkinson's disease, but the benefits were not replicated in a larger clinical trial.

STEM CELLS AND TRANSPLANTATION

The nervous system is traditionally considered to be a nonmitotic organ, in particular with respect to neurons. These concepts have been challenged by the finding that neural progenitor or stem cells exist in the adult CNS that are capable of differentiation, migration over long distances, and extensive axonal arborization and synapse formation with appropriate targets. These capabilities also indicate that the repertoire of factors required for growth, survival, differentiation, and migration of these cells exists in the mature nervous system. In rodents, neural stem cells, defined as progenitor cells capable of differentiating into mature cells of neural or glial lineage, have been experimentally propagated from fetal CNS and neuroectodermal tissues and also from adult germinal matrix and ependyma regions. Human fetal CNS tissue is also capable of differentiation into cells with neuronal, astrocyte, and oligodendrocyte morphology when cultured in the presence of growth factors.

Once the repertoire of signals required for cell type specification is better understood, differentiation into specific neural or glial subpopulations can be directed in vitro; such cells could also be engineered to express therapeutic molecules. Another promising approach is to use

growth factors, such as BDNF, to stimulate endogenous stem cells to proliferate and migrate to areas of neuronal damage.

A major advance has been the development of induced pluripotent stem cells. Using this technique, adult somatic cells such as skin fibroblasts are treated with four pluripotency factors (SOX2, KLF4, cMYC, and Oct4), and this generates induced pluripotent stem cells (iPSCs). These adult-derived stem cells sidestep the ethical issues of using stem cells derived from human embryos. The development of these cells has tremendous promise for both studying disease mechanisms and testing therapeutics. As yet there is no consensus on the best way to generate and differentiate the iPSCs; however, techniques to avoid using viral vectors and use of Cre-lox systems to remove reprogramming factors result in a better match of gene expression profiles with those of embryonic stem cells. Over the years, the field of directed differentiation has used three main strategies to specify neural lineages from human pluripotent stem cells. These strategies are embryoid body formation, coculture on neural-inducing feeders, and direct neural induction. Thus far, iPSCs have been made from patients with all of the major human neurodegenerative diseases, and studies using them are under way.

Although stem cells hold tremendous promise for the treatment of debilitating neurologic diseases, such as Parkinson's disease and spinal cord injury, it should be emphasized that medical application is in its infancy. Major obstacles are the generation of position- and neurotransmitter-defined subtypes of neurons and their isolation as pure populations of the desired cells. This is crucial to avoid persistence of undifferentiated embryonic stem (ES) cells, which can generate tumors. The establishment of appropriate neural connections and afferent control is also critical. For instance, human ES motor neurons will need to be introduced at multiple segments in the neuraxis, and then their axons will need to regenerate from the spinal cord to distal musculature.

Experimental transplantation of human fetal dopaminergic neurons in patients with Parkinson's disease has shown that these transplanted cells can survive within the host striatum; however, some patients developed disabling dyskinesias, and this approach is no longer in clinical development. The possibility that iPSCs will be used in Parkinson's disease was strengthened by studies showing that they can be differentiated into dopaminergic neurons. The dopaminergic neurons were then shown to rescue the parkinsonian phenotype in a MPTP-induced primate model with excellent dopaminergic neuron survival function and lack of neural overgrowth. The correction of tau mutations in iPSC-derived neurons has been shown to reverse the toxic phenotype in dendrite retraction and cell death.

Another new use for iPSCs is to screen drugs as potential treatments for neurodegenerative and other diseases. The feasibility of this has been shown using iPSC-induced macrophages from patients with Gaucher's disease, and verifying the efficacy of protein chaperones in these cells as a means of stabilizing the mutant glucocerebrosidase and increasing its activity and the duration of its effects. Other approaches are to attempt to reduce expression of proteins, such as amyloid, tau, and α -synuclein, implicated in the pathogenesis of neurodegenerative diseases. One difficulty has been that reprogramming cells to iPSCs resets their identity back to an embryonic age, which is a hurdle in modeling of late-onset diseases. One approach to this has been to express a fragment of the mutated gene, such as a portion of lamin A, which causes premature aging in progeria. This approach showed that dendrite degeneration and progressive loss of tyrosine hydroxylase expression, as well as enlarged mitochondria and Lewy body precursor inclusions, were induced in iPSC-derived dopaminergic neurons with progerin-induced aging.

Studies of transplantation for patients with Huntington's disease have also reported encouraging, although very preliminary, results. OPCs transplanted into mice with a dysmyelinating disorder effectively migrated in the new environment, interacted with axons, and mediated myelination; such experiments raise hope that similar transplantation strategies may be feasible in human disorders of myelin such as MS. The promise of stem cells for treatment of both neurodegenerative diseases and neural injury is great, but development has

been slowed by unresolved concerns over safety (including the theoretical risk of malignant transformation of transplanted cells), ethics (particularly with respect to use of fetal tissue), and efficacy.

In developing brain, the extracellular matrix provides stimulatory and inhibitory signals that promote neuronal migration, neurite outgrowth, and axonal extension. After neuronal damage, reexpression of inhibitory molecules such as chondroitin sulfate proteoglycans may prevent tissue regeneration. Chondroitinase degraded these inhibitory molecules and enhanced axonal regeneration and motor recovery in a rat model of spinal cord injury. Several myelin proteins, specifically Nogo, oligodendrocyte myelin glycoprotein (OMGP), and myelin-associated glycoprotein (MAG), may also interfere with axon regeneration. Sialidase, which cleaves one class of receptors for MAG, enhances axonal outgrowth. Antibodies against Nogo promote regeneration after experimental focal ischemia or spinal cord injury. Nogo, OMGP, and MAG all bind to the same neural receptor, the Nogo receptor, which mediates its inhibitory function via the p75 neurotrophin receptor signaling.

CELL DEATH: EXCITOTOXICITY AND APOPTOSIS

Excitotoxicity refers to neuronal cell death caused by activation of excitatory amino acid receptors (Fig. 444e-4). Compelling evidence for a role of excitotoxicity, especially in ischemic neuronal injury, is derived from experiments in animal models. Experimental models of stroke are associated with increased extracellular concentrations of the excitatory amino acid neurotransmitter glutamate, and neuronal damage is attenuated by denervation of glutamate-containing neurons or

the administration of glutamate receptor antagonists. The distribution of cells sensitive to ischemia corresponds closely with that of *N*-methyl-D-aspartate (NMDA) receptors (except for cerebellar Purkinje cells, which are vulnerable to hypoxia-ischemia but lack NMDA receptors); and competitive and noncompetitive NMDA antagonists are effective in preventing focal ischemia. In global cerebral ischemia, non-NMDA receptors (kainic acid and α -amino-3-hydroxy-5-methyl-4-isoxazole-propionate [AMPA]) are activated, and antagonists to these receptors are protective. Experimental brain damage induced by hypoglycemia is also attenuated by NMDA antagonists.

Excitotoxicity is not a single event but rather a cascade of cell injury. Excitotoxicity causes influx of calcium into cells, and much of the calcium is sequestered in mitochondria rather than in the cytoplasm. Increased cytoplasmic calcium causes metabolic dysfunction and free radical generation; activates protein kinases, phospholipases, nitric oxide synthase, proteases, and endonucleases; and inhibits protein synthesis. Activation of nitric oxide synthase generates nitric oxide (NO \bullet), which can react with superoxide (O \bullet_2) to generate peroxynitrite (ONOO $^-$), which may play a direct role in neuronal injury. Another critical pathway is activation of poly-ADP-ribose polymerase, which occurs in response to free radical-mediated DNA damage. Experimentally, mice with knockout mutations of neuronal nitric oxide synthase or poly-ADP-ribose polymerase, or those that overexpress superoxide dismutase, are resistant to focal ischemia.

Another aspect of excitotoxicity is that it has been demonstrated that stimulation of extrasynaptic NMDA receptors mediates cell death, where as stimulation of synaptic receptors is protective. This has been shown to play a role in excitotoxicity in transgenic mouse models of

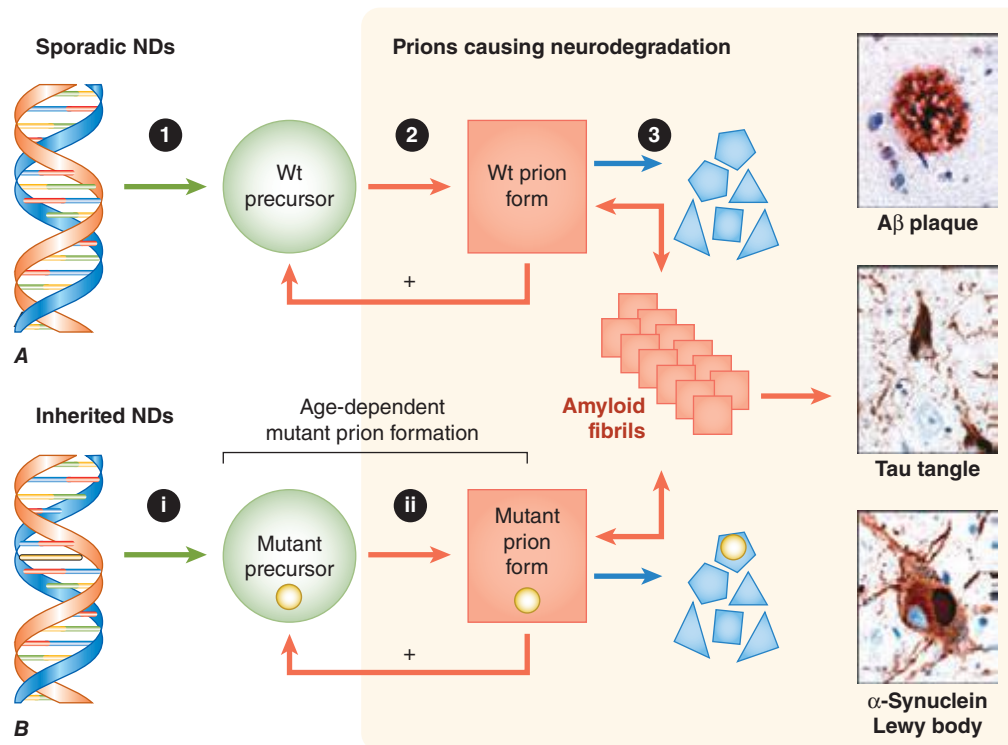


FIGURE 444e-4 Neurodegeneration caused by prions. A. In sporadic neurodegenerative diseases (NDs), wild-type (Wt) prions multiply through self-propagating cycles of posttranslational modification, during which the precursor protein (*green circle*) is converted into the prion form (*red square*), which generally is high in β -sheet content. Pathogenic prions are most toxic as oligomers and less toxic after polymerization into amyloid fibrils. The small polygons (*blue*) represent proteolytic cleavage products of the prion. Depending on the protein, the fibrils coalesce into A β amyloid plaques in Alzheimer's disease (AD), neurofibrillary tangles in AD and Pick's disease, or Lewy bodies in Parkinson's disease and Lewy body dementia. Drug targets for the development of therapeutics include: (1) lowering the precursor protein, (2) inhibiting prion formation, and (3) enhancing prion clearance. **B.** Late-onset heritable neurodegeneration argues for two discrete events: The (*i*) first event is the synthesis of mutant precursor protein (*green circle*), and the (*ii*) second event is the age-dependent formation of mutant prions (*red square*). The *highlighted yellow bar* in the DNA structure represents mutation of a base pair within an exon, and the *small yellow circles* signify the corresponding mutant amino acid substitution. *Green arrows* represent a normal process; *red arrows*, a pathogenic process; and *blue arrows*, a process that is known to occur but unknown whether it is normal or pathogenic. (Micrographs prepared by Stephen J. DeArmond. Reprinted with permission from SB Prusiner: *Biology and genetics of prions causing neurodegeneration. Annu Rev Genet* 47:601, 2013.)

Huntington's disease, in which using low-dose memantine to selectively block the extrasynaptic receptors is beneficial.

Although excitotoxicity is clearly implicated in the pathogenesis of cell death in stroke, to date treatment with NMDA antagonists has not proven to be clinically useful. One approach has been to use an inhibitor of the postsynaptic density-95 protein that uncouples NMDA receptors from neurotoxic pathways, including the generation of nitric oxide. This approach was effective in a primate stroke model and in a phase 2 clinical trial of stroke associated with endovascular repair of cerebral aneurysms. Transient receptor potentials (TRPs) are calcium channels that are activated by oxidative stress in parallel with excitotoxic signal pathways. In addition, glutamate-independent pathways of calcium influx via acid-sensing ion channels have been identified. These channels transport calcium in the setting of acidosis and substrate depletion, and pharmacologic blockade of these channels markedly attenuates stroke injury. These channels offer a potential new therapeutic target for stroke.

Apoptosis, or programmed cell death, plays an important role in both physiologic and pathologic conditions. During embryogenesis, apoptotic pathways operate to destroy neurons that fail to differentiate appropriately or reach their intended targets. There is mounting evidence for an increased rate of apoptotic cell death in a variety of acute and chronic neurologic diseases. Apoptosis is characterized by neuronal shrinkage, chromatin condensation, and DNA fragmentation, whereas necrotic cell death is associated with cytoplasmic and mitochondrial swelling followed by dissolution of the cell membrane. Apoptotic and necrotic cell death can coexist or be sequential events, depending on the severity of the initiating insult. Cellular energy reserves appear to have an important role in these two forms of cell death, with apoptosis favored under conditions in which ATP levels are preserved. Evidence of DNA fragmentation has been found in a number of degenerative neurologic disorders, including AD, Huntington's disease, and ALS. The best characterized genetic neurologic disorder related to apoptosis is infantile spinal muscular atrophy (Werdnig-Hoffmann disease), in which two genes thought to be involved in the apoptosis pathways are causative.

Mitochondria are essential in controlling specific apoptosis pathways. The redistribution of cytochrome *c*, as well as apoptosis-inducing factor (AIF), from mitochondria during apoptosis leads to the activation of a cascade of intracellular proteases known as *caspases*. Caspase-independent apoptosis occurs after DNA damage, activation of poly-ADP-ribose polymerase, and translocation of AIF into the nucleus. Redistribution of cytochrome *c* is prevented by overproduction of the apoptotic protein BCL2 and is promoted by the proapoptotic protein BAX. These pathways may be triggered by activation of a large pore in the mitochondrial inner membrane known as the *permeability transition pore*, although in other circumstances, they occur independently. The permeability transition pore is made up of dimers of ATP synthase and is activated by cyclophilin D, leading to large calcium fluxes across the inner mitochondrial membrane. Certain forms of congenital muscular dystrophies are caused by mutations in collagen VI, which leads to increased activation of the permeability transition pore. Recent studies suggest that blocking the mitochondrial pore reduces both hypoglycemic and ischemic cell death. Mice deficient in cyclophilin D, a key protein involved in opening the permeability transition pore, are resistant to necrosis produced by focal cerebral ischemia.

PROTEIN AGGREGATION AND NEURODEGENERATION

The possibility that protein aggregation plays a role in the pathogenesis of neurodegenerative diseases is a major focus of current research. Protein aggregation is a major histopathologic hallmark of neurodegenerative diseases. Deposition of β -amyloid is strongly implicated in the pathogenesis of AD. Genetic mutations in familial AD cause increased production of β -amyloid with 42 amino acids, which has an increased propensity to aggregate, as compared to β -amyloid with 40 amino acids. Furthermore, mutations in the amyloid precursor protein (APP), which reduce the production of β -amyloid, protect against the development of AD and are associated with preserved

cognition in the elderly. Mutations in genes encoding the MAPT lead to altered splicing of tau and the production of neurofibrillary tangles in frontotemporal dementia and progressive supranuclear palsy. Familial Parkinson's disease is associated with mutations in *leucine-rich repeat kinase 2 (LRRK2)*, *α -synuclein*, *parkin*, *PINK1*, and *DJ-1*. *PINK1* is a mitochondrial kinase (see below), and *DJ-1* is a protein involved in protection from oxidative stress. *Parkin*, which causes autosomal recessive early-onset Parkinson's disease, is a ubiquitin ligase. The characteristic histopathologic feature of Parkinson's disease is the Lewy body, an eosinophilic cytoplasmic inclusion that contains both neurofilaments and α -synuclein. Huntington's disease and cerebellar degenerations are associated with expansions of polyglutamine repeats in proteins, which aggregate to produce neuronal intranuclear inclusions. Familial ALS is associated with superoxide dismutase mutations and cytoplasmic inclusions containing superoxide dismutase. An important finding was the discovery that ubiquitinated inclusions observed in most cases of ALS and the most common form of frontotemporal dementia are composed of TAR DNA binding protein 43 (TDP-43). Subsequently, mutations in the TDP-43 gene, and in the fused in sarcoma gene (*FUS*), were found in familial ALS. Both of these proteins are involved in transcription regulation as well as RNA metabolism. In autosomal dominant neurohypophyseal diabetes insipidus, mutations in vasopressin result in abnormal protein processing, accumulation in the endoplasmic reticulum, and cell death.

Another key mechanism linked to cell death is mitochondrial dynamics, which refers to the processes involved in movement of mitochondria, as well as in mitochondrial fission and fusion, which play a critical role mitochondrial turnover and in replenishment of damaged mitochondria. Mitochondrial dysfunction is strongly linked to the pathogenesis of a number of neurodegenerative diseases such as Friedreich's ataxia, which is caused by mutations in an iron-binding protein that plays an important role in transferring iron to iron-sulfur clusters in aconitase and complex I and II of the electron transport chain. Mitochondrial fission is dependent on the dynamin-related proteins (*Drp1*), which bind to its receptor *Fis*, whereas mitofuscins 1 and 2 (*MF1/2*) and optic atrophy protein 1 (*OPA1*) are responsible for fusion of the outer and inner mitochondrial membrane, respectively. Mutations in *MFN2* cause Charcot-Marie-Tooth neuropathy type 2A, and mutations in *OPA1* cause autosomal dominant optic atrophy. Both β -amyloid and mutant huntingtin protein induce mitochondrial fragmentation and neuronal cell death associated with increased activity of *Drp1*. In addition, mutations in genes causing autosomal recessive Parkinson's disease, *parkin* and *PINK1*, cause abnormal mitochondrial morphology and result in impairment of the ability of the cell to remove damaged mitochondria by autophagy.

One major scientific question is whether protein aggregates directly contribute to neuronal death or whether they are merely secondary bystanders. A current focus in all the neurodegenerative diseases is on small protein aggregates termed *oligomers*. These may be the toxic species of β -amyloid, α -synuclein, and proteins with expanded polyglutamines such as are associated with Huntington's disease. Protein aggregates are usually ubiquitinated, which targets them for degradation by the 26S component of the proteasome. An inability to degrade protein aggregates could lead to cellular dysfunction, impaired axonal transport, and cell death by apoptotic mechanisms.

Autophagy is the degradation of cytosolic components in lysosomes. There is increasing evidence that autophagy plays an important role in degradation of protein aggregates in the neurodegenerative diseases, and it is impaired in AD, Parkinson's disease, and Huntington's disease. Autophagy is particularly important to the health of neurons, and failure of autophagy contributes to cell death. In Huntington's disease, a failure of cargo recognition occurs, contributing to protein aggregates and cell death. Rapamycin, which induces autophagy, exerts beneficial therapeutic effects in transgenic mouse models of AD, Parkinson's disease, and Huntington's disease.

There is other evidence for lysosomal dysfunction and impaired autophagy in Parkinson's disease. Mutations in glucocerebrosidase are associated with 5% of all Parkinson's disease cases as well as 8–9% of patients with dementia with Lewy bodies. Therefore, this is the most

444e-10 important genetic cause of both disorders thus far identified. There appear to be reciprocal interactions between glucocerebrosidase and α -synuclein. It has been shown that glucocerebrosidase concentrations and enzymatic activity are reduced in the substantia nigra of sporadic Parkinson's disease patients. Furthermore, α -synuclein is degraded by chaperone-mediated and macro autophagy. The degradation of α -synuclein has been shown to be impaired in transgenic mice deficient in glucocerebrosidase as well as in mice in which the enzyme has been inhibited. Furthermore, it is known that α -synuclein inhibits the activity of glucocerebrosidase. Therefore, there is bidirectional feedback between α -synuclein and glucocerebrosidase. An attractive therapeutic intervention could be to use protein chaperones to increase the activity and duration of action of glucocerebrosidase. This would also reduce α -synuclein levels and block the degeneration of dopaminergic neurons.

The retromer complex is a conserved membrane-associated protein complex that functions in the endosome-to-Golgi complex. The retromer complex contains a cargo selective complex consisting of VPS35, VPS26, and VPS29, along with a sorting nexin dimer. Recently, mutations in VPS35 were shown to be a cause of late-onset autosomal dominant Parkinson's disease. The retromer also traffics APP away from endosomes, where it is cleaved to generate β -amyloid. Deficiencies of VPS35 and VPS26 were also identified in hippocampal brain tissue from AD. A new therapeutic approach to these diseases might therefore be to use chaperones to stabilize the retromer and reduce the generation of β -amyloid and α -synuclein.

The LRRK2 mutations were shown to have effects on clearance of Golgi-derived vesicles through the autophagy-lysosome system both in vitro and in vivo. LRRK2 mutations also are linked to elevated protein synthesis mediated by ribosomal protein s15 phosphorylation. Blocking this phosphorylation reduces LRRK2-mediated neurite loss and cell death in human dopamine and cortical neurons.

Interestingly, in experimental models of Huntington's disease and cerebellar degeneration, protein aggregates are not well correlated with neuronal death and may be protective. A substantial body of evidence suggests that the mutant proteins with polyglutamine expansions in these diseases bind to transcription factors and that this contributes to disease pathogenesis. In Huntington's disease, there is dysfunction of the transcriptional co-regulator, PGC-1 α , a key regulator of mitochondrial biogenesis. There is evidence that impaired function of PGC-1 α is also important in both Parkinson's disease and AD, making it an attractive target for treatments. Agents that upregulate gene transcription are neuroprotective in animal models of these diseases. A number of compounds have been developed to block β -amyloid production and/or aggregation, and these agents are being studied in early clinical trials in humans. Another approach under investigation is immunotherapy with antibodies that bind β -amyloid, tau, or α -synuclein. These studies have shown efficacy in preventing the spread of amyloid, tau, and α -synuclein in animal studies, raising hopes that this could lead to effective therapies by blocking neuron-to-neuron propagation. Two large clinical trials of β -amyloid immunotherapy, however, did not show efficacy, although this therapeutic strategy is still being studied.

PRIONS AND NEURODEGENERATIVE DISEASES

As we have learned more about the etiology and pathogenesis of the neurodegenerative diseases, it has become clear that the histologic abnormalities that were once curiosities, in fact, are likely to reflect the etiologies. For example, the amyloid plaques in kuru and Creutzfeldt-Jakob disease (CJD) are filled with the PrP^{Sc} prions that have assembled into fibrils. The past three decades have witnessed an explosion of new knowledge about prions. For many years, kuru, CJD, and scrapie of sheep were thought to be caused by slow-acting viruses, but a large body of experimental evidence argues that the infectious pathogens causing these diseases are devoid of nucleic acid. Such pathogens are called prions, which are composed of host-encoded proteins that adopt alternative conformations (Chap. 453e). Prions are self-propagating by imposing their conformations on the normal, precursor protein;

most prions are enriched for β -sheet and can assemble into amyloid fibrils.

Similar to the plaques in kuru and CJD that are composed of PrP prions, the amyloid plaques in AD are filled with A β prions that have polymerized into fibrils. This relationship between the neuropathologic findings and the etiologic prion was strengthened by the genetic linkage between familial CJD and mutations in the PrP gene, as well as (as noted above) between familial AD and mutations in the APP gene. Moreover, a mutation in the APP gene that prevents A β peptide formation was correlated with a decreased incidence of AD in Iceland.

The heritable neurodegenerative diseases offer an important insight into the pathogenesis of the more common, sporadic ones. Although the mutant proteins that cause these disorders are expressed in the brains of people early in life, the diseases do not occur for many decades. Many explanations for the late onset of familial neurodegenerative diseases have been offered, but none are supported by substantial experimental evidence. The late onset might be due to a second event in which a mutant protein, after its conversion into a prion, begins to accumulate at some rather advanced age (Fig. 444e-5). Such a formulation is also consistent with data showing that the protein quality-control mechanisms diminish in efficiency with age. Thus, the prion forms of both wild-type and mutant proteins are likely to be efficiently degraded in younger people but are less well handled in older individuals. This explanation is consistent with the view that neurodegenerative diseases are disorders of the aging nervous system.

A new classification for neurodegenerative diseases can be proposed based on not only the traditional phenotypic presentation and neuropathology, but also the prion etiology (Table 444e-4). Over the past decade, an expanding body of experimental data has accumulated implicating prions in each of these illnesses. In addition to kuru and CJD, Gerstmann-Sträussler-Scheinker disease (GSS) and fatal insomnia in humans are caused by PrP^{Sc} prions. In animals, PrP^{Sc} prions cause scrapie of sheep and goats, bovine spongiform encephalopathy (BSE), chronic wasting disease (CWD) of deer and elk, feline spongiform encephalopathy, and transmissible mink encephalopathy (TME). Similar to PrP, A β , tau, α -synuclein, superoxide dismutase 1 (SOD1), and possibly huntingtin all adopt alternative conformations that become self-propagating, and thus, each protein can become a prion and be transferred to synaptically connected neurons. Moreover, each of these prions causes a distinct constellation of neurodegenerative diseases.

Evidence for a prion etiology of AD comes from a series of transmission experiments initially performed in marmosets and more recently in transgenic (Tg) mice inoculated with a synthetic A β peptide folded into a prion. Studies with the tau protein have shown that it not only features in the pathogenesis of AD, but also causes such illnesses as the frontotemporal dementias including chronic traumatic encephalopathy, which has been reported in both contact sport athletes and military personnel who have suffered traumatic brain injuries. A series of incisive studies using cultured cells and Tg mice has demonstrated that tau can become a prion and multiply in the brain. In contrast to the A β and tau prions, a strain of α -synuclein prions found in the brains of patients who died of multiple system atrophy (MSA) killed the Tg mouse host ~90 days after intracerebral inoculation, whereas α -synuclein prions formed spontaneously in Tg mouse brains killed recipient mice in ~200 days.

For many years, the most frequently cited argument against prions was the existence of strains that produced distinct clinical presentations and different patterns of neuropathologic lesions. Some investigators argued that the biologic information carried in different prion strains could only be encoded within a nucleic acid. Subsequently, many studies demonstrated that strain-specified variation is encoded in the conformation of PrP^{Sc}, but the molecular mechanisms responsible for the storage of biologic information remains enigmatic. The neuroanatomical patterns of prion deposition have been shown to be dependent on the particular strain of prion. Convincing evidence in support of this proposition has been accumulated for PrP, A β , tau, and α -synuclein prions.

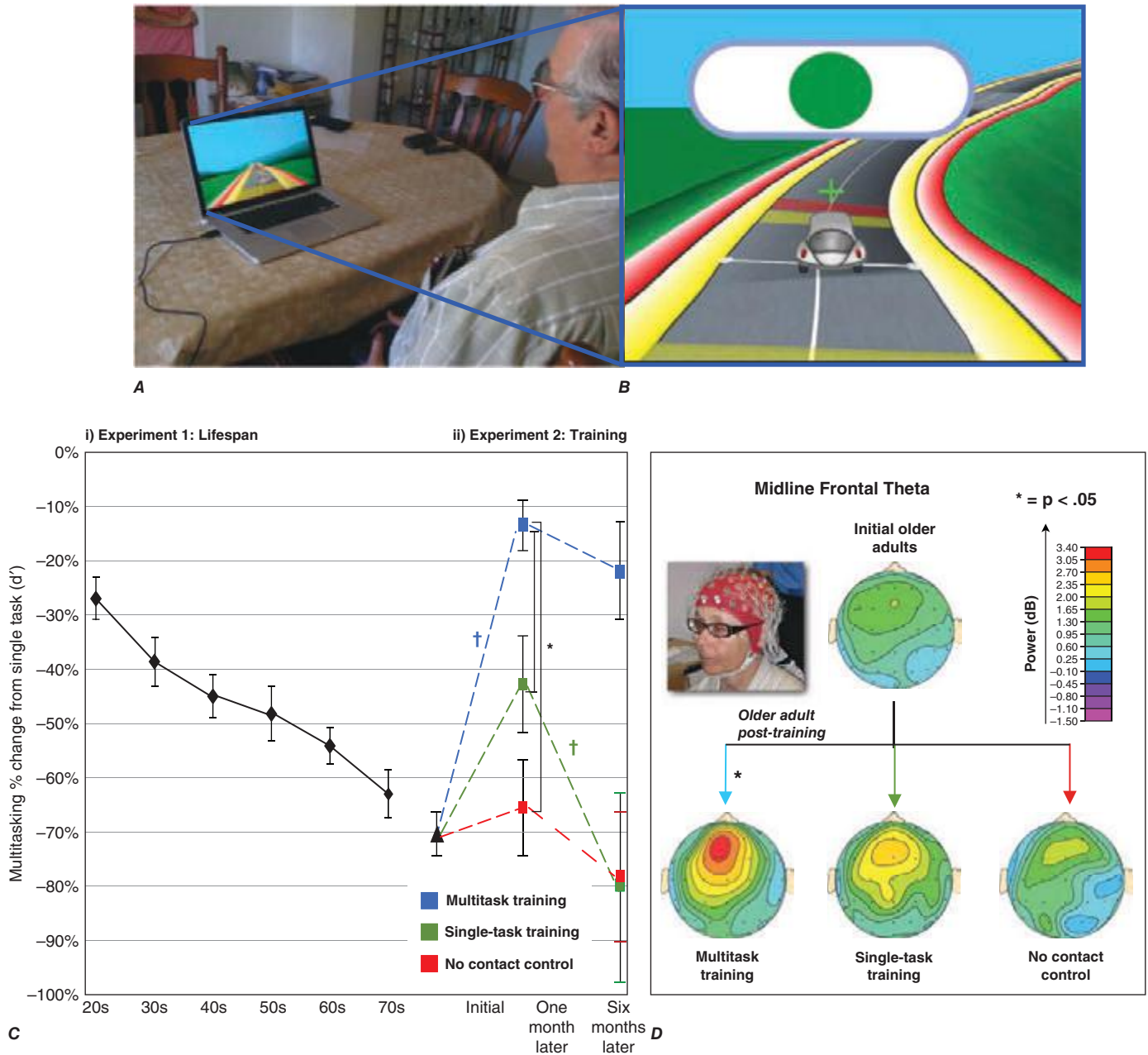


FIGURE 444e-5 Video game training can enhance cognitive performance. **A.** An older participant engaging in NeuroRacer training (driving while responding to target signs), with **(B)** a screen shot of the experimental training session. **C.** NeuroRacer multitasking costs for target discrimination increased (i.e., a larger percentage decrease from single task performance when multitasking) in (i) a linear fashion across the lifespan and with (ii) costs before training 1 month after training and 6 months after training, showing a differential benefit of multitasking training compared to a no-contact control group and a single-task training group. **D.** Midline frontal theta activity obtained with electroencephalogram showed significantly enhanced activity only following multitasking training, mimicking the pattern of change in the behavioral data as well as performance improvements on untrained tests of working memory and sustained attention (not presented). For details, see JA Anguera et al: Nature 501:97, 2013.

Although the number of prions identified in mammals and in fungi continues to expand, the existence of prions in other phylogeny remains undetermined. Some mammalian prions perform vital functions and do not cause disease; such nonpathogenic prions include the cytoplasmic polyadenylation element binding (CPEB) protein, the mitochondrial antiviral-signaling (MAVS) protein, and T cell-restricted intracellular antigen 1 (TIA-1).

All mammalian prion proteins adopt a β -sheet-rich conformation and appear to readily oligomerize as this process becomes self-propagating. Control of the self-propagating state of benign mammalian prions is not well understood but is critical for the well-being of the host. In contrast, pathogenic mammalian prions appear to multiply exponentially, but the mechanisms by which they cause disease are

poorly defined. We do not know if prions multiply as monomers or as oligomers; notably, the ionizing radiation target size of PrP^{Sc} prions seems to suggest it is a trimer. The oligomeric states of pathogenic mammalian prions are thought to be the toxic forms, and assembly into larger polymers, such as amyloid fibrils, seems to be a mechanism for minimizing toxicity.

To date, there is no medication that halts or even slows a human neurodegenerative disease. The development of drugs designed to inhibit the conversion of the normal precursor proteins into prions or to enhance the degradation of prions focuses on the initial step in prion accumulation. Although several drugs that cross the blood-brain barrier have been identified that prolong the lives of mice infected with scrapie prions, none have been identified that extend the lives

TABLE 444e-4 A PRION-BASED CLASSIFICATION OF NEURODEGENERATIVE DISEASES

Neurodegenerative Diseases	Causative Prion Proteins
Creutzfeldt-Jakob disease (CJD)	PrP ^{Sc}
Kuru	
Gerstmann-Sträussler-Scheinker (GSS)	
Fatal insomnia	
Bovine spongiform encephalopathy (BSE)	
Scrapie	
Chronic wasting disease (CWD)	
Feline spongiform encephalopathy	
Transmissible mink encephalopathy	
Alzheimer's disease (AD)	A β \rightarrow tau
Parkinson's disease	α -Synuclein
Frontotemporal dementias (FTDs)	Tau, TDP43, FUS (C9orf72, progranulin)
Posttraumatic FTD, called chronic traumatic encephalopathy	
Amyotrophic lateral sclerosis	SOD1, TDP43, FUS (C9orf72)
Huntington's disease	Huntingtin

of Tg mice that replicate human CJD prions. Despite doubling the length of incubation times in mice inoculated with scrapie prions, all of the mice eventually succumb to illness. Because all of the treated mice develop neurologic dysfunction at the same time, the mutation rate as judged by drug resistance is likely to approach 100%, which is much higher than mutation rates recorded for bacteria and viruses. Mutations in prions seem likely to represent conformational variants that are selected for in mammals where survival becomes limited by the fastest-replicating prions. The results of these studies make it likely that cocktails of drugs that attack a variety of prion conformers will be required for the development of effective therapeutics.

SYSTEMS NEUROSCIENCE

Systems neuroscience refers to study of the functions of neural circuits and how they relate to brain function, behavior, motor activity, and cognition. Brain imaging techniques, primarily functional magnetic resonance imaging (fMRI) and position emission tomography (PET), have made it possible to investigate, noninvasively and in awake individuals, cognitive processes such as perception, making judgments, paying attention, and thinking. This has allowed insights into how networks of neurons operate to produce behavior. Many of these studies at present are based on determining the connectivity of neural circuits and how they operate, and how this can be then modeled to produce improved understanding of physiologic processes. fMRI uses contrast mechanisms related to physiologic changes in tissue, and brain perfusion can be studied by observing the time course of changes in brain water signal as a bolus of injected paramagnetic gadolinium contrast moves through the brain. More recently, to study intrinsic contrast-related local changes in blood oxygenation with brain activity, blood-oxygen-level-dependent (BOLD) contrast has been used to provide a rapid noninvasive approach for functional assessment. These techniques have been reliably used in the field of both behavior and cognitive sciences. One example is the use of fMRI to demonstrate mirror neuron systems, imitative pathways activated when observing actions of others. Mirror neurons are thought to be important for social conditioning and for many forms of learning, and abnormalities in mirror neurons may underlie some autism disorders.

Both structural and functional connectivity methods show large-scale network dysfunction in AD and frontotemporal dementia. The networks targeted have been defined as the default network in AD and the salience network in frontotemporal dementia. The default network is characterized by an area of reduced glucose metabolism in the temporoparietal cortex, which precedes the onset of dementia and which is

an area preferentially affected by amyloid deposition. These networks are now thought to be pathways accounting for the spread of abnormally templated proteins (prions; see above), including β -amyloid, tau, and α -synuclein.

Other examples of the use of fMRI include the study of memory, revealing that not only is hippocampal activity correlated with declarative memory consolidation, but it also involves activation in the ventral medial prefrontal cortex. Consolidation of memory over time results in decreased activity of the hippocampus and progressively stronger activation in the ventral medial prefrontal region associated with retrieval of consolidated memories. fMRI has also been used to identify sequences of brain activation involved in normal movements and alterations in their activation associated with both injury and recovery, to plan neurosurgical operations, and remarkably, to reconstruct actual visual images from the occipital cortex. Noninvasive brain-computer interfaces have extraordinary potential to advance the development of robotics and exoskeleton devices guided by brain activity for patients with a variety of nervous system afflictions. Diffusion tensor imaging is a recently developed MRI technique that can measure macroscopic axonal organization in nervous system tissues; it appears to be useful in assessing myelin and axonal injuries as well as brain development. Advances in understanding neural processing have led to the development of the ability to demonstrate that humans have online voluntary control of human temporal lobe neurons.

Multitasking capabilities, including attention to tasks when faced with distractions, decline as we age due to a decline in medial prefrontal cognitive control system. When faced with a multitasking challenge, video game training can improve cognitive control capabilities by augmenting prefrontal suppression of the default network and, as measured by electroencephalography and fMRI, result in improved performance that is sustained and, importantly, transfers to other cognitive tasks not associated with the training paradigm.

A significant recent advance in neuropathology has discovered that sodium dodecyl sulfate (SDS) detergent treatment can render the brain transparent (CLARITY), removing lipids while preserving most protein and structural elements and providing opportunities to identify brain structures and neural networks with unprecedented detail.

A therapeutic technology that has long-reaching implications for the development of novel interventions for neurologic, including behavioral, conditions has been the development of deep-brain stimulation as a highly effective therapeutic intervention for treating excessively firing neurons in the subthalamic nucleus of patients with Parkinson's disease and the precingulate cortex in patients with depression.

BRAIN RESEARCH THROUGH ADVANCING INNOVATIVE NEUROTECHNOLOGIES (BRAIN) INITIATIVE

The BRAIN initiative, grand in scope, was launched in 2013 to speed development of advances to understand, treat, repair, and prevent common neurologic disorders that, in aggregate, affect more than 1 billion people worldwide. The initial goal of BRAIN is to bring together experts in neurobiology (including optogenetics), engineering, information technology, and other fields to develop novel visualization and electrophysiologic methods to better define and understand neural circuits and all the connections among individual neurons. The announcement of the BRAIN initiative followed just weeks after a similarly ambitious program, the Human Brain Project (HBP), was unveiled by the European Union. The HBP seeks to model individual neurons, neural circuits, and ultimately the entire brain using computer technologies. Its architects also envision layering clinical and biomarker data from large health care databases to identify biosignatures associated with human phenotypes, possibly leading to a fundamental reclassification of disease, a concept also proposed by others, including in a 2011 report of the National Academy of Sciences (Toward Precision Medicine: Building a Knowledge Network for Biomedical Research and a New Taxonomy of Disease). These two ambitious projects are expected to be complementary and over time will hopefully become increasingly

integrated. Emerging discoveries are also certain to stimulate a range of ethical questions, many of which are not unique to the neurosciences but do come into sharpest focus in this area. These include possible risks to the privacy of personal information about our health, cognitive capabilities, or behavioral attributes; the sanctity of our private thoughts; as well as concerns regarding neuroenhancement technologies, including

their potential military use. Both the BRAIN and HBP initiatives have put into place strong ethical components to ensure that these programs are carried out, from the outset and to the fullest extent possible, consistent with guiding ethical principles that include respect for individuals, public beneficence, justice and fairness, democratic deliberation, and transparency.

This page intentionally left blank

445 Seizures and Epilepsy

Daniel H. Lowenstein

A *seizure* (from the Latin *sacire*, “to take possession of”) is a paroxysmal event due to abnormal excessive or synchronous neuronal activity in the brain. Depending on the distribution of discharges, this abnormal brain activity can have various manifestations, ranging from dramatic convulsive activity to experiential phenomena not readily discernible by an observer. Although a variety of factors influence the incidence and prevalence of seizures, ~5–10% of the population will have at least one seizure, with the highest incidence occurring in early childhood and late adulthood.

The meaning of the term *seizure* needs to be carefully distinguished from that of *epilepsy*. *Epilepsy* describes a condition in which a person has *recurrent* seizures due to a chronic, underlying process. This definition implies that a person with a single seizure, or recurrent seizures due to correctable or avoidable circumstances, does not necessarily have epilepsy. Epilepsy refers to a clinical phenomenon rather than a single disease entity, because there are many forms and causes of epilepsy. However, among the many causes of epilepsy there are various *epilepsy syndromes* in which the clinical and pathologic characteristics are distinctive and suggest a specific underlying etiology.

Using the definition of epilepsy as two or more unprovoked seizures, the incidence of epilepsy is ~0.3–0.5% in different populations throughout the world, and the prevalence of epilepsy has been estimated at 5–30 persons per 1000.

CLASSIFICATION OF SEIZURES

Determining the type of seizure that has occurred is essential for focusing the diagnostic approach on particular etiologies, selecting the appropriate therapy, and providing potentially vital information regarding prognosis. The International League against Epilepsy (ILAE) Commission on Classification and Terminology, 2005–2009 has provided an updated approach to classification of seizures (Table 445-1). This system is based on the clinical features of seizures and associated electroencephalographic findings. Other potentially distinctive features such as etiology or cellular substrate are not considered in this classification system, although this will undoubtedly change in the future as more is learned about the pathophysiologic mechanisms that underlie specific seizure types.

A fundamental principle is that seizures may be either focal or generalized. *Focal seizures* originate within networks limited to one

cerebral hemisphere (note that the term *partial seizures* is no longer used). *Generalized seizures* arise within and rapidly engage networks distributed across both cerebral hemispheres. Focal seizures are usually associated with structural abnormalities of the brain. In contrast, generalized seizures may result from cellular, biochemical, or structural abnormalities that have a more widespread distribution. There are clear exceptions in both cases, however.

FOCAL SEIZURES

Focal seizures arise from a neuronal network either discretely localized within one cerebral hemisphere or more broadly distributed but still within the hemisphere. With the new classification system, the subcategories of “simple focal seizures” and “complex focal seizures” have been eliminated. Instead, depending on the presence of cognitive impairment, they can be described as focal seizures with or without dyscognitive features. Focal seizures can also evolve into generalized seizures. In the past this was referred to as *focal seizures with secondary generalization*, but the new system relies on specific descriptions of the type of generalized seizures that evolve from the focal seizure.

The routine interictal (i.e., between seizures) electroencephalogram (EEG) in patients with focal seizures is often normal or may show brief discharges termed *epileptiform spikes*, or *sharp waves*. Because focal seizures can arise from the medial temporal lobe or inferior frontal lobe (i.e., regions distant from the scalp), the EEG recorded during the seizure may be nonlocalizing. However, the seizure focus is often detected using sphenoidal or surgically placed intracranial electrodes.

Focal Seizures Without Dyscognitive Features Focal seizures can cause motor, sensory, autonomic, or psychic symptoms without impairment of cognition. For example, a patient having a focal motor seizure arising from the right primary motor cortex near the area controlling hand movement will note the onset of involuntary movements of the contralateral, left hand. These movements are typically clonic (i.e., repetitive, flexion/extension movements) at a frequency of ~2–3 Hz; pure tonic posturing may be seen as well. Since the cortical region controlling hand movement is immediately adjacent to the region for facial expression, the seizure may also cause abnormal movements of the face synchronous with the movements of the hand. The EEG recorded with scalp electrodes during the seizure (i.e., an ictal EEG) may show abnormal discharges in a very limited region over the appropriate area of cerebral cortex if the seizure focus involves the cerebral convexity. Seizure activity occurring within deeper brain structures is sometimes not detected by the standard EEG, however, and may require intracranial electrodes for its detection.

Three additional features of focal motor seizures are worth noting. First, in some patients, the abnormal motor movements may begin in a very restricted region such as the fingers and gradually progress (over seconds to minutes) to include a larger portion of the extremity. This phenomenon, described by Hughlings Jackson and known as a “Jacksonian march,” represents the spread of seizure activity over a progressively larger region of motor cortex. Second, patients may experience a localized paresis (Todd’s paralysis) for minutes to many hours in the involved region following the seizure. Third, in rare instances, the seizure may continue for hours or days. This condition, termed *epilepsia partialis continua*, is often refractory to medical therapy.

Focal seizures may also manifest as changes in somatic sensation (e.g., paresthesias), vision (flashing lights or formed hallucinations), equilibrium (sensation of falling or vertigo), or autonomic function (flushing, sweating, piloerection). Focal seizures arising from the temporal or frontal cortex may also cause alterations in hearing, olfaction, or higher cortical function (psychic symptoms). This includes the sensation of unusual, intense odors (e.g., burning rubber or kerosene) or sounds (crude or highly complex sounds), or an epigastric

TABLE 445-1 CLASSIFICATION OF SEIZURES

1. Focal seizures	(Can be further described as having motor, sensory, autonomic, cognitive, or other features)
2. Generalized seizures	
a. Absence	Typical
	Atypical
b. Tonic clonic	
c. Clonic	
d. Tonic	
e. Atonic	
f. Myoclonic	
3. May be focal, generalized, or unclear	
	Epileptic spasms

sensation that rises from the stomach or chest to the head. Some patients describe odd, internal feelings such as fear, a sense of impending change, detachment, depersonalization, *déjà vu*, or illusions that objects are growing smaller (*micropsia*) or larger (*macropsia*). These subjective, “internal” events that are not directly observable by someone else are referred to as *auras*.

Focal Seizures with Dyscognitive Features Focal seizures may also be accompanied by a transient impairment of the patient’s ability to maintain normal contact with the environment. The patient is unable to respond appropriately to visual or verbal commands during the seizure and has impaired recollection or awareness of the ictal phase. The seizures frequently begin with an aura (i.e., a focal seizure without cognitive disturbance) that is stereotypic for the patient. The start of the ictal phase is often a sudden behavioral arrest or motionless stare, which marks the onset of the period of impaired awareness. The behavioral arrest is usually accompanied by *automatisms*, which are involuntary, automatic behaviors that have a wide range of manifestations. Automatisms may consist of very basic behaviors such as chewing, lip smacking, swallowing, or “picking” movements of the hands, or more elaborate behaviors such as a display of emotion or running. The patient is typically confused following the seizure, and the transition to full recovery of consciousness may range from seconds up to an hour. Examination immediately following the seizure may show an anterograde amnesia or, in cases involving the dominant hemisphere, a postictal aphasia.

The range of potential clinical behaviors linked to focal seizures is so broad that extreme caution is advised before concluding that stereotypic episodes of bizarre or atypical behavior are not due to seizure activity. In such cases additional, detailed EEG studies may be helpful.

EVOLUTION OF FOCAL SEIZURES TO GENERALIZED SEIZURES

Focal seizures can spread to involve both cerebral hemispheres and produce a generalized seizure, usually of the tonic-clonic variety (discussed below). This evolution is observed frequently following focal seizures arising from a focus in the frontal lobe, but may also be associated with focal seizures occurring elsewhere in the brain. A focal seizure that evolves into a generalized seizure is often difficult to distinguish from a primary generalized-onset tonic-clonic seizure, because bystanders tend to emphasize the more dramatic, generalized convulsive phase of the seizure and overlook the more subtle, focal symptoms present at onset. In some cases, the focal onset of the seizure becomes apparent only when a careful history identifies a preceding aura. Often, however, the focal onset is not clinically evident and may be established only through careful EEG analysis. Nonetheless, distinguishing between these two entities is extremely important, because there may be substantial differences in the evaluation and treatment of epilepsies associated with focal versus generalized seizures.

GENERALIZED SEIZURES

Generalized seizures are thought to arise at some point in the brain but immediately and rapidly engage neuronal networks in both cerebral hemispheres. Several types of generalized seizures have features that place them in distinctive categories and facilitate clinical diagnosis.

Typical Absence Seizures Typical absence seizures are characterized by sudden, brief lapses of consciousness without loss of postural control. The seizure typically lasts for only seconds, consciousness returns as suddenly as it was lost, and there is no postictal confusion. Although the brief loss of consciousness may be clinically inapparent or the sole manifestation of the seizure discharge, absence seizures are usually accompanied by subtle, bilateral motor signs such as rapid blinking of the eyelids, chewing movements, or small-amplitude, clonic movements of the hands.

Typical absence seizures are associated with a group of genetically determined epilepsies with onset usually in childhood (ages 4–8 years) or early adolescence and are the main seizure type in 15–20% of children with epilepsy. The seizures can occur hundreds of times per day, but the child may be unaware of or unable to convey their existence. Because the clinical signs of the seizures are subtle, especially to parents

who may not have had previous experience with seizures, it is not surprising that the first clue to absence epilepsy is often unexplained “daydreaming” and a decline in school performance recognized by a teacher.

The electrophysiologic hallmark of typical absence seizures is a generalized, symmetric, 3-Hz spike-and-wave discharge that begins and ends suddenly, superimposed on a normal EEG background. Periods of spike-and-wave discharges lasting more than a few seconds usually correlate with clinical signs, but the EEG often shows many more brief bursts of abnormal cortical activity than were suspected clinically. Hyperventilation tends to provoke these electrographic discharges and even the seizures themselves and is routinely used when recording the EEG.

Atypical Absence Seizures Atypical absence seizures have features that deviate both clinically and electrophysiologically from typical absence seizures. For example, the lapse of consciousness is usually of longer duration and less abrupt in onset and cessation, and the seizure is accompanied by more obvious motor signs that may include focal or lateralizing features. The EEG shows a generalized, slow spike-and-wave pattern with a frequency of ≤ 2.5 per second, as well as other abnormal activity. Atypical absence seizures are usually associated with diffuse or multifocal structural abnormalities of the brain and therefore may accompany other signs of neurologic dysfunction such as mental retardation. Furthermore, the seizures are less responsive to anticonvulsants compared to typical absence seizures.

Generalized, Tonic-Clonic Seizures Generalized-onset tonic-clonic seizures are the main seizure type in ~10% of all persons with epilepsy. They are also the most common seizure type resulting from metabolic derangements and are therefore frequently encountered in many different clinical settings. The seizure usually begins abruptly without warning, although some patients describe vague premonitory symptoms in the hours leading up to the seizure. This prodrome is distinct from the stereotypic auras associated with focal seizures that generalize. The initial phase of the seizure is usually tonic contraction of muscles throughout the body, accounting for a number of the classic features of the event. Tonic contraction of the muscles of expiration and the larynx at the onset will produce a loud moan or “ictal cry.” Respirations are impaired, secretions pool in the oropharynx, and cyanosis develops. Contraction of the jaw muscles may cause biting of the tongue. A marked enhancement of sympathetic tone leads to increases in heart rate, blood pressure, and pupillary size. After 10–20 s, the tonic phase of the seizure typically evolves into the clonic phase, produced by the superimposition of periods of muscle relaxation on the tonic muscle contraction. The periods of relaxation progressively increase until the end of the ictal phase, which usually lasts no more than 1 min. The postictal phase is characterized by unresponsiveness, muscular flaccidity, and excessive salivation that can cause stridorous breathing and partial airway obstruction. Bladder or bowel incontinence may occur at this point. Patients gradually regain consciousness over minutes to hours, and during this transition, there is typically a period of postictal confusion. Patients subsequently complain of headache, fatigue, and muscle ache that can last for many hours. The duration of impaired consciousness in the postictal phase can be extremely long (i.e., many hours) in patients with prolonged seizures or underlying central nervous system (CNS) diseases such as alcoholic cerebral atrophy.

The EEG during the tonic phase of the seizure shows a progressive increase in generalized low-voltage fast activity, followed by generalized high-amplitude, polyspike discharges. In the clonic phase, the high-amplitude activity is typically interrupted by slow waves to create a spike-and-wave pattern. The postictal EEG shows diffuse slowing that gradually recovers as the patient awakens.

There are a number of variants of the generalized tonic-clonic seizure, including pure tonic and pure clonic seizures. Brief tonic seizures lasting only a few seconds are especially noteworthy since they are usually associated with specific epileptic syndromes having mixed seizure phenotypes, such as the Lennox-Gastaut syndrome (discussed below).

Atonic Seizures Atonic seizures are characterized by sudden loss of postural muscle tone lasting 1–2 s. Consciousness is briefly impaired,

but there is usually no postictal confusion. A very brief seizure may cause only a quick head drop or nodding movement, whereas a longer seizure will cause the patient to collapse. This can be extremely dangerous, because there is a substantial risk of direct head injury with the fall. The EEG shows brief, generalized spike-and-wave discharges followed immediately by diffuse slow waves that correlate with the loss of muscle tone. Similar to pure tonic seizures, atonic seizures are usually seen in association with known epilepsy syndromes.

Myoclonic Seizures Myoclonus is a sudden and brief muscle contraction that may involve one part of the body or the entire body. A normal, common physiologic form of myoclonus is the sudden jerking movement observed while falling asleep. Pathologic myoclonus is most commonly seen in association with metabolic disorders, degenerative CNS diseases, or anoxic brain injury (**Chap. 330**). Although the distinction from other forms of myoclonus is imprecise, myoclonic seizures are considered to be true epileptic events because they are caused by cortical (versus subcortical or spinal) dysfunction. The EEG may show bilaterally synchronous spike-and-wave discharges synchronized with the myoclonus, although these can be obscured by movement artifact. Myoclonic seizures usually coexist with other forms of generalized seizures but are the predominant feature of juvenile myoclonic epilepsy (discussed below).

CURRENTLY UNCLASSIFIABLE SEIZURES

Not all seizure types can be designated as focal or generalized, and they should therefore be labeled as “unclassifiable” until additional evidence allows a valid classification. *Epileptic spasms* are such an example. These are characterized by a briefly sustained flexion or extension of predominantly proximal muscles, including truncal muscles. The EEG in these patients usually shows hypsarrhythmias, which consist of diffuse, giant slow waves with a chaotic background of irregular, multifocal spikes and sharp waves. During the clinical spasm, there is a marked suppression of the EEG background (the “electrodecremental response”). The electromyogram (EMG) also reveals a characteristic rhomboid pattern that may help distinguish spasms from brief tonic and myoclonic seizures. Epileptic spasms occur predominantly in infants and likely result from differences in neuronal function and connectivity in the immature versus mature CNS.

EPILEPSY SYNDROMES

Epilepsy syndromes are disorders in which epilepsy is a predominant feature, and there is sufficient evidence (e.g., through clinical, EEG, radiologic, or genetic observations) to suggest a common underlying mechanism. Three important epilepsy syndromes are listed below; additional examples with a known genetic basis are shown in **Table 445-2**.

JUVENILE MYOCLONIC EPILEPSY

Juvenile myoclonic epilepsy (JME) is a generalized seizure disorder of unknown cause that appears in early adolescence and is usually characterized by bilateral myoclonic jerks that may be single or repetitive. The myoclonic seizures are most frequent in the morning after awakening and can be provoked by sleep deprivation. Consciousness is preserved unless the myoclonus is especially severe. Many patients also experience generalized tonic-clonic seizures, and up to one-third have absence seizures. Although complete remission is relatively uncommon, the seizures usually respond well to appropriate anticonvulsant medication. There is often a family history of epilepsy, and genetic linkage studies suggest a polygenic cause.

LENNOX-GASTAUT SYNDROME

Lennox-Gastaut syndrome occurs in children and is defined by the following triad: (1) multiple seizure types (usually including generalized tonic-clonic, atonic, and atypical absence seizures); (2) an EEG showing slow (<3 Hz) spike-and-wave discharges and a variety of other abnormalities; and (3) impaired cognitive function in most but not all cases. Lennox-Gastaut syndrome is associated with CNS disease or dysfunction from a variety of causes, including *de novo* mutations,

developmental abnormalities, perinatal hypoxia/ischemia, trauma, infection, and other acquired lesions. The multifactorial nature of this syndrome suggests that it is a nonspecific response of the brain to diffuse neural injury. Unfortunately, many patients have a poor prognosis due to the underlying CNS disease and the physical and psychosocial consequences of severe, poorly controlled epilepsy.

MESIAL TEMPORAL LOBE EPILEPSY SYNDROME

Mesial temporal lobe epilepsy (MTLE) is the most common syndrome associated with focal seizures with dyscognitive features and is an example of an epilepsy syndrome with distinctive clinical, electroencephalographic, and pathologic features (**Table 445-3**). High-resolution magnetic resonance imaging (MRI) can detect the characteristic hippocampal sclerosis that appears to be essential in the pathophysiology of MTLE for many patients (**Fig. 445-1**). Recognition of this syndrome is especially important because it tends to be refractory to treatment with anticonvulsants but responds well to surgical intervention. Advances in the understanding of basic mechanisms of epilepsy have come through studies of experimental models of MTLE, discussed below.

THE CAUSES OF SEIZURES AND EPILEPSY

Seizures are a result of a shift in the normal balance of excitation and inhibition within the CNS. Given the numerous properties that control neuronal excitability, it is not surprising that there are many different ways to perturb this normal balance, and therefore many different causes of both seizures and epilepsy. Three clinical observations emphasize how a variety of factors determine why certain conditions may cause seizures or epilepsy in a given patient.

1. *The normal brain is capable of having a seizure under the appropriate circumstances, and there are differences between individuals in the susceptibility or threshold for seizures.* For example, seizures may be induced by high fevers in children who are otherwise normal and who never develop other neurologic problems, including epilepsy. However, febrile seizures occur only in a relatively small proportion of children. This implies there are various underlying *endogenous factors* that influence the threshold for having a seizure. Some of these factors are genetic, as a family history of epilepsy has a clear influence on the likelihood of seizures occurring in otherwise normal individuals. Normal development also plays an important role, because the brain appears to have different seizure thresholds at different maturational stages.
2. *There are a variety of conditions that have an extremely high likelihood of resulting in a chronic seizure disorder.* One of the best examples of this is severe, penetrating head trauma, which is associated with up to a 45% risk of subsequent epilepsy. The high propensity for severe traumatic brain injury to lead to epilepsy suggests that the injury results in a long-lasting pathologic change in the CNS that transforms a presumably normal neural network into one that is abnormally hyperexcitable. This process is known as *epileptogenesis*, and the specific changes that result in a lowered seizure threshold can be considered *epileptogenic factors*. Other processes associated with epileptogenesis include stroke, infections, and abnormalities of CNS development. Likewise, the genetic abnormalities associated with epilepsy likely involve processes that trigger the appearance of specific sets of epileptogenic factors.
3. *Seizures are episodic.* Patients with epilepsy have seizures intermittently and, depending on the underlying cause, many patients are completely normal for months or even years between seizures. This implies there are important provocative or *precipitating factors* that induce seizures in patients with epilepsy. Similarly, precipitating factors are responsible for causing the single seizure in someone without epilepsy. Precipitants include those due to intrinsic physiologic processes such as psychological or physical stress, sleep deprivation, or hormonal changes associated with the menstrual cycle. They also include exogenous factors such as exposure to toxic substances and certain medications.

TABLE 445-2 EXAMPLES OF GENES ASSOCIATED WITH EPILEPSY SYNDROMES^a

Gene (Locus)	Function of Gene	Clinical Syndrome	Comments
<i>CHRNA4</i> (20q13.2)	Nicotinic acetylcholine receptor subunit; mutations cause alterations in Ca ²⁺ flux through the receptor; this may reduce amount of GABA release in presynaptic terminals	Autosomal dominant nocturnal frontal lobe epilepsy (ADNFLE); childhood onset; brief, nighttime seizures with prominent motor movements; often misdiagnosed as primary sleep disorder	Rare; first identified in a large Australian family; other families found to have mutations in <i>CHRNA2</i> or <i>CHRN2</i> , and some families appear to have mutations at other loci
<i>KCNQ2</i> (20q13.3)	Voltage-gated potassium channel subunits; mutation in pore regions may cause a 20–40% reduction of potassium currents, which will lead to impaired repolarization	Benign familial neonatal seizures (BFNS); autosomal dominant inheritance; onset in 1st week of life in infants who are otherwise normal; remission usually within weeks to months; long-term epilepsy in 10–15%	Rare; other families found to have mutations in <i>KCNQ3</i> or an inversion in chromosomal 5; sequence and functional homology to <i>KCNQ1</i> , mutations of which cause long QT syndrome and a cardiac-auditory syndrome
<i>SCN1A</i> (2q24.3)	α-Subunit of a voltage-gated sodium channel; numerous mutations affecting sodium currents that cause either gain or loss of function; network effects appear related to expression in excitatory or inhibitory cells	Generalized epilepsy with febrile seizures plus (GEFS+); autosomal dominant inheritance; presents with febrile seizures at median 1 year, which may persist >6 years, then variable seizure types not associated with fever; numerous other syndromes, including almost 80% of patients with Dravet's syndrome (severe myoclonic epilepsy of infancy) and some cases of Lennox-Gastaut syndrome	Incidence uncertain; GEFS+ identified in other families with mutations in other sodium channel subunits (<i>SCN2B</i> and <i>SCN2A</i>) and GABA _A receptor subunit (<i>GABRG2</i> and <i>GABRA1</i>); significant phenotypic heterogeneity within same family, including members with febrile seizures only
<i>LGII</i> (10q24)	Leucine-rich glioma-inactivated 1 gene; previous evidence for role in glial tumor progression; recent studies suggest an influence in the postnatal development of glutamatergic circuits in the hippocampus	Autosomal dominant partial epilepsy with auditory features (ADPEAF); a form of idiopathic lateral temporal lobe epilepsy with auditory symptoms or aphasia as a major focal seizure manifestation; age of onset usually between 10 and 25 years	Mutations found in up to 50% of families containing two or more subjects with idiopathic localization-related epilepsy with ictal auditory symptoms, suggesting that at least one other gene may underlie this syndrome.
<i>DEPDC5</i> (22q12.2)	Disheveled, Egl-10 and pleckstrin domain containing protein 5; exerts an inhibitory effect on mammalian target of rapamycin (mTOR)-mediated processes, such as cell growth and proliferation	Autosomal dominant familial focal epilepsy with variable foci (FFEVF); family members have seizures originating from different cortical regions; neuroimaging usually normal but may harbor subtle malformations; recent studies also suggest association with benign epilepsy with centrotemporal spikes	Study of families with limited number of affected members revealed mutations in approximately 12% of families; thus may be a relatively common cause of lesion-negative focal epilepsies with suspected genetic basis
<i>CSTB</i> (21q22.3)	Cystatin B, a noncaspase cysteine protease inhibitor; normal protein may block neuronal apoptosis by inhibiting caspases directly or indirectly (via cathepsins), or controlling proteolysis	Progressive myoclonus epilepsy (PME) (Unverricht-Lundborg disease); autosomal recessive inheritance; age of onset between 6 and 15 years, myoclonic seizures, ataxia, and progressive cognitive decline; brain shows neuronal degeneration	Overall rare, but relatively common in Finland and Western Mediterranean (>1 in 20,000); precise role of cystatin B in human disease unknown, although mice with null mutations of cystatin B have similar syndrome
<i>EPM2A</i> (6q24)	Laforin, a protein tyrosine phosphatase (PTP); involved in glycogen metabolism and may have antiapoptotic activity	Progressive myoclonus epilepsy (Lafora's disease); autosomal recessive inheritance; age of onset 6–19 years, death within 10 years; brain degeneration associated with polyglucosan intracellular inclusion bodies in numerous organs	Most common PME in Southern Europe, Middle East, Northern Africa, and Indian subcontinent; genetic heterogeneity; unknown whether seizure phenotype due to degeneration or direct effects of abnormal laforin expression
<i>Doublecortin</i> (Xq21-24)	<i>Doublecortin</i> , expressed primarily in frontal lobes; directly regulates microtubule polymerization and bundling	Classic lissencephaly associated with severe mental retardation and seizures in males; subcortical band heterotopia with more subtle findings in females (presumably due to random X-inactivation); X-linked dominant	Relatively rare but of uncertain incidence; recent increased ascertainment due to improved imaging techniques; relationship between migration defect and seizure phenotype unknown

^aThe first five syndromes listed in the table (ADNFLE, BFNS, GEFS+, ADPEAF, and FFEVF) are examples of idiopathic epilepsies associated with identified gene mutations. The last three syndromes are examples of the numerous Mendelian disorders in which seizures are one part of the phenotype.

Abbreviations: GABA, γ-aminobutyric acid; PME, progressive myoclonus epilepsy.

These observations emphasize the concept that the many causes of seizures and epilepsy result from a dynamic interplay between endogenous factors, epileptogenic factors, and precipitating factors. The potential role of each needs to be carefully considered when determining the appropriate management of a patient with seizures. For example, the identification of predisposing factors (e.g., family history of epilepsy) in a patient with febrile seizures may increase the necessity for closer follow-up and a more aggressive diagnostic evaluation. Finding an epileptogenic lesion may help in the estimation of seizure recurrence and duration of therapy. Finally, removal or modification of a precipitating factor may be an effective and safer method for preventing further seizures than the prophylactic use of anticonvulsant drugs.

CAUSES ACCORDING TO AGE

In practice, it is useful to consider the etiologies of seizures based on the age of the patient, because age is one of the most important factors

determining both the incidence and the likely causes of seizures or epilepsy (Table 445-4). During the *neonatal period and early infancy*, potential causes include hypoxic-ischemic encephalopathy, trauma, CNS infection, congenital CNS abnormalities, and metabolic disorders. Babies born to mothers using neurotoxic drugs such as cocaine, heroin, or ethanol are susceptible to drug-withdrawal seizures in the first few days after delivery. Hypoglycemia and hypocalcemia, which can occur as secondary complications of perinatal injury, are also causes of seizures early after delivery. Seizures due to inborn errors of metabolism usually present once regular feeding begins, typically 2–3 days after birth. Pyridoxine (vitamin B₆) deficiency, an important cause of neonatal seizures, can be effectively treated with pyridoxine replacement. The idiopathic or inherited forms of benign neonatal convulsions are also seen during this time period.

The most common seizures arising in *late infancy and early childhood* are febrile seizures, which are seizures associated with fevers but without evidence of CNS infection or other defined causes. The overall prevalence

TABLE 445-3 CHARACTERISTICS OF THE MESIAL TEMPORAL LOBE EPILEPSY SYNDROME

History	
History of febrile seizures	Rare generalized seizures
Family history of epilepsy	Seizures may remit and reappear
Early onset	Seizures often intractable
Clinical Observations	
Aura common	Postictal disorientation
Behavioral arrest/stare	Memory loss
Complex automatisms	Dysphasia (with focus in dominant hemisphere)
Unilateral posturing	
Laboratory Studies	
Unilateral or bilateral anterior temporal spikes on EEG	
Hypometabolism on interictal PET	
Hypoperfusion on interictal SPECT	
Material-specific memory deficits on intracarotid amobarbital (Wada) test	
MRI Findings	
Small hippocampus with increased signal on T2-weighted sequences	
Small temporal lobe	
Enlarged temporal horn	
Pathologic Findings	
Highly selective loss of specific cell populations within hippocampus in most cases	

Abbreviations: EEG, electroencephalogram; MRI, magnetic resonance imaging; PET, positron emission tomography; SPECT, single-photon emission computed tomography.

is 3–5% and even higher in some parts of the world such as Asia. Patients often have a family history of febrile seizures or epilepsy. Febrile seizures usually occur between 3 months and 5 years of age and have a peak incidence between 18 and 24 months. The typical scenario is a child who has a generalized, tonic-clonic seizure during a febrile illness in the setting of a common childhood infection such as otitis media, respiratory

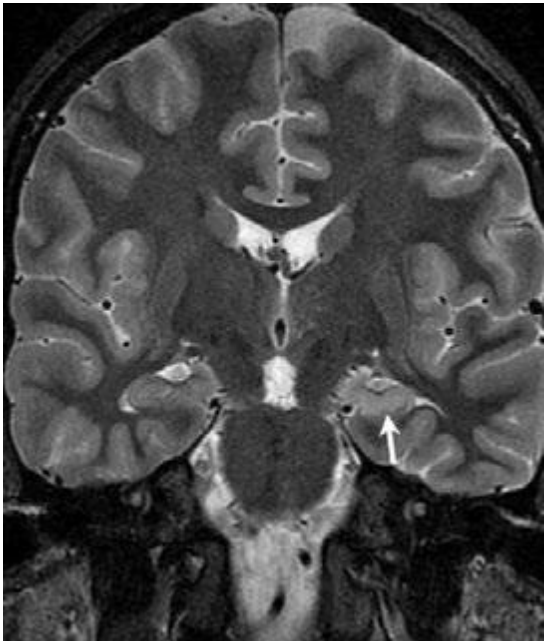


FIGURE 445-1 Mesial temporal lobe epilepsy. The electroencephalogram and seizure semiology were consistent with a left temporal lobe focus. This coronal high-resolution T2-weighted fast spin echo magnetic resonance image obtained at 3 tesla is at the level of the hippocampal bodies, and shows abnormal high signal intensity, blurring of internal laminar architecture, and reduced size of the left hippocampus (*arrow*) relative to the right. This triad of imaging findings is consistent with hippocampal sclerosis.

TABLE 445-4 CAUSES OF SEIZURES

Neonates (<1 month)	Perinatal hypoxia and ischemia Intracranial hemorrhage and trauma CNS infection Metabolic disturbances (hypoglycemia, hypocalcemia, hypomagnesemia, pyridoxine deficiency) Drug withdrawal Developmental disorders Genetic disorders
Infants and children (>1 month and <12 years)	Febrile seizures Genetic disorders (metabolic, degenerative, primary epilepsy syndromes) CNS infection Developmental disorders Trauma
Adolescents (12–18 years)	Trauma Genetic disorders Infection Illicit drug use Brain tumor
Young adults (18–35 years)	Trauma Alcohol withdrawal Illicit drug use Brain tumor Autoantibodies
Older adults (>35 years)	Cerebrovascular disease Brain tumor Alcohol withdrawal Metabolic disorders (uremia, hepatic failure, electrolyte abnormalities, hypoglycemia, hyperglycemia) Alzheimer's disease and other degenerative CNS diseases Autoantibodies

Abbreviation: CNS, central nervous system.

infection, or gastroenteritis. The seizure is likely to occur during the rising phase of the temperature curve (i.e., during the first day) rather than well into the course of the illness. A *simple* febrile seizure is a single, isolated event, brief, and symmetric in appearance. *Complex* febrile seizures are characterized by repeated seizure activity, duration >15 minutes, or by focal features. Approximately one-third of patients with febrile seizures will have a recurrence, but <10% have three or more episodes. Recurrences are much more likely when the febrile seizure occurs in the first year of life. Simple febrile seizures are not associated with an increase in the risk of developing epilepsy, while complex febrile seizures have a risk of 2–5%; other risk factors include the presence of preexisting neurologic deficits and a family history of nonfebrile seizures.

Childhood marks the age at which many of the well-defined epilepsy syndromes present. Some children who are otherwise normal develop idiopathic, generalized tonic-clonic seizures without other features that fit into specific syndromes. Temporal lobe epilepsy usually presents in childhood and may be related to mesial temporal lobe sclerosis (as part of the MTL syndrome) or other focal abnormalities such as cortical dysgenesis. Other types of focal seizures, including those that evolve into generalized seizures, may be the relatively late manifestation of a developmental disorder, an acquired lesion such as head trauma, CNS infection (especially viral encephalitis), or very rarely a CNS tumor.

The period of *adolescence and early adulthood* is one of transition during which the idiopathic or genetically based epilepsy syndromes, including JME and juvenile absence epilepsy, become less common, while epilepsies secondary to acquired CNS lesions begin to predominate. Seizures that arise in patients in this age group may be associated

with head trauma, CNS infections (including parasitic infections such as cysticercosis), brain tumors, congenital CNS abnormalities, illicit drug use, or alcohol withdrawal. Autoantibodies directed against CNS antigens such as potassium channels or glutamate receptors are a newly recognized cause of epilepsy that also begins to appear in this age group (although cases of autoimmunity are being increasingly described in the pediatric population), including patients without an identifiable cancer. This etiology should be suspected when a previously normal individual presents with a particularly aggressive seizure pattern developing over weeks to months and characterized by increasingly frequent and prolonged seizures combined with cognitive decline (Chap. 122).

Head trauma is a common cause of epilepsy in adolescents and adults. The head injury can be caused by a variety of mechanisms, and the likelihood of developing epilepsy is strongly correlated with the severity of the injury. A patient with a penetrating head wound, depressed skull fracture, intracranial hemorrhage, or prolonged post-traumatic coma or amnesia has a 30–50% risk of developing epilepsy, whereas a patient with a closed head injury and cerebral contusion has a 5–25% risk. Recurrent seizures usually develop within 1 year after head trauma, although intervals of >10 years are well known. In controlled studies, mild head injury, defined as a concussion with amnesia or loss of consciousness of <30 min, was found to be associated with only a slightly increased likelihood of epilepsy. Nonetheless, most epileptologists know of patients who have focal seizures within hours or days of a mild head injury and subsequently develop chronic seizures of the same type; such cases may represent rare examples of chronic epilepsy resulting from mild head injury.

The causes of seizures in *older adults* include cerebrovascular disease, trauma (including subdural hematoma), CNS tumors, and degenerative diseases. Cerebrovascular disease may account for ~50% of new cases of epilepsy in patients older than age 65. Acute seizures (i.e., occurring at the time of the stroke) are seen more often with embolic rather than hemorrhagic or thrombotic stroke. Chronic seizures typically appear months to years after the initial event and are associated with all forms of stroke.

Metabolic disturbances such as electrolyte imbalance, hypo- or hyperglycemia, renal failure, and hepatic failure may cause seizures at any age. Similarly, endocrine disorders, hematologic disorders, vasculitides, and many other systemic diseases may cause seizures over a broad age range. A wide variety of medications and abused substances are known to precipitate seizures as well (Table 445-5).

BASIC MECHANISMS

MECHANISMS OF SEIZURE INITIATION AND PROPAGATION

Focal seizure activity can begin in a very discrete region of cortex and then slowly invade the surrounding regions. The hallmark of an established seizure is typically an electrographic “spike” due to intense near-simultaneous firing of a large number of local excitatory neurons, resulting in an apparent hypersynchronization of the excitatory bursts across a relatively large cortical region. The bursting activity in individual neurons (the “paroxysmal depolarization shift”) is caused by a relatively long-lasting depolarization of the neuronal membrane due to influx of extracellular calcium (Ca^{2+}), which leads to the opening of voltage-dependent sodium (Na^+) channels, influx of Na^+ , and generation of repetitive action potentials. This is followed by a hyperpolarizing afterpotential mediated by γ -aminobutyric acid (GABA) receptors or potassium (K^+) channels, depending on the cell type. The synchronized bursts from a sufficient number of neurons result in a so-called spike discharge on the EEG.

The spreading seizure wavefront is slowed and ultimately halted by intact hyperpolarization and a “surround” inhibition created by feed-forward activation of inhibitory neurons. With sufficient activation, there is a recruitment of surrounding neurons via a number of synaptic and nonsynaptic mechanisms, including: (1) an increase in extracellular K^+ , which blunts hyperpolarization and depolarizes neighboring neurons; (2) accumulation of Ca^{2+} in presynaptic terminals, leading to enhanced neurotransmitter release; (3) depolarization-induced

TABLE 445-5 DRUGS AND OTHER SUBSTANCES THAT CAN CAUSE SEIZURES

Alkylating agents (e.g., busulfan, chlorambucil)

Antimalarials (chloroquine, mefloquine)

Antimicrobials/antivirals

β -lactam and related compounds

Quinolones

Acyclovir

Isoniazid

Ganciclovir

Anesthetics and analgesics

Meperidine

Fentanyl

Tramadol

Local anesthetics

Dietary supplements

Ephedra (ma huang)

Ginkgo

Immunomodulatory drugs

Cyclosporine

OKT3 (monoclonal antibodies to T cells)

Tacrolimus

Interferons

Psychotropics

Antidepressants (e.g., bupropion)

Antipsychotics (e.g., clozapine)

Lithium

Radiographic contrast agents

Drug withdrawal

Alcohol

Baclofen

Barbiturates (short-acting)

Benzodiazepines (short-acting)

Zolpidem

Drugs of abuse

Amphetamine

Cocaine

Phencyclidine

Methylphenidate

Flumazenil^a

^aIn benzodiazepine-dependent patients.

activation of the *N*-methyl-D-aspartate (NMDA) subtype of the excitatory amino acid receptor, which causes additional Ca^{2+} influx and neuronal activation; and (4) ephaptic interactions related to changes in tissue osmolarity and cell swelling. The recruitment of a sufficient number of neurons leads to the propagation of excitatory currents into contiguous areas via local cortical connections and to more distant areas via long commissural pathways such as the corpus callosum.

Many factors control neuronal excitability, and thus there are many potential mechanisms for altering a neuron’s propensity to have bursting activity. Mechanisms *intrinsic* to the neuron include changes in the conductance of ion channels, response characteristics of membrane receptors, cytoplasmic buffering, second-messenger systems, and protein expression as determined by gene transcription, translation, and posttranslational modification. Mechanisms *extrinsic* to the neuron include changes in the amount or type of neurotransmitters present at the synapse, modulation of receptors by extracellular ions and other molecules, and temporal and spatial properties of synaptic and nonsynaptic input. Nonneural cells, such as astrocytes and oligodendrocytes, have an important role in many of these mechanisms as well.

Certain recognized causes of seizures are explained by these mechanisms. For example, accidental ingestion of domoic acid, which is an

analogue of glutamate (the principal excitatory neurotransmitter in the brain), causes profound seizures via direct activation of excitatory amino acid receptors throughout the CNS. Penicillin, which can lower the seizure threshold in humans and is a potent convulsant in experimental models, reduces inhibition by antagonizing the effects of GABA at its receptor. The basic mechanisms of other precipitating factors of seizures such as sleep deprivation, fever, alcohol withdrawal, hypoxia, and infection, are not as well understood but presumably involve analogous perturbations in neuronal excitability. Similarly, the endogenous factors that determine an individual's seizure threshold may relate to these properties as well.

Knowledge of the mechanisms responsible for initiation and propagation of most generalized seizures (including tonic-clonic, myoclonic, and atonic types) remains rudimentary and reflects the limited understanding of the connectivity of the brain at a systems level. Much more is understood about the origin of generalized spike-and-wave discharges in absence seizures. These appear to be related to oscillatory rhythms normally generated during sleep by circuits connecting the thalamus and cortex. This oscillatory behavior involves an interaction between GABA_B receptors, T-type Ca²⁺ channels, and K⁺ channels located within the thalamus. Pharmacologic studies indicate that modulation of these receptors and channels can induce absence seizures, and there is good evidence that the genetic forms of absence epilepsy may be associated with mutations of components of this system.

MECHANISMS OF EPILEPTOGENESIS

Epileptogenesis refers to the transformation of a normal neuronal network into one that is chronically hyperexcitable. There is often a delay of months to years between an initial CNS injury such as trauma, stroke, or infection and the first seizure. The injury appears to initiate a process that gradually lowers the seizure threshold in the affected region until a spontaneous seizure occurs. In many genetic and idiopathic forms of epilepsy, epileptogenesis is presumably determined by developmentally regulated events.

Pathologic studies of the hippocampus from patients with temporal lobe epilepsy have led to the suggestion that some forms of epileptogenesis are related to *structural changes in neuronal networks*. For example, many patients with MTLLE have a highly selective loss of neurons that may contribute to inhibition of the main excitatory neurons within the dentate gyrus. There is also evidence that, in response to the loss of neurons, there is reorganization or "sprouting" of surviving neurons in a way that affects the excitability of the network. Some of these changes can be seen in experimental models of prolonged electrical seizures or traumatic brain injury. Thus, an initial injury such as head injury may lead to a very focal, confined region of structural change that causes local hyperexcitability. The local hyperexcitability leads to further structural changes that evolve over time until the focal lesion produces clinically evident seizures. Similar models have provided strong evidence for long-term alterations in *intrinsic, biochemical properties of cells* within the network such as chronic changes in glutamate or GABA receptor function. Recent work has suggested that induction of inflammatory cascades may be a critical factor in these processes as well.

GENETIC CAUSES OF EPILEPSY



The most important recent progress in epilepsy research has been the identification of genetic mutations associated with a variety of epilepsy syndromes (Table 445-2). Although most of the mutations identified to date cause rare forms of epilepsy, their discovery has led to extremely important conceptual advances. For example, it appears that many of the inherited, idiopathic epilepsies (i.e., the relatively "pure" forms of epilepsy in which seizures are the phenotypic abnormality and brain structure and function are otherwise normal) are due to mutations affecting ion channel function. These syndromes are therefore part of the larger group of channelopathies causing paroxysmal disorders such as cardiac arrhythmias, episodic ataxia, periodic weakness, and familial hemiplegic migraine. In contrast, gene mutations observed in symptomatic epilepsies (i.e., disorders in which other neurologic abnormalities such as cognitive impairment coexist with seizures) are proving to be associated with

pathways influencing CNS development or neuronal homeostasis. *De novo* mutations may explain a significant proportion of these syndromes, especially those with onset in early childhood. A current challenge is to identify the multiple susceptibility genes that underlie the more common forms of idiopathic epilepsies. Recent studies suggest that ion channel mutations and copy number variants may contribute to causation in a subset of these patients.

MECHANISMS OF ACTION OF ANTIEPILEPTIC DRUGS

Antiepileptic drugs appear to act primarily by blocking the initiation or spread of seizures. This occurs through a variety of mechanisms that modify the activity of ion channels or neurotransmitters, and in most cases, the drugs have pleiotropic effects. The mechanisms include inhibition of Na⁺-dependent action potentials in a frequency-dependent manner (e.g., phenytoin, carbamazepine, lamotrigine, topiramate, zonisamide, lacosamide, rufinamide), inhibition of voltage-gated Ca²⁺ channels (phenytoin, gabapentin, pregabalin), facilitating the opening of potassium channels (ezogabine), attenuation of glutamate activity (lamotrigine, topiramate, felbamate), potentiation of GABA receptor function (benzodiazepines and barbiturates), increase in the availability of GABA (valproic acid, gabapentin, tiagabine), and modulation of release of synaptic vesicles (levetiracetam). The two most effective drugs for absence seizures, ethosuximide and valproic acid, probably act by inhibiting T-type Ca²⁺ channels in thalamic neurons.

In contrast to the relatively large number of antiepileptic drugs that can attenuate seizure activity, there are currently no drugs known to prevent the formation of a seizure focus following CNS injury. The eventual development of such "antiepileptogenic" drugs will provide an important means of preventing the emergence of epilepsy following injuries such as head trauma, stroke, and CNS infection.

APPROACH TO THE PATIENT:

Seizure

When a patient presents shortly after a seizure, the first priorities are attention to vital signs, respiratory and cardiovascular support, and treatment of seizures if they resume (see "Treatment: Seizures and Epilepsy"). Life-threatening conditions such as CNS infection, metabolic derangement, or drug toxicity must be recognized and managed appropriately.

When the patient is not acutely ill, the evaluation will initially focus on whether there is a history of earlier seizures (Fig. 445-2). If this is the first seizure, then the emphasis will be to: (1) establish whether the reported episode was a seizure rather than another paroxysmal event, (2) determine the cause of the seizure by identifying risk factors and precipitating events, and (3) decide whether anticonvulsant therapy is required in addition to treatment for any underlying illness.

In the patient with prior seizures or a known history of epilepsy, the evaluation is directed toward: (1) identification of the underlying cause and precipitating factors, and (2) determination of the adequacy of the patient's current therapy.

HISTORY AND EXAMINATION

The first goal is to determine whether the event was truly a seizure. An in-depth history is essential, because *in many cases the diagnosis of a seizure is based solely on clinical grounds—the examination and laboratory studies are often normal*. Questions should focus on the symptoms before, during, and after the episode in order to differentiate a seizure from other paroxysmal events (see "Differential Diagnosis of Seizures" below). Seizures frequently occur out-of-hospital, and the patient may be unaware of the ictal and immediate postictal phases; thus, witnesses to the event should be interviewed carefully.

The history should also focus on risk factors and predisposing events. Clues for a predisposition to seizures include a history of febrile seizures, earlier auras or brief seizures not recognized as such, and a family history of seizures. Epileptogenic factors such as prior head trauma, stroke, tumor, or CNS infection should be identified.

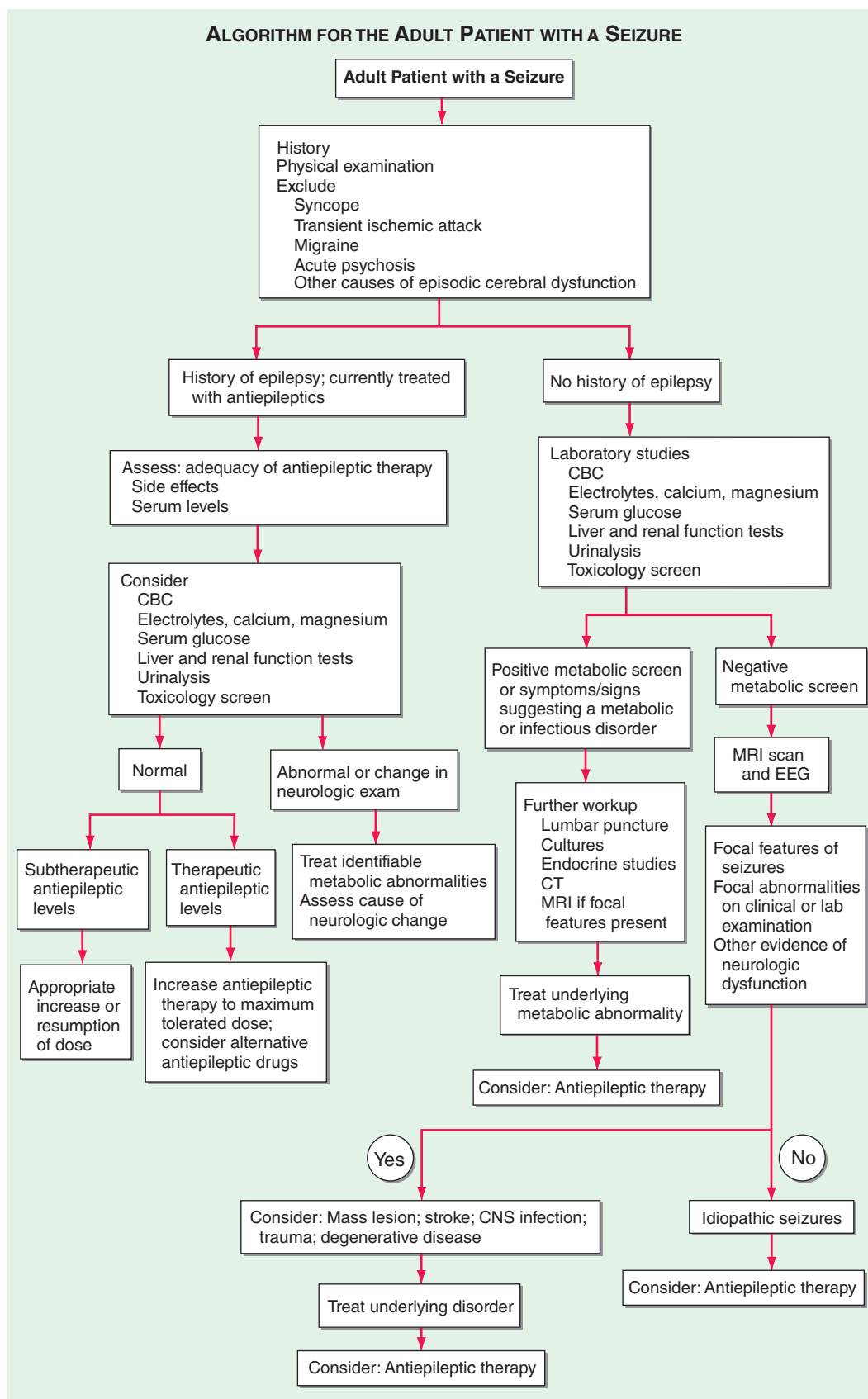


FIGURE 445-2 Evaluation of the adult patient with a seizure. CBC, complete blood count; CNS, central nervous system; CT, computed tomography; EEG, electroencephalogram; MRI, magnetic resonance imaging.

In children, a careful assessment of developmental milestones may provide evidence for underlying CNS disease. Precipitating factors such as sleep deprivation, systemic diseases, electrolyte or metabolic derangements, acute infection, drugs that lower the seizure threshold (Table 445-5), or alcohol or illicit drug use should also be identified.

The general physical examination includes a search for signs of infection or systemic illness. Careful examination of the skin may reveal signs of neurocutaneous disorders such as tuberous sclerosis or neurofibromatosis, or chronic liver or renal disease. A finding of organomegaly may indicate a metabolic storage disease, and limb

asymmetry may provide a clue to brain injury early in development. Signs of head trauma and use of alcohol or illicit drugs should be sought. Auscultation of the heart and carotid arteries may identify an abnormality that predisposes to cerebrovascular disease.

All patients require a complete neurologic examination, with particular emphasis on eliciting signs of cerebral hemispheric disease (**Chap. 437**). Careful assessment of mental status (including memory, language function, and abstract thinking) may suggest lesions in the anterior frontal, parietal, or temporal lobes. Testing of visual fields will help screen for lesions in the optic pathways and occipital lobes. Screening tests of motor function such as pronator drift, deep tendon reflexes, gait, and coordination may suggest lesions in motor (frontal) cortex, and cortical sensory testing (e.g., double simultaneous stimulation) may detect lesions in the parietal cortex.

LABORATORY STUDIES

Routine blood studies are indicated to identify the more common metabolic causes of seizures such as abnormalities in electrolytes, glucose, calcium, or magnesium, and hepatic or renal disease. A screen for toxins in blood and urine should also be obtained from all patients in appropriate risk groups, especially when no clear precipitating factor has been identified. A lumbar puncture is indicated if there is any suspicion of meningitis or encephalitis, and it is mandatory in all patients infected with HIV, even in the absence of symptoms or signs suggesting infection. Testing for autoantibodies in the serum and cerebrospinal fluid (CSF) should be considered in patients presenting with a seemingly aggressive form of epilepsy associated with other abnormalities such as cognitive disturbances.

ELECTROPHYSIOLOGIC STUDIES

All patients who have a possible seizure disorder should be evaluated with an EEG as soon as possible. **Details about the EEG are covered in Chap. 442e.**

In the evaluation of a patient with suspected epilepsy, the presence of *electrographic seizure activity* during the clinically evident event (i.e., abnormal, repetitive, rhythmic activity having a discrete onset and termination) clearly establishes the diagnosis. The absence of electrographic seizure activity does not exclude a seizure disorder, however, because focal seizures may originate from a region of the cortex that cannot be detected by standard scalp electrodes. The EEG is always abnormal during generalized tonic-clonic seizures. Because seizures are typically infrequent and unpredictable, it is often not possible to obtain the EEG during a clinical event. Continuous monitoring for prolonged periods in video-EEG telemetry units for hospitalized patients or the use of portable equipment to record the EEG continuously for ≥ 24 h in ambulatory patients has made it easier to capture the electrophysiologic accompaniments of clinical events. In particular, video-EEG telemetry is now a routine approach for the accurate diagnosis of epilepsy in patients with poorly characterized events or seizures that are difficult to control.

The EEG may also be helpful in the interictal period by showing certain abnormalities that are highly supportive of the diagnosis of epilepsy. Such *epileptiform activity* consists of bursts of abnormal discharges containing spikes or sharp waves. The presence of epileptiform activity is not specific for epilepsy, but it has a much greater prevalence in patients with epilepsy than in normal individuals. However, even in an individual who is known to have epilepsy, the initial routine interictal EEG may be normal up to 60% of the time. Thus, the EEG cannot establish the diagnosis of epilepsy in many cases.

The EEG is also used for classifying seizure disorders and aiding in the selection of anticonvulsant medications. For example, episodic generalized spike-wave activity is usually seen in patients with typical absence epilepsy and may be seen with other generalized epilepsy syndromes. Focal interictal epileptiform discharges would support the diagnosis of a focal seizure disorder such as temporal lobe epilepsy or frontal lobe seizures, depending on the location of the discharges.

The routine scalp-recorded EEG may also be used to assess the prognosis of seizure disorders; in general, a normal EEG implies a better prognosis, whereas an abnormal background or profuse epileptiform activity suggests a poor outcome. Unfortunately, the EEG has

not proved to be useful in predicting which patients with predisposing conditions such as head injury or brain tumor will go on to develop epilepsy, because in such circumstances epileptiform activity is commonly encountered regardless of whether seizures occur.

Magnetoencephalography (MEG) provides another way of looking noninvasively at cortical activity. Instead of measuring electrical activity of the brain, it measures the small magnetic fields that are generated by this activity. The source of epileptiform activity seen on MEG can be analyzed, and its source in the brain can be estimated using a variety of mathematical techniques. These source estimates can then be plotted on an anatomic image of the brain such as an MRI (discussed below), to generate a magnetic source image (MSI). MSI can be useful to localize potential seizure foci.

BRAIN IMAGING

Almost all patients with new-onset seizures should have a brain imaging study to determine whether there is an underlying structural abnormality that is responsible. The only potential exception to this rule is children who have an unambiguous history and examination suggestive of a benign, generalized seizure disorder such as absence epilepsy. MRI has been shown to be superior to computed tomography (CT) for the detection of cerebral lesions associated with epilepsy. In some cases, MRI will identify lesions such as tumors, vascular malformations, or other pathologies that need urgent therapy. The availability of newer MRI methods such as 3-tesla scanners, parallel imaging with multichannel head coils, three-dimensional structural imaging at submillimeter resolution, and widespread use of pulse sequences such as fluid-attenuated inversion recovery (FLAIR), has increased the sensitivity for detection of abnormalities of cortical architecture, including hippocampal atrophy associated with mesial temporal sclerosis, as well as abnormalities of cortical neuronal migration. In such cases, the findings may not lead to immediate therapy, but they do provide an explanation for the patient's seizures and point to the need for chronic antiepileptic drug therapy or possible surgical resection.

In the patient with a suspected CNS infection or mass lesion, CT scanning should be performed emergently when MRI is not immediately available. Otherwise, it is usually appropriate to obtain an MRI study within a few days of the initial evaluation. Functional imaging procedures such as positron emission tomography (PET) and single-photon emission computed tomography (SPECT) are also used to evaluate certain patients with medically refractory seizures (discussed below).

DIFFERENTIAL DIAGNOSIS OF SEIZURES

Disorders that may mimic seizures are listed in **Table 445-6**. In most cases, seizures can be distinguished from other conditions by meticulous attention to the history and relevant laboratory studies. On occasion, additional studies such as video-EEG monitoring, sleep studies, tilt-table analysis, or cardiac electrophysiology, may be required to reach a correct diagnosis. Two of the more common nonepileptic syndromes in the differential diagnosis are detailed below.

SYNCOPE

(**See also Chap. 27**) The diagnostic dilemma encountered most frequently is the distinction between a generalized seizure and syncope. Observations by the patient and bystanders that can help differentiate between the two are listed in **Table 445-7**. Characteristics of a seizure include the presence of an aura, cyanosis, unconsciousness, motor manifestations lasting >15 s, postictal disorientation, muscle soreness, and sleepiness. In contrast, a syncopal episode is more likely if the event was provoked by acute pain or anxiety or occurred immediately after arising from the lying or sitting position. Patients with syncope often describe a stereotyped transition from consciousness to unconsciousness that includes tiredness, sweating, nausea, and tunneling of vision, and they experience a relatively brief loss of consciousness. Headache or incontinence usually suggests a seizure but may on occasion also occur with syncope. A brief period (i.e., 1–10 s) of convulsive motor activity is frequently seen immediately at the onset of a syncopal episode, especially if the patient remains in an upright posture

TABLE 445-6 DIFFERENTIAL DIAGNOSIS OF SEIZURES

Syncope	Transient ischemic attack (TIA)
Vasovagal syncope	Basilar artery TIA
Cardiac arrhythmia	Sleep disorders
Valvular heart disease	Narcolepsy/cataplexy
Cardiac failure	Benign sleep myoclonus
Orthostatic hypotension	Movement disorders
Psychological disorders	Tics
Psychogenic seizure	Nonepileptic myoclonus
Hyperventilation	Paroxysmal choreoathetosis
Panic attack	Special considerations in children
Metabolic disturbances	Breath-holding spells
Alcoholic blackouts	Migraine with recurrent abdominal pain and cyclic vomiting
Delirium tremens	Benign paroxysmal vertigo
Hypoglycemia	Apnea
Hypoxia	Night terrors
Psychoactive drugs (e.g., hallucinogens)	Sleepwalking
Migraine	
Confusional migraine	
Basilar migraine	

after fainting (e.g., in a dentist's chair) and therefore has a sustained decrease in cerebral perfusion. Rarely, a syncopal episode can induce a full tonic-clonic seizure. In such cases, the evaluation must focus on both the cause of the syncopal event as well as the possibility that the patient has a propensity for recurrent seizures.

PSYCHOGENIC SEIZURES

Psychogenic seizures are nonepileptic behaviors that resemble seizures. They are often part of a conversion reaction precipitated by underlying psychological distress. Certain behaviors such as side-to-side turning of the head, asymmetric and large-amplitude shaking movements of the limbs, twitching of all four extremities without loss of consciousness, and pelvic thrusting are more commonly associated with psychogenic rather than epileptic seizures. Psychogenic seizures

TABLE 445-7 FEATURES THAT DISTINGUISH GENERALIZED TONIC-CLONIC SEIZURE FROM SYNCOPÉ

Features	Seizure	Syncopé
Immediate precipitating factors	Usually none	Emotional stress, Valsalva, orthostatic hypotension, cardiac etiologies
Premonitory symptoms	None or aura (e.g., odd odor)	Tiredness, nausea, diaphoresis, tunneling of vision
Posture at onset	Variable	Usually erect
Transition to unconsciousness	Often immediate	Gradual over seconds ^a
Duration of unconsciousness	Minutes	Seconds
Duration of tonic or clonic movements	30–60 s	Never more than 15 s
Facial appearance during event	Cyanosis, frothing at mouth	Pallor
Disorientation and sleepiness after event	Many minutes to hours	<5 min
Aching of muscles after event	Often	Sometimes
Biting of tongue	Sometimes	Rarely
Incontinence	Sometimes	Sometimes
Headache	Sometimes	Rarely

^aMay be sudden with certain cardiac arrhythmias.

often last longer than epileptic seizures and may wax and wane over minutes to hours. However, the distinction is sometimes difficult on clinical grounds alone, and there are many examples of diagnostic errors made by experienced epileptologists. This is especially true for psychogenic seizures that resemble focal seizures with dyscognitive features, because the behavioral manifestations of focal seizures (especially of frontal lobe origin) can be extremely unusual, and in both cases, the routine surface EEG may be normal. Video-EEG monitoring is very useful when historic features are nondiagnostic. Generalized tonic-clonic seizures always produce marked EEG abnormalities during and after the seizure. For suspected focal seizures of temporal lobe origin, the use of additional electrodes beyond the standard scalp locations (e.g., sphenoidal electrodes) may be required to localize a seizure focus. Measurement of serum prolactin levels may also help to distinguish between organic and psychogenic seizures, because most generalized seizures and some focal seizures are accompanied by rises in serum prolactin (during the immediate 30-min postictal period), whereas psychogenic seizures are not. The diagnosis of psychogenic seizures does not exclude a concurrent diagnosis of epilepsy, because the two often coexist.

TREATMENT SEIZURES AND EPILEPSY

Therapy for a patient with a seizure disorder is almost always multimodal and includes treatment of underlying conditions that cause or contribute to the seizures, avoidance of precipitating factors, suppression of recurrent seizures by prophylactic therapy with antiepileptic medications or surgery, and addressing a variety of psychological and social issues. Treatment plans must be individualized, given the many different types and causes of seizures as well as the differences in efficacy and toxicity of antiepileptic medications for each patient. In almost all cases, a neurologist with experience in the treatment of epilepsy should design and oversee implementation of the treatment strategy. Furthermore, patients with refractory epilepsy or those who require polypharmacy with antiepileptic drugs should remain under the regular care of a neurologist.

TREATMENT OF UNDERLYING CONDITIONS

If the sole cause of a seizure is a metabolic disturbance such as an abnormality of serum electrolytes or glucose, then treatment is aimed at reversing the metabolic problem and preventing its recurrence. Therapy with antiepileptic drugs is usually unnecessary unless the metabolic disorder cannot be corrected promptly and the patient is at risk of having further seizures. If the apparent cause of a seizure was a medication (e.g., theophylline) or illicit drug use (e.g., cocaine), then appropriate therapy is avoidance of the drug; there is usually no need for antiepileptic medications unless subsequent seizures occur in the absence of these precipitants.

Seizures caused by a structural CNS lesion such as a brain tumor, vascular malformation, or brain abscess may not recur after appropriate treatment of the underlying lesion. However, despite removal of the structural lesion, there is a risk that the seizure focus will remain in the surrounding tissue or develop *de novo* as a result of gliosis and other processes induced by surgery, radiation, or other therapies. Most patients are therefore maintained on an antiepileptic medication for at least 1 year, and an attempt is made to withdraw medications only if the patient has been completely seizure free. If seizures are refractory to medication, the patient may benefit from surgical removal of the epileptic brain region (see below).

AVOIDANCE OF PRECIPITATING FACTORS

Unfortunately, little is known about the specific factors that determine precisely when a seizure will occur in a patient with epilepsy. Some patients can identify particular situations that appear to lower their seizure threshold; these situations should be avoided. For example, a patient who has seizures in the setting of sleep deprivation should obviously be advised to maintain a normal sleep schedule. Many patients note an association between alcohol

intake and seizures, and they should be encouraged to modify their drinking habits accordingly. There are also relatively rare cases of patients with seizures that are induced by highly specific stimuli such as a video game monitor, music, or an individual's voice ("reflex epilepsy"). Because there is often an association between stress and seizures, stress reduction techniques such as physical exercise, meditation, or counseling may be helpful.

ANTIEPILEPTIC DRUG THERAPY

Antiepileptic drug therapy is the mainstay of treatment for most patients with epilepsy. The overall goal is to completely prevent seizures without causing any untoward side effects, preferably with a single medication and a dosing schedule that is easy for the patient to follow. Seizure classification is an important element in designing the treatment plan, because some antiepileptic drugs have different activities against various seizure types. However, there is considerable overlap between many antiepileptic drugs such that the choice of therapy is often determined more by the patient's specific needs, especially his or her assessment of side effects.

When to Initiate Antiepileptic Drug Therapy Antiepileptic drug therapy should be started in any patient with recurrent seizures of unknown etiology or a known cause that cannot be reversed. Whether to initiate therapy in a patient with a single seizure is controversial. Patients with a single seizure due to an identified lesion such as a CNS tumor, infection, or trauma, in which there is strong evidence that the lesion is epileptogenic, should be treated. The risk of seizure recurrence in a patient with an apparently unprovoked or idiopathic seizure is uncertain, with estimates ranging from 31 to 71% in the first 12 months after the initial seizure. This uncertainty arises from differences in the underlying seizure types and etiologies in various published epidemiologic studies. Generally accepted risk factors associated with recurrent seizures include the following: (1) an abnormal neurologic examination, (2) seizures presenting as status epilepticus, (3) postictal Todd's paralysis, (4) a strong family history of seizures, or (5) an abnormal EEG. Most patients with one or more of these risk factors should be treated. Issues such as employment or driving may influence the decision whether to start medications as well. For example, a patient with a single, idiopathic seizure whose job depends on driving may prefer taking antiepileptic drugs rather than risk a seizure recurrence and the potential loss of driving privileges.

Selection of Antiepileptic Drugs Antiepileptic drugs available in the United States are shown in [Table 445-8](#), and the main

pharmacologic characteristics of commonly used drugs are listed in [Table 445-9](#). Worldwide, older medications such as phenytoin, valproic acid, carbamazepine, phenobarbital, and ethosuximide are generally used as first-line therapy for most seizure disorders because, overall, they are as effective as recently marketed drugs and significantly less expensive overall. Most of the new drugs that have become available in the past decade are used as add-on or alternative therapy, although many are now being used as first-line monotherapy.

In addition to efficacy, factors influencing the choice of an initial medication include the convenience of dosing (e.g., once daily versus three or four times daily) and potential side effects. In this regard, a number of the newer drugs have the advantage of reduced drug-drug interactions and easier dosing. Almost all of the commonly used antiepileptic drugs can cause similar, dose-related side effects such as sedation, ataxia, and diplopia. Long-term use of some agents in adults, especially the elderly, can lead to osteoporosis. Close follow-up is required to ensure these side effects are promptly recognized and reversed. Most of the older drugs and some of the newer ones can also cause idiosyncratic toxicity such as rash, bone marrow suppression, or hepatotoxicity. Although rare, these side effects should be considered during drug selection, and patients must be instructed about symptoms or signs that should signal the need to alert their health care provider. For some drugs, laboratory tests (e.g., complete blood count and liver function tests) are recommended prior to the institution of therapy (to establish baseline values) and during initial dosing and titration of the agent. Importantly, studies have shown that Asian individuals carrying the human leukocyte antigen allele, HLA-B*1502, are at particularly high risk of developing serious skin reactions from carbamazepine and phenytoin. As a result, racial background and genotype are additional factors to consider in drug selection.

ANTIEPILEPTIC DRUG SELECTION FOR FOCAL SEIZURES Carbamazepine (or a related drug, oxcarbazepine), lamotrigine, phenytoin, and levetiracetam are currently the drugs of choice approved for the initial treatment of focal seizures, including those that evolve into generalized seizures. Overall they have very similar efficacy, but differences in pharmacokinetics and toxicity are the main determinants for use in a given patient. For example, an advantage of carbamazepine (which is also available in an extended-release form) is that its metabolism follows first-order pharmacokinetics, which allows for a linear relationship between drug dose, serum levels, and toxicity. Carbamazepine can cause leukopenia, aplastic anemia, or hepatotoxicity and would therefore be contraindicated in patients with predispositions to these problems. Oxcarbazepine has the advantage of being metabolized in a way that avoids an intermediate metabolite associated with some of the side effects of carbamazepine. Oxcarbazepine also has fewer drug interactions than carbamazepine. Lamotrigine tends to be well tolerated in terms of side effects. However, patients need to be particularly vigilant about the possibility of a skin rash during the initiation of therapy. This can be extremely severe and lead to Stevens-Johnson syndrome if unrecognized and if the medication is not discontinued immediately. This risk can be reduced by the use of low initial doses and slow titration. Lamotrigine must be started at lower initial doses when used as add-on therapy with valproic acid, because valproic acid inhibits lamotrigine metabolism and results in a substantially prolonged half-life. Phenytoin has a relatively long half-life and offers the advantage of once or twice daily dosing compared to two or three times daily dosing for many of the other drugs. However, phenytoin shows properties of nonlinear kinetics, such that small increases in phenytoin doses above a standard maintenance dose can precipitate marked side effects. This is one of the main causes of acute phenytoin toxicity. Long-term use of phenytoin is associated with untoward cosmetic effects (e.g., hirsutism, coarsening of facial features, gingival hypertrophy) and effects on bone metabolism. Due to these side effects, phenytoin is often avoided in young patients who are likely to require the drug for many years. Levetiracetam has

TABLE 445-8 SELECTION OF ANTIEPILEPTIC DRUGS

Generalized-Onset Tonic-Clonic	Focal	Typical Absence	Atypical Absence, Myoclonic, Atonic
First-Line			
Lamotrigine	Lamotrigine	Valproic acid	Valproic acid
Valproic acid	Carbamazepine	Ethosuximide	Lamotrigine
	Oxcarbazepine	Lamotrigine	Topiramate
	Phenytoin		
	Levetiracetam		
Alternatives			
Zonisamide ^a	Topiramate	Lamotrigine	Clonazepam
Phenytoin	Zonisamide ^a	Clonazepam	Felbamate
Carbamazepine	Valproic acid		Clobazam
Oxcarbazepine	Tiagabine ^a		Rufinamide
Topiramate	Gabapentin ^a		
Phenobarbital	Lacosamide ^a		
Primidone	Exogabine ^a		
Felbamate	Phenobarbital		
	Primidone		
	Felbamate		

^aAs adjunctive therapy.

TABLE 445-9 DOSAGE AND ADVERSE EFFECTS OF COMMONLY USED ANTIEPILEPTIC DRUGS

Generic Name	Trade Name	Principal Uses	Typical Dose; Dose Interval	Half-Life	Therapeutic Range	Adverse Effects		
						Neurologic	Systemic	Drug Interactions ^a
Carbamazepine	Tegretol ^c	Tonic-clonic Focal-onset	600–1800 mg/d (15–35 mg/kg, child); bid (capsules or tablets), tid-qid (oral suspension)	10–17 h (variable due to autoinduction: complete 3–5 wk after initiation)	4–12 µg/mL	Ataxia Dizziness Diplopia Vertigo	Aplastic anemia Leukopenia Gastrointestinal irritation Hepatotoxicity Hyponatremia	Level decreased by enzyme-inducing drugs ^b Level increased by erythromycin, propoxyphene, isoniazid, cimetidine, fluoxetine
Clobazam	Onfi	Lennox-Gastaut syndrome	10–40 mg/d (5–20 mg/d for patients <30 kg body weight); bid	36–42 h (71–82 h for less active metabolite)	Not established	Fatigue Sedation Ataxia Aggression Insomnia	Constipation Anorexia Skin rash	Level increased by CYP2C19 inhibitors
Clonazepam	Klonopin	Absence Atypical absence Myoclonic	1–12 mg/d; qd-tid	24–48 h	10–70 ng/mL	Ataxia Sedation Lethargy	Anorexia	Level decreased by enzyme-inducing drugs ^b
Ethosuximide	Zarontin	Absence	750–1250 mg/d (20–40 mg/kg); qd-bid	60 h, adult 30 h, child	40–100 µg/mL	Ataxia Lethargy Headache	Gastrointestinal irritation Skin rash Bone marrow suppression	Level decreased by enzyme-inducing drugs ^b Level increased by valproic acid
Ezogabine	Potiga	Focal-onset	800–1200 mg/d; tid	7–11 h	Not established	Dizziness Fatigue Sedation Confusion Vertigo Tremor	Retinal abnormalities Skin discoloration Cardiac conduction (QT interval prolongation) Urinary retention	Level decreased by enzyme-inducing drugs ^b
Felbamate	Felbatol	Focal-onset Lennox-Gastaut syndrome Tonic-clonic	2400–3600 mg/d, tid-qid	16–22 h	30–60 µg/mL	Insomnia Dizziness Sedation Headache	Aplastic anemia Hepatic failure Weight loss Gastrointestinal irritation	Increases phenytoin, valproic acid, active carbamazepine metabolite
Gabapentin	Neurontin	Focal-onset	900–2400 mg/d; tid-qid	5–9 h	2–20 µg/mL	Sedation Dizziness Ataxia Fatigue	Gastrointestinal irritation Weight gain Edema	No known significant interactions
Lacosamide	Vimpat	Focal-onset	200–400 mg/d; bid	13 h	Not established	Dizziness Ataxia Diplopia Vertigo	Gastrointestinal irritation Cardiac conduction (PR interval prolongation)	Level decreased by enzyme-inducing drugs ^b

(Continued)

TABLE 445-9 DOSAGE AND ADVERSE EFFECTS OF COMMONLY USED ANTIEPILEPTIC DRUGS (CONTINUED)

Generic Name	Trade Name	Principal Uses	Typical Dose; Dose Interval	Half-Life	Therapeutic Range	Adverse Effects		Drug Interactions ^a
						Neurologic	Systemic	
Lamotrigine	Lamictal ^c	Focal-onset Tonic-clonic Atypical absence Myoclonic Lennox-Gastaut syndrome	150–500 mg/d; bid (immediate release), daily (extended release) (lower daily dose for regimens with valproic acid; higher daily dose for regimens with an enzyme inducer)	25 h 14 h (with enzyme-inducers), 59 h (with valproic acid)	2.5–20 µg/mL	Dizziness Diplopia Sedation Ataxia Headache	Skin rash Stevens-Johnson syndrome	Level decreased by enzyme-inducing drugs ^b and oral contraceptives Level increased by valproic acid
Levetiracetam	Keppra ^c	Focal-onset	1000–3000 mg/d; bid (immediate release), daily (extended release)	6–8 h	5–45 µg/mL	Sedation Fatigue Incoordination Mood changes	Anemia Leukopenia	No known significant interactions
Oxcarbazepine ^c	Trileptal	Focal-onset Tonic-clonic	900–2400 mg/d (30–45 mg/kg, child); bid	10–17 h (for active metabolite)	10–35 µg/mL	Fatigue Ataxia Dizziness Diplopia Vertigo Headache	See carbamazepine	Level decreased by enzyme-inducing drugs ^b May increase phenytoin
Phenobarbital	Luminal	Tonic-clonic Focal-onset	60–180 mg/d; qd-tid	90 h	10–40 µg/mL	Sedation Ataxia Confusion Dizziness Decreased libido Depression	Skin rash	Level increased by valproic acid, phenytoin
Phenytoin (diphenylhydantoin)	Dilantin	Tonic-clonic Focal-onset	300–400 mg/d (3–6 mg/kg, adult; 4–8 mg/kg, child); qd-tid	24 h (wide variation, dose-dependent)	10–20 µg/mL	Dizziness Diplopia Ataxia Incoordination Confusion	Gingival hyperplasia Lymphadenopathy Hirsutism Osteomalacia Facial coarsening Skin rash	Level increased by isoniazid, sulfonamides, fluoxetine Level decreased by enzyme-inducing drugs ^b Altered folate metabolism
Primidone	Mysoline	Tonic-clonic Focal-onset	750–1000 mg/d; bid-tid	Primidone, 8–15 h Phenobarbital, 90 h	Primidone, 4–12 µg/mL Phenobarbital, 10–40 µg/mL	Same as phenobarbital		Level increased by valproic acid Level decreased by phenytoin (increased conversion to phenobarbital)

Rufinamide	Banzel	Lennox-Gastaut syndrome	3200 mg/d (45 mg/kg, child); bid	6–10 h	Not established	Sedation Fatigue Dizziness Ataxia Headache Diplopia	Gastrointestinal irritation Leukopenia Cardiac conduction (QT interval shortening)	Level decreased by enzyme-inducing drugs ^b Level increased by valproic acid May increase phenytoin
Tiagabine	Gabitril	Focal-onset	32–56 mg/d; bid-qid (as adjunct to enzyme-inducing antiepileptic drug regimen)	2–5 h (with enzyme inducer), 7–9 h (without enzyme inducer)	Not established	Confusion Sedation Depression Dizziness Speech or language problems Paresthesias Psychosis	Gastrointestinal irritation	Level decreased by enzyme-inducing drugs ^b
Topiramate	Topamax	Focal-onset Tonic-clonic Lennox-Gastaut syndrome	200–400 mg/d; bid (immediate release), daily (extended release)	20 h (immediate release), 30 h (extended release)	2–20 µg/mL	Psychomotor slowing Sedation Speech or language problems Fatigue Paresthesias	Renal stones (avoid use with other carbonic anhydrase inhibitors) Glaucoma Weight loss Hypohidrosis	Level decreased by enzyme-inducing drugs ^b
Valproic acid (valproate sodium, divalproex sodium)	Depakene Depakote ^c	Tonic-clonic Absence Atypical absence Myoclonic Focal-onset Atonic	750–2000 mg/d (20–60 mg/kg); bid-qid (immediate and delayed release), daily (extended release)	15 h	50–125 µg/mL	Ataxia Sedation Tremor	Hepatotoxicity Thrombocytopenia Gastrointestinal irritation Weight gain Transient alopecia Hyperammonemia	Level decreased by enzyme-inducing drugs ^b
Zonisamide	Zonegran	Focal-onset Tonic-clonic	200–400 mg/d; qd-bid	50–68 h	10–40 µg/mL	Sedation Dizziness Confusion Headache Psychosis	Anorexia Renal stones Hypohidrosis	Level decreased by enzyme-inducing drugs ^b

^aExamples only; please refer to other sources for comprehensive listings of all potential drug-drug interactions. ^bPhenytoin, carbamazepine, phenobarbital. ^cExtended-release product available.

the advantage of having no known drug-drug interactions, making it especially useful in the elderly and patients on other medications. However, a significant number of patients taking levetiracetam complain of irritability, anxiety, and other psychiatric symptoms. Topiramate can be used for both focal and generalized seizures. Similar to some of the other antiepileptic drugs, topiramate can cause significant psychomotor slowing and other cognitive problems. Additionally, it should not be used in patients at risk for the development of glaucoma or renal stones.

Valproic acid is an effective alternative for some patients with focal seizures, especially when the seizures generalize. Gastrointestinal side effects are fewer when using the delayed-release formulation (Depakote). Laboratory testing is required to monitor toxicity because valproic acid can rarely cause reversible bone marrow suppression and hepatotoxicity. This drug should generally be avoided in patients with preexisting bone marrow or liver disease. Irreversible, fatal hepatic failure appearing as an idiosyncratic rather than dose-related side effect is a relatively rare complication; its risk is highest in children <2 years old, especially those taking other antiepileptic drugs or with inborn errors of metabolism.

Zonisamide, tiagabine, gabapentin, lacosamide, and ezogabine are additional drugs currently used for the treatment of focal seizures with or without evolution into generalized seizures. Phenobarbital and other barbiturate compounds were commonly used in the past as first-line therapy for many forms of epilepsy. However, the barbiturates frequently cause sedation in adults, hyperactivity in children, and other more subtle cognitive changes; thus, their use should be limited to situations in which no other suitable treatment alternatives exist.

ANTIEPILEPTIC DRUG SELECTION FOR GENERALIZED SEIZURES Lamotrigine and valproic acid are currently considered the best initial choice for the treatment of primary generalized, tonic-clonic seizures. Topiramate, zonisamide, phenytoin, carbamazepine, and oxcarbazepine are suitable alternatives. Valproic acid is also particularly effective in absence, myoclonic, and atonic seizures. It is therefore the drug of choice in patients with generalized epilepsy syndromes having mixed seizure types. Importantly, carbamazepine, oxcarbazepine, and phenytoin can worsen certain types of generalized seizures, including absence, myoclonic, tonic, and atonic seizures. Ethosuximide is a particularly effective drug for the treatment of uncomplicated absence seizures, but it is not useful for tonic-clonic or focal seizures. Periodic monitoring of blood cell counts is required since ethosuximide rarely causes bone marrow suppression. Lamotrigine appears to be particularly effective in epilepsy syndromes with mixed, generalized seizure types such as JME and Lennox-Gastaut syndrome. Topiramate, zonisamide, and felbamate may have similar broad efficacy.

INITIATION AND MONITORING OF THERAPY

Because the response to any antiepileptic drug is unpredictable, patients should be carefully educated about the approach to therapy. The goal is to prevent seizures and minimize the side effects of treatment; determination of the optimal dose is often a matter of trial and error. This process may take months or longer if the baseline seizure frequency is low. Most antiepileptic drugs need to be introduced relatively slowly to minimize side effects. Patients should expect that minor side effects such as mild sedation, slight changes in cognition, or imbalance will typically resolve within a few days. Starting doses are usually the lowest value listed under the dosage column in Table 445-9. Subsequent increases should be made only after achieving a steady state with the previous dose (i.e., after an interval of five or more half-lives).

Monitoring of serum antiepileptic drug levels can be very useful for establishing the initial dosing schedule. However, the published therapeutic ranges of serum drug concentrations are only an approximate guide for determining the proper dose for a given patient. The key determinants are the clinical measures of seizure frequency and presence of side effects, not the laboratory values. Conventional assays of serum drug levels measure the total drug (i.e., both free and protein bound). However, it is the concentration of

free drug that reflects extracellular levels in the brain and correlates best with efficacy. Thus, patients with decreased levels of serum proteins (e.g., decreased serum albumin due to impaired liver or renal function) may have an increased ratio of free to bound drug, yet the concentration of free drug may be adequate for seizure control. These patients may have a “subtherapeutic” drug level, but the dose should be changed only if seizures remain uncontrolled, not just to achieve a “therapeutic” level. It is also useful to monitor free drug levels in such patients. In practice, other than during the initiation or modification of therapy, monitoring of antiepileptic drug levels is most useful for documenting adherence.

If seizures continue despite gradual increases to the maximum tolerated dose and documented compliance, then it becomes necessary to switch to another antiepileptic drug. This is usually done by maintaining the patient on the first drug while a second drug is added. The dose of the second drug should be adjusted to decrease seizure frequency without causing toxicity. Once this is achieved, the first drug can be gradually withdrawn (usually over weeks unless there is significant toxicity). The dose of the second drug is then further optimized based on seizure response and side effects. Monotherapy should be the goal whenever possible.

WHEN TO DISCONTINUE THERAPY

Overall, about 70% of children and 60% of adults who have their seizures completely controlled with antiepileptic drugs can eventually discontinue therapy. The following patient profile yields the greatest chance of remaining seizure free after drug withdrawal: (1) complete medical control of seizures for 1–5 years; (2) single seizure type, either focal or generalized; (3) normal neurologic examination, including intelligence; and (4) normal EEG. The appropriate seizure-free interval is unknown and undoubtedly varies for different forms of epilepsy. However, it seems reasonable to attempt withdrawal of therapy after 2 years in a patient who meets all of the above criteria, is motivated to discontinue the medication, and clearly understands the potential risks and benefits. In most cases, it is preferable to reduce the dose of the drug gradually over 2–3 months. Most recurrences occur in the first 3 months after discontinuing therapy, and patients should be advised to avoid potentially dangerous situations such as driving or swimming during this period.

TREATMENT OF REFRACTORY EPILEPSY

Approximately one-third of patients with epilepsy do not respond to treatment with a single antiepileptic drug, and it becomes necessary to try a combination of drugs to control seizures. Patients who have focal epilepsy related to an underlying structural lesion or those with multiple seizure types and developmental delay are particularly likely to require multiple drugs. There are currently no clear guidelines for rational polypharmacy, although in theory a combination of drugs with different mechanisms of action may be most useful. In most cases, the initial combination therapy combines first-line drugs (i.e., carbamazepine, oxcarbazepine, lamotrigine, valproic acid, levetiracetam, and phenytoin). If these drugs are unsuccessful, then the addition of other drugs such as topiramate, zonisamide, lacosamide, or tiagabine is indicated. Patients with myoclonic seizures resistant to valproic acid may benefit from the addition of clonazepam or clobazam, and those with absence seizures may respond to a combination of valproic acid and ethosuximide. The same principles concerning the monitoring of therapeutic response, toxicity, and serum levels for monotherapy apply to polypharmacy, and potential drug interactions need to be recognized. If there is no improvement, a third drug can be added while the first two are maintained. If there is a response, the less effective or less well tolerated of the first two drugs should be gradually withdrawn.

SURGICAL TREATMENT OF REFRACTORY EPILEPSY

Approximately 20–30% of patients with epilepsy continue to have seizures despite efforts to find an effective combination of antiepileptic drugs. For some, surgery can be extremely effective in substantially reducing seizure frequency and even providing complete seizure control. Understanding the potential value of surgery

is especially important when a patient's seizures are not controlled with initial treatment, as such patients often do not respond to subsequent medication trials. Rather than submitting the patient to years of unsuccessful medical therapy and the psychosocial trauma and increased mortality associated with ongoing seizures, the patient should have an efficient but relatively brief attempt at medical therapy and then be referred for surgical evaluation.

The most common surgical procedure for patients with temporal lobe epilepsy involves resection of the anteromedial temporal lobe (temporal lobectomy) or a more limited removal of the underlying hippocampus and amygdala (amygdalohippocampectomy). Focal seizures arising from extratemporal regions may be abolished by a focal neocortical resection with precise removal of an identified lesion (lesionectomy). Localized neocortical resection without a clear lesion identified on MRI is also possible when other tests (e.g. MEG, PET, SPECT) implicate a focal cortical region as a seizure onset zone. When the cortical region cannot be removed, multiple subpial transection, which disrupts intracortical connections, is sometimes used to prevent seizure spread. Hemispherectomy or multilobar resection is useful for some patients with severe seizures due to hemispheric abnormalities such as hemimegalencephaly or other dysplastic abnormalities, and corpus callosotomy has been shown to be effective for disabling tonic or atonic seizures, usually when they are part of a mixed-seizure syndrome (e.g., Lennox-Gastaut syndrome).

Presurgical evaluation is designed to identify the functional and structural basis of the patient's seizure disorder. Inpatient video-EEG monitoring is used to define the anatomic location of the seizure focus and to correlate the abnormal electrophysiologic activity with behavioral manifestations of the seizure. Routine scalp or scalp-sphenoidal recordings and a high-resolution MRI scan are usually sufficient for localization of the epileptogenic focus, especially when the findings are concordant. Functional imaging studies such as SPECT, PET, and MEG are adjunctive tests that may help to reveal or verify the localization of an apparent epileptogenic region. Once the presumed location of the seizure onset is identified, additional studies, including neuropsychological testing, the intracarotid amobarbital test (Wada test), and functional MRI may be used to assess language and memory localization and to determine the possible functional consequences of surgical removal of the epileptogenic region. In some cases, standard noninvasive evaluation is not sufficient to localize the seizure onset zone, and invasive electrophysiologic monitoring, such as implanted depth or subdural electrodes, is required for more definitive localization. The exact extent of the resection to be undertaken can also be determined by performing cortical mapping at the time of the surgical procedure, allowing for a tailored resection. This involves electrocorticographic recordings made with electrodes on the surface of the brain to identify the extent of epileptiform disturbances. If the region to be resected is within or near brain regions suspected of having sensorimotor or language function, electrical cortical stimulation mapping is performed on the awake patient to determine the function of cortical regions in question in order to avoid resection of so-called eloquent cortex and thereby minimize postsurgical deficits.

Advances in presurgical evaluation and microsurgical techniques have led to a steady increase in the success of epilepsy surgery. Clinically significant complications of surgery are <5%, and the use of functional mapping procedures has markedly reduced the neurologic sequelae due to removal or sectioning of brain tissue. For example, about 70% of patients treated with temporal lobectomy will become seizure free, and another 15–25% will have at least a 90% reduction in seizure frequency. Marked improvement is also usually seen in patients treated with hemispherectomy for catastrophic seizure disorders due to large hemispheric abnormalities. Postoperatively, patients generally need to remain on antiepileptic drug therapy, but the marked reduction of seizures following resective surgery can have a very beneficial effect on quality of life.

Not all medically refractory patients are suitable candidates for resective surgery. For example, some patients have seizures arising

from more than one location, making the risk of ongoing seizures or potential harm from the surgery unacceptably high. Vagus nerve stimulation (VNS) has been used in some of these cases, although the results are limited and it is difficult to predict who will benefit. A new implantable device that can detect the onset of a seizure (in some instances before the seizure becomes clinically apparent) and deliver an electrical stimulation (Responsive NeuroStimulation) has recently been approved and may be of benefit in selected patients. Studies are currently evaluating the efficacy of stereotactic radiosurgery, laser thermoablation, and deep brain stimulation (DBS) as other options for surgical treatment of refractory epilepsy.

STATUS EPILEPTICUS

Status epilepticus refers to continuous seizures or repetitive, discrete seizures with impaired consciousness in the interictal period. Status epilepticus has numerous subtypes, including generalized convulsive status epilepticus (GCSE) (e.g., persistent, generalized electrographic seizures, coma, and tonic-clonic movements) and nonconvulsive status epilepticus (e.g., persistent absence seizures or focal seizures with confusion or partially impaired consciousness, and minimal motor abnormalities). The duration of seizure activity sufficient to meet the definition of status epilepticus has traditionally been specified as 15–30 min. However, a more practical definition is to consider status epilepticus as a situation in which the duration of seizures prompts the acute use of anticonvulsant therapy. For GCSE, this is typically when seizures last beyond 5 min.

GCSE is an emergency and must be treated immediately, because cardiorespiratory dysfunction, hyperthermia, and metabolic derangements can develop as a consequence of prolonged seizures, and these can lead to irreversible neuronal injury. Furthermore, CNS injury can occur even when the patient is paralyzed with neuromuscular blockade but continues to have electrographic seizures. The most common causes of GCSE are anticonvulsant withdrawal or noncompliance, metabolic disturbances, drug toxicity, CNS infection, CNS tumors, refractory epilepsy, and head trauma.

GCSE is obvious when the patient is having overt convulsions. However, after 30–45 min of uninterrupted seizures, the signs may become increasingly subtle. Patients may have mild clonic movements of only the fingers or fine, rapid movements of the eyes. There may be paroxysmal episodes of tachycardia, hypertension, and pupillary dilation. In such cases, the EEG may be the only method of establishing the diagnosis. Thus, if the patient stops having overt seizures, yet remains comatose, an EEG should be performed to rule out ongoing status epilepticus. This is obviously also essential when a patient with GCSE has been paralyzed with neuromuscular blockade in the process of protecting the airway.

The first steps in the management of a patient in GCSE are to attend to any acute cardiorespiratory problems or hyperthermia, perform a brief medical and neurologic examination, establish venous access, and send samples for laboratory studies to identify metabolic abnormalities. Anticonvulsant therapy should then begin without delay; a treatment approach is shown in [Fig. 445-3](#).

The treatment of nonconvulsive status epilepticus is thought to be less urgent than GCSE, because the ongoing seizures are not accompanied by the severe metabolic disturbances seen with GCSE. However, evidence suggests that nonconvulsive status epilepticus, especially that caused by ongoing, focal seizure activity, is associated with cellular injury in the region of the seizure focus; therefore this condition should be treated as promptly as possible using the general approach described for GCSE.

BEYOND SEIZURES: OTHER MANAGEMENT ISSUES

INTERICTAL BEHAVIOR

The adverse effects of epilepsy often go beyond clinical seizures, and the extent of these effects largely depends on the etiology of epilepsy, seizure frequency and severity, and side effects from antiepileptic therapy. Many epilepsy patients are completely normal between seizures

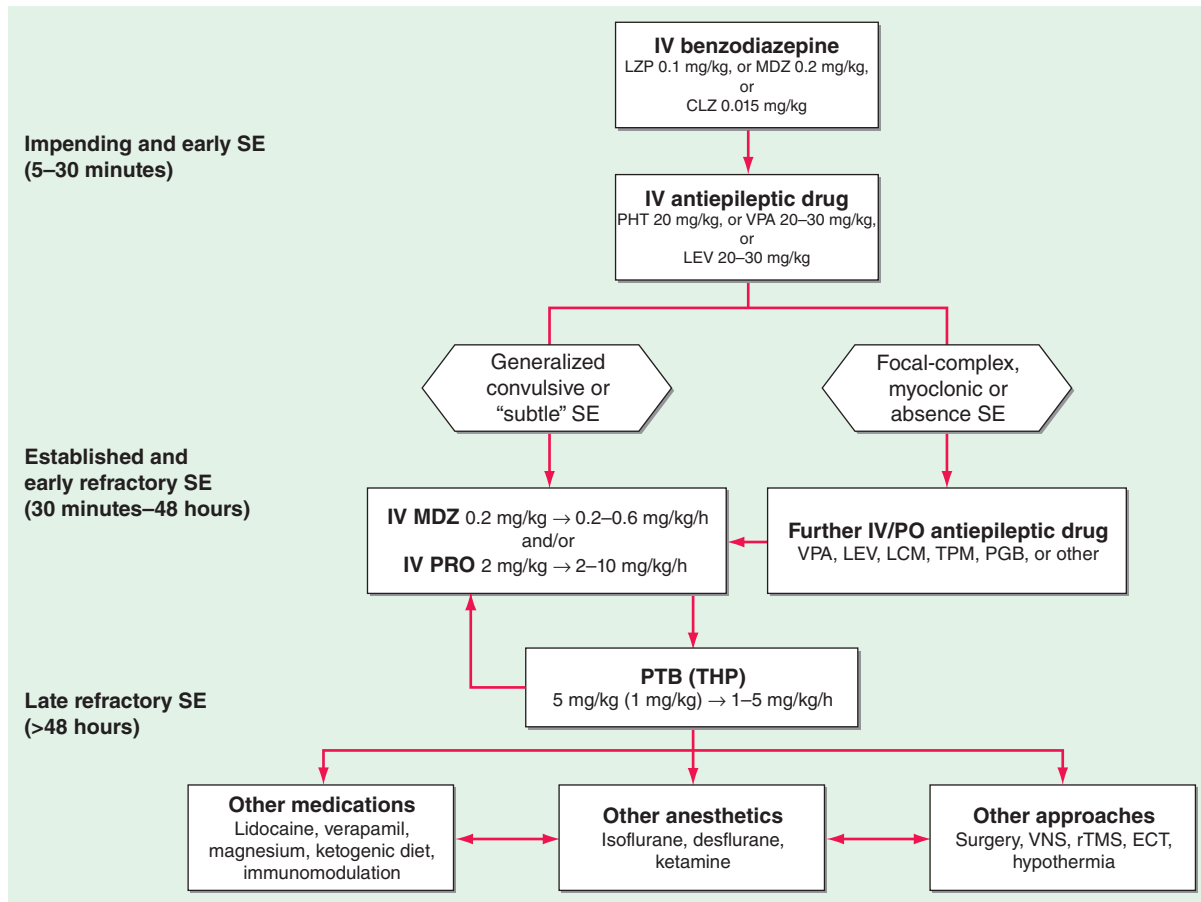


FIGURE 445-3 Pharmacologic treatment of generalized tonic-clonic status epilepticus (SE) in adults. CLZ, clonazepam; ECT, electroconvulsive therapy; LCM, lacosamide; LEV, levetiracetam; LZP, lorazepam; MDZ, midazolam; PGB, pregabalin; PHT, phenytoin or fos-phenytoin; PRO, propofol; PTB, pentobarbital; rTMS, repetitive transcranial magnetic stimulation; THP, thiopental; TPM, topiramate; VNS, vagus nerve stimulation; VPA, valproic acid. (From AO Rossetti, DH Lowenstein: *Lancet Neurol* 10:922, 2011.)

and live highly successful and productive lives. In contrast, patients with seizures secondary to developmental abnormalities or acquired brain injury may have impaired cognitive function and other neurologic deficits. Frequent interictal EEG abnormalities are associated with subtle dysfunction of memory and attention. Patients with many seizures, especially those emanating from the temporal lobe, often note an impairment of short-term memory that may progress over time.

Patients with epilepsy are at risk of developing a variety of psychiatric problems, including depression, anxiety, and psychosis. This risk varies considerably depending on many factors, including the etiology, frequency, and severity of seizures and the patient's age and previous personal or family history of psychiatric disorder. Depression occurs in ~20% of patients, and the incidence of suicide is higher in patients with epilepsy than in the general population. Depression should be treated through counseling or medication. The selective serotonin reuptake inhibitors (SSRIs) typically have minimal effect on seizures, whereas tricyclic antidepressants may lower the seizure threshold. Anxiety can be a seizure symptom, and anxious or psychotic behavior can occur during a postictal delirium. Postictal psychosis is a rare phenomenon that typically occurs after a period of increased seizure frequency. There is usually a brief lucid interval lasting up to a week, followed by days to weeks of agitated, psychotic behavior. The psychosis usually resolves spontaneously but frequently will require short-term treatment with antipsychotic or anxiolytic medications.

There is ongoing controversy as to whether some patients with epilepsy (especially temporal lobe epilepsy) have a stereotypical “interictal personality.” The predominant view is that atypical personality traits occur in diverse epilepsies (e.g., generalized and frontal lobe epilepsy) and may result from an underlying structural brain lesion, antiepileptic drug effects, and psychosocial factors related to suffering from a chronic disease, as well as the epilepsy itself.

MORTALITY OF EPILEPSY

Patients with epilepsy have a risk of death that is roughly two to three times greater than expected in a matched population without epilepsy. Most of the increased mortality is due to the underlying etiology of epilepsy (e.g., tumors or strokes in older adults). However, a significant number of patients die from accidents, status epilepticus, and a syndrome known as *sudden unexpected death in epilepsy* (SUDEP), which usually affects young people with convulsive seizures and tends to occur at night. The cause of SUDEP is unknown; it may result from brainstem-mediated effects of seizures on pulmonary, cardiac, and arousal functions. Recent studies suggest that, in some cases, a genetic mutation may be the cause of both epilepsy and a cardiac conduction defect that gives rise to sudden death.

PSYCHOSOCIAL ISSUES

There continues to be a cultural stigma about epilepsy, although it is slowly declining in societies with effective health education programs. Many patients with epilepsy harbor fears such as the fear of becoming mentally retarded or dying during a seizure. These issues need to be carefully addressed by educating the patient about epilepsy and by ensuring that family members, teachers, fellow employees, and other associates are equally well informed. A useful source of educational material is the Web site www.epilepsy.com.

EMPLOYMENT, DRIVING, AND OTHER ACTIVITIES

Many patients with epilepsy face difficulty in obtaining or maintaining employment, even when their seizures are well controlled. Federal and state legislation is designed to prevent employers from discriminating against patients with epilepsy, and patients should be encouraged to understand and claim their legal rights. Patients in these circumstances

also benefit greatly from the assistance of health providers who act as strong patient advocates.

Loss of driving privileges is one of the most disruptive social consequences of epilepsy. Physicians should be very clear about local regulations concerning driving and epilepsy, because the laws vary considerably among states and countries. In all cases, it is the physician's responsibility to warn patients of the danger imposed on themselves and others while driving if their seizures are uncontrolled (unless the seizures are not associated with impairment of consciousness or motor control). In general, most states allow patients to drive after a seizure-free interval (on or off medications) of between 3 months and 2 years.

Patients with incompletely controlled seizures must also contend with the risk of being in other situations where an impairment of consciousness or loss of motor control could lead to major injury or death. Thus, depending on the type and frequency of seizures, many patients need to be instructed to avoid working at heights or with machinery or to have someone close by for activities such as bathing and swimming.

SPECIAL ISSUES RELATED TO WOMEN AND EPILEPSY

CATAMENIAL EPILEPSY

Some women experience a marked increase in seizure frequency around the time of menses. This is believed to be mediated by either the effects of estrogen and progesterone on neuronal excitability or changes in antiepileptic drug levels due to altered protein binding or metabolism. Some patients may benefit from increases in antiepileptic drug dosages during menses. Natural progestins or intramuscular medroxyprogesterone may be of benefit to a subset of women.

PREGNANCY

Most women with epilepsy who become pregnant will have an uncomplicated gestation and deliver a normal baby. However, epilepsy poses some important risks to a pregnancy. Seizure frequency during pregnancy will remain unchanged in ~50% of women, increase in 30%, and decrease in 20%. Changes in seizure frequency are attributed to endocrine effects on the CNS, variations in antiepileptic drug pharmacokinetics (such as acceleration of hepatic drug metabolism or effects on plasma protein binding), and changes in medication compliance. It is useful to see patients at frequent intervals during pregnancy and monitor serum antiepileptic drug levels. Measurement of the unbound drug concentrations may be useful if there is an increase in seizure frequency or worsening of side effects of antiepileptic drugs.

The overall incidence of fetal abnormalities in children born to mothers with epilepsy is 5–6%, compared to 2–3% in healthy women. Part of the higher incidence is due to teratogenic effects of antiepileptic drugs, and the risk increases with the number of medications used (e.g., 10–20% risk of malformations with three drugs) and possibly with higher doses. A meta-analysis of published pregnancy registries and cohorts found that the most common malformations were defects in the cardiovascular and musculoskeletal system (1.4–1.8%). Valproic acid is strongly associated with an increased risk of adverse fetal outcomes (7–20%). Recent findings from a large pregnancy registry suggest that, other than topiramate, the newer antiepileptic drugs are far safer than valproic acid.

Because the potential harm of uncontrolled convulsive seizures on the mother and fetus is considered greater than the teratogenic effects of antiepileptic drugs, it is currently recommended that pregnant women be maintained on effective drug therapy. When possible, it seems prudent to have the patient on monotherapy at the lowest effective dose, especially during the first trimester. For some women, however, the type and frequency of their seizures may allow for them to safely wean off antiepileptic drugs prior to conception. Patients should also take folate (1–4 mg/d), because the antifolate effects of anticonvulsants are thought to play a role in the development of neural tube defects, although the benefits of this treatment remain unproved in this setting.

Enzyme-inducing drugs such as phenytoin, carbamazepine, oxcarbazepine, topiramate, phenobarbital, and primidone cause a transient and reversible deficiency of vitamin K–dependent clotting factors in ~50% of newborn infants. Although neonatal hemorrhage

is uncommon, the mother should be treated with oral vitamin K (20 mg/d, phylloquinone) in the last 2 weeks of pregnancy, and the infant should receive intramuscular vitamin K (1 mg) at birth.

CONTRACEPTION

Special care should be taken when prescribing antiepileptic medications for women who are taking oral contraceptive agents. Drugs such as carbamazepine, phenytoin, phenobarbital, and topiramate can significantly decrease the efficacy of oral contraceptives via enzyme induction and other mechanisms. Patients should be advised to consider alternative forms of contraception, or their contraceptive medications should be modified to offset the effects of the antiepileptic medications.

BREAST-FEEDING

Antiepileptic medications are excreted into breast milk to a variable degree. The ratio of drug concentration in breast milk relative to serum ranges from ~5% (valproic acid) to 300% (levetiracetam). Given the overall benefits of breast-feeding and the lack of evidence for long-term harm to the infant by being exposed to antiepileptic drugs, mothers with epilepsy can be encouraged to breast-feed. This should be reconsidered, however, if there is any evidence of drug effects on the infant such as lethargy or poor feeding.

446 Cerebrovascular Diseases

Wade S. Smith, S. Claiborne Johnston,
J. Claude Hemphill, III



Cerebrovascular diseases include some of the most common and devastating disorders: ischemic stroke and hemorrhagic stroke. Stroke is the second leading cause of death worldwide, causing 6.2 million deaths in 2011, and is double the rate of heart disease in China. Strokes cause ~200,000 deaths each year in the United States and are a major cause of disability. The incidence of cerebrovascular diseases increases with age, and the number of strokes is projected to increase as the elderly population grows, with a doubling in stroke deaths in the United States by 2030. A stroke, or cerebrovascular accident, is defined as an abrupt onset of a neurologic deficit that is attributable to a focal vascular cause. Thus, the definition of stroke is clinical, and laboratory studies including brain imaging are used to support the diagnosis. The clinical manifestations of stroke are highly variable because of the complex anatomy of the brain and its vasculature. *Cerebral ischemia* is caused by a reduction in blood flow that lasts longer than several seconds. Neurologic symptoms are manifest within seconds because neurons lack glycogen, so energy failure is rapid. If the cessation of flow lasts for more than a few minutes, *infarction* or death of brain tissue results. When blood flow is quickly restored, brain tissue can recover fully and the patient's symptoms are only transient: this is called a *transient ischemic attack* (TIA). The definition of TIA requires that all neurologic signs and symptoms resolve within 24 h without evidence of brain infarction on brain imaging. Stroke has occurred if the neurologic signs and symptoms last for >24 h or brain infarction is demonstrated. A generalized reduction in cerebral blood flow due to systemic hypotension (e.g., cardiac arrhythmia, myocardial infarction, or hemorrhagic shock) usually produces syncope (Chap. 27). If low cerebral blood flow persists for a longer duration, then infarction in the border zones between the major cerebral artery distributions may develop. In more severe instances, *global hypoxia-ischemia* causes widespread brain injury; the constellation of cognitive sequelae that ensues is called *hypoxic-ischemic encephalopathy* (Chap. 330). *Focal ischemia* or infarction, conversely, is usually caused by thrombosis of the cerebral vessels themselves or by emboli from a proximal arterial source or the heart. *Intracranial hemorrhage* is caused by bleeding directly into or around the brain; it produces neurologic symptoms by producing a

APPROACH TO THE PATIENT: Cerebrovascular Disease

Rapid evaluation is essential for use of time-sensitive treatments such as thrombolysis. However, patients with acute stroke often do not seek medical assistance on their own because they are rarely in pain and also may lose the appreciation that something is wrong (anosognosia); it is often a family member or a bystander who calls for help. Therefore, patients and their family members should be counseled to call emergency medical services immediately if they experience or witness the sudden onset of any of the following: loss of sensory and/or motor function on one side of the body (nearly 85% of ischemic stroke patients have hemiparesis); change in vision, gait, or ability to speak or understand; or a sudden, severe headache.

Other causes of sudden-onset neurologic symptoms that may mimic stroke include seizure, intracranial tumor, migraine, and metabolic encephalopathy. An adequate history from an observer that no convulsive activity occurred at the onset usually excludes seizure, although ongoing complex partial seizures without tonic-clonic activity can on occasion mimic stroke. Tumors may present with acute neurologic symptoms due to hemorrhage, seizure, or hydrocephalus. Surprisingly, migraine (Chap. 447) can mimic stroke, even in patients without a significant migraine history. When migraine develops without head pain (*acephalgic migraine*), the diagnosis can be especially difficult. Patients without any prior history of migraine may develop acephalgic migraine even after age 65. A sensory disturbance is often prominent, and the sensory deficit, as well as any motor deficits, tends to migrate slowly across a limb, over minutes rather than seconds as with stroke. The diagnosis of migraine becomes more secure as the cortical disturbance begins to cross vascular boundaries or if typical visual symptoms are present such as scintillating scotomata. At times it may be impossible to make the diagnosis of migraine until there have been multiple episodes with no residual symptoms or signs and no changes on brain magnetic resonance imaging (MRI). Metabolic encephalopathies typically produce fluctuating mental status changes without focal neurologic findings. However, in the setting of prior stroke or brain injury, a patient with fever or sepsis may manifest a recurrent hemiparesis, which clears rapidly when the infection is treated. The metabolic process serves to “unmask” a prior deficit.

Once the diagnosis of stroke is made, a brain imaging study is necessary to determine if the cause of stroke is ischemia or hemorrhage (Fig. 446-1). Computed tomography (CT) imaging of the brain is the standard imaging modality to detect the presence or absence of intracranial hemorrhage (see “Imaging Studies,” below). If the stroke is ischemic, administration of recombinant tissue plasminogen activator (rtPA) or endovascular mechanical thrombectomy may be beneficial in restoring cerebral perfusion (see “Treatment: Acute Ischemic Stroke”). Medical management to reduce the risk of complications becomes the next priority, followed by plans for secondary prevention. For ischemic stroke, several strategies can reduce the risk of subsequent stroke in all patients, while other strategies are effective for patients with specific causes of stroke such as cardiac embolus and carotid atherosclerosis. For hemorrhagic stroke, aneurysmal subarachnoid hemorrhage (SAH) and hypertensive intracerebral hemorrhage are two important causes. The treatment and prevention of hypertensive intracerebral hemorrhage are discussed later in this chapter. SAH is discussed in Chap. 330.

ISCHEMIC STROKE

PATHOPHYSIOLOGY OF ISCHEMIC STROKE

Acute occlusion of an intracranial vessel causes reduction in blood flow to the brain region it supplies. The magnitude of flow reduction is a function of collateral blood flow, and this depends on individual

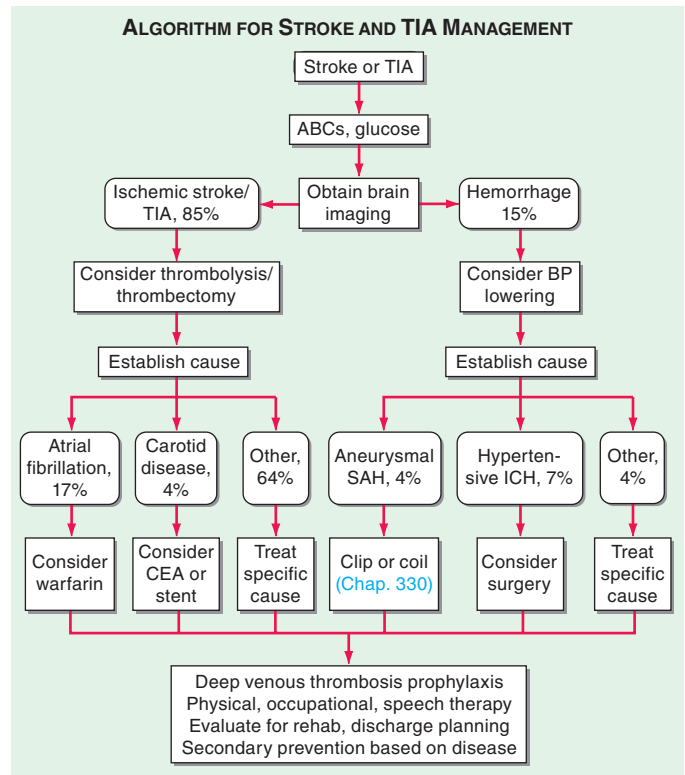


FIGURE 446-1 Medical management of stroke and TIA. Rounded boxes are diagnoses; rectangles are interventions. Numbers are percentages of stroke overall. ABCs, airway, breathing, circulation; BP, blood pressure; CEA, carotid endarterectomy; ICH, intracerebral hemorrhage; SAH, subarachnoid hemorrhage; TIA, transient ischemic attack.

vascular anatomy (which may be altered by disease), the site of occlusion, and systemic blood pressure. A decrease in cerebral blood flow to zero causes death of brain tissue within 4–10 min; values <16–18 mL/100 g tissue per minute cause infarction within an hour; and values <20 mL/100 g tissue per minute cause ischemia without infarction unless prolonged for several hours or days. If blood flow is restored to ischemic tissue before significant infarction develops, the patient may experience only transient symptoms, and the clinical syndrome is called a TIA. Another important concept is the *ischemic penumbra*, defined as the ischemic but reversibly dysfunctional tissue surrounding a core area of infarction. The penumbra can be imaged by perfusion-diffusion imaging using MRI or CT (see below and Figs. 446-15 and 446-16). The ischemic penumbra will eventually progress to infarction if no change in flow occurs, and hence saving the ischemic penumbra is the goal of revascularization therapies.

Focal cerebral infarction occurs via two distinct pathways (Fig. 446-2): (1) a necrotic pathway in which cellular cytoskeletal breakdown is rapid, due principally to energy failure of the cell; and (2) an apoptotic pathway in which cells become programmed to die. Ischemia produces necrosis by starving neurons of glucose and oxygen, which in turn results in failure of mitochondria to produce ATP. Without ATP, membrane ion pumps stop functioning and neurons depolarize, allowing intracellular calcium to rise. Cellular depolarization also causes glutamate release from synaptic terminals; excess extracellular glutamate produces neurotoxicity by activating postsynaptic glutamate receptors that increase neuronal calcium influx. Free radicals are produced by degradation of membrane lipids and mitochondrial dysfunction. Free radicals cause catalytic destruction of membranes and likely damage other vital functions of cells. Lesser degrees of ischemia, as are seen within the ischemic penumbra, favor apoptotic cellular death causing cells to die days to weeks later. Fever dramatically worsens brain injury during ischemia, as does hyperglycemia (glucose >11.1 mmol/L [200 mg/dL]), so it is reasonable to suppress fever and prevent

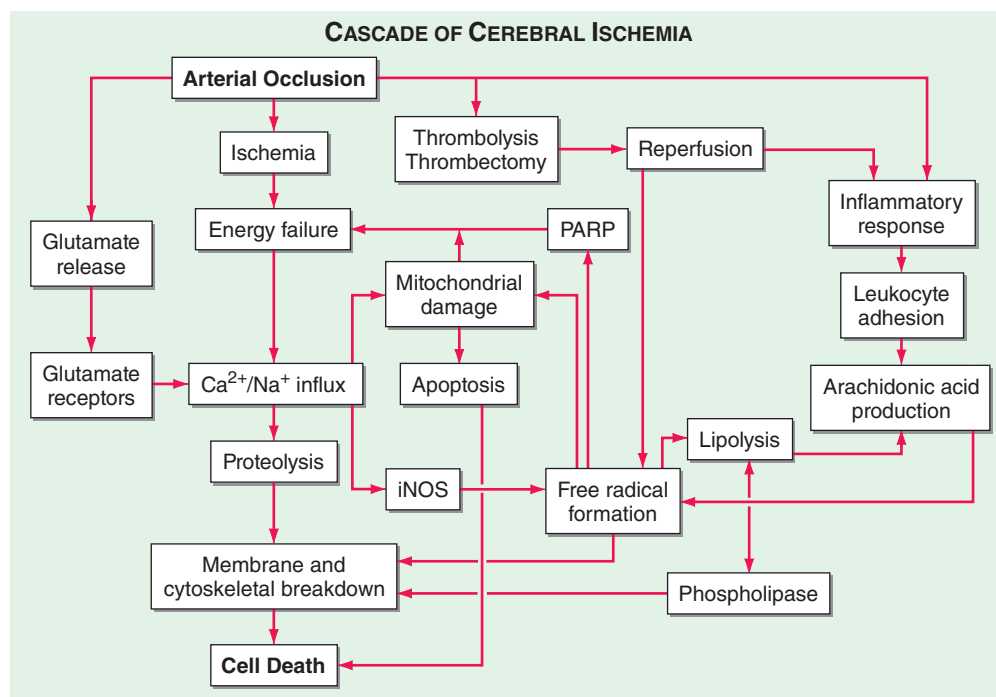


FIGURE 446-2 Major steps in the cascade of cerebral ischemia. See text for details. iNOS, inducible nitric oxide synthase; PARP, poly-A ribose polymerase.

hyperglycemia as much as possible. The value of induced mild hypothermia to improve stroke outcomes is the subject of continuing clinical research.

TREATMENT ACUTE ISCHEMIC STROKE

After the clinical diagnosis of stroke is made, an orderly process of evaluation and treatment should follow (Fig. 446-1). The first goal is to prevent or reverse brain injury. Attend to the patient's airway, breathing, and circulation (ABCs), and treat hypoglycemia or hyperglycemia if identified. Perform an emergency noncontrast head CT scan to differentiate between ischemic stroke and hemorrhagic stroke; there are no reliable clinical findings that conclusively separate ischemia from hemorrhage, although a more depressed level of consciousness, higher initial blood pressure, or worsening of symptoms after onset favor hemorrhage, and a deficit that is maximal at onset, or remits, suggests ischemia. Treatments designed to reverse or lessen the amount of tissue infarction and improve clinical outcome fall within six categories: (1) medical support, (2) IV thrombolysis, (3) endovascular revascularization, (4) antithrombotic treatment, (5) neuroprotection, and (6) stroke centers and rehabilitation.

MEDICAL SUPPORT

When ischemic stroke occurs, the immediate goal is to optimize cerebral perfusion in the surrounding ischemic penumbra. Attention is also directed toward preventing the common complications of bedridden patients—infections (pneumonia, urinary, and skin) and deep venous thrombosis (DVT) with pulmonary embolism. Subcutaneous heparin (unfractionated and low-molecular-weight) is safe and can be used concomitantly. Use of pneumatic compression stockings is of proven benefit in reducing risk of DVT and is a safe alternative to heparin.

Because collateral blood flow within the ischemic brain may be blood pressure dependent, there is controversy about whether blood pressure should be lowered acutely. Blood pressure should be lowered if there is malignant hypertension (Chap. 298) or concomitant myocardial ischemia, or if blood pressure is >185/110 mmHg and thrombolytic therapy is anticipated. When faced with the competing demands of myocardium and brain, lowering the heart rate with a β_1 -adrenergic blocker (such as esmolol) can be a first step to decrease

cardiac work and maintain blood pressure. Routine lowering of blood pressure has been found to worsen outcomes. Fever is detrimental and should be treated with antipyretics and surface cooling. Serum glucose should be monitored and kept at <10.0 mmol/L (180 mg/dL) using an insulin infusion if necessary.

Between 5 and 10% of patients develop enough cerebral edema to cause obtundation or brain herniation. Edema peaks on the second or third day but can cause mass effect for ~10 days. The larger the infarct, the greater the likelihood that clinically significant edema will develop. Water restriction and IV mannitol may be used to raise the serum osmolarity, but hypovolemia should be avoided because this may contribute to hypotension and worsening infarction. Combined analysis of three randomized European trials of hemicraniectomy (craniotomy and temporary removal of part of the skull) shows that hemicraniectomy markedly reduces mortality, and the clinical outcomes of survivors are acceptable. The size of the diffusion-weighted imaging volume of brain infarction during the acute stroke is a predictor of deterioration requiring hemicraniectomy.

Special vigilance is warranted for patients with cerebellar infarction. These strokes may mimic labyrinthitis because of prominent vertigo and vomiting; the presence of head or neck pain should alert the physician to consider cerebellar stroke from vertebral artery dissection. Even small amounts of cerebellar edema can acutely increase intracranial pressure (ICP) by obstructing cerebrospinal fluid (CSF) flow leading to hydrocephalus or by directly compressing the brainstem. The resulting brainstem compression can manifest as coma and respiratory arrest and require emergency surgical decompression. Prophylactic suboccipital decompression of large cerebellar infarcts before brainstem compression, although not tested rigorously in a clinical trial, is practiced at most stroke centers.

INTRAVENOUS THROMBOLYSIS

The National Institute of Neurological Disorders and Stroke (NINDS) rtPA Stroke Study showed a clear benefit for IV rtPA in selected patients with acute stroke. The NINDS study used IV rtPA (0.9 mg/kg to a 90-mg maximum; 10% as a bolus, then the remainder over 60 min) versus placebo in ischemic stroke within 3 h of onset. One-half of the patients were treated within 90 min. Symptomatic intracranial hemorrhage occurred in 6.4% of patients on rtPA and 0.6%

on placebo. In the rTPA group, there was a significant 12% absolute increase in the number of patients with only minimal disability (32% on placebo and 44% on rtPA) and a nonsignificant 4% reduction in mortality (21% on placebo and 17% on rtPA). Thus, despite an increased incidence of symptomatic intracranial hemorrhage, treatment with IV rtPA within 3 h of the onset of ischemic stroke improved clinical outcome.

Three subsequent trials of IV rtPA did not confirm this benefit, perhaps because of the dose of rtPA used, the timing of its delivery, and small sample size. When data from all randomized IV rtPA trials were combined, however, efficacy was confirmed in the <3-h time window, and efficacy likely extended to 4.5 h and possibly to 6 h. Based on these combined results, the European Cooperative Acute Stroke Study (ECASS) III explored the safety and efficacy of rtPA in the 3- to 4.5-h time window. Unlike the NINDS study, patients older than 80 years of age and diabetic patients with a previous stroke were excluded. In this 821-patient randomized study, efficacy was again confirmed, although the treatment effect was less robust than in the 0- to 3-h time window. In the rtPA group, 52.4% of patients achieved a good outcome at 90 days, compared to 45.2% of the placebo group (odds ratio [OR] 1.34, $p = .04$). The symptomatic intracranial hemorrhage rate was 2.4% in the rtPA group and 0.2% in the placebo group ($p = .008$).

Based on these data, rtPA is approved in the 3- to 4.5-h window in Europe and Canada, but is still only approved for 0–3 h in the United States and Canada. Use of IV tPA is now considered a central component of primary stroke centers (see below). It represents the first treatment proven to improve clinical outcomes in ischemic stroke and is cost-effective and cost-saving. Advanced neuroimaging techniques (see neuroimaging section below) may help to select patients beyond the 4.5-h window who will benefit from thrombolysis, but this is currently investigational. The time of stroke onset is defined as the time the patient's symptoms were witnessed to begin or the time the patient was last seen as normal. Patients who awaken with stroke have the onset defined as when they went to bed. **Table 446-1** summarizes eligibility criteria and instructions for administration of IV rtPA.

ENDOVASCULAR REVASULARIZATION

Ischemic stroke from large-vessel intracranial occlusion results in high rates of mortality and morbidity. Occlusions in such large vessels (middle cerebral artery [MCA], intracranial internal carotid artery, and the basilar artery) generally involve a large clot volume and often fail to open with IV rtPA alone. Therefore, there is growing interest in using thrombolytics via an intraarterial route to increase the concentration of drug at the clot and minimize systemic bleeding complications. The Prolyse in Acute Cerebral Thromboembolism (PROACT) II trial found benefit for intraarterial prourokinase in acute MCA occlusions up to the sixth hour following onset of stroke. Intraarterial treatment of basilar artery occlusions may also be beneficial for selected patients. Intraarterial administration of a thrombolytic agent for acute ischemic stroke (AIS) is not approved by the U.S. Food and Drug Administration (FDA); however, many stroke centers offer this treatment based on these data.

Endovascular mechanical thrombectomy has been studied as an alternative or adjunctive treatment of acute stroke in patients who are ineligible for, or have contraindications to, thrombolytics or in those who failed to achieve vascular recanalization with IV thrombolytics (see Fig. 446-15). The Mechanical Embolus Removal in Cerebral Ischemia (MERCi) and multi-MERCi single-arm trials found that an endovascular thrombectomy device restored patency of occluded intracranial vessels within 8 h of ischemic stroke symptoms compared with a historical control group. Recanalization of the target vessel occurred in 48–58% of treated patients and in 60–69% of patients after use of adjuvant endovascular methods, and successful recanalization at 90 days correlated well with favorable outcomes. Based on these nonrandomized data, the FDA approved this device as the first device for revascularization of occluded vessels in AIS even if the patient has been given rtPA and that therapy has failed.

TABLE 446-1 ADMINISTRATION OF INTRAVENOUS RECOMBINANT TISSUE PLASMINOGEN ACTIVATOR (rtPA) FOR ACUTE ISCHEMIC STROKE (AIS)^a

Indication	Contraindication
Clinical diagnosis of stroke	Sustained BP >185/110 mmHg despite treatment
Onset of symptoms to time of drug administration ≤ 4.5 h ^b	Platelets <100,000; HCT <25%; glucose <50 or >400 mg/dL
CT scan showing no hemorrhage or edema of >1/3 of the MCA territory	Use of heparin within 48 h and prolonged PTT, or elevated INR
Age 18 \geq years	Rapidly improving symptoms
Consent by patient or surrogate	Prior stroke or head injury within 3 months; prior intracranial hemorrhage
	Major surgery in preceding 14 days
	Minor stroke symptoms
	Gastrointestinal bleeding in preceding 21 days
	Recent myocardial infarction
	Coma or stupor
Administration of rtPA	
IV access with two peripheral IV lines (avoid arterial or central line placement)	
Review eligibility for rtPA	
Administer 0.9 mg/kg IV (maximum 90 mg) IV as 10% of total dose by bolus, followed by remainder of total dose over 1 h	
Frequent cuff blood pressure monitoring	
No other antithrombotic treatment for 24 h	
For decline in neurologic status or uncontrolled blood pressure, stop infusion, give cryoprecipitate, and reimagine brain emergently	
Avoid urethral catheterization for ≥ 2 h	

^aSee Activase (tissue plasminogen activator) package insert for complete list of contraindications and dosing. ^bDepending on the country, IV rtPA may be approved for up to 4.5 h with additional restrictions.

Abbreviations: BP, blood pressure; CT, computed tomography; HCT, hematocrit; INR, international normalized ratio; MCA, middle cerebral artery; PTT, partial thromboplastin time.

The Penumbra Pivotal Stroke trial tested another mechanical device that showed even higher rates of recanalization and led to FDA clearance of the tested device as well. More recently, two Stentriever devices (nondetachable stents) were shown to significantly improve vascular recanalization compared to the first approved MERCi device, approaching recanalization rates of 90% in most large intracranial vessels.

In 2013, three randomized endovascular trials with nonendovascular controls found no benefits to endovascular therapy. The largest was the Interventional Management of Stroke III trial that randomized 656 AIS patients within 3 h of onset to IV rtPA (0.9 mg/kg) alone versus IV rtPA (0.6 mg/kg) followed by endovascular adjuvant treatment with IA rtPA, or endovascular thrombectomy as soon as possible. Outcomes between these groups were not significantly different, and there were more complications (groin bleeding chiefly) in the endovascular group. The SYNTHESIS trial based in Italy randomized 363 patients to IV rtPA versus intraarterial rtPA for patients within 3 h of stroke onset. No differences were found between the groups at 90 days. These two relatively large trials indicate that endovascular therapy using principally intraarterial rtPA is not better than IV therapy, but many questions remain. Relatively few patients received mechanical clot retraction therapies, and those who did received what we now know were inferior devices. Trials assessing more efficacious thrombectomy devices are currently ongoing.

Because use of endovascular devices in combination with rtPA appears relatively safe, some centers continue to offer endovascular therapy. This applies to patients who are not eligible for IV rtPA (recent surgery, stroke following cardiac catheterization, etc.), and some continue to use thrombectomy because of perceived better

outcomes in patients with more effective devices. Comprehensive stroke centers are now obtaining credentialing to offer this therapy in distinction to primary stroke centers that offer only IV rtPA.

ANTITHROMBOTIC TREATMENT

Platelet Inhibition Aspirin is the only antiplatelet agent that has been proven effective for the acute treatment of ischemic stroke; there are several antiplatelet agents proven for the secondary prevention of stroke (see below). Two large trials, the International Stroke Trial (IST) and the Chinese Acute Stroke Trial (CAST), found that the use of aspirin within 48 h of stroke onset reduced both stroke recurrence risk and mortality minimally. Among 19,435 patients in IST, those allocated to aspirin, 300 mg/d, had slightly fewer deaths within 14 days (9.0 vs 9.4%), significantly fewer recurrent ischemic strokes (2.8 vs 3.9%), no excess of hemorrhagic strokes (0.9 vs 0.8%), and a trend toward a reduction in death or dependence at 6 months (61.2 vs 63.5%). In CAST, 21,106 patients with ischemic stroke received 160 mg/d of aspirin or a placebo for up to 4 weeks. There were very small reductions in the aspirin group in early mortality (3.3 vs 3.9%), recurrent ischemic strokes (1.6 vs 2.1%), and dependency at discharge or death (30.5 vs 31.6%). These trials demonstrate that the use of aspirin in the treatment of AIS is safe and produces a small net benefit. For every 1000 acute strokes treated with aspirin, about 9 deaths or nonfatal stroke recurrences will be prevented in the first few weeks and ~13 fewer patients will be dead or dependent at 6 months.

Clopidogrel is being tested as a way to prevent stroke following TIA and minor ischemic stroke (see below).

Anticoagulation Numerous clinical trials have failed to demonstrate any benefit of anticoagulation in the primary treatment of atherothrombotic cerebral ischemia. Several trials have investigated antiplatelet versus anticoagulant medications given within 12–24 h of the initial event. The U.S. Trial of Organon 10172 in Acute Stroke Treatment (TOAST), an investigational low-molecular-weight heparin (LMWH), failed to show any benefit over aspirin. Use of SC unfractionated heparin versus aspirin was tested in IST. Heparin given SC afforded no additional benefit over aspirin and increased bleeding rates. Several trials of LMWHs have also shown no consistent benefit in AIS. Furthermore, trials generally have shown an excess risk of brain and systemic hemorrhage with acute anticoagulation. A recent meta-analysis of all forms of heparin found no benefit for acute stroke patients at high or low risk of thrombotic events. Therefore, trials do not support the use of heparin or other anticoagulants for patients with atherothrombotic stroke.

NEUROPROTECTION

Neuroprotection is the concept of providing a treatment that prolongs the brain's tolerance to ischemia. Drugs that block the excitatory amino acid pathways have been shown to protect neurons and glia in animals, but despite multiple human trials, they have not yet been proven to be beneficial. Hypothermia is a powerful neuroprotective treatment in patients with cardiac arrest ([Chap. 330](#)) and is neuroprotective in animal models of stroke, but it has not been adequately studied in patients with ischemic stroke and is associated with an increase in pneumonia rates that could adversely impact stroke outcomes.

STROKE CENTERS AND REHABILITATION

Patient care in stroke units followed by rehabilitation services improves neurologic outcomes and reduces mortality. Use of clinical pathways and staff dedicated to the stroke patient can improve care. This includes use of standardized stroke order sets. Stroke teams that provide emergency 24-h evaluation of acute stroke patients for acute medical management and consideration of thrombolysis or endovascular treatments are essential components of primary and comprehensive stroke centers, respectively.

Proper rehabilitation of the stroke patient includes early physical, occupational, and speech therapy. It is directed toward educating the patient and family about the patient's neurologic deficit,

preventing the complications of immobility (e.g., pneumonia, DVT and pulmonary embolism, pressure sores of the skin, and muscle contractures), and providing encouragement and instruction in overcoming the deficit. Use of pneumatic compression stockings is of proven benefit in reducing risk of DVT and is a safe alternative to heparin. The goal of rehabilitation is to return the patient to home and to maximize recovery by providing a safe, progressive regimen suited to the individual patient. Additionally, the use of constrained movement therapy (immobilizing the unaffected side) has been shown to improve hemiparesis following stroke, even years after the stroke, suggesting that physical therapy can recruit unused neural pathways. Newer robotic therapies appear promising as well. The human nervous system is more adaptable than previously thought, and developing physical and pharmacologic strategies to enhance long-term neural recovery is an active area of research.

ETIOLOGY OF ISCHEMIC STROKE

([Figs. 446-1](#) and [446-3](#) and [Table 446-2](#)) Although the initial management of AIS often does not depend on the etiology, establishing a cause is essential to reduce the risk of recurrence. Particular focus should be on atrial fibrillation and carotid atherosclerosis, because these etiologies have proven secondary prevention strategies. The clinical presentation and examination findings often establish the cause of stroke or narrow the possibilities to a few. Judicious use of laboratory testing and imaging studies completes the initial evaluation. Nevertheless, nearly 30% of strokes remain unexplained despite extensive evaluation.

Clinical examination should focus on the peripheral and cervical vascular system (carotid auscultation for bruits and blood pressure), the heart (dysrhythmia, murmurs), extremities (peripheral emboli), and retina (effects of hypertension and cholesterol emboli [Hollenhorst plaques]). A complete neurologic examination is performed to localize the anatomic site of stroke. An imaging study of the brain is nearly always indicated and is required for patients being considered for thrombolysis; it may be combined with CT- or MRI-based angiography to visualize the vasculature of the neck and intracranial vessels (see "Imaging Studies," below). A chest x-ray, electrocardiogram (ECG), urinalysis, complete blood count, erythrocyte sedimentation rate (ESR), serum electrolytes, blood urea nitrogen (BUN), creatinine, blood glucose, serum lipid profile, prothrombin time (PT), and partial thromboplastin time (PTT) are often useful and should be considered in all patients. An ECG may demonstrate arrhythmias or reveal evidence of recent myocardial infarction (MI). Of all these studies, only brain imaging, blood glucose, and perhaps PTT/international normalized ratio (INR) are necessary prior to IV rtPA; the results of other studies should not delay the rapid administration of IV rtPA if the patient is eligible.

Cardioembolic Stroke Cardioembolism is responsible for ~20% of all ischemic strokes. Stroke caused by heart disease is primarily due to embolism of thrombotic material forming on the atrial or ventricular wall or the left heart valves. These thrombi then detach and embolize into the arterial circulation. The thrombus may fragment or lyse quickly, producing only a TIA. Alternatively, the arterial occlusion may last longer, producing stroke. Embolic strokes tend to occur suddenly with maximum neurologic deficit present at onset. With reperfusion following more prolonged ischemia, petechial hemorrhages can occur within the ischemic territory. These are usually of no clinical significance and should be distinguished from frank intracranial hemorrhage into a region of ischemic stroke where the mass effect from the hemorrhage can cause a significant decline in neurologic function.

Emboli from the heart most often lodge in the intracranial internal carotid artery, the MCA, the posterior cerebral artery (PCA), or one of their branches; infrequently, the anterior cerebral artery (ACA) is involved. Emboli large enough to occlude the stem of the MCA (3–4 mm) lead to large infarcts that involve both deep gray and white matter and some portions of the cortical surface and its underlying white matter. A smaller embolus may occlude a small cortical or penetrating arterial branch. The location and size of an infarct within a vascular territory depend on the extent of the collateral circulation.

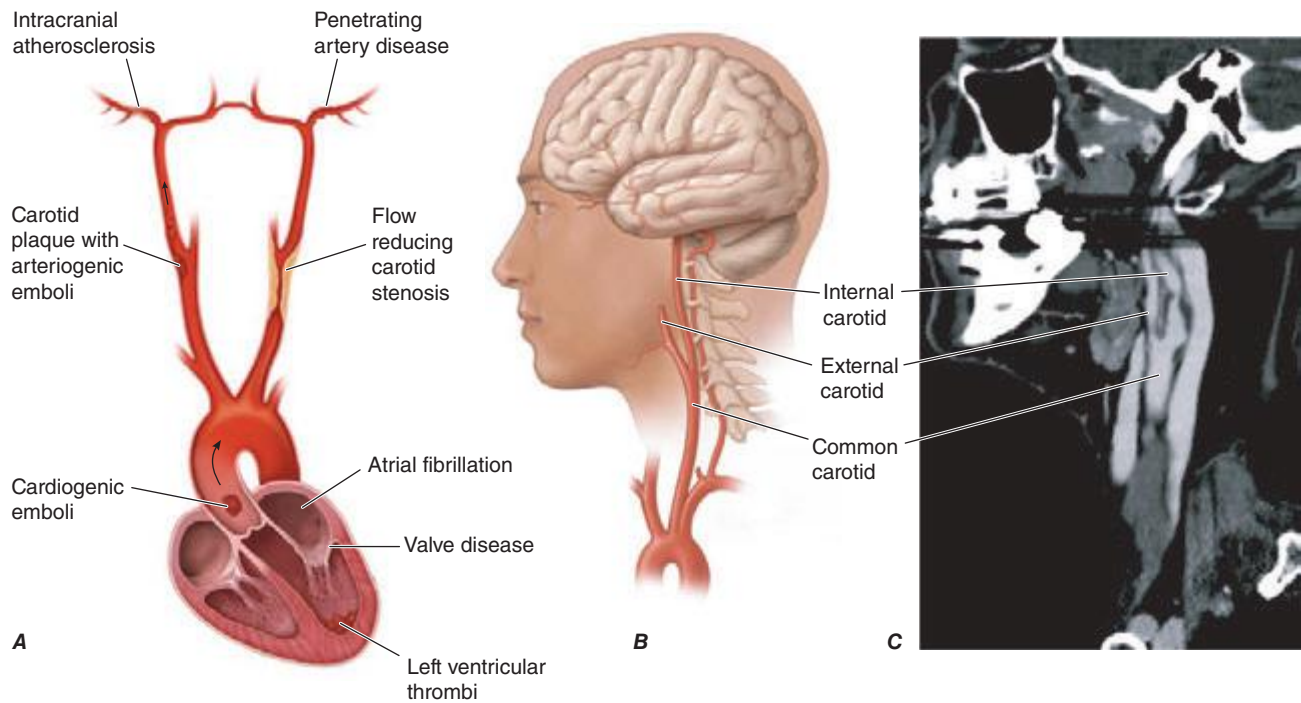


FIGURE 446-3 Pathophysiology of ischemic stroke. **A.** Diagram illustrating the three major mechanisms that underlie ischemic stroke: (1) occlusion of an intracranial vessel by an embolus that arises at a distant site (e.g., cardiogenic sources such as atrial fibrillation or artery-to-artery emboli from carotid atherosclerotic plaque), often affecting the large intracranial vessels; (2) in situ thrombosis of an intracranial vessel, typically affecting the small penetrating arteries that arise from the major intracranial arteries; (3) hypoperfusion caused by flow-limiting stenosis of a major extracranial (e.g., internal carotid) or intracranial vessel, often producing “watershed” ischemia. **B.** and **C.** Diagram and reformatted computed tomography angiogram of the common, internal, and external carotid arteries. High-grade stenosis of the internal carotid artery, which may be associated with either cerebral emboli or flow-limiting ischemia, was identified in this patient.

The most significant causes of cardioembolic stroke in most of the world are nonrheumatic (often called nonvalvular) atrial fibrillation, MI, prosthetic valves, rheumatic heart disease, and ischemic cardiomyopathy (Table 446-2). Cardiac disorders causing brain embolism are discussed in the chapters on heart diseases. A few pertinent aspects are highlighted here.

Nonrheumatic atrial fibrillation is the most common cause of cerebral embolism overall. The presumed stroke mechanism is thrombus formation in the fibrillating atrium or atrial appendage, with subsequent embolization. Patients with atrial fibrillation have an average annual risk of stroke of ~5%. The risk of stroke can be estimated by calculating the CHADS2 score (Table 446-3). Left atrial enlargement is an additional risk factor for formation of atrial thrombi. Rheumatic heart disease usually causes ischemic stroke when there is prominent mitral stenosis or atrial fibrillation. Recent MI may be a source of emboli, especially when transmural and involving the anteroapical ventricular wall, and prophylactic anticoagulation following MI has been shown to reduce stroke risk. Mitral valve prolapse is not usually a source of emboli unless the prolapse is severe.

Paradoxical embolization occurs when venous thrombi migrate to the arterial circulation, usually via a patent foramen ovale or atrial septal defect. Bubble-contrast echocardiography (IV injection of agitated saline coupled with either transthoracic or transesophageal echocardiography) can demonstrate a right-to-left cardiac shunt, revealing the conduit for paradoxical embolization. Alternatively, a right-to-left shunt is implied if immediately following IV injection of agitated saline, the ultrasound signature of bubbles is observed during transcranial Doppler insonation of the MCA; pulmonary arteriovenous malformations should be considered if this test is positive yet an echocardiogram fails to reveal an intracardiac shunt. Both techniques are highly sensitive for detection of right-to-left shunts. Besides venous clot, fat and tumor emboli, bacterial endocarditis, IV air, and amniotic fluid emboli at childbirth may occasionally be responsible for paradoxical embolization. The importance of a patent foramen ovale (PFO) as a

cause of stroke is debated, particularly because they are present in ~15% of the general population. Some studies have suggested that the risk is only elevated in the presence of a coexisting atrial septal aneurysm. The presence of a venous source of embolus, most commonly a deep venous thrombus, may provide confirmation of the importance of a PFO with an accompanying right-to-left shunt in a particular case. Three randomized trials of PFO occlusion for secondary prevention of ischemic stroke were negative, although each lacked sufficient power to be conclusive. At present, there is no supportive evidence to offer percutaneous PFO closure for stroke prevention.

Bacterial endocarditis can be a source of valvular vegetations that give rise to septic emboli. The appearance of multifocal symptoms and signs in a patient with stroke makes bacterial endocarditis more likely. Infarcts of microscopic size occur, and large septic infarcts may evolve into brain abscesses or cause hemorrhage into the infarct, which generally precludes use of anticoagulation or thrombolytics. Mycotic aneurysms caused by septic emboli may also present as SAH or intracerebral hemorrhage.

Artery-to-Artery Embolic Stroke Thrombus formation on atherosclerotic plaques may embolize to intracranial arteries producing an artery-to-artery embolic stroke. Less commonly, a diseased vessel may acutely thrombose. Unlike the myocardial vessels, artery-to-artery embolism, rather than local thrombosis, appears to be the dominant vascular mechanism causing large-vessel brain ischemia. Any diseased vessel may be an embolic source, including the aortic arch, common carotid, internal carotid, vertebral, and basilar arteries.

CAROTID ATHEROSCLEROSIS Atherosclerosis within the carotid artery occurs most frequently within the common carotid bifurcation and proximal internal carotid artery; the carotid siphon (portion within the cavernous sinus) is also vulnerable to atherosclerosis. Male gender, older age, smoking, hypertension, diabetes, and hypercholesterolemia are risk factors for carotid disease, as they are for stroke in general (Table 446-4). Carotid atherosclerosis produces an estimated 10% of

TABLE 446-2 CAUSES OF ISCHEMIC STROKE

Common Causes	Uncommon Causes
Thrombosis	Hypercoagulable disorders
Lacunar stroke (small vessel)	Protein C deficiency ^a
Large-vessel thrombosis	Protein S deficiency ^a
Dehydration	Antithrombin III deficiency ^a
Emboloc occlusion	Antiphospholipid syndrome
Artery-to-artery	Factor V Leiden mutation ^a
Carotid bifurcation	Prothrombin G20210 mutation ^a
Aortic arch	Systemic malignancy
Arterial dissection	Sickle cell anemia
Cardioembolic	β Thalassemia
Atrial fibrillation	Polycythemia vera
Mural thrombus	Systemic lupus erythematosus
Myocardial infarction	Homocysteinemia
Dilated cardiomyopathy	Thrombotic thrombocytopenic purpura
Valvular lesions	Disseminated intravascular coagulation
Mitral stenosis	Dysproteinemias ^a
Mechanical valve	Nephrotic syndrome ^a
Bacterial endocarditis	Inflammatory bowel disease ^a
Paradoxical embolus	Oral contraceptives
Atrial septal defect	Venous sinus thrombosis ^b
Patent foramen ovale	Fibromuscular dysplasia
Atrial septal aneurysm	Vasculitis
Spontaneous echo contrast	Systemic vasculitis (PAN, granulomatosis with polyangiitis [Wegener's], Takayasu's, giant cell arteritis)
Stimulant drugs: cocaine, amphetamine	Primary CNS vasculitis
	Meningitis (syphilis, tuberculosis, fungal, bacterial, zoster)
	Noninflammatory vasculopathy
	Reversible vasoconstriction syndrome
	Fabry's disease
	Angiocentric lymphoma
	Cardiogenic
	Mitral valve calcification
	Atrial myxoma
	Intracardiac tumor
	Marantic endocarditis
	Libman-Sacks endocarditis
	Subarachnoid hemorrhage vasospasm
	Moyamoya disease
	Eclampsia

^aChiefly cause venous sinus thrombosis. ^bMay be associated with any hypercoagulable disorder.

Abbreviations: CNS, central nervous system; PAN, polyarteritis nodosa.

ischemic stroke. **For further discussion of the pathogenesis of atherosclerosis, see Chap. 291e.**

Carotid disease can be classified by whether the stenosis is symptomatic or asymptomatic and by the degree of stenosis (percent narrowing of the narrowest segment compared to a nondiseased segment). Symptomatic carotid disease implies that the patient has experienced a stroke or TIA within the vascular distribution of the artery, and it is associated with a greater risk of subsequent stroke than asymptomatic stenosis, in which the patient is symptom free and the stenosis is detected through screening. Greater degrees of arterial narrowing are generally associated with a higher risk of stroke, except that those with near occlusions are at lower risk of stroke.

OTHER CAUSES OF ARTERY-TO-ARTERY EMBOLIC STROKE Intracranial atherosclerosis produces stroke either by an embolic mechanism or by in situ

thrombosis of a diseased vessel. It is more common in patients of Asian and African-American descent. Recurrent stroke risk is ~15% per year, similar to symptomatic untreated carotid atherosclerosis.

Dissection of the internal carotid or vertebral arteries or even vessels beyond the circle of Willis is a common source of embolic stroke in young (age <60 years) patients. The dissection is usually painful and precedes the stroke by several hours or days. Extracranial dissections do not cause hemorrhage, presumably because of the tough adventitia of these vessels. Intracranial dissections, conversely, may produce SAH because the adventitia of intracranial vessels is thin and pseudoaneurysms may form, requiring urgent treatment to prevent rerupture. Treating asymptomatic pseudoaneurysms following dissection is likely not necessary. The cause of dissection is usually unknown, and recurrence is rare. Ehlers-Danlos type IV, Marfan's disease, cystic medial necrosis, and fibromuscular dysplasia are associated with dissections. Trauma (usually a motor vehicle accident or a sports injury) can cause carotid and vertebral artery dissections. Spinal manipulative therapy is associated with vertebral artery dissection and stroke. Most dissections heal spontaneously, and stroke or TIA is uncommon beyond 2 weeks. Although there are no trials comparing anticoagulation to antiplatelet agents, many physicians treat acutely with anticoagulants and then convert to antiplatelet therapy after demonstration of satisfactory vascular recanalization.

SMALL-VESSEL STROKE

The term *lacunar infarction* refers to infarction following atherothrombotic or lipohyalinotic occlusion of a small artery in the brain. The term *small-vessel stroke* denotes occlusion of such a small penetrating artery and is now the preferred term. Small-vessel strokes account for ~20% of all strokes.

Pathophysiology The MCA stem, the arteries comprising the circle of Willis (A1 segment, anterior and posterior communicating arteries, and P1 segment), and the basilar and vertebral arteries all give rise to 30- to 300-μm branches that penetrate the deep gray and white matter of the cerebrum or brainstem (Fig. 446-4). Each of these small branches can occlude either by atherothrombotic disease at its origin or by the development of lipohyalinotic thickening. Thrombosis of these vessels causes small infarcts that are referred to as *lacunes* (Latin for "lake" of fluid noted at autopsy). These infarcts range in size from 3 mm to 2 cm in diameter. Hypertension and age are the principal risk factors.

Clinical Manifestations The most common small-vessel stroke syndromes are the following: (1) *Pure motor hemiparesis* from an infarct in the posterior limb of the internal capsule or the pons; the face, arm, and leg are almost always involved; (2) *pure sensory stroke* from an infarct in the ventral thalamus; (3) *ataxic hemiparesis* from an infarct in the ventral pons or internal capsule; (4) and *dysarthria and a clumsy hand* or arm due to infarction in the ventral pons or in the genu of the internal capsule.

Transient symptoms (small-vessel TIAs) may herald a small-vessel infarct; they may occur several times a day and last only a few minutes. Recovery from small-vessel strokes tends to be more rapid and complete than recovery from large-vessel strokes; in some cases, however, there is severe permanent disability.

A large-vessel source (either thrombosis or embolism) may manifest initially as a small-vessel infarction. Therefore, the search for embolic sources (carotid and heart) should not be completely abandoned in the evaluation of these patients. Secondary prevention of small-vessel stroke involves risk factor modification, specifically reduction in blood pressure (see "Treatment: Primary and Secondary Prevention of Stroke and TIA," below).

LESS COMMON CAUSES OF STROKE

(Table 446-2) **Hypercoagulable disorders** (Chap. 78) primarily increase the risk of venous, including venous sinus, thrombosis. Systemic lupus erythematosus with Libman-Sacks endocarditis can be a cause of embolic stroke. These conditions overlap with the antiphospholipid syndrome, which probably requires long-term anticoagulation to

TABLE 446-3 RECOMMENDATIONS ON CHRONIC USE OF ANTITHROMBOTICS FOR VARIOUS CARDIAC CONDITIONS

Condition	Recommendation
Nonvalvular atrial fibrillation	Calculate CHADS2 ^a score
• CHADS2 score 0	Aspirin or no antithrombotic
• CHADS2 score 1	Aspirin or OAC
• CHADS2 score >1	OAC
Rheumatic mitral valve disease	
• With atrial fibrillation, previous embolization, or atrial appendage thrombus, or left atrial diameter >55 mm	OAC
• Embolization or appendage clot despite OAC	OAC plus aspirin
Mitral valve prolapse	
• Asymptomatic	No therapy
• With otherwise cryptogenic stroke or TIA	Aspirin
• Atrial fibrillation	OAC
Mitral annular calcification	
• Without atrial fibrillation but systemic embolization, or otherwise cryptogenic stroke or TIA	Aspirin
• Recurrent embolization despite aspirin	OAC
• With atrial fibrillation	OAC
Aortic valve calcification	
• Asymptomatic	No therapy
• Otherwise cryptogenic stroke or TIA	Aspirin
Aortic arch mobile atheroma	
• Otherwise cryptogenic stroke or TIA	Aspirin or OAC
Patent foramen ovale	
• Otherwise cryptogenic ischemic stroke or TIA	Aspirin
• Indication for OAC (deep venous thrombosis or hypercoagulable state)	OAC
Mechanical heart valve	
• Aortic position, bileaflet or Medtronic Hall tilting disk with normal left atrial size and sinus rhythm	VKA INR 2.5, range 2–3
• Mitral position tilting disk or bileaflet valve	VKA INR 3.0, range 2.5–3.5
• Mitral or aortic position, anterior-apical myocardial infarct or left atrial enlargement	VKA INR 3.0, range 2.5–3.5
• Mitral or aortic position, with atrial fibrillation, or hypercoagulable state, or low ejection fraction, or atherosclerotic vascular disease	Aspirin plus VKA INR 3.0, range 2.5–3.5
• Systemic embolization despite target INR	Add aspirin and/or increase INR: prior target was 2.5 increase to 3.0, range 2.5–3.5; prior target was 3.0 increase to 3.5, range 3–4
Bioprosthetic valve	
• No other indication for VKA therapy	Aspirin
Infective endocarditis	Avoid antithrombotic agents
Nonbacterial thrombotic endocarditis	
• With systemic embolization	Full-dose unfractionated heparin or SC LMWH

^aCHADS2 score calculated as follows: 1 point for age >75 years, 1 point for hypertension, 1 point for congestive heart failure, 1 point for diabetes, and 2 points for stroke or TIA; sum of points is the total CHADS2 score.

Note: Dose of aspirin is 50–325 mg/d; target INR for OAC is between 2 and 3 unless otherwise specified.

Abbreviations: INR, international normalized ratio; LMWH, low-molecular-weight heparin; OAC, oral anticoagulant (VKA, thrombin inhibitor, oral factor Xa inhibitors); TIA, transient ischemic attack; VKA, vitamin K antagonist.

Sources: Modified from DE Singer et al: *Chest* 133:546S, 2008; DN Salem et al: *Chest* 133:593S, 2008.

prevent further stroke. Homocysteinemia may cause arterial thromboses as well; this disorder is caused by various mutations in the homocysteine pathways and responds to different forms of cobalamin depending on the mutation.

Venous sinus thrombosis of the lateral or sagittal sinus or of small cortical veins (cortical vein thrombosis) occurs as a complication of oral contraceptive use, pregnancy and the postpartum period, inflammatory bowel disease, intracranial infections (meningitis), and dehydration. It is also seen in patients with laboratory-confirmed thrombophilia including polycythemia, sickle cell anemia, deficiencies

of proteins C and S, factor V Leiden mutation (resistance to activated protein C), antithrombin III deficiency, homocysteinemia, and the prothrombin G20210 mutation. Women who take oral contraceptives and have the prothrombin G20210 mutation may be at particularly high risk for sinus thrombosis. Patients present with headache and may also have focal neurologic signs (especially paraparesis) and seizures. Often, CT imaging is normal unless an intracranial venous hemorrhage has occurred, but the venous sinus occlusion is readily visualized using magnetic resonance (MR) or CT venography or conventional x-ray angiography. With greater degrees of sinus thrombosis, the

TABLE 446-4 RISK FACTORS FOR STROKE

Risk Factor	Relative Risk	Relative Risk Reduction with Treatment	Number Needed to Treat ^a	
			Primary Prevention	Secondary Prevention
Hypertension	2–5	38%	100–300	50–100
Atrial fibrillation	1.8–2.9	68% warfarin, 21% aspirin	20–83	13
Diabetes	1.8–6	No proven effect		
Smoking	1.8	50% at 1 year, baseline risk at 5 years postcessation		
Hyperlipidemia	1.8–2.6	16–30%	560	230
Asymptomatic carotid stenosis	2.0	53%	85	N/A
Symptomatic carotid stenosis (70–99%)		65% at 2 years	N/A	12
Symptomatic carotid stenosis (50–69%)		29% at 5 years	N/A	77

^aNumber needed to treat to prevent one stroke annually. Prevention of other cardiovascular outcomes is not considered here.

Abbreviation: N/A, not applicable.

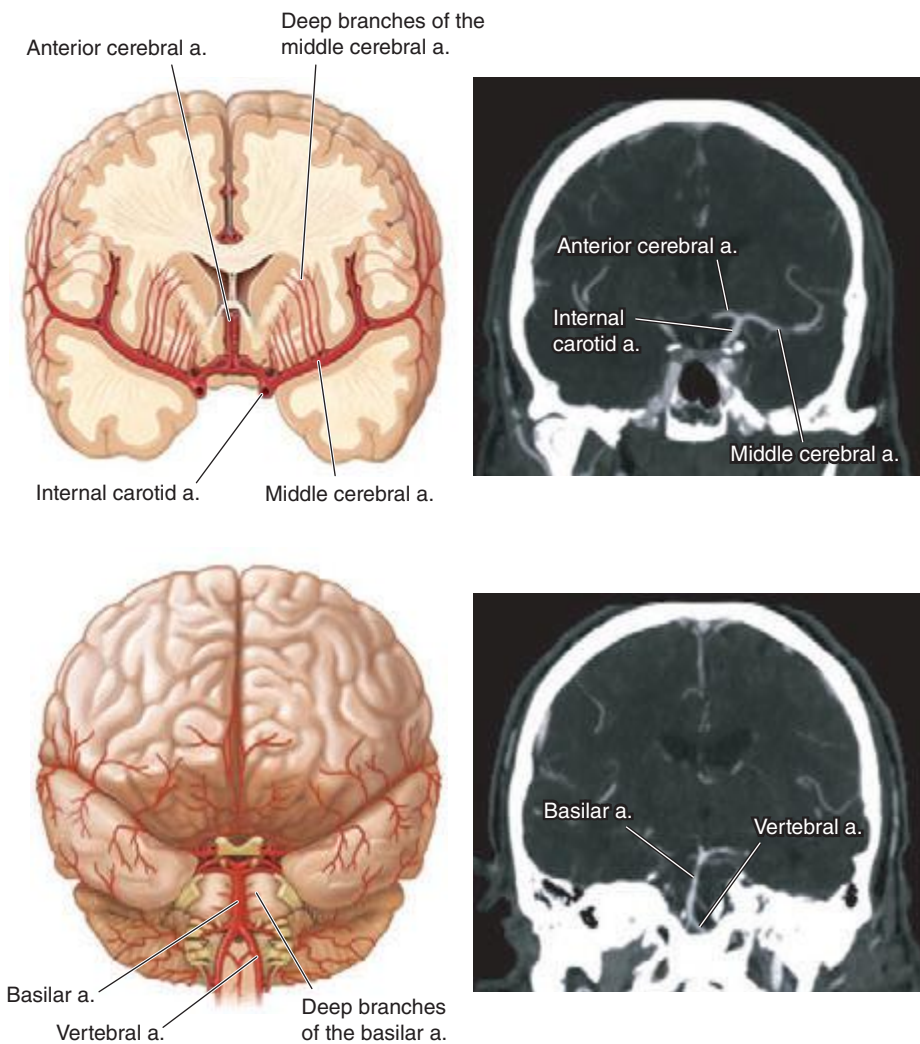


FIGURE 446-4 Diagrams and reformatted computed tomography (CT) angiograms in the coronal section illustrating the deep penetrating arteries involved in small-vessel strokes. In the anterior circulation, small penetrating arteries called *lenticulostriates* arise from the proximal portion of the anterior and middle cerebral arteries and supply deep subcortical structures (**upper panels**). In the posterior circulation, similar arteries arise directly from the vertebral and basilar arteries to supply the brainstem (**lower panels**). Occlusion of a single penetrating artery gives rise to a discrete area of infarct (pathologically termed a “lacune,” or lake). Note that these vessels are too small to be visualized on CT angiography.

patient may develop signs of increased ICP and coma. Intravenous heparin, regardless of the presence of intracranial hemorrhage, reduces morbidity and mortality, and the long-term outcome is generally good. Heparin prevents further thrombosis and reduces venous hypertension and ischemia. If an underlying hypercoagulable state is not found, many physicians treat with vitamin K antagonists (VKAs) for 3–6 months and then convert to aspirin, depending on the degree of resolution of the venous sinus thrombus. Anticoagulation is often continued indefinitely if thrombophilia is diagnosed.

Sickle cell anemia (SS disease) is a common cause of stroke in children. A subset of homozygous carriers of this hemoglobin mutation develop stroke in childhood, and this may be predicted by documenting high-velocity blood flow within the MCAs using transcranial Doppler ultrasonography. In children who are identified to have high velocities, treatment with aggressive exchange transfusion dramatically reduces risk of stroke, and if exchange transfusion is ceased, their stroke rate increases again along with MCA velocities.

Fibromuscular dysplasia affects the cervical arteries and occurs mainly in women. The carotid or vertebral arteries show multiple rings of segmental narrowing alternating with dilatation. Vascular occlusion is usually incomplete. The process is often asymptomatic but occasionally is associated with an audible bruit, TIAs, or stroke. Involvement of the renal arteries is common and may cause hypertension. The cause and natural history of fibromuscular dysplasia are unknown (**Chap. 302**). TIA or stroke generally occurs only when the artery is severely narrowed or dissects. Anticoagulation or antiplatelet therapy may be helpful.

Temporal (giant cell) arteritis (**Chap. 385**) is a relatively common affliction of elderly individuals in which the external carotid system, particularly the temporal arteries, undergo subacute granulomatous inflammation with giant cells. Occlusion of posterior ciliary arteries derived from the ophthalmic artery results in blindness in one or both eyes and can be prevented with glucocorticoids. It rarely causes stroke because the internal carotid artery is usually not inflamed. Idiopathic giant cell arteritis involving the great vessels arising from the aortic arch (*Takayasu's arteritis*) may cause carotid or vertebral thrombosis; it is rare in the Western Hemisphere.

Necrotizing (or granulomatous) arteritis, occurring alone or in association with generalized polyarteritis nodosa or granulomatosis with polyangiitis (Wegener's), involves the distal small

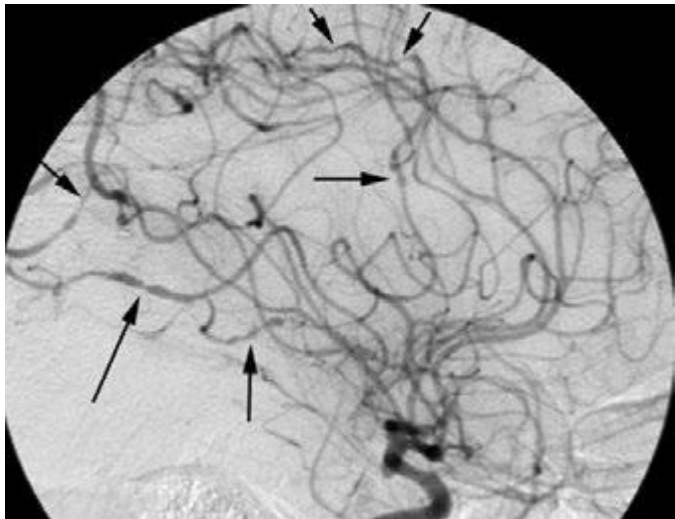


FIGURE 446-5 Cerebral angiogram from a 32-year-old male with central nervous system vasculopathy. Dramatic beading (arrows) typical of vasculopathy is seen.

branches (<2 mm diameter) of the main intracranial arteries and produces small ischemic infarcts in the brain, optic nerve, and spinal cord. The CSF often shows pleocytosis, and the protein level is elevated. *Primary central nervous system vasculitis* is rare; small or medium-sized vessels are usually affected, without apparent systemic vasculitis. The differential diagnosis includes other inflammatory vasculopathies including infection (tuberculous, fungal), sarcoidosis, angiocentric lymphoma, carcinomatous meningitis, and noninflammatory causes such as atherosclerosis, emboli, connective tissue disease, vasospasm, migraine-associated vasculopathy, and drug-associated causes. Some cases develop in the postpartum period and are self-limited.

Patients with any form of vasculopathy may present with insidious progression of combined white and gray matter infarctions, prominent headache, and cognitive decline. Brain biopsy or high-resolution conventional x-ray angiography is usually required to make the diagnosis (Fig. 446-5). An inflammatory profile found on lumbar puncture favors an inflammatory cause. In cases where inflammation is confirmed, aggressive immunosuppression with glucocorticoids, and often cyclophosphamide, is usually necessary to prevent progression; a diligent investigation for infectious causes such as tuberculosis is essential prior to immunosuppression. With prompt recognition and treatment, many patients can make an excellent recovery.

Drugs, in particular amphetamines and perhaps cocaine, may cause stroke on the basis of acute hypertension or drug-induced vasculopathy. No data exist on the value of any treatment. Phenylpropranolamine has been linked with intracranial hemorrhage, as has cocaine and methamphetamine, perhaps related to a drug-induced vasculopathy. *Moyamoya disease* is a poorly understood occlusive disease involving large intracranial arteries, especially the distal internal carotid artery and the stem of the MCA and ACA. Vascular inflammation is absent. The lenticulostriate arteries develop a rich collateral circulation around the occlusive lesion, which gives the impression of a “puff of smoke” (*moyamoya* in Japanese) on conventional x-ray angiography. Other collaterals include transdural anastomoses between the cortical surface branches of the meningeal and scalp arteries. The disease occurs mainly in Asian children or young adults, but the appearance may be identical in adults who have atherosclerosis, particularly in association with diabetes. Intracranial hemorrhage may result from rupture of the transdural and pial anastomotic channels; thus, anticoagulation is risky. Breakdown of dilated lenticulostriate arteries may produce intraparenchymal hemorrhage, and progressive occlusion of large surface arteries can occur, producing large-artery distribution strokes. Surgical bypass of extracranial carotid arteries to the dura or MCAs may prevent stroke and hemorrhage.

Posterior reversible encephalopathy syndrome (PRES) can occur with head injury, seizure, migraine, sympathomimetic drug use, eclampsia, and in the postpartum period (Chap. 463e). The pathophysiology is uncertain but likely involves a hyperperfusion state with widespread segmental vasoconstriction and cerebral edema. Patients complain of headache and manifest fluctuating neurologic symptoms and signs, especially visual symptoms. Sometimes cerebral infarction ensues, but typically the clinical and imaging findings suggest that ischemia reverses completely. MRI findings are characteristic with the edema present within the occipital lobes but can be generalized and do not respect any single vascular territory. A closely related *reversible cerebral vasoconstriction syndrome* (RCVS) typically presents with sudden, severe headache closely mimicking SAH. Patients may experience ischemic infarction and intracerebral hemorrhage and typically have new-onset, severe hypertension. Conventional x-ray angiography reveals changes in the vascular caliber throughout the hemispheres resembling vasculitis, but the process is noninflammatory. Oral calcium channel blockers may be effective in producing remission, and recurrence is rare.

Leukoaraiosis, or *periventricular white matter disease*, is the result of multiple small-vessel infarcts within the subcortical white matter. It is readily seen on CT or MRI scans as areas of white matter injury surrounding the ventricles and within the corona radiata. The pathophysiologic basis of the disease is lipohyalinosis of small penetrating arteries within the white matter, likely produced by chronic hypertension. Patients with periventricular white matter disease may develop a subcortical dementia syndrome, and it is likely that this common form of dementia may be delayed or prevented with antihypertensive medications (Chap. 448).

CADASIL (cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy) is an inherited disorder that presents as small-vessel strokes, progressive dementia, and extensive symmetric white matter changes often including the anterior temporal lobes visualized by MRI. Approximately 40% of patients have migraine with aura, often manifest as transient motor or sensory deficits. Onset is usually in the fourth or fifth decade of life. This autosomal dominant condition is caused by one of several mutations in *Notch-3*, a member of a highly conserved gene family characterized by epidermal growth factor repeats in its extracellular domain. Other monogenic ischemic stroke syndromes include cerebral autosomal recessive arteriopathy with subcortical infarcts and leukoencephalopathy (CARASIL) and hereditary endotheliopathy, retinopathy, nephropathy, and stroke (HERNS). Fabry’s disease also produces both a large-vessel arteriopathy and small-vessel infarctions.

TRANSIENT ISCHEMIC ATTACKS

TIAs are episodes of stroke symptoms that last only briefly; the standard definition of duration is <24 h, but most TIAs last <1 h. If a relevant brain infarction is identified on brain imaging, the clinical entity is now classified as stroke regardless of the duration of symptoms. The causes of TIA are similar to the causes of ischemic stroke, but because TIAs may herald stroke, they are an important risk factor that should be considered separately and urgently. TIAs may arise from emboli to the brain or from in situ thrombosis of an intracranial vessel. With a TIA, the occluded blood vessel reopens and neurologic function is restored.

The risk of stroke after a TIA is ~10–15% in the first 3 months, with most events occurring in the first 2 days. This risk can be directly estimated using the well-validated ABCD² score (Table 446-5). Therefore, urgent evaluation and treatment are justified. Because etiologies for stroke and TIA are identical, evaluation for TIA should parallel that of stroke (Figs. 446-1 and 446-3). The improvement characteristic of TIA is a contraindication to thrombolysis. However, because the risk of subsequent stroke in the first few days after a TIA is high, the opportunity to give rtPA rapidly if a stroke occurs may justify hospital admission for most patients. The combination of aspirin and clopidogrel has been recently reported to prevent stroke following TIA better than aspirin alone in a large Chinese randomized trial and is undergoing similar evaluation in an ongoing National Institutes of Health (NIH)-sponsored trial (POINT study).

TABLE 446-5 RISK OF STROKE FOLLOWING TRANSIENT ISCHEMIC ATTACK: THE ABCD² SCORE

Clinical Factor	Score
A: Age ≥ 60 years	1
B: SBP >140 mmHg or DBP >90 mmHg	1
C: Clinical symptoms	
Unilateral weakness	2
Speech disturbance without weakness	1
D: Duration	
>60 min	2
10–59 min	1
D: Diabetes (oral medications or insulin)	1
Total Score	Sum Each Category
ABCD ² Score Total	3-Month Rate of Stroke (%) ^a
0	0
1	2
2	3
3	3
4	8
5	12
6	17
7	22

^aData ranges are from five cohorts.

Abbreviations: DBP, diastolic blood pressure; SBP, systolic blood pressure.

Source: SC Johnston et al. Validation and refinement of score to predict very early stroke risk after transient ischaemic attack. *Lancet* 369:283, 2007.

TREATMENT PRIMARY AND SECONDARY PREVENTION OF STROKE AND TIA

GENERAL PRINCIPLES

A number of medical and surgical interventions, as well as lifestyle modifications, are available for preventing stroke. Some of these can be widely applied because of their low cost and minimal risk; others are expensive and carry substantial risk but may be valuable for selected high-risk patients. Identification and control of modifiable risk factors, and especially hypertension, is the best strategy to reduce the burden of stroke, and the total number of strokes could be reduced substantially by these means (Table 446-4).

ATHEROSCLEROSIS RISK FACTORS

The relationship of various factors to the risk of atherosclerosis is described in [Chap. 291e](#). Older age, diabetes mellitus, hypertension, tobacco smoking, abnormal blood cholesterol (particularly, low high-density lipoprotein [HDL] and/or elevated low-density lipoprotein [LDL]), and other factors are either proven or probable risk factors for ischemic stroke, largely by their link to atherosclerosis. Risk of stroke is much greater in those with prior stroke or TIA. Many cardiac conditions predispose to stroke, including atrial fibrillation and recent MI. Oral contraceptives and hormone replacement therapy increase stroke risk, and although rare, certain inherited and acquired hypercoagulable states predispose to stroke.

Hypertension is the most significant of the risk factors; in general, all hypertension should be treated to a target of less than 140–150/90 mmHg. However, many vascular neurologists recommend that guidelines for secondary prevention of stroke should aim for blood pressure reduction to 130/80 mmHg or lower. The presence of known cerebrovascular disease is not a contraindication to treatment aimed at achieving normotension. Also, the value of treating systolic hypertension in older patients has been clearly established. Lowering blood pressure to levels below those traditionally defining hypertension appears to reduce the risk of stroke even further. Data are particularly strong in support of thiazide diuretics and angiotensin-converting enzyme inhibitors.

Several trials have confirmed that statin drugs reduce the risk of stroke even in patients without elevated LDL or low HDL. The Stroke Prevention by Aggressive Reduction in Cholesterol Levels (SPARCL) trial showed benefit in secondary stroke reduction for patients with recent stroke or TIA who were prescribed atorvastatin, 80 mg/d. The primary prevention trial, Justification for the Use of Statins in Prevention: An Intervention Trial Evaluating Rosuvastatin (JUPITER), found that patients with low LDL (<130 mg/dL) caused by elevated C-reactive protein benefitted by daily use of this statin. Primary stroke occurrence was reduced by 51% (hazard ratio 0.49, $p = .004$), and there was no increase in the rates of intracranial hemorrhage. Meta-analysis has also supported a primary treatment effect for statins given acutely for ischemic stroke. Therefore, a statin should be considered in all patients with prior ischemic stroke. Tobacco smoking should be discouraged in all patients ([Chap. 470](#)). The use of pioglitazone (an agonist of peroxisome proliferator-activated receptor gamma) in patients with type 2 diabetes and previous stroke may lower risk of recurrent stroke, MI, or vascular death, but no trial sufficiently powered to definitively detect a significant reduction in stroke in the general diabetic population has yet been performed.

ANTIPLATELET AGENTS

Platelet antiaggregation agents can prevent atherothrombotic events, including TIA and stroke, by inhibiting the formation of intraarterial platelet aggregates. These can form on diseased arteries, induce thrombus formation, and occlude or embolize into the distal circulation. Aspirin, clopidogrel, and the combination of aspirin plus extended-release dipyridamole are the antiplatelet agents most commonly used for this purpose. Ticlopidine has been largely abandoned because of its adverse effects but may be used as an alternative to clopidogrel.

Aspirin is the most widely studied antiplatelet agent. Aspirin acetylates platelet cyclooxygenase, which irreversibly inhibits the formation in platelets of thromboxane A₂, a platelet aggregating and vasoconstricting prostaglandin. This effect is permanent and lasts for the usual 8-day life of the platelet. Paradoxically, aspirin also inhibits the formation in endothelial cells of prostacyclin, an antiaggregating and vasodilating prostaglandin. This effect is transient. As soon as aspirin is cleared from the blood, the nucleated endothelial cells again produce prostacyclin. Aspirin in low doses given once daily inhibits the production of thromboxane A₂ in platelets without substantially inhibiting prostacyclin formation. Higher doses of aspirin have not been proven to be more effective than lower doses.

Ticlopidine and clopidogrel block the adenosine diphosphate (ADP) receptor on platelets and thus prevent the cascade resulting in activation of the glycoprotein IIb/IIIa receptor that leads to fibrinogen binding to the platelet and consequent platelet aggregation. Ticlopidine is more effective than aspirin; however, it has the disadvantage of causing diarrhea, skin rash, and, in rare instances, neutropenia and thrombotic thrombocytopenic purpura (TTP). Clopidogrel rarely causes TTP but does not cause neutropenia. The Clopidogrel versus Aspirin in Patients at Risk of Ischemic Events (CAPRIE) trial, which led to FDA approval, found that it was only marginally more effective than aspirin in reducing risk of stroke. The Management of Atherothrombosis with Clopidogrel in High-Risk Patients (MATCH) trial was a large multicenter, randomized, double-blind study that compared clopidogrel in combination with aspirin to clopidogrel alone in the secondary prevention of TIA or stroke. The MATCH trial found no difference in TIA or stroke prevention with this combination, but did show a small but significant increase in major bleeding complications (3 vs 1%). In the Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management, and Avoidance (CHARISMA) trial, which included a subgroup of patients with prior stroke or TIA along with other groups at high risk of cardiovascular events, there was no benefit of clopidogrel combined with aspirin compared to aspirin alone. Lastly, the SPS3 trial looked at the long-term combination of clopidogrel and aspirin versus clopidogrel alone in small-vessel stroke and found no

2570 improvement in stroke prevention and a significant increase in both hemorrhage and death. Thus, the long-term use of clopidogrel in combination with aspirin is not recommended for stroke prevention.

The short-term combination of clopidogrel with aspirin may be effective in preventing second stroke, however. A trial of 5170 Chinese patients enrolled within 24 h of TIA or minor ischemic stroke found that a clopidogrel-aspirin regimen (clopidogrel 300 mg load then 75 mg/d with aspirin 75 mg for the first 21 days) was superior to aspirin (75 mg/d) alone, with 90-day stroke risk decreased from 11.7 to 8.2% ($p < .001$) and no increase in major hemorrhage. An international NIH-sponsored trial of similar design is ongoing.

Dipyridamole is an antiplatelet agent that inhibits the uptake of adenosine by a variety of cells, including those of the vascular endothelium. The accumulated adenosine is an inhibitor of aggregation. At least in part through its effects on platelet and vessel wall phosphodiesterases, dipyridamole also potentiates the antiaggregatory effects of prostacyclin and nitric oxide produced by the endothelium and acts by inhibiting platelet phosphodiesterase, which is responsible for the breakdown of cyclic AMP. The resulting elevation in cyclic AMP inhibits aggregation of platelets. Dipyridamole is erratically absorbed depending on stomach pH, but a newer formulation combines timed-release dipyridamole, 200 mg, with aspirin, 25 mg, and has better oral bioavailability. This combination drug was studied in three trials. The European Stroke Prevention Study (ESPS) II showed efficacy of both 50 mg/d of aspirin and extended-release dipyridamole in preventing stroke, and a significantly better risk reduction when the two agents were combined. The open-label ESPRIT (European/Australasian Stroke Prevention in Reversible Ischaemia Trial) trial confirmed the ESPS-II results. After 3.5 years of follow-up, 13% of patients on aspirin and dipyridamole and 16% on aspirin alone (hazard ratio 0.80, 95% confidence index [CI] 0.66–0.98) met the primary outcome of death from all vascular causes. In the Prevention Regimen for Effectively Avoiding Second Strokes (PROFESS) trial, the combination of extended-release dipyridamole and aspirin was compared directly with clopidogrel with and without the angiotensin receptor blocker telmisartan; there were no differences in the rates of second stroke (9% each) or degree of disability in patients with median follow-up of 2.4 years. Telmisartan also had no effect on these outcomes. This suggests that these antiplatelet regimens are similar and also raises questions about default prescription of agents to block the angiotensin pathway in all stroke patients. The principal side effect of dipyridamole is headache. The combination capsule of extended-release dipyridamole and aspirin is approved for prevention of stroke.

Many large clinical trials have demonstrated clearly that most antiplatelet agents reduce the risk of all important vascular atherothrombotic events (i.e., ischemic stroke, MI, and death due to all vascular causes) in patients at risk for these events. The overall *relative* reduction in risk of nonfatal stroke is about 25–30% and of all vascular events is about 25%. The *absolute* reduction varies considerably, depending on the particular patient's risk. Individuals at very low risk for stroke seem to experience the same relative reduction, but their risks may be so low that the "benefit" is meaningless. Conversely, individuals with a 10–15% risk of vascular events per year experience a reduction to about 7.5–11%.

Aspirin is inexpensive, can be given in low doses, and could be recommended for all adults to prevent both stroke and MI. However, it causes epigastric discomfort, gastric ulceration, and gastrointestinal hemorrhage, which may be asymptomatic or life threatening. Consequently, not every 40- or 50-year-old should be advised to take aspirin regularly because the risk of atherothrombotic stroke is extremely low and is outweighed by the risk of adverse side effects. Conversely, every patient who has experienced an atherothrombotic stroke or TIA and has no contraindication should be taking an antiplatelet agent regularly because the average annual risk of another stroke is 8–10%; another few percent will experience an MI or vascular death. Clearly, the likelihood of benefit far outweighs the risks of treatment.

The choice of antiplatelet agent and dose must balance the risk of stroke, the expected benefit, and the risk and cost of treatment. However, there are no definitive data, and opinions vary. Many authorities believe low-dose (30–75 mg/d) and high-dose (650–1300 mg/d) aspirin are about equally effective. Some advocate very low doses to avoid adverse effects, and still others advocate very high doses to be sure the benefit is maximal. Most physicians in North America recommend 81–325 mg/d, whereas most Europeans recommend 50–100 mg. Clopidogrel and extended-release dipyridamole plus aspirin are being increasingly recommended as first-line drugs for secondary prevention. Similarly, the choice of aspirin, clopidogrel, or dipyridamole plus aspirin must balance the fact that the latter are more effective than aspirin but the cost is higher, and this is likely to affect long-term patient adherence. The use of platelet aggregation studies in individual patients taking aspirin is controversial because of limited data.

ANTICOAGULATION THERAPY AND EMBOLIC STROKE

Several trials have shown that anticoagulation (INR range, 2–3) in patients with chronic nonvalvular (nonrheumatic) atrial fibrillation (NVAF) prevents cerebral embolism and stroke and is safe. For primary prevention and for patients who have experienced stroke or TIA, anticoagulation with a VKA reduces the risk by about 67%, which clearly outweighs the 1–3% risk per year of a major bleeding complication. VKAs are difficult to dose, their effects vary with dietary intake of vitamin K, and they require frequent blood monitoring of the PTT/INR. Several newer oral anticoagulants (OACs) have recently been shown to be more convenient and efficacious for stroke prevention in NVAF. A randomized trial compared the oral thrombin inhibitor dabigatran to VKAs in a noninferiority trial to prevent stroke or systemic embolization in NVAF. Two doses of dabigatran were used: 110 mg/d and 150 mg/d. Both dose tiers of dabigatran were noninferior to VKAs in preventing second stroke and systemic embolization, and the higher dose tier was superior (relative risk 0.66; 95% CI 0.53–0.82; $p < .001$) and the rate of major bleeding was lower in the lower dose tier of dabigatran compared to VKAs. Dabigatran requires no blood monitoring to titrate the dose, and its effect is independent of oral intake of vitamin K. Newer oral factor Xa inhibitors have also been found to be equivalent or safer and more effective than VKAs in NVAF stroke prevention. In the Apixaban for Reduction in Stroke and Other Thromboembolic Events in Atrial Fibrillation (ARISTOTLE) trial, patients were randomized between apixaban, 5 mg twice daily, and dose-adjusted warfarin (INR 2–3). The combined endpoint of ischemic or hemorrhagic stroke or system embolism occurred in 1.27% of patients in the apixaban group and in 1.6% in the warfarin group ($p < .001$ for noninferiority and $p < .01$ for superiority). Major bleeding was 1% less, favoring apixaban ($p < .001$). Similar results were obtained in the Rivaroxaban Once Daily Oral Direct Factor Xa Inhibition Compared with Vitamin K Antagonism for Prevention of Stroke and Embolism Trial in Atrial Fibrillation (ROCKET-AF). Here, patients with NVAF were randomized to rivaroxaban versus warfarin: 1.7% of the factor Xa group and 2.2% of the warfarin group reached the endpoint of stroke and systemic embolism ($p < .001$ for noninferiority); intracranial hemorrhage was also lower with rivaroxaban. Finally, the factor Xa inhibitor edoxaban was also found to be noninferior to warfarin. Thus, oral factor Xa inhibitors are at least a suitable alternative to VKAs, and likely are superior both in efficacy and perhaps compliance.

For patients who cannot take anticoagulant medications, clopidogrel plus aspirin was compared to aspirin alone in the Atrial Fibrillation Clopidogrel Trial with Irbesartan for Prevention of Vascular Events (ACTIVE-A). Clopidogrel combined with aspirin was more effective than aspirin alone in preventing vascular events, principally stroke, but increased the risk of major bleeding (relative risk 1.57, $p < .001$).

The decision to use anticoagulation for primary prevention is based primarily on risk factors (Table 446-3). The history of a TIA or stroke tips the balance in favor of anticoagulation regardless

of other risk factors. Intermittent atrial fibrillation carries the same risk of stroke as chronic atrial fibrillation, and several ambulatory studies of seemingly “cryptogenic” stroke have found evidence of intermittent atrial fibrillation in nearly 20% of patients monitored for a few weeks. Interrogation of implanted pacemakers also confirms an association between subclinical atrial fibrillation and stroke risk. Therefore, for patients with otherwise cryptogenic embolic stroke (no evidence of any other cause for stroke), ambulatory monitoring for 3–4 weeks is a reasonable strategy to determine the best prophylactic therapy.

Because of the high annual stroke risk in untreated rheumatic heart disease with atrial fibrillation, primary prophylaxis against stroke has not been studied in a double-blind fashion. These patients generally should receive long-term anticoagulation. Dabigatran and the oral Xa inhibitors have not been studied in this population.

Anticoagulation also reduces the risk of embolism in acute MI. Most clinicians recommend a 3-month course of anticoagulation when there is anterior Q-wave infarction, substantial left ventricular dysfunction, congestive heart failure, mural thrombosis, or atrial fibrillation. OACs are recommended long-term if atrial fibrillation persists.

Stroke secondary to thromboembolism is one of the most serious complications of prosthetic heart valve implantation. The intensity of anticoagulation and/or antiplatelet therapy is dictated by the type of prosthetic valve and its location. Dabigatran may be less effective than warfarin, and the oral Xa inhibitors have not been studied in this population.

If the embolic source cannot be eliminated, anticoagulation should in most cases be continued indefinitely. Many neurologists recommend combining antiplatelet agents with anticoagulants for patients who “fail” anticoagulation (i.e., have another stroke or TIA), but the evidence basis for this is lacking.

ANTICOAGULATION THERAPY AND NONCARDIOGENIC STROKE

Data do not support the use of long-term VKAs for preventing atherothrombotic stroke for either intracranial or extracranial cerebrovascular disease. The Warfarin-Aspirin Recurrent Stroke Study (WARSS) found no benefit of warfarin sodium (INR 1.4–2.8) over aspirin, 325 mg, for secondary prevention of stroke but did find a slightly higher bleeding rate in the warfarin group; a European study confirmed this finding. The Warfarin and Aspirin for Symptomatic Intracranial Disease (WASID) study (see below) demonstrated no benefit of warfarin (INR 2–3) over aspirin in patients with symptomatic intracranial atherosclerosis and also found a higher rate of bleeding complications.

TREATMENT CAROTID ATHEROSCLEROSIS

Carotid atherosclerosis can be removed surgically (endarterectomy) or mitigated with endovascular stenting with or without balloon angioplasty. Anticoagulation has not been directly compared with antiplatelet therapy for carotid disease.

SURGICAL THERAPY

Symptomatic carotid stenosis was studied in the North American Symptomatic Carotid Endarterectomy Trial (NASCET) and the European Carotid Surgery Trial (ECST). Both showed a substantial benefit for surgery in patients with stenosis of $\geq 70\%$. In NASCET, the average cumulative ipsilateral stroke risk at 2 years was 26% for patients treated medically and 9% for those receiving the same medical treatment plus a carotid endarterectomy. This 17% *absolute* reduction in the surgical group is a 65% *relative* risk reduction favoring surgery (Table 446-4). NASCET also showed a significant, although less robust, benefit for patients with 50–70% stenosis. ECST found harm for patients with stenosis $< 30\%$ treated surgically.

A patient’s risk of stroke and possible benefit from surgery are related to the presence of retinal versus hemispheric symptoms, degree of arterial stenosis, extent of associated medical conditions

(of note, NASCET and ECST excluded “high-risk” patients with significant cardiac, pulmonary, or renal disease), institutional surgical morbidity and mortality, timing of surgery relative to symptoms, and other factors. A recent meta-analysis of the NASCET and ECST trials demonstrated that endarterectomy is most beneficial when performed within 2 weeks of symptom onset. In addition, benefit is more pronounced in patients > 75 years, and men appear to benefit more than women.

In summary, a patient with recent symptomatic hemispheric ischemia, high-grade stenosis in the appropriate internal carotid artery, and an institutional perioperative morbidity and mortality rate of $\leq 6\%$ generally should undergo carotid endarterectomy. If the perioperative stroke rate is $> 6\%$ for any particular surgeon, however, the benefits of carotid endarterectomy are questionable.

The indications for surgical treatment of *asymptomatic carotid disease* have been clarified by the results of the Asymptomatic Carotid Atherosclerosis Study (ACAS) and the Asymptomatic Carotid Surgery Trial (ACST). ACAS randomized asymptomatic patients with $\geq 60\%$ stenosis to medical treatment with aspirin or the same medical treatment plus carotid endarterectomy. The surgical group had a risk over 5 years for ipsilateral stroke (and any perioperative stroke or death) of 5.1%, compared to a risk in the medical group of 11%. Although this demonstrates a 53% *relative* risk reduction, the *absolute* risk reduction is only 5.9% over 5 years, or 1.2% annually (Table 446-4). Nearly one-half of the strokes in the surgery group were caused by preoperative angiograms. ACST randomized asymptomatic patients with $> 60\%$ carotid stenosis to endarterectomy or medical therapy. The 5-year risk of stroke in the surgical group (including perioperative stroke or death) was 6.4%, compared to 11.8% in the medically treated group (46% relative risk reduction and 5.4% absolute risk reduction).

In both ACAS and ACST, the perioperative complication rate was higher in women, perhaps negating any benefit in the reduction of stroke risk within 5 years. It is possible that with longer follow-up, a clear benefit in women will emerge. At present, carotid endarterectomy in asymptomatic women remains particularly controversial.

In summary, the natural history of asymptomatic stenosis is an $\sim 2\%$ per year stroke rate, whereas symptomatic patients experience a 13% per year risk of stroke. Whether to recommend carotid revascularization for an asymptomatic patient is somewhat controversial and depends on many factors, including patient preference, degree of stenosis, age, gender, and comorbidities. Medical therapy for reduction of atherosclerosis risk factors, including cholesterol-lowering agents and antiplatelet medications, is generally recommended for patients with asymptomatic carotid stenosis. As with atrial fibrillation, it is imperative to counsel the patient about TIAs so that therapy can be revised if symptoms develop.

ENDOVASCULAR THERAPY

Balloon angioplasty coupled with stenting is being used with increasing frequency to open stenotic carotid arteries and maintain their patency. These techniques can treat carotid stenosis not only at the bifurcation but also near the skull base and in the intracranial segments. The Stenting and Angioplasty with Protection in Patients at High Risk for Endarterectomy (SAPPHIRE) trial randomized high-risk patients (defined as patients with clinically significant coronary or pulmonary disease, contralateral carotid occlusion, restenosis after endarterectomy, contralateral laryngeal-nerve palsy, prior radical neck surgery or radiation, or age > 80) with symptomatic carotid stenosis $> 50\%$ or asymptomatic stenosis $> 80\%$ to either stenting combined with a distal emboli-protection device or endarterectomy. The risk of death, stroke, or MI within 30 days and ipsilateral stroke or death within 1 year was 12.2% in the stenting group and 20.1% in the endarterectomy group ($p = .055$), suggesting that stenting is at the very least comparable to endarterectomy as a treatment option for this patient group at high risk of surgery. However, the outcomes with both interventions may not have been better than leaving the carotid stenoses untreated, particularly for the asymptomatic patients, and much of the benefit seen in the

stenting group was due to a reduction in periprocedure MI. Two randomized trials comparing stents to endarterectomy in lower-risk patients have been published. The Carotid Revascularization Endarterectomy versus Stenting Trial (CREST) enrolled patients with either asymptomatic or symptomatic stenosis. The 30-day risk of stroke was 4.1% in the stent group and 2.3% in the surgical group, but the 30-day risk of MI was 1.1% in the stent group and 2.3% in the surgery group, suggesting relative equivalence of risk between the procedures. At median follow-up of 2.5 years, the combined endpoint of stroke, MI, and death was the same (7.2% stent vs 6.8% surgery). The rate of restenosis at 2 years was also similar in both groups. The International Carotid Stenting (ICSS) trial randomized symptomatic patients to stents versus endarterectomy and found a different result: At 120 days, the incidence of stroke, MI, or death was 8.5% in the stenting group versus 5.2% in the endarterectomy group ($p = .006$); longer-term follow-up is currently under way. Differences between trial designs, selection of stent, and operator experience may explain these important differences. Until more data are available, there remains controversy as to who should receive a stent or have endarterectomy; it is likely that the procedures carry similar risks if performed by experienced physicians.

BYPASS SURGERY

Extracranial-to-intracranial (EC-IC) bypass surgery has been proven ineffective for atherosclerotic stenoses that are inaccessible to conventional carotid endarterectomy. In patients with recent stroke, an associated carotid occlusion, and evidence of inadequate perfusion of the brain as measured with positron emission tomography, no benefit from EC-IC bypass was found in a trial stopped for futility.

INTRACRANIAL ATHEROSCLEROSIS

The WASID trial randomized patients with symptomatic stenosis (50–99%) of a major intracranial vessel to either high-dose aspirin (1300 mg/d) or warfarin (target INR, 2.0–3.0), with a combined primary endpoint of ischemic stroke, brain hemorrhage, or death from vascular cause other than stroke. The trial was terminated early because of an increased risk of adverse events related to warfarin anticoagulation. With a mean follow-up of 1.8 years, the primary endpoint was seen in 22.1% of patients in the aspirin group and 21.8% of the warfarin group. Death from any cause was seen in 4.3% of the aspirin group and 9.7% of the warfarin group; 3.2% of patients on aspirin experienced major hemorrhage, compared to 8.3% of patients taking warfarin.

Intracranial stenting of intracranial atherosclerosis was found to be dramatically harmful compared to aspirin in the Stenting and Aggressive Medical Management for Preventing Recurrent Stroke in Intracranial Stenosis (SAMMPRIS) trial. This trial enrolled newly symptomatic TIA or minor stroke patients with associated 70–99% intracranial stenosis to primary stenting with a self-expanding stent or to medical management. Both groups received clopidogrel, aspirin, statin, and aggressive control of blood pressure. The endpoint of stroke or death occurred in 14.7% of the stented group and 5.8% of the medically treated groups ($p = .002$). This low rate of second stroke was significantly lower than in the WASID trial and suggests that aggressive medical management had a marked influence on secondary stroke risk.

Dural Sinus Thrombosis Limited evidence exists to support short-term use of anticoagulants, regardless of the presence of intracranial hemorrhage, for venous infarction following sinus thrombosis. The long-term outcome for most patients, even those with intracerebral hemorrhage, is excellent.

STROKE SYNDROMES

A careful history and neurologic examination can often localize the region of brain dysfunction; if this region corresponds to a particular arterial distribution, the possible causes responsible for the syndrome can be narrowed. This is of particular importance when the patient

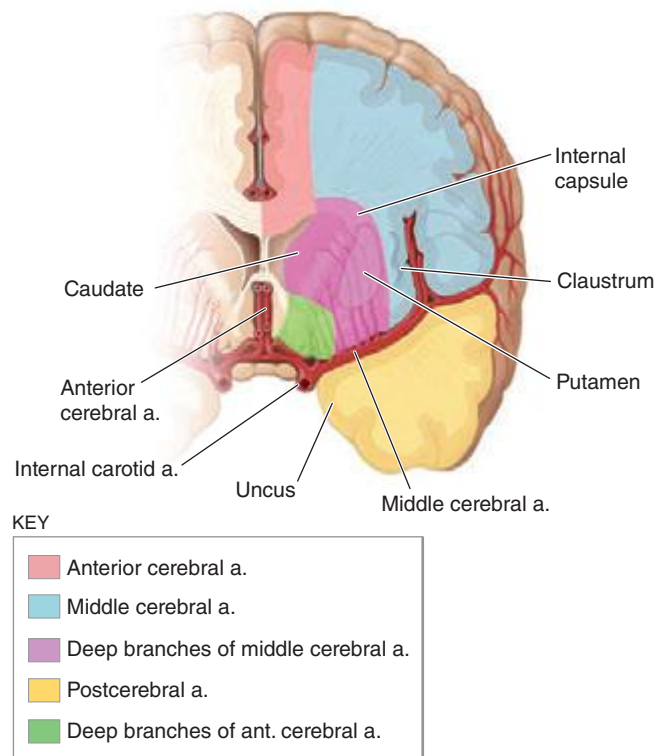


FIGURE 446-6 Diagram of a cerebral hemisphere in coronal section showing the territories of the major cerebral vessels that branch from the internal carotid arteries.

presents with a TIA and a normal examination. For example, if a patient develops language loss and a right homonymous hemianopia, a search for causes of left middle cerebral emboli should be performed. A finding of an isolated stenosis of the right internal carotid artery in that patient, for example, suggests an asymptomatic carotid stenosis, and the search for other causes of stroke should continue. The following sections describe the clinical findings of cerebral ischemia associated with cerebral vascular territories depicted in Figs. 446-4 and 446-6 through 446-14. Stroke syndromes are divided into: (1) large-vessel stroke within the anterior circulation, (2) large-vessel stroke within the posterior circulation, and (3) small-vessel disease of either vascular bed.

Stroke within the Anterior Circulation The internal carotid artery and its branches comprise the anterior circulation of the brain. These vessels can be occluded by intrinsic disease of the vessel (e.g., atherosclerosis or dissection) or by embolic occlusion from a proximal source as discussed above. Occlusion of each major intracranial vessel has distinct clinical manifestations.

MIDDLE CEREBRAL ARTERY Occlusion of the proximal MCA or one of its major branches is most often due to an embolus (artery-to-artery, cardiac, or of unknown source) rather than intracranial atherothrombosis. Atherosclerosis of the proximal MCA may cause distal emboli to the middle cerebral territory or, less commonly, may produce low-flow TIAs. Collateral formation via leptomeningeal vessels often prevents MCA stenosis from becoming symptomatic.

The cortical branches of the MCA supply the lateral surface of the hemisphere except for (1) the frontal pole and a strip along the superomedial border of the frontal and parietal lobes supplied by the ACA, and (2) the lower temporal and occipital pole convolutions supplied by the PCA (Figs. 446-6, 446-7, 446-8, and 446-9).

The proximal MCA (M1 segment) gives rise to penetrating branches (termed *lenticulostriate arteries*) that supply the putamen, outer globus pallidus, posterior limb of the internal capsule, adjacent corona radiata, and most of the caudate nucleus (Fig. 446-6). In the sylvian fissure, the MCA in most patients divides into *superior* and *inferior* divisions

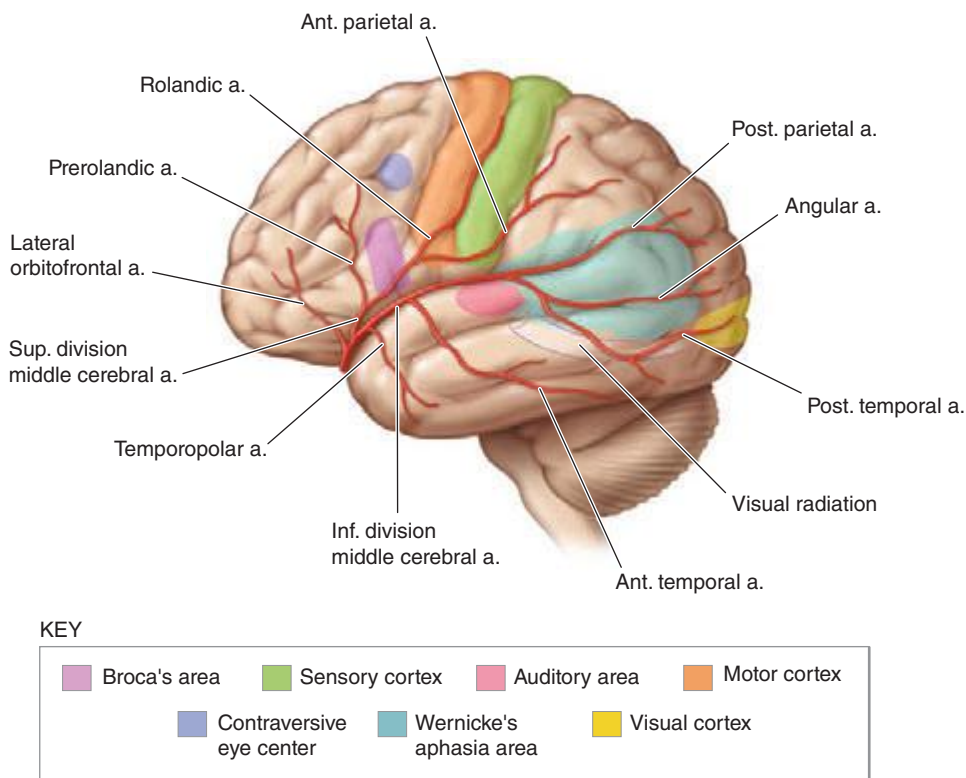


FIGURE 446-7 Diagram of a cerebral hemisphere, lateral aspect, showing the branches and distribution of the middle cerebral artery and the principal regions of cerebral localization. Note the bifurcation of the middle cerebral artery into a superior and inferior division.

Signs and symptoms: Structures involved

Paralysis of the contralateral face, arm, and leg; sensory impairment over the same area (pin-prick, cotton touch, vibration, position, two-point discrimination, stereognosis, tactile localization, barognosis, cutaneographia): *Somatic motor area for face and arm and the fibers descending from the leg area to enter the corona radiata and corresponding somatic sensory system*

Motor aphasia: *Motor speech area of the dominant hemisphere*

Central aphasia, word deafness, anomia, jargon speech, sensory agraphia, acalculia, alexia, finger agnosia, right-left confusion (the last four comprise the Gerstmann syndrome): *Central, suprasylvian speech area and parietooccipital cortex of the dominant hemisphere*

Conduction aphasia: *Central speech area (parietal operculum)*

Apractagnosia of the nondominant hemisphere, anosognosia, hemiasomatognosia, unilateral neglect, agnosia for the left half of external space, dressing "apraxia," constructional "apraxia," distortion of visual coordinates, inaccurate localization in the half field, impaired ability to judge distance, upside-down reading, visual illusions (e.g., it may appear that another person walks through a table): *Nondominant parietal lobe (area corresponding to speech area in dominant hemisphere); loss of topographic memory is usually due to a nondominant lesion, occasionally to a dominant one*

Homonymous hemianopia (often homonymous inferior quadrantanopia): *Optic radiation deep to second temporal convolution*

Paralysis of conjugate gaze to the opposite side: *Frontal contraversive eye field or projecting fibers*

(M2 branches). Branches of the inferior division supply the inferior parietal and temporal cortex, and those from the superior division supply the frontal and superior parietal cortex (Fig. 446-7).

If the entire MCA is occluded at its origin (blocking both its penetrating and cortical branches) and the distal collaterals are limited, the clinical findings are contralateral hemiplegia, hemianesthesia, homonymous hemianopia, and a day or two of gaze preference to the ipsilateral side. Dysarthria is common because of facial weakness. When the dominant hemisphere is involved, global aphasia is present also, and when the nondominant hemisphere is affected, anosognosia, constructional apraxia, and neglect are found (Chap. 36).

Complete MCA syndromes occur most often when an embolus occludes the stem of the artery. Cortical collateral blood flow and differing arterial configurations are probably responsible for the development of many partial syndromes. Partial syndromes may also be due to emboli that enter the proximal MCA without complete occlusion,

occlude distal MCA branches, or fragment and move distally.

Partial syndromes due to embolic occlusion of a single branch include hand, or arm and hand, weakness alone (brachial syndrome) or facial weakness with nonfluent (Broca) aphasia (Chap. 36), with or without arm weakness (frontal opercular syndrome). A combination of sensory disturbance, motor weakness, and nonfluent aphasia suggests that an embolus has occluded the proximal superior division and infarcted large portions of the frontal and parietal cortices (Fig. 446-7). If a fluent (Wernicke's) aphasia occurs without weakness, the inferior division of the MCA supplying the posterior part (temporal cortex) of the dominant hemisphere is probably involved. Jargon speech and an inability to comprehend written and spoken language are prominent features, often accompanied by a contralateral, homonymous superior quadrantanopia. Hemineglect or spatial agnosia without weakness indicates that the inferior division of the MCA in the nondominant hemisphere is involved.

Occlusion of a lenticulostriate vessel produces small-vessel (lacunar) stroke within the internal capsule (Fig. 446-6). This produces pure motor stroke or sensory-motor stroke contralateral to the lesion. Ischemia within the genu of the internal capsule causes primarily facial weakness followed by arm and then leg weakness as the ischemia moves posterior within the capsule. Alternatively, the contralateral hand may become ataxic, and dysarthria will be prominent (clumsy hand, dysarthria lacunar syndrome). Lacunar infarction affecting the globus pallidus and putamen often has few clinical signs, but parkinsonism and hemibalismus have been reported.

ANTERIOR CEREBRAL ARTERY The ACA is divided into two segments: the precommunal (A1) circle of Willis, or stem, which connects the internal carotid artery to the anterior communicating artery, and the postcommunal (A2) segment distal to the anterior communicating artery (Figs. 446-4, 446-6, and 446-8). The A1 seg-

ment gives rise to several deep penetrating branches that supply the anterior limb of the internal capsule, the anterior perforate substance, amygdala, anterior hypothalamus, and the inferior part of the head of the caudate nucleus (Fig. 446-6).

Occlusion of the proximal ACA is usually well tolerated because of collateral flow through the anterior communicating artery and collaterals through the MCA and PCA. Occlusion of a single A2 segment results in the contralateral symptoms noted in Fig. 446-8. If both A2 segments arise from a single anterior cerebral stem (contralateral A1 segment atresia), the occlusion may affect both hemispheres. Profound abulia (a delay in verbal and motor response) and bilateral pyramidal signs with paraparesis or quadriparesis and urinary incontinence result.

ANTERIOR CHOROIDAL ARTERY This artery arises from the internal carotid artery and supplies the posterior limb of the internal capsule and the white matter posterolateral to it, through which pass some of

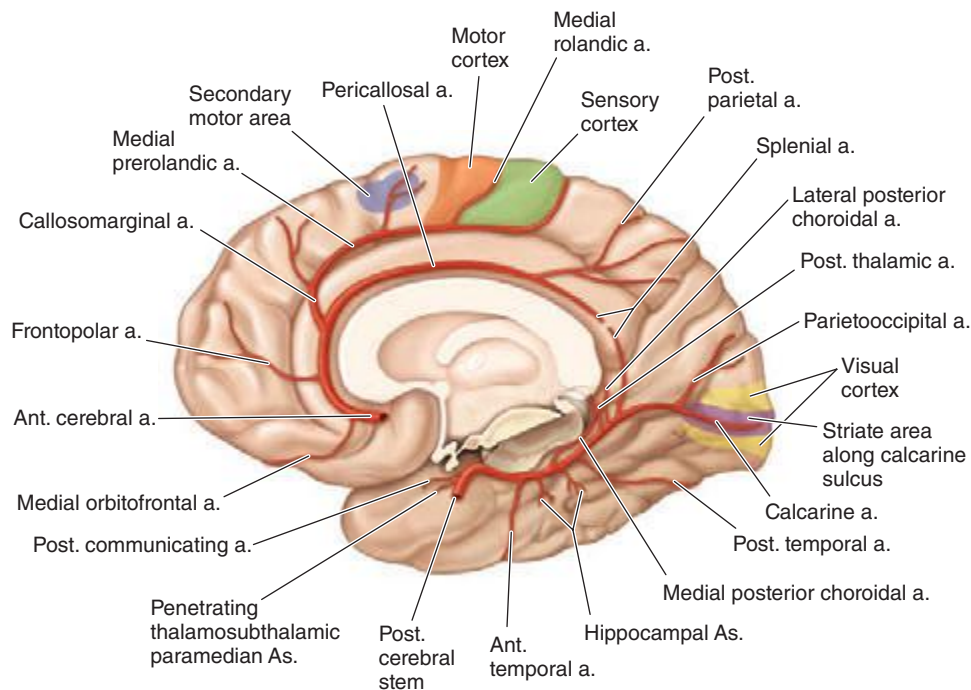


FIGURE 446-8 Diagram of a cerebral hemisphere, medial aspect, showing the branches and distribution of the anterior cerebral artery and the principal regions of cerebral localization.

Signs and symptoms: *Structures involved*

Paralysis of opposite foot and leg: *Motor leg area*

A lesser degree of paresis of opposite arm: *Arm area of cortex or fibers descending to corona radiata*

Cortical sensory loss over toes, foot, and leg: *Sensory area for foot and leg*

Urinary incontinence: *Sensorimotor area in paracentral lobule*

Contralateral grasp reflex, sucking reflex, gegenhalten (paratonic rigidity): *Medial surface of the posterior frontal lobe; likely supplemental motor area*

Abulia (akinetic mutism), slowness, delay, intermittent interruption, lack of spontaneity, whispering, reflex distraction to sights and sounds:

Uncertain localization—probably cingulate gyrus and medial inferior portion of frontal, parietal, and temporal lobes

Impairment of gait and stance (gait apraxia): *Frontal cortex near leg motor area*

Dyspraxia of left limbs, tactile aphasia in left limbs: *Corpus callosum*

the geniculocalcarine fibers (Fig. 446-9). The complete syndrome of anterior choroidal artery occlusion consists of contralateral hemiplegia, hemianesthesia (hypesthesia), and homonymous hemianopia. However, because this territory is also supplied by penetrating vessels of the proximal MCA and the posterior communicating and posterior choroidal arteries, minimal deficits may occur, and patients frequently recover substantially. Anterior choroidal strokes are usually the result of in situ thrombosis of the vessel, and the vessel is particularly vulnerable to iatrogenic occlusion during surgical clipping of aneurysms arising from the internal carotid artery.

INTERNAL CAROTID ARTERY The clinical picture of internal carotid occlusion varies depending on whether the cause of ischemia is propagated thrombus, embolism, or low flow. The cortex supplied by the MCA territory is affected most often. With a competent circle of Willis, occlusion may go unnoticed. If the thrombus propagates up the internal carotid artery into the MCA or embolizes it, symptoms are identical to proximal MCA occlusion (see above). Sometimes there is massive infarction of the entire deep white matter and cortical surface. When the origins of both the ACA and MCA are occluded at the top of the carotid artery, abulia or stupor occurs with hemiplegia, hemianesthesia, and aphasia or anosognosia. When the PCA arises from the internal carotid artery (a configuration called a *fetal posterior cerebral artery*), it may also become occluded and give rise to symptoms referable to its peripheral territory (Figs. 446-8 and 446-9).

In addition to supplying the ipsilateral brain, the internal carotid artery perfuses the optic nerve and retina via the ophthalmic artery. In ~25% of symptomatic internal carotid disease, recurrent transient monocular blindness (amaurosis fugax) warns of the lesion. Patients typically describe a horizontal shade that sweeps down or up across the field of vision. They may also complain that their vision was blurred

in that eye or that the upper or lower half of vision disappeared. In most cases, these symptoms last only a few minutes. Rarely, ischemia or infarction of the ophthalmic artery or central retinal arteries occurs at the time of cerebral TIA or infarction.

A high-pitched prolonged carotid bruit fading into diastole is often associated with tightly stenotic lesions. As the stenosis grows tighter and flow distal to the stenosis becomes reduced, the bruit becomes fainter and may disappear when occlusion is imminent.

COMMON CAROTID ARTERY All symptoms and signs of internal carotid occlusion may also be present with occlusion of the common carotid artery. Jaw claudication may result from low flow in the external carotid branches. Bilateral common carotid artery occlusions at their origin may occur in Takayasu's arteritis ([Chap. 385](#)).

Stroke Within the Posterior Circulation The posterior circulation is composed of the paired vertebral arteries, the basilar artery, and the paired posterior cerebral arteries. The vertebral arteries join to form the basilar artery at the pontomedullary junction. The basilar artery divides into two posterior cerebral arteries in the interpeduncular fossa (Figs. 446-4, 446-8, and 446-9). These major arteries give rise to long and short circumferential branches and to smaller deep penetrating branches that supply the cerebellum, medulla, pons, midbrain, subthalamus, thalamus, hippocampus, and medial temporal and occipital lobes. Occlusion of each vessel produces its own distinctive syndrome.

POSTERIOR CEREBRAL ARTERY In 75% of cases, both PCAs arise from the bifurcation of the basilar artery; in 20%, one has its origin from the ipsilateral internal carotid artery via the posterior communicating artery; in 5%, both originate from the respective ipsilateral internal carotid arteries (Figs. 446-8 and 446-9). The precommunal, or P1, segment of the true posterior cerebral artery is atretic in such cases.

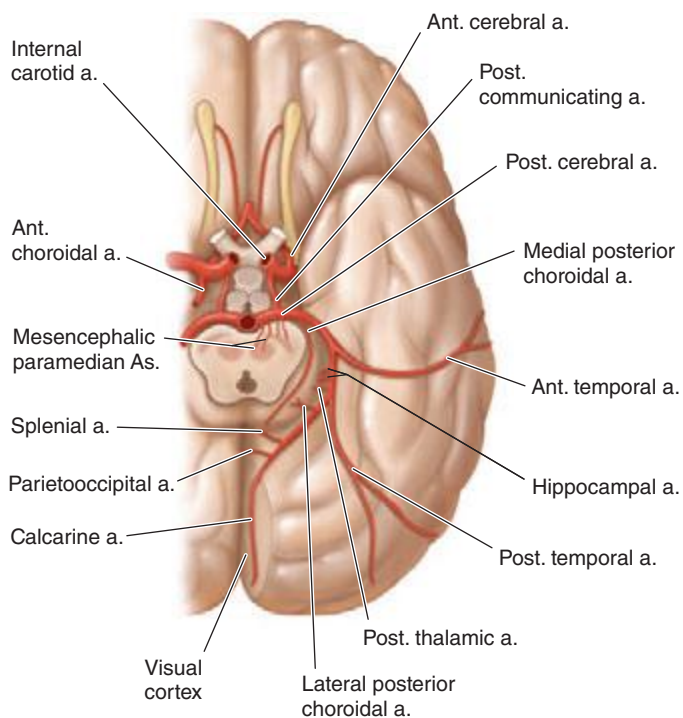


FIGURE 446-9 Inferior aspect of the brain with the branches and distribution of the posterior cerebral artery and the principal anatomic structures shown.

Signs and symptoms: *Structures involved*

Peripheral territory (see also Fig. 446-12). Homonymous hemianopia (often upper quadrantic): *Calcarine cortex or optic radiation nearby*. Bilateral homonymous hemianopia, cortical blindness, awareness or denial of blindness; tactile naming, achromatopia (color blindness), failure to see to-and-fro movements, inability to perceive objects not centrally located, apraxia of ocular movements, inability to count or enumerate objects, tendency to run into things that the patient sees and tries to avoid: *Bilateral occipital lobe with possibly the parietal lobe involved*. Verbal dyslexia without agraphia, color anomia: *Dominant calcarine lesion and posterior part of corpus callosum*. Memory defect: *Hippocampal lesion bilaterally or on the dominant side only*. Topographic disorientation and prosopagnosia: *Usually with lesions of nondominant, calcarine, and lingual gyrus*. Simultanagnosia, hemivisual neglect: *Dominant visual cortex, contralateral hemisphere*. Unformed visual hallucinations, peduncular hallucinosis, metamorphopsia, teleopsia, illusory visual spread, palinopsia, distortion of outlines, central photophobia: *Calcarine cortex*. Complex hallucinations: *Usually nondominant hemisphere*.

Central territory. Thalamic syndrome: sensory loss (all modalities), spontaneous pain and dysesthesias, choreoathetosis, intention tremor, spasms of hand, mild hemiparesis: *Posteroventral nucleus of thalamus; involvement of the adjacent subthalamus body or its afferent tracts*. Thalamoperforate syndrome: crossed cerebellar ataxia with ipsilateral third nerve palsy (Claude's syndrome): *Dentothalamic tract and issuing third nerve*. Weber's syndrome: third nerve palsy and contralateral hemiplegia: *Third nerve and cerebral peduncle*. Contralateral hemiplegia: *Cerebral peduncle*. Paralysis or paresis of vertical eye movement, skew deviation, sluggish pupillary responses to light, slight miosis and ptosis (retraction nystagmus and "tucking" of the eyelids may be associated): *Supranuclear fibers to third nerve, interstitial nucleus of Cajal, nucleus of Darkschewitsch, and posterior commissure*. Contralateral rhythmic, ataxic action tremor; rhythmic postural or "holding" tremor (rubral tremor): *Dentothalamic tract*.

PCA syndromes usually result from atheroma formation or emboli that lodge at the top of the basilar artery; posterior circulation disease may also be caused by dissection of either vertebral artery or fibromuscular dysplasia.

Two clinical syndromes are commonly observed with occlusion of the PCA: (1) *P1 syndrome*: midbrain, subthalamic, and thalamic signs, which are due to disease of the proximal P1 segment of the PCA or its penetrating branches (thalamogeniculate, Percheron, and posterior choroidal arteries); and (2) *P2 syndrome*: cortical temporal and occipital lobe signs, due to occlusion of the P2 segment distal to the junction of the PCA with the posterior communicating artery.

P1 SYNDROMES Infarction usually occurs in the ipsilateral subthalamus and medial thalamus and in the ipsilateral cerebral peduncle and midbrain (Figs. 446-9 and 446-14). A third nerve palsy with contralateral ataxia (Claude's syndrome) or with contralateral hemiplegia (Weber's syndrome) may result. The ataxia indicates involvement of the red nucleus or dentatorubrothalamic tract; the hemiplegia is localized to the cerebral peduncle (Fig. 446-14). If the subthalamic nucleus is involved, contralateral hemiballismus may occur. Occlusion of the artery of Percheron produces paresis of upward gaze and drowsiness and often abulia. Extensive infarction in the midbrain and subthalamus occurring with bilateral proximal PCA occlusion presents as coma, unreactive pupils, bilateral pyramidal signs, and decerebrate rigidity.

Occlusion of the penetrating branches of thalamic and thalamogeniculate arteries produces less extensive thalamic and thalamocapsular lacunar syndromes. The *thalamic Déjérine-Roussy syndrome* consists of contralateral hemisensory loss followed later by an agonizing, searing or burning pain in the affected areas. It is persistent and responds poorly to analgesics. Anticonvulsants (carbamazepine or gabapentin) or tricyclic antidepressants may be beneficial.

P2 SYNDROMES (Figs. 446-8 and 446-9) Occlusion of the distal PCA causes infarction of the medial temporal and occipital lobes. Contralateral homonymous hemianopia with macula sparing is the usual manifestation. Occasionally, only the upper quadrant of visual field is involved. If the visual association areas are spared and only the calcarine cortex is involved, the patient may be aware of visual defects. Medial temporal lobe and hippocampal involvement may cause an acute disturbance in memory, particularly if it occurs in the dominant hemisphere. The defect usually clears because memory has bilateral representation. If the dominant hemisphere is affected and the infarct extends to involve the splenium of the corpus callosum, the patient may demonstrate alexia without agraphia. Visual agnosia for faces, objects, mathematical symbols, and colors and anomia with paraphasic errors (amnestic aphasia) may also occur, even without callosal involvement. Occlusion of the posterior cerebral artery can produce *peduncular hallucinosis* (visual hallucinations of brightly colored scenes and objects).

Bilateral infarction in the distal PCAs produces cortical blindness (blindness with preserved pupillary light reaction). The patient is often unaware of the blindness or may even deny it (*Anton's syndrome*). Tiny islands of vision may persist, and the patient may report that vision fluctuates as images are captured in the preserved portions. Rarely, only peripheral vision is lost and central vision is spared, resulting in "gun-barrel" vision. Bilateral visual association area lesions may result in *Balint's syndrome*, a disorder of the orderly visual scanning of the environment (**Chap. 36**), usually resulting from infarctions secondary to low flow in the "watershed" between the distal PCA and MCA territories, as occurs after cardiac arrest. Patients may experience persistence of a visual image for several minutes despite gazing at another scene (*palinopsia*) or an inability to synthesize the whole of an image (*asimultanagnosia*). Embolic occlusion of the top of the basilar artery can produce any or all of the central or peripheral territory symptoms. The hallmark is the sudden onset of bilateral signs, including ptosis, pupillary asymmetry or lack of reaction to light, and somnolence.

VERTEBRAL AND POSTERIOR INFERIOR CEREBELLAR ARTERIES The vertebral artery, which arises from the innominate artery on the right and the subclavian artery on the left, consists of four segments. The first (V1) extends from its origin to its entrance into the sixth or fifth transverse vertebral foramen. The second segment (V2) traverses the vertebral

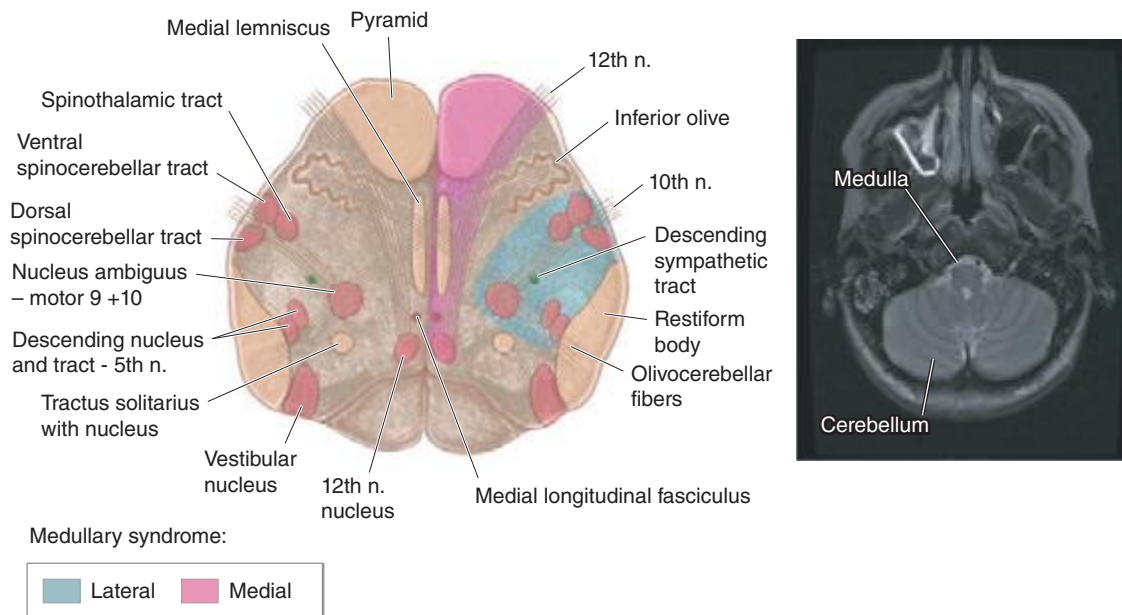


FIGURE 446-10 Axial section at the level of the medulla, depicted schematically on the left, with a corresponding magnetic resonance image on the right. Note that in Figs. 446-10 through 446-14, all drawings are oriented with the dorsal surface at the bottom, matching the orientation of the brainstem that is commonly seen in all modern neuroimaging studies. Approximate regions involved in medial and lateral medullary stroke syndromes are shown.

Signs and symptoms: *Structures involved*

1. Medial medullary syndrome (occlusion of vertebral artery or of branch of vertebral or lower basilar artery)
 - On side of lesion
 - Paralysis with atrophy of one-half half the tongue: *Ipsilateral twelfth nerve*
 - On side opposite lesion
 - Paralysis of arm and leg, sparing face; impaired tactile and proprioceptive sense over one-half the body: *Contralateral pyramidal tract and medial lemniscus*
2. Lateral medullary syndrome (occlusion of any of five vessels may be responsible—vertebral, posterior inferior cerebellar, superior, middle, or inferior lateral medullary arteries)
 - On side of lesion
 - Pain, numbness, impaired sensation over one-half the face: *Descending tract and nucleus fifth nerve*
 - Ataxia of limbs, falling to side of lesion: *Uncertain—restiform body, cerebellar hemisphere, cerebellar fibers, spinocerebellar tract (?)*
 - Nystagmus, diplopia, oscillopsia, vertigo, nausea, vomiting: *Vestibular nucleus*
 - Horner's syndrome (miosis, ptosis, decreased sweating): *Descending sympathetic tract*
 - Dysphagia, hoarseness, paralysis of palate, paralysis of vocal cord, diminished gag reflex: *Issuing fibers ninth and tenth nerves*
 - Loss of taste: *Nucleus and tractus solitarius*
 - Numbness of ipsilateral arm, trunk, or leg: *Cuneate and gracile nuclei*
 - Weakness of lower face: *Geniculate upper motor neuron fibers to ipsilateral facial nucleus*
 - On side opposite lesion
 - Impaired pain and thermal sense over half the body, sometimes face: *Spinothalamic tract*
3. Total unilateral medullary syndrome (occlusion of vertebral artery): Combination of medial and lateral syndromes
4. Lateral pontomedullary syndrome (occlusion of vertebral artery): Combination of lateral medullary and lateral inferior pontine syndrome
5. Basilar artery syndrome (the syndrome of the lone vertebral artery is equivalent): A combination of the various brainstem syndromes plus those arising in the posterior cerebral artery distribution.
 - Bilateral long tract signs (sensory and motor; cerebellar and peripheral cranial nerve abnormalities): *Bilateral long tract; cerebellar and peripheral cranial nerves*
 - Paralysis or weakness of all extremities, plus all bulbar musculature: *Corticobulbar and corticospinal tracts bilaterally*

foramina from C6 to C2. The third (V3) passes through the transverse foramen and circles around the arch of the atlas to pierce the dura at the foramen magnum. The fourth (V4) segment courses upward to join the other vertebral artery to form the basilar artery; only the fourth segment gives rise to branches that supply the brainstem and cerebellum. The posterior inferior cerebellar artery (PICA) in its proximal segment supplies the lateral medulla and, in its distal branches, the inferior surface of the cerebellum.

Atherothrombotic lesions have a predilection for V1 and V4 segments of the vertebral artery. The first segment may become diseased at the origin of the vessel and may produce posterior circulation emboli; collateral flow from the contralateral vertebral artery or the ascending cervical, thyrocervical, or occipital arteries is usually sufficient to prevent low-flow TIAs or stroke. When one vertebral artery is atretic and

an atherothrombotic lesion threatens the origin of the other, the collateral circulation, which may also include retrograde flow down the basilar artery, is often insufficient (Figs. 446-4 and 446-9). In this setting, low-flow TIAs may occur, consisting of syncope, vertigo, and alternating hemiplegia; this state also sets the stage for thrombosis. Disease of the distal fourth segment of the vertebral artery can promote thrombus formation manifest as embolism or with propagation as basilar artery thrombosis. Stenosis proximal to the origin of the PICA can threaten the lateral medulla and posterior inferior surface of the cerebellum.

If the subclavian artery is occluded proximal to the origin of the vertebral artery, there is a reversal in the direction of blood flow in the ipsilateral vertebral artery. Exercise of the ipsilateral arm may increase demand on vertebral flow, producing posterior circulation TIAs, or “subclavian steal.”

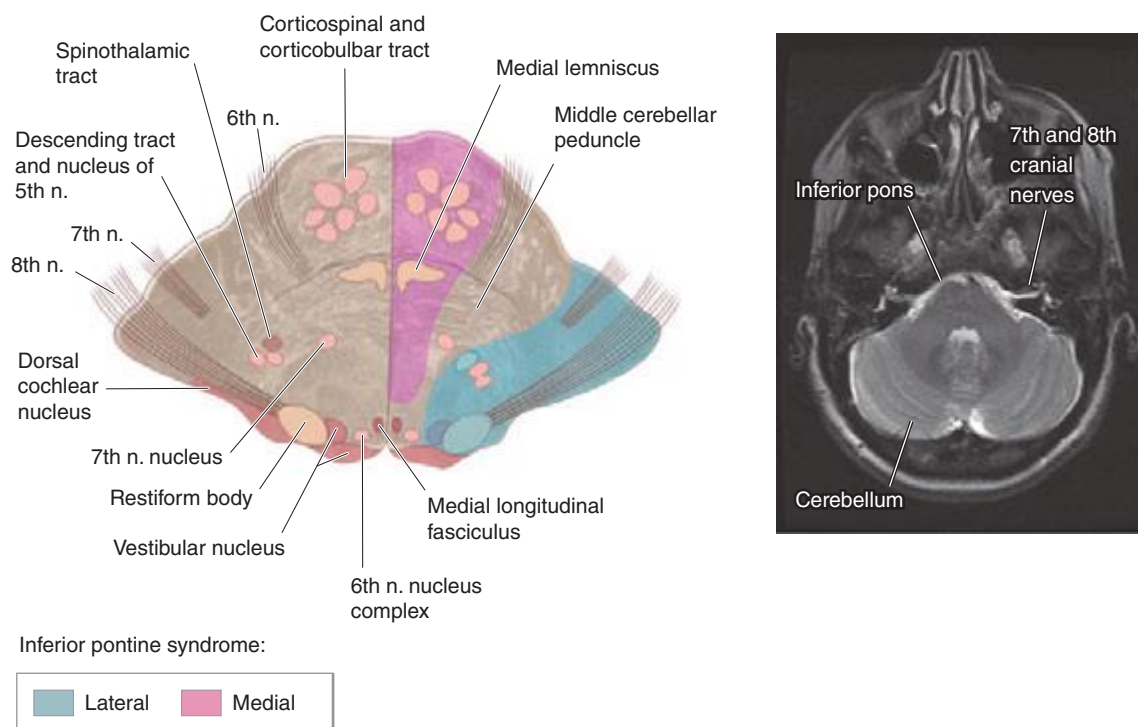


FIGURE 446-11 Axial section at the level of the inferior pons, depicted schematically on the left, with a corresponding magnetic resonance image on the right. Approximate regions involved in medial and lateral inferior pontine stroke syndromes are shown.

Signs and symptoms: Structures involved

1. Medial inferior pontine syndrome (occlusion of paramedian branch of basilar artery)
 - On side of lesion
 - Paralysis of conjugate gaze to side of lesion (preservation of convergence): *Center for conjugate lateral gaze*
 - Nystagmus: *Vestibular nucleus*
 - Ataxia of limbs and gait: Likely *middle cerebellar peduncle*
 - Diplopia on lateral gaze: *Abducens nerve*
 - On side opposite lesion
 - Paralysis of face, arm, and leg: *Corticobulbar and corticospinal tract in lower pons*
 - Impaired tactile and proprioceptive sense over one-half of the body: *Medial lemniscus*
2. Lateral inferior pontine syndrome (occlusion of anterior inferior cerebellar artery)
 - On side of lesion
 - Horizontal and vertical nystagmus, vertigo, nausea, vomiting, oscillopsia: *Vestibular nerve or nucleus*
 - Facial paralysis: *Seventh nerve*
 - Paralysis of conjugate gaze to side of lesion: *Center for conjugate lateral gaze*
 - Deafness, tinnitus: *Auditory nerve or cochlear nucleus*
 - Ataxia: *Middle cerebellar peduncle and cerebellar hemisphere*
 - Impaired sensation over face: *Descending tract and nucleus fifth nerve*
 - On side opposite lesion
 - Impaired pain and thermal sense over one-half the body (may include face): *Spinothalamic tract*

Although atheromatous disease rarely narrows the second and third segments of the vertebral artery, this region is subject to dissection, fibromuscular dysplasia, and, rarely, encroachment by osteophytic spurs within the vertebral foramina.

Embolic occlusion or thrombosis of a V4 segment causes ischemia of the lateral medulla. The constellation of vertigo, numbness of the ipsilateral face and contralateral limbs, diplopia, hoarseness, dysarthria, dysphagia, and ipsilateral Horner's syndrome is called the *lateral medullary (or Wallenberg's) syndrome* (Fig. 446-10). Most cases result from ipsilateral vertebral artery occlusion; in the remainder, PICA occlusion is responsible. Occlusion of the medullary penetrating branches of the vertebral artery or PICA results in partial syndromes. *Hemiparesis is not a feature of vertebral artery occlusion; however, quadriplegia may result from occlusion of the anterior spinal artery.*

Rarely, a *medial medullary syndrome* occurs with infarction of the pyramid and contralateral hemiparesis of the arm and leg, sparing the face. If the medial lemniscus and emerging hypoglossal nerve fibers

are involved, contralateral loss of joint position sense and ipsilateral tongue weakness occur.

Cerebellar infarction with edema can lead to *sudden respiratory arrest* due to raised ICP in the posterior fossa. Drowsiness, Babinski signs, dysarthria, and bifacial weakness may be absent, or present only briefly, before respiratory arrest ensues. Gait unsteadiness, headache, dizziness, nausea, and vomiting may be the only early symptoms and signs and should arouse suspicion of this impending complication, which may require neurosurgical decompression, often with an excellent outcome. Separating these symptoms from those of viral labyrinthitis can be a challenge, but headache, neck stiffness, and unilateral dysmetria favor stroke.

BASILAR ARTERY Branches of the basilar artery supply the base of the pons and superior cerebellum and fall into three groups: (1) paramedian, 7–10 in number, which supply a wedge of pons on either side of the midline; (2) short circumferential, 5–7 in number, that supply

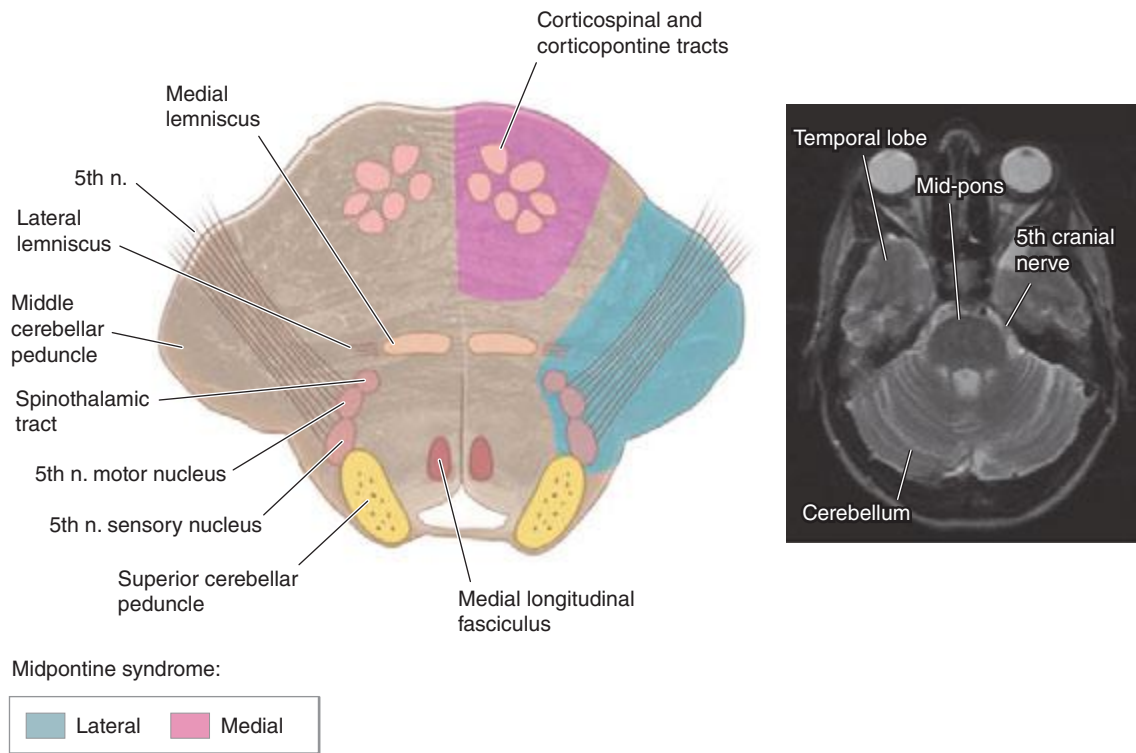


FIGURE 446-12 Axial section at the level of the midpons, depicted schematically on the left, with a corresponding magnetic resonance image on the right. Approximate regions involved in medial and lateral midpontine stroke syndromes are shown.

Signs and symptoms: *Structures involved*

1. Medial midpontine syndrome (paramedian branch of midbasilar artery)
 - On side of lesion
 - Ataxia of limbs and gait (more prominent in bilateral involvement): *Pontine nuclei*
 - On side opposite lesion
 - Paralysis of face, arm, and leg: *Corticobulbar and corticospinal tract*
 - Variable impaired touch and proprioception when lesion extends posteriorly: *Medial lemniscus*
2. Lateral midpontine syndrome (short circumferential artery)
 - On side of lesion
 - Ataxia of limbs: *Middle cerebellar peduncle*
 - Paralysis of muscles of mastication: *Motor fibers or nucleus of fifth nerve*
 - Impaired sensation over side of face: *Sensory fibers or nucleus of fifth nerve*
 - On side opposite lesion
 - Impaired pain and thermal sense on limbs and trunk: *Spinothalamic tract*

the lateral two-thirds of the pons and middle and superior cerebellar peduncles; and (3) bilateral long circumferential (superior cerebellar and anterior inferior cerebellar arteries), which course around the pons to supply the cerebellar hemispheres.

Atheromatous lesions can occur anywhere along the basilar trunk but are most frequent in the proximal basilar and distal vertebral segments. Typically, lesions occlude either the proximal basilar and one or both vertebral arteries. The clinical picture varies depending on the availability of retrograde collateral flow from the posterior communicating arteries. Rarely, dissection of a vertebral artery may involve the basilar artery and, depending on the location of true and false lumen, may produce multiple penetrating artery strokes.

Although atherothrombosis occasionally occludes the distal portion of the basilar artery, emboli from the heart or proximal vertebral or basilar segments are more commonly responsible for “top of the basilar” syndromes.

Because the brainstem contains many structures in close apposition, a diversity of clinical syndromes may emerge with ischemia, reflecting involvement of the corticospinal and corticobulbar tracts, ascending sensory tracts, and cranial nerve nuclei (Figs. 446-11, 446-12, 446-13, and 446-14).

The symptoms of transient ischemia or infarction in the territory of the basilar artery often do not indicate whether the basilar artery itself or one of its branches is diseased, yet this distinction has important

implications for therapy. *The picture of complete basilar occlusion, however, is easy to recognize as a constellation of bilateral long tract signs (sensory and motor) with signs of cranial nerve and cerebellar dysfunction.* A “locked-in” state of preserved consciousness with quadriplegia and cranial nerve signs suggests complete pontine and lower midbrain infarction. The therapeutic goal is to identify *impending* basilar occlusion before devastating infarction occurs. A series of TIAs and a slowly progressive, fluctuating stroke are extremely significant, because they often herald an atherothrombotic occlusion of the distal vertebral or proximal basilar artery.

TIAs in the proximal basilar distribution may produce vertigo (often described by patients as “swimming,” “swaying,” “moving,” “unsteadiness,” or “light-headedness”). Other symptoms that warn of basilar thrombosis include diplopia, dysarthria, facial or circumoral numbness, and hemisensory symptoms. In general, symptoms of basilar branch TIAs affect one side of the brainstem, whereas symptoms of basilar artery TIAs usually affect both sides, although a “herald” hemiparesis has been emphasized as an initial symptom of basilar occlusion. Most often, TIAs, whether due to impending occlusion of the basilar artery or a basilar branch, are short lived (5–30 min) and repetitive, occurring several times a day. The pattern suggests intermittent reduction of flow. Many neurologists treat with heparin to prevent clot propagation.

Atherothrombotic occlusion of the basilar artery with infarction usually causes *bilateral* brainstem signs. A gaze paresis or internuclear

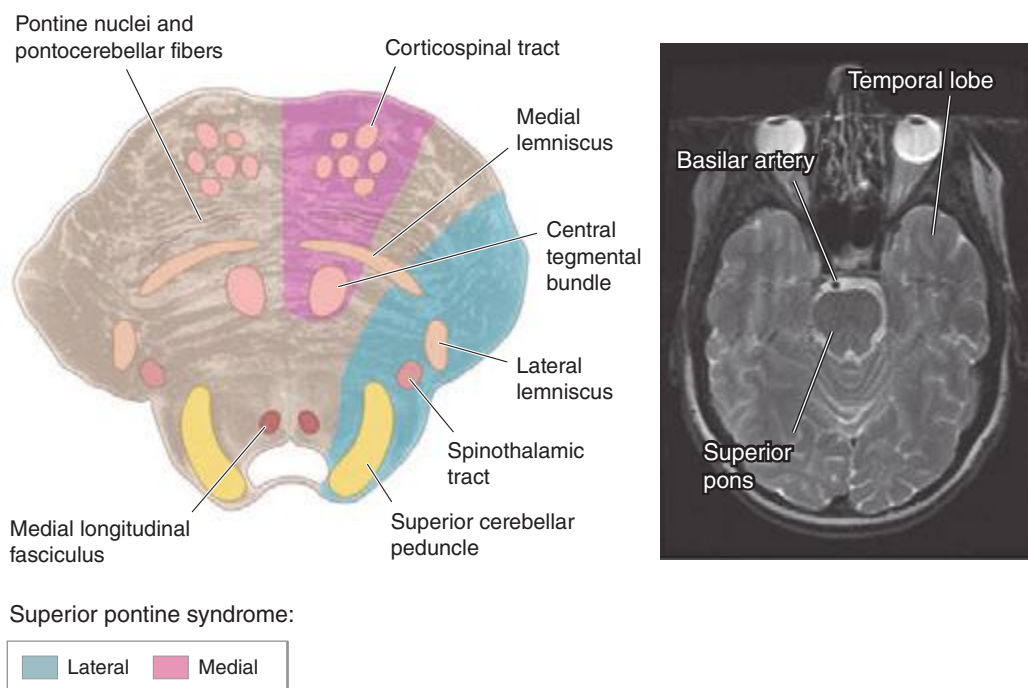


FIGURE 446-13 Axial section at the level of the superior pons, depicted schematically on the left, with a corresponding magnetic resonance image on the right. Approximate regions involved in medial and lateral superior pontine stroke syndromes are shown.

Signs and symptoms: *Structures involved*

1. Medial superior pontine syndrome (paramedian branches of upper basilar artery)

On side of lesion

Cerebellar ataxia (probably): *Superior and/or middle cerebellar peduncle*

Internuclear ophthalmoplegia: *Medial longitudinal fasciculus*

Myoclonic syndrome, palate, pharynx, vocal cords, respiratory apparatus, face, oculomotor apparatus, etc.: *Localization uncertain—central tegmental bundle, dentate projection, inferior olivary nucleus*

On side opposite lesion

Paralysis of face, arm, and leg: *Corticobulbar and corticospinal tract*

Rarely touch, vibration, and position are affected: *Medial lemniscus*

2. Lateral superior pontine syndrome (syndrome of superior cerebellar artery)

On side of lesion

Ataxia of limbs and gait, falling to side of lesion: *Middle and superior cerebellar peduncles, superior surface of cerebellum, dentate nucleus*

Dizziness, nausea, vomiting; horizontal nystagmus: *Vestibular nucleus*

Paresis of conjugate gaze (ipsilateral): *Pontine contralateral gaze*

Skew deviation: *Uncertain*

Miosis, ptosis, decreased sweating over face (Horner's syndrome): *Descending sympathetic fibers*

Tremor: *Localization unclear—Dentate nucleus, superior cerebellar peduncle*

On side opposite lesion

Impaired pain and thermal sense on face, limbs, and trunk: *Spinothalamic tract*

Impaired touch, vibration, and position sense, more in leg than arm (there is a tendency to incongruity of pain and touch deficits): *Medial lemniscus (lateral portion)*

ophthalmoplegia associated with ipsilateral hemiparesis may be the only manifestation of bilateral brainstem ischemia. More often, unequivocal signs of bilateral pontine disease are present. Complete basilar thrombosis carries a high mortality.

Occlusion of a branch of the basilar artery usually causes *unilateral* symptoms and signs involving motor, sensory, and cranial nerves. As long as symptoms remain unilateral, concern over pending basilar occlusion should be reduced.

Occlusion of the superior cerebellar artery results in severe ipsilateral cerebellar ataxia, nausea and vomiting, dysarthria, and contralateral loss of pain and temperature sensation over the extremities, body, and face (spino- and trigeminothalamic tract). Partial deafness, ataxic tremor of the ipsilateral upper extremity, Horner's syndrome, and palatal myoclonus may occur rarely. Partial syndromes occur frequently (Fig. 446-13). With large strokes, swelling and mass effects may compress the midbrain or produce hydrocephalus; these symptoms may evolve rapidly. Neurosurgical intervention may be lifesaving in such cases.

Occlusion of the anterior inferior cerebellar artery produces variable degrees of infarction because the size of this artery and the territory it supplies vary inversely with those of the PICA. The principal symptoms include: (1) ipsilateral deafness, facial weakness, vertigo, nausea and vomiting, nystagmus, tinnitus, cerebellar ataxia, Horner's syndrome, and paresis of conjugate lateral gaze; and (2) contralateral loss of pain and temperature sensation. An occlusion close to the origin of the artery may cause corticospinal tract signs (Fig. 446-11).

Occlusion of one of the short circumferential branches of the basilar artery affects the lateral two-thirds of the pons and middle or superior cerebellar peduncle, whereas occlusion of one of the paramedian branches affects a wedge-shaped area on either side of the medial pons (Figs. 446-11 through 446-13).

IMAGING STUDIES

See also Chap. 440e.

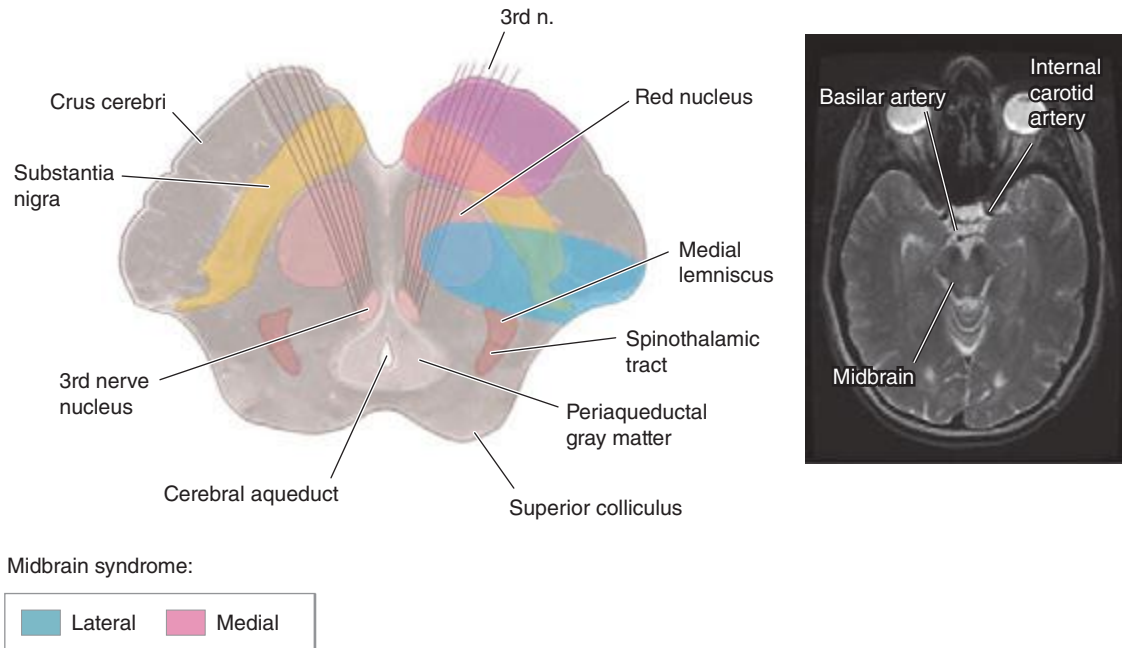


FIGURE 446-14 Axial section at the level of the midbrain, depicted schematically on the left, with a corresponding magnetic resonance image on the right. Approximate regions involved in medial and lateral midbrain stroke syndromes are shown.

Signs and symptoms: *Structures involved*

1. Medial midbrain syndrome (paramedian branches of upper basilar and proximal posterior cerebral arteries)
 - On side of lesion
 - Eye “down and out” secondary to unopposed action of fourth and sixth cranial nerves, with dilated and unresponsive pupil: *Third nerve fibers*
 - On side opposite lesion
 - Paralysis of face, arm, and leg: *Corticobulbar and corticospinal tract descending in crus cerebri*
2. Lateral midbrain syndrome (syndrome of small penetrating arteries arising from posterior cerebral artery)
 - On side of lesion
 - Eye “down and out” secondary to unopposed action of fourth and sixth cranial nerves, with dilated and unresponsive pupil: *Third nerve fibers and/or third nerve nucleus*
 - On side opposite lesion
 - Hemiataxia, hyperkinesias, tremor: *Red nucleus, dentatorubrothalamic pathway*

CT Scans CT radiographic images identify or exclude hemorrhage as the cause of stroke, and they identify extraparenchymal hemorrhages, neoplasms, abscesses, and other conditions masquerading as stroke. Brain CT scans obtained in the first several hours after an infarction generally show no abnormality, and the infarct may not be seen reliably for 24–48 h. CT may fail to show small ischemic strokes in the posterior fossa because of bone artifact; small infarcts on the cortical surface may also be missed.

Contrast-enhanced CT scans add specificity by showing contrast enhancement of subacute infarcts and allow visualization of venous structures. Coupled with multidetector scanners, CT angiography (CTA) can be performed with administration of IV iodinated contrast allowing visualization of the cervical and intracranial arteries, intracranial veins, aortic arch, and even the coronary arteries in one imaging session. Carotid disease and intracranial vascular occlusions are readily identified with this method (Fig. 446-3). After an IV bolus of contrast, deficits in brain perfusion produced by vascular occlusion can also be demonstrated (Fig. 446-15) and used to predict the region of infarcted brain and the brain at risk of further infarction (i.e., the ischemic penumbra, see “Pathophysiology of Ischemic Stroke” above). CT imaging is also sensitive for detecting SAH (although by itself does not rule it out), and CTA can readily identify intracranial aneurysms (Chap. 330). Because of its speed and wide availability, noncontrast head CT is the imaging modality of choice in patients with acute stroke (Fig. 446-1), and CTA and CT perfusion imaging may also be useful and convenient adjuncts.

MRI

MRI reliably documents the extent and location of infarction in all areas of the brain, including the posterior fossa and cortical surface.

It also identifies intracranial hemorrhage and other abnormalities and, using special sequences, can be as sensitive as CT for detecting acute intracerebral hemorrhage. MRI scanners with magnets of higher field strength produce more reliable and precise images. Diffusion-weighted imaging is more sensitive for early brain infarction than standard MR sequences or CT (Fig. 446-16), as is fluid-attenuated inversion recovery (FLAIR) imaging (Chap. 440e). Using IV administration of gadolinium contrast, MR perfusion studies can be performed. Brain regions showing poor perfusion but no abnormality on diffusion provide, compared to CT, an equivalent measure of the ischemic penumbra. MR angiography is highly sensitive for stenosis of extracranial internal carotid arteries and of large intracranial vessels. With higher degrees of stenosis, MR angiography tends to overestimate the degree of stenosis when compared to conventional x-ray angiography. MRI with fat saturation is an imaging sequence used to visualize extra or intracranial arterial dissection. This sensitive technique images clotted blood within the dissected vessel wall. Iron-sensitive imaging (ISI) is helpful to detect cerebral microbleeds that may be present in cerebral amyloid angiopathy and other hemorrhagic disorders.

MRI is more expensive and time consuming than CT and less readily available. Claustrophobia and the logistics of imaging acutely ill patients also limit its application. Most acute stroke protocols use CT because of these limitations. However, MRI is useful outside the acute period by more clearly defining the extent of tissue injury and discriminating new from old regions of brain infarction. MRI may have particular utility in patients with TIA, because it is also more likely to identify new infarction, which is a strong predictor of subsequent stroke.

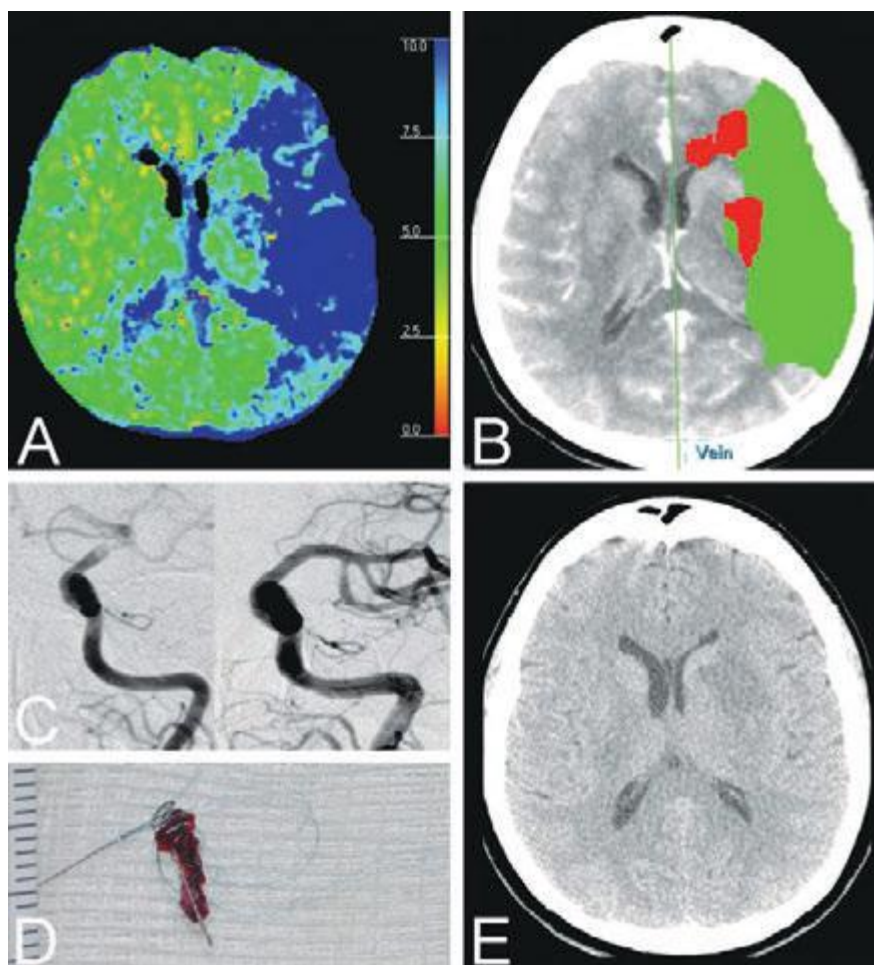


FIGURE 446-15 Acute left middle cerebral artery (MCA) stroke with right hemiplegia but preserved language. **A.** Computed tomography (CT) perfusion mean-transit time map showing delayed perfusion of the left MCA distribution (*blue*). **B.** Predicted region of infarct (*red*) and penumbra (*green*) based on CT perfusion data. **C.** Conventional angiogram showing occlusion of the left internal carotid–MCA bifurcation (*left panel*), and revascularization of the vessels following successful thrombectomy 8 h after stroke symptom onset (*right panel*). **D.** The clot removed with a thrombectomy device (L5, Concentric Medical, Inc.). **E.** CT scan of the brain 2 days later; note infarction in the region predicted in **B** but preservation of the penumbral region by successful revascularization.

Cerebral Angiography Conventional x-ray cerebral angiography is the gold standard for identifying and quantifying atherosclerotic stenoses of the cerebral arteries and for identifying and characterizing other pathologies, including aneurysms, vasospasm, intraluminal thrombi, fibromuscular dysplasia, arteriovenous fistulae, vasculitis, and collateral channels of blood flow. Conventional angiography carries risks of arterial damage, groin hemorrhage, embolic stroke, and renal failure from contrast nephropathy, so it should be reserved for situations where less invasive means are inadequate. As reviewed earlier in this chapter, endovascular stroke therapy has not been proven effective in three randomized trials, and this remains an area of ongoing investigation.

Ultrasound Techniques Stenosis at the origin of the internal carotid artery can be identified and quantified reliably by ultrasonography that combines a B-mode ultrasound image with a Doppler ultrasound assessment of flow velocity (“duplex” ultrasound). Transcranial Doppler (TCD) assessment of MCA, ACA, and PCA flow and of vertebralbasilar flow is also useful. This latter technique can detect stenotic lesions in the large intracranial arteries because such lesions increase systolic flow velocity. TCD can assist thrombolysis and improve large artery recanalization following rtPA administration; the potential clinical benefit of this treatment is the subject of ongoing study. TCD can also detect microemboli from otherwise asymptomatic carotid plaques. In many cases, MR angiography combined with carotid and transcranial ultrasound studies eliminates the need for conventional

x-ray angiography in evaluating vascular stenosis. Alternatively, CT angiography of the entire head and neck can be performed during the initial imaging of acute stroke. Because this images the entire arterial system relevant to stroke, with the exception of the heart, much of the clinician’s stroke workup can be completed with this single imaging study.

Perfusion Techniques Both xenon techniques (principally xenon-CT) and positron emission tomography (PET) can quantify cerebral blood flow. These tools are generally used for research (**Chap. 440e**) but can be useful for determining the significance of arterial stenosis and planning for revascularization surgery. Single-photon emission computed tomography (SPECT) and MR perfusion techniques report relative cerebral blood flow. As noted above, CT imaging is used as the initial imaging modality for acute stroke, and some centers combine both CT angiography and CT perfusion imaging together with the noncontrast CT scan. CT perfusion imaging increases the sensitivity for detecting ischemia and can measure the ischemic penumbra (Fig. 446-15). Alternatively, MR perfusion can be combined with MR diffusion imaging to identify the ischemic penumbra as the mismatch between these two imaging sequences (Fig. 446-16).

INTRACRANIAL HEMORRHAGE

Hemorrhages are classified by their location and the underlying vascular pathology. Bleeding into subdural and epidural spaces is

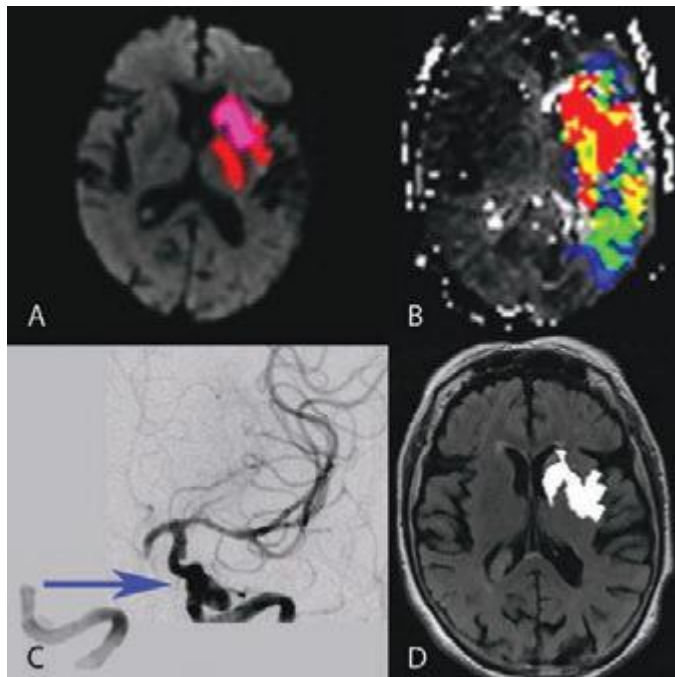


FIGURE 446-16 Magnetic resonance imaging (MRI) of acute stroke. **A.** MRI diffusion-weighted image (DWI) of an 82-year-old woman 2.5 h after onset of right-sided weakness and aphasia reveals restricted diffusion within the left basal ganglia and internal capsule (colored regions). **B.** Perfusion defect within the left hemisphere (colored signal) imaged after administration of an IV bolus of gadolinium contrast. The discrepancy between the region of poor perfusion shown in **B** and the diffusion deficit shown in **A** is called *diffusion-perfusion mismatch* and provides an estimate of the ischemic penumbra. Without specific therapy, the region of infarction will expand into much or all of the perfusion deficit. **C.** Cerebral angiogram of the left internal carotid artery in this patient before (left) and after (right) successful endovascular embolectomy. The occlusion is within the carotid terminus. **D.** Fluid-attenuated inversion recovery image obtained 3 days later showing a region of infarction (coded as white) that corresponds to the initial DWI image in **A**, but not the entire area at risk shown in **B**, suggesting that successful embolectomy saved a large region of brain tissue from infarction. (Courtesy of Gregory Albers, MD, Stanford University; with permission.)

principally produced by trauma. SAH results from trauma or the rupture of an intracranial aneurysm or arteriovenous malformation (AVM) (Chap. 330). Intracerebral and intraventricular hemorrhage will be considered here.

DIAGNOSIS

Intracranial hemorrhage is often discovered on noncontrast CT imaging of the brain during the acute evaluation of stroke. Because CT is more widely available and may be logistically easier, CT imaging is the preferred method for acute stroke evaluation (Fig. 446-1). The location of the hemorrhage narrows the differential diagnosis to a few entities. Table 446-6 lists the causes and anatomic spaces involved in hemorrhages.

EMERGENCY MANAGEMENT

Close attention should be paid to airway management because a reduction in the level of consciousness is common and often progressive. The initial blood pressure should be maintained until the results of the CT scan are reviewed and demonstrate an intracerebral hemorrhage (ICH). In theory, a higher blood pressure should promote hematoma expansion, but it remains unclear if lowering of blood pressure reduces hematoma growth. Recent clinical trials have shown that systolic blood pressure (SBP) can be safely lowered acutely and rapidly to <140 mmHg in patients with spontaneous ICH whose initial SBP was 150–220 mmHg.

The INTERACT2 trial is the only large phase 3 clinical trial to address the effect of acute blood pressure lowering on ICH functional outcome. INTERACT2 randomized patients with spontaneous ICH within 6 h of onset and a baseline SBP of 150–220 mmHg to two different SBP targets (<140 mmHg and <180 mmHg). In those with the target SBP <140 mmHg, 52% had an outcome of death or major disability at 90 days compared with 55.6% of those with a target SBP <180 mmHg ($p = .06$). There was a significant shift to improved outcomes in the lower blood pressure arm, whereas both groups had a similar mortality. This study shows that it is not harmful, and may be modestly beneficial, to lower blood pressure in acute ICH. Thus, it is reasonable to target an SBP <140 mmHg initially in this group of patients. In patients who have higher SBP on presentation or who are deeply comatose with possible elevated ICP, it is unclear whether the INTERACT2 results apply. In patients who have ICP monitors in place, current recommendations are to maintain the cerebral perfusion pressure (mean arterial pressure [MAP] minus ICP) above 60 mmHg. Blood pressure should be lowered with nonvasodilating IV drugs such as nicardipine, labetalol, or esmolol. Patients with cerebellar hemorrhages or with depressed mental status and radiographic evidence of hydrocephalus should undergo urgent neurosurgical evaluation; these patients require close monitoring because they can deteriorate rapidly. Based on the clinical examination and CT findings, further imaging studies may be necessary, including MRI or conventional x-ray angiography. Stuporous or comatose patients with clinical and imaging signs of herniation are generally treated presumptively for elevated ICP, with tracheal intubation, administration of osmotic diuretics such as mannitol or hypertonic saline, and elevation of the head of the bed while surgical consultation is obtained (Chap. 330). Reversal of coagulopathy and consideration of surgical evacuation of the hematoma (detailed below) are two other principal aspects of initial emergency management.

INTRACEREBRAL HEMORRHAGE

ICH accounts for ~10% of all strokes, and about 35–45% of patients die within the first month. Incidence rates are particularly high in Asians and blacks. Hypertension, coagulopathy, sympathomimetic drugs (cocaine, methamphetamine), and cerebral amyloid angiopathy cause the majority of these hemorrhages. Advanced age and heavy alcohol consumption increase the risk, and cocaine and methamphetamine use is one of the most important causes in the young.

Hypertensive Intracerebral Hemorrhage • PATHOPHYSIOLOGY Hypertensive ICH usually results from spontaneous rupture of a small penetrating artery deep in the brain. The most common sites are the basal ganglia (especially the putamen), thalamus, cerebellum, and pons. The small arteries in these areas seem most prone to hypertension-induced vascular injury. When hemorrhages occur in other brain areas or in nonhypertensive patients, greater consideration should be given to other causes such as hemorrhagic disorders, neoplasms, vascular malformations, and cerebral amyloid angiopathy. The hemorrhage may be small, or a large clot may form and compress adjacent tissue, causing herniation and death. Blood may also dissect into the ventricular space, which substantially increases morbidity and may cause hydrocephalus.

Most hypertensive ICHs initially develop over 30–90 min, whereas those associated with anticoagulant therapy may evolve for as long as 24–48 h. However, it is now recognized that about a third of patients even with no coagulopathy may have significant hematoma expansion with the first day. Within 48 h, macrophages begin to phagocytize the hemorrhage at its outer surface. After 1–6 months, the hemorrhage is generally resolved to a slitlike orange cavity lined with glial scar and hemosiderin-laden macrophages.

CLINICAL MANIFESTATIONS ICH generally presents as the abrupt onset of a focal neurologic deficit. Seizures are uncommon. Although clinical symptoms may be maximal at onset, commonly the focal deficit worsens over 30–90 min and is associated with a diminishing level of consciousness and signs of increased ICP such as headache and vomiting.

The putamen is the most common site for hypertensive hemorrhage, and the adjacent internal capsule is usually damaged (Fig. 446-17). Contralateral hemiparesis is therefore the sentinel sign.

TABLE 446-6 CAUSES OF INTRACRANIAL HEMORRHAGE

Cause	Location	Comments
Head trauma	Intraparenchymal: frontal lobes, anterior temporal lobes; subarachnoid; extra-axial (subdural, epidural)	Coup and contrecoup injury during brain deceleration
Hypertensive hemorrhage	Putamen, globus pallidus, thalamus, cerebellar hemisphere, pons	Chronic hypertension produces hemorrhage from small (~30–100 μ m) vessels in these regions
Transformation of prior ischemic infarction	Basal ganglion, subcortical regions, lobar	Occurs in 1–6% of ischemic strokes with predilection for large hemispheric infarctions
Metastatic brain tumor	Lobar	Lung, choriocarcinoma, melanoma, renal cell carcinoma, thyroid, atrial myxoma
Coagulopathy	Any	Risk for ongoing hematoma expansion
Drug	Any, lobar, subarachnoid	Cocaine, amphetamine
Arteriovenous malformation	Lobar, intraventricular, subarachnoid	Risk is ~2–3% per year for bleeding if previously unruptured
Aneurysm	Subarachnoid, intraparenchymal, rarely subdural	Mycotic and nonmycotic forms of aneurysms
Amyloid angiopathy	Lobar	Degenerative disease of intracranial vessels; associated with dementia, rare in patients <60 years
Cavernous angioma	Intraparenchymal	Multiple cavernous angiomas linked to mutations in <i>KRIT1</i> , <i>CCM2</i> , and <i>PDCD10</i> genes
Dural arteriovenous fistula	Lobar, subarachnoid	Produces bleeding by venous hypertension
Capillary telangiectasias	Usually brainstem	Rare cause of hemorrhage

When mild, the face sags on one side over 5–30 min, speech becomes slurred, the arm and leg gradually weaken, and the eyes deviate away from the side of the hemiparesis. The paralysis may worsen until the affected limbs become flaccid or extend rigidly. When hemorrhages are large, drowsiness gives way to stupor as signs of upper brainstem compression appear. Coma ensues, accompanied by deep, irregular, or intermittent respiration, a dilated and fixed ipsilateral pupil, and decerebrate rigidity. In milder cases, edema in adjacent brain tissue may cause progressive deterioration over 12–72 h.

Thalamic hemorrhages also produce a contralateral hemiplegia or hemiparesis from pressure on, or dissection into, the adjacent internal capsule. A prominent sensory deficit involving all modalities is usually present. Aphasia, often with preserved verbal repetition, may occur after hemorrhage into the dominant thalamus, and constructional apraxia or mutism occurs in some cases of nondominant hemorrhage. There may also be a homonymous visual field defect. Thalamic hemorrhages cause several typical ocular disturbances by virtue of extension

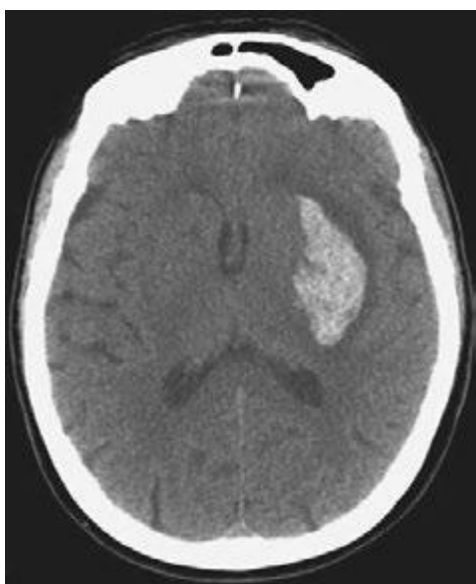


FIGURE 446-17 Hypertensive hemorrhage. Transaxial noncontrast computed tomography scan through the region of the basal ganglia reveals a hematoma involving the left putamen in a patient with rapidly progressive onset of right hemiparesis.

inferiorly into the upper midbrain. These include deviation of the eyes downward and inward so that they appear to be looking at the nose, unequal pupils with absence of light reaction, skew deviation with the eye opposite the hemorrhage displaced downward and medially, ipsilateral Horner's syndrome, absence of convergence, paralysis of vertical gaze, and retraction nystagmus. Patients may later develop a chronic, contralateral pain syndrome (Dérjérine-Roussy syndrome).

In pontine hemorrhages, deep coma with quadriplegia often occurs over a few minutes. Typically, there is prominent decerebrate rigidity and "pinpoint" (1 mm) pupils that react to light. There is impairment of reflex horizontal eye movements evoked by head turning (doll's-head or oculocephalic maneuver) or by irrigation of the ears with ice water (Chap. 328). Hyperpnea, severe hypertension, and hyperhidrosis are common. Most patients with deep coma from pontine hemorrhage ultimately die, but small hemorrhages are compatible with survival.

Cerebellar hemorrhages usually develop over several hours and are characterized by occipital headache, repeated vomiting, and ataxia of gait. In mild cases, there may be no other neurologic signs except for gait ataxia. Dizziness or vertigo may be prominent. There is often paresis of conjugate lateral gaze toward the side of the hemorrhage, forced deviation of the eyes to the opposite side, or an ipsilateral sixth nerve palsy. Less frequent ocular signs include blepharospasm, involuntary closure of one eye, ocular bobbing, and skew deviation. Dysarthria and dysphagia may occur. As the hours pass, the patient often becomes stuporous and then comatose from brainstem compression or obstructive hydrocephalus; immediate surgical evacuation before brainstem compression occurs may be lifesaving. Hydrocephalus from fourth ventricle compression can be relieved by external ventricular drainage, but definitive hematoma evacuation is recommended. If the deep cerebellar nuclei are spared, full recovery is common.

Lobar Hemorrhage The major neurologic deficit with an occipital hemorrhage is hemianopia; with a left temporal hemorrhage, aphasia and delirium; with a parietal hemorrhage, hemisensory loss; and with frontal hemorrhage, arm weakness. Large hemorrhages may be associated with stupor or coma if they compress the thalamus or midbrain. Most patients with lobar hemorrhages have focal headaches, and more than one-half vomit or are drowsy. Stiff neck and seizures are uncommon.

Other Causes of Intracerebral Hemorrhage *Cerebral amyloid angiopathy* is a disease of the elderly in which arteriolar degeneration occurs and amyloid is deposited in the walls of the cerebral arteries. Amyloid angiopathy causes both single and recurrent lobar hemorrhages and is probably the most common cause of lobar hemorrhage in the elderly.

It accounts for some intracranial hemorrhages associated with IV thrombolysis given for MI. This disorder can be suspected in patients who present with multiple hemorrhages (and infarcts) over several months or years or in patients with “microbleeds” seen on brain MRI sequences sensitive for hemosiderin (iron-sensitive imaging), but it is definitively diagnosed by pathologic demonstration of Congo red staining of amyloid in cerebral vessels. The $\epsilon 2$ and $\epsilon 4$ allelic variations of the apolipoprotein E gene are associated with increased risk of recurrent lobar hemorrhage and may therefore be markers of amyloid angiopathy. Currently, there is no specific therapy. OACs are typically avoided.

Cocaine and *methamphetamine* are frequent causes of stroke in young (age <45 years) patients. ICH, ischemic stroke, and SAH are all associated with stimulant use. Angiographic findings vary from completely normal arteries to large-vessel occlusion or stenosis, vasospasm, or changes consistent with vasculopathy. The mechanism of sympathomimetic-related stroke is not known, but cocaine enhances sympathetic activity causing acute, sometimes severe, hypertension, and this may lead to hemorrhage. Slightly more than one-half of stimulant-related intracranial hemorrhages are intracerebral, and the rest are subarachnoid. In cases of SAH, a saccular aneurysm is usually identified. Presumably, acute hypertension causes aneurysmal rupture.

Head injury often causes intracranial bleeding. The common sites are intraparenchymal (especially temporal and inferior frontal lobes) and into the subarachnoid, subdural, and epidural spaces. Trauma must be considered in any patient with an unexplained acute neurologic deficit (hemiparesis, stupor, or confusion), particularly if the deficit occurred in the context of a fall ([Chap. 457e](#)).

Intracranial hemorrhages associated with *anticoagulant therapy* can occur at any location; they are often lobar or subdural. Anticoagulant-related ICHs may continue to evolve over 24–48 h, especially if coagulopathy is insufficiently reversed. Coagulopathy and thrombocytopenia should be reversed rapidly, as discussed below. ICH associated with *hematologic disorders* (leukemia, aplastic anemia, thrombocytopenic purpura) can occur at any site and may present as multiple ICHs. Skin and mucous membrane bleeding may be evident and offers a diagnostic clue.

Hemorrhage into a *brain tumor* may be the first manifestation of neoplasm. Choriocarcinoma, malignant melanoma, renal cell carcinoma, and bronchogenic carcinoma are among the most common metastatic tumors associated with ICH. Glioblastoma multiforme in adults and medulloblastoma in children may also have areas of ICH.

Hypertensive encephalopathy is a complication of malignant hypertension. In this acute syndrome, severe hypertension is associated with headache, nausea, vomiting, convulsions, confusion, stupor, and coma. Focal or lateralizing neurologic signs, either transitory or permanent, may occur but are infrequent and therefore suggest some other vascular disease (hemorrhage, embolism, or atherosclerotic thrombosis). There are retinal hemorrhages, exudates, papilledema (hypertensive retinopathy), and evidence of renal and cardiac disease. In most cases, ICP and CSF protein levels are elevated. MRI brain imaging shows a pattern of typically posterior (occipital > frontal) brain edema that is reversible and termed *reversible posterior leukoencephalopathy*. The hypertension may be essential or due to chronic renal disease, acute glomerulonephritis, acute toxemia of pregnancy, pheochromocytoma, or other causes. Lowering the blood pressure reverses the process, but stroke can occur, especially if blood pressure is lowered too rapidly. Neuropathologic examination reveals multifocal to diffuse cerebral edema and hemorrhages of various sizes from petechial to massive. Microscopically, there are necrosis of arterioles, minute cerebral infarcts, and hemorrhages. The term *hypertensive encephalopathy* should be reserved for this syndrome and not for chronic recurrent headaches, dizziness, recurrent TIAs, or small strokes that often occur in association with high blood pressure.

Primary intraventricular hemorrhage is rare and should prompt investigation for an underlying vascular anomaly. Sometimes bleeding begins within the periventricular substance of the brain and dissects into the ventricular system without leaving signs of intraparenchymal hemorrhage. Alternatively, bleeding can arise from periependymal

veins. Vasculitis, usually polyarteritis nodosa or lupus erythematosus, can produce hemorrhage in any region of the central nervous system; most hemorrhages are associated with hypertension, but the arteritis itself may cause bleeding by disrupting the vessel wall. Nearly one-half of patients with primary intraventricular hemorrhage have identifiable bleeding sources seen using conventional angiography.

Sepsis can cause small petechial hemorrhages throughout the cerebral white matter. *Moyamoya disease*, mainly an occlusive arterial disease that causes ischemic symptoms, may on occasion produce ICH, particularly in the young. Hemorrhages into the spinal cord are usually the result of an AVM, cavernous malformation, or metastatic tumor. *Epidural spinal hemorrhage* produces a rapidly evolving syndrome of spinal cord or nerve root compression ([Chap. 456](#)). Spinal hemorrhages usually present with sudden back pain and some manifestation of myelopathy.

Laboratory and Imaging Evaluation Patients should have routine blood chemistries and hematologic studies. Specific attention to the platelet count and PT/PTT/INR is important to identify coagulopathy. CT imaging reliably detects acute focal hemorrhages in the supratentorial space. Rarely very small pontine or medullary hemorrhages may not be well delineated because of motion and bone-induced artifact that obscure structures in the posterior fossa. After the first 2 weeks, x-ray attenuation values of clotted blood diminish until they become isodense with surrounding brain. Mass effect and edema may remain. In some cases, a surrounding rim of contrast enhancement appears after 2–4 weeks and may persist for months. MRI, although more sensitive for delineating posterior fossa lesions, is generally not necessary for primary diagnosis in most instances. Images of flowing blood on MRI scan may identify AVMs as the cause of the hemorrhage. MRI, CT angiography (CTA), and conventional x-ray angiography are used when the cause of intracranial hemorrhage is uncertain, particularly if the patient is young or not hypertensive and the hematoma is not in one of the usual sites for hypertensive hemorrhage. CTA or postcontrast CT imaging may reveal one or more small areas of enhancement within a hematoma; this “spot sign” is thought to represent ongoing bleeding. The presence of a spot sign is associated with an increased risk of hematoma expansion, increased mortality, and lower likelihood of favorable functional outcome. Some centers routinely perform CT with CTA and postcontrast CT at the time of initial imaging to rapidly identify any macrovascular etiology of the hemorrhage and provide prognostic information at the same time. Because patients typically have focal neurologic signs and obtundation and often show signs of increased ICP, a lumbar puncture is generally unnecessary and should usually be avoided because it may induce cerebral herniation.

TREATMENT INTRACEREBRAL HEMORRHAGE

ACUTE MANAGEMENT

Although about 40% of patients with a hypertensive ICH die, others have a good to complete recovery if they survive the initial hemorrhage. The ICH Score ([Table 446-7](#)) is a validated clinical grading scale that is useful for stratification of mortality risk and clinical outcome. Any identified coagulopathy should be corrected as soon as possible. For patients taking VKAs, rapid correction of coagulopathy can be achieved by infusing prothrombin complex concentrates (PCC), which can be administered quickly, with vitamin K administered concurrently. Fresh frozen plasma is an alternative but generally requires larger fluid volumes and longer time to achieve adequate reversal than PCC. There is no effective antidote to ICH associated with oral thrombin inhibitor dabigatran, although FEIBA (factor VIII inhibitor bypassing activity) and recombinant factor VIIa have been tried in individual cases. PCC may partially reverse the effects of oral factor Xa inhibitors and are reasonable to administer if available. When ICH is associated with thrombocytopenia (platelet count <50,000/ μ L), transfusion of fresh platelets is indicated. The role of platelet transfusions either empirically or based on urgent platelet inhibition assays remains unclear.

TABLE 446-7 THE ICH SCORE

Clinical or Imaging Factor	Point Score	
Age		
<80 years	0	
≥80 years	1	
Hematoma Volume		
<30 cc	0	
≥30 cc	1	
Intraventricular Hemorrhage Present		
No	0	
Yes	1	
Infratentorial Origin of Hemorrhage		
No	0	
Yes	1	
Glasgow Coma Scale Score		
13–15	0	
5–12	1	
3–4	2	
Total Score	Sum of each category above	
ICH Score Total	% Mortality at 30 Days (95% CI)	% Walk Independently at 12 Months (95% CI)
0	0 (0–13)	70 (53–84)
1	13 (4–29)	60 (47–73)
2	26 (11–46)	33 (21–48)
3	72 (53–86)	13 (5–25)
4	97 (82–100)	3 (0–16)
5	100 (54–100)	8 (0–38)

Note: Although an ICH Score of 6 is possible with the scale, this is rarely observed and is considered highly likely to be fatal.

Abbreviations: CI, confidence interval; ICH, intracerebral hemorrhage.

Sources: JC Hemphill et al: *Stroke* 32:891, 2001; JC Hemphill et al: *Neurology* 73:1088, 2009.

Hematomas may expand for several hours following the initial hemorrhage, even in patients without coagulopathy. However, the precise mechanism is unclear. A phase 3 trial of treatment with recombinant factor VIIa reduced hematoma expansion; however, clinical outcomes were not improved, so use of this drug cannot be advocated at present. The theoretical risk of acutely elevated blood pressure on hematoma expansion forms the basis of the consideration for recently completed and ongoing clinical trials of acute blood pressure lowering.

Evacuation of supratentorial hematomas does not appear to improve outcome for most patients. The International Surgical Trial in Intracerebral Haemorrhage (STICH) randomized patients with supratentorial ICH to either early surgical evacuation or initial medical management. No benefit was found in the early surgery arm, although analysis was complicated by the fact that 26% of patients in the initial medical management group ultimately had surgery for neurologic deterioration. The follow-up study STICH-II found that surgery within 24 h of lobar, supratentorial hemorrhage did not improve overall outcome, but might have a role in select severely affected patients. Therefore, existing data do not support routine surgical evacuation of supratentorial hemorrhages in stable patients. However, many centers still consider surgery for patients deemed salvageable and who are having progressive neurologic deterioration due to herniation. Surgical techniques continue to evolve, and minimally invasive endoscopic hematoma evacuation is currently being investigated in clinical trials.

For cerebellar hemorrhages, a neurosurgeon should be consulted immediately to assist with the evaluation; most cerebellar hematomas >3 cm in diameter will require surgical evacuation. If the patient is alert without focal brainstem signs and if the hematoma is <1 cm in diameter, surgical removal is usually unnecessary. Patients with

hematomas between 1 and 3 cm require careful observation for signs of impaired consciousness, progressive hydrocephalus, and precipitous respiratory failure. Hydrocephalus due to cerebellar hematoma should not be treated solely with ventricular drainage.

Tissue surrounding hematomas is displaced and compressed but not necessarily infarcted. Hence, in survivors, major improvement commonly occurs as the hematoma is reabsorbed and the adjacent tissue regains its function. Careful management of the patient during the acute phase of the hemorrhage can lead to considerable recovery.

Surprisingly, ICP is often normal even with large ICHs. However, if the hematoma causes marked midline shift of structures with consequent obtundation, coma, or hydrocephalus, osmotic agents can be instituted in preparation for placement of a ventriculostomy or parenchymal ICP monitor (**Chap. 330**). Once ICP is recorded, CSF drainage (if available), osmotic therapy, and blood pressure management can be tailored to the individual patient to keep cerebral perfusion pressure (MAP minus ICP) above 60 mmHg. For example, if ICP is found to be high, CSF can be drained from the ventricular space and osmotic therapy continued; persistent or progressive elevation in ICP may prompt surgical evacuation of the clot. Alternatively, if ICP is normal or only mildly elevated, interventions such as osmotic therapy may be tapered. Because hyperventilation may actually produce ischemia by cerebral vasoconstriction, induced hyperventilation should be limited to acute resuscitation of the patient with presumptive high ICP and eliminated once other treatments (osmotic therapy or surgical treatments) have been instituted. Glucocorticoids are not helpful for the edema from intracerebral hematoma.

PREVENTION

Hypertension is the leading cause of primary ICH. Prevention is aimed at reducing chronic hypertension, eliminating excessive alcohol use, and discontinuing use of illicit drugs such as cocaine and amphetamines. Patients with amyloid angiopathy should generally avoid OACs, but antiplatelet agents may be administered if there is an indication based on atherothrombotic vascular disease.

VASCULAR ANOMALIES

Vascular anomalies can be divided into congenital vascular malformations and acquired vascular lesions.

CONGENITAL VASCULAR MALFORMATIONS

True *arteriovenous malformations* (AVMs), venous anomalies, and capillary telangiectasias are lesions that usually remain clinically silent through life. AVMs are probably congenital, but cases of acquired lesions have been reported.

True AVMs are congenital shunts between the arterial and venous systems that may present with headache, seizures, and intracranial hemorrhage. AVMs consist of a tangle of abnormal vessels across the cortical surface or deep within the brain substance. AVMs vary in size from a small blemish a few millimeters in diameter to a large mass of tortuous channels composing an arteriovenous shunt of sufficient magnitude to raise cardiac output and precipitate heart failure. Blood vessels forming the tangle interposed between arteries and veins are usually abnormally thin and histologically resemble both arteries and veins. AVMs occur in all parts of the cerebral hemispheres, brainstem, and spinal cord, but the largest ones are most frequently in the posterior half of the hemispheres, commonly forming a wedge-shaped lesion extending from the cortex to the ventricle.

Bleeding, headache, and seizures are most common between the ages of 10 and 30, occasionally as late as the fifties. AVMs are more frequent in men, and rare familial cases have been described. Familial AVM may be a part of the autosomal dominant syndrome of hereditary hemorrhagic telangiectasia (Osler-Rendu-Weber) syndrome due to mutations in either endoglin or activin receptor-like kinase 1, both involved in transforming growth factor (TGF) signaling and angiogenesis.

Headache (without bleeding) may be hemicranial and throbbing, like migraine, or diffuse. Focal seizures, with or without generalization, occur in ~30% of cases. One-half of AVMs become evident as ICHs. In most, the hemorrhage is mainly intraparenchymal with extension into the subarachnoid space in some cases. Blood is usually not deposited in the basal cisterns, and symptomatic cerebral vasospasm is rare. The risk of AVM rupture is strongly influenced by a history of prior rupture. Although unruptured AVMs have a hemorrhage rate of ~2–4% per year, previously ruptured AVMs may have a rate as high as 17% a year, at least for the first year. Hemorrhages may be massive, leading to death, or may be as small as 1 cm in diameter, leading to minor focal symptoms or no deficit. The AVM may be large enough to steal blood away from adjacent normal brain tissue or to increase venous pressure significantly to produce venous ischemia locally and in remote areas of the brain. This is seen most often with large AVMs in the territory of the MCA.

Large AVMs of the anterior circulation may be associated with a systolic and diastolic bruit (sometimes self-audible) over the eye, forehead, or neck and a bounding carotid pulse. Headache at the onset of AVM rupture is generally not as explosive as with aneurysmal rupture. MRI is better than CT for diagnosis, although noncontrast CT scanning sometimes detects calcification of the AVM and contrast may demonstrate the abnormal blood vessels. Once identified, conventional x-ray angiography is the gold standard for evaluating the precise anatomy of the AVM.

Surgical treatment of AVMs presenting with hemorrhage often done in conjunction with preoperative embolization to reduce operative bleeding is usually indicated for accessible lesions. Stereotaxic radiation, an alternative to surgery, can produce a slow sclerosis of the AVM over 2–3 years.

Several angiographic features can be used to help predict future bleeding risk. Paradoxically, smaller lesions seem to have a higher hemorrhage rate. The presence of deep venous drainage, venous outflow stenosis, and intranidal aneurysms may increase rupture risk. Because of the relatively low annual rate of hemorrhage and the risk of complications due to surgical or endovascular treatment, the indication for surgery in asymptomatic AVMs is debated. The ARUBA (A Randomized Trial of Unruptured Brain Arteriovenous Malformations) trial randomized patients to medical management versus intervention (surgery, endovascular embolization, combination embolization and surgery, or gamma-knife). The trial was stopped prematurely for harm, with the medical arm achieving the combined endpoint of death or symptomatic stroke in 10.1% of patients compared to 30.7% in the intervention group at an average follow-up time of 33 months. This highly significant finding argues against routine intervention for patients presenting without hemorrhage, although debate ensues regarding the generalizability of these results.

Venous anomalies are the result of development of anomalous cerebral, cerebellar, or brainstem venous drainage. These structures, unlike AVMs, are functional venous channels. They are of little clinical significance and should be ignored if found incidentally on brain imaging studies. Surgical resection of these anomalies may result in venous infarction and hemorrhage. Venous anomalies may be associated with cavernous malformations (see below), which do carry some bleeding risk.

Capillary telangiectasias are true capillary malformations that often form extensive vascular networks through an otherwise normal brain structure. The pons and deep cerebral white matter are typical locations, and these capillary malformations can be seen in patients with hereditary hemorrhagic telangiectasia (Osler-Rendu-Weber) syndrome. If bleeding does occur, it rarely produces mass effect or significant symptoms. No treatment options exist.

ACQUIRED VASCULAR LESIONS

Cavernous angiomas are tufts of capillary sinusoids that form within the deep hemispheric white matter and brainstem with no normal intervening neural structures. The pathogenesis is unclear. Familial cavernous angiomas have been mapped to several different genes: *KRIT1*, *CCM2*, and *PDCD10*. Both *KRIT1* and *CCM2* have roles

in blood vessel formation, whereas *PDCD10* is an apoptotic gene. Cavernous angiomas are typically <1 cm in diameter and are often associated with a venous anomaly. Bleeding is usually of small volume, causing slight mass effect only. The bleeding risk for single cavernous malformations is 0.7–1.5% per year and may be higher for patients with prior clinical hemorrhage or multiple malformations. Seizures may occur if the malformation is located near the cerebral cortex. Surgical resection eliminates bleeding risk and may reduce seizure risk, but it is usually reserved for those malformations that form near the brain surface. Radiation treatment has not been shown to be of benefit.

Dural arteriovenous fistulas are acquired connections usually from a dural artery to a dural sinus. Patients may complain of a pulse-synchronous cephalic bruit (“pulsatile tinnitus”) and headache. Depending on the magnitude of the shunt, venous pressures may rise high enough to cause cortical ischemia or venous hypertension and hemorrhage, particularly SAH. Surgical and endovascular techniques are usually curative. These fistulas may form because of trauma, but most are idiopathic. There is an association between fistulas and dural sinus thrombosis. Fistulas have been observed to appear months to years following venous sinus thrombosis, suggesting that angiogenesis factors elaborated from the thrombotic process may cause these anomalous connections to form. Alternatively, dural arteriovenous fistulas can produce venous sinus occlusion over time, perhaps from the high pressure and high flow through a venous structure.

447 Migraine and Other Primary Headache Disorders

Peter J. Goadsby, Neil H. Raskin

The general principles around headache as a cardinal symptom are covered elsewhere ([Chap. 21](#)); here we discuss disorders in which headache and associated features occur in the absence of any exogenous cause. The most common are migraine, tension-type headache, and the trigeminal autonomic cephalalgias, notably cluster headache; the complete list is summarized in [Table 447-1](#).

MIGRAINE

Migraine, the second most common cause of headache, and the most common headache-related, and indeed neurologic, cause of disability in the world, afflicts approximately 15% of women and 6% of men over a 1-year period. It is usually an episodic headache associated with certain features such as sensitivity to light, sound, or movement; nausea and vomiting often accompany the headache. A useful description of migraine is a recurring syndrome of headache associated with other symptoms of neurologic dysfunction in varying admixtures ([Table 447-2](#)). Migraine can often be recognized by its activators, referred to as *triggers*.

The brain of the migraineur is particularly sensitive to environmental and sensory stimuli; migraine-prone patients do not habituate easily to sensory stimuli. This sensitivity is amplified in females during the menstrual cycle. Headache can be initiated or amplified by various triggers, including glare, bright lights, sounds, or other afferent stimulation; hunger; let-down from stress; physical exertion; stormy weather or barometric pressure changes; hormonal fluctuations during menses; lack of or excess sleep; and alcohol or other chemical stimulation, such as with nitrates. Knowledge of a patient’s susceptibility to specific triggers can be useful in management strategies involving lifestyle adjustments.

Pathogenesis The sensory sensitivity that is characteristic of migraine is probably due to dysfunction of monoaminergic sensory control systems located in the brainstem and hypothalamus ([Fig. 447-1](#)).

Activation of cells in the trigeminal nucleus results in the release of vasoactive neuropeptides, particularly calcitonin gene-related peptide

TABLE 447-1 PRIMARY HEADACHE DISORDERS, MODIFIED FROM INTERNATIONAL CLASSIFICATION OF HEADACHE DISORDERS-III-BETA (HEADACHE CLASSIFICATION COMMITTEE OF THE INTERNATIONAL HEADACHE SOCIETY, 2013)

1. Migraine	<ul style="list-style-type: none"> 1.1 Migraine without aura 1.2 Migraine with aura <ul style="list-style-type: none"> 1.2.1 Migraine with typical aura <ul style="list-style-type: none"> 1.2.1.1 Typical aura with headache 1.2.1.2 Typical aura without headache 1.2.2 Migraine with brainstem aura 1.2.3 Hemiplegic migraine <ul style="list-style-type: none"> 1.2.3.1 Familial hemiplegic migraine (FHM) <ul style="list-style-type: none"> 1.2.3.1.1 Familial hemiplegic migraine type 1 1.2.3.1.2 Familial hemiplegic migraine type 2 1.2.3.1.3 Familial hemiplegic migraine type 3 1.2.3.2 Sporadic hemiplegic migraine 1.2.4 Retinal migraine 1.3 Chronic migraine 1.4 Complications of migraine <ul style="list-style-type: none"> 1.4.1 Status migrainosus 1.4.2 Persistent aura without infarction 1.4.3 Migrainous infarction 1.4.4 Migraine aura-triggered seizure 1.5 Probable migraine <ul style="list-style-type: none"> 1.5.1 Probable migraine without aura 1.5.2 Probable migraine with aura 1.6 Episodic syndromes that may be associated with migraine <ul style="list-style-type: none"> 1.6.1 Recurrent gastrointestinal disturbance <ul style="list-style-type: none"> 1.6.1.1 Cyclical vomiting syndrome 1.6.1.2 Abdominal migraine 1.6.2 Benign paroxysmal vertigo 1.6.3 Benign paroxysmal torticollis
2. Tension-type headache	<ul style="list-style-type: none"> 2.1 Infrequent episodic tension-type headache 2.2 Frequent episodic tension-type headache 2.3 Chronic tension-type headache
3. Trigeminal autonomic cephalalgias	<ul style="list-style-type: none"> 3.1 Cluster headache <ul style="list-style-type: none"> 3.1.1 Episodic cluster headache 3.1.2 Chronic cluster headache 3.2 Paroxysmal hemicrania <ul style="list-style-type: none"> 3.2.1 Episodic paroxysmal hemicrania 3.2.2 Chronic paroxysmal hemicrania 3.3 Short-lasting unilateral neuralgiform headache attacks <ul style="list-style-type: none"> 3.3.1 Short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing (SUNCT) 3.3.2 Short-lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms (SUNA) 3.4 Hemicrania continua
4. Other primary headache disorders	<ul style="list-style-type: none"> 4.1 Primary cough headache 4.2 Primary exercise headache 4.3 Primary headache associated with sexual activity 4.4 Primary thunderclap headache 4.5 Cold-stimulus headache <ul style="list-style-type: none"> 4.5.1 Headache attributed to external application of a cold stimulus 4.5.2 Headache attributed to ingestion or inhalation of a cold stimulus 4.6 External-pressure headache <ul style="list-style-type: none"> 4.6.1 External-compression headache 4.6.2 External-traction headache 4.7 Primary stabbing headache 4.8 Nummular headache 4.9 Hypnic headache 4.10 New daily persistent headache (NDPH)

TABLE 447-2 SYMPTOMS ACCOMPANYING SEVERE MIGRAINE ATTACKS IN 500 PATIENTS

Symptom	Patients Affected, %
Nausea	87
Photophobia	82
Lightheadedness	72
Scalp tenderness	65
Vomiting	56
Visual disturbances	36
Paresthesias	33
Vertigo	33
Photopsia	26
Alteration of consciousness	18
Diarrhea	16
Fortification spectra	10
Syncope	10
Seizure	4
Confusional state	4

Source: From NH Raskin: *Headache*, 2nd ed. New York, Churchill Livingstone, 1988; with permission.

(CGRP), at vascular terminations of the trigeminal nerve and within the trigeminal nucleus. CGRP receptor antagonists, *gepants*, have now been shown to be effective in the acute treatment of migraine, and monoclonal antibodies to CGRP have been shown effective in two early phase clinical trials. Centrally, the second-order trigeminal neurons cross the midline and project to ventrobasal and posterior nuclei of the thalamus for further processing. Additionally, there are

projections to the periaqueductal gray and hypothalamus, from which reciprocal descending systems have established antinociceptive effects. Other brainstem regions likely to be involved in descending modulation of trigeminal pain include the nucleus locus coeruleus in the pons and the rostroventromedial medulla.

Pharmacologic and other data point to the involvement of the neurotransmitter 5-hydroxytryptamine (5-HT; also known as serotonin) in migraines. Approximately 60 years ago, methysergide was found to antagonize certain peripheral actions of 5-HT and was introduced as the first drug capable of preventing migraine attacks. The *triptans* were designed to stimulate selectively subpopulations of 5-HT receptors; at least 14 different 5-HT receptors exist in humans. The triptans are potent agonists of 5-HT_{1B} and 5-HT_{1D} receptors, and some are active at the 5-HT_{1F} receptors; the latter's exclusive agonists are called *ditans*. Triptans arrest nerve signaling in the nociceptive pathways of the trigeminovascular system, at least in the trigeminal nucleus caudalis and trigeminal sensory thalamus, in addition to cranial vasoconstriction, while *ditans*, now shown conclusively to be effective in acute migraine, act only at neural targets. An interesting range of neural targets is now being actively pursued for the acute and preventive management of migraine.

Data also support a role for dopamine in the pathophysiology of migraine. Most migraine symptoms can be induced by dopaminergic stimulation. Moreover, there is dopamine receptor hypersensitivity in migraineurs, as demonstrated by the induction of yawning, nausea, vomiting, hypotension, and other symptoms of a migraine attack by dopaminergic agonists at doses that do not affect nonmigraineurs. Dopamine receptor antagonists are effective therapeutic agents in migraine, especially when given parenterally or concurrently with other antimigraine agents. Moreover, hypothalamic activation, anterior to that seen in cluster headache, has now been shown in the

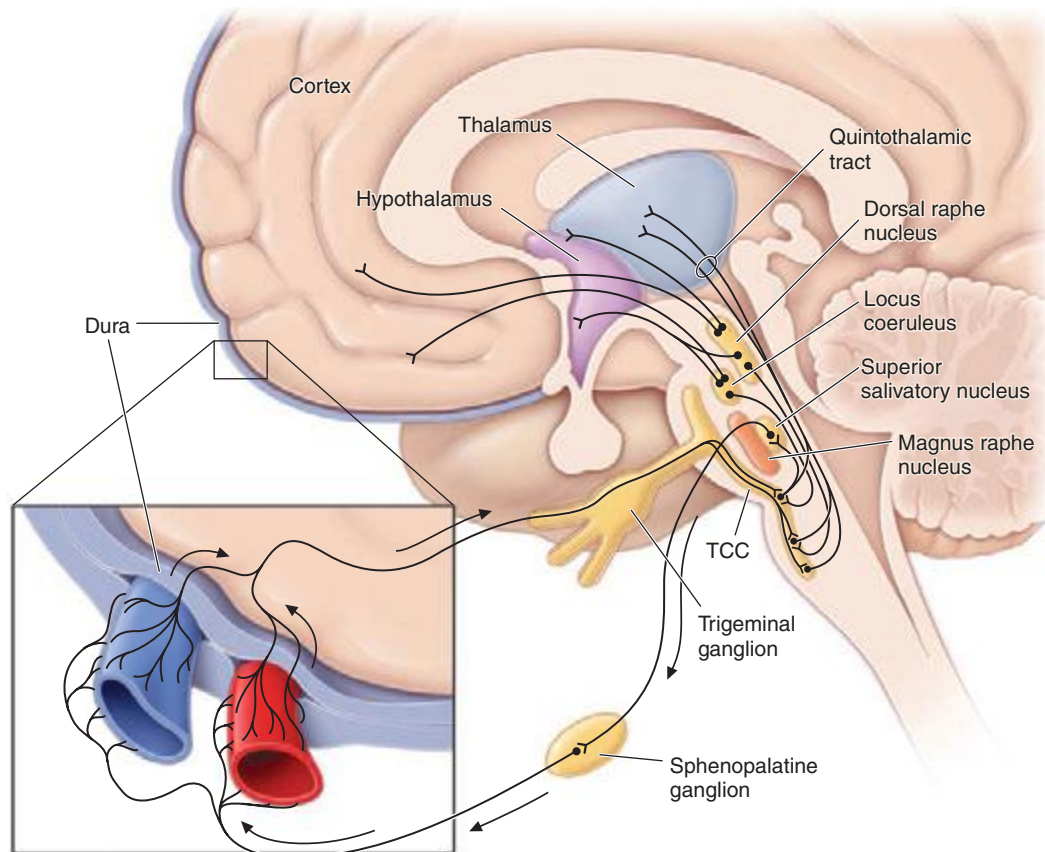


FIGURE 447-1 Brainstem pathways that modulate sensory input. The key pathway for pain in migraine is the trigeminovascular input from the meningeal vessels, which passes through the trigeminal ganglion and synapses on second-order neurons in the trigeminocervical complex (TCC). These neurons in turn project in the quintothalamic tract and, after decussating in the brainstem, synapse on neurons in the thalamus. Important modulation of the trigeminovascular nociceptive input comes from the dorsal raphe nucleus, locus coeruleus, and nucleus raphe magnus.

premonitory phase of migraine using functional imaging, and this may hold a key to understanding some part of the role of dopamine in the disorder.

Migraine genes identified by studying families with familial hemiplegic migraine (FHM) reveal involvement of ion channels, suggesting that alterations in membrane excitability can predispose to migraine. Mutations involving the $\text{Ca}_v2.1$ (P/Q)-type voltage-gated calcium channel *CACNA1A* gene are now known to cause FHM 1; this mutation is responsible for about 50% of FHMs. Mutations in the $\text{Na}^+\text{-K}^+\text{ATPase}$ *ATP1A2* gene, designated FHM 2, are responsible for about 20% of FHMs. Mutations in the neuronal voltage-gated sodium channel *SCN1A* cause FHM 3. Functional neuroimaging has suggested that brainstem regions in migraine (Fig. 447-2) and the posterior hypothalamic gray matter region close to the human circadian pacemaker cells of the suprachiasmatic nucleus in cluster headache (Fig. 447-3) are good candidates for specific involvement in primary headache.

Diagnosis and Clinical Features Diagnostic criteria for migraine headache are listed in Table 447-3. A high index of suspicion is required to diagnose migraine: the migraine aura, consisting of visual disturbances with flashing lights or zigzag lines moving across the visual field or of

other neurologic symptoms, is reported in only 20–25% of patients. A headache diary can often be helpful in making the diagnosis; this is also helpful in assessing disability and the frequency of treatment for acute attacks. Patients with episodes of migraine that occur daily or near-daily are considered to have chronic migraine (see “Chronic Daily Headache” in Chap. 21). Migraine must be differentiated from tension-type headache (discussed below), the most common primary headache syndrome seen in the population. Migraine has several forms that have been defined (Table 447-1): migraine with and without aura and chronic migraine, the latter occurring 15 days or more a month, as the most important. *Migraine at its most basic level is headache with associated features, and tension-type headache is headache that is featureless. Most patients with disabling headache probably have migraine.*

Patients with acephalgic migraine (typical aura without headache, 1.2.1.2 in Table 447-1) experience recurrent neurologic symptoms, often with nausea or vomiting, but with little or no headache. Vertigo can be prominent; it has been estimated that one-third of patients referred for vertigo or dizziness have a primary diagnosis of migraine. Migraine aura can have prominent brainstem symptoms, and the terms *basilar artery* and *basilar-type migraine* have now been replaced by *migraine with brainstem aura* (Table 447-1).

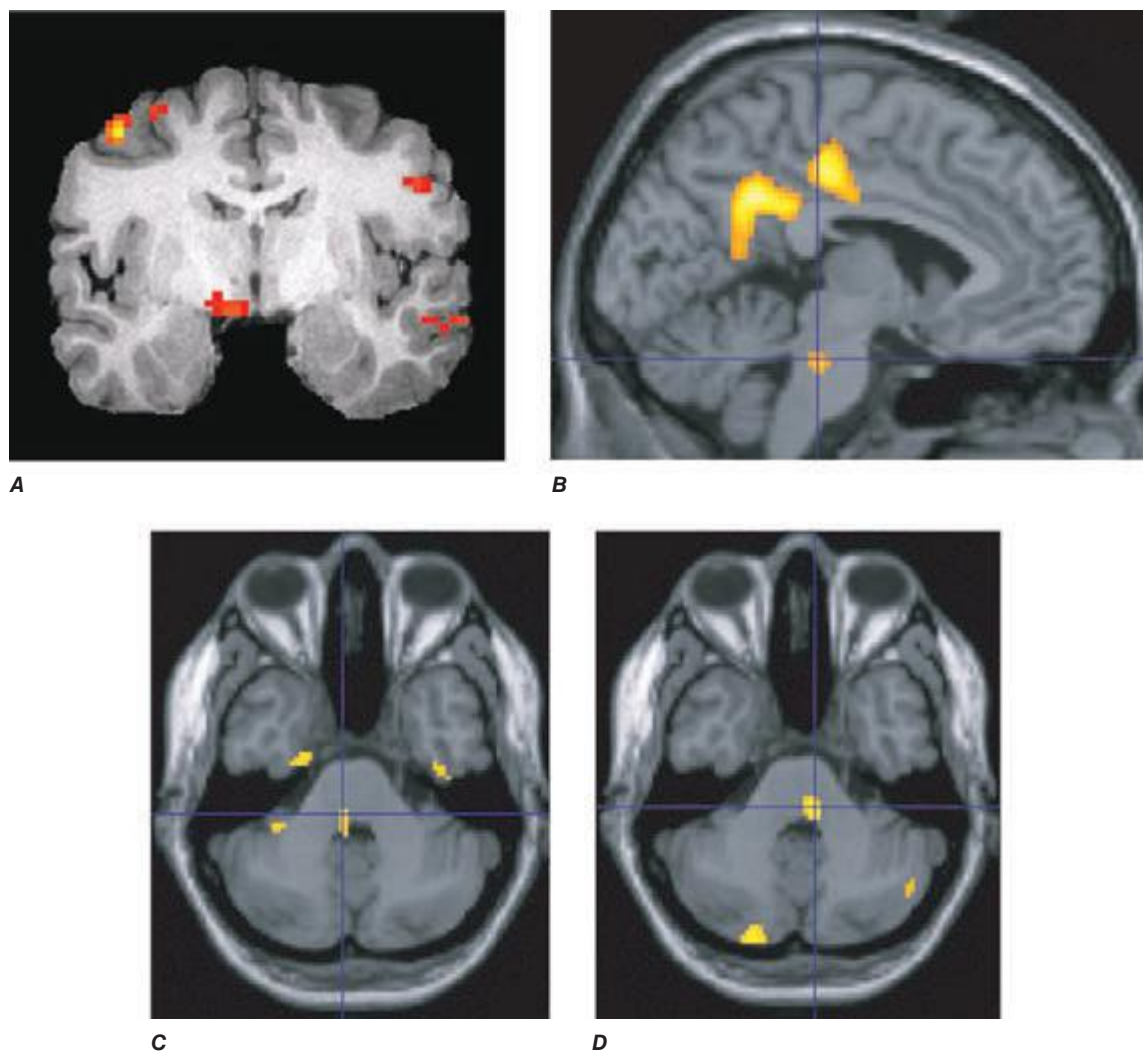


FIGURE 447-2 Positron emission tomography (PET) activation in migraine. Hypothalamic, dorsal midbrain, and dorsolateral pontine activation is seen in triggered attacks in the premonitory phase before pain, whereas in migraine attacks, dorsolateral pontine activation persists, as it does in chronic migraine (not shown). The dorsolateral pontine area, which includes the noradrenergic locus coeruleus, is fundamental to the expression of migraine. Moreover, lateralization of changes in this region of the brainstem correlates with lateralization of the head pain in hemicranial migraine; the scans shown in panels C and D are of patients with acute migraine headache on the right and left side, respectively. (Panel A from FH Maniyar et al: *Brain* 137:232, 2014; panel B from SK Afridi et al: *Arch Neurol* 2005;62:1270; Panels C and D from SK Afridi et al: *Brain* 128:932, 2005.)

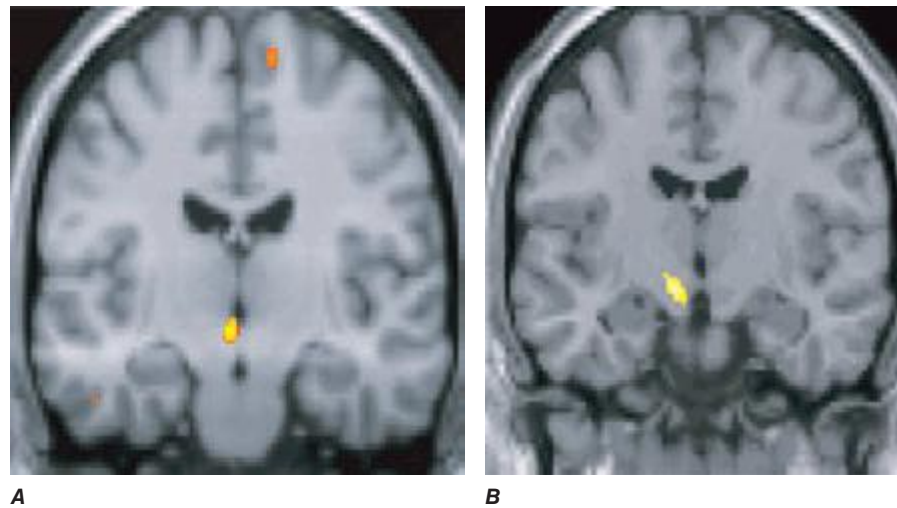


FIGURE 447-3 **A.** Posterior hypothalamic gray matter activation by positron emission tomography in a patient with acute cluster headache. (From A May et al: *Lancet* 352:275, 1998.) **B.** High-resolution T1-weighted magnetic resonance image obtained using voxel-based morphometry demonstrates increased gray matter activity, lateralized to the side of pain in a patient with cluster headache. (From A May et al: *Nat Med* 5:836, 1999.)

TREATMENT MIGRAINE HEADACHE

Once a diagnosis of migraine has been established, it is important to assess the extent of a patient's disease and disability. The Migraine Disability Assessment Score (MIDAS) is a well-validated, easy-to-use tool (Fig. 447-4).

Patient education is an important aspect of migraine management. Information for patients is available at sites such as www.achenet.org, the website of the American Council for Headache Education (ACHE). It is helpful for patients to understand that migraine is an inherited tendency to headache; that migraine can be modified and controlled by lifestyle adjustments and medications, but it cannot be eradicated; and that, except in some occasions in women on oral estrogens or contraceptives, migraine is not associated with serious or life-threatening illnesses.

NONPHARMACOLOGIC MANAGEMENT

Migraine can often be managed to some degree by a variety of nonpharmacologic approaches. Most patients benefit by the identification and avoidance of specific headache triggers. A regulated lifestyle is helpful, including a healthy diet, regular exercise, regular sleep patterns, avoidance of excess caffeine and alcohol, and avoidance of acute changes in stress levels, being particularly wary of the let-down effect.

The measures that benefit a given individual should be used routinely because they provide a simple, cost-effective approach to migraine management. Patients with migraine do not encounter more stress than headache-free individuals; over-responsiveness to changes in stress appears to be the issue. Because the stresses of everyday living cannot be eliminated, lessening one's response to stress by various techniques is helpful for many patients. These may

include yoga, transcendental meditation, hypnosis, and conditioning techniques such as biofeedback. For most patients, this approach is, at best, an adjunct to pharmacotherapy. Nonpharmacologic measures are unlikely to prevent all migraine attacks. If these measures fail to prevent an attack, pharmacologic approaches are then needed to abort an attack.

ACUTE ATTACK THERAPIES FOR MIGRAINE

The mainstay of pharmacologic therapy is the judicious use of one or more of the many medicines that are effective in migraine (Table 447-4). The selection of the optimal regimen for a given patient depends on a number of factors, the most important of which is the severity of the attack. Mild migraine attacks can usually be managed by oral agents; the average efficacy rate is 50–70%. Severe migraine attacks may require parenteral therapy. Most drugs effective in the treatment of migraine are members of one of three major pharmacologic classes: nonsteroidal anti-inflammatory drugs, 5-HT_{1B/1D} receptor agonists, and dopamine receptor antagonists.

In general, an adequate dose of whichever agent is chosen should be used as soon as possible after the onset of an attack. If additional medication is required within 60 min because symptoms return or have not abated, the initial dose should be increased for subsequent attacks or a different class of drug tried as first-line treatment. Migraine therapy must be individualized; a standard approach for all patients is not possible. A therapeutic regimen may need to be constantly refined until one is identified that provides the patient with rapid, complete, and consistent relief with minimal side effects (Table 447-5).

Nonsteroidal Anti-Inflammatory Drugs (NSAIDs) Both the severity and duration of a migraine attack can be reduced significantly by NSAIDs (Table 447-4). Indeed, many undiagnosed migraineurs self-treat with nonprescription NSAIDs. A general consensus is that NSAIDs are most effective when taken early in the migraine attack. However, the effectiveness of these agents in migraine is usually less than optimal in moderate or severe migraine attacks. The combination of acetaminophen, aspirin, and caffeine has been approved for use by the U.S. Food and Drug Administration (FDA) for the treatment of mild to moderate migraine. The combination of aspirin and metoprolamide has been shown to be comparable to a single dose of oral sumatriptan. Important side effects of NSAIDs include dyspepsia and gastrointestinal irritation.

5-HT_{1B/1D} RECEPTOR AGONISTS

Oral Stimulation of 5-HT_{1B/1D} receptors can stop an acute migraine attack. Ergotamine and dihydroergotamine are nonselective receptor

TABLE 447-3 SIMPLIFIED DIAGNOSTIC CRITERIA FOR MIGRAINE

Repeated attacks of headache lasting 4–72 h in patients with a normal physical examination, no other reasonable cause for the headache, and:

At Least 2 of the Following Features:	Plus at Least 1 of the Following Features:
Unilateral pain	Nausea/vomiting
Throbbing pain	Photophobia and phonophobia
Aggravation by movement	
Moderate or severe intensity	

Source: Adapted from the International Headache Society Classification (Headache Classification Committee of the International Headache Society, 2013).

***MIDAS Questionnaire**

INSTRUCTIONS: Please answer the following questions about ALL headaches you have had over the last 3 months. Write zero if you did not do the activity in the last 3 months.

1. On how many days in the last 3 months did you miss work or school because of your headaches? ____ days
2. How many days in the last 3 months was your productivity at work or school reduced by half or more because of your headaches (*do not include days you counted in question 1 where you missed work or school*)?..... ____ days
3. On how many days in the last 3 months did you **not** do household work because of your headaches? ____ days
4. How many days in the last 3 months was your productivity in household work reduced by half or more because of your headaches (*do not include days you counted in question 3 where you did not do household work*)?..... ____ days
5. On how many days in the last 3 months did you miss family, social, or leisure activities because of your headaches? ____ days

A. On how many days in the last 3 months did you have a headache? (*If a headache lasted more than one day, count each day.*)..... ____ days

B. On a scale of 0–10, on average how painful were these headaches? (*Where 0 = no pain at all, and 10 = pain as bad as it can be.*)..... ____

*Migraine Disability Assessment Score
(Questions 1–5 are used to calculate the MIDAS score.)
Grade I—Minimal or Infrequent Disability: 0–5
Grade II—Mild or Infrequent Disability: 6–10
Grade III—Moderate Disability: 11–20
Grade IV—Severe Disability: > 20

© Innovative Medical Research 1997

FIGURE 447-4 The Migraine Disability Assessment Score (MIDAS) Questionnaire.

agonists, whereas the triptans are selective 5-HT_{1B/1D} receptor agonists. A variety of triptans, 5-HT_{1B/1D} receptor agonists—sumatriptan, almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, and zolmitriptan—are now available for the treatment of migraine.

Each drug in the triptan class has similar pharmacologic properties but varies slightly in terms of clinical efficacy. Rizatriptan and eletriptan are the most efficacious of the triptans currently available in the United States. Sumatriptan and zolmitriptan have similar rates of efficacy as well as time to onset, with an advantage of having multiple formulations, whereas almotriptan has a similar rate of efficacy to sumatriptan and is better tolerated, and frovatriptan and naratriptan are somewhat slower in onset and are better tolerated. Clinical efficacy appears to be related more to the t_{max} (time to peak plasma level) than to the potency, half-life, or bioavailability. This observation is consistent with a large body of data indicating that faster-acting analgesics are more effective than slower-acting agents.

Unfortunately, monotherapy with a selective oral 5-HT_{1B/1D} receptor agonist does not result in rapid, consistent, and complete relief of migraine in all patients. Triptans are generally not effective in migraine with aura unless given after the aura is completed and the headache initiated. Side effects are common, although often mild and transient. Moreover, 5-HT_{1B/1D} receptor agonists are contraindicated in individuals with a history of cardiovascular and cerebrovascular disease. Recurrence of headache, within usual time course of an attack, is another important limitation of triptan use and occurs at least occasionally in most patients. Evidence from randomized controlled trials show that coadministration of a longer-acting NSAID, naproxen 500 mg, with sumatriptan will augment the initial effect of sumatriptan and, importantly, reduce rates of headache recurrence.

Ergotamine preparations offer a nonselective means of stimulating 5-HT₁ receptors. A nonnauseating dose of ergotamine should be sought because a dose that provokes nausea is too high and may intensify head pain. Except for a sublingual formulation of ergotamine, oral formulations of ergotamine also contain 100 mg caffeine (theoretically to enhance ergotamine absorption and possibly to

add additional analgesic activity). The average oral ergotamine dose for a migraine attack is 2 mg. Because the clinical studies demonstrating the efficacy of ergotamine in migraine predated the clinical trial methodologies used with the triptans, it is difficult to assess the clinical efficacy of ergotamine versus the triptans. In general, ergotamine appears to have a much higher incidence of nausea than triptans but less headache recurrence.

Nasal Nasal formulations of dihydroergotamine (Migranal), zolmitriptan (Zomig nasal), or sumatriptan can be useful in patients requiring a nonoral route of administration. The nasal sprays result in substantial blood levels within 30–60 min. Although in theory nasal sprays might provide faster and more effective relief of a migraine attack than oral formulations, their reported efficacy is only approximately 50–60%. Studies with a new inhalational formulation of dihydroergotamine indicate that its absorption problems can be overcome to produce rapid onset of action with good tolerability.

Parenteral Administration of drugs by injection, such as dihydroergotamine and sumatriptan, is approved by the FDA for the rapid relief of a migraine attack. Peak plasma levels of dihydroergotamine are achieved 3 min after IV dosing, 30 min after IM dosing, and 45 min after SC dosing. If an attack has not already peaked, SC or IM administration of 1 mg of dihydroergotamine suffices for about 80–90% of patients. Sumatriptan, 4–6 mg SC, is effective in ~50–80% of patients, and can now be administered by a needle-free device.

DOPAMINE RECEPTOR ANTAGONISTS

Oral Oral dopamine receptor antagonists can be considered as adjunctive therapy in migraine. Drug absorption is impaired during migraine because of reduced gastrointestinal motility. Delayed absorption occurs even in the absence of nausea and is related to the severity of the attack and not its duration. Therefore, when oral NSAIDs and/or triptan agents fail, the addition of a dopamine receptor antagonist, such as metoclopramide 10 mg or domperidone 10 mg (not available in the United States), should be considered to

TABLE 447-4 TREATMENT OF ACUTE MIGRAINE

Drug	Trade Name	Dosage
Simple Analgesics		
Acetaminophen, aspirin, caffeine	Excedrin Migraine	Two tablets or caplets q6h (max 8 per day)
NSAIDs		
Naproxen	Aleve, Anaprox, generic	220–550 mg PO bid
Ibuprofen	Advil, Motrin, Nuprin, generic	400 mg PO q3–4h
Tolfenamic acid	Clotam Rapid	200 mg PO; may repeat ×1 after 1–2 h
Diclofenac K	Cambia	50 mg PO with water
5-HT₁ Receptor Agonists		
Oral		
Ergotamine 1 mg, caffeine 100 mg	Cafergot	One or two tablets at onset, then one tablet q½h (max 6 per day, 10 per week)
Naratriptan	Amerge	2.5-mg tablet at onset; may repeat once after 4 h
Rizatriptan	Maxalt Maxalt-MLT	5–10-mg tablet at onset; may repeat after 2 h (max 30 mg/d)
Sumatriptan	Imitrex	50–100-mg tablet at onset; may repeat after 2 h (max 200 mg/d)
Frovatriptan	Frova	2.5-mg tablet at onset, may repeat after 2 h (max 5 mg/d)
Almotriptan	Axert	12.5-mg tablet at onset, may repeat after 2 h (max 25 mg/d)
Eletriptan	Relpax	40 or 80 mg
Zolmitriptan	Zomig Zomig Rapimelt	2.5-mg tablet at onset; may repeat after 2 h (max 10 mg/d)
Nasal		
Dihydroergotamine	Migranal Nasal Spray	Prior to nasal spray, the pump must be primed 4 times; 1 spray (0.5 mg) is administered, followed in 15 min by a second spray
Sumatriptan	Imitrex Nasal Spray	5–20 mg intranasal spray as 4 sprays of 5 mg or a single 20 mg spray (may repeat once after 2 h, not to exceed a dose of 40 mg/d)
Zolmitriptan	Zomig	5 mg intranasal spray as one spray (may repeat once after 2 h, not to exceed a dose of 10 mg/d)
Parenteral		
Dihydroergotamine	DHE-45	1 mg IV, IM, or SC at onset and q1h (max 3 mg/d, 6 mg per week)
Sumatriptan	Imitrex Injection Alsuma Sumavel DosePro	6 mg SC at onset (may repeat once after 1 h for max of 2 doses in 24 h)
Dopamine Receptor Antagonists		
Oral		
Metoclopramide	Reglan, ^a generic ^a	5–10 mg/d
Prochlorperazine	Compazine, ^a generic ^a	1–25 mg/d
Parenteral		
Chlorpromazine	Generic ^a	0.1 mg/kg IV at 2 mg/min; max 35 mg/d
Metoclopramide	Reglan, ^a generic	10 mg IV
Prochlorperazine	Compazine, ^a generic ^a	10 mg IV
Other		
Oral		
Acetaminophen, 325 mg, <i>plus</i> dichloralphenazone, 100 mg, <i>plus</i> isometheptene, 65 mg	Midrin, generic	Two capsules at onset followed by 1 capsule q1h (max 5 capsules)
Nasal		
Butorphanol	Generic	1 mg (1 spray in 1 nostril), may repeat if necessary in 1–2 h
Parenteral		
Opioids	Generic ^a	Multiple preparations and dosages; see Table 18-1

^aNot all drugs are specifically indicated by the FDA for migraine. Local regulations and guidelines should be consulted.

Note: Antiemetics (e.g., domperidone 10 mg or ondansetron 4 or 8 mg) or prokinetics (e.g., metoclopramide 10 mg) are sometimes useful adjuncts.

Abbreviations: 5-HT, 5-hydroxytryptamine; NSAIDs, nonsteroidal anti-inflammatory drugs.

enhance gastric absorption. In addition, dopamine receptor antagonists decrease nausea/vomiting and restore normal gastric motility.

Parenteral Dopamine receptor antagonists (e.g., chlorpromazine, prochlorperazine, metoclopramide) by injection can also provide significant acute relief of migraine; they can be used in combination with parenteral 5-HT_{1B/1D} receptor agonists. A common IV protocol used for

the treatment of severe migraine is the administration over 2 min of a mixture of 5 mg of prochlorperazine and 0.5 mg of dihydroergotamine.

OTHER MEDICATIONS FOR ACUTE MIGRAINE

Oral The combination of acetaminophen, dichloralphenazone, and isometheptene, one to two capsules, has been classified by the FDA

TABLE 447-5 CLINICAL STRATIFICATION OF ACUTE SPECIFIC MIGRAINE TREATMENTS

Clinical Situation	Treatment Options
Failed NSAIDs/analgesics	First tier
	Sumatriptan 50 mg or 100 mg PO
	Almotriptan 12.5 mg PO
	Rizatriptan 10 mg PO
	Eletriptan 40 mg PO
	Zolmitriptan 2.5 mg PO
	Slower effect/better tolerability
	Naratriptan 2.5 mg PO
	Frovatriptan 2.5 mg PO
	Infrequent headache
	Ergotamine/cafeine 1–2/100 mg PO
	Dihydroergotamine nasal spray 2 mg
Early nausea or difficulties taking tablets	Zolmitriptan 5 mg nasal spray
	Sumatriptan 20 mg nasal spray
	Rizatriptan 10 mg MLT wafer
Headache recurrence	Ergotamine 2 mg (most effective PR/ usually with caffeine)
	Naratriptan 2.5 mg PO
	Almotriptan 12.5 mg PO
	Eletriptan 40 mg
Tolerating acute treatments poorly	Naratriptan 2.5 mg
	Almotriptan 12.5 mg
Early vomiting	Zolmitriptan 5 mg nasal spray
	Sumatriptan 25 mg PR
	Sumatriptan 6 mg SC
Menses-related headache	Prevention
	Ergotamine PO at night
	Estrogen patches
	Treatment
	Triptans
Very rapidly developing symptoms	Dihydroergotamine nasal spray
	Zolmitriptan 5 mg nasal spray
	Sumatriptan 6 mg SC
	Dihydroergotamine 1 mg IM

Abbreviation: NSAIDs, nonsteroidal anti-inflammatory drugs.

as “possibly” effective in the treatment of migraine. Because the clinical studies demonstrating the efficacy of this combination analgesic in migraine predated the clinical trial methodologies used with the triptans, it is difficult to compare the efficacy of this sympathomimetic compound to other agents.

Nasal A nasal preparation of butorphanol is available for the treatment of acute pain. As with all opioids, the use of nasal butorphanol has little role in migraine treatment.

Parenteral Opioids are modestly effective in the acute treatment of migraine. For example, IV meperidine (50–100 mg) is given frequently in the emergency room. This regimen “works” in the sense that the pain of migraine is eliminated. However, this regimen is clearly suboptimal for patients with recurrent headache. Opioids do not treat the underlying headache mechanism; rather, they act to alter the pain sensation, and there is evidence their use may decrease the likelihood of a response to triptans in the future. Moreover, in patients taking oral opioids, such as oxycodone or hydrocodone, habituation or addiction can greatly confuse the treatment of migraine. Opioid craving and/or withdrawal can aggravate and accentuate migraine. Therefore, it is recommended that opioid use in migraine be limited to patients with severe, but infrequent, headaches that are unresponsive to

other pharmacologic approaches or who have contraindications to other therapies.

MEDICATION-OVERUSE HEADACHE

Acute attack medications, particularly opioid or barbiturate-containing compound analgesics, have a propensity to aggravate headache frequency and induce a state of refractory daily or near-daily headache called *medication-overuse headache*. This condition is likely not a separate headache entity but a reaction of the migraine patient to a particular medicine. Migraine patients who have two or more headache days a week should be cautioned about frequent analgesic use (see “Chronic Daily Headache” in Chap. 21).

PREVENTIVE TREATMENTS FOR MIGRAINE

Patients with an increasing frequency of migraine attacks or with attacks that are either unresponsive or poorly responsive to abortive treatments are good candidates for preventive agents. In general, a preventive medication should be considered in the subset of patients with four or more attacks a month. Significant side effects are associated with the use of many of these agents; furthermore, determination of dose can be difficult because the recommended doses have been derived for conditions other than migraine. The mechanism of action of these drugs is unclear; it seems likely that the brain sensitivity that underlies migraine is modified. Patients are usually started on a low dose of a chosen treatment; the dose is then gradually increased, up to a reasonable maximum, to achieve clinical benefit.

Drugs that have the capacity to stabilize migraine are listed in Table 447-6. Drugs must be taken daily, and there is usually a lag of between 2 to 12 weeks before an effect is seen. The drugs that have been approved by the FDA for the prophylactic treatment of migraine include propranolol, timolol, sodium valproate, topiramate, and methysergide (not available). In addition, a number of other drugs appear to display prophylactic efficacy. This group includes amitriptyline, nortriptyline, flunarizine, phenelzine, gabapentin, and cyproheptadine. Placebo-controlled trials of onabotulinum toxin type A in episodic migraine were negative, whereas, overall, placebo-controlled trials in chronic migraine were positive. Phenelzine and methysergide are usually reserved for recalcitrant cases because of their serious potential side effects. Phenelzine is a monoamine oxidase inhibitor (MAOI); therefore, tyramine-containing foods, decongestants, and meperidine are contraindicated. Methysergide may cause retroperitoneal or cardiac valvular fibrosis when it is used for >6 months, and thus monitoring is required for patients using this drug; the risk of fibrosis is about 1:1500 and is likely to reverse after the drug is stopped.

The probability of success with any one of the antimigraine drugs is 50–75%. Many patients are managed adequately with low-dose amitriptyline, propranolol, candesartan, topiramate, or valproate. If these agents fail or lead to unacceptable side effects, second-line agents such as methysergide or phenelzine can be used. Once effective stabilization is achieved, the drug is continued for ~6 months and then slowly tapered to assess the continued need. Many patients are able to discontinue medication and experience fewer and milder attacks for long periods, suggesting that these drugs may alter the natural history of migraine.

TENSION-TYPE HEADACHE

Clinical Features The term *tension-type headache* (TTH) is commonly used to describe a chronic head-pain syndrome characterized by bilateral tight, band-like discomfort. The pain typically builds slowly, fluctuates in severity, and may persist more or less continuously for many days. The headache may be episodic or chronic (present >15 days per month).

A useful clinical approach is to diagnose TTH in patients whose headaches are completely without accompanying features such as nausea, vomiting, photophobia, phonophobia, osmophobia, throbbing, and aggravation with movement. Such an approach neatly separates

TABLE 447-6 PREVENTIVE TREATMENTS IN MIGRAINE^a

Drug	Dose	Selected Side Effects
Pizotifen ^b	0.5–2 mg qd	Weight gain Drowsiness
Beta blocker		
Propranolol	40–120 mg bid	Reduced energy
Metoprolol	25–100 mg bid	Tiredness Postural symptoms Contraindicated in asthma
Antidepressants		
Amitriptyline	10–75 mg at night	Drowsiness
Doxepin	25–75 mg at night	
Nortriptyline	25–75 mg at night	Note: Some patients may only need a total dose of 10 mg, although generally 1–1.5 mg/kg body weight is required
Venlafaxine	75–150 mg/d	
Anticonvulsants		
Topiramate	25–200 mg/d	Paresthesias Cognitive symptoms Weight loss Glaucoma Caution with nephrolithiasis
Valproate	400–600 mg bid	Drowsiness Weight gain Tremor Hair loss Fetal abnormalities Hematologic or liver abnormalities
Serotonergic drugs		
Methysergide ^c	1–4 mg qd	Drowsiness Leg cramps Hair loss Retroperitoneal fibrosis (1-month drug holiday is required every 6 months)
Other classes		
Flunarizine ^b	5–15 mg qd	Drowsiness Weight gain Depression Parkinsonism
Candesartan	16 mg daily	Dizziness
Chronic migraine		
Onabotulinum toxin type A	155 U	Loss of brow furrow
No convincing evidence from controlled trials		
Verapamil		
Controlled trials demonstrate <i>no effect</i>		
Nimodipine		
Clonidine		
Selective serotonin reuptake inhibitors: fluoxetine		

^aCommonly used preventives are listed with typical doses and common side effects. Not all listed medicines are approved by the U.S. Food and Drug Administration; local regulations and guidelines should be consulted. ^bNot available in the United States. ^cNot currently available worldwide.

migraine, which has one or more of these features and is the main differential diagnosis, from TTH. The International Headache Society's main definition of TTH allows an admixture of nausea, photophobia, or phonophobia in various combinations, although the appendix definition does not; this illustrates the difficulty in distinguishing these two clinical entities. In clinical practice, dichotomizing patients on the basis of the presence of associated features (migraine) and the absence of associated features (TTH) is highly recommended. Indeed patients whose headaches fit the TTH phenotype and who have migraine at other times, along with a family history of migraine, migrainous illnesses of childhood, or typical migraine triggers to their migraine attacks, may be biologically different from those who have TTH headache with none of the features. TTH may be infrequent (episodic) or occur on 15 days or more a month (chronic).

Pathophysiology The pathophysiology of TTH is incompletely understood. It seems likely that TTH is due to a primary disorder of central nervous system pain modulation alone, unlike migraine, which involves a more generalized disturbance of sensory modulation. Data suggest a genetic contribution to TTH, but this may not be a valid finding; given the current diagnostic criteria, the studies undoubtedly included many migraine patients. The name *tension-type headache* implies that pain is a product of *nervous tension*, but there is no clear evidence for tension as an etiology. Muscle contraction has been considered to be a feature that distinguishes TTH from migraine, but there appear to be no differences in contraction between the two headache types.

TREATMENT TENSION-TYPE HEADACHE

The pain of TTH can generally be managed with simple analgesics such as acetaminophen, aspirin, or NSAIDs. Behavioral approaches including relaxation can also be effective. Clinical studies have demonstrated that triptans in pure TTH are not helpful, although triptans are effective in TTH when the patient also has migraine. For chronic TTH, amitriptyline is the only proven treatment (Table 447-6); other tricyclics, selective serotonin reuptake inhibitors, and the benzodiazepines have not been shown to be effective. There is no evidence for the efficacy of acupuncture. Placebo-controlled trials of onabotulinum toxin type A in chronic TTH were negative.

TRIGEMINAL AUTONOMIC CEPHALALGIAS, INCLUDING CLUSTER HEADACHE

The trigeminal autonomic cephalalgias (TACs) describe a grouping of primary headaches including cluster headache, paroxysmal hemicrania, SUNCT (short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing)/SUNA (short-lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms), and hemicrania continua (Table 447-1). TACs are characterized by relatively short-lasting attacks of head pain associated with cranial autonomic symptoms, such as lacrimation, conjunctival injection, or nasal congestion (Table 447-7). Pain is usually severe and may occur more than once a day. Because of the associated nasal congestion or rhinorrhea, patients are often misdiagnosed with “sinus headache” and treated with decongestants, which are ineffective.

TACs must be differentiated from short-lasting headaches that do not have prominent cranial autonomic syndromes, notably trigeminal neuralgia, primary stabbing headache, and hypnic headache. The cycling pattern and length, frequency, and timing of attacks are useful in classifying patients. Patients with TACs should undergo pituitary imaging and pituitary function tests because there is an excess of TAC presentations in patients with pituitary tumor-related headache.

Cluster Headache Cluster headache is a relatively rare form of primary headache with a population frequency of approximately 0.1%. The pain is deep, usually retroorbital, often excruciating in intensity, non-fluctuating, and explosive in quality. A core feature of cluster headache is periodicity. At least one of the daily attacks of pain recurs at about the same hour each day for the duration of a cluster bout. The typical cluster headache patient has daily bouts of one to two attacks of

TABLE 447-7 CLINICAL FEATURES OF THE TRIGEMINAL AUTONOMIC CEPHALALGIAS

	Cluster Headache	Paroxysmal Hemicrania	SUNCT/SUNA
Gender	M > F	F = M	F ~ M
Pain			
Type	Stabbing, boring	Throbbing, boring, stabbing	Burning, stabbing, sharp
Severity	Excruciating	Excruciating	Severe to excruciating
Site	Orbit, temple	Orbit, temple	Periorbital
Attack frequency	1/alternate day–8/d	1–20/d (>5/d for more than half the time)	3–200/d
Duration of attack	15–180 min	2–30 min	5–240 s
Autonomic features	Yes	Yes	Yes (prominent conjunctival injection and lacrimation) ^a
Migrainous features ^b	Yes	Yes	Yes
Alcohol trigger	Yes	No	No
Cutaneous triggers	No	No	Yes
Indomethacin effect	—	Yes ^c	—
Abortive treatment	Sumatriptan injection or nasal spray Oxygen	No effective treatment	Lidocaine (IV)
Prophylactic treatment	Verapamil Methysergide Lithium	Indomethacin	Lamotrigine Topiramate Gabapentin

^aIf conjunctival injection and tearing are not present, consider SUNA. ^bNausea, photophobia, or phonophobia; photophobia and phonophobia are typically unilateral on the side of the pain. ^cIndicates complete response to indomethacin.

Abbreviations: SUNA, short-lasting unilateral neuralgiform headache attacks with cranial autonomic features; SUNCT, short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing.

relatively short-duration unilateral pain for 8 to 10 weeks a year; this is usually followed by a pain-free interval that averages a little less than 1 year. Cluster headache is characterized as chronic when there is less than 1 month of sustained remission without treatment. Patients are generally perfectly well between episodes. Onset is nocturnal in about 50% of patients, and men are affected three times more often than women. Patients with cluster headache tend to move about during attacks, pacing, rocking, or rubbing their head for relief; some may even become aggressive during attacks. This is in sharp contrast to patients with migraine, who prefer to remain motionless during attacks.

Cluster headache is associated with ipsilateral symptoms of cranial parasympathetic autonomic activation: conjunctival injection or lacrimation, rhinorrhea or nasal congestion, or cranial sympathetic dysfunction such as ptosis. The sympathetic deficit is peripheral and likely to be due to parasympathetic activation with injury to ascending sympathetic fibers surrounding a dilated carotid artery as it passes into the cranial cavity. When present, photophobia and phonophobia are far more likely to be unilateral and on the same side of the pain, rather than bilateral, as is seen in migraine. This phenomenon of unilateral photophobia/phonophobia is characteristic of TACs. Cluster headache is likely to be a disorder involving central pacemaker neurons in the posterior hypothalamic region (Fig. 447-3).

TREATMENT CLUSTER HEADACHE

The most satisfactory treatment is the administration of drugs to prevent cluster attacks until the bout is over. However, treatment of acute attacks is required for all cluster headache patients at some time.

ACUTE ATTACK TREATMENT

Cluster headache attacks peak rapidly, and thus a treatment with quick onset is required. Many patients with acute cluster headache respond very well to oxygen inhalation. This should be given as 100% oxygen at 10–12 L/min for 15–20 min. It appears that high flow and high oxygen content are important. Sumatriptan 6 mg SC is rapid in onset and will usually shorten an attack to 10–15 min; there is no evidence of tachyphylaxis. Sumatriptan (20 mg) and zolmitriptan (5 mg) nasal sprays are both effective in acute cluster

headache, offering a useful option for patients who may not wish to self-inject daily. Oral sumatriptan is not effective for prevention or for acute treatment of cluster headache.

PREVENTIVE TREATMENTS (TABLE 447-8)

The choice of a preventive treatment in cluster headache depends in part on the length of the bout. Patients with long bouts or those with chronic cluster headache require medicines that are safe when taken for long periods. For patients with relatively short bouts, limited courses of oral glucocorticoids or methysergide (not available in the United States) can be very useful. A 10-day course of prednisone, beginning at 60 mg daily for 7 days and followed by a rapid taper, may interrupt the pain bout for many patients. Lithium (400–800 mg/d) appears to be particularly useful for the chronic form of the disorder.

Many experts favor verapamil as the first-line preventive treatment for patients with chronic cluster headache or prolonged bouts. While verapamil compares favorably with lithium in practice, some patients require verapamil doses far in excess of those administered for cardiac disorders. The initial dose range is 40–80 mg twice daily; effective doses may be as high as 960 mg/d. Side effects such as constipation and leg swelling can be problematic. Of paramount concern, however, is the cardiovascular safety of verapamil, particularly

TABLE 447-8 PREVENTIVE MANAGEMENT OF CLUSTER HEADACHE

Short-Term Prevention	Long-Term Prevention
	Episodic Cluster Headache and Prolonged Chronic Cluster Headache
Prednisone 1 mg/kg up to 60 mg qd, tapering over 21 days	Verapamil 160–960 mg/d
Methysergide 3–12 mg/d	Lithium 400–800 mg/d
Verapamil 160–960 mg/d	Methysergide ^a 3–12 mg/d
Greater occipital nerve injection	Topiramate ^b 100–400 mg/d
	Gabapentin ^b 1200–3600 mg/d
	Melatonin ^b 9–12 mg/d

^aNot available worldwide. ^bUnproven but of potential benefit.

at high doses. Verapamil can cause heart block by slowing conduction in the atrioventricular node, a condition that can be monitored by following the PR interval on a standard electrocardiogram (ECG). Approximately 20% of patients treated with verapamil develop ECG abnormalities, which can be observed with doses as low as 240 mg/d; these abnormalities can worsen over time in patients on stable doses. A baseline ECG is recommended for all patients. The ECG is repeated 10 days after a dose change in patients whose dose is being increased above 240 mg daily. Dose increases are usually made in 80-mg increments. For patients on long-term verapamil, ECG monitoring every 6 months is advised.

NEUROSTIMULATION THERAPY

When medical therapies fail in chronic cluster headache, neurostimulation strategies can be used. Deep-brain stimulation of the region of the posterior hypothalamic gray matter has proven successful in a substantial proportion of patients, although its risk-benefit ratio makes it inappropriate with so many other options now available. Favorable results have also been reported with the less-invasive approach of occipital nerve stimulation, with sphenopalatine ganglion stimulation and with a noninvasive vagal nerve stimulator.

PAROXYSMAL HEMICRANIA

Paroxysmal hemicrania (PH) is characterized by frequent unilateral, severe, short-lasting episodes of headache. Like cluster headache, the pain tends to be retroorbital but may be experienced all over the head and is associated with autonomic phenomena such as lacrimation and nasal congestion. Patients with remissions are said to have episodic PH, whereas those with the nonremitting form are said to have chronic PH. The essential features of PH are unilateral, very severe pain; short-lasting attacks (2–45 min); very frequent attacks (usually more than five a day); marked autonomic features ipsilateral to the pain; rapid course (<72 h); and excellent response to indomethacin. In contrast to cluster headache, which predominantly affects males, the male-to-female ratio in PH is close to 1:1.

Indomethacin (25–75 mg tid), which can completely suppress attacks of PH, is the treatment of choice. Although therapy may be complicated by indomethacin-induced gastrointestinal side effects, currently there are no consistently effective alternatives. Topiramate is helpful in some cases. Piroxicam has been used, although it is not as effective as indomethacin. Verapamil, an effective treatment for cluster headache, does not appear to be useful for PH. In occasional patients, PH can coexist with trigeminal neuralgia (PH-tic syndrome); similar to cluster-tic syndrome, each component may require separate treatment.

Secondary PH has been reported with lesions in the region of the sella turcica, including arteriovenous malformation, cavernous sinus meningioma, pituitary pathology and epidermoid tumors. Secondary PH is more likely if the patient requires high doses (>200 mg/d) of indomethacin. In patients with apparent bilateral PH, raised cerebrospinal fluid (CSF) pressure should be suspected. It is important to note that indomethacin reduces CSF pressure. When a diagnosis of PH is considered, magnetic resonance imaging (MRI) is indicated to exclude a pituitary lesion.

SUNCT/SUNA

SUNCT (short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing) is a rare primary headache syndrome characterized by severe, unilateral orbital or temporal pain that is stabbing or throbbing in quality. Diagnosis requires at least 20 attacks, lasting for 5–240 s; ipsilateral conjunctival injection and lacrimation should be present. In some patients, conjunctival injection or lacrimation is missing, and the diagnosis of SUNA (short-lasting unilateral neuralgiform headache attacks with cranial autonomic symptoms) can be made.

DIAGNOSIS The pain of SUNCT/SUNA is unilateral and may be located anywhere in the head. Three basic patterns can be seen: single stabs, which are usually short-lived; groups of stabs; or a longer attack

comprising many stabs between which the pain does not completely resolve, thus giving a “saw-tooth” phenomenon with attacks lasting many minutes. Each pattern may be seen in the context of an underlying continuous head pain. Characteristics that lead to a suspected diagnosis of SUNCT are the cutaneous (or other) triggers of attacks, a lack of refractory period to triggering between attacks, and the lack of a response to indomethacin. Apart from trigeminal sensory disturbance, the neurologic examination is normal in primary SUNCT.

The diagnosis of SUNCT/SUNA is often confused with trigeminal neuralgia (TN) particularly in first-division TN (**Chap. 455**). Minimal or no cranial autonomic symptoms and a clear refractory period to triggering indicate a diagnosis of TN.

SECONDARY (SYMPTOMATIC) SUNCT SUNCT can be seen with posterior fossa or pituitary lesions. All patients with SUNCT/SUNA should be evaluated with pituitary function tests and a brain MRI with pituitary views.

TREATMENT SUNCT/SUNA

ABORTIVE THERAPY

Therapy of acute attacks is not a useful concept in SUNCT/SUNA because the attacks are of such short duration. However, IV lidocaine, which arrests the symptoms, can be used in hospitalized patients.

PREVENTIVE THERAPY

Long-term prevention to minimize disability and hospitalization is the goal of treatment. The most effective treatment for prevention is lamotrigine, 200–400 mg/d. Topiramate and gabapentin may also be effective. Carbamazepine, 400–500 mg/d, has been reported by patients to offer modest benefit.

Surgical approaches such as microvascular decompression or destructive trigeminal procedures are seldom useful and often produce long-term complications. Greater occipital nerve injection has produced limited benefit in some patients. Occipital nerve stimulation is probably helpful in a subgroup of these patients. Complete control with deep-brain stimulation of the posterior hypothalamic region was reported in a single patient. For intractable cases, short-term prevention with IV lidocaine can be effective, as can occipital nerve stimulation.

Hemicrania Continua The essential features of hemicrania continua are moderate and continuous unilateral pain associated with fluctuations of severe pain; complete resolution of pain with indomethacin; and exacerbations that may be associated with autonomic features, including conjunctival injection, lacrimation, and photophobia on the affected side. The age of onset ranges from 11 to 58 years; women are affected twice as often as men. The cause is unknown.

TREATMENT HEMICRANIA CONTINUA

Treatment consists of indomethacin; other NSAIDs appear to be of little or no benefit. The IM injection of 100 mg of indomethacin has been proposed as a diagnostic tool, and administration with a placebo injection in a blinded fashion can be very useful diagnostically. Alternatively, a trial of oral indomethacin, starting with 25 mg tid, then 50 mg tid, and then 75 mg tid, can be given. Up to 2 weeks at the maximal dose may be necessary to assess whether a dose has a useful effect. Topiramate can be helpful in some patients. Occipital nerve stimulation probably has a role in patients with hemicrania continua who are unable to tolerate indomethacin.

OTHER PRIMARY HEADACHES

Primary Cough Headache Primary cough headache is a generalized headache that begins suddenly, lasts for several minutes, sometimes

up to a few hours, and is precipitated by coughing; it is preventable by avoiding coughing or other precipitating events, which can include sneezing, straining, laughing, or stooping. In all patients with this syndrome, serious etiologies must be excluded before a diagnosis of “benign” primary cough headache can be established. A Chiari malformation or any lesion causing obstruction of CSF pathways or displacing cerebral structures can be the cause of the head pain. Other conditions that can present with cough or exertional headache as the initial symptom include cerebral aneurysm, carotid stenosis, and vertebrobasilar disease. Benign cough headache can resemble benign exertional headache (below), but patients with the former condition are typically older.

TREATMENT PRIMARY COUGH HEADACHE

Indomethacin 25–50 mg two to three times daily is the treatment of choice. Some patients with cough headache obtain complete cessation of their attacks with lumbar puncture; this is a simple option when compared to prolonged use of indomethacin, and it is effective in about one-third of patients. The mechanism of this response is unclear.

Primary Exercise Headache Primary exertional headache has features resembling both cough headache and migraine. It may be precipitated by any form of exercise; it often has the pulsatile quality of migraine. The pain, which can last from 5 min to 24 h, is bilateral and throbbing at onset; migrainous features may develop in patients susceptible to migraine. The duration tends to be shorter in adolescents than in older adults. Primary exertional headache can be prevented by avoiding excessive exertion, particularly in hot weather or at high altitude.

The mechanism of primary exertional headache is unclear. Acute venous distension likely explains one syndrome—the acute onset of headache with straining and breath holding, as in weightlifter’s headache. Because exertion can result in headache in a number of serious underlying conditions, these must be considered in patients with exertional headache. Pain from angina may be referred to the head, probably by central connections of vagal afferents, and may present as exertional headache (cardiac cephalgia). The link to exercise is the main clinical clue that headache is of cardiac origin. Pheochromocytoma may occasionally cause exertional headache. Intracranial lesions and stenosis of the carotid arteries are other possible etiologies.

TREATMENT PRIMARY EXERTIONAL HEADACHE

Exercise regimens should begin modestly and progress gradually to higher levels of intensity. Indomethacin at daily doses from 25 to 150 mg is generally effective in benign exertional headache. Indomethacin (50 mg), ergotamine (1 mg orally), dihydroergotamine (2 mg by nasal spray), and methysergide (1–2 mg orally given 30–45 min before exercise) are useful prophylactic measures.

Primary Headache Associated with Sexual Activity Three types of sex headache are reported: a dull bilateral ache in the head and neck that intensifies as sexual excitement increases; a sudden, severe, explosive headache occurring at orgasm; and a postural headache developing after coitus that resembles the headache of low CSF pressure. The last arises from vigorous sexual activity and is a form of low CSF pressure headache (Chap. 21). Headaches developing at the time of orgasm are not always benign; 5–12% of cases of subarachnoid hemorrhage are precipitated by sexual intercourse. Sex headache is reported by men more often than women and may occur at any time during the years of sexual activity. It may develop on several occasions in succession and then not trouble the patient again, even without an obvious change in sexual activity. In patients who stop sexual activity when headache is first noticed, the pain may subside within a period of 5 min to 2 h. In about half of patients, sex headache will subside within 6 months.

About half of patients with sex headache have a history of exertional headaches, but there is no excess of cough headache. Migraine is probably more common in patients with sex headache.

TREATMENT PRIMARY SEX HEADACHE

Benign sex headaches recur irregularly and infrequently. Management can often be limited to reassurance and advice about ceasing sexual activity if a mild, warning headache develops. Propranolol can be used to prevent headache that recurs regularly or frequently, but the dosage required varies from 40 to 200 mg/d. An alternative is the calcium channel-blocking agent diltiazem, 60 mg tid. Ergotamine (1 mg) or indomethacin (25–50 mg) taken 30–45 min prior to sexual activity can also be helpful.

Primary Thunderclap Headache Sudden onset of severe headache may occur in the absence of any known provocation. The differential diagnosis includes the sentinel bleed of an intracranial aneurysm, cervicocephalic arterial dissection, and cerebral venous thrombosis. Headaches of explosive onset may also be caused by the ingestion of sympathomimetic drugs or of tyramine-containing foods in a patient who is taking MAOIs, or they may be a symptom of pheochromocytoma. Whether thunderclap headache can be the presentation of an unruptured cerebral aneurysm is uncertain. When neuroimaging studies and lumbar puncture exclude subarachnoid hemorrhage, patients with thunderclap headache usually do very well over the long term. In one study of patients whose computed tomography (CT) scans and CSF findings were negative, ~15% had recurrent episodes of thunderclap headache, and nearly half subsequently developed migraine or TTH.

The first presentation of any sudden-onset severe headache should be diligently investigated with neuroimaging (CT or, when possible, MRI with MR angiography) and CSF examination. Formal cerebral angiography should be reserved for those cases in which no primary diagnosis is forthcoming and for clinical situations that are particularly suggestive of intracranial aneurysm. Reversible segmental cerebral vasoconstriction may be seen in primary thunderclap headache without an intracranial aneurysm. In the presence of posterior leukoencephalopathy, the differential diagnosis includes cerebral angitis, drug toxicity (cyclosporine, intrathecal methotrexate/cytarabine, pseudoephedrine, or cocaine), posttransfusion effects, and postpartum angiopathy. Treatment with nimodipine may be helpful, although by definition, the vasoconstriction of primary thunderclap headache resolves spontaneously.

Cold-Stimulus Headache This refers to head pain triggered by application or ingestion/inhalation of something cold. It is bought on quickly and typically resolves within 10–30 min of the stimulus being removed. It is best recognized as “brain-freeze” headache or ice-cream headache when due to ingestion. Although cold may be uncomfortable at some level for many people, it is the reliable, severe, and somewhat prolonged nature of these pains that set them apart. The transient receptor potential cation subfamily M member 8 (TRPM8) channel, a known cold temperature sensor, may be a mediator of this syndrome.

External Pressure Headache External pressure from compression or traction on the head can produce a pain that may have some generalized component, although the pain is largely focused around the site of the pressure. It typically resolves within an hour of the stimulus being removed. Examples of stimuli include helmets, swimming goggles, or very long ponytails. Treatment is to recognize the problem and remove the stimulus.

Primary Stabbing Headache The essential features of primary stabbing headache are stabbing pain confined to the head or, rarely, the face, lasting from 1 to many seconds or minutes and occurring as a

2598 single stab or a series of stabs; absence of associated cranial autonomic features; absence of cutaneous triggering of attacks; and a pattern of recurrence at irregular intervals (hours to days). The pains have been variously described as “ice-pick pains” or “jabs and jolts.” They are more common in patients with other primary headaches, such as migraine, the TACs, and hemicrania continua.

TREATMENT PRIMARY STABBING HEADACHE

The response of primary stabbing headache to indomethacin (25–50 mg two to three times daily) is usually excellent. As a general rule, the symptoms wax and wane, and after a period of control on indomethacin, it is appropriate to withdraw treatment and observe the outcome.

Nummular Headache Nummular headache is felt as a round or elliptical discomfort that is fixed in place, ranges in size from 1–6 cm, and may be continuous or intermittent. Uncommonly it may be multifocal. It may be episodic but is more often continuous during exacerbations. Accompanying the pain there may be a local sensory disturbance, such as allodynia or hypesthesia. Local dermatologic or bony lesions need to be excluded by examination and investigation. This condition can be difficult to treat; tricyclics, such as amitriptyline, or anticonvulsants, such as topiramate or valproate, are most often tried.

Hypnic Headache This headache syndrome typically begins a few hours after sleep onset. The headaches last from 15 to 30 min and are typically moderately severe and generalized, although they may be unilateral and can be throbbing. Patients may report falling back to sleep only to be awakened by a further attack a few hours later; up to three repetitions of this pattern occur through the night. Daytime naps can also precipitate head pain. Most patients are female, and the onset is usually after age 60 years. Headaches are bilateral in most, but may be unilateral. Photophobia, phonophobia, and nausea are usually absent. The major secondary consideration in this headache type is poorly controlled hypertension; 24-h blood pressure monitoring is recommended to detect this treatable condition.

TREATMENT HYPNIC HEADACHE

Patients with hypnic headache generally respond to a bedtime dose of lithium carbonate (200–600 mg). For those intolerant of lithium, verapamil (160 mg) or methysergide (1–4 mg at bedtime) may be alternative strategies. One to two cups of coffee or caffeine, 60 mg orally, at bedtime may be effective in approximately one-third of patients. Case reports also suggest that flunarizine, 5 mg nightly, can be effective.

New Daily Persistent Headache Primary new daily persistent headache (NDPH) occurs in both males and females. It can be of the migrainous type, with features of migraine, or it can be featureless, appearing as new-onset TTH. Migrainous features are common and include unilateral headache and throbbing pain; each feature is present in about one-third of patients. Nausea, photophobia, and/or phonophobia occur in about half of patients. Some patients have a previous history of migraine; however, the proportion of NDPH sufferers with preexisting migraine is no greater than the frequency of migraine in the general population. At 24 months, ~86% of patients are headache-free. Treatment of migrainous-type primary NDPH consists of using the preventive therapies effective in migraine (see above). Featureless NDPH is one of the primary headache forms most refractory to treatment. Standard preventive therapies can be offered but are often ineffective. The secondary NDPHs are discussed elsewhere ([Chap. 21](#)).

448 Alzheimer's Disease and Other Dementias

William W. Seeley, Bruce L. Miller

ALZHEIMER'S DISEASE

Approximately 10% of all persons over the age of 70 years have significant memory loss, and in more than half, the cause is Alzheimer's disease (AD). It is estimated that the median annual total cost of caring for a single patient with advanced AD is >\$50,000, while the emotional toll for family members and caregivers is immeasurable. AD can manifest as young as the third decade, but it is the most common cause of dementia in the elderly. Patients most often present with an insidious loss of episodic memory followed by a slowly progressive dementia that evolves over years. In typical amnesic AD, brain imaging reveals atrophy that begins in the medial temporal lobes before spreading to lateral and medial parietal and temporal lobes and lateral frontal cortex. Microscopically, there are neuritic plaques containing amyloid beta (A β), neurofibrillary tangles (NFTs) composed of hyperphosphorylated tau filaments, and A β accumulation of in blood vessel walls in cortex and leptomeninges (see “Pathology,” below). The identification of causative mutations and susceptibility genes for AD has provided a foundation for rapid progress in understanding the biological basis of the disorder. The major genetic risk for AD is apolipoprotein ϵ 4 (Apo ϵ 4). Carrying one E4 allele increases the risk for AD by 2- to 3-fold, whereas two alleles increase the risk 16-fold.

CLINICAL MANIFESTATIONS

The cognitive changes of AD tend to follow a characteristic pattern, beginning with memory impairment and progressing to language and visuospatial deficits. Yet, approximately 20% of patients with AD present with nonmemory complaints such as word-finding, organizational, or navigational difficulty. In other patients, upstream visual processing dysfunction (referred to as posterior cortical atrophy syndrome) or a progressive “logopenic” aphasia are the primary manifestations of AD for years before progressing to involve memory and other cognitive domains. Still other patients may present with an asymmetric akinetic-rigid-dystonic (“corticobasal”) syndrome or a dysexecutive “frontal variant” of AD.

In the early stages of typical amnesic AD, the memory loss may go unrecognized or be ascribed to benign forgetfulness of aging. Once the memory loss becomes noticeable to the patient and spouse and falls 1.5 standard deviations below normal on standardized memory tests, the term mild cognitive impairment (MCI) is applied. This construct provides useful prognostic information, because approximately 50% of patients with MCI (roughly 12% per year) will progress to AD over 4 years. Increasingly, the MCI construct is being replaced by the notion of “early symptomatic AD” to signify that AD is considered the underlying disease (based on clinical or biomarker evidence) in a patient who remains functionally compensated. Even earlier in the course, “prodromal AD” refers to a person with biomarker evidence of AD (amyloid imaging positive with positron emission tomography or low cerebrospinal A β_{42} and mildly elevated tau) in the absence of symptoms. These refinements have been developed in anticipation of early-stage treatment and prevention trials that have already begun in humans. New evidence suggests that partial and sometimes generalized seizures herald AD and can occur even prior to dementia onset.

Eventually, with AD, the cognitive problems begin to interfere with daily activities, such as keeping track of finances, following instructions on the job, driving, shopping, and housekeeping. Some patients are unaware of these difficulties (*anosognosia*), but most remain acutely attuned to their deficits. Changes in environment (travel, relocation, hospitalization) tend to destabilize the patient. Over time patients become lost on walks or while driving. Social graces, routine behavior, and superficial conversation may be surprisingly intact, even into the later stages of the illness.

In the middle stages of AD, the patient is unable to work, is easily lost and confused, and requires daily supervision. Language becomes impaired—first naming, then comprehension, and finally fluency. Word-finding difficulties and circumlocution can be evident in the early stages, even when formal testing demonstrates intact naming and fluency. *Apraxia* emerges, and patients have trouble performing learned sequential motor tasks. Visuospatial deficits begin to interfere with dressing, eating, or even walking, and patients fail to solve simple puzzles or copy geometric figures. Simple calculations and clock reading become difficult in parallel.

In the late stages, some persons remain ambulatory, wandering aimlessly. Loss of judgment and reasoning is inevitable. Delusions are common, usually simple, with common themes of theft, infidelity, or misidentification. Approximately 10% of AD patients develop *Capgras' syndrome*, believing that a caregiver has been replaced by an impostor. In contrast to dementia with Lewy bodies (DLB), where *Capgras' syndrome* is an early feature, in AD this syndrome emerges late. Disinhibition and uncharacteristic belligerence may occur and alternate with passivity and withdrawal. Sleep-wake patterns are disrupted, and nighttime wandering becomes disturbing to the household. Some patients develop a shuffling gait with generalized muscle rigidity associated with slowness and awkwardness of movement. Patients often look parkinsonian (Chap. 449) but rarely have a high-amplitude, low-frequency tremor at rest. There is a strong overlap between Parkinson's disease (PD) and AD, and some AD patients develop more classical PD features.

In the end stages, AD patients become rigid, mute, incontinent, and bedridden, and help is needed with eating, dressing, and toileting. Hyperactive tendon reflexes and myoclonic jerks (sudden brief contractions of various muscles or the whole body) may occur spontaneously or in response to physical or auditory stimulation. Often death results from malnutrition, secondary infections, pulmonary emboli, heart disease, or, most commonly, aspiration. The typical duration of AD is 8–10 years, but the course ranges from 1 to 25 years. For unknown reasons, some patients with AD show a steady decline in function while others have prolonged plateaus without major deterioration.

DIFFERENTIAL DIAGNOSIS

Early in the disease course, other etiologies of dementia should be excluded (see Tables 35-1, 35-3, and 35-4). Neuroimaging studies (computed tomography [CT] and magnetic resonance imaging [MRI]) do not show a single specific pattern with AD and may be normal early in the disease. As AD progresses, more distributed but usually posterior-predominant cortical atrophy becomes apparent, along with atrophy of the medial temporal memory structures (see Chap. 35, Fig. 35-1). The main purpose of imaging is to exclude other disorders, such as primary and secondary neoplasms, vascular dementia, diffuse white matter disease, and normal-pressure hydrocephalus (NPH). Imaging also helps to distinguish AD from other degenerative disorders, such as frontotemporal dementia (FTD) or Creutzfeldt-Jacob disease (CJD), which feature distinctive imaging patterns. Functional imaging studies, such as positron emission tomography (PET), reveal hypometabolism in the posterior temporal-parietal cortex in AD (see Fig. 35-1). PET can also be used to detect the presence of fibrillar amyloid in the brain (see Fig. 35-4), and amyloid PET positivity is becoming required for entry into treatment trials for AD. Barriers to interpretation continue, however, to limit the use of amyloid PET in routine clinical evaluation. Although amyloid binding with PET is typical for AD, many asymptomatic healthy older individuals show amyloid uptake, and the likelihood that these individuals will convert to clinical AD is still under study. Similarly, dementia due to a non-AD disorder can be the underlying etiology in a patient who is amyloid positive on imaging. Electroencephalogram (EEG) is normal or shows nonspecific slowing; prolonged EEG can be used to seek out intermittent non-convulsive seizures. Routine spinal fluid examination is also normal. Cerebrospinal fluid (CSF) $A\beta_{42}$ level is reduced, whereas the tau protein is elevated, but the test characteristics of these assays still make interpretation challenging in individual patients. *Slowly progressive decline in*

memory and orientation, normal results on laboratory tests, and an MRI or CT scan showing only distributed or posteriorly predominant cortical and hippocampal atrophy are highly suggestive of AD. A clinical diagnosis of AD reached after careful evaluation is confirmed at autopsy about 90% of the time, with misdiagnosed cases usually representing one of the other dementing disorders described later in this chapter, a mixture of AD with vascular pathology, or DLB.

Simple clinical clues are useful in the differential diagnosis. Early prominent gait disturbance with only mild memory loss suggests vascular dementia or, rarely, NPH (see below). Resting tremor with stooped posture, bradykinesia, and masked facies suggest PD (Chap. 449). When dementia occurs after a well-established diagnosis of PD, PD dementia (PDD) is usually the correct diagnosis, but many patients with this diagnosis will show a mixture of AD and Lewy body disease at autopsy. The early appearance of parkinsonian features in association with fluctuating alertness, visual hallucinations, or delusional misidentification suggests DLB. Chronic alcoholism should prompt the search for vitamin deficiency. Loss of joint position and vibration sensibility accompanied by Babinski signs suggests vitamin B₁₂ deficiency (Chap. 456). Early onset of a focal seizure suggests a metastatic or primary brain neoplasm (Chap. 118). Previous or ongoing depression raises suspicion for depression-related cognitive impairment, although AD can feature a depressive prodrome. A history of treatment for insomnia, anxiety, psychiatric disturbance, or epilepsy suggests chronic drug intoxication. Rapid progression over a few weeks or months associated with rigidity and myoclonus suggests CJD (Chap. 453e). Prominent behavioral changes with intact navigation and focal anterior-predominant atrophy on brain imaging are typical of FTD. A positive family history of dementia suggests either one of the familial forms of AD or one of the other genetic disorders associated with dementia, such as FTD (see below), HD (see below), prion disease (Chap. 453e), or rare hereditary ataxias (Chap. 450).

EPIDEMIOLOGY

The most important risk factors for AD are old age and a positive family history. The prevalence of AD increases with each decade of adult life, reaching 20–40% of the population over the age of 85. A positive family history of dementia suggests a genetic contribution to AD, although autosomal dominant inheritance occurs in only 2% of patients. Female sex is a risk factor independent of the greater longevity of women, and women who carry an Apo $\epsilon 4$ allele are more susceptible than are male $\epsilon 4$ carriers. A history of head trauma with concussion increases the risk for AD. AD is more common in groups with low educational attainment, but education influences test-taking ability, and it is clear that AD can affect persons of all intellectual levels. One study found that the capacity to express complex written language in early adulthood correlated with a decreased risk for AD. Numerous environmental factors, including aluminum, mercury, and viruses, have been proposed as causes of AD, but rigorous studies have failed to demonstrate to a significant role for any of these exposures. Similarly, several studies suggest that the use of nonsteroidal anti-inflammatory agents is associated with a decreased risk of AD, but this risk has not been confirmed in large prospective studies. Vascular disease, and stroke in particular, seems to lower the threshold for the clinical expression of AD. Also, in many patients with AD, amyloid angiopathy can lead to microhemorrhages, large lobar hemorrhages, ischemic infarctions most often in the subcortical white matter, or in rare cases an inflammatory leukoencephalopathy. Diabetes increases the risk of AD threefold. Elevated homocysteine and cholesterol levels; hypertension; diminished serum levels of folic acid; low dietary intake of fruits, vegetables, and red wine; and low levels of exercise are all being explored as potential risk factors for AD.

PATHOLOGY

At autopsy, the earliest and most severe degeneration is usually found in the medial temporal lobe (entorhinal/perirhinal cortex and hippocampus), lateral temporal cortex, and nucleus basalis of Meynert. The characteristic microscopic findings are neuritic plaques and NFTs (Fig. 448-1). These lesions may accumulate in small numbers during

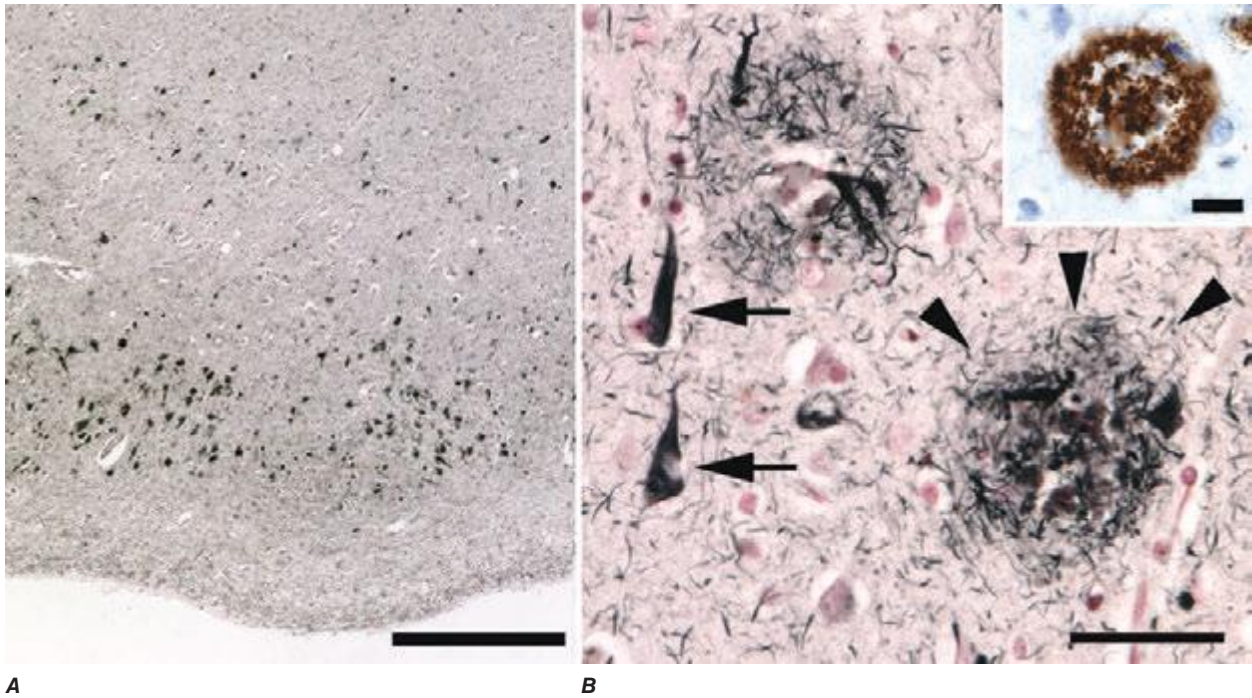
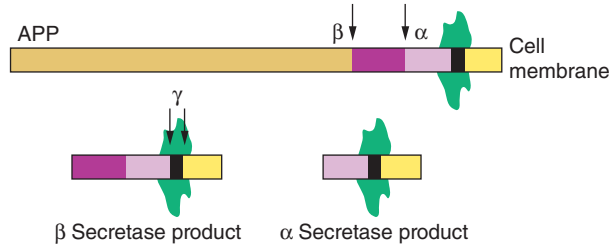


FIGURE 448-1 Neuropathology of Alzheimer's disease. **A.** Early neurofibrillary degeneration, consisting of neurofibrillary tangles and neuropil threads, preferentially affects the medial temporal lobes, especially the stellate pyramidal neurons that compose the layer 2 islands of entorhinal cortex, as shown. **B.** Higher magnification view reveals the fibrillary nature of tangles (*arrows*) and the complex structure of neuritic plaques (*arrowheads*), whose major component is A β (*inset* shows immunohistochemistry for A β). Scale bars are 500 μ m in **A**, 50 μ m in **B**, and 20 μ m in **B** inset.

normal brain aging but dominate the picture in AD. Increasing evidence suggests that soluble amyloid species called *oligomers* may cause cellular dysfunction and represent the early toxic molecule in AD. Eventually, further amyloid polymerization and fibril formation lead to neuritic plaques, which contain a central core of amyloid, proteoglycans, Apo ϵ 4, α -antichymotrypsin, and other proteins. A β is a protein of 39–42 amino acids that is derived proteolytically from a larger transmembrane protein, *amyloid precursor protein* (APP), when APP is cleaved by β and γ secretases (**Fig. 448-2**). The normal

function of the A β peptides remains uncertain. APP has neurotrophic and neuroprotective properties. The plaque core is surrounded by a halo, which contains dystrophic, tau-immunoreactive neurites and activated microglia. The accumulation of A β in cerebral arterioles is termed *amyloid angiopathy*. NFTs are composed of silver-staining neuronal cytoplasmic fibrils composed of abnormally phosphorylated tau protein; they appear as paired helical filaments by electron microscopy. Tau binds to and stabilizes microtubules, supporting axonal transport of organelles, glycoproteins, neurotransmitters, and other important cargoes throughout the neuron. Once hyperphosphorylated, tau can no longer bind properly to microtubules and redistributes from the axon to throughout the neuronal cytoplasm and distal dendrites, compromising function. Finally, patients with AD often show comorbid DLB or vascular pathology. In animal models of AD, diminishing neuronal tau ameliorates the cognitive deficits and seizures, even though A β_{42} continues to accumulate, raising hope for tau-lowering therapies in humans. Biochemically, AD is associated with a decrease in the cortical levels of several proteins and neurotransmitters, especially acetylcholine, its synthetic enzyme choline acetyltransferase, and nicotinic cholinergic receptors. Reduction of acetylcholine reflects degeneration of cholinergic neurons in the nucleus basalis of Meynert that project throughout the cortex. There is also noradrenergic and serotonergic depletion due to degeneration of brainstem nuclei such as the locus coeruleus and dorsal raphe, where tau-immunoreactive neuronal cytoplasmic inclusions can be identified even in individuals lacking entorhinal cortex NFTs.

Step 1: Cleavage by either α or β secretase



Step 2: Cleavage by γ secretase



FIGURE 448-2 Amyloid precursor protein (APP) is catabolized by α , β , and γ secretases. A key initial step is the digestion by either β secretase (BASE) or α secretase (ADAM10 or ADAM17 [TACE]), producing smaller nontoxic products. Cleavage of the β secretase product by γ secretase (Step 2) results in either the toxic A β_{42} or the nontoxic A β_{40} peptide; cleavage of the α secretase product by γ secretase produces the nontoxic P3 peptide. Excess production of A β_{42} is a key initiator of cellular damage in Alzheimer's disease (AD). Therapeutics for AD have focused on attempts to reduce accumulation of A β_{42} by antagonizing β or γ secretases, promoting α secretase, or clearing A β_{42} that has already formed by use of specific antibodies.

GENETIC CONSIDERATIONS

Several genes play an important role in the pathogenesis of AD. One is the *APP* gene on chromosome 21. Adults with trisomy 21 (Down's syndrome) consistently develop the typical neuropathologic hallmarks of AD if they survive beyond age 40 years, and many develop a progressive dementia superimposed on their baseline mental retardation. The extra dose of the *APP* gene on chromosome 21 is the initiating cause of AD in adult Down's syndrome and results in excess cerebral amyloid production. Supporting this hypothesis, some families with early age-of-onset familial AD (FAD) have point

mutations in *APP*. Although very rare, these families were the first examples of single-gene autosomal dominant transmission of AD.

Investigation of large families with multigenerational FAD led to the discovery of two additional AD-causing genes, the *presenilins*. Presenilin-1 (*PS-1*) is on chromosome 14 and encodes a protein called S182. Mutations in this gene cause an early-age-of-onset AD, with onset before the age of 60 and often before age 50, transmitted in an autosomal dominant, highly penetrant fashion. More than 100 different mutations have been found in the *PS-1* gene in families from a wide range of ethnic backgrounds. Presenilin-2 (*PS-2*) is on chromosome 1 and encodes a protein called STM2. A mutation in the *PS-2* gene was first found in a group of American families with Volga German ethnic background. Mutations in *PS-1* are much more common than those in *PS-2*. The presenilins are highly homologous and encode similar proteins that at first appeared to have seven transmembrane domains (hence the designation *STM*), but subsequent studies have suggested eight such domains, with a ninth submembrane region. Both S182 and STM2 are cytoplasmic neuronal proteins that are widely expressed throughout the nervous system. They are homologous to a cell-trafficking protein, sel 12, found in the nematode *Caenorhabditis elegans*. Patients with mutations in the presenilin genes have elevated plasma levels of $A\beta_{42}$, and *PS-1* mutations produce increased $A\beta_{42}$ in the media in cell culture. There is evidence that *PS-1* is involved in the cleavage of APP at the γ secretase site and mutations in either gene (*PS-1* or *APP*) may disturb γ secretase cleavage. Mutations in *PS-1* are the most common cause of early-age-of-onset FAD, representing perhaps 40–70% of all cases. Mutations in *PS-1* tend to produce AD with an earlier age of onset (mean onset 45 years) and a shorter, more rapidly progressive course (mean duration 6–7 years) than the disease caused by mutations in *PS-2* (mean onset 53 years; duration 11 years). Although some carriers of *PS-2* mutations have had onset of dementia after the age of 70, mutations in the presenilins rarely lead to late-age-of-onset AD. Clinical genetic testing for these uncommon mutations is available but likely to be revealing only in early-age-of-onset FAD and should be performed in association with formal genetic counseling.

The Apo ϵ gene on chromosome 19 is involved in the pathogenesis of AD. The protein, apolipoprotein E, participates in cholesterol transport (Chap. 421), and the gene has three alleles: $\epsilon 2$, $\epsilon 3$, and $\epsilon 4$. The Apo $\epsilon 4$ allele confers increased risk of AD in the general population, including sporadic and late-age-of-onset familial forms. Approximately 24–30% of the nondemented white population has at least one $\epsilon 4$ allele (12–15% allele frequency), and about 2% are $\epsilon 4/\epsilon 4$ homozygotes. Among patients with AD, 40–65% have at least one $\epsilon 4$ allele, a highly significant elevation compared with controls. Conversely, many AD patients have no $\epsilon 4$ allele, and $\epsilon 4$ carriers may never develop AD. Therefore, $\epsilon 4$ is neither necessary nor sufficient to cause AD. Nevertheless, the Apo $\epsilon 4$ allele represents the most important genetic risk factor for sporadic AD and acts as a dose-dependent disease modifier, with the earliest age of onset associated with the $\epsilon 4$ homozygosity. Precise mechanisms through which Apo $\epsilon 4$ confers AD risk or hastens onset remain unclear, but $\epsilon 4$ leads to less efficient amyloid clearance and to the production of toxic fragments from cleavage of the molecule. Apo ϵ can be identified in neuritic plaques and may also be involved in neurofibrillary tangle formation, because it binds to tau protein. Apo $\epsilon 4$ decreases neurite outgrowth in dorsal root ganglion neuronal cultures, perhaps indicating a deleterious role in the brain's response to injury. Some evidence suggests that the $\epsilon 2$ allele may reduce AD risk. Use of Apo ϵ testing in AD diagnosis remains controversial. It is not indicated as a predictive test in normal persons because its precise predictive value is unclear, and many individuals with the $\epsilon 4$ allele never develop dementia. Many cognitively normal $\epsilon 4$ heterozygotes and homozygotes show decreased cerebral cortical metabolic function with PET, suggesting presymptomatic abnormalities due to AD or an inherited vulnerability of the AD-targeted network. In demented persons who meet clinical criteria for AD, finding an $\epsilon 4$ allele increases the reliability of diagnosis; however, the absence of an $\epsilon 4$ allele cannot be considered evidence against AD. Furthermore, all patients with dementia, including those with an $\epsilon 4$ allele, require a search for reversible causes of their cognitive impairment. Nevertheless, Apo $\epsilon 4$

remains the single most important biologic marker associated with AD risk, and studies of $\epsilon 4$'s functional role and diagnostic utility are progressing rapidly. The $\epsilon 4$ allele is not associated with risk for FTD, DLB, or CJD, although some evidence suggests that $\epsilon 4$ may exacerbate the phenotype of non-AD degenerative disorders, head trauma, and other brain injuries. Additional genes are also likely to be involved in AD, especially as minor risk alleles for sporadic forms of the disease. Genome-wide association studies have implicated the clusterin (*CLU*), phosphatidylinositol-binding clathrin assembly protein (*PICALM*), and complement component (3b/4b) receptor 1 (*CR1*) genes. *CLU* may play a role in synapse turnover, *PICALM* participates in clathrin-mediated endocytosis, and *CR1* may be involved in amyloid clearance through the complement pathway. *TREM2* is a gene involved with inflammation that increases the likelihood of dementia. Homozygous mutation carriers develop a frontal dementia with bone cysts (Nasu-Hakola disease), whereas heterozygotes are predisposed to the development of AD.

TREATMENT ALZHEIMER'S DISEASE

The management of AD is challenging and gratifying despite the absence of a cure or a robust pharmacologic treatment. The primary focus is on long-term amelioration of associated behavioral and neurologic problems, as well as providing caregiver support.

Building rapport with the patient, family members, and other caregivers is essential to successful management. In the early stages of AD, memory aids such as notebooks and posted daily reminders can be helpful. Family members should emphasize activities that are pleasant while curtailing those that increase stress on the patient. Kitchens, bathrooms, stairways, and bedrooms need to be made safe, and eventually patients will need to stop driving. Loss of independence and change of environment may worsen confusion, agitation, and anger. Communication and repeated calm reassurance are necessary. Caregiver "burnout" is common, often resulting in nursing home placement of the patient or new health problems for the caregiver. Respite breaks for the caregiver help to maintain a successful long-term therapeutic milieu. Use of adult day care centers can be helpful. Local and national support groups, such as the Alzheimer's Association and the Family Caregiver Alliance, are valuable resources. Internet access to these resources has become available to clinicians and families in recent years.

Donepezil (target dose, 10 mg daily), rivastigmine (target dose, 6 mg twice daily or 9.5-mg patch daily), galantamine (target dose 24 mg daily, extended-release), and memantine (target dose, 10 mg twice daily) are approved by the Food and Drug Administration (FDA) for the treatment of AD. Due to hepatotoxicity, tacrine is no longer used. Dose escalations for each of these medications must be carried out over 4–6 weeks to minimize side effects. The pharmacologic action of donepezil, rivastigmine, and galantamine is inhibition of the cholinesterases, primarily acetylcholinesterase, with a resulting increase in cerebral acetylcholine levels. Memantine appears to act by blocking overexcited *N*-methyl-D-aspartate (NMDA) glutamate receptors. Double-blind, placebo-controlled, crossover studies with cholinesterase inhibitors and memantine in moderate to severe AD have shown them to be associated with improved caregiver ratings of patients' functioning and with an apparent decreased rate of decline in cognitive test scores over periods of up to 3 years. The average patient on an anticholinesterase inhibitor maintains his or her mini-mental state examination (MMSE) score for close to a year, whereas a placebo-treated patient declines 2–3 points over the same time period. Memantine, used in conjunction with cholinesterase inhibitors or by itself, slows cognitive deterioration and decreases caregiver burden for patients with moderate to severe AD but is not approved for mild AD. Each of these compounds has only modest efficacy for AD. Cholinesterase inhibitors are relatively easy to administer, and their major side effects are gastrointestinal symptoms (nausea, diarrhea, cramps), altered sleep with unpleasant or vivid dreams, bradycardia (usually benign), and muscle cramps.

In a prospective observational study, the use of estrogen replacement therapy appeared to protect—by about 50%—against development of AD in women. This study seemed to confirm the results of two earlier case-controlled studies. Sadly, a prospective placebo-controlled study of a combined estrogen-progesterone therapy for asymptomatic postmenopausal women increased, rather than decreased, the prevalence of dementia. This study markedly dampened enthusiasm for hormonal treatments to prevent dementia. Additionally, no benefit has been found in the treatment of AD with estrogen alone.

A controlled trial of an extract of *Ginkgo biloba* found modest improvement in cognitive function in subjects with AD and vascular dementia. Unfortunately, a comprehensive 6-year multicenter prevention study using ginkgo found no slowing of progression to dementia in the treated group.

Vaccination against A β_{42} has proved highly efficacious in mouse models of AD, helping clear brain amyloid and preventing further amyloid accumulation. In human trials, this approach led to life-threatening complications, including meningoencephalitis, in a minority of patients. Another experimental approach to AD treatment has been the use of β and γ secretase inhibitors that diminish the production of A β_{42} , but the first two placebo-controlled trials of γ secretase inhibitors, tarenflurbil and semagacestat, were negative, and semagacestat may have accelerated cognitive decline compared to placebo. Passive immunization with monoclonal antibodies against A β_{42} has been tried in mild to moderate AD. These studies were negative, leading some to suggest that the patients treated were too advanced to respond to amyloid-lowering therapies. Therefore, new trials have started in asymptomatic individuals with mild AD, in asymptomatic autosomal dominant forms of AD, and in cognitively normal elderly who are amyloid positive with PET. Medications that modify tau phosphorylation and aggregation, including tau antibodies, are beginning to be studied as possible treatments for both AD and non-AD tau-related disorders including FTD and progressive supranuclear palsy.

Several retrospective studies suggest that nonsteroidal anti-inflammatory agents and 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase inhibitors (statins) may have a protective effect on dementia if used prior to the onset of disease but do not influence clinically symptomatic AD. Finally, there is now a strong interest in the relationship between diabetes and AD, and insulin-regulating studies are being conducted.

Mild to moderate depression is common in the early stages of AD and may respond to antidepressants or cholinesterase inhibitors. Selective serotonin reuptake inhibitors (SSRIs) are commonly used due to their low anticholinergic side effects (for example, escitalopram, target dose 5–10 mg daily). Seizures can be treated with levetiracetam unless the patient had a different regimen that was effective prior to the onset of AD. Agitation, insomnia, hallucinations, and belligerence are especially troublesome characteristics of some AD patients, and these behaviors can lead to nursing home placement. The newer generation of atypical antipsychotics, such as risperidone, quetiapine, and olanzapine, are being used in low doses to treat these neuropsychiatric symptoms. The few controlled studies comparing drugs against behavioral intervention in the treatment of agitation suggest mild efficacy with significant side effects related to sleep, gait, and cardiovascular complications, including an increased risk of death. All antipsychotics carry a black box FDA warning and should be used with caution in the demented elderly; however, careful, daily, nonpharmacologic behavior management is often not available, rendering medications necessary for some patients. Finally, medications with strong anticholinergic effects should be vigilantly avoided, including prescription and over-the-counter sleep aids (e.g., diphenhydramine) or incontinence therapies (e.g., oxybutynin).

VASCULAR DEMENTIA

Dementia associated with cerebrovascular disease can be divided into two general categories: multi-infarct dementia and diffuse white matter disease (also called *leukoaraiosis*, *subcortical arteriosclerotic*

leukoencephalopathy, or *Binswanger's disease*). Cerebrovascular disease appears to be a more common cause of dementia in Asia than in Europe and North America, perhaps due to the increased prevalence of intracranial atherosclerosis. Individuals who have had strokes may develop chronic cognitive deficits, commonly called *multi-infarct dementia*. The strokes may be large or small (sometimes lacunar) and usually involve several different brain regions. The occurrence of dementia depends partly on the total volume of damaged cortex. Patients typically report previous discrete episodes of sudden neurologic deterioration. Many patients with multi-infarct dementia have a history of hypertension, diabetes, coronary artery disease, or other manifestations of widespread atherosclerosis. Physical examination may show focal neurologic deficits such as hemiparesis, a unilateral Babinski sign, a visual field defect, or pseudobulbar palsy. Recurrent strokes result in a stepwise disease progression. Neuroimaging reveals multiple areas of infarction. Thus, the history and neuroimaging findings differentiate this condition from AD; however, both AD and multiple infarctions are common and sometimes co-occur. With normal aging, there is also an accumulation of amyloid in cerebral blood vessels, leading to a condition called *cerebral amyloid angiopathy* (without dementia), which predisposes older persons to lobar hemorrhage and brain microhemorrhages. AD patients appear to be at increased risk for amyloid angiopathy, and this association may explain some of the observed links between AD and stroke.

Some individuals with dementia are discovered on MRI to have bilateral T2 signal hyperintensities in the subcortical white matter, termed *diffuse white matter disease*, often occurring in association with lacunar infarctions (see Fig. 35-2). The dementia may be insidious in onset and progress slowly, features that distinguish it from multi-infarct dementia, but other patients show a stepwise deterioration more typical of multi-infarct dementia. Early symptoms include mild confusion, apathy, anxiety, psychosis, and memory, spatial, or executive deficits. Marked difficulties in judgment and orientation and dependence on others for daily activities develop later. Euphoria, elation, depression, or aggressive behaviors are common as the disease progresses. Pyramidal and cerebellar signs may be present, and a gait disorder is seen in at least half of these patients. With advanced disease, urinary incontinence and dysarthria with or without other pseudobulbar features (e.g., dysphagia, emotional lability) are frequent. Seizures and myoclonic jerks appear in a minority of patients. Often, this disorder results from chronic ischemia due to occlusive disease of small, penetrating cerebral arteries and arterioles (microangiopathy). Any disease-causing stenosis of small cerebral vessels may be the critical underlying factor, although hypertension is the major cause. The term *Binswanger's disease* should be used with caution, because it does not clearly identify a single entity.

Other rare causes of white matter disease also present with dementia, such as adult metachromatic leukodystrophy (arylsulfatase A deficiency) and progressive multifocal leukoencephalopathy (Chap. 164). A dominantly inherited form of white matter disease is known as *cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy* (CADASIL), discussed later in “Other Causes of Dementia.”

Mitochondrial disorders can present with stroke-like episodes and can selectively injure basal ganglia or cortex. Many such patients show other findings suggestive of a neurologic or systemic disorder such as ophthalmoplegia, retinal degeneration, deafness, myopathy, neuropathy, or diabetes. Diagnosis is difficult, but serum or (especially) CSF levels of lactate and pyruvate may be abnormal, and biopsy of affected tissue, preferably muscle, may be diagnostic.

Treatment of vascular dementia must be focused on preventing new ischemic injury by stabilizing or removing the underlying causes, such as hypertension, diabetes, smoking, or lack of exercise. Recovery of lost cognitive function is not likely, although fluctuations with periods of improvement are common.

FRONTOTEMPORAL LOBAR DEGENERATION SPECTRUM

Frontotemporal dementia (FTD) refers to a group of clinical syndromes united by underlying frontotemporal lobar degeneration (FTLD) pathology. FTD most often begins in the fifth to seventh decades and

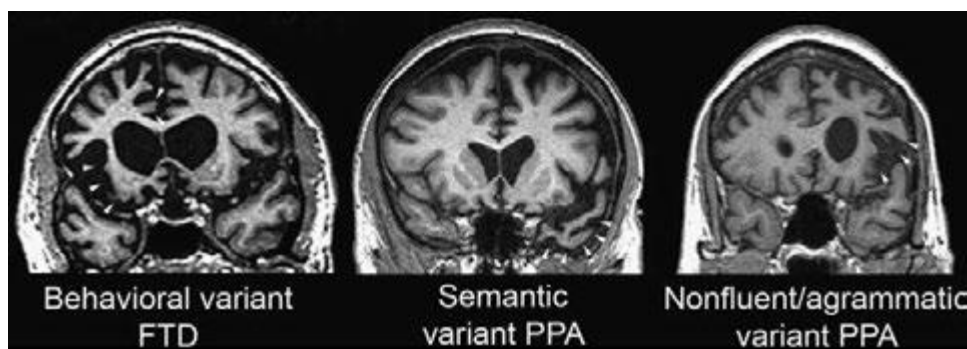


FIGURE 448-3 Three major frontotemporal dementia (FTD) clinical syndromes. Coronal magnetic resonance imaging sections from representative patients with behavioral variant FTD (*left*), semantic dementia (*center*), and progressive nonfluent aphasia (*right*). Areas of early and severe atrophy in each syndrome are highlighted (*white arrowheads*). The behavioral variant features anterior cingulate and frontoinsula atrophy, spreading to orbital and dorsolateral prefrontal cortex. Semantic variant primary progressive aphasia (PPA) shows prominent temporo-polar atrophy, more often on the left. Nonfluent/agrammatic variant PPA is associated with dominant frontal opercular and dorsal insula degeneration.

is nearly as prevalent as AD in this age group. Early studies suggested that FTD may be more common in men than women, although more recent reports cast doubt on this finding. Although a family history of dementia is common, autosomal dominant inheritance is seen in only 10–20% of all FTD cases.

The clinical heterogeneity seen in familial and sporadic FTD is remarkable. Three core clinical syndromes have been described ([Fig. 448-3](#)). In the behavioral variant (bvFTD), the most common FTD syndrome, social and emotional systems dysfunction manifests as apathy, disinhibition, compulsivity, loss of empathy, and overeating, often but not always accompanied by deficits in executive control. Two forms of primary progressive aphasia (PPA), the semantic and nonfluent/agrammatic variants, are commonly due to FTLT and included under the FTD umbrella. In the semantic variant, patients slowly lose the ability to decode word, object, person-specific, and emotion meaning, whereas patients with the nonfluent/agrammatic variant develop profound inability to produce words, often with prominent motor speech impairment. Any of these three clinical syndromes, but most often bvFTD, may be accompanied by motor neuron disease (MND), in which case the term FTD-MND is applied. In addition, the corticobasal syndrome (CBS) and progressive supranuclear palsy syndrome (PSP-S) can be considered part of the FTLT clinical spectrum. Furthermore, patients may evolve from any of the major syndromes described above to have prominent features of another syndrome.

Findings at the bedside are dictated by the anatomic localization of the disorder. Right hemisphere-predominant or symmetric anterior cingulate/medial prefrontal, orbital, and anterior insular degeneration predicts bvFTD. Patients with nonfluent/agrammatic PPA show left (dominant) frontal opercular and precentral gyrus degeneration, whereas left anterior temporal atrophy presents with semantic variant PPA. Visuoconstructive ability, arithmetic calculations, and navigation may remain normal late into any FTD syndrome. Many patients with nonfluent aphasia or bvFTD later develop PSP-S, as disease spreads into diencephalic and brainstem structures, or CBS-like features, as disease moves into dorsal and lateral perirolandic cortices.

The most common autosomal dominantly inherited mutations causing FTD involve the *C9ORF72* (chromosome 9), *GRN* (chromosome 17), and *MAPT* (chromosome 17) genes. Hexanucleotide (GGGGCC) expansions in the noncoding portion of *C9ORF72* are the most recently identified and represent the most common genetic cause of familial or sporadic FTD (usually presenting as bvFTD with or without MND) and amyotrophic lateral sclerosis (ALS). The expansion is associated with reduced *C9ORF72* mRNA expression, nuclear mRNA foci containing transcribed portions of the expansion and other mRNAs, neuronal cytoplasmic inclusions containing dipeptide repeat proteins translated from the repeat mRNA, and transactive response DNA-binding protein of 43 kDa (TDP-43) neuronal cytoplasmic and glial inclusions. The pathogenic significance of these various features

is a topic of vigorous investigation. *MAPT* mutations lead to a change in the alternate splicing of tau or cause loss of function in the tau molecule, thereby altering microtubule binding. With *GRN*, mutations in the coding sequence of the gene encoding progranulin protein result in mRNA degradation due to nonsense-mediated decay, providing a rare example of an autosomal dominant mutation that leads to haploinsufficiency and leads to a ~50% reduction in circulating progranulin protein levels. Intriguingly, a patient with *GRN* mutations on both chromosomes was recently reported to develop neuronal ceroid lipofuscinosis, focusing investigators on the lysosome as a site of molecular dysfunction in *GRN*-related FTD. Progranulin is a growth factor that binds to tumor necrosis factor (TNF) receptors and participates in tissue repair and tumor growth. How progranulin mutations lead to FTD remains unknown, but the most likely mechanisms include lysosomal dysfunction and enhanced neuroinflammation. Both *MAPT* and *GRN* mutations are associated with parkinsonian features, whereas ALS is rare. Infrequently, mutations in the valosin-containing protein (*VCP*, chromosome 9) and charged multivesicular body protein 2b (*CHMP2b*, chromosome 3) genes also lead to autosomal dominant familial FTD. Mutations in the *TARDBP* (encoding TDP-43) and *FUS* (encoding fused in sarcoma [FUS]) genes (see below) cause familial ALS, sometimes in association with an FTD syndrome, although a few patients presenting with FTD alone have been reported.

The gross pathologic hallmark of FTLT is a focal atrophy of frontal, insular, and/or temporal cortex, which can be visualized with neuroimaging studies ([Fig. 448-3](#)) and is often profound at autopsy. Despite the appearance of advanced disease, however, imaging studies suggest that atrophy often begins focally in one hemisphere before spreading to anatomically interconnected regions, including basal ganglia. Loss of cortical serotonergic innervation is seen in many patients. In contrast to AD, the cholinergic system is relatively spared in FTD, which accounts for the poor efficacy of acetylcholinesterase inhibitors in this group.

Although early studies suggested that 15–30% of patients with FTD showed underlying AD at autopsy, progressive refinement in clinical diagnosis has improved pathologic prediction accuracy, and most patients diagnosed with FTD at a dementia clinic with expertise in FTD will show underlying FTLT pathology. Microscopic findings seen across all patients with FTLT include gliosis, microvacuolation, and neuronal loss, but the disease is subtyped according to the protein composition of neuronal and glial inclusions, which contain either tau or TDP-43 in ~90% of patients, with the remaining ~10% showing inclusions containing *FUS* ([Fig. 448-4](#)).

The toxicity and spreading capacity of tau aggregates underlies the pathogenesis of many familial cases and is emerging as a key factor in sporadic tauopathies, although loss of tau microtubule stabilizing function may also play a role. TDP-43 and *FUS*, in contrast, are RNA/DNA binding proteins whose roles in neuronal function are still being actively investigated, but one key role may be the chaperoning of

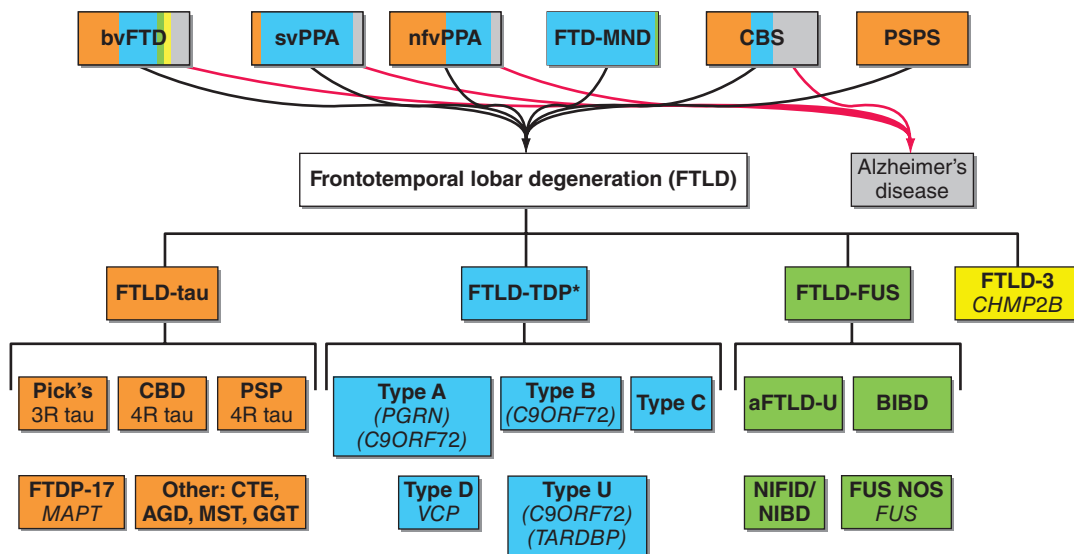


FIGURE 448-4 Frontotemporal dementia syndromes are united by underlying frontotemporal lobar degeneration pathology, which can be divided according to the presence of tau, TDP-43, or FUS-containing inclusions in neurons and glia. Correlations between clinical syndromes and major molecular classes are shown with colored shading. Despite improvements in clinical syndromic diagnosis, a small percentage of patients with some frontotemporal dementia syndromes will show Alzheimer's disease neuropathology at autopsy (*gray shading*). aFTLD-U, atypical frontotemporal lobar degeneration with ubiquitin-positive inclusions; AGD, argyrophilic grain disease; BIBD, basophilic inclusion body disease; bvFTD, behavioral variant frontotemporal dementia; CBD, corticobasal degeneration; CBS, corticobasal syndrome; CTE, chronic traumatic encephalopathy; FTD-MND, frontotemporal dementia with motor neuron disease; FTDP-17, frontotemporal dementia with parkinsonism linked to chromosome 17; FUS, fused in sarcoma; GGT, globular glial tauopathy; MST, multisystem tauopathy; nvfPPA, nonfluent/agrammatic variant primary progressive aphasia; NIBD, neurofilament inclusion body disease; NIFID, neuronal intermediate filament inclusion disease; PSP, progressive supranuclear palsy; PSPS, progressive supranuclear palsy syndrome; svPPA, semantic variant primary progressive aphasia; Type U, unclassifiable type.

mRNAs to the distal neuron for activity-dependent translation within dendritic spines. Because these proteins also form intracellular aggregates and produce similar anatomic progression, protein toxicity and spreading may also factor heavily in the pathogenesis of these FTLD-TDP and FTLD-FUS.

Increasingly, misfolded proteins in neurodegenerative disease are being recognized as having “prion-like” properties in that they can template the misfolding of their natively folded protein counterparts, a process that creates exponential amplification of protein misfolding within a cell and may promote transcellular and even transsynaptic protein propagation between cells. This hypothesis could provide a unifying explanation for the stereotypical patterns of disease spread observed in each syndrome (**Chap. 444e**).

Although the term *Pick's disease* was once used to describe a progressive degenerative disorder characterized by selective involvement of the anterior frontal and temporal neocortex and pathologically by intraneuronal cytoplasmic inclusions (*Pick bodies*), it is now used only in reference to a specific FTLD-tau histopathologic entity. Classical Pick bodies are argyrophilic, staining positively with the Bielschowsky silver method (but not with the Gallyas method) and also with immunostaining for hyperphosphorylated tau. Recognition of the three FTLD major molecular classes has allowed delineation of distinct FTLD subtypes within each class. These subtypes, based on the morphology and distribution of the neuronal and glial inclusions (**Fig. 448-5**), account for the vast majority of patients, and some subtypes show strong clinical or genetic associations (**Fig. 448-4**). Despite this progress, available data do not allow reliable prediction of the underlying FTLD subtype, or even the major molecular class, based on clinical features alone. Molecular PET imaging with ligands chosen to bind misfolded tau protein shows great promise and is already being applied to the study of patients with AD and FTD. Because FTLD-tau and FTLD-TDP account for 90% of FTLD patients, the ability to detect pathologic tau protein deposition *in vivo* will greatly improve prediction accuracy, especially when amyloid PET imaging is negative.

The burden on caregivers of patients with FTD is extremely high, especially when the illness disrupts core emotional and personality

functions of the loved one. Treatment is symptomatic, and there are currently no therapies known to slow progression or improve symptoms. Many of the behaviors that may accompany FTD, such as depression, hyperorality, compulsions, and irritability, can be ameliorated with antidepressants, especially SSRIs. The co-association with motor disorders such as parkinsonism necessitates the careful use of antipsychotics, which can exacerbate this problem.

Progressive supranuclear palsy syndrome (PSP-S; also known as Steele-Richardson-Olszewski syndrome) is a degenerative disorder that involves the brainstem, basal ganglia, limbic structures, and selected areas of cortex. Clinically, PSP-S begins with falls and executive or subtle personality changes (such as mental rigidity, impulsivity, or apathy). Shortly thereafter, a progressive oculomotor syndrome ensues that begins with square wave jerks, followed by slowed saccades (vertical worse than horizontal) before resulting in progressive supranuclear ophthalmoparesis. Dysarthria, dysphagia, and symmetric axial rigidity can be prominent features that emerge at any point in the illness. A stiff, unstable posture with hyperextension of the neck and a slow, jerky, toppling gait are characteristic. Frequent unexplained and sometimes spectacular falls are common secondary to a combination of axial rigidity, inability to look down, and poor judgment. Even once patients have severely limited voluntary eye movements, they retain oculocephalic reflexes (demonstrated using a vertical doll's head maneuver); thus, the oculomotor disorder is supranuclear. The dementia overlaps with bvFTD, featuring apathy, frontal-executive dysfunction, poor judgment, slowed thought processes, impaired verbal fluency, and difficulty with sequential actions and with shifting from one task to another. These features are common at presentation and often precede the motor syndrome. Some patients with a pathologic diagnosis of PSP begin with a nonfluent aphasia or motor speech disorder and progress to classical PSP-S. Response to L-dopa is limited or absent; no other treatments exist. Death occurs within 5–10 years of onset. Like Pick's disease, increasingly the term PSP is used to refer to a specific histopathologic entity within the FTLD-tau class. In PSP, accumulation of hyperphosphorylated 4-repeat tau is seen within neurons and glia. Neuronal inclusions often take the form of NFTs, which may

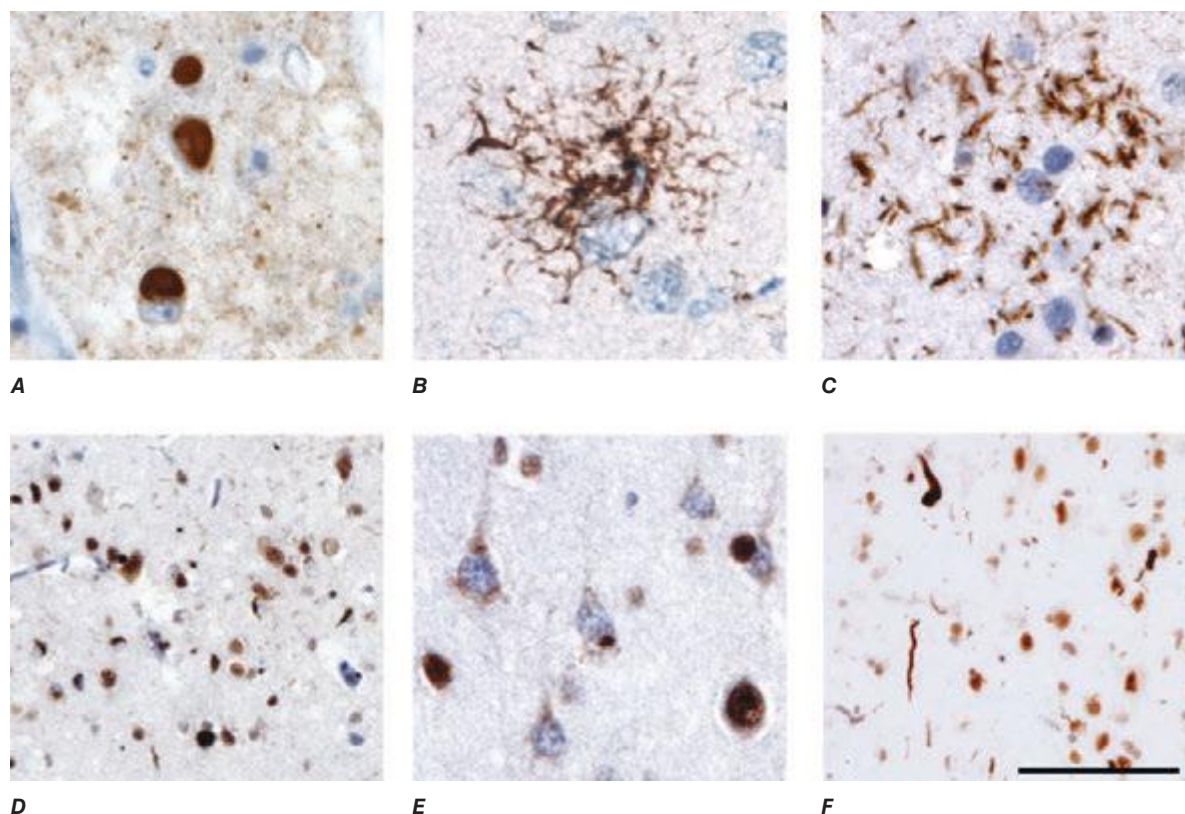


FIGURE 448-5 Neuropathology in frontotemporal lobar degeneration (FTLD). FTLD-tau (**A–C**) and FTLD-TDP (**D–F**) account for over 90% of patients with FTLD, and immunohistochemistry reveals characteristic lesions in each of the major histopathologic subtypes within each class: (**A**) Pick bodies in Pick's disease; (**B**) a tufted astrocyte in progressive supranuclear palsy; (**C**) an astrocytic plaque in corticobasal degeneration; (**D**) small compact or crescentic neuronal cytoplasmic inclusions and short, thin neuropil threads in FTLD-TDP, type A; (**E**) diffuse/granular neuronal cytoplasmic inclusions (with a relative paucity of neuropil threads) in FTLD-TDP, type B; and (**F**) long, tortuous dystrophic neurites in FTLD-TDP, type C. TDP can be seen within the nucleus in neurons lacking inclusions but mislocalizes to the cytoplasm and forms inclusions in FTLD-TDP. Immunostains are 3-repeat tau (**A**), phospho-tau (**B** and **C**), and TDP-43 (**D–F**). Sections are counterstained with hematoxylin. Scale bar applies to all panels and represents 50 μm in **A, B, C,** and **E** and 100 μm in **D** and **F**.

be large, spherical (“globose”), and coarse in brainstem, cerebellar dentate, and diencephalic neurons. Tau deposition is most prominent in subcortical structures (including the subthalamic nucleus, globus pallidus, substantia nigra, locus coeruleus, periaqueductal gray, tectum, oculomotor nuclei, and dentate nucleus of cerebellum). Neocortical NFTs, like those in AD, often take on a more flame-shaped morphology, but on electron microscopy, PSP tangles can be shown to consist of straight tubules rather than the paired helical filaments found in AD. Furthermore, PSP is associated with prominent tau-positive glial pathologies, such as tufted astrocytes (Fig. 448-5), thorny astrocytes, and coiled oligodendroglial inclusions (“coiled bodies”). Most patients with PSP-S show PSP at autopsy, although small numbers will show another tauopathy (corticobasal degeneration [CBD] or Pick's disease; Fig. 448-4).

In addition to its overlap with FTD and CBS (see below), PSP is often confused with idiopathic *Parkinson's disease* (PD). Although elderly patients with PD may have restricted upgaze, they do not develop downgaze paresis or other abnormalities of voluntary eye movements typical of PSP. Dementia occurs in ~20% of patients with PD, often due to the emergence of a full-blown DLB-like syndrome. Furthermore, the behavioral syndromes seen with DLB differ from PSP (see below). Dementia in PD becomes more likely with increasing age, increasing severity of extrapyramidal signs, long disease duration, and the presence of depression. Patients with PD who develop dementia also show cortical atrophy on brain imaging. Neuropathologically, there may be AD-related changes in the cortex, LBD-related α -synuclein inclusions in both the limbic system and cortex, or no specific microscopic changes other than gliosis and neuronal loss. **PD is discussed in detail in Chap. 449.**

Corticobasal syndrome (CBS) is a slowly progressive dementia-movement disorder associated with severe atrophy in perirolandic cortex and basal ganglia (substantia nigra and striatopallidum). Patients typically present with asymmetric onset of rigidity, dystonia, myoclonus, and apraxia of one limb, at times associated with *alien limb* phenomena in which the limb exhibits unintended motor actions such as grasping, groping, drifting, or undoing. Eventually CBS becomes bilateral and leads to dysarthria, slow gait, action tremor, and typically a frontal-predominant dementia. Whereas CBS refers to the clinical syndrome, CBD refers to a specific histopathologic FTLD-tau entity (Fig. 448-4). Although CBS was once thought to be pathognomonic for CBD, increasingly it has been recognized that CBS can be due to CBD, PSP, FTLD-TDP, or even AD. In CBD, the microscopic features include ballooned, achromatic, tau-positive neurons; astrocytic plaques (Fig. 448-5); and other dystrophic glial tau pathomorphologies that overlap with those seen in PSP. Most specifically, CBD features a severe tauopathy burden in the subcortical white matter, consisting of threads and oligodendroglial coiled bodies. As shown in Fig. 448-4, patients with bvFTD, nonfluent/agrammatic PPA, and PSP-S may also show CBD at autopsy, emphasizing the importance of distinguishing clinical and pathologic constructs and terminology. Treatment of CBS remains symptomatic; no disease-modifying therapies are available.

PARKINSON'S DISEASE DEMENTIA AND DEMENTIA WITH LEWY BODIES

The parkinsonian dementia syndromes are under increasing study, with many cases unified by Lewy body and Lewy neurite pathology that ascends from the low brainstem up through the substantia nigra,

limbic system, and cortex. The DLB clinical syndrome is characterized by visual hallucinations, parkinsonism, fluctuating alertness, falls, and often rapid eye movement (REM) sleep behavior disorder (RBD). Dementia can precede or follow the appearance of parkinsonism. Hence, one pathway occurs in patients with long-standing PD without cognitive impairment, who slowly develop a dementia that is associated with visual hallucinations and fluctuating alertness. When this occurs after an established diagnosis of PD, many use the term *Parkinson's disease dementia* (PDD). In others, the dementia and neuropsychiatric syndrome precede or co-emerge with the parkinsonism, and this constellation is referred to as DLB. Both PDD and DLB may be accompanied or preceded by symptoms referable to brainstem pathology below the substantia nigra including constipation, orthostatic lightheadedness, or RBD, and many researchers conceptualize these disorders as points on a spectrum of α -synuclein pathology.

Patients with PDD and DLB are highly sensitive to metabolic perturbations, and in some patients, the first manifestation of illness is a delirium, often precipitated by an infection, new medicine, or other systemic disturbance. A hallucinatory delirium induced by L-dopa, prescribed for parkinsonian symptoms attributed to PD, may likewise provide the initial clue to a PDD or DLB diagnosis. Conversely, patients with mild cognitive deficits and hallucinations may receive typical or atypical antipsychotic medications, which induce profound parkinsonism at low doses due to a subclinical DLB-related nigral dopaminergic neuron loss. Even without an underlying precipitant, fluctuations can be marked in DLB, with episodic confusion or even stupor admixed with lucid intervals. Despite the fluctuating pattern, however, the core clinical features persist, unlike delirium, which resolves following correction of the inciting factor. Cognitively, DLB features relative preservation of memory but more severe visuospatial and executive deficits than seen in patients with early AD.

The key neuropathologic feature in DLB is the presence of Lewy bodies and Lewy neurites throughout specific brainstem nuclei, substantia nigra, amygdala, cingulate gyrus, and, ultimately, the neocortex. Lewy bodies are intraneuronal cytoplasmic inclusions that stain with periodic acid-Schiff (PAS) and ubiquitin but are now identified with antibodies to the presynaptic protein, α -synuclein. Lewy bodies are composed of straight neurofilaments 7–20 nm long with surrounding amorphous material and contain epitopes recognized by antibodies against phosphorylated and nonphosphorylated neurofilament proteins, ubiquitin, and α -synuclein. Lewy bodies are typically found in the substantia nigra of patients with idiopathic PD, where they can be readily seen with hematoxylin-and-eosin staining. A profound cholinergic deficit, owing to basal forebrain and pedunculopontine nucleus involvement, is present in many patients with DLB and may be a factor responsible for the fluctuations, inattention, and visual hallucinations.

Due to the frequent comorbidity with AD and the cholinergic deficit in DLB, cholinesterase inhibitors often provide significant benefit, reducing hallucinosis, stabilizing delusional symptoms, and even helping with RBD in some patients. Exercise programs maximize motor function and protect against fall-related injury. Antidepressants are often necessary. Atypical antipsychotics may be required for psychosis but can worsen extrapyramidal syndromes, even at low doses, and increase risk of death. Patients with DLB are extremely sensitive to dopaminergic medications, which must be carefully titrated; tolerability may be improved by concomitant use of a cholinesterase inhibitor.

OTHER CAUSES OF DEMENTIA

Prion diseases such as *Creutzfeldt-Jakob disease* (CJD) are rare neurodegenerative conditions (prevalence ~1 per million) that produce dementia. CJD is a rapidly progressive disorder associated with dementia, focal cortical signs, rigidity, and myoclonus, causing death <1 year after first symptoms appear. The rapidity of progression seen with CJD is uncommon in AD so that the distinction between the two disorders is usually straightforward. CBD and DLB, more rapid degenerative dementias with prominent movement abnormalities, are more likely to be mistaken for CJD. The differential diagnosis for CJD includes other rapidly progressive dementing conditions such as

viral or bacterial encephalitides, Hashimoto's encephalopathy, central nervous system (CNS) vasculitis, lymphoma, or paraneoplastic/autoimmune syndromes. The markedly abnormal periodic complexes on EEG and cortical ribboning and basal ganglia hyperintensities on fluid-attenuated inversion recovery MRI are diagnostic features of CJD, although rarely, prolonged focal or generalized seizures can produce a similar imaging appearance. **Prion diseases are discussed in detail in Chap. 453e.**

Huntington's disease (HD) (**Chap. 449**) is an autosomal dominant degenerative brain disorder. HD clinical hallmarks include chorea, behavioral disturbance, and executive impairment. Symptoms typically begin in the fourth or fifth decade, but there is a wide range, from childhood to >70 years. Memory is frequently not impaired until late in the disease, but attention, judgment, self-awareness, and executive functions are often deficient at an early stage. Depression, apathy, social withdrawal, irritability, and intermittent disinhibition are common. Delusions and obsessive-compulsive behavior may occur. Disease duration is variable but typically lasts approximately 15 years.

Normal-pressure hydrocephalus (NPH) is a relatively uncommon but treatable syndrome. The clinical, physiologic, and neuroimaging characteristics of NPH must be carefully distinguished from those of other dementias associated with gait impairment. Historically, many patients treated for NPH have suffered from other dementias, particularly AD, vascular dementia, DLB, and PSP. For NPH, the clinical triad includes an abnormal gait (ataxic or apractic), dementia (usually mild to moderate, with an emphasis on executive impairment), and urinary urgency or incontinence. Neuroimaging reveals enlarged lateral ventricles (hydrocephalus) with little or no cortical atrophy, although the sylvian fissures may appear propped open (so-called "boxcarring"), which can be mistaken for perisylvian atrophy. This syndrome is a communicating hydrocephalus with a patent aqueduct of Sylvius (**see Fig. 35-3**), in contrast to aqueductal stenosis, in which the aqueduct is small. Lumbar puncture opening pressure falls in the high-normal range, and the CSF protein, glucose, and cell counts are normal. NPH may be caused by obstruction to normal CSF flow over the cerebral convexities and delayed resorption into the venous system. The indolent nature of the process results in enlarged lateral ventricles with relatively little increase in CSF pressure. Presumed edema, stretching, and distortion of subfrontal white matter tracts may lead to clinical symptoms, but the precise underlying pathophysiology remains unclear. Some patients provide a history of conditions that produce meningeal scarring (blocking CSF resorption) such as previous meningitis, subarachnoid hemorrhage, or head trauma. Others with long-standing but asymptomatic congenital hydrocephalus may have adult-onset deterioration in gait or memory that is confused with NPH. In contrast to AD, the patient with NPH complains of an early and prominent gait disturbance without cortical atrophy on CT or MRI.

Numerous attempts to improve NPH diagnosis with various special studies and predict the success of ventricular shunting have been undertaken. These tests include radionuclide cisternography (showing a delay in CSF absorption over the convexity) and various efforts to monitor and alter CSF flow dynamics, including a constant-pressure infusion test. None has proven to be specific or consistently useful. A transient improvement in gait or cognition may follow lumbar puncture (or serial punctures) with removal of 30–50 mL of CSF, but this finding has also not proved to be consistently predictive of postshunt improvement. Perhaps the most reliable strategy is a period of close inpatient evaluation before, during, and after lumbar CSF drainage. Occasionally, when a patient with AD presents with gait impairment (at times due to comorbid subfrontal vascular injury) and absent or only mild cortical atrophy on CT or MRI, distinguishing NPH from AD can be challenging. Hippocampal atrophy on MRI favors AD, whereas a characteristic "magnetic" gait with external hip rotation, low foot clearance, and short strides, along with prominent truncal sway or instability, favors NPH. The diagnosis of NPH should be avoided when hydrocephalus is not detected on imaging studies, even if the symptoms otherwise fit. Thirty to fifty percent of patients identified by careful diagnosis as having NPH will improve with ventricular shunting. Gait may improve more than cognition, but many reported failures

to improve cognitively may have resulted from comorbid AD. Short-lasting improvement is common. Patients should be carefully selected for shunting, because subdural hematoma, infection, and shunt failure are known complications and can be a cause for early nursing home placement in an elderly patient with previously mild dementia.

Intracranial hypotension, sometimes called sagging brain syndrome, is a disorder caused by low CSF pressure, leading to downward pressure on the subcortical structures and disruption of cerebral function. It presents in a variable manner with headache, often exacerbated by coughing or a Valsalva maneuver or by moving from lying to standing. Other common symptoms include dizziness, vomiting, disruption of sleep-wake cycles, and sometimes a progressive bvFTD-like syndrome. Although sometimes idiopathic, this syndrome can be caused by CSF leaks secondary to lumbar puncture, head trauma, or spinal cord arachnoid cysts. Treatment consists of finding and patching CSF leaks.

Dementia can accompany *chronic alcoholism* (Chap. 467) and may result from associated malnutrition, especially of B vitamins, particularly thiamine. Other poorly defined aspects of chronic alcoholism may, however, also produce cerebral damage. A rare idiopathic syndrome of dementia and seizures with degeneration of the corpus callosum has been reported primarily in male Italian red wine drinkers (Marchiafava-Bignami disease).

Thiamine (vitamin B₁) deficiency causes Wernicke's encephalopathy (Chap. 330). The clinical presentation features a malnourished patient (frequently but not necessarily alcoholic) with confusion, ataxia, and diplopia resulting from inflammation and necrosis of periventricular midline structures, including dorsomedial thalamus, mammillary bodies, midline cerebellum, periaqueductal gray matter, and trochlear and abducens nuclei. Damage to the dorsomedial thalamus correlates most closely with the memory loss. Prompt administration of parenteral thiamine (100 mg intravenously for 3 days followed by daily oral dosage) may reverse the disease if given in the first days of symptom onset. Prolonged untreated thiamine deficiency can result in an irreversible and profound amnesic syndrome (Korsakoff's syndrome) or even death.

In *Korsakoff's syndrome*, the patient is unable to recall new information despite normal immediate memory, attention span, and level of consciousness. Memory for new events is seriously impaired, whereas knowledge acquired prior to the illness remains relatively intact. Patients are easily confused, disoriented, and cannot store information for more than a few minutes. Superficially, they may be conversant, engaging, and able to perform simple tasks and follow immediate commands. Confabulation is common, although not always present. There is no specific treatment because the previous thiamine deficiency has produced irreversible damage to the medial thalamic nuclei and mammillary bodies. Mammillary body atrophy may be visible on MRI in the chronic phase (see Fig. 330-6).

Vitamin B₁₂ deficiency, as can occur in pernicious anemia, causes a megaloblastic anemia and may also damage the nervous system (Chaps. 128 and 456). Neurologically, it most commonly produces a spinal cord syndrome (myelopathy) affecting the posterior columns (loss of vibration and position sense) and corticospinal tracts (hyperactive tendon reflexes with Babinski signs); it also damages peripheral nerves (neuropathy), resulting in sensory loss with depressed tendon reflexes. Damage to myelinated axons may also cause dementia. The mechanism of neurologic damage is unclear but may be related to a deficiency of S-adenosyl methionine (required for methylation of myelin phospholipids) due to reduced methionine synthase activity or accumulation of methylmalonate, homocysteine, and propionate, providing abnormal substrates for fatty acid synthesis in myelin. Use of histamine blockers or metformin, vegan diets, autoimmunity against gastric parietal cells, and various causes of malabsorption are the typical causes for vitamin B₁₂ deficiency. The neurologic sequelae of vitamin B₁₂ deficiency may occur in the absence of hematologic manifestations, making it critical to avoid using the complete blood count (CBC) and blood smear as a substitute for measuring B₁₂ blood levels. Treatment with parenteral vitamin B₁₂ (1000 µg intramuscularly daily for a week, weekly for a month, and monthly for life for pernicious anemia) stops progression of the disease if instituted promptly, but complete reversal of advanced nervous system damage will not occur.

Deficiency of nicotinic acid (*pellagra*) is associated with skin rash over sun-exposed areas, glossitis, and angular stomatitis (Chap. 96e). Severe dietary deficiency of nicotinic acid along with other B vitamins such as pyridoxine may result in spastic paraparesis, peripheral neuropathy, fatigue, irritability, and dementia. This syndrome has been seen in prisoners of war and in concentration camps but should be considered in any malnourished individual. Low serum folate levels appear to be a rough index of malnutrition, but isolated folate deficiency has not been proved as a specific cause of dementia.

CNS infections usually cause delirium and other acute neurologic syndromes. However, some chronic CNS infections, particularly those associated with chronic meningitis (Chap. 165), may produce a dementing illness. The possibility of chronic infectious meningitis should be suspected in patients presenting with a dementia or behavioral syndrome, who also have headache, meningismus, cranial neuropathy, and/or radiculopathy. Between 20 and 30% of patients in the advanced stages of HIV infection become demented (Chap. 226). Cardinal features include psychomotor retardation, apathy, and impaired memory. This syndrome may result from secondary opportunistic infections but can also be caused by direct infection of CNS neurons with HIV. Neurosyphilis (Chap. 206) was a common cause of dementia in the preantibiotic era; it is now uncommon but can still be encountered in patients with multiple sex partners, particularly among patients with HIV. Characteristic CSF changes consist of pleocytosis, increased protein, and a positive Venereal Disease Research Laboratory (VDRL) test.

Primary and metastatic *neoplasms of the CNS* (Chap. 118) usually produce focal neurologic findings and seizures rather than dementia, but if tumor growth begins in the frontal or temporal lobes, the initial manifestations may be memory loss or behavioral changes. A paraneoplastic syndrome of dementia associated with occult carcinoma (often small-cell lung cancer) is termed *limbic encephalitis*. In this syndrome, confusion, agitation, seizures, poor memory, emotional changes, and frank dementia may occur. Paraneoplastic *encephalitis associated with NMDA receptor antibodies* presents as a progressive psychiatric disorder with memory loss and seizures; affected patients are often young women with ovarian teratoma (Chap. 122).

A *nonconvulsive seizure disorder* (Chap. 445) may underlie a syndrome of confusion, clouding of consciousness, and garbled speech. Often, psychiatric disease is suspected, but an EEG demonstrates the epileptic nature of the illness. If recurrent or persistent, the condition may be termed *complex partial status epilepticus*. The cognitive disturbance often responds to anticonvulsant therapy. The etiology may be previous small strokes or head trauma; some cases are idiopathic.

It is important to recognize *systemic diseases* that indirectly affect the brain and produce chronic confusion or dementia. Such conditions include hypothyroidism; vasculitis; and hepatic, renal, or pulmonary disease. Hepatic encephalopathy may begin with irritability and confusion and slowly progress to agitation, lethargy, and coma.

Isolated vasculitis of the CNS (CNS granulomatous angiitis) (Chaps. 385 and 446) occasionally causes a chronic encephalopathy associated with confusion, disorientation, and clouding of consciousness. Headache is common, and strokes and cranial neuropathies may occur. Brain imaging studies may be normal or nonspecifically abnormal. CSF analysis reveals a mild pleocytosis or protein elevation. Cerebral angiography can show multifocal stenoses involving medium-caliber vessels, but some patients have only small-vessel disease that is not revealed on angiography. The angiographic appearance is not specific and may be mimicked by atherosclerosis, infection, or other causes of vascular disease. Brain or meningeal biopsy demonstrates endothelial cell proliferation and mononuclear infiltrates within blood vessel walls. The prognosis is often poor, although the disorder may remit spontaneously. Some patients respond to glucocorticoids or chemotherapy.

Chronic metal exposure represents a rare cause of dementia. The key to diagnosis is to elicit a history of exposure at work or home. Chronic lead poisoning from inadequately fire-glazed pottery has been reported. Fatigue, depression, and confusion may be associated with episodic abdominal pain and peripheral neuropathy. Gray lead lines appear in the gums, usually accompanied by an anemia with basophilic stippling of red blood cells. The clinical presentation can

resemble that of acute intermittent porphyria, including elevated levels of urine porphyrins as a result of the inhibition of δ -aminolevulinic acid dehydrase. The treatment is chelation therapy with agents such as ethylenediamine tetraacetic acid (EDTA). Chronic mercury poisoning produces dementia, peripheral neuropathy, ataxia, and tremulousness that may progress to a cerebellar intention tremor or choreoathetosis. The confusion and memory loss of chronic arsenic intoxication is also associated with nausea, weight loss, peripheral neuropathy, pigmentation and scaling of the skin, and transverse white lines of the fingernails (Mees' lines). Treatment is chelation therapy with dimercaprol (BAL). Aluminum poisoning is rare but was documented with the dialysis dementia syndrome, in which water used during renal dialysis was contaminated with excessive amounts of aluminum. This poisoning resulted in a progressive encephalopathy associated with confusion, nonfluent aphasia, memory loss, agitation, and, later, lethargy and stupor. Speech arrest and myoclonic jerks were common and associated with severe and generalized EEG changes. The condition has been eliminated by the use of deionized water for dialysis.

Recurrent head trauma in professional athletes may lead to a dementia previously referred to as "punch-drunk" syndrome or *dementia pugilistica* but now known as chronic traumatic encephalopathy (CTE) to signify its relevance to contact sport athletes other than boxers. The symptoms can be progressive, beginning late in an athlete's career or, more often, after retirement. Early in the course, a personality change associated with social instability and sometimes paranoia and delusions occurs. Later, memory loss progresses to full-blown dementia, often associated with parkinsonian signs and ataxia or intention tremor. At autopsy, the cerebral cortex shows changes tau-immunoreactive NFTs that are more prominent than amyloid plaques (which are usually diffuse or absent rather than neuritic). NFTs and tau-positive reactive astrocytes are often clustered in the depths of cortical sulci, and TDP-43 inclusions have also been reported, highlighting the overlap with the FTL spectrum. Loss of neurons in the substantia nigra is a variable feature.

Chronic subdural hematoma (Chap. 457e) is also occasionally associated with dementia, often in the context of underlying cortical atrophy from conditions such as AD or HD.

Transient global amnesia (TGA) is characterized by the sudden onset of a severe episodic memory deficit, usually occurring in persons over the age of 50 years. Often the amnesia occurs in the setting of an emotional stimulus or physical exertion. During the attack, the individual is alert and communicative, general cognition seems intact, and there are no other neurologic signs or symptoms. The patient may seem confused and repeatedly ask about his or her location in place and time. The ability to form new memories returns after a period of hours, and the individual returns to normal with no recall for the period of the attack. Frequently no cause is determined, but cerebrovascular disease, epilepsy (7% in one study), migraine, or cardiac arrhythmias have all been implicated. Approximately one-quarter of patients experience recurrent attacks. Rare instances of permanent memory loss have been reported in patients with TGA-like spells, usually representing ischemic infarction of the hippocampus or dorsomedial thalamic nucleus bilaterally. Seizure activity due to AD should always be suspected in this syndrome.

The *ALS/parkinsonian/dementia complex of Guam* is a rare degenerative disease that has occurred in the Chamorro natives on the island of Guam. Individuals may have any combination of parkinsonian features, dementia, and MND. The most characteristic pathologic features are the presence of NFTs in degenerating neurons of the cortex and substantia nigra and loss of motor neurons in the spinal cord, although recent reanalysis has shown that some patients with this illness also show coexisting TDP-43 pathology. Epidemiologic evidence supports a possible environmental cause, such as exposure to a neurotoxin or an infectious agent with a long latency period. One interesting but unproven candidate neurotoxin occurs in the seed of the false palm tree, which Guamanians traditionally used to make flour. The ALS syndrome is no longer present in Guam, but a dementing illness with rigidity continues to be seen.

Rarely, adult-onset leukodystrophies, lysosomal storage diseases, and other genetic disorders can present as a dementia in middle to

late life. Metachromatic leukodystrophy (MLD) causes a progressive psychiatric or dementia syndrome associated with extensive, confluent frontal white matter abnormality. MLD is diagnosed by measuring arylsulfatase A enzyme activity in white blood cells. Adult-onset presentations of adrenoleukodystrophy have been reported in female carriers, and these patients often feature spinal cord and posterior white matter involvement. Adrenoleukodystrophy is diagnosed with measurement of plasma very-long-chain fatty acids. CADASIL is another genetic syndrome associated with white matter disease, often frontally and temporally predominant. Diagnosis is made with skin biopsy, which shows osmophilic granules in arterioles, or, increasingly, through genetic testing for mutations in Notch 3. The neuronal ceroid lipofuscinoses are a genetically heterogeneous group of disorders associated with myoclonus, seizures, vision loss, and progressive dementia. Diagnosis is made by finding eosinophilic curvilinear inclusions within white blood cells or neuronal tissue.

Psychogenic amnesia for personally important memories can be seen. Whether this results from deliberate avoidance of unpleasant memories, outright malingering, or unconscious repression remains unknown and probably depends on the patient. Event-specific amnesia is more likely to occur after violent crimes such as homicide of a close relative or friend or sexual abuse. It may develop in association with severe drug or alcohol intoxication and sometimes with schizophrenia. More prolonged psychogenic amnesia occurs in fugue states that also commonly follow severe emotional stress. The patient with a fugue state suffers from a sudden loss of personal identity and may be found wandering far from home. *In contrast to neurologic amnesia, fugue states are associated with amnesia for personal identity and events closely associated with the personal past.* At the same time, memory for other recent events and the ability to learn and use new information are preserved. The episodes usually last hours or days and occasionally weeks or months while the patient takes on a new identity. On recovery, there is a residual amnesia gap for the period of the fugue. Very rarely does selective loss of autobiographic information reflect a focal injury to the brain areas involved with these functions.

Psychiatric diseases may mimic dementia. Severely depressed or anxious individuals may appear demented, a phenomenon sometimes called *pseudodementia*. Memory and language are usually intact when carefully tested, and a significant memory disturbance usually suggests an underlying dementia, even if the patient is depressed. Patients in this condition may feel confused and unable to accomplish routine tasks. Vegetative symptoms, such as insomnia, lack of energy, poor appetite, and concern with bowel function, are common. Onset is often more abrupt, and the psychosocial milieu may suggest prominent reasons for depression. Such patients respond to treatment of the underlying psychiatric illness. Schizophrenia is usually not difficult to distinguish from dementia, but occasionally the distinction can be problematic. Schizophrenia generally has a much earlier age of onset (second and third decades) than most dementing illnesses and is associated with intact memory. The delusions and hallucinations of schizophrenia are usually more complex, bizarre, and threatening than those of dementia. Some chronic schizophrenics develop an unexplained progressive dementia late in life that is not related to AD. Conversely, FTD, HD, vascular dementia, DLB, AD, or leukoencephalopathy can begin with schizophrenia-like features, leading to the misdiagnosis of a psychiatric condition. Later age of onset, significant deficits on cognitive testing, or the presence of abnormal neuroimaging suggest a degenerative condition. Memory loss may also be part of a *conversion disorder*. In this situation, patients commonly complain bitterly of memory loss, but careful cognitive testing either does not confirm the deficits or demonstrates inconsistent or unusual patterns of cognitive problems. The patient's behavior and "wrong" answers to questions often indicate that he or she understands the question and knows the correct answer.

Clouding of cognition by *chronic drug or medication use*, often prescribed by physicians, is an important cause of dementia. Sedatives, tranquilizers, and analgesics used to treat insomnia, pain, anxiety, or agitation may cause confusion, memory loss, and lethargy, especially in the elderly. Discontinuation of the offending medication often improves mentation.

449 Parkinson's Disease and Other Movement Disorders

C. Warren Olanow, Anthony H.V. Schapira, Jose A. Obeso

PARKINSON'S DISEASE AND RELATED DISORDERS

Parkinson's disease (PD) is the second commonest neurodegenerative disease, exceeded only by Alzheimer's disease (AD). Its cardinal clinical features were first described by the English physician James Parkinson in 1817. It is noteworthy that James Parkinson was a general physician who captured the essence of this condition based on a visual inspection of a mere handful of patients. It is estimated that approximately 1 million persons in the United States, 1 million in Western Europe, and 5 million worldwide suffer from this disorder. PD affects men and women of all races, all occupations, and all countries. The mean age of onset is about 60 years. The frequency of PD increases with aging, but cases can be seen in patients in their 20s and even younger. Based on the aging of the population and projected demographics, it is estimated that the prevalence of the disease will dramatically increase in the next several decades.

Clinically, PD is characterized by rest tremor, rigidity, bradykinesia (slowing), and gait impairment, known as the "cardinal features" of the disease. Additional features can include freezing of gait, postural instability, speech difficulty, autonomic disturbances, sensory alterations, mood disorders, sleep dysfunction, cognitive impairment, and dementia (Table 449-1).

Pathologically, the hallmark features of PD are degeneration of dopaminergic neurons in the substantia nigra pars compacta (SNc), reduced striatal dopamine, and intracytoplasmic proteinaceous inclusions known as Lewy bodies that primarily contain the protein alpha synuclein (Fig. 449-1). While interest has primarily focused on the dopamine system, neuronal degeneration with inclusion body formation can also affect cholinergic neurons of the nucleus basalis of Meynert (NBM), norepinephrine neurons of the locus coeruleus (LC), serotonin neurons in the raphe nuclei of the brainstem, and neurons of the olfactory system, cerebral hemispheres, spinal cord, and peripheral autonomic nervous system. This "nondopaminergic" pathology is likely responsible for the development of nondopaminergic clinical features listed in Table 449-1 characterized by their lack of satisfactory response to dopaminergic replacement therapy. There is evidence that Lewy body pathology first begins in the peripheral autonomic nervous system, olfactory system, and dorsal motor nucleus of the vagus nerve in the lower brainstem, and then spreads in a predictable and sequential manner to affect the upper brainstem and cerebral hemispheres

(Braak staging). These studies suggest that degeneration of dopamine neurons develops in a mid-stage of the disease. Indeed, epidemiologic studies suggest that clinical symptoms reflecting this nondopaminergic degeneration, such as constipation, anosmia, rapid eye movement (REM) behavior sleep disorder, and cardiac denervation, can precede the onset of the classic motor features of PD.

DIFFERENTIAL DIAGNOSIS

Parkinsonism is a generic term that is used to define a syndrome manifest as bradykinesia with rigidity and/or tremor. It has a differential diagnosis (Table 449-2) that reflects damage to different components of the basal ganglia. The basal ganglia are comprised of a group of subcortical nuclei that include the striatum (putamen and caudate nucleus), subthalamic nucleus (STN), globus pallidus pars externa (GPe), globus pallidus pars interna (GPi), and the SNc (Fig. 449-2). Among the different forms of parkinsonism, PD is the most common (approximately 75% of cases). Historically, PD was diagnosed based on the presence of two of three parkinsonian features (tremor, rigidity, bradykinesia). However, postmortem studies found a 24% error rate when diagnosis was based on these criteria. Clinicopathologic correlation studies subsequently determined that parkinsonism associated with rest tremor, asymmetry, and a good response to levodopa was more likely to predict the correct pathologic diagnosis. With these revised criteria (known as the U.K. Brain Bank Criteria), a clinical diagnosis of PD is confirmed pathologically in as many as 99% of cases. A more complete definition of PD is now needed to incorporate the fact that there is widespread pathology beyond the dopaminergic system, nondopamine and nonmotor clinical features, and a premotor stage of the disease.

Imaging of the brain dopamine system in PD with positron emission tomography (PET) or single-photon emission computed tomography (SPECT) shows reduced uptake of striatal dopaminergic markers, particularly in the posterior putamen with relative sparing of the caudate nucleus (Fig. 449-3), reflecting the degeneration of nigrostriatal dopamine neurons. Imaging can be useful in patients where there is diagnostic uncertainty (e.g., dystonic tremor, essential tremor) or in research studies, but is rarely necessary in routine practice because the diagnosis can usually be established on clinical criteria alone. This may change in the future when there is a disease-modifying therapy and it is important to make the diagnosis as early as possible. Genetic testing is not routinely used at present, but can be helpful for identifying at-risk individuals in a research setting. Mutations of the *LRRK2* gene (see below) have attracted particular interest because they are the commonest cause of familial PD and are responsible for approximately 1% of typical sporadic cases of the disease. Mutations in *LRRK2* are a particularly common cause of PD in Ashkenazi Jews and North African Berber Arabs. The penetrance of the most common *LRRK2* mutation ranges from 28 to 74% and is strongly correlated to the age of the carrier, with 50% affected by age 60 years. Mutations in the *parkin* gene should be considered in patients with onset prior to 40 years of age.

Atypical and Secondary Parkinsonism Atypical parkinsonism refers to a group of neurodegenerative conditions that usually are associated with more widespread neurodegeneration than is found in PD (often involvement of striatum and/or globus pallidus as well as the SNc). As a group, they present with parkinsonism (rigidity and bradykinesia) but have a slightly different clinical picture than PD, reflecting differences in the underlying pathology. In these conditions, parkinsonism is typically characterized by early speech and gait impairment, absence of rest tremor, no motor asymmetry, poor or no response to levodopa, and an aggressive clinical course. In the early stages, they may show some modest benefit from levodopa and be difficult to distinguish from PD. Pathologically, neurodegeneration typically involves degeneration of the SNc but occurs without Lewy bodies (see below for individual conditions). Neuroimaging of the dopamine system is usually not helpful, because dopamine depletion can be seen in both PD and atypical parkinsonism. By contrast, metabolic imaging of the basal ganglia/thalamus network (using 2-F-deoxyglucose PET) may be helpful, showing a pattern of decreased activity in the GPi with increased activity in the thalamus, the reverse of what is seen in PD.

TABLE 449-1 CLINICAL FEATURES OF PARKINSON'S DISEASE

Cardinal Motor Features	Other Motor Features	Nonmotor Features
Bradykinesia	Micrographia	Anosmia
Rest tremor	Masked facies (hypomimia)	Sensory disturbances (e.g., pain)
Rigidity	Reduced eye blinking	Mood disorders (e.g., depression)
Gait disturbance/postural instability	Soft voice (hypophonia)	Sleep disturbances (e.g., RBD)
	Dysphagia	Autonomic disturbances
	Freezing	Orthostatic hypotension
		Gastrointestinal disturbances
		Genitourinary disturbances
		Sexual dysfunction
		Cognitive impairment (MCI/dementia)

Abbreviations: RBD, rapid eye movement behavior disorder; MCI, mild cognitive impairment.

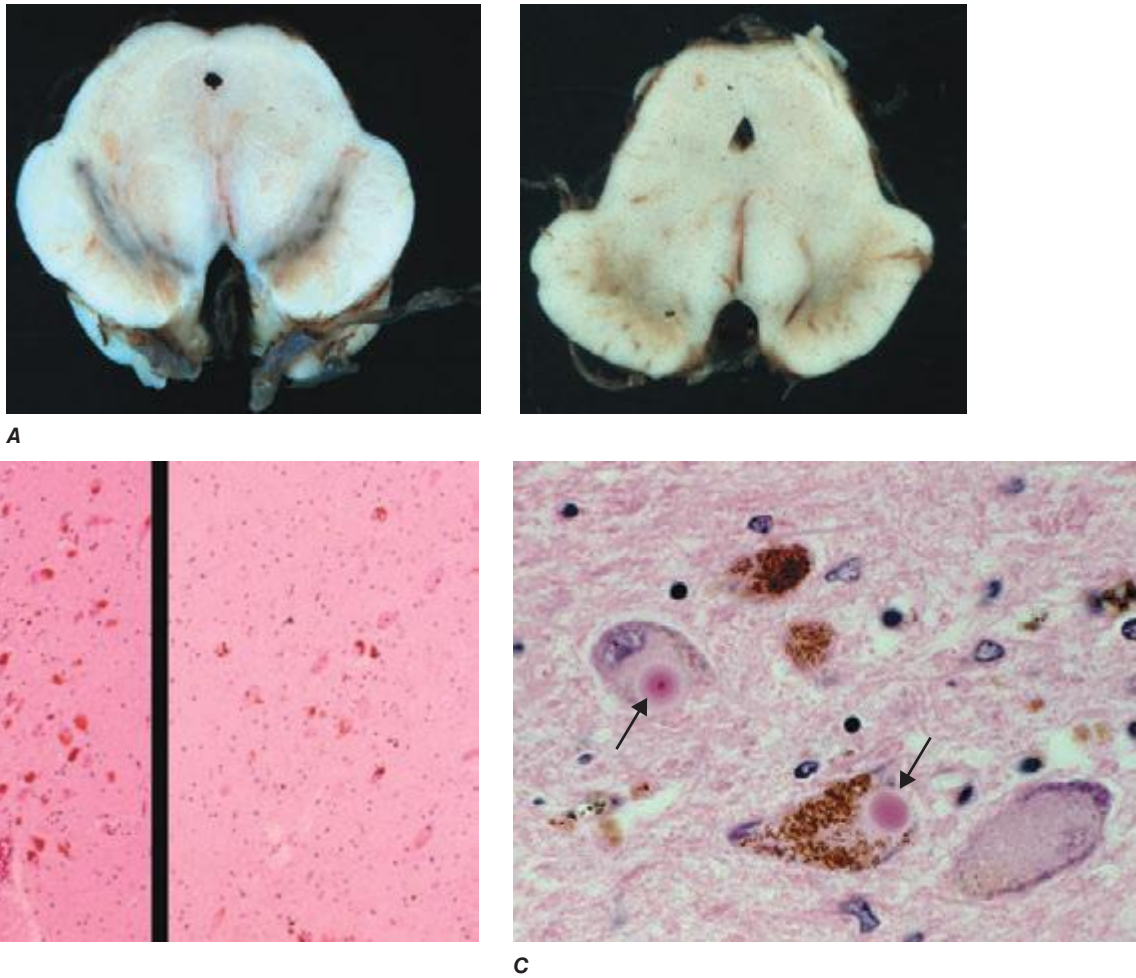


FIGURE 449-1 Pathologic specimens from a patient with Parkinson's disease (PD) compared to a normal control demonstrating (A) reduction of pigment in SNc in PD (*right*) versus control (*left*), (B) reduced numbers of cells in SNc in PD (*right*) compared to control (*left*), and (C) Lewy bodies (*arrows*) within melanized dopamine neurons in PD. SNc, substantia nigra pars compacta.

Multiple-system atrophy (MSA) manifests as a combination of parkinsonian, cerebellar, and autonomic features and can be divided into a predominant parkinsonian (MSA-p) or cerebellar (MSA-c) form. Clinically, MSA is suspected when a patient presents with atypical parkinsonism in conjunction with cerebellar signs and/or early and prominent autonomic dysfunction, usually orthostatic hypotension ([Chap. 454](#)). Pathologically, MSA is characterized by degeneration of the SNc, striatum, cerebellum, and inferior olivary nuclei coupled with characteristic glial cytoplasmic inclusions (GCIs) that stain for α -synuclein. Magnetic resonance imaging (MRI) can show pathologic iron accumulation in the striatum on T2-weighted scans, high signal change in the region of the external surface of the putamen (putaminal rim) in MSA-p, or cerebellar and brainstem atrophy (the pontine “hot

cross buns” sign [[Fig. 454-2](#)]) in MSA-c. Mutations in the *CoQ2* gene encoding parahydroxybenzoate-polyprenyl transferase, an enzyme involved in the biosynthesis of coenzyme Q10 (CoQ10), a cofactor of the mitochondrial respiratory chain, have been identified in familial and sporadic forms of MSA.

Progressive supranuclear palsy (PSP) is a form of atypical parkinsonism that is characterized by slow ocular saccades, eyelid apraxia, and restricted eye movements with particular impairment of downward gaze. Patients frequently experience hyperextension of the neck with early gait disturbance and falls. In later stages, speech and swallowing difficulty and cognitive impairment become evident. MRI may reveal a characteristic atrophy of the midbrain with relative preservation of the pons, the “hummingbird sign” on midsagittal images. Pathologically,

TABLE 449-2 DIFFERENTIAL DIAGNOSIS OF PARKINSONISM

Parkinson's Disease	Atypical Parkinsonism	Secondary Parkinsonism	Other Neurodegenerative Disorders
Genetic	Multiple-system atrophy (MSA)	Drug-induced	Wilson's disease
Sporadic	Cerebellar type (MSA-c)	Tumor	Huntington's disease
Dementia with Lewy bodies	Parkinson type (MSA-p)	Infection	Neurodegeneration with brain iron accumulation
	Progressive supranuclear palsy	Vascular	SCA 3 (spinocerebellar ataxia)
	Corticobasal ganglionic degeneration	Normal-pressure hydrocephalus	Fragile X-associated ataxia-tremor-parkinsonism
	Frontotemporal dementia	Trauma	Prion disease
		Liver failure	Dystonia-parkinsonism (DYT3)
		Toxins (e.g., carbon monoxide, manganese, MPTP, cyanide, hexane, methanol, carbon disulfide)	Alzheimer's disease with parkinsonism

Abbreviations: MPTP, 1-methyl-4-phenyl-1,2,5,6-tetrahydropyridine.

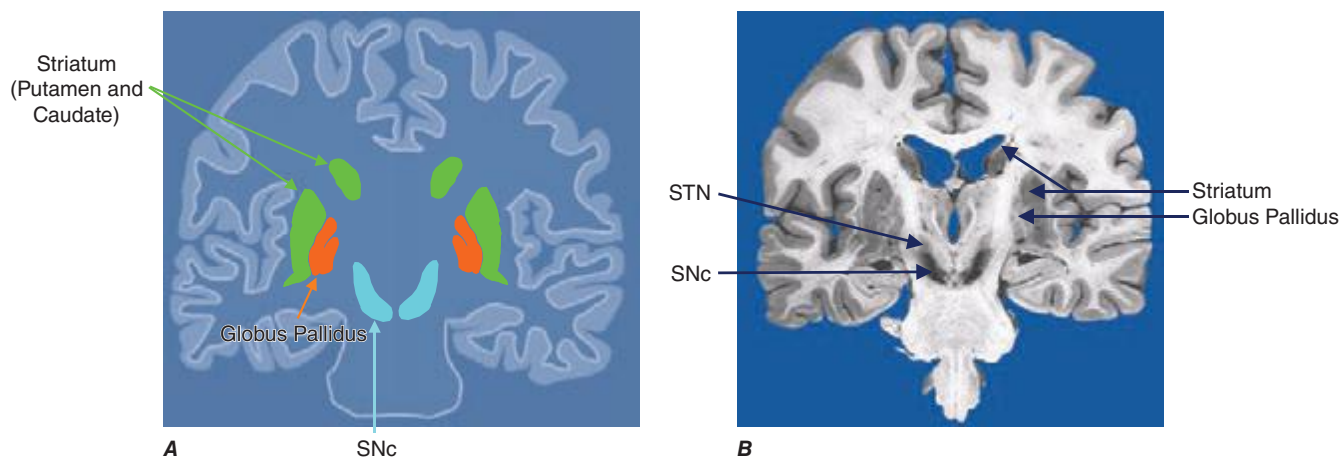


FIGURE 449-2 Basal ganglia nuclei. Schematic (A) and postmortem (B) coronal sections illustrating the various components of the basal ganglia. SNc, substantia nigra pars compacta; STN, subthalamic nucleus.

PSP is characterized by degeneration of the SNc, striatum, subthalamic nucleus, midline thalamic nuclei, and pallidum along with neurofibrillary tangles and inclusions that stain for the tau protein.

Corticobasal ganglionic degeneration is less common and is usually manifest by asymmetric dystonic contractions and clumsiness of one hand coupled with cortical sensory disturbances manifest as apraxia, agnosia, focal limb myoclonus, or alien limb phenomenon (where

the limb assumes a position in space without the patient being aware of it). Dementia may occur at any stage of the disease. Both cortical and basal ganglia features are required to make this diagnosis. MRI frequently shows asymmetric cortical atrophy. Pathologic findings include achromatic neuronal degeneration with tau deposits. Because other disorders such as PSP can present with a similar clinical picture, the term corticobasal ganglia syndrome should be used until a precise diagnosis can be confirmed pathologically.

Secondary parkinsonism can occur as a result of drugs, stroke, tumor, infection, or exposure to toxins such as carbon monoxide or manganese. Dopamine-blocking agents such as the neuroleptics are the commonest cause of secondary parkinsonism. These drugs are most widely used in psychiatry, *but physicians should be aware that drugs such as metoclopramide and chlorpromazine, which are primarily used to treat gastrointestinal problems, are also neuroleptic agents* and common causes of secondary parkinsonism (as well as acute and tardive dyskinesias; see below). Other drugs that can cause secondary parkinsonism include tetrabenazine, calcium channel blockers (flunarizine, cinnarizine), amiodarone, and lithium.

Finally, parkinsonism can be seen as a feature of other degenerative disorders such as Wilson's disease, Huntington's disease (especially the juvenile form known as the Westphal variant), dopa-responsive dystonia, and neurodegenerative disorders with brain iron accumulation such as pantothenate kinase (PANK)-associated neurodegeneration (formerly known as Hallervorden-Spatz disease).

Some features that suggest parkinsonism might be due to a condition other than PD are shown in [Table 449-3](#).

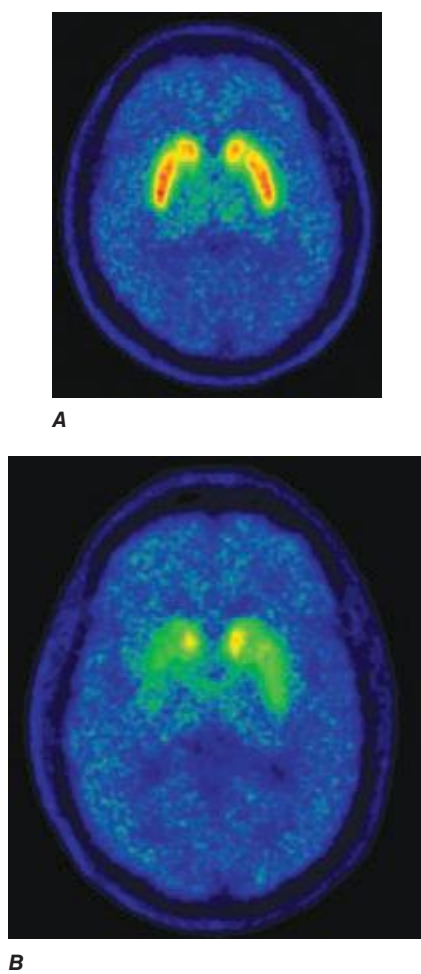


FIGURE 449-3 [^{11}C]Dihydrotetrabenazine positron emission tomography (a marker of VMAT2) in healthy control (A) and Parkinson's disease (B) patient. Note the reduced striatal uptake of tracer, which is most pronounced in the posterior putamen and tends to be asymmetric. (Courtesy of Dr. Jon Stoessl.)

ETIOLOGY AND PATHOGENESIS

Most PD cases occur sporadically (~85–90%) and are of unknown cause. Twin studies suggest that environmental factors likely play an important role in patients older than 50 years, with genetic factors being more important in younger patients. Epidemiologic studies also suggest increased risk with exposure to pesticides, rural living, and drinking well water and reduced risk with cigarette smoking and caffeine. However, no environmental factor has yet been proven to cause typical PD. The environmental hypothesis received support with the demonstration in the 1980s that MPTP (1-methyl-4-phenyl-1,2,5,6-tetrahydropyridine), a byproduct of the illicit manufacture of a heroin-like drug, caused a PD-like syndrome in addicts in northern California. MPTP is transported to the central nervous system, where it is oxidized to form MPP⁺, a mitochondrial toxin that is selectively taken up by, and damages, dopamine neurons. However, MPTP or MPTP-like compounds have not been linked to sporadic PD.

About 10–15% of cases are familial in origin, and multiple specific mutations and gene associations have been identified ([Table 449-4](#)). Genetic factors have also been linked to sporadic cases, with several typical PD cases found to carry the *LRRK2* mutation, and genome-wide association studies (GWAS) implicating alpha synuclein, tau, and

TABLE 449-3 FEATURES SUGGESTING AN ATYPICAL OR SECONDARY CAUSE OF PARKINSONISM

Symptoms/Signs	Alternative Diagnosis to Consider
History	
Early speech and gait impairment (Lack of tremor, lack of motor asymmetry)	Atypical parkinsonism
Exposure to neuroleptics	Drug-induced parkinsonism
Onset prior to age 40	Genetic form of PD
Liver disease	Wilson's disease, non-Wilsonian hepatolenticular degeneration
Early hallucinations and dementia with later development of PD features	Dementia with Lewy bodies
Diplopia, impaired down gaze	PSP
Poor or no response to an adequate trial of levodopa	Atypical or secondary parkinsonism
Physical Exam	
Dementia as first or early feature	Dementia with Lewy bodies
Prominent orthostatic hypotension	MSA-p
Prominent cerebellar signs	MSA-c
Slow saccades with impaired down gaze	PSP
High-frequency (6–10 Hz) symmetric postural tremor with a prominent kinetic component	Essential tremor

Abbreviations: MSA-c, multiple-system atrophy–cerebellar type; MSA-p, multiple-system atrophy–Parkinson's type; PD, Parkinson's disease; PSP, progressive supranuclear palsy.

HLA as risk factors. It has been proposed that most cases of PD may be due to a “double hit” involving an interaction between a gene mutation that induces susceptibility coupled with exposure to a toxic environmental factor that may induce epigenetic or somatic DNA alterations. In this scenario, both factors are required for PD to ensue, while the presence of either one alone is not sufficient to cause the disease.

Several factors have been implicated in the pathogenesis of cell death in PD, including oxidative stress, inflammation, mitochondrial dysfunction, and proteolytic stress. Recent studies have demonstrated that with aging dopamine neurons switch from sodium to calcium pacing through calcium channels, potentially making these high-energy neurons vulnerable to calcium-mediated neurotoxicity. Whatever the pathogenic mechanism, cell death appears to occur, at least in part, by way of a signal-mediated apoptotic or “suicidal” process.

TABLE 449-4 GENETIC CAUSES OF PARKINSON'S DISEASE

Name	Chromosome	Locus	Gene	Inheritance
Park 1	Chr 4	q21-23	<i>α-Synuclein</i>	AD
Park 2	Chr 6	q25-27	<i>Parkin</i>	AR
Park 3	Chr 2	p13	Unknown	AD
Park 4	Chr 4	q21-23	<i>α-Synuclein</i>	AD
Park 5	Chr 4	p14	<i>UCHL-1</i>	AD
Park 6	Chr 1	p35-36	<i>PINK-1</i>	AR
Park 7	Chr 1	p36	<i>DJ-1</i>	AR
Park 8	Chr 12	p11-q13	<i>LRRK2</i>	AR/Sp
Park 9	Chr 1	p36	<i>ATP13A2</i>	AR
Park 10	Chr 1	p32	Unknown	Sp
Park 11	Chr 2	q36-37	<i>GIGYF2</i>	AD
Park 12	Chr X	q21-25	Unknown	Sp
Park 13	Chr 2	p13	<i>Omi/HtrA2</i>	AD
Park 14	Chr 22	q13	<i>PLA2G6</i>	AR
Park 15	Chr 22	q12-13	<i>FBX07</i>	AR
Park 16	Chr 1	q32	Unknown	Sp

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; Chr, chromosome; Sp, sporadic.

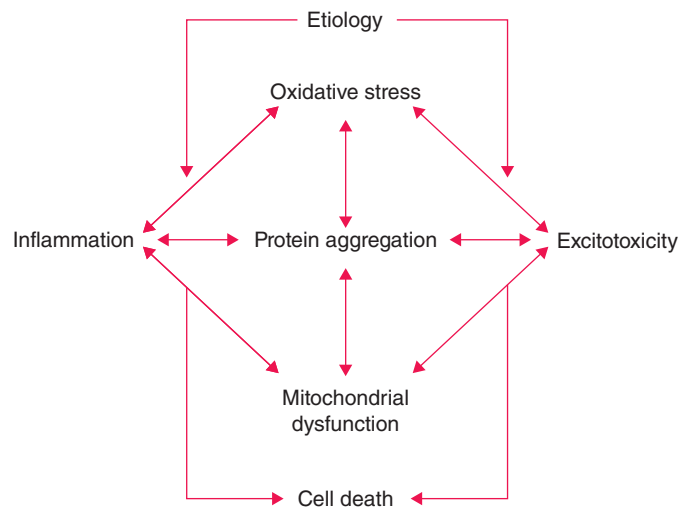


FIGURE 449-4 Schematic representation of how pathogenetic factors implicated in Parkinson's disease interact in a network manner, ultimately leading to cell death. This figure illustrates how interference with any one of these factors may not necessarily stop the cell death cascade. (Adapted from CW Olanow: *Movement Disorders* 22:S-335, 2007.)

Each of these mechanisms offers a potential target for neuroprotective drugs. However, it is not clear which of these factors is primary, if the mechanism is the same in each individual case, if they act by way of a network such that a cocktail of agents might be required to provide neuroprotection, or if the findings to date merely represent epiphenomena unrelated to the true cause of cell death that remains undiscovered (Fig. 449-4).

Gene mutations may not cause all cases of PD, but may be helpful in pointing to specific pathogenic pathways and mechanisms that are central to a neurodegenerative process that might be relevant to all forms of the disease. To date, most interest has focused on pathways implicated by mutations in *α-synuclein*, *LRRK2*, and *PINK1/Parkin*.

Most interest has focused on *α-synuclein*. Mutations in *α-synuclein* cause rare familial forms of PD, and *α-synuclein* constitutes the major component of Lewy bodies in patients with sporadic PD (Fig. 449-1). Furthermore, duplication or triplication of the wild-type *α-synuclein* can also cause a form of PD, indicating that increased production of the normal protein alone can cause the disease. More recently, Lewy pathology was discovered to have developed in healthy embryonic dopamine neurons that had been implanted into the striatum of PD patients, suggesting that the abnormal protein had transferred from affected cells to healthy unaffected dopamine neurons. Based on these findings, it has been proposed that *α-synuclein* is a prion and PD is a prion disorder. Here it is proposed that, like the prion protein PrP^C, *α-synuclein* can misfold to form β -rich sheets, generate toxic oligomers and aggregates, polymerize to form amyloid plaques (i.e., Lewy bodies), cause neurodegeneration, and spread to involve unaffected neurons. Indeed, injection of *α-synuclein* fibrils into the striatum promotes the development of Lewy pathology in host neurons, neurodegeneration, behavioral abnormalities, and the spread of *α-synuclein* pathology to anatomically connected sites. Further support for this hypothesis comes from the demonstration that inoculation of *α-synuclein* derived from human Lewy bodies induces widespread Lewy pathology in mice and primates. Collectively, this evidence supports the possibility that neuroprotective therapies for PD might be developed based on inhibiting accumulation or accelerating removal of *α-synuclein* aggregates.

Mutations in the glucocerebrosidase (*GBA*) gene associated with Gaucher's disease numerically represent the most important risk factor for the development of PD. While the responsible mechanism is not precisely known, it is noteworthy that *GBA* mutations are associated with altered autophagy and lysosomal function and could impair the clearance of *α-synuclein*.

Six different *LRRK2* mutations have been linked to PD, with Gly2019Ser being the most common. The mechanism responsible for cell death with this mutation is not known but is thought to involve changes in kinase activity with altered phosphorylation of target proteins (including autophosphorylation) and possibly lysosomal dysfunction. Kinase inhibitors can block toxicity associated with *LRRK2* mutations in laboratory models, and there has been much interest in developing drugs directed at this target. However, kinase inhibitors are likely to be toxic, the physiologic role of *LRRK2* is not known, and the large majority of PD patients do not carry a *LRRK2* mutation.

Mutations in *PINK1* and *parkin* have implicated mitochondrial dysfunction as a possible cause of PD. Recent studies suggest a role for parkin and PINK1 proteins in the turnover and clearance of damaged mitochondria (mitophagy), and mutations in *parkin* and *PINK1* cause mitochondrial dysfunction in transgenic animals that can be corrected with overexpression of parkin. This is a particularly attractive target because postmortem studies in PD patients show a defect in complex I of the respiratory chain in SNc neurons.

Thus, evidence is accumulating that genetics plays an important role in both familial and “sporadic” forms of PD. It is anticipated that better understanding of the pathways responsible for cell death caused by these mutations will permit the development of more relevant animal models of PD and targets for the development of neuroprotective drugs.

PATHOPHYSIOLOGY OF PD

The classic model of the organization of the basal ganglia in the normal and PD states is provided in Fig. 449-5. With respect to motor function, a series of neuronal circuits or loops link the basal ganglia nuclei with corresponding cortical motor regions in a somatotopic manner. The striatum is the major input region of the basal ganglia, while the GPi and SNr are the major output regions. The input and output regions are connected via direct and indirect pathways that have reciprocal effects on the activity of the output pathway. The output of the basal ganglia provides inhibitory (GABAergic) tone to thalamic and brainstem neurons that in turn connect to motor systems

in the cerebral cortex and spinal cord that control motor function. Physiologically, decreased neuronal activity in the GPi/SNr is associated with movement facilitation and vice versa. Dopaminergic projections from SNc neurons serve to modulate neuronal firing and to stabilize the basal ganglia network. The basal ganglia and similar cortical loops are now thought to also play an important role in regulating normal behavioral, emotional, and cognitive functions.

In PD, dopamine denervation with loss of dopaminergic tone leads to increased firing of neurons in the STN and GPi, excessive inhibition of the thalamus, reduced activation of cortical motor systems, and the development of parkinsonian features (Fig. 449-5). The current role of surgery in the treatment of PD is based on this model, which predicted that lesions or high-frequency stimulation of the STN or GPi might reduce this neuronal overactivity and improve PD features.

TREATMENT PARKINSON'S DISEASE

LEVODOPA

Since its introduction in the late 1960s, levodopa has been the mainstay of therapy for PD. Experiments in the late 1950s by Carlsson demonstrated that blocking dopamine uptake with reserpine caused rabbits to become parkinsonian; this could be reversed with the dopamine precursor, levodopa. Subsequently, Hornykiewicz demonstrated a dopamine deficiency in the striatum of PD patients and suggested the potential benefit of dopaminergic replacement therapy. Dopamine does not cross the blood-brain barrier (BBB), so clinical trials were initiated with levodopa, a precursor of dopamine. Studies over the course of the next decade confirmed the value of levodopa and revolutionized the treatment of PD.

Levodopa is routinely administered in combination with a peripheral decarboxylase inhibitor to prevent its peripheral metabolism to dopamine and the development of nausea and vomiting due to activation of dopamine receptors in the area postrema that are not protected by the BBB. In the United States, levodopa is combined with the decarboxylase inhibitor carbidopa (Sinemet), whereas in

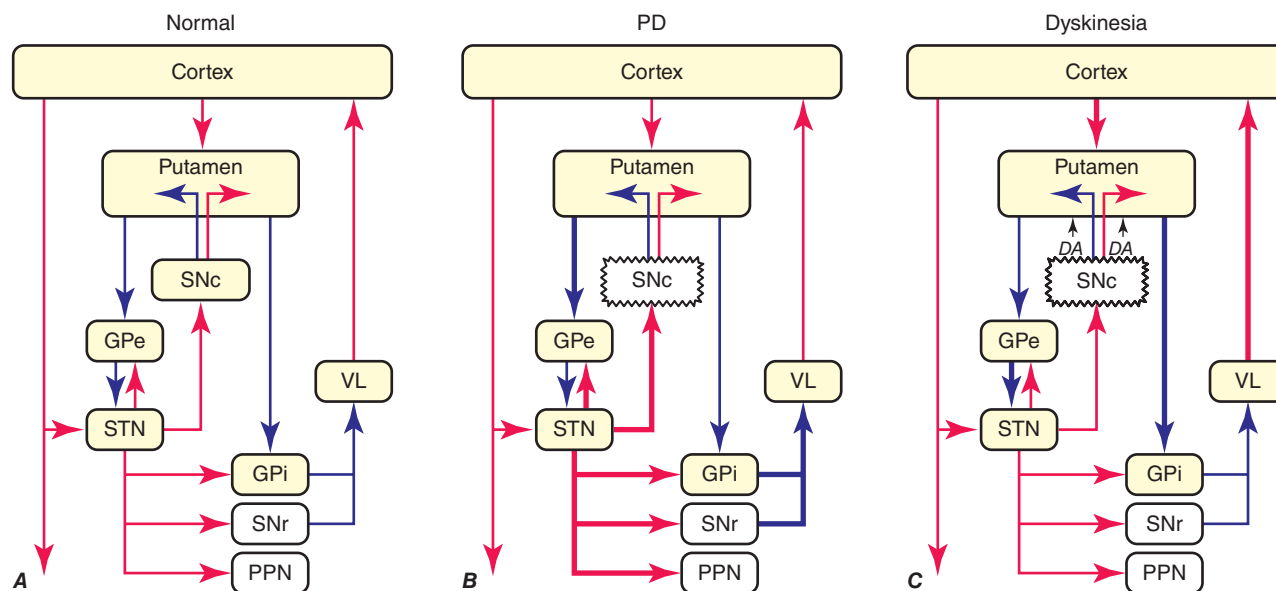


FIGURE 449-5 Basal ganglia organization. Classic model of the organization of the basal ganglia in the normal (A), Parkinson's disease (PD) (B), and levodopa-induced dyskinesia (C) state. Inhibitory connections are shown as blue arrows and excitatory connections as red arrows. The striatum is the major input region and receives its major input from the cortex. The GPi and SNr are the major output regions, and they project to the thalamocortical and brainstem motor regions. The striatum and GPi/SNr are connected by direct and indirect pathways. This model predicts that parkinsonism results from increased neuronal firing in the STN and GPi and that lesions or DBS of these targets might provide benefit. This concept led to the rationale for surgical therapies for PD. The model also predicts that dyskinesia results from decreased firing of the output regions, resulting in excessive cortical activation by the thalamus. This component of the model is not completely correct because lesions of the GPi ameliorate rather than increase dyskinesia in PD, suggesting that firing frequency is just one of the components that lead to the development of dyskinesia. DBS, deep brain stimulation; GPe, external segment of the globus pallidus; GPi, internal segment of the globus pallidus; PPN, pedunculopontine nucleus; SNc, substantia nigra, pars compacta; SNr, substantia nigra, pars reticulata; STN, subthalamic nucleus; VL, ventrolateral thalamus. (Derived from JA Obeso et al: *Trends Neurosci* 23:S8, 2000.)

many other countries, it is combined with benserazide (Madopar). Levodopa is also available in controlled-release formulations as well as in combination with a catechol-*O*-methyltransferase (COMT) inhibitor (see below). Levodopa remains the most effective symptomatic treatment for PD and the gold standard against which new therapies are compared. No current medical or surgical treatment provides antiparkinsonian benefits superior to what can be achieved with levodopa. Levodopa benefits the classic motor features of PD, prolongs independence and employability, improves quality of life, and increases life span. Almost all PD patients experience improvement, and failure to respond to an adequate trial should cause the diagnosis to be questioned.

There are, however, important limitations of levodopa therapy. Acute dopaminergic side effects include nausea, vomiting, and orthostatic hypotension. These are usually transient and can generally be avoided by gradual titration. If they persist, they can be treated with additional doses of a peripheral decarboxylase inhibitor (e.g., carbidopa) or a peripheral dopamine-blocking agent such as domperidone (not available in the United States). More important are motor complications (see below) that develop in the majority of patients treated long-term with levodopa. In addition, the disease continues to progress, and features such as falling, freezing, autonomic dysfunction, sleep disorders, and dementia may emerge with disease progression that are not adequately controlled by levodopa. Indeed, these nondopaminergic features (especially falling and dementia) are the primary source of disability and the main reason for nursing home placement for patients with advanced PD.

Levodopa-induced motor complications consist of fluctuations in motor response (“on” episodes when the drug is working and “off” episodes when parkinsonian features return) and involuntary movements known as dyskinesias (Fig. 449-6). When patients initially take levodopa, benefits are long-lasting (many hours) even though the drug has a relatively short half-life (60–90 min). With continued treatment, however, the duration of benefit following an individual dose becomes progressively shorter until it approaches the half-life of the drug. This loss of benefit is known as the *wearing-off effect*. In more severe cases, patients may experience a delay in turning on (delayed-on) or no response at all to a given dose (non-on). Dyskinesias tend to occur at the time of levodopa peak plasma concentration and maximal clinical benefit (peak-dose dyskinesia). They are usually choreiform in nature but can manifest as dystonic movements, myoclonus, or other movement disorders. They are not troublesome when mild, but can be disabling when severe, and can limit the ability to fully use levodopa to control PD features. In more advanced states, patients may cycle between “on” periods complicated by disabling dyskinesias and “off” periods in which they suffer from severe parkinsonism and painful dystonic postures. Patients may also experience “diphasic dyskinesias,” which occur as the levodopa dose begins to take effect and again as it wears off. These

dyskinesias typically consist of transient, stereotypic, rhythmic movements that predominantly involve the lower extremities and are frequently associated with parkinsonism in other body regions. They can be relieved by increasing the dose of levodopa, although higher doses may induce more severe peak-dose dyskinesia.

The cause of levodopa-induced motor complications is not precisely known. They are more likely to occur in females, younger individuals with more severe disease, and with the use of higher doses (mg/kg) of levodopa. The classic model of the basal ganglia has been useful for understanding the origin of motor features in PD, but has proved less valuable for understanding levodopa-induced dyskinesias (Fig. 449-5). The model predicts that dopamine replacement might excessively inhibit the pallidal output system, thereby leading to increased thalamocortical activity, enhanced stimulation of cortical motor regions, and the development of dyskinesia. However, lesions of the pallidum that completely destroy its output are associated with amelioration rather than induction of dyskinesia as suggested by the classic model. It is now thought that dyskinesia results from levodopa-induced alterations in the GPi neuronal firing pattern (pauses, bursts, synchrony, etc.) and oscillatory activity, and not simply the firing frequency alone. This in turn leads to the transmission of misinformation from pallidum to thalamus/cortex, resulting in dyskinesia. Surgical lesions or high-frequency stimulation might ameliorate dyskinesia by interfering with (blocking or masking) this abnormal neuronal activity and preventing the transfer of misinformation to motor systems.

Current information suggests that altered neuronal firing patterns and motor complications relate to nonphysiologic levodopa replacement. Striatal dopamine levels are normally maintained at a relatively constant level. In PD, dopamine neurons degenerate and striatal dopamine is dependent on the peripheral availability of levodopa. Intermittent doses of short-acting levodopa result in fluctuating plasma levels because of variability in transit of the drug from the stomach to the duodenum where it is absorbed and the short half-life of the drug. This variability results in exposure of dopamine receptors to pathologically high and low concentrations of dopamine. It has been hypothesized that more continuous delivery of levodopa might prevent the development of motor complications. Indeed, a recent controlled study demonstrated that continuous intrainestinal infusion of levodopa/carbidopa intestinal gel (Duodopa) is associated with significant improvement in “off” time and in “on” time without dyskinesia in advanced PD patients compared with optimized standard oral levodopa.

Behavioral alterations can also be encountered in levodopa-treated patients. A dopamine dysregulation syndrome has been described where patients have a craving for levodopa and take frequent and unnecessary doses of the drug in an addictive manner. PD patients taking high doses of levodopa can develop purposeless, stereotyped behaviors such as the meaningless assembly and

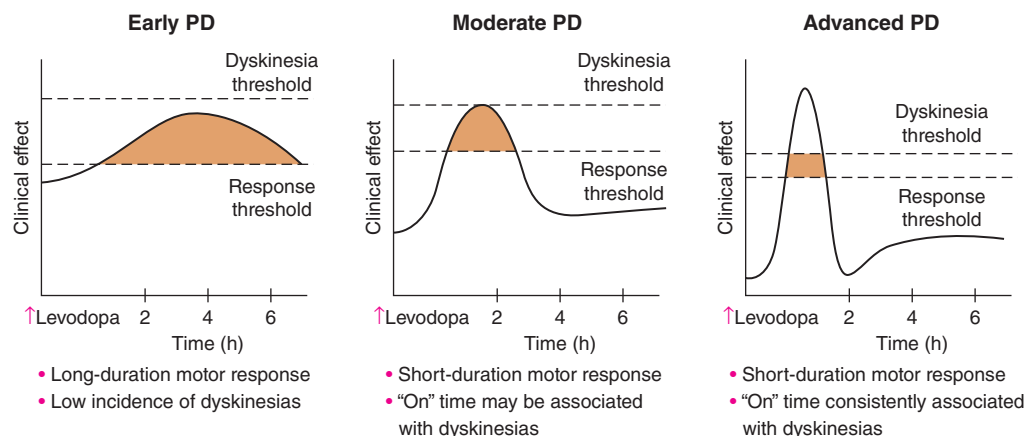


FIGURE 449-6 Changes in motor response associated with chronic levodopa treatment. Levodopa-induced motor complications. Schematic illustration of the gradual shortening of the duration of a beneficial motor response to levodopa (wearing off) and the appearance of dyskinesias complicating “on” time. PD, Parkinson’s disease.

disassembly or collection and sorting of objects. This is known as punding, a term taken from the Swedish description of the meaningless behaviors seen in chronic amphetamine users. Hypersexuality and other impulse-control disorders are occasionally encountered with levodopa, although these are more commonly seen with dopamine agonists.

DOPAMINE AGONISTS

Dopamine agonists are a diverse group of drugs that act directly on dopamine receptors. Unlike levodopa, they do not require metabolism to an active product and do not undergo oxidative metabolism. Initial dopamine agonists were ergot derivatives (e.g., bromocriptine, pergolide, cabergoline) and were associated with ergot-related side effects, including cardiac valvular damage. They have largely been replaced by a second generation of nonergot dopamine agonists (e.g., pramipexole, ropinirole, rotigotine). In general, dopamine agonists do not have comparable efficacy to levodopa. They were initially introduced as adjuncts to levodopa to enhance motor function and reduce "off" time in fluctuating patients. Subsequently, it was shown that dopamine agonists, possibly because they are relatively long-acting, are less prone than levodopa to induce dyskinesia. For this reason, many physicians initiate therapy with a dopamine agonist, although supplemental levodopa is eventually required in virtually all patients. Both ropinirole and pramipexole are available as orally administered immediate (tid) and extended-release (qd) formulations. Rotigotine is administered as a once-daily transdermal patch. Apomorphine is a dopamine agonist with efficacy comparable to levodopa, but it must be administered parenterally and has a very short half-life and duration of activity (45 min). It is generally administered by injection as a rescue agent for the treatment of severe "off" episodes. Apomorphine can also be administered by continuous subcutaneous infusion and has been demonstrated to reduce both "off" time and dyskinesia in advanced patients. However, this approach has not been approved in the United States.

Dopamine agonist use is associated with a variety of side effects. Acute side effects are primarily dopaminergic and include nausea, vomiting, and orthostatic hypotension. As with levodopa, these can usually be avoided by slow titration. Side effects associated with chronic use include hallucinations and cognitive impairment. Sedation with sudden unintended episodes of falling asleep while driving a motor vehicle have been reported. Patients should be informed about this potential problem and should not drive when tired. Dopamine agonists can also be associated with impulse-control disorders, including pathologic gambling, hypersexuality, and compulsive eating and shopping. The precise cause of these problems, and why they appear to occur more frequently with dopamine agonists than levodopa, remains to be resolved, but reward systems associated with dopamine and alterations in the ventral striatum and orbitofrontal regions have been implicated. In general, chronic side effects are dose-related and can be avoided or minimized with lower doses. Injections of apomorphine and patch delivery of rotigotine can be complicated by development of skin lesions at sites of administration.

MAO-B INHIBITORS

Inhibitors of monoamine oxidase type B (MAO-B) block central dopamine metabolism and increase synaptic concentrations of the neurotransmitter. Selegiline and rasagiline are relatively selective suicide inhibitors of the MAO-B enzyme. Clinically, MAO-B inhibitors provide antiparkinsonian benefits when used as monotherapy in early disease and reduced "off" time when used as an adjunct to levodopa in patients with motor fluctuations. MAO-B inhibitors are generally safe and well tolerated. They may increase dyskinesia in levodopa-treated patients, but this can usually be controlled by down-titrating the dose of levodopa. Inhibition of the MAO-A isoform prevents metabolism of tyramine in the gut, leading to a potentially fatal hypertensive reaction known as a "cheese effect" because it can be precipitated by foods rich in tyramine such as some cheeses, aged meats, and red wine. Selegiline and rasagiline

do not functionally inhibit MAO-A and are not associated with a cheese effect with doses typically used in clinical practice. There are theoretical risks of a serotonin reaction in patients receiving concomitant selective serotonin reuptake inhibitor (SSRI) antidepressants, but these are rarely encountered.

Interest in MAO-B inhibitors has also focused on their potential to have disease-modifying effects. MPTP toxicity can be prevented experimentally by coadministration of an MAO-B inhibitor that blocks its conversion to the toxic pyridinium ion MPP⁺. MAO-B inhibitors also have the potential to block the oxidative metabolism of dopamine and prevent oxidative stress. In addition, both selegiline and rasagiline incorporate a propargyl ring within their molecular structure that provides antiapoptotic effects in laboratory models. The DATATOP study showed that selegiline significantly delayed the time until the emergence of disability, necessitating the introduction of levodopa, in untreated PD patients. However, it could not be determined whether this was due to a neuroprotective effect that slowed disease progression or a symptomatic effect that merely masked ongoing neurodegeneration. More recently, the ADAGIO study demonstrated that early treatment with rasagiline 1 mg/d, but not 2 mg/d, provided benefits that could not be achieved when treatment with the same drug was initiated at a later time point, consistent with a disease-modifying effect; however, the long-term significance of these findings is uncertain.

COMT INHIBITORS

When levodopa is administered with a decarboxylase inhibitor, it is primarily metabolized in the periphery by COMT. Inhibitors of COMT increase the elimination half-life of levodopa and enhance its brain availability. Combining levodopa with a COMT inhibitor reduces "off" time and prolongs "on" time in fluctuating patients while enhancing motor scores. Two COMT inhibitors have been approved, tolcapone and entacapone. There is also a combination tablet of levodopa, carbidopa, and entacapone (Stalevo).

Side effects of COMT inhibitors are primarily dopaminergic (nausea, vomiting, increased dyskinesia) and can usually be controlled by down-titrating the dose of levodopa by 20–30%. Severe diarrhea has been described with tolcapone, and to a lesser degree with entacapone, and necessitates stopping the medication in 5–10% of individuals. Cases of fatal hepatic toxicity have been reported with tolcapone, and periodic monitoring of liver function is required. This problem has not been encountered with entacapone. Discoloration of urine can be seen with both COMT inhibitors due to accumulation of a metabolite, but it is of no clinical concern.

It has been proposed that initiating levodopa in combination with a COMT inhibitor to enhance its elimination half-life could provide more continuous levodopa delivery if administered at frequent intervals and reduce the risk of motor complications. While this result has been demonstrated in a preclinical MPTP model, and continuous infusion reduces both "off" time and dyskinesia in advanced PD patients, no benefit of initiating levodopa with a COMT inhibitor compared to levodopa alone was detected in early PD patients in the STRIDE-PD study. This may have been because the combination was not administered at frequent enough intervals to provide continuous levodopa availability. For now, the main value of COMT inhibitors continues to be in patients who experience motor fluctuations.

OTHER MEDICAL THERAPIES

Centrally acting anticholinergic drugs such as trihexyphenidyl and benzotropine were used historically for the treatment of PD, but they lost favor with the introduction of dopaminergic agents. Their major clinical effect is on tremor, although it is not certain that this benefit is superior to what can be obtained with agents such as levodopa and dopamine agonists. Still, they can be helpful in individual patients with severe tremor. Their use is limited particularly in the elderly, due to their propensity to induce a variety of side effects including urinary dysfunction, glaucoma, and particularly cognitive impairment.

TABLE 449-5 DRUGS COMMONLY USED FOR TREATMENT OF PARKINSON'S DISEASE^a

Agent	Available Dosages	Typical Dosing
Levodopa ^a		
Carbidopa/levodopa	10/100, 25/100, 25/250 mg	200–1000 mg levodopa/d 2–4 times/d
Benserazide/levodopa	25/100, 50/200 mg	
Carbidopa/levodopa CR	25/100, 50/200 mg	
Benserazide/levodopa MDS	25/200, 25/250 mg	
Parcopa	10/100, 25/100, 25/250	
Carbidopa/levodopa/entacapone	12.5/50/200, 18.75/75/200, 25/100/200, 31.25/125/200, 37.5/150/200, 50/200/200 mg	
Dopamine agonists		
Pramipexole	0.125, 0.25, 0.5, 1.0, 1.5 mg	0.25–1.0 mg tid
Pramipexole ER	0.375, 0.75, 1.5, 3.0, 4.5 mg	1–3 mg/d
Ropinirole	0.25, 0.5, 1.0, 3.0 mg	6–24 mg/d
Ropinirole XL	2, 4, 6, 8 mg	6–24 mg/d
Rotigotine patch	2-, 4-, 6-, 8-mg patches	4–24 mg/d
Apomorphine SC		2–8 mg
COMT inhibitors		
Entacapone	200 mg	200 mg with each levodopa dose
Tolcapone	100, 200 mg	100–200 mg tid
MAO-B inhibitors		
Selegiline	5 mg	5 mg bid
Rasagiline	0.5, 1.0 mg	1.0 mg QAM

^aTreatment should be individualized. Generally, drugs should be started in low doses and titrated to optimal dose.

Note: Drugs should not be withdrawn abruptly but should be gradually lowered or removed as appropriate.

Abbreviations: COMT, catechol-O-methyltransferase; MAO-B, monoamine oxidase type B; QAM, every morning.

Amantadine also has historical importance. Originally introduced as an antiviral agent, it was appreciated to also have antiparkinsonian effects that are now thought to be due to *N*-methyl-D-aspartate (NMDA) receptor antagonism. While some physicians use amantadine in patients with early disease for its mild symptomatic effects, it is most widely used as an antidyskinesia agent in patients with advanced PD. Indeed, it is the only oral agent that has been demonstrated in controlled studies to reduce dyskinesia without worsening parkinsonian features, although benefits may be relatively transient. Cognitive impairment is a major concern. Other side effects include livido reticularis and weight gain. Amantadine should always be discontinued gradually because patients can experience withdrawal-like symptoms.

Several new classes of drug are currently being investigated in an attempt to enhance antiparkinsonian effects, reduce off time, and treat or prevent dyskinesia. These include adenosine A_{2A} antagonists, nicotinic agonists, glutamate antagonists, and 5-HT_{1A} agonists.

A list of the major drugs and available dosage strengths is provided in [Table 449-5](#).

NEUROPROTECTION

Despite the many therapeutic agents available for the treatment of PD, patients continue to experience disease progression with intolerable disability. A neuroprotective therapy that slows or stops disease progression remains the major unmet therapeutic need in PD. As noted above, trials of certain drugs (e.g., selegiline and rasagiline)

have provided positive results consistent with a disease-modifying effect. However, it is not possible to determine if the positive results were due to neuroprotection with slowing of disease progression or confounding symptomatic effects that mask ongoing progression. CoQ10, a mitochondrial bioenhancer and antioxidant, attracted attention with a positive preliminary trial, but this was not replicated in larger double-blind studies.

SURGICAL TREATMENT

Surgical treatments for PD have been used for more than a century. Lesions placed in the motor cortex improved tremor but were associated with motor deficits, and this approach was abandoned. Subsequently, it was appreciated that lesions placed into the ventral intermediate (VIM) nucleus of the thalamus reduced contralateral tremor without inducing hemiparesis, but these lesions did not meaningfully help other more disabling features of PD. In the 1990s, it was shown that lesions placed in the posteroventral portion of the GPi (motor territory) improved rigidity and bradykinesia as well as tremor. Importantly, pallidotomy was also associated with marked improvement in contralateral dyskinesia. This procedure gained favor with greater understanding of the pathophysiology of PD (see above). However, this procedure is not optimal for patients with bilateral disease, because bilateral lesions are associated with side effects such as dysphagia, dysarthria, and impaired cognition, and has largely been replaced by deep brain stimulation (DBS). Unilateral lesions of the STN are associated with a larger antiparkinsonian benefit and reduced levodopa requirement, but there is a concern about the risk of hemiballismus, and this procedure is not commonly performed.

Most surgical procedures for PD performed today use DBS. Here, an electrode is placed into the target area and connected to a stimulator inserted SC over the chest wall. DBS simulates the effects of a lesion without necessitating making a brain lesion. The precise mechanism whereby DBS works is not fully resolved but may act by disrupting the abnormal signal associated with PD and motor complications. The stimulation variables can be adjusted with respect to electrode configuration, voltage, frequency, and pulse duration in order to maximize benefit and minimize adverse side effects. In cases with intolerable side effects, stimulation can be stopped and the system removed. The procedure does not require making a lesion in the brain and is thus suitable for performing bilateral procedures with relative safety.

DBS for PD primarily targets the STN or the GPi. It provides dramatic results, particularly with respect to reducing “off” time and dyskinesias, but does not improve or prevent the development of features that fail to respond to levodopa such as freezing, falling, and dementia. The procedure is thus primarily indicated for patients who suffer disability resulting from severe tremor, or levodopa-induced motor complications that cannot be satisfactorily controlled with drug manipulation. In such patients, DBS has been shown to improve quality of life in comparison to best medical therapy. Side effects can be seen with respect to the surgical procedure (hemorrhage, infarction, infection), the DBS system (infection, lead break, lead displacement, skin ulceration), or the stimulation itself (ocular and speech abnormalities, muscle twitches, paresthesias, depression, and rarely suicide). Recent studies indicate that benefits following DBS of the STN and GPi are comparable, but that GPi stimulation may be associated with a reduced frequency of depression. Although not all PD patients are candidates, the procedure is profoundly beneficial for many. Studies of DBS in early PD patients show benefits in comparison to medical therapy, but this must be weighed against the cost of the procedure and the risk of side effects. Long-term studies demonstrate continued benefits with respect to the classical motor features of PD, but DBS does not prevent the development of nondopaminergic features, which continue to be a source of disability. Studies continue to evaluate the optimal way to use DBS (low- vs high-frequency stimulation, closed systems, etc.). Comparison of DBS to other therapies aimed at improving motor function without causing dyskinesia, such as

Duodopa and apomorphine infusions, remains to be performed. Studies are examining additional DBS targets that might benefit gait dysfunction, depression, and cognitive impairment in PD patients.

EXPERIMENTAL THERAPIES FOR PD

There has been considerable scientific and public interest in a number of novel interventions that are being investigated as possible treatments for PD. These include cell-based therapies (such as transplantation of fetal nigral dopamine cells or dopamine neurons derived from stem cells), gene therapies, and trophic factors. Transplant strategies are based on the concept of implanting dopaminergic cells into the striatum to replace degenerating SNc dopamine neurons. Fetal nigral mesencephalic cells have been demonstrated to survive implantation, re-innervate the striatum in an organotypic manner, and restore motor function in PD models. However, two double-blind studies failed to show significant benefit of fetal nigral transplantation in comparison to a sham operation with respect to their primary endpoints. Additionally, grafting of fetal nigral cells is associated with a previously unrecognized form of dyskinesia that persists after lowering or even stopping levodopa. This has been postulated to be related to unregulated release of dopamine from serotonin neurons. In addition, there is evidence that after many years, transplanted healthy embryonic dopamine neurons from unrelated donors develop PD pathology and become dysfunctional, suggesting transfer of α -synuclein from affected to unaffected neurons in a prion-like manner (see discussion above). Perhaps most importantly, it is not clear how replacing dopamine cells alone will improve nondopaminergic features such as falling and dementia, which are the major sources of disability for patients with advanced disease. These same concerns apply to dopamine neurons derived from stem cells, which have not yet been properly tested in PD patients and bear the additional concern of tumors and unanticipated side effects. The short-term future for this technology as a treatment for PD, at least in its current state, is therefore not promising, and there is no scientific basis to warrant routine treatment with stem cells as is being marketed in some countries.

Trophic factors are a series of proteins that enhance neuronal growth and restore function to damaged neurons. There are several different trophic factors that have been demonstrated to have beneficial effects on dopamine neurons in laboratory studies. Glial-derived neurotrophic factor (GDNF) and neurturin have attracted particular attention as possible therapies for PD. However, double-blind trials of intraventricular and intraputamenal infusions of GDNF failed to show benefits compared to placebo in PD patients, possibly because of inadequate delivery of the trophic molecule throughout the target region.

Gene delivery offers the potential of providing widespread delivery throughout a target region and long-term expression of a therapeutic protein with a single procedure. Gene therapy involves placing the DNA of a therapeutic protein into a viral vector that can then be delivered to specific target regions. The DNA of the therapeutic protein is then incorporated into the genome of the host cells and released on a continual basis. The AAV2 virus has been most often used as the viral vector because it does not promote an inflammatory response, is not incorporated into the host genome, and is associated with long-lasting transgene expression. Clinical trials of AAV2 delivery of the trophic factor neurturin showed promising results in open label trials but failed in double-blind trials, possibly because axonal damage in PD prevented retrograde transport of the protein to dopamine neurons in the SNc where it is required to induce upregulation of repair genes required for the trophic response. However, a subsequent double-blind trial of AAV2-neurturin delivered into both the putamen and SNc also failed.

Gene delivery is also being explored as a means of delivering aromatic amino acid decarboxylase with or without tyrosine hydroxylase to the striatum to facilitate dopamine production and glutamic acid decarboxylase to the STN to inhibit overactive neuronal firing. None of these procedures has been established to be effective in PD patients. Furthermore, although gene delivery technology has great

potential, this approach also carries the risk of unanticipated side effects, and current approaches directed at the nigrostriatal system do not address the nondopaminergic features of the illness.

MANAGEMENT OF THE NONMOTOR AND NONDOPAMINERGIC FEATURES OF PD

Although PD management has primarily focused on the dopaminergic features of the disease, management of the nondopaminergic features should not be ignored. Some nonmotor features, although not thought to reflect dopaminergic pathology, nonetheless benefit from dopaminergic drugs. For example, problems such as anxiety, panic attacks, depression, sweating, sensory problems, freezing, and constipation all tend to be worse during "off" periods and may improve with better dopaminergic control. Approximately 50% of PD patients suffer depression during the course of the disease, and depression is frequently underdiagnosed and undertreated. Antidepressants should not be withheld, particularly for patients with major depression. Serotonin syndromes have been a theoretical concern with the combined use of SSRIs and MAO-B inhibitors but are rarely encountered. Anxiety can be treated with short-acting benzodiazepines.

Psychosis can be a problem for some PD patients. In contrast to AD, hallucinations are typically visual, formed, and nonthreatening. Importantly, they can limit the use of dopaminergic agents to obtain satisfactory motor control. Psychosis in PD often responds to low doses of atypical neuroleptics and permits higher doses of levodopa to be tolerated. Clozapine is the most effective drug, but it can be associated with agranulocytosis, and regular monitoring is required. For this reason, many physicians start with quetiapine even though it has not been established to be effective in placebo-controlled trials. Hallucinations in PD patients are often a harbinger of a developing dementia.

Dementia in PD (PDD) is common, ultimately affecting as many as 80% of patients. Its frequency increases with aging and, in contrast to AD, primarily affects executive functions and attention, with relative sparing of language, memory, and calculations. When dementia precedes, or develops within 1 year after, the onset of motor dysfunction, it is by convention referred to as dementia with Lewy bodies (DLB; [Chap. 448](#)). These patients are particularly prone to have hallucinations and diurnal fluctuations. Pathologically, DLB is characterized by Lewy bodies distributed throughout the cerebral cortex (especially the hippocampus and amygdala) and is often also associated with AD pathology. It is likely that DLB and PDD represent a PD spectrum rather than separate disease entities. Mild cognitive impairment (MCI) frequently precedes the onset of dementia and is a more reliable index of impending dementia in PD than in the general population. Dopaminergic drugs can worsen cognitive function in demented patients and should be stopped or reduced to try and provide a compromise between antiparkinsonian benefit and preserved cognitive function. Drugs are usually discontinued in the following sequence: anticholinergics, amantadine, dopamine agonists, COMT inhibitors, and MAO-B inhibitors. Eventually, patients with cognitive impairment should be managed with the lowest dose of standard levodopa that provides meaningful antiparkinsonian effects and does not worsen mental function. Anticholinesterase agents such as rivastigmine and donepezil reduce the rate of deterioration of measures of cognitive function and can improve attention, but do not typically improve cognitive function in any meaningful way.

Autonomic disturbances are common and frequently require attention. Orthostatic hypotension can be problematic and contribute to falling. Initial treatment should include adding salt to the diet and elevating the head of the bed to prevent overnight sodium natriuresis. Low doses of fludrocortisone (Florinef) or midodrine provide control for most cases. Vasopressin, erythropoietin, and the norepinephrine precursor 3-O-methyldops can be used in more severe or refractory cases. If orthostatic hypotension is prominent in early disease, MSA should be considered. Sexual dysfunction can be helped with sildenafil or tadalafil. Urinary problems, especially in males, should be treated in consultation with a urologist to exclude

prostate problems. Anticholinergic agents, such as oxybutynin (Ditropan), may be helpful. Constipation can be a very important problem for PD patients. Mild laxatives or enemas can be useful, but physicians should first ensure that patients are drinking adequate amounts of fluid and consuming a diet rich in bulk with green leafy vegetables and bran. Agents that promote gastrointestinal (GI) motility can also be helpful.

Sleep disturbances are common in PD patients, with many experiencing fragmented sleep with excess daytime sleepiness. Restless leg syndrome, sleep apnea, and other sleep disorders should be treated as appropriate. REM behavior disorder (RBD) is a syndrome comprised of violent movements and vocalizations during REM sleep, possibly representing acting out of dreams due to a failure of the normal inhibition of motor movements that typically accompanies REM sleep. Many PD patients have a history of antecedent RBD preceding the onset of the classic motor features of PD, and most cases of RBD go on to develop an α -synucleinopathy (PD or MSA). Low doses of clonazepam (0.5–1 mg at bedtime) are usually effective in controlling this problem. Consultation with a sleep specialist and polysomnography may be necessary to identify and optimally treat sleep problems.

NONPHARMACOLOGIC THERAPY

Gait dysfunction with falling is an important cause of disability in PD. Dopaminergic therapies can help patients whose gait is worse in “off” time, but there are currently no specific therapies available. Canes and walkers may become necessary to increase stability and reduce the risk of falling.

Freezing, where patients suddenly become stuck in place for seconds to minutes as if their feet were glued to the ground, is a major cause of falling. Freezing may occur during “on” or “off” periods. Freezing during “off” periods may respond to dopaminergic therapies, but there are no specific treatments for “on” period freezing. Some patients will respond to sensory cues such as marching in place, singing a song, or stepping over an imaginary line.

Exercise, with a full range of active and passive movements, has been shown to maintain and even improve function for PD patients, and active and passive exercises with full range of motion reduce the risk of arthritis and frozen joints. Some laboratory studies suggest the possibility that exercise might also have neuroprotective effects, but this has not been confirmed in PD. Exercise is generally recommended for all PD patients. It is less clear that physical therapy or specific exercises such as tai chi are required. It is important for patients to maintain social and intellectual activities to the extent possible. Education, assistance with financial planning, social services, and attention to home safety are important elements of the overall care plan. Information is available through numerous PD foundations and on the web, but should be reviewed with physicians to ensure accuracy. The needs of the caregiver should not be neglected. Caring for a person with PD involves a substantial work effort and there is an increased incidence of depression among caregivers. Support groups for patients and caregivers may be useful.

CURRENT MANAGEMENT OF PD

The management of PD should be tailored to the needs of the individual patient, and there is no single treatment approach that is universally accepted and applicable to all individuals. Clearly, if an agent could be demonstrated to have disease-modifying effects, it should be initiated at the time of diagnosis. Indeed, constipation, RBD, and anosmia may represent premotor features of PD and could permit the initiation of a disease-modifying therapy prior to the onset of the classical motor features of the disease. However, no therapy has yet been proved to be disease modifying. For now, physicians must use their judgment in deciding whether or not to introduce rasagiline (see above) or other drugs for their possible disease-modifying effects.

The next important issue to address is when to initiate symptomatic therapy. Several studies now suggest that it may be best to start therapy at the time of diagnosis (or soon after) in order

to preserve beneficial compensatory mechanisms and possibly provide functional benefits even in the early stage of the disease. Levodopa remains the most effective symptomatic therapy for PD, and some recommend starting it immediately using low doses (≤ 400 mg/d), but others prefer to delay levodopa treatment, particularly in younger patients, in order to reduce the risk of inducing motor complications. An alternate approach is to begin with an MAO-B inhibitor and/or a dopamine agonist, and reserve levodopa for later stages when these drugs can no longer provide satisfactory control. In making this decision, the age, degree of disability, and side effect profile of the drug must all be considered. In patients with more severe disability, the elderly, those with cognitive impairment, or those in whom the diagnosis is uncertain, most physicians would initiate therapy with levodopa. Regardless of initial choice, it is important not to deny patients levodopa when they cannot be adequately controlled with alternative medications.

If motor complications develop, patients can initially be treated by manipulating the frequency and dose of levodopa or by combining lower doses of levodopa with a dopamine agonist, a COMT inhibitor, or an MAO-B inhibitor. Amantadine is the only drug that has been demonstrated to treat dyskinesia without worsening parkinsonism, but benefits may be short-lasting, and there are important side effects related to cognitive function. In advanced cases, it may be necessary to consider a surgical therapy such as DBS if the patient is a suitable candidate, but as described above, these procedures have their own set of complications. Continuous intraintestinal infusion of levodopa/carbidopa intestinal gel (Duodopa) appears to offer similar benefits to DBS, but also requires a surgical intervention with potentially serious complications. Continuous infusion of apomorphine is another treatment option and does not require surgery but is associated with potentially troublesome skin nodules. Comparative studies of these approaches in more advanced patients are awaited. There are ongoing efforts aimed at developing a long-acting oral or transdermal formulation of levodopa that mirrors the pharmacokinetic properties of a levodopa infusion. Such a formulation might provide all of the benefits of levodopa without motor complications and avoid the need for polypharmacy and surgical intervention.

A decision tree that considers the various treatment options and decision points for the management of PD is provided in [Fig. 449-7](#).

HYPERKINETIC MOVEMENT DISORDERS

Hyperkinetic movement disorders are characterized by involuntary movements unaccompanied by weakness and occurring in isolation or in combination ([Table 449-6](#)). The major hyperkinetic movement disorders and the diseases with which they are associated are considered in this section.

TREMOR

CLINICAL FEATURES

Tremor consists of alternating contractions of agonist and antagonist muscles in an oscillating, rhythmic manner. It can be most prominent at rest (rest tremor), on assuming a posture (postural tremor), or on actively reaching for a target (kinetic tremor). Tremor is also assessed based on distribution, frequency, and related neurologic dysfunction.

PD is characterized by a resting tremor, essential tremor (ET) by a postural tremor (trying to sustain a posture), and cerebellar disease by an intention or kinetic tremor (on reaching to touch a target). Normal individuals can have a physiologic tremor that typically manifests as a mild, high-frequency (10–12 Hz), postural or action tremor that is usually of no clinical consequence and often is only appreciated with an accelerometer. An enhanced physiologic tremor (EPT) can be seen in up to 10% of the population, often in association with anxiety, fatigue, a metabolic disturbance (e.g., hyperthyroidism, electrolyte abnormalities), drugs (e.g., valproate, lithium), or toxins (e.g., alcohol). Treatment is initially directed to the control of any underlying disorder and, if necessary, can often be improved with a beta blocker.

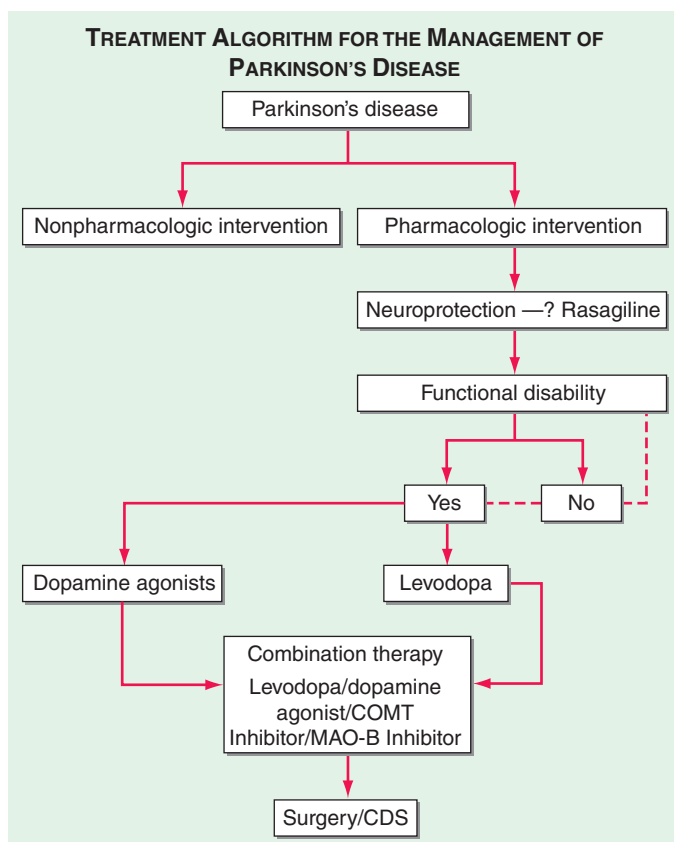


FIGURE 449-7 Treatment options for the management of Parkinson's disease (PD). Decision points for the include: (1) Introduction of a neuroprotective therapy: No drug has been established to have or is currently approved for neuroprotection or disease modification, but there are several agents that have this potential based on laboratory and preliminary clinical studies (e.g., rasagiline 1 mg/d, coenzyme Q10 1200 mg/d, the dopamine agonists ropinirole, and pramipexole). (2) When to initiate symptomatic therapy: There is a trend toward initiating therapy at the time of diagnosis or early in the course of the disease because patients may have some disability even at an early stage, and there is the possibility that early treatment may preserve beneficial compensatory mechanisms; however, some experts recommend waiting until there is functional disability before initiating therapy. (3) What therapy to initiate: Many experts favor starting with a monoamine oxidase type B (MAO-B) inhibitor in mildly affected patients because of the good safety profile of the drug and the potential for a disease-modifying effect; dopamine agonists for younger patients with functionally significant disability to reduce the risk of motor complications; and levodopa for patients with more advanced disease, the elderly, or those with cognitive impairment. Recent studies suggest the early employment of polypharmacy using low doses of multiple drugs to avoid side effects associated with high doses of any one agent. (4) Management of motor complications: Motor complications are typically approached with combination therapy to try and reduce dyskinesia and enhance the “on” time. When medical therapies cannot provide satisfactory control, surgical therapies such as DBS or continuous infusion of levodopa/carbidopa intestinal gel can be considered. (5) Nonpharmacologic approaches: Interventions such as exercise, education, and support should be considered throughout the course of the disease. CDS, continuous dopaminergic stimulation; COMT, catechol-O-methyltransferase. (Adapted from CW Olanow et al: *Neurology* 72:51, 2009.)

ESSENTIAL TREMOR

ET is the commonest movement disorder, affecting approximately 5–10 million persons in the United States. It can present in childhood but dramatically increases in prevalence over the age of 70 years. ET is characterized by a high-frequency tremor (6–10 Hz) that

TABLE 449-6 HYPERKINETIC MOVEMENT DISORDERS

Tremor	Rhythmic oscillation of a body part due to intermittent muscle contractions
Dystonia	Involuntary, patterned, sustained or repeated muscle contractions often associated with twisting movements and abnormal posture
Athetosis	Slow, distal, writhing, involuntary movements with a propensity to affect the arms and hands (this represents a form of dystonia with increased mobility)
Chorea	Rapid, semi-purposeful, graceful, dance-like nonpatterned involuntary movements involving distal or proximal muscle groups. When the movements are of large amplitude and predominant proximal distribution, the term <i>ballism</i> is used.
Myoclonus	Sudden, brief (<100 ms), jerk-like, arrhythmic muscle twitches
Tic	Brief, repeated, stereotyped muscle contractions that can often be suppressed for a short time. These can be simple and involve a single muscle group or complex and affect a range of motor activities.

predominantly affects the upper extremities. The tremor is most often manifest as a postural or action (kinetic) tremor and, in severe cases, can interfere with functions such as eating and drinking. It is typically bilateral and symmetric but may begin on one side and remain asymmetric. Patients with severe ET can have an intention tremor with overshoot and slowness of movement. Tremor involves the head in ~30% of cases, voice in ~20%, tongue in ~20%, face/jaw in ~10%, and lower limbs in ~10%. The tremor is characteristically improved by alcohol and worsened by stress. Subtle impairment of coordination or tandem walking may be present, and disturbances of hearing, cognition, personality, mood, and olfaction have also been described, but usually the neurologic examination is normal aside from tremor. The major differential is a dystonic tremor (see below) or PD. PD can usually be differentiated from ET based on the presence of bradykinesia, rigidity, micrographia, and other parkinsonian features. However, the examiner should be aware that PD patients may have a postural tremor and ET patients may develop a rest tremor. These typically begin after a latency of a few seconds (emergent tremor). The examiner must take care to differentiate the effect of tremor on measurement of tone in ET from the cogwheel rigidity found in PD.

ETIOLOGY AND PATHOPHYSIOLOGY

The etiology and pathophysiology of ET are not known. Approximately 50% of cases have a positive family history with an autosomal dominant pattern of inheritance. Linkage studies have detected loci at chromosomes 3q13 (ETM-1), 2p22-25 (ETM-2), and 6p23 (ETM-3), but no causative genes have been identified to date. GWAS demonstrated an association with the *LINGO1* gene, which is involved in oligodendrocyte differentiation and myelination, particularly in patients with young-onset ET. Recently, a nonsense mutation in the fused in sarcoma (*FUS*) gene was implicated as a cause of ET in a multigenerational family from Canada; this finding is of particular interest because different mutations in *FUS* are a known cause of familial amyotrophic lateral sclerosis (Chap. 452). It is likely that there are many other undiscovered genes for ET. The cerebellum and inferior olives have been implicated as possible sites of a “tremor pacemaker” based on the presence of cerebellar signs and increased metabolic activity and blood flow in these regions in some patients. Some pathologic studies have described cerebellar pathology with a loss of Purkinje cells and axonal torpedoes, but these findings are controversial and the precise pathologic correlate of ET remains to be defined.

TREATMENT

Many cases are mild and require no treatment other than reassurance. Occasionally, tremor can be severe and interfere with eating, writing, and activities of daily living. This is more likely to occur as the patient ages and is often associated with a reduction in tremor frequency. Beta blockers and primidone are the standard drug therapies for ET and help in about 50% of cases. Propranolol (20–120 mg daily, given in

divided doses) is usually effective at relatively low doses, but higher doses may be effective in some patients. The drug is contraindicated in patients with bradycardia or asthma. Hand tremor tends to be most improved, while head tremor is often refractory. Primidone can be helpful but should be started at low doses (12.5 mg) and gradually increased (125–250 mg tid) to avoid sedation. Benefits have also been reported with gabapentin and topiramate. Botulinum toxin injections may be helpful for limb or voice tremor, but treatment can be associated with secondary muscle weakness. Surgical therapies targeting the VIM nucleus of the thalamus can be very effective for severe and drug-resistant cases.

DYSTONIA

CLINICAL FEATURES

Dystonia is a disorder characterized by sustained (>100 ms) or repetitive involuntary muscle contractions frequently associated with twisting and abnormal postures. Dystonia can range from minor contractions in an individual muscle group to severe and disabling involvement of multiple muscle groups. The frequency is estimated to be 300,000 cases in the United States but is likely to be much higher because many cases are not recognized. Dystonia is often brought out by voluntary movements (action dystonia) and can extend to involve muscle groups and body regions not required for a given action (overflow). It can be aggravated by stress and fatigue and attenuated by relaxation and sensory tricks such as touching the affected body part (*geste antagoniste*). Dystonia can be classified according to age of onset (childhood vs adult), distribution (focal, multifocal, segmental, or generalized), or etiology (primary or secondary).

PRIMARY DYSTONIAS

At least 16 gene mutations are associated with dystonia and classified as *DYT1–DYT16*. Idiopathic torsion dystonia (*DYT1*) or Oppenheim's dystonia is predominantly a childhood-onset form of dystonia with an autosomal dominant pattern of inheritance that primarily affects Ashkenazi Jewish families. The majority of patients have an age of onset younger than 26 years (mean 14 years). In young-onset patients, dystonia typically begins in the foot or the arm and in 60–70% progresses to involve other limbs as well as the head and neck. In severe cases, patients can suffer disabling postural deformities that compromise mobility. Severity can vary within family members, with some affected relatives having severe disability and others a mild dystonia that may not even be appreciated. Most childhood-onset cases are linked to a mutation in the *DYT1* gene located on chromosome 9q34, resulting in a trinucleotide GAG deletion with loss of one of a pair of glutamic acid residues in the protein torsin A. *DYT1* mutations are found in 90% of Ashkenazi Jewish patients with *DYT1* dystonia and probably relate to a founder effect that occurred about 350 years ago. There is variable penetrance, with only about 30% of gene carriers expressing a clinical phenotype. Why some gene carriers express dystonia and others do not is not known. The function of torsin A is unknown, but it is a member of the AAA⁺ (ATPase) family that resembles heat-shock proteins and may be related to protein processing and transport. The precise pathology responsible for *DYT1* dystonia is not known.

Dopa-responsive dystonia (DRD) or the Segawa variant (*DYT5*) is a dominantly inherited form of childhood-onset dystonia caused by a mutation in the gene that encodes GTP cyclohydrolase-I, the rate-limiting enzyme for the synthesis of tetrahydrobiopterin. This mutation leads to a defect in the biochemical synthesis of tyrosine hydroxylase, the rate-limiting enzyme in the formation of dopamine. DRD typically presents in early childhood (1–12 years) and is characterized by foot dystonia that interferes with walking. Patients often experience diurnal fluctuations, with worsening of gait as the day progresses and improvement with sleep. DRD is typified by an excellent and sustained response to small doses of levodopa. Some patients may present with parkinsonian features, but can be differentiated from juvenile PD by normal striatal dopamine imaging and the absence of levodopa-induced dyskinesias. DRD may occasionally be confused with cerebral palsy because patients appear to have spasticity, increased reflexes, and Babinski responses (which likely reflect a dystonic contraction rather

than an upper motor neuron lesion). Any patient suspected of having a childhood-onset dystonia should receive a trial of levodopa to exclude this treatable condition.

Mutations in the *THAP1* gene (*DYT6*) on chromosome 8p21q22 have been identified in Amish families and are the cause of as many as 25% of cases of non-*DYT1* young-onset primary torsion dystonia. These patients are more likely to have dystonia beginning in the brachial and cervical muscles, which later can become generalized and associated with speech impairment. Myoclonic dystonia (*DYT11*) results from a mutation in the epsilon-sarcoglycan gene on chromosome 7q21. It typically manifests as a combination of dystonia and myoclonic jerks, frequently accompanied by psychiatric disturbances.

FOCAL DYSTONIAS

These are the most common forms of dystonia. They typically present in the fourth to sixth decades and affect women more than men. The major types are as follows: (1) *blepharospasm*—dystonic contractions of the eyelids with increased blinking that can interfere with reading, watching television, and driving. This can sometimes be so severe as to cause functional blindness. (2) *Oromandibular dystonia* (OMD)—contractions of muscles of the lower face, lips, tongue, and jaw (opening or closing). Meige's syndrome is a combination of OMD and blepharospasm that predominantly affects women older than age 60 years. (3) *Spasmodic dysphonia*—dystonic contractions of the vocal cords during phonation, causing impaired speech. Most cases affect the adductor muscles and cause speech to have a choking or strained quality. Less commonly, the abductors are affected, leading to speech with a breathy or whispering quality. (4) *Cervical dystonia*—dystonic contractions of neck muscles causing the head to deviate to one side (*torticollis*), in a forward direction (*anterocollis*), or in a backward direction (*retrocollis*). Muscle contractions can be painful and associated with a secondary cervical radiculopathy. (5) *Limb dystonias*—these can be present in either arms or legs and are often brought out by task-specific activities such as handwriting (writer's cramp), playing a musical instrument (musician's cramp), or putting (the yips). Focal dystonias can extend to involve other body regions (about 30% of cases) and are frequently misdiagnosed as psychiatric or orthopedic in origin. Their cause is not known, but genetic factors, autoimmunity, and trauma have been suggested. Focal dystonias are often associated with a high-frequency tremor that resembles ET. Dystonic tremor can usually be distinguished from ET because it tends to occur in conjunction with the dystonic contraction and disappears when the dystonia is relieved.

SECONDARY DYSTONIAS

These develop as a consequence of drugs or other neurologic disorders. Drug-induced dystonia is most commonly seen with neuroleptic drugs or after chronic levodopa treatment in PD patients and may be acute or chronic (see below). Secondary dystonia can also be observed following discrete lesions in the striatum and occasionally in the pallidum, thalamus, cortex, and brainstem due to infarction, anoxia, metabolic disorders, trauma, tumor, infection, or toxins such as manganese or carbon monoxide. In these cases, dystonia often assumes a segmental distribution, but it can be generalized when lesions are bilateral or widespread. More rarely, dystonia can develop following peripheral nerve injury and be associated with features of complex regional pain syndrome (**Chap. 454**). A psychogenic origin is responsible for some cases of dystonia presenting with fixed, immobile dystonic postures (see below).

DYSTONIA PLUS SYNDROMES

Dystonia may occur as a part of another neurodegenerative conditions such as Huntington's disease, PD, Wilson's disease, corticobasilar ganglionic degeneration, PSP, the Lubag form of dystonia-parkinsonism (*DYT3*), and mitochondrial encephalopathies. In contrast to the primary dystonias, dystonia is usually not the dominant neurologic feature in these conditions.

PATHOPHYSIOLOGY OF DYSTONIA

The pathophysiologic basis of dystonia is not completely known. The phenomenon is characterized by co-contracting synchronous bursts of agonist and antagonist muscle groups with recruitment of

muscle groups that are not required for a given movement (overflow). Dystonia is characterized by derangement of the basic physiological principle of action-selection, leading to abnormal recruitment of inappropriate muscles for a given action with inadequate inhibition of this undesired motor activity. Physiologically, loss of inhibition is observed at multiple levels of the motor system (e.g., cortex, brainstem, spinal cord) accompanied by increased cortical excitability and reorganization. Attention has focused on the basal ganglia as the site of origin of at least some types of dystonia because there are alterations in blood flow and metabolism in these structures. Further, lesions of the GPi can induce dystonia, and surgical ablation or DBS of the globus pallidus can ameliorate dystonia. The dopamine system has also been implicated, because dopaminergic therapies can both induce and treat some forms of dystonia. Interestingly, no specific pathology has been consistently identified in primary dystonia.

TREATMENT DYSTONIA

Treatment of dystonia is for the most part symptomatic except in rare cases where correction of a primary underlying condition is possible. Wilson's disease should be ruled out in young patients with dystonia. Levodopa should be tried in all cases of childhood-onset dystonia to rule out DRD. High-dose anticholinergics (e.g., trihexyphenidyl 20–120 mg/d) may be beneficial in children, but adults can rarely tolerate high doses because of side effects related to cognitive impairment with hallucinations. Oral baclofen (20–120 mg) may also be helpful, but benefits, if present, are usually modest, and side effects of sedation, weakness, and memory loss can be problematic. Intrathecal infusion of baclofen is more likely to be useful, particularly for leg and trunk dystonia, but benefits are frequently not sustained, and complications can be serious and include infection, seizures, and coma. Tetrabenazine (the usual starting dose is 12.5 mg/d and the average treating dose is 25–75 mg/d) is another consideration, but use may be limited by sedation and the development of parkinsonism. Neuroleptics can improve as well as induce dystonia, but they are typically not recommended because of their potential to induce parkinsonism and other movement disorders, including tardive dystonia. Clonazepam and diazepam are rarely effective.

Botulinum toxin has become the preferred treatment for patients with focal dystonia, particularly where involvement is limited to small muscle groups such as in blepharospasm, torticollis, and spasmodic dysphonia. Botulinum toxin acts by blocking the release of acetylcholine at the neuromuscular junction, leading to reduced dystonic muscle contractions, but excessive weakness may ensue and can be troublesome particularly if it involves neck and swallowing muscles. Two serotypes of botulinum toxin are available (A and B). Both are effective, and it is not clear that there are advantages of one over the other. No systemic side effects are encountered with the doses typically used, but benefits are transient, and repeat injections are required at 2- to 5-month intervals. Some patients fail to respond after having experienced an initial benefit. This has been attributed to antibody formation, but improper muscle selection, injection technique, and inadequate dose should be excluded.

Surgical therapy is an alternative for patients with severe dystonia who are not responsive to other treatments. Peripheral procedures such as rhizotomy and myotomy were used in the past to treat cervical dystonia, but are now rarely used. DBS of the pallidum can provide dramatic benefits for patients with primary DYT1 dystonia. This represents a major therapeutic advance because previously there was no consistently effective therapy, especially for these patients who had severe disability. Benefits tend to be obtained with a lower frequency of stimulation and often occur after a relatively long latency (weeks) in comparison to PD. Better results are typically obtained in younger patients with shorter disease duration. Recent studies suggest that DBS may also be valuable for patients with focal and secondary dystonias, although results are less consistent. Supportive treatments such as physical therapy and education are important and should be a part of the treatment regimen.

Physicians should be aware of dystonic storm, a rare but potentially fatal condition that can occur in response to a stress situation such as surgery in patients with preexisting dystonia. It consists of the acute onset of generalized and persistent dystonic contractions that can involve the vocal cords or laryngeal muscles, leading to airway obstruction. Patients may experience rhabdomyolysis with renal failure and should be managed in an intensive care unit with airway protection if required. Treatment can be instituted with one or a combination of anticholinergics, diphenhydramine, baclofen, benzodiazepines, and dopaminergic agents. Spasms may be difficult to control, and anesthesia with muscle paralysis may be required. Most, if not all, cases of dystonic storm are due to a secondary cause.

CHOREAS

HUNTINGTON'S DISEASE (HD)

HD is a progressive, fatal, highly penetrant autosomal dominant disorder characterized by motor, behavioral, oculomotor, and cognitive dysfunction. The disease is named for George Huntington, a family physician who described cases on Long Island, New York, in the nineteenth century. Onset is typically between the ages of 25 and 45 years (range, 3–70 years) with a prevalence of 2–8 cases per 100,000 and an average age at death of 60 years. It is prevalent in Europe, North and South America, and Australia but is rare in African blacks and Asians. HD is characterized by rapid, nonpatterned, semipurposeful, involuntary choreiform movements, and for this reason was formerly referred to as Huntington's chorea. In the early stages, the chorea tends to be focal or segmental, but progresses over time to involve multiple body regions. Dysarthria, gait disturbance, oculomotor abnormalities, behavioral disturbance, and cognitive impairment with dementia are also common features. With advancing disease, there tends to be a reduction in chorea and the emergence of dystonia, rigidity, bradykinesia, and myoclonus. Functional decline is often predicted by progressive weight loss despite adequate calorie intake. In younger patients (~10% of cases), HD can present as an akinetic-rigid or parkinsonian syndrome (Westphal variant). HD patients eventually develop behavioral and cognitive disturbances, and the majority progress to dementia. Depression with suicidal tendencies, aggressive behavior, and psychosis can be prominent features. HD patients may also develop non-insulin-dependent diabetes mellitus and neuroendocrine abnormalities (e.g., hypothalamic dysfunction). A clinical diagnosis of HD can be strongly suspected in cases of chorea with a positive family history, but genetic testing provides the ultimate confirmation of the diagnosis. The disease predominantly affects the striatum. Progressive atrophy of the heads of the caudate nuclei, which form the lateral margins of the lateral ventricles, can be visualized by MRI (Fig. 449-8), but the putamen can be equally or even more severely affected. More diffuse cortical atrophy is seen in the middle and late stages of the disease. Supportive studies include reduced metabolic activity in the caudate nucleus and putamen. Genetic testing can be used to confirm the diagnosis and to detect at-risk individuals in the family, but must be performed with caution and in conjunction with trained counselors, because positive results can worsen depression and generate suicidal reactions. The neuropathology of HD consists of prominent neuronal loss and gliosis in the caudate nucleus and putamen; similar changes are also widespread in the cerebral cortex. Intra-neuronal inclusions containing aggregates of ubiquitin and the mutant protein huntingtin are found in the nuclei of affected neurons.

In anticipation of developing neuroprotective therapies, there has been an intensive effort to define the premanifest stage of HD. Subtle motor impairment, cognitive alterations, and imaging changes can be detected in at-risk individuals who later go on to develop the manifest form of the disease. Defining the rate of progression of these features is paramount for future studies of putative disease-modifying therapies.

ETIOLOGY

HD is caused by an increase in the number of polyglutamine (CAG) repeats (>40) in the coding sequence of the huntingtin gene located on

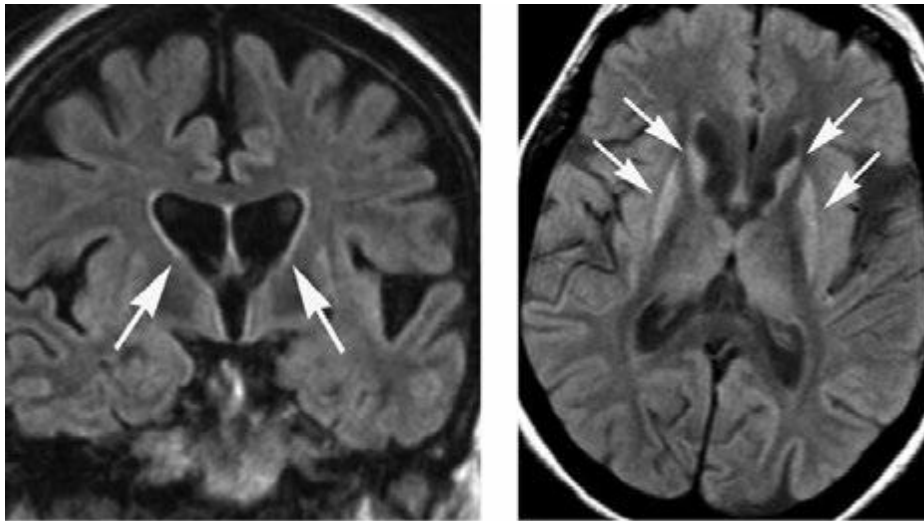


FIGURE 449-8 Huntington's disease. **A.** Coronal fluid attenuated inversion recovery (FLAIR) magnetic resonance imaging shows enlargement of the lateral ventricles reflecting typical atrophy (arrows). **B.** Axial FLAIR image demonstrates abnormal high signal in the caudate and putamen (arrows).

the short arm of chromosome 4. The larger the number of repeats, the earlier the disease is manifest. Intermediate forms of the disease with 36–39 repeats are described in some patients, typically with less severe clinical involvement. Acceleration of the process tends to occur, particularly in males, with subsequent generations having larger numbers of repeats and earlier age of disease onset, a phenomenon referred to as anticipation. The gene encodes the highly conserved cytoplasmic protein huntingtin, which is widely distributed in neurons throughout the central nervous system (CNS) but whose function is not known. Models of HD with striatal pathology can be induced by excitotoxic agents such as kainic acid and 3-nitropropionic acid, which promote calcium entry into the cell and cytotoxicity. Mitochondrial dysfunction has been demonstrated in the striatum and skeletal muscle of symptomatic and presymptomatic individuals. Fragments of the mutant huntingtin protein can be toxic, possibly by translocating into the nucleus and interfering with transcriptional regulation of proteins. Neuronal inclusions found in affected regions in HD may represent a protective mechanism aimed at segregating and facilitating the clearance of these toxic proteins.

TREATMENT HUNTINGTON'S DISEASE

Although the gene for HD was identified more than two decades ago, there is still no disease-modifying therapy for this disorder. Current treatment involves a multidisciplinary approach, with medical, neuropsychiatric, social, and genetic counseling for patients and their families. Dopamine-blocking agents may control the choreatic movements. Tetrabenazine (a presynaptic dopamine depleting agent) has been approved for the treatment of chorea in the United States, but can cause secondary parkinsonism. Neuroleptics are generally not recommended because of their potential to induce other more troubling movement disorders and because HD chorea tends to be self-limited and is usually not disabling. Depression and anxiety can be greater problems, and patients should be treated with appropriate antidepressant and anti-anxiety drugs and monitored for mania and suicidal ideations. Psychosis can be treated with atypical anti-psychotics such as clozapine (50–600 mg/d), quetiapine (50–600 mg/d), and risperidone (2–8 mg/d). There is no adequate treatment for the cognitive or motor decline. A neuroprotective therapy that slows or stops disease progression is the major unmet medical need in HD. Drugs that enhance mitochondrial function and increase the clearance of defective mitochondria are being tested as possible disease-modifying therapies. Antigliutamate agents, dopamine stabilizers, caspase inhibitors, neurotrophic factors, and

transplantation of fetal striatal cells are areas of active research, but none has as yet been demonstrated to have a beneficial effect in HD. The potential to use transcriptional blockade of the mutant huntingtin gene with small interfering RNAs (siRNAs) is an exciting area currently being explored.

HUNTINGTON'S DISEASE-LIKE DISORDERS

A group of rare inherited conditions that can mimic HD, designated HD-like (HDL) disorders, have also been identified. HDL-1, -2, and -4 are autosomal dominant conditions that typically present in adulthood. HDL-1 is due to expansion of an octapeptide repeat in *PRNP*, the gene encoding the prion protein (**Chap. 453e**). Thus HDL-1 is properly considered a prion disease. Patients exhibit onset of personality change in the third or fourth decade, followed by chorea, rigidity, myoclonus, ataxia, and epilepsy. HDL-2 manifests in the third or fourth decade with a variety of movement disorders, including chorea, dystonia, or parkinsonism and dementia. Most patients are of African descent. Acanthocytosis can sometimes be seen in these patients, and this condition must be distinguished from neuroacanthocytosis. HDL-2 is caused by an abnormally expanded CTG/CAG trinucleotide repeat expansion in the *junctophilin-3* (*JPH3*) gene. The pathology of HDL-2 consists of intranuclear inclusions immunoreactive for ubiquitin and expanded polyglutamine repeats. HDL-4, the most common condition in this group, is caused by expansion of trinucleotide repeats in *TBP*, the gene that encodes the TATA box binding protein involved in regulating transcription; this condition is identical to spinocerebellar ataxia (SCA) 17 (**Chap. 451e**), and most patients present primarily with ataxia rather than chorea. Mutations of the *C9orf* gene associated with amyotrophic lateral sclerosis have also been reported in some individuals with an HDL phenotype.

OTHER CHOREAS

Chorea can be seen in a number of additional disorders. Sydenham's chorea (originally called St. Vitus's dance) is more common in females and is typically seen in childhood (5–15 years). It often develops in association with prior exposure to group A streptococcal infection and is thought to be autoimmune in nature. It is characterized by the acute onset of choreiform movements and behavioral disturbances. With the reduction in the incidence of rheumatic fever, the incidence of Sydenham's chorea has fallen, but it can still be seen in developing countries. The chorea generally responds to dopamine-blocking agents, valproic acid, and carbamazepine, but is self-limited, and treatment is generally restricted to those with severe chorea. Chorea may recur in later life, particularly in association with pregnancy (chorea gravidarum) or treatment with sex hormones. Several reports have documented cases of chorea associated with NMDA receptor antibody-positive encephalitis following herpes simplex virus encephalitis.

Chorea-acanthocytosis (neuroacanthocytosis) is a progressive and typically fatal autosomal recessive disorder that is characterized by chorea coupled with red cell abnormalities on peripheral blood smear (acanthocytes). The chorea can be severe and associated with self-mutilating behavior, dystonia, tics, seizures, and a polyneuropathy. Mutations in the *VPS13A* gene encoding chorein have been described. A phenotypically similar X-linked form of the disorder has been described in older individuals who have reactivity with Kell blood group antigens (McLeod syndrome). A benign hereditary chorea of childhood (BHC1) due to mutations in the gene for thyroid transcription factor 1 and a late-onset benign senile chorea (BHC2) have also been described. It is important to ensure that patients with these types of choreas do not have HD.

Chorea may also occur in association with vascular diseases, hypo- and hyperglycemia, and a variety of infections and degenerative disorders. Systemic lupus erythematosus is the most common systemic disorder that causes chorea, which can last for days to years. Chorea can also be seen with hyperthyroidism, autoimmune disorders including Sjögren's syndrome, infectious disorders including HIV disease, metabolic alterations, and polycythemia rubra vera; following open-heart surgery in the pediatric population; and in association with many medications (especially anticonvulsants, cocaine, CNS stimulants, estrogens, and lithium). Chorea is commonly seen in association with chronic levodopa treatment (discussed in the section on PD above). Chorea can also be seen in paraneoplastic syndromes associated with anti-CRMP-5 or anti-Hu antibodies ([Chap. 122](#)).

HEMIBALLISMUS

Hemiballismus is a violent form of chorea comprised of wild, flinging, large-amplitude movements on one side of the body. Proximal limb muscles tend to be predominantly affected. These movements may affect just one limb (monoballismus) or, more exceptionally, both upper or lower limbs (paraballismus). The movements may be so severe as to cause exhaustion, dehydration, local injury, and, in extreme cases, death. Fortunately, dopamine-blocking drugs can be very helpful, and importantly, hemiballismus is usually self-limiting and tends to resolve spontaneously after weeks or months. The most common cause is a partial lesion (infarct or hemorrhage) in the STN, but cases can also be seen with lesions in the putamen, thalamus, and parietal cortex. In extreme cases, pallidotomy can be very effective. Interestingly, surgically induced lesions and DBS of the STN in PD patients are usually not associated with hemiballismus.

TICS

A *tic* is a brief, rapid, recurrent, and seemingly purposeless stereotyped motor contraction. Motor tics can be simple, with movement only affecting an individual muscle group (e.g., blinking, twitching of the nose, jerking of the neck), or complex, with coordinated involvement of multiple muscle groups (e.g., jumping, sniffing, head banging, and echopraxia [mimicking movements]). Phonic (or vocal) tics can also be simple (e.g., grunting) or complex (e.g., echolalia [repeating other people's words], palilalia [repeating one's own words], and coprolalia [expression of obscene words]). Patients may also experience sensory tics, composed of unpleasant focal sensations in the face, head, or neck. These can be mild and of little clinical consequence or severe and disabling to the patient.

TOURETTE'S SYNDROME (TS)

TS is a neurobehavioral disorder named after the French neurologist Georges Gilles de la Tourette. It predominantly affects males, and the prevalence is estimated to be 0.03–1.6%, but it is likely that many mild cases do not come to medical attention. TS is characterized by multiple motor tics often accompanied by vocalizations (phonic tics). Patients characteristically can voluntarily suppress tics for short periods of time, but then experience an irresistible urge to express them. Tics vary in intensity and may be absent for days or weeks only to recur, occasionally in a different pattern. Tics tend to present between ages 2 and 15 years (mean 7 years) and often lessen or even disappear in adulthood. Associated behavioral disturbances include anxiety, depression, attention deficit hyperactivity disorder, and obsessive-compulsive disorder. Patients may experience personality disorders, self-destructive behaviors, difficulties in school, and impaired interpersonal relationships. Tics may present in adulthood and can also be seen in association with a variety of other disorders, including PD, HD, trauma, dystonia, drugs (e.g., levodopa, neuroleptics), and toxins.

Etiology and Pathophysiology TS is thought to be a genetic disorder, but no specific gene mutation has been identified. Current evidence supports a complex inheritance pattern, with one or more major genes, multiple loci, low penetrance, and environmental influences. The risk of a family with one affected child having a second is about 25%. The pathophysiology of TS is not known, but alterations in dopamine

neurotransmission, opioids, and second-messenger systems have been proposed. Some cases of TS may be the consequence of an autoimmune response to β -hemolytic streptococcal infection (pediatric autoimmune neuropsychiatric disorder associated with streptococcal infection [PANDAS]); however, this entity remains controversial.

TREATMENT TOURETTE'S SYNDROME

Patients with mild disease often only require education and counseling (for themselves and family members). Drug treatment is indicated when the tics are disabling and interfere with quality of life. Therapy is individualized, and there is no singular treatment regimen that has been properly evaluated in double-blind trials. Some physicians use the α -agonist clonidine, starting at low doses and gradually increasing the dose and frequency until satisfactory control is achieved. Guanfacine (0.5–2 mg/d) is an α -agonist that is preferred by some because it only requires once-a-day dosing. Other physicians prefer to use neuroleptics. Atypical neuroleptics are usually used initially (risperidone, olanzapine, ziprasidone) because they are thought to be associated with a reduced risk of tardive dyskinesia. If they are not effective, low doses of classical neuroleptics such as haloperidol, fluphenazine, pimozide, or tiapride can be tried because the risk of tardive dyskinesia in young people is relatively low. Botulinum toxin injections can be effective in controlling focal tics that involve small muscle groups. Behavioral features, and particularly anxiety and compulsions, can be a disabling feature of TS and should be treated. The potential value of DBS targeting the anterior portion of the internal capsule, the GPi, or the thalamus is currently being explored.

MYOCLONUS

Myoclonus is a brief, rapid (<100 ms), shock-like, jerky movement consisting of single or repetitive muscle discharges. Myoclonic jerks can be focal, multifocal, segmental, or generalized and can occur spontaneously, in association with voluntary movement (action myoclonus) or in response to an external stimulus (reflex or startle myoclonus). Negative myoclonus consists of a brief loss of muscle activity (e.g., asterixis in hepatic failure). Myoclonic jerks can be severe and interfere with normal movement or benign and of no clinical consequence as is commonly observed in normal people when waking up or falling asleep (hypnogogic jerks).

Myoclonic jerks differ from tics in that they are not typically repetitive, can interfere with normal voluntary movement, and are not suppressible. They can arise in association with abnormal neuronal discharges in cortical, subcortical, brainstem, or spinal cord regions and can be associated with lesions in each of these regions, particularly in association with hypoxemia (especially following cardiac arrest), encephalopathy, and neurodegeneration. Reversible myoclonus can be seen with metabolic disturbances (renal failure, electrolyte imbalance, hypocalcemia), toxins, and many medications. Essential myoclonus is a relatively benign familial condition characterized by multifocal, very brief, lightning-like movements that are frequently alcohol sensitive. A mutation in the epsilon-sarcoglycan gene has been associated with a variety of myoclonus seen in association with dystonia (myoclonic dystonia).

TREATMENT MYOCLONUS

Treatment primarily consists of managing the underlying condition or removing an offending agent. Pharmacologic therapy involves one or a combination of GABAergic agents such as valproic acid (800–3000 mg/d), piracetam (8–20 g/d), clonazepam (2–15 mg/d), levetiracetam (1000–3000 mg/d), or primidone (500–1000 mg/d) and may be associated with striking clinical improvement in chronic cases (e.g., postanoxic myoclonus, progressive myoclonic epilepsy). The serotonin precursor 5-hydroxytryptophan (plus carbidopa) may be useful in some cases of postanoxic myoclonus.

This important group of movement disorders is primarily associated with drugs that block dopamine receptors (neuroleptics) or central dopaminergic transmission. These drugs are widely used in psychiatry, but it is important to appreciate that drugs used in the treatment of nausea or vomiting (e.g., prochlorperazine [Compazine]) or gastroesophageal disorders (e.g., metoclopramide) are neuroleptic agents. Hyperkinetic movement disorders secondary to neuroleptic drugs can be divided into those that present acutely, subacutely, or after prolonged exposure (tardive syndromes). Dopamine-blocking drugs can also be associated with a reversible parkinsonian syndrome for which anticholinergics are often concomitantly prescribed, but there is concern that this may increase the risk of developing a tardive syndrome.

ACUTE

Dystonia is the most common acute hyperkinetic drug reaction. It is typically generalized in children and focal in adults (e.g., blepharospasm, torticollis, or oromandibular dystonia). The reaction can develop within minutes of exposure and can be successfully treated in most cases with parenteral administration of anticholinergics (benztropine or diphenhydramine), benzodiazepines (lorazepam, clonazepam, or diazepam), or dopamine agonists. The abrupt onset of severe spasms may occasionally be confused with a seizure; however, there is no loss of consciousness, automatisms, or postictal features typical of epilepsy. The acute onset of chorea, stereotypic behavior, and tics may also be seen, particularly following exposure to CNS stimulants such as methylphenidate, cocaine, or amphetamines.

SUBACUTE

Akathisia is the commonest reaction in this category. It consists of motor restlessness with a need to move that is alleviated by movement. Therapy consists of removing the offending agent. When this is not possible, symptoms may be ameliorated with benzodiazepines, anticholinergics, beta blockers, or dopamine agonists.

TARDIVE SYNDROMES

These disorders develop months to years after initiation of neuroleptic treatment. Tardive dyskinesia (TD) is most common and typically presents with choreiform movements involving the mouth, lips, and tongue. In severe cases, the trunk, limbs, and respiratory muscles may also be affected. In approximately one-third of patients, TD remits within 3 months of stopping the drug, and most patients gradually improve over the course of several years. Abnormal movements may also develop or worsen after stopping the offending agent. The movements are often mild and more upsetting to the family than to the patient, but they can be severe and disabling, particularly in the context of an underlying psychiatric disorder. Atypical antipsychotics (e.g., clozapine, risperidone, olanzapine, quetiapine, ziprasidone, and aripiprazole) are thought to be associated with a lower risk of TD in comparison to traditional antipsychotics, although this remains to be established in controlled studies. Younger patients have a lower risk of developing neuroleptic-induced TD, whereas the elderly, females, and those with underlying organic cerebral dysfunction have been reported to be at greater risk. Chronic use is associated with increased risk, and specifically, the U.S. Food and Drug Administration has warned that use of metoclopramide for more than 12 weeks increases the risk of TD. Because TD can be permanent and resistant to treatment, antipsychotics should be used judiciously, atypical neuroleptics should be the preferred agent when possible, and the need for continued use should be regularly monitored.

Treatment primarily consists of stopping the offending agent. If the patient is receiving a traditional antipsychotic and withdrawal is not possible, replacement with an atypical antipsychotic should be tried. Abrupt cessation of a neuroleptic should be avoided because acute withdrawal can induce worsening. TD can persist after withdrawal of antipsychotics and can be difficult to treat. Benefits may occasionally be achieved with valproic acid, anticholinergics, or botulinum toxin injections. In refractory cases, catecholamine depleters such as tetrabenazine may be helpful, but this drug can be associated with dose-dependent

sedation and orthostatic hypotension and may induce parkinsonism as a side effect. Other approaches include baclofen (40–80 mg/d), clonazepam (1–8 mg/d), or valproic acid (750–3000 mg/d). In some cases, the abnormal movement is refractory to therapy.

Chronic neuroleptic exposure can also be associated with tardive dystonia, with preferential involvement of axial muscles and characteristic rocking movements of the trunk and pelvis. Tardive dystonia can be more troublesome than tardive dyskinesia and frequently persists despite stopping medication. Valproic acid, anticholinergics, and botulinum toxin may occasionally be beneficial, but patients are frequently refractory to medical therapy. Tardive akathisia, tardive Tourette's, and tardive tremor syndromes are rare but may also occur after chronic neuroleptic exposure.

Neuroleptic medications can also be associated with a neuroleptic malignant syndrome (NMS). NMS is characterized by the acute or subacute onset of muscle rigidity, elevated temperature, altered mental status, hyperthermia, tachycardia, labile blood pressure, renal failure, and markedly elevated creatine kinase levels. Symptoms typically evolve within days or weeks after initiating the drug. NMS can also be precipitated by the abrupt withdrawal of dopaminergic medications in PD patients. Treatment involves immediate cessation of the offending antipsychotic drug and the introduction of a dopaminergic agent (e.g., a dopamine agonist or levodopa), dantrolene, or a benzodiazepine. Treatment may need to be undertaken in an intensive care setting and include supportive measures such as control of body temperature (antipyretics and cooling blankets), hydration, electrolyte replacement, and control of renal function and blood pressure.

Drugs that have serotonin-like activity (tryptophan, MDMA or "ecstasy," meperidine) or that block serotonin reuptake can induce a rare, but potentially fatal, serotonin syndrome that is characterized by confusion, hyperthermia, tachycardia, and coma as well as rigidity, ataxia, and tremor. Myoclonus is often a prominent feature, in contrast to NMS, which it resembles. Patients can be managed with propranolol, diazepam, diphenhydramine, chlorpromazine, or cyproheptadine as well as supportive measures.

A variety of drugs can also be associated with parkinsonism (see above) and hyperkinetic movement disorders. Some examples include phenytoin (chorea, dystonia, tremor, myoclonus), carbamazepine (tics and dystonia), tricyclic antidepressants (dyskinesias, tremor, myoclonus), fluoxetine (myoclonus, chorea, dystonia), oral contraceptives (dyskinesia), β -adrenergics (tremor), buspirone (akathisia, dyskinesias, myoclonus), and digoxin, cimetidine, diazoxide, lithium, methadone, and fentanyl (dyskinesias).

PAROXYSMAL DYSKINESIAS

Paroxysmal dyskinesias are a group of rare disorders characterized by episodic, brief involuntary movements that can manifest as various types of hyperkinetic movements, including chorea, dystonia, tremor, and myoclonus. There are two main categories: (1) *paroxysmal kinesigenic dyskinesia*, where the involuntary movements are triggered by sudden movement, and (2) *paroxysmal nonkinesigenic dyskinesias*, where the attacks are not induced by movement. There are rare cases of *exercise-induced dyskinesia*, where attacks are induced by prolonged exercise.

Paroxysmal kinesigenic dyskinesia (PKD) is characterized by brief, self-limited attacks induced by movement onset such as running but also occasionally by unexpected sound or photic stimulation. Attacks may affect one side of the body, last seconds to minutes at a time, and recur several times a day. They usually manifest as dystonic posturing of a limb but may also become generalized. PKD is most commonly familial with an autosomal dominant pattern of inheritance but may also occur secondary to various brain disorders such as multiple sclerosis or hyperglycemia. PKD is more frequent in males (4:1), and the onset is typically in the first or second decade of life. About 70% report sensory symptoms such as tingling or numbness of the affected limb preceding the attack by a few milliseconds. The evolution is relatively benign, and there is a trend toward resolution of the attacks over time. The cause is not known, but a mutation in the proline-rich transmembrane protein 2 (*PRRT2*) gene that may be involved in

neurotransmitter release has now been identified. Treatment with low-dose anticonvulsant therapy such as carbamazepine or phenytoin is advised when the attacks are frequent and interfere with daily life activities and is effective in about 80% of patients. Some clinical features of PKD (abrupt and short-lasting attacks preceded by an "aura") and its favorable response to anticonvulsant drugs have led to speculation that it is epileptic in origin, but this has not been established.

Paroxysmal nonkinesigenic dyskinesia (PNKD) involves attacks of generalized dyskinesias precipitated by alcohol, caffeine, stress, or fatigue. In comparison to PKD, the episodes have a relatively longer duration (minutes to hours) and are less frequent (one to three per day). PNKD is inherited as autosomal dominant with incomplete penetrance pattern in some 80% of cases. A missense mutation in the myofibrillogenesis regulator (*MR-1*) gene has been identified in several families. Recognition of the condition and elimination of the underlying precipitating factors, where possible, are the first priority. Tetrabenazine, neuroleptics, dopamine-blocking agents, propranolol, clonazepam, and baclofen may be helpful. Treatment may not be required if the condition is mild and self-limited. Most patients with PNKD do not benefit from anticonvulsant drugs, but some may respond to clonazepam or other benzodiazepines.

RESTLESS LEGS SYNDROME

Restless legs syndrome (RLS) is a neurologic disorder that affects approximately 10% of the adult population (it is rare in Asians) and can cause significant morbidity in some. It was first described in the seventeenth century by an English physician (Thomas Willis), but has only recently been recognized as being a bona fide movement disorder. The four core symptoms required for diagnosis are as follows: an urge to move the legs, usually caused or accompanied by an unpleasant sensation in the legs; symptoms that begin or worsen with rest; partial or complete relief by movement; and worsening during the evening or night.

Symptoms most commonly begin in the legs, but can spread to or even begin in the upper limbs. The unpleasant sensation is often described as a creepy-crawly feeling, paresthesia, or burning. In about 80% of patients, RLS is associated with periodic leg movements (PLMs) during sleep and occasionally while awake. These involuntary movements are usually brief, lasting no more than a few seconds, and recur every 5–90 s. The restlessness and PLMs are a major cause of sleep disturbance in patients, leading to poor-quality sleep and daytime sleepiness.

RLS is a heterogeneous condition. Primary RLS is genetic, and several loci have been found with an autosomal dominant pattern of inheritance, although penetrance may be variable. The mean age of onset in genetic forms is 27 years, although pediatric cases are recognized. The severity of symptoms is variable. Secondary RLS may be associated with pregnancy or a range of underlying disorders, including anemia, ferritin deficiency, renal failure, and peripheral neuropathy. The pathogenesis probably involves disordered dopamine function, which may be peripheral or central, in association with an abnormality of iron metabolism. Diagnosis is made on clinical grounds but can be supported by polysomnography and the demonstration of PLMs. The neurologic examination is normal. Secondary RLS should be excluded, and ferritin levels, glucose, and renal function should be measured.

Most RLS sufferers have mild symptoms that do not require specific treatment. General measures to improve sleep hygiene and quality should be attempted first. If symptoms remain intrusive, low doses of dopamine agonists, e.g., pramipexole (0.25–0.5 mg) or ropinirole (1–2 mg), are given 1–2 h before bedtime. Levodopa can be effective but is frequently associated with augmentation (spread and worsening of restlessness and its appearance earlier in the day) or rebound (reappearance sometimes with worsening of symptoms at a time compatible with the drug's short half-life). Other drugs that can be effective include anticonvulsants, analgesics, and opiates. Management of secondary RLS should be directed to correcting the underlying disorder; for example, iron replacement for anemia. Iron infusion may also be helpful for severe primary RLS but requires expert supervision.

DISORDERS THAT MAY PRESENT WITH A COMBINATION OF PARKINSONISM AND HYPERKINETIC MOVEMENTS

WILSON'S DISEASE

Wilson's disease (WD) is an autosomal recessive inherited disorder of copper metabolism that may manifest with neurologic, psychiatric, and liver disorders, alone or in combination. It is caused by mutations in the gene encoding a P-type ATPase. The disease was first comprehensively described by the English neurologist Kinnier Wilson at the beginning of the twentieth century, although at around the same time the German physicians Kayser and Fleischer separately noted the characteristic association of corneal pigmentation with hepatic and neurologic features. WD has a worldwide prevalence of approximately 1 in 30,000, with a gene carrier frequency of 1 in 90. About half of WD patients (especially younger patients) manifest with liver abnormalities. The remainder present with neurologic disease (with or without underlying liver abnormalities), and a small proportion have hematologic or psychiatric problems at disease onset.

Neurologic onset usually manifests in the second decade with tremor and rigidity. The tremor is usually in the upper limbs, bilateral, and asymmetric. Tremor can be on intention or occasionally resting and, in advanced disease, can take on a wing-beating characteristic. Other features include parkinsonism with bradykinesia, dystonia (particularly facial grimacing), dysarthria, and dysphagia. More than half of those with neurologic features have a history of psychiatric disturbances, including depression, mood swings, and overt psychosis. Kayser-Fleischer (KF) rings are seen in 80% of those with hepatic presentations and virtually all with neurologic features. KF rings represent the deposition of copper in Descemet's membrane around the cornea. They consist of a characteristic grayish rim or circle at the limbus of the cornea and are best detected by slit-lamp examination. Neuropathologic examination is characterized by neurodegeneration and astrogliosis in the basal ganglia, particularly in the striatum.

WD should always be considered in the differential diagnosis of a movement disorder in the first decades of life. Low levels of blood copper and ceruloplasmin and high levels of urinary copper may be present, but normal levels do not exclude the diagnosis. A computed tomography (CT) scan usually reveals generalized brain atrophy in established cases, and ~50% have signal hypointensity in the caudate head, putamen, globus pallidum, substantia nigra, and red nucleus on T2-weighted MRI. However, correlation of imaging changes with clinical features is not good. It is very rare for WD patients with neurologic features not to have KF rings, and therefore when the diagnosis is considered, examination by slit-lamp is essential. Liver biopsy with demonstration of high copper levels remains the gold standard for the diagnosis.

In the absence of treatment, the course is progressive and leads to severe neurologic dysfunction and early death. Treatment is directed at reducing tissue copper levels and maintenance therapy to prevent reaccumulation. There is no clear consensus on treatment, and all patients should be managed in a unit with expertise in WD. Penicillamine is frequently used to increase copper excretion, but it may lead to a worsening of symptoms in the initial stages of therapy. Side effects are common and can to some degree be attenuated by coadministration of pyridoxine. Tetrathiomolybdate blocks the absorption of copper and can be used instead of penicillamine. Trientine and zinc are useful drugs for maintenance therapy. Effective treatment can reverse the neurologic features in most patients, particularly when started early. Some patients stabilize, and a few may still progress, especially those with hepatocerebral disease. KF rings tend to decrease after 3–6 months and disappear by 2 years. Adherence to maintenance therapy is a major challenge in long-term care.

NEURODEGENERATION WITH BRAIN IRON ACCUMULATION

Neurodegeneration with brain iron accumulation (NBIA) represents a group of inherited disorders characterized by iron accumulation in the basal ganglia. Clinically, they can manifest as a progressive neurologic disorder manifesting a variety of features including parkinsonism,

dystonia, neuropsychiatric abnormalities, and retinal degeneration. Cognitive disorders and cerebellar dysfunction may also be seen. Presentation is usually in childhood, but adult cases have been described. Multiple genes have been identified to date. Pantothenate kinase-associated neurodegeneration (PKAN) formerly known as Hallervorden-Spatz disease and caused by a mutation in the *PANK2* gene is the most common form of NBIA, accounting for about 50% of cases. Onset is usually in early childhood and is manifest as a combination of dystonia, parkinsonism, and spasticity. MRI shows a characteristic low signal abnormality in the center of the globus pallidus on T2-weighted scans known as the “eye of the tiger” sign caused by iron accumulation. Numerous other gene mutations have been described associated with iron accumulation including mutations in *PLA2G6*, *C19orf12*, *FA2H*, *ATP13A2*, *WDR45*, *FTL*, *CP*, and *DCAF17*. One must be cautious, however, not to assume that all cases with iron accumulation in the basal ganglia represent an NBIA, because iron accumulation in specific basal ganglia regions is normal, and excess iron accumulation may occur in the basal ganglia region as a consequence of neurodegeneration of multiple causes unrelated to a defect in iron metabolism.

OTHER DISORDERS

Acanthocytosis, some hereditary spinocerebellar atrophies and spastic parapareses, and HD can also present with parkinsonian features associated with involuntary movements. Diagnosis in these cases is best established with genetic testing.

PSYCHOGENIC DISORDERS

Virtually all movement disorders including tremor, tics, dystonia, myoclonus, chorea, ballism, and parkinsonism can be psychogenic in origin. Tremor affecting the upper limbs is the most common psychogenic movement disorder. Psychogenic movements can result from a somatoform or conversion disorder, malingering (e.g., seeking financial gain), or a factitious disorder (e.g., seeking psychological gain). Psychogenic movement disorders are common (estimated to be 2–3% of patients seen in a movement disorder clinic), more frequent in women, disabling for the patient and family, and expensive for society (estimated \$20 billion annually). Clinical features suggesting a psychogenic movement disorder include an acute onset and a pattern of abnormal movement that is inconsistent with a known movement disorder. Diagnosis is based on the nonorganic quality of the movement, the absence of findings of an organic disease process, and positive features that specifically point to a psychogenic illness such as variability and distractibility. For example, the magnitude of a psychogenic tremor is increased with attention and diminishes or even disappears when the patient is distracted by being asked to perform a different task or is unaware that he or she is being observed. Other positive features suggesting a psychogenic problem include a tremor frequency that is variable or that entrains with the frequency of a designated movement in the contralateral limb, and a positive response to placebo medication. Associated features can include nonanatomic sensory findings, give-way weakness, astasia-abasia (an odd, gyrating gait; [Chap. 32](#)), and multiple somatic complaints with no underlying pathology (somatoform disorder). Comorbid psychiatric problems such as anxiety, depression, and emotional trauma may be present but are not necessary for the diagnosis of a psychogenic movement disorder to be made. Psychogenic movement disorders can occur as an isolated entity or in association with an underlying organic problem. The diagnosis can often be made based on clinical features alone, and unnecessary tests or medications can be avoided. Underlying psychiatric problems may be present and should be identified and treated, but many patients with psychogenic movement disorders have no obvious psychiatric pathology. Psychotherapy and hypnosis may be of value for patients with conversion reaction, and cognitive behavioral therapy may be helpful for patients with somatoform disorders. Patients with hypochondriasis, factitious disorders, and malingering have a poor prognosis.

450 Ataxic Disorders

Roger N. Rosenberg

APPROACH TO THE PATIENT: Ataxic Disorders

Symptoms and signs of ataxia consist of gait impairment, unclear (“scanning”) speech, visual blurring due to nystagmus, hand incoordination, and tremor with movement. These result from the involvement of the cerebellum and its afferent and efferent pathways, including the spinocerebellar pathways, and the frontopontocerebellar pathway originating in the rostral frontal lobe. True cerebellar ataxia must be distinguished from ataxia associated with vestibular nerve or labyrinthine disease, as the latter results in a disorder of gait associated with a significant degree of dizziness, light-headedness, or the perception of movement ([Chap. 28](#)). True cerebellar ataxia is devoid of these vertiginous complaints and is clearly an unsteady gait due to imbalance. Sensory disturbances can also on occasion simulate the imbalance of cerebellar disease; with sensory ataxia, imbalance dramatically worsens when visual input is removed (Romberg sign). Rarely, weakness of proximal leg muscles mimics cerebellar disease. In the patient who presents with ataxia, the rate and pattern of the development of cerebellar symptoms help to narrow the diagnostic possibilities ([Table 450-1](#)). A gradual and progressive increase in symptoms with bilateral and symmetric involvement suggests a genetic, metabolic, immune, or toxic etiology. Conversely, focal, unilateral symptoms with headache and impaired level of consciousness accompanied by ipsilateral cranial nerve palsies and contralateral weakness imply a space-occupying cerebellar lesion.

SYMMETRIC ATAXIA

Progressive and symmetric ataxia can be classified with respect to onset as acute (over hours or days), subacute (weeks or months), or chronic (months to years). Acute and reversible ataxias include those caused by intoxication with alcohol, phenytoin, lithium, barbiturates, and other drugs. Intoxication caused by toluene exposure, gasoline sniffing, glue sniffing, spray painting, or exposure to methyl mercury or bismuth are additional causes of acute or subacute ataxia, as is treatment with cytotoxic chemotherapeutic drugs such as fluorouracil and paclitaxel. Patients with a postinfectious syndrome (especially after varicella) may develop gait ataxia and mild dysarthria, both of which are reversible ([Chap. 458](#)). Rare infectious causes of acquired ataxia include poliovirus, coxsackievirus, echovirus, Epstein-Barr virus, toxoplasmosis, *Legionella*, and Lyme disease.

The subacute development of ataxia of gait over weeks to months (degeneration of the cerebellar vermis) may be due to the combined effects of alcoholism and malnutrition, particularly with deficiencies of vitamins B₁ and B₁₂. Hyponatremia has also been associated with ataxia. Paraneoplastic cerebellar ataxia is associated with a number of different tumors (and autoantibodies) such as breast and ovarian cancers (anti-Yo), small-cell lung cancer (anti-PQ-type voltage-gated calcium channel), and Hodgkin's disease (anti-Tr) ([Chap. 122](#)). Another paraneoplastic syndrome associated with myoclonus and opsoclonus occurs with breast (anti-Ri) and lung cancers and neuroblastoma. Elevated serum anti-glutamic acid decarboxylase (GAD) antibodies have been associated with a progressive ataxic syndrome affecting speech and gait. For all of these paraneoplastic ataxias, the neurologic syndrome may be the presenting symptom of the cancer. Another immune-mediated progressive ataxia is associated with antigliadin (and antiendomysium) antibodies and the human leukocyte antigen (HLA) DQB1*0201 haplotype; in some affected patients, biopsy of the small intestine reveals villus atrophy consistent with gluten-sensitive enteropathy ([Chap. 349](#)). Finally, subacute progressive ataxia may be caused by a prion disorder,

TABLE 450-1 ETIOLOGY OF CEREBELLAR ATAXIA

Symmetric and Progressive Signs			Focal and Ipsilateral Cerebellar Signs		
Acute (Hours to Days)	Subacute (Days to Weeks)	Chronic (Months to Years)	Acute (Hours to Days)	Subacute (Days to Weeks)	Chronic (Months to Years)
Intoxication: alcohol, lithium, phenytoin, barbiturates (positive history and toxicology screen)	Intoxication: mercury, solvents, gasoline, glue; cytotoxic chemotherapeutic, hemotherapeutic drugs	Paraneoplastic syndrome Antigliadin antibody syndrome Hypothyroidism	Vascular: cerebellar infarction, hemorrhage, or subdural hematoma	Neoplastic: cerebellar glioma or metastatic tumor (positive for neoplasm on MRI/CT)	Stable gliosis secondary to vascular lesion or demyelinating plaque (stable lesion on MRI/CT older than several months)
Acute viral cerebellitis (CSF supportive of acute viral infection)	Alcoholic-nutritional (vitamin B ₁ and B ₁₂ deficiency)	Inherited diseases Tabes dorsalis (tertiary syphilis)	Infectious: cerebellar abscess (mass lesion on MRI/CT, history in support of lesion)	Demyelinating: multiple sclerosis (history, CSF, and MRI are consistent)	Congenital lesion: Chiari or Dandy-Walker malformations (malformation noted on MRI/CT)
Postinfection syndrome	Lyme disease	Phenytoin toxicity Amiodarone		AIDS-related multifocal leukoencephalopathy (positive HIV test and CD4+ cell count for AIDS)	

Abbreviations: CSF, cerebrospinal fluid; CT, computed tomography; MRI, magnetic resonance imaging.

especially when an infectious etiology, such as transmission from contaminated human growth hormone, is responsible (**Chap. 453e**).

Chronic symmetric gait ataxia suggests an inherited ataxia (discussed below), a metabolic disorder, or a chronic infection. Hypothyroidism must always be considered as a readily treatable and reversible form of gait ataxia. Infectious diseases that can present with ataxia are meningovascular syphilis and tabes dorsalis due to degeneration of the posterior columns and spinocerebellar pathways in the spinal cord.

FOCAL ATAXIA

Acute focal ataxia commonly results from cerebrovascular disease, usually ischemic infarction or cerebellar hemorrhage. These lesions typically produce cerebellar symptoms ipsilateral to the injured cerebellum and may be associated with an impaired level of consciousness due to brainstem compression and increased intracranial pressure; ipsilateral pontine signs, including sixth and seventh nerve palsies, may be present. Focal and worsening signs of acute ataxia should also prompt consideration of a posterior fossa subdural hematoma, bacterial abscess, or primary or metastatic cerebellar tumor. Computed tomography (CT) or magnetic resonance imaging (MRI) studies will reveal clinically significant processes of this type. Many of these lesions represent true neurologic emergencies, as sudden herniation, either rostrally through the tentorium or caudal herniation of cerebellar tonsils through the foramen magnum, can occur and is usually devastating. Acute surgical decompression may be required (**Chap. 330**). Lymphoma or progressive multifocal leukoencephalopathy (PML) in a patient with AIDS may present with an acute or subacute focal cerebellar syndrome. Chronic etiologies of progressive ataxia include multiple sclerosis (**Chap. 458**) and congenital lesions such as a Chiari malformation (**Chap. 456**) or a congenital cyst of the posterior fossa (Dandy-Walker syndrome).

THE INHERITED ATAXIAS

These may show autosomal dominant, autosomal recessive, or maternal (mitochondrial) modes of inheritance. A genomic classification (**Chap. 451e**) has now largely superseded previous ones based on clinical expression alone.

Although the clinical manifestations and neuropathologic findings of cerebellar disease dominate the clinical picture, there may also be characteristic changes in the basal ganglia, brainstem, spinal cord, optic nerves, retina, and peripheral nerves. In large families with dominantly inherited ataxias, many gradations are observed from purely cerebellar manifestations to mixed cerebellar and brainstem disorders, cerebellar and basal ganglia syndromes, and spinal cord

or peripheral nerve disease. Rarely, dementia is present as well. The clinical picture may be homogeneous within a family with dominantly inherited ataxia, but sometimes most affected family members show one characteristic syndrome, while one or several members have an entirely different phenotype.

AUTOSOMAL DOMINANT ATAXIAS

The autosomal spinocerebellar ataxias (SCAs) include SCA types 1 through 36, dentatorubropallidoluysian atrophy (DRPLA), and episodic ataxia (EA) types 1 to 7 (**Chap. 451e**). SCA1, SCA2, SCA3 (Machado-Joseph disease [MJD]), SCA6, SCA7, and SCA17 are caused by CAG triplet repeat expansions in different genes. SCA8 is due to an untranslated CTG repeat expansion, SCA12 is linked to an untranslated CAG repeat, and SCA10 is caused by an untranslated pentanucleotide repeat. The clinical phenotypes of these SCAs overlap. The genotype has become the gold standard for diagnosis and classification. CAG encodes glutamine, and these expanded CAG triplet repeat expansions result in expanded polyglutamine proteins, termed *ataxins*, that produce a toxic gain of function with autosomal dominant inheritance. Although the phenotype is variable for any given disease gene, a pattern of neuronal loss with gliosis is produced that is relatively unique for each ataxia. Immunohistochemical and biochemical studies have shown cytoplasmic (SCA2), neuronal (SCA1, MJD, SCA7), and nucleolar (SCA7) accumulation of the specific mutant polyglutamine-containing ataxin proteins. Expanded polyglutamine ataxins with more than ~40 glutamines are potentially toxic to neurons for a variety of reasons including: high levels of gene expression for the mutant polyglutamine ataxin in affected neurons; conformational change of the aggregated protein to a β -pleated structure; abnormal transport of the ataxin into the nucleus (SCA1, MJD, SCA7); binding to other polyglutamine proteins, including the TATA-binding transcription protein and the CREB-binding protein, impairing their functions; altering the efficiency of the ubiquitin-proteasome system of protein turnover; and inducing neuronal apoptosis. An earlier age of onset (anticipation) and more aggressive disease in subsequent generations are due to further expansion of the CAG triplet repeat and increased polyglutamine number in the mutant ataxin. The most common disorders are discussed below.

SCA1

SCA1 was previously referred to as *olivopontocerebellar atrophy*, but genomic data have shown that that entity represents several different genotypes with overlapping clinical features.

Symptoms and Signs SCA1 is characterized by the development in early or middle adult life of progressive cerebellar ataxia of the trunk and limbs, impairment of equilibrium and gait, slowness of voluntary movements, scanning speech, nystagmoid eye movements, and oscillatory tremor of the head and trunk. Dysarthria, dysphagia, and

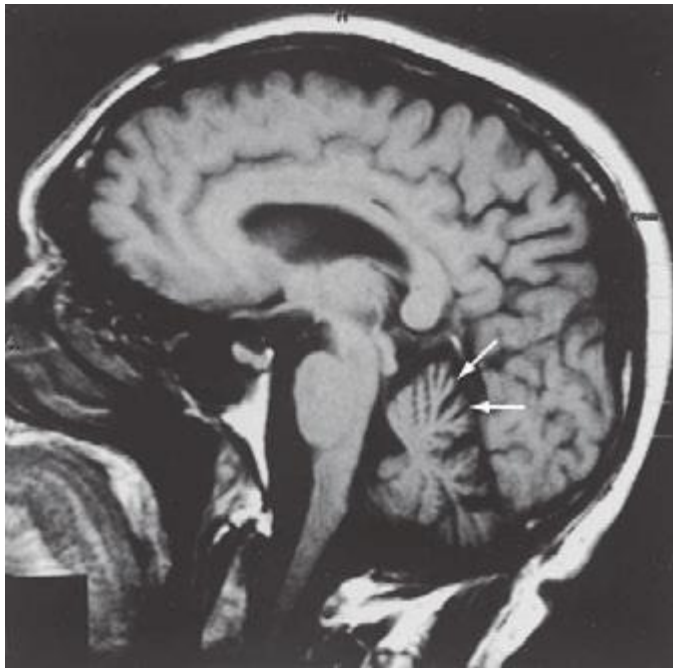


FIGURE 450-1 Sagittal magnetic resonance imaging (MRI) of the brain of a 60-year-old man with gait ataxia and dysarthria due to spinocerebellar ataxia type 1 (SCA1), illustrating cerebellar atrophy (arrows). (Reproduced with permission from RN Rosenberg, P Khemani, in RN Rosenberg, JM Pascual [eds]: *Rosenberg's Molecular and Genetic Basis of Neurological and Psychiatric Disease*, 5th ed. London, Elsevier, 2015.)

oculomotor and facial palsies may also occur. Extrapyramidal symptoms include rigidity, an immobile face, and parkinsonian tremor. The reflexes are usually normal, but knee and ankle jerks may be lost, and extensor plantar responses may occur. Dementia may be noted but is usually mild. Impairment of sphincter function is common, with urinary and sometimes fecal incontinence. Cerebellar and brainstem atrophy are evident on MRI (Fig. 450-1).

Marked shrinkage of the ventral half of the pons, disappearance of the olivary eminence on the ventral surface of the medulla, and atrophy of the cerebellum are evident on gross postmortem inspection of the brain. Variable loss of Purkinje cells, reduced numbers of cells in the molecular and granular layer, demyelination of the middle cerebellar peduncle and the cerebellar hemispheres, and severe loss of cells in the pontine nuclei and olives are found on histologic examination. Degenerative changes in the striatum, especially the putamen, and loss of the pigmented cells of the substantia nigra may be found in cases with extrapyramidal features. More widespread degeneration in the central nervous system (CNS), including involvement of the posterior columns and the spinocerebellar fibers, is often present.

GENETIC CONSIDERATIONS

SCA1 encodes a gene product, called *ataxin-1*, which is a novel protein of unknown function. The mutant allele has 40 CAG repeats located within the coding region, whereas alleles from unaffected individuals have ≤ 36 repeats. A few patients with 38–40 CAG repeats have been described. There is a direct correlation between a larger number of repeats and a younger age of onset for SCA1. Juvenile patients have higher numbers of repeats, and anticipation is present in subsequent generations. Transgenic mice carrying SCA1 developed ataxia and Purkinje cell pathology. Nuclear localization, but not aggregation, of ataxin-1 appears to be required for cell death initiated by the mutant protein.

SCA2

Symptoms and Signs Another clinical phenotype, SCA2, has been described in patients from Cuba and India. Cuban patients probably

are descendants of a common ancestor, and the population may be the largest homogeneous group of patients with ataxia yet described. The age of onset ranges from 2–65 years, and there is considerable clinical variability within families. Although neuropathologic and clinical findings are compatible with a diagnosis of SCA1, including slow saccadic eye movements, ataxia, dysarthria, parkinsonian rigidity, optic disc pallor, mild spasticity, and retinal degeneration, SCA2 is a unique form of cerebellar degenerative disease.

GENETIC CONSIDERATIONS

The gene in SCA2 families also contains CAG repeat expansions coding for a polyglutamine-containing protein, ataxin-2. Normal alleles contain 15–32 repeats; mutant alleles have 35–77 repeats.

MACHADO-JOSEPH DISEASE/SCA3

MJD was first described among the Portuguese and their descendants in New England and California. Subsequently, MJD has been found in families from Portugal, Australia, Brazil, Canada, China, England, France, India, Israel, Italy, Japan, Spain, Taiwan, and the United States. In most populations, it is the most common autosomal dominant ataxia.

Symptoms and Signs MJD has been classified into three clinical types. In type I MJD (amyotrophic lateral sclerosis-parkinsonism-dystonia type), neurologic deficits appear in the first two decades and involve weakness and spasticity of extremities, especially the legs, often with dystonia of the face, neck, trunk, and extremities. Patellar and ankle clonus are common, as are extensor plantar responses. The gait is slow and stiff, with a slightly broadened base and lurching from side to side; this gait results from spasticity, not true ataxia. There is no truncal titubation. Pharyngeal weakness and spasticity cause difficulty with speech and swallowing. Of note is the prominence of horizontal and vertical nystagmus, loss of fast saccadic eye movements, hypermetric and hypometric saccades, and impairment of upward vertical gaze. Facial fasciculations, facial myokymia, lingual fasciculations without atrophy, ophthalmoparesis, and ocular prominence are common early manifestations.

In type II MJD (ataxic type), true cerebellar deficits of dysarthria and gait and extremity ataxia begin in the second to fourth decades along with corticospinal and extrapyramidal deficits of spasticity, rigidity, and dystonia. Type II is the most common form of MJD. Ophthalmoparesis, upward vertical gaze deficits, and facial and lingual fasciculations are also present. Type II MJD can be distinguished from the clinically similar disorders SCA1 and SCA2.

Type III MJD (ataxic-amyotrophic type) presents in the fifth to the seventh decades with a pancerebellar disorder that includes dysarthria and gait and extremity ataxia. Distal sensory loss involving pain, touch, vibration, and position senses and distal atrophy are prominent, indicating the presence of peripheral neuropathy. The deep tendon reflexes are depressed to absent, and there are no corticospinal or extrapyramidal findings.

The mean age of onset of symptoms in MJD is 25 years. Neurologic deficits invariably progress and lead to death from debilitation within 15 years of onset, especially in patients with types I and II disease. Usually, patients retain full intellectual function.

The major pathologic findings are variable loss of neurons and glial replacement in the corpus striatum and severe loss of neurons in the pars compacta of the substantia nigra. A moderate loss of neurons occurs in the dentate nucleus of the cerebellum and in the red nucleus. Purkinje cell loss and granule cell loss occur in the cerebellar cortex. Cell loss also occurs in the dentate nucleus and in the cranial nerve motor nuclei. Sparing of the inferior olives distinguishes MJD from other dominantly inherited ataxias.

GENETIC CONSIDERATIONS

The gene for MJD maps to 14q24.3-q32. Unstable CAG repeat expansions are present in the MJD gene coding for a polyglutamine-containing protein named ataxin-3, or MJD-ataxin. An earlier age of onset is associated with longer repeats. Alleles from normal individuals have between 12 and 37 CAG repeats,

whereas MJD alleles have 60–84 CAG repeats. Polyglutamine-containing aggregates of ataxin-3 (MJD-ataxin) have been described in neuronal nuclei undergoing degeneration. MJD-ataxin codes for a ubiquitin protease, which is inactive due to expanded polyglutamines. Proteasome function is impaired, resulting in altered clearance of proteins and cerebellar neuronal loss.

SCA6

Genomic screening for CAG repeats in other families with autosomal dominant ataxia and vibratory and proprioceptive sensory loss have yielded another locus. Of interest is that different mutations in the same gene for the α_{1A} voltage-dependent calcium channel subunit (CACN1A4; also referred to as the CACNA1A gene) at 19p13 result in different clinical disorders. CAG repeat expansions (21–27 in patients; 4–16 triplets in normal individuals) result in late-onset progressive ataxia with cerebellar degeneration. Missense mutations in this gene result in familial hemiplegic migraine. Nonsense mutations resulting in termination of protein synthesis of the gene product yield hereditary paroxysmal cerebellar ataxia or EA. Some patients with familial hemiplegic migraine develop progressive ataxia and also have cerebellar atrophy.

SCA7

This disorder is distinguished from all other SCAs by the presence of retinal pigmentary degeneration. The visual abnormalities first appear as blue-yellow color blindness and proceed to frank visual loss with macular degeneration. In almost all other respects, SCA7 resembles several other SCAs in which ataxia is accompanied by various noncerebellar findings, including ophthalmoparesis and extensor plantar responses. The genetic defect is an expanded CAG repeat in the SCA7 gene at 3p14-p21.1. The expanded repeat size in SCA7 is highly variable. Consistent with this, the severity of clinical findings varies from essentially asymptomatic to mild late-onset symptoms to severe, aggressive disease in childhood with rapid progression. Marked anticipation has been recorded, especially with paternal transmission. The disease protein, ataxin-7, forms aggregates in nuclei of affected neurons, as has also been described for SCA1 and SCA3/MJD.

SCA8

This form of ataxia is caused by a CTG repeat expansion in an untranslated region of a gene on chromosome 13q21. There is marked maternal bias in transmission, perhaps reflecting contractions of the repeat during spermatogenesis. The mutation is not fully penetrant. Symptoms include slowly progressive dysarthria and gait ataxia beginning at ~40 years of age with a range between 20 and 65 years. Other features include nystagmus, leg spasticity, and reduced vibratory sensation. Severely affected individuals are nonambulatory by the fourth to sixth decades. MRI shows cerebellar atrophy. The mechanism of disease may involve a dominant “toxic” effect occurring at the RNA level, as occurs in myotonic dystrophy.

DENTATORUBROPALLIDOLUYSIAN ATROPHY

DRPLA has a variable presentation that may include progressive ataxia, choreoathetosis, dystonia, seizures, myoclonus, and dementia. DRPLA is due to unstable CAG triplet repeats in the open reading frame of a gene named *atrophin* located on chromosome 12p12-ter. Larger expansions are found in patients with earlier onset. The number of repeats is 49 in patients with DRPLA and ≤ 26 in normal individuals. Anticipation occurs in successive generations, with earlier onset of disease in association with an increasing CAG repeat number in children who inherit the disease from their father. One well-characterized family in North Carolina has a phenotypic variant known as the *Haw River syndrome*, now recognized to be due to the DRPLA mutation.

EPISODIC ATAXIA

EA types 1 and 2 are two rare dominantly inherited disorders that have been mapped to chromosomes 12p (a potassium channel gene) for type 1 and 19p for type 2. Patients with EA-1 have brief episodes of ataxia with myokymia and nystagmus that last only minutes. Startle, sudden change in posture, and exercise can induce episodes. Acetazolamide or

anticonvulsants may be therapeutic. Patients with EA-2 have episodes of ataxia with nystagmus that can last for hours or days. Stress, exercise, or excessive fatigue may be precipitants. Acetazolamide may be therapeutic and can reverse the relative intracellular alkalosis detected by magnetic resonance spectroscopy. Stop codon, nonsense mutations causing EA-2 have been found in the CACNA1A gene, encoding the α_{1A} voltage-dependent calcium channel subunit (see “SCA6,” above).

AUTOSOMAL RECESSIVE ATAXIAS

Friedreich's Ataxia This is the most common form of inherited ataxia, comprising one-half of all hereditary ataxias. It can occur in a classic form or in association with a genetically determined vitamin E deficiency syndrome; the two forms are clinically indistinguishable.

SYMPTOMS AND SIGNS Friedreich's ataxia presents before 25 years of age with progressive staggering gait, frequent falling, and titubation. The lower extremities are more severely involved than the upper ones. Dysarthria occasionally is the presenting symptom; rarely, progressive scoliosis, foot deformity, nystagmus, or cardiopathy is the initial sign.

The neurologic examination reveals nystagmus, loss of fast saccadic eye movements, truncal titubation, dysarthria, dysmetria, and ataxia of trunk and limb movements. Extensor plantar responses (with normal tone in trunk and extremities), absence of deep tendon reflexes, and weakness (greater distally than proximally) are usually found. Loss of vibratory and proprioceptive sensation occurs. The median age of death is 35 years. Women have a significantly better prognosis than men.

Cardiac involvement occurs in 90% of patients. Cardiomegaly, symmetric hypertrophy, murmurs, and conduction defects are reported. Moderate mental retardation or psychiatric syndromes are present in a small percentage of patients. A high incidence of diabetes mellitus (20%) is found and is associated with insulin resistance and pancreatic β -cell dysfunction. Musculoskeletal deformities are common and include pes cavus, pes equinovarus, and scoliosis. MRI of the spinal cord shows atrophy (Fig. 450-2).

The primary sites of pathology are the spinal cord, dorsal root ganglion cells, and the peripheral nerves. Slight atrophy of the cerebellum and cerebral gyri may occur. Sclerosis and degeneration occur

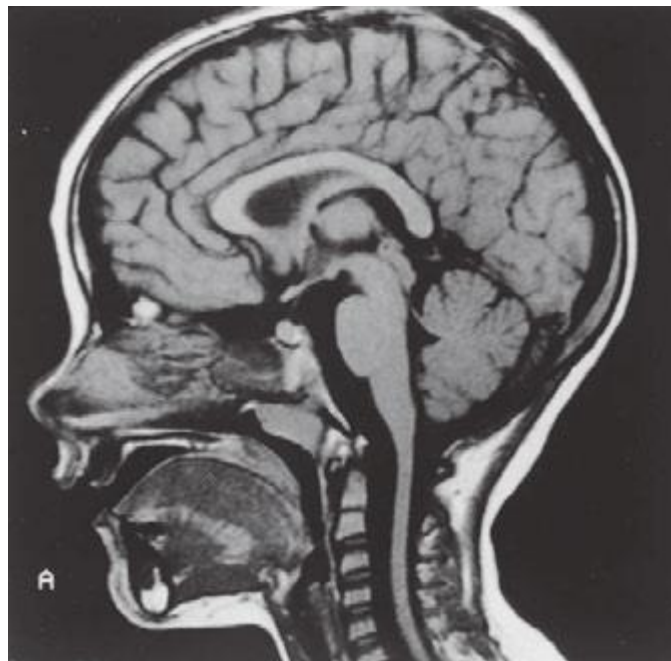


FIGURE 450-2 Sagittal magnetic resonance imaging (MRI) of the brain and spinal cord of a patient with Friedreich's ataxia, demonstrating spinal cord atrophy. (Reproduced with permission from RN Rosenberg, P Khemani, in RN Rosenberg, JM Pascual [eds]: *Rosenberg's Molecular and Genetic Basis of Neurological and Psychiatric Disease*, 5th ed. London, Elsevier, 2015.)

predominantly in the spinocerebellar tracts, lateral corticospinal tracts, and posterior columns. Degeneration of the glossopharyngeal, vagus, hypoglossal, and deep cerebellar nuclei is described. The cerebral cortex is histologically normal except for loss of Betz cells in the precentral gyri. The peripheral nerves are extensively involved, with a loss of large myelinated fibers. Cardiac pathology consists of myocytic hypertrophy and fibrosis, focal vascular fibromuscular dysplasia with subintimal or medial deposition of periodic acid-Schiff (PAS)-positive material, myocytopathy with unusual pleomorphic nuclei, and focal degeneration of nerves and cardiac ganglia.

GENETIC CONSIDERATIONS



The classic form of Friedreich's ataxia has been mapped to 9q13-q21.1, and the mutant gene, *frataxin*, contains expanded GAA triplet repeats in the first intron. There is homozygosity for expanded GAA repeats in >95% of patients. Normal persons have 7–22 GAA repeats, and patients have 200–900 GAA repeats. A more varied clinical syndrome has been described in compound heterozygotes who have one copy of the GAA expansion and the other copy a point mutation in the *frataxin* gene. When the point mutation is located in the region of the gene that encodes the amino-terminal half of frataxin, the phenotype is milder, often consisting of a spastic gait, retained or exaggerated reflexes, no dysarthria, and mild or absent ataxia.

Patients with Friedreich's ataxia have undetectable or extremely low levels of *frataxin* mRNA, as compared with carriers and unrelated individuals; thus, disease appears to be caused by a loss of expression of the frataxin protein. Frataxin is a mitochondrial protein involved in iron homeostasis. Mitochondrial iron accumulation due to loss of the iron transporter coded by the mutant *frataxin* gene results in oxidized intramitochondrial iron. Excess oxidized iron results in turn in the oxidation of cellular components and irreversible cell injury.

Two forms of hereditary ataxia associated with abnormalities in the interactions of vitamin E (α -tocopherol) with very-low-density lipoprotein (VLDL) have been delineated. These are abetalipoproteinemia (Bassen-Kornzweig syndrome) and ataxia with vitamin E deficiency (AVED). Abetalipoproteinemia is caused by mutations in the gene coding for the larger subunit of the microsomal triglyceride transfer protein (MTP). Defects in MTP result in impairment of formation and secretion of VLDL in liver. This defect results in a deficiency of delivery of vitamin E to tissues, including the central and peripheral nervous system, as VLDL is the transport molecule for vitamin E and other fat-soluble substitutes. AVED is due to mutations in the gene for α -tocopherol transfer protein (α -TTP). These patients have an impaired ability to bind vitamin E into the VLDL produced and secreted by the liver, resulting in a deficiency of vitamin E in peripheral tissues. Hence, either absence of VLDL (abetalipoproteinemia) or impaired binding of vitamin E to VLDL (AVED) causes an ataxic syndrome. Once again, a genotype classification has proved to be essential in sorting out the various forms of the Friedreich's disease syndrome, which may be clinically indistinguishable.

Ataxia Telangiectasia • SYMPTOMS AND SIGNS Patients with ataxia telangiectasia (AT) present in the first decade of life with progressive telangiectatic lesions associated with deficits in cerebellar function and nystagmus. The neurologic manifestations correspond to those in Friedreich's disease, which should be included in the differential diagnosis. Truncal and limb ataxia, dysarthria, extensor plantar responses, myoclonic jerks, areflexia, and distal sensory deficits may develop. There is a high incidence of recurrent pulmonary infections and neoplasms of the lymphatic and reticuloendothelial system in patients with AT. Thymic hypoplasia with cellular and humoral (IgA and IgG2) immunodeficiencies, premature aging, and endocrine disorders such as type 1 diabetes mellitus are described. There is an increased incidence of lymphomas, Hodgkin's disease, acute T cell leukemias, and breast cancer.

The most striking neuropathologic changes include loss of Purkinje, granule, and basket cells in the cerebellar cortex as well as of neurons in the deep cerebellar nuclei. The inferior olives of the medulla may also have neuronal loss. There is a loss of anterior horn neurons in the

spinal cord and of dorsal root ganglion cells associated with posterior column spinal cord demyelination. A poorly developed or absent thymus gland is the most consistent defect of the lymphoid system.

GENETIC CONSIDERATIONS



The gene for AT (the *ATM* gene) encodes a protein that is similar to several yeast and mammalian phosphatidylinositol-3'-kinases involved in mitogenic signal transduction, meiotic recombination, and cell cycle control. Defective DNA repair in AT fibroblasts exposed to ultraviolet light has been demonstrated. The discovery of *ATM* permits early diagnosis and identification of heterozygotes who are at risk for cancer (e.g., breast cancer).

MITOCHONDRIAL ATAXIAS

Spinocerebellar syndromes have been identified with mutations in mitochondrial DNA (mtDNA). Thirty pathogenic mtDNA point mutations and 60 different types of mtDNA deletions are known, several of which cause or are associated with ataxia (**Chap. 462e**).

TREATMENT ATAXIC DISORDERS

The most important goal in management of patients with ataxia is to identify treatable disease entities. Mass lesions must be recognized promptly and treated appropriately. Paraneoplastic disorders can often be identified by the clinical patterns of disease that they produce, measurement of specific autoantibodies, and uncovering the primary cancer; these disorders are often refractory to therapy, but some patients improve following removal of the tumor or immunotherapy (**Chap. 122**). Ataxia with antigliadin antibodies and gluten-sensitive enteropathy may improve with a gluten-free diet. Malabsorption syndromes leading to vitamin E deficiency may lead to ataxia. The vitamin E deficiency form of Friedreich's ataxia must be considered, and serum vitamin E levels measured. Vitamin E therapy is indicated for these rare patients. Vitamin B₁₂ levels in serum should be measured, and the vitamins administered to patients having deficient levels. Hypothyroidism is easily treated. The cerebrospinal fluid should be tested for a syphilitic infection in patients with progressive ataxia and other features of tabes dorsalis. Similarly, antibody titers for Lyme disease and *Legionella* should be measured and appropriate antibiotic therapy should be instituted in antibody-positive patients. Aminoacidopathies, leukodystrophies, urea-cycle abnormalities, and mitochondrial encephalomyopathies may produce ataxia, and some dietary or metabolic therapies are available for these disorders. The deleterious effects of phenytoin and alcohol on the cerebellum are well known, and these exposures should be avoided in patients with ataxia of any cause.

There is no proven therapy for any of the autosomal dominant ataxias (SCA1 to SCA36). There is preliminary evidence that idebenone, a free-radical scavenger, can improve myocardial hypertrophy in patients with classic Friedreich's ataxia; there is no current evidence, however, that it improves neurologic function. A small preliminary study in a mixed population of patients with different inherited ataxias raised the possibility that the glutamate antagonist riluzole may offer modest benefit. Iron chelators and antioxidant drugs are potentially harmful in Friedreich's patients because they may increase heart muscle injury. Acetazolamide can reduce the duration of symptoms of episodic ataxia. At present, identification of an at-risk person's genotype, together with appropriate family and genetic counseling, can reduce the incidence of these cerebellar syndromes in future generations (**Chap. 84**).

GENETIC DIAGNOSTIC LABORATORIES

1. Baylor College of Medicine; Houston, Texas, 1-713-798-6522
<http://www.bcm.edu/genetics/index.cfm?pmid=21387>
2. GeneDx
<http://www.genedx.com>
3. Transgenomic, 1-877-274-9432
<http://www.transgenomic.com/labs/neurology>

GLOBAL FEATURES

Ataxias with autosomal dominant, autosomal recessive, X-linked, or mitochondrial forms of inheritance are present on a worldwide basis. Machado-Joseph disease (SCA3) (autosomal dominant) and Friedreich's ataxia (autosomal recessive) are the most common types in most populations. Genetic markers are now commercially available to precisely identify the genetic mutation for correct diagnosis and also for family planning. Early detection of asymptomatic preclinical disease can reduce or eliminate the inherited form of ataxia in some families on a global, worldwide basis.

Ataxias with autosomal dominant, autosomal recessive, X-linked, or mitochondrial forms of inheritance are present on a worldwide

basis. Machado-Joseph disease (SCA3) (autosomal dominant) and Friedreich's ataxia (autosomal recessive) are the most common types in most populations. Mutation markers are now commercially available to identify carriers at risk in their families, which allows for precise identification of the genetic mutation for correct diagnosis and also for family planning. Identification of positive mutation carriers with family planning has allowed for early detection of asymptomatic pre-clinical disease to reduce or eliminate the inherited form of ataxia in specific families on a global, worldwide basis.

TABLE 451e-1 CLASSIFICATION OF THE SPINOCEREBELLAR ATAXIAS (SCA)

Name	Locus	Phenotype
SCA1 (autosomal dominant type 1)	6p22-p23 with CAG repeats (exonic); leucine-rich acidic nuclear protein (LANP), region-specific interaction protein Ataxin-1	Ataxia with ophthalmoparesis, pyramidal and extrapyramidal findings; genetic testing is available; 6% of all autosomal dominant (AD) cerebellar ataxia
SCA2 (autosomal dominant type 2)	12q23-q24.1 with CAG repeats (exonic) Ataxin-2	Ataxia with slow saccades and minimal pyramidal and extrapyramidal findings; genetic testing available; 13% of all AD cerebellar ataxia
Machado-Joseph disease/ SCA3 (autosomal dominant type 3)	14q24.3-q32 with CAG repeats (exonic); codes for ubiquitin protease (inactive with polyglutamine expansion); altered turnover of cellular proteins due to proteasome dysfunction MJD-ataxin-3	Ataxia with ophthalmoparesis and variable pyramidal, extrapyramidal, and amyotrophic signs; dementia (mild); 23% of all AD cerebellar ataxia; genetic testing available
SCA4 (autosomal dominant type 4)	16q22.1-ter; pleckstrin homology domain-containing protein, family G, member 4 (PLEKHG4; puratrophin-1: Purkinje cell atrophy associated protein-1, including spectrin repeat and the guanine-nucleotide exchange factor, GEF for Rho GTPases)	Ataxia with normal eye movements, sensory axonal neuropathy, and pyramidal signs; genetic testing available
SCA5 (autosomal dominant type 5)	11p12-q12; β -III spectrin mutations; (SPTBN2); stabilizes glutamate transporter EAAT4; descendants of President Abraham Lincoln	Ataxia and dysarthria; genetic testing available
SCA6 (autosomal dominant type 6)	19p13.2 with CAG repeats in α_{1v} -voltage-dependent calcium channel gene (exonic); CACNA1A protein, P/Q type calcium channel subunit	Ataxia and dysarthria, nystagmus, mild proprioceptive sensory loss; genetic testing available
SCA7 (autosomal dominant type 7)	3p14.1-p21.1 with CAG repeats (exonic); ataxin-7; subunit of GCN5, histone acetyltransferase-containing complexes; ataxin-7 binding protein; Cbl-associated protein (CAP; SH3D5)	Ophthalmoparesis, visual loss, ataxia, dysarthria, extensor plantar response, pigmentary retinal degeneration; genetic testing available
SCA8 (autosomal dominant type 8)	13q21 with CTG repeats; noncoding; 3' untranslated region of transcribed RNA; KLHL1AS	Gait ataxia, dysarthria, nystagmus, leg spasticity, and reduced vibratory sensation; genetic testing available
SCA10 (autosomal dominant type 10)	22q13; pentanucleotide repeat ATTCT repeat; noncoding, intron 9	Gait ataxia, dysarthria, nystagmus; partial complex and generalized motor seizures; polyneuropathy; genetic testing available
SCA11 (autosomal dominant type 11)	15q14-q21.3 by linkage	Slowly progressive gait and extremity ataxia, dysarthria, vertical nystagmus, hyperreflexia
SCA12 (autosomal dominant type 12)	5q31-q33 by linkage; CAG repeat; protein phosphatase 2A, regulatory subunit B, (PPP2R2B); protein PP2A, serine/threonine phosphatase	Tremor, decreased movement, increased reflexes, dystonia, ataxia, dysautonomia, dementia, dysarthria; genetic testing available
SCA13 (autosomal dominant type 13)	19q13.3-q14.4; potassium channel voltage-gated; KCNC3	Ataxia, legs > arms; dysarthria, horizontal nystagmus; delayed motor development; mental developmental delay; tendon reflexes increased; MRI: cerebellar and pontine atrophy; genetic testing available
SCA14 (autosomal dominant type 14)	19q-13.4; protein kinase Cy (PRKCG), missense mutations including in-frame deletion and a splice site mutation among others; serine/threonine kinase	Gait ataxia; leg > arm ataxia; dysarthria; pure ataxia with later onset; myoclonus; tremor of head and extremities; increased deep tendon reflexes at ankles; occasional dystonia and sensory neuropathy; genetic testing available
SCA15 (autosomal dominant type 15)	3p24.2-3pter; inositol 1,4,5- triphosphate receptor type 1 (ITPR1)	Gait and extremity ataxia, dysarthria; nystagmus; MRI: superior vermis atrophy; sparing of hemispheres and tonsils
SCA16 (autosomal dominant type 16)	8q22.1-24.1	Pure cerebellar ataxia and head tremor, gait ataxia, and dysarthria; horizontal gaze-evoked nystagmus; MRI: cerebellar atrophy; no brainstem changes
SCA17 (autosomal dominant type 17)	6q27; CAG expansion in the TATA-binding protein (<i>TBP</i>) gene	Gait ataxia, dementia, parkinsonism, dystonia, chorea, seizures; hyperreflexia; dysarthria and dysphagia; MRI shows cerebral and cerebellar atrophy; genetic testing available
SCA18 (autosomal dominant type 18)	7q22-q32	Ataxia; motor/sensory neuropathy; head tremor; dysarthria; extensor plantar responses in some patients; sensory axonal neuropathy; EMG denervation; MRI: cerebellar atrophy
SCA19 (autosomal dominant type 19)	1p21-q21; KCND3; missense mutations; T352P; M373I; S390N; allelic with SCA22; overlaps with the locus of SCA22	Ataxia, tremor, cognitive impairment, myoclonus; MRI: atrophy of cerebellum
SCA20 (autosomal dominant)	11p13-q11; 260 kb duplication	Dysarthria; gait ataxia; ocular gaze-evoked saccades; palatal tremor; dentate calcification on CT; MRI: cerebral atrophy

(Continued)

TABLE 451e-1 CLASSIFICATION OF THE SPINOCEREBELLAR ATAXIAS (SCA) (CONTINUED)

Name	Locus	Phenotype
SCA21 (autosomal dominant)	7p21.3-p15.1	Ataxia, dysarthria, extrapyramidal features of akinesia, rigidity, tremor, cognitive defect; reduced deep tendon reflexes; MRI: cerebellar atrophy, normal basal ganglia and brainstem
SCA22 (autosomal dominant)	1p21-q23; deletion (in frame); V338E; G345V; T377M; allelic with SCA19; KCND3; Kv4.3 channels	Pure cerebellar ataxia; dysarthria; dysphagia; nystagmus; MRI: cerebellar atrophy
SCA23 (autosomal dominant)	20p13-12.3; prodynorphin (PDYN protein); missense R138S; L211S; R212W; R215C	Gait ataxia; dysarthria; extremity ataxia; ocular nystagmus, dysmetria; leg vibration loss; extensor plantar responses; MRI: cerebellar atrophy
SCA25 (autosomal dominant)	2p15-p21	Ataxia, nystagmus; vibratory loss in the feet; pain loss in some; abdominal pain; nausea and vomiting may be prominent; absent ankle reflexes; sensory nerve action potentials are absent; MRI: cerebellar atrophy, normal brainstem
SCA26 (autosomal dominant)	19p13.3	Gait ataxia; extremity ataxia; dysarthria; nystagmus; MRI: cerebellar atrophy
SCA27 (autosomal dominant)	13q34; fibroblast growth factor 14 protein; mutation F145S; produces reduced protein stability	Tremor in extremities and head and orofacial dyskinesia; ataxia of arms > legs, gait ataxia; dysarthria; nystagmus; psychiatric symptoms; cognitive defect; MRI: cerebellar atrophy; genetic testing available
SCA28 (autosomal dominant)	18p11.22-q11.2; ATPase family gene 3- like 2 (AFG3L2 protein) mutations: N432T; S674L; E691K; A694E; R702Q	Extremity and gait ataxia; dysarthria; nystagmus; ophthalmoparesis; leg hyperreflexia and extensor plantar responses; MRI: cerebellar atrophy
SCA30 (autosomal dominant)	4q34.3-q35.1; candidate gene <i>ODZ3</i>	Candidate gene <i>ODZ3</i> ; gait ataxia, dysarthria, saccades; nystagmus, brisk tendon reflexes in legs; MRI: cerebellar atrophy
SCA31 (autosomal dominant)	16q22.1; associated with NEDD4 (BEAN)	Pentanucleotide (TGGAA) _n repeat insertions; previously called SCA4; gait ataxia; limb dysmetria; MRI: cerebellar atrophy
SCA32 (autosomal dominant)	7q32-q33	Ataxia, azoospermia, mental retardation; absent germ cells on testicular biopsy
SCA35 (autosomal dominant)	20p13; TGM6 protein; transglutaminase 6	Ataxia; ocular dysmetria; upper motor neuron signs; extensor plantars; onset fifth decade
SCA36 (autosomal dominant)	20p13; large intronic expansion of GGCCTG (1500–2500); also phe265leu mutation; RNA gain of function; microRNA; MIR 1292 suppression	Ataxia; onset fifth to sixth decades; motor neuron disorder; grouped atrophy (muscle biopsy) fasciculations; increased reflexes; flexor plantars
Prion disease (autosomal dominant)	20p13; pro102leu; ala 117 val mutations; proteinase k resistant form PrP27-30 accumulates in brain; eponym: Gerstmann-Straüssler-Scheinker disease Glu200Lys mutation; increased octapeptide repeats; eponym: Creutzfeldt-Jakob disease	Ataxia; dementia third to seventh decades Ataxia; dementia; rigidity
Multiple hamartoma syndrome (autosomal dominant)	10q23.31; phosphatase and tensin homolog (PTEN); Cowden's; Lhermitte-Duclos syndrome	Skin hamartomas; ataxia; mental retardation; increased intracranial pressure; epilepsy
Cerebellar ataxia, deafness, and narcolepsy (autosomal dominant)	19p13.2; exon 21; missense ala570val; val606phe mutations	Ataxia; deafness; narcolepsy cataplexy; REM sleep disorder
Cerebellar ataxia (nonprogressive) mental retardation (autosomal dominant)	1p36.31-p36.23	Ataxia, mental retardation
Familial dementia with amyloid angiopathy and spastic ataxia (autosomal dominant)	13q14.2; integral membrane protein 2B (ITM2B)	Ataxia; dementia; amyloid angiopathy
Dentatorubropallidolusian atrophy (autosomal dominant)	12p13.31 with CAG repeats (exonic) Atrophin 1	Ataxia, choreoathetosis, dystonia, seizures, myoclonus, dementia; genetic testing available
Friedreich's ataxia (autosomal recessive)	9q13-q21.1 with intronic GAA repeats, in intron at end of exon 1 Frataxin defective; abnormal regulation of mitochondrial iron metabolism; iron accumulates in mitochondria in yeast mutants	Ataxia, areflexia, extensor plantar responses, position sense deficits, cardiomyopathy, diabetes mellitus, scoliosis, foot deformities; optic atrophy; late-onset form, as late as 50 years with preserved deep tendon reflexes, slower progression, reduced skeletal deformities, associated with an intermediate number of GAA repeats and missense mutations in one allele of frataxin; genetic testing available
Vitamin E deficiency syndrome (autosomal recessive)	8q13.1-q13.3 (α-TTP deficiency)	Same as phenotype that maps to 9q but associated with vitamin E deficiency; genetic testing available
Sensory ataxic neuropathy and ophthalmoparesis (SANDO) with dysarthria (autosomal recessive)	15q25; mutations in DNA polymerase-gamma (<i>POLG</i>) gene that leads to mtDNA deletions	Young adult-onset ataxia, sensory neuropathy, ophthalmoparesis, hearing loss, gastric symptoms; a variant of progressive external ophthalmoplegia; MRI: cerebellar and thalamic abnormalities; mildly increased lactate and creatine kinase
Von Hippel-Lindau syndrome (autosomal dominant)	3p26-p25	Cerebellar hemangioblastoma; pheochromocytoma

TABLE 451e-1 CLASSIFICATION OF THE SPINOCEREBELLAR ATAXIAS (SCA) (CONTINUED)

Name	Locus	Phenotype
Baltic myoclonus (Unverricht-Lundborg) (recessive)	21q22.3; cystatin B; extra repeats of 12–base pair tandem repeats	Myoclonus epilepsy; late-onset ataxia; responds to valproic acid, clonazepam; phenobarbital
Marinesco-Sjögren syndrome (recessive)	5q31; SIL 1 protein, nucleotide exchange factor for the heat-shock protein 70 (HSP70); chaperone HSPA5; homozygous 4-nucleotide duplication in exon 6; also compound heterozygote	Ataxia, dysarthria; nystagmus; retarded motor and mental maturation; rhabdomyolysis after viral illness; weakness; hypotonia; areflexia; cataracts in childhood; short stature; kyphoscoliosis; contractures; hypogonadism
Autosomal recessive spastic ataxia of Charlevoix-Saguenay (ARSACS)	Chromosome 13q12; SACS gene; loss of saccin peptide activity	Childhood onset of ataxia, spasticity, dysarthria, distal muscle wasting, foot deformity, retinal striations, mitral valve prolapse
Kearns-Sayre syndrome (sporadic)	mtDNA deletion and duplication mutations	Ptosis, ophthalmoplegia, pigmentary retinal degeneration, cardiomyopathy, diabetes mellitus, deafness, heart block, increased CSF protein, ataxia
Myoclonic epilepsy and ragged red fiber syndrome (MERRF) (maternal inheritance)	Mutation in mtDNA of the tRNA ^{lys} at 8344; also mutation at 8356	Myoclonic epilepsy, ragged red fiber myopathy, ataxia
Mitochondrial encephalopathy, lactic acidosis, and stroke syndrome (MELAS) (maternal inheritance)	tRNA ^{leu} mutation at 3243; also at 3271 and 3252	Headache, stroke, lactic acidosis, ataxia
Neuropathy; ataxia; retinitis pigmentosa (NARP)	ATPase6 (Complex 5); mtDNA point mutation at 8993	Neuropathy; ataxia; retinitis pigmentosa; dementia; seizures
Episodic ataxia, type 1 (EA-1) (autosomal dominant)	12p13; potassium voltage-gated channel gene, <i>KCNA1</i> ; Phe249Leu mutation; variable syndrome	Episodic ataxia for minutes; provoked by startle or exercise; with facial and hand myokymia; cerebellar signs are not progressive; choreoathetotic movements; responds to phenytoin; genetic testing available
Episodic ataxia, type 2 (EA-2) (autosomal dominant)	19p-13 (<i>CACNA1A</i>) (allelic with SCA6) (α_{1A} -voltage-dependent calcium channel subunit); point mutations or small deletions; allelic with SCA6 and familial hemiplegic migraine	Episodic ataxia for days; provoked by stress, fatigue; with down-gaze nystagmus; vertigo; vomiting; headache; cerebellar atrophy results; progressive cerebellar signs; responds to acetazolamide; genetic testing available
Episodic ataxia, type 3 (autosomal dominant)	1q42	Episodic ataxia; 1 min to over 6 h; induced by movement; vertigo and tinnitus; headache; responds to acetazolamide
Episodic ataxia, type 4 (autosomal dominant)	Not mapped	Episodic ataxia; vertigo; diplopia; ocular slow pursuit defect; no response to acetazolamide
Episodic ataxia, type 5 (autosomal dominant)	2q22-q23; <i>CACNB4</i> protein	Episodic ataxia; hours to weeks; seizures
Episodic ataxia type 6 with seizures, migraine, and alternating hemiplegia (autosomal dominant)	<i>SLC1A3</i> ; 5p13; <i>EAAT1</i> protein; missense mutations; glial glutamate transporter (<i>GLAST</i>); 1047 C to G; proline to arginine	Ataxia, duration 2–4 days; episodic hypotonia; delayed motor milestones; seizures; migraine; alternating hemiplegia; mild truncal ataxia; coma; febrile illness as a trigger; MRI: cerebellar atrophy
Episodic ataxia, type 7 (autosomal dominant)	19q13	Episodic ataxia; vertigo, weakness; less than 24 h
Episodic ataxia with paroxysmal choreoathetosis and spasticity (dystonia-9) (<i>DYT9</i> ; CSE) (autosomal dominant)	1p	Ataxia; involuntary movements; dystonia; headache; spastic paraplegia; responds on occasion to acetazolamide
Fragile X tremor/ataxia syndrome (FXTAS) X-linked dominant	Xq27.3; CGG premutation expansion in <i>FMR1</i> gene; expansions of 55–200 repeats in 5' UTR of the <i>FMR-1</i> mRNA; presumed dominant toxic RNA effect	Late-onset ataxia with tremor, cognitive impairment, occasional parkinsonism; males typically affected, although affected females also reported; syndrome is of high concern if affected male has grandson with mental retardation (fragile X syndrome); MRI shows increased T2 signal in middle cerebellar peduncles, cerebellar atrophy, and occasional widespread brain atrophy; genetic testing available
Ataxia telangiectasia (autosomal recessive)	11q22-23; <i>ATM</i> gene for regulation of cell cycle; mitogenic signal transduction and meiotic recombination; elevated serum alpha-fetoprotein level; immunoglobulin deficiency	Telangiectasia, ataxia, dysarthria, pulmonary infections, neoplasms of lymphatic system; IgA and IgG deficiencies; diabetes mellitus, breast cancer; genetic testing available; chorea; dystonia
Early-onset cerebellar ataxia with retained deep tendon reflexes (autosomal recessive)	13q11-12	Ataxia; neuropathy; preserved deep tendon reflexes; impaired cognitive and visuospatial functions; MRI, cerebellar atrophy
Ataxia with oculomotor apraxia (AOA1) (autosomal recessive)	9p21; protein is member of histidine triad superfamily, role in DNA repair; elevation of serum LDL cholesterol and low serum albumin level; <i>APTX</i> , aprataxin	Ataxia; dysarthria; limb dysmetria; dystonia; oculomotor apraxia; optic atrophy; motor neuropathy; late sensory loss (vibration); genetic testing available
Ataxia with oculomotor apraxia 2 (AOA2) (autosomal recessive)	9q34; senataxin protein, involved in RNA maturation and termination; helicase superfamily 1; elevated serum alpha-fetoprotein level; <i>SETX</i> , senataxin	Gait ataxia; choreoathetosis; dystonia; oculomotor apraxia; neuropathy, vibration loss, position sense loss, and mild light touch loss; absent leg deep tendon reflexes; extensor plantar response; genetic testing available

(Continued)

TABLE 451e-1 CLASSIFICATION OF THE SPINOCEREBELLAR ATAXIAS (SCA) (CONTINUED)

Name	Locus	Phenotype
Cerebellar ataxia with muscle coenzyme Q10 deficiency (autosomal recessive)	9p13	Ataxia; hypotonia; seizures; mental retardation; increased deep tendon reflexes; extensor plantar responses; coenzyme Q10 levels reduced with about 25% of patients with a block in transfer of electrons to complex 3; may respond to coenzyme 10
Refsum's disease (autosomal recessive)	10pter; elevated serum phytanic acid level; phytanoyl-COH hydroxylase and PEX7	Retinitis pigmentosa; ataxia; sensorineural deafness; demyelinating neuropathy
Cerebrotendinous xanthomatosis (autosomal recessive)	2p33; elevated cholesterol level; CYP27; sterol 27 hydroxylase	Spastic ataxia; mental retardation; dementia; tendon xanthomas; diarrhea; cataracts
Joubert's syndrome (autosomal recessive)	9q34.3	Ataxia; ptosis; mental retardation; oculomotor apraxia; nystagmus; retinopathy; rhythmic tongue protrusion; episodic hyperpnea or apnea; dimples at wrists and elbows; telecanthus; micrognathia
Sideroblastic anemia and spinocerebellar ataxia (X-linked recessive)	Xq13; ATP-binding cassette 7 (ABC7; ABC7) transporter; mitochondrial inner membrane; iron homeostasis; export from matrix to the intermembrane space	Ataxia; elevated free erythrocyte protoporphyrin levels; ring sideroblasts in bone marrow; heterozygous females may have mild anemia but not ataxia
Infantile-onset spinocerebellar ataxia of Nikali et al (autosomal recessive)	10q23.3-q24.1; <i>twinkle</i> protein (gene); homozygous for Tyr508Cys missense mutations	Infantile ataxia, sensory neuropathy; athetosis, hearing deficit, reduced deep tendon reflexes; ophthalmoplegia, optic atrophy; seizures; primary hypogonadism in females
Hypoceruloplasminemia with ataxia and dysarthria (autosomal recessive)	Ceruloplasmin gene; 3q23-q25 (trp 858 ter)	Gait ataxia and dysarthria; hyperreflexia; cerebellar atrophy by MRI; iron deposition in cerebellum, basal ganglia, thalamus, and liver; onset in the fourth decade
Spinocerebellar ataxia with neuropathy (SCAN1) (autosomal recessive)	Tyrosyl-DNA phosphodiesterase-1 (TDP-1) 14q31-q32	Onset in second decade; gait ataxia, dysarthria, seizures, cerebellar vermis atrophy on MRI, dysmetria
Cerebellar ataxia type 1 (autosomal recessive)	6p25 SCAR8; SYNE1; spectrin repeats-nuclear envelope 1	Pure ataxia
Cerebellar ataxia type 2 (autosomal recessive)	1q42; ADCK3 (CABC1); aarf-domain containing kinase 3; elevation of serum lactate and decreased coenzyme Q10 level	Ataxia; mental retardation; myoclonus; epilepsy; exercise intolerance; stroke or transient ischemic-like episodes
Niemann-Pick type C disease	18q11; NPC1; NPCH1 and 2; skin biopsy (filipin staining)	Ataxia; vertical supranuclear ophthalmoplegia; splenomegaly; dystonia; impaired cognition

Abbreviations: CSF, cerebrospinal fluid; CT, computed tomography; EMG, electromyogram; LDL, low-density lipoprotein; MRI, magnetic resonance imaging; REM, rapid eye movement; UTR, untranslated region.

TABLE 452-1 ETIOLOGY OF MOTOR NEURON DISORDERS

Diagnostic Category	Investigation
Structural lesions	MRI scan of head (including foramen magnum and cervical spine)
Parasagittal or foramen magnum tumors	
Cervical spondylosis	
Chiari malformation of syrinx	
Spinal cord arteriovenous malformation	
Infections	CSF exam, culture
Bacterial—tetanus, Lyme	Lyme titer
Viral—poliomyelitis, herpes zoster	Anti-viral antibody
Retroviral—myelopathy	HTLV-1 titers
Intoxications, physical agents	24-h urine for heavy metals
Toxins—lead, aluminum, others	Serum lead level
Drugs—strychnine, phenytoin	
Electric short, x-irradiation	
Immunologic mechanisms	Complete blood count ^a
Plasma cell dyscrasias	Sedimentation rate ^a
Autoimmune polyradiculopathy	Total protein ^a
Motor neuropathy with conduction block	Anti-GM1 antibodies ^a
Paraneoplastic	Anti-Hu antibody
Paracarcinomatous	MRI scan, bone marrow biopsy
Metabolic	Fasting blood sugar ^a
Hypoglycemia	Routine chemistries including calcium ^a
Hyperparathyroidism	PTH
Hyperthyroidism	Thyroid function ^a
Deficiency of folate, vitamin B ₁₂ , vitamin E	Vitamin B ₁₂ , vitamin E, folate ^a
Malabsorption	Serum zinc, copper ^a
Deficiency of copper, zinc	24-h stool fat, carotene, prothrombin time
Mitochondrial dysfunction	Fasting lactate, pyruvate, ammonia
	Consider mtDNA
Hyperlipidemia	Lipid electrophoresis
Hyperglycinuria	Urine and serum amino acids
	CSF amino acids
Hereditary disorders	WBC DNA for mutational analysis
Superoxide dismutase	
TDP43	
FUS/TLS	
Androgen receptor defect (Kennedy's disease)	
Hexosaminidase deficiency	
Infantile a-glucosidase deficiency (Pompe's disease)	

^aShould be obtained in all cases.

Abbreviations: CSF, cerebrospinal fluid; FUS/TLS, fused in sarcoma/translocated in liposarcoma; HTLV-1, human T-cell lymphotropic virus; MRI, magnetic resonance imaging; PTH, parathyroid; WBC, white blood cell.

452 Amyotrophic Lateral Sclerosis and Other Motor Neuron Diseases

Robert H. Brown, Jr.

AMYOTROPHIC LATERAL SCLEROSIS

Amyotrophic lateral sclerosis (ALS) is the most common form of progressive motor neuron disease. It is a prime example of a neurodegenerative disease and is arguably the most devastating of the neurodegenerative disorders.

PATHOLOGY

The pathologic hallmark of motor neuron degenerative disorders is death of lower motor neurons (consisting of anterior horn cells in the spinal cord and their brainstem homologues innervating bulbar muscles) and upper, or corticospinal, motor neurons (originating in layer five of the motor cortex and descending via the pyramidal tract to synapse with lower motor neurons, either directly or indirectly via interneurons) (Chap. 30). Although at its onset ALS may involve selective loss of function of only upper or lower motor neurons, it ultimately causes progressive loss of both categories of motor neurons. Indeed, in the absence of clear involvement of both motor neuron types, the diagnosis of ALS is questionable. In a subset of cases, ALS arises concurrently with frontotemporal dementia (Chap. 448); in these instances, there is degeneration of frontotemporal cortical neurons and corresponding cortical atrophy.

Other motor neuron diseases involve only particular subsets of motor neurons (Tables 452-1 and 452-2). Thus, in bulbar palsy and spinal muscular atrophy (SMA; also called *progressive muscular atrophy*),

the lower motor neurons of brainstem and spinal cord, respectively, are most severely involved. By contrast, pseudobulbar palsy, primary lateral sclerosis (PLS), and familial spastic paraplegia (FSP) affect only upper motor neurons innervating the brainstem and spinal cord.

In each of these diseases, the affected motor neurons undergo shrinkage, often with accumulation of the pigmented lipid (lipofuscin) that normally develops in these cells with advancing age. In ALS, the motor neuron cytoskeleton is typically affected early in the illness. Focal enlargements are frequent in proximal motor axons; ultrastructurally, these "spheroids" are composed of accumulations of neurofilaments and other proteins. Commonly in both sporadic and familial

Chronic	Entity
Upper and lower motor neuron	Amyotrophic lateral sclerosis
Predominantly upper motor neuron	Primary lateral sclerosis
Predominantly lower motor neuron	Multifocal motor neuropathy with conduction block
	Motor neuropathy with paraproteinemia or cancer
	Motor predominant peripheral neuropathies
Other	
	Associated with other neurodegenerative disorders
	Secondary motor neuron disorders (see Table 452-1)
Acute	
	Poliomyelitis
	Herpes zoster
	Coxsackie virus
	West Nile virus

ALS, the affected neurons demonstrate ubiquitin-positive aggregates, typically associated with the protein TDP43 (see below). Also seen is proliferation of astroglia and microglia, the inevitable accompaniment of all degenerative processes in the central nervous system (CNS).

The death of the peripheral motor neurons in the brainstem and spinal cord leads to denervation and consequent atrophy of the corresponding muscle fibers. Histochemical and electrophysiologic evidence indicates that in the early phases of the illness denervated muscle can be reinnervated by sprouting of nearby distal motor nerve terminals, although reinnervation in this disease is considerably less extensive than in most other disorders affecting motor neurons (e.g., poliomyelitis, peripheral neuropathy). As denervation progresses, muscle atrophy is readily recognized in muscle biopsies and on clinical examination. This is the basis for the term *amyotrophy*. The loss of cortical motor neurons results in thinning of the corticospinal tracts that travel via the internal capsule (Fig. 452-1) and brainstem to the lateral and anterior white matter columns of the spinal cord. The loss of fibers in the lateral columns and resulting fibrillary gliosis impart a particular firmness (*lateral sclerosis*). A remarkable feature of the disease is the selectivity of neuronal cell death. By light microscopy, the entire sensory apparatus, the regulatory mechanisms for the control and coordination of movement, remains intact. Except in cases of frontotemporal dementia, the components of the brain required for cognitive processing are also preserved. However, immunostaining indicates that neurons bearing ubiquitin, a marker for degeneration, are also detected in nonmotor systems. Moreover, studies of glucose metabolism in the illness also indicate that there is neuronal dysfunction outside of the motor system. Within the motor system, there is some selectivity of involvement. Thus, motor neurons required for ocular motility remain unaffected, as do the parasympathetic neurons in the sacral spinal cord (the nucleus of Onufrowicz, or Onuf) that innervate the sphincters of the bowel and bladder.

CLINICAL MANIFESTATIONS

The manifestations of ALS are somewhat variable depending on whether corticospinal neurons or lower motor neurons in the brainstem and spinal cord are more prominently involved. With lower motor neuron dysfunction and early denervation, typically the first evidence of the disease is insidiously developing asymmetric weakness, usually first evident distally in one of the limbs. A detailed history often discloses recent development of cramping with volitional movements, typically in the early hours of the morning (e.g., while stretching in bed). Weakness caused by denervation is associated with progressive wasting and atrophy of muscles and, particularly early in the illness, spontaneous twitching of motor units, or fasciculations. In the hands, a

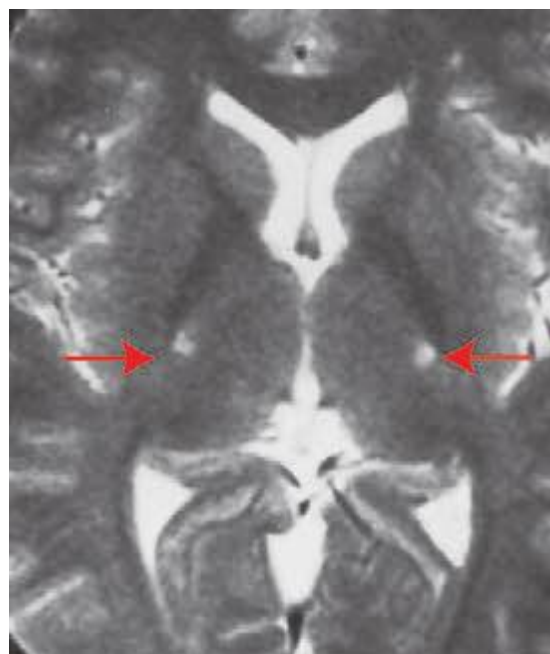


FIGURE 452-1 Amyotrophic lateral sclerosis. Axial T2-weighted magnetic resonance imaging (MRI) scan through the lateral ventricles of the brain reveals abnormal high signal intensity within the corticospinal tracts (arrows). This MRI feature represents an increase in water content in myelin tracts undergoing Wallerian degeneration secondary to cortical motor neuronal loss. This finding is commonly present in ALS, but can also be seen in AIDS-related encephalopathy, infarction, or other disease processes that produce corticospinal neuronal loss in a symmetric fashion.

preponderance of extensor over flexor weakness is common. When the initial denervation involves bulbar rather than limb muscles, the problem at onset is difficulty with chewing, swallowing, and movements of the face and tongue. Early involvement of the muscles of respiration may lead to death before the disease is far advanced elsewhere. With prominent corticospinal involvement, there is hyperactivity of the muscle-stretch reflexes (tendon jerks) and, often, spastic resistance to passive movements of the affected limbs. Patients with significant reflex hyperactivity complain of muscle stiffness often out of proportion to weakness. Degeneration of the corticobulbar projections innervating the brainstem results in dysarthria and exaggeration of the motor expressions of emotion. The latter leads to involuntary excess in weeping or laughing (pseudobulbar affect).

Virtually any muscle group may be the first to show signs of disease, but, as time passes, more and more muscles become involved until ultimately the disorder takes on a symmetric distribution in all regions. It is characteristic of ALS that, regardless of whether the initial disease involves upper or lower motor neurons, both will eventually be implicated. Even in the late stages of the illness, sensory, bowel and bladder, and cognitive functions are preserved. Even when there is severe brainstem disease, ocular motility is spared until the very late stages of the illness. As noted, in some cases (particularly those that are familial), ALS develops concurrently with frontotemporal dementia, characterized by early behavioral abnormalities with prominent behavioral features indicative of frontal lobe dysfunction.

A committee of the World Federation of Neurology has established diagnostic guidelines for ALS. Essential for the diagnosis is simultaneous upper and lower motor neuron involvement with progressive weakness and the exclusion of all alternative diagnoses. The disorder is ranked as “definite” ALS when three or four of the following are involved: bulbar, cervical, thoracic, and lumbosacral motor neurons. When two sites are involved, the diagnosis is “probable,” and when only one site is implicated, the diagnosis is “possible.” An exception is made for those who have progressive upper and lower motor neuron

signs at only one site and a mutation in the gene encoding superoxide dismutase (SOD1; see below).

EPIDEMIOLOGY

The illness is relentlessly progressive, leading to death from respiratory paralysis; the median survival is from 3–5 years. There are very rare reports of stabilization or even regression of ALS. In most societies, there is an incidence of 1–3 per 100,000 and a prevalence of 3–5 per 100,000. It is striking that about 1 in 1000 adult deaths in North America and Western Europe (and probably elsewhere) are due to ALS; this finding predicts that some 300,000 individuals now alive in the United States will die of ALS. Several endemic foci of higher prevalence exist in the western Pacific (e.g., in specific regions of Guam or Papua New Guinea). In the United States and Europe, males are somewhat more frequently affected than females. Epidemiologic studies have incriminated risk factors for this disease including exposure to pesticides and insecticides, smoking, and, in one report, service in the military. Although ALS is overwhelmingly a sporadic disorder, some 5–10% of cases are inherited as an autosomal dominant trait.

FAMILIAL ALS

Several forms of selective motor neuron disease are inheritable (Table 452-3). Familial ALS (FALS) involves both corticospinal and lower motor neurons. Apart from its inheritance as an autosomal dominant trait, it is clinically indistinguishable from sporadic ALS. Genetic studies have identified mutations in multiple genes, including those encoding the protein C9orf72 (open reading frame 72 on chromosome 9), cytosolic enzyme SOD1 (superoxide dismutase), the RNA binding proteins TDP43 (encoded by the TAR DNA binding protein gene), and FUS/TLS (fused in sarcoma/translocated in liposarcoma), as the most common causes of FALS. Mutations in C9orf72 account for ~45–50% of FALS and perhaps 4–5% of sporadic ALS cases. Mutations in SOD1 explain another 20% of cases of FALS, whereas TDP43 and FUS/TLS each represent about 5% of familial cases. It has recently been reported that ~1–2% of cases are caused by mutations in genes encoding the proteins optineuron and profilin-1 as well.

Rare mutations in other genes are also clearly implicated in ALS-like diseases. Thus, a familial, dominantly inherited motor disorder that in some individuals closely mimics the ALS phenotype arises from mutations in a gene that encodes a vesicle-binding protein. A predominantly lower motor neuron disease with early hoarseness due to laryngeal dysfunction has been ascribed to mutations in the gene encoding the cellular accessory motor protein dynactin. Mutations in *senataxin*, a helicase, cause an early adult-onset, slowly evolving ALS variant. Kennedy's syndrome is an X-linked, adult-onset disorder that may mimic ALS, as described below.

Genetic analyses are also beginning to illuminate the pathogenesis of some childhood-onset motor neuron diseases. For example, a slowly disabling degenerative, predominantly upper motor neuron disease that starts in the first decade is caused by mutations in a gene that expresses a novel signaling molecule with properties of a guanine-exchange factor, termed *alsin*.

DIFFERENTIAL DIAGNOSIS

Because ALS is currently untreatable, it is imperative that potentially remediable causes of motor neuron dysfunction be excluded (Table 452-1). This is particularly true in cases that are atypical by virtue of (1) restriction to either upper or lower motor neurons, (2) involvement of neurons other than motor neurons, and (3) evidence of motor neuronal conduction block on electrophysiologic testing. Compression of the cervical spinal cord or cervicomedullary junction from tumors in the cervical regions or at the foramen magnum or from cervical spondylosis with osteophytes projecting into the vertebral canal can produce weakness, wasting, and fasciculations in the upper limbs and spasticity in the legs, closely resembling ALS. The absence of cranial nerve involvement may be helpful in differentiation, although some foramen magnum lesions may compress the twelfth cranial (hypoglossal) nerve, with resulting paralysis of the tongue. Absence of

pain or of sensory changes, normal bowel and bladder function, normal roentgenographic studies of the spine, and normal cerebrospinal fluid (CSF) all favor ALS. Where doubt exists, magnetic resonance imaging (MRI) scans and contrast myelography should be performed to visualize the cervical spinal cord.

Another important entity in the differential diagnosis of ALS is *multifocal motor neuropathy with conduction block* (MMCB), discussed below. A diffuse, lower motor axonal neuropathy mimicking ALS sometimes evolves in association with hematopoietic disorders such as lymphoma or multiple myeloma. In this clinical setting, the presence of an M-component in serum should prompt consideration of a bone marrow biopsy. Lyme disease (Chap. 210) may also cause an axonal, lower motor neuropathy, although typically with intense proximal limb pain and a CSF pleocytosis.

Other treatable disorders that occasionally mimic ALS are chronic lead poisoning and thyrotoxicosis. These disorders may be suggested by the patient's social or occupational history or by unusual clinical features. When the family history is positive, disorders involving the genes encoding C9orf72, cytosolic SOD1, TDP43, FUS/TLS, and adult hexosaminidase A or α -glucosidase deficiency must be excluded (Chap. 432e). These are readily identified by appropriate laboratory tests. Benign fasciculations are occasionally a source of concern because on inspection they resemble the fascicular twitchings that accompany motor neuron degeneration. The absence of weakness, atrophy, or denervation phenomena on electrophysiologic examination usually excludes ALS or other serious neurologic disease. Patients who have recovered from poliomyelitis may experience a delayed deterioration of motor neurons that presents clinically with progressive weakness, atrophy, and fasciculations. Its cause is unknown, but it is thought to reflect sublethal prior injury to motor neurons by poliovirus (Chap. 228).

Rarely, ALS develops concurrently with features indicative of more widespread neurodegeneration. Thus, one infrequently encounters otherwise typical ALS patients with a parkinsonian movement disorder or frontotemporal dementia, particularly in instances of C9orf72 mutations, which strongly suggests that the simultaneous occurrence of two disorders is a direct consequence of the gene mutation. As another example, prominent amyotrophy has been described as a dominantly inherited disorder in individuals with bizarre behavior and a movement disorder suggestive of parkinsonism; many such cases have now been ascribed to mutations that alter the expression of tau protein in brain (Chap. 448). In other cases, ALS develops simultaneously with a striking frontotemporal dementia. An ALS-like disorder has also been described in some individuals with chronic traumatic encephalopathy, associated with deposition of TDP43 and neurofibrillary tangles in motor neurons.

PATHOGENESIS

The cause of sporadic ALS is not well defined. Several mechanisms that impair motor neuron viability have been elucidated in mice and rats induced to develop motor neuron disease by SOD1 transgenes with ALS-associated mutations. One may loosely group the genetic causes of ALS into three categories. In one group, the primary problem is inherent instability of the mutant proteins, with subsequent perturbations in protein degradation (SOD1, ubiquilin-1 and -2, p62). In the second, most rapidly growing category, the causative mutant genes perturb RNA processing, transport, and metabolism (C9orf73, TDP43, FUS). In the case of C9orf72, the molecular pathology is an expansion of an intronic hexanucleotide repeat (-GGGGCC-) beyond an upper normal of 30 repeats to hundreds or more repeats. As observed in other intronic repeat disorders such as myotonic dystrophy (Chap. 462e) and spinocerebellar atrophy type 8 (Chap. 450), data suggest that the expanded intronic repeats generate expanded RNA repeats that form intranuclear foci and confer toxicity by sequestering transcription factors or by undergoing noncanonical protein translation across all possible reading frames of the expanded RNA tracts. TDP43 and FUS are multifunctional proteins that bind RNA and DNA and shuttle between the nucleus and the cytoplasm, playing multiple roles in the control of cell proliferation, DNA repair and transcription,

Disease	Locus	Gene	Inheritance	Usual Onset	Gene Function	Unusual Features
I. Upper and Lower Motor Neurons (Familial ALS)						
ALS1	21q	Superoxide dismutase	AD	Adult	Protein antioxidant	
ALS2	2q	Alsin	AR	Juvenile	GEF signaling	Severe corticobulbar, corticospinal features
ALS4	9q	Senataxin	AD	Late juvenile	DNA helicase	Late childhood onset
ALS6	16p	FUS/TLS	AD	Adult	DNA, RNA binding	
ALS8	20q	Vesicle associated protein B	AD	Adult	Vesicular trafficking	
ALS9	14q	Angiogenin	AD	Adult	RNAse, angiogenesis	
ALS10	1q	TDP43	AD	Adult	DNA, RNA binding	
ALS12	10p	Optineurin	AD/AR	Adult	Attenuates NF- κ B	
ALS13	12q	Ataxin-2	AD	Adult	Cytotoxic expanded CAG repeat	
ALS14	9p	Valosin-containing protein	AD	Adult	ATPase	
ALS18	17p	Profilin-1	AD	Adult	Involved in actin polymerization	
ALS19	2q	ErbB4	AD	Adult	Signaling molecule	
ALS20	12q	HNRNPA1	AD	Adult	Heteronuclear RNA binding protein	
ALS21	5q	MTR3	AD	Adult	Nuclear matrix protein	Early vocal/bulbar involvement
ALS	2p	Dynactin	AD	Adult	Axonal transport	
ALS	17q	Paraoxonases 1-3	AD	Adult	Detoxify intoxicants	
ALS	mtDNA	Cytochrome c oxidase		Adult	ATP generation	
ALS	mtDNA	tRNA-isoleucine		Adult	ATP generation	
II. Lower Motor Neurons						
Spinal muscular atrophies	5q	Survival motor neuron	AR	Infancy	RNA metabolism	
GM2-gangliosidosis						
1. Sandhoff's disease	5q	Hexosaminidase B	AR	Childhood	Ganglioside recycling	
2. AB variant	5q	GM2-activator protein	AR	Childhood	Ganglioside recycling	
3. Adult Tay-Sachs disease	15q	Hexosaminidase A	AR	Childhood	Ganglioside recycling	
X-linked spino-bulbar muscular atrophy	Xq	Androgen receptor	XR	Adult	Nuclear signaling	
III. Upper Motor Neuron (Selected FSPs)						
SPG3A	14q	Atlastin	AD	Childhood	GTPase—vesicle recycling	
SPG4	2p	Spastin	AD	Early adulthood	ATPase family—microtubule associate	Some sensory loss
SPG6	15q	NIPA1	AD	Early adulthood	Membrane transporter or receptor	Deleted in Prader-Willi, Angelman's
SPG8	8q	Strumpellin	AD	Early adulthood	Ubiquitous, spectrin-like	
SPG10	12q	Kinesin heavy chain KIF5A	AD	Second–third decade	Motor-associated protein	\pm Peripheral neuropathy, retardation
SPG12	19q	Reticulon 2	AD	Childhood	ER protein, interacts with spastin	
SPG13	2q	Heat shock protein 60	AD	Early adulthood	Chaperone protein	
SPG17	11q	Silver (BSCL2)	AD	Variable	Membrane protein in ER	Amyotrophy hands, feet
SPG31	2p	REEP1	AD	Early	Mitochondrial protein	Rarely, amyotrophy
SPG33	10q	ZFYVE27	AD	Adult	Interacts with spastin	Pes equinus
SPG42	3q	Acetyl-CoA-transporter	AD	Variable	Solute carrier	
SPG72	5q	REEP2	AD	Childhood	ER protein	
SPG5	8q	Cytochrome P450	AR	Variable	Degrades endogenous substances	Sensory loss
SPG7	16q	Paraplegin	AR	Variable	Mitochondrial protein	Rarely, optic atrophy, ataxia
SPG11	15q	Spatacsin	AR	Childhood	Cytosolic, ? membrane-associated	Some sensory loss, thin corpus callosum
SPG15	14q	Spastizin	AR	Childhood	Zinc finger protein	Some amyotrophy, some CNS features

(Continued)

TABLE 452-3 GENETIC MOTOR NEURON DISEASES (CONTINUED)

Disease	Locus	Gene	Inheritance	Usual Onset	Gene Function	Unusual Features
SPG20	13q	Spartin	AR	Childhood	Endosomal trafficking protein	
SPG21	15q	Maspardin	AR	Childhood	Endosomal trafficking protein	
SPG35	16q	Fatty acid 2 hydrolase	AR	Childhood	Membrane protein	Multiple CNS features
SPG39	19p	Neuropathy target esterase	AR	Early childhood	Esterase	
SPG44	1q	Connexin 47	AR	Childhood	Gap junction protein	Possible mild CNS features
SPG46	9p	β -Glucosidase 2	AR	Childhood	Glycoside hydrolase	Thin corpus callosum, mental retardation
SPG2	Xq	Proteolipid protein	XR	Early childhood	Myelin protein	Sometimes multiple CNS features
SPG1	Xq	L1-CAM	XR	Infancy	Cell adhesion molecule	
SPG22	Xq	SLC16A2	XR	Infancy	Monocarboxylic acid transporter	
	Xq	Adrenoleukodystrophy	XR	Early adulthood	ATP binding transporter protein	Possible adrenal insufficiency, CNS inflammation
IV. ALS-Plus Syndromes						
ALS with fronto-temporal dementia, Parkinson's disease	9p	C9orf72				
Amyotrophy with behavioral disorders Parkinsonism	17q	Tau protein				

Abbreviations: ALS, amyotrophic lateral sclerosis; BSCL2, Bernadelli-Seip congenital lipodystrophy 2B; AD, autosomal dominant; AR, autosomal recessive; CNS, central nervous system; FSP, familial spastic paraplegia; FUS/TLS, fused in sarcoma/translocated in liposarcoma; TDP43, Tar DNA binding protein 43 kd; XR, X-linked recessive.

and gene translation, both in the cytoplasm and locally in dendritic spines in response to electrical activity. How mutations in FUS/TLS provoke motor neuron cell death is not clear, although this may represent loss of function of FUS/TLS in the nucleus or an acquired, toxic function of the mutant proteins in the cytosol. In the third group of ALS genes, the primary problem is defective axonal cytoskeleton and transport (dynactin, profilin-1). It is striking that variants in other genes (e.g., EphA4) influence survival in ALS but not ALS susceptibility. Beyond the upstream, primary defects, it is also evident that the ultimate neuronal cell death process is complex involving multiple cellular processes that accelerate cell death. These include but are not limited to excitotoxicity, impairment of axonal transport, oxidative stress, activation of endoplasmic reticulum stress and the unfolded protein response, and mitochondrial dysfunction.

Multiple recent studies have convincingly demonstrated that non-neuronal cells importantly influence the disease course, at least in ALS transgenic mice. A striking additional finding in neurodegenerative disorders is that miscreant proteins arising from gene defects in familial forms of these diseases are often implicated in sporadic forms of the same disorder. For example, germline mutations in the genes encoding β -amyloid and α -synuclein cause familial forms of Alzheimer's and Parkinson's diseases, and posttranslational, noninherited abnormalities in these proteins are also central to sporadic Alzheimer's and Parkinson's diseases. Analogously, recent reports propose that nonheritable, posttranslational modifications in SOD1 are pathogenic in sporadic ALS.

TREATMENT AMYOTROPHIC LATERAL SCLEROSIS

No treatment arrests the underlying pathologic process in ALS. The drug riluzole (100 mg/d) was approved for ALS because it produces a modest lengthening of survival. In one trial, the survival rate at 18 months with riluzole was similar to placebo at 15 months. The mechanism of this effect is not known with certainty; riluzole may reduce excitotoxicity by diminishing glutamate release. Riluzole is generally well tolerated; nausea, dizziness, weight loss,

and elevated liver enzymes occur occasionally. Pathophysiologic studies of mutant SOD1-related ALS in mice have disclosed diverse targets for therapy; consequently, multiple therapies are presently in clinical trials for ALS including experimental trials of small molecules, mesenchymal stem cells, and immunosuppression. Interventions such as antisense oligonucleotides (ASO) that diminish expression of mutant SOD1 protein prolong survival in transgenic ALS mice and rats and are also nearing trial now for SOD1-mediated ALS.

In the absence of a primary therapy for ALS, a variety of rehabilitative aids may substantially assist ALS patients. Foot-drop splints facilitate ambulation by obviating the need for excessive hip flexion and by preventing tripping on a floppy foot. Finger extension splints can potentiate grip. Respiratory support may be life-sustaining. For patients electing against long-term ventilation by tracheostomy, positive-pressure ventilation by mouth or nose provides transient (several weeks) relief from hypercarbia and hypoxia. Also extremely beneficial for some patients is a respiratory device (Cough Assist Device) that produces an artificial cough. This is highly effective in clearing airways and preventing aspiration pneumonia. When bulbar disease prevents normal chewing and swallowing, gastrostomy is uniformly helpful, restoring normal nutrition and hydration. Fortunately, an increasing variety of speech synthesizers are now available to augment speech when there is advanced bulbar palsy. These facilitate oral communication and may be effective for telephone use.

In contrast to ALS, several of the disorders (Tables 452-1 and 452-3) that bear some clinical resemblance to ALS are treatable. For this reason, a careful search for causes of secondary motor neuron disease is warranted.

OTHER MOTOR NEURON DISEASES

SELECTED LOWER MOTOR NEURON DISORDERS

In these motor neuron diseases, the peripheral motor neurons are affected without evidence of involvement of the corticospinal motor system (Tables 452-1, 452-2, and 452-3).

X-Linked Spinobulbar Muscular Atrophy (Kennedy's Disease) This is an X-linked lower motor neuron disorder in which progressive weakness and wasting of limb and bulbar muscles begins in males in mid-adult life and is conjoined with androgen insensitivity manifested by gynecomastia and reduced fertility (**Chap. 411**). In addition to gynecomastia, which may be subtle, two findings distinguishing this disorder from ALS are the absence of signs of pyramidal tract disease (spasticity) and the presence of a subtle sensory neuropathy in some patients. The underlying molecular defect is an expanded trinucleotide repeat (-CAG-) in the first exon of the androgen receptor gene on the X chromosome. DNA testing is available. An inverse correlation appears to exist between the number of -CAG- repeats and the age of onset of the disease.

Adult Tay-Sachs Disease Several reports have described adult-onset, predominantly lower motor neuropathies arising from deficiency of the enzyme β -hexosaminidase (hex A). These tend to be distinguishable from ALS because they are very slowly progressive; dysarthria and radiographically evident cerebellar atrophy may be prominent. In rare cases, spasticity may also be present, although it is generally absent (**Chap. 432e**).

Spinal Muscular Atrophy The SMAs are a family of selective lower motor neuron diseases of early onset. Despite some phenotypic variability (largely in age of onset), the defect in the majority of families with SMA maps to a locus on chromosome 5 encoding a putative motor neuron survival protein (SMN, for survival motor neuron) that is important in the formation and trafficking of RNA complexes across the nuclear membrane. Neuropathologically these disorders are characterized by extensive loss of large motor neurons; muscle biopsy reveals evidence of denervation atrophy. Several clinical forms exist.

Infantile SMA (SMA I, Werdnig-Hoffmann disease) has the earliest onset and most rapidly fatal course. In some instances it is apparent even before birth, as indicated by decreased fetal movements late in the third trimester. Though alert, afflicted infants are weak and floppy (hypotonic) and lack muscle stretch reflexes. Death generally ensues within the first year of life. *Chronic childhood SMA* (SMA II) begins later in childhood and evolves with a more slowly progressive course. *Juvenile SMA* (SMA III, Kugelberg-Welander disease) manifests during late childhood and runs a slow, indolent course. Unlike most denervating diseases, in this chronic disorder, weakness is greatest in the proximal muscles; indeed, the pattern of clinical weakness can suggest a primary myopathy such as limb-girdle dystrophy. Electrophysiologic and muscle biopsy evidence of denervation distinguish SMA III from the myopathic syndromes. There is no primary therapy for SMA, although remarkable recent experimental data indicate that it may be possible to deliver the missing SMN gene to motor neurons using intravenously or intrathecally delivered adeno-associated viruses (e.g., AAV9) immediately after birth.

Multifocal Motor Neuropathy with Conduction Block In this disorder lower motor neuron function is regionally and chronically disrupted by remarkably focal blocks in conduction. Many cases have elevated serum titers of mono- and polyclonal antibodies to ganglioside GM1; it is hypothesized that the antibodies produce selective, focal, paranodal demyelination of motor neurons. MMCB is not typically associated with corticospinal signs. In contrast with ALS, MMCB may respond dramatically to therapy such as IV immunoglobulin or chemotherapy; thus, it is imperative that MMCB be excluded when considering a diagnosis of ALS.

Other Forms of Lower Motor Neuron Disease In individual families, other syndromes characterized by selective lower motor neuron dysfunction in an SMA-like pattern have been described. There are rare X-linked and autosomal dominant forms of apparent SMA. There is an ALS variant of juvenile onset, the Fazio-Londe syndrome, that involves mainly the musculature innervated by the brainstem. A component of lower motor neuron dysfunction is also found in degenerative disorders such as Machado-Joseph disease and the related olivopontocerebellar degenerations (**Chap. 450**).

SELECTED DISORDERS OF THE UPPER MOTOR NEURON

Primary Lateral Sclerosis This exceedingly rare disorder arises sporadically in adults in mid to late life. Clinically PLS is characterized by progressive spastic weakness of the limbs, preceded or followed by spastic dysarthria and dysphagia, indicating combined involvement of the corticospinal and corticobulbar tracts. Fasciculations, amyotrophy, and sensory changes are absent; neither electromyography nor muscle biopsy shows denervation. On neuropathologic examination, there is selective loss of the large pyramidal cells in the precentral gyrus and degeneration of the corticospinal and corticobulbar projections. The peripheral motor neurons and other neuronal systems are spared. The course of PLS is variable; although long-term survival is documented, the course may be as aggressive as in ALS, with ~3-year survival from onset to death. Early in its course, PLS raises the question of multiple sclerosis or other demyelinating diseases such as adrenoleukodystrophy as diagnostic considerations (**Chap. 458**). A myelopathy suggestive of PLS is infrequently seen with infection with the retrovirus human T cell lymphotropic virus 1 (HTLV-1) (**Chap. 456**). The clinical course and laboratory testing will distinguish these possibilities.

Familial Spastic Paraplegia In its pure form, FSP is usually transmitted as an autosomal trait; most adult-onset cases are dominantly inherited. Symptoms usually begin in the third or fourth decade, presenting as progressive spastic weakness beginning in the distal lower extremities; however, there are variants with onset so early that the differential diagnosis includes cerebral palsy. FSP typically has a long survival, presumably because respiratory function is spared. Late in the illness, there may be urinary urgency and incontinence and sometimes fecal incontinence; sexual function tends to be preserved.

In pure forms of FSP, the spastic leg weakness is often accompanied by posterior column (vibration and position) abnormalities and disturbance of bowel and bladder function. Some family members may have spasticity without clinical symptoms.

By contrast, particularly when recessively inherited, FSP may have complex or complicated forms in which altered corticospinal and dorsal column function is accompanied by significant involvement of other regions of the nervous system, including amyotrophy, mental retardation, optic atrophy, and sensory neuropathy.

Neuropathologically, in FSP, there is degeneration of the corticospinal tracts, which appear nearly normal in the brainstem but show increasing atrophy at more caudal levels in the spinal cord; in effect, the pathologic picture is of a dying-back or distal axonopathy of long neuronal fibers within the CNS.

Defects at numerous loci underlie both dominantly and recessively inherited forms of FSP (Table 452-3). More than 30 FSP genes have now been identified. The gene most commonly implicated in dominantly inherited FSP is *spastin*, which encodes a microtubule interacting protein. The most common childhood-onset dominant form arises from mutations in the *atlastin* gene. A kinesin heavy-chain protein implicated in microtubule motor function was found to be defective in a family with dominantly inherited FSP of variable-onset age.

An infantile-onset form of X-linked, recessive FSP arises from mutations in the gene for myelin proteolipid protein. This is an example of rather striking allelic variation, as most other mutations in the same gene cause not FSP but Pelizaeus-Merzbacher disease, a widespread disorder of CNS myelin. Another recessive variant is caused by defects in the *paraplegin* gene. Paraplegin has homology to metalloproteases that are important in mitochondrial function in yeast.

WEBSITES

Several websites provide valuable information on ALS including those offered by the Muscular Dystrophy Association (www.mdausa.org), the Amyotrophic Lateral Sclerosis Association (www.alsa.org), and the World Federation of Neurology and the Neuromuscular Unit at Washington University in St. Louis (www.neuro.wustl.edu).

453e Prion Diseases

Stanley B. Prusiner, Bruce L. Miller

Prions are proteins that adopt an alternative conformation, which becomes self-propagating. Some prions cause degeneration of the central nervous system (CNS). Once relegated to causing a group of rare disorders of the CNS such as Creutzfeldt-Jakob disease (CJD), prions—as mounting evidence shows—also appear to play a key role in more common illnesses such as Alzheimer’s disease (AD) and Parkinson’s disease (PD). While CJD is caused by the accumulation of PrP^{Sc}, increasing data argue that Aβ prions cause AD, α-synuclein prions cause PD, and tau prions cause the frontotemporal dementias (FTDs). In this chapter, we confine our discussion to CJD, which typically presents with a rapidly progressive dementia as well as motor abnormalities. The illness is relentlessly progressive and generally causes death within 9 months of onset. Most CJD patients are between 50 and 75 years of age; however, patients as young as 17 and as old as 83 have been recorded.

CJD is one malady in a group of disorders caused by prions composed of the prion protein (PrP). PrP prions reproduce by binding to the normal, cellular isoform of the prion protein (PrP^C) and stimulating conversion of PrP^C into the disease-causing isoform PrP^{Sc}. PrP^C is rich in α-helix and has little β-structure, whereas PrP^{Sc} has less α-helix and a high amount of β-structure (Fig. 453e-1). This α-to-β structural transition in PrP is the fundamental event underlying this group of prion diseases (Table 453e-1).

Four new concepts have emerged from studies of prions: (1) Prions are the only known transmissible pathogens that are devoid of nucleic acid; all other infectious agents possess genomes composed of either RNA or DNA that direct the synthesis of their progeny. (2) Prion diseases may be manifest as infectious, genetic, and sporadic disorders; no other group of illnesses with a single etiology presents with such a wide spectrum of clinical manifestations. (3) Prion diseases result from the accumulation of PrP^{Sc}, the conformation of which differs substantially from that of its precursor, PrP^C. (4) Distinct strains of prions exhibit different biologic properties, which are epigenetically inherited. In other words, PrP^{Sc} can exist in a variety of different conformations, many of which seem to specify particular disease phenotypes.

How a specific conformation of a PrP^{Sc} molecule is imparted to PrP^C during prion replication to produce nascent PrP^{Sc} with the same conformation is unknown. Additionally, it is unclear what factors determine where in the CNS a particular PrP^{Sc} molecule will be deposited.

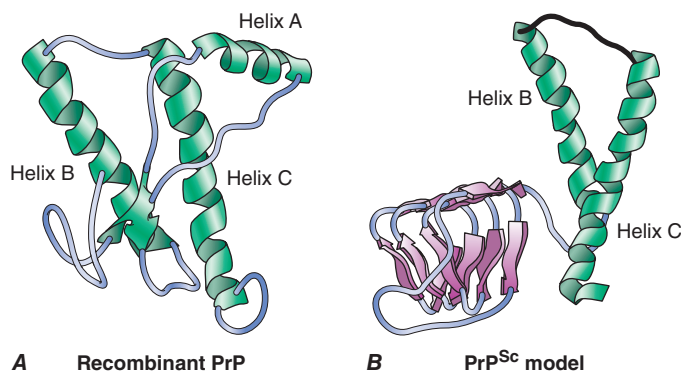


FIGURE 453e-1 Structures of prion proteins. **A.** NMR structure of Syrian hamster recombinant (rec) PrP(90–231). Presumably, the structure of the α-helical form of recPrP(90–231) resembles that of PrP^C. recPrP(90–231) is viewed from the interface where PrP^{Sc} is thought to bind to PrP^C. Shown are: α-helices A (residues 144–157), B (172–193), and C (200–227). Flat ribbons depict β-strands S1 (129–131) and S2 (161–163). **B.** Structural model of PrP^{Sc}. The 90–160 region has been modeled onto a β-helical architecture while the COOH terminal helices B and C are preserved as in PrP^C.

TABLE 453e-1 GLOSSARY OF PRION TERMINOLOGY

Prion	Proteinaceous infectious particle that lacks nucleic acid. Prions are composed entirely of alternatively folded proteins that undergo self-propagation. Distinct strains of prions exhibit different biologic properties, which are epigenetically heritable. PrP prions cause scrapie in sheep and goats, mad cow disease, and related neurodegenerative diseases of humans such as Creutzfeldt-Jakob disease (CJD).
PrP ^{Sc}	Disease-causing isoform of the prion protein. This protein is the only identifiable macromolecule in purified preparations of scrapie prions.
PrP ^C	Cellular isoform of the prion protein. PrP ^C is the precursor of PrP ^{Sc} .
PrP 27-30	A fragment of PrP ^{Sc} , generated by truncation of the NH ₂ -terminus by limited digestion with proteinase K. PrP 27-30 retains prion infectivity and polymerizes into amyloid.
PRNP	PrP gene located on human chromosome 20.
Prion rod	An aggregate of prions composed largely of PrP 27-30 molecules. Created by detergent extraction and limited proteolysis of PrP ^{Sc} . Morphologically and histochemically indistinguishable from many amyloids.
PrP amyloid	Amyloid containing PrP in the brains of animals or humans with prion disease; often accumulates as plaques.

SPECTRUM OF PRION DISEASES

The sporadic form of CJD is the most common prion disorder in humans. Sporadic CJD (sCJD) accounts for ~85% of all cases of human PrP prion disease, whereas inherited prion diseases account for 10–15% of all cases (Table 453e-2). Familial CJD (fCJD), Gerstmann-Sträussler-Scheinker (GSS) disease, and fatal familial insomnia (FFI) are all dominantly inherited prion diseases that are caused by mutations in the PrP gene.



Although infectious PrP prion diseases account for <1% of all cases and infection does not seem to play an important role in the natural history of these illnesses, the transmissibility of

TABLE 453e-2 THE PrP PRION DISEASES

Disease	Host	Mechanism of Pathogenesis
Human		
Kuru	Fore people	Infection through ritualistic cannibalism
iCJD	Humans	Infection from prion-contaminated hGH, dura mater grafts, etc.
vCJD	Humans	Infection from bovine prions
fCJD	Humans	Germline mutations in <i>PRNP</i>
GSS	Humans	Germline mutations in <i>PRNP</i>
FFI	Humans	Germline mutation in <i>PRNP</i> (D178N, M129)
sCJD	Humans	Somatic mutation or spontaneous conversion of PrP ^C into PrP ^{Sc} ?
sFI	Humans	Somatic mutation or spontaneous conversion of PrP ^C into PrP ^{Sc} ?
Animal		
Scrapie	Sheep, goats	Infection in genetically susceptible sheep
BSE	Cattle	Infection with prion-contaminated MBM
TME	Mink	Infection with prions from sheep or cattle
CWD	Mule deer, elk	Unknown
FSE	Cats	Infection with prion-contaminated beef
Exotic ungulate encephalopathy	Greater kudu, nyala, or oryx	Infection with prion-contaminated MBM

Abbreviations: BSE, bovine spongiform encephalopathy; CJD, Creutzfeldt-Jakob disease; CWD, chronic wasting disease; fCJD, familial Creutzfeldt-Jakob disease; FFI, fatal familial insomnia; FSE, feline spongiform encephalopathy; GSS, Gerstmann-Sträussler-Scheinker disease; hGH, human growth hormone; iCJD, iatrogenic Creutzfeldt-Jakob disease; MBM, meat and bone meal; sCJD, sporadic Creutzfeldt-Jakob disease; sFI, sporadic fatal insomnia; TME, transmissible mink encephalopathy; vCJD, variant Creutzfeldt-Jakob disease.

prions is an important biologic feature. Kuru of the Fore people of New Guinea is thought to have resulted from the consumption of brains from dead relatives during ritualistic cannibalism. With the cessation of ritualistic cannibalism in the late 1950s, kuru has nearly disappeared, with the exception of a few recent patients exhibiting incubation periods of >40 years. Iatrogenic CJD (iCJD) seems to be the result of the accidental inoculation of patients with prions. Variant CJD (vCJD) in teenagers and young adults in Europe is the result of exposure to tainted beef from cattle with bovine spongiform encephalopathy (BSE). Although occasional cases of iatrogenic CJD still occur, this form of CJD is currently on the decline due to public health measures aimed at preventing the spread of PrP prions.

Six diseases of animals are caused by prions (Table 453e-2). Scrapie of sheep and goats is the prototypic prion disease. Mink encephalopathy, BSE, feline spongiform encephalopathy, and exotic ungulate encephalopathy are all thought to occur after the consumption of prion-infected foodstuffs. The BSE epidemic emerged in Britain in the late 1980s and was shown to be due to industrial cannibalism. Whether BSE began as a sporadic case of BSE in a cow or started with scrapie in sheep is unknown. The origin of chronic wasting disease (CWD), a prion disease endemic in deer and elk in regions of North America, is uncertain. In contrast to other prion diseases, CWD is highly communicable. Feces from asymptomatic, infected cervids contain prions that are likely to be responsible for the spread of CWD.

EPIDEMIOLOGY

CJD is found throughout the world. The incidence of sCJD is approximately one case per million population, and thus it accounts for approximately 1 in every 10,000 deaths. Because sCJD is an age-dependent neurodegenerative disease, its incidence is expected to increase steadily as older segments of populations in developed and developing countries continue to expand. Although many geographic clusters of CJD have been reported, each has been shown to segregate with a PrP gene mutation. Attempts to identify common exposure to some etiologic agent have been unsuccessful for both the sporadic and familial cases. Ingestion of scrapie-infected sheep or goat meat as a cause of CJD in humans has not been demonstrated by epidemiologic studies, although speculation about this potential route of inoculation continues. Of particular interest are deer hunters who develop CJD, because up to 90% of culled deer in some game herds have been shown to harbor CWD prions. Whether prion disease in deer or elk has passed to cows, sheep, or directly to humans remains unknown. Studies with rodents demonstrate that oral infection with prions can occur, but the process is inefficient compared to intracerebral inoculation.

PATHOGENESIS

The human prion diseases were initially classified as neurodegenerative disorders of unknown etiology on the basis of pathologic changes being confined to the CNS. With the transmission of kuru and CJD to apes, investigators began to view these diseases as infectious CNS illnesses caused by slow viruses. Even though the familial nature of a subset of CJD cases was well described, the significance of this observation became more obscure with the transmission of CJD to animals. Eventually the meaning of heritable CJD became clear with the discovery of mutations in the *PRNP* gene of these patients. The prion concept explains how a disease can manifest as a heritable as well as an infectious illness. Moreover, the hallmark of all prion diseases, whether sporadic, dominantly inherited, or acquired by infection, is that they involve the aberrant metabolism of PrP.

A major feature that distinguishes prions from viruses is the finding that both PrP isoforms are encoded by a chromosomal gene. In humans, the PrP gene is designated *PRNP* and is located on the short arm of chromosome 20. Limited proteolysis of PrP^{Sc} produces a smaller, protease-resistant molecule of ~142 amino acids designated PrP 27-30; PrP^C is

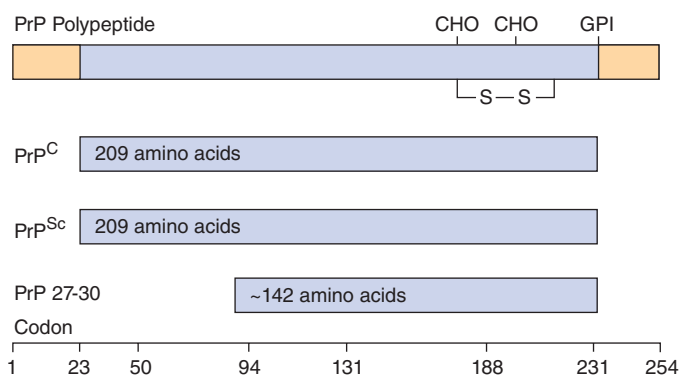


FIGURE 453e-2 Prion protein isoforms. Bar diagram of Syrian hamster PrP, which consists of 254 amino acids. After processing of the NH₂ and COOH termini, both PrP^C and PrP^{Sc} consist of 209 residues. After limited proteolysis, the NH₂ terminus of PrP^{Sc} is truncated to form PrP 27–30 composed of ~142 amino acids. GPI, glycosylphosphatidyl inositol anchor attachment site; S—S, disulfide bond; CHO, N-linked sugars.

completely hydrolyzed under the same conditions (Fig. 453e-2). In the presence of detergent, PrP 27-30 polymerizes into amyloid. Prion rods formed by limited proteolysis and detergent extraction are indistinguishable from the filaments that aggregate to form PrP amyloid plaques in the CNS. Both the rods and the PrP amyloid filaments found in brain tissue exhibit similar ultrastructural morphology and green-gold birefringence after staining with Congo red dye.

Prion Strains Distinct strains of prions exhibit different biologic properties, which are epigenetically heritable. The existence of prion strains raised the question of how heritable biologic information can be enciphered in a molecule other than nucleic acid. Various strains of prions have been defined by incubation times and the distribution of neuronal vacuolation. Subsequently, the patterns of PrP^{Sc} deposition were found to correlate with vacuolation profiles, and these patterns were also used to characterize prion strains.

Persuasive evidence that strain-specific information is enciphered in the tertiary structure of PrP^{Sc} comes from transmission of two different inherited human prion diseases to mice expressing a chimeric human-mouse PrP transgene. In FFI, the protease-resistant fragment of PrP^{Sc} after deglycosylation has a molecular mass of 19 kDa, whereas in fCJD and most sporadic prion diseases, it is 21 kDa (Table 453e-3). This difference in molecular mass was shown to be due to different sites of proteolytic cleavage at the NH₂ termini of the two human PrP^{Sc} molecules, reflecting different tertiary structures. These distinct conformations were not unexpected because the amino acid sequences of the PrPs differ.

Extracts from the brains of patients with FFI transmitted disease into mice expressing a chimeric human-mouse PrP transgene and induced formation of the 19-kDa PrP^{Sc}, whereas brain extracts from

TABLE 453e-3 DISTINCT PRION STRAINS GENERATED IN HUMANS WITH INHERITED PRION DISEASES AND TRANSMITTED TO TRANSGENIC MICE^a

Inoculum	Host Species	Host PrP Genotype	Incubation Time [days ± SEM] (n/n ₀)	PrP ^{Sc} (kDa)
None	Human	FFI(D178N, M129)		19
FFI	Mouse	Tg(MHu2M)	206 ± 7 (7/7)	19
FFI → Tg(MHu2M)	Mouse	Tg(MHu2M)	136 ± 1 (6/6)	19
None	Human	fCJD(E200K)		21
fCJD	Mouse	Tg(MHu2M)	170 ± 2 (10/10)	21
fCJD → Tg(MHu2M)	Mouse	Tg(MHu2M)	167 ± 3 (15/15)	21

^aTg(MHu2M) mice express a chimeric mouse-human PrP gene.

Notes: Clinicopathologic phenotype is determined by the conformation of PrP^{Sc} in accord with the results of the transmission of human prions from patients with FFI to transgenic mice.

Abbreviations: fCJD, familial Creutzfeldt-Jakob disease; FFI, fatal familial insomnia; SEM, standard error of the mean.

fCJD and sCJD patients produced the 21-kDa PrP^{Sc} in mice expressing the same transgene. On second passage, these differences were maintained, demonstrating that chimeric PrP^{Sc} can exist in two different conformations based on the sizes of the protease-resistant fragments, even though the amino acid sequence of PrP^{Sc} is invariant.

This analysis was extended when patients with sporadic fatal insomnia (sFI) were identified. Although they did not carry a *PRNP* gene mutation, the patients demonstrated a clinical and pathologic phenotype that was indistinguishable from that of patients with FFI. Furthermore, 19-kDa PrP^{Sc} was found in their brains, and on passage of prion disease to mice expressing a chimeric human-mouse PrP transgene, 19-kDa PrP^{Sc} was also found. These findings indicate that the disease phenotype is dictated by the conformation of PrP^{Sc} and not the amino acid sequence. PrP^{Sc} acts as a template for the conversion of PrP^C into nascent PrP^{Sc}. On the passage of prions into mice expressing a chimeric hamster-mouse PrP transgene, a change in the conformation of PrP^{Sc} was accompanied by the emergence of a new strain of prions.

Many new strains of prions were generated using recombinant (rec) PrP produced in bacteria; recPrP was polymerized into amyloid fibrils and inoculated into transgenic mice expressing high levels of wild-type mouse PrP^C; approximately 500 days later, the mice died of prion disease. The incubation times of the “synthetic prions” in mice were dependent on the conditions used for polymerization of the amyloid fibrils. Highly stable amyloids gave rise to stable prions with long incubation times; low-stability amyloids led to prions with short incubation times. Amyloids of intermediate stability gave rise to prions with intermediate stabilities and intermediate incubation times. Such findings are consistent with earlier studies showing that the incubation times of synthetic and naturally occurring prions are directly proportional to the stability of the prion.

Species Barrier Studies on the role of the primary and tertiary structures of PrP in the transmission of prion disease have given new insights into the pathogenesis of these maladies. The amino acid sequence of PrP encodes the species of the prion, and the prion derives its PrP^{Sc} sequence from the last mammal in which it was passed. While the primary structure of PrP is likely to be the most important or even sole determinant of the tertiary structure of PrP^C, PrP^{Sc} seems to function as a template in determining the tertiary structure of nascent PrP^{Sc} molecules as they are formed from PrP^C. In turn, prion diversity appears to be enciphered in the conformation of PrP^{Sc}, and thus prion strains seem to represent different conformers of PrP^{Sc}.

In general, transmission of PrP prion disease from one species to another is inefficient, in that not all intracerebrally inoculated animals develop disease, and those that fall ill do so only after long incubation times that can approach the natural life span of the animal. This “species barrier” to transmission is correlated with the degree of similarity between the amino acid sequences of PrP^C in the inoculated host and of PrP^{Sc} in the prion inoculum. The importance of sequence similarity between the host and donor PrP argues that PrP^C directly interacts with PrP^{Sc} in the prion conversion process.

SPORADIC AND INHERITED PrP PRION DISEASES

Several different scenarios might explain the initiation of sporadic prion disease: (1) A somatic mutation may be the cause and thus follow a path similar to that for germline mutations in inherited disease. In this situation, the mutant PrP^{Sc} must be capable of targeting wild-type PrP^C, a process known to be possible for some mutations but less likely for others. (2) The activation energy barrier separating wild-type PrP^C from PrP^{Sc} could be crossed on rare occasions when viewed in the context of a population. Most individuals would be spared, while presentations in the elderly with an incidence of ~1 per million would be seen. (3) PrP^{Sc} may be present at low levels in some normal cells, where it performs some important, as yet unknown, function. The level of PrP^{Sc} in such cells is hypothesized to be sufficiently low as to be not detected by routine bioassay. In some altered metabolic states, the cellular mechanisms for clearing PrP^{Sc} might become compromised, and the rate of PrP^{Sc} formation would then begin to exceed the capacity of

the cell to clear it. The third possible mechanism is attractive because it suggests PrP^{Sc} is not simply a misfolded protein, as proposed for the first and second mechanisms, but that it is an alternatively folded molecule with a function. Moreover, the multitude of conformational states that PrP^{Sc} can adopt, as described above, raises the possibility that PrP^{Sc} or another prion-like protein might function in a process like short-term memory where information storage occurs in the absence of new protein synthesis.

More than 40 different mutations resulting in nonconservative substitutions in the human *PRNP* gene have been found to segregate with inherited human prion diseases. Missense mutations and expansions in the octapeptide repeat region of the gene are responsible for familial forms of prion disease. Five different mutations of the *PRNP* gene have been linked genetically to heritable prion disease.

Although phenotypes may vary dramatically within families, specific phenotypes tend to be observed with certain mutations. A clinical phenotype indistinguishable from typical sCJD is usually seen with substitutions at codons 180, 183, 200, 208, 210, and 232. Substitutions at codons 102, 105, 117, 198, and 217 are associated with the GSS variant of prion disease. The normal human PrP sequence contains five repeats of an eight-amino-acid sequence. Insertions from two to nine extra octarepeats frequently cause variable phenotypes ranging from a condition indistinguishable from sCJD to a slowly progressive dementing illness of many years in duration to an early-age-of-onset disorder that is similar to AD. A mutation at codon 178 resulting in substitution of asparagine for aspartic acid produces FFI if a methionine is encoded at the polymorphic residue 129 on the same allele. Typical CJD is seen if the D178N mutation occurs with a valine encoded at position 129 of the same allele.

HUMAN PRNP GENE POLYMORPHISMS

Polymorphisms influence the susceptibility to sporadic, inherited, and infectious forms of prion disease. The methionine/valine polymorphism at position 129 not only modulates the age of onset of some inherited prion diseases but can also determine the clinical phenotype. The finding that homozygosity at codon 129 predisposes to sCJD supports a model of prion production that favors PrP interactions between homologous proteins.

Substitution of the basic residue lysine at position 218 in mouse PrP produced dominant-negative inhibition of prion replication in transgenic mice. This same lysine at position 219 in human PrP has been found in 12% of the Japanese population, and this group appears to be resistant to prion disease. Dominant-negative inhibition of prion replication was also found with substitution of the basic residue arginine at position 171; sheep with arginine were resistant to scrapie prions but were susceptible to BSE prions that were inoculated intracerebrally.

INFECTIOUS PrP PRION DISEASES

IATROGENIC CJD

Accidental transmission of CJD to humans appears to have occurred with corneal transplantation, contaminated electroencephalogram (EEG) electrode implantation, and surgical procedures. Corneas from donors with unsuspected CJD have been transplanted to apparently healthy recipients who developed CJD after variable incubation periods. The same improperly decontaminated EEG electrodes that caused CJD in two young patients with intractable epilepsy caused CJD in a chimpanzee 18 months after their experimental implantation.

Surgical procedures may have resulted in accidental inoculation of patients with prions, presumably because some instrument or apparatus in the operating theater became contaminated when a CJD patient underwent surgery. Although the epidemiology of these studies is highly suggestive, no proof for such episodes exists.

Dura Mater Grafts More than 160 cases of CJD after implantation of dura mater grafts have been recorded. All of the grafts appear to have been acquired from a single manufacturer whose preparative procedures were inadequate to inactivate human prions. One case of CJD occurred after repair of an eardrum perforation with a pericardium graft.

Human Growth Hormone and Pituitary Gonadotropin Therapy The transmission of CJD prions from contaminated human growth hormone (hGH) preparations derived from human pituitaries has been responsible for fatal cerebellar disorders with dementia in >180 patients ranging in age from 10 to 41 years. These patients received injections of hGH every 2–4 days for 4–12 years. If it is thought that these patients developed CJD from injections of prion-contaminated hGH preparations, the possible incubation periods range from 4 to 30 years. Only recombinant hGH is now used therapeutically so that possible contamination with prions is no longer an issue. Four cases of CJD have occurred in women receiving human pituitary gonadotropin.

VARIANT CJD

The restricted geographic occurrence and chronology of vCJD raised the possibility that BSE prions had been transmitted to humans through the consumption of tainted beef. More than 190 cases of vCJD have occurred, with >90% of these in Britain. vCJD has also been reported in people either living in or originating from France, Ireland, Italy, Netherlands, Portugal, Spain, Saudi Arabia, United States, Canada, and Japan.

The steady decline in the number of vCJD cases over the past decade argues that there will not be a prion disease epidemic in Europe, similar to those seen for BSE and kuru. What is certain is that prion-tainted meat should be prevented from entering the human food supply.

The most compelling evidence that vCJD is caused by BSE prions was obtained from experiments in mice expressing the bovine PrP transgene. Both BSE and vCJD prions were efficiently transmitted to these transgenic mice and with similar incubation periods. In contrast to sCJD prions, vCJD prions did not transmit disease efficiently to mice expressing a chimeric human-mouse PrP transgene. Earlier studies with nontransgenic mice suggested that vCJD and BSE might be derived from the same source because both inocula transmitted disease with similar but very long incubation periods.

Attempts to determine the origin of BSE and vCJD prions have relied on passaging studies in mice, some of which are described above, as well as studies of the conformation and glycosylation of PrP^{Sc}. One scenario suggests that a particular conformation of bovine PrP^{Sc} was selected for heat resistance during the rendering process and was then reselected multiple times as cattle infected by ingesting prion-contaminated meat and bone meal (MBM) were slaughtered and their offal rendered into more MBM. Variant CJD cases have virtually disappeared with protection of the beef supply in Europe.

NEUROPATHOLOGY

Frequently the brains of patients with CJD have no recognizable abnormalities on gross examination. Patients who survive for several years have variable degrees of cerebral atrophy.

On light microscopy, the pathologic hallmarks of CJD are spongiform degeneration and astrocytic gliosis. The lack of an inflammatory response in CJD and other prion diseases is an important pathologic feature of these degenerative disorders. Spongiform degeneration is characterized by many 1- to 5- μ m vacuoles in the neuropil between nerve cell bodies. Generally the spongiform changes occur in the cerebral cortex, putamen, caudate nucleus, thalamus, and molecular layer of the cerebellum. Astrocytic gliosis is a constant but nonspecific feature of prion diseases. Widespread proliferation of fibrous astrocytes is found throughout the gray matter of brains infected with CJD prions. Astrocytic processes filled with glial filaments form extensive networks.

Amyloid plaques have been found in ~10% of CJD cases. Purified CJD prions from humans and animals exhibit the ultrastructural and histochemical characteristics of amyloid when treated with detergents during limited proteolysis. In first passage from some human Japanese CJD cases, amyloid plaques have been found in mouse brains. These plaques stain with antibodies raised against PrP.

The amyloid plaques of GSS disease are morphologically distinct from those seen in kuru or scrapie. GSS plaques consist of a central dense core of amyloid surrounded by smaller globules of amyloid. Ultrastructurally, they consist of a radiating fibrillar network of amyloid

fibrils, with scant or no neuritic degeneration. The plaques can be distributed throughout the brain but are most frequently found in the cerebellum. They are often located adjacent to blood vessels. Congophilic angiopathy has been noted in some cases of GSS disease.

In vCJD, a characteristic feature is the presence of “florid plaques.” These are composed of a central core of PrP amyloid, surrounded by vacuoles in a pattern suggesting petals on a flower.

CLINICAL FEATURES

Nonspecific prodromal symptoms occur in approximately a third of patients with CJD and may include fatigue, sleep disturbance, weight loss, headache, anxiety, vertigo, malaise, and ill-defined pain. Most patients with CJD present with deficits in higher cortical function. Similarly, psychiatric symptoms, such as depression, psychosis, and visual hallucinations, are often the defining features of the illness. These deficits almost always progress over weeks or months to a state of profound dementia characterized by memory loss, impaired judgment, and a decline in virtually all aspects of intellectual function. A few patients present with either visual impairment or cerebellar gait and coordination deficits. Frequently the cerebellar deficits are rapidly followed by progressive dementia. Visual problems often begin with blurred vision and diminished acuity, rapidly followed by dementia.

Other symptoms and signs include extrapyramidal dysfunction manifested as rigidity, masklike facies, or (less commonly) choreo-athetoid movements; pyramidal signs (usually mild); seizures (usually major motor) and, less commonly, hypoesthesia; supranuclear gaze palsy; optic atrophy; and vegetative signs such as changes in weight, temperature, sweating, or menstruation.

Myoclonus Most patients (~90%) with CJD exhibit myoclonus that appears at various times throughout the illness. Unlike other involuntary movements, myoclonus persists during sleep. Startle myoclonus elicited by loud sounds or bright lights is frequent. It is important to stress that myoclonus is neither specific nor confined to CJD and tends to occur later in the course of CJD. Dementia with myoclonus can also be due to AD (**Chap. 448**), dementia with Lewy bodies (**Chap. 448**), corticobasal degeneration (**Chap. 448**), cryptococcal encephalitis (**Chap. 239**), or the myoclonic epilepsy disorder Unverricht-Lundborg disease (**Chap. 445**).

Clinical Course In documented cases of accidental transmission of CJD to humans, an incubation period of 1.5–2 years preceded the development of clinical disease. In other cases, incubation periods of up to 40 years have been suggested. Most patients with CJD live 6–12 months after the onset of clinical signs and symptoms, whereas some live for up to 5 years.

DIAGNOSIS

The constellation of dementia, myoclonus, and periodic electrical bursts in an afebrile 60-year-old patient generally indicates CJD. Clinical abnormalities in CJD are confined to the CNS. Fever, elevated sedimentation rate, leukocytosis in blood, or a pleocytosis in cerebrospinal fluid (CSF) should alert the physician to another etiology to explain the patient’s CNS dysfunction, although there are rare cases of CJD in which mild CSF pleocytosis is observed.

Variations in the typical course appear in inherited and transmitted forms of the disease. fCJD has an earlier mean age of onset than sCJD. In GSS disease, ataxia is usually a prominent and presenting feature, with dementia occurring late in the disease course. GSS disease presents earlier than CJD (mean age 43 years) and is typically more slowly progressive than CJD; death usually occurs within 5 years of onset. FFI is characterized by insomnia and dysautonomia; dementia occurs only in the terminal phase of the illness. Rare sporadic cases have been identified. vCJD has an unusual clinical course, with a prominent psychiatric prodrome that may include visual hallucinations and early ataxia, whereas frank dementia is usually a late sign of vCJD.

DIFFERENTIAL DIAGNOSIS

Many conditions mimic CJD. Dementia with Lewy bodies (**Chap. 448**) is the most common disorder to be mistaken for CJD. It can present

in a subacute fashion with delirium, myoclonus, and extrapyramidal features. Other neurodegenerative disorders (Chap. 448) to consider include AD, FTD, corticobasal degeneration, progressive supranuclear palsy, ceroid lipofuscinosis, and myoclonic epilepsy with Lafora bodies (Chap. 445). The absence of abnormalities on diffusion-weighted and fluid-attenuated inversion recovery (FLAIR) magnetic resonance imaging (MRI) will almost always distinguish these dementing conditions from CJD.

Hashimoto's encephalopathy, which presents as a subacute progressive encephalopathy with myoclonus and periodic triphasic complexes on the EEG, should be excluded in every case of suspected CJD. It is diagnosed by the finding of high titers of antithyroglobulin or antithyroid peroxidase (antimicrosomal) antibodies in the blood and improves with glucocorticoid therapy. Unlike CJD, fluctuations in severity typically occur in Hashimoto's encephalopathy.

Intracranial vasculitides (Chap. 385) may produce nearly all of the symptoms and signs associated with CJD, sometimes without systemic abnormalities. Myoclonus is exceptional with cerebral vasculitis, but focal seizures may confuse the picture. Prominent headache, absence of myoclonus, stepwise change in deficits, abnormal CSF, and focal white matter changes on MRI or angiographic abnormalities all favor vasculitis.

Paraneoplastic conditions (Chap. 122), particularly limbic encephalitis and cortical encephalitis, can also mimic CJD. In many of these patients, dementia appears prior to the diagnosis of a tumor, and in some, no tumor is ever found. Detection of the paraneoplastic antibodies is often the only way to distinguish these cases from CJD.

Other diseases that can simulate CJD include neurosyphilis (Chap. 206), AIDS dementia complex (Chap. 226), progressive multifocal leukoencephalopathy (Chap. 164), subacute sclerosing panencephalitis, progressive rubella panencephalitis, herpes simplex encephalitis (Chap. 164), diffuse intracranial tumor (gliomatosis cerebri; Chap. 118), anoxic encephalopathy, dialysis dementia, uremia, hepatic encephalopathy, voltage-gated potassium channel (VGKC) autoimmune encephalopathy, and lithium or bismuth intoxication.

LABORATORY TESTS

The only specific diagnostic tests for CJD and other human prion diseases measure PrP^{Sc}. The most widely used method involves limited proteolysis that generates PrP 27-30, which is detected by immunoassay after denaturation. The conformation-dependent immunoassay (CDI) is based on immunoreactive epitopes that are exposed in PrP^C but buried in PrP^{Sc}. In humans, the diagnosis of CJD can be established by brain biopsy if PrP^{Sc} is detected. If no attempt is made to measure PrP^{Sc}, but the constellation of pathologic changes frequently found in CJD is seen in a brain biopsy, then the diagnosis is reasonably secure (see "Neuropathology," above). The high sensitivity and specificity of cortical ribboning and basal ganglia hyperintensity on FLAIR and diffusion-weighted MRI for the diagnosis of CJD have greatly diminished the need for brain biopsy in patients with suspected CJD. Because PrP^{Sc} is not uniformly distributed throughout the CNS, the apparent absence of PrP^{Sc} in a limited sample such as a biopsy does not rule out prion disease. At autopsy, sufficient brain samples should be taken for both PrP^{Sc} immunoassay, preferably by CDI, and immunohistochemistry of tissue sections.

To establish the diagnosis of either sCJD or familial prion disease, sequencing the *PRNP* gene must be performed. Finding the wild-type *PRNP* gene sequence permits the diagnosis of sCJD if there is no history to suggest infection from an exogenous source of prions. The identification of a mutation in the *PRNP* gene sequence that encodes a nonconservative amino acid substitution argues for familial prion disease.

CT may be normal or show cortical atrophy. MRI is valuable for distinguishing sCJD from most other conditions. On FLAIR sequences and diffusion-weighted imaging, ~90% of patients show increased intensity in the basal ganglia and cortical ribboning (Fig. 453e-3). This pattern is not seen with other neurodegenerative disorders but has been seen infrequently with viral encephalitis, paraneoplastic syndromes, or seizures. When the typical MRI pattern is present, in the

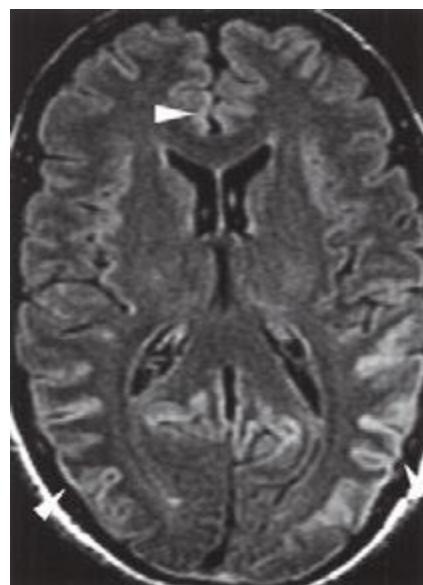


FIGURE 453e-3 T2-weighted (fluid-attenuated inversion recovery) magnetic resonance imaging showing hyperintensity in the cortex in a patient with sporadic CJD. This so-called “cortical ribboning” along with increased intensity in the basal ganglia on T2- or diffusion-weighted imaging can aid in the diagnosis of Creutzfeldt-Jakob disease.

proper clinical setting, diagnosis is facilitated. However, some cases of sCJD do not show this typical pattern, and other early diagnostic approaches are still needed.

CSF is nearly always normal but may show protein elevation and, rarely, mild pleocytosis. Although the stress protein 14-3-3 is elevated in the CSF of some patients with CJD, similar elevations of 14-3-3 are found in patients with other disorders; thus this elevation is not specific. Similarly, elevations of CSF neuron-specific enolase and tau occur in CJD but lack specificity for diagnosis.

The EEG is often useful in the diagnosis of CJD, although only approximately 60% of individuals show the typical pattern. During the early phase of CJD, the EEG is usually normal or shows only scattered theta activity. In most advanced cases, repetitive, high-voltage, triphasic, and polyphasic sharp discharges are seen, but in many cases their presence is transient. The presence of these stereotyped periodic bursts of <200 ms in duration, occurring every 1–2 s, makes the diagnosis of CJD very likely. These discharges are frequently but not always symmetric; there may be a one-sided predominance in amplitude. As CJD progresses, normal background rhythms become fragmentary and slower.

CARE OF CJD PATIENTS

Although CJD should not be considered either contagious or communicable, it is transmissible. The risk of accidental inoculation by aerosols is very small; nonetheless, procedures producing aerosols should be performed in certified biosafety cabinets. Biosafety level 2 practices, containment equipment, and facilities are recommended by the Centers for Disease Control and Prevention and the National Institutes of Health. The primary problem in caring for patients with CJD is the inadvertent infection of health care workers by needle and stab wounds. Electroencephalographic and electromyographic needles should not be reused after studies on patients with CJD have been performed.

There is no reason for pathologists or other morgue employees to resist performing autopsies on patients whose clinical diagnosis was CJD. Standard microbiologic practices outlined here, along with specific recommendations for decontamination, seem to be adequate precautions for the care of patients with CJD and the handling of infected specimens.

Prions are extremely resistant to common inactivation procedures, and there is some disagreement about the optimal conditions for sterilization. Some investigators recommend treating CJD-contaminated materials once with 1 *N* NaOH at room temperature, but we believe this procedure may be inadequate for sterilization. Autoclaving at 134°C for 5 h or treatment with 2 *N* NaOH for several hours is recommended for sterilization of prions. The term *sterilization* implies complete destruction of prions; any residual infectivity can be hazardous. Recent studies show that sCJD prions bound to stainless steel surfaces are resistant to inactivation by autoclaving at 134°C for 2 h; exposure of bound prions to an acidic detergent solution prior to autoclaving rendered prions susceptible to inactivation.

PREVENTION AND THERAPEUTICS

There is no known effective therapy for preventing or treating CJD. The finding that phenothiazines and acridines inhibit PrP^{Sc} formation in cultured cells led to clinical studies of quinacrine in CJD patients. Unfortunately, quinacrine failed to slow the rate of cognitive decline in CJD, possibly because therapeutic concentrations in the brain were not achieved. Although inhibition of the P-glycoprotein (Pgp) transport system resulted in substantially increased quinacrine levels in the brains of mice, the prion incubation times were not extended by treatment with the drug. Whether such an approach can be used to treat CJD remains to be established.

Like the acridines, anti-PrP antibodies have been shown to eliminate PrP^{Sc} from cultured cells. Additionally, such antibodies in mice, either administered by injection or produced from a transgene, have been shown to prevent prion disease when prions are introduced by a peripheral route, such as intraperitoneal inoculation. Unfortunately, the antibodies were ineffective in mice inoculated intracerebrally with prions. Several drugs, including pentosan polysulfate as well as porphyrin and phenylhydrazine derivatives, delay the onset of disease in animals inoculated intracerebrally with prions if the drugs are given intracerebrally beginning soon after inoculation.

DIFFERENT PRIONS CAUSING OTHER NEURODEGENERATIVE DISEASES

There is a rapidly expanding body of literature demonstrating that in addition to PrP, other proteins including amyloid beta (A β), tau, α -synuclein, and huntingtin can all become prions ([Chap. 444e](#)).

Experimental studies have shown that transgenic mice expressing mutant amyloid precursor protein (APP) develop amyloid plaques containing fibrils composed of the A β peptide approximately a year after inoculation with synthetic A β peptides polymerized into amyloid fibrils or extracts prepared from the brains of patients with AD. Mutant tau aggregates in transgenic mice and cultured cells can trigger the aggregation of tau into fibrils that resemble those found in neurofibrillary tangles and Pick bodies. Such tangles have been found in AD, FTDs, Pick's disease, and some cases of posttraumatic brain injury, all of which are likely to be caused by the prion isoforms of A β and/or tau.

In patients with advanced PD who received grafts of fetal substantia nigral neurons, Lewy bodies containing β -sheet-rich α -synuclein were identified in grafted cells approximately 10 years after transplantation, arguing for the axonal transport of misfolded α -synuclein crossing into grafted neurons, where it initiated aggregation of nascent α -synuclein into fibrils that coalesced into Lewy bodies. These findings combined with studies of multiple system atrophy (MSA) argue that the synucleinopathies are caused by prions. Brain homogenates from MSA patients injected into transgenic mice transmitted lethal neurodegeneration in approximately 3 months; moreover, recombinant synuclein injected into wild-type mice initiated the deposition of synuclein fibrils.

In summary, a wealth of evidence continues to accumulate arguing that proteins causing AD, PD, FTDs, ALS, and even HD acquire alternative conformations that become self-propagating. Each of these neurodegenerative diseases is caused by a different protein that undergoes a conformational transformation to become a prion. Prions explain many of the features that the neurodegenerative diseases have in common: (1) incidence increases with age, (2) steady progression over years, (3) spread from one region of the CNS to another, (4) protein deposits consisting of amyloid fibrils, and (5) late onset of inherited forms of the neurodegenerative diseases. Notably, amyloid plaques containing PrP^{Sc} are a nonobligatory feature of PrP prion disease in humans and animals. Furthermore, amyloid plaques in AD do not correlate with the level of dementia; however, the level of soluble (oligomeric) A β peptide does correlate with memory loss and other intellectual deficits.

454 Disorders of the Autonomic Nervous System

Phillip A. Low, John W. Engstrom

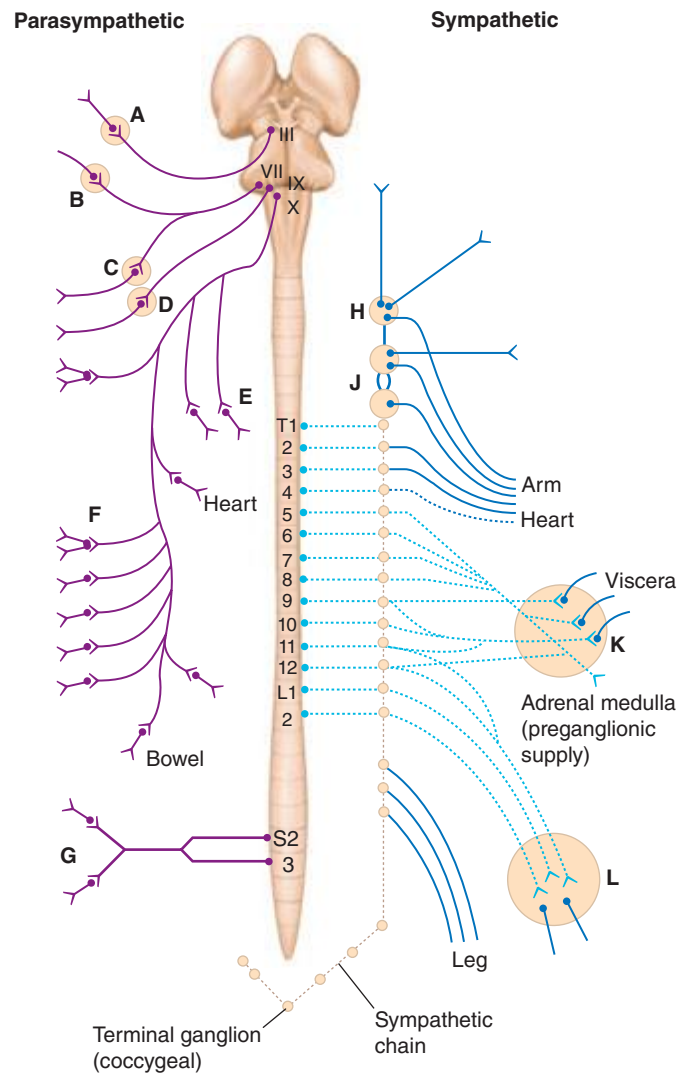
The autonomic nervous system (ANS) innervates the entire neuraxis and influences all organ systems. It regulates blood pressure (BP), heart rate, sleep, and bladder and bowel function. It operates automatically; its full importance becomes recognized only when ANS function is compromised, resulting in dysautonomia. **Hypothalamic disorders that cause disturbances in homeostasis are discussed in Chaps. 23 and 401e.**

ANATOMIC ORGANIZATION

The activity of the ANS is regulated by central neurons responsive to diverse afferent inputs. After central integration of afferent information, autonomic outflow is adjusted to permit the functioning of the major organ systems in accordance with the needs of the whole organism. Connections between the cerebral cortex and the autonomic centers in the brainstem coordinate autonomic outflow with higher mental functions.

The preganglionic neurons of the parasympathetic nervous system leave the central nervous system (CNS) in the third, seventh, ninth, and tenth cranial nerves as well as the second and third sacral nerves, while the preganglionic neurons of the sympathetic nervous system exit the spinal cord between the first thoracic and the second lumbar segments (Fig. 454-1). These are thinly myelinated. The postganglionic neurons, located in ganglia outside the CNS, give rise to the postganglionic unmyelinated autonomic nerves that innervate organs and tissues throughout the body. Responses to sympathetic and parasympathetic stimulation are frequently antagonistic (Table 454-1), reflecting highly coordinated interactions within the CNS; the resultant changes in parasympathetic and sympathetic activity provide more precise control of autonomic responses than could be achieved by the modulation of a single system.

Acetylcholine (ACh) is the preganglionic neurotransmitter for both divisions of the ANS as well as the postganglionic neurotransmitter



Parasympathetic system
from cranial nerves III, VII, IX, X
and from sacral nerves 2 and 3

Sympathetic system

from T1-L2
Preganglionic fibers
Postganglionic fibers ———

- | | |
|--|---|
| A Ciliary ganglion | H Superior cervical ganglion |
| B Sphenopalatine (pterygopalatine) ganglion | J Middle cervical ganglion and inferior cervical (stellate) ganglion including T1 ganglion |
| C Submandibular ganglion | K Coeliac and other abdominal ganglia |
| D Otic ganglion | L Lower abdominal sympathetic ganglia |
| E Vagal ganglion cells in the heart wall | |
| F Vagal ganglion cells in bowel wall | |
| G Pelvic ganglia | |

FIGURE 454-1 Schematic representation of the autonomic nervous system. (From M Moskowitz: *Clin Endocrinol Metab* 6:77, 1977.)

of the parasympathetic neurons; the preganglionic receptors are nicotinic, and the postganglionic are muscarinic in type. Norepinephrine (NE) is the neurotransmitter of the postganglionic sympathetic neurons, except for cholinergic neurons innervating the eccrine sweat glands.

CLINICAL EVALUATION

CLASSIFICATION

Disorders of the ANS may result from pathology of either the CNS or the peripheral nervous system (PNS) (Table 454-2). Signs and symptoms may result from interruption of the afferent limb, CNS

TABLE 454-1 FUNCTIONAL CONSEQUENCES OF NORMAL ANS ACTIVATION

	Sympathetic	Parasympathetic
Heart rate	Increased	Decreased
Blood pressure	Increased	Mildly decreased
Bladder	Increased sphincter tone	Voiding (decreased tone)
Bowel motility	Decreased motility	Increased
Lung	Bronchodilation	Bronchoconstriction
Sweat glands	Sweating	—
Pupils	Dilation	Constriction
Adrenal glands	Catecholamine release	—
Sexual function	Ejaculation, orgasm	Erection
Lacrimal glands	—	Tearing
Parotid glands	—	Salivation

processing centers, or efferent limb of reflex arcs controlling autonomic responses. For example, a lesion of the medulla produced by a posterior fossa tumor can impair BP responses to postural changes and result in orthostatic hypotension (OH). OH can also be caused by lesions of the spinal cord or peripheral vasomotor nerve fibers (e.g., diabetic autonomic neuropathy). Lesions of the efferent limb cause the most consistent and severe OH. The site of reflex interruption is usually established by the clinical context in which the dysautonomia arises, combined with judicious use of ANS testing and neuroimaging studies. The presence or absence of CNS signs, association with sensory or motor polyneuropathy, medical illnesses, medication use, and family history are often important considerations. Some syndromes do not fit easily into any classification scheme.

SYMPTOMS OF AUTONOMIC DYSFUNCTION

Clinical manifestations can result from loss of function, overactivity, or dysregulation of autonomic circuits. Disorders of autonomic function should be considered in patients with unexplained OH, syncope, sleep dysfunction, altered sweating (hyperhidrosis or hypohidrosis), impotence, constipation or other gastrointestinal symptoms (bloating,

TABLE 454-2 CLASSIFICATION OF CLINICAL AUTONOMIC DISORDERS

I. Autonomic disorders with brain involvement	d. Cardiac arrhythmias
A. Associated with multisystem degeneration	e. Central sleep apnea
1. Multisystem degeneration: autonomic failure clinically prominent	f. Baroreflex failure
a. Multiple system atrophy (MSA)	g. Horner's syndrome
b. Parkinson's disease with autonomic failure	h. Vertebrobasilar and lateral medullary (Wallenberg's) syndromes
c. Diffuse Lewy body disease (some cases)	i. Brainstem encephalitis
2. Multisystem degeneration: autonomic failure clinically not usually prominent	II. Autonomic disorders with spinal cord involvement
a. Parkinson's disease	A. Traumatic quadriplegia
b. Other extrapyramidal disorders (inherited spinocerebellar atrophies, progressive supranuclear palsy, corticobasal degeneration, Machado-Joseph disease, fragile X syndrome [FXTAS])	B. Syringomyelia
B. Unassociated with multisystem degeneration (focal CNS disorders)	C. Subacute combined degeneration
1. Disorders mainly due to cerebral cortex involvement	D. Multiple sclerosis and neuromyelitis optica
a. Frontal cortex lesions causing urinary/bowel incontinence	E. Amyotrophic lateral sclerosis
b. Focal seizures (temporal lobe or anterior cingulate)	F. Tetanus
c. Cerebral infarction of the insula	G. Stiff-person syndrome
2. Disorders of the limbic and paralimbic circuits	H. Spinal cord tumors
a. Shapiro's syndrome (agenesis of corpus callosum, hyperhidrosis, hypothermia)	III. Autonomic neuropathies
b. Autonomic seizures	A. Acute/subacute autonomic neuropathies
c. Limbic encephalitis	1. Subacute autoimmune autonomic ganglionopathy (AAG)
3. Disorders of the hypothalamus	a. Subacute paraneoplastic autonomic neuropathy
a. Thiamine deficiency (Wernicke-Korsakoff syndrome)	b. Guillain-Barré syndrome
b. Diencephalic syndrome	c. Botulism
c. Neuroleptic malignant syndrome	d. Porphyrria
d. Serotonin syndrome	e. Drug induced autonomic neuropathies-stimulants, drug withdrawal, vasoconstrictor, vasodilators, beta-receptor antagonists, beta-agonists
e. Fatal familial insomnia	f. Toxin-induced autonomic neuropathies
f. Antidiuretic hormone (ADH) syndromes (diabetes insipidus, inappropriate ADH secretion)	g. Subacute cholinergic neuropathy
g. Disturbances of temperature regulation (hyperthermia, hypothermia)	B. Chronic peripheral autonomic neuropathies
h. Disturbances of sexual function	1. Distal small fiber neuropathy
i. Disturbances of appetite	2. Combined sympathetic and parasympathetic failure
j. Disturbances of BP/HR and gastric function	a. Amyloid
k. Horner's syndrome	b. Diabetic autonomic neuropathy
4. Disorders of the brainstem and cerebellum	c. Autoimmune autonomic ganglionopathy (paraneoplastic and idiopathic)
a. Posterior fossa tumors	d. Sensory neuronopathy with autonomic failure
b. Syringobulbia and Arnold-Chiari malformation	e. Familial dysautonomia (Riley-Day syndrome)
c. Disorders of BP control (hypertension, hypotension)	f. Diabetic, uremic, or nutritional deficiency
	g. Dysautonomia of old age
	3. Disorders of reduced orthostatic intolerance: reflex syncope, POTS, associated with prolonged bed rest, associated with space flight, chronic fatigue

Abbreviations: BP, blood pressure; CNS, central nervous system; HR, heart rate; POTS, postural orthostatic tachycardia syndrome.

TABLE 454-3 SYMPTOMS OF ORTHOSTATIC INTOLERANCE

Lightheadedness (dizziness)	88%
Weakness or tiredness	72%
Cognitive difficulty (thinking/ concentrating)	47%
Blurred vision	47%
Tremulousness	38%
Vertigo	37%
Pallor	31%
Anxiety	29%
Palpitations	26%
Clammy feeling	19%
Nausea	18%

Source: PA Low et al: Mayo Clin Proc 70:617, 1995.

nausea, vomiting of old food, diarrhea), or bladder disorders (urinary frequency, hesitancy, or incontinence). Symptoms may be widespread or regional in distribution. An autonomic history focuses on systemic functions (BP, heart rate, sleep, fever, sweating) and involvement of individual organ systems (pupils, bowel, bladder, sexual function). The autonomic symptom profile is a self-report questionnaire that can be used for formal assessment. It is also important to recognize the modulating effects of age. For example, OH typically produces lightheadedness in the young, whereas cognitive slowing is more common in the elderly. Specific symptoms of orthostatic intolerance are diverse (Table 454-3). Autonomic symptoms may vary dramatically, reflecting the dynamic nature of autonomic control over homeostatic function. For example, OH might be manifest only in the early morning, following a meal, with exercise, or with raised ambient temperature, depending on the regional vascular bed affected by the dysautonomia.

Early symptoms may be overlooked. Impotence, although not specific for autonomic failure, often heralds autonomic failure in men and may precede other symptoms by years (Chap. 67). A decrease in the frequency of spontaneous early morning erections may occur months before loss of nocturnal penile tumescence and development of total impotence. Bladder dysfunction may appear early in men and women, particularly in those with a CNS etiology. Cold feet may indicate increased peripheral vasomotor constriction. Brain and spinal cord disease above the level of the lumbar spine results first in urinary frequency and small bladder volumes and eventually in incontinence (upper motor neuron or spastic bladder). By contrast, PNS disease of autonomic nerve fibers results in large bladder volumes, urinary frequency, and overflow incontinence (lower motor neuron flaccid bladder). Measurement of bladder volume (postvoid residual) is a useful bedside test for distinguishing between upper and lower motor neuron bladder dysfunction in the early stages of dysautonomia. Gastrointestinal autonomic dysfunction typically presents as severe constipation. Diarrhea may develop (typically in diabetes mellitus) due to rapid transit of contents or uncoordinated small-bowel motor activity, or on an osmotic basis from bacterial overgrowth associated with small-bowel stasis. Impaired glandular secretory function may cause difficulty with food intake due to decreased salivation or eye irritation due to decreased lacrimation. Occasionally, temperature elevation and vasodilation can result from anhidrosis because sweating is normally important for heat dissipation (Chap. 23). Lack of sweating after a hot bath, during exercise, or on a hot day can suggest sudomotor failure.

OH (also called *orthostatic or postural hypotension*) is perhaps the most disabling feature of autonomic dysfunction. The prevalence of OH is relatively high, especially when OH associated with aging and diabetes mellitus is included (Table 454-4). OH can cause a variety of symptoms, including dimming or loss of vision, lightheadedness, diaphoresis, diminished hearing, pallor, and weakness. Syncope results when the drop in BP impairs cerebral perfusion. Other manifestations of impaired baroreflexes are supine hypertension, a heart rate that is fixed regardless of posture, postprandial hypotension, and an excessively high nocturnal BP. Many patients with OH have a preceding

TABLE 454-4 PREVALENCE OF ORTHOSTATIC HYPOTENSION IN DIFFERENT DISORDERS

Disorder	Prevalence
Aging	14–20%
Diabetic neuropathy	10%
Other autonomic neuropathies	10–50 per 100,000
Multiple system atrophy	5–15 per 100,000
Pure autonomic failure	10–30 per 100,000

diagnosis of hypertension or have concomitant supine hypertension, reflecting the great importance of baroreflexes in maintaining postural and supine normotension. The appearance of OH in patients receiving antihypertensive treatment may indicate overtreatment or the onset of an autonomic disorder. The most common causes of OH are not neurologic in origin; these must be distinguished from the neurogenic causes (Table 454-5). **Neurocardiogenic and cardiac causes of syncope are considered in Chap. 27.**

APPROACH TO THE PATIENT:

Orthostatic Hypotension and Other ANS Disorders

The first step in the evaluation of symptomatic OH is the exclusion of treatable causes. The history should include a review of medications that may affect the ANS (Table 454-6). The main classes of drugs that may cause OH are diuretics, antihypertensives, antidepressants, ethanol, narcotics, insulin, dopamine agonists, barbiturates, and calcium channel-blocking agents. However, the precipitation of OH by medications may also be the first sign of an underlying autonomic disorder. The history may reveal an underlying cause for symptoms (e.g., diabetes, Parkinson's disease) or specific underlying mechanisms (e.g., cardiac pump failure, reduced intravascular volume). The relationship of symptoms to meals (splanchnic pooling), standing on awakening in the morning (intravascular volume depletion), ambient warming (vasodilatation), or exercise (muscle arteriolar vasodilatation)

TABLE 454-5 NONNEUROGENIC CAUSES OF ORTHOSTATIC HYPOTENSION

Cardiac pump failure	Venous pooling
Myocardial infarction	Alcohol
Myocarditis	Postprandial dilation of splanchnic vessel beds
Constrictive pericarditis	Vigorous exercise with dilation of skeletal vessel beds
Aortic stenosis	Heat: hot environment, hot showers and baths, fever
Tachyarrhythmias	Prolonged recumbency or standing
Bradyarrhythmias	Sepsis
Salt-losing nephropathy	
Adrenal insufficiency	Medications
Diabetes insipidus	Antihypertensives
Venous obstruction	Diuretics
Reduced intravascular volume	Vasodilators: nitrates, hydralazine
Straining or heavy lifting, urination, defecation	Alpha- and beta-blocking agents
Dehydration	Central nervous system sedatives: barbiturates, opiates
Diarrhea, emesis	Tricyclic antidepressants
Hemorrhage	Phenothiazines
Burns	
Metabolic	
Adrenocortical insufficiency	
Hypoadosteronism	
Pheochromocytoma	
Severe potassium depletion	

TABLE 454-6 SOME DRUGS THAT AFFECT AUTONOMIC FUNCTION

Symptom	Drug Class	Specific Examples
Impotence	Opioids	Tylenol #3
	Anabolic steroids	—
	Some antiarrhythmics	Prazosin
	Some antihypertensives	Clonidine
	Some diuretics	Benazepril
Urinary retention	Some SSRIs	Venlafaxine
	Opioids	Fentanyl
Diaphoresis	Decongestants	Brompheniramine Diphenhydramine
	Some antihypertensives	Amlodipine
	Some SSRIs	Citalopram
Hypotension	Opioids	Morphine
	Tricyclics	Amitriptyline
	Beta blockers	Propranolol
	Diuretics	HCTZ
	CCBs	Verapamil

Abbreviations: CCBs, calcium channel blockers; HCTZ, hydrochlorothiazide; SSRIs, selective serotonin reuptake inhibitors.

should be sought. Standing time to first symptom and to presyncope (**Chap. 27**) should be followed for management.

Physical examination includes measurement of supine and standing pulse and BP. OH is defined as a sustained drop in systolic (≥ 20 mmHg) or diastolic (≥ 10 mmHg) BP after 2–3 min of standing. In nonneurogenic causes of OH (such as hypovolemia), the BP drop is accompanied by a compensatory increase in heart rate of >15 beats/min. A clue that the patient has neurogenic OH is the aggravation or precipitation of OH by autonomic stressors (a meal, hot bath, or exercise). Neurologic examination should include mental status (neurodegenerative disorders), cranial nerves (impaired downgaze with progressive supranuclear palsy; abnormal pupils with Horner's or Adie's syndrome), motor tone (Parkinson's disease and parkinsonism), and sensation (polyneuropathies). In patients without a clear diagnosis initially, follow-up evaluations every few months or whenever symptoms worsen may reveal the underlying cause.

Disorders of autonomic function should be considered in patients with symptoms of altered sweating (hyperhidrosis or hypohidrosis), gastroparesis (bloating, nausea, vomiting of old food), impotence, constipation, or bladder disturbances (urinary frequency, hesitancy, or incontinence).

AUTONOMIC TESTING

Autonomic function tests are helpful to document abnormalities when findings on history and examination are inconclusive; to detect subclinical involvement; or to follow the course of an autonomic disorder.

Heart Rate Variation with Deep Breathing This tests the parasympathetic component of cardiovascular reflexes via the vagus nerve. Results are influenced by multiple factors including the subject's position (recumbent, sitting, or standing), rate and depth of respiration (6 breaths per minute and a forced vital capacity [FVC] >1.5 L are optimal), age, medications, weight, and degree of hypocapnia. Interpretation of results requires comparison of test data with results from age-matched controls collected under identical test conditions. For example, the lower limit of normal heart rate variation with deep breathing in persons <20 years is >15 – 20 beats/min, but for persons over age 60 it is 5–8 beats/min. Heart rate variation with deep breathing (respiratory sinus arrhythmia) is abolished by the muscarinic ACh receptor antagonist atropine but is unaffected by sympathetic postganglionic blockade (e.g., propranolol).

Valsalva Response This response (**Table 454-7**) assesses the integrity of the baroreflex control of heart rate (parasympathetic) and BP (adrenergic). Under normal conditions, increases in BP at the carotid bulb trigger a reduction in heart rate (increased vagal tone), and decreases in BP trigger an increase in heart rate (reduced vagal tone). The Valsalva response is tested in the supine position. The subject exhales against a closed glottis (or into a manometer maintaining a constant expiratory pressure of 40 mmHg) for 15 s while measuring changes in heart rate and beat-to-beat BP. There are four phases of the BP and heart rate response to the Valsalva maneuver. Phases I and III are mechanical and related to changes in intrathoracic and intraabdominal pressure. In early phase II, reduced venous return results in a fall in stroke volume and BP, counteracted by a combination of reflex tachycardia and increased total peripheral resistance. Increased total peripheral resistance arrests the BP drop ~ 5 – 8 s after the onset of the maneuver. Late phase II begins with a progressive rise in BP toward or above baseline. Venous return and cardiac output return to normal in phase IV. Persistent peripheral arteriolar vasoconstriction and increased cardiac adrenergic tone result in a temporary BP overshoot and phase IV bradycardia (mediated by the baroreceptor reflex).

Autonomic function during the Valsalva maneuver can be measured using beat-to-beat BP or heart rate changes. The *Valsalva ratio* is defined as the maximum phase II tachycardia divided by the minimum phase IV bradycardia (**Table 454-8**). The ratio reflects the integrity of the entire baroreceptor reflex arc and of sympathetic efferents to blood vessels.

Sudomotor Function Sweating is induced by release of ACh from sympathetic postganglionic fibers. The quantitative sudomotor axon reflex test (QSART) is a measure of regional autonomic function mediated by ACh-induced sweating. A reduced or absent response indicates a lesion of the postganglionic sudomotor axon. For example, sweating may be reduced in the feet as a result of distal polyneuropathy (e.g., diabetes). The thermoregulatory sweat test (TST) is a qualitative measure of regional sweat production in response to an elevation of body temperature under controlled conditions. An indicator powder placed on the anterior surface of the body changes color with sweat production during temperature

TABLE 454-7 NORMAL BLOOD PRESSURE AND HEART RATE CHANGES DURING THE VALSALVA MANEUVER

Phase	Maneuver	Blood Pressure	Heart Rate	Comments
I	Forced expiration against a partially closed glottis	Rises; aortic compression from raised intrathoracic pressure	Decreases	Mechanical
II early	Continued expiration	Falls; decreased venous return to the heart	Increases (reflex tachycardia)	Reduced vagal tone
II late	Continued expiration	Rises; reflex increase in peripheral vascular resistance	Increases at slower rate	Requires intact efferent sympathetic response
III	End of expiration	Falls; increased capacitance of pulmonary bed	Increases further	Mechanical
IV	Recovery	Rises; persistent vasoconstriction and increased cardiac output	Compensatory bradycardia	Requires intact efferent sympathetic response

TABLE 454-8 NEURAL PATHWAYS UNDERLYING SOME STANDARDIZED AUTONOMIC TESTS

Test Evaluated	Procedure	Autonomic Function
HRDB	6 deep breaths/min	Cardiovascular function
Valsalva ratio	Expiratory pressure, 40 mmHg for 10–15 s	Cardiovascular function
QSART	Axon-reflex test 4 limb sites	Postganglionic sudomotor function
BP _{bb} to VM	BP _{bb} response to VM	Adrenergic function: baroreflex adrenergic control of vagal and vasomotor function
HUT	BP _{bb} and heart rate response to HUT	Adrenergic and cardiovascular responses to HUT

Abbreviations: BP_{bb}, beat-to-beat blood pressure; HRDB, heart rate response to deep breathing; HUT, head-up tilt; QSART, quantitative sudomotor axon reflex test; VM, Valsalva maneuver.

elevation. The pattern of color change is a measure of regional sweat secretion. A postganglionic lesion is present if both QSART and TST show absent sweating. In a preganglionic lesion, the QSART is normal but TST shows anhidrosis.

Orthostatic BP Recordings Beat-to-beat BP measurements determined in supine, 70° tilt, and tilt-back positions are useful to quantitate orthostatic failure of BP control. Allow a 20-min period of rest in the supine position before assessing changes in BP during tilting. The BP change combined with heart rate monitoring is useful for the evaluation of patients with suspected OH or unexplained syncope.

Tilt Table Testing for Syncope The great majority of patients with syncope do not have autonomic failure. Tilt table testing can be used to make the diagnosis of vasovagal syncope with sensitivity, specificity, and reproducibility. A standardized protocol is used that specifies the tilt apparatus, angle and duration of tilt, and procedure for provocation of vasodilation (e.g., sublingual or spray nitroglycerin). A positive nitroglycerin-stimulated test predicts recurrence of syncope. Recommendations for the performance of tilt studies for syncope have been incorporated in consensus guidelines.

SPECIFIC SYNDROMES OF ANS DYSFUNCTION

MULTIPLE SYSTEM ATROPHY (CHAP. 449)

Multiple system atrophy (MSA) is an entity that comprises autonomic failure (OH or a neurogenic bladder) and either parkinsonism (MSA-p) or a cerebellar syndrome (MSA-c). MSA-p is the more common form; the parkinsonism is atypical in that it is usually unassociated with significant tremor or a response to levodopa. Symptomatic OH within 1 year of onset of parkinsonism predicts eventual development of MSA-p in 75% of patients. There is a very high frequency of impotence in men. Although autonomic abnormalities are common in advanced Parkinson's disease (Chap. 449), the severity and distribution of autonomic failure are more severe and generalized in MSA. Brain magnetic resonance imaging (MRI) is a useful diagnostic adjunct: in MSA-p, iron deposition in the striatum may be evident as T2 hypointensity, and in MSA-c, cerebellar atrophy is present with a characteristic T2 hyperintense signal ("hot cross buns sign") in the pons (Fig. 454-2). Cardiac postganglionic adrenergic innervation, measured by uptake of fluorodopamine on positron emission tomography, is markedly impaired in the dysautonomia of Parkinson's disease (PD) but is usually normal in MSA. Neuropathologic changes include neuronal loss and gliosis in many CNS regions, including the brainstem, cerebellum, striatum, and intermediolateral cell column of the thoracolumbar spinal cord.

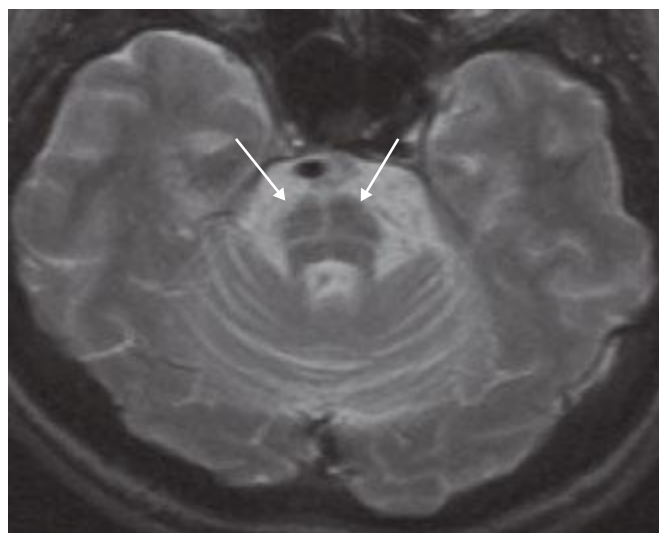


FIGURE 454-2 Multiple system atrophy, cerebellar type (MSA-c).

Axial T2-weighted magnetic resonance image at the level of the pons shows a characteristic hyperintense signal, the "hot cross buns" sign. This appearance can also be seen in some spinocerebellar atrophies, as well as other neurodegenerative conditions affecting the brainstem.

MSA is uncommon, with a prevalence estimated at 2–5 per 100,000 individuals. Onset is typically in the mid-fifties, men are slightly more often affected than women, and most cases are sporadic. The diagnosis should be considered in adults over the age of 30 years who present with OH or urinary incontinence and either parkinsonism that is poorly responsive to dopamine replacement or a cerebellar syndrome. MSA generally progresses relentlessly to death 7–10 years after onset, but survival beyond 15 years has been reported. Factors that predict a worse prognosis include rapid progression of disability, bladder dysfunction, female gender, the MSA-p subtype, and an older age at onset. Attempts to slow the progression of MSA have thus far been unsuccessful, including trials of lithium, growth hormone, riluzole, rasagiline, minocycline, and a recent trial of rifampicin.

Management is symptomatic for neurogenic OH (see below), sleep disorders including laryngeal stridor, and gastrointestinal (GI) and urinary dysfunction. GI management includes frequent small meals, soft diet, stool softeners, and bulk agents. Gastroparesis is difficult to treat; metoclopramide stimulates gastric emptying but worsens parkinsonism by blocking central dopamine receptors. The peripheral dopamine (D₂ and D₃) receptor antagonist domperidone has been used in patients with various GI conditions in many countries and is now available in the United States through the U.S. Food and Drug Administration's (FDA) Expanded Access to Investigational Drugs program.

Autonomic dysfunction is also a common feature in dementia with Lewy bodies (Chap. 448); the severity is usually less than that found in MSA or PD. In multiple sclerosis (MS; Chap. 458), autonomic complications reflect the CNS location of MS involvement and generally worsen with disease duration and disability.

SPINAL CORD

Spinal cord lesions from any cause may result in focal autonomic deficits or autonomic hyperreflexia (e.g., spinal cord transection or hemisection) affecting bowel, bladder, sexual, temperature-regulation, or cardiovascular functions. Quadriparetic patients exhibit both supine hypertension and OH after upward tilting. *Autonomic dysreflexia* describes a dramatic increase in BP in patients with traumatic spinal cord lesions above the T6 level, often in response to stimulation of the bladder, skin, or muscles. A distended or obstructed bladder, suprapubic palpation, catheter insertion, and urinary infection are common triggers. Associated symptoms can include facial flushing, headache, hypertension, or piloerection. Potential complications

include intracranial vasospasm or hemorrhage, cardiac arrhythmia, and death. Awareness of the syndrome, identifying the trigger, and careful monitoring of BP during procedures in patients with acute or chronic spinal cord injury are essential. In patients with supine hypertension, BP can be lowered by tilting the head upward or sitting the patient up. Vasodilator drugs may be used to treat acute elevations in BP. Clonidine can be used prophylactically to reduce the hypertension resulting from bladder stimulation. Dangerous increases or decreases in body temperature may result from an inability to experience the sensory accompaniments of heat or cold exposure or control peripheral vasoconstriction or sweating below the level of the spinal cord injury.

PERIPHERAL NERVE AND NEUROMUSCULAR JUNCTION DISORDERS

Peripheral neuropathies (**Chap. 459**) are the most common cause of chronic autonomic insufficiency. Polyneuropathies that affect small myelinated and unmyelinated fibers of the sympathetic and parasympathetic nerves commonly occur in diabetes mellitus, amyloidosis, chronic alcoholism, porphyria, and Guillain-Barré syndrome. Neuromuscular junction disorders with autonomic involvement include botulism and Lambert-Eaton syndrome (**Chap. 461**).

Diabetes Mellitus Autonomic neuropathy in patients with diabetes increases the mortality rate 1.5- to 3-fold, even after adjusting for other cardiovascular risk factors. Estimates of 5-year mortality risk among these patients range from 15 to 53%. Although many deaths are due to secondary vascular disease, there are patients who specifically suffer cardiac arrest due to autonomic neuropathy. The autonomic involvement is also predictive of other complications including renal disease, stroke, and sleep apnea. **Diabetes mellitus is discussed in Chaps. 417–419.**

Amyloidosis Autonomic neuropathy occurs in both sporadic and familial forms of amyloidosis (**Chap. 137**). The AL (immunoglobulin light chain) type is associated with primary amyloidosis or amyloidosis secondary to multiple myeloma. The ATTR type, with transthyretin as the primary protein component, is responsible for the most common form of inherited amyloidosis. Although patients usually present with a distal painful polyneuropathy accompanied by sensory loss, autonomic insufficiency can precede the development of the polyneuropathy or occur in isolation. The diagnosis can be made by protein electrophoresis of blood and urine, tissue biopsy (abdominal fat pad, rectal mucosa, or sural nerve) to search for amyloid deposits, and genetic testing for transthyretin mutations in familial cases. Treatment of familial cases with liver transplantation can be successful. The response of primary amyloidosis to melphalan and stem cell transplantation has been mixed. Death is usually due to cardiac or renal involvement. Postmortem studies reveal amyloid deposition in many organs, including two sites that contribute to autonomic failure: intraneural blood vessels and autonomic ganglia. Pathologic examination reveals a loss of both unmyelinated and myelinated nerve fibers.

Alcoholic Neuropathy Abnormalities in parasympathetic vagal and efferent sympathetic function are usually mild in alcoholic polyneuropathy. OH is usually due to brainstem involvement, rather than injury to the PNS. Impotence is a major problem, but concurrent gonadal hormone abnormalities may play a role in this symptom. Clinical symptoms of autonomic failure generally appear only when the stocking-glove polyneuropathy is severe, and there is usually coexisting Wernicke's encephalopathy (**Chap. 330**). Autonomic involvement may contribute to the high mortality rates associated with alcoholism (**Chap. 467**).

Porphyria (Chap. 430) Autonomic dysfunction is most extensively documented in acute intermittent porphyria but can also occur with variegate porphyria and hereditary coproporphyrin. Autonomic symptoms include tachycardia, sweating, urinary retention, abdominal pain, nausea and vomiting, insomnia, hypertension, and (less commonly) hypotension. Another prominent symptom is anxiety. Abnormal autonomic function can occur both during acute attacks

and during remissions. Elevated catecholamine levels during acute attacks correlate with the degree of tachycardia and hypertension that is present.

Guillain-Barré Syndrome (Chap. 460) BP fluctuations and arrhythmias from autonomic instability can be severe. It is estimated that between 2 and 10% of patients with severe Guillain-Barré syndrome suffer fatal cardiovascular collapse. GI autonomic involvement, sphincter disturbances, abnormal sweating, and pupillary dysfunction can also occur. Demyelination has been described in the vagus and glossopharyngeal nerves, the sympathetic chain, and the white rami communicantes. Interestingly, the degree of autonomic involvement appears to be independent of the severity of motor or sensory neuropathy. Acute autonomic and sensory neuropathy is a variant that spares the motor system and presents with neurogenic OH and varying degrees of sensory loss. It is treated similarly to Guillain-Barré syndrome, but prognosis is less favorable, with persistent severe sensory deficits and variable degrees of OH in many patients.

Autoimmune Autonomic Ganglionopathy (AAG) This disorder presents with the subacute development of autonomic disturbances including OH, enteric neuropathy (gastroparesis, ileus, constipation/diarrhea), flaccid bladder, and cholinergic failure (e.g., loss of sweating, sicca complex, and a tonic pupil). A chronic form of AAG resembles pure autonomic failure (see below). Autoantibodies against the ganglionic ACh receptor (A_3 AChR), which are present in approximately half of patients, are considered diagnostic of AAG. Pathology shows preferential involvement of small unmyelinated nerve fibers, with sparing of larger myelinated ones. Onset of the neuropathy follows a viral infection in approximately half of cases. Up to one-third of untreated patients experience significant functional improvement over time. Immunotherapies that have been reported to be helpful include plasmapheresis, intravenous immune globulin, glucocorticoids, azathioprine, rituximab, and mycophenolate mofetil. OH, gastroparesis, and sicca symptoms can be managed symptomatically.

AAG can also occur on a paraneoplastic basis, with adenocarcinoma or small-cell carcinoma of the lung, lymphoma, or thymoma being the most common (**Chap. 122**). In the paraneoplastic cases, distinctive additional features, such as cerebellar involvement or dementia, may be present (see **Tables 122-1, 122-2, and 122-3**). The neoplasm may be occult and possibly suppressed by the autoantibody.

Botulism Botulinum toxin binds presynaptically to cholinergic nerve terminals and, after uptake into the cytosol, blocks ACh release. This acute cholinergic neuropathy presents as motor paralysis and autonomic disturbances that include blurred vision, dry mouth, nausea, unreactive or sluggishly reactive pupils, constipation, and urinary retention (**Chap. 178**).

PURE AUTONOMIC FAILURE (PAF)

This sporadic syndrome consists of postural hypotension, impotence, bladder dysfunction, and impaired sweating. The disorder begins in midlife and occurs in women more often than men. The symptoms can be disabling, but the disease does not shorten life span. The clinical and pharmacologic characteristics suggest primary involvement of postganglionic sympathetic neurons. A severe reduction in the density of neurons within sympathetic ganglia results in low supine plasma NE levels and noradrenergic supersensitivity. Some patients who are initially labeled with this diagnosis subsequently go on to develop AAG or MSA. Skin biopsies can demonstrate phosphorylated α -synuclein inclusions in postganglionic sympathetic adrenergic and cholinergic nerve fibers from some individuals with PAF, distinguishing them from AAG and suggesting that PAF is a synucleinopathy; patients with PD also have α -synuclein inclusions in sympathetic nerve biopsies.

POSTURAL ORTHOSTATIC TACHYCARDIA SYNDROME (POTS)

This syndrome is characterized by symptomatic orthostatic intolerance without OH, accompanied by either an increase in heart rate to >120 beats/min or an increase of 30 beats/min with standing that subsides on sitting or lying down. Women are affected approximately five

times more often than men, and most develop the syndrome between the ages of 15 and 50. Presyncopal symptoms (lightheadedness, weakness, blurred vision) combined with symptoms of autonomic overactivity (palpitations, tremulousness, nausea) are common. Recurrent unexplained episodes of dysautonomia and fatigue also occur. The pathogenesis is unclear, but there is increasing evidence for sympathetic denervation distally in the legs with preserved cardiovascular function. Hypovolemia, venous pooling, impaired brainstem regulation, or increased sympathetic activity may play a role. Optimal treatment is uncertain, but expansion of fluid volume with water, salt, and fludrocortisone can be helpful as initial interventions. If this approach is inadequate, then midodrine, pyridostigmine, phenobarbital, beta blockers, or clonidine can be tried. Reconditioning and a sustained exercise program are important adjuncts to treatment.

INHERITED DISORDERS

There are five known hereditary sensory and autonomic neuropathies (HSAN I–V). The most important autonomic variants are HSAN I and HSAN III. HSAN I is dominantly inherited and often presents as a distal small-fiber neuropathy (burning feet syndrome) associated with sensory loss and foot ulcers. The most common responsible gene, on chromosome 9q, is *SPTLC1*. *SPTLC1* is an important enzyme in the regulation of ceramide. Cells from HSAN I patients with the mutation produce higher-than-normal levels of glucosyl ceramide, perhaps triggering apoptosis. HSAN III (Riley-Day syndrome; familial dysautonomia) is an autosomal recessive disorder of Ashkenazi Jewish children and adults and is much less prevalent than HSAN I. Decreased tearing, hyperhidrosis, reduced sensitivity to pain, areflexia, absent fungiform papillae on the tongue, and labile BP may be present. Episodic abdominal crises and fever are common. Pathologic examination of nerves reveals a loss of sympathetic, parasympathetic, and sensory neurons. The defective gene, *IKBKAP*, may prevent normal transcription of important molecules in neural development.

PRIMARY HYPERHIDROSIS

This syndrome presents with excess sweating of the palms of the hands and soles of the feet beginning in childhood or early adulthood. The condition tends to improve with age. The disorder affects 0.6–1.0% of the population. The etiology is unclear, but there may be a genetic component because 25% of patients have a positive family history. The condition can be socially embarrassing (e.g., shaking hands) or even disabling (e.g., inability to write without soiling the paper). Topical antiperspirants are occasionally helpful. More useful are potent anticholinergic drugs such as glycopyrrolate (1–2 mg PO tid). T2 ganglionectomy or sympathectomy is successful in >90% of patients with palmar hyperhidrosis. The advent of endoscopic transaxillary T2 sympathectomy has lowered the complication rate of the procedure. The most common postoperative complication is compensatory hyperhidrosis, which improves spontaneously over months. Other potential complications include recurrent hyperhidrosis (16%), Horner's syndrome (<2%), gustatory sweating, wound infection, hemothorax, and intercostal neuralgia. Local injection of botulinum toxin has also been used to block cholinergic, postganglionic sympathetic fibers to sweat glands in patients with palmar hyperhidrosis. This approach is limited by the need for repetitive injections (the effect usually lasts 4 months before waning).

ACUTE SYMPATHETIC OVERACTIVITY SYNDROMES

The physician may be confronted occasionally with an acute state of sympathetic overactivity.

An *autonomic storm* is an acute state of sustained sympathetic surge that results in variable combinations of alterations in BP and heart rate, body temperature, respiration, and sweating. Causes of autonomic storm include brain and spinal cord injury, toxins and drugs, autonomic neuropathy, and chemodectomas (e.g., pheochromocytoma). Brain injury is the most common cause of autonomic storm and typically follows severe head trauma and postresuscitation encephalopathy anoxic-ischemic brain injury. Autonomic storm can also occur with other acute intracranial lesions such as hemorrhage,

cerebral infarction, rapidly expanding tumors, subarachnoid hemorrhage, hydrocephalus, or (less commonly) an acute spinal cord lesion. The most consistent setting is that of an acute intracranial catastrophe of sufficient size and rapidity to produce a massive catecholaminergic surge. The surge can cause seizures, neurogenic pulmonary edema, and myocardial injury. Manifestations include fever, tachycardia, hypertension, tachypnea, hyperhidrosis, pupillary dilatation, and flushing. Lesions of the afferent limb of the baroreflex can result in milder recurrent autonomic storms; many of these follow neck irradiation.

Drugs and toxins may also be responsible, including sympathomimetics such as phenylpropanolamine, cocaine, amphetamines, and tricyclic antidepressants; tetanus; and, less often, botulinum toxin. Cocaine, including “crack,” can cause a hypertensive state with CNS hyperstimulation. Tricyclic overdose, such as from amitriptyline, can cause flushing, hypertension, tachycardia, fever, mydriasis, anhidrosis, and a toxic psychosis. The hyperadrenergic state associated with Guillain-Barré syndrome can produce a moderate autonomic storm. Pheochromocytoma presents with a paroxysmal or sustained hyperadrenergic state, headache, hyperhidrosis, palpitations, anxiety, tremulousness, and hypertension. *Neuroleptic malignant syndrome* refers to a syndrome of muscle rigidity, hyperthermia, and hypertension in psychotic patients treated with phenothiazines (Chap. 449). Management of autonomic storm includes ruling out other causes of autonomic instability, including malignant hyperthermia, porphyria, and seizures. Sepsis and encephalitis need to be excluded with appropriate studies. An electroencephalogram (EEG) should be done to search for seizure activity; MRI of the brain and spine is often necessary. The patient should be managed in an intensive care unit. Management with morphine sulphate (10 mg every 4 h) and labetalol (100–200 mg twice daily) may be helpful. Supportive treatment may need to be maintained for several weeks. For chronic and milder autonomic storm, propranolol and/or clonidine can be effective.

MISCELLANEOUS

Other conditions associated with autonomic failure include infections, malignancy, poisoning (organophosphates), and aging. Disorders of the hypothalamus can affect autonomic function and produce abnormalities in temperature control, satiety, sexual function, and circadian rhythms (Chap. 403).

REFLEX SYMPATHETIC DYSTROPHY AND CAUSALGIA

The failure to identify a primary role of the ANS in the pathogenesis of these disorders has resulted in a change of nomenclature. The terms complex regional pain syndrome (CRPS) types I and II are now used in place of reflex sympathetic dystrophy (RSD) and causalgia.

CRPS type I is a regional pain syndrome that often develops after tissue injury and most commonly affects one limb. Examples of associated injury include minor shoulder or limb trauma, fractures, myocardial infarction, or stroke. *Allodynia* (the perception of a nonpainful stimulus as painful), *hyperpathia* (an exaggerated pain response to a painful stimulus), and spontaneous pain occur. The symptoms are unrelated to the severity of the initial trauma and are not confined to the distribution of a single peripheral nerve. CRPS type II is a regional pain syndrome that develops after injury to a specific peripheral nerve, often a major nerve trunk. Spontaneous pain initially develops within the territory of the affected nerve but eventually may spread outside the nerve distribution.

Pain (usually burning or electrical in quality) is the primary clinical feature of CRPS. Vasomotor dysfunction, sudomotor abnormalities, or focal edema may occur alone or in combination but must be present for diagnosis. Limb pain syndromes that do not meet these criteria are best classified as “limb pain—not otherwise specified.” In CRPS, localized sweating (increased resting sweat output) and changes in blood flow may produce temperature differences between affected and unaffected limbs.

CRPS type I (RSD) has been classically divided into three clinical phases. Phase I consists of pain and swelling in the distal extremity occurring within weeks to 3 months after the precipitating event. The

pain is diffuse, spontaneous, and either burning, throbbing, or aching in quality. The involved extremity is warm and edematous, and the joints are tender. Increased sweating and hair growth develop. In phase II (3–6 months after onset), thin, shiny, cool skin appears. After an additional 3–6 months (phase III), atrophy of the skin and subcutaneous tissue plus flexion contractures complete the clinical picture. Autonomic testing or bone scans are occasionally useful when the diagnosis is in doubt.

The natural history of typical CRPS may be more benign and more variable than previously recognized. A variety of surgical and medical treatments have been developed, with conflicting reports of efficacy. Clinical trials suggest that early mobilization with physical therapy or a brief course of glucocorticoids may be helpful for CRPS type I or II. Other medical treatments include the use of adrenergic blockers, nonsteroidal anti-inflammatory drugs, calcium channel blockers, phenytoin, opioids, and calcitonin. Stellate ganglion blockade is a commonly used invasive technique that often provides temporary pain relief, but the efficacy of repetitive blocks is uncertain.

TREATMENT AUTONOMIC FAILURE

Management of autonomic failure is aimed at specific treatment of the cause and alleviation of symptoms. Of particular importance is the removal of drugs or amelioration of underlying conditions that cause or aggravate the autonomic symptoms, especially in the elderly. For example, OH can be caused or aggravated by angiotensin-converting enzyme inhibitors, calcium channel-blocking agents, tricyclic antidepressants, levodopa, alcohol, or insulin. A summary of drugs that can cause OH by class, putative mechanism, and magnitude of the BP drop is shown in Table 454-6.

PATIENT EDUCATION

Only a minority of patients with OH require drug treatment. All patients should be taught the mechanisms of postural normotension (volume status, resistance and capacitance bed, autoregulation) and the nature of orthostatic stressors (time of day and the influence of meals, heat, standing, and exercise). Patients should learn to recognize orthostatic symptoms early (especially subtle cognitive symptoms, weakness, and fatigue) and to modify or avoid activities that provoke episodes. Other helpful measures may include keeping a BP log and dietary education (salt/fluids). Learning physical counter-manuevers that reduce standing OH and practicing postural and resistance training are helpful measures.

SYMPTOMATIC TREATMENT

Nonpharmacologic approaches are summarized in Table 454-9. Adequate intake of salt and fluids to produce a voiding volume between 1.5 and 2.5 L of urine (containing >170 meq/L of Na⁺) each 24 h is essential. Sleeping with the head of the bed elevated will minimize the effects of supine nocturnal hypertension. Prolonged recumbency should be avoided when possible. Patients are advised to sit with legs dangling over the edge of the bed for several minutes before attempting to stand in the morning; other postural stresses should be similarly approached in a gradual manner. One

maneuver that can reduce OH is leg-crossing with maintained contraction of leg muscles for 30 s; this compresses leg veins and increases systemic resistance. Compressive garments, such as compression stockings or abdominal binders, are helpful on occasion but uncomfortable for many patients. For transient worsening of OH, drinking two 250-mL (8-oz) glasses of water can raise standing BP 20–30 mmHg for about 2 h, beginning ~20 min after the fluid load. The patient can increase intake of salt and fluids (bouillon treatment), increase use of physical counter-manuevers (elevate the legs when supine), or temporarily resort to a full-body stocking (compression pressure 30–40 mmHg).

Anemia should be corrected with erythropoietin, administered subcutaneously at doses of 25–75 U/kg three times per week. The hematocrit increases after 2–6 weeks. A weekly maintenance dose is usually necessary. However, the increased intravascular volume that accompanies the rise in hematocrit can exacerbate supine hypertension.

If these measures are not sufficient, pharmacologic treatment may be necessary. Midodrine, a directly acting α_1 -agonist that does not cross the blood-brain barrier, is effective. It has a duration of action of 2–4 h. The usual dose is 5–10 mg orally tid, but some patients respond best to a decremental dose (e.g., 15 mg on awakening, 10 mg at noon, and 5 mg in the afternoon). Midodrine should not be taken after 6:00 P.M. Side effects include pruritus, uncomfortable piloerection, and supine hypertension especially at higher doses. Droxidopa (Northera) was recently approved by the FDA for treatment of neurogenic OH associated with PAF, PD, or MSA; oral droxidopa is converted to NE and in short-term clinical trials was effective in decreasing symptoms of OH. Pyridostigmine (Mestinon) appears to improve OH without aggravating supine hypertension by enhancing ganglionic transmission (maximal when orthostatic, minimal when supine). Fludrocortisone will reduce OH but aggravates supine hypertension. At doses between 0.1 mg/d and 0.3 mg bid orally, it enhances renal sodium conservation and increases the sensitivity of arterioles to NE. Susceptible patients may develop fluid overload, congestive heart failure, supine hypertension, or hypokalemia. Potassium supplements are often necessary with chronic administration of fludrocortisone. Sustained elevations of supine BP >180/110 mmHg should be avoided. Supine hypertension (>180/110 mmHg) can be self-treated by avoiding the supine position (e.g., sleeping in a recumbent chair) and reducing fludrocortisone. A daily glass of wine, if requested by the patient, can be taken shortly before bedtime. If these simple measures are not adequate, drugs to be considered include oral hydralazine (25 mg qhs), oral nifedipine (Procardia; 10 mg qhs), or a nitroglycerin patch.

A promising drug combination (atomoxetine and yohimbine) has been studied for use in human subjects with severe OH not responsive to other agents, as can occur in some patients with diabetes and severe autonomic neuropathy not responsive to other medications. The atomoxetine blocks the NE reuptake transporter, and yohimbine blocks α_2 receptors that mediate the sympathetic feedback loop for downregulation of BP in response to atomoxetine. The result is a dramatic increase in BP and standing tolerance. This combination is not FDA approved for this purpose. It is possible that the limited drug duration of action can be used to withdraw drug treatment when the patient anticipates becoming supine (e.g., before sleep).

Postprandial OH may respond to several measures. Frequent, small, low-carbohydrate meals may diminish splanchnic shunting of blood after meals and reduce postprandial OH. Prostaglandin inhibitors (ibuprofen or indomethacin) taken with meals or midodrine (10 mg with the meal) can be helpful. The somatostatin analogue octreotide can be useful in the treatment of postprandial syncope by inhibiting the release of GI peptides that have vasodilator and hypotensive effects. The subcutaneous dose ranges from 25 μ g bid to 200 μ g tid.

TABLE 454-9 INITIAL TREATMENT OF ORTHOSTATIC HYPOTENSION (OH)

Patient education: mechanisms and stressors of OH
High-salt diet (10–20 g/d)
High-fluid intake (2 L/d)
Elevate head of bed 10 cm (4 in.) to minimize supine hypertension
Maintain postural stimuli
Learn physical counter-manuevers
Compression garments
Correct anemia

455 Trigeminal Neuralgia, Bell's Palsy, and Other Cranial Nerve Disorders

M. Flint Beal, Stephen L. Hauser

Symptoms and signs of cranial nerve pathology are common in internal medicine. They often develop in the context of a widespread neurologic disturbance, and in such situations, cranial nerve involvement may represent the initial manifestation of the illness. In other disorders, involvement is largely restricted to one or several cranial nerves; these distinctive disorders are reviewed in this chapter. **Disorders of ocular movement are discussed in Chap. 39, disorders of hearing in Chap. 43, and vertigo and disorders of vestibular function in Chap. 28.**

FACIAL PAIN OR NUMBNESS

ANATOMIC CONSIDERATIONS

The trigeminal (fifth cranial) nerve supplies sensation to the skin of the face and anterior half of the head (Fig. 455-1). The motor part innervates the muscles involved in chewing (including masseters and pterygoids) as well as the tensor tympani of the middle ear (hearing especially for high-pitched tones). It is the largest of the cranial nerves. It exits in the lateral midpons and traverses the middle cranial fossa to the semilunar (gasserian, trigeminal) ganglion in Meckel's cave, where the nerve divides into three divisions (ophthalmic [V1], maxillary [V2], and mandibular [V3]). V1 and V2 traverse the cavernous sinus to exit in the superior orbital fissure and foramen rotundum, located above and below the eye socket respectively; V3 exits through the foramen ovale. The trigeminal nerve is predominantly sensory, and motor innervation is exclusively carried in V3. The cornea is primarily innervated by V1, although an inferior crescent may be V2. Upon

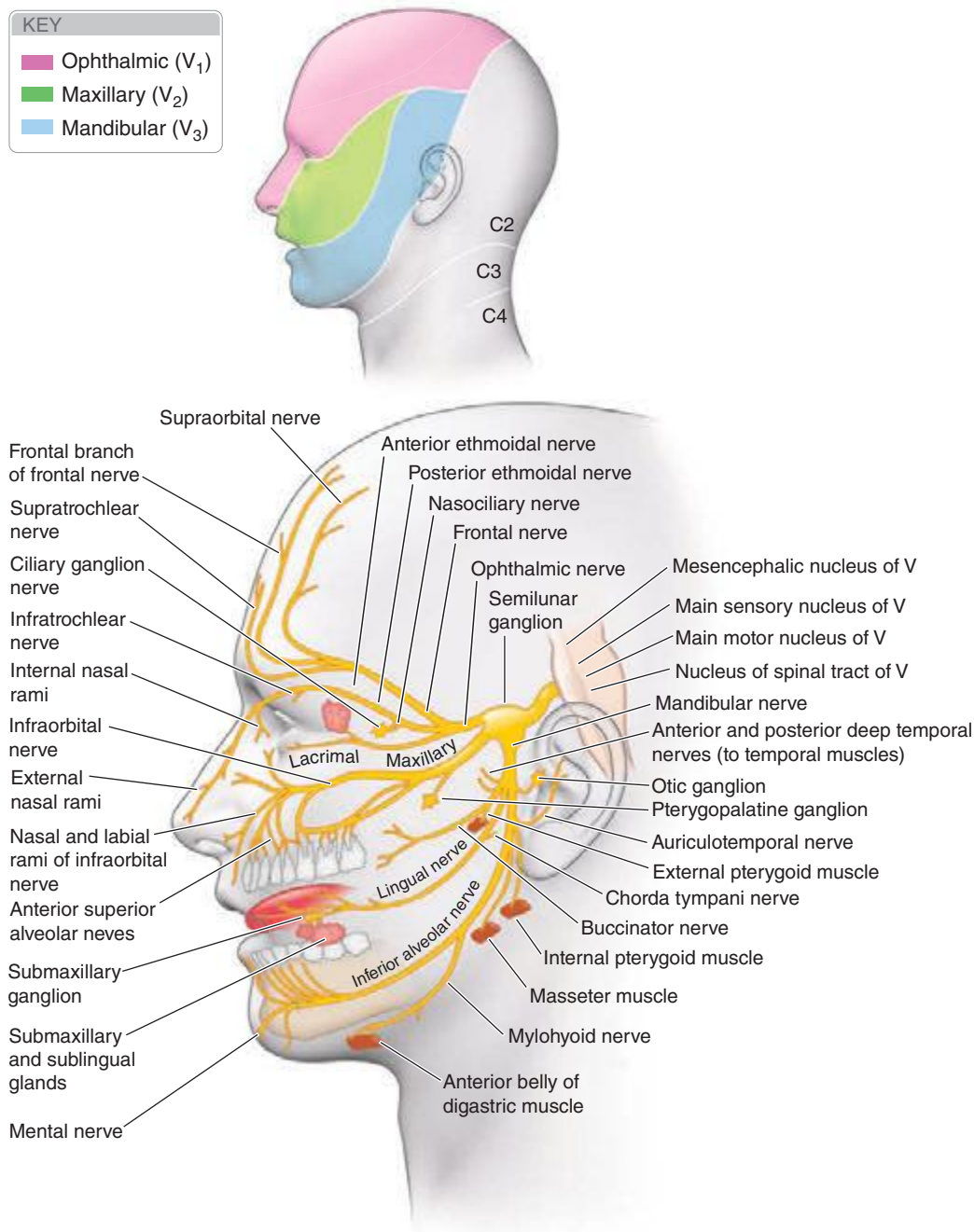


FIGURE 455-1 The trigeminal nerve and its branches and sensory distribution on the face. The three major sensory divisions of the trigeminal nerve consist of the ophthalmic, maxillary, and mandibular nerves. (Adapted from Waxman SG: *Clinical Neuroanatomy*, 26th ed. <http://www.accessmedicine.com>. Copyright © The McGraw-Hill Companies, Inc. All rights reserved.)

entering the pons, pain and temperature fibers descend ipsilaterally to the upper cervical spinal cord as the spinal tract of V, before synapsing with the spinal nucleus of V; this accounts for the facial numbness that can occur with spinal cord lesions above C2. In the brainstem, the spinal tract of V is also located adjacent to crossed ascending fibers of the spinothalamic tract, producing a “crossed” sensory loss for pain and temperature (ipsilateral face, contralateral arm/trunk/leg) with lesions of the lateral lower brainstem. CN V is also ensheathed by oligodendrocyte-derived, rather than Schwann cell–derived, myelin for up to 7 mm after it leaves the brainstem, unlike just a few millimeters for other cranial and spinal nerves; this may explain the high frequency of trigeminal neuralgia in multiple sclerosis (**Chap. 458**), a disorder of oligodendrocyte myelin.

TRIGEMINAL NEURALGIA (TIC DOULOUREUX)

Clinical Manifestations Trigeminal neuralgia is characterized by excruciating paroxysms of pain in the lips, gums, cheek, or chin and, very rarely, in the distribution of the ophthalmic division of the fifth nerve. The pain seldom lasts more than a few seconds or a minute or two but may be so intense that the patient winces, hence the term tic. The paroxysms, experienced as single jabs or clusters, tend to recur frequently, both day and night, for several weeks at a time. They may occur spontaneously or with movements of affected areas evoked by speaking, chewing, or smiling. Another characteristic feature is the presence of trigger zones, typically on the face, lips, or tongue, that provoke attacks; patients may report that tactile stimuli—e.g., washing the face, brushing the teeth, or exposure to a draft of air—generate excruciating pain. An essential feature of trigeminal neuralgia is that objective signs of sensory loss cannot be demonstrated on examination.

Trigeminal neuralgia is relatively common, with an estimated annual incidence of 4–8 per 100,000 individuals. Middle-aged and elderly persons are affected primarily, and ~60% of cases occur in women. Onset is typically sudden, and bouts tend to persist for weeks or months before remitting spontaneously. Remissions may be long-lasting, but in most patients, the disorder ultimately recurs.

Pathophysiology Symptoms result from ectopic generation of action potentials in pain-sensitive afferent fibers of the fifth cranial nerve root just before it enters the lateral surface of the pons. Compression or other pathology in the nerve leads to demyelination of large myelinated fibers that do not themselves carry pain sensation but become hyperexcitable and electrically coupled with smaller unmyelinated or poorly myelinated pain fibers in close proximity; this may explain why tactile stimuli, conveyed via the large myelinated fibers, can stimulate paroxysms of pain. Compression of the trigeminal nerve root by a blood vessel, most often the superior cerebellar artery or on occasion a tortuous vein, is now believed to be the source of trigeminal neuralgia in most patients. In cases of vascular compression, age-related brain sagging and increased vascular thickness and tortuosity may explain the prevalence of trigeminal neuralgia in later life.

Differential Diagnosis Trigeminal neuralgia must be distinguished from other causes of face and head pain (**Chap. 21**) and from pain arising from diseases of the jaw, teeth, or sinuses. Pain from migraine or cluster headache tends to be deep-seated and steady, unlike the superficial stabbing quality of trigeminal neuralgia; rarely, cluster headache is associated with trigeminal neuralgia, a syndrome known as *cluster-tic*. In temporal arteritis, superficial facial pain is present but is not typically shock-like, the patient frequently complains of myalgias and other systemic symptoms, and an elevated erythrocyte sedimentation rate (ESR) is usually present (**Chap. 385**). When trigeminal neuralgia develops in a young adult or is bilateral, multiple sclerosis (MS) is a key consideration, and in such cases, the cause is a demyelinating plaque near the root entry zone of the fifth nerve in the pons; often, evidence of facial sensory loss can be found on careful examination. Cases that are secondary to mass lesions—such as aneurysms, neurofibromas, acoustic schwannomas, or meningiomas—usually produce objective signs of sensory loss in the trigeminal nerve distribution (trigeminal neuropathy, see below).

Laboratory Evaluation An ESR is indicated if temporal arteritis is suspected. In typical cases of trigeminal neuralgia, neuroimaging studies

are usually unnecessary but may be valuable if MS is a consideration or in assessing overlying vascular lesions in order to plan for decompression surgery.

TREATMENT TRIGEMINAL NEURALGIA

Drug therapy with carbamazepine is effective in ~50–75% of patients. Carbamazepine should be started as a single daily dose of 100 mg taken with food and increased gradually (by 100 mg daily in divided doses every 1–2 days) until substantial (>50%) pain relief is achieved. Most patients require a maintenance dose of 200 mg qid. Doses >1200 mg daily provide no additional benefit. Dizziness, imbalance, sedation, and rare cases of agranulocytosis are the most important side effects of carbamazepine. If treatment is effective, it is usually continued for 1 month and then tapered as tolerated. Oxcarbazepine (300–1200 mg bid) is an alternative to carbamazepine that has less bone marrow toxicity and probably is equally efficacious. If these agents are not well tolerated or are ineffective, lamotrigine, 400 mg daily, and phenytoin, 300–400 mg daily, are other options. Baclofen may also be tried, either alone or in combination with an anticonvulsant. The initial dose is 5–10 mg tid, gradually increasing as needed to 20 mg qid.

If drug treatment fails, surgical therapy should be offered. The most widely used method is currently microvascular decompression to relieve pressure on the trigeminal nerve as it exits the pons. This procedure requires a suboccipital craniotomy. This procedure appears to have a >70% efficacy rate and a low rate of pain recurrence in responders; the response is better for classic tic-like symptoms than for nonlancinating facial pains. In a small number of cases, there is perioperative damage to the eighth or seventh cranial nerves or to the cerebellum or a postoperative cerebrospinal fluid leak syndrome. High-resolution magnetic resonance angiography is useful preoperatively to visualize the relationships between the fifth cranial nerve root and nearby blood vessels.

Gamma knife radiosurgery of the trigeminal nerve root is also used for treatment and results in complete pain relief, sometimes delayed in onset, in more than two-thirds of patients and a low risk of persistent facial numbness; the response is sometimes long-lasting, but recurrent pain develops over 2–3 years in half of patients. Compared with surgical decompression, gamma knife surgery appears to be somewhat less effective but has few serious complications.

Another procedure, radiofrequency thermal rhizotomy, creates a heat lesion of the trigeminal ganglion or nerve. It is used less often now than in the past. Short-term relief is experienced by >95% of patients; however, long-term studies indicate that pain recurs in up to one-third of treated patients. Postoperatively, partial numbness of the face is common, masseter (jaw) weakness may occur especially following bilateral procedures, and corneal denervation with secondary keratitis can follow rhizotomy for first-division trigeminal neuralgia.

TRIGEMINAL NEUROPATHY

A variety of diseases can affect the trigeminal nerve (**Table 455-1**). Most present with sensory loss on the face or with weakness of the jaw muscles. Deviation of the jaw on opening indicates weakness of the pterygoids on the side to which the jaw deviates. Some cases are due to Sjögren’s syndrome or a collagen-vascular disease such as systemic lupus erythematosus, scleroderma, or mixed connective tissue disease. Among infectious causes, herpes zoster (acute or postherpetic) and leprosy should be considered. Tumors of the middle cranial fossa (meningiomas), of the trigeminal nerve (schwannomas), or of the base of the skull (metastatic tumors) may cause a combination of motor and sensory signs. Lesions in the cavernous sinus can affect the first and second divisions of the trigeminal nerve, and lesions of the superior orbital fissure can affect the first (ophthalmic) division; the accompanying corneal anesthesia increases the risk of ulceration (neurokeratitis).

Isolated sensory loss over the chin (mental neuropathy) can be the only manifestation of systemic malignancy. Rarely, an idiopathic form of trigeminal neuropathy is observed. It is characterized by numbness

TABLE 455-1 TRIGEMINAL NERVE DISORDERS

Nuclear (brainstem) lesions	Peripheral nerve lesions
Multiple sclerosis	Nasopharyngeal carcinoma
Stroke	Trauma
Syringobulbia	Guillain-Barré syndrome
Glioma	Sjögren's syndrome
Lymphoma	Collagen-vascular diseases
Preganglionic lesions	Sarcoidosis
Acoustic neuroma	Leprosy
Meningioma	Drugs (stilbamidine, trichloroethylene)
Metastasis	Idiopathic trigeminal neuropathy
Chronic meningitis	
Cavernous carotid aneurysm	
Gasserian ganglion lesions	
Trigeminal neuroma	
Herpes zoster	
Infection (spread from otitis media or mastoiditis)	

and paresthesias, sometimes bilaterally, with loss of sensation in the territory of the trigeminal nerve but without weakness of the jaw. Gradual recovery is the rule. Tonic spasm of the masticatory muscles, known as trismus, is symptomatic of tetanus (**Chap. 177**) or may occur in patients treated with phenothiazines.

FACIAL WEAKNESS

ANATOMIC CONSIDERATIONS

(**Fig. 455-2**) The seventh cranial nerve supplies all the muscles concerned with facial expression. The sensory component is small (the nervus intermedius); it conveys taste sensation from the anterior

two-thirds of the tongue and probably cutaneous impulses from the anterior wall of the external auditory canal. The motor nucleus of the seventh nerve lies anterior and lateral to the abducens nucleus. After leaving the pons, the seventh nerve enters the internal auditory meatus with the acoustic nerve. The nerve continues its course in its own bony channel, the facial canal, and exits from the skull via the stylomastoid foramen. It then passes through the parotid gland and subdivides to supply the facial muscles.

A complete interruption of the facial nerve at the stylomastoid foramen paralyzes all muscles of facial expression. The corner of the mouth droops, the creases and skinfolds are effaced, the forehead is unfurrowed, and the eyelids will not close. Upon attempted closure of the lids, the eye on the paralyzed side rolls upward (Bell's phenomenon). The lower lid sags and falls away from the conjunctiva, permitting tears to spill over the cheek. Food collects between the teeth and lips, and saliva may dribble from the corner of the mouth. The patient complains of a heaviness or numbness in the face, but sensory loss is rarely demonstrable and taste is intact.

If the lesion is in the middle-ear portion, taste is lost over the anterior two-thirds of the tongue on the same side. If the nerve to the stapedius is interrupted, there is hyperacusis (sensitivity to loud sounds). Lesions in the internal auditory meatus may affect the adjacent auditory and vestibular nerves, causing deafness, tinnitus, or dizziness. Intrapontine lesions that paralyze the face usually affect the abducens nucleus as well, and often the corticospinal and sensory tracts.

If the peripheral facial paralysis has existed for some time and recovery of motor function is incomplete, a continuous diffuse contraction of facial muscles may appear. The palpebral fissure becomes narrowed, and the nasolabial fold deepens. Attempts to move one group of facial muscles may result in contraction of all (associated movements, or synkinesis). Facial spasms, initiated by movements of the face, may develop (hemifacial spasm). Anomalous regeneration of seventh nerve fibers may result in other troublesome phenomena. If fibers originally connected with the orbicularis oculi come to innervate the orbicularis oris, closure of the lids may cause a retraction of the mouth, or if fibers originally connected with muscles of the face later innervate the lacrimal gland, anomalous tearing ("crocodile tears") may occur with any activity of the facial muscles, such as eating. Another facial synkinesia is triggered by jaw opening, causing closure of the eyelids on the side of the facial palsy (jaw-winking).

BELL'S PALSY

The most common form of facial paralysis is Bell's palsy. The annual incidence of this idiopathic disorder is ~25 per 100,000 annually, or about 1 in 60 persons in a lifetime. Risk factors include pregnancy and diabetes mellitus.

Clinical Manifestations The onset of Bell's palsy is fairly abrupt, with maximal weakness being attained by 48 h as a general rule. Pain behind the ear may precede the paralysis for a day or two. Taste sensation may be lost unilaterally, and hyperacusis may be present. In some cases, there is mild cerebrospinal fluid lymphocytosis. Magnetic resonance imaging (MRI) may reveal swelling and uniform enhancement of the geniculate ganglion and facial nerve and, in some cases, entrapment of the swollen nerve in the temporal bone. Approximately 80% of patients recover within a few weeks or months. Electromyography may be of some prognostic value; evidence of denervation after 10 days indicates there has been axonal degeneration, that there will be a long delay (3 months as a rule) before regeneration occurs, and that it may be incomplete. The presence of incomplete paralysis in the first week is the most favorable prognostic sign. Recurrences are reported in approximately 7% of cases.

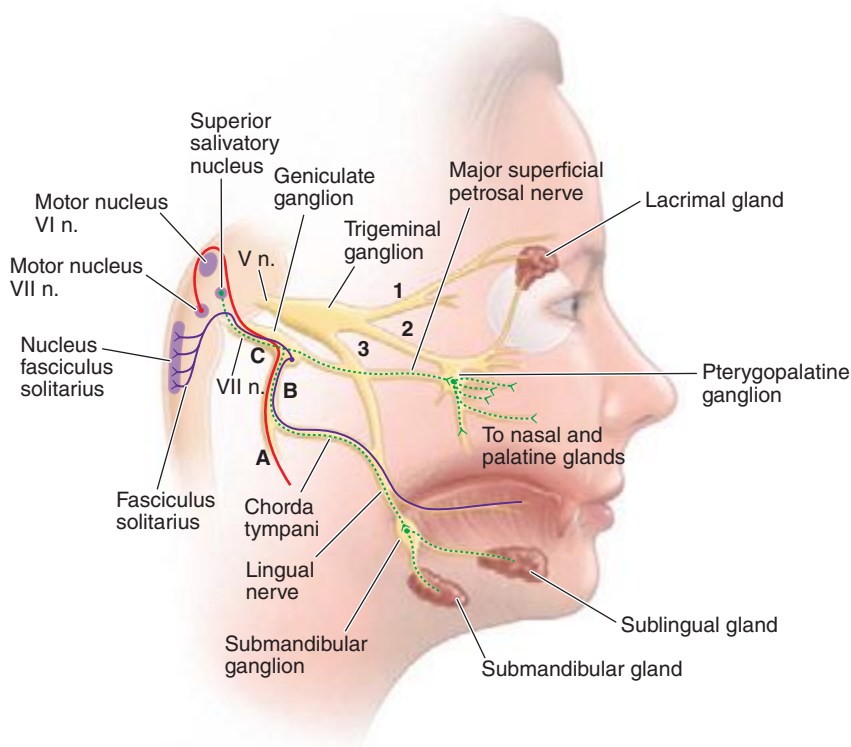


FIGURE 455-2 The facial nerve. A, B, and C denote lesions of the facial nerve at the stylomastoid foramen, distal and proximal to the geniculate ganglion, respectively. Green lines indicate the parasympathetic fibers, red line indicates motor fibers, and purple lines indicate visceral afferent fibers (taste). (Adapted from MB Carpenter: *Core Text of Neuroanatomy*, 2nd ed. Williams & Wilkins, 1978.)

2648 Pathophysiology In acute Bell's palsy, there is inflammation of the facial nerve with mononuclear cells, consistent with an infectious or immune cause. Herpes simplex virus (HSV) type 1 DNA was frequently detected in endoneurial fluid and posterior auricular muscle, suggesting that a reactivation of this virus in the geniculate ganglion may be responsible for most cases. Reactivation of varicella-zoster virus is associated with Bell's palsy in up to one-third of cases and may represent the second most frequent cause. A variety of other viruses have also been implicated less commonly. An increased incidence of Bell's palsy was also reported among recipients of inactivated intranasal influenza vaccine, and it was hypothesized that this could have resulted from the *Escherichia coli* enterotoxin used as adjuvant or reactivation of latent virus.

Differential Diagnosis There are many other causes of acute facial palsy that must be considered in the differential diagnosis of Bell's palsy. Lyme disease can cause unilateral or bilateral facial palsies; in endemic areas, 10% or more of cases of facial palsy are likely due to infection with *Borrelia burgdorferi* (Chap. 210). The Ramsay Hunt syndrome, caused by reactivation of herpes zoster in the geniculate ganglion, consists of a severe facial palsy associated with a vesicular eruption in the external auditory canal and sometimes in the pharynx and other parts of the cranial integument; often the eighth cranial nerve is affected as well. Facial palsy that is often bilateral occurs in sarcoidosis (Chap. 390) and in Guillain-Barré syndrome (Chap. 460). Leprosy frequently involves the facial nerve, and facial neuropathy may also occur in diabetes mellitus, connective tissue diseases including Sjögren's syndrome, and amyloidosis. The rare Melkersson-Rosenthal syndrome consists of recurrent facial paralysis; recurrent—and eventually permanent—facial (particularly labial) edema; and, less constantly, plication of the tongue. Its cause is unknown. Acoustic neuromas frequently involve the facial nerve by local compression. Infarcts, demyelinating lesions of MS, and tumors are the common pontine lesions that interrupt the facial nerve fibers; other signs of brainstem involvement are usually present. Tumors that invade the temporal bone (carotid body, cholesteatoma, dermoid) may produce a facial palsy, but the onset is insidious and the course progressive.

All these forms of nuclear or peripheral facial palsy must be distinguished from the supranuclear type. In the latter, the frontalis and orbicularis oculi muscles of the forehead are involved less than those of the lower part of the face, since the upper facial muscles are innervated by corticobulbar pathways from both motor cortices, whereas the lower facial muscles are innervated only by the opposite hemisphere. In supranuclear lesions, there may be a dissociation of emotional and voluntary facial movements, and often some degree of paralysis of the arm and leg or an aphasia (in dominant hemisphere lesions) is present.

Laboratory Evaluation The diagnosis of Bell's palsy can usually be made clinically in patients with (1) a typical presentation, (2) no risk factors or preexisting symptoms for other causes of facial paralysis, (3) absence of cutaneous lesions of herpes zoster in the external ear canal, and (4) a normal neurologic examination with the exception of the facial nerve. Particular attention to the eighth cranial nerve, which courses near to the facial nerve in the pontomedullary junction and in the temporal bone, and to other cranial nerves is essential. In atypical or uncertain cases, an ESR, testing for diabetes mellitus, a Lyme titer, angiotensin-converting enzyme and chest imaging studies for possible sarcoidosis, a lumbar puncture for possible Guillain-Barré syndrome, or MRI scanning may be indicated. MRI often shows swelling and enhancement of the facial nerve in idiopathic Bell's palsy (Fig. 455-3).

TREATMENT BELL'S PALSY

Symptomatic measures include (1) the use of paper tape to depress the upper eyelid during sleep and prevent corneal drying, and (2) massage of the weakened muscles. A course of glucocorticoids, given as prednisone 60–80 mg daily during the first 5 days and then tapered over the next 5 days, modestly shortens the recovery period and improves the functional outcome. Although large and well-controlled randomized trials found no added benefit of the antiviral agents valacyclovir (1000 mg daily for 5–7 days) or acyclovir (400 mg five times daily for 10 days) compared to glucocorticoids alone, some earlier data suggested that combination therapy with prednisone plus valacyclovir might be marginally better than prednisone alone, especially in patients with severe clinical presentations. For patients with permanent paralysis from Bell's palsy, a number of cosmetic surgical procedures have been used to restore a relatively symmetric appearance to the face.

OTHER MOTOR DISORDERS OF THE FACE

Hemifacial spasm consists of painless irregular involuntary contractions on one side of the face. Most cases appear related to vascular compression of the exiting facial nerve in the pons. Other cases develop as a sequela to Bell's palsy or are secondary to compression and/or demyelination of the nerve by tumor, infection, or MS. Mild cases can be treated with carbamazepine, gabapentin, or, if these drugs fail, baclofen. Local injections of botulinum toxin into affected muscles can relieve spasms for 3–4 months, and the injections can be repeated. Refractory cases due to vascular compression usually respond to surgical decompression of the facial nerve. Blepharospasm is an involuntary recurrent spasm of both eyelids that usually occurs in elderly persons as an isolated phenomenon or with varying degrees of spasm of other

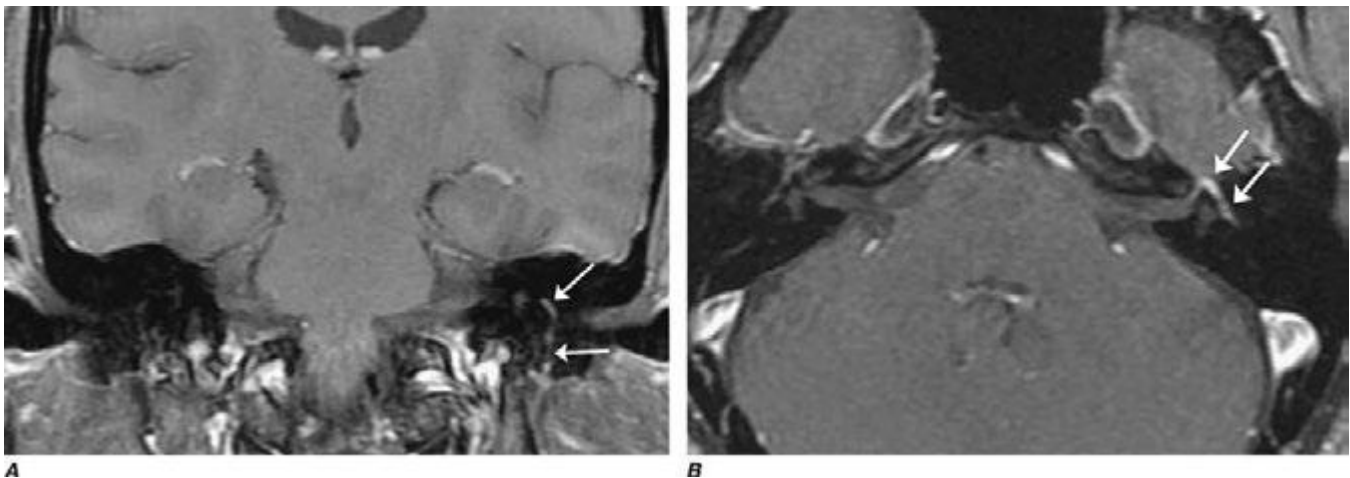


FIGURE 455-3 Axial and coronal T1-weighted images after gadolinium with fat suppression demonstrate diffuse smooth linear enhancement of the left facial nerve, involving the genu, tympanic, and mastoid segments within the temporal bone (arrows), without evidence of mass lesion. Although highly suggestive of Bell's palsy, similar findings may be seen with other etiologies such as Lyme disease, sarcoidosis, and perineural malignant spread.

facial muscles. Severe, persistent cases of blepharospasm can be treated by local injection of botulinum toxin into the orbicularis oculi. Facial myokymia refers to a fine rippling activity of the facial muscles; it may be caused by MS or follow Guillain-Barré syndrome (Chap. 460).

Facial hemiatrophy occurs mainly in women and is characterized by a disappearance of fat in the dermal and subcutaneous tissues on one side of the face. It usually begins in adolescence or the early adult years and is slowly progressive. In its advanced form, the affected side of the face is gaunt, and the skin is thin, wrinkled, and brown. The facial hair may turn white and fall out, and the sebaceous glands become atrophic. Bilateral involvement may occur. A limited form of systemic sclerosis (scleroderma) may be the cause of some cases. Treatment is cosmetic, consisting of transplantation of skin and subcutaneous fat.

OTHER CRANIAL NERVE DISORDERS

GLOSSOPHARYNGEAL NEURALGIA

This form of neuralgia involves the ninth (glossopharyngeal) and sometimes portions of the tenth (vagus) cranial nerves. It resembles trigeminal neuralgia in many respects but is much less common. The pain is intense and paroxysmal; it originates on one side of the throat, approximately in the tonsillar fossa. In some cases, the pain is localized in the ear or may radiate from the throat to the ear because of involvement of the tympanic branch of the glossopharyngeal nerve. Spasms of pain may be initiated by swallowing or coughing. There is no demonstrable motor or sensory deficit; the glossopharyngeal nerve supplies taste sensation to the posterior third of the tongue and, together with the vagus nerve, sensation to the posterior pharynx. Cardiac symptoms—bradycardia or asystole, hypotension, and fainting—have been reported. Glossopharyngeal neuralgia can result from vascular compression, MS, or tumors, but many cases are idiopathic. Medical therapy is similar to that for trigeminal neuralgia, and carbamazepine is generally the first choice. If drug therapy is unsuccessful, surgical procedures—including microvascular decompression if vascular compression is evident—or rhizotomy of glossopharyngeal and vagal fibers in the jugular bulb is frequently successful.

Glossopharyngeal neuropathy in conjunction with vagus and accessory nerve palsies may occur with herpes zoster infection or with a tumor or aneurysm in the posterior fossa or in the jugular foramen. Hoarseness due to vocal cord paralysis, some difficulty in swallowing, deviation of the soft palate to the intact side, anesthesia of the posterior wall of the pharynx, and weakness of the upper part of the trapezius and sternocleidomastoid muscles make up the jugular foramen syndrome (Table 455-2).

DYSPHAGIA AND DYSPHONIA

When the intracranial portion of one vagus (tenth cranial) nerve is interrupted, the soft palate droops ipsilaterally and does not rise in phonation. There is loss of the gag reflex on the affected side, as well as of the “curtain movement” of the lateral wall of the pharynx, whereby the faucial pillars move medially as the palate rises in saying “ah.” The voice is hoarse and slightly nasal, and the vocal cord lies immobile midway between abduction and adduction. Loss of sensation at the external auditory meatus and the posterior pinna may also be present.

The pharyngeal branches of both vagal nerves may be affected in diphtheria; the voice has a nasal quality, and regurgitation of liquids through the nose occurs during swallowing.

Injury to the vagus nerve in the carotid sheath can also occur with carotid dissection or following endarterectomy. The vagus nerve may be involved at the meningeal level by neoplastic and infectious processes and within the medulla by tumors, vascular lesions (e.g., the lateral medullary syndrome), and motor neuron disease. This nerve may be involved by infection with varicella zoster virus. Polymyositis and dermatomyositis, which cause hoarseness and dysphagia by direct involvement of laryngeal and pharyngeal muscles, may be confused with diseases of the vagus nerves. Dysphagia is also a symptom in some patients with myotonic dystrophy. **Nonneurologic causes of dysphagia are discussed in Chap. 53.**

The recurrent laryngeal nerves, especially the left, are most often damaged as a result of intrathoracic disease. Aneurysm of the aortic

TABLE 455-2 CRANIAL NERVE SYNDROMES

Site	Cranial Nerves	Usual Cause
Sphenoid fissure (superior orbital)	III, IV, first division V, VI	Invasive tumors of sphenoid bone; aneurysms
Lateral wall of cavernous sinus	III, IV, first division V, VI, often with VI	Infection, thrombosis, aneurysm, or fistula of cavernous sinus; invasive tumors from sinuses and sella turcica; benign granuloma responsive to glucocorticoids
Retrosphenoid space	II, III, IV, V, VI	Large tumors of middle cranial fossa
Apex of petrous bone	V, VI	Petrositis; tumors of petrous bone
Internal auditory meatus	VII, VIII	Tumors of petrous bone (dermoids, etc.); infectious processes; acoustic neuroma
Pontocerebellar angle	V, VII, VIII, and sometimes IX	Acoustic neuroma; meningioma
Jugular foramen	IX, X, XI	Tumors and aneurysms
Posterior laterocondylar space	IX, X, XI, XII	Tumors of parotid gland and carotid body and metastatic tumors
Posterior retroparotid space	IX, X, XI, XII, and Horner's syndrome	Tumors of parotid gland, carotid body, lymph nodes; metastatic tumor; tuberculous adenitis

arch, an enlarged left atrium, and tumors of the mediastinum and bronchi are much more frequent causes of an isolated vocal cord palsy than are intracranial disorders. However, a substantial number of cases of recurrent laryngeal palsy remain idiopathic.

When confronted with a case of laryngeal palsy, the physician must attempt to determine the site of the lesion. If it is intramedullary, there are usually other signs, such as ipsilateral cerebellar dysfunction, loss of pain and temperature sensation over the ipsilateral face and contralateral arm and leg, and an ipsilateral Horner's syndrome. If the lesion is extramedullary, the glossopharyngeal and spinal accessory nerves are frequently involved (jugular foramen syndrome). If it is extracranial in the posterior laterocondylar or retroparotid space, there may be a combination of ninth, tenth, eleventh, and twelfth cranial nerve palsies and a Horner's syndrome (Table 455-2). If there is no sensory loss over the palate and pharynx and no palatal weakness or dysphagia, the lesion is below the origin of the pharyngeal branches, which leave the vagus nerve high in the cervical region; the usual site of disease is then the mediastinum.

NECK WEAKNESS

Isolated involvement of the accessory (eleventh cranial) nerve can occur anywhere along its route, resulting in partial or complete paralysis of the sternocleidomastoid and trapezius muscles. More commonly, involvement occurs in combination with deficits of the ninth and tenth cranial nerves in the jugular foramen or after exit from the skull (Table 455-2). An idiopathic form of accessory neuropathy, akin to Bell's palsy, has been described, and it may be recurrent in some cases. Most but not all patients recover.

TONGUE PARALYSIS

The hypoglossal (twelfth cranial) nerve supplies the ipsilateral muscles of the tongue. The nucleus of the nerve or its fibers of exit may be involved by intramedullary lesions such as tumor, poliomyelitis, or most often motor neuron disease. Lesions of the basal meninges and the occipital bones (platybasia, invagination of occipital condyles, Paget's disease) may compress the nerve in its extramedullary course or in the hypoglossal canal. Isolated lesions of unknown cause can occur. Atrophy and fasciculation of the tongue develop weeks to months after interruption of the nerve.

Several cranial nerves may be affected by the same disease process. In this situation, the main clinical problem is to determine whether the lesion lies within the brainstem or outside it. Lesions that lie on the surface of the brainstem are characterized by involvement of adjacent cranial nerves (often occurring in succession) and late and rather slight involvement of the long sensory and motor pathways and segmental structures lying within the brainstem. The opposite is true of primary lesions within the brainstem. The extramedullary lesion is more likely to cause bone erosion or enlargement of the foramina of exit of cranial nerves. The intramedullary lesion involving cranial nerves often produces a crossed sensory or motor paralysis (cranial nerve signs on one side of the body and tract signs on the opposite side).

Involvement of multiple cranial nerves outside the brainstem is frequently the result of trauma, localized infections including varicella-zoster virus, infectious and noninfectious (especially carcinomatous) causes of meningitis (Chaps. 164 and 165), granulomatous diseases such as Wegener's granulomatosis, Behçet's disease, vascular disorders including those associated with diabetes, enlarging aneurysms, or locally infiltrating tumors. Among the tumors, nasopharyngeal cancers, lymphomas, neurofibromas, meningiomas, chordomas, cholesteatomas, carcinomas, and sarcomas have all been observed to involve a succession of lower cranial nerves. Owing to their anatomic relationships, the multiple cranial nerve palsies form a number of distinctive syndromes, listed in Table 455-2. Sarcoidosis is the cause of some cases of multiple cranial neuropathy; tuberculosis, the Chiari malformation, platybasia, and basilar invagination of the skull are additional causes. A purely motor disorder without atrophy always raises the question of myasthenia gravis (Chap. 461). As noted above, Guillain-Barré syndrome commonly affects the facial nerves bilaterally. In the Fisher variant of the Guillain-Barré syndrome, oculomotor paresis occurs with ataxia and areflexia in the limbs (Chap. 460). Wernicke's encephalopathy can cause a severe ophthalmoplegia combined with other brainstem signs (Chap. 330).

The cavernous sinus syndrome (Fig. 455-4) is a distinctive and frequently life-threatening disorder. It often presents as orbital or facial pain; orbital swelling and chemosis due to occlusion of the ophthalmic veins; fever; oculomotor neuropathy affecting the third, fourth, and sixth cranial nerves; and trigeminal neuropathy affecting the ophthalmic (V1) and occasionally the maxillary (V2) divisions of the trigeminal nerve. Cavernous sinus thrombosis, often secondary to infection from orbital cellulitis (frequently *Staphylococcus aureus*), a cutaneous source on the face, or sinusitis (especially with mucormycosis in diabetic patients), is the most frequent cause; other etiologies include

aneurysm of the carotid artery, a carotid-cavernous fistula (orbital bruit may be present), meningioma, nasopharyngeal carcinoma, other tumors, or an idiopathic granulomatous disorder (Tolosa-Hunt syndrome). The two cavernous sinuses directly communicate via intercavernous channels; thus, involvement on one side may extend to become bilateral. Early diagnosis is essential, especially when due to infection, and treatment depends on the underlying etiology.

In infectious cases, prompt administration of broad-spectrum antibiotics, drainage of any abscess cavities, and identification of the offending organism are essential. Anticoagulant therapy may benefit cases of primary thrombosis. Repair or occlusion of the carotid artery may be required for treatment of fistulas or aneurysms. The Tolosa-Hunt syndrome generally responds to glucocorticoids. A dramatic improvement in pain is usually evident within a few days; oral prednisone (60 mg daily) is usually continued for 2 weeks and then gradually tapered over a month, or longer if pain recurs. Occasionally an immunosuppressive medication, such as azathioprine or methotrexate, needs to be added to maintain an initial response to glucocorticoids.

An idiopathic form of multiple cranial nerve involvement on one or both sides of the face is occasionally seen. The syndrome consists of a subacute onset of boring facial pain, followed by paralysis of motor cranial nerves. The clinical features overlap those of the Tolosa-Hunt syndrome and appear to be due to idiopathic inflammation of the dura mater, which may be visualized by MRI. The syndrome is usually responsive to glucocorticoids.

456 Diseases of the Spinal Cord

Stephen L. Hauser, Allan H. Ropper

Diseases of the spinal cord are frequently devastating. They produce quadriplegia, paraplegia, and sensory deficits far beyond the damage they would inflict elsewhere in the nervous system because the spinal cord contains, in a small cross-sectional area, almost the entire motor output and sensory input of the trunk and limbs. Many spinal cord diseases are reversible if recognized and treated at an early stage (Table 456-1); thus, they are among the most critical of neurologic emergencies. The efficient use of diagnostic procedures, guided by knowledge of the anatomy and the clinical features of spinal cord diseases, is required to maximize the likelihood of a successful outcome.

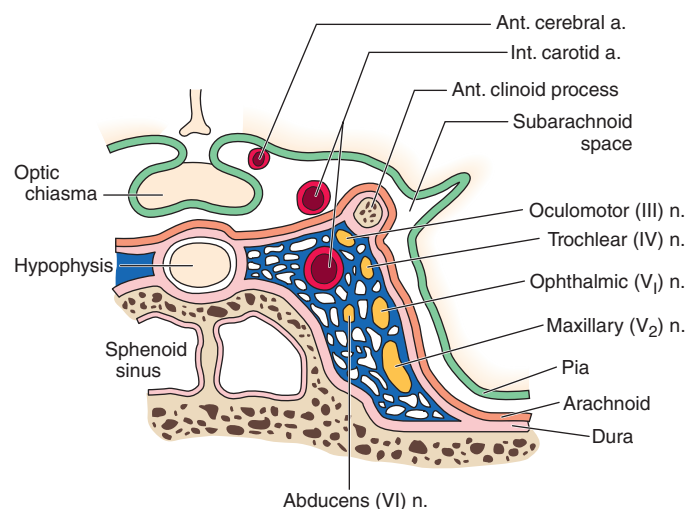


FIGURE 455-4 Anatomy of the cavernous sinus in coronal section, illustrating the location of the cranial nerves in relation to the vascular sinus, internal carotid artery (which loops anteriorly to the section), and surrounding structures.

APPROACH TO THE PATIENT: Spinal Cord Disease

SPINAL CORD ANATOMY RELEVANT TO CLINICAL SIGNS

The spinal cord is a thin, tubular extension of the central nervous system contained within the bony spinal canal. It originates at the medulla and continues caudally to the conus medullaris at the lumbar level; its fibrous extension, the filum terminale, terminates at the coccyx. The adult spinal cord is ~46 cm (18 in.) long, oval in shape, and enlarged in the cervical and lumbar regions, where neurons that innervate the upper and lower extremities, respectively, are located. The white matter tracts containing ascending sensory and descending motor pathways are located peripherally, whereas nerve cell bodies are clustered in an inner region of gray matter shaped like a four-leaf clover that surrounds the central canal (anatomically an extension of the fourth ventricle). The membranes that cover the spinal cord—the pia, arachnoid, and dura—are continuous with those of the brain, and the cerebrospinal fluid is contained within the subarachnoid space between the pia and arachnoid.

The spinal cord has 31 segments, each defined by an exiting ventral motor root and entering dorsal sensory root. During

TABLE 456-1 TREATABLE SPINAL CORD DISORDERS

Compressive	
Epidural, intradural, or intramedullary neoplasm	
Epidural abscess	
Epidural hemorrhage	
Cervical spondylosis	
Herniated disk	
Posttraumatic compression by fractured or displaced vertebra or hemorrhage	
Vascular	
Arteriovenous malformation and dural fistula	
Antiphospholipid syndrome and other hypercoagulable states	
Inflammatory	
Multiple sclerosis	
Neuromyelitis optica	
Transverse myelitis	
Sarcoidosis	
Sjögren-related myelopathy	
Systemic lupus erythematosus-related myelopathy	
Vasculitis	
Infectious	
Viral: VZV, HSV-1 and -2, CMV, HIV, HTLV-1, others	
Bacterial and mycobacterial: <i>Borrelia</i> , <i>Listeria</i> , syphilis, others	
<i>Mycoplasma pneumoniae</i>	
Parasitic: schistosomiasis, toxoplasmosis	
Developmental	
Syringomyelia	
Meningocele	
Tethered cord syndrome	
Metabolic	
Vitamin B ₁₂ deficiency (subacute combined degeneration)	
Copper deficiency	

Abbreviations: CMV, cytomegalovirus; HSV, herpes simplex virus; HTLV, human T cell lymphotropic virus; VZV, varicella-zoster virus.

embryologic development, growth of the cord lags behind that of the vertebral column, and the mature spinal cord ends at approximately the first lumbar vertebral body. The lower spinal nerves take an increasingly downward course to exit via intervertebral foramina. The first seven pairs of cervical spinal nerves exit above the same-numbered vertebral bodies, whereas all the subsequent nerves exit below the same-numbered vertebral bodies because of the presence of eight cervical spinal cord segments but only seven cervical vertebrae. The relationship between spinal cord segments and the corresponding vertebral bodies is shown in [Table 456-2](#). These relationships assume particular importance for localization of lesions that cause spinal cord compression. Sensory loss below the circumferential level of the umbilicus, for example, corresponds to the T10 cord segment but indicates involvement of the cord adjacent to the seventh or eighth thoracic vertebral body (see [Figs. 31-2 and 31-3](#)). In addition, at every level, the main ascending and descending tracts are somatotopically organized with a laminated distribution that reflects the origin or destination of nerve fibers.

TABLE 456-2 SPINAL CORD LEVELS RELATIVE TO THE VERTEBRAL BODIES

Spinal Cord Level	Corresponding Vertebral Body
Upper cervical	Same as cord level
Lower cervical	1 level higher
Upper thoracic	2 levels higher
Lower thoracic	2 to 3 levels higher
Lumbar	T10-T12
Sacral	T12-L1

Determining the Level of the Lesion The presence of a horizontally defined level below which sensory, motor, and autonomic function is impaired is a hallmark of a lesion of the spinal cord. This *sensory level* is sought by asking the patient to identify a pinprick or cold stimulus applied to the proximal legs and lower trunk and successively moved up toward the neck on each side. Sensory loss below this level is the result of damage to the spinothalamic tract on the opposite side, one to two segments higher in the case of a unilateral spinal cord lesion, and at the level of a bilateral lesion. The discrepancy in the level of a unilateral lesion is the result of the course of the second-order sensory fibers, which originate in the dorsal horn, and ascend for one or two levels as they cross anterior to the central canal to join the opposite spinothalamic tract. Lesions that transect the descending corticospinal and other motor tracts cause paraplegia or quadriplegia with heightened deep tendon reflexes, Babinski signs, and eventual spasticity (the upper motor neuron syndrome). Transverse damage to the cord also produces autonomic disturbances consisting of absent sweating below the implicated cord level and bladder, bowel, and sexual dysfunction.

The uppermost level of a spinal cord lesion can also be localized by attention to the *segmental signs* corresponding to disturbed motor or sensory innervation by an individual cord segment. A band of altered sensation (hyperalgesia or hyperpathia) at the upper end of the sensory disturbance, fasciculations or atrophy in muscles innervated by one or several segments, or a muted or absent deep tendon reflex may be noted at this level. These signs also can occur with focal root or peripheral nerve disorders; thus, they are most useful when they occur together with signs of long tract damage. With severe and acute transverse lesions, the limbs initially may be flaccid rather than spastic. This state of “spinal shock” lasts for several days, rarely for weeks, and may be mistaken for extensive damage to the anterior horn cells over many segments of the cord or for an acute polyneuropathy.

The main features of transverse damage at each level of the spinal cord are summarized below.

CERVICAL CORD Upper cervical cord lesions produce quadriplegia and weakness of the diaphragm. The uppermost level of weakness and reflex loss with lesions at C5-C6 is in the biceps; at C7, in finger and wrist extensors and triceps; and at C8, finger and wrist flexion. Horner’s syndrome (miosis, ptosis, and facial hypohidrosis) may accompany a cervical cord lesion at any level.

THORACIC CORD Lesions here are localized by the sensory level on the trunk and, if present, by the site of midline back pain. Useful markers of the sensory level on the trunk are the nipples (T4) and umbilicus (T10). Leg weakness and disturbances of bladder and bowel function accompany the paralysis. Lesions at T9-T10 paralyze the lower—but not the upper—abdominal muscles, resulting in upward movement of the umbilicus when the abdominal wall contracts (*Beevor’s sign*).

LUMBAR CORD Lesions at the L2-L4 spinal cord levels paralyze flexion and adduction of the thigh, weaken leg extension at the knee, and abolish the patellar reflex. Lesions at L5-S1 paralyze only movements of the foot and ankle, flexion at the knee, and extension of the thigh, and abolish the ankle jerks (S1).

SACRAL CORD/CONUS MEDULLARIS The conus medullaris is the tapered caudal termination of the spinal cord, comprising the sacral and single coccygeal segments. The distinctive conus syndrome consists of bilateral saddle anesthesia (S3-S5), prominent bladder and bowel dysfunction (urinary retention and incontinence with lax anal tone), and impotence. The bulbocavernosus (S2-S4) and anal (S4-S5) reflexes are absent ([Chap. 437](#)). Muscle strength is largely preserved. By contrast, lesions of the cauda equina, the nerve roots derived from the lower cord, are characterized by low back and radicular pain, asymmetric leg weakness and sensory loss, variable areflexia in the lower extremities, and relative sparing of bowel and bladder function. Mass lesions in the lower spinal

canal often produce a mixed clinical picture with elements of both cauda equina and conus medullaris syndromes. **Cauda equina syndromes are also discussed in Chap. 22.**

Special Patterns of Spinal Cord Disease The location of the major ascending and descending pathways of the spinal cord are shown in Fig. 456-1. Most fiber tracts—including the posterior columns and the spinocerebellar and pyramidal tracts—are situated on the side of the body they innervate. However, afferent fibers mediating pain and temperature sensation ascend in the spinothalamic tract contralateral to the side they supply. The anatomic configurations of these tracts produce characteristic syndromes that provide clues to the underlying disease process.

BROWN-SEQUARD HEMICORD SYNDROME This consists of ipsilateral weakness (corticospinal tract) and loss of joint position and vibratory sense (posterior column), with contralateral loss of pain and temperature sense (spinothalamic tract) one or two levels below the lesion. Segmental signs, such as radicular pain, muscle atrophy, or loss of a deep tendon reflex, are unilateral. Partial forms are more common than the fully developed syndrome.

CENTRAL CORD SYNDROME This syndrome results from selective damage to the gray matter nerve cells and crossing spinothalamic tracts surrounding the central canal. In the cervical cord, the central cord syndrome produces arm weakness out of proportion to leg weakness and a “dissociated” sensory loss, meaning loss of pain and temperature sensations over the shoulders, lower neck, and upper trunk (cape distribution), in contrast to preservation of light touch, joint position, and vibration sense in these regions. Spinal trauma, syringomyelia, and intrinsic cord tumors are the main causes.

ANTERIOR SPINAL ARTERY SYNDROME Infarction of the cord is generally the result of occlusion or diminished flow in this artery. The result

is bilateral tissue destruction at several contiguous levels that spares the posterior columns. All spinal cord functions—motor, sensory, and autonomic—are lost below the level of the lesion, with the striking exception of retained vibration and position sensation.

FORAMEN MAGNUM SYNDROME Lesions in this area interrupt decussating pyramidal tract fibers destined for the legs, which cross caudal to those of the arms, resulting in weakness of the legs (*crural paresis*). Compressive lesions near the foramen magnum may produce weakness of the ipsilateral shoulder and arm followed by weakness of the ipsilateral leg, then the contralateral leg, and finally the contralateral arm, an “around the clock” pattern that may begin in any of the four limbs. There is typically suboccipital pain spreading to the neck and shoulders.

INTRAMEDULLARY AND EXTRAMEDULLARY SYNDROMES It is useful to differentiate *intramedullary* processes, arising within the substance of the cord, from *extramedullary* ones that lie outside the cord and compress the spinal cord or its vascular supply. The differentiating features are only relative and serve as clinical guides. With extramedullary lesions, radicular pain is often prominent, and there is early sacral sensory loss and spastic weakness in the legs with incontinence due to the superficial location of the corresponding sensory and motor fibers in the spinothalamic and corticospinal tracts (Fig. 456-1). Intramedullary lesions tend to produce poorly localized burning pain rather than radicular pain and to spare sensation in the perineal and sacral areas (“sacral sparing”), reflecting the laminated configuration of the spinothalamic tract with sacral fibers outermost; corticospinal tract signs appear later. Regarding extramedullary lesions, a further distinction is made between extradural and intradural masses, as the former are generally malignant and the latter benign (neurofibroma being a common cause). Consequently, a long duration of symptoms favors an intradural origin.

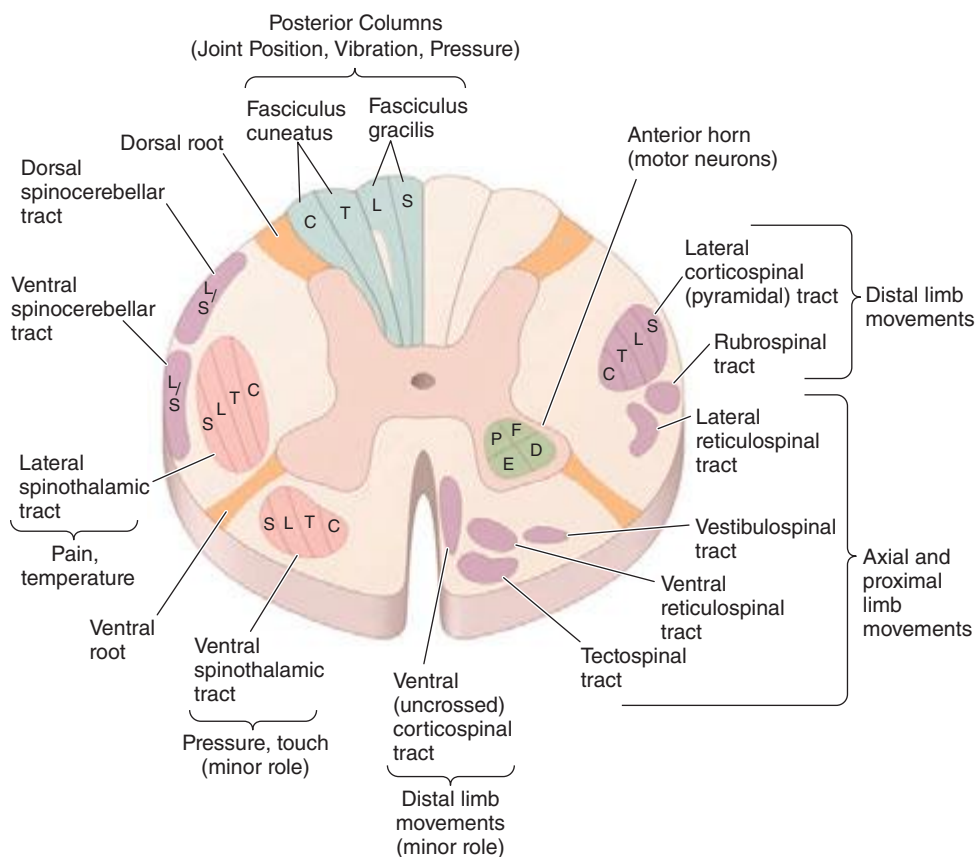


FIGURE 456-1 Transverse section through the spinal cord, composite representation, illustrating the principal ascending (*left*) and descending (*right*) pathways. The lateral and ventral spinothalamic tracts ascend contralateral to the side of the body that is innervated. C, cervical; D, distal; E, extensors; F, flexors; L, lumbar; P, proximal; S, sacral; T, thoracic.

ACUTE AND SUBACUTE SPINAL CORD DISEASES

The initial symptoms of structural diseases of the cord that evolve over days or weeks are focal neck or back pain, followed by various combinations of paresthesias, sensory loss, motor weakness, and sphincter disturbance. There may be only mild sensory symptoms or a devastating functional transection of the cord. Partial lesions selectively involve the posterior columns or anterior spinothalamic tracts or are limited to one side of the cord. Paresthesias or numbness typically begins in the feet and ascend symmetrically or asymmetrically. These symptoms simulate a polyneuropathy, but a sharply demarcated spinal cord level indicates the myelopathic nature of the process.

In severe and abrupt cases, areflexia reflecting spinal shock may be present, but hyperreflexia supervenes over days or weeks; persistent areflexic paralysis with a sensory level usually indicates necrosis over multiple segments of the spinal cord.

APPROACH TO THE PATIENT: Compressive and Noncompressive Myelopathy

DISTINGUISHING COMPRESSIVE FROM NONCOMPRESSIVE MYELOPATHY

The first priority is to exclude a treatable compression of the cord by a mass that may be amenable to treatment. The common causes are tumor, epidural abscess or hematoma, herniated disk, and vertebral pathology. Epidural compression due to malignancy or abscess often causes warning signs of neck or back pain, bladder disturbances, and sensory symptoms that precede the development of paralysis. Spinal subluxation, hemorrhage, and noncompressive etiologies such as infarction are more likely to produce myelopathy without antecedent symptoms. Magnetic resonance imaging (MRI) with gadolinium, centered on the clinically suspected level, is the initial diagnostic procedure if it is available; in some cases, it is appropriate to image the entire spine (cervical through sacral regions) to search for additional clinically inapparent lesions. Once compressive lesions have been excluded, noncompressive causes of acute myelopathy that are intrinsic to the cord are considered, primarily vascular, inflammatory, and infectious etiologies.

COMPRESSIVE MYELOPATHIES

Neoplastic Spinal Cord Compression In adults, most neoplasms are epidural in origin, resulting from metastases to the adjacent vertebral column. The propensity of solid tumors to metastasize to the vertebral column probably reflects the high proportion of bone marrow located in the axial skeleton. Almost any malignant tumor can metastasize to the spinal column, with breast, lung, prostate, kidney, lymphoma, and myeloma being particularly frequent. The thoracic spinal column is most commonly involved; exceptions are metastases from prostate and ovarian cancer, which occur disproportionately in the sacral and lumbar vertebrae, probably from spread through Batson's plexus, a network of veins along the anterior epidural space. Retroperitoneal neoplasms (especially lymphomas or sarcomas) enter the spinal canal laterally through the intervertebral foramina and produce radicular pain with signs of weakness that corresponds to the level of involved nerve roots.

Pain is usually the initial symptom of spinal metastasis; it may be aching and localized or sharp and radiating in quality and typically worsens with movement, coughing, or sneezing and characteristically awakens patients at night. A recent onset of persistent back pain, particularly if in the thoracic spine (which is uncommonly involved by spondylosis), should prompt consideration of vertebral metastasis. Rarely, pain is mild or absent. Plain radiographs of the spine and radionuclide bone scans have a limited role in diagnosis because they do not identify 15–20% of metastatic vertebral lesions and fail to detect paravertebral masses that reach the epidural space through the intervertebral foramina. MRI provides excellent anatomic resolution of the extent of spinal tumors (Fig. 456-2) and is able to distinguish between malignant lesions and other masses—epidural abscess, tuberculoma,

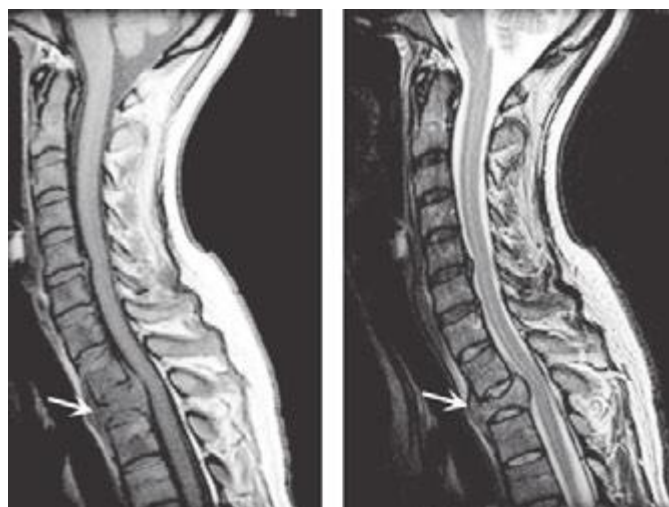


FIGURE 456-2 Epidural spinal cord compression due to breast carcinoma. Sagittal T1-weighted (A) and T2-weighted (B) magnetic resonance imaging scans through the cervicothoracic junction reveal an infiltrated and collapsed second thoracic vertebral body with posterior displacement and compression of the upper thoracic spinal cord. The low-intensity bone marrow signal in A signifies replacement by tumor.

lipoma, or epidural hemorrhage, among others—that present in a similar fashion. Vertebral metastases are usually hypointense relative to a normal bone marrow signal on T1-weighted MRI; after the administration of gadolinium, contrast enhancement may deceptively “normalize” the appearance of the tumor by increasing its intensity to that of normal bone marrow. Infections of the spinal column (osteomyelitis and related disorders) are distinctive in that, unlike tumor, they often cross the disk space to involve the adjacent vertebral body.

If spinal cord compression is suspected, imaging should be obtained promptly. If there are radicular symptoms but no evidence of myelopathy, it may be safe to defer imaging for 24–48 h. Up to 40% of patients who present with cord compression at one level are found to have asymptomatic epidural metastases elsewhere; thus, the length of the spine is often imaged when epidural malignancy is in question.

TREATMENT NEOPLASTIC SPINAL CORD COMPRESSION

Management of cord compression includes glucocorticoids to reduce cord edema, local radiotherapy (initiated as early as possible) to the symptomatic lesion, and specific therapy for the underlying tumor type. Glucocorticoids (dexamethasone, up to 40 mg daily) can be administered before an imaging study if there is clinical suspicion of cord compression and the medication is continued at a lower dose until definitive treatment with radiotherapy (generally 3000 cGy administered in 15 daily fractions) and/or surgical decompression is completed. In one randomized controlled trial, initial management with surgery followed by radiotherapy was more effective than radiotherapy alone for patients with a single area of spinal cord compression by extradural tumor; however, patients with recurrent cord compression, brain metastases, radiosensitive tumors, or severe motor symptoms of >48 h in duration were excluded from this study. Radiotherapy alone may be effective even for some typically radioresistant metastases. A good response to therapy can be expected in individuals who are ambulatory at presentation. Treatment usually prevents new weakness, and some recovery of motor function occurs in up to one-third of patients. Motor deficits (paraplegia or quadriplegia), once established for >12 h, do not usually improve, and beyond 48 h the prognosis for substantial motor recovery is poor. Although most patients do not experience recurrences in the months following radiotherapy, with survival beyond 2 years, recurrence becomes

2654 increasingly likely and can be managed with additional radiotherapy. Newer techniques such as stereotactic radiosurgery can deliver high doses of focused radiation and with similar rates of response compared to traditional radiotherapy. Biopsy of the epidural mass is unnecessary in patients with known primary cancer, but it is indicated if a history of underlying cancer is lacking. Surgery, either decompression by laminectomy or vertebral body resection, is also indicated when signs of cord compression worsen despite radiotherapy, when the maximum-tolerated dose of radiotherapy has been delivered previously to the site, or when a vertebral compression fracture or spinal instability contributes to cord compression.

In contrast to tumors of the epidural space, most intradural mass lesions are slow-growing and benign. Meningiomas and neurofibromas account for most of these, with occasional cases caused by chordoma, lipoma, dermoid, or sarcoma. Meningiomas (Fig. 456-3) are often located posterior to the thoracic cord or near the foramen magnum, although they can arise from the meninges anywhere along the spinal canal. Neurofibromas are benign tumors of the nerve sheath that typically arise from the posterior root; when multiple, neurofibromatosis is the likely etiology. Symptoms usually begin with radicular sensory symptoms followed by an asymmetric, progressive spinal cord syndrome. Therapy is surgical resection.

Primary intramedullary tumors of the spinal cord are uncommon. They present as central cord or hemicord syndromes, often in the cervical region. There may be poorly localized burning pain in the extremities and sparing of sacral sensation. In adults, these lesions are ependymomas, hemangioblastomas, or low-grade astrocytomas (Fig. 456-4). Complete resection of an intramedullary ependymoma is often possible with microsurgical techniques. Debulking of an intramedullary astrocytoma can also be helpful, as these are often slowly growing lesions; the value of adjunctive radiotherapy and chemotherapy is uncertain. Secondary (metastatic) intramedullary tumors also occur, especially in patients with advanced metastatic disease (Chap. 118), although these are not nearly as frequent as brain metastases.



FIGURE 456-3 Magnetic resonance imaging of a thoracic meningioma. Coronal T1-weighted postcontrast image through the thoracic spinal cord demonstrates intense and uniform enhancement of a well-circumscribed extramedullary mass (arrows) that displaces the spinal cord to the left.



FIGURE 456-4 Magnetic resonance imaging of an intramedullary astrocytoma. Sagittal T1-weighted postcontrast image through the cervical spine demonstrates expansion of the upper cervical spine by a mass lesion emanating from within the spinal cord at the cervicomedullary junction. Irregular peripheral enhancement occurs within the mass (arrows).

Spinal Epidural Abscess Spinal epidural abscess presents with midline back or neck pain, fever, and progressive limb weakness. Prompt recognition of this distinctive process may prevent permanent sequelae. Aching pain is almost always present, either over the spine or in a radicular pattern. The duration of pain prior to presentation is generally ≤ 2 weeks but may on occasion be several months or longer. Fever is typically but not invariably present, accompanied by elevated white blood cell count, sedimentation rate, and C-reactive protein. As the abscess expands, further spinal cord damage results from venous congestion and thrombosis. Once weakness and other signs of myelopathy appear, progression may be rapid and irreversible. A more chronic sterile granulomatous form of abscess is also known, usually after treatment of an acute epidural infection.

Risk factors include an impaired immune status (HIV, diabetes mellitus, renal failure, alcoholism, malignancy), intravenous drug abuse, and infections of the skin or other tissues. Two-thirds of epidural infections result from hematogenous spread of bacteria from the skin (furunculosis), soft tissue (pharyngeal or dental abscesses; sinusitis), or deep viscera (bacterial endocarditis). The remainder arises from direct extension of a local infection to the subdural space; examples of local predisposing conditions are vertebral osteomyelitis, decubitus ulcers, lumbar puncture, epidural anesthesia, or spinal surgery. Most cases are due to *Staphylococcus aureus*; gram-negative bacilli, *Streptococcus*, anaerobes, and fungi can also cause epidural abscesses. Tuberculosis from an adjacent vertebral source (Pott's disease) remains an important cause in the developing world.

MRI (Fig. 456-5) localizes the abscess and excludes other causes of myelopathy. Blood cultures are positive in more than half of cases, but direct aspiration of the abscess at surgery is often required for a microbiologic diagnosis. Lumbar puncture is only required if encephalopathy or other clinical signs raise the question of associated meningitis, a feature that is found in $<25\%$ of cases. The level of the puncture should be planned to minimize the risk of meningitis due to passage of the needle through infected tissue. A high cervical tap is sometimes the safest approach. Cerebrospinal fluid (CSF) abnormalities in epidural and subdural abscess consist of pleocytosis with a preponderance of polymorphonuclear cells, an elevated protein level, and a reduced glucose level, but the responsible organism is not cultured unless there is associated meningitis.

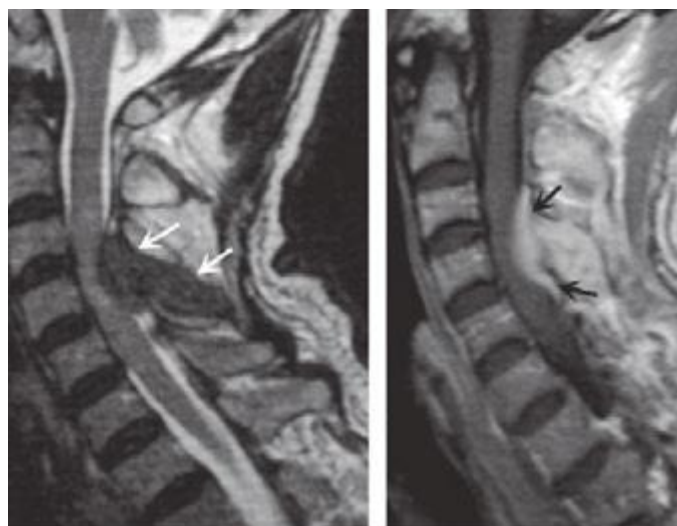


FIGURE 456-5 Magnetic resonance (MR) imaging of a spinal epidural abscess due to tuberculosis. **A.** Sagittal T2-weighted free spin-echo MR sequence. A hypointense mass replaces the posterior elements of C3 and extends epidurally to compress the spinal cord (arrows). **B.** Sagittal T1-weighted image after contrast administration reveals a diffuse enhancement of the epidural process (arrows) with extension into the epidural space.

TREATMENT SPINAL EPIDURAL ABSCESS

Treatment is by decompressive laminectomy with debridement combined with long-term antibiotic treatment. Surgical evacuation prevents development of paralysis and may improve or reverse paralysis in evolution, but it is unlikely to improve deficits of more than several days in duration. Broad-spectrum antibiotics should be started empirically before surgery and then modified on the basis of culture results; medication is generally continued for at least 6 weeks. If surgery is contraindicated or if there is a fixed paraplegia or quadriplegia that is unlikely to improve following surgery, long-term administration of systemic and oral antibiotics can be used; in such cases, the choice of antibiotics may be guided by results of blood cultures. Surgical management remains the treatment of choice unless the abscess is limited in size and causes few or no neurologic signs.

With prompt diagnosis and treatment of spinal epidural abscess, up to two-thirds of patients experience significant recovery.

Spinal Epidural Hematoma Hemorrhage into the epidural (or subdural) space causes acute focal or radicular pain followed by variable signs of a spinal cord or conus medullaris disorder. Therapeutic anticoagulation, trauma, tumor, or blood dyscrasias are predisposing conditions. Rare cases complicate lumbar puncture or epidural anesthesia. MRI and computed tomography (CT) confirm the clinical suspicion and can delineate the extent of the bleeding. Treatment consists of prompt reversal of any underlying clotting disorder and surgical decompression. Surgery may be followed by substantial recovery, especially in patients with some preservation of motor function preoperatively. Because of the risk of hemorrhage, lumbar puncture should be avoided whenever possible in patients with severe thrombocytopenia or other coagulopathies.

Hematomyelia Hemorrhage into the substance of the spinal cord is a rare result of trauma, intraparenchymal vascular malformation (see below), vasculitis due to polyarteritis nodosa or systemic lupus erythematosus (SLE), bleeding disorders, or a spinal cord neoplasm. Hematomyelia presents as an acute painful transverse myelopathy. With large lesions, extension into the subarachnoid space results in subarachnoid hemorrhage (Chap. 330). Diagnosis is by MRI or CT.

TABLE 456-3 EVALUATION OF ACUTE TRANSVERSE MYELOPATHY

1. MRI of spinal cord with and without contrast (exclude compressive causes).
2. CSF studies: Cell count, protein, glucose, IgG index/synthesis rate, oligoclonal bands, VDRL; Gram's stain, acid-fast bacilli, and India ink stains; PCR for VZV, HSV-2, HSV-1, EBV, CMV, HHV-6, enteroviruses, HIV; antibody for HTLV-1, *Borrelia burgdorferi*, *Mycoplasma pneumoniae*, and *Chlamydia pneumoniae*; viral, bacterial, mycobacterial, and fungal cultures.
3. Blood studies for infection: HIV; RPR; IgG and IgM enterovirus antibody; IgM mumps, measles, rubella, group B arbovirus, *Brucella melitensis*, *Chlamydia psittaci*, *Bartonella henselae*, schistosomal antibody; cultures for *B. melitensis*. Also consider nasal/pharyngeal/anal cultures for enteroviruses; stool O&P for *Schistosoma* ova.
4. Immune-mediated disorders: ESR; ANA; ENA; dsDNA; rheumatoid factor; anti-SSA; anti-SSB, complement levels; antiphospholipid and anticardiolipin antibodies; p-ANCA; antimicrobial and antithyroglobulin antibodies; if Sjögren's syndrome suspected, Schirmer test, salivary gland scintigraphy, and salivary/lacrimal gland biopsy.
5. Sarcoidosis: Serum angiotensin-converting enzyme; serum Ca; 24-h urine Ca; chest x-ray; chest CT; total-body gallium scan; lymph node biopsy.
6. Demyelinating disease: Brain MRI scan, evoked potentials, CSF oligoclonal bands, neuromyelitis optica antibody (anti-aquaporin-4 [NMO] antibody).
7. Vascular causes: MRI, CT myelogram; spinal angiogram.

Abbreviations: ANA, antinuclear antibodies; CMV, cytomegalovirus; CSF, cerebrospinal fluid; CT, computed tomography; EBV, Epstein-Barr virus; ENA, epithelial neutrophil-activating peptide; ESR, erythrocyte sedimentation rate; HHV, human herpes virus; HSV, herpes simplex virus; HTLV, human T cell leukemia/lymphoma virus; MRI, magnetic resonance imaging; O&P, ova and parasites; p-ANCA, perinuclear antineutrophilic cytoplasmic antibodies; PCR, polymerase chain reaction; RPR, rapid plasma reagin (test); VDRL, Venereal Disease Research Laboratory; VZV, varicella-zoster virus.

Therapy is supportive, and surgical intervention is generally not useful. An exception is hematomyelia due to an underlying vascular malformation, for which spinal angiography and endovascular occlusion may be indicated, or surgery to evacuate the clot and remove the underlying vascular lesion.

NONCOMPRESSIVE MYELOPATHIES

The most frequent causes of noncompressive acute transverse myelopathy are spinal cord infarction; systemic inflammatory disorders, including SLE and sarcoidosis; demyelinating diseases, including multiple sclerosis (MS); neuromyelitis optica (NMO); postinfectious or idiopathic transverse myelitis, which is presumed to be an immune condition related to acute disseminated encephalomyelitis (Chap. 458); and infectious (primarily viral) causes. After spinal cord compression is excluded, the evaluation generally requires a lumbar puncture and a search for underlying systemic disease (Table 456-3).

Spinal Cord Infarction The cord is supplied by three arteries that course vertically over its surface: a single anterior spinal artery and paired posterior spinal arteries. The anterior spinal artery originates in paired branches of the vertebral arteries at the craniocervical junction and is fed by additional radicular vessels that arise at C6, at an upper thoracic level, and, most consistently, at T11-L2 (artery of Adamkiewicz). At each spinal cord segment, paired penetrating vessels branch from the anterior spinal artery to supply the anterior two-thirds of the cord; the posterior spinal arteries, which often become less distinct below the midthoracic level, supply the posterior columns.

Spinal cord ischemia can occur at any level; however, the presence of the artery of Adamkiewicz below, and the anterior spinal artery circulation above, creates a region of marginal blood flow in the upper thoracic segments. With hypotension or cross-clamping of the aorta, cord infarction typically occurs at the level of T3-T4, and also at boundary zones between the anterior and posterior spinal artery territories. The latter may result in a rapidly progressive syndrome over hours of weakness and spasticity with little sensory change.

Acute infarction in the territory of the *anterior spinal artery* produces paraplegia or quadriplegia, dissociated sensory loss affecting pain and temperature sense but sparing vibration and position sense, and loss of sphincter control (“anterior cord syndrome”). Onset may be sudden but more typically is progressive over minutes or a few hours, quite unlike stroke in the cerebral hemispheres. Sharp midline

or radiating back pain localized to the area of ischemia is frequent. Areflexia due to spinal shock is often present initially; with time, hyperreflexia and spasticity appear. Less common is infarction in the territory of the *posterior spinal arteries*, resulting in loss of posterior column function either on one side or bilaterally.

Causes of spinal cord infarction include aortic atherosclerosis, dissecting aortic aneurysm, vertebral artery occlusion or dissection in the neck, aortic surgery, or profound hypotension from any cause. A “surfer’s myelopathy” in the cervical region is probably vascular in origin. Cardiogenic emboli, vasculitis (**Chap. 385**), and collagen vascular disease (particularly SLE [**Chap. 378**], Sjögren’s syndrome [**Chap. 383**], and the antiphospholipid antibody syndrome [**Chap. 379**]) are other etiologies. Occasional cases develop from *embolism of nucleus pulposus* material into spinal vessels, usually from local spine trauma. In a substantial number of cases, no cause can be found, and thromboembolism in arterial feeders is suspected. MRI may fail to demonstrate infarctions of the cord, especially in the first day, but often the imaging becomes abnormal at the affected level.

In cord infarction due to presumed thromboembolism, acute anticoagulation is not indicated, with the possible exception of the unusual transient ischemic attack or incomplete infarction with a stuttering or progressive course. The antiphospholipid antibody syndrome is treated with anticoagulation (**Chap. 379**). Lumbar drainage of spinal fluid has reportedly been successful in some cases of cord infarction and has been used prophylactically during aortic surgery, but it has not been studied systematically.

Inflammatory and Immune Myelopathies (Myelitis) This broad category includes the demyelinating conditions MS, NMO, and postinfectious myelitis, as well as sarcoidosis and systemic autoimmune disease. In approximately one-quarter of cases of myelitis, no underlying cause can be identified. Some will later manifest additional symptoms of an immune-mediated disease. *Recurrent episodes of myelitis* are usually due to one of the immune-mediated diseases or to infection with herpes simplex virus (HSV) type 2 (below).

MULTIPLE SCLEROSIS MS may present with acute myelitis, particularly in individuals of Asian or African ancestry. In Caucasians, MS attacks rarely cause a transverse myelopathy (i.e., attacks of bilateral sensory disturbances, unilateral or bilateral weakness, and bladder or bowel symptoms), but it is among the most common causes of a partial cord syndrome. MRI findings in MS-associated myelitis typically consist of mild swelling of the cord and diffuse or multifocal “shoddy” areas of abnormal signal on T2-weighted sequences. Contrast enhancement, indicating disruption in the blood-brain barrier associated with inflammation, is present in many acute cases. A brain MRI is most helpful in gauging the likelihood that a case of myelitis represents an initial attack of MS. A normal scan indicates that the risk of evolution to MS is low, ~10–15% over 5 years; in contrast, the finding of multiple periventricular T2-bright lesions indicates a much higher risk, >50% over 5 years and >90% by 14 years. The CSF may be normal, but more often there is a mild mononuclear cell pleocytosis, with normal or mildly elevated CSF protein levels; the presence of oligoclonal bands is variable, but when they are found, a diagnosis of MS is more likely.

There are no adequate trials of therapy for MS-associated transverse myelitis. Intravenous methylprednisolone (500 mg qd for 3 days) followed by oral prednisone (1 mg/kg per day for several weeks, then gradual taper) has been used as initial treatment. A course of plasma exchange may be indicated for severe cases if glucocorticoids are ineffective. **MS is discussed in Chap. 458.**

NEUROMYELITIS OPTICA NMO is an immune-mediated demyelinating disorder consisting of a severe myelopathy that is typically longitudinally extensive, meaning that the lesion spans three or more vertebral segments. NMO is associated with optic neuritis that is often bilateral and may precede or follow myelitis by weeks or months, and also by brainstem and, in some cases, hypothalamic involvement. Recurrent myelitis without optic nerve involvement can also occur in NMO; affected individuals are usually female and often of Asian ancestry. CSF studies reveal a variable mononuclear pleocytosis of up to several hundred cells per microliter; unlike MS, oligoclonal bands are

generally absent. Diagnostic serum autoantibodies against the water channel protein aquaporin-4 are present in 60–70% of patients with NMO. NMO has also been associated with SLE and antiphospholipid antibodies (see below) as well as with other systemic autoimmune diseases; rare cases are paraneoplastic in origin. Treatment is with glucocorticoids and, for refractory cases, plasma exchange (as for MS, above). Preliminary studies suggest that treatment with azathioprine, mycophenolate, or anti-CD20 (anti-B cell) monoclonal antibody may protect against subsequent relapses; treatment for 5 years or longer is generally recommended. **NMO is discussed in Chap. 458.**

SYSTEMIC IMMUNE-MEDIATED DISORDERS Myelitis occurs in a small number of patients with SLE, many cases of which are associated with antibodies to antiphospholipids and/or to aquaporin-4. Patients with aquaporin-4 antibodies are likely to have longitudinally extensive myelitis by MRI, are considered to have an NMO-spectrum disorder, and are at high risk of developing future episodes of myelitis and/or optic neuritis. The CSF in SLE myelitis is usually normal or shows a mild lymphocytic pleocytosis; oligoclonal bands are a variable finding. Although there are no systematic trials of therapy for SLE myelitis, based on limited data, high-dose glucocorticoids followed by cyclophosphamide have been recommended. Acute severe episodes of transverse myelitis that do not initially respond to glucocorticoids are often treated with a course of plasma exchange. Sjögren’s syndrome (**Chap. 383**) can also be associated with NMO spectrum disorder and also with cases of acute transverse or chronic progressive myelopathy. Other immune-mediated myelitides include antiphospholipid antibody syndrome (**Chap. 379**), mixed connective tissue disease (**Chap. 382**), Behçet’s syndrome (**Chap. 387**), and vasculitis related to polyarteritis nodosa, perinuclear antineutrophilic cytoplasmic (p-ANCA) antibodies, or primary central nervous system vasculitis (**Chap. 385**).

Another important consideration in this group is sarcoid myelopathy that may present as a slowly progressive or relapsing disorder. MRI reveals an edematous swelling of the spinal cord that may mimic tumor; there is almost always gadolinium enhancement of active lesions and in some cases nodular enhancement of the adjacent surface of the cord; lesions may be single or multiple, and on axial images, enhancement of the central cord is usually present. The typical CSF profile consists of a mild lymphocytic pleocytosis and mildly elevated protein level; in a minority of cases, reduced glucose and oligoclonal bands are found. The diagnosis is particularly difficult when systemic manifestations of sarcoid are minor or absent (nearly 50% of cases) or when other typical neurologic manifestations of the disease—such as cranial neuropathy, hypothalamic involvement, or meningeal enhancement visualized by MRI—are lacking. A slit-lamp examination of the eye to search for uveitis, chest x-ray and CT to assess pulmonary involvement and mediastinal lymphadenopathy, serum or CSF angiotensin-converting enzyme (ACE; CSF values elevated in only a minority of cases), serum calcium, and a gallium scan may assist in the diagnosis. The usefulness of spinal fluid ACE is uncertain. Initial treatment is with oral glucocorticoids; immunosuppressant drugs, including the tumor necrosis factor α inhibitor infliximab, have been used for resistant cases. **Sarcoidosis is discussed in Chap. 390.**

POSTINFECTIOUS MYELITIS Many cases of myelitis, termed *postinfectious* or *postvaccinal*, follow an infection or vaccination. Numerous organisms have been implicated, including Epstein-Barr virus (EBV), cytomegalovirus (CMV), mycoplasma, influenza, measles, varicella, rubeola, and mumps. As in the related disorder acute disseminated encephalomyelitis (**Chap. 458**), postinfectious myelitis often begins as the patient appears to be recovering from an acute febrile infection, or in the subsequent days or weeks, but an infectious agent cannot be isolated from the nervous system or CSF. The presumption is that the myelitis represents an autoimmune disorder triggered by infection and is not due to direct infection of the spinal cord. No randomized controlled trials of therapy exist; treatment is usually with glucocorticoids or, in fulminant cases, plasma exchange.

ACUTE INFECTIOUS MYELITIS Many viruses have been associated with an acute myelitis that is infectious in nature rather than postinfectious. Nonetheless, the two processes are often difficult to distinguish. Herpes

zoster is the best characterized viral myelitis, but HSV types 1 and 2, EBV, CMV, and rabies virus are other well-described causes. HSV-2 (and less commonly HSV-1) produces a distinctive syndrome of recurrent sacral cauda equina neuritis in association with outbreaks of genital herpes (Elsberg's syndrome). Poliomyelitis is the prototypic viral myelitis, but it is more or less restricted to the anterior gray matter of the cord containing the spinal motoneurons. A polio-like syndrome can also be caused by a large number of enteroviruses (including enterovirus 71 and coxsackie), and with West Nile virus and other flaviviruses. Recently, cases of paralysis in children and adolescents were associated with enterovirus D-68 infection but a causal role for this virus has not been established. Chronic viral myelitic infections, such as those due to HIV or human T cell lymphotropic virus type 1 (HTLV-1), are discussed below.

Bacterial and mycobacterial myelitis (most are essentially abscesses) are less common than viral causes and much less frequent than cerebral bacterial abscess. Almost any pathogenic species may be responsible, including *Borrelia burgdorferi* (Lyme disease), *Listeria monocytogenes*, *Mycobacterium tuberculosis*, and *Treponema pallidum* (syphilis). *Mycoplasma pneumoniae* may be a cause of myelitis, but its status is uncertain because many cases are more properly classified as postinfectious.

Schistosomiasis (Chap. 259) is an important cause of parasitic myelitis in endemic areas. The process is intensely inflammatory and granulomatous, caused by a local response to tissue-digesting enzymes from the ova of the parasite, typically *Schistosoma mansoni*. Toxoplasmosis (Chap. 253) can occasionally cause a focal myelopathy, and this diagnosis should especially be considered in patients with AIDS (Chap. 226).

In cases of suspected viral myelitis, it may be appropriate to begin specific therapy pending laboratory confirmation. Herpes zoster, HSV, and EBV myelitis are treated with intravenous acyclovir (10 mg/kg q8h) or oral valacyclovir (2 g tid) for 10–14 days; CMV is treated with ganciclovir (5 mg/kg IV bid) plus foscarnet (60 mg/kg IV tid) or cidofovir (5 mg/kg per week for 2 weeks).

High-Voltage Electrical Injury Spinal cord injuries are prominent following electrocution from lightning strikes or other accidental electrical exposures. The syndrome consists of transient weakness acutely (often with an altered sensorium and focal cerebral disturbances), followed several days or even weeks later by a myelopathy that can be severe and permanent. This is a rare injury type, and limited data incriminate a vascular pathology involving the anterior spinal artery and its branches in some cases. Therapy is supportive.

CHRONIC MYELOPATHIES

SPONDYLOTIC MYELOPATHY

Spondylotic myelopathy is one of the most common causes of chronic cord compression and of gait difficulty in the elderly. Neck and shoulder pain with stiffness are early symptoms; impingement of bone and soft tissue overgrowth on nerve roots results in radicular arm pain, most often in a C5 or C6 distribution. Compression of the cervical cord, which occurs in fewer than one-third of cases, produces a slowly progressive spastic paraparesis, at times asymmetric and often accompanied by paresthesias in the feet and hands. Vibratory sense is diminished in the legs, there is a Romberg sign, and occasionally there is a sensory level for vibration or pinprick on the upper thorax. In some cases, coughing or straining produces leg weakness or radiating arm or shoulder pain. Dermatomal sensory loss in the arms, atrophy of intrinsic hand muscles, increased deep-tendon reflexes in the legs, and extensor plantar responses are common. Urinary urgency or incontinence occurs in advanced cases, but there are many alternative causes of these problems in older individuals. A tendon reflex in the arms is often diminished at some level; most often at the biceps (C5–C6). In individual cases, radicular, myelopathic, or combined signs may predominate. The diagnosis should be considered in appropriate cases of progressive cervical myelopathy, paresthesias of the feet and hands, or wasting of the hands.

Diagnosis is usually made by MRI and may be suspected from CT images; plain x-rays are less helpful. Extrinsic cord compression and

deformation are appreciated on axial MRI views, and T2-weighted sequences may reveal areas of high signal intensity within the cord adjacent to the site of compression. A cervical collar may be helpful in milder cases, but definitive therapy consists of surgical decompression. Posterior laminectomy or an anterior approach with resection of the protruded disk and bony material may be required. **Cervical spondylosis and related degenerative diseases of the spine are discussed in Chap. 22.**

VASCULAR MALFORMATIONS OF THE CORD AND DURA

Vascular malformations of the cord and overlying dura are treatable causes of progressive myelopathy. Most common are fistulas located within the dura or posteriorly along the surface of the cord. Most dural arteriovenous (AV) fistulas are located at or below the midthoracic level, usually consisting of a direct connection between a radicular feeding artery in the nerve root sleeve with dural veins. The typical presentation is a middle-aged man with a progressive myelopathy that worsens slowly or intermittently and may have periods of remission, sometimes mimicking MS. Acute deterioration due to hemorrhage into the spinal cord (hematomyelia) or subarachnoid space may also occur but is rare. A saltatory progression is most common and appears to be the result of local ischemia and edema from venous congestion. Most patients have incomplete sensory, motor, and bladder disturbances. The motor disorder may predominate and produce a mixture of upper and restricted lower motor neuron signs, simulating amyotrophic lateral sclerosis (ALS). Pain over the dorsal spine, dysesthesias, or radicular pain may be present. Other symptoms suggestive of AV malformation (AVM) or dural fistula include intermittent claudication; symptoms that change with posture, exertion, Valsalva maneuver, or menses; and fever.

Less commonly, AVM disorders are intramedullary rather than dural. One unusual disorder is a progressive thoracic myelopathy with paraparesis developing over weeks or months, characterized pathologically by abnormally thick, hyalinized vessels within the cord (subacute necrotic myelopathy, or Foix-Alajouanine syndrome).

Spinal bruits are infrequent but may be sought at rest and after exercise in suspected cases. A vascular nevus on the overlying skin may indicate an underlying vascular malformation as occurs with Klippel-Trenaunay-Weber syndrome. High-resolution MRI with contrast administration detects the draining vessels of many but not all AVMs (Fig. 456-6). An uncertain proportion may be visualized by CT myelography as enlarged vessels along the surface of the cord. Definitive diagnosis requires selective spinal angiography, which defines the feeding vessels and the extent of the malformation. Endovascular embolization of the major feeding vessels may stabilize a progressive neurologic deficit or allow for gradual recovery. Some lesions, especially small dural fistulas, can be resected surgically.

RETROVIRUS-ASSOCIATED MYELOPATHIES

The myelopathy associated with HTLV-1, formerly called tropical spastic paraparesis, is a slowly progressive spastic syndrome with variable sensory and bladder disturbance. Approximately half of patients have mild back or leg pain. The neurologic signs may be asymmetric, often lacking a well-defined sensory level; the only sign in the arms may be hyperreflexia after several years of illness. The onset is insidious, and the illness is slowly progressive at a variable rate; most patients are unable to walk within 10 years of onset. This presentation may resemble primary progressive MS or a thoracic AVM. Diagnosis is made by demonstration of HTLV-1-specific antibody in serum by enzyme-linked immunosorbent assay (ELISA), confirmed by radioimmunoprecipitation or Western blot analysis. Especially in endemic areas, a finding of HTLV-1 seropositivity in a patient with myelopathy does not necessarily prove that HTLV-1 is causative. The CSF/serum antibody index may provide support by establishing intrathecal synthesis of antibodies favoring HTLV-1 myelopathy over asymptomatic carriage. Measuring proviral DNA by polymerase chain reaction (PCR) in serum and CSF cells can be useful as an ancillary part of diagnosis, because proviral DNA levels may be higher in patients with myelopathy. The myelopathy appears to result from an

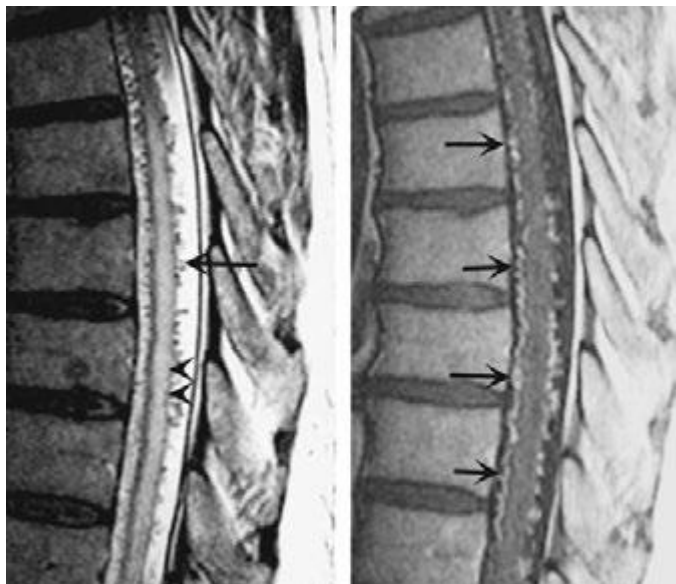


FIGURE 456-6 Arteriovenous malformation. Sagittal magnetic resonance scans of the thoracic spinal cord: T2 fast spin-echo technique (*left*) and T1 postcontrast image (*right*). On the T2-weighted image (*left*), abnormally high signal intensity is noted in the central aspect of the spinal cord (*arrowheads*). Numerous punctate flow voids indent the dorsal and ventral spinal cord (*arrow*). These represent the abnormally dilated venous plexus supplied by a dural arteriovenous fistula. After contrast administration (*right*), multiple, serpentine, enhancing veins (*arrows*) on the ventral and dorsal aspect of the thoracic spinal cord are visualized, diagnostic of arteriovenous malformation. This patient was a 54-year-old man with a 4-year history of progressive paraparesis.

immune-mediated attack on the spinal cord rather than the result of direct viral infection. There is no effective treatment, but symptomatic therapy for spasticity and bladder symptoms may be helpful.

A progressive myelopathy may also result from HIV infection (**Chap. 226**). It is characterized by vacuolar degeneration of the posterior and lateral tracts, resembling subacute combined degeneration (see below).

SYRINGOMYELIA

Syringomyelia is a developmental cavity of the cervical cord that may enlarge and produce progressive myelopathy or may remain asymptomatic. Symptoms begin insidiously in adolescence or early adulthood, progress irregularly, and may undergo spontaneous arrest for several years. Many young patients acquire a cervical-thoracic scoliosis. More than half of all cases are associated with Chiari type 1 malformations in which the cerebellar tonsils protrude through the foramen magnum and into the cervical spinal canal. The pathophysiology of syrinx expansion is controversial, but some interference with the normal flow of CSF seems likely, perhaps by the Chiari malformation. Acquired cavitations of the cord in areas of necrosis are also termed *syrinx cavities*; these follow trauma, myelitis, necrotic spinal cord tumors, and chronic arachnoiditis due to tuberculosis and other etiologies.

The presentation is a central cord syndrome consisting of a regional dissociated sensory loss (loss of pain and temperature sensation with sparing of touch and vibration) and areflexic weakness in the upper limbs. The sensory deficit has a distribution that is “suspended” over the nape of the neck, shoulders, and upper arms (cape distribution) or in the hands. Most cases begin asymmetrically with unilateral sensory loss in the hands that leads to injuries and burns that are not appreciated by the patient. Muscle wasting in the lower neck, shoulders, arms, and hands with asymmetric or absent reflexes in the arms reflects expansion of the cavity in the gray matter of the cord. As the cavity



FIGURE 456-7 Magnetic resonance imaging of syringomyelia associated with a Chiari malformation. Sagittal T1-weighted image through the cervical and upper thoracic spine demonstrates descent of the cerebellar tonsils below the level of the foramen magnum (*black arrows*). Within the substance of the cervical and thoracic spinal cord, a cerebrospinal fluid collection dilates the central canal (*white arrows*).

enlarges and compresses the long tracts, spasticity and weakness of the legs, bladder and bowel dysfunction, and a Horner’s syndrome appear. Some patients develop numbness and sensory loss from damage to the descending tract of the trigeminal nerve (C2 level or above). In cases with Chiari malformations, cough-induced headache and neck, arm, or facial pain may be reported. Extension of the syrinx into the medulla, syringobulbia, causes palatal or vocal cord paralysis, dysarthria, horizontal or vertical nystagmus, episodic dizziness or vertigo, and tongue weakness with atrophy.

MRI accurately identifies developmental and acquired syrinx cavities and their associated spinal cord enlargement (**Fig. 456-7**). Images of the brain and the entire spinal cord should be obtained to delineate the full longitudinal extent of the syrinx, assess posterior fossa structures for the Chiari malformation, and determine whether hydrocephalus is present.

TREATMENT SYRINGOMYELIA

Treatment of syringomyelia is generally unsatisfactory. The Chiari tonsillar herniation may be decompressed, generally by suboccipital craniectomy, upper cervical laminectomy, and placement of a dural graft. Fourth ventricular outflow is reestablished by this procedure. If the syrinx cavity is large, some surgeons recommend direct decompression or drainage by one of a number of methods, but the added benefit of this procedure is uncertain, and complications are common. With Chiari malformations, shunting of hydrocephalus generally precedes any attempt to correct the syrinx. Surgery may stabilize the neurologic deficit, and some patients improve. Patients with few symptoms and signs from the syrinx do not require surgery and are followed by serial clinical and imaging examinations.

Syrinx cavities secondary to trauma or infection, if symptomatic, are treated with a decompression and drainage procedure in which a small shunt is inserted between the cavity and subarachnoid space; alternatively, the cavity can be fenestrated. Cases due to intramedullary spinal cord tumor are generally managed by resection of the tumor.

CHRONIC MYELOPATHY OF MULTIPLE SCLEROSIS

A chronic progressive myelopathy is the most frequent cause of disability in both primary progressive and secondary progressive forms of MS. Involvement is typically bilateral but asymmetric and produces motor, sensory, and bladder/bowel disturbances. Fixed motor disability appears to result from extensive loss of axons in the corticospinal tracts. Diagnosis is facilitated by identification of earlier attacks such as optic neuritis. MRI, CSF, and evoked response testing are confirmatory. Disease-modifying therapy is indicated for patients with progressive myelopathy who also have coexisting MS relapses. Therapy is sometimes offered to patients who have a progressive course without relapses but with “active” MRI scans (e.g., the presence of new focal demyelinating lesions) despite the lack of evidence supporting the value of treatment in this setting. **MS is discussed in Chap. 458.**

SUBACUTE COMBINED DEGENERATION (VITAMIN B₁₂ DEFICIENCY)

This treatable myelopathy presents with subacute paresthesias in the hands and feet, loss of vibration and position sensation, and a progressive spastic and ataxic weakness. Loss of reflexes due to an associated peripheral neuropathy in a patient who also has Babinski signs is an important diagnostic clue. Optic atrophy and irritability or other cognitive changes may be prominent in advanced cases and are occasionally the presenting symptoms. The myelopathy of subacute combined degeneration tends to be diffuse rather than focal; signs are generally symmetric and reflect predominant involvement of the posterior and lateral tracts, including Romberg’s sign. The diagnosis is confirmed by the finding of macrocytic red blood cells, a low serum B₁₂ concentration, elevated serum levels of homocysteine and methylmalonic acid, and in uncertain cases, testing for anti-parietal cell antibodies and a Schilling test. Treatment is by replacement therapy, beginning with 1000 µg of intramuscular vitamin B₁₂ repeated at regular intervals or by subsequent oral treatment (**Chap. 128**).

HYPOCUPRIC MYELOPATHY

This myelopathy is similar to subacute combined degeneration (described above), except there is no neuropathy, and explains cases with normal serum levels of B₁₂. Low levels of serum copper are found, and often there is also a low level of serum ceruloplasmin. Some cases follow gastrointestinal procedures, particularly bariatric surgery, that result in impaired copper absorption; others have been associated with excess zinc from health food supplements or, until recently, zinc-containing denture creams, all of which impair copper absorption via induction of metallothionein, a copper-binding protein. Many cases are idiopathic. Improvement or at least stabilization may be expected with reconstitution of copper stores by oral supplementation. There is microcytic or macrocytic anemia. The pathophysiology and pathology of the idiopathic form are not known.

TABES DORSALIS

The classic syphilitic syndromes of tabes dorsalis and meningovascular inflammation of the spinal cord are now less frequent than in the past but must be considered in the differential diagnosis of spinal cord disorders. The characteristic symptoms of tabes are fleeting and repetitive lancinating pains, primarily in the legs or less often in the back, thorax, abdomen, arms, and face. Ataxia of the legs and gait due to loss of position sense occurs in half of patients. Paresthesias, bladder disturbances, and acute abdominal pain with vomiting (visceral crisis) occur in 15–30% of patients. The cardinal signs of tabes are loss of reflexes in the legs; impaired position and vibratory sense; Romberg’s sign; and, in almost all cases, bilateral Argyll Robertson pupils, which fail to constrict to light but accommodate. Diabetic polyradiculopathy may simulate tabes.

FAMILIAL SPASTIC PARAPLEGIA

Many cases of slowly progressive myelopathy are genetic in origin (**Chap. 452**). More than 30 different causative loci have been identified, including autosomal dominant, autosomal recessive, and X-linked forms. Especially for the recessive and X-linked forms, a

family history of myelopathy may be lacking. Most patients present with almost imperceptibly progressive spasticity and weakness in the legs, usually but not always symmetrical. Sensory symptoms and signs are absent or mild, but sphincter disturbances may be present. In some families, additional neurologic signs are prominent, including nystagmus, ataxia, or optic atrophy. The onset may be as early as the first year of life or as late as middle adulthood. Only symptomatic therapies are available.

ADRENOMYELONEUROPATHY

This X-linked disorder is a variant of adrenoleukodystrophy. Most affected males have a history of adrenal insufficiency and then develop a progressive spastic (or ataxic) paraparesis beginning in early or sometimes middle adulthood; some patients also have a mild peripheral neuropathy. Female heterozygotes may develop a slower, insidiously progressive spastic myelopathy beginning later in adulthood and without adrenal insufficiency. Diagnosis is usually made by demonstration of elevated levels of very-long-chain fatty acids in plasma and in cultured fibroblasts. The responsible gene encodes the adrenoleukodystrophy protein (ADLP), a peroxisomal membrane transporter involved in carrying long-chain fatty acids to peroxisomes for degradation. Corticosteroid replacement is indicated if hypoadrenalism is present, and bone marrow transplantation and nutritional supplements have been attempted for this condition without clear evidence of efficacy.

OTHER CHRONIC MYELOPATHIES

Primary lateral sclerosis (**Chap. 452**) is a degenerative disorder characterized by progressive spasticity with weakness, eventually accompanied by dysarthria and dysphonia; bladder symptoms occur in approximately half of patients. Sensory function is spared. The disorder resembles ALS and is considered a variant of the motor neuron degenerations, but without the characteristic lower motor neuron disturbance. Some cases may represent familial spastic paraplegia, particularly autosomal recessive or X-linked varieties in which a family history may be absent.

Tethered cord syndrome is a developmental disorder of the lower spinal cord and nerve roots that rarely presents in adulthood as low back pain accompanied by a progressive lower spinal cord and/or nerve root syndrome. Some patients have a small leg or foot deformity indicating a long-standing process, and in others, a dimple, patch of hair, or sinus tract on the skin overlying the lower back is the clue to a congenital lesion. Diagnosis is made by MRI, which demonstrates a low-lying conus medullaris and thickened filum terminale. The MRI may also reveal diastematomyelia (division of the lower spinal cord into two halves), lipomas, cysts, or other congenital abnormalities of the lower spine coexisting with the tethered cord. Treatment is with surgical release.

There are a number of rare toxic causes of spastic myelopathy, including lathyrism due to ingestion of chickpeas containing the excitotoxin β-N-oxalylamino-L-alanine (BOAA), seen primarily in the developing world, and nitrous oxide inhalation producing a myelopathy identical to subacute combined degeneration. SLE, Sjögren’s syndrome, and sarcoidosis may each cause a myelopathy without overt evidence of systemic disease. Cancer-related causes of chronic myelopathy, besides the common neoplastic compressive myelopathy discussed earlier, include radiation injury (**Chap. 118**) and rare paraneoplastic myelopathies. The last of these are most often associated with lung or breast cancer and anti-Hu antibodies (**Chap. 122**) or with lymphoma that causes a syndrome of destruction of anterior horn cells; NMO (**Chap. 458**) can also rarely be paraneoplastic in origin. Metastases to the cord are probably more common than either of these in patients with cancer. Often, a cause of intrinsic myelopathy can be identified only through periodic reassessment.

REHABILITATION OF SPINAL CORD DISORDERS

The prospects for recovery from an acute destructive spinal cord lesion fade after ~6 months. There are currently no effective means to

TABLE 456-4 EXPECTED NEUROLOGIC FUNCTION FOLLOWING COMPLETE CORD LESIONS

Level	Self-Care	Transfers	Maximum Mobility
High quadriplegia (C1-C4)	Dependent on others; requires respiratory support	Dependent on others	Motorized wheelchair
Low quadriplegia (C5-C8)	Partially independent with adaptive equipment	May be dependent or independent	May use manual wheelchair, drive an automobile with adaptive equipment
Paraplegia (below T1)	Independent	Independent	Ambulates short distances with aids

Source: Adapted from JF Ditunno, CS Formal: Chronic spinal cord injury. *N Engl J Med* 330:550, 1994; with permission.

promote repair of injured spinal cord tissue; promising but entirely experimental approaches include the use of factors that influence reinnervation by axons of the corticospinal tract, nerve and neural sheath graft bridges, forms of electrical stimulation at the site of injury, and the local introduction of stem cells. The disability associated with irreversible spinal cord damage is determined primarily by the level of the lesion and by whether the disturbance in function is complete or incomplete (Table 456-4). Even a complete high cervical cord lesion may be compatible with a productive life. The primary goals are development of a rehabilitation plan framed by realistic expectations and attention to the neurologic, medical, and psychological complications that commonly arise.

Many of the usual symptoms associated with medical illnesses, especially somatic and visceral pain, may be lacking because of the destruction of afferent pain pathways. Unexplained fever, worsening of spasticity, or deterioration in neurologic function should prompt a search for infection, thrombophlebitis, or an intraabdominal pathology. The loss of normal thermoregulation and inability to maintain normal body temperature can produce recurrent fever (*quadriplegic fever*), although most episodes of fever are due to infection of the urinary tract, lung, skin, or bone.

Bladder dysfunction generally results from loss of supraspinal innervation of the detrusor muscle of the bladder wall and the sphincter musculature. Detrusor spasticity is treated with anticholinergic drugs (oxybutynin, 2.5–5 mg qid) or tricyclic antidepressants with anticholinergic properties (imipramine, 25–200 mg/d). Failure of the sphincter muscle to relax during bladder emptying (urinary dyssynergia) may be managed with the α -adrenergic blocking agent terazosin hydrochloride (1–2 mg tid or qid), with intermittent catheterization, or, if that is not feasible, by use of a condom catheter in men or a permanent indwelling catheter. Surgical options include the creation of an artificial bladder by isolating a segment of intestine that can be catheterized intermittently (enterocystoplasty) or can drain continuously to an external appliance (urinary conduit). Bladder areflexia due to acute spinal shock or conus lesions is best treated by catheterization. Bowel regimens and disimpaction are necessary in most patients to ensure at least biweekly evacuation and avoid colonic distention or obstruction.

Patients with acute cord injury are at risk for venous thrombosis and pulmonary embolism. Use of calf-compression devices and anticoagulation with low-molecular-weight heparin is recommended. In cases of persistent paralysis, anticoagulation should probably be continued for 3 months.

Prophylaxis against decubitus ulcers should involve frequent changes in position in a chair or bed, the use of special mattresses, and cushioning of areas where pressure sores often develop, such as the sacral prominence and heels. Early treatment of ulcers with careful cleansing, surgical or enzyme debridement of necrotic tissue, and appropriate dressing and drainage may prevent infection of adjacent soft tissue or bone.

Spasticity is aided by stretching exercises to maintain mobility of joints. Drug treatment is effective but may result in reduced function, as some patients depend on spasticity as an aid to stand, transfer, or walk. Baclofen (up to 240 mg/d in divided doses) is effective; it acts by facilitating γ -aminobutyric acid–mediated inhibition of motor reflex arcs. Diazepam acts by a similar mechanism and is useful for leg spasms that interrupt sleep (2–4 mg at bedtime). Tizanidine (2–8 mg tid), an α_2 adrenergic agonist that increases presynaptic inhibition

of motor neurons, is another option. For nonambulatory patients, the direct muscle inhibitor dantrolene (25–100 mg qid) may be used, but it is potentially hepatotoxic. In refractory cases, intrathecal baclofen administered via an implanted pump, botulinum toxin injections, or dorsal rhizotomy may be required to control spasticity.

Despite the loss of sensory function, many patients with spinal cord injury experience chronic pain sufficient to diminish their quality of life. Randomized controlled studies indicate that gabapentin or pregabalin is useful in this setting. Epidural electrical stimulation and intrathecal infusion of pain medications have been tried with some success. **Management of chronic pain is discussed in Chap. 18.**

A paroxysmal autonomic hyperreflexia may occur following lesions above the major splanchnic sympathetic outflow at T6. Headache, flushing, and diaphoresis above the level of the lesion, as well as hypertension with bradycardia or tachycardia, are the major symptoms. The trigger is typically a noxious stimulus—for example, bladder or bowel distention, a urinary tract infection, or a decubitus ulcer—below the level of the cord lesion. Treatment consists of removal of offending stimuli; ganglionic blocking agents (mecamylamine, 2.5–5 mg) or other short-acting antihypertensive drugs are useful in some patients.

Attention to these details allows longevity and a productive life for patients with complete transverse myelopathies.

457e Concussion and Other Traumatic Brain Injuries

Allan H. Ropper

This is a digital-only chapter. It is available on the DVD that accompanies this book, as well as on Access Medicine/Harrison's Online, and the eBook and "app" editions of HPIM 19e.

Almost 10 million head injuries occur annually in the United States, about 20% of which are serious enough to cause brain damage. Among men <35 years, accidents, usually motor vehicle collisions, are the chief cause of death and >70% of these involve head injury. Furthermore, minor head injuries are so common that almost all physicians will be called upon to provide immediate care or to see patients who are suffering from various sequelae.

Medical personnel caring for head injury patients should be aware that (1) spinal injury often accompanies head injury, and care must be taken in handling the patient to prevent compression of the spinal cord due to instability of the spinal column; (2) intoxication is frequently associated with traumatic brain injury, and thus testing for drugs and alcohol should be carried out when appropriate; and (3) additional injuries, including rupture of abdominal organs, may produce vascular collapse, shock, or respiratory distress that requires immediate attention.

457e Concussion and Other Traumatic Brain Injuries

Allan H. Ropper

Almost 10 million head injuries occur annually in the United States, about 20% of which are serious enough to cause brain damage. Among men <35 years, accidents, usually motor vehicle collisions, are the chief cause of death and >70% of these involve head injury. Furthermore, minor head injuries are so common that almost all physicians will be called upon to provide immediate care or to see patients who are suffering from various sequelae.

Medical personnel caring for head injury patients should be aware that (1) spinal injury often accompanies head injury, and care must be taken in handling the patient to prevent compression of the spinal cord due to instability of the spinal column; (2) intoxication is frequently associated with traumatic brain injury, and thus testing for drugs and alcohol should be carried out when appropriate; and (3) additional injuries, including rupture of abdominal organs, may produce vascular collapse, shock, or respiratory distress that requires immediate attention.

TYPES OF HEAD INJURIES

CONCUSSION

This form of minor head injury had in the past referred to an immediate and transient loss of consciousness that was associated with a short period of amnesia. Many patients, however, do not lose consciousness after a minor head injury but instead are dazed or confused, or feel stunned or “star struck,” and the term *concussion* is now applied to all such cognitive and perceptual changes experienced after a blow to the head. Severe concussion may precipitate a brief convulsion or autonomic signs such as facial pallor, bradycardia, faintness with mild hypotension, or sluggish pupillary reaction, but most patients quickly return to a neurologically normal state.

The mechanics of a typical concussion involve sudden deceleration of the head when hitting a blunt stationary object. This creates an anterior-posterior movement of the brain within the skull due to inertia and rotation of the cerebral hemispheres on the fulcrum of the relatively fixed upper brainstem. Loss of consciousness in concussion is believed to result from a transient electrophysiologic dysfunction of the reticular activating system in the upper midbrain that is at the site of rotation (Chap. 328). The transmission of a wave of kinetic energy throughout the brain is an alternative explanation for the disruption in consciousness.

Gross and light-microscopic changes in the brain are usually absent following concussion, but biochemical and ultrastructural changes, such as mitochondrial ATP depletion and local disruption of the blood-brain barrier, may be transient abnormalities. Computed tomography (CT) and magnetic resonance imaging (MRI) scans are usually normal; however, a small number of patients will be found to have a skull fracture, an intracranial hemorrhage, or a brain contusion.

A brief period of both retrograde and anterograde amnesia is characteristic of concussion, and it recedes rapidly in alert patients. Memory loss spans the moments before impact but may encompass the previous days or weeks (rarely months). With severe injuries, the extent of retrograde amnesia roughly correlates with the severity of injury. Memory is regained erratically from the most distant to more recent memories, with islands of amnesia occasionally remaining. The mechanism of amnesia is not known. Hysterical posttraumatic amnesia is not uncommon after head injury and should be suspected when inexplicable behavioral abnormalities occur, such as recounting events that cannot be recalled on later testing, a bizarre affect, forgetting one's own name, or a persistent anterograde deficit that is excessive in comparison with the degree of injury. **Amnesia is discussed in Chap. 36.**

A single, uncomplicated concussion only infrequently produces permanent neurobehavioral changes in patients who are free of

preexisting psychiatric and neurologic problems. Nonetheless, residual problems in memory and concentration may have an anatomic correlate in microscopic cerebral lesions (see below).

The mechanisms by which a blast injury affects the brain and causes symptoms that are associated with concussion, a problem mainly in military medicine, are not known. The energy of a blast wave can enter the cranium through the openings of the orbits, auditory canals, and foramen magnum. There are not consistent changes in cerebral imaging studies but more subtle indications of tissue disruption have been found, comparable to those of mild concussion. It has been difficult to separate the direct effects of the blast from the consequences of being thrown against fixed objects or injured by flying debris.

CONTUSION, BRAIN HEMORRHAGE, AND AXONAL SHEARING LESIONS

These pathologic changes are the result of severe cranial trauma. A surface bruise of the brain, or *contusion*, consists of varying degrees of petechial hemorrhage, edema, and tissue destruction. Contusions and deeper hemorrhages result from mechanical forces that displace and compress the hemispheres forcefully and by deceleration of the brain against the inner skull, either under a point of impact (coup lesion) or, as the brain swings back, in the antipolar area (contrecoup lesion). Trauma sufficient to cause prolonged unconsciousness usually produces some degree of contusion. Blunt deceleration impact, as occurs against an automobile dashboard or from falling forward onto a hard surface, causes contusions on the orbital surfaces of the frontal lobes and the anterior and basal portions of the temporal lobes. With lateral forces, as from impact on an automobile door frame, contusions are situated on the lateral convexity of the hemisphere. The clinical signs of contusion are determined by the location and size of the lesion; often, there are no focal neurologic abnormalities, but these injured regions are later the sites of gliotic scars that may produce seizures. A hemiparesis or gaze preference is fairly typical of moderately sized contusions. Large bilateral contusions produce stupor with extensor posturing, while those limited to the frontal lobes cause a taciturn state. Contusions in the temporal lobe may cause delirium or an aggressive, combative syndrome.

Acute contusions are easily visible on CT and MRI scans, appearing as inhomogeneous hyperdensities on CT and as hyperintensities on T2 and fluid-attenuated inversion recovery (FLAIR) MRI sequences; there is usually surrounding localized brain edema (Fig. 457e-1) and some subarachnoid bleeding. Blood in the cerebrospinal fluid (CSF) due to trauma may provoke a mild inflammatory reaction. Over a few days, contusions acquire a surrounding contrast enhancement and edema that may be mistaken for tumor or abscess. Glial and macrophage reactions result in chronic, scarred, hemosiderin-stained depressions on the cortex (*plaques jaunes*) that are the main source of posttraumatic epilepsy.



FIGURE 457e-1 Traumatic cerebral contusion. Noncontrast computed tomography scan demonstrating a hyperdense hemorrhagic region in the anterior temporal lobe.

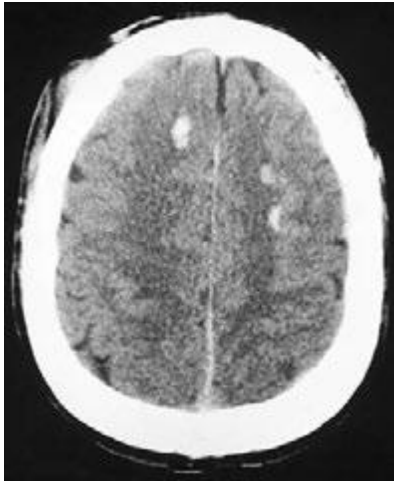


FIGURE 457e-2 Multiple small areas of hemorrhage and tissue disruption in the white matter of the frontal lobes on noncontrast computed tomography scan. These appear to reflect an extreme type of the diffuse axonal shearing lesions that occur with closed head injury.

Torsional or shearing forces within the brain cause hemorrhages of the basal ganglia and other deep regions. Large hemorrhages after minor trauma suggest that there is a bleeding diathesis or cerebrovascular amyloidosis. For unexplained reasons, deep cerebral hemorrhages may not develop until several days after injury. Sudden neurologic deterioration in a comatose patient or a sudden rise in intracranial pressure (ICP) suggests this complication has occurred and should therefore prompt investigation with a CT scan.

A special type of deep white matter lesion consists of widespread mechanical disruption, or shearing, of axons at the time of impact. Most characteristic are small areas of tissue injury in the corpus callosum and dorsolateral pons. The presence of widespread multifocal axonal damage in both hemispheres, a state called *diffuse axonal injury* (DAI), has been proposed to explain persistent coma and the vegetative state after closed head injury (**Chap. 328**), but small ischemic-hemorrhagic lesions in the midbrain and thalamus are an alternative explanation. Only severe shearing lesions that contain blood are visualized by CT, usually in the corpus callosum and centrum semiovale (**Fig. 457e-2**); however, special MRI sequences that detect small amounts of blood and diffusion tensor imaging can demonstrate numerous such lesions throughout the white matter.

SKULL FRACTURES

A blow to the skull that exceeds the elastic tolerance of the bone causes a fracture. Intracranial lesions accompany roughly two-thirds of skull fractures, and the presence of a fracture increases manyfold the chances of an underlying subdural or epidural hematoma. Consequently, fractures are primarily markers of the site and severity of injury. If the underlying arachnoid membrane has been torn, fractures also provide potential pathways for entry of bacteria to the CSF with a risk of meningitis and for leakage of CSF outward through the dura. If there is leakage of CSF, severe orthostatic headache results from lowered pressure in the spinal fluid compartment.

Most fractures are linear and extend from the point of impact toward the base of the skull. Basilar skull fractures are often extensions of adjacent linear fractures over the convexity of the skull but may occur independently owing to stresses on the floor of the middle cranial fossa or occiput. Basilar fractures are usually parallel to the petrous bone or along the sphenoid bone and directed toward the sella turcica and ethmoidal groove. Although most basilar fractures are uncomplicated, they can cause CSF leakage, pneumocephalus, and delayed cavernous-carotid fistulas. Hemotympanum (blood behind the tympanic membrane), ecchymosis over the mastoid process (Battle sign), and periorbital ecchymosis (“raccoon sign”) are associated with basilar fractures. Because routine x-ray examination may fail

to disclose basilar fractures, they should be suspected if these clinical signs are present.

CSF may leak through the cribriform plate or the adjacent sinus and cause CSF rhinorrhea (a watery discharge from the nose). Persistent rhinorrhea and recurrent meningitis usually require surgical repair of torn dura underlying the fracture. The site of the leak is often difficult to determine, but useful diagnostic tests include the instillation of water-soluble contrast into the CSF followed by CT with the patient in various positions, or injection of radionuclide compounds or fluorescein into the CSF and the insertion of absorptive nasal pledgets. The location of an intermittent leak is infrequently delineated, and many resolve spontaneously.

Sellar fractures, even those associated with serious neuroendocrine dysfunction, may be radiologically occult or evident only by an air-fluid level in the sphenoid sinus. Fractures of the dorsum sella cause sixth or seventh nerve palsies or optic nerve damage.

Petrous bone fractures, especially those oriented along the long axis of the bone, may be associated with facial palsy, disruption of ear ossicles, and CSF otorrhea. Transverse petrous fractures are less common; they almost always damage the cochlea or labyrinths and often the facial nerve as well. External bleeding from the ear is usually from local abrasion of the external canal but can also result from petrous fracture.

Fractures of the frontal bone are usually depressed, involving the frontal and paranasal sinuses and the orbits. Depressed skull fractures are typically compound, but they may be asymptomatic because the impact energy is dissipated in breaking the bone; some have underlying brain contusions. Debridement and exploration of compound fractures are required in order to avoid infection; simple fractures usually do not require surgery.

CRANIAL NERVE INJURIES

The cranial nerves most often injured with head trauma are the olfactory, optic, oculomotor, and trochlear; the first and second branches of the trigeminal nerve; and the facial and auditory nerves. Anosmia and an apparent loss of taste (actually a loss of perception of aromatic flavors, with retained elementary taste perception) occur in ~10% of persons with serious head injuries, particularly from falls on the back of the head. This is the result of displacement of the brain and shearing of the fine olfactory nerve filaments that course through the cribriform bone. At least partial recovery of olfactory and gustatory function is expected, but if bilateral anosmia persists for several months, the prognosis is poor. Partial optic nerve injuries from closed trauma result in blurring of vision, central or paracentral scotomas, or sector defects. Direct orbital injury may cause short-lived blurred vision for close objects due to reversible iridoplegia. Diplopia limited to downward gaze and corrected when the head is tilted away from the side of the affected eye indicates trochlear (fourth nerve) nerve damage. It occurs frequently as an isolated problem after minor head injury or may develop for unknown reasons after a delay of several days. Facial nerve injury caused by a basilar fracture is present immediately in up to 3% of severe injuries; it may also be delayed for 5–7 days. Fractures through the petrous bone, particularly the less common transverse type, are liable to produce facial palsy. Delayed facial palsy occurring up to a week after injury, the mechanism of which is unknown, has a good prognosis. Injury to the eighth cranial nerve from a fracture of the petrous bone causes loss of hearing, vertigo, and nystagmus immediately after injury. Deafness from eighth nerve injury is rare and must be distinguished from blood in the middle ear or disruption of the middle ear ossicles. Dizziness, tinnitus, and high-tone hearing loss occur from cochlear concussion, most typically after blast injury.

SEIZURES

Convulsions are surprisingly uncommon immediately after a head injury, but a brief period of tonic extensor posturing or a few clonic movements of the limbs just after the moment of impact can occur. However, the cortical scars that evolve from contusions are highly epileptogenic and may later manifest as seizures, even after many months or years (**Chap. 445**). The severity of injury roughly determines the

risk of future seizures. It has been estimated that 17% of individuals with brain contusion, subdural hematoma, or prolonged loss of consciousness will develop a seizure disorder and that this risk extends for an indefinite period of time, whereas the risk is $\leq 2\%$ after mild injury. The majority of convulsions in the latter group occur within 5 years of injury but may be delayed for decades. Penetrating injuries have a much higher rate of subsequent epilepsy.

SUBDURAL AND EPIDURAL HEMATOMAS

Hemorrhages beneath the dura (subdural) or between the dura and skull (epidural) have characteristic clinical and imaging features. They are sometimes associated with underlying contusions and other injuries, often making it difficult to determine the relative contribution of each component to the clinical state. The mass effect and raised ICP caused by these hematomas can be life threatening, making it imperative to identify them rapidly by CT or MRI scan and to remove them when appropriate.

Acute Subdural Hematoma (Fig. 457e-3) Direct cranial trauma may be minor and is not required for acute subdural hemorrhage to occur, especially in the elderly and those taking anticoagulant medications. Acceleration forces alone, as from whiplash, are sometimes sufficient to produce subdural hematoma. Up to one-third of patients have a lucid interval lasting minutes to hours before coma supervenes, but most are drowsy or comatose from the moment of injury. A unilateral headache and slightly enlarged pupil on the side of the hematoma are frequently, but not invariably, present. Stupor or coma, hemiparesis, and unilateral pupillary enlargement are signs of larger hematomas. In an acutely deteriorating patient, burr (drainage) holes or an emergency craniotomy are required. Small subdural hematomas may be asymptomatic and usually do not require evacuation if they do not enlarge.

A subacutely evolving syndrome due to subdural hematoma occurs days or weeks after injury with drowsiness, headache, confusion, or mild hemiparesis, usually in alcoholics and in the elderly and often after only minor trauma. On imaging studies, subdural hematomas appear as crescentic collections over the convexity of one or both hemispheres, most commonly in the frontotemporal region, and less often in the inferior middle fossa or over the occipital poles (Fig. 457e-3). Interhemispheric, posterior fossa, or bilateral convexity hematomas are less frequent and are difficult to diagnose clinically, although drowsiness and the neurologic signs expected from damage in each region can usually be detected. The bleeding that causes larger hematomas is primarily venous in origin, although additional arterial bleeding sites are sometimes found at operation, and a few large hematomas have a purely arterial origin.



FIGURE 457e-3 Acute subdural hematoma. Noncontrast computed tomography scan reveals a hyperdense clot that has an irregular border with the brain and causes more horizontal displacement (mass effect) than might be expected from its thickness. The disproportionate mass effect is the result of the large rostral-caudal extent of these hematomas. Compare to Fig. 457e-4.

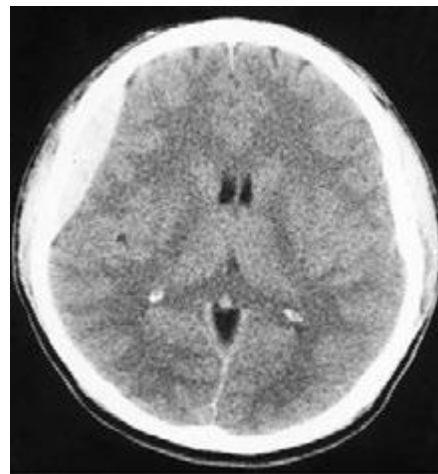


FIGURE 457e-4 Acute epidural hematoma. The tightly attached dura is stripped from the inner table of the skull, producing a characteristic lenticular-shaped hemorrhage on noncontrast computed tomography scan. Epidural hematomas are usually caused by tearing of the middle meningeal artery following fracture of the temporal bone.

Epidural Hematoma (Fig. 457e-4) These usually evolve more rapidly than subdural hematomas and are correspondingly more treacherous. They occur in up to 10% of cases of severe head injury but are associated with underlying cortical damage less often than for subdural hematomas. Most patients are unconscious when first seen. A “lucid interval” of several minutes to hours before coma supervenes is most characteristic of epidural hemorrhage, but it is still uncommon, and epidural hemorrhage is not the only cause of this temporal sequence. Rapid surgical evacuation and ligation or cauterization of the damaged vessel is indicated, usually the middle meningeal artery that has been lacerated by an overlying skull fracture.

Chronic Subdural Hematoma (Fig. 457e-5) A subacutely evolving syndrome due to subdural hematoma occurs days or weeks after injury with drowsiness, headache, confusion, or mild hemiparesis, usually in alcoholics and in the elderly and often after only minor or unnoticed trauma. On imaging studies, chronic subdural hematomas appear as crescentic clots over the convexity of one or both hemispheres, most commonly in the frontotemporal region (Fig. 457e-3). A history of

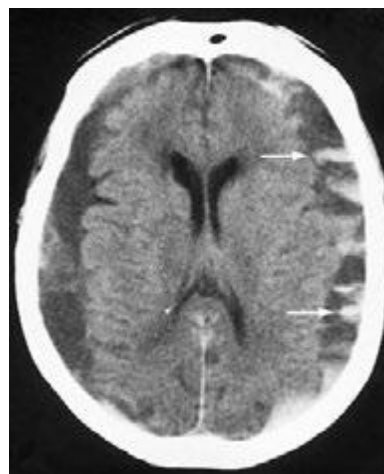


FIGURE 457e-5 Computed tomography scan of chronic bilateral subdural hematomas of different ages. The collections began as acute hematomas and have become hypodense in comparison to the adjacent brain after a period during which they were isodense and difficult to appreciate. Some areas of resolving blood are contained on the more recently formed collection on the left (arrows).

trauma may or may not be elicited in relation to chronic subdural hematoma; the injury may have been trivial and forgotten, particularly in the elderly and those with clotting disorders. Headache is common but not invariable. Additional features that may appear weeks later include slowed thinking, vague change in personality, seizure, or a mild hemiparesis. Headache fluctuates in severity, sometimes with changes in head position.

Bilateral chronic subdural hematomas produce perplexing clinical syndromes, and the initial clinical impression may be of a stroke, brain tumor, drug intoxication, depression, or a dementing illness. Drowsiness, inattentiveness, and incoherence of thought are generally more prominent than focal signs such as hemiparesis. Rarely, chronic hematomas cause brief episodes of hemiparesis or aphasia that are indistinguishable from transient ischemic attacks. Patients with undetected bilateral subdural hematomas have a low tolerance for surgery, anesthesia, and drugs that depress the nervous system; drowsiness or confusion persists for long periods postoperatively.

CT without contrast initially shows a low-density mass over the convexity of the hemisphere (Fig. 457e-5). Between 2 and 6 weeks after the initial bleeding, the clot becomes isodense compared to adjacent brain and may be inapparent. Many subdural hematomas that are several weeks in age contain areas of blood and intermixed serous fluid. Bilateral chronic hematomas may fail to be detected because of the absence of lateral tissue shifts; this circumstance in an older patient is suggested by a “hypernormal” CT scan with fullness of the cortical sulci and small ventricles. Infusion of contrast material demonstrates enhancement of the vascular fibrous capsule surrounding the collection. MRI reliably identifies subacute and chronic hematomas.

Clinical observation coupled with serial imaging is a reasonable approach to patients with few symptoms, such as headache alone, and in those with small chronic subdural collections. Treatment of minimally symptomatic chronic subdural hematoma with glucocorticoids is favored by some clinicians, but surgical evacuation is more often successful. The fibrous membranes that grow from the dura and encapsulate the collection require removal to prevent recurrent fluid accumulation. Small hematomas are resorbed, leaving only the organizing membranes. On imaging studies, very chronic subdural hematomas are difficult to distinguish from hygromas, which are collections of CSF from a rent in the arachnoid membrane.

CLINICAL SYNDROMES AND TREATMENT OF HEAD INJURY

MINOR INJURY

The patient who has briefly lost consciousness or been stunned after a minor head injury usually becomes fully alert and attentive within minutes but may complain of headache, dizziness, faintness, nausea, a single episode of emesis, difficulty with concentration, a brief amnesic period, or slight blurring of vision. This typical concussion syndrome has a good prognosis with little risk of subsequent deterioration. Children are particularly prone to drowsiness, vomiting, and irritability, symptoms that are sometimes delayed for several hours after apparently minor injuries. Vasovagal syncope that follows injury may cause undue concern. Generalized or frontal headache is common in the following days. It may be migrainous (throbbing and hemicranial) in nature or aching and bilateral. After several hours of observation, patients with minor injury may be accompanied home and observed for a day by a family member or friend, with written instructions to return if symptoms worsen.

Persistent severe headache and repeated vomiting in the context of normal alertness and no focal neurologic signs is usually benign, but CT should be obtained and a longer period of observation is appropriate. The decision to perform imaging tests also depends on clinical signs that indicate that the impact was severe (e.g., persistent confusion, periorbital or mastoid hematoma, repeated vomiting, palpable skull fracture), on the seriousness of other bodily injuries, and on the degree of surveillance that can be anticipated after discharge. Two studies have indicated that older age, two or more episodes of vomiting, >30 min of retrograde or persistent anterograde amnesia, seizure, and concurrent drug or alcohol intoxication are sensitive

(but not specific) indicators of intracranial hemorrhage that justify CT scanning. It may be appropriate to be more liberal in obtaining CT scans in children because a small number, even without loss of consciousness, will have intracranial traumatic lesions but this exposes the child to radiation.

Concussion in Sports In the current absence of adequate data, a common sense approach to athletic concussion has been to remove the individual from play immediately and avoid contact sports for at least several days after a mild injury and for a longer period if there are more severe injuries or if there are protracted neurologic symptoms such as headache and difficulty concentrating. No individual should return to play unless all symptoms have resolved and an assessment has been made by a health care professional who has experience with treatment of concussion. Once cleared, the individual can then begin a graduated program of increasing activity. Younger athletes are particularly likely to experience protracted concussive symptoms, and a slower return to play in this age group may be reasonable. These guidelines are designed in part to avoid a perpetuation of symptoms but also to prevent the rare *second impact syndrome*, in which diffuse and fatal cerebral swelling follows a second minor head injury.

In the past, mental decline in boxers late in their careers had been called *dementia pugilistica*. There is some evidence that repeated concussions from other sports are associated with a similar delayed and progressive cognitive disorder that is due mainly to the deposition of tau protein in cortical neurons. The brains of these patients display deposition of tau protein in the superficial cortical layers, and particularly in the depths of sulci within the frontal cortices, a pattern named *chronic traumatic encephalopathy* (CTE) that is quite unlike other degenerative conditions. CTE is an intensively studied and provocative entity. Its contribution, if any, to late-life dementia and parkinsonism in former athletes, soldiers, or others who have sustained repeated concussive injuries is unknown. **CTE is also discussed in Chap. 444.**

INJURY OF INTERMEDIATE SEVERITY

Patients who are not fully alert or have persistent confusion, behavioral changes, extreme dizziness, or focal neurologic signs such as hemiparesis should be admitted to the hospital and have cerebral imaging. A cerebral contusion or hematoma will usually be found. Common syndromes include: (1) delirium with a disinclination to be examined or moved, expletive speech, and resistance if disturbed (anterior temporal lobe contusions); (2) a quiet, disinterested, slowed mental state (abulia) alternating with irascibility (inferior frontal and frontopolar contusions); (3) a focal deficit such as aphasia or mild hemiparesis (due to subdural hematoma or convexity contusion or, less often, carotid artery dissection); (4) confusion and inattention, poor performance on simple mental tasks, and fluctuating orientation (associated with several types of injuries, including those described above, and with medial frontal contusions and interhemispheric subdural hematoma); (5) repetitive vomiting, nystagmus, drowsiness, and unsteadiness (labyrinthine concussion, but occasionally due to a posterior fossa subdural hematoma or vertebral artery dissection); and (6) diabetes insipidus (damage to the median eminence or pituitary stalk). Injuries of this degree are often complicated by drug or alcohol intoxication, and clinically inapparent cervical spine injury may be present. Blast injuries are often accompanied by rupture of the tympanic membranes.

After surgical removal of hematomas, most patients in this category improve over weeks. During the first week, the state of alertness, memory, and other cognitive functions often fluctuate, and agitation and somnolence are common. Behavioral changes tend to be worse at night, as with many other encephalopathies, and may be treated with small doses of antipsychotic medications. Subtle abnormalities of attention, intellect, spontaneity, and memory return toward normal weeks or months after the injury, sometimes abruptly. Persistent cognitive problems are discussed below.

SEVERE INJURY

Patients who are comatose from the moment of injury require immediate neurologic attention and resuscitation. After intubation, with care taken to immobilize the cervical spine, the depth of coma,

pupillary size and reactivity, limb movements, and Babinski responses are assessed. As soon as vital functions permit and cervical spine x-rays and a CT scan have been obtained, the patient should be transported to a critical care unit. Hypoxia should be reversed, and normal saline used as the resuscitation fluid in preference to albumin. The finding of an epidural or subdural hematoma or large intracerebral hemorrhage is usually an indication for prompt surgery and intracranial decompression in an otherwise salvageable patient. Measurement of ICP with a ventricular catheter or fiberoptic device in order to guide treatment has been favored by many units but has not improved outcome. Hyperosmolar intravenous solutions are used in various regimens to limit intracranial pressure. The inherently appealing approach of removing portions of the skull in order to decompress the intracranial contents, as has been successful for brain swelling after cerebral infarction, has so far not proven effective for traumatic brain injury. The use of prophylactic antiepileptic medications has been recommended, but there is little supportive data. **Management of raised ICP, a frequent feature of severe head injury, is discussed in Chap. 330.**

GRADING AND PROGNOSIS

In severe head injury, the clinical features of eye opening, motor responses of the limbs, and verbal output have been found to be generally predictive of outcome. These three responses are assessed by the Glasgow Coma Scale; a score between 3 and 15 is assigned (Table 457e-1). Over 85% of patients with aggregate scores of <5 die within 24 h. However, a number of patients with slightly higher scores, including a few without pupillary light responses, survive, suggesting that an initially aggressive approach is justified in most patients. Patients <20 years old, particularly children, may make remarkable recoveries after having grave early neurologic signs. In one large study of severe head injury, 55% of children had a good outcome at 1 year, compared with 21% of adults. Older age, increased ICP, early hypoxia or hypotension, compression of the brainstem on CT or MRI, and a delay in the evacuation of large intracranial hemorrhages are indicators of a poor prognosis.

POSTCONCUSSION SYNDROME

The *postconcussion syndrome* refers to a state following minor head injury consisting of combinations of fatigue, dizziness, headache, and difficulty in concentration. The syndrome simulates asthenia and anxious depression. Based on experimental models, it has been proposed that subtle axonal shearing lesions or as yet undefined biochemical alterations account for the cognitive symptoms. In moderate and severe trauma, neuropsychological changes such as difficulty with attention and memory and other cognitive deficits are undoubtedly present, sometimes severe, but many problems identified by formal

TABLE 457e-1 GLASGOW COMA SCALE FOR HEAD INJURY

Eye Opening (E)		Verbal Response (V)	
Spontaneous	4	Oriented	5
To loud voice	3	Confused, disoriented	4
To pain	2	Inappropriate words	3
Nil	1	Incomprehensible sounds	2
		Nil	1
Best Motor Response (M)			
Obeys	6		
Localizes	5		
Withdraws (flexion)	4		
Abnormal flexion posturing	3		
Extension posturing	2		
Nil	1		

Note: Coma score = E + M + V. Patients scoring 3 or 4 have an 85% chance of dying or remaining vegetative, whereas scores >11 indicate only a 5–10% likelihood of death or vegetative state and 85% chance of moderate disability or good recovery. Intermediate scores correlate with proportional chances of recovery.

testing do not affect daily functioning. Test scores tend to improve rapidly during the first 6 months after injury and then more slowly for years.

Management of the postconcussive syndrome requires the identification and treatment of each separate element of depression, sleeplessness, anxiety, persistent headache, and dizziness. A clear explanation of the problems that may follow concussion has been shown to reduce subsequent complaints. Care is taken to avoid prolonged use of drugs that produce dependence. Headache may initially be treated with acetaminophen and small doses of amitriptyline. Vestibular exercises (Chap. 28) and small doses of vestibular suppressants such as promethazine (Phenergan) may be helpful when dizziness is the main problem. Patients who after minor or moderate injury have difficulty with memory or with complex cognitive tasks at work may be reassured that these problems usually improve over 6–12 months, and workload may be reduced in the interim. It is sometimes helpful to obtain serial and quantified neuropsychological testing in order to adjust the work environment to the patient's abilities and to document improvement over time. Whether cognitive exercises are useful in contrast to rest and a reduction in mental challenges is uncertain. Previously energetic and resilient individuals usually have the best recoveries. In patients with persistent symptoms, the possibility exists of malingering or prolongation as a result of litigation.

458 Multiple Sclerosis and Other Demyelinating Diseases

Stephen L. Hauser, Douglas S. Goodin

Demyelinating disorders are immune-mediated conditions characterized by preferential destruction of central nervous system (CNS) myelin. The peripheral nervous system (PNS) is spared, and most patients have no evidence of an associated systemic illness. Multiple sclerosis, the most common disease in this category, is second only to trauma as a cause of neurologic disability beginning in early to middle adulthood.

MULTIPLE SCLEROSIS

Multiple sclerosis (MS) is an autoimmune disease of the CNS characterized by chronic inflammation, demyelination, gliosis (scarring), and neuronal loss; the course can be relapsing-remitting or progressive. Lesions of MS typically develop at different times and in different CNS locations (i.e., MS is said to be disseminated in time and space). Approximately 350,000 individuals in the United States and 2.5 million individuals worldwide are affected. The clinical course can be extremely variable, ranging from a benign condition to a rapidly evolving and incapacitating disease requiring profound lifestyle adjustments.

PATHOGENESIS

Pathology New MS lesions begin with perivenular cuffing by inflammatory mononuclear cells, predominantly T cells and macrophages, which also infiltrate the surrounding white matter. At sites of inflammation, the blood-brain barrier (BBB) is disrupted, but unlike vasculitis, the vessel wall is preserved. Involvement of the humoral immune system is also evident; small numbers of B lymphocytes also infiltrate the nervous system, myelin-specific autoantibodies are present on degenerating myelin sheaths, and complement is activated. Demyelination is the hallmark of the pathology, and evidence of myelin degeneration is found at the earliest time points of tissue injury. A remarkable feature of MS plaques is that oligodendrocyte precursor cells survive—and in many lesions are present in even greater numbers than in normal tissue—but these cells fail to differentiate into mature myelin-producing cells. In some lesions, surviving oligodendrocytes or those that differentiate from precursor cells partially remyelinate the surviving naked axons, producing so-called *shadow plaques*. As lesions evolve, there is prominent astrocytic proliferation (gliosis). Over time, ectopic lymphocyte follicle-like structures, consisting of aggregates of T and B cells resembling secondary lymphoid tissue, appear in the meninges and especially overlying deep cortical sulci and also in perivascular spaces. Although relative sparing of axons is typical of MS, partial or total axonal destruction can also occur, especially within highly inflammatory lesions. Thus MS is not solely a disease of myelin, and neuronal pathology is increasingly recognized as a major contributor to irreversible neurologic disability. Inflammation, demyelination, and plaque formation are also present in the cerebral cortex, and significant axon loss indicating death of neurons is widespread, especially in advanced cases (see “Neurodegeneration,” below).

Physiology Nerve conduction in myelinated axons occurs in a saltatory manner, with the nerve impulse jumping from one node of Ranvier to the next without depolarization of the axonal membrane underlying the myelin sheath between nodes (Fig. 458-1). This produces considerably faster conduction velocities (~70 m/s) than the slow velocities (~1 m/s) produced by continuous propagation in unmyelinated nerves. Conduction block occurs when the nerve impulse is unable to traverse the demyelinated segment. This can happen when the resting axon membrane becomes hyperpolarized due to the exposure of voltage-dependent potassium channels that are normally buried underneath the myelin sheath. A temporary conduction block often follows a demyelinating event before sodium channels (originally concentrated at the nodes) redistribute along the naked

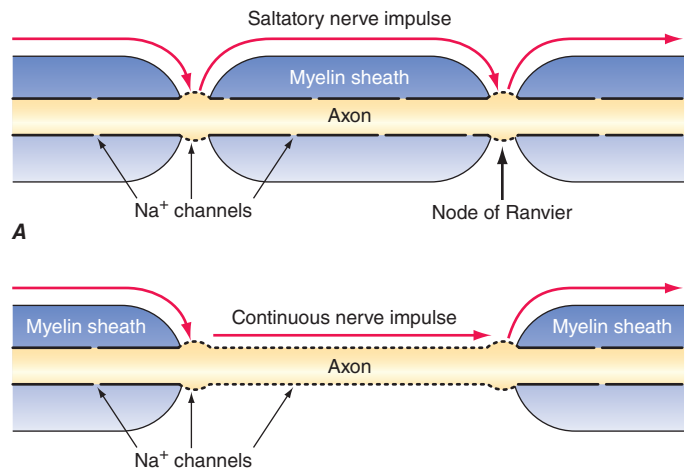


FIGURE 458-1 Nerve conduction in myelinated and demyelinated axons. **A.** Saltatory nerve conduction in myelinated axons occurs with the nerve impulse jumping from one node of Ranvier to the next. Sodium channels (shown as breaks in the solid black line) are concentrated at the nodes where axonal depolarization occurs. **B.** Following demyelination, additional sodium channels are redistributed along the axon itself, thereby allowing continuous propagation of the nerve action potential despite the absence of myelin.

axon (Fig. 458-1). This redistribution ultimately allows continuous propagation of nerve action potentials through the demyelinated segment. Conduction block may be incomplete, affecting high- but not low-frequency volleys of impulses. Variable conduction block can occur with raised body temperature or metabolic alterations and may explain clinical fluctuations that vary from hour to hour or appear with fever or exercise. Conduction slowing occurs when the demyelinated segments of the axonal membrane are reorganized to support continuous (slow) nerve impulse propagation.

Epidemiology MS is approximately threefold more common in women than men. The age of onset is typically between 20 and 40 years (slightly later in men than in women), but the disease can present across the lifespan. Approximately 10% of cases begin before age 18 years of age, and a small percentage of cases begin before the age of 10 years.

Geographical gradients have been repeatedly observed in MS, with the highest known prevalence for MS (250 per 100,000) in the Orkney Islands, located north of Scotland. In other temperate zone areas (e.g., northern North America, northern Europe, southern Australia, and southern New Zealand), the prevalence of MS is 0.1–0.2%. By contrast, in the tropics (e.g., Asia, equatorial Africa, and the Middle East), the prevalence is often 10- to 20-fold less.

The prevalence of MS has increased steadily (and dramatically) in several regions around the world over the past half-century, presumably reflecting the impact of some environmental shift. Moreover, the fact that this increase has occurred primarily (or exclusively) in women indicates that women are more responsive to this environmental change.

Well-established risk factors for MS include vitamin D deficiency, exposure to Epstein-Barr virus (EBV) after early childhood, and cigarette smoking.

Vitamin D deficiency is associated with an increase in MS risk, and data suggest that ongoing deficiency may also increase disease activity after MS begins. Immunoregulatory effects of vitamin D could explain these apparent relationships. Exposure of the skin to ultraviolet-B (UVB) radiation from the sun is essential for the biosynthesis of vitamin D, and this endogenous production is the most important source of vitamin D in most individuals; a diet rich in fatty fish represents another source of vitamin D. At high latitudes, the amount of UVB radiation reaching the earth's surface is often insufficient, particularly during winter months, and consequently, low serum levels of vitamin D are common in temperate zones. The common practice to avoid direct

sun exposure and the widespread use of sun block, which (at sun protection factor [SPF] 15) blocks 94% of the incoming UVB radiation, would be expected to exacerbate any population-wide vitamin D deficiency.

Evidence of a remote EBV infection playing some role in MS is supported by numerous epidemiologic and laboratory studies. A higher risk of infectious mononucleosis (associated with relatively late EBV infection) and higher antibody titers to latency-associated EBV nuclear antigen have been repeatedly associated with MS risk, although a causal role for EBV has not been established.

A history of cigarette smoking has also been associated with MS risk. Interestingly, in an animal model of MS, the lung was identified as a critical site for activation of pathogenic T lymphocytes responsible for autoimmune demyelination.

Recent data in MS models have also shown that high levels of dietary sodium activate pathogenic autoreactive T lymphocytes, suggesting that consumption of a high-salt diet, now widespread in the Western world, might be part of the explanation for the observed increase in the prevalence of MS in recent years.

GENETIC CONSIDERATIONS



Whites are inherently at higher risk for MS than Africans or Asians, even when residing in a similar environment. MS also aggregates within some families, and adoption, half-sibling, twin, and spousal studies indicate that familial aggregation is due to genetic, and not environmental, factors (Table 458-1).

Susceptibility to MS is polygenic, with each gene contributing a relatively small amount to the overall risk. The strongest susceptibility signal in genome-wide studies maps to the HLA-DRB1 gene in the class II region of the major histocompatibility complex (MHC), and this association accounts for approximately 10% of the disease risk. This HLA association, which was first described several decades ago, suggests that MS, at its core, is an antigen-specific autoimmune disease. Whole-genome association studies have now identified approximately 110 other MS susceptibility variants, each of which individually has only a modest effect on MS risk. Most of these MS-associated genes have known roles in the adaptive immune system, for example the genes for the interleukin (IL) 7 receptor (CD127), IL-2 receptor (CD25), and T cell costimulatory molecule LFA-3 (CD58); some variants also influence susceptibility to other autoimmune diseases in addition to MS. The variants identified so far all lack specificity and sensitivity for MS; thus, at present, they are not useful for diagnosis or to predict the future course of the disease.

IMMUNOLOGY A proinflammatory autoimmune response directed against a component of CNS myelin, and perhaps other neural elements as well, remains the cornerstone of current concepts of MS pathogenesis.

AUTOREACTIVE T LYMPHOCYTES Myelin basic protein (MBP), an intracellular protein involved in myelin compaction, is an important T cell antigen in experimental allergic encephalomyelitis (EAE), a laboratory model, and probably also in human MS. Activated MBP-reactive T cells have been identified in the blood, in cerebrospinal fluid (CSF), and within MS lesions. Moreover, *DRB1*15:01* may influence the autoimmune response because it binds with high affinity to a fragment of MBP (spanning amino acids 89–96), stimulating T cell responses to this self-protein. Two different populations of proinflammatory T cells are likely to mediate autoimmunity in MS. T-helper type 1 (T_H1)

cells producing interferon γ (IFN- γ) are one key effector population, and more recently, a role for highly proinflammatory T_H17 T cells has been established. T_H17 cells are induced by transforming growth factor β (TGF- β) and IL-6 and are amplified by IL-21 and IL-23. T_H17 cells, and levels of their corresponding cytokine IL-17, are increased in MS lesions and also in the circulation of people with active MS. High circulating levels of IL-17 may also be a marker of a more severe course of MS. T_H1 cytokines, including IL-2, tumor necrosis factor (TNF)- α , and IFN- γ , also play key roles in activating and maintaining autoimmune responses, and TNF- α and IFN- γ may directly injure oligodendrocytes or the myelin membrane.

HUMORAL AUTOIMMUNITY B cell activation and antibody responses also appear to be necessary for the full development of demyelinating lesions to occur, both in experimental models and in human MS. Clonally restricted populations of activated, antigen-experienced, memory B cells and plasma cells are present in MS lesions, in lymphoid follicle-like structures in the meninges overlying the cerebral cortex, and in the CSF. Similar or identical clonal populations are found in each compartment, indicating that a highly focused B cell response is occurring locally within the CNS in MS. Myelin-specific autoantibodies, some directed against an extracellular myelin protein, myelin oligodendrocyte glycoprotein (MOG), have been detected bound to vesiculated myelin debris in MS plaques. In the CSF, elevated levels of locally synthesized immunoglobulins and oligoclonal antibodies, derived from clonally restricted CNS B cells and plasma cells, are also characteristic of MS. The pattern of oligoclonal banding is unique to each individual, and attempts to identify the targets of these antibodies have been largely unsuccessful.

TRIGGERS Serial magnetic resonance imaging (MRI) studies in early relapsing-remitting MS reveal that bursts of focal inflammatory disease activity occur far more frequently than would have been predicted by the frequency of relapses. Thus, early in MS, most disease activity is clinically silent. Although the triggers causing these bursts are unknown, molecular mimicry between environmental agents, presumably pathogens, and myelin antigens activating pathogenic T cells may be responsible (Chap. 377e).

Neurodegeneration Axonal damage occurs in every newly formed MS lesion, and cumulative axonal loss is considered to be one important cause of irreversible neurologic disability in MS. As many as 70% of axons are lost from the lateral corticospinal (e.g., motor) tracts in patients with advanced paraparesis from MS, and longitudinal MRI studies suggest there is progressive axonal loss over time within established, inactive lesions. Demyelination can result in reduced trophic support for axons, redistribution of ion channels, and destabilization of action potential membrane potentials. Axons can adapt initially to these injuries, but over time, distal and retrograde degeneration often occurs. Therefore, promoting remyelination remains an important therapeutic goal.

In progressive MS, a key unresolved question is whether the primary neurodegenerative process occurs primarily in the cerebral cortex, the white matter, or in some combination of the two sites. As noted above, meningeal infiltrates of B and T cells are particularly prominent in progressive MS cases, and these “lymphoid follicles” are associated with underlying microglial activation, gray matter plaques, and loss of cortical neurons. White matter lesions may also contribute to late progressive MS; inactive plaques are often noninflammatory at the center, but at the edges, microglia and macrophages and evidence of ongoing axonal injury can be found. This suggests that a simmering, and possibly concentrically expanding, axonopathy may be present, even in the most chronic cases. In addition, a diffuse low-grade inflammation across large areas of white matter may be present, associated with reduced myelin staining and axonal injury (“dirty white matter”). Another characteristic of progressive MS is that inflammation is often present without a concomitant disruption of the BBB; possibly, this feature might explain the failure of immunotherapies not capable of crossing the BBB to benefit patients with progressive MS.

TABLE 458-1 RISK OF DEVELOPING MULTIPLE SCLEROSIS (MS)

1 in 3	If an identical twin has MS
1 in 15	If a fraternal twin has MS
1 in 25	If a sibling has MS
1 in 50	If a parent or half-sibling has MS
1 in 100	If a first cousin has MS
1 in 1000	If a spouse has MS
1 in 1000	If no one in the family has MS

Evidence supports a role of one, or more likely several, of the following mechanisms in progressive MS. Axonal and neuronal death may result from glutamate-mediated excitotoxicity, oxidative injury, iron accumulation, and/or mitochondrial failure either occurring as a consequence of free-radical damage or due to accumulation of deletions in mitochondrial DNA.

CLINICAL MANIFESTATIONS

The onset of MS may be abrupt or insidious. Symptoms may be severe or seem so trivial that a patient may not seek medical attention for months or years. Indeed, at autopsy, approximately 0.1% of individuals who were asymptomatic during life will be found, unexpectedly, to have pathologic evidence of MS. Similarly, in the modern era, an MRI scan obtained for an unrelated reason may show evidence of asymptomatic MS. Symptoms of MS are extremely varied and depend on the location and severity of lesions within the CNS (Table 458-2). Examination often reveals evidence of neurologic dysfunction, often in asymptomatic locations. For example, a patient may present with symptoms in one leg but signs in both.

Weakness of the limbs may manifest as loss of strength, speed, or dexterity, as fatigue, or as a disturbance of gait. Exercise-induced weakness is a characteristic symptom of MS. The weakness is of the upper motor neuron type (Chap. 30) and is usually accompanied by other pyramidal signs such as spasticity, hyperreflexia, and Babinski signs. Occasionally a tendon reflex may be lost (simulating a lower motor neuron lesion) if an MS lesion disrupts the afferent reflex fibers in the spinal cord (see Fig. 30-2).

Spasticity (Chap. 30) is commonly associated with spontaneous and movement-induced muscle spasms. More than 30% of MS patients have moderate to severe spasticity, especially in the legs. This is often accompanied by painful spasms interfering with ambulation, work, or self-care. Occasionally spasticity provides support for the body weight during ambulation, and in these cases, treatment of spasticity may actually do more harm than good.

Optic neuritis (ON) presents as diminished visual acuity, dimness, or decreased color perception (desaturation) in the central field of vision. These symptoms can be mild or may progress to severe visual loss. Rarely, there is complete loss of light perception. Visual symptoms are generally monocular but may be bilateral. Periorbital pain (aggravated by eye movement) often precedes or accompanies the visual loss. An afferent pupillary defect (Chap. 39) is usually present. Funduscopic examination may be normal or reveal optic disc swelling (papillitis). Pallor of the optic disc (optic atrophy) commonly follows ON. Uveitis is uncommon and should raise the possibility of alternative diagnoses such as sarcoid or lymphoma.

Visual blurring in MS may result from ON or diplopia (double vision); if the symptom resolves when either eye is covered, the cause is diplopia.

Diplopia may result from internuclear ophthalmoplegia (INO) or from palsy of the sixth cranial nerve (rarely the third or fourth). An INO consists of impaired adduction of one eye due to a lesion in the ipsilateral medial longitudinal fasciculus (Chaps. 41e and 42).

Prominent nystagmus is often observed in the abducting eye, along with a small skew deviation. A bilateral INO is particularly suggestive of MS. Other common gaze disturbances in MS include (1) a horizontal gaze palsy, (2) a “one and a half” syndrome (horizontal gaze palsy plus an INO), and (3) acquired pendular nystagmus.

Sensory symptoms are varied and include both paresthesias (e.g., tingling, prickling sensations, formications, “pins and needles,” or painful burning) and hypesthesia (e.g., reduced sensation, numbness, or a “dead” feeling). Unpleasant sensations (e.g., feelings that body parts are swollen, wet, raw, or tightly wrapped) are also common. Sensory impairment of the trunk and legs below a horizontal line on the torso (a sensory level) indicates that the spinal cord is the origin of the sensory disturbance. It is often accompanied by a bandlike sensation of tightness around the torso. Pain is a common symptom of MS, experienced by >50% of patients. Pain can occur anywhere on the body and can change locations over time.

Ataxia usually manifests as cerebellar tremors (Chap. 450). Ataxia may also involve the head and trunk or the voice, producing a characteristic cerebellar dysarthria (scanning speech).

Bladder dysfunction is present in >90% of MS patients, and in a third of patients, dysfunction results in weekly or more frequent episodes of incontinence. During normal reflex voiding, relaxation of the bladder sphincter (α -adrenergic innervation) is coordinated with contraction of the detrusor muscle in the bladder wall (muscarinic cholinergic innervation). *Detrusor hyperreflexia*, due to impairment of suprasegmental inhibition, causes urinary frequency, urgency, nocturia, and uncontrolled bladder emptying. *Detrusor sphincter dyssynergia*, due to loss of synchronization between detrusor and sphincter muscles, causes difficulty in initiating and/or stopping the urinary stream, producing hesitancy, urinary retention, overflow incontinence, and recurrent infection.

Constipation occurs in >30% of patients. Fecal urgency or *bowel incontinence* is less common (<15%) but can be socially debilitating.

Cognitive dysfunction can include memory loss; impaired attention; difficulties in executive functioning, memory, and problem solving; slowed information processing; and problems shifting between cognitive tasks. Euphoria (elevated mood) was once thought to be characteristic of MS but is actually uncommon, occurring in <20% of patients. Cognitive dysfunction sufficient to impair activities of daily living is rare.

Depression, experienced by approximately half of patients, can be reactive, endogenous, or part of the illness itself and can contribute to fatigue.

Fatigue (Chap. 29) is experienced by 90% of patients; this symptom is the most common reason for work-related disability in MS. Fatigue can be exacerbated by elevated temperatures, depression, expending exceptional effort to accomplish basic activities of daily living, or sleep disturbances (e.g., from frequent nocturnal awakenings to urinate).

Sexual dysfunction may manifest as decreased libido, impaired genital sensation, impotence in men, and diminished vaginal lubrication or adductor spasms in women.

Facial weakness due to a lesion in the pons may resemble idiopathic Bell's palsy (Chap. 455). Unlike Bell's palsy, facial weakness in MS is usually not associated with ipsilateral loss of taste sensation or retroauricular pain.

Vertigo may appear suddenly from a brainstem lesion, superficially resembling acute labyrinthitis (Chap. 28). *Hearing loss* (Chap. 43) may also occur in MS but is uncommon.

Ancillary Symptoms *Heat sensitivity* refers to neurologic symptoms produced by an elevation of the body's core temperature. For example, unilateral visual blurring may occur during a hot shower or with physical exercise (*Uhthoff's symptom*). It is also common for MS symptoms to worsen transiently, sometimes dramatically, during febrile illnesses (see “Acute Attacks or Initial Demyelinating Episodes,” below). Such heat-related symptoms probably result from transient conduction block (see above).

Lhermitte's symptom is an electric shock-like sensation (typically induced by flexion or other movements of the neck) that radiates down

TABLE 458-2 INITIAL SYMPTOMS OF MS

Symptom	Percentage of Cases	Symptom	Percentage of Cases
Sensory loss	37	Lhermitte	3
Optic neuritis	36	Pain	3
Weakness	35	Dementia	2
Paresthesias	24	Visual loss	2
Diplopia	15	Facial palsy	1
Ataxia	11	Impotence	1
Vertigo	6	Myokymia	1
Paroxysmal attacks	4	Epilepsy	1
Bladder	4	Falling	1

Source: After WB Matthews et al: *McAlpine's Multiple Sclerosis*. New York, Churchill Livingstone, 1991.

the back into the legs. Rarely, it radiates into the arms. It is generally self-limited but may persist for years. Lhermitte's symptom can also occur with other disorders of the cervical spinal cord (e.g., cervical spondylosis).

Paroxysmal symptoms are distinguished by their brief duration (10 s to 2 min), high frequency (5–40 episodes per day), lack of any alteration of consciousness or change in background electroencephalogram during episodes, and a self-limited course (generally lasting weeks to months). They may be precipitated by hyperventilation or movement. These syndromes may include Lhermitte's symptom; tonic contractions of a limb, face, or trunk (tonic seizures); paroxysmal dysarthria and ataxia; paroxysmal sensory disturbances; and several other less well-characterized syndromes. Paroxysmal symptoms probably result from spontaneous discharges, arising at the edges of demyelinated plaques and spreading to adjacent white matter tracts.

Trigeminal neuralgia, hemifacial spasm, and glossopharyngeal neuralgia (Chap. 455) can occur when the demyelinating lesion involves the root entry (or exit) zone of the fifth, seventh, and ninth cranial nerve, respectively. Trigeminal neuralgia (tic douloureux) is a very brief lancinating facial pain often triggered by an afferent input from the face or teeth. Most cases of trigeminal neuralgia are not MS related; however, atypical features such as onset before age 50 years, bilateral symptoms, objective sensory loss, or nonparoxysmal pain should raise the possibility that MS could be responsible.

Facial myokymia consists of either persistent rapid flickering contractions of the facial musculature (especially the lower portion of the orbicularis oculus) or a contraction that slowly spreads across the face. It results from lesions of the corticobulbar tracts or brainstem course of the facial nerve.

DISEASE COURSE

Four clinical types of MS exist (Fig. 458-2):

1. *Relapsing/remitting MS (RRMS)* accounts for 85% of MS cases at onset and is characterized by discrete attacks that generally evolve over days to weeks (rarely over hours). With initial attacks, there is often substantial or complete recovery over the ensuing weeks to months, but as attacks continue over time recovery may be less evident (Fig. 458-2A). Between attacks, patients are neurologically stable.
2. *Secondary progressive MS (SPMS)* always begins as RRMS (Fig. 458-2B). At some point, however, the clinical course changes so that the patient experiences a steady deterioration in function

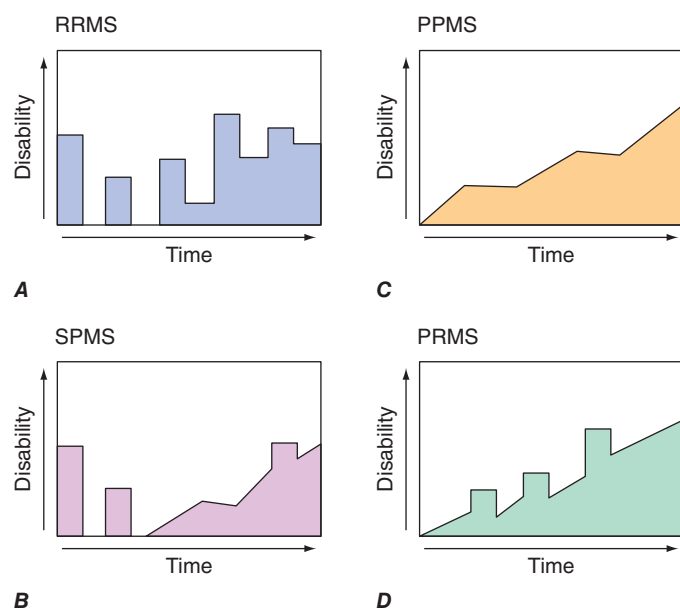


FIGURE 458-2 Clinical course of multiple sclerosis (MS).

A. Relapsing/remitting MS (RRMS). B. Secondary progressive MS (SPMS). C. Primary progressive MS (PPMS). D. Progressive/relapsing MS (PRMS).

unassociated with acute attacks (which may continue or cease during the progressive phase). SPMS produces a greater amount of fixed neurologic disability than RRMS. For a patient with RRMS, the risk of developing SPMS is ~2% each year, meaning that the great majority of RRMS ultimately evolves into SPMS. SPMS appears to represent a late stage of the same underlying illness as RRMS.

3. *Primary progressive MS (PPMS)* accounts for ~15% of cases. These patients do not experience attacks but only a steady functional decline from disease onset (Fig. 458-2C). Compared to RRMS, the sex distribution is more even, the disease begins later in life (mean age ~40 years), and disability develops faster (at least relative to the onset of the first clinical symptom). Despite these differences, PPMS appears to represent the same underlying illness as RRMS.
4. *Progressive/relapsing MS (PRMS)* overlaps PPMS and SPMS and accounts for ~5% of MS patients. Like patients with PPMS, these patients experience a steady deterioration in their condition from disease onset. However, like SPMS patients, they experience occasional attacks superimposed upon their progressive course (Fig. 458-2D).

DIAGNOSIS

There is no definitive diagnostic test for MS. Diagnostic criteria for clinically definite MS require documentation of two or more episodes of symptoms and two or more signs that reflect pathology in anatomically noncontiguous white matter tracts of the CNS (Table 458-3). Symptoms must last for >24 h and occur as distinct episodes that are separated by a month or more. In patients who have only one of the two required signs on neurologic examination, the second may be documented by abnormal tests such as MRI or evoked potentials (EPs). Similarly, in the most recent diagnostic scheme, the second clinical event (in time) may be supported solely by MRI findings, consisting of either the development of new focal white matter lesions on MRI or the simultaneous presence of both an enhancing lesion and a nonenhancing lesion in an asymptomatic location. In patients whose course is progressive from onset for ≥ 6 months without superimposed relapses, documentation of intrathecal IgG synthesis may be used to support a diagnosis of PPMS.

DIAGNOSTIC TESTS

Magnetic Resonance Imaging MRI has revolutionized the diagnosis and management of MS (Fig. 458-3); characteristic abnormalities are found in >95% of patients, although more than 90% of the lesions visualized by MRI are asymptomatic. An increase in vascular permeability from a breakdown of the BBB is detected by leakage of intravenous gadolinium (Gd) into the parenchyma. Such leakage occurs early in the development of an MS lesion and serves as a useful marker of inflammation. Gd enhancement typically persists for approximately 1 month, and the residual MS plaque remains visible indefinitely as a focal area of hyperintensity (a lesion) on spin-echo (T2-weighted) and proton-density images. Lesions are frequently oriented perpendicular to the ventricular surface, corresponding to the pathologic pattern of perivenous demyelination (Dawson's fingers). Lesions are multifocal within the brain, brainstem, and spinal cord. Lesions larger than 6 mm located in the corpus callosum, periventricular white matter, brainstem, cerebellum, or spinal cord are particularly helpful diagnostically. Current criteria for the use of MRI in the diagnosis of MS are shown in Table 458-3.

The total volume of T2-weighted signal abnormality (the "burden of disease") shows a significant (albeit weak) correlation with clinical disability, as do measures of brain atrophy. Approximately one-third of T2-weighted lesions appear as hypointense lesions (black holes) on T1-weighted imaging. Black holes may be a marker of irreversibly demyelination and axonal loss, although even this measure depends on the timing of the image acquisition (e.g., most acute Gd-enhancing T2 lesions are T1 dark).

Newer MRI methods such as magnetization transfer ratio (MTR) imaging and proton magnetic resonance spectroscopic imaging (MRSI) may ultimately serve as surrogate markers of clinical disability. MRSI can quantitate molecules such as *N*-acetyl aspartate, which

TABLE 458-3 DIAGNOSTIC CRITERIA FOR MULTIPLE SCLEROSIS (MS)

Clinical Presentation	Additional Data Needed for MS Diagnosis
2 or more attacks; objective clinical evidence of 2 or more lesions or objective clinical evidence of 1 lesion with reasonable historical evidence of a prior attack	None
2 or more attacks; objective clinical evidence of 1 lesion	Dissemination in space, demonstrated by <ul style="list-style-type: none"> • ≥ 1 T2 lesion on MRI in at least 2 out of 4 MS-typical regions of the CNS (periventricular, juxtacortical, infratentorial, or spinal cord) OR <ul style="list-style-type: none"> • Await a further clinical attack implicating a different CNS site
1 attack; objective clinical evidence of 2 or more lesions	Dissemination in time, demonstrated by <ul style="list-style-type: none"> • Simultaneous presence of asymptomatic gadolinium-enhancing and nonenhancing lesions at any time OR <ul style="list-style-type: none"> • A new T2 and/or gadolinium-enhancing lesion(s) on follow-up MRI, irrespective of its timing with reference to a baseline scan OR <ul style="list-style-type: none"> • Await a second clinical attack
1 attack; objective clinical evidence of 1 lesion (clinically isolated syndrome)	Dissemination in space and time, demonstrated by: For dissemination in space <ul style="list-style-type: none"> • ≥ 1 T2 lesion in at least 2 out of 4 MS-typical regions of the CNS (periventricular, juxtacortical, infratentorial, or spinal cord) OR <ul style="list-style-type: none"> • Await a second clinical attack implicating a different CNS site AND For dissemination in time <ul style="list-style-type: none"> • Simultaneous presence of asymptomatic gadolinium-enhancing and nonenhancing lesions at any time OR <ul style="list-style-type: none"> • A new T2 and/or gadolinium-enhancing lesion(s) on follow-up MRI, irrespective of its timing with reference to a baseline scan OR <ul style="list-style-type: none"> • Await a second clinical attack
Insidious neurologic progression suggestive of MS (PPMS)	1 year of disease progression (retrospectively or prospectively determined) PLUS 2 out of the 3 following criteria Evidence for dissemination in space in the brain based on ≥ 1 T2+ lesions in the MS-characteristic periventricular, juxtacortical, or infratentorial regions Evidence for dissemination in space in the spinal cord based on ≥ 2 T2+ lesions in the cord Positive CSF (isoelectric focusing evidence of oligoclonal bands and/or elevated IgG index)

Abbreviations: CNS, central nervous system; CSF, cerebrospinal fluid; MRI, magnetic resonance imaging; PPMS, primary progressive multiple sclerosis.

Source: From CH Polman et al: Diagnostic Criteria for Multiple Sclerosis: 2010 Revisions to the "McDonald Criteria." *Ann Neurol* 69:292, 2011.

is a marker of axonal integrity, and MTR may be able to distinguish demyelination from edema.

Evoked Potentials EP testing assesses function in afferent (visual, auditory, and somatosensory) or efferent (motor) CNS pathways. EPs use computer averaging to measure CNS electric potentials evoked

by repetitive stimulation of selected peripheral nerves or of the brain. These tests provide the most information when the pathways studied are clinically uninvolved. For example, in a patient with a remitting and relapsing spinal cord syndrome with sensory deficits in the legs, an abnormal somatosensory EP following posterior tibial nerve stimulation provides little new information. By contrast, an abnormal visual EP in this circumstance would permit a diagnosis of clinically definite MS (Table 458-3). Abnormalities on one or more EP modalities occur in 80–90% of MS patients. EP abnormalities are not specific to MS, although a marked delay in the latency of a specific EP component (as opposed to a reduced amplitude or distorted wave-shape) is suggestive of demyelination.

Cerebrospinal Fluid CSF abnormalities found in MS include a mononuclear cell pleocytosis and an increased level of intrathecally synthesized IgG. The total CSF protein is usually normal. Various formulas distinguish intrathecally synthesized IgG from IgG that may have entered the CNS passively from the serum. One formula, the CSF IgG index, expresses the ratio of IgG to albumin in the CSF divided by the same ratio in the serum. The IgG synthesis rate uses serum and CSF IgG and albumin measurements to calculate the rate of CNS IgG synthesis. The measurement of oligoclonal bands (OCBs) by agarose gel electrophoresis in the CSF also assesses intrathecal production of IgG. Two or more discrete OCBs, not present in a paired serum sample, are found in >75% of patients with MS. OCBs may be absent at the onset of MS, and in individual patients, the number of bands may increase with time.

A mild CSF pleocytosis (>5 cells/ μ L) is present in ~25% of cases, usually in young patients with RRMS. A pleocytosis of >75 cells/ μ L, the presence of polymorphonuclear leukocytes, or a protein concentration >1 g/L (>100 mg/dL) in CSF should raise concern that the patient may not have MS.

DIFFERENTIAL DIAGNOSIS

No single clinical sign or test is diagnostic of MS. The diagnosis is readily made in a young adult with relapsing and remitting symptoms involving different areas of CNS white matter. The possibility of an alternative diagnosis should always be considered (Table 458-4), particularly when (1) symptoms are localized exclusively to the posterior fossa, craniocervical junction, or spinal cord; (2) the patient is <15 or >60 years of age; (3) the clinical course is progressive from onset; (4) the patient has never experienced visual, sensory, or bladder symptoms; or (5) laboratory findings (e.g., MRI, CSF, or EPs) are atypical. Similarly, uncommon or rare symptoms in MS (e.g., aphasia, parkinsonism, chorea, isolated dementia, severe muscular atrophy, peripheral neuropathy, episodic loss of consciousness, fever, headache, seizures, or coma) should increase concern about an alternative diagnosis. Diagnosis is also difficult in patients with a rapid or explosive (stroke-like) onset or with mild symptoms and a normal neurologic examination. Rarely, intense inflammation and swelling may produce a mass lesion that mimics a primary or metastatic tumor. In the current era, the disorders most likely to be mistaken for MS are neuromyelitis optica (see below), sarcoid, vascular disorders (including antiphospholipid syndrome and vasculitis), and rarely CNS lymphoma. The specific tests required to exclude alternative diagnoses will vary with each clinical situation; however, an erythrocyte sedimentation rate, serum B_{12} level, ANA, and treponemal antibody should probably be obtained in all patients with suspected MS.

PROGNOSIS

Most patients with clinically evident MS ultimately experience progressive neurologic disability. In older studies conducted mostly before disease-modifying therapies for MS were widely available, 15 years after onset, only 20% of patients had no functional limitation, and between one-third and one-half progressed to SPMS and required assistance with ambulation; furthermore, 25 years after onset, ~80% of MS patients reached this level of disability. The long-term prognosis for untreated MS appears to have improved in recent years, at least in part because of the development of therapies for the early relapsing form of the disease. Although the prognosis in an individual is

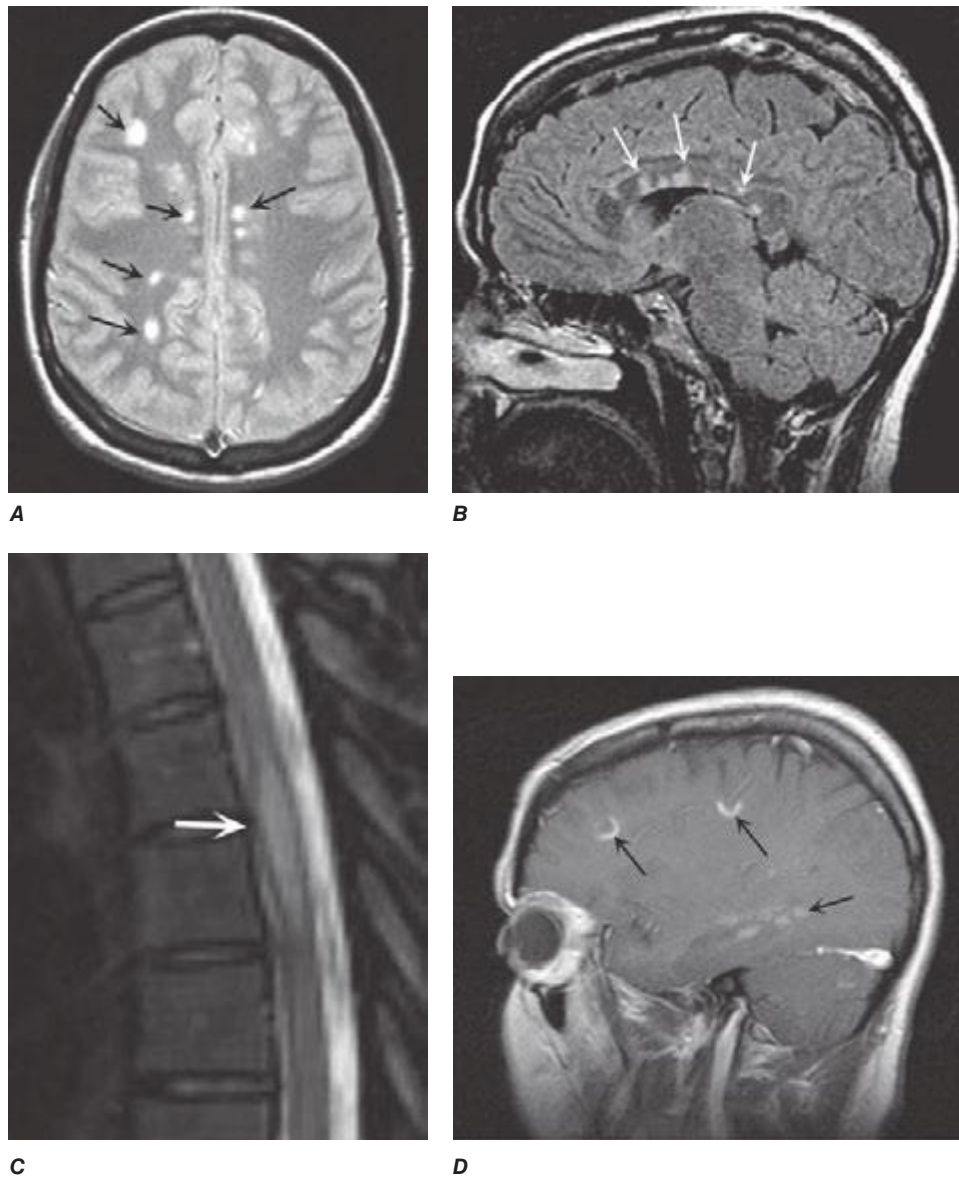


FIGURE 458-3 Magnetic resonance imaging findings in multiple sclerosis (MS). **A.** Axial first-echo image from T2-weighted sequence demonstrates multiple bright signal abnormalities in white matter, typical for MS. **B.** Sagittal T2-weighted fluid-attenuated inversion recovery (FLAIR) image in which the high signal of cerebrospinal fluid (CSF) has been suppressed. CSF appears dark, whereas areas of brain edema or demyelination appear high in signal as shown here in the corpus callosum (*arrows*). Lesions in the anterior corpus callosum are frequent in MS and rare in vascular disease. **C.** Sagittal T2-weighted fast spin echo image of the thoracic spine demonstrates a fusiform high-signal-intensity lesion in the midthoracic spinal cord. **D.** Sagittal T1-weighted image obtained after the intravenous administration of gadolinium DTPA reveals focal areas of blood-brain barrier disruption, identified as high-signal-intensity regions (*arrows*).

difficult to establish, certain clinical features suggest a more favorable prognosis. These include ON or sensory symptoms at onset; fewer than two relapses in the first year of illness; and minimal impairment after 5 years. By contrast, patients with truncal ataxia, action tremor, pyramidal symptoms, or a progressive disease course are more likely to become disabled. Patients with a long-term favorable course are likely to have developed fewer MRI lesions during the early years of disease, and vice versa. Importantly, some MS patients have a benign variant of MS and never develop neurologic disability. The likelihood of having benign MS is thought to be <20%. Patients with benign MS 15 years after onset who have entirely normal neurologic examinations are likely to maintain their benign course

In patients with their first demyelinating event (i.e., a clinically isolated syndrome), the brain MRI provides prognostic information. With three or more typical T2-weighted lesions, the risk of developing MS after 20 years is ~80%. Conversely, with a normal brain MRI, the likelihood of developing MS is <20%. Similarly, the presence of two or more Gd-enhancing lesions at baseline is highly predictive of future

MS, as is the appearance of either new T2-weighted lesions or new Gd enhancement ≥ 3 months after the initial episode.

Mortality as a direct consequence of MS is uncommon, although it has been estimated that the 25-year survival is only 85% of expected. Death can occur during an acute MS attack, although this is distinctly rare. More commonly, death occurs as a complication of MS (e.g., pneumonia in a debilitated individual). Death can also result from suicide. Early disease-modifying therapy seems to reduce this excess mortality.

Effect of Pregnancy Pregnant MS patients experience fewer attacks than expected during gestation (especially in the last trimester), but more attacks than expected in the first 3 months postpartum. When considering the pregnancy year as a whole (i.e., 9 months of pregnancy plus 3 months postpartum), the overall disease course is unaffected. Decisions about childbearing should thus be made based on (1) the mother's physical state, (2) her ability to care for the child, and (3) the availability of social support. Disease-modifying therapy is

TABLE 458-4 DISORDERS THAT CAN MIMIC MULTIPLE SCLEROSIS (MS)

Acute disseminated encephalomyelitis (ADEM)
Antiphospholipid antibody syndrome
Behçet's disease
Cerebral autosomal dominant arteriopathy, subcortical infarcts, and leukoencephalopathy (CADASIL)
Congenital leukodystrophies (e.g., adrenoleukodystrophy, metachromatic leukodystrophy)
Human immunodeficiency virus (HIV) infection
Ischemic optic neuropathy (arteritic and nonarteritic)
Lyme disease
Mitochondrial encephalopathy with lactic acidosis and stroke (MELAS)
Neoplasms (e.g., lymphoma, glioma, meningioma)
Sarcoid
Sjögren's syndrome
Stroke and ischemic cerebrovascular disease
Syphilis
Systemic lupus erythematosus and related collagen vascular disorders
Tropical spastic paraparesis (HTLV-1/2 infection)
Vascular malformations (especially spinal dural AV fistulas)
Vasculitis (primary CNS or other)
Vitamin B ₁₂ deficiency

Abbreviations: AV, arteriovenous; CNS, central nervous system; HTLV, human T cell lymphotropic virus.

generally discontinued during pregnancy, although the actual risk from the interferons and glatiramer acetate (see below) appears to be low.

TREATMENT MULTIPLE SCLEROSIS

Therapy for MS can be divided into several categories: (1) treatment of acute attacks, (2) treatment with disease-modifying agents that reduce the biologic activity of MS, and (3) symptomatic therapy. Treatments that promote remyelination or neural repair do not currently exist, but several promising approaches are being actively investigated.

The Expanded Disability Status Score (EDSS) is a widely used measure of neurologic impairment in MS (Table 458-5). Most patients with EDSS scores <3.5 have RRMS, walk normally, and are generally not disabled; by contrast, patients with EDSS scores >5.5 have progressive MS (SPMS or PPMS), are gait-impaired, and, typically, are occupationally disabled.

ACUTE ATTACKS OR INITIAL DEMYELINATING EPISODES

When patients experience acute deterioration, it is important to consider whether this change reflects new disease activity or a "pseudoexacerbation" resulting from an increase in ambient temperature, fever, or an infection. When the clinical change is thought to reflect a pseudoexacerbation, glucocorticoid treatment is inappropriate. Glucocorticoids are used to manage either first attacks or acute exacerbations. They provide short-term clinical benefit by reducing the severity and shortening the duration of attacks. Whether treatment provides any long-term benefit on the course of the illness is less clear. Therefore, mild attacks are often not treated. Physical and occupational therapy can help with mobility and manual dexterity.

Glucocorticoid treatment is usually administered as intravenous methylprednisolone, 500–1000 mg/d for 3–5 days, either without a taper or followed by a course of oral prednisone beginning at a dose of 60–80 mg/d and gradually tapered over 2 weeks. Orally administered methylprednisolone or dexamethasone (in equivalent dosages) can be substituted for the intravenous portion of the therapy, although gastrointestinal complications are more common by this route. Outpatient treatment is almost always possible.

Side effects of short-term glucocorticoid therapy include fluid retention, potassium loss, weight gain, gastric disturbances, acne,

and emotional lability. Concurrent use of a low-salt, potassium-rich diet and avoidance of potassium-wasting diuretics are advisable. Lithium carbonate (300 mg orally bid) may help to manage emotional lability and insomnia associated with glucocorticoid therapy. Patients with a history of peptic ulcer disease may require cimetidine (400 mg bid) or ranitidine (150 mg bid). Proton pump inhibitors such as pantoprazole (40 mg orally bid) may reduce the likelihood of gastritis, especially when large doses are administered orally. Plasma exchange (five to seven exchanges: 40–60 mL/kg per exchange, every other day for 14 days) may benefit patients with fulminant attacks of demyelination that are unresponsive to glucocorticoids. However, the cost is high, and conclusive evidence of efficacy is lacking.

DISEASE-MODIFYING THERAPIES FOR RELAPSING FORMS OF MS (RRMS, SPMS WITH EXACERBATIONS)

Ten such agents are approved by the U.S. Food and Drug Administration (FDA): (1) IFN- β -1a (Avonex), (2) IFN- β -1a (Rebif), (3) IFN- β -1b (Betaseron or Extavia), (4) glatiramer acetate (Copaxone), (5) natalizumab (Tysabri), (6) fingolimod (Gilenya), (7) dimethyl fumarate (Tecfidera), (8) teriflunomide (Aubagio), (9) mitoxantrone (Novantrone), and (10) alemtuzumab (Lemtrada). Several other promising agents are in varying stages of product development. Each of these treatments can also be used in SPMS patients who continue to experience attacks, both because SPMS can be difficult to distinguish from RRMS and because the available clinical trials, although not definitive, suggest that such patients may sometimes derive therapeutic benefit. Thus, in several phase 3 clinical trials, recipients of each of these agents experienced fewer clinical exacerbations and fewer new MRI lesions compared to placebo recipients (Table 458-6). Because of its potential toxicity as an immunosuppressant, mitoxantrone is generally reserved for patients with progressive disability who have failed other treatments. When considering the data in Table 458-6, however, it is important to note that the relative efficacy of the different agents cannot be determined by cross-trial comparisons. Relative efficacy can only be assessed from nonbiased head-to-head clinical trials.

Interferon- β IFN- β is a class I interferon originally identified by its antiviral properties. Efficacy in MS probably results from immunomodulatory properties including (1) downregulating expression of MHC molecules on antigen-presenting cells, (2) reducing proinflammatory and increasing regulatory cytokine levels, (3) inhibiting T cell proliferation, and (4) limiting the trafficking of inflammatory cells in the CNS. IFN- β reduces the attack rate and improves disease severity measures such as EDSS progression and MRI-documented disease burden. IFN- β should be considered in patients with either RRMS or SPMS with superimposed relapses. In patients with SPMS but without relapses, efficacy has not been established. Head-to-head trials suggest that dosing IFN- β more frequently and at higher doses has better efficacy but is also more likely to induce neutralizing antibodies (see below). IFN- β -1a (Avonex), 30 μ g, is administered by intramuscular injection once every week. IFN- β -1a (Rebif), 44 μ g, is administered by subcutaneous injection three times per week. IFN- β -1b (Betaseron or Extavia), 250 μ g, is administered by subcutaneous injection every other day.

Common side effects of IFN- β therapy include flulike symptoms (e.g., fevers, chills, and myalgias) and mild abnormalities on routine laboratory evaluation (e.g., elevated liver function tests or lymphopenia). Rarely, more severe hepatotoxicity may occur. Subcutaneous IFN- β also causes reactions at the injection site (e.g., pain, redness, induration, or, rarely, skin necrosis). Side effects can usually be managed with concomitant nonsteroidal anti-inflammatory medications. Depression, increased spasticity, and cognitive changes have been reported, although these symptoms can also be due to the underlying disease. In any event, side effects due to IFN- β therapy usually subside over time.

Kurtzke Expanded Disability Status Score (EDSS)	
0.0 = Normal neurologic exam (all grade 0 in functional status [FS])	6.0 = Unilateral assistance required to walk about 100 m with or without resting
1.0 = No disability, minimal signs in one FS (i.e., grade 1)	6.5 = Constant bilateral assistance required to walk about 20 m without resting
1.5 = No disability, minimal signs in more than one FS (more than one grade 1)	7.0 = Unable to walk beyond about 5 m even with aid; essentially restricted to wheelchair; wheels self and transfers alone
2.0 = Minimal disability in one FS (one FS grade 2, others 0 or 1)	7.5 = Unable to take more than a few steps; restricted to wheelchair; may need aid to transfer
2.5 = Minimal disability in two FS (two FS grade 2, others 0 or 1)	8.0 = Essentially restricted to bed or chair or perambulated in wheelchair, but out of bed most of day; retains many self-care functions; generally has effective use of arms
3.0 = Moderate disability in one FS (one FS grade 3, others 0 or 1) or mild disability in three or four FS (three/four FS grade 2, others 0 or 1) although fully ambulatory	8.5 = Essentially restricted to bed much of the day; has some effective use of arm(s); retains some self-care functions
3.5 = Fully ambulatory but with moderate disability in one FS (one grade 3) and one or two FS grade 2; or two FS grade 3; or five FS grade 2 (others 0 or 1)	9.0 = Helpless bed patient; can communicate and eat
4.0 = Ambulatory without aid or rest for ~500 m	9.5 = Totally helpless bed patient; unable to communicate or eat
4.5 = Ambulatory without aid or rest for ~300 m	10.0 = Death due to MS
5.0 = Ambulatory without aid or rest for ~200 m	
5.5 = Ambulatory without aid or rest for ~100 m	
Functional Status (FS) Score	
A. Pyramidal functions	4 = Marked decrease in touch or pain or loss of proprioception, alone or combined, in 1 or 2 limbs or moderate decrease in touch or pain and/or severe proprioceptive decrease in more than 2 limbs
0 = Normal	5 = Loss (essentially) of sensation in 1 or 2 limbs or moderate decrease in touch or pain and/or loss of proprioception for most of the body below the head
1 = Abnormal signs without disability	6 = Sensation essentially lost below the head
2 = Minimal disability	E. Bowel and bladder functions
3 = Mild or moderate paraparesis or hemiparesis, or severe monoparesis	0 = Normal
4 = Marked paraparesis or hemiparesis, moderate quadriparesis, or monoplegia	1 = Mild urinary hesitancy, urgency, or retention
5 = Paraplegia, hemiplegia, or marked quadriparesis	2 = Moderate hesitancy, urgency, retention of bowel or bladder, or rare urinary incontinence
6 = Quadriplegia	3 = Frequent urinary incontinence
B. Cerebellar functions	4 = In need of almost constant catheterization
0 = Normal	5 = Loss of bladder function
1 = Abnormal signs without disability	6 = Loss of bowel and bladder function
2 = Mild ataxia	F. Visual (or optic) functions
3 = Moderate truncal or limb ataxia	0 = Normal
4 = Severe ataxia all limbs	1 = Scotoma with visual acuity (corrected) better than 20/30
5 = Unable to perform coordinated movements due to ataxia	2 = Worse eye with scotoma with maximal visual acuity (corrected) of 20/30 to 20/59
C. Brainstem functions	3 = Worse eye with large scotoma, or moderate decrease in fields, but with maximal visual acuity (corrected) of 20/60 to 20/99
0 = Normal	4 = Worse eye with marked decrease of fields and maximal acuity (corrected) of 20/100 to 20/200; grade 3 plus maximal acuity of better eye of 20/60 or less
1 = Signs only	5 = Worse eye with maximal visual acuity (corrected) less than 20/200; grade 4 plus maximal acuity of better eye of 20/60 or less
2 = Moderate nystagmus or other mild disability	6 = Grade 5 plus maximal visual acuity of better eye of 20/60 or less
3 = Severe nystagmus, marked extraocular weakness, or moderate disability of other cranial nerves	G. Cerebral (or mental) functions
4 = Marked dysarthria or other marked disability	0 = Normal
5 = Inability to swallow or speak	1 = Mood alteration only (does not affect EDSS score)
D. Sensory functions	2 = Mild decrease in mentation
0 = Normal	3 = Moderate decrease in mentation
1 = Vibration or figure-writing decrease only, in 1 or 2 limbs	4 = Marked decrease in mentation
2 = Mild decrease in touch or pain or position sense, and/or moderate decrease in vibration in 1 or 2 limbs, or vibratory decrease alone in 3 or 4 limbs	5 = Chronic brain syndrome—severe or incompetent
3 = Moderate decrease in touch or pain or position sense, and/or essentially lost vibration in 1 or 2 limbs, or mild decrease in touch or pain, and/or moderate decrease in all proprioceptive tests in 3 or 4 limbs	

Source: Adapted from JF Kurtzke: Rating neurologic impairment in multiple sclerosis: An expanded disability status scale (EDSS). *Neurology* 33:1444, 1983.

Approximately 2–10% of IFN- β -1a (Avonex) recipients, 15–25% of IFN- β -1a (Rebif) recipients, and 30–40% of IFN- β -1b (Betaseron/Extavia) recipients develop neutralizing antibodies to IFN- β , which may disappear over time. Two very large randomized trials (one with >2000 patients) provide unequivocal evidence that neutralizing antibodies reduce efficacy as determined by several MRI outcomes. Paradoxically, however, these same trials, despite abundant statistical power, failed to demonstrate any concomitant impact on the clinical outcomes of disability and relapse rate. The reason for this clinical-radiologic dissociation is unresolved. For a patient doing

well on therapy, the presence of antibodies should not affect treatment. Conversely, for a patient doing poorly on therapy, alternative treatment should be considered, even if there are no detectable antibodies.

Glatiramer Acetate Glatiramer acetate is a synthetic, random polypeptide composed of four amino acids (L-glutamic acid, L-lysine, L-alanine, and L-tyrosine). Its mechanism of action may include (1) induction of antigen-specific suppressor T cells; (2) binding to MHC molecules, thereby displacing bound MBP; or (3) altering

TABLE 458-6 TWO-YEAR OUTCOMES FOR FDA-APPROVED THERAPIES FOR MULTIPLE SCLEROSIS^a

Dose, Route, and Schedule	Clinical Outcomes ^b		MRI Outcomes ^c	
	Attack Rate, Mean	Change in Disease Severity	New T2 Lesions ^d	Total Burden of Disease
IFN-β-1b, 250 μg SC qod	-34% ^e	-29% (NS)	-83% ^f	-17% ^e
IFN-β-1a, 30 μg IM qw	-18% ^g	-37% ^g	-36% ^f	-4% (NS)
IFN-β-1a, 44 μg SC tiw	-32% ^e	-30% ^g	-78% ^e	-15% ^e
GA, 20 mg SC qd	-29% ^f	-12% (NS)	-38% ^f	-8% ^f
MTX, 12 mg/m ² IV q3mo	-66% ^e	-75% ^g	-79% ^g	NR
NTZ, 300 mg IV qmo	-68% ^e	-42% ^e	-83% ^e	-18% ^e
FGM, 0.5 mg PO qd	-55% ^e	-34% ^f	-74% ^e	-23% ^e
DMF, 240 mg PO bid	-52% ^e	-40% ^f	-71% ^e	NR
TF, 14 mg PO qd	-31% ^e	-26% ^g	-70% ^e	-20% ^g

^aPercentage reductions (or increases) have been calculated by dividing the reported rates in the treated group by the comparable rates in the placebo group, except for magnetic resonance imaging (MRI) disease burden, which was calculated as the difference in the median percent change between the treated and placebo groups. ^bSeverity = 1 point Expanded Disability Status Score progression, sustained for 3 months (in the IFN-β-1a 30 μg qw trial, this change was sustained for 6 months; in the IFN-β-1b trial, this was over 3 years). ^cDifferent studies measured these MRI measures differently, making comparisons difficult (numbers for new T2 represent the best case scenario for each trial). ^dNew lesions seen on T2-weighted MRI. ^e*p* = .001. ^f*p* = .01. ^g*p* = .05.

Abbreviations: DMF, dimethyl fumarate; FDA, U.S. Food and Drug Administration; FGM, fingolimod; GA, glatiramer acetate; IFN-β, interferon β; IM, intramuscular; IV, intravenous; MTX, mitoxantrone; NR, not reported; NS, not significant; NTZ, natalizumab; PO, oral; q3mo, once every 3 months; qd, daily; qmo, once per month; qod, every other day; qw, once per week; qyr, once per year; SC, subcutaneous; TF, teriflunomide; tiw, three times per week.

the balance between proinflammatory and regulatory cytokines. Glatiramer acetate reduces the attack rate (whether measured clinically or by MRI) in RRMS. Glatiramer acetate also benefits disease severity measures, although, for clinical disability, this is less well established than for IFN-β. Nevertheless, two very large head-to-head trials demonstrated that the impact of glatiramer acetate on clinical relapse rates and disability was comparable to high-dose, high-frequency IFN-β. Therefore, glatiramer acetate should be considered as an equally effective alternative to IFN-β in RRMS patients. Its usefulness in progressive disease is unknown. Glatiramer acetate is administered by subcutaneous injection of either 20 mg every day or 40 mg thrice weekly. Injection-site reactions also occur with glatiramer acetate. Initially, these were thought to be less severe than with IFN-β, although two recent head-to-head comparisons of high-dose, high-frequency IFN-β to daily glatiramer acetate did not bear out this impression. In addition, approximately 15% of patients experience one or more episodes of flushing, chest tightness, dyspnea, palpitations, and anxiety after injection. This systemic reaction is unpredictable, brief (duration <1 h), and tends not to recur. Finally, some patients experience lipoatrophy, which, on occasion, can be disfiguring and require cessation of treatment.

Natalizumab Natalizumab is a humanized monoclonal antibody directed against the α₄ subunit of α₄β₁ integrin, a cellular adhesion molecule expressed on the surface of lymphocytes. It prevents lymphocytes from binding to endothelial cells, thereby preventing lymphocytes from penetrating the BBB and entering the CNS. Natalizumab is highly effective in reducing the attack rate and significantly improves all measures of disease severity in MS (both clinical and MRI). Moreover, it is well-tolerated, and the dosing schedule of monthly intravenous infusions makes it very convenient for patients. However, progressive multifocal leukoencephalopathy (PML), a life-threatening condition resulting from infection by the John Cunningham (JC) virus, has occurred in approximately 0.3% of patients treated with natalizumab. The incidence of PML is very low in the first year of treatment but then rises by the second year to reach a level of about 2 cases per 1000 patients per year. Nevertheless, the measurement of antibodies against the JC virus in the serum can be used to stratify this risk. Thus, in patients who do not have these antibodies, the risk of PML is either minimal or nonexistent (as long as they remain JC antibody free). Conversely, in patients who have these antibodies (especially those who have them in high titer), the risk may be as high as 0.6% or greater. The risk is also high in patients who have previously received immunosuppressive therapy. Natalizumab is currently recommended only for JC antibody-negative patients, unless they have failed

alternative therapies or if they have a particularly aggressive disease course. Head-to-head data show that natalizumab is superior to low-dose (weekly) IFN-β-1a in RRMS. However, its relative efficacy compared to other agents has not been established conclusively.

Natalizumab, 300 mg, is administered by IV infusion each month. Treatment with natalizumab is, in general, well tolerated. A small percentage (<10%) of patients experience hypersensitivity reactions (including anaphylaxis), and ~6% develop neutralizing antibodies to the molecule (only half of which persist).

The major concern with long-term treatment is the risk of PML. Approximately half of the adult population is JC antibody positive, indicating that they experienced an asymptomatic infection with the JC virus at some time in the past. Nevertheless, because the risk is extremely low during the first year of treatment with natalizumab (regardless of antibody status), natalizumab can still be used safely in JC antibody-positive patients for a period of 12 months. After this time, in antibody-positive patients, a change to another disease-modifying therapy should be strongly considered. By contrast, persistently antibody-negative patients can be continued on treatment indefinitely. Up to 2% of seronegative MS patients undergoing treatment with natalizumab seroconvert annually; thus it is recommended that JC antibody status be assessed at 6-month intervals in all patients receiving treatment with this agent.

Fingolimod Fingolimod is a sphingosine-1-phosphate (S1P) inhibitor that prevents the egress of lymphocytes from the secondary lymphoid organs such as the lymph nodes and spleen. Its mechanism of action is probably due, in part, to the trapping of lymphocytes in the periphery and inhibiting their trafficking to the CNS. Fingolimod reduces the attack rate and significantly improves all measures of disease severity in MS. It is well tolerated, and the daily oral dosing schedule makes it very convenient for patients. A large head-to-head phase 3 randomized study demonstrated the superiority of fingolimod over low-dose (weekly) IFN-β-1a. However, its relative efficacy compared to other agents has not been established conclusively.

Fingolimod, 0.5 mg, is administered orally each day. Treatment with fingolimod is also, in general, well tolerated. Mild abnormalities on routine laboratory evaluation (e.g., elevated liver function tests or lymphopenia) are more common than in controls, sometimes requiring discontinuation of the medication. First- and second-degree heart block and bradycardia can also occur when fingolimod therapy is initiated. A 6-h period of observation (including electrocardiogram monitoring) is recommended for all patients receiving their first dose, and individuals with preexisting cardiac disease should probably not be treated with this agent. Other side effects

include macular edema and, rarely, disseminated varicella-zoster virus (VZV) infection; prior to initiating therapy with fingolimod, an ophthalmic exam and VZV vaccination for seronegative individuals are indicated.

Dimethyl Fumarate (DMF) Although the precise mechanisms of action of DMF are not fully understood, it seems to have anti-inflammatory effects through its modulation of the expression of proinflammatory and anti-inflammatory cytokines. Also, DMF inhibits the ubiquitylation and degradation of nuclear factor E2-related factor 2 (Nrf2)—a transcription factor that binds to the antioxidant response elements (AREs) located on the DNA and thereby induces the transcription of several antioxidant proteins. DMF reduces the attack rate and significantly improves all measures of disease severity in MS patients. However, its twice-daily oral dosing schedule makes it somewhat less convenient for patients than daily oral therapies. In addition, compliance is likely to be less with a twice-daily dosing regimen—a factor that could be of concern given the observation (in a small clinical trial) that once-daily DMF lacks efficacy. A head-to-head trial provided evidence that DMF was superior to glatiramer acetate on some outcome measures.

DMF, 240 mg, is administered orally twice each day. Gastrointestinal side effects (abdominal discomfort, nausea, vomiting, flushing, and diarrhea) are common at the start of therapy but generally subside with continued administration. Other adverse events included mild decreases in neutrophil and lymphocyte counts and mild elevations in liver enzymes. Nevertheless, in general, treatment with DMF is well tolerated after an initial period of adjustment. Following the release of DMF, four cases of PML were reported in patients receiving other products (not Tecfidera) that contained DMF. Each of these patients was lymphocytopenic, and most had received previous immunosuppressant therapy so that the relationship of DMF to the PML (if any) in these cases is uncertain. Nevertheless, these reports underscore the fact, stated previously, that long-term safety can never be guaranteed by the results of short-term trials. In the case of DMF for MS, only time and experience will tell us whether or not there is any cause for concern.

Teriflunomide Teriflunomide inhibits the mitochondrial enzyme dihydro-orotate dehydrogenase, which is a key part of the pathway for de novo pyrimidine biosynthesis from carbamoyl phosphate and aspartate. It is the active metabolite of the drug leflunomide (FDA-approved for rheumatoid arthritis), and it exerts its anti-inflammatory effects by limiting the proliferation of rapidly dividing T and B cells. This enzyme is not involved in the so-called “salvage pathway,” by which existing pyrimidine pools are recycled for DNA and RNA synthesis in resting and homeostatically proliferating cells. Consequently, teriflunomide is considered to be cytostatic rather than cytotoxic. Teriflunomide reduces the attack rate and significantly improves all measures of disease severity in MS patients. It is well tolerated, and its daily oral dosing schedule makes it very convenient for patients. A head-to-head trial suggested the equivalence, but not superiority, of teriflunomide and high-dose (thrice-weekly) IFN- β -1a. Teriflunomide, either 7 or 14 mg, is administered orally each day. In the pivotal clinical trials, mild hair thinning and gastrointestinal symptoms (nausea and diarrhea) were more common than in controls, but in general, treatment with teriflunomide was well tolerated. As with any new agent, the long-term safety is not guaranteed by the results of short-term trials. A major limitation, especially in women of childbearing age, is its possible teratogenicity (pregnancy category X); teriflunomide can remain in the bloodstream for 2 years, and it is recommended that exposed men and women who wish to conceive receive cholestyramine or activated charcoal to eliminate residual drug.

Mitoxantrone Hydrochloride Mitoxantrone, an anthracenedione, exerts its antineoplastic action by (1) intercalating into DNA and producing both strand breaks and interstrand cross-links, (2) interfering with RNA synthesis, and (3) inhibiting topoisomerase II (involved in DNA repair). The FDA approved mitoxantrone on the basis of a

single (relatively small) phase 3 clinical trial in Europe, in addition to an even smaller phase 2 study completed earlier. Mitoxantrone received (from the FDA) the broadest indication of any current treatment for MS. Thus, mitoxantrone is indicated for use in SPMS, in PRMS, and in patients with worsening RRMS (defined as patients whose neurologic status remains significantly abnormal between MS attacks). Despite this broad indication, however, the data supporting its efficacy are weaker than for other approved therapies.

Mitoxantrone can be cardiotoxic (e.g., cardiomyopathy, reduced left ventricular ejection fraction, and irreversible congestive heart failure). As a result, a cumulative dose <140 mg/m² is not recommended. At currently approved doses (12 mg/m² every 3 months), the maximum duration of therapy can be only 2–3 years. Furthermore, $>40\%$ of women will experience amenorrhea, which may be permanent. Finally, there is risk of acute leukemia from mitoxantrone, estimated as at least a 1% lifetime risk, and this complication has been reported in several mitoxantrone-treated MS patients.

Because of these risks, and a growing list of alternative therapies, mitoxantrone is now only rarely used for MS. It should not be used as a first-line agent in either RRMS or relapsing SPMS, but might be considered in selected patients with a progressive course who have failed other therapies.

Alemtuzumab Alemtuzumab is a humanized monoclonal antibody directed against the CD52 antigen, which is expressed on both monocytes and lymphocytes. It causes lymphocyte depletion (of both B and T cells) and a change in the composition of lymphocyte subsets. Both of these changes, particularly the impact on lymphocyte subsets, are long lasting. In preliminary trials, alemtuzumab markedly reduced the attack rate and significantly improved all measures of disease severity in MS patients. In two phase 3 trials, however, its impact on clinical disability was less convincing. Notably, both trials used the active comparator of thrice-weekly, high-dose IFN- β -1a. The European and Canadian drug agencies were the first to approve this agent for use in RRMS; the FDA has also approved alemtuzumab, but only after an appeal following initial disapproval. The reasons for the initial disapproval were based on a perceived lack of a convincing disability effect and concerns over potential toxicity. The toxicities of concern were the occurrence (during the trial or thereafter) of (1) autoimmune diseases including thyroiditis, Graves' disease, thrombocytopenia, hemolytic anemia, pancytopenia, antiglomerular basement membrane disease, and membranous glomerulonephritis; (2) malignancies including thyroid cancer, melanoma, breast cancer, and human papillomavirus (HPV)-related cancers; (3) serious infections; and (4) infusion reactions.

Initiating and Changing Treatment Previously, most patients with relapsing forms of MS received injectable agents (IFN- β or glatiramer acetate) as first-line therapy. However, with the introduction of effective and probably safe oral agents, including DMF, fingolimod, and teriflunomide, this has begun to change. In addition, the monthly infusion therapy natalizumab, which is highly effective, well tolerated, and apparently safe in JC antibody-negative patients, provides an attractive option in many cases. As noted above, with the exception of the first-generation injectable agents, long-term safety data are not available, and for the most part, comparative data are lacking. The value of combination therapy is also largely unknown, although a recent clinical trial demonstrated no added benefit to the combination of glatiramer acetate with low-dose, once-weekly IFN- β -1a.

Despite these unknowns, clinicians need to make decisions based on the best available evidence, coupled with practical considerations. One reasonable approach stratifies initial decision-making based on two levels of disease aggressiveness ([Fig. 458-4](#)).

MILD INITIAL COURSE In the case of recent onset, normal exam or minimal impairment (EDSS ≤ 2.5 or less), or low disease activity, either an injectable (IFN- β or glatiramer acetate) or an oral (DMF,

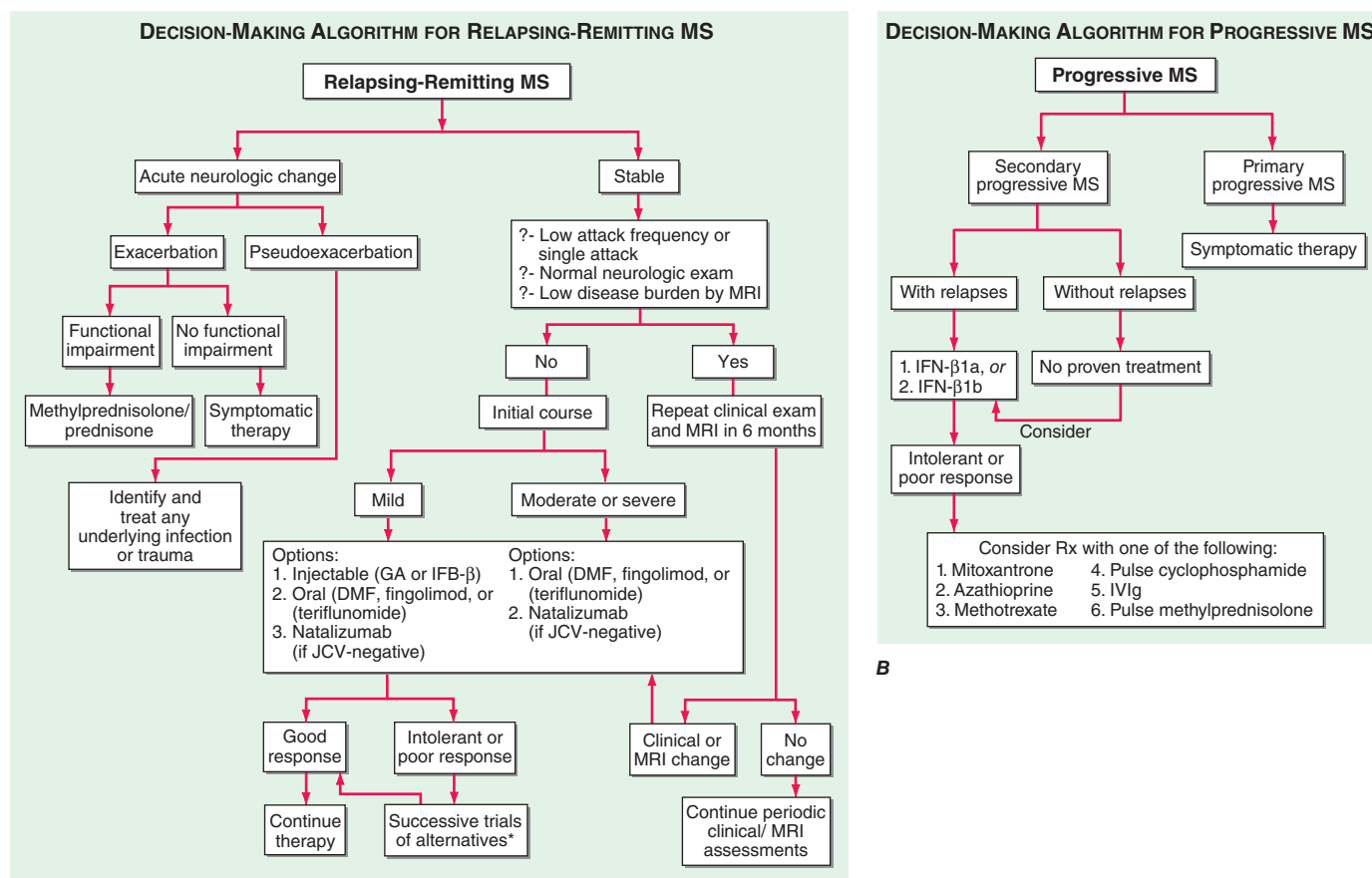


FIGURE 458-4 Therapeutic decision-making for multiple sclerosis (MS). *Can include trials of different preparations of interferon β (IFN- β), particularly advancing from once-weekly (Avonex) to a more frequent (e.g., Rebif, Betaseron/Extavia) dosing regimen. Options also include use of natalizumab in JC virus–positive patients. MRI, magnetic resonance imaging.

fingolimod, or teriflunomide) agent is reasonable. Although head-to-head comparisons are not available, natalizumab is thought to be more effective than these other agents, and therefore, this therapy can be considered even in minimally affected, JCV antibody–seronegative patients. The injectable agents (IFN- β and glatiramer acetate) have a superb track record for safety but have a high nuisance factor due to the need for frequent injections, as well as bothersome side effects that contribute to noncompliance. Some of the oral agents (DMF and fingolimod) are probably more effective than the injectables, but long-term risks are mostly unknown; DMF produces bothersome gastrointestinal symptoms in many patients at least initially (can be mitigated by beginning at one-quarter strength and gradually advancing to full dose), and fingolimod can lead to bradycardia and other cardiac disturbances of unclear clinical significance. Teriflunomide may be less effective than the other oral agents, and there are concerns about its possible long-lasting pregnancy risks. Nevertheless, its long-term safety has likely been established because of its extensive human exposure as the active metabolite of leflunomide—a drug long approved by the FDA.

MODERATE OR SEVERE INITIAL COURSE In highly active disease or moderate impairment (EDSS >2.5), either a highly effective oral agent (DMF or fingolimod) or, if the patient is JC virus antibody seronegative, infusion therapy with natalizumab is recommended.

Regardless of which agent is chosen first, treatment should probably be changed in patients who continue to have relapses, progressive neurologic impairment or, arguably, ongoing evidence of subclinical MRI activity (Fig. 458-4).

The long-term impact of these treatments on the disease course remains controversial, although several recent studies have shown that these agents improve the long-term outcome of MS including a prolongation of the time to reach certain disability outcomes

(e.g., SPMS and requiring assistance to ambulate) and reduction in MS-related mortality. These benefits seem most conspicuous when treatment begins early in the RRMS stage of the illness. Unfortunately, however, already established progressive symptoms do not respond well to treatment with these disease-modifying therapies. Because progressive symptoms are likely to result from accumulated axonal and neuronal loss, many experts now believe that very early treatment with a disease-modifying drug is appropriate for most MS patients. It may also be reasonable to delay initiating treatment in patients with (1) normal neurologic exams, (2) a single attack or a low attack frequency, and (3) a low burden of disease as assessed by brain MRI. Untreated patients, however, should be followed closely with periodic brain MRI scans; the need for therapy is reassessed if scans reveal evidence of ongoing, subclinical disease. Finally, vitamin D deficiency should be corrected in all patients with MS, and generally this requires oral supplementation with vitamin D₃, 4000 to 5000 IU daily.

DISEASE-MODIFYING THERAPIES FOR PROGRESSIVE MS

SPMS High-dose IFN- β probably has a beneficial effect in patients with SPMS who are still experiencing acute relapses. IFN- β is probably ineffective in patients with SPMS who are not having acute attacks. All of the other agents have not yet been studied in this patient population. Although mitoxantrone has been approved for patients with progressive MS, this is not the population studied in the pivotal trial. Therefore no evidence-based recommendation can be made with regard to its use in this setting.

PPMS No therapies have been convincingly shown to modify the course of PPMS. A phase 3 clinical trial of glatiramer acetate in PPMS was stopped because of lack of efficacy. A phase 2/3 trial of the monoclonal antibody rituximab (anti-CD20) in PPMS was

also negative, but in a preplanned secondary analysis, treatment appeared to modestly slow disability progression in patients with Gd-enhancing lesions at entry; the results of a follow-up trial with a fully humanized monoclonal anti-CD20 therapy (ocrelizumab) will soon be available.

OFF-LABEL TREATMENT OPTIONS FOR RRMS AND SPMS

Azathioprine (2–3 mg/kg per day) has been used primarily in SPMS. Meta-analysis of published trials suggests that azathioprine is marginally effective at lowering relapse rates, although a benefit on disability progression has not been demonstrated.

Methotrexate (7.5–20 mg/week) was shown in one study to slow the progression of upper extremity dysfunction in SPMS. Because of the possibility of developing irreversible liver damage, some experts recommend a blind liver biopsy after 2 years of therapy.

Cyclophosphamide (700 mg/m², every other month) may be helpful for treatment-refractory patients who are (1) otherwise in good health, (2) ambulatory, and (3) <40 years of age. Because cyclophosphamide can be used for periods in excess of 3 years, it may be preferable to mitoxantrone in these circumstances.

Intravenous immunoglobulin (IVIg), administered in monthly pulses (up to 1 g/kg) for up to 2 years, appears to reduce annual exacerbation rates. However, its use is limited because of its high cost, questions about optimal dose, and uncertainty about its having any impact on long-term disability.

Methylprednisolone, administered in one study as monthly high-dose intravenous pulses, reduced disability progression (see above).

OTHER THERAPEUTIC CLAIMS

Many purported treatments for MS have never been subjected to scientific scrutiny. These include dietary therapies (e.g., the Swank diet, in addition to others), megadose vitamins, calcium orotate, bee stings, cow colostrum, hyperbaric oxygen, procarin (a combination of histamine and caffeine), chelation, acupuncture, acupressure, various Chinese herbal remedies, and removal of mercury-amalgam tooth fillings, among many others. Patients should avoid costly or potentially hazardous unproven treatments. Many such treatments lack biologic plausibility. For example, no reliable case of mercury poisoning resembling typical MS has ever been described.

Although potential roles for EBV, human herpesvirus (HHV) 6, or chlamydia have been suggested for MS, these reports are unconfirmed, and treatment with antiviral agents or antibiotics is not recommended.

Most recently, chronic cerebrospinal insufficiency (CCSVI) has been proposed as a cause of MS, and vascular-surgical intervention is recommended. However, the failure of independent investigators to even approximate the initial claims of 100% sensitivity and 100% specificity for the diagnostic procedure have raised considerable doubt that CCSVI is a real entity. Certainly, any potentially dangerous surgery should be avoided until more rigorous science is available.

SYMPTOMATIC THERAPY

For all patients, it is useful to encourage attention to a healthy lifestyle, including maintaining an optimistic outlook, a healthy diet, and regular exercise as tolerated (swimming is often well-tolerated because of the cooling effect of cold water). It is reasonable also to correct vitamin D deficiency with oral vitamin D and to recommend dietary supplementation with long-chain (omega-3) unsaturated fatty acids (present in oily fish such as salmon), because of their biologic plausibility for MS pathogenesis, safety, and general health benefits.

Ataxia/tremor is often intractable. Clonazepam, 1.5–20 mg/d; primidone, 50–250 mg/d; propranolol, 40–200 mg/d; or ondansetron, 8–16 mg/d, may help. Wrist weights occasionally reduce tremor in the arm or hand. Thalamotomy or deep-brain stimulation has been tried with mixed success.

Spasticity and *spasms* may improve with physical therapy, regular exercise, and stretching. Avoidance of triggers (e.g., infections, fecal impactions, bed sores) is extremely important. Effective medications

include baclofen (20–120 mg/d), diazepam (2–40 mg/d), tizanidine (8–32 mg/d), dantrolene (25–400 mg/d), and cyclobenzaprine hydrochloride (10–60 mg/d). For severe spasticity, a baclofen pump (delivering medication directly into the CSF) can provide substantial relief.

Weakness can sometimes be improved with the use of potassium channel blockers such as 4-aminopyridine (10–40 mg/d) and 3,4-diaminopyridine (40–80 mg/d), particularly in the setting where lower extremity weakness interferes with the patient's ability to ambulate. The FDA has approved 4-aminopyridine (at 20 mg/d), and this can be obtained either as dalfampridine (Ampyra) or, more cheaply, through a compounding pharmacy. The principle concern with the use of these agents is the possibility of inducing seizures at high doses.

Pain is treated with anticonvulsants (carbamazepine, 100–1000 mg/d; phenytoin, 300–600 mg/d; gabapentin, 300–3600 mg/d; or pregabalin, 50–300 mg/d), antidepressants (amitriptyline, 25–150 mg/d; nortriptyline, 25–150 mg/d; desipramine, 100–300 mg/d; or venlafaxine, 75–225 mg/d), or antiarrhythmics (mexiletine, 300–900 mg/d). If these approaches fail, patients should be referred to a comprehensive pain management program.

Bladder dysfunction management is best guided by urodynamic testing. Evening fluid restriction or frequent voluntary voiding may help *detrusor hyperreflexia*. If these methods fail, propantheline bromide (10–15 mg/d), oxybutynin (5–15 mg/d), hyoscyamine sulfate (0.5–0.75 mg/d), tolterodine tartrate (2–4 mg/d), or solifenacin (5–10 mg/d) may help. Coadministration of pseudoephedrine (30–60 mg) is sometimes beneficial.

Detrusor/sphincter dyssynergia may respond to phenoxybenzamine (10–20 mg/d) or terazosin hydrochloride (1–20 mg/d). Loss of reflex bladder wall contraction may respond to bethanechol (30–150 mg/d). However, both conditions often require catheterization.

Urinary tract infections should be treated promptly. Patients with large postvoid residual urine volumes are predisposed to infections. Prevention by urine acidification (with cranberry juice or vitamin C) inhibits some bacteria. Prophylactic administration of antibiotics is sometimes necessary but may lead to colonization by resistant organisms. Intermittent catheterization may help to prevent recurrent infections.

Treatment of *constipation* includes high-fiber diets and fluids. Natural or other laxatives may help. *Fecal incontinence* may respond to a reduction in dietary fiber.

Depression should be treated. Useful drugs include the selective serotonin reuptake inhibitors (fluoxetine, 20–80 mg/d, or sertraline, 50–200 mg/d), the tricyclic antidepressants (amitriptyline, 25–150 mg/d; nortriptyline, 25–150 mg/d; or desipramine, 100–300 mg/d), and the nontricyclic antidepressants (venlafaxine, 75–225 mg/d).

Fatigue may improve with assistive devices, help in the home, or successful management of spasticity. Patients with frequent nocturia may benefit from anticholinergic medication at bedtime. Primary MS fatigue may respond to amantadine (200 mg/d), methylphenidate (5–25 mg/d), or modafinil (100–400 mg/d).

Cognitive problems may respond to the cholinesterase inhibitor donepezil hydrochloride (10 mg/d).

Paroxysmal symptoms respond dramatically to low-dose anticonvulsants (acetazolamide, 200–600 mg/d; carbamazepine, 50–400 mg/d; phenytoin, 50–300 mg/d; or gabapentin, 600–1800 mg/d).

Heat sensitivity may respond to heat avoidance, air-conditioning, or cooling garments.

Sexual dysfunction may be helped by lubricants to aid in genital stimulation and sexual arousal. Management of pain, spasticity, fatigue, and bladder/bowel dysfunction may also help. Sildenafil (50–100 mg), tadalafil (5–20 mg), or vardenafil (5–20 mg), taken 1–2 h before sex, is now the standard treatment for maintaining erections.

PROMISING EXPERIMENTAL THERAPIES

Numerous clinical trials are currently under way. These include studies on (1) monoclonal antibodies against CD20 to deplete B cells and against the IL-2 receptor; (2) selective oral

sphingosine-1-phosphate receptor antagonists to sequester lymphocytes in secondary lymphoid organs; (3) estriol to induce a pregnancy-like state; (4) molecules to promote remyelination; and (4) bone marrow transplantation.

CLINICAL VARIANTS OF MS

Acute MS (Marburg's variant) is a fulminant demyelinating process that in some cases progresses inexorably to death within 1–2 years. Typically, there are no remissions. When acute MS presents as a solitary, usually cavitary, lesion, a brain tumor is often suspected. In such cases, a brain biopsy is usually required to establish the diagnosis. An antibody-mediated process appears to be responsible for most cases. Marburg's variant does not seem to follow infection or vaccination, and it is unclear whether this syndrome represents an extreme form of MS or another disease altogether. No controlled trials of therapy exist; high-dose glucocorticoids, plasma exchange, and cyclophosphamide have been tried, with uncertain benefit.

NEUROMYELITIS OPTICA

Neuromyelitis optica (NMO; Devic's disease) is an aggressive inflammatory disorder characterized by recurrent attacks of ON and myelitis (Table 458-7). NMO is more frequent in women than men (>3:1), typically begins in childhood or early adulthood but can arise at any age, and is uncommon in whites compared with individuals of Asian and African ancestry. Attacks of ON can be bilateral (rare in MS) or unilateral; myelitis can be severe and transverse (rare in MS) and is typically longitudinally extensive, involving three or more contiguous vertebral segments. Also in contrast to MS, progressive symptoms do not occur in NMO. The brain MRI was earlier thought to be normal in NMO, but it is now recognized that in approximately half of cases, there are lesions involving the hypothalamus causing an endocrinopathy; the lower brainstem presenting as intractable hiccoughs or vomiting from involvement of the area postrema in the lower medulla; or the cerebral hemispheres producing focal symptoms, encephalopathy, or seizures. Large MRI lesions in the cerebral hemispheres can be asymptomatic, sometimes have a "cloud-like" appearance and, unlike MS lesions, are often not destructive, and can resolve completely. Spinal cord MRI lesions typically consist of focal enhancing areas of swelling and tissue destruction, extending over three or more spinal cord segments, and on axial sequences, these are centered on the gray matter of the cord. CSF findings include pleocytosis greater than that observed in MS, with neutrophils and eosinophils present in some cases; OCBs are uncommon, occurring in fewer than 30% of NMO patients. The pathology of NMO is a distinctive astrocytopathy with inflammation, a loss of astrocytes, and an absence of staining of the water channel protein aquaporin-4 by immunohistochemistry, plus thickened blood vessel walls, demyelination, and deposition of antibody and complement.

NMO is best understood as a syndrome with diverse causes. Up to 40% of patients have a systemic autoimmune disorder, often systemic lupus erythematosus, Sjögren's syndrome, perinuclear antineutrophil cytoplasmic antibody (p-ANCA)-associated vasculitis, myasthenia gravis, Hashimoto's thyroiditis, or mixed connective tissue disease. In others, onset may be associated with acute infection with VZV, EBV, HIV, or tuberculosis. Rare cases appear to be paraneoplastic and

associated with breast, lung, or other cancers. NMO is often idiopathic, however. NMO is usually disabling over time; in one series, respiratory failure from cervical myelitis was present in one-third of patients, and 8 years after onset, 60% of patients were blind and more than half had permanent paralysis of one or more limbs.

A highly specific autoantibody directed against aquaporin-4 is present in the sera of approximately two-thirds of patients with a clinical diagnosis of NMO. Seropositive patients have a very high risk for future relapses; more than half will relapse within 1 year if untreated. Aquaporin-4 is localized to the foot processes of astrocytes in close apposition to endothelial surfaces, as well as at paranodal regions near nodes of Ranvier. It is likely that aquaporin-4 antibodies are pathogenic, as passive transfer of antibodies from NMO patients into laboratory animals reproduce histologic features of the disease.

When MS affects individuals of African or Asian ancestry, there is a propensity for demyelinating lesions to involve predominantly the optic nerve and spinal cord, an MS subtype termed *opticospinal MS*. Interestingly, some individuals with opticospinal MS are seropositive for aquaporin-4 antibodies, suggesting that such cases represent an NMO spectrum disorder.

TREATMENT NEUROMYELITIS OPTICA

Disease-modifying therapies have not been rigorously studied in NMO. Acute attacks of NMO are usually treated with high-dose glucocorticoids (solumedrol 1–2 g/d for 5–10 days followed by a prednisone taper). Plasma exchange (typically 7 qod exchanges of 1.5 plasma volumes) has also been used empirically for acute episodes that fail to respond to glucocorticoids. Given the unfavorable natural history of untreated NMO, prophylaxis against relapses is recommended for most patients using one of the following regimens: mycophenylate mofetil (250 mg bid gradually increasing to 1000 mg bid); B cell depletion with anti-CD20 monoclonal antibody (rituximab); or a combination of glucocorticoids (500 mg IV methylprednisolone daily for 5 days; then oral prednisone 1 mg/kg per day for 2 months, followed by slow taper) plus azathioprine (2 mg/kg per day started on week 3). Available evidence suggests that use of IFN- β is ineffective and paradoxically may increase the risk of NMO relapses.

ACUTE DISSEMINATED ENCEPHALOMYELITIS

Acute disseminated encephalomyelitis (ADEM) has a monophasic course and is most frequently associated with an antecedent infection (postinfectious encephalomyelitis); approximately 5% of ADEM cases follow immunization (postvaccinal encephalomyelitis). ADEM is far more common in children than adults, and many adult cases initially thought to represent ADEM subsequently experience late relapses qualifying as either MS or another chronic inflammatory disorder such as vasculitis, sarcoid, or lymphoma. The hallmark of ADEM is the presence of widely scattered small foci of perivenular inflammation and demyelination, in contrast to larger confluent demyelinating lesions typical of MS. In the most explosive form of ADEM, acute hemorrhagic leukoencephalitis, the lesions are vasculitic and hemorrhagic, and the clinical course is devastating.

Postinfectious encephalomyelitis is most frequently associated with the viral exanthems of childhood. Infection with measles virus is the most common antecedent (1 in 1000 cases). Worldwide, measles encephalomyelitis is still common, although use of the live measles vaccine has dramatically reduced its incidence in developed countries. An ADEM-like illness rarely follows vaccination with live measles vaccine (1–2 in 10⁶ immunizations). ADEM is now most frequently associated with varicella (chickenpox) infections (1 in 4000–10,000 cases). It may also follow infection with rubella, mumps, influenza, parainfluenza, EBV, HHV-6, HIV, other viruses, and *Mycoplasma pneumoniae*. Some patients may have a nonspecific upper respiratory infection or no known antecedent illness. In addition to measles, postvaccinal encephalomyelitis may also follow the administration

TABLE 458-7 DIAGNOSTIC CRITERIA FOR NEUROMYELITIS OPTICA

Required:

1. Optic neuritis
2. Acute transverse myelitis

Supportive (2 of 3 criteria required):

1. Longitudinally extensive cord lesion extending over 3 or more vertebral segments
2. Brain magnetic resonance imaging normal or not meeting criteria for multiple sclerosis
3. Aquaporin-4 seropositivity

Source: Adapted from DM Wingerchuk et al: *Neurology* 66:1485, 2006.

2674 of vaccines for smallpox (5 cases per million), the Semple rabies, and Japanese encephalitis. Modern vaccines that do not require viral culture in CNS tissue have reduced the ADEM risk.

All forms of ADEM presumably result from a cross-reactive immune response to the infectious agent or vaccine that then triggers an inflammatory demyelinating response. Autoantibodies to MBP and to other myelin antigens have been detected in the CSF from some patients with ADEM. Attempts to demonstrate direct viral invasion of the CNS have been unsuccessful.

CLINICAL MANIFESTATIONS

In severe cases, onset is abrupt and progression rapid (hours to days). In postinfectious ADEM, the neurologic syndrome generally begins late in the course of the viral illness as the exanthem is fading. Fever reappears, and headache, meningismus, and lethargy progressing to coma may develop. Seizures are common. Signs of disseminated neurologic disease are consistently present (e.g., hemiparesis or quadriparesis, extensor plantar responses, lost or hyperactive tendon reflexes, sensory loss, and brainstem involvement). In ADEM due to chickenpox, cerebellar involvement is often conspicuous. CSF protein is modestly elevated (0.5–1.5 g/L [50–150 mg/dL]). Lymphocytic pleocytosis, generally 200 cells/ μ L or greater, occurs in 80% of patients. Occasional patients have higher counts or a mixed polymorphonuclear-lymphocytic pattern during the initial days of the illness. Transient CSF oligoclonal banding has been reported. MRI usually reveals extensive changes in the brain and spinal cord, consisting of white matter hyperintensities on T2 and fluid-attenuated inversion recovery sequences with Gd enhancement on T1-weighted sequences.

DIAGNOSIS

The diagnosis is most reliably established when there is a history of recent vaccination or viral exanthematous illness. In severe cases

with predominantly cerebral involvement, acute encephalitis due to infection with herpes simplex or other viruses including HIV may be difficult to exclude (**Chap. 164**); other considerations include hypercoagulable states including the antiphospholipid antibody syndrome, vasculitis, neurosarcoïd, primary CNS lymphoma, or metastatic cancer. An explosive presentation of MS can mimic ADEM, and especially in adults, it may not be possible to distinguish these conditions at onset. The simultaneous onset of disseminated symptoms and signs is common in ADEM and rare in MS. Similarly, meningismus, drowsiness, coma, and seizures suggest ADEM rather than MS. Unlike MS, in ADEM, optic nerve involvement is generally bilateral and transverse myelopathy complete. MRI findings that favor ADEM include extensive and relatively symmetric white matter abnormalities, basal ganglia or cortical gray matter lesions, and Gd enhancement of all abnormal areas. By contrast, OCBs in the CSF are more common in MS. In one study of adult patients initially thought to have ADEM, 30% experienced additional relapses over a follow-up period of 3 years, and they were reclassified as having MS. Occasional patients with “recurrent ADEM” have also been reported, especially children; however, it is not possible to distinguish this entity from atypical MS.

TREATMENT ACUTE DISSEMINATED ENCEPHALOMYELITIS

Initial treatment is with high-dose glucocorticoids as for exacerbations of NMO (see above); depending on the response, treatment may need to be continued for 8 weeks. Patients who fail to respond within a few days may benefit from a course of plasma exchange or intravenous immunoglobulin. The prognosis reflects the severity of the underlying acute illness. In recent case series of presumptive ADEM in adults, mortality rates of 5–20% are reported, and many survivors have permanent neurologic sequelae.

SECTION 3 NERVE AND MUSCLE DISORDERS

459 Peripheral Neuropathy

Anthony A. Amato, Richard J. Barohn

Peripheral nerves are composed of sensory, motor, and autonomic elements. Diseases can affect the cell body of a neuron or its peripheral processes, namely the axons or the encasing myelin sheaths. Most peripheral nerves are mixed and contain sensory and motor as well as autonomic fibers. Nerves can be subdivided into three major classes: large myelinated, small myelinated, and small unmyelinated. Motor axons are usually large myelinated fibers that conduct rapidly (approximately 50 m/s). Sensory fibers may be any of the three types. Large-diameter sensory fibers conduct proprioception and vibratory sensation to the brain, while the smaller-diameter myelinated and unmyelinated fibers transmit pain and temperature sensation. Autonomic nerves are also small in diameter. Thus, peripheral neuropathies can impair sensory, motor, or autonomic function, either singly or in combination. Peripheral neuropathies are further classified into those that primarily affect the cell body (e.g., neuronopathy or ganglionopathy), myelin (myelinopathy), and the axon (axonopathy). These different classes of peripheral neuropathies have distinct clinical and electrophysiologic features. This chapter discusses the clinical approach to a patient suspected of having a peripheral neuropathy, as well as specific neuropathies, including hereditary and acquired neuropathies. **The inflammatory neuropathies are discussed in Chap. 460.**

GENERAL APPROACH

In approaching a patient with a neuropathy, the clinician has three main goals: (1) identify where the lesion is, (2) identify the cause, and (3) determine the proper treatment. The first goal is accomplished by obtaining a thorough history, neurologic examination, and electrodiagnostic and other laboratory studies (**Fig. 459-1**). While gathering this information, seven key questions are asked (**Table 459-1**), the answers to which can usually identify the category of pathology that is present (**Table 459-2**). Despite an extensive evaluation, in approximately half of patients, no etiology is ever found; these patients typically have a predominately sensory polyneuropathy and have been labeled as having idiopathic or cryptogenic sensory polyneuropathy (CSPN).

INFORMATION FROM THE HISTORY AND PHYSICAL EXAMINATION: SEVEN KEY QUESTIONS (TABLE 459-1)

1. What Systems are Involved? It is important to determine if the patient’s symptoms and signs are motor, sensory, autonomic, or a combination of these. If the patient has only weakness without any evidence of sensory or autonomic dysfunction, a motor neuropathy, neuromuscular junction abnormality, or myopathy should be considered. Some peripheral neuropathies are associated with significant autonomic nervous system dysfunction. Symptoms of autonomic involvement include fainting spells or orthostatic lightheadedness; heat intolerance; or any bowel, bladder, or sexual dysfunction (**Chap. 454**). There will typically be an orthostatic fall in blood pressure without an appropriate increase in heart rate. Autonomic dysfunction in the absence of diabetes should

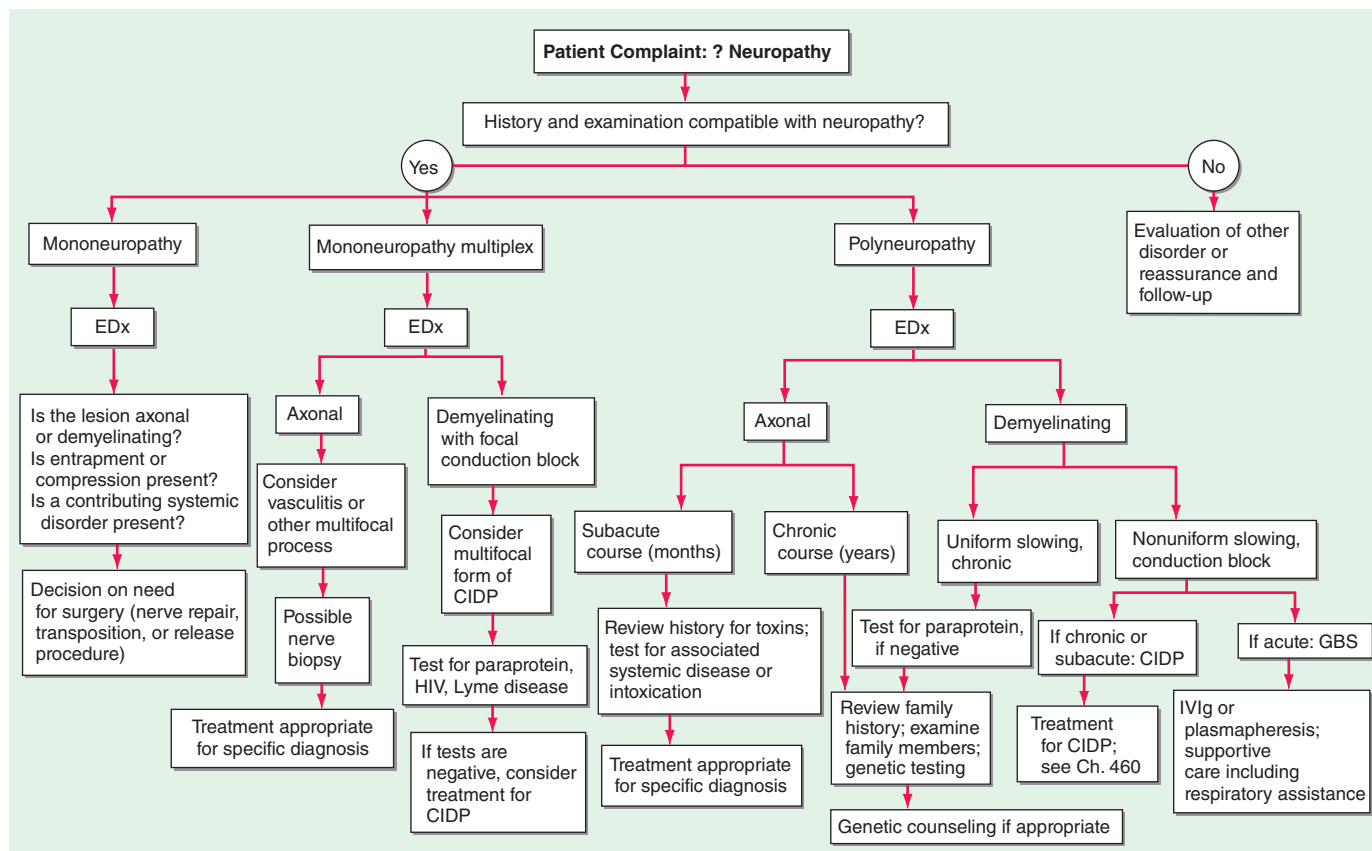


FIGURE 459-1 Approach to the evaluation of peripheral neuropathies. CIDP, chronic inflammatory demyelinating polyradiculoneuropathy; EDx, electrodiagnostic; GBS, Guillain-Barré syndrome; IVIg, intravenous immunoglobulin.

alert the clinician to the possibility of amyloid polyneuropathy. Rarely, a pandysautonomic syndrome can be the only manifestation of a peripheral neuropathy without other motor or sensory findings. The majority of neuropathies are predominantly sensory in nature.

TABLE 459-1 APPROACH TO NEUROPATHIC DISORDERS: SEVEN KEY QUESTIONS

1. What systems are involved?

- Motor, sensory, autonomic, or combinations

2. What is the distribution of weakness?

- Only distal versus proximal and distal
- Focal/asymmetric versus symmetric

3. What is the nature of the sensory involvement?

- Temperature loss or burning or stabbing pain (e.g., small fiber)
- Vibratory or proprioceptive loss (e.g., large fiber)

4. Is there evidence of upper motor neuron involvement?

- Without sensory loss
- With sensory loss

5. What is the temporal evolution?

- Acute (days to 4 weeks)
- Subacute (4–8 weeks)
- Chronic (>8 weeks)
- Monophasic, progressive, or relapsing-remitting

6. Is there evidence for a hereditary neuropathy?

- Family history of neuropathy
- Lack of sensory symptoms despite sensory signs

7. Are there any associated medical conditions?

- Cancer, diabetes mellitus, connective tissue disease or other autoimmune diseases, infection (e.g., HIV, Lyme disease, leprosy)
- Medications including over-the-counter drugs that may cause a toxic neuropathy
- Preceding events, drugs, toxins

2. What is the Distribution of Weakness? Delineating the pattern of weakness, if present, is essential for diagnosis, and in this regard two additional questions should be answered: (1) Does the weakness only involve the distal extremity, or is it both proximal and distal? and (2) Is the weakness focal and asymmetric, or is it symmetric? Symmetric proximal and distal weakness is the hallmark of acquired immune demyelinating polyneuropathies, both the acute form (acute inflammatory demyelinating polyneuropathy [AIDP], also known as Guillain-Barré syndrome [GBS]) and the chronic form (chronic inflammatory demyelinating polyneuropathy [CIDP]). The importance of finding symmetric proximal and distal weakness in a patient who presents with both motor and sensory symptoms cannot be over-emphasized because this identifies the important subset of patients who may have a treatable acquired demyelinating neuropathic disorder (i.e., AIDP or CIDP).

Findings of an asymmetric or multifocal pattern of weakness narrow the differential diagnosis. Some neuropathic disorders may present with unilateral extremity weakness. In the absence of sensory symptoms and signs, such weakness evolving over weeks or months would be worrisome for motor neuron disease (e.g., amyotrophic lateral sclerosis [ALS]), but it would be important to exclude multifocal motor neuropathy that may be treatable (Chap. 452). In a patient presenting with asymmetric subacute or acute sensory and motor symptoms and signs, radiculopathies, plexopathies, compressive mononeuropathies, or multiple mononeuropathies (e.g., mononeuropathy multiplex) must be considered.

3. What is the Nature of the Sensory Involvement? The patient may have loss of sensation (numbness), altered sensation to touch (hyperpathia or allodynia), or uncomfortable spontaneous sensations (tingling, burning, or aching) (Chap. 31). Neuropathic pain can be burning, dull, and poorly localized (protopathic pain), presumably transmitted by polymodal C nociceptor fibers, or sharp and lancinating (epicritic pain), relayed by A-delta fibers. If pain and temperature perception are lost, while vibratory and position sense are preserved along with

Pattern 1: Symmetric proximal and distal weakness with sensory loss

Consider: inflammatory demyelinating polyneuropathy (GBS and CIDP)

Pattern 2: Symmetric distal sensory loss with or without distal weakness

Consider: cryptogenic or idiopathic sensory polyneuropathy (CSPN), diabetes mellitus and other metabolic disorders, drugs, toxins, familial (HSAN), CMT, amyloidosis, and others

Pattern 3: Asymmetric distal weakness with sensory loss

With involvement of multiple nerves

Consider: multifocal CIDP, vasculitis, cryoglobulinemia, amyloidosis, sarcoid, infectious (leprosy, Lyme, hepatitis B, C, or E, HIV, CMV), HNPP, tumor infiltration

With involvement of single nerves/regions

Consider: may be any of the above but also could be compressive mononeuropathy, plexopathy, or radiculopathy

Pattern 4: Asymmetric proximal and distal weakness with sensory loss

Consider: polyradiculopathy or plexopathy due to diabetes mellitus, meningeal carcinomatosis or lymphomatosis, hereditary plexopathy (HNPP, HNA), idiopathic

Pattern 5: Asymmetric distal weakness without sensory loss

With upper motor neuron findings

Consider: motor neuron disease

Without upper motor neuron findings

Consider: progressive muscular atrophy, juvenile monomelic amyotrophy (Hirayama's disease), multifocal motor neuropathy, multifocal acquired motor axonopathy

Pattern 6: Symmetric sensory loss and distal areflexia with upper motor neuron findings

Consider: Vitamin B₁₂, vitamin E, and copper deficiency with combined system degeneration with peripheral neuropathy, hereditary leukodystrophies (e.g., adrenomyeloneuropathy)

Pattern 7: Symmetric weakness without sensory loss

With proximal and distal weakness

Consider: SMA

With distal weakness

Consider: hereditary motor neuropathy ("distal" SMA) or atypical CMT

Pattern 8: Asymmetric proprioceptive sensory loss without weakness

Consider causes of a sensory neuronopathy (ganglionopathy):

Cancer (paraneoplastic)

Sjögren's syndrome

Idiopathic sensory neuronopathy (possible GBS variant)

Cisplatin and other chemotherapeutic agents

Vitamin B₆ toxicity

HIV-related sensory neuronopathy

Pattern 9: Autonomic symptoms and signs

Consider neuropathies associated with prominent autonomic dysfunction:

Hereditary sensory and autonomic neuropathy

Amyloidosis (familial and acquired)

Diabetes mellitus

Idiopathic pandysautonomia (may be a variant of Guillain-Barré syndrome)

Porphyria

HIV-related autonomic neuropathy

Vincristine and other chemotherapeutic agents

Abbreviations: CIDP, chronic inflammatory demyelinating polyneuropathy; CMT, Charcot-Marie-Tooth disease; CMV, cytomegalovirus; GBS, Guillain-Barré syndrome; HIV, human immunodeficiency virus; HNA, hereditary neuralgic amyotrophy; HNPP, hereditary neuropathy with liability to pressure palsies; HSAN, hereditary sensory and autonomic neuropathy; SMA, spinal muscular atrophy.

muscle strength, deep tendon reflexes, and normal nerve conduction studies, a small-fiber neuropathy is likely. This is important, because the most likely cause of small-fiber neuropathies, when one is identified, is diabetes mellitus or glucose intolerance. Amyloid neuropathy should be considered as well in such cases, but most of these small-fiber neuropathies remain idiopathic in nature despite extensive evaluation.

Severe proprioceptive loss also narrows the differential diagnosis. Affected patients will note imbalance, especially in the dark. A neurologic examination revealing a dramatic loss of proprioception with vibration loss and normal strength should alert the clinician to consider a sensory neuronopathy/ganglionopathy (Table 459-2, Pattern 8). In particular, if this loss is asymmetric or affects the arms more than the legs, this pattern suggests a non-length-dependent process as seen in sensory neuronopathies.

4. Is There Evidence of Upper Motor Neuron Involvement? If the patient presents with symmetric distal sensory symptoms and signs suggestive of a distal sensory neuropathy, but there is additional evidence of symmetric upper motor neuron involvement (**Chap. 30**), the physician should consider a disorder such as combined system degeneration with neuropathy. The most common cause for this pattern is vitamin B₁₂ deficiency, but other causes of combined system degeneration with neuropathy should be considered (e.g., copper deficiency, HIV infection, severe hepatic disease, adrenomyeloneuropathy).

5. What is the Temporal Evolution? It is important to determine the onset, duration, and evolution of symptoms and signs. Does the disease have an acute (days to 4 weeks), subacute (4–8 weeks), or chronic (>8 weeks) course? Is the course monophasic, progressive, or relapsing? Most neuropathies are insidious and slowly progressive in nature. Neuropathies with acute and subacute presentations include GBS, vasculitis, and radiculopathies related to diabetes or Lyme disease. A relapsing course can be present in CIDP and porphyria.

6. Is There Evidence for a Hereditary Neuropathy? In patients with slowly progressive distal weakness over many years with very little in the way of sensory symptoms yet with significant sensory deficits on clinical examination, the clinician should consider a hereditary neuropathy (e.g., Charcot-Marie-Tooth disease [CMT]). On examination, the feet may show arch and toe abnormalities (high or flat arches, hammer-toes); scoliosis may be present. In suspected cases, it may be necessary to perform both neurologic and electrophysiologic studies on family members in addition to the patient.

7. Does the Patient Have Any Other Medical Conditions? It is important to inquire about associated medical conditions (e.g., diabetes mellitus, systemic lupus erythematosus); preceding or concurrent infections (e.g., diarrheal illness preceding GBS); surgeries (e.g., gastric bypass and nutritional neuropathies); medications (toxic neuropathy), including over-the-counter vitamin preparations (B₆); alcohol; dietary habits; and use of dentures (e.g., fixatives contain zinc that can lead to copper deficiency).

PATTERN RECOGNITION APPROACH TO NEUROPATHIC DISORDERS

Based on the answers to the seven key questions, neuropathic disorders can be classified into several patterns based on the distribution or pattern of sensory, motor, and autonomic involvement (Table 459-2). Each pattern has a limited differential diagnosis. A final diagnosis is established by using other clues such as the temporal course, presence of other disease states, family history, and information from laboratory studies.

ELECTRODIAGNOSTIC STUDIES

The electrodiagnostic (EDx) evaluation of patients with a suspected peripheral neuropathy consists of nerve conduction studies (NCS) and needle electromyography (EMG). In addition, studies of autonomic function can be valuable. The electrophysiologic data provide additional information about the distribution of the neuropathy that will support or refute the findings from the history and physical examination; they can confirm whether the neuropathic disorder is a mononeuropathy, multiple mononeuropathy (mononeuropathy multiplex), radiculopathy, plexopathy, or generalized polyneuropathy. Similarly, EDx evaluation can ascertain whether the process involves only sensory fibers, motor fibers, autonomic fibers, or a combination of these. Finally, the electrophysiologic data can help distinguish axonopathies from myelinopathies as well as axonal degeneration secondary to ganglionopathies from the more common length-dependent axonopathies.

TABLE 459-3 ELECTROPHYSIOLOGIC FEATURES: AXONAL DEGENERATION VERSUS SEGMENTAL DEMYELINATION

	Axonal Degeneration	Segmental Demyelination
Motor Nerve Conduction Studies		
CMAP amplitude	Decreased	Normal (except with CB or distal dispersion)
Distal latency	Normal	Prolonged
Conduction velocity	Normal	Slow
Conduction block	Absent	Present
Temporal dispersion	Absent	Present
F wave	Normal or absent	Prolonged or absent
H reflex	Normal or absent	Prolonged or absent
Sensory Nerve Conduction Studies		
SNAP amplitude	Decreased	Normal or decreased
Distal latency	Normal	Prolonged
Conduction velocity	Normal	Slow
Needle EMG		
Spontaneous activity		
Fibrillations	Present	Absent
Fasciculations	Present	Absent
Motor unit potentials		
Recruitment	Decreased	Decreased
Morphology	Long duration/ polyphasic	Normal

Abbreviations: CB, conduction block; CMAP, compound motor action potential; EMG, electromyography; SNAP, sensory nerve action potential.

NCS are most helpful in classifying a neuropathy as being due to axonal degeneration or segmental demyelination (Table 459-3). In general, low-amplitude potentials with relatively preserved distal latencies, conduction velocities, and late potentials, along with fibrillations on needle EMG, suggest an axonal neuropathy. On the other hand, slow conduction velocities, prolonged distal latencies and late potentials, relatively preserved amplitudes, and the absence of fibrillations on needle EMG imply a primary demyelinating neuropathy. The presence of nonuniform slowing of conduction velocity, conduction block, or temporal dispersion further suggests an acquired demyelinating neuropathy (e.g., GBS or CIDP) as opposed to a hereditary demyelinating neuropathy (e.g., CMT type 1).

Autonomic studies are used to assess small myelinated (A-delta) or unmyelinated (C) nerve fiber involvement. Such testing includes heart rate response to deep breathing, heart rate, and blood pressure response to both the Valsalva maneuver and tilt-table testing and quantitative sudomotor axon reflex testing (Chap. 454). These studies are particularly useful in patients who have pure small-fiber neuropathy or autonomic neuropathy in which routine NCS are normal.

OTHER IMPORTANT LABORATORY INFORMATION

In patients with generalized symmetric peripheral neuropathy, a standard laboratory evaluation should include a complete blood count, basic chemistries including serum electrolytes and tests of renal and hepatic function, fasting blood glucose (FBS), HbA_{1c}, urinalysis, thyroid function tests, B₁₂, folate, erythrocyte sedimentation rate (ESR), rheumatoid factor, antinuclear antibodies (ANA), serum protein electrophoresis (SPEP) and immunoelectrophoresis or immunofixation, and urine for Bence Jones protein. Quantification of the concentration of serum free light chains and the kappa/lambda ratio is more sensitive than SPEP, immunoelectrophoresis, or immunofixation in looking for a monoclonal gammopathy and therefore should be done if one suspects amyloidosis. A skeletal survey should be performed in patients with acquired demyelinating neuropathies and M-spikes to look for osteosclerotic or lytic lesions. Patients with monoclonal gammopathy should also be referred to a hematologist for consideration of a bone marrow biopsy. An oral glucose tolerance test is indicated in patients

with painful sensory neuropathies even if FBS and HbA_{1c} are normal, as the test is abnormal in about one-third of such patients. In addition to the above tests, patients with a mononeuropathy multiplex pattern of involvement should have a vasculitis workup, including antineutrophil cytoplasmic antibodies (ANCA), cryoglobulins, hepatitis serology, Western blot for Lyme disease, HIV, and occasionally a cytomegalovirus (CMV) titer.

There are many autoantibody panels (various antiganglioside antibodies) marketed for screening routine neuropathy patients for a treatable condition. These autoantibodies have no proven clinical utility or added benefit beyond the information obtained from a complete clinical examination and detailed EDx. A heavy metal screen is also not necessary as a screening procedure, unless there is a history of possible exposure or suggestive features on examination (e.g., severe painful sensorimotor and autonomic neuropathy and alopecia—thallium; severe painful sensorimotor neuropathy with or without gastrointestinal [GI] disturbance and Mee's lines—arsenic; wrist or finger extensor weakness and anemia with basophilic stippling of red blood cells—lead).

In patients with suspected GBS or CIDP, a lumbar puncture is indicated to look for an elevated cerebral spinal fluid (CSF) protein. In idiopathic cases of GBS and CIDP, there should not be pleocytosis in the CSF. If cells are present, one should consider HIV infection, Lyme disease, sarcoidosis, or lymphomatous or leukemic infiltration of nerve roots. Some patients with GBS and CIDP have abnormal liver function tests. In these cases, it is important to also check for hepatitis B and C, HIV, CMV, and Epstein-Barr virus (EBV) infection. In patients with an axonal GBS (by EMG/NCS) or those with a suspicious coinciding history (e.g., unexplained abdominal pain, psychiatric illness, significant autonomic dysfunction), it is reasonable to screen for porphyria.

In patients with a severe sensory ataxia, a sensory ganglionopathy or neuronopathy should be considered. The most common causes of sensory ganglionopathies are Sjögren's syndrome and a paraneoplastic neuropathy. Neuropathy can be the initial manifestation of Sjögren's syndrome. Thus, one should always inquire about dry eyes and mouth in patients with sensory signs and symptoms. Further, some patients can manifest sicca complex without full-blown Sjögren's syndrome. Thus, patients with sensory ataxia should have a senile systemic amyloidosis (SSA) and single strand binding (SSB) in addition to the routine ANA. To work up a possible paraneoplastic sensory ganglionopathy, antineuronal nuclear antibodies (e.g., anti-Hu antibodies) should be obtained (Chap. 122). These antibodies are most commonly seen in patients with small-cell carcinoma of the lung but are seen also in breast, ovarian, lymphoma, and other cancers. Importantly, the paraneoplastic neuropathy can precede the detection of the cancer, and detection of these autoantibodies should lead to a search for malignancy.

NERVE BIOPSIES

Nerve biopsies are now rarely indicated for evaluation of neuropathies. The primary indication for nerve biopsy is suspicion for amyloid neuropathy or vasculitis. In most instances, the abnormalities present on biopsies do not help distinguish one form of peripheral neuropathy from another (beyond what is already apparent by clinical examination and the NCS). Nerve biopsies should only be done if the NCS are abnormal. The sural nerve is most commonly biopsied because it is a pure sensory nerve and biopsy will not result in loss of motor function. In suspected vasculitis, a combination biopsy of a superficial peroneal nerve (pure sensory) and the underlying peroneus brevis muscle obtained from a single small incision increases the diagnostic yield. Tissue can be analyzed by frozen section and paraffin section to assess the supporting structures for evidence of inflammation, vasculitis, or amyloid deposition. Semithin plastic sections, teased fiber preparations, and electron microscopy are used to assess the morphology of the nerve fibers and to distinguish axonopathies from myelinopathies.

SKIN BIOPSIES

Skin biopsies are sometimes used to diagnose a small-fiber neuropathy. Following a punch biopsy of the skin in the distal lower extremity, immunologic staining can be used to measure the density of small

unmyelinated fibers. The density of these nerve fibers is reduced in patients with small-fiber neuropathies in whom NCS and routine nerve biopsies are often normal. This technique may allow for an objective measurement in patients with mainly subjective symptoms. However, it adds little to what one already knows from the clinical examination and EDx.

SPECIFIC DISORDERS

HEREDITARY NEUROPATHIES

Charcot-Marie-Tooth (CMT) disease is the most common type of hereditary neuropathy. Rather than one disease, CMT is a syndrome of several genetically distinct disorders (Table 459-4). The various subtypes of CMT are classified according to the nerve conduction velocities and predominant pathology (e.g., demyelination or axonal degeneration), inheritance pattern (autosomal dominant, recessive, or X-linked), and the specific mutated genes. Type 1 CMT (or CMT1) refers to inherited demyelinating sensorimotor neuropathies, whereas the axonal sensory neuropathies are classified as CMT2. By definition, motor conduction velocities in the arms are slowed to less than 38 m/s in CMT1 and are greater than 38 m/s in CMT2. However, most cases of CMT1 actually have motor nerve conduction velocities (NCVs) between 20 and 25 m/s. CMT1 and CMT2 usually begin in childhood or early adult life; however, onset later in life can occur, particularly in CMT2. Both are associated with autosomal dominant inheritance, with a few exceptions. CMT3 is an autosomal dominant neuropathy that appears in infancy and is associated with severe demyelination or hypomyelination. CMT4 is an autosomal recessive neuropathy that typically begins in childhood or early adult life. There are no medical therapies for any of the CMTs, but physical and occupational therapy can be beneficial, as can bracing (e.g., ankle-foot orthotics for foot-drop) and other orthotic devices.

CMT1 CMT1 is the most common form of hereditary neuropathy, with the ratio of CMT1:CMT2 being approximately 2:1. Affected individuals usually present in the first to third decade of life with distal leg weakness (e.g., footdrop), although patients may remain asymptomatic even late in life. People with CMT generally do not complain of numbness or tingling, which can be helpful in distinguishing CMT from acquired forms of neuropathy in which sensory symptoms usually predominate. Although usually asymptomatic in this regard, reduced sensation to all modalities is apparent on examination. Muscle stretch reflexes are unobtainable or reduced throughout. There is often atrophy of the muscles below the knee (particularly the anterior compartment), leading to so-called inverted champagne bottle legs.

Motor NCVs are usually in the 20–25 m/s range. Nerve biopsies usually are not performed on patients suspected of having CMT1, because the diagnosis usually can be made by less invasive testing (e.g., NCS and genetic studies). However, when done, the biopsies reveal reduction of myelinated nerve fibers with a predilection for the loss of the large-diameter fibers and Schwann cell proliferation around thinly or demyelinated fibers, forming so-called onion bulbs.

CMT1A is the most common subtype of CMT1, representing 70% of cases, and is caused by a 1.5-megabase (Mb) duplication within chromosome 17p11.2-12 wherein the gene for peripheral myelin protein-22 (PMP-22) lies. This results in patients having three copies of the PMP-22 gene rather than two. This protein accounts for 2–5% of myelin protein and is expressed in compact portions of the peripheral myelin sheath. Approximately 20% of patients with CMT1 have CMT1B, which is caused by mutations in the myelin protein zero (MPZ). CMT1B is for the most part clinically, electrophysiologically, and histologically indistinguishable from CMT1A. MPZ is an integral myelin protein and accounts for more than half of the myelin protein in peripheral nerves. Other forms of CMT1 are much less common and again indistinguishable from one another clinically and electrophysiologically.

CMT2 CMT2 tends to present later in life compared to CMT1. Affected individuals usually become symptomatic in the second decade of life; some cases present earlier in childhood, whereas others remain asymptomatic into late adult life. Clinically, CMT2 is for the most part indistinguishable from CMT1. NCS are helpful in this regard;

in contrast to CMT1, the velocities are normal or only slightly slowed. The most common cause of CMT2 is a mutation in the gene for mitofusin 2 (MFN2), which accounts for one-third of CMT2 cases overall. MFN2 localizes to the outer mitochondrial membrane, where it regulates the mitochondrial network architecture by fusion of mitochondria. The other genes associated with CMT2 are much less common.

CMTDI In dominant-intermediate CMTs (CMTDIs), the NCVs are faster than usually seen in CMT1 (e.g., >38 m/s) but slower than in CMT2.

CMT3 CMT3 was originally described by Dejerine and Sottas as a hereditary demyelinating sensorimotor polyneuropathy presenting in infancy or early childhood. Affected children are severely weak. Motor NCVs are markedly slowed, typically 5–10 m/s or less. Most cases of CMT3 are caused by point mutations in the genes for PMP-22, MPZ, or ERG-2, which are also the genes responsible for CMT1.

CMT4 CMT4 is extremely rare and is characterized by a severe, childhood-onset sensorimotor polyneuropathy that is usually inherited in an autosomal recessive fashion. Electrophysiologic and histologic evaluations can show demyelinating or axonal features. CMT4 is genetically heterogenic (Table 459-4).

CMT1X CMT1X is an X-linked dominant disorder with clinical features similar to CMT1 and CMT2, except that the neuropathy is much more severe in men than in women. CMT1X accounts for approximately 10–15% of CMT overall. Men usually present in the first two decades of life with atrophy and weakness of the distal arms and legs, areflexia, pes cavus, and hammertoes. Obligate women carriers are frequently asymptomatic, but can develop signs and symptoms. Onset in women is usually after the second decade of life, and the neuropathy is milder in severity.

NCS reveal features of both demyelination and axonal degeneration that are more severe in men compared to women. In men, motor NCVs in the arms and legs are moderately slowed (in the low to mid 30-m/s range). About 50% of men with CMT1X have motor NCVs between 15 and 35 m/s with about 80% of these falling between 25 and 35 m/s (intermediate slowing). In contrast, about 80% of women with CMT1X have NCVs in the normal range and 20% have NCVs in the intermediate range. CMT1X is caused by mutations in the connexin 32 gene. Connexins are gap junction structural proteins that are important in cell-to-cell communication.

Hereditary Neuropathy with Liability to Pressure Palsies (HNPP) HNPP is an autosomal dominant disorder related to CMT1A. Although CMT1A is usually associated with a 1.5-Mb duplication in chromosome 17p11.2 that results in an extra copy of PMP-22 gene, HNPP is caused by inheritance of the chromosome with the corresponding 1.5-Mb deletion of this segment, and thus affected individuals have only one copy of the PMP-22 gene. Patients usually manifest in the second or third decade of life with painless numbness and weakness in the distribution of single peripheral nerves, although multiple mononeuropathies can occur. Symptomatic mononeuropathy or multiple mononeuropathies are often precipitated by trivial compression of nerve(s) as can occur with wearing a backpack, leaning on the elbows, or crossing one's legs for even a short period of time. These pressure-related mononeuropathies may take weeks or months to resolve. In addition, some affected individuals manifest with a progressive or relapsing, generalized and symmetric, sensorimotor peripheral neuropathy that resembles CMT.

Hereditary Neuralgic Amyotrophy (HNA) HNA is an autosomal dominant disorder characterized by recurrent attacks of pain, weakness, and sensory loss in the distribution of the brachial plexus often beginning in childhood. These attacks are similar to those seen with idiopathic brachial plexitis (see below). Attacks may occur in the postpartum period, following surgery, or at other times of stress. Most patients recover over several weeks or months. Slightly dysmorphic features, including hypotelorism, epicanthal folds, cleft palate, syndactyly, micrognathia, and facial asymmetry, are evident in some individuals. EDx demonstrate an axonal process. HNA is caused by mutations in septin 9 (*SEPT9*). Septins may be important in formation of the

TABLE 459-4 CLASSIFICATION OF CHARCOT-MARIE-TOOTH DISEASE AND RELATED NEUROPATHIES

Name	Inheritance	Gene Location	Gene Product
CMT1			
CMT1A	AD	17p11.2	PMP-22 (usually duplication of gene)
CMT1B	AD	1q21-23	MPZ
CMT1C	AD	16p13.1-p12.3	LITAF
CMT1D	AD	10q21.1-22.1	ERG2
CMT1E (with deafness)	AD	17p11.2	Point mutations in PMP 22 gene
CMT1F	AD	8p13-21	Neurofilament light chain
CMT1G	AD	14q32.33	INF2
CMT1X	X-linked dominant	Xq13	Connexin-32
HNPP	AD	17p11.2 1q21-23	PMP-22 MPZ
CMT dominant-intermediate (CMTD1)			
CMTD1A	AD	10q24.1-25.1	?
CMTD1B	AD	19p12-13.2	Dynamin 2
CMTD1C	AD	1p35	YARS
CMTD1D	AD	1q22	MPZ
CMT2			
CMT2A2 (allelic to HMSN VI with optic atrophy)	AD	1p36.2	MFN2
CMT2B	AD	3q13-q22	RAB7
CMT2B1 (allelic to LGMD 1B)	AR	1q21.2	Lamin A/C
CMT2B2	AR and AD	19q13	MED25 for AR Unknown for AD
CMT2C (with vocal cord and diaphragm paralysis)	AD	12q23-24	TRPV4
CMT2D (allelic to distal SMA5)	AD	7p14	Glycine tRNA synthetase
CMT2E (allelic to CMT 1F)	AD	8p21	Neurofilament light chain
CMT2F	AD	7q11-q21	Heat-shock 27-kDa protein-1
CMT2G	AD	12q23	Unknown
CMT2I (allelic to CMT1B)	AD	1q22	MPZ
CMT2J	AD	1q22	MPZ
CMT2H, CMT2K (allelic to CMT4A)	AD	8q13-q21	GDAP1
CMT2L (allelic to distal hereditary motor neuropathy type 2)	AD	12q24	Heat-shock protein 8
CMT2M	AD	16q22	Dynamin-2
CMT2N	AD	16q22.1	AARS
CMT2O	AD	14q32.31	DYNC1H1
CMT2P	AD	9q34.13	LRSAM1
CMT2P-Okinawa (HSMN2P)	AD	3q13-q14	TFG
CMT2X	X-linked	Xq22-24	PRPS1
CMT3			
(Dejerine-Sottas disease, congenital hypomyelinating neuropathy)	AD	17p11.2 1q21-23	PMP-22 MPZ
	AR	10q21.1-22.1	ERG2
	AR	19q13	Periaxon
CMT4			
CMT4A	AR	8q13-21.1	GDAP1
CMT4B1	AR	11q23	MTMR2
CMT4B2	AR	11p15	MTMR13
CMT4C	AR	5q23-33	SH3TC2
CMT4D (HMSN-Lom)	AR	8q24	NDRG1
CMT4E (congenital hypomyelinating neuropathy)	AR	Multiple	Includes PMP22, MPZ, and ERG-2
CMT4F	AR	19q13.1-13.3	Periaxon
CMT4G	AR	10q23.2	HKI
CMT4H	AR	12q12-q13	Frabin
CMT4J	AR	6q21	FIG4
HNA	AD	17q24	SEPT9
HSAN1A	AD	9q22	SPTLC1
HSAN1B	AD	3q21	RAB7
HSAN1C	AD	14q24.3	SPTLC2
HSAN1D	AD	14q21.3	ATL1
HSAN1E	AD	19p13.2	DNMT1

(Continued)

TABLE 459-4 CLASSIFICATION OF CHARCOT-MARIE-TOOTH DISEASE AND RELATED NEUROPATHIES (CONTINUED)

Name	Inheritance	Gene Location	Gene Product
HSAN2A	AR	12p13.33	PRKWNK1
HSAN2B	AR	5p15.1	FAM134B
HSAN2C	AR	12q13.13	KIF1A
HSAN3	AR	9q21	IKAP
HSAN4	AR	3q	trkA/NGF receptor
HSAN5	AD or AR	1p11.2-p13.2	NGFb
HSAN6	AR	6p12.1	Dystonin

Abbreviations: AARS, alanyl-tRNA synthetase; AD, autosomal dominant; AR, autosomal recessive; ATL, atlastin; CMT, Charcot-Marie-Tooth; DNMT1, DNA methyltransferase 1; DYNC1H1, cytoplasmic dynein 1 heavy chain 1; ERG2, early growth response-2 protein; FAM134B, family with sequence similarity 134, member B; FIG4, FDG1-related F actin-binding protein; GDAP1, ganglioside-induced differentiation-associated protein-1; HK1, hexokinase 1; HMSN-P, hereditary motor and sensory neuropathy proximal; HNA, hereditary neuralgic amyotrophy; HNPP, hereditary neuropathy with liability to pressure palsies; HSAN, hereditary sensory and autonomic neuropathy; IFN2, inverted formin-2; IKAP, β kinase complex-associated protein; LGMD, limb girdle muscular dystrophy; LITAF, lipopolysaccharide-induced tumor necrosis factor α factor; LRSAM1, E3 ubiquitin-protein ligase; MED25, mediator 25; MFN2, mitochondrial fusion protein mitofusin 2 gene; MPZ, myelin protein zero protein; MTMR2, myotubularin-related protein-2; NDRG1, N-myc downstream regulated 1; PMP-22, peripheral myelin protein-22; PRKWNK1, protein kinase, lysine deficient 1; PRPS1, phosphoribosylpyrophosphate synthetase 1; RAB7, Ras-related protein 7; SEPT9, Septin 9; SH3TC2, SH3 domain and tetratricopeptide repeats 2; SMA, spinal muscular atrophy; SPTLC, serine palmitoyltransferase long-chain base; TFG, TRK-fused gene; TrkA/NGF, tyrosine kinase A/nerve growth factor; tRNA, transfer ribonucleic acid; TRPV4, transient receptor potential cation channel, subfamily V, member 4; WNK1, WNK lysine deficient; YARS, tyrosyl-tRNA synthetase.

Source: Modified from AA Amato, J Russell: *Neuromuscular Disease*. New York, McGraw-Hill, 2008.

neuronal cytoskeleton and have a role in cell division, but the mechanism of causing HNA is unclear.

Hereditary Sensory and Autonomic Neuropathy (HSAN) The HSANs are a very rare group of hereditary neuropathies in which sensory and autonomic dysfunction predominates over muscle weakness, unlike CMT, in which motor findings are most prominent (Table 459-4). Nevertheless, affected individuals can develop motor weakness and there can be overlap with CMT. There are no medical therapies available to treat these neuropathies, other than prevention and treatment of mutilating skin and bone lesions.

Of the HSANs, only HSAN1 typically presents in adults. HSAN1 is the most common of the HSANs and is inherited in an autosomal dominant fashion. Affected individuals with HSAN1 usually manifest in the second through fourth decades of life. HSAN1 is associated with the degeneration of small myelinated and unmyelinated nerve fibers leading to severe loss of pain and temperature sensation, deep dermal ulcerations, recurrent osteomyelitis, Charcot joints, bone loss, gross foot and hand deformities, and amputated digits. Although most people with HSAN1 do not complain of numbness, they often describe burning, aching, or lancinating pains. Autonomic neuropathy is not a prominent feature, but bladder dysfunction and reduced sweating in the feet may occur. HSAN1A, which is most common, is caused by mutations in the serine palmitoyltransferase long-chain base 1 (*SPTLC1*) gene.

OTHER HEREDITARY NEUROPATHIES (TABLE 459-5)

FABRY'S DISEASE

Fabry's disease (angiokeratoma corporis diffusum) is an X-linked dominant disorder. Although men are more commonly and severely affected, women can also show severe signs of the disease. Angiokeratomas are reddish-purple maculopapular lesions that are usually found around the umbilicus, scrotum, inguinal region, and perineum. Burning or lancinating pain in the hands and feet often develops in males in late childhood or early adult life. However, the neuropathy is usually overshadowed by complications arising from the associated premature atherosclerosis (e.g., hypertension, renal failure, cardiac disease, and stroke) that often lead to death by the fifth decade of life. Some patients also manifest primarily with a dilated cardiomyopathy.

Fabry's disease is caused by mutations in the α -galactosidase gene that leads to the accumulation of ceramide trihexoside in nerves and blood vessels. A decrease in α -galactosidase activity is evident in leukocytes and cultured fibroblasts. Glycolipid granules may be appreciated in ganglion cells of the peripheral and sympathetic nervous systems and in perineurial cells. Enzyme replacement therapy with α -galactosidase B can improve the neuropathy if patients are treated early, before irreversible nerve fiber loss.

ADRENOLEUKODYSTROPHY/ADRENOMYELONEUROPATHY

Adrenoleukodystrophy (ALD) and adrenomyeloneuropathy (AMN) are allelic X-linked dominant disorders caused by mutations in the

peroxisomal transmembrane adenosine triphosphate-binding cassette (ABC) transporter gene. Patients with ALD manifest with central nervous system (CNS) abnormalities. However, 30% with mutations in this gene present with the AMN phenotype that typically manifests in the third to fifth decade of life with mild to moderate peripheral neuropathy combined with progressive spastic paraplegia (**Chap. 456**). Rare patients present with an adult-onset spinocerebellar ataxia or only with adrenal insufficiency.

EDx is suggestive of a primary axonopathy with secondary demyelination. Nerve biopsies demonstrate a loss of myelinated and unmyelinated nerve fibers with lamellar inclusions in the cytoplasm of Schwann cells. Very long chain fatty acid (VLCFA) levels (C24, C25, and C26) are increased in the urine. Laboratory evidence of adrenal insufficiency is evident in approximately two-thirds of patients. The diagnosis can be confirmed by genetic testing.

Adrenal insufficiency is managed by replacement therapy; however, there is no proven effective therapy for the neurologic manifestations

TABLE 459-5 RARE HEREDITARY NEUROPATHIES

Hereditary Disorders of Lipid Metabolism
Metachromatic leukodystrophy
Krabbe's disease (globoid cell leukodystrophy)
Fabry's disease
Adrenoleukodystrophy/adrenomyeloneuropathy
Refsum's disease
Tangier disease
Cerebrotendinous xanthomatosis
Hereditary Ataxias with Neuropathy
Friedreich's ataxia
Vitamin E deficiency
Spinocerebellar ataxia
Abetalipoproteinemia (Bassen-Kornzweig disease)
Disorders of Defective DNA Repair
Ataxia-telangiectasia
Cockayne's syndrome
Giant Axonal Neuropathy
Porphyria
Acute intermittent porphyria (AIP)
Hereditary coproporphyrin (HCP)
Variante porphyria (VP)
Familial Amyloid Polyneuropathy (FAP)
Transthyretin-related
Gelsolin-related
Apolipoprotein A1-related

of ALD/AMN. Diets low in VLCFAs and supplemented with Lorenzo's oil (erucic and oleic acids) reduce the levels of VLCFAs and increase the levels of C22 in serum, fibroblasts, and liver; however, several large, open-label trials of Lorenzo's oil failed to demonstrate efficacy.

REFSUM'S DISEASE

Refsum's disease can manifest in infancy to early adulthood with the classic tetrad of (1) peripheral neuropathy, (2) retinitis pigmentosa, (3) cerebellar ataxia, and (4) elevated CSF protein concentration. Most affected individuals develop progressive distal sensory loss and weakness in the legs leading to footdrop by their 20s. Subsequently, the proximal leg and arm muscles may become weak. Patients may also develop sensorineural hearing loss, cardiac conduction abnormalities, ichthyosis, and anosmia.

Serum phytanic acid levels are elevated. Sensory and motor NCS reveal reduced amplitudes, prolonged latencies, and slowed conduction velocities. Nerve biopsy demonstrates a loss of myelinated nerve fibers, with remaining axons often thinly myelinated and associated with onion bulb formation.

Refsum disease is genetically heterogeneous but autosomal recessive in nature. Classical Refsum disease with childhood or early adult onset is caused by mutations in the gene that encodes for phytanoyl-CoA α -hydroxylase (*PAHX*). Less commonly, mutations in the gene encoding peroxin 7 receptor protein (*PRX7*) are responsible. These mutations lead to the accumulation of phytanic acid in the central and peripheral nervous systems. Refsum's disease is treated by removing phytanic precursors (phytols: fish oils, dairy products, and ruminant fats) from the diet.

TANGIER DISEASE

Tangier disease is a rare autosomal recessive disorder that can present as (1) asymmetric multiple mononeuropathies, (2) a slowly progressive symmetric polyneuropathy predominantly in the legs, or (3) a pseudo-syringomyelia pattern with dissociated sensory loss (i.e., abnormal pain/temperature perception but preserved position/vibration in the arms [Chap. 456]). The tonsils may appear swollen and yellowish-orange in color, and there may also be splenomegaly and lymphadenopathy.

Tangier disease is caused by mutations in the ATP-binding cassette transporter 1 (*ABC1*) gene, which leads to markedly reduced levels of high-density lipoprotein (HDL) cholesterol levels, whereas triacylglycerol levels are increased. Nerve biopsies reveal axonal degeneration with demyelination and remyelination. Electron microscopy demonstrates abnormal accumulation of lipid in Schwann cells, particularly those encompassing unmyelinated and small myelinated nerves. There is no specific treatment.

PORPHYRIA

Porphyria is a group of inherited disorders caused by defects in heme biosynthesis (Chap. 430). Three forms of porphyria are associated with peripheral neuropathy: acute intermittent porphyria (AIP), hereditary coproporphyrin (HCP), and variegate porphyria (VP). The acute neurologic manifestations are similar in each, with the exception that a photosensitive rash is seen with HCP and VP but not in AIP. Attacks of porphyria can be precipitated by certain drugs (usually those metabolized by the P450 system), hormonal changes (e.g., pregnancy, menstrual cycle), and dietary restrictions.

An acute attack of porphyria may begin with sharp abdominal pain. Subsequently, patients may develop agitation, hallucinations, or seizures. Several days later, back and extremity pain followed by weakness ensues, mimicking GBS. Weakness can involve the arms or the legs and can be asymmetric, proximal, or distal in distribution, as well as affecting the face and bulbar musculature. Dysautonomia and signs of sympathetic overactivity are common (e.g., pupillary dilation, tachycardia, and hypertension). Constipation, urinary retention, and incontinence can also be seen.

The CSF protein is typically normal or mildly elevated. Liver function tests and hematologic parameters are usually normal. Some patients are hyponatremic due to inappropriate secretion of antidiuretic hormone

(Chap. 401e). The urine may appear brownish in color secondary to the high concentration of porphyrin metabolites. Accumulation of intermediary precursors of heme (i.e., δ -aminolevulinic acid, porphobilinogen, uroporphobilinogen, coproporphyrinogen, and protoporphyrinogen) is found in urine. Specific enzyme activities can also be measured in erythrocytes and leukocytes. The primary abnormalities on EDx are marked reductions in compound motor action potential (CMAP) amplitudes and signs of active axonal degeneration on needle EMG.

The porphyrias are inherited in an autosomal dominant fashion. AIP is associated with porphobilinogen deaminase deficiency, HCP is caused by defects in coproporphyrin oxidase, and VP is associated with protoporphyrinogen oxidase deficiency. The pathogenesis of the neuropathy is not completely understood. Treatment with glucose and hematin may reduce the accumulation of heme precursors. Intravenous glucose is started at a rate of 10–20 g/h. If there is no improvement within 24 h, intravenous hematin 2–5 mg/kg per day for 3–14 days should be given.

FAMILIAL AMYLOID POLYNEUROPATHY

Familial amyloid polyneuropathy (FAP) is phenotypically and genetically heterogeneous and is caused by mutations in the genes for transthyretin (TTR), apolipoprotein A1, or gelsolin (Chap. 137). The majority of patients with FAP have mutations in the TTR gene. Amyloid deposition may be evident in abdominal fat pad, rectal, or nerve biopsies. The clinical features, histopathology, and EDx reveal abnormalities consistent with a generalized or multifocal, predominantly axonal but occasionally demyelinating, sensorimotor polyneuropathy.

Patients with TTR-related FAP usually develop insidious onset of numbness and painful paresthesias in the distal lower limbs in the third to fourth decade of life, although some patients develop the disorder later in life. Carpal tunnel syndrome (CTS) is common. Autonomic involvement can be severe, leading to postural hypotension, constipation or persistent diarrhea, erectile dysfunction, and impaired sweating. Amyloid deposition also occurs in the heart, kidneys, liver, and corneas. Patients usually die 10–15 years after the onset of symptoms from cardiac failure or complications from malnutrition. Because the liver produces much of the body's TTR, liver transplantation has been used to treat FAP related to TTR mutations. Serum TTR levels decrease after transplantation, and improvement in clinical and EDx features has been reported.

Patients with apolipoprotein A1-related FAP (Van Allen type) usually present in the fourth decade with numbness and painful dysesthesias in the distal limbs. Gradually, the symptoms progress, leading to proximal and distal weakness and atrophy. Although autonomic neuropathy is not severe, some patients develop diarrhea, constipation, or gastroparesis. Most patients die from systemic complications of amyloidosis (e.g., renal failure) 12–15 years after the onset of the neuropathy.

Gelsolin-related amyloidosis (Finnish type) is characterized by the combination of lattice corneal dystrophy and multiple cranial neuropathies that usually begin in the third decade of life. Over time, a mild generalized sensorimotor polyneuropathy develops. Autonomic dysfunction does not occur.

ACQUIRED NEUROPATHIES

PRIMARY OR AL AMYLOIDOSIS (SEE CHAP. 137)

Besides FAP, amyloidosis can also be acquired. In primary or AL amyloidosis, the abnormal protein deposition is composed of immunoglobulin light chains. AL amyloidosis occurs in the setting of multiple myeloma, Waldenström's macroglobulinemia, lymphoma, other plasmacytomas, or lymphoproliferative disorders, or without any other identifiable disease.

Approximately 30% of patients with AL primary amyloidosis present with a polyneuropathy, most typically painful dysesthesias and burning sensations in the feet. However, the trunk can be involved, and some patients manifest with a mononeuropathy multiplex pattern. CTS occurs in 25% of patients and may be the initial manifestation. The neuropathy is slowly progressive, and eventually weakness

develops along with large-fiber sensory loss. Most patients develop autonomic involvement with postural hypertension, syncope, bowel and bladder incontinence, constipation, impotence, and impaired sweating. Patients generally die from their systemic illness (renal failure, cardiac disease).

The monoclonal protein may be composed of IgG, IgA, IgM, or only free light chain. Lambda (λ) is more common than κ light chain (>2:1) in AL amyloidosis. The CSF protein is often increased (with normal cell count), and thus the neuropathy may be mistaken for CIDP (Chap. 460). Nerve biopsies reveal axonal degeneration and amyloid deposition in either a globular or diffuse pattern infiltrating the perineurial, epineurial, and endoneurial connected tissue and in blood vessel walls.

The median survival of patients with primary amyloidosis is less than 2 years, with death usually from progressive congestive heart failure or renal failure. Chemotherapy with melphalan, prednisone, and colchicine, to reduce the concentration of monoclonal proteins, and autologous stem cell transplantation may prolong survival, but whether the neuropathy improves is controversial.

DIABETIC NEUROPATHY

Diabetes mellitus (DM) is the most common cause of peripheral neuropathy in developed countries. DM is associated with several types of polyneuropathy: distal symmetric sensory or sensorimotor polyneuropathy, autonomic neuropathy, diabetic neuropathic cachexia, polyradiculoneuropathies, cranial neuropathies, and other mononeuropathies. Risk factors for the development of neuropathy include long-standing, poorly controlled DM and the presence of retinopathy and nephropathy.

Diabetic Distal Symmetric Sensory and Sensorimotor Polyneuropathy (DSPN)

DSPN is the most common form of diabetic neuropathy and manifests as sensory loss beginning in the toes that gradually progresses over time up the legs and into the fingers and arms. When severe, a patient may develop sensory loss in the trunk (chest and abdomen), initially in the midline anteriorly and later extending laterally. Tingling, burning, deep aching pains may also be apparent. NCS usually show reduced amplitudes and mild to moderate slowing of conduction velocities (CVs). Nerve biopsy reveals axonal degeneration, endothelial hyperplasia,

and, occasionally, perivascular inflammation. Tight control of glucose can reduce the risk of developing neuropathy or improve the underlying neuropathy. A variety of medications have been used with variable success to treat painful symptoms associated with DSPN, including antiepileptic medications, antidepressants, sodium channel blockers, and other analgesics (Table 459-6).

Diabetic Autonomic Neuropathy Autonomic neuropathy is typically seen in combination with DSPN. The autonomic neuropathy can manifest as abnormal sweating, dysfunctional thermoregulation, dry eyes and mouth, pupillary abnormalities, cardiac arrhythmias, postural hypotension, GI abnormalities (e.g., gastroparesis, postprandial bloating, chronic diarrhea or constipation), and genitourinary dysfunction (e.g., impotence, retrograde ejaculation, incontinence). Tests of autonomic function are generally abnormal, including sympathetic skin responses and quantitative sudomotor axon reflex testing. Sensory and motor NCS generally demonstrate features described above with DSPN.

Diabetic Radiculoplexus Neuropathy (Diabetic Amyotrophy or Bruns-Garland Syndrome)

Diabetic radiculoplexus neuropathy is the presenting manifestation of DM in approximately one-third of patients. Typically, patients present with severe pain in the low back, hip, and thigh in one leg. Rarely, the diabetic polyradiculoneuropathy begins in both legs at the same time. Atrophy and weakness of proximal and distal muscles in the affected leg become apparent within a few days or weeks. The neuropathy is often accompanied or heralded by severe weight loss. Weakness usually progresses over several weeks or months, but can continue to progress for 18 months or more. Subsequently, there is slow recovery but many are left with residual weakness, sensory loss, and pain. In contrast to the more typical lumbosacral radiculoplexus neuropathy, some patients develop thoracic radiculopathy or, even less commonly, a cervical polyradiculoneuropathy. CSF protein is usually elevated, while the cell count is normal. ESR is often increased. EDx reveals evidence of active denervation in affected proximal and distal muscles in the affected limbs and in paraspinous muscles. Nerve biopsies may demonstrate axonal degeneration along with perivascular inflammation. Patients with severe pain are sometimes treated in the acute period with glucocorticoids, although a randomized controlled trial

TABLE 459-6 TREATMENT OF PAINFUL SENSORY NEUROPATHIES

Therapy	Route	Dose	Side Effects
First-Line			
Lidoderm 5% patch	Apply to painful area	Up to 3 patches qd	Skin irritation
Tricyclic antidepressants (e.g., amitriptyline, nortriptyline)	PO	10–100 mg qhs	Cognitive changes, sedation, dry eyes and mouth, urinary retention, constipation
Gabapentin	PO	300–1200 mg tid	Cognitive changes, sedation, peripheral edema
Pregabalin	PO	50–100 mg tid	Cognitive changes, sedation, peripheral edema
Duloxetine	PO	30–60 mg qd	Cognitive changes, sedation, dry eyes, diaphoresis, nausea, diarrhea, constipation
Second-Line			
Carbamazepine	PO	200–400 mg q 6–8 h	Cognitive changes, dizziness, leukopenia, liver dysfunction
Phenytoin	PO	200–400 mg qhs	Cognitive changes, dizziness, liver dysfunction
Venlafaxine	PO	37.5–150 mg/d	Asthenia, sweating, nausea, constipation, anorexia, vomiting, somnolence, dry mouth, dizziness, nervousness, anxiety, tremor, and blurred vision as well as abnormal ejaculation/orgasm and impotence
Tramadol	PO	50 mg qid	Cognitive changes, gastrointestinal upset
Third-Line			
Mexiletine	PO	200–300 mg tid	Arrhythmias
Other Agents			
EMLA cream 2.5% lidocaine 2.5% prilocaine	Apply cutaneously	qid	Local erythema
Capsaicin 0.025–0.075% cream	Apply cutaneously	qid	Painful burning skin

Source: Modified from AA Amato, J Russell: *Neuromuscular Disease*. New York, McGraw-Hill, 2008.

has yet to be performed, and the natural history of this neuropathy is gradual improvement.

Diabetic Mononeuropathies or Multiple Mononeuropathies The most common mononeuropathies are median neuropathy at the wrist and ulnar neuropathy at the elbow, but peroneal neuropathy at the fibular head, and sciatic, lateral femoral, cutaneous, or cranial neuropathies also occur. In regard to cranial mononeuropathies, seventh nerve palsies are relatively common but may have other, nondiabetic etiologies. In diabetics, a third nerve palsy is most common, followed by sixth nerve, and, less frequently, fourth nerve palsies. Diabetic third nerve palsies are characteristically pupil-sparing (**Chap. 39**).

HYPOTHYROIDISM

Hypothyroidism is more commonly associated with a proximal myopathy, but some patients develop a neuropathy, most typically CTS. Rarely, a generalized sensory polyneuropathy characterized by painful paresthesias and numbness in both the legs and hands can occur. Treatment is correction of the hypothyroidism.

SJÖGREN'S SYNDROME

Sjögren's syndrome, characterized by the sicca complex of xerophthalmia, xerostomia, and dryness of other mucous membranes, can be complicated by neuropathy (**Chap. 383**). Most common is a length-dependent axonal sensorimotor neuropathy characterized mainly by sensory loss in the distal extremities. A pure small-fiber neuropathy or a cranial neuropathy, particularly involving the trigeminal nerve, can also be seen. Sjögren's syndrome is also associated with sensory neuronopathy/ganglionopathy. Patients with sensory ganglionopathies develop progressive numbness and tingling of the limbs, trunk, and face in a non-length-dependent manner such that symptoms can involve the face or arms more than the legs. The onset can be acute or insidious. Sensory examination demonstrates severe vibratory and proprioceptive loss leading to sensory ataxia.

Patients with neuropathy due to Sjögren's syndrome may have ANAs, SS-A/Ro, and SS-B/La antibodies in the serum, but most do not. NCS demonstrate reduced amplitudes of sensory studies in the affected limbs. Nerve biopsy demonstrates axonal degeneration. Nonspecific perivascular inflammation may be present, but only rarely is there necrotizing vasculitis. There is no specific treatment for neuropathies related to Sjögren's syndrome. When vasculitis is suspected, immunosuppressive agents may be beneficial. Occasionally, the sensory neuronopathy/ganglionopathy stabilizes or improves with immunotherapy, such as IVIg.

RHEUMATOID ARTHRITIS

Peripheral neuropathy occurs in at least 50% of patients with rheumatoid arthritis (RA) and may be vasculitic in nature (**Chap. 380**). Vasculitic neuropathy can present with a mononeuropathy multiplex, a generalized symmetric pattern of involvement, or a combination of these patterns (**Chap. 385**). Neuropathies may also be due to drugs used to treat the RA (e.g., tumor necrosis blockers, leflunomide). Nerve biopsy often reveals thickening of the epineurial and endoneurial blood vessels as well as perivascular inflammation or vasculitis, with transmural inflammatory cell infiltration and fibrinoid necrosis of vessel walls. The neuropathy often is responsive to immunomodulating therapies.

SYSTEMIC LUPUS ERYTHEMATOSUS (SLE)

Between 2 and 27% of individuals with SLE develop a peripheral neuropathy (**Chap. 378**). Affected patients typically present with a slowly progressive sensory loss beginning in the feet. Some patients develop burning pain and paresthesias with normal reflexes, and NCS suggest a pure small-fiber neuropathy. Less common are multiple mononeuropathies presumably secondary to necrotizing vasculitis. Rarely, a generalized sensorimotor polyneuropathy meeting clinical, laboratory, electrophysiologic, and histologic criteria for either GBS or CIDP may occur. Immunosuppressive therapy is beneficial in SLE patients with neuropathy due to vasculitis. Immunosuppressive agents are less likely

to be effective in patients with a generalized sensory or sensorimotor polyneuropathy without evidence of vasculitis. Patients with a GBS or CIDP-like neuropathy should be treated accordingly (**Chap. 385**).

SYSTEMIC SCLEROSIS (SCLERODERMA)

A distal symmetric, mainly sensory, polyneuropathy complicates 5–67% of scleroderma cases (**Chap. 382**). Cranial mononeuropathies can also develop, most commonly of the trigeminal nerve, producing numbness and dysesthesias in the face. Multiple mononeuropathies also occur. The EDx and histologic features of nerve biopsy are those of an axonal sensory greater than motor polyneuropathy.

MIXED CONNECTIVE TISSUE DISEASE (MCTD)

A mild distal axonal sensorimotor polyneuropathy occurs in approximately 10% of patients with MCTD.

SARCOIDOSIS

The peripheral or central nervous system is involved in about 5% of patients with sarcoidosis (**Chap. 390**). The most common cranial nerve involved is the seventh nerve, which can be affected bilaterally. Some patients develop radiculopathy or polyradiculopathy. With a generalized root involvement, the clinical presentation can mimic GBS or CIDP. Patients can also present with multiple mononeuropathies or a generalized, slowly progressive, sensory greater than motor polyneuropathy. Some have features of a pure small-fiber neuropathy. EDx reveals an axonal neuropathy. Nerve biopsy can reveal noncaseating granulomas infiltrating the endoneurium, perineurium, and epineurium along with lymphocytic necrotizing angiitis. Neurosarcoidosis may respond to treatment with glucocorticoids or other immunosuppressive agents.

HYPEREOSINOPHILIC SYNDROME

Hypereosinophilic syndrome is characterized by eosinophilia associated with various skin, cardiac, hematologic, and neurologic abnormalities. A generalized peripheral neuropathy or a mononeuropathy multiplex occurs in 6–14% of patients.

CELIAC DISEASE (GLUTEN-INDUCED ENTEROPATHY OR NONTROPICAL SPRUE)

Neurologic complications, particularly ataxia and peripheral neuropathy, are estimated to occur in 10% of patients with celiac disease (**Chap. 349**). A generalized sensorimotor polyneuropathy, pure motor neuropathy, multiple mononeuropathies, autonomic neuropathy, small-fiber neuropathy, and neuromyotonia have all been reported in association with celiac disease or antigliadin/antiendomysial antibodies. Nerve biopsy may reveal a loss of large myelinated fibers. The neuropathy may be secondary to malabsorption of vitamins B₁₂ and E. However, some patients have no appreciable vitamin deficiencies. The pathogenic basis for the neuropathy in these patients is unclear but may be autoimmune in etiology. The neuropathy does not appear to respond to a gluten-free diet. In patients with vitamin B₁₂ or vitamin E deficiency, replacement therapy may improve or stabilize the neuropathy.

INFLAMMATORY BOWEL DISEASE

Ulcerative colitis and Crohn's disease may be complicated by GBS, CIDP, generalized axonal sensory or sensorimotor polyneuropathy, small-fiber neuropathy, or mononeuropathy (**Chap. 351**). These neuropathies may be autoimmune, nutritional (e.g., vitamin B₁₂ deficiency), treatment related (e.g., metronidazole), or idiopathic in nature. An acute neuropathy with demyelination resembling GBS, CIDP, or multifocal motor neuropathy may occur in patients treated with tumor necrosis factor α blockers.

UREMIC NEUROPATHY

Approximately 60% of patients with renal failure develop a polyneuropathy characterized by length-dependent numbness, tingling, allodynia, and mild distal weakness. Rarely, a rapidly progressive weakness and sensory loss very similar to GBS can occur that improves with an increase in the intensity of renal dialysis or with transplantation. Mononeuropathies can also occur, the most common of which is CTS.

Ischemic monomelic neuropathy (see below) can complicate arteriovenous shunts created in the arm for dialysis. EDx in uremic patients reveals features of a length-dependent, primarily axonal, sensorimotor polyneuropathy. Sural nerve biopsies demonstrate a loss of nerve fibers (particularly large myelinated nerve fibers), active axonal degeneration, and segmental and paranodal demyelination. The sensorimotor polyneuropathy can be stabilized by hemodialysis and improved with successful renal transplantation.

CHRONIC LIVER DISEASE

A generalized sensorimotor neuropathy characterized by numbness, tingling, and minor weakness in the distal aspects of primarily the lower limbs commonly occurs in patients with chronic liver failure. EDx studies are consistent with a sensory greater than motor axonopathy. Sural nerve biopsy reveals both segmental demyelination and axonal loss. It is not known if hepatic failure in isolation can cause peripheral neuropathy, as the majority of patients have liver disease secondary to other disorders, such as alcoholism or viral hepatitis, which can also cause neuropathy.

CRITICAL ILLNESS POLYNEUROPATHY

The most common causes of acute generalized weakness leading to admission to a medical intensive care unit (ICU) are GBS and myasthenia gravis (Chap. 461). However, weakness developing in critically ill patients while in the ICU is usually caused by critical illness polyneuropathy (CIP) or critical illness myopathy (CIM) or, much less commonly, by prolonged neuromuscular blockade. From a clinical and EDx standpoint, it can be quite difficult to distinguish these disorders. Most specialists suggest that CIM is more common. Both CIM and CIP develop as a complication of sepsis and multiple organ failure. They usually present as an inability to wean a patient from a ventilator. A coexisting encephalopathy may limit the neurologic exam, in particular the sensory examination. Muscle stretch reflexes are absent or reduced.

Serum creatine kinase (CK) is usually normal; an elevated serum CK would point to CIM as opposed to CIP. NCS reveal absent or markedly reduced amplitudes of motor and sensory studies in CIP, whereas sensory studies are relatively preserved in CIM. Needle EMG usually reveals profuse positive sharp waves and fibrillation potentials, and it is not unusual in patients with severe weakness to be unable to recruit motor unit action potentials. The pathogenic basis of CIP is not known. Perhaps circulating toxins and metabolic abnormalities associated with sepsis and multiorgan failure impair axonal transport or mitochondrial function, leading to axonal degeneration.

LEPROSY (HANSEN'S DISEASE)



Leprosy, caused by the acid-fast bacteria *Mycobacterium leprae*, is the most common cause of peripheral neuropathy in Southeast Asia, Africa, and South America (Chap. 203).

Clinical manifestations range from tuberculoid leprosy at one end to lepromatous leprosy at the other end of the spectrum, with borderline leprosy in between. Neuropathies are most common in patients with borderline leprosy. Superficial cutaneous nerves of the ears and distal limbs are commonly affected. Mononeuropathies, multiple mononeuropathies, or a slowly progressive symmetric sensorimotor polyneuropathy may develop. Sensory NCS are usually absent in the lower limb and are reduced in amplitude in the arms. Motor NCS may demonstrate reduced amplitudes in affected nerves but occasionally can reveal demyelinating features. Leprosy is usually diagnosed by skin lesion biopsy. Nerve biopsy can also be diagnostic, particularly when there are no apparent skin lesions. The tuberculoid form is characterized by granulomas, and bacilli are not seen. In contrast, with lepromatous leprosy, large numbers of infiltrating bacilli, T_H2 lymphocytes, and organism-laden, foamy macrophages with minimal granulomatous infiltration are evident. The bacilli are best appreciated using the Fite stain, where they can be seen as red-staining rods often in clusters free in the endoneurium, within macrophages, or within Schwann cells.

Patients are generally treated with multiple drugs: dapsone, rifampin, and clofazimine. Other medications that are used include thalidomide,

pefloxacin, ofloxacin, sparfloxacin, minocycline, and clarithromycin. Patients are generally treated for 2 years. Treatment is sometimes complicated by the so-called reversal reaction, particularly in borderline leprosy. The reversal reaction can occur at any time during treatment and develops because of a shift to the tuberculoid end of the spectrum, with an increase in cellular immunity during treatment. The cellular response is upregulated as evidenced by an increased release of tumor necrosis factor α , interferon γ , and interleukin 2, with new granuloma formation. This can result in an exacerbation of the rash and the neuropathy as well as in appearance of new lesions. High-dose glucocorticoids blunt this adverse reaction and may be used prophylactically at treatment onset in high-risk patients. Erythema nodosum leprosum (ENL) is also treated with glucocorticoids or thalidomide.

LYME DISEASE

Lyme disease is caused by infection with *Borrelia burgdorferi*, a spirochete usually transmitted by the deer tick *Ixodes dammini* (Chap. 210). Neurologic complications may develop during the second and third stages of infection. Facial neuropathy is most common and is bilateral in about half of cases, which is rare for idiopathic Bell's palsy. Involvement of nerves is frequently asymmetric. Some patients present with a polyradiculoneuropathy or multiple mononeuropathies. EDx is suggestive of a primary axonopathy. Nerve biopsies can reveal axonal degeneration with perivascular inflammation. Treatment is with antibiotics (Chap. 210).

DIPHTHERITIC NEUROPATHY

Diphtheria is caused by the bacteria *Corynebacterium diphtheriae* (Chap. 175). Infected individuals present with flulike symptoms of generalized myalgias, headache, fatigue, low-grade fever, and irritability within a week to 10 days of the exposure. About 20–70% of patients develop a peripheral neuropathy caused by a toxin released by the bacteria. Three to 4 weeks after infection, patients may note decreased sensation in their throat and begin to develop dysphagia, dysarthria, hoarseness, and blurred vision due to impaired accommodation. A generalized polyneuropathy may manifest 2 or 3 months following the initial infection, characterized by numbness, paresthesias, and weakness of the arms and legs and occasionally ventilatory failure. CSF protein can be elevated with or without lymphocytic pleocytosis. EDx suggests a diffuse axonal sensorimotor polyneuropathy. Antitoxin and antibiotics should be given within 48 h of symptom onset. Although early treatment reduces the incidence and severity of some complications (i.e., cardiomyopathy), it does not appear to alter the natural history of the associated peripheral neuropathy. The neuropathy usually resolves after several months.

HUMAN IMMUNODEFICIENCY VIRUS (HIV)

HIV infection can result in a variety of neurologic complications, including peripheral neuropathies (Chap. 226). Approximately 20% of HIV-infected individuals develop a neuropathy either as a direct result of the virus itself, other associated viral infections (e.g., CMV), or neurotoxicity secondary to antiviral medications (see below). The major presentations of peripheral neuropathy associated with HIV infection include (1) distal symmetric polyneuropathy, (2) inflammatory demyelinating polyneuropathy (including both GBS and CIDP), (3) multiple mononeuropathies (e.g., vasculitis, CMV-related), (4) polyradiculopathy (usually CMV-related), (5) autonomic neuropathy, and (6) sensory ganglionitis.

HIV-Related Distal Symmetric Polyneuropathy (DSP) DSP is the most common form of peripheral neuropathy associated with HIV infection and usually is seen in patients with AIDS. It is characterized by numbness and painful paresthesias involving the distal extremities. The pathogenic basis for DSP is unknown but is not due to actual infection of the peripheral nerves. The neuropathy may be immune mediated, perhaps caused by the release of cytokines from surrounding inflammatory cells. Vitamin B₁₂ deficiency may contribute in some instances but is not a major cause of most cases of DSP. Some antiretroviral agents (e.g., dideoxycytidine, dideoxyinosine, stavudine) are also neurotoxic and can cause a painful sensory neuropathy.

HIV-Related Inflammatory Demyelinating Polyradiculoneuropathy Both AIDP and CIDP can occur as a complication of HIV infection. AIDP usually develops at the time of seroconversion, whereas CIDP can occur any time in the course of the infection. Clinical and EDx features are indistinguishable from idiopathic AIDP or CIDP (discussed in [Chap. 460](#)). In addition to elevated protein levels, lymphocytic pleocytosis is evident in the CSF, a finding that helps distinguish this HIV-associated polyradiculoneuropathy from idiopathic AIDP/CIDP.

HIV-Related Progressive Polyradiculopathy An acute, progressive lumbosacral polyradiculoneuropathy usually secondary to CMV infection can develop in patients with AIDS. Patients present with severe radicular pain, numbness, and weakness in the legs, which is usually asymmetric. CSF is abnormal, demonstrating an increased protein along with reduced glucose concentration and notably a neutrophilic pleocytosis. EDx studies reveal features of active axonal degeneration. The polyradiculoneuropathy may improve with antiviral therapy.

HIV-Related Multiple Mononeuropathies Multiple mononeuropathies can also develop in patients with HIV infection, usually in the context of AIDS. Weakness, numbness, paresthesias, and pain occur in the distribution of affected nerves. Nerve biopsies can reveal axonal degeneration with necrotizing vasculitis or perivascular inflammation. Glucocorticoid treatment is indicated for vasculitis directly due to HIV infection.

HIV-Related Sensory Neuronopathy/Ganglionopathy Dorsal root ganglionitis is a very rare complication of HIV infection, and neuronopathy can be the presenting manifestation. Patients develop sensory ataxia similar to idiopathic sensory neuronopathy/ganglionopathy. NCS reveal reduced amplitudes or absence of sensory nerve action potentials (SNAPs).

HERPES VARICELLA-ZOSTER VIRUS

Peripheral neuropathy from herpes varicella-zoster (HVZ) infection results from reactivation of latent virus or from a primary infection ([Chap. 217](#)). Two-thirds of infections in adults are characterized by dermal zoster in which severe pain and paresthesias develop in a dermatomal region followed within a week or two by a vesicular rash in the same distribution. Weakness in muscles innervated by roots corresponding to the dermatomal distribution of skin lesions occurs in 5–30% of patients. Approximately 25% of affected patients have continued pain (postherpetic neuralgia [PHN]). A large clinical trial demonstrated that vaccination against zoster reduces the incidence of HVZ among vaccine recipients by 51% and reduces the incidence of PHN by 67%. Treatment of PHN is symptomatic (Table 459-6).

CYTOMEGALOVIRUS

CMV can cause an acute lumbosacral polyradiculopathy and multiple mononeuropathies in patients with HIV infection and in other immune deficiency conditions ([Chap. 219](#)).

EPSTEIN-BARR VIRUS

EBV infection has been associated with GBS, cranial neuropathies, mononeuropathy multiplex, brachial plexopathy, lumbosacral radiculoplexopathy, and sensory neuronopathies ([Chap. 218](#)).

HEPATITIS VIRUSES

Hepatitis B and C can cause multiple mononeuropathies related to vasculitis, AIDP, or CIDP ([Chap. 362](#)).

NEUROPATHIES ASSOCIATED WITH MALIGNANCY

Patients with malignancy can develop neuropathies due to (1) a direct effect of the cancer by invasion or compression of the nerves, (2) remote or paraneoplastic effect, (3) a toxic effect of treatment, or (4) as a consequence of immune compromise caused by immunosuppressive medications. The most common associated malignancy is lung cancer, but neuropathies also complicate carcinoma of the breast, ovaries, stomach, colon, rectum, and other organs, including the lymphoproliferative system.

PARANEOPLASTIC SENSORY NEURONOPATHY/GANGLIONOPATHY

Paraneoplastic encephalomyelitis/sensory neuronopathy (PEM/SN) usually complicates small-cell lung carcinoma ([Chap. 122](#)). Patients usually present with numbness and paresthesias in the distal extremities that are often asymmetric. The onset can be acute or insidiously progressive. Prominent loss of proprioception leads to sensory ataxia. Weakness can be present, usually secondary to an associated myelitis, motor neuronopathy, or concurrent Lambert-Eaton myasthenic syndrome (LEMS). Many patients also develop confusion, memory loss, depression, hallucinations or seizures, or cerebellar ataxia. Polyclonal antineuronal antibodies (IgG) directed against a 35- to 40-kDa protein or complex of proteins, the so-called Hu antigen, are found in the sera or CSF in the majority of patients with paraneoplastic PEM/SN. CSF may be normal or may demonstrate mild lymphocytic pleocytosis and elevated protein. PEM/SN is probably the result of antigenic similarity between proteins expressed in the tumor cells and neuronal cells, leading to an immune response directed against both cell types. Treatment of the underlying cancer generally does not affect the course of PEM/SN. However, occasional patients may improve following treatment of the tumor. Unfortunately, plasmapheresis, intravenous immunoglobulin, and immunosuppressive agents have not shown benefit.

NEUROPATHY SECONDARY TO TUMOR INFILTRATION

Malignant cells, in particular leukemia and lymphoma, can infiltrate cranial and peripheral nerves, leading to mononeuropathy, mononeuropathy multiplex, polyradiculopathy, plexopathy, or even a generalized symmetric distal or proximal and distal polyneuropathy. Neuropathy related to tumor infiltration is often painful; it can be the presenting manifestation of the cancer or the heralding symptom of a relapse. The neuropathy may improve with treatment of the underlying leukemia or lymphoma or with glucocorticoids.

NEUROPATHY AS A COMPLICATION OF BONE MARROW TRANSPLANTATION

Neuropathies may develop in patients who undergo bone marrow transplantation (BMT) because of the toxic effects of chemotherapy, radiation, infection, or an autoimmune response directed against the peripheral nerves. Peripheral neuropathy in BMT is often associated with graft-versus-host disease (GVHD). Chronic GVHD shares many features with a variety of autoimmune disorders, and it is possible that an immune-mediated response directed against peripheral nerves is responsible. Patients with chronic GVHD may develop cranial neuropathies, sensorimotor polyneuropathies, multiple mononeuropathies, and severe generalized peripheral neuropathies resembling AIDP or CIDP. The neuropathy may improve by increasing the intensity of immunosuppressive or immunomodulating therapy and resolution of the GVHD.

LYMPHOMA

Lymphomas may cause neuropathy by infiltration or direct compression of nerves or by a paraneoplastic process. The neuropathy can be purely sensory or motor, but most commonly is sensorimotor. The pattern of involvement may be symmetric, asymmetric, or multifocal, and the course may be acute, gradually progressive, or relapsing and remitting. EDx can be compatible with either an axonal or demyelinating process. CSF may reveal lymphocytic pleocytosis and an elevated protein. Nerve biopsy may demonstrate endoneurial inflammatory cells in both the infiltrative and the paraneoplastic etiologies. A monoclonal population of cells favors lymphomatous invasion. The neuropathy may respond to treatment of the underlying lymphoma or immunomodulating therapies.

MULTIPLE MYELOMA

Multiple myeloma (MM) usually presents in the fifth to seventh decade of life with fatigue, bone pain, anemia, and hypercalcemia ([Chap. 136](#)). Clinical and EDx features of neuropathy occur in as many as 40% of patients. The most common pattern is that of a distal, axonal, sensory, or sensorimotor polyneuropathy. Less frequently, a chronic demyelinating polyradiculoneuropathy may develop (see POEMS, [Chap. 460](#)). MM can be complicated by amyloid polyneuropathy and should be considered in patients with painful paresthesias, loss of pinprick and

temperature discrimination, and autonomic dysfunction (suggestive of a small-fiber neuropathy) and CTS. Expanding plasmacytomas can compress cranial nerves and spinal roots as well. A monoclonal protein, usually composed of γ or μ heavy chains or κ light chains, may be identified in the serum or urine. EDx usually shows reduced amplitudes with normal or only mildly abnormal distal latencies and conduction velocities. A superimposed median neuropathy at the wrist is common. Abdominal fat pad, rectal, or sural nerve biopsy can be performed to look for amyloid deposition. Unfortunately, the treatment of the underlying MM does not usually affect the course of the neuropathy.

NEUROPATHIES ASSOCIATED WITH MONOCLONAL GAMMOPATHY OF UNCERTAIN SIGNIFICANCE (SEE CHAP. 460)

Toxic Neuropathies Secondary to Chemotherapy Many of the commonly used chemotherapy agents can cause a toxic neuropathy (Table 459-7). The mechanisms by which these agents cause toxic neuropathies vary, as does the specific type of neuropathy produced. The risk of developing a toxic neuropathy or more severe neuropathy appears to be greater in patients with a preexisting neuropathy (e.g., Charcot-Marie-Tooth disease, diabetic neuropathy) and those who also take other potentially neurotoxic drugs (e.g., nitrofurantoin, isoniazid, disulfiram, pyridoxine). Chemotherapeutic agents usually cause a sensory greater than motor length-dependent axonal neuropathy or neuropathy/ganglionopathy.

OTHER TOXIC NEUROPATHIES

Neuropathies can develop as complications of toxic effects of various drugs and other environmental exposures (Table 459-8). The more common neuropathies associated with these agents are discussed here.

CHLOROQUINE AND HYDROXYCHLOROQUINE

Chloroquine and hydroxychloroquine can cause a toxic myopathy characterized by slowly progressive, painless, proximal weakness and atrophy, which is worse in the legs than the arms. In addition, neuropathy can also develop with or without the myopathy leading to sensory loss and distal weakness. The “neuromyopathy” usually appears in patients taking 500 mg daily for a year or more but has been reported with doses as low as 200 mg/d. Serum CK levels are usually elevated due to the superimposed myopathy. NCS reveal mild slowing of motor and sensory NCVs with a mild to moderate reduction in the amplitudes, although NCS may be normal in patients with only the myopathy. EMG demonstrates myopathic muscle action potentials (MUAPs), increased insertional activity in the form of positive sharp waves, fibrillation potentials, and occasionally myotonic potentials, particularly in the proximal muscles. Neurogenic MUAPs and reduced recruitment are found in more distal muscles. Nerve biopsy demonstrates autophagic vacuoles within Schwann cells. Vacuoles may also be evident in muscle biopsies. The pathogenic basis of the neuropathy is not known but may be related to the amphiphilic properties of the drug. These agents contain both hydrophobic and hydrophilic regions

TABLE 459-7 TOXIC NEUROPATHIES SECONDARY TO CHEMOTHERAPY

Drug	Mechanism of Neurotoxicity	Clinical Features	Nerve Histopathology	EMG/NCS
Vinca alkaloids (vincristine, vinblastine, vindesine, vinorelbine)	Interfere with axonal microtubule assembly; impairs axonal transport	Symmetric, S-M, large-/small-fiber PN; autonomic symptoms common; infrequent cranial neuropathies	Axonal degeneration of myelinated and unmyelinated fibers; regenerating clusters, minimal segmental demyelination	Axonal sensorimotor PN; distal denervation on EMG; abnormal QST, particularly vibratory perception
Cisplatin	Preferential damage to dorsal root ganglia; ? binds to and cross-links DNA ? inhibits protein synthesis ? impairs axonal transport	Predominant large-fiber sensory neuropathy; sensory ataxia	Loss of large > small myelinated and unmyelinated fibers; axonal degeneration with small clusters of regenerating fibers; secondary segmental demyelination	Low-amplitude or unobtainable SNAPs with normal CMAPs and EMG; abnormal QST, particularly vibratory perception
Taxanes (paclitaxel, docetaxel)	Promotes axonal microtubule assembly; interferes with axonal transport	Symmetric, predominantly sensory PN; large-fiber modalities affected more than small-fiber	Loss of large > small myelinated and unmyelinated fibers; axonal degeneration with small clusters of regenerating fibers; secondary segmental demyelination	Axonal sensorimotor PN; distal denervation on EMG; abnormal QST, particularly vibratory perception
Suramin				
Axonal PN	Unknown; ? inhibition of neurotrophic growth factor binding; ? neuronal lysosomal storage	Symmetric, length-dependent, sensory-predominant PN	None described	Abnormalities consistent with an axonal S-M PN
Demyelinating PN	Unknown; ? immunomodulating effects	Subacute, S-M PN with diffuse proximal and distal weakness; areflexia; increased CSF protein	Loss of large and small myelinated fibers with primary demyelination and secondary axonal degeneration; occasional epi- and endoneurial inflammatory cell infiltrates	Features suggestive of an acquired demyelinating sensorimotor PN (e.g., slow CVs, prolonged distal latencies and F-wave latencies, conduction block, temporal dispersion)
Cytarabine (ARA-C)	Unknown; ? selective Schwann cell toxicity; ? immunomodulating effects	GBS-like syndrome; pure sensory neuropathy; brachial plexopathy	Loss of myelinated nerve fibers; axonal degeneration; segmental demyelination; no inflammation	Axonal, demyelinating, or mixed S-M PN; denervation on EMG
Etoposide (VP-16)	Unknown; ? selective dorsal root ganglia toxicity	Length-dependent, sensory-predominant PN; autonomic neuropathy	None described	Abnormalities consistent with an axonal S-M PN
Bortezomib (Velcade)	Unknown	Length-dependent, sensory, predominantly small-fiber PN	Not reported	Abnormalities consistent with an axonal sensory neuropathy with early small-fiber involvement (abnormal autonomic studies)

Abbreviations: CMAP, compound motor action potential; CSF, cerebrospinal fluid; CVs, conduction velocities; EMG, electromyography; GBS, Guillain-Barré syndrome; NCS, nerve conduction studies; PN, polyneuropathy; QST, quantitative sensory testing; S-M, sensorimotor; SNAP, sensory nerve action potential.

Source: From AA Amato, J Russell: *Neuromuscular Disease*. New York, McGraw-Hill, 2008.

TABLE 459-8 TOXIC NEUROPATHIES

Drug	Mechanism of Neurotoxicity	Clinical Features	Nerve Histopathology	EMG/NCS
Misonidazole	Unknown	Painful paresthesias and loss of large- and small-fiber sensory modalities and sometimes distal weakness in length-dependent pattern	Axonal degeneration of large myelinated fibers; axonal swellings; segmental demyelination	Low-amplitude or unobtainable SNAPs with normal or only slightly reduced CMAPs amplitudes
Metronidazole	Unknown	Painful paresthesias and loss of large- and small-fiber sensory modalities and sometimes distal weakness in length-dependent pattern	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal CMAPs
Chloroquine and hydroxychloroquine	Amphiphilic properties may lead to drug-lipid complexes that are indigestible and result in accumulation of autophagic vacuoles	Loss of large- and small-fiber sensory modalities and distal weakness in length-dependent pattern; superimposed myopathy may lead to proximal weakness	Axonal degeneration with autophagic vacuoles in nerves as well as muscle fibers	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes; distal denervation on EMG; irritability and myopathic-appearing MUAPs proximally in patients with superimposed toxic myopathy
Amiodarone	Amphiphilic properties may lead to drug-lipid complexes that are indigestible and result in accumulation of autophagic vacuoles	Paresthesias and pain with loss of large- and small-fiber sensory modalities and distal weakness in length-dependent pattern; superimposed myopathy may lead to proximal weakness	Axonal degeneration and segmental demyelination with myeloid inclusions in nerves and muscle fibers	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes; can also have prominent slowing of CVs; distal denervation on EMG; irritability and myopathic-appearing MUAPs proximally in patients with superimposed toxic myopathy
Colchicine	Inhibits polymerization of tubulin in microtubules and impairs axoplasmic flow	Numbness and paresthesias with loss of large-fiber modalities in a length-dependent fashion; superimposed myopathy may lead to proximal in addition to distal weakness	Nerve biopsy demonstrates axonal degeneration; muscle biopsy reveals fibers with vacuoles	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes; irritability and myopathic-appearing MUAPs proximally in patients with superimposed toxic myopathy
Podophyllin	Binds to microtubules and impairs axoplasmic flow	Sensory loss, tingling, muscle weakness, and diminished muscle stretch reflexes in length-dependent pattern; autonomic neuropathy	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Thalidomide	Unknown	Numbness, tingling, and burning pain and weakness in a length-dependent pattern	Axonal degeneration; autopsy studies reveal degeneration of dorsal root ganglia	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Disulfiram	Accumulation of neurofilaments and impaired axoplasmic flow	Numbness, tingling, and burning pain in a length-dependent pattern	Axonal degeneration with accumulation of neurofilaments in the axons	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Dapsone	Unknown	Distal weakness that may progress to proximal muscles; sensory loss	Axonal degeneration and segmental demyelination	Low-amplitude or unobtainable CMAPs with normal or reduced SNAP amplitudes
Leflunomide	Unknown	Paresthesias and numbness in a length-dependent pattern	Unknown	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Nitrofurantoin	Unknown	Numbness, painful paresthesias, and severe weakness that may resemble GBS	Axonal degeneration; autopsy studies reveal degeneration of dorsal root ganglia and anterior horn cells	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Pyridoxine (vitamin B ₆)	Unknown	Dysesthesias and sensory ataxia; impaired large-fiber sensory modalities on examination	Marked loss of sensory axons and cell bodies in dorsal root ganglia	Reduced amplitudes or absent SNAPs
Isoniazid	Inhibits pyridoxal phosphokinase leading to pyridoxine deficiency	Dysesthesias and sensory ataxia; impaired large-fiber sensory modalities on examination	Marked loss of sensory axons and cell bodies in dorsal root ganglia and degeneration of the dorsal columns	Reduced amplitudes or absent SNAPs and, to a lesser extent, CMAPs
Ethambutol	Unknown	Numbness with loss of large-fiber modalities on examination	Axonal degeneration	Reduced amplitudes or absent SNAPs
Antinucleosides	Unknown	Dysesthesia and sensory ataxia; impaired large-fiber sensory modalities on examination	Axonal degeneration	Reduced amplitudes or absent SNAPs
Phenytoin	Unknown	Numbness with loss of large-fiber modalities on examination	Axonal degeneration and segmental demyelination	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Lithium	Unknown	Numbness with loss of large-fiber modalities on examination	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes

(Continued)

TABLE 459-8 TOXIC NEUROPATHIES (CONTINUED)

Drug	Mechanism of Neurotoxicity	Clinical Features	Nerve Histopathology	EMG/NCS
Acrylamide	Unknown; may be caused by impaired axonal transport	Numbness with loss of large-fiber modalities on examination; sensory ataxia; mild distal weakness	Degeneration of sensory axons in peripheral nerves and posterior columns, spinocerebellar tracts, mammillary bodies, optic tracts, and corticospinal tracts in the CNS	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Carbon disulfide	Unknown	Length-dependent numbness and tingling with mild distal weakness	Axonal swellings with accumulation of neurofilaments	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Ethylene oxide	Unknown; may act as alkylating agent and bind DNA	Length-dependent numbness and tingling; may have mild distal weakness	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Organophosphates	Bind and inhibit neuropathy target esterase	Early features are those of neuromuscular blockade with generalized weakness; later axonal sensorimotor PN ensues	Axonal degeneration along with degeneration of gracile fasciculus and corticospinal tracts	Early: repetitive firing of CMAPs and decrement with repetitive nerve stimulation; late: axonal sensorimotor PN
Hexacarbons	Unknown; may lead to covalent cross-linking between neurofilaments	Acute, severe sensorimotor PN that may resemble GBS	Axonal degeneration and giant axons swollen with neurofilaments	Features of a mixed axonal and/or demyelinating sensorimotor axonal PN—reduced amplitudes, prolonged distal latencies, conduction block, and slowing of CVs
Lead	Unknown; may interfere with mitochondria	Encephalopathy; motor neuropathy (often resembles radial neuropathy with wrist and finger drop); autonomic neuropathy; bluish-black discoloration of gums	Axonal degeneration of motor axons	Reduction of CMAP amplitudes with active denervation on EMG
Mercury	Unknown; may combine with sulfhydryl groups	Abdominal pain and nephrotic syndrome; encephalopathy; ataxia; paresthesias	Axonal degeneration; degeneration of dorsal root ganglia, calcarine, and cerebellar cortex	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Thallium	Unknown	Encephalopathy; painful sensory symptoms; mild loss of vibration; distal or generalized weakness may also develop; autonomic neuropathy; alopecia	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes
Arsenic	Unknown; may combine with sulfhydryl groups	Abdominal discomfort, burning pain, and paresthesias; generalized weakness; autonomic insufficiency; can resemble GBS	Axonal degeneration	Low-amplitude or unobtainable SNAPs with normal or reduced CMAP amplitudes; may have demyelinating features: prolonged distal latencies and slowing of CVs
Gold	Unknown	Distal paresthesias and reduction of all sensory modalities	Axonal degeneration	Low-amplitude or unobtainable SNAPs

Abbreviations: CMAP, compound motor action potential; CVs, conduction velocities; EMG, electromyography; GBS, Guillain-Barré syndrome; MUAP, muscle action potential; NCS, nerve conduction studies; PN, polyneuropathy; S-M, sensorimotor; SNAP, sensory nerve action potential

Source: From AA Amato, J Russell: *Neuromuscular Disease*. New York, McGraw-Hill, 2008.

that allow them to interact with the anionic phospholipids of cell membranes and organelles. The drug-lipid complexes may be resistant to digestion by lysosomal enzymes, leading to the formation of autophagic vacuoles filled with myeloid debris that may in turn cause degeneration of nerves and muscle fibers. The signs and symptoms of the neuropathy and myopathy are usually reversible following discontinuation of medication.

AMIODARONE

Amiodarone can cause a neuromyopathy similar to chloroquine and hydroxychloroquine. The neuromyopathy typically appears after patients have taken the medication for 2–3 years. Nerve biopsy demonstrates a combination of segmental demyelination and axonal loss. Electron microscopy reveals lamellar or dense inclusions in Schwann cells, pericytes, and endothelial cells. The inclusions in muscle and nerve biopsies have persisted as long as 2 years following discontinuation of the medication.

COLCHICINE

Colchicine can also cause a neuromyopathy. Patients usually present with proximal weakness and numbness and tingling in the distal extremities. EDx reveals features of an axonal polyneuropathy. Muscle biopsy reveals a vacuolar myopathy, whereas sensory nerves demonstrate

axonal degeneration. Colchicine inhibits the polymerization of tubulin into microtubules. The disruption of the microtubules probably leads to defective intracellular movement of important proteins, nutrients, and waste products in muscle and nerves.

THALIDOMIDE

Thalidomide is an immunomodulating agent used to treat multiple myeloma, GVHD, leprosy, and other autoimmune disorders. Thalidomide is associated with severe teratogenic effects as well as peripheral neuropathy that can be dose-limiting. Patients develop numbness, painful tingling, and burning discomfort in the feet and hands and less commonly muscle weakness and atrophy. Even after stopping the drug for 4–6 years, as many as 50% patients continue to have significant symptoms. NCS demonstrate reduced amplitudes or complete absence of SNAPs, with preserved conduction velocities when obtainable. Motor NCS are usually normal. Nerve biopsy reveals a loss of large-diameter myelinated fibers and axonal degeneration. Degeneration of dorsal root ganglion cells has been reported at autopsy.

PYRIDOXINE (VITAMIN B₆) TOXICITY

Pyridoxine is an essential vitamin that serves as a coenzyme for transamination and decarboxylation. However, at high doses (116 mg/d), patients can develop a severe sensory neuropathy with dysesthesias

and sensory ataxia. NCS reveal absent or markedly reduced SNAP amplitudes with relatively preserved CMAPs. Nerve biopsy reveals axonal loss of fiber at all diameters. Loss of dorsal root ganglion cells with subsequent degeneration of both the peripheral and central sensory tracts have been reported in animal models.

ISONIAZID

One of the most common side effects of isoniazid (INH) is peripheral neuropathy. Standard doses of INH (3–5 mg/kg per day) are associated with a 2% incidence of neuropathy, whereas neuropathy develops in at least 17% of patients taking in excess of 6 mg/kg per day. The elderly, malnourished, and “slow acetylators” are at increased risk for developing the neuropathy. INH inhibits pyridoxal phosphokinase, resulting in pyridoxine deficiency and the neuropathy. Prophylactic administration of pyridoxine 100 mg/d can prevent the neuropathy from developing.

ANTIRETROVIRAL AGENTS

The nucleoside analogues zalcitabine (dideoxycytidine or ddC), didanosine (dideoxyinosine or ddi), stavudine (d4T), lamivudine (3TC), and antiretroviral nucleoside reverse transcriptase inhibitor (NRTI) are used to treat HIV infection. One of the major dose-limiting side effects of these medications is a predominantly sensory, length-dependent, symmetrically painful neuropathy. Zalcitabine (ddC) is the most extensively studied of the nucleoside analogues, and at doses greater than 0.18 mg/kg per day, it is associated with a subacute onset of severe burning and lancinating pains in the feet and hands. NCS reveal decreased amplitudes of the SNAPS with normal motor studies. The nucleoside analogues inhibit mitochondrial DNA polymerase, which is the suspected pathogenic basis for the neuropathy. Because of a “coasting effect,” patients can continue to worsen even 2–3 weeks after stopping the medication. Following dose reduction, improvement in the neuropathy is seen in most patients after several months (mean time about 10 weeks).

HEXACARBONS (*n*-HEXANE, METHYL *n*-BUTYL KETONE)/GLUE SNIFFER'S NEUROPATHY

n-Hexane and methyl *n*-butyl ketone are water-insoluble industrial organic solvents that are also present in some glues. Exposure through inhalation, accidentally or intentionally (glue sniffing), or through skin absorption can lead to a profound subacute sensory and motor polyneuropathy. NCS demonstrate decreased amplitudes of the SNAPS and CMAPs with slightly slow CVs. Nerve biopsy reveals a loss of myelinated fibers and giant axons that are filled with 10-nm neurofilaments. Hexacarbon exposure leads to covalent cross-linking between axonal neurofilaments that result in their aggregation, impaired axonal transport, swelling of the axons, and eventual axonal degeneration.

LEAD

Lead neuropathy is uncommon, but it can be seen in children who accidentally ingest lead-based paints in older buildings and in industrial workers exposed to lead-containing products. The most common presentation of lead poisoning is an encephalopathy; however, symptoms and signs of a primarily motor neuropathy can also occur. The neuropathy is characterized by an insidious and progressive onset of weakness usually beginning in the arms, in particular involving the wrist and finger extensors, resembling a radial neuropathy. Sensation is generally preserved; however, the autonomic nervous system can be affected. Laboratory investigation can reveal a microcytic hypochromic anemia with basophilic stippling of erythrocytes, an elevated serum lead level, and an elevated serum coproporphyrin level. A 24-h urine collection demonstrates elevated levels of lead excretion. The NCS may reveal reduced CMAP amplitudes, while the SNAPS are typically normal. The pathogenic basis may be related to abnormal porphyrin metabolism. The most important principle of management is to remove the source of the exposure. Chelation therapy with calcium disodium ethylene-diaminetetraacetic acid (EDTA), British anti-Lewisite (BAL), and penicillamine also demonstrates variable efficacy.

MERCURY

Mercury toxicity may occur as a result of exposure to either organic or inorganic mercurials. Mercury poisoning presents with paresthesias in hands and feet that progress proximally and may involve the face and tongue. Motor weakness can also develop. CNS symptoms often overshadow the neuropathy. EDx shows features of a primarily axonal sensorimotor polyneuropathy. The primary site of neuromuscular pathology appears to be the dorsal root ganglia. The mainstay of treatment is removing the source of exposure.

THALLIUM

Thallium can exist in a monovalent or trivalent form and is primarily used as a rodenticide. The toxic neuropathy usually manifests as burning paresthesias of the feet, abdominal pain, and vomiting. Increased thirst, sleep disturbances, and psychotic behavior may be noted. Within the first week, patients develop pigmentation of the hair, an acne-like rash in the malar area of the face, and hyperreflexia. By the second and third week, autonomic instability with labile heart rate and blood pressure may be seen. Hyporeflexia and alopecia also occur but may not be evident until the third or fourth week following exposure. With severe intoxication, proximal weakness and involvement of the cranial nerves can occur. Some patients require mechanical ventilation due to respiratory muscle involvement. The lethal dose of thallium is variable, ranging from 8 to 15 mg/kg body weight. Death can result in less than 48 h following a particularly large dose. NCS demonstrate features of a primarily axonal sensorimotor polyneuropathy. With acute intoxication, potassium ferric ferrocyanide II may be effective in preventing absorption of thallium from the gut. However, there may be no benefit once thallium has been absorbed. Unfortunately, chelating agents are not very efficacious. Adequate diuresis is essential to help eliminate thallium from the body without increasing tissue availability from the serum.

ARSENIC

Arsenic is another heavy metal that can cause a toxic sensorimotor polyneuropathy. The neuropathy manifests 5–10 days after ingestion of arsenic and progresses for several weeks, sometimes mimicking GBS. The presenting symptoms are typically an abrupt onset of abdominal discomfort, nausea, vomiting, pain, and diarrhea followed within several days by burning pain in the feet and hands. Examination of the skin can be helpful in the diagnosis as the loss of the superficial epidermal layer results in patchy regions of increased or decreased pigmentation on the skin several weeks after an acute exposure or with chronic low levels of ingestion. Mee's lines, which are transverse lines at the base of the fingernails and toenails, do not become evident until 1 or 2 months after the exposure. Multiple Mee's lines may be seen in patients with long fingernails who have had chronic exposure to arsenic. Mee's lines are not specific for arsenic toxicity as they can also be seen following thallium poisoning. Because arsenic is cleared from blood rapidly, the serum concentration of arsenic is not diagnostically helpful. However, arsenic levels are increased in the urine, hair, and fingernails of patients exposed to arsenic. Anemia with stippling of erythrocytes is common, and occasionally pancytopenia and aplastic anemia can develop. Increased CSF protein levels without pleocytosis can be seen; this can lead to misdiagnosis as GBS. NCS are usually suggestive of an axonal sensorimotor polyneuropathy; however, demyelinating features can be present. Chelation therapy with BAL has yielded inconsistent results; therefore, it is not generally recommended.

NUTRITIONAL NEUROPATHIES

COBALAMIN (VITAMIN B₁₂)

Pernicious anemia is the most common cause of cobalamin deficiency. Other causes include dietary avoidance (vegetarians), gastrectomy, gastric bypass surgery, inflammatory bowel disease, pancreatic insufficiency, bacterial overgrowth, and possibly histamine-2 blockers and proton pump inhibitors. An underappreciated cause of cobalamin deficiency is food-cobalamin malabsorption. This typically occurs in older individuals and results from an inability to adequately absorb

cobalamin in food protein. No apparent cause of deficiency is identified in a significant number of patients with cobalamin deficiency. The use of nitrous oxide as an anesthetic agent or as a recreational drug can produce acute cobalamin deficiency neuropathy and subacute combined degeneration.

Complaints of numb hands typically appear before lower extremity paresthesias are noted. A preferential large-fiber sensory loss affecting proprioception and vibration with sparing of small-fiber modalities is present; an unsteady gait reflects sensory ataxia. These features, coupled with diffuse hyperreflexia and absent Achilles reflexes, should always focus attention on the possibility of cobalamin deficiency. Optic atrophy and, in severe cases, behavioral changes ranging from mild irritability and forgetfulness to severe dementia and frank psychosis may appear. The full clinical picture of subacute combined degeneration is uncommon. CNS manifestations, especially pyramidal tract signs, may be missing, and in fact some patients may only exhibit symptoms of peripheral neuropathy.

EDx shows an axonal sensorimotor neuropathy. CNS involvement produces abnormal somatosensory and visual evoked potential latencies. The diagnosis is confirmed by finding reduced serum cobalamin levels. In up to 40% of patients, anemia and macrocytosis are lacking. Serum methylmalonic acid and homocysteine, the metabolites that accumulate when cobalamin-dependent reactions are blocked, are elevated. Antibodies to intrinsic factor are present in approximately 60%, and antiparietal cell antibodies in about 90%, of individuals with pernicious anemia.

Cobalamin deficiency can be treated with various regimens of cobalamin. One typical regimen consists of 1000 µg cyanocobalamin IM weekly for 1 month and monthly thereafter. Patients with food cobalamin malabsorption can absorb free cobalamin and therefore can be treated with oral cobalamin supplementation. An oral cobalamin dose of 1000 µg per day should be sufficient. Treatment for cobalamin deficiency usually does not completely reverse the clinical manifestations, and at least 50% of patients exhibit some permanent neurologic deficit.

THIAMINE DEFICIENCY

Thiamine (vitamin B₁) deficiency is an uncommon cause of peripheral neuropathy in developed countries. It is now most often seen as a consequence of chronic alcohol abuse, recurrent vomiting, total parenteral nutrition, and bariatric surgery. Thiamine deficiency polyneuropathy can occur in normal, healthy young adults who do not abuse alcohol but who engage in inappropriately restrictive diets. Thiamine is water-soluble. It is present in most animal and plant tissues, but the greatest sources are unrefined cereal grains, wheat germ, yeast, soybean flour, and pork. Beriberi means “I can’t, I can’t” in Singhaese, the language of natives of what was once part of the Dutch East Indies (now Sri Lanka). *Dry beriberi* refers to neuropathic symptoms. The term *wet beriberi* is used when cardiac manifestations predominate (in reference to edema). Beriberi was relatively uncommon until the late 1800s when it became widespread among people for whom rice was a dietary mainstay. This epidemic was due to a new technique of processing rice that removed the germ from the rice shaft, rendering the so-called polished rice deficient in thiamine and other essential nutrients.

Symptoms of neuropathy follow prolonged deficiency. These begin with mild sensory loss and/or burning dysesthesias in the toes and feet and aching and cramping in the lower legs. Pain may be the predominant symptom. With progression, patients develop features of a nonspecific generalized polyneuropathy, with distal sensory loss in the feet and hands.

Blood and urine assays for thiamine are not reliable for diagnosis of deficiency. Erythrocyte transketolase activity and the percentage increase in activity (in vitro) following the addition of thiamine pyrophosphate (TPP) may be more accurate and reliable. EDx shows nonspecific findings of an axonal sensorimotor polyneuropathy. When a diagnosis of thiamine deficiency is made or suspected, thiamine replacement should be provided until proper nutrition is restored. Thiamine is usually given intravenously or intramuscularly at a dose of 100 mg/d. Although cardiac manifestations show a striking response to

thiamine replacement, neurologic improvement is usually more variable and less dramatic.

VITAMIN E DEFICIENCY

The term *vitamin E* is usually used for α -tocopherol, the most active of the four main types of vitamin E. Because vitamin E is present in animal fat, vegetable oils, and various grains, deficiency is usually due to factors other than insufficient intake. Vitamin E deficiency usually occurs secondary to lipid malabsorption or in uncommon disorders of vitamin E transport. One hereditary disorder is abetalipoproteinemia, a rare autosomal dominant disorder characterized by steatorrhea, pigmentary retinopathy, acanthocytosis, and progressive ataxia. Patients with cystic fibrosis may also have vitamin E deficiency secondary to steatorrhea. There are genetic forms of isolated vitamin E deficiency not associated with lipid malabsorption. Vitamin E deficiency may also occur as a consequence of various cholestatic and hepatobiliary disorders as well as short-bowel syndromes resulting from the surgical treatment of intestinal disorders.

Clinical features may not appear until many years after the onset of deficiency. The onset of symptoms tends to be insidious, and progression is slow. The main clinical features are spinocerebellar ataxia and polyneuropathy, thus resembling Friedreich’s ataxia or other spinocerebellar ataxias. Patients manifest progressive ataxia and signs of posterior column dysfunction, such as impaired joint position and vibratory sensation. Because of the polyneuropathy, there is hyporeflexia, but plantar responses may be extensor as a result of the spinal cord involvement. Other neurologic manifestations may include ophthalmoplegia, pigmented retinopathy, night blindness, dysarthria, pseudoathetosis, dystonia, and tremor. Vitamin E deficiency may present as an isolated polyneuropathy, but this is very rare. The yield of checking serum vitamin E levels in patients with isolated polyneuropathy is extremely low, and this test should not be part of routine practice.

Diagnosis is made by measuring α -tocopherol levels in the serum. EDx shows features of an axonal neuropathy. Treatment is replacement with oral vitamin E, but high doses are not needed. For patients with isolated vitamin E deficiency, treatment consists of 1500–6000 IU/d in divided doses.

VITAMIN B₆ DEFICIENCY

Vitamin B₆, or pyridoxine, can produce neuropathic manifestations from both deficiency and toxicity. Vitamin B₆ toxicity was discussed above. Vitamin B₆ deficiency is most commonly seen in patients treated with isoniazid or hydralazine. The polyneuropathy of vitamin B₆ is nonspecific, manifesting as a generalized axonal sensorimotor polyneuropathy. Vitamin B₆ deficiency can be detected by direct assay. Vitamin B₆ supplementation with 50–100 mg/d is suggested for patients being treated with isoniazid or hydralazine. This same dose is appropriate for replacement in cases of nutritional deficiency.

PELLAGRA (NIACIN DEFICIENCY)

Pellagra is produced by deficiency of niacin. Although pellagra may be seen in alcoholics, this disorder has essentially been eradicated in most Western countries by means of enriching bread with niacin. Nevertheless, pellagra continues to be a problem in a number of underdeveloped regions, particularly in Asia and Africa, where corn is the main source of carbohydrate. Neurologic manifestations are variable; abnormalities can develop in the brain and spinal cord as well as peripheral nerves. When peripheral nerves are involved, the neuropathy is usually mild and resembles beriberi. Treatment is with niacin 40–250 mg/d.

COPPER DEFICIENCY

A syndrome that has only recently been described is myeloneuropathy secondary to copper deficiency. Most patients present with lower limb paresthesias, weakness, spasticity, and gait difficulties. Large-fiber sensory function is impaired, reflexes are brisk, and plantar responses are extensor. In some cases, light touch and pinprick sensation are affected, and NCS indicate sensorimotor axonal polyneuropathy in addition to myelopathy.

Hematologic abnormalities are a known complication of copper deficiency; these can include microcytic anemia, neutropenia, and occasionally pancytopenia. Because copper is absorbed in the stomach and proximal jejunum, many cases of copper deficiency are in the setting of prior gastric surgery. Excess zinc is an established cause of copper deficiency. Zinc upregulates enterocyte production of metallothionein, which results in decreased absorption of copper. Excessive dietary zinc supplements or denture cream containing zinc can produce this clinical picture. Other potential causes of copper deficiency include malnutrition, prematurity, total parenteral nutrition, and ingestion of copper-chelating agents.

Following oral or IV copper replacement, some patients show neurologic improvement, but this may take many months or not occur at all. Replacement consists of oral copper sulfate or gluconate 2 mg one to three times a day. If oral copper replacement is not effective, elemental copper in the copper sulfate or copper chloride forms can be given as 2 mg IV daily for 3–5 days, then weekly for 1–2 months until copper levels normalize. Thereafter, oral daily copper therapy can be resumed. In contrast to the neurologic manifestations, most of the hematologic indices completely normalize in response to copper replacement therapy.

NEUROPATHY ASSOCIATED WITH GASTRIC SURGERY

Polynuropathy may occur following gastric surgery for ulcer, cancer, or weight reduction. This usually occurs in the context of rapid, significant weight loss and recurrent, protracted vomiting. The clinical picture is one of acute or subacute sensory loss and weakness. Neuropathy following weight loss surgery usually occurs in the first several months after surgery. Weight reduction surgical procedures include gastrojejunostomy, gastric stapling, vertical banded gastroplasty, and gastrectomy with Roux-en-Y anastomosis. The initial manifestations are usually numbness and paresthesias in the feet. In many cases, no specific nutritional deficiency factor is identified.

Management consists of parenteral vitamin supplementation, especially including thiamine. Improvement has been observed following supplementation, parenteral nutritional support, and reversal of the surgical bypass. The duration and severity of deficits before identification and treatment of neuropathy are important predictors of final outcome.

CRYPTOGENIC (IDIOPATHIC) SENSORY AND SENSORIMOTOR POLYNEUROPATHY

CSPN is a diagnosis of exclusion, established after a careful medical, family, and social history; neurologic examination; and directed laboratory testing. Despite extensive evaluation, the cause of polyneuropathy in as many as 50% of all patients is idiopathic. CSPN should be considered a distinct diagnostic subset of peripheral neuropathy. The onset of CSPN is predominantly in the sixth and seventh decades. Patients complain of distal numbness, tingling, and often burning pain that invariably begins in the feet and may eventually involve the fingers and hands. Patients exhibit a distal sensory loss to pinprick, touch, and vibration in the toes and feet, and occasionally in the fingers. It is uncommon to see significant proprioception deficits, even though patients may complain of gait unsteadiness. However, tandem gait may be abnormal in a minority of cases. Neither subjective nor objective evidence of weakness is a prominent feature. Most patients have evidence of both large- and small-fiber loss on neurologic exam and EDx. Approximately 10% of patients have only evidence of small-fiber involvement. The ankle muscle stretch reflex is frequently absent, but in cases with predominantly small-fiber loss, this may be preserved. The EDx findings range from isolated sensory nerve action potential abnormalities (usually with loss of amplitude), to evidence for an axonal sensorimotor neuropathy, to a completely normal study (if primarily small fibers are involved). Therapy primarily involves the control of neuropathic pain (Table 459-6) if present. These drugs should not be used if the patient has only numbness and tingling but no pain.

Although no treatment is available that can reverse an idiopathic distal peripheral neuropathy, the prognosis is good. Progression often

does not occur or is minimal, with sensory symptoms and signs progressing proximally up to the knees and elbows. The disorder does not lead to significant motor disability over time. The relatively benign course of this disorder should be explained to patients.

MONONEUROPATHIES/PLEXOPATHIES/RADICULOPATHIES

MEDIAN NEUROPATHY

CTS is a compression of the median nerve in the carpal tunnel at the wrist. The median nerve enters the hand through the carpal tunnel by coursing under the transverse carpal ligament. The symptoms of CTS consist of numbness and paresthesias variably in the thumb, index, middle, and half of the ring finger. At times, the paresthesias can include the entire hand and extend into the forearm or upper arm or can be isolated to one or two fingers. Pain is another common symptom and can be located in the hand and forearm and, at times, in the proximal arm. CTS is common and often misdiagnosed as thoracic outlet syndrome. The signs of CTS are decreased sensation in the median nerve distribution; reproduction of the sensation of tingling when a percussion hammer is tapped over the wrist (Tinel's sign) or the wrist is flexed for 30–60 s (Phalen's sign); and weakness of thumb opposition and abduction. EDx is extremely sensitive and shows slowing of sensory and, to a lesser extent, motor median potentials across the wrist. Treatment options consist of avoidance of precipitating activities; control of underlying systemic-associated conditions if present; nonsteroidal anti-inflammatory medications; neutral (volar) position wrist splints, especially for night use; glucocorticoid/anesthetic injection into the carpal tunnel; and surgical decompression by dividing the transverse carpal ligament. The surgical option should be considered if there is a poor response to nonsurgical treatments; if there is thenar muscle atrophy and/or weakness; and if there are significant denervation potentials on EMG.

Other proximal median neuropathies are very uncommon and include the pronator teres syndrome and anterior interosseous neuropathy. These often occur as a partial form of brachial plexitis.

ULNAR NEUROPATHY AT THE ELBOW—"CUBITAL TUNNEL SYNDROME"

The ulnar nerve passes through the condylar groove between the medial epicondyle and the olecranon. Symptoms consist of paresthesias, tingling, and numbness in the medial hand and half of the fourth and the entire fifth fingers, pain at the elbow or forearm, and weakness. Signs consist of decreased sensation in an ulnar distribution, Tinel's sign at the elbow, and weakness and atrophy of ulnar-innervated hand muscles. The Froment sign indicates thumb adductor weakness and consists of flexion of the thumb at the interphalangeal joint when attempting to oppose the thumb against the lateral border of the second digit. EDx may show slowing of ulnar motor NCV across the elbow with prolonged ulnar sensory latencies. Treatment consists of avoiding aggravating factors, using elbow pads, and surgery to decompress the nerve in the cubital tunnel. Ulnar neuropathies can also rarely occur at the wrist in the ulnar (Guyon) canal or in the hand, usually after trauma.

RADIAL NEUROPATHY

The radial nerve winds around the proximal humerus in the spiral groove and proceeds down the lateral arm and enters the forearm, dividing into the posterior interosseous nerve and superficial nerve. The symptoms and signs consist of wristdrop; finger extension weakness; thumb abduction weakness; and sensory loss in the dorsal web between the thumb and index finger. Triceps and brachioradialis strength is often normal, and triceps reflex is often intact. Most cases of radial neuropathy are transient compressive (neuropraxic) injuries that recover spontaneously in 6–8 weeks. If there has been prolonged compression and severe axonal damage, it may take several months to recover. Treatment consists of cock-up wrist and finger splints, avoiding further compression, and physical therapy to avoid flexion contracture. If there is no improvement in 2–3 weeks, an EDx study is recommended to confirm the clinical diagnosis and determine the degree of severity.

The lateral femoral cutaneous nerve arises from the upper lumbar plexus (spinal levels L2/3), crosses through the inguinal ligament near its attachment to the iliac bone, and supplies sensation to the anterior lateral thigh. The neuropathy affecting this nerve is also known as meralgia paresthetica. Symptoms and signs consist of paresthesias, numbness, and occasionally pain in the lateral thigh. Symptoms are increased by standing or walking and are relieved by sitting. There is normal strength, and knee reflexes are intact. The diagnosis is clinical, and further tests usually are not performed. EDx is only needed to rule out lumbar plexopathy, radiculopathy, or femoral neuropathy. If the symptoms and signs are classic, EMG is not necessary. Symptoms often resolve spontaneously over weeks or months, but the patient may be left with permanent numbness. Treatment consists of weight loss and avoiding tight belts. Analgesics in the form of a lidocaine patch, nonsteroidal agents, and occasionally medications for neuropathic pain can be used (Table 459-6). Rarely, locally injecting the nerve with an anesthetic can be tried. There is no role for surgery.

FEMORAL NEUROPATHY

Femoral neuropathies can arise as complications of retroperitoneal hematoma, lithotomy positioning, hip arthroplasty or dislocation, iliac artery occlusion, femoral arterial procedures, infiltration by hematogenous malignancy, penetrating groin trauma, pelvic surgery including hysterectomy and renal transplantation, and diabetes (a partial form of lumbosacral diabetic plexopathy); some cases are idiopathic. Patients with femoral neuropathy have difficulty extending their knee and flexing the hip. Sensory symptoms occurring either on the anterior thigh and/or medial leg occur in only half of reported cases. A prominent painful component is the exception rather than the rule, may be delayed, and is often self-limited in nature. The quadriceps (patellar) reflex is diminished.

SCIATIC NEUROPATHY

Sciatic neuropathies commonly complicate hip arthroplasty, pelvic procedures in which patients are placed in a prolonged lithotomy position, trauma, hematomas, tumor infiltration, and vasculitis. In addition, many sciatic neuropathies are idiopathic. Weakness may involve all motions of the ankles and toes as well as flexion of the leg at the knee; abduction and extension of the thigh at the hip are spared. Sensory loss occurs in the entire foot and the distal lateral leg. The ankle jerk and on occasion the internal hamstring reflex are diminished or more typically absent on the affected side. The peroneal subdivision of the sciatic nerve is typically involved disproportionately to the tibial counterpart. Thus, patients may have only ankle dorsiflexion and eversion weakness with sparing of knee flexion, ankle inversion, and plantar flexion; these features can lead to misdiagnosis of a common peroneal neuropathy.

PERONEAL NEUROPATHY

The sciatic nerve divides at the distal femur into the tibial and peroneal nerve. The common peroneal nerve passes posterior and laterally around the fibular head, under the fibular tunnel. It then divides into the superficial peroneal nerve, which supplies the ankle evertor muscles and sensation over the anterolateral distal leg and dorsum of the foot, and the deep peroneal nerve, which supplies ankle dorsiflexors and toe extensor muscles and a small area of sensation dorsally in the area of the first and second toes.

Symptoms and signs consist of footdrop (ankle dorsiflexion, toe extension, and ankle eversion weakness) and variable sensory loss, which may involve the superficial and deep peroneal pattern. There is usually no pain. Onset may be on awakening in the morning. Peroneal neuropathy needs to be distinguished from L5 radiculopathy. In L5 radiculopathy, ankle invertors and evertors are weak and needle EMG reveals denervation. EDx can help localize the lesion. Peroneal motor conduction velocity shows slowing and amplitude drop across the fibular head. Management consists of rapid weight loss and avoiding leg crossing. Footdrop is treated with an ankle brace. A knee pad can

be worn over the lateral knee to avoid further compression. Most cases spontaneously resolve over weeks or months.

RADICULOPATHIES

Radiculopathies are most often due to compression from degenerative joint disease and herniated disks, but there are a number of unusual etiologies (Table 459-9). Degenerative spine disease affects a number of different structures, which narrow the diameter of the neural foramen or canal of the spinal column and compromise nerve root integrity; these are discussed in detail in Chap. 22.

PLEXOPATHIES

BRACHIAL PLEXUS

The brachial plexus is composed of three trunks (upper, middle, and lower), with two divisions (anterior and posterior) per trunk (Fig. 459-2). Subsequently, the trunks divide into three cords (medial, lateral, and posterior), and from these arise the multiple terminal nerves innervating the arm. The anterior primary rami of C5 and C6 fuse to form the upper trunk; the anterior primary ramus of C7 continues as the middle trunk, while the anterior rami of C8 and T1 join to form the lower trunk. There are several disorders commonly associated with brachial plexopathy.

Immune-Mediated Brachial Plexus Neuropathy Immune-mediated brachial plexus neuropathy (IBPN) goes by various terms, including *acute brachial plexitis*, *neuralgic amyotrophy*, and *Parsonage-Turner syndrome*. IBPN usually presents with an acute onset of severe pain in the shoulder region. The intense pain usually lasts several days to a few weeks, but a dull ache can persist. Individuals who are affected may not appreciate weakness of the arm early in the course because the pain limits movement. However, as the pain dissipates, weakness and often sensory loss are appreciated. Attacks can occasionally recur.

Clinical findings are dependent on the distribution of involvement (e.g., specific trunk, divisions, cords, or terminal nerves). The most common pattern of IBPN involves the upper trunk or a single or multiple mononeuropathies primarily involving the suprascapular, long thoracic, or axillary nerves. Additionally, the phrenic and anterior interosseous nerves may be concomitantly affected. Any of these nerves may also be affected in isolation. EDx is useful to confirm and localize the site(s) of involvement. Empirical treatment of severe pain with glucocorticoids is often used in the acute period.

Brachial Plexopathies Associated with Neoplasms Neoplasms involving the brachial plexus may be primary nerve tumors, local cancers expanding into the plexus (e.g., Pancoast lung tumor or lymphoma),

TABLE 459-9 CAUSES OF RADICULOPATHY

- Herniated nucleus pulposus
- Degenerative joint disease
- Rheumatoid arthritis
- Trauma
- Vertebral body compression fracture
- Pott's disease
- Compression by extradural mass (e.g., meningioma, metastatic tumor, hematoma, abscess)
- Primary nerve tumor (e.g., neurofibroma, schwannoma, neurinoma)
- Carcinomatous meningitis
- Perineurial spread of tumor (e.g., prostate cancer)
- Acute inflammatory demyelinating polyradiculopathy
- Chronic inflammatory demyelinating polyradiculopathy
- Sarcoidosis
- Amyloidoma
- Diabetic radiculopathy
- Infection (Lyme disease, herpes zoster, cytomegalovirus, syphilis, schistosomiasis, strongyloides)

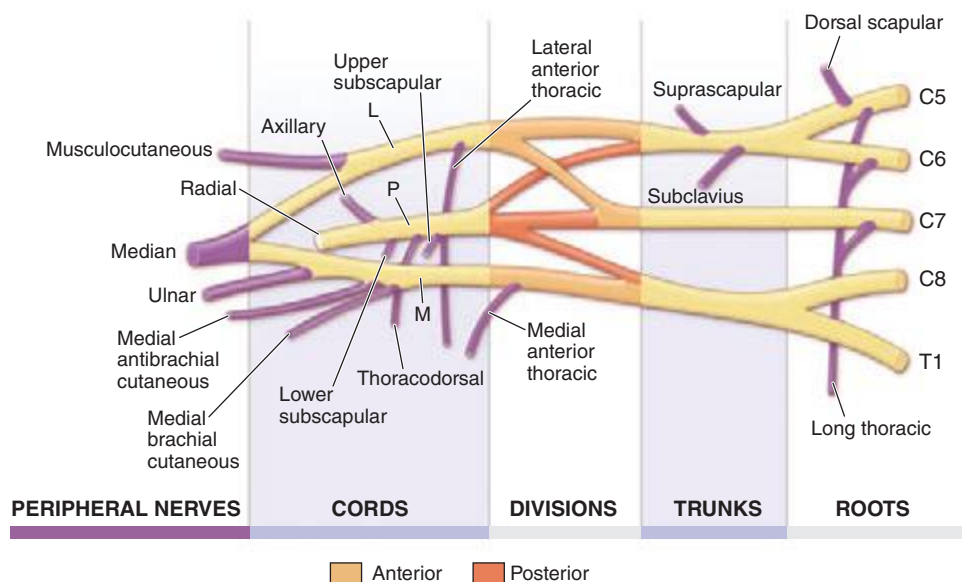


FIGURE 459-2 Brachial plexus anatomy. L, lateral; M, medial; P, posterior. (From J Goodgold: *Anatomical Correlates of Clinical Electromyography*. Baltimore, Williams and Wilkins, 1974, p. 126, with permission.)

and metastatic tumors. Primary brachial plexus tumors are less common than the secondary tumors and include schwannomas, neurofibromas, and neurofibromas. Secondary tumors affecting the brachial plexus are more common and are always malignant. These may arise from local tumors, expanding into the plexus. For example, a Pancoast tumor of the upper lobe of the lung may invade or compress the lower trunk, whereas a primary lymphoma arising from the cervical or axillary lymph nodes may also infiltrate the plexus. Pancoast tumors typically present as an insidious onset of pain in the upper arm, sensory disturbance in the medial aspect of the forearm and hand, and weakness and atrophy of the intrinsic hand muscles along with an ipsilateral Horner's syndrome. Chest computed tomography (CT) scans or magnetic resonance imaging (MRI) can demonstrate extension of the tumor into the plexus. Metastatic involvement of the brachial plexus may occur with spread of breast cancer into the axillary lymph nodes with local spread into the nearby nerves.

Perioperative Plexopathies (Median Sternotomy) The most common surgical procedures associated with brachial plexopathy as a complication are those that involve median sternotomies (e.g., open-heart surgeries and thoracotomies). Brachial plexopathies occur in as many as 5% of patients following a median sternotomy and typically affect the lower trunk. Thus, individuals manifest with sensory disturbance affecting the medial aspect of forearm and hand along with weakness of the intrinsic hand muscles. The mechanism is related to the stretch of the lower trunk, so most individuals who are affected recover within a few months.

Lumbosacral Plexus The lumbar plexus arises from the ventral primary rami of the first to the fourth lumbar spinal nerves (Fig. 459-3). These nerves pass downward and laterally from the vertebral column within the psoas major muscle. The femoral nerve derives from the dorsal branches of the second to the fourth lumbar ventral rami. The obturator nerve arises from the ventral branches of the same lumbar rami. The lumbar plexus communicates with the sacral plexus by the lumbosacral trunk, which contains some fibers from the fourth and all of the fibers from the fifth lumbar ventral rami (Fig. 459-4).

The sacral plexus is the part of the lumbosacral plexus that is formed by the union of the lumbosacral trunk with the ventral rami of the first to fourth sacral nerves. The plexus lies on the posterior and posterolateral wall of the pelvis with its components converging toward the sciatic notch. The lateral trunk of the sciatic nerve (which forms the common peroneal nerve) arises from the union of the dorsal branches of the lumbosacral trunk (L4, L5) and the dorsal branches of the S1 and S2 spinal nerve ventral rami. The medial trunk of the sciatic nerve

(which forms the tibial nerve) derives from the ventral branches of the same ventral rami (L4-S2).

LUMBOSACRAL PLEXOPATHIES

Plexopathies are typically recognized when motor, sensory, and if applicable, reflex deficits occur in multiple nerve and segmental distributions confined to one extremity. If localization within the lumbosacral plexus can be accomplished, designation as a lumbar plexopathy, a sacral plexopathy, a lumbosacral trunk lesion, or a plexopathy is the best localization that can be expected. Although lumbar plexopathies may be bilateral, usually occurring in a stepwise and chronologically dissociated manner, sacral plexopathies are more likely to behave in this manner due to their closer anatomic proximity. The differential diagnosis of plexopathy includes disorders of the conus medullaris and cauda equina (polyradiculopathy). If there is a paucity of pain and sensory involvement, motor neuron disease should be considered as well.

The causes of lumbosacral plexopathies are listed in Table 459-10. Diabetic radiculopathy (discussed above) is a fairly common cause of painful leg weakness. Lumbosacral plexopathies are a well-recognized

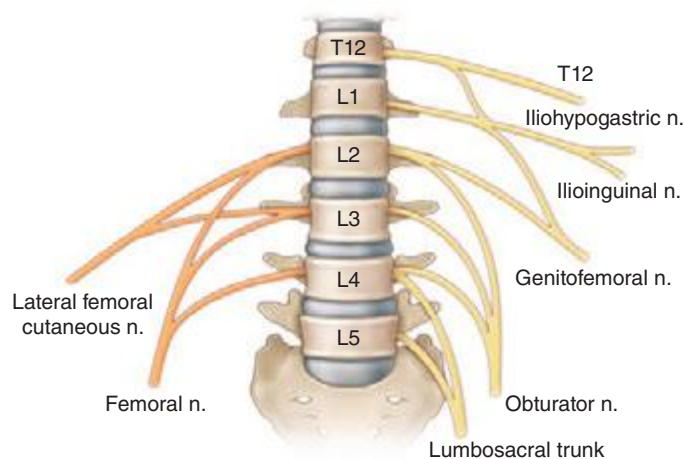


FIGURE 459-3 Lumbar plexus. Posterior divisions are in orange, and anterior divisions are in yellow. (From J Goodgold: *Anatomical Correlates of Clinical Electromyography*. Baltimore, Williams and Wilkins, 1974, p. 126, with permission.)

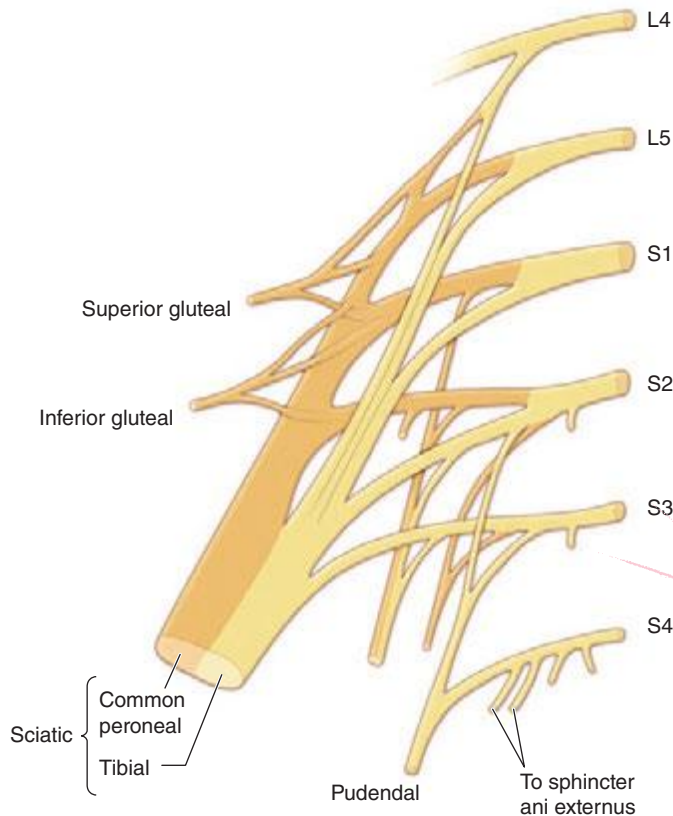


FIGURE 459-4 Lumbosacral plexus. Posterior divisions are in orange, and anterior divisions are in yellow. (From J Goodgold: *Anatomical Correlates of Clinical Electromyography*. Baltimore, Williams and Wilkins, 1974, p. 126, with permission.)

complication of retroperitoneal hemorrhage. Various primary and metastatic malignancies can affect the lumbosacral plexus as well; these include carcinoma of the cervix, endometrium, and ovary; osteosarcoma; testicular cancer; multiple myeloma; lymphoma; acute myelogenous leukemia; colon cancer; squamous cell carcinoma of the rectum; adenocarcinoma of unknown origin; and intraneural spread of prostate cancer.

RECURRENT NEOPLASTIC DISEASE OR RADIATION-INDUCED PLEXOPATHY

The treatment for various malignancies is often radiation therapy, the field of which may include parts of the brachial plexus. It can be difficult in such situations to determine if a new brachial or lumbosacral plexopathy is related to tumor within the plexus or from radiation-induced nerve damage. Radiation can be associated with

microvascular abnormalities and fibrosis of surrounding tissues, which can damage the axons and the Schwann cells. Radiation-induced plexopathy can develop months or years following therapy and is dose dependent.

Tumor invasion is usually painful and more commonly affects the lower trunk, whereas radiation injury is often painless and affects the upper trunk. Imaging studies such as MRI and CT scans are useful but can be misleading with small microscopic invasion of the plexus. EMG can be informative if myokymic discharges are appreciated, as this finding strongly suggests radiation-induced damage.

EVALUATION AND TREATMENT OF PLEXOPATHIES

Most patients with plexopathies will undergo both imaging with MRI and EDx evaluations. Severe pain from acute idiopathic lumbosacral plexopathy may respond to a short course of glucocorticoids.

460 Guillain-Barré Syndrome and Other Immune-Mediated Neuropathies

Stephen L. Hauser, Anthony A. Amato

GUILLAIN-BARRÉ SYNDROME

Guillain-Barré syndrome (GBS) is an acute, frequently severe, and fulminant polyradiculoneuropathy that is autoimmune in nature. It occurs year-round at a rate of between 1 and 4 cases per 100,000 annually; in the United States, ~5000–6000 cases occur per year. Males are at slightly higher risk for GBS than females, and in Western countries, adults are more frequently affected than children.

Clinical Manifestations GBS manifests as a rapidly evolving areflexic motor paralysis with or without sensory disturbance. The usual pattern is an ascending paralysis that may be first noticed as rubbery legs. Weakness typically evolves over hours to a few days and is frequently accompanied by tingling dysesthesias in the extremities. The legs are usually more affected than the arms, and facial diparesis is present in 50% of affected individuals. The lower cranial nerves are also frequently involved, causing bulbar weakness with difficulty handling secretions and maintaining an airway; the diagnosis in these patients may initially be mistaken for brainstem ischemia. Pain in the neck, shoulder, back, or diffusely over the spine is also common in the early stages of GBS, occurring in ~50% of patients. Most patients require hospitalization, and in different series, up to 30% require ventilatory assistance at some time during the illness. The need for mechanical ventilation is associated with more severe weakness on admission, a rapid tempo of progression, and the presence of facial and/or bulbar weakness during the first week of symptoms. Fever and constitutional symptoms are absent at the onset and, if present, cast doubt on the diagnosis. Deep tendon reflexes attenuate or disappear within the first few days of onset. Cutaneous sensory deficits (e.g., loss of pain and temperature sensation) are usually relatively mild, but functions subserved by large sensory fibers, such as deep tendon reflexes and proprioception, are more severely affected. Bladder dysfunction may occur in severe cases but is usually transient. If bladder dysfunction is a prominent feature and comes early in the course, diagnostic possibilities other than GBS should be considered, particularly spinal cord disease. Once clinical worsening stops and the patient reaches a plateau (almost always within 4 weeks of onset), further progression is unlikely.

Autonomic involvement is common and may occur even in patients whose GBS is otherwise mild. The usual manifestations are loss of vasomotor control with wide fluctuation in blood pressure, postural

TABLE 459-10 LUMBOSACRAL PLEXOPATHIES: ETIOLOGIES

- Retroperitoneal hematoma
- Psoas abscess
- Malignant neoplasm
- Benign neoplasm
- Radiation
- Amyloid
- Diabetic radiculoplexus neuropathy
- Idiopathic radiculoplexus neuropathy
- Sarcoidosis
- Aortic occlusion/surgery
- Lithotomy positioning
- Hip arthroplasty
- Pelvic fracture
- Obstetric injury

TABLE 460-1 SUBTYPES OF GUILLAIN-BARRÉ SYNDROME (GBS)

Subtype	Features	Electrodiagnosis	Pathology
Acute inflammatory demyelinating polyneuropathy (AIDP)	Adults affected more than children; 90% of cases in Western world; recovery rapid; anti-GM1 antibodies (<50%)	Demyelinating	First attack on Schwann cell surface; widespread myelin damage, macrophage activation, and lymphocytic infiltration; variable secondary axonal damage
Acute motor axonal neuropathy (AMAN)	Children and young adults; prevalent in China and Mexico; may be seasonal; recovery rapid; anti-GD1a antibodies	Axonal	First attack at motor nodes of Ranvier; macrophage activation, few lymphocytes, frequent periaxonal macrophages; extent of axonal damage highly variable
Acute motor sensory axonal neuropathy (AMSAN)	Mostly adults; uncommon; recovery slow, often incomplete; closely related to AMAN	Axonal	Same as AMAN, but also affects sensory nerves and roots; axonal damage usually severe
Miller Fisher syndrome (MFS)	Adults and children; ophthalmoplegia, ataxia, and areflexia; anti-GQ1b antibodies (90%)	Axonal or demyelinating	Few cases examined; resembles AIDP

hypotension, and cardiac dysrhythmias. These features require close monitoring and management and can be fatal. Pain is another common feature of GBS; in addition to the acute pain described above, a deep aching pain may be present in weakened muscles that patients liken to having overexercised the previous day. Other pains in GBS include dysesthetic pain in the extremities as a manifestation of sensory nerve fiber involvement. These pains are self-limited and often respond to standard analgesics (Chap. 18).

Several subtypes of GBS are recognized, as determined primarily by electrodiagnostic (Edx) and pathologic distinctions (Table 460-1). The most common variant is acute inflammatory demyelinating polyneuropathy (AIDP). Additionally, there are two axonal variants, which are often clinically severe—the acute motor axonal neuropathy (AMAN) and acute motor sensory axonal neuropathy (AMSAN) subtypes. In addition, a range of limited or regional GBS syndromes are also encountered. Notable among these is the Miller Fisher syndrome (MFS), which presents as rapidly evolving ataxia and areflexia of limbs without weakness, and ophthalmoplegia, often with pupillary paralysis. The MFS variant accounts for ~5% of all cases and is strongly associated with antibodies to the ganglioside GQ1b (see “Immunopathogenesis,” below). Other regional variants of GBS include (1) pure sensory forms; (2) ophthalmoplegia with anti-GQ1b antibodies as part of severe motor-sensory GBS; (3) GBS with severe bulbar and facial paralysis, sometimes associated with antecedent cytomegalovirus (CMV) infection and anti-GM2 antibodies; and (4) acute pandysautonomia (Chap. 454).

Antecedent Events Approximately 70% of cases of GBS occur 1–3 weeks after an acute infectious process, usually respiratory or gastrointestinal. Culture and seroepidemiologic techniques show that 20–30% of all cases occurring in North America, Europe, and Australia are preceded by infection or reinfection with *Campylobacter jejuni*. A similar proportion is preceded by a human herpes virus infection, often CMV or Epstein-Barr virus. Other viruses (e.g., HIV, hepatitis E) and also *Mycoplasma pneumoniae* have been identified as agents involved in antecedent infections, as have recent immunizations. The swine influenza vaccine, administered widely in the United States in 1976, is the most notable example. Influenza vaccines in use from 1992 to 1994, however, resulted in only one additional case of GBS per million persons vaccinated, and the more recent seasonal influenza vaccines appear to confer a GBS risk of <1 per million. Epidemiologic studies looking at H1N1 vaccination demonstrated at most only a slight increased risk of GBS. Meningococcal vaccinations (Menactra) does not appear to carry an increased risk. Older-type rabies vaccine, prepared in nervous system tissue, is implicated as a trigger of GBS in developing countries where it is still used; the mechanism is presumably immunization against neural antigens. GBS also occurs more frequently than can be attributed to chance alone in patients with lymphoma (including Hodgkin’s disease), in HIV-seropositive individuals, and in patients with systemic lupus erythematosus (SLE). *C. jejuni* has also been implicated in summer

outbreaks of AMAN among children and young adults exposed to chickens in rural China.

Immunopathogenesis Several lines of evidence support an autoimmune basis for acute inflammatory demyelinating polyneuropathy (AIDP), the most common and best-studied type of GBS; the concept extends to all of the subtypes of GBS (Table 460-1).

It is likely that both cellular and humoral immune mechanisms contribute to tissue damage in AIDP. T cell activation is suggested by the finding that elevated levels of cytokines and cytokine receptors are present in serum (interleukin [IL] 2, soluble IL-2 receptor) and in cerebrospinal fluid (CSF) (IL-6, tumor necrosis factor α , interferon γ). AIDP is also closely analogous to an experimental T cell-mediated immunopathy designated *experimental allergic neuritis* (EAN). EAN is induced in laboratory animals by immune sensitization against protein fragments derived from peripheral nerve proteins, and in particular against the P2 protein. Based on analogy to EAN, it was initially thought that AIDP was likely to be primarily a T cell-mediated disorder; however, abundant data now suggest that autoantibodies directed against nonprotein determinants may be central to many cases.

Circumstantial evidence suggests that all GBS results from immune responses to nonself antigens (infectious agents, vaccines) that misdirect to host nerve tissue through a resemblance-of-epitope (molecular mimicry) mechanism (Fig. 460-1). The neural targets are likely to be glycoconjugates, specifically gangliosides (Table 460-2; Fig. 460-2). Gangliosides are complex glycosphingolipids that contain one or more sialic acid residues; various gangliosides participate in cell-cell interactions (including those between axons and glia), modulation of receptors, and regulation of growth. They are typically exposed on the plasma membrane of cells, rendering them susceptible to an antibody-mediated attack. Gangliosides and other glycoconjugates are present in large quantity in human nervous tissues and in key sites, such as nodes of Ranvier. Antiganglioside antibodies, most frequently to GM1, are common in GBS (20–50% of cases), particularly in AMAN and AMSAN and in those cases preceded by *C. jejuni* infection. Furthermore, isolates of *C. jejuni* from stool cultures of patients with GBS have surface glycolipid structures that antigenically cross react with gangliosides, including GM1, concentrated in human nerves. Sialic acid residues from pathogenic *C. jejuni* strains can also trigger activation of dendritic cells via signaling through a toll-like receptor (TLR4), promoting B cell differentiation and further amplifying humoral autoimmunity. Another line of evidence is derived from experience in Europe with parenteral use of purified bovine brain gangliosides for treatment of various neuropathic disorders. Between 5 and 15 days after injection, some recipients developed acute motor axonal GBS with high titers of anti-GM1 antibodies that recognized epitopes at nodes of Ranvier and motor endplates. Experimentally, anti-GM1 antibodies can trigger complement-mediated injury at paranodal axon-glia junctions, disrupting the clustering of sodium channels and likely contributing to conduction block (see “Pathophysiology,” below).

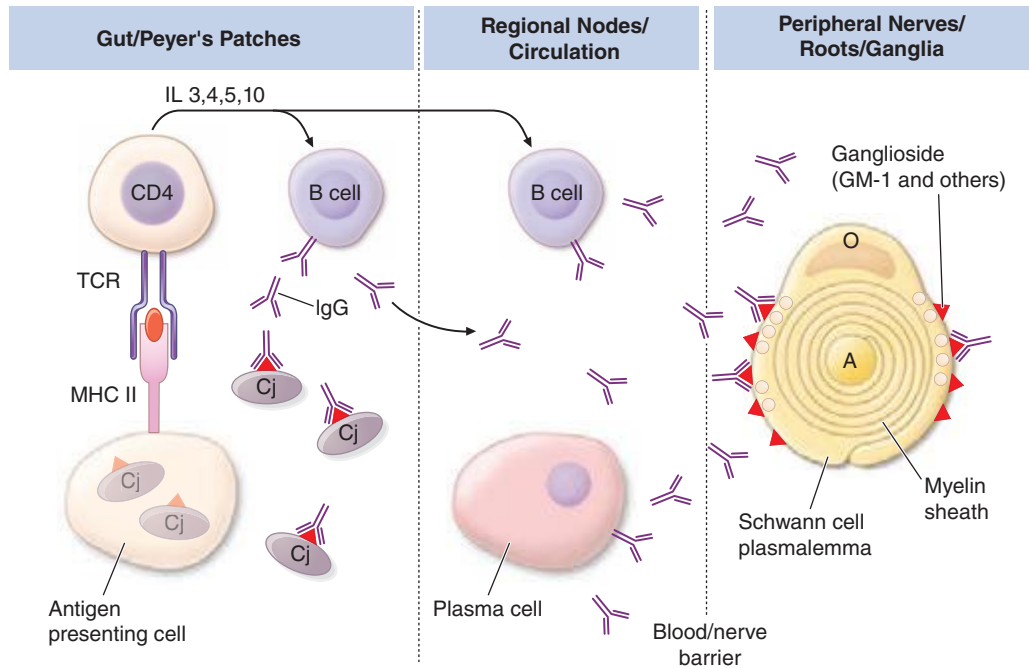


FIGURE 460-1 Postulated immunopathogenesis of Guillain-Barré syndrome (GBS) associated with *Campylobacter jejuni* infection. B cells recognize glycoconjugates on *C. jejuni* (Cj) (triangles) that cross-react with ganglioside present on Schwann cell surface and subjacent peripheral nerve myelin. Some B cells, activated via a T cell–independent mechanism, secrete primarily IgM (not shown). Other B cells (upper left side) are activated via a partially T cell–dependent route and secrete primarily IgG; T cell help is provided by CD4 cells activated locally by fragments of Cj proteins that are presented on the surface of antigen-presenting cells (APCs). A critical event in the development of GBS is the escape of activated B cells from Peyer’s patches into regional lymph nodes. Activated T cells probably also function to assist in opening of the blood-nerve barrier, facilitating penetration of pathogenic autoantibodies. The earliest changes in myelin (right) consist of edema between myelin lamellae and vesicular disruption (shown as circular blebs) of the outermost myelin layers. These effects are associated with activation of the C5b-C9 membrane attack complex and probably mediated by calcium entry; it is possible that the macrophage cytokine tumor necrosis factor (TNF) also participates in myelin damage. A, axon; B, B cell; MHC II, class II major histocompatibility complex molecule; O, oligodendrocyte; TCR, T cell receptor.

TABLE 460-2 PRINCIPAL ANTIGLYCOLIPID ANTIBODIES IMPLICATED IN IMMUNE NEUROPATHIES		
Clinical Presentation	Antibody Target	Usual Isotype
Acute Immune Neuropathies (Guillain-Barré Syndrome)		
Acute inflammatory demyelinating polyneuropathy (AIDP)	No clear patterns	IgG (polyclonal)
Acute motor axonal neuropathy (AMAN)	GM1 most common GD1a, GM1, GM1b, GalNAc–GD1a (<50% for any)	IgG (polyclonal)
Miller Fisher syndrome (MFS)	GQ1b (>90%)	IgG (polyclonal)
Acute pharyngeal cervicobrachial neuropathy (APCBN)	GT1a (? most)	IgG (polyclonal)
Chronic Immune Neuropathies		
Chronic inflammatory demyelinating polyneuropathy (CIDP) (75%)	P0, myelin P2 protein, PMP22, neurofascin	No clear pattern
CIDP-M (MGUS associated) (25%)	Neural binding sites	IgG, IgA (monoclonal)
Chronic sensory > motor neuropathy	SPGP, SGLPG (on MAG) (50%) Uncertain (50%)	IgM (monoclonal)
Multifocal motor neuropathy (MMN)	GM1, GalNAc–GD1a, others (25–50%)	IgM (polyclonal, monoclonal)
Chronic sensory ataxic neuropathy	GD1b, GQ1b, and other b-series gangliosides	IgM (monoclonal)

Abbreviations: CIDP-M, CIDP with a monoclonal gammopathy; MAG, myelin-associated glycoprotein; MGUS, monoclonal gammopathy of undetermined significance.
Source: Modified from HJ Willison, N Yuki: Brain 125:2591, 2002.

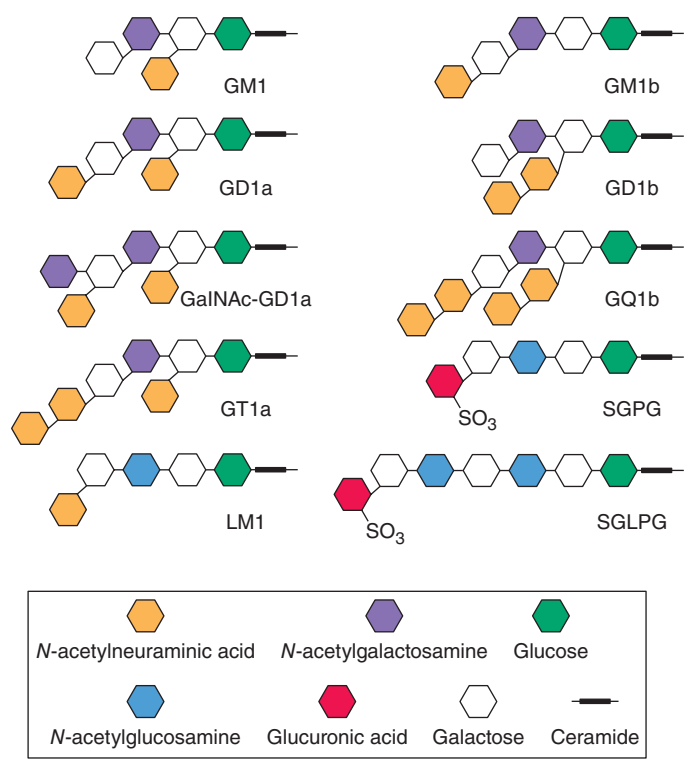


FIGURE 460-2 Glycolipids implicated as antigens in immune-mediated neuropathies. (Modified from HJ Willison, N Yuki: Brain 125: 2591, 2002.)

Anti-GQ1b IgG antibodies are found in >90% of patients with MFS (Table 460-2; Fig. 460-2), and titers of IgG are highest early in the course. Anti-GQ1b antibodies are not found in other forms of GBS unless there is extraocular motor nerve involvement. A possible explanation for this association is that extraocular motor nerves are enriched in GQ1b gangliosides in comparison to limb nerves. In addition, a monoclonal anti-GQ1b antibody raised against *C. jejuni* isolated from a patient with MFS blocked neuromuscular transmission experimentally.

Taken together, these observations provide strong but still inconclusive evidence that autoantibodies play an important pathogenic role in GBS. Although antiganglioside antibodies have been studied most intensively, other antigenic targets may also be important. One report identified IgG antibodies against Schwann cells and neurons (nerve growth cone region) in some GBS cases. Proof that these antibodies are pathogenic requires that they be capable of mediating disease following direct passive transfer to naïve hosts; this has not yet been demonstrated, although one case of possible maternal-fetal transplacental transfer of GBS has been described.

In AIDP, an early step in the induction of tissue damage appears to be complement deposition along the outer surface of the Schwann cell. Activation of complement initiates a characteristic vesicular disintegration of the myelin sheath and also leads to recruitment of activated macrophages, which participate in damage to myelin and axons. In AMAN, the pattern is different in that complement is deposited along with IgG at the nodes of Ranvier along large motor axons. Interestingly, in cases of AMAN, antibodies against GD1a appear to have a fine specificity that favors binding to motor rather than sensory nerve roots, even though this ganglioside is expressed on both fiber types.

Pathophysiology In the demyelinating forms of GBS, the basis for flaccid paralysis and sensory disturbance is conduction block. This finding, demonstrable electrophysiologically, implies that the axonal connections remain intact. Hence, recovery can take place rapidly as remyelination occurs. In severe cases of demyelinating GBS, secondary axonal degeneration usually occurs; its extent can be estimated electrophysiologically. More secondary axonal degeneration correlates with a slower rate of recovery and a greater degree of residual disability. When a severe primary axonal pattern is encountered electrophysiologically, the implication is that axons have degenerated and become disconnected from their targets, specifically the neuromuscular junctions, and must therefore regenerate for recovery to take place. In motor axonal cases in which recovery is rapid, the lesion is thought to be localized to preterminal motor branches, allowing regeneration and reinnervation to take place quickly. Alternatively, in mild cases, collateral sprouting and reinnervation from surviving motor axons near the neuromuscular junction may begin to reestablish physiologic continuity with muscle cells over a period of several months.

Laboratory Features CSF findings are distinctive, consisting of an elevated CSF protein level (1–10 g/L [100–1000 mg/dL]) without accompanying pleocytosis. The CSF is often normal when symptoms have been present for ≤48 h; by the end of the first week, the level of protein is usually elevated. A transient increase in the CSF white cell count (10–100/μL) occurs on occasion in otherwise typical GBS; however, a sustained CSF pleocytosis suggests an alternative diagnosis (viral myelitis) or a concurrent diagnosis such as unrecognized HIV infection, leukemia or lymphoma with infiltration of nerves, or neurosarcoidosis. Edx features are mild or absent in the early stages of GBS and lag behind the clinical evolution. In AIDP, the earliest features are prolonged F-wave latencies, prolonged distal latencies, and reduced amplitudes of compound muscle action potentials (CMAPs), probably owing to the predilection for involvement of nerve roots and distal motor nerve terminals early in the course. Later, slowing of conduction velocity, conduction block, and temporal dispersion may be appreciated (Table 460-1). Occasionally, sensory nerve action potentials (SNAPs) may be normal in the feet (e.g., sural nerve) when abnormal in the arms. This is also a sign that the patient does not have one of the more typical “length-dependent” polyneuropathies. In cases with

primary axonal pathology, the principal Edx finding is reduced amplitude of CMAPs (and also SNAPs with AMSAN) without conduction slowing or prolongation of distal latencies.

Diagnosis GBS is a descriptive entity. The diagnosis of AIDP is made by recognizing the pattern of rapidly evolving paralysis with areflexia, absence of fever or other systemic symptoms, and characteristic antecedent events. In 2011, the Brighton Collaboration developed a new set of case definitions for GBS in response to needs of epidemiologic studies of vaccination and assessing risks of GBS (Table 460-3). These criteria have subsequently been validated. Other disorders that may enter into the differential diagnosis include acute myelopathies (especially with prolonged back pain and sphincter disturbances); diphtheria (early oropharyngeal disturbances); Lyme polyradiculitis and other tick-borne paralyzes; porphyria (abdominal pain, seizures, psychosis); vasculitic neuropathy (check erythrocyte sedimentation rate, described below); poliomyelitis (fever and meningismus common); West Nile virus; CMV polyradiculitis (in immunocompromised patients); critical illness neuropathy or myopathy; neuromuscular junction disorders such as myasthenia gravis and botulism (pupillary reactivity lost early); poisonings with organophosphates, thallium, or arsenic; paralytic shellfish poisoning; or severe hypophosphatemia (rare). Laboratory tests are helpful primarily to exclude mimics of GBS. Edx features may be minimal, and the CSF protein level may not rise until the end of the first week. If the diagnosis is strongly suspected, treatment should be initiated without waiting for evolution of the characteristic Edx and CSF findings to occur. GBS patients with risk factors for HIV or with CSF pleocytosis should have a serologic test for HIV.

TREATMENT GUILLAIN-BARRÉ SYNDROME

In the vast majority of patients with GBS, treatment should be initiated as soon after diagnosis as possible. Each day counts; ~2 weeks after the first motor symptoms, it is not known whether immunotherapy is still effective. If the patient has already reached the plateau stage, then treatment probably is no longer indicated, unless the patient has severe motor weakness and one cannot exclude the possibility that an immunologic attack is still ongoing. Either high-dose intravenous immune globulin (IVIg) or plasmapheresis can be initiated, as they are equally effective for typical GBS. A combination of the two therapies is not significantly better than either alone. IVIg is often the initial therapy chosen because of its ease of administration and good safety record. Anecdotal data have also suggested that IVIg may be preferable to plasma exchange (PE) for the AMAN and MFS variants of GBS. IVIg is administered as five daily infusions for a total dose of 2 g/kg body weight. There is some evidence that GBS autoantibodies are neutralized by anti-idiotypic antibodies present in IVIg preparations, perhaps accounting for the therapeutic effect. A course of plasmapheresis usually consists of ~40–50 mL/kg PE four to five times over a week. Meta-analysis of randomized clinical trials indicates that treatment reduces the need for mechanical ventilation by nearly half (from 27% to 14% with PE) and increases the likelihood of full recovery at 1 year (from 55% to 68%). Functionally significant improvement may occur toward the end of the first week of treatment or may be delayed for several weeks. The lack of noticeable improvement following a course of IVIg or PE is not an indication to treat with the alternate treatment. However, there are occasional patients who are treated early in the course of GBS and improve, who then relapse within a month. Brief retreatment with the original therapy is usually effective in such cases. Glucocorticoids have not been found to be effective in GBS. Occasional patients with very mild forms of GBS, especially those who appear to have already reached a plateau when initially seen, may be managed conservatively without IVIg or PE.

In the worsening phase of GBS, most patients require monitoring in a critical care setting, with particular attention to vital capacity, heart rhythm, blood pressure, nutrition, deep vein thrombosis prophylaxis, cardiovascular status, early consideration (after 2 weeks

Clinical case definitions for diagnosis of GBS*Level 1 of diagnostic certainty*

Bilateral AND flaccid weakness of the limbs

AND

Decreased or absent deep tendon reflexes in weak limbs

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

Electrophysiologic findings consistent with GBS

AND

Cytoalbuminologic dissociation (i.e., elevation of CSF protein level above laboratory normal value AND CSF total white cell count <50 cells/ μ L)

AND

Absence of an identified alternative diagnosis for weakness

Level 2 of diagnostic certainty

Bilateral AND flaccid weakness of the limbs

AND

Decreased or absent deep tendon reflexes in weak limbs

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

CSF total white cell count <50 cells/ μ L (with or without CSF protein elevation above laboratory normal value)

OR

If CSF not collected or results not available, electrophysiologic studies consistent with GBS

AND

Absence of identified alternative diagnosis for weakness

Level 3 of diagnostic certainty

Bilateral and flaccid weakness of the limbs

AND

Decreased or absent deep tendon reflexes in weak limbs

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

Absence of identified alternative diagnosis for weakness

Clinical case definitions for diagnosis of Miller Fisher syndrome*Level 1 of diagnostic certainty*

Bilateral ophthalmoparesis and bilateral reduced or absent tendon reflexes, and ataxia

AND

Absence of limb weakness

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

Cytoalbuminologic dissociation (i.e., elevation of cerebrospinal protein above the laboratory normal and total CSF white cell count <50 cells/ μ L)

AND

Nerve conduction studies are normal, OR indicate involvement of sensory nerves only

AND

No alterations in consciousness or corticospinal tract signs

AND

Absence of identified alternative diagnosis

Level 2 of diagnostic certainty

Bilateral ophthalmoparesis and bilateral reduced or absent tendon reflexes and ataxia

AND

Absence of limb weakness

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

CSF with a total white cell count <50 cells/ μ L (with or without CSF protein elevation above laboratory normal value)

OR

Nerve conduction studies are normal, OR indicate involvement of sensory nerves only

AND

No alterations in consciousness or corticospinal tract signs

AND

Absence of identified alternative diagnosis

Level 3 of diagnostic certainty

Bilateral ophthalmoparesis and bilateral reduced or absent tendon reflexes and ataxia

AND

Absence of limb weakness

AND

Monophasic illness pattern and interval between onset and nadir of weakness between 12 h and 28 days and subsequent clinical plateau

AND

No alterations in consciousness or corticospinal tract signs

AND

Absence of identified alternative diagnosis

Abbreviation: CSF, cerebrospinal fluid.**Source:** From JJ Sejvar et al: Guillain-Barré syndrome and Fisher syndrome: case definitions and guidelines for collection, analysis, and presentation of immunization safety data. *Vaccine* 29:599, 2011. Validation study published by C Fokke et al: Diagnosis of Guillain-Barré syndrome and validation of Brighton criteria. *Brain* 137:33, 2014.

of intubation) of tracheotomy, and chest physiotherapy. As noted, ~30% of patients with GBS require ventilatory assistance, sometimes for prolonged periods of time (several weeks or longer). Frequent turning and assiduous skin care are important, as are daily range-of-motion exercises to avoid joint contractures and daily reassurance as to the generally good outlook for recovery.

Prognosis and Recovery Approximately 85% of patients with GBS achieve a full functional recovery within several months to a year, although minor findings on examination (such as areflexia) may persist and patients often complain of continued symptoms, including fatigue. The mortality rate is <5% in optimal settings; death usually results from secondary pulmonary complications. The outlook

is worst in patients with severe proximal motor and sensory axonal damage. Such axonal damage may be either primary or secondary in nature (see "Pathophysiology," above), but in either case successful regeneration cannot occur. Other factors that worsen the outlook for recovery are advanced age, a fulminant or severe attack, and a delay in the onset of treatment. Between 5 and 10% of patients with typical GBS have one or more late relapses; such cases are then classified as chronic inflammatory demyelinating polyneuropathy (CIDP).

CHRONIC INFLAMMATORY DEMYELINATING POLYNEUROPATHY

CIDP is distinguished from GBS by its chronic course. In other respects, this neuropathy shares many features with the common demyelinating form of GBS, including elevated CSF protein levels

and the Edx findings of acquired demyelination. Most cases occur in adults, and males are affected slightly more often than females. The incidence of CIDP is lower than that of GBS, but due to the protracted course, the prevalence is greater.

Clinical Manifestations Onset is usually gradual over a few months or longer, but in a few cases, the initial attack is indistinguishable from that of GBS. An acute-onset form of CIDP should be considered when GBS deteriorates >9 weeks after onset or relapses at least three times. Symptoms are both motor and sensory in most cases. Weakness of the limbs is usually symmetric but can be strikingly asymmetric in multifocal acquired demyelinating sensory and motor (MADSAM) neuropathy variant (Lewis-Sumner syndrome) in which discrete peripheral nerves are involved. There is considerable variability from case to case. Some patients experience a chronic progressive course, whereas others, usually younger patients, have a relapsing and remitting course. Some have only motor findings, and a small proportion present with a relatively pure syndrome of sensory ataxia. Tremor occurs in ~10% and may become more prominent during periods of subacute worsening or improvement. A small proportion have cranial nerve findings, including external ophthalmoplegia. CIDP tends to ameliorate over time with treatment; the result is that many years after onset, nearly 75% of patients have reasonable functional status. Death from CIDP is uncommon.

Diagnosis The diagnosis rests on characteristic clinical, CSF, and electrophysiologic findings. The CSF is usually acellular with an elevated protein level, sometimes several times normal. As with GBS, a CSF pleocytosis should lead to the consideration of HIV infection, leukemia or lymphoma, and neurosarcoidosis. Edx findings reveal variable degrees of conduction slowing, prolonged distal latencies, distal and temporal dispersion of CMAPs, and conduction block as the principal features. In particular, the presence of conduction block is a certain sign of an acquired demyelinating process. Evidence of axonal loss, presumably secondary to demyelination, is present in >50% of patients. Serum protein electrophoresis with immunofixation is indicated to search for monoclonal gammopathy and associated conditions (see “Monoclonal Gammopathy of Undetermined Significance,” below). In all patients with presumptive CIDP, it is also reasonable to exclude vasculitis, collagen vascular disease (especially SLE), chronic hepatitis, HIV infection, amyloidosis, and diabetes mellitus. Other associated conditions include inflammatory bowel disease and lymphoma.

Pathogenesis Although there is evidence of immune activation in CIDP, the precise mechanisms of pathogenesis are unknown. Biopsy typically reveals little inflammation and onion-bulb changes (imbricated layers of attenuated Schwann cell processes surrounding an axon) that result from recurrent demyelination and remyelination (Fig. 460-1). The response to therapy suggests that CIDP is immune-mediated; CIDP responds to glucocorticoids, whereas GBS does not. Passive transfer of demyelination into experimental animals has been accomplished using IgG purified from the serum of some patients with CIDP, lending support for a humoral autoimmune pathogenesis. A minority of patients have serum antibodies against P0, myelin P2 protein, PMP22, or neurofascin. It is also of interest that a CIDP-like illness developed spontaneously in the nonobese diabetic (NOD) mouse when the immune co-stimulatory molecule B7-2 (CD86) was genetically deleted; this suggests that CIDP can result from altered triggering of T cells by antigen-presenting cells.

Approximately 25% of patients with clinical features of CIDP also have a monoclonal gammopathy of undetermined significance (MGUS). Cases associated with monoclonal IgA or IgG kappa usually respond to treatment as favorably as cases without a monoclonal gammopathy. Patients with IgM monoclonal gammopathy and antibodies directed against myelin-associated glycoprotein (MAG) have a distinct polyneuropathy, tend to have more sensory findings and a more protracted course, and usually have a less satisfactory response to treatment.

TREATMENT CHRONIC INFLAMMATORY DEMYELINATING POLYNEUROPATHY

Most authorities initiate treatment for CIDP when progression is rapid or walking is compromised. If the disorder is mild, management can be expectant, awaiting spontaneous remission. Controlled studies have shown that high-dose IVIg, PE, and glucocorticoids are all more effective than placebo. Initial therapy is usually with IVIg, administered as 2.0 g/kg body weight given in divided doses over 2–5 days; three monthly courses are generally recommended before concluding a patient is a treatment failure. If the patient responds, the infusion intervals can be gradually increased or the dosage decreased (e.g., 1 g/kg per month). PE, which appears to be as effective as IVIg, is initiated at two to three treatments per week for 6 weeks; periodic re-treatment may also be required. Treatment with glucocorticoids is another option (60–80 mg prednisone PO daily for 1–2 months, followed by a gradual dose reduction of 10 mg per month as tolerated), but long-term adverse effects including bone demineralization, gastrointestinal bleeding, and cushingoid changes are problematic. As many as one-third of patients with CIDP fail to respond adequately to the initial therapy chosen; a different treatment should then be tried. Patients who fail therapy with IVIg, PE, and glucocorticoids may benefit from treatment with immunosuppressive agents such as azathioprine, methotrexate, cyclosporine, and cyclophosphamide, either alone or as adjunctive therapy. Early experience with anti-CD20 (rituximab) has also shown promise. Use of these therapies requires periodic reassessment of their risks and benefits. In patients with a CIDP-like neuropathy who fail to respond to treatment, it is important to evaluate for POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, monoclonal gammopathy, skin changes; see below).

MULTIFOCAL MOTOR NEUROPATHY

Multifocal motor neuropathy (MMN) is a distinctive but uncommon neuropathy that presents as slowly progressive motor weakness and atrophy evolving over years in the distribution of selected nerve trunks, associated with sites of persistent focal motor conduction block in the same nerve trunks. Sensory fibers are relatively spared. The arms are affected more frequently than the legs, and >75% of all patients are male. Some cases have been confused with lower motor neuron forms of amyotrophic lateral sclerosis (Chap. 452). Less than 50% of patients present with high titers of polyclonal IgM antibody to the ganglioside GM1. It is uncertain how this finding relates to the discrete foci of persistent motor conduction block, but high concentrations of GM1 gangliosides are normal constituents of nodes of Ranvier in peripheral nerve fibers. Pathology reveals demyelination and mild inflammatory changes at the sites of conduction block.

Most patients with MMN respond to high-dose IVIg (dosages as for CIDP, above); periodic re-treatment is required (usually at least monthly) to maintain the benefit. Some refractory patients have responded to rituximab or cyclophosphamide. Glucocorticoids and PE are not effective.

NEUROPATHIES WITH MONOCLONAL GAMMOPATHY

MULTIPLE MYELOMA

Clinically overt polyneuropathy occurs in ~5% of patients with the commonly encountered type of multiple myeloma, which exhibits either lytic or diffuse osteoporotic bone lesions. These neuropathies are sensorimotor, are usually mild and slowly progressive but may be severe, and generally do not reverse with successful suppression of the myeloma. In most cases, Edx and pathologic features are consistent with a process of axonal degeneration.

In contrast, myeloma with osteosclerotic features, although representing only 3% of all myelomas, is associated with polyneuropathy in one-half of cases. These neuropathies, which may also occur with solitary plasmacytoma, are distinct because they (1) are usually

demyelinating in nature and resemble CIDP; (2) often respond to radiation therapy or removal of the primary lesion; (3) are associated with different monoclonal proteins and light chains (almost always lambda as opposed to primarily kappa in the lytic type of multiple myeloma); (4) are typically refractory to standard treatments of CIDP; and (5) may occur in association with other systemic findings including thickening of the skin, hyperpigmentation, hypertrichosis, organomegaly, endocrinopathy, anasarca, and clubbing of fingers. These are features of the POEMS syndrome (*polyneuropathy, organomegaly, endocrinopathy, monoclonal gammopathy, and skin changes*). Levels of vascular endothelial growth factor (VEGF) are increased in the serum, and this factor is felt to somehow play a pathogenic role in this syndrome. Treatment of the neuropathy is best directed at the osteosclerotic myeloma using surgery, radiotherapy, chemotherapy, or autologous peripheral blood stem cell transplantation.

Neuropathies are also encountered in other systemic conditions with gammopathy, including Waldenström's macroglobulinemia, primary systemic amyloidosis, and cryoglobulinemic states (mixed essential cryoglobulinemia, some cases of hepatitis C).

MONOCLONAL GAMMOPATHY OF UNDETERMINED SIGNIFICANCE

Chronic polyneuropathies occurring in association with MGUS are usually associated with the immunoglobulin isotypes IgG, IgA, and IgM. Most patients present with isolated sensory symptoms in their distal extremities and have Edx features of an axonal sensory or sensorimotor polyneuropathy. These patients otherwise resemble idiopathic sensory polyneuropathy, and the MGUS might just be coincidental. They usually do not respond to immunotherapies designed to reduce the concentration of the monoclonal protein. Some patients, however, present with generalized weakness and sensory loss and Edx studies indistinguishable from CIDP without monoclonal gammopathy (see "Chronic Inflammatory Demyelinating Polyneuropathy," above), and their response to immunosuppressive agents is also similar. An exception is the syndrome of IgM kappa monoclonal gammopathy associated with an indolent, longstanding, sometimes static sensory neuropathy, frequently with tremor and sensory ataxia. Most patients are male and older than age 50 years. In the majority, the monoclonal IgM immunoglobulin binds to a normal peripheral nerve constituent, MAG, found in the paranodal regions of Schwann cells. Binding appears to be specific for a polysaccharide epitope that is also found in other normal peripheral nerve myelin glycoproteins, P0 and PMP22, and also in other normal nerve-related glycosphingolipids (Fig. 460-1). In the MAG-positive cases, IgM paraprotein is incorporated into the myelin sheaths of affected patients and widens the spacing of the myelin lamellae, thus producing a distinctive ultrastructural pattern. Demyelination and remyelination are the hallmarks of the lesions, but axonal loss develops over time. These anti-MAG polyneuropathies are typical refractory to immunotherapy. In a small proportion of patients (30% at 10 years), MGUS will in time evolve into frankly malignant conditions such as multiple myeloma or lymphoma.

VASCULITIC NEUROPATHY

Peripheral nerve involvement is common in polyarteritis nodosa (PAN), appearing in half of all cases clinically and in 100% of cases at postmortem studies (Chap. 385). The most common pattern is multifocal (asymmetric) motor-sensory neuropathy (mononeuropathy multiplex) due to ischemic lesions of nerve trunks and roots; however, some cases of vasculitic neuropathy present as a distal, symmetric sensorimotor polyneuropathy. Symptoms of neuropathy are a common presenting complaint in patients with PAN. The Edx findings are those of an axonal process. Small- to medium-sized arteries of the vasa nervorum, particularly the epineural vessels, are affected in PAN, resulting in a widespread ischemic neuropathy. A high frequency of neuropathy occurs in allergic angiitis and granulomatosis (Churg-Strauss syndrome [CSS]).

Systemic vasculitis should always be considered when a subacute or chronically evolving mononeuropathy multiplex occurs in conjunction with constitutional symptoms (fever, anorexia, weight loss, loss of energy, malaise, and nonspecific pains). Diagnosis of suspected vasculitic neuropathy is made by a combined nerve and muscle biopsy, with serial section or skip-serial techniques.

Approximately one-third of biopsy-proven cases of vasculitic neuropathy are "nonsystemic" in that the vasculitis appears to affect only peripheral nerves. Constitutional symptoms are absent, and the course is more indolent than that of PAN. The erythrocyte sedimentation rate may be elevated, but other tests for systemic disease are negative. Nevertheless, clinically silent involvement of other organs is likely, and vasculitis is frequently found in muscle biopsied at the same time as nerve.

Vasculitic neuropathy may also be seen as part of the vasculitis syndrome occurring in the course of other connective tissue disorders (Chap. 385). The most frequent is rheumatoid arthritis, but ischemic neuropathy due to involvement of vasa nervorum may also occur in mixed cryoglobulinemia, Sjögren's syndrome, granulomatosis with polyangiitis (Wegener's), hypersensitivity angiitis, systemic lupus erythematosus, and progressive systemic sclerosis.

Some vasculitides are associated with antineutrophil cytoplasmic antibodies (ANCA), which in turn, are subclassified as cytoplasmic (cANCA) or perinuclear (pANCA). cANCAs are directed against proteinase 3 (PR3), whereas pANCAs target myeloperoxidase (MPO). PR3/cANCAs are associated with granulomatosis with polyangiitis (Wegener's), whereas MPO/pANCAs are typically associated with microscopic polyangiitis, CSS, and less commonly PAN. Of note, MPO/pANCA has also been seen in minocycline-induced vasculitis.

Management of these neuropathies, including the "nonsystemic" vasculitic neuropathy, consists of treatment of the underlying condition as well as the aggressive use of glucocorticoids and cyclophosphamide. Use of these immunosuppressive agents has resulted in dramatic improvements in outcome, with 5-year survival rates now greater than 80%. Recent clinical trials found that the combination of rituximab and glucocorticoids is not inferior to cyclophosphamide and glucocorticoids. Thus, combination therapy with glucocorticoids and rituximab is increasingly recommended as the standard initial treatment, particularly for ANCA-associated vasculitis.

ANTI-Hu PARANEOPLASTIC NEUROPATHY (CHAP. 122)

This uncommon immune-mediated disorder manifests as a sensory neuronopathy (i.e., selective damage to sensory nerve bodies in dorsal root ganglia). The onset is often asymmetric with dysesthesias and sensory loss in the limbs that soon progress to affect all limbs, the torso, and the face. Marked sensory ataxia, pseudoathetosis, and inability to walk, stand, or even sit unsupported are frequent features and are secondary to the extensive deafferentation. Subacute sensory neuronopathy may be idiopathic, but more than half of cases are paraneoplastic, primarily related to lung cancer, and most of those are small-cell lung cancer (SCLC). Diagnosis of the underlying SCLC requires awareness of the association, testing for the paraneoplastic antibody, and often positron emission tomography (PET) scanning for the tumor. The target antigens are a family of RNA-binding proteins (HuD, HuC, and Hel-N1) that in normal tissues are only expressed by neurons. The same proteins are usually expressed by SCLC, triggering in some patients an immune response characterized by antibodies and cytotoxic T cells that cross-react with the Hu proteins of the dorsal root ganglion neurons, resulting in immune-mediated neuronal destruction. An encephalomyelitis may accompany the sensory neuronopathy and presumably has the same pathogenesis. Neurologic symptoms usually precede, by ≤ 6 months, the identification of SCLC. The sensory neuronopathy runs its course in a few weeks or months and stabilizes, leaving the patient disabled. Most cases are unresponsive to treatment with glucocorticoids, IVIg, PE, or immunosuppressant drugs.

461 Myasthenia Gravis and Other Diseases of the Neuromuscular Junction

Daniel B. Drachman, Anthony A. Amato

Myasthenia gravis (MG) is a neuromuscular disorder characterized by weakness and fatigability of skeletal muscles. The underlying defect is a decrease in the number of available acetylcholine receptors (AChRs) at neuromuscular junctions due to an antibody-mediated autoimmune attack. Treatment now available for MG is highly effective, although a specific cure has remained elusive.

PATHOPHYSIOLOGY

At the neuromuscular junction (Fig. 461-1, Video 461-1), acetylcholine (ACh) is synthesized in the motor nerve terminal and stored in vesicles (quanta). When an action potential travels down a motor nerve and reaches the nerve terminal, ACh from 150 to 200 vesicles is released and combines with AChRs that are densely packed at the peaks of postsynaptic folds. The AChR consists of five subunits (2α , 1β , 1δ , and 1γ or ϵ) arranged around a central pore. When ACh combines with the binding sites on the α subunits of the AChR, the channel in the AChR opens, permitting the rapid entry of cations, chiefly sodium, which produces depolarization at the end-plate region of the muscle fiber. If the depolarization is sufficiently large, it initiates an action potential that is propagated along the muscle fiber, triggering muscle contraction. This process is rapidly terminated by hydrolysis of ACh by acetylcholinesterase (AChE), which is present within the synaptic folds, and by diffusion of ACh away from the receptor.

In MG, the fundamental defect is a decrease in the number of available AChRs at the postsynaptic muscle membrane. In addition, the postsynaptic folds are flattened, or “simplified.” These changes result in decreased efficiency of neuromuscular transmission. Therefore, although ACh is released normally, it produces small end-plate potentials that may fail to trigger muscle action potentials. Failure of transmission at many neuromuscular junctions results in weakness of muscle contraction.

The amount of ACh released per impulse normally declines on repeated activity (termed *presynaptic rundown*). In the myasthenic

patient, the decreased efficiency of neuromuscular transmission combined with the normal rundown results in the activation of fewer and fewer muscle fibers by successive nerve impulses and hence increasing weakness, or *myasthenic fatigue*. This mechanism also accounts for the decremental response to repetitive nerve stimulation seen on electrodiagnostic testing.

The neuromuscular abnormalities in MG are caused by an autoimmune response mediated by specific anti-AChR antibodies. The anti-AChR antibodies reduce the number of available AChRs at neuromuscular junctions by three distinct mechanisms: (1) accelerated turnover of AChRs by a mechanism involving cross-linking and rapid endocytosis of the receptors; (2) damage to the postsynaptic muscle membrane by the antibody in collaboration with complement; and (3) blockade of the active site of the AChR, i.e., the site that normally binds ACh. An immune response to muscle-specific kinase (MuSK), a protein involved in AChR clustering at neuromuscular junctions, can also result in MG, with reduction of AChRs demonstrated experimentally. Anti-MuSK antibody occurs in about 40% of patients without AChR antibody. A small proportion of patients whose sera are negative for both AChR and MuSK antibodies have antibodies to another protein at the neuromuscular junction—low-density lipoprotein receptor-related protein 4 (lrp4)—that is important for clustering of AChRs. The pathogenic antibodies are IgG and are T cell dependent. Thus, immunotherapeutic strategies directed against either the antibody-producing B cells or helper T cells are effective in this antibody-mediated disease.

How the autoimmune response is initiated and maintained in MG is not completely understood, but the thymus appears to play a role in this process. The thymus is abnormal in ~75% of patients with AChR antibody-positive MG; in ~65% the thymus is “hyperplastic,” with the presence of active germinal centers detected histologically, although the hyperplastic thymus is not necessarily enlarged. An additional 10% of patients have thymic tumors (thymomas). Muscle-like cells within the thymus (myoid cells), which express AChRs on their surface, may serve as a source of autoantigen and trigger the autoimmune reaction within the thymus gland.

CLINICAL FEATURES

MG is not rare, having a prevalence as high as 2–7 in 10,000. It affects individuals in all age groups, but peaks of incidence occur in women in their twenties and thirties and in men in their fifties and sixties. Overall, women are affected more frequently than men, in a ratio of ~3:2. The cardinal features are *weakness* and *fatigability* of muscles.

The weakness increases during repeated use (fatigue) or late in the day and may improve following rest or sleep. The course of MG is often variable. Exacerbations and remissions may occur, particularly during the first few years after the onset of the disease. Remissions are rarely complete or permanent. Unrelated infections or systemic disorders can lead to increased myasthenic weakness and may precipitate “crisis” (see below).

The distribution of muscle weakness often has a characteristic pattern. The cranial muscles, particularly the lids and extraocular muscles, are typically involved early in the course of MG; diplopia and ptosis are common initial complaints. Facial weakness produces a “snarling” expression when the patient attempts to smile. Weakness in chewing is most noticeable after prolonged effort, as in chewing meat. Speech may have a nasal timbre caused by weakness of the palate or a dysarthric “mushy” quality due to tongue weakness. Difficulty in swallowing may occur as a result of weakness of the palate, tongue, or pharynx, giving rise to nasal regurgitation or aspiration of liquids or food. Bulbar weakness is especially prominent in MuSK antibody-positive MG. In ~85% of

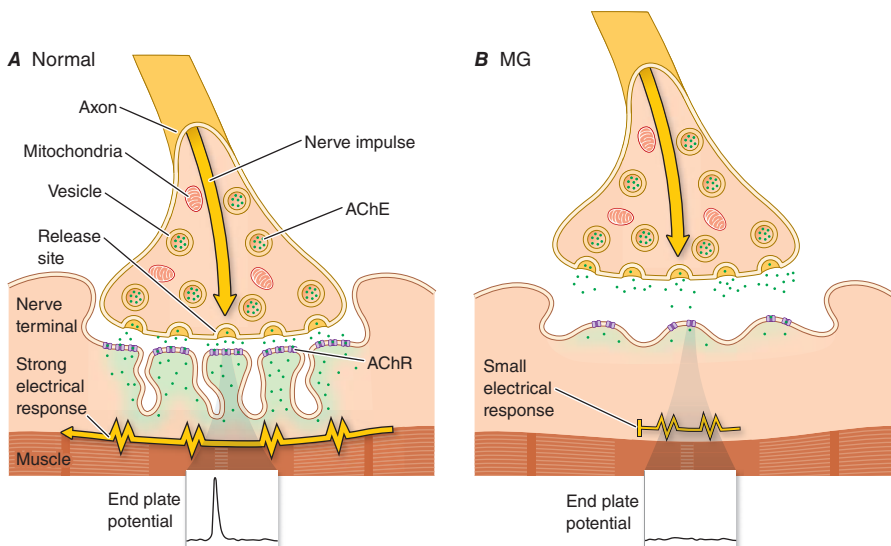


FIGURE 461-1 Diagrams of (A) normal and (B) myasthenic neuromuscular junctions.

AChE, acetylcholinesterase. See text for description of normal neuromuscular transmission. The myasthenia gravis (MG) junction demonstrates a normal nerve terminal; a reduced number of acetylcholine receptors (AChRs) (stippling); flattened, simplified postsynaptic folds; and a widened synaptic space. See Video 461-1 also. (Modified from DB Drachman: *N Engl J Med* 330:1797, 1994; with permission.)

History
Diplopia, ptosis, dysarthria, dysphagia, dyspnea
Weakness in characteristic distribution: proximal limbs, neck extensors, generalized
Fluctuation and fatigue: worse with repeated activity, improved by rest
Effects of previous treatments
Physical examination
Ptosis, diplopia
Motor power survey: quantitative testing of muscle strength
Forward arm abduction time (5 min)
Vital capacity measurement
Absence of other neurologic signs
Laboratory testing
Anti-AChR radioimmunoassay: ~85% positive in generalized MG; 50% in ocular MG; definite diagnosis if positive; negative result does not exclude MG; ~40% of AChR antibody–negative patients with generalized MG have anti-MuSK antibodies
Repetitive nerve stimulation: decrement of >15% at 3 Hz: highly probable
Single-fiber electromyography: blocking and jitter, with normal fiber density; confirmatory, but not specific
Edrophonium chloride (Enlon®) 2 mg + 8 mg IV; highly probable diagnosis if unequivocally positive
For ocular or cranial MG: exclude intracranial lesions by CT or MRI

Abbreviations: AChR, acetylcholine receptor; CT, computed tomography; MRI, magnetic resonance imaging; MuSK, muscle-specific tyrosine kinase.

patients, the weakness becomes generalized, affecting the limb muscles as well. If weakness remains restricted to the extraocular muscles for 3 years, it is likely that it will not become generalized, and these patients are said to have *ocular MG*. The limb weakness in MG is often proximal and may be asymmetric. Despite the muscle weakness, deep tendon reflexes are preserved. If weakness of respiration becomes so severe as to require respiratory assistance, the patient is said to be in *crisis*.

DIAGNOSIS AND EVALUATION

(Table 461-1) The diagnosis is suspected on the basis of weakness and fatigability in the typical distribution described above, without loss of reflexes or impairment of sensation or other neurologic function. The suspected diagnosis should always be confirmed definitively before treatment is undertaken; this is essential because (1) other treatable conditions may closely resemble MG and (2) the treatment of MG may involve surgery and the prolonged use of drugs with potentially adverse side effects.

Antibodies to AChR, MuSK, or lrp4 As noted above, anti-AChR antibodies are detectable in the serum of ~85% of all myasthenic patients but in only about 50% of patients with weakness confined to the ocular muscles. The presence of anti-AChR antibodies is virtually diagnostic of MG, but a negative test does not exclude the disease. The measured level of anti-AChR antibody does not correspond well with the severity of MG in different patients. However, in an individual patient, a treatment-induced fall in the antibody level often correlates with clinical improvement, whereas a rise in the level may occur with exacerbations. Antibodies to MuSK have been found to be present in ~40% of AChR antibody–negative patients with generalized MG, and their presence is a useful diagnostic test in these patients. MuSK antibodies are rarely present in AChR antibody–positive patients or in patients with MG limited to ocular muscles. These antibodies may interfere with clustering of AChRs at neuromuscular junctions, as MuSK is known to do during early development. A small proportion of MG patients without antibodies to AChR or MuSK may have antibodies to lrp4, although a test for lrp4 antibodies is not yet commercially available. Finally, antibodies against agrin have recently been found in some patients with MG. Agrin is a protein derived from motor nerves that normally binds to lrp4 and thus may also interfere with clustering of AChRs at neuromuscular junctions. There may well be other—yet undefined—antibodies that impair neuromuscular transmission.

Electrodiagnostic Testing Repetitive nerve stimulation may provide helpful diagnostic evidence of MG. Anti-AChE medication is stopped 6–24 h before testing. It is best to test weak muscles or proximal muscle groups. Electric shocks are delivered at a rate of two or three per second to the appropriate nerves, and action potentials are recorded from the muscles. In normal individuals, the amplitude of the evoked muscle action potentials does not change at these rates of stimulation. However, in myasthenic patients, there is a rapid reduction of >10–15% in the amplitude of the evoked responses.

Anticholinesterase Test Drugs that inhibit the enzyme AChE allow ACh to interact repeatedly with the limited number of AChRs in MG, producing improvement in muscle strength. Edrophonium is used most commonly for diagnostic testing because of the rapid onset (30 s) and short duration (~5 min) of its effect. An objective end point must be selected to evaluate the effect of edrophonium, such as weakness of extraocular muscles, impairment of speech, or the length of time that the patient can maintain the arms in forward abduction. An initial IV dose of 2 mg of edrophonium is given. If definite improvement occurs, the test is considered positive and is terminated. If there is no change, the patient is given an additional 8 mg IV. The dose is administered in two parts because some patients react to edrophonium with side effects such as nausea, diarrhea, salivation, fasciculations, and rarely with severe symptoms of syncope or bradycardia. Atropine (0.6 mg) should be drawn up in a syringe and ready for IV administration if these symptoms become troublesome. The edrophonium test is now reserved for patients with clinical findings that are suggestive of MG but who have negative antibody and electrodiagnostic test results. False-positive tests occur in occasional patients with other neurologic disorders, such as amyotrophic lateral sclerosis, and in placebo-reactors. False-negative or equivocal tests may also occur. In some cases, it is helpful to use a longer-acting drug such as neostigmine (15 mg PO), because this permits more time for detailed evaluation of strength.

Inherited Myasthenic Syndromes The congenital myasthenic syndromes (CMS) comprise a heterogeneous group of disorders of the neuromuscular junction that are not autoimmune but rather are due to genetic mutations in which virtually any component of the neuromuscular junction may be affected. Alterations in function of the presynaptic nerve terminal, in the various subunits of the AChR, AChE, or the other molecules involved in end-plate development or maintenance, have been identified in the different forms of CMS. These disorders share many of the clinical features of autoimmune MG, including weakness and fatigability of skeletal muscles, in some cases involving extraocular muscles (EOMs), lids, and proximal muscles, similar to the distribution in autoimmune MG. CMS should be suspected when symptoms of myasthenia have begun in infancy or childhood and AChR antibody tests are consistently negative. By far the most common genetic defects occur in the AChR or other postsynaptic molecules (67% in the Mayo Clinic series of 350 CMS patients), with about equal frequencies of abnormalities in AChE (13%) and the various maintenance molecules (DOK7, GFPT, etc.; ~14%). In the forms that involve the AChR, a wide variety of mutations have been identified in each of the subunits, but the ϵ subunit is affected in ~75% of these cases. In most of the recessively inherited forms of CMS, the mutations are heteroallelic; that is, different mutations affecting each of the two alleles are present. Features of the four most common forms of CMS are summarized in **Table 461-2**. Although clinical features and electrodiagnostic and pharmacologic tests may suggest the correct diagnosis, molecular analysis is required for precise elucidation of the defect; this may lead to helpful treatment as well as genetic counseling.

Differential Diagnosis Other conditions that cause weakness of the cranial and/or somatic musculature include the nonautoimmune CMS discussed above, drug-induced myasthenia, Lambert-Eaton myasthenic syndrome (LEMS), neurasthenia, hyperthyroidism (Graves' disease), botulism, intracranial mass lesions, oculopharyngeal dystrophy, and mitochondrial myopathy (Kearns-Sayre syndrome, progressive external ophthalmoplegia). Treatment with penicillamine (used for scleroderma or rheumatoid arthritis) may result in true autoimmune

TABLE 461-2 THE CONGENITAL MYASTHENIC SYNDROMES

Type	Clinical Features	Electrophysiology	Genetics	End-Plate Effects	Treatment
Slow channel	Most common; weak forearm extensors; onset second to third decade; variable severity	Repetitive muscle response on nerve stimulation; prolonged channel opening and MEPP duration	Autosomal dominant; α , β , ϵ ; AChR mutations	Excitotoxic end-plate myopathy; decreased AChRs; postsynaptic damage	Quinidine, fluoxetine: decrease end-plate damage; made worse by anti-AChE
Low-affinity fast channel	Onset early; moderately severe; ptosis, EOM involvement; weakness and fatigue	Brief and infrequent channel openings; opposite of slow channel syndrome	Autosomal recessive; may be heteroallelic	Normal end-plate structure	3,4-DAP; anti-AChE
Severe AChR deficiencies	Early onset; variable severity; fatigue; typical MG features	Decremental response to repetitive nerve stimulation; decreased MEPP amplitudes	Autosomal recessive; ϵ mutations most common; many different mutations	Increased length of end plates; variable synaptic folds	Anti-AChE; 3,4-DAP
AChE deficiency	Early onset; variable severity; scoliosis; may have normal EOM, absent pupillary responses	Decremental response to repetitive nerve stimulation	Mutant gene for AChE's collagen anchor (COLQ)	Small nerve terminals; degenerated junctional folds	Worse with anti-AChE drugs; use albuterol, ephedrine, 3,4-DAP

Abbreviations: AChE, acetylcholinesterase; AChR, acetylcholine receptor; EOM, extraocular muscles; MEPP, miniature end-plate potentials; MG, myasthenia gravis; 3,4-DAP, 3,4-diaminopyridine.

MG, but the weakness is usually mild, and recovery occurs within weeks or months after discontinuing its use. Aminoglycoside antibiotics or procainamide can cause exacerbation of weakness in myasthenic patients; very large doses can cause neuromuscular weakness in normal individuals.

LEMS is a presynaptic disorder of the neuromuscular junction that can cause weakness similar to that of MG. The proximal muscles of the lower limbs are most commonly affected, but other muscles may be involved as well. Cranial nerve findings, including ptosis of the eyelids and diplopia, occur in up to 70% of patients and resemble features of MG. However, the two conditions are usually readily distinguished, because patients with LEMS have depressed or absent reflexes and experience autonomic changes such as dry mouth and impotence. Nerve stimulation produces an initial low-amplitude response and, at low rates of repetitive stimulation (2–3 Hz), decremental responses like those of MG; however, at high rates (50 Hz), or following exercise, incremental responses occur. LEMS is caused by autoantibodies directed against P/Q-type calcium channels at the motor nerve terminals, which can be detected in ~85% of LEMS patients by radioimmunoassay. These autoantibodies result in impaired release of ACh from nerve terminals. Many patients with LEMS have an associated malignancy, most commonly small-cell carcinoma of the lung, which may express calcium channels that stimulate the autoimmune response. The diagnosis of LEMS may signal the presence of a tumor long before it would otherwise be detected, permitting early removal. Treatment of LEMS involves plasmapheresis and immunosuppression, as for MG. 3,4-Diaminopyridine (3,4-DAP) and pyridostigmine may also be symptomatically helpful. 3,4-DAP acts by blocking potassium channels, which results in prolonged depolarization of the motor nerve terminals and thus enhances ACh release. Pyridostigmine prolongs the action of ACh, allowing repeated interactions with AChRs.

Botulism (Chap. 178) is due to potent bacterial toxins produced by any of eight different strains of *Clostridium botulinum*. The toxins enzymatically cleave specific proteins essential for the release of ACh from the motor nerve terminal, thereby interfering with neuromuscular transmission. Most commonly, botulism is caused by ingestion of improperly prepared food containing toxin. Rarely, the nearly ubiquitous spores of *C. botulinum* may germinate in wounds. In infants, the spores may germinate in the gastrointestinal (GI) tract and release toxin, causing muscle weakness. Patients present with myasthenia-like bulbar weakness (e.g., diplopia, dysarthria, dysphagia) and lack sensory symptoms and signs. Weakness may generalize to the limbs and may result in respiratory failure. Reflexes are present early, but they may be diminished as the disease progresses. Mentation is normal. Autonomic findings include paralytic ileus, constipation, urinary retention, dilated or poorly reactive pupils, and dry mouth. The demonstration of toxin in serum by bioassay is definitive, but the results usually take a relatively long time to be completed and may be negative. Nerve

stimulation studies reveal findings of presynaptic neuromuscular blockade with reduced compound muscle action potentials (CMAPs) that increase in amplitude following high-frequency repetitive stimulation. Treatment includes ventilatory support and aggressive inpatient supportive care (e.g., nutrition, deep vein thrombosis prophylaxis) as needed. Antitoxin should be given as early as possible to be effective and can be obtained through the Centers for Disease Control and Prevention. A preventive vaccine is available for laboratory workers or other highly exposed individuals.

Neurasthenia is the historic term for a myasthenia-like fatigue syndrome without an organic basis. These patients may present with subjective symptoms of weakness and fatigue, but muscle testing usually reveals the “give-away weakness” characteristic of nonorganic disorders; the complaint of fatigue in these patients means tiredness or apathy rather than decreasing muscle power on repeated effort. Hyperthyroidism is readily diagnosed or excluded by tests of thyroid function, which should be carried out routinely in patients with suspected MG. Abnormalities of thyroid function (hyper- or hypothyroidism) may increase myasthenic weakness. Diplopia resembling that in MG may occasionally be due to an intracranial mass lesion that compresses nerves to the EOMs (e.g., sphenoid ridge meningioma), but magnetic resonance imaging (MRI) of the head and orbits usually reveals the lesion.

Progressive external ophthalmoplegia is a rare condition resulting in weakness of the EOMs, which may be accompanied by weakness of the proximal muscles of the limbs and other systemic features. Most patients with this condition have mitochondrial disorders that can be detected on muscle biopsy (Chap. 462e).

Search for Associated Conditions (Table 461-3) Myasthenic patients have an increased incidence of several associated disorders. Thymic abnormalities occur in ~75% of AChR antibody-positive patients, as noted above. Neoplastic change (thymoma) may produce enlargement of the thymus, which is detected by computed tomography (CT) scanning of the anterior mediastinum. A thymic shadow on CT scan may normally be present through young adulthood, but enlargement of the thymus in a patient age >40 years is highly suspicious of thymoma. Hyperthyroidism occurs in 3–8% of patients and may aggravate the myasthenic weakness. Thyroid function tests should be obtained in all patients with suspected MG. Because of the association of MG with other autoimmune disorders, blood tests for rheumatoid factor and antinuclear antibodies should also be carried out. Chronic infection of any kind can exacerbate MG and should be sought carefully. Finally, measurements of ventilatory function are valuable because of the frequency and seriousness of respiratory impairment in myasthenic patients.

Because of the side effects of glucocorticoids and other immunosuppressive agents used in the treatment of MG, a thorough medical

TABLE 461-3 DISORDERS ASSOCIATED WITH MYASTHENIA GRAVIS AND RECOMMENDED LABORATORY TESTS

Associated disorders

Disorders of the thymus: thymoma, hyperplasia

Other autoimmune disorders: Hashimoto's thyroiditis, Graves' disease, rheumatoid arthritis, lupus erythematosus, skin disorders, family history of autoimmune disorder

Disorders or circumstances that may exacerbate myasthenia gravis: hyperthyroidism or hypothyroidism, occult infection, medical treatment for other conditions (see Table 461-4)

Disorders that may interfere with therapy: tuberculosis, diabetes, peptic ulcer, gastrointestinal bleeding, renal disease, hypertension, asthma, osteoporosis, obesity

Recommended laboratory tests or procedures

CT or MRI of chest

Tests for lupus erythematosus, antinuclear antibody, rheumatoid factor, antithyroid antibodies

Thyroid function tests

PPD skin test

Fasting blood glucose, hemoglobin A1c

Pulmonary function tests

Bone densitometry

Abbreviations: CT, computed tomography; MRI, magnetic resonance imaging; PPD, purified protein derivative.

investigation should be undertaken, searching specifically for evidence of chronic or latent infection (such as tuberculosis or hepatitis), hypertension, diabetes, renal disease, and glaucoma.

TREATMENT MYASTHENIA GRAVIS

The prognosis has improved strikingly as a result of advances in treatment. Nearly all myasthenic patients can be returned to full productive lives with proper therapy. The most useful treatments for MG include anticholinesterase medications, immunosuppressive agents, thymectomy, and plasmapheresis or intravenous immunoglobulin (IVIg) (Fig. 461-2).

ANTICHOLINESTERASE MEDICATIONS

Anticholinesterase medication produces at least partial improvement in most myasthenic patients, although improvement is complete in only a few. Patients with anti-MuSK MG generally obtain less benefit from anticholinesterase agents than those with AChR antibodies. Pyridostigmine is the most widely used anticholinesterase drug. The beneficial action of oral pyridostigmine begins within 15–30 min and lasts for 3–4 h, but individual responses vary. Treatment is begun with a moderate dose, e.g., 30–60 mg three to four times daily. The frequency and amount of the dose should be tailored to the patient's individual requirements throughout the day. For example, patients with weakness in chewing and swallowing may benefit by taking the medication before meals so that peak strength coincides with mealtimes. Long-acting pyridostigmine may occasionally be useful to get the patient through the night but should not be used for daytime medication because of variable absorption. The maximum useful dose of pyridostigmine rarely exceeds 120 mg every 4–6 h during daytime. Overdosage with anticholinesterase medication may cause increased weakness and other side effects. In some patients, muscarinic side effects of the anticholinesterase medication (diarrhea, abdominal cramps, salivation, nausea) may limit the dose tolerated. Atropine/diphenoxylate or loperamide is useful for the treatment of GI symptoms.

THYMECTOMY

Two separate issues should be distinguished: (1) surgical removal of thymoma, and (2) thymectomy as a treatment for MG. Surgical removal of a thymoma is necessary because of the possibility of local tumor spread, although most thymomas are histologically benign.

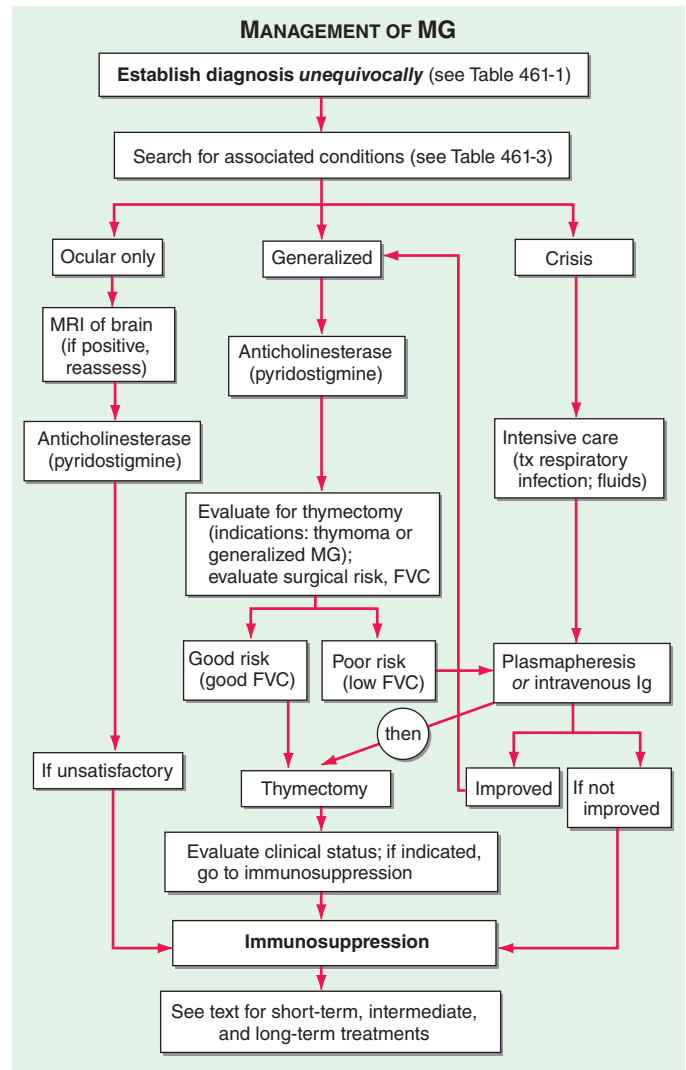


FIGURE 461-2 Algorithm for the management of myasthenia gravis. FVC, forced vital capacity; MRI, magnetic resonance imaging.

In the absence of a tumor, the available evidence suggests that up to 85% of patients experience improvement after thymectomy; of these, ~35% achieve drug-free remission. However, the improvement is typically delayed for months to years. The advantage of thymectomy is that it offers the possibility of long-term benefit, in some cases diminishing or eliminating the need for continuing medical treatment. Review of the published studies showed that following thymectomy, MG patients were 1.7 times as likely to improve and twice as likely to attain remission as those who did not have surgical thymectomy. In view of these potential benefits and of the negligible risk in skilled hands, thymectomy has gained widespread acceptance in the treatment of MG. It is the consensus that thymectomy should be carried out in all patients with generalized MG who are between the ages of puberty and at least 55 years. Whether thymectomy should be recommended in children, in adults >55 years of age, and in patients with weakness limited to the ocular muscles is still a matter of debate. There is also evidence that patients with MuSK antibody–positive MG respond less well to thymectomy than those with AChR antibody. Thymectomy must be carried out in a hospital where it is performed regularly and where the staff is experienced in the pre- and postoperative management, anesthesia, and surgical techniques of total thymectomy. Thymectomy should never be carried out as an emergency procedure, but only when the patient is adequately prepared. If necessary, treatment with IVIg or plasmapheresis may be used before surgery, but it is helpful to try to avoid immunosuppressive agents because of the risk of infection.

IMMUNOSUPPRESSION

Immunosuppression using one or more of the available agents is effective in nearly all patients with MG. The choice of drugs or other immunomodulatory treatments should be guided by the relative benefits and risks for the individual patient and the urgency of treatment. It is helpful to develop a treatment plan based on short-term, intermediate-term, and long-term objectives. For example, if immediate improvement is essential either because of the severity of weakness or because of the patient's need to return to activity as soon as possible, IVIg should be administered or plasmapheresis should be undertaken. For the intermediate term, glucocorticoids and cyclosporine or tacrolimus generally produce clinical improvement within a period of 1–3 months. The beneficial effects of azathioprine and mycophenolate mofetil usually begin after many months (as long as a year), but these drugs have advantages for the long-term treatment of patients with MG. There is a growing body of evidence that rituximab is effective in many MG patients, especially those with MuSK antibody.

Glucocorticoid Therapy Glucocorticoids, when used properly, produce improvement in myasthenic weakness in the great majority of patients. To minimize adverse side effects, prednisone should be given in a single dose rather than in divided doses throughout the day. The initial dose should be relatively low (15–25 mg/d) to avoid the early weakening that occurs in up to one-third of patients treated initially with a high-dose regimen. The dose is increased stepwise, as tolerated by the patient (usually by 5 mg/d at 2- to 3-day intervals), until there is marked clinical improvement or a dose of 50–60 mg/d is reached. This dose is maintained for 1–3 months and then is gradually modified to an alternate-day regimen over the course of an additional 1–3 months; the goal is to reduce the dose on the “off day” to zero or to a minimal level. Generally, patients begin to improve within a few weeks after reaching the maximum dose, and improvement continues to progress for months or years. The prednisone dosage may gradually be reduced, but usually months or years may be needed to determine the minimum effective dose, and close monitoring is required. Few patients are able to do without immunosuppressive agents entirely. Patients on long-term glucocorticoid therapy must be followed carefully to prevent or treat adverse side effects. The most common errors in glucocorticoid treatment of myasthenic patients include (1) insufficient persistence—improvement may be delayed and gradual; (2) tapering the dosage too early, too rapidly, or excessively; and (3) lack of attention to prevention and treatment of side effects.

The management of patients treated with glucocorticoids is discussed in Chap. 406.

Other Immunosuppressive Drugs Mycophenolate mofetil, azathioprine, cyclosporine, tacrolimus, rituximab, and occasionally cyclophosphamide are effective in many patients, either alone or in various combinations.

Mycophenolate mofetil has become one of the most widely used drugs in the treatment of MG because of its effectiveness and relative lack of side effects. A dose of 1–1.5 g bid is recommended. Its mechanism of action involves inhibition of purine synthesis by the *de novo* pathway. Since lymphocytes have only the *de novo* pathway, but lack the alternative salvage pathway that is present in all other cells, mycophenolate inhibits proliferation of lymphocytes but not proliferation of other cells. It does not kill or eliminate preexisting autoreactive lymphocytes, and therefore clinical improvement may be delayed for many months to a year, until the preexisting autoreactive lymphocytes die spontaneously. The advantage of mycophenolate lies in its relative lack of adverse side effects, with only occasional production of GI symptoms, rare development of leukopenia, and very small risks of malignancy or progressive multifocal leukoencephalopathy inherent in nearly all immunosuppressive treatments. Although two published studies did not show positive outcomes, most experts attribute the negative results to flaws in the trial designs, and mycophenolate is widely used for long-term treatment of myasthenic patients.

Until recently, azathioprine has been the most commonly used immunosuppressive agent for MG because of its relative safety in most patients and long track record. Its therapeutic effect may add to that of glucocorticoids and/or allow the glucocorticoid dose to be reduced. However, up to 10% of patients are unable to tolerate azathioprine because of idiosyncratic reactions consisting of flu-like symptoms of fever and malaise, bone marrow suppression, or abnormalities of liver function. An initial dose of 50 mg/d should be used for several days to test for these side effects. If this dose is tolerated, it is increased gradually to about 2–3 mg/kg of total body weight, or until the white blood count falls to 3000–4000/ μ L. The beneficial effect of azathioprine takes 3–6 months to begin and even longer to peak. In patients taking azathioprine, allopurinol should never be used to treat hyperuricemia. Because the two drugs share a common degradation pathway; the result may be severe bone marrow suppression due to increased effects of the azathioprine.

The calcineurin inhibitors cyclosporine and tacrolimus (FK506) are approximately as effective as azathioprine and are being used increasingly in the management of MG. Their beneficial effect appears more rapidly than that of azathioprine. Either drug may be used alone, but they are usually used as an adjunct to glucocorticoids to permit reduction of the glucocorticoid dose. The usual dose of cyclosporine is 4–5 mg/kg per day, and the average dose of tacrolimus is 0.07–0.1 mg/kg per day, given in two equally divided doses (to minimize side effects). Side effects of these drugs include hypertension and nephrotoxicity, which must be closely monitored. “Trough” blood levels are measured 12 h after the evening dose. The therapeutic range for the trough level of cyclosporine is 150–200 ng/L, and for tacrolimus, it is 5–15 ng/L.

Rituximab (Rituxan) is a monoclonal antibody that binds to the CD20 molecule on B lymphocytes. It has been widely used for the treatment of B cell lymphomas and has also proven successful in the treatment of several autoimmune diseases including rheumatoid arthritis, pemphigus, and some IgM-related neuropathies. There is now an extensive literature on the benefit of rituximab in MG. It is particularly effective in MuSK antibody-positive MG, although some patients with AChR antibody MG respond to it as well. The usual dose is 375 mg/m², given IV in 4 weekly infusions, or 1 g, given IV on two occasions 2 weeks apart.

For the occasional patient with MG that is genuinely refractory to optimal treatment with conventional immunosuppressive agents, a course of high-dose cyclophosphamide may induce long-lasting benefit by “rebooting” the immune system. At high doses, cyclophosphamide eliminates mature lymphocytes but spares hematopoietic precursors (stem cells), because they express the enzyme aldehyde dehydrogenase, which hydrolyzes cyclophosphamide. At present, this procedure is reserved for refractory patients and should be administered only in a facility fully familiar with this approach. Maintenance immunotherapy after rebooting is usually required to sustain the beneficial effect.

PLASMAPHERESIS AND INTRAVENOUS IMMUNOGLOBULIN

Plasmapheresis has been used therapeutically in MG. Plasma, which contains the pathogenic antibodies, is mechanically separated from the blood cells, which are returned to the patient. A course of five exchanges (3–4 L per exchange) is generally administered over a 10- to 14-day period. Plasmapheresis produces a short-term reduction in anti-AChR antibodies, with clinical improvement in many patients. It is useful as a temporary expedient in seriously affected patients or to improve the patient's condition prior to surgery (e.g., thymectomy).

The indications for the use of IVIg are the same as those for plasma exchange: to produce rapid improvement to help the patient through a difficult period of myasthenic weakness or prior to surgery. This treatment has the advantages of not requiring special equipment or large-bore venous access. The usual dose is 2 g/kg, which is typically administered over 5 days (400 mg/kg per day). If tolerated, the total dose of IVIg can be given over a 3- to 4-day period. Improvement occurs in ~70% of patients, beginning during

treatment, or within a week, and continuing for weeks to months. The mechanism of action of IVIg is not known; the treatment has no consistent effect on the measurable amount of circulating AChR antibody. Adverse reactions are generally not serious but may include headache, fluid overload, and rarely aseptic meningitis or renal failure. IVIg should rarely be used as a long-term treatment in place of rationally managed immunosuppressive therapy. Unfortunately, there is a tendency for physicians unfamiliar with immunosuppressive treatments to rely on repeated IVIg infusions, which usually produce only intermittent benefit, do not reduce the underlying autoimmune response, and are very costly. The intermediate and long-term treatment of myasthenic patients requires other methods of therapy outlined earlier in this chapter.

MANAGEMENT OF MYASTHENIC CRISIS

Myasthenic crisis is defined as an exacerbation of weakness sufficient to endanger life; it usually consists of respiratory failure caused by diaphragmatic and intercostal muscle weakness. Crisis rarely occurs in properly managed patients. Treatment should be carried out in intensive care units staffed with teams experienced in the management of MG, respiratory insufficiency, infectious disease, and fluid and electrolyte therapy. The possibility that deterioration could be due to excessive anticholinesterase medication (“cholinergic crisis”) is best excluded by temporarily stopping anticholinesterase drugs. The most common cause of crisis is intercurrent infection. This should be treated immediately, because the mechanical and immunologic defenses of the patient can be assumed to be compromised. The myasthenic patient with fever and early infection should be treated like other immunocompromised patients. Early and effective antibiotic therapy, respiratory assistance (preferably noninvasive, using bilevel positive airway pressure), and pulmonary physiotherapy are essentials of the treatment program. As discussed above, plasmapheresis or IVIg is frequently helpful in hastening recovery.

DRUGS TO AVOID IN MYASTHENIC PATIENTS

Many drugs have been reported to exacerbate weakness in patients with MG (Table 461-4), but not all patients react adversely to all of

TABLE 461-4 DRUGS WITH INTERACTIONS IN MYASTHENIA GRAVIS (MG)

Drugs That May Exacerbate MG
Antibiotics
Aminoglycosides: e.g., streptomycin, tobramycin, kanamycin
Quinolones: e.g., ciprofloxacin, levofloxacin, ofloxacin, gatifloxacin
Macrolides: e.g., erythromycin, azithromycin
Nondepolarizing muscle relaxants for surgery
D-Tubocurarine (curare), pancuronium, vecuronium, atracurium
Beta-blocking agents
Propranolol, atenolol, metoprolol
Local anesthetics and related agents
Procaine, Xylocaine in large amounts
Procainamide (for arrhythmias)
Botulinum toxin
Botox exacerbates weakness
Quinine derivatives
Quinine, quinidine, chloroquine, mefloquine (Lariam)
Magnesium
Decreases acetylcholine release
Penicillamine
May cause MG
Drugs with Important Interactions in MG
Cyclosporine
Broad range of drug interactions, which may raise or lower cyclosporine levels.
Azathioprine
Avoid allopurinol—combination may result in myelosuppression.

Myasthenia Gravis Worksheet				
History				
General	Normal	Good	Fair	Poor
Diplopia	None	Rare	Occasional	Constant
Ptosis	None	Rare	Occasional	Constant
Arms	Normal	Slightly limited	Some ADL impairment	Definitely limited
Legs	Normal	Walks/runs fatigues	Can walk limited distances	Minimal walking
Speech	Normal	Dysarthric	Severely dysarthric	Unintelligible
Voice	Normal	Fades	Impaired	Severely impaired
Chew	Normal	Fatigue on normal foods	Fatigue on soft foods	Feeding tube
Swallow	Normal	Normal foods	Soft foods only	Feeding tube
Respiration	Normal	Dyspnea on unusual effort	Dyspnea on any effort	Dyspnea at rest

Examination

BP _____	Pulse _____	Wt _____	Arm abduction time R _____ L _____
Edema _____			Deltoids R _____ L _____
Vital capacity _____			Biceps R _____ L _____
Cataracts? R _____ L _____			Triceps R _____ L _____
EOMS _____			Grip R _____ L _____
Ptosis time _____			Iliopsoas R _____ L _____
Face _____			Quadriceps R _____ L _____
			Hamstrings R _____ L _____
			Other R _____ L _____

FIGURE 461-3 Abbreviated interval assessment form for use in evaluating treatment for myasthenia gravis.

these. Conversely, not all “safe” drugs can be used with impunity in patients with MG. As a rule, the listed drugs should be avoided *whenever possible*, and myasthenic patients should be followed closely when *any new drug* is introduced.

PATIENT ASSESSMENT

To evaluate the effectiveness of treatment as well as drug-induced side effects, it is important to assess the patient’s clinical status systematically at baseline and on repeated interval examinations. Because of the variability of symptoms of MG, the interval history and physical findings on examination must be taken into account. The most useful clinical tests include forward arm abduction time (up to a full 5 min), spirometry with determination of forced vital capacity, range of eye movements, and time to development of ptosis on upward gaze. Manual muscle testing or, preferably, quantitative dynamometry of limb muscles, especially proximal muscles, is also important. An interval form can provide a succinct summary of the patient’s status and a guide to treatment results; an abbreviated form is shown in Fig. 461-3. A progressive reduction in the patient’s AChR antibody level also provides clinically valuable confirmation of the effectiveness of treatment; conversely, a rise in AChR antibody levels during tapering of immunosuppressive medication may predict clinical exacerbation. For reliable quantitative measurement of AChR antibody levels, it is best to compare antibody levels from prior frozen serum aliquots with current serum samples in simultaneously run assays.

462e Muscular Dystrophies and Other Muscle Diseases

Anthony A. Amato, Robert H. Brown, Jr.

Skeletal muscle diseases, or myopathies, are disorders with structural changes or functional impairment of muscle. These conditions can be differentiated from other diseases of the motor unit (e.g., lower motor neuron or neuromuscular junction pathologies) by characteristic clinical and laboratory findings.

Myasthenia gravis and related disorders are discussed in Chap. 461; dermatomyositis, polymyositis, and inclusion body myositis are discussed in Chap. 388.

CLINICAL FEATURES

Most myopathies present with proximal, symmetric limb weakness (arms or legs) with preserved reflexes and sensation. However, asymmetric and predominantly distal weakness can be seen in some myopathies. An associated sensory loss suggests injury to a peripheral nerve or the central nervous system (CNS) rather than myopathy. On occasion, disorders affecting the motor nerve cell bodies in the spinal cord (anterior horn cell disease), the neuromuscular junction, or peripheral nerves can mimic findings of myopathy.

Muscle Weakness Symptoms of muscle weakness can be either intermittent or persistent. Disorders causing *intermittent weakness* (Fig. 462e-1) include myasthenia gravis, periodic paralyses (hypokalemic, hyperkalemic, and paramyotonia congenita), and metabolic energy deficiencies of glycolysis (especially myophosphorylase deficiency), fatty acid utilization (carnitine palmitoyltransferase deficiency), and some mitochondrial myopathies. The states of energy deficiency cause activity-related muscle breakdown accompanied by myoglobinuria, appearing as light-brown- to dark-brown-colored urine.

Most muscle disorders cause *persistent weakness* (Fig. 462e-2). In the majority of these, including most types of muscular dystrophy, polymyositis, and dermatomyositis, the proximal muscles are weaker than the distal and are symmetrically affected, and the facial muscles are spared, a pattern referred to as *limb-girdle*. The differential diagnosis is more restricted for other patterns of weakness. Facial weakness (difficulty with eye closure and impaired smile) and scapular winging

(Fig. 462e-3) are characteristic of facioscapulohumeral dystrophy (FSHD). Facial and distal limb weakness associated with hand grip myotonia is virtually diagnostic of myotonic dystrophy type 1. When other cranial nerve muscles are weak, causing ptosis or extraocular muscle weakness, the most important disorders to consider include neuromuscular junction disorders, oculopharyngeal muscular dystrophy, mitochondrial myopathies, or some of the congenital myopathies (Table 462e-1). A pathognomonic pattern characteristic of inclusion body myositis is atrophy and weakness of the flexor forearm (e.g., wrist and finger flexors) and quadriceps muscles that is often asymmetric. Less frequently, but important diagnostically, is the presence of a dropped head syndrome indicative of selective neck extensor muscle weakness. The most important neuromuscular diseases associated with this pattern of weakness include myasthenia gravis, amyotrophic lateral sclerosis, late-onset nemaline myopathy, hyperparathyroidism, focal myositis, and some forms of inclusion body myopathy. A final pattern, recognized because of preferential distal extremity weakness, is typical of a unique category of muscular dystrophy, the distal myopathies.

It is important to examine functional capabilities to help disclose certain patterns of weakness (Table 462e-2). The Gowers' sign (Fig. 462e-4) is particularly useful. Observing the gait of an individual may disclose a lordotic posture caused by combined trunk and hip weakness, frequently exaggerated by toe walking (Fig. 462e-5). A waddling gait is caused by the inability of weak hip muscles to prevent hip drop or hip dip. Hyperextension of the knee (genu recurvatum or back-kneeing) is characteristic of quadriceps muscle weakness; and a steppage gait, due to footdrop, accompanies distal weakness.

Any disorder causing muscle weakness may be accompanied by *fatigue*, referring to an inability to maintain or sustain a force (pathologic fatigability). This condition must be differentiated from *asthenia*, a type of fatigue caused by excess tiredness or lack of energy. Associated symptoms may help differentiate asthenia and pathologic fatigability. Asthenia is often accompanied by a tendency to avoid physical activities, complaints of daytime sleepiness, necessity for frequent naps, and difficulty concentrating on activities such as reading. There may be feelings of overwhelming stress and depression. Thus, asthenia is not a myopathy. In contrast, pathologic fatigability occurs in disorders of neuromuscular transmission and in disorders altering energy production, including defects in glycolysis, lipid metabolism, or mitochondrial energy production. Pathologic fatigability also occurs in chronic myopathies because of difficulty accomplishing a task with less

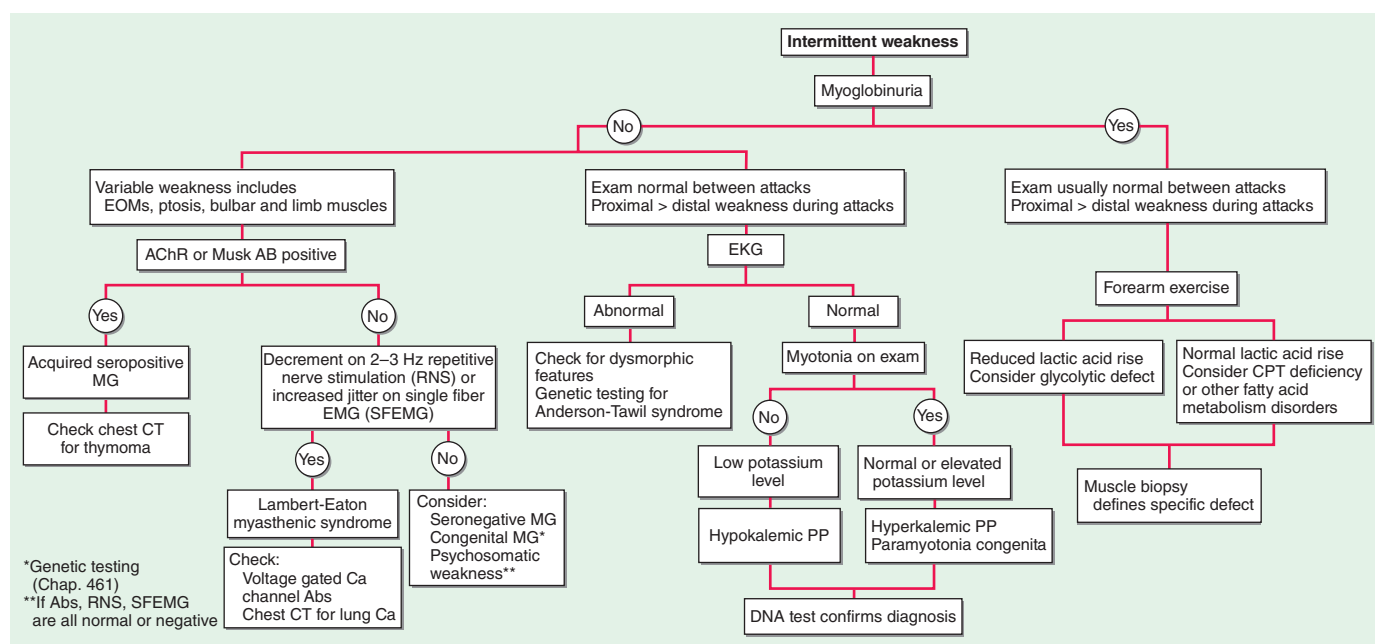


FIGURE 462e-1 Diagnostic evaluation of intermittent weakness. AChR AB, acetylcholine receptor antibody; CPT, carnitine palmitoyltransferase; EOMs, extraocular muscles; MG, myasthenia gravis; PP, periodic paralysis.

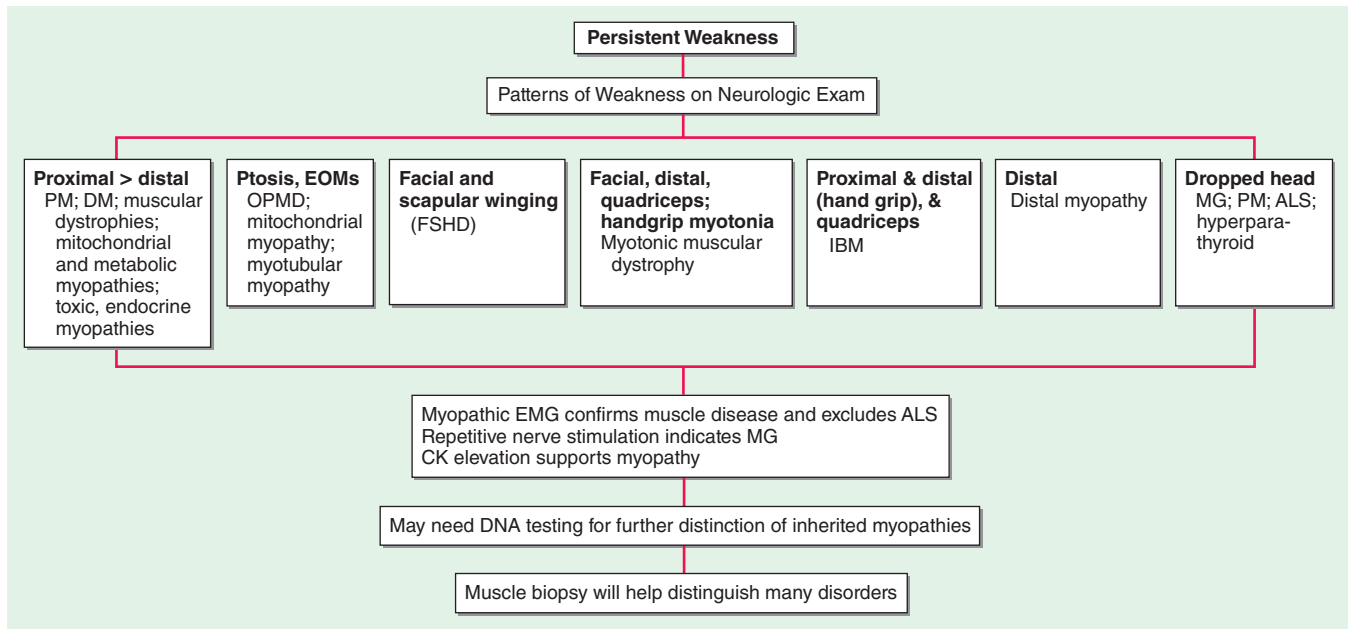


FIGURE 462e-2 Diagnostic evaluation of persistent weakness. Examination reveals one of seven patterns of weakness. The pattern of weakness in combination with the laboratory evaluation leads to a diagnosis. ALS, amyotrophic lateral sclerosis; CK, creatine kinase; DM, dermatomyositis; EMG, electromyography; EOMs, extraocular muscles; FSHD, facioscapulohumeral dystrophy; IBM, inclusion body myositis; MG, myasthenia gravis; OPMD, oculopharyngeal muscular dystrophy; PM, polymyositis.

muscle. Pathologic fatigability is accompanied by abnormal clinical or laboratory findings. Fatigue without those supportive features almost never indicates a primary muscle disease.

Muscle Pain (Myalgias), Cramps, and Stiffness Muscle pain can be associated with cramps, spasms, contractures, and stiff or rigid muscles. In distinction, true myalgia (muscle aching), which can be localized or generalized, may be accompanied by weakness, tenderness to palpation, or swelling. Certain drugs cause true myalgia ([Table 462e-3](#)).

There are two painful muscle conditions of particular importance, neither of which is associated with muscle weakness. *Fibromyalgia* is a common, yet poorly understood, type of myofascial pain syndrome. Patients complain of severe muscle pain and tenderness and have specific painful trigger points, sleep disturbances, and easy fatigability. Serum creatine kinase (CK), erythrocyte sedimentation rate (ESR), electromyography (EMG), and muscle biopsy are normal ([Chap. 396](#)). *Polymyalgia rheumatica* occurs mainly in patients >50 years and is characterized by stiffness and pain in the shoulders, lower back, hips, and thighs ([Chap. 385](#)). The ESR is elevated, while serum CK, EMG, and muscle biopsy are normal. Temporal arteritis, an inflammatory



FIGURE 462e-3 Facioscapulohumeral dystrophy with prominent scapular winging.

TABLE 462e-1 NEUROMUSCULAR CAUSES OF PTOSIS OR OPHTHALMOPLEGIA

Peripheral Neuropathy

Guillain-Barré syndrome
Miller Fisher syndrome

Neuromuscular Junction

Botulism
Lambert-Eaton syndrome
Myasthenia gravis
Congenital myasthenia

Myopathy

Mitochondrial myopathies
Kearns-Sayre syndrome
Progressive external ophthalmoplegia
Oculopharyngeal and oculopharyngodistal muscular dystrophy
Myotonic dystrophy (ptosis only)
Congenital myopathy
Myotubular
Nemaline (ptosis only)
Hyperthyroidism/Graves' disease (ophthalmoplegia without ptosis)
Hereditary inclusion body myopathy type 3

disorder of medium- and large-sized arteries, usually involving one or more branches of the carotid artery, may accompany polymyalgia rheumatica. Vision is threatened by ischemic optic neuritis. Glucocorticoids can relieve the myalgias and protect against visual loss.

Localized muscle pain is most often traumatic. A common cause of sudden abrupt-onset pain is a ruptured tendon, which leaves the muscle belly appearing rounded and shorter in appearance compared to the normal side. The biceps brachii and Achilles tendons are particularly vulnerable to rupture. Infection or neoplastic infiltration of the muscle is a rare cause of localized muscle pain.

A *muscle cramp* or *spasm* is a painful, involuntary, localized, muscle contraction with a visible or palpable hardening of the muscle. Cramps are abrupt in onset, short in duration, and may cause abnormal posturing of the joint. The EMG shows firing of motor units, reflecting an origin from spontaneous neural discharge. Muscle cramps often occur

TABLE 462e-2 OBSERVATIONS ON EXAMINATION THAT DISCLOSE MUSCLE WEAKNESS

Functional Impairment	Muscle Weakness
Inability to forcibly close eyes	Upper facial muscles
Impaired pucker	Lower facial muscles
Inability to raise head from prone position	Neck extensor muscles
Inability to raise head from supine position	Neck flexor muscles
Inability to raise arms above head	Proximal arm muscles (may be only scapular stabilizing muscles)
Inability to walk without hyperextending knee (back-kneeing or genu recurvatum)	Knee extensor muscles
Inability to walk with heels touching the floor (toe walking)	Shortening of the Achilles tendon
Inability to lift foot while walking (steppage gait or footdrop)	Anterior compartment of leg
Inability to walk without a waddling gait	Hip muscles
Inability to get up from the floor without climbing up the extremities (Gowers' sign)	Hip, thigh, and trunk muscles
Inability to get up from a chair without using arms	Hip muscles

in neurogenic disorders, especially motor neuron disease (Chap. 452), radiculopathies, and polyneuropathies (Chap. 459), but are not a feature of most primary muscle diseases. Duchenne muscular dystrophy is an exception because calf muscle complaints are a common complaint. Muscle cramps are also common during pregnancy.

A *muscle contracture* is different from a muscle cramp. In both conditions, the muscle becomes hard, but a contracture is associated with energy failure in glycolytic disorders. The muscle is unable to relax after an active muscle contraction. The EMG shows electrical silence. Confusion is created because contracture also refers to a muscle that cannot be passively stretched to its proper length (fixed contracture) because of fibrosis. In some muscle disorders, especially in Emery-Dreifuss muscular dystrophy and Bethlem myopathy, fixed contractures occur early and represent distinctive features of the disease.

Muscle stiffness can refer to different phenomena. Some patients with inflammation of joints and periarticular surfaces feel stiff. This condition is different from the disorders of hyperexcitable motor nerves causing stiff or rigid muscles. In *stiff-person syndrome*, spontaneous discharges of the motor neurons of the spinal cord cause involuntary muscle contractions mainly involving the axial (trunk) and proximal lower extremity muscles. The gait becomes stiff and labored, with hyperlordosis of the lumbar spine. Superimposed episodic muscle spasms are precipitated by sudden movements, unexpected noises, and emotional upset. The muscles relax during sleep. Serum antibodies against glutamic acid decarboxylase are present in approximately two-thirds of cases. In *neuromyotonia* (Isaacs' syndrome), there is hyperexcitability of the peripheral nerves manifesting as continuous muscle fiber activity. *Myokymia* (groups of fasciculations associated with continuous undulations of muscle) and impaired muscle relaxation are the result. Muscles of the leg are stiff, and the constant contractions of the muscle cause increased sweating of the extremities. This peripheral nerve hyperexcitability is mediated by antibodies that target voltage-gated potassium channels. The site of origin of the spontaneous nerve discharges is principally in the distal portion of the motor nerves.

Myotonia is a condition of prolonged muscle contraction followed by slow muscle relaxation. It always follows muscle activation (action myotonia), usually voluntary, but may be elicited by mechanical stimulation (percussion myotonia) of the muscle. Myotonia typically causes difficulty in releasing objects after a firm grasp. In myotonic muscular dystrophy type 1 (DM1), distal weakness usually accompanies myotonia, whereas in DM2, proximal muscles are more affected; thus

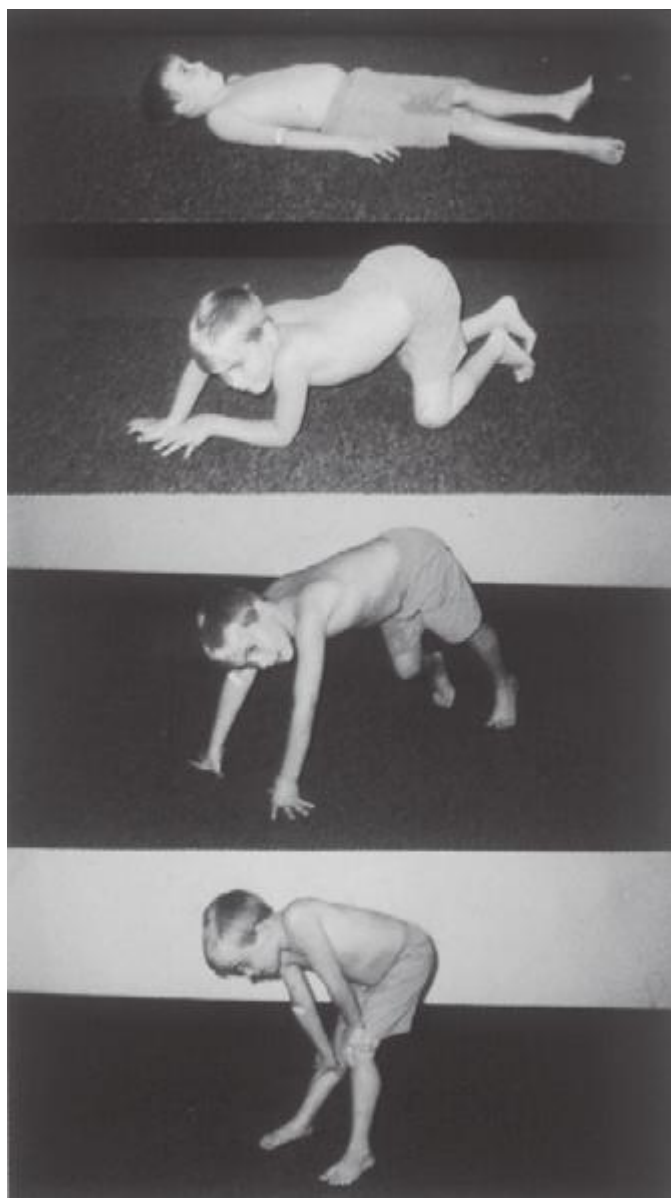


FIGURE 462e-4 Gowers' sign showing a patient using his arms to climb up the legs in attempting to get up from the floor.

the related term *proximal myotonic myopathy* (PROMM) is used to describe this condition. Myotonia also occurs with *myotonia congenita* (a chloride channel disorder), but in this condition muscle weakness is not prominent. Myotonia may also be seen in individuals with sodium channel mutations (*hyperkalemic periodic paralysis* or *potassium-sensitive myotonia*). Another sodium channelopathy, *paramyotonia congenita*, also is associated with muscle stiffness. In contrast to other disorders associated with myotonia in which the myotonia is eased by repetitive activity, paramyotonia congenita is named for a paradoxical phenomenon whereby the myotonia worsens with repetitive activity.

Muscle Enlargement and Atrophy In most myopathies muscle tissue is replaced by fat and connective tissue, but the size of the muscle is usually not affected. However, in many limb-girdle muscular dystrophies (and particularly the dystrophinopathies), enlarged calf muscles are typical. The enlargement represents true muscle hypertrophy; thus the term *pseudohypertrophy* should be avoided when referring to these patients. The calf muscles remain very strong even late in the course of these disorders. Muscle enlargement can also result from infiltration by sarcoid granulomas, amyloid deposits, bacterial and parasitic infections, and focal myositis. In contrast, muscle atrophy is characteristic of other myopathies. In dysferlinopathies (LGMD2B)

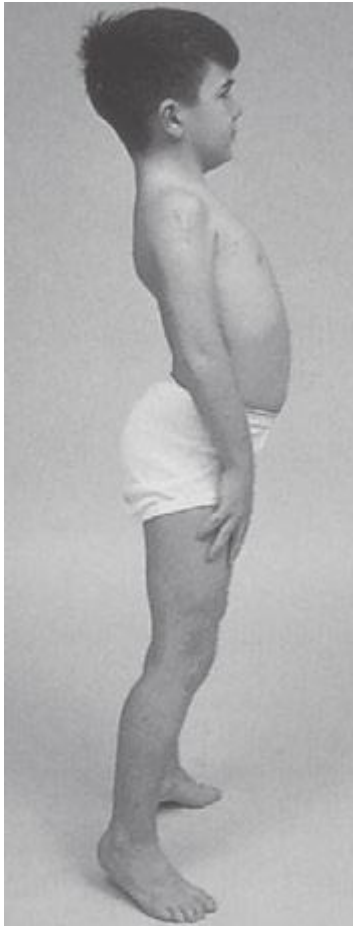


FIGURE 462e-5 Lordotic posture, exaggerated by standing on toes, associated with trunk and hip weakness.

and anoctaminopathies (LGMD2L), there is a predilection for early atrophy of the gastrocnemius muscles, particularly the medial aspect. Atrophy of the humeral muscles is characteristic of FSHD.

LABORATORY EVALUATION

A limited battery of tests can be used to evaluate a suspected myopathy. Nearly all patients require serum enzyme level measurements and electrodiagnostic studies as screening tools to differentiate muscle disorders from other motor unit diseases. The other tests described—DNA studies, the forearm exercise test, and muscle biopsy—are used to diagnose specific types of myopathies.

Serum Enzymes CK is the preferred muscle enzyme to measure in the evaluation of myopathies. Damage to muscle causes the CK to leak

TABLE 462e-3 DRUGS THAT CAUSE TRUE MYALGIA

Cimetidine
Cocaine
Cyclosporine
Danazol
Emetine
Gold
Heroin
Labetalol
Methadone
D-Penicillamine
Statins and other cholesterol-lowering agents
L-Tryptophan
Zidovudine

from the muscle fiber to the serum. The MM isoenzyme predominates in skeletal muscle, whereas creatine kinase-myocardial bound (CK-MB) is the marker for cardiac muscle. Serum CK can be elevated in normal individuals without provocation, presumably on a genetic basis or after strenuous activity, minor trauma (including the EMG needle), a prolonged muscle cramp, or a generalized seizure. Aspartate aminotransferase (AST), alanine aminotransferase (ALT), aldolase, and lactic dehydrogenase (LDH) are enzymes sharing an origin in both muscle and liver. Problems arise when the levels of these enzymes are found to be elevated in a routine screening battery, leading to the erroneous assumption that liver disease is present when in fact muscle could be the cause. An elevated γ -glutamyl transferase (GGT) helps to establish a liver origin because this enzyme is not found in muscle.

Electrodiagnostic Studies EMG, repetitive nerve stimulation, and nerve conduction studies ([Chap. 442e](#)) are essential methods for evaluation of the patient with suspected muscle disease. In combination, they provide the information necessary to differentiate myopathies from neuropathies and neuromuscular junction diseases. Routine nerve conduction studies are typically normal in myopathies but reduced amplitudes of compound muscle action potentials may be seen in atrophied muscles. The needle EMG may reveal irritability on needle placement suggestive of a necrotizing myopathy (inflammatory myopathies, dystrophies, toxic myopathies, myotonic myopathies), whereas a lack of irritability is characteristic of long-standing myopathic disorders (muscular dystrophies, endocrine myopathies, disuse atrophy, and many of the metabolic myopathies). In addition, the EMG may demonstrate myotonic discharges that will narrow the differential diagnosis ([Table 462e-4](#)). Another important EMG finding is the presence of short-duration, small-amplitude, polyphasic motor unit action potentials (MUAPs). Such MUAPs can be seen in both myopathic and neuropathic disorders; however, the recruitment or firing pattern is different. In myopathies, the MUAPs fire early but at a normal rate to compensate for the loss of individual muscle fibers, whereas in neurogenic disorders the MUAPs fire faster. The EMG is usually normal in steroid or disuse myopathy, both of which are associated with type 2 fiber atrophy; this is because the EMG preferentially assesses the physiologic function of type 1 fibers. The EMG can also be invaluable in helping to choose an appropriately affected muscle to sample for biopsy.

DNA Analysis This serves as an important tool for the definitive diagnosis of many muscle disorders. Nevertheless, there are a number of limitations in currently available molecular diagnostics. For example, in Duchenne and Becker dystrophies, two-thirds of patients have deletion or duplication mutations in the dystrophin gene that are easy to detect, while the remainder have point mutations that are much more difficult to find. For patients without identifiable gene defects, the muscle biopsy remains the main diagnostic tool.

TABLE 462e-4 MYOTONIC DISORDERS

Myotonic dystrophy type 1
Myotonic dystrophy type 2/proximal myotonic myopathy
Myotonia congenita
Paramyotonia congenita
Hyperkalemic periodic paralysis
Chondrodystrophic myotonia (Schwartz-Jampel syndrome)
Centronuclear/myotubular myopathy ^a
Drug-induced
Cholesterol-lowering agents (statin medications, fibrates)
Cyclosporine
Chloroquine
Glycogen storage disorders ^a (Pompe's disease, branching enzyme deficiency, debranching enzyme deficiency)
Myofibrillar myopathies (MFM) ^a

^aAssociated with myotonic discharges on electromyography but no clinical myotonia.

Forearm Exercise Test In myopathies with intermittent symptoms, and especially those associated with myoglobinuria, there may be a defect in glycolysis. Many variations of the forearm exercise test exist. For safety, the test should not be performed under ischemic conditions to avoid an unnecessary insult to the muscle, causing rhabdomyolysis. The test is performed by placing a small indwelling catheter into an antecubital vein. A baseline blood sample is obtained for lactic acid and ammonia. The forearm muscles are exercised by asking the patient to vigorously open and close the hand for 1 min. Blood is then obtained at intervals of 1, 2, 4, 6, and 10 min for comparison with the baseline sample. A three- to fourfold rise of lactic acid is typical. The simultaneous measurement of ammonia serves as a control, because it should also rise with exercise. In patients with myophosphorylase deficiency or other glycolytic defects, the lactic acid rise will be absent or below normal, while the rise in ammonia will reach control values. If there is lack of effort, neither lactic acid nor ammonia will rise. Patients with selective failure to increase ammonia may have myoadenylate deaminase deficiency. This condition has been reported to be a cause of myoglobinuria, but deficiency of this enzyme in asymptomatic individuals makes interpretation controversial.

Muscle Biopsy Muscle biopsy is an important step in establishing the diagnosis of a suspected myopathy. The biopsy is usually obtained from a quadriceps or biceps brachii muscle, less commonly from a deltoid muscle. Evaluation includes a combination of techniques—light microscopy, histochemistry, immunocytochemistry with a battery of antibodies, and electron microscopy. Not all techniques are needed for every case. A specific diagnosis can be established in many disorders. Endomysial inflammatory cells surrounding and invading muscle

fibers are seen in polymyositis; similar endomysial infiltrates associated with muscle fibers containing rimmed vacuoles and amyloid deposits consisting of SMI-31-, p62-, and TDP-43-positive inclusions within fibers are characteristic of inclusion body myositis; and perivascular, perimysial inflammation associated with perifascicular atrophy is a feature of dermatomyositis. In addition, the congenital myopathies have distinctive light and electron microscopy features essential for diagnosis. Mitochondrial and metabolic (e.g., glycogen and lipid storage diseases) myopathies also demonstrate distinctive histochemical and electron-microscopic profiles. Biopsied muscle tissue can be sent for metabolic enzyme or mitochondrial DNA analyses. A battery of antibodies is available for the identification of abnormal proteins to help diagnose specific types of muscular dystrophies. Western blot analysis on muscle specimens can be performed to determine whether specific muscle proteins are reduced in quantity or are of abnormal size.

HEREDITARY MYOPATHIES

Muscular dystrophy refers to a group of hereditary progressive diseases each with unique phenotypic and genetic features (Tables 462e-5, 462e-6, and 462e-7).

DUCHENNE MUSCULAR DYSTROPHY

This X-linked recessive disorder, sometimes also called *pseudohypertrophic muscular dystrophy*, has an incidence of ~1 per 5200 live-born males.

Clinical Features Duchenne dystrophy is present at birth, but the disorder usually becomes apparent between ages 3 and 5 years. The boys fall frequently and have difficulty keeping up with friends when

TABLE 462e-5 PROGRESSIVE MUSCULAR DYSTROPHIES

Type	Inheritance	Defective Gene/Protein	Onset Age	Clinical Features	Other Organ Systems Involved
Duchenne	XR	Dystrophin	Before 5 years	Progressive weakness of girdle muscles Unable to walk after age 12 Progressive kyphoscoliosis Respiratory failure in second or third decade	Cardiomyopathy Mental impairment
Becker	XR	Dystrophin	Early childhood to adult	Progressive weakness of girdle muscles Able to walk after age 15 Respiratory failure may develop by fourth decade	Cardiomyopathy
Limb-girdle	AD/AR	Several (Tables 462e-6, 462e-7)	Early childhood to early adult	Slow progressive weakness of shoulder and hip girdle muscles	± Cardiomyopathy
Emery-Dreifuss	XR/AD	Emerin, lamin A/C Nesprin-1, nesprin-2, TMEM43	Childhood to adult	Elbow/knee/ankle contractures, humeral and peroneal weakness	Cardiomyopathy
Congenital	AR	Several	At birth or within first few months	Hypotonia, contractures, delayed milestones Progression to respiratory failure in some; static course in others	CNS abnormalities (hypomyelination, malformation) Eye abnormalities
Myotonic ^a (DM1, DM2)	AD	DM1: Expansion CTG repeat DM2: Expansion CCTG repeat	Childhood to adult; possibly infancy if mother affected (DM1 only)	Slowly progressive weakness of face, shoulder girdle, and foot dorsiflexion Preferential proximal weakness in DM2	Cardiac conduction defects Mental impairment Cataracts Frontal baldness Gonadal atrophy
FSHD1	AD	DUX4 4q	Childhood to adult	Slowly progressive weakness of face, shoulder girdle, and foot dorsiflexion	Deafness
FSHD2	AD	SMCHD1			Coats' (eye) disease
Oculopharyngeal	AD	Expansion, poly-A RNA binding protein	Fifth to sixth decade	Slowly progressive weakness of extraocular, pharyngeal, and limb muscles	—

^aTwo forms of myotonic dystrophy, DM1 and DM2, have been identified. Many features overlap (see text).

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; CNS, central nervous system; XR, X-linked recessive.

TABLE 462e-6 AUTOSOMAL DOMINANT LIMB-GIRDLE MUSCULAR DYSTROPHIES (LGMDs)

Disease	Clinical Features	Laboratory Features	Abnormal Protein
LGMD1A	Onset second to eighth decade Muscle weakness affects proximal and distal limb muscles, vocal cords, and pharyngeal muscles	Serum CK 2x normal EMG myopathic and may have pseudotonic discharges Muscle biopsy: features of MFM	Myotilin
LGMD1B	Onset first or second decade Proximal lower limb weakness and cardiomyopathy with conduction defects Some cases indistinguishable from Emery-Dreifuss muscular dystrophy with joint contractures	Serum CK 3–5x normal EMG myopathic	Lamin A/C
LGMD1C	Onset in early childhood Proximal weakness Gowers' sign, calf hypertrophy, rippling muscles Exercise-related muscle cramps	Serum CK 4–25x normal EMG myopathic	Caveolin-3
LGMD1D	Onset second to sixth decade Proximal and distal muscle weakness	Serum CK 2–3x normal EMG myopathic Muscle biopsy: features of MFM	DNAJB6
LGMD1E	Onset first to sixth decade Proximal or distal muscle weakness Cardiomyopathy and arrhythmias	Serum CK 2–4x normal EMG myopathic and may have pseudotonic discharges Muscle biopsy: features of MFM	Desmin
LGMD1F	Onset infancy to sixth decade Proximal or distal weakness May have early contractures resembling Emery-Dreifuss syndrome	Serum CK normal to 20x normal EMG myopathic Muscle biopsy may show enlarged nuclei with central pallor, rimmed vacuoles, and filamentous inclusions	TNPO3

Abbreviations: CK, creatine kinase; EMG, electromyography; MFM, myofibrillar myopathy; NCS, nerve conduction studies.

playing. Running, jumping, and hopping are invariably abnormal. By age 5 years, muscle weakness is obvious by muscle testing. On getting up from the floor, the patient uses his hands to climb up himself (Gowers' maneuver [Fig. 462e-4]). Contractures of the heel cords and iliotibial bands become apparent by age 6 years, when toe walking is associated with a lordotic posture. Loss of muscle strength is progressive, with predilection for proximal limb muscles and the neck flexors; leg involvement is more severe than arm involvement. Between ages 8 and 10 years, walking may require the use of braces; joint contractures and limitations of hip flexion, knee, elbow, and wrist extension are made worse by prolonged sitting. Prior to the use of glucocorticoids, most boys became wheelchair dependent by 12 years of age. Contractures become fixed, and a progressive scoliosis often develops that may be associated with pain. The chest deformity with scoliosis impairs pulmonary function, which is already diminished by muscle weakness. By age 16–18 years, patients are predisposed to serious, sometimes fatal pulmonary infections. Other causes of death include aspiration of food and acute gastric dilation.

A cardiac cause of death is uncommon despite the presence of a cardiomyopathy in almost all patients. Congestive heart failure seldom occurs except with severe stress such as pneumonia. Cardiac arrhythmias are rare. The typical electrocardiogram (ECG) shows an increased net RS in lead V_1 ; deep, narrow Q waves in the precordial leads; and tall right precordial R waves in V_1 . Intellectual impairment in Duchenne dystrophy is common; the average intelligence quotient (IQ) is ~1 standard deviation (SD) below the mean. Impairment of intellectual function appears to be nonprogressive and affects verbal ability more than performance.

Laboratory Features Serum CK levels are invariably elevated to between 20 and 100 times normal. The levels are abnormal at birth but decline late in the disease because of inactivity and loss of muscle mass. EMG demonstrates features typical of myopathy. The muscle biopsy shows muscle fibers of varying size as well as small groups of necrotic and regenerating fibers. Connective tissue and fat replace lost muscle fibers. A definitive diagnosis of Duchenne dystrophy can be established on the basis of dystrophin deficiency in a biopsy of

muscle tissue or mutation analysis on peripheral blood leukocytes, as discussed below.

Duchenne dystrophy is caused by a mutation of the gene that encodes dystrophin, a 427-kDa protein localized to the inner surface of the sarcolemma of the muscle fiber. The dystrophin gene is >2000 kb in size and thus is one of the largest identified human genes. It is localized to the short arm of the X chromosome at Xp21. The most common gene mutation is a deletion. The size varies but does not correlate with disease severity. Deletions are not uniformly distributed over the gene but rather are most common near the beginning (5' end) and middle of the gene. Less often, Duchenne dystrophy is caused by a gene duplication or point mutation. Identification of a specific mutation allows for an unequivocal diagnosis, makes possible accurate testing of potential carriers, and is useful for prenatal diagnosis.

A diagnosis of Duchenne dystrophy can also be made by Western blot analysis of muscle biopsy specimens, revealing abnormalities in the quantity and molecular weight of dystrophin protein. In addition, immunocytochemical staining of muscle with dystrophin antibodies can be used to demonstrate absence or deficiency of dystrophin localizing to the sarcolemmal membrane. Carriers of the disease may demonstrate a mosaic pattern, but dystrophin analysis of muscle biopsy specimens for carrier detection is not reliable.

Pathogenesis Dystrophin is part of a large complex of sarcolemmal proteins and glycoproteins (Fig. 462e-6). Dystrophin binds to F-actin at its amino terminus and to β -dystroglycan at the carboxyl terminus. β -Dystroglycan complexes to α -dystroglycan, which binds to laminin in the extracellular matrix (ECM). Laminin has a heterotrimeric molecular structure arranged in the shape of a cross with one heavy chain and two light chains, β_1 and γ_1 . The laminin heavy chain of skeletal muscle is designated laminin α_2 . Collagen proteins IV and VI are also found in the ECM. Like β -dystroglycan, the transmembrane sarcoglycan proteins also bind to dystrophin; these five proteins (designated α - through ϵ -sarcoglycan) complex tightly with each other. More recently, other membrane proteins implicated in muscular dystrophy have been found to be loosely affiliated with constituents of the dystrophin complex. These include caveolin-3, α , integrin, and collagen VI.

TABLE 462e-7 AUTOSOMAL RECESSIVE LIMB-GIRDLE MUSCULAR DYSTROPHIES (LGMDs)

Disease	Clinical Features	Laboratory Features	Abnormal Protein
LGMD2A	Onset first or second decade Scapular winging; no calf hypertrophy; no cardiac or respiratory muscle weakness Proximal and distal weakness; may have contractures at elbows, wrists, and fingers	Serum CK 3–15× normal EMG myopathic Muscle biopsy may show lobulated muscle fibers	Calpain-3
LGMD2B	Onset second or third decade Proximal muscle weakness at onset, later distal (calf) muscles affected Miyoshi's myopathy is variant of LGMD2B with calf muscles affected at onset	Serum CK 3–100× normal EMG myopathic Inflammation on muscle biopsy may simulate polymyositis	Dysferlin
LGMD2C–F	Onset in childhood to teenage years Clinical condition similar to Duchenne and Becker muscular dystrophies Cognitive function normal	Serum CK 5–100× normal EMG myopathic	γ, α, β, δ sarcoglycans
LGMD2G	Onset age 10 to 15 Proximal and distal muscle weakness	Serum CK 3–17× normal EMG myopathic Muscle biopsy may show rimmed vacuoles	Telethonin
LGMD2H	Onset first to third decade Proximal muscle weakness	Serum CK 2–25× normal EMG myopathic	<i>TRIM32</i> gene
LGMD2I	Onset first to third decade Clinical condition similar to Duchenne or Becker dystrophies Cardiomyopathy and respiratory failure may occur early before significant weakness Cognitive function normal	Serum CK 10–30× normal EMG myopathic	Fukutin-related protein
LGMD2J ^a	Onset first to third decade Proximal lower limb weakness Mild distal weakness Progressive weakness causes loss of ambulation	Serum CK 1.5–2× normal EMG myopathic Muscle biopsy reveals rimmed vacuoles	Titin
LGMD2K	Usually presents in infancy as Walker-Warburg syndrome but can present in early adult life with proximal weakness and only minor CNS abnormalities	CK 10–20× normal EMG myopathic	POMT1
LGMD2L	Presents in childhood or adult life May manifest with quadriceps atrophy and myalgia Some present with early involvement of the calves in the second decade of life, resembling Miyoshi's myopathy (dysferlinopathy)	CK 8–20× normal EMG myopathic	Anoctamin 5
LGMD2M	Usually presents in infancy as Fukuyama's congenital muscular dystrophy but can present in early adult life with proximal weakness and only minor CNS abnormalities	CK 10–50× normal EMG myopathic	Fukutin
LGMD2N	Usually presents in infancy as muscle-eye-brain disease but can present in early adult life with proximal weakness and only minor CNS abnormalities	CK 5–20× normal EMG myopathic	POMGnT1
LGMD2O	Usually presents in infancy as Walker-Warburg syndrome but can present in early adult life with proximal weakness and only minor CNS abnormalities	CK 5–20× normal EMG myopathic	POMT2
LGMD2P	One case reported presenting in early childhood	CK >10× normal	α-Dystroglycan
LGMD2Q	Onset in infancy to fourth decade; proximal weakness; may have ptosis and extraocular weakness; epidermolysis bullosa (also considered a congenital myasthenic syndrome)	CK variable, but usually only mildly elevated EMG myopathic Repetitive nerve stimulation may show decrement	Plectin 1
LGMD2R	See LGMD1E (Table 462e-6)	See LGMD1E	Desmin
LGMD2S	Onset in infancy to sixth decade Proximal weakness Eye abnormalities common; truncal ataxia and chorea Mild to moderate intellectual disability Hutterite descent	CK 1.5–20× normal	TRAPC11

^aUdd's type distal myopathy is a form of titin deficiency with only distal muscle weakness (see Table 462e-9).

Abbreviations: CK, creatine kinase; EMG, electromyography; NCS, nerve conduction studies; POMT1, protein-O-mannosyltransferase 1; POMT2, protein-O-mannosyltransferase 2; POMGnT1, O-linked mannose beta 1,2-N-acetylglucosaminyltransferase; TNPO3, transportin 3; TRAPC11, transport (trafficking) protein particle complex, subunit 11.

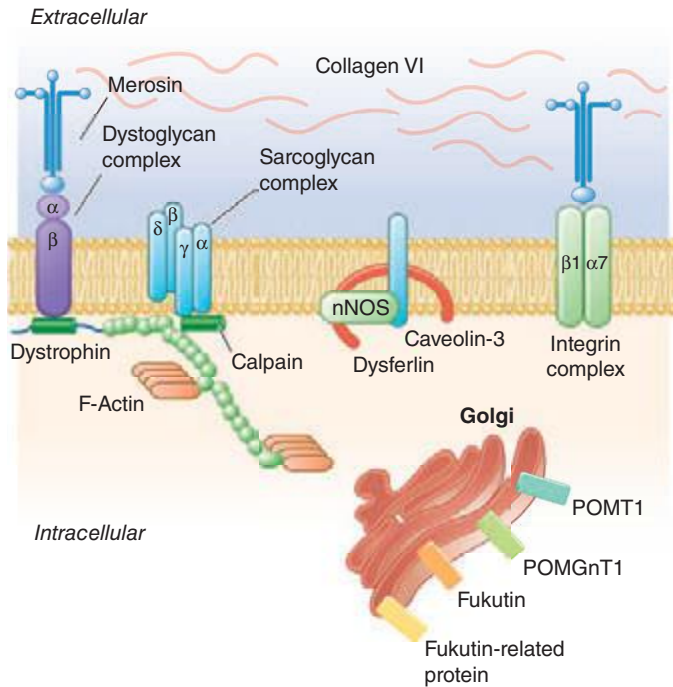


FIGURE 462e-6 Selected muscular dystrophy–associated proteins in the cell membrane and Golgi complex.

Dystrophin localizes to the cytoplasmic face of the muscle cell membrane. It complexes with two transmembrane protein complexes, the dystroglycans and the sarcoglycans. The dystroglycans bind to the extracellular matrix protein merosin, which is also complexed with β_1 and α_7 integrins (Tables 462e-5, 462e-6, and 462e-7). Dysferlin complexes with caveolin-3 (which binds to neuronal nitric oxide synthase, or nNOS) but not with the dystrophin-associated proteins or the integrins. In some of the congenital dystrophies and limb-girdle muscular dystrophies (LGMDs), there is loss of function of different enzymes that glycosylate α -dystroglycan, which thereby inhibits proper binding to merosin: POMT1, POMT2, POMGnT1, Fukutin, Fukutin-related protein, and LARGE.

The dystrophin-glycoprotein complex appears to confer stability to the sarcolemma, although the function of each individual component of the complex is incompletely understood. Deficiency of one member of the complex may cause abnormalities in other components. For example, a primary deficiency of dystrophin (Duchenne dystrophy) may lead to secondary loss of the sarcoglycans and dystroglycan. The primary loss of a single sarcoglycan (see “Limb-Girdle Muscular Dystrophy,” below) results in a secondary loss of other sarcoglycans in the membrane without uniformly affecting dystrophin. In either instance, disruption of the dystrophin-glycoprotein complexes weakens the sarcolemma, causing membrane tears and a cascade of events leading to muscle fiber necrosis. This sequence of events occurs repeatedly during the life of a patient with muscular dystrophy.

TREATMENT DUCHENNE MUSCULAR DYSTROPHY

Glucocorticoids, administered as prednisone in a dose of 0.75 mg/kg per day, significantly slow progression of Duchenne dystrophy for up to 3 years. Some patients cannot tolerate glucocorticoid therapy; weight gain and increased risk of fractures in particular represent a significant deterrent for some boys. As in other recessively inherited dystrophies presumed to arise from loss of function of a critical muscle gene, there is optimism that Duchenne disease may benefit from novel therapies that either replace the defective gene or missing protein or implement downstream corrections (e.g., skipping mutated exons or reading through mutations that introduce stop codons).

BECKER MUSCULAR DYSTROPHY

This less severe form of X-linked recessive muscular dystrophy results from allelic defects of the same gene responsible for Duchenne dystrophy. Becker muscular dystrophy is ~10 times less frequent than Duchenne.

Clinical Features The pattern of muscle wasting in Becker muscular dystrophy closely resembles that seen in Duchenne. Proximal muscles, especially of the lower extremities, are prominently involved. As the disease progresses, weakness becomes more generalized. Significant facial muscle weakness is not a feature. Hypertrophy of muscles, particularly in the calves, is an early and prominent finding.

Most patients with Becker dystrophy first experience difficulties between ages 5 and 15 years, although onset in the third or fourth decade or even later can occur. By definition, patients with Becker dystrophy walk beyond age 15, whereas patients with Duchenne dystrophy are typically in a wheelchair by the age of 12. Patients with Becker dystrophy have a reduced life expectancy, but most survive into the fourth or fifth decade.

Mental retardation may occur in Becker dystrophy, but it is not as common as in Duchenne. Cardiac involvement occurs in Becker dystrophy and may result in heart failure; some patients manifest with only heart failure. Other less common presentations are asymptomatic hyper-CK-emia, myalgias without weakness, and myoglobinuria.

Laboratory Features Serum CK levels, results of EMG, and muscle biopsy findings closely resemble those in Duchenne dystrophy. The diagnosis of Becker muscular dystrophy requires Western blot analysis of muscle biopsy samples, demonstrating a reduced amount or abnormal size of dystrophin or mutation analysis of DNA from peripheral blood leukocytes. Genetic testing reveals deletions or duplications of the dystrophin gene in 65% of patients with Becker dystrophy, approximately the same percentage as in Duchenne dystrophy. In both Becker and Duchenne dystrophies, the size of the DNA deletion does not predict clinical severity; however, in ~95% of patients with Becker dystrophy, the DNA deletion does not alter the translational reading frame of messenger RNA. These “in-frame” mutations allow for production of some dystrophin, which accounts for the presence of altered rather than absent dystrophin on Western blot analysis.

TREATMENT BECKER MUSCULAR DYSTROPHY

The use of glucocorticoids has not been adequately studied in Becker dystrophy.

LIMB-GIRDLE MUSCULAR DYSTROPHY

The syndrome of LGMD represents more than one disorder. Both males and females are affected, with onset ranging from late in the first decade to the fourth decade. The LGMDs typically manifest with progressive weakness of pelvic and shoulder girdle musculature. Respiratory insufficiency from weakness of the diaphragm may occur, as may cardiomyopathy.

A systematic classification of LGMD is based on autosomal dominant (LGMD1) and autosomal recessive (LGMD2) inheritance. Superimposed on the backbone of LGMD1 and LGMD2, the classification uses a sequential alphabetical lettering system (LGMD1A, LGMD2A, etc.). Disorders receive letters in the order in which they are found to have chromosomal linkage. This results in an ever-expanding list of conditions summarized in Tables 462e-6 and 462e-7. None of the conditions is as common as the dystrophinopathies; however, prevalence data for the LGMDs have not been systematically gathered for any large heterogeneous population. In referral-based clinical populations, Fukutin-related protein (FKRP) deficiency (LGMD2I), calpainopathy (LGMD2A), anoctaminopathy (LGMD2L), and to a lesser extent dysferlinopathy (LGMD2B) have emerged as the most common disorders.

EMERY-DREIFUSS MUSCULAR DYSTROPHY

There are at least five genetically distinct forms of Emery-Dreifuss muscular dystrophy (EDMD). Emerin mutations are the most common cause of X-linked EDMD, although mutations in *FHL1* may also be associated with a similar phenotype, which is X-linked as well. Mutations involving the gene for lamin A/C are the most common cause of autosomal dominant EDMD (also known as LGMD1B) and are also a common cause of hereditary cardiomyopathy. Less commonly, autosomal dominant EDMD has been reported with mutations in nesprin-1, nesprin-2, and *TMEM43*.

Clinical Features Prominent contractures can be recognized in early childhood and teenage years, often preceding muscle weakness. The contractures persist throughout the course of the disease and are present at the elbows, ankles, and neck. Muscle weakness affects humeral and peroneal muscles at first and later spreads to a limb-girdle distribution. The cardiomyopathy is potentially life threatening and may result in sudden death. A spectrum of atrial rhythm and conduction defects includes atrial fibrillation and paralysis and atrioventricular heart block. Some patients have a dilated cardiomyopathy. Female carriers of the X-linked variant may have cardiac manifestations that become clinically significant.

Laboratory Features Serum CK may be elevated two- to tenfold. EMG is myopathic. Muscle biopsy usually shows nonspecific dystrophic features, although cases associated with *FHL1* mutations have features of myofibrillar myopathy. Immunohistochemistry reveals absent emerin staining of myonuclei in X-linked EDMD due to emerin mutations. ECGs demonstrate atrial and atrioventricular rhythm disturbances.

X-linked EDMD usually arises from defects in the emerin gene encoding a nuclear envelope protein. *FHL1* mutations are also a cause of X-linked scapuloperoneal dystrophy, but can also present with an X-linked form of EDMD. The autosomal dominant disease can be caused by mutations in the *LMNA* gene encoding lamin A and C; in the synaptic nuclear envelope protein 1 (*SYNE1*) or 2 (*SYNE2*) encoding nesprin-1 and nesprin-2, respectively; and most recently in *TMEM43* encoding LUMA. These proteins are essential components of the filamentous network underlying the inner nuclear membrane. Loss of structural integrity of the nuclear envelope from defects in emerin, lamin A/C, nesprin-1, nesprin-2, and LUMA accounts for overlapping phenotypes (Fig. 462e-7).

TREATMENT EMERY-DREIFUSS MUSCULAR DYSTROPHY

Supportive care should be offered for neuromuscular disability, including ambulatory aids, if necessary. Stretching of contractures is difficult. Management of cardiomyopathy and arrhythmias (e.g., early use of a defibrillator or cardiac pacemaker) may be life saving.

CONGENITAL MUSCULAR DYSTROPHY (CMD)

This is not one entity but rather a group of disorders with varying degrees of muscle weakness, CNS impairment, and eye abnormalities.

Clinical Features As a group, CMDs present at birth or in the first few months of life with hypotonia and proximal or generalized muscle weakness. Calf muscle hypertrophy is seen in some patients. Facial muscles may be weak, but other cranial nerve-innervated muscles are spared (e.g., extraocular muscles are normal). Most patients have joint contractures of varying degrees at elbows, hips, knees, and ankles. Contractures present at birth are referred to as *arthrogryposis*. Respiratory failure may be seen in some cases.

The CNS is affected in some forms of CMD. In merosin and FKR deficiency, cerebral hypomyelination may be seen by magnetic resonance imaging (MRI), although only a small number of patients have mental retardation and seizures. Three forms of congenital muscular dystrophy have severe brain impairment. These include Fukuyama's congenital muscular dystrophy (FCMD), muscle-eye-brain (MEB) disease, and Walker-Warburg syndrome (WWS). Patients are severely disabled in all three of these conditions. In MEB disease and WWS, but

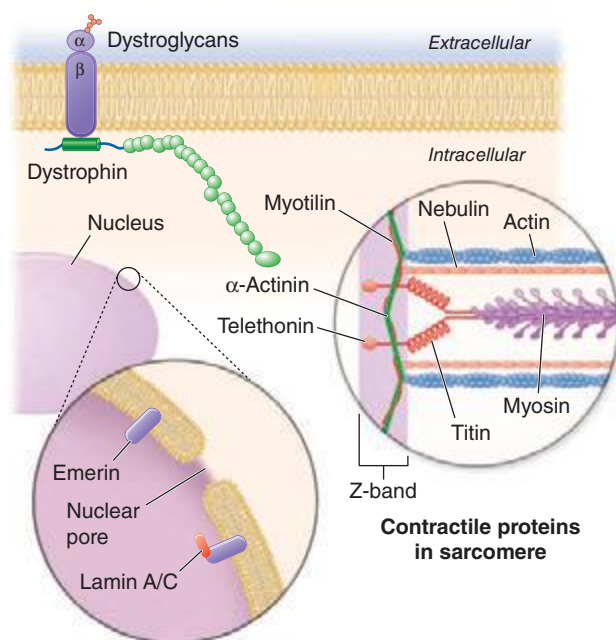


FIGURE 462e-7 Selected muscular dystrophy–associated proteins in the nuclear membrane and sarcomere. As shown in the exploded view, emerin and lamin A/C are constituents of the inner nuclear membrane. Several dystrophy-associated proteins are represented in the sarcomere including titin, nebulin, calpain, telethonin, actinin, and myotilin. The position of the dystrophin-dystroglycan complex is also illustrated.

not in FCMD, ocular abnormalities impair vision. WWS is the most severe congenital muscular dystrophy, causing death by 1 year of age.

Laboratory Features Serum CK is markedly elevated in all of these conditions. The EMG is myopathic and muscle biopsies show nonspecific dystrophic features. Merosin, or laminin α_2 chain (a basal lamina protein), is deficient in surrounding muscle fibers in merosin deficiency. Skin biopsies can also demonstrate defects in laminin α_2 chain. In the other disorders (FKRP deficiency, FCMD, MEB disease, WWS), there is abnormal α -dystroglycan staining in muscle. In merosin deficiency, cerebral hypomyelination is common, and a host of brain malformations are seen in FCMD, MEB disease, and WWS.

All forms of CMD are inherited as autosomal recessive disorders. Chromosomal linkage and specific gene defects are presented in Table 462e-8. With the exception of merosin, the other gene defects affect posttranslational glycosylation of α -dystroglycan. This abnormality is thought to impair binding with merosin and leads to weakening of the dystrophin-glycoprotein complex, instability of the muscle membrane, and/or abnormalities in muscle contraction. CMDs with brain and eye phenotypes probably involve defective glycosylation of additional proteins, accounting for the more extensive phenotypes.

TREATMENT CONGENITAL MUSCULAR DYSTROPHY

There is no specific treatment for CMD. Proper wheelchair seating is important. Management of epilepsy and cardiac manifestations is necessary for some patients.

MYOTONIC DYSTROPHY

Myotonic dystrophy is also known as *dystrophia myotonica* (DM). The condition is composed of at least two clinical disorders with overlapping phenotypes and distinct molecular genetic defects: myotonic dystrophy type 1 (DM1), the classic disease originally described by Steinert, and myotonic dystrophy type 2 (DM2), also called *proximal myotonic myopathy* (PROMM).

TABLE 462e-8 CONGENITAL MUSCULAR DYSTROPHIES^a

Disease	Clinical Features	Laboratory Features	Abnormal Protein
Merosin deficiency	Onset at birth with hypotonia, joint contractures, delayed milestones, generalized muscle weakness Cerebral hypomyelination, less often cortical dysplasia Normal intelligence usually, some with MR (~6%) and seizures (~8%) Partial deficiency leads to milder phenotype (LGMD picture)	Serum CK 5–35× normal EMG myopathic NCS abnormal in some cases	Laminin α_2 chain
Fukutin-related protein deficiency ^b	Onset at birth or shortly after Hypotonia and feeding problems Weakness of proximal muscles, especially shoulder girdles Hypertrophy of leg muscles Joint contractures Cognition normal	Serum CK 10–50× normal EMG myopathic NCS normal	Fukutin-related protein
Fukuyama's congenital muscular dystrophy ^b	Onset at birth Hypotonia, joint contractures Generalized muscle weakness Hypertrophy of calf muscles Seizures, MR Cardiomyopathy	Serum CK 10–50× normal EMG myopathic NCS normal MRI shows hydrocephalus and periventricular and frontal hypomyelination	Fukutin
Muscle-eye-brain disease	Onset at birth, hypotonia Eye abnormalities include: progressive myopia, cataracts, and optic nerve, glaucoma, retinal pigmentary changes Progressive muscle weakness Joint contractures Seizures, MR	Serum CK 5–20× normal MRI shows hydrocephalus, cobblestone lissencephaly, corpus callosum and cerebellar hypoplasia, cerebral hypomyelination	N-Acetyl-glucosaminyl transferase (POMGnT1)
Walker-Warburg syndrome ^b	Onset at birth, hypotonia Generalized muscle weakness Joint contractures Microphthalmos, retinal dysplasia, buphthalmos, glaucoma, cataracts Seizures, MR	Serum CK 5–20× normal MRI shows cobblestone lissencephaly, hydrocephalus, encephalocele, absent corpus callosum	O-Mannosyl-transferase-1 (POMT1)

^aAll are inherited as recessive traits. ^bThere is phenotypic overlap between disorders related to defective glycosylation. In muscle, this is a consequence of altered glycosylation of dystroglycans; in brain/eye, other glycosylated proteins are involved. Clinically, Walker-Warburg syndrome is more severe, with death by 1 year.

Abbreviations: CK, creatine kinase; EMG, electromyography; LGMD, limb-girdle muscular dystrophy; MR, mental retardation; MRI, magnetic resonance imaging; NCS, nerve conduction studies.

Clinical Features The clinical expression of DM1 varies widely and involves many systems other than muscle. Affected patients have a typical “hatchet-faced” appearance due to temporalis, masseter, and facial muscle atrophy and weakness. Frontal baldness is also characteristic of the disease. Neck muscles, including flexors and sternocleidomastoids, and distal limb muscles are involved early. Weakness of wrist extensors, finger extensors, and intrinsic hand muscles impairs function. Ankle dorsiflexor weakness may cause footdrop. Proximal muscles remain stronger throughout the course, although preferential atrophy and weakness of quadriceps muscles occur in many patients. Palatal, pharyngeal, and tongue involvement produce a dysarthric speech, nasal voice, and swallowing problems. Some patients have diaphragm and intercostal muscle weakness, resulting in respiratory insufficiency.

Myotonia, which usually appears by age 5 years, is demonstrable by percussion of the thenar eminence, the tongue, and wrist extensor muscles. Myotonia causes a slow relaxation of hand grip after a forced voluntary closure. Advanced muscle wasting makes myotonia more difficult to detect.

Cardiac disturbances occur commonly in patients with DM1. ECG abnormalities include first-degree heart block and more extensive conduction system involvement. Complete heart block and sudden death can occur. Congestive heart failure occurs infrequently but may result from cor pulmonale secondary to respiratory failure. Mitral valve prolapse also occurs commonly. Other associated features include intellectual impairment, hypersomnia, posterior subcapsular cataracts, gonadal atrophy, insulin resistance, and decreased esophageal and colonic motility.

Congenital myotonic dystrophy is a more severe form of DM1 and occurs in ~25% of infants of affected mothers. It is characterized by severe facial and bulbar weakness, transient neonatal respiratory insufficiency, and mental retardation.

DM2, or PROMM, has a distinct pattern of muscle weakness affecting mainly proximal muscles. Other features of the disease overlap with DM1, including cataracts, testicular atrophy, insulin resistance, constipation, hypersomnia, and cognitive defects. Cardiac conduction defects occur but are less common, and the hatchet face and frontal baldness are less consistent features. A very striking difference is the failure to clearly identify a congenital form of DM2.

Laboratory Features The diagnosis of myotonic dystrophy can usually be made on the basis of clinical findings. Serum CK levels may be normal or mildly elevated. EMG evidence of myotonia is present in most cases of DM1 but may be more patchy in DM2. Muscle biopsy shows muscle atrophy, which selectively involves type 1 fibers in 50% of cases, and ringed fibers in DM1 but not in DM2. Typically, numerous internalized nuclei can be seen in individual muscle fibers as well as atrophic fibers with pyknotic nuclear clumps in both DM1 and DM2. Necrosis of muscle fibers and increased connective tissue, common in other muscular dystrophies, are less apparent in myotonic dystrophy.

DM1 and DM2 are both autosomal dominant disorders. New mutations do not appear to contribute to the pool of affected individuals. DM1 is transmitted by an intronic mutation consisting of an unstable expansion of a CTG trinucleotide repeat in a serine-threonine protein

kinase gene (named *DMPK*) on chromosome 19q13.3. An increase in the severity of the disease phenotype in successive generations (genetic anticipation) is accompanied by an increase in the number of trinucleotide repeats. A similar type of mutation has been identified in fragile X syndrome (Chap. 451e). The unstable triplet repeat in myotonic dystrophy can be used for prenatal diagnosis. Congenital disease occurs almost exclusively in infants born to affected mothers; it is possible that sperm with greatly expanded triplet repeats do not function well.

DM2 is caused by a DNA expansion mutation consisting of a CCTG repeat in intron 1 of the *ZNF9* gene located at chromosome 3q13.3-q24. The gene is believed to encode an RNA-binding protein expressed in many different tissues, including skeletal and cardiac muscle.

The DNA expansions in DM1 and DM2 almost certainly impair muscle function by a toxic gain of function of the mutant mRNA. In both DM1 and DM2, the mutant RNA appears to form intranuclear inclusions composed of aberrant RNA. These RNA inclusions sequester RNA-binding proteins essential for proper splicing of a variety of other mRNAs. This leads to abnormal transcription of multiple proteins in a variety of tissues/organ systems, in turn causing the systemic manifestations of DM1 and DM2.

TREATMENT MYOTONIC DYSTROPHY

The myotonia in DM1 rarely warrants treatment, although some patients with DM2 are significantly bothered by the discomfort related to the associated muscle stiffness. Phenytoin and mexiletine are the preferred agents for the occasional patient who requires an antimyotonia drug; other agents, particularly quinine and procainamide, may worsen cardiac conduction. A cardiac pacemaker should be considered for patients with unexplained syncope, advanced conduction system abnormalities with evidence of second-degree heart block, or trifascicular conduction disturbances with marked prolongation of the PR interval. Molded ankle-foot orthoses help stabilize gait in patients with foot drop. Excessive daytime somnolence with or without sleep apnea is not uncommon. Sleep studies, noninvasive respiratory support (biphasic positive airway pressure [BiPAP]), and treatment with modafinil may be beneficial.

FACIOSCAPULOHUMERAL (FSH) MUSCULAR DYSTROPHY

This form of muscular dystrophy has a prevalence of ~1 in 20,000. There are two forms of FSHD that have similar pathogenesis, as will be discussed. Most patients have FSHD type 1 (95%), whereas approximately 5% have FSHD2. FSHD1 and FSHD2 are clinically and histopathologically identical. FSHD is not to be confused with the genetically distinct scapuloperoneal dystrophies.

Clinical Features The condition typically has an onset in childhood or young adulthood. In most cases, facial weakness is the initial manifestation, appearing as an inability to smile, whistle, or fully close the eyes. Weakness of the shoulder girdles, rather than the facial muscles, usually brings the patient to medical attention. Loss of scapular stabilizer muscles makes arm elevation difficult. Scapular winging (Fig. 462e-3) becomes apparent with attempts at abduction and forward movement of the arms. Biceps and triceps muscles may be severely affected, with relative sparing of the deltoid muscles. Weakness is invariably worse for wrist extension than for wrist flexion, and weakness of the anterior compartment muscles of the legs may lead to footdrop.

In most patients, the weakness remains restricted to facial, upper extremity, and distal lower extremity muscles. In 20% of patients, weakness progresses to involve the pelvic girdle muscles, and severe functional impairment and possible wheelchair dependency result.

Characteristically, patients with FSHD do not have involvement of other organ systems, although labile hypertension is common, and there is an increased incidence of nerve deafness. *Coats' disease*, a disorder consisting of telangiectasia, exudation, and retinal detachment, also occurs.

Laboratory Features The serum CK level may be normal or mildly elevated. EMG usually indicates a myopathic pattern. The muscle biopsy

shows nonspecific features of a myopathy. A prominent inflammatory infiltrate, which is often multifocal in distribution, is present in some biopsy samples. The cause or significance of this finding is unknown.

An autosomal dominant inheritance pattern with almost complete penetrance has been established, but each family member should be examined for the presence of the disease, since ~30% of those affected are unaware of involvement. FSHD1 is associated with deletions of tandem 3.3-kb repeats at 4q35. The deletion reduces the number of repeats to a fragment of <35 kb in most patients. Within these repeats lies the *DUX4* gene, which usually is not expressed. In patients with FSHD1 these deletions in the setting of a specific polymorphism leads to hypomethylation of the region and toxic expression of the *DUX4* gene. In patients with FSHD2, there is no deletion, but a mutation in *SMCHD1*. Interestingly, in the setting of the same polymorphism, there again is seen hypomethylation of the region and the permissive expression of the *DUX4* gene. In both FSHD1 and FSHD2, there is overexpression of the *DUX4* transcript.

TREATMENT FACIOSCAPULOHUMERAL MUSCULAR DYSTROPHY

No specific treatment is available; ankle-foot orthoses are helpful for footdrop. Scapular stabilization procedures improve scapular winging but may not improve function.

OCULOPHARYNGEAL DYSTROPHY

This form of muscular dystrophy represents one of several disorders characterized by progressive external ophthalmoplegia, which consists of slowly progressive ptosis and limitation of eye movements with sparing of pupillary reactions for light and accommodation. Patients usually do not complain of diplopia, in contrast to patients having conditions with a more acute onset of ocular muscle weakness (e.g., myasthenia gravis).

Clinical Features Oculopharyngeal muscular dystrophy has a late onset; it usually presents in the fourth to sixth decade with ptosis and/or dysphagia. The extraocular muscle impairment is less prominent in the early phase but may be severe later. The swallowing problem may become debilitating and result in pooling of secretions and repeated episodes of aspiration. Mild weakness of the neck and extremities also occurs.

Laboratory Features The serum CK level may be two to three times normal. Myopathic EMG findings are typical. On biopsy, muscle fibers are found to contain rimmed vacuoles, which by electron microscopy are shown to contain membranous whorls, accumulation of glycogen, and other nonspecific debris related to lysosomes. A distinct feature of oculopharyngeal dystrophy is the presence of tubular filaments, 8.5 nm in diameter, in muscle cell nuclei.

Oculopharyngeal dystrophy has an autosomal dominant inheritance pattern with complete penetrance. The incidence is high in French-Canadians and in Spanish-American families of the southwestern United States. Large kindreds of Italian and of eastern European Jewish descent have been reported. The molecular defect in oculopharyngeal muscular dystrophy is a subtle expansion of a modest polyalanine repeat tract in a poly-RNA-binding protein (PABP2) in muscle.

TREATMENT OCULOPHARYNGEAL DYSTROPHY

Dysphagia can lead to significant undernourishment and inanition, making oculopharyngeal muscular dystrophy a potentially life-threatening disease. Cricopharyngeal myotomy may improve swallowing, although it does not prevent aspiration. Eyelid crutches can improve vision when ptosis obstructs vision; candidates for ptosis surgery must be carefully selected—those with severe facial weakness are not suitable.

A group of muscle diseases, the distal myopathies, are notable for their preferential distal distribution of muscle weakness in contrast to most muscle conditions associated with proximal weakness. The major distal myopathies are summarized in [Table 462e-9](#).

Clinical Features *Welander's*, *Udd's*, and *Markesbery-Griggs type distal myopathies* are all late-onset, dominantly inherited disorders of distal limb muscles, usually beginning after age 40 years. *Welander's* distal myopathy preferentially involves the wrist and finger extensors, whereas the others are associated with anterior tibial weakness leading to progressive footdrop. *Laing's distal myopathy* is also a dominantly inherited disorder heralded by tibial weakness; however, it is distinguished by onset in childhood or early adult life. *Nonaka's distal myopathy* and *Miyoshi's myopathy* are distinguished by autosomal recessive inheritance and onset in the late teens or twenties. *Nonaka's* and *Williams' myopathy* entails anterior tibial weakness, whereas *Miyoshi's myopathy* is unique in that gastrocnemius muscles are preferentially affected at onset. Finally, the *myofibrillar myopathies* (MFMs) are a clinically and genetically heterogeneous group of disorders that can be associated with prominent distal weakness; they can be inherited in an autosomal dominant or recessive pattern. Of note, *Markesbery-Griggs myopathy* (caused by mutations in *ZASP*) and *LGMD1B* (caused by mutations in *myotilin*) are in fact subtypes of myofibrillar myopathy.

Laboratory Features Serum CK level is particularly helpful in diagnosing *Miyoshi's myopathy* because it is very elevated. In the other conditions, serum CK is only slightly increased. EMGs are myopathic. In the MFMs, myotonic or pseudomyotonic discharges are common. Muscle biopsy shows nonspecific dystrophic features and, with the exception of *Laing's* and *Miyoshi's myopathies*, often shows rimmed vacuoles. MFM is associated with the accumulation of dense inclusions, as well as amorphous material best seen on Gomori trichrome and myofibrillar disruption on electron microscopy. Immune staining sometimes demonstrates accumulation of desmin and other proteins in MFM, large deposits of myosin heavy chain in the subsarcolemmal region of type 1 muscle fibers in *Laing's myopathy*, and reduced or absent dysferlin in *Miyoshi's myopathy*.

The affected genes and their gene products are listed in [Table 462e-9](#).

TREATMENT **DISTAL MYOPATHIES**

Occupational therapy is offered for loss of hand function; ankle-foot orthoses can support distal lower limb muscles. The MFMs can be associated with cardiomyopathy (congestive heart failure or arrhythmias) and respiratory failure that may require medical management. *Laing's-type distal myopathy* can also be associated with a cardiomyopathy.

CONGENITAL MYOPATHIES

These rare disorders are distinguished from muscular dystrophies by the presence of specific histochemical and structural abnormalities in muscle. Although primarily disorders of infancy or childhood, three forms that may present in adulthood are described here: central core disease, nemaline (rod) myopathy, and centronuclear (myotubular) myopathy. Sarcotubular myopathy is caused by mutations in *TRIM-32* and is identical to *LGMD2H*. Other types, such as minicore myopathy (multi-minicore disease), fingerprint body myopathy, and cap myopathy, are not discussed.

CENTRAL CORE DISEASE

Patients with central core disease may have decreased fetal movements and breech presentation. Hypotonia and delay in motor milestones, particularly in walking, are common. Later in childhood, patients develop problems with stair climbing, running, and getting up from the floor. On examination, there is mild facial, neck-flexor, and proximal-extremity muscle weakness. Legs are more affected than arms. Skeletal abnormalities include congenital hip dislocation, scoliosis,

and pes cavus; clubbed feet also occur. Most cases are nonprogressive, but exceptions are well documented. Susceptibility to malignant hyperthermia must be considered as a potential risk factor for patients with central core disease. Recent series have demonstrated that many cases of late-onset axial myopathy in which patients manifest with bent spine (camptocormia) or neck extensor weakness (neck extensor myopathy) are caused by mutations in the ryanodine receptor gene (*RYR1*). This illustrates the interesting spectrum of *RYR1* mutations.

The serum CK level is usually normal. Needle EMG demonstrates a myopathic pattern. Muscle biopsy shows fibers with single or multiple central or eccentric discrete zones (*cores*) devoid of oxidative enzymes. Cores occur preferentially in type 1 fibers and represent poorly aligned sarcomeres associated with Z disk streaming.

Autosomal dominant inheritance is characteristic; sporadic cases also occur. As alluded above, this myopathy is caused by point mutations of *RYR1*, encoding the calcium-release channel of the sarcoplasmic reticulum of skeletal muscle; mutations of this gene also account for some cases of inherited malignant hyperthermia ([Chap. 23](#)). Malignant hyperthermia is an allelic condition; C-terminal mutations of the *RYR1* gene predispose to this complication.

Specific treatment is not required, but establishing a diagnosis of central core disease is extremely important because these patients have a known predisposition to malignant hyperthermia during anesthesia.

NEMALINE MYOPATHY

The term *nemaline* refers to the distinctive presence in muscle fibers of rods or threadlike structures (Greek *nema*, “thread”). Nemaline myopathy is clinically heterogeneous. A severe neonatal form presents with hypotonia and feeding and respiratory difficulties, leading to early death. Nemaline myopathy usually presents in infancy or childhood with delayed motor milestones. The course is nonprogressive or slowly progressive. The physical appearance is striking because of the long, narrow facies, high-arched palate, and open-mouthed appearance due to a prognathous jaw. Other skeletal abnormalities include pectus excavatum, kyphoscoliosis, pes cavus, and clubfoot deformities. Facial and generalized muscle weakness, including respiratory muscle weakness, is common. An adult-onset disorder with progressive proximal or distal weakness may be seen. Myocardial involvement is occasionally present in both the childhood and adult-onset forms. The serum CK level is usually normal or slightly elevated. The EMG demonstrates a myopathic pattern. Muscle biopsy shows clusters of small rods (nemaline bodies), which occur preferentially, but not exclusively, in the sarcoplasm of type 1 muscle fibers. Occasionally, the rods are also apparent in myonuclei. The muscle often shows type 1 muscle fiber predominance. Rods originate from the Z disk material of the muscle fiber.

Six genes have been associated with nemaline myopathy. Five of these code for thin filament-associated proteins, suggesting disturbed assembly or interplay of these structures as a pivotal mechanism. Mutations of the nebulin (*NEB*) gene account for most cases, including both severe neonatal and early childhood forms, inherited as autosomal recessive disorders. Neonatal and childhood cases, inherited as predominantly autosomal dominant disorders, are caused by mutations of the skeletal muscle α -actinin (*ACTA1*) gene. In milder forms of the disease with autosomal dominant inheritance, mutations have been identified in both the slow α -tropomyosin (*TPM3*) and β -tropomyosin (*TPM2*) genes accounting for <3% of cases. Muscle troponin T (*TNNT1*) gene mutations appear to be limited to the Amish population in North America. Mutations may also be seen in *NEM6* that encodes a putative BTB/Kelch protein. No specific treatment is available.

CENTRONUCLEAR (MYOTUBULAR) MYOPATHY

Three distinct variants of centronuclear myopathy occur. A neonatal form, also known as *myotubular myopathy*, presents with severe hypotonia and weakness at birth. The late infancy-early childhood form presents with delayed motor milestones. Later, difficulty with running and stair climbing becomes apparent. A marfanoid, slender body habitus, long narrow face, and high-arched palate are typical. Scoliosis and

TABLE 462e-9 DISTAL MYOPATHIES

Disease	Clinical Features	Laboratory Features	Abnormal Protein
Welander's distal myopathy	Onset in fifth decade Weakness begins in hands Slow progression with spread to distal lower extremities Lifespan normal	Serum CK 2–3× normal EMG myopathic NCS normal Muscle biopsy shows dystrophic features and rimmed vacuoles	AD Chromosome 2p13 TIA1
Tibial muscular dystrophy (Udd's)	Onset fourth to eighth decade Distal lower extremity weakness (tibial distribution) Upper extremities usually normal Lifespan normal	Serum CK 2–4× normal EMG myopathic NCS normal Muscle biopsy shows dystrophic features and rimmed vacuoles Titin absent in M-line of muscle	AD Titin AR (associated with more proximal weakness—LGMD2J)
Markesbery-Griggs distal myopathy	Onset fourth to eighth decade Distal lower extremity weakness (tibial distribution) with progression to distal arms and proximal muscles	Serum CK is usually mildly elevated EMG reveals irritative myopathy Muscle biopsies demonstrate rimmed vacuoles and features of MFM	AD Z-band alternatively spliced PDX motif-containing protein (ZASP)
Laing's distal myopathy	Onset childhood to third decade Distal lower extremity weakness (anterior tibial distribution) and neck flexors affected early May have cardiomyopathy	Serum CK is normal or slightly elevated Muscle biopsies do not typically show rimmed vacuoles, but may show hyaline bodies with accumulation of myosin Large deposits of myosin heavy chain are seen in type 1 muscle fibers	AD Myosin heavy chain 7
Nonaka's distal myopathy (autosomal recessive hereditary inclusion body myopathy)	Onset: second to third decade Distal lower extremity weakness (anterior tibial distribution) Mild distal upper limb weakness may be present early Progression to other muscles sparing quadriceps Ambulation may be lost in 10–15 y	Serum CK 3–10× normal EMG myopathic NCS normal Dystrophic features on muscle biopsy plus rimmed vacuoles and 15- to 19-nm filaments within vacuoles	AR GNE gene: UDP-N-acetylglucosamine 2-epimerase/N-acetylmannosamine kinase Allelic to hereditary inclusion body myopathy
Miyoshi's myopathy ^a	Onset: second to third decade Lower extremity weakness in posterior compartment muscles Progression leads to weakness in other muscle groups Ambulation lost after 10–15 y in about one-third of cases	Serum CK 20–100× normal EMG myopathic NCS normal Muscle biopsy shows nonspecific dystrophic features often with prominent inflammatory cell infiltration; no rimmed vacuoles	AR Allelic to LGMD2B (see Table 462e-7) Dysferlin
Williams' myopathy	Distal lower extremity weakness (anterior tibial distribution)	Muscle biopsy may show rimmed vacuoles and features of MFM	X-linked Filamin-C
Myofibrillar myopathies	Onset from early childhood to late adult life Weakness may be distal, proximal, or generalized Cardiomyopathy and respiratory involvement is not uncommon	Serum CKs can be normal or moderately elevated EMG is myopathic and often associated with myotonic discharges Muscle biopsy demonstrates abnormal accumulation of desmin and other proteins, rimmed vacuoles, and myofibrillar degeneration	Genetically heterogeneous AD Myotilin (also known as LGMD 1A) ZASP (see Markesbery-Griggs distal myopathy) Filamin-C Desmin Alpha B crystallin Bag3 Titin DNAJB6 TNPO3 AR Desmin X-linked FHL1

^aMiyoshi's myopathy phenotype may also be seen with mutations in ANO-5 that encodes for anoctamin 5 (allelic to LGMD2L).**Abbreviations:** AD, autosomal dominant; AR, autosomal recessive; CK, creatine kinase; EMG, electromyography; MFM, myofibrillar myopathy; NCS, nerve conduction studies.

462e-14 clubbed feet may be present. Most patients exhibit progressive weakness, some requiring wheelchairs. Progressive external ophthalmoplegia with ptosis and varying degrees of extraocular muscle impairment are characteristic of both the neonatal and the late-infantile forms. A third variant, the late childhood–adult form, has an onset in the second or third decade. Patients have full extraocular muscle movements and rarely exhibit ptosis. There is mild, slowly progressive limb weakness that may be distally predominant (some of these patients have been classified as having Charcot-Marie-Tooth disease type 2 [CMT2; Chap. 459]).

Normal or slightly elevated CK levels occur in each of the forms. Nerve conduction studies may reveal reduced amplitudes of distal compound muscle action potentials, in particular in adult-onset cases that resemble CMT2. EMG studies often give distinctive results, showing positive sharp waves and fibrillation potentials, complex and repetitive discharges, and rarely myotonic discharges. Muscle biopsy specimens in longitudinal section demonstrate rows of central nuclei, often surrounded by a halo. In transverse sections, central nuclei are found in 25–80% of muscle fibers.

A gene for the neonatal form of centronuclear myopathy has been localized to Xq28; this gene encodes myotubularin, a protein tyrosine phosphatase. Missense, frameshift, and splice-site mutations predict loss of myotubularin function in affected individuals. Carrier identification and prenatal diagnosis are possible. Autosomal recessive forms are caused by mutations in *BIB1* that encodes for amphiphysin-2, whereas some autosomal dominant cases, which are allelic to a form of CMT2, are associated with mutations in the gene that encodes dynamin-2. No specific medical treatments are available at this time.

DISORDERS OF MUSCLE ENERGY METABOLISM

There are two principal sources of energy for skeletal muscle—fatty acids and glucose. Abnormalities in either glucose or lipid utilization can be associated with distinct clinical presentations that can range from an acute, painful syndrome with rhabdomyolysis and myoglobinuria to a chronic, progressive muscle weakness simulating muscular dystrophy.

GLYCOGEN STORAGE AND GLYCOLYTIC DEFECTS

Disorders of Glycogen Storage Causing Progressive Weakness • α -GLUCOSIDASE, OR ACID MALTASE, DEFICIENCY (POMPE'S DISEASE) Three clinical forms of α -glucosidase, or acid maltase, deficiency (*type II glycogenosis*) can be distinguished. The infantile form is the most common, with onset of symptoms in the first 3 months of life. Infants develop severe muscle weakness, cardiomegaly, hepatomegaly, and respiratory insufficiency. Glycogen accumulation in motor neurons of the spinal cord and brainstem contributes to muscle weakness. Death usually occurs by 1 year of age. In the childhood form, the picture resembles muscular dystrophy. Delayed motor milestones result from proximal limb muscle weakness and involvement of respiratory muscles. The heart may be involved, but the liver and brain are unaffected. The adult form usually begins in the third or fourth decade but can present as late as the seventh decade. Respiratory failure and diaphragmatic weakness are often initial manifestations, heralding progressive proximal muscle weakness. The heart and liver are not involved.

The serum CK level is 2–10 times normal in infantile or childhood-onset Pompe's disease but can be normal in adult-onset cases. EMG examination demonstrates a myopathic pattern, but other features are especially distinctive, including myotonic discharges, trains of fibrillation and positive waves, and complex repetitive discharges. EMG discharges are very prominent in the paraspinal muscles. The muscle biopsy in infants typically reveals vacuoles containing glycogen and the lysosomal enzyme acid phosphatase. Electron microscopy reveals membrane-bound and free tissue glycogen. However, muscle biopsies in late-onset Pompe's disease may demonstrate only nonspecific abnormalities. Enzyme analysis of dried blood spots is a sensitive technique to screen for Pompe's disease. A definitive diagnosis is established by enzyme assay in muscle or cultured fibroblasts or by genetic testing.

Pompe's disease is inherited as an autosomal recessive disorder caused by mutations of the α -glucosidase gene. Enzyme replacement therapy (ERT) with IV recombinant human α -glucosidase has been shown to be beneficial in infantile-onset Pompe's disease. Clinical benefits in the infantile disease include reduced heart size, improved muscle function, reduced need for ventilatory support, and longer life. In late-onset cases, ERT has not been associated with the dramatic response that can be seen in classic infantile Pompe's disease, yet it appears to stabilize the disease process.

OTHER GLYCOGEN STORAGE DISEASES WITH PROGRESSIVE WEAKNESS In *debranching enzyme deficiency (type III glycogenosis)*, a slowly progressive form of muscle weakness can develop after puberty. Rarely, myoglobinuria may be seen. Patients are usually diagnosed in infancy, however, because of hypotonia and delayed motor milestones, hepatomegaly, growth retardation, and hypoglycemia. *Branching enzyme deficiency (type IV glycogenosis)* is a rare and fatal glycogen storage disease characterized by failure to thrive and hepatomegaly. Hypotonia and muscle wasting may be present, but the skeletal muscle manifestations are minor compared to liver failure.

Disorders of Glycolysis Causing Exercise Intolerance Several glycolytic defects are associated with recurrent myoglobinuria: *myophosphorylase deficiency (type V glycogenosis)*, *phosphofructokinase deficiency (type VII glycogenosis)*, *phosphoglycerate kinase deficiency, phosphorylase kinase deficiency (type IX glycogenosis)*, *phosphoglycerate mutase deficiency (type X glycogenosis)*, *lactate dehydrogenase deficiency (glycogenosis type XI)*, and *β -enolase deficiency*. Myophosphorylase deficiency, also known as *McArdle's disease*, is by far the most common of the glycolytic defects associated with exercise intolerance. These glycolytic defects result in a common failure to support energy production at the initiation of exercise, although the exact site of energy failure remains controversial.

Clinical muscle manifestations in these conditions usually begin in adolescence. Symptoms are precipitated by brief bursts of high-intensity exercise such as running or lifting heavy objects. A history of myalgia and muscle stiffness usually precedes the intensely painful muscle contractures, which may be followed by myoglobinuria. Acute renal failure accompanies significant pigmenturia.

Certain features help distinguish some enzyme defects. In McArdle's disease, exercise tolerance can be enhanced by a slow induction phase (warm-up) or brief periods of rest, allowing for the start of the "second-wind" phenomenon (switching to utilization of fatty acids). Varying degrees of hemolytic anemia accompany deficiencies of both phosphofructokinase (mild) and phosphoglycerate kinase (severe). In phosphoglycerate kinase deficiency, the usual clinical presentation is a seizure disorder associated with mental retardation; exercise intolerance is an infrequent manifestation.

In all of these conditions, the serum CK levels fluctuate widely and may be elevated even during symptom-free periods. CK levels >100 times normal are expected, accompanying myoglobinuria. All patients with suspected glycolytic defects leading to exercise intolerance should undergo a forearm exercise test. An impaired rise in venous lactate is highly indicative of a glycolytic defect. In lactate dehydrogenase deficiency, venous levels of lactate do not increase, but pyruvate rises to normal. A definitive diagnosis of glycolytic disease is made by muscle biopsy and subsequent enzyme analysis or by genetic testing.

Myophosphorylase deficiency, phosphofructokinase deficiency, and phosphoglycerate mutase deficiency are inherited as autosomal recessive disorders. Phosphoglycerate kinase deficiency is X-linked recessive. Mutations can be found in the respective genes encoding the abnormal proteins in each of these disorders.

Training may enhance exercise tolerance, perhaps by increasing perfusion to muscle. Dietary intake of free glucose or fructose prior to activity may improve function but care must be taken to avoid obesity from ingesting too many calories.

LIPID AS AN ENERGY SOURCE AND ASSOCIATED DEFECTS

Lipid is an important muscle energy source during rest and during prolonged, submaximal exercise. Fatty acids are derived from

circulating very-low-density lipoprotein (VLDL) in the blood or from triglycerides stored in muscle fibers. Oxidation of fatty acids occurs in the mitochondria. To enter the mitochondria, a fatty acid must first be converted to an “activated fatty acid,” acyl-CoA. The acyl-CoA must be linked with carnitine by the enzyme carnitine palmitoyltransferase (CPT) I for transport into the mitochondria. CPT I is present on the inner side of the outer mitochondrial membrane. Carnitine is removed by CPT II, an enzyme attached to the inside of the inner mitochondrial membrane, allowing transport of acyl-CoA into the mitochondrial matrix for β -oxidation.

Carnitine Palmitoyltransferase Deficiency CPT II deficiency is the most common recognizable cause of recurrent myoglobinuria, more common than the glycolytic defects. Onset is usually in the teenage years or early twenties. Muscle pain and myoglobinuria typically occur after prolonged exercise but can also be precipitated by fasting or infections; up to 20% of patients do not exhibit myoglobinuria, however. Strength is normal between attacks. In contrast to disorders caused by defects in glycolysis, in which muscle cramps follow short, intense bursts of exercise, the muscle pain in CPT II deficiency does not occur until the limits of utilization have been exceeded and muscle breakdown has already begun. Episodes of rhabdomyolysis may produce severe weakness. In young children and newborns, CPT II deficiency can present with a very severe clinical picture including hypoketotic hypoglycemia, cardiomyopathy, liver failure, and sudden death.

Serum CK levels and EMG findings are both usually normal between episodes. A normal rise of venous lactate during forearm exercise distinguishes this condition from glycolytic defects, especially myophosphorylase deficiency. Muscle biopsy does not show lipid accumulation and is usually normal between attacks. The diagnosis requires direct measurement of muscle CPT or genetic testing.

CPT II deficiency is much more common in men than women (5:1); nevertheless, all evidence indicates autosomal recessive inheritance. A mutation in the gene for CPT II (chromosome 1p36) causes the disease in some individuals. Attempts to improve exercise tolerance with frequent meals and a low-fat, high-carbohydrate diet, or by substituting medium-chain triglycerides in the diet, have not proven to be beneficial.

Myoadenylate Deaminase Deficiency The muscle enzyme myoadenylate deaminase converts adenosine-5'-monophosphate (5'-AMP) to inosine monophosphate (IMP) with liberation of ammonia. Myoadenylate deaminase may play a role in regulating adenosine triphosphate (ATP) levels in muscles. Most individuals with myoadenylate deaminase deficiency have no symptoms. There have been a few reports of patients with this disorder who have exercise-exacerbated myalgia and myoglobinuria. Many questions have been raised about the clinical effects of myoadenylate deaminase deficiency, and, specifically, its relationship to exertional myalgia and fatigability, but there is no consensus.

MITOCHONDRIAL MYOPATHIES

In 1972, Olson and colleagues recognized that muscle fibers with significant numbers of abnormal mitochondria could be highlighted with the modified trichrome stain; the term *ragged red fibers* was coined. By electron microscopy, the mitochondria in ragged red fibers are enlarged and often bizarrely shaped and have crystalline inclusions. Since that seminal observation, the understanding of these disorders of muscle and other tissues has expanded ([Chap. 82](#)).

Mitochondria play a key role in energy production. Oxidation of the major nutrients derived from carbohydrate, fat, and protein leads to the generation of reducing equivalents. The latter are transported through the respiratory chain in the process known as *oxidative phosphorylation*. The energy generated by the oxidation-reduction reactions of the respiratory chain is stored in an electrochemical gradient coupled to ATP synthesis.

A novel feature of mitochondria is their genetic composition. Each mitochondrion possesses a DNA genome that is distinct from that of the nuclear DNA. Human mitochondrial DNA (mtDNA) consists of a double-strand, circular molecule comprising 16,569 base pairs. It

codes for 22 transfer RNAs, 2 ribosomal RNAs, and 13 polypeptides of the respiratory chain enzymes. The genetics of mitochondrial diseases differ from the genetics of chromosomal disorders. The DNA of mitochondria is directly inherited from the cytoplasm of the gametes, mainly from the oocyte. The sperm contributes very little of its mitochondria to the offspring at the time of fertilization. Thus, mitochondrial genes are derived almost exclusively from the mother, accounting for maternal inheritance of some mitochondrial disorders.

Patients with mitochondrial myopathies have clinical manifestations that usually fall into three groups: chronic progressive external ophthalmoplegia (CPEO), skeletal muscle–CNS syndromes, and pure myopathy simulating muscular dystrophy or metabolic myopathy.

PROGRESSIVE EXTERNAL OPHTHALMOPLÉGIA SYNDROMES WITH RAGGED RED FIBERS

The single most common sign of a mitochondrial myopathy is CPEO, occurring in >50% of all mitochondrial myopathies. Varying degrees of ptosis and weakness of extraocular muscles are seen, usually in the absence of diplopia, a point of distinction from disorders with fluctuating eye weakness (e.g., myasthenia gravis).

KEARNS-SAYRE SYNDROME (KSS)

KSS is a widespread multiorgan system disorder with a defined triad of clinical findings: onset before age 20, CPEO, and pigmentary retinopathy, plus one or more of the following features: complete heart block, cerebrospinal fluid (CSF) protein >1 g/L (100 mg/dL), or cerebellar ataxia. Some patients with CPEO and ragged red fibers may not fulfill all of the criteria for KSS. The cardiac disease includes syncopal attacks and cardiac arrest related to the abnormalities in the cardiac conduction system: prolonged intraventricular conduction time, bundle branch block, and complete atrioventricular block. Death attributed to heart block occurs in ~20% of the patients. Varying degrees of progressive limb muscle weakness and easy fatigability affect activities of daily living. Endocrine abnormalities are common, including gonadal dysfunction in both sexes with delayed puberty, short stature, and infertility. Diabetes mellitus is a cardinal sign of mitochondrial disorders and is estimated to occur in 13% of KSS patients. Other less common endocrine disorders include thyroid disease, hyperaldosteronism, Addison's disease, and hypoparathyroidism. Both mental retardation and dementia are common accompaniments to this disorder. Serum CK levels are normal or slightly elevated. Serum lactate and pyruvate levels may be elevated. EMG is myopathic. Nerve conduction studies may be abnormal related to an associated neuropathy. Muscle biopsies reveal ragged red fibers, highlighted in oxidative enzyme stains, many showing defects in cytochrome oxidase. By electron microscopy, there are increased numbers of mitochondria that often appear enlarged and contain paracrystalline inclusions.

KSS is a sporadic disorder. The disease is caused by single mtDNA deletions presumed to arise spontaneously in the ovum or zygote. The most common deletion, occurring in about one-third of patients, removes 4977 bp of contiguous mtDNA. Monitoring for cardiac conduction defects is critical. Prophylactic pacemaker implantation is indicated when ECGs demonstrate a bifascicular block. In KSS, no benefit has been shown for supplementary therapies, including multivitamins or coenzyme Q10. Of all the proposed options, exercise might be the most applicable but must be approached cautiously because of defects in the cardiac conduction system.

PROGRESSIVE EXTERNAL OPHTHALMOPLÉGIA (PEO)

This condition is caused by nuclear DNA mutations affecting mtDNA copy number and integrity and is thus inherited in a Mendelian fashion. Onset is usually after puberty. Fatigue, exercise intolerance, and complaints of muscle weakness are typical. Some patients notice swallowing problems. The neurologic examination confirms the ptosis and ophthalmoplegia, usually asymmetric in distribution. A sensorineural hearing loss may be encountered. Mild facial, neck flexor, and proximal weakness are typical. Rarely, respiratory muscles may be progressively affected and may be the direct cause of death. Serum CK is normal or mildly elevated. The resting lactate level is normal or

462e-16 slightly elevated but may rise excessively after exercise. CSF protein is normal. The EMG is myopathic, and nerve conduction studies are usually normal. Ragged red fibers are prominently displayed in the muscle biopsy. Southern blots of muscle reveal a normal mtDNA band at 16.6 kb and several additional mtDNA deletion bands with genomes varying from 0.5 to 10 kb.

This autosomal dominant form of CPEO has been linked to loci on three chromosomes: 4q35, 10q24, and 15q22–26. In the chromosome 4q-related form of disease, mutations of the gene encoding the heart and skeletal muscle-specific isoform of the adenine nucleotide translocator 1 (*ANT1*) gene are found. This highly abundant mitochondrial protein forms a homodimeric inner mitochondrial channel through which adenosine diphosphate (ADP) enters and ATP leaves the mitochondrial matrix. In the chromosome 10q-related disorder, mutations of the gene *C10orf2* are found. Its gene product, *twinkle*, co-localizes with the mtDNA and is named for its punctate, starlike staining properties. The function of *twinkle* is presumed to be critical for lifetime maintenance of mitochondrial integrity. In the cases mapped to chromosome 15q, a mutation affects the gene encoding mtDNA polymerase (*POLG*), an enzyme important in mtDNA replication. Autosomal recessive PEO has also been described with mutations in the *POLG* gene. Point mutations have been identified within various mitochondrial tRNA (Leu, Ile, Asn, Trp) genes in families with maternal inheritance of PEO.

Exercise may improve function but will depend on the patient's ability to participate.

MITOCHONDRIAL DNA SKELETAL MUSCLE–CENTRAL NERVOUS SYSTEM SYNDROMES

Myoclonic Epilepsy with Ragged Red Fibers (MERRF) The onset of MERRF is variable, ranging from late childhood to middle adult life. Characteristic features include myoclonic epilepsy, cerebellar ataxia, and progressive muscle weakness. The seizure disorder is an integral part of the disease and may be the initial symptom. Cerebellar ataxia precedes or accompanies epilepsy. It is slowly progressive and generalized. The third major feature of the disease is muscle weakness in a limb-girdle distribution. Other more variable features include dementia, peripheral neuropathy, optic atrophy, hearing loss, and diabetes mellitus.

Serum CK levels are normal or slightly increased. The serum lactate may be elevated. EMG is myopathic, and in some patients nerve conduction studies show a neuropathy. The electroencephalogram is abnormal, corroborating clinical findings of epilepsy. Typical ragged red fibers are seen on muscle biopsy. MERRF is caused by maternally inherited point mutations of mitochondrial tRNA genes. The most common mutation found in 80% of MERRF patients is an A to G substitution at nucleotide 8344 of tRNA lysine (A8344G tRNA^{Lys}). Other tRNA^{Lys} mutations include base-pair substitutions T8356C and G8363A. Only supportive treatment is possible, with special attention to epilepsy.

Mitochondrial Myopathy, Encephalopathy, Lactic Acidosis, and Strokelike Episodes (MELAS)

MELAS is the most common mitochondrial encephalomyopathy. The term *strokelike* is appropriate because the cerebral lesions do not conform to a strictly vascular distribution. The onset in the majority of patients is before age 20. Seizures, usually partial motor or generalized, are common and may represent the first clearly recognizable sign of disease. The cerebral insults that resemble strokes cause hemiparesis, hemianopia, and cortical blindness. A presumptive stroke occurring before age 40 should place this mitochondrial encephalomyopathy high in the differential diagnosis. Associated conditions include hearing loss, diabetes mellitus, hypothalamic pituitary dysfunction causing growth hormone deficiency, hypothyroidism, and absence of secondary sexual characteristics. In its full expression, MELAS leads to dementia, a bedridden state, and a fatal outcome. Serum lactic acid is typically elevated. The CSF protein is also increased but is usually ≤ 1 g/L (100 mg/dL). Muscle biopsies show ragged red fibers. Neuroimaging demonstrates basal ganglia calcification in a high percentage of cases. Focal lesions that mimic infarction

are present predominantly in the occipital and parietal lobes. Strict vascular territories are not respected, and cerebral angiography fails to demonstrate lesions of the major cerebral blood vessels.

MELAS is caused by maternally inherited point mutations of mitochondrial tRNA genes. Most of the tRNA mutations are lethal, accounting for the paucity of multigeneration families with this syndrome. The A3243G point mutation in tRNA^{Leu(UUR)} is the most common, occurring in ~80% of MELAS cases. About 10% of MELAS patients have other mutations of the tRNA^{Leu(UUR)} gene, including 3252G, 3256T, 3271C, and 3291C. Other tRNA gene mutations have also been reported in MELAS, including G583A tRNA^{Phe}, G1642A tRNA^{Val}, G4332A tRNA^{Glu}, and T8316C tRNA^{Lys}. Mutations have also been reported in mtDNA polypeptide-coding genes. Two mutations were found in the ND5 subunit of complex I of the respiratory chain. A missense mutation has been reported at mtDNA position 9957 in the gene for subunit III of cytochrome C oxidase. No specific treatment is available. Supportive treatment is essential for the strokelike episodes, seizures, and endocrinopathies.

PURE MYOPATHY SYNDROMES

Muscle weakness and fatigue can be the predominant manifestations of mtDNA mutations. When the condition affects exclusively muscle (pure myopathy), the disorder becomes difficult to recognize. Occasionally, mitochondrial myopathies can present with recurrent myoglobinuria without fixed weakness and thus resemble a glycogen storage disorder or CPT deficiency.

Mitochondrial DNA Depletion Syndromes Mitochondrial DNA depletion syndrome (MDS) is a heterogeneous group of disorders that are inherited in an autosomal recessive fashion and can present in infancy or adults. MDS can be caused by mutations in genes (*TK2*, *DGUOK*, *RRM2B*, *TYMP*, *SUCLA1*, and *SUCLA2*) that lead to depletion of pools of mitochondrial deoxyribonucleotide (dNTP) pools necessary for mtDNA replication. The other major cause of MDS is a set of mutations in genes essential for mtDNA replication (e.g., *POLG1* and *C10orf2*). The clinical phenotypes associated with MDS vary. Patients may develop a severe encephalopathy (e.g., Leigh's syndrome), PEO, an isolated myopathy, myo-neuro-gastrointestinal-encephalopathy (MNGIE), and a sensory neuropathy with ataxia.

DISORDERS OF MUSCLE MEMBRANE EXCITABILITY

Muscle membrane excitability is affected in a group of disorders referred to as *channelopathies*. The heart may also be involved, resulting in life-threatening complications (Table 462e-10).

CALCIUM CHANNEL DISORDERS OF MUSCLE

Hypokalemic Periodic Paralysis (HypoKPP) Onset occurs at adolescence. Men are more often affected because of decreased penetrance in women. Episodic weakness with onset after age 25 is almost never due to periodic paralyzes, with the exception of thyrotoxic periodic paralysis (see below). Attacks are often provoked by meals high in carbohydrates or sodium and may accompany rest following prolonged exercise. Weakness usually affects proximal limb muscles more than distal. Ocular and bulbar muscles are less likely to be affected. Respiratory muscles are usually spared, but when they are involved, the condition may prove fatal. Weakness may take as long as 24 h to resolve. Life-threatening cardiac arrhythmias related to hypokalemia may occur during attacks. As a late complication, patients commonly develop severe, disabling proximal lower extremity weakness.

Attacks of thyrotoxic periodic paralysis resemble those of primary HypoKPP. Despite a higher incidence of thyrotoxicosis in women, men, particularly those of Asian descent, are more likely to manifest this complication. Attacks abate with treatment of the underlying thyroid condition.

A low serum potassium level during an attack, excluding secondary causes, establishes the diagnosis. Interattack muscle biopsies show the presence of single or multiple centrally placed vacuoles or tubular aggregates. Provocative tests with glucose and insulin to establish a diagnosis are usually not necessary and are potentially hazardous.

TABLE 462e-10 CLINICAL FEATURES OF PERIODIC PARALYSIS AND NONDYSTROPHIC MYOTONIAS

Feature	Calcium Channel		Sodium Channel		Potassium Channel
	Hypokalemic PP	Hyperkalemic PP	Paramyotonia Congenita	Andersen-Tawil Syndrome ^a	
Mode of inheritance	AD	AD	AD	AD	AD
Age of onset	Adolescence	Early childhood	Early childhood	Early childhood	Early childhood
Myotonia ^b	No	Yes	Yes	No	No
Episodic weakness	Yes	Yes	Yes	Yes	Yes
Frequency of attacks of weakness	Daily to yearly	May be 2–3/d	With cold, usually rare	Daily to yearly	Daily to yearly
Duration of attacks of weakness	2–12 h	From 1–2 h to >1 d	2–24 h	2–24 h	2–24 h
Serum K ⁺ level during attacks of weakness	Decreased	Increased or normal	Usually normal	Variable	Variable
Effect of K ⁺ loading	No change	Increased myotonia, then weakness	Increased myotonia	No change	No change
Effect of muscle cooling	No change	Increased myotonia	Increased myotonia, then weakness	No change	No change
Fixed weakness	Yes	Yes	Yes	Yes	Yes

^aDysmorphic features and cardiac arrhythmias are distinguishing features (see text). ^bMay be paradoxical in paramyotonia congenita.

Abbreviations: AD, autosomal dominant; PP, periodic paralysis.

In the midst of an attack of weakness, motor conduction studies may demonstrate reduced amplitudes, whereas EMG may show electrical silence in severely weak muscles. In between attacks, the EMG and routine nerve conduction studies are normal. However, a long exercise test may demonstrate a decrementing amplitude, and myopathic MUAPs may be seen on EMG in patients with fixed weakness.

HypoKPP is caused by mutations in either of two genes. HypoKPP type 1, the most common form, is inherited as an autosomal dominant disorder with incomplete penetrance. These patients have mutations in the voltage-sensitive, skeletal muscle calcium channel gene, *CALCLIA3* (Fig. 462e-8). Approximately 10% of cases are HypoKPP type 2, arising from mutations in the voltage-sensitive sodium channel gene (*SCN4A*). In either instance, the mutations lead to an abnormal gating pore current that predisposes the muscle cell to depolarize when potassium levels are low. It is also now recognized that some cases of thyrotoxic HypoKPP are caused by genetic variants in a potassium channel (Kir 2.6), whose expression is regulated by thyroid hormone.

The chloride channel is envisioned to have 10 membrane-spanning domains. The positions of mutations causing dominantly and recessively inherited myotonia congenita are indicated, along with mutations that cause this disease in mice and goats.

TREATMENT HYPOKALEMIC PERIODIC PARALYSIS

The acute paralysis improves after the administration of potassium. Muscle strength and ECG should be monitored. Oral KCl (0.2–0.4 mmol/kg) should be given every 30 min. Only rarely is IV therapy necessary (e.g., when swallowing problems or vomiting is present). Administration of potassium in a glucose solution should be avoided because it may further reduce serum potassium levels. Mannitol is the preferred vehicle for administration of IV potassium. The long-term goal of therapy is to avoid attacks. This may reduce late-onset, fixed weakness. Patients should be made aware of the importance of a low-carbohydrate, low-sodium diet and consequences of intense exercise. Prophylactic administration of acetazolamide (125–1000 mg/d in divided doses) reduces or may abolish attacks in HypoKPP type 1. Paradoxically the potassium is lowered, but this is offset by the beneficial effect of metabolic acidosis. If attacks persist on acetazolamide, oral KCl should be added. Some patients require treatment with triamterene (25–100 mg/d) or spironolactone (25–100 mg/d). However, in patients with HypoKPP type 2, attacks of weakness can be exacerbated with acetazolamide.

SODIUM CHANNEL DISORDERS OF MUSCLE

Hyperkalemic Periodic Paralysis (HyperKPP) The term *hyperkalemic* is misleading because patients are often normokalemic during attacks.

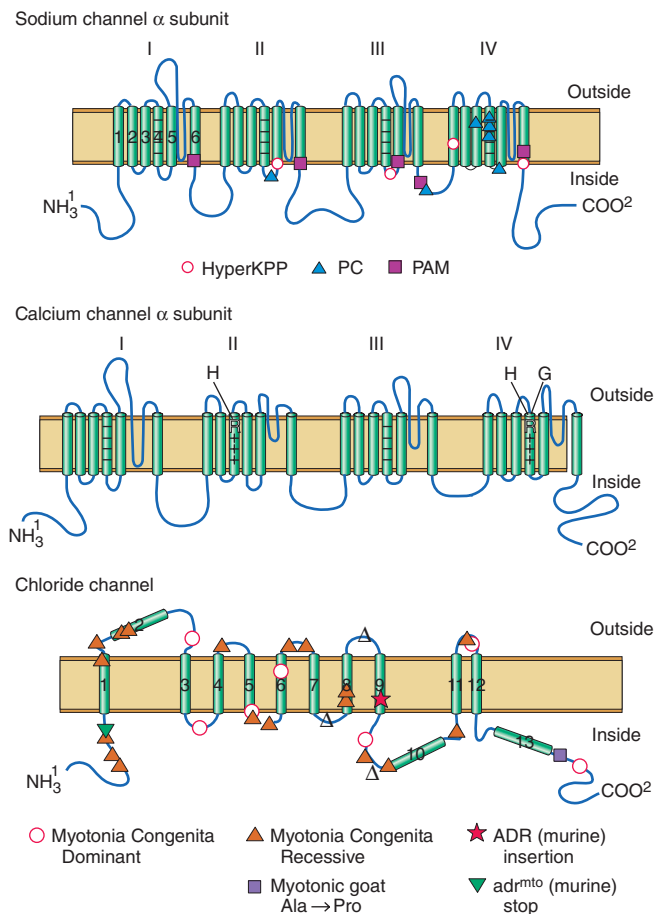


FIGURE 462e-8 The sodium and calcium channels are depicted here as containing four homologous domains, each with six membrane-spanning segments. The fourth segment of each domain bears positive charges and acts as the “voltage sensor” for the channel. The association of the four domains is thought to form a pore through which ions pass. Sodium channel mutations are shown along with the phenotype that they confer. HyperKPP, hyperkalemic periodic paralysis; PC, paramyotonia congenita; PAM, potassium-aggravated myotonia. See text for details.

462e-18 The fact that attacks are precipitated by potassium administration best defines the disease. The onset is in the first decade; males and females are affected equally. Attacks are brief and mild, usually lasting 30 min to 4 ho. Weakness affects proximal muscles, sparing bulbar muscles. Attacks are precipitated by rest following exercise and fasting. In a variant of this disorder, the predominant symptom is myotonia without weakness (*potassium-aggravated myotonia*). The symptoms are aggravated by cold, and myotonia makes the muscles stiff and painful. This disorder can be confused with paramyotonia congenita, myotonia congenita, and proximal myotonic myopathy (DM2).

Potassium may be slightly elevated but may also be normal during an attack. As in HypoKPP, nerve conduction studies in HyperKPP muscle may demonstrate reduced motor amplitudes and the EMG may be silent in very weak muscles. In between attacks of weakness, the conduction studies are normal. The EMG will often demonstrate myotonic discharges during and between attacks.

The muscle biopsy shows vacuoles that are smaller, less numerous, and more peripheral compared to the hypokalemic form or tubular aggregates. Provocative tests by administration of potassium can induce weakness but are usually not necessary to establish the diagnosis. HyperKPP and potassium-aggravated myotonia are inherited as autosomal dominant disorders. Mutations of the voltage-gated sodium channel *SCN4A* gene (Fig. 462e-8) cause these conditions. For patients with frequent attacks, acetazolamide (125–1000 mg/d) is helpful. We have found mexiletine to be helpful in patients with significant myotonia.

Paramyotonia Congenita In paramyotonia congenita (PC), the attacks of weakness are cold-induced or occur spontaneously and are mild. Myotonia is a prominent feature but worsens with muscle activity (paradoxical myotonia). This is in contrast to classic myotonia in which exercise alleviates the condition. Attacks of weakness are seldom severe enough to require emergency room treatment. Over time patients develop interattack weakness as they do in other forms of periodic paralysis. PC is usually associated with normokalemia or hyperkalemia.

Serum CK is usually mildly elevated. Routine sensory and motor nerve conduction studies are normal. Short exercise test may be abnormal however, and cooling of the muscle often dramatically reduces the amplitude of the compound muscle action potentials. EMG reveals diffuse myotonic potentials in PC. Upon local cooling of the muscle, the myotonic discharges disappear as the patient becomes unable to activate MUAPs.

PC is inherited as an autosomal dominant condition; voltage-gated sodium channel mutations (Fig. 462e-8) are responsible, and thus this disorder is allelic with HyperKPP and potassium-aggravated myotonia. Patients with PC seldom seek treatment during attacks. Oral administration of glucose or other carbohydrates hastens recovery. Because interattack weakness may develop after repeated episodes, prophylactic treatment is usually indicated. Thiazide diuretics (e.g., chlorothiazide, 250–1000 mg/d) and mexiletine (slowly increase dose from 450 mg/d) are reported to be helpful. Patients should be advised to increase carbohydrates in their diet.

POTASSIUM CHANNEL DISORDERS

Andersen-Tawil Syndrome This rare disease is characterized by episodic weakness, cardiac arrhythmias, and dysmorphic features (short stature, scoliosis, clinodactyly, hypertelorism, small or prominent low-set ears, micrognathia, and broad forehead). The cardiac arrhythmias are potentially serious and life threatening. They include long QT, ventricular ectopy, bidirectional ventricular arrhythmias, and tachycardia. For many years, the classification of this disorder was uncertain because episodes of weakness are associated with elevated, normal, or reduced levels of potassium during an attack. In addition, the potassium levels differ among kindreds but are consistent within a family. Inheritance is autosomal dominant, with incomplete penetrance and variable expressivity. The disease is caused by mutations of the inwardly rectifying potassium channel (*Kir 2.1*) gene that heighten muscle cell excitability. The treatment is similar to that for other forms of periodic paralysis and must include cardiac monitoring. The episodes of weakness may differ between patients because of potassium

variability. Acetazolamide may decrease the attack frequency and severity.

CHLORIDE CHANNEL DISORDERS

Two forms of this disorder, autosomal dominant (*Thomsen's disease*) and autosomal recessive (*Becker disease*), are related to the same gene abnormality. Symptoms are noted in infancy and early childhood. The severity lessens in the third to fourth decade. Myotonia is worsened by cold and improved by activity. The gait may appear slow and labored at first but improves with walking. In Thomsen's disease, muscle strength is normal, but in Becker disease, which is usually more severe, there may be muscle weakness. Muscle hypertrophy is usually present. Myotonic discharges are prominently displayed by EMG recordings.

Serum CK is normal or mildly elevated. The muscle biopsy shows hypertrophied fibers. The disease is inherited as dominant or recessive and is caused by mutations of the chloride channel gene (Fig. 462e-8) that increase muscle cell excitability. Many patients will not require treatment and learn that the symptoms improve with activity. Medications that can be used to decrease myotonia include quinine, phenytoin, and mexiletine.

ENDOCRINE AND METABOLIC MYOPATHIES

Many endocrine disorders cause weakness. Muscle fatigue is more common than true weakness. The cause of weakness in these disorders is not well defined. It is not even clear that weakness results from disease of muscle as opposed to another part of the motor unit, since the serum CK level is often normal (except in hypothyroidism) and the muscle histology is characterized by atrophy rather than destruction of muscle fibers. Nearly all endocrine myopathies respond to treatment.

THYROID DISORDERS

(See also Chap. 405) Abnormalities of thyroid function can cause a wide array of muscle disorders. These conditions relate to the important role of thyroid hormones in regulating the metabolism of carbohydrates and lipids as well as the rate of protein synthesis and enzyme production. Thyroid hormones also stimulate calorogenesis in muscle, increase muscle demand for vitamins, and enhance muscle sensitivity to circulating catecholamines.

Hypothyroidism Patients with hypothyroidism have frequent muscle complaints, and proximal muscle weakness occurs in about one-third of them. Muscle cramps, pain, and stiffness are common. Some patients have enlarged muscles. Features of slow muscle contraction and relaxation occur in 25% of patients; the relaxation phase of muscle stretch reflexes is characteristically prolonged and best observed at the ankle or biceps brachii reflexes. The serum CK level is often elevated (up to 10 times normal), even when there is minimal clinical evidence of muscle disease. EMG is typically normal. The cause of muscle enlargement has not been determined, and muscle biopsy shows no distinctive morphologic abnormalities.

Hyperthyroidism Patients who are thyrotoxic commonly have proximal muscle weakness and atrophy on examination, but they rarely complain of myopathic symptoms. Activity of deep tendon reflexes may be enhanced. Bulbar, respiratory, and even esophageal muscles may occasionally be affected, causing dysphagia, dysphonia, and aspiration. When bulbar involvement occurs, it is usually accompanied by chronic proximal limb weakness, but occasionally it presents in the absence of generalized thyrotoxic myopathy. Fasciculations may be apparent and, when coupled with increased muscle stretch reflexes, may lead to an erroneous diagnosis of amyotrophic lateral sclerosis. A form hypokalemic periodic paralysis can occur in patients who are thyrotoxic. Recently, mutations in the *KCNJ18* gene that encodes for the inwardly rectifying potassium channel, Kir 2.6, have been discovered in up to a third of cases. Other neuromuscular disorders that occur in association with hyperthyroidism include myasthenia gravis (Chap. 461) and a progressive ocular myopathy associated with proptosis (*Graves' ophthalmopathy*). Serum CK levels are not elevated in thyrotoxic myopathy, the EMG is normal, and muscle histology usually shows only atrophy of muscle fibers.

PARATHYROID DISORDERS

(See also Chap. 424)

Hyperparathyroidism Muscle weakness is an integral part of primary and secondary hyperparathyroidism. Proximal muscle weakness, muscle wasting, and brisk muscle stretch reflexes are the main features of this endocrinopathy. Some patients develop neck extensor weakness (part of the dropped head syndrome). Serum CK levels are usually normal or slightly elevated. Serum parathyroid hormone levels are elevated. Serum calcium and phosphorus levels show no correlation with the clinical neuromuscular manifestations. Muscle biopsies show only varying degrees of atrophy without muscle fiber degeneration.

Hypoparathyroidism An overt myopathy due to hypocalcemia rarely occurs. Neuromuscular symptoms are usually related to localized or generalized tetany. Serum CK levels may be increased secondary to muscle damage from sustained tetany. Hyporeflexia or areflexia is usually present and contrasts with the hyperreflexia in hyperparathyroidism.

ADRENAL DISORDERS

(See also Chap. 406) Conditions associated with glucocorticoid excess cause a myopathy; in fact, steroid myopathy is the most commonly diagnosed endocrine muscle disease. Glucocorticoid excess, either endogenous or exogenous (see “Drug-Induced Myopathies,” below), produces various degrees of proximal limb weakness. Muscle wasting may be striking. A cushingoid appearance usually accompanies clinical signs of myopathy. Histologic sections demonstrate muscle fiber atrophy, preferentially affecting type 2b fibers, rather than degeneration or necrosis of muscle fibers. Adrenal insufficiency commonly causes muscle fatigue. The degree of weakness may be difficult to assess but is typically mild. In primary hyperaldosteronism (*Conn’s syndrome*), neuromuscular complications are due to potassium depletion. The clinical picture is one of persistent muscle weakness. Long-standing hyperaldosteronism may lead to proximal limb weakness and wasting. Serum CK levels may be elevated, and a muscle biopsy may demonstrate degenerating fibers, some with vacuoles. These changes relate to hypokalemia and are not a direct effect of aldosterone on skeletal muscle.

PITUITARY DISORDERS

(See also Chap. 403) Patients with acromegaly usually have mild proximal weakness without muscle atrophy. Muscles often appear enlarged but exhibit decreased force generation. The duration of acromegaly, rather than the serum growth hormone levels, correlates with the degree of myopathy.

DIABETES MELLITUS

(See also Chap. 417) Neuromuscular complications of diabetes mellitus are most often related to neuropathy, with cranial and peripheral nerve palsies or distal sensorimotor polyneuropathy. *Diabetic amyotrophy* is a clumsy term because the condition represents a neuropathy affecting the proximal major nerve trunks and lumbosacral plexus. More appropriate terms for this disorder include *diabetic proximal neuropathy* and *lumbosacral radiculoplexus neuropathy*.

The only notable myopathy of diabetes mellitus is ischemic infarction of leg muscles, usually involving one of the thigh muscles but on occasion affecting the distal leg. This condition occurs in patients with poorly controlled diabetes and presents with abrupt onset of pain, tenderness, and edema of one thigh. The area of muscle infarction is hard and indurated. The muscles most often affected include the vastus lateralis, thigh adductors, and biceps femoris. Computed tomography (CT) or MRI can demonstrate focal abnormalities in the affected muscle. Diagnosis by imaging is preferable to muscle biopsy, if possible, as hemorrhage into the biopsy site can occur.

VITAMIN DEFICIENCY

Vitamin D deficiency (Chap. 96e) due to decreased intake, decreased absorption, or impaired vitamin D metabolism (as occurs in renal disease) may lead to chronic muscle weakness. Pain reflects the underlying bone disease (*osteomalacia*). Vitamin E deficiency may result from malabsorption. Clinical manifestations include ataxic neuropathy due to loss of proprioception and myopathy with proximal weakness.

Progressive external ophthalmoplegia is a distinctive finding. It has not been established that deficiency of other vitamins causes a myopathy.

MYOPATHIES OF SYSTEMIC ILLNESS

Systemic illnesses such as chronic respiratory, cardiac, or hepatic failure are frequently associated with severe muscle wasting and complaints of weakness. Fatigue is usually a more significant problem than weakness, which is typically mild.

Myopathy may be a manifestation of chronic renal failure (CRF), independent of the better known uremic polyneuropathy. Abnormalities of calcium and phosphorus homeostasis and bone metabolism in chronic renal failure result from a reduction in 1,25-dihydroxyvitamin D, leading to decreased intestinal absorption of calcium. Hypocalcemia, further accentuated by hyperphosphatemia due to decreased renal phosphate clearance, leads to secondary hyperparathyroidism. Renal osteodystrophy results from the compensatory hyperparathyroidism, which leads to osteomalacia from reduced calcium availability and to osteitis fibrosa from the parathyroid hormone excess. The clinical picture of the myopathy of CRF is identical to that of primary hyperparathyroidism and osteomalacia. There is proximal limb weakness with bone pain.

Gangrenous calcification represents a separate, rare, and sometimes fatal complication of CRF. In this condition, widespread arterial calcification occurs and results in ischemia. Extensive skin necrosis may occur, along with painful myopathy and even myoglobinuria.

DRUG-INDUCED MYOPATHIES

Drug-induced myopathies are relatively uncommon in clinical practice with the exception of those caused by the cholesterol-lowering agents and glucocorticoids. Others impact practice to a lesser degree but are important to consider in specific situations. Table 462e-11 provides a comprehensive list of drug-induced myopathies with their distinguishing features.

MYOPATHY FROM LIPID-LOWERING AGENTS

All classes of lipid-lowering agents have been implicated in muscle toxicity, including fibrates (clofibrate, gemfibrozil), HMG-CoA reductase inhibitors (referred to as *statins*), niacin (nicotinic acid), and ezetimibe. Myalgia, malaise, and muscle tenderness are the most common manifestations. Muscle pain may be related to exercise. Patients may exhibit proximal weakness. Varying degrees of muscle necrosis are seen, and in severe reactions rhabdomyolysis and myoglobinuria occur. Concomitant use of statins with fibrates and cyclosporine is more likely to cause adverse reactions than use of one agent alone. Elevated serum CK is an important indication of toxicity. Muscle weakness is accompanied by a myopathic EMG, and muscle necrosis is observed by muscle biopsy. Severe myalgias, muscle weakness, significant elevations in serum CK (>three times baseline), and myoglobinuria are indications for stopping the drug. Patients usually improve with drug cessation, although this may take several weeks. Rare cases continue to progress after the offending agent is discontinued. It is possible that in such cases the statin may have triggered an immune-mediated necrotizing myopathy, as these individuals require aggressive immunotherapy (e.g., prednisone and sometimes other agents) to improve and often relapse when these therapies are discontinued. Interestingly, antibodies directed against the 100-kDa HMG-CoA reductase receptor on muscle fibers have been identified in many of these cases.

GLUCOCORTICOID-RELATED MYOPATHIES

Glucocorticoid myopathy occurs with chronic treatment or as “acute quadriplegic” myopathy secondary to high-dose IV glucocorticoid use. Chronic administration produces proximal weakness accompanied by cushingoid manifestations, which can be quite debilitating; the chronic use of prednisone at a daily dose of ≥ 30 mg/d is most often associated with toxicity. Patients taking fluorinated glucocorticoids (triamcinolone, betamethasone, dexamethasone) appear to be at especially high risk for myopathy. In chronic steroid myopathy, the serum CK is

TABLE 462e-11 DRUG-INDUCED MYOPATHIES

Drugs	Major Toxic Reaction
Lipid-lowering agents Fibric acid derivatives HMG-CoA reductase inhibitors Niacin (nicotinic acid)	Drugs belonging to all three of the major classes of lipid-lowering agents can produce a spectrum of toxicity: asymptomatic serum creatine kinase elevation, myalgias, exercise-induced pain, rhabdomyolysis, and myoglobinuria.
Glucocorticoids	Acute, high-dose glucocorticoid treatment can cause acute quadriplegic myopathy. These high doses of steroids are often combined with nondepolarizing neuromuscular blocking agents but the weakness can occur without their use. Chronic steroid administration produces predominantly proximal weakness.
Nondepolarizing neuromuscular blocking agents	Acute quadriplegic myopathy can occur with or without concomitant glucocorticoids.
Zidovudine	Mitochondrial myopathy with ragged red fibers
Drugs of abuse Alcohol Amphetamines Cocaine Heroin Phencyclidine Meperidine	All drugs in this group can lead to widespread muscle breakdown, rhabdomyolysis, and myoglobinuria. Local injections cause muscle necrosis, skin induration, and limb contractures.
Autoimmune toxic myopathy D-Penicillamine	Use of this drug may cause polymyositis and myasthenia gravis.
Amphophilic cationic drugs Amiodarone Chloroquine Hydroxychloroquine	All amphophilic drugs have the potential to produce painless, proximal weakness associated with autophagic vacuoles in the muscle biopsy.
Antimicrotubular drugs Colchicine	This drug produces painless, proximal weakness especially in the setting of renal failure. Muscle biopsy shows autophagic vacuoles.

usually normal. Serum potassium may be low. The muscle biopsy in chronic cases shows preferential type 2 muscle fiber atrophy; this is not reflected in the EMG, which is usually normal.

Patients receiving high-dose IV glucocorticoids for status asthmaticus, chronic obstructive pulmonary disease, organ transplantation, or other indications may develop severe generalized weakness (critical illness myopathy). This myopathy, also known as acute quadriplegic myopathy, can also occur in the setting of sepsis. Involvement of the diaphragm and intercostal muscles causes respiratory failure and requires ventilatory support. In these settings, the use of glucocorticoids in combination with nondepolarizing neuromuscular blocking agents potentiates this complication. In critical illness myopathy, the muscle biopsy is abnormal, showing a distinctive loss of thick filaments (myosin) by electron microscopy. By light microscopy, there is focal loss of ATPase staining in central or paracentral areas of the muscle fiber. Calpain stains show diffusely reactive atrophic fibers. Withdrawal of glucocorticoids will improve the chronic myopathy. In acute quadriplegic myopathy, recovery is slow. Patients require supportive care and rehabilitation.

DRUG-INDUCED MITOCHONDRIAL MYOPATHY

Zidovudine, used in the treatment of HIV infection, is a thymidine analog that inhibits viral replication by interrupting reverse transcriptase.

Myopathy is a well-established complication of this agent. Patients present with myalgias, muscle weakness, and atrophy affecting the thigh and calf muscles. The complication occurs in about 17% of patients treated with doses of 1200 mg/d for 6 months. The introduction of protease inhibitors for treatment of HIV infection has led to lower doses of zidovudine therapy and a decreased incidence of myopathy. Serum CK is elevated and EMG is myopathic. Muscle biopsy shows ragged red fibers with minimal inflammation; the lack of inflammation serves to distinguish zidovudine toxicity from HIV-related myopathy. If the myopathy is thought to be drug related, the medication should be stopped or the dosage reduced.

DRUGS OF ABUSE AND RELATED MYOPATHIES

Myotoxicity is a potential consequence of addiction to alcohol and illicit drugs. Ethanol is one of the most commonly abused substances with potential to damage muscle. Other potential toxins include cocaine, heroin, and amphetamines. The most deleterious reactions occur from overdosing leading to coma and seizures, causing rhabdomyolysis, myoglobinuria, and renal failure. Direct toxicity can occur from cocaine, heroin, and amphetamines causing muscle breakdown and varying degrees of weakness. The effects of alcohol are more controversial. Direct muscle damage is less certain, since toxicity usually occurs in the setting of poor nutrition and possible contributing factors such as hypokalemia and hypophosphatemia. Alcoholics are also prone to neuropathy (Chap. 467).

Focal myopathies from self-administration of meperidine, heroin, and pentazocine can cause pain, swelling, muscle necrosis, and hemorrhage. The cause is multifactorial; needle trauma, direct toxicity of the drug or vehicle, and infection may all play a role. When severe, there may be overlying skin induration and contractures with replacement of muscle by connective tissue. Elevated serum CK and myopathic EMG are characteristic of these reactions. The muscle biopsy shows widespread or focal areas of necrosis. In conditions leading to rhabdomyolysis, patients need adequate hydration to reduce serum myoglobin and protect renal function. In all of these conditions, counseling is essential to limit drug abuse.

DRUG-INDUCED AUTOIMMUNE MYOPATHIES

As mentioned previously, an autoimmune necrotizing myopathy associated with autoantibodies directed against HMG-CoA rarely occurs in the setting of statin use. An inflammatory myopathy also may occur with D-penicillamine, sometimes used in the treatment of Wilson's disease scleroderma, rheumatoid arthritis, and primary biliary cirrhosis. The incidence of this inflammatory muscle disease is about 1%. Myasthenia gravis is also induced by D-penicillamine, with a higher incidence estimated at 7%. These disorders resolve with drug withdrawal, although immunosuppressive therapy may be warranted in severe cases.

Scattered reports of other drugs causing an inflammatory myopathy are rare and include a heterogeneous group of agents: cimetidine, phenytoin, procainamide, and propylthiouracil. In most cases, a cause-and-effect relationship is uncertain. A complication of interest was related to L-tryptophan. In 1989 an epidemic of eosinophilia-myalgia syndrome (EMS) in the United States was caused by a contaminant in the product from one manufacturer. The product was withdrawn, and incidence of EMS diminished abruptly following this action.

OTHER DRUG-INDUCED MYOPATHIES

Certain drugs produce painless, largely proximal, muscle weakness. These drugs include the amphophilic cationic drugs (amiodarone, chloroquine, hydroxychloroquine) and antimicrotubular drugs (colchicine) (Table 462e-11). Muscle biopsy can be useful in the identification of toxicity because autophagic vacuoles are prominent pathologic features of these toxins.

463e Special Issues in Inpatient Neurologic Consultation

S. Andrew Josephson, Martin A. Samuels

Inpatient neurologic consultations usually involve questions regarding specific disease processes or prognostication after various cerebral injuries. Common reasons for neurologic consultation include stroke (Chap. 446), seizures (Chap. 445), altered mental status (Chap. 34), headache (Chap. 21), and management of coma and other neurocritical care conditions (Chaps. 328 and 330). This chapter focuses on additional common reasons for consultation that are not addressed elsewhere in the text.

CONSULTATIONS REGARDING CENTRAL NERVOUS SYSTEM DYSFUNCTION

HYPERPERFUSION STATES LEADING TO POSTERIOR REVERSIBLE ENCEPHALOPATHY SYNDROME

A group of neurologic disorders shares the common feature of hyperperfusion, probably related to endothelial dysfunction, playing a key role in pathogenesis. These seemingly diverse syndromes include hypertensive encephalopathy, eclampsia, postcarotid endarterectomy syndrome, and toxicity from calcineurin-inhibitor and other medications. Modern imaging techniques and experimental models suggest that vasogenic edema is typically the primary process leading to neurologic dysfunction; therefore, prompt recognition and management of this condition should allow for clinical recovery as long as superimposed hemorrhage or infarction has not occurred.

The brain's autoregulatory capability successfully maintains a fairly stable cerebral blood flow in adults despite alterations in systemic mean arterial pressure (MAP) ranging from 50 to 150 mmHg (Chap. 330). In patients with chronic hypertension, this cerebral autoregulation curve is shifted, resulting in autoregulation working over a much higher range of pressures (e.g., 70–175 mmHg). In these hypertensive patients, cerebral blood flow is kept steady at higher MAP, but a rapid lowering of pressure can lead to ischemia on the lower end of the autoregulatory curve, even at values typically thought of as normotensive. This autoregulatory phenomenon is achieved through both myogenic and neurogenic influences causing small arterioles to contract and dilate. When the systemic blood pressure exceeds the limits of this mechanism, breakthrough of autoregulation occurs, resulting in hyperperfusion via increased cerebral blood flow, capillary leakage into the interstitium, and resulting edema. The predilection of all of the hyperperfusion disorders to affect the posterior rather than anterior portions of the brain may be due to a lower threshold for autoregulatory breakthrough in the posterior circulation or a vasculopathy that is more common in these blood vessels.

Although elevated or relatively elevated blood pressure is common in many of these disorders, some hyperperfusion states such as calcineurin-inhibitor toxicity occur with no apparent pressure rise. In these cases, vasogenic edema is likely due primarily to dysfunction of the capillary endothelium itself, leading to breakdown of the blood-brain barrier. It is useful to separate disorders of hyperperfusion into those caused primarily by increased pressure and those due mostly to endothelial dysfunction from a toxic or autoimmune etiology (Table 463e-1). In reality, both of these pathophysiologic processes likely play some role in each of these disorders.

The clinical presentation of all of the hyperperfusion syndromes is similar with prominent headaches, seizures, or focal neurologic deficits. Headaches have no specific characteristics, range from mild to severe, and may be accompanied by alterations in consciousness ranging from confusion to coma. Seizures may be present, and these can be of multiple types depending on the severity and location of the edema. Nonconvulsive seizures have been described in hyperperfusion states; therefore, a low threshold for obtaining an electroencephalogram (EEG) in these patients should be maintained. The typical focal deficit

TABLE 463e-1 SOME COMMON ETIOLOGIES OF HYPERPERFUSION SYNDROME

Disorders in which increased capillary pressure dominates the pathophysiology
Hypertensive encephalopathy, including secondary causes such as renovascular hypertension, pheochromocytoma, cocaine use, etc.
Postcarotid endarterectomy syndrome
Preeclampsia/eclampsia
High-altitude cerebral edema
Disorders in which endothelial dysfunction dominates the pathophysiology
Calcineurin-inhibitor toxicity
Chemotherapeutic agent toxicity (e.g., cytarabine, azathioprine, 5-fluorouracil, cisplatin, methotrexate, tumor necrosis factor α antagonists)
Glucocorticoids
Erythropoietin
HELLP syndrome (hemolysis, elevated liver enzyme levels, low platelet count)
Thrombotic thrombocytopenic purpura (TTP)
Hemolytic-uremic syndrome (HUS)
Systemic lupus erythematosus (SLE)
Granulomatosis with polyangiitis (Wegener's)

in hyperperfusion states is cortical visual loss, given the tendency of the process to involve the occipital lobes. However, any focal deficit can occur depending on the area affected, as evidenced by patients who, after carotid endarterectomy, exhibit neurologic dysfunction referable to the ipsilateral newly reperfused hemisphere. In conditions where increased cerebral blood flow plays a role, examination of the inpatient vital signs record will usually reveal a systemic blood pressure that is increased above the patient's baseline. It appears as if the rapidity of rise, rather than the absolute value of pressure, is the most important risk factor.

The diagnosis in all of these conditions is clinical. The symptoms of these disorders are common and nonspecific, so a long differential diagnosis should be entertained, including consideration of other causes of confusion, focal neurologic deficits, headache, and seizures. Magnetic resonance imaging (MRI) has improved the ability of clinicians to diagnose hyperperfusion syndromes, although cases have been reported with normal imaging. Patients classically exhibit the high T2 signal of edema primarily in the posterior occipital lobes, not respecting any single vascular territory (Fig. 463e-1). Diffusion-weighted images are typically normal, emphasizing the vasogenic rather than cytotoxic nature of this edema. Imaging with computed tomography (CT) is less sensitive but may show a pattern of patchy hypodensity in the involved territory. Previously this classic radiographic appearance had been termed *reversible posterior leukoencephalopathy* (RPLE). However, this term has fallen out of favor because none of its elements are completely accurate. The radiographic and clinical changes are not always reversible; the territory involved is not uniquely posterior; and gray matter may be affected as well, rather than purely white matter as the term "leukoencephalopathy" intimates. The now more commonly used radiologic term *posterior reversible encephalopathy syndrome* (PRES) suffers from many of these same limitations. Vessel imaging may demonstrate narrowing of the cerebral vasculature, especially in the posterior circulation; whether this noninflammatory vasculopathy is a primary cause of the edema or occurs as a secondary phenomenon remains unclear. Other ancillary studies such as cerebrospinal fluid (CSF) analysis often yield nonspecific results. It should be noted that many of the substances that have been implicated, such as cyclosporine, can cause this syndrome even at low doses or after years of treatment. Therefore, normal serum levels of these medications do not exclude them as inciting agents.

In cases of hyperperfusion syndromes, treatment should commence urgently once the diagnosis is considered. Hypertension plays a key role commonly, and judicious lowering of the blood pressure with IV agents such as labetalol or nicardipine is advised along with continuous cardiac and blood pressure monitoring, often through an arterial line. It is reasonable to lower MAP by ~20% initially, as further

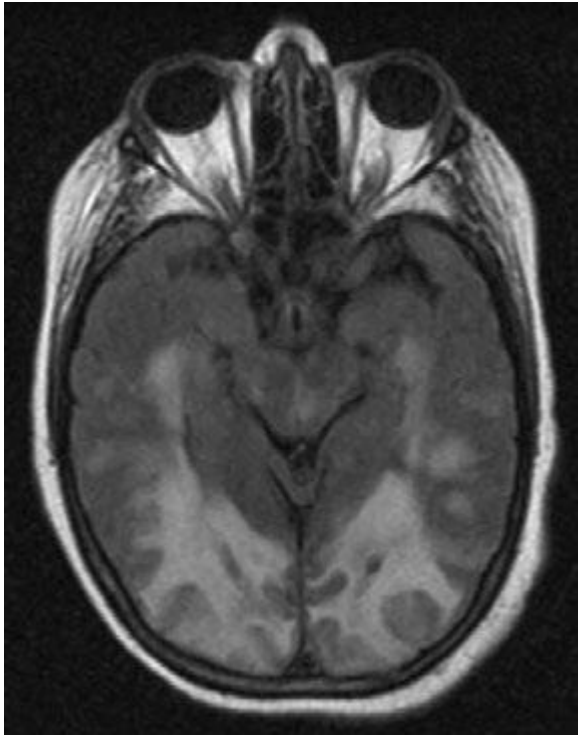


FIGURE 463e-1 Axial fluid-attenuated inversion recovery (FLAIR) magnetic resonance imaging (MRI) of the brain in a patient taking cyclosporine after liver transplantation, who presented with seizures, headache, and cortical blindness. Increased signal is seen bilaterally in the occipital lobes predominantly involving the white matter, consistent with a hyperperfusion state secondary to calcineurin-inhibitor exposure.

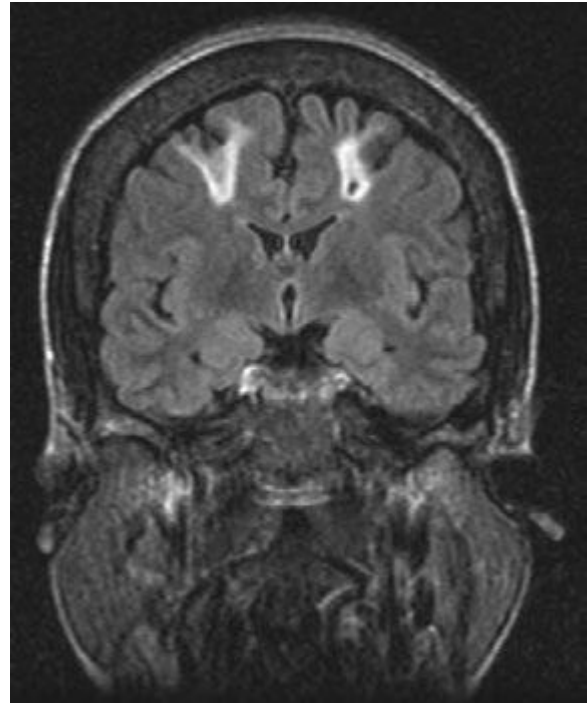


FIGURE 463e-2 Coronal fluid-attenuated inversion recovery (FLAIR) magnetic resonance imaging (MRI) of the brain in a patient presenting with altered mental status after an episode of hypotension during coronary artery bypass grafting (CABG). Increased signal is seen in the border zones bilaterally between the middle cerebral artery and anterior cerebral artery territories. Diffusion-weighted MRI sequences demonstrated restricted diffusion in these same locations, suggesting acute infarction.

lowering of the pressure may cause secondary ischemia and possibly infarction as pressure drops below the lower range of the patient's autoregulatory capability. In cases where there is an identified cause of the syndrome, these etiologies should be treated promptly, including discontinuation of offending substances such as calcineurin inhibitors in toxic processes, treatment of immune-mediated disorders such as thrombotic thrombocytopenic purpura (TTP), and prompt delivery of the fetus in eclampsia. Seizures must be identified and controlled, often necessitating continuous EEG monitoring. Anticonvulsants are effective when seizure activity is identified, but in the special case of eclampsia, there is evidence to support the use of magnesium sulfate for seizure control.

POST-CARDIAC BYPASS BRAIN INJURY

Central nervous system (CNS) injuries following open heart or coronary artery bypass grafting (CABG) surgery are common and include acute encephalopathy, stroke, and a chronic syndrome of cognitive impairment. Hypoperfusion and embolic disease are frequently involved in the pathogenesis of these syndromes, although multiple mechanisms may be involved in these critically ill patients who are at risk for various metabolic and polypharmaceutical complications.

The frequency of hypoxic injury secondary to inadequate blood flow intraoperatively has been markedly decreased by the use of modern surgical and anesthetic techniques. Despite these advances, some patients still experience neurologic complications from cerebral hypoperfusion or may suffer focal ischemia from carotid or focal intracranial stenoses in the setting of regional hypoperfusion. Postoperative infarcts in the border zones between vascular territories commonly are blamed on systemic hypotension, although these infarcts can also result from embolic disease (Fig. 463e-2).

Embolic disease is likely the predominant mechanism of cerebral injury during cardiac surgery as evidenced by diffusion-weighted MRI and intraoperative transcranial Doppler studies. It should be noted that some of the emboli that are found histologically in these

patients are too small to be detected by standard imaging sequences; therefore, a negative MRI after surgery does not exclude the diagnosis of emboli-related complications. Thrombus in the heart itself as well as atheromas in the aortic arch can become dislodged during cardiac surgeries, releasing a shower of particulate matter into the cerebral circulation. Cross-clamping of the aorta, manipulation of the heart, extracorporeal circulation techniques ("bypass"), arrhythmias such as atrial fibrillation, and introduction of air through suctioning have all been implicated as potential sources of emboli. Histologic studies indicate that literally millions of tiny emboli may be released, even using modern surgical techniques.

This shower of microemboli results in a number of clinical syndromes. Occasionally, a single large embolus leads to an isolated large-vessel stroke that presents with obvious clinical focal deficits. More commonly, the emboli released are multiple and smaller. When there is a high burden of these small emboli, an acute encephalopathy can occur postoperatively, presenting as either a hyperactive or hypoactive confusional state, the latter of which is frequently and incorrectly ascribed to depression or a sedative-induced delirium. When the burden of microemboli is lower, no acute syndrome is recognized, but the patient may suffer a chronic cognitive deficit. Cardiac surgery can be viewed, like delirium, as a "stress test for the brain." Some patients with a low cerebral reserve due to underlying cerebrovascular disease or an early neurodegenerative process will develop a chronic, cognitive deficit, whereas others with higher reserves may remain asymptomatic despite a similar dose of microemboli. In this manner, cardiac surgery may serve to unmask the early manifestations of neurodegenerative disorders such as Alzheimer's disease.

Since modern techniques have successfully minimized hypoperfusion complications during these surgeries, much attention is now focused on reducing this inevitable shower of microemboli. Off-pump CABG surgeries have the advantages of reducing length of stay and perioperative complications; however, off-pump CABG probably does not preserve cognitive function compared with on-pump CABG.

Filters placed in the aortic arch may have some promise in capturing these emboli, although convincing evidence is lacking. Development of successful endovascular operative approaches may provide a reasonable alternative to conventional CABG procedures, especially for patients at high risk of developing cognitive dysfunction after surgery due to advanced age, previous stroke, underlying neurodegenerative disorders, or severe atheromatous disease of the carotid arteries or aortic arch.

POST-SOLID ORGAN TRANSPLANT BRAIN INJURY

Patients who have undergone solid organ transplantation are at risk for neurologic injury in the postoperative period and for months to years thereafter. Neurologic consultants should view these patients as a special population at risk for both unique neurologic complications as well as for the usual disorders found in any critically ill inpatient.

Immunosuppressive medications are administered in high doses to patients after solid organ transplant, and many of these compounds have well-described neurologic complications. In patients with headache, seizures, or focal neurologic deficits taking calcineurin inhibitors, the diagnosis of hyperperfusion syndrome should be considered, as discussed above. This neurotoxicity occurs mainly with cyclosporine and tacrolimus and can present even in the setting of normal serum drug levels. Treatment primarily involves lowering the drug dosage or discontinuing the drug. Sirolimus has very few recorded cases of neurotoxicity and may be a reasonable alternative for some patients. Other examples of immunosuppressive medications and their neurologic complications include OKT3-associated akinetic mutism and the leukoencephalopathy seen with methotrexate, especially when it is administered intrathecally or with concurrent radiotherapy. In any solid organ transplant patient with neurologic complaints, a careful examination of the medication list is required to search for these possible drug effects.

Cerebrovascular complications of solid organ transplant are often first recognized in the immediate postoperative period. Border zone territory infarctions can occur, especially in the setting of systemic hypotension during cardiac transplant surgery. Embolic infarctions classically complicate cardiac transplantation, but all solid organ transplant procedures place patients at risk for systemic emboli. When cerebral embolization accompanies renal or liver transplantation surgery, a careful search for right-to-left shunting should include evaluation of the heart with agitated saline echocardiography (i.e., “bubble study”), as well as looking for intrapulmonary shunting. Renal and some cardiac transplant patients often have advanced atherosclerosis, providing yet another mechanism for stroke. Imaging with CT or MRI with diffusion is advised when cerebrovascular complications are suspected to confirm the diagnosis and to exclude intracerebral hemorrhage, which most often occurs in the setting of coagulopathy secondary to liver failure or after cardiac bypass procedures.

Given that patients with solid organ transplants are chronically immunosuppressed, infections are a common concern (Chap. 169). In any transplant patient with new CNS signs or symptoms such as seizure, confusion, or focal deficit, the diagnosis of a CNS infection should be considered and evaluated through imaging (usually MRI) and possibly lumbar puncture. The most common pathogens responsible for CNS infections in these patients vary based on time since transplant. In the first month posttransplant, common pathogens include the usual bacterial organisms associated with surgical procedures and indwelling catheters. Starting in the second month posttransplant, opportunistic infections of the CNS become more common, including *Nocardia* and *Toxoplasma* species as well as fungal infections such as aspergillosis. Viral infections that can affect the brain of the immunosuppressed patient, such as herpes simplex virus, cytomegalovirus, human herpesvirus type 6 (HHV-6), and varicella, also become more common after the first month posttransplant. After 6 months posttransplant, immunosuppressed patients still remain at risk for these opportunistic bacterial, fungal, and viral infections but can also suffer late CNS infectious complications such as progressive multifocal leukoencephalopathy (PML) associated with JC virus and Epstein-Barr virus–driven clonal expansions of B cells resulting in posttransplant lymphoproliferative disorder or CNS lymphoma.

COMMON NEUROLOGIC COMPLICATIONS OF ELECTROLYTE DISTURBANCES

A wide variety of neurologic conditions can result from abnormalities in serum electrolytes, and consideration of electrolyte disturbances should be part of any inpatient neurologic consultation. **A complete general discussion of fluid and electrolyte imbalance and homeostasis can be found in Chap. 63.**

HYPERNATREMIA AND HYPEROSMOLALITY

The normal range of serum osmolality is around 275–295 mOsm/kg, but neurologic manifestations are usually seen only at levels >325 mOsm/kg. Hyperosmolality is usually due to hypernatremia, hyperglycemia, azotemia, or the addition of extrinsic osmoles such as mannitol, which is commonly used in critically ill neurologic patients. Hyperosmolality itself can lead to a generalized encephalopathy that is nonspecific and without focal findings; however, an underlying lesion such as a mass can become symptomatic under the metabolic stress of a hyperosmolar state, producing focal neurologic signs. Some patients with hyperosmolality from severe hyperglycemia can present, for unclear reasons, with generalized seizures or unilateral movement disorders, which usually respond to lowering of the serum glucose. The treatment of all forms of hyperosmolality involves calculation of apparent water losses and slow replacement so that the serum sodium declines no faster than 2 mmol/L (2 meq/L) per hour.

Hypernatremia leads to the loss of intracellular water, leading to cell shrinkage. In the cells of the brain, solutes such as glutamine and urea are generated under these conditions in order to minimize this shrinkage. Despite this corrective mechanism, when hypernatremia is severe (serum sodium >160 mmol/L [>160 meq/L]) or occurs rapidly, cellular metabolic processes fail and encephalopathy will result. There are many etiologies of hypernatremia including, most commonly, renal and extrarenal losses of water. Causes of neurologic relevance include central diabetes insipidus, where hyperosmolality is accompanied by submaximal urinary concentration due to inadequate release of arginine vasopressin (AVP) from the posterior pituitary, resulting often from pituitary injury in the setting of surgery, hemorrhage, infiltrative processes, or cerebral herniation.

HYPONATREMIA

Hyponatremia is commonly defined as a serum sodium <135 mmol/L (<135 meq/L). Neurologic symptoms occur at different levels of low sodium, depending not only on the absolute value but also on the rate of fall. In patients with hyponatremia that develops over hours, life-threatening seizures and cerebral edema may occur at values as high as 125 mmol/L. In contrast, some patients with more chronic hyponatremia that has slowly developed over months to years may be asymptomatic even with serum levels <110 mmol/L. Correction of hyponatremia, especially when chronic, must take place slowly in order to avoid additional neurologic complications. Cells in the brain swell in hypotonic hyponatremic states but may compensate over time by excreting solute into the extracellular space, leading to restoration of cell volume when water follows the solute out of the cells. If treatment of hyponatremia results in a rapid rise in serum sodium, cells in the brain may quickly shrink, leading to osmotic demyelination, a process that previously was thought to be limited exclusively to the brainstem (central pontine myelinolysis; see Fig. 330-6), but now has been described elsewhere in the CNS.

Treatment of hyponatremia is dependent on the cause. Hypertonic hyponatremia treatment focuses on correcting the underlying condition, such as hyperglycemia. Isovolemic hyponatremia (syndrome of inappropriate antidiuretic hormone [SIADH]) is managed with water restriction or administration of AVP antagonists. The management of choice for patients with hypervolemic hypotonic hyponatremia is free-water restriction and treatment of the underlying edematous disorder, such as hepatic failure, nephrotic syndrome, or congestive heart failure. Finally, in hypovolemic hypotonic hyponatremia, volume is replaced with isotonic saline while underlying conditions of the kidneys, adrenals, and gastrointestinal tract are addressed.

One neurologic cause of hypovolemic hypotonic hyponatremia is the cerebral salt-wasting syndrome that accompanies subarachnoid hemorrhage and, less commonly, other cerebral processes such as meningitis, stroke, or traumatic brain injury. In these cases, the degree of renal sodium excretion can be remarkable, and large amounts of saline, hypertonic saline, or oral sodium may need to be given in a judicious fashion in order to avoid complications from cerebral edema.

HYPOKALEMIA

Hypokalemia, defined as a serum potassium level < 3.5 mmol/L (< 3.5 meq/L), occurs either because of excessive potassium losses (from the kidneys or gut) or due to an abnormal potassium distribution between the intracellular and extracellular spaces. At very low levels (< 1.5 mmol/L), hypokalemia may be life threatening due to the risk of cardiac arrhythmia and may present neurologically with severe muscle weakness and paralysis. Hypokalemic periodic paralysis is a rare disorder caused by excessive intracellular potassium uptake in the setting of a calcium or sodium channel mutation. Treatment of hypokalemia is dependent on the etiology but usually includes replacement of potassium through oral or IV routes as well as correcting the cause of potassium balance problems (e.g., eliminating β_2 -adrenergic agonist medications or treating the underlying cause of severe diarrhea).

HYPERKALEMIA

Hyperkalemia is defined as a serum potassium level > 5.5 mmol/L (> 5.5 meq/L) and can neurologically present as muscle weakness with or without paresthesias. Hyperkalemia becomes life threatening when it produces electrocardiographic abnormalities such as peaked T waves or a widened QRS complex. In these cases, prompt treatment is essential and consists of strategies that protect the heart against arrhythmias (calcium gluconate administration); promote potassium redistribution into cells (with glucose, insulin, and β_2 -agonist medications); and increase potassium removal (through sodium polystyrene sulfonate, loop diuretics, or hemodialysis).

CALCIUM DISTURBANCES

Hypercalcemia usually occurs in the setting of either hyperparathyroidism or systemic malignancy. Neurologic manifestations include encephalopathy as well as muscle weakness due to reduced neuromuscular excitability. Seizures can occur but are more common in states of low calcium.

Hypocalcemia in adults often follows surgical treatment of the thyroid or parathyroid. Seizures and altered mental status dominate the neurologic picture and usually resolve with calcium repletion. Tetany is due to spontaneous, repetitive action potentials in peripheral nerves and remains the classic sign of symptomatic hypocalcemia.

MAGNESIUM DISTURBANCES

Disorders of magnesium are difficult to correlate with serum levels because a very small amount of total-body magnesium is located in the extracellular space. Hypomagnesemia presents neurologically with seizures, tremor, and myoclonus. When intractable seizures occur in the setting of hypomagnesemia, only administration of magnesium will lead to their resolution.

High levels of magnesium, in contrast, lead to CNS depression. Hypermagnesemia usually only occurs in the setting of renal failure or magnesium administration and can lead to confusion and muscular paralysis when severe.

CONSULTATIONS REGARDING PERIPHERAL NERVOUS SYSTEM DYSFUNCTION

ENTRAPMENT NEUROPATHIES

Polyneuropathy is a common cause of outpatient neurologic consultation (Chap. 459). In the inpatient setting, however, mononeuropathies are more frequent, especially the entrapment neuropathies that complicate many surgical procedures and medical conditions. Median neuropathy at the wrist (carpal tunnel syndrome) is the most frequent entrapment neuropathy by far, but it is rarely a cause for inpatient

consultation. Mechanisms for perioperative mononeuropathy include traction, compression, and ischemia of the nerve. Imaging with MRI, including neurography, may allow these causes to be distinguished. In all cases of mononeuropathy, the diagnosis can be made through the clinical examination and then confirmed with electrodiagnostic studies in the subacute period, if necessary. Treatment consists mainly of avoidance of repetitive trauma but may also include surgical approaches to relieve pressure or repair the nerve.

RADIAL NEUROPATHY

Radial nerve injury classically presents with weakness of extension of the wrist and fingers ("wrist drop") with or without more proximal weakness of extensor muscles of the upper extremity, depending on the site of injury. Sensory loss is in the distribution of the radial nerve, which includes the dorsum of the hand (Fig. 463e-3A). Compression at the level of the axilla, e.g., resulting from use of crutches, leads to weakness of the triceps, brachioradialis, and supinator muscles in addition to wrist drop. A more common site of compression occurs in the spiral groove of the upper arm in the setting of a humerus fracture or from sleeping with the arm draped over a bench or chair ("Saturday night palsy"). Sparing of the triceps is the rule when the nerve is injured in this location. Because extensors of the upper extremity are injured preferentially in radial nerve injury, these lesions may be mistaken for the pyramidal distribution of weakness that accompanies upper motor neuron lesions from brain or spinal cord processes, prompting neuroimaging to exclude acute stroke or mass lesion.

ULNAR NEUROPATHY

Compression of the ulnar nerve is the second most common entrapment neuropathy after carpal tunnel syndrome. The most frequent site of compression is at the elbow where the nerve passes superficially in the ulnar groove. Symptoms usually begin with tingling in the ulnar distribution, including the fourth and fifth digits of the hand (Fig. 463e-3B). Sensory symptoms may be worsened by elbow flexion due to increased pressure on the nerve, hence the tendency of patients to complain of increasing paresthesias at night when the arm is flexed at the elbow during sleep. Motor dysfunction can be disabling and involves most of the intrinsic hand muscles, limiting dexterity and strength of grasp and pinch. Etiologies of ulnar entrapment include trauma to the nerve (hitting the "funny bone"), malpositioning during anesthesia for surgical procedures, and chronic arthritis of the elbow. When a perioperative ulnar nerve injury is considered, stretch injury or trauma to the lower trunk of the brachial plexus should be entertained as well since its symptoms can mimic those of an ulnar neuropathy. If the clinical examination is equivocal, electrodiagnostic studies can definitively distinguish between plexus and ulnar nerve lesions a few weeks after the injury. Conservative methods of treatment are often the first step including bracing of the elbow to prevent flexion while sleeping. A variety of surgical approaches may also be effective in refractory or recurrent cases, including anterior ulnar nerve transposition and release of the flexor carpi ulnaris aponeurosis.

PERONEAL NEUROPATHY

The peroneal (also known as the fibular) nerve winds around the head of the fibula in the leg below the lateral aspect of the knee, and its superficial location at this site makes it vulnerable to trauma. Patients present with weakness of foot dorsiflexion ("foot drop") as well as with weakness in eversion but not inversion at the ankle. Sparing of inversion, which is a function of muscles innervated by the tibial nerve, helps to distinguish peroneal neuropathies from L5 radiculopathies. Sensory loss involves the lateral aspect of the leg as well as the dorsum of the foot (Fig. 463e-3C). Fractures of the fibular head may be responsible for peroneal neuropathies, but in the perioperative setting, poorly applied braces exerting pressure on the nerve while the patient is unconscious, often in lithotomy position, are more often responsible. Tight-fitting stockings or casts of the upper leg can also cause a peroneal neuropathy, and thin individuals and those with recent weight loss are at increased risk.

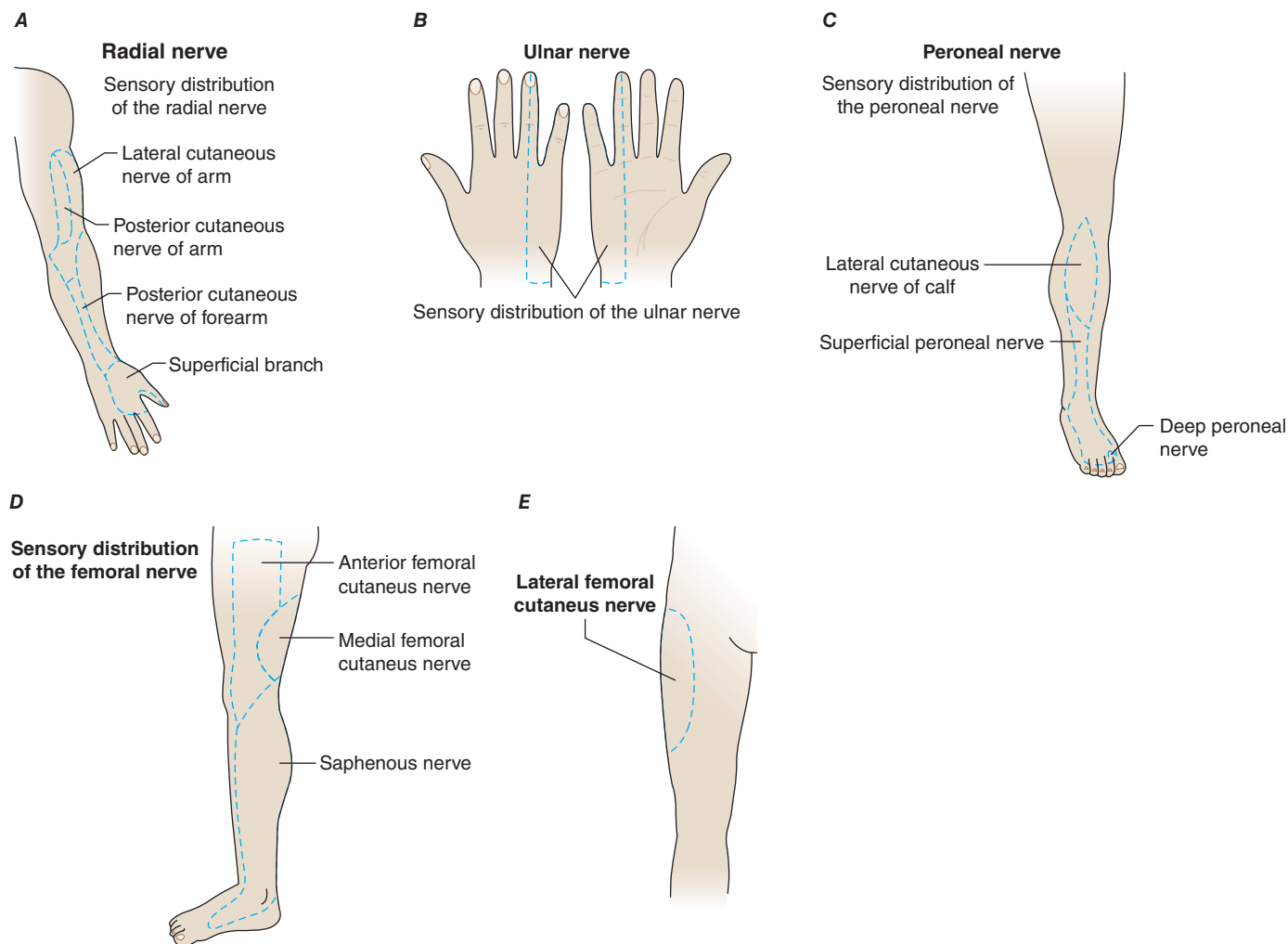


FIGURE 463e-3 Sensory distribution of peripheral nerves commonly affected by entrapment neuropathies. **A.** Radial nerve. **B.** Ulnar nerve. **C.** Peroneal nerve. **D.** Femoral nerve. **E.** Lateral femoral cutaneous nerve.

PROXIMAL FEMORAL NEUROPATHY

Lesions of the proximal femoral nerve are relatively uncommon but may present dramatically with weakness of hip flexion, quadriceps atrophy, weakness of knee extension (often manifesting with leg-buckling falls), and an absent patellar reflex. Adduction of the thigh is spared as these muscles are supplied by the obturator nerve, thereby distinguishing a femoral neuropathy from a more proximal lumbosacral plexus lesion. The sensory loss found is in the distribution of the femoral nerve sensory branches including the anterior part of the thigh (Fig. 463e-3D). Compressive lesions from retroperitoneal hematomas or masses are common, and a CT of the pelvis should be obtained in all cases of femoral neuropathy to exclude these conditions. Bleeding into the pelvis resulting in hematoma can occur spontaneously, following trauma, or after intrapelvic surgeries such as renal transplantation. In intoxicated or comatose patients, stretch injuries to the femoral nerve are seen following prolonged, extreme hip flexion or extension. Rarely, attempts at femoral vein or arterial puncture can be complicated by injury to this nerve.

LATERAL FEMORAL CUTANEOUS NERVE

The symptoms of lateral femoral cutaneous nerve entrapment, commonly known as “meralgia paresthetica,” include sensory loss, pain, and dysesthesia in part of the area supplied by the nerve (Fig. 463e-3E). There is no motor component to the nerve, and therefore weakness is

not a part of this syndrome. Symptoms often are worsened by standing or walking. Compression of the nerve occurs where it enters the leg near the inguinal ligament, usually in the setting of tight-fitting belts, pants, corsets, or recent weight gain, including that of pregnancy. The differential diagnosis of these symptoms includes hip problems such as trochanteric bursitis.

OBSTETRIC NEUROPATHIES

Pregnancy and delivery place women at special risk for a variety of nerve injuries. Radiculopathy due to a herniated lumbar disc is not common during pregnancy, but compressive injuries of the lumbosacral plexus do occur secondary to either the fetal head passing through the pelvis or the use of forceps during delivery. These plexus injuries are more frequent with cephalopelvic disproportion and often present with a painless unilateral foot drop which must be distinguished from a peroneal neuropathy caused by pressure on the nerve while in lithotomy position during delivery. Other compressive mononeuropathies of pregnancy include meralgia paresthetica, carpal tunnel syndrome, femoral neuropathy when the thigh is abducted severely in an effort to facilitate delivery of the fetal shoulder, and isolated obturator neuropathy during lithotomy positioning. The latter presents with medial thigh pain that may be accompanied by weakness of thigh adduction. There is also a clear association between pregnancy and an increased frequency of idiopathic facial palsy (Bell’s palsy).

SECTION 4 CHRONIC FATIGUE SYNDROME

464e Chronic Fatigue Syndrome

Gijs Bleijenberg, Jos W. M. van der Meer

DEFINITION

Chronic fatigue syndrome (CFS) is a disorder characterized by persistent and unexplained fatigue resulting in severe impairment in daily functioning. Besides intense fatigue, most patients with CFS report concomitant symptoms such as pain, cognitive dysfunction, and unrefreshing sleep. Additional symptoms can include headache, sore throat, tender lymph nodes, muscle aches, joint aches, feverishness, difficulty sleeping, psychiatric problems, allergies, and abdominal cramps. Criteria for the diagnosis of CFS have been developed by the U.S. Centers for Disease Control and Prevention ([Table 464e-1](#)).

EPIDEMIOLOGY



CFS is seen worldwide, with adult prevalence rates varying between 0.2% and 0.4%. In the United States, the prevalence is higher among women (~75% of cases), members of minority groups (African and Native Americans), and individuals with lower levels of education and occupational status. The mean age of onset is between 29 and 35 years. Many patients probably go undiagnosed and/or do not seek help.

ETIOLOGY

There are numerous hypotheses about the etiology of CFS; there is no definitively identified cause. Distinguishing between predisposing, precipitating, and perpetuating factors in CFS helps to provide a framework for understanding this complex condition ([Table 464e-2](#)).

Predisposing Factors Physical inactivity and trauma in childhood tend to increase the risk of CFS in adults. Neuroendocrine dysfunction may be associated with childhood trauma, reflecting a biological correlate of vulnerability. Psychiatric illness and physical hyperactivity in adulthood raise the risk of CFS in later life. Twin studies suggest a familial predisposition to CFS, but no causative genes have been identified.

Precipitating Factors Physical or psychological stress may elicit the onset of CFS. Most patients report an infection (usually a flulike illness or infectious mononucleosis) as the trigger of their fatigue. Relatively high percentages of CFS cases follow Q fever and Lyme disease.

TABLE 464e-1 DIAGNOSTIC CRITERIA FOR CHRONIC FATIGUE SYNDROME

Characteristic Persistent or Relapsing Unexplained Chronic Fatigue

Fatigue lasts for at least 6 months.

Fatigue is of new or definite onset.

Fatigue is not the result of an organic disease or of continuing exertion.

Fatigue is not alleviated by rest.

Fatigue results in a substantial reduction in previous occupational, educational, social, and personal activities.

Four or more of the following symptoms are concurrently present for 6 months: impaired memory or concentration, sore throat, tender cervical or axillary lymph nodes, muscle pain, pain in several joints, new headaches, unrefreshing sleep, or malaise after exertion.

Exclusion Criteria

Medical condition explaining fatigue

Major depressive disorder (psychotic features) or bipolar disorder

Schizophrenia, dementia, or delusional disorder

Anorexia nervosa, bulimia nervosa

Alcohol or substance abuse

Severe obesity (body mass index >40)

TABLE 464e-2 PREDISPOSING, PRECIPITATING, AND PERPETUATING FACTORS IN CHRONIC FATIGUE SYNDROME

Predisposing Factors

Childhood trauma (sexual, physical, emotional abuse; emotional and physical neglect)

Physical inactivity during childhood

Premorbid psychiatric illness or psychopathology

Premorbid hyperactivity



Precipitating Factors

Somatic events: infection (e.g., mononucleosis, Q fever, Lyme disease), surgery, pregnancy

Psychosocial stress, life events



Perpetuating Factors

Non-acknowledgment by physician

Negative self-efficacy

Strong physical attributions

Strong focus on bodily symptoms

Fear of fatigue

Lack of social support

Low physical activity pattern

However, no differences in Epstein-Barr virus load and immunologic reactivity were found between individuals who developed CFS and those who did not. While antecedent infections are associated with CFS, a direct microbial causality is unproven and unlikely. One study identified a murine leukemia virus–related retrovirus (XMRV); however, several subsequent studies have established this virus as a laboratory artifact. Patients also often report other precipitating somatic events such as serious injury, surgery, pregnancy, or childbirth. Serious life events, such as the loss of a loved one or a job, military combat, and other stressful situations, may also precipitate CFS. One-third of all patients cannot recall a trigger.

Perpetuating Factors Once CFS has developed, numerous factors may impede recovery. Physicians may contribute to chronicity by ordering unnecessary diagnostic procedures, by persistently suggesting psychological causes, and by not acknowledging CFS as a diagnosis.

A patient's focus on illness and avoidance of activities may perpetuate symptoms. A firm belief in a physical cause, a strong focus on bodily sensations, and a poor sense of control over symptoms may also prolong or exacerbate the fatigue and functional impairment. In most patients, inactivity is caused by negative illness perceptions rather than by poor physical fitness. Solicitous behavior of others may reinforce a patient's illness-related perceptions and behavior. A lack of social support is another known perpetuating factor.

PATHOPHYSIOLOGY

The pathophysiology of CFS is unclear. Neuroimaging studies have found that CFS is associated with reduced gray matter volume, which in turn is associated with a decline in physical activity; these changes have been partially reversed following cognitive behavioral therapy (CBT). In addition, functional MRI data have suggested that abnormal patterns of activation correlate with self-reported problems with information processing. Neurophysiologic studies have shown altered CNS activation patterns during muscle contraction.

Evidence for immunologic dysfunction is inconsistent. Modest elevations in titers of antinuclear antibodies, reductions in immunoglobulin subclasses, deficiencies in mitogen-driven lymphocyte proliferation, reductions in natural killer cell activity, disturbances

How have you felt during the last two weeks?

Please rate all four statements and per statement check the box that reflects your situation best.

1. I feel tired	Yes, that is true	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	No, that is not true
2. I tire easily	Yes, that is true	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	No, that is not true
3. I feel fit	Yes, that is true	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	No, that is not true
4. Physically I feel exhausted	Yes, that is true	<input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/> <input type="checkbox"/>	No, that is not true

Scoring:

1, 2 and 4: Yes, that is true

7	6	5	4	3	2	1
---	---	---	---	---	---	---

 No, that is not true 3: Reversed

Sum scores >18 indicate severe fatigue

FIGURE 464e-1 Shortened fatigue questionnaire.

in cytokine production, and shifts in lymphocyte subsets have been described. None of these immune findings appears in most patients, nor does any correlate with the severity of CFS. In theory, symptoms of CFS could result from excessive production of a cytokine, such as interleukin 1, that induces asthenia and other flulike symptoms; however, compelling data in support of this hypothesis are lacking. There is some evidence that CFS patients have mild hypocortisolism, the degree of which is associated with a poorer response to CBT. Discrepancies in perceived and actual cognitive performance are consistent findings in patients with CFS.

DIAGNOSIS

In addition to a thorough history, a systematic physical examination is warranted to exclude disorders causing fatigue (e.g., endocrine disorders, neoplasms, heart failure). The heart rate of CFS patients is often slightly above normal. Laboratory tests serve primarily to exclude other diagnoses; no test can diagnose CFS. The following laboratory screen usually suffices: complete blood count; erythrocyte sedimentation rate; C-reactive protein; serum creatinine, electrolytes, calcium, and iron; blood glucose; creatine kinase; liver function tests; thyroid-stimulating hormone; anti-gliadin antibodies; and urinalysis. Serology for viral or bacterial infections usually is not helpful. No specific abnormalities have been identified on MRI or CT scans. The decrease in gray matter volume observed at a population level by MRI is not useful for diagnosis in the individual patient. Extensive, unfocused, and expensive testing in a search for the “hidden” cause of the fatigue is not productive. CFS is a constellation of symptoms with no pathognomonic features and remains a diagnosis of exclusion.

Bipolar disorders, schizophrenia, and substance abuse exclude a diagnosis of CFS, as do eating disorders, unless these issues have been resolved ≥ 5 years before symptom onset. In addition, CFS is excluded if the chronic fatigue developed immediately after a depressive episode. Depression developing in the course of the fatigue, however, does not preclude CFS. Concurrent psychiatric disorders, especially anxiety and mood disorders, are present in 30–60% of cases.

INITIAL MANAGEMENT

In cases of suspected CFS, the clinician should acknowledge the impact of the patient’s symptoms on daily functioning. Disbelief or denial can provoke an exacerbation of genuine symptoms, which in turn strengthens the clinician’s disbelief, leading to an unfortunate cycle of miscommunication. The possibility of CFS should be considered if

a patient fulfills all criteria (Table 464e-1) and if other diagnoses have been excluded.

The patient should be asked to describe the symptoms (fatigue and accompanying symptoms) and their duration as well as their consequences (reduction in daily activities). To assess symptom severity and the extent of daily-life impairment, the patient should describe a typical day, from waking to retiring, and, for comparison, an average day prior to symptom onset. Next, potential fatigue-precipitating factors are sought. The severity of fatigue is commonly difficult to assess quantitatively; a brief questionnaire is often helpful (Fig. 464e-1).

The patient should be informed of the current understanding of precipitating and perpetuating factors and effective treatments and should be offered general advice about disease management. If CBT for CFS is not available as an initial option (see below) and depression and anxiety are present, these symptoms should be treated. For patients with headache, diffuse pain, and feverishness, nonsteroidal anti-inflammatory drugs may be helpful. Even modest improvements in symptoms can make an important difference in the patient’s degree of self-sufficiency and ability to appreciate life’s pleasures.

Controlled therapeutic trials have established that acyclovir, fludrocortisone, galantamine, modafinil, and IV immunoglobulin, among other agents, offer no significant benefit in CFS. Countless anecdotes circulate regarding other traditional and nontraditional therapies. It is important to guide patients away from those therapeutic modalities that are toxic, expensive, or unreasonable.

The patient should be encouraged to maintain regular sleep patterns, to remain as active as possible, and to gradually return to previous levels of exercise and other activity (work).

TREATMENT CHRONIC FATIGUE SYNDROME

CBT and graded exercise therapy (GET) have been found to be the only beneficial interventions in CFS. Some patient groups argue against these approaches because of the implication that CFS is a purely mental disorder. CBT is a psychotherapeutic approach directed at changing unhealthy disease-perpetuating patterns of thoughts and behaviors. It includes educating the patient about the etiologic model, setting goals, restoring fixed bedtimes and wake-up times, challenging and changing fatigue- and activity-related concerns, reducing a focus on symptoms, spreading activities

evenly throughout the day, gradually increasing physical activity, planning a return to work, and resuming other activities. The intervention, which typically consists of 12–14 sessions spread over 6 months, helps CFS patients gain control over their symptoms.

GET targets deconditioning and exercise intolerance and usually involves a home exercise program that continues for 3–5 months. Walking or cycling is systematically increased, with set goals for maximal heart rates. Evidence that deconditioning is the basis for symptoms in CFS is lacking, however. CBT and GET appear to improve fatigue primarily by changing the patient's perception of the fatigue and also by reducing the focus on symptoms. In general, CBT is the more multifaceted treatment, which may explain why CBT studies tend to yield better improvement rates than GET trials.

Not all patients benefit from CBT or GET. Predictors of poor outcome are medical (including psychiatric) comorbidities, current disability claims, and severe pain. CBT offered in an early stage of the illness reduces the burden of CFS for the patient as well as for society in terms of decreased medical and disability-related costs.

PROGNOSIS

Full recovery from untreated CFS is rare: the median annual recovery rate is 5% (range, 0–31%), and the median improvement rate is 39% (range, 8–63%). Patients with an underlying psychiatric disorder and those who continue to attribute their symptoms to an undiagnosed medical condition have poorer outcomes.

SECTION 5 PSYCHIATRIC AND ADDICTION DISORDERS

465e **Biology of Psychiatric Disorders**

Robert O. Messing, Eric J. Nestler

Psychiatric disorders are central nervous system diseases characterized by disturbances in emotion, cognition, motivation, and socialization. They are highly heritable, with genetic risk comprising 20–90% of disease vulnerability. As a result of their prevalence, early onset, and persistence, they contribute substantially to the burden of illness worldwide. All psychiatric disorders are broad heterogeneous syndromes that currently lack well-defined neuropathology and *bona fide* biologic markers. Therefore, diagnoses continue to be made solely from clinical observations using criteria in the *Diagnostic and Statistical Manual of Mental Disorders* (DSM) of the American Psychiatric Association, which is in its fifth edition as of 2013.

There is increasing agreement that the classification of psychiatric illnesses in DSM does not accurately reflect the underlying biology of these disorders. Uncertainties in diagnosis make it extremely difficult to study the neurobiologic and genetic basis of mental illness. This has led to the development of an alternative diagnostic scheme, termed Research Domain Criteria (RDoCs), which classifies mental illness on the basis of core abnormalities—e.g., psychosis (loss of reality) or anhedonia (decreased ability to experience pleasure), which are common symptoms of several illnesses—with the idea that such classifications will assist in defining the biologic basis of at least key symptoms. Other factors that have impeded progress in understanding mental illness include the lack of access to pathologic brain tissue except upon death and the inherent limitations of animal models for disorders defined largely by behavioral abnormalities (e.g., hallucinations, delusions, guilt, suicidality) that are inaccessible in animals.

Despite these limitations, the past decade has seen significant advances. Neuroimaging methods are beginning to provide evidence of brain pathology, genome-wide association studies and high-throughput sequencing are at last revealing genes that confer risk for severe forms of mental illness, and investigations using better validated animal models are offering new insight into the molecular, cellular, and circuit mechanisms of disease pathogenesis. There is also excitement in the potential utility of neuron-like cells induced in vitro from a patient's peripheral tissues (e.g., fibroblasts) providing novel ways of studying disease pathophysiology and screening for new treatments. There is consequently justified optimism that the field of psychiatry will transition from behaviorally defined syndromes to true biologic disease entities and that such advances will drive the development of improved treatments and eventually cures and preventive measures. This chapter describes several examples of recent discoveries in basic neuroscience that have informed our current understanding of disease mechanisms in psychiatry.

NEUROGENETICS

Because the human brain can only be examined indirectly during life, genome analyses have been extremely important for obtaining molecular clues about the pathogenesis of psychiatric disorders. A wealth of new information has been made possible by recent technological developments that have permitted affordable, large-scale genome-wide association studies and fine-scale sequencing. As an example, significant progress has been made in the genetics of autism spectrum disorders (ASDs), which are a heterogeneous group of neurodevelopmental diseases that share clinical features of impaired reciprocal social communication and interaction and restricted, repetitive patterns of behavior. ASDs are highly heritable; concordance rates in monozygotic twins (~60–90%) are roughly 10-fold higher than in

dizygotic twins and siblings, whereas first-degree relatives show about 50-fold increased risk compared with the general population. ASDs are also genetically heterogeneous. More than 100 known mutations account for up to 20% of cases, although none individually accounts for more than 1% (Table 465e-1). It appears that most cases result from complex genetic mechanisms, including inheritance of multiple genetic risk variants and epigenetic modifications. For example, up to 10% of patients with autism have large (>500 kb) *de novo* copy number variations scattered across the genome, suggesting that hundreds of different genes can influence autism risk.

Amid this genetic heterogeneity, however, some common themes have emerged that inform pathogenesis of ASDs. For instance, many identified mutations are in genes that encode proteins involved in synaptic function and early transcriptional regulation (Table 465e-1) and have a clear relationship to activity-dependent neural responses that can affect the development of neural systems underlying cognition and social behaviors. Mutations in these genes may be detrimental by altering the balance of excitatory versus inhibitory synaptic signaling in local and extended circuits and by altering mechanisms that control brain growth. Some mutations affect genes (e.g., *PTEN*, *TSC1*, and *TSC2*) that negatively regulate signaling from several types of extracellular stimuli, including those transduced by receptor tyrosine kinases. Their dysregulation can alter neuronal growth as well as synaptic development and function.

With further understanding of pathogenesis and the definition of specific ASD subtypes, there is reason to believe that effective therapies will be identified. Work in mouse models has already demonstrated that some autism-like behaviors can be reversed, even in fully developed adult animals, by modifying the underlying pathology; these results encourage hope for many affected individuals. Treatments that target excitation-inhibition imbalance or altered mRNA translation appear to offer early promise. For example, the genes *TSC1*, *TSC2*, and *PTEN* are negative regulators of signaling through the target of rapamycin complex 1 (TORC1), which regulates protein synthesis. Rapamycin, a selective inhibitor of TORC1, can reverse several behavioral and synaptic defects in mice carrying null mutations in these genes. Another example is fragile X syndrome, which is the leading cause of inherited autism and mental disability and is due to mutations in *FMR1* that result in loss of the encoded fragile X mental retardation protein (FMRP). FMRP is a polyribosome-associated mRNA-binding protein that represses the translation of a subset (~5%) of all mRNAs, several of which encode proteins that comprise the postsynaptic density, including the metabotropic glutamate receptor 5 (mGluR5). Treatment of *Fmr1* knockout mice with mGluR5 antagonists reduces several behavioral and morphologic abnormalities in these mice; these promising preclinical results have led to ongoing trials of mGluR5 antagonists in humans with fragile X syndrome.

The ability to catalog common genetic variants and assay them on array-based platforms and, more recently, to carry out whole-exome sequencing has allowed investigators to collect sample sizes sufficient to detect genetic risk loci for schizophrenia with genome-wide significance. Several of the identified genes are parts of molecular complexes, such as voltage-gated calcium channels (in particular, *CACNA1C* and *CACNB2*) and the postsynaptic density of excitatory synapses. As in ASDs, copy number variants, single-nucleotide polymorphisms, and small insertions and deletions are common in schizophrenia. Genes that promote risk for drug addiction have also begun to emerge from large family and population studies. The best-established susceptibility loci are regions on chromosomes 4 and 5 containing GABA_A receptor gene clusters linked to alcoholism and the *CHRNA5-A3-B4* nicotinic acetylcholine receptor gene cluster on chromosome 15 associated with nicotine and alcohol addiction.

A recurrent theme that has emerged from genetic studies of psychiatric disorders is pleiotropy, namely, that many genes are associated

TABLE 465e-1 EXAMPLES OF GENES IMPLICATED IN AUTISM

Gene Symbol	Gene Name	Function
<i>PTEN</i>	Phosphatase and tensin homolog	Signal transduction Synaptic function
<i>TSC1</i>	Tuberous sclerosis 1	Signal transduction Translation and protein stability Synaptic function
<i>TSC2</i>	Tuberous sclerosis 2	Signal transduction Translation and protein stability Synaptic function
<i>DYRK1A</i>	Dual-specificity tyrosine-(Y)-phosphorylation regulated kinase	Signal transduction
<i>FMR1</i>	Fragile X mental retardation 1	Translation and protein stability Synaptic function
<i>UBE3A</i>	Ubiquitin protein ligase E3A	Translation and protein stability Synaptic function
<i>CNTN3</i>	Contactin 3	Synaptic function
<i>CNTN4</i>	Contactin 4	Synaptic function
<i>CNTNAP2</i>	Contactin-associated protein-like 2	Synaptic function
<i>NLGN3</i>	Neuroigin 3	Synaptic function
<i>NLGN4</i>	Neuroigin 4	Synaptic function
<i>NRXN1</i>	Neurexin 1	Synaptic function
<i>PCDH10</i>	Protocadherin 10	Synaptic function
<i>SHANK3</i>	Shank 3	Synaptic function
<i>SLC6A4</i>	Serotonin transporter	Neurotransmitter signaling
<i>AVPR1</i>	Arginine vasopressin receptor 1	Neurotransmitter signaling
<i>OXTR</i>	Oxytocin receptor	Neurotransmitter signaling
<i>CACNA1C</i>	Voltage-gated calcium channel- α 1C subunit	Ion channel
<i>CACNA1H</i>	Voltage-gated calcium channel- α 1H subunit	Ion channel
<i>SCN1A</i>	Sodium channel, voltage-gated, type I, α subunit	Ion channel
<i>SCN2A</i>	Sodium channel, voltage-gated, type II, α subunit	Ion channel
<i>SLC9A9</i>	Sodium/hydrogen exchanger	Transporter
<i>GRIN2B</i>	Glutamate receptor, ionotropic, <i>N</i> -methyl-D-aspartate 2B	Neurotransmitter signaling
<i>DHCR7</i>	7-Dehydrocholesterol reductase	Metabolism
<i>PAH</i>	Phenylalanine hydroxylase	Metabolism
<i>ARX</i>	Arx transcription factor	Gene expression
<i>EN2</i>	Engrailed 2 transcription factor	Gene expression
<i>MECP2</i>	Methyl CpG-binding protein 2 (Rett's syndrome)	Gene expression
<i>RNF8</i>	Ring finger protein 8	Gene expression
<i>CHD8</i>	Chromodomain helicase DNA binding protein	Gene expression
<i>TBR1</i>	T-box, brain 1	Gene expression

with multiple psychiatric syndromes. For example, mutations in *MECP2*, *FMR1*, and *TSC1* and *TSC2* (see Table 465e-1 for abbreviations) can cause mental retardation without ASD, others in *MECP2* can cause obsessive-compulsive and attention-deficit hyperactivity disorders, some alleles of *NRXN1* are associated with symptoms of both ASD and schizophrenia, and common polymorphisms in *CACNA1C* are strongly associated with both schizophrenia and bipolar disorder. Likewise, duplication of chromosome 16p is associated with both schizophrenia and autism, whereas DiGeorge's (velocardiofacial) syndrome region deletions and the *DISC1* (disrupted in schizophrenia 1) locus on chromosome 22 are associated with schizophrenia, autism, and bipolar disorder. The association of genes with multiple syndromes attests to the complexity of psychiatric disorders and the influence of additional factors that combine to specify the ultimate phenotype, including regulatory variants that determine cell-type specificity and timing of gene expression, protective variants, and epigenetic effects.

SIGNAL TRANSDUCTION

Studies of signal transduction have revealed numerous intracellular signaling pathways that are perturbed in psychiatric disorders, and such research has provided insight into development of new therapeutic agents. For example, lithium is a highly effective drug for bipolar disorder and competes with magnesium to inhibit numerous magnesium-dependent enzymes, including the enzyme GSK3 β and several enzymes involved in phosphoinositide signaling that lead to activation of protein kinase C. These findings have led to discovery programs focused on developing GSK3 β or protein kinase C inhibitors as potential novel treatments for mood disorders.

The observations that tricyclic antidepressants (e.g., imipramine) inhibit serotonin and/or norepinephrine reuptake and that monoamine oxidase inhibitors (e.g., tranylcypromine) are effective antidepressants initially led to the view that depression is caused by a deficiency of these monoamines. However, this hypothesis has not been substantiated. A cardinal feature of these drugs is that long-term administration is needed for their antidepressant effects. This means that their short-term actions, namely promotion of serotonin or norepinephrine function, are not per se antidepressant but rather induce a cascade of adaptations in the brain that underlie their clinical effects. The nature of these therapeutic drug-induced adaptations has not been identified with certainty. One theory holds that, in a subset of depressed patients who display upregulation of the hypothalamic-pituitary-adrenal (HPA) axis characterized by increased secretion of corticotropin-releasing factor (CRF) and glucocorticoids, excessive glucocorticoids cause atrophy of hippocampal neurons, which is associated with reduced hippocampal volumes seen clinically. Chronic antidepressant administration might reverse this atrophy by increasing brain-derived neurotrophic factor (BDNF) in hippocampus. A role for stress-induced decreases in the generation of newly born hippocampal granule cell neurons, and its reversal by antidepressants through BDNF and other growth factors, has also been suggested.

A major advance in recent years has been the identification of several rapidly acting antidepressants with non-monoamine-based mechanisms of action. The best established is ketamine, a noncompetitive antagonist of *N*-methyl-D-aspartate (NMDA) glutamate receptors, which exerts rapid (hours) and robust antidepressant effects in severely depressed patients who have not responded to other treatments. Ketamine, which at higher doses is psychotomimetic and anesthetic, exerts these antidepressant effects at low doses with minimal side effects. However, the response to ketamine is transient, which has led to several approaches to maintain treatment response, such as repeated ketamine delivery. The mechanism underlying ketamine's antidepressant action is not known, but its striking clinical efficacy has stimulated animal research on the role of glutamate neurotransmission and synaptic plasticity in key limbic regions. Recent evidence supports a role for TORC1 activation because administration of rapamycin blocks the antidepressant-like effects of ketamine in animal models. Mechanisms by which ketamine activates TORC1 are currently an active area of investigation.

A major goal in the field of drug abuse has been to identify neuroadaptive mechanisms that lead from recreational use to addiction. Such research has determined that repeated intake of abused drugs induces specific changes in cellular signal transduction, leading to changes in synaptic strength (long-term potentiation or depression) and neuronal structure (altered dendritic branching or cell soma size) within the brain's reward circuitry. These modifications are mediated in part by changes in gene expression, achieved by drug regulation of transcription factors (e.g., CREB [cAMP response element-binding protein] and Δ FosB [a Fos family protein]) and their target genes. Such alterations in gene expression are associated with lasting alterations in epigenetic modifications, including histone acetylation and methylation and DNA methylation. These adaptations provide opportunities for developing treatments targeted to drug-addicted individuals. The fact that the spectrum of these adaptations partly differs depending on the particular addictive substance used creates opportunities for treatments that are specific for different classes of addictive drugs and that may, therefore, be less likely to disturb basic mechanisms that govern normal motivation and reward.

Increasingly, causal relationships are being established between individual molecular and cellular adaptations and specific behavioral abnormalities that characterize the addicted state. For example, acute activation of μ -opioid receptors by morphine or other opiates activates $G_{i/o}$ proteins, leading to inhibition of adenylyl cyclase, resulting in reduced cyclic AMP (cAMP) production, protein kinase A (PKA) activation, and activation of the transcription factor CREB. Repeated administration of these drugs (Fig. 465e-1) evokes a homeostatic response involving upregulation of adenylyl cyclases and PKA and increased activation of CREB. Such upregulation of cAMP-CREB signaling has been identified in the locus coeruleus, periaqueductal gray, ventral tegmental area (VTA), nucleus accumbens, and several other CNS regions, and contributes to opiate craving and signs of opiate withdrawal. The fact that endogenous opioid peptides do not produce tolerance and dependence, while morphine and heroin do, may relate to the observation that, unlike endogenous opioids, morphine and heroin are weak inducers of μ -opioid receptor desensitization and endocytosis. Therefore, these drugs cause prolonged receptor activation and inhibition of adenylyl cyclases, which provides a powerful stimulus for the upregulation of cAMP-CREB signaling that characterizes the opiate-dependent state.

SYSTEMS NEUROSCIENCE

The study of interconnected brain circuits that drive behavior has been greatly advanced through newer methods in brain imagining that have documented abnormalities in neural function and connectivity in psychiatric disorders. The past decade has also witnessed the development of revolutionary new techniques—optogenetics and designer receptors and ligands—that provide unprecedented temporal and spatial control of neural circuits and permit detection of neural activity in real time in awake, behaving animals.

Positron emission tomography (PET), diffusion tensor imaging (DTI), and functional magnetic resonance imaging (fMRI) have identified neural circuits that contribute to psychiatric disorders, for example, defining the neural circuitry of mood within the brain's limbic system (Fig. 465e-2). Integral to this system are the nucleus accumbens (important also for brain reward—see below), amygdala, hippocampus, and regions of prefrontal cortex. Recent optogenetic research in animals, where the activity of specific types of neurons in defined circuits can be controlled with light, has confirmed the importance of this limbic circuitry in controlling depression-related behavioral abnormalities. Given that many symptoms of depression (so-called neurovegetative symptoms) involve physiologic functions, a key role for the hypothalamus is also presumed. A subset of depressed individuals shows a small reduction in hippocampal size, as noted above. In addition, brain imaging investigations have revealed increased activation of the amygdala by negative stimuli and reduced activation of the nucleus accumbens by rewarding stimuli. There is also evidence for altered activity in prefrontal cortex, such as hyperactivity of subgenual area 25 in anterior cingulate cortex. Such findings

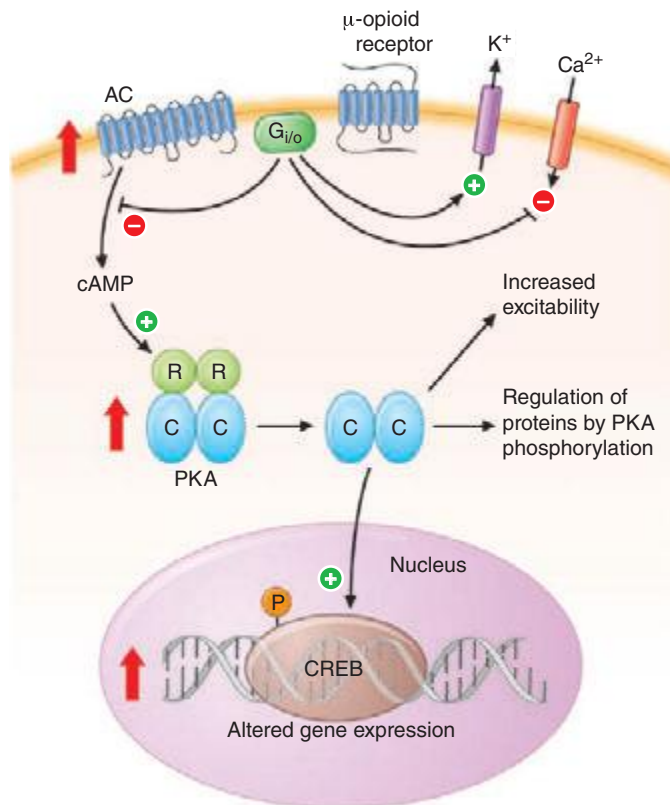


FIGURE 465e-1 Opiate action in the locus coeruleus (LC). Binding of opiate agonists to μ -opioid receptors catalyzes nucleotide exchange on G_i and G_o proteins, leading to inhibition of adenylyl cyclase, neuronal hyperpolarization via activation of K^+ channels, and inhibition of neurotransmitter release via inhibition of Ca^{2+} channels. Inhibition of adenylyl cyclase (AC) reduces protein kinase A (PKA) activity and phosphorylation of several PKA substrate proteins, thereby altering their function. For example, opiates reduce phosphorylation of the cAMP response element-binding protein (CREB), which appears to initiate longer term changes in neuronal function. Chronic administration of opiates increases levels of AC isoforms, PKA catalytic (C) and regulatory (R) subunits, and the phosphorylation of several proteins, including CREB (indicated by red arrows). These changes contribute to the altered phenotype of the drug-addicted state. For example, the excitability of LC neurons is increased by enhanced cAMP signaling, although the ionic basis of this effect remains unknown. Activation of CREB causes upregulation of AC isoforms and tyrosine hydroxylase, the rate-limiting enzyme in catecholamine biosynthesis.

have led to trials of deep brain stimulation (DBS) of either the nucleus accumbens or subgenual area 25, which appears to be therapeutic in some severely depressed individuals.

In schizophrenia, structural and functional imaging studies have identified a 3% loss of brain volume, most of which is in gray matter. This loss is progressive, and cortical gray matter appears to be particularly affected over time. The temporal lobes, particularly the left superior temporal gyrus, Heschl gyrus, and planum temporale, are often the most severely affected. The rate of loss in these regions as well as in frontal and parietal lobes appears to be greatest early in the course of the disease. Functional imaging studies provide evidence of reduced metabolic (presumably neural) activity in the dorsolateral prefrontal cortex at rest and when performing tests of executive function, including working memory. There is also evidence for impaired structural and task-related functional connectivity, mainly in frontal and temporal lobes.

These neuroimaging findings in schizophrenia have been confirmed in pathologic studies that show enlargement of the ventricular

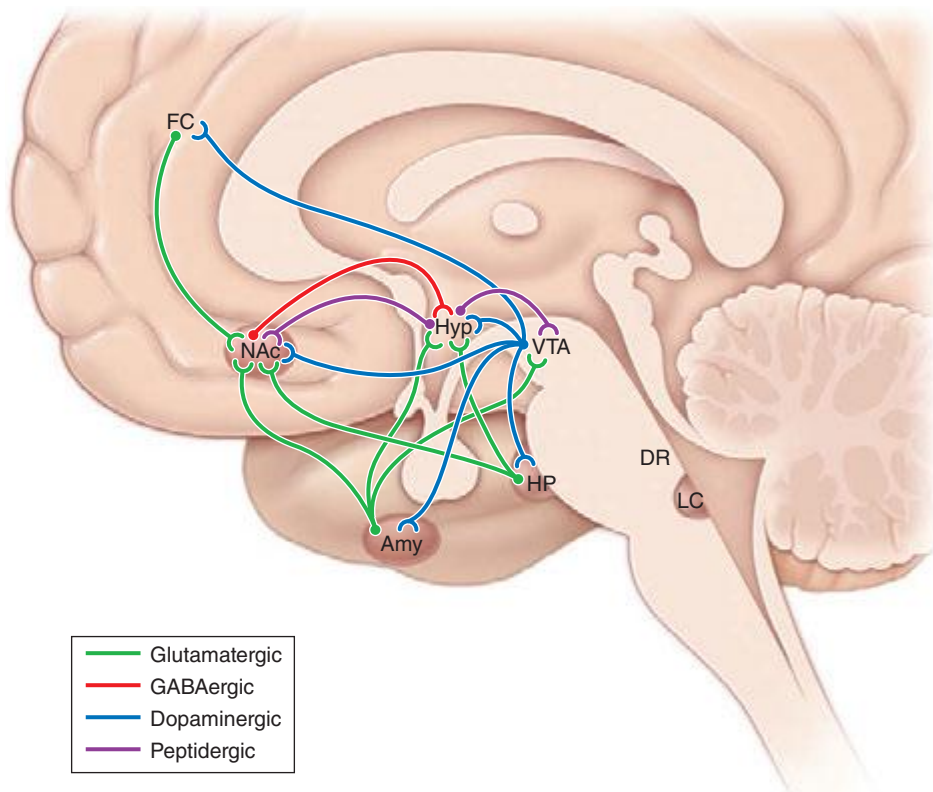


FIGURE 465e-2 Neural circuitry of depression and addiction. The figure shows a simplified summary of a series of limbic circuits in brain that regulate mood and motivation and are implicated in depression and addiction. Shown in the figure are the hippocampus (HP) and amygdala (Amy) in the temporal lobe, regions of prefrontal cortex, nucleus accumbens (NAc), and hypothalamus (Hyp). Only a subset of the known interconnections among these brain regions is shown. Also shown is the innervation of several of these brain regions by monoaminergic neurons. The ventral tegmental area (VTA) provides dopaminergic input to each of the limbic structures. Norepinephrine (from the locus coeruleus [LC]) and serotonin (from the dorsal raphe [DR] and other raphe nuclei) innervate all of the regions shown. In addition, there are strong connections between the hypothalamus and the VTA-NAc pathway. Important peptidergic projections from the hypothalamus include those from the arcuate nucleus that release β -endorphin and melanocortin and from the lateral hypothalamus that release orexin.

system and reduction of cortical and subcortical gray matter in frontal and temporal lobes and in the limbic system. The reduction in cortical thickness is associated with increased cell packing density and reduced neuropil (defined as axons, dendrites, and glial cell processes) without an apparent change in neuronal cell number. Specific classes of interneurons in prefrontal cortex consistently show reduced expression of the gene encoding the enzyme glutamic acid decarboxylase 1 (*GAD1*), which synthesizes γ -aminobutyric acid (GABA), the principal inhibitory neurotransmitter in the brain. Neuregulin 1 (*NRG1*), a member of the epidermal growth factor (EGF) family of growth factors, and its receptor *ERBB4*, have been implicated in schizophrenia, and they serve important roles in the maturation of GABAergic interneurons in cerebral cortex; loss of *NRG1-ERBB4* in mice leads to a reduced neuropil, thus phenocopying a pathologic finding in schizophrenia. These findings are consistent with one working hypothesis of schizophrenia as a developmental neurodegenerative disorder due in part to loss of cortical interneurons in frontal and temporal lobes.

Work in rodent and nonhuman primate models of addiction has established the brain's reward regions as key neural substrates for the acute actions of drugs of abuse and for addiction induced by repeated drug administration (Fig. 465e-2). Midbrain dopamine neurons in the VTA function normally as rheostats of reward: they are activated by natural rewards (food, sex, social interaction) or even by the expectation of such rewards, and many are suppressed by the absence of an expected reward or by aversive stimuli. These neurons thereby transmit crucial survival signals to the rest of the limbic brain to promote reward-related behavior, including motor responses to seek and obtain

the rewards (nucleus accumbens), memories of reward-related cues (amygdala, hippocampus), and executive control of obtaining rewards (prefrontal cortex).

Drugs of abuse alter neurotransmission through initial actions at different classes of ion channels, neurotransmitter receptors, or neurotransmitter transporters (Table 465e-2). Studies in animal models have demonstrated that although the initial targets differ, the actions of these drugs converge on the brain's reward circuitry by promoting dopamine neurotransmission in the nucleus accumbens and other limbic targets of the VTA. In addition, some drugs promote activation of opioid and cannabinoid receptors, which modulate this reward circuitry. By these mechanisms, drugs of abuse produce powerful rewarding signals, which, after repeated drug administration, corrupt a vulnerable brain's reward circuitry in ways that promote addiction. Three major pathologic adaptations have been described. First, drugs produce tolerance and dependence in reward circuits, which promote escalating drug intake and a negative emotional state during drug withdrawal that promotes relapse. Second, sensitization to the rewarding effects of the drugs and associated cues is seen during prolonged abstinence and also triggers relapse. Third, executive function is impaired in such a way as to increase impulsivity and compulsivity, both of which promote relapse.

Imaging studies in humans confirm that addictive drugs, as well as craving for them, activate the brain's reward circuitry. In addition, patients who abuse alcohol or psychostimulants show reduced gray matter in the prefrontal cortex as well as reduced activity in anterior cingulate and

orbitofrontal cortex during tasks of attention and inhibitory control. It is thought that damage to these cortical areas contributes to addiction by impairing decision-making and increasing impulsivity.

NEUROINFLAMMATION

There is increasing evidence for the involvement of inflammatory mechanisms in a subset of depressed patients. These individuals display elevated blood levels of interleukin 6 (IL-6), tumor necrosis factor α (TNF- α), and other cytokines. Moreover, rodents exposed to chronic stress exhibit similar increases in peripheral cytokines, and peripheral or central delivery of those cytokines to normal rodents increases their susceptibility to chronic stress. These findings have led to the novel idea of using peripheral cytokines as biomarkers of a subtype of depression and the potential utility of developing new antidepressants that oppose cytokine action.

Recent evidence has also linked proinflammatory signaling in the brain to addiction, particularly to alcohol. Human alcoholism is associated with impaired innate immunity, increases in circulating proinflammatory cytokines, and an increase in brain levels of the cytokine monocyte chemoattractant protein-1 (MCP-1, also referred to as CCL2). Many of these cytokines are produced by astrocytes and microglia, and by neurons under certain pathologic conditions, where they play important roles in modifying neuronal function and plasticity. For example MCP-1 modulates the release of certain neurotransmitters and, when administered into the VTA, increases neuronal excitability, promotes dopamine release, and increases locomotor activity. Recent gene expression array studies of alcohol drinking in mice have

TABLE 465e-2 INITIAL ACTIONS OF DRUGS OF ABUSE

Drug	Neurotransmitter Affected	Drug Target (Action)
Opiates	Endorphins, enkephalins	μ - and δ -opioid receptors (agonist)
Psychostimulants (cocaine, amphetamine, methamphetamine)	Dopamine	Dopamine transporter (antagonist—cocaine; reverse transport—amphetamine, methamphetamine)
Nicotine	Acetylcholine	Nicotinic cholinergic receptors (agonist)
Ethanol	GABA	GABA _A receptors (positive allosteric modulator)
	Glutamate	NMDA glutamate receptors (antagonist)
	Acetylcholine	Nicotinic cholinergic receptors (allosteric modulator)
	Serotonin	5HT-3 receptor (positive allosteric modulator)
	—	Calcium-activated K ⁺ channel (activator)
Marijuana	Endocannabinoids (anandamide, 2-arachidonoylglycerol)	CB ₁ receptor (agonist)
Phencyclidine	Glutamate	NMDA glutamate receptor (antagonist)

identified a network of regulated cytokines in brain, and a role in regulation of alcohol consumption has been recently validated for several of them, including IL-6. A major focus of current research is to define the site and mechanism by which proinflammatory cytokines impair brain function to elicit a depressive episode or promote drug abuse.

CONCLUSIONS

This brief narrative illustrates the substantial progress that is being made in understanding the genetic and neurobiologic basis of mental illness. It is anticipated that biologic measures will be used increasingly to diagnosis and subtype a psychiatric disorder and that targeted therapeutics will become available to treat them.

This page intentionally left blank

466 Mental Disorders

Victor I. Reus

Mental disorders are common in medical practice and may present either as a primary disorder or as a comorbid condition. The prevalence of mental or substance use disorders in the United States is approximately 30%, but only one-third of affected individuals are currently receiving treatment. Global burden of disease statistics indicate that 4 of the 10 most important causes of morbidity and attendant health care costs worldwide are psychiatric in origin.

Changes in health care delivery underscore the need for primary care physicians to assume responsibility for the initial diagnosis and treatment of the most common mental disorders. Prompt diagnosis is essential to ensure that patients have access to appropriate medical services and to maximize the clinical outcome. Validated patient-based questionnaires have been developed that systematically probe for signs and symptoms associated with the most prevalent psychiatric diagnoses and guide the clinician into targeted assessment. The Primary Care Evaluation of Mental Disorders (PRIME-MD; and a self-report form, the Patient Health Questionnaire) and the Symptom-Driven Diagnostic System for Primary Care (SDDS-PC) are inventories that require only 10 min to complete and link patient responses to the formal diagnostic criteria of anxiety, mood, somatoform, and eating disorders and to alcohol abuse or dependence.

A physician who refers patients to a psychiatrist should know not only when doing so is appropriate but also how to refer, because societal misconceptions and the stigma of mental illness impede the process. Primary care physicians should base referrals to a psychiatrist on the presence of signs and symptoms of a mental disorder and not simply on the absence of a physical explanation for a patient's complaint. The physician should discuss with the patient the reasons for requesting the referral or consultation and provide reassurance that he or she will continue to provide medical care and work collaboratively with the mental health professional. Consultation with a psychiatrist or transfer of care is appropriate when physicians encounter evidence of psychotic symptoms, mania, severe depression, or anxiety; symptoms of posttraumatic stress disorder

(PTSD); suicidal or homicidal preoccupation; or a failure to respond to first-order treatment. This chapter reviews the clinical assessment and treatment of some of the most common mental disorders presenting in primary care and is based on the *Diagnostic and Statistical Manual of Mental Disorders*, Fifth Edition (DSM-5), the framework for categorizing psychiatric illness used in the United States. **Eating disorders are discussed later in this chapter, and the biology of psychiatric and addictive disorders is discussed in Chap. 465e.**

GLOBAL CONSIDERATIONS



The DSM-5 and the tenth revision of the International Classification of Diseases (ICD-10), which is used more commonly worldwide, have taken somewhat differing approaches to the diagnosis of mental illness, but considerable effort has been expended to provide an operational translation between the two nosologies. Both systems are in essence purely descriptive and emphasize clinical pragmatism, in distinction to the Research Domain Criteria (RDOC) proposed by National Institute of Mental Health, which aspires to provide a causal framework for classification of behavioral disturbance. None of these diagnostic systems has as yet achieved adequate validation. The Global Burden of Disease Study 2010, using available epidemiologic data, nevertheless has reinforced the conclusion that, regardless of nosologic differences, mental and substance abuse disorders are the major cause of life-years lost to disability among all medical illnesses. There is general agreement that high-income countries will need to build capacity in professional training in low- and middle-income countries in order to provide an adequate balanced care model for the delivery of evidence-based therapies for mental disorders. Recent surveys that indicate a dramatic increase in mental disorder prevalence in rapidly developing countries, such as China, may reflect both an increased recognition of the issue, but also the consequence of social turmoil, stigma, and historically inadequate resources. The need for improved prevention strategies and for more definitive and effective interventional treatments remains a global concern.

ANXIETY DISORDERS

Anxiety disorders, the most prevalent psychiatric illnesses in the general community, are present in 15–20% of medical clinic patients. Anxiety, defined as a subjective sense of unease, dread, or foreboding, can indicate a primary psychiatric condition or can be a component of, or reaction to, a primary medical disease. The primary anxiety disorders are classified according to their duration and course and the existence and nature of precipitants.

When evaluating the anxious patient, the clinician must first determine whether the anxiety antedates or postdates a medical illness or is due to a medication side effect. Approximately one-third of patients presenting with anxiety have a medical etiology for their psychiatric symptoms, but an anxiety disorder can also present with somatic symptoms in the absence of a diagnosable medical condition.

PANIC DISORDER

Clinical Manifestations Panic disorder is defined by the presence of recurrent and unpredictable panic attacks, which are distinct episodes of intense fear and discomfort associated with a variety of physical symptoms, including palpitations, sweating, trembling, shortness of breath, chest pain, dizziness, and a fear of impending doom or death. Paresthesias, gastrointestinal distress, and feelings of unreality are also common. Diagnostic criteria require at least 1 month of concern or worry about the attacks or a change in behavior related to them. The lifetime prevalence of panic disorder is 2–3%. Panic attacks have a sudden onset, developing within 10 min and usually resolving over the course of an hour, and they occur in an unexpected fashion. Some may occur when waking from sleep. The frequency and severity of panic

attacks vary, ranging from once a week to clusters of attacks separated by months of well-being. The first attack is usually outside the home, and onset is typically in late adolescence to early adulthood. In some individuals, anticipatory anxiety develops over time and results in a generalized fear and a progressive avoidance of places or situations in which a panic attack might recur. *Agoraphobia*, which occurs commonly in patients with panic disorder, is an acquired irrational fear of being in places where one might feel trapped or unable to escape. It may, however, be diagnosed even if panic disorder is not present. Typically, it leads the patient into a progressive restriction in lifestyle and, in a literal sense, in geography. Frequently, patients are embarrassed that they are housebound and dependent on the company of others to go out into the world and do not volunteer this information; thus, physicians will fail to recognize the syndrome if direct questioning is not pursued.

Differential Diagnosis A diagnosis of panic disorder is made after a medical etiology for the panic attacks has been ruled out. A variety of cardiovascular, respiratory, endocrine, and neurologic conditions can present with anxiety as the chief complaint. Patients with true panic disorder will often focus on one specific feature to the exclusion of others. For example, 20% of patients who present with syncope as a primary medical complaint have a primary diagnosis of a mood, anxiety, or substance abuse disorder, the most common being panic disorder. The differential diagnosis of panic disorder is complicated by a high rate of comorbidity with other psychiatric conditions, especially alcohol and benzodiazepine abuse, which patients initially use in an attempt at self-medication. Some 75% of panic disorder patients will also satisfy criteria for major depression at some point in their illness.

When the history is nonspecific, physical examination and focused laboratory testing must be used to rule out anxiety states resulting from medical disorders such as pheochromocytoma, thyrotoxicosis, or hypoglycemia. Electrocardiogram (ECG) and echocardiogram may detect some cardiovascular conditions associated with panic such as paroxysmal atrial tachycardia and mitral valve prolapse. In two studies, panic disorder was the primary diagnosis in 43% of patients with chest pain who had normal coronary angiograms and was present in 9% of all outpatients referred for cardiac evaluation. Panic disorder has also been diagnosed in many patients referred for pulmonary function testing or with symptoms of irritable bowel syndrome.

Etiology and Pathophysiology The etiology of panic disorder is unknown but appears to involve a genetic predisposition, altered autonomic responsiveness, and social learning. Panic disorder shows familial aggregation; the disorder is concordant in 30–45% of monozygotic twins, and genome-wide screens have identified suggestive risk loci. Acute panic attacks appear to be associated with increased noradrenergic discharges in the locus coeruleus. Intravenous infusion of sodium lactate evokes an attack in two-thirds of panic disorder patients, as do the α_2 -adrenergic antagonist yohimbine, cholecystokinin tetrapeptide (CCK-4), and carbon dioxide inhalation. It is hypothesized that each of these stimuli activates a pathway involving noradrenergic neurons in the locus coeruleus and serotonergic neurons in the dorsal raphe. Agents that block serotonin reuptake can prevent attacks. Patients with panic disorder have a heightened sensitivity to somatic symptoms, which triggers increasing arousal, setting off the panic attack; accordingly, therapeutic intervention involves altering the patient's cognitive interpretation of anxiety-producing experiences as well as preventing the attack itself.

TREATMENT PANIC DISORDER

Achievable goals of treatment are to decrease the frequency of panic attacks and to reduce their intensity. The cornerstone of drug therapy is antidepressant medication (Tables 466-1 through 466-3). Selective serotonin reuptake inhibitors (SSRIs) benefit the majority of panic disorder patients and do not have the adverse effects of tricyclic antidepressants (TCAs). Fluoxetine, paroxetine, sertraline, and the selective serotonin-norepinephrine reuptake inhibitor (SNRI) venlafaxine have received approval from the U.S. Food and Drug Administration (FDA) for this indication. These drugs should be started at one-third to one-half of their usual antidepressant dose (e.g., 5–10 mg

fluoxetine, 25–50 mg sertraline, 10 mg paroxetine, venlafaxine 37.5 mg). Monoamine oxidase inhibitors (MAOIs) are also effective and may specifically benefit patients who have comorbid features of atypical depression (i.e., hypersomnia and weight gain). Insomnia, orthostatic hypotension, and the need to maintain a low-tyramine diet (avoidance of cheese and wine) have limited their use, however. Antidepressants typically take 2–6 weeks to become effective, and doses may need to be adjusted based on the clinical response.

Because of anticipatory anxiety and the need for immediate relief of panic symptoms, benzodiazepines are useful early in the course of treatment and sporadically thereafter (Table 466-4). For example, alprazolam, starting at 0.5 mg qid and increasing to 4 mg/d in divided doses, is effective, but patients must be monitored closely, as some develop dependence and begin to escalate the dose of this medication. Clonazepam, at a final maintenance dose of 2–4 mg/d, is also helpful; its longer half-life permits twice-daily dosing, and patients appear less likely to develop dependence on this agent.

Early psychotherapeutic intervention and education aimed at symptom control enhance the effectiveness of drug treatment. Patients can be taught breathing techniques, be educated about physiologic changes that occur with panic, and learn to expose themselves voluntarily to precipitating events in a treatment program spanning 12–15 sessions. Homework assignments and monitored compliance are important components of successful treatment. Once patients have achieved a satisfactory response, drug treatment should be maintained for 1–2 years to prevent relapse. Controlled trials indicate a success rate of 75–85%, although the likelihood of complete remission is somewhat lower.

GENERALIZED ANXIETY DISORDER

Clinical Manifestations Patients with generalized anxiety disorder (GAD) have persistent, excessive, and/or unrealistic worry associated with muscle tension, impaired concentration, autonomic arousal, feeling “on edge” or restless, and insomnia (Table 466-5). Onset is usually before age 20 years, and a history of childhood fears and social inhibition may be present. The lifetime prevalence of GAD is 5–6%; the risk is higher in first-degree relatives of patients with the diagnosis. Interestingly, family studies indicate that GAD and panic disorder segregate independently. More than 80% of patients with GAD also suffer from major depression, dysthymia, or social phobia. Comorbid substance abuse is common in these patients, particularly alcohol and/or sedative/hypnotic abuse. Patients with GAD worry excessively over minor matters, with life-disrupting effects; unlike in panic disorder, complaints of shortness of breath, palpitations, and tachycardia are relatively rare.

Etiology and Pathophysiology All anxiogenic agents act on the γ -aminobutyric acid (GABA)_A receptor/chloride ion channel complex, implicating this neurotransmitter system in the pathogenesis of anxiety and panic attacks. Benzodiazepines are thought to bind two separate GABA_A receptor sites: type I, which has a broad neuroanatomic distribution, and type II, which is concentrated in the hippocampus, striatum, and neocortex. The antianxiety effects of the various benzodiazepines are influenced by their relative binding to alpha 2 and 3 subunits of the GABA_A receptor, and sedation and memory impairment to the alpha 1 subunit. Serotonin (5-hydroxytryptamine [5-HT]) and 3 α -reduced neuroactive steroids (allosteric modulators of GABA_A) also appear to have a role in anxiety, and buspirone, a partial 5-HT_{1A} receptor agonist, and certain 5-HT_{2A} and 5-HT_{2C} receptor antagonists (e.g., nefazodone) may have beneficial effects.

TREATMENT GENERALIZED ANXIETY DISORDER

A combination of pharmacologic and psychotherapeutic interventions is most effective in GAD, but complete symptomatic relief is rare. A short course of a benzodiazepine is usually indicated, preferably lorazepam, oxazepam, or alprazolam. (The first two of these agents are metabolized via conjugation rather than oxidation and thus do not accumulate if hepatic function is impaired; the latter

Name	Usual Daily Dose, mg	Side Effects	Comments	
SSRIs				
Fluoxetine (Prozac)	10–80	Headache; nausea and other GI effects; jitteriness; insomnia; sexual dysfunction; can affect plasma levels of other medicines (except sertraline); akathisia rare	Once-daily dosing, usually in the morning; fluoxetine has very long half-life; must not be combined with MAOIs	
Sertraline (Zoloft)	50–200			
Paroxetine (Paxil)	20–60			
Fluvoxamine (Luvox)	100–300			
Citalopram (Celexa)	20–60			
Escitalopram (Lexapro)	10–30			
TCAs				
Amitriptyline (Elavil)	150–300	Anticholinergic (dry mouth, tachycardia, constipation, urinary retention, blurred vision); sweating; tremor; postural hypotension; cardiac conduction delay; sedation; weight gain	Once-daily dosing, usually qhs; blood levels of most TCAs available; can be lethal in overdose (lethal dose = 2 g); nortriptyline best tolerated, especially by elderly	
Nortriptyline (Pamelor)	50–200			
Imipramine (Tofranil)	150–300			
Desipramine (Norpramin)	150–300			
Doxepin (Sinequan)	150–300			
Clomipramine (Anafranil)	150–300			FDA approved for OCD
Mixed Norepinephrine/Serotonin Reuptake Inhibitors (SNRI) and Receptor Blockers				
Venlafaxine (Effexor)	75–375	Nausea; dizziness; dry mouth; headaches; increased blood pressure; anxiety and insomnia	Bid-tid dosing (extended release available); lower potential for drug interactions than SSRIs; contraindicated with MAOIs	
Desvenlafaxine (Pristiq)	50–400	Nausea, dizziness, insomnia	Primary metabolite of venlafaxine; no increased efficacy with higher dosing	
Duloxetine (Cymbalta)	40–60	Nausea, dizziness, headache, insomnia, constipation	May have utility in treatment of neuropathic pain and stress incontinence	
Mirtazapine (Remeron)	15–45	Somnolence, weight gain; neutropenia rare	Once a day dosing	
Vilazodone (Viibryd)	40	Nausea, diarrhea, headache; dosage adjustment if given with CYP3A4 inhibitor/stimulator	Also 5-HT _{1a} receptor partial agonist	
Vortioxetine (Brintellix)	5–20	Nausea, diarrhea, sweating, headache; low incidence of sedation or weight gain	No specific p450 effects; 5-HT _{3a} and 5-HT ₇ receptor antagonist, 5-HT _{1b} partial agonist, and 5-HT _{1a} agonist	
Levomilnacipran (Fetzima)	40–120	Nausea, constipation, sweating; rare increase in blood pressure/pulse	Most noradrenergic of SNRIs	
Mixed-Action Drugs				
Bupropion (Wellbutrin)	250–450	Jitteriness; flushing; seizures in at-risk patients; anorexia; tachycardia; psychosis	Tid dosing, but sustained release also available; fewer sexual side effects than SSRIs or TCAs; may be useful for adult ADD	
Trazodone (Desyrel)	200–600	Sedation; dry mouth; ventricular irritability; postural hypotension; priapism rare	Useful in low doses for sleep because of sedating effects with no anticholinergic side effects	
Trazodone extended release (Oleptro)	150–375	Daytime somnolence, dizziness, nausea		
Amoxapine (Asendin)	200–600	Sexual dysfunction	Lethality in overdose; EPS possible	
MAOIs				
Phenelzine (Nardil)	45–90	Insomnia; hypotension; edema; anorgasmia; weight gain; neuropathy; hypertensive crisis; toxic reactions with SSRIs; narcotics	May be more effective in patients with atypical features or treatment-refractory depression	
Tranylcypromine (Parnate)	20–50			
Isocarboxazid (Marplan)	20–60			Less weight gain and hypotension than phenelzine
Transdermal selegiline (Emsam)	6–12			Local skin reaction hypertension

Abbreviations: ADD, attention deficit disorder; EPS, extrapyramidal symptoms; FDA, U.S. Food and Drug Administration; GI, gastrointestinal; MAOIs, monoamine oxidase inhibitors; OCD, obsessive-compulsive disorder; SSRIs, selective serotonin reuptake inhibitors; TCAs, tricyclic antidepressants.

also has limited active metabolites.) Treatment should be initiated at the lowest dose possible and prescribed on an as-needed basis as symptoms warrant. Benzodiazepines differ in their milligram per kilogram potency, half-life, lipid solubility, metabolic pathways, and presence of active metabolites. Agents that are absorbed rapidly and are lipid soluble, such as diazepam, have a rapid onset of action and a higher abuse potential. Benzodiazepines should generally not be prescribed for >4–6 weeks because of the development of tolerance and the risk of abuse and dependence. Withdrawal must be closely monitored as relapses can occur. It is important to warn patients that concomitant use of alcohol or other sedating drugs may exacerbate side effects and impair their ability to function. An optimistic approach that encourages the patient to clarify environmental precipitants, anticipate his or her reactions, and plan effective response strategies is an essential element of therapy.

Adverse effects of benzodiazepines generally parallel their relative half-lives. Longer-acting agents, such as diazepam, chlordiazepoxide, flurazepam, and clonazepam, tend to accumulate active metabolites, with resultant sedation, impairment of cognition, and poor psychomotor performance. Shorter-acting compounds, such as alprazolam, lorazepam, and oxazepam, can produce daytime anxiety, early morning insomnia, and, with discontinuation, rebound anxiety and insomnia. Although patients develop tolerance to the sedative effects of benzodiazepines, they are less likely to habituate to the adverse psychomotor effects. Withdrawal from the longer half-life benzodiazepines can be accomplished through gradual, stepwise dose reduction (by 10% every 1–2 weeks) over 6–12 weeks. It is usually more difficult to taper patients off shorter-acting benzodiazepines. Physicians may need to switch the patient to a benzodiazepine with a longer half-life or use an

TABLE 466-2 MANAGEMENT OF ANTIDEPRESSANT SIDE EFFECTS

Symptoms	Comments and Management Strategies
Gastrointestinal	
Nausea, loss of appetite	Usually short-lived and dose-related; consider temporary dose reduction or administration with food and antacids
Diarrhea	Famotidine, 20–40 mg/d
Constipation	Wait for tolerance; try diet change, stool softener, exercise; avoid laxatives
Sexual dysfunction	Consider dose reduction; drug holiday
Anorgasmia/impotence; impaired ejaculation	Bethanechol, 10–20 mg, 2 h before activity, or cyproheptadine, 4–8 mg 2 h before activity, or bupropion, 100 mg bid, or amantadine, 100 mg bid/tid
Orthostasis	Tolerance unlikely; increase fluid intake, use calf exercises/support hose; fludrocortisone, 0.025 mg/d
Anticholinergic	Wait for tolerance
Dry mouth, eyes	Maintain good oral hygiene; use artificial tears, sugar-free gum
Tremor/jitteriness	Antiparkinsonian drugs not effective; use dose reduction/slow increase; lorazepam, 0.5 mg bid, or propranolol, 10–20 mg bid
Insomnia	Schedule all doses for the morning; trazodone, 50–100 mg qhs
Sedation	Caffeine; schedule all dosing for bedtime; bupropion, 75–100 mg in afternoon
Headache	Evaluate diet, stress, other drugs; try dose reduction; amitriptyline, 50 mg/d
Weight gain	Decrease carbohydrates; exercise; consider fluoxetine
Loss of therapeutic benefit over time	Related to tolerance? Increase dose or drug holiday; add amantadine, 100 mg bid, buspirone, 10 mg tid, or pindolol, 2.5 mg bid

adjunctive medication such as a beta blocker or carbamazepine, before attempting to discontinue the benzodiazepine. Withdrawal reactions vary in severity and duration; they can include depression, anxiety, lethargy, diaphoresis, autonomic arousal, and, rarely, seizures.

TABLE 466-3 POSSIBLE DRUG INTERACTIONS WITH SELECTIVE SEROTONIN REUPTAKE INHIBITORS

Agent	Effect
Monoamine oxidase inhibitors	Serotonin syndrome—absolute contraindication
Serotonergic agonists, e.g., tryptophan, fenfluramine, tryptans	Potential serotonin syndrome
Drugs that are metabolized by P450 isoenzymes: tricyclics, other SSRIs, antipsychotics, beta blockers, codeine, triazolobenzodiazepines, calcium channel blockers	Delayed metabolism resulting in increased blood levels and potential toxicity
Drugs that are bound tightly to plasma proteins, e.g., warfarin	Increased bleeding secondary to displacement
Drugs that inhibit the metabolism of SSRIs by P450 isoenzymes, e.g., quinidine	Increased SSRI side effects

Abbreviation: SSRIs, selective serotonin reuptake inhibitors.

Buspirone is a nonbenzodiazepine anxiolytic agent. It is non-sedating, does not produce tolerance or dependence, does not interact with benzodiazepine receptors or alcohol, and has no abuse or disinhibition potential. However, it requires several weeks to take effect and requires thrice-daily dosing. Patients who were previously responsive to a benzodiazepine are unlikely to rate buspirone as equally effective, but patients with head injury or dementia who have symptoms of anxiety and/or agitation may do well with this agent. Escitalopram, paroxetine, and venlafaxine are FDA approved for the treatment of GAD, usually at doses that are comparable to their efficacy in major depression, and may be preferable to usage of benzodiazepines in the treatment of chronic anxiety. Benzodiazepines are contraindicated during pregnancy and breast-feeding.

Anticonvulsants with GABAergic properties may also be effective against anxiety. Gabapentin, oxcarbazepine, tiagabine, pregabalin, and divalproex have all shown some degree of benefit in a variety of anxiety-related syndromes in off-label usage. Agents that selectively target GABA_A receptor subtypes are currently under development, and it is hoped that these will lack the sedating, memory-impairing, and addicting properties of benzodiazepines.

TABLE 466-4 ANXIOLYTICS

Name	Equivalent PO Dose, mg	Onset of Action	Half-Life, h	Comments
Benzodiazepines				
Diazepam (Valium)	5	Fast	20–70	Active metabolites; quite sedating
Flurazepam (Dalmane)	15	Fast	30–100	Flurazepam is a prodrug; metabolites are active; quite sedating
Triazolam (Halcion)	0.25	Intermediate	1.5–5	No active metabolites; can induce confusion and delirium, especially in elderly
Lorazepam (Ativan)	1	Intermediate	10–20	No active metabolites; direct hepatic glucuronide conjugation; quite sedating; FDA approved for anxiety with depression
Alprazolam (Xanax)	0.5	Intermediate	12–15	Active metabolites; not too sedating; FDA approved for panic disorder and anxiety with depression; tolerance and dependence develop easily; difficult to withdraw
Chlordiazepoxide (Librium)	10	Intermediate	5–30	Active metabolites; moderately sedating
Oxazepam (Serax)	15	Slow	5–15	No active metabolites; direct glucuronide conjugation; not too sedating
Temazepam (Restoril)	15	Slow	9–12	No active metabolites; moderately sedating
Clonazepam (Klonopin)	0.5	Slow	18–50	No active metabolites; moderately sedating; FDA approved for panic disorder
Clorazepate (Tranxene)	15	Fast	40–200	Low sedation; unreliable absorption
Nonbenzodiazepines				
Buspirone (BuSpar)	7.5	2 weeks	2–3	Active metabolites; tid dosing—usual daily dose 10–20 mg tid; nonsedating; no additive effects with alcohol; useful for controlling agitation in demented or brain-injured patients

Abbreviation: FDA, U.S. Food and Drug Administration.

TABLE 466-5 DIAGNOSTIC CRITERIA FOR GENERALIZED ANXIETY DISORDER

- A. Excessive anxiety and worry (apprehensive expectation), occurring more days than not for at least 6 months, about a number of events or activities (such as work or school performance).
- B. The individual finds it difficult to control the worry.
- C. The anxiety and worry are associated with three (or more) of the following six symptoms (with at least some symptoms present for more days than not for the past 6 months): (1) restlessness or feeling keyed up or on edge; (2) being easily fatigued; (3) difficulty concentrating or mind going blank; (4) irritability; (5) muscle tension; (6) sleep disturbance (difficulty falling or staying asleep, or restless, unsatisfying sleep).
- D. The anxiety, worry, or physical symptoms cause clinically significant distress or impairment in social, occupational, or other important areas of functioning.
- E. The disturbance is not attributable to the physiologic effects of a substance (e.g., a drug of abuse, a medication) or another medical condition (e.g., hyperthyroidism).
- F. The disturbance is not better explained by another mental disorder (e.g., anxiety or worry about having panic attacks in panic disorder, negative evaluation in social anxiety disorder [social phobia], contamination or other obsessions in obsessive-compulsive disorder, separation from attachment figures in separation anxiety disorder, reminders of traumatic events in posttraumatic stress disorder, gaining weight in anorexia nervosa, physical complaints in somatic symptom disorder, perceived appearance flaws in body dysmorphic disorder, having a serious illness in illness anxiety disorder, or the content of delusional beliefs in schizophrenia or delusional disorder).

Source: *Diagnostic and Statistical Manual of Mental Disorders*, 5th ed. Washington, DC, American Psychiatric Association, 2013.

PHOBIC DISORDERS

Clinical Manifestations The cardinal feature of phobic disorders is a marked and persistent fear of objects or situations, exposure to which results in an immediate anxiety reaction. The patient avoids the phobic stimulus, and this avoidance usually impairs occupational or social functioning. Panic attacks may be triggered by the phobic stimulus or may occur spontaneously. Unlike patients with other anxiety disorders, individuals with phobias usually experience anxiety only in specific situations. Common phobias include fear of closed spaces (claustrophobia), fear of blood, and fear of flying. Social phobia is distinguished by a specific fear of social or performance situations in which the individual is exposed to unfamiliar individuals or to possible examination and evaluation by others. Examples include having to converse at a party, use public restrooms, and meet strangers. In each case, the affected individual is aware that the experienced fear is excessive and unreasonable given the circumstance. The specific content of a phobia may vary across gender, ethnic, and cultural boundaries.

Phobic disorders are common, affecting ~7–9% of the population. Twice as many females are affected than males. Full criteria for diagnosis are usually satisfied first in early adulthood, but behavioral avoidance of unfamiliar people, situations, or objects dating from early childhood is common.

In one study of female twins, concordance rates for agoraphobia, social phobia, and animal phobia were found to be 23% for monozygotic twins and 15% for dizygotic twins. A twin study of fear conditioning, a model for the acquisition of phobias, demonstrated a heritability of 35–45%. Animal studies of fear conditioning have indicated that processing of the fear stimulus occurs through the lateral nucleus of the amygdala, extending through the central nucleus and projecting to the periaqueductal gray region, lateral hypothalamus, and paraventricular hypothalamus.

TREATMENT PHOBIC DISORDERS

Beta blockers (e.g., propranolol, 20–40 mg orally 2 h before the event) are particularly effective in the treatment of “performance anxiety” (but not general social phobia) and appear to work by blocking the peripheral manifestations of anxiety such as perspiration, tachycardia, palpitations, and tremor. MAOIs alleviate

social phobia independently of their antidepressant activity, and paroxetine, sertraline, and venlafaxine have received FDA approval for treatment of social anxiety. Benzodiazepines can be helpful in reducing fearful avoidance, but the chronic nature of phobic disorders limits their usefulness.

Behaviorally focused psychotherapy is an important component of treatment because relapse rates are high when medication is used as the sole treatment. Cognitive-behavioral strategies are based on the finding that distorted perceptions and interpretations of fear-producing stimuli play a major role in perpetuation of phobias. Individual and group therapy sessions teach the patient to identify specific negative thoughts associated with the anxiety-producing situation and help to reduce the patient’s fear of loss of control. In desensitization therapy, hierarchies of feared situations are constructed, and the patient is encouraged to pursue and master gradual exposure to the anxiety-producing stimuli.

Patients with social phobia, in particular, have a high rate of comorbid alcohol abuse, as well as of other psychiatric conditions (e.g., eating disorders), necessitating the need for parallel management of each disorder if anxiety reduction is to be achieved.

STRESS DISORDERS

Clinical Manifestations Patients may develop anxiety after exposure to extreme traumatic events such as the threat of personal death or injury or the death of a loved one. The reaction may occur shortly after the trauma (*acute stress disorder*) or be delayed and subject to recurrence (PTSD) (Table 466-6). In both syndromes, individuals experience associated symptoms of detachment and loss of emotional responsiveness. The patient may feel depersonalized and unable to recall specific aspects of the trauma, although typically it is reexperienced through intrusions in thought, dreams, or flashbacks, particularly when cues of the original event are present. Patients often actively avoid stimuli that precipitate recollections of the trauma and demonstrate a resulting increase in vigilance, arousal, and startle response. Patients with stress disorders are at risk for the development of other disorders related to anxiety, mood, and substance abuse (especially alcohol). Between 5 and 10% of Americans will at some time in their life satisfy criteria for PTSD, with women more likely to be affected than men.

Risk factors for the development of PTSD include a past psychiatric history and personality characteristics of high neuroticism and extroversion. Twin studies show a substantial genetic influence on all symptoms associated with PTSD, with less evidence for an environmental effect.

Etiology and Pathophysiology It is hypothesized that in PTSD there is excessive release of norepinephrine from the locus coeruleus in response to stress and increased noradrenergic activity at projection sites in the hippocampus and amygdala. These changes theoretically facilitate the encoding of fear-based memories. Greater sympathetic responses to cues associated with the traumatic event occur in PTSD, although pituitary adrenal responses are blunted.

TREATMENT STRESS DISORDERS

Acute stress reactions are usually self-limited, and treatment typically involves the short-term use of benzodiazepines and supportive/expressive psychotherapy. The chronic and recurrent nature of PTSD, however, requires a more complex approach using drug and behavioral treatments. PTSD is highly correlated with peritraumatic dissociative symptoms and the development of an acute stress disorder at the time of the trauma. The SSRIs (paroxetine and sertraline are FDA approved for PTSD), venlafaxine, and topiramate can all reduce anxiety, symptoms of intrusion, and avoidance behaviors, as can prazosin, an α_1 antagonist. Propranolol and opiates such as morphine, given during the acute stress period, may have beneficial effects in preventing the development of PTSD, and adjunctive naltrexone can be effective when comorbid alcoholism is present. Trazodone, a sedating antidepressant, is frequently used at night

TABLE 466-6 DIAGNOSTIC CRITERIA FOR POSTTRAUMATIC STRESS DISORDER

- A. Exposure to actual or threatened death, serious injury, or sexual violence in one (or more) of the following ways:
1. Directly experiencing the traumatic event(s).
 2. Witnessing, in person, the event(s) as it occurred to others.
 3. Learning that the traumatic event(s) occurred to a close family member or close friend. In cases of actual or threatened death of a family member or friend, the event(s) must have been violent or accidental.
 4. Experiencing repeated or extreme exposure to aversive details of the traumatic event(s) (e.g., first responders collecting human remains; police officers repeatedly exposed to details of child abuse).
- B. Presence of one (or more) of the following intrusion symptoms associated with the traumatic event(s), beginning after the traumatic event(s) occurred:
1. Recurrent, involuntary, and intrusive distressing memories of the traumatic event(s).
 2. Recurrent distressing dreams in which the content and/or affect of the dream are related to the traumatic event(s).
 3. Dissociative reactions (e.g., flashbacks) in which the individual feels or acts as if the traumatic event(s) were recurring. (Such reactions may occur on a continuum, with the most extreme expression being a complete loss of awareness of present surroundings.)
 4. Intense or prolonged psychological distress at exposure to internal or external cues that symbolize or resemble an aspect of the traumatic event(s).
 5. Marked physiologic reactions to internal or external cues that symbolize or resemble an aspect of the traumatic event(s).
- C. Persistent avoidance of stimuli associated with the traumatic event(s), beginning after the traumatic event(s) occurred, as evidenced by one or both of the following:
1. Avoidance of or efforts to avoid distressing memories, thoughts, or feelings about or closely associated with the traumatic event(s).
 2. Avoidance of or efforts to avoid external reminders (people, places, conversations, activities, objects, situations) that arouse distressing memories, thoughts, or feelings about or closely associated with the traumatic event(s).
- D. Negative alterations in cognitions and mood associated with the traumatic event(s), beginning or worsening after the traumatic event(s) occurred as evidenced by two (or more) of the following:
1. Inability to remember an important aspect of the traumatic event(s) (typically due to dissociative amnesia and not to other factors such as head injury, alcohol, or drugs).
 2. Persistent and exaggerated negative beliefs or expectations about oneself, others, or the world (e.g., “I am bad,” “No one can be trusted,” “The world is completely dangerous,” “My whole nervous system is permanently ruined”).
 3. Persistent, distorted cognitions about the cause or consequences of the traumatic event(s) that lead the individual to blame himself/herself or others.
 4. Persistent negative emotional state (e.g., fear, horror, anger, guilt, or shame).
 5. Markedly diminished interest or participation in significant activities.
 6. Feelings of detachment or estrangement from others.
 7. Persistent inability to experience positive emotions (e.g., inability to experience happiness, satisfaction, or loving feelings).
- E. Marked alterations in arousal and reactivity associated with the traumatic event(s), beginning or worsening after the traumatic event(s) occurred, as evidenced by two (or more) of the following:
1. Irritable behavior and angry outbursts (with little or no provocation) typically expressed as verbal or physical aggression toward people or objects.
 2. Reckless or self-destructive behavior.
 3. Hypervigilance.
 4. Exaggerated startle response.
 5. Problems with concentration.
 6. Sleep disturbance (e.g., difficulty falling or staying asleep or restless sleep).
- F. Duration of the disturbance (criteria B, C, D, and E) is more than 1 month.
- G. The disturbance causes clinically significant distress or impairment in social, occupational, or other important areas of functioning.
- H. The disturbance is not attributable to the physiologic effects of a substance (e.g., medication, alcohol) or another medical condition.

Source: *Diagnostic and Statistical Manual of Mental Disorders*, 5th ed. Washington, DC, American Psychiatric Association, 2013.

to help with insomnia (50–150 mg qhs). Carbamazepine, valproic acid, and alprazolam have also independently produced improvement in uncontrolled trials. Psychotherapeutic strategies for PTSD help the patient overcome avoidance behaviors and demoralization and master fear of recurrence of the trauma; therapies that encourage the patient to dismantle avoidance behaviors through stepwise focusing on the experience of the traumatic event, such as trauma-focused cognitive-behavioral therapy, exposure therapy, and eye movement desensitization and reprocessing, are the most effective.

OBSESSIVE-COMPULSIVE DISORDER

Clinical Manifestations Obsessive-compulsive disorder (OCD) is characterized by obsessive thoughts and compulsive behaviors that impair everyday functioning. Fears of contamination and germs are common, as are handwashing, counting behaviors, and having to check and recheck such actions as whether a door is locked. The degree to which the disorder is disruptive for the individual varies, but in all cases, obsessive-compulsive activities take up >1 h per day and are

undertaken to relieve the anxiety triggered by the core fear. Patients often conceal their symptoms, usually because they are embarrassed by the content of their thoughts or the nature of their actions. Physicians must ask specific questions regarding recurrent thoughts and behaviors, particularly if physical clues such as chafed and reddened hands or patchy hair loss (from repetitive hair pulling, or trichotillomania) are present. Comorbid conditions are common, the most frequent being depression, other anxiety disorders, eating disorders, and tics. OCD has a lifetime prevalence of 2–3% worldwide. Onset is usually gradual, beginning in early adulthood, but childhood onset is not rare. The disorder usually has a waxing and waning course, but some cases may show a steady deterioration in psychosocial functioning.

Etiology and Pathophysiology A genetic contribution to OCD is suggested by twin studies, but no susceptibility gene for OCD has been identified to date. Family studies show an aggregation of OCD with Tourette’s disorder, and both are more common in males and in first-born children.

The anatomy of obsessive-compulsive behavior is thought to include the orbital frontal cortex, caudate nucleus, and globus pallidus. The caudate nucleus appears to be involved in the acquisition

TREATMENT **OBSESSIVE-COMPULSIVE DISORDER**

Clomipramine, fluoxetine, fluvoxamine, and sertraline are approved for the treatment of OCD in adults (fluvoxamine is also approved for children). Clomipramine is a TCA that is often tolerated poorly owing to anticholinergic and sedative side effects at the doses required to treat the illness (25–250 mg/d); its efficacy in OCD is unrelated to its antidepressant activity. Fluoxetine (5–60 mg/d), fluvoxamine (25–300 mg/d), and sertraline (50–150 mg/d) are as effective as clomipramine and have a more benign side effect profile. Only 50–60% of patients with OCD show adequate improvement with pharmacotherapy alone. In treatment-resistant cases, augmentation with other serotonergic agents such as buspirone, or with a neuroleptic or benzodiazepine, may be beneficial, and in severe cases, deep brain stimulation has been found to be effective. When a therapeutic response is achieved, long-duration maintenance therapy is usually indicated.

For many individuals, particularly those with time-consuming compulsions, behavior therapy will result in as much improvement as that afforded by medication. Effective techniques include the gradual increase in exposure to stressful situations, maintenance of a diary to clarify stressors, and homework assignments that substitute new activities for compulsive behaviors.

MOOD DISORDERS

Mood disorders are characterized by a disturbance in the regulation of mood, behavior, and affect. Mood disorders are subdivided into (1) depressive disorders, (2) bipolar disorders, and (3) depression in association with medical illness or alcohol and substance abuse (Chaps. 467 through 471e). Major depressive disorder (MDD) is differentiated from bipolar disorder by the absence of a manic or hypomanic episode. The relationship between pure depressive syndromes and bipolar disorders is not well understood; MDD is more frequent in families of bipolar individuals, but the reverse is not true. In the Global Burden of Disease Study conducted by the World Health Organization, unipolar major depression ranked fourth among all diseases in terms of disability-adjusted life-years and was projected to rank second by the year 2020. In the United States, lost productivity directly related to mood disorders has been estimated at \$55.1 billion per year.

DEPRESSION IN ASSOCIATION WITH MEDICAL ILLNESS

Depression occurring in the context of medical illness is difficult to evaluate. Depressive symptomatology may reflect the psychological stress of coping with the disease, may be caused by the disease process itself or by the medications used to treat it, or may simply coexist in time with the medical diagnosis.

Virtually every class of medication includes some agent that can induce depression. Antihypertensive drugs, anticholesterolemic agents, and antiarrhythmic agents are common triggers of depressive symptoms. Iatrogenic depression should also be considered in patients receiving glucocorticoids, antimicrobials, systemic analgesics, antiparkinsonian medications, and anticonvulsants. To decide whether a causal relationship exists between pharmacologic therapy and a patient's change in mood, it may sometimes be necessary to undertake an empirical trial of an alternative medication.

Between 20 and 30% of cardiac patients manifest a depressive disorder; an even higher percentage experience depressive symptomatology when self-reporting scales are used. Depressive symptoms following unstable angina, myocardial infarction, cardiac bypass surgery, or heart transplant impair rehabilitation and are associated with higher rates of mortality and medical morbidity. Depressed patients often show decreased variability in heart rate (an index of reduced parasymp-

athetic nervous system activity), which may predispose individuals to ventricular arrhythmia and increased morbidity. Depression also appears to increase the risk of developing coronary heart disease, possibly through increased platelet aggregation. TCAs are contraindicated in patients with bundle branch block, and TCA-induced tachycardia is an additional concern in patients with congestive heart failure. SSRIs appear not to induce ECG changes or adverse cardiac events and thus are reasonable first-line drugs for patients at risk for TCA-related complications. SSRIs may interfere with hepatic metabolism of anticoagulants, however, causing increased anticoagulation.

In patients with cancer, the mean prevalence of depression is 25%, but depression occurs in 40–50% of patients with cancers of the pancreas or oropharynx. This association is not due to the effect of cachexia alone, as the higher prevalence of depression in patients with pancreatic cancer persists when compared to those with advanced gastric cancer. Initiation of antidepressant medication in cancer patients has been shown to improve quality of life as well as mood. Psychotherapeutic approaches, particularly group therapy, may have some effect on short-term depression, anxiety, and pain symptoms.

Depression occurs frequently in patients with neurologic disorders, particularly cerebrovascular disorders, Parkinson's disease, dementia, multiple sclerosis, and traumatic brain injury. One in five patients with left-hemisphere stroke involving the dorsolateral frontal cortex experiences major depression. Late-onset depression in otherwise cognitively normal individuals increases the risk of a subsequent diagnosis of Alzheimer's disease. All classes of antidepressant agents are effective against these depressions, as are, in some cases, stimulant compounds.

The reported prevalence of depression in patients with diabetes mellitus varies from 8 to 27%, with the severity of the mood state correlating with the level of hyperglycemia and the presence of diabetic complications. Treatment of depression may be complicated by effects of antidepressive agents on glycemic control. MAOIs can induce hypoglycemia and weight gain, whereas TCAs can produce hyperglycemia and carbohydrate craving. SSRIs and SNRIs, like MAOIs, may reduce fasting plasma glucose, but they are easier to use and may also improve dietary and medication compliance.

Hypothyroidism is frequently associated with features of depression, most commonly depressed mood and memory impairment. Hyperthyroid states may also present in a similar fashion, usually in geriatric populations. Improvement in mood usually follows normalization of thyroid function, but adjunctive antidepressant medication is sometimes required. Patients with subclinical hypothyroidism can also experience symptoms of depression and cognitive difficulty that respond to thyroid replacement.

The lifetime prevalence of depression in HIV-positive individuals has been estimated at 22–45%. The relationship between depression and disease progression is multifactorial and likely to involve psychological and social factors, alterations in immune function, and central nervous system (CNS) disease. Chronic hepatitis C infection is also associated with depression, which may worsen with interferon- α treatment.

Some chronic disorders of uncertain etiology, such as chronic fatigue syndrome (Chap. 464e) and fibromyalgia (Chap. 396), are strongly associated with depression and anxiety; patients may benefit from antidepressant treatment or anticonvulsant agents such as pregabalin.

DEPRESSIVE DISORDERS

Clinical Manifestations Major depression is defined as depressed mood on a daily basis for a minimum duration of 2 weeks (Table 466-7). An episode may be characterized by sadness, indifference, apathy, or irritability and is usually associated with changes in sleep patterns, appetite, and weight; motor agitation or retardation; fatigue; impaired concentration and decision making; feelings of shame or guilt; and thoughts of death or dying. Patients with depression have a profound loss of pleasure in all enjoyable activities, exhibit early morning awakening, feel that the dysphoric mood state is qualitatively different from sadness, and often notice a diurnal variation in mood (worse in morning hours). Patients experiencing bereavement or grief may exhibit many of the same signs and symptoms of major depression, although the emphasis is usually on feelings of emptiness and loss, rather than

TABLE 466-7 CRITERIA FOR A MAJOR DEPRESSIVE EPISODE

- A. Five (or more) of the following symptoms have been present during the same 2-week period and represent a change from previous functioning; at least one of the symptoms is either (1) depressed mood or (2) loss of interest or pleasure. **Note:** Do not include symptoms that are clearly attributable to another medical condition.
1. Depressed mood most of the day, nearly every day, as indicated by either subjective report (e.g., feels sad, empty, hopeless) or observation made by others (e.g., appears tearful).
 2. Markedly diminished interest or pleasure in all, or almost all, activities most of the day, nearly every day (as indicated by either subjective account or observation).
 3. Significant weight loss when not dieting or weight gain (e.g., a change of >5% of body weight in a month), or decrease or increase in appetite nearly every day.
 4. Insomnia or hypersomnia nearly every day.
 5. Psychomotor agitation or retardation nearly every day (observable by others, not merely subjective feelings of restlessness or being slowed down).
 6. Fatigue or loss of energy nearly every day.
 7. Feelings of worthlessness or excessive or inappropriate guilt (which may be delusional) nearly every day (not merely self-reproach or guilt about being sick).
 8. Diminished ability to think or concentrate, or indecisiveness, nearly every day (either by subjective account or as observed by others).
 9. Recurrent thoughts of death (not just fear of dying), recurrent suicidal ideation without a specific plan, or a suicide attempt or a specific plan for committing suicide
- B. The symptoms cause clinically significant distress or impairment in social, occupational, or other important areas of functioning
- C. The episode is not attributable to the physiologic effects of a substance or to another medical condition.
- D. The occurrence of the major depressive episode is not better explained by seasonal affective disorder, schizophrenia, schizophreniform disorder, delusional disorder, or other specified and unspecified schizophrenia spectrum and other psychotic disorders.
- E. There has never been a manic episode or a hypomanic episode.

Source: *Diagnostic and Statistical Manual of Mental Disorders*, 5th ed. Washington, DC, American Psychiatric Association, 2013.

anhedonia and loss of self-esteem, and the duration is usually limited. In certain cases, however, the diagnosis of major depression may be warranted even in the context of a significant loss.

Approximately 15% of the population experiences a major depressive episode at some point in life, and 6–8% of all outpatients in primary care settings satisfy diagnostic criteria for the disorder. Depression is often undiagnosed, and even more frequently, it is treated inadequately. If a physician suspects the presence of a major depressive episode, the initial task is to determine whether it represents unipolar or bipolar depression or is one of the 10–15% of cases that are secondary to general medical illness or substance abuse. Physicians should also assess the risk of suicide by direct questioning, as patients are often reluctant to verbalize such thoughts without prompting. If specific plans are uncovered or if significant risk factors exist (e.g., a past history of suicide attempts, profound hopelessness, concurrent medical illness, substance abuse, or social isolation), the patient must be referred to a mental health specialist for immediate care. The physician should specifically probe each of these areas in an empathic and hopeful manner, being sensitive to denial and possible minimization of distress. The presence of anxiety, panic, or agitation significantly increases near-term suicidal risk. Approximately 4–5% of all depressed patients will commit suicide; most will have sought help from physicians within 1 month of their deaths.

In some depressed patients, the mood disorder does not appear to be episodic and is not clearly associated with either psychosocial dysfunction or change from the individual's usual experience in life. Persistent depressive disorder (*dysthymic disorder*) consists of a pattern of chronic (at least 2 years), ongoing depressive symptoms that are usually less severe and/or less numerous than those found in major

depression, but the functional consequences may be equivalent to or even greater; the two conditions are sometimes difficult to separate and can occur together (“double depression”). Many patients who exhibit a profile of pessimism, disinterest, and low self-esteem respond to antidepressant treatment. Persistent and chronic depressive disorders occur in approximately 2% of the general population.

Depression is approximately twice as common in women as in men, and the incidence increases with age in both sexes. Twin studies indicate that the liability to major depression of early onset (before age 25) is largely genetic in origin. Negative life events can precipitate and contribute to depression, but genetic factors influence the sensitivity of individuals to these stressful events. In most cases, both biologic and psychosocial factors are involved in the precipitation and unfolding of depressive episodes. The most potent stressors appear to involve death of a relative, assault, or severe marital or relationship problems.

Unipolar depressive disorders usually begin in early adulthood and recur episodically over the course of a lifetime. The best predictor of future risk is the number of past episodes; 50–60% of patients who have a first episode have at least one or two recurrences. Some patients experience multiple episodes that become more severe and frequent over time. The duration of an untreated episode varies greatly, ranging from a few months to ≥ 1 year. The pattern of recurrence and clinical progression in a developing episode are also variable. Within an individual, the nature of episodes (e.g., specific presenting symptoms, frequency and duration) may be similar over time. In a minority of patients, a severe depressive episode may progress to a psychotic state; in elderly patients, depressive symptoms may be associated with cognitive deficits mimicking dementia (“pseudodementia”). A seasonal pattern of depression, called *seasonal affective disorder*, may manifest with onset and remission of episodes at predictable times of the year. This disorder is more common in women, whose symptoms are anergy, fatigue, weight gain, hypersomnia, and episodic carbohydrate craving. The prevalence increases with distance from the equator, and improvement may occur by altering light exposure.

Etiology and Pathophysiology Although evidence for genetic transmission of unipolar depression is not as strong as in bipolar disorder, monozygotic twins have a higher concordance rate (46%) than dizygotic siblings (20%), with little support for any effect of a shared family environment.

Neuroendocrine abnormalities that reflect the neurovegetative signs and symptoms of depression include: (1) increased cortisol and corticotropin-releasing hormone (CRH) secretion, (2) an increase in adrenal size, (3) a decreased inhibitory response of glucocorticoids to dexamethasone, and (4) a blunted response of thyroid-stimulating hormone (TSH) level to infusion of thyroid-releasing hormone (TRH). Antidepressant treatment leads to normalization of these abnormalities. Major depression is also associated with changes in levels of pro-inflammatory cytokines and neurotrophins.

Diurnal variations in symptom severity and alterations in circadian rhythmicity of a number of neurochemical and neurohumoral factors suggest that biologic differences may be secondary to a primary defect in regulation of biologic rhythms. Patients with major depression show consistent findings of a decrease in rapid eye movement (REM) sleep onset (REM latency), an increase in REM density, and, in some subjects, a decrease in stage IV delta slow-wave sleep.

Although antidepressant drugs inhibit neurotransmitter uptake within hours, their therapeutic effects typically emerge over several weeks, implicating adaptive changes in second messenger systems and transcription factors as possible mechanisms of action.

The pathogenesis of depression is discussed in detail in Chap. 465e.

TREATMENT DEPRESSIVE DISORDERS

Treatment planning requires coordination of short-term strategies to induce remission combined with longer term maintenance designed to prevent recurrence. The most effective intervention

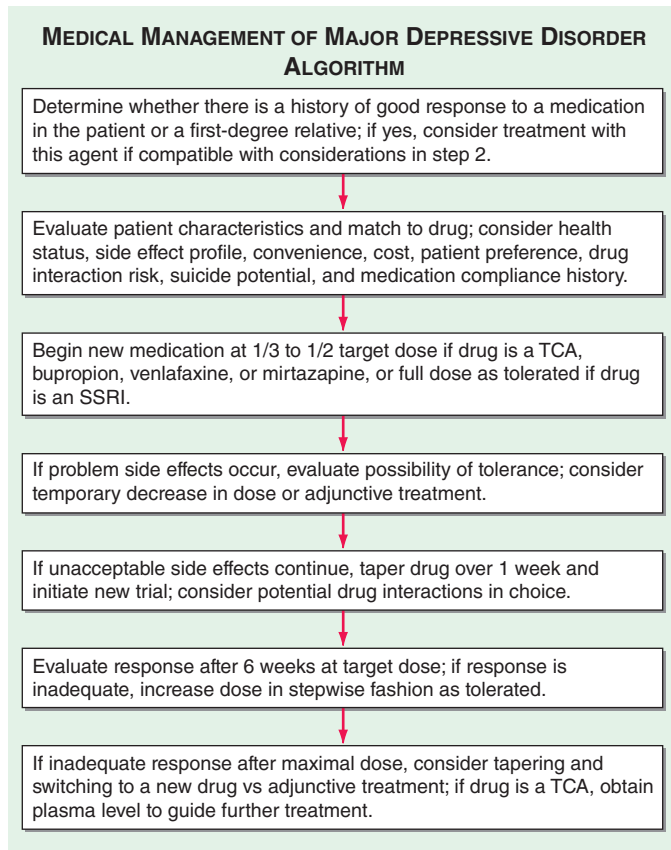


FIGURE 466-1 A guideline for the medical management of major depressive disorder. SSRI, selective serotonin reuptake inhibitor; TCA, tricyclic antidepressant.

for achieving remission and preventing relapse is medication, but combined treatment, incorporating psychotherapy to help the patient cope with decreased self-esteem and demoralization, improves outcome (Fig. 466-1). Approximately 40% of primary care patients with depression drop out of treatment and discontinue medication if symptomatic improvement is not noted within a month, unless additional support is provided. Outcome improves with (1) increased intensity and frequency of visits during the first 4–6 weeks of treatment, (2) supplemental educational materials, and (3) psychiatric consultation as indicated. Despite the widespread use of SSRIs and other second-generation antidepressants, there is no convincing evidence that these classes of antidepressants are more efficacious than TCAs. Between 60 and 70% of all depressed patients respond to any drug chosen, if it is given in a sufficient dose for 6–8 weeks.

A rational approach to selecting which antidepressant to use involves matching the patient's preference and medical history with the metabolic and side effect profile of the drug (Tables 466-4 and 466-5). A previous response, or a family history of a positive response, to a specific antidepressant often suggests that that drug be tried first. Before initiating antidepressant therapy, the physician should evaluate the possible contribution of comorbid illnesses and consider their specific treatment. In individuals with suicidal ideation, particular attention should be paid to choosing a drug with low toxicity if taken in overdose. Newer antidepressant drugs are distinctly safer in this regard; nevertheless, the advantages of TCAs have not been completely superseded. The existence of generic equivalents makes TCAs relatively cheap, and for secondary tricyclics, particularly nortriptyline and desipramine, well-defined relationships among dose, plasma level, and therapeutic response exist. The steady-state plasma level achieved for a given drug dose can vary more than 10-fold between individuals, and plasma levels may help in interpreting apparent resistance to treatment

and/or unexpected drug toxicity. The principal side effects of TCAs are antihistamine (sedation) and anticholinergic (constipation, dry mouth, urinary hesitancy, blurred vision). TCAs are contraindicated in patients with serious cardiovascular risk factors, and overdoses of tricyclic agents can be lethal, with desipramine carrying the greatest risk. It is judicious to prescribe only a 10-day supply when suicide is a risk. Most patients require a daily dose of 150–200 mg of imipramine or amitriptyline or its equivalent to achieve a therapeutic blood level of 150–300 ng/mL and a satisfactory remission; some patients show a partial effect at lower doses. Geriatric patients may require a low starting dose and slow escalation. Ethnic differences in drug metabolism are significant, with Hispanic, Asian, and black patients generally requiring lower doses than whites to achieve a comparable blood level. P450 profiling using genetic chip technology may be clinically useful in predicting individual sensitivity.

Second-generation antidepressants are similar to tricyclics in their effect on neurotransmitter reuptake, although some also have specific actions on catecholamine and indolamine receptors as well. Amoxapine is a dibenzoxazepine derivative that blocks norepinephrine and serotonin reuptake and has a metabolite that shows a degree of dopamine blockade. Long-term use of this drug carries a risk of tardive dyskinesia. Maprotiline is a potent noradrenergic reuptake blocker that has little anticholinergic effect but may produce seizures. Bupropion is a novel antidepressant whose mechanism of action is thought to involve enhancement of noradrenergic function. It has no anticholinergic, sedating, or orthostatic side effects and has a low incidence of sexual side effects. It may, however, be associated with stimulant-like side effects, may lower seizure threshold, and has an exceptionally short half-life, requiring frequent dosing. An extended-release preparation is available.

SSRIs such as fluoxetine, sertraline, paroxetine, citalopram, and escitalopram cause a lower frequency of anticholinergic, sedating, and cardiovascular side effects but a possibly greater incidence of gastrointestinal complaints, sleep impairment, and sexual dysfunction than do TCAs. Akathisia, involving an inner sense of restlessness and anxiety in addition to increased motor activity, may also be more common, particularly during the first week of treatment. One concern is the risk of "serotonin syndrome," which is thought to result from hyperstimulation of brainstem 5-HT_{1A} receptors and characterized by myoclonus, agitation, abdominal cramping, hyperpyrexia, hypertension, and potentially death. Serotonergic agonists taken in combination should be monitored closely for this reason. Considerations such as half-life, compliance, toxicity, and drug-drug interactions may guide the choice of a particular SSRI. Fluoxetine and its principal active metabolite, norfluoxetine, for example, have a combined half-life of almost 7 days, resulting in a delay of 5 weeks before steady-state levels are achieved and a similar delay for complete drug excretion once its use is discontinued. All the SSRIs may impair sexual function, resulting in diminished libido, impotence, or difficulty in achieving orgasm. Sexual dysfunction frequently results in noncompliance and should be asked about specifically. Sexual dysfunction can sometimes be ameliorated by lowering the dose, by instituting weekend drug holidays (two or three times a month), or by treatment with amantadine (100 mg tid), bethanechol (25 mg tid), buspirone (10 mg tid), or bupropion (100–150 mg/d). Paroxetine appears to be more anticholinergic than either fluoxetine or sertraline, and sertraline carries a lower risk of producing an adverse drug interaction than the other two. Rare side effects of SSRIs include angina due to vasospasm and prolongation of the prothrombin time. Escitalopram is the most specific of currently available SSRIs and appears to have no specific inhibitory effects on the P450 system.

Venlafaxine, desvenlafaxine, duloxetine, vilazodone, vortioxetine, and levomilnacipran block the reuptake of both norepinephrine and serotonin but produce relatively little in the way of traditional tricyclic side effects. Unlike the SSRIs, venlafaxine and vortioxetine have relatively linear dose-response curves. Patients on immediate release venlafaxine should be monitored for a possible increase in diastolic blood pressure, and multiple daily dosing is required

because of the drug's short half-life. An extended-release form is available and has a somewhat lower incidence of gastrointestinal side effects. Mirtazapine is a TCA that has a unique spectrum of activity. It increases noradrenergic and serotonergic neurotransmission through a blockade of central α_2 -adrenergic receptors and postsynaptic 5-HT₂ and 5-HT₃ receptors. It is also strongly antihistaminic and, as such, may produce sedation. Levomilnacipran is the most noradrenergic of the SNRIs and theoretically may be appropriate for patients with more severe fatigue and anergia.

With the exception of citalopram and escitalopram, each of the SSRIs may inhibit one or more cytochrome P450 enzymes. Depending on the specific isoenzyme involved, the metabolism of a number of concomitantly administered medications can be dramatically affected. Fluoxetine and paroxetine, for example, by inhibiting 2D6, can cause dramatic increases in the blood level of type 1C antiarrhythmics, whereas sertraline, by acting on 3A4, may alter blood levels of carbamazepine or digoxin. Depending on drug specificity for a particular CYP enzyme for its own metabolism, concomitant medications or dietary factors, such as grapefruit juice, may in turn affect the efficacy or toxicity of the SSRI.

The MAOIs are highly effective, particularly in atypical depression, but the risk of hypertensive crisis following intake of tyramine-containing food or sympathomimetic drugs makes them inappropriate as first-line agents. Transdermal selegiline may avert this risk at low dose. Common side effects include orthostatic hypotension, weight gain, insomnia, and sexual dysfunction. MAOIs should not be used concomitantly with SSRIs, because of the risk of serotonin syndrome, or with TCAs, because of possible hyperadrenergic effects.

Electroconvulsive therapy is at least as effective as medication, but its use is reserved for treatment-resistant cases and delusional depressions. Transcranial magnetic stimulation (TMS) is approved for treatment-resistant depression and has been shown to have efficacy in several controlled trials. Vagus nerve stimulation (VNS) has also recently been approved for treatment-resistant depression, but its degree of efficacy is controversial. Deep brain stimulation and ketamine, a glutamatergic antagonist, are experimental approaches for treatment-resistant cases.

Regardless of the treatment undertaken, the response should be evaluated after ~2 months. Three-quarters of patients show improvement by this time, but if remission is inadequate, the patient should be questioned about compliance, and an increase in medication dose should be considered if side effects are not troublesome. If this approach is unsuccessful, referral to a mental health specialist is advised. Strategies for treatment then include selection of an alternative drug, combinations of antidepressants, and/or adjunctive treatment with other classes of drugs, including lithium, thyroid hormone, atypical antipsychotic agents, and dopamine agonists. A large randomized trial (STAR-D) was unable to show preferential efficacy, but the addition of certain atypical antipsychotic drugs (quetiapine extended-release; aripiprazole) has received FDA approval, as has usage of a combined medication, olanzapine and fluoxetine (Symbyax). Patients whose response to an SSRI wanes over time may benefit from the addition of buspirone (10 mg tid) or pindolol (2–5 mg tid) or small amounts of a TCA such as desipramine (25 mg bid or tid). Most patients will show some degree of response, but aggressive treatment should be pursued until remission is achieved, and drug treatment should be continued for at least 6–9 more months to prevent relapse. In patients who have had two or more episodes of depression, indefinite maintenance treatment should be considered.

It is essential to educate patients both about depression and the benefits and side effects of medications they are receiving. Advice about stress reduction and cautions that alcohol may exacerbate depressive symptoms and impair drug response are helpful. Patients should be given time to describe their experience, their outlook, and the impact of the depression on them and their families. Occasional empathic silence may be as helpful for the treatment alliance as verbal reassurance. Controlled trials have shown that cognitive-behavioral and interpersonal therapies are effective

in improving psychological and social adjustment and that a combined treatment approach is more successful than medication alone for many patients.

BIPOLAR DISORDER

Clinical Manifestations Bipolar disorder is characterized by unpredictable swings in mood from mania (or hypomania) to depression. Some patients suffer only from recurrent attacks of *mania*, which in its pure form is associated with increased psychomotor activity; excessive social extroversion; decreased need for sleep; impulsivity and impairment in judgment; and expansive, grandiose, and sometimes irritable mood (Table 466-8). In severe mania, patients may experience delusions and paranoid thinking indistinguishable from schizophrenia. One-half of patients with bipolar disorder present with a mixture of psychomotor agitation and activation with dysphoria, anxiety, and irritability. It may be difficult to distinguish *mixed mania* from *agitated depression*. In some bipolar patients (*bipolar II disorder*), the full criteria for mania are lacking, and the requisite recurrent depressions are separated by periods of mild activation and increased energy (hypomania). In *cyclothymic disorder*, there are numerous hypomanic periods, usually of relatively short duration, alternating with clusters of depressive symptoms that fail, either in severity or duration, to meet the criteria of major depression. The mood fluctuations are chronic and should be present for at least 2 years before the diagnosis is made.

Manic episodes typically emerge over a period of days to weeks, but onset within hours is possible, usually in the early morning hours. An untreated episode of either depression or mania can be as short as several weeks or last as long as 8–12 months, and rare patients have an unremitting chronic course. The term *rapid cycling* is used for patients who have four or more episodes of either depression or mania in a given year. This pattern occurs in 15% of all patients, almost all of whom are women. In some cases, rapid cycling is linked to an underlying thyroid dysfunction, and in others, it is iatrogenically triggered by prolonged antidepressant treatment. Approximately one-half of patients have sustained difficulties in work performance and psychosocial functioning, with depressive phases being more responsible for impairment than mania.

TABLE 466-8 CRITERIA FOR A MANIC EPISODE

- A. A distinct period of abnormally and persistently elevated, expansive, or irritable mood and abnormally and persistently increased goal-directed activity or energy, lasting at least 1 week and present most of the day, nearly every day (or any duration if hospitalization is necessary).
- B. During the period of the mood disturbance and increased energy or activity, three (or more) of the following symptoms (four if the mood is only irritable) are present to a significant degree and represent a noticeable change from usual behavior:
 1. Inflated self-esteem or grandiosity.
 2. Decreased need for sleep (e.g., feels rested after only 3 h of sleep).
 3. More talkative than usual or pressure to keep talking.
 4. Flight of ideas or subjective experience that thoughts are racing.
 5. Distractibility (i.e., attention too easily drawn to unimportant or irrelevant external stimuli), as reported or observed.
 6. Increase in goal-directed activity (either socially, at work or school, or sexually) or psychomotor agitation (i.e., purposeless non-goal-directed activity).
 7. Excessive involvement in activities that have a high potential for painful consequences (e.g., engaging in unrestrained buying sprees, sexual indiscretions, or foolish business investments).
- C. The mood disturbance is sufficiently severe to cause marked impairment in social or occupational functioning or to necessitate hospitalization to prevent harm to self or others, or there are psychotic features.
- D. The episode is not attributable to the physiologic effects of a substance (e.g., a drug of abuse, a medication, or other treatment) or another medical condition.

Source: *Diagnostic and Statistical Manual of Mental Disorders*, 5th ed. Washington, DC, American Psychiatric Association, 2013.

Bipolar disorder is common, affecting ~1.5% of the population in the United States. Onset is typically between 20 and 30 years of age, but many individuals report premorbid symptoms in late childhood or early adolescence. The prevalence is similar for men and women; women are likely to have more depressive and men more manic episodes over a lifetime.

Differential Diagnosis The differential diagnosis of mania includes secondary mania induced by stimulant or sympathomimetic drugs, hyperthyroidism, AIDS, and neurologic disorders such as Huntington's or Wilson's disease and cerebrovascular accidents. Comorbidity with alcohol and substance abuse is common, either because of poor judgment and increased impulsivity or because of an attempt to self-treat the underlying mood symptoms and sleep disturbances.

Etiology and Pathophysiology Genetic predisposition to bipolar disorder is evident from family studies; the concordance rate for monozygotic twins approaches 80%. Patients with bipolar disorder also appear to have altered circadian rhythmicity, and lithium may exert its therapeutic benefit through a resynchronization of intrinsic rhythms keyed to the light/dark cycle.

TREATMENT BIPOLAR DISORDER

(Table 466-9) Lithium carbonate is the mainstay of treatment in bipolar disorder, although sodium valproate and carbamazepine, as well as a number of second-generation antipsychotic agents (aripiprazole, asenapine, olanzapine, quetiapine, risperidone, ziprasidone), also have FDA approval for the treatment of acute mania. Oxcarbazepine is not FDA approved, but appears to enjoy carbamazepine's spectrum of efficacy. The response rate to lithium carbonate

is 70–80% in acute mania, with beneficial effects appearing in 1–2 weeks. Lithium also has a prophylactic effect in prevention of recurrent mania and, to a lesser extent, in the prevention of recurrent depression. A simple cation, lithium is rapidly absorbed from the gastrointestinal tract and remains unbound to plasma or tissue proteins. Some 95% of a given dose is excreted unchanged through the kidneys within 24 h.

Serious side effects from lithium are rare, but minor complaints such as gastrointestinal discomfort, nausea, diarrhea, polyuria, weight gain, skin eruptions, alopecia, and edema are common. Over time, urine-concentrating ability may be decreased, but significant nephrotoxicity does not usually occur. Lithium exerts an antithyroid effect by interfering with the synthesis and release of thyroid hormones. More serious side effects include tremor, poor concentration and memory, ataxia, dysarthria, and incoordination. There is suggestive, but not conclusive, evidence that lithium is teratogenic, inducing cardiac malformations in the first trimester.

In the treatment of acute mania, lithium is initiated at 300 mg bid or tid, and the dose is then increased by 300 mg every 2–3 days to achieve blood levels of 0.8–1.2 meq/L. Because the therapeutic effect of lithium may not appear until after 7–10 days of treatment, adjunctive usage of lorazepam (1–2 mg every 4 h) or clonazepam (0.5–1 mg every 4 h) may be beneficial to control agitation. Antipsychotics are indicated in patients with severe agitation who respond only partially to benzodiazepines. Patients using lithium should be monitored closely, since the blood levels required to achieve a therapeutic benefit are close to those associated with toxicity.

Valproic acid may be better than lithium for patients who experience rapid cycling (i.e., more than four episodes a year) or who present with a mixed or dysphoric mania. Tremor and weight gain are the most common side effects; hepatotoxicity and pancreatitis are rare toxicities.

The recurrent nature of bipolar mood disorder necessitates maintenance treatment. A sustained blood lithium level of at least 0.8 meq/L is important for optimal prophylaxis and has been shown to reduce the risk of suicide, a finding not yet apparent for other mood stabilizers. Combinations of mood stabilizers together or with atypical antipsychotic drugs are sometimes required to maintain mood stability. Quetiapine extended release, olanzapine, risperidone, and lamotrigine have been approved for maintenance treatment as sole agents, in combination with lithium and with aripiprazole and ziprasidone as adjunctive drugs. Lurasidone, olanzapine/fluoxetine, and quetiapine are also approved to treat acute depressive episodes in bipolar disorder. Compliance is frequently an issue and often requires enlistment and education of concerned family members. Efforts to identify and modify psychosocial factors that may trigger episodes are important, as is an emphasis on lifestyle regularity. Antidepressant medications are sometimes required for the treatment of severe breakthrough depressions, but their use should generally be avoided during maintenance treatment because of the risk of precipitating mania or accelerating the cycle frequency. Loss of efficacy over time may be observed with any of the mood-stabilizing agents. In such situations, an alternative agent or combination therapy is usually helpful.

TABLE 466-9 CLINICAL PHARMACOLOGY OF MOOD STABILIZERS

Agent and Dosing	Side Effects and Other Effects
Lithium	Common Side Effects
Starting dose: 300 mg bid or tid	Nausea/anorexia/diarrhea, fine tremor, thirst, polyuria, fatigue, weight gain, acne, folliculitis, neutrophilia, hypothyroidism
Therapeutic blood level: 0.8–1.2 meq/L	Blood level is increased by thiazides, tetracyclines, and NSAIDs
	Blood level is decreased by bronchodilators, verapamil, and carbonic anhydrase inhibitors
	<i>Rare side effects:</i> Neurotoxicity, renal toxicity, hypercalcemia, ECG changes
Valproic Acid	Common Side Effects
Starting dose: 250 mg tid	Nausea/anorexia, weight gain, sedation, tremor, rash, alopecia
Therapeutic blood level: 50–125 µg/mL	Inhibits hepatic metabolism of other medications
	<i>Rare side effects:</i> Pancreatitis, hepatotoxicity, Stevens-Johnson syndrome
Carbamazepine/Oxcarbazepine	Common Side Effects
Starting dose: 200 mg bid for carbamazepine, 150 mg bid for oxcarbazepine	Nausea/anorexia, sedation, rash, dizziness/ataxia
Therapeutic blood level: 4–12 µg/mL for carbamazepine	Carbamazepine, but not oxcarbazepine, induces hepatic metabolism of other medications
	<i>Rare side effects:</i> Hyponatremia, agranulocytosis, Stevens-Johnson syndrome
Lamotrigine	Common Side Effects
Starting dose: 25 mg/d	Rash, dizziness, headache, tremor, sedation, nausea
	<i>Rare side effect:</i> Stevens-Johnson syndrome

Abbreviations: ECG, electrocardiogram; NSAIDs, nonsteroidal anti-inflammatory drugs.

SOMATIC SYMPTOM DISORDER

Many patients presenting in general medical practice, perhaps as many as 5–7%, will experience a somatic symptom(s) as particularly distressing and preoccupying, to the point that it comes to dominate their thoughts, feelings, and beliefs and interferes to a varying degree with everyday functioning. Although the absence of a medical explanation for these complaints was historically emphasized as a diagnostic element, it has been recognized that the patient's interpretation and elaboration of the experience is the critical defining factor and that patients with well-established medical causation may qualify for the diagnosis. Multiple complaints are typical, but severe single symptoms can occur as well. Comorbidity with depressive and anxiety disorders

is common and may affect the severity of the experience and its functional consequences. Personality factors may be a significant risk factor, as may a low level of educational or socioeconomic status or a history of recent stressful life events. Cultural factors are relevant as well and should be incorporated into the evaluation. Individuals who have persistent preoccupations about having or acquiring a serious illness, but who do not have a specific somatic complaint, may qualify for a related diagnosis—illness anxiety disorder. The diagnosis of conversion disorder (functional neurologic symptom disorder) is used to specifically identify those individuals whose somatic complaints involve one or more symptoms of altered voluntary motor or sensory function that cannot be medically explained and that causes significant distress or impairment or requires medical evaluation.

In *factitious illnesses*, the patient consciously and voluntarily produces physical symptoms of illness. The term *Munchausen's syndrome* is reserved for individuals with particularly dramatic, chronic, or severe factitious illness. In true factitious illness, the sick role itself is gratifying. A variety of signs, symptoms, and diseases have been either simulated or caused by factitious behavior, the most common including chronic diarrhea, fever of unknown origin, intestinal bleeding or hematuria, seizures, and hypoglycemia. Factitious disorder is usually not diagnosed until 5–10 years after its onset, and it can produce significant social and medical costs. In *malingering*, the fabrication derives from a desire for some external reward such as a narcotic medication or disability reimbursement.

TREATMENT SOMATIC SYMPTOM DISORDER AND RELATED DISORDERS

Patients with somatic symptom disorder are frequently subjected to many diagnostic tests and exploratory surgeries in an attempt to find their “real” illness. Such an approach is doomed to failure and does not address the core issue. Successful treatment is best achieved through behavior modification, in which access to the physician is tightly regulated and adjusted to provide a sustained and predictable level of support that is less clearly contingent on the patient’s level of presenting distress. Visits can be brief and should not be associated with a need for a diagnostic or treatment action. Although the literature is limited, some patients may benefit from antidepressant treatment.

Any attempt to confront the patient usually creates a sense of humiliation and causes the patient to abandon treatment from that caregiver. A better strategy is to introduce psychological causation as one of a number of possible explanations in the differential diagnoses that are discussed. Without directly linking psychotherapeutic intervention to the diagnosis, the patient can be offered a face-saving means by which the pathologic relationship with the health care system can be examined and alternative approaches to life stressors developed. Specific medical treatments also may be indicated and effective in treating some of the functional consequences of conversion disorder.

FEEDING AND EATING DISORDERS

CLINICAL MANIFESTATIONS

Feeding and eating disorders constitute a group of conditions in which there is a persistent disturbance of eating or associated behaviors that significantly impair an individual’s physical health or psychosocial functioning. In DSM-5 the described categories (with the exception of pica) are defined to be mutually exclusive in a given episode, based on the understanding that although they are phenotypically similar in some ways, they differ in course, prognosis, and effective treatment interventions. Compared with DSM-IV-TR, three disorders (i.e., avoidant/restrictive food intake disorder, rumination disorder, pica) that were previously classified as disorders of infancy or childhood have been grouped together with the disorders of anorexia and bulimia nervosa. Binge-eating disorder is also now included as a formal diagnosis; the intent of each of these modifications is to encourage clinicians to be more specific in their codification of eating and feeding pathology.

PICA

Pica is diagnosed when the individual, over age 2, eats one or more nonnutritive, nonfood substances for a month or more and requires medical attention as a result. There is usually no specific aversion to food in general but a preferential choice to ingest substances such as clay, starch, soap, paper, or ash. The diagnosis requires the exclusion of specific culturally approved practices and has not been commonly found to be caused by a specific nutritional deficiency. Onset is most common in childhood but the disorder can occur in association with other major psychiatric conditions in adults. An association with pregnancy has been observed, but the condition is only diagnosed when medical risks are increased by the behavior.

RUMINATION DISORDER

In this condition, individuals who have no demonstrable associated gastrointestinal or other medical condition repeatedly regurgitate their food after eating and then either re chew or swallow it or spit it out. The behavior typically occurs on a daily basis and must persist for at least 1 month. Weight loss and malnutrition are common sequelae, and individuals may attempt to conceal their behavior, either by covering their mouth or through social avoidance while eating. In infancy, the onset is typically between 3 to 12 months, and the behavior may remit spontaneously, although in some it appears to be recurrent.

AVOIDANT/RESTRICTIVE FOOD INTAKE DISORDER

The cardinal feature of this disorder is avoidance or restriction of food intake, usually stemming from a lack of interest in or distaste of food and associated with weight loss, nutritional deficiency, dependency on nutritional supplementation, or marked impairment in psychosocial functioning, either alone or in combination. Culturally approved practices, such as fasting, or a lack of available food must be excluded as possible causes. The disorder is distinguished from anorexia nervosa by the presence of emotional factors, such as a fear of gaining weight and distortion of body image in the latter condition. Onset is usually in infancy or early childhood, but avoidant behaviors may persist into adulthood. The disorder is equally prevalent in males and females and is frequently comorbid with anxiety and cognitive and attention-deficit disorders and situations of familial stress. Developmental delay and functional deficits may be significant if the disorder is long-standing and unrecognized.

ANOREXIA NERVOSA

Individuals are diagnosed with anorexia nervosa if they restrict their caloric intake to a degree that their body weight deviates significantly from age, gender, health, and developmental norms and if they also exhibit a fear of gaining weight and an associated disturbance in body image. The condition is further characterized by differentiating those who achieve their weight loss predominantly through restricting intake or by excessive exercise (restricting type) from those who engage in recurrent binge eating and/or subsequent purging, self-induced vomiting, and usage of enemas, laxatives, or diuretics (binge-eating/purging type). Such subtyping is more state than trait specific, as individuals may transition from one profile to the other over time. Determination of whether an individual satisfies the primary criterion of significant low weight is complex and must be individualized, using all available historical information and comparison of body habitus to international body mass norms and guidelines.

Individuals with anorexia nervosa frequently lack insight into their condition and are in denial about possible medical consequences; they often are not comforted by their achieved weight loss and persist in their behaviors despite having met previously self-designated weight goals. Recent research has identified alterations in the circuitry of reward sensitivity and executive function in anorexia and implicated disturbances in frontal cortex and anterior insula regulation of interoceptive awareness of satiety and hunger. Neurochemical findings, including the role of ghrelin, remain controversial.

Onset is most common in adolescence, although onset in later life can occur. Many more females than males are affected, with a lifetime prevalence in women of up to 4%. The disorder appears

most prevalent in postindustrialized and urbanized countries and is frequently comorbid with preexisting anxiety disorders. The medical consequences of prolonged anorexia nervosa are multisystemic and can be life-threatening in severe presentations. Changes in blood chemistry include leukopenia with lymphocytosis, elevations in blood urea nitrogen, and metabolic alkalosis and hypokalemia when purging is present. History and physical examination may reveal amenorrhea in females, skin abnormalities (petechiae, lanugo hair, dryness), and signs of hypometabolic function, including hypotension, hypothermia, and sinus bradycardia. Endocrine effects include hypogonadism, growth hormone resistance, and hypercortisolemia. Osteoporosis is a longer-term concern.

The course of the disorder is variable, with some individuals recovering after a single episode, while others exhibit recurrent episodes or a chronic course. Untreated anorexia has a mortality of 5.1/1000, the highest among psychiatric conditions. Maudsley family-based therapy has proven to be an effective therapy in younger individuals, with strict behavioral contingencies used when weight loss becomes critical. No pharmacologic intervention has proven to be specifically beneficial, but comorbid depression and anxiety should be treated. Weight gain should be undertaken gradually with a goal of 0.5 to 1 pound per week to prevent refeeding syndrome. Most individuals are able to achieve remission within 5 years of the original diagnosis.

BULIMIA NERVOSA

Bulimia nervosa describes individuals who engage in recurrent and frequent (at least once a week for 3 months) periods of binge eating and who then resort to compensatory behaviors, such as self-induced purging, enemas, use of laxatives, or excessive exercise to avoid weight gain. Binge eating itself is defined as excessive food intake in a prescribed period of time, usually <2 h. As in anorexia nervosa, disturbances in body image occur and promote the behavior, but unlike in anorexia, individuals are of normal weight or even somewhat overweight. Subjects typically describe a loss of control and express shame about their actions, and often relate that their episodes are triggered by feelings of negative self-esteem or social stresses. The lifetime prevalence in women is approximately 2%, with a 10:1 female-to-male ratio. The disorder typically begins in adolescence and may be persistent over a number of years. Transition to anorexia occurs in only 10–15% of cases. Many of the medical risks associated with bulimia nervosa parallel those of anorexia nervosa and are a direct consequence of purging, including fluid and electrolyte disturbances and conduction abnormalities. Physical examination often results in no specific findings, but dental erosion and parotid gland enlargement may be present. Effective treatment approaches include SSRI antidepressants, usually in combination with cognitive-behavioral, emotion regulation, or interpersonal-based psychotherapies.

BINGE-EATING DISORDER

Binge-eating disorder is distinguished from bulimia nervosa by the absence of compensatory behaviors to prevent weight gain after an episode and by a lack of effort to restrict weight gain between episodes. Other features are similar, including distress over the behavior and the experience of loss of control, resulting in eating more rapidly or in greater amounts than intended or eating when not hungry. The 12-month prevalence in females is 1.6%, with a much lower female-to-male ratio than bulimia nervosa. Little is known about the course of the disorder, given its recent categorization, but its prognosis is markedly better than for other eating disorders, both in terms of its natural course and response to treatment. Transition to other eating disorder conditions is thought to be rare.

PERSONALITY DISORDERS

CLINICAL MANIFESTATIONS

Personality disorders are characteristic patterns of thinking, feeling, and interpersonal behavior that are relatively inflexible and cause significant functional impairment or subjective distress for the individual. The observed behaviors are not secondary to another mental disorder, nor are they precipitated by substance abuse or a general medical condition.

This distinction is often difficult to make in clinical practice, because personality change may be the first sign of serious neurologic, endocrine, or other medical illness. Patients with frontal lobe tumors, for example, can present with changes in motivation and personality while the results of the neurologic examination remain within normal limits. Individuals with personality disorders are often regarded as “difficult patients” in clinical medical practice because they are seen as excessively demanding and/or unwilling to follow recommended treatment plans. Although DSM-5 portrays personality disorders as qualitatively distinct categories, there is an alternative perspective that personality characteristics vary as a continuum between normal functioning and formal mental disorder.

Personality disorders have been grouped into three overlapping clusters. *Cluster A* includes paranoid, schizoid, and schizotypal personality disorders. It includes individuals who are odd and eccentric and who maintain an emotional distance from others. Individuals have a restricted emotional range and remain socially isolated. Patients with schizotypal personality disorder frequently have unusual perceptual experiences and express magical beliefs about the external world. The essential feature of paranoid personality disorder is a pervasive mistrust and suspiciousness of others to an extent that is unjustified by available evidence. *Cluster B* disorders include antisocial, borderline, histrionic, and narcissistic types and describe individuals whose behavior is impulsive, excessively emotional, and erratic. *Cluster C* incorporates avoidant, dependent, and obsessive-compulsive personality types; enduring traits are anxiety and fear. The boundaries between cluster types are to some extent artificial, and many patients who meet criteria for one personality disorder also meet criteria for aspects of another. The risk of a comorbid major mental disorder is increased in patients who qualify for a diagnosis of personality disorder.

TREATMENT PERSONALITY DISORDERS

Dialectical behavior therapy (DBT) is a cognitive-behavioral approach that focuses on behavioral change while providing acceptance, compassion, and validation of the patient. Several randomized trials have demonstrated the efficacy of DBT in the treatment of personality disorders. Antidepressant medications and low-dose antipsychotic drugs have some efficacy in cluster A personality disorders, whereas anticonvulsant mood-stabilizing agents and MAOIs may be considered for patients with cluster B diagnoses who show marked mood reactivity, behavioral dyscontrol, and/or rejection hypersensitivity. Anxious or fearful cluster C patients often respond to medications used for axis I anxiety disorders (see above). It is important that the physician and the patient have reasonable expectations vis-à-vis the possible benefit of any medication used and its side effects. Improvement may be subtle and observable only over time.

SCHIZOPHRENIA

CLINICAL MANIFESTATIONS

Schizophrenia is a heterogeneous syndrome characterized by perturbations of language, perception, thinking, social activity, affect, and volition. There are no pathognomonic features. The syndrome commonly begins in late adolescence, has an insidious (and less commonly acute) onset, and, often, a poor outcome, progressing from social withdrawal and perceptual distortions to recurrent delusions and hallucinations. Patients may present with positive symptoms (such as conceptual disorganization, delusions, or hallucinations) or negative symptoms (loss of function, anhedonia, decreased emotional expression, impaired concentration, and diminished social engagement) and must have at least two of these for a 1-month period and continuous signs for at least 6 months to meet formal diagnostic criteria. Disorganized thinking or speech and grossly disorganized motor behavior, including catatonia, may also be present. As individuals age, positive psychotic symptoms tend to attenuate, and some measure of social and occupational function may be regained. “Negative” symptoms predominate in one-third of the schizophrenic population and are associated with

a poor long-term outcome and a poor response to drug treatment. However, marked variability in the course and individual character of symptoms is typical.

The term *schizophreniform disorder* describes patients who meet the symptom requirements but not the duration requirements for schizophrenia, and *schizoaffective disorder* is used for those who manifest symptoms of schizophrenia and independent periods of mood disturbance. The terms “schizotypal” and “schizoid” refer to specific personality disorders and are discussed in that section. The diagnosis of delusional disorder is used for individuals who have delusions of various content for at least 1 month but who otherwise do not meet criteria for schizophrenia. Patients who experience a sudden onset of a brief (<1 month) alteration in thought processing, characterized by delusions, hallucinations, disorganized speech, or gross motor behavior, are most appropriately designated as having a brief psychotic disorder. Catatonia is recognized as a nonspecific syndrome that can occur as a consequence of other severe psychiatric/medical disorders and is diagnosed by the documentation of three or more of a cluster of motor and behavioral symptoms, including stupor, cataplexy, mutism, waxy flexibility, and stereotypy, among others. Prognosis depends not on symptom severity but on the response to antipsychotic medication. A permanent remission without recurrence does occasionally occur. About 10% of schizophrenic patients commit suicide.

Schizophrenia is present in 0.85% of individuals worldwide, with a lifetime prevalence of ~1–1.5%. An estimated 300,000 episodes of acute schizophrenia occur annually in the United States, resulting in direct and indirect costs of \$62.7 billion.

DIFFERENTIAL DIAGNOSIS

The diagnosis is principally one of exclusion, requiring the absence of significant associated mood symptoms, any relevant medical condition, and substance abuse. Drug reactions that cause hallucinations, paranoia, confusion, or bizarre behavior may be dose-related or idiosyncratic; parkinsonian medications, clonidine, quinacrine, and procaine derivatives are the most common prescription medications associated with these symptoms. Drug causes should be ruled out in any case of newly emergent psychosis. The general neurologic examination in patients with schizophrenia is usually normal, but motor rigidity, tremor, and dyskinesias are noted in one-quarter of untreated patients.

EPIDEMIOLOGY AND PATHOPHYSIOLOGY

Epidemiologic surveys identify several risk factors for schizophrenia, including genetic susceptibility, early developmental insults, winter birth, and increasing parental age. Genetic factors are involved in at least a subset of individuals who develop schizophrenia. Schizophrenia is observed in ~6.6% of all first-degree relatives of an affected proband. If both parents are affected, the risk for offspring is 40%. The concordance rate for monozygotic twins is 50%, compared to 10% for dizygotic twins. Schizophrenia-prone families are also at risk for other psychiatric disorders, including schizoaffective disorder and *schizotypal* and *schizoid personality disorders*, the latter terms designating individuals who show a lifetime pattern of social and interpersonal deficits characterized by an inability to form close interpersonal relationships, eccentric behavior, and mild perceptual distortions.

TREATMENT SCHIZOPHRENIA

Antipsychotic agents (Table 466-10) are the cornerstone of acute and maintenance treatment of schizophrenia and are effective in the treatment of hallucinations, delusions, and thought disorders, regardless of etiology. The mechanism of action involves, at least in part, binding to dopamine D₂/D₃ receptors in the ventral striatum; the clinical potencies of traditional antipsychotic drugs parallel their affinities for the D₂ receptor, and even the newer “atypical” agents exert some degree of D₂ receptor blockade. All neuroleptics induce expression of the immediate-early gene *c-fos* in the nucleus accumbens, a dopaminergic site connecting prefrontal and limbic cortices. The clinical efficacy of newer atypical neuroleptics, however, may

involve *N*-methyl-D-aspartate (NMDA) receptor blockade, α₁- and α₂-noradrenergic activity, altering the relationship between 5-HT₂ and D₂ receptor activity, and faster dissociation of D₂ binding and effects on neuroplasticity.

Conventional neuroleptics differ in their potency and side effect profile. Older agents, such as chlorpromazine and thioridazine, are more sedating and anticholinergic and more likely to cause orthostatic hypotension, whereas higher potency antipsychotics, such as haloperidol, perphenazine, and thiothixene, are more likely to induce extrapyramidal side effects. The model “atypical” antipsychotic agent is *clozapine*, a dibenzodiazepine that has a greater potency in blocking the 5-HT₂ than the D₂ receptor and a much higher affinity for the D₄ than the D₂ receptor. Its principal disadvantage is a risk of blood dyscrasias. Paliperidone is a recently approved agent that is a metabolite of risperidone and shares many of its properties. Unlike other antipsychotics, clozapine does not cause a rise in prolactin level. Approximately 30% of patients who do not benefit from conventional antipsychotic agents will have a better response to this drug, which also has a demonstrated superiority to other antipsychotic agents in preventing suicide; however, its side effect profile makes it most appropriate for treatment-resistant cases. *Risperidone*, a benzisoxazole derivative, is more potent at 5-HT₂ than D₂ receptor sites, like clozapine, but it also exerts significant α₂ antagonism, a property that may contribute to its perceived ability to improve mood and increase motor activity. Risperidone is not as effective as clozapine in treatment-resistant cases but does not carry a risk of blood dyscrasias. *Olanzapine* is similar neurochemically to clozapine but has a significant risk of inducing weight gain. *Quetiapine* is distinct in having a weak D₂ effect but potent α₁ and histamine blockade. *Ziprasidone* causes minimal weight gain and is unlikely to increase prolactin but may increase QT prolongation. *Aripiprazole* also has little risk of weight gain or prolactin increase but may increase anxiety, nausea, and insomnia as a result of its partial agonist properties. Asenapine is associated with minimal weight gain and anticholinergic effect but may have a higher than expected risk of extrapyramidal symptoms.

Antipsychotic agents are effective in 70% of patients presenting with a first episode. Improvement may be observed within hours or days, but full remission usually requires 6–8 weeks. The choice of agent depends principally on the side effect profile and cost of treatment or on a past personal or family history of a favorable response to the drug in question. Atypical agents appear to be more effective in treating negative symptoms and improving cognitive function. An equivalent treatment response can usually be achieved with relatively low doses of any drug selected (i.e., 4–6 mg/d of haloperidol, 10–15 mg of olanzapine, or 4–6 mg/d of risperidone). Doses in this range result in >80% D₂ receptor blockade, and there is little evidence that higher doses increase either the rapidity or degree of response. Maintenance treatment requires careful attention to the possibility of relapse and monitoring for the development of a movement disorder. Intermittent drug treatment is less effective than regular dosing, but gradual dose reduction is likely to improve social functioning in many schizophrenic patients who have been maintained at high doses. If medications are completely discontinued, however, the relapse rate is 60% within 6 months. Long-acting injectable preparations (risperidone, paliperidone, olanzapine, aripiprazole) are considered when noncompliance with oral therapy leads to relapses but should not be considered interchangeable, because the agents differ in their indications, injection intervals and sites/volumes, and possible adverse reactions, among other factors. In treatment-resistant patients, a transition to clozapine usually results in rapid improvement, but a prolonged delay in response in some cases necessitates a 6- to 9-month trial for maximal benefit to occur.

Antipsychotic medications can cause a broad range of side effects, including lethargy, weight gain, postural hypotension, constipation, and dry mouth. Extrapyramidal symptoms such as dystonia, akathisia, and akinesia are also frequent with first-generation agents and may contribute to poor adherence if not specifically

Name	Usual PO Daily Dose, mg	Side Effects	Sedation	Comments
First-Generation Antipsychotics				
Low potency				
Chlorpromazine (Thorazine)	100–1000	Anticholinergic effects; orthostasis; photosensitivity; cholestasis; QT prolongation	+++	EPSEs usually not prominent; can cause anticholinergic delirium in elderly patients
Thioridazine (Mellaril)	100–600			
Midpotency				
Trifluoperazine (Stelazine)	2–50	Fewer anticholinergic side effects	++	Well tolerated by most patients
Perphenazine (Trilafon)	4–64	Fewer EPSEs than with higher potency agents	++	
Loxapine (Loxitane)	30–100	Frequent EPSEs	++	
Molindone (Moban)	30–100	Frequent EPSEs	0	Little weight gain
High potency				
Haloperidol (Haldol)	5–20	No anticholinergic side effects; EPSEs often prominent	0/+	Often prescribed in doses that are too high; long-acting injectable forms of haloperidol and fluphenazine available
Fluphenazine (Prolixin)	1–20	Frequent EPSEs	0/+	
Thiothixene (Navane)	2–50	Frequent EPSEs	0/+	
Second-Generation Antipsychotics				
Clozapine (Clozaril)	150–600	Agranulocytosis (1%); weight gain; seizures; drooling; hyperthermia	++	Requires weekly WBC count for first 6 months, then biweekly if stable
Risperidone (Risperdal)	2–8	Orthostasis	+	Requires slow titration; EPSEs observed with doses >6 mg qd
Olanzapine (Zyprexa)	10–30	Weight gain	++	Mild prolactin elevation
Quetiapine (Seroquel)	350–800	Sedation; weight gain; anxiety	+++	Bid dosing
Ziprasidone (Geodon)	120–200	Orthostatic hypotension	+ /+++	Minimal weight gain; increases QT interval
Aripiprazole (Abilify)	10–30	Nausea, anxiety, insomnia	0/+	Mixed agonist/antagonist
Paliperidone (Invega)	3–12	Restlessness, EPSEs, increased prolactin, headache	+	Active metabolite of risperidone
Iloperidone (Fanapt)	12–24	Dizziness, hypotension	0/+	Requires dose titration; long acting injectable available
Asenapine (Saphris)	10–20	Dizziness, anxiety, EPSEs, minimal weight gain	++	Sublingual tablets; bid dosing
Lurasidone (Latuda)	40–80	Nausea, EPSEs	++	Uses CYP3A4

Abbreviations: EPSEs, extrapyramidal side effects; WBC, white blood cell.

addressed. Anticholinergic and parkinsonian symptoms respond well to trihexyphenidyl, 2 mg bid, or bethtropine mesylate, 1–2 mg bid. Akathisia may respond to beta blockers. In rare cases, more serious and occasionally life-threatening side effects may emerge, including hyperprolactinemia, ventricular arrhythmias, gastrointestinal obstruction, retinal pigmentation, obstructive jaundice, and neuroleptic malignant syndrome (characterized by hyperthermia, autonomic dysfunction, muscular rigidity, and elevated creatine phosphokinase levels). The most serious adverse effects of clozapine are agranulocytosis, which has an incidence of 1%, and induction of seizures, which has an incidence of 10%. Weekly white blood cell counts are required, particularly during the first 3 months of treatment.

The risk of type 2 diabetes mellitus appears to be increased in schizophrenia, and second-generation agents as a group produce greater adverse effects on glucose regulation, independent of effects on obesity, than traditional agents. Clozapine, olanzapine, and quetiapine seem more likely to cause hyperglycemia, weight gain, and hypertriglyceridemia than other atypical antipsychotic drugs. Close monitoring of plasma glucose and lipid levels are indicated with the use of these agents.

A serious side effect of long-term use of first-generation antipsychotic agents is *tardive dyskinesia*, characterized by repetitive, involuntary, and potentially irreversible movements of the tongue and lips (bucco-linguo-masticatory triad) and, in approximately half of cases, choreoathetosis. Tardive dyskinesia has an incidence of 2–4% per year of exposure and a prevalence of 20% in chronically treated patients. The prevalence increases with age, total dose, and duration

of drug administration. The risk associated with second-generation agents appears to be much lower. The cause may involve formation of free radicals and perhaps mitochondrial energy failure. Vitamin E may reduce abnormal involuntary movements if given early in the syndrome.

The CATIE study, a large-scale investigation of the effectiveness of antipsychotic agents in “real-world” patients, revealed a high rate of discontinuation of treatment over 18 months. Olanzapine showed greater effectiveness than quetiapine, risperidone, perphenazine, or ziprasidone but also a higher discontinuation rate due to weight gain and metabolic effects. Surprisingly, perphenazine, a first-generation agent, showed little evidence of inferiority to newer drugs.

Drug treatment of schizophrenia is by itself insufficient. Educational efforts directed toward families and relevant community resources have proved to be necessary to maintain stability and optimize outcome. A treatment model involving a multidisciplinary case-management team that seeks out and closely follows the patient in the community has proved particularly effective.

ASSESSMENT AND EVALUATION OF VIOLENCE

Primary care physicians may encounter situations in which family, domestic, or societal violence is discovered or suspected. Such an awareness can carry legal and moral obligations; many state laws mandate reporting of child, spousal, and elder abuse. Physicians are frequently the first point of contact for both victim and abuser. Approximately 2 million older Americans and 1.5 million U.S. children are thought to experience some form of physical maltreatment

each year. Spousal abuse is thought to be even more prevalent. An interview study of 24,000 women in 10 countries found a lifetime prevalence of physical or sexual violence that ranged from 15 to 71%; these individuals are more likely to suffer from depression, anxiety, and substance abuse and to have attempted suicide. In addition, abused individuals frequently express low self-esteem, vague somatic symptomatology, social isolation, and a passive feeling of loss of control. Although it is essential to treat these elements in the victim, the first obligation is to ensure that the perpetrator has taken responsibility for preventing any further violence. Substance abuse and/or dependence and serious mental illness in the abuser may contribute to the risk of harm and require direct intervention. Depending on the situation, law enforcement agencies, community resources such as support groups and shelters, and individual and family counseling can be appropriate components of a treatment plan. A safety plan should be formulated with the victim, in addition to providing information about abuse, its likelihood of recurrence, and its tendency to increase in severity and frequency. Antianxiety and antidepressant medications may sometimes be useful in treating the acute symptoms, but only if independent evidence for an appropriate psychiatric diagnosis exists.

MENTAL HEALTH PROBLEMS IN THE HOMELESS

There is a high prevalence of mental disorders and substance abuse among homeless and impoverished individuals. Depending on the definition used, estimates of the total number of homeless individuals in the United States range from 800,000 to 2 million, one-third of whom qualify as having a serious mental disorder. Poor hygiene and nutrition, substance abuse, psychiatric illness, physical trauma, and exposure to the elements combine to make the provision of medical care challenging. Only a minority of these individuals receives formal mental health care; the main points of contact are outpatient medical clinics and emergency departments. Primary care settings represent a critical site in which housing needs, treatment of substance dependence, and evaluation and treatment of psychiatric illness can most efficiently take place. Successful intervention is dependent on breaking down traditional administrative barriers to health care and recognizing the physical constraints and emotional costs imposed by homelessness. Simplifying health care instructions and follow-up, allowing frequent visits, and dispensing medications in limited amounts that require ongoing contact are possible techniques for establishing a successful therapeutic relationship.

467 Alcohol and Alcoholism

Marc A. Schuckit

Alcohol (beverage ethanol) distributes throughout the body, affecting almost all systems and altering nearly every neurochemical process in the brain. This drug is likely to exacerbate most medical problems, affect medications metabolized in the liver, and temporarily mimic many medical (e.g., diabetes) and psychiatric (e.g., depression) conditions. The lifetime risk for repetitive alcohol problems is almost 20% for men and 10% for women, regardless of a person's education or income. Although low doses of alcohol might have healthful benefits, greater than three standard drinks per day enhances the risk for cancer and vascular disease, and alcohol use disorders decrease the life span by about 10 years. Unfortunately, most clinicians have had only limited training regarding alcohol-related disorders. This chapter presents a brief overview of clinically useful information about alcohol use and problems.

PHARMACOLOGY AND NUTRITIONAL IMPACT OF ETHANOL

Ethanol blood levels are expressed as milligrams or grams of ethanol per deciliter (e.g., 100 mg/dL = 0.10 g/dL), with values of ~0.02 g/dL

resulting from the ingestion of one typical drink. In round figures, a standard drink is 10–12 g, as seen in 340 mL (12 oz) of beer, 115 mL (4 oz) of nonfortified wine, and 43 mL (1.5 oz) (a shot) of 80-proof beverage (e.g., whisky); 0.5 L (1 pint) of 80-proof beverage contains ~160 g of ethanol (about 16 standard drinks), and 750 mL of wine contains ~60 g of ethanol. These beverages also have additional components (*congeners*) that affect the drink's taste and might contribute to adverse effects on the body. Congeners include methanol, butanol, acetaldehyde, histamine, tannins, iron, and lead. Alcohol acutely decreases neuronal activity and has similar behavioral effects and cross-tolerance with other depressants, including benzodiazepines and barbiturates.

Alcohol is absorbed from mucous membranes of the mouth and esophagus (in small amounts), from the stomach and large bowel (in modest amounts), and from the proximal portion of the small intestine (the major site). The rate of absorption is increased by rapid gastric emptying (as seen with carbonation); by the absence of proteins, fats, or carbohydrates (which interfere with absorption); and by dilution to a modest percentage of ethanol (maximum at ~20% by volume).

Between 2% (at low blood alcohol concentrations) and 10% (at high blood alcohol concentrations) of ethanol is excreted directly through the lungs, urine, or sweat, but most is metabolized to acetaldehyde, primarily in the liver. The most important pathway occurs in the cell cytosol where alcohol dehydrogenase (ADH) produces acetaldehyde, which is then rapidly destroyed by aldehyde dehydrogenase (ALDH) in the cytosol and mitochondria (Fig. 467-1). A second pathway occurs in the microsomes of the smooth endoplasmic reticulum (the microsomal ethanol-oxidizing system, or MEOS) that is responsible for ≥10% of ethanol oxidation at high blood alcohol concentrations.

Although a drink contains ~300 kJ, or 70–100 kcal, these are devoid of minerals, proteins, and vitamins. In addition, alcohol interferes with absorption of vitamins in the small intestine and decreases their storage in the liver with modest effects on folate (folacin or folic acid), pyridoxine (B₆), thiamine (B₁), nicotinic acid (niacin, B₃), and vitamin A.

Heavy drinking in a fasting, healthy individual can produce transient hypoglycemia within 6–36 h, secondary to the acute actions of ethanol on gluconeogenesis. This can result in temporary abnormal glucose tolerance tests (with a resulting erroneous diagnosis of diabetes mellitus) until the alcoholic has abstained for 2–4 weeks. Alcohol ketoacidosis, probably reflecting a decrease in fatty acid oxidation coupled with poor diet or recurrent vomiting, can be misdiagnosed as diabetic ketosis. With the former, patients show an increase in serum ketones along with a mild increase in glucose but a large anion gap, a mild to moderate increase in serum lactate, and a β-hydroxybutyrate/lactate ratio of between 2:1 and 9:1 (with normal being 1:1).

In the brain, alcohol affects almost all neurotransmitter systems, with acute effects that are often the opposite of those seen following desistance after a period of heavy drinking. The most prominent actions relate to boosting γ-aminobutyric acid (GABA) activity,

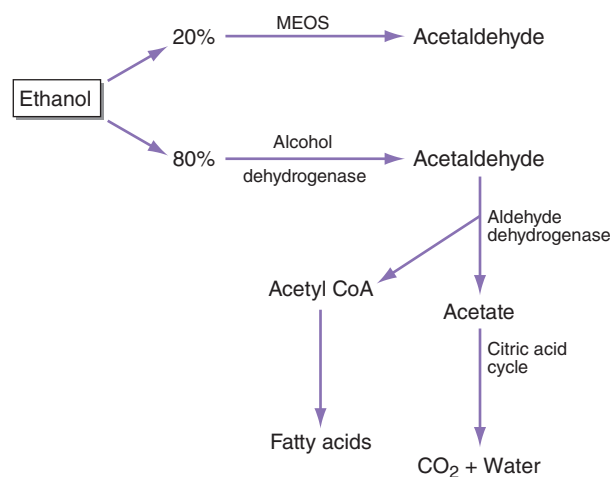


FIGURE 467-1 The metabolism of alcohol. CoA, coenzyme A; MEOS, microsomal ethanoloxidizing system.

especially at GABA_A receptors. Enhancement of this complex chloride channel system contributes to anticonvulsant, sleep-inducing, antianxiety, and muscle relaxation effects of all GABA-boosting drugs. Acutely administered alcohol produces a release of GABA, and continued use increases density of GABA_A receptors, whereas alcohol withdrawal states are characterized by decreases in GABA-related activity. Equally important is the ability of acute alcohol to inhibit postsynaptic *N*-methyl-D-aspartate (NMDA) excitatory glutamate receptors, whereas chronic drinking and desistance are associated with an upregulation of these excitatory receptor subunits. The relationships between greater GABA and diminished NMDA receptor activity during acute intoxication and diminished GABA with enhanced NMDA actions during alcohol withdrawal explain much of intoxication and withdrawal phenomena.

As with all pleasurable activities, alcohol acutely increases dopamine levels in the ventral tegmentum and related brain regions, and this effect plays an important role in continued alcohol use, craving, and relapse. The changes in dopamine pathways are also linked to increases in “stress hormones,” including cortisol and adrenocorticotropic hormone (ACTH) during intoxication and withdrawal. Such alterations are likely to contribute to both feelings of reward during intoxication and depression during falling blood alcohol concentrations. Also closely linked to alterations in dopamine (especially in the nucleus accumbens) are alcohol-induced changes in opioid receptors, with acute alcohol causing release of beta endorphins.

Additional neurochemical changes include increases in synaptic levels of serotonin during acute intoxication and subsequent upregulation of serotonin receptors. Acute increases in nicotinic acetylcholine systems contribute to the impact of alcohol in the ventral tegmental region, which occurs in concert with enhanced dopamine activity. In the same regions, alcohol impacts on cannabinol receptors, with resulting release of dopamine, GABA, and glutamate as well as subsequent effects on brain reward circuits.

BEHAVIORAL EFFECTS, TOLERANCE, AND WITHDRAWAL

The acute effects of a drug depend on the dose, the rate of increase in plasma, the concomitant presence of other drugs, and past experience with the agent. “Legal intoxication” with alcohol in most states requires a blood alcohol concentration of 0.08 g/dL, but levels of 0.04 are cited in some other countries. However, behavioral, psychomotor, and cognitive changes are seen at 0.02–0.04 g/dL (i.e., after one to two drinks) (Table 467-1). Deep but disturbed sleep can be seen at twice the legal intoxication level, and death can occur with levels between 0.30 and 0.40 g/dL. Beverage alcohol is probably responsible for more overdose deaths than any other drug.

Repeated use of alcohol contributes to acquired tolerance, a phenomenon involving at least three compensatory mechanisms. (1) After 1–2 weeks of daily drinking, *metabolic or pharmacokinetic tolerance* can be seen, with up to 30% increases in the rate of hepatic ethanol metabolism. This alteration disappears almost as rapidly as it develops. (2) *Cellular or pharmacodynamic tolerance* develops through neurochemical changes that maintain relatively normal physiologic functioning despite the presence of alcohol. Subsequent decreases in blood levels contribute to symptoms of withdrawal. (3) Individuals learn to adapt their behavior so that they can function better than expected under influence of the drug (*learned or behavioral tolerance*).

TABLE 467-1 EFFECTS OF BLOOD ALCOHOL LEVELS IN THE ABSENCE OF TOLERANCE

Blood Level, g/dL	Usual Effect
0.02	Decreased inhibitions, a slight feeling of intoxication
0.08	Decrease in complex cognitive functions and motor performance
0.20	Obvious slurred speech, motor incoordination, irritability, and poor judgment
0.30	Light coma and depressed vital signs
0.40	Death

The cellular changes caused by chronic ethanol exposure may not resolve for several weeks or longer following cessation of drinking. Rapid decreases in blood alcohol levels before that time can produce a withdrawal syndrome, which is most intense during the first 5 days, but some symptoms (e.g., disturbed sleep and anxiety) can take up to 4–6 months to resolve.

THE EFFECTS OF ETHANOL ON ORGAN SYSTEMS

Relatively low doses of alcohol (one or two drinks per day) have potential beneficial effects of increasing high-density lipoprotein cholesterol and decreasing aggregation of platelets, with a resulting decrease in risk for occlusive coronary disease and embolic strokes. Red wine has additional potential health-promoting qualities at relatively low doses due to flavinols and related substances. Modest drinking might also decrease the risk for vascular dementia and, possibly, Alzheimer’s disease. However, any potential healthful effects disappear with the regular consumption of three or more drinks per day, and knowledge about the deleterious effects of alcohol can both help the physician to identify patients with an alcohol use disorder and to supply them with information that might help motivate a change in behavior.

NERVOUS SYSTEM

Approximately 35% of drinkers (and a much higher proportion of alcoholics) experience a *blackout*, an episode of temporary anterograde amnesia, in which the person forgets all or part of what occurred during a drinking evening. Another common problem, one seen after as few as one or two drinks shortly before bedtime, is disturbed sleep. Although alcohol might initially help a person fall asleep, it disrupts sleep throughout the rest of the night. The stages of sleep are altered, and time spent in rapid eye movement (REM) and deep sleep is reduced. Alcohol relaxes muscles in the pharynx, which can cause snoring and exacerbate sleep apnea; symptoms of the latter occur in 75% of alcoholic men older than age 60 years. Patients may also experience prominent and sometimes disturbing dreams later in the night. All of these sleep problems are more pronounced in alcoholics, and their persistence may contribute to relapse.

Another common consequence of alcohol use is impaired judgment and coordination, increasing the risk of injury. In the United States, ~40% of drinkers have at some time driven while intoxicated. Heavy drinking can also be associated with headache, thirst, nausea, vomiting, and fatigue the following day, a hangover syndrome that is responsible for much missed time and temporary cognitive deficits at work and school.

Chronic high doses cause *peripheral neuropathy* in ~10% of alcoholics: similar to diabetes, patients experience bilateral limb numbness, tingling, and paresthesias, all of which are more pronounced distally. Approximately 1% of alcoholics develop *cerebellar degeneration or atrophy*, producing a syndrome of progressive unsteady stance and gait often accompanied by mild nystagmus; neuroimaging studies reveal atrophy of the cerebellar vermis. Fortunately, very few alcoholics (perhaps as few as 1 in 500 for the full syndrome) develop *Wernicke’s* (ophthalmoparesis, ataxia, and encephalopathy) and *Korsakoff’s* (retrograde and anterograde amnesia) *syndromes*, although a higher proportion have one or more neuropathologic findings related to these conditions. These result from low levels of thiamine, especially in predisposed individuals with transketolase deficiencies. Alcoholics can manifest *cognitive problems* and temporary memory impairment lasting for weeks to months after drinking heavily for days or weeks. Brain atrophy, evident as ventricular enlargement and widened cortical sulci on magnetic resonance imaging (MRI) and computed tomography (CT) scans, occurs in ~50% of chronic alcoholics; these changes are usually reversible if abstinence is maintained. There is no single alcoholic dementia syndrome; rather, this label describes patients who have irreversible cognitive changes (possibly from diverse causes) in the context of chronic alcoholism.

Psychiatric Comorbidity As many as two-thirds of individuals with alcohol use disorders meet the criteria for another psychiatric syndrome in the fifth edition of the *Diagnostic and Statistical Manual of Mental Disorders* (DSM-5) of the American Psychiatric Association

(Chap. 466). Half of these relate to a preexisting antisocial personality manifesting as impulsivity and disinhibition that contribute to both alcohol and drug use disorders. The lifetime risk is 3% in males, and $\geq 80\%$ of such individuals demonstrate alcohol- and/or drug-related conditions. Another common comorbidity occurs with problems regarding illicit substances. The remainder of alcoholics with psychiatric syndromes have preexisting conditions such as schizophrenia or manic-depressive disease and anxiety syndromes such as panic disorder. The comorbidities of alcoholism with independent psychiatric disorders might represent an overlap in genetic vulnerabilities, impaired judgment in the use of alcohol from the independent psychiatric condition, or an attempt to use alcohol to alleviate symptoms of the disorder or side effects of medications.

Many psychiatric syndromes can be seen *temporarily* during heavy drinking and subsequent withdrawal. These alcohol-induced conditions include an intense *sadness* lasting for days to weeks in the midst of heavy drinking seen in 40% of alcoholics, which tends to disappear over several weeks of abstinence (alcohol-induced mood disorder); temporary severe *anxiety* in 10–30% of alcoholics, often beginning during alcohol withdrawal, which can persist for a month or more after cessation of drinking (alcohol-induced anxiety disorder); and auditory *hallucinations* and/or paranoid delusions in a person who is alert and oriented, seen in 3–5% of alcoholics (*alcohol-induced psychotic disorder*).

Treatment of all forms of alcohol-induced psychopathology includes helping patients achieve abstinence and offering supportive care, as well as reassurance and “talk therapy” such as cognitive-behavioral approaches. However, with the exception of short-term antipsychotics or similar drugs for substance-induced psychoses, substance-induced psychiatric conditions only rarely require medications. Recovery is likely within several days to 4 weeks of abstinence. Conversely, because alcohol-induced conditions are temporary and do not indicate a need for long-term pharmacotherapy, a history of alcohol intake is an important part of the workup for any patient with one of these psychiatric symptoms.

THE GASTROINTESTINAL SYSTEM

Esophagus and Stomach Alcohol can cause inflammation of the esophagus and stomach causing epigastric distress and gastrointestinal bleeding, making alcohol one of the most common causes of hemorrhagic gastritis. Violent vomiting can produce severe bleeding through a Mallory-Weiss lesion, a longitudinal tear in the mucosa at the gastroesophageal junction.

Pancreas and Liver The incidence of acute pancreatitis (~25 per 1000 per year) is almost threefold higher in alcoholics than in the general population, accounting for an estimated 10% or more of the total cases. Alcohol impairs gluconeogenesis in the liver, resulting in a fall in the amount of glucose produced from glycogen, increased lactate production, and decreased oxidation of fatty acids. This contributes to an increase in fat accumulation in liver cells. In healthy individuals these changes are reversible, but with repeated exposure to ethanol, especially daily heavy drinking, more severe changes in the liver occur, including alcohol-induced hepatitis, perivenular sclerosis, and cirrhosis, with the latter observed in an estimated 15% of alcoholics (Chap. 363). Perhaps through an enhanced vulnerability to infections, alcoholics have an elevated rate of hepatitis C, and drinking in the context of that disease is associated with more severe liver deterioration.

CANCER

As few as 1.5 drinks per day increases a woman’s risk of breast cancer 1.4-fold. For both genders, four drinks per day increases the risk for oral and esophageal cancers approximately threefold and rectal cancers by a factor of 1.5; seven to eight or more drinks per day produces an approximately fivefold increased risk for many cancers. These consequences may result directly from cancer-promoting effects of alcohol and acetaldehyde or indirectly by interfering with immune homeostasis.

HEMATOPOIETIC SYSTEM

Ethanol causes an increase in red blood cell size (mean corpuscular volume [MCV]), which reflects its effects on stem cells. If heavy

drinking is accompanied by folic acid deficiency, there can also be hypersegmented neutrophils, reticulocytopenia, and a hyperplastic bone marrow; if malnutrition is present, sideroblastic changes can be observed. Chronic heavy drinking can decrease production of white blood cells, decrease granulocyte mobility and adherence, and impair delayed-hypersensitivity responses to novel antigens (with a possible false-negative tuberculin skin test). Associated immune deficiencies can contribute to vulnerability toward infections, including hepatitis and HIV, and interfere with their treatment. Finally, many alcoholics have mild thrombocytopenia, which usually resolves within a week of abstinence unless there is hepatic cirrhosis or congestive splenomegaly.

CARDIOVASCULAR SYSTEM

Acutely, ethanol decreases myocardial contractility and causes peripheral vasodilation, with a resulting mild decrease in blood pressure and a compensatory increase in cardiac output. Exercise-induced increases in cardiac oxygen consumption are higher after alcohol intake. These acute effects have little clinical significance for the average healthy drinker but can be problematic when persisting cardiac disease is present.

The consumption of three or more drinks per day results in a dose-dependent increase in blood pressure, which returns to normal within weeks of abstinence. Thus, heavy drinking is an important factor in mild to moderate hypertension. Chronic heavy drinkers also have a sixfold increased risk for coronary artery disease, related, in part, to increased low-density lipoprotein cholesterol, and carry an increased risk for cardiomyopathy through direct effects of alcohol on heart muscle. Symptoms of the latter include unexplained arrhythmias in the presence of left ventricular impairment, heart failure, hypocontractility of heart muscle, and dilation of all four heart chambers with associated mural thrombi and mitral valve regurgitation. Atrial or ventricular arrhythmias, especially paroxysmal tachycardia, can also occur temporarily after heavy drinking in individuals showing no other evidence of heart disease—a syndrome known as the “holiday heart.”

GENITOURINARY SYSTEM CHANGES, SEXUAL FUNCTIONING, AND FETAL DEVELOPMENT

Drinking in adolescence can affect normal sexual development and reproductive onset. At any age, modest ethanol doses (e.g., blood alcohol concentrations of 0.06 g/dL) can increase sexual drive but also decrease erectile capacity in men. Even in the absence of liver impairment, a significant minority of chronic alcoholic men show irreversible testicular atrophy with shrinkage of the seminiferous tubules, decreases in ejaculate volume, and a lower sperm count (Chap. 411).

The repeated ingestion of high doses of ethanol by women can result in amenorrhea, a decrease in ovarian size, absence of corpora lutea with associated infertility, and an increased risk of spontaneous abortion. Heavy drinking during pregnancy results in the rapid placental transfer of both ethanol and acetaldehyde, which may contribute to a range of consequences known as fetal alcohol spectrum disorder (FASD). One severe result is the *fetal alcohol syndrome* (FAS), seen in ~5% of children born to heavy-drinking mothers, which can include any of the following: facial changes with epicanthal eye folds; poorly formed ear concha; small teeth with faulty enamel; cardiac atrial or ventricular septal defects; an aberrant palmar crease and limitation in joint movement; and microcephaly with mental retardation. Less pervasive FASD conditions include combinations of low birth weight, a lower intelligence quotient (IQ), hyperactive behavior, and some modest cognitive deficits. The amount of ethanol required and the time of vulnerability during pregnancy have not been defined, making it advisable for pregnant women to abstain completely.

OTHER EFFECTS

Between one-half and two-thirds of alcoholics have skeletal muscle weakness caused by acute *alcoholic myopathy*, a condition that improves but which might not fully remit with abstinence. Effects of repeated heavy drinking on the *skeletal system* include changes in calcium metabolism, lower bone density, and decreased growth in the epiphyses, leading to an increased risk for fractures and osteonecrosis of the femoral head. *Hormonal changes* include an increase in cortisol

levels, which can remain elevated during heavy drinking; inhibition of vasopressin secretion at rising blood alcohol concentrations and enhanced secretion at falling blood alcohol concentrations (with the final result that most alcoholics are likely to be slightly overhydrated); a modest and reversible decrease in serum thyroxine (T_4); and a more marked decrease in serum triiodothyronine (T_3). Hormone irregularities should be reevaluated because they may disappear after a month of abstinence.

ALCOHOLISM (ALCOHOL USE DISORDER)

Because many drinkers occasionally imbibe to excess, temporary alcohol-related problems are common in nonalcoholics, especially in the late teens to the late twenties. However, repeated problems in multiple life areas can indicate an alcohol use disorder as defined in DSM-5.

DEFINITIONS AND EPIDEMIOLOGY

An *alcohol use disorder* is defined as repeated alcohol-related difficulties in at least 2 of 11 life areas that cluster together in the same 12-month period (Table 467-2). Ten of the 11 items were taken directly from the 7 dependence and 4 abuse criteria in DSM-IV, after deleting legal problems and adding craving. Severity of an alcohol use disorder is based on the number of items endorsed: mild is two or three items; moderate is four or five; and severe is six or more of the criterion items. The new diagnostic approach is similar enough to DSM-IV that the following descriptions of associated phenomena are still accurate.

The lifetime risk for an alcohol use disorder in most Western countries is about 10–15% for men and 5–8% for women. Rates are similar in the United States, Canada, Germany, Australia, and the United Kingdom, tend to be lower in most Mediterranean countries, such as Italy, Greece, and Israel, and may be higher in Ireland, France, and Scandinavia. An even higher lifetime prevalence has been reported for most native cultures, including American Indians, Eskimos, Maori groups, and aboriginal tribes of Australia. These differences reflect both cultural and genetic influences, as described below. In Western countries, the typical alcoholic is more often a blue- or white-collar worker or homemaker. The lifetime risk for alcoholism among physicians is similar to that of the general population.

GENETICS

Approximately 60% of the risk for alcohol use disorders is attributed to genes, as indicated by the fourfold higher risk in children of alcoholics (even if adopted early in life and raised by nonalcoholics) and a higher risk in identical twins compared to fraternal twins of alcoholics. The genetic variations operate primarily through intermediate

characteristics that subsequently combine with environmental influences to alter the risk for heavy drinking and alcohol problems. These include genes relating to a high risk for all substance use disorders that operate through impulsivity, schizophrenia, and bipolar disorder. Another characteristic, an intense flushing response when drinking, decreases the risk for only alcohol use disorders through gene variations for several alcohol-metabolizing enzymes, especially aldehyde dehydrogenase (a mutation only seen in Asians), and to a lesser extent, variations in ADH.

An additional genetically influenced characteristic, a low sensitivity to alcohol, affects the risk for heavy drinking and may operate, in part, through variations in genes relating to calcium and potassium channels, GABA, nicotinic, and serotonin systems. A low response per drink is observed early in the drinking career and before alcohol use disorders develop. All follow-up studies have demonstrated that this need for higher doses of alcohol to achieve effects predicts future heavy drinking, alcohol problems, and alcohol use disorders. The impact of a low response to alcohol on adverse drinking outcomes is partially mediated by a range of environmental influences, including the selection of heavier-drinking friends, more positive expectations of the effects of high doses of alcohol, and suboptimal ways of coping with stress.

NATURAL HISTORY

Although the age of the first drink (~15 years) is similar in most alcoholics and nonalcoholics, a slightly earlier onset of regular drinking and drunkenness, especially in the context of conduct problems, is associated with a higher risk for later alcohol use disorders. By the mid-twenties, most nonalcoholic men and women moderate their drinking (perhaps learning from problems), whereas alcoholics are likely to escalate their patterns of drinking despite difficulties. The first major life problem from alcohol often appears in the late teens to early twenties, and a pattern of multiple alcohol difficulties by the midtwenties. Once established, the course of alcoholism is likely to include exacerbations and remissions, with little difficulty in temporarily stopping or controlling alcohol use when problems develop, but without help, desistance usually gives way to escalations in alcohol intake and subsequent problems. Following treatment, between half and two-thirds of alcoholics maintain abstinence for years, and often permanently. Even without formal treatment or self-help groups, there is at least a 20% chance of spontaneous remission with long-term abstinence. However, should the alcoholic continue to drink heavily, the life span is shortened by ~10 years on average, with the leading causes of death being heart disease, cancer, accidents, and suicide.

TREATMENT

The approach to treating alcohol-related conditions is relatively straightforward: (1) recognize that at least 20% of all patients have an alcohol use disorder; (2) learn how to identify and treat acute alcohol-related conditions; (3) know how to help patients begin to address their alcohol problems; and (4) know enough about treating alcoholism to appropriately refer patients for additional help.

IDENTIFICATION OF THE ALCOHOLIC

Even in affluent locales, ~20% of patients have an alcohol use disorder. These men and women can be identified by asking questions about *alcohol problems* and noting laboratory test results that can reflect regular consumption of six to eight or more drinks per day. The two blood tests with ≥60% sensitivity and specificity for heavy alcohol consumption are γ -glutamyl transferase (GGT) (>35 U) and carbohydrate-deficient transferrin (CDT) (>20 U/L or >2.6%); the combination of the two is likely to be more accurate than either alone. The values for these serologic markers are likely to return toward normal within several weeks of abstinence. Other useful blood tests include high-normal MCVs (≥91 μm^3) and serum uric acid (>416 mol/L, or 7 mg/dL).

The diagnosis of an alcohol use disorder ultimately rests on the documentation of a pattern of repeated difficulties associated with alcohol (Table 467-2). Thus, in screening, it is important to probe for marital or job problems, legal difficulties, histories of accidents, medical problems, evidence of tolerance, and so on, and then attempt to tie in use

TABLE 467-2 DIAGNOSTIC AND STATISTICAL MANUAL OF MENTAL DISORDERS, FIFTH EDITION, CLASSIFICATION OF ALCOHOL USE DISORDER (AUD)

Criteria

Two or more of the following items occurring in the same 12-month period must be endorsed for the diagnosis of an alcohol use disorder^a:

- Drinking resulting in recurrent failure to fulfill role obligations
- Recurrent drinking in hazardous situations
- Continued drinking despite alcohol-related social or interpersonal problems
- Tolerance
- Withdrawal, or substance use for relief/avoidance of withdrawal
- Drinking in larger amounts or for longer than intended
- Persistent desire/unsuccessful attempts to stop or reduce drinking
- Great deal of time spent obtaining, using, or recovering from alcohol
- Important activities given up/reduced because of drinking
- Continued drinking despite knowledge of physical or psychological problems caused by alcohol
- Alcohol craving

^aMild AUD: 2–3 criteria required; Moderate AUD: 4–5 items endorsed; severe AUD: 6 or more items endorsed.

TABLE 467-3 THE ALCOHOL USE DISORDERS IDENTIFICATION TEST (AUDIT)^a

Item	5-Point Scale (Least to Most)
1. How often do you have a drink containing alcohol?	Never (0) to 4+ per week (4)
2. How many drinks containing alcohol do you have on a typical day?	1 or 2 (0) to 10+ (4)
3. How often do you have six or more drinks on one occasion?	Never (0) to daily or almost daily (4)
4. How often during the last year have you found that you were not able to stop drinking once you had started?	Never (0) to daily or almost daily (4)
5. How often during the last year have you failed to do what was normally expected from you because of drinking?	Never (0) to daily or almost daily (4)
6. How often during the last year have you needed a first drink in the morning to get yourself going after a heavy drinking session?	Never (0) to daily or almost daily (4)
7. How often during the last year have you had a feeling of guilt or remorse after drinking?	Never (0) to daily or almost daily (4)
8. How often during the last year have you been unable to remember what happened the night before because you had been drinking?	Never (0) to daily or almost daily (4)
9. Have you or someone else been injured as a result of your drinking?	No (0) to yes, during the last year (4)
10. Has a relative, friend, doctor or other health worker been concerned about your drinking or suggested that you should cut down?	No (0) to yes, during the last year (4)

^aThe AUDIT is scored by simply summing the values associated with the endorsed response. A score ≥ 8 may indicate harmful alcohol use.

of alcohol or another substance. Some standardized questionnaires can be helpful, including the 10-item Alcohol Use Disorders Identification Test (AUDIT) (Table 467-3), but these are only screening tools, and a face-to-face interview is still required for a meaningful diagnosis.

TREATMENT ALCOHOL-RELATED CONDITIONS

ACUTE INTOXICATION

The first priority in treating severe intoxication is to assess vital signs and manage respiratory depression, cardiac arrhythmias, or blood pressure instability, if present. The possibility of intoxication with other drugs should be considered by obtaining toxicology screens for other central nervous system (CNS) depressants such as benzodiazepines and for opioids. Aggressive behavior should be handled by offering reassurance but also by considering a possible show of force with an intervention team. If the aggressive behavior continues, relatively low doses of a short-acting benzodiazepine such as lorazepam (e.g., 1–2 mg PO or IV) may be used and can be repeated as needed, but care must be taken not to destabilize vital signs or worsen confusion. An alternative approach is to use an antipsychotic medication (e.g., 0.5–5 mg of haloperidol PO or IM every 4–8 h as needed, or olanzapine 2.5–10 mg IM repeated at 2 and 6 h, if needed).

INTERVENTION

There are two main elements to intervention in a person with alcoholism: motivational interviewing and brief interventions. During motivational interviewing, the clinician helps the patient to think through the assets (e.g., comfort in social situations) and liabilities (e.g., health- and interpersonal-related problems) of the current pattern of drinking. The patient's responses are key, and the clinician should listen empathetically, helping to weigh options and encouraging the patient to take responsibility for needed changes. Patients should be reminded that only they can decide to avoid

the consequences that will occur without changes in drinking. The process of brief intervention has been summarized by the acronym FRAMES: Feedback to the patient; Responsibility to be taken by the patient; Advice, rather than orders, on what needs to be done; Menus of options that might be considered; Empathy for understanding the patient's thoughts and feelings; and Self-efficacy, i.e., offering support for the capacity of the patient to make changes.

Once the patient begins to consider change, the emphasis shifts to brief interventions designed to help them understand more about potential actions. Discussions focus on consequences of high alcohol consumption, suggested approaches to stopping drinking, and help in recognizing and avoiding situations likely to lead to heavy drinking. Both motivational interviewing and brief interventions can be carried out in 15-min sessions, but because patients do not always change behavior immediately, multiple meetings are often required to explain the problem, discuss optimal treatments, and explain the benefits of abstinence.

ALCOHOL WITHDRAWAL

If the patient agrees to stop drinking, sudden decreases in alcohol intake can produce withdrawal symptoms, many of which are the opposite of those produced by intoxication. Features include tremor of the hands (shakes); agitation and anxiety; autonomic nervous system overactivity including an increase in pulse, respiratory rate, sweating, and body temperature; and insomnia. These symptoms usually begin within 5–10 h of decreasing ethanol intake, peak on day 2 or 3, and improve by day 4 or 5, although mild levels of these problems may persist for 4–6 months as a protracted abstinence syndrome.

About 2% of alcoholics experience a withdrawal seizure, with the risk increasing in the context of concomitant medical problems, misuse of additional drugs, and higher alcohol quantities. The same risk factors also contribute to a similar rate of *delirium tremens* (DTs), where the withdrawal includes delirium (mental confusion, agitation, and fluctuating levels of consciousness) associated with a tremor and autonomic overactivity (e.g., marked increases in pulse, blood pressure, and respirations). The risks for seizures and DTs can be diminished by identifying and treating any underlying medical conditions early in the course of withdrawal.

Thus, the first step is a thorough physical examination in all alcoholics considering abstinence, including a search for evidence of liver failure, gastrointestinal bleeding, cardiac arrhythmia, infection, and glucose or electrolyte imbalances. It is also important to offer adequate nutrition and oral multiple B vitamins, including 50–100 mg of thiamine daily for a week or more. Because most alcoholics who enter withdrawal are either normally hydrated or mildly overhydrated, IV fluids should be avoided unless there is a relevant medical problem or significant recent bleeding, vomiting, or diarrhea.

The next step is to recognize that because withdrawal symptoms reflect the rapid removal of a CNS depressant, alcohol, the symptoms can be controlled by administering any depressant in doses that decrease symptoms (e.g., a rapid pulse and tremor) and then tapering the dose over 3–5 days. Although most depressants are effective, benzodiazepines (Chap. 466) have the highest margin of safety and lowest cost and are, therefore, the preferred class of drugs. Short-half-life benzodiazepines can be considered for patients with serious liver impairment or evidence of significant brain damage, but they must be given every 4 h to avoid abrupt blood-level fluctuations that may increase the risk for seizures. Therefore, most clinicians use drugs with longer half-lives (e.g., chlorthalidoxepoxide), adjusting the dose if signs of withdrawal escalate, and withholding the drug if the patient is sleeping or has orthostatic hypotension. The average patient requires 25–50 mg of chlorthalidoxepoxide or 10 mg of diazepam given PO every 4–6 h on the first day, with doses then decreased to zero over the next 5 days. Although alcohol withdrawal can be treated in a hospital, patients in good physical condition who demonstrate mild signs of withdrawal despite low blood alcohol concentrations and who have no prior history of DTs or withdrawal seizures can be considered for outpatient detoxification. For the next

2728 4 or 5 days, these patients should return daily for evaluation of vital signs and can be hospitalized if signs and symptoms of withdrawal escalate.

Treatment of patient with DTs can be challenging, and the condition is likely to run a course of 3–5 days regardless of the therapy used. The focus of care is to identify and correct medical problems and to control behavior and prevent injuries. Many clinicians recommend the use of high doses of a benzodiazepine (as much as 800 mg/d of chlordiazepoxide has been reported), a treatment that will decrease agitation and raise the seizure threshold but probably does little to improve the confusion. Other clinicians recommend the use of antipsychotic medications, such as haloperidol or olanzapine as discussed above, although these drugs have not been directly evaluated for DTs. Antipsychotics are less likely to exacerbate confusion but may increase the risk of seizures; they have no place in the treatment of mild withdrawal symptoms.

Generalized withdrawal seizures rarely require more than giving an adequate dose of benzodiazepines. There is little evidence that anticonvulsants such as phenytoin or gabapentin are more effective in drug-withdrawal seizures, and the risk of seizures has usually passed by the time effective drug levels are reached. The rare patient with status epilepticus must be treated aggressively (Chap. 445).

REHABILITATION OF ALCOHOLICS

An Overview After completing alcoholic rehabilitation, ≥60% of alcoholics, especially middle-class patients, maintain abstinence for at least a year, and many achieve lifetime sobriety. The core of treatment uses cognitive-behavioral approaches to help patients recognize the need to change, while working with them to alter their behaviors to enhance compliance. A key step is to optimize motivation toward abstinence through education about alcoholism and instructions to family members to stop protecting the patient from problems caused by alcohol. After years of heavy drinking, many patients also need counseling, some require vocational or avocational help to structure their days, and all should try self-help groups such as Alcoholics Anonymous (AA) to help them develop a sober peer group and learn how to deal with life's stresses while sober. A third component, *relapse prevention*, helps the patient identify situations in which a return to drinking is likely, formulate ways of managing these risks, and develop coping strategies that increase the chances of a return to abstinence if a slip occurs.

Although many can be treated as outpatients, more intense interventions are more effective, and some alcoholics do not respond to AA or outpatient groups. Whatever the setting, subsequent contact with outpatient treatment staff should be maintained for at least 6 months and preferably a year after abstinence. Counseling focuses on areas of improved functioning in the absence of alcohol (i.e., why it is a good idea to continue abstinence) and helping the patient to manage free time without alcohol, develop a nondrinking peer group, and handle stresses.

The physician serves an important role in identifying the alcoholic, diagnosing and treating associated medical and psychiatric syndromes, overseeing detoxification, referring the patient to rehabilitation programs, providing counseling, and, if appropriate, selecting which (if any) medication might be needed. For insomnia, patients should be reassured that troubled sleep is normal after alcohol withdrawal and will improve over subsequent weeks. They should be taught the elements of "sleep hygiene" including maintaining consistent schedules for bedtime and awakening. Sleep medications have the danger of being misused and of rebound insomnia when stopped. Sedating antidepressants (e.g., trazodone) should not be used because they interfere with cognitive functioning the next morning and disturb the normal sleep architecture, but occasional use of over-the-counter sleeping medications (sedating antihistamines) can be considered. Anxiety can be addressed by increasing the patient's insight into the temporary nature of the symptoms and helping the patient to develop strategies to achieve relaxation by using forms of cognitive therapy.

Medications for Rehabilitation Several medications have modest benefits when used for the first 6 months of recovery. The opioid antagonist, naltrexone, 50–150 mg/d orally, may shorten subsequent relapses, whether used in the oral form or as a once-per-month 380-mg injection, especially in individuals with the G allele of the A118G polymorphism of the μ opioid receptor. By blocking opioid receptors, naltrexone decreases activity in the dopamine-rich ventral tegmental reward system and decreases the feeling of pleasure if alcohol is imbibed. A second medication, acamprosate (Campral) at ~2 g/d divided into three oral doses, has similar modest effects; acamprosate inhibits NMDA receptors, decreasing mild symptoms of protracted withdrawal. Several trials of combined naltrexone and acamprosate have reported that the combination may be superior to either drug alone, although not all studies agree.

It is more difficult to establish the asset-to-liability ratio of a third drug, disulfiram, an ALDH inhibitor, used at doses of 250 mg/d. This drug produces vomiting and autonomic nervous system instability in the presence of alcohol as a result of rapidly rising blood levels of acetaldehyde. This reaction can be dangerous, especially for patients with heart disease, stroke, diabetes mellitus, or hypertension. The drug itself carries potential risks of depression, psychotic symptoms, peripheral neuropathy, and liver damage. Disulfiram is best given under supervision by someone (such as a spouse), especially during high-risk drinking situations (such as the Christmas holiday). Other drugs under investigation include another opioid antagonist nalmefene, the nicotinic receptor agonist varenicline, the serotonin antagonist ondansetron, the α -adrenergic agonist prazosin, the GABA_B receptor agonist baclofen, the anticonvulsant topiramate, and cannabinol receptor antagonists. At present, there are insufficient data to determine the asset-to-liability ratio for these medications in treating alcoholism and, therefore, no data to offer solid support for their use in routine clinical settings.

GLOBAL CONSIDERATIONS



As described above, rates of alcohol use disorders differ across sex, age, ethnicity, and country. There are also differences across countries regarding the definition of a standard drink (e.g., 10–12 g of ethanol in the United States and 8 g in the United Kingdom) and the definition of being legally drunk. The preferred alcoholic beverage also varies across groups, even within countries. That said, regardless of sex, ethnicity, or country, the actual drug in the drink is still ethanol, and the risks for problems, course of alcohol use disorders, and approaches to treatment are similar across the world.

468e Opioid-Related Disorders

Thomas R. Kosten, Colin N. Haile

This is a digital-only chapter. It is available on the DVD that accompanies this book, as well as on Access Medicine/Harrison's Online, and the eBook and "app" editions of HPIM 19e.



Opiate analgesics have been abused since at least 300 B.C. Nipenthe (Greek "free from sorrow") helped the hero of the *Odyssey*, but widespread opium smoking in China and the Near East has caused harm for centuries. Since the first chemical isolation of opium and codeine 200 years ago, a wide range of synthetic opioids have been developed, and opioid receptors were cloned in the 1990s. Two of the most important adverse effects of all these agents are the development of opioid use disorder and overdose. The 0.1% annual prevalence of heroin dependence in the United States is only about one-third the rate of prescription opiate use and is substantially lower than the 2% rate of morphine users in Southeast and Southwest Asia.

468e Opioid-Related Disorders

Thomas R. Kosten, Colin N. Haile



Opiate analgesics have been abused since at least 300 B.C. Nephenthe (Greek “free from sorrow”) helped the hero of the *Odyssey*, but widespread opium smoking in China and the Near East has caused harm for centuries. Since the first chemical isolation of opium and codeine 200 years ago, a wide range of synthetic opioids have been developed, and opioid receptors were cloned in the 1990s. Two of the most important adverse effects of all these agents are the development of opioid use disorder and overdose. The 0.1% annual prevalence of heroin dependence in the United States is only about one-third the rate of prescription opiate use and is substantially lower than the 2% rate of morphine users in Southeast and Southwest Asia. Prescription opiates are primarily used for pain management, but due to ease of availability, adolescents procure and use these drugs with dire consequences. In 2011, for example, 11 million individuals in the United States used nonmedically prescribed pain killers that were linked to over 420,000 emergency department visits and nearly 17,000 overdose deaths. Although these rates are low relative to other abused substances, their disease burden is substantial, with high rates of morbidity and mortality; disease transmission; increased health care, crime, and law enforcement costs; and less tangible costs of family distress and lost productivity.

The terms “dependence” and “addiction” are no longer used to describe substance use disorders. Opioid-related disorders encompass opioid use disorder, opioid intoxication, and opioid withdrawal. The diagnosis of opioid use disorder as defined in the *Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition* (DSM-5) requires the repeated use of the opiate while producing problems in two or more areas in a 12-month period. The areas include tolerance, withdrawal, use of greater amounts of opiates than intended, craving, and use despite adverse consequences. This new definition of opiate use disorder, reducing the criteria for diagnosis from three problem areas to two, is not expected to change the rates of these disorders because most individuals using these substances meet more than three criteria.

A striking recent aspect of illicit opiate use has been its marked increase as the gateway to illicit drugs in the United States. Since 2007, prescription opiates have surpassed marijuana as the most common illicit drug that adolescents initially use, although overall rates of opiate dependence are far lower than marijuana. The most commonly used opiates are diverted prescriptions for oxycodone and hydrocodone, followed by heroin and morphine, and—among health professionals—meperidine and fentanyl. Heroin is derived from morphine and acts as a prodrug that more readily penetrates the brain and is converted rapidly to morphine in the body. Two opiate maintenance treatment agents—methadone and buprenorphine—are also misused, but at substantially lower rates, and the partial opiate agonists such as butorphanol, tramadol, and pentazocine are misused even less frequently. Because the chemistry and general pharmacology of these agents are covered in major pharmacology texts, this chapter focuses on the neurobiology and pharmacology relevant to dependence and its treatments. Although the neurobiology of abuse involves all four of the known opiate receptors—mu, kappa, delta, and nociceptin/orphanin—this discussion focuses on the mu receptor, at which most of the clinically used opiates are active.

NEUROBIOLOGY

The neurobiology of opiates and their effects not only include opiate receptors, but also the downstream intracellular messenger systems and ion channels that the receptors regulate. The different functional activities of opiate receptors are summarized in [Table 468e-1](#). Abuse liability of opiates is primarily associated with the mu receptor. All opiate receptors are G protein-linked and coupled to the cyclic adenosine monophosphate (cAMP) second messenger system and to G protein-coupled, inwardly rectifying potassium channels (GIRKs). Opiates activate GIRKs, increasing permeability to potassium ions to cause

TABLE 468e-1 ACTIONS OF OPIOID RECEPTORS

Receptor Type	Actions
Mu (μ) (e.g., morphine, buprenorphine)	Analgesia, reinforcement euphoria, cough and appetite suppression, decreased respirations, decreased GI motility, sedation, hormone changes, dopamine and acetylcholine release
Kappa (κ) (e.g., butorphanol)	Dysphoria, decreased GI motility, decreased appetite, decreased respiration, psychotic symptoms, sedation, diuresis, analgesia
Delta (δ) (e.g., etorphine)	Analgesia, euphoria, physical dependence, Hormone changes, appetite suppression, dopamine release
Nociceptin/orphanin (e.g., buprenorphine)	Analgesia, appetite, anxiety, tolerance to opioids, hypotension, decreased GI motility, 5-HT and NE release

Abbreviations: GI, gastrointestinal; 5-HT, serotonin; NE, norepinephrine.

hyperpolarization, which inhibits the production of action potentials. Thus, opiates inhibit the activity of diverse and widely distributed neuronal types. The major effects of opiates, such as analgesia, sedation, and drug reinforcement are produced through this inhibition of neurons that belong to specific brain pathways.

Many opiate actions are related to the specific neuroanatomic locations of mu receptors. Reinforcing and euphoric effects of opiates occur in the mesolimbic dopaminergic pathway from the ventral tegmental area (VTA) to the nucleus accumbens (NAc), where opiates increase synaptic levels of dopamine. This increase is due to inhibition of GABAergic neurons that inhibit both the activity of neurons within the VTA and the NAc. The positive subjective effects of opiate drugs also include mu receptor desensitization and internalization, potentially related to stimulation of beta-arrestin signaling pathways. However, the “high” only occurs when the *rate of change* in dopamine is fast. Large, rapidly administered doses of opiates block γ -aminobutyric acid (GABA) inhibition and produce a burst of VTA dopamine neuron activity that is associated with “high” in all abused drugs. Therefore, routes of administration that slowly increase opiate blood and brain levels, such as oral and transdermal routes, are effective for analgesia and sedation but do not produce an opiate “high” that follows smoking and intravenous routes. Other acute effects such as analgesia and respiratory depression involve opiate receptors located in other brain areas such as the locus coeruleus (LC).

Opiate tolerance and withdrawal are chronic effects related to the cAMP-protein kinase A (PKA)-cAMP response-element binding protein (CREB) intracellular cascade ([Fig. 468e-1](#)). These effects are also reflective of genetic risk factors for developing opiate use disorder, with estimates of up to 50% of the risk for dependence due to polygenic inheritance. Specific functional polymorphisms in the mu opiate receptor gene appear to be associated with this risk for opiate abuse, including one producing a threefold increase in this receptor’s affinity for opiates and the endogenous ligand beta endorphin. Epigenetic methylation changes also occur on the DNA of the mu receptor gene of opiate addicts, inhibiting gene transcription. This molecular cascade links acute intoxication and sedation to opiate tolerance and withdrawal mediated by the LC. Noradrenergic neurons in the LC mediate activation of the cortical hemispheres. When large opiate doses saturate and activate all of its mu receptors, action potentials cease. When this direct inhibitory effect is sustained over weeks and months of opiate use, a secondary set of adaptive changes occur that lead to tolerance and withdrawal symptoms ([Fig. 468e-1](#)). Withdrawal symptoms reflect, in part, overactivity of norepinephrine (NE) neurons in the LC. This molecular model of NE neuronal activation during withdrawal has had important treatment implications, such as the use of the alpha-2 agonist clonidine to treat opioid withdrawal. Other contributors to withdrawal include deficits within the dopamine reward system.

PHARMACOLOGY

Tolerance and withdrawal commonly occur with chronic daily use, developing as quickly as 6–8 weeks depending on dose concentration

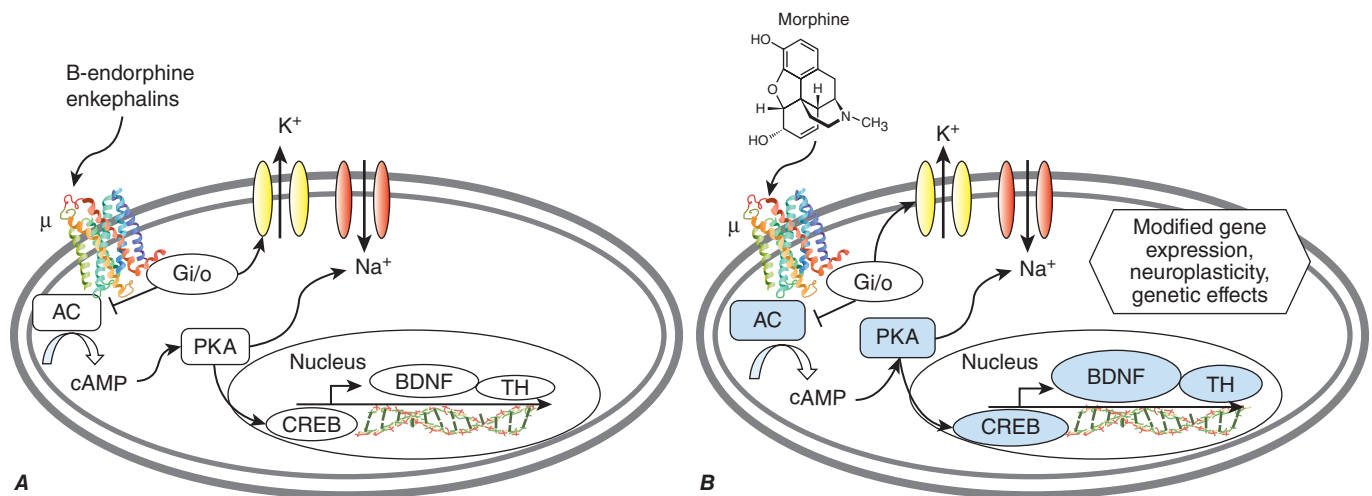


FIGURE 468e-1 Normal mu-receptor activation by endogenous opioids inhibits the cyclic adenosine monophosphate (cAMP)-protein kinase A (PKA)-cAMP response-element binding protein (CREB) cascade in noradrenergic neurons within the locus coeruleus (**A**) through inhibitory Gi/o protein influence on adenylyl cyclase (AC). Similarly, acute exposure to opiates (e.g., morphine) inhibits this system, whereas chronic exposure to opiates (**B**) leads to upregulation of the cAMP pathway in an attempt to oppose opiate-induced inhibitory influence. Upregulation of this system is involved in opiate tolerance, and when the opiate is removed, unopposed noradrenergic neurotransmission is involved in opiate withdrawal. Upregulated PKA phosphorylates CREB, initiating the expression of various genes such as tyrosine hydroxylase (*TH*) and brain-derived neurotrophic factor (*BDNF*). *BDNF* is implicated in long-term neuroplastic changes in response to chronic opiates.

and dosing frequency. Tolerance appears to be primarily a pharmacodynamic rather than pharmacokinetic effect, with relatively limited induction of cytochrome P450 or other liver enzymes. The metabolism of opiates occurs in the liver primarily through the cytochrome P450 systems of 2D6 and 3A4. They then are conjugated to glucuronic acid and excreted in small amounts in feces. The plasma half-lives generally range from 2.5 to 3 h for morphine and more than 22 h for methadone. The shortest half-lives of several minutes are for fentanyl-related opiates and the longest are for buprenorphine and its active metabolites, which can block opiate withdrawal for up to 3 days after a single dose. Tolerance to opioids leads to the need for increasing amounts of drugs to sustain the desired euphoric effects—as well as to avoid the discomfort of withdrawal. This combination has the expected consequence of strongly reinforcing dependence once it has started. Methadone taken chronically at maintenance doses is stored in the liver, which may reduce the occurrence of withdrawal between daily doses. The role of endogenous opioid peptides in tolerance and withdrawal is uncertain.

The clinical features of abuse are tied to route of administration and the rapidity of an opiate bolus in reaching the brain. Intravenous and smoked administration rapidly produces a bolus of high drug concentration in the brain. This bolus produces a “rush,” followed by euphoria, a feeling of tranquility, and sleepiness (“the nod”). Heroin produces effects that last 3–5 h, and several doses a day are required to forestall manifestations of withdrawal in chronic users. Symptoms of opioid withdrawal begin 8–10 h after the last dose; lacrimation, rhinorrhea, yawning, and sweating appear first. Restless sleep followed by weakness, chills, gooseflesh (“cold turkey”), nausea and vomiting, muscle aches, and involuntary movements (“kicking the habit”), hyperpnea, hyperthermia, and hypertension occur in later stages of the withdrawal syndrome. The acute course of withdrawal may last 7–10 days. A secondary phase of protracted abstinence lasts for 26–30 weeks and is characterized by hypotension, bradycardia, hypothermia, mydriasis, and decreased responsiveness of the respiratory center to carbon dioxide.

Besides the brain effects of opioids on sedation and euphoria and the combined brain and peripheral nervous system effects on analgesia, a wide range of other organs can be affected. The cough reflex is inhibited through the brain, leading to the use of some opiates as an antitussive, and nausea and vomiting are due to effects on the medulla. The release of several pituitary hormones is inhibited, including corticotropin-releasing factor (CRF) and luteinizing hormone, which reduces levels of cortisol and sex hormones and can lead to impaired

stress responses and reduced libido. An increase in prolactin also contributes to the reduced sex drive in males. Two other hormones affected are thyrotropin, which is reduced, and growth hormone, which is increased. Respiratory depression results from opiate-induced insensitivity of brainstem neurons to increases in carbon dioxide, and in patients with pulmonary disease, this can result in clinically significant complications. In overdoses, aspiration pneumonia is common due to loss of the gag reflex. Opiates reduce gut motility, which is helpful for treating diarrhea, but can lead to nausea, constipation, and anorexia with weight loss. Deaths occurred in early methadone maintenance programs due to severe constipation and toxic megacolon. Opiates such as methadone may prolong QT intervals and lead to sudden death in some patients. Orthostatic hypotension may occur due to histamine release and peripheral blood vessel dilation, which is an opiate effect usefully applied to managing acute myocardial infarction. During opiate maintenance, interactions with other medications are of concern; these include inducers of the cytochrome P450 system (usually CYP3A4) such as rifampin and carbamazepine.

Heroin users in particular tend to use opiates intravenously and are likely to be polydrug users, also using alcohol, sedatives, cannabinoids, and stimulants. None of these other drugs are substitutes for opioids, but they have desired additive effects. Therefore, one needs to be sure that the person undergoing a withdrawal reaction is not also withdrawing from alcohol or sedatives, which might be more dangerous and more difficult to manage.

Intravenous opiate use carries with it the risk of serious complications. The common sharing of hypodermic syringes can lead to infections with hepatitis B and HIV/AIDS, among others. Bacterial infections can lead to septic complications such as meningitis, osteomyelitis, and abscesses in various organs. Off-target effects of opiates synthesized in illicit drug labs can lead to serious toxicity. For example, attempts to illicitly manufacture meperidine in the 1980s resulted in the production of a highly specific neurotoxin, MPTP, which produced parkinsonism in users ([Chap. 449](#)).

Lethal overdose is a relatively common complication of opiate use disorder. Rapid recognition and treatment with naloxone, a highly specific reversal agent that is relatively free of complications, is essential. The diagnosis is based on recognition of characteristic signs and symptoms, including shallow and slow respirations, pupillary miosis (mydriasis does not occur until significant brain anoxia supervenes), bradycardia, hypothermia, and stupor or coma. Blood or urine toxicology studies can confirm a suspected diagnosis, but immediate

management must be based on clinical criteria. If naloxone is not administered, progression to respiratory and cardiovascular collapse leading to death occurs. At autopsy, cerebral edema and sometimes frothy pulmonary edema are generally found. Opiates generally do not produce seizures except for unusual cases of polydrug use with the opiate meperidine, with high doses of tramadol, or in the newborn.

TREATMENT OPIOID OVERDOSE

Beyond the acute treatment of opiate overdose with naloxone, clinicians have two general treatment options: opioid maintenance or detoxification. Opioid agonist and partial agonist medications are commonly used for both maintenance and detoxification purposes. Alpha-2-adrenergic agonists are primarily used for detoxification. Antagonists are used to accelerate detoxification and then continued after detoxification to prevent relapse. Only the residential medication-free programs have had success that comes close to matching that of the medication-based programs. Success of the various treatment approaches is assessed as retention in treatment and reduced opioid and other drug use; secondary outcomes, such as reduced HIV risk behaviors, crime, psychiatric symptoms, and medical comorbidity, also indicate successful treatment.

Stopping opiates is much easier than preventing relapse. Long-term relapse prevention for opioid-dependent persons requires combined pharmacologic and psychosocial approaches. Chronic users tend to prefer pharmacologic approaches; those with shorter histories of drug abuse are more amenable to detoxification and psychosocial interventions.

OPIATE OVERDOSE

Managing overdose requires naloxone and support of vital functions, including intubation if needed (Table 468e-2). If the overdose is due to buprenorphine, then naloxone might be required at total doses of 10 mg or greater, but primary buprenorphine overdose is nearly impossible because this agent is a partial opiate agonist, meaning that as the dose of buprenorphine is increased it has greater opiate antagonist than agonist activity. Thus, a 0.2-mg buprenorphine dose leads to analgesia and sedation, while a hundred times greater 20-mg dose produces profound opiate antagonism, precipitating opiate withdrawal in a person who was opiate dependent on morphine or methadone. It is important to recognize that the goal is to reverse the respiratory depression and not to administer so much naloxone that it precipitates opiate withdrawal. Because naloxone only lasts a few hours and most opiates last considerably longer, an IV naloxone drip with close monitoring is frequently employed to provide a continuous level of antagonism for 24–72 h depending on the opiate used in the overdose (e.g., morphine vs methadone). Whenever naloxone has only a limited effect, other sedative drugs that produce significant overdoses must be considered. The most common are benzodiazepines, which have produced overdoses and deaths in combination with buprenorphine. A specific antagonist for benzodiazepines—flumazenil at 0.2 mg/min—can be given to a maximum of 3 g/h, but it may precipitate seizures and increase intracranial pressure. Like naloxone, administration for a prolonged period is usually required because most benzodiazepines remain active for considerably longer than flumazenil. Support of vital functions may include oxygen and positive-pressure breathing, IV fluids, pressor agents for hypotension, and cardiac monitoring to detect QT prolongation, which might require specific treatment. Activated

TABLE 468e-2 MANAGEMENT OF OPIOID OVERDOSE

Establish airway. Intubation and mechanical ventilation may be necessary.
Naloxone 0.4–2.0 mg (IV, IM, or endotracheal tube). Onset of action with IV is approximately 1–2 min.
Repeat doses of naloxone if needed to restore adequate respiration or a continuous infusion of naloxone can be used.
One-half to two-thirds of the initial naloxone dose that reversed the respiratory depression is administered on an hourly basis (note: naloxone dosing is not necessary if the patient has been intubated).

charcoal and gastric lavage may be helpful for oral ingestions, but intubation will be needed if the patient is stuporous.

OPIATE WITHDRAWAL

The principles of detoxification are the same for all drugs: to substitute a longer-acting, orally active, pharmacologically equivalent medication for the substance being used, stabilize the patient on that medication, and then gradually withdraw the substituted medication. Methadone or buprenorphine are the two medications used to treat opioid use disorder. Clonidine, a centrally acting sympatholytic agent, has also been used for detoxification in the United States. By reducing central sympathetic outflow, clonidine mitigates many of the signs of sympathetic overactivity but typically requires augmentation with other agents. Clonidine has no narcotic action and is not addictive. Lofexidine, a clonidine analogue with less hypotensive effect, is not yet approved in the United States.

Methadone for Detoxification Dose-tapering regimens for detoxification using methadone range from 2–3 weeks to as long as 180 days, but this approach is controversial given the relative effectiveness of methadone maintenance and the low success rates of detoxification. Unfortunately, the vast majority of patients tend to relapse to heroin or other opiates during or after the detoxification period, indicative of the chronic and relapsing nature of opioid use disorder.

Buprenorphine for Detoxification Buprenorphine does not appear to lead to better outcomes than methadone but is superior to clonidine in reducing symptoms of withdrawal, retaining patients in a withdrawal protocol, and in completing treatment.

Alpha-2-Adrenergic Agonists for Detoxification Several alpha-2-adrenergic agonists have relieved opioid withdrawal by suppressing brain noradrenergic hyperactivity. Clonidine relieves some signs and symptoms of opiate withdrawal such as lacrimation, rhinorrhea, muscle pain, joint pain, restlessness, and gastrointestinal symptoms. Related agents are lofexidine, guanfacine, and guanabenz acetate. Lofexidine can be dosed up to ~2 mg/d and appears to be associated with fewer adverse effects. Clonidine or lofexidine is typically administered orally, in three or four doses per day, with dizziness, sedation, lethargy, and dry mouth as the primary adverse side effects. Outpatient-managed withdrawal will require close follow-up often with naltrexone maintenance to prevent relapse.

Rapid and Ultrarapid Opiate Detoxification The opioid antagonist naltrexone typically combined with an alpha-2-adrenergic agonist has been purported to shorten the duration of withdrawal without significantly increasing patient discomfort. Completion rates using naltrexone and clonidine range from 75 to 81% compared to 40 to 65% for methadone or clonidine alone. Ultrarapid opiate detoxification is an extension of this approach using anesthetics but is highly controversial due to the medical risks and mortality associated with it.

Opioid Agonist Medications for Maintenance Methadone maintenance substitutes a once-daily oral opioid dose for three- to four-times daily heroin. Methadone saturates the opioid receptors and, by inducing a high level of opiate tolerance, blocks the euphoria from additional opiates. Buprenorphine, a partial opioid agonist, also can be given once daily at sublingual doses of 4–32 mg daily, and in contrast to methadone, it can be given in an office-based primary care setting.

METHADONE MAINTENANCE Methadone's slow onset of action when taken orally, long elimination half-life (24–36 h), and production of cross-tolerance at doses from 80 to 150 mg are the basis for its efficacy in treatment retention and reductions in IV drug use, criminal activity, and HIV risk behaviors and mortality. Methadone can prolong the QT interval at rates as high as 16% above the rates in non-methadone-maintained, drug-injecting patients, but it has been used safely in the treatment of opioid dependence for 40 years.

BUPRENORPHINE MAINTENANCE While France and Australia have had sublingual buprenorphine maintenance since 1996, it was first

approved by the U.S. Food and Drug Administration (FDA) in 2002 as a Schedule III drug for managing opioid use disorder. Unlike the full agonist methadone, buprenorphine is a partial agonist of mu-opioid receptors with a slow onset and long duration of action. Its partial agonism reduces the risk of unintentional overdose but limits its efficacy to patients who need the equivalent of only 60–70 mg of methadone, and many patients in methadone maintenance require higher doses up to 150 mg daily. Buprenorphine is combined with naloxone at a 4:1 ratio in order to reduce its abuse liability. Because of pediatric exposures and diversion of buprenorphine to illicit use, a new formulation, using mucosal films rather than sublingual pills that were crushed and snorted, is now marketed. A subcutaneous buprenorphine implant that lasts up to 6 months has also been tested and is pending FDA approval as a formulation improvement to prevent pediatric exposures and illicit diversion and enhance compliance.

In the United States, the ability of primary care physicians to prescribe buprenorphine for opioid use disorder represents an important opportunity to improve access and quality of treatment as well as reduce social harm. Europe, Asia, and Australia have found reduced opioid-related deaths and drug-injection-related medical morbidity with buprenorphine available in primary care. Retention in office-based buprenorphine treatment has been high as 70% at 6-month follow-ups.

Opioid Antagonist Medications The rationale for using narcotic antagonist therapy is that blocking the action of self-administered opioids should eventually extinguish the habit, but this therapy is poorly accepted by patients. Naltrexone, a long-acting orally active pure opioid antagonist, can be given three times a week at doses of 100–150 mg. Because it is an antagonist, the patient must first be detoxified from opioid dependence before starting naltrexone. It is safe even when taken chronically for years, is associated with few side effects (headache, nausea, abdominal pain), and can be given to patients infected with hepatitis B or C without producing hepatotoxicity. However, most providers refrain from prescribing naltrexone if liver function tests are three times above normal levels. Naltrexone maintenance

combined with psychosocial therapy is effective in reducing heroin use, but medication adherence is low. Depot injection formulations lasting up to 4 weeks markedly improve adherence, retention, and drug use. Subcutaneous naltrexone implants in Russia, China, and Australia have doubled treatment retention and reduced relapse to half that of oral naltrexone. In the United States, a depot naltrexone formulation is available for monthly use and maintains blood levels equivalent to 25 mg of daily oral use.

Medication-Free Treatment Most opiate addicts enter medication-free treatments in inpatient, residential, or outpatient settings, but 1- to 5-year outcomes are very poor compared to pharmacotherapy except for residential settings lasting 6 to 18 months. The residential programs require full immersion in a regimented system with progressively increasing levels of independence and responsibility within a controlled community of fellow drug abusers. These medication-free programs, as well as the pharmacotherapy programs, also include counseling and behavioral treatments designed to teach interpersonal and cognitive skills for coping with stress and for avoiding situations leading to easy access to drugs or to craving. Relapse is prevented by having the individual very gradually reintroduced to greater responsibilities and to the working environment outside of the protected therapeutic community.

PREVENTION

Preventing opiate abuse represents a critically important challenge for physicians. Opiate prescriptions are the most common source of drugs accessed by adolescents who begin a pattern of illicit drug use. The major sources of these drugs are family members, not drug dealers or the Internet. Pain management involves providing sufficient opiates to relieve the pain over as short a period of time as the pain warrants (**Chap. 18**). The patient then needs to dispose of any remaining opiates, not save them in the medicine cabinet, because this behavior leads to diversion by adolescents. Finally, physicians should never prescribe opiates for themselves.

469e Cocaine and Other Commonly Abused Drugs

Nancy K. Mello¹, Jack H. Mendelson¹

The abuse of cocaine and other psychostimulants reflects a complex interaction between the pharmacology of the drug, the personality and expectations of the user, and the environmental context in which the drug is used. Polydrug abuse involving the concurrent use of several drugs with different pharmacologic effects is increasingly common. Sometimes one drug is used to enhance the effects of another, as with the combined use of cocaine and nicotine, benzodiazepines and methadone, or cocaine and heroin in methadone-maintained patients. Some forms of polydrug abuse, such as the combined use of IV heroin and cocaine, are especially dangerous and account for many hospital emergency room visits.



Chronic cocaine and psychostimulant abuse may cause a number of adverse health consequences and may exacerbate preexisting disorders such as hypertension and cardiac disease. The combined use of two or more drugs may accentuate medical complications associated with abuse of one drug. Chronic drug abuse is often associated with immune system dysfunction and increased vulnerability to infections, including risk for HIV infection. In addition, concurrent use of cocaine and opiates (the “speedball”) is frequently associated with needle sharing by IV drug users. IV drug abusers continue to be the largest single group of persons with HIV infection in several major metropolitan areas in the United States as well as in many parts of Europe and Asia.

Stimulants and hallucinogens have been used to induce euphoria and alter consciousness for centuries. Cocaine and marijuana are two of the most commonly abused drugs today. Synthetic variations of marijuana and a variety of hallucinogens have become popular recently, and new drugs are continually being developed. This chapter describes the subjective and adverse medical effects of cocaine, marijuana, and lysergic acid diethylamide (LSD), as well as methamphetamine, 3,4-methylenedioxy-N-methamphetamine (MDMA), synthetic cathinones (bath salts), phencyclidine (PCP), *Salvia divinorum*, and other drugs of abuse (flunitrazepam, γ -hydroxybutyric acid [GHB], ketamine). Some options for medical management of severe adverse effects are also described.

COCAINE

Cocaine is a stimulant and a local anesthetic with potent vasoconstrictor properties. The leaves of the *coca* plant (*Erythroxylum coca*) contain ~0.5–1% cocaine. The drug produces physiologic and behavioral effects after oral, intranasal, IV, or inhalation/smoking routes of administration. The reinforcing effects of cocaine are related to activation of dopaminergic neurons in the mesolimbic system (**Chap. 465e**). Cocaine increases synaptic concentrations of the monoamine neurotransmitters dopamine, norepinephrine, and serotonin by binding to transporter proteins in presynaptic neurons and blocking reuptake.

PREVALENCE OF COCAINE USE

Cocaine is widely available and is abused in virtually all social and economic strata of society. In 2012, an estimated 1.6 million persons in the United States used cocaine, and 1.1 million abused or were dependent on cocaine. Emergency room admissions involving cocaine totaled 505,224 in 2011. Cocaine abuse is prevalent in the general population and in heroin-dependent persons, including those in methadone maintenance programs. IV cocaine is often used concurrently with IV heroin in a combination called a “speedball.” This combination purportedly attenuates the postcocaine “crash” and substitutes a cocaine “high” for the heroin “high” blocked by methadone.

ACUTE AND CHRONIC INTOXICATION

There has been an increase in both IV administration and inhalation of pyrolyzed cocaine via smoking. Following intranasal administration, changes in mood and sensation are perceived within 3–5 min, and peak effects occur at 10–20 min. These effects rarely last more than 1 h. Inhalation of pyrolyzed materials includes inhaling crack/cocaine or smoking coca paste, a product made by extracting cocaine preparations with flammable solvents, and cocaine free-base smoking. Free-base cocaine, including the free-base prepared with sodium bicarbonate (crack), has become increasingly popular because of its relative high potency and rapid onset of action (8–10 seconds following smoking).

Cocaine produces a brief, dose-related stimulation and euphoria and an increase in cardiac rate and blood pressure. Body temperature usually increases following cocaine administration, and high doses of cocaine may induce lethal pyrexia or hypertension. Because cocaine inhibits reuptake of catecholamines at adrenergic nerve endings, it potentiates sympathetic nervous system activity. Cocaine has a short plasma half-life of approximately 45–60 min. Cocaine is metabolized by plasma esterases, and cocaine metabolites are excreted in urine. The brief duration of the euphorogenic effects of cocaine reported by chronic abusers is probably due to both acute and chronic tolerance. Cocaine may be used as often as two to three times per hour. Alcohol is often used to modulate both the cocaine high and the dysphoria associated with the abrupt disappearance of cocaine’s effects. A metabolite of cocaine, cocaethylene, has been detected in blood and urine of persons who concurrently abuse alcohol and cocaine. Cocaethylene induces changes in cardiovascular function similar to those of cocaine alone, and the pathophysiologic consequences of the concurrent abuse of alcohol plus cocaine may be additive.

Cocaine may cause serious medical consequences by any route of administration. The prevalent assumption that cocaine inhalation or IV administration is relatively safe is contradicted by reports of death from respiratory depression, cardiac arrhythmias, and convulsions associated with cocaine use. In addition to generalized seizures, neurologic complications may include headache, ischemic or hemorrhagic stroke, or subarachnoid hemorrhage. Disorders of cerebral blood flow and perfusion in cocaine-dependent persons have been detected with magnetic resonance spectroscopy (MRS). Inhalation of crack cocaine may lead to severe pulmonary disease due to the direct effects of cocaine and to residual contaminants in the smoked material. Hepatic necrosis may occur following chronic crack/cocaine use. Protracted cocaine abuse may also cause paranoid ideation and visual and auditory hallucinations, a state that resembles alcoholic hallucinosis.

Although men and women who abuse cocaine may report that the drug enhances libidinal drive, chronic cocaine use causes significant loss of libido and adversely affects sexual function. Impotence and gynecomastia have been observed in male cocaine abusers, and these abnormalities often persist for long periods following cessation of drug use. Cocaine abuse may produce major derangements in menstrual cycle function including galactorrhea, amenorrhea, and infertility in women and in a rhesus monkey model of cocaine self-administration. Chronic cocaine abuse may cause persistent hyperprolactinemia as a consequence of disordered dopaminergic inhibition of prolactin secretion by the anterior pituitary. Cocaine abuse by pregnant women, particularly crack smoking, has been associated with both an increased risk of congenital malformations in the fetus and perinatal cardiovascular and cerebrovascular disease in the mother. However, cocaine abuse per se is probably not the sole cause of these perinatal disorders, because maternal cocaine abuse is often associated with poor nutrition and prenatal health care as well as polydrug abuse that may contribute to the risk for perinatal disease.

Psychological dependence on cocaine, indicated by inability to abstain from frequent compulsive use, has been reported. Although the occurrence of withdrawal syndromes involving psychomotor agitation and autonomic hyperactivity remains controversial, severe depression (“crashing”) following cocaine intoxication may accompany drug withdrawal.

¹Deceased

Treatment of cocaine overdose is a medical emergency that is best managed in an intensive care unit. Cocaine toxicity produces a hyperadrenergic state characterized by hypertension, tachycardia, tonic-clonic seizures, dyspnea, and ventricular arrhythmias. IV diazepam in doses up to 0.5 mg/kg administered over an 8-h period has been shown to be effective for control of seizures. Ventricular arrhythmias have been managed successfully by administration of 0.5–1.0 mg of propranolol IV. Because many instances of cocaine-related mortality have been associated with concurrent use of other illicit drugs (particularly heroin), the physician must be prepared to institute effective emergency treatment for multiple drug toxicities.

Treatment of chronic cocaine abuse requires the combined efforts of primary care physicians, psychiatrists, and psychosocial care providers. Early abstinence from cocaine use is often complicated by symptoms of depression and guilt, insomnia, and anorexia, which may be as severe as those observed in major affective disorders. Individual and group psychotherapy, family therapy, and peer group assistance programs are often useful for inducing prolonged remission from drug use. Although psychotherapy may be helpful, no specific form of psychotherapy or behavioral modification is uniquely beneficial.

A number of medications used for the treatment of various medical and psychiatric disorders have been administered to reduce the duration and severity of cocaine abuse and dependence. The search for a medication that is both safe and highly effective for cocaine detoxification or maintenance of abstinence is continuing. Clinical trials of buspirone (BuSpar), a nonbenzodiazepine anxiolytic with dopamine D₃ and D₄ receptor antagonist properties, are ongoing. Buspirone reduced use of cocaine, nicotine, and cocaine plus nicotine in combination in a nonhuman primate model of stimulant addiction.

Another approach to reducing cocaine abuse is the development of vaccines to actively immunize against cocaine or to functionally antagonize cocaine by preventing it from reaching the brain. Cocaine is converted into inactive metabolites by the plasma enzyme, butyrylcholinesterase (BChE). When this enzyme is modified to increase its catalytic efficiency and accelerate cocaine metabolism, it can both prevent and reverse cocaine-induced toxicity in animals. Importantly, it remains effective even when high doses of cocaine are administered. Ongoing development of this approach includes cocaine hydrolase gene therapy. Vaccines for both cocaine and nicotine have been designed and shown to be safe and somewhat effective in clinical trials. Individual variability in antibody titers and difficulties in determining the optimally effective antibody titer that will neutralize responses to increasing doses of cocaine and have a relatively long duration of action are among the challenges that remain to be resolved.

MARIJUANA AND CANNABIS COMPOUNDS

Cannabis sativa contains >400 compounds in addition to the psychoactive substance, delta-9-tetrahydrocannabinol (THC). Marijuana cigarettes are prepared from the leaves and flowering tops of the plant, and a typical marijuana cigarette contains 0.5–1 g of plant material. The usual THC concentration varies between 10 and 40 mg, but concentrations <100 mg per cigarette have been detected. Hashish is prepared from concentrated resin of *C. sativa* and contains a THC concentration of between 8 and 12% by weight. “Hash oil,” a lipid-soluble plant extract, may contain THC between 25 and 60% and may be added to marijuana or hashish to enhance its THC concentration. Smoking is the most common mode of marijuana or hashish use. During pyrolysis, <150 compounds in addition to THC are released in the smoke. Although most of these compounds do not have psychoactive properties, they may have physiologic effects.

THC is quickly absorbed from the lungs into blood and then rapidly sequestered in tissues. THC is metabolized primarily in the liver, where

it is converted to 11-hydroxy-THC, a psychoactive compound, and >20 other metabolites. Many THC metabolites are excreted through the feces at a relatively slow rate of clearance compared with most other psychoactive drugs.

Specific cannabinoid receptors (CB₁ and CB₂) have been identified in the central and peripheral nervous system. High densities of cannabinoid receptors have been found in the cerebral cortex, basal ganglia, and hippocampus. T and B lymphocytes also contain cannabinoid receptors, and these appear to mediate the anti-inflammatory and immunoregulatory properties of cannabinoids. A naturally occurring THC-like ligand has been identified and is widely distributed in the nervous system.

Herbal marijuana alternatives are also available. These are usually a combination of several herbs and synthetic cannabinoids. “Spice” and “K2” are among the best known, but many formulations exist, and marijuana is undetectable by the usual methods. These compounds are marketed on the Internet as containing no illegal ingredients. However a number of synthetic cannabinoids are now classified as Schedule I by the Drug Enforcement Administration due to reports of toxicity.

PREVALENCE OF USE

Marijuana is the most commonly used illegal drug in the United States. In 2012, an estimated 18.9 million people reported using marijuana within the past month. An estimated 7.2% of adolescents age 12 to 17 years reported current use of marijuana. Marijuana is relatively inexpensive and is often considered to be less hazardous than other controlled drugs and substances. Very potent forms of marijuana (sinsemilla) are widely available, and concurrent use of marijuana with other drugs such as cocaine is not uncommon. Due in part to the difficulty of detecting herbal marijuana alternatives, the prevalence of use is unknown.

ACUTE AND CHRONIC INTOXICATION

Acute intoxication from marijuana and cannabis compounds is related to both the dose of THC and the route of administration. THC is absorbed more rapidly from marijuana smoking than from orally ingested cannabis compounds. Acute marijuana intoxication may produce a perception of relaxation and mild euphoria resembling mild to moderate alcohol intoxication. This condition is usually accompanied by some impairment in thinking, concentration, and perceptual and psychomotor function. Higher doses of cannabis may produce more pronounced impairment in concentration and perception, as well as greater sedation. Although the acute effects of marijuana intoxication are relatively benign in normal users, the drug can precipitate severe emotional disorders in individuals who have antecedent psychotic or neurotic problems. Like other psychoactive compounds, both the user’s expectations and the environmental context are important determinants of the type and severity of the effects of marijuana intoxication.

As with abuse of cocaine, opioids, and alcohol, chronic marijuana abusers may lose interest in common socially desirable goals and devote progressively more time to drug acquisition and use. However, THC does not cause a specific and unique “amotivational syndrome.” The range of symptoms sometimes attributed to marijuana use is difficult to distinguish from mild to moderate depression and the maturational dysfunctions often associated with protracted adolescence. Chronic marijuana use has also been reported to increase the risk of psychotic symptoms in individuals with a past history of schizophrenia. Persons who begin marijuana smoking before the age of 17 may have more pronounced cognitive deficits and also may be at higher risk for polydrug and alcohol abuse problems in later life, but the role of marijuana in this sequence is uncertain.

The acute effects of herbal marijuana alternatives are based primarily on case reports and include anxiety, agitation, delusions, paranoia, and psychosis. The extent to which these symptoms reflect drug effects or exacerbation of an underlying psychiatric disorder is often difficult to determine.

PHYSICAL EFFECTS

Conjunctival injection and tachycardia are the most frequent immediate physical concomitants of smoking marijuana. Tolerance for

marijuana-induced tachycardia develops rapidly among regular users. However, marijuana smoking may precipitate angina in persons with a history of coronary insufficiency. Exercise-induced angina may increase after marijuana use to a greater extent than after tobacco cigarette smoking. Patients with cardiac disease should be strongly advised not to smoke marijuana or use cannabis compounds.

Significant decrements in pulmonary vital capacity have been found in regular daily marijuana smokers. Because marijuana smoking typically involves deep inhalation and prolonged retention of marijuana smoke, chronic bronchial irritation may develop. Impairment of single-breath carbon monoxide diffusion capacity (DL_{CO}) is greater in persons who smoke both marijuana and tobacco than in tobacco smokers.

Although marijuana has also been associated with a number of other adverse effects, many of these studies await replication and confirmation. A reported correlation between chronic marijuana use and decreased testosterone levels in males has not been confirmed. Decreased sperm count and sperm motility and morphologic abnormalities of spermatozoa following marijuana use have been reported. Prospective studies found a correlation between impaired fetal growth and development and heavy marijuana use during pregnancy. Marijuana has also been implicated in derangements of the immune system; in chromosomal abnormalities; and in inhibition of DNA, RNA, and protein synthesis; however, these findings have not been confirmed or related to any specific physiologic effect in humans. Herbal marijuana alternatives produce many of the effects of marijuana including conjunctival injection and tachycardia.

TOLERANCE AND PHYSICAL DEPENDENCE

Habitual marijuana users may develop tolerance to the psychoactive effects of marijuana, and then smoke more frequently and try to acquire more potent cannabis compounds. Tolerance for the physiologic effects of marijuana develops at different rates; e.g., tolerance develops rapidly for marijuana-induced tachycardia but more slowly for marijuana-induced conjunctival injection. Tolerance for both behavioral and physiologic effects of marijuana decreases rapidly upon cessation of marijuana use.

A distinct withdrawal syndrome has been documented in chronic cannabis users, and the severity of symptoms is related to dosage and duration of use. These symptoms typically reach their peak several days after cessation of chronic use and include irritability, anorexia, and sleep disturbances. Withdrawal signs and symptoms observed in chronic marijuana users are usually relatively mild in comparison to those observed in heavy opioid or alcohol users and rarely require medical or pharmacologic intervention. However, more severe and protracted abstinence syndromes may occur after sustained use of high-potency cannabis compounds. As yet there have been no systematic studies of tolerance and physical dependence to the herbal marijuana alternatives. The large number of synthetic cannabinoids available for combination with about 20 herbs presents a daunting challenge for analysis.

THERAPEUTIC USE OF MARIJUANA

Marijuana, administered as cigarettes or as a synthetic oral cannabinoid (dronabinol), is thought to have a number of clinically useful medicinal properties. These include antiemetic effects in chemotherapy recipients, appetite-promoting effects in AIDS patients, reduction of intraocular pressure in glaucoma, and reduction of spasticity in multiple sclerosis and other neurologic disorders. With the possible exception of AIDS-related cachexia, none of these attributes of marijuana compounds is clearly superior to other readily available therapies.

METHAMPHETAMINE

Methamphetamine is also referred to as “meth,” “speed,” “crank,” “chalk,” “ice,” “glass,” or “crystal.” Methamphetamine is a mixed-action monoamine releaser with activity at dopamine, serotonin, and norepinephrine systems. Methamphetamine was considered second only to cocaine as a drug threat to society by the U.S. Department

of Justice in 2009. Hospital admissions for methamphetamine treatment more than doubled between 1998 and 2007, and young adults (age 18–25) have the highest use rates. In 2011, an estimated 439,000 people reported current use of methamphetamine in the United States, and emergency room admissions involving amphetamines/methamphetamine totaled 160,000. Persistent abuse of methamphetamine continues despite drug seizures, closures of clandestine laboratories that produce methamphetamine illegally, and an increase in methamphetamine abuse prevention programs.

Methamphetamine can be used by smoking, snorting, IV injection, or oral administration. Methamphetamine abusers report that drug use induces feelings of euphoria and decreased fatigue. Adverse consequences of methamphetamine use include headache, difficulty concentrating, diminished appetite, abdominal pain, vomiting or diarrhea, disordered sleep, paranoid or aggressive behavior, and psychosis. Chronic methamphetamine abuse can result in severe dental caries, described as blackened, rotting, crumbling teeth. Severe, life-threatening methamphetamine toxicity may include hypertension, cardiac arrhythmia or cardiac failure, subarachnoid hemorrhage, ischemic stroke, intracerebral hemorrhage, convulsions, or coma.

Methamphetamines increase the release of monoamine neurotransmitters (dopamine, norepinephrine, and serotonin) from presynaptic neurons. It is thought that the euphoric and reinforcing effects of this class of drugs are mediated through dopamine and the mesolimbic system, whereas the cardiovascular effects are related to norepinephrine. MRS studies of the brain suggest that chronic abusers have neuronal damage in the frontal areas and basal ganglia.

Treatment of acute methamphetamine overdose is largely symptomatic. Ammonium chloride may be useful to acidify the urine and enhance clearance of the drug. Hypertension may respond to sodium nitroprusside or α -adrenergic antagonists. Sedatives may reduce agitation and other signs of central nervous system hyperactivity. Treatment of chronic methamphetamine dependence may be accomplished in either an inpatient or outpatient setting using strategies similar to those described earlier for cocaine abuse.

MDMA is a derivative of methamphetamine also called *Ecstasy* or *Molly*. Reported use of MDMA in the United States has increased from 615,000 persons in 2005, to an estimated 869,000 people in 2012. Emergency ward admissions involving MDMA totaled more than 22,000 in 2011. Ecstasy is usually taken orally but may be injected or inhaled, and its effects last for 3–6 h. MDMA has amphetamine-like effects including vivid visual and auditory hallucinations and other perceptual distortions. Recent studies indicate that MDMA use is associated with cognitive and memory impairment. MDMA can induce hyperthermia and elevated blood pressure, seizures, coma, and death. Withdrawal symptoms after cessation of use may include teeth grinding, anxiety, loss of appetite, insomnia, and fever. The long-term consequences of recreational use of MDMA by young persons are poorly understood.

SYNTHETIC CATHINONES (BATH SALTS)

The rapid emergence of synthetic cathinone abuse during 2010 was accompanied by numerous reports of adverse medical and psychiatric effects, suicides, and deaths. Reports to poison centers and health agencies increased from about 300 in 2010 to over 6000 in 2011. In 2011, the Drug Enforcement Administration classified three commonly abused synthetic cathinones (mephedrone [4-methyl methcathinone], MDPV [3,4-methylenedioxy pyrovalerone], and methylene) as Schedule I compounds with no accepted medical use and a high potential for abuse. However, synthetic cathinones are readily available on the Internet as well as in convenience stores, gas stations, and head shops. These drugs are merchandised under a variety of names such as Vanilla Sky, Purple Wave, Blue Silk, White Lightning, and Snow Leopard. Regulatory constraints are evaded by labeling the products as plant food, insecticides, pond cleaner, and bath salts with the qualifier, “not for human consumption.”

Cathinone is the primary psychoactive ingredient in khat leaves. Chewing the leaves of the khat shrub (*Catha edulis*) produces mild

stimulant and euphoric effects and remains a common practice in east Africa that has persisted for centuries. Cathinone is structurally similar to amphetamine, and mephedrone is structurally similar to methamphetamine. Cathinones, like amphetamines, inhibit dopamine, serotonin, and norepinephrine transporters to varying degrees, and this probably accounts for variations in the behavioral effects observed. The effects of cathinone derivatives are often described as similar to the effects of MDMA or Ecstasy. Synthetic cathinones can be inhaled, snorted, injected, or taken orally. These drugs may be taken repeatedly over several hours in episodes lasting for hours or days. The onset of effects after oral ingestion is relatively rapid for MDPV (15–30 min) and slightly slower for mephedrone and methylone (30–45 min). The duration of action varies from 2 to 5 or 7 h. After mephedrone inhalation, effects occur within minutes and only last for 1 h or less, but mood changes may persist for several days. Evaluation of the neurotoxic effects of prolonged synthetic cathinone abuse is just beginning, and the long-term consequences are unknown.

The reported positive subjective effects of synthetic cathinones include euphoria, improved energy, alertness, sociability, and increased sensitivity to music and other sensory experiences. The reported negative subjective effects include agitation, visual and auditory hallucinations, anxiety and panic attacks, paranoid delusions, disorientation, depression, and suicidal ideation. Observers report irritability, aggression, violent behavior, tremors, and seizures. Medical evidence of adverse effects includes cardiovascular dysfunction and cardiac arrest, hypertension, hyperthermia, nausea and vomiting, and anorexia. There is no specific antagonist for synthetic cathinone intoxication. Patients with severe hyperthermia, seizures, and arrhythmia are medical emergencies and should be treated in a hospital. Sedation with benzodiazepines can be useful for managing agitation, seizures, aggression, and other related symptoms. Antipsychotic medications may be necessary for management of severe and persistent psychiatric symptoms.

LYSERGIC ACID DIETHYLAMIDE (LSD)

Discovery of the psychedelic effects of LSD led to an epidemic of LSD abuse during the 1960s. Imposition of stringent constraints on the manufacture and distribution of LSD (classified as a Schedule I substance by the U.S. Food and Drug Administration [FDA]) and public recognition that psychedelic experiences induced by LSD were a health hazard have resulted in a reduction in LSD abuse. LSD still remains popular among adolescents and young adults, and there are indications that LSD use among young persons has been increasing in some areas in the United States. In 2011, an estimated 358,000 persons used LSD, whereas 200,000 and 271,000 persons reported LSD use in 2003 and 2007, respectively.

LSD is a very potent hallucinogen; oral doses as low as 20 µg may induce profound psychological and physiologic effects. Tachycardia, hypertension, pupillary dilation, tremor, and hyperpyrexia occur within minutes following oral administration of 0.5–2 µg/kg. A variety of bizarre and often conflicting perceptual and mood changes, including visual illusions, synesthesias, and extreme lability of mood, usually occur within 30 min after LSD intake. These effects of LSD may persist for 12–18 h, even though the half-life of the drug is only 3 h.

Emergency ward visits involving LSD totaled nearly 5000 in 2011. The most frequent acute medical emergency associated with LSD use is a panic episode (the “bad trip”), which may persist up to 24 h. Management of this problem is best accomplished by supportive reassurance (“talking down”) and, if necessary, administration of small doses of anxiolytic drugs. Adverse consequences of chronic LSD use include an enhanced risk for schizophreniform psychosis and derangements in memory function, problem solving, and abstract thinking. Treatment of these disorders is best carried out in specialized psychiatric facilities.

Tolerance develops rapidly for LSD-induced changes in psychological function when the drug is used one or more times per day for >4 days. Abrupt abstinence following continued use does not produce withdrawal signs or symptoms. There have been no clinical reports of death caused by the direct effects of LSD.

PHENCYCLIDINE (PCP)

PCP, a cyclohexylamine derivative, is widely used in veterinary medicine to briefly immobilize large animals and is sometimes described as a dissociative anesthetic. PCP binds to ionotropic *N*-methyl-D-aspartate (NMDA) receptors in the nervous system, blocking ion current through these channels. PCP is easily synthesized and is abused primarily by young people and polydrug users. It is used orally, by smoking, by snorting, or by IV injection. It is also used as an adulterant in THC, LSD, amphetamine, or cocaine. The most common street preparation, *angel dust*, is a white granular powder that contains 50–100% of the drug. Low doses (5 mg) produce agitation, excitement, impaired motor coordination, dysarthria, and analgesia. Physical signs of intoxication may include horizontal or vertical nystagmus, flushing, diaphoresis, and hyperacusis. Behavioral changes include distortions of body image, disorganization of thinking, and feelings of estrangement. Higher doses of PCP (5–10 mg) may produce profuse salivation, vomiting, myoclonus, fever, stupor, or coma. PCP doses of ≥10 mg cause convulsions, opisthotonus, and decerebrate posturing that may be followed by prolonged coma.

In 2011, more than 75,000 emergency ward admissions involved PCP. The diagnosis of PCP overdose is difficult because the patient's initial symptoms (anxiety, paranoia, delusions, and hallucinations) may suggest an acute schizophrenic reaction. Confirmation of PCP use is possible by determination of PCP levels in serum or urine. PCP assays are available at most toxicologic centers. PCP remains in urine for 1–5 days following high-dose intake.

PCP overdose requires emergency life-support measures that may involve treatment of coma, convulsions, and respiratory depression in an intensive care unit. There is no specific antidote or antagonist for PCP. PCP excretion from the body can be enhanced by gastric lavage and acidification of urine. Death from PCP overdose may occur as a consequence of some combination of pharyngeal hypersecretion, hyperthermia, respiratory depression, severe hypertension, seizures, hypertensive encephalopathy, and intracerebral hemorrhage.

Acute psychosis associated with PCP use is a psychiatric emergency because patients may be at high risk for suicide or extreme violence toward others. Phenothiazines should not be used for treatment because these drugs potentiate PCP's anticholinergic effects. Haloperidol (5 mg IM) has been administered on an hourly basis to induce suppression of psychotic behavior. PCP, like LSD and mescaline, produces vasospasm of cerebral arteries at relatively low doses. Chronic PCP use has been shown to induce insomnia, anorexia, severe changes in behavior, and, in some cases, chronic schizophrenia.

SALVIA DIVINORUM

This naturally occurring herb is a recent entry into the spectrum of hallucinogens. Like PCP and ecstasy, this drug can produce profound alterations in mood, hallucinations, and distorted perceptions. This drug is available on the Internet and is known by a variety of names including *magic mint*, *mystic sage*, *Mariana Pastora*, and *purple sticky*. The drug was first added to the annual National Surveys on Drug Use and Health in 2006, and its use is increasing. Between 2006 and 2011, the number of estimated users in the United States nearly tripled to more than 5000.

The active ingredient is salvinorin A, a selective kappa opioid receptor agonist that has a range of effects including hallucinations, sedation, analgesia, and depression. The hallucinatory symptoms may be associated with intense anxiety and severe agitation that can be managed with benzodiazepines. Importantly, this kappa opioid receptor agonist does not produce respiratory depression, and no significant change in blood pressure or heart rate was reported in a clinical study with healthy subjects.

Salvinorin A extract or crushed leaves of the *Salvia divinorum* plant can be chewed and absorbed through the buccal membrane or inhaled during smoking. The onset of the acute “high” is within 5–10 min after chewing and 30 s after inhalation. The duration of the effect is relatively brief, usually 15–20 min. However, if the drug is taken with alcohol or other hallucinogens, the duration and intensity of adverse

effects may be increased. The effects of the drug are reported to be similar to those of ketamine, LSD, and marijuana.

OTHER DRUGS OF ABUSE

A number of other pharmacologically diverse drugs of abuse are often referred to as “club drugs” because these are frequently used in bars, at concerts, and at rave parties. Commonly abused club drugs include flunitrazepam, GHB, and ketamine and are described below. Methamphetamine, MDMA, and LSD are also considered club drugs and were described earlier in this chapter. Abuse of club drugs at high doses, especially in combination with alcohol, can be lethal and should be treated as a medical emergency. GHB and ketamine can be identified in blood, and flunitrazepam can be identified in urine and hair samples. Flunitrazepam and GHB toxicity can be treated with antagonists at benzodiazepine and γ -aminobutyric acid B (GABA_B) receptors, respectively.

Flunitrazepam (Rohypnol) is a benzodiazepine derivative primarily used to treat insomnia, but it has significant abuse potential because of its strong hypnotic, anxiolytic, and amnesia-producing effects. It is a club drug commonly referred to as a “date-rape drug” or “roofies.” The drug enhances GABA_A receptor activity, and overdose can be treated with flumazenil, a benzodiazepine receptor antagonist. Flunitrazepam is typically used orally but can be snorted or injected. Concomitant use of alcohol or opioids is common, and this enhances the sedative and hypnotic effects of flunitrazepam and also the risk of motor vehicle accidents. Overdose can produce life-threatening respiratory depression and coma. Abrupt cessation after chronic use may result in a benzodiazepine withdrawal syndrome consisting of anxiety, insomnia, disordered thinking, and seizures.

GHB (Xyrem) is a sedative drug that is approved by the FDA for the treatment of narcolepsy. It is classified as a club drug, is sometimes used in combination with alcohol or other drugs of abuse, and has been implicated in cases of date rape. It is also used by body builders as a growth hormone stimulant. GHB is usually available as a liquid, is taken orally, and has no distinctive color or odor. Its stimulant properties are attributed to agonist activity at the GHB receptor, but it also has sedative effects at high doses that reflect its activity at GABA_B receptors. GABA_B antagonists can reverse GHB’s sedative effects, and opioid antagonists (naloxone, naltrexone) can attenuate GHB effects on dopamine release. Low doses of GHB may produce euphoria and disinhibition, whereas high doses result in nausea, agitation, convulsions, and sedation that can lead to unconsciousness and death from respiratory depression. In 2011, more than 2400 emergency ward admissions involved GHB.

Ketamine (Ketaset, Ketalar) is a dissociative anesthetic, similar to PCP. In veterinary medicine, it is used for brief immobilization. In clinical medicine, it is used for sedation, analgesia, and to supplement anesthesia. Ketamine increases heart rate and blood pressure, with less respiratory depression than other anesthetics. Ketamine’s popularity as a club drug appears to reflect its ability to induce a dissociative state and feelings of depersonalization, accompanied by intense hallucinations and subsequent amnesia. It can be administered orally, by smoking (usually in combination with tobacco and/or marijuana), or by IV or IM injection. Like PCP, it binds to NMDA receptors and acts as a noncompetitive NMDA antagonist. In 2011, ketamine accounted for 1550 emergency ward admissions. Ketamine has a complex profile of action and appears to be useful as an antidepressant in treatment-resistant patients and as an analgesic in patients with chronic pain.

The extent to which chronic recreational use leads to memory impairment remains controversial.

POLYDRUG ABUSE

Although some drug abusers may prefer a particular drug, the concurrent use of multiple drugs is common. Polydrug abuse often involves substances that may have different pharmacologic effects from the preferred drug. For example, concurrent use of such dissimilar compounds as stimulants and opioids or stimulants and alcohol is common. The diversity of reported drug use combinations suggests that achieving a change in subjective state, rather than any particular direction of change (stimulation or sedation), may be the primary reinforcer in polydrug abuse. There is also evidence that intoxication with alcohol, opiates, and cocaine is associated with increased tobacco smoking. Nicotine and cocaine enhance each other’s effects in clinical laboratory studies, and this drug combination maintains significantly higher levels of self-administration than either drug alone in preclinical models of addiction. There are relatively few controlled studies of multiple drug interactions. However, the combined use of cocaine, heroin, and alcohol increases the risk for toxic effects and adverse medical consequences. Similarly, some hallucinogens (MDMA, LSD) and club drugs (GHB, ketamine, flunitrazepam) are used in various combinations with an associated increase in toxic consequences.

One determinant of polydrug use patterns is the relative availability and cost of the drugs. For example, alcohol abuse, with its attendant medical complications, is one of the most serious problems encountered in former heroin addicts participating in methadone maintenance programs. Cocaine abuse often increases during methadone maintenance.

The physician must recognize that perpetuation of polydrug abuse and drug dependence is not necessarily a symptom of an underlying emotional disorder. Neither alleviation of anxiety nor reduction of depression accounts for initiation and perpetuation of polydrug abuse. Severe depression and anxiety are the consequences of polydrug abuse as frequently as they are the antecedents. Interestingly, some adverse consequences of drug use may be reinforcing and contribute to the continuation of polydrug abuse.

Adequate treatment of polydrug abuse, as well as other forms of drug abuse, requires innovative intervention programs. The first step in successful treatment is detoxification, a process that may be difficult when several drugs with different pharmacologic actions (e.g., alcohol, opiates, and cocaine) have been abused. Because patients may not recall or may deny simultaneous multiple drug use, diagnostic evaluation should always include urinalysis for qualitative detection of psychoactive substances and their metabolites. Treatment of polydrug abuse often requires hospitalization or inpatient residential care during detoxification and the initial phase of drug abstinence. When possible, specialized facilities for the care and treatment of drug-dependent persons should be used. Outpatient detoxification of polydrug abuse patients is unlikely to be effective and may increase risk for dangerous medical consequences.

Drug abuse disorders often respond to effective treatment, but periods of relapse may occur unpredictably. The physician should continue to assist patients during episodes of relapse with compassion and understanding. The physician and the patient must recognize that occasional recurrent drug use is not unusual in this complex behavioral disorder.

The aggregate of particulate matter, after subtracting nicotine and moisture, is referred to as tar.

The alkaline pH of smoke from blends of tobacco used for pipes and cigars allows sufficient absorption of nicotine across the oral mucosa to satisfy the smoker's need for this drug. Therefore, smokers of pipes and cigars tend not to inhale the smoke into the lung, confining the toxic and carcinogenic exposure (and the increased rates of disease) largely to the upper airway for most users of these products. The acidic pH of smoke generated by the tobacco used in cigarettes dramatically reduces absorption of nicotine in the mouth, necessitating inhalation of the smoke into the larger surface of the lungs in order to absorb quantities of nicotine sufficient to satisfy the smoker's addiction. The shift to using tobacco as cigarettes, with resultant increased deposition of smoke in the lung, has created the epidemic of heart disease, lung disease, and lung cancer that dominates the current disease manifestations of tobacco use.

Several genes have been associated with nicotine addiction. Some reduce the clearance of nicotine, and others have been associated with an increased likelihood of becoming dependent on tobacco and other drugs as well as a higher incidence of depression. Rates of smoking cessation have increased, and rates of nicotine addiction have decreased dramatically, since the mid-1950s, suggesting that factors other than genetics are important. It is likely that genetic susceptibility can influence the probability that adolescent experimentation with tobacco will lead to addiction as an adult.

Adult cigarette smoking prevalence has declined to about 19% in the United States, with 20–40% of those smokers not smoking every day. Male smoking prevalence is falling but remains high in most Asian countries, with increasing smoking prevalence among women in those countries. The highest rates of smoking and least cessation are observed in eastern European countries. Of particular concern is the rapidly rising smoking rate observed in the developing world. The World Health Organization Framework Convention on Tobacco Control is encouraging effective tobacco control approaches in these countries with the hope of preventing a future epidemic of tobacco-related illness.

DISEASE MANIFESTATIONS OF CIGARETTE SMOKING

More than 400,000 individuals die prematurely each year in the United States from cigarette use; this represents almost one of every five deaths in the United States. Approximately 40% of cigarette smokers will die prematurely due to cigarette smoking unless they are able to quit.

The major diseases caused by cigarette smoking are listed in [Table 470-1](#). The ratio of smoking-related disease rates in smokers compared to never smokers (relative risk) is greater at younger ages, particularly for coronary artery disease and stroke. At older ages, the background rate of disease in nonsmokers increases, diminishing the fractional contribution of smoking and the relative risk; however, absolute excess rates of disease mortality found in smokers compared to nonsmokers increase with increasing age. The organ damage caused by smoking and the number of smokers who die from smoking are both greater among the elderly, as one would expect from a process of cumulative injury.

CARDIOVASCULAR DISEASES

Cigarette smokers are more likely than nonsmokers to develop both large-vessel atherosclerosis and small-vessel disease. Approximately 90% of peripheral vascular disease in the nondiabetic population can be attributed to cigarette smoking, as can ~50% of aortic aneurysms. In contrast, 20–30% of coronary artery disease and ~10% of ischemic and hemorrhagic strokes are caused by cigarette smoking. There is a multiplicative interaction between cigarette smoking and other cardiac risk factors such that the increment in risk produced by smoking among individuals with hypertension or elevated serum lipids is substantially greater than the increment in risk produced by smoking for individuals without these risk factors.

In addition to its role in promoting atherosclerosis, cigarette smoking also increases the likelihood of myocardial infarction and sudden cardiac death by promoting platelet aggregation and vascular occlusion. Reversal of these effects on coagulation may explain the rapid

470 Nicotine Addiction

David M. Burns

The use of tobacco leaf to create and satisfy nicotine addiction was introduced to Columbus by Native Americans and spread rapidly to Europe. Use of tobacco as cigarettes, however, only became popular in the twentieth century and so is a modern phenomenon, as is the epidemic of disease caused by this form of tobacco use.

Nicotine is the principal constituent of tobacco responsible for its addictive character, but other smoke constituents and behavioral associations contribute to the strength of the addiction. Addicted smokers regulate their nicotine intake by adjusting the frequency and intensity of their tobacco use both to obtain the desired psychoactive effects and avoid withdrawal.

Unburned cured tobacco used orally contains nicotine, carcinogens, and other toxicants capable of causing gum disease, oral and pancreatic cancers, and an increase in the risk of heart disease. When tobacco is burned, the resultant smoke contains, in addition to nicotine, more than 7000 other compounds that result from volatilization, pyrolysis, and pyrosynthesis of tobacco and various chemical additives used in making different tobacco products. The smoke is composed of a fine aerosol and a vapor phase; aerosolized particles are of a size range that results in deposition in the airways and alveolar surfaces of the lungs.

¹Deceased

TABLE 470-1 RELATIVE RISKS FOR CURRENT SMOKERS OF CIGARETTES

Disease or Condition	Current Smokers	
	Males	Females
Coronary heart disease		
Age 35–64	2.8	3.1
Age ≥65	1.5	1.6
Cerebrovascular disease		
Age 35–64	3.3	4
Age ≥65	1.6	1.5
Aortic aneurysm	6.2	7.1
Chronic airway obstruction	10.6	13.1
Cancer		
Lung	23.3	12.7
Larynx	14.6	13
Lip, oral cavity, pharynx	10.9	5.1
Esophagus	6.8	7.8
Bladder, other urinary organs	3.3	2.2
Kidney	2.7	1.3
Pancreas	2.3	2.3
Stomach	2	1.4
Liver	1.7	1.7
Colorectal	1.2	1.2
Cervix		1.6
Acute myeloid leukemia	1.4	1.4
Sudden infant death syndrome		2.3
Infant respiratory distress syndrome		1.3
Low birth weight at delivery		1.8

benefit of smoking cessation for a new coronary event demonstrable among those who have survived a first myocardial infarction. This effect may also explain the substantially higher rates of graft occlusion among continuing smokers following vascular bypass surgery for cardiac or peripheral vascular disease.

Cessation of cigarette smoking reduces the risk of a second coronary event within 6–12 months; rates of first myocardial infarction and death from coronary heart disease also decline within the first few years following cessation among those with no prior cardiovascular history. After 15 years of abstinence, the risk of a new myocardial infarction or death from coronary heart disease in former smokers is similar to that for those who have never smoked.

CANCER

Tobacco smoking causes cancer of the lung; oral cavity; naso-, oro-, and hypopharynx; nasal cavity and paranasal sinuses; larynx; esophagus; stomach; pancreas; liver (hepatocellular); colon and rectum; kidney (body and pelvis); ureter; urinary bladder; and uterine cervix, and also causes myeloid leukemia. There is evidence suggesting that cigarette smoking may play a role in increasing the risk of breast cancer. There does not appear to be a causal link between cigarette smoking and cancer of the endometrium, and there is a lower risk of uterine cancer among postmenopausal women who smoke. The risks of cancer increase with the increasing number of cigarettes smoked per day and with increasing duration of smoking. Additionally, there are synergistic interactions between cigarette smoking and alcohol use for cancer of the oral cavity and esophagus. Several occupational exposures synergistically increase lung cancer risk among cigarette smokers, most notably occupational asbestos and radon exposure.

Cessation of cigarette smoking reduces the risk of developing cancer relative to continuing smoking, but even 20 years after cessation, there is a modest persistent increased risk of developing lung cancer.

RESPIRATORY DISEASE

Cigarette smoking is responsible for 90% of chronic obstructive pulmonary disease. Within 1–2 years of beginning to smoke regularly,

many young smokers will develop inflammatory changes in their small airways, although lung function measures of these changes do not predict development of chronic airflow obstruction. After 20 years of smoking, pathophysiologic changes in the lungs develop and progress proportional to smoking intensity and duration. Chronic mucous hyperplasia of the larger airways results in a chronic productive cough in as many as 80% of smokers >60 years of age. Chronic inflammation and narrowing of the small airways and/or enzymatic digestion of alveolar walls resulting in pulmonary emphysema can result in reduced expiratory airflow sufficient to produce clinical symptoms of respiratory limitation in ~15–25% of smokers.

Changes in the small airways of young smokers will reverse after 1–2 years of cessation. There may also be a small increase in measures of expiratory airflow following cessation among individuals who have developed chronic airflow obstruction, but the major change following cessation is a slowing of the rate of decline in lung function with advancing age rather than a return of lung function toward normal.

PREGNANCY

Cigarette smoking is associated with several maternal complications of pregnancy: premature rupture of membranes, abruptio placentae, and placenta previa; there is also a small increase in the risk of spontaneous abortion among smokers. Infants of smoking mothers are more likely to experience preterm delivery, have a higher perinatal mortality rate, be small for their gestational age, and have higher rates of infant respiratory distress syndrome; they are more likely to die of sudden infant death syndrome and appear to have a developmental lag for at least the first several years of life.

OTHER CONDITIONS

Smoking delays healing of peptic ulcers; increases the risk of developing diabetes, active tuberculosis, rheumatoid arthritis, osteoporosis, senile cataracts, and neovascular and atrophic forms of macular degeneration; and results in premature menopause, wrinkling of the skin, gallstones and cholecystitis in women, and male impotence.

ENVIRONMENTAL TOBACCO SMOKE

Long-term exposure to environmental tobacco smoke increases the risk of lung cancer and coronary artery disease among nonsmokers. It also increases the incidence of respiratory infections, chronic otitis media, and asthma in children and causes exacerbation of asthma in children. Some evidence suggests that environmental tobacco smoke exposure may increase the risk of premenopausal breast cancer. Patients who continue to smoke during treatment for cancer with chemotherapy or radiation have poorer outcomes and reduced survival.

PHARMACOLOGIC INTERACTIONS

Cigarette smoking may interact with a variety of other drugs (Table 470-2). Cigarette smoking induces the cytochrome P450 system, which may alter the metabolic clearance of drugs such as theophylline. This may result in inadequate serum levels in smokers as outpatients when the dosage is established in the hospital under nonsmoking conditions. Correspondingly, serum levels may rise when smokers are hospitalized and not allowed to smoke. Smokers may also have higher first-pass clearance for drugs such as lidocaine, and the stimulant effects of nicotine may reduce the effect of benzodiazepines or beta blockers.

OTHER FORMS OF TOBACCO USE

Other major forms of tobacco use are moist snuff deposited between the cheek and gum, chewing tobacco, pipes and cigars, and recently bidi (tobacco wrapped in tendu or temburni leaf; commonly used in India), clove cigarettes, and water pipes. Oral tobacco use leads to gum disease and can result in oral and pancreatic cancer as well as heart disease, with dramatic differences in the risks evident for products used in Africa and Asia as compared to those in the United States and Europe.

TABLE 470-2 INTERACTIONS OF SMOKING AND PRESCRIPTION DRUGS

Drug	Interaction
Amitriptyline	Increased clearance
Benzodiazepines	Less sedation
Beta blockers	Reduced lowering of heart rate and blood pressure
Caffeine	Faster metabolic clearance
Chlorpromazine	Decreased serum concentrations ^a
Clomipramine	Decreased serum concentrations ^a
Clozapine	Decreased serum concentrations ^a
Dextropropoxyphene	Less analgesia
Estrogens (oral)	Increased hepatic clearance
Flecainide	Increased first-pass clearance
Fluphenazine	Decreased serum concentrations ^a
Fluvoxamine	Decreased serum concentrations ^a
Haloperidol	Decreased serum concentrations ^a
Heparin	Faster clearance
Imipramine	Decreased serum concentrations ^a
Insulin	Delayed absorption due to skin vasoconstriction
Lidocaine	Increased first-pass clearance
Mexiletine	Increased first-pass clearance
Naratriptan	Increased clearance
Olanzapine	Faster clearance
Pentazocine	Less analgesia, possible increased clearance
Propranolol	Increased first-pass clearance
Propoxyphene	Increased hepatic metabolism
Rivastigmine	Increased clearance
Tacrine	Increased clearance
Tacrine	Faster metabolic clearance
Theophylline	Faster metabolic clearance
Thiothixene	Faster metabolic clearance
Trazodone	Decreased serum concentrations ^a
Verapamil	Increased clearance

^aClinical implications uncertain.

All forms of burned tobacco generate toxic and carcinogenic smoke similar to that of cigarette smoke. The differences in disease consequences of use relate to frequency of use and depth of inhalation. The risk of upper airway cancers is similar among cigarette, pipe, and cigar smokers, whereas those who have smoked only pipes and cigars have a much lower risk of lung cancer, heart disease, and chronic obstructive pulmonary disease. However, cigarette smokers who switch to pipes or cigars do tend to inhale the smoke, increasing their risk; and it is likely that comparable inhalation and frequency of exposure to tobacco smoke from any of these forms of tobacco use will lead to comparable disease outcomes.

A resurgence of cigar, bidi, and water pipe use among adolescents of both genders has raised concerns that these older forms of tobacco use are once again causing a public health problem. A variety of devices are currently sold that deliver nicotine by electronically heating materials containing nicotine, the so-called electronic cigarettes. Although these devices are marketed as substitutes for cigarettes and as cessation tools, the composition of the vapor and nicotine delivery varies widely from product to product, raising questions of both safety and efficacy in the absence of regulatory oversight.

LOWER TAR AND NICOTINE CIGARETTES

Filtered cigarettes with lower machine-measured yields of tar and nicotine commonly use ventilation holes in the filters and other engineering designs to artificially lower the machine measurements. Smokers compensate for the lowered nicotine delivery by changing the manner in which they puff on the cigarette or the number of cigarettes smoked per day, and tar and nicotine deliveries are not reduced with use of these products. Cigarette design changes that reduce machine-measured tar

and nicotine lead to deeper inhalation of the smoke and an increase in the carcinogenicity of the smoke inhaled by smokers. The presentation of more carcinogenic smoke to the alveolar portions of the lung has resulted in an increase in the risk of lung cancer, and possibly chronic obstructive pulmonary disease, among smokers over the past six decades. This change in cigarette product is also one cause of the dramatic rise in rates of adenocarcinoma of the lung observed over the past half century. There has been no increase in risk of lung cancer or adenocarcinoma of the lung in never smokers over time.

CESSATION

The process of stopping smoking is commonly a cyclical one, with the smoker sometimes making multiple attempts to quit and failing before finally being successful. Approximately 70–80% of smokers would like to quit smoking. More than one-half of current smokers attempted to quit in the last year, but only 6% quit for 6 months, and only 3% remain abstinent for 2 years. Clinician-based smoking interventions should repeatedly encourage smokers to try to quit and to use different forms of cessation assistance with each new cessation attempt rather than focusing exclusively on immediate cessation at the time of the first visit.

Advice from a physician to quit smoking, particularly at the time of an acute illness, is a powerful trigger for cessation attempts, with up to half of patients who are advised to quit making a cessation effort. Other triggers include the cost of cigarettes, media campaigns, and changes in rules to restrict smoking in the workplace.

PHYSICIAN INTERVENTION (TABLE 470-3)

All patients should be asked whether they smoke, how much they smoke, how long they have smoked, their past experience with quitting, and whether they are currently interested in quitting. Intensity of smoking and smoking within 30 min of waking are useful measures of the intensity of nicotine addiction. Even those who are not interested in quitting should be encouraged and motivated to quit; provided a clear, strong, and personalized message by the clinician that smoking

TABLE 470-3 CLINICAL PRACTICE GUIDELINES

Physician Actions
Ask: Systematically identify all tobacco users at every visit
Advise: Strongly urge all smokers to quit
Identify smokers willing to quit
Assist the patient in quitting
Arrange follow-up contact
Effective Pharmacologic Interventions ^a
First-line therapies
Nicotine gum (1.5)
Nicotine patch (1.9)
Nicotine nasal inhaler (2.7)
Nicotine oral inhaler (2.5)
Nicotine lozenge (2.0)
Bupropion (2.1)
Varenicline (2.7)
Second-line therapies
Clonidine (2.1)
Nortriptyline (3.2)
Other Effective Interventions ^a
Physician or other medical personnel counseling (10 min) (1.3)
Intensive smoking cessation programs (at least 4–7 sessions of 20- to 30-min duration lasting at least 2 and preferably 8 weeks) (2.3)
Clinic-based smoking status identification system (3.1)
Counseling by nonclinicians and social support by family and friends
Telephone counseling (1.2)

^aNumerical value following the intervention is the multiple for cessation success compared to no intervention.

is an important health concern; and offered assistance if they become interested in quitting in the future. Many of those not currently expressing an interest in quitting may nevertheless make an attempt to quit in the subsequent year. For those interested in quitting, a quit date should be negotiated, usually not the day of the visit but within the next few weeks, and a follow-up contact by office staff around the time of the quit date should be provided. There is a relationship between the amount of assistance a patient is willing to accept and the success of the cessation attempt.

There are a variety of nicotine-replacement products, including over-the-counter nicotine patches, gum, and lozenges, as well as nicotine nasal and oral inhalers available by prescription. These products can be used for up to 3–6 months, and some products are formulated to allow a gradual step-down in dosage with increasing duration of abstinence. Antidepressants such as bupropion (300 mg in divided doses for up to 6 months) have also been shown to be effective, as has varenicline, a partial agonist for the nicotinic acetylcholine receptor (initial dose 0.5 mg daily increasing to 1 mg twice daily at day 8; treatment duration up to 6 months). Severe psychiatric symptoms, including suicidal ideation, have been reported with varenicline, resulting in a U.S. Food and Drug Administration–mandated warning and a recommendation for closer therapeutic supervision, but evidence to establish the frequency of these responses and the specificity of their association with varenicline remains unclear. Some evidence supports the combined use of nicotine-replacement therapy (NRT) and antidepressants as well as the use of gum or lozenges for acute cravings in patients using patches. Pretreatment with antidepressants or varenicline is recommended for 1–2 weeks prior to the quit date, and pretreatment with nicotine-replacement products is also being explored, as is longer duration of nicotine replacement as a maintenance therapy for those who are unsuccessful in quitting with a shorter duration of use. NRT is provided in different dosages, with higher doses being recommended for more intense smokers. Clonidine or nortriptyline may be useful for patients who have failed on first-line pharmacologic treatment or who are unable to use other therapies. Antidepressants are more effective in those with a history of depression symptoms.

Current recommendations are to offer pharmacologic treatment, usually with NRT or varenicline, to all who will accept it and to provide counseling and other support as a part of the cessation attempt. There are some data to suggest that longer term use of NRT may enable cessation in some smokers who are unable to quit with shorter duration use and that some individuals are able to achieve abstinence from tobacco through use of NRT chronically. Cessation advice alone by a physician or his or her staff is likely to increase success compared with no intervention; a more comprehensive approach with advice, pharmacologic assistance, and counseling can increase cessation success nearly threefold.

Incorporation of cessation assistance into a practice requires a change of the care delivery infrastructure. Simple changes include (1) adding questions about smoking and interest in cessation on patient-intake questionnaires, (2) asking patients whether they smoke as part of the initial vital sign measurements made by office staff, (3) listing smoking as a problem in the medical record, and (4) automating follow-up contact with the patient on the quit date. These changes are essential to institutionalizing smoking intervention within the practice setting; without this institutionalization, the best intentions of physicians to intervene with their patients who smoke are often lost in the time crush of a busy practice.

PREVENTION

Approximately 85% of individuals who become cigarette smokers initiate the behavior during adolescence. Factors that promote adolescent initiation are parental or older-sibling cigarette smoking, tobacco advertising and promotional activities, the availability of cigarettes, and the social acceptability of smoking. The need for an enhanced self-image and to imitate adult behavior is greatest for those adolescents who have the least external validation of their self-worth, which may explain in part the enormous differences in adolescent smoking prevalence by socioeconomic and school performance strata.

Prevention of smoking initiation must begin early, preferably in the elementary school years. Physicians who treat adolescents should be sensitive to the prevalence of this problem even in the pre-teen population. Physicians should ask all adolescents whether they have experimented with tobacco or currently use tobacco, reinforce the fact that most adolescents and adults do not smoke, and explain that all forms of tobacco are both addictive and harmful.

471e Neuropsychiatric Illnesses in War Veterans

Charles W. Hoge

Neuropsychiatric sequelae are common in combat veterans. Advances in personal protective body armor, armored vehicles, battlefield resuscitation, and the speed of evacuation to tertiary care have considerably improved the survivability of battlefield injuries, resulting in a greater awareness of the “silent wounds” associated with service in a combat zone. Although psychiatric and neurologic problems have been well documented in veterans of prior wars, the conflicts in Iraq and Afghanistan that began after September 11, 2001, were unique in terms of the level of commitment by the U.S. Department of Defense (DoD) and Department of Veterans Affairs (VA), Veterans Health Administration (VHA) to support research as the wars unfolded and to use that knowledge to guide population-level screening, evaluation, and treatment initiatives.

The Iraq and Afghanistan conflicts produced over 2.5 million combat veterans, many of whom have received or will need care in government and civilian medical facilities in the future. Studies clearly showed that service in the Iraq and Afghanistan theaters was associated with significantly elevated rates of mental disorders. Two conditions in particular have been labeled the signature injuries related to these wars: posttraumatic stress disorder (PTSD) and mild traumatic brain injury (mTBI)—also known as concussion. Although particular emphasis will be given in this chapter to PTSD and concussion/mTBI, it is important to understand that the neuropsychiatric sequelae of war are much broader than these two conditions. Wartime service is associated with a number of health concerns that coexist and overlap, and a multidisciplinary patient-centered approach to care is necessary.

EPIDEMIOLOGY OF WAR-RELATED PSYCHOLOGICAL AND NEUROLOGIC CONDITIONS

Service members involved in the Iraq and Afghanistan wars faced multiple deployments to two very different high-intensity combat theaters, and for many veterans, the cumulative strain negatively impacted health, marriages, parenting, educational goals, and civilian occupations. The stresses of service in these conflicts also led to a significant increase in rates of suicide in personnel from the two branches of service involved in the greatest level of ground combat (U.S. Army, Marines).

Service in a war zone can involve extreme physical stress in austere environments, prolonged sleep deprivation, physical injury, exposure to highly life-threatening events, and hazards such as explosive devices, sniper fire, ambushes, indirect fire from rockets and mortars, and chemical pollutants. Certain events, such as loss of a close friend in combat, leave indelible scars. All of these experiences have additive effects on health, likely mediated through physiologic mechanisms involving dysregulation of neuroendocrine and autonomic nervous system (ANS) functions.

Veterans of virtually all wars have reported elevated rates of generalized and multisystem physical, cognitive, and psychological health concerns that often become the focus of treatment months or years after returning home. These multisystem health concerns include sleep disturbance, memory and concentration problems, headaches, musculoskeletal pain, gastrointestinal symptoms (including gastroesophageal reflux), residual effects of wartime injuries, fatigue, anger, hyperarousal symptoms, high blood pressure, rapid heart rate (sometimes associated with panic symptoms), sexual problems, and symptoms associated with PTSD and depression. In order to provide optimal care to veterans with these symptoms, it is important to understand how the symptoms interrelate and to consider the possibility that there may be underlying combat-related physiologic effects.

POSTWAR SYMPTOMS

The overlapping and multisystem health concerns reported by warriors from every generation have been given different labels and have led to debates among medical professionals as to whether these are mediated primarily by physical or psychological causes. For example, World War I produced extensive debate about whether “shell shock,” diagnosed in more than 80,000 British soldiers, was neurologic (“commotional” from the brain being shaken in the skull by concussive blasts) or psychological (“emotional” or “neurasthenia”) in origin. World War II veterans were said to suffer from “battle fatigue,” Korean War veterans developed “combat stress reactions,” and Vietnam veterans developed the “post-Vietnam syndrome.” The role of environmental exposure (e.g., Agent Orange) and psychological causes (e.g., PTSD, depression, substance use disorders) continue to be debated.

Gulf War I (Operation Desert Storm), following the Iraqi invasion of Kuwait in 1990, led to extensive debates as to whether Gulf War syndrome, also known as multisystem illness, was best explained by environmental exposures (e.g., oil fires, depleted uranium, nerve gas, pesticides, multiple vaccinations) or the psychological stress of deployment to a war zone where there was anticipation of high casualty rates from chemical and biologic weapons, repeated stressful alerts, and training exercises involving the use of impermeable full-body protective uniforms (made from rubber, vinyl, charcoal-impregnated polyurethane, and other materials) in desert conditions under extreme temperatures. Although no clinical syndrome was ever definitively confirmed among the nearly 1 million service members who deployed in 1990–1991, studies consistently found that military personnel who served in the Gulf experienced elevations in generalized symptoms across all health domains (e.g., physical, cognitive, neurologic, psychological) compared with service members who deployed elsewhere or did not deploy. In addition, there is good evidence that deployment to the Persian Gulf region during this period was associated with subsequent development of PTSD; other psychiatric disorders including generalized anxiety disorder, depression, and substance use disorders ([Chap. 467](#)); functional gastrointestinal symptoms such as irritable bowel syndrome ([Chap. 352](#)); and chronic fatigue syndrome ([Chap. 464e](#)).

The conflicts in Iraq and Afghanistan led to similar debates as to whether postwar symptoms such as headaches, irritability, sleep disturbance, dizziness, and concentration problems are best attributed to concussion/mTBI or to PTSD. Numerous studies showed that either PTSD or depression explained the majority of the postdeployment “postconcussive” symptoms attributed to concussion/mTBI, a finding not well received by many experts in traumatic brain injury (TBI) but consistent with civilian studies on risk factors for developing persistent symptoms after concussion. As in past wars, the polarized nature of the debate largely focused on only the two conditions of PTSD and concussion/mTBI, which has interfered with full appreciation of how the large spectrum of deployment-related health concerns interrelate and of the clinical implications for designing effective evaluation and treatment strategies.

Veterans understandably may become angry at the suggestion that their postwar health concerns could be “stress-related” or psychological, and thus it is necessary for primary care professionals to be able to discuss the physical toll that war-zone service has on the body, the generalized nature of war-related health concerns, and the likely underlying physiologic neuroendocrine and ANS contributors. Mental health specialists also need to be able to reinforce this message and understand the important role they have in promoting physical health through addressing comorbid health concerns.

PTSD

PTSD ([Chap. 466](#)) is the most common mental disorder documented following war-zone service. Studies from the conflicts in Iraq and Afghanistan found PTSD prevalence rates of 2–6% before deployment (comparable to civilian general population samples) and rates of 6–20% after deployment, depending primarily on the level of combat

frequency and intensity. Many other veterans experience subclinical PTSD symptoms after war-zone service, sometimes termed posttraumatic stress (PTS) or combat stress. These subclinical symptoms can contribute to distress and affect health, even if overall functioning is not as impaired as in the full disorder.

The definition of PTSD was modified in the fifth edition of the American Psychiatric Association's *Diagnostic and Statistical Manual of Mental Disorders* (2013), although most individuals who had PTSD diagnosed according to the previous criteria also meet the definition under the new criteria. PTSD is defined as persistent (>1 month) symptoms occurring after a traumatic event (involving exposure to actual or threatened death, serious injury, or sexual assault). The symptoms must be associated with significant distress or impairment in social or occupational functioning. Symptoms are grouped into four categories: (1) intrusion/reexperiencing symptoms in which the person has nightmares, flashbacks, or intrusive (often involuntary) memories connected with the traumatic event; (2) avoidance symptoms where the person avoids distressing memories or people, places, situations, or other stimuli that serve as reminders of the traumatic event (for example, a crowded mall that triggers heightened alertness to threat); (3) negative alterations of cognitions or mood (for example, feeling detached or losing interest in things that previously brought enjoyment); and (4) hyperarousal symptoms in which the person is physiologically revved up, hyperalert, startles easily, and experiences sleep disturbance, anger, and/or concentration problems. Although PTSD is a clinical symptom-based case definition, it is best to think of PTSD not as an emotional or psychological/psychiatric condition, but rather as a physiologically-based response to life-threatening trauma that is associated with physical, cognitive, emotional, and psychological symptoms.

PTSD has strong biologic correlates, based in fear-conditioning responses to threat and responses to extreme stress involving neuroendocrine dysregulation and ANS reactivity. Numerous studies have shown that PTSD is highly correlated with generalized physical and cognitive symptoms—including hypertension, chronic pain, and cardiovascular disease—as well as cell-mediated immune dysfunction and shortened life expectancy. PTSD is frequently comorbid with other mental disorders such as major depressive disorder, generalized anxiety, substance use disorders, and risky behaviors (e.g., aggression, accidents); it has been estimated that up to 80% of patients with PTSD exhibit one or more comorbid conditions. Misuse of alcohol or substances is most prevalent, often reflecting self-medication. PTSD is also associated with tolerance and withdrawal symptoms related to prescription pain and sleep medications, as well as nicotine dependence (Chap. 470).

Clinicians should understand how to provide meaningful psychological education in a way that resonates with veterans who may have PTSD symptoms as a result of their military service. There is an important occupational context to consider, which is also applicable to trauma exposures that occur in other first responder professions, such as law enforcement officers and firefighters. Service members and other first responders are trained to respond to traumatic events and effectively learn to override automatic fight-or-flight reflexes in order to carry out their duties. Reactions that are labeled as symptoms of PTSD are based on adaptive survival responses that are beneficial in a combat environment. For example, physiologic hyperarousal, use of anger, and being able to shut down other emotions are very useful skills in combat and can be present even prior to traumatic events during tough realistic training. It is natural for these responses to persist after returning home, and the label of a “disorder” only gets applied when the responses that persist significantly impair functioning.

CONCUSSION/mTBI

TBI (Chap. 457e) gained increased recognition during the conflicts in Iraq and Afghanistan because of the widespread exposure of troops to improvised explosive devices. Many veterans of Iraq and Afghanistan reported experiencing multiple concussions during deployments, and many also reported ignoring concussions and not seeking treatment

at the time of injury in order to remain with their unit. However, these legitimate concerns were also counterbalanced and challenged by high prevalence estimates of deployment-related TBI that did not distinguish concussion/mTBI from moderate or severe TBI; data from animal models of blast exposure that did not necessarily extrapolate to human experiences on the battlefield; neuroimaging studies (e.g., diffusion tensor imaging) that attributed putative abnormalities to blast exposure but lacked adequate control comparisons; and fear-provoking speculation that repetitive blast exposure may lead to future dementia, based largely on case series of professional athletes (e.g., boxers, football players) exposed to highly repetitive injuries linked to chronic traumatic encephalopathy (previously termed *dementia pugilistica*) (Chap. 444e).

TBI includes closed and penetrating head injuries; closed head injuries are categorized as mild (mTBI or concussion), moderate, or severe based on the duration of loss of consciousness, duration of posttraumatic amnesia, and the Glasgow coma score (GCS) (see Table 457e-2). Several studies have estimated that 10–20% of all military personnel deployed to Iraq or Afghanistan sustained one or more concussion/mTBI events during deployment, most commonly from exposure to blasts; however, concussion injuries are also common in nondeployed environments from sports, training (e.g., hand-to-hand combatives), and accidents.

Although there is a neurophysiologic continuum of injury, there are stark clinical and epidemiologic distinctions between concussion/mTBI and moderate or severe TBI (Table 471e-1). Concussion/mTBI is defined as a blow or jolt to the head that results in brief loss of consciousness (LOC) for <30 min (most commonly, only a few seconds to minutes), posttraumatic amnesia (PTA) of <24 h (most commonly <1 h), or transient alteration in consciousness (AOC) without LOC. The majority of concussions in Iraq or Afghanistan involved AOC without LOC or PTA (which soldiers may refer to as getting their “bell rung”). GCSs in concussion/mTBI are usually normal (15 out of 15). Concussion is treated with rest to allow the brain time to heal, and

TABLE 471e-1 COMPARISON BETWEEN CONCUSSION/MILD TRAUMATIC BRAIN INJURY (TBI) AND MODERATE/SEVERE TBI

	Mild TBI (concussion)	Moderate/Severe TBI
Clinical case definition		
Loss of consciousness	<30 min (usually a few seconds to minutes)	≥30 min to indefinite
Altered consciousness	<24 h (usually <30 min)	≥24 h to indefinite
Posttraumatic amnesia	<24 h (usually <30 min)	≥24 h to indefinite
Glasgow coma score	13–15 (usually 15)	As low as 3
Focal neurologic signs	None or transient	Frequently present
Traditional neuroimaging (CT/MRI)	Usually negative	Diagnostic
Clinical usefulness of neurocognitive testing after acute injury period	Usually inconclusive	Essential and valuable
Neuronal cell damage	Metabolic/ionic processes associated with axonal swelling, which can lead to disconnection	Direct injury effects plus metabolic/ionic effects
Sequelae, natural history, and recovery	Full recovery expected in majority of individuals; no consensus on natural history; the percentage who develop persistent symptoms is debated	Based directly on injury characteristics; may be severely disabling
Predictors of persistent postconcussive symptoms or disability	Intensely debated; risk factors found to be most predictive include psychiatric conditions (e.g., depression, PTSD) and negative expectations	Not debated; predictors are directly related to injury severity and clinical progress with rehabilitation treatment

Abbreviations: CT, computed tomography; MRI, magnetic resonance imaging; PTSD, post-traumatic stress disorder.

it almost never resulted in air evacuation from Iraq or Afghanistan unless there were other associated injuries.

In contrast, moderate, severe, or penetrating TBI, which is estimated to account for <1% of all battlefield head injuries in Iraq and Afghanistan, is characterized by LOC \geq 30 min (up to permanent coma), PTA \geq 24 h (also may be permanent), and GCSs as low as 3 (the minimum value). These virtually always result in air evacuation from the battlefield and carry a significant risk of severe long-term neurologic impairment and requirement for rehabilitative care.

Symptoms following concussion/mTBI can include headache; fatigue; concentration, memory, or attention problems; sleep disturbance; irritability; balance difficulties; and tinnitus, among other symptoms. Recovery is usually rapid, with symptoms usually resolving in a few hours to days, but in a small percentage of patients, symptoms may persist for a longer period or become chronic (referred to as persistent “postconcussive symptoms” [PCS]).

Establishing a clear causal connection between a deployment concussion injury and persistent PCS months or years after return from deployment has been difficult and often confounded by other postwar conditions that are associated with the same symptoms, including injuries not involving the head, other medical disorders, sleep disorders, PTSD, depression, grief, substance use disorders, chronic pain, and the generalized physiologic effects of wartime service. Contributing to the difficulty in establishing causation is the fact that the concussion/mTBI case definition refers only to the acute injury event and lacks symptoms, time course, or impairment; case definitions for persistent postconcussion syndrome have failed tests of validation. Numerous studies found that PTSD and depression were much stronger predictors of PCS and objective neuropsychological impairment after combat deployment than concussions/mTBIs, and one study even found that bereavement (particularly related to the death of a team member) was as strong a predictor of postdeployment symptoms and poor general health as were symptoms of depression or PTSD. These data do not minimize the importance of concussion/mTBI per se, but highlight the complex interrelationships of war-related health problems and the relatively lower importance of concussion/mTBI in overall postdeployment health than is generally thought.

Studies of veterans who sustained concussions in Iraq or Afghanistan have suggested that blast mechanisms produce similar clinical outcomes as nonblast mechanisms, in contrast to expectations based on some animal models. An explosion can produce serious injury from rapid atmospheric pressure changes (primary blast wave mechanism), as well as from munition fragments/flying debris (secondary blast mechanism) or being thrown into a hard object (tertiary blast mechanism). Secondary and tertiary mechanisms are similar to other mechanical mechanisms of concussions sustained during accidents. It is likely that blast physics explains differences between human clinical studies and experimental animal studies. Because the distribution of munition fragments usually extends well beyond the distribution of the primary blast wave in most explosions, the possibility of a unique head injury solely from the primary blast wave in otherwise uninjured service members appears to be very low.

Multisystem health problems that lack clear case definitions do not lend themselves well to uniform public health strategies such as screening. Nevertheless, mass population screening for concussion/mTBI was mandated for all U.S. service members returning from Iraq or Afghanistan and all veterans presenting for care at VA health care facilities. These screening processes attempt to apply the acute concussion case definition (lacking symptoms, time course, or impairment) months or years after injury, and often involve questions that encourage patients and clinicians to make a direct link between current symptoms and past head injuries that likely have very little to do with the current symptoms. These screening approaches led to sharp criticism that they were encouraging clinicians to misattribute common postwar symptoms to concussion/mTBI. Nevertheless, the screening processes have persisted and are part of an extensive specialty structure of care erected in both the DoD and VA to address health concerns attributed to concussions/mTBIs.

Management of postwar physical and cognitive health concerns is largely symptom focused and ideally carried out within primary care-based structures of care. Studies suggest that optimal strategies for treatment of multisymptom health concerns include regularly scheduled primary care visits with brief physical exam at each visit, protecting patients from unnecessary diagnostic tests and non-evidence-based interventions, judicious use of consultations that protects patients from unnecessary specialty referrals, care/case management, and communication that enhances positive expectations for recovery. Concussion research has shown that negative expectations are one of the most important risk factors for persistent symptoms.

Although many questions remain regarding the long-term health effects of concussions (particularly multiple concussions) sustained during deployment, these are important battlefield injuries that require careful attention. However, they need to be addressed within the context of a much broader approach to other war-related health concerns.

STIGMA AND BARRIERS TO CARE

Stigma and other barriers to care add to the complexity of treating veterans. Despite extensive education efforts among military leaders and service members, perceptions of stigma showed little change over the many years of war; warriors are often concerned that they will be perceived as weak by peers or leaders if they seek care. Studies have shown that less than one-half of service members and veterans with serious mental health problems receive needed care, and upwards of half of those who begin treatment drop out before receiving an adequate number of encounters. Many factors contribute to this, including the pervasive nature of stigma in society in general (particularly among men), the critical importance of group cohesiveness of military teams, the nature of avoidance symptoms in PTSD, perceptions of self-sufficiency (e.g., “I can handle problems on my own”), and sometimes negative perceptions of mental health care and skepticism that mental health professionals will be able to help.

APPROACH TO THE PATIENT:

Evaluation of Veterans with Neuropsychiatric Health Concerns

Evaluation should begin with a careful occupational history as part of the routine medical evaluation; this includes the number of years served, military occupation, deployment locations and dates, illnesses or injuries resulting from service, and significant combat traumatic experiences that may be continuing to affect the individual (Table 471e-2). The clinician should evaluate the degree to which the patient’s current difficulties reflect the normal course of readjusting after the intense occupational experience of combat. It is helpful to reinforce the many strengths associated with being a professional in the military: courage, honor, service to country, resiliency in combat, leadership, ability to work in a cohesive workgroup with peers, and demonstrated skills in handling extreme stress, as well as the fact that reactions that interfere with functioning back home may have their roots in beneficial adaptive physiologic processes.

One of the challenges with current medical practice is that there may be multiple providers with different clinical perspectives. Care should be coordinated through the primary care clinician, with the assistance of a care manager if needed. It is particularly important to continually evaluate all medications prescribed by other practitioners and assess each for possible long-term side effects, dependency, or drug-drug interactions. Particular attention should be given to the level of chronic pain and sleep disturbance, self-medication with alcohol or substances, chronic use of nonsteroidal anti-inflammatory agents (which can contribute to rebound headaches or pain), chronic use of sedative-hypnotic agents, chronic use of narcotic pain medications, and the impact of war-related health concerns on social and occupational functioning.

TABLE 471e-2 SPECIFIC CONSIDERATIONS IN THE MEDICAL EVALUATION OF VETERANS

Occupational context of health concerns	Deployment locations and dates, combat experiences or other deployment stressors, frequent moves, separations from family, impact of deployment on civilian occupation (for reservists)
Medical problems during deployment	History of deployment-related injuries (including concussions), environmental exposures, sleep pattern during deployment, use of caffeine/energy drinks, use of other substances
Current medical history	Current symptoms, level of chronic pain, sleep problems, evidence of persistent physiologic hyperarousal (hypertension, tachycardia, panic symptoms, concentration/memory problems, irritability/anger, sleep disturbance), chronic use of caffeine or energy drinks, chronic use of nonsteroidal anti-inflammatory medications, chronic use of narcotic pain medications, chronic use of nonbenzodiazepine sedative-hypnotic medications, chronic use of benzodiazepines for sleep or anxiety
Mental health assessment	Screen for PTSD, major depressive disorder; ask about suicidal or homicidal ideation, intent, or plans, as well as access to firearms
Alcohol/substance use	Screen for alcohol and substance use disorders, quantity and frequency of use, and evidence of tolerance; inquire about "self-medication" (e.g., use of alcohol to sleep, "calm down," or "forget" war-zone experiences)
Functional impairment	Impact of current symptoms on social and occupational functioning; high-risk behaviors (e.g., drinking and driving, reckless driving, aggression)
Social support, impact of military service on marriage and family	Level of social support; readjustment stress on spouse, children, or other family members

Abbreviation: PTSD, posttraumatic stress disorder.

Screening for PTSD, depression, and alcohol misuse should be performed routinely in all combat veterans. Three screening tools, which are in the public domain, have been validated for use in primary care, and have been used frequently in veterans: the four-question Primary Care PTSD Screen (PC-PTSD), the two-question Patient Health Questionnaire (PHQ-2), and the three-question Alcohol Use Disorders Identification Test-Consumption module (AUDIT-C) (Table 471e-3).

Because the clinical definition of an acute concussion/mTBI does not include symptoms, time course, or impairment, there is currently no clinically validated screening process for use months or years after injury. However, it is important to gather information about all injuries sustained during deployment, including any that resulted in loss or alteration of consciousness or loss of memory around the time of the event. If concussion injuries have occurred, the clinician should assess the number of such injuries, the duration of time unconscious, and injury mechanisms. This should be followed by an assessment of any PCS immediately following the injury event (e.g., headaches, dizziness, tinnitus, nausea, irritability, insomnia, and concentration or memory problems) and the severity and duration of such symptoms.

TREATMENT NEUROPSYCHIATRIC ILLNESSES IN WAR VETERANS

Given the interrelationship of postwar health concerns, care needs to be carefully coordinated. Specific techniques that have been found to be helpful include scheduling regular primary care visits instead of as-needed visits, establishing care management, using good risk-communication principles, establishing a consultative step care approach that draws on the expertise of specialists in a collaborative manner (instead of immediately referring the patient to a specialist and relying on the specialist to provide care), and having behavioral health support directly within primary care clinics (both

for referrals and to provide education and support to primary care professionals prescribing treatment for depression or PTSD).

It is important not to implicitly or explicitly convey the message that physical or cognitive symptoms are psychological or due to "stress." Even if depression or anxiety plays a large role in the etiology of physical health symptoms, the treatment approach should be designed within a patient-centered primary care structure, and referrals should be managed from within this framework. For example, it might help to explain that the primary goal of referral to a mental health professional is to improve sleep and reduce physiologic hyperarousal, which in turn will help with treatment of war-related chronic headaches, concentration problems, or chronic fatigue. If, however, the primary care professional conveys the message that the cause of headaches or concentration problems is anxiety or depression, and this conflicts with the patient's own viewpoint, then this could damage therapeutic rapport and in turn exacerbate the symptoms.

Specific questions related to military service (Table 471e-2) combined with screening for depression, PTSD, and alcohol use disorders (Table 471e-3) should be a routine part of care for all veterans. A positive screen for depression or PTSD should prompt follow-up questions related to these disorders (or use of a longer screening tool such as the nine-question Patient Health Questionnaire or National Center for PTSD Checklist), as well as risk assessment for suicide or homicide. It is important to assess the impact of depression or PTSD symptoms on occupational functioning and interpersonal relationships.

A positive screen for alcohol misuse should prompt a brief motivational intervention that includes bringing attention to the elevated level of drinking, informing the veteran about the effects of alcohol on health, recommending limiting use or abstaining, exploring and setting goals related to drinking behavior, and follow-up and referral to specialty care if needed. This type of brief primary care intervention has been found to be effective and should be incorporated into routine practice. One way to facilitate dialogue about this topic with veterans is to point out how hyperarousal associated with combat service can lead to increased craving for alcohol as the body searches for ways to modulate this. Veterans may consciously or unconsciously drink more to help with sleep, reduce arousal, or avoid thinking about events that happened "downrange." A key educational strategy is to help the veteran to learn that drinking to get to sleep actually damages sleep architecture and makes sleep worse (e.g., reduces rapid eye movement [REM] sleep initially followed by rebound REM activity and early morning waking).

SPECIFIC TREATMENT STRATEGIES FOR PTSD AND COMORBID DEPRESSION

PTSD and depression are highly comorbid in combat veterans, and the evidence-based treatments are similar, involving antidepressant medications, cognitive behavioral therapy (CBT), or both. Psychoeducation that assists veterans to understand that their symptoms of PTSD have a basis in adaptive survival mechanisms and skills they exhibited in combat can facilitate therapeutic rapport. Remaining hypervigilant to threat, being able to shut down emotions, being able to function on less sleep, and using anger to help focus and control fear are all adaptive beneficial survival skills in a combat environment. Therefore, PTSD for warriors is both a medical disorder and a set of reactions that have their roots in the physiologic adaptation and skills they successfully applied in combat.

It is important to know that combat is not the only important trauma in a war-zone environment. Rape, assault, and accidents also occur. Rape or assault by a fellow service member, which affects a greater number of women veterans, but also occurs in men, can be particularly devastating because it destroys the vital feeling of safety that individuals derive from their own unit peers in a war environment.

The treatments for PTSD considered by most consensus guideline committees to have an A level of evidence include CBTs and medications, specifically selective serotonin reuptake inhibitors (SSRIs) and serotonin norepinephrine reuptake inhibitors (SNRIs), with the strongest evidence from double-blind, placebo-controlled studies

TABLE 471e-3 PRIMARY CARE MENTAL HEALTH SCREENING TOOLS**PC-PTSD Screen**

1. Have you ever had any experience that was so frightening, horrible, or upsetting that, in the past month, you:

Have had nightmares about it or thought about it when you did not want to?	Yes	No
Tried hard not to think about it or went out of your way to avoid situations that remind you of it?	Yes	No
Were constantly on guard, watchful, or easily startled?	Yes	No
Felt numb or detached from others, activities, or your surroundings?	Yes	No

Note: Two or more "yes" responses (three or more a more specific cutoff) are considered a positive screen.**Source:** A Prins et al: The Primary Care PTSD Screen (PC-PTSD): Development and operating characteristics. Prim Care Psychiatr 9:9, 2004.**PHQ-2 Depression Screen**

2. Over the last 2 weeks, how often have you been bothered by any of the following problems?	Not at all (0)	Few or several days (1)	More than half the days (2)	Nearly every day (3)
Little interest or pleasure in doing things.	0	1	2	3
Feeling down, depressed, or hopeless.	0	1	2	3

Note: If either (or both) questions are marked 2 or 3 (more than half the days or higher), this is considered a positive screen for depression.**Source:** K Kroenke et al: The Patient Health Questionnaire-2: Validity of a two-item depression screener. Med Care 41:1284, 2003.**AUDIT-C Alcohol Screen**

3a. How often do you have a drink containing alcohol?

Never (0)	Monthly or less (1)	Two or four times a month (2)	Two to three times per week (3)	Four or more times a week (4)
-----------	---------------------	-------------------------------	---------------------------------	-------------------------------

3b. How many drinks containing alcohol do you have on a typical day when you are drinking?

1 or 2 (0)	3 or 4 (1)	5 or 6 (2)	7 or 9 (3)	10 or more (4)
------------	------------	------------	------------	----------------

3c. How often do you have six or more drinks on one occasion?

Never (0)	Less than Monthly (1)	Monthly (2)	Two to three times per week (3)	Four or more times a week (4)
-----------	-----------------------	-------------	---------------------------------	-------------------------------

Note: A positive AUDIT-C screen is defined as a total score for men ≥ 4 ; for women ≥ 3 . A report of drinking 6 or more drinks on one occasion should prompt an in-depth assessment of drinking.**Source:** K Bush et al: The AUDIT Alcohol Consumption Questions (AUDIT-C): An effective brief screening test for problem drinking. Arch Intern Med 158:1789, 1998.

for sertraline, paroxetine, fluoxetine, and venlafaxine (of which paroxetine and sertraline received U.S. Food and Drug Administration approval for PTSD). (See Table 466-3 for recommended dosages.) Prazosin has also gained very strong evidence recently through randomized placebo-controlled studies for its effectiveness in controlling nightmares as well as global PTSD symptoms, through modulation of the physiologic processes associated with PTSD.

CBT interventions include narrative therapy (often called "imaginal exposure"), in vivo exposure focused on retraining the body not to react to stimuli related to traumatic reminders (e.g., a crowded mall), and techniques to modulate physiologic hyperarousal (e.g., diaphragmatic breathing, progressive muscle relaxation). A number of complementary alternative medicine approaches including acupuncture, mindfulness meditation, yoga, and massage are also being used in PTSD. Although not evidence-based treatments per se, if they facilitate a relaxation response and alleviation of hyperarousal or sleep symptoms, they can be considered useful adjunctive modalities.

There have been no head-to-head comparisons of medication compared with psychotherapy for treatment of PTSD. It is reasonable for primary care clinicians to consider initiating treatment for mild to moderate PTSD symptoms with an SSRI and to refer patients to a mental health professional if there are more severe symptoms, significant comorbidity, safety concerns, or limited response to initial treatment. All PTSD treatments are associated with a sizable proportion of individuals who fail to respond adequately, and it is often necessary to add modalities or switch treatment. SNRIs may be useful alternatives to SSRIs if there has been nonresponse or side effects with SSRIs or if there is comorbid pain (duloxetine, in particular, has indications for pain). Both SSRIs and SNRIs can increase anxiety initially; patients should be warned about this possibility, and treatment should be initiated with the lowest recommended dose (or even one-half of the lowest dose for a few days) and gradually

increased thereafter. Antidepressants also are likely to be useful in comorbid depression, which is common in veterans with PTSD. All antidepressants have potential drug-drug interactions that must be considered.

Many other medications have been used in PTSD, including tricyclic antidepressants, benzodiazepines, atypical antipsychotics, and anticonvulsants. In general, these should be prescribed in conjunction with psychiatric consultation because of their greater side effects and risks. Benzodiazepines, in particular, should be avoided in the treatment of PTSD. Studies have shown that they do not reduce core PTSD symptoms, are likely to exacerbate substance use disorders that are common in veterans with PTSD, and may produce significant rebound anxiety and anger. Individuals with PTSD often report symptomatic relief upon initiation of a benzodiazepine, but this is generally short lived and associated with a high risk of tolerance and dependence that can worsen recovery. Atypical antipsychotics, which have gained widespread popularity as adjunctive treatment for depression, anxiety, or sleep problems, have significant long-term side effects, including metabolic effects (e.g., glucose dysregulation), weight gain, and cardiovascular risks.

Sleep disturbance should be addressed initially with sleep hygiene education, followed by consideration of an antihistamine, trazodone, low-dose mirtazapine, or nonbenzodiazepine sedative-hypnotic such as zolpidem, eszopiclone, or zaleplon. However, the nonbenzodiazepine sedative-hypnotics should be used with caution in veterans because they can lead to tolerance and rebound sleep problems similar to those seen with benzodiazepine use.

TREATMENT STRATEGIES FOR CONCUSSION/mTBI AND POSTDEPLOYMENT POSTCONCUSSIVE SYMPTOMS

Concussion/mTBI is best treated at the time of injury with education and rest to allow time for the brain to heal and protect against a second impact syndrome (a rare but life-threatening event involving

471e-6 brain swelling that can occur when a second concussion occurs before the brain has adequately healed from an initial event). Randomized trials have shown that education regarding concussion that informs the patient of what to expect and promotes the expectation of recovery is the most effective treatment in preventing persistent symptoms.

Once service members return from deployment and seek care for postwar health problems, treatment is largely symptom focused, following the principles of patient-centered and collaborative care models. Cognitive rehabilitation, which is very useful in moderate and severe TBI to improve memory, attention, and concentration, has generally not been shown to be effective for mTBI in randomized clinical studies, although consensus groups have supported its use.

General recommendations for the clinical management of persistent, chronic PCS include treating physical and cognitive health problems based on symptom presentation, coexisting health problems, and individual preferences; and addressing coexisting depression, PTSD, substance use disorders, or other factors that may be contributing to symptom persistence. Headache is the most common

symptom associated with concussion/mTBI, and the evaluation and treatment of headache parallels that for other causes of headache (**Chaps. 21 and 447**). Stimulant medications for alleviating neurocognitive effects attributed to concussion/mTBI are not recommended. Clinicians should be aware of the potential for cognitive or sedative side effects of certain medications that may be prescribed for depression, anxiety, sleep, or chronic pain.

Treatment of neuropsychiatric problems must be coordinated with care for other war-related health concerns, with the goal of treatment to reduce the severity of symptoms, improve social and occupational functioning, and prevent long-term disability. Understanding the occupational context of war-related health concerns is important in communicating with veterans and developing a comprehensive treatment strategy.

DISCLOSURE

The opinions or assertions contained herein are the private views of the author and are not to be construed as official, or as reflecting the views of the Department of the Army or the Department of Defense.

472e Heavy Metal Poisoning

Howard Hu

Metals pose a significant threat to health through low-level environmental as well as occupational exposures. One indication of their importance relative to other potential hazards is their ranking by the U.S. Agency for Toxic Substances and Disease Registry, which maintains an updated list of all hazards present in toxic waste sites according to their prevalence and the severity of their toxicity. The first, second, third, and seventh hazards on the list are heavy metals: lead, mercury, arsenic, and cadmium, respectively (<http://www.atsdr.cdc.gov/spl/>). Specific information pertaining to each of these metals, including sources and metabolism, toxic effects produced, diagnosis, and the appropriate treatment for poisoning, is summarized in [Table 472e-1](#).

Metals are inhaled primarily as dusts and fumes (the latter defined as tiny particles generated by combustion). Metal poisoning can also result from exposure to vapors (e.g., mercury vapor in creating dental amalgams). When metals are ingested in contaminated food or drink or by hand-to-mouth activity (implicated especially in children), their gastrointestinal absorption varies greatly with the specific chemical form of the metal and the nutritional status of the host. Once a metal is absorbed, blood is the main medium for its transport, with the precise kinetics dependent on diffusibility, protein binding, rates of biotransformation, availability of intracellular ligands, and other factors. Some organs (e.g., bone, liver, and kidney) sequester metals in relatively high concentrations for years. Most metals are excreted through renal clearance and gastrointestinal excretion; some proportion is also excreted through salivation, perspiration, exhalation, lactation, skin exfoliation, and loss of hair and nails. The intrinsic stability of metals facilitates tracing and measurement in biologic material, although the clinical significance of the levels measured is not always clear.

Some metals, such as copper and selenium, are essential to normal metabolic function as trace elements ([Chap. 96e](#)) but are toxic at high levels of exposure. Others, such as lead and mercury, are xenobiotic and theoretically are capable of exerting toxic effects at any level of exposure. Indeed, much research is currently focused on the contribution of low-level xenobiotic metal exposure to chronic diseases and to subtle changes in health that may have significant public health consequences. Genetic factors, such as polymorphisms that encode for variant enzymes with altered properties in terms of metal binding, transport, and effects, also may modify the impact of metals on health and thereby account, at least in part, for individual susceptibility to metal effects.

The most important component of treatment for metal toxicity is the termination of exposure. *Chelating agents* are used to bind metals into stable cyclic compounds with relatively low toxicity and to enhance their excretion. The principal chelating agents are dimercaprol (British anti-Lewisite [BAL]), ethylenediamine tetraacetic acid (EDTA), succimer (dimercaptosuccinic acid [DMSA]), and penicillamine; their specific use depends on the metal involved and the clinical circumstances. Activated charcoal does not bind metals and thus is of limited usefulness in cases of acute metal ingestion.

In addition to the information provided in [Table 472e-1](#), several other aspects of exposure, toxicity, or management are worthy of discussion with respect to the four most hazardous toxicants (arsenic, cadmium, lead, and mercury).

Arsenic, even at moderate levels of exposure, has been clearly linked with increased risks for cancer of the skin, bladder, renal pelvis, ureter, kidney, liver, and lung. These risks appear to be modified by smoking, folate and selenium status, genetic traits (such as ability to methylate arsenic), and other factors. Studies in community-based populations are beginning to demonstrate that arsenic exposure is a risk factor for increased coronary heart disease and stroke. Evidence is also emerging

that low-level arsenic may cause neurodevelopmental delays in children and possibly diabetes, but the evidence remains uneven.

Serious *cadmium* poisoning from the contamination of food and water by mining effluents in Japan contributed to the 1946 outbreak of “itai-itai” (“ouch-ouch”) disease, so named because of cadmium-induced bone toxicity that led to painful bone fractures. Modest exposures from environmental contamination have recently been associated in some studies with a lower bone density, a higher incidence of fractures, and a faster decline in height in both men and women, effects that may be related to cadmium’s calciuric effect on the kidney. There is some evidence for synergy between the adverse impacts of cadmium and lead on kidney function. Environmental exposures have also been linked to lower lung function (even after adjusting for smoking cigarettes, which contain cadmium) as well as increased risk of cardiovascular disease and mortality, stroke, and heart failure. Several studies have also raised concerns that cadmium may be carcinogenic and contribute to elevated risks of prostate, breast, and pancreatic cancer. Overall, this growing body of research indicates that cadmium exposure may be contributing significantly to morbidity and mortality rates in the general population.

Advances in our understanding of *lead* toxicity have recently benefited by the development of K x-ray fluorescence (KXRF) instruments for making safe in vivo measurements of lead levels in bone (which, in turn, reflect cumulative exposure over many years, as opposed to blood lead levels, which mostly reflect recent exposure). Higher bone lead levels measured by KXRF have been linked to increased risk of hypertension and accelerated declines in cognition in both men and women from an urban population. Upon reviewing these studies in conjunction with other epidemiologic and toxicologic studies, a recent federal expert panel concluded that the impact of lead exposure on hypertension and cognition in adults was causal. Prospective studies have also demonstrated that higher bone lead levels are a major risk factor for increased cardiovascular morbidity and mortality rates in both community-based and occupational-exposed populations. Lead exposure at community levels has also been recently associated with increased risks of hearing loss, Parkinson’s disease, and amyotrophic lateral sclerosis. With respect to pregnancy-associated risks, high maternal bone lead levels were found to predict lower birth weight, head circumference, birth length, and neurodevelopmental performance in offspring by age 2 years. In a randomized trial, calcium supplementation (1200 mg daily) was found to significantly reduce the mobilization of lead from maternal bone into blood during pregnancy.

The toxicity of low-level organic *mercury* exposure (as manifested by neurobehavioral performance) is of increasing concern based on studies of the offspring of mothers who ingested mercury-contaminated fish. With respect to whether the consumption of fish by women during pregnancy is good or bad for offspring neurodevelopment, balancing the trade-offs of the beneficial effects of the omega-3-fatty acids (FAs) in fish versus the adverse effects of mercury contamination in fish has led to some confusion and inconsistency in public health recommendations. Overall, it would appear that it would be best for pregnant women to either limit fish consumption to those species known to be low in mercury contamination but high in omega-3-FAs (such as sardines or mackerel) or to avoid fish and obtain omega-3-FAs through supplements or other dietary sources. Current evidence has not supported the recent contention that ethyl mercury, used as a preservative in multiuse vaccines administered in early childhood, has played a significant role in causing neurodevelopmental problems such as autism. With regard to adults, there is conflicting evidence as to whether mercury exposure is associated with increased risk of hypertension and cardiovascular disease. At this point, conclusions cannot be drawn.

Heavy metals pose risks to health that are especially burdensome in selected parts of the world. For example, *arsenic* exposure from natural contamination of shallow tube wells inserted for drinking water is a major environmental problem for millions of



TABLE 472e-1 HEAVY METALS

Main Sources	Metabolism	Toxicity	Diagnosis	Treatment
Arsenic				
Smelting and micro-electronics industries; wood preservatives, pesticides, herbicides, fungicides; contaminant of deep-water wells; folk remedies; and coal; incineration of these products.	Organic arsenic (arsenobetaine, arsenocholine) is ingested in seafood and fish, but is nontoxic; inorganic arsenic is readily absorbed (lung and GI); sequesters in liver, spleen, kidneys, lungs, and GI tract; residues persist in skin, hair, and nails; biomethylation results in detoxification, but this process saturates.	Acute arsenic poisoning results in necrosis of intestinal mucosa with hemorrhagic gastroenteritis, fluid loss, hypotension, delayed cardiomyopathy, acute tubular necrosis, and hemolysis. Chronic arsenic exposure causes diabetes, vasospasm, peripheral vascular insufficiency and gangrene, peripheral neuropathy, and cancer of skin, lung, liver (angiosarcoma), bladder, and kidney. Lethal dose: 120–200 mg (adults); 2 mg/kg (children).	Nausea, vomiting, diarrhea, abdominal pain, delirium, coma, seizures; garlicky odor on breath; hyperkeratosis, hyperpigmentation, exfoliative dermatitis, and Mees' lines (transverse white striae of the fingernails); sensory and motor polyneuritis, distal weakness. Radiopaque sign on abdominal x-ray; ECG—QRS broadening, QT prolongation, ST depression, T-wave flattening; 24-h urinary arsenic >67 $\mu\text{mol/d}$ or 50 $\mu\text{g/d}$; (no seafood \times 24 h); if recent exposure, serum arsenic >0.9 $\mu\text{mol/L}$ (7 $\mu\text{g/dL}$). High arsenic in hair or nails.	If acute ingestion, ipecac to induce vomiting, gastric lavage, activated charcoal with a cathartic. Supportive care in ICU. Dimercaprol 3–5 mg/kg IM q4h \times 2 days; q6h \times 1 day, then q12h \times 10 days; alternative: oral succimer.
Cadmium				
Metal-plating, pigment, smelting, battery, and plastics industries; tobacco; incineration of these products; ingestion of food that concentrates cadmium (grains, cereals).	Absorbed through ingestion or inhalation; bound by metallothionein, filtered at the glomerulus, but reabsorbed by proximal tubules (thus, poorly excreted). Biologic half-life: 10–30 y. Binds cellular sulfhydryl groups, competes with zinc, calcium for binding sites. Concentrates in liver and kidneys.	Acute cadmium inhalation causes pneumonitis after 4–24 h; acute ingestion causes gastroenteritis. Chronic exposure causes anosmia, yellowing of teeth, emphysema, minor LFT elevations, microcytic hypochromic anemia unresponsive to iron therapy, proteinuria, increased urinary β_2 -microglobulin, calciuria, leading to chronic renal failure, osteomalacia, and fractures. Possible risks of cardiovascular disease and cancer.	With inhalation: pleuritic chest pain, dyspnea, cyanosis, fever, tachycardia, nausea, noncardiogenic pulmonary edema. With ingestion: nausea, vomiting, cramps, diarrhea. Bone pain, fractures with osteomalacia. If recent exposure, serum cadmium >500 nmol/L (5 $\mu\text{g/dL}$). Urinary cadmium >100 nmol/L (10 $\mu\text{g/g}$ creatinine) and/or urinary β_2 -microglobulin >750 $\mu\text{g/g}$ creatinine (but urinary β_2 -microglobulin also increased in other renal diseases such as pyelonephritis).	There is no effective treatment for cadmium poisoning (chelation not useful; dimercaprol can exacerbate nephrotoxicity). Avoidance of further exposure, supportive therapy, vitamin D for osteomalacia.
Lead				
Manufacturing of auto batteries, lead crystal, ceramics, fishing weights, etc.; demolition or sanding of lead-painted houses, bridges; stained glass-making, plumbing, soldering; environmental exposure to paint chips, house dust (in homes built <1975), firing ranges (from bullet dust), food or water from improperly glazed ceramics, lead pipes; contaminated herbal remedies, candies; exposure to the combustion of leaded fuels.	Absorbed through ingestion or inhalation; organic lead (e.g., tetraethyl lead) absorbed dermally. In blood, 95–99% sequestered in RBCs—thus, must measure lead in whole blood (not serum). Distributed widely in soft tissue, with half-life \sim 30 days; 15% of dose sequestered in bone with half-life of >20 years. Excreted mostly in urine, but also appears in other fluids including breast milk. Interferes with mitochondrial oxidative phosphorylation, ATPases, calcium-dependent messengers; enhances oxidation and cell apoptosis.	Acute exposure with blood lead levels (BPb) of >60–80 $\mu\text{g/dL}$ can cause impaired neurotransmission and neuronal cell death (with central and peripheral nervous system effects); impaired hematopoiesis and renal tubular dysfunction. At higher levels of exposure (e.g., BPb >80–120 $\mu\text{g/dL}$), acute encephalopathy with convulsions, coma, and death may occur. Subclinical exposures in children (BPb 25–60 $\mu\text{g/dL}$) are associated with anemia; mental retardation; and deficits in language, motor function, balance, hearing, behavior, and school performance. Impairment of IQ appears to occur at even lower levels of exposure with no measurable threshold above the limit of detection in most assays of 1 $\mu\text{g/dL}$. In adults, chronic subclinical exposures (BPb >40 $\mu\text{g/dL}$) are associated with an increased risk of anemia, demyelinating peripheral neuropathy (mainly motor), impairments of reaction time and hearing, accelerated declines in cognition, hypertension, ECG conduction delays, higher risk of cardiovascular disease and death, interstitial nephritis and chronic renal failure, diminished sperm counts, and spontaneous abortions.	Abdominal pain, irritability, lethargy, anorexia, anemia, Fanconi's syndrome, pyuria, azotemia in children with blood lead level (BPb) >80 $\mu\text{g/dL}$; may also see epiphyseal plate "lead lines" on long bone x-rays. Convulsions, coma at BPb >120 $\mu\text{g/dL}$. Noticeable neurodevelopmental delays at BPb of 40–80 $\mu\text{g/dL}$; may also see symptoms associated with higher BPb levels. Screening of all U.S. children when they begin to crawl (\sim 6 months) is recommended by the CDC; source identification and intervention is begun if the BPb >10 $\mu\text{g/dL}$. In adults, acute exposure causes similar symptoms as in children as well as headaches, arthralgias, myalgias, depression, impaired short-term memory, loss of libido. Physical exam may reveal a "lead line" at the gingiva-tooth border, pallor, wrist drop, and cognitive dysfunction (e.g., declines on the mini-mental state exam); lab tests may reveal a normocytic, normochromic anemia, basophilic stippling, an elevated blood protoporphyrin level (free erythrocyte or zinc), and motor delays on nerve conduction. U.S. OSHA requires regular testing of lead-exposed workers with removal if BPb >40 $\mu\text{g/dL}$. New guidelines have been proposed recommending that BPb be maintained at <10 $\mu\text{g/dL}$, removal of workers if BPb >20 $\mu\text{g/dL}$, and monitoring of cumulative exposure parameters.	Identification and correction of exposure sources is critical. In some U.S. states, screening and reporting to local health boards of children with BPb >10 $\mu\text{g/dL}$ and workers with BPb >40 $\mu\text{g/dL}$ is required. In the highly exposed individual with symptoms, chelation is recommended with oral DMSA (succimer); if acutely toxic, hospitalization and IV or IM chelation with ethylenediamine tetraacetic acid calcium disodium (CaEDTA) may be required, with the addition of dimercaprol to prevent worsening of encephalopathy. It is uncertain whether children with asymptomatic lead exposure (e.g., BPb 20–40 $\mu\text{g/dL}$) benefit from chelation; a recent randomized trial showed no benefit. Correction of dietary deficiencies in iron, calcium, magnesium, and zinc will lower lead absorption and may also improve toxicity. Vitamin C is a weak but natural chelating agent. Calcium supplements (1200 mg at bedtime) have been shown to lower blood lead levels in pregnant women.

(Continued)

TABLE 472e-1 HEAVY METALS (CONTINUED)

Main Sources	Metabolism	Toxicity	Diagnosis	Treatment
Mercury				
<p>Metallic, mercurous, and mercuric mercury (Hg, Hg⁺, Hg²⁺) exposures occur in some chemical, metal-processing, electrical-equipment, automotive industries; they are also in thermometers, dental amalgams, batteries.</p> <p>Mercury is dispersed by waste incineration. Environmental bacteria convert inorganic to organic mercury, which then bioconcentrates up the aquatic food chain to contaminate tuna, swordfish, and other pelagic fish.</p>	<p>Elemental mercury (Hg) is not well absorbed; however, it will volatilize into highly absorbable vapor. Inorganic mercury is absorbed through the gut and skin. Organic mercury is well absorbed through inhalation and ingestion. Elemental and organic mercury cross the blood-brain barrier and placenta. Mercury is excreted in urine and feces and has a half-life in blood of ~60 days; however, deposits will remain in the kidney and brain for years. Exposure to mercury stimulates the kidney to produce metallothionein, which provides some detoxification benefit. Mercury binds sulfhydryl groups and interferes with a wide variety of critical enzymatic processes.</p>	<p>Acute inhalation of Hg vapor causes pneumonitis and noncardiogenic pulmonary edema leading to death, CNS symptoms, and polyneuropathy.</p> <p>Chronic high exposure causes CNS toxicity (mercurial <i>erethism</i>; see Diagnosis); lower exposures impair renal function, motor speed, memory, coordination.</p> <p>Acute ingestion of inorganic mercury causes gastroenteritis, the nephritic syndrome, or acute renal failure, hypertension, tachycardia, and cardiovascular collapse, with death at a dose of 10–42 mg/kg.</p> <p>Ingestion of organic mercury causes gastroenteritis, arrhythmias, and lesions in the basal ganglia, gray matter, and cerebellum at doses >1.7 mg/kg.</p> <p>High exposure during pregnancy causes derangement of fetal neuronal migration resulting in severe mental retardation.</p> <p>Mild exposures during pregnancy (from fish consumption) are associated with declines in neurobehavioral performance in offspring.</p> <p>Dimethylmercury, a compound only found in research labs, is “supertoxic”—a few drops of exposure via skin absorption or inhaled vapor can cause severe cerebellar degeneration and death.</p>	<p>Chronic exposure to metallic mercury vapor produces a characteristic intention tremor and mercurial <i>erethism</i>: excitability, memory loss, insomnia, timidity, and delirium (“mad as a hatter”). On neurobehavioral tests: decreased motor speed, visual scanning, verbal and visual memory, visuomotor coordination.</p> <p>Children exposed to mercury in any form may develop <i>acrodyndia</i> (“pink disease”): flushing, itching, swelling, tachycardia, hypertension, excessive salivation or perspiration, irritability, weakness, morbilliform rashes, desquamation of palms and soles.</p> <p>Toxicity from elemental or inorganic mercury exposure begins when blood levels >180 nmol/L (3.6 µg/dL) and urine levels >0.7 µmol/L (15 µg/dL). Exposures that ended years ago may result in a >20-µg increase in 24-h urine after a 2-g dose of succimer.</p> <p>Organic mercury exposure is best measured by levels in blood (if recent) or hair (if chronic); CNS toxicity in children may derive from fetal exposures associated with maternal hair Hg >30 nmol/g (6 µg/g).</p>	<p>Treat acute ingestion of mercuric salts with induced emesis or gastric lavage and polythiol resins to bind mercury in the GI tract). Chelate with dimercaprol (up to 24 mg/kg per day IM in divided doses), DMSA (succimer), or penicillamine, with 5-day courses separated by several days of rest. If renal failure occurs, treat with peritoneal dialysis, hemodialysis, or extracorporeal regional complexing hemodialysis and succimer.</p> <p>Chronic inorganic mercury poisoning is best treated with <i>N</i>-acetyl penicillamine.</p>

Abbreviations: ATPase, adenosine triphosphatase; BPb, blood lead; CDC, Centers for Disease Control and Prevention; CNS, central nervous system; DMSA, dimercaptosuccinic acid; ECG, electrocardiogram; GI, gastrointestinal; ICU, intensive care unit; IQ, intelligence quotient; LFT, liver function tests; OSHA, Occupational Safety and Health Administration; RBC, red blood cell.

residents in parts of Bangladesh and Western India. Contamination was formerly considered only a problem with deep wells; however, the geology of this region allows most residents only a few alternatives for potable drinking water. The combustion of leaded gasoline with resulting contamination of air and soil with *lead oxide* remains a problem in some countries of Central Asia, Southeast Asia, Africa, and the Middle East. Populations living in the Arctic have been shown to have particularly high exposures to *mercury* due to long-range transport patterns that concentrate mercury in the polar regions, as well as the traditional dependence of Arctic peoples on the consumption of fish and other wildlife that bioconcentrate methylmercury.

A few additional metals deserve brief mention but are not covered in Table 472e-1 because of the relative rarity of their being clinically encountered or the uncertainty regarding their potential toxicities. *Aluminum* contributes to the encephalopathy in patients with severe renal disease, who are undergoing dialysis (Chap. 424). High levels of aluminum are found in the neurofibrillary tangles in the cerebral cortex and hippocampus of patients with Alzheimer’s disease, as well as in the drinking water and soil of areas with an unusually high incidence of Alzheimer’s. The experimental and epidemiologic evidence for the aluminum–Alzheimer’s disease link remains relatively weak, however, and it cannot be concluded that aluminum is a causal agent or a contributing factor in neurodegenerative disease. Hexavalent chromium is corrosive and sensitizing. Workers in the chromate and chrome pigment production industries have consistently had a greater risk of lung cancer. The introduction of *cobalt* chloride as a fortifier in beer led to outbreaks of fatal cardiomyopathy among heavy consumers. Occupational exposure (e.g., of miners, dry-battery manufacturers, and arc welders) to *manganese* can cause a parkinsonian syndrome within 1–2 years, including gait disorders; postural instability; a

masked, expressionless face; tremor; and psychiatric symptoms. With the introduction of methylcyclopentadienyl manganese tricarbonyl (MMT) as a gasoline additive, there is concern for the toxic potential of environmental manganese exposure. For example, a recent study found a high prevalence of parkinsonian disorders in a community with risks proportionate to estimated manganese exposures emitted by local ferroalloy industries. Epidemiologic studies have also suggested that manganese may interfere with early childhood neurodevelopment in ways similar to that of lead. *Nickel* exposure induces an allergic response, and inhalation of nickel compounds with low aqueous solubility (e.g., nickel subsulfide and nickel oxide) in occupational settings is associated with an increased risk of lung cancer. Overexposure to selenium may cause local irritation of the respiratory system and eyes, gastrointestinal irritation, liver inflammation, loss of hair, depigmentation, and peripheral nerve damage. Workers exposed to certain organic forms of *tin* (particularly trimethyl and triethyl derivatives) have developed psychomotor disturbances, including tremor, convulsions, hallucinations, and psychotic behavior.

Thallium, which is a component of some insecticides, metal alloys, and fireworks, is absorbed through the skin as well as by ingestion and inhalation. Severe poisoning follows a single ingested dose of >1 g or >8 mg/kg. Nausea and vomiting, abdominal pain, and hematemesis precede confusion, psychosis, organic brain syndrome, and coma. Thallium is radiopaque. Induced emesis or gastric lavage is indicated within 4–6 h of acute ingestion; Prussian blue prevents absorption and is given orally at 250 mg/kg in divided doses. Unlike other types of metal poisoning, thallium poisoning may be less severe when activated charcoal is used to interrupt its enterohepatic circulation. Other measures include forced diuresis, treatment with potassium chloride (which promotes renal excretion of thallium), and peritoneal dialysis.

Poisoning refers to the development of dose-related adverse effects following exposure to chemicals, drugs, or other xenobiotics. To paraphrase Paracelsus, the dose makes the poison. In excessive amounts, substances that are usually innocuous, such as oxygen and water, can cause toxicity. Conversely, in small doses, substances commonly regarded as poisons, such as arsenic and cyanide, can be consumed without ill effect. Although most poisons have predictable dose-related effects, individual responses to a given dose may vary because of genetic polymorphism, enzymatic induction or inhibition in the presence of other xenobiotics, or acquired tolerance. Poisoning may be local (e.g., skin, eyes, or lungs) or systemic depending on the route of exposure, the chemical and physical properties of the poison, and its mechanism of action. The severity and reversibility of poisoning also depend on the functional reserve of the individual or target organ, which is influenced by age and preexisting disease.

EPIDEMIOLOGY

More than 5 million poison exposures occur in the United States each year. Most are acute, are accidental (unintentional), involve a single agent, occur in the home, result in minor or no toxicity, and involve children <6 years of age. Pharmaceuticals are involved in 47% of exposures and in 84% of serious or fatal poisonings. Unintentional exposures can result from the improper use of chemicals at work or play; label misreading; product mislabeling; mistaken identification of unlabeled chemicals; uninformed self-medication; and dosing errors by nurses, pharmacists, physicians, parents, and the elderly. Excluding the recreational use of ethanol, attempted suicide (deliberate self-harm) is the most common reported reason for intentional poisoning. Recreational use of prescribed and over-the-counter drugs for psychotropic or euphoric effects (*abuse*) or excessive self-dosing (*misuse*) is increasingly common and may also result in unintentional self-poisoning.

About 20–25% of exposures require bedside health-professional evaluation, and 5% of all exposures require hospitalization. Poisonings account for 5–10% of all ambulance transports, emergency department visits, and intensive care unit admissions. Up to 30% of psychiatric admissions are prompted by attempted suicide via overdosage. Overall, the mortality rate is low: <1% of all exposures. It is much higher (1–2%) among hospitalized patients with intentional (suicidal) overdose, who account for the majority of serious poisonings. Acetaminophen is the pharmaceutical agent most often implicated in fatal poisoning. Overall, carbon monoxide is the leading cause of death from poisoning, but this prominence is not reflected in hospital or poison center statistics because patients with such poisoning are typically dead when discovered and are referred directly to medical examiners.

DIAGNOSIS

Although poisoning can mimic other illnesses, the correct diagnosis can usually be established by the history, physical examination, routine and toxicologic laboratory evaluations, and characteristic clinical course.

HISTORY

The *history* should include the time, route, duration, and circumstances (location, surrounding events, and intent) of exposure; the name and amount of each drug, chemical, or ingredient involved; the time of onset, nature, and severity of symptoms; the time and type of first-aid measures provided; and the medical and psychiatric history.

In many cases the patient is confused, comatose, unaware of an exposure, or unable or unwilling to admit to one. Suspicious circumstances include unexplained sudden illness in a previously healthy person or a group of healthy people; a history of psychiatric problems (particularly depression); recent changes in health, economic status,

or social relationships; and onset of illness during work with chemicals or after ingestion of food, drink (especially ethanol), or medications. When patients become ill soon after arriving from a foreign country or being arrested for criminal activity, “body packing” or “body stuffing” (ingesting or concealing illicit drugs in a body cavity) should be suspected. Relevant information may be available from family, friends, paramedics, police, pharmacists, physicians, and employers, who should be questioned regarding the patient’s habits, hobbies, behavioral changes, available medications, and antecedent events. A search of clothes, belongings, and place of discovery may reveal a suicide note or a container of drugs or chemicals. The imprint code on pills and the label on chemical products may be used to identify the ingredients and potential toxicity of a suspected poison by consulting a reference text, a computerized database, the manufacturer, or a regional poison information center (800-222-1222). Occupational exposures require review of any available material safety data sheet (MSDS) from the worksite. Because of increasing globalization, unfamiliar poisonings may result in local emergency department evaluation. Pharmaceuticals, industrial chemicals, or drugs of abuse from foreign countries may be identified with the assistance of a regional poison center or via the World Wide Web.

PHYSICAL EXAMINATION AND CLINICAL COURSE

The *physical examination* should focus initially on vital signs, the cardiopulmonary system, and neurologic status. The neurologic examination should include documentation of neuromuscular abnormalities such as dyskinesia, dystonia, fasciculations, myoclonus, rigidity, and tremors. The patient should also be examined for evidence of trauma and underlying illnesses. Focal neurologic findings are uncommon in poisoning, and their presence should prompt evaluation for a structural central nervous system (CNS) lesion. Examination of the eyes (for nystagmus and pupil size and reactivity), abdomen (for bowel activity and bladder size), and skin (for burns, bullae, color, warmth, moisture, pressure sores, and puncture marks) may reveal findings of diagnostic value. When the history is unclear, all orifices should be examined for the presence of chemical burns and drug packets. The odor of breath or vomitus and the color of nails, skin, or urine may provide important diagnostic clues.

The diagnosis of poisoning in cases of unknown etiology primarily relies on pattern recognition. The first step is to assess the pulse, blood pressure, respiratory rate, temperature, and neurologic status and to characterize the overall physiologic state as stimulated, depressed, discordant, or normal (Table 473e-1). Obtaining a complete set of vital signs and reassessing them frequently are critical. Measuring core temperature is especially important, even in difficult or combative patients, since temperature elevation is the most reliable prognosticator of poor outcome in poisoning or drug withdrawal. The next step is to consider the underlying causes of the physiologic state and to attempt to identify a pathophysiologic pattern or toxic syndrome (*toxidrome*) based on the observed findings. Assessing the severity of physiologic derangements (Table 473e-2) is useful in this regard and also for monitoring the clinical course and response to treatment. The final step is to attempt to identify the particular agent involved by looking for unique or relatively poison-specific physical or ancillary test abnormalities. Distinguishing among toxidromes on the basis of the physiologic state is summarized next.

The Stimulated Physiologic State Increased pulse, blood pressure, respiratory rate, temperature, and neuromuscular activity characterize the *stimulated* physiologic state, which can reflect sympathetic, antimuscarinic (anticholinergic), or hallucinogen poisoning or drug withdrawal (Table 473e-1). Other features are noted in (Table 473e-2). Mydriasis, a characteristic feature of all stimulants, is most marked in antimuscarinic (anticholinergic) poisoning since pupillary reactivity relies on muscarinic control. In sympathetic poisoning (e.g., due to cocaine), pupils are also enlarged, but some reactivity to light remains. The antimuscarinic (anticholinergic) toxidrome is also distinguished by hot, dry, flushed skin; decreased bowel sounds; and urinary retention. Other stimulant syndromes increase sympathetic activity and cause diaphoresis, pallor, and increased bowel activity with varying

TABLE 473e-1 DIFFERENTIAL DIAGNOSIS OF POISONING BASED ON PHYSIOLOGIC STATE

Stimulated	Depressed	Discordant	Normal
Sympathetics	Sympatholytics	Asphyxiants	Nontoxic exposure
Sympathomimetics	α ₁ -Adrenergic antagonists	Cytochrome oxidase inhibitors	Psychogenic illness
Ergot alkaloids	α ₂ -Adrenergic agonists	Inert gases	“Toxic time-bombs”
Methylxanthines	ACE inhibitors	Irritant gases	Slow absorption
Monoamine oxidase inhibitors	Angiotensin receptor blockers	Methemoglobin inducers	Anticholinergics
Thyroid hormones	Antipsychotics	Oxidative phosphorylation inhibitors	Carbamazepine
Anticholinergics	β-Adrenergic blockers	AGMA inducers	Concretion formers
Antihistamines	Calcium channel blockers	Alcohol (ketoacidosis)	Extended-release phenytoin sodium capsules (Dilantin Kapseals)
Antiparkinsonian agents	Cardiac glycosides	Ethylene glycol	Drug packets
Antipsychotics	Cyclic antidepressants	Iron	Enteric-coated pills
Antispasmodics	Cholinergics	Methanol	Diphenoxylate-atropine (Lomotil)
Belladonna alkaloids	Acetylcholinesterase inhibitors	Salicylate	Opioids
Cyclic antidepressants	Muscarinic agonists	Toluene	Salicylates
Mushrooms and plants	Nicotinic agonists	CNS syndromes	Sustained-release pills
Hallucinogens	Opioids	Extrapyramidal reactions	Valproate
Cannabinoids (marijuana)	Analgesics	Hydrocarbon inhalation	Slow distribution
LSD and analogues	GI antispasmodics	Isoniazid	Cardiac glycosides
Mescaline and analogues	Heroin	Lithium	Lithium
Mushrooms	Sedative-hypnotics	Neuroleptic malignant syndrome	Metals
Phencyclidine and analogues	Alcohols	Serotonin syndrome	Salicylate
Withdrawal syndromes	Anticonvulsants	Strychnine	Valproate
Barbiturates	Barbiturates	Membrane-active agents	Toxic metabolite
Benzodiazepines	Benzodiazepines	Amantadine	Acetaminophen
Ethanol	GABA precursors	Antiarrhythmics	Carbon tetrachloride
Opioids	Muscle relaxants	Antihistamines	Cyanogenic glycosides
Sedative-hypnotics	Other agents	Antipsychotics	Ethylene glycol
Sympatholytics	GHB products	Carbamazepine	Methanol
		Cyclic antidepressants	Methemoglobin inducers
		Local anesthetics	Mushroom toxins
		Opioids (some)	Organophosphate insecticides
		Orphenadrine	Paraquat
		Quinoline antimalarials	Metabolism disruptors
			Antineoplastic agents
			Antiviral agents
			Colchicine
			Hypoglycemic agents
			Immunosuppressive agents
			MAO inhibitors
			Metals
			Salicylate
			Warfarin

Abbreviations: ACE, angiotensin-converting enzyme; AGMA, anion-gap metabolic acidosis; CNS, central nervous system; GABA, γ-aminobutyric acid; GHB, γ-hydroxybutyrate; GI, gastrointestinal; LSD, lysergic acid diethylamide; MAO, monoamine oxidase.

TABLE 473e-2 SEVERITY OF PHYSIOLOGIC STIMULATION AND DEPRESSION IN POISONING AND DRUG WITHDRAWAL

Physiologic Stimulation	
Grade 1	Anxious, irritable, tremulous; vital signs normal; diaphoresis, flushing or pallor, mydriasis, and hyperreflexia sometimes present
Grade 2	Agitated; may have confusion or hallucinations but can converse and follow commands; vital signs mildly to moderately increased
Grade 3	Delirious; unintelligible speech, uncontrollable motor hyperactivity; moderately to markedly increased vital signs; tachyarrhythmias possible
Grade 4	Coma, seizures, cardiovascular collapse
Physiologic Depression	
Grade 1	Awake, lethargic, or sleeping but arousable by voice or tactile stimulation; able to converse and follow commands; may be confused
Grade 2	Responds to pain but not voice; can vocalize but not converse; spontaneous motor activity present; brainstem reflexes intact
Grade 3	Unresponsive to pain; spontaneous motor activity absent; brainstem reflexes depressed; motor tone, respirations, and temperature decreased
Grade 4	Unresponsive to pain; flaccid paralysis; brainstem reflexes and respirations absent; cardiovascular vital signs decreased

degrees of nausea, vomiting, abnormal distress, and occasionally diarrhea. The absolute and relative degree of vital-sign changes and neuromuscular hyperactivity can help distinguish among stimulant toxidromes. Since sympathetics stimulate the peripheral nervous system more directly than do hallucinogens or drug withdrawal, markedly increased vital signs and organ ischemia suggest sympathetic poisoning. Findings helpful in suggesting the particular drug or class causing physiologic stimulation include reflex bradycardia from selective α -adrenergic stimulants (e.g., decongestants), hypotension from selective β -adrenergic stimulants (e.g., asthma therapeutics), limb ischemia from ergot alkaloids, rotatory nystagmus from phencyclidine and ketamine (the only physiologic stimulants that cause this finding), and delayed cardiac conduction from high doses of cocaine and some anticholinergic agents (e.g., antihistamines, cyclic antidepressants, and antipsychotics). Seizures suggest a sympathetic etiology, an anticholinergic agent with membrane-active properties (e.g., cyclic antidepressants, orphenadrine, phenothiazines), or a withdrawal syndrome. Close attention to core temperature is critical in patients with grade 4 physiologic stimulation (Table 473e-2).

The Depressed Physiologic State Decreased pulse, blood pressure, respiratory rate, temperature, and neuromuscular activity are indicative of the *depressed* physiologic state caused by “functional” sympatholytics (agents that decrease cardiac function and vascular tone as well as sympathetic activity), cholinergic (muscarinic and nicotinic) agents, opioids, and sedative-hypnotic γ -aminobutyric acid (GABA)-ergic agents (Table 473e-1 and 473e-2). Miosis is also common and is most pronounced in opioid and cholinergic poisoning. Miosis is distinguished from other depressant syndromes by muscarinic and nicotinic signs and symptoms (Table 473e-1). Pronounced cardiovascular depression in the absence of significant CNS depression suggests a direct or peripherally acting sympatholytic. In contrast, in opioid and sedative-hypnotic poisoning, vital-sign changes are secondary to depression of CNS cardiovascular and respiratory centers (or consequent hypoxemia), and significant abnormalities in these parameters do not occur until there is a marked decrease in the level of consciousness (grade 3 or 4 physiologic depression; [Table 473e-2]). Other clues that suggest the cause of physiologic depression include cardiac arrhythmias and conduction disturbances (due to antiarrhythmics, β -adrenergic antagonists, calcium channel blockers, digitalis glycosides, propoxyphene, and cyclic antidepressants), mydriasis (due to tricyclic antidepressants, some antiarrhythmics, meperidine, and diphenoxylate-atropine [Lomotil]), nystagmus (due to sedative-hypnotics), and seizures (due to cholinergic agents, propoxyphene, and cyclic antidepressants).

The Discordant Physiologic State The *discordant* physiologic state is characterized by mixed vital-sign and neuromuscular abnormalities, as observed in poisoning by asphyxiants, CNS syndromes, membrane-active agents, and anion-gap metabolic acidosis (AGMA) inducers (Table 473e-1). In these conditions, manifestations of physiologic stimulation and physiologic depression occur together or at different times during the clinical course. For example, membrane-active agents can cause simultaneous coma, seizures, hypotension, and tachyarrhythmias. Alternatively, vital signs may be normal while the patient has an altered mental status or is obviously sick or clearly symptomatic. Early, pronounced vital-sign and mental-status changes suggest asphyxiant or membrane-active agent poisoning; the lack of such abnormalities suggests an AGMA inducer; and marked neuromuscular dysfunction without significant vital-sign abnormalities suggests a CNS syndrome.

The Normal Physiologic State A *normal* physiologic status and physical examination may be due to a nontoxic exposure, psychogenic illness, or poisoning by “toxic time-bombs”: agents that are slowly absorbed, are slowly distributed to their sites of action, require metabolic activation, or disrupt metabolic processes (Table 473e-1). Because so many medications have now been reformulated into a once-a-day preparations for the patient’s convenience and adherence, toxic time-bombs are increasingly common. Diagnosing a nontoxic exposure requires that the identity of the exposure agent be known or that a toxic

time-bomb exposure be excluded and the time since exposure exceed the longest known or predicted interval between exposure and peak toxicity. Psychogenic illness (fear of being poisoned, mass hysteria) may also follow a nontoxic exposure and should be considered when symptoms are inconsistent with exposure history. Anxiety reactions resulting from a nontoxic exposure can cause mild physiologic stimulation (Table 473e-2) and be indistinguishable from toxicologic causes without ancillary testing or a suitable period of observation.

LABORATORY ASSESSMENT

Laboratory assessment may be helpful in the differential diagnosis. Increased AGMA is most common in advanced methanol, ethylene glycol, and salicylate intoxication but can occur with any poisoning that results in hepatic, renal, or respiratory failure; seizures; or shock. The serum lactate concentration is more commonly low (less than the anion gap) in the former and high (nearly equal to the anion gap) in the latter. An abnormally low anion gap can be due to elevated blood levels of bromide, calcium, iodine, lithium, or magnesium. An increased osmolal gap—a difference of >10 mmol/L between serum osmolality (measured by freezing-point depression) and osmolality calculated from serum sodium, glucose, and blood urea nitrogen levels—suggests the presence of a low-molecular-weight solute such as acetone; an alcohol (benzyl, ethanol, isopropanol, methanol); a glycol (diethylene, ethylene, propylene); ether (ethyl, glycol); or an “unmeasured” cation (calcium, magnesium) or sugar (glycerol, mannitol, sorbitol). Ketosis suggests acetone, isopropyl alcohol, salicylate poisoning, or alcoholic ketoacidosis. Hypoglycemia may be due to poisoning with β -adrenergic blockers, ethanol, insulin, oral hypoglycemic agents, quinine, and salicylates, whereas hyperglycemia can occur in poisoning with acetone, β -adrenergic agonists, caffeine, calcium channel blockers, iron, theophylline, or *N*-3-pyridylmethyl-*N'*-*p*-nitrophenylurea (PNU [Vacor]). Hypokalemia can be caused by barium, β -adrenergic agonists, caffeine, diuretics, theophylline, or toluene; hyperkalemia suggests poisoning with an α -adrenergic agonist, a β -adrenergic blocker, cardiac glycosides, or fluoride. Hypocalcemia may be seen in ethylene glycol, fluoride, and oxalate poisoning.

The *electrocardiogram* (ECG) can be useful for rapid diagnostic purposes. Bradycardia and atrioventricular block may occur in patients poisoned by α -adrenergic agonists, antiarrhythmic agents, beta blockers, calcium channel blockers, cholinergic agents (carbamate and organophosphate insecticides), cardiac glycosides, lithium, or tricyclic antidepressants. QRS- and QT-interval prolongation may be caused by hyperkalemia, various antidepressants, and other membrane-active drugs (Table 473e-1). Ventricular tachyarrhythmias may be seen in poisoning with cardiac glycosides, fluorides, membrane-active drugs, methylxanthines, sympathomimetics, antidepressants, and agents that cause hyperkalemia or potentiate the effects of endogenous catecholamines (e.g., chloral hydrate, aliphatic and halogenated hydrocarbons).

Radiologic studies may occasionally be useful. Pulmonary edema (adult respiratory distress syndrome (ARDS) can be caused by poisoning with carbon monoxide, cyanide, an opioid, paraquat, phencyclidine, a sedative-hypnotic, or salicylate; by inhalation of irritant gases, fumes, or vapors (acids and alkali, ammonia, aldehydes, chlorine, hydrogen sulfide, isocyanates, metal oxides, mercury, phosgene, polymers); or by prolonged anoxia, hyperthermia, or shock. Aspiration pneumonia is common in patients with coma, seizures, and petroleum distillate aspiration. The presence of radiopaque densities on abdominal x-rays suggests the ingestion of calcium salts, chloral hydrate, chlorinated hydrocarbons, heavy metals, illicit drug packets, iodinated compounds, potassium salts, enteric-coated tablets, or salicylates.

Toxicologic analysis of urine and blood (and occasionally of gastric contents and chemical samples) can sometimes confirm or rule out suspected poisoning. Interpretation of laboratory data requires knowledge of the qualitative and quantitative tests used for screening and confirmation (enzyme-multiplied, fluorescence polarization, and radio-immunoassays; colorimetric and fluorometric assays; thin-layer, gas-liquid, or high-performance liquid chromatography; gas chromatography; mass spectrometry), their sensitivity (limit of detection) and specificity, the preferred biologic specimen for analysis, and the

473e-4 optimal time of specimen sampling. Personal communication with the hospital laboratory is essential to an understanding of institutional testing capabilities and limitations.

Rapid *qualitative* hospital-based urine tests for drugs of abuse are only screening tests that cannot confirm the exact identity of the detected substance and should not be considered diagnostic or used for forensic purposes: False-positive and false-negative results are common. A positive screen may result from other pharmaceuticals that interfere with laboratory analysis (e.g., fluoroquinolones commonly cause “false-positive” opiate screens). Confirmatory testing with gas chromatography/mass spectrometry can be requested, but it often takes weeks to obtain a reported result. A negative screening result may mean that the responsible substance is not detectable by the test used or that its concentration is too low for detection at the time of sampling. For instance, recent new drugs of abuse that often result in emergency department evaluation for unexpected complications, such as synthetic cannabinoids (spice), cathinones (bath salts), and opiate substitutes (kratom), are not detectable by hospital-based tests. In cases where a drug concentration is too low to be detected early during clinical evaluation, repeating the test at a later time may yield a positive result. Patients symptomatic from drugs of abuse often require immediate management based on the history, physical examination, and observed toxidrome without laboratory confirmation (e.g., apnea from opioid intoxication). When the patient is asymptomatic or when the clinical picture is consistent with the reported history, qualitative screening is neither clinically useful nor cost-effective. Thus, qualitative drug screens are of greatest value for the evaluation of patients with severe or unexplained toxicities, such as coma, seizures, cardiovascular instability, metabolic or respiratory acidosis, and nonsinus cardiac rhythms. In contrast to qualitative drug screens, *quantitative* serum tests are useful for evaluation of patients poisoned with acetaminophen (**Chap. 361**), alcohols (including ethylene glycol and methanol), anticonvulsants, barbiturates, digoxin, heavy metals, iron, lithium, salicylate, and theophylline as well as for the presence of carboxyhemoglobin and methemoglobin. The serum concentration in these cases guides clinical management, and results are often available within an hour.

The *response to antidotes* is sometimes useful for diagnostic purposes. Resolution of altered mental status and abnormal vital signs within minutes of IV administration of dextrose, naloxone, or flumazenil is virtually diagnostic of hypoglycemia, opioid poisoning, and benzodiazepine intoxication, respectively. The prompt reversal of dystonic (extrapyramidal) signs and symptoms following an IV dose of benztropine or diphenhydramine confirms a drug etiology. Although complete reversal of both central and peripheral manifestations of anticholinergic poisoning by physostigmine is diagnostic of this condition, physostigmine may cause some arousal in patients with CNS depression of any etiology.

TREATMENT POISONING AND DRUG OVERDOSE

GENERAL PRINCIPLES

Treatment goals include support of vital signs, prevention of further poison absorption (decontamination), enhancement of poison elimination, administration of specific antidotes, and prevention of reexposure (**Table 473e-3**). Specific treatment depends on the identity of the poison, the route and amount of exposure, the time of presentation relative to the time of exposure, and the severity of poisoning. Knowledge of the offending agents’ pharmacokinetics and pharmacodynamics is essential.

During the *pretoxic phase*, prior to the onset of poisoning, decontamination is the highest priority, and treatment is based solely on the history. The maximal potential toxicity based on the greatest possible exposure should be assumed. Since decontamination is more effective when accomplished soon after exposure and when the patient is asymptomatic, the initial history and physical examination should be focused and brief. It is also advisable to establish IV access and initiate cardiac monitoring, particularly in patients with potentially serious ingestions or unclear histories.

TABLE 473e-3 FUNDAMENTALS OF POISONING MANAGEMENT

Supportive Care	
Airway protection	Treatment of seizures
Oxygenation/ventilation	Correction of temperature abnormalities
Treatment of arrhythmias	Correction of metabolic derangements
Hemodynamic support	Prevention of secondary complications
Prevention of Further Poison Absorption	
Gastrointestinal decontamination	Decontamination of other sites
Gastric lavage	Eye decontamination
Activated charcoal	Skin decontamination
Whole-bowel irrigation	Body cavity evacuation
Dilution	
Endoscopic/surgical removal	
Enhancement of Poison Elimination	
Multiple-dose activated charcoal administration	Extracorporeal removal
	Hemodialysis
Alteration of urinary pH	Hemoperfusion
Chelation	Hemofiltration
	Plasmapheresis
	Exchange transfusion
	Hyperbaric oxygenation
Administration of Antidotes	
Neutralization by antibodies	Metabolic antagonism
Neutralization by chemical binding	Physiologic antagonism
Prevention of Reexposure	
Adult education	Notification of regulatory agencies
Child-proofing	Psychiatric referral

When an accurate history is not obtainable and a poison causing delayed toxicity (i.e., a toxic time-bomb) or irreversible damage is suspected, blood and urine should be sent for appropriate toxicologic screening and quantitative analysis. During poison absorption and distribution, blood levels may be greater than those in tissue and may not correlate with toxicity. However, high blood levels of agents whose metabolites are more toxic than the parent compound (acetaminophen, ethylene glycol, or methanol) may indicate the need for additional interventions (antidotes, dialysis). Most patients who remain asymptomatic or who become asymptomatic 6 h after ingestion are unlikely to develop subsequent toxicity and can be discharged safely. Longer observation will be necessary for patients who have ingested toxic time-bombs.

During the *toxic phase*—the interval between the onset of poisoning and its peak effects—management is based primarily on clinical and laboratory findings. *Effects after an overdose usually begin sooner, peak later, and last longer than they do after a therapeutic dose.* A drug’s published pharmacokinetic profile in standard references such as the *Physician’s Desk Reference* (PDR) is usually different from its toxicokinetic profile in overdose. Resuscitation and stabilization are the first priority. Symptomatic patients should have an IV line placed and should undergo oxygen saturation determination, cardiac monitoring, and continuous observation. Baseline laboratory, ECG, and x-ray evaluation may also be appropriate. Intravenous glucose (unless the serum level is documented to be normal), naloxone, and thiamine should be considered in patients with altered mental status, particularly those with coma or seizures. Decontamination should also be considered, but it is less likely to be effective during this phase than during the *pretoxic phase*.

Measures that enhance poison elimination may shorten the duration and severity of the toxic phase. However, they are not without risk, which must be weighed against the potential benefit. Diagnostic certainty (usually via laboratory confirmation) is generally a prerequisite. Intestinal (gut) dialysis with repetitive doses of activated charcoal (see “Multiple-Dose Activated Charcoal,” later)

can enhance the elimination of selected poisons such as theophylline or carbamazepine. Urinary alkalinization may enhance the elimination of salicylates and a few other poisons. Chelation therapy can enhance the elimination of selected metals. Extracorporeal elimination methods are effective for many poisons, but their expense and risk make their use reasonable only in patients who would otherwise have an unfavorable outcome.

During the *resolution phase* of poisoning, supportive care and monitoring should continue until clinical, laboratory, and ECG abnormalities have resolved. Since chemicals are eliminated sooner from the blood than from tissues, blood levels are usually lower than tissue levels during this phase and again may not correlate with toxicity. This discrepancy applies particularly when extracorporeal elimination procedures are used. Redistribution from tissues may cause a rebound increase in the blood level after termination of these procedures. When a metabolite is responsible for toxic effects, continued treatment may be necessary in the absence of clinical toxicity or abnormal laboratory studies.

SUPPORTIVE CARE

The goal of supportive therapy is to maintain physiologic homeostasis until detoxification is accomplished and to prevent and treat secondary complications such as aspiration, bedsores, cerebral and pulmonary edema, pneumonia, rhabdomyolysis, renal failure, sepsis, thromboembolic disease, coagulopathy, and generalized organ dysfunction due to hypoxemia or shock.

Admission to an intensive care unit is indicated for the following: patients with severe poisoning (coma, respiratory depression, hypotension, cardiac conduction abnormalities, cardiac arrhythmias, hypothermia or hyperthermia, seizures); those needing close monitoring, antidotes, or enhanced elimination therapy; those showing progressive clinical deterioration; and those with significant underlying medical problems. Patients with mild to moderate toxicity can be managed on a general medical service, on an intermediate care unit, or in an emergency department observation area, depending on the anticipated duration and level of monitoring needed (intermittent clinical observation versus continuous clinical, cardiac, and respiratory monitoring). Patients who have attempted suicide require continuous observation and measures to prevent self-injury until they are no longer suicidal.

Respiratory Care Endotracheal intubation for protection against the aspiration of gastrointestinal contents is of paramount importance in patients with CNS depression or seizures as this complication can increase morbidity and mortality rates. Mechanical ventilation may be necessary for patients with respiratory depression or hypoxemia and for facilitation of therapeutic sedation or paralysis of patients in order to prevent or treat hyperthermia, acidosis, and rhabdomyolysis associated with neuromuscular hyperactivity. Since clinical assessment of respiratory function can be inaccurate, the need for oxygenation and ventilation is best determined by continuous pulse oximetry or arterial blood-gas analysis. The gag reflex is not a reliable indicator of the need for intubation. A patient with CNS depression may maintain airway patency while being stimulated but not if left alone. Drug-induced pulmonary edema is usually noncardiac rather than cardiac in origin, although profound CNS depression and cardiac conduction abnormalities suggest the latter. Measurement of pulmonary artery pressure may be necessary to establish the cause and direct appropriate therapy. Extracorporeal measures (membrane oxygenation, venoarterial perfusion, cardiopulmonary bypass) and partial liquid (perfluorocarbon) ventilation may be appropriate for severe but reversible respiratory failure.

Cardiovascular Therapy Maintenance of normal tissue perfusion is critical for complete recovery to occur once the offending agent has been eliminated. If hypotension is unresponsive to volume expansion, treatment with norepinephrine, epinephrine, or high-dose dopamine may be necessary. Intraaortic balloon pump counterpulsation and venoarterial or cardiopulmonary perfusion techniques should be considered for severe but reversible cardiac failure. For patients with

a return of spontaneous circulation after resuscitative treatment for cardiopulmonary arrest secondary to poisoning, therapeutic hypothermia should be used according to protocol. Bradyarrhythmias associated with hypotension generally should be treated as described in [Chaps. 274 and 275](#). Glucagon, calcium, and high-dose insulin with dextrose may be effective in beta blocker and calcium channel blocker poisoning. Antibody therapy may be indicated for cardiac glycoside poisoning.

Supraventricular tachycardia associated with hypertension and CNS excitation is almost always due to agents that cause generalized physiologic excitation (Table 473e-1). Most cases are mild or moderate in severity and require only observation or nonspecific sedation with a benzodiazepine. In severe cases or those associated with hemodynamic instability, chest pain, or ECG evidence of ischemia, specific therapy is indicated. When the etiology is sympathetic hyperactivity, treatment with a benzodiazepine should be prioritized. Further treatment with a combined alpha and beta blocker (labetalol), a calcium channel blocker (verapamil or diltiazem), or a combination of a beta blocker and a vasodilator (esmolol and nitroprusside) may be considered for cases refractory to high doses of benzodiazepines. Treatment with an α -adrenergic antagonist (phentolamine) alone may sometimes be appropriate. If the cause is anticholinergic poisoning, physostigmine alone can be effective. Supraventricular tachycardia without hypertension is generally secondary to vasodilation or hypovolemia and responds to fluid administration.

For ventricular tachyarrhythmias due to tricyclic antidepressants and other membrane-active agents (Table 473e-1), sodium bicarbonate is indicated, whereas class IA, IC, and III antiarrhythmic agents are contraindicated because of similar electrophysiologic effects. Although lidocaine and phenytoin are historically safe for ventricular tachyarrhythmias of any etiology, sodium bicarbonate should be considered first for any ventricular arrhythmia suspected to have a toxicologic etiology. Intravenous emulsion therapy has shown benefit for treatment of arrhythmias and hemodynamic instability from various membrane-active agents. Beta blockers can be hazardous if the arrhythmia is due to sympathetic hyperactivity. Magnesium sulfate and overdrive pacing (by isoproterenol or a pacemaker) may be useful in patients with torsades des pointes and prolonged QT intervals. Magnesium and anti-digoxin antibodies should be considered in patients with severe cardiac glycoside poisoning. Invasive (esophageal or intracardiac) ECG recording may be necessary to determine the origin (ventricular or supraventricular) of wide-complex tachycardias ([Chaps. 276 and 277](#)). If the patient is hemodynamically stable, however, it is reasonable to simply observe him or her rather than to administer another potentially proarrhythmic agent. Arrhythmias may be resistant to drug therapy until underlying acid-base, electrolyte, oxygenation, and temperature derangements are corrected.

Central Nervous System Therapies Neuromuscular hyperactivity and seizures can lead to hyperthermia, lactic acidosis, and rhabdomyolysis and should be treated aggressively. Seizures caused by excessive stimulation of catecholamine receptors (sympathomimetic or hallucinogen poisoning and drug withdrawal) or decreased activity of GABA (isoniazid poisoning) or glycine (strychnine poisoning) receptors are best treated with agents that enhance GABA activity, such as benzodiazepine or barbiturates. Since benzodiazepines and barbiturates act by slightly different mechanisms (the former increases the frequency and the latter increases the duration of chloride channel opening in response to GABA), therapy with both may be effective when neither is effective alone. Seizures caused by isoniazid, which inhibits the synthesis of GABA at several steps by interfering with the cofactor pyridoxine (vitamin B₆), may require high doses of supplemental pyridoxine. Seizures resulting from membrane destabilization (beta blocker or cyclic antidepressant poisoning) require GABA enhancers (benzodiazepines first, barbiturates second). Phenytoin is contraindicated in toxicologic seizures: Animal and human data demonstrate worse outcomes after phenytoin loading, especially

in theophylline overdose. For poisons with central dopaminergic effects (methamphetamine, phencyclidine) manifested by psychotic behavior, a dopamine receptor antagonist, such as haloperidol, may be useful. In anticholinergic and cyanide poisoning, specific antidotal therapy may be necessary. The treatment of seizures secondary to cerebral ischemia or edema or to metabolic abnormalities should include correction of the underlying cause. Neuromuscular paralysis is indicated in refractory cases. Electroencephalographic monitoring and continuing treatment of seizures are necessary to prevent permanent neurologic damage. Serotonergic receptor overstimulation in serotonin syndrome may be treated with cyproheptadine.

Other Measures Temperature extremes, metabolic abnormalities, hepatic and renal dysfunction, and secondary complications should be treated by standard therapies.

PREVENTION OF POISON ABSORPTION

Gastrointestinal Decontamination Whether or not to perform gastrointestinal decontamination and which procedure to use depends on the time since ingestion; the existing and predicted toxicity of the ingested; the availability, efficacy, and contraindications of the procedure; and the nature, severity, and risk of complications. The efficacy of all decontamination procedures decreases with time, and data are insufficient to support or exclude a beneficial effect when they are used >1 h after ingestion. The average time from ingestion to presentation for treatment is >1 h for children and >3 h for adults. Most patients will recover from poisoning uneventfully with good supportive care alone, but complications of gastrointestinal decontamination, particularly aspiration, can prolong this process. Hence, gastrointestinal decontamination should be performed selectively, not routinely, in the management of overdose patients. It is clearly unnecessary when predicted toxicity is minimal or the time of expected maximal toxicity has passed without significant effect.

Activated charcoal has comparable or greater efficacy; has fewer contraindications and complications; and is less aversive and invasive than ipecac or gastric lavage. Thus it is the preferred method of gastrointestinal decontamination in most situations. Activated charcoal suspension (in water) is given orally via a cup, straw, or small-bore nasogastric tube. The generally recommended dose is 1 g/kg body weight because of its dosing convenience, although in vitro and in vivo studies have demonstrated that charcoal adsorbs $\geq 90\%$ of most substances when given in an amount equal to 10 times the weight of the substance. Palatability may be increased by adding a sweetener (sorbitol) or a flavoring agent (cherry, chocolate, or cola syrup) to the suspension. Charcoal adsorbs ingested poisons within the gut lumen, allowing the charcoal-toxin complex to be evacuated with stool. Charged (ionized) chemicals such as mineral acids, alkalis, and highly dissociated salts of cyanide, fluoride, iron, lithium, and other inorganic compounds are not well adsorbed by charcoal. In studies with animals and human volunteers, charcoal decreases the absorption of ingestants by an average of 73% when given within 5 min of ingestant administration, 51% when given at 30 min, and 36% when given at 60 min. For this reason, charcoal given before hospital arrival increases the potential clinical benefit. Side effects of charcoal include nausea, vomiting, and diarrhea or constipation. Charcoal may also prevent the absorption of orally administered therapeutic agents. Complications include mechanical obstruction of the airway, aspiration, vomiting, and bowel obstruction and infarction caused by inspissated charcoal. Charcoal is not recommended for patients who have ingested corrosives because it obscures endoscopy.

Gastric lavage should be considered for life-threatening poisons that cannot be treated effectively with other decontamination, elimination, or antidotal therapies (e.g., colchicine). Gastric lavage is performed by sequentially administering and aspirating ~ 5 mL of fluid per kilogram of body weight through a no. 40 French orogastric tube (no. 28 French tube for children). Except in infants, for whom normal saline is recommended, tap water is acceptable. The patient should be placed in Trendelenburg and left lateral

decubitus positions to prevent aspiration (even if an endotracheal tube is in place). Lavage decreases ingestant absorption by an average of 52% if performed within 5 min of ingestion administration, 26% if performed at 30 min, and 16% if performed at 60 min. Significant amounts of ingested drug are recovered from <10% of patients. Aspiration is a common complication (occurring in up to 10% of patients), especially when lavage is performed improperly. Serious complications (esophageal and gastric perforation, tube misplacement in the trachea) occur in $\sim 1\%$ of patients. For this reason, the physician should personally insert the lavage tube and confirm its placement, and the patient must be cooperative during the procedure. Gastric lavage is contraindicated in corrosive or petroleum distillate ingestions because of the respective risks of gastroesophageal perforation and aspiration pneumonia. It is also contraindicated in patients with a compromised unprotected airway and those at risk for hemorrhage or perforation due to esophageal or gastric pathology or recent surgery. Finally, gastric lavage is absolutely contraindicated in combative patients or those who refuse, as most published complications involve patient resistance to the procedure.

Syrup of ipecac, an emetogenic agent that was once the substance most commonly used for decontamination, no longer has a role in poisoning management. Even the American Academy of Pediatrics—traditionally the strongest proponent of ipecac—issued a policy statement in 2003 recommending that ipecac should no longer be used in poisoning treatment. Chronic ipecac use (by patients with anorexia nervosa or bulimia) has been reported to cause electrolyte and fluid abnormalities, cardiac toxicity, and myopathy.

Whole-bowel irrigation is performed by administering a bowel-cleansing solution containing electrolytes and polyethylene glycol (Golytely, Colyte) orally or by gastric tube at a rate of 2 L/h (0.5 L/h in children) until rectal effluent is clear. The patient must be in a sitting position. Although data are limited, whole-bowel irrigation appears to be as effective as other decontamination procedures in volunteer studies. It is most appropriate for those who have ingested foreign bodies, packets of illicit drugs, and agents that are poorly adsorbed by charcoal (e.g., heavy metals). This procedure is contraindicated in patients with bowel obstruction, ileus, hemodynamic instability, and compromised unprotected airways.

Cathartics are salts (disodium phosphate, magnesium citrate and sulfate, sodium sulfate) or saccharides (mannitol, sorbitol) that historically have been given with activated charcoal to promote the rectal evacuation of gastrointestinal contents. However, no animal, volunteer, or clinical data have ever demonstrated any decontamination benefit from cathartics. Abdominal cramps, nausea, and occasional vomiting are side effects. Complications of repeated dosing include severe electrolyte disturbances and excessive diarrhea. Cathartics are contraindicated in patients who have ingested corrosives and in those with preexisting diarrhea. Magnesium-containing cathartics should not be used in patients with renal failure.

Dilution (i.e., drinking water, another clear liquid, or milk at a volume of 5 mL/kg of body weight) is recommended only after the ingestion of corrosives (acids, alkali). It may increase the dissolution rate (and hence absorption) of capsules, tablets, and other solid ingestants and should *not* be used in these circumstances.

Endoscopic or surgical removal of poisons may be useful in rare situations, such as ingestion of a potentially toxic foreign body that fails to transit the gastrointestinal tract, a potentially lethal amount of a heavy metal (arsenic, iron, mercury, thallium), or agents that have coalesced into gastric concretions or bezoars (heavy metals, lithium, salicylates, sustained-release preparations). Patients who become toxic from cocaine due to its leakage from ingested drug packets require immediate surgical intervention.

Decontamination of Other Sites Immediate, copious flushing with water, saline, or another available clear, drinkable liquid is the initial treatment for topical exposures (exceptions include alkali metals, calcium oxide, phosphorus). Saline is preferred for eye irrigation. A triple wash (water, soap, water) may be best for dermal decontamination.

Inhalational exposures should be treated initially with fresh air or oxygen. The removal of liquids from body cavities such as the vagina or rectum is best accomplished by irrigation. Solids (drug packets, pills) should be removed manually, preferably under direct visualization.

ENHANCEMENT OF POISON ELIMINATION

Although the elimination of most poisons can be accelerated by therapeutic interventions, the pharmacokinetic efficacy (removal of drug at a rate greater than that accomplished by intrinsic elimination) and clinical benefit (shortened duration of toxicity or improved outcome) of such interventions are often more theoretical than proven. Accordingly, the decision to use such measures should be based on the actual or predicted toxicity and the potential efficacy, cost, and risks of therapy.

Multiple-Dose Activated Charcoal Repetitive oral dosing with charcoal can enhance the elimination of previously absorbed substances by binding them within the gut as they are excreted in the bile, are secreted by gastrointestinal cells, or passively diffuse into the gut lumen (*reverse absorption* or *enterocapillary exsorption*). Doses of 0.5–1 g/kg of body weight every 2–4 h, adjusted downward to avoid regurgitation in patients with decreased gastrointestinal motility, are generally recommended. Pharmacokinetic efficacy approaches that of hemodialysis for some agents (e.g., phenobarbital, theophylline). Multiple-dose therapy should be considered only for selected agents (theophylline, phenobarbital, carbamazepine, dapsone, quinine). Complications include intestinal obstruction, pseudo-obstruction, and nonocclusive intestinal infarction in patients with decreased gut motility. Because of electrolyte and fluid shifts, sorbitol and other cathartics are absolutely contraindicated when multiple doses of activated charcoal are administered.

Urinary Alkalinization Ion trapping via alteration of urine pH may prevent the renal reabsorption of poisons that undergo excretion by glomerular filtration and active tubular secretion. Since membranes are more permeable to non-ionized molecules than to their ionized counterparts, acidic (low- pK_a) poisons are ionized and trapped in alkaline urine, whereas basic ones become ionized and trapped in acid urine. Urinary alkalinization (producing a urine pH ≥ 7.5 and a urine output of 3–6 mL/kg of body weight per hour by the addition of sodium bicarbonate to an IV solution) enhances the excretion of chlorophenoxyacetic acid herbicides, chlorpropamide, diflunisal, fluoride, methotrexate, phenobarbital, sulfonamides, and salicylates. Contraindications include congestive heart failure, renal failure, and cerebral edema. Acid-base, fluid, and electrolyte parameters should be monitored carefully. Although acid diuresis may make theoretical sense for some overdoses (amphetamines), it is *never* indicated and is potentially harmful.

Extracorporeal Removal Hemodialysis, charcoal or resin hemoperfusion, hemofiltration, plasmapheresis, and exchange transfusion are capable of removing any toxin from the bloodstream. Agents most amenable to enhanced elimination by dialysis have low molecular mass (<500 Da), high water solubility, low protein binding, small volumes of distribution (<1 L/kg of body weight), prolonged elimination (long half-life), and high dialysis clearance relative to total-body clearance. Molecular weight, water solubility, and protein binding do not limit the efficacy of the other forms of extracorporeal removal.

Dialysis should be considered in cases of severe poisoning due to carbamazepine, ethylene glycol, isopropyl alcohol, lithium, methanol, theophylline, salicylates, and valproate. Although hemoperfusion may be more effective in removing some of these poisons, it does not correct associated acid-base and electrolyte abnormalities, and most hospitals no longer have hemoperfusion cartridges readily available. Fortunately, recent advances in hemodialysis technology make it as effective as hemoperfusion for removing poisons such as caffeine, carbamazepine, and theophylline. Both techniques require central venous access and systemic anticoagulation and may result in transient hypotension. Hemoperfusion may also cause hemolysis, hypocalcemia, and thrombocytopenia.

Peritoneal dialysis and exchange transfusion are less effective but may be used when other procedures are unavailable, contraindicated, or technically difficult (e.g., in infants). Exchange transfusion may be indicated in the treatment of severe arsine- or sodium chlorate-induced hemolysis, methemoglobinemia, and sulfhemoglobinemia. Although hemofiltration can enhance elimination of aminoglycosides, vancomycin, and metal-chelate complexes, the roles of hemofiltration and plasmapheresis in the treatment of poisoning are not yet defined.

Candidates for extracorporeal removal therapies include patients with severe toxicity whose condition deteriorates despite aggressive supportive therapy; those with potentially prolonged, irreversible, or fatal toxicity; those with dangerous blood levels of toxins; those who lack the capacity for self-detoxification because of liver or renal failure; and those with a serious underlying illness or complication that will adversely affect recovery.

Other Techniques The elimination of heavy metals can be enhanced by chelation, and the removal of carbon monoxide can be accelerated by hyperbaric oxygenation.

ADMINISTRATION OF ANTIDOTES

Antidotes counteract the effects of poisons by neutralizing them (e.g., antibody-antigen reactions, chelation, chemical binding) or by antagonizing their physiologic effects (e.g., activation of opposing nervous system activity, provision of a competitive metabolic or receptor substrate). Poisons or conditions with specific antidotes include acetaminophen, anticholinergic agents, anticoagulants, benzodiazepines, beta blockers, calcium channel blockers, carbon monoxide, cardiac glycosides, cholinergic agents, cyanide, drug-induced dystonic reactions, ethylene glycol, fluoride, heavy metals, hypoglycemic agents, isoniazid, membrane-active agents, methemoglobinemia, opioids, sympathomimetics, and a variety of envenomations. Intravenous lipid emulsion has been shown to be a successful antidote for poisoning from various anesthetics and membrane-active agents (e.g., cyclic antidepressants), but the exact mechanism of benefit is still under investigation. Antidotes can significantly reduce morbidity and mortality rates but are potentially toxic if used for inappropriate reasons. Since their safe use requires correct identification of a specific poisoning or syndrome, details of antidotal therapy are discussed with the conditions for which they are indicated (Table 473e-4).

PREVENTION OF REEXPOSURE

Poisoning is a preventable illness. Unfortunately, some adults and children are poison-prone, and recurrences are common. Unintentional polypharmacy poisoning has become especially common among adults with developmental delays, among the growing population of geriatric patients who are prescribed a large number of medications, and among adolescents and young adults experimenting with pharmaceuticals for recreational euphoria. Adults with unintentional exposures should be instructed regarding the safe use of medications and chemicals (according to labeling instructions). Confused patients may need assistance with the administration of their medications. Errors in dosing by health care providers may require educational efforts. Patients should be advised to avoid circumstances that result in chemical exposure or poisoning. Appropriate agencies and health departments should be notified in cases of environmental or workplace exposure. The best approach to young children and patients with intentional overdose (deliberate self-harm or attempted suicide) is to limit their access to poisons. In households where children live or visit, alcoholic beverages, medications, household products (automotive, cleaning, fuel, pet-care, and toiletry products), inedible plants, and vitamins should be kept out of reach or in locked or child-proof cabinets. Depressed or psychotic patients should undergo psychiatric assessment, disposition, and follow-up. They should be given prescriptions for a limited supply of drugs with a limited number of refills and should be monitored for compliance and response to therapy.

TABLE 473e-4 PATHOPHYSIOLOGIC FEATURES AND TREATMENT OF SPECIFIC TOXIC SYNDROMES AND POISONINGS

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
Stimulated				
Sympathetics ^a				
Sympathomimetics	α_1 -Adrenergic agonists (decongestants): phenylephrine, phenylpropranolamine β_2 -Adrenergic agonists (bronchodilators): albuterol, terbutaline Nonspecific adrenergic agonists: amphetamines, cocaine, ephedrine	Stimulation of central and peripheral sympathetic receptors directly or indirectly (by promoting release or inhibiting reuptake of norepinephrine and sometimes dopamine)	Physiologic stimulation (Table 473e-2). Reflex bradycardia can occur with selective α_1 agonists; β agonists can cause hypotension and hypokalemia.	Phentolamine, a nonselective α_1 -adrenergic receptor antagonist, for severe hypertension due to α_1 -adrenergic agonists; propranolol, a nonselective β blocker, for hypotension and tachycardia due to β_2 agonists; <i>either</i> labetalol, a β blocker with α -blocking activity, <i>or</i> phentolamine with esmolol, metoprolol, or another cardioselective β blocker for hypertension with tachycardia due to nonselective agents (β blockers, if used alone, can exacerbate hypertension and vasospasm due to unopposed α stimulation.); benzodiazepines; propofol
Ergot alkaloids	Ergotamine, methysergide, bromocriptine, pergolide	Stimulation and inhibition of serotonergic and α -adrenergic receptors; stimulation of dopamine receptors	Physiologic stimulation (Table 473e-2); formication; vasospasm with limb (isolated or generalized), myocardial, and cerebral ischemia progressing to gangrene or infarction. Hypotension, bradycardia, and involuntary movements can also occur.	Nitroprusside or nitroglycerine for severe vasospasm; prazosin (an α_1 blocker), captopril, nifedipine, and cyproheptadine (a serotonin receptor antagonist) for mild-to-moderate limb ischemia; dopamine receptor antagonists (antipsychotics) for hallucinations and movement disorders
Methylxanthines	Caffeine, theophylline	Inhibition of adenosine synthesis and adenosine receptor antagonism; stimulation of epinephrine and norepinephrine release; inhibition of phosphodiesterase resulting in increased intracellular cyclic adenosine and guanosine monophosphate	Physiologic stimulation (Table 473e-2); pronounced gastrointestinal symptoms and β agonist effects (see above). Toxicity occurs at lower drug levels in chronic poisoning than in acute poisoning.	Propranolol, a nonselective β blocker, for tachycardia with hypotension; any β blocker for supraventricular or ventricular tachycardia without hypotension; elimination enhanced by multiple-dose charcoal, hemoperfusion, and hemodialysis. Indications for hemoperfusion or hemodialysis include unstable vital signs, seizures, and a theophylline level of 80–100 $\mu\text{g}/\text{mL}$ after an acute overdose and 40–60 $\mu\text{g}/\text{mL}$ with chronic exposure.
Monoamine oxidase inhibitors	Phenelzine, tranylcypromine, selegiline	Inhibition of monoamine oxidase resulting in impaired metabolism of endogenous catecholamines and exogenous sympathomimetic agents	Delayed or slowly progressive physiologic stimulation (Table 473e-2); terminal hypotension and bradycardia in severe cases	Short-acting agents (e.g., nitroprusside, esmolol) for severe hypertension and tachycardia; direct-acting sympathomimetics (e.g., norepinephrine, epinephrine) for hypotension and bradycardia
Anticholinergics				
Antihistamines	Diphenhydramine, doxylamine, pyrilamine	Inhibition of central and postganglionic parasympathetic muscarinic cholinergic receptors. At high doses, amantadine, diphenhydramine, orphenadrine, phenothiazines, and tricyclic antidepressants have additional nonanticholinergic activity (see below).	Physiologic stimulation (Table 473e-2); dry skin and mucous membranes, decreased bowel sounds, flushing, and urinary retention; myoclonus and picking activity. Central effects may occur without significant autonomic dysfunction.	Physostigmine, an acetylcholinesterase inhibitor (see below), for delirium, hallucinations, and neuromuscular hyperactivity. Contraindications include asthma and nonanticholinergic cardiovascular toxicity (e.g., cardiac conduction abnormalities, hypotension, and ventricular arrhythmias).
Antipsychotics	Chlorpromazine, olanzapine, quetiapine, thioridazine	Inhibition of α -adrenergic, dopaminergic, histaminergic, muscarinic, and serotonergic receptors. Some agents also inhibit sodium, potassium, and calcium channels.	Physiologic depression (Table 473e-2), miosis, anticholinergic effects (see above), extrapyramidal reactions (see below), tachycardia	Sodium bicarbonate for ventricular tachydyrhythmias associated with QRS prolongation; magnesium, isoproterenol, and overdrive pacing for torsades des pointes. Avoid class IA, IC, and III antiarrhythmics.
Belladonna alkaloids	Atropine, hyoscyamine, scopolamine	Inhibition of central and postganglionic parasympathetic muscarinic cholinergic receptors	Physiologic stimulation (Table 473e-2); dry skin and mucous membranes, decreased bowel sounds, flushing, and urinary retention; myoclonus and picking activity. Central effects may occur without significant autonomic dysfunction.	Physostigmine, an acetylcholinesterase inhibitor (see below), for delirium, hallucinations, and neuromuscular hyperactivity. Contraindications include asthma and nonanticholinergic cardiovascular toxicity (e.g., cardiac conduction abnormalities, hypotension, and ventricular arrhythmias).

(Continued)

TABLE 473e-4 PATHOPHYSIOLOGIC FEATURES AND TREATMENT OF SPECIFIC TOXIC SYNDROMES AND POISONINGS (CONTINUED)

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
Cyclic antidepressants	Amitriptyline, doxepin, imipramine	Inhibition of α -adrenergic, dopaminergic, GABA-ergic, histaminergic, muscarinic, and serotonergic receptors; inhibition of sodium channels (see membrane-active agents); inhibition of norepinephrine and serotonin reuptake	Physiologic depression (Table 473e-2), seizures, tachycardia, cardiac conduction delays (increased PR, QRS, JT, and QT intervals; terminal QRS right-axis deviation) with aberrancy and ventricular tachydysrhythmias; anticholinergic toxidrome (see above)	Hypertonic sodium bicarbonate (or hypertonic saline) for ventricular tachydysrhythmias associated with QRS prolongation. Use of phenytoin is controversial. Avoid class IA, IC, and III antiarrhythmics. IV emulsion therapy may be beneficial in some cases.
Mushrooms and plants	<i>Amanita muscaria</i> and <i>A. pantherina</i> , henbane, jimson weed, nightshade	Inhibition of central and postganglionic parasympathetic muscarinic cholinergic receptors	Physiologic stimulation (Table 473e-2); dry skin and mucous membranes, decreased bowel sounds, flushing, and urinary retention; myoclonus and picking activity. Central effects may occur without significant autonomic dysfunction.	Physostigmine, an acetylcholinesterase inhibitor (see below), for delirium, hallucinations, and neuromuscular hyperactivity. Contraindications include asthma and nonanticholinergic cardiovascular toxicity (e.g., cardiac conduction abnormalities, hypotension, and ventricular arrhythmias).
Depressed				
Sympatholytics				
α_2 -Adrenergic agonists	Clonidine, guanabenz, tetrahydrozoline and other imidazoline decongestants, tizanidine and other imidazoline muscle relaxants	Stimulation of α_2 -adrenergic receptors leading to inhibition of CNS sympathetic outflow. Activity at nonadrenergic imidazoline binding sites also contributes to CNS effects.	Physiologic depression (Table 473e-2), miosis. Transient initial hypertension may be seen.	Dopamine and norepinephrine for hypotension; atropine for symptomatic bradycardia; naloxone for CNS depression (inconsistently effective)
Antipsychotics	Chlorpromazine, clozapine, haloperidol, risperidone, thioridazine	Inhibition of α -adrenergic, dopaminergic, histaminergic, muscarinic, and serotonergic receptors. Some agents also inhibit sodium, potassium, and calcium channels.	Physiologic depression (Table 473e-2), miosis, anticholinergic effects (see above), extrapyramidal reactions (see below), tachycardia. Cardiac conduction delays (increased PR, QRS, JT, and QT intervals) with ventricular tachydysrhythmias, including torsades des pointes, can sometimes develop.	Sodium bicarbonate for ventricular tachydysrhythmias associated with QRS prolongation; magnesium, isoproterenol, and overdrive pacing for torsades des pointes. Avoid class IA, IC, and III antiarrhythmics.
β -Adrenergic blockers	Cardioselective (β_1) blockers: atenolol, esmolol, metoprolol Nonselective (β_1 and β_2) blockers: nadolol, propranolol, timolol Partial β agonists: acebutolol, pindolol α_1 Antagonists: carvedilol, labetalol Membrane-active agents: acebutolol, propranolol, sotalol	Inhibition of β -adrenergic receptors (class II antiarrhythmic effect). Some agents have activity at additional receptors or have membrane effects (see below).	Physiologic depression (Table 473e-2), atrioventricular block, hypoglycemia, hyperkalemia, seizures. Partial agonists can cause hypertension and tachycardia. Sotalol can cause increased QT interval and ventricular tachydysrhythmias. Onset may be delayed after sotalol and sustained-release formulation overdose.	Glucagon for hypotension and symptomatic bradycardia. Atropine, isoproterenol, dopamine, dobutamine, epinephrine, and norepinephrine may sometimes be effective. High-dose insulin (with glucose and potassium to maintain euglycemia and normokalemia), electrical pacing, and mechanical cardiovascular support for refractory cases
Calcium channel blockers	Diltiazem, nifedipine and other dihydropyridine derivatives, verapamil	Inhibition of slow (type L) cardiovascular calcium channels (class IV antiarrhythmic effect)	Physiologic depression (Table 473e-2), atrioventricular block, organ ischemia and infarction, hyperglycemia, seizures. Hypotension is usually due to decreased vascular resistance rather than to decreased cardiac output. Onset may be delayed for ≥ 12 h after overdose of sustained-release formulations.	Calcium and glucagon for hypotension and symptomatic bradycardia. Dopamine, epinephrine, norepinephrine, atropine, and isoproterenol are less often effective but can be used adjunctively. High-dose insulin (with glucose and potassium to maintain euglycemia and normokalemia), IV lipid emulsion therapy, electrical pacing, and mechanical cardiovascular support for refractory cases
Cardiac glycosides	Digoxin, endogenous cardioactive steroids, foxglove and other plants, toad skin secretions (<i>Bufo</i> spp.)	Inhibition of cardiac Na^+ , K^+ -ATPase membrane pump	Physiologic depression (Table 473e-2); gastrointestinal, psychiatric, and visual symptoms; atrioventricular block with or without concomitant supraventricular tachyarrhythmia; ventricular tachyarrhythmias; hyperkalemia in acute poisoning. Toxicity occurs at lower drug levels in chronic poisoning than in acute poisoning.	Digoxin-specific antibody fragments for hemodynamically compromising dysrhythmias, Mobitz II or third-degree atrioventricular block, hyperkalemia (>5.5 meq/L; in acute poisoning only). Temporizing measures include atropine, dopamine, epinephrine, and external cardiac pacing for bradydysrhythmias and magnesium, lidocaine, or phenytoin, for ventricular tachydysrhythmias. Internal cardiac pacing and cardioversion can increase ventricular irritability and should be reserved for refractory cases.

(Continued)

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
Cyclic antidepressants	Amitriptyline, doxepin, imipramine	Inhibition of α -adrenergic, dopaminergic, GABA-ergic, histaminergic, muscarinic, and serotonergic receptors; inhibition of sodium channels (see membrane-active agents); inhibition of norepinephrine and serotonin reuptake	Physiologic depression (Table 473e-2), seizures, tachycardia, cardiac conduction delays (increased PR, QRS, JT, and QT intervals; terminal QRS right-axis deviation) with aberrancy and ventricular tachydysrhythmias; anticholinergic toxidrome (see above)	Hypertonic sodium bicarbonate (or hypertonic saline) for ventricular tachydysrhythmias associated with QRS prolongation. Use of phenytoin is controversial. Avoid class IA, IC, and III antiarrhythmics. IV emulsion therapy may be beneficial in some cases.
Cholinergics				
Acetylcholinesterase inhibitors	Carbamate insecticides (aldicarb, carbaryl, propoxur) and medicinals (neostigmine, physostigmine, tacrine); nerve gases (sarin, soman, tabun, VX); organophosphate insecticides (diazinon, chlorpyrifos-ethyl, malathion)	Inhibition of acetylcholinesterase leading to increased synaptic acetylcholine at muscarinic and nicotinic cholinergic receptor sites	Physiologic depression (Table 473e-2). Muscarinic signs and symptoms: seizures, excessive secretions (lacrimation, salivation, bronchorrhea and wheezing, diaphoresis), and increased bowel and bladder activity with nausea, vomiting, diarrhea, abdominal cramps, and incontinence of feces and urine. Nicotinic signs and symptoms: hypertension, tachycardia, muscle cramps, fasciculations, weakness, and paralysis. Death is usually due to respiratory failure. Cholinesterase activity in plasma and red cells is <50% of normal in acetylcholinesterase inhibitor poisoning.	Atropine for muscarinic signs and symptoms; 2-PAM, a cholinesterase reactivator, for nicotinic signs and symptoms due to organophosphates, nerve gases, or an unknown anticholinesterase
Muscarinic agonists	Bethanechol, mushrooms (<i>Boletus</i> , <i>Clitocybe</i> , <i>Inocybe</i> spp.), pilocarpine	Stimulation of CNS and postganglionic parasympathetic cholinergic (muscarinic) receptors		
Nicotinic agonists	Lobeline, nicotine (tobacco)	Stimulation of preganglionic sympathetic and parasympathetic and striated muscle (neuromuscular junction) cholinergic (nicotine) receptors		
Sedative-hypnotics ^a				
Anticonvulsants	Carbamazepine, ethosuximide, felbamate, gabapentin, lamotrigine, levetiracetam, oxcarbazepine, phenytoin, tiagabine, topiramate, valproate, zonisamide	Potential of the inhibitory effects of GABA by binding to the neuronal GABA-A chloride channel receptor complex and increasing the frequency or duration of chloride channel opening in response to GABA stimulation. Baclofen and, to some extent, GHB act at the GABA-B receptor complex. Meprobamate, its metabolite carisoprodol, felbamate, and orphenadrine antagonize NDMA excitatory receptors. Ethosuximide, valproate, and zonisamide decrease conduction through T-type calcium channels. Valproate decreases GABA degradation, and tiagabine blocks GABA reuptake. Carbamazepine, lamotrigine, oxcarbazepine, phenytoin, topiramate, valproate, and zonisamide slow the rate of recovery of inactivated sodium channels. Some agents also have α_2 agonist, anticholinergic, and sodium channel-blocking activity (see above and below).	Physiologic depression (Table 473e-2), nystagmus. Delayed absorption can occur with carbamazepine, phenytoin, and valproate. Myoclonus, seizures, hypertension, and tachyarrhythmias can occur with baclofen, carbamazepine, and orphenadrine.	Benzodiazepines, barbiturates, or propofol for seizures.
Barbiturates	Short-acting: butabarbital, pentobarbital, secobarbital			Hemodialysis and hemoperfusion may be indicated for severe poisoning by some agents (see "Extracorporeal Removal," in text).
Benzodiazepines	Long-acting: phenobarbital, primidone Ultrashort-acting: estazolam, midazolam, temazepam, triazolam Short-acting: alprazolam, flunitrazepam, lorazepam, oxazepam Long-acting: chlordiazepoxide, clonazepam, diazepam, flurazepam Pharmacologically related agents: zaleplon, zolpidem		Tachyarrhythmias can also occur with chloral hydrate. AGMA, hypernatremia, hyperosmolality, hyperammonemia, chemical hepatitis, and hypoglycemia can be seen in valproate poisoning. Carbamazepine and oxcarbazepine may produce hyponatremia from SIADH.	See above and below for treatment of anticholinergic and sodium channel (membrane)-blocking effects.
GABA precursors	γ -Hydroxybutyrate (sodium oxybate; GHB), γ -butyrolactone (GBL), 1,4-butanediol		Some agents can cause anticholinergic and sodium channel (membrane) blocking effects (see above and below).	
Muscle relaxants	Baclofen, carisoprodol, cyclobenzaprine, etomidate, metaxalone, methocarbamol, orphenadrine, propofol, tizanidine and other imidazoline muscle relaxants			

(Continued)

TABLE 473e-4 PATHOPHYSIOLOGIC FEATURES AND TREATMENT OF SPECIFIC TOXIC SYNDROMES AND POISONINGS (CONTINUED)

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
Other agents	Chloral hydrate, ethchlorvynol, glutethimide, meprobamate, methaqualone, methyprylon			
Discordant				
Asphyxiants				
Cytochrome oxidase inhibitors	Cyanide, hydrogen sulfide	Inhibition of mitochondrial cytochrome oxidase, with consequent blockage of electron transport and oxidative metabolism. Carbon monoxide also binds to hemoglobin and myoglobin and prevents oxygen binding, transport, and tissue uptake. (Binding to hemoglobin shifts the oxygen dissociation curve to the left.)	Signs and symptoms of hypoxemia with initial physiologic stimulation and subsequent depression (Table 473e-2); lactic acidosis; normal P_{O_2} and calculated oxygen saturation but decreased oxygen saturation by co-oximetry. (That measured by pulse oximetry is falsely elevated but is less than normal and less than the calculated value.) Headache and nausea are common with carbon monoxide. Sudden collapse may occur with cyanide and hydrogen sulfide exposure. A bitter almond breath odor may be noted with cyanide ingestion, and hydrogen sulfide smells like rotten eggs.	High-dose oxygen; IV hydroxocobalamin or IV sodium nitrite and sodium thiosulfate (Lilly cyanide antidote kit) for coma, metabolic acidosis, and cardiovascular dysfunction in cyanide poisoning
Methemoglobin inducers	Aniline derivatives, dapsone, local anesthetics, nitrates, nitrites, nitrogen oxides, nitro- and nitrosohydrocarbons, phenazopyridine, primaquine-type antimalarials, sulfonamides	Oxidation of hemoglobin iron from ferrous (Fe^{2+}) to ferric (Fe^{3+}) state prevents oxygen binding, transport, and tissue uptake. (Methemoglobinemia shifts oxygen dissociation curve to the left.) Oxidation of hemoglobin protein causes hemoglobin precipitation and hemolytic anemia (manifesting as Heinz bodies and "bite cells" on peripheral-blood smear).	Signs and symptoms of hypoxemia with initial physiologic stimulation and subsequent depression (Table 473e-2), gray-brown cyanosis unresponsive to oxygen at methemoglobin fractions >15–20%, headache, lactic acidosis (at methemoglobin fractions >45%), normal P_{O_2} and calculated oxygen saturation but decreased oxygen saturation and increased methemoglobin fraction by co-oximetry (Oxygen saturation by pulse oximetry may be falsely increased or decreased but is less than normal and less than the calculated value.)	High-dose oxygen; IV methylene blue for methemoglobin fraction >30%, symptomatic hypoxemia, or ischemia (contraindicated in G6PD deficiency); exchange transfusion and hyperbaric oxygen for severe or refractory cases
AGMA inducers	Ethylene glycol	Ethylene glycol causes CNS depression and increased serum osmolality. Metabolites (primarily glycolic acid) cause AGMA, CNS depression, and renal failure. Precipitation of oxalic acid metabolite as calcium salt in tissues and urine results in hypocalcemia, tissue edema, and crystalluria.	Initial ethanol-like intoxication, nausea, vomiting, increased osmolar gap, calcium oxalate crystalluria; delayed AGMA, back pain, renal failure; coma, seizures, hypotension, ARDS in severe cases	Sodium bicarbonate to correct acidemia; thiamine, folinic acid, magnesium, and high-dose pyridoxine to facilitate metabolism; ethanol or fomepizole for AGMA, crystalluria or renal dysfunction, ethylene glycol level >3 mmol/L (20 mg/dL), and ethanol-like intoxication or increased osmolar gap if level not readily obtainable; hemodialysis for persistent AGMA, lack of clinical improvement, and renal dysfunction; hemodialysis also useful for enhancing ethylene glycol elimination and shortening duration of treatment when ethylene glycol level is >8 mmol/L (50 mg/dL)
AGMA inducers	Iron	Hydration of ferric (Fe^{3+}) ion generates H^+ . Non-transferrin-bound iron catalyzes formation of free radicals that cause mitochondrial injury, lipid peroxidation, increased capillary permeability, vasodilation, and organ toxicity.	Initial nausea, vomiting, abdominal pain, diarrhea; AGMA, cardiovascular and CNS depression, hepatitis, coagulopathy, and seizures in severe cases. Radiopaque iron tablets may be seen on abdominal x-ray.	Whole-bowel irrigation for large ingestions; endoscopy and gastrotomy if clinical toxicity and large number of tablets are still visible on x-ray; IV hydration; sodium bicarbonate for acidemia; IV deferoxamine for systemic toxicity, iron level >90 μ mol/L (500 μ g/dL)

(Continued)

TABLE 473e-4 PATHOPHYSIOLOGIC FEATURES AND TREATMENT OF SPECIFIC TOXIC SYNDROMES AND POISONINGS (CONTINUED)

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
	Methanol	Methanol causes ethanol-like CNS depression and increased serum osmolality. Formic acid metabolite causes AGMA and retinal toxicity.	Initial ethanol-like intoxication, nausea, vomiting, increased osmolar gap; delayed AGMA, visual (clouding, spots, blindness) and retinal (edema, hyperemia) abnormalities; coma, seizures, cardiovascular depression in severe cases; possible pancreatitis	Gastric aspiration for recent ingestion; sodium bicarbonate to correct acidemia; high-dose folic acid or folate to facilitate metabolism; ethanol or fomepizole for AGMA, visual symptoms, methanol level >6 mmol/L (20 mg/dL), and ethanol-like intoxication or increased osmolal gap if level not readily obtainable; hemodialysis for persistent AGMA, lack of clinical improvement, and renal dysfunction; hemodialysis also useful for enhancing methanol elimination and shortening duration of treatment when methanol level is >15 mmol/L (50 mg/dL)
	Salicylate	Increased sensitivity of CNS respiratory center to changes in and stimulates respiration. Uncoupling of oxidative phosphorylation, inhibition of Krebs cycle enzymes, and stimulation of carbohydrate and lipid metabolism generate unmeasured endogenous anions and cause AGMA.	Initial nausea, vomiting, hyperventilation, alkalemia, alkaluria; subsequent alkalemia with both respiratory alkalosis and AGMA and paradoxical aciduria; late acidemia with CNS and respiratory depression; cerebral and pulmonary edema in severe cases. Hypoglycemia, hypocalcemia, hypokalemia, and seizures can occur.	IV hydration and supplemental glucose; sodium bicarbonate to correct acidemia; urinary alkalinization for systemic toxicity; hemodialysis for coma, cerebral edema, seizures, pulmonary edema, renal failure, progressive acid-base disturbances or clinical toxicity, salicylate level >7 mmol/L (100 mg/dL) following acute overdose
CNS syndromes				
Extrapyramidal reactions	Antipsychotics (see above), some cyclic antidepressants and antihistamines	Decreased CNS dopaminergic activity with relative excess of cholinergic activity	Akathisia, dystonia, parkinsonism	Oral or parenteral anticholinergic agent such as benztropine or diphenhydramine
Isoniazid		Interference with activation and supply of pyridoxal-5-phosphate, a cofactor for glutamic acid decarboxylase, which converts glutamic acid to GABA, results in decreased levels of this inhibitory CNS neurotransmitter; complexation with and depletion of pyridoxine itself; inhibition of nicotine adenine dinucleotide-dependent lactate and hydroxybutyrate dehydrogenases, resulting in substrate accumulation	Nausea, vomiting, agitation, confusion; coma, respiratory depression, seizures, lactic and ketoacidosis in severe cases	High-dose IV pyridoxine (vitamin B ₆) for agitation, confusion, coma, and seizures; diazepam or barbiturates for seizures
Lithium		Interference with cell membrane ion transport, adenylate cyclase and Na ⁺ , K ⁺ -ATPase activity, and neurotransmitter release	Nausea, vomiting, diarrhea, ataxia, choreoathetosis, encephalopathy, hyperreflexia, myoclonus, nystagmus, nephrogenic diabetes insipidus, falsely elevated serum chloride with low anion gap, tachycardia; coma, seizures, arrhythmias, hyperthermia, and prolonged or permanent encephalopathy and movement disorders in severe cases; delayed onset after acute overdose, particularly with delayed-release formulations. Toxicity occurs at lower drug levels in chronic poisoning than in acute poisoning.	Whole-bowel irrigation for large ingestions; IV hydration; hemodialysis for coma, seizures, encephalopathy or neuromuscular dysfunction (severe, progressive, or persistent), peak lithium level >4 meq/L following acute overdose

(Continued)

TABLE 473e-4 PATHOPHYSIOLOGIC FEATURES AND TREATMENT OF SPECIFIC TOXIC SYNDROMES AND POISONINGS (CONTINUED)

Physiologic Condition, Causes	Examples	Mechanism of Action	Clinical Features	Specific Treatments
Serotonin syndrome	Amphetamines, cocaine, dextromethorphan, meperidine, MAO inhibitors, selective serotonin (5-HT) reuptake inhibitors, tricyclic antidepressants, tramadol, triptans, tryptophan	Promotion of serotonin release, inhibition of serotonin reuptake, or direct stimulation of CNS and peripheral serotonin receptors (primarily 5-HT-1a and 5-HT-2), alone or in combination	Altered mental status (agitation, confusion, mutism, coma, seizures), neuromuscular hyperactivity (hyperreflexia, myoclonus, rigidity, tremors), and autonomic dysfunction (abdominal pain, diarrhea, diaphoresis, fever, flushing, labile hypertension, mydriasis, tearing, salivation, tachycardia). Complications include hyperthermia, lactic acidosis, rhabdomyolysis, and multisystem organ failure.	Discontinue the offending agent(s); the serotonin receptor antagonist cyproheptadine may be helpful in severe cases.
Membrane-active agents	Amantadine, anti-arrhythmics (class I and III agents; some beta blockers), antipsychotics (see above), antihistamines (particularly diphenhydramine), carbamazepine, local anesthetics (including cocaine), opioids (meperidine, propoxyphene), orphenadrine, quinoline antimalarials (chloroquine, hydroxychloroquine, quinine), cyclic antidepressants (see above)	Blockade of fast sodium membrane channels prolongs phase 0 (depolarization) of the cardiac action potential, which prolongs QRS duration and promotes reentrant (monomorphic) ventricular tachycardia. Class Ia, Ic, and III antiarrhythmics also block potassium channels during phases 2 and 3 (repolarization) of the action potential, prolonging the JT interval and promoting early after-depolarizations and polymorphic (torsades des pointes) ventricular tachycardia. Similar effects on neuronal membrane channels cause CNS dysfunction. Some agents also block α -adrenergic and cholinergic receptors or have opioid effects (see above and Chap. 468e).	QRS and JT prolongation (or both) with hypotension, ventricular tachyarrhythmias, CNS depression, seizures; anticholinergic effects with amantadine, antihistamines, carbamazepine, disopyramide, antipsychotics, and cyclic antidepressants (see above); opioid effects with meperidine and propoxyphene (see Chap. 468e); cinchonism (hearing loss, tinnitus, nausea, vomiting, vertigo, ataxia, headache, flushing, diaphoresis) and blindness with quinoline antimalarials	Hypertonic sodium bicarbonate (or hypertonic saline) for cardiac conduction delays and monomorphic ventricular tachycardia; lidocaine for monomorphic ventricular tachycardia (except when due to class Ib antiarrhythmics); magnesium, isoproterenol, and overdrive pacing for polymorphic ventricular tachycardia; physostigmine for anticholinergic effects (see above); naloxone for opioid effects (see Chap. 468e); extracorporeal removal for some agents (see text).

^aSee above and [Chap. 469e](#). ^bSee above and [Chap. 468e](#).

Abbreviations: AGMA, anion-gap metabolic acidosis; ARDS, adult respiratory distress syndrome; CNS, central nervous system; GABA, γ -aminobutyric acid; GBL, γ -butyrolactone; GHB, γ -hydroxybutyrate; G6PD, glucose-6-phosphate dehydrogenase; MAO, monoamine oxidase; NDMA, *N*-methyl-*D*-aspartate; 2-PAM, pralidoxime; SIADH, syndrome of inappropriate antidiuretic hormone secretion.

SPECIFIC TOXIC SYNDROMES AND POISONINGS

Table 473e-4 summarizes the pathophysiology, clinical features, and treatment of toxidromes and poisonings that are common, produce life-threatening toxicity, or require unique therapeutic interventions. In all cases, treatment should include attention to the general principles discussed above and, in particular, supportive care. Poisonings not covered in this chapter are discussed in specialized texts.

Alcohol, cocaine, hallucinogen, and opioid poisoning and alcohol and opioid withdrawal are discussed in Chaps. 467–469e;

nicotine addiction is discussed in Chap. 470; acetaminophen poisoning is discussed in Chap. 361; the neuroleptic malignant syndrome is discussed in Chap. 449; and heavy metal poisoning is discussed in Chap. 472e.

ACKNOWLEDGMENT

The author acknowledges the contributions of Christopher H. Linden and Michael J. Burns to this chapter in previous editions of this text.

474 Disorders Caused by Venomous Snakebites and Marine Animal Exposures

Charles Lei, Natalie J. Badowski, Paul S. Auerbach,
Robert L. Norris

This chapter outlines general principles for the evaluation and management of victims of envenomation and poisoning by venomous snakes and marine animals. Because the incidence of serious bites and stings is relatively low in developed nations, there is a paucity of relevant clinical research; as a result, therapeutic decision making often is based on anecdotal information.

VENOMOUS SNAKEBITE

EPIDEMIOLOGY

The venomous snakes of the world belong to the families Viperidae (subfamily Viperinae: Old World vipers; subfamily Crotalinae: New World and Asian pit vipers), Elapidae (including cobras, coral snakes, sea snakes, kraits, and all Australian venomous snakes), Lamprophiidae (subfamily Atractaspidinae: burrowing asps), and Colubridae (a large family in which most species are nonvenomous and only a few are dangerously toxic to humans). Most snakebites occur in developing countries with temperate and tropical climates in which populations subsist on agriculture and fishing. Recent estimates indicate that somewhere between 1.2 million and 5.5 million snakebites occur worldwide each year, with 421,000–1,841,000 envenomations and 20,000–94,000 deaths. Such wide-ranging estimates reflect the challenges of collecting accurate data in the regions most affected by venomous snakes; many victims do not seek hospital treatment, and reporting and record keeping are generally poor.

SNAKE ANATOMY/IDENTIFICATION

The typical snake venom delivery apparatus consists of bilateral venom glands situated below and behind the eyes and connected by ducts to hollow anterior maxillary fangs. In viperids (vipers and pit vipers), these fangs are long and highly mobile; they are retracted against the roof of the mouth when the snake is at rest and brought to an upright position for a strike. In elapids, the fangs are smaller and are relatively fixed in an erect position. Approximately 20% of pit viper bites and higher percentages of other snakebites (up to 75% for sea snakes) are “dry” bites, meaning no venom is released. Significant envenomation probably occurs in ~50% of all venomous snakebites.

Differentiation of venomous from nonvenomous snake species can be difficult. Viperids are characterized by somewhat triangular heads (a feature shared with many harmless snakes), elliptical pupils (also seen in some nonvenomous snakes, such as boas and pythons), enlarged maxillary fangs, and, in pit vipers, heat-sensing pits (foveal organs) on each side of the head that assist with locating prey and aiming strikes. The New World rattlesnakes possess a series of interlocking keratin plates (the rattle) on the tip of the tail that emits a buzzing sound when the snake rapidly vibrates its tail; this sound serves as a warning signal to perceived threats. Identifying venomous snakes by color pattern is notoriously misleading, as many harmless snakes have color patterns that closely mimic those of venomous snakes found in the same region.

VENOMS AND CLINICAL MANIFESTATIONS

Snake venoms are highly variable and complex mixtures of enzymes, low-molecular-weight polypeptides, glycoproteins, and other constituents. Among the deleterious components are hemorrhagins that promote vascular leakage and cause both local and systemic bleeding.

2734 Proteolytic enzymes cause local tissue necrosis, affect the coagulation pathway at various steps, and impair organ function. Hyaluronidases promote the spread of venom through connective tissue. Myocardial depressant factors reduce cardiac output, and bradykinins cause vasodilation and hypotension. Neurotoxins act either pre- or postsynaptically to block transmission at the neuromuscular junction, causing muscle paralysis. Most snake venoms have multisystem effects on their victims.

After a venomous snakebite, the time to symptom onset and clinical presentation can be quite variable and depend on the species involved, the anatomic location of the bite, and the amount of venom injected. Envenomations by most viperids and some elapids with necrotizing venoms cause progressive local pain, swelling, ecchymosis (Fig. 474-1), and (over a period of hours to days) hemorrhagic or serum-filled vesicles and bullae. In serious bites, tissue loss can be significant (Figs. 474-2 and 474-3). Systemic findings are extremely variable and can include tachycardia or bradycardia, hypotension, generalized weakness, changes in taste, mouth numbness, muscle fasciculations, pulmonary edema, renal dysfunction, and spontaneous hemorrhage (from essentially any anatomic site). Envenomations by neurotoxic elapids such as kraits (*Bungarus* species), many Australian elapids (e.g., death adders [*Acanthopis* species] and tiger snakes [*Notechis* species]), some cobras (*Naja* species), and some viperids (e.g., the South American rattlesnake [*Crotalus durissus*] and some Indian Russell's vipers [*Daboia russelii*]) cause neurologic dysfunction. Early findings may consist of nausea and vomiting, headache, paresthesias or numbness, and altered mental status. Victims may develop cranial nerve abnormalities (e.g., ptosis, difficulty swallowing) followed by peripheral motor weakness. Severe envenomation may result in muscle paralysis, including the muscles of respiration, and lead to death from respiratory failure and aspiration. Sea snake envenomation results in



A



B

FIGURE 474-1 Northern Pacific rattlesnake (*Crotalus oreganus oreganus*) envenomations. **A.** Moderately severe envenomation. Note edema and early ecchymosis 2 h after a bite to the finger. **B.** Severe envenomation. Note extensive ecchymosis 5 days after a bite to the ankle.



FIGURE 474-2 Early stages of severe, full-thickness necrosis 5 days after a Russell's viper (*Daboia russelii*) bite in southwestern India.

local pain (variable), generalized myalgias, trismus, rhabdomyolysis, and progressive flaccid paralysis; these manifestations can be delayed for several hours.

TREATMENT VENOMOUS SNAKEBITE

FIELD MANAGEMENT

The most important aspect of prehospital care of a person bitten by a venomous snake is rapid transport to a medical facility equipped to provide supportive care (airway, breathing, and circulation) and antivenom therapy. Most of the first-aid measures recommended in the past are of little benefit, and some actually worsen outcome. It is reasonable to apply a splint to the bitten extremity to lessen bleeding and discomfort and, if possible, to keep the extremity at approximately heart level. In developing countries, indigenous people should be encouraged to seek immediate care at a health care



FIGURE 474-3 Severe necrosis 10 days after a pit viper bite in a young child in Colombia. (Courtesy of Jay R. Stanka; with permission.)

facility equipped with antivenom instead of consulting traditional healers and thus incurring significant delays in reaching appropriate care. Attempting to capture and transport the offending snake, alive or dead, is not advised; instead, digital photographs of the snake taken from a safe distance may assist with snake identification and treatment decisions.

Incising and/or applying suction to the bite site should be avoided, as these measures are ineffective and exacerbate local tissue damage. Similarly ineffective and potentially harmful are the application of poultices, ice, and electric shock. Techniques or devices used in an effort to limit venom spread (e.g., lymphoocclusive bandages or tourniquets) are ineffective and may result in greater local tissue damage by restricting the spread of potentially necrotizing venom. Tourniquet use can result in loss of function and amputation even in the absence of envenomation.

Elapid venoms that are primarily neurotoxic and have no significant effects on local tissue may be localized by pressure-immobilization, a technique in which the entire limb is wrapped immediately with a bandage (e.g., crepe or elastic) and then immobilized. For this technique to be effective, the wrap pressure must be precise (40–70 mmHg in upper-extremity application and 55–70 mmHg in lower-extremity application) and the victim must be carried out of the field because walking generates muscle-pumping activity that—regardless of the anatomic site of the bite—will disperse venom into the systemic circulation. Pressure-immobilization should be used only in cases in which the offending snake is reliably identified and known to be primarily neurotoxic, the rescuer is skilled in pressure-wrap application, the necessary supplies are readily available, and the victim can be fully immobilized and carried to medical care—an uncommon combination of conditions, particularly in the regions of the world where such bites are most common.

HOSPITAL MANAGEMENT

In the hospital, the victim should be closely monitored (vital signs, cardiac rhythm, oxygen saturation, urine output) while a history is quickly obtained and a rapid, thorough physical examination is performed. To objectively evaluate the progression of local envenomation, the level of swelling in the bitten extremity should be marked and limb circumference should be measured every 15 min until the swelling has stabilized. During this period of observation, the extremity should be positioned at approximately heart level. Measures applied in the field (such as bandages or wraps) should be removed once IV access has been obtained, with cognizance that the release of such ligatures may result in hypotension or dysrhythmias when stagnant acidotic blood containing venom is released into the systemic circulation. Two large-bore IV lines should be established in unaffected extremities. Because of the potential for coagulopathy, venipuncture attempts should be minimized, and noncompressible sites (e.g., a subclavian vein) should be avoided. Early hypotension is due to pooling of blood in the pulmonary and splanchnic vascular beds. Later, systemic bleeding, hemolysis, and loss of intravascular volume into the soft tissues may play important roles. Fluid resuscitation with isotonic saline (20–40 mL/kg IV) should be initiated if there is any evidence of hemodynamic instability, and a trial of 5% albumin (10–20 mL/kg IV) may be given if the response to saline infusion is inadequate. Only after aggressive volume resuscitation and antivenom administration (see below) are accomplished should vasopressors (e.g., dopamine) be added. Invasive hemodynamic monitoring (central venous and/or continuous arterial pressures) can be helpful in such cases, although obtaining central vascular access is risky if coagulopathy has developed. Victims of neurotoxic envenomation should be watched carefully for evidence of cranial nerve dysfunction (e.g., ptosis) that may precede the onset of difficulty swallowing or respiratory insufficiency that necessitates definitive airway protection by endotracheal intubation.

Blood should be drawn for typing and cross-matching and for laboratory evaluation as soon as possible. Important studies include a complete blood count to determine the degree of hemorrhage or hemolysis and to identify thrombocytopenia; studies of renal

and hepatic function; coagulation studies to diagnose consumptive coagulopathy; creatine kinase for suspected rhabdomyolysis; and testing of urine for blood or myoglobin. In developing regions, the 20-min whole-blood clotting test can be used to reliably diagnose coagulopathy. A few milliliters of fresh blood are placed in a new, clean, plain glass receptacle (e.g., a test tube) and left undisturbed for 20 min. The tube is then tipped once to 45° to determine whether a clot has formed. If it has not, coagulopathy is diagnosed. Arterial blood gas studies, electrocardiography, and chest radiography may be helpful in severe envenomations or when there is significant comorbidity. Any arterial puncture in the setting of coagulopathy requires great caution and must be performed at an anatomic site amenable to direct-pressure tamponade. After antivenom therapy (see below), laboratory values should be rechecked every 6 h until clinical stability is achieved. If initial laboratory values are normal, the complete blood count and coagulation studies should be repeated every hour until it is clear that no systemic envenomation has occurred.

The mainstay of treatment of a venomous snakebite resulting in significant envenomation is prompt administration of specific antivenom. Antivenoms are produced by injecting animals (generally horses or sheep) with venoms from medically important snakes. Once the stock animals develop antibodies to the venoms, their serum is harvested and the antibodies are isolated for antivenom preparation, which may involve varying degrees of digestion and purification of the IgG molecules. The goal of antivenom administration is to allow antibodies (or antibody fragments) to bind and deactivate circulating venom components before they can attach to target tissues and cause deleterious effects. Antivenoms may be monospecific (directed against a particular snake species) or polyspecific (covering several medically important species in the region) but rarely offer cross-protection against snake species other than those used in their production unless the species are known to have homologous venoms. Thus, antivenom selection must be specific for the offending snake; if the antivenom chosen does not contain antibodies to that snake's venom components, it will provide no benefit and may lead to unnecessary complications (see below). In the United States, assistance in finding appropriate antivenom can be obtained from regional poison control centers.

For victims of bites by viperids or cytotoxic elapids, indications for antivenom administration include significant progressive local findings (e.g., soft tissue swelling crossing a joint or involving more than half the bitten limb) and any evidence of systemic envenomation (systemic symptoms or signs, laboratory abnormalities). Caution must be used in determining the significance of isolated soft tissue swelling after the bite of an unidentified snake because the saliva of some relatively harmless species can cause mild edema at the bite site; in such bites, antivenoms are useless and potentially harmful. Antivenoms have limited efficacy in preventing tissue damage caused by necrotizing venoms, as venom components bind to local tissues very quickly, before antivenom administration can be initiated. Nevertheless, antivenom should be administered as soon as the need for it is identified to limit further tissue damage and systemic effects. Antivenom administration after bites by neurotoxic elapids is indicated at the first sign of any evidence of neurotoxicity (e.g., cranial nerve dysfunction, peripheral neuropathy). In general, antivenom is effective only in reversing active venom toxicity; it is of no benefit in reversing effects that already have been established (e.g., renal failure, established paralysis) and that will improve only with time and other supportive therapies.

Specific comments related to the management of venomous snakebites in the United States and Canada appear in [Table 474-1](#). The package insert for the selected antivenom can be consulted regarding species covered, method of administration, starting dose, and need (if any) for re-dosing. The information in antivenom package inserts, however, is not always accurate and reliable. Whenever possible, it is advisable for treating physicians to seek advice from experts in snakebite management regarding indications for and dosing of antivenom.

Pit viper bites (rattlesnakes [*Crotalus* and *Sistrurus* spp.], cottonmouth water moccasins [*Agkistrodon piscivorus*], and copperheads [*Agkistrodon contortrix*])

- Stabilize airway, breathing, and circulation.
- Institute monitoring (vital signs, cardiac rhythm, and oxygen saturation).
- Establish two large-bore IV lines.
 - If the patient is hypotensive, administer a normal saline bolus (20–40 mL/kg IV).
 - If hypotension persists, consider 5% albumin (10–20 mL/kg IV).
- Take rapid history and perform thorough physical examination.
- Identify offending snake if possible.
- Measure and record circumference of bitten extremity every 15 min until swelling has stabilized.
- Send laboratory studies (CBC, metabolic panel, PT/INR/PTT, fibrinogen level, FDP, blood type and cross-matching, urinalysis).
 - If normal, repeat CBC and coagulation studies every hour until it is clear that no systemic envenomation has occurred.
 - If abnormal, repeat 6 h after antivenom administration (see below).
- Determine severity of envenomation.
 - None: fang marks only (“dry” bite)
 - Mild: local findings only (e.g., pain, ecchymosis, nonprogressive swelling)
 - Moderate: swelling that is clearly progressing, systemic symptoms or signs, and/or laboratory abnormalities
 - Severe: neurologic dysfunction, respiratory distress, and/or cardiovascular instability/shock
- Contact regional poison control center.
- Locate and administer antivenom as indicated: Crotalidae Polyvalent Immune Fab (CroFab) (Ovine) (BTG International Inc., West Conshohocken, PA).
 - Starting dose
 - Based on severity of envenomation
 - None or mild: none
 - Moderate: 4–6 vials
 - Severe: 6 vials
 - Dilute reconstituted vials in 250 mL of normal saline.
 - No pretesting for potential hypersensitivity; no pretreatment
 - Infuse IV over 1 h (with physician in close attendance).
 - If acute reaction to antivenom:
 - Stop infusion.
 - Treat with standard doses of epinephrine (IM or IV; latter route only in setting of severe hypotension), antihistamines (IV), and glucocorticoids (IV).
 - When reaction is controlled, restart antivenom as soon as possible (may further dilute in larger volume of normal saline).
 - Monitor clinical status over 1 h.
 - Stabilized or improved: Admit to hospital.
 - Progressing or unimproved: Repeat starting dose (and continue this pattern until patient’s condition is stable or improved).
- Blood products are rarely needed; if required, they should be given only after antivenom administration.
- Update tetanus immunization as needed.
- Prophylactic antibiotics are unnecessary unless prehospital care included incision or mouth suction.
- Pain management: acetaminophen and/or narcotics as needed; avoidance of salicylates and nonsteroidal anti-inflammatory agents
- Admit to hospital. (If no evidence of envenomation, monitor for 8 h before discharge.)
 - Give additional CroFab (2 vials every 6 h for 3 additional doses, with close monitoring).
 - Monitor for evidence of rising intracompartmental pressures (see text).
 - Provide wound care (see text).
 - Start physical therapy (see text).
- At discharge, warn patient of possible recurrent coagulopathy and symptoms/signs of delayed serum sickness.

Coral snakebites (*Micrurus* spp. and *Micruroides euryxanthus*)

- Stabilize airway, breathing, and circulation.
- Institute monitoring (vital signs, cardiac rhythm, and oxygen saturation).
- Establish one large-bore IV line and initiate normal saline infusion.
- Take rapid history and perform thorough physical examination.
- Identify offending snake if possible.
- Laboratory studies are unlikely to be helpful.
- Contact regional poison control center.
- Locate and administer antivenom as indicated: Antivenin (*Micrurus fulvius*) (equine) (commonly referred to as North American Coral Snake Antivenin; Wyeth Pharmaceuticals, New York, NY).^b
 - Refer to antivenom package insert.
 - Dilute 3–5 reconstituted vials of antivenom in 250 mL of normal saline.
 - Infuse IV over 1 h (with physician in close attendance).
 - If signs of envenomation progress despite initial antivenom, repeat the starting dose (up to 10 vials total may be required).

(Continued)

TABLE 474-1 MANAGEMENT OF VENOMOUS SNAKEBITE IN THE UNITED STATES AND CANADA^a (CONTINUED)

- If acute adverse reaction to antivenom:
 - Stop infusion.
 - Treat with standard doses of epinephrine (IM or IV; latter route only in setting of severe hypotension), antihistamines (IV), and glucocorticoids (IV).
 - When reaction is controlled, restart antivenom as soon as possible (may further dilute in larger volume of normal saline).
- If any evidence of neurologic dysfunction (e.g., any cranial nerve abnormalities such as ptosis):
 - Trial of acetylcholinesterase inhibitors (see Table 474-2)
 - With any evidence of difficulty swallowing or breathing, proceed with endotracheal intubation and ventilatory support (may be required for days or weeks).
- Update tetanus immunization as needed.
- Prophylactic antibiotics are unnecessary unless prehospital care included incision or mouth suction.
- Admit to hospital (intensive care unit) even if there is no evidence of envenomation; monitor for at least 24 h.

^aThese recommendations are specific to the care of victims of venomous snakebites in the United States and Canada and should not be applied to bites in other regions of the world. ^bAt the time of publication, a single lot of antivenom remains, with an extended expiration date of April 30, 2015.

Abbreviations: CBC, complete blood count; FDP, fibrin degradation products; PT/INR/PTT, prothrombin time/international normalized ratio/partial thromboplastin time.

Antivenom should be administered only by the IV route, and the infusion should be started slowly, with the physician at the bedside ready to intervene immediately at the first signs of an acute adverse reaction. In the absence of an adverse reaction, the rate of infusion can be increased gradually until the full starting dose has been administered (over a total period of ~1 h). Further antivenom may be necessary if the patient's acute clinical condition worsens or fails to stabilize or if venom effects that were initially controlled recur. The decision to administer further antivenom to a stabilized patient should be based on clinical evidence of persistent circulation of unbound venom components. For viperid bites, antivenom administration generally should be continued until the victim shows definite improvement (e.g., stabilized vital signs, reduced pain, restored coagulation). Neurotoxicity from elapid bites may be more difficult to reverse with antivenom. Once neurotoxicity is established and endotracheal intubation is required, further doses of antivenom are unlikely to be beneficial. In such cases, the victim must be maintained on mechanical ventilation until recovery, which may take days or weeks.

Adverse reactions to antivenom administration include immediate (nonallergic and, less commonly, allergic anaphylaxis) and delayed-type hypersensitivity reactions (serum sickness). Clinical manifestations of immediate hypersensitivity include urticaria, laryngeal edema, bronchospasm, and hypotension. Skin testing for potential hypersensitivity, although recommended by some antivenom manufacturers, is insensitive and nonspecific and should be omitted. Worldwide, the quality of antivenoms is highly variable. Rates of acute nonallergic anaphylactic reactions to some of these products exceed 50%. For this reason, some authorities have recommended pretreatment with IV antihistamines (e.g., diphenhydramine, 1 mg/kg to a maximum of 100 mg; and cimetidine, 5–10 mg/kg to a maximum of 300 mg) or even a prophylactic SC or IM dose of epinephrine (0.01 mg/kg, up to 0.3 mg). Further research is necessary, however, to determine whether any pretreatment measures are truly beneficial. Modest expansion of the patient's intravascular volume with crystalloids may blunt acute adverse blood pressure decline. Epinephrine and airway equipment should always be immediately available during antivenom infusion. An acute anaphylactic reaction may be heralded by a single hive or mild itching or may present as bronchospasm or acute cardiovascular collapse. If the patient develops an acute reaction to antivenom, the infusion should be temporarily stopped and the reaction immediately treated with IM epinephrine and IV antihistamines and glucocorticoids. Once the reaction has been controlled, if the severity of the envenomation warrants additional antivenom, the dose should be diluted further in isotonic saline and restarted as soon as possible. Rarely, in cases of recalcitrant hypotension, a concomitant IV infusion of epinephrine may be initiated and titrated to clinical effect while antivenom is administered. The patient must be monitored very closely during such therapy, preferably in an intensive care setting. Serum sickness typically develops 1–2 weeks after antivenom

administration and may present as fever, chills, urticaria, myalgias, arthralgias, lymphadenopathy, and renal or neurologic dysfunction. Treatment of serum sickness consists of systemic glucocorticoids (e.g., oral prednisone, 1–2 mg/kg daily) until all symptoms have resolved, followed by a taper over 1–2 weeks. Oral antihistamines and analgesics may provide additional relief of symptoms.

Blood products are rarely necessary in the management of an envenomed patient. The venoms of many snake species can deplete coagulation factors and cause a decrease in platelet count or hematocrit. Nevertheless, these components usually rebound within hours after administration of adequate antivenom. If the need for blood products is thought to be great (e.g., a dangerously low platelet count in a hemorrhaging patient), these products should be given only after adequate antivenom administration to avoid fueling ongoing consumptive coagulopathy.

Rhabdomyolysis and hemolysis should be managed in standard fashion. Victims who develop acute renal failure should be evaluated by a nephrologist and referred for hemodialysis or peritoneal dialysis as needed. Such renal failure, which usually is due to acute tubular necrosis, is frequently reversible. If bilateral cortical necrosis occurs, however, the prognosis for renal recovery is less favorable, and long-term dialysis with possible renal transplantation may be necessary.

Acetylcholinesterase inhibitors (e.g., edrophonium and neostigmine) may promote neurologic improvement in patients bitten by snakes with postsynaptic neurotoxins. Snakebite victims with objective evidence of neurologic dysfunction should receive a test dose of acetylcholinesterase inhibitors, as outlined in [Table 474-2](#). If they exhibit improvement, additional doses of long-acting neostigmine can be administered as needed. Close monitoring is required to prevent aspiration if repetitive dosing of neostigmine is used in an

TABLE 474-2 USE OF ACETYLCHOLINESTERASE INHIBITORS IN NEUROTOXIC SNAKE ENVENOMATION

1. Patients with clear, objective evidence of neurotoxicity (e.g., ptosis or inability to maintain upward gaze) should receive a test dose of edrophonium (if available) or neostigmine.
 - a. Pretreat with atropine: 0.6 mg IV (children, 0.02 mg/kg with a minimum of 0.1 mg)
 - b. Treat with:
 - Edrophonium: 10 mg IV (children, 0.25 mg/kg)
 - or
 - Neostigmine: 1.5–2.0 mg IM (children, 0.025–0.08 mg/kg)
2. If objective improvement is evident after 5 min, treat with:
 - a. Neostigmine: 0.5 mg IV or SC (children, 0.01 mg/kg) every 30 min as needed
 - b. Atropine: 0.6 mg IV continuous infusion over 8 h (children, 0.02 mg/kg over 8 h)
3. Closely monitor the airway and perform endotracheal intubation as needed.

2738 attempt to obviate endotracheal intubation. Acetylcholinesterase inhibitors are not a substitute for administration of an appropriate antivenom when available.

Care of the bite wound includes simple cleansing with soap and water; application of a dry, sterile dressing; and splinting of the affected extremity with padding between the digits. Once antivenom therapy has been initiated, the extremity should be elevated above heart level to reduce swelling. Tetanus immunization should be updated as appropriate. Prophylactic antibiotics are generally unnecessary after bites by North American snakes, as the incidence of secondary infection is low. In some regions, secondary bacterial infection is more common and the consequences are dire; in these regions, prophylactic antibiotics (e.g., cephalosporins) are used commonly. Antibiotics may also be considered if misguided first aid efforts have included incision or mouth suction of the bite site. Pain control should be achieved with acetaminophen or narcotic analgesics. Salicylates and nonsteroidal anti-inflammatory agents should be avoided because of their effects on blood clotting.

Most snake envenomations involve SC deposition of venom. On occasion, however, venom can be injected more deeply into muscle compartments, particularly if the offending snake was large and the bite occurred on the lower leg, forearm, or hand. Intramuscular swelling of the affected extremity may be accompanied by severe pain, decreased strength, altered sensation, cyanosis, and apparent pulselessness—signs suggesting a muscle compartment syndrome. If there is clinical concern that subfascial muscle edema may be impeding tissue perfusion, intracompartmental pressures should be measured by a minimally invasive technique (e.g., wick catheter or digital readout device). If the intracompartmental pressure is high (>30–40 mmHg), the extremity should be kept elevated while antivenom is administered. A dose of IV mannitol (1 g/kg) can be given in an effort to reduce muscle edema if the patient is hemodynamically stable. If the intracompartmental pressure remains elevated after 1 h of such therapy, a surgical consultation should be obtained for possible fasciotomy. Although evidence from animal studies suggests that fasciotomy may actually worsen myonecrosis, compartmental decompression is still necessary to preserve nerve function. Fortunately, the incidence of compartment syndrome is very low after a snakebite, with fasciotomies required in <1% of cases. Nevertheless, vigilance is essential. If a fasciotomy is deemed necessary, it should be undertaken with the patient's informed consent whenever possible.

Wound care in the days after the bite should include careful aseptic debridement of clearly necrotic tissue once coagulation has been restored. Intact serum-filled vesicles or hemorrhagic blebs should be left undisturbed. If ruptured, they should be debrided with sterile technique. Any debridement of damaged muscle should be conservative because there is evidence that such muscle may recover to a significant degree after antivenom therapy.

Physical therapy should be started as soon as possible so that the victim can return to a functional state. The incidence of long-term loss of function (e.g., reduced range of motion, impaired sensory function) is unclear but is probably quite high (>30%), particularly after viperid bites.

Any patient with signs of envenomation should be observed in the hospital for at least 24 h. In North America, a patient with an apparently “dry” viperid bite should be watched for at least 8 h before discharge, as significant toxicity occasionally develops after a delay of several hours. The onset of systemic symptoms commonly is delayed for a number of hours after bites by several of the elapids (including coral snakes, *Micrurus* species), some non-North American viperids (e.g., the hump-nosed pit viper [*Hypnale hypnale*]), and sea snakes. Patients bitten by these snakes should be observed in the hospital for at least 24 h. Patients whose condition is not stable should be admitted to an intensive care setting.

At hospital discharge, victims of venomous snakebites should be warned about symptoms and signs of wound infection, antivenom-related serum sickness, and potential long-term sequelae, such as pituitary insufficiency from Russell's viper (*D. russelii*) bites. If coagulopathy


developed in the acute stages of envenomation, it can recur during the first 2–3 weeks after the bite. In such cases, victims should be warned to avoid elective surgery or activities posing a high risk of trauma during this period. Outpatient analgesic treatment, wound management, and physical therapy should be provided.

MORBIDITY AND MORTALITY

The overall mortality rates for victims of venomous snakebites are low in regions with rapid access to medical care and appropriate antivenoms. In the United States, for example, the mortality rate is <1% for victims who receive antivenom. Eastern and western diamondback rattlesnakes (*Crotalus adamanteus* and *Crotalus atrox*, respectively) are responsible for the majority of snakebite deaths in the United States. Snakes responsible for large numbers of deaths in other countries include cobras (*Naja* spp.), carpet and saw-scaled vipers (*Echis* spp.), Russell's vipers (*D. russelii*), large African vipers (*Bitis* spp.), lancehead pit vipers (*Bothrops* spp.), and tropical rattlesnakes (*C. durissus*).

The incidence of morbidity—defined as permanent functional loss in a bitten extremity—is difficult to estimate but is substantial. Morbidity may be due to muscle, nerve, or vascular injury or to scar contracture. Such morbidity can have devastating consequences for victims in the developing world when they lose the ability to work and provide for their families. In the United States, functional loss tends to be more common and severe after rattlesnake bites than after bites by copperheads (*Agkistrodon contortrix*) or water moccasins (*Agkistrodon piscivorus*).

GLOBAL CONSIDERATIONS

 In many developing countries where snakebites are common, scarce access to medical care and antivenom resources contributes to high rates of morbidity and mortality. In many countries, the available antivenoms are inappropriate and ineffective against the venoms of medically important indigenous snakes. In those regions, further research is necessary to determine the actual impact of venomous snakebites and the specific antivenom needs in terms of both quantity and spectrum of coverage. Without accurate statistics, it is difficult to persuade antivenom manufacturers to begin and sustain production of appropriate antisera in developing nations. There is evidence that antivenoms can be produced in much more cost-effective ways than those currently being used. Just as important as getting the correct antivenoms into underserved regions is the need to educate populations about snakebite prevention and to train medical care providers in proper management approaches. Local protocols written with significant input from experienced providers in the region of concern should be developed and distributed. Appropriate antivenoms must be available at the likely first point of medical contact for patients (e.g., primary health centers) in order to minimize the common practice of referring victims to more distant, higher levels of care for the initiation of antivenom therapy. Those who care for snakebite victims in these often-remote clinics must have the skills and confidence required to begin antivenom treatment (and to treat possible reactions) as soon as possible when needed.

MARINE ENVENOMATIONS

Much of the management of envenomation by marine creatures is supportive in nature. A specific marine antivenom can be used when appropriate.

INVERTEBRATES

Cnidarians The Golgi apparatus of the cnidoblast cells within cnidarians, such as hydroids, fire coral, jellyfish, Portuguese men-of-war, and sea anemones, secretes specialized living stinging organelles called *cnidae* (also referred to as *cnidocysts*, a term that encompasses nematocysts, ptychocysts, and spirocysts). Within each organelle resides a stinging mechanism (“thread tube”) and venom. In the stinging process, cnidocysts are released and discharged upon mechanosensory stimulation. The venoms from these organisms contain bioactive substances such as tetramine, 5-hydroxytryptamine, histamine,

serotonin, and high-molecular-weight toxins, all of which can, among other effects, change the permeability of cells to ions. Victims usually report immediate prickling or burning, pruritus, paresthesias, and painful throbbing with radiation. The skin becomes reddened, darkened, edematous, and blistered and may show signs of superficial necrosis. A legion of neurologic, cardiovascular, respiratory, rheumatologic, gastrointestinal, renal, and ocular symptoms has been described, especially following stings from anemones, *Physalia* species, and scyphozoans. Anaphylaxis is possible. Hundreds of deaths have been reported, many of them caused by *Chironex fleckeri*, *Stomolophus nomurai*, *Physalia physalis*, and *Chiropsalmus quadrumanus*. Irukandji syndrome, associated with the Australian jellyfish *Carukia barnesi* and other species, is a potentially fatal condition that most commonly is characterized by hypertension; severe back, chest, and abdominal pain; nausea and vomiting; headache; sweating; and, in the most serious cases, myocardial troponin leak, pulmonary edema, and ultimately hypotension. This syndrome is thought to be mediated, at least in part, by the release of endogenous catecholamines followed by cytokines and nitric oxide.

Rescuers should note that envenomations by different cnidarians (typified by jellyfish) may respond differently to similar topical therapies; thus, the recommendations in this chapter must be tailored to local species and clinical practices. During stabilization, the skin should be decontaminated immediately with a generous application of lidocaine (up to 4%), an all-purpose agent that appears to be useful for relieving pain caused by a large number of species. Vinegar (5% acetic acid), rubbing alcohol (40–70% isopropyl alcohol), baking soda (sodium bicarbonate, especially for sea nettle stings), papain (unseasoned meat tenderizer), fresh lemon or lime juice, olive oil, or sugar may be effective, depending on the species of stinging creature. Household ammonia may in and of itself cause skin irritation.

The pressure-immobilization technique is no longer recommended for venom containment in the setting of any jellyfish sting. For the sting of the venomous box jellyfish (*C. fleckeri*), vinegar should be used. Local application of heat (up to 45°C/113°F), commonly by immersion in hot water, may be as effective. A baking soda slurry (50% baking soda, 50% water) has been recommended for *Cyanea* and *Chrysaora* species. Commercial (chemical) cold packs or real ice packs applied over a thin dry cloth or a plastic membrane have been shown to be effective in alleviating mild or moderate *Physalia utriculus* (blue-bottle jellyfish) stings but may be less effective than application of heat. Perfume, aftershave lotion, and high-proof ethanol are not efficacious and may be detrimental; formalin, ether, gasoline, and other organic solvents should not be used. Shaving the skin helps remove remaining nematocysts. Freshwater irrigation and rubbing lead to further stinging by adherent nematocysts and should be avoided.

After decontamination, topical application of an anesthetic ointment (lidocaine, benzocaine), an antihistamine (diphenhydramine), or a glucocorticoid (hydrocortisone) may be helpful. Persistent severe pain after decontamination may be treated with morphine, meperidine, fentanyl, or another narcotic analgesic. Muscle spasms may respond to diazepam (2–5 mg, titrated upward as necessary) or 10% calcium gluconate (5–10 mL) given IV. An ovine-derived IgG antivenom is available from Commonwealth Serum Laboratories (see “Sources of Antivenoms and Other Assistance,” below) for stings from the box jellyfish found in Australian and Indo-Pacific waters. However, despite its reported clinical efficacy, one school of thought holds that perhaps the antivenom is unable to bind the venom rapidly enough to account for its effects. Until further notice, current recommendations for its use apply. Treatment for Irukandji syndrome may require administration of opioid analgesics and MgSO₄ as well as aggressive treatment (phentolamine, 5 mg IV) of hypertension. All victims with systemic reactions should be observed for at least 6–8 h for rebound from any therapy, and all elderly adults should be checked for cardiac arrhythmias. Patients may suffer postinflammatory hyperpigmentation and persistent cutaneous hypersensitivity in areas of skin contact.

Safe Sea, a “jellyfish-safe” sunblock (www.nidaria.com) applied to the skin before an individual enters the water, inactivates the recognition and discharge mechanisms of nematocysts, has been tested

successfully against a number of marine stingers, and may prevent or diminish the effects of coelenterate stings. Whenever possible, a dive skin or wet suit should be worn when entering ocean waters.

Sea Sponges Many sponges produce crinotoxins that are present on their surface or in their internal secretions. As a result, touching a sea sponge may result in dermatitis or “sponge diver’s disease,” a necrotic skin reaction. Irritant dermatitis may result if small spicules of silica or calcium carbonate penetrate the skin. It is impossible to distinguish between the allergic and spicule reactions, so the treatment is the same for both. Afflicted skin should be gently dried and adhesive tape used to remove embedded spicules. Vinegar should be applied immediately and then for 10–30 min three or four times a day. Rubbing alcohol may be used if vinegar is unavailable. After spicule removal and skin decontamination, glucocorticoid or antihistamine cream may be applied to the skin. Severe vesiculation should be treated with a 2-week tapering course of systemic glucocorticoids. Mild reactions subside in 3–7 days, while involvement of large areas of the skin may result in systemic symptoms of fever, dizziness, nausea, muscle cramps, and formication.

Annelid Worms Annelid worms (bristleworms) possess rows of soft, cactus-like spines capable of inflicting painful stings. Contact results in symptoms similar to those of nematocyst envenomation. Without treatment, pain usually subsides over several hours, but inflammation may persist for up to a week (Fig. 474-4). Victims should resist the urge to scratch because scratching may fracture retrievable spines. Visible bristles should be removed with forceps and adhesive tape or a commercial facial peel; alternatively, a thin layer of rubber cement can be used to entrap and then peel away the spines. Use of vinegar or rubbing alcohol or a brief application of lidocaine or unseasoned meat tenderizer (papain) may provide additional relief. Local inflammation should be treated with topical or systemic glucocorticoids.

Sea Urchins Venomous sea urchins possess either hollow, venom-filled calcified spines or triple-jawed, globiferous pedicellariae with venom glands. Venom may also be found within the integumentary sheath on the external spine surface of certain species. The venom contains toxic components, including steroid glycosides, hemolysins, proteases, serotonin, and cholinergic substances. Contact with either venom apparatus produces immediate and intensely painful stings. One or more spines entering a joint can cause synovitis that may, over time, progress to arthritis if the spine(s) remain in or near the joint. If multiple spines penetrate the skin, the patient may develop systemic symptoms, including nausea, vomiting, numbness, muscular paralysis, and respiratory distress. A delayed hypersensitivity reaction 7–10 days after resolution of primary symptoms has been described.

The affected part should be immersed immediately in hot water to tolerance (up to 45°C/113°F). Pedicellariae should be removed by shaving so that envenomation cannot continue. Accessible embedded spines should be removed but may break off and remain lodged in



FIGURE 474-4 Rash on the hand of a diver from the spines of a bristleworm. (Courtesy of Paul Auerbach, with permission.)

2740 the victim. Residual dye from the surface of a spine remaining after the spine's removal may mimic a retained spine but is otherwise of no consequence. Soft tissue radiography or MRI can confirm the presence of retained spines, which may warrant referral for attempted surgical removal if the spines are near vital structures (e.g., joints, neurovascular bundles). Retained spines may cause the formation of granulomas that are amenable to excision or to intralesional injection with triamcinolone hexacetonide (5 mg/mL). Chronic granulomatous arthritis of the proximal interphalangeal joints has been treated with synovectomy and removal of granulation tissue. Erbium-YAG laser ablation has been deployed to destroy multiple sea urchin spines embedded in the foot and identified visually at the surface level without causing thermal necrosis of the adjacent tissues. Eosinophilic pneumonia and local and diffuse neuropathies have been observed separately after penetration by multiple spines of the black sea urchin (presumed *Diadema* species). The pathophysiologies of these phenomena have not been determined.

Starfish The crown-of-thorns *Acanthaster planci* produces venom in glandular tissue underneath the epidermis, which is released via its spiny surfaces (Fig. 474-5). Skin puncture causes pain, bleeding, and local edema, usually with remission over 30–180 min. Multiple punctures may result in reactions such as local muscle paralysis; retained fragments may cause granulomatous lesions and synovitis. There has also been a case report of elevated liver enzymes after *A. planci* envenomation. Envenomed persons benefit from acute immersion therapy in hot water, local anesthesia, wound cleansing, and possible exploration to remove foreign material.

Sea Cucumbers Sea cucumbers produce holothurin (a cantharin-like liquid toxin) in their body walls. This toxin is concentrated in the tentacular organs that are projected when the animal is threatened. Underwater, holothurin induces minimal contact dermatitis in the skin but can cause significant corneal and conjunctival irritation. A severe reaction can lead to blindness. Skin should be detoxified with 5% acetic acid (vinegar), papain, or isopropyl alcohol. The eye should be anesthetized with one or two drops of 0.5% proparacaine and irrigated with 100–250 mL of normal saline, with subsequent slit-lamp examination to identify corneal defects.

Cone Snails Cone snails use a detachable dartlike tooth to inject conotoxins into prey, inducing tetanus followed by paralysis. In an unknowing handler, stings result in small, burning punctate wounds followed by local ischemia, cyanosis, and numbness. Dysphagia, syncope, dysarthria, ptosis, blurred vision, and pruritus have also been documented. Some envenomations induce paralysis leading to respiratory failure, coma, and death. There is no antivenom. Pressure-immobilization (see “Octopuses,” below), hot-water soaks, and local anesthetics have been used empirically with success. The wound should be inspected for a foreign body. Edrophonium has been recommended as therapy for paralysis if a Tensilon test is positive.

Octopuses Serious envenomations and deaths have followed bites of Australian blue-ringed octopuses (*Octopus maculosus* and *Octopus*

lunulata). Although these animals rarely exceed 20 cm in length, their salivary venom contains a potent neurotoxin (maculotoxin) that inhibits peripheral nerve transmission by blocking sodium conductance. Oral numbness and facial numbness develop within several minutes of a serious envenomation and rapidly progress to total flaccid paralysis, including failure of respiratory muscles. Immediately after envenomation, a circumferential pressure-immobilization dressing 15 cm wide should be applied over a gauze pad (~7 × 7 × 2 cm) that has been placed directly over the sting. The dressing should be applied at venous-lymphatic pressure, with the preservation of distal arterial pulses. The limb should then be splinted. Once the victim has been transported to the nearest medical facility, the bandage can be released. Because there is no antidote and passive immunotherapy (rabbit IgG antibody) has been proven effective only against tetrodotoxin in mice, treatment is supportive. Patients with respiratory failure may need to be mechanically ventilated. If respirations are assisted, the victim may remain awake although completely paralyzed. Even with serious envenomations, significant recovery often takes place within 4–10 h, although complete recovery may require 2–4 days. Sequelae are uncommon unless related to hypoxia.

VERTEBRATES

As for all penetrating injuries, first-aid care should be undertaken. In addition, consideration must be given to local wound infection by marine *Vibrio* species and freshwater *Aeromonas hydrophila* as well as other “aquatic bacteria,” particularly if spines and needles remain embedded.

Stingrays A stingray injury is both an envenomation and a traumatic wound. Thoracic and cardiac penetration, major vessel laceration, and compartment syndrome have all been observed. The venom, which contains serotonin, 5'-nucleotidase, and phosphodiesterase, causes immediate, intense pain that may last up to 48 h. The wound is very painful (with the pain peaking at 30–60 min and lasting up to 48 h), often becomes ischemic in appearance, and heals poorly, with adjacent soft tissue swelling and prolonged disability. Systemic effects include weakness, diaphoresis, nausea, vomiting, diarrhea, dysrhythmias, syncope, hypotension, muscle cramps, fasciculations, paralysis, and (in rare cases) death. Because of differences in the toxins present on the tissues covering the stingers, freshwater stingrays may cause more severe injuries than marine stingrays.

Scorpionfish The designation *scorpionfish* encompasses members of the family Scorpaenidae and includes not only scorpionfish but also lionfish and stonefish. A complex venom with neuromuscular toxicity is delivered through 12 or 13 dorsal, 2 pelvic, and 3 anal spines. In general, the sting of a stonefish is regarded as the most serious (severe to life-threatening); that of the scorpionfish is of intermediate seriousness; and that of the lionfish is the least serious. Like that of a stingray, the sting of a scorpionfish is immediately and intensely painful. Pain from a stonefish envenomation may last for days. Systemic manifestations of scorpionfish stings are similar to those of stingray envenomations but may be more pronounced, particularly in the case of a stonefish sting, which may cause severe local tissue necrosis in addition to vital organ failure. The rare deaths that follow stonefish envenomation usually occur within 6–8 h. There is a commercially available stonefish antivenom.

Other Fish Three species of marine catfish—*Plotosus lineatus* (oriental catfish), *Bagre marinus* (sail catfish), and *Galeichthys felis* (common sea catfish)—as well as several species of freshwater catfish are capable of stinging humans. Venom is delivered through a single dorsal spine and two pectoral spines. Clinically, a catfish sting is comparable to that of a stingray, although marine catfish envenomations are generally more severe than those of their freshwater counterparts. Surgeonfish (doctorfish, tang), weeverfish, ratfish, and horned venomous sharks have also envenomed humans.

Platypus The platypus is a venomous mammal. The male has a keratinous spur on each hind limb; the spur is connected to a venom gland within the upper thigh. Skin puncture causes soft tissue edema and



FIGURE 474-5 Spines on the crown-of-thorns sea star (*Acanthaster planci*). (Courtesy of Paul Auerbach, with permission.)

pain that may last for days or weeks. Care is supportive, and hot-water therapy does not appear to benefit the victim.

TREATMENT MARINE VERTEBRATE STINGS

The stings of all marine vertebrates are treated in a similar fashion. Except for stonefish and serious scorpionfish envenomations (see below), no antivenom is available. The affected part should be immersed immediately in nonscalding hot water (45°C/113°F) for 30–90 min or until there is significant relief from pain. Recurrent pain may respond to repeated hot-water treatment. Cryotherapy is contraindicated, and no data support the use of antihistamines or steroids. Opiates will help alleviate the pain, as will local wound infiltration or regional nerve block with 1% lidocaine, 0.5% bupivacaine, and sodium bicarbonate mixed in a 5:5:1 ratio. After soaking and anesthetic administration, the wound must be explored and debrided. Radiography (in particular, MRI) may be helpful in identification of foreign bodies. After exploration and debridement, the wound should be irrigated vigorously with warm sterile water, saline, or 1% povidone-iodine in solution. Bleeding usually can be controlled by sustained local pressure for 10–15 min. In general, wounds should be left open to heal by secondary intention or treated by delayed primary closure. Tetanus immunization should be updated. Antibiotic treatment should be considered for serious wounds and for envenomation in immunocompromised hosts. The initial antibiotics should cover *Staphylococcus* and *Streptococcus* species. If the victim is immunocompromised, if a wound is primarily repaired and is more than minor, or if an infection develops, antibiotic coverage should be broadened to include *Vibrio* species. Infection with *Aeromonas* species is of similar concern for wounds associated with natural freshwater.

APPROACH TO THE PATIENT: Marine Envenomations

It is useful to be familiar with the local marine fauna and to recognize patterns of injury.

A large puncture wound or jagged laceration (particularly on the lower extremity) that is more painful than one would expect from the size and configuration of the wound is likely to be a stingray envenomation. Smaller punctures, as described above, represent the activity of a sea urchin (Fig. 474-6) or starfish. Stony corals cause rough abrasions and, in rare instances, lacerations or puncture wounds.

Coelenterate (marine invertebrate) stings sometimes create diagnostic skin patterns. A diffuse urticarial rash on exposed skin is often indicative of exposure to fragmented hydroids or larval anemones. A linear, whiplike print pattern appears where a jellyfish tentacle has contacted the skin. In the case of the dreaded box jellyfish, a cross-hatched appearance, followed by development of dark purple coloration within a few hours of the sting, heralds skin necrosis. A frosted appearance may be created by aluminum salt-based remedies applied to the wound. An encounter with fire coral causes immediate pain and swollen red skin irritation in the pattern of contact, similar to but more severe than the imprint left by exposure to an intact feather hydroid. Seabather's eruption, caused by thimble jellyfishes and larval anemones, may produce a diffuse rash that consists of clusters of erythematous macules or raised papules and is accompanied by intense itching (Fig. 474-7). Toxic sponges create a burning and painful red rash on exposed skin, which may blister and later desquamate. Because virtually all marine stingers invoke the sequelae of inflammation, local erythema, swelling, and adenopathy are fairly nonspecific.

SOURCES OF ANTIVENOMS AND OTHER ASSISTANCE

The best way to locate a specific antivenom in the United States is to call a regional poison control center and ask for assistance.

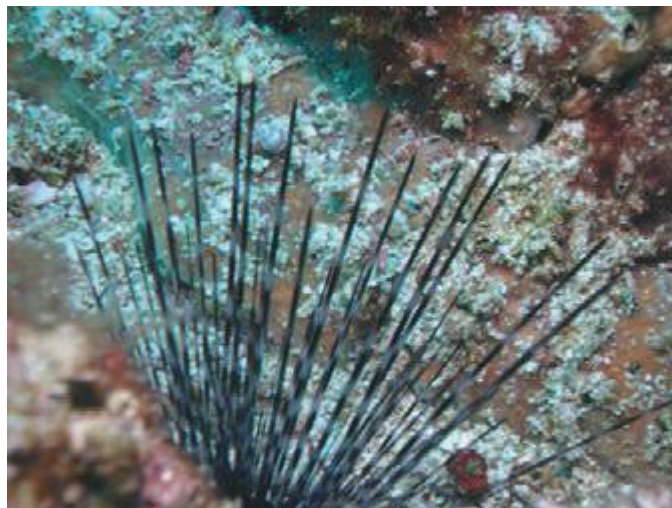


FIGURE 474-6 Spiny sea urchins. (Courtesy of Dr. Paul Auerbach, with permission.)

Divers Alert Network, a nonprofit organization designed to assist in the care of injured divers, also may help with the treatment of marine injuries. The network can be reached on the Internet at www.diversalertnetwork.org or by telephone 24 h a day at (919) 684-9111. An antivenom for the box jellyfish (*C. fleckeri*) and another for stonefish



FIGURE 474-7 Erythematous, papular rash typical of seabather's eruption caused by thimble jellyfish and larval anemones.

2742 (and severe scorpionfish) envenomation are made in Australia by the Commonwealth Serum Laboratories (CSL; 45 Poplar Road, Parkville, Victoria, Australia 3052; www.csl.com.au; 61-3-9389-1911). When administering the box jellyfish antivenom, time is of the essence. For cardiac or respiratory decompensation, give a minimum of one ampoule and up to six ampoules consecutively IV, preferably in a 1:10 dilution with normal saline. For stonefish (or severe scorpionfish) envenomation, give one ampoule of specific antivenom IM for every one or two punctures, to a maximum of three ampoules.

MARINE POISONINGS

CIGUATERA

Epidemiology and Pathogenesis Ciguatera poisoning is the most common nonbacterial food poisoning associated with fish in the United States; most U.S. cases occur in Florida and Hawaii, although, with transportation of imported fish nationwide, all clinicians need to be aware of ciguatera. The poisoning almost exclusively involves tropical and semitropical marine coral reef fish common in the Indian Ocean, the South Pacific, and the Caribbean Sea. Global estimates predict that 20,000–50,000 people may be affected by this poisoning each year. More than 400 different fish have been associated with ciguatera toxicity, but 75% of poisonings involve the reef-dwelling barracuda, snapper, jack, or grouper. Ciguatera toxin is created by warm-water ocean reef microalgae of the genus *Gambierdiscus toxicus*, whose consumption by grazing fish allows the toxin to bioaccumulate in the food chain. Three major ciguatoxins are found in the flesh and viscera of ciguateric fish: CTX-1, -2, and -3. Recent research suggests that CTX-1 activates astrocytes and astroglia. In addition, TRPV1, a nonselective cation channel expressed in nociceptive neurons, may play a role in the neurologic disturbances unique to ciguatera poisoning. Most, if not all, ciguatoxins are unaffected by freeze-drying, heat, cold, and gastric acid. None of the toxins affects the odor, color, or taste of fish. Cooking methods may alter the relative concentrations of the various toxins.

Clinical Manifestations The onset of symptoms may come within 15–30 min of ingestion and typically takes place within 2–6 h. Symptoms increase in severity over the ensuing 4–6 h. Most victims develop symptoms within 12 h of ingestion, and virtually all are afflicted within 24 h. The more than 150 signs and symptoms reported include those shown in [Table 474-3](#). Diarrhea, vomiting, and abdominal pain usually develop 3–6 h after ingestion of a ciguateric fish. Symptoms may persist for 48 h and then generally resolve (even without treatment). A pathognomonic symptom is the reversal of hot and cold tactile perception, which develops in some persons after 3–5 days and may last for months. More severe reactions tend to occur in persons previously stricken with the disease. Persons who have ingested parrotfish (scaritoxin) may develop classic ciguatera poisoning as well as a “second-phase” syndrome (after 5–10 days’ delay) of disequilibrium

with locomotor ataxia, dysmetria, and resting or kinetic tremor. This syndrome may persist for 2–6 weeks.

Diagnosis The differential diagnosis of ciguatera includes paralytic shellfish poisoning, eosinophilic meningitis, type E botulism, organophosphate insecticide poisoning, tetrodotoxin poisoning, and psychogenic hyperventilation. At present, the diagnosis of ciguatera poisoning is made on clinical grounds because no routinely used laboratory test detects ciguatoxin in human blood. Liquid chromatography–mass spectrometry is available for ciguatoxins but is of limited clinical value because most health care institutions do not have the equipment needed to perform the test. A ciguatoxin enzyme immunoassay or radioimmunoassay may be used to test small portions of the suspected fish, but even these tests may not detect the very small amount of toxin (0.1 ppb) necessary to render fish flesh toxic. A newer neuroblastoma assay may be sufficiently sensitive to detect small amounts of toxin but is not readily available for clinical use.

TREATMENT CIGUATERA POISONING

Therapy is supportive and based on symptoms. Nausea and vomiting may be controlled with an antiemetic such as ondansetron (4–8 mg IV). Syrup of ipecac and activated charcoal are not recommended for ciguatera poisoning. Hypotension may require the administration of IV crystalloid and, in rare cases, a pressor drug. Bradyarrhythmias that lead to cardiac insufficiency and hypotension generally respond well to atropine (0.5 mg IV, up to 2 mg). Goal-directed combination cardiovascular fluid and pressor therapy may be required. Cool showers or the administration of hydroxyzine (25 mg PO every 6–8 h) may relieve pruritus. Amitriptyline (25 mg PO twice a day) reportedly alleviates pruritus and dysesthesias. In three cases unresponsive to amitriptyline, tocainide has appeared to be efficacious. Nifedipine has been used to treat headache and poor circulation in order to prevent hypotension, but only after the initial acute phase of the poisoning has passed. IV infusion of mannitol may be beneficial in moderate or severe cases in fluid-repleted patients, particularly for the relief of distressing neurologic or cardiovascular symptoms, although the efficacy of this therapy has been challenged and has not been definitively proved. The infusion is rendered initially as 1 g/kg per day over 45–60 min during the acute phase (days 1–5). If symptoms improve, a second dose may be given within 3–4 h and a third dose may be administered the next day. Care must be taken to avoid dehydration in a treated patient. The mechanism of the benefit against ciguatera intoxication is perhaps hyperosmotic water-drawing action, which reverses ciguatoxin-induced Schwann cell edema. Mannitol may also act in some fashion as a “hydroxyl scavenger” or may competitively inhibit ciguatoxin at the cell membrane.

During recovery from ciguatera poisoning, the victim should exclude the following from the diet for 6 months: fish (fresh or preserved), fish sauces, shellfish, shellfish sauces, alcoholic beverages, nuts, and nut oils. Consumption of fish in ciguatera-endemic regions should be avoided. All oversized fish of any predacious reef species should be suspected of harboring ciguatoxin. Neither moray eels nor the viscera of tropical marine fish should ever be eaten.

DIARRHETIC SHELLFISH POISONING

Diarrhetic shellfish poisoning occurs with consumption of shellfish producing diarrhetic illness. The first suspected incident, which occurred in the Netherlands in 1961, was followed by outbreaks in Japan, the United Kingdom, and (most recently) China. The causative agents are the lipophilic compound okadaic acid and the dinophysistoxins, which inhibit serine and threonine protein phosphatases, with consequent protein accumulation and continued secretion of fluid in intestinal cells leading to diarrhea. Shellfish acquire these toxins by feeding on dinoflagellates, particularly of the genera *Dinophysis* and *Prorocentrum*.

Symptoms include diarrhea, nausea, vomiting, abdominal pain, and chills. Onset occurs within 30 min to 12 h. The illness is usually

TABLE 474-3 REPRESENTATIVE SIGNS AND SYMPTOMS OF CIGUATERA POISONING

System	Signs/Symptoms
Gastrointestinal	Abdominal pain, nausea, vomiting, diarrhea
Neurologic	Paresthesias, pruritus, tongue and throat numbness or burning, sensation of “carbonation” during swallowing, odontalgia or dental dysesthesias, dysphagia, tremor, fasciculations, athetosis, meningismus, aphonia, ataxia, vertigo, pain and weakness in the lower extremities, visual blurring, transient blindness, hyporeflexia, seizures, coma
Dermatologic	Conjunctivitis, maculopapular rash, skin vesiculations, dermatographism
Cardiovascular	Bradycardia, heart block, hypotension, central respiratory failure ^a
Other	Chills, dysuria, dyspnea, dyspareunia, weakness, fatigue, nasal congestion and dryness, insomnia, sialorrhea, diaphoresis, headache, arthralgias, myalgias

^aTachycardia and hypertension may occur after potentially severe transient bradycardia and hypotension. Death is rare.

self-limited; most patients recover in 3 or 4 days, and only a few require hospitalization. Treatment is supportive and focused on hydration. Toxins can be detected in food samples by a mouse bioassay, an immunoassay, and high-performance liquid chromatography with fluorometric detection (HPLC-FLD).

PARALYTIC SHELLFISH POISONING

Paralytic shellfish poisoning is induced by ingestion of any of a variety of feral or aquacultured filter-feeding organisms, including clams, oysters, scallops, mussels, chitons, limpets, starfish, and sand crabs. The origin of their toxicity is the chemical toxin they accumulate and concentrate by feeding on various planktonic dinoflagellates (e.g., *Protogonyaulax*, *Ptychodiscus*, and *Gymnodinium*) and protozoan organisms. The unicellular phytoplanktonic organisms form the foundation of the food chain, and in warm summer months these organisms “bloom” in nutrient-rich coastal temperate and semitropical waters. In the United States, paralytic shellfish poisoning is acquired primarily from seafood harvested in the Northeast, the Pacific Northwest, and Alaska. These planktonic species can release massive amounts of toxic metabolites into the water and cause mortality in bird and marine populations. The paralytic shellfish toxins are water soluble as well as heat and acid stable; they cannot be destroyed by ordinary cooking or freezing. Contaminated seafood looks, smells, and tastes normal. The best-characterized, most potent, and most frequently identified paralytic shellfish toxin is saxitoxin, which takes its name from the Alaska butter clam *Saxidomus giganteus*. Saxitoxin appears to block sodium conductance, inhibiting neuromuscular transmission at the axonal and muscle membrane levels. A toxin concentration of >75 µg/100 g of foodstuff is considered hazardous to humans. In the 1972 New England “red tide,” the concentration of saxitoxin in blue mussels exceeded 9000 µg/100 g of foodstuff.

The onset of intraoral and perioral paresthesias (notably of the lips, tongue, and gums) comes within minutes to a few hours after ingestion of contaminated shellfish, and these paresthesias progress rapidly to involve the neck and distal extremities. The tingling or burning sensation later changes to numbness. Other symptoms rapidly develop and include light-headedness, disequilibrium, incoordination, weakness, hyperreflexia, incoherence, dysarthria, sialorrhea, dysphagia, thirst, diarrhea, abdominal pain, nausea, vomiting, nystagmus, dysmetria, headache, diaphoresis, loss of vision, chest pain, and tachycardia. Flaccid paralysis and respiratory insufficiency may follow 2–12 h after ingestion. In the absence of hypoxia, the victim often remains alert but paralyzed. Up to 12% of patients die.

TREATMENT PARALYTIC SHELLFISH POISONING

Treatment is supportive and based on symptoms. If the victim comes to medical attention within the first few hours after poison ingestion, the stomach should be emptied by gastric lavage and then irrigated with 2 L (in 200-mL aliquots) of a solution of 2% sodium bicarbonate; this intervention has not been proved to be of benefit but is based on the notion that gastric acidity may enhance the potency of saxitoxin. Because breathing difficulty can be rapid in onset, induction of emesis is not advised. The administration of activated charcoal (50–100 g) and a cathartic (sorbitol, 20–50 g) makes empirical sense because these shellfish toxins are believed to bind well to charcoal. Some authors advise against administration of magnesium-based solutions (e.g., certain cathartics), cautioning that hypermagnesemia may contribute to suppression of nerve conduction.

The most serious problem is respiratory paralysis. The victim should be closely observed for respiratory distress for at least 24 h in a hospital. With prompt recognition of ventilatory failure, endotracheal intubation, and assisted ventilation, anoxic myocardial and brain injury may be prevented. If the patient survives for 18 h, the prognosis is good for a complete recovery.

A direct human serum assay to identify the toxin responsible for paralytic shellfish poisoning is not yet clinically available; the mouse bioassay in widespread use may be replaced by an automated

tissue-culture bioassay. A polyclonal enzyme-linked immunosorbent assay (ELISA) to measure specific toxins is under development, as is HPLC-FLD. In addition, an inhibition immunoassay that may be able to simultaneously detect paralytic shellfish, diarrhetic shellfish, and amnesic shellfish toxins is being investigated.

DOMOIC ACID INTOXICATION (AMNESTIC SHELLFISH POISONING)

In late 1987 in eastern Canada, an outbreak of gastrointestinal and neurologic symptoms (amnesic shellfish poisoning) was documented in persons who had consumed mussels found to be contaminated with domoic acid. In this outbreak, the source of the toxin was *Nitzschia pungens*, a diatom ingested by the mussels. Since the Canadian outbreak, the toxin has been found in shellfish from the United States, the United Kingdom, and Spain. In 1991, an epidemic of domoic acid poisoning in the state of Washington was attributed to the consumption of razor clams. A water-soluble, heat-stable neuroexcitatory amino acid with biochemical analogues of kainic acid and glutamic acid, domoic acid binds to the kainate type of glutamate receptor with three times the affinity of kainic acid and is 20 times as powerful a toxin. Shellfish can be tested for domoic acid by mouse bioassay and HPLC. The regulatory limit for domoic acid in shellfish is 20 parts per million.

The abnormalities noted within 24 h of ingesting contaminated mussels (*Mytilus edulis*) include arousal, confusion, disorientation, and memory loss. The median time of onset is 5.5 h. Other prominent signs and symptoms include severe headache, nausea, vomiting, diarrhea, abdominal cramps, hiccups, arrhythmias, hypotension, seizures, ophthalmoplegia, pupillary dilation, piloerection, hemiparesis, mutism, grimacing, agitation, emotional lability, coma, copious bronchial secretions, and pulmonary edema. Histologic study of brain tissue taken at autopsy has shown neuronal necrosis or cell loss and astrogliosis, most prominently in the hippocampus and the amygdaloid nucleus—findings similar to those in animals poisoned with kainic acid. Several months after the primary intoxication, victims still display chronic residual memory deficits and motor neuronopathy or axonopathy. Nonneurologic illness does not persist.

TREATMENT DOMOIC ACID INTOXICATION

Therapy is supportive and based on symptoms. Because kainic acid neuropathology seems to be nearly entirely seizure mediated, the emphasis should be on anticonvulsive therapy, for which diazepam appears to be as effective as any other drug.

SCOMBROID POISONING

Scombroid fish poisoning may be the most common type of seafood poisoning worldwide. It follows consumption of scombroid (mackerel-like) fish, which include albacore, bluefin, and yellowfin tuna; mackerel; saury; needlefish; wahoo; skipjack; and bonito, as well as nonscombroid fish, such as dolphinfish (Hawaiian mahimahi, *Coryphaena hippurus*), kahawai, sardine, black marlin, pilchard, anchovy, herring, amberjack, and Australian ocean salmon. In the northeastern and mid-Atlantic United States, bluefish (*Pomatomus saltatrix*) has been linked to scombroid poisoning. Because greater numbers of nonscombroid fish are being recognized as scombrototoxic, the syndrome may more appropriately be called *pseudoallergic fish poisoning*.

Under conditions of inadequate preservation or refrigeration, the musculature of these dark- or red-fleshed fish undergoes decomposition by *Proteus morganii* and *Klebsiella pneumoniae* bacteria, with consequent decarboxylation of the amino acid L-histidine to histamine, histamine phosphate, and histamine hydrochloride. Histamine levels of 20–50 mg/100 g are noted in toxic fish, with levels >400 mg/100 g on occasion. However, it is possible that some other compound may be responsible for this intoxication, because large doses of oral histamine do not reproduce the affliction. It is proposed that this unknown agent works by inhibiting the metabolism of histamine, promoting degranulation of mast cells to release endogenous histamine, or acting as a histamine receptor agonist. Whatever toxin or toxins are involved

2744 are heat stable and are not destroyed by domestic or commercial cooking. Affected fish typically have a sharply metallic or peppery taste; however, they may be normal in appearance, color, and flavor. Not all persons who eat a contaminated fish necessarily become ill, perhaps because of uneven distribution of decay within the fish.

Symptoms develop within 15–90 min of ingestion. Most cases are mild, with tingling of lips and mouth, mild abdominal discomfort, and nausea. The more severe and commonly described presentation includes flushing (sharply demarcated; exacerbated by ultraviolet exposure; particularly pronounced on the face, neck, and upper trunk), a sensation of warmth without elevated core temperature, conjunctival hyperemia, pruritus, urticaria, and angioneurotic edema. This syndrome may progress to bronchospasm, nausea, vomiting, diarrhea, epigastric pain, abdominal cramps, dysphagia, headache, thirst, pharyngitis, gingival burning, palpitations, tachycardia, dizziness, and hypotension. Without treatment, the symptoms generally resolve within 8–12 h. Because of blockade of gastrointestinal tract histaminase, the reaction may be more severe in a person who is concurrently ingesting isoniazid.

TREATMENT SCOMBROID POISONING

Therapy is directed at reversing the histamine effect with antihistamines, either H-1 or H-2. If bronchospasm is severe, an inhaled bronchodilator—or in rare, extremely severe circumstances, injected epinephrine—may be used. Glucocorticoids are of no proven benefit. Protracted nausea and vomiting, which may empty the stomach of toxin, may be controlled with a specific antiemetic, such as ondansetron or prochlorperazine. The persistent headache of scombroid poisoning may respond to cimetidine or a similar antihistamine if standard analgesics are not effective.

475 Ectoparasite Infestations and Arthropod Injuries

Richard J. Pollack, Scott A. Norton

Ectoparasites include arthropods and creatures from other phyla that infest the skin or hair of animals; the host animals provide them with sustenance and shelter. The ectoparasites may penetrate within or beneath the surface of the host or may attach by mouthparts and specialized claws. These organisms may inflict direct mechanical injury, consume blood or nutrients, induce hypersensitivity reactions, inoculate toxins, transmit pathogens, and incite fear or disgust. Humans are the sole or obligate hosts for many kinds of ectoparasites and serve as facultative or paratenic (accidental) hosts for many others.

Arthropods that are ectoparasitic or otherwise cause injury include insects (such as lice, fleas, bedbugs, wasps, ants, bees, and flies), arachnids (spiders, scorpions, mites, and ticks), millipedes, and centipedes. Certain nematodes (helminths), such as the hookworms (Chap. 256), are ectoparasitic in that they penetrate and migrate through the skin. Infrequently encountered ectoparasites in other phyla include the tentacles (tongue worms) and leeches.

Arthropods may also cause injury when they attempt to take a blood meal or as they defend themselves by biting, stinging, or exuding venoms. Various arachnids (spiders and scorpions), insects (bees, hornets, wasps, ants, flies, true bugs, caterpillars, and beetles), millipedes, and centipedes produce ill effects during these behaviors. Similarly, certain ectoparasites (e.g., ticks, biting mites, and fleas) that typically infest nonhuman animals can be medically significant. In the United States, lesions caused by arthropod bites and stings are so diverse and variable that it is rarely possible to identify the precise causative organism without a bona fide specimen and taxonomic expertise.

SCABIES

The human itch mite, *Sarcoptes scabiei* var. *hominis*, is a common cause of itching dermatosis, infesting ~300 million persons worldwide at any one time. Gravid female mites (~0.3 mm in length) burrow superficially within the stratum corneum, depositing three or fewer eggs per day. Six-legged larvae mature to eight-legged nymphs and then to adults. Gravid adult females emerge to the surface of the skin about 8 days later and then (re)invade the skin of the same or another host. Newly fertilized female mites are transferred from person to person mainly by direct skin-to-skin contact; transfer is facilitated by crowding, poor hygiene, and sex with multiple partners. Generally, these mites die within a day or so in the absence of host contact. Transmission via sharing of contaminated bedding or clothing occurs far less frequently than is often thought. In the United States, scabies may account for up to 5% of visits to dermatologists. Outbreaks occur in preschools, hospitals, nursing homes, and other residential institutions.

The itching and rash associated with scabies derive from a sensitization reaction to the mites and their secretions/excretions. A person's initial infestation remains asymptomatic for up to 6 weeks before the onset of intense pruritus, but a reinfestation produces a hypersensitivity reaction without delay. Burrows become surrounded by inflammatory infiltrates composed of eosinophils, lymphocytes, and histiocytes, and a generalized hypersensitivity rash later develops in remote sites. Immunity and associated scratching limit most infestations to <15 mites per person. Hyperinfestation with thousands of mites, a condition known as *crusted scabies* (formerly termed *Norwegian scabies*), may result from glucocorticoid use, immunodeficiency, and neurologic or psychiatric illnesses that limit the itch and/or the scratch response.

Pruritus typically intensifies at night and after hot showers. Classic burrows are often difficult to find because they are few in number and may be obscured by excoriations. Burrows appear as dark wavy lines in the upper epidermis and are 3–15 mm long. Scabetic lesions are most common on the volar wrists and along the digital web spaces. In males, the penis and scrotum become involved. Small papules and vesicles, often accompanied by eczematous plaques, pustules, or nodules, appear symmetrically at those sites; within intertriginous areas; around the navel and belt line; in the axillae; and on the buttocks and upper thighs. Except in infants, the face, scalp, neck, palms, and soles are usually spared. Crusted scabies often resembles psoriasis: both are characterized by widespread thick keratotic crusts, scaly plaques, and dystrophic nails. Characteristic burrows are not seen in crusted scabies, and patients usually do not itch, although their infestations are highly contagious and have been responsible for outbreaks of classic scabies in hospitals.

Scabies should be considered in patients with pruritus and symmetric superficial, excoriated, papulovesicular skin lesions in characteristic locations, particularly if there is a history of household contact with an infested person. Burrows should be sought and unroofed with a sterile needle or scalpel blade, and the scrapings should be examined microscopically for mites, eggs, and fecal pellets. Examination of skin biopsies (including superficial cyanoacrylate biopsy) or scrapings, dermatoscopic imaging of papulovesicular lesions, and microscopic inspection of clear cellophane tape lifted from lesions also may be diagnostic. In the absence of identifiable mites or eggs, the diagnosis is based on a history of pruritus, a clinical examination, and an epidemiologic link. Diverse kinds of dermatitis from other causes frequently are misdiagnosed as scabies, particularly in presumed “outbreak” situations. Scabies mites of other animals may cause transient irritation, but they do not reside or reproduce in human hosts.

TREATMENT SCABIES

Permethrin cream (5%) is less toxic than 1% lindane preparations and is effective against lindane-tolerant infestations. Scabicides are applied thinly but thoroughly behind the ears and from the neck down after bathing—with careful application to interdigital spaces and the umbilicus and under the fingernails—and are removed

8–14 h later with soap and water. Successful treatment of crusted scabies requires preapplication of a keratolytic agent such as 6% salicylic acid and then of scabicides to the scalp, face, and ears. Repeated treatments or the sequential use of several agents may be necessary. Ivermectin has not been approved by the U.S. Food and Drug Administration (FDA) for treatment of any form of scabies, but a single oral dose (200 µg/kg) is effective in otherwise healthy persons; patients with crusted scabies may require two doses separated by an interval of 1–2 weeks. All FDA-approved scabicides are available solely by prescription.

Within 1 day of effective treatment, scabies infestations become noncommunicable, but the pruritic hypersensitivity dermatitis induced by the now-dead mites and their remnant products frequently persists for weeks. Unnecessary re-treatment with topical agents may provoke contact dermatitis. Antihistamines, salicylates, and calamine lotion relieve itching during treatment, and topical glucocorticoids are useful for pruritus that lingers after effective treatment. To prevent reinfestations, bedding and clothing should be washed and dried on high heat or heat-pressed. Close contacts of confirmed cases, even if asymptomatic, should be treated simultaneously.

CHIGGERS AND OTHER BITING MITES

Chiggers are the larvae of trombiculid (harvest) mites that normally feed on mice in grassy or brush-covered sites in tropical, subtropical, and (less frequently) temperate areas during warm months. They reside on low vegetation and attach themselves to passing mammalian hosts. While feeding, larvae secrete saliva with proteolytic enzymes to create a tube-like invagination in the host's skin; this *stylostome* allows the mite to imbibe tissue fluids. The stylostomal saliva is highly antigenic and causes exceptionally pruritic papular, papulovesicular, or papulourticarial lesions (≤2 cm in diameter). In people previously sensitized to salivary antigens, the papules develop within hours of attachment. While attached, mites appear as tiny red vesicles on the skin. Generally, lesions vesiculate and develop a hemorrhagic base. Scratching, however, invariably destroys the body of a mite. Itching and burning often persist for weeks. The rash is common on the ankles and areas where clothing obstructs the further wanderings of the mites. Repellents are useful for preventing chigger bites.

Many kinds of mites that are associated with peridomestic birds and rodents are particularly bothersome when they invade homes and bite people. In North America, the northern fowl mite, chicken mite, tropical rat mite, and house mouse mite normally feed on poultry, various songbirds, and small mammals and are abundant in and around their hosts' nests. After their natural hosts die or leave the nest, these mites frequently invade human habitations. Although the mites are rarely seen because of their small size, their bites can be painful and pruritic. Once confirmed as the cause of irritation, rodent- and bird-associated mites are best eliminated by excluding their hosts, removing the nests, and cleaning and treating the nesting area with appropriate acaricides. *Pyemotes* and other mites that infest grain, straw, cheese, hay, or other products occasionally produce similar episodes of rash and discomfort and may produce a unique dermatologic "comet sign" lesion—a paisley-shaped urticarial plaque.

Diagnosis of mite-induced dermatitides (including those caused by chiggers) relies on confirmation of the mite's identity or elicitation of a history of exposure to the mite's source. Oral antihistamines or topical steroids may suppress mite-induced pruritus temporarily but do not eliminate the mites.

TICK BITES AND TICK PARALYSIS

Ticks attach superficially to skin and feed painlessly; blood is their only food. Their salivary secretions are biologically active and can produce local reactions, induce fevers, and cause paralysis in addition to transmitting diverse pathogens. The two main families of ticks are the hard (*ixodid*) and soft (*argasid*) ticks. Generally, soft ticks attach for <1 h, leaving red macules after they drop off. Some species in Africa, the western United States, and Mexico produce painful hemorrhagic

lesions. Hard ticks are much more common and transmit most of the tick-borne infections that are familiar to physicians and patients. Hard ticks attach to the host and feed for several days or sometimes for >1 week. At the site of hard-tick bites, small areas of induration, often purpuric, develop and may be surrounded by an erythematous rim. A necrotic eschar, called a *tâche noire*, occasionally develops. Chronic nodules (persistent tick-bite granulomas) can be several centimeters in diameter and may linger for months after the feeding tick has been removed. These granulomas can be treated with injected intralesional glucocorticoids or by surgical excision. Tick-induced fever, unassociated with transmission of any pathogen, is often accompanied by headache, nausea, and malaise but usually resolves ≤36 h after the tick is removed.

Tick paralysis, an acute ascending flaccid paralysis that resembles Guillain-Barré syndrome, is believed to be caused by one or more toxins in tick saliva that block neuromuscular transmission and decrease nerve conduction. This rare complication has followed the bites of more than 60 kinds of ticks, although in the United States dog and wood ticks (*Dermacentor* species) are most commonly involved. Weakness begins symmetrically in the lower extremities ≤6 days after the tick's attachment, ascends symmetrically during several days, and may culminate in complete paralysis of the extremities and cranial nerves. Deep tendon reflexes are diminished or absent, but sensory examination and findings on lumbar puncture are typically normal. Removal of the tick generally leads to rapid improvement within a few hours and complete recovery after several days, although the patient's condition may continue to deteriorate for a full day. Failure to remove the tick may lead to dysarthria, dysphagia, and ultimately death from aspiration or respiratory paralysis. Diagnosis depends on finding the tick, which is often hidden beneath scalp hair. An antiserum to the saliva of *Ixodes holocyclus*, the usual cause of tick paralysis in Australia, effectively reverses paralysis caused by these ticks.

Removal of hard ticks during the first 36 h of attachment nearly always prevents transmission of the agents of Lyme disease, babesiosis, anaplasmosis, and ehrlichiosis, although several tick-borne viruses may be transmitted more quickly. Ticks should be removed by traction with fine-tipped forceps placed firmly around the tick's mouthparts. Careful handling (to avoid rupture of ticks) and use of gloves may avert accidental contamination with pathogens contained in tick fluids. Use of occlusive dressings, heat, or other substances (in an attempt to induce the tick to detach) merely delay tick removal. Afterward, the site of attachment should be disinfected. Tick mouthparts sometimes remain in the skin but generally are shed spontaneously within days without excision. Although somewhat controversial, current guidelines from the Centers for Disease Control and Prevention suggest that, rather than awaiting the onset of erythema migrans, the results of tick testing, or seroconversion to antigens diagnostic for Lyme disease, administering prophylaxis with a single oral dose of doxycycline (200 mg) within 72 h of tick removal is appropriate in adult patients with bites thought to be associated with deer ticks (Fig. 475-1) in Lyme disease–endemic areas from Maryland to Maine and in Wisconsin and Minnesota.

LOUSE INFESTATION (PEDICULIASIS AND PHTHIRIASIS)

Nymphs and adults of all three kinds of human lice feed at least once a day, ingesting human blood exclusively. Head lice (*Pediculus capitis*) infest mainly the hair of the scalp, body lice (*Pediculus humanus*) the clothing, and crab or pubic lice (*Phthirus pubis*) mainly the hair of the pubis. The saliva of lice produces a pruritic morbilliform or urticarial rash in some sensitized persons. Female head and pubic lice cement their eggs (nits) firmly to hair, whereas female body lice cement their eggs to clothing, particularly to threads along clothing seams. After ~10 days of development within the egg, a nymph hatches. Empty eggs may remain affixed for months thereafter.

In North America, head lice infest ~1% of elementary school-age children. Head lice are transmitted mainly by direct head-to-head contact rather than by fomites such as shared headgear, bed linens, hairbrushes, and other grooming implements. Chronic infestations by head lice tend to be asymptomatic. Pruritus, due mainly to hypersensitivity to the louse's saliva, generally is transient and mild. Head lice



FIGURE 475-1 Deer ticks (*Ixodes scapularis*, black-legged ticks) on a U.S. penny: larva (below ear), nymph (right), adult male (above), and adult female (left).

removed from a person succumb to desiccation and starvation within ~1 day. Head lice are not known to serve as a natural vector for any pathogens.

Body lice remain on clothing except when feeding and generally succumb in ≤ 2 days if separated from their host. In most Western countries, body lice are generally found on a small proportion of indigent persons but may become increasingly prevalent after upheaval associated with natural or human-caused disasters, when homeless victims are in close contact with infested individuals with whom they share accommodations. Body lice are acquired by direct contact or by sharing of infested clothing and bedding. These lice are vectors for the agents of louse-borne (epidemic) typhus (Chap. 211), louse-borne relapsing fever (Chap. 209), and trench fever (Chap. 197). Pruritic lesions from their bites are particularly common around the neckline. Chronic infestations result in a postinflammatory hyperpigmentation and thickening of skin known as *vagabond's disease*.

The crab or pubic louse is transmitted mainly by sexual contact. These lice occur predominantly on pubic hair and less frequently on axillary or facial hair, including the eyelashes. Children and adults may acquire pubic lice by sexual or close nonsexual contact. Intensely pruritic, bluish macules ~3 mm in diameter (*maculae ceruleae*) develop at the site of bites. Blepharitis commonly accompanies infestations of the eyelashes.

Pediculiasis is often suspected upon the detection of nits firmly cemented to hairs or in clothing. Many bona fide nits, however, are dead or hatched relics of prior infestation, and pseudo-nits are frequently misconstrued to be signs of a louse infestation. Confirmation of a louse infestation, therefore, best relies on the discovery of a live louse.

TREATMENT LOUSE INFESTATION

Generally, treatment is warranted only if live lice are discovered. The presence of nits alone is evidence of a former—not necessarily current—infestation. Mechanical removal of lice and their eggs with a fine-toothed louse or nit comb (Fig. 475-2) often fails to eliminate infestations. Treatment of newly identified active infestations generally relies on a 10-min topical application of ~1% permethrin or pyrethrins, with a second application ~10 days later. Lice persisting after this treatment may be resistant to pyrethroids. Chronic infestations may be treated for ≤ 12 h with 0.5% malathion. Lindane is applied for just 4 min but seems less effective and may pose a greater risk of adverse reactions, particularly when misused. Resistance of head lice to permethrin, malathion, and lindane has been reported. Newer FDA-approved topical pediculicides contain benzyl alcohol, dimethicone, spinosad, and ivermectin. Although children infested by head lice—or those who simply have remnant nits from a prior infestation—are frequently isolated or excluded



FIGURE 475-2 Adult female human head louse (*Pediculus capitis*) on a nit (louse-egg) comb.

from school, this practice increasingly is seen as unjustified and ineffective.

Body lice usually are eliminated by bathing and by changing to laundered clothes. Application of topical pediculicides from head to foot may be necessary for hirsute patients. Clothes and bedding are effectively deloused by heating in a clothes dryer at $\geq 55^\circ\text{C}$ ($\geq 131^\circ\text{F}$) for 30 min or by heat-pressing. Emergency mass delousing of persons and clothing may be warranted during periods of civil strife and after natural disasters to reduce the risk of pathogen transmission by body lice.

Pubic louse infestations are treated with topical pediculicides, except for eyelid infestations (*pthiriasis palpebrum*), which generally respond to a coating of petrolatum applied for 3–4 days.

MYIASIS (FLY INFESTATION)

Myiasis refers to infestations by several kinds of fly larvae (maggots) that invade living or necrotic tissues or body cavities and produce different clinical syndromes, depending on the species of fly.

In forested parts of Central and South America, larvae of the human botfly (*Dermatobia hominis*) produce furuncular (boil-like) papules or subcutaneous nodules ≤ 3 cm in diameter. A gravid adult female botfly captures a mosquito or another bloodsucking insect and deposits her eggs on its abdomen. When the carrier insect attacks a human or bovine host several days later, the warmth and moisture of the host's skin stimulate the eggs to hatch. The emerging larvae promptly penetrate intact skin. After 6–12 weeks of development, mature larvae emerge from the skin and drop to the ground to pupate and then become adults.

The African tumbu fly (*Cordylobia anthropophaga*) deposits its eggs on damp sand or leaf litter or on drying laundry, particularly that contaminated by urine or sweat. Larvae hatch from eggs upon contact with a host's body and penetrate the skin, producing boil-like lesions from which mature larvae emerge ~9 days later. Furuncular myiasis is suggested by uncomfortable lesions with a central breathing pore that emits bubbles when submerged in water. A sensation of movement under the patient's skin may cause severe emotional distress.

Larvae that cause furuncular myiasis may be induced to emerge if the air pore is coated with petrolatum or another occlusive substance. Removal may be facilitated by injection of a local anesthetic into the surrounding tissue, but surgical excision is sometimes necessary because upward-pointing spines of some species hold the larvae firmly in place.

Other fly larvae cause nonfuruncular myiasis. For example, larvae of the horse botfly (*Gasterophilus intestinalis*) emerge from eggs deposited on the horse's flanks and may come into contact with and infest human beings. After penetrating human skin, these larvae rarely mature but instead may migrate for weeks in the dermis. The resulting pruritic and serpiginous eruption resembles cutaneous larva migrans caused by canine or feline hookworms (Chap. 256). Larvae of rabbit and rodent botflies (*Cuterebra* species) occasionally cause dermal or tracheopulmonary myiasis.

Certain flies are attracted to blood and pus, laying their eggs on open or draining sores. Newly hatched larvae enter wounds or diseased skin. Larvae of several types of green bottle flies (*Lucilia/Phaenicia* species) usually remain superficial and confined to necrotic tissue. Specially raised, sterile “surgical maggots” are sometimes used intentionally for wound debridement. Larvae of screwworm flies, *Cochliomyia*, and the flesh fly invade viable tissues more deeply and produce large suppurating lesions. Larvae that infest wounds also may enter body cavities such as the mouth, nose, ears, sinuses, anus, vagina, and lower urinary tract, particularly in unconscious or otherwise debilitated patients. The consequences range from harmless colonization to destruction of the nose, meningitis, and deafness. Treatment involves removal of maggots and debridement of tissue.

The maggots responsible for furuncular and wound myiasis also may cause ophthalmomyiasis. Sequelae include nodules in the eyelid, retinal detachment, and destruction of the globe. Most instances in which maggots are found in human feces result from deposition of eggs or larvae by flies on recently passed stools, not from an intestinal maggot infestation.

PENTASTOMIASIS

Pentastomids (tongue worms) inhabit the respiratory passages of reptiles and carnivorous mammals. Human infestation by *Linguatula serrata* is common in the Middle East and results from the consumption of encysted larval stages in raw liver or lymph nodes of sheep and goats, which are true intermediate hosts for the tongue worms. Larvae migrate to the nasopharynx and produce an acute self-limiting syndrome—known as *halzoun* or *marrara*—characterized by pain and itching of the throat and ears, coughing, hoarseness, dysphagia, and dyspnea. Severe edema may cause obstruction that requires tracheostomy. In addition, ocular invasion has been described. Diagnostic larvae measuring ≤ 10 mm in length appear in copious nasal discharge or vomitus. Individuals become infected with another type of tongue worm, *Armillifer armillatus*, by consuming its eggs in contaminated food or drink or after handling the definitive host, the African python. Larvae encyst in various organs but rarely cause symptoms. Cysts may require surgical removal as they enlarge during molting, but they usually are encountered as an incidental finding at autopsy. Parasite-induced lesions may be misinterpreted as a malignancy, with the correct diagnosis confirmed histopathologically. Cutaneous larva migrans-type syndromes of other pentastomes have been reported from Southeast Asia and Central America.

LEECH INFESTATIONS

Medically important leeches are annelid worms that attach to their hosts with chitinous cutting jaws and draw blood through muscular suckers. The medicinal leech (*Hirudo medicinalis*) is still used occasionally for medical purposes to reduce venous congestion in surgical flaps or replanted body parts. This practice has been complicated by intractable bleeding, wound infections, myonecrosis, and sepsis due to *Aeromonas hydrophila*, which colonizes the gullets of commercially available leeches.

Ubiquitous aquatic leeches that parasitize fish, frogs, and turtles readily attach to the skin of human beings and avidly suck blood. More notorious are arboreal land leeches that live among moist vegetation of tropical rain forests. Attachment is usually painless, and the leeches will detach themselves when satiated with a blood meal. Hirudin, a powerful anticoagulant secreted by the leech, causes continued bleeding after the leech has detached. Healing of a leech-bite wound is slow, and bacterial infections are not uncommon. Several kinds of aquatic leeches in Africa, Asia, and southern Europe can enter the mouth, nose, and genitourinary tract and attach to mucosal surfaces at sites as deep as the esophagus and trachea. Externally attached leeches generally drop off after they have engorged, but removal is hastened by gentle scraping aside of the anterior and posterior suckers the leech uses for attachment and feeding. Some authorities dispute the wisdom of removing leeches with alcohol, salt, vinegar, insect repellent, a flame or heated instrument, or applications of other noxious substances.

Internally attached leeches may detach on exposure to gargled saline or may be removed by forceps.

SPIDER BITES

Of the more than 30,000 recognized species of spiders, only ~ 100 defend themselves aggressively and have fangs sufficiently long to penetrate human skin. The venom that some spiders use to immobilize and digest their prey can cause necrosis of skin and systemic toxicity. Whereas the bites of most spiders are painful but not harmful, envenomations by recluse or fiddleback spiders (*Loxosceles* species) and widow spiders (*Latrodectus* species) may be life-threatening. Identification of the offending spider is important because specific treatments exist for bites of widow spiders and because injuries attributed to spiders are frequently due to other causes. Except in cases where the patient actually observes a spider immediately associated with the bite or fleeing from the site, lesions reported as spider-bite reactions are most often due to other injuries or to infections with bacteria such as methicillin-resistant *Staphylococcus aureus* (MRSA).

Recluse Spider Bites and Necrotic Arachnidism Brown recluse spiders live mainly in the south-central United States and have close relatives in Central and South America, Africa, and the Middle East. Bites by brown recluse spiders usually cause only minor injuries, with edema and erythema. Envenomation, however, occasionally causes severe necrosis of skin and subcutaneous tissue and more rarely causes systemic hemolysis. These spiders are not aggressive toward humans and will bite only if threatened or pressed against the skin. They hide under rocks and logs or in caves and animal burrows. They invade homes and seek dark and undisturbed hiding spots in closets, in folds of clothing, or under furniture and rubbish in storage rooms, garages, and attics. Despite their impressive abundance in some homes, these spiders rarely bite humans. Bites tend to occur while the victim is dressing and are sustained primarily on the hands, arms, neck, and lower abdomen.

Initially, the bite is painless or may produce a stinging sensation. Within the next few hours, the site becomes painful and pruritic, with central induration surrounded by a pale ischemic zone that itself is encircled by a zone of erythema. In most cases, the lesion resolves without treatment in just a few days. In severe cases, the erythema spreads, and the center of the lesion becomes hemorrhagic or necrotic with an overlying bulla. A black eschar forms and sloughs several weeks later, leaving an ulcer that eventually may create a depressed scar. Healing usually takes place in ≤ 6 months but may take as long as 3 years if adipose tissue is involved. Local complications include injury to nerves and secondary bacterial infection. Fever, chills, weakness, headache, nausea, vomiting, myalgia, arthralgia, maculopapular rash, and leukocytosis may develop ≤ 72 h after the bite. Reports of deaths attributed to bites of North American brown recluse spiders have not been verified.

TREATMENT RECLUSE SPIDER BITES

Initial management includes RICE (rest, ice, compression, elevation). Analgesics, antihistamines, antibiotics, and tetanus prophylaxis should be administered if indicated. Early debridement or surgical excision of the wound without closure delays healing. Routine use of antibiotics or dapsone is unnecessary. Patients should be monitored closely for signs of hemolysis, renal failure, and other systemic complications.

Widow Spider Bites The black widow spider, common in the southeastern United States, measures ≤ 1 cm in body length and 5 cm in leg span and is shiny black with a red hourglass marking on the ventral abdomen. Other dangerous *Latrodectus* species occur elsewhere in temperate and subtropical parts of the world. The bites of the female widow spiders are notorious for their potent neurotoxins.

Widow spiders spin their webs under stones, logs, plants, or rock piles and in dark spaces in barns, garages, and outhouses. Bites are most common in the summer and early autumn and occur when a web is disturbed or a spider is trapped or provoked. The initial bite

2748 is perceived as a sharp pinprick or may go unnoticed. Fang-puncture marks are uncommon. The venom that is injected does not produce local necrosis, and some persons experience no other symptoms. α -Latrotoxin, the most active component of the venom, binds irreversibly to presynaptic nerve terminals and causes release and eventual depletion of acetylcholine, norepinephrine, and other neurotransmitters from those terminals. Painful cramps may spread within 60 min from the bite site to large muscles of the extremities and trunk. Extreme rigidity of the abdominal muscles and excruciating pain may suggest peritonitis, but the abdomen is not tender on palpation and surgery is not warranted. The pain begins to subside during the first 12 h but may recur during several days or weeks before resolving spontaneously. A wide range of other neurologic sequelae may include salivation, diaphoresis, vomiting, hypertension, tachycardia, labored breathing, anxiety, headache, weakness, fasciculations, paresthesia, hyperreflexia, urinary retention, uterine contractions, and premature labor. Rhabdomyolysis and renal failure have been reported, and respiratory arrest, cerebral hemorrhage, or cardiac failure may end fatally, especially in very young, elderly, or debilitated persons.

TREATMENT WIDOW SPIDER BITES

Treatment consists of RICE and tetanus prophylaxis. Hypertension that does not respond to analgesics and antispasmodics (e.g., benzodiazepines or methocarbamol) requires specific antihypertensive medication. The efficacy and safety of antivenoms for black widow and redback spiders are controversial because of concerns about potential anaphylaxis or serum sickness.

Tarantulas and Other Spiders Tarantulas are hairy spiders of which 30 species are found in the United States, mainly in the Southwest. The tarantulas that have become popular household pets are usually imported from Central or South America. Tarantulas bite only when threatened and usually cause no more harm than a bee sting, but on occasion the venom causes deep pain and swelling. Several species of tarantulas are covered with urticating hairs that are brushed off in the thousands when a threatened spider rubs its hind legs across its dorsal abdomen. These hairs can penetrate human skin and produce pruritic papules that may persist for weeks. Failure to wear gloves or to wash the hands after handling the Chilean Rose tarantula, a popular pet spider, has resulted in transfer of hairs to the eye with subsequent devastating ocular inflammation. Treatment of bites includes local washing and elevation of the bitten area, tetanus prophylaxis, and analgesic administration. Antihistamines and topical or systemic glucocorticoids are given for exposure to urticating hairs.

Atrax robustus, a funnel-web spider of Australia, and *Phoneutria* species, the South American banana spiders, are among the most dangerous spiders in the world because of their aggressive behavior and potent neurotoxins. Envenomation by *A. robustus* causes a rapidly progressive neuromotor syndrome that can be fatal within 2 h. The bite of a banana spider causes severe local pain followed by profound systemic symptoms and respiratory paralysis that can lead to death within 2–6 h. Specific antivenoms for use after bites by each of these spiders are available. Yellow sac spiders (*Cheiracanthium* species) are common in homes worldwide. Their bites, though painful, generally lead to only minor erythema, edema, and pruritus.

SCORPION STINGS

Scorpions are arachnids that feed on ground-dwelling arthropods and small lizards. They paralyze their prey and defend themselves by injecting venom from a stinger on the tip of the tail. Painful but relatively harmless scorpion stings need to be distinguished from the potentially lethal envenomations that are produced by ~30 of the ~1000 known species and that cause more than 5000 deaths worldwide each year. Scorpions are nocturnal and remain hidden during the day in crevices or burrows or under wood, loose bark, or rocks. They occasionally enter houses and tents and may hide in shoes, clothing, or bedding. Scorpions sting humans only when threatened.

Of the 40 or so scorpion species in the United States, only bark scorpions—e.g., *Centruroides sculpturatus* (*C. exilicauda*) in the Southwest—produce venom that is potentially lethal to humans. This venom contains neurotoxins that cause sodium channels to remain open. Such envenomations usually are associated with little swelling, but prominent pain, paresthesia, and hyperesthesia can be accentuated by tapping on the affected area (the *tap test*). These symptoms soon spread to other locations; dysfunction of cranial nerves and hyperexcitability of skeletal muscles develop within hours. Patients present with restlessness, blurred vision, abnormal eye movements, profuse salivation, lacrimation, rhinorrhea, slurred speech, difficulty in handling secretions, diaphoresis, nausea, and vomiting. Muscle twitching, jerking, and shaking may be mistaken for a seizure. Complications include tachycardia, arrhythmias, hypertension, hyperthermia, rhabdomyolysis, and acidosis. Symptoms progress to maximal severity in ~5 h and subside within a day or two, although pain and paresthesia can last for weeks. Fatal respiratory arrest is most common among young children and the elderly.

Envenomations by *Leiurus quinquestriatus* in the Middle East and North Africa, by *Mesobuthus tamulus* in India, by *Androctonus* species along the Mediterranean littoral and in North Africa and the Middle East, and by *Tityus serrulatus* in Brazil cause massive release of endogenous catecholamines with hypertensive crises, arrhythmias, pulmonary edema, and myocardial damage. Acute pancreatitis occurs with stings of *Tityus trinitatis* in Trinidad, and central nervous toxicity complicates stings of *Parabuthus* and *Buthotus* scorpions of South Africa. Tissue necrosis and hemolysis may follow stings of the Iranian *Hemiscorpius lepturus*.

Stings of most other species cause immediate sharp local pain followed by edema, ecchymosis, and a burning sensation. Symptoms typically resolve within a few hours, and skin does not slough. Allergic reactions to the venom sometimes develop.

TREATMENT SCORPION STINGS

Identification of the offending scorpion helps to determine the course of treatment. Stings of nonlethal species require at most ice packs, analgesics, or antihistamines. Because most victims experience only local discomfort, they can be managed at home with instructions to return to the emergency department if signs of cranial-nerve or neuromuscular dysfunction develop. Aggressive supportive care and judicious use of antivenom can reduce or eliminate deaths from more severe envenomations. Keeping the patient calm and applying pressure dressings and cold packs to the sting site are measures that decrease the absorption of venom. A continuous IV infusion of midazolam controls the agitation, flailing, and involuntary muscle movements produced by scorpion stings. Close monitoring during treatment with this drug and other sedatives or narcotics is necessary for persons with neuromuscular symptoms because of the risk of respiratory arrest. Hypertension and pulmonary edema respond to nifedipine, nitroprusside, hydralazine, or prazosin. Dangerous bradyarrhythmias can be controlled with atropine.

Commercially prepared antivenoms are available in several countries for some of the most dangerous species. An FDA-approved *C. sculpturatus* antivenom in horse serum is now available. IV administration of antivenom rapidly reverses cranial-nerve dysfunction and muscular symptoms. Although effective, cost analyses suggest that antivenoms should be reserved for only the most severe envenomations.

HYMENOPTERA STINGS

Insects that sting in defense or to subdue their prey belong to the order Hymenoptera, which includes bees, wasps, hornets, yellow jackets, and ants. Their venoms contain a wide array of amines, peptides, and enzymes that cause local and systemic reactions. Although the toxic effect of multiple stings can be fatal to a human, nearly all of the ≥ 100 deaths due to hymenopteran stings in the United States each year result from allergic reactions.

Bee and Wasp Stings The stinger of the honeybee (*Apis mellifera*) is unique in being barbed. When a bee stings a foe, its stinging apparatus and attached venom sac tear loose from its body. Muscular contraction of the venom sac continues to inject venom into the skin. Other kinds of bees, ants, and wasps have smooth stinging mechanisms and can sting numerous times in succession. Honeybees, bumblebees, and social wasps generally attack only when a colony is disturbed. Africanized honeybees (now present in South and Central America and the southern and western United States) respond to minimal intrusions more aggressively. Whereas the sting of an Africanized bee contains less venom than that of its non-Africanized relatives, victims tend to sustain far more stings and therefore to receive a far greater overall volume of venom. Most patients who report having sustained a “bee sting,” however, are more likely to have encountered stinging wasps instead.

The venoms of different species of hymenopterans are biochemically and immunologically distinct. Direct toxic effects are mediated by mixtures of low-molecular-weight compounds such as serotonin, histamine, acetylcholine, and several kinins. Polypeptide toxins in honeybee venom include mellitin that damages cell membranes, mast cell-degranulating protein that causes histamine release, the neurotoxin apamin, and the anti-inflammatory compound adolapin. Enzymes in venom include hyaluronidase and phospholipases. There appears to be little cross-sensitization between the venoms of honeybees and wasps.

Uncomplicated hymenopteran stings cause immediate pain, a wheal-and-flare reaction, and local edema, all of which usually subside in a few hours. Multiple stings can lead to vomiting, diarrhea, generalized edema, dyspnea, hypotension, and non-anaphylactic circulatory collapse. Rhabdomyolysis and intravascular hemolysis may cause renal failure. Death from the direct (nonallergic) effects of venom has followed stings of several hundred honeybees. Stings to the tongue or mouth may induce life-threatening edema of the upper airways.

Large local reactions accompanied by erythema, edema, warmth, and tenderness that spread ≥ 10 cm around the sting site over 1–2 days are not uncommon. These reactions may resemble bacterial cellulitis but are caused by hypersensitivity rather than by secondary infection. Such reactions tend to recur on subsequent exposure but are seldom accompanied by anaphylaxis and are not prevented by venom immunotherapy.

An estimated 0.4–4.0% of the U.S. population exhibits clinical immediate-type hypersensitivity to hymenopteran stings, and 15% may have asymptomatic sensitization manifested by positive skin tests. Persons who experience severe allergic reactions are likely to have similar reactions after subsequent stings by the same or closely related species. Occasionally, persons who have had mild reactions earlier in life will experience more serious reactions to subsequent stings. Mild anaphylactic reactions from insect stings, as from other causes, consist of nausea, abdominal cramping, generalized urticaria, flushing, and angioedema. Serious reactions, including upper airway edema, bronchospasm, hypotension, and shock, may be rapidly fatal. Severe reactions usually begin within 10 min of the sting and only rarely develop after 5 h.

TREATMENT BEE AND WASP STINGS

Honeybee stingers embedded in the skin should be removed as soon as possible to limit the quantity of venom delivered. The stinger and venom sac may be scraped off with a blade, the edge of a credit card, or a fingernail or may be removed with forceps. The site should be cleansed and disinfected and ice packs applied to slow the spread of venom. Elevation of the affected site and administration of analgesics, oral antihistamines, and topical calamine lotion help relieve symptoms. Large local reactions may require a short course of oral therapy with glucocorticoids. Patients with numerous stings should be monitored for 24 h for evidence of renal failure or coagulopathy.

Anaphylaxis is treated with SC injection of 0.3–0.5 mL of epinephrine hydrochloride in a 1:1000 dilution; treatment is repeated every 20–30 min as necessary. IV epinephrine (2–5 mL of a 1:10,000

solution administered by slow push) is indicated for profound shock. A tourniquet may slow the spread of venom. Parenteral antihistamines, fluid resuscitation, bronchodilators, supplemental oxygen, intubation, and vasopressors may be required. Patients should be observed for 24 h for recurrent anaphylaxis.

Persons with a history of allergy to insect stings should carry an anaphylaxis kit with a preloaded syringe containing epinephrine for self-administration. These patients should seek medical attention immediately after using the kit.

Repeated injections of purified venom produce a blocking IgG antibody response to venom and reduce the incidence of recurrent anaphylaxis. Honeybee, wasp, and yellow jacket venoms are commercially available for desensitization and for skin testing. Results of skin tests and venom-specific radioallergosorbent tests (RASTs) aid in the selection of patients for immunotherapy and guide the design of such treatment.

STINGING ANTS

Stinging fire ants are an important medical problem in the United States. Imported fire ants (*Solenopsis* species) infest southern states from Texas to North Carolina, with colonies now established in California, New Mexico, Arizona, and Virginia. Slight disturbances of their mound nests have provoked massive outpourings of ants and as many as 10,000 stings on a single person. Elderly and immobile persons are at high risk for attacks when fire ants invade dwellings.

Fire ants attach to skin with powerful mandibles and rotate their bodies while repeatedly injecting venom with posteriorly situated stingers. The alkaloid venom consists of cytotoxic and hemolytic piperidines and several proteins with enzymatic activity. The initial wheal-and-flare reaction, burning, and itching resolve in ~ 30 min, and a sterile pustule develops within 24 h. The pustule ulcerates over the next 48 h and then heals in ≥ 1 week. Large areas of erythema and edema lasting several days are not uncommon and in extreme cases may compress nerves and blood vessels. Anaphylaxis occurs in fewer than 2% of victims; seizures and mononeuritis have been reported. Stings are treated with ice packs, topical glucocorticoids, and oral antihistamines. Pustules should be cleansed and then covered with bandages and antibiotic ointment to prevent bacterial infection. Epinephrine administration and supportive measures are indicated for anaphylactic reactions. Fire ant whole-body extracts are available for skin testing and immunotherapy, which appears to lower the rate of anaphylactic reactions.

European fire (red) ants (*Myrmica rubra*) have recently become public health pests in the northeastern United States and southern Canada. The western United States is home to harvester ants (*Pogonomyrmex* species). The painful local reaction that follows harvester ant stings often extends to lymph nodes and may be accompanied by anaphylaxis. The bullet or conga ant (*Paraponera clavata*) of South America is known locally as *hormiga veinticuatro* (“24-hour ant”), a designation that refers to the 24 h of throbbing, excruciating pain following a sting that delivers the potent paralyzing neurotoxin poneratoxin.

DIPTERAN (FLY AND MOSQUITO) BITES

In the process of feeding on vertebrate blood and tissue fluids, adults of certain flies inflict painful bites, produce local allergic reactions, and may transmit pathogenic agents. Bites of mosquitoes, tiny “no-see-um” (ceratopogonid) midges, and phlebotomine sand flies typically produce a wheal and a pruritic papule. Small humpbacked black flies (simuliids) lacerate skin, resulting in a lesion with serosanguineous discharge that is often painful and pruritic. Regional lymphadenopathy, fever, or anaphylaxis occasionally ensues. The widely distributed deerflies and horseflies as well as the tsetse flies of Africa are stout flies measuring ≤ 25 mm in length that attack during the day and produce large and painful bleeding punctures. Houseflies (*Musca domestica*) do not consume blood but use rasping mouthparts to scarify skin and feed upon tissue fluids and salt. Beyond direct injury from bites of any kind

2750 of fly, risks include transmission of diverse pathogens and secondary infection of the lesion.

TREATMENT FLY AND MOSQUITO BITES

Treatment of fly bites is symptom based. Topical application of antipruritic agents, glucocorticoids, or antiseptic lotions may relieve itching and pain. Allergic reactions may require oral antihistamines. Antibiotics may be necessary for the treatment of large bite wounds that become secondarily infected.

FLEA BITES

Common human-biting fleas include the dog and cat fleas (*Ctenocephalides* species) and the rat flea (*Xenopsylla cheopis*), which infest their respective hosts and the hosts' nests and resting sites. Sensitized persons develop erythematous pruritic papules (*papular urticaria*) and occasionally vesicles and bacterial superinfection at the site of the bite. Symptom-based treatment consists of antihistamines, topical glucocorticoids, and topical antipruritic agents.

Flea infestations are eliminated by removal and treatment of animal nests, frequent cleaning of pet bedding, and application of contact and systemic insecticides to pets and the dwelling. Flea infestations in the home may be abated or prevented if pets are regularly treated with veterinary antiparasitic agents and insect growth regulators.

Tunga penetrans, like other fleas, is a wingless, laterally flattened insect that feeds on blood. Also known as the chigoe flea, sand flea, or jigger (not to be confused with the chigger), it occurs in tropical regions of Africa and the Americas. Adult female chigoes live in sandy soil and burrow under the skin, usually between toes, under nails, or on the soles of bare feet. Gravid chigoes engorge on the host's blood and grow from pinpoint to pea size during a 2-week interval. They produce lesions that resemble a white pustule with a central black depression and that may be pruritic or painful. Occasional complications include tetanus, bacterial infections, and autoamputation of toes (*ainhum*). Tungiasis is treated by removal of the intact flea with a sterile needle or scalpel, tetanus vaccination, and topical application of antibiotics.

HEMIPTERAN (TRUE BUG) BITES

Several true bugs of the family Reduviidae inflict bites that produce allergic reactions and are sometimes painful. The cone-nose bugs, so called because of their elongated heads, include the assassin and wheel bugs, which feed on other insects and bite vertebrates only in self-defense, and the kissing bugs, which routinely feed on vertebrate blood. The bites of the night-feeding kissing bugs are painless. Reactions to such bites depend on prior sensitization and include tender and pruritic papules, vesicular or bullous lesions, extensive urticaria, fever, lymphadenopathy, and (rarely) anaphylaxis. Bug bites are treated with topical antipruritics or oral antihistamines. Persons with anaphylactic reactions to reduviid bites should keep an epinephrine kit available. Some reduviids transmit *Trypanosoma cruzi*, the agent of New World trypanosomiasis (also called *Chagas disease*) (Chap. 252).

The cosmopolitan bedbugs (*Cimex* species) hide in crevices of mattresses, bed frames and other furniture, walls, and picture frames and under loose wallpaper. Bedbug populations have resurged, recently attaining levels and spreading to an extent not encountered since the mid-twentieth century. These bugs are now a common pest in homes, dormitories, and hotels; on cruise ships; and even in medical facilities. Generally, the bugs hide during the day and take blood meals at night. Their bite is painless, but minutes to days later, sensitized persons develop erythema, itching, and wheals around a central hemorrhagic punctum. Bedbugs are not known to transmit pathogens.

CENTIPEDE BITES AND MILLIPEDE DERMATITIS

The fangs of centipedes of the genus *Scolopendra* can penetrate human skin and deliver a venom that produces intense burning pain, swelling, erythema, and sterile lymphangitis. Dizziness, nausea, and anxiety are described occasionally, and rhabdomyolysis and renal failure have

been reported. Treatment includes washing of the site, application of cold dressings, oral analgesic administration or local lidocaine infiltration, and tetanus prophylaxis.

Millipedes are docile and do not bite, but some secrete defensive fluids that may burn and discolor human skin. Affected skin turns brown overnight and may blister and exfoliate. Secretions in the eye cause intense pain and inflammation that can result in corneal ulcers and even blindness. Management includes irrigation with copious amounts of water or saline, use of analgesics, and local care of denuded skin.

CATERPILLAR STINGS AND DERMATITIS

Caterpillars of several moth species are covered with hairs or spines that produce mechanical irritation and may contain or be coated with venom. Contact with these caterpillars or their hairs may lead to lepidopterism or caterpillar envenomation. The response typically consists of an immediate burning sensation followed by local swelling and erythema and occasionally by regional lymphadenopathy, nausea, vomiting, and headache. A rare reaction to a South American caterpillar, *Lonomia obliqua*, can cause disseminated coagulopathy and fatal hemorrhagic shock. In the United States, dermatitis is most often associated with caterpillars of the io, puss, saddleback, and brown-tail moths. Even contact with detached hairs of other caterpillars, such as gypsy moth larvae, can later produce a pruritic urticarial or papular rash called *erucism*. Spines may be deposited on tree trunks or drying laundry or may be airborne and cause irritation of the eyes and upper airways. Treatment of caterpillar stings consists of repeated application of adhesive or cellophane tape to remove the hairs, which can then be identified microscopically. Local ice packs, topical glucocorticoids, and oral antihistamines relieve symptoms.

BEETLE VESICATION AND DERMITIS

Several families of beetles have independently developed the ability to produce chemically unrelated vesicating toxins. When disturbed, blister beetles (family Meloidae) extrude cantharidin, a low-molecular-weight toxin that produces thin-walled blisters (≤ 5 cm in diameter) 2–5 h after contact. The blisters are not painful or pruritic unless broken and resolve without treatment in ≤ 10 days. Nephritis may follow unusually heavy cantharidin exposure. Contact occurs when individuals sit on the ground, work in the garden, or deliberately handle the beetles. The hemolymph of certain rove beetles (Staphylinidae) contains pederin, a potent vesicant. When these beetles are crushed or brushed against the skin, the released fluid causes painful, red, flaccid bullae. These beetles occur worldwide but are most numerous and problematic in parts of Africa (where they are called "Nairobi fly") and southwestern Asia. Ocular lesions may develop after impact with flying beetles at night or unintentional transfer of the vesicant on the fingers. Treatment is rarely necessary, although ruptured blisters should be kept clean and bandaged.

Larvae of common carpet beetles are adorned with dense arrays of ornate hairs called *hastisetae*. Contact with these larvae or their setae results in delayed dermal reactions in sensitized individuals. The lesions are commonly mistaken as bites of bedbugs.

DELUSIONAL INFESTATIONS

The groundless conviction that one is infested with arthropods or other parasites (Ekbohm's syndrome, delusory parasitosis, delusions of parasitosis, and perhaps Morgellons syndrome) is extremely difficult to treat and, unfortunately, is not uncommon. Patients describe uncomfortable sensations of something moving in or on their skin. Excoriations and self-induced ulcerations typically accompany the pruritus, dysesthesias, and imaginary insect bites. Patients often believe that some invisible or as yet undescribed creatures are infesting their skin, clothing, homes, or environment in general. Frequently, patients submit as evidence of infestation specimens that consist of plant-feeding and nonbiting peridomestic arthropods, pieces of skin, vegetable matter, lint, and other inanimate detritus. When evaluating a patient with possible delusional parasitosis, it is imperative to rule out true infestations and bites by arthropods, endocrinopathies, sensory disorders due to neuropathies, opiate and other drug use, environmental irritants

(e.g., fiberglass threads), and other causes of tingling or prickling sensations. Frequently, such patients repeatedly seek medical consultations, resist alternative explanations for their symptoms, and exacerbate their discomfort by self-treatment. Long-term pharmacotherapy with pimozide or other psychotropic agents has been more helpful than psychotherapy in treating this disorder. Patients with delusional parasitosis often develop the unshakeable conviction that they are

infested by a previously unknown pathogen, while their personal lives, family support, and employment collapse around them. **2751**

ACKNOWLEDGMENT

The substantial contributions of Andrew Spielman and James H. Maguire to this chapter in previous editions are gratefully acknowledged.

This page intentionally left blank

476e

Altitude Illness

Buddha Basnyat, Geoffrey Tabin

EPIDEMIOLOGY

Mountains cover one-fifth of the earth's surface; 38 million people live permanently at altitudes ≥ 2400 m, and 100 million people travel to high-altitude locations each year. Skiers in the Alps or Aspen; religious pilgrims to Lhasa or Kailash; trekkers and climbers to Kilimanjaro, Aconcagua, or Everest; and military personnel deployed to high-altitude locations are all at risk of developing acute mountain sickness (AMS), high-altitude cerebral edema (HACE), high-altitude pulmonary edema (HAPE), and other altitude-related problems. AMS is the benign form of altitude illness, whereas HACE and HAPE are life-threatening. Altitude illness is likely to occur above 2500 m but has been documented even at 1500–2500 m. In the Mount Everest region of Nepal, ~50% of trekkers who walk to altitudes >4000 m over ≥ 5 days develop AMS, as do 84% of people who fly directly to 3860 m. The incidences of HACE and HAPE are much lower than that of AMS, with estimates in the range of 0.1–4%.

PHYSIOLOGY

Ascent to a high altitude subjects the body to a decrease in barometric pressure that results in a decreased partial pressure of oxygen in the inspired gas in the lungs. This change leads in turn to less pressure driving oxygen diffusion from the alveoli and throughout the oxygen cascade. A normal initial “struggle response” to such an ascent includes increased ventilation—the cornerstone of acclimation—mediated by the carotid bodies. Hyperventilation may cause respiratory alkalosis and dehydration. Alkalosis may depress the ventilatory drive during sleep, with consequent periodic breathing and hypoxemia. During early acclimation, renal suppression of carbonic anhydrase and excretion of dilute alkaline urine combat alkalosis and tend to bring the pH of the blood to normal. Other physiologic changes during normal acclimation include increased sympathetic tone; increased erythropoietin levels, leading to increased hemoglobin levels and red blood cell mass; increased tissue capillary density and mitochondrial numbers; and higher levels of 2,3-bisphosphoglycerate, enhancing oxygen utilization. Even with normal acclimation, however, ascent to a high altitude decreases maximal exercise capacity (by ~1% for every 100 m gained above 1500 m) and increases susceptibility to cold injury due to peripheral vasoconstriction. Finally, if the ascent is made faster than the body can adapt to the stress of hypobaric hypoxemia, altitude-related disease states can result.

GENETICS



Hypoxia-inducible factor, which is important in high-altitude adaptation, controls transcriptional responses to hypoxia throughout the body and is involved in the release of vascular endothelial growth factor (VEGF) in the brain, erythropoiesis, and other pulmonary and cardiac functions at high altitudes. In particular, the gene *EPAS1*, which codes for transcriptional regulator hypoxia-inducible factor 2 α , appears to play an important role in the adaptation of Tibetans living at high altitude, resulting in lower hemoglobin concentrations than are found in the Han Chinese. For acute altitude illness, a single gene variant is unlikely to be found, but the differences in the susceptibility of individuals and populations, familial clustering of cases, and a positive association of some genetic variants all clearly support a role for genetics. Approximately 58 candidate genes have been tested, and at least one variant from 17 of these genes is associated with altitude illness.

ACUTE MOUNTAIN SICKNESS AND HIGH-ALTITUDE CEREBRAL EDEMA

AMS is a neurologic syndrome characterized by nonspecific symptoms (headache, nausea, fatigue, and dizziness), with a paucity of physical findings, developing 6–12 h after ascent to a high altitude. AMS is a clinical diagnosis. For uniformity in research studies, the Lake Louise Scoring System, created at the 1991 International Hypoxia Symposium, is generally used. AMS must be distinguished from exhaustion, dehydration, hypothermia, alcoholic hangover, and hyponatremia. AMS and HACE are thought to represent opposite ends of a continuum of altitude-related neurologic disorders. HACE (but not AMS) is an encephalopathy whose hallmarks are ataxia and altered consciousness with diffuse cerebral involvement but generally without focal neurologic deficits. Progression to these signal manifestations can be rapid. Papilledema and, more commonly, retinal hemorrhages may also develop. In fact, retinal hemorrhages occur frequently at ≥ 5000 m, even in individuals without clinical symptoms of AMS or HACE. It is unclear whether retinal hemorrhage and cerebral hemorrhage at high altitude are caused by the same mechanism.

Risk Factors The most important risk factors for the development of altitude illness are the rate of ascent and a prior history of high-altitude illness. Exertion is a risk factor, but lack of physical fitness is not. An attractive but still speculative hypothesis proposes that AMS develops in people who have inadequate cerebrospinal capacity to buffer the brain swelling that occurs at high altitude. Children and adults seem to be equally affected, but people >50 years of age may be less likely to develop AMS than younger people. Aging appears to be associated with blunting of cardiac chronotropic function and an increase in ventilatory response leading to maintenance of arterial oxygen saturation in hypoxia. Most studies reveal no gender difference in AMS incidence. A recent study showed that, in women, adaptive responses to hypoxia with aging are blunted by menopause but can be maintained with endurance training. Sleep desaturation—a common phenomenon at high altitude—is associated with AMS. Debilitating fatigue consistent with severe AMS on descent from a summit is also an important risk factor for death in mountaineers. A recently published prospective study involving trekkers and climbers who ascended to altitudes between 4000 m and 8848 m showed that high oxygen desaturation and low ventilatory response to hypoxia during exercise are independent predictors of severe altitude illness. However, because there may be overlap between groups of susceptible and nonsusceptible individuals, accurate cutoff values are hard to define. Prediction is made more difficult because the pretest probabilities of HAPE and HACE are low. Neck irradiation or surgery damaging the carotid bodies, respiratory tract infections, and dehydration appear to be other potential risk factors for altitude illness.

Pathophysiology The exact mechanisms causing AMS and HACE are unknown. Evidence points to a central nervous system process. MRI studies have suggested that vasogenic (interstitial) cerebral edema is a component of the pathophysiology of HACE. In the setting of high-altitude illness, the MRI findings shown in [Fig. 476e-1](#) are confirmatory of HACE, with increased signal in the white matter and particularly in the splenium of the corpus callosum. Quantitative analysis in a 3-tesla MRI study revealed that hypoxia is associated with mild vasogenic cerebral edema irrespective of AMS. This finding is in keeping with case reports of suddenly symptomatic brain tumors and of cranial nerve palsies without AMS at high altitudes. Vasogenic edema may become cytotoxic (intracellular) in severe HACE.

Impaired cerebral autoregulation in the presence of hypoxic cerebral vasodilation and altered permeability of the blood-brain barrier due to hypoxia-induced chemical mediators like histamine, arachidonic acid, and VEGF may all contribute to brain edema. In 1995, VEGF was first proposed as a potent promoter of capillary leakage

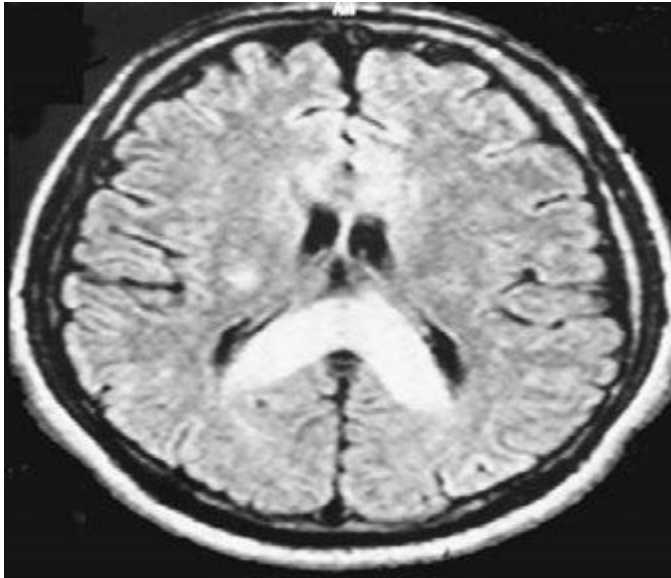


FIGURE 476e-1 T2 magnetic resonance image of the brain of a patient with high-altitude cerebral edema (HACE) shows marked swelling and a hyperintense posterior body and splenium of the corpus callosum (area with dense opacity). The patient, a climber, went on to climb Mount Everest about 9 months after this episode of HACE. (With permission from *Wilderness Environ Med* 15:53–55, 2004.)

in the brain at high altitude, and studies in mice have borne out this role. Although preliminary studies of VEGF in climbers have yielded inconsistent results regarding its association with altitude illness, indirect evidence of a role for this growth factor in AMS and HACE comes from the observation that dexamethasone, when used in the prevention and treatment of these conditions, blocks hypoxic upregulation of VEGF. Other factors in the development of cerebral edema may be the release of calcium-mediated nitric oxide and neuronally mediated adenosine, which may promote cerebral vasodilation.

Increased sympathetic activity triggered by hypoxia may also contribute to blood-brain barrier leakage. Enhanced optic-nerve sheath diameter with increasing severity of AMS has been noted and suggests an important role for increased intracranial pressures in the pathophysiology of AMS. Microhemorrhage formation caused by cytokines or damage through increased hydrostatic pressure is an important feature of HACE. Lesions in the globus pallidum (which is sensitive to hypoxia) leading to Parkinson's disease have also been reported to be complications of HACE. Finally, the effect of hypoxia on reactive oxygen species and the role of these species in clinical AMS are unclear.

The pathophysiology of the most common and prominent symptom of AMS—headache—remains unclear because the brain itself is an insensate organ; only the meninges contain trigeminal sensory nerve fibers. The cause of high-altitude headache is multifactorial. Various chemicals and mechanical factors activate a final common pathway, the trigeminovascular system. In the genesis of high-altitude headache, the response to nonsteroidal anti-inflammatory drugs and glucocorticoids provides indirect evidence for involvement of the arachidonic acid pathway and inflammation. Although the International Headache Society acknowledges that high altitude may be a trigger for migraine, it is unclear whether high-altitude headache and migraine share the same pathophysiology.

Prevention and Treatment (Table 476e-1) Gradual ascent, with adequate time for acclimation, is the best method for the prevention of altitude illness. Even though there may be individual variation in the rate of acclimation, a graded ascent of ≤ 400 m from the previous day's sleeping altitude is recommended above 3000 m, and taking every third day of gain in sleeping altitude as an extra day for acclimation is helpful. Spending one night at an intermediate altitude before proceeding to a higher altitude may enhance acclimation and attenuate the risk of AMS. Another protective factor in AMS is high-altitude exposure

TABLE 476e-1 MANAGEMENT OF ALTITUDE ILLNESS

Condition	Management
Acute mountain sickness (AMS), mild ^a	Discontinuation of ascent Treatment with acetazolamide (250 mg q12h) Descent ^b
AMS, moderate ^a	Immediate descent for worsening symptoms Use of low-flow oxygen if available Treatment with acetazolamide (250 mg q12h) and/or dexamethasone (4 mg q6h) ^c Hyperbaric therapy ^d
High-altitude cerebral edema (HACE)	Immediate descent or evacuation Administration of oxygen (2–4 L/min) Treatment with dexamethasone (8 mg PO/IM/IV; then 4 mg q6h) Hyperbaric therapy if descent is not possible
High-altitude pulmonary edema (HAPE)	Immediate descent or evacuation Minimization of exertion while patient is kept warm Administration of oxygen (4–6 L/min) to bring O ₂ saturation to >90% Adjunctive therapy with nifedipine ^e (30 mg, extended-release, q12h) Hyperbaric therapy if descent is not possible

^aCategorization of cases as mild or moderate is a subjective judgment based on the severity of headache and the presence and severity of other manifestations (nausea, fatigue, dizziness, insomnia). ^bNo fixed altitude is specified; the patient should descend to a point below that at which symptoms developed. ^cAcetazolamide treats and dexamethasone masks symptoms. For prevention (as opposed to treatment), 125–250 mg of acetazolamide q12h or (when acetazolamide is contraindicated—e.g., in people with sulfa allergy) 4 mg of dexamethasone q12h may be used. ^dIn hyperbaric therapy, the patient is placed in a portable altitude chamber or bag to simulate descent. ^eNifedipine at this dose is also effective for the prevention of HAPE, as is salmeterol (125 mg inhaled twice daily), tadalafil (10 mg twice daily), or dexamethasone (8 mg twice daily).

during the preceding 2 months; for example, the incidence and severity of AMS at 4300 m are reduced by 50% with an ascent after 1 week at an altitude of ≥ 2000 m rather than with an ascent from sea level. Studies have examined whether exposure to a normobaric hypoxic environment (in a room or a tent) before an ascent can provide protection against AMS. In double-blind placebo-controlled trials, repeated intermittent exposure (60–90 min) to normobaric hypoxia (up to 4500 m) or continuous exposure to 3000 m during 8 h of sleep for 7 consecutive days failed to reduce the incidence of AMS at altitudes of 4300–4559 m.

Clearly, a flexible itinerary that permits additional rest days will be helpful. Sojourners to high-altitude locations must be aware of the symptoms of altitude illness and should be encouraged not to ascend further if these symptoms develop. Any hint of HAPE (see below) or HACE mandates descent. Finally, proper hydration (but not overhydration) in high-altitude trekking and climbing, aimed at countering fluid loss due to hyperventilation and sweating, may also play a role in avoiding AMS. Pharmacologic prophylaxis at the time of travel to high altitudes is warranted for people with a history of AMS or when a graded ascent and acclimation are not possible—e.g., when rapid ascent is necessary for rescue purposes or when flight to a high-altitude location is required. Acetazolamide is the drug of choice for AMS prevention. It inhibits renal carbonic anhydrase, causing a prompt bicarbonate diuresis that leads to metabolic acidosis and hyperventilation. Acetazolamide (125–250 mg twice a day), administered for 1 day before ascent and continued for 2 or 3 days, is effective. Higher doses are not required. A meta-analysis limited to randomized controlled trials revealed that 125 mg of acetazolamide twice daily was effective in the prevention of AMS, with a relative-risk reduction of $\sim 48\%$ from values obtained with placebo. Paresthesia and a tingling sensation are common side effects of acetazolamide. This drug is a nonantibiotic sulfonamide that has low-level cross-reactivity with sulfa antibiotics; as a result, severe reactions are rare. Dexamethasone (8 mg/d in divided doses) is also effective. A large-scale, randomized, double-blind,

placebo-controlled trial in partially acclimated trekkers has clearly shown that *Ginkgo biloba* is ineffective in the prevention of AMS. Recently, ibuprofen (600 mg three times daily) was shown to be beneficial in the prevention of AMS, but more definitive studies and proper gastrointestinal-bleeding risk assessment need to be conducted before ibuprofen can be routinely recommended for AMS prevention. Many drugs, including spironolactone, medroxyprogesterone, magnesium, calcium channel blockers, and antacids, confer no benefit in the prevention of AMS. Similarly, no efficacy studies are available for coca leaves (a weak form of cocaine), which are offered to high-altitude travelers in the Andes, or for *soroche* pills, which contain aspirin, caffeine, and acetaminophen and are sold over the counter in Bolivia and Peru. Finally, a word of caution applies in the pharmacologic prevention of altitude illness. A fast-growing population of climbers in pursuit of a summit are using prophylactic drugs such as glucocorticoids in an attempt to improve their performance; the outcome can be tragic because of potentially severe side effects of these drugs.

For the treatment of mild AMS, rest alone with analgesic use may be adequate. Descent and the use of acetazolamide and (if available) oxygen are sufficient to treat most cases of moderate AMS. Even a minor descent (400–500 m) may be adequate for symptom relief. For moderate AMS or early HACE, dexamethasone (4 mg orally or parenterally) is highly effective. For HACE, immediate descent is mandatory. When descent is not possible because of poor weather conditions or darkness, a simulation of descent in a portable hyperbaric chamber is effective and, like dexamethasone administration, “buys time.” Thus, in certain high-altitude locations (e.g., remote pilgrimage sites), the decision to bring along the light-weight hyperbaric chamber may prove life-saving. Like nifedipine, phosphodiesterase-5 inhibitors have no role in the treatment of AMS or HACE.

HIGH-ALTITUDE PULMONARY EDEMA

Risk Factors and Manifestations Unlike HACE (a neurologic disorder), HAPE is primarily a pulmonary problem and therefore is not necessarily preceded by AMS. HAPE develops within 2–4 days after arrival at high altitude; it rarely occurs after more than 4 or 5 days at the same altitude, probably because of remodeling and adaptation that render the pulmonary vasculature less susceptible to the effects of hypoxia. A rapid rate of ascent, a history of HAPE, respiratory tract infections, and cold environmental temperatures are risk factors. Men are more susceptible than women. People with abnormalities of the cardiopulmonary circulation leading to pulmonary hypertension—e.g., large patent foramen ovale, mitral stenosis, primary pulmonary hypertension, and unilateral absence of the pulmonary artery—are at increased risk of HAPE, even at moderate altitudes. For example, patent foramen ovale is four times more common among HAPE-susceptible individuals than in the general population. Echocardiography is recommended when HAPE develops at relatively low altitudes (<3000 m) and whenever cardiopulmonary abnormalities predisposing to HAPE are suspected.

The initial manifestation of HAPE may be a reduction in exercise tolerance greater than that expected at the given altitude. Although a dry, persistent cough may presage HAPE and may be followed by the production of blood-tinged sputum, cough in the mountains is almost universal and the mechanism is poorly understood. Tachypnea and tachycardia, even at rest, are important markers as illness progresses. Crackles may be heard on auscultation but are not diagnostic. HAPE may be accompanied by signs of HACE. Patchy or localized opacities (Fig. 476e-2) or streaky interstitial edema may be noted on chest radiography. In the past, HAPE was mistaken for pneumonia due to the cold or for heart failure due to hypoxia and exertion. Kerley B lines or a bat-wing appearance are not seen on radiography. Electrocardiography may reveal right ventricular strain or even hypertrophy. Hypoxemia and respiratory alkalosis are consistently present unless the patient is taking acetazolamide, in which case metabolic acidosis may supervene. Assessment of arterial blood gases is not necessary in the evaluation of HAPE; an oxygen saturation reading with a pulse oximeter is generally adequate. The existence of a subclinical form of HAPE has been suggested by an increased alveolar-arterial oxygen gradient in Everest

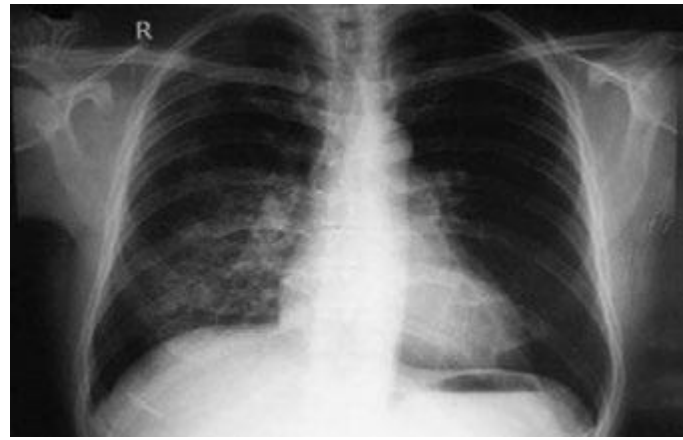


FIGURE 476e-2 Chest radiograph of a patient with high-altitude pulmonary edema shows opacity in the right middle and lower zones simulating pneumonic consolidation. The opacity cleared almost completely in 2 days with descent and supplemental oxygen.

climbers near the summit, but hard evidence correlating this abnormality with the development of clinically relevant HAPE is lacking.

Pathophysiology HAPE is a noncardiogenic pulmonary edema characterized by patchy pulmonary vasoconstriction that leads to overperfusion in some areas. This abnormality leads in turn to increased pulmonary capillary pressure (>18 mmHg) and capillary “stress” failure. The exact mechanism for the vasoconstriction is unknown. Endothelial dysfunction due to hypoxia may play a role by impairing the release of nitric oxide, an endothelium-derived vasodilator. At high altitude, HAPE-prone persons have reduced levels of exhaled nitric oxide. The effectiveness of phosphodiesterase-5 inhibitors in alleviating altitude-induced pulmonary hypertension, decreased exercise tolerance, and hypoxemia supports the role of nitric oxide in the pathogenesis of HAPE. One study demonstrated that prophylactic use of tadalafil, a phosphodiesterase-5 inhibitor, decreases the risk of HAPE by 65%. In contrast, the endothelium also synthesizes endothelin-1, a potent vasoconstrictor whose concentrations are higher than average in HAPE-prone mountaineers. Bosentan, an endothelin receptor antagonist, attenuates hypoxia-induced pulmonary hypertension, but further field studies with this drug are necessary.

Exercise and cold lead to increased pulmonary intravascular pressure and may predispose to HAPE. In addition, hypoxia-triggered increases in sympathetic drive may lead to pulmonary vasoconstriction and extravasation into the alveoli from the pulmonary capillaries. Consistent with this concept, phentolamine, which elicits α -adrenergic blockade, improves hemodynamics and oxygenation in HAPE more than do other vasodilators. The study of tadalafil cited above also investigated dexamethasone in the prevention of HAPE. Surprisingly, dexamethasone reduced the incidence of HAPE by 78%—a greater decrease than with tadalafil. Besides possibly increasing the availability of endothelial nitric oxide, dexamethasone may have altered the excessive sympathetic activity associated with HAPE: the heart rate of participants in the dexamethasone arm of the study was significantly lowered. Finally, people susceptible to HAPE also display enhanced sympathetic activity during short-term hypoxic breathing at low altitudes.

Because many patients with HAPE have fever, peripheral leukocytosis, and an increased erythrocyte sedimentation rate, inflammation has been considered an etiologic factor in HAPE. However, strong evidence suggests that inflammation in HAPE is an epiphenomenon rather than the primary cause. Nevertheless, inflammatory processes (e.g., those elicited by viral respiratory tract infections) do predispose persons to HAPE—even those who are constitutionally resistant to its development.

Another proposed mechanism for HAPE is impaired transepithelial clearance of sodium and water from the alveoli. β -Adrenergic agonists

476e-4 upregulate the clearance of alveolar fluid in animal models. In a double-blind, randomized, placebo-controlled study of HAPE-susceptible mountaineers, prophylactic inhalation of the β -adrenergic agonist salmeterol reduced the incidence of HAPE by 50%. Other effects of β agonists may also contribute to the prevention of HAPE, but these findings are in keeping with the concept that alveolar fluid clearance may play a pathogenic role in this illness.

Prevention and Treatment (Table 476e-1) Allowing sufficient time for acclimation by ascending gradually (as discussed above for AMS and HACE) is the best way to prevent HAPE. Sustained-release nifedipine (30 mg), given once or twice daily, prevents HAPE in people who must ascend rapidly or who have a history of HAPE. Other drugs for the prevention of HAPE are listed in Table 476e-1 (footnote *e*). Although dexamethasone is listed for prevention, its adverse-effect profile requires close monitoring. Acetazolamide has been shown to blunt hypoxic pulmonary vasoconstriction in animal models, and this observation warrants further study in HAPE prevention. However, one large study failed to show a decrease in pulmonary vasoconstriction in partially acclimated individuals given acetazolamide.

Early recognition is paramount in the treatment of HAPE, especially when it is not preceded by the AMS symptoms of headache and nausea. Fatigue and dyspnea at rest may be the only initial manifestations. Descent and the use of supplementary oxygen (aimed at bringing oxygen saturation to >90%) are the most effective therapeutic interventions. Exertion should be kept to a minimum, and the patient should be kept warm. Hyperbaric therapy in a portable altitude chamber may be used if descent is not possible and oxygen is not available. Oral sustained-release nifedipine (30 mg once or twice daily) can be used as adjunctive therapy. Inhaled β agonists, which are safe and convenient to carry, are useful in the prevention of HAPE and may be effective in its treatment, although no trials have yet been carried out. Inhaled nitric oxide and expiratory positive airway pressure may also be useful therapeutic measures but may not be available in high-altitude settings. No studies have investigated phosphodiesterase-5 inhibitors in the treatment of HAPE, but reports have described their use in clinical practice. The mainstays of treatment remain descent and (if available) oxygen.

In AMS, if symptoms abate (with or without acetazolamide), the patient may reascend gradually to a higher altitude. Unlike that in acute respiratory distress syndrome (another noncardiogenic pulmonary edema), the architecture of the lung in HAPE is usually well preserved, with rapid reversibility of abnormalities (Fig. 476e-2). This fact has allowed some people with HAPE to reascend slowly after a few days of descent and rest. In HACE, reascend after a few days may not be advisable during the same trip.

OTHER HIGH-ALTITUDE PROBLEMS

Sleep Impairment The mechanisms underlying sleep problems, which are among the most common adverse reactions to high altitude, include increased periodic breathing; changes in sleep architecture, with increased time in lighter sleep stages; and changes in rapid eye movement sleep. Sojourners should be reassured that sleep quality improves with acclimation. In cases where drugs do need to be used, acetazolamide (125 mg before bedtime) is especially useful because this agent decreases hypoxemic episodes and alleviates sleeping disruptions caused by excessive periodic breathing. Whether combining acetazolamide with temazepam or zolpidem is more effective than administering acetazolamide alone is unknown. In combinations, the doses of temazepam and zolpidem should not be increased by >10 mg at high altitudes. Limited evidence suggests that diazepam causes hypoventilation at high altitudes and therefore is contraindicated. For trekkers with obstructive sleep apnea who are using a continuous positive airway pressure (CPAP) machine, the addition of acetazolamide, which will decrease centrally mediated sleep apnea, may be helpful. There is evidence to show that obstructive sleep apnea at high altitude may decrease and “convert” to central sleep apnea.

Gastrointestinal Issues High-altitude exposure may be associated with increased gastric and duodenal bleeding, but further studies

are required to determine whether there is a causal effect. Because of decreased atmospheric pressure and consequent intestinal gas expansion at high altitudes, many sojourners experience abdominal bloating and distension as well as excessive flatus expulsion. In the absence of diarrhea, these phenomena are normal, if sometimes uncomfortable. Accompanying diarrhea, however, may indicate the involvement of bacteria or *Giardia* parasites, which are common at many high-altitude locations in the developing world. Prompt treatment with fluids and empirical antibiotics may be required to combat dehydration in the mountains. Finally, hemorrhoids are common on high-altitude treks; treatment includes hot soaks, application of hydrocortisone ointment, and measures to avoid constipation.

High-Altitude Cough High-altitude cough can be debilitating and is sometimes severe enough to cause rib fracture, especially at >5000 m. The etiology is probably multifactorial. Although high-altitude cough has been attributed to inspiration of cold dry air, this explanation appears not to be sufficient by itself; in long-duration studies in hypobaric chambers, cough has occurred despite controlled temperature and humidity. The implication is that hypoxia also plays a role. Exercise can precipitate cough at high altitudes, possibly because of water loss from the respiratory tract. Long-acting β agonists and glucocorticoids prevent bronchoconstriction that otherwise may be brought on by cold and exercise. In general, infection does not seem to be a common etiology. Anecdotal reports have described the efficacy of an inhaled combination of fluticasone and salmeterol in the treatment of high-altitude cough; however, a placebo-controlled, randomized trial failed to support this beneficial effect. Furthermore, anecdotal evidence supports the utility of the proton pump inhibitor omeprazole in preventing gastroesophageal reflux in some trekkers and climbers. In most situations, cough resolves upon descent.

High-Altitude Neurologic Events Unrelated to “Altitude Illness” Transient ischemic attacks (TIAs) and strokes have been well described in high-altitude sojourners outside the setting of altitude sickness. However, these descriptions are not based on cause (hypoxia) and effect. In general, symptoms of AMS present gradually, whereas many of these neurologic events happen suddenly. The population that suffers strokes and TIAs at sea level is generally an older age group with other risk factors, whereas those so afflicted at high altitudes are generally younger and probably have fewer risk factors for atherosclerotic vascular disease. Other mechanisms (e.g., migraine, vasospasm, focal edema, hypocapnic vasoconstriction, hypoxia in the watershed zones of minimal cerebral blood flow, or cardiac right-to-left shunt) may be operative in TIAs and strokes at high altitude.

Subarachnoid hemorrhage, transient global amnesia, delirium, and cranial nerve palsies (e.g., lateral rectus palsy) occurring at high altitudes but outside the setting of altitude sickness have also been well described. Syncope is common at moderately high altitudes, generally occurs shortly after ascent, usually resolves without descent, and appears to be a vasovagal event related to hypoxemia. Seizures occur rarely with HACE, but hypoxemia and hypocapnia, which are prevalent at high altitudes, are well-known triggers that may contribute to new or breakthrough seizures in predisposed individuals. Nevertheless, the consensus among experts is that sojourners with well-controlled seizure disorders can ascend to high altitudes. Ophthalmologic problems, such as cortical blindness, amaurosis fugax, and retinal hemorrhage with macular involvement and compromised vision, are well recognized. Visual problems from previous refractive surgery and blurred monocular vision—due either to the use of a transdermal scopolamine patch (touching the eye after touching the patch) or to dry-eye syndrome—may also occur in the field at high altitudes and may be confused with neurologic conditions. Finally, persons with hypercoagulable conditions (e.g., antiphospholipid syndrome, protein C deficiency) who are asymptomatic at sea level may experience cerebral venous thrombosis (possibly due to the enhanced blood viscosity triggered by polycythemia and dehydration) at high altitudes. Proper history taking, examination, and prompt investigations where possible will help define these conditions as entities separate from altitude sickness. Administration of oxygen (where available)

and prompt descent are the cornerstones of treatment of most of these neurologic conditions.

Psychological/Psychiatric Problems Delirium characterized by a sudden change in mental status, a short attention span, disorganized thinking, and an agitated state during the period of confusion has been well described in mountain climbers and trekkers without a prior history. In addition, anxiety attacks, often triggered at night by excessive periodic breathing, are well documented. The contribution of hypoxia to these conditions is unknown. Expedition medical kits need to include antipsychotic injectable drugs to control psychosis in patients in remote high-altitude locations.

PREEXISTING MEDICAL ISSUES

Because travel to high altitudes is increasingly popular, common conditions such as hypertension, coronary artery disease, and diabetes are more frequently encountered among high-altitude sojourners. This situation is of particular concern for the thousands of elderly pilgrims with medical problems who visit high-altitude sacred areas (e.g., in the Himalayas) each year. In recent years, high-altitude travel has attracted intrepid trekkers who are taking immunosuppressive medications (e.g., kidney transplant recipients or patients undergoing chemotherapy). Recommended vaccinations and other precautions (e.g., hand washing) may be especially important for this group. Although most of these medical conditions do not appear to influence susceptibility to altitude illness, they may be exacerbated by ascent to altitude, exertion in cold conditions, and hypoxemia. Advice regarding the advisability of high-altitude travel and the impact of high-altitude hypoxia on these preexisting conditions is becoming increasingly relevant, but there are no evidence-based guidelines. In addition, recommendations made for relatively low altitudes (~3000 m) may not hold true for higher altitudes (>4000 m), where hypoxic stress is greater. Personal risks and benefits must be clearly thought through before ascent.

Hypertension At high altitudes, enhanced sympathetic activity may lead to a transient rise in blood pressure. Occasionally, nonhypertensive, healthy, asymptomatic trekkers have pathologically high blood pressure at high altitude that rapidly normalizes without medicines on descent. Sojourners should continue to take their antihypertensive medications at high altitudes. Hypertensive patients are not more likely than others to develop altitude illness. Because the probable mechanism of high-altitude hypertension is α -adrenergic activity, anti- α -adrenergic drugs like prazosin have been suggested for symptomatic patients and those with labile hypertension. It is best to start taking the drug several weeks before the trip and to carry a sphygmomanometer if a trekker has labile hypertension. Sustained-release nifedipine may also be useful. For a common problem like hypertension, there is clearly inadequate knowledge on which to base appropriate recommendations.

Coronary Artery Disease Myocardial oxygen demand and maximal heart rate are reduced at high altitudes because the VO_2 max (maximal oxygen consumption) decreases with increasing altitude. This effect may explain why signs of cardiac ischemia or dysfunction usually are not seen in healthy persons at high altitudes. Asymptomatic, fit individuals with no risk factors need not undergo any tests for coronary artery disease before ascent. For persons with ischemic heart disease, previous myocardial infarction, angioplasty, and/or bypass surgery, an exercise treadmill test is indicated. A strongly positive treadmill test is a contraindication for high-altitude trips. Patients with poorly controlled arrhythmias should avoid high-altitude travel, but patients with arrhythmias that are well controlled with antiarrhythmic medications do not seem to be at increased risk. Sudden cardiac deaths are not noted with a greater frequency in the Alps than at lower altitudes; although sudden cardiac deaths are encountered every trekking season in the higher Himalayan range, accurate documentation is lacking.

Asthma Although cold air and exercise may provoke acute bronchoconstriction, asthmatic patients usually have fewer problems at high than at low altitudes, possibly because of decreased allergen levels and increased circulating catecholamine levels. Nevertheless, asthmatic individuals should carry all their medications, including oral

glucocorticoids, with proper instructions for use in case of an exacerbation. Severely asthmatic persons should be cautioned against ascending to high altitudes.

Pregnancy In general, low-risk pregnant women ascending to 3000 m are not at special risk except for the relative unavailability of medical care in many high-altitude locations, especially in developing countries. Despite the lack of firm data on this point, venturing higher than 3000 m to altitudes at which oxygen saturation drops steeply seems inadvisable for pregnant women.

Obesity Although living at a high altitude has been suggested as a means of controlling obesity, obesity has also been reported to be a risk factor for AMS, probably because nocturnal hypoxemia is more pronounced in obese individuals. Hypoxemia may also lead to greater pulmonary hypertension, thus possibly predisposing the trekker to HAPE.

Sickle Cell Disease High altitude is one of the rare environmental exposures that occasionally provokes a crisis in persons with the sickle cell trait. Even when traversing mountain passes as low as 2500 m, people with sickle cell disease have been known to have a vasoocclusive crisis. Sickle cell disease needs to be considered when persons traveling to high altitudes become unwell and develop left-upper-quadrant pain. Patients with known sickle cell disease who need to travel to high altitudes should use supplemental oxygen and travel with caution.

Diabetes Mellitus Trekking at high altitudes may enhance sugar uptake. Thus, high-altitude travel may not pose problems for persons with diabetes that is well controlled with oral hypoglycemic agents. An eye examination before travel may be useful. Patients taking insulin may require lower doses on trekking/climbing days than on rest days. Because of these variations, diabetic patients need to carry a reliable glucometer and use fast-acting insulin. Ready access to sweets is also essential. It is important for companions of diabetic trekkers to be fully aware of potential problems like hypoglycemia.

Chronic Lung Disease Depending on disease severity and access to medical care, preexisting lung disease may not always preclude high-altitude travel. A proper pretravel evaluation must be conducted. Supplemental oxygen may be required if the predicted PaO_2 for the altitude is <50–55 mmHg. Preexisting pulmonary hypertension may also need to be assessed in these patients. If the result is positive, patients should be discouraged from ascending to high altitudes; if such travel is necessary, treatment with sustained-release nifedipine (20 mg twice a day) should be considered. Small-scale studies have revealed that when patients with bullous disease reach ~5000 m, bullous expansion and pneumothorax are not noted. Compared with information on chronic obstructive pulmonary disease, fewer data exist about the safety of travel to high altitude for people with pulmonary fibrosis, but acute exacerbation of pulmonary fibrosis has been seen at high altitude. A handheld pulse oximeter can be useful to check for oxygen saturation.

Chronic Kidney Disease Patients with chronic kidney disease can tolerate short-term stays at high altitudes, but theoretical concern persists about progression to end-stage renal disease. Acetazolamide, the drug most commonly used for altitude sickness, should be avoided by anyone with preexisting metabolic acidosis, which can be exacerbated by this drug. In addition, the acetazolamide dosage should be adjusted when the glomerular filtration rate falls below 50 mL/min, and the drug should not be used at all if this value falls below 10 mL/min.

CHRONIC MOUNTAIN SICKNESS AND HIGH-ALTITUDE PULMONARY HYPERTENSION

Chronic mountain sickness (*Monge's disease*) is a disease of long-term residents of altitudes above 2500 m that is characterized by excessive erythrocytosis with moderate to severe pulmonary hypertension leading to cor pulmonale. This condition was originally described in South America and has also been documented in Colorado and in the Han Chinese population in Tibet. Migration to a low altitude results in the resolution of chronic mountain illness. Venesection and acetazolamide are helpful.

476e-6 High-altitude pulmonary hypertension is also a subacute disease of long-term high-altitude residents. Unlike Monge's disease, this syndrome is characterized primarily by pulmonary hypertension (not erythrocytosis) leading to heart failure. Indian soldiers living at extreme altitudes for prolonged periods and Han Chinese infants born

in Tibet have presented with the adult and infantile forms, respectively. High-altitude pulmonary hypertension bears a striking pathophysiologic resemblance to brisket disease in cattle. Descent to a lower altitude is curative.

477e Hyperbaric and Diving Medicine

Michael H. Bennett, Simon J. Mitchell

WHAT IS HYPERBARIC AND DIVING MEDICINE?

Hyperbaric medicine is the treatment of health disorders using whole-body exposure to pressures greater than 101.3 kPa (1 atmosphere or 760 mmHg). In practice, this almost always means the administration of *hyperbaric oxygen therapy* (HBO₂T). The Undersea and Hyperbaric Medical Society (UHMS) defines HBO₂T as: “a treatment in which a patient breathes 100% oxygen ... while inside a treatment chamber at a pressure higher than sea level pressure (i.e., >1 atmosphere absolute or ATA).” The treatment chamber is an airtight vessel variously called a hyperbaric chamber, recompression chamber, or decompression chamber, depending on the clinical and historical context. Such chambers may be capable of compressing a single patient (a monoplace chamber) or multiple patients and attendants as required (a multiplace chamber) (Figs. 477e-1 and 477e-2). Historically, these compression chambers were first used for the treatment of divers and compressed air workers suffering decompression sickness (DCS; “the bends”). Although the prevention and treatment of disorders arising after decompression in diving, aviation, and space flight has developed into a specialized field of its own, it remains closely linked to the broader practice of hyperbaric medicine.

Despite an increased understanding of mechanisms and an improving evidence basis, hyperbaric medicine has struggled to achieve widespread recognition as a “legitimate” therapeutic measure. There are several contributing factors, but high among them are a poor grounding in general oxygen physiology and oxygen therapy at medical schools and a continuing tradition of charlatans advocating hyperbaric therapy (often using air) as a panacea. Funding for both basic and clinical research has been difficult in an environment where the pharmacologic agent under study is abundant, cheap, and unpatentable. Recently, however, there are signs of an improved appreciation of the potential importance of HBO₂T with significant National Institutes of Health (NIH) funding for mechanisms research and from the U.S. military for clinical investigation.

MECHANISMS OF HYPERBARIC OXYGEN

Increased hydrostatic pressure will reduce the volume of any bubbles present within the body (see “Diving Medicine”), and this is partly responsible for the success of prompt recompression in DCS and



FIGURE 477e-1 A monoplace chamber. (Prince of Wales Hospital, Sydney.)



FIGURE 477e-2 A chamber designed to treat multiple patients. (Karolinska University Hospital.)

arterial gas embolism. Supplemental oxygen breathing has a dose-dependent effect on oxygen transport, ranging from improvement in hemoglobin oxygen saturation when a few liters per minute are delivered by simple mask at 101.3 kPa (1 ATA) to raising the dissolved plasma oxygen sufficiently to sustain life without the need for hemoglobin at all when 100% oxygen is breathed at 303.9 kPa (3 ATA). Most HBO₂T regimens involve oxygen breathing at between 202.6 kPa and 283.6 kPa (2 and 2.8 ATA), and the resultant increase in arterial oxygen tensions to >133.3 kPa (1000 mmHg) has widespread physiologic and pharmacologic consequences (Fig. 477e-3).

One direct consequence of such high intravascular tension is to increase greatly the effective capillary-tissue diffusion distance for oxygen such that oxygen-dependent cellular processes can resume in hypoxic tissues. Important as this may be, the mechanism of action is not limited to this restoration of oxygenation in hypoxic tissue. Indeed, there are pharmacologic effects that are profound and long-lasting. Although removal from the hyperbaric chamber results in a rapid return of poorly vascularized tissues to their hypoxic state, even a single dose of HBO₂T produces changes in fibroblast, leukocyte, and angiogenic functions and antioxidant defenses that persist many hours after oxygen tensions are returned to pretreatment levels.

It is widely accepted that oxygen in high doses produces adverse effects due to the production of reactive oxygen species (ROS) such as superoxide (O₂⁻) and hydrogen peroxide (H₂O₂). It has become increasingly clear over the last decade that both ROS and reactive nitrogen species (RNS) such as nitric oxide (NO) participate in a wide range of intracellular signaling pathways involved in the production of a range of cytokines, growth factors, and other inflammatory and repair modulators. Such mechanisms are complex and at times apparently paradoxical. For example, when used to treat chronic hypoxic wounds, HBO₂T has been shown to enhance the clearance of cellular debris and bacteria by providing the substrate for macrophage phagocytosis; stimulate growth factor synthesis by increased production and stabilization of hypoxia-inducible factor 1 (HIF-1); inhibit leukocyte activation and adherence to damaged endothelium; and mobilize CD34+ pluripotent vasculogenic progenitor cells from the bone marrow. The interactions between these mechanisms remain a very active field of investigation. One exciting development is the concept of *hyperoxic preconditioning* in which a short exposure to HBO₂ can induce tissue protection against future hypoxic/ischemic insult, most likely through an inhibition of mitochondrial permeability transition pore (MPTP) opening and the release of cytochrome c. By targeting these mechanisms of cell death during reperfusion events, HBO₂ has potential applications in a variety of settings including organ transplantation. One randomized clinical trial suggested that HBO₂T prior to coronary artery bypass grafting reduces biochemical markers of ischemic stress and improves neurocognitive outcomes.

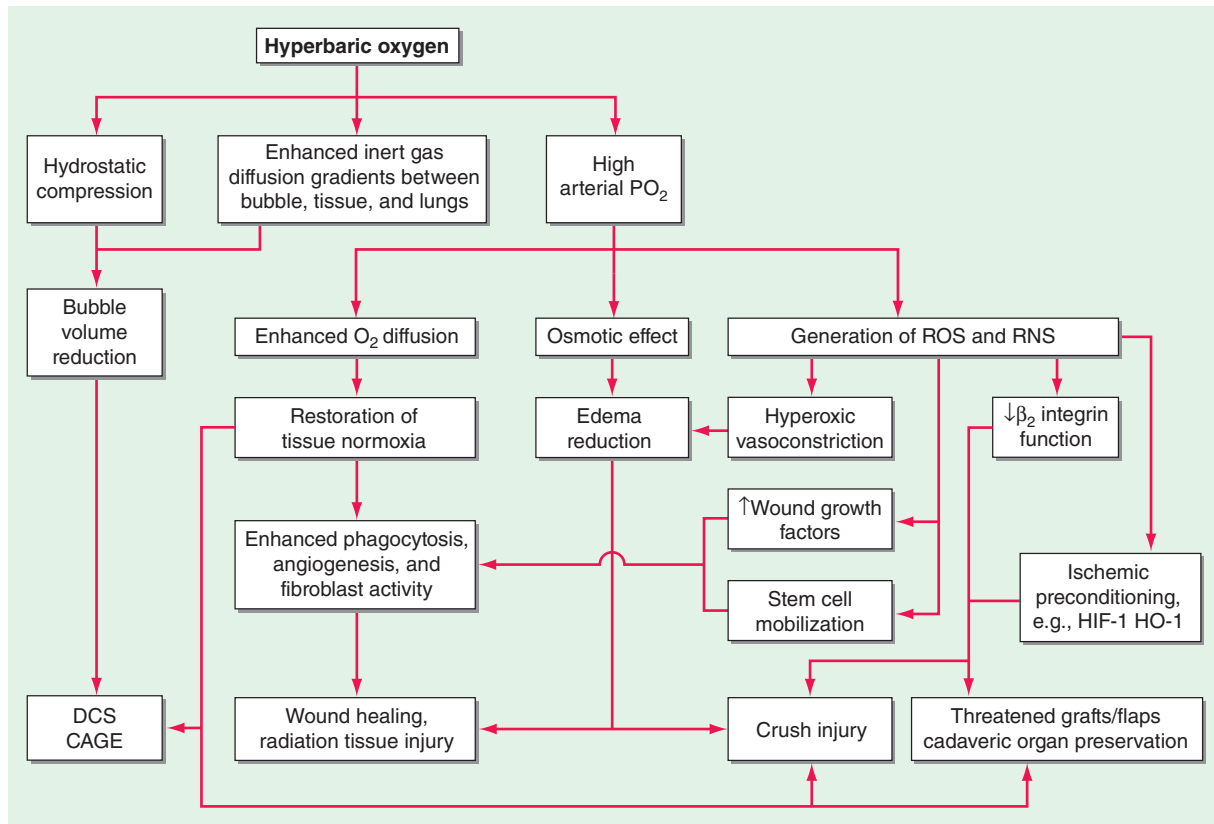


FIGURE 477e-3 Mechanisms of action of hyperbaric oxygen. There are many consequences of compression and oxygen breathing. The cell-signaling effects of HBO₂T are the least understood but potentially most important. Examples of indications for use are shown in the shaded boxes. CAGE, cerebral arterial gas embolism; DCS, decompression sickness; HIF-1, hypoxia inducible factor-1; HO-1, hemoxygenase 1; RNS, reactive nitrogen species; ROS, reactive oxygen species.

ADVERSE EFFECTS OF THERAPY

HBO₂T is generally well tolerated and safe in clinical practice. Adverse effects are associated with both alterations in pressure (barotrauma) and the administration of oxygen.

BAROTRAUMA

Barotrauma occurs when any noncompliant gas-filled space within the body does not equalize with environmental pressure during compression or decompression. About 10% of patients complain of some difficulty equalizing middle-ear pressure early in compression, and although most of these problems are minor and can be overcome with training, 2–5% of conscious patients require middle-ear ventilation tubes or formal grommets across the tympanic membrane. Unconscious patients cannot equalize and should have middle-ear ventilation tubes placed prior to compression if possible. Other less common sites for barotrauma of compression include the respiratory sinuses and dental caries. The lungs are potentially vulnerable to barotrauma of decompression as described below in the section on diving medicine, but the decompression following HBO₂T is so slow that pulmonary gas trapping is extremely rare in the absence of an undrained pneumothorax or lesions such as bullae.

OXYGEN TOXICITY

The practical limit to the dose of oxygen, either in a single treatment session or in a series of daily sessions, is oxygen toxicity. The most common acute manifestation is a seizure, often preceded by anxiety and agitation, during which time a switch from oxygen to air breathing may avoid the convulsion. Hyperoxic seizures are typically generalized tonic-clonic seizures followed by a variable postictal period. The cause is an overwhelming of the antioxidant defense systems within the brain. Although clearly dose-dependent, onset is very variable both between individuals and within the same individual on different days. In routine clinical hyperbaric practice, the incidence is about 1:1500 to 1:2000 compressions.

Chronic oxygen poisoning most commonly manifests as myopic shift. This is due to alterations in the refractive index of the lens following oxidative damage that reduces the solubility of lenticular proteins in a process similar to that associated with senescent cataract formation. Up to 75% of patients show deterioration in visual acuity after a course of 30 treatments at 202.6 kPa (2 ATA). Although most return to pretreatment values 6–12 weeks after cessation of treatment, a small proportion do not recover. A more rapid maturation of preexisting cataracts has occasionally been associated with HBO₂T. Although a theoretical problem, the development of pulmonary oxygen toxicity over time does not seem to be problematic in practice—probably due to the intermittent nature of the exposure.

CONTRAINDICATIONS TO HYPERBARIC OXYGEN

There are few absolute contraindications to HBO₂T. The most commonly encountered is an untreated pneumothorax. A pneumothorax may expand rapidly on decompression and come under tension. Prior to any compression, patients with a pneumothorax should have a patent chest drain in place. The presence of other obvious risk factors for pulmonary gas trapping such as bullae should trigger a very cautious analysis of the risks of treatment versus benefit. Prior bleomycin treatment deserves special mention because of its association with a partially dose-dependent pneumonitis in about 20% of people. These individuals appear to be at particular risk for rapid deterioration of ventilatory function following exposure to high oxygen tensions. The relationship between distant bleomycin exposure and subsequent risk of pulmonary oxygen toxicity is uncertain, however late pulmonary fibrosis is a potential complication of bleomycin, and any patient with a history of receiving this drug should be carefully counseled prior to exposure to HBO₂T. For those recently exposed to doses above 300,000 IU (200 mg) and whose course was complicated by a respiratory reaction to bleomycin, compression should be avoided except in a life-threatening situation.

INDICATIONS FOR HYPERBARIC OXYGEN

The appropriate indications for HBO₂T are controversial and evolving. Practitioners in this area are in an unusual position. Unlike most branches of medicine, hyperbaric physicians do not deal with a range of disorders within a defined organ system (e.g., cardiology), nor are they masters of a therapy specifically designed for a single category of disorders. Inevitably, the encroachment of hyperbaric physicians into other medical fields generates suspicion from specialist practitioners in those fields. At the same time this relatively benign therapy, the prescription and delivery of which requires no medical license in most jurisdictions (including the United States), attracts both charlatans and well-motivated proselytizers who tout the benefits of oxygen for a plethora of chronic incurable diseases. This battle on two fronts has meant that mainstream hyperbaric physicians have been particularly careful to claim effectiveness only for those conditions where there is a reasonable body of supporting evidence.

In 1977, the UHMS systematically examined claims for the use of HBO₂T in more than 100 disorders and found sufficient evidence to support routine use in only 12. The Hyperbaric Oxygen Therapy Committee of that organization has continued to update this list periodically with an increasingly formalized system of appraisal for new indications and emerging evidence (Table 477e-1). Around the world, other relevant medical organizations have generally taken a similar approach, although indications vary considerably—particularly those recommended by hyperbaric medical societies in Russia and China where HBO₂T has gained much wider support than in the United States, Europe, and Australasia. Recently, several Cochrane reviews have examined the randomized trial evidence for many putative indications, including attempts to examine the cost-effectiveness of HBO₂T. Table 477e-2 is a synthesis of these two approaches and lists the estimated cost of attaining health outcomes with the use of HBO₂T. Any savings associated with alternative treatment strategies avoided as a result of HBO₂T are not accounted for in these estimates (e.g., the avoidance of lower leg amputation in diabetic foot ulcers). Following are short reviews of three important indications currently accepted by the UHMS.

LATE RADIATION TISSUE INJURY

Radiotherapy is a well-established treatment for suitable malignancies. In the United States alone, approximately 300,000 individuals annually will become long-term survivors of cancer treated by irradiation. Serious radiation-related complications developing months or years after treatment (late radiation tissue injury [LRTI]) will significantly affect between 5 and 15% of those long-term survivors, although inci-

dence varies widely with dose, age, and site. LRTI is most common in the head and neck, chest wall, breast, and pelvis.

Pathology and Clinical Course With time, tissues undergo a progressive deterioration characterized by a reduction in the density of small blood vessels (reduced vascularity) and the replacement of normal tissue with dense fibrous tissue (fibrosis). An alternative model of pathogenesis suggests that rather than a primary hypoxia, the principle trigger is an overexpression of inflammatory cytokines that promote fibrosis, probably through oxidative stress and mitochondrial dysfunction, and a secondary tissue hypoxia. Ultimately, and often triggered by a further physical insult such as surgery or infection, there may be insufficient oxygen to sustain normal function, and the tissue becomes necrotic (radiation necrosis). LRTI may be life-threatening and significantly reduce quality of life. Historically, the management of these injuries has been unsatisfactory. Conservative treatment is usually restricted to symptom management, whereas definitive treatment traditionally entails surgery to remove the affected part and extensive repair. Surgical intervention in an irradiated field is often disfiguring and associated with an increased incidence of delayed healing, breakdown of a surgical wound, or infection. HBO₂T may act by several mechanisms to improve this situation, including edema reduction, vasculogenesis, and enhancement of macrophage activity (Fig. 477e-3). The intermittent application of HBO₂ is the only intervention shown to increase the microvascular density in irradiated tissue.

Clinical Evidence The typical course of HBO₂T consists of 30 once-daily compressions to 202.6–243.1 kPa (2–2.4 ATA) for 1.5–2 h each session, often bracketed around surgical intervention if required. Although HBO₂T has been used for LRTI since at least 1975, most clinical studies have been limited to small case series or individual case reports. In a review, Feldmeier and Hampson located 71 such reports involving a total of 1193 patients across eight different tissues. There were clinically significant improvements in the majority of patients, and only 7 of 71 reports indicated a generally poor response to HBO₂T. A Cochrane systematic review with meta-analysis included six randomized trials published since 1985 and drew the following conclusions (see Table 477e-2 for numbers needed to treat): HBO₂T improves healing in radiation proctitis (relative risk [RR] of healing with HBO₂T 2.7; 95% confidence interval [CI] 1.2–6) and after hemimandibulectomy and reconstruction of the mandible (RR 1.4; 95% CI 1.1–1.8); HBO₂T improves the probability of achieving mucosal coverage (RR 1.4; 95% CI 1.2–1.6) and the restoration of bony continuity with osteoradionecrosis (ORN) (RR 1.4; 95% CI 1.1–1.8); HBO₂T prevents the development of ORN following tooth extraction from a radiation field (RR 1.4; 95% CI 1.08–1.7) and reduces the risk of wound dehiscence following grafts and flaps in the head and neck (RR 4.2; 95% CI 1.1–16.8). Conversely, there was no evidence of benefit in established radiation brachial plexus lesions or brain injury.

SELECTED PROBLEM WOUNDS

A problem wound is any cutaneous ulceration that requires a prolonged time to heal, does not heal, or recurs. In general, wounds referred to hyperbaric facilities are those where sustained attempts to heal by other means have failed. Problem wounds are common and constitute a significant health problem. It has been estimated that 1% of the population of industrialized countries will experience a leg ulcer at some time. The global cost of chronic wound care may be as high as U.S. \$25 billion per year.

Pathology and Clinical Course By definition, chronic wounds are indolent or progressive and resistant to the wide array of treatments applied. Although there are many contributing factors, most commonly these wounds arise in association with one or more comorbidities such as diabetes, peripheral venous or arterial disease, or prolonged pressure (decubitus ulcers). First-line treatments are aimed at correction of the underlying pathology (e.g., vascular reconstruction, compression bandaging, or normalization of blood glucose level), and HBO₂T is an adjunctive therapy to good general wound care practice to maximize the chance of healing.

TABLE 477e-1 CURRENT LIST OF INDICATIONS FOR HYPERBARIC OXYGEN THERAPY

1. Air or gas embolism (includes diving-related, iatrogenic, and accidental causes)
2. Carbon monoxide poisoning (including poisoning complicated by cyanide poisoning)
3. Clostridial myositis and myonecrosis (gas gangrene)
4. Crush injury, compartment syndrome, and acute traumatic ischemias
5. Decompression sickness
6. Arterial insufficiency
 - Central retinal artery occlusion
 - Enhancement of healing in selected problem wounds
7. Exceptional blood loss (where transfusion is refused or impossible)
8. Intracranial abscess
9. Necrotizing soft tissue infections (e.g., Fournier's gangrene)
10. Osteomyelitis (refractory to other therapy)
11. Delayed radiation injury (soft-tissue injury and bony necrosis)
12. Skin grafts and flaps (compromised)
13. Thermal burns
14. Sudden sensorineural hearing loss

Source: The Undersea and Hyperbaric Medical Society (2013).

TABLE 477e-2 SELECTED INDICATIONS FOR WHICH THERE IS PROMISING EFFICACY FOR THE APPLICATION OF HYPERBARIC OXYGEN THERAPY

Diagnosis	Outcome (number of sessions)	NNT 95% CI	Estimated Cost to Produce One Extra Favorable Outcome 95% CI (USD)	Comments and Recommendations
Radiation tissue injury	More information is required on the subset of disease severity, the affected tissue type that is most likely to benefit, and the time over which benefit may persist.			
	Resolved proctitis (30)	3	22,392	Large ongoing multicenter trial
		2–11	14,928–82,104	
	Healed mandible (30)	4	29,184	Based on one poorly reported study
		2–8	14,592–58,368	
	Mucosal cover in ORN (30)	3	29,888	Based on one poorly reported study
		2–4	14,592–29,184	
	Bony continuity in ORN (30)	4	29,184	Based on one poorly reported study
	2–8	14,592–58,368		
Prevention of ORN after dental extraction (30)	4	29,184	Based on a single study	
	2–13	14,592–94,848		
Prevention of dehiscence (30)	5	36,480	Based on one poorly reported study	
	3–8	21,888–58,368		
Chronic wounds	More information is required on the subset of disease severity or classification most likely to benefit, the time over which benefit may persist, and the most appropriate oxygen dose. Economic analysis is required.			
	Diabetic ulcer healed at 1 year (30)	2	14,928	Based on one small study, more research required
		1–5	7464–37,320	
	Diabetic ulcer, major amputation avoided (30)	4	29,856	Three small studies; outcome over a longer time period required
	3–11	22,392–82,104		
ISSNHL	No evidence of benefit more than 2 weeks after onset. More research is required to define the role (if any) of HBO₂T in routine therapy.			
	Improvement of 25% in hearing loss within 2 weeks of onset (15)	5	18,240	Some improvement in hearing, but functional significance unknown
		3–20	10,944–72,960	
Acute coronary syndrome	More information is required on the subset of disease severity and timing of therapy most likely to result in benefit. Given the potential of HBO₂T in modifying ischemia-reperfusion injury, attention should be given to the combination of HBO₂T and thrombolysis in early management and in the prevention of restenosis after stent placement.			
	Episode of MACE (5)	4	4864	Based on a single small study; more research required
		3–10	3648–12,160	
		6	7296	
	Incidence of significant dysrhythmia (5)	3–24	3648–29,184	Based on a single moderately powered study in the 1970s
Traumatic brain injury	Limited evidence that for acute injury HBO₂T reduces mortality but not functional morbidity. Routine use not yet justified.			
	Mortality (15)	7	34,104	Based on four heterogeneous studies
		4–22	19,488–58,464	
Enhancement of radiotherapy	There is some evidence that HBO₂T improves local tumor control, reduces mortality for cancers of the head and neck, and reduces the chance of local tumor recurrence in cancers of the head, neck, and uterine cervix.			
	Head and neck cancer:	5	14,592	Based on trials performed in the 1970s and 1980s. There may be some confounding by radiation fractionation schedule
	5-year mortality (12)	3–14	8755–40,858	
	Local recurrence 1 year (12)	5	14,592	May no longer be relevant to therapy
		4–8	11,674–23,347	
	Cancer of uterine cervix:	5	24,320	As above
	Local recurrence at 2 years (20)	4–8	19,456–38,912	
Decompression illness ^a	Reasonable evidence for reduced number of HBO₂T sessions but similar outcomes when NSAID added.			
	Reduction of HBO ₂ T treatment requirement by 1	5	N/R	Single appropriately powered randomized trial
	3–18			

^aTenoxicam used as an adjunct to recompression on oxygen.

Abbreviations: CI, confidence interval; HBO₂T, hyperbaric oxygen therapy; ISSNHL, idiopathic sudden sensorineural hearing loss; MACE, major adverse cardiac events; N/R, not remarkable; NNT, number needed to treat; NSAID, nonsteroidal anti-inflammatory drug; ORN, osteoradionecrosis; USD, U.S. dollars.

Source: M Bennett: The evidence-basis of diving and hyperbaric medicine—a synthesis of the high level evidence with meta-analysis. <http://unsworks.unsw.edu.au/vital/access/manager/Repository/unsworks:949>.

For most indolent wounds, hypoxia is a major contributor to failure to heal. Many guidelines to patient selection for HBO₂T include the interpretation of transcutaneous oxygen tensions around the wound while breathing air and oxygen at pressure (Fig. 477e-4). Wound healing is a complex and incompletely understood process. While it appears that in acute wounds healing is stimulated by the initial hypoxia, low pH, and high lactate concentrations found in freshly injured tissue, some elements of tissue repair are extremely oxygen dependent, for example, collagen elaboration and deposition by fibroblasts, and bacterial killing by macrophages. In this complicated interaction between wound hypoxia and peri-wound oxygenation, successful healing relies on adequate tissue oxygenation in the area surrounding the fresh wound. Certainly, wounds that lie in hypoxic tissue beds are those that most often display poor or absent healing. Some causes of tissue hypoxia will be reversible with HBO₂T, whereas some will not (e.g., in the presence of severe large vessel disease). When tissue hypoxia can be overcome by a high driving pressure of oxygen in the arterial blood, this can be demonstrated by measuring the tissue partial pressure of oxygen using an implantable oxygen electrode or, more commonly, a modified transcutaneous Clarke electrode.

The intermittent presentation of oxygen to those hypoxic tissues facilitates a resumption of healing. These short exposures to high oxygen tensions have long-lasting effects (at least 24 h) on a wide range of healing processes (Fig. 477e-3). The result is a gradual improvement in oxygen tension around the wound that reaches a plateau in experimental studies at about 20 treatments over 4 weeks. Improvements in oxygenation are associated with an eight- to ninefold increase in vascular density over both normobaric oxygen and air-breathing controls.

Clinical Evidence The typical course of HBO₂T consists of 20–30 once-daily compressions to 2–2.4 ATA for 1.5–2 h each session, but is highly dependent on the clinical response. There are many case series in the literature supporting the use of HBO₂T for a wide range of problem wounds. Both retrospective and prospective cohort studies suggest that 6 months after a course of therapy, about 70% of indolent ulcers will be substantially improved or healed. Often these ulcers have been present for many months or years, suggesting that the application of HBO₂T has a profound effect, either primarily or as a facilitator of other strategies. A recent Cochrane review included nine randomized controlled trials (RCTs) and concluded that the chance of ulcer healing improved about fivefold with HBO₂T (RR 5.20; 95% CI 1.25–21.66; $p = .02$).

Although there was a trend to benefit with HBO₂T, there was no statistically significant difference in the rate of major amputations (RR 0.36; 95% CI 0.11–1.18).

CARBON MONOXIDE POISONING

Carbon monoxide (CO) is a colorless, odorless gas formed during incomplete hydrocarbon combustion. Although CO is an essential endogenous neurotransmitter linked to NO metabolism and activity, it is also a leading cause of poisoning death, and in the United States alone results in more than 50,000 emergency department visits per year and about 2000 deaths. Although there are large variations from country to country, about half of nonlethal exposures are due to self-harm. Accidental poisoning is commonly associated with defective or improperly installed heaters, house fires, and industrial exposures. The motor vehicle is by far the most common source of intentional poisoning.

Pathology and Clinical Course The pathophysiology of carbon monoxide exposure is incompletely understood. CO binds to hemoglobin with an affinity more than 200 times that of oxygen, directly reducing the oxygen-carrying capacity of blood, and further promoting tissue hypoxia by shifting the oxyhemoglobin dissociation curve to the left. CO is also an anesthetic agent that inhibits evoked responses and narcotizes experimental animals in a dose-dependent manner. The associated loss of airway patency together with reduced oxygen carriage in blood may cause death from acute arterial hypoxia in severe poisoning. CO may also cause harm by other mechanisms including direct disruption of cellular oxidative processes, binding to myoglobin and hepatic cytochromes, and peroxidation of brain lipids.

The brain and heart are the most sensitive target organs due to their high blood flow, poor tolerance of hypoxia, and high oxygen requirements. Minor exposure may be asymptomatic or present with vague constitutional symptoms such as headache, lethargy, and nausea, whereas higher doses may present with poor concentration and cognition, short-term memory loss, confusion, seizures, and loss of consciousness. While carboxyhemoglobin (COHb) levels on admission do not necessarily reflect the severity or the prognosis of CO poisoning, cardiorespiratory arrest carries a very poor prognosis. Over the longer term, surviving patients commonly have neuropsychological sequelae. Motor disturbances, peripheral neuropathy, hearing loss, vestibular abnormalities, dementia, and psychosis have all been reported. Risk

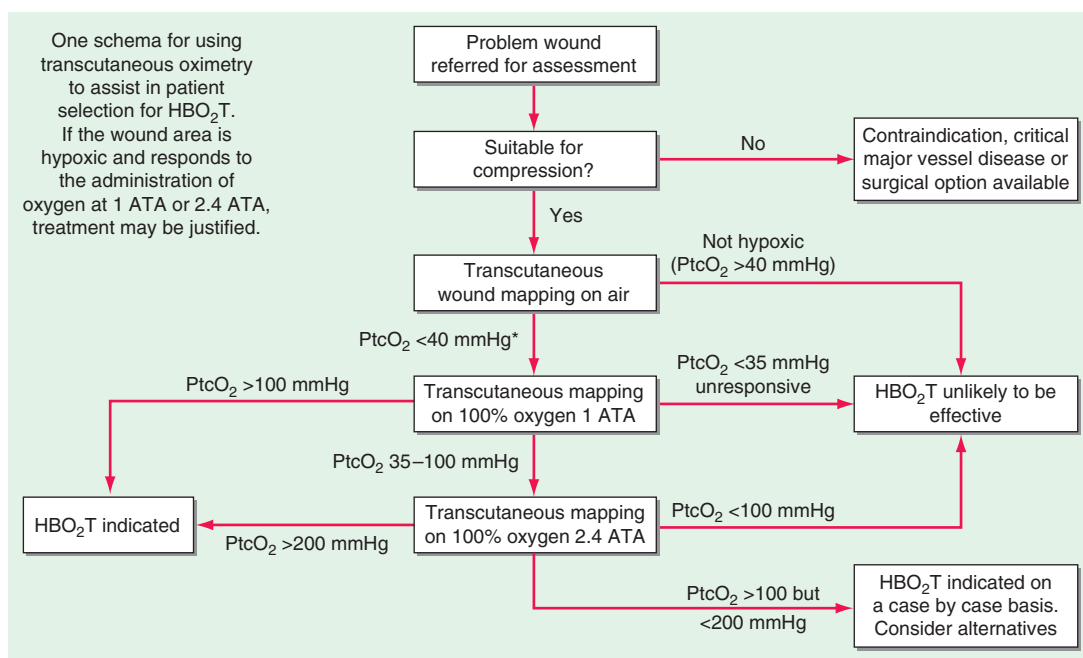


FIGURE 477e-4 Determining suitability for hyperbaric oxygen therapy (HBO₂T) guided by transcutaneous oximetry around the wound bed. *In diabetic patients, <50 mmHg may be more appropriate. PtcO₂, transcutaneous oxygen pressure.

477e-6 factors for poor outcome are age >35 years, exposure for >24 h, acidosis, and loss of consciousness.

Clinical Evidence The typical course of HBO₂T consists of two to three compressions to 2–2.8 ATA for 1.5–2 h each session. It is common for the first two compressions to be delivered within 24 h of the exposure. CO poisoning is one of the longest-standing indications for HBO₂T—based largely on the obvious connection between exposure, tissue hypoxia, and the ability of HBO₂T rapidly to overcome this hypoxia. CO is eliminated rapidly via the lungs on application of HBO₂T, with a half-life of about 21 min at 2.0 ATA versus 5.5 h breathing air and 71 min breathing oxygen at sea level. In practice, however, it seems unlikely that HBO₂T can be delivered in time to prevent either acute hypoxic death or irreversible global cerebral hypoxic injury. If HBO₂T is beneficial in CO poisoning, it must reduce the likelihood of persisting and/or delayed neurocognitive deficit through a mechanism other than the simple reversal of arterial hypoxia due to high levels of COHb. The difficulty in accurately assessing neurocognitive deficit has been one of the primary sources of controversy surrounding the clinical evidence in this area. To date there have been six randomized controlled trials of HBO₂T for CO poisoning, although only four have been reported in full. While a Cochrane review suggested that overall there is insufficient evidence to confirm a beneficial effect of HBO₂T on the chance of persisting neurocognitive deficit following poisoning (34% of patients treated with oxygen at 1 atmosphere vs 29%, of those treated with HBO₂T; odds ratio [OR] 0.78; 95% CI 0.54–1.1), this may have more to do with poor reporting and inadequate follow-up than with evidence that HBO₂T is not effective. The interpretation of the literature has much to do with how one defines neurocognitive deficit. In the most methodologically rigorous of these studies (Weaver et al.), a professionally administered battery of validated neuropsychological tests and a definition based on the deviation of individual subtest scores from the age-adjusted normal values was used; if the patient complained of memory, attention, or concentration difficulties, the required decrement was decreased. Using this approach, 6 weeks after poisoning, 46% of patients treated with normobaric oxygen alone had cognitive sequelae compared to 25% of those who received HBO₂T ($p = .007$; number needed to treat [NNT] = 5; 95% CI 3–16). At 12 months, the difference remained significant (32% vs 18%; $p = .04$; NNT = 7; 95% CI 4–124) despite considerable loss to follow-up.

On this basis, HBO₂T remains widely advocated for the routine treatment of patients with moderate to severe poisoning—in particular in those older than 35 years, presenting with a metabolic acidosis on arterial blood-gas analysis, exposed for lengthy periods, or with a history of unconsciousness. Conversely, many toxicologists remain unconvinced about the place of HBO₂T in this situation and call for further well-designed studies.

DIVING MEDICINE

INTRODUCTION

Underwater diving is both a popular recreational activity and a means of employment in a range of tasks from underwater construction to military operations. It is a complex activity with unique hazards and medical complications arising mainly as a consequence of the dramatic changes in pressure associated with both descent and ascent through the water column. For every 10.13 m increase in depth of seawater, the ambient pressure (P_{amb}) increases by 101.3 kPa (1 atmosphere) so that a diver at 20 m depth is exposed to a P_{amb} of approximately 303.9 kPa (3 ATA), made up of 1 ATA due to atmospheric pressure and 2 ATA generated by the water column.

BREATHING EQUIPMENT

Most diving is undertaken using a self-contained underwater breathing apparatus (scuba) consisting of one or more cylinders of compressed gas connected to a pressure-reducing regulator and a demand valve activated by inspiratory effort. Some divers use “rebreathers,” which are scuba devices that are closed or semiclosed circle systems with a carbon dioxide scrubber and a system designed to maintain a safe inspired PO₂. Exhaled gas is recycled, and gas consumption

is limited to little more than the oxygen metabolized by the diver. Rebreathers are therefore popular for deep dives where expensive helium is included in the respired mix (see below). Occupational divers frequently use “surface supply” equipment where gas, along with other utilities such as communications and power, is supplied via an umbilical from the surface.

All these systems must supply gas to the diver at the P_{amb} of the surrounding water or inspiration would be impossible against the surrounding water pressure. For most recreational diving, the respired gas is air. Pure oxygen is rarely used because oxygen may provoke seizures above an inspired PO₂ of 162 kPa (1.6 ATA) in aquatic environments, limiting the practical safe depth to 6 m. This is a conspicuously lower PO₂ than routinely used for hyperbaric therapy, reflecting a higher risk of both seizures and pulmonary toxicity during immersion. For the same reason, very deep diving requires the use of oxygen fractions lower than in air (FO₂ 0.21). This is because breathing air at 66 m means inspiring 1.6 ATA of oxygen, the maximum allowable pressure. To dive any deeper, breathing gases must contain less oxygen than air. Deep-diving gases often include helium instead of nitrogen to reduce both the narcotic effect and high gas density that result from breathing nitrogen at high pressures.

SUITABILITY FOR DIVING

The most common reason for physician consultation in relation to diving is for the evaluation of suitability for diver training or after a health event. Occupational diver candidates are usually compelled to see doctors with specialist training in the field, both at entry to the industry and periodically thereafter, and their medical evaluations are usually conducted according to legally mandated standards. In contrast, in most jurisdictions prospective recreational diver candidates simply complete a self-assessment medical questionnaire prior to diver training. If there are no positive responses, the candidate proceeds directly to training, but positive responses mandate the candidate see a doctor for evaluation of the identified medical issue. Prospective divers will often present to their family medicine practitioner for this purpose. In the modern era, such consultations have evolved from a simple proscriptive exercise of excluding those with potential contraindications to a more “risk analysis” approach in which each case is evaluated on its own merits. Such analyses require integration of diving physiology, the impact of associated medical problems, and a detailed knowledge of the specific medical condition of the candidate. A detailed discussion of the subject is beyond the scope of this chapter, but a few important principles are outlined below.

There are three primary questions that should be answered: (1) Could the underlying condition be exacerbated by diving? (2) Could the condition make a diving medical problem more likely? (3) Could the condition prevent the diver from meeting the functional requirements of diving? As examples, epilepsy is usually considered a contraindication because there are epileptogenic stimuli encountered in diving that could make a seizure more likely (such as thermal stress and exercise). Active asthma is a relative contraindication because it could predispose to pulmonary barotrauma (see below), and untreated ischemic heart disease is a contraindication because it could prevent a diver from exercising sufficiently to get out of a difficult situation such as being caught in a current. It can be a complex matter to recognize the relevant interactions between diving and medical conditions and to determine the impact on suitability for diving. Physicians interested in regularly conducting such evaluations should obtain relevant training. Short courses providing relevant training are offered by specialist groups in most countries.

BAROTRAUMA

The problem of middle-ear barotrauma (MEBT) with diving is similar to the problem that may occur during descent from altitude in an airplane, but difficulties with equalizing pressure in the middle ear are exaggerated underwater by both the rapidity and magnitude of pressure change as a diver descends or ascends. Failure to periodically insufflate the middle-ear spaces via the eustachian tubes during descent results in increasing pain. As the P_{amb} increases, the tympanic membrane (TM)

may be bruised or even ruptured as it is pushed inward. Negative pressure in the middle ear results in engorgement of blood vessels in the mucous membranes and leads to effusion or bleeding, which can be associated with a *conductive* hearing loss. MEBT is much less common during ascent because expanding gas in the middle-ear space tends to open the eustachian tube easily and “automatically.” Barotrauma may also affect the respiratory sinuses, although the sinus ostia are usually widely patent and allow automatic pressure equalization without the need for specific maneuvers. If equalization fails, pain usually results in termination of the dive. Difficulty with equalizing ears or sinuses may respond to oral or nasal decongestants.

Much less commonly the inner ear may suffer barotrauma (IEBT). Several explanations have been proposed, of which the most favored holds that forceful attempts to insufflate the middle-ear space by Valsalva maneuvers during descent cause sudden transmission of pressure to the perilymph via the cochlear aqueduct and outward rupture of the round window already under tension because of negative middle ear pressure. The clinician should be alerted to possible IEBT after diving by a *sensorineural* hearing loss or true vertigo (which is often accompanied by nausea, vomiting, nystagmus, and ataxia). These manifestations can also occur in vestibulocochlear DCS (see below) but should never be attributed to MEBT. Immediate review by an expert diving physician is recommended, and urgent referral to an otologist will often follow.

The lungs are also vulnerable to barotrauma but are at most risk during ascent. If expanding gas becomes trapped in the lungs as P_{amb} falls, this may rupture alveoli and associated vascular tissue. Gas trapping may occur if divers intentionally or involuntarily hold their breath during ascent or if there are bullae. The extent to which asthma predisposes to pulmonary barotrauma is debated, but the presence of active bronchoconstriction must increase risk. For this reason, asthmatics who regularly require bronchodilator medications or whose airways are sensitive to exercise or cold air are usually discouraged from diving. While possible consequences of pulmonary barotrauma include pneumothorax and mediastinal emphysema, the most feared is the introduction of gas into the pulmonary veins leading to cerebral arterial gas embolism (CAGE). Manifestations of CAGE include loss of consciousness, confusion, hemiplegia, visual disturbances, and speech difficulties, appearing immediately or within minutes after surfacing. The management is the same as for DCS described below. It is notable that the natural history of CAGE often includes substantial or complete resolution of symptoms early after the event. This is probably the clinical correlate of bubble involution and redistribution with consequent restoration of flow. Patients exhibiting such remissions should still be reviewed at specialist diving medical centers because secondary deterioration or reembolization can occur. Unsurprisingly, these events can be misdiagnosed as typical strokes or transient ischemic attacks (TIAs) (Chap. 455) when patients are seen by those unfamiliar with diving medicine. *All patients presenting with neurologic symptoms after diving should have their symptoms discussed with a specialist in diving medicine and be considered for recompression therapy.*

DECOMPRESSION SICKNESS

DCS is caused by the formation of bubbles from dissolved inert gas (usually nitrogen) during or after ascent (decompression) from a compressed gas dive. Bubble formation is also possible following decompression for extravehicular activity during space flight and with ascent to altitude in unpressurized aircraft. DCS in the latter scenarios is probably rare in comparison with diving, where the incidence is approximately 1:10,000 recreational dives.

Breathing at elevated P_{amb} results in increased uptake of inert gas into blood and then into tissues. The rate at which tissue inert gas equilibrates with the inspired inert gas pressure is proportional to tissue blood flow and the blood-tissue partition coefficient for the gas. Similar factors dictate the kinetics of inert gas washout during ascent. If the rate of gas washout from tissues does not match the rate of decline in P_{amb} , then the sum of dissolved gas pressures in the tissue will exceed P_{amb} , a condition referred to as “supersaturation.” This is the prerequisite for bubbles to form during decompression, although other

less well-understood factors are also involved. Deeper and longer dives result in greater inert gas absorption and greater likelihood of tissue supersaturation during ascent. Divers control their ascent for a given depth and time exposure using algorithms that often include periods where ascent is halted for a prescribed period at different depths to allow time for gas washout (“decompression stops”). Although a breach of these protocols increases the risk of DCS, adherence does not guarantee against it. DCS should be considered in any diver manifesting symptoms not readily explained by an alternative mechanism.

Bubbles may form within tissues themselves, where they cause symptoms by mechanical distraction of pain-sensitive or functionally important structures. They also appear in the venous circulation as blood passes through supersaturated tissues. Some venous bubbles are tolerated without symptoms and are filtered from the circulation in the pulmonary capillaries. However, in sufficiently large numbers, these bubbles are capable of inciting inflammatory and coagulation cascades, damaging endothelium, activating formed elements of blood such as platelets, and causing symptomatic pulmonary vascular obstruction. Moreover, if there is a right-to-left shunt such as through a patent foramen ovale (PFO) or an intrapulmonary shunt, then venous bubbles may enter the arterial circulation (25% of adults have a probe-patent PFO). The risk of cerebral, spinal cord, inner ear, and skin manifestations appears higher in the presence of significant shunts, suggesting that these “arterialized” venous bubbles can cause harm, perhaps by disrupting flow in the microcirculation of target organs. Circulating endothelial microparticles, which are elevated in number and size after diving, are currently under investigation as indicators of decompression stress or possibly as injurious agents in their own right. How they arise, and their role in DCS, remain unclear.

Table 477e-3 lists manifestations of DCS grouped according to organ system. The majority of cases present with mild symptoms, including musculoskeletal pain, fatigue, and minor neurologic manifestations such as patchy paresthesias. Serious presentations are much less common. Pulmonary and cardiovascular manifestations can be

TABLE 477e-3 MANIFESTATIONS OF DECOMPRESSION SICKNESS

Organ System	Manifestations
Musculoskeletal	Limb pain
Neurologic	
Cerebral	Confusion Visual disturbances Speech disturbances
Spinal	Muscular weakness Paralysis Upper motor neuron signs Bladder and sphincter dysfunction Dermatomal sensory disturbances Abdominal pain Girdle pain
Vestibulocochlear	Hearing loss Vertigo and ataxia Nausea and vomiting
Peripheral	Patchy nondermatomal sensory disturbance
Pulmonary	Cough Dyspnea Pulmonary edema (rare)
Cardiovascular	Chest pain Arrhythmia Hemoconcentration Coagulopathy Hypotension
Cutaneous	Rash, itch
Lymphatic	Soft tissue edema, often relatively localized
Constitutional	Fatigue and malaise

477e-8 life-threatening, and spinal cord involvement frequently results in permanent disability. Latency is variable. Serious DCS usually manifests within minutes of surfacing, but mild symptoms may not appear for several hours. Symptoms arising more than 24 h after diving are very unlikely to be DCS. The presentation may be confusing and nonspecific, and there are as yet no useful diagnostic investigations. Diagnosis is based on integration of findings from examination of the dive profile, the nature and temporal relationship of symptoms, and the clinical examination. Some DCS presentations may be difficult to separate from CAGE following pulmonary barotrauma, but from a clinical perspective the distinction is unimportant because the first aid and definitive management of both conditions are the same.

TREATMENT DIVING MEDICINE

First aid includes horizontal positioning (especially if there are cerebral manifestations), intravenous fluids if available, and sustained 100% oxygen administration. The latter accelerates inert gas wash-out from tissues and promotes resolution of bubbles. Definitive treatment of DCS or CAGE with recompression and hyperbaric oxygen is justified in most instances, although some mild or marginal DCS cases may be managed with first aid measures, an option that may be invoked under various circumstances, but especially if evacuation for recompression is hazardous or extremely difficult.

Long-distance evacuations are usually undertaken using a helicopter flying at low altitude or a fixed wing air ambulance pressurized to 1 atmosphere pressure.

Recompression reduces bubble volume in accordance with Boyle's law and increases the inert gas partial pressure difference between a bubble and surrounding tissue. At the same time, oxygen administration markedly increases the inert gas partial pressure difference between alveoli and tissue. The net effect is to significantly increase the rate of inert gas diffusion from bubble to tissue and tissue to blood, thus accelerating bubble resolution. Hyperbaric oxygen also helps oxygenate compromised tissues and appears to ameliorate some of the proinflammatory effects of bubbles. Various recompression protocols have been advocated, but there are no data that define the optimum approach. It typically begins with oxygen administered at 2.8 atmospheres absolute, the maximum pressure at which the risk of oxygen toxicity remains acceptable in a hyperbaric chamber. There follows a stepwise decompression over variable periods adjusted to symptom response. The most widely used algorithm is the U.S. Navy Table 6, whose shortest format lasts 4 h and 45 min. Typically, shorter "follow-up" recompressions are repeated daily while symptoms persist and appear responsive to treatment. Adjuncts to recompression include intravenous fluids and other supportive care as necessary. Occasionally, very sick divers require high-level intensive care.

HYPOTHERMIA

Accidental hypothermia occurs when there is an unintentional drop in the body's core temperature below 35°C (95°F). At this temperature, many of the compensatory physiologic mechanisms that conserve heat begin to fail. *Primary accidental hypothermia* is a result of the direct exposure of a previously healthy individual to the cold. The mortality rate is much higher for patients who develop *secondary hypothermia* as a complication of a serious systemic disorder.

CAUSES

Primary accidental hypothermia is geographically and seasonally pervasive. Although most cases occur in the winter months and in colder climates, this condition is surprisingly common in warmer regions as well. Multiple variables render individuals at the extremes of age—both the elderly and neonates—particularly vulnerable to hypothermia (Table 478e-1). The elderly have diminished thermal perception and are more susceptible to immobility, malnutrition, and systemic illnesses that interfere with heat generation or conservation. Dementia, psychiatric illness, and socioeconomic factors often compound these problems by impeding adequate measures to prevent hypothermia. Neonates have high rates of heat loss because of their increased surface-to-mass ratio and their lack of effective shivering and adaptive behavioral responses. At all ages, malnutrition can contribute to heat loss because of diminished subcutaneous fat and as a result of depleted energy stores used for thermogenesis.

Individuals whose occupations or hobbies entail extensive exposure to cold weather are at increased risk for hypothermia. Military history is replete with hypothermic tragedies. Hunters, sailors, skiers, and climbers also are at great risk of exposure, whether it involves injury, changes in weather, or lack of preparedness.

Ethanol causes vasodilation (which increases heat loss), reduces thermogenesis and gluconeogenesis, and may impair judgment or lead to obtundation. Phenothiazines, barbiturates, benzodiazepines, heterocyclic antidepressants, and many other medications reduce centrally mediated vasoconstriction. Up to 25% of patients admitted to an intensive care unit because of drug overdose are hypothermic. Anesthetics can block shivering responses; their effects are compounded when patients are not insulated adequately in the operating or recovery units.

TABLE 478e-1 RISK FACTORS FOR HYPOTHERMIA

Age extremes	Endocrine-related
Elderly	Diabetes mellitus
Neonates	Hypoglycemia
Environmental exposure	Hypothyroidism
Occupational	Adrenal insufficiency
Sports-related	Hypopituitarism
Inadequate clothing	Neurologic
Immersion	Cerebrovascular accident
Toxicologic and pharmacologic	Hypothalamic disorders
Ethanol	Parkinson's disease
Phenothiazines	Spinal cord injury
Barbiturates	Multisystemic
Anesthetics	Trauma
Neuromuscular blockers	Sepsis
Antidepressants	Shock
Insufficient fuel	Hepatic or renal failure
Malnutrition	Carcinomatosis
Marasmus	Burns and exfoliative dermatologic disorders
Kwashiorkor	Immobility or debilitation

Several types of endocrine dysfunction can lead to hypothermia. Hypothyroidism—particularly when extreme, as in myxedema coma—reduces the metabolic rate and impairs thermogenesis and behavioral responses. Adrenal insufficiency and hypopituitarism also increase susceptibility to hypothermia. Hypoglycemia, most commonly caused by insulin or oral hypoglycemic drugs, is associated with hypothermia, in part because of neuroglycopenic effects on hypothalamic function. Increased osmolality and metabolic derangements associated with uremia, diabetic ketoacidosis, and lactic acidosis can lead to altered hypothalamic thermoregulation.

Neurologic injury from trauma, cerebrovascular accident, subarachnoid hemorrhage, and hypothalamic lesion increases susceptibility to hypothermia. Agenesis of the corpus callosum (*Shapiro syndrome*) is one cause of episodic hypothermia; in this syndrome, profuse perspiration is followed by a rapid fall in temperature. Acute spinal cord injury disrupts the autonomic pathways that lead to shivering and prevents cold-induced reflex vasoconstrictive responses.

Hypothermia associated with sepsis is a poor prognostic sign. Hepatic failure causes decreased glycogen storage and gluconeogenesis as well as a diminished shivering response. In acute myocardial infarction associated with low cardiac output, hypothermia may be reversed after adequate resuscitation. With extensive burns, psoriasis, erythrodermas, and other skin diseases, increased peripheral-blood flow leads to excessive heat loss.

THERMOREGULATION

Heat loss occurs through five mechanisms: radiation (55–65% of heat loss), conduction (10–15% of heat loss, much increased in cold water), convection (increased in the wind), respiration, and evaporation; both of the latter two mechanisms are affected by the ambient temperature and the relative humidity.

The preoptic anterior hypothalamus normally orchestrates thermoregulation (Chap. 23). The immediate defense of thermoneutrality is via the autonomic nervous system, whereas delayed control is mediated by the endocrine system. Autonomic nervous system responses include the release of norepinephrine, increased muscle tone, and shivering, leading to thermogenesis and an increase in the basal metabolic rate. Cutaneous cold thermoreception causes direct reflex vasoconstriction to conserve heat. Prolonged exposure to cold also stimulates the thyroid axis, leading to an increased metabolic rate.

CLINICAL PRESENTATION

In most cases of hypothermia, the history of exposure to environmental factors (e.g., prolonged exposure to the outdoors without adequate clothing) makes the diagnosis straightforward. In urban settings, however, the presentation is often more subtle, and other disease processes, toxin exposures, or psychiatric diagnoses should be considered.

After initial stimulation by hypothermia, there is progressive depression of all organ systems. The timing of the appearance of these clinical manifestations varies widely (Table 478e-2). Without knowing the core temperature, it can be difficult to interpret other vital signs. For example, tachycardia disproportionate to the core temperature suggests secondary hypothermia resulting from hypoglycemia, hypovolemia, or a toxin overdose. Because carbon dioxide production declines progressively, the respiratory rate should be low; persistent hyperventilation suggests a central nervous system (CNS) lesion or one of the organic acidoses. A markedly depressed level of consciousness in a patient with mild hypothermia raises suspicion of an overdose or CNS dysfunction due to infection or trauma.

Physical examination findings can also be altered by hypothermia. For instance, the assumption that areflexia is solely attributable to hypothermia can obscure and delay the diagnosis of a spinal cord lesion. Patients with hypothermia may be confused or combative; these symptoms abate more rapidly with rewarming than with chemical or physical restraint. A classic example of maladaptive behavior in patients with hypothermia is paradoxical undressing, which involves the inappropriate removal of clothing in response to a cold stress. The cold-induced ileus and abdominal rectus spasm can mimic or mask the presentation of an acute abdomen (Chap. 20).

TABLE 478e-2 PHYSIOLOGIC CHANGES ASSOCIATED WITH ACCIDENTAL HYPOTHERMIA

Severity	Body Temperature	Central Nervous System	Cardiovascular	Respiratory	Renal and Endocrine	Neuromuscular
Mild	35°C (95°F)–32.2°C (90°F)	Linear depression of cerebral metabolism; amnesia; apathy; dysarthria; impaired judgment; maladaptive behavior	Tachycardia, then progressive bradycardia; cardiac cycle prolongation; vasoconstriction; increase in cardiac output and blood pressure	Tachypnea, then progressive decrease in respiratory minute volume; declining oxygen consumption; bronchorrhea; bronchospasm	Diuresis; increase in catecholamines, adrenal steroids, triiodothyronine, and thyroxine; increase in metabolism with shivering	Increased preshivering muscle tone, then fatiguing
Moderate	<32.2°C (90°F)–28°C (82.4°F)	EEG abnormalities; progressive depression of level of consciousness; pupillary dilation; paradoxical undressing; hallucinations	Progressive decrease in pulse and cardiac output; increased atrial and ventricular arrhythmias; suggestive (J-wave) ECG changes	Hypoventilation; 50% decrease in carbon dioxide production per 8°C (46°F) drop in temperature; absence of protective airway reflexes	50% increase in renal blood flow; renal autoregulation intact; impaired insulin action	Hyporeflexia; diminishing shivering-induced thermogenesis; rigidity
Severe	<28°C (<82.4°F)	Loss of cerebrovascular autoregulation; decline in cerebral blood flow; coma; loss of ocular reflexes; progressive decrease in EEG abnormalities	Progressive decrease in blood pressure, heart rate, and cardiac output; reentrant dysrhythmias; maximal risk of ventricular fibrillation; asystole	Pulmonic congestion and edema; 75% decrease in oxygen consumption; apnea	Decrease in renal blood flow that parallels decrease in cardiac output; extreme oliguria; poikilothermia; 80% decrease in basal metabolism	No motion; decreased nerve-conduction velocity; peripheral areflexia; no corneal or oculocephalic reflexes

Abbreviations: ECG, electrocardiogram; EEG, electroencephalogram.

Source: Modified from DF Danz, RS Pozos: *N Engl J Med* 331:1756, 1994.

When a patient in hypothermic cardiac arrest is first discovered, cardiopulmonary resuscitation is indicated unless (1) a do-not-resuscitate status is verified, (2) obviously lethal injuries are identified, or (3) the depression of a frozen chest wall is not possible. As the resuscitation proceeds, the prognosis is grave if there is evidence of widespread cell lysis, as reflected by potassium levels >10–12 mmol/L (10–12 meq/L). Other findings that may preclude continuing resuscitation include a core temperature <10–12°C (<50–54°F), a pH <6.5, and evidence of intravascular thrombosis with a fibrinogen value <0.5 g/L (<50 mg/dL). The decision to terminate resuscitation before rewarming the patient past 33°C (91°F) should be predicated on the type and severity of the precipitants of hypothermia. There are no validated prognostic indicators for recovery from hypothermia. A history of asphyxia with secondary cooling is the most important negative predictor of survival.

DIAGNOSIS AND STABILIZATION

Hypothermia is confirmed by measurement of the core temperature, preferably at two sites. Rectal probes should be placed to a depth of 15 cm and not adjacent to cold feces. A simultaneous esophageal probe should be placed 24 cm below the larynx; it may read falsely high during heated inhalation therapy. Relying solely on infrared tympanic thermography is not advisable.

After a diagnosis of hypothermia is established, cardiac monitoring should be instituted, along with attempts to limit further heat loss. If the patient is in ventricular fibrillation, it is unclear at what core temperature ventricular defibrillation (2 J/kg) should first be attempted. One attempt below 30°C is warranted. Further defibrillation attempts should be deferred until some rewarming (1°–2°C) is achieved and ventricular fibrillation is coarser. Although cardiac pacing for hypothermic bradycardias is rarely indicated, the transthoracic technique is preferable.

Supplemental oxygenation is always warranted, since tissue oxygenation is affected adversely by the leftward shift of the oxyhemoglobin dissociation curve. Pulse oximetry may be unreliable in patients with vasoconstriction. If protective airway reflexes are absent, gentle endotracheal intubation should be performed. Adequate preoxygenation will prevent ventricular arrhythmias.

Insertion of a gastric tube prevents dilation secondary to decreased bowel motility. Indwelling bladder catheters facilitate monitoring of cold-induced diuresis. Dehydration is encountered commonly with chronic hypothermia, and most patients benefit from an intravenous or intraosseous bolus of crystalloid. Normal saline is preferable to

lactated Ringer's solution, as the liver in hypothermic patients inefficiently metabolizes lactate. The placement of a pulmonary artery catheter can cause perforation of the less compliant pulmonary artery. Insertion of a central venous catheter deeply into the cold right atrium should be avoided since this procedure can precipitate arrhythmias.

Arterial blood gases should not be corrected for temperature (Chap. 66). An uncorrected pH of 7.42 and a Pco₂ of 40 mmHg reflect appropriate alveolar ventilation and acid-base balance at any core temperature. Acid-base imbalances should be corrected gradually, since the bicarbonate buffering system is inefficient. A common error is overzealous hyperventilation in the setting of depressed CO₂ production. When the Pco₂ decreases by 10 mmHg at 28°C (82°F), it doubles the pH increase of 0.08 that occurs at 37°C (99°F).

The severity of anemia may be underestimated because the hematocrit increases 2% for each 1°C drop in temperature. White blood cell sequestration and bone marrow suppression are common, potentially masking an infection. Although hypokalemia is more common in chronic hypothermia, hyperkalemia also occurs; the expected electrocardiographic changes can be obscured by hypothermia. Patients with renal insufficiency, metabolic acidoses, or rhabdomyolysis are at greatest risk for electrolyte disturbances.

Coagulopathies are common because cold inhibits the enzymatic reactions required for activation of the intrinsic cascade. In addition, thromboxane B₂ production by platelets is temperature dependent, and platelet function is impaired. The administration of platelets and fresh-frozen plasma is therefore not effective. The prothrombin or partial thromboplastin times or the international normalized ratio can be deceptively normal and contrast with the observed *in vivo* coagulopathy. This contradiction occurs because all coagulation tests are routinely performed at 37°C (99°F), and the enzymes are thus rewarmed.

REWARMING STRATEGIES

The key initial decision is whether to rewarm the patient passively or actively. *Passive external rewarming* simply involves covering and insulating the patient in a warm environment. With the head also covered, the rate of rewarming is usually 0.5°–2°C (1.10°–4.4°F) per hour. This technique is ideal for previously healthy patients who develop acute, mild primary accidental hypothermia. The patient must have sufficient glycogen to support endogenous thermogenesis.

The application of heat directly to the extremities of patients with chronic severe hypothermia should be avoided because it can induce peripheral vasodilation and precipitate core temperature “afterdrop,” a

response characterized by a continual decline in the core temperature after removal of the patient from the cold. Truncal heat application reduces the risk of afterdrop.

Active rewarming is necessary under the following circumstances: core temperature $<32^{\circ}\text{C}$ ($<90^{\circ}\text{F}$) (*poikilothermia*), cardiovascular instability, age extremes, CNS dysfunction, hormone insufficiency, and suspicion of secondary hypothermia. *Active external rewarming* is best accomplished with forced-air heating blankets. Other options include devices that circulate water through external heat exchange pads, radiant heat sources, and hot packs. Monitoring a patient with hypothermia in a heated tub is extremely difficult. Electric blankets should be avoided because vasoconstricted skin is easily burned.

There are numerous widely available options for *active core rewarming*. Airway rewarming with heated humidified oxygen ($40^{\circ}\text{--}45^{\circ}\text{C}$ [$104^{\circ}\text{--}113^{\circ}\text{F}$]) via mask or endotracheal tube is a convenient option. Although airway rewarming provides less heat than do some other forms of active core rewarming, it eliminates respiratory heat loss and adds $1^{\circ}\text{--}2^{\circ}\text{C}$ ($1.1^{\circ}\text{--}4.4^{\circ}\text{F}$) to the overall rewarming rate. Crystalloids should be heated to $40^{\circ}\text{--}42^{\circ}\text{C}$ ($104^{\circ}\text{--}108^{\circ}\text{F}$), but the quantity of heat provided is significant only during massive volume resuscitation. The most efficient method for heating and delivering fluid or blood is with a countercurrent in-line heat exchanger. Heated irrigation of the gastrointestinal tract or bladder transfers minimal heat because of the limited available surface area. These methods should be reserved for patients in cardiac arrest and then used in combination with all available active rewarming techniques.

Closed thoracic lavage is far more efficient in severely hypothermic patients with cardiac arrest. The hemithoraxes are irrigated through two inserted large-bore thoracostomy tubes. Thoracostomy tubes should not be placed in the left chest of a spontaneously perfusing patient for purposes of rewarming. Peritoneal lavage with the dialysate at $40^{\circ}\text{--}45^{\circ}\text{C}$ ($104^{\circ}\text{--}113^{\circ}\text{F}$) efficiently transfers heat when delivered through two catheters with outflow suction. Like peritoneal dialysis, standard hemodialysis is especially useful for patients with electrolyte abnormalities, rhabdomyolysis, or toxin ingestion. Another option involves the central venous insertion of a rapid endovascular warming device.

Extracorporeal blood rewarming options (Table 478e-3) should be considered in severely hypothermic patients, especially those with *primary accidental hypothermia*. Cardiopulmonary bypass should be considered in nonperfusing patients without documented contraindications to resuscitation. Circulatory support may be the only effective option in patients with completely frozen extremities or those with significant tissue destruction coupled with rhabdomyolysis. There is no evidence that extremely rapid rewarming improves survival in perfusing patients. The best strategy is usually a combination of passive, truncal active, and active core rewarming techniques.

TREATMENT HYPOTHERMIA

When a patient is hypothermic, target organs and the cardiovascular system respond minimally to most medications. Generally, IV medications are withheld below 30°C (86°F). In contrast to antiarrhythmics, low-dose vasopressor medications may improve the intra-arrest rates of return of spontaneous circulation. Because of increased binding of drugs to proteins as well as impaired metabolism and excretion, either a lower dose or a longer interval between doses should be used to avoid toxicity. As an example, the administration of repeated doses of digoxin or insulin would be ineffective while the patient is hypothermic, and the residual drugs would be potentially toxic during rewarming.

Achieving a mean arterial pressure of at least 60 mmHg should be an early objective. If the hypotension does not respond to crystalloid/colloid infusion and rewarming, low-dose dopamine support ($2\text{--}5\ \mu\text{g}/\text{kg}$ per min) should be considered. Perfusion of the vasoconstricted cardiovascular system also may be improved with low-dose IV nitroglycerin.

Atrial arrhythmias should be monitored initially without intervention, as the ventricular response should be slow and, unless preexistent, most will convert spontaneously during rewarming. The role of

TABLE 478e-3 OPTIONS FOR EXTRACORPOREAL BLOOD REWARMING

Extracorporeal Rewarming Technique	Considerations
Continuous venovenous (CV) rewarming	Circuit: CV catheter to CV, dual-lumen CV, or peripheral catheter No oxygenator/circulatory support Flow rates 150–400 mL/min ROR $2^{\circ}\text{--}3^{\circ}\text{C}$ ($36^{\circ}\text{--}37^{\circ}\text{F}$)/h
Hemodialysis	Circuit: single- or dual-vessel cannulation Stabilizes electrolyte or toxicologic abnormalities Exchange cycle volumes 200–500 mL/min ROR $2^{\circ}\text{--}3^{\circ}\text{C}$ ($36^{\circ}\text{--}37^{\circ}\text{F}$)/h
Continuous arteriovenous rewarming (CAVR)	Circuit: percutaneous 8.5-Fr femoral catheters Requires systolic blood pressure of 60 mmHg No perfusionist/pump/anticoagulation Flow rates 225–375 mL/min ROR $3^{\circ}\text{--}4^{\circ}\text{C}$ ($37^{\circ}\text{--}39^{\circ}\text{F}$)/h
Cardiopulmonary bypass (CPB)	Circuit: full circulatory support with pump and oxygenator Perfusate-temperature gradient ($5^{\circ}\text{--}10^{\circ}\text{C}$ [$41^{\circ}\text{--}50^{\circ}\text{F}$]) Flow rates 2–7 L/min (average 3–4 L/min) ROR up to 9.5°C (49°F)/h
Extracorporeal membrane oxygenation (ECMO)	Decreased risk of post-rewarming cardiorespiratory failure

Abbreviations: CV, central venous; ROR, rate of rewarming.

prophylaxis and treatment of ventricular arrhythmias is problematic. Preexisting ventricular ectopy may be suppressed by hypothermia and reappear during rewarming. None of the class I agents has proved to be safe and efficacious. There is also insufficient evidence that the class III ventricular antiarrhythmic amiodarone is safe.

Initiating empirical therapy for adrenal insufficiency usually is not warranted unless the history suggests steroid dependence or hypoadrenalism or efforts to rewarm with standard therapy fail. The administration of parenteral levothyroxine to euthyroid patients with hypothermia, however, is potentially hazardous. Because laboratory results can be delayed and confounded by the presence of the sick euthyroid syndrome (Chap. 405), historic clues or physical findings suggestive of hypothyroidism should be sought. When myxedema is the cause of hypothermia, the relaxation phase of the Achilles reflex is prolonged more than is the contraction phase.

Hypothermia obscures most of the symptoms and signs of infection, notably fever and leukocytosis. Shaking rigors from infection may be mistaken for shivering. Except in mild cases, extensive cultures and repeated physical examinations are essential. Unless an infectious source is identified, empirical antibiotic prophylaxis is most warranted in the elderly, neonates, and immunocompromised patients.

Preventive measures should be discussed with high-risk individuals, such as the elderly and people whose work frequently exposes them to extreme cold. The importance of layered clothing and headgear, adequate shelter, increased caloric intake, and the avoidance of ethanol should be emphasized, along with access to rescue services.

FROSTBITE

Peripheral cold injuries include both freezing and nonfreezing injuries to tissue. Tissue freezes quickly when in contact with thermal conductors such as metal and volatile solutions. Other predisposing factors include constrictive clothing or boots, immobility, and vasoconstrictive medications. Frostbite occurs when the tissue temperature drops below 0°C (32°F). Ice-crystal formation subsequently distorts and

478e-4 destroys the cellular architecture. Once the vascular endothelium is damaged, stasis progresses rapidly to microvascular thrombosis. After the tissue thaws, there is progressive dermal ischemia. The microvasculature begins to collapse, arteriovenous shunting increases tissue pressures, and edema forms. Finally, thrombosis, ischemia, and superficial necrosis appear. The development of mummification and demarcation may take weeks to months.

CLINICAL PRESENTATION

The initial presentation of frostbite can be deceptively benign. The symptoms always include a sensory deficiency affecting light touch, pain, and temperature perception. The acral areas and distal extremities are the most common insensate areas. Some patients describe a clumsy or “chunk of wood” sensation in the extremity.

Deep frostbitten tissue can appear waxy, mottled, yellow, or violaceous-white. Favorable presenting signs include some warmth or sensation with normal color. The injury is often superficial if the subcutaneous tissue is pliable or if the dermis can be rolled over bony prominences.

Frostnip may precede frostbite. Frostnip is a nonfreezing cold injury resulting from intense vasoconstriction of exposed acral skin.

Clinically, it is most practical to classify frostbite as superficial or deep. Superficial frostbite does not entail tissue loss but rather causes only anesthesia and erythema. The appearance of vesiculation surrounded by edema and erythema implies deeper involvement (Fig. 478e-1). Hemorrhagic vesicles reflect a serious injury to the microvasculature and indicate severe frostbite. Damages in subcuticular, muscular, or osseous tissues may result in amputation.

The two most common nonfreezing peripheral cold injuries are *chilblain (pernio)* and *immersion (trench) foot*. Chilblain results from neuronal and endothelial damage induced by repetitive exposure to

dry cold. Young females, particularly those with a history of Raynaud’s phenomenon, are at greatest risk. Persistent vasospasticity and vasculitis can cause erythema, mild edema, and pruritus. Eventually plaques, blue nodules, and ulcerations develop. These lesions typically involve the dorsa of the hands and feet. In contrast, immersion foot results from repetitive exposure to wet cold above the freezing point. The feet initially appear cyanotic, cold, and edematous. The subsequent development of bullae is often indistinguishable from frostbite. This vesiculation rapidly progresses to ulceration and liquefaction gangrene. Patients with milder cases report hyperhidrosis, cold sensitivity, and painful ambulation for many years.

TREATMENT FROSTBITE

When frostbite accompanies hypothermia, hydration may improve vascular stasis. Frozen tissue should be thawed rapidly and completely by immersion in circulating water at 37°–40°C (99°–104°F). Rapid rewarming often produces an initial hyperemia. The early formation of large clear distal blebs is more favorable than that of smaller proximal dark hemorrhagic blebs. A common error is the premature termination of thawing, since the reestablishment of perfusion is intensely painful. Parenteral narcotics will be necessary with deep frostbite. If cyanosis persists after rewarming, the tissue compartment pressures should be monitored carefully.

Many antithrombotic and vasodilatory primary and adjunctive treatment regimens have been evaluated. The prostacyclin analogue iloprost in combination with aspirin may prove useful. There is no conclusive evidence that sympathectomy, steroids, calcium channel blockers, or hyperbaric oxygen salvages tissue.

Patients who have deep frostbite injuries with the potential for significant morbidity should be considered for intravenous or intraarterial thrombolytic therapy. Angiography or pyrophosphate scanning should help evaluate the injury and monitor the progress of tissue plasminogen activator therapy. Heparin is recommended as adjunctive therapy. Intraarterial thrombolysis may reduce the need for digital and more proximal amputations when administered within 24 h of severe injuries. A treatment protocol for frostbite is summarized in (Table 478e-4).

Unless infection develops, any decision regarding debridement or amputation should generally be deferred. Angiography or



FIGURE 478e-1 Frostbite with vesiculation, surrounded by edema and erythema.

TABLE 478e-4 TREATMENT FOR FROSTBITE

Before Thawing	During Thawing	After Thawing
Remove from environment.	Consider parenteral analgesia and ketorolac.	Gently dry and protect part; elevate; place pledgets between toes, if macerated.
Prevent partial thawing and refreezing.	Administer ibuprofen (400 mg PO).	If clear vesicles are intact, aspirate sterilely; if broken, debride and dress with antibiotic or sterile aloe vera ointment.
Stabilize core temperature and treat hypothermia.	Immerse part in 37°–40°C (99°–104°F) (thermometer-monitored) circulating water containing an antiseptic soap until distal flush (10–45 min).	Leave hemorrhagic vesicles intact to prevent desiccation and infection.
Protect frozen part—no friction or massage.	Encourage patient to gently move part.	Continue ibuprofen (400 mg PO [12 mg/kg per day] q12h).
Address medical or surgical conditions.	If pain is refractory, reduce water temperature to 35°–37°C (95°–99°F) and administer parenteral narcotics.	Consider tetanus and streptococcal prophylaxis; elevate part. Administer hydrotherapy at 37°C (99°F). Consider dextran or phenoxybenzamine or, in severe cases, thrombolysis

technetium-99 bone scan may assist in the determination of surgical margins. Magnetic resonance angiography may also demonstrate the line of demarcation earlier than does clinical demarcation.

The most common symptomatic sequelae reflect neuronal injury and persistently abnormal sympathetic tone, including paresthesias, thermal misperception, and hyperhidrosis. Delayed findings

include nail deformities, cutaneous carcinomas, and epiphyseal damage in children.

Management of the chilblain syndrome is usually supportive. With refractory perniosis, alternatives include nifedipine, steroids, and limaprost, a prostaglandin E₁ analogue.

479e Heat-Related Illnesses

Daniel F. Danzl

Heat-related illnesses include a spectrum of disorders ranging from heat syncope, muscle cramps, and heat exhaustion to medical emergencies such as heatstroke. The core body temperature is normally maintained within a very narrow range. Although significant levels of hypothermia are tolerated (Chap. 478e), multiorgan dysfunction occurs rapidly at temperatures $>41^{\circ}$ – 43° C. In contrast to severe hyperthermia, the far more common sign of fever reflects intact thermoregulation.

THERMOREGULATION

Humans are capable of significant heat generation. Strenuous exercise can increase heat generation twentyfold. The heat load from metabolic heat production and environmental heat absorption is balanced by a variety of heat dissipation mechanisms. These dissipation pathways are orchestrated by the central thermostat, which is located in the preoptic nucleus of the anterior hypothalamus. Efferent signals sent via the autonomic nervous system trigger cutaneous vasodilation and diaphoresis to facilitate heat loss.

Normally, the body dissipates heat into the environment via four mechanisms. The *evaporation* of skin moisture is the single most efficient mechanism of heat loss but becomes progressively ineffective as the relative humidity rises to $>70\%$. The *radiation* of infrared electromagnetic energy directly into the surrounding environment occurs continuously. (Conversely, radiation is a major source of heat gain in hot climates.) *Conduction*—the direct transfer of heat to a cooler object—and *convection*—the loss of heat to air currents—become ineffective when the environmental temperature exceeds the skin temperature.

Factors that interfere with the evaporation of diaphoresis significantly increase the risk of heat illness. Examples include dripping of sweat off the skin, constrictive or occlusive clothing, dehydration, and excessive humidity. While air is an effective insulator, the thermal conductivity of water is 25 times greater than that of air at the same temperature. The *wet-bulb globe temperature* is a commonly used index to assess the environmental heat load. This calculation considers the ambient air temperature, the relative humidity, and the degree of radiant heat.

The regulation of this heat load is complex and involves the central nervous system (CNS), thermosensors, and thermoregulatory effectors. The central thermostat activates the effectors that produce peripheral vasodilation and sweating. The skin surface is in effect the radiator and the principal location of heat loss, since skin blood flow can increase 25–30 times over the basal rate. This dramatic increase in skin blood flow, coupled with the maintenance of peripheral vasodilation, efficiently radiates heat. At the same time, there is a compensatory vasoconstriction of the splanchnic and renal beds.

Acclimatization to heat reflects a constellation of physiologic adaptations that permit the body to lose heat more efficiently. This process often requires one to several weeks of exposure and work in a hot environment. During acclimatization, the thermoregulatory set point is altered, and this alteration affects the onset, volume, and content of diaphoresis. The threshold for the initiation of sweating is lowered, and the amount of sweat increases, with a lowered salt concentration. Sweating rates can be 1–2 L/h in acclimated individuals during heat stress. Plasma volume expansion also occurs and improves cutaneous vascular flow. The heart rate lowers, with a higher stroke volume. After the individual leaves the hot environment, improved tolerance to heat stress dissipates rapidly, the plasma volume decreases, and de-acclimatization occurs within weeks.

PREDISPOSING FACTORS AND DIFFERENTIAL DIAGNOSIS

When there is an excessive heat load, unacclimated individuals can develop a variety of heat-related illnesses. Heat waves exacerbate the

mortality rate, particularly among the elderly and poor and among persons lacking adequate nutrition and access to air-conditioned environments. Secondary vascular events, including cerebrovascular accidents and myocardial infarctions, occur at least 10 times more often in conditions of extreme heat.

Exertional heat illness continues to occur when laborers, military personnel, or athletes exercise strenuously in the heat. A variety of common factors predispose to heat illness. In addition to the very young and very old, preadolescents and teenagers are at risk since they may use poor judgment when vigorously exercising in high humidity and heat. Other risk factors include obesity, poor conditioning and lack of acclimatization, and mild dehydration.

Cardiovascular inefficiency is a common feature of heat illness. Any physiologic or pharmacologic impediment to cutaneous perfusion will impair heat loss. Many patients are unaware of the heat risk associated with their medications. Anticholinergic agents impair sweating and blunt the normal cardiovascular response to heat. Phenothiazines also have anticholinergic properties that interfere with the function of the preoptic nucleus of the anterior hypothalamus due to central depletion of dopamine.

Calcium channel blockers, beta blockers, and various stimulants also inhibit sweating by reducing peripheral blood flow. To maintain the mean arterial blood pressure, increased cardiac output must be capable of compensating for progressive dehydration. A variety of stimulants and substances of abuse also increase muscle activity and heat production. Careful consideration of the differential diagnosis is important in the evaluation of a patient for a potential heat-related illness. The clinical setting may suggest other etiologies, such as malignant hyperthermia after general anesthesia or neuroleptic malignant syndrome in a patient taking certain antipsychotic medications. A variety of infectious and endocrine disorders as well as conditions with toxicologic or CNS etiologies may initially mimic heatstroke (Table 479e-1).

MINOR HEAT-EMERGENCY SYNDROMES

Heat edema is characterized by mild swelling of the hands, feet, and ankles during the first few days of significant heat exposure. The principal mechanism involves cutaneous vasodilation and pooling of interstitial fluid in response to heat stress. Heat also increases the secretion of antidiuretic hormone and aldosterone. Systemic causes of edema, including cirrhosis, nephrotic syndrome, and congestive heart failure, can usually be excluded by the history and physical examination. Heat edema generally resolves without treatment in several days. Diuretics are not effective and, in fact, predispose to volume depletion and the development of more serious heat-related illnesses.

Prickly heat (*miliaria rubra*, *lichen tropicus*) is a maculopapular, pruritic, erythematous rash that commonly occurs in clothed areas. Blockage of the sweat pores by debris from macerated stratum corneum causes inflammation in the sweat ducts. As the ducts dilate, they rupture and produce superficial vesicles. The predominant symptom is pruritus. In addition to antihistamines, chlorhexidine may provide some relief. Localized areas may benefit from 1% salicylic acid, with caution taken to avoid salicylate intoxication. Clothing should be clean and loose fitting, and activities or environments that induce diaphoresis should be avoided.

Heat syncope (exercise-associated collapse) can follow endurance exercise or can occur in the elderly. Other common clinical scenarios include prolonged standing while stationary in the heat and sudden standing after prolonged exposure to heat. Heat stress routinely causes relative volume depletion, decreased vasomotor tone, and peripheral vasodilation. The cumulative effect of this decrease in venous return is postural hypotension, especially in nonacclimated elderly individuals. Many of those affected also have comorbidities. Therefore, other cardiovascular, neurologic, and metabolic causes of syncope should be considered. After removal from the heat source, most patients will recover promptly with cooling and rehydration.

Hyperventilation tetany occurs in some individuals when exposure to heat stimulates hyperventilation, producing respiratory alkalosis, paresthesias, and carpopedal spasm. Unlike heat cramps, heat tetany causes very little muscle-compartment pain. Treatment includes

TABLE 479e-1 HEAT-RELATED ILLNESS: PREDISPOSING FACTORS AND DIFFERENTIAL DIAGNOSIS

Illness	Predisposing Factors
Cardiovascular inefficiency	Age extremes Beta/calcium channel blockade Congestive heart failure Dehydration Diuresis Obesity Poor physical fitness
Central nervous system illness	Cerebral hemorrhage Hypothalamic cerebrovascular accident Psychiatric disorders Status epilepticus
Impaired heat loss	Antihistamines Heterocyclic antidepressants Occlusive clothing Skin abnormalities
Endocrine-related illness	Diabetic ketoacidosis Pheochromocytoma Thyroid storm
Excessive heat load	Environmental conditions Exertion Fever Hypermetabolic state Lack of acclimatization
Infectious illness	Cerebral abscess Encephalitis Malaria Meningitis Sepsis syndrome Tetanus Typhoid
Toxicologic illness	Amphetamines Anticholinergic toxidrome Cocaine Dietary supplements Hallucinogens Malignant hyperthermia Neuroleptic malignant syndrome Salicylates Serotonin syndrome Strychnine Sympathomimetics Withdrawal syndromes (ethanol, hypnotics)

providing reassurance, moving the patient out of the heat, and addressing the hyperventilation.

HEAT CRAMPS

Heat cramps (exercise-associated muscle cramps) are intermittent, painful, and involuntary spasmodic contractions of skeletal muscles. They typically occur in an unacclimated individual who is at rest after vigorous exertion in a humid, hot environment. In contrast, cramps that occur in athletes during exercise last longer, are relieved by stretching and massage, and resolve spontaneously.

Of note, not all muscle cramps are related to exercise, and the differential diagnosis includes many other disorders. A variety of medications, myopathies, endocrine disorders, and sickle cell trait are other possible causes.

The typical patient with heat cramps is usually profusely diaphoretic and has been replacing fluid losses with copious water or other

hypotonic fluids. Roofers, firefighters, military personnel, athletes, steel workers, and field workers are commonly affected. Other predisposing factors include insufficient sodium intake before intense activity in the heat and lack of heat acclimatization, resulting in sweat with a high salt concentration.

The precise pathogenesis of heat cramps is unclear but appears to involve a relative deficiency of sodium, potassium, and fluid at the intracellular level. Coupled with copious hypotonic fluid ingestion, large amounts of sodium in the diaphoresis cause hyponatremia and hypochloremia, resulting in muscle cramps due to calcium-dependent muscle relaxation. Total-body depletion of potassium may be observed during the period of heat acclimatization. Rhabdomyolysis is very rare with routine exercise-associated muscle cramps.

Heat cramps that are not accompanied by significant dehydration can be treated with commercially available electrolyte solutions. Although the flavored electrolyte solutions are far more palatable, two 650-mg salt tablets dissolved in 1 quart of water produces a 0.1% saline solution. Individuals should avoid the ingestion of undissolved salt tablets, which are a gastric irritant and may induce vomiting.

HEAT EXHAUSTION

The physiologic hallmarks of heat exhaustion—in contrast to heatstroke—are the maintenance of thermoregulatory control and CNS function. The core temperature is usually elevated but is generally <40.5°C (<105°F). The two physiologic precipitants are water depletion and sodium depletion, which often occur in combination. Laborers, athletes, and elderly individuals exerting themselves in hot environments, without adequate fluid intake, tend to develop *water-depletion heat exhaustion*. Persons working in the heat frequently consume only two-thirds of their net water loss and are voluntarily dehydrated. In contrast, *salt-depletion heat exhaustion* occurs more slowly in unacclimated persons who have been consuming large quantities of hypotonic solutions.

Heat exhaustion is usually a diagnosis of exclusion because of the multitude of nonspecific symptoms. If any signs of heatstroke are present, rapid cooling and crystalloid resuscitation should be initiated immediately during stabilization and evaluation. Mild neurologic and gastrointestinal influenza-like symptoms are common. These symptoms may include headache, vertigo, ataxia, impaired judgment, malaise, dizziness, nausea, and muscle cramps. Orthostatic hypotension and sinus tachycardia develop frequently. More significant CNS impairment suggests heatstroke or other infectious, neurologic, or toxicologic diagnoses.

Hemoconcentration does not always develop, and rapid infusion of isotonic IV fluids should be guided by frequent electrolyte determinations and perfusion requirements. Most cases of heat exhaustion reflect mixed sodium and water depletion. Sodium-depletion heat exhaustion is characterized by hyponatremia and hypochloremia. Hepatic aminotransferases are mildly elevated in both types of heat exhaustion. Urinary sodium and chloride concentrations are usually low.

Some patients with heat exhaustion develop heatstroke after removal from the heat-stress environment. Aggressive cooling of nonresponders is indicated until their core temperature is 39°C (102.2°F). Except in mild cases, free water deficits should be replaced slowly over 24–48 h to avoid a decrease of serum osmolality by >2 mOsm/h.

The disposition of younger, previously healthy heat-exhaustion patients who have no major laboratory abnormalities may include hospital observation and discharge after IV rehydration. Older patients with comorbidities (including cardiovascular disease) or predisposing factors often require inpatient fluid and electrolyte replacement, monitoring, and reassessment.

HEATSTROKE

The clinical manifestations of heatstroke reflect a total loss of thermoregulatory function. Typical vital-sign abnormalities include tachypnea, various tachycardias, hypotension, and a widened pulse pressure. Although there is no single specific diagnostic test, the historical and physical triad of exposure to a heat stress, CNS dysfunction, and a core temperature >40.5°C helps establish the preliminary diagnosis.

TABLE 479e-2 TYPICAL MANIFESTATIONS OF HEATSTROKE

Classic	Exertional
Older patient	Younger patient
Predisposing health factors/medications	Healthy condition
Epidemiology (heat waves)	Sporadic cases
Sedentary	Exercising
Anhidrosis (possible)	Diaphoresis (common)
Central nervous system dysfunction	Myocardial/hepatic injury
Oliguria	Acute renal failure
Coagulopathy (mild)	Disseminated intravascular coagulation
Mild lactic acidosis	Marked lactic acidosis
Mild creatine kinase elevation	Rhabdomyolysis
Normoglycemia/calcemia	Hypoglycemia/calcemia
Normokalemia	Hyperkalemia

The definitive diagnosis should be reserved until the other potential causes of hyperthermia are excluded. Many of the usual laboratory abnormalities seen with heatstroke overlap with other conditions. If the patient's mental status does not improve with cooling, toxicologic screening may be indicated, and cranial CT and spinal fluid analysis can be considered.

The premonitory clinical characteristics may be nonspecific and include weakness, dizziness, disorientation, ataxia, and gastrointestinal or psychiatric symptoms. These prodromal symptoms often resemble heat exhaustion. The sudden onset of heatstroke occurs when the maintenance of adequate perfusion requires peripheral vasoconstriction to stabilize the mean arterial blood pressure. As a result, the cutaneous radiation of heat ceases. At this juncture, the core temperature rises dramatically. Since many patients with heatstroke also meet the criteria for systemic inflammatory response syndrome and have a broad differential diagnosis, rapid cooling is essential during the extensive diagnostic evaluation (Table 479e-1).

There are two forms of heatstroke with significantly different manifestations (Table 479e-2). Classic (epidemic) heatstroke (CHS) usually occurs during long periods of high ambient temperature and humidity, as during summer heat waves. Patients with CHS commonly have chronic diseases that predispose to heat-related illness, and they may have limited access to oral fluids. Heat dissipation mechanisms are overwhelmed by both endogenous heat production and exogenous heat stress. Patients with CHS are often compliant with prescribed medications that can impair tolerance to a heat stress. In many of these dehydrated CHS patients, sweating has ceased and the skin is hot and dry. If cooling is delayed, severe hepatic dysfunction, renal failure, disseminated intravascular coagulation, and fulminant multisystem organ failure may occur. Hepatocytes are very heat sensitive. On presentation, the serum level of aspartate aminotransferase (AST) is routinely elevated. Eventually, levels of both AST and alanine aminotransferase (ALT) often increase to >100 times the normal values. Coagulation studies commonly demonstrate decreased platelets, fibrinogen, and prothrombin. Most patients with CHS require cautious crystalloid resuscitation, electrolyte monitoring, and—in certain refractory cases—consideration of central venous pressure (CVP) measurements. Hypernatremia is secondary to dehydration in CHS. Many patients exhibit significant stress leukocytosis, even in the absence of infection.

Patients with exertional heatstroke (EHS), in contrast to those with CHS, are often young and previously healthy, and their diagnosis is usually more obvious from the history. Athletes, laborers, and military recruits are common victims. Unlike those with CHS, many EHS patients present profusely diaphoretic despite significant dehydration. As a result of muscular exertion, rhabdomyolysis and acute renal failure are more common in EHS. Studies to detect rhabdomyolysis and its complications, including hypocalcemia and hyperphosphatemia, should be considered. Hyponatremia, hypoglycemia, and coagulopathies are

frequent findings. Elevated creatine kinase and lactate dehydrogenase levels also suggest EHS. Oliguria is a common finding. Renal failure can result from direct thermal injury, untreated rhabdomyolysis, or volume depletion. Common urinalysis findings include microscopic hematuria, myoglobinuria, and granular or red cell casts.

With both CHS and EHS, heat-related reversible increases in cardiac biomarker levels are often present. Heatstroke often causes thermal cardiomyopathy. As a result, the CVP may be elevated despite significant dehydration. In addition, the patient often presents with potentially deceptive noncardiogenic pulmonary edema and basilar rales despite being significantly hypovolemic. The electrocardiogram commonly displays a variety of tachyarrhythmias, nonspecific ST-T wave changes, and heat-related ischemia or infarction. Rapid cooling—not the administration of antiarrhythmic medications—is essential.

Above 42°C (107.6° F), heat can rapidly produce direct cellular injury. Thermosensitive enzymes become nonfunctional, and eventually there is irreversible uncoupling of oxidative phosphorylation. The production of heat-shock proteins increases, and cytokines mediate a systemic inflammatory response. The vascular endothelium is also damaged, and this injury activates the coagulation cascade. Significant shunting away from the splanchnic circulation produces gastrointestinal ischemia. Endotoxins further impair normal thermoregulation. As a result, if cooling is delayed, severe hepatic dysfunction, permanent renal failure, disseminated intravascular coagulation, and fulminant multisystem organ failure may occur.

COOLING STRATEGIES

Before cooling is initiated, endotracheal intubation, CVP determination, and continuous core-temperature monitoring should be considered. Hypoglycemia is a frequent finding and can be addressed by glucose infusion. Since peripheral vasoconstriction delays heat dissipation, repeated administration of discrete boluses of isotonic crystalloid for hypotension is preferable to the administration of α -adrenergic agonists.

Evaporative cooling is usually the most practical and effective technique. Rapid cooling is essential in both CHS and EHS, and an immediate improvement in vital signs and mental status may prove valuable for diagnostic purposes. Cool water (15°C [60° F]) is sprayed on the exposed skin while fans direct continuous airflow over the moistened skin. Cold packs applied to the axillae and groin are a useful cooling adjunct. If cardiac electrodes will not adhere, they can be applied to the patient's back. To avoid “overshoot hypothermia,” active cooling should be terminated at ~38°–39°C (100.4°–102.2°F).

Immersion cooling in cold water is an alternative option in EHS but induces peripheral vasoconstriction and intense shivering. This technique presents significant monitoring and resuscitation challenges in most clinical settings. The safety of immersion cooling is best established for young, previously healthy patients with EHS (but not for those with CHS).

Cooling with commercially available cooling blankets should not be the sole technique used, since the rate of cooling is far too slow. Other methods are less efficacious and rarely indicated, such as IV infusion of cold fluids and cold irrigation of the bladder or gastrointestinal tract. Cold thoracic and peritoneal lavage are efficient maneuvers but are invasive and rarely necessary. Cardiopulmonary bypass provides fast and effective cooling but is labor intensive and is rarely available on a stat basis.

RESUSCITATION

Aspiration and seizures commonly occur in heatstroke, and endotracheal intubation is usually necessary. The metabolic demands are high, and supplemental oxygenation is essential due to hypoxemia induced by thermal stress and pulmonary dysfunction. Pneumonitis, pulmonary infarction, hemorrhage, edema, and acute respiratory distress syndrome occur frequently in heatstroke patients.

The circulatory fluid requirements, particularly in CHS, may be deceptively modest. Aggressive cooling and modest volume repletion usually elevate the CVP to 12–14 mmHg. The reading, however,

479e-4 may be deceptive. Many patients present with a thermally induced hyperdynamic circulation accompanied by a high cardiac index, low peripheral vascular resistance, and an elevated CVP caused by right-sided heart failure. Rarely, wedge pressures measured via a pulmonary artery catheter may be necessary to guide resuscitation. In contrast, most patients with EHS require far more zealous isotonic crystalloid resuscitation.

The hypotension that is initially common among patients with heatstroke results from both dehydration and high-output cardiac failure caused by peripheral vasodilation. Inotropes causing α -adrenergic stimulation (e.g., norepinephrine) can impede cooling by causing significant vasoconstriction. Vasoactive catecholamines such as dopamine or dobutamine may be necessary if the cardiac output remains depressed despite an elevated CVP, particularly in patients with a hyperdynamic circulation.

A wide variety of tachyarrhythmias are routinely observed on presentation and usually resolve during cooling. The administration of atrial or ventricular antiarrhythmic medications is rarely indicated during cooling. Anticholinergic medications (including atropine) that inhibit sweating should not be given, and electrical cardioversion of the hyperthermic myocardium should be avoided except when there is ventricular fibrillation.

Significant shivering, discomfort, or extreme agitation is preferably mitigated with short-acting benzodiazepines, which are ideal due to their renal clearance. On the other hand, chlorpromazine may lower the seizure threshold, has anticholinergic properties, and can exacerbate the hypotension or cause neuroleptic malignant syndrome. Because of likely hepatic dysfunction, barbiturates should be avoided and seizures should be treated with benzodiazepines.

Coagulopathies more commonly occur after the first day of illness. After cooling, the patient should be monitored for laboratory signs of disseminated intravascular coagulation, and replacement therapy with fresh-frozen plasma and platelets should be considered.

There is no therapeutic role for antipyretics in the control of environmentally induced hyperthermia; these drugs block the actions of pyrogens at hypothalamic receptor sites. Salicylates can further uncouple oxidative phosphorylation in heatstroke and exacerbate coagulopathies. Acetaminophen may further stress hepatic function. The safety and efficacy of other medications, including dantrolene and aminocaproic acid, are not established.

DISPOSITION

Most patients with minor heat-emergency syndromes (including heat edema, heat syncope, and heat cramps) require only stabilization and treatment with outpatient follow-up. Although there are no decision rules to guide disposition choices in heat exhaustion, many of these patients have multiple predisposing factors and comorbidities that will require prolonged observation or hospital admission.

Essentially all patients with actual heatstroke require admission to a monitored setting, and most require intensive care. Many of these patients also require prolonged tracheal intubation, invasive hemodynamic monitoring, and support for various degrees of multiorgan dysfunction syndrome. The prognosis worsens if the initial core temperature exceeds 42°C (107.6°F) or if there was a prolonged period during which the core temperature exceeded this level. Other features of a negative prognosis include acute renal failure, massively elevated liver enzymes, and significant hyperkalemia. As expected, the number of dysfunctional organ systems also correlates directly with mortality risk.

This page intentionally left blank

APPENDIX: Laboratory Values of Clinical Importance

Alexander Kratz, Michael A. Pesce, Robert C. Basner,
Andrew J. Einstein

This Appendix contains tables of reference values for common laboratory tests. A variety of factors can influence reference values. Such variables include the population studied, the duration and means of specimen transport, laboratory methods and instrumentation, and even the type of container used for the collection of the specimen. The reference or “normal” ranges given in this appendix may therefore not be appropriate for all laboratories, and these values should only

be used as general guidelines. Whenever possible, reference values provided by the laboratory performing the testing should be used in the interpretation of laboratory data. Values supplied in this Appendix reflect typical reference ranges in nonpregnant adults. Pediatric reference ranges and values in pregnant patients may vary significantly from the data presented in the Appendix.

In preparing the Appendix, the authors have taken into account the fact that the system of international units (SI, système international d’unités) is used in most countries and in some medical journals. However, clinical laboratories may continue to report values in “traditional” or conventional units. Therefore, both systems are provided in the Appendix. The dual system is also used in the text except for those instances in which the numbers remain the same and only the terminology is changed (mmol/L for meq/L or IU/L for mIU/mL), when only the SI units are given.

TABLE 1 HEMATOLOGY AND COAGULATION

Analyte	Specimen	SI Units	Conventional Units
Activated clotting time	WB	70–180 s	70–180 s
Activated protein C resistance (factor V Leiden)	P	Not applicable	Ratio >2.1
ADAMTS13 activity	P	≥0.67	≥67%
ADAMTS13 inhibitor activity	P	Not applicable	≤0.4 U
ADAMTS13 antibody	P	Not applicable	≤18 U
Alpha ₂ antiplasmin	P	0.87–1.55	87–155%
Antiphospholipid antibody panel			
PTT-LA (lupus anticoagulant screen)	P	Negative	Negative
Platelet neutralization procedure	P	Negative	Negative
Dilute viper venom screen	P	Negative	Negative
Anticardiolipin antibody	S		
IgG		0–15 arbitrary units	0–15 GPL
IgM		0–15 arbitrary units	0–15 MPL
Beta-2 glycoprotein 1 antibodies	S		
IgG		0–20 arbitrary units	0–20 SGU
IgM		0–20 arbitrary units	0–20 SMU
Antithrombin III	P		
Antigenic		220–390 mg/L	22–39 mg/dL
Functional		0.7–1.30 U/L	70–130 %
Anti-Xa assay (heparin assay)	P		
Unfractionated heparin		0.3–0.7 kIU/L	0.3–0.7 IU/mL
Low-molecular-weight heparin		0.5–1.0 kIU/L	0.5–1.0 IU/mL
Danaparoid (Orgaran)		0.5–0.8 kIU/L	0.5–0.8 IU/mL
Autohemolysis test	WB	0.004–0.045	0.4–4.50%
Autohemolysis test with glucose	WB	0.003–0.007	0.3–0.7%
Bleeding time (adult)		<7.1 min	<7.1 min
C4 binding protein	P	305–695 mg/L	30.5–69.5 mg/dL
	S	275–604 mg/L	27.5–60.4 mg/dL
Clot retraction	WB	0.50–1.00/2 h	50–100%/2 h
Cryofibrinogen	P	Negative	Negative
D-dimer	P	220–740 ng/mL FEU	220–740 ng/mL FEU
Differential blood count	WB		
Relative counts:			
Neutrophils		0.40–0.70	40–70%
Bands		0.0–0.05	0–5%
Lymphocytes		0.20–0.50	20–50%
Monocytes		0.04–0.08	4–8%
Eosinophils		0.0–0.6	0–6%
Basophils		0.0–0.02	0–2%

(Continued)

TABLE 1 HEMATOLOGY AND COAGULATION (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Absolute counts:			
Neutrophils		1.42–6.34 × 10 ⁹ /L	1420–6340/mm ³
Bands		0–0.45 × 10 ⁹ /L	0–450/mm ³
Lymphocytes		0.71–4.53 × 10 ⁹ /L	710–4530/mm ³
Monocytes		0.14–0.72 × 10 ⁹ /L	140–720/mm ³
Eosinophils		0–0.54 × 10 ⁹ /L	0–540/mm ³
Basophils		0–0.18 × 10 ⁹ /L	0–180/mm ³
Erythrocyte count	WB		
Adult males		4.30–5.60 × 10 ¹² /L	4.30–5.60 × 10 ⁶ /mm ³
Adult females		4.00–5.20 × 10 ¹² /L	4.00–5.20 × 10 ⁶ /mm ³
Erythrocyte life span	WB		
Normal survival		120 days	120 days
Chromium labeled, half-life (t _{1/2})		25–35 days	25–35 days
Erythrocyte sedimentation rate	WB		
Females		0–20 mm/h	0–20 mm/h
Males		0–15 mm/h	0–15 mm/h
Euglobulin lysis time	P	7200–14400 s	120–240 min
Factor II, prothrombin	P	0.50–1.50	50–150%
Factor V	P	0.50–1.50	50–150%
Factor VII	P	0.50–1.50	50–150%
Factor VIII	P	0.50–1.50	50–150%
Factor IX	P	0.50–1.50	50–150%
Factor X	P	0.50–1.50	50–150%
Factor XI	P	0.50–1.50	50–150%
Factor XII	P	0.50–1.50	50–150%
Factor XIII screen	P	Not applicable	Present
Factor inhibitor assay	P	<0.5 Bethesda Units	<0.5 Bethesda Units
Fibrin(ogen) degradation products	P	0–1 mg/L	0–1 µg/mL
Fibrinogen	P	2.33–4.96 g/L	233–496 mg/dL
Glucose-6-phosphate dehydrogenase (erythrocyte)	WB	<2400 s	<40 min
Ham's test (acid serum)	WB	Negative	Negative
Hematocrit	WB		
Adult males		0.388–0.464	38.8–46.4
Adult females		0.354–0.444	35.4–44.4
Hemoglobin			
Plasma	P	6–50 mg/L	0.6–5.0 mg/dL
Whole blood:	WB		
Adult males		133–162 g/L	13.3–16.2 g/dL
Adult females		120–158 g/L	12.0–15.8 g/dL
Hemoglobin electrophoresis	WB		
Hemoglobin A		0.95–0.98	95–98%
Hemoglobin A ₂		0.015–0.031	1.5–3.1%
Hemoglobin F		0–0.02	0–2.0%
Hemoglobins other than A, A ₂ , or F		Absent	Absent
Heparin-induced thrombocytopenia antibody	P	Negative	Negative
Immature platelet fraction (IPF)	WB	0.011–0.061	1.1–6.1%
Joint fluid crystal	JF	Not applicable	No crystals seen
Joint fluid mucin	JF	Not applicable	Only type I mucin present
Leukocytes			
Alkaline phosphatase (LAP)	WB	0.2–1.6 µkat/L	13–100 µ/L
Count (WBC)	WB	3.54–9.06 × 10 ⁹ /L	3.54–9.06 × 10 ³ /mm ³
Mean corpuscular hemoglobin (MCH)	WB	26.7–31.9 pg/cell	26.7–31.9 pg/cell
Mean corpuscular hemoglobin concentration (MCHC)	WB	323–359 g/L	32.3–35.9 g/dL
Mean corpuscular hemoglobin of reticulocytes (CH)	WB	24–36 pg	24–36 pg
Mean corpuscular volume (MCV)	WB	79–93.3 fL	79–93.3 µm ³
Mean platelet volume (MPV)	WB	9.00–12.95 fL	9.00–12.95

(Continued)

TABLE 1 HEMATOLOGY AND COAGULATION (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Osmotic fragility of erythrocytes	WB		
Direct		0.0035–0.0045	0.35–0.45%
Indirect		0.0030–0.0065	0.30–0.65%
Partial thromboplastin time, activated	P	26.3–39.4 s	26.3–39.4 s
Plasminogen	P		
Antigen		84–140 mg/L	8.4–14.0 mg/dL
Functional		0.70–1.30	70–130%
Plasminogen activator inhibitor 1	P	4–43 µg/L	4–43 ng/mL
Platelet aggregation	PRP	Not applicable	>65% aggregation in response to adenosine diphosphate, epinephrine, collagen, ristocetin, and arachidonic acid
Platelet count	WB	165–415 × 10 ⁹ /L	165–415 × 10 ³ /mm ³
Platelet, mean volume	WB	6.4–11 fL	6.4–11.0 µm ³
Prekallikrein assay	P	0.50–1.5	50–150%
Prekallikrein screen	P		No deficiency detected
Protein C	P		
Total antigen		0.70–1.40	70–140%
Functional		0.70–1.30	70–130%
Protein S	P		
Total antigen		0.70–1.40	70–140%
Functional		0.65–1.40	65–140%
Free antigen		0.70–1.40	70–140%
Prothrombin gene mutation G20210A	WB	Not applicable	Not present
Prothrombin time	P	12.7–15.4 s	12.7–15.4 s
Protoporphyrin, free erythrocyte	WB	0.28–0.64 µmol/L of red blood cells	16–36 µg/dL of red blood cells
Red cell distribution width	WB	<0.145	<14.5%
Reptilase time	P	16–23.6 s	16–23.6 s
Reticulocyte count	WB		
Adult males		0.008–0.023 red cells	0.8–2.3% red cells
Adult females		0.008–0.020 red cells	0.8–2.0% red cells
Reticulocyte hemoglobin content	WB	>26 pg/cell	>26 pg/cell
Ristocetin cofactor (functional von Willebrand factor)	P		
Blood group O		0.75 mean of normal	75% mean of normal
Blood group A		1.05 mean of normal	105% mean of normal
Blood group B		1.15 mean of normal	115% mean of normal
Blood group AB		1.25 mean of normal	125% mean of normal
Serotonin release assay	S	<0.2 release	<20% release
Sickle cell test	WB	Negative	Negative
Sucrose hemolysis	WB	<0.1	<10% hemolysis
Thrombin time	P	15.3–18.5 s	15.3–18.5 s
Thrombin-antithrombin (TAT) complex	P	<4 µg/L	<4 ng/mL
Total eosinophils	WB	150–300 × 10 ⁶ /L	150–300/mm ³
Transferrin receptor	S, P	9.6–29.6 nmol/L	9.6–29.6 nmol/L
Viscosity			
Plasma	P	1.7–2.1	1.7–2.1
Serum	S	1.4–1.8	1.4–1.8
von Willebrand factor (VWF) antigen (factor VIII:R antigen)	P		
Blood group O		0.75 mean of normal	75% mean of normal
Blood group A		1.05 mean of normal	105% mean of normal
Blood group B		1.15 mean of normal	115% mean of normal
Blood group AB		1.25 mean of normal	125% mean of normal
von Willebrand factor collagen binding	P	590–2490 units/L	59–249 units/dL
von Willebrand factor multimers	P	Normal distribution	Normal distribution
White blood cells: see “Leukocytes”			

Abbreviations: JF, joint fluid; P, plasma; PRP, platelet-rich plasma; S, serum; WB, whole blood.

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY

Analyte	Specimen	SI Units	Conventional Units
Acetoacetate	P	49–294 $\mu\text{mol/L}$	0.5–3.0 mg/dL
Adrenocorticotropin (ACTH)	P	1.3–16.7 pmol/L	6.0–76.0 pg/mL
Alanine aminotransferase (ALT, SGPT)	S	0.12–0.70 $\mu\text{kat/L}$	7–41 U/L
Albumin	S	40–50 g/L	4.0–5.0 mg/dL
Aldolase	S	26–138 nkat/L	1.5–8.1 U/L
Aldosterone (adult)			
Supine, normal sodium diet	S, P	<443 pmol/L	<16 ng/dL
Upright, normal.	S, P	111–858 pmol/L	4–31 ng/dL
Alpha fetoprotein (adult)	S	0–8.5 $\mu\text{g/L}$	0–8.5 ng/mL
Alpha-1-acid glycoprotein	S	0.50–1.2 g/L	50–120 mg/dL
Alpha ₁ antitrypsin	S	1.0–2.0 g/L	100–200 mg/dL
Ammonia, as NH_3	P	11–35 $\mu\text{mol/L}$	19–60 $\mu\text{g/dL}$
Amylase (method dependent)	S	0.34–1.6 $\mu\text{kat/L}$	20–96 U/L
Androstenedione (adult)	S		
Males		0.81–3.1 nmol/L	23–89 ng/dL
Females			
Premenopausal		0.91–7.5 nmol/L	26–214 ng/dL
Postmenopausal		0.46–2.9 nmol/L	13–82 ng/dL
Angiotensin-converting enzyme (ACE)	S	0.15–1.1 $\mu\text{kat/L}$	9–67 U/L
Anion gap	S	7–16 mmol/L	7–16 mmol/L
Apolipoprotein A-1	S		
Male		0.94–1.78 g/L	94–178 mg/dL
Female		1.01–1.99 g/L	101–199 mg/dL
Apolipoprotein B	S		
Male		0.55–1.40 g/L	55–140 mg/dL
Female		0.55–1.25 g/L	55–125 mg/dL
Arterial blood gases	WB		
(HCO_3^-)		22–30 mmol/L	22–30 meq/L
Pco_2 (Sea Level, Fio_2 0.21)		4.7–6.0 kPa	35–45 mmHg
pH		7.35–7.45	7.35–7.45
Po_2 (Sea Level, Fio_2 0.21, age related)		8.9–13.8 kPa	67–104 mmHg
Carboxyhemoglobin and methemoglobin at pH 7.40 and 37°C		≤ 0.01	$\leq 1\%$
Aspartate aminotransferase (AST, SGOT)	S	0.20–0.65 $\mu\text{kat/L}$	12–38 U/L
Autoantibodies	S		
Anti-centromere antibody IgG		≤ 29 AU/mL	≤ 29 AU/mL
Anti-double-strand (native) DNA		<25 IU/L	<25 IU/L
Anti-glomerular basement membrane antibodies			
Qualitative IgG, IgA		Negative	Negative
Quantitative IgG antibody		≤ 19 AU/mL	≤ 19 AU/mL
Anti-histone antibodies		<1.0 U	<1.0 U
Anti-Jo-1 antibody		≤ 29 AU/mL	≤ 29 AU/mL
Anti-mitochondrial antibody		Not applicable	<20 Units
Anti-neutrophil cytoplasmic autoantibodies		Not applicable	<1:20
Serine proteinase 3 antibodies		≤ 19 AU/mL	≤ 19 AU/mL
Myeloperoxidase antibodies		≤ 19 AU/mL	≤ 19 AU/mL
Antinuclear antibody		Not applicable	Negative at 1:40
Anti-parietal cell antibody		Not applicable	None detected
Anti-RNP antibody		Not applicable	<1.0 U
Anti-Scl 70 antibody		Not applicable	<1.0 U
Anti-Smith antibody		Not applicable	<1.0 U
Anti-smooth muscle antibody		Not applicable	<1.0 U
Anti-SSA antibody		Not applicable	<1.0 U
Anti-SSB antibody		Not applicable	Negative
Anti-thyroglobulin antibody		<40 KIU/mL	<40 IU/mL
Anti-thyroid peroxidase antibody		<35 KIU/L	<35 IU/L

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
B-type natriuretic peptide (BNP)	P	Age and gender specific: <100 ng/L	Age and gender specific: <100 pg/mL
Bence Jones protein, serum qualitative	S	Not applicable	None detected
Bence Jones protein, serum quantitative	S		
Free kappa		3.3–19.4 mg/L	0.33–1.94 mg/dL
Free lambda		5.7–26.3 mg/L	0.57–2.63 mg/dL
K/L ratio		0.26–1.65	0.26–1.65
Beta-2-microglobulin	S	1.1–2.4 mg/L	1.1–2.4 mg/L
Bile acids	S		
Cholic acid		0–1.9 µmol/L	0–1.9 µmol/L
Chenodeoxycholic acid		0–3.4 µmol/L	0–3.4 µmol/L
Deoxycholic acid		0–2.5 µmol/L	0–2.5 µmol/L
Ursodeoxycholic acid		0–1.0 µmol/L	0–1.0 µmol/L
Total		0–7.0 µmol/L	0–7.0 µmol/L
Bilirubin	S		
Total		5.1–22 µmol/L	0.3–1.3 mg/dL
Direct		1.7–6.8 µmol/L	0.1–0.4 mg/dL
Indirect		3.4–15.2 µmol/L	0.2–0.9 mg/dL
C peptide	S	0.27–1.19 nmol/L	0.8–3.5 ng/mL
C1-esterase-inhibitor protein	S	210–390 mg/L	21–39 mg/dL
CA 125	S	<35 kU/L	<35 U/mL
CA 19-9	S	<37 kU/L	<37 U/mL
CA 15-3	S	<33 kU/L	<33 U/mL
CA 27-29	S	0–40 kU/L	0–40 U/mL
Calcitonin	S		
Male		0–7.5 ng/L	0–7.5 pg/mL
Female		0–5.1 ng/L	0–5.1 pg/mL
Calcium	S	2.2–2.6 mmol/L	8.7–10.2 mg/dL
Calcium, ionized	WB	1.12–1.32 mmol/L	4.5–5.3 mg/dL
Carbon dioxide content (TCO ₂)	P (sea level)	22–30 mmol/L	22–30 meq/L
Carboxyhemoglobin (carbon monoxide content)	WB		
Nonsmokers (in a nonsmoking environment)		0.0–0.025	0–2.5% of total hemoglobin (Hgb) value
Smokers		0.04–0.09	4–9% of total Hgb value
Loss of consciousness and death		>0.50	>50% of total Hgb value
Carcinoembryonic antigen (CEA)	S		
Nonsmokers		0.0–3.0 µg/L	0.0–3.0 ng/mL
Smokers		0.0–5.0 µg/L	0.0–5.0 ng/mL
Ceruloplasmin	S	250–630 mg/L	25–63 mg/dL
Chloride	S	102–109 mmol/L	102–109 meq/L
Cholesterol (LCL, Total, HDL): Ranges depend on individual patient factors; see 2013 ACC/AHA Guideline on the Treatment of Blood Cholesterol			
Cholinesterase	S	5–12 kU/L	5–12 U/mL
Chromogranin A	S	0–95 µg/L	0–95 ng/mL
Complement	S		
C3		0.83–1.77 g/L	83–177 mg/dL
C4		0.16–0.47 g/L	16–47 mg/dL
Complement total		60–144 CAE units	60–144 CAE units
Cortisol			
Fasting, 8 A.M.–12 noon	S	138–690 nmol/L	5–25 µg/dL
12 noon–8 P.M.		138–414 nmol/L	5–15 µg/dL
8 P.M.–8 A.M.		0–276 nmol/L	0–10 µg/dL
C-reactive protein	S	<10 mg/L	<10 mg/L
C-reactive protein, high sensitivity	S	Cardiac risk	Cardiac risk
		Low: <1.0 mg/L	Low: <1.0 mg/L
		Average: 1.0–3.0 mg/L	Average: 1.0–3.0 mg/L
		High: >3.0 mg/L	High: >3.0 mg/L

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Creatine kinase (total)	S		
Females		0.66–4.0 μ kat/L	39–238 U/L
Males		0.87–5.0 μ kat/L	51–294 U/L
Creatine kinase-MB	S		
Mass		0.0–5.5 μ g/L	0.0–5.5 ng/mL
Fraction of total activity (by electrophoresis)		0–0.04	0–4.0%
Creatinine	S		
Female		44–80 μ mol/L	0.5–0.9 mg/dL
Male		53–106 μ mol/L	0.6–1.2 mg/dL
Cryoglobulins	S	Not applicable	None detected
Cyclic citrullinated peptide (CCP) antibody (IgG)	S	Negative: <20 Units Weak positive: 20–39 Units Moderate positive: 40–59 Units Strong positive: \geq 60 Units	Negative: <20 Units Weak positive: 20–39 Units Moderate positive: 40–59 Units Strong positive: \geq 60 Units
Cystatin C	S	0.5–1.0 mg/L	0.5–1.0 mg/L
Deamidated gliadin peptide (DGP) antibody, IgA	S		
Negative		\leq 19 Units	\leq 19 Units
Weak positive		20–30 Units	20–30 Units
Positive		\geq 31 Units	\geq 31 Units
Deamidated gliadin peptide (DGP) antibody, IgG	S		
Negative		\leq 19 Units	\leq 19 Units
Weak positive		20–30 Units	20–30 Units
Positive		\geq 31 Units	\geq 31 Units
Dehydroepiandrosterone (DHEA) (adult)			
Male	S	6.2–43.4 nmol/L	180–1250 ng/dL
Female		4.5–34.0 nmol/L	130–980 ng/dL
Dehydroepiandrosterone (DHEA) sulfate	S		
Male (adult)		100–6190 μ g/L	10–619 μ g/dL
Female (adult, premenopausal)		120–5350 μ g/L	12–535 μ g/dL
Female (adult, postmenopausal)		300–2600 μ g/L	30–260 μ g/dL
11-Deoxycortisol (adult) (compound S)	S	0.34–4.56 nmol/L	12–158 ng/dL
Dihydrotestosterone			
Male	S, P	1.03–2.92 nmol/L	30–85 ng/dL
Female		0.14–0.76 nmol/L	4–22 ng/dL
Dopamine	P	0–130 pmol/L	0–20 pg/mL
Endomysial antibody, IgA	S	<1:10	<1:10
Endomysial antibody, IgG	S	<1:10	<1:10
Epinephrine	P		
Supine (30 min)		<273 pmol/L	<50 pg/mL
Sitting		<328 pmol/L	<60 pg/mL
Standing (30 min)		<491 pmol/L	<90 pg/mL
Erythropoietin	S	4–27 U/L	4–27 U/L
Estradiol	S, P		
Female			
Menstruating:			
Follicular phase		74–532 pmol/L	<20–145 pg/mL
Midcycle peak		411–1626 pmol/L	112–443 pg/mL
Luteal phase		74–885 pmol/L	<20–241 pg/mL
Postmenopausal		217 pmol/L	<59 pg/mL
Male		74 pmol/L	<20 pg/mL
Estrone	S, P		
Female			
Menstruating:			
Follicular phase		<555 pmol/L	<150 pg/mL
Luteal phase		<740 pmol/L	<200 pg/mL
Postmenopausal		11–118 pmol/L	3–32 pg/mL
Male		33–133 pmol/L	9–36 pg/mL

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Fatty acids, free (nonesterified)	P	0.1–0.6 mmol/L	2.8–16.8 mg/dL
Ferritin	S		
Female		10–150 µg/L	10–150 ng/mL
Male		29–248 µg/L	29–248 ng/mL
Follicle-stimulating hormone (FSH)	S, P		
Female			
Menstruating			
Follicular phase		3.0–20.0 IU/L	3.0–20.0 mIU/mL
Ovulatory phase		9.0–26.0 IU/L	9.0–26.0 mIU/mL
Luteal phase		1.0–12.0 IU/L	1.0–12.0 mIU/mL
Postmenopausal		18.0–153.0 IU/L	18.0–153.0 mIU/mL
Male		1.0–12.0 IU/L	1.0–12.0 mIU/mL
Fructosamine	S	<285 µmol/L	<285 µmol/L
Galectin-3	S		
Low risk		≤17.8 µg/L	≤17.8 ng/mL
Intermediate risk		17.9–25.9 µg/L	17.9–25.9 ng/mL
Higher risk		>25.9 µg/L	>25.9 ng/mL
Gamma glutamyltransferase	S	0.15–0.99 µkat/L	9–58 U/L
Gastrin	S	<100 ng/L	<100 pg/mL
Glucagon	P	40–130 ng/L	40–130 pg/mL
Glucose	WB	3.6–5.3 mmol/L	65–95 mg/dL
Glucose (fasting)	P		
Normal		4.2–5.6 mmol/L	75–100 mg/dL
Increased risk for diabetes		5.6–6.9 mmol/L	100–125 mg/dL
Diabetes mellitus		Fasting ≥7.0 mmol/L A 2-h level of ≥11.1 mmol/L during an oral glucose tolerance test A random glucose level of ≥11.1 mmol/L in patients with symptoms of hyperglycemia	Fasting ≥126 mg/dL A 2-h level of ≥200 mg/dL during an oral glucose tolerance test A random glucose level of ≥200 mg/dL in patients with symptoms of hyperglycemia
Growth hormone	S	0–5 µg/L	0–5 ng/mL
Hemoglobin A _{1c}	WB	0.04–0.06 Hgb fraction	4.0–5.6%
Prediabetes		0.057–0.064 Hgb fraction	5.7–6.4%
Diabetes mellitus		A hemoglobin A _{1c} level of ≥0.065 Hgb fraction as suggested by the American Diabetes Association	A hemoglobin A _{1c} level of ≥6.5% as suggested by the American Diabetes Association
Hemoglobin A _{1c} with estimated average glucose (eAG)	WB	eAg mmol/L = 1.59 × HbA _{1c} – 2.59	eAg (mg/dL) = 28.7 × HbA _{1c} – 46.7
Homocysteine	P	4.4–10.8 µmol/L	4.4–10.8 µmol/L
Human chorionic gonadotropin (HCG)	S		
Nonpregnant female		<5 IU/L	<5 mIU/mL
1–2 weeks postconception		9–130 IU/L	9–130 mIU/mL
2–3 weeks postconception		75–2600 IU/L	75–2600 mIU/mL
3–4 weeks postconception		850–20,800 IU/L	850–20,800 mIU/mL
4–5 weeks postconception		4000–100,200 IU/L	4000–100,200 mIU/mL
5–10 weeks postconception		11,500–289,000 IU/L	11,500–289,000 mIU/mL
10–14 weeks postconception		18,300–137,000 IU/L	18,300–137,000 mIU/mL
Second trimester		1400–53,000 IU/L	1400–53,000 mIU/mL
Third trimester		940–60,000 IU/L	940–60,000 mIU/mL
Human epididymis protein 4 (HE-4)	S	0–150 pmol/L	0–150 pmol/L
β-Hydroxybutyrate	P	60–170 µmol/L	0.6–1.8 mg/dL
17-Hydroxyprogesterone (adult)	S		
Male		<4.17 nmol/L	<139 ng/dL
Female			
Follicular phase		0.45–2.1 nmol/L	15–70 ng/dL
Luteal phase		1.05–8.7 nmol/L	35–290 ng/dL
Immunofixation	S	Not applicable	No bands detected

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Immunoglobulin, quantitation (adult)			
IgA	S	0.70–3.50 g/L	70–350 mg/dL
IgD	S	0–140 mg/L	0–14 mg/dL
IgE	S	1–87 KIU/L	1–87 IU/mL
IgG	S	7.0–17.0 g/L	700–1700 mg/dL
IgG ₁	S	2.7–17.4 g/L	270–1740 mg/dL
IgG ₂	S	0.3–6.3 g/L	30–630 mg/dL
IgG ₃	S	0.13–3.2 g/L	13–320 mg/dL
IgG ₄	S	0.11–6.2 g/L	11–620 mg/dL
IgM	S	0.50–3.0 g/L	50–300 mg/dL
Inhibin A			
Males		≤2.0 ng/L	≤2.0 pg/mL
Females			
Early follicular phase		1.8–17.3 ng/L	1.8–17.3 pg/mL
Mid follicular phase		3.5–31.7 ng/L	3.5–17.3 pg/mL
Late follicular phase		9.8–90.3 ng/L	9.8–90.3 pg/mL
Midcycle		16.9–91.8 ng/L	16.9–91.8 pg/mL
Early luteal phase		16.1–97.5 ng/L	16.1–97.5 pg/mL
Mid luteal phase		3.9–87.7 ng/L	3.9–87.7 pg/mL
Late luteal phase		2.7–47.1 ng/L	2.7–47.1 pg/mL
Postmenopausal		<1.0–2.1 ng/L	<1.0–2.1 pg/mL
Insulin	S, P	14.35–143.5 pmol/L	2–20 μU/mL
Iron	S	7–25 μmol/L	41–141 μg/dL
Iron-binding capacity	S	45–73 μmol/L	251–406 μg/dL
Iron-binding capacity saturation	S	0.16–0.35	16–35%
Ischemia modified albumin	S	<85 KU/L	<85 U/mL
Joint fluid crystal	JF	Not applicable	No crystals seen
Joint fluid mucin	JF	Not applicable	Only type I mucin present
Ketone (acetone)	S	Negative	Negative
Lactate	P, arterial	0.5–1.6 mmol/L	4.5–14.4 mg/dL
	P, venous	0.5–2.2 mmol/L	4.5–19.8 mg/dL
Lactate dehydrogenase	S	2.0–3.8 μkat/L	115–221 U/L
Lamellar body count			
Immature	AMF	<15,000/μL	<15,000/μL
Indeterminate		15,000–50,000/μL	15,000–50,000/μL
Mature		>50,000/μL	>50,000/μL
Lecithin/sphingomyelin (L/S) ratio			
Immature	AMF	≤1.5	≤1.5
Transitional		1.5–1.9	1.5–1.9
Mature		2.0–2.5 or greater	2.0–2.5 or greater
Lipase	S	0.51–0.73 μkat/L	3–43 U/L
Lipoprotein (a)	S	0–300 mg/L	0–30 mg/dL
Lipoprotein associated phospholipase A2	S, P	0–234 μg/L	0–234 ng/mL
Luteinizing hormone (LH)			
Female			
Menstruating			
Follicular phase		2.0–15.0 U/L	2.0–15.0 mIU/mL
Ovulatory phase		22.0–105.0 U/L	22.0–105.0 mIU/mL
Luteal phase		0.6–19.0 U/L	0.6–19.0 mIU/mL
Postmenopausal		16.0–64.0 U/L	16.0–64.0 mIU/mL
Male			
		2.0–12.0 U/L	2.0–12.0 mIU/mL
Magnesium	S	0.62–0.95 mmol/L	1.5–2.3 mg/dL
Metanephrine	P	<0.5 nmol/L	<100 pg/mL
Methemoglobin	WB	0.0–0.01	0–1% of total Hgb value

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Myoglobin	S		
Male		20–71 µg/L	20–71 µg/L
Female		25–58 µg/L	25–58 µg/L
Norepinephrine	P		
Supine (30 min)		650–2423 pmol/L	110–410 pg/mL
Sitting		709–4019 pmol/L	120–680 pg/mL
Standing (30 min)		739–4137 pmol/L	125–700 pg/mL
N-telopeptide (cross-linked), NTx	S		
Female, premenopausal		6.2–19.0 nmol BCE	6.2–19.0 nmol BCE
Male		5.4–24.2 nmol BCE	5.4–24.2 nmol BCE
BCE = bone collagen equivalent			
NT-proBNP	S, P	<125 ng/L up to 75 years <450 ng/L >75 years	<125 pg/mL up to 75 years <450 pg/mL >75 years
5' Nucleotidase	S	0.00–0.19 µkat/L	0–11 U/L
Osmolality	P	275–295 mOsmol/kg serum water	275–295 mOsmol/kg serum water
Osteocalcin	S	11–50 µg/L	11–50 ng/mL
Oxygen content (age and gender related)	WB		
Arterial (sea level)		17–21 mL/dL	17–21 vol%
Venous (sea level)		10–16 mL/dL	10–16 vol%
Oxygen saturation (sea level)	WB	Fraction:	Percent:
Arterial		0.91–1.0	91–100%
Venous, arm		0.60–0.85	60–85%
Parathyroid hormone (intact)	S	8–51 ng/L	8–51 pg/mL
Phosphatase, alkaline	S	0.56–1.63 µkat/L	33–96 U/L
Phosphatase, alkaline bone	S		
Male		≤20 µg/L	≤20 ng/mL
Female			
Premenopausal		≤14 µg/L	≤14 ng/mL
Postmenopausal		≤22 µg/L	≤22 ng/mL
Phosphorus, inorganic	S	0.81–1.4 mmol/L	2.5–4.3 mg/dL
Potassium	S	3.5–5.0 mmol/L	3.5–5.0 meq/L
Prealbumin (transthyretin)	S	170–340 mg/L	17–34 mg/dL
Procalcitonin	S	<0.1 µg/L	<0.1 ng/mL
Progesterone	S, P		
Female: Follicular		<3.18 nmol/L	<1.0 ng/mL
Midluteal		9.54–63.6 nmol/L	3–20 ng/mL
Male		<3.18 nmol/L	<1.0 ng/mL
Prolactin	S		
Male		53–360 mg/L	2.5–17 ng/mL
Female		40–530 mg/L	1.9–25 ng/mL
Prostate-specific antigen (PSA)	S	0.0–4.0 µg/L	0.0–4.0 ng/mL
Prostate-specific antigen, free	S	With total PSA between 4 and 10 µg/L and when the free PSA is: >0.25 decreased risk of prostate cancer <0.10 increased risk of prostate cancer	With total PSA between 4 and 10 ng/mL and when the free PSA is: >25% decreased risk of prostate cancer <10% increased risk of prostate cancer
Protein fractions:	S		
Albumin		35–55 g/L	3.5–5.5 g/dL (50–60%)
Globulin		20–35 g/L	2.0–3.5 g/dL (40–50%)
Alpha ₁		2–4 g/L	0.2–0.4 g/dL (4.2–7.2%)
Alpha ₂		5–9 g/L	0.5–0.9 g/dL (6.8–12%)
Beta		6–11 g/L	0.6–1.1 g/dL (9.3–15%)
Gamma		7–17 g/L	0.7–1.7 g/dL (13–23%)
Protein, total	S	67–86 g/L	6.7–8.6 g/dL
Pyruvate	P	40–130 µmol/L	0.35–1.14 mg/dL
Retinol-binding protein	S	0.71–2.9 µmol/L	1.5–6.0 mg/dL
Rheumatoid factor	S	<15 kIU/L	<15 IU/mL
Serotonin	WB	0.28–1.14 µmol/L	50–200 ng/mL

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Serum protein electrophoresis	S	Not applicable	Normal pattern
Sex hormone-binding globulin (adult)	S		
Male		11–80 nmol/L	11–80 nmol/L
Female		30–135 nmol/L	30–135 nmol/L
Sodium	S	136–146 mmol/L	136–146 meq/L
Somatomedin-C (IGF-1) (adult)	S		
16 years		226–903 µg/L	226–903 ng/mL
17 years		193–731 µg/L	193–731 ng/mL
18 years		163–584 µg/L	163–584 ng/mL
19 years		141–483 µg/L	141–483 ng/mL
20 years		127–424 µg/L	127–424 ng/mL
21–25 years		116–358 µg/L	116–358 ng/mL
26–30 years		117–329 µg/L	117–329 ng/mL
31–35 years		115–307 µg/L	115–307 ng/mL
36–40 years		119–204 µg/L	119–204 ng/mL
41–45 years		101–267 µg/L	101–267 ng/mL
46–50 years		94–252 µg/L	94–252 ng/mL
51–55 years		87–238 µg/L	87–238 ng/mL
56–60 years		81–225 µg/L	81–225 ng/mL
61–65 years		75–212 µg/L	75–212 ng/mL
66–70 years		69–200 µg/L	69–200 ng/mL
71–75 years		64–188 µg/L	64–188 ng/mL
76–80 years		59–177 µg/L	59–177 ng/mL
81–85 years		55–166 µg/L	55–166 ng/mL
Somatostatin	P	<25 ng/L	<25 pg/mL
Testosterone, free			
Female, adult	S	10.4–65.9 pmol/L	3–19 pg/mL
Male, adult		312–1041 pmol/L	90–300 pg/mL
Testosterone, total,	S		
Female		0.21–2.98 nmol/L	6–86 ng/dL
Male		9.36–37.10 nmol/L	270–1070 ng/dL
Thyroglobulin	S	13–318 µg/L	1.3–31.8 ng/mL
Thyroid-binding globulin	S	13–30 mg/L	1.3–3.0 mg/dL
Thyroid-stimulating hormone (thyrotropin)	S	0.34–4.25 mIU/L	0.34–4.25 µIU/mL
Thyrotropin receptor antibody	S	≤1.75 IU/L	≤1.75 mIU/mL
Thyroxine, free (fT ₄)	S	9.0–16 pmol/L	0.7–1.24 ng/dL
Thyroxine, total (T ₄)	S	70–151 nmol/L	5.4–11.7 µg/dL
Thyroxine index (free)	S	6.7–10.9	6.7–10.9
Tissue transglutaminase (tTG) Antibody, IgA	S		
Negative		<4.0 Units/mL	<4.0 Units/mL
Weak positive		4.0–10.0 Units/mL	4.0–10.0 Units/mL
Positive		>10.0 Units/mL	>10.0 Units/mL
Tissue transglutaminase (tTG) antibody, IgG	S		
Negative		<6.0 Units/mL	<6.0 Units/mL
Weak positive		6.0–9.0 Units/mL	6.0–9.0 Units/mL
Positive		>9.0 Units/mL	>9.0 Units/mL
Transferrin	S	2.0–4.0 g/L	200–400 mg/dL
Transferrin, carbohydrate deficient, for alcohol use	S	0.017	1.7%
Triglycerides	S	0.34–2.26 mmol/L	30–200 mg/dL
Triiodothyronine, free (fT ₃)	S	3.7–6.5 pmol/L	2.4–4.2 pg/mL
Triiodothyronine, total (T ₃)	S	1.2–2.1 nmol/L	77–135 ng/dL
Troponin I	S, P		
99th percentile of a healthy population		Method-dependent	Method-dependent
Troponin T	S, P		
99th percentile of a healthy population		0–14 ng/L	0–14 ng/L
Urea nitrogen	S	2.5–7.1 mmol/L	7–20 mg/dL

(Continued)

TABLE 2 CLINICAL CHEMISTRY AND IMMUNOLOGY (CONTINUED)

Analyte	Specimen	SI Units	Conventional Units
Uric acid	S		
Females		0.15–0.33 mmol/L	2.5–5.6 mg/dL
Males		0.18–0.41 mmol/L	3.1–7.0 mg/dL
Vasoactive intestinal polypeptide	P	0–60 ng/L	0–60 pg/mL
Zinc protoporphyrin	WB	0–400 µg/L	0–40 µg/dL
Zinc protoporphyrin (ZPP)-to-heme ratio	WB	0–69 µmol ZPP/mol heme	0–69 µmol ZPP/mol heme

Abbreviations: AMF, amniotic fluid; P, plasma; S, serum; WB, whole blood.

Source: NJ Stone et al: J Am Coll Cardiol 63:2889-2934, 2014.

TABLE 3 TOXICOLOGY AND THERAPEUTIC DRUG MONITORING

Drug	Therapeutic Range		Toxic Level	
	SI Units	Conventional Units	SI Units	Conventional Units
Acetaminophen	66–199 µmol/L	10–30 µg/mL	>1320 µmol/L	>200 µg/mL
Amikacin				
Peak	34–51 µmol/L	20–30 µg/mL	>60 µmol/L	>35 µg/mL
Trough	0–17 µmol/L	0–10 µg/mL	>17 µmol/L	>10 µg/mL
Amitriptyline/nortriptyline (total drug)	430–900 nmol/L	120–250 ng/mL	>1800 nmol/L	>500 ng/mL
Amphetamine	150–220 nmol/L	20–30 ng/mL	>1500 nmol/L	>200 ng/mL
Bromide	9.4–18.7 mmol/L	75–150 mg/dL	>18.8 mmol/L	>150 mg/dL
Mild toxicity			6.4–18.8 mmol/L	51–150 mg/dL
Severe toxicity			>18.8 mmol/L	>150 mg/dL
Lethal			>37.5 mmol/L	>300 mg/dL
Caffeine	25.8–103 µmol/L	5–20 µg/mL	>206 µmol/L	>40 µg/mL
Carbamazepine	17–42 µmol/L	4–10 µg/mL	> 85 µmol/L	>20 µg/mL
Chloramphenicol				
Peak	31–62 µmol/L	10–20 µg/mL	>77 µmol/L	>25 µg/mL
Trough	15–31 µmol/L	5–10 µg/mL	>46 µmol/L	>15 µg/mL
Chlordiazepoxide	1.7–10 µmol/L	0.5–3.0 µg/mL	>17 µmol/L	>5.0 µg/mL
Clonazepam	32–240 nmol/L	10–75 ng/mL	>320 nmol/L	>100 ng/mL
Clozapine	0.6–2.1 µmol/L	200–700 ng/mL	>3.7 µmol/L	>1200 ng/mL
Cocaine			>3.3 µmol/L	>1.0 µg/mL
Codeine	43–110 nmol/mL	13–33 ng/mL	>3700 nmol/mL	>1100 ng/mL (lethal)
Cyclosporine				
Renal transplant				
0–6 months	208–312 nmol/L	250–375 ng/mL	>312 nmol/L	>375 ng/mL
6–12 months after transplant	166–250 nmol/L	200–300 ng/mL	>250 nmol/L	>300 ng/mL
>12 months	83–125 nmol/L	100–150 ng/mL	>125 nmol/L	>150 ng/mL
Cardiac transplant				
0–6 months	208–291 nmol/L	250–350 ng/mL	>291 nmol/L	>350 ng/mL
6–12 months after transplant	125–208 nmol/L	150–250 ng/mL	>208 nmol/L	>250 ng/mL
>12 months	83–125 nmol/L	100–150 ng/mL	>125 nmol/L	150 ng/mL
Lung transplant				
0–6 months	250–374 nmol/L	300–450 ng/mL	>374 nmol/L	>450 ng/mL
Liver transplant				
Initiation	208–291 nmol/L	250–350 ng/mL	>291 nmol/L	>350 ng/mL
Maintenance	83–166 nmol/L	100–200 ng/mL	>166 nmol/L	>200 ng/mL
Desipramine	375–1130 nmol/L	100–300 ng/mL	>1880 nmol/L	>500 ng/mL
Diazepam (and metabolite)				
Diazepam	0.7–3.5 µmol/L	0.2–1.0 µg/mL	>7.0 µmol/L	>2.0 µg/mL
Nordiazepam	0.4–6.6 µmol/L	0.1–1.8 µg/mL	>9.2 µmol/L	>2.5 µg/mL
Digoxin	0.64–1.6 nmol/L	0.5–1.2 ng/mL	>3.1 nmol/L	>2.4 ng/mL
Disopyramide	5.3–14.7 µmol/L	2–5 µg/mL	>20.6 µmol/L	>7 µg/mL
Doxepin and nordoxepin				
Doxepin	0.36–0.98 µmol/L	101–274 ng/mL	>1.8 µmol/L	>503 ng/mL
Nordoxepin	0.38–1.04 µmol/L	106–291 ng/mL	>1.9 µmol/L	>531 ng/mL

(Continued)

TABLE 3 TOXICOLOGY AND THERAPEUTIC DRUG MONITORING (CONTINUED)

Drug	Therapeutic Range		Toxic Level	
	SI Units	Conventional Units	SI Units	Conventional Units
Ethanol				
Behavioral changes			>4.3 mmol/L	>20 mg/dL
Legal limit			≥17 mmol/L	≥80 mg/dL
Critical with acute exposure			>54 mmol/L	>250 mg/dL
Ethylene glycol				
Toxic			>2 mmol/L	>12 mg/dL
Lethal			>20 mmol/L	>120 mg/dL
Ethosuximide	280–700 μmol/L	40–100 μg/mL	>700 μmol/L	>100 μg/mL
Everolimus (trough)				
Kidney transplant	3.1–8.34 nmol/L	3–8 ng/mL	>12.5 nmol/L	>12 ng/mL
Oncology	5.2–10.4 nmol/L	5.0–10.0 ng/mL	>12.5 nmol/L	>12 ng/mL
Felbamate (trough)	126–252 μmol/L	30–60 μg/mL	Toxic range not well established	
Flecainide	0.5–2.4 μmol/L	0.2–1.0 μg/mL	>3.6 μmol/L	>1.5 μg/mL
Gabapentin (trough)	11.7–116.8 μmol/L	2–20 μg/mL	Toxic range not well established	
Gentamicin				
Peak	10–21 μmol/mL	5–10 μg/mL	>25 μmol/mL	>12 μg/mL
Trough	0–4.2 μmol/mL	0–2 μg/mL	>4.2 μmol/mL	>2 μg/mL
Heroin (diacetyl morphine)			>700 μmol/L	>200 ng/mL (as morphine)
Ibuprofen	49–243 μmol/L	10–50 μg/mL	>970 μmol/L	>200 μg/mL
Imipramine (and metabolites)				
Desimipramine	375–1130 nmol/L	100–300 ng/mL	>1880 nmol/L	>500 ng/mL
Total imipramine + desimipramine	563–1130 nmol/L	150–300 ng/mL	>1880 nmol/L	>500 ng/mL
Isopropanol			>8.3 mmol/L	>50 mg/dL
Lamotrigine	11.7–54.7 μmol/L	3–14 μg/mL	>58.7 μmol/L	>15 μg/mL
Levetiracetam (Keppra)	29–176 μmol/L	5–30 μg/mL	Toxic range not well established	
Lidocaine	5.1–21.3 μmol/L	1.2–5.0 μg/mL	>38.4 μmol/L	>9.0 μg/mL
Lithium	0.5–1.3 mmol/L	0.5–1.3 meq/L	>2 mmol/L	>2 meq/L
Methadone	1.0–3.2 μmol/L	0.3–1.0 μg/mL	>6.5 μmol/L	>2 μg/mL
Methamphetamine	0.07–0.34 μmol/L	0.01–0.05 μg/mL	>3.35 μmol/L	>0.5 μg/mL
Methanol			>6 mmol/L	>20 mg/dL
Methotrexate				
Low-dose	0.01–0.1 μmol/L	0.01–0.1 μmol/L	>0.1 μmol/L	>0.1 μmol/L
High-dose (24h)	<5.0 μmol/L	<5.0 μmol/L	>5.0 μmol/L	>5.0 μmol/L
High-dose (48h)	<0.50 μmol/L	<0.50 μmol/L	>0.5 μmol/L	>0.5 μmol/L
High-dose (72h)	<0.10 μmol/L	<0.10 μmol/L	>0.1 μmol/L	>0.1 μmol/L
Morphine	232–286 μmol/L	65–80 ng/mL	>720 μmol/L	>200 ng/mL
Mycophenolic acid	3.1–10.9 μmol/L	1.0–3.5 ng/mL	>37 μmol/L	>12 ng/mL
Nitroprusside (as thiocyanate)	103–499 μmol/L	6–29 μg/mL	860 μmol/L	>50 μg/mL
Nortriptyline	190–569 nmol/L	50–150 ng/mL	>1900 nmol/L	>500 ng/mL
Phenobarbital	65–172 μmol/L	15–40 μg/mL	>258 μmol/L	>60 μg/mL
Phenytoin	40–79 μmol/L	10–20 μg/mL	>158 μmol/L	>40 μg/mL
Phenytoin, free	4.0–7.9 μg/mL	1–2 μg/mL	>13.9 μg/mL	>3.5 μg/mL
% Free	0.08–0.14	8–14%		
Primidone and metabolite				
Primidone	23–55 μmol/L	5–12 μg/mL	>69 μmol/L	>15 μg/mL
Phenobarbital	65–172 μmol/L	15–40 μg/mL	>215 μmol/L	>50 μg/mL
Procainamide				
Procainamide	17–42 μmol/L	4–10 μg/mL	>43 μmol/L	>10 μg/mL
NAPA (N-acetylprocainamide)	22–72 μmol/L	6–20 μg/mL	>126 μmol/L	>35 μg/mL
Quinidine	6.2–15.4 μmol/L	2.0–5.0 μg/mL	>19 μmol/L	>6 μg/mL
Salicylates	145–2100 μmol/L	2–29 mg/dL	>2900 μmol/L	>40 mg/dL
Sirolimus (trough level)				
Kidney transplant	4.4–15.4 nmol/L	4–14 ng/mL	>16 nmol/L	>15 ng/mL

(Continued)

TABLE 3 TOXICOLOGY AND THERAPEUTIC DRUG MONITORING (CONTINUED)

Drug	Therapeutic Range		Toxic Level	
	SI Units	Conventional Units	SI Units	Conventional Units
Tacrolimus (FK506) (trough)				
Kidney and liver transplant				
Initiation	12–19 nmol/L	10–15 ng/mL	>25 nmol/L	>20 ng/mL
Maintenance	6–12 nmol/L	5–10 ng/mL	>25 nmol/L	>20 ng/mL
Heart transplant				
Initiation	19–25 nmol/L	15–20 ng/mL		
Maintenance	6–12 nmol/L	5–10 ng/mL		
Theophylline	56–111 µg/mL	10–20 µg/mL	>168 µg/mL	>30 µg/mL
Thiocyanate				
After nitroprusside infusion	103–499 µmol/L	6–29 µg/mL	860 µmol/L	>50 µg/mL
Nonsmoker	17–69 µmol/L	1–4 µg/mL		
Smoker	52–206 µmol/L	3–12 µg/mL		
Tobramycin				
Peak	11–22 µg/L	5–10 µg/mL	>26 µg/L	>12 µg/mL
Trough	0–4.3 µg/L	0–2 µg/mL	>4.3 µg/L	>2 µg/mL
Topiramate (trough)	5.9–58.9 µmol/L	2.0–20.0 µg/mL	Toxic range not well established	
Valproic acid	346–693 µmol/L	50–100 µg/mL	>693 µmol/L	>100 µg/mL
Vancomycin				
Peak	14–28 µmol/L	20–40 µg/mL	>55 µmol/L	>80 µg/mL
Trough	3.5–10.4 µmol/L	5–15 µg/mL	>14 µmol/L	>20 µg/mL
Zonisamide (trough)	47–188 µmol/L	10–40 µg/mL	Toxic range not well established	

TABLE 4 VITAMINS AND SELECTED TRACE MINERALS

Specimen	Analyte	Reference Range	
		SI Units	Conventional Units
Aluminum	S	<0.2 µmol/L	<5.41 µg/L
Arsenic	WB	0.0–0.17 µmol/L	0–13 µg/L
Cadmium	WB	<44.5 nmol/L	<5.0 µg/L
Coenzyme Q10 (ubiquinone)	P	433–1532 µg/L	433–1532 µg/L
β-Carotene	S	0.07–1.43 µmol/L	4–77 µg/dL
Copper	S	11–22 µmol/L	70–140 µg/dL
Folic acid	RC	340–1020 nmol/L cells	150–450 ng/mL cells
Folic acid	S	12.2–40.8 nmol/L	5.4–18.0 ng/mL
Lead	S		
Adult		<0.5 µmol/L	<10 µg/dL
Children		<0.25 µmol/L	<5 µg/dL
Mercury	WB	0–50 µmol/L	0–10 µg/L
Selenium	S	0.8–2.0 µmol/L	63–160 µg/L
Vitamin A	S	0.7–3.5 µmol/L	20–100 µg/dL
Vitamin B ₁ (thiamine)	S	0–75 nmol/L	0–2 µg/dL
Vitamin B ₂ (riboflavin)	S	106–638 nmol/L	4–24 µg/dL
Vitamin B ₆	P	20–121 nmol/L	5–30 ng/mL
Vitamin B ₁₂	S	206–735 pmol/L	279–996 pg/mL
Vitamin C (ascorbic acid)	S	23–57 µmol/L	0.4–1.0 mg/dL
Vitamin D 1,25-dihydroxy, total	S, P	36–180 pmol/L	15–75 pg/mL
Vitamin D 25-hydroxy, total	P	75–250 nmol/L	30–100 ng/mL
Vitamin E	S	12–42 µmol/L	5–18 µg/mL
Vitamin K	S	0.29–2.64 nmol/L	0.13–1.19 ng/mL
Zinc	S	11.5–18.4 µmol/L	75–120 µg/dL

Abbreviations: P, plasma; RC, red cells; S, serum; WB, whole blood.

TABLE 5 CEREBROSPINAL FLUID^a

Constituent	Reference Range	
	SI Units	Conventional Units
Osmolarity	292–297 mmol/kg water	292–297 mOsm/L
Electrolytes		
Sodium	137–145 mmol/L	137–145 meq/L
Potassium	2.7–3.9 mmol/L	2.7–3.9 meq/L
Calcium	1.0–1.5 mmol/L	2.1–3.0 meq/L
Magnesium	1.0–1.2 mmol/L	2.0–2.5 meq/L
Chloride	116–122 mmol/L	116–122 meq/L
CO ₂ content	20–24 mmol/L	20–24 meq/L
Pco ₂	6–7 kPa	45–49 mmHg
pH	7.31–7.34	
Glucose	2.22–3.89 mmol/L	40–70 mg/dL
Lactate	1–2 mmol/L	10–20 mg/dL
Total protein:		
Lumbar	0.15–0.5 g/L	15–50 mg/dL
Cisternal	0.15–0.25 g/L	15–25 mg/dL
Ventricular	0.06–0.15 g/L	6–15 mg/dL
Albumin	0.066–0.442 g/L	6.6–44.2 mg/dL
IgG	0.009–0.057 g/L	0.9–5.7 mg/dL
IgG index ^b	0.29–0.59	
Oligoclonal bands (OGB)	<2 bands not present in matched serum sample	
Ammonia	15–47 μmol/L	25–80 μg/dL
Creatinine	44–168 μmol/L	0.5–1.9 mg/dL
Myelin basic protein	<4 μg/L	
Cerebrospinal fluid pressure		50–180 mmH ₂ O
Cerebrospinal fluid volume (adult)	~150 mL	
Red blood cells	0	0
Leukocytes		
Total	0–5 mononuclear cells per μL	
Differential		
Lymphocytes	60–70%	
Monocytes	30–50%	
Neutrophils	None	

^aSince cerebrospinal fluid (CSF) concentrations are equilibrium values, measurements of the same parameters in blood plasma obtained at the same time are recommended. However, there is a time lag in attainment of equilibrium, and cerebrospinal levels of plasma constituents that can fluctuate rapidly (such as plasma glucose) may not achieve stable values until after a significant lag phase. ^bIgG index = CSF IgG (mg/dL) × serum albumin (g/dL) / serum IgG (g/dL) × CSF albumin (mg/dL).

TABLE 6 URINE ANALYSIS AND RENAL FUNCTION TESTS

	Reference Range	
	SI Units	Conventional Units
Acidity, titratable	20–40 mmol/d	20–40 meq/d
Aldosterone	Normal diet: 6–25 µg/d Low-salt diet: 17–44 µg/d High-salt diet: 0–6 µg/d	Normal diet: 6–25 µg/d Low-salt diet: 17–44 µg/d High-salt diet: 0–6 µg/d
Aluminum	0.19–1.11 µmol/L	5–30 µg/L
Ammonia	30–50 mmol/d	30–50 meq/d
Amylase		4–400 U/L
Amylase/creatinine clearance ratio ($[(Cl_{am}/Cl_{cr}) \times 100]$)	1–5	1–5
Arsenic	0.07–0.67 µmol/d	5–50 µg/d
Bence Jones protein, urine, qualitative	Not applicable	None detected
Bence Jones protein, urine, quantitative		
Free kappa	1.4–24.2 mg/L	0.14–2.42 mg/dL
Free lambda	0.2–6.7 mg/L	0.02–0.67 mg/dL
K/L ratio	2.04–10.37	2.04–10.37
Calcium (10 meq/d or 200 mg/d dietary calcium)	<7.5 mmol/d	<300 mg/d
Chloride	140–250 mmol/d	140–250 mmol/d
Citrate	320–1240 mg/d	320–1240 mg/d
Copper	<0.95 µmol/d	<60 µg/d
Coproporphyrins (types I and III)	0–20 µmol/mol creatinine	0–20 µmol/mol creatinine
Cortisol, free	55–193 nmol/d	20–70 µg/d
Creatine, as creatinine		
Female	<760 µmol/d	<100 mg/d
Male	<380 µmol/d	<50 mg/d
Creatinine	8.8–14 mmol/d	1.0–1.6 g/d
Dopamine	392–2876 nmol/d	60–440 µg/d
Eosinophils	<100 eosinophils/mL	<100 eosinophils/mL
Epinephrine	0–109 nmol/d	0–20 µg/d
Glomerular filtration rate	>60 mL/min/1.73 m ² For African Americans, multiply the result by 1.21	>60 mL/min/1.73 m ² For African Americans, multiply the result by 1.21
Glucose (glucose oxidase method)	0.3–1.7 mmol/d	50–300 mg/d
5-Hydroindoleacetic acid (5-HIAA)	0–78.8 µmol/d	0–15 mg/d
Hydroxyproline	53–328 µmol/d	53–328 µmol/d
Iodine, spot urine		
WHO classification of iodine deficiency:		
Not iodine deficient	>100 µg/L	>100 µg/L
Mild iodine deficiency	50–100 µg/L	50–100 µg/L
Moderate iodine deficiency	20–49 µg/L	20–49 µg/L
Severe iodine deficiency	<20 µg/L	<20 µg/L
Ketone (acetone)	Negative	Negative
17 Ketosteroids	3–12 mg/d	3–12 mg/d
Metanephrines		
Metanephrine	30–350 µg/d	30–350 µg/d
Normetanephrine	50–650 µg/d	50–650 µg/d
Microalbumin		
Normal	0.0–0.03 g/d	0–30 mg/d
Microalbuminuria	0.03–0.30 g/d	30–300 mg/d
Clinical albuminuria	>0.3 g/d	>300 mg/d
Microalbumin/creatinine ratio		
Normal	0–3.4 g/mol creatinine	0–30 µg/mg creatinine
Microalbuminuria	3.4–34 g/mol creatinine	30–300 µg/mg creatinine
Clinical albuminuria	>34 g/mol creatinine	>300 µg/mg creatinine
β ₂ -Microglobulin	0–160 µg/L	0–160 µg/L
Norepinephrine	89–473 nmol/d	15–80 µg/d

(Continued)

TABLE 6 URINE ANALYSIS AND RENAL FUNCTION TESTS (CONTINUED)

	Reference Range	
	SI Units	Conventional Units
<i>N</i> -telopeptide (cross-linked), NTx		
Female, premenopausal	17–94 nmol BCE/mmol creatinine	17–94 nmol BCE/mmol creatinine
Female, postmenopausal	26–124 nmol BCE/mmol creatinine	26–124 nmol BCE/mmol creatinine
Male	21–83 nmol BCE/mmol creatinine	21–83 nmol BCE/mmol creatinine
Osmolality	100–800 mosm/kg	100–800 mosm/kg
Oxalate		
Male	80–500 μ mol/d	7–44 mg/d
Female	45–350 μ mol/d	4–31 mg/d
pH	5.0–9.0	5.0–9.0
Phosphate (phosphorus) (varies with intake)	12.9–42.0 mmol/d	400–1300 mg/d
Porphobilinogen	None	None
Potassium (varies with intake)	25–100 mmol/d	25–100 meq/d
Protein	<0.15 g/d	<150 mg/d
Protein/creatinine ratio	Male: 15–68 mg/g Female: 10–107 mg/g	Male: 15–68 mg/g Female: 10–107 mg/g
Sediment		
Red blood cells	0–2/high-power field	
White blood cells	0–2/high-power field	
Bacteria	None	
Crystals	None	
Bladder cells	None	
Squamous cells	None	
Tubular cells	None	
Broad casts	None	
Epithelial cell casts	None	
Granular casts	None	
Hyaline casts	0–5/low-power field	
Red blood cell casts	None	
Waxy casts	None	
White cell casts	None	
Sodium (varies with intake)	100–260 mmol/d	100–260 meq/d
Specific gravity:		
After 12-h fluid restriction	>1.025	>1.025
After 12-h deliberate water intake	\leq 1.003	\leq 1.003
Tubular reabsorption, phosphorus	0.79–0.94 of filtered load	79–94% of filtered load
Urea nitrogen	214–607 mmol/d	6–17 g/d
Uric acid (normal diet)	1.49–4.76 mmol/d	250–800 mg/d
Vanillylmandelic acid (VMA)	<30 μ mol/d	<6 mg/d

Abbreviation: BCE = bone collagen equivalent.

ACKNOWLEDGMENT

The contributions of Drs. Daniel J. Fink, Patrick M. Sluss, James L. Januzzi, Kent B. Lewandrowski, Amudha Palanisamy, and Scott Fink to this chapter in previous editions are gratefully acknowledged. We also express our gratitude to Drs. Alex Rai and Jeffrey Jhang for helpful suggestions.

480e The Clinical Laboratory in Modern Health Care

Anthony A. Killeen

Modern medicine relies extensively on the clinical laboratory as a key component of health care. It is estimated that, in current practice, at least 60–70% of all clinical decisions rely to some extent on a laboratory result. For many diseases, the clinical laboratory provides essential diagnostic information. As an example, histopathologic analysis provides basic information about histologic type and classification of tumors and their degree of invasion into adjacent tissues. Microbiologic testing is required to identify infectious organisms and determine antibiotic susceptibility. For many common diseases, expert groups have produced standard guidelines for diagnosis that rely on defined clinical laboratory values, e.g., blood glucose or hemoglobin A_{1c} levels form the basis for diagnosis of diabetes mellitus; the presence of specific serum antibodies is required for diagnosis of many rheumatologic diseases; and serum levels of cardiac markers are a mainstay in diagnosis of acute coronary syndromes.

With their ever-increasing number and scope, clinical laboratory tests provide the clinician with a powerful set of tools but pose challenges in terms of appropriate selection and judicious, cost-effective use to deliver effective patient care.

RATIONALE FOR PERFORMING CLINICAL LABORATORY TESTS

DIAGNOSIS OF DISEASE

One of the most frequent reasons for performing clinical laboratory tests is to support, confirm, or refute a diagnosis of disease that is suspected on the basis of other information sources, such as history, physical examination findings, and imaging studies. The following questions need to be considered: Which clinical laboratory tests may be of value in supporting, confirming, or excluding the clinical impression? What is the most efficient test-ordering strategy? Will a positive test result confirm the clinical impression or even definitively establish the diagnosis? Will a negative result disprove the clinical suspicion, and, if so, what further testing or alternative approach will be needed? What are the known sources of false-positive and false-negative results, and how are these misleading results recognized?

SCREENING FOR DISEASE

Another reason for ordering clinical laboratory tests is to screen for disease in asymptomatic individuals ([Chap. 4](#)). Perhaps the most common examples of this application are the newborn screening programs now routinely used in most developed countries. Their purpose is to identify newborns with treatable conditions for which early intervention—even before clinical symptoms develop—is known to be beneficial. In adults, screening tests for diabetes mellitus, renal disease, prostate cancer (measurement of serum prostate-specific antigen [PSA] levels), and colorectal cancer (testing for occult blood in stool), for example, are widely applied to apparently healthy individuals on the grounds that early diagnosis and intervention lead to improved long-term outcomes.

Differences between Screening Tests and Confirmatory Tests It is important to distinguish between clinical laboratory tests that can be used to *screen* for disease and those that offer a *confirmatory result*. Screening tests are generally less expensive and more widely available than are confirmatory tests, which often require more specialized equipment or personnel. As a general principle, screening tests are designed to identify all individuals who have the disease of interest, even if, in the process, some healthy individuals are misidentified as possibly having the disease. In other words, maximization of the diagnostic sensitivity of screening tests inevitably comes at the expense of reduced diagnostic specificity. Confirmatory testing is intended to correctly separate those individuals who have a disease from those who do not.

These principles are exemplified by screening for hepatitis C virus (HCV) infection. A common approach is to test first for the presence of antibodies to HCV in serum. A positive result generally indicates either a current infection or a previous infection that the patient's immune system has successfully cleared. In the latter situation, antibodies may persist at detectable levels for life. However, a small proportion of patients have false-positive results in the serologic screening test for HCV. To resolve uncertainty in these instances, a positive serologic screening test should be followed by confirmatory identification of HCV RNA with molecular techniques. This confirmatory testing can provide evidence of current viral infection and can identify patients who are not infected.

RISK ASSESSMENT OF FUTURE DISEASE

Another reason for clinical laboratory testing is to assess a patient's risk of developing disease in the future. A number of diseases are associated with well-established clinical laboratory-defined risk factors that, if present, indicate the need for more frequent monitoring for the disease in question. The need for risk assessment is even clearer if there are useful interventions that decrease the risk of developing disease. For example, hypercholesterolemia is a well-established risk factor for coronary artery disease that may be modified by pharmacologic intervention ([Chap. 291e](#)). Many genetic mutations are known to be associated with increased risk of cancer. For example, hereditary mutations in the *BRCA1* and *BRCA2* genes predispose to breast and/or ovarian cancer. Individuals who are known to carry these mutations require more vigilant monitoring for early signs of cancer and may even opt for prophylactic surgery in an attempt to prevent cancer ([Chap. 84](#)). Individuals with factor V Leiden are at increased risk of developing deep venous thrombosis and may benefit from prophylactic anticoagulation in the perioperative period.

MONITORING DISEASE AND THERAPY

Many clinical laboratory tests offer useful information on the progress of disease and the response to therapy. One example is the measurement of viral load in HIV-1-infected patients who are taking antiretroviral agents. According to current guidelines from the Centers for Disease Control and Prevention, a successful antiretroviral response is defined by a fall in plasma HIV-1 levels of 0.5 log₁₀ copies/mL, and a key goal of treatment is a reduction in the viral load to below the level of detection, which is typically in the range of 40–75 copies/mL. Other examples of the use of clinical laboratory testing for monitoring disease include measurement of tumor markers such as PSA, especially following surgical removal of tumors. In this situation, the expectation is that successful treatment of a tumor will cause a decrease in the level of the tumor marker. A later increase in the level of the tumor marker suggests a recurrence of the disease. Finally, the clinical laboratory offers direct monitoring of levels of some therapeutic agents, such as drugs. This monitoring is important if a drug has a defined therapeutic concentration range above which it is toxic and below which it is ineffective. Monitoring of drug levels in this situation facilitates optimal dosing and avoidance of toxicity.

CRITICAL VALUES

It is common practice for clinical laboratories to establish a list of “critical values.” These are values of test results that indicate immediate risk to the health or life of the patient and therefore require urgent communication with the patient's physician so that appropriate medical intervention may be initiated. Critical values are reported regardless of whether the test was ordered as a “stat” or a routine test. The critical values themselves are generally created by the clinical laboratory medical director in conjunction with the medical staff. A representative list of critical values is shown in [Table 480e-1](#).

“STAT” ORDERS

Tests that are ordered on a “stat” basis receive priority in the clinical laboratory's testing queue; testing of specimens from other patients may be delayed while a stat specimen is analyzed. Ordering a test stat

TABLE 480e-1 SELECTED EXAMPLES OF LABORATORY CRITICAL VALUES

Value	Less Than	Greater Than
Chemistry		
Ammonia		>100 $\mu\text{mol/L}$
Calcium, ionized	<3 mg/dL	>7 mg/dL
Calcium, total	<6 mg/dL	>14 mg/dL
Carboxyhemoglobin		>10%
Creatine kinase, total		>1000 U/L
CO ₂ , total	<11 mol/L	>45 mmol/L
Digoxin		>2.5 $\mu\text{g/L}$
Glucose	<40 mg/dL	>500 mg/dL
Glucose—CSF	< 40 mg/dL	
Ketone		>1.5 mmol/L
Lithium		> 2.0 mmol/L
Magnesium		>7 mg/dL
Methemoglobin		>35%
Osmolality	<250 mmol/L	>340 mmol/L
Phosphorus	<1 mg/dL	
pH	<7.1	>7.6
Pco ₂	<20 mmHg	>75 mmHg
Po ₂ , arterial	<40 mmHg	
Po ₂ , capillary	<30 mmHg	
Bicarbonate	<11 mol/L	>45 mol/L
Potassium	<2.7 mmol/L	>6 mmol/L
Salicylate		>30 mg/L
Sodium	<120 mmol/L	>160 mmol/L
Troponin		>0.120 $\mu\text{g/L}$
Hematology		
INR		>6.0
PTT		>105 secs
Hemoglobin	<7 g/dL	
WBCs	<1 $\times 10^9/\text{L}$	>50 $\times 10^9/\text{L}$
Platelets	<50 $\times 10^9/\text{L}$	>1000 $\times 10^9/\text{L}$
Microbiology^a		
Acid fast culture or smear		
Blood culture		
CSF Gram stain/culture		
CSF cryptococcal antigen		
Malarial smear		

^aAny positive result is critical.

Abbreviations: CO₂, carbon dioxide; CSF, cerebrospinal fluid; INR, international normalized ratio; Pco₂, partial pressure of carbon dioxide; Po₂, partial pressure of oxygen; PTT, partial thromboplastin time; WBCs, white blood cells.

should be reserved for situations in which a result is needed for urgent medical care—a judgment that must be made by the ordering physician. Stat testing should not be used merely for the convenience of either the patient or the health care provider.

SENSITIVITY AND SPECIFICITY IN THE CLINICAL LABORATORY

The commonly used metrics of a clinical laboratory test are the diagnostic sensitivity, specificity, and positive and negative predictive values. These concepts are discussed in [Chap. 3](#). In the clinical laboratory, the terms *sensitivity* and *specificity* have alternative meanings that are applied to tests, and the different uses of these terms may cause confusion. *Analytic sensitivity* can refer to the lowest detectable concentration of analyte that can be measured with some defined certainty or to the rate of change of signal intensity as analyte concentration changes. For example, newer generations of laboratory assays frequently exhibit improved sensitivity over that of earlier generations, i.e., they can detect lower concentrations of the analyte—a feature that is often of value in disease diagnosis. *Analytic specificity* refers to the extent to

which other substances in the test system interfere with measurement of the analyte of interest. This concept is frequently applied to immunoassays, in which a detection antibody may also bind with compounds that have a structure similar to the substance sought. For instance, immunoassays for drugs may show cross-reactivity with drug metabolites, and immunoassays for glucocorticoids may show cross-reactivity with other glucocorticoids of similar structure. Certain chemical assays are also subject to nonspecificity. For example, the Jaffe reaction, a chemical method commonly used to measure creatinine, is subject to positive interference from a number of other compounds, including glucose, certain ketones, and cephalosporin antibiotics. Elevated concentrations of bilirubin, free hemoglobin, or turbidity in plasma or serum specimens may also interfere with some assays. The clinical laboratory should be able to provide advice about the presence or magnitude of these effects in the assays it performs.

CLINICAL LABORATORY DIAGNOSTIC PRINCIPLES

Clinical laboratory diagnosis, like all diagnosis, is based on observation of disease-related changes from normality.

1. Tissue injury or necrosis allows leakage of intracellular components into the circulation, with consequent detectable rises in blood levels of these components. Many intracellular molecules are common across tissue types and are therefore not indicative of injury to a specific tissue. Other constituents are selectively expressed in relatively high concentrations—or are even uniquely present—in certain tissues. Therefore, their presence in the blood is evidence of injury to that tissue. This principle forms the rationale for measurement of blood levels of, for example, liver enzymes in evaluating liver disease ([Chap. 358](#)), cardiac troponins in acute coronary syndromes ([Chap. 295](#)), and myoglobin in muscle injury. The extent of the rise in blood levels of these markers generally correlates with the extent of tissue damage, although there are exceptions; for example, liver enzyme levels may fall in end-stage liver disease.
2. An increase in blood levels of some analytes indicates failure of normal excretory processes. This principle is illustrated by elevations in conjugated bilirubin that accompany obstruction of the biliary system, by elevations in ammonia in advanced or metabolic liver disease, by rises in creatinine and potassium levels in renal failure, and by increases in Pco₂ in some pulmonary diseases.
3. Increases in the blood concentration of tissue-specific markers may result from expansion of the total volume of that tissue. This principle forms the basis for the measurement of levels of many tumor markers such as PSA (prostate cancer), CA-125 (ovarian cancer), CEA (colon cancer), and CA-19-9 (pancreatic cancer). In practice, the usefulness of these markers varies with the degree to which they are produced by a tumor and by the tumor's size. Small colon cancers, for example, may not produce a significant rise in CEA levels, whereas small prostate cancers often produce detectable rises in PSA concentrations.
4. Disease processes often manifest characteristic patterns of coincident changes in levels of several analytes. These patterns of change can be understood by consideration of the underlying pathophysiology. For example, acute intravascular hemolysis is characterized by a fall in levels of hemoglobin and haptoglobin and by a rise in unconjugated bilirubin. In endocrine diseases, there are often changes in concentrations of several hormones because of disturbance of feedback loops. Primary hyperthyroidism, as an example, is characterized by increases in thyroxine and by suppression of thyroid-stimulating hormone. In diabetic ketoacidosis caused by insulin deficiency, there are concomitant elevations of plasma glucose, ketones, and (frequently) potassium. In response to metabolic acidosis, levels of bicarbonate are typically reduced.
5. Genetic changes underlie many diseases, both inherited and acquired. In the era of molecular medicine, there is increasing recognition of the contribution of hereditary factors to many common diseases. Often, the epidemiology of common diseases such as hypertension is characterized by a minority of families that have

mutations in recognized genes, whereas the genetic basis of the same disease phenotype in the larger population is unclear. The search for the genetic factors that contribute to many common diseases remains a topic of intense research interest. It is now clear that essentially all tumors have genetic abnormalities. Although there is an inherited predisposition in some families, most of these genetic changes are acquired. Identification of the genetic abnormalities in cancer offers new tools for clinical laboratory diagnosis and classification of tumors in ways that surpass traditional histopathology and also provides insights into cellular processes that may be targets for treatment.

- Clinical laboratory results should always be interpreted in the context of the patient's history and physical examination as well as any other relevant information (e.g., imaging studies). The clinician should avoid treating laboratory results rather than the patient.
- Recommended clinical laboratory tests change with time. As new markers of disease emerge, they may replace older markers. For example, measurement of serum creatine kinase (CK) levels was introduced for diagnosis of acute myocardial infarction in the 1980s. Use of the cardiac-specific isoenzyme CK-MB later became widespread in clinical practice. Today, cardiac troponins are replacing CK (or CK-MB) measurements in recommended guidelines. Many other assays have fallen out of use as better assays have become available. Measurement of urine 17-ketosteroids (arising from androgens) and of urine 17-hydroxycorticosteroids (arising from glucocorticoids) has been supplanted by immunoassays or mass spectroscopy determinations of specific steroid hormones. Today, many steroid hormones are measured by mass spectrometry, often with better analytic specificity than is provided by immunoassays. As new tests are introduced, it is essential that they be evaluated critically before adoption for clinical use. At a minimum, consideration needs to be given to questions of clinical validation, specimen stability, diagnostic sensitivity and specificity, positive and negative predictive values, analytic accuracy and precision, and relative costs.

REFERENCE RANGES

In the interpretation of clinical laboratory results, comparison is usually made to a reference range (sometimes called a normal range) that defines the values seen in health or considered to be desirable for health. Several common methods are used to describe reference ranges in the clinical laboratory.

- For many quantitative clinical laboratory tests, the range of observed values in a healthy population shows an approximately Gaussian distribution. The factors that contribute to this range include the inter- and intraindividual variation in the concentration of the analyte and the analytic imprecision. When there is an approximately Gaussian distribution of values in the population, the reference range is commonly defined as being the central 95% of the range of distribution of those values. According to this method, 2.5% of the population will have a measured value that is below the reference range for the analyte, and 2.5% will have a value that is above the reference range. The fact that 5% of healthy individuals will have a test value that is outside the reference range has important implications when multiple tests are ordered. If N independent tests are performed on a specimen, then the probability that at least one result will be outside the reference range is $(1 - 0.95^N)$. The greater the number of tests ordered (even for a healthy individual), the greater is the likelihood of an abnormal result (Fig. 480e-1). If 20 independent tests are performed on a healthy subject, the probability of at least one abnormal result is almost two-thirds.

In some settings, a narrower range of values is considered to be abnormal. For example, current American Heart Association guidelines recommend the use of a serum level of cardiac troponins that is greater than the 99th percentile of values found in a healthy population as evidence of acute myocardial infarction.

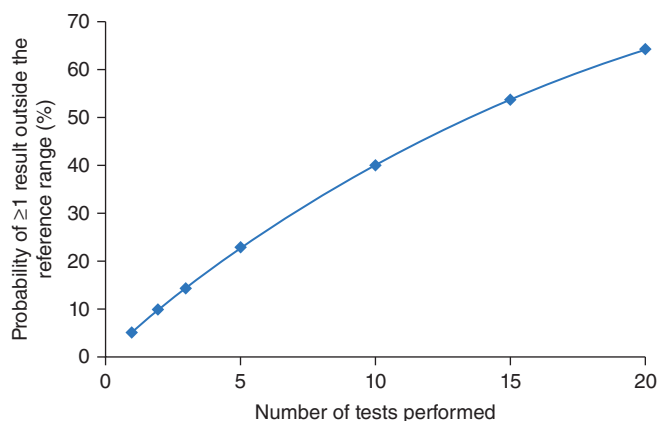


FIGURE 480e-1 Probability that at least one laboratory result will be abnormal in a healthy individual as an increasing number of independent tests are performed. The reference range is the central 95% of values measured in a healthy population.

- An alternative approach to using population means and standard deviations is to define a range of analyte values that is judged to be consistent with health on the basis of expert consensus opinion. These ranges are often referred to as *decision limits*. Examples of reference ranges established in this way include those for total, high-density, and low-density cholesterol (Table 480e-2). Such ranges may deviate considerably from those that would be established if the analyte concentrations of the population (mean \pm 2 standard deviations) were used as a basis for establishing the reference range. For example, the “desirable” total cholesterol value according to the National Cholesterol Education Program is <200 mg/dL. This value is actually very close to the mean concentration among U.S. adults; in fact, almost one-half of U.S. adults have a total cholesterol concentration that is above the “desirable” range. If the central 95% of cholesterol concentrations in the population were taken as the reference range, the upper end of that range would be ~240 mg/dL, well beyond what is considered desirable.

Reference ranges may vary with age, gender, ethnic background, and physiologic state (e.g., pregnancy, high-altitude adaptation). Some examples of these variations are shown in the Appendix. The existence of different reference ranges poses challenges for interpretation of results. In particular, creatinine stands out as an analyte for which conventional reference ranges are not always easy to apply in clinical practice. Plasma levels of creatinine vary with age, gender, and ethnic group. This fact makes it difficult in practice to use a simple reference

TABLE 480e-2 NATIONAL CHOLESTEROL EDUCATION PROGRAM ADULT TREATMENT PANEL III GUIDELINES FOR CHOLESTEROL

Cholesterol Type	Level (mg/dL)
LDL	
Optimal	<100
Near optimal/above optimal	100–129
Borderline high	130–159
High	160–189
Very high	\geq 190
Total	
Desirable	<200
Borderline high	200–239
High	\geq 240
HDL	
Low	<40
High	\geq 60

Abbreviations: HDL, high-density lipoprotein; LDL, low-density lipoprotein.

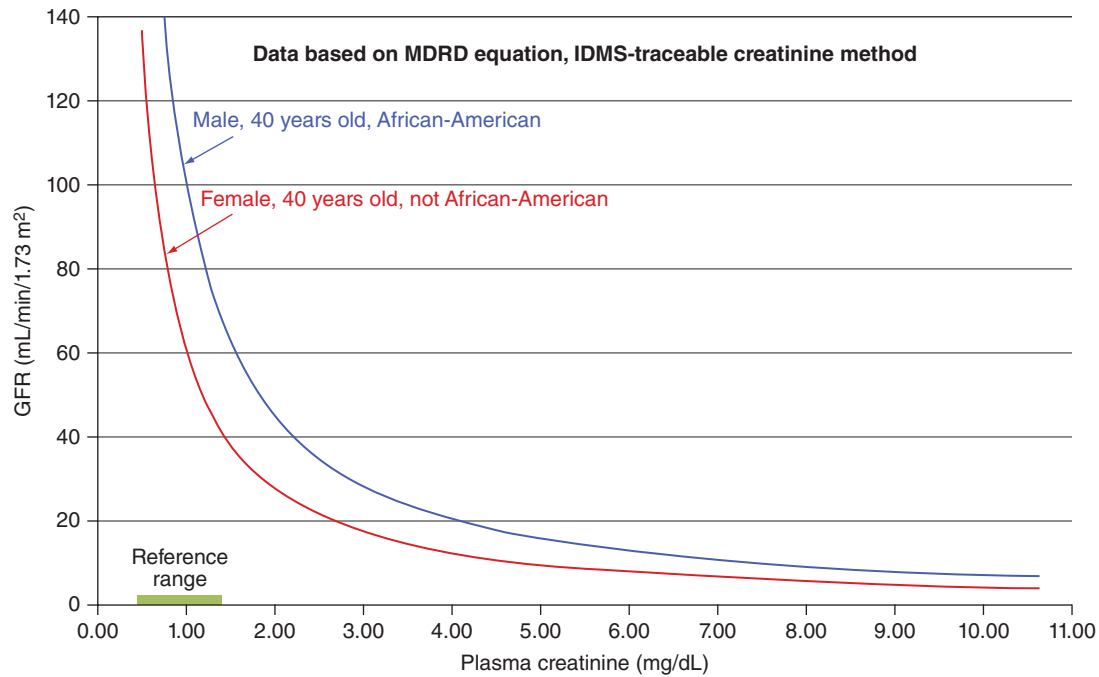


FIGURE 480e-2 Relationship between plasma creatinine and estimated glomerular filtration rate (eGFR) using the 4-parameter Modification of Diet in Renal Disease (MDRD) equation. IDMS, isotope dilution mass spectrometry.

range for this analyte when attempting to gauge a patient's renal function. A large decrease in glomerular filtration rate (GFR) is associated with slight increases in the plasma creatinine concentration within the typical reference range provided by many laboratories (Fig. 480e-2). A 60-year-old white woman with a serum creatinine level of 1.00 mg/dL, which is well within the typical reference range, has an estimated GFR of only 57 mL/min per 1.73 m², whereas the same creatinine concentration in a 20-year-old African-American male is consistent with normal renal function. To better estimate the GFR, which is widely considered to be the most useful index of overall renal function, it has become customary to use equations that incorporate plasma creatinine with other parameters. The most widely used of these equations in current practice is the 4-parameter Modification of Diet in Renal Disease (MDRD) equation that incorporates plasma creatinine, age, gender, and ethnic group (African American or not African American). The more recent CKD-EPI equation, which uses the same 4 parameters, is beginning to replace the MDRD equation. Recommended clinical laboratory practice is to report the estimated GFR with all creatinine measurements in adults. This addition provides more useful information than would a creatinine reference range alone.

SOURCES OF ERROR IN CLINICAL LABORATORY TESTING

Errors can arise at all stages of the testing process, from specimen collection to result interpretation. An error arising at any stage may adversely affect patient care. In clinical laboratory practice, it is customary to divide the testing process into three phases: preanalytic, analytic, and postanalytic. Examples of each type of error are shown in Table 480e-3. The most common error in the testing process is specimen mislabeling, in which a specimen from one patient is placed in a container labeled with another patient's name or identifiers. Specimen mislabeling errors may have very serious consequences for a patient. For example, if erroneous typing of a patient's blood group results from specimen mislabeling and is followed by transfusion of a mismatched unit of blood, the outcome may be fatal. A mislabeled biopsy specimen can lead either to an erroneous diagnosis and inappropriate therapy or to a failure to make a diagnosis and institute appropriate therapy.

In addition to errors, many preanalytic factors can influence clinical laboratory results. Posture (i.e., recumbent versus upright), exercise, diet, recently ingested food, and use of prescribed or recreational drugs (including tobacco, alcohol, caffeine, and herbal supplements)

can influence a variety of analyte concentrations. After blood has been collected, certain analytes undergo changes in their concentration during storage or transportation. Glucose levels fall as a result of red cell metabolism. Ammonia levels rise as a result of protein breakdown. Increasing permeability and breakdown of red cell membranes leads to increases in plasma potassium and free hemoglobin levels. Bacterial contamination can lead to overgrowth of specimens. To minimize

TABLE 480e-3 EXAMPLES OF PREANALYTIC, ANALYTIC, AND POSTANALYTIC ERRORS DURING THE LABORATORY TESTING PROCESS

Preanalytic Sources of Error

Test selection
Inappropriate test for the clinical need
Lack of clinical usefulness, regardless of possible results
Test order misunderstood or not communicated
Specimen collection
Incorrect time of collection
Patient not prepared for collection (e.g., not fasting)
Incorrect specimen type (e.g., wrong anticoagulant, wrong tissue fixative)
Use of incorrect specimen container
Insufficient specimen collected
Contamination of specimen by IV fluids, drugs, or bacteria
Specimen mislabeled or unlabeled
Important clinical information not provided
Delays in transportation to the lab, leading to alterations in specimen constituents

Analytic Sources of Error

Incorrect storage conditions prior to analysis
Specimen misidentification in the laboratory
Wrong test performed
Assay interferences
Assay failure (e.g., assay out of control)

Postanalytic Sources of Error

Delay in communication of assay results
Results not communicated to correct person
Incorrect result communicated
Misinterpretation of result

these precollection alterations, specimens should be processed or transported to the clinical laboratory as soon as possible after collection. The list of known preanalytic variables and their effects is extensive, and the reader is referred to the compendium on this subject (see Young DS: *Effects of Preanalytical Variables on Clinical Laboratory Tests*, 3rd ed. Washington, DC, AACC Press, 2007).

POINT-OF-CARE TESTING

The great majority of tests continue to be performed in dedicated clinical laboratory facilities, but for several decades there has been a trend toward point-of-care testing. This change has been made possible by the development of portable analytic devices, including single-purpose instruments such as glucometers and oxygen saturation monitors, and multifunction instruments that can perform a wider variety of analyses, particularly in chemistry and hematology but also in some areas of microbiology. The use of these devices is driven largely by the convenience of faster result availability. In some settings (e.g., in rural areas and developing countries), there may be no easily accessible clinical laboratory and a point-of-care device may be the best or only option for testing. However, the per-specimen cost of point-of-care testing, in terms both of reagents and supplies and of personnel, is often greater than that of centralized testing. Other concerns relate to the adequacy of personnel training for point-of-care testing, the quality of the results, and the incorporation of results into the medical record.

HOME TESTING BY PATIENTS

One of the largest markets for point-of-care testing is home testing by patients, which has long been an important element in the management of persons with diabetes who monitor their own blood glucose levels. Over-the-counter kits for home pregnancy testing have been available for decades. More recently, kits have become available for home testing of the international normalized ratio or prothrombin time by patients taking oral anticoagulants. Kits are also available for cholesterol monitoring, fecal occult-blood detection, and hemoglobin measurement. In these areas, there is often little information on the quality of test performance, the accuracy of the results, or the correctness of result interpretation.

ISSUES SPECIFIC TO GENETIC TESTING



The principles of genetic medicine in clinical practice are discussed in [Chaps. 82–84](#). Here we will concentrate on issues related to clinical laboratory testing for genetic disease.

The distinction between genetic testing for inherited disorders and that for acquired disorders affects the type of tissue that should be obtained for analysis. In inherited disorders, all nucleated cells are expected to carry the inherited mutation; thus white blood cells or buccal cells (obtained by scraping the inside of the cheek) are convenient sources of DNA for clinical laboratory testing. For prenatal testing of the fetus, chorionic villi or amniocytes are commonly used. In tests for acquired genetic disorders (e.g., in tumors), the tissue of interest that contains a suspected mutation must be sampled. It is often useful to compare tumor DNA with the patient's normal DNA in order to identify acquired mutations (e.g., testing for microsatellite instability in colorectal cancer; [Chap. 101e](#)).

INFORMED CONSENT FOR GENETIC TESTING

Although it is assumed that all clinical laboratory testing is performed with the consent of the patient (or, in the case of minors, the parents), regulations may require formal written consent for genetic testing. Such regulations vary among jurisdictions, and the practicing clinician should be aware of local regulations. In some jurisdictions, there are regulations on the storage and use of genetic information and on the maximal period for which genetic specimens may be stored.

For some late-onset genetic diseases, such as Huntington's disease ([Chap. 449](#)), genetic testing allows a prediction about the future development of the disease. The degree of certainty that is possible

on the basis of this testing surpasses that associated with identification of more traditional disease risk factors (e.g., hyperlipidemia as a risk factor for future myocardial infarction). When deciding to undertake predictive genetic testing, it is important for the patient to consider the broad implications of a positive or negative test result, to be made aware of any support and counseling that is available, and to understand the implications of a result for other family members. In dealing with these issues, genetic counselors play an important role ([Chap. 84](#)). Their expertise includes the ability to explain genetic disorders at an understandable level to patients and their families, to arrange for support services, and to provide genetic risk assessments to members of families with genetic disorders.

When testing for genetic disorders, the clinical laboratory will use different analytic approaches according to the disease of interest. Some disorders, such as sickle cell anemia, are caused by single-point mutations. Testing for these disorders involves mere assessment for one or a few mutations in a single gene. Other disorders (e.g., hyperphenylalaninemia) may be caused by numerous mutations in a single gene. Still others (e.g., hereditary breast cancer) may be caused by mutations in many genes. The number of possible mutations and genes that underlie a clinical phenotype affects the cost of and time required for clinical laboratory testing as well as the likelihood of finding a disease-causing mutation.

If a disease phenotype can be caused by many mutations, a clinical laboratory result that is negative should be interpreted with care. For example, it is common to screen healthy pregnant women (and their partners) for mutations in the *CFTR* gene, which is mutated in patients with cystic fibrosis (CF). The goal of this screening is to identify women who are carriers of a *CFTR* mutation and therefore are at increased risk of having a baby with CF. Because CF is an autosomal recessive disorder, a fetus has a 1:4 chance of being affected if both parents are carriers of disease-causing *CFTR* mutations. The screening test approach that is commonly used to identify mutations in carriers detects 80–85% of all known disease-causing *CFTR* mutations in Caucasians and up to 97% of mutations among Ashkenazi Jews. A negative screening result therefore does not completely eliminate the possibility that a woman (or her partner) actually has a mutation. What can be inferred from a negative test result is that the risk of having a CF-affected baby has decreased significantly to an extent that depends on the woman's ethnic group and the mutations that were examined. The clinical laboratory should calculate and report the woman's new risk of being a carrier if the screening result is negative.

The increasing availability of large-scale (next-generation) sequencing of a patient's whole genome or exome will greatly affect genetic testing over the next decade, with implications for the number of mutations that can be detected and the increased complexity of result interpretation.

LIMITATIONS TO MOLECULAR GENETIC TESTING

Genetic testing has limitations that are often unique to this field. Results may be inconclusive. For example, a search for mutations in a gene that is suspected of causing a disease may fail to reveal any known disease-causing mutations. A mutation may be discovered that is of unknown clinical significance. In this situation, consideration of the predicted change in the amino acid sequence of the encoded protein may suggest a biologic effect—e.g., replacement of a charged amino acid by one of the opposite charge or by a neutral amino acid; replacement of an amino acid by one of a different size; or replacement of an amino acid that is conserved across multiple species. Further information may be obtained by determining whether the mutation is found in healthy individuals. Even with all of these considerations, it is not uncommon that the biologic significance of an identified mutation remains uncertain, and further research may be needed to assess its significance.

It is also important to understand the limitations of the clinical laboratory approach used to detect mutations. At this time, next-generation sequencing remains an impractical undertaking for financial reasons, although extensive sequence analysis has become the

480e-6 standard of care for a few genes (e.g., analysis of *BRCA1* and *BRCA2* in assessing the risk of breast and ovarian cancer in individuals with a strong family history of these disorders). As sequencing technologies become less expensive, they can be expected to be more commonly used for both identifying mutations in patients with genetic disorders and screening asymptomatic individuals at risk of genetic disease.

Another unique aspect of genetic testing is the concern that genetic information about individuals may be used to discriminate against them by employers or by insurance companies. In the United States, the Genetic Information Nondiscrimination Act of 2008 (GINA) prohibits the use of genetic information by employers in making decisions related to employment and by health insurance companies in issuing insurance policies or setting premium rates based on knowledge of the applicant's genetic status. GINA does not cover disability insurance, long-term care insurance, or life insurance policies.

Although public attention has been most closely focused on DNA testing, other clinical laboratory investigations that are not usually thought of as genetic may provide important genetic information about the person being tested. For example, serum protein electrophoresis may reveal α -1 antitrypsin deficiency. Depending on the clinical laboratory technology used, measurement of hemoglobin A_{1c}, commonly used for monitoring diabetes control, may reveal a hemoglobin variant such as HbS (sickle cell). Measurement of cholesterol and triglyceride levels may reveal any of a number of hereditary disorders. All of these results are types of genetic information.

REGULATION OF THE CLINICAL LABORATORY

In the United States, all clinical laboratory testing performed for clinical purposes (but not for research purposes) is regulated by the federal Clinical Laboratory Improvement Amendments Act of 1988 (CLIA). Home monitoring by patients who are testing their own specimens is not covered by CLIA. The statute and the regulations, which are administered by the Centers for Medicare and Medicaid Services, apply to all laboratories, whether located in a physician's office, a large hospital, or a reference laboratory; and all laboratories are required to hold a valid CLIA certificate that is appropriate for the highest complexity level of the tests they perform. The U.S. Food and Drug Administration is responsible for assigning the complexity level of commercial tests. The lowest category of complexity is the "waived" category, which is followed (in order of increasing complexity) by the categories of "provider-performed microscopy," "moderate-complexity testing," and "high-complexity testing." The category of provider-performed microscopy is used to cover tests such as the use of potassium hydroxide preparations on skin scrapings to examine for fungi, fern tests, and sperm motility tests; it does not encompass histopathology that falls into the high-complexity category. Even if a clinical laboratory is performing only testing in the "waived" category, it must still hold a valid CLIA certificate. Laboratories that hold certificates for nonwaived tests are required to participate in proficiency testing and are regularly inspected to monitor their performance.

481e Clinical Procedure Tutorial: Central Venous Catheter Placement

Maria A. Yialamas, William E. Corcoran,
Gyorgy Frenzl, Kurt Fink

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these *Harrison's* **Chaps. 481e–486e**, video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

482e Clinical Procedure Tutorial: Thoracentesis

Charles A. Morris, Andrea Wolf

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these new *Harrison's* **Chaps. 481e–486e**, video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

483e Clinical Procedure Tutorial: Abdominal Paracentesis

Maria A. Yialamas, Anna E. Rutherford, Lindsay King

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these new *Harrison's* **Chaps. 481e–486e** video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

484e Clinical Procedure Tutorial: Endotracheal Intubation

Charles A. Morris, Emily Page Nelson

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these new *Harrison's* **Chaps. 481e–486e**, video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

**485e**

**Clinical Procedures Tutorial:
Percutaneous Arterial Blood
Gas Sampling**

Christian D. Becker

Medical Editors: Sean Sadikot, Jeremy Matloff

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these *Harrison's* [Chaps. 481e–486e](#), video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

486e Clinical Procedures Tutorial: Lumbar Puncture

Beth Rapaport, Stephen Krieger, Corey McGraw
Medical Editors: Sean Sadikot, Jeremy Matloff

Clinical procedures are an important component of medical student and resident training, and some are required for board and hospital certification. In these *Harrison's* **Chaps. 481e–486e**, video tutorials are presented for performing abdominal paracentesis, thoracentesis, endotracheal intubation, central venous catheter placement, percutaneous arterial blood gas sampling, and lumbar puncture. These videos have been created specifically for *Harrison's*. Each includes the indications, contraindications, equipment, potential complications, and related patient safety considerations. Additional video tutorials covering clinical procedures such as breast biopsy, IV line insertion, phlebotomy, arterial line insertion, arthrocentesis, bone marrow biopsy, pelvic examination, thyroid aspiration, basic suturing, and urethral catheterization are available to subscribers of *Harrison's Online* and AccessMedicine (available at www.accessmedicine.com).

Page numbers in **bold** indicate the start of the main discussion of the topic. Page numbers followed by “f” or “t” refer to the page location of figures and tables, respectively, and a “v” indicates a video on the DVD. Numbers containing “e-” followed by a number (e.g., 91e-3) refer to pages in the e-chapters on the DVD.

- A**
- AA amyloidosis, **723**
 characteristics of, 719
 clinical features of, 720t, 723, 1835t
 diagnosis of, 720f, 1845
 etiology of, 723
 incidence of, 723
 pathogenesis of, 1845
 treatment of, 723, 1845
- AAG (autoimmune autonomic ganglionopathy), 2642
- AAS abuse. *See* Anabolic-androgenic steroid (AAS) abuse
- AAV (adeno-associated virus), 91e-1t, 91e-3, 214e-4, 214e-9
- AAV-LPL, 91e-3
- AAV-RPE65, 91e-3
- Abacavir
 adverse effects of
 genetic considerations in, 373e-7, 373e-8t, 378, 1241
 in HIV infection, 1260, 1274t, 1277
 hypersensitivity, 382, 1277. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 genetic variations in response to, 38, 39t, 41
 for HIV infection, 1274t, 1277–1278
 molecular structure of, 1276f, 1277
 resistance to, 1278, 1279f
- A band, of sarcomere, 265e-5
- Abatacept, 2146t, 2147
- ABCA1 deficiency (Tangier disease), 1443, 2445, 2681
- ABCA1 gene, 291e-2
- ABCB11 gene mutations, 284
- ABCDEs, in nevi evaluation, 494
- ABCD rule, 76e-6
- ABCG5/G8 hemitransporter, 2075
- Abciximab
 action of, 746f, 748–749
 adverse effects of, 727, 727t, 749
 discontinuation before lumbar puncture, 443e-1
 dosage of, 749
 genetic variations in response to, 277t, 743t
 indications for, 749
 for myocardial infarction, 1598t
 in PCI, 296e-1
 pharmacology of, 749t
- Abdomen
 paradoxical movement of, 47e-3
 physical examination of, 286, 1443
- Abdominal aortic aneurysm. *See* Aortic aneurysm, abdominal
- Abdominal pain, **103**
 in acute intermittent porphyria, 2526
 in appendicitis, 1986–1987, 1987t
 approach to the patient, 103t, 105–107
 differential diagnosis of, 106t, 1876, 1877t
 in Fabry's disease, 432e-1
 history in, 106
 imaging in, 106–107
 in immunocompromised patients, 105
 in intestinal ischemia, 1979–1980
 in intestinal obstruction, 1983
 in irritable bowel syndrome, 1965
 in malabsorption syndromes, 1946t
 mechanisms of
 abdominal wall disorders, 104–105
 mechanical obstruction, 103–104, 104t
 metabolic, 104t, 105
 neurogenic, 105
 neurologic/psychiatric, 104t
 parietal peritoneal inflammation, 103, 104t
 toxic, 104t
 vascular disturbances, 104
 in pancreatitis, 106, 106t, 2092, 2099, 2101
 in pelvic inflammatory disease, 877
 in peptic ulcer disease, 1918
 in peritonitis, 846, 1989
 physical examination in, 106
 in pregnancy, 50
 referred, 105, 105t
- Abdominal paracentesis, 483e-1v
- Abdominal reflexes, 2539
- Abdominal swelling/distention, **285**, 286f, 1983.
See also Ascites
- Abdominal wall disorders, 104–105
- Abdominojugular reflex, 1444
- Abducens nerve, 2537
- Abducens nerve palsy, 209
- Abducens nerve paresis, 208
- Abduction and external rotation test, 1647
- A β amyloidosis, 719
- A β fibers, 87
- Abetalipoproteinemia (acanthocytosis), **2444**
 clinical features of, 269, 436e-4f, 2444
 genetic considerations in, 2444, 2630
 pathophysiology of, 1939, 2444, 2630
 peripheral blood smear in, 81e-4f
 small-intestinal mucosal biopsy in, 1939, 1940t
 treatment of, 96e-8, 2444
 vitamin E deficiency in, 96e-8
- A β_2 M amyloidosis, 719, 720t, 724. *See also* Amyloidosis
- A β peptides, 2600, 2600f
- A β /tau protein, 172t, 175
- ABI (ankle-brachial index), 1616, 1643
- Abiotrophia* spp., 821, 971
- Abiraterone, 103e-15, 586
- ABL gene mutations, 101e-3, 101e-3t, 689
- ABO blood group system, 138e-1
- Abortion
 infections following, 1165
 septic, 1060, 1099
 spontaneous, 83e-4
 threatened, 338
- ABPA. *See* Allergic bronchopulmonary aspergillosis (ABPA)
- Abruptio placenta, 46, 49
- Abscess
 anorectal, 1977–1978, 1977f
 brain. *See* Brain abscess
 epidural. *See* Epidural abscess
 formation of, 145e-10
 intraabdominal. *See* Intraabdominal abscess/infections
 periapical, 236
 peritonsillar, 233, 235, 1097
 perivalvular, 818
 retropharyngeal, 235
 subperiosteal, 231
 tuboovarian, 1089, 1099
- Abscopal effect, 412
- Absence seizures, 2543
- Absent (vanishing) testis syndrome, 2353
- Absolute risk reduction, 26
- Absorption
 of antibacterial drugs, 934–935
 of carbohydrates, 1937, 1937t
 disorders of. *See* Malabsorption syndromes of drugs, 33–34
 of lipids, 62f, 1935–1936, 1935t
 nutrient, **1933**
 of proteins, 1937
- Absorption spectrum, 385
- Abstract thought, assessment of, 2537
- Abulia, 1771, 2574f
- Acalculia, 179
- Acamprosate, 2059, 2728
- Acanthamoeba* spp., 245e-4t, 1367
- Acanthamoeba* spp. infections
 chronic meningitis, 908t
 clinical features of, 1367
 diagnosis of, 245e-4t, 245e-6t, 1367–1368
 epidemiology of, 245e-4t
 keratitis, 1368, 1368f
 meningoencephalitis, 896
 treatment of, 1368
- Acanthocytes, 81e-1
- Acanthocytosis. *See* Abetalipoproteinemia
- Acantholysis, 370
- Acanthosis, 372e-24t, 387, 1654f
- Acanthosis nigricans
 anti-insulin receptor antibodies in, 2348
 in diabetes mellitus, 2405, 2430
 disorders associated with, 76e-15, 359, 360t
 in gastric cancer, 535
 hyperpigmentation in, 76e-16f, 359
 in metabolic syndrome, 2452
 in obesity, 415e-6
- Acarbose, 1926, 2414t
- Accelerated idioventricular rhythm, 278e-12f, 1609
- Accelerated junctional rhythm, 1480–1481, 1609
- Accessory genome, 972
- Accessory pathways (APs)
 atrial fibrillation and, 1483
 in AV reentry tachycardia, 1481
 definition of, 1481
 in preexcited tachycardias, 1482, 1483f
 treatment of, 1482–1489
 in Wolff-Parkinson-White syndrome, 1481.
See also Wolff-Parkinson-White syndrome
- Accidents
 deaths from, 6e-1t, 6e-2f, 56t
 in health care industry, 12e-1, 12e-1f
- ACE. *See* Angiotensin-converting enzyme (ACE)
- Acebutolol, 473e-9t, 1480t, 1589t
- Aceclofenac, 1923
- ACE inhibitors. *See* Angiotensin-converting enzyme (ACE) inhibitors
- Acephalgic migraine, 2560, 2589
- Aceruloplasminemia, 96e-10
- Acetaminophen
 adverse effects of
 hepatotoxicity, **2027**
 clinical features of, 2025t
 dose-related, 2028f
 liver biopsy in, 366e-4f
 pathogenesis of, 43–44
 metabolic acidosis, 320
 nephrotoxicity, 92
 thrombocytopenia, 727t
 for back pain, 118, 119t
 drug interactions of, 103e-10t, 205e-3, 2232t
 for fever, 126
 for influenza, 1214
 metabolism of, 2027
 for migraine, 2592t
 for osteoarthritis, 2231, 2232t
 overdose/poisoning with, 2027, 2764t
 for pain, 60, 91–92, 91t
 for SLE-related pain, 2132
 for spinal stenosis, 116
 for tension-type headache, 2594
 therapeutic range of, 2764t

- Acetaminophen-aspirin-caffeine, 2590, 2592t
 Acetate, 98e-5t
 Acetazolamide
 action of, 332e-7
 for acute angle-closure glaucoma, 200
 adverse effects of, 297, 380
 for altitude illness, 476e-2, 476e-2t, 476e-4, 476e-5
 for episodic ataxia, 2630
 for hyperkalemia, 64e-3
 for hyperkalemic periodic paralysis, 462e-18
 for hypokalemic periodic paralysis, 462e-17
 for metabolic alkalosis, 322
 for paroxysmal symptoms in MS, 2672
 for pseudotumor cerebri, 203
 for raised CSF pressure headache, 110
 for salicylate-induced acidosis, 319
 for tumoral calcinosis, 426e-8
 for uric acid nephropathy, 431e-5
 Acetoacetate, 318
 Acetohydroxamic acid, 1871
 Acetylcholine (ACh)
 action of, 444e-3t, 2637
 clinical aspects of, 444e-3t
 deficiency of, in delirium, 167
 in neural control of gastrointestinal system, 264
 structure of, 444e-3t
 in swallowing, 255
 Acetylcholine receptor (AChR)
 in congenital myasthenic syndromes, 2702, 2703t
 in myasthenia gravis, 372e-24t, 377e-4, 2701, 2701f
 Acetylcholinesterase (AChE)
 deficiencies of, 2702, 2703t
 in nerve agent exposure, 262e-4, 262e-4f
 test, in myasthenia gravis, 2702
 Acetylcholinesterase (AChE) inhibitors
 for gastroparesis, 260t, 261
 overdose/poisoning with, 473e-10t
 for snakebite treatment, 2737, 2737t
 N-Acetylglutamate synthase deficiency, 434e-3t
 N-Acetyltransferase, 34t
 ACh. *See* Acetylcholine (ACh)
 Achalasia, **1904**
 chronic, esophageal cancer and, 532
 clinical features of, 1904
 diagnosis of, 1904, 1905f
 differential diagnosis of, 1904
 dysphagia in, 256
 LES relaxation impairment in, 1904, 1905f
 subtypes of, 1905f
 treatment of, 258, 1905
 AChE. *See* Acetylcholinesterase (AChE)
 Achilles bursitis, 2247
 Achlorhydria, 350e-1
 Achondrodysplasia, 426e-7
 Achondroplasia, 2512
 AChR. *See* Acetylcholine receptor (AChR)
 Achromatopsia, 182, 197
Achromobacter xylooxidans, 183e-2, 183e-2t
 Acid anhydride exposure, 1692, 1692t
 Acid-base disorders, **315**. *See also specific disorders*
 anion gap in, 64e-8, 317
 approach to the patient, 317, 317t
 mixed, 64e-1, 64e-8, 316-317, 316t
 simple, 315-316
 Acid-base homeostasis, 315
 Acid-base nomogram, 316, 316f
 Acidemia, 309, 434e-1, 2092
 Acid-fast bacteria, 150e-1-2
 Acid fume exposure, 1692t
 Acid maltase (acid α -glucosidase) deficiency.
 See Pompe disease
 Aciduria, orotic, 431e-5t, 431e-6, 431e-6f
Acinetobacter spp., 150e-2t, 934, 1036
Acinetobacter spp. infections, **1036**
 antibiotic resistance in, 1037
 approach to the patient, 1037
 burn-wound, 166e-1
 in cancer patient, 486t, 487
 clinical features of, 1037
 complications of, 1038
 diagnosis of, 1037
 epidemiology of, 1036
 etiology of, 1036
 global considerations in, 1036, 1037
 health care-associated pneumonia, 803t, 914
 pathogenesis of, 1036-1037
 prevention of, 1038
 prognosis of, 1038
 in returning war veterans, 152e-5t, 152e-9
 treatment of, 152e-5t, 152e-9, 940t, 1037-1038, 1037t
 urinary tract, 862
 ventilator-associated pneumonia, 811, 915
 ACIP (Advisory Committee on Immunization Practices), 786
 Acitretin
 adverse effects of, 348t, 354, 380
 for pityriasis rubra pilaris, 354t
 for psoriasis, 348, 348t
 aCL (anticardiolipin antibodies), 2131, 2135t, 2754t
 ACLS (advanced cardiac life support), 1769-1770, 1769f
 ACMV (assist control ventilation), 1741, 1742t.
 See also Mechanical ventilation
 Acneiform eruptions, 356, 356t
 Acne rosacea, **352**
 clinical features of, 76e-1f, 341t, 342f, 352, 352f
 treatment of, 352
 Acne vulgaris, **351**
 in 21-hydroxylase deficiency, 2355
 clinical features of, 76e-1f, 341t, 342f, 351-352, 352f
 drugs exacerbating, 378
 folliculitis in, 829
 pathophysiology of, 351
 treatment of, 352
 Acotiamide, 263
 Acoustic neuroma. *See* Vestibular schwannoma (acoustic neuroma)
 Acoustic reflex, 222
 Acoustic reflex decay, 223
 ACP Journal Club, 25t
 Acral angiodermatitis, 367
 Acral erythema, drug-induced, 380
 Acrocephalopolysyndactyly (Pfeiffer syndrome), 435
 Acrochordon, 341t, 342f, 359
 Acrocyanosis, 1442, 1648f, **1649**, 1754
 Acrodermatitis chronica atrophicans, 1151
 Acrodermatitis enteropathica, 96e-9
 Acroline exposure, 1692t
 Acromegaly, **2269**
 clinical features of, 359, 462e-19, 2269, 2270f
 arthropathy, 2240-2241
 cardiac, 290e-1t, 290e-3
 hirsutism, 332
 hypertension, 1620
 oral, 236
 diagnosis of, 2269
 etiology of, 2240, 2269, 2269t
 familial, 2266
 in GRFoma, 571
 laboratory investigation of, 2269
 vs. pachydermoperiostosis, 426e-7
 paraneoplastic, 609t
 screening tests for, 2262t
 treatment of, 2269-2271, 2270f
 Acro-osteolysis, 2160, 2160f
 Acropachy, thyroid, 2245, 2295, 2295f
 Acroparesthesia, in Fabry's disease, 432e-1
 Acrylamide, 2688t
 Acta, 982
 ACTA1 gene, 462e-12
 ACTH. *See* Adrenocorticotropic hormone (ACTH)
 ACTHoma, 559t, 571
 ACTH stimulation test, 2310
 Actin, 265e-5, 265e-5f, 372e-24t, 1555t, 1556f
 Actinic dermatitis, 389
 Actinic keratosis
 clinical features of, 76e-6f, 341t, 342f, 501f
 malignant conversion of, 387, 501
 sun avoidance and, 476
 treatment of, 386
Actinobacillus spp. *See* *Aggregatibacter* spp.
Actinomyces spp., 150e-2t, 1088. *See also* Actinomycosis
 Actinomycosis, 1089
 Actinomyces-like organisms, 1089
Actinomyces pyogenes, 981
 Actinomycetoma, nocardial, 1085-1086, 1086f, 1087t. *See also* *Nocardia* spp. infections
 Actinomycosis, **1088**
 clinical features of
 abdominal, 1089
 CNS, 907t, 1089
 disseminated, 1090
 in dog-bite wound, 167e-1
 hepatic-splenic, 1089, 1090f
 musculoskeletal and soft tissue, 1089-1090
 oral-cervicofacial, 239t, 241, 1088, 1089f, 1097
 pelvic, 1089, 1090f
 thoracic, 1088-1089, 1089f
 diagnosis of, 150e-1, 150e-2t, 1090
 epidemiology of, 1088
 etiology of, 1088
 pathogenesis of, 1088
 treatment of, 1090-1091, 1091t
 Action Plan to Prevent Healthcare-Associated Infections, 912, 913t
 Action potentials, cardiac
 electrophysiology of, 273e-1-2, 273e-1f
 phases of, 265e-6
 regional differences in, 1466, 1466f, 1467f
 Action spectrum, 385
 Activase. *See* Alteplase
 Activated charcoal
 for indigestion, 263
 for paralytic shellfish poisoning, 2743
 for poisoning/drug overdose, 473e-6, 473e-7
 Activated partial thromboplastin time (aPTT)
 causes of abnormal values, 406t
 in coagulation disorders, 732, 733t, 735
 coagulation factors tested in, 405-406, 406f, 733f
 LA-PTT, 406
 for monitoring heparin therapy, 750-751
 in preoperative testing, 405
 prolonged, 406t
 reagent composition of, 406
 reference values for, 2755t
 Activated protein C, recombinant, 1758
 Activator protein-1, in asthma, 1674
 Active transport, 332e-6, 1933
 Activin, 401e-4
 Activities of daily living (ADLs), 72t, 78t
 Acupressure, 14e-1t, 261, 1880
 Acupuncture, 14e-1t, 14e-3, 120, 338
 Acustimulation, 1880
 Acute brain failure. *See* Delirium
 Acute chest syndrome, 634, 635-636
 Acute confusional state. *See* Delirium
 Acute coronary syndromes
 diagnosis of, 1599, 1599f
 evaluation and management of, 1595, 1595f
 hyperbaric oxygen therapy for, 477e-4t
 myocardial infarction
 non-ST-segment elevation. *See* Non-ST-segment elevation acute coronary syndrome (NSTE-ACS)
 ST-segment elevation. *See* ST-segment elevation myocardial infarction (STEMI)
 PCI for, 296e-4. *See also* Percutaneous coronary interventions (PCI)
 pulmonary edema in, 1764
 unstable angina, 96, 96t, 97, 1593. *See also* Non-ST-segment elevation acute coronary syndrome (NSTE-ACS)
 Acute disseminated encephalomyelitis (ADEM), 441e-26f, **2673**

- Acute febrile illness, 779
 approach to the patient, 779
 clinical presentations of, 764, 780t, 781–783
 diagnostic workup in, 779
 history in, 779
 physical examination in, 779
 rashes associated with, 25e-1
 treatment of, 779–781, 780t
- Acute febrile neutrophilic dermatosis. *See* Sweet syndrome
- Acute generalized eruptive/exanthematous pustulosis (AGEP)
 clinical features of, 132t, 384t
 pustular lesions, 356, 382, 382f
 rash, 25e-12f, 132t
 differential diagnosis of, 135, 382
 drugs associated with, 382
 immune pathways of, 378t
 nondrug related causes of, 384t
- Acute HIV syndrome, 1228, 1249, 1249f. *See also* HIV infection
- Acute inflammatory demyelinating polyneuropathy (AIDP), 2695, 2695t, 2696t
- Acute intermittent porphyria (AIP), 2526
 clinical features of, 2522t, 2526, 2529, 2681
 diagnosis of, 2522t, 2529
 genetic considerations in, 2681
 global considerations in, 2521
 pathophysiology of, 2522f, 2681
 treatment of, 367e-3, 2529–2530
 unsafe drugs in, 2526, 2527–2529t
- Acute interstitial nephritis (AIN), 1856
 allergic, 1804, 1857, 1857f
 clinical features of, 62e-10f, 289t, 1806t, 1856, 1857
 in crystal deposition disorders, 1858–1859
 diagnosis of, 289t, 1806t
 etiology of, 1856t
 granulomatous, 1857f, 1858
 hyperkalemia in, 307
 hypovolemia in, 297
 idiopathic, 1858
 IgG4-related, 391e-2, 1858
 infection-associated, 1858
 in light chain cast nephropathy, 62e-7f, 1859, 1859f
 in lymphomatous infiltration of the kidney, 1859
 in obstructive tubulopathies, 1858–1859
 renal biopsy in, 62e-10f
 in Sjögren's syndrome, 1857
 in SLE, 1858. *See also* Lupus nephritis
 treatment of, 1857f, 1857t
 in tubulointerstitial nephritis with uveitis, 1857–1858, 1858f
- Acute interstitial pneumonia (Hamman-Rich syndrome), 1712
- Acute kidney injury (AKI), 1799
 adaptive response to, 333e-5
 after liver transplantation, 2072
 approach to the patient, 290f, 291
 clinical features of, 289t, 1805, 1806t
 fluid volume disturbances, 297
 hyperkalemia, 304, 307
 hyponatremia, 299
 complications of, 1808–1809
 anemia and bleeding, 1809, 1810
 fluid volume disturbances, 1808, 1810
 hyperkalemia, 1808
 hyperphosphatemia, 1807, 1808–1809, 1810
 hypocalcemia, 1807
 malnutrition, 1809, 1810
 metabolic acidosis, 1808, 1810
 uremia, 1808
 in critically ill patient, 1735
 diagnosis of
 biomarkers, 1808
 blood laboratory findings, 291t, 1805, 1806t, 1807
 history and physical examination, 1805
 imaging, 1808
 renal biopsy, 62e-10f, 1808
 renal failure indices, 1807–1808
 urine findings, 291t, 1805, 1806t, 1807f
 epidemiology of, 1799
 etiology and pathophysiology of
 envenomation, 2737
 hypokalemia, 307
 intrinsic, 292, 1800, 1800f, 1802f
 ischemia-associated, 292, 1801–1802, 1803f, 1806t
 nephrotoxin-associated, 292, 1803–1804, 1806t
 postrenal, 291–292, 1804–1805, 1804f, 1806t
 prerenal, 291, 1800–1801, 1800f, 1801f, 1806t
 sepsis-associated, 1755, 1806t
 uric acid nephropathy, 431e-4, 431e-5
 global considerations in, 1799
 incidence of, 1799
 in leptospirosis, 1144
 maladaptive response to, 333e-5
 in malaria, 1374, 1374t, 1381
 preconditioning and, 333e-3–4, 333e-4t
 prevention of, 1809
 prognosis of, 1811
 in sarcoidosis, 2208
 in scleroderma renal crisis, 62e-9f, 2158, 2162, 2164
 in shock, 1746
 in transplant recipient, 1829
 treatment of
 in cirrhosis and hepatorenal syndrome, 1809–1810
 dialysis indications and modalities, 1810–1811
 intrinsic, 1810
 postrenal, 1810
 prerenal azotemia, 1809
 principles of, 1809, 1809t
 supportive measures, 1809t, 1810
- Acute lung injury (ALI). *See also* Acute respiratory distress syndrome (ARDS)
 pathophysiology of, 1736, 1737f
 in shock, 1746
 transfusion-related, 138e-5
 ventilator-induced, 1738
- Acute lymphoid leukemia (ALL)
 approach to the patient, 699–700
 classification of, 695, 695t
 epidemiology of, 695, 696f
 etiology of, 695
 genetic considerations in, 101e-3t, 699t
 hyperleukocytosis in, 1793
 immunology of, 698f
 infections in, 484t, 485, 486f, 486t, 488, 489
 late consequences of, 622, 624t
 lymphadenopathy in, 408
 peripheral blood smear in, 81e-9f
 precursor B cell, 701, 701f
 precursor T cell, 702t, 707
 prevention of, 643
 prognosis of, 700
 treatment of, 103e-22, 139e-5–6, 139e-5t
- Acute megakaryoblastic leukemia, 135e-7
- Acute motor axonal neuropathy (AMAN), 2695, 2695t, 2696t
- Acute motor sensory axonal neuropathy (AMSAN), 2695, 2695t, 2696t
- Acute mountain sickness (AMS), 476e-1–3, 476e-2t
- Acute myeloid leukemia (AML), 678. *See also* Acute promyelocytic leukemia (APL)
 cancer stem cells in, 102e-11
 classification of, 678–679, 679t
 clinical features of, 681
 diseases associated with, 678
 etiology of, 678
 genetic considerations in, 101e-2t, 373e-5, 679, 679t
 hyperleukocytosis in, 1793
 incidence of, 678
 infections in, 484t, 486t, 488, 685–686
 monocytopenia in, 422
 with mutated CEBPA, 679
 with mutated nucleophosmin, 679
 myelodysplasia and, 669
 peripheral blood smear in, 81e-9f, 681, 682f
 pretreatment evaluation of, 681–682, 683t
 prognosis of, 679–681, 680t
 as second malignancy in cancer survivors, 623, 678
 skin manifestations of, 366, 369
 treatment of
 agents under study for, 685t
 algorithm for, 682–684, 684f
 HSCT, 139e-5, 139e-5t, 684–685
 late consequences of, 624t
 postremission therapy, 684
 relapse, 686
 supportive care, 685–686
- Acute myelomonocytic leukemia, 237, 239t
- Acute myocardial infarction (AMI). *See* Myocardial infarction (MI)
- Acute necrotic collection, pancreatic, 2093t
- Acute necrotizing myopathy, 614t, 619, 1784
- Acute nephritic syndromes, 1835, 1835t, 1837
 ANCA small vessel vasculitis, 1835t, 1840
See also Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome); Granulomatosis with polyangiitis (Wegener's); Microscopic polyangiitis
 antiglomerular basement membrane disease. *See* Antiglomerular basement membrane (anti-GBM) disease
 cryoglobulinemia. *See* Cryoglobulinemia
 endocarditis-associated glomerulonephritis, 1835t, 1837–1838
 Goodpasture's syndrome. *See* Goodpasture's syndrome
 IgA nephropathy. *See* IgA nephropathy
 IgA vasculitis. *See* IgA vasculitis (Henoch-Schönlein)
 lupus nephritis. *See* Lupus nephritis
 membranoproliferative glomerulonephritis. *See* Membranoproliferative glomerulonephritis
 poststreptococcal glomerulonephritis. *See* Poststreptococcal glomerulonephritis
 renal biopsy in, 62e-1–6f
- Acute pancreatitis, 2091
 in alcohol use disorder, 2091, 2725
 approach to the patient, 2086, 2092
 clinical features of, 2094
 abdominal pain, 106, 106t, 2092
 acute kidney injury, 1803
 fat necrosis, 366
 hypocalcemia, 2482
 hypovolemia, 298
 nausea and vomiting, 258
 complications of, 2096–2097, 2097t
 diagnosis of
 comparison of imaging methods, 2087t, 2088–2089, 2094–2095
 CT, 2093t, 2094f, 2095f
 endoscopy, 1899, 2087t
 ERCP, 2095f
 laboratory tests, 2087, 2087t, 2092–2093
 differential diagnosis of, 2093
 drug-related, 1273, 2092
 epidemiology of, 2086
 etiology of, 2091–2092, 2091t
 gallstone, 1893–1894
 genetic considerations in, 2092
 in HIV infection, 1258, 2097–2098
 incidence of, 2091
 interstitial, 2092, 2093t, 2095
 mild, 2094
 morphologic features of, 2093t
 necrotizing. *See* Necrotizing pancreatitis
 pancreatic enzyme activation in, 2092
 pathogenesis of, 2092
 phases of, 2093–2094
 physical examination in, 2092
 recurrent, 2097
 severe, 2094, 2094f, 2095f, 2095t
 treatment of, 2095–2096

- Acute pharyngeal cervicobrachial neuropathy, 2696t
- Acute promyelocytic leukemia (APL)
chemotherapy-associated, 103e-13
chromosome abnormalities in, 102e-7, 431t, 679
peripheral blood smear in, 81e-9f
treatment of, 102e-5t, 102e-7, 103e-16, 686-687
- Acute quadriplegic neuropathy, 462e-20
- Acute radiation syndrome, 263e-2-4, 263e-4f
- Acute renal failure. *See* Acute kidney injury (AKI)
- Acute respiratory distress syndrome (ARDS), **1736**
clinical course of, 1736-1738, 1736f, 1737f, 1738f
diagnosis of, 1736t
chest radiography, 308e-10f, 1737, 1737f
CT, 308e-10f, 1737, 1738f
disorders associated with, 1736, 1736t
etiology of, 1736
functional recovery after, 1740
HSV infection and, 1180
incidence of, 1736
pathophysiology of, 1736-1738, 1737f
pressure-volume relationships in, 1732f
prognosis of, 1739-1740
in sepsis/septic shock, 1755
in shock, 1746
treatment of, 1738-1739, 1739f, 1739t. *See also* Mechanical ventilation
- Acute rheumatic fever. *See* Rheumatic fever
- Acute tubular necrosis (ATN). *See* Acute kidney injury (AKI)
- Acyclovir
action of, 214e-4, 215e-6, 896-897
adverse effects of
gastrointestinal, 897
hematologic, 897
musculoskeletal, 2218t
neurologic, 215e-7, 897, 2547t
renal, 215e-7, 1181, 1804
for Bell's palsy, 2648
global considerations in, 1183
for herpes zoster, 215e-3-4t, 215e-7, 832t, 1186, 2657
for HSV infections
encephalitis, 215e-2t, 1179, 1181, 1182t
esophagitis, 1182t, 1910
genital, 215e-2t, 215e-7, 832t, 1182t
mucocutaneous, 215e-7, 233, 1182t
in neonates, 1180, 1182t
ocular, 200
pharyngitis, 233
as prophylaxis in HIV infection, 1253t
as prophylaxis in transplant recipient, 139e-4t, 215e-6, 921, 925t, 927
for oral hairy leukoplakia, 1190
pharmacology of, 215e-7
resistance to, 215e-7, 897, 1182-1183
for varicella prevention, 917
for varicella treatment, 215e-2t, 215e-7, 1185-1186
for viral encephalitis, 896-897
for viral meningitis, 893
for viral myelitis, 2657
for VZV prophylaxis in HIV infection, 1252t
for VZV prophylaxis in transplant recipient, 139e-4
- AD. *See* Alzheimer's disease (AD)
- Adalimumab
adverse effects of, 348t, 422, 2146t, 2173
for ankylosing spondylitis, 2172-2173
for autoimmune and inflammatory diseases, 372e-30
for IBD, 1961
monitoring during treatment with, 2146t
for psoriasis/psoriatic arthritis, 348t, 2177
for rheumatoid arthritis, 2145, 2146t
for sarcoidosis, 2212
- Adamantinoma, 119e-3
- ADAMTS13, 729, 1863, 2754t
- ADAMTS13 metalloproteinase, 1848
- Adaptive immune system, 372e-2, **372e-16**. *See also* Immune response/immune system
cells triggering, 372e-7t
definition of, 372e-1
immunoglobulins in. *See* Immunoglobulin(s)
innate immune system and, 372e-2
intercellular interactions of, 372e-9f
primary immunodeficiencies of
B lymphocyte deficiencies. *See* B cell(s), deficiencies of
T lymphocyte deficiencies. *See* T cell(s), deficiencies of
PRRs in modulation of, 372e-6t
response to microbes, 372e-23, 372e-26
T cells in. *See* T cell(s)
- Adaptive senescence, 94e-3
- Adaptive servoventilation, 1727
- ADCC (antibody-dependent cell-mediated cytotoxicity), 372e-1, 372e-6, 1244
- Addison's disease. *See also* Adrenal insufficiency
anemia in, 630
APS-1 and, 2345, 2346t
APS-2 and, 2346, 2346t, 2347t
autoantibodies in, 372e-24t, 2346
clinical features of, 2324, 2326f
diarrhea in, 268
genetic considerations in, 2346
hyperkalemia in, 310
hypoglycemia in, 2433
oral manifestations of, 240t
skin manifestations of, 359, 360
thymic tumors in, 2348
- Adefovir
action of, 215e-9
adverse effects of, 215e-9, 2035
for chronic HBV infection, 215e-4t, 215e-9, 2035, 2038f, 2074
for HBV prophylaxis in transplant recipient, 927
resistance to, 2040t
- A-delta (A δ) fibers, 87, 88
- ADEM (acute disseminated encephalomyelitis), 441e-26f, **2673**
- Adenine, 426
- Adenine nucleotide translator (ANT), 372e-24t
- Adenine phosphoribosyltransferase (APRT)
deficiency, 431e-3t, 431e-5-6
- Adeno-associated virus (AAV), 91e-1t, 91e-3, 214e-4, 214e-9
- Adenocarcinoma
bladder, 575. *See also* Bladder cancer
cervix, 595. *See also* Cervical cancer
colon. *See* Colorectal cancer
endometrial. *See* Endometrial cancer
esophagus, 532-533, 533t. *See also* Esophageal cancer
gastric. *See* Gastric adenocarcinoma
lung, 507, 507f. *See also* Lung cancer
small intestinal, 537
of unknown primary, 120e-3, 120e-3f
- Adenocarcinoma in situ, 515-516
- Adenoidectomy, 230
- Adenoma
ACTH-producing. *See* Cushing's syndrome
aldosterone-producing adrenal, **2318**. *See also* Mineralocorticoid excess
clinical features of, 462e-19
diagnosis of, 1619
etiology of, 2318-2319, 2318t
incidence of, 306, 2321t
hepatic, 553-554
parathyroid, 2470. *See also* Hyperparathyroidism
pituitary. *See* Pituitary tumors (adenomas)
polypoid, 536
small intestinal, 536
thyroid, 2303-2304
- Adenoma sebaceum, 359, 364
- Adenomyomatosis, 2083
- Adenosine
adverse effects of, 1467t, 1471t, 1480t, 1484
for arrhythmias, 1479t
dosage of, 1479t
pharmacology of, 1479t
for preexcited tachycardias, 1482
for supraventricular tachycardia, 35, 47, 1483-1484
as vasoconstrictor, 332e-2
for ventricular tachycardia, 1494
- Adenosine deaminase, 288
- Adenosine deaminase deficiency, 91e-2, 375e-1, 431e-3t, 2108
- Adenosine kinase deficiency, 434e-2t
- 5' Adenosine monophosphate-activated protein kinase (AMPK), 94e-5, 102e-13
- Adenosine receptors, 276, 742
- S-Adenosylhomocysteine hydrolase deficiency, 434e-2
- Adenovirus(es)
adherence of, 145e-2t
characteristics of, 1208
enteric, 1286f, 1286t, 1289
fidelity of replication in, 214e-5
gene expression and regulation in, 214e-4
for gene therapy delivery, 91e-1t, 214e-4, 214e-9
host defense inhibition by, 214e-8
laboratory identification of, 214e-9
structure of, 214e-2t, 214e-3f
- Adenovirus infections, **1208**
clinical features of, 1208
conjunctivitis, 199, 1208
diarrhea, 266, 1208, 1289
rash, 362, 362t
upper respiratory, 226, 231, 234, 1202t
diagnosis of, 1208
epidemiology of, 1208
in immunocompromised patient, 1208
prevention of, 1209
in transplant recipient, 920t, 925, 1208
treatment of, 1208
- Adenovirus vaccine, 1209
- Adenylate kinase (AK) deficiency, 654t
- Adenylosuccinate lyase deficiency, 431e-3t, 431e-6
- Adequate intake, 95e-1, 95e-3
- ADH (antidiuretic hormone). *See* Arginine vasopressin (AVP)
- Adhesins, 145e-1
bacterial, 145e-2-3, 145e-2f, 145e-2t, 1043
fungal, 145e-2t, 145e-3
protozoa, 145e-2t, 145e-3
viral, 145e-1-2, 145e-2t
- Adhesive arachnoiditis, lumbar, 117
- Adhesive capsulitis, 2249
- ADHH (autosomal dominant hypocalcemic hypercalciuria), 2483
- Adie's syndrome, 196
- Adipocytes, 415e-2, 415e-3f, 2405
- Adiponectin, 415e-2, 2452
- Adipose stem cells, 453t
- Adipose tissue, 415e-2
- Adipostat, 415e-5
- Adipsin, 415e-2
- ADLs (activities of daily living), 72t, 78t
- Adnexal mass/tumor, 338, 364, 364t, 593, 595
- ADNFLE (autosomal dominant nocturnal frontal lobe epilepsy), 2545t
- Adolescence
ankylosing spondylitis in, 2170
iron requirements in, 626
seizures in, 2546-2547, 2546t
smoking in, 475
sudden cardiac death in, 1767
- Ado-trastuzumab emtansine, 102e-4, 102e-5t, 103e-22
- ADPEAF (autosomal dominant partial epilepsy with auditory features), 2545t
- ADPKD (autosomal dominant polycystic kidney disease). *See* Polycystic kidney disease, autosomal dominant
- ADP receptor antagonists, 747-748. *See also* Clopidogrel; Prasugrel; Ticagrelor
- Adrenalectomy
for adenoma, 1619
for breast cancer, 103e-3, 530t
for Cushing's syndrome, 2273, 2318

- for ectopic ACTH syndrome, 611
for mineralocorticoid excess, 2319
for pheochromocytoma, 2330
- Adrenal gland**
anatomy and development of, 2309–2310
disorders of
adrenal insufficiency. *See* Adrenal insufficiency
congenital adrenal hyperplasia. *See* Congenital adrenal hyperplasia (CAH)
Cushing's syndrome. *See* Cushing's syndrome
masses
adrenocortical carcinoma. *See* Adrenocortical carcinoma
classification of, 2321t
incidentally discovered, 2321, 2321t, 2322f
in MEN 1, 2339
pheochromocytoma. *See* Pheochromocytoma
mineralocorticoid excess. *See* Mineralocorticoid excess
steroidogenesis in, 2309, 2309f
activation/inactivation of, 2311–2313, 2312f, 2313f
regulatory control of, 2310–2311, 2310f, 2311f, 2314f
- Adrenal hyperplasia**
congenital. *See* Congenital adrenal hyperplasia (CAH)
hypokalemia in, 306
macronodular, 609t
primary, 306, 1619
- Adrenal hypoplasia congenita**, 2324t, 2364
- Adrenal insufficiency**, 2323
acute, 2324
AV conduction block in, 1471t
in cancer, 1795
chronic, 2324
clinical features of, 2324, 2325t, 2326f
fatigue, 152
hyperkalemia, 64e-2, 310
hyponatremia, 299
hypothermia, 478e-1, 478e-3
muscle weakness, 462e-19
shock, 1750
diagnosis of, 2325, 2327f
epidemiology of, 2323
etiology of, 2323–2324, 2324t, 2325t
HLA gene association with, 373e-8t
primary, 2323–2324, 2324t. *See also* Addison's disease
secondary, 2324, 2325t
in sepsis/septic shock, 1755, 1757
treatment of, 2325–2327, 2327f
- Adrenal vein sampling**, 1619, 2319
- Adrenarche**, 2357
- Adrenergic crisis**, 1626t, 1627
- Adrenocortical carcinoma**
aldosteronism in, 1619
classification of, 2322
diagnosis of, 2320, 2323f
genetic considerations in, 2318t, 2321–2322
incidence of, 2321, 2321t
in MEN 1, 2339
staging of, 2322, 2323t
treatment of, 103e-16, 2322–2323
- Adrenocorticotrophic hormone (ACTH)**, 401e-4
action of, 400e-2, 401e-4
in adrenal steroidogenesis control, 2310–2311, 2310f, 2312f
autoantibodies to, 372e-24t
deficiency of, 372e-24t, 2259t, 2260
ectopic production of, 610
clinical features of, 64e-8, 611
diagnosis of, 611, 2272t
epidemiology of, 2314t
etiology of, 64e-8, 609t, 610–611, 2314
in GI-NETs, 564
in lung cancer, 511
skin manifestations of, 360
treatment of, 611, 2318
eutopic production of, 608
excess of. *See* Cushing's syndrome
expression and regulation of, 401e-1t
laboratory evaluation of, 2258t, 2272, 2325
secretion of, 401e-4, 2310
in shock, 1745
stimulation test, 1750
synthesis of, 401e-4, 2310
- Adrenoleukodystrophy**
clinical features of, 2324t, 2608, 2680
diagnosis of, 441e-41f, 2325, 2608
gene therapy for, 91e-2–3
genetic considerations in, 2323, 2324t, 2680
Adrenomyeloneuropathy, 2659, 2680–2681
- ADT. See** Androgen deprivation therapy (ADT)
- Adult bile ductopenia**, 284
- Adult growth hormone deficiency**, 2259–2260, 2259t, 2260f
- Adult stem cells**, 453
- Adult T-cell leukemia/lymphoma**, 707
acute, 225e-4–5
chronic, 225e-5
clinical features of, 225e-4–5, 707
diagnosis of, 81e-10f, 707, 707f
genetic considerations in, 101e-3, 101e-3t
HTLV-1 and, 214e-7, 225e-4, 697, 707
lymphomatous, 225e-5
paraneoplastic syndromes in, 609
skin manifestations of, 366, 707
smoldering, 225e-5, 707
Strongyloides infections and, 225e-4
treatment of, 225e-5, 707
tumor markers in, 473t
- Advance care directives**, 58, 66, 475
- Advance care planning**, 58–59, 59t, 475
- Advanced cardiac life support (ACLS)**, 1769–1770, 1769f
- Advanced glycosylation end products**, 2424
- Advanced sleep-wake phase disorder**, 193
- Adventitia**, 265e-1, 265e-1f
- Adverse drug reactions**, 43. *See also* Drug-induced illness; *specific drugs*
classification, 43
cytotoxic, 43–44
diagnosis of, 44
in elderly, 77–78
identification of, 32
idiosyncratic hepatotoxicity, 2023
immunologic mechanisms of, 44
incidence of, 43
treatment of, 44
- Adverse events**, 12e-1, 792
- Advisory Committee on Immunization Practices (ACIP)**, 786
- AE1 (red cell anion exchanger)**, 309
- AED (automated external defibrillation)**, 1769
- Aedes spp. mosquitos**
in chikungunya virus infection, 1313
climate change and habitat of, 151e-5, 151e-6f
in dengue fever, 1318–1319, 1322
in Eastern equine encephalitis, 1316
in Jamestown Canyon virus infection, 1315
in La Crosse virus infection, 1315
in lymphatic filariasis, 1418t
in Rift Valley fever, 1322
in yellow fever, 1322
in Zika virus infection, 1314
- AEDs. See** Antiepileptic drugs (AEDs)
- Aerobic capacity**, 73, 74f
- Aerobic glycolysis**, 102e-12, 102e-13f
- Aeromonas spp. infections**
in animal bite-wound, 167e-1
cellulitis, 830
clinical features of, 183e-2, 266t
epidemiology of, 183e-2, 266
in leech infestations, 2747
treatment of, 183e-2, 183e-2t, 830
- Aerophagia**, 285
- Aerophobia**, 896, 1301
- AF. See** Atrial fibrillation (AF)
- Afamelanotide**, 2533
- AF amyloidosis**, 719, 723–724. *See also* Amyloidosis
- Afatinib**
action and targets of, 102e-5t, 103e-16f, 103e-17t, 103e-19
adverse effects of, 103e-17t, 103e-26
for lung cancer, 103e-17t, 103e-19, 521, 521t
- Affected sib and relative pair analysis**, 442t
- Afferent loop syndromes**, 1926
- Affirm test**, 873
- Affordable Care Act**, 15e-2, 59
- Aflatoxin**, 102e-2
- Aflatoxin B₁**, 544, 545t
- Aflibercept**, 205
- AFP. See** a fetoprotein (AFP)
- African eye worm (*Loa loa*) loiasis. *See* *Loa loa* (African eye worm) loiasis**
- Africanized honeybees**, 2749
- African python**, 2747
- African tick-bite fever**, 1155t, 1157
- African trypanosomiasis, human. *See* Sleeping sickness (human African trypanosomiasis)**
- African tumbu fly**, 2747
- Afterdepolarization**, 273e-3–4, 273e-3f
- Afterload**, 265e-8, 265e-9t, 265e-10f
- Agammaglobulinemia**
genetic considerations in, 2111
pathophysiology of, 2110f, 2111
small-intestinal mucosal biopsies in, 1939, 1940t
X-linked, 377e-2
- Ageism**, 16e-4
- Agent 15 (BZ)**, 262e-1t
- AGEP. *See* Acute generalized eruptive/exanthematous pustulosis (AGEP)**
- Aggregatibacter spp.**, 183e-1, 183e-2t. *See also* Infective endocarditis, HACEK group
- Aggrenox**, 748
- Aging**, 70, 94e-1. *See also* Elderly
biologic theories of, 77
cellular processes in, 94e-3
death rates and, 94e-1f
definitions of, 94e-1
disease susceptibility and, 94e-1, 94e-1f
evolutionary theories of, 94e-1–3, 94e-2f
genetic influences on, 94e-4–5
impact on medicine, 94e-1
longevity dividend and, 94e-1
metabolic syndrome and, 2449
mitochondrial DNA mutations and, 85, 85e-8–9, 85e-8f, 94e-3, 94e-5
nutrient-sensing pathways and, 94e-4–5
olfactory function and, 214, 214f, 217
oral health and, 242
osteoarthritis risk and, 2228
skin effects of, 387
sleep patterns and, 185, 185f
strategies to delay
caloric restriction, 85, 94e-5–6
exercise and physical activity, 94e-7
hormesis, 94e-7
periodic fasting, 94e-6
pharmacologic interventions, 94e-6–7, 94e-6f
systemic effects of. *See also* Frailty
body composition, 71, 73t
energetics, 73t, 74f
homeostatic regulation, 73t, 74f, 75
hormone levels, 75t
interaction of, 73f, 76–77
neurodegeneration, 73t, 75–76
tissue changes in, 94e-4
weight regulation and, 275
world demography of, 93e-1
dependency and caregiving ratios, 93e-1t, 93e-4–5, 93e-4f
economic implications, 93e-4
educational attainment trends, 93e-3
historical trends in, 93e-1–2
life expectancy at age 60, 93e-1t
life expectancy trends, 93e-3–4, 93e-3f
numbers and percentages aged 60+, 93e-1t, 93e-2, 93e-2f
numbers and percentages aged 80+, 70f, 71, 71f, 93e-2–3, 93e-3f

- Aging, world demography of (*Cont.*):
 population aging trends, 70–71, 70f, 71f
 population size projections, 93e–2–3
- Agitation, in terminally ill patient, 69t
- Agnetogenic myeloid metaplasia. *See* Primary myelofibrosis
- Agnosia, visual object, 182, 2575
- Agonal gasps, 1775
- Agoraphobia, 151, 2709
- Agouti-related peptide (AgRP), 415e–2, 415e–4, 415e–4f, 415e–4t
- Agrammatism, 178
- Agranulocytosis, 237
- Agraphesthesia, 160
- Agraphia, 177
- agr* gene, 955
- Agurin, 2702
- Agrobacterium* spp., 487
- AH amyloidosis. *See* Amyloidosis
- AH interval, 1474, 1474f
- Ahlstrom's syndrome, 415e–5t
- AHO. *See* Albright's hereditary osteodystrophy (AHO)
- aHUS (atypical hemolytic uremic syndrome), 657
- AIDP (acute inflammatory demyelinating polyneuropathy), 2695, 2695t, 2696t
- AIDS. *See also* HIV infection
 deaths from, 16e–1f, 1224f, 1227f
 definition of, 1215–1216, 1216t, 1217t
 incidence of, 1223–1225, 1227f
- AIDS dementia complex (HIV-associated dementia), 1263, 1264t, 1265f
- AIM2 inflammasome, 145e–8f
- AIN. *See* Acute interstitial nephritis (AIN)
- Ainhum, 2750
- AION (anterior ischemic optic neuropathy), 202, 202f
- AIP. *See* Acute intermittent porphyria (AIP)
- AIP gene mutations, 2340
- Air-conditioner lung, 1681t
- Air-conduction threshold, 222
- AIRE gene mutations, 377e–2, 1343, 2113, 2323, 2345, 2483
- Air flow
 in COPD, 1701–1702, 1704t
 in lung, 306e–1–2, 306e–2f, 306e–3f, 306e–6
- Air hunger, 47e–1, 47e–1t
- Air pollution
 asthma and, 1670, 1671
 COPD and, 1703
 lung disease and, 1693
- Air travel
 fitness for, 794, 799
 in pregnancy, 797–798
 venous thrombosis risk in, 744
- Airway, rewarming of, 478e–3
- Airway clearance, in cardiopulmonary resuscitation, 1768
- Airway hyperresponsiveness
 in asthma, 1672, 1675
 in COPD, 1702–1703
 in sarcoidosis, 2207
- Airway obstruction
 in cancer, 1794, 1794f
 in deep neck infections, 235
 dyspnea in, 47e–1t
- Airway pressure release ventilation, 1742
- Airway remodeling, in asthma, 1675
- Airway resistance (R_{aw})
 increased, dyspnea in, 47e–2
 measurement of, 306e–6
 in obstructive respiratory disorders, 306e–5f
 in restrictive respiratory disorders, 306e–5f
- AIS. *See* Androgen insensitivity syndrome (AIS)
- Akathisia, 2624, 2716, 2722
- AK (adenylate kinase) deficiency, 654t
- AKI. *See* Acute kidney injury (AKI)
- Akinetic mutism, 1771
- AKR1C2 gene mutations, 2353, 2354t
- AKT gene mutations
 cancers associated with, 101e–2t
 in Cowden's syndrome, 2344
 in lung cancer, 507t, 509f
 in melanoma, 496
- AKT pathway, in melanoma, 496, 496f
- ALA (5-aminolevulinic acid), 386
- ALA (5-aminolevulinic acid) dehydrase, 2523t, 2524
- ALA (5-aminolevulinic acid)-dehydratase deficient porphyria, 2521, 2522f, 2522t, 2523t, 2526
- ALA (5-aminolevulinic acid) synthase, 2522, 2523t, 2524–2525
- ALAD gene, 2523t
- AL amyloidosis, 720. *See also* Amyloidosis
 clinical features of, 721, 721f, 2642
 diagnosis of, 365, 720f, 721, 722f, 1845
 differential diagnosis of, 1835t
 etiology and incidence of, 720–722
 pathogenesis of, 1845
 treatment of, 721–723, 1845
- Alanine aminotransferase (ALT)
 in acute viral hepatitis, 2008f, 2010f, 2016
 in alcoholic liver disease, 2053, 2053t
 in autoimmune hepatitis, 2050
 in chronic HBV infection, 2033
 in liver function evaluation, 1997, 1998t
 in SLE, 2131
- ALARA principle, 263e–4
- ALAS genes, 2523t, 2524, 2526
- Albendazole
 action of, 246e–1
 adverse effects of, 246e–1, 246e–3t
 for ascariasis, 1413–1414
 for capillariasis, 1417
 for cutaneous larva migrans, 1411t
 for cysticercosis, 1432
 drug interactions of, 246e–1, 246e–3t
 for echinococcosis, 1433, 1434
 for enterobiasis, 1416
 for *Giardia* infections, 1406
 for gnathostomiasis, 1411t
 for hookworm infections, 1415
 indications for, 246e–3t
 for loiasis, 1422
 for lymphatic filariasis, 1419
 for neurocysticercosis, 903
 pharmacology of, 246e–1
 in pregnancy and lactation, 246e–3t
 for strongyloidiasis, 1416
 for trichinellosis, 1411t
 for trichuriasis, 1416
 for visceral larva migrans, 1411t
- Albers-Schonberg disease, 426e–4
- Albinism
 ocular, 434e–2t
 oculocutaneous, 357, 357t, 434e–2t
 skin cancer in, 500
- Albright's hereditary osteodystrophy (AHO)
 genetic considerations in, 430, 437, 2284t, 2486, 2486t
 hypothyroidism in, 2284t
- Albumin
 serum
 causes of abnormal values, 464t
 in edema, 253
 in hypercalcemia, 314
 in liver function evaluation, 1997–1998
 in nutritional assessment, 462, 464t
 urinary, 293, 1813. *See also* Microalbuminuria
- Albuterol, 312, 473e–8t
- Alcaligenes xylosoxidans*. *See* *Achromobacter xylosoxidans*
- Alcohol, 2723. *See also* Alcohol use; Alcohol use disorder (alcoholism)
 absorption of, 2058, 2723
 acetaminophen toxicity and, 44
 action of, 444e–4, 465e–5t
 behavioral effects of, 2724, 2724t, 2765t
 drug interactions of, 939t
 metabolism of, 2723, 2723f
 pharmacology of, 2723–2724
 tolerance of, 2724
- Alcohol dehydrogenase (ADH), 2058
- Alcohol hand rub, 913
- Alcoholic ketoacidosis, 299, 318–319
- Alcoholic liver disease, 2052
 acetaminophen hepatotoxicity and, 366e–4f
 cirrhosis, 2058–2059, 2058f, 2075
 clinical features of, 2053, 2058–2059
 diagnosis of, 1998t, 2053, 2053t, 2059, 2517t
 etiology of, 2052
 global considerations in, 2052
 pathogenesis of, 2052, 2058, 2725
 pathology of, 2053
 prognosis of, 2053
 risk factors for, 1991–1992, 1992t, 2052, 2052t
 treatment of, 2053, 2059
 alcohol abstinence, 1995
 algorithm for, 2054f
 glucocorticoids, 2053, 2054f
 liver transplantation, 2068, 2075
- Alcohol septal ablation, 1569
- Alcohol use
 breast cancer and, 524, 2725
 cardiovascular effects of, 2725
 esophageal cancer and, 532–533, 2725
 genitourinary effects of, 2725
 head and neck cancer and, 502, 2725
 hematopoietic effects of, 2725
 hepatocellular carcinoma and, 545, 545t
 hypertension and, 1622–1623, 1622t
 hyperuricemia and, 431e–3
 lipoprotein disorders and, 2438
 liver disease and. *See* Alcoholic liver disease
 nervous system effects of, 2724
 porphyria cutanea tarda and, 2530
 in pregnancy, 2725
 prevalence of, 2058
 sleep disturbances and, 190, 2724
 testicular dysfunction and, 2365
- Alcohol use disorder (alcoholism), 2726. *See also* Alcohol withdrawal syndrome
 acetaminophen toxicity in, 44, 2027
 acute intoxication, 164
 anemia in, 398
 cancer risk and, 477t, 2725
 cardiovascular disease in, 1562, 2725
 cognitive dysfunction in, 2607, 2724
 definition of, 2726, 2726t
 dental disease and, 236
 diagnosis of, 2725–2726, 2726t
 epidemiology of, 2726, 2728
 erectile dysfunction in, 326
 folate deficiency in, 647
 gastrointestinal effects of, 277, 2725
 genetic considerations in, 2726
 global considerations in, 2728
 hepatitis in, 283
 hormonal disturbances in, 2725–2726
 hypoglycemia in, 2433
 hyponatremia in, 300
 hypothermia and, 478e–1
 involuntary weight loss in, 275
 lifetime risk for, 2723
 liver disease in. *See* Alcoholic liver disease
 myopathy in, 462e–20, 462e–20t, 2725
 natural history of, 2726
 nausea and vomiting in, 259
 nervous system effects of, 2724
 neuropathy in, 2642, 2724
 ocular involvement in, 211
 olfactory/taste dysfunction in, 215
 palpitations in, 254
 pancreatic insufficiency in, 269
 pancreatitis in, 2091, 2725
 prevalence of, 398, 2058
 psychiatric comorbidity with, 2724–2725
 rehabilitation for, 2728
 screening for, 29t, 1992, 1992t
 testicular defects in, 2365
 thiamine deficiency in, 96e–1, 461
 treatment of, 2726–2728
 in war veterans, 471e–4, 471e–5t
 in women, 6e–5

- Alcohol withdrawal syndrome
clinical features of, 2727
delirium in, 166, 169
hypokalemia in, 64e-7, 305
treatment of, 2727-2728
- Aldehyde dehydrogenase (ALDH), 2058, 2726
- ALDEN (Algorithm of Drug Causality for Epidermal Necrolysis), 384
- Aldesleukin. *See* Interleukin-2 (IL-2) therapy
- Aldolase deficiency, 654t
- Aldosterone
action of, 296, 304-305, 306, 332e-5f, 332e-9
excess of. *See* Glucocorticoid-remediable aldosteronism (GRA);
Hyperaldosteronism
in potassium regulation, 304-305
ratio to plasma renin activity, 1618
reference values for, 2757t
in shock, 1745-1746
in sodium regulation, 332e-11
synthesis of, 2309f, 2313, 2314f
- Aldosterone antagonists, 1623, 1624t
- Aldosterone escape, 332e-11
- Aldosterone-renin-ratio (ARR), 1618, 2319, 2320t
- Aldosterone synthase, 434
- Alemtuzumab
action and targets of, 102e-5t, 103e-21t, 2670
adverse effects of, 488, 1798, 2670
for CLL, 102e-5t, 703
for graft-versus-host disease prevention, 921
for hypereosinophilic syndromes, 135e-8
for immunosuppression, 1828
for MS, 2670
for myelodysplasia, 671
- Alendronate
adverse effects of, 2499
for osteoporosis management/prevention, 2383, 2499, 2500f, 2504
for Paget's disease of bone, 426e-3, 426e-3t
- Alenquer virus, 1309t
- Alexander technique, 14e-1t
- Alexia, 177, 177t, 178-179
- Alfentanil, 246e-3t, 1741
- Alfimeprase, 760
- ALG. *See* Antilymphocyte globulin (ALG)
- Alginic acid, 263
- Algorithm of Drug Causality for Epidermal Necrolysis (ALDEN), 384
- ALI. *See* Acute lung injury (ALI)
- Alien hand (limb) syndrome, 179, 2605
- Alipogene tiparovec, 2440
- Aliskiren, 1512, 1623, 1624t, 1625
- Alkaline phosphatase (ALP)
in hypophosphatasia, 426e-5
in jaundiced patient, 283
in liver abscess, 850
in liver function evaluation, 1996f, 1997, 1998t
in Paget's disease of bone, 426e-1, 426e-3
tissue nonspecific, 426e-5
- Alkali therapy, 320, 321. *See also* Bicarbonate therapy
- Alkaptonuria, 434e-2t, 434e-4-5
- ALK gene mutations, 507t, 508, 509f, 521
- Alkylating agents, 103e-8, 103e-9t. *See also* specific drugs
action of, 103e-8
adverse effects of, 103e-8, 103e-9t
anemia, 398, 664t
in essential thrombocytosis, 678
hair loss, 380
infertility, 623
late, 620t, 621, 670, 678
pulmonary, 621
seizures, 2547t
for breast cancer, 530
carcinogenicity of, 477t, 678
- ALL. *See* Acute lymphoid leukemia (ALL)
- Allele(s), 435
association of, 442
exclusion of, 439
frequency of, 442
heterogeneity of, 433f, 435, 436t
segregation of, 437, 438
types of, 435
- Allele-sharing methods, 442t
- Allelic exclusion, 372e-20, 710
- Allelic loss, 538
- Allen's test, 1445
- Allergic angitis and granulomatosis. *See* Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
- Allergic bronchopulmonary aspergillosis (ABPA)
bronchiectasis in, 1695
clinical features of, 1346t, 1347
in cystic fibrosis, 1699
diagnosis of, 1681, 1686
etiology of, 1686
genetic considerations in, 1346
treatment of, 1349t, 1681, 1686
- Allergic conjunctivitis, 199
- Allergic contact dermatitis. *See* Contact dermatitis, allergic
- Allergic fungal sinusitis, 228
- Allergic granulomatosis, 366
- Allergic interstitial nephritis. *See* Acute interstitial nephritis (AIN), allergic
- Allergic reactions. *See* Hypersensitivity reactions
- Allergic rhinitis, 2121, 2123f
- Allergic vasculitis, 422
- Allergy, 2114
in asthma, 1669-1670, 1671
atopic, 2114
eosinophils in, 422
pathophysiology of, 2114-2116, 2115f
- Alligator bite-wound, 167e-1
- Alloantibodies, 138e-1-2, 138e-1t
- Alloantibody screen, 138e-2
- Allodynia, 87, 158, 2643
- Alloimmunization, 138e-1, 138e-2, 138e-5
- Allopurinol
adverse effects of
cutaneous, 353, 361, 2235
fever, 138, 2705
genetic considerations in, 378
hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
serious, 2235
SJS or TEN, 383. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
vasculitis, 382, 2193, 2218t
drug interactions of, 42, 42t, 2234-2235
azathioprine, 103e-10t, 2705
6-mercaptopurine, 103e-10t
theophylline, 1677
for gout, 2234-2235
for hyperuricemia in myeloproliferative diseases, 674, 676
pyrimidine metabolism and, 431e-6
for uric acid nephrolithiasis, 431e-5, 1871
for uric acid nephropathy, 431e-5
- Alloreactivity, 373e-5
- Allotaxis, 186
- Allotypes, of immunoglobulins, 710
- All-trans-retinoic acid (ATRA). *See also* Retinoic acid
action and targets of, 102e-5t, 102e-7, 103e-16, 103e-17t
for acute promyelocytic leukemia, 102e-5t, 102e-7, 103e-17t, 686
adverse effects of, 103e-16, 103e-17t, 366
- Allylamines, 350
- Almotriptan, 2591, 2592t, 2593t
- Alogliptin, 291e-7, 2414t
- Alopecia, 354
androgenetic, 354, 355t
androgenic, 331
definition of, 340t
drug-induced, 103e-25, 354, 379-380
etiology of, 354-355, 355t
nonscarring, 354, 355t
scarring, 354, 355t
in tinea capitis, 350
traumatic, 354, 355t
treatment of, 355t
- Alopecia areata, 76e-3f, 354, 355t
- Alosetron, 272, 566, 1970, 1970t
- ALP. *See* Alkaline phosphatase (ALP)
- α_1 -adrenergic agonists, 473e-8t
- α_2 -adrenergic agonists, 54, 200, 468e-3, 473e-9t
- α -adrenergic antagonists
adverse effects of, 146, 241, 253t
aldosterone-renin ratio effects of, 2320t
drug interactions of, 587
for hypertension, 1624, 1624t
for lower urinary tract symptoms, 7e-4, 587
overdosage/poisoning with, 473e-3, 473e-9t
- α -adrenergic receptors, 1613, 1745
- α chains, 2506
- Alpha coma, 1775
- Alpha diversity, 86e-1t, 86e-4-5
- α fetoprotein (AFP)
in carcinoma of unknown primary, 120e-1, 120e-2t
ectopic production of, 608
in hepatocellular carcinoma, 546
in testicular cancer, 588
as tumor marker, 473t
- Alpha-granules, 275, 741
- α heavy chain disease (Seligman's disease), 537, 718-719
- α -hemolytic streptococci, 963, 964t, 971. *See also* *Streptococcus pneumoniae*
- α -lipoic acid, 216, 239, 2348
- α motor neurons, 155f
- Alpha (α) radiation, 263e-1
- Alpharadin (radium-223 chloride), 119e-4, 586, 587
- α -synuclein, 453e-6, 2612
- Alphaviruses, 91e-1t, 1306t
- Alphavirus (togavirus) infections
Barmah Forest virus infection, 837, 1313
chikungunya virus infection. *See* Chikungunya virus infection
clinical syndromes of, 1306t, 1313
Eastern equine encephalitis, 1316
geographic distribution of, 1311-1312t
Ross River virus infection (epidemic polyarthritides), 837, 1313
Sindbis virus infection, 1314
Venezuelan equine encephalitis, 1317
Western equine encephalitis, 1317
Zika virus infection, 1314
- Alport's syndrome, 2513
clinical features of, 220, 1836, 1847
diagnosis of, 1847, 2514
forms of, 2513-2514
genetic considerations in, 1832, 1847, 2514
hearing loss in, 220, 222t
incidence of, 2514
renal biopsy in, 62e-8f, 1847
treatment of, 1847
- Alprazolam
adverse effects of, 64
for anxiety disorders, 2711t
for depression, 64
drug interactions of, 246e-3t
overdosage/poisoning with, 473e-10t
for panic disorder, 2709
pharmacology of, 2711t
- Alprostadil, 329
- ALPS (autoimmune lymphoproliferative syndrome), 2112
- ALS. *See* Amyotrophic lateral sclerosis (ALS)
- Alsin gene, 2633, 2634t
- ALS/parkinsonian/dementia complex of Guam, 2608
- ALT. *See* Alanine aminotransferase (ALT)
- Alteplase (rtPA)
action of, 759
for acute limb ischemia, 1646
catheter-directed therapy with, 1636
contraindications to, 1605
domain structure of, 759, 759f
indications for, 759
for ischemic stroke, 2561-2562, 2562t
for myocardial infarction, 1605
for occluded catheter, 98e-7
for pulmonary embolism, 1636

- Alternative splicing, 444e-1
 Alternative systems of medicine, 14e-1. *See also* Complementary and alternative medicine
 Altitude illness. *See* High-altitude illness
 Altretamine, 103e-9t, 103e-12
 Aluminum hydroxide, 1920
 Aluminum toxicity
 erythropoietin therapy response and, 631
 etiology of, 320
 hypercalcemia due to, 2479
 neurologic, 472e-3, 2608
 pulmonary, 1691
 Alveolar collapse, 1738
 Alveolar flooding, 1731
 Alveolar gas equation, 306e-3
 Alveolar hemorrhage, diffuse, 245
 Alveolar hypoventilation, 1732
 Alveolar proteinosis, primary, 375e-2
 Alvimopan, 92
 Alzheimer's disease (AD), **2598**
 aluminum and, 472e-3
 clinical features of, 172, **2598**
 aphasia, 178, 179–180
 Bálint's syndrome, 182
 depression, 2714
 frontal lobe syndromes, 184
 involuntary weight loss, 275
 memory loss, 183
 olfactory dysfunction, 211, 215
 deaths from, 6e-1t, 56t
 dementia with Lewy bodies and, 2606. *See also* Dementia with Lewy bodies
 differential diagnosis of, 171t, 2599
 in Down's syndrome, 2600–2601
 epidemiology of, 2599
 familial, 2600–2601
 functional anatomy of, 170
 genetic considerations in, 444e-9, 2600–2601
 imaging in, 174f, 175f, 2599
 incidence of, 2598
 pathology of, 172t, 2599–2600, 2600f
 pathophysiology of
 A β peptide, 453e-6, 719
 amyloid plaques, 76, 444e-9
 genetic factors, 172t, 448
 molecular, 172t
 prions, 444e-8f, 444e-10, 453e-1, 453e-6
 signaling pathways, 444e-4
 PD and, 2599
 presenilins in, 2601
 treatment of, 176, 2601–2602
 with and without other dementia, 94e-1
 in women, 6e-1–2, 6e-2f
 AMA (antimitochondrial antibodies), 2060
 Amalgam tattoo, 240t
 AMAN (acute motor axonal neuropathy), 2695, 2695t, 2696t
 Amanita mushrooms, 267, 283, 473e-9t
 Amantadine
 action of, 214e-9, 215e-5–6
 adverse effects of, 215e-6, 1213–1214, 2616
 for fatigue in MS, 2672
 for influenza prophylaxis, 215e-2t, 215e-5, 1213t, 1214, 1252t
 for influenza treatment, 215e-2t, 1213–1214, 1213t
 overdosage/poisoning with, 473e-8t, 473e-13t
 for PD, 2616, 2618
 pharmacology of, 215e-5
 resistance to, 215e-2t, 215e-5
 for SSRI-related sexual dysfunction, 2716
 Amaurosis, congenital. *See* Leber's hereditary optic neuropathy (congenital amaurosis)
 Amaurosis fugax, 40e-4, 200–202, 2574
 Amblyomma spp. ticks, 1152, 1154. *See also* Tick-borne diseases
 Amblyopia, 208
 Ambrisentan, 1660, 1660t
 Ameba, free-living. *See* *Acanthamoeba*; *Balamuthia*; *Naegleria*
metabolism of, disorders of, 434e-1, 434e-2–3t
renal transport of, 332e-3f, 332e-8
in specialized nutritional support formulas, 98e-5
types of, 434e-1
 Aminoaciduria(s), 435e-1t, 435e-2
 Aminoacyl-tRNA histidyl synthetase, 372e-24t
 Aminoacyl-tRNA synthetase, 372e-24t
 Aminobenzoic acid esters, 389t
 ϵ -Aminocaproic acid
 adverse effects of, 309, 2199
 for CIINH deficiency, 2120
 for DIC, 738
 for hemophilia, 734
 for thrombocytosis, 678
 for von Willebrand disease, 731–732
 Aminoglutethimide, 103e-16, 2273
 Aminoglycosides
 action of, 930, 931t, 932f, 942
 adverse effects of, 938t
 hearing loss, 219
 hypokalemia, 306
 in myasthenia gravis, 2703
 nephrotoxicity, 1803
 for anaerobic bacterial infections, 1101t
 for cystic fibrosis, 1699
 for enterococcal infections, 974, 975t
 indications for, 942
 for infections in cancer patient, 491
 for infective endocarditis, 821
 for osteomyelitis, 840t
 for postpartum endomyometritis, 50
 resistance to, 821, 931t, 934, 976, 1118
 for tuberculosis, 205e-2t
 5-Aminolevulinic acid (ALA), 386
 5-Aminolevulinic acid (ALA) dehydrase, 2523t, 2524
 5-Aminolevulinic acid (ALA)-dehydratase deficient porphyria, 2521, 2522f, 2522t, 2523t, **2526**
 5-Aminolevulinic acid (ALA) synthase, 2522, 2523t, 2524–2525
 Aminophylline, 254, 324, 1677, 1679, 2117
 4-Aminopyridine, 2672
 Aminosalicilic acid, 727t
 Aminosidine. *See* Paromomycin
 Aminotransferase(s), 283, 1997, 1998t
 Amiodarone
 action of, 273e-6t
 adverse effects of
 in chronic lung disease, 36
 cutaneous, 361, 379
 hepatotoxicity, 2025, 2029
 hyperthyroxinemia, 2286t
 interstitial lung disease, 1714
 myopathy, 462e-20, 462e-20t, 2199, 2218t
 neuropathy, 2687t, 2688
 nonarrhythmic, 1480t, 1499
 optic neuropathy, 203
 phototoxicity, 389t
 proarrhythmic, 1480t, 1499
 secondary parkinsonism, 2611
 thrombocytopenia, 727t
 thyrotoxicosis, 2300
 for arrhythmias, 1479t, 1480t
 for atrial fibrillation, 1489
 for cardiac arrest, 1769f, 1770
 dosage of, 1479t, 1480t
 drug interactions of
 calcium channel blockers, 34t
 digoxin, 42t
 protease inhibitors, 34t
 quinidine, 42t, 246e-6t
 statins, 2199
 warfarin, 34t, 42t
 ECG changes caused by, 1458f
 for focal atrial tachycardia, 1479
 monitoring during treatment with, 1499
 for multifocal atrial tachycardia, 1485
 pharmacology of, 1480t
 thyroid function and, 2300
 for ventricular arrhythmias, 1494, 1495, 1498–1499

- for ventricular fibrillation, 1498
for ventricular tachycardia, 1608
- Amitriptyline**
adverse effects of, 2710t, 2716
 cardiac, 1467t
 drowsiness, 2594t
 hypotension, 2640t
 smell and taste distortion, 216
for chronic daily headache, 109
for ciguatera poisoning, 2742
for cough, 245
for depression, 473, 2672, 2710t, 2716
for fibromyalgia, 2240, 2240t
for insomnia, 191
for interstitial cystitis/bladder pain syndrome, 60e-3
for migraine prevention, 2593, 2594t
monitoring during treatment with, 2764t
for nausea and vomiting, 260t
for new daily persistent headache, 111
for olfactory dysfunction, 216
overdosage/poisoning with, 473e-9t, 473e-10t, 2764t
for pain, 91t, 2672
for tension-type headache, 2594
- AML.** See Acute myeloid leukemia (AML)
- Amlodipine**, 1513, 1589, 1589t, 1649, 2640t
- Ammonia**, as urinary buffer, 332e-7
- Ammonia/ammonium**, 320, 1996
- Ammonia exposure**, 1692t
- Ammoniogenesis**, 320, 332e-7, 333e-3
- Amnesia**, **182**
after head injury, 457e-1
anterograde, 182
caring for patient with, 184
clinical examination in, 37e-1v, 182–183
etiology of, 183
psychogenic, 2608
retrograde, 182
transient global, 183, 2608
- Amnestic shellfish poisoning (domoic acid intoxication)**, 2547–2548, 2743
- Amnestic state**, 182
- AMN gene mutations**, 645
- Amniocentesis**, 83e-3, 444
- Amnionless**, 640
- Amniotic fluid-derived stem cells**, 92e-1, 453t
- Amniotic fluid embolism**, 40e-4, 49, 1632
- Amniotic infection**, 50
- Amodiaquine**
action of, 246e-1, 246e-7
adverse effects of, 246e-1t, 246e-7, 1380t, 1383
for malaria, 1378t, 1380t
pharmacology of, 246e-1, 246e-7, 1380t
- Amoxapine**, 2710t, 2716
- Amoxicillin**
for actinomycosis, 1090–1091, 1091t
adverse effects of, 377, 1923
for anthrax, 261e-5t
for *C. trachomatis* infection, 1171
for community-acquired pneumonia, 808t
for endocarditis prophylaxis, 826t
for enteric fever, 1052t
for gonococcal arthritis, 835
in *H. pylori* eradication, 1042t, 1922t, 1923
indications for, 937–938, 940t
for leptospirosis, 1145t
for Lyme disease, 836, 1153, 1153f
for nontyphoidal *Salmonella* infections, 1054t
for osteomyelitis, 841t
for otitis media, 229t, 230, 953
for pharyngitis, 232f
for pneumonia, 952
in pregnancy and lactation, 937t
prophylactic, 945t
resistance to, 940t
for rheumatic fever, 2152
for sinusitis, 227, 227t
- Amoxicillin-clavulanate**
adverse effects of, 2023, 2025t, 2027–2028
for bite-wound infections, 167e-2, 167e-3t, 832t
for community-acquired pneumonia, 808t
for febrile neutropenia, 103e-23
for lung abscess, 815
for multidrug-resistant TB, 205e-7
for *Nocardia* infections, 1087
for osteomyelitis, 841t
for otitis media, 229t
for pelvic inflammatory disease, 878
prophylactic, 945t
for sinusitis, 227, 227t
- AmpC**, 933
- AmpC β -lactamase**, 1026–1027, 1033–1034
- Amphetamine(s)**
action of, 444e-4, 465e-5t
adverse effects of
 alopecia, 354
 cardiomyopathy, 1562
 myopathy, 462e-20, 462e-20t
 palpitations, 254
 vasculitis, 2218t
for fatigue, 153
overdosage/poisoning with, 473e-8t, 473e-13t, 2764t
therapeutic monitoring of, 2764t
- Amphiphsin**, 372e-25t
- Amphotericin B**
adverse effects of, 1330
 frequent, 246e-2t, 1390
 hypokalemia, 306
 myopathy, 2199
 nephrotoxicity, 1803
 occasional, 246e-2t
 rare, 1390
 thrombocytopenia, 727t
for *Aspergillus* infections, 1348, 1349t
for *Blastomyces* infections, 1338, 1338t
for *Candida* infections
 disseminated, 1344t, 1830
 endocarditis, 824
 liver abscess, 851
 prophylactic, 1345
 thrush, 233
for *Coccidioides* infections, 899, 1336, 1336t
for *Cryptococcus* infections, 899, 1264, 1341
drug interactions of, 246e-2t
for febrile neutropenia, 491
for *Fusarium* infections, 1356
for *Histoplasma* infections, 899, 1334, 1334t
for *Leishmania* infections, 1390–1391
lipid formulations of, 1330
for mucormycosis, 1352t, 1353
for mucosal leishmaniasis, 1393
for paracoccidioidomycosis, 1354, 1354t
for penicilliosis, 1354t
in pregnancy and lactation, 246e-2t
in sepsis/septic shock, 1757t
for sinusitis, 227
for *Sporothrix* infections, 1354t
for visceral leishmaniasis, 1390–1391
- Ampicillin**
adverse effects of, 25e-2f, 284, 362, 727t
for clostridial infections, 991t
drug interactions of, 246e-1t, 1921
for enterococcal infections, 975t, 976
for HACEK infections, 183e-2t
indications for, 937–938, 940t
for infective endocarditis, 822t, 823t, 824, 826t, 975t
for *L. monocytogenes* infections, 983
for leptospirosis, 1145t
for meningitis, 887t, 889, 889t, 975t, 1011
for necrotizing fasciitis, 832t
for nontyphoidal *Salmonella* infections, 1054t
in pregnancy and lactation, 937t
resistance to, 940t, 976
 in gram-negative bacteria, 1033, 1034, 1035
 in *H. influenzae*, 235
for streptococcal infections in pregnancy, 50
for urinary tract infections, 975t
- Ampicillin-clavulanate**, 227t, 938–939
- Ampicillin-sulbactam**
for anaerobic bacterial infections, 1101t
for bite-wound infections, 167e-2, 167e-3t, 832t
for *Capnocytophaga* infections, 183e-2t, 183e-3
for cellulitis, 832t
for community-acquired pneumonia, 767t
for deep neck infections, 235
for epiglottitis, 235
for HACEK infections, 183e-2t, 822t
for health care-associated pneumonia, 812t
indications for, 938–939
for infective endocarditis, 822t
for Lemierre's disease, 234
for osteomyelitis, 841t
resistance to, 1101, 1102
for skin and soft tissue infections, 832
- AMPK (5' adenosine monophosphate-activated protein kinase)**, 94e-5, 102e-13
- Amplatzer septal occluder**, 297e-7f, 297e-8f
- Emprenavir**, 1276f, 1278
- Ampulla of Vater, carcinoma of**, 553
- Ampullary carcinoma**, 285
- Amputation**
for frostbite, 478e-4–5
risks for, in diabetes mellitus, 2428
- AMS.** See Acute mountain sickness (AMS)
- AMSAN (acute motor sensory axonal neuropathy)**, 2695, 2695t, 2696t
- Amsel criteria**, for bacterial vaginosis, 874
- Amur/Soochong virus**, 1307t
- Amygdalohippocampectomy**, for epilepsy, 2557
- Amylase**
ascitic fluid, 288, 2087t, 2088
in pancreatic disease evaluation, 2087–2088, 2087t, 2089t
pleural fluid, 2088
- Amyl/butyl nitrate**, 328
- Amylin agonists**, 2414t
- Amyl nitrite**, for cyanide poisoning, 262e-7–8
- Amyloid angiopathy, cerebral**, 2583–2584, 2583t, 2600
- Amyloid beta (A β) peptide**, 453e-6
- Amyloidosis**, **719**
AA. See AA amyloidosis
A β , 719
A β M, 719, 720t, 724
AF, 719, 723–724
AH, 720t
AL. See AL amyloidosis
amyloid fibril proteins in, 719, 720t
ATTR, 720t, 723–724
cardiac, 290e-1t, **1566**
 AV conduction block in, 1471, 1471t
 clinical features of, 1566
 diagnosis of
 biopsy, 1567, 1567f
 CMR, 270e-21, 270e-21f, 271e-6f, 1566–1567
 echocardiography, 271e-6f, 271e-6v, 1566–1567, 1567f
 epidemiology of, 1566
 SA node dysfunction in, 1467
 transthyretin-mediated, 271e-6f, 1566
 treatment of, 1567
classification of, 719, 720t
clinical features of, **719**
 autonomic neuropathy, 2642
 bleeding, 403
 cutaneous, 364t, 365, 367
 diarrhea, 269
 gastroparesis, 258
 hepatic, 367e-3
 hyperkalemia, 307
 neuropathy, 2681–2682
 ocular, 196
 proteinuria, 293
 renal, 62e-6f, 1835t, 1845
 splenomegaly, 411
diagnosis of, 719, 720f, 2642
epidemiology of, 367e-3
familial, 720t, 723–724
in familial Mediterranean fever, 723, 1845, 2214
in IBD, 1959
in lepromatous leprosy, 1126

- Amyloidosis (Cont.):**
 lichenoid, 365
 localized, 720t
 macular, 365
 in monoclonal gammopathy of uncertain significance, 721
 in multiple myeloma, 721, 1567
 nodular, 365
 pathogenesis of, 721, 2642
 prognosis of, 1567
 senile, 1467, 1566
 small-intestinal mucosal biopsies in, 1939, 1940t
 systemic, 720t
 treatment of, 721–723, 724, 1845
- Amyloid plaques**
 in AD, 76, 172t, 444e–8f, 444e–10
 in Creutzfeldt-Jakob disease, 453e–4
- Amyloid polyneuropathy, familial,** 2681
- Amyloid precursor protein (APP)**
 in AD, 444e–10, 453e–6, 2600, 2600f
 in amyloidosis, 719
 in neurodegenerative disease, 444e–9
- Amyolytic enzymes,** 2090
- Amyotrophic lateral sclerosis (ALS), 2631**
 clinical features of, 2632–2633
 diagnosis of, 2632–2633
 differential diagnosis of, 2633
 dysphagia in, 256, 257
 epidemiology of, 2633
 familial, 437, 444e–9, 2633, 2634t
 frontotemporal dementia and, 2633, 2635t
 genetic considerations in, 444e–1, 444e–9, 2634t
 imaging in, 2632f
 juvenile-onset, 2636
 parkinsonism and, 2633, 2635t
 pathogenesis of, 2633, 2635
 pathology of, 2631–2632, 2632f
 treatment of, 90e–3, 2633
- Amyotrophic lateral sclerosis (ALS)/parkinsonian/dementia complex of Guam,** 2608
- Amyotrophy,** 2632
- ANA. See Antinuclear antibodies (ANA)**
- Anabolic-androgenic steroid (AAS) abuse**
 adverse effects of, 7e–2–4, 7e–3t, 2374
 cardiovascular, 2374
 cholestasis, 284
 edema, 253t
 hepatotoxicity, 2029
 hirsutism, 380
 long-term, 2374
 diagnosis of, 7e–3–4, 7e–3t, 2374
 epidemiology of, 7e–2, 2374
 treatment of, 7e–4
 withdrawal symptoms, 2374
- Anaerobic bacteria, 1094**
 antibiotic resistance in, 1101–1102, 1101t
 antibiotic susceptibility testing of, 1101
 in normal human microbiota, 1094–1095, 1095t
 specimen handling, 1100
 virulence factors of, 1096
- Anaerobic bacterial infections, mixed, 1094**
 approach to the patient, 1096–1097
 clinical features of
 bacteremia, 1100
 bone and joint, 1099–1100
 in burn wounds, 166e–1
 with clostridial involvement, 990–991, 991t
 CNS, 1098
 in dog-bite wounds, 167e–1
 endocarditis and pericarditis, 1100
 in human bite-wounds, 167e–2
 intraabdominal, 1099, 1102
 mouth, head, and neck, 1097, 1102
 pelvic, 1099
 pleuropulmonary, 1098–1099, 1098f
 skin and soft tissue, 1099
 definitions in, 1094
 diagnosis of, 150e–5t, 1100
 epidemiology of, 1097
 etiology of, 1095–1096
 pathogenesis of, 1096
 supportive measures, 1102
 treatment of, 1101–1102, 1101t
- Anaerobic glycolysis,** 461
- Anagen, 331**
- Anagen effluvium,** 354, 379
- Anagliptin,** 2414t
- Anagrelide,** 674, 678
- Anajatuba virus,** 1307t
- Anakinra**
 action of, 372e–30
 adverse effects of, 2146t
 for calcium apatite deposition disease, 2237
 for calcium pyrophosphate deposition disease, 2236
 for familial Mediterranean fever, 2214
 for fever of unknown origin, 141
 for gout, 2234
 indications for, 372e–8t, 2147
 for rheumatoid arthritis, 2146t, 2147
- Anal cancer,** 351, 543–544, 1198, 1199
- Anal fissures, 1978**
- Anal fistula,** 1111, 1978
- Analgesia (sensory abnormality),** 158
- Analgesics**
 adverse effects of, 1861, 1861f
 for lumbar puncture, 443e–1–2
 medication-overuse headache and, 109, 2593
 urinary, 867
- Anal intercourse, HIV infection risk and,** 1221
- Anal reflex,** 2539
- Anal sphincter,** 265f
- Anaphylactic reactions. See also Hypersensitivity reactions**
 to dialyzer, 1824
 drug-induced, 381
 to insect stings, 2749
 to transfusion, 138e–5
- Anaphylaxis, 2116**
 clinical features of, 2116
 definition of, 2116
 diagnosis of, 2116–2117
 drug-induced, 381
 etiology of, 2116
 exercise-induced, 2118, 2119
 pathophysiology of, 2116
 predisposing factors for, 2116
 prevention of, 2117
 respiratory acidosis in, 322–323
 treatment of, 2117, 2749
- Anaplasma spp. infections,** 888, 1152, 1160.
See also Human granulocytic anaplasmosis
- Anaplastic large T/null cell lymphoma,** 102e–5t, 699t, 702t, 707
- Anaplastic thyroid cancer,** 2307
- Anasarca,** 251
- Anastrozole,** 103e–15
- Anatomic dead space,** 306e–3
- ANCA. See Antineutrophil cytoplasmic antibodies (ANCA)**
- Anchoring heuristic,** 19
- Ancylostoma braziliense,** 245e–3t, 1415. *See also* Larva migrans, cutaneous
- Ancylostoma caninum,** 1415. *See also* Larva migrans, cutaneous
- Ancylostoma duodenale,** 245e–3t, 1414, 1414t.
See also Hookworm infections
- Andersen syndrome (type IV GSD),** 433e–2t, 462e–14
- Andersen-Tawil syndrome,** 462e–17t, 462e–18
- Andersson lesions,** 840
- Andes virus,** 1307t
- Androgen(s)**
 abuse of. *See* Anabolic-androgenic steroid (AAS) abuse
 action and metabolism of, 2359, 2359f.
See also Androgen insensitivity syndrome (AIS)
 adrenal, for adrenal insufficiency, 2327
 adverse effects of, 378
 body fat distribution and, 6e–4
 excess of, 332–333
 in hair growth and differentiation, 331–332
 synthesis of
 disorders of, 2353, 2354t
 pathways of, 2309f, 2355f, 2358–2359, 2358f, 2359f
- Androgen blockade. See Androgen deprivation therapy (ADT)**
- Androgen deficiency**
 age-related, 7e–1
 approach to the patient, 2368f, 2369
 diagnosis of, 2369
 gynecomastia and, 2366
 in rheumatoid arthritis, 2139
 treatment of, 2369. *See also* Testosterone therapy
- Androgen-depletion syndrome,** 585
- Androgen deprivation therapy (ADT)**
 adverse effects of, 7e–5–6, 7e–5f, 585
 approach to the patient, 7e–6, 7e–6t
 for hirsutism, 334
 intermittent use of, 586
 for prostate cancer, 103e–15, 585–586
- Androgen insensitivity syndrome (AIS), 2353**
 amenorrhea in, 335
 clinical features of, 2354–2355, 2354t
 complete, 2353–2354
 genetic considerations in, 335, 431t, 2354, 2366
 gynecomastia due to, 2366
 partial, 2355
 testicular dysfunction due to, 2366
 treatment of, 336
- Androgen receptors,** 102e–7, 431t, 2354t, 2359
- Androgen therapy. See also Testosterone therapy**
 adverse effects of, 380
 carcinogenicity, 477t
 cholestasis, 2025t
 cutaneous, 352
 monitoring of, 2372–2374, 2373t
 for aplastic anemia, 667
 for breast cancer, 530t
 contraindications to, 2372, 2372t
 for female sexual dysfunction, 331
 indications for, 2371–2372
 novel formulations of, 2371
 recommended regimens, 2372
- Andropause,** 2367
- Androstenedione,** 332, 2757t
- Anemia, 393. See also specific types**
 in acute kidney injury, 1807, 1810
 after kidney transplantation, 1831
 after liver transplantation, 2072
 approach to the patient, 393
 from blood loss, 399, 662
 chemotherapy-related, 103e–24–25
 in chronic kidney disease. *See* Chronic kidney disease (CKD), hematologic abnormalities in
 classification of, 398, 398f
 clinical features of, 393
 in colorectal cancer, 541
 in critically ill patient, 1735
 diagnosis of, 393
 bone marrow examination, 397
 initial workup, 393–394, 394t
 iron supply and storage tests, 397
 peripheral blood smear, 394, 394–396f.
See also Peripheral blood smear reticulocyte count in, 394, 396–397, 396t, 397t
 dyspnea in, 47e–1t, 47e–2, 47e–2t
 in endocrine deficiency states, 630
 endoscopic findings in, 1896, 1897f
 erythropoietin in, 392f
 in gastrointestinal bleeding, 276
 in heart failure, 1514
 hemolytic. *See* Hemolytic anemia
 history in, 393
 in HIV infection, 1262
 in hypometabolic states, 630–631, 630t
 hypoproliferative, 398, 625, 630, 631.
See also Aplastic anemia; Iron-deficiency anemia

- in hypothermia, 478e-2
hypoxia in, 248
in inflammatory states, 628, 628t, 629–630, 630f
lactic acidosis in, 318
in liver disease, 631
macrocytic, 81e-8f
in malabsorption syndromes, 1945
from maturation disorders, 398–399
in multiple myeloma, 714
myelophthisic, 671–672
normocytic, in rheumatoid arthritis, 2138
physical examination in, 393
in protein starvation, 630–631
refractory, 669t. *See also* Myelodysplasia
sideroblastic, 399, 451e-4t, 628t
in SLE, 2131
in thalassemia, 637
treatment of, 399
- Anesthesia (sensory abnormality), 158
Anesthetics, 54, 64, 64t
Aneuploidy, 83e-1, 83e-4
Aneurysm
 aortic. *See* Aortic aneurysm
 aortic sinus of Valsalva, 1523
 coronary artery, 271e-4f, 2192–2193
 fusiform, 1637
 intracranial, 1784–1785. *See also*
 Subarachnoid hemorrhage
 left ventricular, 1610
 mycotic. *See* Mycotic aneurysm
 popliteal artery, 1647
 renal artery, 1630f
 saccular. *See* Saccular aneurysm
 true, 1637
- Ang1/Ang2. *See* Angiopoietins
Angelman's syndrome, 83e-7, 439, 444e-2
Angiitis
 allergic. *See* Eosinophilic granulomatosis with
 polyangiitis (Churg-Strauss syndrome)
 cutaneous leukocytoclastic, 2119, 2191
 isolated granulomatous, chronic meningitis
 in, 909t
- Angina
 microvascular, 1579
 Prinzmetal's variant, 1456, 1579, 1588, **1598**
 stable, **1580**. *See also* Ischemic heart disease
 in aortic stenosis, 1530
 atherosclerosis and, 291e-4
 atypical, 1581
 exertional, 1581
 history in, 1580
 myocardial infarction probability with, 99f
 in myocardial ischemia, 95–97, 96t, 99–100,
 99f, 1439, 1581, 1594
 in orthostatic hypotension, 146
 physical examination in, 1581–1582
 severity classification of, 1581, 1581t
 threshold for development of, 1581
 treatment of, 296e-3, 1586–1587. *See also*
 Ischemic heart disease, treatment of
 unstable, 96, 96t, 97, 1593. *See also*
 Non-ST-segment elevation acute
 coronary syndrome (NSTEMI-ACS)
- Angina decubitus, 1581
Anginal equivalents, 1581
Angina pectoris, 95. *See also* Angina, stable;
 Angina, unstable
Angiocentric immunoproliferative lesion/
 angiocentric lymphoma. *See*
 Extranodal NK/T-cell lymphoma,
 nasal type
Angiodermatitis, acral, 367
Angioedema, **2117**
 classification of, 2117, 2118t
 clinical features of, 363, 2119
 definition of, 2117
 diagnosis of, 2119
 drug-induced, 381, 384t, 1623
 etiology of, 363t, 2117–2118, 2118t
 hereditary, 372e-25t, 2118, 2119, 2120
 pathophysiology of, 2119
 predisposing factors for, 2117–2118, 2118t
 treatment of, 381, 2119–2120
 vibratory, 2118
Angiofibroma, 364, 364t
Angiofibromas, 101e-6t, 2336t, 2340
Angiogenesis, 265e-4
 in gastric mucosa, 1912
 inhibition of, **102e-15**
 gene therapy for, 91e-4
 mechanisms of, 102e-16f
 therapeutic strategies, 102e-17f
 tumor, 102e-1t, 102e-13
 inhibition of, 102e-15, 102e-16f, 102e-17
 signaling pathways in, 102e-14–15,
 102e-15f, 102e-16f
 vessel formation mechanisms in,
 102e-13–15
Angiogenic switch, 102e-13
Angiography
 cerebral
 in CNS vasculitis, 441e-19f
 in CNS vasculopathy, 2568f
 complications of, 440e-10
 indications for, 440e-10
 in stroke, 2581, 2581f
 coronary. *See* Coronary angiography
 CT. *See* Computed tomography (CT)
 angiography (CTA)
 magnetic resonance. *See* Magnetic resonance
 angiography (MRA)
 pulmonary, 1634, 1666–1667
 renal intraarterial, 1628t
 spinal, 440e-10–11
Angioimmunoblastic T-cell lymphoma, 135e-4
 to 135e-5, 708
Angiokeratoma, 364t, 365, 432e-1, 436e-7f
Angioma
 cavernous, 2586
 cherry, 341t
 senile, 342f
 small intestinal, 536
 spider, 286, 1992, 2058, 2059f
Angiomyolipoma, renal, 101e-6t, 1854
Angiopoietins, 89e-3, 102e-14–15, 102e-15f,
 102e-17f
Angiosarcoma, 119e-2, 364t, 367
Angiostrongylus spp. infections
 abdominal, 1417
 clinical features of, 908t, 1412
 diagnosis of, 245e-3t, 245e-6t, 765t, 1412
 epidemiology of, 245e-3t
 life cycle of parasite, 245e-3t, 1412, 1412f
 pathogenesis of, 1412
 treatment of, 1411t, 1412
Angiotensin-converting enzyme (ACE)
 in glomerular filtration rate regulation,
 332e-2, 332e-2f
 reference values for, 2757t
 in sarcoidosis, 2209
 in sodium absorption, 332e-11
Angiotensin-converting enzyme (ACE)
 inhibitors
 action of, 1623
 adverse effects of
 acid-base disorders, 321t
 anaphylaxis, 44
 angioedema, 1623, 2118, 2119
 cough, 244, 1623
 cross-sensitivity and, 385
 cutaneous, 353, 378, 379, 381
 erectile dysfunction, 326
 hyperkalemia, 64e-2, 310
 lupus syndrome, 2218t
 renal, 291, 1623, 1800, 1801f, 1815
 transfusion-related hypotension, 138e-5–6
 for aortic aneurysm in Marfan's syndrome,
 1639
 for aortic dissection, 1641
 for diabetic nephropathy, 2426
 for heart failure, 1511, 1511f, 1512t
 for hypertension, 1623, 1624t
 in chronic kidney disease, 1817, 1821
 comparison to other agents, 1625
 in metabolic syndrome, 291e-7, 2454
 for hypokalemia, 308
 for ischemic heart disease, 1590
 metabolism of, 35
 for myocardial infarction, 1596, 1598, 1607
 for peripheral artery disease, 1645
 preoperative discontinuation of, 54
 for pulmonary edema, 1761f, 1763
 for radiation-induced cardiovascular disease,
 621
 for renal artery stenosis, 1618
 for scleroderma renal crisis, 1865, 2164
Angiotensin II
 aldosterone levels and, 306
 in blood pressure regulation, 1613–1614
 in glomerular filtration rate regulation,
 332e-2, 332e-2f, 1800
 in shock, 1745
 in sodium reabsorption, 296, 332e-11
 in urinary tract obstruction, 1872
Angiotensinogen, 1613
Angiotensin receptor blockers (ARBs)
 adverse effects of, 1623
 cutaneous, 381
 hyperkalemia, 64e-2, 310
 renal, 1800, 1801f, 1815
 aldosterone-renin ratio effects of, 2320t
 for diabetic nephropathy, 2426
 for heart failure, 1507, 1511–1512, 1512t
 for hypertension, 1623, 1624t
 in chronic kidney disease, 1817, 1821
 comparison to other agents, 1625
 in metabolic syndrome, 291e-7, 2454
 for Marfan syndrome, 445
 metabolism of, 40
 for myocardial infarction, 1607
 preoperative discontinuation of, 54
 for Raynaud's phenomenon, 2164
 for renal artery stenosis, 1618
Angle of Louis, 1443
Anidulafungin, 233, 1331, 1344t, 1352t
Aniline dyes, 575
Animal bite-wounds. *See* Bite-wounds
"Animal Rule," 261e-10
Anion gap, 317, 2757t
Anisakis/anisakiasis, 765t, 1416–1417
Anisimus, 274
Anisocoria, 196–197
Anisocytosis, 81e-1, 394, 413
Anistreplase, 758
ANK gene mutations, 653t
ANKH gene mutations, 2235
Ankle, pulse measurement in, 1446f
Ankle-brachial index (ABI), 1616, 1643
Ankylosing spondylitis, **2169**
 in adolescence, 2170
 classification of, 2169, 2169t
 clinical features of, **2170**
 aortitis, 290e-3, 1642
 murmur, 51e-6, 1443
 neck pain, 122
 ocular, 2171
 pseudoarthrosis, 2171
 sacroiliitis, 2171, 2172f
 skeletal, 1443
 spinal inflammation, 2170–2171, 2172f
 complications of, 2171
 diagnosis of, 2169t, 2171–2172, 2172f
 epidemiology of, 2169
 HLA gene association with, 373e-7, 373e-8t
 IBD and, 1958–1959, 2178–2179
 KIRs with, 372e-15t
 pathogenesis of, 2170
 pathology of, 2169–2170
 prognosis of, 2171
 treatment of, 2172–2173
 vs. vertebral osteomyelitis, 840
Annelid worm stings, 2739, 2739f
Annexin-2, 402
Annular pancreas, 2102
Annular skin lesion, 340t
Annuloaortic ectasia, 51e-6
Anogenital warts. *See* Condyloma acuminata
 (anogenital warts)

- Anomia, 177, 179, 197
Anopheles mosquitoes
 climate change and, 151e-4
 in lymphatic filariasis, 1418t
 in malaria transmission, 1368–1370, 1369f, 1371
- Anorectal (rectoanal) angle, 265, 265f
 Anorectal disorders, **1973**
 anal fissures, 1978
 anorectal abscess, 1977–1978, 1977f
C. trachomatis proctitis, 1168
 in Crohn's disease, 1953
 fecal incontinence. *See* Fecal incontinence
 fistula in ano, 1111, 1978
 hemorrhoidal disease. *See* Hemorrhoidal
 disease
 in HIV infection, 1257–1258, 1258f
 perianal skin tags, in IBD, 1958
 rectal prolapse. *See* Rectal prolapse
 solitary rectal ulcer syndrome, 277, 1957
 Anorectal manometry, 274
 Anorectum, 265, 265f
 Anorexia
 in appendicitis, 1987
 in elderly, 81
 enteral tube feeding in, 98e-3
 in gastric cancer, 534
 in terminally ill patient, 69t
 Anorexia nervosa, 259, **2719**
 Anorexiants, 1617t, 2396–2397, 2453
 Anosmia. *See* Olfactory disorders
 Anosognosia, 180, 2560, 2598
 Anovulation, 335
 Anoxic cerebral injury, in critically ill patient,
 1735
- ANP (atrial natriuretic peptide), 251, 310,
 332e-6f, 332e-9, 511
- Anserine bursitis, 2222, 2248
- Ant(s), stinging, 2749
- Antacids
 adverse effects of, 267, 269, 321
 drug interactions of, 41, 42t, 246e-1t, 246e-3t,
 939t
 for GERD, 263
 for internal radionuclide contamination,
 263e-6, 263e-6t
 for peptic ulcer disease, 1919–1920, 1920t
- Antagonistic pleiotropy, 94e-2
- Anterior cerebral artery
 lesion in territory of, 156, 178
 occlusion of, 2573, 2575f
 territory of, 2572f, 2574f
- Anterior chamber, eye, 40e-1
- Anterior choroidal artery, 2573–2574, 2575f
- Anterior dens dislocation, 441e-31f
- Anterior horn cell disorders, 154, 157
- Anterior ischemic optic neuropathy (AION),
 202, 202f
- Anterior pituitary. *See also* Pituitary gland
 disorders of
 insufficiency. *See* Hypopituitarism
 tumors. *See* Pituitary tumors (adenomas)
 hormone expression and regulation in,
 401e-1–4, 401e-1t. *See also specific
 hormones*
- Anterior spinal artery syndrome, 2652, 2655
- Anterocollis, 2620
- Anterograde amnesia, 182
- Anterolateral system, 158
- Anthracyclines
 adverse effects of, 103e-25, 380–381, 620t,
 621, 1562
 for breast cancer, 530
 for soft tissue sarcoma, 119e-2
 for thymoma, 123e-2
- Anthralin, 355t
- Anthrax
 in bioterrorism, 261e-2–3, 261e-2f
 chest x-ray in, 261e-3, 261e-4f
 clinical features of, 261e-2–3, 261e-3f, 261e-
 5t, 784
 cutaneous, 134t, 135, 261e-2, 261e-3f, 829
 diagnosis of, 261e-5t
 gastrointestinal, 261e-2
 inhalational, 261e-2–3, 784
 prevention of, 261e-3, 261e-5t
 treatment of, 261e-3, 261e-5t
- Anthrax vaccine, 261e-3
 Anthrax vaccine adsorbed, 261e-3
 Anthropometrics, 462
 Anthroposophic medicine, 14e-1t
- Anti-AChR antibodies, 372e-24t, 377e-4, 614t,
 2701, 2702t
- Anti-AMPA receptor antibodies, 617
- Anti-amphiphysin, 614t
- Antiandrogens. *See* Androgen deprivation
 therapy (ADT)
- Antiarrhythmics, **273e-5**. *See also specific drugs*
 actions of, 273e-5, 273e-6t
 adverse effects of, 267, 1467t, 1471t, 1480t
 classification of, 273e-5, 273e-6t
 intravenous dose range/primary indication,
 1479t
 oral dosing/primary indications, 1480t
 overdosage/poisoning with, 473e-13t
 for pain, 91t, 94
- Antibacterial drugs, **930**. *See also specific drugs
 and diseases*
 absorption of, 934–935
 actions of, 930–932, 931t, 932f
 adverse effects of, 936
C. difficile infections, 858
 in G6PD deficiency, 656t
 allergy to, 936
 for anaerobic gram-negative rods, 1101t
 approach to therapy with, 935–936
 concentration-dependent, 935
 distribution of, 935
 drug–drug interactions of, 936
 effects on microbiota, 86e-7
 empirical therapy with
 for acutely ill febrile patient, 779, 780t
 for community-acquired pneumonia, 807,
 808t
 for febrile neutropenia, 103e-23
 for fever of unknown origin, 141
 for infective endocarditis, 824
 for sepsis/septic shock, 1756, 1757t
 excretion of, 935
 inappropriate use of, 918, 1026
 for infections in cancer patient, 490–491
 metabolism of, 935
 pharmacodynamics of, 935, 935f
 pharmacokinetics of, 934–935, 935f
 prophylactic use of, 945, 945t
 for animal bite-wounds, 167e-3, 167e-3t
 for bronchiectasis, 1696
 for cancer patient, 492
 for chronic neutropenia, 423
 for dental procedures, 242
 for infective endocarditis, 826–827, 826t
 for transplant recipient, 925t, 927
 resistance to, **933**. *See also specific drugs*
 in anaerobic bacteria, 1101, 1101t
 genomics in detection of, 769t, 773–775
 in gram-negative bacteria, 807, 1026–1027
 in health care–associated infections, 918
 mechanisms of, 933
 in *S. pneumoniae*, 807
 selection of, 775
 stewardship of, 918, 945–946, 1026, 1027
 susceptibility testing, 150e-8
 time-dependent, 935
- Anti- β_2 -glycoprotein antibodies, 2131,
 2134–2135, 2135t, 2754t
- Antibody(ies). *See also* Immunoglobulin(s);
specific antibodies
 affinity maturation of, 372e-21
 blood group, 138e-1–2, 138e-1t
 for cancer treatment. *See* Cancer biologic
 therapy
 cytotoxic reactions of, 372e-29
 definition of, 372e-1
 drug-dependent, 727
 in neurologic paraneoplastic syndromes, 614,
 614t
 in parasitic infections, 245e-6, 245e-6t
 to T and B cells, 372e-30–31
- Antibody-dependent cell-mediated cytotoxicity
 (ADCC), 372e-1, 372e-6, 1244
- Antibody enhancement, 1244
- Antibody isotope switching, 372e-22
- Anticardiolipin antibodies (aCL), 2131, 2135t,
 2754t
- Anti-Caspr2 antibodies, 614t, 617
- Anti-CCP antibodies, 2141, 2142f
- Anticholinergics. *See also specific drugs*
 adverse effects of, 81, 167, 241, 1969
 for asthma, 1676–1677
 for COPD, 1705
 for irritable bowel syndrome, 1969
 for lower urinary tract symptoms, 7e-4
 for nausea and vomiting, 260, 260t
 overdosage/poisoning with, 473e-1, 473e-8t
- Anticholinesterase agents. *See* Cholinesterase
 inhibitors
- Anticipation, 440
- Anticoagulant(s), **749**
 for acute limb ischemia, 1646
 for brain tumors, 598
 dental procedures and, 241–242
 drug interactions of, 43
 for DVT, 47, 1634–1635, 1635t
 intracranial hemorrhage due to, 2584
 for myocardial infarction, 1597, 1598t, 1606
 in noncardiogenic stroke prevention, 2571
 oral, 754, 756t, 1889t. *See also specific drugs*
 parenteral. *See specific drugs*
 for pulmonary embolism, 1634–1635, 1635t
 reversal of, 738
 for stroke prevention in cardiac disease, 1485t,
 1487, 1488, 2566t, 2570–2901
 for suppurative thrombophlebitis, 906
- Anticonvulsants. *See* Antiepileptic drugs (AEDs)
- Anti-CRMP5 antibody, 614t, 616
- Anti-dandruff shampoo, 347
- Antidepressants. *See also specific drugs*
 actions of, 465e-2
 adverse effects management, 2711t
 adverse effects of, 65, 81, 267, 378, 380, 2710t
 for depression, 2710t, 2716–2717, 2716f
 for fatigue, 153
 for insomnia, 65, 191
 for irritable bowel syndrome, 1969
 for pain, 91t, 93–94, 93t, 2672
 for personality disorders, 2720
- Antidiuretic hormone (ADH). *See* Arginine
 vasopressin (AVP)
- Antidotes, 473e-7. *See also specific drugs/poisons*
- Antidote Treatment Nerve Agent Autoinjector
 (ATNAA), 262e-6, 262e-6f
- Anti-DPPX antibodies, 614t, 617
- Antidromic AV reentry, 1482, 1483f
- Anti-dsNDA antibody, 2126t, 2131
- Antiemetics, 260, 260t
- Antiepileptic drugs (AEDs). *See also specific
 drugs*
 action of, 2548
 adverse effects of
 cobalamin malabsorption, 644t
 cross-sensitivity and, 385
 genetic considerations in, 385
 hair loss, 380
 hypersensitivity, 382. *See also* Drug-induced
 hypersensitivity syndrome (DIHS)
 incidence of, 377
 neurologic, 2553–2555t
 osteoporosis, 2492
 SJS/TEN, 383. *See also* Stevens-Johnson
 syndrome (SJS); Toxic epidermal
 necrolysis (TEN)
 systemic, 2553–2555t
 vitamin D deficiency, 2485
 for brain abscess, 902
 for brain tumors, 598
 breast-feeding and, 2559
 discontinuation of, 2556
 dosage monitoring, 36
 drug interactions of, 246e-10, 1793,
 2553–2555t, 2559
 for focal seizures, 2552, 2556
 for generalized seizures, 2556

- initiation of, 2552
 monitoring of, 2556
 for neurocysticercosis, 903
 overdose/poisoning with, 473e-10t
 for pain, 91t, 94
 pharmacology of, 2553–2555t
 in pregnancy, 2559
 for refractory epilepsy, 2556
 selection of, 2552, 2552t
- Antierythrocyte antibody, 2126t
- Antiestrogens, 530, 530t
- Anti-factor Xa assay, 407, 750, 752
- Antifibrinolytic drugs, 734
- Antifungal agents, **1330**. *See also specific drugs*
 prophylactic, in transplant recipients, 921
 in sepsis/septic shock, 1756, 1757t
 susceptibility testing, 150e-8
 topical, 1331–1332
- Anti-GABA antibodies, 614t, 617
- Anti-GAD antibodies, 614t, 617
- Antigen(s)
 B cell receptor for, 372e-1
 definition of, 372e-1
 macroscopic detection, 150e-2
 minor histocompatibility, 373e-6
 polysaccharide, selective antibody deficiency to, 2112
 red blood cell, 138e-1–2, 138e-1t
 T cell recognition of, 372e-18–19, 372e-18f
 TCR for, 372e-2
- Antigenic drift, 1209–1210, 1209t
- Antigenic shift, 1209, 1209t
- Antigen-presenting cells, 103e-22, 103e-22f
- Antiglomerular basement membrane (anti-GBM) disease
 clinical features of, 1835t, 1839
 genetic considerations in, 1833
 renal biopsy in, 62e-6f, 1839
- Antiglycolipid antibodies, 2695, 2696f
- Antihistamines. *See also specific drugs*
 adverse effects of, 273e-4
 for allergic rhinitis, 2122, 2123f
 for angioedema, 2119
 for atopic dermatitis, 354t
 for contact dermatitis, 354t
 for cough, 244
 for insect stings, 2749
 for insomnia, 191
 for nausea and vomiting, 260
 for otitis media, 230
 overdose/poisoning with, 473e-8t, 473e-13t
 for pityriasis rosea, 349
 as premedication for antivenom, 2737
 for pruritus, 345
 for scombroid poisoning, 2744
 for systemic mastocytosis, 2121
 for vertigo, 151t
- Antihistone antibody, 2126t
- Anti-Hu antibody, 614t, 616
- Anti-Hu paraneoplastic neuropathy, 2700
- Anti-insulin receptor antibodies, 2347t, 2348
- Anti-Jo-1, 2196
- Anti-La antibody, 2126t
- Antileukotrienes, 1678, 1680
- Anti-LGI1 antibodies, 614t, 617
- Antilymphocyte globulin (ALG), 1828
- Antimalarial drugs. *See also specific drugs*
 adverse effects of, 353, 1380t, 2132
 cutaneous, 347, 349, 360, 378, 379, 381
 in GP6D deficiency, 656, 656t
 seizure, 2547t
 for discoid lupus erythematosus, 375
 drug interactions of, 246e-1–2t, 798
 for malaria prophylaxis, 1383t
 for malaria treatment, 1378t
 overdose/poisoning with, 473e-11t, 473e-13t
 in pregnancy and lactation, 246e-1–2t
 properties of, 1380t
 resistance to, 14–15, 1370f, 1379, 1381f, 1384
 for SLE, 2132
 for subacute cutaneous lupus erythematosus, 375
- Anti-Ma proteins, 614t, 616, 616f
- Antimetabolites. *See also specific drugs*
 antibacterial, 931t, 932, 932f
 in cancer chemotherapy, 103e-10–11t, 103e-13–14
- Antimicrobial peptides, 372e-1, 372e-2, 372e-4t
- Antimicrobial stewardship, 918, 945–946, 1026, 1027
- Antimitochondrial antibodies (AMA), 2060
- Antimitotic agents, 103e-11–12t, 103e-14–15.
See also specific drugs
- Antimonial compounds. *See* Meglumine
 antimonate; Pentavalent antimonials
- Anti-müllerian hormone (AMH), 332, 594, 2376, 2378
- Antimycobacterial agents, **205e-1**. *See also*
 Tuberculosis, treatment of
- Antimyeloperoxidase (MPO) antibodies, 1840, 2180, 2193
- Antineuronal antibodies, 614, 614t, 2126t
- Antineutrophil antibodies, 418
- Antineutrophil cytoplasmic antibodies (ANCA)
 cytoplasmic, 2180
 in drug-induced vasculitis, 2193
 in eosinophilic granulomatosis with polyangiitis, 1685, 2180
 in granulomatosis with polyangiitis, 2180, 2184
 in microscopic polyangiitis, 2186
 perinuclear, 2180
 in relapsing polychondritis, 2204
 in vasculitis, 1840, 2180
- Anti-NMDA receptor antibodies, 614t, 615f, 617
- Antinuclear antibodies (ANA)
 in musculoskeletal disease, 2223, 2223t
 in SLE, 2126–2127, 2126t, 2129t, 2131, 2223t
 in systemic sclerosis, 2163
- Antiparasitic agents, **246e-1**, 246e-1–7t. *See also specific drugs*
- Anti-PD-1 agents, 499
- Antiphospholipid antibody(ies)
 classification of, 2135t
 laboratory evaluation of, 406, 2754t
 in SLE, 2126t, 2131
- Antiphospholipid syndrome (APS), **2134**
 in cancer, 613
 classification of, 2135t
 clinical features of, 1835t, 1865, 2135–2136, 2136t
 definition of, 2134–2135
 diagnosis of, 2135t, 2136
 differential diagnosis of, 2136
 epidemiology of, 2135
 high-altitude symptoms of, 476e-4
 pathogenesis of, 372e-24t, 377e-4, 1865, 2135
 renal manifestations of, 1865
 skin manifestations of, 369
 SLE and, 2133
 treatment of, 1865, 2136
- Anti-PL antibodies (aPL), 2135
- α_2 -Antiplasmin, 402f, 403, 758, 758f
- Antiplatelet agents, **745**
 action of, 745–746
 during and after PCI, 296e-2
 chronic use in cardiac disease, 2566t
 in development, 749
 discontinuation before dental procedures, 241–242
 discontinuation before lumbar puncture, 443e-1
 for ischemic heart disease, 1590
 for ischemic stroke, 2563
 for myocardial infarction, 1596, 1598, 1598t, 1606
 perioperative, 54
 for peripheral artery disease, 1644
 for stroke/TIA prevention, 2566t, 2569–2570
- Antiplatelet antibody, 2126t
- Antiporters (exchangers), 332e-6
- Antiprogestins, 2391
- Antiproteinase 3 antibodies, 1840
- Antipsychotics
 action of, 2721
 for acute alcohol intoxication, 2727
 adverse effects of, 2624, 2721–2722, 2722t
- for dementia, 175–176, 2602
 drug interactions of, 2711t
 overdose/poisoning with, 473e-8t, 473e-9t, 473e-12t, 473e-13t
 for schizophrenia, 2721–2722, 2722t
- Antipyretic agents, 126
- Antireflux surgery, 263
- Antiretroviral agents. *See* Combination antiretroviral therapy (cART)
- Anti-Ri antibody, 614t, 618
- Antiribosomal P antibody, 2126t
- Anti-RNP antibody, 2126t, 2757t
- Anti-Ro antibody, 2126t, 2131, 2133
- Anti-Saccharomyces cerevisiae antibodies, 2194
- Anti-Sm antibody, 2126t, 2131
- Antithrombin
 action of, 402, 402f, 405
 activation of, 749, 750f, 752
 deficiency of, 744
- Antithrombotic drugs, 745, 745f
 anticoagulants. *See* Anticoagulant(s)
 antiplatelet agents. *See* Antiplatelet agents
 fibrinolytic drugs. *See* Fibrinolytic drugs
- Antithrombotic mechanisms, 402, 402f
- Antithymocyte globulin (ATG)
 adverse effects of, 2071
 for aplastic anemia, 666–667
 in HSCT preparation, 139e-2
 for immunosuppression after liver transplantation, 2071
 for myelodysplasia, 671
 for paroxysmal nocturnal hemoglobinuria, 662
- Antithyroid drugs, 2296–2297. *See also specific drugs*
- Anti-TNF- α antibodies. *See* Tumor necrosis factor- α (TNF- α) inhibitors
- Antitoxin
 botulinum, 261e-9, 989
 diphtheria, 979
- Anti-Tr antibody, 614t, 618
- α_1 -Antitrypsin, 1706, 1945
- α_1 -Antitrypsin deficiency
 clinical features of
 cutaneous, 366
 granulomatosis with polyangiitis, 1840
 hepatic, 367e-2, 1992, 2062
 pulmonary, 1695, 1700–1701, 1703, 1705
 diagnosis of, 366e-4f, 367e-2, 1993t
 early interventions for, 450t
 epidemiology of, 367e-2
 genetic considerations in, 367e-2, 435
 genetic testing for, 443, 450t
 hepatocellular carcinoma and, 545, 545t
 treatment of, 367e-2
- Antitumor antibiotics, 103e-9–10t, 103e-13
- Antitussive agents, 245
- Antiuricosuric compounds, 431e-1
- Antivenom
 adverse effects of, 2737
 indications for, 2735
 for marine envenomations, 2739
 for scorpion stings, 2748
 for snakebite treatment, 2735, 2736t
 sources of, 2741–2742
- Anti-VGCC antibodies, 614t
- Antiviral therapy, **215e-1**. *See also* Combination antiretroviral therapy (cART); *specific drugs and diseases*
 for hepatitis virus infections, 215e-4–5t, 215e-9–11
 for herpesvirus infections, 215e-2–3t, 215e-6–9
 for respiratory infections, 215e-1, 215e-2t, 215e-5–6
- Anti-Yo antibody, 614t, 618
- Anton's syndrome, 2575
- Antrectomy, for duodenal ulcers, 1925, 1925f
- Anuria, 292
- Anxiety
 definition of, 2708
 erectile dysfunction and, 326
 female sexual dysfunction and, 330
 in respiratory acidosis, 322
 vs. syncope, 147

- Anxiety disorders, **2708**
 alcohol-induced, 2725
 dyspnea in, 47e-1
 gait disorders in, 164
 generalized anxiety disorder. *See* Generalized anxiety disorder
 at high altitudes, 476e-5
 insomnia in, 190
 nausea and vomiting in, 259
 obsessive-compulsive disorder, **2712**
 palpitations in, 254
 panic disorder, 96t, 101, **2708**
 phobic disorders, **2712**
 stress disorders, **2712**. *See also* Posttraumatic stress disorder (PTSD)
 vertigo in, 151
 in women, 6e-5
- Anxiolytics. *See* Benzodiazepine(s)
- AOA1 (ataxia with oculomotor apraxia), 451e-3t
- AOA2 (ataxia with oculomotor apraxia), 451e-3t
- Aorta
 acute occlusion of, 1642
 aneurysm of. *See* Aortic aneurysm
 atherosclerosis of, 1638, 1642
 coarctation of, **1525**, 1620
 congenital anomalies of, 1637
 dissection of. *See* Aortic dissection
 IgG4-related disease of, 391e-1t, 391e-2
 inflammatory disease of. *See* Aortitis
 rupture of, 289e-3, 1637
- Aortic aneurysm, **1637**
 abdominal, **1639**
 clinical features of, 118, 286, 1639
 diagnosis of, 118, 1639
 epidemiology of, 1639
 physical examination in, 1443, 1446
 rupture of, 1639
 screening for, 29t, 1639
 treatment of, 1640, 1640f
 chest pain in, 96t, 97
 cystic medial necrosis in, 1638
 etiology of, 1637-1638, 1638t
 familial, 1638
 infectious causes of, 1638
 in late syphilis, 1134, 1638
 monitoring expansion of, 1638
 mycotic. *See* Mycotic aneurysm
 rupture of, 104, 106t, 118, 1639
 thoracic, 386e-4f, 1638-1639, 1639f
 traumatic, 1638
 treatment of, 1639, 1640
- Aortic arch syndrome. *See* Takayasu's arteritis
- Aortic dissection, **1640**
 aortic regurgitation in, 1641
 chest pain in, 96t, 97, 99
 classification of, 1640, 1641f
 clinical features of, 1641
 diagnosis of, 19-20, 101, 1641
 intramural hematoma and, 97, 1640
 murmur in, 51e-6
 penetrating atherosclerotic ulcer and, 97, 1640
 risk factors for, 1640-1641
 treatment of, 1626t, 1641-1642
- Aortic occlusion, 1642
- Aortic regurgitation, **1534**
 in aortic dissection, 1641
 with aortic stenosis, 1551
 in aortitis, 1642
 carotid pulse in, 1445, 1445f
 clinical features of, 254, 1535-1536
 diagnosis of
 cardiac catheterization, 1536
 chest radiograph, 1536
 CMR, 270e-18, 270e-18f, 270e-27v
 ECG, 1536
 echocardiography, 270e-17-18, 270e-17f, 1536
 etiology of, 1529t, 1534-1535
 heart sounds/murmur in, 51e-1, 51e-1f, 51e-5-6, 1448f, 1449, 1536
 history in, 1535
 with mitral stenosis, 1551
 pathophysiology of, 1535
 in pregnancy, 46-47
 in relapsing polychondritis, 2204
 in rheumatic heart disease, 2151
 treatment of, **1536**, 1537f, 1538f
- Aortic root disease, 51e-5, 1535, 1538, 1538f, 2442
- Aortic root-to-right-heart shunts, 1523-1524
- Aortic stenosis, **1529**
 with aortic regurgitation, 1551
 in bicuspid aortic valve, 1529
 carotid pulse in, 1445, 1445f
 clinical features of, 1530-1531
 congenital, 1524, 1529
 diagnosis of
 cardiac catheterization, 1531-1532
 chest radiograph, 1531
 ECG, 1531
 echocardiography, 270e-16-17, 270e-17f, 271e-5f, 271e-5v, 297e-6f, 1531
 hemodynamic measurements, 1462, 1462f
 valve area, 1463
 etiology of, 1529, 1529t
 heart sounds/murmur in, 1449
 characteristics of, 1447-1448, 1447f, 1531
 duration, 51e-1
 intensity and radiation, 51e-2f, 51e-3
 systemic vascular resistance and, 51e-7
 with mitral regurgitation, 1551
 myocardial ischemia in, 1579
 natural history of, 1532
 palpitations in, 254
 pathogenesis of, 1529, 1530f
 pathophysiology of, 1530
 in pregnancy, 46-47
 rheumatic, 1529
 stroke prevention in, 2566t
 subaortic, 1525
 supraaortic, 1525
 treatment of
 balloon valvuloplasty, 296e-5, 1525, 1533
 medical, 1524-1525, 1532
 strategy for, 1533f
 valve replacement. *See* Aortic valve replacement
- Aortic valve
 bicuspid, 1529
 sclerosis, 51e-4
 stenosis. *See* Aortic stenosis
- Aortic valve replacement
 indications for, 1524-1525, 1532, 1533f, 1537, 1537f
 with mitral valve replacement, 1552
 paravalvular abscess following, 271e-10f
 surgical, 1532, 1532t
 transcatheter
 with mitral valve repair, 1552
 outcomes of, 296e-5, 297e-6f, 297e-7, 1534t
 patient selection for, 297e-6f, 1533-1534, 1533f, 1537f
 procedure for, 297e-7, 297e-7v
 valve models for, 296e-5, 297e-6f, 1534f
- Aortic valvuloplasty, 1533
- Aortitis, **1642**
 aneurysm in, 1638
 etiology of, 1638t
 in giant cell arteritis. *See* Giant cell arteritis (temporal arteritis)
 idiopathic, 1643
 IgG4-related, 391e-1t, 391e-2
 imaging in, 270e-24, 270e-25f
 infective, 1642-1643
 rheumatic, 1642
 syphilitic, 1134, 1136, 1643
 in Takayasu's arteritis. *See* Takayasu's arteritis
- Aortobifemoral bypass, 1645
- Aortocolic fistula, 277
- Aortoenteric fistula, 277
- Aortography, 1464
- Aortoiliac disease, 1643. *See also* Peripheral artery disease
- APACHE II scoring system, 1729, 1730f, 1730t
- APAF-1 gene, 102e-10
- APAF-1 protein, 102e-9f
- APC gene mutations
 in colorectal cancer, 101e-1f, 538, 539
 in familial adenomatous polyposis, 101e-5, 101e-6t
 in Gardner's syndrome, 599t
 germline and somatic, 101e-7f
 in thyroid cancer, 2305t
- APECED (autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy). *See* Autoimmune polyendocrine (polyglandular) syndrome-1 (APS-1)
- Apeú virus, 1309t
- Aphagia, 254
- Aphasia, 37e-1v, **177**
 in AD, 2598
 Broca's, 177t, 178
 caring for patient with, 184
 clinical examination of, 177
 clinical features of, 177t
 conduction, 177t, 178
 fluent transcortical, 177t, 178
 global, 177t, 178
 isolation, 177t, 178
 jargon, 177
 nonfluent transcortical, 177t, 178
 progressive, 37e-1v, 179-180
 pure alexia without agraphia, 177t, 178-179
 pure word deafness, 177t, 178
 subcortical, 178
 Wernicke's, 177-178, 177t
- Aphemia, 179
- Apheresis, 138e-2
- Aphthous ulcers
 etiology of, 237, 417
 in HIV infection, 1255-1256, 1256f
 in IBD, 1958
 prevention of, 423
 recurrent, 239t
- Apical ballooning syndrome, 289e-3
- Apixaban
 adverse effects of, 757
 for DVT/PE, 1635
 indications for, 756-757
 management before endoscopy, 1889t
 pharmacology of, 756t
 for stroke prevention in atrial fibrillation/atrial flutter, 757, 1485t, 1488, 2570
 for VTE prophylaxis, 1637t
- APL. *See* Acute promyelocytic leukemia (APL)
- aPL (anti-PL antibodies), 2135
- Aplasia cutis, 48, 2297
- Aplastic anemia, **663**
 acquired, 663t
 after liver transplantation, 2072
 classification of, 663t
 definition of, 663
 diagnosis of, 81e-7f, 665f, 666
 epidemiology of, 663
 etiology of
 chemical-induced, 663, 664t
 drug-induced, 664, 664t
 immunologic, 664
 infectious, 664
 inherited, 663t, 664-665
 paroxysmal nocturnal hemoglobinuria, 660, 664
 pregnancy, 664
 radiation-induced, 663
 history in, 665
 monocytopenia in, 422
 pathophysiology of, 665
 physical examination in, 666
 prognosis of, 666
 treatment of, 139e-2, 139e-5, 139e-5t, 666-667
- Aplastic crisis, 639
- Apnea
 sleep. *See* Sleep apnea
 in terminally ill patient, 69t
- Apnea testing, 1776
- APOBEC, 225e-2
- APOBEC3G, 214e-5
- Apobec-3G* gene, in HIV infection, 1240t
- APOBEC proteins, 1217
- Apo e4 allele, 2601

- APOE* gene, 94e-4, 435, 1241, 1241t, 2443
Apo ε gene, 2601
 Apoferritin, 625t
APOL1 gene mutations, 1811, 1832, 1847–1848
APOLI/MYH9 gene, 1241, 1241t, 1630
 Apolipoprotein(a), 2436t, 2437
 Apolipoprotein(s), 2436–2437, 2436t
 Apolipoprotein A-I (ApoA-I), 2436t, 2445, 2757t
 Apolipoprotein A-V (ApoA-V), 2436t, 2440, 2440t
 Apolipoprotein B, 1241t, 2436–2437, 2444, 2757t
 Apolipoprotein B-48, 2436t, 2437
 Apolipoprotein B-100, 2436t, 2437, 2440t, 2442
 Apolipoprotein C, 1241t
 Apolipoprotein C-II (ApoC-II), 2436t
 Apolipoprotein C-II (ApoC-II) deficiency, 2091, 2439–2440, 2440t
 Apolipoprotein E, 2436t, 2437, 2443
 Apomorphine, 263e-6, 2615, 2616t
 Aponeurotic ptosis, 207
 Apoplexy, pituitary, 2257
 Apoptosis
 of cancer cells, 102e-1t, 102e–9, 102e-9f, 103e-6, 103e-7f
 cellular and molecular control of, 372e-23, 372e-26f, 372e-27t
 definition of, 372e-1, 372e-4, 444e-9
 of foam cells, 291e-2
 in HIV infection, 1234
 mitochondria in, 444e-9
 in neurologic disease, 444e-9
 pathways of, 102e-8–9, 102e-9f, 372e-26f
 of red blood cells, 392
 in smoking, 1701
 in viral infection, 214e-6
 Apoptosis proteins, 372e-27t
 Apoptotic bodies, 103e-6
 Apotransferrin, 625
 APP. *See* Amyloid precursor protein (APP)
 Appendectomy, 1988
 Appendiceal fecalith, 1987, 1988f
 Appendiceal neuroendocrine tumors
 (carcinoids), 561t, 563, 567. *See also*
 Gastrointestinal neuroendocrine
 tumors (GI-NETs)
 Appendicitis, **1985**
 clinical features of, 1986–1987
 abdominal pain, 103, 106, 106t, 1986–1987, 1987t
 anorexia, 1987
 atypical patterns, 1987
 classic, 1987t
 nausea and vomiting, 258, 1987, 1987t
 complications of, 1985
 diagnosis of, 1987–1988
 differential diagnosis of, 877, 1986, 1986t
 in the elderly, 1987
 epidemiology of, 1985
 in immunocompromised patients, 1988
 pathogenesis of, 1986
 peritonitis in, 847
 in pregnancy, 1988
 radiographic studies in, 1988f
 treatment of, 1988
 Appendix, 1986, 1986f
 Appetite, 95e-5, 415e-2, 415e-2f
 Appetite suppressants. *See* Anorexiant
APP gene mutations, 172t, 2600–2601
 Apraxia
 in AD, 2599
 buccofacial, 179
 construction, 182
 definition of, 154
 dressing, 182
 gait, 163
 ideational, 179
 ideomotor, 179
 limb, 179
 limb-kinetic, 179
 oculomotor, 181
 of speech, 179
 Apremilast, 2177
 Aprepitant, 103e-25, 109, 260t, 474
 Apriso, 1963
 Aprosodia, 179
 APRT (adenine phosphoribosyltransferase)
 deficiency, 431e-5–6, 431e-5t
 APS. *See* Antiphospholipid syndrome (APS)
 APS-1. *See* Autoimmune polyendocrine
 (polyglandular) syndrome-1 (APS-1)
 APS-2. *See* Autoimmune polyendocrine
 (polyglandular) syndrome-2 (APS-2)
 aPTT. *See* Activated partial thromboplastin time
 (aPTT)
 APUDomas, 557. *See also* Gastrointestinal
 neuroendocrine tumors (GI-NETs)
 Aquagenic urticaria, 2118
 Aquaporins
 AVP action and, 2275, 2276f
 in collecting duct, 332e-9
 in loop of Henle, 332e-8
 in nephrogenic diabetes insipidus, 303, 1872
 in neuromyelitis optica, 2673
 in proximal tubule, 332e-6
 in water balance, 295–296, 296f, 332e-10, 332e-10f
 Araçatuba, 220e-1t, 220e-2
 Arachnidism, necrotic, 2747
 Arachnodactyly, 2512
 Arachnoid cysts, 2264
 Arachnoiditis, lumbar, 117
 Araraquara virus, 1307t
 Araucária virus, 1307t
 Arboviruses, 214e-6, 894
 Arbovirus infections, 892, 894
 ARBs. *See* Angiotensin receptor blockers (ARBs)
Arcanobacterium haemolyticum infections, 129t, 363, 981
 Archaea, in human microbiome, 86e-4
Arcobacter spp. infections, 1059, 1059t
 Arcus corneae, 436e-4f
 ARDS. *See* Acute respiratory distress syndrome
 (ARDS)
 Area postrema (chemoreceptor trigger zone), 258
 Arenaviruses
 global considerations in, 782
 reservoirs and vectors of, 1305, 1306t
 structure of, 214e-2t, 214e-3f, 1305
 Arenavirus infections
 clinical syndromes of, 1306t, 1312t
 geographic distribution of, 1311–1312t
 Junin/Argentinian hemorrhagic fever, 1320
 Lassa fever. *See* Lassa fever
 lymphocytic choriomeningitis (LCM), 891, 893, 908t, 1317–1318
 Machupo/Bolivian hemorrhagic fever, 1320
 Argasid (soft) tick, 2745
 Argatroban
 action of, 754
 discontinuation before lumbar puncture, 443e-1
 for heparin-induced thrombocytopenia, 728, 1461
 monitoring treatment with, 754
 pharmacology of, 753t
 Argentine hemorrhagic fever, 1320
 Arginase deficiency, 434e-3t
 Arginine, 434e-5
 Arginine metabolism disorders, 434e-3t
 Arginine vasopressin (AVP), **2274**
 action of, 2275, 2276f
 in ascites, 286
 for cardiac arrest, 1769, 1769f
 deficiency of
 in diabetes insipidus. *See* Diabetes insipidus
 (DI)
 in hypodipsic hypernatremia, **2279**, 2279f
 ectopic production of. *See* Syndrome of
 inappropriate antidiuretic hormone
 (SIADH)
 in edema, 250
 in heart failure, 1502f
 metabolism of, 2275
 secretion of, 2274–2275
 for septic shock, 1757
 for shock, 1750, 1751
 structure of, 2275f
 synthesis of, 295, 2274
 in thirst, 2275
 in water balance regulation, 64e-6, 295–296, 296f
 Arginine vasopressin (AVP) receptor-2
 antagonists, 2281–2282, 2282f
 Argininosuccinic acidemia, 434e-3t
 Argyll Robertson pupil, 195, 1136
 Argyria, 361
 Aripiprazole, 2721, 2722t
Aristolochia, 14e-4, 1861
 Aristolochic acid nephropathy, 577, 1804, 1856t, 1861
 Armodafinil, 153, 194
 Aromatase deficiency, 337
 Aromatase inhibitors
 action of, 102e-7
 adverse effects of, 103e-15, 2218t, 2492–2493
 for breast cancer, 102e-7, 103e-15, 528–529, 529t, 530t
 for breast cancer prevention, 478, 531
 for male precocious puberty, 2362
 for ovulatory induction, 2389
 Aromatic amines, 477t, 1692
 Aromatic L-amino acid decarboxylase, 372e-24t
 Arousal level, in coma, 1774
 ARR (aldosterone-renin-ratio), 2320t
 Arrhythmogenic right ventricular
 cardiomyopathy (ARVC)
 clinical features of, 1564
 diagnosis of, 1564, 1564f
 genetic considerations in, 1555t
 ventricular arrhythmias in, 1493f, 1495, 1495t
 ARSACS (autosomal recessive spastic ataxia of
 Charlevoix-Saguenay), 451e-3t
 Arsenic exposure/poisoning
 carcinogenicity of, 472e-1, 477t, 506
 clinical features of, 472e-2t, 2608, 2689
 diagnosis of, 472e-2t
 diarrhea in, 267
 global considerations in, 472e-2
 hyperpigmentation in, 361
 laboratory evaluation of, 2766t
 neuropathies in, 2688t, 2689
 optic neuropathy in, 203
 pathophysiology of, 472e-2t
 sources of, 472e-2t
 treatment of, 472e-2t
 Arsenic trioxide, 103e-19t, 686–687
 Arsine, 262e-1t
 Arteether, 246e-2t, 246e-7
 Artemether
 action of, 246e-7
 adverse effects of, 246e-2t, 1380t
 for malaria, 1378t, 1379, 1380t
 pharmacology of, 246e-7, 1380t
 in pregnancy and lactation, 246e-2t
 resistance to, 1379
 Artemisinin derivatives. *See* Artemether;
 Artesunate
 Arterial blood gases
 abnormal, approach to, 317
 clinical procedure tutorial, 485e-1v
 in hypothermia, 478e-2
 in interstitial lung diseases, 1710
 reference values for, 2757t
 in respiratory disease evaluation, 306e-7, 1663
 Arterial diseases, of extremities
 acrocyanosis, 1442, 1648f, **1649**, 1754
 acute limb ischemia, 1646
 arteriovenous fistula. *See* Arteriovenous fistula
 atheroembolism, 1646, 1646f
 erythromelalgia (erythralgia), 674, 677, 1648f, **1650**
 fibromuscular dysplasia, 1617, 1627, 1645, 2567
 frostbite. *See* Frostbite
 livedo reticularis, 68, 369, 1648f, **1649**, 2135
 peripheral artery disease. *See* Peripheral artery
 disease
 pernio (chilblain), 478e-4, 1648f, **1649**
 popliteal artery aneurysm, 1647

- Arterial diseases, of extremities (*Cont.*):
 popliteal artery entrapment, 1647
 Raynaud's phenomenon. *See* Raynaud's phenomenon
 thoracic outlet compression syndrome, 1646–1647
 thromboangiitis obliterans, 1645–1646, 1649
- Arterial O₂ concentration, 1748t
- Arterial spin labeling, in MRI, 440e-8
- Arterial thoracic outlet syndrome, 122
- Arterial thrombosis, **275, 741**
 in acute limb ischemia, 1646
 in antiphospholipid syndrome, 2135–2136, 2135t
 genetic considerations in, 277, 277t, 742–743, 743t
 lupus anticoagulant in, 740
 mesenteric, 1979–1980, 1980t
 platelets in, 275–276, 275f, 741–742, 741f, 745–746
 risk factors for, 405t, 745
 vascular disease and, 275, 741
 vs. venous thrombosis, 744–745
- Arterial volume, effective, reduction in, 250
- Arteries. *See* Blood vessels
- Arteriovenous fistula
 acquired, 1647
 clinical features of, 367, 1647
 congenital, 1647
 coronary, 1523
 diagnosis of, 1647
 for dialysis access, 1823
 dural, 2586
 pulmonary, cyanosis in, 249
 treatment of, 1647
- Arteriovenous malformation (AVM)
 cerebral, 2585–2586
 hemoptysis in, 245
 in Osler-Weber-Rendu syndrome, 732
 pulmonary, 308e-17f, 1666
 spinal cord, 2657, 2658f
- Arteriovenous O₂ difference, 1748t
- Arteritis
 giant cell (temporal). *See* Giant cell arteritis (temporal arteritis)
 necrotizing, 2567–2568
 obliterative, 391e-2–3
 Takayasu's. *See* Takayasu's arteritis
- Artery of Percheron, occlusion of, 2575
- Artery-to-artery embolic stroke, 2564–2565. *See also* Ischemic stroke
- Artesunate
 action of, 246e-7
 adverse effects of, 1380t
 drug interactions of, 246e-2t
 for malaria, 780t, 1378t, 1379, 1380t
 pharmacology of, 246e-7, 1380t
 in pregnancy and lactation, 246e-2t
 resistance to, 1379
- Arthritis/arthropathy. *See also* Osteoarthritis (OA)
 in acromegaly, 2240–2241
 in ankylosing spondylitis, 2170
 back pain in, 117
 bacterial. *See* Infectious (septic) arthritis
 in complex regional pain syndrome, 2246
 costochondritis and, 2246
 crystal-induced, 2233, 2233t
 calcium apatite deposition disease, 2236–2237, 2236t, 2237f
 calcium oxalate deposition disease, 2237, 2237f
 calcium pyrophosphate deposition disease, 2235–2236, 2235t, 2236f
 gout. *See* Gout
 differential diagnosis of, 833t
 enteropathic. *See* Enteropathic arthritis
 in familial Mediterranean fever, 2213
 gonococcal. *See* Gonococcal infections, disseminated
 in hemochromatosis, 2241
 in hemophilia, 2241–2242
 in HIV infection, 1261
 in hyperlipidemia, 2242–2243
 hypertrophic. *See* Hypertrophic osteoarthropathy
 in IBD, 1958
 infectious. *See* Infectious (septic) arthritis
 juvenile, 373e-7–8, 373e-8t
 in Lyme disease, 1151
 monoarticular, 833t
 in myofascial pain syndrome, 93, 237, 2246
 neoplasias and, 2246–2247
 neuropathic, 1136, 2243–2244, 2243f, 2243t
 new-onset, in hospitalized patient, 2219
 ochronotic, 434e-4
 paraneoplastic, 2247
 polyarticular, 833t
 postinfectious, 837. *See also* Reactive arthritis (Reiter's syndrome)
 psoriatic. *See* Psoriatic arthritis
 with rash, 1312t, 1313
 in relapsing polychondritis, 2203
 in sickle cell disease, 2242, 2242t
 in Sjögren's syndrome, 2167, 2167t
 in SLE, 2127, 2129t
 in thalassemia, 2242
 in Tietze syndrome, 2246
- Arthroconidia, 1335
- Arthrogryposis, 462e-9
- Arthropathy. *See* Musculoskeletal disorders
- Arthroplasty, 2233. *See also* Knee replacement
- Arthropod-borne viruses, **1304**. *See also specific viruses*
 diagnosis of, 1310
 disease syndromes of. *See also specific diseases*
 arthritis and rash, 1312t, 1313–1314
 encephalitis, 1312t, 1314–1317
 fever and myalgia, 1312t, 1314–1319
 hemorrhagic fevers. *See* Viral hemorrhagic fevers
 pulmonary disease, 1312t
 epidemiology of, 1310
 geographic distribution of, 1311–1312t
 reservoirs and vectors of, 1306–1310t
 transmission of, 1304
- Arthropod injuries, 2744. *See also specific organisms*
- Arthroscopy, 2233
- Artificial saliva, 216
- Artificial tears, 199
- Art of medicine, 1
- ARVC. *See* Arrhythmogenic right ventricular cardiomyopathy (ARVC)
- ARX gene mutations, 2354t
- Asacol, 1959
- Asbestos, 477t, 1688–1689, 1688t
- Asbestos-related disease
 asbestosis, 1689, 1689f
 chest imaging in, 1689, 1689f
 CT in, 308e-15f
 lung cancer, 506, 1689. *See also* Lung cancer mesothelioma, 1689. *See also* Mesothelioma risk of, 1689
 soft tissue sarcoma, 119e-1
- Ascaris lumbricoides*/ascariasis, **1413**
 clinical features of, 1413, 1414t, 1686–1687
 diagnosis of, 245e-3t, 245e-5t, 245e-6t, 765t, 1413, 2084
 epidemiology of, 245e-3t, 765t, 1413
 life cycle of parasite, 245e-3t, 1413
 treatment of, 246e-8, 246e-11–12, 1413–1414, 1414t
- Ascites, **286**
 in cardiovascular disease, 1443
 chylous, 287
 in cirrhosis, 250, 286, 2064, 2065f
 clinical features of, 2065
 complications of, 288, 2065–2066
 definition of, 251, 2064
 diagnosis of, 287–288, 287f, 2065
 etiology of, 285, 287
 in heart failure, 1504
 in hepatocellular carcinoma, 545
 with jaundice, 283
 in liver disease, 1992
 malignant, 288, 474
 in ovarian cancer, 593
 pancreatic, 288, 2087t, 2088
 pathogenesis of, 286–287, 287f, 2064–2065, 2065f
 peritonitis and, 288, 846, 2065–2066
 refractory, 2065, 2065f
 treatment of, 288, 2065, 2065f
- Ascorbic acid. *See* Vitamin C
- ASD. *See* Atrial septal defect (ASD)
- Asenapine, 2721, 2722t
- Asherman's syndrome, 335
- Ashkenazi Jewish population
 BRCA gene testing in, 101e-6
 dystonia in, 2620
 factor XI deficiency in, 736
 Gaucher's disease in, 432e-5. *See also* Gaucher's disease
 hearing loss in, 220
 PD in, 2609
 Tay-Sachs disease in, 432e-1. *See also* Tay-Sachs disease
- Ash leaf spot, 344, 359
- Asialoglycoprotein receptor, 372e-24t
- Asimultanagnosia, 2575
- Askin's tumor, 119e-4
- Asparaginase, 103e-11t, 103e-14, 607t, 1798
- Aspartate aminotransferase (AST)
 in acute viral hepatitis, 2016
 in alcoholic liver disease, 2053, 2053t
 in autoimmune hepatitis, 2050
 in chronic HBV infection, 2033
 in liver function evaluation, 1997, 1998t
 reference values for, 2757t
 in SLE, 2131
- Aspartylglucosaminuria, 432e-3t
- Aspergilloma, 1347, 1349t
- Aspergillus*
 aflatoxin B₁ and, 544
 antifungal susceptibility testing, 150e-8
 ecology of, 1345
 epidemiology of, 1345
 laboratory identification of, 1330, 1348
- Aspergillus* infections, **1345**
 in burn wounds, 166e-1, 166e-3
 in cancer patient, 486t, 488, 488t, 491
 clinical features of, 489, **1346**
 aspergilloma, 1347
 cerebral, 441e-9–10f, 1346–1347, 1346t
 chronic meningitis, 908t
 cutaneous, 25e-10f, 76e-12f, 135, 1346t, 1347
 endocarditis, 1346t, 1347
 keratitis, 1348
 otitis externa, 229, 1348
 respiratory tract. *See below*
 diagnosis of, 489, 1346t, 1348
 disseminated, 25e-10f, 1346
 epidemiology of, 1345, 1346t
 genetic considerations in, 1346
 in HIV infection, 1255
 pathogenesis of, 1345–1346
 prevention of, in hospitals, 491, 492, 845
 prognosis of, 1349
 prophylaxis for, 1349
 respiratory tract
 allergic bronchopulmonary. *See* Allergic bronchopulmonary aspergillosis (ABPA)
 allergic sinusitis, 1348
 in asthma, 1348
 in cancer patient, 489
 chronic pulmonary disease, 1347, 1347f
 chronic sinusitis, 228, 1347
 invasive pulmonary disease, 1346, 1346t
 invasive sinusitis, 441e-10–11f, 1346
 tracheobronchitis, 1346
 risk factors for, 1345–1346
 in transplant recipient, 320e-4, 920, 925, 926
 treatment of, 1348–1349, 1349f, 1349t
- Aspergillus terreus*, 491
- Asphyxiant/blood agents, 262e-1t, 262e-2t. *See also* Cyanide poisoning

- Aspiration
 cough in, 244
 in dysphagia, 257
 dyspnea in, 47e-4
 with enteral feeding, 98e-8–9
 in opioid use disorders, 468e-2
 pneumonia and, 804
 pneumonitis and, 1098. *See also* Pneumonitis of vomitus, 259
- Aspirin, **746**
 action of, 746, 746f
 for acute pericarditis, 1573
 adverse effects of
 anaphylaxis, 44
 asthma exacerbation, 1680
 bleeding, 404, 746
 cutaneous, 377, 381
 eosinophilia, 422
 gastric irritation/bleeding, 90, 746, 1590
 gout, 2218t
 immunologic factors in, 377
 nephropathy, 1861
 Reye's syndrome, 1184, 1185, 1213
 uricosuria, 431e-5
 allergy to, 746
 for antiphospholipid syndrome, 2136
 for atherosclerosis prevention, 291e-9
 for Behçet's syndrome, 2194
 for breast cancer prevention, 524
 for cardiac catheterization pretreatment, 1461
 chronic use in cardiac disease, 2566t
 with clopidogrel, 747
 for colorectal cancer prevention, 477, 539
 for coronary heart disease prevention, 29t, 291e-9
 dosage of, 746
 for erythromelalgia, 1650
 for fever, 126
 genetic variation in response to, 277, 277t, 742, 743t
 for giant cell arteritis, 2189
 indications for, 746
 for ischemic heart disease, 1590
 for ischemic stroke, 2563
 for Kawasaki disease, 2193
 low-dose, 34, 241, 277
 management of
 before endoscopic procedures, 1889t
 before lumbar puncture, 443e-1
 perioperative, 54
 for medication-overuse headache, 109
 metabolism of, 34
 for migraine, 2592t
 for myocardial infarction
 NSTEMI-ACS, 1596, 1598t
 as secondary prevention, 1610
 STEMI, 1603, 1606, 1610
 for niacin-associated flushing, 96e-4
 for pain, 90–91, 91t
 in PCI, 296e-1, 296e-2, 1591
 for peripheral artery disease, 1645
 for preeclampsia prevention in high-risk women, 45
 for rheumatic fever, 2152–2153
 for stroke/TIA prevention, 291e-9, 2569–2570
 for subacute thyroiditis, 2299
 for tension-type headache, 2594
 in women, 6e-3
- Aspirin resistance, 1596
 Aspirin-sensitive asthma, 1680
 Asplenia, 375e-1, 413, 788f, 2104–2105
 Assassin bug bite, 2750
 Assist control ventilation (ACMV), 1741, 1742t.
 See also Mechanical ventilation
- Assisted living, 83, 83t
 Assisted reproductive technologies, 2360, 2389
 Association cortex, 176, 176f
 Association study, 442, 442t
 Associative prosopagnosia, 182
 AST. *See* Aspartate aminotransferase (AST)
 Asstasia-abasia, 164
- Asteatotic eczema, 342f, 346
 Astereognosis, 160
 Asterixis, 1773
 Asthenia, 462e-1, 1212
 Asthma, **1669**
 acute severe, 1679
 air pollution and, 1670
 allergies and, 1669–1670, 1671
 aspirin-sensitive, 1680
 atopy and, 1669–1670
 brittle, 1680
 bronchopulmonary aspergillosis in, 1681
 cardiac, 1504
 chronic, 1678–1679
 clinical features of, 1675
 corticosteroid-resistant, 1680
 cough in, 244
 cough-variant, 244
 diagnosis of, 1675–1676, 1675f
 dietary factors in, 1670
 differential diagnosis of, 1676
 drug-induced, 1671
 dyspnea in, 47e-1t, 47e-2, 47e-2t, 47e-3
 in elderly, 1680
 environmental factors in, 1670–1671
 exercise-induced, 1671
 with fungal sensitization, 1348
 genetic considerations in, 1670
 high-altitude travel and, 476e-5
 hygiene hypothesis of, 1670
 infections and, 1670, 1671
 inflammatory mediators in, 372e-29t
 intrinsic, 1671
 noncompliance with medication for, 1679–1680
 occupational, 1670
 pathophysiology of, 306e-3, **1671**, 1673f
 airway hyperresponsiveness, 1675
 airway inflammation, 1672–1673, 1672f
 airway remodeling, 1675
 inflammatory effects, 1674–1675
 inflammatory mediators, 1673–1674, 1673f
 pathology, 1671–1672, 1672f
 physiology, 1675
 in pregnancy, 1680
 prevalence of, 1669
 pulmonary function values in, 306e-5, 306e-5f
 refractory, 1679–1680
 respiratory acidosis in, 322
 risk factors for, 1670t, 1671
 smoking and, 1680
 in surgical patient, 1680–1681
 treatment of, **1676**
 in acute severe disease, 1679
 alternative therapies, 1678
 anticholinergics, 1676–1677
 antileukotrienes, 1678
 β_2 -agonists, 1676
 bronchodilators, 1676
 in chronic disease, 1678–1679
 cromones, 1678
 future directions in, 1678
 goals of, 1676t, 1678–1679, 1679t
 immunotherapy, 1678
 inhaled corticosteroids, 1677, 1678f
 omalizumab, 1678
 patient education, 1679
 in refractory disease, 1680
 stepwise approach to, 1679, 1679f
 steroid-sparing therapies, 1678
 systemic corticosteroids, 1677–1678
 theophylline, 1677, 1677t
 triggers of, 1670t, 1671
- Astigmatism, 195
 Astrakhan spotted fever, 1157
 Astrocytomas
 glioblastoma, 599–600, 600f
 high-grade, 600
 intramedullary, 2654, 2654f
 low-grade, 599–600
 pilocytic, 441e-39f
 Astroviruses, 1286f, 1286t, 1289
- Asunaprevir, 2049
 ASXL1 gene mutations, 135e-6, 680, 680t
 Asymptomatic bacteriuria, 861, 863, 866, 868
 Asystole, 1769f. *See also* Cardiac arrest
 AT. *See* Atrial tachycardia (AT)
- Ataxia, **2626**
 approach to the patient, 2626–2627
 autosomal dominant
 dentatorubropallidolysian atrophy, 440t, 451e-2t, 2629
 episodic ataxia, 444e-2t, 451e-3t, 2629
 spinocerebellar ataxia. *See* Spinocerebellar ataxia (SCA)
 autosomal recessive. *See* Ataxia telangiectasia (AT); Friedreich's ataxia
 cerebellar. *See* Cerebellar ataxia
 channelopathies in, 444e-2t
 chemotherapy-induced, 607t
 clinical features of, 33e-1v, 2626
 diseases associated with, 2626–2627
 focal, 2627
 global considerations in, 2631
 mitochondrial, 2630
 in MS, 2663, 2672
 optic, 181
 pathophysiology of, 87e-2, 87e-3t
 postinfectious, 2626
 sensory, 158, 163, 164t
 symmetric, 2626–2627
 toxin-induced, 2626
 treatment of, 2630, 2672
 video library, 33e-1v
 with vitamin E deficiency, 2630
- Ataxia telangiectasia (AT)
 clinical features of, 451e-3t, 2109, 2630
 diagnosis of, 2109
 genetic considerations in
 ATM gene mutations, 101e-6t, 434, 451e-3t, 2630
 DNA repair defects, 375e-1, 434, 2630
 incidence of, 2109
 lymphoid malignancies in, 375e-1, 2109, 2630
 pancreatic cancer risk in, 554
 pathology of, 2630
 treatment of, 2630
- Ataxia-telangiectasia mutated (ATM), 102e-2
 Ataxia with oculomotor apraxia (AOA1), 451e-3t
 Ataxia with oculomotor apraxia (AOA2), 451e-3t
 Ataxin-1, 2628
 Ataxins, 2627
 Atazanavir
 adverse effects of, 1258, 1259, 1275t, 1278, 1280
 drug interactions of, 1278, 1280, 1921
 for HIV infection, 1275t, 1278
 molecular structure of, 1276f
 resistance to, 1278, 1280f
- Atelectasis, 1732
 Atenolol
 for adrenergic symptoms in Graves' disease, 2296
 for arrhythmias, 1480t
 dosage of, 1480t
 for hypertension, 1624t
 for ischemic heart disease, 1589t
 overdosage/poisoning with, 473e-9t
- ATF1 gene mutations, 101e-3t
 ATG. *See* Antithymocyte globulin (ATG)
 Atherectomy, 296e-2
 Athroembolism, 1629, 1646, 1647f, 1806t
 Atherosclerosis, **291e-1**
 aging and, 94e-4
 aortic, 1638, 1642
 cardiovascular risk reduction in, 291e-6t, 291e-7f, 291e-8f, 291e-8t
 carotid. *See* Carotid stenosis
 consequences of, 291e-4
 coronary. *See* Coronary artery disease (CAD)
 endothelium in, 265e-2
 in familial hypercholesterolemia, 2441–2442
 hypertension and, 1614
 inflammatory mediators in, 372e-29t
 intracranial, 2572

- Atherosclerosis (*Cont.*):
 lipoprotein disorders and, 2435. *See also*
 Lipoprotein disorders
 pathogenesis of, **291e-1**
 atheroma evolution in, 291e-1f, 291e-2-3
 foam-cell formation in, 291e-2
 initiation of, 291e-1-2, 291e-1f
 leukocyte recruitment in, 291e-2
 microvessel formation in, 291e-3
 plaque formation in, 291e-3, 292e-1v,
 1579
 plaque instability and rupture in, 291e-3f,
 291e-4, 291e-5f, 292e-1v, 1579, 1599.
See also Myocardial infarction (MI)
 plaque stabilization in, 291e-3f, 292e-1v
 peripheral. *See* Peripheral artery disease
 prevention of, 291e-8t, 291e-9, 291e-9f
 renal, 1835t, 1847. *See also* Renovascular
 disease
 risk factors for, 291e-6t, 2569
 assessment of, 291e-9-10
 diabetes mellitus, 291e-6-7
 dysregulated coagulation and fibrinolysis,
 291e-8-9
 gender, 291e-7-8
 hypertension, 291e-6
 inflammation, 291e-9, 291e-9f
 lipid disorders, 291e-4-6, 291e-7t
 postmenopausal state, 291e-7-8
 Atherothrombosis, 291e-3
 Athetosis, 2619t
 Athlete(s)
 androgen abuse in. *See* Anabolic-androgenic
 steroid (AAS) abuse
 concussion in, 457e-4
 sudden cardiac death in, 1767
Atlastin gene, 2634t, 2636
 ATM (ataxia-telangiectasia mutated), 102e-2
 ATM gene mutations
 actions of, 102e-2, 102e-2f, 434
 in ataxia-telangiectasia, 101e-6t, 434, 451e-3t,
 2630
 breast cancer risk and, 101e-6t
 pancreatic cancer risk and, 554
 in T-cell prolymphocytic leukemia, 135e-3
 ATN (acute tubular necrosis). *See* Acute kidney
 injury (AKI)
 ATNAA (Antidote Treatment Nerve Agent
 Autoinjector), 262e-6, 262e-6f
 Atomic bomb survivors, 678
 Atomoxetine, 2644
 Atonic seizures, 2543-2544
 Atopic dermatitis, **344**
 allergic contact dermatitis and, 345
 asthma and, 1669-1670
 clinical features of, 344t, 354t, 2119
 in children, 344
 distribution of skin lesions, 341t, 342f, 344,
 354t
 in infants, 344
 lichenification and scaling, 76e-1f, 344f
 morphology of skin lesions, 341t, 354t
 pruritus, 344
 diagnosis of, 354t
 differential diagnosis of, 2119
 erythroderma, 344t
 etiology of, 344
 genetic considerations in, 344
 immunopathology of, 372e-24t
 inflammatory mediators in, 372e-29t
 treatment of, 344-345, 354t, 391
Atopobium vaginae, 874
 Atopy, 1669-1670, 2114
 Atorvastatin, 42, 42t, 2218t, 2447, 2448t
 Atovaquone
 action of, 246e-7
 adverse effects of, 246e-2t, 1380t
 for *Babesia* infection, 780t, 1386, 1386t
 drug interactions of, 246e-2t
 for malaria, 1380t
 for malaria prophylaxis, 1382, 1383, 1383t
 for PCB, 1361t
 for PCP prophylaxis, 1251t, 1362t
 pharmacology of, 246e-7-8, 1380t
 in pregnancy and lactation, 246e-2t
 for toxoplasmosis prophylaxis in HIV
 infection, 1251t
ATP1A1 gene mutations, 306, 2589
ATP2B3 gene mutations, 306
ATP7B gene mutations, 367e-2, 2519
ATP8B1 gene mutations, 284
 ATP III (National Cholesterol Education Project
 Adult Treatment Panel III), 480e-3t
 ATRA. *See* All-trans-retinoic acid (ATRA)
Atrax robustus, 2748
 Atresia
 biliary, 2068, 2083
 esophageal, 1904
 Atrial fibrillation (AF), **1485**
 clinical features of, 1485-1486
 CT pulmonary valve mapping in, 271e-10f
 early onset, 436t
 ECG in, 278e-9f, 278e-10f, 278e-11f, 1487f
 epidemiology of, 1486
 in heart failure, 1514
 in hypertrophic cardiomyopathy, 1570
 mechanism of, 273e-2, 273e-3t
 in mitral stenosis, 1539
 palpitations in, 254
 paroxysmal, 1486, 1486f
 persistent, 1486, 1486f
 preexcited, 1482, 1483f
 in pulmonary edema, 1763
 risk factors for, 1486
 in SA node dysfunction, 1468
 stroke and, 1485t, 1488, 2564, 2567t, 2570
 syncope in, 1487, 1487f
 in thyrotoxicosis, 2294
 treatment of, **1487**
 cardioversion and anticoagulation, 1487
 catheter and surgical ablation, 1488-1489
 rate control, 1487-1488
 rhythm control, 1488-1489
 stroke prevention, 1485t, 1488, 2566t
 Atrial flutter
 common right, 1484, 1484f
 ECG in, 278e-8f, 278e-9f, 1484f
 treatment of, 1484
 vs. tremor artifact, 278e-5f
 Atrial natriuretic peptide (ANP)
 action of, 332e-6f, 332e-9
 in edema, 251
 in hyporeninemic hypoaldosteronism, 310
 in lung cancer, 511
 Atrial septal defect (ASD), **1520**
 AV conduction block in, 1472
 clinical features of, 297e-9, 1520-1521
 complications of, 1521
 diagnosis of
 cardiac catheterization, 1463
 ECG, 269e-11f, 1521
 echocardiography, 270e-25, 270e-25f, 297e-
 7f, 297e-8f, 1521, 1522f
 heart sounds/murmur, 51e-1, 51e-3-4, 51e-
 6, 1447f, 1448, 1521
 physical examination, 1521
 differential diagnosis of, 1521
 locations of, 297e-7f, 1521f
 vs. mitral stenosis, 1540
 ostium primum, 51e-4, 297e-7f, 1472, 1520
 ostium secundum, 51e-4, 297e-7f, 1520
 sinus venosus, 51e-4, 1520
 treatment of, 296e-4, 297e-7-9, 297e-7f, 297e-
 8f, 297e-9v, 1521
 Atrial tachycardia (AT)
 ECG in, 278e-5f, 278e-6f, 278e-7f, 1479f
 focal, 1477, 1478f, 1479, 1479f
 macroreentrant, 1485-1486
 mechanism of, 273e-3
 multifocal, 278e-5f, 1485, 1486f
 treatment of, 1485
 Atriofascicular pathways, 1481
 Atrioventricular (AV) bundle, 1470-1471
 Atrioventricular (AV) conduction block, **1471**
 in cardiac mesothelioma, 289e-2
 CMR in, 270e-22f, 271e-7f
 complete, 1471, 1472
 congenital, 1471t, 1472
 diagnosis of, 1473-1474
 electrophysiology of, 1472-1473
 epidemiology of, 1466
 etiology of, 1471-1472, 1471t
 first-degree, 269e-6f, 269e-16f, 1472-1473,
 1472f
 high-grade, 1473, 1473f, 1474f
 iatrogenic, 1472
 in Lyme disease, 278e-4f, 1151, 1472
 mechanism of, 273e-3t
 in myocardial infarction, 1609
 paroxysmal, 1473, 1473f
 second-degree, 278e-2f, 278e-3f, 278e-4f,
 1472-1473, 1472f
 treatment of, 1474-1476, 1475t, 1476t
 Atrioventricular (AV) nodal reentry tachycardia,
 278e-7f, 1478f, 1479-1480, 1481f
 Atrioventricular (AV) node, 273e-1, 1466,
 1470-1471
 Atrioventricular (AV) reentry tachycardia, 1478f,
 1481
 Atrophie blanche (livedoid vasculopathy), 369,
 369t, 1651
 Atrophie blanche en plaque, 1649. *See also*
 Livedo reticularis
 Atrophy, muscle, 462e-3-4
 Atropine
 adverse effects of, 254
 for AV conduction block, 1473, 1474
 for bradycardia, 1770
 for ciguatera poisoning, 2742
 for nerve agent exposure, 262e-5t, 262e-6,
 262e-6f, 473e-10t
 overdose/poisoning with, 473e-8t
 for SA node dysfunction, 1469
 for snakebite treatment, 2737t
ATRX gene mutations, 2354t
ATRX pathway, 632-633, 638
 Attention, assessment of, 168
 Attentional deficit, 183
 ATTR amyloidosis, 720t, 723-724. *See also*
 Amyloidosis
 Atypical adenomatous hyperplasia, 515
 Atypical chronic myeloid leukemia (aCML),
 135e-5-6, 135e-5t
 Atypical hemolytic uremic syndrome (aHUS),
 657, 730, 1832
 Atypical lipomatous tumor, 119e-1. *See also* Soft
 tissue sarcoma
 AUA Symptom Index, 587t
 AUC/MIC, 935, 935f
 Audiogram, 222
 Audiometry, 221-222
 AUDIT (Alcohol Use Disorder Screening Test),
 2727, 2727t
 AUDIT-C (Alcohol Use Disorders Identification
 Test-Consumption), 471e-5t
 Auditory brainstem responses, 223
 Auditory cortex, 176f
 Auditory nerve, 217
 Auer rod, 81e-9f, 681, 682f
 Aura, 2543, 2589
 Auricular cellulitis, 228
 Auricular chondritis, 2203, 2204f
 Auscultation
 abdominal, 286
 cardiac. *See* Heart murmurs; Heart sounds
 pulmonary, 47e-3, 1662
 Austin Flint murmur, 51e-6, 1449, 1537, 1540
 Australian bat lyssavirus, 1304
 Australian blue-ringed octopus envenomation,
 2740
 Australian jellyfish sting, 2739
 Autism spectrum disorders, 444e-1, 444e-6,
 465e-1, 465e-2t
 Autoantibody(ies)
 autoantigens and, 372e-24-25t
 blood group, 138e-1
 drug-induced, 44
 in inflammatory myopathies, 2196
 islet cells, 2404
 pathogenicity of, 377e-3t, 377e-4
 in SLE, 377e-5, 2125, 2126t, 2131
 Autocrine regulation/signaling, 102e-3, 400e-6

- Autofluorescence bronchoscopy, 1668
- Autoimmune autonomic ganglionopathy (AAG), 2642
- Autoimmune diseases, **377e-1**. *See also specific diseases*
- after HSCT, 139e-4
 - aging and, 94e-4
 - apoptosis protein defects and, 372e-27t, 377e-2
 - autoantigens associated with, 372e-23, 372e-24–25t
 - as cause of fever of unknown origin, 137t
 - cell- or organ-specific, 372e-24t
 - clinical features of, 377e-4, 377e-4t
 - cutaneous, **370**
 - cytokine defects and, 372e-27t
 - definition of, 372e-1
 - fever in, 126, 126t
 - genetic considerations in, 377e-3
 - in HIV infection, 1234
 - HLA molecule defects and, 372e-27t
 - immunopathogenesis of, 377e-3–4, 377e-3t, 377e-4t
 - immunotherapy for, 372e-30–31
 - mechanisms of, 377e-1–3, 377e-1t
 - molecular defects of immune system and, 372e-27t
 - myasthenia gravis and, 2703
 - organ-specific vs. systemic, 377e-4–5, 377e-5t
 - Sjögren's syndrome associated with, 2166, 2166t
 - systemic, 372e-24t
 - tissue damage in, 377e-3t, 377e-4
 - treatment of, 377e-5
 - types of, 377e-5t
 - in women, 6e-4
- Autoimmune encephalitis, 896
- Autoimmune hemolytic anemia, 377e-4, **658**
- Autoimmune hepatitis. *See* Hepatitis, autoimmune
- Autoimmune hypothyroidism, 2290–2292. *See also* Hypothyroidism
- Autoimmune lymphoproliferative syndrome (ALPS), 2112
- Autoimmune polyendocrine (polyglandular) syndrome-1 (APS-1)
- vs. APS-2, 2345t
 - autoantigens in, 372e-24t, 2345
 - clinical features of, 1343, 2113, 2323, 2324t, 2345, 2366
 - diagnosis of, 2345, 2346t
 - diseases associated with, 2344t, 2346t
 - etiology of, 2113, 2323
 - genetic considerations in, 377e-2, 2113, 2323, 2324t, 2345
 - treatment of, 2345
- Autoimmune polyendocrine (polyglandular) syndrome-2 (APS-2)
- vs. APS-1, 2345t
 - autoantigens in, 2347t
 - clinical features of, 2323, 2324t
 - diagnosis of, 2346–2347, 2346t
 - diseases associated with, 2344t, 2346, 2346t, 2347t
 - genetic considerations in, 2323, 2324t, 2346
 - treatment of, 2347
- Autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED). *See* Autoimmune polyendocrine (polyglandular) syndrome-1 (APS-1)
- Autoimmunity, **377e-1**
- cell- or organ-specific, 372e-24t
 - cytokine, 372e-25t
 - manifestations of, 377e-4, 377e-4t
 - mechanisms of, 372e-23, 377e-1–3, 377e-1t
 - paraneoplastic, 372e-25t
 - plasma protein, 372e-25t
 - prevention of, 377e-1, 377e-1t
 - primary immune deficiency diseases and, 2113
 - systemic, 372e-24t
- Autoinflammatory diseases. *See also specific diseases*
- definition of, 372e-1, 2212
 - fever in, 126, 126t
 - genetic considerations in, 372e-4, 372e-8t
 - hereditary, 2215
 - pyogenic arthritis, pyoderma gangrenosum, and acne syndrome, 372e-8t, 422, 2215
 - recurrent fever syndromes. *See* Hereditary recurrent fever syndromes
- Autolysins, 930
- Automated external defibrillation (AED), 1769
- Automaticity, cardiac, 273e-2–4
- Automatisms, 2543
- Autonomic dysfunction, **2637**
- approach to the patient, 2639–2641, 2640t
 - classification of, 2637–2638, 2638t
 - clinical features of, 2638–2639, 2639t
 - diagnosis of, 2640–2641, 2640t
 - drug-related, 2639, 2640t
 - syndromes of, **2641**
 - acute overactivity, 2643
 - in complex regional pain syndrome, 2643–2644
 - in diabetes mellitus, 2426, 2642
 - hereditary and sensory autonomic neuropathies, 2643
 - in hypoglycemia, 2432, 2432f
 - in multiple system atrophy, 2641. *See also* Multiple system atrophy (MSA)
 - orthostatic hypotension. *See* Orthostatic hypotension
 - paraneoplastic, 614t, 619
 - in peripheral nerve and neuromuscular junction disorders, 2642
 - postural orthostatic tachycardia syndrome, 1477, 2642–2643
 - primary hyperhidrosis, 2643
 - pure autonomic failure, 2642
 - with spinal cord lesions, 2641–2642
 - treatment of, 2644
- Autonomic dysreflexia, 2641–2642, 2660
- Autonomic nervous system
- anatomy of, 2637, 2637f
 - in blood pressure regulation, 1613
 - disorders of. *See* Autonomic dysfunction
 - normal activation of, 2638t
- Autonomic storm, 2643
- AutoPEEP, 1733, 1734f
- Autophagy
- in cancer cells, 102e-10, 103e-6
 - impaired, aging and, 94e-3
 - in microbial infection, 145e-7
 - in neurodegenerative diseases, 444e-9
- Autosomal dominant disorders, 436t, 437–438, 438f. *See also specific disorders*
- Autosomal dominant hypercholesterolemia, 2440t, 2442
- Autosomal dominant hypocalcemic hypercalciuria (ADHH), 2483
- Autosomal dominant nocturnal frontal lobe epilepsy (ADNFLE), 2545t
- Autosomal dominant partial epilepsy with auditory features (ADPEAF), 2545t
- Autosomal recessive disorders, 436t, 437, 437f. *See also specific disorders*
- Autosomal recessive hypercholesterolemia, 2440t, 2442
- Autosomal recessive spastic ataxia of Charlevoix-Saguenay (ARSACS), 451e-3t
- Autosome, 426
- Availability heuristic, 19
- Avalon virus, 1308t
- Avanafil
- action of, 325f, 326f
 - adverse effects of, 328, 329t
 - characteristics of, 329t
 - contraindications to, 328, 329t
 - for erectile dysfunction, 328
- AV conduction block. *See* Atrioventricular (AV) conduction block
- Avian influenza A viruses, 145e-3, 784, 917, 1210
- AVM. *See* Arteriovenous malformation (AVM)
- Avobenzon, 391, 391t
- Avoidant/restrictive food intake disorder, 2719
- AVP. *See* Arginine vasopressin (AVP)
- AVP-NP/II gene, 2276
- Axial osteomalacia, 426e-6
- Axial spondyloarthritis, 2169, 2169t. *See also* Ankylosing spondylitis
- Axillofemoral bypass, 1645
- Axitinib
- action and targets of, 102e-5t, 102e-17, 103e-18t
 - adverse effects of, 103e-18t
 - for renal cell carcinoma, 102e-17, 579
- Ayurvedic medicine, 14e-1t
- 5-Azacytidine
- action of, 103e-20
 - adverse effects of, 103e-11t, 671
 - for AML, 683
 - for myelodysplasia, 671
- 5-Aza-2'-deoxycytidine, 102e-8, 102e-8f
- Azathioprine
- action of, 943, 1829t
 - adverse effects of, 103e-10t, 1829t, 1960, 2130t, 2132
 - bone marrow suppression, 1831, 1960
 - carcinogenicity, 477t
 - carcinoma, 1962
 - major, 2182t
 - neutropenia, 417
 - pancreatitis, 1960
 - prevention of, 2182
 - pure red cell aplasia, 668t
 - for autoimmune hepatitis, 2051–2052
 - for Behçet's syndrome, 2194
 - for bullous pemphigoid, 372
 - for cold agglutinin disease, 659
 - drug interactions of, 42t, 103e-10t, 1828, 2705
 - for epidermolysis bullosa acquisita, 374
 - genetic variations in response to, 39t, 40, 445
 - for granulomatosis with polyangiitis, 2185
 - for IBD, 1960
 - for idiopathic pulmonary fibrosis, 1712
 - for immunosuppression, 1828, 1829t
 - for inflammatory myopathies, 2201
 - for interstitial lung diseases, 1711, 1712
 - metabolism of, 34t, 103e-10t
 - for MS, 2672
 - for mucous membrane pemphigoid, 374
 - for myasthenia gravis, 2705
 - for neuromyelitis optica, 2673
 - for pemphigus vulgaris, 371
 - for photoallergy, 390
 - for reactive arthritis, 2175
 - for sarcoidosis, 2211, 2212t
 - for SLE, 2130t, 2132
 - for vasculitis, 2182
- Azelaic acid, 352
- Azelastine nasal spray, 2122
- AZF gene mutations, 439
- Azidothymidine. *See* Zidovudine (AZT)
- Azithromycin
- action of, 205e-8, 931, 931t, 943
 - adverse effects of, 246e-4t, 943
 - for *Babesia* infections, 780t, 1386–1387, 1386t
 - for bite-wound infections, 167e-2
 - for bronchiectasis prophylaxis, 1696
 - for *C. trachomatis* infections, 1171, 1172
 - for cat-scratch disease, 1081t
 - for chancroid, 880t, 1013
 - for cholera, 1064
 - for community-acquired pneumonia, 767t, 808t
 - for COPD exacerbation prophylaxis, 1706
 - for donovanosis, 198e-2, 198e-2t
 - drug interactions of, 246e-4t
 - for endemic treponematoses, 207e-2
 - for enteric fever, 1052t
 - for gonococcal infections, 1008, 1009t
 - indications for, 940t, 943
 - for infective endocarditis prophylaxis, 826t
 - for *Legionella* infections, 1020t
 - for leptospirosis, 1145t
 - for *M. pneumoniae* infections, 1164
 - for mucopurulent cervicitis, 876
 - for nontuberculous mycobacterial infections, 205e-8
 - for otitis media, 229t
 - for pertussis, 1024t
 - for pharyngitis, 232f, 233

- Azithromycin (*Cont.*):
 for pneumonia, 953
 in pregnancy and lactation, 246e-4t, 937t
 prophylactic
 for *Bartonella* infection in HIV infection, 1252t
 for MAC infection in HIV infection, 1251t
 for meningitis, 889
 for reactive arthritis, 2175
 resistance to
 mechanisms of, 931t
 in nontuberculous mycobacteria, 205e-8
 prevalence of, 940t
 prevention of, 226
 in *S. pyogenes*, 233
 in *T. pallidum*, 1138
 for scrub typhus, 1159
 for *Shigella* infections, 1058t
 for streptococcal pharyngitis, 966
 for syphilis, 1138
 for trachoma, 1172
 for travelers' diarrhea, 797, 857t
 for urethritis, 871, 872t
- Azole antifungals, 1330–1331. *See also specific drugs*
- Azotemia, **290**. *See also* Acute kidney injury (AKI)
 approach to the patient, 290f, 291
 glomerular filtration rate and, 290–291
 in malabsorption syndrome, 1945
 with parenteral nutrition, 98e-7t
 in urinary tract obstruction, 1872
- AZT. *See* Zidovudine (AZT)
- Aztreonam
 indications for, 940t, 941–942
 for infections in cancer patient, 491
 resistance to, 940t
 for sepsis/septic shock, 1757t
- Azul. *See* Pinta
- Azurophil granules, 415
- B**
- BAALC overexpression, 680, 680t
- Babesia* spp.
 coinfections of, 781
 geographic distribution of, 245e-4t
 laboratory identification of, 250e-1, 250e-4f, 1386f
 life-cycle hosts of, 245e-4t
 transmission of, 245e-4t, 1384
- Babesia* spp. infections, **1384**
 in cancer patient, 484, 484t
 clinical features of, 1384–1385
 co-infection with Lyme disease, 1152, 1385
 complications of, 781
 diagnosis of, 150e-8, 245e-4t, 245e-6t, 250e-4f, 1385–1386, 1386f
 epidemiology of, 245e-4t, 1384
 etiology of, 780t, 781
 global considerations in, 1384, 1385f
 in immunocompromised patient, 1387
 pathogenesis of, 1385
 prevention of, 1387
 in splenectomized patient, 413
 transmission of, 138e-6, 1384
 treatment of, 780t, 1386–1387, 1386t
- Babes nodules, 1301
- Babinski sign, 154, 154t, 1772, 1775, 2539
- Bacillary angiomatosis, **1082**
 clinical features of, 133t, 1082, 1083f
 diagnosis of, 1083
 epidemiology of, 1082
 etiology of, 1082
 in HIV infection, 365, 1082, 1083f, 1268
 pathology of, 1082–1083
 prevention of, 1083
 skin manifestations of, 133t, 364t, 365
 treatment of, 832t, 1081t, 1083
- Bacillary peliosis, 1081t, 1082–1083
- Bacille Calmette-Guérin (BCG) vaccine, 1120, 1128
- Bacillus anthracis*, 261e-2. *See also* Anthrax
- Bacillus Calmette-Guérin (BCG), for bladder cancer, 576
- Bacillus cereus* infections
 clinical features of, 258, 266t, 853
 food-borne, 856, 856t
 in transplant recipient, 919
- Bacitracin, 930, 931t
- Back pain, **111**
 in ankylosing spondylitis, 2170, 2172
 in cancer patient, 117, 118, 530, 588, 1790–1792, 1791f
 causes of, **114**, 115t
 arthritis, 117
 autoimmune inflammatory disease, 118
 congenital anomalies, 118
 degenerative, 115–116, 116f
 infections/inflammation, 117, 784, 839
 lumbar disk disease, 113f, 114–115, 115f
 metabolic, 117–118, 426e-2
 metastatic disease, 2653
 postural, 118
 psychiatric, 118
 spondylolisthesis, 117
 spondylosis, 117
 traumatic, 117
 chronic, 119–120
 costs of, 111
 diagnosis of, 113–114, 440e-10
 in hemolysis, 393
 idiopathic, 118
 local, 111
 patient education in, 119
 physical examination in, 112–113, 114t
 radicular, 112
 referred from other locations, 111, 118
 risk factors for serious cause, 113t
 of spine origin, 111
 treatment of, **118**
 acute pain without radiculopathy, 119
 chronic pain without radiculopathy, 119–120
 clinical care systems view, 118–119
 nerve root and epidural spinal injections, 440e-10
 pain with radiculopathy, 120–121
 types of, 111–112
- Baclofen
 for dystonia, 2621
 for GERD, 263
 overdose/poisoning with, 473e-10t
 for pain, 61
 for spasticity, 2660, 2672
 for stiff-person syndrome, 618
 for tardive dyskinesia, 2624
 for trigeminal neuralgia, 2646
- Bacteremia
A. baumannii, 1037
 anaerobic bacterial, 1100
 chronic *B. quintana*, 1081t, 1082
 clostridial, 992
 definition of, 1751t
E. coli, 1029
Enterobacter, 1034
Klebsiella, 1033
L. monocytogenes, 983
 nontyphoidal *Salmonella*, 1054, 1054t
P. aeruginosa, 1044, 1045t
Proteus, 1033
 streptococcal, 968
- Bacteria
 adherence of, 145e-2–3, 145e-2t, 852
 antibiotic resistance in. *See* Antibacterial drugs, resistance to
 avoidance of host defenses by, 145e-4–7, 145e-6f
 colonic, 265
 culture-independent studies of, 86e-2–4, 86e-3f
 genome sequences of, 768, 770f, 776
 gram-negative. *See* Gram-negative bacteria
 gram-positive. *See* Gram-positive bacteria
 growth after entry, 145e-4
 host receptors of, 145e-3
 in human microbiome, 86e-2–3
 laboratory identification of, **150e-1**
 blood culture for, 150e-3
 gas-liquid chromatography for, 150e-3
 genomics in, 769t, 770–772
 limitations of, 770
 macroscopic antigen detection for, 150e-2
 microscopy for, 150e-1
 multiplex sequencing in, 86e-2–4, 86e-3f
 specimen-processing algorithm for, 150e-6f
 staining in, 150e-1–2
 susceptibility testing, 150e-8
 tissue invasion by, 145e-7–8, 145e-9–10
 toxin production by, 145e-9, 852–853
 viruses infecting, 86e-4
- Bacterial endocarditis. *See* Infective endocarditis
- Bacterial infections. *See also specific bacteria and bacterial types*
 anaerobic, mixed. *See* Anaerobic bacterial infections, mixed
 as cause of fever of unknown origin, 137t. *See also* Acute febrile illness
 in pregnancy, 49–50
 transfusion-transmitted, 138e-6
 in transplant recipient
 after HSCT, 919–920, 920t
 after kidney transplantation, 924–926, 924t, 1829–1831, 1830t
 after liver transplantation, 924t, 927–928, 1341f, 2072–2073
- Bacterial meningitis. *See* Meningitis, bacterial
- Bacterial overgrowth syndromes, **1944**
 cobalamin absorption in, 350e-1, 350e-1t
 diagnosis of, 271, 1941t, 1944
 diarrhea in, 269, 1944
 etiology of, 269, 1944
 pathogenesis of, 1944
 small-intestinal, 263, 1940t
 steatorrhea in, 1944
 in systemic sclerosis, 2162, 2164
 treatment of, 1944
- Bacterial pods, 145e-5
- Bacterial vaginosis
 clinical features of, 873t
 complications of, 873
 diagnosis of, 873t, 874, 874f, 875f
 etiology of, 873t, 874, 1099
 HIV transmission and, 1221
 treatment of, 873t, 875
- Bactericidal drugs, 930. *See also* Antibacterial drugs
- Bactericidal/permeability-increasing protein (Bpi), 372e-24t
- Bacteriostatic drugs, 930. *See also* Antibacterial drugs
- Bacteriuria. *See* Urinary tract infections (UTIs)
- Bacteroides* spp., 1094–1095
- Bacteroides fragilis*
 antibiotic resistance in, 1101
 gut immune functions of, 86e-8
 in normal human microbiota, 1094–1096, 1095t
 virulence factors of, 849
- Bacteroides fragilis* infections. *See also* Anaerobic bacterial infections, mixed
 bacteremia, 1100
 female genital tract, 849, 1099
 head and neck, 1097
 intraabdominal, 1099
 intraperitoneal abscess, 849
 liver abscess, 850
 pathogenesis of, 1096
 peritonitis, 847
 treatment of, 940t, 1101
- “Bad news” situation, communication in, 57–58, 57t
- BAEPs (brainstem auditory evoked potentials), 223, 442e-3
- Bagassosis, 1681t
- Baker's (popliteal) cyst, 1632, 2222
- Balamuthia* spp., 245e-4t, 1367, 1368
- Balamuthia* spp. infections
 in chronic meningitis, 908t
 clinical features of, 1368
 diagnosis of, 245e-4t, 245e-6t, 1368
 encephalitis, 896, 1368, 1368f
 epidemiology of, 245e-4t
 treatment of, 1368

- Balance
 anatomy and physiology of, 162
 disorders of, 162, 164. *See also* Fall(s) standing, 162
- Balanitis, 1343, 2174
- Balantidium coli*/balantidiasis, 246e-9, 1409
- Bald tongue, 241t
- Bálint's syndrome, 181, 209, 2575
- Balkan nephropathy, 577, 1804, 1856t, 1861
- Balloon angioplasty
 carotid, 2571–2572
 coronary, 296e-1, 296e-1f. *See also* Percutaneous coronary interventions (PCI)
- Balloon expulsion test, 274
- Balloon tamponade, 2064
- Balloon valvuloplasty, 296e-4, 1526
- Baló's concentric sclerosis, 441e-27–28f. *See also* Multiple sclerosis (MS)
- Balsalazide, 1959, 1959t
- Baltic myoclonus (Unverricht-Lundborg disease), 451e-3t
- “Bamboo spine,” 2169
- Banana spider, 2748
- Band(s), in peripheral blood smear, 81e-1–2
- Bangladesh, 13e-3f, 13e-6
- Bangui virus, 1306t
- Banna virus, 1310t
- Bannwarth's syndrome, 907t, 1150
- Barbiturates
 adverse effects of
 cross-sensitivity and, 385
 cutaneous, 360, 361
 folate deficiency, 646t
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 hypothermia, 478e-1
 mitochondrial defects and, 85e-9
 drug interactions of, 42t, 103e-9t
 for epilepsy, 2556
 overdose/poisoning with, 473e-10t
- Bar-coding, of medications, 12e-2
- BARDA (Biomedical Advanced Research and Development Authority), 261e-10
- Bardet-Biedl syndrome
 clinical features of, 415e-4, 1850t, 1855, 2256
 genetic considerations in, 415e-4, 1850t, 1855
 pathophysiology of, 1851f
- Bariatric surgery
 complications of, 2398, 2691
 hypoglycemia after, 569, 2434
 for metabolic syndrome, 2453
 for NALFD, 2057
 outcomes of, 2398
 procedures for, 2398, 2398f
 for type 2 diabetes mellitus, 2417
- Baricitinib, 2218t
- Barium radiography
 in colorectal cancer, 541f
 for colorectal cancer screening, 540
 in constipation, 273
 in dysphagia, 257
 in malabsorption syndromes, 1938, 1939f
 in nausea and vomiting, 260
 in peptic ulcer disease, 1919, 1919f
- Bark scorpion, 2748
- Barlow's syndrome. *See* Mitral valve prolapse
- Barmah Forest virus infection, 837, 1306t, 1313
- Baroreceptors, 142, 143f, 332e-10, 1613
- Baroreflex, 1613
- Barotrauma
 in divers, 477e-6–7
 in hyperbaric oxygen therapy, 477e-2
 in mechanical ventilation, 1743
 sinusitis and, 226
- Barrel chest deformity, 1443
- Barrett's esophagus/metaplasia
 dysplasia in, 1907, 1908f
 endoscopic findings in, 1881, 1882f, 1894–1895, 1907, 1907f
 endoscopic treatment of, 346e-1v
 esophageal cancer and, 262, 533, 1907–1908, 1907f
- GERD and, 262, 1894–1895, 1907
 in systemic sclerosis, 2161
- Barth's Index, 78t
- Barth's syndrome, 375e-1
- Bartonella* spp., 765t, 1078, 1079t, 1083
- Bartonella* spp. infections
 arthritis, 834
 bacillary angiomatosis. *See* Bacillary angiomatosis
 bacillary peliosis, 1081t, 1082–1083
 Carrión's disease, 1081t, 1083
 cat-scratch disease. *See* Cat-scratch disease
 endocarditis, 816, 817, 823t, 824, 1081t, 1082
 global considerations in, 1082
 in HIV infection, 1252t
 Oroya fever, 1081t, 1083
 trench fever, 1080–1082, 1081t
 verruga peruana, 829, 1081t, 1083
- Barter's syndrome
 antenatal, 306
 classic, 306
 clinical features of, 295, 306, 332e-8, 2483
 differential diagnosis of, 321
 genetic considerations in, 306, 332e-7t, 332e-8, 2461
 with sensorineural deafness, 332e-7t
 subtypes of, 332e-7t, 2483
- Basal cell carcinoma (BCC)
 clinical features of
 color, 364–365, 364t
 distribution, 341t, 342f, 500
 morphology, 76e-5f, 341t, 500, 501f
 epidemiology of, 500
 etiology of, 500
 genetic considerations in, 101e-6t, 500
 metastatic, 501
 micronodular, 500
 morpheiform (fibrosing), 500
 natural history of, 501
 nodular, 500
 pathophysiology of, 387, 500, 500f
 pigmented, 500
 prevention of, 502
 risk factors for, 500
 screening for, 483
 sun exposure and, 387, 500
 superficial, 500
 treatment of, 102e-5t, 386, 500, 500f, 501–502
- Basal cell nevus syndrome. *See* Gorlin's syndrome (nevoid basal cell carcinoma syndrome)
- Basal energy expenditure, 462
- Basal ganglia, 2611f, 2613, 2613f
- Basal meningitis, 906
- Basal (resting) metabolic rate, 73, 74f, 415e-2, 461
- BASDAI (Bath Ankylosing Spondylitis Disease Activity Index), 2171
- Baseline Dyspnea Index, 47e-2
- Basement membrane syndromes, 1835t, 1836, 1846
 Alport's syndrome. *See* Alport's syndrome
 antglomerular basement membrane disease. *See* Antiglomerular basement membrane (anti-GBM) disease
 Goodpasture's syndrome. *See* Goodpasture's syndrome
 nail-patella syndrome, 1835t, 1847
 pathogenesis of, 1846–1847
 thin basement membrane disease, 294, 1835t, 1847
- BASFI (Bath Ankylosing Spondylitis Functional Index), 2171
- Basic activities of daily living (ADLs), 72t
- Basic fibroblast growth factor (bFGF), 102e-14, 102e-15f, 1912, 2265
- Basic life support, for cardiac arrest, 1768
- Basilar artery occlusion, 220, 2577–2579, 2577–2580f
- Basilar skull fracture, 457e-2
- Basiliximab, 372e-30, 1517
- Basket cells, 702
- Basophil(s)
 in adaptive immunity, 372e-7t
 hematopoietic differentiation of, 89e-3f
 in innate immunity, 372e-7t, 372e-8–9, 372e-14f
 mediators released from, 372e-15t
 morphology of, 416f
 in peripheral blood smear, 81e-1, 81e-6f
- Basophilic stippling, 657, 659
- Basophil kallikrein of anaphylaxis (BK-A), 372e-15t
- Bassen-Kornzweig disease, 206, 2630
- Batai virus, 1308t
- Bath Ankylosing Spondylitis Disease Activity Index (BASDAI), 2171
- Bath Ankylosing Spondylitis Functional Index (BASFI), 2171
- Bath salts (synthetic cathinones), 469e-3–4
- Battlefield injuries. *See* War veterans
- Battle sign, 457e-2
- BAX, 103e-6, 103e-23, 444e-9
- Bayes' rule (theorem), 3, 21–22, 23f
- Baylisascaris procyonis* infections
 chronic meningitis, 908t
 diagnosis of, 245e-3t, 765t
 encephalitis, 896
 epidemiology of, 245e-3t, 765t
 life-cycle hosts of, 245e-3t
 visceral larva migrans in, 1411
- Bayou virus, 1307t
- Bazedoxifene, 2383, 2499
- Bazex-Dupré-Christol syndrome, 500
- BCC. *See* Basal cell carcinoma (BCC)
- B cell(s)
 deficiencies of, 375e-2, 2104t, 2110f, 2111
 in agammaglobulinemia. *See* Agammaglobulinemia
 in common variable immunodeficiency, 2111
 diagnosis of, 2111
 in hyper-IgM syndrome, 2110, 2110f, 2111
 in IgA deficiency, 2111–2112
 in selective antibody deficiency to polysaccharide antigens, 2112
 definition of, 372e-1
 development of, 372e-17f, 372e-20
 differentiation of, 89e-3f, 697–698, 698f, 710
 functions of, 372e-19–21
 in HIV infection, 1237–1238
 Ig gene rearrangement and, 372e-20–21
 locations of, 372e-19
 monoclonal antibodies to, 372e-30–31
 in rheumatoid arthritis, 2143
- B-cell activating factor, 2166
- B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma, 701
 anemia in, 393
 approach to the patient, 700
 clinical features of, 702t
 diagnosis of, 695t, 701–702
 epidemiology of, 695, 696f
 genetic considerations in, 699, 699t
 hypogammaglobulinemia in, 492
 immunology of, 698f, 699
 infections in, 484t, 485, 486t, 488
 lymphadenopathy in, 408
 lymphoid disorders that can present as, 695t
 mu heavy chain disease in, 719
 paraneoplastic syndromes in, 618
 peripheral blood smear in, 81e-10f, 702f
 prognosis of, 700
 pure red cell aplasia in, 668
 renal involvement in, 1859
 skin manifestations of, 372
 splenomegaly in, 411
 staging of, 700t
 treatment of, 703
 biologic therapy, 103e-21
 gene therapy, 91e-4
 HSCT, 139e-5t, 139e-6
 rituximab, 102e-5t, 102e-6, 103e-21, 703
 tumor markers in, 135e-2t
- B-cell prolymphocytic leukemia, 135e-1, 135e-2t
- B-cell prolymphocytic lymphoma, 706

- B cell receptors
activation of, 372e-19–20, 372e-20f
for antigen, 372e-1
defective rearrangements of, 2108
- BCG (Bacille Calmette-Guérin) vaccine, 1120, 1128
- BCL1* gene mutations, 101e-3t
- BCL2* gene mutations, 101e-3t
- BCL2* proteins, 102e-9–10, 102e-9f, 103e-6, 444e-9
- BCL6* gene mutations, 703
- BCNU. *See* Carmustine (BCNU)
- BCR-ABL1* oncogene, 41, 687–688, 688f, 689–690. *See also* Chronic myeloid leukemia (CML)
- BCX4430, 1328
- bdNA (branched-chain DNA) tests, 150e-7–8, 1247, 1247t
- BDNF (brain-derived neurotrophic factor), 444e-7t
- Beau's line, 380
- Beck Depression Inventory, 93
- Becker's disease, 462e-18
- Becker's muscular dystrophy, 462e-8
clinical features of, 462e-5t, 462e-8
diagnosis of, 462e-8
early interventions for, 450t
genetic considerations in, 436, 462e-5t
genetic testing for, 450t
treatment of, 462e-8
- Beck's triad, 1573
- Beckwith-Wiedemann syndrome, 83e-7, 439
- Beclomethasone dipropionate, 1679
- Bedaquiline, 205e-7, 1115, 1118–1119
- Bedbugs, 2750
- Bed nets, insecticide-treated, 14, 796
- Beef tapeworm. *See* *Taenia saginata*
- Beer potomania, 300, 301, 302
- Bee sting, 2116, 2748–2749
- Beetle vesication/dermatitis, 2750
- Bevor's sign, 2651
- Behavioral therapy. *See* Cognitive behavioral therapy (CBT)
- Behçet's syndrome, 2194
clinical features of, 2194
aortic aneurysm, 1638
chronic meningitis, 909t
cutaneous, 356t, 2194
diarrhea, 270
dysuria, 60e-1
neurologic, 441e-11f, 2194
ocular, 200, 2194
oral, 237, 239t
vasculitis, 2192, 2194
definition of, 2194
diagnosis of, 2194, 2194t
epidemiology of, 2194
genetic considerations in, 2194
HLA gene association with, 373e-7, 373e-8t
inflammasome activity in, 372e-8t
KIRs with, 372e-15t
pathogenesis of, 2194
treatment of, 2194
- Bejel. *See* Endemic syphilis
- Belatacept, 1829, 1829t
- Belimumab
adverse effects of, 2130t
for SLE, 2128f, 2130t, 2132, 2133
- Belladonna alkaloids, 473e-8t
- Bellini (collecting) duct tumors, 578, 578t. *See also* Renal cell carcinoma (RCC)
- Bell's palsy, 2647
antiviral therapy for, 1178
clinical features of, 2647
diagnosis of, 2648
differential diagnosis of, 2648
etiology of, 1178
imaging in, 441e-24f, 2648f
incidence of, 2647
oral pain in, 237
pathophysiology of, 2648
in pregnancy, 49
in relapsing fever, 1147
taste disorders in, 215
treatment of, 2648
- Bell's phenomenon, 2647
- Bence Jones proteins
laboratory evaluation of, 293, 1859, 1859f
in multiple myeloma, 293, 712, 715
reference values for, 2758t
- Bendamustine, 103e-8, 703, 704, 718
- Benedikt's syndrome, 208
- Benevolence, 17e-2
- Benign familial hematuria, 1847
- Benign familial neonatal seizures, 2545t
- Benign forgetfulness of the elderly, 171
- Benign hereditary chorea of childhood, 2622
- Benign paroxysmal positional vertigo, 149t, 150, 150f
- Benign prostatic hypertrophy (BPH)
clinical features of, 7e-4, 60e-1, 587, 1872
diagnosis of, 7e-4, 587, 587t
PSA levels in, 581
treatment of, 587
- Benign recurrent intrahepatic cholestasis (BRIC), 284, 2003t, 2004, 2076
- Benserazide/levodopa, 2614, 2616t
- Benzathine penicillin
for endemic treponematoses, 207e-3
for pharyngitis, 233, 965t
for rheumatic fever, 2152
for syphilis, 880t
- Benzbromarone, 2234
- Benzene exposure, 477t, 663, 678
- Benznidazole
action of, 246e-8
adverse effects of, 246e-3t, 246e-8, 1396, 1560
for Chagas disease, 246e-8, 1266, 1396, 1560
pharmacology of, 246e-8
- Benzoate, 434e-5
- Benzodiazepine(s). *See also* specific drugs
for acute alcohol intoxication, 2727
adverse effects of, 168, 478e-1
for alcohol withdrawal, 2727
for anxiety disorders, 2709–2710, 2711t
for delirium, 170
for dyspnea, 62
for insomnia, 62, 191
in mechanical ventilation, 1741
for nausea and vomiting, 260t, 261
in opiate overdose management, 468e-3
overdosage/poisoning with, 473e-10t
pharmacology of, 2711t
for restless legs syndrome, 192
for stress disorders, 2712
withdrawal from, 169, 2710–2551
- Benzonatate, 245
- Benzoyl peroxide, 352
- Benzphetamine, 2396
- Benztropine, 65, 2615, 2722
- Benzylpenicillin, 991
- BEP regimen, 590, 594
- Bereavement, 64, 69, 275
- Berg Balance Scale, 78t
- Beriberi. *See also* Thiamine (vitamin B₁) deficiency
cardiac manifestations of, 252, 290e-1–2, 1563
clinical features of, 96e-1t, 96e-2–3
dry, 96e-3, 2690
wet, 96e-2–3, 463t, 2690
- Berkelium exposure, 263e-7
- Berlin Questionnaire, 1721
- Bermejo virus, 1307t
- Bernard-Soulier syndrome, 729, 730
- Bernoulli effect, 306e-2
- Berry aneurysm, 1784–1785. *See also* Subarachnoid hemorrhage
- Beryllium, 1688t, 1691
- Beryllium disease/toxicity, 367, 373e-7, 1691
- Beryllium lymphocyte proliferation test, 1691
- β -adrenergic agonists
actions of, 265e-4, 265e-6, 265e-7f, 1676
adverse effects of, 65, 267, 305, 312, 1676
for asthma, 1676, 1679, 1679t
for COPD, 1705, 1707
for cough, 245
for high-altitude pulmonary edema, 476e-4
for hyperkalemia, 312
overdosage/poisoning with, 473e-3, 473e-8t
safety of, 1676
tolerance to, 1676
- β -adrenergic antagonists (beta blockers)
for accessory pathway-mediated tachycardia, 1483
for acute angle-closure glaucoma, 200
adverse effects of, 36, 1588–1589
alopecia, 354
arthralgias, 2218t
in asthma, 1671
AV conduction block, 1471t
cutaneous, 347, 353, 378
erectile dysfunction, 326
hair loss, 380
heat-related illness, 479e-1
Raynaud's phenomenon, 1649
SA node dysfunction, 1467t, 1469
xerostomia, 241
aldosterone-renin ratio effects of, 2320t
for aortic aneurysm, 1639, 1640
for aortic dissection, 1641
for atrial fibrillation, 1487–1488
for AV nodal reentrant tachycardia, 1479
contraindications to, 1588, 1603–1604, 2117
for cyclic vomiting syndrome, 261
drug interactions of, 43, 246e-10, 2711t
genetic variations in response to, 40
for glaucoma, 205
for heart failure, 1511, 1511f, 1512t
for hypertension, 1623–1624, 1624t
for hypertrophic cardiomyopathy, 1570
for inappropriate sinus tachycardia, 1477
for ischemic heart disease, 1588–1590, 1589t
for migraine prevention, 2593
for myocardial infarction, 1596, 1597t, 1603–1604, 1607, 1610
overdosage/poisoning with, 473e-9t, 473e-13t
for palpitations, 254
perioperative, 53–54
in peripheral artery disease, 1644, 1645
for pheochromocytoma, 2331
for phobic disorders, 2712
for radiation-induced cardiovascular disease, 621
for respiratory alkalosis, 324
for syncope, 145
for variceal bleeding, 277, 2063
for ventricular tachycardia, 1498
- β -adrenergic receptors, 37f, 40, 1613, 1745
- Beta blockers. *See* β -adrenergic antagonists (beta blockers)
- β -carotene
cancer and, 96e-7
for erythropoietic protoporphyria, 390, 2533
excess of, 463t
for lung cancer prevention, 477
for macular degeneration, 205
- Beta diversity, 86e-1t, 86e-5
- β -globin gene mutations, 433f, 435
- β -glucan test, 1330
- β_2 -glycoprotein, 372e-24t. *See also* Anti- β_2 -glycoprotein antibodies
- β -hemolytic streptococcal infections. *See* Group A *Streptococcus* (GAS) infections; Group B *Streptococcus* (GBS) infections
- β -IV spectrin, 372e-25t
- β -lactam antibiotics
action of, 930, 931t, 932f, 936–937
adverse effects of, 138, 938t, 942
for anaerobic bacterial infections, 1101
for cystic fibrosis, 1699
for cystitis, 867, 867t
for infective endocarditis, 963
for *P. aeruginosa* infections, 1044
for pneumococcal infections, 952
for pneumonia, 767t
resistance to
in anaerobic bacteria, 1101
detection of, 807
in *Klebsiella*, 1033
mechanisms of, 931t, 933, 976
 β -lactamase, 933, 1026, 1037

- Betamethasone, 347, 462e-19
 Beta (β) radiation, 263e-1
 Betaxolol, 1589t
 Bethanechol, 473e-10t, 2673, 2716
 Bethesda assay, 735, 740
 Bevacizumab
 action and targets of, 102e-5t, 102e-16f, 102e-17, 103e-21, 103e-21t
 adverse effects of, 102e-17, 103e-21, 103e-26, 520
 hemoptysis, 1794
 hypersensitivity reaction, 1798
 nephrotoxicity, 1804, 1864
 neurologic, 607, 607t
 for brain metastases, 605
 for breast cancer, 102e-17, 530-531
 for cholangiocarcinoma, 553
 for colorectal cancer, 102e-5t, 102e-17
 for esophageal cancer, 533
 for gastric cancer, 535
 for glioblastoma, 102e-5t, 600, 601f
 for hepatocellular carcinoma, 550-551
 for lung cancer, 102e-17, 520, 520t
 for macular degeneration, 205
 for ovarian cancer, 594
 for renal cell carcinoma, 102e-5t
 Bexarotene, 103e-17t, 103e-20
 Bezold-Jarisch reflex, 100
 bFGF (basic fibroblast growth factor), 102e-14, 102e-15f, 2265
 Bhanja virus, 1309t
 Bias, in screening tests, 479-480
BIB1 gene, 462e-14
 Bicalutamide, 103e-15, 585
 Bicarbonate
 in pancreatic secretion, 2090
 renal transport of, 332e-3f, 332e-6
 Bicarbonate therapy
 for DKA, 2419
 for hyperkalemia, 312
 for hypovolemia, 298
 for internal radionuclide contamination, 263e-6t
 for lactic acidosis, 318
 for metabolic acidosis, 318
 for salicylate-induced acidosis, 319
 for sepsis/septic shock, 1757
 for uremic acidosis, 320
 Bicarbonaturia, 299, 305
 Bicipital tendinitis and rupture, 2248
 Bicuspid aortic valve disease, 1529
 Bifascicular block, 1455, 1476, 1476t. *See also* Atrioventricular (AV) conduction block
 Bifid pulse, 1445
 Bifid uvula, 1443
 Biguanides. *See also* Metformin
 advantages of, 2414t
 adverse effects of, 318
 contraindications to, 2414t
 for diabetes mellitus, 2413, 2414t, 2416
 for metabolic syndrome, 2454
 pharmacology of, 2414t
 Bilateral simultaneous stimulation
 test, 160
 Bilberry, 404t
 Bile
 composition of, 2075
 effects on esophageal epithelium, 1907
 functions of, 1875
 hepatic, concentration within gallbladder, 2076
 leak, from duct of Luschka, 1885f
 limey, cholecystitis and, 2082
 regulation of flow of, 2075-2076
 secretion of, 2075-2076
 volume and electrolyte content of, 98e-4t
 Bile acid(s), 1934-1935, 1934f, 1934t, 2076, 2758t
 Bile acid diarrhea, 268, 1934-1935, 1935t
 Bile acid malabsorption, 268, 272
 Bile acid pool, 2076
 Bile acid sequestrants
 adverse effects of, 2415, 2448t, 2453
 drug interactions of, 42t
 for lipoprotein disorders, 2448t, 2449
 for metabolic syndrome, 2453
 for type 2 diabetes mellitus, 2415
 Bile duct(s)
 common, 2075
 imaging of, 346e-1v, 1892, 1895f
 obstruction of
 ERCP in, 1885f, 1892
 malignant, 1790
 vitamin K deficiency in, 96e-8
 resection, for cholangiocarcinoma, 553
 Bile duct disease, **2083**
 back pain in, 118
 choledocholithiasis. *See* Choledocholithiasis
 congenital anomalies, 2083
 extrinsic compression, 2084
 hemobilia, 2084
 hepatobiliary parasitism, 2084
 in HIV infection, 1258
 primary sclerosing cholangitis. *See* Primary sclerosing cholangitis (PSC)
 stones, 346e-1v, 1881, 1884f
 strictures, 2084
 trauma-related, 2084
 Bile reflux gastropathy, surgery-related, 1926-1927
 Bile salt excretory protein, 2004
 Bile salt export pump, 2075
 Biliary atresia, 2068, 2083
 Biliary cirrhosis, **2060**
 vs. autoimmune hepatitis, 2051
 clinical features of, 284, 2061
 coagulation disorders in, 739
 diagnosis of, 284, 366e-3f, 1992t, 2061
 diseases associated with, 545, 545t
 epidemiology of, 2060
 pathology of, 366e-3f, 2060-2061
 pathophysiology of, 372e-24t
 recurrence of, after liver transplantation, 2073-2074
 secondary, choledocholithiasis and, 2084
 skin manifestations of, 360, 360t, 365, 2061
 treatment of, 2061
 Biliary colic
 abdominal pain in, 103, 106, 262
 differential diagnosis of, 262
 in gallbladder disease, 2079-2080
 nausea and vomiting in, 258
 treatment of, 263
 Biliary dyskinesia, postcholecystectomy, 2082
 Biliary ectasia, congenital, 2083
 Biliary (hepatic) flukes
 cholangiocarcinoma and, 552
 clinical features of, 1428-1429
 diagnosis of, 245e-2t, 245e-5t, 245e-6t, 1429, 2084
 geographic distribution of, 1423t
 life cycle hosts of, 245e-2t
 transmission of, 1423t
 treatment of, 1428t, 1429
 Biliary hypoplasia, 2083
 Biliary sludge, 2077-2078
 Biliopancreatic diversion, 2398, 2398f
 Biliprotein, 280
 Bilirubin. *See also* Jaundice
 biliary excretion of, 2000
 conjugated, 280, 1998t, 1999-2000, 2000f
 delta fraction, 280
 extrahepatic disposition of, 2000
 in gut, 2000
 hepatocellular transport of, 1999-2000, 1999f
 metabolism of, 280, 1999-2000
 production of, 280, 2000
 renal excretion of, 2000
 serum, 280
 elevation of. *See* Hyperbilirubinemia
 in liver function evaluation, 283, 1995-1996, 1996f, 1998t
 measurement of, 280
 normal value, 280
 unconjugated, 280, 1996
 urine, 280, 1996, 1998t, 2000
 Bilirubinuria, 279
 Biliverdin, 280
 Biliverdin reductase, 280
 Bill & Melinda Gates Foundation, 8
 Billowing mitral leaflet syndrome. *See* Mitral valve prolapse
 Billroth procedures, 1925-1926, 1925f
 Binet staging system, for lymphoid malignancies, 700t
 Binge-eating disorder, **2720**
 Binswanger's disease (diffuse white matter disease), 163, 2568, 2602
 Bioavailability, 33-34, 33f
 Biocompatibility, 1822
 Biodegradable vascular scaffold (BVS) stent, 296e-2
 Biodosimetry Assessment Tool, 263e-7
 Biofeedback training, 274, 2590
 Biofilm, 145e-4, 166e-1
 Bioinformatics, 425
 Biolimus, 296e-2
 Biologic signal, 150e-1
 Biologic systems, 87e-1-2, 87e-2f. *See also* Systems biology
 Biologic therapy. *See also* Gene therapy
 cancer. *See* Cancer biologic therapy
 for IBD, 1961-1962
 for rheumatoid arthritis, 2145, 2146-2147t, 2147
 Biomass smoke-induced illness, 1693-1694, 1694f
 Biomedical Advanced Research and Development Authority (BARDA), 261e-10
 Bioprinting, 92e-2, 92e-3v
 Bioprosthetic heart valve. *See* Prosthetic heart valve
 Biopsy. *See also specific types and sites*
 in cancer diagnosis, 103e-1-2, 103e-1t, 468
 core-needle, 103e-2
 excisional, 103e-2
 fine-needle aspiration, 103e-2
 incisional, 103e-2
 specimen collection and transport, 150e-5t
 Bioterrorism, **261e-1**
 Category A agents, 261e-1, 261e-1t, 261e-2
 Category B agents, 261e-1, 261e-1t, 261e-9
 Category C agents, 261e-1, 261e-1t, 261e-9
 exposure in war veterans, 152e-8
 features of effective agents, 261e-1, 261e-7t
 preparedness for, 918
 prevention of and preparedness for, 261e-9-10
 weaponization of agents, 261e-1
 Biotin, 95e-2t, 96e-5, 98e-5t
 Bipolar cell antibodies, 614t
 Bipolar disorder, 2717, 2717t, 2718t
 Bipolar II disorder, 2717
Bipolaris spp., 228, 491
 Birbeck granules, 135e-6
 Bird fancier's lung, 1681, 1681t, 1682
 Birth weight, low, 333e-1
 Bisacodyl, 62t, 274
 Bischloromethyl ether, 506
 Bismuth-containing preparations, 1920t, 1921, 1923
 Bismuth poisoning, 240t
 Bismuth subsalicylate
 adverse effects of, 268, 379, 1921
 for diarrhea, 268
 in *H. pylori* eradication, 263, 1042t, 1922t, 1923
 for microscopic colitis, 270
 for travelers' diarrhea, 797, 857t
 for travelers' diarrhea prophylaxis, 268, 857
 Bisoprolol, 1512t, 1589t
 2,3-Bisphosphoglycerate, 632
 Bisphosphonates
 action of, 102e-11, 2501
 adverse effects of
 actinomycotic osteomyelitis, 1088, 1089f
 atypical femur fracture, 2501, 2501f
 osteonecrosis of jaw, 242, 623, 717, 2218t, 2501

- Bisphosphonates (Cont.):**
 for bone pain, 61, 119e-4
 for fibrous dysplasia, 426e-7
 for hypercalcemia, 314, 610, 2480t, 2481
 for metastatic bone disease, 119e-4
 for metastatic breast cancer, 530
 for multiple myeloma, 717
 for osteoporosis treatment, 2499–2501, 2500f, 2504
 for Paget's disease of bone, 426e-3–4, 426e-3t
 prophylactic
 in breast cancer, 529
 in osteoporosis, 2383
 in prostate cancer, 586
- Bite cells, 655**
- Bite-wounds, 167e-1**
 antibiotic prophylaxis, 167e-3
 approach to the patient, 945t
 cat. *See* Cat(s), bite-wounds
 dog. *See* Dog(s), bite-wounds
 epidemiology of, 167e-1
 human, 167e-2, 167e-3t, 830
 infections of, 167e-1–2, 167e-3t
 other animals, 167e-1–2
 rabies and tetanus prophylaxis, 167e-3
 treatment of, 167e-2–3, 167e-3t, 832t
- Bithionol, 389t**
- Bitot's spots, 96e-6**
- Bitter taste sensation, 212**
- Bivalirudin**
 action of, 754
 discontinuation before lumbar puncture, 443e-1
 vs. heparin, 754
 for heparin-induced thrombocytopenia, 728, 1461
 for myocardial infarction, 1597, 1598t
 in PCI, 296e-1
 pharmacology of, 753t
- BK-A (basophil kallikrein of anaphylaxis), 372e-15t**
- BK virus infection**
 in cancer patient, 490
 in transplant recipient
 after HSCT, 920t, 923
 after kidney transplant, 924t, 925, 926, 1831
- Black Creek Canal virus, 1307t**
- Black Death, 261e-5**
- Black hairy tongue, 240t**
- Blackhead, 351**
- Blackout, 2724**
- Black sea urchin envenomation, 2740**
- Black widow spider bite, 105, 2747–2748**
- Bladder**
 distention of, 286
 dysfunction of, 2660, 2663
 obstruction of, 104
 pain in, **60e-1**. *See also* Interstitial cystitis/
 bladder pain syndrome
- Bladder cancer, 575**
 adenocarcinoma, 575
 carcinoma of unknown primary and, 120e-2t
 chemotherapy-related, 575, 622–623
 clinical features of, 576
 deaths from, 467t, 575
 diagnosis of, 576
 epidemiology of, 575
 incidence of, 467t, 468f, 575
 pathogenesis of, 575–576
 pathology of, 575
 prognosis of, 577t
 risk factors for, 575
 staging of, 575, 576, 576f
 superficial, 577t
 transitional cell, 575
 treatment of, 576–577, 577t
- Bladder neck obstruction, 1805**
- Blastic NK cell lymphoma, 135e-4**
- Blastic phase, in CML, 693–694**
- Blast injury, 457e-1, 457e-2**
- Blastocystis hominis infection, 1409**
- Blastomyces spp., 145e-2t, 145e-3, 1338, 1338f**
Blastomyces spp. infections, **1337**
 approach to the patient, 1338
 clinical features of, 234, 829, 836, 908t, 1338
 diagnosis of, 1338
 epidemiology of, 1337
 in immunocompromised patient, 925t
 pathogenesis of, 1337–1338
 prognosis of, 1338
 treatment of, 1338–1339, 1338t
- Blau's syndrome, 2215**
- Bleeding, 400**
 approach to the patient, **403**
 in chronic kidney disease, 1818
 in DIC, 737
 drug-induced, 404, 404t
 fondaparinux, 753
 heparin, 751
 low-molecular-weight heparin, 753
 warfarin, 756
 gastrointestinal. *See* Gastrointestinal bleeding
 in hemophilia, 734
 history in, 403–404
 laboratory evaluation of, 405–407
 in liver disease, 739t
 systemic diseases that cause/exacerbate, 404
 variceal. *See* Variceal bleeding
- Bleeding time, 405**
- Bleomycin**
 action of, 103e-13
 adverse effects of, 103e-9t, 103e-13, 590
 cutaneous, 379, 2218t
 gonadal dysfunction, 103e-25
 hair loss, 380
 hemolytic-uremic syndrome, 1796
 in hyperbaric oxygen therapy, 477e-2
 hyperpigmentation, 360, 360t
 nausea and vomiting, 103e-25
 neurologic, 1794
 pulmonary, 489, 590, 621, 1797
 pulmonary fibrosis, 103e-13
 Raynaud's phenomenon, 590, 623, 1649
 for gestational trophoblastic disease, 597
 for ovarian germ cell tumors, 594
 for pericardial effusion, 474
 for pleural effusion, 474
 for testicular cancer, 590
- Blepharitis, 199, 2746**
- Blepharoptosis, 207**
- Blepharospasm, 2620, 2648–2649**
- Blinatumomab, 103e-22**
- Blind loop syndrome. See Bacterial overgrowth syndromes**
- Blindness. See Visual loss**
- Blind spot, 197**
- Blink reflexes, 442e-6**
- Blister beetle, 2750**
- Blister cells, 655**
- Blistering/vesicant agents, 262e-1t, 262e-2t. See also Sulfur mustard**
- BLM gene mutations, 101e-6t**
- Blomstrand's lethal chondrodysplasia, 2476f**
- Blood**
 donation, directed/designated, 138e-6
 laboratory evaluation of
 automated microbial detection, 150e-3
 culture for pathogens, 150e-3, 150e-4t
 for parasite detection, 245e-1–5, 245e-6t
 peripheral blood smear. *See* Peripheral blood smear
 reference values for, 2754–2756t
 transfusions. *See* Transfusion(s)
- Blood-brain barrier, 33, 37f**
- Blood culture, 806, 819–820, 824**
- Blood flukes. See Schistosoma spp.**
- Blood gases. See Arterial blood gases**
- Blood glucose monitoring, 2409–2410. See also Glycemic control**
- Blood group antigens and antibodies, 138e-1, 138e-1t**
- Blood pressure. See also Hypertension**
 age-related changes in, 1611
 control of, 265e-3
 goals for, 1616, 1616t, 1626
 measurement of, 1444–1445, 1616, 1621–1622
- in pregnancy, 45
 regulation of, **1612**
 arterial pressure in, 1612f
 autonomic nervous system in, 1613
 intravascular volume in, 1612–1613
 renin-angiotensin-aldosterone system in, 1613–1614, 1614f
 vascular mechanisms in, 1614–1615
 response to Valsalva maneuver, 2640, 2640t
 screening, 29t
- Blood products, contamination of**
 bacterial, 138e-6
 hepatitis viruses, 138e-3t, 138e-6, 735
 HIV, 138e-3t, 138e-6, 735, 1220t, 1222
 infectious, 138e-3t, 138e-6
 malaria, 1374
 screening for, 1222
T. cruzi, 1394
Yersinia, 1077
- Blood substitutes, 662**
- Blood urea nitrogen (BUN)**
 causes of abnormal values, 464t
 in hyponatremia, 301
 in hypovolemia, 298
 in nutritional assessment, 464t
 in prerenal azotemia, 298
- Blood vessels**
 biology of, **265e-1**
 engineered tissue for replacement of, 92e-3t
 origin of vascular cells, 265e-1
 pharmacogenetics of, 265e-4–5
 regeneration of, 265e-4
 smooth muscles of, 265e-1f, 265e-3–4
 in tumor angiogenesis, 102e-13–15, 102e-14f, 102e-16f
 vascular cell biology of, 265e-1–4
- Blood volume, 332e-10–11, 332e-10f**
- Bloom's syndrome, 101e-6t, 375e-1, 427**
- "Blueberry muffin syndrome," 230e-1, 230e-2t. See also Rubella (German measles), congenital**
- "Blue bloaters," 1704**
- Bluebottle jellyfish sting, 2739**
- Blue halo effect, 328**
- Blue light, for actinic keratosis, 386**
- Blue macules, 360**
- Blue nevus, 364t, 367**
- Blue River virus, 1307t**
- Blue rubber bleb syndrome, 364t, 367**
- "Blue toe" syndrome, 1646, 1647f**
- Blumer's shelf, 534**
- B lymphocytes. See B cell(s)**
- BMI. See Body mass index (BMI)**
- Bmi-1 gene, 89e-2, 89e-4**
- BNP. See B-type natriuretic peptide (BNP)**
- Bocaviruses, 1197**
- Boceprevir**
 action of, 215e-11
 adverse effects of, 2047
 for chronic HCV infection, 2044, 2045–2046t, 2047–2048
 drug interactions of, 2047
- Body composition, 71, 73t, 74f**
- Body fat, 6e-4, 462. See also Obesity**
- Body fluids**
 composition of, 295–297
 detection of parasites in, 245e-1–2, 245e-6t
 specimen collection and transport, 150e-4–5t
- Body image, 7e-2**
- Body lice (Pediculus humanus), 1146, 2745–2746**
- Body mass index (BMI)**
 calculation of, 415e-1, 415e-1f, 462, 2392
 nutritional status and, 98e-1t
 reference values for, 2393t
- Body plethysmography, 306e-6**
- Body temperature, 123–124**
- Boerhaave's syndrome, 98, 1910**
- Bohr effect, 323, 393, 632, 632f, 636**
- Bolivian (Machupo) hemorrhagic fever, 1320**
- Bombay phenotype, 138e-1**
- Bone**
 growth of, 2456
 metabolism of, 2454–2456, 2455f
 remodeling of, 2456, 2456f, 2467

- resorption of
 biochemical markers of, 2494–2495
 estrogen status and, 2491–2492, 2492f
 mechanism of, 2489–2490, 2491f
 structure of, 2454
- Bone age, 2257
- Bone-anchored hearing aid, 223
- Bone-conduction threshold, 222
- Bone disease
 anaerobic bacterial infections, 1099–1100
 in chronic kidney disease, 1815
 in cirrhosis, 2067
 with defective mineralization, 426e-5–6
 dysplasias, 426e-7
 extraskelatal calcification and ossification, 426e-7–8
 fibrous dysplasia, 426e-6, 426e-6f
 in Gaucher disease, 432e-5
 in IBD, 1958
 metastases from other sites, 102e-10–11, **119e-4**
 in multiple myeloma. *See* Multiple myeloma, clinical features of
 in osteogenesis imperfecta, 2508
 Paget's disease. *See* Paget's disease of bone
 primary immunodeficiencies with, 375e-2
 sarcoma. *See* Bone sarcoma
 sclerosing, 426e-4–5
 in tuberculosis, 1110–1111
- Bone loss. *See* Osteoporosis
- Bone marrow/bone marrow examination
 in anemia, 397
 in aplastic anemia, 81e-7f, 665f, 666
 cellularity in, 81e-7f, 663t
 in CML, 689, 689t
 collection for transplantation, 139e-1
 differential diagnosis of "dry tap," 135e-2t
 erythroid hyperplasia in, 81e-8f, 397f
 hypocellular, 663t
 iron stores, 81e-8f, 628, 628t
 in megaloblastic anemia, 643, 644f
 in myelodysplastic syndromes, 668f, 670–671
 in myelofibrosis, 675, 675f. *See also* Primary myelofibrosis
 myeloid hyperplasia in, 81e-8f, 397f
 normal, 81e-7f, 397f
 in sarcoidosis, 2208
 stem cell niche sites in, 89e-2
- Bone marrow failure, **662**
 in aplastic anemia. *See* Aplastic anemia
 bleeding tendencies in, 404
 differential diagnosis of, 663t
 in eosinophilic fasciitis, 376, 664
 pathognomic cells in, 668f
- Bone marrow transplantation. *See* Hematopoietic stem cell transplantation
- Bone mass/strength
 aging and, 71
 gut microbiota and, 86e-7
 in IBD, 1958
 measurement of, 2493, 2493f, 2493t
 sex differences in, 6e-4
- Bone pain
 at end of life, 119e-4
 in Gaucher disease, 432e-5
 in malabsorption syndromes, 1945
 in metastatic bone disease, 119e-4
 in sickle cell anemia, 634, 635
 treatment of, 61, 119e-4
- Bone sarcoma, **119e-3**
 chondrosarcoma, 101e-6t, **119e-3**
 classification of, 119e-3
 epidemiology of, 119e-3
 Ewing's sarcoma. *See* Ewing's sarcoma
 incidence of, 119e-3
 osteosarcoma, 101e-2t, **119e-3**, 236
 staging of, 119e-3, 119e-3t
- Bone scan, 426e-2–3, 582
- Bone spicules, 206, 206f
- Bone spurs, 2493
- Bordetella* spp., 1021
- Bordetella pertussis*, 145e-2t, 1021–1022. *See also* Pertussis
- Bornholm disease (pleurodynia), 1291
- Boron, 96e-9t
- Borrelia* spp., 150e-1, 150e-8, 1146, 1146t. *See also* Relapsing fever
- Borrelia burgdorferi*, 145e-3, 1149–1150, 1561.
See also Lyme borreliosis (Lyme disease)
- BOR (branchio-oto-renal) syndrome, 222t
- Bortezomib
 action and targets of, 102e-5t, 102e-6, 103e-18t, 103e-19
 adverse effects of, 103e-18t, 103e-19, 607, 607t, 716–717, 2686t
 for amyloidosis, 723
 for kidney transplant rejection, 1829
 for mantle cell lymphoma, 704
 for multiple myeloma, 102e-5t, 102e-6, 103e-19, 716–717
 for Waldenström's macroglobulinemia, 718
- Bosentan
 for high-altitude pulmonary edema, 476e-3
 for ischemic ulcers, 2164
 for pulmonary arterial hypertension, 1513, 1659–1660, 1660t, 2164
- Boston exanthem, 1292
- Bosutinib
 action and targets of, 102e-3, 102e-5t, 103e-17t
 adverse effects of, 103e-17t, 692
 for CML, 691–692, 691t
 resistance to, 103e-16
- Botflies, 2746–2747
- Botryomycosis, 1087, 1090
- Botswana, 13e-6–7
- Botulinum antitoxin, 989
- Botulinum toxin
 for achalasia, 1904
 for anal fissure, 1978
 as bioterrorism agent, 261e-8, 988. *See also* Botulism
 botulism and, 988
 for dystonia, 2621
 for essential tremor, 2620
 for gastroparesis, 261
 for hemifacial spasm, 2648
 for hyperhidrosis, 2643
 for ischemic ulcers, 2164
 for migraine prevention, 2594t
 for tics, 2623
- Botulinum toxoid, 261e-9
- Botulism, **987**
 antitoxin, 989
 in bioterrorism, 261e-8
 clinical features of, 261e-5t, **988**
 neurologic, 261e-9, 1784, 2642, 2703
 diagnosis of, 261e-5t, 988–989
 differential diagnosis of, 989
 diplopia in, 208
 epidemiology of, 987–988
 etiology of, 987
 food-borne, 987–988
 global considerations in, 987
 iatrogenic, 988
 infant, 987, 988, 1784
 inhalational, 988
 intentional, 988
 intestinal, 987, 988
 notification of, 989
 pathogenesis of, 987, 2703
 prevention of, 261e-5t, 261e-9, 989
 treatment of, 261e-5t, 261e-9, 989, 2703
 wound, 987, 988, 1784
- Bouba. *See* Yaws
- Bouchard's nodes, 2226f
- Bound morphemes, 178
- Bourneville's disease. *See* Tuberous sclerosis (Bourneville's disease)
- Boutonneuse fever (Mediterranean spotted fever), 128t, 782, 1155t, 1157
- Boutonnière deformity, 2136
- Bovine papular stomatitis, 220e-1t
- Bovine spongiform encephalopathy (BSE), 453e-1t, 453e-2
- Bowel obstruction. *See* Intestinal obstruction
- Bowen's disease, 351, 353t, 501
- Bowlegs, 2221, 2228, 2228f
- Bowman's capsule, 332e-1
- Box jellyfish envenomation, 2739, 2741–2742
- Bpi (bactericidal/permeability-increasing protein), 372e-24t
- Brachial neuritis, 122
- Brachial plexopathy
 imaging in, 441e-30–31f
 immune-mediated, 2692
 neoplastic, 618, 2692–2693
- Brachial plexus, 122, 2693, 2694f
- Brachytherapy
 for cholangiocarcinoma, 553
 definition of, 103e-4
 for prostate cancer, 584
 for soft tissue sarcoma, 119e-2
- Bradyarrhythmias
 with asystole, 1769f
 in AV conduction block. *See* Atrioventricular (AV) conduction block
 etiology of, 763t, 1466, 1470
 junctional bradycardia, 278e-3f
 mechanism of, 1466, 1466f, 1467f
 pacemaker for, 273e-7
 relative bradycardia, 126
 in SA node dysfunction. *See* Sinoatrial (SA) node disease/dysfunction
 syncope in. *See* Cardiac syncope
- Bradykinesia, 154, 2609
- Bradykinin, 88f
- Bradyrhizobium enterica*, 772
- BRAF gene mutations
 in colorectal cancer, 101e-1f
 as drug target, 32, 101e-8, 102e-6, 103e-19
 in hairy cell leukemia, 135e-2, 706
 in lung cancer, 507t, 509, 509f, 521
 in melanoma, 496, 499
 in thyroid cancer, 2304, 2305t
 tumors associated with, 101e-2t
- BRAF inhibitors
 action of, 102e-3, 103e-19
 adverse effects of, 103e-17t, 380
 for melanoma, 499, 501
- Brain
 atrophy of, 75–76, 76f
 cerebral hemispheres, 176f, 2572f, 2573f, 2574f
 coronal section of, 2572f
 gut microbiota and, 86e-7–8
 inferior aspect of, 2575f
 lateral aspect of, 2573f
 medial aspect of, 2574f
 metastases to, 1792
- Brain abscess, **900**
 anaerobic bacterial, 1097–1098
 approach to the patient, 883–884, 884–885f
 in cancer patient, 488, 488t
 clinical features of, 783, 901
 definition of, 900
 diagnosis of, 440e-5f, 901, 902f
 differential diagnosis of, 902
 epidemiology of, 900
 etiology of, 780t, 900–901, 1097–1098
 global considerations in, 900
 histopathology of, 901
 nodocardial, 1085, 1085f, 1087
 pathogenesis of, 901
 prognosis of, 902
 treatment of, 780t, 902
- Brain biopsy, 175, 386e-6f, 895
- Brain death, 442e-2, 1735, 1776
- Brain-derived neurotrophic factor (BDNF), 444e-7t
- Brain drain, 13–14
- Brain edema, 1777
- Brainerd diarrhea, 267
- BRAIN (Brain Research through Advancing Innovative Neurotechnologies) Initiative, 444e-12–13
- Brain injury
 post-cardiac bypass, 463e-2–3, 463e-2f
 post-solid organ transplantation, 463e-3
 stem cell therapy in, 90e-3
 traumatic. *See* Head injury

- Brain natriuretic peptide. *See* B-type natriuretic peptide (BNP)
- Brainstem accessory optic system, 195
- Brainstem auditory evoked potentials (BAEPs), 223, 442e-3
- Brainstem disorders/lesions
clinical features of, 2540t
ischemia, 2577-2580f, 2578
sensory abnormalities in, 161
weakness in, 156
- Brainstem reflexes, 1774-1775, 1774f
- Brain tumors, **598**
amnesia in, 183
approach to the patient, 598
clinical features of, 598, 598t
germ cell tumors, 2264
headache in, 107t, 108, 598
imaging in, 440e-9f, 598
intracranial hemorrhage in, 2583t, 2584
metastatic, **604**
diagnosis of, 604-605, 605f
epidural. *See* Spinal cord compression, neoplastic
increased intracranial pressure and, 1792
leptomeningeal, 605, 606f
sources of, 604, 604t
treatment of, 605
- in neurofibromatosis type 1, 604-605. *See also* Neurofibromatosis type 1 (NF1)
- in neurofibromatosis type 2, 605. *See also* Neurofibromatosis type 2 (NF2)
- primary, **599**
extrinsic "benign," **602**
colloid cysts, 603
craniopharyngiomas. *See* Craniopharyngiomas
dermoid cysts, 603
dysembryoplastic neuroepithelial tumors, 603
epidermoid cysts, 603
meningiomas. *See* Meningiomas
pituitary tumors. *See* Pituitary tumors (adenomas)
schwannomas, 602-603. *See also* Vestibular schwannoma (acoustic neuroma)
genetic syndromes associated with, 599t
intrinsic malignant
astrocytomas. *See* Astrocytomas
brainstem gliomas, 441e-39f, 601
ependymoma, 441e-39f, 601
gangliomas, 601
gliosarcomas, 601
medulloblastomas, 602
oligodendroglioma, 600-601
pineal region tumors, 602
pleomorphic xanthoastrocytomas, 601
primary central nervous system lymphoma. *See* Primary central nervous system lymphoma (PCNSL)
pathogenesis of, 599
treatment of, 598-599
- in tuberous sclerosis, 604. *See also* Tuberous sclerosis (Bourneville's disease)
- Branched-chain DNA (bdNA) tests, 150e-7-8, 1247, 1247t
- Branching enzyme deficiency (type IV glycogenosis), 462e-14
- Branchio-oto-renal (BOR) syndrome, 222t, 1855
- Branch retinal vein occlusion, 201
- Brazil, 13e-6, 13e-6f
- BRC (benign recurrent cholestasis), 284
- BRCA gene mutations
in breast cancer, 523-524
cancers associated with, 101e-6t
early interventions in patients with, 450t
epigenetic silencing of, 102e-7
genetic testing for, 101e-6, 450t, 480, 523
in ovarian cancer, 592
in pancreatic cancer, 554
synthetic lethality and, 102e-7
variable expression of, 446
- Break-bone fever. *See* Dengue fever
- Breakpoint method, 150e-9
- Breast cancer, **523**
biology of, 526
carcinoma of unknown primary and, 120e-1, 120e-2t, 120e-3
in contralateral breast, 531
deaths from
in 2014, 467t, 523
by age group, 6e-2f
by racial and ethnic group, 471t
trends in, 469f, 470t
diagnosis of, 524-526, 525f, 526f
epidemiology of, 524
epigenetic silencing in, 102e-7
estrogen and progesterone receptor status of, 526, 528
follow-up, 532, 532t
gene expression profiles in, 448, 526
genetic considerations in, 101e-2t, 101e-3, 101e-6t, 446, 447f, 523-524
global considerations in, 524
hereditary, 447, 450t. *See also* BRCA gene mutations
histologic classification of, 528
incidence of, 467t, 468, 468f, 471t, 523
infections in, 484t
lifetime risk of, 6e-1, 27t, 524
male, 523, 532
metastatic
to bone, 119e-4
bone marrow examination in, 81e-7f
to CNS, 604, 604t, 606f, 2653, 2653f
diagnosis of, 529
to eye, 206
to heart, 289e-2
to liver, 553
lymphadenopathy in, 408
prognosis of, 529
timing of, 529
treatment of, 529-531
noninvasive, 531-532
oral contraceptives and, 2391t
ovarian cancer and, 592
paraneoplastic syndromes in
endocrinologic, 609, 609t, 614t
hematologic, 612, 612t
neurologic, 618
as perceived health problem, 6e-2f
pleural effusion in, 474
during pregnancy or lactation, 124e-1t, 124e-3-4, 124e-3t, 525-526
- prevention of
lifestyle modifications for, 476
pharmacologic, 478, 531
prophylactic surgery for, 103e-3, 450t, 479
tamoxifen for, 478
- prognosis of, 528, 528t
radiation-induced, 103e-4, 524, 623
recurrent, 103e-16, 528
- risk factors for
alcohol use, 524, 2725
diet, 476, 524
hormone use, 524, 2383, 2384f
obesity, 95e-5, 524
- screening for
benefits of, 27, 28t, 480, 526
in elderly, 84
recommendations for high-risk women, 480, 526
recommendations for normal-risk women, 29t, 480, 481t, 526
- second malignancies in survivors of, 623
staging of, 526, 527t, 528
subtypes of, 526
treatment of, **526**
ado-trastuzumab emtansine, 102e-4, 102e-5t, 103e-22
biologic therapy, 103e-21, 103e-21t
chemotherapy, 528-531, 529t
hormone receptor-directed, 102e-7, 103e-15-16, 530, 530t
HSCT, 139e-6, 531
late consequences of, 622, 624t
lymphedema after, 1654
- in metastatic disease, 529-530
in postmenopausal women, 528-529, 529t
in premenopausal women, 528, 529t
radiation therapy, 527
sentinel lymph node biopsy, 528
in stage III disease, 531
surgical, 526-528
tamoxifen, 102e-7
targeted chemotherapy, 102e-5t, 102e-6, 102e-16, 103e-19
trastuzumab, 102e-4, 102e-5t, 102e-6, 102e-10, 103e-21, 103e-21t
tumor growth rate measures in, 528
tumor markers in, 473t, 526
- Breast development, 2379, 2379t
- Breast-feeding
AED use during, 2559
antibacterial therapy during, 936, 937t
HIV transmission by, 50, 1223
during SLE therapy, 2133
water requirement during, 95e-1
- Breast mass
benign, 526
cyst, 525f, 526
evaluation of, 524-526, 525f, 526f
palpable, 524-525, 525f
in pregnant or lactating woman, 124e-3, 525-526
- Breast milk jaundice, 2001
- Breast self-examination, 480, 481t, 526
- Breath sounds, 1662
- Brenner tumor, 592
- Brentuximab vedotin, 102e-4, 102e-5t, 103e-22
- Breslow thickness, 496
- BRIC (benign recurrent intrahepatic cholestasis), 284, 2003t, 2004, 2076
- Brill-Zinsser disease, 152e-5t, 1154, 1158
- Brim sign, 426e-2
- Brincidofovir (CMX-001), 215e-7, 261e-6
- Bristleworm sting, 2739, 2739f
- Brittle asthma, 1680
- Broadbent's sign, 1576
- Broad casts, urinary, 294
- Broca's aphasia, 177t, 178
- Broca's area, 177, 180f
- Brodalumab, 2177
- Brodie-Trendelenburg test, 1651f
- Bromides, 366, 2764t
- Bromocriptine
for acromegaly, 2271
adverse effects of, 1650, 2268
overdosage/poisoning with, 473e-8t
for prolactinoma, 2268
for prolactin-secreting tumors, 103e-16
for type 2 diabetes mellitus, 2415
- Bronchial artery embolization, 247
- Bronchial aspirate, 150e-4t
- Bronchial hygiene, 1696
- Bronchial neuroendocrine tumors (carcinoids), 269, 560t, 563-564
- Bronchiectasis, **1694**
approach to the patient, 1695
breath sounds in, 1662
chest radiograph in, 308e-11f
clinical features of, 1695
clubbing/hypertrophic osteoarthropathy in, 250
complications of, 1696
CT in, 308e-3f, 308e-12f
definition of, 1694
diagnosis of, 1694t, 1695, 1695f
epidemiology of, 1695
etiology of, 1694-1695, 1695t
global considerations in, 1695
hemoptysis in, 245
imaging in, 308e-11-12f, 1695, 1695f
lung transplantation for, 320e-1t
nontuberculous mycobacterial, 205e-8, 1129, 1130, 1694, 1696
pathogenesis of, 1695
prevention of, 1696
prognosis of, 1696
traction, 1695
treatment of, 1696

- Bronchiolitis
 HMPV, 1206
 HRSV, 1205
 interstitial lung disease associated with, 1713
 parainfluenza virus, 1207
 Bronchiolitis obliterans, 320e-4, 372
 Bronchitis
 chronic, 244, 1692t, 1701. *See also* Chronic obstructive pulmonary disease (COPD)
 hemoptysis in, 245
 pathophysiology of, 306e-3
 Bronchoalveolar lavage
 in interstitial lung diseases, 1710, 1711t
 procedure, 1667
 in sarcoidosis, 2210, 2210f
 Bronchocentric granulomatosis, 1716
 Bronchodilators. *See* β -adrenergic agonists
 Bronchoscopy
 autofluorescence, 1667
 flexible fiberoptic, 1667
 in hypersensitivity pneumonitis, 1683
 in lung cancer diagnosis, 512
 in respiratory disease evaluation, 1663, 1667
 therapeutic, 1667
 virtual, 1666, 1666f
 Bronzing, in hemochromatosis, 2516
 Brown adipose tissue, 415e-2
 Brown recluse spider bite, 2747
 Brown-Séquard syndrome, 156, 161, 2652
 Brown tumor, 1815
Brucella spp., 194e-1, 194e-2
Brucella spp. infections, 194e-1
 chronic meningitis, 907t
 clinical features of, 152e-3t, 194e-2-3
 diagnosis of, 194e-3-4, 194e-3f
 epidemiology of, 194e-1
 global considerations in, 194e-1
 immune response to, 194e-1-2
 incubation period for, 152e-3t, 194e-2
 monocytosis in, 421
 osteomyelitis, 839
 pathogenesis of, 194e-1-2
 prevention of, 152e-3t, 194e-4-5
 prognosis of, 194e-4
 in returning war veterans, 152e-3t, 152e-8
 skin manifestations of, 366
 treatment of, 152e-3t, 194e-4
 vs. tuberculosis, 194e-2, 194e-2t
 Brudzinski's sign, 883
 Brugada's sign, 310
 Brugada syndrome
 clinical features of, 147, 1497
 ECG in, 273e-5, 1493f, 1497
 genetic considerations in, 147
 treatment of, 1497-1498
 ventricular arrhythmias in, 1495t
Brugia malayi, 245e-3t, 1417, 1418t. *See also* Lymphatic filariasis
Brugia timori, 1417, 1418t. *See also* Lymphatic filariasis
 Brunner's gland adenoma, 536
 Bruns-Garland syndrome, 2682-2683
 Brush border, 332e-6
 Bruxism, 192, 237
 Bryostatins, 102e-4f
 BSE (bovine spongiform encephalopathy), 453e-1t, 453e-2
 BTK gene, 2111
 B-type natriuretic peptide (BNP)
 in amyloidosis, 720
 in chest pain evaluation, 101
 in edema, 251
 in heart failure, 101, 287, 1505
 in pulmonary edema, 1763
 in pulmonary hypertension, 1657, 2161
 reference values for, 2758t
 in shock, 1746
 Bubble study, 1655
 Bubo, 261e-4, 829, 1072, 1072f
 Buccofacial apraxia, 179
 Bucindolol, 40
 Budd-Chiari syndrome (hepatic vein thrombosis)
 abdominal pain in, 106t
 ascites in, 287
 in hepatocellular carcinoma, 545, 2027
 liver transplantation for, 2068
 in paroxysmal nocturnal hemoglobinuria, 660
 in polycythemia, 399, 673
 recurrence of, after liver transplantation, 2074
 splenomegaly in, 411
 Budesonide, 270, 1960
 Buerger's disease (thromboangiitis obliterans), 1645-1646, 1649
 Bufadienolide, 310
 Buffalo hump, 1259, 1260f
 Buffalopox, 220e-1t, 220e-2
Bufo marinus (cane toad), 310
 Bufonidae spp., 473e-9t
 "Building-associated illness," 1663
 Bulbospinal pathway, 155f
 "Bulge sign," 2219
 Bulimia nervosa
 clinical features of, 2720
 hypokalemia in, 307
 nausea and vomiting in, 259
 oral manifestations of, 236
 treatment of, 2720
 Bullae. *See also* Vesicles/bullae
 characteristics of, 127, 827f
 description of, 339t, 340f
 etiology of, 828t, 829
 Bullet ant, 2749
 Bullous dermatosis of hemodialysis, 361t, 362
 Bullous impetigo, 349, 967. *See also* Impetigo
 Bullous pemphigoid. *See* Pemphigoid, bullous
 Bumetanide, 322, 1510t, 1763
 BUN. *See* Blood urea nitrogen (BUN)
 Bundobugo virus disease, 1325f, 1326f, 1327t
 Bundle branch reentry, 273e-4, 1495
 Bundle of His. *See* His bundle
 Bunyamwera virus, 1308t
 Bunyaviruses
 gene expression and replication in, 214e-4
 geographic distribution of, 1311t
 global considerations in, 782
 reservoirs and vectors of, 1305, 1306t
 structure of, 214e-2t, 214e-3f
 Bunyavirus infections. *See also* Hantavirus infections
 California meningoencephalitis, 1314
 clinical syndromes of, 1306t, 1312t, 1318
 Crimean-Congo hemorrhagic fever, 1321
 hemorrhagic fever with renal syndrome, 1321
 Orthobunyavirus group C serogroup, 1318
 Orthobunyavirus Simbu serogroup, 1318
 Rift valley fever. *See* Rift Valley fever
 sandfly fever, 152e-5t, 1318
 severe fever with thrombocytopenia syndrome, 1322
 Buprenorphine, 91t, 94, 468e-3-4
 Bupropion
 adverse effects of, 64, 507, 2710t, 2716
 for depression, 64, 2710t, 2716
 with naltrexone, for obesity, 2397
 for smoking cessation therapy, 507, 1705, 2732
 for SSRI-related sexual dysfunction, 2716
Burkholderia cepacia infections, 487, 1045t, 1048, 1697
Burkholderia mallei, 1049
Burkholderia pseudomallei, 152e-4t, 1045t, 1048
 Burkitt's lymphoma, 706
 clinical features of, 702t, 706
 diagnosis of, 81e-9f, 81e-11f, 706, 706f
 EBV infection and, 101e-7, 477t, 706
 endemic, 706
 genetic considerations in, 101e-3, 101e-3t, 698, 699t, 706
 in HIV infection, 706, 1270
 immunology of, 698, 698f
 malaria and, 1375
 oral manifestations of, 236
 during pregnancy, 124e-4
 sporadic, 706
 treatment of, 706
 tumor lysis syndrome in, 1795
 Burnett's syndrome, 2479
 Burning mouth syndrome (glossodynia; glossalgia), 215, 239
 Burnout syndrome, 475
 Burn patient
 acute kidney injury in, 1803
 complications in, 166e-1. *See also* Burn-wound infections
 hypocalcemia in, 315
 hypothermia in, 478e-1
 hypovolemia in, 298
 respiratory acidosis in, 323
 Burn-wound infections, 166e-1
 classification of, 166e-2
 clinical features of, 166e-1-2
 diagnosis of, 166e-2
 epidemiology of, 166e-1
 immunosuppression caused by, 166e-2
P. aeruginosa, 166e-1, 166e-2f, 1043
 pathophysiology of, 166e-1
 treatment of, 166e-3
 Burow's solution, 228
 Burrowing asps, 2733. *See also* Snakebites
 Bursitis, 2247
 Achilles, 2247
 anserine, 2222, 2248
 diagnosis of, 2247
 iliopsoas, 2247
 ischial, 2247
 olecranon, 2247
 pathophysiology of, 2247
 prepatellar, 2248
 retrocalcaneal, 2247
 subacromial, 2221, 2222
 treatment of, 2247
 trochanteric, 2222, 2222f, 2247
 Buschke-Ollendorff syndrome (dermatofibrosis lenticularis disseminata), 426e-5
 Buspirone
 for anxiety disorders, 2711, 2711t
 for cocaine abuse/dependence, 469e-2
 drug interactions of, 246e-3t
 for functional dyspepsia, 263
 for SSRI-related sexual dysfunction, 2716
 Busulfan
 adverse effects of, 694
 hepatic, 2025
 hyperpigmentation, 361, 379
 nausea and vomiting, 103e-25
 neurologic, 607
 pulmonary, 1797
 for CML, 694
 drug interactions of, 246e-3t
 genetic variations in response to, 39t
 in HSCT preparation, 139e-2
 Butabarbital, 473e-10t
 Butorphanol
 action of, 94
 for migraine, 2592t, 2593
 for pain, 91t, 92
 Butyrate, for stimulation of fetal hemoglobin, 639
 Bwamba virus, 1308t
 Byssinosis, 1688t, 1691
 Bystander killing, 1244
 C
 C1 inhibitor (C1INH) deficiency, 105, 372e-25t, 2118, 2119, 2120
 C1q, 372e-25t
 C3 glomerulopathy, 62e-4f
 C9ORF72 gene mutations, 2603
 Ca²⁺/calmodulin pathway, 1106
 CA 19-9, 473t, 553, 555
 CA-125
 in cholangiocarcinoma, 552
 in ovarian cancer, 482, 593
 reference values for, 2758t
 as tumor marker, 473t
 Cabazitaxel, 103e-15, 586
 Cabergoline, 565, 2267-2268, 2271
 CABG. *See* Coronary artery bypass grafting (CABG)

- c-abl* gene mutations, 83e-7
- Cabozantinib
 action and targets of, 102e-5t, 103e-18t
 adverse effects of, 103e-18t
 for lung cancer, 521
 for medullary thyroid carcinoma, 234z
- Cachectic myopathy, 1784
- Cache valley virus, 1308t
- Cachexia (chronic disease-related malnutrition), 459-460, 1504
- CACNA1 gene mutations, 465e-2, 2589, 2629
- CAD. *See* Coronary artery disease (CAD)
- CADASIL (cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy), 441e-18f, 2568, 2608
- Cadherins, 102e-11f
- Cadmium exposure/poisoning, 472e-1, 472e-2t, 1692t, 1856t, 1862, 2766t
- Ca-DTPA, 263e-6t, 263e-7
- Caenorhabditis elegans*, 255e-1
- Cafe-au-lait macules/spots, 359, 360t, 426e-6
- Caffeine
 action of, 186
 adverse effects of, 65, 194, 254, 305, 1861
 for hypnic headache, 2598
 for low CSF volume headache, 110
 for migraine, 2592t
 overdose/poisoning with, 473e-8t, 2764t
 for post-lumbar puncture headache, 443e-3
 for shift-work disorder, 194
 therapeutic range of, 2764t
- CagA protein, 1039
- CAGE questionnaire, 1992, 1992t
- CAH. *See* Congenital adrenal hyperplasia (CAH)
- Calabar swellings, 1421
- Calcification
 of coronary arteries, 270e-4, 270e-7f, 270e-11f, 271e-2f, 1585
 dystrophic, 363, 426e-7t, 426e-8
 extraskelatal, 426e-7-8, 426e-7t
 metastatic, 426e-7, 426e-7t
 of solitary pulmonary nodule, 515
- Calcific emboli, 40e-4
- Calcific tendinitis, 2248
- Calcinosis
 in calcium apatite deposition disease, 2236
 in dermatomyositis, 2202
 treatment of, 2237
 tumoral, 426e-7t, 426e-8
- Calcinosis circumscripta, 426e-8
- Calcinosis cutis
 clinical features of, 363, 364t, 2160f
 metastatic, 364t
 in systemic sclerosis, 376, 2160, 2160f
- Calcinosis universalis, 426e-8
- Calciphylaxis, 363, 1816, 1816f
- Calcipotriene, 348
- Calcitonin, **2469**
 actions of, 2469, 2501
 adverse effects of, 426e-4
 for bone pain, 61
 circulating level of, 2469
 ectopic production of, 609t
 for hypercalcemia, 610, 2480t, 2481
 hypocalcemic activity of, 2469
 in medullary thyroid carcinoma, 2341
 for osteoporosis management/prevention, 2501
 for Paget's disease of bone, 426e-3t, 426e-4
 pNET secreting, 559t
 reference values for, 2758t
 sources of, 2469
 as tumor marker, 473t
- Calcitonin gene(s), 2469
- Calcitonin gene-related peptide (CGRP), 88, 444e-3, 2469
- Calcitriol, 315, 1816, 2466, 2484, 2488
- Calcium
 absorption of, 2457
 in cardiac contraction, 265e-5f, 265e-6-7, 1563
 in chronic kidney disease, 333e-3, 1816-1817, 1816f
 deficiency of. *See* Hypocalcemia
 excess of. *See* Hypercalcemia
 extracellular, 313, 313f
 homeostasis of, 2456-2457, 2457f, 2467
 intake of, 2457, 2497t
 malabsorption of, postgastroectomy, 1927
 metabolism of, 2208, 2456-2458, 2457f
 nephrolithiasis risk and intake of, 1867, 1870
 osteoporosis risk and intake of, 2491
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-3t, 2496-2497, 2496t
 reference values for, 2758t
 renal transport of, 332e-4f, 332e-8
 supplements. *See* Calcium supplements
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t
 urinary excretion of, 1867
 in vascular smooth-muscle cell function, 265e-3-4, 265e-3f
- Calcium apatite deposition disease, 2236-2237, 2236t, 2237f
- Calcium carbonate, 1920
- Calcium channel(s)
 cardiac, 273e-2, 273e-2f
 muscle
 characteristics of, 265e-3-4, 265e-3f, 462e-17f
 disorders of, 462e-16-17, 462e-17f, 462e-17t
 in seizure initiation, 2547
- Calcium channel blockers (CCBs)
 adverse effects of, 326-327, 1625
 AV conduction block, 1471t
 edema, 253t, 1443, 1651
 erythromelalgia, 1650
 gingival hyperplasia, 46e-1f, 237
 heat-related illness, 479e-1
 lupus syndrome, 2218t
 peripheral edema, 2392
 SA node dysfunction, 1467t, 1469
 secondary parkinsonism, 2611
 urinary incontinence, 81
 xerostomia, 241
- aldosterone-renin ratio effects of, 2320t
 for aortic dissection, 1641
 drug interactions of, 42t, 2711t
 for focal atrial tachycardia, 1479
 for heart failure, 1513
 for hypertension, 1624t, 1625, 1821
 for hypertrophic cardiomyopathy, 1569
 for inappropriate sinus tachycardia, 1477
 for ischemic heart disease, 1589-1590, 1589t
 metabolism of, 34t, 42
 for NSTEMI-ACS, 1596, 1597t
 overdose/poisoning with, 473e-9t
 perioperative prophylactic, 54
 for Prinzmetal's variant angina, 1589, 1598
 for Raynaud's phenomenon, 1649, 2165
 for supraventricular tachycardia in pregnancy, 47
 for ventricular tachycardia, 1498
- Calcium docusate, 62t
- Calcium gluconate, 312, 315, 2739
- Calcium oxalate crystals/stones
 in ethylene glycol poisoning, 64e-9, 1858-1859
 renal, 62e-10f, 1866, 1868f, 1870. *See also* Nephrolithiasis
- Calcium oxalate deposition disease, 2237, 2237f
- Calcium phosphate product, elevated, 363
- Calcium phosphate stones, 1866, 1870. *See also* Nephrolithiasis
- Calcium polycarboxophil, for irritable bowel syndrome, 1970t
- Calcium pyrophosphate deposition (CPPD) disease, 372e-8t, 2235-2236, 2235t, 2236f
- Calcium-sensing receptors, 313, 372e-24t
- Calcium signaling defects, 2109
- Calcium sparks, 265e-3
- Calcium supplements
 absorption of, 2457
 for colorectal cancer prevention, 477-478, 539
 in elderly, 84
 for hyperkalemia, 312
 for hypocalcemia, 315
 for hypoparathyroidism, 2484
 for osteoporosis management/prevention, 2497
 for vitamin D deficiency, 2466
- Calcofluor white, 150e-1
- Calculation ability, assessment of, 2537
- Calculus, 46e-4f, 46e-5f, 236
- Caliciviruses. *See also* Norovirus (Norwalk virus) infections
 characteristics of, 1285
 gene expression and replication in, 214e-4
 structure of, 214e-2t, 214e-3f, 1286f, 1286t
- California encephalitis virus, 1309t, 1314
- California meningoencephalitis, 1314
- Californium exposure, 263e-7
- Calor, 414f, 1753
- Caloric irrigation, 209, 1775
- Caloric needs, 95e-1, 462
- Caloric restriction, aging effects and, 85, 94e-4-6
- Calorimetry, indirect, 462
- Calretinin, 120e-2t, 120e-4, 508
- CALR gene mutations, 675
- Calsequestrin, 265e-7
- Calstabin 2, 265e-6
- Calymmatobacter granulosus*, 198e-1. *See also* Donovanosis
- CAMP factor, 969
- cAMP response-element binding protein. *See* CREB (cAMP response-element binding protein)
- Camptothecin, 103e-13
- Campylobacter* spp., 1058-1059
- Campylobacter* spp. infections, **1058**
 clinical features of, 266t, 856t, 1059t, 1060
 colitis, vs. IBD, 1956
 complications of, 1060
 diagnosis of, 150e-2t, 150e-6, 856, 1060
 differential diagnosis of, 1060-1061
 epidemiology of, 855t, 1059
 food-borne, 266, 855t, 856t, 857, 1059
 Guillain-Barré syndrome and, 2695, 2696f
 in HIV infection, 1257
 immunoproliferative small intestinal disease, 719
 pathogenesis of, 1059-1060
 prognosis of, 1061
 treatment of, 857, 1061
- Camurati-Engelmann disease (progressive diaphyseal dysplasia), 426e-5
- Canadian Cardiovascular Society, classification of cardiovascular disease, 1581t
- Canagliflozin, 2414t
- Canakinumab, 2214, 2234
- Canasa suppositories, 1960
- c-ANCA (cytoplasmic antineutrophil cytoplasmic antibody), 377e-4, 2180
- Cancer
 anemia in, 630
 approach to the patient, **467**
 chromosome abnormalities in, 83e-7, 471. *See also* Cancer genetics
 chronic meningitis in, 909t
 clonal origin of, 101e-1, 101e-1f
 death and dying issues, 475
 deaths from
 by age, 470t
 by race and ethnicity, 16e-1f, 470f, 471t
 by sex, 6e-1t, 6e-2f, 467t, 469f, 470t
 by site, 467t, 469f
 trends in, 467, 469f
 in United States and Britain, 56t
 worldwide, 472f
 definition of, 103e-1
 depression in, 2714
 diagnosis of, 103e-1-2, 103e-1t, 468
 effusions in, 474
 emergencies in
 adrenal insufficiency, 1795
 airway obstruction, 1794, 1794f
 biliary obstruction, 1790
 hemolytic-uremic syndrome, 1796-1797

- hemoptysis, 1793–1794
 hemorrhagic cystitis, 1798
 human antibody infusion reactions, 1796
 hypersensitivity reactions to antineoplastic drugs, 1798
 hypocalcemia. *See* Hypocalcemia
 hypoglycemia, 1794–1795
 increased intracranial pressure, 1792
 intestinal obstruction, 1789–1790
 lactic acidosis, 1794
 neoplastic meningitis, 1792–1793
 neutropenic enterocolitis, 488, 1797–1798, 1798f
 pericardial effusion/tamponade, 474, 1788–1789
 pneumonia, 488–489, 489t, 1797
 pneumonitis, 489, 489t, 621–622, 1797
 pulmonary and intracerebral leukostasis, 1793
 pulmonary infiltrates, 1797
 seizures, 1793
 SIADH. *See* Syndrome of inappropriate antidiuretic hormone (SIADH)
 spinal cord compression. *See* Spinal cord compression, neoplastic
 superior vena cava syndrome. *See* Superior vena cava syndrome
 tumor lysis syndrome. *See* Tumor lysis syndrome
 urinary obstruction, 1790
 extent of disease, 470–471
 fatigue in, 152
 fever of unknown origin in, 136t, 137t, 138
 genetic considerations in. *See* Cancer genetics
 genetic testing in, 101e-6–7, 101e-8f, 450t
 global burden of, 16
 global considerations in, 467–468, 490
 immunization in, 485t, 492
 incidence of, 467, 467t, 468f, 471t
 infections in, **484**
 antibacterial therapy for, 490–491
 antibiotic prophylaxis for, 492
 antifungal therapy for, 491
 antiviral therapy for, 491–492
 cardiovascular, 489
 catheter-related, 487, 487t
 central nervous system, 488, 488t
 cytokine therapy, 492
 endocrine, 489
 febrile neutropenia and, 103e-23, 490, 490f
 gastrointestinal tract, 487–488, 1797–1798
 granulocyte transfusions for, 492
 musculoskeletal, 489–490
 predisposing factors in, 484–485, 484t, 490
 prevention of, 492
 pulmonary, 488–489, 489t, 1797
 renal-ureteral, 490
 skin, 76e-12f, 485–487, 486f
 infectious causes of, 761
 in inflammatory myopathies, 2196
 involuntary weight loss in, 275
 late complications of, 473, **620**, 620t, 624t
 in low- and middle-income countries, 16, 16f
 lymphatic vessels of, 102e-15
 metastasis of, 102e-1t. *See also* Metastatic disease
 new symptoms in, 472
 nutritional status of patient, 474
 obesity and, 415e-6
 as organ that ignores its niche, 102e-2–3
 pain in, 473–474
 pathogenesis of. *See* Cancer cells
 patient management in, **468**
 physiologic reserve of patient, 470–471, 472t
 during pregnancy, **124e-1**
 prevention of. *See* Cancer prevention
 prognosis of, 470–471
 progressive disease, 472
 recurrent, 473
 risk factors for, 467, 2725, 2730, 2730t
 screening for. *See* Cancer screening
 second malignancy, 103e-4, 623
 by cancer type, 623
 chemotherapy-related, 623, 709
 hormonal therapy-related, 623
 immunosuppressive therapy-related, 623–624
 radiation therapy-related, 623, 709
 in transplant recipient, 928
 self-renewing capacity of, 89e-4
 stable disease, 472
 staging of, 103e-2, 470
 survivorship issues, 475, 620, 622, 624, 624t
 treatment of
 assessing response to, 472–473, 473t
 biologic therapy. *See* Cancer biologic therapy
 cancer stem cells as target of, 102e-11
 chemotherapy. *See* Cancer chemotherapy
 complementary and alternative medicine, 473
 complete response to, 472
 gene therapy, 91e-2f, 91e-3–4, 91e-3t
 genetic considerations in, 101e-8–9
 goal of, 103e-2
 late consequences of, 624t
 long-term follow-up, 473, 624
 palliative, 103e-3, 472
 partial response to, 472
 planning, 471–472
 principles of, **103e-1**
 psychosocial support, 474–475
 radiation therapy. *See* Radiation therapy
 salvage treatment plan, 473
 spectrum of, 103e-1t, 103e-2
 supportive care, **473**
 surgical. *See* Cancer surgery
 tumor growth rates in, 102e-2–3, 103e-2, 103e-3f
 tumor microenvironment in, 102e-13
 viruses and, 101e-7
Cancer biologic therapy, 103e-20
 action and targets of, 103e-5, 103e-5t, 103e-7f
 adverse effects of, 103e-21
 antibody-mediated, 103e-20–21
 antiangiogenic, 102e-15, 102e-16f, 102e-17, 102e-17f, 102e-21t
 conjugates, 102e-4, 102e-5t, 103e-22
 immunoregulatory, 102e-5t, 102e-17–18, 102e-18f, 103e-21–22, 103e-21t, 103e-22f
 tumor-regulatory, 102e-5t, 103e-21, 103e-21t
 cytokines, 103e-22
 immune cell-mediated, 102e-17–18, 103e-20
 ligand receptor-directed constructs, 103e-22
 principles of, 102e-17–18, 103e-20
 T cell-mediated, 103e-20
Cancer cells
 apoptosis in, 102e-1t, 102e-8–10, 102e-9f, 103e-6, 103e-7f
 autophagy in, 102e-10
 biology of, **102e-1**
 cell cycle in, 102e-1–2, 103e-8
 chromatin structure alterations in, 102e-7–8, 102e-8f
 evasion of immune system, 102e-1t
 failure to differentiate, 102e-1t
 gene expression in, 102e-1t, 102e-7–8, 102e-8f
 Gompertzian growth curve/kinetics in, 102e-2, 103e-2, 103e-3f
 induction of differentiation in, 103e-6
 loss of replicative senescence in, 102e-1t, 102e-3
 metabolism in, 102e-12–13, 102e-13f
 mutations in, 101e-8, 102e-1
 necrosis in, 102e-10
 oncogene addiction in, 102e-6
 phenotypic characteristics of, 102e-1t
 plasticity of, 102e-12
 resistance of, 102e-12
 signal transduction pathways in, 102e-3–7, 102e-4f
 synthetic lethality in, 102e-6f
 telomerase activation in, 102e-3
 tissue invasion by, 102e-1t, 102e-10
 in tumor angiogenesis. *See* Angiogenesis, tumor
Cancer chemotherapy, 103e-5. See also specific drugs and cancers
 action of, 103e-6–7, 103e-7f
 adjuvant, 103e-7
 cell/tumor response to, 103e-6–7
 complete response to, 103e-5
 complications of, 472, 489
 cardiovascular, 270e-21–22
 neurologic, 607, 607t
 neuropathies, 2686, 2686t
 curability of cancers with, 103e-6–7, 103e-8t
 cytotoxic
 antimetabolites, 103e-10–11t, 103e-13–14
 antimitotic agents, 103e-11–12t, 103e-14–15
 antitumor antibiotics, 103e-9–10t, 103e-13
 DNA-interactive agents, 103e-8, 103e-9t, 103e-12
 topoisomerase inhibitors, 103e-9–10t, 103e-13
 development of, 103e-5–6, 103e-6f
 dose-limiting toxicity, 103e-5
 gene therapy to enhance, 91e-3–4
 high-dose regimens, 103e-7, 103e-8t
 intraperitoneal, 593
 late complications of, 620t
 cardiovascular, 270e-21–22, 621, 1562–1563
 hematologic, 678
 hepatic, 622
 neurologic, 622
 ocular, 623
 pulmonary, 621–622
 Raynaud's phenomenon, 623
 renal/urinary, 622–623
 reproductive/endocrine, 623, 2365
 maximum-tolerated dose, 103e-5
 metabolism modulators for, 102e-13
 nausea and vomiting in. *See* Nausea and vomiting, chemotherapy-induced
 neoadjuvant, 103e-7
 palliative, 103e-7, 103e-8t
 partial response to, 103e-5
 during pregnancy, 103e-26, 124e-1, 124e-2
 principles for use, 103e-5–8
 resistance to, 102e-12, 103e-23
 response to, 103e-5–6
 supportive care during, **103e-23**
 alopecia, 103e-25
 diarrhea, 103e-25
 gonadal dysfunction, 103e-25–26
 mucositis, 103e-25
 myelosuppression, 103e-23–25, 103e-24t
 nausea and vomiting. *See* Nausea and vomiting, chemotherapy-induced
 targeted, 102e-5t, **103e-15**, 103e-17–19t
 action of, 103e-3–4, 103e-5t, 103e-6, 103e-7f, 103e-16f
 adverse effects of, 103e-25–26
 arsenic trioxide, 103e-19t
 combination, 102e-4, 102e-6
 development of, 103e-6
 diagnostically guided, 101e-8–9, 103e-16, 103e-17t
 DNA methyltransferase inhibitors, 102e-8, 102e-8f, 103e-20
 histone deacetylase inhibitors, 102e-7–8, 102e-8f, 103e-18t, 103e-19–20
 hormone receptor-directed, 102e-7, 103e-5, 103e-15–16
 mTOR inhibitors, 102e-3, 103e-18t
 multikinase inhibitors, 102e-3–4, 102e-4f, 103e-18t, 103e-19
 nondiagnostically guided, 103e-17t
 principles of, 103e-5
 proteasome inhibitors, 102e-6, 102e-9f, 103e-18t, 103e-19
 vismodegib, 102e-5t, 103e-19t
 timing of, 194
Cancer genetics, 101e-1. See also specific cancers
 cancer gene classes, 101e-1–2. *See also* Oncogene(s); Tumor-suppressor genes
 chromosomal instability in solid tumors, 101e-3–4
 clonal origin and multistep nature of cancer, 101e-1, 101e-1f

- Cancer genetics (*Cont.*):
 CpG islands in, 430, 431
 cytogenetic analysis in, 83e-7
 DNA methylation in, 431
 epigenetic control mechanisms, 101e-4
 epigenetic influences on gene transcription, 102e-7-8, 102e-8f
 familial cancer syndromes, 101e-6-7, 101e-6t, 101e-8f
 future of, 101e-9
 gene expression and, 101e-7, 101e-8f
 genomewide mutational profiling, 101e-7-8, 101e-9f
 historical perspective, 101e-1
 microRNAs in, 101e-7
 somatic mutations in, 440
 treatment strategy and, 101e-8-9, 425
 viruses in, 101e-7
- Cancer Genome Atlas, 101e-8
- CancerNet, 471
- Cancer prevention, 17, 475. *See also specific cancers*
 chemoprevention, 476-478
 diet modification, 476
 education and healthful habits, 475-476
 energy balance, 476
 physical activity, 476
 smoking cessation, 475-476
 sun avoidance, 476
 surgical methods, 103e-3, 479
 vaccines and, 478-479
- Cancer screening, 27, 479. *See also specific cancers*
 accuracy of, 479, 479t
 assessment of tests for, 480
 biases of tests for, 479-480
 recommendations for asymptomatic normal-risk subjects, 481t
 risks from, 480
- Cancer stem cells, 102e-11, 102e-12f, 103e-2, 508
- Cancer surgery, 103e-3
 curative, 103e-3
 diagnostic, 103e-3
 late complications of, 620, 620t
 palliative, 103e-3
 preventive, 103e-3, 479
- Cancer syndromes, familial, 101e-6-7, 101e-6t, 101e-8f, 450t
- Cancrum oris (noma), 236, 487, 1097
- Candesartan, 1512t, 1624t, 2594t
- Candida* spp.
 adherence of, 145e-2t, 145e-3
 antifungal susceptibility testing of, 150e-8
 characteristics of, 1342
 epidemiology of, 1342
 laboratory identification of, 150e-8, 1330
 tissue invasion by, 145e-8
- Candida* spp. infections, 1342
 in burn-wounds, 166e-1
 in cancer patient, 486, 486t, 487-488, 489
 catheter-related bloodstream, 98e-7, 915
 in diabetes mellitus, 2429
 diagnosis of, 341, 1343-1344
 disseminated/deeply invasive, 1343f
 brain parenchyma, 441e-8f
 chronic meningitis, 908t
 endocarditis, 817t, 824
 gastrointestinal, 488, 850
 liver abscess, 850
 in newborn, 441e-8f
 ocular, 1343, 1343f, 1345
 osteomyelitis, 839
 renal abscess, 851
 risk factors for, 1342, 1342t
 skin manifestations of, 25e-10f, 135, 1343, 1343f
 treatment of, 1344-1345, 1344t
 urinary tract, 914
 epidemiology of, 1342
- esophageal
 complications of, 1910
 diagnosis of, 1256f, 1896f, 1909f
 in HIV infection, 1256, 1909
 treatment of, 1344t, 1910
- etiology of, 350, 350t
 in HIV infection, 1253t
 in immunocompromised patient, 351
 joint, 836
 mucocutaneous, 350
 chronic, 1343, 2110-2111, 2345, 2346t
 clinical features of, 1342
 appearance, 350t, 351
 color, 351
 distribution, 341t
 intertriginous, 351
 perianal, 1343
 prevention of, 423
 treatment of, 350t, 351, 1344t
- oropharyngeal. *See* Thrush
- pathogenesis of, 1342
- prophylaxis for, 1253t, 1345
 in transplant recipient, 920
- urinary, 868
- vulvovaginal
 clinical features of, 873t, 875, 1342
 diagnosis of, 873t, 875
 in HIV infection, 1259
 in systemic illness, 873
 treatment of, 873t, 875, 1344t
- Candidate genes, 441
- Candidatus* Neoehrlichia mikurensis, 1160
- Candiru virus, 1309t
- CANDLE (chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature), 2215
- Cane toad (*Bufo marinus*), 310
- Cangrelor, 749
- Cannabinoid(s), 260t, 261, 469e-2. *See also* Marijuana
- Cannabinoid hyperemesis syndrome, 259
- Cantagalo, 220e-1t, 220e-2
- Cantharidin, 2750
- CAPD. *See* Continuous ambulatory peritoneal dialysis (CAPD)
- Capecitabine
 action of, 103e-11t
 adverse effects of, 103e-11t, 574
 for breast cancer, 531
 for colorectal cancer, 543
 for gastric cancer, 535
 genetic variations in response to, 39t
 for pancreatic cancer, 557, 557t
 for pNETS, 574
- Capgras' syndrome, 172, 2599
- Capillaria philippinensis*/capillariasis, 245e-3t, 765t, 1417
- Capillaries. *See* Blood vessels
- Capillaritis, 367
- Capillary leak syndrome, 46, 265e-1
- Capillary telangiectasias, 2586
- Capitulation, 15e-2, 21
- Caplan's syndrome, 1690, 2138
- Capnocytophaga* spp. infections
 in cancer patient, 484, 484t
 clinical features of, 183e-2-3, 782
 complications of, 782, 1755
 in dog-bite wound, 167e-1, 183e-3
 etiology of, 782
 treatment of, 167e-3, 167e-3t, 183e-2t, 183e-3
- Capreomycin, 205e-6, 1115
- CAPS (cryopyrin-associated periodic syndromes), 141, 2213t, 2215
- Capsaicin, 2232t, 2682t
- Capsid, viral, 214e-1
- Capsular polysaccharide complex (CPC), 849
- Capsule endoscopy
 in celiac sprue, 1897f
 in Crohn's disease, 1954, 1954f
 in gastrointestinal bleeding, 279
 indications for, 1878, 1879t
 of jejunal vascular ectasia, 1883f
 small-bowel, 1881, 1883f
- Capsulitis, adhesive, 2249
- Captopril
 adverse effects of, 371, 380, 727t
 for heart failure, 1511-1512, 1512t
 for hypertension, 1624t
 for hypertensive emergencies, 1627
 for pulmonary edema, 1761f
- Captopril renography, 1628, 1628t
- Caput medusae, 286, 1992
- CAR (chimeric antigen receptor), 91e-4
- Caraparú virus, 1309t
- Carate. *See* Pinta
- Carbamazepine
 for acroparesthesia, 432e-1
 adverse effects of, 2646
 cholestasis, 2023
 common, 2718t
 cross-sensitivity and, 385
 cutaneous, 353, 361
 fever, 138
 genetic considerations in, 378
 hair loss, 380
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 liver dysfunction, 2682t
 lupus syndrome, 2218t
 nail disorders, 380
 neurologic, 2553t, 2682t
 obesity, 2392
 rare, 2718t
 systemic, 2553t
 thrombocytopenia, 727t
 for bipolar disorder, 2718, 2718t
 for chorea, 2153
 contraindications to, 2552
 drug interactions of, 42t, 205e-3, 246e-3t, 2553t
 for epilepsy, 2552, 2553t
 genetic variations in response to, 38, 39t
 for neuromyotonia, 619
 for neuropathy, 2682t
 overdose/poisoning with, 473e-10t, 473e-13t, 2764t
 for pain, 61, 91t, 94, 2672
 for paroxysmal symptoms in MS, 2672
 pharmacology of, 2553t, 2718t
 for SUNCT/SUNA prevention, 2596
 therapeutic monitoring of, 2764t
 for trigeminal neuralgia, 2646
- Carbamoylphosphate synthase-1 deficiency, 434e-3t
- Carbapenem(s)
 action of, 930, 931t
 for anaerobic bacterial infections, 1101, 1101t
 drug interactions of, 939t
 for gram-negative bacterial infections, 1026-1027
 indications for, 941
 for multidrug-resistant TB, 205e-7
 resistance to, 941
 in *Acinetobacter*, 1037
 in anaerobic bacteria, 1101
 detection of, 769t, 774
 in gram-negative bacteria, 1027
 mechanisms of, 931t, 933
 for staphylococcal infections, 961
- Carbapenemases, 933, 1026, 1027
- Carbenicillin, 306, 322
- Carbenoxolone, 64e-8
- Carbidopa/levodopa, 2614, 2616t
- Carbimazole, 354, 2199, 2296
- Carbinolamine dehydratase deficiency, 434e-2t
- Carbohydrate(s)
 digestion and absorption of, 1937, 1937t
 disorders of metabolism of
 fructose metabolism disorders, 433e-1f, 433e-3t, 433e-5
 galactose metabolism disorders, 433e-1f, 433e-3t, 433e-5
 glycogen storage diseases. *See* Glycogen storage diseases
 high intake of, lipoprotein disorders and, 2438
 malabsorption of, 269
 metabolism of, 433e-1, 433e-1f
 requirement for, 95e-1
 in specialized nutritional support formulas, 98e-5
- Carbon dioxide (CO₂), 151e-1, 151e-2f, 151e-2t

- Carbon disulfide, 2688t
- Carbonic anhydrase, 332e-3f, 332e-6-7
- Carbonic anhydrase II, 372e-24t
- Carbonic anhydrase inhibitors, 64e-3, 205, 246e-6t
- Carbon monoxide poisoning
- clinical features of, 248, 477e-5
 - complications of, 1782
 - epidemiology of, 477e-5
 - hypoxia in, 248
 - lactic acidosis in, 318
 - optic neuropathy in, 203
 - pathophysiology of, 477e-5-6, 639
 - polycythemia in, 400
 - treatment of, 477e-6, 1782
- Carbon tetrachloride, 2023, 2025t
- Carboplatin
- adverse effects of, 103e-9t, 103e-12, 103e-25, 1798, 1804
 - for carcinoma of unknown primary, 120e-4
 - for head and neck cancer, 505
 - for lung cancer, 518t, 520, 521t, 522
 - for melanoma, 499
 - for ovarian cancer, 593
 - as substitute for cisplatin, 622
 - for testicular cancer, 590, 591
 - for thymoma, 123e-2
- Carboxyhemoglobin, 400, 639
- Carboxypeptidase E, 415e-4, 415e-4t
- Carcinoembryonic antigen (CEA), 473t
- in cholangiocarcinoma, 552
 - in colorectal cancer, 76, 542t
 - reference values for, 2758t
- Carcinogen(s), 387, 476
- Carcinogenesis
- field, 477
 - initiation step in, 476
 - Knudson two-hit hypothesis of, 101e-2, 440
 - multistep nature of, 101e-1, 101e-1f, 440, 475
 - promotion, 476
 - radiation in, 103e-4
 - in skin, 387-388
- Carcinoid(s). *See* Gastrointestinal neuroendocrine tumors (GI-NETS)
- Carcinoid crisis, 565
- Carcinoid syndrome, 564
- atypical, 565
 - clinical features of, 96e-4, 269, 559t, 563t, 564-565
 - complications of, 565
 - diagnosis of, 356, 560f, 566
 - diarrhea in, 269, 855t
 - heart disease in, 565, 567, 1568
 - neuroendocrine tumors associated with, 558, 559t, 560f
 - pathobiology of, 559t, 560f, 565
 - pulmonic stenosis/regurgitation in, 1550
 - skin manifestations of, 356, 357, 360
 - treatment of, 272, 566-567, 566f
 - tumor locations in, 559t
- Carcinoma, 102e-1, 103e-1
- Carcinoma in situ, bladder, 575, 576
- Carcinoma of unknown primary, 120e-1, 503
- biology of, 120e-1
 - cytogenetics in, 120e-1
 - genetic considerations in, 120e-1
 - imaging in, 120e-1
 - immunohistochemical analysis in, 120e-2-3, 120e-2t
 - incidence of, 120e-1
 - molecular profiling in, 120e-2-3
 - pathology of, 120e-1-3, 120e-1t
 - prognosis of, 120e-3
 - treatment of, 120e-3-4, 120e-3f, 120e-4f
 - tumor markers in, 120e-1, 120e-2f
- carD* gene, 1106
- CARD15* gene mutations. *See* *NOD2* gene mutations
- Cardiac arrest, 1768
- clinical features of, 1768
 - death after successful resuscitation, 1768
 - definition of, 1764, 1765t
 - etiology of, 1764, 1765t
 - hypothermic, 478e-2
 - in-hospital, 1768, 1770
 - long-term management of survivors of, 1770
 - out-of-hospital, 1768
 - prevention of, 1767
 - primary, 1768
 - prodromal symptoms, 1768
 - prognosis of, 1768
 - progression to biologic death after, 1768
 - secondary, 1768
 - treatment of, 1768
 - advanced cardiac life support, 1769-1770
 - algorithm for, 1769f
 - automated external defibrillation, 1769
 - basic life support, 1768-1769
 - initial response, 1768
 - postresuscitation care, 1770
 - ventricular tachycardia and, 1491
- Cardiac arrhythmias. *See also specific arrhythmias*
- approach to the patient, 273e-4-5
 - bradyarrhythmias. *See* Bradyarrhythmias
 - diagnosis of, 273e-5
 - in heatstroke, 479e-2, 479e-3, 479e-4
 - in hyperkalemia, 310
 - hypertension and, 1615
 - in hypokalemia, 307
 - in hypomagnesemia, 2462
 - in hypothermia, 478e-3
 - in ischemic heart disease, 1580
 - mechanisms of, 273e-2, 273e-3t
 - abnormal impulse conduction, 273e-3t, 273e-4, 273e-4f
 - afterdepolarizations and trigger automaticity, 273e-2-3, 273e-3f, 273e-3t
 - alterations in impulse initiation, 273e-2-3, 273e-3t
 - in myocardial infarction, 1608-1610
 - physical examination in, 273e-5
 - in respiratory alkalosis, 323
 - respiratory sinus, 278e-1f
 - in sarcoidosis, 2208
 - syncope in. *See* Cardiac syncope
 - tachyarrhythmias. *See* Supraventricular tachyarrhythmias; Tachycardia
 - treatment of, 273e-5-7, 273e-6f, 273e-6t
 - warning arrhythmias, 1608
- Cardiac asthma, 1504
- Cardiac auscultation. *See* Heart murmurs; Heart sounds
- Cardiac biomarkers
- in myocardial infarction, 1594, 1596t, 1600-1601, 1601f
 - in VTE, 1633
- Cardiac cachexia, 1504
- Cardiac catheterization, 1460
- in aortic regurgitation, 1536
 - in aortic stenosis, 1531-1532
 - aortography during, 1464
 - in cardiogenic shock, 1760
 - coronary angiography during. *See* Coronary angiography
 - hemodynamic assessment in, 1461
 - cardiac output, 1462-1463
 - in constrictive/restrictive conditions, 1462, 1463t
 - intracardiac shunts, 1463
 - normal waveforms, 1462f
 - reference values for, 1461t
 - valve area, 1463
 - in valvular disease, 1462, 1462f
 - vascular resistance, 1463
 - indications for, 1460, 1460t
 - in mitral stenosis, 1541
 - postprocedure care, 1465
 - preprocedure management of, 1461
 - in pulmonary hypertension, 1657, 2161
 - racial/ethnic disparities in referral for, 16e-1
 - risks of, 1460-1461
 - in STEMI, 1605
 - technique of, 1461
 - vascular access for, 1461
 - ventriculography during, 1463-1464, 1464f
- Cardiac conduction system
- automaticity of, 273e-2-3
 - components of, 1451, 1451f
 - embryonic development of, 265e-12
- Cardiac contraction, 265e-5
- cardiac activation in, 265e-6-7, 265e-8f
 - cardiac ultrastructure and, 265e-5
 - contractile process in, 265e-5-6, 265e-5f, 265e-6f, 265e-7f
 - excitation-contraction coupling in, 265e-7f
 - sliding filament model for, 265e-5-6
- Cardiac glycosides, 473e-9t
- Cardiac imaging, 270e-1
- after myocardial infarction, 270e-19-20
 - in CAD evaluation. *See* Coronary artery disease (CAD), diagnosis of
 - in cardiac mass evaluation, 270e-23-24, 270e-23f, 270e-24f
 - in cardiac structure and function assessment, 270e-6-7
 - in chest pain evaluation. *See* Chest discomfort/pain, approach to the patient
 - in congenital heart disease, 270e-24-25, 270e-25f, 270e-26f. *See also specific defects*
 - contrast agents for, 270e-9
 - CT. *See* Computed tomography (CT), in cardiac disease
 - CT angiography. *See* Computed tomography (CT) angiography (CTA)
 - diagnostic accuracy of, 270e-9t
 - echocardiography. *See* Echocardiography
 - in infectious and inflammatory disease, 270e-24, 270e-24f, 270e-25f
 - MRI. *See* Cardiac magnetic resonance imaging (CMR)
 - in new-onset heart failure, 270e-21-22, 270e-21f, 270e-22f, 270e-27v
 - nuclear, 270e-2-4, 270e-2f. *See also* Myocardial perfusion imaging
 - in pericardial disease, 270e-22-23, 270e-22f, 271e-8f, 271e-8v
 - radiation exposure in, 270e-7-9
 - in valvular heart disease, 270e-15-16. *See also specific conditions*
- Cardiac index, 1748t
- Cardiac jelly, 265e-11
- Cardiac magnetic resonance imaging (CMR)
- after myocardial infarction, 270e-19-20, 270e-20f
 - in aortic regurgitation, 270e-18, 270e-18f, 270e-27v, 1536
 - in atrial myxoma, 289e-2f
 - in CAD evaluation, 270e-12-13, 270e-13f, 270e-26v
 - in cardiac amyloidosis, 270e-21, 270e-21f, 271e-6f
 - in cardiac masses, 270e-23-24, 270e-23f, 270e-24f
 - in chest pain evaluation, 102, 270e-15, 270e-15f, 270e-26v
 - in complete heart block, 271e-7f
 - in congenital heart disease, 270e-25, 270e-26f
 - in cor pulmonale, 1506
 - in hypertrophic cardiomyopathy, 270e-21, 270e-21f, 271e-7f, 271e-7v
 - in iron overload, 270e-21, 270e-27v
 - in left ventricular dysfunction evaluation, 270e-6, 270e-6v
 - in metastatic lung cancer, 271e-9f, 271e-9v
 - in mitral regurgitation, 270e-19
 - in myocardial infarction evaluation, 271e-4f, 271e-4v, 1601-1602
 - in myocarditis, 1559f
 - in patient with pacemaker/defibrillator, 270e-9
 - in pericarditis, 270e-23, 270e-23f, 270e-27v, 271e-8f, 1576f
 - principles of, 270e-4, 270e-6
 - pulse sequences for, 270e-6, 270e-6t
 - in right ventricular dysfunction, 270e-7, 270e-26v
 - stress testing with, 51e-8, 271e-3f, 271e-3v, 1585
 - in tetralogy of Fallot, 1527

- Cardiac magnetic resonance imaging (CMR)
(*Cont.*):
in valvular heart disease, 270e-6
in ventricular tachycardia, 1491, 1493f
- Cardiac mass, 270e-23–24, 270e-23f. *See also*
Cardiac tumors
- Cardiac murmurs. *See* Heart murmurs
- Cardiac output
calculation and normal values, 1748t
control of, **265e-7**
measurement of, 1462–1463
in pregnancy, 45
in sepsis/septic shock, 1755
in shock, 1729, 1745, 1747, 1748t
- Cardiac resynchronization therapy, 273e-7, 1514
- Cardiac risk assessment, 51–53, 52f, 52t, 53f, 53t
- Cardiac-specific troponins. *See* Troponins
- Cardiac stem cells, 453t
- Cardiac syncope
in arrhythmias, 145t, 146–147, 273e-4, 273e-5
ECG in, 148
pattern of, 1439
in structural heart disease, 145t, 147
treatment of, 147
- Cardiac tamponade, **1573**
cardiogenic shock in. *See* Cardiogenic shock
clinical features of, 1573, 1573t
diagnosis of, 1574
differential diagnosis of, 1462, 1463t, 1573t, 1762t
dyspnea in, 47e-2
ECG in, 1458, 1459f, 1573, 1573t
low-pressure, 1574
malignant, 1788–1789
paradoxical pulse in, 1574
pathophysiology of, 1573
treatment of, 1574
- Cardiac thrombus, 270e-23, 270e-23f
- Cardiac transplantation, **1516**
contraindications to, 1517
donor allocation system for, 1516–1517
global considerations in, 1516
immunosuppression for, 1517, 2764t, 2766t
indications for, 1517
LVADs as bridge to, 1518–1519, 1519f
posttransplant complications
allograft CAD, 1517–1518
infection, 924t, 926, 1518
malignancy, 1518
results of, 1517, 1517f
surgical technique, 1516
- Cardiac trauma, **289e-3**
aortic rupture in, 289e-3, 1637
blunt, 289e-3
myocardial contusions, 289e-3
nonpenetrating, 289e-3
penetrating, 289e-3
pericarditis following, 1575
treatment of, 289e-3
- Cardiac tumors, **289e-1**
benign, 289e-1–2
clinical features of, 289e-1–2
CMR in, 289e-2f
echocardiography in, 270e-23–24, 270e-24f, 289e-1f
metastatic, 270e-24, 271e-8f, 271e-8v, 289e-2
primary, 270e-24, 289e-1–2. *See also* Myxoma sarcoma, 289e-2
- Cardiobacterium hominis*, 183e-1, 183e-2t
- Cardioembolic stroke, 2563–2564. *See also*
Ischemic stroke
- Cardio-fascio-cutaneous syndrome, 436t
- Cardiogenic shock, **1759**
clinical features of, 1760
compressive, **1750**
diagnosis of, 1759–1760, 1761f
etiology of, 1759, 1760t
in free wall rupture, 1762
hemodynamic patterns in, 1762t
iatrogenic, 1764
incidence of, 1759
in mitral regurgitation, 1762
in myocardial infarction, 1608
in myocarditis, 1762
pathophysiology of, 1759, 1759f
patient profile in, 1759
physiologic characteristics of, 1748t
prognosis of, 1762
in right ventricular failure, 1762
timing of, 1759
treatment of, 1760–1762, 1761f
in ventricular septal rupture, 1762
- Cardiolipin, 372e-24t, 406
- Cardiomyocytes, 90e-2, 265e-5, 265e-5f
- Cardiomyopathy, **1553**
cardiac arrest risk factors in, 1767
cardiogenic shock in. *See* Cardiogenic shock
in Chagas disease, 1395, 1396
classification of, 1553, 1554t
clinical features of, 1553, 1554t
definition of, 1553
dilated. *See* Dilated cardiomyopathy
dyspnea in, 47e-2
genetic considerations in, 1553–1556, 1555t, 1556f
HIV-associated, 1255, 1560
hypertrophic. *See* Hypertrophic cardiomyopathy
idiopathic, 1565
initial evaluation of, 1554t
iron-overload, 270e-21, 270e-27v
ischemic, 1580
in low- and middle-income countries, 15–16
in metabolic disorders, 1565–1566
overlapping types of, 1558t, 1565–1566
peripartum, 47, 1562
radiation-related, 1567
restrictive. *See* Restrictive cardiomyopathy
in rheumatoid arthritis, 2138
syncope in. *See* Cardiac syncope
Takotsubo (stress-related), 97, 289e-3, 1565
ventricular tachycardia in, 1495, 1495t, 1498
- Cardiopulmonary bypass
acute kidney injury and, 1802–1803
brain injury after, 463e-2–3, 463e-2f
platelet dysfunction in, 730
for rewarming, 478e-3t
- Cardiopulmonary resuscitation, 66, 478e-2, 1768
- Cardiorenal syndrome, 1508
- Cardiovascular collapse, 1764, 1765t. *See also*
Cardiac arrest
- Cardiovascular disease
in alcohol use disorder, 2725
in amyloidosis, 721, 722
in Andersen-Tawil syndrome, 462e-18
approach to the patient, **1439**
arrhythmias in. *See* Cardiac arrhythmias
biology of, **265e-1**
in carcinoid syndrome, 564–565, 564–567
cardiac symptoms in, 1439
congenital. *See* Congenital heart disease (CHD)
coronary artery. *See* Coronary artery disease (CAD)
cyanosis in, 249
deaths from, 56t, 467
estimated 2010–2030, 266e-3t
vs. other causes, 266e-2f
by race and Hispanic origin, 16e-1f
by sex, 6e-1t
in women, 6e-1t, 6e-2f
worldwide by income group, 11t, 266e-3, 266e-3f
worldwide by region, 266e-4, 266e-4f
- depression in, 2714
in diabetes mellitus, 290e-1, 290e-1t
atherosclerosis, 291e-6–7
autoimmune pancreatitis and, 2099
cardiomyopathy, 1563
global considerations in, 266e-4
ischemic heart disease, 1587
morbidity and mortality data, 2427
risk factors for, 2428
treatment and prevention of, 2427–2428
- diagnosis of, 1439–1440
imaging tests for. *See* Cardiac imaging
physical examination. *See* Cardiovascular system, physical examination of
- pitfalls in, 1440–1441
in women vs men, 6e-2–3
- diet and, 266e-4
dyspnea in, 47e-1t, 47e-2, 47e-2t
edema in, 251–252, 252t
epidemiologic transition of, 266e-1–2, 266e-1t
epidemiology of, **266e-1**, 266e-3, 266e-3f
erectile dysfunction in, 326
exercise and, 266e-4
family history in, 1440
functional classification of, 1439–1440, 1440t, 1501, 1501t, 1581t
gene therapy in, 91e-2f, 91e-4
genetic considerations in, 291e-10
global burden of, 11t, 15–16, 266e-2–3, 266e-3f, 266e-3t
heart failure. *See* Heart failure (HF)
in hemochromatosis, 270e-21, 290e-1t, 1471, 1564, 2517
hypercholesterolemia and, 266e-4
in hypothermia, 478e-2t
in IBD, 1959
immunizations recommendations for, 788f
infections in cancer patient, 489
ischemic. *See* Ischemic heart disease
ischemic stroke in, 2564, 2566t
in Kearns-Sayre syndrome, 462e-15
as late consequence of cancer/cancer therapy, 621, 1562–1563
in low- and middle-income countries, 11t, 15–16
in megaloblastic anemia, 643
metabolic syndrome and, 1439, 2450
in muscular dystrophy, 462e-6, 462e-9, 462e-10
myocardial infarction. *See* Myocardial infarction (MI)
natural history of, 1439
obesity and, 266e-4, 415e-6
in Paget's disease of bone, 426e-2
in pregnancy, 46–47
prevention of, 17, 266e-4–5, 1441
pulmonary edema in, 47e-4
rheumatic. *See* Rheumatic fever, cardiovascular complications of
in rheumatoid arthritis, 2138
risk factors for, 266e-4, 291e-4, 291e-6t, 2725. *See also* Atherosclerosis
in sarcoidosis, 2208
in SLE, 27, 2129t
smoking and, 266e-4, 2729–2730, 2730t
syncope in. *See* Cardiac syncope
in syphilis. *See* Syphilis, secondary
in systemic sclerosis, 2158, 2162
testosterone therapy and, 2373–2374
tissue engineering strategies in, 92e-2, 92e-3v
in travelers, 799
treatment of, 1441
valvular. *See* Valvular heart disease
in women, 6e-1t, 6e-2–3, 6e-2f
- Cardiovascular system
biology of, **265e-1**
assessment of cardiac function, 265e-9–11, 265e-10f
blood vessels. *See* Blood vessels
cardiac contraction. *See* Cardiac contraction
control of cardiac performance and output, 265e-7–8, 265e-9f, 265e-9t
developmental, 265e-11–12, 265e-11f
gut microbiota and, 86e-7
in hypoxia, 247
imaging of. *See* Cardiac imaging
physical examination of, **1442**
abdomen, 1443
approach to, 1442
arterial pulses, 1445–1446, 1445f, 1446f
blood pressure, 1444–1445
cardiac auscultation
dynamic, 1449–1450, 1450t
heart murmurs, 1448–1449, 1448f, 1449f. *See also* Heart murmurs
heart sounds, 1447–1448, 1447f. *See also* Heart sounds
in pericardial disease, 1450
of prosthetic valves, 1450

- chest, 1443
 extremities, 1443
 head and neck, 1443
 heart inspection and palpation, 1446–1447
 jugular venous pressure and waveform, 1443–1444, 1444f
 skin, 1442–1443
 in sleep, 187
- Cardioversion
 anticoagulation before, 1485, 1485t, 1487
 for atrial fibrillation, 1487
 for atrial flutter, 1485
 for cardiac arrest, 1769
 for preexcited tachycardias, 1482
- Care coordination, for elderly, 83–84, 83t
- Caregiver(s), 65–66, 93e–4–5
- Caregiving ratios, 93e–4–5
- Caretaker genes, 101e–2, 101e–5
- Carey–Coombs murmur, 51e–6
- Carfilzomib, 102e–5t, 103e–18t, 103e–19, 717
- Caries, dental, 46e–4f, 236, 1097
- Carisoprodol, 473e–10t
- Carmustine (BCNU)
 adverse effects of, 103e–9t, 607t, 1714
 drug interactions of, 103e–10t
 for melanoma, 499
- Carney's syndrome/complex (LAMB/NAME syndrome)
 adrenal nodules in, 2314, 2321, 2343
 clinical features of, 2265t, 2266, 2343
 genetic considerations in, 2266, 2337t, 2344
 myxoma in, 289e–1
 skin manifestations of, 359, 360t, 1443
- Carnitine deficiency, 435e–1t
- Carnitine metabolism disorders, 1563, 1565
- Carnitine palmitoyltransferase (CPT), 462e–15
- L-Carnitine therapy, 63
- Caroli's disease/syndrome, 552, 1853, 2068
- Carotenemia, 96e–7
- Carotenoderma, 279
- Carotenoids, 96e–6, 96e–7
- Carotid artery
 embryonic development of, 265e–11
 intima media thickness, 1241
 occlusion of, 386e–4f, 2574, 2574f, 2575f.
See also Carotid stenosis
- Carotid artery pulse, 1445, 1445f
- Carotid body, 47e–1
- Carotid cavernous fistula, 207
- Carotid sinus hypersensitivity, 1467t, 1469, 1470, 1471, 1471t
- Carotid sinus massage, 148
- Carotid sinus pressure, PSVT, 1484
- Carotid sinus stimulation, 1625
- Carotid stenosis. *See also* Atherosclerosis
 asymptomatic, surgical treatment of, 2571
 risk factors for, 2569
 stroke due to, 2565, 2567t
 treatment of, 296e–5, 296e–5f, 2571
- Carpal tunnel syndrome
 in acromegaly, 2241
 in amyloidosis, 724
 conditions associated with, 2221
 neck pain in, 122
 in pregnancy, 49
 in systemic sclerosis, 2162
 wrist pain in, 2221
- Carpenter's syndrome, 415e–5t
- Carriers, 332e–6
- Carrión's disease, 1081t, 1083
- cART. *See* Combination antiretroviral therapy (cART)
- Cartilage
 engineered tissue for replacement of, 92e–3t
 joint failure related to, 2227, 2227f
 regeneration of, for osteoarthritis, 2233
- Cartilage hair hypoplasia, 375e–2, 418, 2109
- Carvajal syndrome, 1555t
- Carvalho's sign, 51e–3, 51e–4, 1506, 1540
- Carvedilol
 genetic variations in response to, 39t, 40
 for heart failure, 1512t
 for hypertension, 1624, 1624t
 metabolism of, 34t
 overdose/poisoning with, 473e–9t
- Casal's necklace, 96e–4
- Cascara, 273
- Case-control study, 442t
- Caseous necrosis, 1107
- Case reports, 25
- CASPAR (Classification Criteria for Psoriatic Arthritis), 2177, 2177t
- Caspase, 102e–8, 102e–9f, 372e–23, 444e–9
- Caspase-8, 102e–8, 102e–10
- Caspofungin
 for *Candida* infections, 233, 1344t, 1345, 1910
 for febrile neutropenia, 491
 indications for, 1331
 for infections in cancer patient, 686
 for mucormycosis, 1352t
 for sepsis/septic shock, 1757t
- CaSR gene mutations, 2092, 2340, 2475
- Cast(s), urinary, 62e–11–12f, 294
- Castell's method, splenic percussion, 411
- Castelo dos Sonhos virus, 1307t
- Castleman's disease
 vs. AA amyloidosis, 723
 HHV-8 and, 214e–7, 697, 924, 1194
 vs. lymphoma, 710
 multicentric, in HIV infection, 1271
 paraneoplastic pemphigus and, 372
 treatment of, 710
 vs. tuberculosis, 1109
- Cat(s)
 bite-wounds, 167e–1
 approach to the patient, 167e–2
 cellulitis in, 830
 infections from, 167e–1, 183e–3
 microbiology of, 167e–1
 prophylactic therapy, 167e–3
 treatment of, 167e–2–3, 167e–3t, 832t
- cat-scratch disease. *See* Cat-scratch disease
- flea, 1078, 2750
 in *T. gondii* life cycle, 1399, 1399f
 tapeworms, 1435
- Catagen, 331
- Catalepsy, 1771
- Catalogue of Somatic Mutations in Cancer (COSMIC), 101e–8
- Catamenial epilepsy, 2559
- Cataplexy, 147, 190, 194v
- Cataract
 etiology of, 204
 in HSCT recipient, 139e–3
 radiation-induced, 103e–4, 623
 sunflower, 2519
 surgical treatment of, 204
 tamoxifen use and, 531
 visual loss in, 204
- Catatonia, 1771
- Catecholamine(s)
 in cardiovascular regulation, 1613
 in pheochromocytoma, 2329, 2330t
 as tumor marker, 473t
- Catecholaminergic polymorphic ventricular tachycardia, 147, 1498
- Catechol-O-methyltransferase (COMT)
 inhibitors, 2615, 2616t
- Caterpillar stings/dermatitis, 2750
- Cat eye syndrome, 83e–6
- Catfish sting, 2740
- Cathartics, 473e–6
- Catheter ablation
 for accessory pathway, 1483
 for atrial fibrillation, 1489
 for atrial flutter, 1485
 for AV nodal reentrant tachycardia, 1480
 for cardiac arrhythmias, 273e–5–7, 273e–6f
 for focal atrial tachycardia, 1475
 for ventricular arrhythmias, 1477, 1499–1500
- Catheter-related bloodstream infections
 in cancer patient, 487, 487t, 489
Candida, 98e–7, 915
 diagnosis of, 916
 enterococcal, 973
 exit-site, 487, 487t
 gram-negative bacteria in, 183e–3
 with parenteral nutrition, 98e–7–8
 pathophysiology of, 915
 prevention of, 12e–2, 914t, 916
 in transplant recipient, 919, 928
 treatment of, 916
 tunnel-site, 487, 487t
- Catheter-related urinary tract infections, 868, 913–914, 914t
- Cathinones, synthetic (bath salts), 469e–3–4
- Cation-exchange resin, for hyperkalemia, 312
- Cation transport disorders, 653
- Cat-scratch disease
 approach to the patient, 1079–1080
 clinical features of, 1079, 1080–1081f
 diagnosis of, 1079
 epidemiology of, 1078
 etiology of, 1078
 in HIV infection, 1268
 lymphadenopathy in, 408
 ocular involvement in, 1081f
 pathogenesis of, 1079
 prevention of, 1080
 skin manifestations of, 829, 1080f
 treatment of, 1080, 1081t
- Catu virus, 1309t
- Cauda equina syndrome
 clinical features of, 115, 1791
 diagnosis of, 115
 treatment of, 115
 weakness in, 156, 157
- Caudal dysplasia, 48
- Causalgia, 90, 105
- Cautious gait, 163
- Cavernomas, 441e–38f
- Cavernous angiomas, 2586
- Cavernous malformations, 440e–7f
- Cavernous sinus syndrome, 2650, 2650f
- Cavernous sinus thrombosis, septic, 783, 906
- CBC (complete blood count), 393, 394t
- CBFA2 gene mutations, 431t
- CBG (corticosteroid-binding globulin), 2378
- CBG (cortisol-binding globulin), 2312
- CBL gene mutations, 135e–6
- CBT. *See* Cognitive behavioral therapy (CBT)
- CBX2 gene mutations, 2354t
- CCBs. *See* Calcium channel blockers (CCBs)
- C-C chemokine family, 372e–14, 372e–14t
- CCI (corrected count increment), 138e–3
- CCK (cholecystokinin), 415e–2, 2090
- CCK (cholecystokinin)-releasing factor, 2091
- CCL2 gene, 1240t, 1241
- CCL3 gene, 1240t, 1241
- CCL4L gene, 1240t
- CCL5 gene, 1240t
- CCL11. *See* Eotaxin (CCL11)
- CCL17 (TARC), 372e–12t, 1674, 1674f
- CCL22 (MDC), 372e–12t, 1674, 1674f
- CCNU (lomustine), 103e–9t, 601
- CCR1, 372e–13t
- CCR2, 372e–13t
- CCR2 receptor, 1240t
- CCR3, 372e–13t, 1674
- CCR4, 372e–13t
- CCR5 receptor
 assays of, 1248–1249
 in HIV infection
 pathogenesis, 145e–1, 145e–2t, 145e–4, 372e–31, 1235–1236, 1236f, 1238
 susceptibility and response, 1239, 1240t, 1241t
 viral replication cycle, 1216, 1218f
 ligands, cell types, and disease connection, 372e–13t
 maraviroc response and, 39t, 1239, 1280f
- CCR6, 372e–13t
- CCR7, 372e–13t
- CCR8, 372e–13t
- CCR9, 372e–13t
- CCR10, 372e–13t
- CD1a (T6, HTA-1), 372e–3t
- CD1b, 372e–3t
- CD1c, 372e–3t
- CD1d, 372e–3t
- CD2 (T12, LFA-2), 372e–3t

- CD3 (T3, Leu-4), 372e-3t
 CD4 (T4, Leu-3), 372e-3t
 CD4 surface antigen, 1216, 1218f
 CD4+ T cells
 in adaptive immunity, 372e-9f, 372e-22
 functions of, 372e-17
 helper, 372e-14f, 1673
 in HIV infection
 dysfunction and depletion, 1227–1228, 1231, 1236–1237
 early effects, 1228, 1229f
 HIV-specific, 1244
 latently infected, resting, 1230, 1230f
 for monitoring immunologic competence, 1247–1248, 1248f
 viral invasion, 1216–1217, 1217f, 1229–1230
 in immunity to abscesses, 849
 in *M. tuberculosis* infection, 1107
 in nontuberculous mycobacterial infections, 1129
 in response to viral infections, 214e-8
 in rheumatoid arthritis, 2141, 2142f
 in *T. gondii* infection, 1401
 in transplant rejection, 1827, 1828f
 CD4+ T lymphocytopenia, idiopathic, 1272
 CD7 (3A1, Leu-9), 372e-3t
 CD8 (T8, Leu-2), 372e-3t
 CD8+ cytolytic T lymphocytes (CTLs), in HIV infection, 1229–1230, 1237, 1244
 CD8+ T cells
 in adaptive immunity, 372e-9f
 in COPD, 1701
 in HIV infection, 1237, 1244–1245
 in *L. monocytogenes* infection, 982
 in *M. tuberculosis* infection, 1107
 in response to viral infections, 214e-8
 in *T. gondii* infection, 1401
 in transplant rejection, 1827, 1828f
 CD14 (LPS-receptor)
 in immune response, 372e-2, 372e-3t
 in phagocytosis, 145e-5, 145e-6f
 CD16 (FcyRIIIa), 372e-3t, 372e-6
 CD18 gene mutations, 419
 CD19 (B4), 372e-3t
 CD20 (B1), 372e-3t
 CD21 (B2, C2, EBV-R, C3dR), 372e-3t
 CD22 (BL-CAM), 372e-3t
 CD23 (FceRII), 372e-3t, 372e-8
 CD25, as tumor marker, 473t
 CD28, 372e-3t
 CD30, as tumor marker, 473t
 CD32 (FcyRII), 372e-3t
 CD34 antibodies, 139e-1
 CD40, 372e-3t
 CD40 deficiency, 418, 2110
 CD40 ligand, 276, 742, 2110
 CD40 surface antigen, 276, 742
 CD45 (LCA, T200, B220), 372e-3t
 CD45RA, 372e-3t
 CD45RB, 372e-3t
 CD45RC, 372e-3t
 CD45RO, 372e-3t
 CD56, 372e-6
 CD64 (FcyRI), 372e-3t
 CD80 (B7-1, BB1), 372e-3t
 CD86 (B7-2, B70), 372e-3t
 CD89 (FCaR), 372e-3t
 CD95 (APO-1, Fas), 372e-3t
 CD152 (CTLA-4), 372e-3t
 CD154 (CD40L), 372e-3t
 CD279 (PD-1), 372e-3t
 CDC. *See* Centers for Disease Control and Prevention (CDC)
 CDC73 gene mutations, 2340
 CDCA (chenodeoxycholic acid), 2076
 CD (cluster of differentiation) classification, human leukocyte surface antigens, 372e-1, 372e-3–4t
 CDH1 gene mutations, 101e-6t, 534
 CDK4 gene mutations, 101e-2t, 494
 CDKN2A gene mutations, 494, 554
 CDNK1B gene mutations, 2336t, 2342
 CDX-2, in carcinoma of unknown primary, 120e-2, 120e-2t
 CEA. *See* Carcinoembryonic antigen (CEA)
 CEAP classification, chronic venous insufficiency, 1651, 1652t
 CEBPA gene mutations, 679, 680, 680t
 Cecitis. *See* Enterocolitis, neutropenic
 Cefaclor, 363, 941
 Cefadroxil, 939, 941
 Cefamandole, 941
 Cefazolin
 for cellulitis, 228, 832t
 indications for, 940t, 941, 945t
 for infective endocarditis, 822t, 826t
 for intraabdominal infections, 767t
 prophylactic, 945t
 resistance to, 940t
 for staphylococcal infections, 962t
 Cefdinir, 229t, 941
 Cefepime
 adverse effects of, 942
 for *Citrobacter* infections, 1035
 for community-acquired pneumonia, 808t
 for *Enterobacter* infections, 1034
 for febrile neutropenia, 491
 for gram-negative bacterial infection, 1026
 for health care–associated pneumonia, 767t, 812t
 indications for, 940t, 941
 for intraabdominal infections, 767t
 for meningitis, 887t, 889, 889t
 for osteomyelitis, 840t
 for *P. aeruginosa* infections, 808t, 1045t
 resistance to, 933, 940t
 for sepsis/septic shock, 767t, 780t, 1757t
 Cefixime, 937t, 941, 1009t
 Cefoperazone, 941
 Cefotaxime
 for community-acquired pneumonia, 767t, 808t
 for epiglottitis, 235, 1011
 for gonococcal infections, 1009t
 indications for, 941
 for infective endocarditis, 821, 823
 for intraabdominal infections, 767t
 for leptospirosis, 1145t
 for Lyme disease, 1153, 1153f
 for meningitis, 887t, 889, 889t, 952, 1011
 for meningococcal infections, 1000
 for nongonococcal bacterial arthritis, 835
 for peritonitis, 288, 847, 2066
 for pneumococcal infections, 952
 resistance to, 933
 for sepsis/septic shock, 1757t
 Cefotetan
 for anaerobic bacterial infections, 1101
 indications for, 940t, 941
 for pelvic inflammatory disease, 878t, 879
 prophylactic, 945t
 resistance to, 940t, 1026, 1101
 Cefoxitin
 for anaerobic bacterial infections, 1101
 for bite-wound infections, 167e-2, 167e-3t
 for gas gangrene, 832t, 991t
 for gonococcal infections, 1009t
 indications for, 940t, 941
 for intraabdominal infections, 767t
 for nontuberculous mycobacterial infections, 205e-8
 for pelvic inflammatory disease, 878, 878t, 879
 for peritonitis, 848
 prophylactic, 945t
 resistance to, 940t, 1026, 1101
 for skin and soft tissue infections, 832
 Cefpirome, 941
 Cefpodoxime, 808t, 937t, 941
 Cefprozil, 941
 Ceftaroline
 action of, 961
 indications for, 940t, 941
 for infective endocarditis, 823
 for MRSA infections, 933, 961, 962t
 resistance to, 940t
 Cefazidime
 for febrile neutropenia, 103e-23, 491, 1045t
 for glanders, 1045t
 for health care–associated pneumonia, 767t, 812t
 indications for, 940t, 941
 for intraabdominal infections, 767t
 for melioidosis, 1045t
 for meningitis, 887t
 for nongonococcal bacterial arthritis, 835
 for osteomyelitis, 840t
 for otitis externa, 229
 for *P. aeruginosa* infections, 1045t
 resistance to, 940t
 Ceftriaxone, 1009t, 1091t
 Ceftriaxone
 for actinomycosis, 1091t
 for *Bartonella* endocarditis, 1081t
 for chancroid, 880t, 1013
 for community-acquired pneumonia, 767t, 808t
 for deep neck infections, 235
 for donovanosis, 198e-2
 drug interactions of, 939t
 empirical uses of, 767t, 780t
 for enteric fever, 1052t
 for enterococcal infections, 975t, 976
 for epididymitis, 872
 for epidural abscess, 767t, 780t
 for epiglottitis, 235, 1011
 for gonococcal infections, 835, 872, 1008, 1009t
 for health care–associated pneumonia, 812t
 indications for, 940t
 for infective endocarditis, 767t, 780t
 Bartonella, 823t
 HACEK, 183e-1, 183e-2t, 822t
 streptococcal, 822t, 823
 for intraabdominal infections, 767t
 for intracranial infections, 780t
 for leptospirosis, 1145t
 for Lyme disease, 836, 1153, 1153f
 for meningitis, 767t, 780t, 887t, 889, 889t, 952, 1011
 for meningococcal infections, 1000
 for meningococcal prophylaxis, 1003
 for *Neisseria meningitidis* infections, 1087
 for nongonococcal bacterial arthritis, 835
 for nontyphoidal *Salmonella* infections, 1054t
 for osteomyelitis, 840t, 841t
 for otitis media, 229t
 for pelvic inflammatory disease, 878t
 for peritonitis, 847, 848
 for pneumococcal infections, 952
 for postsplenectomy sepsis, 780t
 in pregnancy and lactation, 937t
 prophylactic, 826t, 889
 for purpura fulminans, 780t
 for relapsing fever, 1148, 1148f
 resistance to, 872f, 933, 940t
 for sepsis/septic shock, 1757t
 for *Shigella* infections, 1058t
 for syphilis, 1139
 for Whipple's disease, 1093–1094
 Cefuroxime
 for bite-wound infections, 167e-2
 for community-acquired pneumonia, 808t
 for epiglottitis, 235
 indications for, 941
 for intraabdominal infections, 767t
 for Lyme disease, 1153, 1153f
 for otitis media, 229t
 in pregnancy and lactation, 937t
 Celecoxib
 adverse effects of, 92
 drug interactions of, 43
 genetic variations in response to, 39t
 for gout, 2234
 for osteoarthritis, 2232
 for pain, 91–92, 91t
 for polyposis syndromes, 539
 Celiac disease (gluten-induced enteropathy), 1940
 clinical features of, 1940
 ataxia, 2626
 diarrhea, 269, 1940, 1942

- neuropathy, 2683
 oral, 236, 237
 cobalamin malabsorption in, 645
 complications of, 1942, 2347, 2366
 diagnosis of, 1941t, 1942
 barium radiography in, 1939f
 endoscopic, 1897f
 small-intestinal mucosal biopsies in, 1939, 1940t, 1941f, 1942
 tTg antibody test in, 2347
 diseases associated with, 1942
 APS-2, 2346t, 2347, 2347t
 dermatitis herpetiformis, 373, 1942
 diabetes, 1942
 enteropathy-type T-cell lymphoma and, 135e-3
 environmental factors in, 1940
 etiology of, 1940–1942
 genetic considerations in, 373e-7, 373e-8t, 1941–1942, 2347t
 gluten restriction in, 1942
 immunology of, 372e-23, 372e-24t
 incidence of, 269, 1940
 pathology of, 269
 refractory, 1942
 treatment of, 263, 272
 vitamin A deficiency in, 96e-7
 vitamin K deficiency in, 96e-8
 Celiac sprue. *See* Celiac disease
 Cell cycle
 in cancer cells, 102e-1–2, 102e-1t
 checkpoints, 101e-3, 102e-1–2
 DNA replication in, 427
 segregation of mitochondrial DNA in, 85e-2
 Cell free fetal DNA screening, 83e-4
 Cell lysis, in hypothermia, 478e-2
 Cell-mediated immunity
 destruction of tumor cells by, 103e-20
 in HIV infection, 1243t, 1244–1245
 in tuberculosis, 1106
 Cell necrosis, 372e-1
 Cellular metabolism, genetic defects of, 1555t
 Cellular senescence, 102e-3, 102e-8
 Cellular transport, 332e-3
 Cellulitis
 anaerobic bacterial, 995, 1099
 auricular, 228
 burn-wound, 166e-2, 166e-2f
 in cancer patient, 485–486
 diagnosis of, 830
 differential diagnosis of, 366
 E. coli, 1028–1029
 etiology of, 828t, 830
 H. influenzae, 1011
 Nocardia, 1085, 1087t
 pathophysiology of, 827f, 830
 periorbital, 207, 830
 streptococcal, 830, 967–968
 treatment of, 832t, 965t
 Cell wall synthesis inhibitors, 930, 931t, 932f
 Cementum, 236
 Centers for Disease Control and Prevention (CDC)
 infection-control guidance and oversight, 912t, 913
 Vaccines & Immunizations site, 791
 Centipede bite, 2750
 Central cord syndrome, 2652
 Central core disease, 462e-12
 Central cyanosis, 249, 249t. *See also* Cyanosis
 Central fever, 1773
 Central hypoventilation syndrome, 1722
 Central nervous system (CNS)
 cytokine production in, 124
 in hypothermia, 478e-2t
 in hypoxia, 247
 paraneoplastic disorders of. *See* Paraneoplastic syndromes, neurologic
 vasculitis of. *See* Vasculitis, CNS
 Central nervous system (CNS) cancer. *See also* specific types and sites
 brain tumors. *See* Brain tumors
 deaths from, 467t, 470t
 incidence of, 467t
 lymphoma. *See* Primary central nervous system lymphoma (PCNSL)
 metastatic, 1792–1793
 Central nervous system (CNS) infections
 actinomycosis, 1089
 after solid organ transplantation, 463e-3
 anaerobic bacterial, 1098, 1102
 Aspergillus, 441e-9–10f, 1346–1347
 brain abscess. *See* Brain abscess
 in cancer patient, 488, 488t
 Candida, 441e-8f
 cranial epidural abscess, 904–905, 904f
 dementia due to, 2607
 empyema. *See* Subdural empyema
 encephalitis. *See* Encephalitis
 intracranial, 780t, 784
 L. monocytogenes, 983
 M. tuberculosis, 1111
 meningitis. *See* Meningitis
 mucormycosis, 1351, 1351f
 mumps, 231e-2–3
 neurocysticercosis. *See* Neurocysticercosis
 Nocardia, 1085
 nontyphoidal *Salmonella*, 1054
 P. aeruginosa, 1045t, 1046
 rabies. *See* Rabies
 sarcoidosis, 909t
 sporotrichosis, 1354, 1354t
 T. gondii, 1401–1402, 1402f, 1403t
 T. whipplei, 1092
 treatment of, 884–885f, 1102
 VZV, 1183. *See also* Herpes zoster (shingles)
 Central pontine myelinolysis, 301, 1782, 1782f
 Central precocious puberty, 2361. *See also* Precocious puberty
 Central retinal artery occlusion, 201, 201f
 Central retinal vein occlusion, 201–202, 203f
 Central sensitization, 87
 Central sleep apnea, 189, 1727
 Central transtentorial herniation, 1772, 1772f
 Central venous catheter
 clinical procedure tutorial, 481e-1v
 infections related to. *See* Catheter-related bloodstream infections
 for parenteral nutrition, 98e-6
 superior vena cava syndrome and, 1788
 Central venous pressure, 1443, 1748t, 1757
 Centromere-associated proteins, 372e-24t, 2157t
 Centronuclear (myotubular) myopathy, 462e-12, 462e-14
 Cepacia syndrome, 487, 1045t, 1048, 1697
 Cephalixin
 for endocarditis prophylaxis, 826t
 for impetigo, 967
 indications for, 939, 941
 for otitis externa, 228
 for pharyngitis, 232f
 in pregnancy and lactation, 937t
 for staphylococcal skin/soft tissue infections, 345, 768t
 Cephalosporins
 action of, 930, 931t
 adverse effects of, 356, 371, 377, 422, 491
 cross-sensitivity to, 385
 for gonococcal infections, 1008
 indications for, 939–940
 for infections in cancer patient, 491
 for nongonococcal bacterial arthritis, 835
 for plague, 261e-5
 in pregnancy and lactation, 937t
 resistance to
 in *E. coli*, 1026
 in gonococcal infections, 1005
 in gram-negative bacteria, 1026
 in *Klebsiella*, 1033
 mechanisms of, 931t, 933
 for staphylococcal infections, 961
 Cerebellar arteries, occlusion of, 2575–2576, 2575f
 Cerebellar ataxia
 channelopathies in, 444e-2t
 early-onset, 451e-3t
 etiology of, 2626, 2627t
 gait disorders in, 163, 164t
 with muscle coenzyme Q deficiency, 451e-3t
 type 1, 451e-4t
 type 2, 451e-4t
 Cerebellar ataxia, deafness, and narcolepsy, 451e-2t
 Cerebellar ataxia mental retardation, 451e-2t
 Cerebellar degeneration/atrophy
 in alcohol use disorder, 2724
 disease-modifying protein identification in, 87e-2, 87e-3t
 falls in, 147, 164
 gait disorders in, 163, 164t
 paraneoplastic, 372e-25t, 614t, 617–618
 Cerebral amyloid angiopathy, 2583–2584, 2583t, 2602
 Cerebral angiography. *See* Angiography, cerebral
 Cerebral arterial gas embolism, 477e-7
 Cerebral artery(ies)
 anatomy of, 2572f
 aneurysm of, 440e-2f
 middle. *See* Middle cerebral artery
 Cerebral autosomal dominant arteriopathy
 with subcortical infarcts and leukoencephalopathy (CADASIL), 441e-18f, 2568, 2608
 Cerebral blood flow, 1615, 1627, 1778, 1778f
 Cerebral edema
 after subarachnoid hemorrhage, 1785
 high-altitude. *See* High-altitude cerebral edema (HACE)
 in hyponatremia, 300
 Cerebral hemispheres. *See* Brain
 Cerebral ischemia, 2559, 2561f
 Cerebral palsy, 163
 Cerebral perfusion pressure, 1779, 1779f
 Cerebral salt wasting, 299
 Cerebral venous sinuses, 905–906, 905f
 Cerebritis, 901
 Cerebrospinal fluid (CSF)
 leakage of, 110, 110f, 457e-2, 2607
 raised pressure. *See* Intracranial pressure, increased
 Cerebrospinal fluid (CSF) analysis
 in bacterial meningitis, 764, 766t, 887–888, 888t
 in chronic meningitis, 907–909t, 909–910
 in coma, 1775
 in delirium, 170
 in encephalitis, 764, 766t
 in fever of unknown origin, 138
 in HIV infection, 1264
 in MS, 2665
 normal values, 443e-3t, 443e-4, 766t
 in parasitic infections, 245e-6t
 in progressive multifocal leukoencephalopathy, 899
 reference values for, 2767t
 specimen collection and transport, 150e-4t
 in subacute meningitis, 898
 in viral encephalitis, 894
 in viral meningitis, 891
 Cerebrotendinous xanthomatosis, 451e-4t
 Cerebrovascular disease, 2559. *See also* Stroke
 approach to the patient, 2560, 2560f
 deaths from, 6e-1t, 11t
 definition of, 2559
 dementia associated with, 170, 172, 173t, 2602
 global considerations in, 11t, 2559
 imaging studies of. *See* Neuroimaging
 incidence of, 2559
 seizures due to, 2547
 types of, 2559
 vascular anomalies, 2585–2586
 in women, 6e-2f
 Cerivastatin, 246e-3t
 Certolizumab
 adverse effects of, 422, 2146t
 for ankylosing spondylitis, 2172–2173
 for enteropathic arthritis, 2179
 for IBD, 1961
 for rheumatoid arthritis, 2145, 2146t
 Ceruloplasmin, 2519, 2520t
 Cerumen impaction, 220
 Cervical adenopathy, 120e-4, 503, 503f

- Cervical angina syndrome, 122
- Cervical cancer, 595
- clinical features of, 595
 - deaths from, 470t, 471t
 - epidemiology of, 595, 1198
 - etiology of, 595
 - global considerations in, 595, 1198
 - in HIV infection, 1271–1272
 - HIV infection and, 1198
 - HPV in. *See* Human papillomavirus (HPV) infection, cervical cancer and
 - incidence of, 468, 471t, 595
 - KIRs with, 372e–15t
 - paraneoplastic syndromes in, 609t
 - during pregnancy, 124e–1t, 124e–2–3
 - prevention of, 478–479, 595
 - prognosis of, 593t
 - risk factors for, 27t, 595
 - screening for, 28, 29t, 84, 480, 481t, 482, 595, 1200
 - staging of, 593t, 596, 596f
 - in transplant recipient, 926
 - treatment of, 595–596
- Cervical collar, 123
- Cervical dystonia, 2620
- Cervical spine
- atlantoaxial involvement in rheumatoid arthritis, 2137
 - disk disease, 121t, 122
 - lesions of, 2651
 - trauma to, 121–122, 441e–31f
- Cervical stenosis, 338
- Cervical venous hum, 1449
- Cervicitis, mucopurulent
- C. trachomatis*, 1168–1169, 1170t
 - diagnosis of, 876, 1005
 - etiology of, 870t, 876, 876f
 - gonococcal, 1005
 - treatment of, 876
- Cervix, ectopy of, 876
- Cesarean section, 50
- CESD (cholesteryl ester storage disease), 432e–4t, 432e–5, 2443
- Cesium exposure, 263e–3t, 263e–6t
- Cestodes/cestode infections, 255e–1–2, 1430
- coenurosis, 1435
 - Diphyllobothrium latum* (fish tapeworm), 245e–2t, 645, 1434
 - Dipylidium caninum*/dipylidiasis, 1435
 - Echinococcus*. *See* Echinococcosis (hydatid disease)
 - Hymenolepis diminuta*, 1434
 - Hymenolepis nana* (dwarf tapeworm), 245e–2t, 246e–11, 1434
 - sparganosis, 1435
 - Taenia*. *See* *Taenia saginata* (beef tapeworm) infection; *Taenia solium* (pork tapeworm) infection
- CETP (cholesteryl ester transfer protein), 2438, 2438f
- CETP (cholesteryl ester transfer protein) deficiency, 2446
- CETP (cholesteryl ester transfer protein) inhibitors, 291e–6
- Cetuximab
- action and targets of, 102e–5t, 103e–21, 103e–21t
 - adverse effects of, 103e–26, 380, 505, 543, 1798
 - for colorectal cancer, 102e–5t, 103e–21, 103e–21t, 543
 - genetic variations in response to, 39t, 41
 - for head and neck cancer, 505
 - for Ménétrier's disease, 1932
 - for squamous cell carcinoma, 102e–5t
- Cevimeline, 241
- CF. *See* Cystic fibrosis (CF)
- C fibers, 87, 88
- CFM (confocal fluorescent laser microscopy), 1668
- CFTR gene mutations
- in *Aspergillus* infections, 1346
 - in cystic fibrosis, 145e–5, 429f, 435, 480e–5, 1697–1698, 1698f
 - genetic testing for, 480e–5
 - in pancreatitis, 2092, 2098–2099
 - types of, 435
 - typhoid fever resistance and, 145e–3, 442
- CFTR protein
- in cystic fibrosis, 1697, 1698f, 1700v
 - P. aeruginosa* and, 145e–4f, 145e–5, 1043
 - potentiation of, 1699
- CGDs. *See* Chronic granulomatous diseases (CGDs)
- C gene, of HBV, 2005f, 2006
- CGH (comparative genomic hybridization), 83e–3, 83e–3t
- CGRP (calcitonin gene-related peptide), 88, 444e–3, 2469
- CHADS2 score, 2564, 2566t
- Chagas' disease
- cardiac manifestations of, 1471, 1471t, 1495, 1560
 - clinical features of, 1394f, 1395
 - diagnosis of, 245e–2, 245e–4t, 245e–6t, 1395
 - differential diagnosis of, 1904
 - epidemiology of, 245e–4t, 1395
 - in HIV infection, 1265
 - pathology of, 1394, 1394f
 - prevention of, 1396
 - transfusion-associated, 138e–6, 1394
 - treatment of, 246e–8, 246e–11, 1395–1396, 1560
- Chagoma, 1395
- Chagas virus, 1309t
- Chalazion, 199
- Chamomile, 404t
- Chancre
- syphilitic, 25e–4f, 76e–9f, 237, 238t, 1134, 1134f
 - trypanosomal, 25e–12f, 1397
- Chancroid
- clinical features of, 76e–11f, 879f, 880t, 1012–1013, 1012f
 - diagnosis of, 881, 1013
 - epidemiology of, 879, 1012
 - etiology of, 1012
 - global considerations in, 879
 - in HIV infection, 1259
 - lymphadenopathy in, 408
 - treatment of, 880t, 881, 1013
- Chandipura virus, 1310t, 1316
- Channelopathy(ies)
- in cardiomyopathy, 1554, 1555t
 - muscular, 462e–16–18, 462e–17f, 462e–17t
 - neurologic, 444e–2, 444e–2t
- Chao1, 86e–4
- Chapare virus, 1306t
- Charcot-Leyden crystals, 270, 422, 1348
- Charcot-Marie-Tooth (CMT) disease
- clinical features of, 2678
 - diagnosis of, 2678
 - genetic considerations in, 434, 435, 436t, 444e–1, 2678
 - pathogenesis of, 2678
 - treatment of, 2678
- Charcot's foot, 845, 845f
- Charcot's joint (neuropathic joint disease), 1136, 2243–2244, 2243f, 2243t
- Charcot's triad, 1891, 2083
- CHARGE (coloboma of the eye, heart anomaly, choanal atresia, retardation, genital and ear anomalies) syndrome, 375e–1, 2108
- Charter on Medical Professionalism, 5
- Chart review, 25
- CHD. *See* Congenital heart disease (CHD)
- CHD7 deficiency, 375e–1, 2108
- CHD7 gene mutations, 2363, 2364t
- Chédiak-Higashi syndrome
- cellular/molecular defects in, 372e–7, 419, 420t
 - clinical features of, 236, 357t, 375e–2, 419, 420t
 - diagnosis of, 81e–2, 81e–7f, 420t, 421f
 - treatment of, 96e–5
- Cheek, traumatic lesion inside of, 46e–3f
- “Cheese effect,” 2615
- Cheese washer's lung, 1681t
- Cheilitis, 46e–2f, 501
- Cheilosis, 237, 627, 1945
- CHEK2 gene mutations, 446
- Chelation therapy, 263e–6t, 263e–7, 472e–1
- Chemical burn, oral, 239t
- Chemical restraints, 170
- Chemical terrorism, 262e–1
- characteristics and effects of, 262e–1t
 - decontamination and treatment of, 262e–2t
 - nerve agents. *See* Nerve agents vesicants. *See* Sulfur Mustard
- Chemical worker's lung, 1681, 1681t
- Chemo brain, 622
- Chemo cap, 103e–25
- Chemokine(s)
- in asthma, 1674
 - cell types, 372e–13t
 - definition of, 372e–1
 - disease connections of, 372e–13t
 - functions of, 416–417
 - in HIV infection, 1240t. *See also* CCR5 receptor
 - in pneumonia, 804
 - receptors, 372e–13t
- Chemoreceptors, 47e–1
- Chemoreceptor trigger zone (CTZ), 61, 103e–25, 258
- Chenodeoxycholic acid (CDCA), 2076
- Cherry angioma, 341t
- Cherry hemangioma, 76e–4f, 364t, 365
- Cherry red spot, 40e–4, 40e–4f, 436e–7f
- Chest compression, for cardiac arrest, 1768
- Chest discomfort/pain, 95
- approach to the patient, 98, 98t
 - cardiac biomarkers, 101
 - chest radiography, 101
 - clinical algorithms, 101, 102f
 - CMR, 102, 270e–15, 270e–15f, 270e–26v
 - CT angiography, 102, 270e–14, 270e–14f, 270e–15, 270e–16f
 - ECG, 100–101
 - echocardiography, 102, 271e–1f, 271e–1v
 - in the emergency department, 270e–15
 - history, 99–100, 99f
 - in patients with known CAD, 270e–14–15
 - in patients without known CAD, 270e–13–14, 270e–14f
 - physical examination, 100
 - stress testing, 101–102, 270e–13–14, 270e–14f
- of cardiopulmonary origin, 96t
- aortic disease, 97, 99
 - exertional, 1439
 - lung cancer, 510, 510t
 - mechanisms of, 97
 - myocardial ischemia. *See* Angina; Myocardial infarction
 - non-ischemic myocardial disease, 97
 - pericardial disease, 97, 99, 1571, 1580, 1600, 1610
 - pneumothorax, 98
 - pulmonary disease, 98
 - pulmonary embolism, 97–98
 - respiratory disease, 98, 1661
- critical pathways for, 102–103
- with dysphagia, 256
 - of emotional/psychiatric origin, 96t, 98
 - epidemiology of, 95, 95f
 - of gastrointestinal origin, 96t
 - esophageal disorders, 98, 100, 1901, 1905
 - GERD, 98, 99, 262, 1907
 - hepatobiliary disorders, 98, 100
 - pancreatitis, 98
- incidence of, 95
- of musculoskeletal origin, 96t
 - costochondritis, 98
 - herpes zoster, 98
- myocardial infarction probability with, 99f
- natural history of, 95
 - outpatient evaluation of, 103
 - in pericarditis, 1571
 - in sickle cell anemia, 634
- Chest pain units, 102–103, 1595
- Chest physiotherapy, 1696, 1699

- Chest radiography, **308e-1**
 in alveolar processes, 308e-10f
 in anthrax, 261e-3, 261e-4f
 in ARDS, 1737f
 in asthma, 1675
 in bronchiectasis, 308e-11f
 in chest pain evaluation, 101
 in dyspnea, 47e-4
 in hemoptysis, 246
 in hypersensitivity pneumonitis, 1682
 in interstitial lung disease, 308e-7-9f, 1710
 in Legionnaires' disease, 1016, 1017f
 in lung cancer, 308e-15f
 in lung cancer screening, 509-510, 510t
 in lymphadenopathy, 409
 normal, 308e-1f
 in parenchymal loss, 308e-5-6f
 in pleural effusion, 308e-14f
 in pneumococcal pneumonia, 951f
 in pneumothorax, 308e-13f
 in pulmonary edema, 47e-5
 in pulmonary hypertension, 308e-18f, 1656, 1658f
 in respiratory disease evaluation, 1663, 1664f, 1665f
 in volume loss, 308e-3-4
- Chest tightness, 47e-1, 47e-1t
- Chest tube drainage, for pleural effusion, 474
- Chest wall
 abnormality of, ventilatory restriction due to, 306e-5, 1721
 compliance of, 306e-1-2, 1733
 disorders of, 47e-2
 pressure-volume curves of, 306e-1f
 tenderness in, 100
 trauma to, 51e-2
- Chewing tobacco, 476
- Cheyne-Stokes respiration
 in central sleep apnea, 1726f, 1727
 in coma, 1775
 in heart failure, 1504
 in terminally ill patient, 68, 69t
- CHF (congestive heart failure). *See* Heart failure (HF)
- Chiari malformation, 2658, 2658f
- Chiasmal tumor, 206-207
- Chiasmata, 428
- Chickenpox. *See* Varicella
- Chief cells, 1914
- Chiggers, 2745
- Chigoe flea, 2750
- Chikungunya virus infection
 climate change and, 151e-4-5
 clinical features of, 130t, 134, 837, 1313
 epidemiology of, 130t, 134, 151e-4-5, 802, 894, 1313
 reservoirs and vectors of, 1306t, 1313
 treatment of, 1313
- Chilblain (pernio), 478e-4, 1648f, **1649**
- Child abuse, 17e-1, 167e-2, 2722
- Child-Pugh classification, of cirrhosis, 1994-1995, 1995t
- Children
 atopic dermatitis in, 344
 ECG in, 269e-14f
 fever treatment in, 126
 GH deficiency in, 2257-2258
 Graves' disease in, 2296
 HIV infection in, 1226, 1226f
 liver transplantation in, 2068, 2068t
 malaria in, 1374
 mortality rates in, 9, 13e-3f, 13e-6f, 93e-1-2
 rheumatic fever in, 2149
 seizures in, 2545-2546, 2546t
 smoking in, 475
- Child-Turcotte-Pugh (CTP) score, 2069
- Chile, 13e-6
- Chilean Rose tarantula, 2748
- Chimeric antigen receptor (CAR), 91e-4
- Chimpanzee, as HIV reservoir, 1219
- China, 13e-7, 13e-7f
- Chinese herbal nephropathy, 577, 1804, 1856t, 1861
- Chios virus, 1309t
- CHIPS, 956
- Chironex fleckeri* sting, 76e-11f
- Chiropractic, 14e-1t, 14e-2-3
- Chlamydia* spp., 1165-1166, 1166f
- Chlamydia psittaci*, 1172-1173
- Chlamydia trachomatis*, 1165-1166
- Chlamydia trachomatis* infections
 anorectal, 881
 genitourinary, **1167**
 diagnosis of, 150e-6, 771, 1170-1171, 1170t
 epidemiology of, 869, 1167
 epididymitis, 872, 1168
 incidence of, 869
 lymphogranuloma venereum. *See* Lymphogranuloma venereum
 mucopurulent cervicitis, 1168-1169
 partner-delivered therapy for, 1171
 pathogenesis of, 1166-1167
 proctitis, 1168, 1169
 screening for, 29t, 882, 1172
 spectrum of, 1167
 treatment of, 871, 872t, 1171
 urethral syndrome in women, 1169
 urethritis, 870, 872, 1167-1168
 in immunocompromised patients, 489, 489t
 in neonates, 1169, 1170t, 1171
 pelvic inflammatory disease. *See* Pelvic inflammatory disease (PID)
 perihepatitis, 1169
 in pregnancy, 1169
 prevention of, 1171-1172
 reactive arthritis, 1168
 trachoma, 199, **1172**
- Chlamydophila pneumoniae* infections, **1173**
- Chloral hydrate, 473e-11t
- Chlorambucil, 103e-8, 103e-9t
- Chloramphenicol
 action of, 931t, 932
 adverse effects of, 938t, 945
 aplastic anemia, 664t
 bone marrow failure, 678
 neutropenia, 417
 optic neuropathy, 203
 pure red cell aplasia, 668t
 for anaerobic bacterial infections, 1101, 1102
 for *Campylobacter* infections, 1061
 for Carrión's disease, 1081t
 for donovanosis, 198e-2
 drug interactions of, 945
 for enteric fever, 1052t
 indications for, 941t, 944-945
 for meningitis, 1011
 for meningococcal infections, 1000
 for plague, 261e-5, 261e-5t, 1074t
 in pregnancy and lactation, 937t
 for relapsing fever, 1148f
 resistance to, 931t, 934
 for scrub typhus, 1159
 therapeutic monitoring of, 2764t
 for tick-borne spotted fevers, 1157
 for tularemia, 261e-5t, 261e-7
 for Whipple's disease, 1945
- Chlordiazepoxide
 for alcohol withdrawal, 2727, 2728
 for anxiety disorders, 2711t
 overdose/poisoning with, 473e-10t, 2764t
 pharmacology of, 2711t
 therapeutic monitoring of, 2764t
- Chlorhexidine bath, 918
- Chlorhexidine mouthwash, 216
- Chloride
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference values for, 2758t
 renal transport of, 332e-3f, 332e-7-8
- Chloride channel activators. *See* Lubiprostone
- Chloride channel disorders, of muscle, 462e-18
- Chloridorrhea, congenital, 254
- Chlorine, as chemical terrorism agent, 262e-1
- Chlornaphazine, 575
- 2-Chlorodeoxyadenosine, 103e-10t, 103e-14
- Chloroma, cutaneous (granulocytic sarcoma), 366
- Chloroquine
 action of, 246e-8
 adverse effects of
 bone marrow failure, 664t
 common, 1380t
 cutaneous, 353, 381
 hair discoloration, 380
 myopathy, 462e-20, 462e-20t, 2199
 neuropathy, 2686, 2687t
 occasional, 246e-1t, 1383
 ocular, 206, 1383
 rare, 246e-1t, 246e-8, 1380t
 drug interactions of, 246e-1t
 genetic variations in response to, 39t
 for hypercalcemia, 314
 for malaria, 1378t, 1379, 1380t
 for malaria prophylaxis, 796t, 1383, 1383t
 overdose/poisoning with, 473e-13t
 pharmacology of, 246e-8, 1380t
 for porphyria cutanea tarda, 390, 2530
 resistance to, 1370f, 1379
 for SLE, 2132
 for Whipple's disease, 1094
- Chlorothiazide, 462e-18
- Chlorproguanil, 246e-4t
- Chlorpromazine
 adverse effects of, 2721, 2722t
 cholestatic hepatitis, 284
 chronic cholestasis, 284
 cutaneous, 379
 lupus syndrome, 2218t
 photoallergy, 389t
 secondary parkinsonism, 2611
 for delirium, 64t, 65
 drug interactions of, 42
 for dyspnea, 62
 for medication-overuse headache, 109
 for migraine, 2592, 2592t
 overdose/poisoning with, 473e-8t, 473e-9t
 parenteral, 2592, 2592t
 for schizophrenia, 2721, 2722t
- Chlorpropamide, 82, 727t, 2413
- Chlorthalidone, 1624t
- Choclo virus, 1307t
- Choking/pulmonary-damaging agents, 262e-1t, 262e-2t
- Cholangiocarcinoma, **552**
 clinical features of, 552
 diagnosis of, 552-553
 etiology of, 552
 extrahepatic cholestasis in, 285
 in sclerosing cholangitis, 2086
 sites of origin of, 552
 treatment of, 553, 1885f
- Cholangiopancreatography
 endoscopic. *See* Endoscopic retrograde cholangiopancreatography (ERCP)
 magnetic resonance. *See* Magnetic resonance cholangiopancreatography (MRCP)
- Cholangitis
 abdominal pain in, 106t
 ascending, 1892-1893
 in choledocholithiasis, 1884f, 2083
 immunoglobulin G4-associated, 2085, 2099
 primary sclerosing. *See* Primary sclerosing cholangitis (PSC)
 in transplant recipient, 924t, 927
- Cholecystectomy
 complications of, 2080, 2082-2083
 for hyperplastic cholecystoses, 2083
 indications for, 2080
 with splenectomy, in hereditary spherocytosis, 652
- Cholecystitis, **2080**
 acalculous, 2081
 acute, 2080-2081
 chronic, 2081
 clinical features of, 2080
 abdominal pain, 106, 106t, 2080
 chest pain, 98
 nausea and vomiting, 258, 2080
 complications of, 2081-2082
 diagnosis of, 107, 2081
 emphysematous, 2081

- Cholecystitis (*Cont.*):
 fasting-induced, 415e-6
 gallbladder cancer and, 553
 treatment of, 2080, 2082
- Cholecystokinin (CCK), 415e-2, 1913f, 2090
- Cholecystokinin (CCK)-releasing factor, 2091
- Cholecystopathy, acalculous, 2081
- Cholecystosis(es), hyperplastic, 2083
- Choledochal cysts, 552, 2083
- Choledocholithiasis, **2083**
 in cholangiocarcinoma, 552
 clinical features of, 285, 2083
 complications of, 2083–2084
 diagnosis of, 2084, 2085t
 pathophysiology of, 2083
 treatment of, 2084
- Cholelithiasis. *See* Gallstone(s)
- Cholera, **1061**
 clinical features of, 266t, 856t, **1063**, 1063f
 diagnosis of, 1063
 differential diagnosis of, 266t, 856t
 epidemiology of
 climate change and, 151e-8, 151e-8f
 environmental factors, 1061
 genomics in, 777, 777f
 historical studies, 776
 global considerations in, 1061–1062, 1062f
 pancreatic, 269
 pathogenesis of, 856t, 1062–1063. *See also* *Vibrio cholerae*
 prevention of, 1064–1065. *See also* Cholera vaccine
 treatment of, 1063–1064, 1064t
- Cholera gravis, 1061, 1063
- Cholera vaccine
 killed whole-cell, 1064–1065
 live attenuated viral, 1065
 for travelers, 795, 795t
- Cholestasis
 benign postoperative, 284
 benign recurrent intrahepatic, 284, 2003t, 2004, 2076
 drug-induced, 284, 2023, 2025, 2025t, 2026t, 2030
 extrahepatic, 283, 284t, 285
 imaging in, 283–284
 intrahepatic, 284, 284t
 jaundice in, 283–285
 in lipoprotein disorders, 2443
 liver function tests in, 1997, 1998t
 parental nutrition and, 2030
 of pregnancy, 49, 284, 2029, 2078
 progressive familial intrahepatic, 284, 2003t, 2004
- Cholestatic diseases, 1990, 1990t. *See also* Liver disease/failure
- Cholesteatoma, 218, 230
- Cholesterol, reverse transport of, 291e-2
- Cholesterol absorption inhibitors, 2448–2449, 2448t
- Cholesterol embolism
 in kidney, 62e-9f, 1835t, 1848
 retinal, 40e-4
 skin manifestations of, 368
- Cholesterolemia, 2083
- Cholesterol side-chain cleavage enzyme (CYP11a), 372e-24t
- Cholesterol stones
 gallbladder, 2076–2077, 2077f, 2078t. *See also* Gallstone(s)
 in short bowel syndrome, 1943
- Cholesteryl ester storage disease (CESD), 432e-4t, 432e-5, 2443
- Cholesteryl ester transfer protein (CETP), 2438, 2438f
- Cholesteryl ester transfer protein (CETP) deficiency, 2446
- Cholesteryl ester transfer protein (CETP) inhibitors, 291e-6
- Cholestyramine
 adverse effects of, 269, 2448t
 drug interactions of, 42t
 for irritable bowel syndrome, 1970t
 for lipoprotein disorders, 2448t, 2449
 for metabolic syndrome, 2453
 for pruritus, 2086
 vitamin A absorption and, 96e-6
- Cholic acid, 2076
- Choline, 95e-2t, **96e-5**
- Cholinergic urticaria, 363, 363t, 2118, 2119
- Cholinesterase inhibitors
 for AD, 2601
 adverse effects of, 2601, 2704
 for dementia, 176
 for dementia with Lewy bodies, 2606
 for myasthenia gravis, 2704
- Chondritis, 166e-2, 2203, 2203f
- Chondrocalcinosis, 306, 2235
- Chondrocyte(s), 2227, 2227f, 2233
- Chondrodysplasia(s), **2511**
 classification of, 2511
 diagnosis of, 2512
 genetic considerations in, 2511–2512
 incidence of, 2511
 treatment of, 2512
- Chondrodysplasia punctata, fetal, 47
- Chondroitin, 2232
- Chondromatosis, synovial, 2246–2247
- Chondrosarcoma, 101e-6f, **119e-3**
- Choose My Plate, 95e-5, 95e-6t
- CHOP regimen, 705–706
- Chordoma, 119e-3, 2264
- Chorea(s), **2621**
 clinical features of, 2619t
 disorders associated with, 2622–2623
 Huntington's disease. *See* Huntington's disease
 paraneoplastic, 614t
 Sydenham's, 49, 377e-1, 2151, 2622
 treatment of, 2153
- Chorea-acanthocytosis, 2622
- Chorea gravidarum, 49
- Choriocarcinoma. *See also* Gestational trophoblastic disease
 incidence of, 124e-5
 ovarian, 594–595
 paraneoplastic syndromes in, 609t
 parasellar, 2264
 testicular, 588
- Choriomeningitis, lymphocytic, 837, 908t
- Chorionic villus sampling (CVS), 83e-3–4, 444
- Chorioretinopathy, central serous, 205–206
- Choroid, 40e-1
 gyrate atrophy of, 434e-3t
 mass lesions of, 40e-5–6, 40e-5f
 melanoma of, 206, 206f
- Choroidal artery, occlusion of, 2573–2574, 2575f
- CHRNA4 gene mutations, 2545t
- Chromatin
 in cancer cells, 102e-7–8, 102e-8f, 103e-6
 in DNA, 427, 427f
- Chromatin immunoprecipitation, 455
- Chromium, **96e-10**
 carcinogenicity of, 96e-10, 472e-3, 506
 deficiency of, 96e-9t
 dietary sources of, 96e-10
 functions of, 96e-10
 parental supplementation of, 98e-6t
 recommended intake of, 95e-3t
 supplements, 96e-10
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t, 477t
- Chromobacterium violaceum*, 183e-3
- Chromoblastomycosis, 1355
- Chromogranin
 in carcinoma of unknown primary, 120e-2t, 120e-4
 in GI-NETs, 566
 in lung cancer, 507, 508
 in nonfunctional pNETs, 571
- Chromophobe tumors, renal, 578, 578t
- Chromophores, 386
- Chromosomal instability, 101e-3–4, 101e-5. *See also* Cancer genetics
- Chromosomal mosaicism, 439
- Chromosomal sex, 2349, 2349f
- Chromosome(s)
 acrocentric, 83e-5
 analysis of. *See* Cytogenetic analysis;
 Cytogenomic analysis
 crossing-over and recombination of, 428, 430f
 imprinting in, 430, 432f
 replication of, 426
 staining techniques for, 83e-1
 structure of, 426, 427f
- Chromosome abnormalities/disorders, **83e-1**.
See also Genetic disorders; *specific diseases*
 in cancer, 83e-7, 101e-3t
 contiguous gene syndromes, 432, 437
 copy number variations, 427, 430f
 deletions, 83e-6, 434
 duplications, 83e-6
 imprinting, 83e-7
 insertions, 434
 inversions, 83e-6
 numerical, 83e-1, 83e-4–5
 in radiation-exposed patient, 263e-7
 in sex chromosomes, 2350t. *See also* Disorders of sex development (DSDs)
 single-nucleotide polymorphisms. *See* Single-nucleotide polymorphisms (SNPs)
 structural, 83e-1, 83e-5–7, 83e-5f
 translocation. *See* Translocation, chromosome
 unequal crossing-over, 434
 uniparental disomy, 83e-6–7, 83e-6f
- Chromosome banding, 83e-1
- Chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature (CANDLE), 2215
- Chronic beryllium disease/toxicity, 367, 373e-7, 1691
- Chronic bronchitis, 244, 1692t, 1701
- Chronic Care Model, 12e-3, 12e-3f
- Chronic disease–related malnutrition. *See* Cachexia (chronic disease–related malnutrition)
- Chronic eosinophilic leukemia, 135e-7–8, 135e-7t, 422, 672
- Chronic fatigue syndrome, **464e-1**
 clinical features of, 464e-1, 2199
 definition of, 464e-1
 diagnosis of, 464e-1t, 464e-2, 464e-2f
 EBV infection and, 1188
 epidemiology of, 464e-1
 etiology of, 464e-1, 464e-1t
 vs. Lyme disease, 1152
 pathophysiology of, 464e-1–2
 prognosis of, 464e-3
 treatment of, 464e-2–3
- Chronic granulomatous diseases (CGDs)
 cellular/molecular defects in, 419, 419t, 421, 2106
 clinical features of, 419t, 421, 2106
 complications of, 421
 diagnosis of, 419t
 treatment of, 372e-30, 423, 2106
 X-linked, 375e-1, 2106
- Chronic infantile neurologic cutaneous and articular syndrome (CINCA), 372e-8t
- Chronic inflammatory demyelinating polyneuropathy (CIDP), 2696t, **2698**
- Chronic kidney disease (CKD), **1811**
 in amyloidosis, 719, 721. *See also* Amyloidosis
 approach to the patient
 establishment of diagnosis and etiology, 1820
 history and physical examination, 1819
 imaging, 1820
 laboratory studies, 1813, 1813t, 1819–1820, 2768–2769t
 renal biopsy, 1820
 staging, 1812–1813
 calcium and phosphate metabolism disorders in
 bone manifestations of, 1815
 calciphylaxis, 1816, 1816f
 cardiovascular manifestations of, 1815
 hyperphosphatemia, 2460, 2485
 hypocalcemia, 333e-3, 462e-19, 2484–2485

- pathophysiology of, 333e-3, 333e-3f
 potassium regulation, 333e-2-3, 1814
 secondary hyperparathyroidism, 1815, 2478-2479
 treatment of, 1816, 2478-2479, 2485
 tumoral calcinosis, 1815, 1815f
- cardiovascular disorders in
 epidemiology of, 1816, 1816f
 gangrenous calcification, 462e-19
 heart failure, 1817
 hypertension, 201f, 1615, 1617, 1817
 ischemic vascular disease, 1816-1817
 left ventricular hypertrophy, 1817
 pericardial disease, 1575, 1817-1818
 risk factors for, 1825
 treatment of, 1817
- classification of, 1811, 1812f
 clinical features of, 289t
 contrast agent use in, 1461
 deaths from, 6e-1t, 11t, 56t
 diagnosis of, 289t
 drug concentration and response and, 37
 endocrine-metabolic disturbances in, 1819, 2366
 epidemiology of, 1813
 etiology of, 1813, 1813t
 diabetes mellitus. *See* Diabetic nephropathy
 glomerulonephritis, 1813. *See also* Glomerulonephritis
 urinary tract obstruction, 1873
- in Fabry's disease, 432e-1
 fluid, electrolyte, and acid-base disorders in
 acid-base regulation, 333e-3
 edema, 252, 252t
 hyperkalemia, 64e-2, 310
 hypokalemia, 307, 1814
 hyponatremia, 299, 1814
 metabolic acidosis, 318, 320, 1814
 sodium regulation, 333e-3, 1814
 treatment of, 320, 1815
 water regulation, 1814
- gastrointestinal and nutritional abnormalities in, 1819
 genetic considerations in, 332e-7t, 1811-1812
 global considerations in, 1821, 1822, 1825
 glomerular filtration rate in, 290-291, 1813t
 hematologic abnormalities in
 abnormal hemostasis, 1818
 bleeding, 404
 etiology of, 1818, 1818t
 hypoproliferative anemia, 398, 630, 630t
 high-altitude travel and, 476e-5-6
 immunization in, 788f
 lipoprotein disorders in, 2441
 in multiple myeloma, 714
 neuromuscular disorders in, 1818-1819
 back pain, 118
 coma, 1773
 delirium, 169
 fatigue, 152
 gait disorders, 164
 myopathy, 462e-19
 neuropathy, 1819, 2683-2684
 pathophysiology of, 1811, 1813f
 counterbalance, 333e-2
 inflammation, 1814
 nephron hyperfunction, 333e-1, 333e-1f, 1812f
 tubular hypertrophy, 333e-1, 333e-1f
 uremic syndrome, 1811, 1813, 1814
- in pregnancy, 46, 1819
 renal response to, 333e-2-3, 333e-2f, 333e-2t, 1812f
 risk factors for, 1811-1812
 in Sjögren's syndrome, 64e-7
 skin manifestations of, 362, 363, 462e-19, 1819
 in SLE. *See* Lupus nephritis
 stage 5 (end-stage renal disease)
 cardiovascular disease and, 1825
 in HIV infection, 1259
 hypertension in, 1847-1848
 incidence of, 1822
- treatment of, 1822. *See also* Hemodialysis; Kidney transplantation; Peritoneal dialysis
 uremic syndrome in, 1811, 1813
- treatment of
 antihypertensive therapy, 1820-1821
 diabetic nephropathy. *See* Diabetic nephropathy
 erythropoietin, 631
 medication dose adjustment, 1821
 patient education, 1821
 preparation for renal replacement therapy, 1821. *See also* Hemodialysis; Kidney transplantation; Peritoneal dialysis
 specialized nutritional support, 98e-4
 timing of renal replacement therapy, 1821
 tuberculous, 1110, 1110f, 1119-1120
- Chronic lymphoid leukemia. *See* B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma
- Chronic mountain sickness, 248, 476e-5
- Chronic myeloid leukemia (CML), **687**
 accelerated blastic phase of, 690
 atypical (*BCR-ABL1* negative), 135e-5-6, 135e-5t
 blastic phase of, 690
 cancer stem cells in, 102e-11
 chromosome abnormalities in, 687-689, 688f
 amplifications, 101e-2t
 translocations, 83e-7, 101e-3, 101e-3t, 101e-4f, 672, 687-688
 clinical features of, 689, 689t
 diagnosis of, 689-690
 epidemiology of, 687
 etiology of, 687
 global considerations in, 694-695, 694f
 incidence of, 687
 pathophysiology of, 687-689
 peripheral blood smear in, 81e-10f
 prognosis of, 690-691, 690f
 skin manifestations of, 369
 splenomegaly in, 411
 treatment of, **691**, 693t
 in accelerated or blastic phases, 693-694
 chemotherapy, 694
 goals for, 692
 HSCT, 139e-5t, 139e-6, 692-693, 693t
 interferon, 691, 694
 leukapheresis, 694
 monitoring, 693
 in pregnancy, 694
 splenectomy, 412, 694
 tyrosine kinase inhibitors, 101e-3, 102e-3-4, 102e-5t, 103e-16, 691-692, 691t, 693t
- Chronic myelomonocytic leukemia (CMML), 102e-3, 135e-5t, 135e-6, 669
- Chronic neutrophilic leukemia, 135e-5-6, 135e-6t, 672
- Chronic nonspherocytic hemolytic anemia, 656
- Chronic obstructive pulmonary disease (COPD)
 vs. asthma, 1676
 cor pulmonale in, 1506, 1704. *See also* Cor pulmonale
 CT in, 1705, 1705f
 deaths from, 1700
 definition of, 1700
 dyspnea in, 47e-1t, 47e-2, 47e-2t, 47e-3, 1704
 ECG in, 269e-12f, 278e-5f
 exacerbations of, 1706-1707
H. influenzae infections in, 1011
M. catarrhalis infections in, 1013, 1014f
 mechanical ventilation for, 1707
 noninvasive ventilation for, 1707, 1740
 genetic considerations in, 1703
 history in, 1704
 incidence of, 1700
 laboratory findings in, 1704-1705, 1704t
 natural history of, 1703-1704, 1704f
 pathogenesis of, 1700-1701, 1700f
 pathology of, 1701
- pathophysiology of, 41-42
 physical findings in, 1704
 risk factors for, 1702-1703, 1702f
 severity of, GOLD criteria for, 1704t
 in travelers, 799
 treatment of
 lung transplantation, 320e-1t, 1706
 lung volume reduction surgery, 1706
 pharmacotherapy, 1705-1706
 pulmonary rehabilitation, 1706
 smoking cessation, 1704
- Chronic pancreatitis, **2098**
 approach to the patient, 2088f
 clinical features of, 269, 284, 366, 2099-2100
 cobalamin absorption in, 350e-1, 350e-1t
 complications of, 2100, 2100t
 diagnosis of
 approach to, 2086, 2088f
 endoscopic, 1900
 imaging, 2087t, 2088-2089, 2089t
 imaging in, 2099-2100, 2100f
 Schilling test, 1941t
 epidemiology of, 2086
 etiology of, 2098-2099, 2098t
 genetic considerations in, 2098-2099
 pathophysiology of, 2098
 treatment of, 2100-2101, 2101t
- Chronic progressive external ophthalmoplegia. *See* Progressive external ophthalmoplegia
- Chronic Respiratory Disease Questionnaire, 47e-2
- Chronic traumatic encephalopathy, 173, 457e-4, 471e-2, 2608
- Chronic venous disease, **1650**
 anatomic considerations, 1650
 classification of, 1651, 1652t
 clinical features of, 1651, 1651f
 diagnosis of, 1651-1652
 differential diagnosis of, 1651
 epidemiology of, 1650
 pathophysiology of, 1650-1651
 treatment of, 1651-1652
- Chronic wasting disease (CWD), 453e-1t, 453e-2
- Chronotropic incompetence, 1468
- Chryseobacterium indologenes* infections, 183e-2t, 183e-3
- Chryseobacterium meningosepticum* infections. *See* *Elizabethkingia meningoseptica* infections
- Chrysiasis, 361
- Churg-Strauss syndrome. *See* Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
- Chvostek's sign, 315
- Chylomicron(s), 1241t, 1936, 2437, 2437f
- Chylomicronemia syndromes, familial, 2439-2440, 2440t
- Chylothorax, 1718
- Ciclopirox, 347, 1358
- Cidofovir
 action of, 215e-7, 897
 for adenovirus, 1208
 adverse effects of, 897, 1804
 for CMV infection, 215e-2t, 215e-7, 922, 1194, 2657
 for CMV infection prophylaxis in HIV infection, 1252t
 for HHV-6 infections, 922
 for HSV infections, 1182-1183
 for molluscum contagiosum, 220e-1
 pharmacology of, 215e-7
 for smallpox, 261e-6
 for viral encephalitis, 897
- CIDP (chronic inflammatory demyelinating polyneuropathy), 2696t, **2698**
- Cigarette smoking. *See* Smoking
- Cigar smoking, 476
- Ciguatera poisoning, 267, 2742, 2742t
- CIITA, 373e-3
- Cilansetron, 1970
- Cilium, 1851f
- Cilostazol, 1645

- Cimetidine
 action of, 332e-8
 adverse effects of
 arthralgias, 2218t
 cholestatic hepatitis, 284
 erectile dysfunction, 1920
 myopathy, 462e-20
 SA node dysfunction, 1467t
 thrombocytopenia, 727t
 for carcinoid syndrome, 566
 drug interactions of, 42t
 altretamine, 103e-9t
 chloroquine, 246e-1t
 macrolides, 939t
 mebendazole, 246e-3t
 metronidazole, 246e-5t
 PDE-5 inhibitors, 328
 phenytoin, 1920
 praziquantel, 246e-12
 quinidine, 246e-6t
 quinine, 246e-13
 theophylline, 1920
 warfarin, 1920
 for gastrinoma, 2337
 for GERD, 263
 for peptic ulcer disease, 1920, 1920t
 as premedication for antivenom, 2737
 as premedication for taxanes, 103e-15
- Cinacalcet, 1816
- CINCA (chronic infantile neurologic cutaneous and articular syndrome), 372e-8t
- Cingulate cortex, 180
- Cinnarizine, 2611
- Ciprofloxacin
 action of, 931t, 932-933, 943
 adverse effects of, 203, 246e-6t, 943, 2023, 2025t
 for *Aeromonas* infections, 183e-2, 183e-2t
 for anthrax, 261e-3, 261e-5t
 for *Bartonella* infections, 1081t
 for bite-wound infections, 167e-3, 167e-3t, 832t
 for bronchiectasis prophylaxis, 1696
 for burn-wound infections, 166e-3
 for *Campylobacter* infections, 1061
 for chancroid, 880t, 1013
 for cholera, 1064
 for clostridial infections, 991t
 for community-acquired pneumonia, 767t, 808t
 drug interactions of, 246e-6t
 for enteric fever, 1052t
 for febrile neutropenia, 103e-23
 for gonococcal arthritis, 835
 gut microbiota and, 86e-7
 for health care-associated pneumonia, 812t
 indications for, 246e-6t, 941t, 943
 for infections in cancer patient, 491
 for infectious diarrhea, 268, 857t
 for intraabdominal infections, 767t
 for *Legionella* infections, 1020t
 for necrotizing fasciitis, 832t
 for nongonococcal bacterial arthritis, 835
 for nontyphoidal *Salmonella* infections, 1054t
 for osteomyelitis, 840t, 841t
 for otitis externa, 229
 for *P. aeruginosa* infections, 808t, 1045t
 for perichondritis, 228
 for peritonitis prophylaxis, 847
 for plague prophylaxis, 1075t
 for plague treatment, 261e-5, 261e-5t, 1075
 for pouchitis in IBD, 1960
 for pyelonephritis, 867
 for Q fever, 1162
 resistance to
 in *Campylobacter* spp., 1061
 in *E. coli*, 862
 mechanisms of, 931t
 rates of, 941t
 in *S. pneumoniae*, 807
 for *Salmonella* infection prophylaxis, in HIV infection, 1252t
 for sepsis/septic shock, 1757t
 for *Shigella* infections, 1058, 1058t
 for tularemia, 261e-5t, 261e-7
- Circadian cycle, 400e-6
- Circadian pacemaker, 193
- Circadian rhythmicity, 186-187, 194, 2310, 2310f
- Circadian rhythm sleep disorders, 193-194
- CIRCI (critical illness-related corticosteroid deficiency), 1755, 1757
- Circinate balanitis, 2174
- Circle of Willis aneurysm, 1525
- Circulating endothelial precursors, 102e-14f
- Circulatory integrity, maintenance of, 296-297, 297f
- Circumcision, HIV infection risk and, 1221, 1285
- Cirrhosis
 alcoholic, 2058-2059, 2059f, 2075. *See also* Alcoholic liver disease
 in α_1 -AT deficiency, 2062
 autoimmune hepatitis and, 2060
 biliary. *See* Biliary cirrhosis
 cardiac, 2062
 clinical features of, 2058
 ascites, 286f, 287
 clubbing, 250
 cyanosis, 249
 diarrhea, 269
 edema, 252, 252t
 esophageal varices, 277
 gynecomastia, 2366
 hyponatremia, 299
 jaundice, 284
 pleural effusion, 1717
 splenomegaly, 411
 testicular atrophy, 2366
 coagulation disorders in, 739
 complications of, 2062, 2062t
 acute kidney injury, 1800, 1809-1810
 ascites. *See* Ascites
 bone disease, 2067
 coagulopathy, 2067
 encephalopathy. *See* Encephalopathy, hepatic
 hematologic abnormalities, 2067
 hepatorenal syndrome, 1800, 1809-1810, 2066
 hypersplenism, 2064
 malnutrition, 2066
 portal hypertension. *See* Portal hypertension
 splenomegaly, 2064
 spontaneous bacterial peritonitis, 2065-2066
 cryptogenic, vs. autoimmune hepatitis, 2051
 in cystic fibrosis, 2062
 definition of, 2031, 2058
 etiology of, 2058, 2059t
 global considerations in, 2059-2060
 HBV infection and, 2031, 2060
 HCV infection and, 2041, 2059-2060
 in hematocromatosis, 2062
 hepatocellular carcinoma and, 545, 545t
 Laennec's, 282
 liver biopsy in, 366e-3f, 366e-4f
 liver transplantation for, 2068, 2068t
 in NAFLD/NASH, 2054-2055, 2060
 pathogenesis of, 286
 pathology of, 2058
 physical examination in, 282-283
 postnecrosis, vs. autoimmune hepatitis, 2051
 recurrent, after liver transplantation, 2073-2074
 staging of, 1994-1995, 1995t
 treatment of, 1995, 2060
 tumor markers in, 473t
 in Wilson's disease, 2062
- Cisapride, 42t
- Cisplatin
 action of, 103e-12
 adverse effects of, 103e-9t, 103e-12, 590
 cardiotoxicity, 1563
 electrolyte disturbances, 64e-8, 103e-12, 306
 gonadal dysfunction, 103e-25
 hearing loss, 219, 622
 hemolytic-uremic syndrome, 1796
 nausea and vomiting, 103e-12, 103e-25, 474
- nephrotoxicity, 1804
 neurologic, 607, 607t, 622, 1793, 2686t
 neutropenic enterocolitis, 1798
 Raynaud's phenomenon, 103e-12, 1649
 for bladder cancer, 577
 for carcinoma of unknown primary, 120e-4
 for cervical cancer, 596
 for cholangiocarcinoma, 553
 for gastric cancer, 535
 for gestational trophoblastic disease, 597
 for head and neck cancer, 505
 for hepatocellular carcinoma, 550
 for lung cancer, 517-518, 518t, 520t, 521t, 522
 for melanoma, 499
 for osteosarcoma, 119e-3
 for ovarian germ cell tumors, 594
 for salivary gland tumor, 506
 for squamous cell carcinoma, 502
 for testicular cancer, 590, 591
 for thymoma, 123e-2
- Cisternography, CT, 440e-2
- Citalopram
 adverse effects of, 2640t, 2710t, 2716
 for depression, 64, 2710t, 2716
 for irritable bowel syndrome, 1969
- Citrate
 in transfusions, 138e-5
 urinary, 1867, 1870
- Citrobacter* spp. infections, 486t, 862, 1034-1035
- Citrulline, 435e-2
- Citrullinemia, 545, 545t
- Citrullinemia type 2 (citrin deficiency), 434e-3t, 435e-1t, 435e-2
- CJD. *See* Creutzfeldt-Jakob disease (CJD)
- CK. *See* Creatine kinase (CK)
- CKD. *See* Chronic kidney disease (CKD)
- c-kit*, 2114, 2115-2116
- c-kit* mutations
 in gastrointestinal stromal tumor, 119e-1
 imatinib response and, 39t, 536
 in melanoma, 499
 in ovarian germ cell tumors, 594
 in thymoma, 123e-2
- Cladophialophora*, 1355
- Cladribine, 135e-2, 682, 706, 718, 2121
- CLAH (congenital lipoid adrenal hyperplasia), 2324t
- Clarithromycin
 action of, 205e-8, 931, 931t, 943
 adverse effects of, 205e-8, 943
 for *Bartonella* infection prophylaxis in HIV infection, 1252t
 for community-acquired pneumonia, 808t
 drug interactions of, 42t, 205e-8, 246e-3t, 943, 2235
 for endocarditis prophylaxis, 826t
 in *H. pylori* eradication, 1042t, 1922t, 1923
 indications for, 940t, 943
 for *Legionella* infections, 1020t
 for MAC infection prophylaxis in HIV infection, 1251t
 for nontuberculous mycobacterial infections, 205e-8
 for pertussis, 1024t
 for pneumonia, 953
 in pregnancy and lactation, 937t
 resistance to, 931t, 940t, 1042
- Clasp-knife phenomenon, 154
- Classism, 16e-4
- Class switch, 710
- Claude's syndrome, 208, 2575, 2575f
- Claudication
 in aortic occlusion, 1642
 intermittent, 291e-4, 1616, 1643. *See also* Peripheral artery disease
 neurogenic, 115
 physical examination in, 1446, 1446f
 in thromboangiitis obliterans, 1645
- Claustrophobia, 2712
- Clavulanate, 933
- Clavulanic acid, 284
- CLCN5* gene mutations, 332e-8, 2459
- CLDN16* gene mutations, 332e-8
- Clearance, of drugs, 35

- Clear cell carcinoma, renal, 578, 578t. *See also* Renal cell carcinoma (RCC)
- Clear liquid diet, 95e-5
- Cleidocranial dysplasia, 2455
- Clenched-fist injury, 167e-2, 167e-3t
- CLIA (Clinical Laboratory Improvement Amendments Act), 480e-6
- Climate change, **151e-1**
 adaptation to, 151e-9
 arbovirus infections and, 151e-4–5
 average global temperatures and, 151e-1–2
 El Niño events and, 151e-4, 151e-4f, 151e-8, 151e-8f
 greenhouse gases and, 151e-1, 151e-2f, 151e-2t
 hurricane severity and, 151e-4
 Lyme disease and, 151e-5–6, 151e-6f
 mitigation of, 151e-9
 overview of, 151e-1, 151e-1f
 population displacement and, 151e-8
 population migration and conflict and, 151e-3–4
 precipitation patterns and, 151e-3–4
 sea levels and, 151e-4
 vector-borne disease and, 151e-4–6, 151e-5f, 151e-6f
 waterborne disease and, 151e-6–8, 151e-7f
- Clindamycin
 for acne vulgaris, 352
 for actinomycosis, 1091t
 action of, 931, 931t, 943
 adverse effects of, 246e-3t, 938t, 943
 for anaerobic bacterial infections, 1101, 1102
 for anthrax, 261e-3, 261e-5t
 for asymptomatic pharyngeal GAS carriage, 966
 for *Babesia* infections, 780t, 1386–1387, 1386t
 for bacterial vaginosis, 873t, 875
 for bite-wound infections, 167e-2, 167e-3t, 832t
 for *Capnocytophaga* infections, 183e-3
 for cellulitis, 832, 832t
 for clostridial infections, 991t, 994
 for clostridial myonecrosis, 780t
 for CNS toxoplasmosis, 903
 for deep neck infections, 235
 for endocarditis prophylaxis, 826t
 for gas gangrene, 832t, 991t
 indications for, 246e-3t, 940t, 943
 for Lemierre's disease, 234
 for lung abscess, 815
 for *M. hominis* infections, 926
 for malaria treatment, 1378t
 for MRSA infections, 345, 962t
 for necrotizing fasciitis, 780t, 832t
 for osteomyelitis, 840t, 841t
 for otitis media, 229t
 for PCP, 1361t
 for pelvic inflammatory disease, 879
 for pharyngitis, 232f
 for pneumococcal pneumonia, 953
 in pregnancy and lactation, 246e-3t, 937t
 prophylactic, 945t
 resistance to
 in anaerobic bacteria, 1101
 in group A streptococci, 969
 inducible, 345
 mechanisms of, 931t, 934
 prevalence of, 940t
 for sepsis/septic shock, 1757t
 for sinusitis, 227t
 for skin and soft tissue infections, 768t, 832, 962t
 for streptococcus group B, 50
 for toxic shock syndrome, 780t, 963, 965t, 969
 for toxoplasmosis prophylaxis in HIV infection, 1251t
 for Vincent's angina, 233
- Clinical breast examination, 480, 481t, 524
- Clinical decision-making, 3, 16e-4. *See also* Clinical reasoning
- Clinical Evidence (directory), 25t
- Clinical Laboratory Improvement Amendments Act (CLIA), 480e-6
- Clinical laboratory tests, **480e-1**
 accuracy of, 21, 21t
 analytic sensitivity of, 480e-2
 analytic specificity of, 480e-2
 batteries of, 2
 critical values, 480e-2, 480e-2t
 diagnostic principles of, 480e-2–3
 for disease and treatment monitoring, 480e-1
 for disease diagnosis, 480e-1
 for disease screening, 480e-1
 error sources in, 480e-4–5, 480e-4t
 genetic. *See* Genetic testing
 home, 480e-5
 physician's evaluation and use of, 2–3
 point-of-care, 480e-5
 predictive value of, 22, 479, 479t
 receiver operative characteristic curves for, 21, 22f
 reference values for, 480e-3–4, 480e-3f
 cerebrospinal fluid, 2766t
 cholesterol, 480e-3, 480e-4f
 clinical chemistry and immunology, 2757–2764t
 hematology and coagulation, 2754–2756t
 renal function, 480e-3–4, 480e-4f
 toxicology and therapeutic drug monitoring, 2764–2766t
 urine analysis and renal function, 2768–2769t
 vitamins and trace minerals, 2766t
 regulation of laboratories performing, 480e-6
 for risk assessment, 480e-1
 sensitivity of, 21, 21t, 479, 479t
 specificity of, 21, 21t, 479, 479t
 "stat" orders for, 480e-2–3
- Clinical microbiology laboratory, 150e-1. *See also* Infectious diseases, laboratory diagnosis of; Virus(es), laboratory identification of
- Clinical practice guidelines, 3, 17e-3, 26
- Clinical Pulmonary Infection Score (CPIS), 811, 811t. *See also* Ventilator-associated pneumonia (VAP)
- Clinical reasoning, **18**
 clinical expertise in, 18
 cognitive shortcuts in, 19
 decision support tools, 24
 diagnostic test interpretation and, 21–24. *See also* Clinical laboratory tests
 diagnostic vs. therapeutic, 20
 economic incentives and, 21
 evidence-based medicine and, 24–26
 hypothetico-deductive model of, 20
 influences on, 20–21
 intuitive vs. analytic, 18–20
 pattern recognition in, 18–19
 practice setting and, 21
 practice style and, 20–21
- Clinical research. *See* Research
- Clinical skills, 1–3
- Clinical trials, 25–26, 43, 2025
- Clinical uncertainty, 16e-4
- Clinton Foundation HIV/AIDS Initiative, 12
- CLIP staging, of hepatocellular carcinoma, 546, 546t
- Clitoral engorgement, 330
- Clitoral vacuum device, 331
- CLL. *See* B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma
- Clobazam, 2553t
- Clobetasol, 347
- Clofarabine, 103e-11t, 683
- Clofazimine
 action of, 205e-7
 adverse effects of, 205e-7, 361, 379, 1126, 1263
 for erythema nodosum leprosum, 1128
 for leprosy, 1126–1127, 1127t
 for TB, 205e-7
- Clofibrate, 2199, 2218t
- Clomiphene citrate, 7e-4, 337, 2389
- Clomipramine, 190, 2710t, 2714
- Clonal eosinophilia, 135e-7
- Clonality, of stem cells, 454
- Clonazepam
 adverse effects of, 2553t
 for anxiety disorders, 2711t
 for ataxia/tremor, 2672
 for dyspnea, 62t
 for epilepsy, 2553t
 for glossodynia, 239
 for myoclonus, 2623
 overdose/poisoning with, 473e-10t, 2764t
 for pain, 91t
 for panic disorder, 2709
 pharmacology of, 2711t
 for posthypoxic myoclonus, 1782
 for REM sleep behavior disorder, 193
 for sleep disorders in PD, 2618
 for status epilepticus, 2558f
 for tardive dyskinesia, 2624
 therapeutic monitoring of, 2764t
 for vertigo, 151t
- Clonidine
 adverse effects of, 253t, 1467t, 2640t
 for ascites, 288
 for diabetic diarrhea, 272
 drug interactions of, 41
 for hot flashes, 623
 for hypertension, 1624t
 for hypertensive emergencies, 1627
 for inappropriate sinus tachycardia, 1477
 for menopausal symptoms, 2383
 for opioid withdrawal syndrome, 468e-3
 overdose/poisoning with, 473e-9t
 for pain, 61
 perioperative, 54
 for smoking cessation therapy, 507, 2732
 for Tourette's syndrome, 2623
- Clonidine suppression test, 2330
- Cloning, 441, 454
- Clonorchis sinensis*/clonorchiasis. *See also* Biliary (hepatic) flukes
 clinical features of, 1428
 diagnosis of, 245e-2t, 245e-5t, 245e-6t, 765t
 epidemiology of, 245e-2t, 765t, 1428
 life-cycle hosts of, 245e-2t
- Clopidogrel
 action of, 746f, 747
 adverse effects of, 404, 729, 747, 1864
 with aspirin, 747, 2570
 for cardiac catheterization pretreatment, 1461
 dosage of, 747
 drug interactions of, 205e-3, 1921
 genetic variations in response to, 39t, 40, 43, 277, 277t, 742–743, 743t, 1596
 indications for, 747l
 for ischemic heart disease, 1590
 management before endoscopic procedures, 1889t
 management before lumbar puncture, 443e-1
 metabolism of, 34t, 35
 for myocardial infarction
 NSTEMI-ACS, 1596, 1598, 1598t
 as secondary prevention, 1610
 STEMI, 1606
 in PCI, 296e-1, 296e-2, 1596
 in peripheral artery disease, 1644
 resistance to, 747
 for stroke/TIA prevention, 2569–2570
- Clorazepate, 2711t
- Closed-loop obstruction, 1982. *See also* Intestinal obstruction
- Clostridial myonecrosis. *See* Gas gangrene (clostridial myonecrosis)
- Clostridium* spp., 990, 990f, 1096
- Clostridium baratii*, 987
- Clostridium botulinum*, 261e-8, 987. *See also* Botulism
- Clostridium butyricum*, 987
- Clostridium difficile*, 145e-9, 857–858
- Clostridium difficile*-associated disease
 antibiotics associated with, 858
 in cancer, 488, 1798
 clinical features of, 266t, 859
 diagnosis of, 770–771, 856, 859, 859t
 with enteral feeding, 98e-9
 epidemiology of, 855, 857–858

- Clostridium difficile*–associated disease (Cont.):
 etiology of, 857–858
 fulminant, 861
 global considerations in, 859
 health care–associated, 266, 913, 917
 pathogenesis of, 858–859, 858f
 PPIs and, 858, 1921
 prevention and control of, 861, 913, 946
 prognosis of, 861
 recurrent, 860–861
 in transplant recipient, 919, 920t, 924t
 treatment of, 86e-10, 859–860, 860t
- Clostridium* spp. infections
 bacteremia, 992, 1100
 cardiac, 1561
 enteric, 991
 epidemiology of, 990
 global considerations in, 990
 vs. IBD, 1956
 polymicrobial infections involving, 990–991, 991t
 skin and soft tissue, 992. *See also* Gas gangrene (clostridial myonecrosis)
 surgical-site, 915
 treatment of, 991t
- Clostridium perfringens* infections
 bacteremia, 992
 in cancer patient, 490
 clinical features of, 266t
 food-borne, 856, 856t, 991
 gas gangrene. *See* Gas gangrene (clostridial myonecrosis)
 necrotizing enterocolitis, 991
 necrotizing fasciitis, 783. *See also* Necrotizing fasciitis
- Clostridium septicum* infections
 bacteremia, 992
 gas gangrene. *See* Gas gangrene (clostridial myonecrosis)
 intraabdominal, 1099
 toxins in, 993
- Clostridium sordellii* infections, 782, 831, 994–995
- Clostridium tertium* infections, 992, 994
- Clostridium tetani*, 984–985. *See also* Tetanus
- Clotrimazole, 233, 349, 873t, 875
- Clotting factors. *See also specific factors*
 defective, 367
 laboratory evaluation of, 733f
 in normal hemostasis, 401, 401f
 specific factor assays, 406
- Cloxacillin, 937t
- Clozapine
 adverse effects of, 2617, 2721, 2722, 2722t
 overdose/poisoning with, 473e-9t, 2764t
 for psychosis in Huntington's disease, 2622
 for psychosis in PD, 2617
 for schizophrenia, 2721, 2722t
 therapeutic monitoring of, 2764t
- CLRs (C-type lectin receptors), 372e-5t
- Clubbing, of digits, **250**
 in cardiovascular disease, 1443, 2245
 disorders associated with, 250, 2245, 2245t
 in Graves' disease, 2295, 2295f
 in hypertrophic osteoarthropathy, 250, 426e-7, 2244, 2244f, 2245
 in lung cancer, 246, 511
 in patent ductus arteriosus, 1524f
 in respiratory disease, 47e-3, 1662
 in thyroid disease, 2245
- “Club drugs,” 469e-5
- Clue cells, 873, 874, 875f
- Cluster headache, **2594**
 clinical features of, 2594–2595, 2595t
 imaging in, 2590f
 incidence of, 107t
 prevention of, 2595–2596, 2595t
 treatment of, 2595, 2595t
- Cluster of differentiation (CD) classification, 372e-1, 372e-3–4t
- Cluster-tic, 2646
- Clutton's joints, 836, 1137
- CMDs (congenital muscular dystrophies), 462e-9, 462e-10t
- CML. *See* Chronic myeloid leukemia (CML)
- CMML (chronic myelomonocytic leukemia), 102e-3, 135e-5t, 135e-6, 669
- CMR. *See* Cardiac magnetic resonance imaging (CMR)
- CMS (congenital myasthenic syndromes), 2702, 2703t
- CMT disease. *See* Charcot-Marie-Tooth (CMT) disease
- CMV. *See* Cytomegalovirus (CMV)
- CMV infections. *See* Cytomegalovirus (CMV) infections
- CMX-001 (brincidofovir), 215e-7, 261e-6
- c-myc* gene, 706
- cMYC* gene mutations, 101e-2t, 2305t
- Cnidae (cnidocyst) envenomation, 2738–2739
- CNS. *See* Central nervous system (CNS)
- Co-activators, 428
- Coagulase-negative staphylococci (CoNS), 954, 960, 961. *See also* Staphylococcal infections
- Coagulation
 fibrin clot formation in, 401, 401f
 platelet plug formation in, 400–401, 725
 in sepsis/septic shock, 1753
 in shock, 1746
- Coagulation cascade
 extrinsic pathway, 401, 733f
 intrinsic pathway, 401, 401f, 733f
 overview, 744f
 overview of, 743–744
 red blood cells fate in, 661f
- Coagulation disorders, **732**
 acquired factor inhibitors and, 740
 atherosclerosis and, 291e-8–9
 in chronic kidney disease, 1818
 in cirrhosis, 2067
 disseminated intravascular coagulation. *See* Disseminated intravascular coagulation (DIC)
 familial multiple coagulation deficiencies, 736
 genetic characteristics of, 733t
 in heatstroke, 479e-4
 hemophilia. *See* Hemophilia
 in hypothermia, 478e-2
 laboratory evaluation of, 733f, 733t, 2754–2756t
 in liver failure, 739, 739t
 in vitamin K deficiency. *See* Vitamin K deficiency
- Coagulation factor(s)
 in hepatobiliary diseases, 1998, 1998t
 multiple deficiencies of, 736
 in sepsis/septic shock, 1753
 vitamin K–dependent, 736, 736f
- Coagulation factor inhibitors, 732, 734–735, 740
- Coagulation tests, 405–406, 406t, 2754–2756t
- Coal dust, 1688t
- Coal tar, 354t
- Coal worker's pneumoconiosis, 1690
- Coarctation of the aorta, **1525**, 1620
- Coats' disease, 462e-11
- Cobalamin (vitamin B₁₂), **640**
 absorption of, 350e-1, 640, 645
 deficiency of. *See* Cobalamin (vitamin B₁₂) deficiency
 dietary sources of, 640, 644
 folate and, 641
 functions of, 96e-2f, 640
 for hypogammaglobulinemia, 375e-2
 malabsorption of, 216, 350e-1, 350e-1t, 644–645
 metabolism of, disorders of, 646
 parenteral requirement for, 98e-5t
 postgastroctomy levels of, 1927
 reference range for, 2766t
 requirement/recommended intake of, 95e-2t, 640
 serum, 647
 structure of, 96e-2f
 transport in blood, 640
- Cobalamin (vitamin B₁₂) deficiency. *See also* Megaloblastic anemia
 bacterial overgrowth syndromes in, 1944.
See also Bacterial overgrowth syndromes
 clinical features of, 96e-1t, 360, 463t, 642–643, 2607
 anemia, 398
 cardiovascular, 290e-2, 643
 dementia, 2607
 frontal lobe syndrome, 184
 gait disorders, 163
 myelopathy, 2607, 2659
 neuropathies in, 2690
 olfactory dysfunction, 216
 optic neuropathy, 203
 peripheral neuropathy, 643
 in *D. latum* infections, 1434
 diagnosis of, 647, 2690
 etiology of, 644–646, 644t, 2607, 2689–2690
 hematologic findings in, 81e-3f, 643–644, 644f
 in HIV infection, 1262
 in infants, 644, 646
 postgastroctomy, 645
 PPIs and, 568
 in pregnancy, 642
 risk factors for, 96e-1t
 small-intestinal mucosal biopsies in, 1940t
 treatment of, 648, 2607, 2690
- Cobalt exposure, 263e-3t, 472e-3, 1691
- Cobblestone papillae, 199
- Cobras, 2734, 2738. *See also* Snakebites
- Cocaine, **469e-1**
 action and effects of, 444e-4, 465e-5t, 469e-1
 adverse effects of, **469e-1**
 autonomic overactivity, 2643
 cardiomyopathy, 1562
 erectile dysfunction, 327t
 hypertension, 1617t
 intracranial hemorrhage, 2568, 2584
 myopathy, 462e-20, 462e-20t, 2218t
 nasal and auricular, 2184, 2204
 palpitations, 254
 vasculitis, 2218t
 overdose and abuse of
 clinical features of, 473e-3, 473e-8t, 473e-13t
 laboratory evaluation of, 2764t
 prevalence of, 469e-1
 treatment of, 469e-2, 473e-8t, 473e-13t
- Coccidioides* spp., 1329–1330, 1334, 1335f, 1336
- Coccidioides* spp. infections, **1334**
 arthritis, 836
 chronic meningitis, 441e-7f, 908t
 clinical features of, 1335–1336, 1336t
 complications of, 1336
 diagnosis of, 765t, 1336
 disseminated, 1335–1336
 epidemiology of, 1334–1335
 in HIV infection, 1252t, 1255
 pathogenesis of, 1335
 prevention of, 1337
 skin manifestations of, 366, 829, 1335
 subacute meningitis, 898–899
 treatment of, 1336–1337, 1336t
- Cochlea, 217, 218f
- Cochlear implant, 219, 223–224, 224f
- Cochrane Collaboration Complementary Medicine Reviews, 14e-4t
- Cochrane Database of Systematic Reviews, 25, 25t
- Cockayne syndrome, 94e-4
- Cockcroft-Gault equation, 291
- Codeine
 for diarrhea, 272
 drug interactions of, 2711t
 for dyspnea, 62t
 genetic variations in response to, 39t, 40
 metabolism of, 34t, 35
 for pain, 60, 91t
- Codman's triangle, 119e-3
- Codons, 426
- Coenurosis, 1435
- Coenzyme Q10, 2610, 2766t

- Coeur en sabot*, 1527
 Coffee worker's lung, 1681t
 Cogan's syndrome, 1638, 2192, 2204
 Cognitive behavioral therapy (CBT)
 for alcohol use disorder, 2728
 for back pain, 120
 for chronic fatigue syndrome, 464e-2-3
 for depression, 2717
 for fatigue, 153
 for female sexual dysfunction, 331
 for fibromyalgia, 2240
 for glossodynia, 239
 for insomnia, 191
 for metabolic syndrome, 2453
 for obesity, 2396
 for PTSD, 471e-5
 Cognitive dysfunction
 alcohol use and, 2724
 assessment of, 37e-1v
 caring for patient with, 184
 delirium and, 166
 hypertension-related, 1615
 in MS, 2663
 treatment of, 444e-11f
 Cognitive evoked potentials, 442e-3
 Cognitive function
 assessment of, 167
 postmenopausal hormone therapy and, 6e-2, 2384t, 2385-2386
 Cognitive rehabilitation procedures, 184
 Cogwheel rigidity, 2538
 Cohen's syndrome, 415e-5t
COL1A1/2 gene mutations
 in chondrodysplasias, 2511
 in Ehlers-Danlos syndrome, 2510t, 2511
 in osteogenesis imperfecta, 435, 2454, 2508t, 2509
COL4A4/5 gene mutations, 2514
COL17A1 gene mutations, 434
 Colchicine
 for acute idiopathic pericarditis, 1573
 adverse effects of, 2214
 alopecia, 354
 cobalamin malabsorption, 644t
 diarrhea, 269, 1573, 2234
 myopathy, 462e-20, 462e-20t, 2199, 2218t
 neuropathy, 2687t, 2688
 for Behçet's syndrome, 2194
 for calcium apatite deposition disease, 2236
 for calcium pyrophosphate deposition disease, 2236
 contraindications to, 1573
 drug interactions of, 2235
 for familial Mediterranean fever, 723, 2214
 for fever of unknown origin, 141
 for gout, 2234
 Cold abscess, 423
 Cold agglutinin disease, 659
 Cold exposure, cyanosis in, 249
 Cold-induced autoinflammatory syndrome, familial, 422
 Cold sore, 233
 Cold-stimulus headache, 2597
 Cold urticaria, 363, 363t, 2118-2119
 Colectomy, 274, 479, 539, 861
 Colesevelam, 2415, 2448t, 2449
 Colestipol, 2448t, 2449, 2453
 Colic, biliary. *See* Biliary colic
 Colic acid, 2076
 Colistin. *See* Polymyxin E (colistin)
 Colitis
 amebic, 1365
 bleeding in, 277
 CMV, in HIV infection, 1257
 collagenous, 270, 1957
 colonoscopy in, 1881, 1882f, 1899f
 cord-blood stem cell-associated, 772
 in Crohn's disease, 1953
 diarrhea in, 270
 diversion, 1957
 etiology of, 1882f
 vs. IBD, 1956-1957
 ischemic. *See* Ischemic colitis
 lymphocytic, 1957
 microscopic, 270
 primary immune deficiency diseases and, 2113
 pseudomembranous, 855, 858, 858f, 1882f. *See also* *Clostridium difficile*-associated disease
 sexually acquired, 881
 Collagen
 autoantibodies to, 372e-24t
 in connective tissues, 2505
 fibrillar, 2506-2507, 2506f
 Collagenoma, 2336t, 2340
 Collagenous colitis, 1957
 Collagenous sprue, 1942
 Collagen type IV-3 α -chain, 372e-24t
 Collagen vascular disease, 47e-4, 1467t, 1911
 Collecting duct
 disorders involving, 332e-7t
 functions of, 332e-5f, 332e-6f, 332e-8-9
 tumors of, 578, 578t. *See also* Renal cell carcinoma (RCC)
 Collectins, 145e-5, 372e-2
 Colles' fracture, 2488f
 Collier's sign, 210
 Colloid cysts, 603
 Colloid goiter, 2301
 Colon
 bleeding from, 277
 blood supply to, 1979, 1979f
 cathartic, 273
 diverticula of. *See* Diverticular disease
 functions of, 1875, 1933-1934
 length of, 1933
 obstruction of. *See* Intestinal obstruction
 physiology of, 264-265, 264t
 polyps of, 482, 1897, 1897f. *See also* Colorectal cancer
 storage and salvage in, 265
 transit time, measurement of, 273
 Colonic obstruction. *See* Intestinal obstruction
 Colonic pseudoobstruction. *See* Intestinal pseudoobstruction
 Colonization, 145e-1
 Colonization factors, 145e-2
 Colonoscopy, **1881**
 after polyp removal, 538
 in colitis, 1881, 1882f
 for colorectal cancer screening
 adenocarcinoma appearance in, 1883f
 in average-risk patients, 29t, 481t, 482, 1898t
 costs of, 30
 vs. flexible sigmoidoscopy, 540-541
 in inflammatory bowel disease, 1898t
 with personal or family history, 1898t
 polyp appearance in, 1877, 1882f, 1883f, 1897f, 1899f
 sensitivity of, 1881
 strategies for, 1897-1898
 in constipation, 273
 in diarrhea, 267, 272
 in gastrointestinal bleeding, 278-279
 indications for, 1878, 1879t
 in intestinal ischemia, 1980
 in nausea and vomiting, 260
 open-access, 1900
 patient preparation for, 1900
 polyp removal during, 346e-1v
 in ulcerative colitis, 1952, 1953f
 virtual, 540, 1898-1899, 1899f
 Colon stem cells, 453t
 Colony-stimulating factors, 413
 Colorado tick fever virus, 1306t, 1319
 Color anomia, 179, 197
 Color blindness, 197, 2629
 Colorectal cancer, **537**
 adenomatous polyps and, 537-538
 back pain in, 118
 carcinoma of unknown primary and, 120e-4
 clinical features of, 540-541, 541f
 deaths from, 467t, 469f, 471t, 537
 epigenetic silencing in, 102e-7
 etiology of, 538-539, 538t
 genetic considerations in
 chromosome abnormalities, 101e-1f, 101e-2t, 101e-3t, 101e-5, 537-538
 familial syndromes, 101e-1f, 101e-5, 101e-6t, 538-539, 538t
 genomewide profiling, 101e-9f
 IBD and, 1964-1965, 1965f
 incidence of, 467t, 468, 468f, 471t, 537
 infections in, 486t, 539
 insulin resistance and, 538
 lifetime cumulative risk of, 27t
 metastatic, 542, 553, 604t
 pathogenesis of, 101e-1f, 537-538
 postmenopausal hormone therapy and, 2384t, 2385
 prevention of, 476, 477-478, 479, 539
 prognosis of, 541-542, 542f, 542t
 recurrence of, 542
 risk factors for, 538-539, 538t
 screening for. *See also* Colonoscopy, for colorectal cancer screening
 benefits of, 27, 30, 540
 in elderly, 84
 in inflammatory bowel disease, 1898t
 with personal or family history, 1898t
 recommendations for normal-risk adults, 29t, 481t, 482, 540-541, 1898t
 techniques of, 540-541, 540t
 spread of, 542
 staging of, 541-542, 542f
 treatment of, 543-544
 biologic therapy, 102e-5t, 103e-21, 103e-21t, 543
 chemotherapy, 543
 endoscopic dissection, 346e-1v, 1887f
 endoscopic stent placement, 1895f
 late consequences of, 624t
 radiation therapy, 543
 surgery, 542-543
 targeted chemotherapy, 103e-19, 543
 tumor markers in, 473t
 Color flow Doppler, 270e-2, 270e-2f. *See also* Echocardiography
 Color vision, assessment of, 197
 Coltiviruses, 1305, 1306t
 Coltivirus infections, 1306t, 1319
 Coma, **1771**
 anatomy of, 329e-1v, 1771-1772
 approach to the patient
 brainstem reflexes, 1774-1775, 1774f
 history, 1773
 level of arousal, 1774
 neurologic examination, 41e-1v, 1773-1774
 physical examination, 329e-1v, 1773
 brain death determination in, 1776
 definition of, 1771
 diagnosis of, 329e-1v, 442e-2, 1772f, 1775
 differential diagnosis of, 1775-1776, 1776t
 in metabolic disorders, 1772-1773
 pathophysiology of, 1772-1773, 1772f
 prognosis of, 1777
 treatment of, 1776-1777
 Coma bullae, 361t, 362
 Combat veterans. *See* War veterans
 Combination antiretroviral therapy (cART). *See also specific drugs*
 action of, 214e-5
 adverse effects of, 6e-4, 1274-1275t, 2030, 2689
 in developing countries, 12, 13f
 entry inhibitors, **1275t**, 1277f, 1280f, 1281
 formulations of, 1277t
 in HBV virus infection, 1273
 indications for changing, 1282, 1282t
 initiation of, 770, 1282, 1282t
 integrase inhibitors, **1275t**, 1277f, 1280f, 1281
 principles of, 1273, 1281-1283, 1282t
 protease inhibitors, **1275t**
 action of, 1278
 adverse effects of, 1278, 2218t, 2422
 drug interactions of, 34t, 41, 798, 1278
 for HCV infection, 2044, 2045-2046t, 2046-2047
 for HIV infection, 1275t, 1278

- Combination antiretroviral therapy (cART), protease inhibitors (*Cont.*):
 molecular structure of, 1276f
 resistance to, 1278, 1280f
 resistance testing for, 1248
 reverse transcriptase inhibitors, **1274–1275t**
 action of, 1273, 1278
 adverse effects of, 318, 1273, 1278
 drug interactions of, 798
 for HIV infection, 1273, 1274–1275t, 1278
 molecular structure of, 1276f
 resistance to, 1279f
 selection of initial drugs, 1282, 1282t
- Combined pituitary hormone deficiency (CPHD), 2325t
- Comedones, 351
- Commensals, 86e-1
- COMMIT (Community Intervention Trial for Smoking Cessation), 476
- Common carotid artery, 2574
- Common cold, 225–226
 adenovirus, 1208
 clinical features of, 1202t
 coronavirus, 1204
 HMPV, 1206
 HRSV, 1205
 parainfluenza virus, 1207
 rhinovirus, 1203
 treatment of, 1203
- Common variable immunodeficiency (CVID), 2111
- Commonwealth Serum Laboratories, 2742
- Commotio cordis, 289e-3
- Communication
 of bad news, 57–58, 57t
 racial/ethnic factors in, 16e-3–4, 16e-4f
- Community-acquired pneumonia. *See* Pneumonia, community-acquired
- Community Intervention Trial for Smoking Cessation (COMMIT), 476
- Comparative genomic hybridization (CGH), 83e-3, 83e-3t
- Comparative genomics, 426
- Compensatory enlargement, 291e-4
- COMP gene mutations, 2512
- Complementary and alternative medicine, **14e-1**
 for asthma, 1678
 for back pain, 120
 cancer treatment, 473
 definitions and scope of, 14e-1–2
 herbal medicines. *See* Herbal medicines/supplements
 information sources on, 14e-4, 14e-4t
 licensure and accreditation in, 14e-2
 patterns of use of, 14e-2
 vitamin and mineral supplements. *See* Dietary supplements
- Complement factor H, 205, 1832
- Complement system
 deficiencies of, 372e-28, 372e-29t, 2106–2107
 definition of, 372e-1
 in HIV infection, 1244
 innate immune system and, 372e-4t, 372e-9
 pathways of, 372e-9, 372e-16f
 in shock, 1746
- Complete blood count (CBC), 393, 394t
- Complete heart block, 278e-4f, 278e-10f, 1472. *See also* Atrioventricular (AV) conduction block
- Complete response, to chemotherapy, 103e-5
- Complex IV (COX), 94e-3
- Complex partial status epilepticus, 2607
- Complex regional pain syndrome (CRPS)
 clinical features of, 90, 2246, 2644
 treatment of, 2644
 type I, 90, 2246, 2643
 type II, 90, 2643–2644
- Complex sleep apnea, 1727
- Complex systems, 87e-1
- Composite tissue transplantation, 928
- Compound heterozygote, 435
- Comprehension, assessment of, 177
- Compression of morbidity, 94e-1
- Compression stockings
 for chronic venous disease, 1652
 for DVT prophylaxis, 1634
 for lymphedema, 1655
 for stasis dermatitis/ulceration, 344t, 346
- Compression ultrasonography, of leg veins, 613
- Computed tomography (CT)
 adrenal, 1619
 in adrenocortical carcinoma, 2323f
 in appendicitis, 1988, 1988f
 in ARDS, 308e-10f, 1738f
 in asbestosis, 308e-15f
 in *Aspergillus* infections, 1348
 in bile duct evaluation, 2085t
 in bone mass measurement, 2493
 in bronchiectasis, 308e-12f
 in CAH, 2328f
 in cardiac disease
 angiography. *See* Computed tomography (CT) angiography (CTA)
 CAD, 270e-10, 270e-11f, 270e-12, 270e-12f, 270e-13f, 1585
 for calcium scoring, 270e-4, 270e-7f, 270e-11f
 contrast agents in, 270e-9
 cor pulmonale, 1506
 infective endocarditis, 270e-24, 270e-25f
- chest, **308e-2**
 air bronchograms, 308e-11f
 normal, 308e-2f
 parenchymal loss, 308e-5–6f
 volume loss, 308e-3–4f
 in cirrhosis, 286, 286f
 in Cushing's syndrome, 2317, 2317f
 in diffuse axonal injury, 457e-2f
 in eosinophilic granulomatosis with polyangiitis, 1685
 in granulomatosis with polyangiitis, 2183f
 halo sign on, 1348
 in hearing loss evaluation, 223
 helical, 1664, 1665f
 in hemoptysis, 246–247
 high-resolution, 1664
 in hypersensitivity pneumonitis, 1682, 1682f
 in interstitial lung diseases, 308e-7–9f, 1710, 1710f
 in intestinal obstruction, 1984, 1984f
 in lung cancer diagnosis, 308e-15–16f
 in lung cancer screening, 509–510, 510t
 in lung cancer staging, 513
 in mineralocorticoid excess, 2319
 multidetector, 1664–1666
 in musculoskeletal disease evaluation, 2224–2225, 2224t, 2225f
 in nephrolithiasis, 1869f
 in NETs, 572, 572f
 in neurologic disease. *See also* Neuroimaging, CT
 adrenoleukodystrophy, 441e-41f
 anterior dens dislocation, 441e-31f
 cavernous malformation, 440e-7f
 cerebral contusion, 457e-1f
 CNS vasculitis, 441e-19f
 coma, 1775
 epidural hematoma, 441e-31f, 457e-3f
 Huntington's disease, 441e-23f
 intracerebral hemorrhage, 2584
 Jefferson fracture, 441e-32f
 lacunar infarction, 441e-17f
 ligament injury, 441e-33f
 neurologic critical illness, 1779
 seizures, 2550
 stroke, 2580, 2581f
 subarachnoid hemorrhage, 1786, 1786f
 subdural hematoma, 457e-3f, 457e-4
 superior sagittal sinus thrombosis, 441e-20f
 vertebral fractures, 441e-31f, 441e-33f, 441e-34f
 viral encephalitis, 895
 in osteomyelitis, 842, 842f
 in pancreatic cancer diagnosis, 555, 555f
 in pancreatic disease evaluation, 2087t, 2088–2089
- in pancreatitis, 2094–2095, 2094f, 2095f, 2099–2100, 2100f
 in pneumothorax, 308e-14f
 in pulmonary arteriovenous malformation, 308e-17f
 in pulmonary embolism, 308e-17f, 1633, 1633f
 in pulmonary hypertension, 308e-18f, 1656
 in respiratory disease evaluation, 1663, 1664–1666, 1664f, 1665f
 in sarcoidosis, 2206, 2206f, 2208f, 2209f
 of temporal bone, 220, 223
 in tracheomalacia, 308e-13f
 in urinary tract obstruction, 1873
- Computed tomography (CT) angiography (CTA)
 in cerebral artery aneurysm, 440e-2f
 coronary
 in anomalous right coronary artery, 271e-4f
 in aortic aneurysm, 1640f
 in aortic stenosis, 270e-17
 in CAD evaluation, 270e-5f, 271e-2f, 1594–1595, 1594f
 in chest pain evaluation, 102, 270e-14, 270e-14f, 270e-15, 270e-16f
 contrast agents in, 270e-9
 in coronary aneurysm, 271e-4f
 in murmur evaluation, 51e-8
 principles of, 270e-4
 neurologic, 440e-1
 in cerebral artery aneurysm, 440e-2f
 in cerebral artery occlusion, 440e-3f
 in small-vessel stroke, 2567f
 in subarachnoid hemorrhage, 1786f
 in pulmonary hypertension, 1656
 of renal arteries, 1628, 1628t, 1630f
- Computed tomography (CT) cisternography, 440e-2
- Computed tomography (CT) colonography, 481t, 482
- Computed tomography (CT) myelography, 440e-2, 440e-9–10
- Computerized physician order entry (CPOE), 12e-2
- COMT (catechol-O-methyltransferase) inhibitors, 2615, 2616t
- Concentration gradients, 332e-6
- Concussion, **457e-1**
 clinical features of, 471e-3
 definition of, 457e-1
 mechanics of, 457e-1
 sports-related, 457e-4
 in war veterans, 471e-2–3, 471e-3t
- Condoms, 1009, 1285
- Conduction aphasia, 177t, 178
- Condyloma acuminata (anogenital warts)
 anal cancer and, 544
 clinical features of, 76e-11f, 1198, 1199f
 in HIV infection, 1263
 treatment of, 215e-4t, 351, 1200–1201, 1201f
- Condylomata lata, 25e-5f, 76e-10f, 1259
- Cone(s), 195, 197
- Cone-nose bug bite, 2750
- Cone snail envenomation, 2740
- Confabulation, 182
- Confidentiality, in patient-physician relationship, 17e-1
- Conflicts of interest, 17e-3–4
- Confocal fluorescent laser microscopy (CFM), 1668
- Confusion, 166, 1771. *See also* Delirium
- Confusion Assessment Method, 167, 167t
- Conga ant, 2749
- Congeniters, 2723
- Congenital absence of vagina, 2357
- Congenital adrenal hyperplasia (CAH), **2327**
 clinical features of
 ambiguous genitalia, 437, 2355
 cutaneous, 356t, 2355
 hirsutism, 331, 2355
 hypokalemia, 306
 precocious puberty, 2362
 sex differences in, 2328
 diagnosis of, 332–333, 2328, 2328f
 etiology of, 2355

- genetic considerations in
CYP gene mutations, 2318t, 2327–2328, 2328t, 2355, 2356t
 gene conversion, 434
 sex-limited factors, 437, 2328
 HLA gene association with, 373e-8t
 treatment of, 334, 2328–2329, 2356–2357
 variants of, 2327–2328, 2328t
- Congenital amaurosis. *See* Leber's hereditary optic neuropathy (congenital amaurosis)
- Congenital central hypoventilation syndrome, 1722
- Congenital erythropoietic porphyria (CEP), 2521, 2522t, 2532
- Congenital generalized lipodystrophy, 436e-8f
- Congenital heart disease (CHD), 1519. *See also specific defects*
 AV conduction block in, 1472
 brain abscess and, 901
 care of survivor of surgical care of, 1528
 chest wall deformity in, 1443
 clubbing/hypertrophic osteoarthropathy in, 250
 complex, 1521
 cyanosis in, 249
 endocarditis prophylaxis in, 1528
 head and neck features of, 1443
 imaging in, 270e-24–25, 270e-25f, 270e-26f
 incidence of, 1519
 intermediate complexity, 1520t
 intracardiac shunt in, 1463
 percutaneous interventions for, 296e-4
 polycythemia in, 400
 pregnancy and, 47, 1519, 1522, 1528
 simple, 1520t
 in Turner's syndrome, 2351–2352
- Congenital hypothyroidism, 2283, 2284t, 2289–2290
- Congenital infections. *See* Neonate(s), infections in
- Congenital lipoid adrenal hyperplasia (CLAH), 2324t
- Congenitally corrected transposition, 1528
- Congenital muscular dystrophy(ies) (CMDs), 462e-9, 462e-10t
- Congenital myasthenic syndromes (CMS), 2702, 2703t
- Congenital rubella syndrome. *See* Rubella, congenital
- Congenital syphilis. *See* Syphilis, congenital
- Congenital vascular malformations, 2585–2586
- Congestive heart failure. *See* Heart failure (HF)
- Conivaptan, 302, 610, 2281
- Conjugate eye movement, 1774, 1774f
- Conjunctivitis, 199
 acute hemorrhagic, 1293, 1293f
 allergic, 199
 atopic, 199
C. trachomatis, 1169, 1170t, 1171, 1172
 gonococcal, 1006, 1009t
Onchocerca, 1421
 in tularemia, 1068
- Connective tissue(s), 2505–2506, 2505t, 2507f
- Connective tissue disorders, 2504
 Alport's syndrome. *See* Alport's syndrome
 chondrodysplasia. *See* Chondrodysplasia(s)
 classification of, 2504–2505
 Ehlers-Danlos syndrome. *See* Ehlers-Danlos syndrome
 elastin-related diseases, 2513
 epidermolysis bullosa, 257, 434, 2513
 interstitial lung disease in, 1713–1714
 Marfan's syndrome. *See* Marfan's syndrome
 osteogenesis imperfecta. *See* Osteogenesis imperfecta
- Connexin(s), 220, 444e-4
- Conn's syndrome. *See* Adenoma, aldosterone-producing adrenal
- CoNS (coagulase-negative staphylococci), 954, 960, 961. *See also* Staphylococcal infections
- Consciousness, assessment of, 2537
- Constipation, 272
 abdominal swelling in, 285
 approach to the patient, 273, 273f
 assessment of, 61
 definition of, 272
 differential diagnosis of, 1877
 epidemiology of, 264
 etiology of, 61, 272, 272t
 idiopathic, 272
 in irritable bowel syndrome, 1965
 opioid-induced, 61, 92
 rectal prolapse and, 1974
 severe, 273–274
 in terminally ill patient, 61
 treatment of, 61, 62t, 85, 92, 274
- Construction apraxia, 182
- Consumption. *See* Tuberculosis (TB)
- Contact dermatitis, 345
 allergic, 76e-2f, 345, 354t
 clinical features of
 acute phase, 343f
 chronic phase, 343f
 distribution of skin lesions, 341t, 342f, 354t
 erythroderma, 354t
 morphology of skin lesions, 341t, 354t
 vesicles/bullae, 361, 361t
 diagnosis of, 344, 354t
 drug-induced, 44, 381
 inflammasome activity in, 372e-8t
 irritant, 345, 354t
 treatment of, 345, 354t
- Contact sensitivity, 2119
- Contact urticaria, 2118
- Contig, 86e-4, 769t, 773f
- Contiguous gene syndromes, 432, 437
- Continuing education, 7
- Continuous ambulatory peritoneal dialysis (CAPD). *See also* Peritoneal dialysis
 for acute kidney injury, 1811
 for chronic kidney disease, 1824
 peritonitis in, 848, 1825
- Continuous arteriovenous rewarming, 478e-3t
- Continuous positive airway pressure (CPAP)
 adverse effects of, 1727t
 for obesity hypoventilation syndrome, 1722
 principles of, 1742
 for sleep apnea, 1620, 1727
- Continuous quality improvement, 12e-2–3
- Continuous renal replacement therapy, for acute kidney injury, 1811
- Continuous wave Doppler, 270e-2, 270e-2f. *See also* Echocardiography
- Contraception, 2389
 AEDs and, 2559
 barrier methods, 2390, 2390t
 effectiveness of, 2390, 2390t
 hormonal methods
 injection, 2390t, 2391
 long-term, 2390t, 2391
 male, 2372
 oral contraceptives. *See* Oral contraceptives
 patch, 2390t, 2391
 intrauterine devices, 2390, 2390t
 obesity effects on, 2391
 postcoital, 2391
 sterilization, 2390t
- Contractures
 definition of, 2219t
 etiology of, 2220
 in muscular dystrophy, 462e-6, 462e-9
 in myopathies, 462e-3
- Contrast agents
 adverse effects of
 allergic reactions, 440e-4, 440e-5, 1461
 diuresis, 295
 immediate cutaneous, 377
 nephrogenic systemic fibrosis, 379, 440e-5, 1819
 nephropathy, 270e-9, 440e-2, 1461, 1803, 1806t, 1820
 neurologic, 440e-10
 pustular eruptions, 382. *See also* Acute generalized eruptive/exanthematous pustulosis (AGEP)
- seizure, 2547t
 SJS/TEN, 383. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
 uricosuria, 431e-5
 urticaria/anaphylaxis, 381
 for cardiac imaging, 270e-9
 for MRI, 440e-5
 precautionary measures for, 440e-4, 440e-4t, 440e-10, 1820
- Contrast enema. *See* Barium radiography
- Control, of disease, 785
- Control chart, 12e-3
- Contusion, cerebral, 457e-1, 457e-1f
- Conus medullaris syndrome, 2651
- Convergence-projection hypothesis, of referred pain, 89f
- Conversion disorders, 2608, 2719
- Co-occurrence analysis, 86e-5
- Cooling blankets, 126, 479e-3
- Cooling strategies, in heat-related illness, 479e-3
- Coombs test, 138e-4, 138e-4f
- Coordination, assessment of, 2539
- COPD. *See* Chronic obstructive pulmonary disease (COPD)
- Copper, 96e-10
 deficiency of. *See* Copper deficiency
 functions of, 96e-10
 liver, 2520t
 parental supplementation of, 98e-6t
 recommended intake of, 95e-3t
 reference values for, 2766t
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t, 96e-10. *See also* Wilson's disease
 urine, 2519, 2520t
- Copper deficiency
 clinical features of, 96e-9t, 2690–2691
 diagnosis of, 96e-10
 etiology of, 96e-10, 1927, 2691
 genetic considerations in, 96e-10
 treatment of, 2691
- Copperhead bite, 2736t, 2738. *See also* Snakebites
- Copper-silver ionization, of water supply, 1020
- Coprolalia, 2623
- Coproporphyrin, hereditary. *See* Hereditary coproporphyrin (HCP)
- Copy number variations (CNVs), 427, 430f, 444e-1
- Coral snake bite, 2737t, 2738. *See also* Snakebites
- Cord blood stem cells, 89e-3–4, 90e-2, 139e-2, 453t
- Cord blood transplantation, 139e-2, 919
- Cord colitis, 772
- Core-needle biopsy, 103e-2
- Co-repressors, 428
- CoreValve ReValving system, 296e-5
- Cori syndrome (type IIIa GSD), 433e-2t, 433e-3
- “Corkscrew” esophagus, 1906, 1906f
- Corneal abrasions, 199
- Corneal beading, 1125
- Corneal blink reflex, 1774f, 1775
- Corneal light reflex, 197
- Corollary discharge, 47e-1
- Coronary angiography
 in cardiogenic shock, 1760
 coronary artery visualization on, 1464–1465, 1464f
 CT. *See* Computed tomography (CT)
 angiography (CTA)
 intravascular ultrasound in, 1465, 1465f
 optical coherence tomography during, 1465, 1465f
 in patent ductus arteriosus, 1524f
 in Prinzmetal's variant angina, 1598
 in STEMI, 1605
- Coronary arteriography, 1585–1586
- Coronary arteriovenous fistula, 1523
- Coronary artery(ies)
 aneurysm of, 271e-4f
 calcification of, 270e-4, 270e-7f, 270e-11f, 271e-2f, 1585
 CT angiography of. *See* Computed tomography (CT) angiography (CTA)

- CT of. *See* Computed tomography (CT), in cardiac disease
- embryonic development of, 265e-12
- fistula of, 1523
- MRI of. *See* Cardiac magnetic resonance imaging (CMR)
- normal, 1464, 1464f
- perfusion imaging of. *See* Myocardial perfusion imaging
- stenosis of, 1465, 1465f
- Coronary artery bypass grafting (CABG)
- brain injury after, 463e-2-3, 463e-2f
- for cardiogenic shock, 1762
- complications of, 1602t
- for heart failure, 1514-1515
- for ischemic heart disease, **1591**
- indications for, 1591-1592
- minimally invasive, 1592
- vs. PCI, 296e-3-4, 296e-3f, 1592, 1593f
- results of, 1592
- for myocardial infarction, 1597, 1598t
- perioperative, 53
- pleural effusion after, 1718
- racial/ethnic disparities in referral for, 16e-1, 16e-2
- survival benefit of, 1592
- in women, 6e-3
- Coronary artery disease (CAD). *See also* Atherosclerosis; Ischemic heart disease
- in alcohol use disorder, 2725
- in allografts, 1517-1518
- AV conduction block in, 1472
- cardiac arrest/sudden cardiac death and, 1765
- clinical features of, 1439
- angina pectoris. *See* Angina pectoris
- cardiac symptoms, 1439
- dyspnea, 47e-2
- oral pain, 237
- palpitations, 254
- deaths from, 6e-1t, 6e-2f
- diagnosis of
- CMR, 270e-12-13, 270e-13f, 270e-20f, 270e-26v
- coronary angiography, 1465, 1465f
- C-reactive protein, 291e-9
- CT, 270e-10, 270e-11f, 270e-12, 270e-12f, 270e-13f, 1585
- ECG, 269e-7f
- imaging tests, 270e-9-10, 270e-9t
- myocardial perfusion imaging, 270e-10, 270e-10f, 270e-11f, 270e-20f, 271e-2f
- pre- and posttest probability analysis, 22, 23f, 270e-9
- sex differences in, 6e-3
- stress echocardiography, 270e-10, 270e-26v, 271e-1f, 271e-1v
- diet and, 95e-5
- estrogen therapy and, 291e-7-8
- heart failure in, 1501. *See also* Heart failure (HF)
- high-altitude travel and, 476e-5
- hypertension and, 1615
- lipoprotein disorders and, 291e-5-6, 2442. *See also* Lipoprotein disorders
- metabolic syndrome and, 2450
- oral contraceptives and, 2391, 2391t
- pathophysiology of, 372e-24t, 1579-1580
- PCI for. *See* Percutaneous coronary interventions (PCI), for revascularization
- periodontal disease and, 236
- peripheral artery disease and, 1644
- postmenopausal hormone therapy and, 2384t, 2385
- preoperative testing for, 52f, 53
- prevention of, 291e-6t, 291e-9
- radiation-associated, 621
- SA node dysfunction in, 1467, 1467t
- vasoconstrictor use in, 242
- ventricular tachycardia in, 1494-1495
- in women, 6e-1t, 6e-2-3, 6e-2f
- Coronary artery spasm (Prinzmetal's variant angina), 1456, 1579, 1588, **1598**
- Coronary atherosclerosis. *See* Coronary artery disease (CAD)
- Coronary blood flow, 1579
- Coronary care units, 1605-1606
- Coronary revascularization. *See* Coronary artery bypass grafting (CABG); Percutaneous coronary interventions (PCI)
- Coronaviruses
- characteristics of, 1203-1204
- gene expression and replication in, 214e-4
- structure of, 214e-2t, 214e-3f, 1204
- Coronavirus infections, **1203**
- clinical features of, 225, 231, 234, 1202t, 1204
- diagnosis of, 1204-1205
- diarrhea in, 266
- epidemiology of, 1204
- MERS. *See* Middle East respiratory syndrome (MERS)
- pathogenesis of, 1204
- prevention of, 1205
- SARS. *See* Severe acute respiratory syndrome (SARS)
- in transplant recipient, 923
- treatment of, 1205
- Coronin A deficiency, 375e-1
- Cor pulmonale, **1505**
- clinical features of, 1506
- vs. constrictive pericarditis, 1576
- in COPD, 1704
- diagnosis of, 1454, 1506
- etiology of, 1505, 1506f
- pathophysiology of, 1505-1506
- in polycythemia, 400
- right ventricular dysfunction in, 270e-7
- treatment of, 320e-2
- Corpus callosum, agenesis of, 478e-1
- Corpus luteum, 2376f, 2379
- Corrected count increment (CCI), 138e-3
- Corrigan's (water-hammer) pulse, 51e-6, 1445, 1535
- Corrosive esophagitis, 1910
- Corticobasal degeneration syndrome
- clinical features of, 163, 173, 179, 2605, 2611
- neuropathology of, 2604f, 2605, 2605f, 2611
- Corticospinal pathway, 155f
- Corticosteroid-binding globulin (CBG), 2378
- Corticosteroids. *See* Glucocorticoid(s)
- Corticotropin-releasing hormone, ectopic production of, 609t, 611
- Cortisol
- circadian rhythm of, 2310, 2310f
- in hypoglycemia, 2431t
- inactivation of, 2313, 2313f
- laboratory evaluation of, 2310
- in shock, 1745
- synthesis of, 2309f
- Cortisol-binding globulin (CBG), 2312
- Cortisone acetate, 2259t
- Corynebacterium* spp., 980-981
- Corynebacterium diphtheriae*, 150e-2t, 231, 977. *See also* Diphtheria
- Corynebacterium minutissimum*, 343, 981
- Corynebacterium ulcerans*, 231, 980
- COSMIC (Catalogue of Somatic Mutations in Cancer), 101e-8
- Cost-effectiveness, of screening tests, 30
- Co-stimulatory molecules, 372e-1, 372e-19
- Costochondritis, 96t, 98, 100, 2246
- Costoclavicular maneuver, 1647
- Cosyntropin test, 2310, 2325, 2346
- Cotransporters (symporters), 332e-6
- Cotton dust exposure, 1688t, 1691
- Cottonmouth water moccasin bite, 2736t. *See also* Snakebites
- Cotton-wool spots
- retinal, 40e-3, 40e-3f, 201f
- in skull radiographs, 426e-2, 426e-2f
- Cough, **243**
- assessment of, 244, 1661
- chronic, 244-245
- duration of, 244
- etiology of, 243-245
- in GERD, 262
- high-altitude, 476e-4
- impaired, 243, 243t
- in lung cancer, 510, 510t
- mechanism of, 243, 243f
- in *Nocardia* infections, 1084
- nocturnal, 1503-1504
- in pertussis, 1022-1023, 1023t. *See also* Pertussis
- physical examination in, 244
- in respiratory disease, 1661
- with swallowing, 256
- symptomatic, 243-244
- treatment of, 245
- in tuberculosis, 1108
- Cough Assist Device, 2635
- Cough headache, primary, 2596-2597
- Cough hypersensitivity syndrome, 245
- Coumarin, herbs that contain, 404t
- Counter-current in-line heat exchanger, 478e-3
- Counter-current multiplication, 332e-8
- Courvoisier's sign, 555
- Cover test, 197, 208
- Cowden's syndrome (multiple hamartoma syndrome)
- clinical features of, 101e-6t, 364t, 365, 451e-2t, 2344
- genetic considerations in, 101e-6t, 365, 446, 451e-2t, 599t, 2337t, 2344
- tumors associated with, 365, 599t
- Cowpox, 220e-1t, 220e-2
- COWS mnemonic, 1775
- COX. *See* Cyclooxygenase (COX)
- COX (complex IV), 94e-3
- Coxiella burnetii*, 1154, 1161. *See also* Q fever
- COX inhibitors. *See* Cyclooxygenase (COX) inhibitors
- Coxsackievirus, 145e-2t
- Coxsackievirus infections. *See also* Enterovirus infections
- acute hemorrhagic conjunctivitis, 1293, 1293f
- clinical syndromes associated with, 1291t
- diagnosis of, 1293
- hand-foot-and-mouth disease. *See* Hand-foot-and-mouth disease
- herpangina, 231, 238t, 1293
- vs. herpes zoster, 1185
- laryngitis, 234
- meningitis, 890
- myocarditis and pericarditis, 1292
- oral manifestations of, 238t
- pharyngitis, 231
- pleurodynia (Bornholm disease), 1291
- skin manifestations of, 362, 362t, 1292
- maculopapular eruptions, 128t
- peripheral eruptions. *See* Hand-foot-and-mouth disease
- petechial rash, 133t
- treatment of, 1293-1294
- CPAP. *See* Continuous positive airway pressure (CPAP)
- CPC (capsular polysaccharide complex), 849
- cpe* gene, 991
- C peptide, 2402, 2434, 2758t
- CpG islands, 102e-7, 102e-8, 430, 431
- CPHD (combined pituitary hormone deficiency), 2325t
- CPIS (Clinical Pulmonary Infection Score), 811, 811t. *See also* Ventilator-associated pneumonia (VAP)
- CPOX gene, 2523t
- CPPD (calcium pyrophosphate deposition) disease, 372e-8t, 2235-2236, 2235t, 2236f
- CPT (carnitine palmitoyltransferase), 462e-15
- Crab lice (*Pthirus pubis*), 2745-2746
- Crab yaws, 207e-2
- Crack, 469e-1. *See also* Cocaine
- Crackles, 1662
- Cradle cap, 347, 1357
- Cramps, muscle, 462e-2-3
- Cranial epidural abscess, 904-905, 904f
- Cranial nerve disorders
- in botulism, 988
- dysphagia and dysphonia in, 2649
- etiology of, 2649t

- facial nerve, 2648–2649. *See also* Bell's palsy
glossopharyngeal neuralgia, 237, 2649
in Guillain-Barré syndrome, 2650, 2694
injuries, 457e-2
multiple, 2650
neck weakness in, 2649
in subarachnoid hemorrhage, 1785
tongue paralysis in, 2649
trigeminal nerve, 215, 2646–2647, 2647t. *See also* Trigeminal neuralgia
- Cranial nerve examination, 2537–2538
- Cranioectodermal dysplasia (Sensenbrenner's syndrome), 1850t
- Craniofacial synostosis (Crouzon's syndrome), 435
- Craniopharyngiomas, 198, 415e-4, 603, 2262, 2264
- “Crazy paving,” 1690, 1690f
- C-reactive protein (CRP)
age-related changes in levels of, 75, 75f
in cardiovascular disease, 291e-9, 291e-9f
in chronic kidney disease, 1816–1817
in community-acquired pneumonia, 806
in fever evaluation, 125
high-sensitivity, 1249t, 2758t
in immune response to pneumococci, 950
in infectious diseases, 764
reference values for, 2758t
in sepsis/septic shock, 1752
in ulcerative colitis, 1952
- Creatine deficiency, 435e-1t
- Creatine kinase (CK)
myocardial band (CK-MB)
in cardiac trauma, 289e-3
in myocardial infarction, 1594, 1600–1601, 1601f
reference values for, 2759t
- serum
in distal myopathies, 462e-12, 462e-13t
in glycogen storage disorders, 462e-14
in glycolytic defects, 462e-14
in inflammatory myopathies, 2199, 2199t
in limb-girdle muscular dystrophies, 462e-6t, 462e-7t
in muscle disease, 462e-4
in progressive muscular dystrophies, 462e-5t, 462e-6, 462e-8, 462e-9
reference values for, 2759t
- Creatine phosphate, 265e-11
- Creatinine
serum, 291t
in acute kidney injury, 1805
causes of abnormal values, 464t
in estimation of glomerular filtration rate, 480e-3–4, 480e-3f, 1813t
in hypovolemia, 298
in nutritional assessment, 464t
as surrogate for glomerular filtration rate, 291
urine, 291t
causes of abnormal values, 464t
in nutritional assessment, 464t
- Creatinine clearance, 37, 38, 46, 290–291
- Creatinine-height index, 460
- CREB (cAMP response-element binding protein)
action of, 444e-4
diseases associated with, 431t
in opiate addiction, 465e-3, 465e-3f, 468e-1, 468e-2f
- Cremasteric reflex, 2539
- Crepitus, 2219t, 2220
- “Crescent sign,” 489
- CREST syndrome, 357, 376, 2155, 2223. *See also* Systemic sclerosis (scleroderma)
- Cretinism, 236, 2285
- Creutzfeldt-Jakob disease (CJD)
after dura matter grafts, 453e-3
after growth hormone therapy, 453e-4
after pituitary gonadotropin therapy, 453e-4
clinical features of, 173, 173t, 453e-1, 453e-4, 2606
diagnosis of, 442e-1f, 442e-2, 453e-4
differential diagnosis of, 453e-4–5, 2606
- familial, 453e-1t, 453e-4
genetic considerations in, 172t, 453e-3
iatrogenic, 453e-1t, 453e-2, 453e-3
infection control in, 453e-5–6
laboratory tests in, 453e-5, 453e-5f
molecular basis of, 172t
neuropathology of, 172t, 453e-4
prevention of, 453e-6
as prion disease, 453e-2
treatment of, 453e-6
variant, 453e-1t, 453e-2, 453e-4
- Cricopharyngeal bar, 256
- Cri-du-chat syndrome, 83e-6
- Crigler-Najjar syndrome, 2001
clinical features of, 282, 1995, 2001–2002, 2001t
diagnosis of, 2001–2002, 2001t
genetic considerations in, 2001–2002
subtypes of, 282, 2001t
treatment of, 2068
- Crimean-Congo hemorrhagic virus, 1308t, 1321
- CRISPRs, 636, 776
- Critical care medicine, 1729. *See also* Critically ill patient
- Critical illness-related corticosteroid deficiency (CIRCI), 1755, 1757
- Critically ill patient
acute kidney injury in, 1735
anemia in, 1735
approach to, 1729
assessment of illness severity in, 1729, 1730f, 1730t
cholestasis in, 284–285
DVT in, 1734. *See also* Deep vein thrombosis (DVT)
glycemic control in, 1734
hypoglycemia in, 2433
hypogonadism in, 2364
monitoring of, 1733–1734, 1734f
multiorgan system failure in, 1733
neurologic disorders in, 1777, 1778t. *See also* Coma
approach to the patient, 1779
diagnostic studies, 1779
increased ICP monitoring and treatment, 1779–1780, 1780t. *See also* Intracranial pressure (ICP), increased
secondary brain insults, 1780–1781
brain death, 1735
central pontine myelinolysis, 301, 1782, 1782f
coma. *See* Coma
delirium. *See* Delirium
hypoxic-ischemic encephalopathy, 1735, 1773, 1780–1781, 1781f
metabolic encephalopathies, 178, 1782
myopathy, 462e-19, 1784
neuromuscular transmission disorders, 1784
pathophysiology of
brain edema, 1777
cerebral perfusion and autoregulation, 1778, 1778f
CSF and intracranial pressure, 1778–1779, 1779f
ischemic cascade and cellular injury, 1777–1778, 2561f
peripheral nervous system, 1778t, 1783–1784
polynuropathy, 1755, 1783–1784, 2684
respiratory evaluation in, 1783
sepsis-associated encephalopathy, 1782
status epilepticus. *See* Status epilepticus
stroke. *See* Stroke
subarachnoid hemorrhage. *See* Subarachnoid hemorrhage
Wernicke's disease/encephalopathy. *See* Wernicke's disease/encephalopathy
nutritional status of, 1734
prevention of complications in, 1734–1735
respiratory failure in, 1731–1732, 1732f4. *See also* Acute respiratory distress syndrome (ARDS)
sepsis in, 1734
- shock in, 1729–1731, 1731f. *See also* Shock
specialized nutritional support, 98e-3
stress-related ulcers in, 1734, 1930
weakness in, 1734
withholding/withdrawing care from, 1735–1736
- Critical values, in laboratory tests, 480e-2, 480e-2t
- Crizotinib
action and targets of, 101e-8, 102e-3, 102e-5t, 103e-16f, 103e-17t, 103e-19
adverse effects of, 103e-17t
for lung cancer, 103e-19, 521
- CroFab (Crotalidae polyvalent immune fab), 2736t
- Crohn's disease. *See also* Inflammatory bowel disease (IBD)
cancer risk in, 1965
clinical features of
gastrointestinal disease, 262, 1954
gastrointestinal bleeding, 277
ileocolitis, 1953
jejunoileitis, 1953
nausea and vomiting, 258
perianal disease, 1953–1954
colorectal cancer in, 1964–1965
complications of, 1954
diagnosis of
endoscopic, 1954, 1954f
MRI in, 1954, 1955f
radiology in, 1939f, 1954
small-intestinal mucosal biopsies in, 1939, 1940t
differential diagnosis of, 1061, 1168, 1956–1957, 1956t
epidemiology of, 1947, 1947t
etiology of, 1947, 1948f
genetic considerations in, 1947, 1949, 1950t
immune regulation in, 1950–1951
inflammation activity in, 372e-8t
inflammatory mediators in, 372e-29t, 1950–1951
pathogenesis of, 372e-23, 372e-24t, 1948f, 1949–1951
pathology of, 1951–1952, 1951f, 1952f
in pregnancy, 49, 1964
racial and ethnic differences in, 1947–1948
serologic markers in, 1954–1956
smoking effects on, 1947
spondyloarthritis and, 2178–2179
“string sign” in, 1953, 1954
treatment of, 1963–1964, 1963t
vitamin K deficiency in, 96e-8
- Cromolyn sodium, 1678, 2122, 2123f
- Cronkite-Canada syndrome, 270
- Cross-bridge cycling, in cardiac contraction, 265e-5f, 265e-6
- Cross-cultural education, 16e-5–6
- Crossed straight-leg raising sign, 113
- Crossing-over, 428, 430f
- Cross-matching, 138e-2
- Cross-sensitivity, drug, 384–385
- Cross-talk, nuclear-mitochondrial, 85e-7, 85e-7f
- Croup, 234, 1206, 1207
- Crouzon's syndrome (craniofacial synostosis), 435
- Crow-Fukase syndrome. *See* POEMS syndrome (Crow-Fukase syndrome)
- Crown, tooth, 236
- CRP. *See* C-reactive protein (CRP)
- CRPS. *See* Complex regional pain syndrome (CRPS)
- Cruise ship passenger, 266
- Crust (skin lesion), 340t, 827f, 828t, 829
- Crusted (Norwegian) scabies, 1263, 2744
- Cryoablation, for ventricular tachycardia, 1500
- Cryofibrinogenemia, 369
- Cryoglobulin(s), 2190
- Cryoglobulinemia
in chronic HCV infection, 2041, 2048, 2180, 2191
clinical features of, 1835t, 2191
essential mixed, 2190
hand arteriogram in, 386e-5f

- Cryoglobulinemia (*Cont.*):
 monoclonal, 368
 pathogenesis of, 2191
 treatment of, 2191
 vasculitis and, 2190–2191
 vessel wall disorders in, 732
- Cryohydrocytosis, 653
- Cryomapping, 273e-7
- Cryoprecipitate, 138e-2, 138e-2t, 138e-3
 for coagulation disorders in chronic kidney disease, 1818
 for coagulation disorders in liver disease, 739
 for DIC, 738
 for hemophilia, 734
- Cryopyrin-associated periodic syndromes (CAPS), 141, 2213t, 2215
- Cryopyrin/NALP3 inflammasome, 145e-7
- Cryosurgery, 103e-5, 501
- Cryotherapy, 351, 1201, 1201t
- Cryptic epitopes, 377e-2
- Cryptic miliary tuberculosis, 1112
- Cryptococcomas, 1340
- Cryptococcus* spp., 765t, 1329–1330, 1339
- Cryptococcus* spp. infections, **1339**
 approach to the patient, 1340
 in cancer patient, 488, 488t, 491
 chronic meningitis in, 908t
 clinical features of, 1340
 complications of, 1341–1342
 diagnosis of, 150e-1, 765t, 1340–1341
 disseminated, 25e-9f, 135
 epidemiology of, 765t, 1339
 in HIV infection, 1252t, 1264, 1341–1342
 pathogenesis of, 1340, 1340f
 prevention of, 1342
 prognosis of, 1341–1342
 skin manifestations of, 25e-9f, 356t, 1340, 1341f
 subacute meningitis, 898–899
 in transplant recipient, 924t, 926
 treatment of, 899, 1341
- Cryptogenic organizing pneumonia, 1712–1713
- Cryptogenic sensory and sensorimotor polyneuropathy (CSPN), 2691
- Cryptorchidism, 588, 2355, 2365
- Cryptosporidium*/cryptosporidiosis, **1407**
 clinical features of, 152e-5t, 266, 266t, **1407**
 diagnosis of, **1407**
 duodenal aspirate or jejunal biopsy, 245e-5t
 enzyme immunoassay, 245e-6t
 stool studies, 152e-5t, 245e-1, 245e-4t, 1406t
 differential diagnosis of, 266t
 epidemiology of, 245e-4t, 1407
 in HIV infection, 1257
 incubation period for, 152e-5t
 life cycle of, 245e-4t, 1407
 pathogenesis of, 1407
 in returning war veterans, 152e-5t, 152e-8
 in travelers, 855t
 treatment of, 246e-11, 1407
- Crypt stem cells, 453t
- Crystal(s)
 arthritis induced by. *See* Arthritis, crystal-induced
 renal tubular obstruction by, 1858–1859
 urinary, 1868f, 1869. *See also* Nephrolithiasis
- Crystalluria, in ethylene glycol poisoning, 64e-9
- Csendes' procedure, 1926
- CSF. *See* Cerebrospinal fluid (CSF)
- CSF3R gene mutations, 135e-5
- CSPN (cryptogenic sensory and sensorimotor polyneuropathy), 2691
- CSTB gene mutations, 2545t
- CT. *See* Computed tomography (CT)
- CTCL (cutaneous T-cell lymphoma).
See Mycosis fungoides
- Ctenocephalides felis*, 1158
- CTLA-4 antibodies, 102e-17, 372e-30, 498, 499f
- CTLA-4 gene mutations, 103e-22, 103e-22f, 2125
- CTLA-4 protein, 372e-31
- CTLs (CD8+ cytolytic T lymphocytes), in HIV infection, 1229–1230, 1237, 1244
- CTNNT1 gene mutations, 101e-2t, 2304, 2305t
- CTRC gene mutations, 2092
- C-type lectin receptors (CLRs), 372e-5t
- C-type lectins, 372e-6t
- CTZ (chemoreceptor trigger zone), 61, 258
- Cubilin, 640
- Culex* mosquitoes, 1315, 1317, 1418t
- Culicoides paraensis*, 1318
- Culiseta* mosquitoes, 1316
- Cullen's sign, 2092
- Cultural competence, 16e-7
- Cultured stem cells, 453, 453t
- Culture-independent analysis, 86e-1t, 86e-2–3, 86e-3f
- Curanderismo, 14e-1t
- CURB-65 criteria, for pneumonia, 806–807
- Curium exposure, 263e-6t, 263e-7
- Curling's ulcer, 1930
- Curvularia* spp., 228
- Cushing's disease
 vs. Cushing's syndrome, 2313
 definition of, 2313
 pathogenesis of, 2251
 screening tests for, 2262t
 skin manifestations of, 356t
 treatment of, 2318
- Cushing's response, 1467
- Cushing's syndrome, **2271**, **2313**
 in Carney's complex, 2343
 caused by ectopic ACTH production. *See* Adrenocorticotropic hormone (ACTH), ectopic production of
 clinical features of, 2271, 2272t, 2315–2316, 2315t
 bleeding, 403
 cutaneous, 359, 367, 436e-2f, 2315f
 hirsutism, 332
 hypertension, 1619
 hypokalemia, 64e-9
 lipoprotein disorders, 2439
 obesity/abdominal fat, 415e-4, 436e-2f, 2315f
 vessel wall disorders, 732
 vs. Cushing's disease, 2313
 diagnosis of, 415e-4, 1619, 2271–2272, 2316–2317, 2316f, 2317f
 differential diagnosis of, 2272t, 2316f, 2317, 2317f
 epidemiology of, 2313–2314
 etiology of, 2271, 2314–2315, 2314t
 genetic considerations in, 2318t
 GI-NETs and, 564
 glucocorticoid withdrawal and, 103e-15
 hormonal rhythms in, 400e-6
 inferior petrosal venous sampling in, 2272
 laboratory evaluation of, 2271–2272
 in lung cancer, 511
 periodic, 2343
 pNETs and, 571
 prevalence of, 2271
 in Sertoli-Leydig tumors, 594
 treatment of, 64e-9, 2272–2273, 2273f, 2316f, 2318
- Cushing's ulcer, 1930
- Cutaneous CD30+ T-cell lymphoma, 135e-4
- Cutaneous diphtheria, 977, 978, 978f, 979
- Cutaneous emboli, 361t, 362
- Cutaneous larva migrans. *See* Larva migrans, cutaneous
- Cutaneous leukocytoclastic angiitis, 2119, 2191
- Cutaneous lymphoid hyperplasia (pseudolymphoma), 353, 364t, 365, 366
- Cutaneous reflexes, 154, 2538–2539
- Cutaneous T-cell lymphoma. *See* Mycosis fungoides
- Cutaneous ulcers, **369**
 chronic
 hyperbaric oxygen therapy for, 477e-4t, 477e-5f, 477e-6
 pathophysiology of, 477e-3, 477e-6
 etiology of, 369, 369t, 828t, 829–830
 with fever, 127, 134t, 135
 venous, 1651, 1651f
 Cutaneous vasculitis. *See* Vasculitis
- Cutis verticis gyrata, 2245
- CVID (common variable immunodeficiency), 2111
- CVS (chorionic villus sampling), 83e-3–4, 444
- c-v wave, 51e-4
- CWD (chronic wasting disease), 453e-1t, 453e-2
- CX₃CR1, 372e-13t
- CX₃CR1 receptor, 1240t
- CXC chemokine family, 372e-14, 372e-14t
- CXCL12, 89e-1, 89e-2, 139e-1
- CXCR1, 372e-13t
- CXCR1 gene mutations, 863
- CXCR2, 372e-13t
- CXCR3-A, 372e-13t
- CXCR3-B, 372e-13t
- CXCR4 gene, 2111
- CXCR4 receptor
 in hematopoietic stem cells, 89e-1, 139e-1
 in HIV infection, 1216, 1235, 1236f
 HIV pathogenesis and, 145e-1
 ligands, cell types, and disease connection, 372e-13t
- CXCR5, 372e-13t
- CXCR6, 372e-13t
- CXCR6 receptor, 1240t
- Cyanide poisoning
 in chemical terrorism, 262e-1, 262e-1t, 262e-7
 clinical features of, 262e-7, 473e-11t
 complications of, 1782
 differential diagnosis of, 262e-7
 mechanisms of, 248, 262e-7, 473e-11t
 prognosis of, 262e-8
 treatment of, 262e-7–8, 262e-8t, 473e-11t
- Cyanocobalamin, for cobalamin deficiency, 648
- Cyanosis, **248**
 approach to the patient, 249
 central, 1442
 in cor pulmonale, 1506
 degree of, 248
 differential diagnosis of, 248
 etiology of, 248–249t
 pathophysiology of, 248–249
 peripheral (acrocyanosis), 1442
- CYB5A gene mutations, 2354t
- Cyclic adenosine 5'-monophosphate (cyclic AMP), 125, 125f
- Cyclic GMP, in male sexual function, 324, 325f
- Cyclic neutropenia, 236, 418
- Cyclic vomiting syndrome, 259, 261
- Cyclin B1, 372e-25t
- Cyclin-dependent kinase(s), 102e-1
- Cyclin-dependent kinase inhibitors, 102e-1
- Cyclobenzaprine, 119, 473e-10t, 2672
- Cyclobutane dimers, 386
- Cyclooxygenase (COX), 126, 1912, 1914f
- Cyclooxygenase (COX) inhibitors, 92–93, 126, 746f. *See also* Aspirin
- Cyclooxygenase 2 (COX-2) inhibitors. *See also* Nonsteroidal anti-inflammatory drugs (NSAIDs)
 adverse effects of, 92, 310, 1912
 in asthma, 1680
 for colorectal cancer prevention, 477
 drug interactions of, 43
 for fever, 126
 in NSAID-induced ulceration prevention, 1923–1924
 for osteoarthritis, 2232, 2232t
 for pain, 91–92
 in peptic ulcer disease, 277
 for polyposis syndromes, 539
- Cyclooxygenase 3 (COX-3), 126
- Cyclopentolate hydrochloride, for red or painful eye, 199
- Cyclophosphamide
 action of, 103e-8
 adverse effects of, 103e-8, 103e-9t, 2130t, 2132, 2163
 bladder cancer, 575, 622–623, 2182
 cardiovascular, 621, 1563
 cutaneous, 361, 379
 cystitis, 622–623, 1798
 infertility, 623, 1838, 2365
 leukopenia, 2182

- major, 2182t
 nail disorders, 380
 nausea and vomiting, 103e-2
 neurologic, 1793
 ovarian failure, 2132
 pulmonary, 621
 for Behçet's syndrome, 2194
 for breast cancer, 530
 for bullous pemphigoid, 372
 for Burkitt's lymphoma, 706
 for cerebellar degeneration, 618
 for CLL, 703
 for cold agglutinin disease, 659
 drug interactions of, 2130t
 for eosinophilic granulomatosis with polyangiitis, 1685, 2187
 for Ewing's sarcoma, 119e-4
 for gastric lymphoma, 536
 for gestational trophoblastic disease, 124e-5, 597
 for granulomatosis with polyangiitis, 2184-2185
 hepatotoxicity, 2025
 in HSCT preparation, 139e-2
 for inflammatory myopathies, 2201
 for interstitial lung diseases, 1711
 for lung cancer, 522
 for lupus nephritis, 1838
 for malignant pheochromocytoma, 2331
 for MS, 2672
 for mucous membrane pemphigoid, 374
 for myasthenia gravis, 2705
 for paraneoplastic encephalitis, 617
 for pemphigus vulgaris, 371
 for photoallergy, 390
 in pregnancy, 124e-4
 for SLE, 2130t, 2132
 for systemic sclerosis, 2163
 for testicular cancer, 591
 for thrombotic thrombocytopenic purpura, 730
 for vasculitis, 2182
- Cyclops*, 1412, 1422, 1435
 Cyclosarin, 262e-4. *See also* Nerve agents
 Cycloserine, 96e-4, 205e-6, 1116t
Cyclospora cayentanensis/cyclosporiasis
 clinical features of, 266, 1407
 diagnosis of, 245e-1, 245e-4t, 245e-6t, 1406t, 1407
 epidemiology of, 266, 1407
 treatment of, 1408
- Cyclosporine
 action of, 1828, 1829t, 1960, 2071
 adverse effects of, 348t, 667, 1829t, 1961, 2071
 carcinogenicity, 477t
 gingival hyperplasia, 237
 gout, 2218t
 hirsutism, 332, 380
 hyperkalemia, 310
 hyperperfusion syndrome, 463e-2f
 hypertension, 1617t, 2705
 myopathy, 2218t
 nephrotoxicity, 1828-1829, 1856t, 1862, 1864, 2071, 2705
 neurologic, 1793
 osteoporosis, 2492
 seizure, 2547t
 seizures, 1961
 skin cancer, 388
 thrombocytopenia, 729
 for angioedema, 2120
 for aplastic anemia, 666-667
 for atopic dermatitis, 354t
 for Behçet's syndrome, 2194
 contraindications to, 348
 drug interactions of, 34t
 azithromycin, 246e-4t
 azoles, 246e-3t
 calcium channel blockers, 42
 chloroquine, 246e-1t
 digoxin, 42t
 St. John's wort, 14e-4
 statins, 2199
- for epidermolysis bullosa acquisita, 374
 for IBD, 1960-1961
 for immunosuppression
 after kidney transplantation, 1828, 1829t
 after liver transplantation, 2071
 for inflammatory myopathies, 2201
 metabolism of, 34t, 42
 for myasthenia gravis, 2705
 for paroxysmal nocturnal hemoglobinuria, 662
 P-glycoprotein and, 34t
 for photoallergy, 390
 for psoriasis, 348, 348t, 354t
 for psoriatic arthritis, 2177
 for SJS/TEN, 383
 for SLE, 2133
 therapeutic monitoring of, 2764t
 for urticaria, 2120
- Cyclothymic disorder, 2717
 CYP1A1, 507
 CYP2C9
 in drug metabolism pathways, 34t
 drug responses and, 36, 39t, 41, 42
 warfarin dose requirements and, 754, 755, 755t
- CYP2C19
 in drug metabolism pathways, 34t
 drug responses and, 39t, 40, 277, 742-743, 747
- CYP2D6
 in autoimmune hepatitis, 372e-24t
 in drug metabolism pathways, 34t, 38f
 drug responses and, 34t, 39t, 40, 42
- CYP3A, 34t, 42, 277, 742
 CYP3A4, 40, 42, 935
 CYP3A5, 39t, 40
 CYP7A1 gene mutations, 2077
 CYP11A1, 2311
 CYP11A1 gene mutations, 2353, 2354t
 CYP11B1 gene mutations, 306, 434, 2356t
 CYP11B2 gene mutations, 306, 434, 2357
 CYP17 (17 α -hydroxylase) gene mutations, 372e-24t, 2311, 2353, 2354t
- CYP19 gene mutations, 2356t, 2357, 2366
 CYP21A1P pseudogene, 434
 CYP21A2 (21-hydroxylase gene) mutations
 in CAH, 2327, 2328t, 2355, 2356t
 gene conversion, 434
- CYP11A (cholesterol side-chain cleavage enzyme), 372e-24t
- Cyproheptadine
 for angioedema, 2119
 for carcinoid syndrome, 566
 for Cushing's syndrome, 2273
 for cyclic vomiting syndrome, 261
 for ergot alkaloid overdose/poisoning, 473e-8t
 for migraine prevention, 2593
 for serotonin syndrome, 473e-13t
 for urticaria, 2119
- Cyproterone acetate
 adverse effects of, 7e-6, 334
 for hirsutism, 334
 for hot flushes with androgen deprivation therapy, 7e-6
 for polycystic ovarian syndrome, 337
- CysLT (cysteinyl leukotrienes), 2114
 CysLT₁ receptor antagonist. *See* Montelukast; Zafirlukast
- Cyst, 340t. *See also specific types and sites*
 Cystathionine metabolism disorders, 434e-2t
 Cystathionine synthase, 642f, 643
 Cystathioninuria, 434e-2t
 Cystatin C, 291, 2759t
 Cystectomy, for bladder cancer, 576-577, 577t
 Cysteinyl-leukotrienes, 1678
 Cystic duct stump syndrome, 2082
- Cysticercosis
 chronic meningitis in, 908t
 clinical features of, 1430-1431
 diagnosis of, 245e-6t, 1431-1432, 1431t
 etiology of, 902
 neurologic. *See* Neurocysticercosis
 treatment of, 246e-1, 1432
- Cystic fibrosis (CF), 1697
 bronchiectasis in, 1695
 chest radiograph in, 308e-11f
 clinical features of
 clubbing, 2245
 gastrointestinal, 269, 1697
 hemoptysis, 245
 hepatic, 367e-2, 1697, 2062
 other organ systems, 1697
 pancreatic, 1697
 respiratory, 1697
 sinusitis, 226
 diagnosis of, 1698-1699
 disease spectrum of, 1699
 genetic considerations in
 CFTR gene mutations, 145e-5, 429f, 435, 480e-5, 1697-1698, 1698f
 heterozygote advantage, 442
 genetic testing for, 443, 450t
 incidence of, 1699
 infections in
Aspergillus, 1686
 nontuberculous mycobacterial, 776-777, 1128
P. aeruginosa, 145e-4f, 145e-5, 1044
 transmission of, 776-777
 newborn screening for, 1700
 pathogenesis of
 airflow resistance, 306e-3
 autoantigens, 372e-24t
 CFTR deficiency, 1697, 1698f, 1700v
 immunodeficiency, 375e-2
 pulmonary inflammation and remodeling, 1697
 treatment of
 CFTR defect correction, 1699
 early interventions for, 450t
 effectiveness of, 1700
 ivacaftor, 1699
 lung transplantation, 320e-1t, 320e-2
 personalized molecular therapies, 1699
 sequelae management, 1699
 typhoid fever resistance in, 145e-3
 vitamin E deficiency in, 96e-8
- Cystic fibrosis transmembrane conductance regulator. *See* CFTR protein
- Cystic medial necrosis, 1638
 Cystine metabolism disorders, 434e-2t
 Cystine stones, 1868, 1869f, 1871. *See also* Cystinuria; Nephrolithiasis
- Cystinosis, 434e-2t, 435e-1t
 Cystinuria
 clinical features of, 435e-1t, 435e-2, 1868, 1937
 genetic considerations in, 332e-7t, 332e-8, 435e-1t, 435e-2
 non-type I, 332e-7t
 treatment of, 435e-2
 type I, 332e-7t
- Cystitis. *See also* Urinary tract infections (UTIs)
 alkaline-encrusted, 981
 bacterial, 60e-1
 chemotherapy-induced, 103e-8
 clinical features of, 60e-1, 863
 definition of, 861
 diagnosis of, 865, 865f
E. coli, 60e-1, 1028
 epidemiology of, 861
 hematuria in, 294
 hemorrhagic, 139e-3, 1208, 1798
 interstitial. *See* Interstitial cystitis/bladder pain syndrome
 in men, 866
 prognosis of, 868
 recurrent, 868, 945t
 sexually transmitted, 870t
 treatment of, 60e-1, 866-867, 867t
- Cystoisospora belli*/cystoisosporiasis
 clinical features of, 266, 765t, 1407, 1957
 diagnosis of, 245e-4t, 245e-5t, 1406t, 1407
 epidemiology of, 245e-4t, 765t
 in HIV infection, 1257, 1407, 1957
 life-cycle hosts of, 245e-4t
 treatment of, 246e-9, 246e-12, 1407
- Cystopathy, diabetic, 2427

- Cystoscopy, 60e-4, 587
- Cytarabine
action of, 682
adverse effects of, 380, 607t, 682–683, 2686t
for AML, 682–683, 684
for neoplastic meningitis, 1793
for primary central nervous system lymphoma, 602
- Cytidine deaminase, 103e-23
- Cytoadherence, 1371
- Cytochrome b5 (CYB5A), 2353
- Cytochrome b5 reductase deficiency, 654t
- Cytochrome c, 102e-8–9
- Cytochrome c oxidase deficiency, 85e-3–4, 85e-5f, 85e-7
- Cytochrome oxidase inhibitors, 473e-11t
- Cytochrome P450 monooxygenase, 33
- Cytogenetic analysis
molecular, 83e-2, 83e-2f. *See also* Fluorescence in situ hybridization (FISH)
postnatal indications for, 83e-4t
prenatal indications for, 83e-3–4, 83e-4t
standard, 83e-1, 83e-1f
- Cytogenomic analysis
comparative genomic hybridization, 83e-3, 83e-3t
indications for, 83e-3–4, 83e-4t
SNP array, 83e-2f, 83e-3, 83e-3t. *See also* Single-nucleotide polymorphisms (SNPs)
- Cytokeratins
in carcinoma of unknown primary, 120e-1, 120e-2, 120e-2f, 120e-2t
in lung cancer, 508
- Cytokine(s), 372e-9
in asthma, 1673–1674
in atherosclerosis, 291e-2
in autoimmune and inflammatory diseases, 372e-25t, 372e-27t, 372e-30
biologic activity of, 372e-10–13t
definition of, 372e-1
expression of, 2114–2115
of innate immune system, 372e-2, 372e-4t
production in CNS, 125
proinflammatory
in anemia, 630, 630f
in asthma, 1673–1674
cellular signaling pathways for, 145e-5, 145e-6f
in HIV infection, 1234–1235
in host response to pathogens, 145e-10, 372e-26
in metabolic syndrome, 2452
pyrogenic, 124–125, 125f
receptors of, 372e-10–13t
secretion of, 372e-14f
in sepsis/septic shock, 1753
signaling deficiency, in SCID, 2108
sources and targets of, 372e-10–13t
structural families of, 372e-14t
in thyroid-associated ophthalmopathy, 2294
in thyroid hormone synthesis, 2285
- Cytokine blockers, 377e-5
- Cytokine receptor–linked kinase, 400e-1t
- Cytokine receptors, 372e-15–16, 400e-3–4, 400e-3f. *See also specific receptors*
- Cytokine release syndrome, 1796
- Cytokine storm, 956
- Cytokine therapy, 103e-22, 487, 492
- Cytomegalovirus (CMV), 214e-4, 214e-9, 1190
- Cytomegalovirus (CMV) immune globulin, 1193
- Cytomegalovirus (CMV) infections, 1190
in burn patient, 166e-3
in cancer patient, 489, 489t, 492
congenital, 50, 1191, 1191t
diagnosis of, 150e-7–8, 1192–1193
diarrhea in, 266
differential diagnosis of, 1190t
epidemiology of, 1190–1191
esophagitis, 1896f, 1910
graft-versus-host disease and, 921–922
in HIV infection, 1191t, 1192, 1252t, 1257, 1267
in liver allograft, 366e-3f
in lung allograft, 320e-4
lymphadenopathy in, 408
mononucleosis, 1191–1192
nausea and vomiting in, 258
neuropathy in, 2685
pathogenesis of, 1191
pathology of, 1191
perinatal, 1191
pharyngitis, 231
pneumonia, 1192
in pregnancy, 50
prevention of, 1193
proctitis, vs. IBD, 1956
retinitis
clinical features of, 1192, 1192f
in HIV infection, 1267
treatment of, 215e-7, 215e-8, 1193
skin manifestations of, 362, 362t
transfusion-transmitted, 138e-6, 1193
in transplant recipient
after heart transplant, 926
after HSCT, 920t, 921–922, 921t, 1191t
after kidney transplant, 925–926
after lung transplant, 927
clinical features of, 1191t, 1192
herpesviruses and, 927
pathophysiology of, 921, 926
prevention of, 1191t
prophylaxis for, 139e-4, 139e-4t, 215e-8, 1831
sites of, 924t
treatment of, 320e-4, 922, 926, 1191t, 1831
treatment of, 215e-2t, 215e-7–9, 1193–1194, 2657
- Cytoplasmic antineutrophil cytoplasmic antibody (c-ANCA), 377e-4, 2180
- Cytosine, 429, 431f
- Cytosine arabinoside, 103e-11t, 103e-14, 103e-25, 620t, 649
- Cytoskeletal proteins, 1553, 1555t, 1556f
- Cytotoxic edema, 1777
- D**
- d4T (stavudine), 1273, 1276f, 1279f
- Dabigatran
adverse effects of, 757, 1488
dosage of, 757
drug interactions of, 42
for DVT/PE, 1635
indications for, 379, 756–757
management before endoscopic procedures, 1889t
monitoring treatment with, 407, 757
pharmacology of, 756t
in pregnancy, 757
for stroke prevention in atrial fibrillation/atrial flutter, 1485t, 1488, 2570
for VTE prophylaxis, 1637t
- Dabrafenib
action and targets of, 102e-5t, 103e-16f, 103e-17t, 103e-19
adverse effects of, 103e-17t
for melanoma, 103e-17t, 103e-19, 498, 499
- Dacarbazine (DTIC)
action of, 103e-12
adverse effects of, 103e-9t, 103e-12, 103e-25, 389t, 474, 607t
for malignant pheochromocytoma, 2331
for melanoma, 499
for soft tissue sarcoma, 119e-2
- Daclatasvir, 2049
- Daclizumab, 1517
- Dacryocystitis, 199
- Dacryocytes, 81e-1
- Dactinomycin
adverse effects of, 103e-9t
for Ewing's sarcoma, 119e-4
for gestational trophoblastic disease, 124e-5, 597
- Dactylitis
characteristics of, 829, 2176f, 2220
etiology of, 829, 2220
in psoriatic arthritis, 2176, 2176f
in reactive arthritis, 2174
in sickle cell disease, 2242
- DADs (delayed afterdepolarizations), 273e-3, 273e-3f
- Dakin's solution, 262e-4
- Dalbavancin
action of, 930, 931t, 961
adverse effects of, 938t, 942
indications for, 940t, 942
in pregnancy and lactation, 937t
resistance to, 931t, 933–934
for staphylococcal infections, 961, 962t
- Dalfampridine, 2672
- Dalfopristin. *See* Quinupristin-dalfopristin
- Dalteparin, 1635t, 1637t
- DALY (disability-adjusted life years), 9, 10f, 11t
- Damage-associated molecular patterns (DAMPs), 145e-8f, 377e-2, 1747, 2227f
- Damocles syndrome, 475
- Danaparoid, 728
- Danazol, 225e-5, 727t
- Dandruff, 1357
- Danon's disease (LAMP2 deficiency), 432e-4t, 433e-5, 1566
- Dantrolene, 2660, 2672
- Dapagliflozin, 2414t
- Dapsone
for acne vulgaris, 352
adverse effects of
frequent, 246e-4t
in G6PD deficiency, 656, 656t, 1126, 1262
hemolysis, 373
hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
methemoglobinemia, 373, 1384
neuropathy, 2687t
occasional, 246e-4t, 1126
for dermatitis herpetiformis, 373
drug interactions of, 246e-4t
for epidermolysis bullosa acquisita, 374
for leprosy, 1126–1127, 1127t
for linear IgA disease, 373
for lupus dermatitis, 2134
for mucous membrane pemphigoid, 374
overdosage/poisoning with, 473e-11t
for PCP, 1362t
for PCP prophylaxis, 1251t, 1362t
in pregnancy and lactation, 246e-4t
for relapsing polychondritis, 2204
for toxoplasmosis prophylaxis in HIV infection, 1251t
- Daptomycin
action of, 931t, 932f, 933, 961
adverse effects of, 938t, 942, 976
for enterococcal infections, 975t, 976
indications for, 940t, 942
for infective endocarditis, 823
for MRSA infections, 962t, 963
for osteomyelitis, 840t
resistance to, 931t, 934, 940t, 961
for staphylococcal infections, 961, 962t
- Darbepoetin alfa, 631
- DARC receptor, 1240t
- Darier's sign, 367, 2120
- Darifenacin, 81
- Darmbrand (enteritis necroticans), 991
- Dartmouth Atlas of Health Care, 12e-3
- Darunavir
adverse effects of, 1275t, 1281
for HIV infection, 1275t, 1276f, 1280–1281
resistance to, 1280f
- DAS181, for influenza, 215e-6
- Dasatinib
action and targets of, 102e-3, 102e-5t, 103e-16, 103e-17t
adverse effects of, 103e-17t, 103e-26, 692
for CML, 102e-3, 102e-5t, 691–692, 691t
genetic variations in response to, 39t, 103e-16
- DASH diet, 1622t, 1623
- Daunorubicin
action of, 103e-13
adverse effects of, 103e-10t, 103e-13, 354, 379
for AML, 682, 683
drug interactions of, 103e-10t
liposomal, for Kaposi's sarcoma, 1270
David procedure, 1538f

- DAX1, 401e-1, 401e-1t, 2310
DAX1 gene mutations, 435, 2349, 2354t, 2364
 Day-care centers, 83–84, 83t, 266
 Daytime sleepiness, 188, 188t, 1726
DAZ gene mutations, 437
 DBS. *See* Deep brain stimulation (DBS)
 DBT (dialectical behavior therapy), 2720
 DCA (dichloroacetate), 102e-13
DCC gene, 538
 DC-CK1/PARC, 372e-12t
 DCP (des- γ -carboxy prothrombin), 546, 547
 DC-SIGN, 1216
 DDAVP. *See* Desmopressin (DDAVP)
 ddC (zalcitabine), 1276f
 ddI (didanosine), 1276f, 1278, 1279f, 2023, 2030
DDR2 gene mutations, 507t, 509
 Deafness. *See* Hearing loss
 Death
 among international travelers, 793
 from cancer, 467, 467t, 469f, 470t
 cardinal signs of, 68–69
 childhood, 9
 infectious disease–related, 761, 762f
 leading causes of, 6e-1
 age-specific, 30–31t
 cardiovascular disease, 266e-1, 266e-3, 266e-3f, 266e-3t
 epidemiologic transition in, 266e-1
 in men, 7e-1
 by sex, 6e-1t
 in United States and Britain, 56t
 in women, 6e-1t, 6e-2f
 worldwide, 9, 11t, 266e-2f
 rates of, by age group, 94e-1f
 site of, 55–56, 56f
 of spouse, 69
 sudden cardiac. *See* Sudden cardiac death
 sudden natural, 1764
 Death adders, 2734
 Death rattle, 68, 69t
 Death receptors, 102e-8–9, 102e-9f, 102e-10, 372e-23, 372e-26f
 Debranching enzyme deficiency (type IIIa/b GSD), 433e-2t, 433e-3, 436e-6f, 462e-14
 DEC. *See* Diethylcarbamazine (DEC)
 Deception, 17e-1
 Decerebrate rigidity, 1773–1774
 Decision analysis, 24, 24f
 Decision limits, 480e-3
 Decision-making. *See* Clinical reasoning
 Dectabine
 action of, 103e-20
 adverse effects of, 671
 for AML, 683
 for myelodysplasia, 671
 for sickle cell anemia, 636
 Declaration of Alma Ata, 13e-1
 Decompression illness, 477e-4t
 Decompression sickness, 477e-4t, 477e-7–8, 477e-7t
 Deconditioning, 47e-1t, 47e-2, 47e-2t
 Decongestants
 adverse effects of, 1617t
 for nasal congestion, 2122, 2123f
 for otitis media, 230
 overdosage/poisoning with, 473e-3, 473e-8t
 for rhinovirus infections, 1203
 for sinusitis, 227
 for upper respiratory infections, 226
 Decorporation, 263e-6
 Decorticate rigidity, 1773–1774
 Decremental conduction, 1471
 Dectin-1, 1342, 1752
 Decubitus (pressure) ulcers, 830, 1743, 2660
 Deep brain stimulation (DBS)
 for depression, 2717
 for dystonia, 2621
 for headache, 2596
 for obsessive-compulsive disorder, 2714
 for PD, 2616–2617
 Deep breathing exercises, preoperative, 55
 Deep neck infections, 235
 Deep vein thrombosis (DVT), 1631
 classification of, 1632
 in critically ill patient, 1734
 diagnosis of
 blood tests, 1632
 clinical decision rules, 1632t
 clinical evaluation, 1632
 clinical pearls, 1632
 imaging algorithms, 1632f, 1634f
 MRI, 1634
 venous ultrasonography, 1633, 1633f
 differential diagnosis of, 1632, 1633t
 epidemiology of, 743
 history in, 404–405
 in HIV infection, 1262
 paraneoplastic, 612t, 613
 perioperative prophylaxis of, 55
 postmenopausal hormone therapy and, 2384t
 in pregnancy, 47
 prevention of, 1634, 1636–1637, 1637t, 1734, 1743, 1757–1758, 1785
 risk factors for, 404–405, 405f
 treatment of, 1634
 anticoagulation, 1634–1635, 1635t
 duration of anticoagulation, 1635–1636
 inferior vena cava filter, 1636
 in pregnancy, 47
 Deepox, 220e-1t
 Deer tick. *See* *Ixodes* spp. ticks
 DEET-containing products, 796
 Defecation, 265, 265f
 Defecography, 274
 Defensins, 415
 Defensive medicine, 20
 Deferasirox, 639, 1353, 2518
 Deferoxamine, 473e-11t, 639, 2518
 Defibrillation, 1769
 Defibrillator
 automated external, 1769
 implantable. *See* Implantable cardioverter-defibrillator (ICD)
 Deficiency of the IL-1 receptor antagonist (DIRA), 2215
 Deficiency of the IL-36 receptor antagonist (DITRA), 2215
 Deformity, joint, 2219t, 2220
 Degarelix, 585
 Deglutitive inhibition, 255
 Dehalogenase 1, 2284t
 Dehydration
 in cholera, 1063–1064, 1064t
 delirium in, 169
 in diarrhea, 267
 in hypothermia, 478e-2
 mechanisms of, 332e-9, 332e-10f
 in terminally ill patient, 69t
 Dehydroemetine. *See* Emetine
 Dehydroepiandrosterone (DHEA)
 for adrenal insufficiency, 2327
 adverse effects of, 2130t
 in hirsutism, 332
 reference values for, 2759t
 for SLE, 2130t
 synthesis of, 2309f
 Deiodinase(s), 2283f, 2286
 Déjérine-Roussy syndrome, 161, 2575, 2583
 Delamanid, 205e-7, 1115, 1119
 Delavirdine, 1276f, 1278
 Delayed afterdepolarizations (DADs), 273e-3, 273e-3f
 Delayed puberty
 anosmia in, 215, 2380
 female, 2380–2381, 2381t
 male, 2361t, 2362–2363
 Delayed sleep-wake phase disorder, 193
 Delayed-type hypersensitivity reaction, 372e-29–30, 377–378, 1107
 Deletion
 chromosome, 83e-1, 83e-6
 gene, 432
 Delirium, 166
 approach to the patient, 167–170
 assessment of, 64, 167, 167t
 clinical features of, 166
 in critically ill patient, 166, 170, 1735, 1782
 dementia and, 79, 166
 diagnosis of, 169–170, 169t
 drug-induced, 167, 168t, 1606
 in elderly, 79, 79f, 166
 epidemiology of, 166
 etiology of, 64, 168–169, 168t
 at high altitudes, 476e-5
 history in, 167
 pathogenesis of, 166–167
 physical examination in, 168–169
 postoperative/ICU, 1735
 prevention of, 170
 reversibility of, 166
 risk factors for, 79, 166
 in sepsis/septic shock, 1755
 in terminally ill patient, 64, 64t
 treatment of, 64–65, 65t, 170, 1782
 Delirium tremens, 166, 2727, 2728
 Delta fraction, 280
 Delta receptors, 468e-1t
 Deltaretroviruses. *See* Human T-cell lymphotropic virus (HTLV-1)
 Delusional infestations, 2750–2751
 Delzicol, 1963
 Demecarium bromide, 197
 Demeclocycline
 adverse effects of, 64e-5, 302
 for SIADH, 64e-5, 302, 511, 610, 2282
 Dementia, 170
 aging and, 94e-1
 in alcoholism, 2607
 in ALS/parkinsonian/dementia complex of Guam, 2608
 aphasia in, 179–180
 approach to the patient, 171–175
 in chronic traumatic encephalopathy, 2608
 clinical features of, 173, 173t
 in CNS infections, 2607
 in CNS neoplasms, 2607
 in CNS vasculitis, 2607
 in cobalamin deficiency, 2607
 cognitive examination in, 173–174
 definition of, 170
 degenerative
 Alzheimer's disease. *See* Alzheimer's disease
 Creutzfeldt-Jakob disease. *See* Creutzfeldt-Jakob disease (CJD)
 dementia with Lewy bodies. *See* Dementia with Lewy bodies
 frontotemporal temporal dementia. *See* Frontotemporal dementia (FTD)
 molecular basis of, 172t
 types of, 172
 delirium and, 166
 diagnosis of, 172t, 174
 differential diagnosis of, 171t
 EEG in, 442e-2
 etiology of, 171, 171t, 2606–2608
 familial, 451e-2t
 functional anatomy of, 170
 history in, 171–172, 172t
 HIV-associated, 1263, 1264t, 1265f
 in Huntington's disease, 2606
 imaging in, 174f, 175f
 insomnia in, 191
 intracranial hypotension and, 2607
 in Korsakoff's syndrome, 2607
 metal exposure and, 2607–2608
 multi-infarct, 2602
 neurologic examination in, 173, 173t
 in normal-pressure hydrocephalus, 175f, 2606–2607
 in PD. *See* Parkinson's disease dementia (PDD)
 in pellagra, 2607
 physical examination in, 172t, 173
 postmenopausal hormone therapy and, 2384t, 2385–2386
 prevalence of, 170
 in psychiatric diseases, 2608
 in seizure disorders, 2607
 semantic, 182
 in substance abuse, 2608

- Dementia (*Cont.*):
 in thiamine deficiency, 2607
 transient global amnesia and, 2608
 treatment of, 175–176
 vascular, 170, 172, 173t, **2602**
- Dementia pugilistica. *See* Chronic traumatic encephalopathy
- Dementia with Lewy bodies
 AD and, 2606
 clinical features of, 166, 172, 173, 173t, 2606
 vs. Creutzfeldt-Jakob disease, 453e-4–5
 molecular basis of, 172t
 neuropathology of, 2606
 olfactory dysfunction in, 214–215
 vs. Parkinson's disease dementia, 2617
 prions in, 444e-8f
 treatment of, 2606
- Demethylation, of DNA, 102e-8
- Demographic Transition, 93e-1
- de Montagnana, Petrus, 2f
- Dendritic cells
 in asthma, 1672
 definition of, 372e-1
 functions of, 372e-5f, 372e-6, 372e-7t, 372e-14f
 hematopoietic differentiation of, 89e-3f, 372e-6, 372e-9f
 in HIV infection, 1238
 myeloid, 372e-6, 372e-7t
 neoplasms of, 135e-8–9
 plasmacytoid 1, 372e-6, 372e-7t
- Dengue fever
 climate change and, 151e-4, 151e-5f, 151e-6f, 151e-8, 151e-8f
 clinical features of, 129t, 362, 783, 1319
 diagnosis of, 1319
 epidemiology of, 134, 151e-4, 151e-5f, 151e-8, 151e-8f, 783, 1318–1319
 prevention of, 796
 in returned traveler, 802, 802t
 severe, 783, 1322
- Dengue viruses, 1306t
- Dennie-Morgan folds, 344
- Denosumab
 action of, 102e-5t, 102e-11, 2501
 adverse effects of, 2501
 for hypercalcemia, 2481
 indications for, 2501
 for metastatic bone disease, 119e-4
 for osteoporosis management/prevention, 2501, 2502f
 for Paget's disease of bone, 426e-4
 prophylactic, in prostate cancer, 586
- Dense deposit disease, 62e-4f, 1841. *See also* Membranoproliferative glomerulonephritis
- Dense-granules, 275, 741
- Dental care/procedures
 in medically complex patient, 241–242
 preventive, 236
 prophylactic antibiotics for, 242, 826–827, 826t
- Dental disease
 brain abscess and, 901
 calculus, 46e-4f, 46e-5f
 caries, 46e-4f, 236, 1097
 gingivitis. *See* Gingivitis
 involuntary weight loss in, 275
 lymphadenopathy in, 408
 periodontitis. *See* Periodontal disease (periodontitis)
 plaque, 46e-4f, 236, 1097
 root cavity, 46e-4f
- Dental pulp stem cells, 453t
- Dentatorubropallidolusian atrophy, 440t, 451e-2t, 2629
- Dentin, 236
- Dent's disease, 332e-7t, 332e-8, 2459
- Dentures, 46e-1f, 46e-2f, 242
- Denys-Drash syndrome, 2353
- Deoxycoformycin, 103e-10t, 103e-14, 135e-2
- Deoxycorticosterone (DOC), 2309f, 2313, 2319
- Deoxyhemoglobin, 632, 632f
- DEPDC5 gene mutations, 2545t
- Dependence
 alcohol. *See* Alcohol use disorder (alcoholism)
 substance. *See* Substance use disorders
- Dependency ratios, 93e-4–5
- Depilatory, 334
- Depression/depressive disorders, **2714**
 agitated, 2717
 assessment of, 63
 in cancer, 473, 524, 2714
 in cardiovascular disease, 1514, 2714
 chemosensory alterations in, 215
 in chronic disease, 2714
 clinical features of, 2714–2715, 2715t
 in diabetes mellitus, 2714
 diagnosis of, 2715t
 dizziness in, 151
 drug-related, 2714
 etiology of, 63, 2715
 headache and, 108
 in HIV infection, 2714
 in hypothyroidism, 2714
 insomnia in, 190
 involuntary weight loss in, 275
 major depression, 2715, 2715t, 2716f
 nausea and vomiting in, 259
 in neurologic disorders, 2714
 AD, 2602
 epilepsy, 2558
 MS, 2663, 2672
 PD, 2617
 pain and, 93
 pathophysiology of, 465e-3–4, 465e-4f, 2715
 persistent, 2715
 postpartum, 6e-5
 screening for, 29t
 sexual dysfunction in, 326, 330
 in terminally ill patient, 63–64
 treatment of, **2715**
 algorithm for, **2716f**
 antidepressants, 2710t, 2711t, 2716–2717
 in cerebral function disorders, 184
 St. John's wort, 14e-4
 supportive, 2715–2716
 in terminally ill patient, 63–64
 unipolar, 2715
 in war veterans, 471e-4–5, 471e-5t
 in women, 6e-5, 2715
- de Quervain's tenosynovitis, 2220–2221, 2220f, 2248
- de Quervain's (subacute) thyroiditis, 239, 2298–2299, 2298f, 2298t
- Dermacentor* spp. ticks, 1154, 1319, 2745. *See also* Tick-borne diseases
- Dermal papillae, 827f
- Dermatitis, **344**
 atopic. *See* Atopic dermatitis
 contact. *See* Contact dermatitis
 in malabsorption syndromes, 1945
 seborrheic. *See* Seborrheic dermatitis
 stasis. *See* Stasis dermatitis
- Dermatitis herpetiformis, **373**
 celiac disease and, 373, 1942
 clinical features of, **373**
 distribution of lesions, 76e-14f, 340, 343f, 370t
 papules/vesicles, 76e-14f, 370t
 vesicles/bullae, 361t
 diagnosis of, 373
 histology of, 370t, 373
 HLA gene association with, 373, 373e-8t
 immunopathology of, 370t, 373
 treatment of, 373
- Dermatofibroma, 341t, 342f
- Dermatofibrosarcoma protuberans, 119e-2
- Dermatofibrosis lenticularis disseminata (Buschke-Ollendorff syndrome), 426e-5
- Dermatographism. *See* Dermographism
- Dermatoheliosis, 387
- Dermatomes, 160f, 161f
- Dermatomes, 160f, 161f
- Dermatomyositis, **2194**
 clinical features of, 157, 374, 2194–2196, 2195t
 calcinosis, 2202
 cutaneous, 374–375
 on face, 374f
 on hands, 76e-14f, 374f
 rash, 76e-13f
 telangiectasias, 356t, 357
 myocarditis, 1562
 Raynaud's phenomenon, 1649
 diagnosis of, 375, 2199–2200, 2199t, 2200f
 differential diagnosis of, 2197–2199
 immunopathogenesis of, 372e-24t, 2196, 2197f
 interstitial lung disease in, 1714
 malignancies associated with, 2196
 prognosis of, 2202
 treatment of, 2200–2202
- Dermatomyositis sine myositis, 374, 2195
- Dermatophagoides farinae*, 2121
- Dermatophagoides pteronyssinus*, 1669, 1671, 2121
- Dermatophytosis, **349**
 clinical features of, **349**
 distribution of lesions, 341t
 papulosquamous lesions, 347t, 350t
 diagnosis of, 350, 1357
 epidemiology of, 1357
 etiology of, 349, 350t, 1357
 histologic features of, 347t
 pathogenesis of, 1357
 treatment of, 350, 350t, 1357–1358, 1358t
- Dermographism, 76e-4f, 363, 363t, 2118, 2118f
- Dermoid cysts, 594, 603
- Dermopathy, thyroid, 2295, 2295f, 2297–2298
- DES (diethylstilbestrol), 103e-15, 477t, 585
- Descending perineum syndrome, 273
- Desensitization, 384
- Desipramine
 adverse effects of, 2710t, 2716
 for depression, 473, 2672, 2710t, 2716
 for irritable bowel syndrome, 1969
 for neuropathic pain, 94
 for pain, 91t, 93, 2672
 therapeutic monitoring of, 2764t
- Desmin
 in autoimmune disorders, 372e-24t
 functions of, 1553–1554
 mutations in, 1554
- Desmogleins, 371, 372e-24t, 373e-7, 377e-4
- Desmoplakin, 372e-25t
- Desmopressin (DDAVP)
 for adult hypopituitarism, 2259t
 adverse effects of, 731
 for central diabetes insipidus, 304
 for coagulation disorders in chronic kidney disease, 1818
 for diabetes insipidus, 2278–2279, 2279f
 for gestational diabetes insipidus, 303
 for hemophilia, 734
 for hypernatremia, 304
 for hypodipsic hypernatremia, 2280
 for hyponatremia, 64e-6–7, 303
 for nephrogenic diabetes insipidus, 64e-3, 304
 for orthostatic hypotension, 146
 for platelet function disorders, 730
 for sleep enuresis, 192
 structure of, 2275f
 for von Willebrand disease, 731
- Desmosome complexes, 1555t, 1556f, 1564
- Desmoteplase, 759, 759f
- Desquamative interstitial pneumonia, 1713
- Desvenlafaxine, 2710t
- Des- γ -carboxy prothrombin (DCP), 546, 547
- Detergent worker's lung, 1681t
- Detumescence, 325, 325f
- Developing world. *See* Low- and middle-income countries
- Device-related infections. *See* Health care-associated infections
- Devic's disease (neuromyelitis optica), 2656, 2673, 2673t
- DEXA (dual-energy x-ray absorptiometry), 2067, 2472, 2493
- Dexamethasone
 action of, 936
 adverse effects of, 462e-19
 for altitude illness, 476e-2–3, 476e-2t, 476e-4
 for brain abscess, 902

- for brain metastases, 1792
- for CAH, 334, 2329
- for cerebral edema, 598
- drug interactions of, 246e-3t
- for enteric fever, 1052
- for fatigue, 63
- for hypoadrenal shock, 1751
- for increased intracranial pressure, 1780t
- for meningitis, 780t, 889, 890, 1011
- for multiple myeloma, 716-717
- for nausea and vomiting, 62, 103e-25, 474
- for neurocysticercosis, 903
- for pain, 61
- as premedication for taxanes, 103e-15
- for prenatal treatment of CAH, 2356
- for spinal cord compression, 2653
- for tuberculous meningitis, 899
- Dexamethasone suppression test, 332, 415e-4, 2310
- Dexlansoprazole, 263, 1920, 1920t
- Dexmedetomidine, 1782
- Dexrazoxane, 103e-13, 621
- Dextroamphetamine, 61, 63, 190
- Dextrocardia, 269e-16f, 1528
- Dextromethorphan, 245, 473e-13t
- Dextrose, 298, 304, 312
- DF (discriminant function) value, 2059
- DGM (diphosphoglycerate mutase) deficiency, 654t
- DHCR7* gene mutations, 2354t
- DHEA. *See* Dehydroepiandrosterone (DHEA)
- DHF (dihydrofolate) reductase, 641, 642f, 931t, 934
- DHF (dihydrofolate) reductase inhibitors, 246e-4t, 246e-12
- DHH* gene mutations, 2354t
- Dhori virus, 1310t
- DHPR (dihydropteridine reductase) deficiency, 434e-2t
- DHT (dihydrotestosterone), 332, 2358, 2359f. *See also* Testosterone
- Diabetes insipidus (DI), **2275**
 - adipsic, 304
 - differential diagnosis of, 2277-2278, 2278f
 - etiology of, 2275-2276, 2277t
 - gestational, 303, 2276
 - nephrogenic
 - clinical features of, 64e-3, 294-295, 303
 - diagnosis of, 294f, 304
 - early interventions for, 450t
 - etiology of, 294f, 295, 297, 303, 2276, 2277t
 - genetic considerations in, 332e-7t
 - genetic testing for, 450t
 - lithium-associated, 64e-3-4
 - pathogenesis of, 303
 - treatment of, 64e-4, 304
 - pathophysiology of, 2276-2277
 - pituitary (central, neurohypophyseal)
 - clinical features of, 294-295, 303
 - diagnosis of, 294f, 304
 - early interventions for, 450t
 - etiology of, 294f, 295, 297, 2277t
 - genetic considerations in, 435
 - genetic forms of, 2276, 2277t
 - genetic testing for, 450t
 - treatment of, 304
 - treatment of, 2278-2279, 2279f
- Diabetes mellitus (DM), **2399**
 - abdominal pain in, 106, 106t
 - anemia in, 398
 - approach to the patient, 2406-2407
 - celiac disease and, 1942
 - classification of, 2399-2400, 2399f, 2399t, 2407
 - complications of, **2422**, 2422t
 - cardiovascular. *See* Cardiovascular disease, in diabetes mellitus
 - cutaneous, 2429-2430
 - detection and prevention of, 2408, 2408t
 - diabetic ketoacidosis. *See* Diabetic ketoacidosis (DKA)
 - dyslipidemia, 2428
 - erectile dysfunction, 326, 2427
 - female sexual dysfunction, 330, 2427
 - gastrointestinal, 258, 2427
 - genitourinary, 862, 2427
 - glycemic control and, 2422-2423, 2423f
 - hyperglycemic hypermolar state, 2417, 2417t, 2420
 - hyperkalemia, 309
 - hypertension, 2428
 - hypoglycemia. *See* Hypoglycemia, in diabetes mellitus
 - infections, 2429
 - lactic acidosis, 318
 - lower extremity
 - osteomyelitis, 845, 845f
 - pathophysiology of, 839, 2428
 - prevention of, 2428-2429
 - treatment of, 477e-4t, 2429
 - mechanisms of, 2423-2424
 - neuromuscular involvement, 462e-19
 - neuropathy. *See* Diabetic neuropathy
 - ocular
 - abducens nerve palsy, 209
 - cataracts, 204
 - epiretinal membrane, 206
 - retinopathy. *See* Diabetic retinopathy
 - tonic pupil, 196
 - vitreous degeneration, 204
 - renal. *See* Diabetic nephropathy
 - retinopathy. *See* Diabetic retinopathy
 - contrast agent risk in, 1461, 2425
 - cystic fibrosis-related, 1697
 - deaths from, 6e-1t, 56t
 - definition of, 2399
 - depression in, 2714
 - in developing countries, 15, 266e-4
 - diagnosis of, 2401, 2401f, 2401t
 - diarrhea in, 270, 272, 855t
 - in elderly, 82
 - epidemiology of, 2400
 - erythropoietin deficiency in, 398
 - etiology of, 2399-2400, 2400t
 - frailty and, 77, 82
 - fulminant, 2400
 - genetic considerations in, 431t, 441, 441t, **2404**
 - gestational, 6e-3-4, 48, 2400, 2422
 - global considerations in, 15, 266e-4, 2400, 2400f
 - gut microbiota and, 86e-8-9
 - HCV infection and, 2041
 - in hemochromatosis, 2516
 - high-altitude travel and, 476e-5
 - history in, 2406
 - immunization recommendations for, 788f
 - incidence of, 2400
 - in Kearns-Sayre syndrome, 462e-15
 - latent autoimmune in the adult, 2407
 - lipodystrophic, 2422
 - maturity-onset of the young (MODY), 431t, 441t, 2400, 2406
 - nausea and vomiting in, 258
 - obstructive sleep apnea and, 1724
 - oral manifestations of, 236
 - otitis externa in, 228-229
 - pancreatic cancer and, 554
 - parenteral nutrition in patient with, 98e-7, 2421
 - physical examination in, 2406-2407
 - polyuria in, 295
 - in pregnancy, 47-48
 - preoperative evaluation and management of, 55
 - prevalence of, 2400, 2400f
 - prevention of, 17
 - screening for, 29t, 84, 2401t, 2402
 - skin manifestations of, 359, 361t, 362, 363, 365, 436e-4f, 2429-2430
 - stroke risk in, 2567t
 - systems biology applied to, 87e-3t
 - taste disorders in, 215
 - treatment and management of, **2407**
 - blood glucose monitoring in, 2409-2410
 - complication detection and prevention, 2408, 2408t
 - in elderly, 82
 - emerging therapies, 2417
 - exercise, 2408
 - during glucocorticoid therapy, 2421-2422
 - glycemic control monitoring, 2410
 - glycemic control targets, 2410
 - goals and principles of, 2407-2408, 2408t
 - in hospitalized patients, 2420-2421
 - medical nutrition therapy in, 2408-2409, 2409t
 - multidisciplinary team in, 2407-2408
 - ongoing aspects of, 2407-2408, 2408t
 - outpatient, improvement of, 12e-3
 - during parenteral nutrition therapy, 2421
 - patient education in, 2408-2409
 - in pregnancy, 2422. *See also* Gestational diabetes mellitus
 - stem cell therapy for, 90e-2-3
 - type 1, **2403**
 - APS-1 and, 2346t
 - APS-2 and, 2346, 2347t
 - environmental factors in, 2404
 - genetic considerations in, 2403
 - HLA gene association with, 373e-8, 373e-8t
 - hypertension management, 2426
 - immunologic markers in, 2404
 - immunopathogenesis of, 372e-24t, 372e-27t, 372e-28, 2403, 2403f
 - inflammatory mediators in, 372e-29t
 - KIRs with, 372e-15t
 - pathophysiology of, 2403-2404
 - prevention of, 2404
 - treatment and management of, **2410**
 - adverse effects of, 2417
 - emerging therapies, 2417
 - goals of, 2408t, 2410-2411
 - insulin preparations, 2411, 2411t
 - insulin regimens, 2411-2412, 2412f
 - intensive management, 2411
 - pramlintide, 2412-2413
 - type 2, **2404**
 - bariatric surgery and, 2398
 - genetic considerations in, 440-441, 441t, 2404
 - hypertension management, 2426
 - insulin resistance syndromes in, 2405-2406
 - ketosis-prone, 2407
 - metabolic abnormalities in, 2404-2405, 2405f
 - metabolic syndrome and, 291e-6-7, 2450, 2452
 - obesity and, 415e-6
 - pathophysiology of, 2404, 2405f
 - postmenopausal hormone therapy and, 2384t
 - prevalence of, 2254t
 - prevention of, 2406
 - risk factors for, 2401, 2401t
 - in schizophrenia, 2722
 - screening/testing for, 2254t
 - treatment and management of, **2413**, 2413f, 2414t
 - adverse effects of, 2417
 - α -glucosidase inhibitors, 2415
 - biguanides, 2413
 - bile acid-binding resins, 2415
 - bromocriptine, 2415
 - combination therapy, 2416-2417
 - emerging therapies, 2417
 - insulin secretagogues, 2413-2415
 - insulin therapy, 2415-2416
 - selection of initial agents, 2416, 2416f
 - sodium-glucose co-transporter 2 inhibitors, 2415
 - thiazolidinediones, 2415
 - in women, 6e-2f, 6e-3-4
 - Diabetic diarrhea, 270, 272, 855t, 1942
 - Diabetic ketoacidosis (DKA), **2418**
 - clinical features of, 2418, 2418t
 - complications of, 2420
 - diagnosis of, 64e-1, 2418, 2418t
 - hypokalemia in, 305
 - hyponatremia in, 299
 - pathophysiology of, 318, 2418, 2418t

- Diabetic ketoacidosis (DKA) (*Cont.*):
prevention of, 2420
treatment of, 64e-1-2, 318, 2419-2420, 2419t
- Diabetic nephropathy, **2424**
clinical features of, 1835t
epidemiology of, 1844
glycemic control and, 2423
microalbuminuria in, 1844, 2425, 2425f
natural history of, 1844, 2425, 2425f
pathogenesis of, 1813, 1832, 1844, 2424-2425
renal biopsy in, 62e-8f, 1844
treatment of, 1845, 2425-2426
in type 1 vs. type 2 diabetes mellitus, 1844-1845, 2425
- Diabetic neuropathy, **2426**
autonomic, 2426, 2427, 2642, 2682
constipation in, 274
distal symmetric sensory and sensorimotor polyneuropathy, 2682
mononeuropathy, 2426, 2683
pathophysiology of, 89
polyneuropathy, 2426, 2683
polyradiculopathy, 2426
prevalence of, 2426
proximal, 462e-19
radiculoplexus neuropathy, 2682-2683
risk factors for, 2426
screening for, 2407
treatment of, 93, 93t, 2426-2427, 2682t
- Diabetic retinopathy
advanced, 206
clinical features of, 40e-2f, 40e-4f, 2424, 2424f
glycemic control and, 2423, 2423f
treatment of, 206, 2424
- "Diabetic skin spots," 2429
- Diacetyl exposure, 1692
- Diagnostic hypothesis, 19-20
- Diagnostic imperative, 19-20
- Diagnostic tests. *See* Clinical laboratory tests
- Dialectical behavior therapy (DBT), 2720
- Dialysate, 1822
- Dialysis. *See* Hemodialysis; Peritoneal dialysis
- Dialysis access, 1823
- Dialyzer, 1822, 1824
- Diamines, 205e-7
- 3,4-Diaminopyridine, 2672, 2703
- Diamond-Blackfan anemia, 668, 669t
- Diapedesis, 416, 417f
- Diaper rash, 1343
- Diaphyseal aclasia (multiple exostoses), 426e-7
- Diarrhea
acute, **265**. *See also* Diarrhea, infectious
approach to the patient, 267, 267f
etiology of, 265-267
treatment of, 267-268
after cholecystectomy, 2083
antibiotic-associated, 267, 991, 1936. *See also* *Clostridium difficile*-associated disease
in bacterial overgrowth syndrome, 1945
bile acid, 1934-1935, 1935t
bloody, 266, 266t, 267
Brainerd, 267
in carcinoid syndrome, 564
in celiac disease, 1942
chemotherapy-related, 103e-25, 103e-26
chronic, **268**
approach to the patient, 270-272, 271f, 271t
definition of, 265
endoscopic findings in, 1899, 1899f
etiology of, 268-270, 268t
in returning war veterans, 152e-8
treatment of, 272
in Crohn's disease, 1953
definition of, 265
diabetic, 270, 272, 855t, 1942
differential diagnosis of, 1877, 1877t, 1932-1933
with enteral feeding, 98e-9
epidemiology of, 264
fatty acid, 1935, 1935t
in HIV infection, 1257, 1257f
hypernatremia in, 303
hypokalemia in, 305
hyponatremia in, 298
hypovolemia in, 298
infectious, 266, **852**. *See also* Gastroenteritis;
Travelers' diarrhea
adenovirus, 266, 1208
approach to the patient, 853-854, 854f
B. fragilis, 1099
bacterial vs. viral, 1287t. *See also* Gastroenteritis
Campylobacter. *See* *Campylobacter* spp.
infections
clinical features of, 266, 266t, 1287t
complications of, 854, 855t
diagnosis of, 1031
E. coli. *See* *Escherichia coli* infections,
intestinal
epidemiology of, 266, 855-856
food-related, 266, 855-856, 856t. *See also* Food-related illness
Giardia, 1406
host defenses against, 853
in immunodeficient persons, 266
inflammatory, 852t
in institutionalized persons, 266
laboratory evaluation of, 856
noncholera *Vibrio*, 1065-1066, 1065t
noninflammatory, 852t
pathogenesis of, 265-266, 852-853, 852t
penetrating, 852t
prevention of, 857
protozoal, 1405-1409
Salmonella. *See* *Salmonella* spp. infections
Shigella. *See* *Shigella* spp. infections
with systemic manifestations, 266-267
T. whipplei, 1092
treatment of, 856-857, 857t, 1031
V. cholera. *See* Cholera
in irritable bowel syndrome, 1965
in malabsorption syndromes, 1932-1933, 1937, 1946t
metabolic acidosis in, 298, 320
osmotic, 303, 1933
overflow, 265
paraneoplastic, 609t
persistent, 265
postvagotomy, 1926
in protein-losing enteropathy, 1945
secretory, 303, 1933
in short bowel syndrome, 1943
traveler's. *See* Travelers' diarrhea
in tropical sprue, 1942
in ulcerative colitis, 1952, 1952t
in VIPoma, 570
water loss in, 95e-1
in Whipple's disease, 1945
in ZES, 568, 1927
- Diarrhetic shellfish poisoning, 2741-2742
- Diarylquinolines, 205e-7
- Diascopy, 342
- Diastolic blood pressure, 1616t
- Diastolic dysfunction, 265e-10, 265e-10f, 270e-7, 270e-8f
- Diastolic failure. *See* Heart failure (HF)
- Diastolic murmur. *See* Heart murmurs
- Diatrizoate meglumine, 727t
- Diazepam
for alcohol withdrawal, 2727
for anxiety disorders, 2711t
for cocaine toxicity, 469e-2
drug interactions of, 246e-3t, 1921
for insomnia, 65
for marine envenomations, 2739
for nerve agent exposure, 262e-5t, 262e-6-7, 262e-6f
overdose/poisoning with, 473e-10t, 2764t
pharmacology of, 2711t
for sedation after myocardial infarction, 1606
for spasticity, 2660, 2672
for stiff-person syndrome, 618
therapeutic monitoring of, 2764t
for vertigo, 151t
- Diazoxide, 380, 569, 2434
- Dibasic aminoaciduria (lysinuric protein intolerance), 332e-7t, 435e-1t, 435e-2
- Dibutyl phthalate, 1963
- DIC. *See* Disseminated intravascular coagulation (DIC)
- Dicarboxylic aminoaciduria, 435e-1t
- DICER gene mutations, 594
- Dichloralphenazone, 2592, 2592t
- Dichloroacetate (DCA), 102e-13
- Dichromat, 197
- Dichuchwa. *See* Endemic syphilis
- Dick test, 966
- Diclofenac
advantages of, 1923
adverse effects of, 389t, 727t
for gout, 2234
for osteoarthritis, 2232t
for pain, 91
- Dicloxacillin
adverse effects of, 284, 306
for auricular cellulitis, 228
for impetigo, 967
indications for, 937
for otitis externa, 228
for staphylococcal skin/soft tissue infections, 345, 768t, 962t
- Didanosine (ddI)
adverse effects of, 2023, 2030
drug interactions of, 1278, 2030
molecular structure of, 1276f
resistance to, 1279f
- Dideoxynucleosides, 1267
- DIDMOAD (Wolfram's) syndrome, 2276, 2348
- Dientamoeba fragilis*, 246e-9, 765t, 1409
- Diet
asthma and, 1670, 1671
breast cancer and, 524
for cancer prevention, 476
cardiovascular disease and, 266e-4
colorectal cancer and, 538
composition of, 95e-4
in elderly, 84-85
energy density of, 2395
global considerations in, 266e-4
infectious disease exposure and, 763
low-bacteria, 492
lung cancer and, 506
nephrolithiasis and, 1867, 1870
peptic ulcer disease and, 1917
- Dietary assessment. *See* Nutritional assessment
- Dietary fiber
cancer and, 476
colorectal cancer and, 538
for constipation, 273, 274
for irritable bowel syndrome, 1969
- Dietary reference intakes, 95e-1
- Dietary supplements, 14e-3-4, 14e-4t, 95e-3
- Dietary Supplements Health and Education Act (DSHEA), 14e-3
- Diet history, 95e-5
- Diethylcarbamazine (DEC)
action of, 246e-8
adverse effects of, 246e-6t, 1419
for loiasis, 1422
for lymphatic filariasis, 1419
pharmacology of, 246e-8
for tropical pulmonary eosinophilia, 1420
- Diethylene glycol poisoning, 64e-9
- Diethylpropion, 2396
- Diethylstilbestrol (DES), 103e-15, 477t, 585
- Diet therapy. *See* Nutritional support/therapy
- Dieulafoy's lesion, 277, 346e-1v, 1891, 1891f
- Differentiation, in cancer cells, 103e-6
- Diffuse alveolar hemorrhage, 245
- Diffuse axonal injury, 457e-2, 457e-2f
- Diffuse esophageal spasm, 256, 1905-1906, 1906f
- Diffuse fasciitis with eosinophilia, 376
- Diffuse idiopathic skeletal hyperostosis (DISH), 2172
- Diffuse infiltrative lymphocytosis syndrome (DILS), 1260-1261
- Diffuse large B-cell lymphoma, **705**
clinical features of, 702t, 705
diagnosis of, 705, 705f
genetic considerations in, 698-699, 699t, 705
in HIV infection, 1270
lymph node examination in, 81e-10f, 705f

- in pregnancy, 124e-4
 primary mediastinal, 705
 treatment of, 705-706
- Diffuse nontoxic goiter, 2301
- Diffuse white matter disease (leukoaraiosis), 174f, 2568, 2602
- Diffusion capacity of the lung for carbon monoxide (DL_{CO}), 306e-5f, 306e-6, 1710
- Diffusion tensor imaging (DTI), 440e-8, 440e-9f
- Diffunisal, 724
- DiGeorge (velocardiofacial) syndrome, 83e-6, 375e-1, 465e-2, 2113, 2483
- Digitalis
 action of, 265e-6
 adverse effects of, 44, 203, 273e-3t, 1471t, 1477, 2365
 ECG effects of, 1457
 for pulmonary edema, 1763
 for radiation-induced cardiovascular disease, 621
 for SA node dysfunction, 1469
- Digital necrosis, in systemic sclerosis, 2159, 2159f
- Digital rectal examination, 482-483, 482t, 540, 580
- Digit span tests, 168, 183
- Digoxin
 adverse effects of, 81, 727t, 1480t
 for arrhythmias, 1479t, 1480t
 for atrial fibrillation, 1487-1488
 distribution of, 34
 dosage of, 1479t, 1480t
 drug interactions of, 34t, 42, 42t
 azithromycin, 246e-4t
 azoles, 246e-3t
 PPIs, 1921
 pyronaridine, 246e-12
 for heart failure, 1512
 metabolism of, 34t, 86e-9
 overdose/poisoning with, 309-310, 473e-9t, 2764t
 pharmacology of, 1479t, 1480t
 for supraventricular arrhythmias, 1609
 therapeutic monitoring of, 2764t
 toxicity of, 42, 307, 312
- DIHS. *See* Drug-induced hypersensitivity syndrome (DIHS)
- Dihydroartemisinin, 246e-2t, 246e-7
- Dihydroergotamine
 for exertional headache, 2597
 for medication-overuse headache, 109
 for migraine, 2591, 2592t, 2593t
 nasal spray, 2591, 2592t
 parenteral, 2591, 2592, 2592t
- Dihydrofolate (DHF) reductase
 in antibiotic resistance, 931t, 934
 in DNA synthesis, 641, 642f
- Dihydrofolate (DHF) reductase inhibitors, 246e-4t, 246e-12
- Dihydropteridine reductase (DHPR) deficiency, 434e-2t
- Dihydropteroate synthetase, 931t, 932, 934
- Dihydropyrimidinase deficiency, 431e-5t
- Dihydropyrimidine dehydrogenase, 39t
- Dihydropyrimidine dehydrogenase deficiency, 431e-5t, 431e-6, 431e-6f
- Dihydrorhodamine oxidation test, 423
- Dihydrotestosterone (DHT), 332, 2358, 2359f, 2759t. *See also* Testosterone
- 1,25-Dihydroxyvitamin D. *See also* Vitamin D
 action of, 313, 313f, 2464-2465
 overproduction of, 313, 314, 609t, 610
 synthesis of, 387
- Diisocyanate exposure, 1692
- Dilated cardiomyopathy, 1556
 alcohol-related, 1562
 chemotherapy-related, 1562-1563
 chest imaging in, 47e-4
 in chronic kidney disease, 1817
 clinical features of, 1554t
 diagnosis of, 1554t, 1559-1560, 1559f
 echocardiography in, 1557f
 familial, 1555t, 1564-1565, 1564f
 genetic considerations in, 436t, 1498, 1554, 1555t
 heart failure in, 1501. *See also* Heart failure (HF)
 in hemochromatosis, 1564, 1564f
 in HIV infection, 1560
 idiopathic, 1565
 metabolic causes of, 1558t, 1563-1564
 myocarditis and. *See* Myocarditis
 natural history of, 1439
 pathology of, 1557f
 pathophysiology of, 372e-24t, 1556-1557, 1559f
 Takotsubo (stress-related), 97, 289e-3, 1565
 toxic, 1558t, 1562-1563
 ventricular tachycardia in, 1495
- DILIN (Drug-Induced Liver Injury Network), 2025
- Diloxanide furoate, 246e-3t, 246e-8
- DILS (diffuse infiltrative lymphocytosis syndrome), 1260-1261
- Diltiazem
 for accessory pathway-mediated tachycardia, 1483
 adverse effects of, 1589t, 2218t
 for aortic dissection, 1641
 for arrhythmias, 1479t, 1480t
 for atrial fibrillation, 1487
 for AV nodal reentrant tachycardia, 1479
 dosage of, 1479t, 1480t
 drug interactions of, 246e-6t, 1590
 for hypertension, 1624t, 1625
 for hypertension in chronic kidney disease, 1821
 for ischemic heart disease, 1589, 1589t
 for NSTEMI-ACS, 1596
 overdose/poisoning with, 473e-9t
 pharmacology of, 1479t, 1480t
 for Raynaud's phenomenon, 1649
 for sex headache prevention, 2597
 for supraventricular arrhythmia, 1609
- Dilute Russell's viper venom test, 406, 740
- Dimenhydrinate, 151t, 260, 260t
- Dimercaprol, 472e-2-3t
- D-Dimer test, 403, 407
 in DIC, 737
 in HIV infection, 1249t
 in VTE/PE, 1632
- 1,3-Dimethylamylamine (DMAA), 14e-3
- Dimethyl fumarate (DMF), 2669t, 2670, 2671
- Dimethyl sulfoxide, 60e-4
- Dimorphic fungi, 1329, 1353
- Dipeptidyl peptidase-4 (DPP-IV) inhibitors, 2414-2415, 2414t, 2416
- Diphenhydramine
 adverse effects of, 2640t
 for anaphylaxis, 2117
 for carcinoid syndrome, 566
 for insomnia, 191
 overdose/poisoning with, 473e-8t, 473e-13t
 as premedication for antivenom, 2737
 as premedication for taxanes, 103e-15
 for prevention of dystonic reactions, 103e-25
 for pruritus, 345
- Diphenoxylate, 272
- Diphenylhydantoin, 407, 569
- Diphosphoglycerate mutase (DGM) deficiency, 654t
- [^{99m}Tc]Diphosphonate scintigraphy, 2224. *See also* Radionuclide scans/scintigraphy
- Diphtheria, 977
 approach to the patient, 978
 clinical features of, 978-979, 978f, 1471t, 1561, 2633
 complications of, 979
 cutaneous, 977, 978, 978f, 979
 diagnosis of, 979
 differential diagnosis of, 978
 epidemiology of, 977
 etiology of, 977
 global considerations in, 977, 979
 incidence pre- and postvaccine, U.S., 785t, 977
 pathogenesis of, 977-978
 pathology of, 978
 prevention of, 787f, 980
 prognosis of, 980
 prophylaxis for close contacts, 980
 treatment of, 979-980
- Diphtheria antitoxin, 979
- Diphtheria-tetanus-pertussis vaccine (DTaP), 929, 929t, 980, 1024. *See also* Tetanus, diphtheria, acellular pertussis (Tdap) vaccine
- Diphtheria-tetanus vaccine. *See* Tetanus-diphtheria vaccine
- Diphtheria vaccine, for travelers, 794
- Diphyllbothrium latum* (fish tapeworm), 245e-2t, 645, 1434
- Diploid, 426, 427
- Diplopia, 208
 in abducens nerve palsy, 209
 in horizontal gaze disorders, 209-210
 monocular vs. binocular, 208
 in multiple ocular nerve palsies, 209
 in myasthenia gravis, 208. *See also* Myasthenia gravis
 in nystagmus. *See* Nystagmus
 in oculomotor nerve palsy, 208
 in opsoclonus, 211
 in supranuclear disorders of gaze, 209
 in trochlear nerve palsy, 209
- Dipsogenic polydipsia, 2276, 2279
- Dipteran bite, 2749-2750
- Dipylidium caninum*/dipylidiasis, 1435
- Dipyridamole, 748, 748f, 1889t, 2570
- DIRA (deficiency of the IL-1 receptor antagonist), 2215
- Direct alloreactivity, 373e-5
- Direct antiglobulin (direct Coombs) test, 138e-4, 138e-4f
- Directional atherectomy catheter, 296e-2
- Direct thrombin inhibitors, parenteral, 443e-1, 753, 753t. *See also* Argatroban; Bivalirudin; Lepirudin
- Dirofilaria* infections, 1423
- Disability
 assessment of, 1693
 frailty and, 78-79
 osteoarthritis-related, 2226
 prevalence, in older population, 93e-4-5, 93e-4f
- Disability-adjusted life years (DALY), 9, 10f, 11t
- DISC1 gene mutations. *See* DiGeorge (velocardiofacial) syndrome
- Discectomy with fusion, cervical, 123
- Discoid (chronic cutaneous) lupus erythematosus
 alopecia in, 355, 355t
 clinical features of, 375
 diagnosis of, 375
 oral manifestations of, 237
 skin manifestations of, 25e-3f, 76e-13f, 375, 376f, 2127-2128, 2129t
 telangiectasias in, 357
 treatment of, 375
- Discriminant function (DF) value, 2059
- Discrimination, 16e-4
- Disease
 classification of, 87e-3, 87e-4f
 definition of, 145e-1
 genotype-phenotype relationships in, 87e-3, 87e-4f
 germ theory of, 761
 networks, 87e-3-4, 87e-5f, 87e-6f, 87e-7f
 probability, measures of, 21-22
- Disease Control Priorities in Developing Countries*, 9, 10
- Disease-modifying antirheumatic drugs (DMARDs), 2145-2147, 2146-2147t
- Disequilibrium, 164. *See also* Balance
- DISH (diffuse idiopathic skeletal hyperostosis), 2172
- Disk/agar diffusion method, 150e-9
- Diskography, 440e-10
- Disk-space infection. *See* Osteomyelitis, vertebral
- Dislocation, 2219, 2219t

- Disopyramide
adverse effects of, 1480t
for arrhythmias, 1479, 1480t, 1489, 1498
dosage of, 1480t
pharmacology of, 1480t
therapeutic monitoring of, 2764t
- Disorders of sex development (DSDs), **2349**
classification of, 2350t
global considerations in, 2357
sex chromosome disorders. *See* Sex chromosome disorders
46,XX disorders. *See* 46,XX disorders of sex development
46,XY disorders. *See* 46,XY disorders of sex development
- Disposable soma theory of aging, 94e-2
- Disseminated intravascular coagulation (DIC), **736**
chronic, 738
clinical features of, 737
diagnosis of, 737
etiology of, 737, 737t
in filovirus infections, 1327
in leptospirosis, 1142
pathophysiology of, 737, 737f
in pregnancy, 49
in sepsis/septic shock, 1754, 1757
skin manifestations of, 368, 782
treatment of, 738
- Distal convoluted tubule, 332e-4f, 332e-7t, 332e-8
- Distal myopathies, 462e-12, 462e-13t
- Disulfiram
adverse effects of, 203, 2687t
in alcoholism management, 2728
drug interactions of, 246e-5t
- DITRA (deficiency of the IL-36 receptor antagonist), 2215
- Diuresis
for hypercalcemia, 2480t, 2481
postobstructive, 1874
water, 2275, 2276f
- Diuretics
abuse of, 307
action of, 297
adverse effects of
acid-base disorders, 322
gout, 2218t
hypokalemia, 305–306, 307
hyponatremia, 299
hypovolemia, 297
renal, 291
urinary incontinence, 81
xerostomia, 241
- aldosterone-renin ratio effects of, 2320t
for ascites, 288
for chronic venous disease, 1652
for heart failure, 1508, 1509f, 1510t, 1512–1513, 1608
for hyperkalemia, 312
for hypertension, 1623, 1624t
for Ménière's disease, 219
for pulmonary edema, 1763
for vertigo, 151t
- Divalent metal transporter, 626
- Divers Alert Network, 2741
- Diversion colitis, 1957
- Diversity, 86e-1t
- Diverticular disease, **1971**
anatomy of, 1971
bleeding in, 277, 1971
clinical features of, 1971–1972, 1972t
diagnosis of, 1891, 1893f, 1971–1972
epidemiology of, 1971
esophageal, 1903, 1903f
incidence of, 1971
jejunal, 1939f
pathophysiology of, 1971, 1971f
recurrent, 1973
treatment of, 346e-1v, 1893f, 1972–1973, 1973f, 1973t
Zenker's, 256, 259, 1903, 1903f
- Diverticulitis
clinical features of, 106t, 267, 1971–1972, 1972t
diagnosis of, 1971–1972
Hinchey classification of, 1972, 1972f
vs. IBD, 1957
- Diving medicine, **477e-6**
barotrauma, 477e-6–7
breathing equipment, 477e-6
decompression sickness, 477e-4t, 477e-7–8, 477e-7t
suitability evaluation, 477e-6
- Dix-Hallpike maneuver, 149
- Dizziness, **148**. *See also* Vertigo
approach to the patient, 148–149
diagnostic testing in, 149
etiology of, 148
physical examination in, 149
psychosomatic, 151
- DJ-1, 444e-9
- DKA. *See* Diabetic ketoacidosis (DKA)
- DKC1 gene mutations, 664
- DKK-1, 2143
- DL_{CO} (diffusion capacity of the lung for carbon monoxide), 306e-5f, 306e-6, 1710
- DLT (dose-limiting toxicity), 103e-5
- DM. *See* Diabetes mellitus (DM)
- DMAA (1,3-dimethylamylamine), 14e-3
- DMARDs (disease-modifying antirheumatic drugs), 2145–2147, 2146–2147t
- DMD gene mutations, 434, 436, 1555t
- DMF (dimethyl fumarate), 2669t, 2670, 2671
- DMSA (succimer), 472e-2–3t
- DNA
demethylation of, 102e-8
epigenetic modifications of, 429–431, 431f
human, 425
methylation of, 101e-4, 429–430, 431f, 432f
mitochondrial. *See* Mitochondrial DNA (mtDNA)
repetitive, 427
replication of, 427, 428f
structure of, 426–427, 427f
unstable sequences, 434
- DNA-dependent nucleoside-stimulated ATPase, 372e-24t
- DNA gyrase, 932
- DNA-interactive agents, 103e-8, 103e-9t, 103e-12. *See also* specific drugs
- DNA methyltransferase inhibitors, 103e-20
- DNA repair, 375e-1–2, 386, 434, 2109
- DNase aerosols, 1699
- DNA sequencing
in genetic testing, 427, 447. *See also* Genetic testing
multiplex, 86e-2, 86e-3f, 447
of stem cells, 455
- DNA topoisomerase II, 372e-25t
- DNA virus(es), 214e-2t, 214e-3f, 214e-4–5. *See also* specific viruses
- DNMT3A gene mutations, 680, 680t
- Dobrava-Belgrade virus, 1307t
- Dobutamine
for cardiogenic shock, 1761, 1761f
for heart failure, 1509–1510, 1510t
for pulmonary edema, 1761f, 1763
for sepsis/septic shock, 1757
for shock, 1751
in stress testing, 270e-3
- DOC (deoxycorticosterone), 2309f, 2313, 2319
- Docetaxel
action of, 103e-14–15
adverse effects of, 103e-12t, 103e-15, 1797, 2218t, 2686t
for breast cancer, 529
drug interactions of, 246e-3t
for gastric cancer, 535
for head and neck cancer, 505
for lung cancer, 520, 520t, 521
for melanoma, 499
premedication for, 103e-12t, 103e-15
for prostate cancer, 586
for soft tissue sarcoma, 119e-2
- DOCK8 deficiency, 420t, 423
- Docosanol, 215e-3t
- Docusate, 61, 62t
- Dofetilide
action of, 273e-6t
adverse effects of, 1480t, 1489, 1498
for arrhythmias, 1480t
for atrial fibrillation, 1489
for atrial flutter, 1485
dosage in kidney disease, 37
dosage monitoring, 36
dosage of, 1480t
drug interactions of, 43, 246e-3t
pharmacology of, 1480t
for ventricular arrhythmias, 1498
- Dog(s)
bite-wounds, **167e-1, 782**
antibiotic prophylaxis for, 167e-3
approach to the patient, 167e-2
cellulitis in, 830
infections from, 167e-1, 183e-3
microbiology of, 167e-1
rabies prophylaxis for, 167e-3
treatment of, 167e-2–3, 167e-3t, 832t
- filarial infections, 1423
- fleas, 2750
- tapeworms, 1435
- ticks, 1154
- Döhle bodies, 81e-2, 81e-7f, 415, 415f
- Dolasetron, 62, 103e-25
- "Doll's eyes," 1775
- Dolor, 414f, 1753
- Dolutegravir, 1275t, 1281
- Domains of life, 86e-1t
- Domestic abuse/violence
human bite-wounds in, 167e-2
incidence of, 6e-5, 2723
lifetime cumulative risk of, 27t
mandated reporting of, 17e-1
screening for, 29t, 2722–2723
- Domoic acid intoxication (amnesic shellfish poisoning), 2547–2548, 2743
- Domperidone
adverse effects of, 261–262
for gastroparesis, 260, 260t, 2164, 2641
for migraine, 2591
for nausea, 109
- Donath-Landsteiner antibodies, 138e-2, 659
- Donepezil
for AD, 176, 217, 2601
for cognitive problems in MS, 2672
for opioid-induced drowsiness, 61
for Parkinson's disease dementia, 2617
- Dong quai, 404t
- Do-not-resuscitate (DNR) orders, 67
- Donovan bodies, 198e-1, 198e-1f
- Donovanosis, **198e-1**
clinical features of, 198e-1, 198e-1f, 880t
diagnosis of, 198e-1–2, 198e-1f
differential diagnosis of, 198e-2
epidemiology of, 198e-1, 879
etiology of, 198e-1
global considerations in, 198e-1
prevention and control of, 198e-2
treatment of, 198e-2, 198e-2t
- L-Dopa. *See* Levodopa
- Dopamine
action of, 444e-3t
for cardiogenic shock, 1761, 1761f
in cardiovascular regulation, 1613
clinical aspects of, 444e-3t
for hypothermia, 478e-3
in migraine, 2588
for pulmonary edema, 1763
reference values for, 2759t
for shock, 1750, 1751
structure of, 444e-3t
- Dopamine agonists
for acromegaly, 2271
adverse effects of, 192, 2268, 2615
for ovulatory dysfunction, 2389
for PD, 2615, 2616t
for prolactinoma, 2267–2268
for restless legs syndrome, 192

- Dopamine receptor antagonists
adverse effects of, 565
for migraine, 2588, 2591–2592, 2592t
for nausea and vomiting, 260, 260t
parenteral, 2592, 2592t
- Dopa-responsive dystonia, 434e-2t, 2620
- Doppler ultrasound, 270e-2, 270e-2f. *See also* Echocardiography
- Droripenem
for anaerobic bacterial infections, 1101
indications for, 941
for intraabdominal infections, 767t
for *P. aeruginosa* infections, 1045t
resistance to, 933
- Dormancy, in cancer cells, 103e-6
- Dornase, 1696
- Dorsal midbrain syndrome (Parinaud's syndrome). *See* Parinaud's syndrome/complex
- Dorsal respiratory group, 1720
- Dorsal root ganglia, 87, 87f, 88
- Dorsal root ganglionopathy, 618
- Dose-limiting toxicity (DLT), 103e-5
- Dosulepin, 2594t
- DOTS/DOTS-Plus strategy, for tuberculosis, 12, 1121
- Double-balloon endoscopy/enteroscopy
indications for, 1878, 1879t
small-bowel, 346e-1v, 1881, 1883f, 1897f
- Doublecortin, 2545t
- Double-dip sign, 226
- Double minute, 101e-3
- Downbeat nystagmus, 211
- Down's syndrome (trisomy 21)
AD and, 2600–2601
AML in, 678
autoimmune polyendocrine deficiencies in, 2348
chromosome abnormalities in, 83e-1, 83e-4
clinical features of, 83e-4, 1654
myelodysplasia in, 670
oral manifestations of, 236
transient myeloproliferative disorder in, 135e-7
- Doxazosin, 1624t, 1649
- Doxepin
adverse effects of, 2710t
for angioedema, 2119
for insomnia, 191
overdosage/poisoning with, 473e-9t, 473e-10t, 2764t
for pain, 91t
for pruritus, 345
therapeutic monitoring of, 2764t
for urticaria, 2119
- Doxorubicin
action of, 103e-13
adverse effects of, 103e-10t, 103e-13
acral erythema, 380
cardiotoxicity, 103e-13, 1563, 1567
late, 620t, 621
myelosuppression, 103e-13
neurologic, 1793
for bladder cancer, 577
for breast cancer, 528, 529
drug interactions of, 103e-10t
for Ewing's sarcoma, 119e-4
for gastric lymphoma, 536
for hepatocellular carcinoma, 550
for insulinoma, 2338
for lung cancer, 522
for osteosarcoma, 119e-3
for ovarian cancer, 594
for pNETS, 574
in pregnancy, 124e-4
for salivary gland tumors, 506
for soft tissue sarcoma, 119e-2
- Doxycycline
for acne rosacea, 352
for acne vulgaris, 352
for actinomycosis, 1091t
action of, 930, 931t, 943
adverse effects of, 361, 943, 1380t
for anthrax, 261e-3, 261e-5t
for bacillary angiomatosis/peliosis, 832t, 1081t
for *Bartonella* endocarditis, 1081t
for *Bartonella* infection prophylaxis in HIV infection, 1252t
for bite-wound infections, 167e-3, 167e-3t, 832t
for *Brucella* infections, 194e-4
for *C. trachomatis* infections, 1171, 1172
for cat-scratch disease, 1081t
for cholera, 1064
for community-acquired pneumonia, 808t
for donovanosis, 198e-2t
for ehrlichioses, 1160
for endemic treponematoses, 207e-2
for epidemic typhus, 1159
for gonococcal infections, 1009t
for human granulocytotropic anaplasmosis, 1161
indications for, 940t, 943
for infective endocarditis, 823t
for *Legionella* infections, 1020t
for leptospirosis, 1145t
for Lyme disease, 836, 1153, 1153f
for Lyme disease prophylaxis, 2745
for lymphatic filariasis, 1419
for *M. pneumoniae* infections, 1164
for malaria, 1378t, 1380t
for malaria prophylaxis, 796t, 1383, 1383t
for MRSA infections, 345, 962t
for mucopurulent cervicitis, 876
for onchocerciasis, 1421
for pelvic inflammatory disease, 878, 878t, 879
for pericardial effusion, 474
for plague prophylaxis, 1075t
for plague treatment, 261e-5, 261e-5t, 261e-6, 1075
for pleural effusion, 474
prophylactic, 945t
for Q fever, 1162
for reactive arthritis, 2175
for relapsing fever, 1148, 1148f
resistance to, 931t, 940t
for rickettsialpox, 1158
for Rocky Mountain spotted fever, 780t, 782, 1157
for scrub typhus, 1159
for sinusitis, 227t
for staphylococcal skin/soft tissue infections, 962t
for syphilis, 1139, 1139t
for tularemia, 261e-5t, 261e-7
for Whipple's disease, 1094
- Doxylamine, 261, 473e-8t
- DPP-IV (dipeptidyl peptidase-4) inhibitors, 2414–2415, 2414t, 2416
- DQB genes, 373e-7–8
- Dracunculus medinensis* (guinea worm)/dracunculiasis
clinical features of, 829, 837, 1422
diagnosis of, 245e-3t, 1422
epidemiology of, 1422
life cycle of, 245e-3t, 1422
prevention of, 1423
treatment of, 246e-13, 1423
- Dravet's syndrome, 2545t
- Drawer sign, 2222
- DRB genes, 373e-2, 373e-7
- DRESS (drug reaction with eosinophilia and systemic symptoms). *See* Drug-induced hypersensitivity syndrome (DIHS)
- Dressing apraxia, 182
- Dressings, for chronic venous disease, 1652
- Driving ability/privileges, 188, 2559
- Dronabinol, 81, 469e-3
- Dronedarone
adverse effects of, 1480t, 1489
for arrhythmias, 1480t, 1489, 1499, 1514
dosage of, 1480t
drug interactions of, 205e-3
pharmacology of, 1480t
- Drop attacks, 165. *See also* Fall(s)
- Dropped head syndrome, 462e-1
- Drosiprenone, 310, 334
- Drowsiness, 61, 1771
- Droxidopa, 2644
- Drp1, 444e-9
- Drug(s)
adverse reactions to. *See* Adverse drug reactions; Drug-induced illness
benefits of, 32
bioavailability of, 33
clearance of, 35
development of
for cancer, 103e-5–6, 103e-6f
genomics in, 775–776
systems biology and, 87e-2–3
targeted agents, 103e-6
target molecules in, 32
distribution of, 34–35
dose-response relations, 32, 32f
dose selection, 36
elimination of, 35
failure of efficacy of, 36
genetic determinants of response to, 38
half-life of, 34, 35f
indications for, 32
interactions between. *See* Drug interactions
intravenous administration of, 35
medication errors, 3
metabolism of
active metabolites in, 35
enzymes in, 935
first-pass effect, 33–34, 33f, 935
genetic variations in, 38–41, 38f, 39t
gut microbiota and, 86e-7
new, skews in research for, 13e-4
overdosage of. *See* Poisoning/drug overdose
plasma concentration of, 33–34, 33f, 36–37, 37f
prescription, use in U.S., 377
risks of, 32
slow-release formulation of, 33
sustained-release formulation of, 33
therapeutic monitoring of, 2764–2766t
time-of-day effects on, 194
- Drug-dependent antibodies, 727
- Drug disposition. *See* Drug(s), metabolism of
- Drug dosage
dosing interval, 34
in elderly, 38
in heart failure and shock, 37
in kidney disease, 37
in liver disease, 37
loading dose, 35, 35f
maintenance dose, 36
plasma concentration as guide to therapy, 35
selection of, 36–37
- Drug fever, 370
- Drug history, 44
- Drug-induced hypersensitivity syndrome (DIHS)
clinical features of, 131t, 134, 138, 370, 382
diagnosis of, 382, 384t
drugs associated with, 382
etiology of, 131t, 353, 382
genetic considerations in, 378
immune pathways of, 378, 378t
skin manifestations of, 25e-12f, 131t
time of onset, 382
- Drug-induced illness. *See also* Adverse drug reactions; Poisoning/drug overdose
anaphylaxis, 381
autonomic dysfunction, 2639, 2640t
cutaneous, 103e-26, 127, 377
cross-sensitivity in, 384–385
delayed hypersensitivity, 377–378
desensitization for prevention of, 384
diagnosis of, 383, 383t
drug selection after, 383–384
exacerbation or induction of dermatologic disease, 378–379
exanthematous eruption, 25e-2f, 128t
exanthems, 362, 362t
fixed drug eruption, 381, 382f

- Drug-induced illness, cutaneous (*Cont.*):
 genetic considerations in, 378
 global considerations in, 378
 hair disorders, 379–380
 hyperpigmentation, 360t, 361
 hypersensitivity reactions, 384t, 909t
 hypersensitivity syndrome. *See* Drug-induced hypersensitivity syndrome (DIHS)
 immediate, 377
 immune complex–dependent, 377
 incidence of, 377
 irritant/contact dermatitis, 381
 maculopapular eruptions, 380–381
 management of, 383–384
 nail disorders, 380
 overlap hypersensitivity syndromes, 383
 pathogenesis of, 377–378
 photosensitivity eruptions, 379, 379f, 389–390, 389t
 pigmentation changes, 379
 pruritus, 381
 pustular eruptions, 382. *See also* Acute generalized eruptive/exanthematous pustulosis (AGEP)
 reporting of, 385
 severe. *See* Stevens–Johnson syndrome; Toxic epidermal necrolysis
 “treating through” an eruption, 384
 urticaria/angioedema, 381
 vasculitis, 382, 2193
 vesicles/bullae, 361
 fatigue, 152
 fever of unknown origin, 138, 139
 liver injury. *See* Liver disease/failure, drug-induced
 lupus-like disease, 2134
 movement disorders, **2624**
 musculoskeletal conditions, 2218t
 primary immunodeficiencies, 375e–2
 vasculitis, 2218t
- Drug-Induced Liver Injury Network (DILIN), 2025
- Drug interactions, **36, 41**
 in elderly, 77–78
 with herbal/dietary supplements, 14e-4, 14e-4t
 in HIV infection, 798
 not mediated by changes in drug disposition, 43
 pharmacokinetic, 41–43, 42t
- Drug reaction with eosinophilia and systemic symptoms (DRESS). *See* Drug-induced hypersensitivity syndrome (DIHS)
- Drug-receptor interaction, 36
- Drug-target interactions, 40–41
- Drug-target network analysis, 87e-3
- Drusen
 vs. hard exudates, 40e-3
 in macular degeneration, 205, 205f
 optic disc, 203–204, 203f
- Dry beriberi, 96e-3, 2690
- Dry eye. *See* Keratoconjunctivitis sicca
- Dry mouth. *See* Xerostomia
- “Dry tap,” in bone marrow aspiration, 135e-1, 135e-2t
- DSDs. *See* Disorders of sex development (DSDs)
- DSGI/2* gene mutations, 2483
- DSHEA (Dietary Supplements Health and Education Act), 14e-3
- DTaP. *See* Diphtheria-tetanus-pertussis vaccine (DTaP)
- DTIC. *See* Dacarbazine (DTIC)
- DTPA, for internal radionuclide contamination, 263e-6t, 263e-7
- Dual-energy x-ray absorptiometry (DEXA), 2067, 2472, 2493
- Dual-process theory, 18
- Dual-task walking, 76, 162
- Dubin–Johnson syndrome, 282, 2003–2004, 2003t
- Duchenne’s muscular dystrophy, **462e-5**
 clinical features of, 462e-5–6, 462e-5t
 diagnosis of, 462e-6
 early interventions for, 450t
 genetic considerations in, 436, 462e-5t, 462e-6
 genetic testing for, 450t
 incidence of, 462e-5
 pathogenesis of, 462e-6, 462e-8, 462e-8f
 treatment of, 462e-8
- Duck fever, 1681t
- Ductal carcinoma in situ, 531
- Duct of Luschka, bile leak from, 1885f
- Duffy-binding protein, 145e-3
- Duffy blood group system, 138e-1t, 138e-2
- Dugbe virus, 1308t
- Duke criteria, for infective endocarditis, 819, 819t
- Dulaglutide, 2414t
- Duloxetine
 adverse effects of, 2682t, 2710t
 for depression, 2710t
 for fibromyalgia, 2240, 2240t
 for neuropathic pain, 94–95, 2427, 2682t
 for osteoarthritis, 2232
 for pain, 91t, 94
- Dumping syndrome, 1926
- Duodenal neuroendocrine tumors, 563. *See also* Gastrointestinal neuroendocrine tumors (GI-NETs)
- Duodenal ulcers
 clinical features of, 1918
 complications of, 1918
 diagnosis of, 1881, 1881f, 1889f, 1918–1919, 1919f
 differential diagnosis of, 1918
 epidemiology of, 1914
 gastroduodenal mucosal defense and, 1912–1913, 1912–1914
H. pylori and. *See Helicobacter pylori* infection
 NSAID-induced, 1923–1924, 1923t
 pathology of, 1914
 pathophysiology of, 1915
 recurrence of, 1926
 treatment of, 346e-1v, 1890, 1924–1927, 1925f.
See also Peptic ulcer disease, treatment of
- Duodenitis, 277
- DUOX, 2285
- DUOXA, 2285
- Duplication, chromosome, 83e-1, 83e-6
- Duplication cyst, 277
- Dura, vascular malformations of, 2586, 2657, 2658f
- Durable power of attorney for health care, 59
- Dural sinus thrombosis, 2572
- Dura matter grafts, Creutzfeldt–Jakob disease after, 453e-4
- Duroziez’s sign, 411
- Dutasteride, 7e-4, 478, 580
- Duvenhage virus, 1304
- DUX4* gene, 462e-2f
- DVT. *See* Deep vein thrombosis (DVT)
- Dwarf tapeworm (*Hymenolepis nana*), 245e-2t, 246e-11, 765t, 1434
- Dynamic visual acuity, 149
- Dysbetalipoproteinemia, familial, 2440t, 2443
- Dysbiosis, 86e-1t
- Dyschondroplasia (Ollier’s disease), 426e-7, 594
- Dyschromic macules, 207e-3
- Dysembryoplastic neuroepithelial tumors, 603
- Dysentery
 in amebiasis. *See* Amebiasis
 etiology of, 266, 853
 in *Shigella* infections. *See Shigella* spp. infections
 treatment of, 268, 857
V. parahaemolyticus, 1065
- Dysesthesia, 158. *See also* Sensation, abnormalities of
- Dysferlin, 462e-8, 462e-8f
- Dysfibrinogenemia, 736, 739
- Dysfunctional uterine bleeding, 335
- Dysgerminoma
 brain, 2264
 ovary, 594–595
- Dysgeusia. *See* Taste disorders
- Dysgraphia, 179
- Dyshematopoietic syndrome, 729
- Dyshidrotic eczema. *See* Eczema, dyshidrotic
- Dyskeratosis congenita (Hoyeraal–Hreidarsson syndrome)
 clinical features of, 102e-3, 359–360, 360t, 664
 genetic considerations in, 375e-1, 664, 2109
 pathogenesis of, 102e-3
- Dyskerin, 664
- Dyskinesia(s)
 diphasic, levodopa-related, 2614
 exercise-induced, 2624
 paroxysmal, 2624–2625
 pathophysiology of, 2613f
 tardive, 163, 261, 2624, 2722
- Dyslipedemias. *See* Lipoprotein disorders
- Dysmenorrhea, 338–339
- Dyspareunia, 330
- Dyspepsia. *See also* Indigestion
 clinical features of, 258, 1894
 endoscopy in, 1894
 functional (essential), 261, 1040, 1918
 new-onset, approach to, 1924, 1924f
 nonulcer, 1894, 1895
- Dysphagia, **254**
 approach to the patient, 256–257, 256f
 in cranial nerve disorders, 2649
 diagnosis of, 257, 1895–1896, 1896f, 1897f
 esophageal, 256, 1901
 in esophageal cancer, 262, 533
 in gastric cancer, 534
 in GERD, 1907
 history in, 256–257
 involuntary weight loss and, 275
 in myasthenia gravis, 2701
 nutritional support in, 98e-3
 oropharyngeal, 256
 pathophysiology of, 255–256
 physical examination in, 257
 propulsive (motor), 255–256
 radiation therapy-related, 505
 with solids vs. liquids, 256
 structural, 255
 in terminally ill patient, 69t
 transfer, 1895
 treatment of, 258
- Dysphonia, 2649
- Dysplasia(s)
 of bone. *See* Bone disease
 fibromuscular, 1617, 1627, 1645, 2567
 pituitary, 2255–2256
 skeletal. *See* Chondrodysplasia(s)
- Dysplastic nevus, 76e-7f, 493–494, 493f
- Dyspnea, **47e-1**
 anxiety and, 47e-1
 in aortic stenosis, 1530
 approach to the patient, **47e-3, 47e-4f**
 assessment of, 47e-1–2, 62, 1661
 in cardiovascular disease, 47e-1t, 47e-2, 47e-4, 1439, 1503–1504
 in COPD, 1704
 in cor pulmonale, 1506
 definition of, 47e-1
 differential diagnosis of, 47e-2, 47e-4
 efferent-afferent mismatch in, 47e-1f
 exertional, 47e-4
 history in, 47e-3
 imaging in, 47e-4
 intensity and quality of, 47e-1–2, 47e-1t
 laboratory studies in, 47e-4
 nocturnal, 47e-3
 in orthostatic hypotension, 146
 pathophysiology of, 47e-1, 47e-1f
 physical examination in, 47e-3–4
 in pulmonary hypertension, 1655
 in respiratory acidosis, 322
 in respiratory disease, 47e-2, 47e-2t, 47e-4, 1661
 in terminally ill patient, 62, 62t
 treatment of, 47e-4, 62, 62t
- Dysreflexia, autonomic, 2641–2642
- Dysthymic disorder, 2715
- Dystonia, **2620**
 clinical features of, 163, 2619t, 2620
 drug-induced, 65, 2624
 focal, 2620

- pathophysiology of, 2620–2621
 primary, 2620
 secondary, 2620
 tardive, 2624
 treatment of, 2621
 in Wilson's disease, 2519
- Dystonia plus syndromes, 2620
 Dystonic storm, 2621
 Dystroglycans, 462e-8, 462e-8f, 462e-9f
 Dystrophia myotonica. *See* Myotonic dystrophy
 Dystrophic calcification, 363, 426e-7t, 426e-8
 Dystrophin, 436, 462e-6, 462e-8, 462e-8f, 462e-9f, 1556f
- Dysuria, **60e-1**
DYT gene mutations, 2620
- E**
 EAEC (enteroaggregative *E. coli*), 1028t, 1031
 EAP (extracellular adherence protein), 956
 Ear. *See also* Hearing loss
 anatomy and physiology of, 217, 218f
 chondritis of, 2203, 2204f
 infections of
 auricular cellulitis, 228
 vs. auricular chondritis, 2204
 otitis externa, 228–229, 1045t, 1046–1047, 1348
 otitis media. *See* Otitis media
P. aeruginosa, 1046–1047
 perichondritis, 228
- Early after depolarizations (EADs), 273e-3–4, 273e-3f
 Early repolarization syndrome, 1498
 Eastern Cooperative Oncology Group (ECOG) fatigue assessment, 63
 performance scale, 103e-7, 472, 472t
 Eastern equine encephalitis virus, 1306t, 1316
- Eating disorders
 anorexia nervosa, 259, **2719**
 avoidant/restrictive food intake disorder, 2719
 binge-eating disorder, **2720**
 bulimia nervosa. *See* Bulimia nervosa
 constipation in, 273–274
 diarrhea in, 270
 hypokalemia in, 307
 sleep-related, 192
- Eaton-Lambert syndrome. *See* Lambert-Eaton myasthenic syndrome (LEMS)
- Ebola virus, 261e-8, 1324, 1324f
 Ebola virus disease
 clinical features of, 783, 1327–1328, 1327t
 complications of, 1328
 control and prevention of, 1328–1329
 diagnosis of, 1328
 epidemiology of, 782–783, 1324, 1325f, 1326f
 pathogenesis of, 1326–1327
 prognosis of, 1328
 treatment of, 1328
- Ebstein's anomaly, 1527
 Eburnation, 1443
 EBV. *See* Epstein-Barr virus (EBV)
 EBV infections. *See* Epstein-Barr virus (EBV) infections
- E-cadherin, in gastric cancer, 534
 Ecallantide, 2120
 Ecamsule, 391, 391t
 Ecchymoses, 127, 403, 721, 721f, 1442
 ECF-A (eosinophil chemotactic factor of anaphylaxis), 372e-15t, 422
- ECG. *See* Electrocardiography (ECG)
 Echinocandins, 1331, 1344t, 1352t, 1353, 1910
 Echinococcosis (hydatid disease)
 clinical features of, 152e-4t, 1432–1433, 1561
 diagnosis of, 245e-2t, 245e-6t, 1433
 eosinophilia in, 245e-6t
 epidemiology of, 152e-4t, 1432
 etiology of, 245e-2t, 1432
 incubation period for, 152e-4t
 joint involvement in, 837
 prevention of, 152e-4t, 1434
 in returning war veterans, 152e-4t
 staging of, 1433, 1433f
 treatment of, 152e-4t, 1433–1434
- Echinococcus granulosus*, 245e-2t, 1432. *See also* Echinococcosis (hydatid disease)
Echinococcus multilocularis, 245e-2t, 1432. *See also* Echinococcosis (hydatid disease)
- Echinocytes, 81e-1
 Echocardiography, **270e-1**
 after myocardial infarction, 270e-19–20, 270e-20f
 in aortic dissection, 1641
 in atrial septal defect, 270e-25, 270e-25f, 297e-7f, 297e-8f, 1521, 1522f
 in cardiac amyloidosis, 271e-6f, 271e-6v
 in cardiac tumors, 270e-23–24, 270e-24f, 289e-1, 289e-1f
 in chest pain evaluation, 102, 271e-1f, 271e-1v
 contrast agents in, 270e-9
 in cor pulmonale, 1506
 Doppler, 270e-1–2, 270e-2f
 hand-held equipment for, 270e-2, 270e-3f
 in heart murmurs, 51e-8, 1440
 in hypertrophic cardiomyopathy, 271e-6f, 271e-6v
 in infective endocarditis, 270e-24, 270e-24f, 820, 820f, 821f
 in multiple/mixed valvular heart disease, 1552
 in myocardial function assessment, 265e-9
 in myocardial infarction, 271e-1f, 271e-1v, 1601
 in pericardial effusion, 270e-22–23, 270e-22f, 271e-8f, 271e-8v, 1572f
 principles of, 270e-1–2, 270e-1f
 in pulmonary embolism evaluation, 1634
 in pulmonary hypertension, 271e-9f, 271e-9v, 1655–1656, 1657f
 in pulsus paradoxus, 271e-8f, 271e-8v
 in rheumatic heart disease, 2151f, 2151t, 2154v
- stress
 advantages of, 270e-10
 in CAD evaluation, 270e-10, 270e-26v, 271e-1f, 271e-1v
 indications for, 270e-2
 in ischemic heart disease evaluation, 1585
 for risk stratification, 270e-14, 270e-14f
 three-dimensional, 270e-1, 270e-2f
 transesophageal. *See* Transesophageal echocardiography
- two-dimensional, 270e-1, 270e-1f
 in valvular heart disease, 270e-15–16
 aortic regurgitation, 270e-17–18, 270e-17f, 1536
 aortic stenosis, 270e-16–17, 270e-17f, 271e-5f, 271e-5v, 297e-6f, 1531
 mitral regurgitation, 270e-18–19, 270e-18f, 270e-19f, 1545, 2154v
 mitral stenosis, 270e-19, 270e-19f, 1540, 2154v
 mitral valve prolapse, 270e-16, 1546–1547
 tricuspid regurgitation, 1548
 tricuspid stenosis, 1548
 in ventricular septal defect, 270e-25
- Echolalia, 177t, 178, 2623
 Echo-planar magnetic resonance imaging, 440e-8
- Echoviruses, 1289
 Echovirus infections
 chronic meningitis, 908t
 rash in, 128t, 135, 362, 362t, 1292
- Eclampsia, 49, 1626t
 ECMO. *See* Extracorporeal membrane oxygenation (ECMO)
- ECOG. *See* Eastern Cooperative Oncology Group
- Econazole, 233
 Economic issues
 health and, 9
 in medical care, **15e-1**, 21
 clinical decision-making and, 15e-2–3, 21
 health insurance and use of services, 15e-2–3
 pricing structures, 15e-3
 selection, 15e-1–2
 in population aging, 93e-4
- Ecstasy (MDMA), 168, 301, 469e-3
- Ectasias, vascular. *See* Vascular ectasias
 Ecthyma, 349
 Ecthyma gangrenosum
 in burn patient, 166e-2
 in cancer patient, 486
 clinical features of, 76e-12f, 132t, 135, 1044
 epidemiology of, 132t, 135
 etiology of, 132t, 183e-2, 782, 830, 1044
 in sepsis/septic shock, 1754
 skin manifestations of, 25e-9f, 132t, 369, 782
- Ectopia lentis, 436e-6f
 Ectopic ossification, 426e-7t, 426e-8
 Ectopic pregnancy, 106t, 338
 Ectopy, cervical, 876
 Ectropion, 199, 1977
 Eculizumab
 for familial hemolytic-uremic syndrome, 657, 1864
 for kidney transplant rejection, 1829
 for membranoproliferative glomerulonephritis, 1841
 for paroxysmal nocturnal hemoglobinuria, 661–662, 661f
- Eczema, **344**. *See also* Atopic dermatitis
 asteatotic, 342f, 346
 clinical features of, 344
 dyshidrotic, 76e-2f, 341t, 342f, 346, 346f
 hand, 342f, 346, 346f
 nummular, 346
- Eczema herpeticum, 1178
 ED. *See* Erectile dysfunction (ED)
 EDD (extended daily dialysis), 1811
- Edema, **250**
 approach to the patient, 253
 brain, 1777
 in cardiovascular disease, 1439
 in cirrhosis, 252, 252t
 clinical features of, 251
 cytotoxic, 1777
 distribution of, 253
 drug-induced, 252, 253t
 etiology of, 251–253, 252t, 253t
 in heart failure, 251–252, 252t, 1504
 heat-related, 479e-1
 hypoalbuminemia and, 294
 inflammatory, 250
 in kidney disease, 252, 252t
 localized, 253
 localized hypoxia and, 248
 lower extremity, 1443
 in nephrotic syndrome, 252, 252t
 of nutritional origin, 252–253
 pathogenesis of, 250–251, 251f, 332e-10f
 pitting, 251, 1443
 in pneumonia, 804
 pulmonary. *See* Pulmonary edema
 refeeding, 253
 stasis dermatitis and, 346
 vasogenic, 1777
- Edentulousness, 242
 Edinger-Westphal nuclei, 195
 Edoxaban, 756–757, 756t, 1635, 2570
- Edrophonium
 for anticholinesterase testing
 in botulism, 989
 false-positive results from, 2702
 in myasthenia gravis, 208, 2702, 2702t
 in snakebite, 2737t
 for snakebite treatment, 2737, 2737t
- EDSS (Expanded Disability Status Score), 2668t
 EDTA (ethylenediaminetetraacetic acid), 2237
Edwardsiella tarda infections, 1025, 1035–1036
 EEG. *See* Electroencephalography (EEG)
- Efalizumab, 372e-31
 Efavirenz
 adverse effects of, 1274t, 1278
 contraindications to, 1273
 for HIV infection, 1274t, 1278
 metabolism of, 34t
 molecular structure of, 1276f
 resistance to, 773, 1278, 1279f
- Eflornithine
 action of, 246e-8
 adverse effects of, 246e-3t

- Eflornithine (*Cont.*):
 cream, for hirsutism, 334
 pharmacology of, 246e-8-9
 in pregnancy and lactation, 246e-3t
 for sleeping sickness, 1398, 1398t
 EGF (epidermal growth factor), 400e-1t, 1912, 1913f, 2285
 EGFR (epidermal growth factor receptor), 39t, 41, 103e-16, 507, 1932
 EGFR (epidermal growth factor receptor) antibodies, 103e-21, 103e-21t
 EGFR gene mutations
 in autism spectrum disorders, 599
 in head and neck cancer, 503
 in lung cancer, 507t, 508, 509, 509f, 521
 EGFR inhibitors. *See* Epidermal growth factor receptor (EGFR) inhibitors
Eggerthella lenta, 86e-9
 Egophony, 1662
 EGPA. *See* Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
 Egyptian roussettes, 1324
 EHEC. *See* Enterohemorrhagic *E. coli* (EHEC)
 Ehlers-Danlos syndrome, 2510
 classification of, 2510, 2510t
 clinical features of
 bleeding, 403
 cardiac, 290e-1t, 1638, 1641, 2511
 cutaneous, 367, 2510-2511
 hernias, 2511
 ligament and joint changes, 2511
 pes planus, 2511
 scoliosis, 2511
 vascular disorders, 732
 diagnosis of, 2511
 genetic considerations in, 732, 2510t, 2511
 incidence of, 2510
 in pregnancy, 47
 treatment of, 2511
Ehrlichia spp., 150e-8, 1152, 1159
Ehrlichia chaffeensis, 129t, 888, 1159-1160.
 See also Human monocytotropic ehrlichiosis
Ehrlichia ewingii, 1160
 Ehrlichiosis, 1155t, 1159-1160, 1159f
 ELA (enzyme immunoassay), 150e-2, 1245
 EIEC (enteroinvasive *E. coli*), 1028t, 1031.
 See also *Escherichia coli* infections, intestinal
Eikenella corrodens, 167e-1, 167e-2, 183e-1, 183e-2t. *See also* Infective endocarditis, HACEK group
 Eisenmenger's syndrome
 clinical features of, 400, 1522, 1523
 heart-lung transplantation for, 320e-2
 hypoxia in, 248
 in pregnancy, 47, 1522
 treatment of, 1523
 Ejaculation, 325, 327
 Ejection fraction, 265e-9
 Ekbohm's syndrome, 2750-2751
 Ekiri syndrome, 1056
 ELANE gene, 2104
 Elapids, 2734, 2737. *See also* Snakebites
 Elastase: antielastase hypothesis, emphysema, 1700-1701
 Elastase, stool, 2087t, 2090
 Elastin, 1700, 2507, 2513
 Elbow
 bursitis of, 2247
 epicondylitis of, 2249
 neuropathy at. *See* Ulnar neuropathy
 ELC/MIP-3 β , 372e-12t
 Elderly, 70. *See also* Aging
 appendicitis in, 1987
 asthma in, 1680
 benign forgetfulness of, 171
 care coordination for, 83-84, 83t
 cerumen impaction in, 220
 clinical decision-making for, 82
 comorbidity of chronic diseases in, 71, 72t, 77-78, 82
 delirium in, 79, 79f, 166. *See also* Delirium
 diabetes mellitus in, 82. *See also* Diabetes mellitus
 dietary guidelines for, 84-85
 disability in, 71, 72t, 93e-4-5, 93e-4f
 drug-induced illness in, 38
 drug use in, 38
 elder abuse, 17e-1
 exercise in, 84
 falls in, 76, 79-80, 80f, 162, 164. *See also* Fall(s)
 frailty in. *See* Frailty
 functional assessment of, 78-79, 78t
 health care settings for, 82-83, 83t
 hearing loss in, 219, 219f
 heat-related illness in, 479e-1
 hemoglobin/hematocrit in, 394t
 hypernatremia in, 303
 hypothermia in, 478e-1, 478e-3
 involuntary weight loss in, 81, 274-275
 levothyroxine dosage in, 2293
 liver abscess in, 850
 musculoskeletal disease in, 2219, 2228
 olfactory dysfunction in, 214, 214f, 217
 oral health, 242
 pain management in, 80
 PD in, 82. *See also* Parkinson's disease
 polypharmacy in, 77-78
 preventive care for, 84
 principles of care for, 4
 rheumatoid arthritis in, 2148-2149
 screening tests in, 31t, 84
 seizures in, 2546t, 2547
 undernutrition and anorexia in, 81
 urinary incontinence in, 80-81, 81f
 vaccines for, 84
 vitamin D deficiency in, 75, 2465
 water requirement of, 95e-1
 weight regulation in, 275
 Electrical alternans, 1457-1458, 1572, 1572f
 Electrical injury, 1649, 2657
 Electrical remodeling, 273e-4
 Electrocardiography (ECG), 1450. *See also* specific disorders
 ambulatory, 273e-5
 in chest pain evaluation, 100-101
 clinical interpretation of, 1440, 1458-1459
 computerized, 1459
 electrophysiology in, 273e-1, 1451, 1451f
 intervals of, 1451, 1451f
 leads of, 1451-1452, 1452f
 normal, 269e-14f, 269e-15f, 273e-1, 278e-1f, 1452-1453, 1453f
 in overdose/poisoning, 473e-3
 P wave, 1452
 QRS complex, 273e-1, 1451, 1452-1453, 1453f
 in respiratory sinus arrhythmia, 278e-1f
 sensitivity and specificity of, 1451
 stress
 in arrhythmia evaluation, 273e-5
 in chest pain evaluation, 101-102, 270e-13-14
 contraindications to, 1582
 in ischemic heart disease evaluation, 1582, 1583f, 1595
 pharmacologic, 1582, 1585
 protocols for, 1585t
 ST segment elevations on, 1457, 1457f, 1457t
 T wave, 273e-1, 1453
 U wave, 1453
 waveforms of, 1451, 1451f
 Electrocautery, 247
 Electrochemical potential, 332e-6
 Electrocochleography, 223
 Electroconvulsive therapy, 2717
 Electrodiagnostic studies, 442e-1
 electroencephalography. *See*
 Electroencephalography (EEG)
 electrophysiologic studies. *See*
 Electrophysiologic studies
 evoked potentials. *See* Evoked potentials
 magnetoencephalography, 442e-3, 2550
 in myasthenia gravis, 2702
 in myopathies, 462e-4, 462e-4t
 nerve conduction studies. *See* Nerve conduction studies
 in peripheral neuropathy, 2676-2677, 2677t
 Electroencephalography (EEG), 442e-1, 442e-1f
 in brain death, 1776
 in coma, 442e-2, 1775
 continuous monitoring, 442e-2-3
 in delirium, 167
 in dementia, 442e-2
 in encephalitis, 442e-2
 in epilepsy, 442e-1-2, 442e-2f
 in neurally mediated syncope, 144f
 in neurologic critical illness, 1779
 in seizures, 2550
 Electrogenic transport, 332e-6
 Electrolysis, for hair removal, 334
 Electrolyte(s)
 for diarrhea, 267
 in enteric secretions, 98e-4t
 pancreatic secretion of, 2090
 parenteral requirement for, 98e-5t
 in specialized nutritional support formulas, 98e-4
 in stool, measurement of, 1933
 toxicity of, in transfusions, 138e-5
 Electrolyte disturbances. *See also* specific conditions
 delirium in, 169
 in hypothermia, 478e-2
 neurologic complications of, 463e-3-4
 Electromyography (EMG)
 in back pain, 114
 in fecal incontinence, 274
 in inflammatory myopathies, 2199-2200, 2199t
 in myasthenia gravis, 2702t
 in peripheral neuropathy, 2676-2677, 2677t
 principles of, 442e-4, 442e-4f
 single-fiber, 442e-6
 Electronutral transport, 332e-6
 Electronic medical records, 3, 30
 Electrophoresis, of serum proteins, 710-712, 712f
 Electrophysiologic studies, 442e-4
 blink reflexes, 442e-6
 cardiac. *See* Electrocardiography (ECG)
 electromyography. *See* Electromyography (EMG)
 F-wave studies, 442e-5
 H-reflex studies, 442e-5
 muscle response to repetitive nerve stimulation, 442e-5-6
 nerve conduction studies. *See* Nerve conduction studies
 in seizures, 2550
 Elementary body, 1166, 1166f
 Elephantiasis, 1418, 1419f. *See also* Lymphatic filariasis
 Eletriptan, 2591, 2592t, 2593t
 Eleuthero, 404t
 Elimination
 of disease, 785
 of drugs, 35
 ELISA (enzyme-linked immunosorbent assay), 214e-8
Elizabethkingia meningoseptica infections, 183e-2t, 183e-3
 Elliptocytes, 81e-1, 81e-4f
 Elliptocytosis, 81e-4f, 652f, 653, 653t
 El Moro Canyon virus, 1308t
 ELN gene mutations, 2513
 El Niño events
 disease outbreaks and, 151e-8, 151e-8f
 weather effects of, 151e-4, 151e-4f
 Elongin (pVHL), 2343
 Elsberg's syndrome, 2657
 Eltrombopag, 729
 Elvitegravir, 1275t, 1281
 Emanuel syndrome, 83e-5
 Embsen-Meyerhof pathway, defects of, 399
 Embolectomy, 1636, 1980, 1980t
 Embolic protection device, 296e-2
 Embolic stroke. *See* Ischemic stroke
 Embolism
 in acute limb ischemia, 1646
 cerebral arterial gas, 477e-7
 cutaneous, 361t, 362

- infectious, 369
 in infective endocarditis, 76e-12f, 818, 818f, 825
 mesenteric arterial, 1980, 1980t
 platelet, 40e-4
 pulmonary. *See* Pulmonary embolism (PE)
 retinal, 40e-4
 talc, 40e-4
- Embospheres, 550
- Embryologic development
 of blood vessels, 265e-1
 of cardiovascular system, 265e-11-12, 265e-11f
 of heart, 1520
 of kidney, 332e-1-2, 332e-1f, 333e-1
- Embryonal carcinoma
 ovarian, 594-595
 paraneoplastic syndromes in, 609t
 parasellar, 2264
 testicular, 588
- Embryonic carcinoma cells, 453t
- Embryonic germ cells, 453, 453t
- Embryonic stem cells, 454-455
 ethical issues, 90e-4
 genes in regulation of, 455
 patient-specific, 454
 potency of, 454f
 sources of, 90e-1-2, 453t
 in tissue engineering, 92e-1
 for tissue repair, 90e-1-2
- Emergent behavior, in biologic systems, 87e-2
- Emerin proteins, 1554
- Emery-Dreifuss muscular dystrophy, **462e-9**
 clinical features of, 462e-5t, 462e-9
 diagnosis of, 462e-9
 genetic considerations in, 435, 436t, 462e-5t
 pathogenesis of, 462e-9, 462e-9f
 treatment of, 462e-9
- Emesis, 103e-25, 258. *See also* Nausea and vomiting
- Emetine, 203, 246e-4t, 246e-8, 2199
- EMG. *See* Electromyography (EMG)
- Emmetropia, 195
- Emollients, 344, 346
- Empagliflozin, 2414t
- Empyema. *See also* Chronic obstructive pulmonary disease (COPD)
 centriacinar, 1701
 chest radiograph in, 308e-5f
 CT in, 308e-5f
 cyanosis in, 249
 diagnosis of, 1664f
 panacinar, 1701
 pathophysiology of, 306e-3
 pulmonary function values in, 306e-5-6, 306e-5f
- Empyematous cholecystitis, 2081
- Employee health service, 918
- Empty sella, 2257
- Empyema
 abdominal pain in, 106t
 actinomycotic, 1089, 1089f
 anaerobic bacterial, 1098-1099
 in cholecystitis, 2081
 definition of, 1717
 in pneumococcal pneumonia, 951
 in streptococcal pneumonia, 968
 subdural. *See* Subdural empyema
 tuberculous, 1110
- EMS. *See* Eosinophilia-myalgia syndrome (EMS)
- Emtricitabine (FTC)
 adverse effects of, 1274t
 for HIV infection, 1274t, 1277
 molecular structure of, 1276f, 1277
 resistance to, 1277, 1279f
- Enalapril, 371, 1507, 1512t
- Enalaprilat, 1627t, 1641
- Enamel, mottled, 236
- Enamel hypoplasia, 236
- Encephalitis
 amebic, 896
 with antibodies to cell-surface or synaptic proteins, 614t, 617, 617f
 approach to the patient, 883-884, 884-885f
 autoimmune, 896
 brainstem, 614t, 616
 in cancer patient, 488, 488t
 chronic
 progressive multifocal
 leukoencephalopathy. *See* Progressive multifocal leukoencephalopathy (PML)
 progressive rubella panencephalitis, 900
 subacute sclerosing panencephalitis, 900, 1298
 cortical, 616
 delirium in, 169
 EEG in, 442e-1f
 focal, 614t, 616-617, 616f
 global considerations in, 896
 granulomatous amebic, 1368
 hyponatremia in, 299
 in influenza, 1212
L. monocytogenes, 983
 limbic, 441e-1f, 616, 616f, 2607
M. pneumoniae, 1164
 measles-associated, 1298
 pinworm, 896
 in rabies, 1301
 in Rocky Mountain spotted fever, 1156
 toxoplasmic, 1401-1402, 1402f
 viral, **893**
 approach to the patient, 883-884, 884-885f
 arboviral, 1314-1317. *See also* West Nile virus infection
 clinical features of, 893
 CMV, 1192
 definition of, 893
 diagnosis of, 766t, 894-895, 895f, 895t
 differential diagnosis of, 895-896
 enteroviral, 1291, 1291t
 epidemiology of, 896
 etiology of, 891t, 893-894
 HHV-6, 1194
 HSV. *See* Herpes simplex virus (HSV) infections, encephalitis
 mumps-associated, 231e-2, 231e-3, 893
 sequelae of, 898
 tick-borne, 1316-1317
 treatment of, 896-898
- Encephalitozoon* spp., 245e-4t, 246e-9, 1408
- Encephalomyelitis
 acute disseminated (ADEM), 441e-26f, **2673**
 experimental allergic, 444e-5, 444e-6f
 paraneoplastic, 372e-25t, 614t, 616-617
 postinfectious, 1298, 2673
 with rigidity, 617
- Encephalopathy. *See also* Delirium
 chemotherapy-induced, 607, 607t
 chronic traumatic, 457e-4
 Hashimoto's, 441e-29f, 453e-5
 hepatic, **2066**
 asterix in, 1782
 MRI in, 441e-42f
 pathophysiology of, 1773
 physical examination in, 1992
 high-altitude. *See* High-altitude cerebral edema (HACE)
 HIV. *See* HIV infection, neurologic disease in
 hypertensive, 1615, 1626-1627, 2584
 hyponatremic, 300
 hypoxic-ischemic, 1773, 1781-1782, 1782f, 2559
 in Lyme disease, 1151
 in malaria, 780t, 783-784, 1373, 1373f, 1374t
 metabolic, 178, 1772-1773, 1782
 posterior reversible, 2568
 sepsis-associated, 1782
- Enchondromatosis (Ollier's disease), 426e-7, 594
- Enderterectomy
 aortoiliac, 1645
 carotid, 2571
- Endemic goiter, 2301
- Endemic syphilis, 207e-2, 207e-2f, 207e-2t, 207e-3
- Endemic (murine) typhus
 clinical features of, 127, 128t, 1155t, 1159
 diagnosis of, 1159
 epidemiology of, 1155t, 1158-1159
- etiology of, 1155t
 treatment of, 1159
- Endocarditis
 acute, 816
 infective. *See* Infective endocarditis
 Libman-Sacks, 2136
 marantic, 138, 367, 816
 nonbacterial thrombotic, 816, 2566t
 subacute, 816
- Endocrine, 2251
- Endocrine disorders, **2251**. *See also specific disorders*
 clinical evaluation of, 2252-2253
 fatigue in, 152
 hypothermia in, 478e-1, 478e-2t
 in lung cancer, 511
 myopathy in, 462e-18-19
 paraneoplastic. *See* Paraneoplastic syndromes, endocrinologic
 pathologic mechanisms of, 2251-2252
 in pregnancy, 47-48
 prevalence of, 2254t
 screening/testing for, 2253, 2254t
 types of, 2251, 2252t
- Endocrine glands, 2251. *See also specific glands*
- Endocrine tumors
 gastrointestinal. *See* Gastrointestinal neuroendocrine tumors (GI-NETS)
 pancreatic. *See* Pancreatic neuroendocrine tumors (pNETs)
- Endocrinology, **2251**
- Endodermal sinus tumor, testicular, 588
- End-of-life care, **55**. *See also* Terminally ill patient
 advance care planning in, 58-59, 59t
 assessment of patient in, 56-57
 for cancer patient, 119e-4, 475
 care during last hours, 68-69, 68f, 69t
 communicating bad news to patient in, 57--58, 57t
 continuous goal assessment in, 48
 epidemiology of, 55-56, 56f
 euthanasia in, 67-68, 67t
 futile care in, 67
 future directions, 70
 hospice care, 56, 70
 outcome measures for, 70
 physician-assisted suicide, 67-68, 67t
 principles of, 4
 withdrawing/withholding life-sustaining treatment during, 66-67. *See also* Withdrawal/withholding of care
- Endolymphatic hydrops. *See* Ménière's disease
- Endometrial cancer, **596**
 back pain in, 118
 clinical features of, 596-597
 deaths from, 467t, 469f, 596
 epidemiology of, 596
 genetic considerations in, 596
 granulosa cell tumor and, 594
 incidence of, 467t, 468f
 oral contraceptives and, 2391t
 postmenopausal hormone therapy and, 2383, 2384t, 2386
 prognosis of, 593t, 597
 risk factors for, 596
 as second malignancy in breast cancer survivor, 623
 staging of, 593t, 596
 tamoxifen use and, 531, 623
 treatment of, 597
- Endometriosis
 clinical features of, 118, 338, 2388
 definition of, 2388
 diagnosis of, 338
 etiology of, 338
 infertility due to, 2388
 vs. interstitial cystitis/bladder pain syndrome, 60e-3
 pulmonary, 246
 treatment of, 2389
- Endometritis
 actinomycotic, 1089
 alone, vs. with salpingitis, 877

- Endometritis (*Cont.*):
 anaerobic bacterial, 1909
C. trachomatis, 1169
 clinical features of, 338
 clostridial, 991t, 994
 etiology of, 338
- Endomyocardial biopsy, 1559, 1560f
- Endomyocardial fibrosis, 1567–1568
- Endomyometritis, postpartum, 50
- Endophthalmitis
Candida, 1343, 1343f, 1345
 clinical features of, 200, 201f
 etiology of, 200
Nocardia, 1086
P. aeruginosa, 1045t, 1046
- β-Endorphin, 386
- Endoscopic retrograde
 cholangiopancreatography (ERCP)
 in acute biliary obstruction, 1892
 in bile duct evaluation, 2085t
 in bile duct leak, 1885f
 in bile duct stones, 1884f, 1892
 in cholangiocarcinoma, 552–553, 1885f
 in cholestasis, 283–284
 in gallstone pancreatitis, 1893–1894
 indications for, 1878, 1879t, 1881
 in liver disease, 1993
 in pancreatic cancer, 555, 555f, 1894f
 in pancreatic disease, 2089
 in pancreatitis, 2095f
 pancreatitis after, 2091
 in primary sclerosing cholangitis, 1958, 2061
 procedure, 1881
 risks of, 1884
- Endoscopic secretin test, 2087t
- Endoscopic sphincterotomy, 346e-1f, 1881, 1884f
- Endoscopic therapy. *See also* Gastrointestinal endoscopy
 for achalasia, 1905
 for chronic pancreatitis, 2101
 for colonic pseudoobstruction, 1894f
 for esophageal varices, 277, 1890–1891, 1890f, 2063–2064, 2064f
 for gastric outlet obstruction, 1893f, 1894f
 for gastrointestinal bleeding, 277, 278f, 1890f
 for gastrointestinal malignancies, 346e-1v, 1885f, 1886f, 1887f, 1896
 for ileal stricture, 1884f
 indications for, 1880
 for intestinal obstruction, 1895f
 for Mallory-Weiss tears, 277
 mucosal dissection, 346e-1v, 1883–1884
 natural orifice transluminal endoscopic (NOTES) surgery, 346e-1v, 1881, 1883, 1886f
 for peptic ulcer disease, 277, 1890f
 sphincterotomy, 1884f
 submucosal dissection, 346e-1v, 1883–1884, 1887f, 1896
 suturing, 346e-1v, 1884, 1887f, 1888f
 for vascular ectasia, 1897, 1897f
- Endoscopic ultrasound (EUS)
 of bile duct, 1892, 1895f
 in bile duct evaluation, 2085t
 in chronic pancreatitis, 2100
 in esophageal disease, 1901
 with fine-needle aspiration, 1881, 1886f, 1900
 for gastrointestinal cancer staging, 1878, 1881, 1885f
 indications for, 1878, 1879t, 1881
 in pancreatic cancer, 554, 555
 in pancreatic disease evaluation, 2087t, 2089, 2089t
 for pancreatic necrosis, 346e-1v
 in pNETs, 572
- Endoscopy. *See* Gastrointestinal endoscopy
- Endosomes, 373e-5
- Endospores, 1335, 1335f
- Endostatin, 102e-17f
- Endothelial lipase, 2438
- Endothelial nitric oxide synthase (eNOS)
 in bicuspid aortic valve disease, 1529
 in brush cells, 213
 in cirrhosis, 286
 in female sexual response, 330
 in vascular function, 291e-2
- Endothelin(s)
 in cardiovascular function, 265e-2, 265e-4
 in edema, 251
 in male sexual function, 324, 325f
 in shock, 1745
 in thyroid function, 2285
- Endothelin receptor antagonists, 1615, 1659–1660, 1660t
- Endothelium
 antithrombotic effects of, 402
 biology of, 265e-1–2
 clinical assessment of, 265e-2f
 functions of, 265e-1–2, 265e-2t
 injury in sepsis/septic shock, 1754
 lymphocyte interactions with, 372e-26–28, 372e-29t
 in modulation of vascular tone, 1615
 in tumor angiogenesis, 102e-14–15, 102e-14f, 102e-15f, 102e-16f
- Endothelium-derived hyperpolarizing factor, 265e-1, 265e-4
- Endotoxin, 145e-9, 1752
- Endotoxin challenge test, 423
- Endotracheal intubation. *See also* Mechanical ventilation
 in cardiac arrest, 1769
 clinical procedure tutorial, 484e-1v
 for hemoptysis, 247
- Endotracheal suctioning, 810, 1467, 1467t
- Endovascular procedures. *See* Percutaneous interventions
- End-stage renal disease (ESRD). *See* Chronic kidney disease (CKD), stage 5
- End-systolic left ventricular pressure-volume relationship, 265e-10, 265e-10f
- Enema
 barium. *See* Barium radiography
 for constipation, 61, 62t
- Energy balance, 95e-1
 in aging, 73, 74f, 75
 assessment of, 73t
 physiologic regulation of, 415e-2, 415e-2f
- Energy density, of food, 2395
- Energy expenditure, 95e-1, 415e-2, 415e-5
- Energy requirements, 95e-1, 98e-4–5, 461, 462–463
- Enfuvirtide
 action of, 214e-3, 214e-9, 1281
 adverse effects of, 1275t, 1281
 for HIV infection, 214e-9, 1275t, 1281
 molecular structure of, 1277f
 resistance to, 1280f, 1281
- Enhanced external counterpulsation, 1513, 1592
- Enolase, 947f
- β-Enolase deficiency, 433e-2t, 462e-14
- Enophthalmos, 207
- eNOS. *See* Endothelial nitric oxide synthase (eNOS)
- Enoxaparin
 discontinuation before lumbar puncture, 443e-1
 for myocardial infarction, 1597, 1598t
 in PCI, 296e-1
 for VTE prophylaxis, 1637t
 for VTE treatment, 1635t
- Entacapone, 2615, 2616t
- Entamoeba dispar*, 1363
- Entamoeba histolytica*. *See also* Amebiasis
 adherence of, 145e-2t, 145e-3
 entry into host, 145e-8
 epidemiology of, 245e-4t
 immune response to, 1365
 laboratory identification of, 245e-1, 245e-3, 245e-4t, 245e-5t, 245e-6, 1365–1366
 life-cycle hosts of, 245e-4t
 life cycle of, 1363, 1364f
 phagocyte interaction with, 145e-7
 tissue invasion by, 145e-8, 1364, 1364f
 virulence factors of, 1364
- Entamoeba moshkovskii*, 1363
- Entecavir
 action of, 215e-9
 for chronic HBV infection, 215e-4t, 215e-10, 2036, 2037t, 2038f
 for HBV prophylaxis in transplant recipient, 927
 pharmacology of, 215e-9
 resistance to, 215e-10, 2040t
- Enteral feeding tubes, 98e-8, 98e-8t
- Enteral nutrition, **98e-1**. *See also* Specialized nutritional support
 combined with parenteral feeding, 98e-3
 complications of, 98e-8–9
 definition of, 98e-1
 disadvantages of, 98e-1, 98e-3
 in esophageal cancer, 533
 formulas for, 98e-8t
 for IBD, 1962
 indications for, 98e-3
 patient monitoring in, 98e-8
 for sepsis/septic shock, 1757
 tube placement for, 98e-1, 98e-8
- Enteric (typhoid) fever
 vs. *Brucella* infection, 194e-2
 clinical features of, 129t, 853, 1050–1051, 1051f
 diagnosis of, 1051
 epidemiology of, 134, 1050, 1050f
 etiology of, 1050
 global considerations in, 1050, 1050f, 1052
 multidrug-resistant, 1050, 1052, 1052t
 pathogenesis of, 853, 1049
 prevention and control of, 1052–1053
 resistance to, 145e-3
 skin manifestations of, 25e-3f, 362, 362t, 1051f
 treatment of, 1052, 1052t
- Enteric hyperoxaluria, 1943
- Enteric nervous system, 264
- Enteritis
 drug-induced, 1940t
 radiation, 103e-4, 1940t
 sexually acquired, 881
- Enteritis necroticans, 991
- Enterogaggressive *E. coli* (EAEC), 1028t, 1031
- Enterobacter* spp., 1034
- Enterobacter* spp. infections, **1034**
 antibiotic resistance in, 807, 933, 1027, 1034
 in cancer patient, 486t
 clinical features of, 1034
 diagnosis of, 1034
 endocarditis, 823
 epidemiology of, 1025
 health care–associated pneumonia, 803t
 osteomyelitis, 840t, 841t
 risk factors for, 805
 sepsis and septic shock, 1752
 sinusitis, 226
 treatment of, 1034
 ventilator-associated pneumonia, 915
- Enterobius vermicularis*/enterobiasis (pinworm), 245e-3t, 245e-5t, 1414t, 1416
- Enterocoele, 274
- Enterococcal infections, **971**
 bacteremia, 973, 975t
 bone and joint, 974
 in cancer patient, 486t
 endocarditis, 817t, 821–823, 822t, 973–974, 975t
 epidemiology of, 973
 etiology of, 971–972, 972f
 global considerations in, 973
 health care–associated, 973
 intraabdominal, 974
 meningitis. *See* Meningitis, bacterial
 neonatal, 974
 osteomyelitis, 841t
 pathogenesis of, 972–973
 soft tissue, 974
 in transplant recipient, 923, 927
 treatment of, 974–976, 975t
 urinary tract, 862, 973, 975t
 vancomycin-resistant. *See* Vancomycin-resistant enterococci (VRE)

- Enterococci. *See also* Enterococcal infections
antimicrobial resistance in, 150e-9, 941t, 976
characteristics of, 971-972
genome of, 776, 972-973
laboratory identification of, 971-972, 972f
virulence factors of, 972
- Enterococcus faecalis*, 941t, 972, 976. *See also*
Enterococcal infections
- Enterococcus faecium*, 941t, 972, 976. *See also*
Enterococcal infections
- Enterocolic fistula, 268
- Enterocolitis
necrotizing, 991
neutropenic, 488, 1099, 1797-1798, 1798f
radiation, 270
sexually acquired, 881
- Enterocytozoon bienersi*, 1257
- Enteroglucagon, pNET secretion of, 559t, 571
- Enterohemorrhagic *E. coli* (EHEC). *See also*
Escherichia coli infections, intestinal
characteristics of, 852, 1028t, 1030
epidemiology of, 1028t, 1030
food-borne, 856t, 857
global considerations in, 1030
- Enteroinvasive *E. coli* (EIEC), 1028t, 1031.
See also Escherichia coli infections,
intestinal
- Enterokinase, 2090
- Enterokinase deficiency, 1937
- Enteropathic arthritis, **2178**
clinical features of, 2178-2179
diagnosis of, 2179
epidemiology of, 2178
historic background of, 2178
pathology of, 2178
treatment of, 2179
- Enteropathogenic *E. coli* (EPEC), 852, 1028t,
1030-1031. *See also Escherichia coli*
infections, intestinal
- Enteropathy
gluten-induced. *See* Celiac disease (gluten-
induced enteropathy)
HIV, 1257
protein-losing, **1945**
- Enteropathy-type T-cell lymphoma, 135e-3, 708
- Enteroscopy, 346e-1v, 1881, 1883f
- Enterotoxigenic *E. coli* (ETEC). *See also*
Escherichia coli infections, intestinal
adherence of, 852
characteristics of, 1028t, 1030
epidemiology of, 1028t, 1030
food-borne, 856t
toxin production in, 853
- Enterotype analysis, 86e-5-6
- Enterovirus(es), 1289-1290. *See also*
Coxsackievirus
- Enterovirus infections, **1290**. *See also*
Coxsackievirus infections
acute hemorrhagic conjunctivitis, 1293, 1293f
clinical syndromes associated with, 1291t,
1293
diagnosis of, 1293
encephalitis, 1291
epidemiology of, 1290, 1293
exanthems, 1292, 1292f
generalized disease of the newborn, 1291
global considerations in, 1290, 1292
hand-foot-and-mouth disease. *See* Hand-foot-
and-mouth disease
herpangina, 1293
immunity to, 1290
meningitis, 892, 1291
myelitis, 2657
neurologic complications of, 1293
nonspecific febrile illness (summer grippé),
1291
pathogenesis of, 1290
pleurodynia, 1291
poliovirus. *See* Poliovirus infection
rash in, 127, 133t, 1292
treatment of, 1293-1294
upper respiratory, 226, 1202t, 1293
- Entesitis, 2219t
- Entomologic inoculation rate, 1371
- Entrance block, 273e-3
- Entrapment neuropathy, 49, 463e-4
- Entropion, 199, 1172
- Entry inhibitors. *See* Combination antiretroviral
therapy (cART)
- Enuresis, sleep, 192
- Envelope protein, HIV, 1216, 1219
- env* gene, 225e-1, 225e-2, 225e-3f, 1219, 1219f
- Environmental tobacco smoke. *See* Smoking,
passive exposure to
- Enzalutamide, 103e-15, 586
- Enzyme(s). *See specific enzymes*
- Enzyme immunoassay (EIA), 150e-2, 1245
- Enzyme-linked immunosorbent assay (ELISA),
214e-8
- Eosinopenia, 423
- Eosinophil(s), **422**
in adaptive immunity, 372e-7t
in asthma, 1673
disorders of, 422-423
functions of, 422
hematopoietic differentiation of, 89e-3f
in inflammation, 372e-8-9
in innate immunity, 372e-7t, 372e-8, 372e-14f
morphology of, 416f
in parasitic infections, 245e-4, 245e-6t, 422
in peripheral blood smear, 81e-6f
- Eosinophil chemotactic factor of anaphylaxis
(ECF-A), 372e-15t, 422
- Eosinophil granules, 422
- Eosinophilia. *See also* Hypereosinophilic
syndromes (HES)
clonal, 135e-7
diseases associated with, 422
drug-induced, 422
FGFR1-mutated, 135e-8
infectious causes of, 764, 765t
paraneoplastic, 612t, 613
parasites associated with, 245e-4, 245e-6t, 255e-1
PDGFR-mutated, 135e-8
primary, 135e-7, 135e-7t
tropical pulmonary, 422
- Eosinophilia-myalgia syndrome (EMS)
clinical features of, 376, 423, 2155, 2195
pathophysiology of, 423
L-tryptophan ingestion and, 14e-3, 376, 423,
462e-20, 2155
- Eosinophilic esophagitis, **1908**
clinical features of, 1909
diagnosis of, 1895, 1897f, 1909, 1909f
dysphagia in, 257, 1895
epidemiology of, 1908
etiology of, 1908
GERD and, 1908
histopathology of, 1909, 1909f
treatment of, 257, 1909
- Eosinophilic fasciitis, 376, 664, 2165
- Eosinophilic gastritis, 1931-1932
- Eosinophilic gastroenteritis, 270, 1939, 1940t
- Eosinophilic granulomatosis with polyangiitis
(Churg-Strauss syndrome), **2186**
ANCA in, 1685, 2180
clinical features of, 1685, 1840, 2186
diagnosis of, 386e-1, 386e-3f, 1685, 1840,
2186-2187
incidence and prevalence of, 2186
pathology and pathogenesis of, 2180, 2186
pericardial effusion in, 386e-3f
prognosis of, 1686
renal involvement in, 1840
treatment of, 1685-1686, 2187
- Eosinophilic leukemia, chronic, 135e-7-8,
135e-7t, 422, 672
- Eosinophilic myocarditis, 1562
- Eosinophilic myofasciitis, 2198
- Eosinophilic pneumonia. *See* Pulmonary
infiltrates with eosinophilia
- Eosinophilic pustular folliculitis, 1263
- Eotaxin (CCL11), 372e-12t, 1674, 1674f
- EPB41* gene mutations, 653t
- EPB42* gene mutations, 653t
- EPEC (enteropathogenic *E. coli*), 852, 1028t,
1030-1031. *See also Escherichia coli*
infections, intestinal
- Ependymomas, 441e-39f, 601
- Ephedra sinica* (ma huang), 14e-3, 2547t
- Ephedrine, 305, 473e-8t
- Ephelides, 359, 360t
- Ephrins, 102e-15, 102e-15f
- Epicondylitis, 2219t, 2249
- Epidemic polyarthritis (Ross River virus
infection), 837, 1313
- Epidemic (louse-borne) typhus
clinical features of, 128t, 782, 1155t, 1158
diagnosis of, 1158
epidemiology of, 127, 128t, 782, 1155t, 1158
global considerations in, 1158
prevention of, 1158
skin manifestations of, 128t
treatment of, 1158
- Epidemiologic transition, **266e-1**, 266e-1t
in United States, 266e-1-2
worldwide variations in, 93e-5, 93e-5f, 93e-5t,
266e-2
- Epidermal growth factor (EGF), 400e-1t, 1912,
1913f, 2285
- Epidermal growth factor receptor (EGFR), 39t,
41, 103e-16, 507, 1932. *See also EGFR*
gene mutations
- Epidermal growth factor receptor (EGFR)
antibodies, 103e-21, 103e-21t
- Epidermal growth factor receptor (EGFR)
inhibitors
action of, 103e-16, 103e-17t
adverse effects of, 103e-26, 356, 356t,
378-379
for head and neck cancer, 505
for lung cancer, 103e-16, 103e-17t, 507, 521,
521t
resistance to, 103e-23
- Epidermoid cysts, 76e-4f, 364, 603
- Epidermolysis bullosa, 257, 434, 2513
- Epidermolysis bullosa acquisita, 361t, 362, 370t,
373
- Epididymitis, 870t, 871-872, 1168, 1170t
- Epididymoorchitis, 194e-3, 231e-2
- Epidural abscess
anaerobic bacterial, 1098
clinical features of, 784
cranial, 904-905, 904f
etiology of, 780t
intracranial, 784
spinal, 117, 780t, 784, 2654, 2655f
treatment of, 767t, 780t, 2655
- Epidural anesthesia, 54
- Epidural hematoma, 441e-32f, 457e-3, 457e-3f,
2655
- Epidural metastases, 606, 606f, 1790
- Epidural spinal cord compression, 2653, 2653f
- Epidural spinal hemorrhage, 2584
- Epigenetic events
in cancer cells, 102e-7-8, 102e-8f
gene expression and, 101e-4, 429-430, 431f
- Epigenetic marks, 455
- Epigenetics, 102e-7, 444e-1-2
- Epigenome, 429
- Epigenomics, 6
- Epiglottitis, **234**, 235f, 1011, 1207
- Epilepsy, 334
- Epilepsia partialis continua, 616, 2542
- Epilepsy, **2542**. *See also* Seizure(s)
autonomic, 147
catamenial, 2559
coma in, 1773
continuous seizures in. *See* Status epilepticus
definition of, 2542
EEG in, 442e-1-2, 442e-2f
employment and, 2558-2559
etiology of, 2544-2545
age-related, 2545-2547, 2546t
channelopathies, 444e-2, 444e-2t
drug-related, 2547t
genetic factors, 444e-2t, 2545t, 2548
incidence of, 2542
interictal behavior in, 2557-2558
loss of driving privileges in, 2559
magnetoencephalography in, 442e-3
mortality data, 2558

- Epilepsy (*Cont.*):
 pregnancy and, 49, 2559
 psychiatric disorders associated with, 2558
 psychosocial issues in, 2558
 refractory, 2556–2557
 in schistosomiasis, 1427
 syndromes, **2544**
 age-related, 2545–2547
 genetic, 2545t
 juvenile myoclonic epilepsy, 2544
 Lennox-Gastaut syndrome, 2544
 mesial temporal lobe epilepsy, 2544, 2546f, 2546t
 treatment of, **2551**
 antiepileptic drug therapy. *See* Antiepileptic drugs (AEDs); *specific drugs*
 avoidance of precipitating factors in, 2551–2552
 management of underlying conditions, 2551
 in refractory disease, 2556–2557
 surgical, 2556–2557
 in women, 2559
- Epileptiform activity, 442e-1, 2550
- Epileptogenesis, 2548
- Epinephrine
 for anaphylaxis, 2117, 2749
 for brittle asthma, 1680
 for cardiac arrest, 1769, 1769f
 in cardiovascular regulation, 1613
 for drug-induced urticaria/angioedema, 381
 drug interactions of, 381
 in hypoglycemia, 2431t
 in local anesthetics, 242
 for parainfluenza virus infection, 1208
 as premedication for antivenom, 2737
 reference values for, 2759t
 for scambroid poisoning, 2744
 in shock, 1745
 in vascular smooth-muscle cells, 265e-4
- Epinephrine challenge test, 423
- Epirretinal membrane, 206
- Epirubicin, 103e-10t, 124e-4, 535
- Episcleritis, 200, 1958
- Episodic ataxia, 444e-2t, 451e-3t, 2629
- Epistaxis, 440
- Epistaxis
 approach to the patient, 403
 vs. hemoptysis, 245
 in nasopharyngeal cancer, 503
 in Osler-Weber-Rendu disease, 732
 in von Willebrand disease, 730
- Epistemic cells, 453t
- Epithelial cells/epithelium
 in adaptive immunity, 372e-7t
 airway, in asthma, 1673, 1674
 gastric, 1911–1914
 high-resistance, 332e-9
 intestinal, 86e-7, 1933
 pathogen encounters with, 145e-4–5, 145e-4f
 permeability of, 332e-3, 372e-23, 372e-25f
- Epithelial-mesenchymal transition, 102e-10, 102e-11f
- Epithelioid hemangi endothelioma, 552
- Epitopes, 377e-2
- Eplerenone
 adverse effects of, 64e-2
 drug interactions of, 205e-3
 for heart failure, 1511, 1512t
 for hypertension, 1623, 1624t, 1625
 for mineralocorticoid excess, 2319
- Epley maneuver, 150, 150f
- EPM2A* gene mutations, 2545t
- EPOCH regimen, 135e-2
- Epoprostenol, 1659, 1660t, 2164
- Epothilones, 103e-15, 531
- EPP. *See* Erythropoietic protoporphyria (EPP)
- Eprodiasate, 723
- Epsilometer (E-test), 150e-9
- Epsilon wave, 1495
- Epstein-Barr virus (EBV), 145e-2t, 214e-1, 214e-4, 1186–1187
- Epstein-Barr virus (EBV) infections, **1187**
 B-cell lymphoproliferative disease and, 214e-7, 623–624, 922, 1188
 Burkitt's lymphoma and, 101e-7, 477t, 695, 1188, 1375
 cancer and, 477t
 chronic fatigue syndrome and, 1188
 clinical features of, 1186–1187
 complications of, 1188
 diagnosis of, 150e-2, 150e-7, 233, 1188, 1189, 1189f, 1190t
 encephalitis, 894, 895t
 epidemiology of, 1187
 Hodgkin's lymphoma and, 214e-7, 697, 1189
 infectious mononucleosis. *See* Infectious mononucleosis
 lymphadenopathy in, 408
 lymphoid malignancies and, 135e-3, 135e-4, 214e-7, 488, 601, 697, 697t, 1188–1189
 meningitis, 892
 myelitis, 2657
 nasopharyngeal cancer and, 214e-7, 502, 503, 1188–1189
 neuropathy in, 2685
 oral hair leukoplakia, 1188, 1188f
 pathogenesis of, 1187
 pharyngitis, 231
 primary effusion lymphoma and, 135e-2
 reactivation in hypersensitivity syndrome, 382
 rheumatoid arthritis and, 2140
 skin manifestation of, 362, 362t
 SLE and, 2126
 systems biology applied to, 87e-3t
 thymoma and, 123e-2
 in transplant recipient
 after HSCT, 920t, 921t, 922–923
 after solid organ transplant, 924t, 925, 926
 treatment of, 926
 treatment of, 1189–1190
 vasculitis, 2193
- Epstein's syndrome, 729
- Eptifibatid
 action of, 746f, 748–749
 adverse effects of, 727t, 749
 discontinuation before lumbar puncture, 443e-1
 dosage of, 749
 genetic variations in response to, 277t, 743t
 indications for, 749
 for myocardial infarction, 1598t
 for PCI, 296e-1
 pharmacology of, 749t
- Epulis (gingival hypertrophy), 46e-2f
- Epulis fissuratum, 46e-2f
- Epworth sleepiness scale, 1721
- Equine antitoxin, for tetanus, 986
- Equine immunoglobulin, hyperimmune, 1328
- Eradication, of disease, 785
- erbB2* gene mutations
 in breast cancer, 101e-3, 524, 528, 531
 tumors associated with, 101e-2t
- ERBB4* gene mutations, 465e-4
- Erb's palsy, 48
- ERCC-6* gene, 94e-4
- ERCC-8* gene, 94e-4
- ERCP. *See* Endoscopic retrograde cholangiopancreatography (ERCP)
- Erectile dysfunction (ED), **326**
 after cystectomy, 577
 after prostatectomy, 583
 after radiation therapy, 583, 623
 alcohol use and, 2725
 approach to the patient, 327–328, 327f
 autonomic dysfunction and, 2639
 cocaine use and, 469e-1
 in diabetes mellitus, 2427
 drug-related, 326–327, 327t
 epidemiology of, 326
 pathophysiology of, 326–327
 prevalence of, 2254t
 screening/testing for, 2254t
 in systemic sclerosis, 2162
 treatment of, 328–330, 329t
- Erection, 324, 325f
- Ergot alkaloids, 473e-3, 473e-8t, 1649
- Ergotamine
 adverse effects of, 2591
 for exertional headache, 2597
 for migraine, 2591, 2592t, 2593t
 overdose/poisoning with, 473e-8t
 for sex headache prevention, 2597
- ERG* overexpression, 680, 680t
- Erlotinib
 action and targets of, 102e-3, 102e-5t, 103e-16, 103e-16f, 103e-17t, 103e-19
 adverse effects of, 103e-17t, 103e-26, 380, 380f, 1797
 for cholangiocarcinoma, 553
 for head and neck cancer, 505
 for hepatocellular carcinoma, 551
 for lung cancer, 102e-3, 102e-5t, 103e-16, 103e-19, 520t, 521, 521t
 for pancreatic cancer, 102e-5t, 557, 557t
 resistance to, 103e-23
- ermB* gene, 807
- Erosio interdigitalis blastomycetica, 1343
- Ertapenem
 for anaerobic bacterial infections, 1101
 for community-acquired pneumonia, 808t
 for health care-associated pneumonia, 812t
 indications for, 940t, 941
 in pregnancy and lactation, 937t
 resistance to, 933, 940t
- Erucism, 2750
- Erve virus, 1308t
- Erysipelas
 etiology of, 828t, 967
 pathophysiology of, 827f, 830
 skin manifestations of, 25e-4f, 76e-8f, 364t, 830, 967, 967f
 treatment of, 830, 965t, 967
- Erysipelothrix rhusiopathiae*, cellulitis, 830
- Erythema, 386
- Erythema annulare centrifugum, 355, 356t
- Erythema elevatum diutinum, 364t, 367
- Erythema gyratum repens, 355, 356f
- Erythema induratum, 366
- Erythema infectiosum (fifth disease)
 clinical features of, 128t, 1196
 diagnosis of, 1196t
 epidemiology of, 128t
 etiology of, 128t
 skin manifestations of
 buttocks, 25e-1f, 76e-8f
 facial, 25e-1f, 1196f
 progression of, 127, 128t, 362, 362t
- Erythema marginatum, 129t, 134, 356, 2152.
See also Rheumatic fever
- Erythema migrans. *See also* Lyme borreliosis (Lyme disease)
 clinical features of, 25e-2f, 76e-9f, 134, 355–356, 1150f
 evolution of, 1150
- Erythema multiforme
 in cancer patient, 487
 classification of, 361
 clinical features of, 340, 343f, 361, 361t
 differential diagnosis of, 130t
 epidemiology of, 130t
 etiology of, 361
 HSV infection and, 1178, 1182t
 oral manifestations of, 239t
 skin manifestations of, 25e-6f, 76e-14f, 130t, 134
- Erythema nodosum
 clinical features of, 132t, 366
 epidemiology of, 132t
 etiology of, 132t
 in IBD, 1957
 skin manifestations of, 25e-10f, 76e-15f, 132t, 135
- Erythema nodosum leprosum, 1124, 1124f, 1128
- Erythrasma, 343, 981
- Erythroblastosis fetalis, 236
- Erythrocyte sedimentation rate (ESR)
 in fever evaluation, 125
 in giant cell arteritis, 108

- in infectious diseases, 764, 766t
reference values for, 2755t
- Erythrocytosis, 399. *See also* Polycythemia
in cyanotic CHD, 1523
etiology of, 673, 673t, 1873
in hemoglobinopathy, 637
paraneoplastic, 612, 612t
stress (spurious), 673
testosterone therapy and, 2372–2373
- Erythroderma, 353
drug-induced, 353–354
etiology of, 353–354, 353t, 354t
in toxic shock syndrome, 780t, 782. *See also*
Toxic shock syndrome (TSS)
- Erythroid hyperplasia, 397, 397f, 626, 637
- Erythroleukemia, 81e-9f, 639
- Erythromelalgia (erythromalgia), 674, 677, 1648f, 1650
- Erythromycin
for acne vulgaris, 352
for actinomycosis, 1091t
action of, 931, 931t
adverse effects of, 261, 284, 2023, 2029
for bacillary angiomatosis, 832t, 1081t
for bite-wound infections, 167e-3t
for bronchiectasis prophylaxis, 1696
for *C. pneumoniae* infections, 1174
for *C. trachomatis* infection, 1171
for *Campylobacter* infections, 1061
for cellulitis, 832t
for chancroid, 1013
for cholera, 1064
for diphtheria, 979
for donovanosis, 198e-2t
drug interactions of, 34t, 943
azoles, 246e-3t
cisapride, 42t
cyclosporine, 42t
digoxin, 42t
PDE-5 inhibitors, 328
statins, 34t
theophylline, 1677
for gastroparesis, 260, 260t
indications for, 940t
for Lyme disease, 1153, 1153f
for pertussis, 1024t
in pregnancy and lactation, 937t
prophylactic, 945t
for psittacosis, 1173
for relapsing fever, 1148, 1148f
resistance to, 931t, 940t, 966
- Erythron, 392
- Erythrophagocytosis, in aggressive lymphoma, 81e-11f
- Erythroplakia, 237, 240t, 503
- Erythropoiesis
ineffective, 281, 398, 2000
iron-deficient, 627, 627f
megaloblastic, 81e-8f
physiology of, 392–393, 392f
- Erythropoietic protoporphyria (EPP)
clinical features of, 363, 390, 2522t, 2532–2533, 2533f
diagnosis of, 390, 2522t, 2533
global considerations in, 2521
pathophysiology of, 2522f
treatment of, 390, 2533
- Erythropoietin
action of, 392
in anemia, 392–393, 392f, 629–630
deficiency of, 398
ectopic production of, 400, 612
elevated, 400
in hematopoietic differentiation, 89e-3f
plasma, 392
pNET secreting, 559t, 571
regulation of production of, 392–393, 392f
- Erythropoietin therapy
adverse effects of, 103e-25, 668t
for anemia, 399
for anemia in chronic kidney disease, 1818
for chemotherapy-induced anemia, 103e-24–25
depot formulations, 103e-25
- for hypoproliferative anemia, 631
iron demand and, 631
for orthostatic hypotension, 146, 2644
- ESBL (extended-spectrum β -lactamase), 933, 1026
- Escape mutants, in HBV, 2008
- Eschar
in anthrax, 134t, 261e-2, 261e-3f
burn, 166e-2, 166e-2f
characteristics of, 127, 827f
in ecthyma gangrenosum, 132t
in rickettsialpox, 132t, 135
in scrub typhus, 128t, 135
- Escharate virus, 1309t
- Escherichia coli*
adherence of, 145e-2, 145e-2t, 852, 863
antibiotic resistance in, 807, 862, 934, 941t, 1026–1027
commensal strains, 1027
epidemiology of, 1025
extraintestinal pathogenic strains, 1027–1028
genome of, 776
host interactions of, 1025–1026, 1025t
intestinal pathogenic strains, 1028t, 1029–1030
enteroaggregative, 1031
enterohemorrhagic. *See* Enterohemorrhagic *E. coli* (EHEC)
enteroinvasive, 1031
enteropathogenic, 852, 1030–1031
enterotoxigenic. *See* Enterotoxigenic *E. coli* (ETEC)
Shiga toxin–producing. *See* Shiga toxin–producing *E. coli* (STEC)
laboratory identification of, 150e-2t
pili of, 145e-2, 863
structure and function of, 1025
toxin production by, 853
virulence factors of, 863, 1025
- Escherichia coli* infections, 1027
burn-wound, 166e-1
in cancer patient, 486, 486f, 486t
epidemiology of, 1025
extraintestinal
abdominal/pelvic, 846, 1028
bacteremia, 1029
cellulitis, 1028–1029
ecthyma gangrenosum, 369
endovascular, 1029
intrauterine, 50
meningitis, 1028
osteomyelitis, 839
perinephric abscess, 851
peritonitis, 288
pneumonia, 1028
treatment of, 1029
urinary tract, 862, 863, 866, 1028
- intestinal. *See also* Diarrhea, infectious;
Travelers' diarrhea
clinical features of, 266, 1028t
diagnosis of, 267, 1031
enterohemorrhagic: strain O157:H7, 267, 856, 1030, 1031, 1863. *See also*
Hemolytic-uremic syndrome (HUS)
epidemiology of, 855
etiology of, 1030–1031
vs. IBD, 1956
Shiga toxin–producing, 1864
transmission of, 1030
treatment of, 1031
intraabdominal, 1099
sepsis and septic shock, 1752
in transplant recipient, 919, 924
treatment of, 940t
- Escitalopram
adverse effects of, 2710t
for depression, 64, 2602, 2710t, 2716
for generalized anxiety disorder, 2711
- Esmolol
for aortic dissection, 1641
for arrhythmias, 1479t
dosage of, 1479t
for hypertension in subarachnoid hemorrhage, 1787
- for hypertensive emergencies, 1627t
for ischemic heart disease, 1589t
overdosage/poisoning with, 473e-9t
- Esomeprazole
for GERD, 263
for peptic ulcer disease, 1920, 1920t
for ZES, 1929
- Esophageal atresia, 1904
- Esophageal cancer, 532
adenocarcinoma, 532–533, 533t
Barrett's esophagus and. *See* Barrett's esophagus/metaplasia
clinical features of, 257, 533, 1904
deaths from, 467t, 532, 1903
diagnosis of, 533
endoscopic appearance of, 1907, 1907f
etiology of, 532–533, 532t, 533t
global considerations in, 532
H. pylori infection and, 1040
incidence of, 467t, 468, 532–533, 1903
prevention of, 476
prognosis of, 533
risk factors for, 262, 2725, 2730t
smokeless tobacco and, 476
squamous cell, 532–533, 532t
staging of, 1885f
treatment of, 91e-4, 533
- Esophageal disease, 1900
clinical features of
chest pain, 96t, 98, 1901, 1905
dysphagia, 256–257, 1901, 1904
globus sensation, 1901
heartburn, 262, 1876, 1900, 1901
indigestion, 262. *See also* Indigestion
odynophagia, 1901
regurgitation, 1901
water brash, 1901
congenital anomalies, 1904
dermatologic disorders and, 1911
diagnosis of, 1901–1902, 1902f
diverticula, 256, 259, 1903, 1903f
esophagitis. *See* Esophagitis
GERD. *See* Gastroesophageal reflux disease (GERD)
hiatal hernia, 1902, 1902f
iatrogenic injury, 1910
mechanical trauma, 1910–1911
motility disorders, 1904
achalasia. *See* Achalasia
diffuse esophageal spasm, 256, 1905–1906, 1906f
obstruction, 1891–1892, 1893f
rings, 1902, 1902f
Schatzki's ring, 256, 1895, 1896f, 1902
stricture, 256, 1895, 1896f
structural disorders, 1902–1904, 1902f, 1903f
in systemic sclerosis, 2161
tumors, 1904. *See also* Esophageal cancer
varices. *See* Esophageal varices
webs, 1902–1903
- Esophageal manometry
in diffuse esophageal spasm, 1906, 1906f
in dysphagia, 257
in GERD, 262
indications for, 1901
normal, 1902f
- Esophageal pseudodiverticulosis, 1903, 1903f
- Esophageal spasm
chest pain in, 96t, 98
diffuse, 256, 1905–1906, 1906f
jackhammer esophagus, 1907f
- Esophageal varices
bleeding in, 276, 276t, 277, 2063–2064, 2064f
endoscopic findings in, 1890, 1890f
portal hypertension and, 2063
in superior vena cava syndrome, 1788
treatment of, 346e-1v, 1890–1891, 1890f, 2063–2064, 2064f
- Esophageal web, 532
- Esophagectomy, for esophageal cancer, 533
- Esophagitis
in cancer patient, 488
clinical features of, 106t, 256–257, 1909
corrosive, 1910

- Esophagitis (*Cont.*):
 diagnosis of, 1894
 endoscopic findings in, 1896f
 eosinophilic. *See* Eosinophilic esophagitis
 in GERD. *See* Gastroesophageal reflux disease (GERD)
 in HIV infection, 1256, 1256f, 1909–1910, 1909f
 infectious, 1909
 Candida. *See* *Candida* spp. infections, esophageal
 CMV, 1896f, 1910
 HSV, 1179, 1182t, 1896f, 1910
 peptic, 1907f
 pill-induced, 257, 1910
 radiation, 1910
 treatment of, 263
- Esophagogastroduodenoscopy, 1881, 1881f, 1882f, 1901
- Esophagogastrosomy, 257
- Esophagoscopy, 533
- Esophagus
 anatomy of, 255, 1900
 Barrett's. *See* Barrett's esophagus/metaplasia
 "corkscrew," 1905, 1907f
 foreign bodies in, 1891–1892, 1893f, 1911
 functions of, 1900
 gastric juice effects on, 1906–1907
 jackhammer, 1907f
 scleroderma, 1911
 systemic disease effects on, 1911
- ESR. *See* Erythrocyte sedimentation rate (ESR)
- ESRD (end-stage renal disease). *See* Chronic kidney disease (CKD), stage 5
- Essential fructosuria, 433e-3t, 433e-5
- Essential mixed cryoglobulinemia, 2189.
See also Cryoglobulinemia
- Essential thrombocytosis (ET), 672, 676
- Essential tremor, 2612t, 2619–2620
- Estazolam, 473e-10t
- Esthesioneuroblastoma, 119e-4
- Estimated average requirement, 95e-1
- Estramustine, 103e-12t, 103e-15
- Estrogen(s)
 actions of, 2377f
 bone remodeling and, 2491–2492, 2492f, 2498
 breast cancer and, 524
 carcinogenicity of, 477t
 cardioprotective action of, 6e-2–3
 deficiency of, 6e-4
 excess of, 2366
 neuroprotective action of, 6e-2
 in perimenopause, 2381, 2381f
 production of, 2359f, 2377, 2377f
 reference values for, 2759
- Estrogen receptors
 action of, 2378, 2498
 in breast cancer, 102e-7, 526, 528
 in carcinoma of unknown primary, 120e-2t
- Estrogen therapy. *See also* Postmenopausal hormone therapy
 for adult hypopituitarism, 2259t
 adverse effects of, 2498, 2498f
 cholestasis, 2023, 2025t
 cutaneous, 353
 edema, 253t
 erectile dysfunction, 327
 hypertension, 1617t
 lipoprotein disorders, 2443–2444
 for amenorrhea, 337
 colorectal cancer and, 478, 539
 for contraception. *See* Contraception
 coronary heart disease and, 291e-7–8
 endometrial cancer and, 596
 for female sexual dysfunction, 331
 for hirsutism, 334
 for hypogonadism, 2261
 for migraine prevention, 2593t
 for osteoporosis, 2497–2498
 for Turner's syndrome, 2352
 vaginal, 2383, 2386
- Eszopiclone, 191, 215
- ET. *See* Essential thrombocytosis (ET)
- Etanercept
 action of, 372e-30
 adverse effects of, 348t, 422, 834, 2146t, 2173
 for ankylosing spondylitis, 2172–2173
 monitoring during treatment with, 2146t
 for psoriasis/psoriatic arthritis, 348t, 2177
 for rheumatoid arthritis, 2145, 2146t
 for sarcoidosis, 2211–2212
- ETEC. *See* Enterotoxigenic *E. coli* (ETEC)
- E-test, 150e-9
- Ethacrynic acid, 322, 1624t
- Ethambutol
 action of, 205e-4
 adverse effects of, 205e-4
 gout, 2218t
 neuropathy, 2687t
 optic neuritis, 205e-4
 optic neuropathy, 203
 thrombocytopenia, 727t
 zinc deficiency, 96e-9
 dosage of, 205e-4
 for MAC infection prophylaxis, 1251t
 for nontuberculous mycobacterial infections, 205e-4, 205e-8, 1131–1132, 1696
 pharmacology of, 205e-4
 resistance to, 205e-4, 1118
 for TB, 205e-2t, 205e-4, 1115–1117, 1115t, 1116t
 for tuberculous meningitis, 898
- Ethanol
 beverage. *See* Alcohol
 for ethylene glycol poisoning, 64e-9, 319
 injection, for hepatocellular carcinoma, 549
- Ethchlorvynol, 203, 473e-11t
- Ethical issues, 17e-1
 acting in best interests of patient, 17e-2
 avoiding deception, 17e-1
 clinical practice guidelines and, 17e-3
 confidentiality, 17e-1
 conflicts between patient's choices and their best interests, 17e-2
 conflicts of interest, 17e-3–4
 futile care, 67
 in genetic testing, 445
 in global health field experiences, 17e-3
 informed consent, 17e-1
 justice, 17e-2
 new technologies and, 7
 patient autonomy, 17e-1
 patient who lacks decision-making capacity, 17e-1–2
 personal values and, 17e-2–3
 professional oaths and codes and, 17e-2
 in social media use, 17e-3
 in stem cell therapy, 90e-4
 in team-based care, 17e-3
 virtue ethics, 17e-2
 work hour limitations and, 17e-3
- Ethinyl estradiol, 334, 2382, 2391. *See also* Estrogen therapy
- Ethionamide, 205e-6, 1116t
- Ethnic groups. *See also* Racial/ethnic disparities in health care
 drug responses in, genetic basis, 40
 waist circumference values in, 2394t
- Ethosuximide, 473e-10t, 2553t, 2556, 2765t
- Ethylendiaminetetraacetic acid (EDTA), 2237
- Ethylene glycol poisoning
 acidosis in, 64e-9, 319
 acute kidney injury in, 1804
 clinical features of, 473e-11t
 diagnosis of, 64e-9, 1805
 laboratory evaluation of, 2765t
 mechanism of, 473e-11t
 optic neuropathy in, 203
 treatment of, 64e-9, 319, 473e-11t
- Ethylene oxide, 2688t
- Ethynodiol diacetate, 334
- Etidronate
 for hypercalcemia, 610, 2481
 for osteoporosis management/prevention, 2499
 for Paget's disease of bone, 426e-3, 426e-3t
- Etomidate
 for Cushing's syndrome, 2273, 2318
 in mechanical ventilation, 1741
 overdose/poisoning with, 473e-10t
- Etoposide
 action of, 103e-13
 adverse effects of, 103e-10t, 103e-13, 590
 acute leukemia, 103e-13
 gonadal dysfunction, 103e-25
 nausea and vomiting, 103e-25
 neurologic, 1793, 2686t
 for breast cancer, 531
 drug interactions of, 1793
 for Ewing's sarcoma, 119e-4
 for gestational trophoblastic disease, 124e-5, 597
 for lung cancer, 522
 for ovarian germ cell tumors, 594
 for soft tissue sarcoma, 119e-2
 for testicular cancer, 590, 591
- Etravirine
 adverse effects of, 1274t, 1278
 for HIV, 1278
 for HIV infection, 1274t
 molecular structure of, 1276f
 resistance to, 1279f
- Etretinate, 2199
- Eukaryotes, in human microbiome, 86e-4
- Eumycetoma, 1355, 1356
- Eunuchoid proportions, 2360
- European bat lyssaviruses, 1304
- European fire ants, 2749
- EUS. *See* Endoscopic ultrasound (EUS)
- Eustachian tube dysfunction, 217
- Euthanasia, 67–68, 67t
- Euthyroid hyperthyroxinemia, 2286, 2286t
- EV1 overexpression, 680, 680t
- Evacuation disorders, 272
- Evaporative cooling, 479e-3
- Everglades virus, 1306t
- Everolimus
 action and targets of, 102e-5t, 103e-16f, 103e-18t, 103e-19, 1829t
 adverse effects of, 102e-5t, 103e-18t, 103e-19, 103e-26, 574, 1829t, 2071
 for breast cancer, 102e-5t, 103e-19, 530
 in drug-eluting stents, 296e-2
 for immunosuppression, 1829, 1829t, 2071
 for insulinoma, 569
 for NETs, 562, 574–575
 for reducing skin cancer risk in transplant recipients, 388
 for renal cell carcinoma, 102e-5t, 579
 therapeutic monitoring of, 2765t
- Evidence-based medicine, 3, 18, 24–26
- Evidence-Based Medicine Reviews, 25t
- Evoked potentials, 442e-3–4, 2665, 2665t
- Ewingii ehrlichiosis, 1155t, 1160
- Ewing's sarcoma, 119e-4
 diagnosis of, 119e-4
 genetic considerations in, 101e-3t, 119e-4
 incidence of, 119e-4
 oral manifestations of, 236
 treatment of, 119e-4
 tumor markers in, 473t
- Exanthems. *See* Rash
- Exanthem subitum (roseola), 25e-2f, 127, 128t, 1194
- Exchangers (antiporters), 332e-6
- Exchange transfusion, 636, 1386t, 1387
- Excitable gap reentry, 273e-3, 273e-4, 273e-4f
- Excitation-contraction coupling, genetic defects of, 1555t
- Excitotoxicity, 444e-8–9
- Executive functions, 183
- Exemestane, 103e-15, 478
- Exenatide, 2414, 2414t
- Exercise
 anaphylaxis due to, 2118
 for atherosclerosis prevention, 291e-8t, 291e-9
 for AV conduction block, 1473
 for back pain, 119, 120
 for cancer prevention, 476
 cardiac response to, 265e-8, 265e-9f

- cardiovascular disease and, 266e-4
 for diabetes mellitus, 2409
 for dysmenorrhea, 338
 for elderly, 84, 94e-7
 energy requirements for common activities, 1587, 1587t
 for fatigue, 153
 gonadotropin deficiency due to, 2364
 for heart failure, 1513
 for hypertension, 1622, 1622t
 for intermittent claudication, 1644
 for metabolic syndrome, 2453
 for osteoarthritis, 2231
 for osteoporosis management/prevention, 2497
 for PD, 2618
 for prevention of falls, 165
 tissue oxygen requirements in, 248
 for type 2 diabetes mellitus prevention/delay, 2406
 for weight loss, 2396
- Exercise-induced asthma, 1671
- Exercise-induced dyskinesia, 2624
- Exercise testing
 in chest pain evaluation, 101–102, 270e-13–14, 270e-14f. *See also* Echocardiography, stress; Electrocardiography (ECG), stress
 in heart failure, 1505
 in interstitial lung diseases, 1710
- Exercise tolerance
 in glycolytic defect disorders, 462e-14
 in ischemic heart disease, 1587
 preoperative, 51, 52t
- Exertional headache, primary, 107t, 2597
- Exertional heatstroke, 479e-3
- Exfoliatin, 829
- Exfoliative erythroderma syndrome, 25e-7f, 130t, 134
- Exocrine, 2251
- Exons, 427
- Exophthalmos, 207
- Exostoses, multiple, 101e-6t, 426e-7
- Exotoxin A, 1043, 1043t
- Expanded Disability Status Score (EDSS), 2668t
- Expedited partner therapy, 883
- Experimental allergic encephalomyelitis, 444e-5, 444e-6f
- Experimental allergic neuritis, 2695
- Exposomics, 6
- Expressivity, variable, 436–437
- Exserohilum rostratum* infection, 836
- EXT1/2* gene mutations, 101e-6t, 426e-7
- Extended daily dialysis (EDD), 1811
- Extended-spectrum β -lactamase (ESBL), 933, 1026
- Extensive metabolizers, 38f, 40
- Extensor plantar response, 2539
- External beam radiation therapy, 119e-2, 583
- External pressure headache, 2597
- Extinction (sensory phenomenon), 160, 180
- Extracellular adherence protein (EAP), 956
- Extracellular blood volume, 332e-10f
- Extracellular fluid, 295
- Extracorporeal blood rewarming, 478e-3, 478e-3t
- Extracorporeal membrane oxygenation (ECMO)
 for ARDS, 1739, 1739t
 clinical trials, 1742–1743
 for pulmonary edema, 1761, 1764
- Extracorporeal shockwave lithotripsy, 1869
- Extraembryonic endoderm cells, 453t
- Extramammary Paget disease, 502
- Extramedullary syndromes, 2652
- Extranodal NK/T-cell lymphoma, nasal type, 135e-3, 697, 708, 2184
- Extravascular space, 295
- Extremities
 arterial diseases of. *See* Arterial diseases, of extremities
 in cardiovascular examination, 1443, 1446, 1446f
 lymphatic disorders of. *See* Lymphedema
 venous disorders of. *See* Venous disorders, of extremities
- Extrinsic allergic alveolitis. *See* Hypersensitivity pneumonitis
- Extrinsic incubation, 1304
- Extrinsic pathway, coagulation, 401
- Extubation, of terminally ill patient, 66–67
- Eyach virus, 1306t
- Eye(s). *See also* Ocular disorders
 alignment of, 197
 “cherry red” spot in, 436e-7f
 dry. *See* Keratoconjunctivitis sicca
 injury to, 262e-3, 262e-3f, 262e-4–5
 melanoma of, 206, 206f
 movements of. *See* Ocular movements
 in nutritional deficiencies, 463t
 painful or red, 199–200
 structure of, 40e-1
 tumors of, 206–207
- Eye fields, frontal, 180, 180f
- Eyelids, disorders of, 199, 207–208
- Ezetimibe, 462e-19, 2448–2449, 2448t, 2453
- Ezogabine, 2553t
- F**
- Fab exchange, in IgG4-related disease, 391e-3
- Fabry’s disease, 432e-1
 cardiomyopathy in, 1565, 1566f
 carriers of, 437, 1846
 cerebrovascular disease in, 432e-1, 2568
 clinical features of, 432e-1, 432e-3t, 1835, 1846, 2680
 enzyme deficiency in, 432e-3t
 genetic considerations in, 432e-1, 1554, 1846, 2680
 neuropathy in, 2680
 pathogenesis of, 1846
 prevalence of, 432e-1
 renal biopsy in, 62e-7f, 1846
 skin manifestations of, 364t, 365, 432e-1, 436e-7f
 treatment of, 432e-1, 1565, 1846, 2680
- Face recognition, 182
- Facet fracture, 441e-31f
- Facet joint, 111
- Facet joint hypertrophy, 116–117
- Facial flushing, 352
- Facial hemiatrophy, 2649
- Facial myokymia, 2649, 2664
- Facial nerve
 anatomy of, 213f, 2647, 2647f
 examination of, 2538
 imaging of, 441e-24f
 injuries of, 220, 457e-2
 involvement in otitis externa, 229
 in taste sensation, 213
- Facial pain/numbness, 2645–2646
- Facial palsy, 157, 1152
- Facial weakness/paralysis
 anatomic considerations in, 2647, 2647f
 in Bell’s palsy. *See* Bell’s palsy
 in facial hemiatrophy, 2649
 in facial myokymia, 2649
 in hemifacial spasm, 2648
 in MS, 2663
- Facilitated diffusion, 332e-6
- Facioscapulohumeral muscular dystrophy (FSHD), 462e-1, 462e-2f, 462e-5t, 462e-11
- F-actin, 372e-24t
- Factitious illness
 characteristics of, 2719
 fever, 138
 hypoglycemia, 2434
 visual loss, 204
- Factor V
 deficiency of, 404, 733t, 736
 inhibitors, 732, 740
 screening assays, 405, 406f, 2755t
- Factor V Leiden (factor Va)
 early interventions for, 450t
 in fibrin clot formation, 401
 genetic testing for, 450t
 in pregnancy, 47
 protein C resistance and, 402
 thrombosis risk in, 404, 405f, 744
- Factor VII
 in coagulation cascade, 743
 deficiency of, 404, 736
 screening assays, 405, 406f, 2755t
- Factor VII deficiency, 404
- Factor VIII
 deficiency of, 403, 404, 736. *See also* Hemophilia
 inhibitors, 734–735, 740
 in pregnancy, 735
 replacement, for hemophilia, 734
 screening assays, 405, 406f, 2755t
 in von Willebrand disease, 730
- Factor IX
 deficiency of, 403, 404. *See also* Hemophilia
 inhibitors, 734–735, 740
 in pregnancy, 735
 replacement, for hemophilia, 734
 screening assays, 2755t
- Factor X
 deficiency of, 403, 404, 733t, 736
 inhibitors, 740
 screening assays, 405, 406f, 2755t
- Factor Xa, 401, 743, 752
- Factor XI
 deficiency of, 733t, 736
 in fibrin clot formation, 401, 401f
 inhibitors, 736, 740
- Factor XII deficiency, 733t
- Factor XIII deficiency, 733t, 736
- Facultative bacteria, 1094
- FADD (Fas-associated death domain), 102e-8, 102e-9f, 372e-26f
- Faget’s sign (relative bradycardia), 763, 763t
- Fall(s), 164
 in elderly, 76, 79–80, 80f, 162, 164–165
 epidemiology of, 162, 164
 hyponatremia and, 301
 mechanical, 164
 patterns of, 165
 prevention of, 165
 recurrent, 165
 reducing injury risk in, 165
 risk factors for, 76, 165, 165t
 sensory deficits and, 164
 vs. syncope, 147
- Fallopian tube cancer, 595
- False-negative tests, 21, 22
- False-positive tests, 21, 22
- Famciclovir
 adverse effects of, 215e-8
 for herpes zoster, 215e-3–4t, 215e-8, 832t, 1186
 for HSV infections, 215e-3t, 215e-8, 832t, 1182t, 1910
 for HSV prophylaxis in HIV infection, 1253t
 pharmacology of, 215e-8
 resistance to, 215e-8
 for varicella, 1185–1186
 for viral meningitis, 893
- Familial adenomatous polyposis (polyposis coli)
 clinical features of, 539
 colonic polyps in, 1897f
 colorectal cancer screening in, 1898t
 early interventions for, 450t, 451
 genetic considerations in, 101e-5, 101e-6t, 101e-7f, 538–539
 genetic testing for, 449, 450t
 polyp characteristics in, 538t
 screening for, 539
 treatment of, 539
- Familial amyloid polyneuropathy, 2681
- Familial amyotrophic lateral sclerosis, 2633, 2634t
- Familial benign hypocalciuric hypercalcemia (FBHH). *See* Familial hypocalciuric hypercalcemia (FHH)
- Familial chylomicronemia syndromes, 2439–2440, 2440t
- Familial cold autoinflammatory syndrome (FCAS), 372e-8t, 2213t, 2215
- Familial combined hyperlipidemia (FCHL), 2439
- Familial defective ApoB-100, 2440t, 2442
- Familial dementia with amyloid angiopathy and spastic ataxia, 451e-2t
- Familial dysalbuminemic hyperthyroxinemia (FDH), 2286, 2286t
- Familial dysbetalipoproteinemia, 2440t, 2443

- Familial focal epilepsy with variable foci, 2545t
 Familial gingival fibromatosis, 237
 Familial glucocorticoid deficiency, 2324t
 Familial hepatic lipase deficiency, 2440t
 Familial Hibernian fever. *See* TRAPS (tumor necrosis factor- α receptor-associated periodic syndrome)
 Familial hyperaldosteronism, 306. *See also* Glucocorticoid-remediable aldosteronism (GRA)
 Familial hypercholesterolemia, 436e-4f, 2440t, 2441–2442, 2449
 Familial hypertriglyceridemia, 2440
 Familial hypobetalipoproteinemia, 2444
 Familial hypocalciuric hypercalcemia (FHH)
 etiology of, 313
 genetic considerations in, 2340, 2467, 2475
 genetic testing in, 450t
 Familial inflammatory inclusion body myositis, 2195
 Familial isolated hyperparathyroidism (FIHP), 2340
 Familial Mediterranean fever, **2212**
 ascites in, 287
 clinical features of, 2213–2214, 2213t
 abdominal pain, 105, 106t, 2213
 in acute attacks, 2213–2214
 amyloidosis, 723, 1845, 2214
 arthritis, 2213
 cutaneous, 2214
 fever pattern/duration, 126, 2213–2214
 diagnosis of, 2214
 differential diagnosis of, 2214
 early interventions for, 450t
 genetic considerations in, 2212–2213, 2213t
 genetic testing for, 450t, 2214
 inflammasome activity in, 372e-8t
 interleukin 1 in, 422
 treatment of, 141, 723, 2213t, 2214
 Familial spastic paraplegia (FSP), **2636**
 clinical features of, 163, 2636, 2659
 genetic considerations in, 2634–2635t, 2636, 2659
 Family history, 1
 Family structure, 93e-4–5
 Famotidine, 263, 727t, 1920, 1920t
 FAN1 gene mutations, 1855, 1861
 Fanconi-Bickel syndrome (type IX GSD; glucose transporter 2 deficiency), 332e-7t, 433e-2t, 435e-1t, 2434–2435
 Fanconi's anemia, 434, 664
 Fanconi's syndrome, 320, 714, 2519
 Farber disease, 432e-4t
 Farmer's lung, 1681, 1681t, 1682, 1688t, 1691
 Farnesoid X receptor, 2076
 Farnesyl transferase inhibitors, 102e-4f
 Fas-associated death domain (FADD), 102e-8, 102e-9f, 372e-26f
 Fasciculation potentials, EMG, 442e-4
 Fasciculations, 154, 154t, 262e-5
Fasciculus Medicinæ, 2f
 Fasciitis
 eosinophilic, 376, 664, 2165
 necrotizing. *See* Necrotizing fasciitis
 palmar, 2247
 plantar, 2249–2250
Fasciola/fascioliasis. *See also* Biliary (hepatic) flukes
 clinical features of, 1429
 diagnosis of, 245e-2t, 245e-5t, 245e-6t, 765t
 epidemiology of, 245e-2t, 765t, 1428–1429
 life-cycle hosts of, 245e-2t
 treatment of, 246e-13, 1428t
Fasciolopsis buski, 245e-2t, 1428t, 1429. *See also* Intestinal flukes
 FAS gene mutations, 101e-6t
 FASL gene mutations, 101e-6t
 Fas ligand, 102e-8
 Fasting, periodic, 94e-6
 Fat(s)
 dietary, cancer and, 476, 538, 580
 malabsorption of, 269. *See also* Steatorrhea
 metabolism of, in type 2 diabetes mellitus, 2404–2405
 requirement for, 95e-1
 in specialized nutritional support formulas, 98e-4–5
 Fatal familial insomnia (FFI), 191, 453e-1t, 453e-4
 Fat embolism, 40e-4, 368, 1632
 fat gene, 415e-4, 415e-4t
 Fatigability, 152, 154
 Fatigue, **151**. *See also* Chronic fatigue syndrome
 approach to the patient, 153
 assessment of, 62, 464e-2f
 in biliary cirrhosis, 2061
 in chronic hepatitis, 2033, 2041
 definition of, 151–152, 462e-1
 differential diagnosis of, 152–153
 epidemiology of, 152
 etiology of, 63, 152
 in fibromyalgia, 2238
 global considerations in, 152
 health care errors and, 12e-1
 in heart failure, 1503
 in liver disease, 1991
 in MS, 2663, 2672
 in myasthenia gravis, 2701
 in myopathies, 462e-1–2
 prognosis of, 153
 radiation-induced, 103e-4
 in sarcoidosis, 2209
 vs. sleepiness, 188
 in terminally ill patient, 63–64, 69t
 treatment of, 64, 153
 Fat mass, aging and, 71
 Fat necrosis, 366
 Fatty acid(s), 1935, 1935t. *See also* Free fatty acids (FFAs); Lipid(s)
 Fatty acid diarrhea, 1935, 1935t
 Fatty liver disease
 alcoholic, 2053, 2053t. *See also* Alcoholic liver disease
 drug-induced, 2026t
 in IBD, 1958
 of pregnancy, 49
 Fatty streak, 291e-1, 291e-1f
 Favism, 655, 656f, 657t
 Fazio-Londe syndrome, 2636
 FBHH (familial benign hypocalciuric hypercalcemia). *See* Familial hypocalciuric hypercalcemia (FHH)
FBN1 gene mutations, 2513
 FCAS (familial cold autoinflammatory syndrome), 372e-8t, 2213t, 2215
 F cells, 632
FCGRIIA gene mutations, 2125
 Fc receptors. *See* Ig Fc receptors
 FDH (familial dysalbuminemic hyperthyroxinemia), 2286, 2286t
 Feather hydroid envenomation, 2741
 Febrile neutrophilic dermatosis. *See* Sweet syndrome
 Febrile nonhemolytic transfusion reaction, 138e-4–5
 Febrile seizures, 126, 2545–2546
 Febuxostat, 431e-5, 1871, 2235
 Fecal DNA testing, 481t
 Fecal immunochemical testing, 481t, 482
 Fecal impaction, 265, 268
 Fecal incontinence, **1975**
 clinical features of, 1975
 epidemiology of, 1975
 evaluation of, 1975–1976
 pathophysiology of, 265, 1975, 1975t
 rectal prolapse and, 1974
 in systemic sclerosis, 2164
 treatment of, 1976
 Fecalith, appendiceal, 1987, 1988f
 Fecal microbiota transplantation, 86e-10, 860, 860t
 Fecal occult blood
 in colorectal cancer screening, 28, 29t, 279, 481t, 482, 540
 endoscopic evaluation of, 1896–1897, 1897f
 Fecal patina, 916, 918
 Feces. *See* Stool
FECH gene, 2523t
FECH gene mutations, 2521
 Fechtner's syndrome, 729
 Fecundability, 2387
 Feedback control, 400e-5–6, 400e-5f
 Fee-for-service reimbursement, 21
 Felbamate, 473e-10t, 2553t, 2765t
 Feline esophagus, 1895
 Felodipine
 adverse effects of, 1589t
 for heart failure, 1513
 for ischemic heart disease, 1589, 1589t
 for Raynaud's phenomenon, 1649
 Felty's syndrome, 411, 418, 2138
 Female orgasmic disorder, 330
 Female pseudohermaphroditism. *See* 46,XX disorders of sex development
 Female reproductive system disorders
 cancer. *See specific organs*
 infertility. *See* Infertility
 menstrual disorders. *See* Menstrual disorders
 obesity and, 415e-6
 ovarian disorders. *See* Ovary, disorders of pelvic inflammatory disease. *See* Pelvic inflammatory disease (PID)
 sex chromosome disorders. *See* Sex chromosome disorders
 sexual dysfunction, 330–331, 330t, 2427
 tubal disease, 2388
 46,XX disorders. *See* 46,XX disorders of sex development
 Femoral artery
 for PCI access, 296e-1
 percutaneous intervention for occlusion of, 296e-5, 296e-6f
 pulse, in aortic regurgitation, 1536–1537
 Femoral-femoral bypass, 1645
 Femoral neuropathy, 2692
 Femoral-popliteal artery disease, 1643. *See also* Peripheral artery disease
 Femur
 atypical fracture of, 2501, 2501f
 ochronotic pigmentation of, 436e-7f
 FeNa (fractional excretion of sodium), 1807
 Fenfluramine, 2199, 2711t
 Fenofibrate, 431e-5, 2448t, 2453
 Fenopropfen, 91t
 Fentanyl
 adverse effects of, 2640t
 in mechanical ventilation, 1741
 for pain, 91t, 92
 transdermal, 92
 Fenugreek, 404t
FERM3 gene mutations, 419
 Ferriman and Gallwey scale for hirsutism, 332, 333f
 Ferritin
 formation of, 625
 serum, 393
 in alcoholic liver disease, 2517t
 factors affecting, 397
 in hemochromatosis, 2516f, 2517, 2517t
 in hypoproliferative anemia, 398, 630t
 in iron-deficiency states, 627f, 630t
 iron stores and, 628t
 in microcytic anemia, 628t
 normal, 397, 627f, 628, 628t, 2760t
 Ferroportin, 626
 Ferrous salts, for iron-deficiency anemia, 628–629, 629t
 Fertility, rates of, 93e-1–2
 Fesoterodine, for urinary incontinence, 81
 Fetal alcohol spectrum disorder, 2725
 Fetal alcohol syndrome, 2725
 Fetal hemoglobin
 developmental biology of, 632, 633
 hereditary persistence of, 633, 633t, 639
 manipulation of, for hemoglobinopathy, 639
 Fetal nigral dopamine cells, 2617
 Fetal posterior cerebral artery, 2574
 Fetal hepatic, 1992
 FEV₁ (forced expiratory volume in 1 s), 306e-5f, 306e-6

- Fever, **123**. *See also* Acute febrile illness; Fever of unknown origin (FUO)
 approach to the patient, 125
 in autoimmune/autoinflammatory disease, 126, 126t
 central, 1773
 in coma, 1773
 factitious, 138
 fraudulent, 138
 vs. hyperthermia, 124
 hypothalamic, 124
 hypoxia and, 248
 with myalgia, 1312t, 1317–1319
 pathogenesis of, 124–125, 125f
 Pel-Ebstein, 126
 in pregnancy, 50
 quadriplegic, 2660
 with rash, **25e-1**, **127**, 128–134t, 369–370, 998.
See also Rash
 in recipient of anticytokine therapy, 125
 recurrent, 126, 139–140, 140t. *See also*
 Hereditary recurrent fever syndromes
 relapsing. *See* Relapsing fever
 in returned traveler, 801–802, 802t
 in rheumatic fever, 2152
 transfusion-related, 138e-4–5
 treatment of, 126
 water loss in, 95e-1
- Feverfew, 404t
- Fever of unknown origin (FUO). *See also* Acute febrile illness
 approach to the patient, 139f
 first-stage diagnostic tests, 138–139
 later-stage diagnostic tests, 141
 PET, 140–141
 in recurrent fever, 139–140
 scintigraphy, 140
 definition of, 135
 differential diagnosis of, 136–138
 etiology of, 135–136, 136t
 global considerations in, 136, 136t
 prognosis of, 142
 recurrent, 140t
 treatment of, 141
- Fever syndrome, 372e-8t
- FFAs (free fatty acids), 265e-10
- FFI (fatal familial insomnia), 191, 453e-1t, 453e-4
- FGF8* gene mutations, 2364t
- FGF23* (fibroblast growth factor 23), 333e-3, 612, 1815
- FGFR1* gene mutations
 in eosinophilia, 135e-7, 135e-7t
 in hypogonadotropic hypogonadism, 2363, 2364t
 in lung cancer, 507t, 509
- FGFR1*-mutated eosinophilia, 135e-8
- FGFR2* gene mutations, 435
- FGFR3* gene mutations, 426e-7, 2512
- FGFR23* gene mutations, 2458
- FHH. *See* Familial hypocalciuric hypercalcemia (FHH)
- Fialuridine, 2025
- Fiber, dietary. *See* Dietary fiber
- Fiberoptic bronchoscopy, 512, 1667, 1710
- Fibric acid derivatives (fibrates)
 adverse effects of, 462e-19, 462e-20t, 2447, 2448t
 for hypertriglyceridemia, 2447, 2448t
 for metabolic syndrome, 2453
- Fibrillar, 372e-24t, 2157t
- Fibrillary-immunotactoid glomerulopathy, 1835t, 1845
- Fibrillation potentials, EMG, 155, 442e-4, 442e-4f
- Fibrillin aggregates, 2507
- Fibrin, 402, 725, 745, 746f
- Fibrin clot, 401, 401f
- Fibrin degradation products, 737
- Fibrin(ogen) degradation products, 402, 402f, 407
- Fibrinogen
 deficiency of, 404, 733t
 functions of, 402, 746f
- Fibrinolysis
 atherosclerosis and, 8–9
 in DIC, 737, 737f
 regulation of, 402–403, 402f
 thrombosis and, 744
- Fibrinolytic drugs, 757
 action of, 757f, 758
 for acute limb ischemia, 1646
 adverse effects of, 1605
 alteplase. *See* Alteplase (rtPA)
 anistreplase, 758
 contraindications to, 1605, 1636
 in development, 759–760
 indications for, 757
 for myocardial infarction, 1604f, 1605
 for pulmonary embolism, 1636
 reteplase. *See* Reteplase
 streptokinase. *See* Streptokinase
 tenecteplase. *See* Tenecteplase
 urokinase, 758–759
- Fibroblast growth factor-19 (FGF-19), 268
- Fibroblast growth factor-23 (FGF-23), 333e-3, 1815, 2455, 2458
- Fibroblasts, 102e-3
- Fibrocystic disease, of breast, 526
- Fibrocystin/polyductin, 1851f, 1853
- Fibrodysplasia ossificans progressiva, 426e-7t, 426e-8
- Fibroelastoma, papillary, 289e-2
- Fibrogenesis imperfecta ossium, 426e-6
- Fibroids, uterine (leiomyoma), 596
- Fibrolamellar hepatocellular carcinoma, 552
- Fibromas
 cardiac, 270e-24, 270e-24f, 289e-2
 ovarian, 594
 paraneoplastic syndromes in, 609t
- Fibromuscular dysplasia, 1617, 1627, 1645, 2567
- Fibromyalgia, **2238**
 approach to the patient, 2240
 clinical features of, 2199, 2221, 2238–2239, 2238f
 definition of, 2238
 diagnosis of, 2239, 2239t
 differential diagnosis of, 462e-2, 1152, 2239, 2239t
 epidemiology of, 2238
 genetic considerations in, 2239–2240
 in HIV infection, 1261
 overlapping syndromes, 2239
 physiology of, 2240
 tender point assessment in, 2238, 2238f
 treatment of, 2240, 2240t
- Fibronectin, 145e-3, 372e-24t
- Fibronectin-binding protein, 817
- Fibrosis
 airway, in asthma, 1674
 pulmonary. *See* Pulmonary fibrosis
 in systemic sclerosis, 2157
- Fibrous dysplasia, 426e-6–7, 426e-6f. *See also*
 McCune-Albright syndrome
- Fibrous histiocytoma, malignant, 119e-1.
See also Soft tissue sarcoma
- Fick method, cardiac output measurement, 1462–1463
- Fidaxomicin, 860, 860t
- Fiduciary obligations, 17e-2
- Fifth disease. *See* Erythema infectiosum (fifth disease)
- Figurate skin lesions, 354–355, 356t
- FIHP (familial isolated hyperparathyroidism), 2340
- Filaggrin, 344
- Filamentous hemagglutinin, 1021, 1022
- Filaria dance sign, 1419
- Filariasis, lymphatic. *See* Lymphatic filariasis
- Filiform warts, 351
- Filoviruses
 gene expression and replication in, 214e-4
 genome of, 1323
 global considerations in, 782
 phylogeny/evolution of, 1324f
 structure of, 214e-2t, 214e-3f, 1323
 taxonomy of, 1323f
- Filovirus infections. *See also* Ebola virus disease; Marburg virus disease
 clinical features of, 1327–1328, 1327t
 complications of, 1328
 control and prevention of, 1328–1356
 diagnosis of, 1328
 pathogenesis of, 1326–1327
 prognosis of, 1328
 treatment of, 1328
- FIM (Functional Independence Measure), 78t
- Fimbriae. *See* Pili (fimbriae), bacterial
- Financial incentives, 17e-3
- Finasteride
 for androgenetic alopecia, 355t
 for hirsutism, 335
 lower urinary tract symptoms, 7e-4
 for prostate cancer prevention, 478, 580
- Fine-needle aspiration biopsy, 103e-2, 409
- Finger(s)
 clubbing of. *See* Clubbing, of digits
 in gouty arthritis, 436e-2f
 in psoriatic arthritis, 2176, 2176f
- Finger anomia, 179
- Finger-to-nose testing, 2539
- Fingolimod, 2669–2670, 2669t
- Finkelstein's sign/test, 2220–2221, 2248
- Fire ants, 2116, 2749
- Fire coral envenomation, 2741
- First-pass elimination/effect, 33–34, 33f, 935
- FISH. *See* Fluorescence in situ hybridization (FISH)
- Fish, nematode infections from, 1412, 1416
- Fisher's syndrome, 41e-1v, 209
- Fish-eye disease, 2445
- Fish meal worker's lung, 1681t
- Fish odor syndrome (trimethylaminuria), 96e-6
- Fish oils. *See* Omega-3 fatty acids (fish oils)
- Fish tapeworm (*Diphyllobothrium latum*), 245e-2t, 645, 1434
- Fissure(s), anal, **1978**
- Fissured (scrotal) tongue, 46e-5f, 241t
- Fistula(s). *See also* Arteriovenous fistula
 coronary artery, 1523
 gallbladder, 2081
 hepatobronchial, 1365
 pancreaticopleural, 2095f
- Fistula in ano, 1111, **1978**
- Fitz-Hugh–Curtis syndrome (perihepatitis), 877, 1169
- Fitzpatrick skin types, 386, 386t
- 5-day fever. *See* Trench fever
- Fixed drug eruption, 381, 382f
- Flaccidity, 154
- Flagella
 of bacteria, 145e-2f, 145e-3, 145e-5
 of *P. aeruginosa*, 1043, 1043t
- Flail leaflet, 51e-4
- FLAIR. *See* Fluid-attenuated inversion recovery (FLAIR) MRI
- Flatus/flatulence, 285, 1945, 1965–1966
- Flat warts, 351
- Flatworm infections. *See* Cestodes/cestode infections; Trematodes/trematode infections
- Flaviviruses
 gene expression and replication in, 214e-4
 global considerations in, 782
 reservoirs and vectors of, 1305, 1306–1307t
 structure of, 214e-2t, 214e-3f
- Flavivirus infections
 clinical syndromes of, 1306–1307t
 dengue fever. *See* Dengue fever
 geographic distribution of, 1311–1312t
 Japanese encephalitis, 1315
 St. Louis encephalitis, 1315
 tick-borne viral encephalitis, 1315–1316
 viral hemorrhagic fevers, 1322–1323
 West Nile virus. *See* West Nile virus infection
 yellow fever, 8, 1322–1323
- Flavoenzymes, 96e-3
- Flavonoids, **96e-6**
- Flavopiridol, 102e-4f
- Flavor enhancers, 216

- Flea(s)
 bites, 2750
 cat, 1078, 2750
 dog, 2750
 in endemic typhus. *See* Endemic (murine) typhus
 in plague, 261e-4
 Flea-borne spotted fever, 1155t
 Flecainide
 action of, 273e-6t
 adverse effects of, 1480t, 1489
 for arrhythmias, 1480t
 for atrial fibrillation, 1489
 for AV nodal reentry tachycardia, 1480
 dosage of, 1480t
 for focal atrial tachycardia, 1479
 metabolism of, 34t
 pharmacology of, 1480t
 therapeutic monitoring of, 2765t
 for ventricular arrhythmias, 1498
 Flesh fly, 2747
 Flexible sigmoidoscopy. *See* Sigmoidoscopy
FLI1 gene mutations, 101e-3t, 119e-4
 Flinders Island spotted fever, 1157
FLNC gene mutations, 135e-1
 Floaters, 204
 Floppy-valve syndrome. *See* Mitral valve prolapse
 Flow limit, of lung, 306e-2
 Flow-volume loop, 306e-3f, 306e-6, 1675, 1675f
 Floxuridine, 2023
FLT3-ITD gene mutations, 679, 680, 680t, 685
FLT3-KTD mutation, 680t
 Fluclxacillin, 822t, 937, 2023
 Fluconazole
 advantages of, 1330
 adverse effects of, 246e-3t, 727t, 1330
 for *Blastomyces* infections, 1338, 1339t
 for *Candida* infections, 233
 disseminated, 1344t
 endocarditis, 1345
 esophagitis, 1344t, 1910
 liver abscess, 851
 thrush, 233, 487
 urinary tract, 868
 vulvovaginal, 873t, 875, 1332, 1344t
 for *Coccidioides* infections, 899, 1336
 for *Cryptococcus* infections, 899, 1264, 1341
 for cutaneous leishmaniasis, 1393
 drug interactions of, 34t, 246e-3t
 for febrile neutropenia, 491
 indications for, 1330
 for *Malassezia* infections, 1357
 in pregnancy and lactation, 246e-3t
 prophylactic use
 for *Candida* infections, 1345
 for *Cryptococcus* infections, 1341
 for fungal infections, 1330
 in HIV infection, 1252–1253t
 in transplant recipients, 139e-4, 139e-4t
 susceptibility testing for, 150e-8
 for tinea versicolor, 350
 Flucytosine
 action of, 1331
 adverse effects of, 417, 1331
 for *Candida* endocarditis, 824
 for *Candida* endophthalmitis, 1345
 for cryptococcal meningitis, 1264, 1331, 1341
 for *Cryptococcus* infections, 899
 indications for, 1331
 susceptibility testing for, 150e-8
 Fludarabine
 action of, 103e-14
 adverse effects of, 103e-11t, 103e-14, 103e-25, 488, 607, 607t
 for CLL, 703
 in HSCT preparation, 139e-2
 for Waldenström's macroglobulinemia, 718
 Fluidocortisone
 for adrenal insufficiency, 2326
 for orthostatic hypotension, 146, 2617, 2644
 for postural orthostatic tachycardia syndrome, 1477
 for SIADH, 2282
 for syncope, 145
 Fluidocortisone suppression test, 2319
 Fluid-attenuated inversion recovery (FLAIR) MRI
 in brain tumors, 440e-9f, 598, 600f
 characteristics of, 440e-4
 in Hashimoto's encephalopathy, 441e-29f
 in Huntington's disease, 441e-23f, 2622f
 in lacunar infarction, 441e-17f
 in limbic encephalitis, 441e-1f
 in multiple sclerosis, 441e-27f, 441e-35f
 in neurofibromatosis type 1, 441e-36f
 in seizures, 2550
 in viral encephalitis, 888, 895, 895f, 895t
 Fluid requirements, 95e-1, 98e-4
 Fluid restriction, 98e-4, 1739
 Fluid retention, 98e-4, 98e-7, 138e-5
 Fluid therapy
 for alcoholic ketoacidosis, 319
 for constipation, 61
 for diabetic ketoacidosis, 318
 for diarrhea, 267
 for DKA, 2419
 for HHS, 2420
 for hypercalcemia, 314, 2480t
 for hypovolemia, 298
 for hypovolemic shock, 1749
 for prevention of contrast-induced nephropathy, 1461
 for sepsis/septic shock, 1757
 Flukes. *See* Trematodes/trematode infections
 Flumazenil, in opiate overdose management, 468e-3
 Flunarizine
 adverse effects of, 2594t, 2611
 for hypnic headache, 2598
 for migraine prevention, 2593, 2594t
 Flunitrazepam, 469e-5, 473e-10t
 Fluorescence in situ hybridization (FISH)
 interphase, 83e-2, 83e-2f, 83e-3t
 metaphase, 83e-2, 83e-2f, 83e-3t
 for mutational analyses, 444
 probes for, 83e-2
 Fluorescent auramine microscopy, 245e-1
 Fluorescent treponemal antibody-absorbed (FTA-ABS) test, 1137, 1137f
 Fluoridation, 236
 Fluoride
 deficiency of, 96e-9t
 functions of, 96e-10
 for osteoporosis management/prevention, 2503
 recommended intake of, 95e-3t
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t, 96e-10, 310
 Fluorochrome stains, 150e-2
 Fluorocytosine, 431e-6
 Fluoroxyuridine, 431e-6
 Fluoroquinolones
 absorption of, 205e-6
 action of, 931t, 932–933, 943
 adverse effects of, 205e-6, 867, 867t, 938t, 943
 onycholysis, 380
 photosensitivity, 379, 389t
 vesicles/bullae, 361
 for anaerobic bacterial infections, 1101
 for bite-wound infections, 167e-2
 for *C. trachomatis* infections, 1171
 for cystitis, 867, 867t
 drug interactions of, 939t
 empirical uses of, 767t
 for *Enterobacter* infections, 1034
 for gram-negative bacterial infections, 183e-2t
 indications for, 943
 for *M. pneumoniae* infections, 1164
 in pregnancy and lactation, 937t
 prophylactic, 945t
 for pyelonephritis, 867
 resistance to
 in anaerobic bacteria, 1101
 in *Campylobacter* spp., 1061
 detection of, 773, 807
 in *E. coli*, 1029
 in gram-negative bacteria, 1027
 in *Klebsiella* spp., 1033
 mechanisms of, 931t, 943
 in mycobacteria, 205e-6, 1118
 in *Proteus* spp., 1033
 in *Shigella*, 1057
 in *V. cholera*, 1064
 for TB, 205e-6
 Fluoroscopy, in dysphagia, 256
 Fluorosis, 96e-10
 5-Fluorouracil (5-FU)
 action of, 103e-14
 adverse effects of, 103e-11t, 103e-14, 389t
 cardiotoxicity, 1563
 cutaneous, 361, 380
 diarrhea, 103e-25
 gastrointestinal, 1798
 nail disorders, 380
 nausea and vomiting, 103e-25
 neurologic, 607t, 1793
 for basal cell carcinoma, 502
 for breast cancer, 530
 for colorectal cancer, 543
 drug interactions of, 103e-11t
 for gastric cancer, 535
 for gastrinoma, 2337
 genetic variations in response to, 39t
 for head and neck cancer, 505
 for insulinoma, 2338
 for pancreatic cancer, 556t, 557
 in pregnancy, 124e-4
 pyrimidine metabolism and, 431e-6
 Fluoxetine
 adverse effects of, 2710t, 2716
 for cataplexy, 190
 for depression, 64, 473, 2672, 2710t, 2716
 drug interactions of, 34t, 40, 42, 42t
 metabolism of, 34t, 40
 for obsessive-compulsive disorder, 2714
 for pain, 94
 for panic disorder, 2709
 pharmacology of, 2716, 2717
 for Raynaud's phenomenon, 2164
 Fluoxymesterone, 103e-15, 2371
 Fluphenazine, 2722t
 Flurazepam, 473e-10t, 2711t
 Flushing
 in acne rosacea, 352
 in carcinoid syndrome, 564, 565
 differential diagnosis of, 566
 nicotinic acid therapy and, 96e-4
 Flutamide, 103e-15, 335, 585
 Fluvastatin, 2448t
 Flvoxamine, 2710t, 2714
 Fly bite, 2749–2750
 Fly infestation, 2746–2747
FMRI gene mutations, 336, 465e-1, 465e-2t
 Foam cells, 291e-2, 365
 Focal ataxia, 2627
 Focal atrial tachycardia, 1477, 1478f, 1479, 1479f
 Focal nodular hyperplasia, of liver, 553–554
 Focal segmental glomerulosclerosis
 clinical features of, 1835t, 1842
 genetic considerations in, 1832
 HIV-related, 1849
 pathogenesis of, 1842, 1842f, 1842t
 renal biopsy in, 62e-1f, 62e-2f, 1842
 treatment of, 1842–1843
 Focal seizures. *See* Seizure(s), focal
 FODMAPs, 1968–1969, 1968t
 Fogo salvagem, 371
 Folate, 640
 absorption of, 640–641
 for alcohol-induced acidosis, 64e-9
 cobalamin and, 641
 deficiency of. *See* Folate deficiency
 dietary sources of, 640
 functions of, 96e-2f, 641, 641t, 642f
 parenteral requirement for, 98e-5t
 red cell, 648
 reference values for, 2766t
 requirement/recommended intake of, 95e-2t, 641
 serum, 647–648

- structure of, 96e-2f
 supplements. *See* Folate supplements
 systems analysis of effects of, 87e-3
 transport in blood, 641
- Folate deficiency. *See also* Megaloblastic anemia
 anemia in, 398
 causes of, 646-647, 646f
 clinical features of, 96e-1t, 461, 642-643
 diagnosis of, 647-648
 hematologic findings in, 643-644
 neural tube defects and, 642-643
 neutrophil segmentation in, 415
 optic neuropathy in, 203
 peripheral blood smear in, 81e-3f
 in pregnancy, 265e-12, 642-643, 646
 in premature infants, 647
 prophylaxis for, 647-648
 psychiatric disturbance in, 643
 skin manifestations of, 360
 small-intestinal mucosal biopsies in, 1940t
 treatment of, 461, 647
- Folate supplements
 for colorectal cancer prevention, 539
 in infants and children, 649
 for methotrexate toxicity prophylaxis, 2182
 for paroxysmal nocturnal hemoglobinuria, 661
 in pregnancy, 29t, 47, 642, 648-649
 prophylactic, 648
 for pyruvate kinase deficiency, 654
 for thalassemia, 638
 for tropical sprue, 1943
- Folate synthesis antagonists, 931t, 932, 932f
- FOLFIRINOX regimen, 557, 557t
- FOLFIRI regimen, 543
- FOLFOX regimen, 543
- Folic acid, 640. *See also* Folate
- Folinic acid
 for folate deficiency, 648
 for megaloblastic anemia prophylaxis, 903
 for methotrexate toxicity prophylaxis, 2182
 for pancreatic cancer, 556t, 557
- Follicle(s)
 Graafian, 2376f
 ovarian, 2375f, 2376, 2376f
- Follicle-stimulating hormone (FSH)
 action of, 401e-4
 deficiency of, 2259t
 expression and regulation of, 401e-1t
 in females, age-related changes in levels of, 2376-2377, 2377f
 laboratory evaluation of, 2258t, 2760t
 in male pubertal development, 2358
 in menstrual cycle regulation, 2378-2379, 2378f, 2379f
 in ovarian follicle development, 2375f, 2376
 in perimenopause, 2381, 2381f
 secretion of, 401e-4
 in spermatogenesis, 2359-2360
 synthesis of, 401e-4
- Follicle-stimulating hormone (FSH) receptor, 437
- Follicle-stimulating hormone (FSH) therapy, 337, 2369
- Follicular dendritic cell(s), 135e-9, 1232
- Follicular lymphoma, 704
 clinical features of, 702t, 704
 epidemiology of, 704
 genetic considerations in, 101e-3t, 699t, 704
 immunology of, 698, 698f
 lymph node examination in, 81e-10f, 704f
 transformation to diffuse B-cell lymphoma, 705
 treatment of, 704-705
 tumor markers in, 135e-2t
- Folliculitis, 356
Candida, 1343
 clinical features of, 341t, 342f
 eosinophilic, 356
 etiology of, 828t, 829
 in HIV infection, 1262-1263
 hot-tub, 131t, 134, 356, 829, 1047
Malassezia, 1357
- Folliculitis decalvans, 355, 355t
- Fomepizole, 64e-9, 319, 1777
- Fomivirsen, 215e-2t, 215e-7-8, 215e-8
- Fondaparinux, 752
 action of, 750f, 753
 adverse effects of, 753
 for antiphospholipid antibody syndrome, 2136
 for DVT/PE treatment, 1635, 1635t
 for heparin-induced thrombocytopenia, 728
 vs. low-molecular-weight heparin, 753t
 for myocardial infarction, 1597, 1598t
 pharmacology of, 753
 for VTE prophylaxis, 55, 1637t
- Fonsecaea*, 1355
- Food exchange list, 95e-5
- Food frequency questionnaire, 95e-5
- Food guide, 95e-5
- Food insecurity, 461
- Food intake, in obesity, 415e-5
- Food-related illness
 bacterial, 855-856, 856t
B. cereus, 856, 856t
C. botulinum, 987-988
C. jejuni. *See* *Campylobacter* spp. infections
C. perfringens, 856, 856t, 991
E. coli. *See* *Escherichia coli* infections, intestinal
L. monocytogenes, 266, 983
 non-01/039 *V. cholera*, 1065-1066, 1065t
S. aureus. *See* *Staphylococcus aureus* infections, food-borne
Salmonella. *See* *Salmonella* spp. infections, gastroenteritis
Shigella. *See* *Shigella* spp. infections
V. cholerae. *See* Cholera
V. parahaemolyticus, 856t, 1065, 1065t
V. vulnificus, 782, 1065t, 1066
 ciguatera poisoning, 2742
 diarrhea in, 266, 855-856, 856t. *See also* Diarrhea, infectious
 diarrhetic shellfish poisoning, 2742-2743
 differential diagnosis of, 266, 266t
 domoic acid intoxication, 2743
 nematode-borne. *See* Nematodes/nematode infections
 paralytic shellfish poisoning, 2743
 scombroid poisoning, 2743-2744
 viral, 266t, 855
- Food webs, 86e-5
- Foot drop, 463e-4, 1124
- Foot examination, in diabetes mellitus, 2407
- Foot osteomyelitis, 845, 845f
- Foramen magnum syndrome, 2652
- Foramen ovale, 1520
- Foraminal herniation, 1772, 1772f
- Forbes syndrome (type IIIa GSD), 433e-2t, 433e-3
- Forced expiratory volume in 1 s (FEV₁), 306e-5f, 306e-6
- Forced vital capacity (FVC), 306e-5f, 306e-6
- Forchheimer spots, 127
- Fordyce spots, 127, 240t
- Forearm exercise test, 462e-5
- Foreign body
 aspiration of, 1768
 in esophagus, 1891-1892, 1893f, 1911
 in eye, 199
 granuloma, 367
- Formaldehyde exposure, 345, 1692t
- Formalin-ether sedimentation, 152e-1
- Formiminoglutamic aciduria, 434e-2t
- Formoterol, 1705
- 10-Formyl (CHO)tetrahydrofolate, 641t
- Fortification spectra, 204, 2535
- Fort Sherman virus, 1308t
- "Forward type," 138e-2
- Fosamprenavir, 1278, 1280f
- Foscarnet
 action of, 215e-8, 897
 adverse effects of
 electrolyte disturbances, 215e-8-9, 1193-1194
 hypokalemia, 306
 nephrotoxicity, 215e-8, 897, 1193-1194, 1804
- for CMV infection, 215e-2t, 215e-8, 922, 1193-1194, 2657
 for CMV prophylaxis in HIV infection, 1252t
 for HHV-6 infections, 215e-8, 922
 for HSV infections, 215e-8, 1182, 1182t
 for viral encephalitis, 897
- Fosfomycin
 action of, 930, 931t, 944
 adverse effects of, 938t, 944
 for *Citrobacter* infections, 1035
 indications for, 941t, 944
 for *Proteus* infections, 1034
 resistance to, 931t, 944
 for urinary tract infections, 866, 867t, 975t, 1029
- FOS gene mutations, 101e-2t
- Fosphenytoin, 1735
- Foster Kennedy syndrome, 216
- Founder effect, 435
- Fournier's gangrene, 831, 1099, 2199
- Fovea, 40e-1, 195
- Foveation, 195
- Foville's syndrome, 209
- Foxglove, 310, 473e-9t
- FOXL2 gene mutations, 594
- FOXO3A gene, 94e-4
- FOXP3, 377e-2, 2113, 2141, 2348
- FOXP3 gene mutations, 2348
- Fractalkine, 372e-13t
- Fractional excretion of sodium (FeNA), 1807
- Fractional flow reserve (FFR), 296e-4, 1465
- Fracture(s)
 atypical femur, 2501, 2501f
 Colles', 2488f
 hip. *See* Hip(s), fracture of
 hyponatremia and, 301
 IBD-related, 1958
 Lisfranc, 2244
 odontoid, 441e-33f
 in osteogenesis imperfecta, 2508
 osteoporotic. *See* Osteoporosis, fractures in
 Paget's disease of bone and, 426e-2
 pelvic, 2489
 risk assessment for, 2495-2496, 2496f
 skull, 457e-2
 teardrop, 441e-34f
 temporal bone, 220
 vertebral. *See* Vertebral fractures
 wrist, 2489
- Fragile X pre-mutation, 163
- Fragile X syndrome, 336, 434, 440t, 465e-1
- Fragile X tremor/ataxia syndrome, 451e-3t
- Frailty
 assessment of, 78-79, 78t
 consequences of, 77-79
 definition of, 77
 falls and, 163
 model of, 73f
 severity of, 77
- Framboesia. *See* Yaws
- Frameshift mutation, 432
- Framework Convention on Tobacco Control (WHO), 17
- Francisella* spp., 1066-1067
- Francisella tularensis*, 261e-7, 1066-1067. *See also* Tularemia
- Franklin's disease (gamma heavy chain disease), 718
- Frasier's syndrome, 2353
- Frataxin* gene, 2630
- Fraudulent fever, 138
- FRAX tool, for fracture risk assessment, 2495-2496, 2495f
- FRC (functional residual capacity), 306e-2, 306e-5f, 306e-6, 1733
- Freckles (ephelides), 359
- Free erythrocyte protoporphyrin, 2525
- Free fatty acids (FFAs)
 in cardiac metabolism, 265e-10
 in glucose utilization, 2405
 in insulin resistance, 2450, 2451f
 reference values for, 2760t
- Free radical theory of aging, 94e-3
- Free T₃ index, 2288

- Free T₄ index, 2288
 Free water deficit, 64e-4
 Freezing gait, 163, 165. *See also* Parkinson's disease
 Frenzel eyeglasses, 149
 Fresh-frozen plasma, 138e-2, 138e-2t, 736, 738, 739
 Friedreich's ataxia
 clinical features of, 451e-2t, 2629–2630
 genetic considerations in, 440t, 444e-9, 451e-2t, 1555t, 2630
 imaging in, 2629, 2629f
 SA node dysfunction in, 1467t
 treatment of, 2630
 Frontal abulic syndrome, 183–184
 Frontal bone fractures, 457e-2
 Frontal disinhibition syndrome, 183–184
 Frontal eye field, 209
 Frontal gait disorder, 163, 164t
 Frontal network syndrome, 184
 Frontotemporal dementia (FTD), **2602**
 ALS and, 2633
 behavioral variant, 183–184, 2603, 2603f
 clinical features of, 172, 173, 173t, 2602–2603
 functional anatomy of, 170
 genetic considerations in, 172t, 444e-1, 444e-9, 2603
 imaging in, 2603f
 molecular basis of, 172t
 nonfluent/agrammatic variant, 2603, 2603f
 pathogenesis of, 172t, 2603–2604, 2604f
 prions in, 453e-1
 semantic variant, 2603, 2603f
 syndromes
 corticobasal. *See* Corticobasal degeneration syndrome
 neuropathology of, 2604f
 in PD, 2605
 Pick's disease, 444e-8f, 2604, 2604f, 2605f
 progressive supranuclear palsy. *See* Progressive supranuclear palsy syndrome
 treatment of, 2604
 Frontotemporal lobar degeneration (FTLD), 2603–2604, 2604f
 Frostbite, **478e-3**
 clinical features of, 76e-5f, 478e-4, 478e-4f, 1648f, 1650
 pathophysiology of, 478e-4
 treatment of, 478e-4–5, 478e-4t, 1650
 Frostnip, 478e-4
 Frovatriptan, 2591, 2592t, 2593t
 "Frozen shoulder," 2249
 Fructose, 433e-1
 Fructose 1,6-biphosphate aldolase A deficiency, 433e-2t
 Fructose 1,6-biphosphate aldolase B deficiency, 433e-1f, 433e-3t, 433e-5
 Fructose 1,6-diphosphatase deficiency, 433e-1f, 433e-3t, 433e-5
 Fructose intolerance, hereditary, 433e-3t
 Fructose metabolism disorders, 433e-1f, 433e-3t, 433e-5
 Fructosuria, 433e-3t
 FSH. *See* Follicle-stimulating hormone (FSH)
 FSH β gene mutations, 2364, 2364t
 FSHD (facioscapulohumeral muscular dystrophy), 462e-1, 462e-2f, 462e-5t, 462e-11
 FSP. *See* Familial spastic paraplegia (FSP)
 FS (Functional Status) Score, 2668t
 FTA-ABS (fluorescent treponemal antibody-absorbed) test, 1137, 1137f
 FTC. *See* Emtricitabine (FTC)
 FTD. *See* Frontotemporal dementia (FTD)
 FTLD (frontotemporal lobar degeneration), 2603–2604, 2604f
 Fucosidosis, 432e-3t
 Fugue states, 2608
 Fukutin-related protein deficiency, 462e-9, 462e-10t
 Fukuyama congenital muscular dystrophy, 462e-9, 462e-10t
 Full liquid diet, 95e-5
 Fulvestrant, 530
 Fumagillin, 246e-4t, 246e-9
 Fumarate reductase, 246e-13
 Functional genomics, 426, 428
 Functional Independence Measure (FIM), 78t
 Functional MRI, 90f, 440e-8, 444e-12
 Functional redundancy, 86e-6
 Functional residual capacity (FRC), 306e-2, 306e-5f, 306e-6, 1733
 Functional somatic syndromes, 60e-2
 Functional status, 52t
 Functional Status (FS) Score, 2668t
 Fundoplication, 263, 1908
 Fundus, 40e-1
 in cardiovascular disease, 1443
 flat pigmented lesions of, 40e-6, 40e-6f, 40e-6t
 Fungal balls, 1347, 1348, 1349t
 Fungal infections, **1329**. *See also specific diseases*
 arthritis, 836
 in burn wounds, 166e-1, 166e-3
 in cancer patient, 491
 as cause of fever of unknown origin, 137t
 classification of, 1329
 diagnosis of, 1330
 endemic, 1329, 1329t
 epidemiology of, 1329
 infection control measures, 917
 mucocutaneous, 1357–1358, 1358t
 myocarditis, 1561
 opportunistic, 1329, 1329t, 1355–1357
 septic shock in, 781, 1752t
 terminology of, 1329
 in transplant recipient
 after HSCT, 920–921, 920t
 after solid organ transplant, 139e-4, 139e-4t, 926, 927, 1830–1831, 2072–2073
 treatment overview, 1330–1332. *See also specific drugs*
 Fungi
 adherence of, 145e-2t, 145e-3
 entry into host, 145e-1
 growth after entry, 145e-4
 in human microbiome, 86e-4
 laboratory evaluation of
 specimen collection and transport in, 150e-4t, 150e-5t
 specimen-processing algorithm for, 150e-6f
 staining in, 150e-2
 susceptibility testing in, 150e-8
 tissue invasion by, 145e-8, 145e-9
 Funnel chest (pectus excavatum), 1443, 2512
 Funnel-web spider, 2748
 FUO. *See* Fever of unknown origin (FUO)
 Furazolidone, 246e-4t, 246e-9
 Furosemide
 for acute kidney injury, 1810
 adverse effects of, 138, 322, 362, 379, 389t, 727t
 for ascites, 288, 2065
 for dyspnea, 47e-4
 for heart failure, 1510t, 1608
 for hypercalcemia of malignancy, 610
 for hypertension, 1624t
 for pulmonary edema, 62, 1761f, 1763
 for sepsis/septic shock, 1757
 for SIADH, 64e-5, 302, 610
 for uric acid nephropathy, 431e-5
 Furrier's lung, 1681t
 Furuncular myiasis, 2746
 Furunculosis, 228, 349
Fusarium spp., 1355
Fusarium spp. infections
 in cancer patient, 491
 clinical features of, 1356, 1356f
 diagnosis of, 1356
 epidemiology of, 1355–1356
 in transplant recipient, 920
 treatment of, 1356
 FUS gene mutations
 in ALS, 444e-1, 444e-9
 in essential tremor, 2619
 in frontotemporal dementia, 2603, 2604f
 Fusidic acid, 840t, 841t
 Fusiform aneurysm, 1637
Fusobacterium spp., 1094
Fusobacterium spp. infections. *See also*
 Anaerobic bacterial infections, mixed
 in cancer patient, 487
 in dog-bite wound, 167e-1
 in human bite-wounds, 167e-2
 infectious arthritis, 1100
 Lemierre's disease/syndrome. *See* Lemierre's disease/syndrome
 in lung abscess, 814
 peritonsillar abscess, 233
 pharyngitis, 231
 treatment of, 233
 FUS protein, 172t
 FUS/TLS gene, 2633, 2634t
 Futile care, 67, 1735
 FVC (forced vital capacity), 306e-5f, 306e-6
 F-wave studies, 442e-5
 FXR deficiency, 86e-9
 G
 G_{5a}, 2305t
 G6PC3 (glucose-6-phosphatase) deficiency, 375e-1
 G6PD deficiency. *See* Glucose-6-phosphate dehydrogenase (G6PD) deficiency
 G20210A mutation, 745
 GABA. *See* γ -aminobutyric acid (GABA)
 Gabapentin
 adverse effects of, 61, 2392, 2553t, 2682t
 for anxiety disorders, 2711
 for cluster headache prevention, 2595t
 for cough, 245
 for epilepsy, 2553t
 for fibromyalgia, 2240, 2240t
 for menopausal symptoms, 2383
 for migraine prevention, 2593
 for neuropathy, 61, 2682t
 for new daily persistent headache, 111
 overdosage/poisoning with, 473e-10t, 2765t
 for pain, 91t, 94, 2672
 for paroxysmal symptoms in MS, 2672
 pharmacology of, 2553t
 for restless legs syndrome, 192
 for SUNCT/SUNA prevention, 2596
 therapeutic monitoring of, 2765t
 GAD1 gene mutations, 465e-4
 GAD65 (glutamate decarboxylase), 372e-24t
 Gadolinium contrast, 270e-9, 379, 440e-5, 1819.
 See also Contrast agents
 Gadolinium exposure, 263e-3t
 gag gene, 225e-1, 225e-2, 225e-3f, 1218, 1219, 1219f
 Gag protein, 214e-5
 Gag reflex, 2538
 Gaisböck's syndrome, 400, 673
 Gait
 anatomy and physiology of, 162
 cerebellar, 616
 disorders, **162**
 approach to the patient, 164
 cautious gait, 163
 cerebellar ataxia, 163, 164t
 in elderly, 79–80, 80f
 etiology of, 162t
 freezing gait, 163, 165. *See also* Parkinson's disease
 frontal, 163, 164t
 in neuromuscular disease, 163–164
 psychogenic, 164
 sensory ataxia, 163, 164t
 stiff-legged, 163
 in toxic and metabolic disorders, 164
 video library, 33e-1v
 examination of, 2539
 Gait apraxia, 163
 Gait cycle, 162
 Galactokinase deficiency, 433e-1f, 433e-3t, 433e-5
 Galactomannan test, 1330, 1348
 Galactorrhea, 2267
 Galactose, 433e-1
 Galactose metabolism disorders, 433e-1f, 433e-3t, 433e-5

- Galactosemia, 336, 433e-3t, 433e-5
 α -Galactosidase A gene mutations. *See* Fabry's disease
- Galantamine, 176, 2601
- Gallavardin effect, 51e-1, 51e-3, 1450, 1531
- Gallbladder
 bile concentration within, 2076
 fistula formation in, 2081
 gangrene of, 2081
 hypomotility of, 2077
 perforations of, 2081
 porcelain, 2082
 strawberry, 2083
- Gallbladder disease, **2076**
 abdominal pain in, 103
 cancer, 285, 553
 chest discomfort/pain in, 96t, 98
 cholecystitis. *See* Cholecystitis
 clinical features of, 2079–2080
 congenital anomalies, 2076
 diagnosis of, 2078–2079, 2079f, 2079t
 gallstones. *See* Gallstone(s)
 hyperplastic cholecystoses, 2083
 natural history of, 2080
 postmenopausal hormone therapy and, 2384t, 2385
 in pregnancy, 49
- Gallium nitrate, 314, 607t
- Gallstone(s), **2076**
 asymptomatic, 2080
 biliary sludge and, 2077–2078
 cholesterol stones, 2076–2078, 2077f, 2078t
 clinical features of, 2079–2080
 complications of, 2080
 diagnosis of, 2078–2079, 2079t
 epidemiology of, 2076
 genetic considerations in, 2076–2077
 in hemolytic anemia, 399
 jaundice in, 281–282
 natural history of, 2080
 obesity and, 415e-6
 pathogenesis of, 2076
 pigment stones, 2078, 2078t
 prevention of, during weight loss, 2396
 treatment of, 2080
- Gallstone ileus, 1985, 2081–2082
- Gallstone pancreatitis, 1893–1894, 2096
- GALNT3* gene mutations, 2460
- GALT (gut-associated lymphoid tissue), 1228, 1232
- γ -aminobutyric acid (GABA)
 action of, 444e-3t
 alcohol and, 2723–2724
 clinical aspects of, 444e-3t
 metabolism, disorders of, 434e-2t
 in sleep promotion, 185, 186f
 structure of, 444e-3t
- γ -aminobutyric acid (GABA) agonists, 263
- γ -carboxylase, 736
- γ -glutamyl transpeptidase (GGTP), 1997, 2053, 2053t
- Gamma heavy chain disease (Franklin's disease), 718
- γ -hydroxybutyric acid (GHB)
 abuse of, 168, 469e-5
 action of, 469e-5
 for narcolepsy, 190
 overdose/poisoning with, 473e-10t
- Gamma knife surgery, 1792, 2263, 2646
- γ motor neurons, 155f
- γ radiation, 103e-4, 263e-1
- Ganciclovir
 action of, 214e-4, 215e-8, 897
 adverse effects of, 215e-8, 897, 2547t
 for CMV infection, 1193
 prophylaxis, 215e-8
 in transplant recipient, 139e-4t, 922, 926, 927
 treatment, 215e-2t, 215e-8, 2657
 pharmacology of, 215e-8
 resistance to, 215e-8
 for viral encephalitis, 897
- Gan Gan virus, 1306t, 1313
- Gangliocytomas, 2264
- Gangliogliomas, 601
- Ganglionopathy, dorsal root, 618, 2685
- Gangosa, 207e-2
- Gangrene
 anaerobic bacterial, 1099
 Fournier's, 831
 gallbladder, 2081
 gas. *See* Gas gangrene (clostridial myonecrosis)
 hemolytic streptococcal. *See* Necrotizing fasciitis
- GAPD (glyceraldehyde-3-phosphate dehydrogenase) deficiency, 654t
- Gap junctions, 444e-4
- Gardner-Diamond syndrome, 368
- Gardnerella vaginalis*, 150e-6. *See also* Bacterial vaginosis
- Gardner's syndrome
 clinical features of, 538t, 539, 599t
 genetic considerations in, 599t
 polyp distribution and characteristics in, 538t
 skin manifestations of, 364, 364t
 small bowel adenomas in, 536
- Garlic, 404t
- Gas, intestinal. *See* Flatus/flatulence
- Gas cysts, intestinal, 991
- Gas exchange
 in COPD, 1702
 diffusion of oxygen and carbon dioxide, 306e-3–4
 measurement of, 306e-6
 physiology of, 306e-3–4
 ventilation-perfusion heterogeneity, 306e-4, 306e-4f
- Gas gangrene (clostridial myonecrosis), **992**
 of bowel, 991
 clinical features of, 783, 992, 993f
 diagnosis of, 993, 2429
 etiology and pathogenesis of, 780t, 783, 831, 992–993, 993f
 histologic findings in, 993, 994f
 nontraumatic (spontaneous), 992, 993f
 prevention of, 994
 prognosis of, 994
 traumatic, 992
 treatment of, 780t, 832t, 833, 991t, 993–994
- GAS infections. *See* Group A *Streptococcus* (GAS) infections
- Gas-liquid chromatography, 150e-3
- Gastrectomy
 cobalamin deficiency after, 645
 complications of, 1926–1927
 for gastric adenocarcinoma, 535
 for gastric lymphoma, 536
 for peptic ulcer disease, 1925–1926
- Gastric acid, 853, 1912–1913, 1914f, 2090
- Gastric adenocarcinoma, **534**
 clinical features of, 534–535
 deaths from, 469f, 534
 diagnosis of, 535
 diffuse type, 534
 epidemiology of, 534
 etiology of, 534, 534t
 genetic considerations in, 101e-2t, 101e-6t, 1039
 global considerations in, 534
H. pylori infection and, 1039–1040, 1039f, 1922
 incidence of, 468, 534
 indigestion in, 262
 intestinal type, 534
 iron-deficiency anemia and, 535
 metastatic, 535
 pathology of, 534
 postgastrectomy, 1927
 spread by direct extension, 534
 staging of, 535t, 1885f
 treatment of, 535
 ulcers and, 534
- Gastric antral vascular ectasia, 2161, 2164
- Gastric bypass surgery. *See* Bariatric surgery
- Gastric emptying, delayed. *See* Gastroparesis
- Gastric fundic varices, 1891, 1891f
- Gastric glands, 1911, 1911f
- Gastric inhibitory peptide, ectopic production of, 609t, 611
- Gastric lavage, 319, 473e-6
- Gastric lymphoma, 535–536. *See also* Mucosa-associated lymphoid tissue (MALT) lymphoma
- Gastric neuroendocrine tumors (carcinoids), 560t, 564. *See also* Gastrointestinal neuroendocrine tumors (GI-NETs)
- Gastric outlet obstruction, 1892, 1893f, 1894f, 1918
- Gastric ulcers. *See* Peptic ulcer disease
- Gastric varices, 277
- Gastrin, 1928, 1928t
- Gastrinoma. *See also* Zollinger-Ellison syndrome (ZES)
 clinical features of, 269, 568
 in MEN 1, 1927–1928, 2336–2337, 2336t
 screening tests for, 1928, 1928t, 2337t
 treatment of, 272, 1929, 2337
- Gastritis, **1930**
 abdominal pain in, 106t
 acute, 1930, 1930t
 after cholecystectomy, 2083
 autoimmune, 372e-24t
 bleeding in, 277
 chronic, 1930–1931, 1930t
 classification of, 1930, 1930t
 eosinophilic, 1931–1932
 granulomatous, 1932
 halitosis in, 242
 lymphocytic, 1931
 multifocal atrophic, 1931
 stress-related, 277, 1929–1930
 treatment of, 1931
 type A, 1931
 type B, 1931, 1931f
 varioliform, 1931
- Gastroenteritis. *See also* Diarrhea, infectious vs. appendicitis, 1987
 eosinophilic. *See* Eosinophilic gastroenteritis
 food-related. *See* Food-related illness
Salmonella. *See* *Salmonella* spp. infections, gastroenteritis
 viral, **1285**
 adenovirus, 1286f, 1286t, 1289
 astrovirus, 1286f, 1286t, 1289
 vs. bacterial, 1287t
 calicivirus, 1286f, 1286t. *See also* Norovirus (Norwalk virus) infections
 clinical features of, 1287t
 diagnosis of, 1286t
 picobirnavirus, 1286f, 1289
 rotavirus. *See* Rotavirus infections
 sapovirus, 1285, 1286t
 torovirus, 1286f, 1289
 treatment of, 1287t
- Gastroesophageal junction, 1902, 1902f
- Gastroesophageal reflux disease (GERD), **1906**
 approach to the patient, 261–262
 asthma and, 1671
 clinical features of
 alarm symptoms, 262, 262t, 1894
 chest pain, 96t, 98, 99
 cough, 244
 dysphagia, 256
 esophagitis, 1894, 1907f
 extraesophageal, 1907
 indigestion/heartburn, 262, 1900, 1907
 nausea and vomiting, 259
 oral, 236
 complications of, 1907–1908, 1907f. *See also* Barrett's esophagus/metaplasia
 diagnosis of, 1894, 1896f, 1901–1902
 differential diagnosis of, 261, 1907
 eosinophilic esophagitis and, 1908
 epidemiology of, 261, 1906
H. pylori and, 1040
 mechanisms of, 261
 pathophysiology of, 1906–1907
 in systemic sclerosis, 2161, 2164
 treatment of, 263, 1908

- Gastrointestinal bleeding, 276
 approach to the patient, 278–279, 278f, 279f, 1885–1886
 aspirin-induced, 746
 in bleeding disorders, 404
 colonic diverticula and, 1891, 1893f
 in Crohn's disease, 1953
 diagnosis of, 278–279
 in Dieulafoy's lesion, 1891, 1891f
 differential diagnosis of, 1877t
 diverticular, 1971
 endoscopic evaluation of, 1885–1886
 from esophageal varices. *See* Esophageal varices
 in malabsorption syndromes, 1946t
 from Mallory-Weiss tears, 346e-1v, 1891, 1892f
 minor, endoscopic findings in, 1899, 1899f
 in peptic ulcer disease. *See* Peptic ulcer disease, complications of
 sources of
 colonic, 277, 1877. *See also* Hemorrhoidal disease; Rectal bleeding
 obscure, 279
 occult, 276, 279, 1896–1897, 1897f
 small intestinal, 277, 1877
 upper gastrointestinal, 276–277, 276t, 1877
 vascular ectasias and, 1891, 1892f
- Gastrointestinal cancer
 anal, 351, 543–544, 1198, 1199
 colorectal. *See* Colorectal cancer
 esophageal. *See* Esophageal cancer
 gastric. *See* Gastric adenocarcinoma
 gastric lymphoma, 535–536. *See also* Mucosa-associated lymphoid tissue (MALT) lymphoma
 gastric sarcoma, 536
 small bowel, 536–537
- Gastrointestinal disease, 1875. *See also specific diseases*
 approach to the patient, 1878–1879, 1879t. *See also* Gastrointestinal endoscopy
 classification of, 1875–1876
 clinical features of, 1876–1877, 1877t
 abdominal pain, 1877. *See also* Abdominal pain
 altered bowel habits, 1877. *See also* Constipation; Diarrhea
 ascites, 286
 gastrointestinal bleeding, 1877. *See also* Gastrointestinal bleeding
 heartburn, 1876
 involuntary weight loss, 275. *See also* Involuntary weight loss
 jaundice, 1877. *See also* Jaundice
 nausea and vomiting, 258, 1876–1877. *See also* Nausea and vomiting
 in diabetes mellitus, 2427
 genetic considerations in, 1876
 at high altitudes, 476e-4
 infections in cancer patient, 487–488
 in nerve agent exposure, 262e-5
 in pregnancy, 49
 in sepsis/septic shock, 1755
 in SLE, 2129t, 2131
 in sulfur mustard exposure, 262e-4
 in travelers. *See* Travelers' diarrhea
 treatment of, 98e-3, 1878–1879. *See also* Endoscopic therapy
- Gastrointestinal endoscopy, 1880. *See also specific types, e.g.,* Capsule endoscopy
 in anemia, 1896
 antibiotic prophylaxis for, 1884, 1888t
 antithrombotic drug management before, 1884, 1889t
 in Barrett's esophagus, 1881, 1882f, 1894–1895
 in biliary obstruction, 1885f, 1892
 in cancer staging, 1885f, 1900
 in celiac disease. *See* Celiac disease (gluten-induced enteropathy)
 in colonic pseudo-obstruction, 1892, 1894f
 for colorectal cancer screening. *See* Colonoscopy
 in diarrhea, 267, 1899
 in dyspepsia, 1894
 in dysphagia, 257, 1895–1896, 1896f
 in esophageal disease, 1901
 in gastric outlet obstruction, 1892, 1893f
 in gastrointestinal bleeding. *See* Gastrointestinal bleeding
 in gastrointestinal obstruction, 1891–1892, 1893–1894f
 in GERD, 1894, 1896f
 in hematochezia, 1899, 1899f
 indications for, 1878, 1879t
 in nausea and vomiting, 260
 in nonulcer dyspepsia, 1895
 open-access, 1900
 in pancreatitis, 1899–1900
 patient preparation for, 1900
 in peptic ulcer disease
 advantages of, 1919
 anesthesia for, 1881
 benign disease, 1919f
 duodenal ulcer, 1881f
 gastric ulcers, 1881f
 with hemorrhage, 1886, 1889f, 1890, 1890f
 procedure, 1881
 risks of, 1884
 therapeutic. *See* Endoscopic therapy
 upper tract, 262, 1881, 1881f, 1882f
 urgent, 1885
- Gastrointestinal hemorrhage. *See* Gastrointestinal bleeding
- Gastrointestinal motility test, 260
- Gastrointestinal neuroendocrine tumors (GI-NETs), 557. *See also* Pancreatic neuroendocrine tumors (pNETs)
 appendiceal, 561t, 563, 567
 carcinoid syndrome and. *See* Carcinoid syndrome
 cardiac manifestations of, 290e-1t, 290e-2–3, 1568
 classification of, 558, 560
 clinical features of, 268–269, 559t
 diagnosis of, 566, 2339
 gastric, 564
 general characteristics of, 557, 558t
 genetic syndromes associated with, 562, 562t
 hemoptysis in, 246
 incidence of, 559t, 560–561
 malignant behavior of, 559t, 560t, 561–562
 in MEN 1, 2336t, 2339
 metastatic, 269
 paraneoplastic syndromes in, 609t, 610, 619
 prognosis of, 561t, 573f
 rectal, 560t, 563, 567
 sites of, 537, 560, 560t
 small intestinal, 560t, 563, 567
 syndromes of, 559t. *See also* Carcinoid syndrome
 with systemic symptoms due to secreted products, 564
 treatment of, 567, 572–575, 2339
 tumor localization in, 571–572, 572f
 without carcinoid syndrome, 564
- Gastrointestinal obstruction. *See* Intestinal obstruction
- Gastrointestinal pseudoobstruction. *See* Intestinal pseudoobstruction
- Gastrointestinal stromal tumor (GIST)
 genetic considerations in, 102e-3, 119e-1
 metastatic, 119e-2
 pathology of, 119e-1
 treatment of
 angiogenesis inhibitors, 102e-17
 endoscopic resection, 346e-1v, 1886f
 imatinib, 102e-3, 536
 targeted chemotherapy, 102e-5t, 103e-19
- Gastrointestinal tract
 anaerobic flora of, 1094–1095, 1095t
 anatomy of, 1875
 extrinsic modulation of, 1875
 functions of, 1875
 microbiota in, 86e-7
 Gastrojejunostomy, 98e-8t
- Gastroparesis
 abdominal pain in, 262
 in chronic pancreatitis, 2101
 in diabetes mellitus, 2427
 diagnosis of, 263
 etiology of, 258
 indigestion in, 261, 262
 in mechanically ventilated patient, 1743
 nausea and vomiting in, 258
 in systemic sclerosis, 2161
 treatment of, 261, 1743
- Gastropathy, prolapse, 277
- Gastrostomy tube feeding, 98e-8t, 258
 GATA1 gene mutations, 135e-7, 729
 GATA2 deficiency, 420t, 2105–2106, 2105f
 GATA2 gene mutations, 670
 GATA4 gene mutations, 2354t
 GATA6 gene mutations, 2406
 Gatekeeper genes, 101e-5
 Gates Foundation, 8
 Gatifloxacin, 1115
 Gaucher cells, 432e-5
 Gaucher's disease, 432e-5
 cardiomyopathy in, 1565
 clinical features of, 432e-3t, 432e-5
 diagnosis of, 432e-5
 enzyme deficiency in, 432e-3t
 genetic considerations in, 432e-5
 myelofibrosis in, 672
 splenomegaly in, 411
 treatment of, 432e-5
 type 2, 432e-5
 type 3, 432e-5
- Gaze disorders
 horizontal, 209–210, 210f
 in MS, 2663
 supranuclear, 209
 vertical, 210
- Gaze stabilization reflex, 195
 GBA1 gene, 432e-5
 GBA gene mutations, 2612
 G-banding, 83e-1, 83e-1f, 83e-3t
 GBD 2010 (Global Burden of Diseases, Injuries, and Risk Factors Study 2010), 9
 GBS. *See* Guillain-Barré syndrome (GBS)
 GBS infections. *See* Group B *Streptococcus* (GBS) infections
 GB virus C (hepatitis G virus) infection, 1258
 GC regimen, for bladder cancer, 577
 G-CSF (granulocyte colony-stimulating factor), 372e-11t, 414f, 804
 G-CSF therapy. *See* Granulocyte colony-stimulating factor (G-CSF) therapy
 GDNF (glial-derived neurotrophic factor), 444e-7
- Gefitinib
 action and targets of, 103e-16, 103e-17t
 adverse effects of, 103e-16, 103e-17t, 380, 1797
 drug interactions of, 1793
 for head and neck cancer, 505
 for lung cancer, 102e-3, 102e-16, 520t, 521
 resistance to, 103e-23
- Gegenhalten (paratonia), 154
- Gemcitabine
 action of, 103e-14
 adverse effects of, 103e-11t, 103e-14, 577, 621, 1796, 1797, 1864
 for bladder cancer, 576, 577
 for breast cancer, 531
 for carcinoma of unknown primary, 120e-4
 for cholangiocarcinoma, 553
 for fibrolamellar hepatocellular carcinoma, 552
 for lung cancer, 519–520, 520t
 for ovarian cancer, 594
 for pancreatic cancer, 555, 556t, 557, 557t
 for soft tissue sarcoma, 119e-2
- Gemella morbillorum*, 821, 822t, 971
- Gemfibrozil
 adverse effects of, 2448t
 drug interactions of, 42t, 2199, 2449
 for hypertriglyceridemia, 2448t
 for metabolic syndrome, 2453

- Gemifloxacin
 action of, 931t, 932–933
 for community-acquired pneumonia, 767t, 808t
 for gonococcal infections, 1008
 indications for, 943
 resistance to, 931t
- Gemigliptin, 2414t
- Gemtuzumab, 683, 1795, 1796, 1798
- Gender bias, 16e-4
- Gender differences, 6e-1
- Gene(s). *See also specific genes*
 assortment and segregation during meiosis, 427–428, 430f
 definition of, 427
 mutations. *See* Mutation(s)
 regulatory regions of, 427, 428f
 structure of, 427
- Gene conversion, 434
- Gene expression
 alterations of, 91e-2t, 94e-3, 102e-7–8, 102e-8f
 in cancer cells, 101e-7
 epigenetic regulation of, 429–431, 431f
 regulation by transcription factors, 428–429, 428f, 431t
- Gene Expression Omnibus, 101e-7
- Gene network, 87e-3, 87e-6f, 87e-7f
- Generalized anxiety disorder, 2709, 2712t
- Generalized epilepsy with febrile seizures, 2545t
- Generalized seizures. *See* Seizure(s), generalized
- Gene silencing, 101e-4
- Gene therapy, 91e-1
 for adenosine deaminase deficiency, 91e-2
 for adrenoleukodystrophy, 91e-2–3
 for cancer, 91e-2f, 91e-3t, 91e-4
 for cardiovascular disease, 91e-2f, 91e-4
 complications of, 91e-2, 91e-2t
 delivery vehicles for, 91e-1t
 for genetic disorders, 91e-1–3
 for heart failure, 1515
 for hemophilia, 91e-3
 for immunodeficiency disorders, 91e-1–2, 423, 2108
 for Leber's congenital amaurosis, 91e-3
 for leukemia, 91e-4
 for LPL deficiency, 2440
 for macular degeneration, 91e-3
 for PD, 2617
 for sickle cell anemia, 635, 639
 strategies for, 91e-1
 taking history from subjects in clinical trials, 91e-5, 91e-5t
 for thalassemia, 639
 for therapeutic angiogenesis, 91e-4
 for vascular disease, 91e-4
 viral vectors for, 91e-1t, 214e-9, 225e-2
- Genetic counseling, 449
 directive, 449
 ethical issues in, 445
 in familial cancer syndromes, 101e-6–7, 101e-8f
 indications for, 449, 449t
 in mitochondrial DNA diseases, 85e-9
 nondirective, 449
 pretest, 449–450
- Genetic disorders. *See also specific disorders*
 adult-onset
 family history in, 446–447, 447f
 genetic testing for, 447–449, 448f. *See also* Genetic testing
 inheritance patterns in, 446
 therapeutic interventions for, 450–451, 450t
 animal models of, 431–432
 approach to the patient, 442, 444f
 chromosome disorders. *See* Chromosome abnormalities/disorders
 complex, 440–441
 databases/websites, 426t
 diagnosis of, 425
 family history in, 442–443
 identifying disease-causing gene in, 442t, 443
- Mendelian. *See* Mendelian disorders
 mitochondrial DNA. *See* Mitochondrial DNA diseases
 multifactorial, 425, 440–441, 441t, 446
 mutational analysis in, 444–445
 penetrance of, 446
 polygenic, 426+, 441t
 prenatal diagnosis, 444
 transmission of. *See* Mutation(s)
 variable expression of, 446
 X-linked. *See* X-linked disorders
 Y-linked, 439
- Genetic education, 449–450
- Genetic Information Nondiscrimination Act (GINA), 101e-6, 445, 449, 480e-6
- Genetic linkage, 441–442, 442t
- Genetic privacy, 445
- Genetics, 425
 applications in clinical medicine, 446
 databases/websites, 426t
 impact on medical practice, 425
 population, 442
- Genetic testing
 approach to, 448–449, 448f
 ethical issues in, 445
 for familial cancer syndromes, 101e-6–7, 101e-8f
 family relationships and, 448–449
 follow-up care after, 449
 informed consent in, 443, 449, 480e-5
 limitations to, 480e-5–6
 methods for, 444–445, 447–449
 predisposition testing, 448
 presymptomatic, 448
 screening programs, 443–444
- Gene transfer, horizontal, 769t
- Genistein, 580
- Genital tract, anaerobic flora of, 1094, 1095t
- Genital tract infections
 abscesses, 849
 anaerobic bacterial, 1099
 HSV. *See* Herpes simplex virus (HSV) infections, genital
 tuberculosis, 1110
 warts, 351, 829, 1198, 1199f, 1200–1201. *See also* Condyloma acuminata (anogenital warts)
- Genital ulcer(s)
 chancroid. *See* Chancroid
 clinical features of, 880t
 diagnosis of, 879–880
 donovanosis. *See* Donovanosis
 etiology of, 870t, 879
 HIV transmission and, 1221, 1221f
 HSV. *See* Herpes simplex virus (HSV) infections, genital
 lymphogranuloma venereum. *See* Lymphogranuloma venereum
 treatment of, 879t, 880
- Genome, 425, 769t. *See also* Human genome
- Genome-wide association studies (GWAS)
 advantages and limitations of, 442t
 of complex traits, 425, 442, 442f
 in neurologic diseases, 444e-1
 in pharmacogenomics, 38–39
- Genomic imprinting. *See* Imprinting
- Genomic medicine, 443
- Genomics, 5–6, 425. *See also* Pharmacogenomics
 comparative, 2
 databases/websites, 426t
 functional, 426, 428
 global health and, 445
 impact on medical practice, 425
 in infectious disease. *See* Infectious diseases, genomics of
 stem cell, 455
 terminology of, 769t
- Genotoxicity, 91e-2t
- Genotype, 435
- Genotype-phenotype correlation
 in common disease types, 87e-4f
 definition of, 435
 in sickle cell anemia, 87e-3, 87e-5f
 in von Hippel–Lindau disease, 435
- Genotyping, 40, 41
- Gentamicin
 action of, 930, 931t
 adverse effects of, 821, 942
 for *Bartonella* endocarditis, 1081t
 for *Brucella* infections, 194e-4
 for burn-wound infection, 166e-3
 for *Campylobacter* infections, 1061
 for donovanosis, 198e-2
 for health care–associated pneumonia, 767t, 812t
 indications for, 940t
 for infective endocarditis, 821, 822t, 823, 823t, 824, 963
 intratympanic, for Ménière's disease, 219
 for *L. monocytogenes* infections, 983
 for meningitis, 887t, 889, 889t
 for necrotizing fasciitis, 780t
 for pelvic inflammatory disease, 878t, 879
 for plague, 261e-5, 261e-5t, 1074–1075, 1074t
 in pregnancy and lactation, 937t
 prophylactic, 945t
 resistance to, 821, 931t, 934, 940t, 976
 for septic shock, 780t
 for skin and soft tissue infections, 832
 therapeutic monitoring of, 2765t
 for trench fever, 1081t
 for tularemia, 261e-5t, 261e-7, 1069–1070
- Genu valgum, 2221, 2228, 2228f
- Genu varum, 2221, 2228, 2228f
- Geographic tongue, 46e-3f, 241t, 342f
- Gephyrin, 372e-25t
- Gerbil bite-wound, 167e-1
- GERD. *See* Gastroesophageal reflux disease (GERD)
- Geriatric syndromes, 73f, 76, 79. *See also* Aging; Elderly
- German measles. *See* Rubella (German measles)
- Germ cell(s)
 ovarian, 2349, 2375–2376, 2376f
 primordial, 2349, 2375, 2375f
- Germ cell tumor
 carcinoma of unknown primary and, 120e-2t, 120e-4
 extragonadal, 120e-4, 591
 ovarian, 594–595
 testicular. *See* Testicular cancer
 tumor markers in, 473t
- Germinoma, 609t, 2264–2265
- Germiston virus, 1308t
- Germline stem cells, 453t
- Germ theory of disease, 761
- Gerstmann's syndrome, 179, 2573f
- Gerstmann–Sträussler–Scheinker (GSS) disease, 444e-10, 453e-1t, 453e-4
- Gestational diabetes insipidus, 303, 2276
- Gestational diabetes mellitus, 6e-3–4, 48, 2400, 2422
- Gestational hypertension, 46
- Gestational trophoblastic disease, 597
 clinical features of, 124e-5, 597
 diagnosis of, 124e-5
 global considerations in, 597
 incidence of, 124e-5
 inflammasome activity in, 372e-8t
 paraneoplastic syndromes in, 609t
 risk factors for, 597
 treatment of, 124e-5, 597
 tumor markers in, 473t
- Gfi-1, 89e-4
- GFR. *See* Glomerular filtration rate (GFR)
- GGTP (γ -glutamyl transpeptidase), 1997, 2053, 2053t
- GH. *See* Growth hormone (GH)
- GHB. *See* γ -hydroxybutyric acid (GHB)
- GH gene mutations, 434
- GH-N gene, 2258
- Ghrelin, 415e-2
 functions of, 1912–1913, 2357–2358
 GH release and, 401e-3
 pNET secreting, 559t
- GHRH (growth hormone-releasing hormone), 401e-3, 609t, 2258, 2265

- Giant cell arteritis (temporal arteritis), 2188
 as cause of fever of unknown origin, 138, 141
 clinical features of, 108, 2188
 aortic aneurysm, 386e-4f, 1638
 aortitis, 1642
 headache, 108
 ocular, 202, 202f, 2188
 oral pain, 239
 definition of, 2188
 diagnosis of, 2188–2189
 arteriogram, 386e-4f
 erythrocyte sedimentation rate, 108
 temporal artery biopsy, 386e-6f, 2188–2189
 incidence and prevalence of, 2188
 ischemic stroke in, 2567
 pathogenesis of, 1638, 2188
 treatment of, 141, 2189
- Giant cell myocarditis, 1561–1562
- Giant cell tumor, 119e-3, 609t
- Giant melanocytic nevus, 493
- Giardia* spp., 245e-4t, 1405–1406, 1405f
- Giardia* spp. infections, 1405
 clinical features of, 152e-4t, 266t, 1406
 acute disease, 266
 chronic disease, 269, 797
 diagnosis of, 152e-4t
 alternative procedures, 245e-5t
 blood and tissue, 245e-4t
 duodenal content analysis, 245e-1
 laboratory tests, 245e-4t
 serologic and molecular tests, 245e-6t, 267
 small-intestinal mucosal biopsies, 1939, 1941f
 stool analysis, 1406, 1406t
 differential diagnosis of, 266t
 epidemiology of, 266, 855, 1405–1406
 incubation period for, 152e-4t
 pathogenesis of, 1406
 prevention of, 1407
 in returning war veterans, 152e-8
 in travelers, 855t
 treatment of, 152e-4t, 1406
 albendazole, 1406
 furazolidone, 246e-9
 metronidazole, 268, 1406
 nitazoxanide, 246e-11, 1406
 quinacrine, 246e-12
 tinidazole, 246e-13
- Gibbus, 1110
- Giemsa stain, 83e-1, 245e-2, 342
- Gigantism. *See* Acromegaly
- Gilbert's syndrome, 2002
 clinical features of, 282, 2002–2003, 2003t
 diagnosis of, 1995, 1996f, 1998t, 2003t
 drug sensitivity in, 103e-13
 genetic considerations in, 40, 282, 2002–2003
 unconjugated bilirubin in, 280
- GINA (Genetic Information Nondiscrimination Act), 101e-6, 445, 449, 480e-6
- GI-NETs. *See* Gastrointestinal neuroendocrine tumors (GI-NETs)
- Ginger, 261, 404t, 1880
- Gingival erythema, linear, 237
- Gingival fibromatosis, idiopathic familial, 237
- Gingival hyperplasia, 237
- Gingival hypertrophy (epulis), 46e-2f
- Gingival recession, 46e-4f
- Gingivitis
 acute necrotizing ulcerative (Vincent's angina), 231, 233, 236, 238t, 1097
 clinical features of, 46e-3f, 46e-4f, 236
 etiology of, 1097
 prevention of, 423
 as risk factor for pneumonia, 805
- Gingivostomatitis, herpetic. *See* Herpes simplex virus (HSV) infections, oral-facial
- Ginkgo biloba*, 404t, 2547t, 2602
- Ginseng, 63, 404t
- GIRK4 channels, 306
- GIST. *See* Gastrointestinal stromal tumor (GIST)
- Gitelman's syndrome
 clinical features of, 306, 332e-8
 diagnosis of, 306, 307
 differential diagnosis of, 321
 genetic considerations in, 306, 332e-7t, 332e-8, 2461
- GJB2* gene mutations, 220
- Glanders, 1045t, 1049
- Glanzmann's thrombasthenia, 730
- Glasgow Coma Scale, 457e-5, 457e-5t, 1730t, 1777
- Glucocorticoids, 2668–2669, 2669t
- Glaucoma
 acute angle-closure, 200, 205
 clinical features of, 205
 headache in, 108
 low-tension, 205
 pathophysiology of, 204–205, 205f
 treatment of, 205
 visual field assessment in, 197–199
 visual loss in, 204–205
- Gleason grading system, for prostate cancer, 582
- Gliadin, 373e-7
- Glial-derived neurotrophic factor (GDNF), 444e-7
- Glibornuride, 2414t
- Gliclazide, 2414t
- Glimepiride, 2413, 2414t
- Glioblastoma(s), 440e-9f, 599, 600, 600f, 601f
- Glioblastoma multiforme, 91e-4
- Gliomas
 brainstem, 441e-39f, 601
 cerebral, 440e-9f
 EEG in, 442e-1f
 genetic considerations in, 101e-2t
 hypothalamic, 2264
 imaging in, 2262
 ocular signs in, 198
 optic, 2264
- Gliomatosis cerebri, 600
- Gliosarcomas, 601
- Glipizide, 34t, 40, 2413, 2414t
- Gliquidone, 2414t
- Global aphasia, 177t, 178
- Global burden of disease. *See* Global health
- Global Burden of Diseases, Injuries, and Risk Factors Study 2010 (GBD 2010), 9, 761
- Global Fund to Fight AIDS, Tuberculosis, and Malaria, 8
- Global health, 7. *See also* Low- and middle-income countries
A. baumannii infections, 1036, 1037
 acyclovir effectiveness, 1183
 addressing problems in, 17–18
 alcoholic liver disease, 2052
 alcohol use disorder, 2726, 2728
 amebiasis, 1363
 angiostrongyliasis, 1412
 antibiotic resistance, 966, 1101
 aplastic anemia, 663
 aspergillosis, 1345, 1346t
 ataxias, 2631
 azithromycin resistance, 233
B. mallei infections, 1049
B. pseudomallei infections, 1048
Babesia infections, 1384, 1385f
Bartonella infections, 1082
 biomass smoke-induced illness, 1693–1694, 1694f
 bioterrorism, 918
 blastomycosis, 1337
 botulism, 987
 brain abscess, 900
 brain drain, 13–14
 bronchiectasis, 1695
Brucella infections, 194e-1
C. difficile infections, 859
C. jejuni infections, 1059
C. trachomatis infections, 1167
 cancer, 16, 16f, 467–468, 472f, 490
Candidatus *Neoehrlichia mikurensis* infection, 1160
 cardiac transplantation, 1516
 cardiovascular disease, 15–16, 266e-2–3, 266e-3f, 266e-3t
 Carrión's disease, 1083
 cervical cancer, 595, 1198
 cholera, 1061–1062, 1062f
 chronic fatigue syndrome, 464e-1
 chronic kidney disease, 1821, 1822, 1825
 chronic myeloid leukemia, 694–695, 694f
 cirrhosis, 2059–2060
 climate change and. *See* Climate change
 clostridial infections, 990
 congenital syphilis, 1136
 cryptococcosis, 1339, 1341
 cryptosporidiosis, 1407
 cystic fibrosis, 1700
 diabetes mellitus, 15, 266e-4, 2400, 2400f
 dietary assessment, 95e-6
 diphtheria, 977, 979
 disease burden
 causes of, 10f, 11t
 health workforce and, 13e-4f
 by income, 13e-2f, 93e-5f, 93e-5t
 mortality and, 9, 11t, 266e-2f
 by region, 11t, 93e-5f, 93e-5t
 risk factors, 9–10
 disease eradication, 785
 disorders of sex development, 2357
 donovanosis, 198e-1
 drug hypersensitivity, 378
 EBV infections, 1187
 economics of, 8, 9
 encephalitis, 896
 endemic treponematoses, 207e-1, 207e-1f
 endemic typhus, 1158–1159
 enteric (typhoid) fever, 1050, 1050f, 1052
 enteritis necroticans, 991
 enterococcal infections, 973
 enterovirus infections, 1290, 1292
 epidemic typhus, 1158
 epidemiologic transition, 266e-1–2
 esophageal cancer, 532
 familial hypercholesterolemia, 2441
 fatigue, 152
 fever of unknown origin, 136, 136t
 field experiences in, ethical issues in, 17e-3
 flea-borne spotted fever, 1158
 fungal meningitis, 898
 gastric adenocarcinoma, 534
 genomics and, 445, 778
 glucose-6-phosphate dehydrogenase deficiency, 655, 655f
 glycogen storage diseases, 433e-5
 gnathostomiasis, 1412
 gonococcal infections, 1003, 1004
 gram-negative bacterial infections, 1026–1027
 group A *Streptococcus* infections, 233, 964, 964f
H. influenzae infections, 1010, 1012
H. pylori infection, 1042
 HBV infection, 2060
 HCV infection, 2059–2060
 head and cancer, 502
 heart failure, 1500–1501, 1516
 heavy metal poisoning, 472e-1, 472e-3
 helminthic infections, 255e-2
 hemoglobinopathies, 633
 hepatitis E infections, 1991
 hepatocellular carcinoma, 544–545, 544t
HFE gene mutations, 2519
 HHV-6 infections, 1194
 HHV-8 infections, 1194
 histoplasmosis, 1332
 HIV infection, 12, 12f, 13f, 1219–1220, 1220f, 1223–1225, 1224f
 HPV infection, 1198
 HSV infection, 1176
 HTLV-1 infections, 225e-4, 225e-4f
 HTLV-2 infections, 225e-5–6
 human granulocytotropic anaplasmosis, 1160
 hyperlipidemia, 266e-4
 hyperparathyroidism, 315
 hypersensitivity pneumonitis, 1683
 hypertension, 266e-4
 hyponatremia, 2283
 infectious disease-associated
 glomerulonephritis, 1847–1850
 inflammatory bowel disease, 1947
 inflammatory myopathies, 2197
 influenza, 1209, 1210

- institutions for, 8
 interstitial lung diseases, 1716
 iodine deficiency, 2285, 2285f
 iron-deficiency anemia, 626
 ischemic heart disease, 1578
 jaundice, 285
K. pneumoniae infections, 1031–1032
 kidney disease, 1799, 1862–1863
 laryngitis, 234
Legionella spp. infections, 1014, 1015
 leprosy, 1123, 1124, 2684
 leptospirosis, 1141
 liver abscess, 850
 Lyme borreliosis, 1151
 lymphatic filariasis, 1417, 1420
 lymphogranuloma venereum, 1169
 lyssaviruses, 1304
M. pneumoniae infections, 1164
 malaria, 14–15, 151e-4, 1370–1371, 1370f
 mastoiditis, 230
 maternal mortality, 51
 measles, 1295
 Mediterranean spotted fever, 1157
 meningococcal disease, 995–996, 997f
 meningococcal vaccines, 1001, 1002
 meningococcemia, 781–782
 mental disorders, 17, 2708
 Mesoamerican nephropathy, 1855–1856
 monkeypox, 220e-1–2
 MRSA infections, 918
 multiple myeloma, 713
 mumps, 231e-1, 231e-1f, 231e-3
 neurologic disease, 2535, 2535t
 nocardiosis, 1084
 noma, 1097
 nonalcoholic fatty liver disease, 2057
 nontuberculous mycobacterial infections, 1128–1129, 1132
 nontyphoidal *Salmonella* infections, 1053
 norovirus infections, 1285
 nutritional issues, 461
 obesity, 17, 266e-4
 opioid use disorders, 468e-1
 osteoarthritis, 2228
P. aeruginosa infections, 1043, 1045
 Paget's disease of bone, 426e-1
 parenteral nutrition, 98e-9
 parvovirus B19 infections, 1196
 PCP pneumonia, 1358
 pellagra, 2690
 peptic ulcer disease, 1042
 pertussis, 1021, 1021f, 1024
 plague, 1070–1071, 1071f, 1073
 pneumococcal infections, 949
 pneumococcal vaccine, 947
 poliovirus infections, 896, 1294, 1294t
 porphyrias, 2521, 2531
 prion diseases, 453e-1–2, 453e-1t
 pulmonary infiltrates with eosinophilia, 1687
 pulmonary paragonimiasis, 245
 Q fever, 1161
 rabies, 1300
 relapsing fever, 1146
 rheumatic fever, 2149
 rheumatic heart disease, 964f, 1501, 1528–1529, 2149
 rheumatoid arthritis, 1304, 2139, 2139f, 2149
 rickettsial diseases, 782, 1157
 Rocky Mountain spotted fever, 1154
 rotavirus infections, 1287, 1288f, 1289
 rubella, 230e-1, 230e-2–3, 230e-3f
 sarcoidosis, 2205
 schistosomiasis, 1424, 1425, 1425f
 scrub typhus, 1159
 selenium deficiency, 96e-10
 sexually transmitted infections, 869, 879
Shigella infections, 1055, 1057
 sleeping sickness, 1397
 snakebite care, 2738
Spirillum infection, 167e-2
 syphilis, 1133
 systemic lupus erythematosus, 2134
T. cruzi infections, 1396
T. gondii infections, 1404
 tapeworm infections, 1430
 testicular cancer, 588
 tetanus, 984, 986–987
 tobacco use, 17
 trachoma, 1172
 travel medicine, 793, 855
 trematode infections, 1424
 trichinellosis, 1410
 tropical pulmonary eosinophilia, 1420
 tropical sprue, 1942
 tuberculosis, 12–13, 205e-1, 778, 1103–1105, 1104f
 tularemia, 1067
 urinary tract infections, 862, 866
 vaccine-preventable disease, 785, 786
 valvular heart disease, 1528–1529
 vancomycin resistance rates, 973
Vibrio spp. infections, 1065, 1065t
 viral arthritis, 837
 viral hemorrhagic fevers, 782–783
 vitamin A deficiency, 96e-6
 vitamin D deficiency, 315
 Whipple's disease, 1091
 Wilson's disease, 2521
Yersinia spp. infections, 1076, 1077, 1078
 Global health equity, 10, 13e-4–5, 17
 Global Health Initiative, 13e-8
 Global Malaria Action Plan (GMAP), 14
 Globi, 1124
 Globin genes, 631, 631f, 632–633
 Globus pharyngeus, 254
 Globus sensation, 1901
 Glomerular diseases, **1831**. *See also specific diseases*
 acute nephritic syndromes. *See* Acute nephritic syndromes
 approach to the patient, 1834–1836, 1834t
 basement membrane syndromes. *See* Basement membrane syndromes
 clinical syndromes, 1835–1836, 1835–1836t
 genetic considerations in, 1832
 glomerular-vascular syndromes, 1835t, 1836, 1847–1848
 glomerulonephritis. *See* Glomerulonephritis
 hematuria in, 294, 1834, 1835t
 infectious disease-associated syndromes, 1836, 1836t, 1848–1850
 microalbuminuria in, 1834t
 nephrotic syndromes. *See* Nephrotic syndromes
 pathogenesis of, 1832–1834, 1833f
 progression of, 1834
 proteinuria in, 1834, 1834t, 1835t
 pulmonary-renal syndromes, 1835t, 1836, 1846
 renal biopsy in, 62e-1–6f, 1836–1837
 Glomerular filtration rate (GFR)
 age-related decline in, 1812
 assessment of, 290–291, 1812, 1813t
 in kidney disease, 290–291
 in pregnancy, 46
 reduced, 290–291
 regulation of, 332e-2–3, 1800, 1801f
 serum creatinine and, 290–291
 Glomerulonephritis
 acute, 1806t, 1836
 C3, 62e-4f
 chronic, 1836
 classification of, 1836–1837
 endocarditis-associated, 1835t, 1837–1838
 hematuria in, 294, 1834
 membranoproliferative. *See* Membranoproliferative glomerulonephritis
 membranous. *See* Membranous glomerulonephritis
 mesangioproliferative, 1835t, 1841
 pathogenesis of, 1832–1833
 pauci-immune, 1840
 poststreptococcal. *See* Poststreptococcal glomerulonephritis
 progression of, 62e-9–10f, 1834
 rapidly progressive (crescentic), 1836, 1846f
 in relapsing polychondritis, 2204
 renal biopsy in, 62e-1–6f, 1836–1837
 in Sjögren's syndrome, 2167
 in SLE. *See* Lupus nephritis
 tubulointerstitial abnormalities with, 1860–1861
 Glomerulosclerosis
 age-related, 1837
 focal segmental. *See* Focal segmental glomerulosclerosis
 Glomerulus (glomeruli)
 anatomy of, 1812f, 1831–1832, 1832f
 embryologic development of, 332e-1
 filtration by, 333e-1
 histology of, 1813f
 Glossitis, 241t, 463t, 1946t
 Glossodynia (burning mouth syndrome; glossalgia), 215, 239
 Glossopharyngeal nerve, 213, 213f, 2538
 Glossopharyngeal neuralgia, 237, 2649, 2664
 GLP-1. *See* Glucagon-like peptide-1 (GLP-1)
 Glucagon
 for bradycardia due to poisoning/overdose, 473e-9t
 for food impaction dislodgement, 1911
 in hypoglycemia, 2431t
 in shock, 1745
 Glucagon cell adenomatosis, 569
 Glucagon-like peptide-1 (GLP-1), 559t, 572, 2402
 Glucagon-like peptide-1 (GLP-1) receptor agonists, 2414t, 2416
 Glucagonoma, **569**
 clinical features of, 559t, 569, 2338
 diagnosis of, 570
 incidence of, 561
 in MEN 1, 2338
 skin manifestations of, 356t
 treatment of, 570, 2338
 Glucocorticoid(s)
 for ACTH deficiency, 2260
 actions of, 1829t
 for adrenal insufficiency, 2325–2326
 adverse effects of, 598, 1677, 1831, 2130t, 2667
 carcinogenicity, 477t
 cataracts, 204
 cutaneous, 352, 356, 379, 381, 732
 depression, 63
 edema, 253t
 eosinopenia, 423
 hypertrichosis, 380
 late, 620t
 major, 2182t
 in myocardial infarction, 1604
 myopathy, 462e-19–20, 462e-20t, 2199, 2218t
 osteoporosis, 2492, 2503–2504
 in pregnancy, 373
 protection from, 622, 2667
 psychosis, 2129
 for alcoholic liver disease, 2053, 2054f, 2059
 for allergic conjunctivitis, 199
 for anaphylaxis, 2117
 for angioedema, 2119–2120
 for anterior ischemic optic neuropathy, 202
 for autoimmune hemolytic anemia, 658
 for autoimmune hepatitis, 2051–2052
 for autoimmune pancreatitis, 2099
 for Behçet's syndrome, 2194
 for bronchiectasis, 1696
 for CAH, 2328–2329
 for cancer, 103e-15
 for cerebellar degeneration, 618
 for chronic inflammatory demyelinating neuropathy, 2699
 for cluster headache prevention, 2595, 2595t
 for contact dermatitis, 354t
 diabetes mellitus and, 2421–2422
 drug interactions of, 2130t
 for dyspnea, 62
 for ectopic ACTH syndrome, 611
 for eosinophilia, 613
 for eosinophilic granulomatosis with polyangiitis, 1685, 1686, 2187
 epidural, for back pain, 120

- Glucocorticoid(s) (*Cont.*):
 for episcleritis and scleritis, 200
 equipotency, 2326
 for fatigue, 63
 for fever, 126
 for fever of unknown origin, 141
 genomics of response to, 775
 for giant cell arteritis, 2189
 for gout, 2234
 for Graves' ophthalmopathy, 2297
 half-lives of, 400e-2
 for hypercalcemia, 314, 610, 2480t, 2481
 for hyperesinophilic syndromes, 423, 1686
 for IBD, 1960
 for idiopathic cutaneous vasculitis, 2192
 for IgA vasculitis, 2190
 for immune thrombocytopenic purpura, 728
 for immunosuppression, 1828, 1829t
 for inflammatory myopathies, 2200–2201
 inhaled
 actions of, 1677, 1678f
 adverse effects of, 1677, 1705
 for asthma, 1677, 1678f
 for COPD, 1705
 for cough, 245
 intraarticular, for osteoarthritis, 2232, 2232t
 intralesional
 for alopecia areata, 355t
 for mucous membrane pemphigoid, 374
 for pemphigus foliaceus, 372
 intranasal
 adverse effects of, 2122
 for allergic rhinitis, 2122, 2123f
 for sinusitis, 216, 227, 228
 for nausea and vomiting, 62, 260t
 for neck pain, 123
 for optic neuritis, 202
 for orbital pseudotumor, 207
 for otitis externa, 228
 for pain, 61
 for parainfluenza virus infection, 1208
 resistance to, 1680, 2318t
 for rheumatic fever, 2153
 for rheumatoid arthritis, 2145
 for Sweet syndrome, 487
 synthesis of, 2309, 2309f, 2355f
 systemic
 adverse effects of, 1677–1678, 1706, 2130t, 2132, 2182t
 for asthma, 1677–1678
 for atopic dermatitis, 345
 for bullous pemphigoid, 372
 for cerebral edema, 598
 for COPD exacerbation, 1707
 for drug-induced hypersensitivity syndrome, 382
 for drug-induced urticaria/angioedema, 381
 drug interactions of, 2130t
 for Duchenne's muscular dystrophy, 462e-8
 for epidermolysis bullosa acquisita, 374
 for erythema nodosum leprosum, 1128
 for hypercalcemia, 314
 for IgG4-related disease, 391e-3
 for interstitial lung diseases, 1711
 for lichen simplex chronicus, 345
 for lupus nephritis, 1838
 for Ménière's disease, 219
 for minimal change disease, 1842
 for MS, 2667
 for myasthenia gravis, 2705
 for neuromyelitis optica, 2673
 for pemphigoid gestationis, 373
 for pemphigus foliaceus, 372
 for pemphigus vulgaris, 371
 for phototoxicity, 390
 for sarcoidosis, 2210–2211, 2212t
 for serum sickness, 2737
 for sinusitis, 214
 for SJS or TEN, 383
 for SLE, 2130t, 2132
 for subacute cutaneous lupus erythematosus, 375
 for systemic mastocytosis, 2121
 for systemic sclerosis, 2164–2165
 for Takayasu's arteritis, 2190
 for thrombotic thrombocytopenic purpura, 730
 topical
 adverse effects of, 381, 2130t
 for atopic dermatitis, 345
 for bullous pemphigoid, 372
 for candidiasis, 351
 for contact dermatitis, 345
 for dermatitis, 354t
 for dermatomyositis, 375
 for hand eczema, 346
 for herpes simplex keratitis, 200
 for lichen planus, 349
 for lichen simplex chronicus, 345
 for morbilliform drug eruption, 381
 for mucous membrane pemphigoid, 374
 for pemphigoid gestationis, 373
 for pemphigus foliaceus, 372
 for phototoxicity, 390
 for pityriasis rosea, 349
 for psoriasis, 348, 354t
 for seborrheic dermatitis, 347
 for SLE, 2130t
 for stasis dermatitis, 346
 for sulfur mustard exposure, 262e-4
 for uveitis, 200
 for vitiligo, 358t
 for tuberculous meningitis, 775
 for urticaria, 2119–2120
 for vasculitis, 2181–2182
 withdrawal from, 103e-15, 378
 Glucocorticoid receptor gene mutations, 2356t
 Glucocorticoid-remediable aldosteronism (GRA;
 familial hyperaldosteronism type I)
 clinical features of, 1619, 1620t
 complications of, 2319
 diagnosis of, 307, 1619
 genetic considerations in, 307, 434, 1620t, 2318–2319, 2318t
 treatment of, 1619, 2320
 Gluconogenesis, 95e-1, 461, 2430
 Glucosamine, 2232
 Glucose
 in ascitic fluid, 287–288
 balance and counterregulation of, 2430, 2431f
 for hyperkalemia, 312
 metabolism in cancer cells, 102e-12
 plasma levels of, 2401, 2401f, 2401t, 2760t
 regulation of, 2399f, 2401, 2402–2403, 2402f
 renal transport of, 332e-3f, 332e-8
 in specialized nutritional support formulas, 98e-4–5, 98e-7
 Glucose-6-phosphatase deficiency (type Ia GSD;
 Von Gierke's disease), 375e-1, 433e-1, 433e-2t, 433e-3
 Glucose-6-phosphate dehydrogenase (G6PD)
 deficiency, 654
 clinical features of, 393, 655–656
 diagnosis of, 656, 656f, 2755t
 drug contraindications in, 656, 656t
 dapson, 1126, 1262
 furazolidone, 246e-9
 primaquine, 246e-12, 1378t, 1380t, 1381, 1384
 rasburicase, 1795
 drug responses in, 39t, 40
 early interventions for, 450t
 epidemiology of, 393, 655, 655f
 genetic considerations in, 654t, 655
 genetic testing for, 450t
 global considerations in, 655, 655f
 jaundice in, 281
 malaria resistance and, 656, 1371
 pathophysiology of, 650f, 654–655, 654f
 red cell enzyme abnormalities in, 654t
 Rocky Mountain spotted fever in, 1156
 treatment of, 656–657
 Glucose-6-phosphate isomerase, 372e-24t
 Glucose-6-phosphate isomerase (GPI)
 deficiency, 650f, 654t
 Glucose–galactose, malabsorption of, 435e-1t
 Glucose tolerance, 2401, 2401t, 2451, 2454
 Glucose-transport defect, 435e-1t
 Glucose transporter 1 (GLUT-1), 225e-3, 247
 Glucose transporter 2 (GLUT-2), 247
 Glucose transporter 2 (GLUT-2) deficiency
 (Fanconi-Bickel syndrome; type XI
 GSD), 332e-7t, 433e-2t, 435e-1t, 2434–2435
 Glucose transporter 9 (GLUT9), 431e-1, 431e-2f, 431e-3
 α-Glucosidase inhibitors, 2414t, 2415, 2416
 Glue sniffer's neuropathy, 2688t, 2689
 GLUR (glutamate receptor), 372e-24t
 GLUT9 (glucose transporter 9), 431e-1, 431e-2f, 431e-3
 Glutamate, 88, 444e-3t
 Glutamate decarboxylase (GAD65), 372e-24t
 Glutamate receptor (GLUR), 372e-24t
 Glutamine metabolism disorders, 434e-3t
 Glutamine synthase deficiency, 434e-3t
 γ-Glutamylcarboxylase, 736, 736f
 Glutaric acidemia, 434e-3t
 Glutathione peroxidase deficiency, 657
 Glutathione-S-transferase proteins, 280
 Glutathione synthase deficiency, 654t
 Gluten-free diet
 for ataxic disorders, 2630
 for celiac disease, 1942
 for dermatitis herpetiformis, 373
 Gluten-induced enteropathy. *See* Celiac disease (gluten-induced enteropathy)
 Gluten intolerance, nonceliac, 269
 Glutethimide, 41, 42t, 473e-11t
 Glyburide, 48, 727t, 2414t
 Glycan shield, 1244
 Glycemic control
 in critically ill patient, 1734
 in diabetes mellitus, 2409–2410
 in diabetic nephropathy, 2425–2426
 diabetic retinopathy and, 2423f
 long-term, assessment of, 2410
 in metabolic syndrome, 291e-7, 2454
 perioperative, 55
 in pregnancy, 48
 in sepsis/septic shock, 1758
 Glycereraldehyde-3-phosphate dehydrogenase (GAPD) deficiency, 654t
 Glycine, 444e-3t
 Glycine encephalopathy, 434e-2t
 Glycine metabolism disorders, 434e-2t
 Glycine N-methyltransferase deficiency, 434e-2t
 Glycine receptor antibodies, 614t, 617
 Glycogen, 433e-1
 Glycogenesis Ib disease, 375e-1
 Glycogenolysis, myocardial, 265e-10
 Glycogen storage diseases (GSD), **433e-1**
 clinical features of, 433e-2–3t, 2198, 2434–2435
 global considerations in, 433e-5
 with hepatomegaly and hypoglycemia
 type 0, 433e-2t
 type Ia (von Gierke's disease), 432e-2t, 433e-1, 433e-3, 436e-5f
 type Ib, 432e-2t, 433e-3
 type III, 432e-2t, 433e-3, 436e-6f, 462e-14, 1565
 type VI (Hers disease), 433e-2t, 433e-4
 type IX, 332e-7t, 432e-2t, 433e-4, 462e-14
 type X, 462e-14
 type XI (Fanconi-Bickel syndrome), 332e-7t, 433e-2t, 433e-4, 462e-14
 hyperuricemia in, 431e-2
 IBD and, 1949t
 late-onset, 432e-4t
 with liver cirrhosis (type IV), 433e-2t, 462e-14
 with muscle-energy impairment
 β-enolase deficiency, 432e-2t, 462e-14
 lactate dehydrogenase deficiency, 432e-2t, 462e-14
 pyruvate kinase deficiency, 281, 433e-2t, 650f, 653–654, 654t
 type V (McCardle's disease), 433e-2t, 433e-4, 462e-14
 type VII (Tarui's disease), 433e-2t, 462e-14, 654t
 type X, 433e-2t, 462e-14

- pathogenesis of, 433e-1, 433e-1f
with progressive skeletal myopathy and/or cardiomyopathy
 Danon's disease, 432e-2t, 432e-4t, 433e-5, 1566
 PRKAG2 deficiency, 433e-2t, 433e-5
 type II. *See* Pompe disease (acid α -1,4 glucosidase deficiency)
- Glycolysis
 abnormalities of, 462e-14, 650f, 653–654
 anaerobic bacterial, 461
 in heart muscle, 265e-10
- Glycopeptides, 930, 931t, 942
- Glycoprotein gp120, 214e-3
- Glycoprotein hormones, 400e-1
- Glycoprotein Ib/IX, 372e-25t
- Glycoprotein Ib-IX-V complex, 275–276, 741–742
- Glycoprotein IIb/IIIa, 401, 745, 746f, 748
- Glycoprotein IIb/IIIa antagonists. *See also specific drugs*
 adverse effects of, 749
 discontinuation before lumbar puncture, 443e-1
 dosage of, 749
 indications for, 749
 for myocardial infarction, 1596, 1606
 in PCI, 296e-1
 pharmacology of, 749t
- Glycoprotein IIb/IIIa receptors, 276, 725, 742
- Glycoprotein IIb/IIIg, 372e-25t
- Glycoproteinosis(es), 432e-3t
- Glycopyrrolate, 2643
- Glycosphingolipidoses, neutral, 432e-3t
- Glycosuria, 299, 332e-7t
- Glycylcyclines. *See* Tigecycline
- Glycyrrhizic acid, 64e-8, 306, 307
- GMAP (Global Malaria Action Plan), 14
- GM-CSF (granulocyte-macrophage colony-stimulating factor), 372e-11t, 414f, 416–417
- GM-CSF therapy. *See* Granulocyte-macrophage colony-stimulating factor (GM-CSF) therapy
- GM gangliosides, 372e-24t
- GNA11* gene mutations, 2475
- GNAS1* gene mutations, 426e-6, 430, 437, 2344
- GNAS* gene mutations, 2315, 2486
- Gnathostoma spinigerum*/gnathostomiasis
 clinical features of, 908t, 1412–1413
 diagnosis of, 245e-6t, 765t, 1413
 epidemiology of, 765t, 1412
 treatment of, 1411t, 1413
- Gnotobiotic, 86e-1t, 86e-7
- GnRH. *See* Gonadotropin-releasing hormone (GnRH)
- GnRH1* gene mutations, 2363
- GnRHR* gene mutations, 2363, 2364t
- GOBI, 8
- GOBI-FFF, 8
- Goblet cells, 86e-8
- Goiter, **2301**
- Goitrogens, 2301
- Goitrous thyroiditis. *See* Hashimoto's thyroiditis (autoimmune hypothyroidism)
- Gold, adverse effects of, 349, 361, 379, 727t, 2193, 2688t
- Golden Rice, 96e-7
- Golgin, 372e-24t
- Golimumab
 adverse effects of, 348t, 2146t, 2173
 for ankylosing spondylitis, 2172–2173
 for autoimmune and inflammatory diseases, 372e-30
 for IBD, 1961
 for psoriasis/psoriatic arthritis, 348t, 2177
 for rheumatoid arthritis, 2145, 2146t
- Gompertzian growth curve/kinetics, 102e-2, 103e-3f
- Gonadal development, 2349f, 2351f
- Gonadal dysgenesis, 2353
- Gonadal mosaicism, 433
- Gonadal sex, 2349, 2349f
- Gonadectomy, 336
- Gonadotropin(s), **401e-4**. *See also* Follicle-stimulating hormone (FSH); Luteinizing hormone (LH)
 action of, 401e-4
 deficiency of. *See* Hypogonadism
 in male reproductive function, 2360
 for ovulatory induction, 2389
 pituitary tumor production of, 2273–2274
 secretion of, 401e-4
 synthesis of, 401e-4
- Gonadotropin-releasing hormone (GnRH)
 action of, 401e-2f, 401e-4
 deficiency of. *See* Hypogonadotropic hypogonadism
 for hypogonadotropic hypogonadism, 337
 hypothalamic, 401e-1, 401e-2f
 in male pubertal development, 2357
 in menstrual cycle regulation, 2378
 in ovarian function regulation, 2376–2377, 2377f
 pulse frequency of, 400e-6, 401e-4
- Gonadotropin-releasing hormone (GnRH) analogues, 2362
- Gonadotropin-releasing hormone (GnRH) stimulation test, 2360
- Gonioscopy, 200
- Gonococcal arthritis. *See* Gonococcal infections, disseminated
- Gonococcal infections, **1003**
 anorectal, 1005–1006
 cephalosporin intermediate/resistant, 1005
 in children, 1006, 1008
 diagnosis of, 150e-6, 150e-8, 1007–1008
 disseminated
 clinical features of, 133t, 835, 1006, 1007f
 complications of, 1006
 epidemiology of, 133t, 1006
 joint involvement in, 1006, 1007t
 oral manifestations of, 238t
 risk factors for, 1006
 skin manifestations of, 25e-11f, 76e-13f, 133t, 135, 369, 1007f
 treatment of, 1008, 1009t
 epidemiology of, 869, 1003
 epididymitis, 872
 global considerations in, 1003, 1004
 in HIV infection, 1006–1007
 in men, 870–871, 881, 1005, 1005f
 in neonates, 1006, 1009t
 ocular, 1006, 1009t
 oral manifestations of, 238t
 pathogenesis of, 1003–1004. *See also* *Neisseria gonorrhoeae*
 pelvic inflammatory disease. *See* Pelvic inflammatory disease (PID)
 pharyngeal, 231, 1006, 1008, 1009t
 in pregnancy, 50, 1006, 1008
 prevention and control of, 1009
 quinolone-resistant, 1004
 screening for, 29t, 882
 spectinomycin-resistant, 1004–1005
 treatment of, 871, 872t, 1008, 1009t
 in women, 872, 1005
- Go-no go test, 183
- Gonorrhea. *See* Gonococcal infections
- Goodpasture's syndrome
 clinical features of, 377e-4, 1715, 1835t, 1836, 1839
 diagnosis of, 1839
 hemoptysis in, 245
 HLA gene association with, 373e-8t
 pathophysiology of, 372e-24t, 377e-4, 1715, 1833
 prognosis of, 1839
 treatment of, 1715, 1839
- Good's syndrome, 123e-1
- Gordon's syndrome. *See* Pseudohypaldosteronism type II (Gordon's syndrome)
- Gorlin formula, valve area measurement, 1463
- Gorlin's syndrome (nevroid basal cell carcinoma syndrome)
 clinical features of, 364t, 365, 599t
 genetic considerations in, 101e-6t, 599t, 602
- nonmelanoma skin cancer in, 500
 ovarian fibroma in, 594
- Goserelin, 103e-15, 585
- Gottron's sign, 374, 374f, 2195
- Gout, **2233**
 clinical features of, 2233
 diagnosis of, 2233–2234, 2234f
 drug-induced, 2218t
 genetic considerations in, 431e-3, 431e-3t
 hyperuricemia in, 431e-4
 inflammasome activity in, 372e-8t
 nephropathy in, 431e-4, 1862
 obesity and, 415e-6
 prevalence of, 431e-1, 431e-4
 tophaceous, 365, 436e-2f
 treatment of, 2234–2235
- Gou virus, 1307t
- Gowers' sign, 462e-1, 462e-3f
- gp41, 1230, 1244
- gp120, 1216, 1230, 1237
- gp160, 1237
- GPC regimen, for bladder cancer, 577
- GPCRs (G protein-coupled receptors), 212, 400e-1, 400e-1t, 2252
- GPI (glucose-6-phosphate isomerase) deficiency, 650f, 654t
- GPIHP1 deficiency, 2440, 2440t
- G protein(s), 400e-3, 400e-3f, 468e-1, 1613
- G protein-coupled receptors (GPCRs), 212, 400e-1, 400e-1t, 2252
- GRA. *See* Glucocorticoid-remediable aldosteronism (GRA)
- Graafian follicle, 2376f
- Graded exercise therapy, 153, 464e-2–3
- Gradenigo's syndrome, 209, 906
- Graft-versus-host disease (GVHD)
 chronic, 139e-4
 clinical features of, 139e-3
 CMV infection and, 921–922
 cobalamin malabsorption in, 646
 diarrhea in, 267, 270
 grading and staging of, 139e-3–4, 139e-3t
 incidence of, 139e-1
 prevention of, 138e-5, 139e-4
 skin manifestations of, 139e-3, 349, 353, 361, 362, 362t
 transfusion-associated, 138e-5, 664
 treatment of, 139e-4
- Graft-versus-tumor effect, 139e-1
- Graham Steell murmur, 51e-6, 1540, 1550
- Grain dust exposure, 1688t, 1691
- Gramicidin, 945t
- Grammar, assessment of, 177
- Gram-negative bacteria. *See also specific bacteria*
 antibiotic resistance in, 807, 933, 1026–1027
 antibiotic targets in, 932f
 diagnosis of, 150e-1, 150e-2t, 1026
 epidemiology of, 1025
 global considerations in, 1026–1027
 host interactions of, 1025, 1025t
 infections caused by, 1026, 1752t
 in sepsis/septic shock, 1752t
 structure and function of, 1025
- Gram-positive bacteria. *See also specific bacteria*
 antibiotic resistance in, 933
 antibiotic targets in, 932f
 diagnosis of, 150e-1, 150e-2t
 in sepsis/septic shock, 1752t
- Gram's stain, **150e-1**, 150e-2t, 806, 1085f
- Grandmother hypothesis, of aging, 94e-3
- Granisetron, 62, 103e-25, 109, 260, 260t
- Granular casts, 62e-11f
- Granular lymphocytes, 81e-2
- Granulicatella* spp., 821, 971
- Granulocyte(s), 81e-6f, 89e-3f, 419t. *See also* Eosinophil(s); Neutrophil(s)
- Granulocyte colony-stimulating factor (G-CSF), 372e-11t, 414f, 804
- Granulocyte colony-stimulating factor (G-CSF) therapy
 adverse effects of, 366, 379
 for AML, 686
 for aplastic anemia, 667
 in HSCT, 89e-1, 139e-1–2

- Granulocyte colony-stimulating factor (G-CSF) therapy (*Cont.*):
 indications for, 103e-23–24, 103e-24t
 for myelodysplasia, 671
 for neutropenia, 423
 for sulfur mustard exposure, 262e-4
- Granulocyte-macrophage colony-stimulating factor (GM-CSF), 372e-11t, 414f, 416–417
- Granulocyte-macrophage colony-stimulating factor (GM-CSF) therapy
 for AML, 686
 in HSCT, 139e-1–2
 indications for, 103e-23–24, 103e-24t
- Granulocytic sarcoma (cutaneous chloroma), 365
- Granulocytopenia, in cancer patient, 486, 486t, 488
- Granulocytosis, 612, 612t, 634
- Granuloma(s)
Candida, 1343
 diagnosis of, 342
 formation of, 145e-10
 hepatic, 367e-3, 2026t
 palisading, 364
 periapical, 236
 pyogenic, 237
 in sarcoidosis, 2205. *See also* Sarcoidosis
 in schistosomiasis, 1426
 tick-bite, 2745
 in vasculitis, 2180–2181
- Granuloma annulare, 355, 356t, 2430
- Granuloma inguinale. *See* Donovanosis
- Granulomatosis
 bronchocentric, 1716
 eosinophilic, with allergic angitis. *See* Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
 lymphomatoid, 135e-3, 1716, 2184
- Granulomatosis infantiseptica, 983
- Granulomatosis with polyangiitis (Wegener's), 2182
 clinical features of, 1840, 2183–2184, 2183t
 bacterial pneumonia, 386e-2f
 bilateral nodular infiltrates, 386e-1f
 chronic meningitis, 909t
 cutaneous, 366, 2184
 hemoptysis, 245
 intracranial infarcts, 2568–2569
 ocular, 200, 386e-2f, 2184
 oral, 237
 renal, 1835t, 1840, 2184
 saddle-nose deformity, 1443, 2183
 sinusitis, 226, 386e-3f
 definition of, 2182
 diagnosis of, 2184
 ANCA, 2184
 chest imaging, 386e-1, 386e-1f
 CT, 386e-1f, 386e-2f, 386e-3f, 2183f
 lung biopsy, 386e-1, 386e-5f
 lung histology, 2182, 2183f
 renal biopsy, 62e-5f, 386e-1, 386e-6f, 1840
 differential diagnosis of, 2184, 2204
 incidence and prevalence of, 2182
 pathogenesis of, 377e-4, 2180, 2182–2183
 pathology of, 2182–2183, 2183f
 treatment of
 cyclophosphamide induction, 2185
 methotrexate induction, 2185
 organ-specific, 2185–2186
 principles of, 2181–2182, 2184–2185
 remission maintenance, 2185
 rituximab induction, 2185
 toxicities of, 2182t
 trimethoprim-sulfamethoxazole, 2185
- Granulomatous (necrotizing) arteritis, 2567–2568
- Granulomatous diseases, chronic. *See* Chronic granulomatous diseases (CGDs)
- Granulomatous gastritis, 1932
- Granulomatous interstitial nephritis, 1857f, 1858
- Granulomatous myocarditis, 1561
- Granulomatous (subacute) thyroiditis, 2298–2299, 2298f
- Granulosa cell(s), 2375
- Granulosa cell tumor, 594
- Grapefruit juice, 42, 246e-9, 1590
- Graphesthesia, 160, 2539
- Grasp reflex, 2539
- Graves' disease, 2293. *See also* Thyrotoxicosis
 anti-*Yersinia* antibodies in, 1077
 APS-2 and, 2346, 2347t
 in children, 2297
 clinical course of, 2296
 clinical features of, 2294
 goiter, 2301
 ophthalmopathy. *See* Graves' ophthalmopathy
 pretibial myxedema, 76e-16f
 signs and symptoms, 2294t
 thyroid acropachy, 2245, 2295f
 thyroid dermopathy, 2295f
 diagnosis of, 2254t, 2288, 2295, 2295f
 differential diagnosis of, 2295–2296
 edema in, 253
 epidemiology of, 2293
 genetic considerations in, 2294
 insulin autoimmune syndrome in, 2348
 pathogenesis of, 372e-24t, 2251, 2294
 in pregnancy, 2297
 prevalence of, 2254t
 radiation therapy-related, 623
 remission rates, 2296
 thymic tumors in, 2348
 treatment of, 2296–2297
- Graves' ophthalmopathy
 clinical course of, 2296
 clinical features of, 41e-1v, 207, 2294–2295, 2295f
 diagnosis of, 207
 pathogenesis of, 462e-18, 2294
 treatment of, 207, 2297
- Gray (Gy), 103e-4, 263e-1
- Gray hepatization, 804
- Great saphenous vein, ligation of, 1652–1653
- Green bottle fly, 2747
- Greenhouse gases, 151e-1, 151e-2f, 151e-2t
- Green nail syndrome, 1047
- Green tea extracts, 351
- GRFoma (growth hormone–releasing factor tumor), 559t, 571
- Griffiths' point, 1979, 1979f
- Griselli syndrome, 375e-2
- Griseofulvin, 350, 350t, 355t, 1331
- GRN gene mutations, 172t, 2603
- Gross cystic disease fibrous protein, 120e-2, 120e-2t
- Ground-glass opacity
 in ARDS, 308e-10f
 bleomycin-related, 621
 in granulomatosis with polyangiitis, 386e-2f
 pulmonary, 515–516, 517f, 1689
- Group A *Streptococcus* (GAS) infections, 963.
See also *Streptococcus pyogenes*
 asymptomatic carrier of, 966
 in cancer patient, 486
 cellulitis, 830, 967–968
 complications of. *See* Poststreptococcal glomerulonephritis; Rheumatic fever
 diagnosis of, 150e-2, 150e-6, 232–233
 ecthyma, 349
 endocarditis. *See* Infective endocarditis
 epidemiology of, 963–964
 epiglottitis, 234
 erysipelas. *See* Erysipelas
 global considerations in, 964, 964f, 966
 health care–associated, 913
 impetigo. *See* Impetigo
 infection control measures, 917
 joints, 834. *See also* Infectious (septic) arthritis
 laryngitis, 234
 myositis, 831, 968
 necrotizing fasciitis. *See* Necrotizing fasciitis
 pathogenesis of, 964–965
 pharyngitis. *See* Pharyngitis, streptococcal
 pneumonia, 968
 prevention of, 969
- puerperal, 968
- pyomyositis, 831
- rheumatic fever and, 963–964, 2152. *See also* Rheumatic fever
 scarlet fever. *See* Scarlet fever
 surgical-site, 915, 968
 toxic shock syndrome. *See* Toxic shock syndrome (TSS), streptococcal
 treatment of, 965–966, 965t
- Group B *Streptococcus* (GBS) infections
 cellulitis, 830, 970
 clinical syndromes, 970
 diagnosis of, 150e-8, 969
 endocarditis. *See* Infective endocarditis
 epidemiology of, 969
 myocarditis, 992
 neonatal, 970
 pharyngitis, 231
 in pregnancy, 50, 970
 prevention of, 970
 treatment of, 50, 970
- Group C *Streptococcus* infections, 969
- Group D *Streptococcus* infections, 486t, 490, 539, 971
- Group G *Streptococcus* infections, 231, 969
- Growth
 disorders of, 2257–2258. *See also* Short stature
 hormones in, 400e-5. *See also* Growth hormone (GH)
 somatic, skeletal maturation and, 2257
 in type Ia glycogen storage disease, 436e-5f
- Growth hormone (GH), 401e-3
 action of, 401e-3
 Creutzfeldt-Jakob disease after, 453e-3
 ectopic production of, 609t
 effectors and signaling pathways of, 400e-1t
 expression and regulation of, 401e-1t
 insensitivity, 2258
 laboratory evaluation of, 2258t
 in obesity, 415e-4
 for osteoporosis management/prevention, 2503
 in puberty, 2379
 recombinant
 for adult growth hormone deficiency, 2260, 2260f
 adverse effects of, 2260
 for growth hormone deficiency in children, 2259
 secretion of, 401e-3
 synthesis of, 401e-3
- Growth hormone (GH) deficiency
 in adults, 2259–2260, 2259t, 2260f
 in children, 2257–2258, 2259t, 2433
 idiopathic, 2258
 laboratory investigation of, 2259
- Growth hormone (GH) gene mutations, 434
- Growth hormone (GH) receptor antagonists, for acromegaly, 2271
- Growth hormone–releasing factor tumors (GRFomas), 559t, 571
- Growth hormone-releasing hormone (GHRH), 401e-3, 609t, 2258, 2265
- Growth hormone-releasing hormone (GHRH) agonists, 327
- GRO- α /MGSA, 372e-12t
- GRO- β /MIP-2 α , 372e-12t
- GSD. *See* Glycogen storage diseases (GSD)
- GSS (Gerstmann-Sträussler-Scheinker) disease, 453e-1t, 453e-4
- GSTP1* gene promoter, 580
- GTP cyclohydrolase I deficiency, 434e-2t
- Guama virus, 1309t
- Guanabenz, 473e-9t
- Guanarito virus, 1306t
- Guanethidine, 253t
- Guanfacine, 468e-3, 1624t, 2623
- Guanine, 426
- Guided imagery, 14e-1t
- Guillain-Barré syndrome (GBS), 2694
 after infectious diarrhea, 855t
 antecedent events, 2695
 vs. botulism, 989

- vs. chronic inflammatory demyelinating polyneuropathy, 2699
- clinical features of, 2694–2695, 2698t
 autonomic dysfunction, 2642, 2643, 2695–2696
 cranial nerve palsies, 2650
 dyspnea, 47e-2
 facial nerve disorders, 2648
 weakness, 156, 157, 2694
- in CMV mononucleosis, 1192
- diagnosis of, 2695t, 2697
- imaging in, 441e-42f
- immunopathogenesis of, 372e-24t, 2695, 2696f, 2696t, 2697
- in influenza, 1212–1213
- influenza vaccine and, 1214, 2695
- Miller Fisher variant, 989
- paraneoplastic, 614t, 618
- pathophysiology of, 2697
- prevalence of, 2694
- prognosis of, 2698
- subtypes of, 2695, 2695t
- treatment of, 2697–2698
- Guinea worm infection. *See* *Dracunculus medinensis* (guinea worm)/dracunculiasis
- Gulf War syndrome, 152e-8–9
- Gumboil, 236
- Gummas, 1134, 1136
- Gums, strawberry, 237
- Gut-associated lymphoid tissue (GALT), 1228, 1232
- Gut microbiota
 bone mass and, 86e-7
 brain function and, 86e-7–8
 cardiovascular system and, 86e-7
 diabetes and, 86e-8–9
 in IBD, 1948f, 1949
 immune function and, 86e-8
 obesity and, 86e-8
- Guttate psoriasis, 347. *See also* Psoriasis
- GVHD. *See* Graft-versus-host disease (GVHD)
- GWAS. *See* Genome-wide association studies (GWAS)
- Gy (Gray), 103e-4, 263e-1
- Gynecomastia, **2366**
 breast cancer risk and, 532
 cocaine use and, 469e-1
 definition of, 2366
 diagnosis of, 2254t, 2366, 2367f
 etiology of, 2366
 in Klinefelter's syndrome, 2350
 in liver disease, 286
 obesity and, 415e-6
 prevalence of, 2254t, 2366
 prevention of, in androgen deprivation therapy, 7e-6
 in testicular cancer, 588
 treatment of, 2366–2367
- Gyrate atrophy, choroid and retina, 434e-3t
- H**
- H₁ receptor antagonists. *See* Antihistamines
- H₂ receptor antagonists. *See* Histamine H₂ receptor antagonists
- H6PDH (hexose-6-phosphate dehydrogenase), 2312
- HAART (highly active antiretroviral therapy). *See* Combination antiretroviral therapy (cART)
- HACE. *See* High-altitude cerebral edema (HACE)
- HACEK group, 183e-1
- HACEK group infections, 183e-1, 183e-2t. *See also* Infective endocarditis, HACEK group
- HAD (HIV-associated dementia), 1263, 1264t, 1265f
- Haemophilus ducreyi*, 1012. *See also* Chancroid
- Haemophilus influenzae*
 antibiotic resistance in, 1012
 host response to, 1010
 laboratory identification of, 150e-2t, 1011
 microbiology of, 1010
 nontypable strains, 1010, 1010t
 type b, 1010, 1010t
- Haemophilus influenzae* infections, **1010**
 in cancer patient, 484, 484t, 486t, 488
 chemoprophylaxis for, 1012
 clinical features of, 1010t, 1011
 cellulitis, 830, 1011
 epiglottitis, 234–235, 1011
 hemoptysis, 245
 mastoiditis, 231
 meningitis, 1011. *See also* Meningitis
 otitis media, 229–230, 1011
 pneumonia, 915, 1011
 sinusitis, 226
- in cystic fibrosis, 1697
 diagnosis of, 1011
 epidemiology of, 1010, 1010f
 global considerations in, 1010, 1012
 in human bite-wound, 167e-2
 incidence pre- and postvaccine, U.S., 785t
 pathogenesis of, 1010
 pneumonia, 1212
 in postsplenectomy sepsis, 413, 781
 prevention of. *See* *Haemophilus influenzae* vaccine
 in transplant recipient, 920t
 treatment of, 230, 940t, 1011–1012
- Haemophilus influenzae* vaccine
 global considerations in, 1012
 recommended administration of
 in cancer patient, 485t, 492
 in children, 1012
 in splenectomized patient, 413, 635
 in transplant recipient, 929, 929t
- Haemophilus parainfluenzae*, 183e-1, 183e-2t, 234. *See also* Infective endocarditis, HACEK group
- Haemophilus* vaginitis. *See* Bacterial vaginosis
- Hair
 drug-induced disorders of, 103e-25, 354, 379–380
 growth cycle of, 331–332
 loss of. *See* Alopecia
 in nutritional deficiencies, 463t
 removal of, 334
 terminal, 331
 vellus, 331
- Hair follicle stem cells, 453t
- Hairy cell leukemia
 clinical features of, 135e-1, 706, 2247
 infections in, 484t, 486t, 706
 monocytopenia in, 422
 skin manifestations of, 369, 2187, 2193
 splenomegaly in, 411
 treatment of, 135e-2, 412, 706
 tumor markers in, 135e-1–2, 135e-2t, 473t
- Hakki formula, aortic valve area measurement, 1463
- Half-life
 circulating hormone, 400e-2
 of drugs, 34, 35f, 935
- Half-normal saline, 298. *See also* Saline
- Halide exposure, 1692t
- Halitophobia, 226
- Halitosis, 226, 233, 242
- Hallervorden-Spatz disease. *See* Pantothenate kinase-associated neurodegeneration (PKAN)
- Hallucinations
 chemosensory, 215
 hypnagogic, 189
 hypnopompic, 189
 in PD, 2617
 in respiratory acidosis, 322
- Hallucinoses, peduncular, 2575
- Halocarbons, 151e-2t
- Halofantrine
 action of, 246e-9
 adverse effects of, 246e-2t, 1380t
 drug interactions of, 246e-2t, 246e-5t, 246e-10, 798
 for malaria, 1380t
 pharmacology of, 246e-9, 1380t
 in pregnancy and lactation, 246e-2t
- Haloperidol
 for acute alcohol intoxication, 2727
 adverse effects of, 2721, 2722t
 for delirium, 64, 64t, 79, 1782
 drug interactions of, 42
 for nausea and vomiting, 61, 103e-25
 overdosage/poisoning with, 473e-9t
 for PCP-induced psychosis, 469e-4
 for schizophrenia, 2722t
- Halo sign, 1348
- Halzoun, 2747
- HAM (HTLV-1-associated myelopathy), 163, 225e-5, 697, 707, 2657
- Hamartin, 562, 604, 1854
- Hamartoma
 colonic, 538t
 hypothalamic, 2264
 iris, 2343
 pulmonary, 515
 small intestinal, 536
- Hamman-Rich syndrome (acute interstitial pneumonia), 1712
- Hamman's sign, 1720
- Hampton's hump, 101, 1633
- HAND (HIV-associated neurocognitive disorders), 1241, 1263, 1264t, 1265f
- Hand(s)
 cryoglobulinemia in, 386e-5f
 eczema of, 342f, 346, 346f
 injury, bite-wounds, 167e-1–2. *See also* Bite-wounds
 mechanic's, 2195
 osteoarthritis of, 2226, 2226f
 pain in, 2220–2221, 2220f
- Hand-foot-and-mouth disease
 epidemiology of, 1292–1293
 global considerations in, 1292–1293
 oral manifestations of, 25e-6f, 76e-8f, 238t, 1292, 1292f
 skin manifestations of, 130t, 134, 361t, 1292, 1292f
- Hand-foot syndrome, 103e-26, 634, 2242
- Hand hygiene, 86e-7, 913, 916
- Hand-Schüller-Christian disease, 2264
- Hansen's disease. *See* Leprosy
- Hantaan virus, 1307t
- Hantaviruses, 1307–1308t
- Hantavirus infections
 epidemiology of, 151e-8f, 784, 1319
 hemorrhagic fever with renal syndrome, 784, 1321
 pulmonary syndrome, 784, 1319–1320
- HAPE. *See* High-altitude pulmonary edema (HAPE)
- Haploid, 426
- Haploinsufficiency, 434
- Haplotype, 373e-2, 427, 429f, 435
- Haplotype block, 429f
- HapMap project, 425, 426t, 427, 435, 442
- Haptocorrins, 640
- Hard (ixodid) tick, 2745. *See also* Tick(s)
- Harris-Benedict equations, 462
- Hartmann's procedure, 1973, 1973f
- Hartnup's disease/syndrome, 96e-4, 435e-1t, 435e-2, 1937
- Harvester ants, 2749
- Hashimoto's encephalopathy, 441e-29f, 453e-5
- Hashimoto's thyroiditis (autoimmune hypothyroidism)
 after drug-induced hypersensitivity syndrome, 382
 clinical features of, 2290t, 2291, 2291f, 2299, 2301
 diagnosis of, 2291, 2292f
 differential diagnosis of, 2291
 pathogenesis of, 372e-24t, 377e-4–5, 2290
 prevalence of, 2290
 thyroid lymphoma in, 2307
 treatment of, 2292–2293
- Hashish, 469e-2. *See also* Marijuana
- Hastisetæ, 2750
- HAV (hepatitis A virus), 2004–2005, 2005f, 2006t. *See also* Hepatitis A virus (HAV) infection

- Haverhill fever (rat-bite fever), 130t, 167e-1-2
 Haversian systems, 2455, 2456
 HAV infections. *See* Hepatitis A virus (HAV) infection
 Hawkinsinuria, 434e-2t
 Haw River syndrome, 2629
 HAX-1 gene, 375e-1
 Hayflick limit, 94e-3
 HAZMATs, 262e-1
 HbA_{1c}. *See* Hemoglobin A_{1c} (HbA_{1c})
 HBAT (heptavalent botulinum antitoxin), 261e-9
 HBcAg (hepatitis B core antigen), 2006, 2006t
 HBeAg (hepatitis B e antigen), 2006, 2006t
 HBsAg (hepatitis B surface antigen), 2005-2006, 2006t
 HBV. *See* Hepatitis B virus (HBV)
 HBV infection. *See* Hepatitis B virus (HBV) infection
 HBxAG (hepatitis B x antigen), 2005f, 2007
 HCC. *See* Hepatocellular carcinoma (HCC)
 hCG. *See* Human chorionic gonadotropin (hCG)
 HCOM. *See* Hypertrophic cardiomyopathy
 HCP. *See* Hereditary coproporphyrin (HCP)
 HCP5 gene, HIV infection and, 1240t, 1241
 HCV (hepatitis C virus), 2005f, 2006t, 2009-2010, 2009f, 2010f. *See also* Hepatitis C virus (HCV) infection
 HDAC. *See* Histone deacetylase (HDAC)
 HDL. *See* High-density lipoprotein (HDL)
 HDR (hypoparathyroidism, deafness, and renal dysplasia) syndrome, 2483
 HDV (hepatitis D virus), 2005f, 2006t, 2008-2009. *See also* Hepatitis D virus (HDV) infection
 Headache, **107, 2586**
 acute, new-onset, 107-108, 108t, 1785
 in acute mountain sickness, 476e-2
 after head injury, 457e-4
 anatomy and physiology of, 107
 with autonomic symptoms, 107
 in brain abscess, 901
 in brain tumor, 107t, 108, 598
 chronic daily, **109**
 approach to the patient, 109-110
 classification of, 109, 109t
 management of, 109
 medication-overuse, 109, 2593
 cluster. *See* Cluster headache
 cold-stimulus, 2597
 depression and, 108
 external pressure, 2597
 in giant cell arteritis, 108
 in glaucoma, 108
 hemicrania continua, 109t, 2596
 in hyperperfusion syndrome, 463e-1
 hypertensive, 1621
 hypnic, 2598
 medication-overuse headache, **109**
 in meningitis, 108
 migraine. *See* Migraine
 new daily persistent, **109**
 clinical features of, 110, 2598
 differential diagnosis of, 109t
 low CSF volume headache, 110, 110f
 posttraumatic, 107t, 110-111
 raised CSF pressure headache, 110
 treatment of, 111, 2598
 nummular, 2598
 in papilledema, 203
 paroxysmal hemicrania, 109t, 2595t, 2596
 post-lumbar puncture, 443e-3, 443e-3f, 443e-3t
 in pregnancy, 49
 primary, 107, 107t, 108, 2587t
 primary care management of, 111
 primary cough, 2597-2598
 primary exertional, 2597
 primary sex, 2597
 primary stabbing, 2597-2598
 primary thunderclap, 2597
 in respiratory acidosis, 322
 secondary, 107, 107t, 108
 in SLE, 2129, 2129t
 in subarachnoid hemorrhage, 107t, 108, 1785
 SUNCT/SUNA, 109t, 2595t, 2596
 symptoms of serious underlying disorder, 108t
 tension-type, 107t, 2587t, 2593-2594
 Head and neck cancer, **502**
 carcinoma of unknown primary and, 120e-4
 clinical features of, 503
 diagnosis of, 503-505, 503f
 epidemiology of, 502
 etiology of, 502-503
 genetic considerations in, 503
 global considerations in, 502
 histopathology of, 503
 incidence of, 502
 infections in, 484t
 lymphadenopathy in, 408
 paraneoplastic syndromes in, 609t
 premalignant lesions, 503
 second malignancies in, 503
 second malignancies in survivors of, 623
 staging of, 504, 504f
 treatment of
 complications of, 505-506
 gene therapy, 91e-4
 late consequences of, 623, 624t
 localized disease, 505
 locally or regionally advanced disease, 505
 recurrent/metastatic disease, 505
 Head drop, 462e-1
 Head impulse test, 149
 Head injury, **457e-1**
 amnesia in, 183
 aphasia in, 178
 autonomic storm following, 2643
 brain abscess and, 901
 clinical syndromes of, 457e-4
 concussion in. *See* Concussion
 contusion, 457e-1, 457e-1f
 cranial nerve injuries, 457e-2
 dementia and, 173, 2608
 diffuse axonal injury, 457e-2, 457e-2f
 emergency care of, 457e-1
 epidural hematoma in. *See* Epidural hematoma
 epilepsy and, 2544, 2547
 frontal lobe syndromes and, 183-184
 grading of, 457e-5, 457e-5t
 headache in, 107t, 110-111
 hyperbaric oxygen therapy for, 477e-4t
 hypokalemia in, 305
 hyponatremia in, 299
 incidence of, 457e-1
 of intermediate severity, 457e-4
 intracranial hemorrhage in, 457e-2, 2584
 minor, 457e-4
 neurogenic shock in, 1750
 postconcussion syndrome, 457e-5, 471e-3, 471e-5-6
 prognosis of, 457e-5
 recurrent, 173, 457e-4, 471e-2, 2608
 respiratory acidosis in, 322
 seizures and, 457e-2-3, 2547
 severe, 457e-4-5
 skull fractures, 457e-2
 subdural hematoma. *See* Subdural hematoma
 in war veterans, 152e-9, 471e-2-3, 471e-3t, 471e-5-6
 Head lice (*Pediculus capitis*), 2745-2746, 2746f
 Head tilt test, 209
 Head-up tilt test, 273e-5, 2641, 2641t
 Health care-associated infections, **911**
A. baumannii, 1037
 antibiotic-resistant, 918
C. difficile, 858, 913. *See also* *Clostridium difficile*-associated disease
 in cancer patient, 492
 central venous catheter-related. *See* Catheter-related bloodstream infections
 chickenpox, 913
 in critically ill patient, 1734
 diarrhea, 855
 endocarditis, 819
 enterococcal, 973
 epidemiology of, 913
 fungal, 917
 HIV, 1223
 HRSV, 1205
Klebsiella spp., 1031
 measles, 1296
 Medicare coverage of, 912t
 pneumonia, 767t, 803, 803t, 809, 914-915, 958. *See also* Ventilator-associated pneumonia
 prevention of, 12e-3, 492, 913, 913t, 914t, 916, 1734
S. aureus, 955
 sinusitis, 226
 surgical-site. *See* Surgical-site infections
 surveillance, 911-912
 urinary tract, 868, 913-914, 914t
 Health care costs
 financing of, 15e-1-2
 as global health issue, 9
 pricing issues and, 15e-3
 of terminally ill patient, 65
 Health care disparities. *See* Racial/ethnic disparities in health care
 Health care proxy, 59-60
 Health care quality. *See* Quality in health care
 Health care safety. *See* Safety in health care
 Health care workers
 brain drain, 13-14
 errors of, 12e-1
 global burden of disease and, 13e-3, 13e-4f
 hand hygiene practices of, 913
 health risks of, 17e-4
 employee health service issues and, 918
 HBV exposure/transmission, 1284
 HIV exposure/transmission, 918, 1222-1223, 1283-1284
 human bite-wounds, 167e-2
 shift-work disorder, 193-194
 tuberculosis, 1284
 immunizations in, 918
 minority, 16e-6
 Health insurance
 administered prices in, 15e-3
 Affordable Care Act and, 15e-2
 employment-based, 15e-1-2
 medical care usage and, 15e-2-3
 moral hazard and, 15e-2-3
 selection issues in, 15e-1-2
 Health systems reform
 leadership and, 13e-5-6
 primary care in, 13e-1
 public policy and, 13e-5
 service delivery approaches for, 13e-5
 universal coverage and, 13e-4-5, 13e-5f
 Healthy volunteer effect, 480
 Hearing, 217, 218f
 Hearing aids, 219, 223-224
 Hearing loss, **217**
 age-related, 219, 219f
 in Alport's syndrome, 2514
 approach to the patient, 219f, 220-221
 audiologic assessment of, 221-222
 from barotrauma, 477e-7
 channelopathies in, 444e-2, 444e-2t
 conductive, 217-219
 drug-induced, 219
 evoked responses in, 223
 genetic causes of, 85e-6, 85e-6t, 220, 221t, 222t, 444e-2t, 444e-4
 imaging studies in, 223
 mixed, 220
 mumps-associated, 231e-3
 noise-induced, 224-225, 225t
 in osteogenesis imperfecta, 2509
 physical examination in, 220
 prevention of, 224-225
 sensorineural, 85e-6t, 219-220
 sudden, 220
 treatment of, 223-224, 224f
 Heart
 control of performance and output of, 265e-7-8, 265e-8f, 265e-8t
 diastolic function of, 265e-10, 265e-10f
 electrophysiology of, **273e-1**
 embryogenesis of, 265e-11-12, 265e-11f, 1520

- fetal development of, 1520
functional assessment of, 265e-9-10, 265e-10f
metabolism in, 265e-10-11
physical examination of, 1447-1448. *See also*
Heart murmurs; Heart sounds
regenerative potential of, 265e-12
tissue engineering strategies, 92e-2, 92e-3v
- Heart block. *See* Atrioventricular (AV)
conduction block
- Heartburn, 261, 262, 1876, 1900. *See also*
Gastroesophageal reflux disease
(GERD); Indigestion
- Heart disease. *See* Cardiovascular disease
- Heart failure (HF), **1500, 1507**
anemia in, 1514
atrial arrhythmias in, 1514
cardiac cirrhosis in, 2062
cardiogenic shock in. *See* Cardiogenic shock
in chronic kidney disease, 1817
clinical features of
cerebral, 1504
Cheyne-Stokes respiration, 1503-1504
dyspnea, 47e-3, 1503, 1661
edema, 251-252, 253
fatigue, 152, 1503
gastrointestinal, 1504
hemoptysis, 246
hypokalemia, 307
hyponatremia, 299, 302
hypoxia, 248
orthopnea, 1503
paroxysmal nocturnal dyspnea, 1503-1504
pleural effusion, 1716-1717
splenomegaly, 411
definition of, 1500
depression in, 1514
diagnosis of, 1504-1505
biomarkers, 1505
chest radiograph, 308e-7f, 1505
echocardiography, 1505
laboratory tests, 1504-1505
left ventricular function, 1505
in new-onset disease, 270e-20-21, 270e-21f,
270e-22f, 270e-27v
stress testing, 1505
differential diagnosis of, 1505
effect on drug concentration and response, 37
epidemiology of, 15, 1500-1501
etiology of, 1501, 1501t
folate deficiency in, 647
frailty and, 77
functional classification of, 1439-1440, 1440t,
1501, 1501t, 1581t
global considerations in, 15, 1500-1501
hypertension and, 1615, 1625
in infective endocarditis, 818, 825
in ischemic heart disease, 1579, 1590
mitral regurgitation in, 1515
myocardial infarction and, 1607-1608
pathogenesis of
depressed ejection fraction, 1501-1503,
1501t, 1502f
left ventricular remodeling, 1502-1503,
1503t
preserved ejection fraction, 1501, 1501t,
1508f
physical examination in, 1504
prognosis of, 1501
in rheumatic fever, 2153
sleep-disordered breathing in, 1514
sudden cardiac death in, 1514
treatment of
in acute decompensated disease, **1507**
inotropic therapy, 1509-1510, 1510t
neurohormonal antagonists, 1510
principles of, 1507-1508, 1509f
vascular therapy, 1509, 1510t
volume management, 1508-1509, 1510t
in association with STEMI, 1578
cardiac resynchronization therapy, 1514
cardiac transplantation. *See* Cardiac
transplantation
cellular and gene-based, 1515
comorbidity management, 1514
global considerations in, 1516
implantable cardiac defibrillator, 1514,
1515t, 1770
with preserved ejection fraction, 1507,
1508f
racial/ethnic disparities in, 16e-1
with reduced ejection fraction, **1510**
anticoagulants and antiplatelets, 1513
ARBs, 1511-1512
arteriovenous vasodilators, 1512
calcium channel antagonists, 1513
digoxin, 1512
diuretics, 1512-1513
enhanced external counterpulsation,
1513
exercise, 1513
fish oil, 1513
heart rate modification, 1512
inflammation, 1513
micronutrients, 1513
neurohormonal antagonists, 1511
novel neurohormonal antagonists, 1513
principles of, 1510-1511, 1511f
statins, 1513
sleep disordered-breathing, 1514
specialized nutritional support, 98e-4
supportive care, 1515-1516
surgical, 1514-1515
ventricular assist devices, 1518-1519, 1519f
- Heartland virus, 1309t
- HeartMate II LVAD, 1518, 1519f
- Heart murmurs
in aortic regurgitation, 51e-1, 51e-1f, 51e-7,
1448f, 1449, 1536
in aortic root-to-right-heart shunts, 1523
in aortic stenosis, 51e-1, 51e-2f, 51e-3, 51e-7,
1448, 1448f, 1531
approach to the patient, **51e-1**, 1441, 1441f
in atrial septal defect, 51e-1, 51e-3-4, 51e-6,
1448, 1521
auscultation of, 51e-6-8, 51e-7f, 51e-7t,
1448-1449, 1448f
benign, 1449
character of, 51e-1
clinical context of, 51e-8
continuous, 51e-1f, 51e-2t, 51e-6, 51e-7f,
1449
diastolic, 51e-1f, 51e-2t, 51e-5-6, 51e-6f,
1449
duration of, 51e-1
dynamic evaluation of, 51e-6-8, 51e-6f,
1449-1450, 1450t
echocardiography in, 51e-8, 1440
etiology of, 51e-1, 51e-2t
in hypertrophic cardiomyopathy, 1448, 1569
in infective endocarditis, 818
intensity of, 51e-2, 1450, 1450t
location and radiation of, 51e-1-2, 51e-2f
in mitral regurgitation, 51e-1-5, 51e-1f,
51e-2f, 1448, 1448f, 1545
in mitral stenosis, 51e-1, 51e-1f, 51e-6, 51e-6f,
1448f, 1449, 1540
in mitral valve prolapse, 51e-2, 51e-3, 51e-3f,
51e-4, 51e-7, 1449, 1449f, 1546
in patent ductus arteriosus, 51e-2f, 1449
in prosthetic valves, 1450
in pulmonic regurgitation, 51e-5-6, 1449,
1550
in pulmonic stenosis, 51e-1f, 51e-2f, 51e-3,
51e-4f, 1448, 1525
systolic, 51e-1f, 51e-2-5, 51e-2t
early, 51e-2-3
holosystolic, 51e-1f, 51e-4-5, 51e-5f
late, 51e-4
mid-systolic, 51e-3-4, 51e-3f
in tricuspid regurgitation, 1448, 1548
in tricuspid stenosis, 51e-1f, 1449, 1547
in ventricular septal defect, 51e-1f, 51e-2-5,
51e-2f, 1449
- Heart rate
deep breathing and, 2640, 2641t
intrinsic, 1468
in pregnancy, 45
resting, 1468
in shock, 1730t
during sleep, 187
Valsalva maneuver and, 2640, 2640t
- HEART score, 102f
- Heart sounds. *See also* Heart murmurs
in aortic regurgitation, 1536
in aortic stenosis, 1531
in atrial septal defect, 1446f, 1447, 1521
diastolic, 1447-1448
in heart failure, 1504
in hypertension, 1622
in ischemic heart disease, 1581-1582
in mitral regurgitation, 1545
in mitral stenosis, 1540
in mitral valve prolapse, 1546
in myocardial infarction, 1600
in myxoma, 51e-6, 289e-1
normal, 1446f, 1447
in pericardial disease, 1450
in prosthetic valves, 1450
in pulmonary hypertension, 1655
in pulmonic stenosis, 1525
second, splitting of, 51e-7, 51e-7f, 1446f, 1447
systolic, 1447
in tricuspid regurgitation, 1548
in tricuspid stenosis, 1547
- Heart transplantation. *See* Cardiac
transplantation
- HeartWare Ventricular Assist System, 1519
- Heartworm, 1423
- Heat conservation, 124
- Heat cramps, 479e-2
- Heat edema, 479e-1
- Heat exhaustion, 479e-2
- Heat-labile enterotoxin, *E. coli*, 853
- Heat loss, 124, 478e-1
- Heat production, 124
- Heat-related illness, **479e-1**
cooling strategies for, 479e-3
differential diagnosis of, 124, 479e-1, 479e-2t
heat cramps, 479e-2
heat exhaustion, 479e-2
heatstroke, 124, 479e-2-3, 479e-3t
minor syndromes of, 479e-1
prognosis of, 479e-4
resuscitation in, 479e-3-4
risk factors for, 479e-1, 479e-2t
- Heat sensitivity, in MS, 2663
- Heat shock protein, 372e-24t
- Heat stability test, hemoglobin, 637
- Heat-stable enterotoxin, *E. coli*, 853
- Heatstroke, 124, 479e-2-3, 479e-3t
- Heat syncope, 479e-1
- Heat therapy, for dysmenorrhea, 338
- Heat waves, 151e-2
- Heavy breathing, 47e-1t
- Heavy chain diseases, 718-719
- Heavy metal pigmentation, 240t, 361
- Heavy metal poisoning, **472e-1**, 1862. *See also*
specific metals
- Heberden's nodes, 2226f
- Hedgehog pathway, 387, 500, 500f
- Heel
bursitis of, 2247
plantar fasciitis of, 2249-2250
Heel-knee-shin maneuver, 2539
- Heimlich maneuver, 1768
- Heinz bodies
after splenectomy, 413
in G6PD deficiency, 655
laboratory identification of, 81e-1, 81e-5f
in sickle cell syndromes, 636
in splenomegaly, 410
unstable hemoglobin variants and, 637
- HELEX septal occluder, 297e-7f
- Helicobacter cinaedi* infection, 1059, 1059t
- Helicobacter fennelliae* infection, 1059, 1059t
- Helicobacter pylori*, 1038-1039, 1915
- Helicobacter pylori* infections
antibiotic resistance in, 934
breath odor in, 242
clinical features of, 1039-1040
diagnosis of, 262, 1040, 1040t, 1919, 1920t
epidemiology of, 1038, 1915

- Helicobacter pylori* infections (Cont.):
 gastric cancer and, 534, 1039–1040, 1931
 gastritis and, 1930, 1931, 1931f
 host factors in, 1916, 1916f
 indigestion and, 261
 MALT lymphoma and, 536, 697, 703, 1038
 natural history of, 1916, 1917f
 pathophysiology of, 1038–1039, 1039f, 1915–1916, 1916f
 in peptic ulcer disease, **1915**
 bacterial factors in, 1916f
 diagnosis of, 1919, 1920t
 epidemiology of, 1039, 1039f, 1915
 host factors in, 1916f
 mechanisms of, 1916, 1916f
 natural history of, 1916, 1917f
 pathophysiology of, 1915–1916
 recurrent bleeding in, 277
 in polycythemia vera, 673
 prevention of, 1042
 thrombocytopenic purpura and, 728
 treatment of
 adverse effects of, 1042, 1923
 antibiotic resistance in, 1042, 1922
 approach to, 1041, 1041f, 1924
 failure in, 1042, 1924
 in gastric lymphoma, 536
 indications for, 1922
 investigational approaches to, 1923
 quadruple therapy, 1922, 1922t
 regimens, 263, 1042t, 1922, 1922t
 reinfection after, 1923
 “test and treat” approach, 262, 1922
 triple therapy, 1922, 1922t
- Heller myotomy, for achalasia, 1905
- HELLP syndrome, 46, 49, 1865, 2019
- Helminthic infections, 255e-1–2. *See also specific infections*
- Helminthic parasites, 255e-1–2. *See also specific organisms*
- Helper T cells. *See* CD4+ T cells
- Hemagglutinin, filamentous, 1021, 1022
- Hemangioblastoma, 578, 612t
- Hemangioendothelioma, 119e-3, 552
- Hemangioma
 cardiac, 289e-2
 cherry, 76e-4f, 364t, 365
 of liver, 553
 retinal, renal cell carcinoma and, 578
 of small intestine, 536
- Hemangiopericytoma, 609t
- Hemarthrosis, 403, 2241–2242
- Hematemesis, 245, 259, 276. *See also*
 Gastrointestinal bleeding
- Hematin, 2529
- Hematochezia, 276, 1899, 1899f. *See also*
 Gastrointestinal bleeding
- Hematocrit
 age-related changes in, 394t
 elevated, 393, 399, 405
 normal values, 392, 394t, 2755t
 in pregnancy, 394t
- Hematologic disease
 in cirrhosis, 2067
 fatigue in, 152
 paraneoplastic. *See* Paraneoplastic syndromes, hematologic
 in pregnancy, 49
 reference values for, 2754–2756t
 in rheumatoid arthritis, 2138
 in SLE, 2129t, 2131
- Hematoma
 epidural, 441e-32f, 457e-3, 457e-3f, 2655
 in hemophilia, 733, 734
 muscle, spontaneous, 404
 spinal epidural, 2655
 subdural. *See* Subdural hematoma
- Hematomyelia, 2655
- Hematopoiesis, 81e-5f, 392, 392f, 410
- Hematopoietic differentiation, 89e-3f
- Hematopoietic growth factor (type 1) receptor, 372e-15–16
- Hematopoietic stem cells, **89e-1**, 392
 cell division in, 89e-1
 in circulation, 89e-1–2
 developmental biology of, 89e-1
 differentiation of, 89e-2, 89e-3f
 excess capacity of, 89e-2
 features of, in HSCT, 139e-1. *See also* Hematopoietic stem cell transplantation (HSCT)
 functions of, 89e-1, 89e-1f
 microenvironment of, 89e-2
 mobility of, 89e-1–2
 quiescence of, 89e-2
 self-renewal of, 89e-2–4, 89e-4f
 in tumor angiogenesis, 102e-14f, 102e-15f
- Hematopoietic stem cell transplantation (HSCT), **139e-1**
 for ALL, 139e-5–6, 139e-5t
 allogenic, 103e-7, 139e-1, 919
 for AML, 139e-5, 139e-5t, 684–685
 for amyloidosis, 722
 for aplastic anemia, 139e-2, 139e-5, 139e-5t, 666
 for autoimmune and inflammatory diseases, 372e-31
 autologous, 103e-7, 139e-1, 919
 bone marrow collection for, 139e-1, 139e-2
 for breast cancer, 139e-6, 531
 categories of, 139e-1–2
 for CLL, 139e-5t, 139e-6, 703
 for CML, 139e-5t, 139e-6, 692–693, 693t
 complications of
 early direct chemoradiotoxicities, 139e-3
 graft failure, 139e-3
 graft-versus-host disease. *See* Graft-versus-host disease (GVHD)
 hemorrhagic cystitis, 1798
 infections, **919**
 bacterial, 919–920, 1084
 CMV, 1191t, 1192
 fungal, 920–921
 parasitic, 921
 prophylaxis for, 139e-4, 139e-4t
 risks of, by type of transplant, 920t
 sources and timing of, 139e-4, 920t
 viral, 921–923, 921t, 1185, 1208
 late, 623–624, 624t
 late direct complications, 139e-3
 microangiopathic kidney injury, 1864, 1864t
 neuropathy, 2685
 cord blood, 139e-2, 919
 diffuse alveolar hemorrhage following, 245
 for diffuse large B-cell lymphoma, 706
 engraftment, 139e-2–3
 for hemoglobinopathies, 139e-5, 639
 for Hodgkin’s lymphoma, 139e-5t, 139e-6
 immunization in recipient, 928–929, 929t
 for immunodeficiency disorders, 139e-5
 jaundice following, 285
 for mucopolysaccharidoses, 432e-6
 for multiple myeloma, 139e-5t, 139e-6, 717
 for myelodysplasia, 139e-5t, 139e-6, 671
 for non-Hodgkin’s lymphoma, 139e-5t, 139e-6
 for nonmalignant diseases, 139e-5
 for osteopetrosis, 426e-4
 for paroxysmal nocturnal hemoglobinuria, 662
 peripheral blood stem cells for, 139e-1–2
 for phagocyte disorders, 423
 posttransplant relapse, 139e-6
 preparative regimens, **139e-2**
 for primary myelofibrosis, 676
 procedure for, 139e-2
 for pyruvate kinase deficiency, 654
 for SCID, 139e-2, 139e-5, 139e-5t, 2108
 for sickle cell anemia, 139e-2, 139e-5, 636
 survival rates following, 139e-5t
 for testicular cancer, 139e-6
 for thalassemia, 139e-2, 139e-5, 139e-5t, 639
- Hematopoietins, 372e-14, 372e-14t
- Hematorporphyrins, 103e-5
- Hematuria, **294**
 in Alport’s syndrome, 2514
 approach to the patient, 292f
 benign familial, 1847
 in bladder cancer, 576
 in bleeding disorders, 404
 in glomerular disease, 62e-11f, 294, 1834, 1835–1836t
 gross, 294
 in hemophilia, 733
 isolated, 294
 microscopic, 294
 in nephrolithiasis, 1868
 in polycystic kidney disease, 1851–1852
 in renal cell carcinoma, 578
- Heme
 biosynthesis of
 enzymes and genes in, 625, 2523t, 2524f
 pathway for, 2521–2522, 2522f, 2523–2524, 2524f
 regulation of, 2523t, 2525
 characteristics of, 390, 631
- Heme oxygenase, 280
- Hemianopia, 198, 198f, 199
- Hemiatrophy, facial, 2649
- Hemiballismus, 2623
- Hemicrania continua, 109t, 2596
- Hemidesmosomal protein, 372e-24t
- Hemifacial spasm, 2648, 2664
- Hemilaminectomy, 121
- Hemin, 2529
- Hemiparesis, 156, 156f
- Hemipteran bite, 2750
- Hemispatial neglect, 180–182, 180f, 181f
- Hemispherectomy, 2557
- Hemizygous, 435
- Hemobilia, 277, 2084
- Hemocult test. *See* Fecal occult blood
- Hemochromatosis, **2514**
 classification of, 2514, 2514t
 clinical features of, 450, 2516–2517
 arthropathy, 2220, 2241, 2517–2518
 cardiac, 270e-21, 290e-1t, 1471, 1564, 2518
 cutaneous, 360, 360t
 diabetes mellitus, 2517
 hepatic, 367e-1–2, 367e-1t, 367e-2t, 1992, 2062, 2517
 hypogonadism, 2365, 2518
 diagnosis of, 2517
 CMR, 270e-21, 270e-27v
 endomyocardial biopsy, 1564, 1564f
 laboratory tests, 1993t, 2517, 2517t
 liver biopsy, 366e-4f, 366e-5f, 2517
 early interventions for, 450t
 environmental factors in, 437
 in females vs. males, 437
 hepatocellular carcinoma and, 366e-4f, 545
 hereditary, **367e-1**
 clinical features of, 367e-1, 367e-1t, 367e-2t, 2062
 diagnosis of, 367e-1, 2062
 gene mutations in, 367e-1t, 2514t, 2515
 genetic testing for, 367e-1, 450, 450t, 1992
 treatment of, 367e-2
 pathophysiology of, 2515–2516, 2516f
 precirrhotic (early), 2514
 prevalence of, 2514
 prognosis of, 2518
 screening for, 2517–2518
 secondary. *See* Iron overload
 treatment of, 1564, 2220, 2518
V. vulnificus infection in, 1066
- Hemodialysis
 for acute kidney injury, 1810–1811
 bullous dermatosis of, 361t, 362
 for chronic kidney disease, **1822**
 access for, 1823
 complications of, 1824
 components of, 1822–1823, 1823f
 dose of, 1824
 goals of, 1823–1824
 patient education, 1821

- preparation for, 1821
 timing of, 1821
 in diabetes mellitus, 2426
 for drug- and toxin-induced acidosis, 64e-9, 319
 folate deficiency in, 647
 for hypercalcemia, 314, 610, 2480t, 2481
 for hyperkalemia, 312
 for hypermagnesemia, 2463
 oxalosis in, 2237
 pericarditis in, 1575
 for poisoning/drug overdose, 473e-7, 473e-8t, 473e-9t
 for rewarming, 478e-3, 478e-3t
- Hemodynamic assessment
 calculation and normal values, 1461t, 1748t, 1762t
 in cardiac catheterization, 1461–1463
 in cardiac tamponade, 1463t
 in myocardial infarction, 1607, 1762t
 normal waveforms, 1461t
 in pericarditis, 1463t
 in restrictive cardiomyopathy, 1463t
 in shock, 1762t
 in valvular heart disease, 1462, 1462f
- Hemoglobin, **631**
 age-related changes in, 394t
 biosynthesis of, 632–633
 developmental biology of, 632
 disorders of. *See* Hemoglobinopathies
 electrophoresis of, 633, 634t
 elevated, 399
 functions of, 632
 genetics of, 632–633
 heme-heme interaction, 632
 high-affinity, 636
 increased, in hypoxia, 248
 iron content of, 625t
 in iron metabolism, 625
 low-affinity, 636
 in microcytic anemia, 628t
 normal, 393, 394t, 2755t
 oxygen-carrying capacity, 1748t
 in pregnancy, 394t
 reduced, in cyanosis, 248–249
 structure of, 631, 631f
 testosterone therapy and, 2372–2373
- Hemoglobin A, 631, 633
 Hemoglobin A_{1c} (HbA_{1c})
 goals in diabetes, 2408t, 2410
 goals in elderly, 82
 reference values for, 2760t
- Hemoglobin A₂, 631
 Hemoglobin-based oxygen carriers (HBOC), 662
 Hemoglobin C, 634, 636t
 Hemoglobin C disease, 81e-4f
 Hemoglobin E, 636t, 638–639
 Hemoglobinemia, 81e-12f
 Hemoglobin H disease, 638, 638t
 Hemoglobin Kansas, 636, 636t
 Hemoglobin Köln, 636t
 Hemoglobin Lepore, 638
 Hemoglobin M. Iwata, 636t
- Hemoglobinopathies, **633**
 acquired, 633, 633t, 639
 α-globin, 633
 aplastic crisis in, 639
 arthropathies associated with, 2242, 2242t
 β-globin, 633
 classification of, 633, 633t
 detection and characterization of, 633–634, 636–637
 epidemiology of, 633
 genetic considerations in, 633
 global considerations in, 633
 with hemoglobins with altered oxygen affinity, 636, 636t
 HSCT for, 139e-5
 in pregnancy, 49
 red cell indices in, 399
 structural, 633t, 634, 634t. *See also* Sickle cell anemia
- thalassemias, 633, 633t. *See also* Thalassemia syndromes
 with unstable hemoglobins, 636–637, 636t
 Hemoglobin-oxygen dissociation curve, 247, 393, 632, 632f
- Hemoglobin S, 636t. *See also* Sickle cell anemia
 Hemoglobin SC disease, 634t, 635
- Hemoglobinuria
 march, 657, 657t
 paroxysmal cold, 657t, 659
 paroxysmal nocturnal. *See* Paroxysmal nocturnal hemoglobinuria
- Hemoglobin Yakima, 636, 636t
 Hemojuvelin, 2515, 2515f
- Hemolysins, 145e-7
- Hemolysis
 acute immune-mediated, 138e-3–4
 compensated, 651
 delayed immune-mediated, 138e-4
 extravascular, 658, 659f
 hyperbilirubinemia and, 2000
 intravascular, 657t, 658
- Hemolytic anemia, **649**
 acquired, 649t, 657, 657t
 immune, 658, 658t
 autoimmune hemolytic anemia, 377e-4, 658, 659f
 cold agglutinin disease, 659
 drug-induced, 658t
 microangiopathic, 81e-4f, 282, 657, 657t, 726f
 paroxysmal cold hemoglobinuria, 657t, 659
 paroxysmal nocturnal hemoglobinuria. *See* Paroxysmal nocturnal hemoglobinuria
 in infections, 657t, 658
 from mechanical destruction of red cells, 657
 microangiopathic, 657, 657t
 from toxic agents and drugs, 659–660
 in alcoholic liver disease, 2059
 aplastic crisis in, 639
 in babesiosis, 1385
 bone marrow examination in, 81e-8f
 chronic nonspherocytic, 656
 clinical features of, 393, 399, 649–650, 649t
 vs. compensated hemolysis, 650–651
 diagnosis of, 399, 649–650, 649t
 drug-induced, 39, 44
 folate deficiency in, 647
 in G6PD deficiency, 39t, 40
 inherited, 649t
 with cation transport disorders, 653, 653t
 with enzyme abnormalities, 653–654, 654t
 pyruvate kinase deficiency, 281, 650f, 653–654, 654t
 familial hemolytic-uremic syndrome, 657
 with red cell membrane-cytoskeleton abnormalities, 651, 653t
 hereditary elliptocytosis, 652f, 653, 653t
 hereditary spherocytosis. *See* Hereditary spherocytosis
 with redox metabolism disorders, 654f, 657
 glucose-6-phosphate dehydrogenase deficiency. *See* Glucose-6-phosphate dehydrogenase (G6PD) deficiency
 pyrimidine-5'-nucleotidase (P5N) deficiency, 431e-5t, 431e-6f, 654t, 657
 pathophysiology of, 650–651, 650f
 peripheral blood smear in, 81e-4f
 in sickle cell anemia, 634
- Hemolytic streptococcal gangrene. *See* Necrotizing fasciitis
- Hemolytic-uremic syndrome (HUS)
 acute kidney injury in, 292, 1806t
 after infectious diarrhea, 855t, 1864
 atypical, 657, 730, 1832
 in cancer, 1796
 clinical features of, 133t, 1863
 acute nephritis syndrome, 292
 chemotherapy-related, 1796
 jaundice, 282
- rash, 135
 Shiga toxin-related, 1057
 diagnosis of, 1807
E. coli infections and, 267, 856, 1030, 1031, 1863
 etiology of, 133t, 267
 familial (atypical), 657, 1863
 genetic considerations in, 1848
 not associated with diarrhea, 730
 pathogenesis of, 265e-2, 730, 1030, 1796, 1848, 1863
 renal biopsy in, 62e-9f, 1848
 Shiga toxin-related, 1057
Shigella infections and, 1056–1057, 1864
 in SLE, 2133–2134
 vs. thrombotic thrombocytopenic purpura, 729, 1863
 treatment of, 730, 1796–1797, 1848, 1864
- Hemophagocytic lymphohistiocytosis, 375e-2, 2112
- Hemophagocytic syndrome, 135e-4, 135e-9
- Hemophilia, **732**
 arthropathy in, 2241–2242
 autosomal, 731. *See also* von Willebrand disease
 bleeding in, 403
 carriers of, 735
 clinical features of, 732–733
 complications of, 734–735
 diagnosis of, 733t
 genetic considerations in, 732–733, 733t
 pathogenesis of, 732–733
 treatment of, 91e-3, 733t, 734
- Hemophilia A, 403, 437, 450t, 733. *See also* Hemophilia
- Hemophilia B, 450t, 733. *See also* Hemophilia
- Hemoptysis, **245**
 in cancer, 1793–1794
 etiology of, 245–246
 evaluation of, 246–247, 246f
 in lung cancer, 510, 510t
 massive, 246
 in mitral stenosis, 1540
 in pulmonary embolism, 100
 in respiratory disease, 1661
 treatment of, 247
 in tuberculosis, 1108
- Hemorrhage
 anemia of, 662
 epidural spinal, 2584
 gastrointestinal. *See* Gastrointestinal bleeding
 hypovolemia in, 298
 intracranial. *See* Intracranial hemorrhage
 retinal, 40e-2–3, 40e-2f, 40e-3f
 splinter, 1443
 subconjunctival, 199
- Hemorrhagic cystitis, 1208, 1798
- Hemorrhagic fever, viral. *See* Viral hemorrhagic fevers
- Hemorrhagic fever with renal syndrome, 1321
- Hemorrhagic telangiectasia, hereditary. *See* Osler-Weber-Rendu syndrome
- Hemorrhagic thrombocytopenia. *See* Essential thrombocytosis (ET)
- Hemorrhoidal cushions, 1976
- Hemorrhoidal disease, **1976**
 classification of, 1976, 1976t
 clinical features of, 1976–1977
 diagnosis of, 1977
 epidemiology of, 1976
 etiology of, 273
 at high altitudes, 476e-4
 internal, endoscopic findings in, 1899, 1899f
 pathophysiology of, 1976
 treatment of, 1976t, 1977
- Hemorrhoidectomy, 1976t, 1977
- Hemosiderin, 346, 397, 2514
- Hemosiderosis, 639, 2514
- Hemostasis, **400**
 antithrombotic mechanisms, 402
 antithrombotic mechanisms in, 402f
 disorders of, 403–404, 403t, 406t. *See also* specific disorders

- Hemostasis (*Cont.*):
 endothelium in, 265e-2
 fibrin clot formation, 401
 fibrin clot formation in, 401f
 fibrinolytic system in, 401f, 403
 platelet plug formation in, 400–401
 tests of, 732, 733f
- Hemostatic sealants, 740
- Hemosuccus pancreaticus, 277
- Hemothorax, 1718
- Henbane, 473e-9t
- Henderson-Hasselbalch equation, 315
- Hennekam's syndrome, 1654
- Henoch-Schönlein purpura. *See* IgA vasculitis (Henoch-Schönlein)
- Hensin, 332e-9
- Hepadnaviruses, 214e-2t, 214e-3f. *See also* Hepatitis B virus (HBV)
- Heparan proteoglycans, 402
- Heparan sulfate glycosaminoglycans, 265e-2
- Heparin, 749
 action of, 749–750, 750f
 for acute limb ischemia, 1646
 adverse effects of
 alopecia, 354
 bleeding, 751
 osteoporosis/osteopenia, 752, 2218t
 skin necrosis, 379
 thrombocytopenia. *See* Heparin-induced thrombocytopenia
 transaminase elevation, 752
 in cardiac catheterization, 1461
 coagulation test abnormalities and, 407
 for DIC, 738
 dosage of, 751
 drug interactions of, 44, 103e-10t
 for DVT/PE, 47, 1635, 1635t
 for frostbite, 478e-4
 limitations of, 751, 751t
 vs. low-molecular-weight heparin, 752t. *See also* Low-molecular-weight heparin (LMWH)
 management of
 before endoscopic procedures, 1889t
 before lumbar puncture, 443e-1
 treatment monitoring, 750–751
 monitoring treatment with, 2754t
 for myocardial infarction, 1597, 1598t, 1606
 in parenteral formula, 98e-6
 in PCI, 296e-1
 pharmacology of, 750, 751t
 for thrombophlebitis, 613
 for VTE prophylaxis, 1637t
 for warfarin necrosis, 379
- Heparin-binding proteins, 750
- Heparin-induced thrombocytopenia, 727
 clinical features of, 751, 751t
 delayed onset, 727
 diagnosis of, 727–728, 752
 pathogenesis of, 44
 skin manifestations of, 368
 time course of, 727, 727f
 treatment of, 728, 752, 752t
- Hepatic artery embolization, 567, 2338
- Hepatic encephalopathy. *See* Encephalopathy, hepatic
- Hepatic hydrothorax, 288, 1717
- Hepatic lipase deficiency, familial, 2440t, 2443
- Hepatic necrosis, drug-induced, 43–44
- Hepatic steatosis. *See* Fatty liver disease
- Hepatic stem cells, 453t
- Hepatic vein thrombosis. *See* Budd-Chiari syndrome
- Hepatitis
 acute viral, 2004
 vs. autoimmune hepatitis, 2051
 clinical features of, 283, 2013t, 2015–2016
 complications and sequelae of, 2012, 2018–2019
 diagnosis of
 algorithm for, 2017–2018, 2018t
 liver biopsy, 366e-1f, 366e-6f
 liver function tests, 283, 1998t, 2016
 serologic tests, 2016–2017
 differential diagnosis of, 2019
 etiology of. *See specific hepatitis viruses*
 extrahepatic manifestations of, 2011–2012
 fulminant, 2018
 HIV infection and, 1258
 in military forces, 152e-7
 niacin-associated, 96e-4
 pathogenesis of, 2010–2011
 pathology of, 2012
 prognosis of, 2018
 prophylaxis for, 2020
 risk factors for, 1991
 treatment of, 2020
 alcoholic, 283, 2053, 2053t. *See also* Alcoholic liver disease
 aplastic anemia and, 664
 autoimmune, 2049
 cirrhosis due to, 2060
 clinical features of, 283, 2050–2051
 definition of, 2049
 diagnosis of, 366e-2–3f, 1993t, 2031t, 2050–2051
 differential diagnosis of, 2051
 immunopathogenesis of, 372e-24t, 2049–2050
 prognosis of, 2049
 treatment of, 2051–2052
 type I, 2050
 type II, 2050
 type III, 2050
 chronic, 2031. *See also* Hepatitis B virus (HBV) infection, chronic; Hepatitis C virus (HCV) infection, chronic
 classification of, 2031, 2031t, 2032t
 grading of, 2031, 2032t
 laboratory evaluation of, 2019, 2031t
 staging of, 2031, 2031t
 coagulation disorders in, 739
 drug-induced. *See* Liver disease/failure, drug-induced
 fibrosing cholestatic, 284, 2011, 2074
 fulminant, 2018, 2020
 HSV, 1180
 inflammatory mediators in, 372e-29t
 jaundice in, 283
 neuropathy in, 2685
 physical examination in, 282–283
 radiation, 551
 toxic, 2023, 2025t, 2026t. *See also* Liver disease/failure, drug-induced
 in transplant recipient, 927
 tumor markers in, 473t
- Hepatitis A virus (HAV), 2004–2005, 2005f, 2006t
- Hepatitis A virus (HAV) infection
 clinical features of, 152e-2t, 2005f, 2013t, 2015–2016
 complications and sequelae of, 2018
 diagnosis of, 152e-2t, 1993t, 2005, 2005f, 2016, 2018t
 epidemiology of, 2012, 2013t
 etiology of. *See* Hepatitis A (HAV) virus
 global considerations in, 2012
 incidence pre- and postvaccine, U.S., 785t
 incubation period for, 152e-2t
 in military forces, 152e-2t, 152e-7
 prevention of. *See* Hepatitis A virus (HAV) vaccine
 recurrence of, after liver transplantation, 2074
 risk factors for, 1991
 serology in, 2016
 transmission of, 2012
 treatment of, 2020
- Hepatitis A virus (HAV) vaccine
 for postexposure prophylaxis, 2020–2021
 precautions for, 790t
 recommended administration of
 in adults, 787–788f, 2021, 2021t
 in cancer patients, 485t
 in children, 2021, 2021t
 in HIV infection, 1252t
 in liver disease, 1995, 2021
 in transplant recipients, 929, 929t
 in travelers, 795, 795t
- Hepatitis B core antigen (HBcAg), 2006, 2006t
- Hepatitis B e antigen (HBeAg), 2006, 2006t
- Hepatitis B surface antigen (HBsAg), 2005–2006, 2006t
- Hepatitis B virus (HBV)
 extrahepatic sites of, 2008
 gene expression and replication in, 214e-5
 genomic structure of, 2005, 2005f
 molecular variants of, 2008
 proteins and particles of, 2005–2007, 2005f, 2006t
 serologic and viral markers of, 2007–2008, 2007f, 2008f
 structure of, 214e-2t
- Hepatitis B virus (HBV) infection
 chronic, 2031
 after kidney transplantation, 927, 1831
 cirrhosis due to, 2060
 clinical features of, 2033
 diagnosis of, 366e-2f, 2031t, 2033
 global considerations in, 2060
 HBeAg-negative, 2032–2033, 2035, 2038, 2039t
 HBeAg-reactive, 2032, 2038, 2039t
 HDV infection with, 2014, 2040
 histology of, 2031–2032
 nonreplicative phase, 2032
 prognosis of, 2032
 renal involvement in, 1849
 replicative phase of, 2032
 treatment of, 2033
 adefovir, 215e-4t, 215e-9, 2035, 2037t, 2038f
 combination therapy, 215e-4t, 215e-9, 2037–2038
 entecavir, 215e-4t, 215e-9–10, 2036, 2037t, 2038f
 IFN- α , 2033
 IFN- α 2a, 2036
 IFN- α 2b, 215e-4t, 215e-10, 2035
 lamivudine, 215e-4t, 215e-9, 2034–2035, 2037t, 2038f
 liver transplantation, 2068
 novel antivirals and strategies, 2038
 PEG IFN, 215e-4t, 215e-10, 2035–2036, 2037t, 2038f, 2039, 2040t
 recommendations for, 2038–2040, 2039t, 2040t
 telbivudine, 215e-4t, 215e-10, 2036, 2037t, 2038f
 tenofovir, 215e-4t, 215e-9, 2037, 2037t, 2038f
 clinical features of, 152e-2t, 2013t, 2015–2016
 cholestasis, 284
 polyarteritis nodosa, 2187
 rash, 132t, 135
 serum sickness–like, 837
 time course of, 2007f
 complications and sequelae of, 2018–2019
 diagnosis of, 2016
 algorithm for, 2017–2018, 2018t
 laboratory tests in, 1993t
 quantitative nucleic acid test in, 150e-7–8
 serology in, 152e-2t, 2007–2008, 2008f, 2016–2017, 2017t, 2018t
 epidemiology of, 2012–2014, 2013t
 escape mutants in, 2008
 etiology of. *See* Hepatitis B virus (HBV)
 extrahepatic manifestations of, 2011–2012
 genotypes in, 2005–2006
 global considerations in, 2013–2014
 HDV infection with, 2009, 2011, 2039
 in health care workers, 1284
 hepatocellular carcinoma and, 214e-7, 544, 545t
 HIV infection and, 1258, 1273, 2030
 incidence of, 785t, 869
 incubation period for, 152e-2t
 in military forces, 152e-2t, 152e-7
 pathogenesis of, 2010–2011
 in pregnancy, 49
 prevention of, 152e-2t, 1284. *See also* Hepatitis B virus (HBV) vaccine
 prognosis of, 2018

- recurrence of, after liver transplantation, 2068, 2074
- risk factors for, 1991, 2014t
- screening for, 882–883, 2013–2014, 2014t
- transfusion-transmitted, 138e-3t, 138e-6
- transmission of, 2012–2013, 2013t
- treatment of, 2020
- urticarial vasculitis in, 363, 363t
- Hepatitis B virus (HBV) vaccine
- adverse effects of, 2218t
- contraindications to, 790t
- liver cancer risk and, 478
- for postexposure prophylaxis, 2021–2022
- recommended administration of
- in adults, 787–788f, 883, 2021, 2022t
- in cancer patients, 485t
- in hemodialysis patients, 2022t
- in HIV infection, 788f, 1252t
- in infants and children, 2021, 2022t
- in liver disease, 1995
- in transplant recipients, 929, 929t
- in travelers, 795, 795t
- Hepatitis B x antigen (HBxAg), 2005f, 2007
- Hepatitis C virus (HCV), 2006t, 2009–2010, 2009f, 2010f
- Hepatitis C virus (HCV) infection
- chronic, **2040**
- alcoholic liver disease associated with, 2052, 2052t, 2725
- aminotransferase activity in, 2041
- cirrhosis due to, 2040–2041, 2059–2060
- clinical features of, 2041
- arthritis, 837
- cardiomyopathy, 1560
- cholestasis, 284
- cryoglobulinemic vasculitis, 2041, 2048, 2180, 2191
- depression, 2714
- lichen planus, 349, 353
- osteosclerosis, 426e-5
- porphyria cutanea tarda, 390
- urticarial vasculitis, 363, 363t
- diagnosis of, 366e-1–3f, 2031t, 2041
- global considerations in, 2059–2060
- iron overload in, 2518
- liver disease progression in, 2040–2041
- prognosis of, 2041
- renal involvement in, 1849
- treatment of, **2041**
- after liver transplantation, 2048
- after relapse or nonresponse, 2048
- antiviral therapy, 2044, 2045–2046t, 2047–2049
- boceprevir, 2044, 2046–2047
- combination therapy, 215e-4t, 215e-6, 215e-10–11, 2041–2043, 2042f
- genotypes and, 2042, 2047–2048
- in HIV infection, 2048–2049
- IFN-alfacon, 215e-4t, 215e-10
- IFN- α , 2041–2042
- investigative agents, 215e-11, 2047
- liver transplantation, 2068
- PEG IFN- α 2a, 215e-4t, 215e-10, 2043–2044, 2043t
- PEG IFN- α 2b, 215e-4t, 215e-10, 2043–2044, 2043t
- PEG IFN and ribavirin, 2041–2042, 2042f
- recommendations for, 2045–2046t, 2047
- in renal failure, 2049
- ribavirin, 215e-4t, 215e-6, 2041–2042
- simeprevir, 215e-5t, 215e-11, 2047
- sofosbuvir, 215e-5t, 215e-10–11, 2047
- telaprevir, 2044, 2046–2047
- clinical features of, 152e-6t, 2013t, 2015–2016
- complications and sequelae of, 2018–2019
- diagnosis of, 150e-7–8, 152e-6t, 480e-1, 1993t, **2017**, 2018t
- epidemiology of, 2013t, 2014–2015
- etiology of. *See* Hepatitis C virus (HCV)
- genetic considerations in, 2011
- global features of, 2014–2015
- in hemophiliacs, 734, 735
- hepatocellular carcinoma and, 214e-7, 544–545, 545t, 550
- HIV infection and, 1258, 2030
- incubation period for, 152e-6t
- KIRs with, 372e-15t
- lymphoid malignancies and, 697
- pathogenesis of, 2011
- prevention of, 2022
- prognosis of, 2018
- recurrent, after liver transplantation, 2068, 2074–2075
- risk factors for, 1991, 2015t
- screening for, 29t, 49, 2015, 2015t
- transfusion-transmitted, 138e-3t, 138e-6
- transmission of, 2013t, 2014–2015
- in transplant recipient, 927
- treatment of, 152e-6t, 2020
- Hepatitis D virus (HDV), 2005f, 2006t, 2008–2009
- Hepatitis D virus (HDV) infection
- chronic, 2031t, **2040**
- clinical features of, 2013t, 2015–2016
- complications and sequelae of, 2018–2019
- diagnosis of, 1993t, 2017
- epidemiology of, 2013t, 2014
- etiology of. *See* Hepatitis D virus (HDV)
- global considerations in, 2014
- HBV infection with, 2011, 2014
- pathology of, 2012
- prevention of, 2022
- prognosis of, 2018
- transmission of, 2013t, 2014
- treatment of, 215e-5t, 2020
- Hepatitis E virus (HEV), 2005f, 2006t, 2010
- Hepatitis E virus (HEV) infection
- clinical features of, 152e-2t, 2013t, 2015–2016
- diagnosis of, 152e-2t, 1993t
- epidemiology of, 152e-2t, 1991, 2013t, 2015
- etiology of. *See* Hepatitis E virus (HEV)
- global considerations in, 1991, 2015
- incubation period for, 152e-2t
- pathology of, 2012
- prevention of, 2022
- prognosis of, 2018
- in returning war veterans, 152e-2t
- risk factors for, 1991
- transmission of, 2013t, 2015
- treatment of, 2020
- Hepatitis G virus (GB virus C) infection, 1258
- Hepatocellular carcinoma (HCC), **544**
- approach to the patient, 547
- biopsy in, 547
- clinical features of, 545–546, 545t
- deaths from, 467t, 469f, 471t, 544
- diagnosis of, 366e-4f, 547, 1993t
- epidemiology of, 544
- etiology of, 102e-2, 214e-7, 544–545, 545t
- fibrolamellar, 552
- global considerations in, 544–545, 544t
- in hemophilia, 735
- hepatitis and, 101e-7, 214e-7, 477t, 544–545, 545t
- history in, 547
- imaging in, 547
- immunopathology of, 372e-25t, 547
- incidence of, 467t, 468, 468f, 471t, 544, 544t
- paraneoplastic syndromes in, 545–546, 609t
- physical examination in, 547
- prevention of, 548
- risk factors for, 544–545, 545t, 2530
- screening for, 547–548
- staging of, 546–547, 546t
- treatment of, 102e-5t, 102e-17, **548**
- adjuvant therapy, 550
- algorithm for, 548f
- chemotherapy, 550
- liver transplantation, 549–550, 2068
- new directions in, 551, 551t
- options for, 549t, 552
- response to, 551–552
- in stages I and II, 548–549, 549t
- in stages III and IV, 550–551, 551t
- targeted chemotherapy, 550–551, 551t
- transhepatic artery chemoembolization, 549, 549t
- tumor markers in, 473t
- Hepatocellular diseases, 1990, 1990t. *See also* Liver disease/failure
- Hepatocyte growth factor/c-Met pathway, 102e-11f
- Hepatocytes, 1990
- Hepatomegaly
- abdominal swelling with, 286
- in *Brucella* infections, 194e-3
- in Gaucher disease, 432e-5
- in heart failure, 1504
- in hepatocellular carcinoma, 545
- in liver disease, 1992
- in lysosomal storage diseases, 432e-5
- in schistosomiasis, 1426f, 1427
- Hepatopulmonary syndrome, 47e-3, 1992
- Hepatorenal syndrome, 1800, 1809–1810, 2066
- Hepatosplenic $\gamma\delta$ T-cell lymphoma, 135e-3, 708, 1962
- Hepcidin, 398, 630
- Hephaestin, 626
- Heptavalent botulinum antitoxin (HBAT), 261e-9
- HER2 gene mutations
- in breast cancer, 101e-3, 526, 528
- in gastric adenocarcinoma, 534
- in lung cancer, 507t, 509, 509f
- HER2/neu
- in breast cancer, 102e-10, 103e-19, 103e-21, 528
- drug response and, 39t
- in GI-NETs, 562
- Herbal medicines/supplements
- causing increased bleeding, 404, 404t
- drug interactions of, 14e-4, 14e-4t, 43, 44
- hepatotoxicity of, 2030
- nephrotoxicity of, 577, 1804, 1861
- regulation of, 14e-3
- toxicity of, 14e-3–4
- Hereditary coproporphyrria (HCP)
- clinical features of, 2522t, 2531, 2681
- diagnosis of, 2522t, 2531
- genetic considerations in, 2681
- pathophysiology of, 2522f, 2681
- treatment of, 2531, 2681
- unsafe drugs in, 2526, 2527–2529t
- Hereditary elliptocytosis, 81e-4f, 652f, 653, 653t
- Hereditary endotheliopathy, retinopathy, nephropathy, and stroke (HERNs), 2568
- Hereditary hemochromatosis. *See* Hemochromatosis, hereditary
- Hereditary hemorrhagic telangiectasia. *See* Osler-Weber-Rendu syndrome
- Hereditary inclusion body myopathy, 2195–2196
- Hereditary liability to pressure palsy, 434
- Hereditary nephritis, 294
- Hereditary neuralgic amyotrophy, 2678, 2679t
- Hereditary neuropathy(ies). *See* Peripheral neuropathy, hereditary
- Hereditary neuropathy with liability to pressure palsies (HNPP), 2678, 2679t
- Hereditary nonpolyposis colon cancer (HNPCC; Lynch's syndrome)
- clinical features of, 538t
- early interventions for, 450t, 451
- endometrial cancer in, 596
- epigenetic silencing in, 102e-7
- genetic considerations in, 101e-6t, 539
- APC tumor-suppressor gene, 101e-5
- DNA mismatch/repair, 434
- somatic mutational steps, 101e-1, 101e-1f
- genetic testing for, 450t
- ovarian cancer in, 592
- pancreatic cancer risk in, 554
- polyp distribution and characteristics in, 538t
- screening for, 434, 539
- type 2, 592
- Hereditary persistence of fetal hemoglobin, 633, 639
- Hereditary recurrent fever syndromes
- clinical features of, 2213t
- familial cold autoinflammatory syndrome, 372e-8t, 2215

- Hereditary recurrent fever syndromes (*Cont.*):
 familial Mediterranean fever. *See* Familial Mediterranean fever
 genetic considerations in, 2213t
 hyperimmunoglobulinemia D with periodic fever syndrome, 2215
 Muckle-Wells syndrome. *See* Muckle-Wells syndrome (MWS)
 neonatal-onset multisystem inflammatory disease, 2215
 treatment of, 2213t
 tumor necrosis factor- α receptor-associated periodic syndrome, 141, 422, 2214–2215
- Hereditary sensory and autonomic neuropathy (HSAN), 2643, 2679–2680t, 2680
- Hereditary spherocytosis, **651**
 anemia in, 399
 clinical features of, 651–652
 diagnosis of, 652
 genetic considerations in, 651, 652f, 653t
 peripheral blood smear in, 652f
 splenomegaly in, 411
 treatment of, 652
- Hereditary stomatocytosis, 81e-4f, 653f, 653t
- Hermansky-Pudlak syndrome, 375e-2, 730, 1715, 1949t
- Hermaphroditism. *See* Ovotesticular disorders of sexual development
- Hernia(s)
 hiatal, 1902, 1902f
 paraesophageal, 1902
- Herniation, cerebral, 1772, 1772f
- HERNS (hereditary endotheliopathy, retinopathy, nephropathy, and stroke), 2568
- Heroin
 action of, 468e-1
 adverse effects of, 462e-20, 462e-20t, 2199
 systems effects of, 468e-2
 toxic levels of, 2765t
- Herpangina, 231, 238t, 1293
- Herpes gestationis (pemphigoid gestationis), 370t, 372–373
- Herpes gladiatorum, 829, 1178
- Herpes labialis, 215e-7, 215e-8, 238t, 342f, 829
- Herpes simplex virus (HSV)
 adherence of, 145e-1–2, 145e-2t
 gene expression and regulation in, 214e-4
 for gene therapy delivery, 91e-1t
 genome of, 1175
 host response to, 1176
 laboratory identification of, 214e-8, 1180–1181
 ligand-receptor interactions of, 1175
 reactivation mechanisms of, 1175
 structure of, 214e-1f, 1175
- Herpes simplex virus (HSV) infections, **1175**
 actinomyces and, 1088
 anorectal, 881, 1178, 1956
 “asymptomatic,” 1177
 in burn wounds, 166e-1
 in cancer patient, 487, 488
 diagnosis of, 342, 1180–1181
 disseminated
 clinical features of, 132t, 134, 258, 266
 in immunosuppressed patients, 1176, 1179–1180
 rash in, 132t
 treatment of, 1181, 1182t
- encephalitis
 clinical features of, 1179, 1179f
 complications of, 1179
 differential diagnosis of, 894–896, 895t
 genetic considerations in, 1179
 vs. meningitis, 888
 MRI in, 440e-6f
 pathogenesis of, 1178–1179
 treatment of, 215e-2t, 1179, 1181, 1182t
- epidemiology of, 1176–1177
 genital, 341t
 clinical features of, 880f, 880t, 1178, 1178f
 diagnosis of, 881
 epidemiology of, 869, 879, 1177
- lymphadenopathy in, 408
 in pregnancy, 50
 prophylaxis for, 215e-7, 215e-8, 1181
 recurrence rates of, 1178
 treatment of, 215e-3t, 215e-7, 215e-8–9, 832t, 1181, 1182t
- herpes gladiatorum, 829, 1178
- herpetic whitlow, 1178, 1182t, 1263
- in HIV infection
 perirectal, 1257–1258, 1258f, 1263
 prophylaxis for, 1253t
 treatment of, 1181, 1182t, 1263
- in immunocompromised patients, 1176
- keratitis, 200, 215e-3t, 215e-9, 1178, 1182t
- latent, 1176
- meningitis, 892, 908t, 1179
- neonatal, 50, 215e-2t, 1180, 1182t
- oral-facial
 clinical features of, 238t, 1177–1178
 course of, 238t
 pathogenesis of, 233, 827, 829, 1177–1178
 treatment of, 215e-3t, 215e-7, 233, 1182t
- pathogenesis of, 1175–1176
- in pregnancy, 50, 1180
- prevention of, 1183
- reactivation of, 1176
- skin manifestations of
 distribution, 341t, 342f
 morphology, 341t
 vesicles/bullae, 132t, 238t, 361, 361t, 827, 829
- in transplant recipient
 clinical features of, 921t
 prophylaxis for, 139e-4t, 215e-6, 921
 timing of, 920t
 treatment of, 1181–1183, 1182t
- visceral, 1179–1180
 esophagitis, 1179, 1182t, 1896f, 1910
 liver, 1180
 pharyngitis, 231, 1177
 pneumonitis, 1179–1180
 upper respiratory, 1202t
- Herpesvirus(es). *See also specific viruses*
 gene expression and regulation in, 214e-4
 host defense inhibition by, 214e-8
 microRNAs in, 214e-6
 protein functions of, 214e-6
 structure of, 214e-1, 214e-1f, 214e-2t, 214e-3f
 transmission of, 214e-6
 viral assembly and egress in, 214e-5
- Herpes zoster (shingles), **1184**
 clinical features of, 341t, 1184–1185
 abdominal pain, 105
 chest discomfort, 96t, 98
 glossopharyngeal neuropathy, 2649
 neck pain, 122
 neuropathy, 2685
 ocular, 200
 oral manifestations, 237, 238t
 postherpetic neuralgia, 237, 1185, 1186, 2685
 skin manifestations, 25e-8f, 76e-8f, 827, 1184–1185, 1184f
 trigeminal neuropathy, 2646
- diagnosis of, 1185
 differential diagnosis of, 1185
 epidemiology of, 1184
 in HIV infection, 1185, 1185f, 1252t, 1263
 pathogenesis of, 89, 1183
 prevention of, 485t, 929, 1186, 1186t, 1252t
 in transplant recipient, 921, 921t, 925, 1185
 treatment of, 215e-3–4t, 215e-6–7, 215e-8, 832t, 1186
- Herpes zoster ophthalmicus, 200, 215e-4t, 1184
- Herpes zoster vaccine
 contraindications to, 790t
 recommended administration of, 29t, 84, 787–788f, 893
 storage and handling of, 791
- Herpetic whitlow, 1178, 1182t, 1263
- Herpetiform lesions, 340t
- Hers disease (type VI GSD), 433e-2t, 433e-4
- Hertel exophthalmometer, 207
- HES. *See* Hypereosinophilic syndromes (HES)
- HESX1 gene, 2256
- HESX1 gene mutations, 2325t, 2364, 2364t
- Heterophyes heterophyes, 245e-2t, 765t, 1428t, 1429. *See also* Intestinal flukes
- Heteroplasmy/heteroplasmia, 85e-3–4, 85e-5f, 439
- Heterosexual precocity, 2380
- Heterosexual sexual precocity, 2362
- Heterotropia, 197
- Heterozygote advantage, 438, 442
- Heterozygous, 433f, 435
- Heuristics, 19
- HEV (hepatitis E virus), 2005f, 2006t, 2010. *See also* Hepatitis E virus (HEV) infection
- HEVs (high endothelial venules), 372e-26
- Hexacarbonyls, 2688t, 2689
- Hexagonal-phase phospholipids test, 740
- Hexokinase (HK) deficiency, 654t
- Hexosaminidase A deficiency. *See* Tay-Sachs disease
- Hexose-6-phosphate dehydrogenase (H6PDH), 2312
- Hexoses, genetic disorders of, 435e-1t
- Heyde's syndrome, 731
- HF. *See* Heart failure (HF)
- HFE gene mutations
 global considerations in, 2519
 in hemochromatosis, 367e-1, 367e-1t, 367e-3
 in liver disease, 1992, 2515, 2518
 in porphyria cutanea tarda, 390, 2515, 2530
- 2-HG (2-hydroxyglutarate), 102e-13
- hGH-N gene, 401e-3
- hGH-V gene, 401e-3
- HHIP gene mutations, 1703
- HHS (hyperglycemic hyperosmolar state), 2417t, 2420
- HHV-6 infection. *See* Human herpesvirus-6 (HHV-6) infection
- HHV-7 infection. *See* Human herpesvirus-7 (HHV-7) infection
- HHV-8 infection. *See* Human herpesvirus-8 (HHV-8) infection
- Hiatal hernia, 1902, 1902f
- hic (pneumococcal H inhibitor), 947
- Hidradenitis, neutrophilic eccrine, 366, 380
- Hidradenitis suppurativa, 1099
- HIDS (hyperimmunoglobulinemia D with periodic fever syndrome), 2213t, 2215
- HIF-1 (hypoxia-inducible factor-1), 247
- HIF (hypoxia-inducible factor) 1 α , 333e-4, 392
- High-altitude cerebral edema (HACE)
 clinical features of, 247, 476e-1
 epidemiology of, 476e-1
 pathophysiology of, 476e-1–2, 476e-2f
 prevention of, 476e-2
 risk factors for, 476e-1
 treatment of, 476e-2–3, 476e-2t
- High-altitude illness, **476e-1**
 acute mountain sickness. *See* Acute mountain sickness (AMS)
 cerebral edema. *See* High-altitude cerebral edema (HACE)
 chronic mountain sickness, 248, 476e-5
 cough, 476e-4
 epidemiology of, 476e-1
 gastrointestinal, 476e-4
 genetic considerations in, 476e-1
 hypoxia in, 248
 neurologic events, 476e-4–5
 physiology of, 476e-1
 polycythemia, 399
 preexisting medical issues and, 476e-5
 psychological/psychiatric problems, 476e-5
 pulmonary edema. *See* High-altitude pulmonary edema (HAPE)
 pulmonary hypertension, 476e-6
 sleep impairment, 476e-4
- High-altitude pulmonary edema (HAPE)
 clinical features of, 476e-3, 476e-3f
 pathophysiology of, 47e-4, 247, 476e-3–4
 prevention of, 476e-4, 1764
 risk factors for, 476e-3
 treatment of, 476e-2t, 476e-4, 1764
- High-amplitude propagated contractions, 265

- High-density lipoprotein (HDL)
atherosclerosis and, 291e-2, 291e-6
composition of, 2436-2437, 2436f, 2436t
low levels of, 291e-6, 2445-2446, 2451. *See also* Lipoprotein disorders
in metabolic syndrome, 2454
metabolism of, 2437-2438, 2438f
very high levels of, 2446
- High endothelial venules (HEVs), 372e-26
- High-frequency ventilation, 1738, 1739t, 1742.
See also Mechanical ventilation
- Highly active antiretroviral therapy (HAART).
See Combination antiretroviral therapy (cART)
- High-molecular-weight kininogen deficiency, 733t
- High-sensitivity C-reactive protein (hs-CRP), in HIV infection, 1249t
- Hinchey classification, of diverticulitis, 1972, 1972f
- Hip(s)
bursitis of, 2222, 2222f, 2247
fracture of
 bisphosphonates for prevention of, 2499-2500, 2500f
 denosumab for prevention of, 2501, 2502f
 in elderly women, 6e-4
 epidemiology of, 2488-2489, 2489f
 lifetime cumulative risk of, 27t
 management of, 2495
 PPIs and, 1921
 risk screening for, 28
 osteoarthritis of, 2228, 2230
 pain in, 112, 2222, 2222f, 2230
- Hip replacement, 2233
- Hirata's syndrome (insulin autoimmune syndrome), 2348
- Hirsutism, **331**
after menopause, 332
approach to the patient, 332, 334f
drug-induced, 331t, 332, 380
etiology of, 331, 331t
hormonal evaluation in, 332-333
in 21-hydroxylase deficiency, 2355
prevalence of, 2254t
scoring scale for, 332, 333f
screening/testing for, 2254t
treatment of, 334-335
- Hirudin, 2747
- His bundle, 273e-1, 1471
- His bundle electrogram, 1472f, 1474
- Histamine, 88f, 372e-15t
- Histamine H₁ receptor antagonists. *See* Antihistamines
- Histamine H₂ receptor antagonists
adverse effects of, 1920
for angioedema, 2119
drug interactions of, 42t, 246e-3t
for gastritis, 277
for GERD, 263, 1908
for peptic ulcer disease, 1920, 1920t
for premedication in chemotherapy, 1798
for systemic mastocytosis, 2121
for urticaria, 2119
for ZES, 568
- Histidine metabolism disorders, 434e-2t
- Histidinemia, 434e-2t
- Histidinuria, 435e-1t
- Histiocytosis, 135e-9, 441e-16f
- Histiocytosis X (Langerhans cell histiocytosis), 135e-9, 1713, 2257, 2264
- Histone(s), 102e-7, 430, 431f
- Histone acetylase, 102e-7
- Histone deacetylase (HDAC), 102e-7, 102e-8f
- Histone deacetylase-2, 1677, 1701
- Histone deacetylase (HDAC) inhibitors, 102e-7-8, 102e-8f, 103e-18t, 103e-19-20
- Histone H2A-H2B-DNA, 372e-24t
- Histoplasma capsulatum*, 145e-3, 1330, 1332, 1332f, 1333f
- Histoplasma* spp. infections, **1332**
clinical features of, 1333
 arthritis, 836, 837f
 cavitary, 1333
 chronic meningitis, 908t
 cutaneous, 366
 laryngitis, 234
 lymphadenopathy, 408, 1333
 mediastinitis, 1333
 oral, 239t
 subacute meningitis, 898-899
 complications of, 1333
 diagnosis of, 152e-4t, 1332f, 1333, 1333f
 disseminated
 clinical course of, 1333
 in HIV infection, 1252t, 1255, 1268
 treatment of, 1334, 1334t
 epidemiology of, 1332
 incubation period for, 152e-4t
 pathogenesis of, 1332
 of pons, 441e-6f
 prevention of, 152e-4t
 in returning war veterans, 152e-4t
 in transplant recipient, 925t
 treatment of, 152e-4t, 899, 1333-1334, 1334t
- Histotoxic hypoxia, 248
- HIV
adherence of, 145e-1, 145e-2t
antibodies to, 1243-1244, 1243f
binding and fusion with target cell, 1216-1217, 1218f
cell membrane interactions of, 214e-1, 214e-3
cellular targets of, 1236
gene expression and replication in, 214e-4
genome of, 1218-1219, 1219f
genotypes of, 773, 774
geographic distribution of, 1220f
molecular heterogeneity of, 1219-1220
mutations in, 214e-5
persistent infections, mechanism of, 214e-7
phylogeny of, 1216, 1216f
receptors and co-receptors of, 1234-1235, 1236f
replication cycle of, 1216-1218, 1218f
resistance to, 145e-4
structure of, 225e-1-2, 225e-2f, 1216, 1217f
subtypes of, 1216, 1216f
viral release of, 145e-9
- HIV-1, 1216, 1216f, 1219-1220, 1219f, 1220f. *See also* HIV; HIV infection
- HIV-2, 1216, 1216f
- HIV-associated dementia (HAD), 1263, 1264t, 1265f
- HIV-associated neurocognitive disorders (HAND), 1241, 1263, 1264t, 1265f
- HIV infection
acute syndrome, 1228, 1249, 1249f
arthritis syndromes in, 837
asymptomatic stage (clinical latency), 1231, 1249-1250
cardiovascular disease in, 290e-1t, 1255, 1560, 1574
as chronic disease, 13-14
cobalamin deficiency in, 645-646
depression in, 2714
diagnosis of, 150e-7-8, **1245**
 algorithm for, 1245-1246, 1246f
 bDNA assay, 1247, 1247t
 enzyme immunoassay, 1245
 NABSA, 1247, 1247t
 p24 antigen capture assay, 770, 1243f, 1246-1247, 1247t
 point-of-care tests, 1246
 RT-PCR, 1247, 1247t
 Western blot, 1245, 1246f
drug allergies in, 384, 1260
drug interactions in, 798
drug resistance in, 773
drug toxicity in, 2030
elite nonprogressor, 1250
endocrine and metabolic disorders in
 adrenal insufficiency, 310
 hypogonadism, 1260, 2364
 lipodystrophy, 2450
 metabolic syndrome, 1259
 osteonecrosis, 1261
 osteoporosis, 1261
- SIADH, 1259
- thyroid disorders, 1259-1260
- epidemiology of
 global, 8, 12, 869, 1219-1220, 1220f, 1223-1225, 1224f
 in infants and children, 1226, 1226f, 1227f
 racial/ethnic disparities in, 1225-1226, 1226f, 1227f
 in U.S., 12, 1225-1227, 1225f, 1226f, 1227f
- gastrointestinal disease in, **1256**
 diarrhea, 266, 1257, 1257f
 esophagitis, 1256, 1256f, 1909-1910, 1910f
 HIV enteropathy, 1257
 infections, 1256-1257
 proctocolitis, 266
- generalized wasting in, 1268
- genetic considerations in
 abacavir-associated hypersensitivity, 1241
 chemokine receptors, 1239, 1240t
 chemokines, 1240t
 cytokines, 1240t
 gene-gene interactions, 1241t
 MHC locus genes, 1239
 phenotypes of susceptibility and response, 1238-1239
 specific conditions and, 1241, 1241t
- genitourinary tract infections in, 1258-1259
 chancroid, 1012
 gonococcal, 1006-1007
 syphilis, 1138, 1138t
 warts, 1198
- health care workers and, 918, 1222-1223, 1283-1284
- hearing loss in, 219
- hematopoietic system disease in
 anemia, 1262
 bone marrow suppression, 1261-1262, 1261t
 lymphadenopathy, 408, 1232, 1270
 monoclonal gammopathy of undetermined significance, 1262
 monocyte dysfunction, 422
 neutropenia, 1262
 thrombocytopenia, 727, 728, 1262
 VTE, 1262
- in hemophiliacs, 734, 735
- hepatobiliary disease in, 1258
 cholangiopathy, 285, 2086
 drug-induced, 2030
 hepatitis, 1258, 1273, 2039, 2048-2049
 liver transplantation for, 2069
 pancreatitis, 1258, 2097-2098
- hyperkalemia in, 64e-2, 310
- idiopathic CD4+ T lymphocytopenia in, 1272
- immune reconstitution inflammatory syndrome in, 205e-1, 1112-1113, 1254, 1261, 1261t
- immune response to, **1242**, 1243f
 cell-mediated, 1243t, 1244-1245
 humoral, 1243-1244, 1243f, 1243t, 1244f
- immunizations in, 788f, 798, 1252-1253t
- immunologic and rheumatologic disease in, 1260-1261
- in infants and children, 1226, 1226f
- KIRs with, 372e-15t
- laboratory monitoring of
 CD4+ T cell counts, 1247-1248, 1248f
 co-receptor tropism assays, 1248-1249
 D-dimer, 1249t
 HIV resistance testing, 1248
 HIV RNA determinations, 1248
 hs-CRP, 1249t
 IL-6, 1249t
- long-term nonprogressor, 1250
- long-term survivor, 1231
- natural history of, 771f
- neoplastic disease in
 anal cancer, 544, 1198, 1271-1272
 cervical cancer, 1198
 HPV-related, 1198
 invasive cervical cancer, 1271-1272
 Kaposi's sarcoma. *See* Kaposi's sarcoma
 lymphomas, 1270-1271, 1271f
 multicentric Castleman's disease, 1271

- HIV infection, neoplastic disease in (*Cont.*):
 primary central nervous system lymphoma, 601
 primary effusion lymphoma, 135e-2
 neurologic disease in, 1263, 1263t
 aseptic meningitis, 1264
 Chagas disease, 1265
 chronic meningitis, 907, 908t, 911
 cryptococcal meningitis, 1264, 1341
 distal sensory polyneuropathy, 1267, 2684
 focal neurologic deficits, 1265
 HIV-associated dementia, 1263, 1264t, 1265f
 HIV-associated neurocognitive disorders, 1241, 1263, 1264t, 1265f
 inflammatory demyelinating polyradiculoneuropathy, 2685
 meningitis, 892–893, 899
 mononeuropathies, 2685
 myelopathy, 1266–1267
 myopathy, 1266–1267
 pathogenesis of, 1241–1242, 1263–1264
 peripheral neuropathy, 1267, 2684–2685
 polyradiculopathy, 2685
 progressive multifocal leukoencephalopathy, 899–900, 1266
 seizures, 1265, 1265t
 sensory neuronopathy/ganglionopathy, 2685
 stroke, 1266
 toxoplasmosis, 1265–1266, 1266f, 1401–1402, 1402f
- ophthalmologic disease in, 1267
 choroid lesions, 1267–1268
 CMV retinitis, 215e-7–8, 1192, 1267
 progressive outer retinal necrosis, 1267
 opportunistic/disseminated infections in
 actinomycosis, 1091
B. dermatitidis, 1338
Bartonella, 366, 1082, 1083f, 1268
 CMV, 1192, 1193
Cryptococcus, 1264, 1341
Cryptosporidium, 1407
Histoplasma, 1255, 1268
 incidence of, 1250, 1253f
 JC virus, 1266
Leishmania, 798, 1268, 1388, 1391
 malaria, 798, 1268, 1374
Nocardia, 1084
 nontuberculous mycobacterial, 1129
P. aeruginosa, 1047
P. marneffeii, 1268, 1354, 1354t
 prophylaxis for, 1251–1253t
T. cruzi, 1266
T. gondii, 1265–1266, 1266f, 1401–1402, 1402f, 1403, 1404, 1561
T. whipplei, 1091
- oropharyngeal disease in, 241t, 1255
 aphthous ulcers, 237, 1255–1256, 1256f
 dysphagia, 257
 hairy leukoplakia, 1188, 1188f, 1255, 1256f
 linear gingival erythema, 236
 parotid enlargement, 241
 periodontitis, 236
 pharyngitis, 231
 during primary infection, 238t
 sicca syndrome, 2167t
 thrush, 1255, 1256f
 pathogenesis of, 1227
 advanced disease, 1231
 apoptosis in, 1234
 autoimmune phenomenon in, 1234
 B cells defects in, 1237–1238
 CD4+ T cell defects in, 1236–1237
 CD8+ T cell defects in, 1237
 cellular targets, 1236
 chronic infection
 clinical latency vs. microbiologic latency, 1231
 immune system evasion, 1229–1230
 persistent virus replication, 1229, 1229f
 reservoirs of HIV-infected cells, 1230, 1230f
 viral dynamics, 1230–1231, 1231f
 clinical course in untreated patient, 1228f
 in CNS, 1241–1242
 cytokine network in, 1234–1235
 dendritic cell defects in, 1238
 immune activation, inflammation and, 1233–1234, 1233t
 in long-term nonprogressors, 1231–1232
 in long-term survivors, 1231–1232
 lymphocyte turnover in, 1235
 lymphoid organs and, 1232–1233, 1232f, 1233f
 macrophage defects in, 1238
 monocyte defects in, 1238
 NK cell defects in, 1238
 primary infection, 1228–1229, 1229f
 viral receptors and co-receptors in, 1234–1235, 1236f
- postexposure prophylaxis, 918, 1283
 in pregnancy, 50, 1223
 prevention of, 1284–1285
 renal disease in, 62e-1f, 1258–1259, 1849, 1864
- respiratory disease in, 1250
 bacterial pneumonia, 1250
 fungal infections, 1254–1255
 MAC infections, 1251t, 1254
Pneumocystis pneumonia (PCP), 1250, 1251t, 1253, 1358, 1361
 pulmonary hypertension, 1255, 1658
R. equi infection, 1254
 tuberculosis. *See* Tuberculosis (TB), in HIV infection
- in returning war veterans, 152e-6t, 152e-8
 screening for, 24, 24f, 29t, 883, 1245, 1284–1285
- sepsis/septic shock in, 1757t
 skin manifestations of, 1262
 bacillary angiomatosis, 365, 1082, 1083f, 1268
 condyloma acuminatum, 1263
 cutaneous drug reactions, 377, 384, 1263
 folliculitis, 1262–1263
 herpes zoster, 1185, 1185f, 1263
 HSV infections, 1263
 perirectal, 1257–1258, 1258f, 1263
 treatment of, 1181, 1182t, 1263
- ichthyosis, 1263
 molluscum contagiosum, 220e-1, 1263
 onychomycosis, 350
 in primary infection, 25e-2f, 127, 362, 362t
- psoriasis, 1263
 seborrheic dermatitis, 347, 1262
 skin cancer, 500
- stage 3. *See* AIDS
 staging, 1215–1216, 1216t
 symptomatic disease, 1250
 thyroid hormone levels in, 2300
 transmission of
 in blood and blood products, 138e-3t, 138e-6, 735, 1220t, 1222–1223
 by breast-feeding, 50, 1223
 by human bite, 1220t, 1223
 mother-to-child, 50, 1223
 occupational, 1222–1223
 patterns, 1226–1227, 1226f
 prevention of, 1284–1285
 in saliva, 1220t, 1223
 sexual, 1220–1221, 1220t, 1221f
- traveler with, 798
 treatment of
 combination antiretroviral therapy. *See* Combination antiretroviral therapy (cART)
 with HCV co-infection, 2048–2049
 HSCT, 372e-31, 1239
 initial evaluation, 1272–1273, 1273t
 Internet resources for, 1273t
 milestones in, 771f
 in pregnancy, 50
 principles of, 1281–1283, 1282t
 vaccine development for, 1284
 weakness in, 156
 in women, 6e-2f, 6e-4
- H⁺K⁺-ATPase, 372e-24t, 1913–1914
 HK (hexokinase) deficiency, 654t
 HLA-B gene, 1239, 1240t
 HLA-B8 gene mutations, 2346
 HLA-B locus, 38, 39t, 41, 373e-1, 373e-7, 378.
See also Human leukocyte antigen (HLA) complex
 HLA-C gene, 1240t, 1241
 HLA complex. *See* Human leukocyte antigen (HLA) complex
 HLA-DR3 gene mutations, 2323, 2346
 HLA-DR4, 2202
 HLA-DRB1 gene, 2139
 HMBS gene, 2523t
 HMBS gene mutations, 2526, 2529
 HMB synthase, 2523t, 2524
 hMG (human menopausal gonadotropin), 337
 HMG-CoA reductase inhibitors. *See* Statins
 hMLH1 gene mutations, 539, 554
 HMPV (human metapneumovirus) infections, 1202t, 1206–1207
 hMSH2 gene mutations, 539, 554
 HNF gene mutations, 431t, 2406
 HNPCC. *See* Hereditary nonpolyposis colon cancer (HNPCC)
 HNPP (hereditary neuropathy with liability to pressure palsies), 2678, 2679t
- Hoarseness
 with dysphagia, 256
 in GERD, 262
 in laryngeal cancer, 503
 in laryngitis, 234
 in lung cancer, 510
- Hockey puck sign, 1014
 Hodgkin's lymphoma, 708
 approach to the patient, 700
 classical, 708–809
 classification of, 696t
 clinical features of, 708
 diagnosis of, 708–709
 EBV infection and, 214e-7, 697, 1189
 epidemiology of, 697
 fever in, 126
 infections in, 484t, 486t, 489
 lymphocyte-depleted, 708
 vs. mediastinal B-cell lymphoma, 135e-2
 mixed cellularity, 697, 708, 709f
 nodular lymphocyte-predominant, 709
 nodular sclerosing, 81e-11f, 697, 708
 paraneoplastic syndromes in, 612, 612t, 617, 618
 peripheral blood smear in, 81e-11f
 during pregnancy, 124e-1t, 124e-4
 radiation therapy for, breast cancer following, 103e-4, 524, 623
 second malignancies in survivors of, 623
 skin manifestations of, 366
 splenomegaly in, 410
 staging of, 413, 700, 701t, 709
 treatment of, 709
 brentuximab vedotin, 102e-4, 102e-5t, 103e-22
 HSCT, 139e-5t, 139e-6
 late consequences of, 623, 624t, 709, 2365
 tumor markers in, 473t
 “Holiday heart,” 2725
- Hollenhorst plaques, 40e-4, 201, 201f, 1443
 Holobiont, 86e-1, 86e-1t
 Holter monitoring, 273e-5
 Holt-Oram syndrome, 431t, 1443
 Homan's sign, 1443
 Home health care, 83, 83t
 Homeless, mental health problems in, 2723
 Homeopathy, 14e-1t, 14e-3
 Homeostasis, 400e-5
 Homer-Wright rosettes, 602
 Homocitrullinuria, 434e-3t, 435e-1t
 Homocysteine, 96e-4, 647
 Homocystinuria(s), 434e-1
 cardiovascular disease in, 643
 classic, 434e-1
 clinical features of, 434e-2t, 434e-4
 enzyme defects in, 434e-2t
 folate deficiency in, 643, 647

- pathways, enzymes, and coenzymes involved in, 434e-4, 434e-4f
treatment of, 434e-4
- Homogeneous staining region, 101e-3
- Homogentisic acid oxidase, 434e-4
- Homoplasmy, 85e-2-3, 85e-7-8
- Homozygous, 433f, 435
- Honeybee sting/venom, 2116, 2749
- Hookworm infections
clinical features of, 1414, 1414t
diagnosis of, 245e-3t, 245e-5t, 245e-6t, 765t, 1414-1415
eosinophilia in, 245e-6t, 1687
epidemiology of, 245e-3t, 765t, 1414
life-cycle hosts of, 245e-3t, 1414
skin manifestations of, 829
in travelers, 797
treatment of, 1414t, 1415
- Hoover's sign, 1704
- Hordeolum (sty), 199
- Horizontal gene transfer, 769t
- Hormesis, 94e-7
- Hormiga veinticuatro*, 2749
- Hormone(s), **400e-1**. *See also specific hormones and types*
age-related changes in levels of, 75t, 94e-4
autocrine control, 400e-6
classes of, 400e-1
deficiency of, 2252, 2252t
degradation of, 400e-3
ectopic production of, 608-612, 609t
eutopic production of, 608
excess, 2251-2252, 2252t
feedback regulatory systems, 400e-5-6, 400e-5f
functions of, 400e-4-5
half-life of, 400e-2
measurements of, 2253
paracrine control, 400e-6
resistance to, 2252, 2252t
role of, 2251
secretion of, 400e-2
serum-binding proteins and, 400e-2
stimulation tests, 2253
suppression tests, 2253
synthesis and processing of, 400e-2
transport of, 400e-2-3
unbound, 400e-2-3
- Hormone receptor(s)
characteristics of, 400e-1-2
families of, 400e-1-2, 400e-1t
membrane receptors, 400e-3-4, 400e-3f
nuclear receptors, 400e-4, 400e-4f
- Hormone receptor-directed therapy, 102e-7, 103e-5, 103e-15-16. *See also specific drugs*
- Hormone replacement therapy
for hypopituitarism, 2257, 2259t
postmenopausal. *See* Postmenopausal hormone therapy
- Hormone resistance syndromes, 2252
- Horner's syndrome
in cervical cord lesions, 2651
in lung cancer, 510, 519
ocular manifestations of, 196, 208
- Horse botfly, 2746
- Horse chestnut, 404t
- Hospice care, 56, 70, 103e-8. *See also* End-of-life care
- Hospital(s)
medical care in, 5
racial/ethnic disparities in care by, 16e-2f
- Hospital-acquired pneumonia, 767t, 813, 1015, 1020, 1028
- Hospital Compare* (website), 12e-4
- Hospitalists, 5
- Hospital Quality Initiative, 12e-3-4
- "Hot cross buns sign," 2641, 2641f
- "Hot potato" voice, 233
- Hot-tub folliculitis, 131t, 134, 356, 829, 1047
- Hot-tub lung, 1129, 1681, 1681t
- House dust mite, in asthma, 1669, 1671
- Howell-Jolly bodies
in asplenia, 2105
laboratory identification of, 81e-1, 81e-5f, 395f
- postsplenectomy, 413
in splenomegaly, 410
- Howship's lacunae, 2456, 2472
- HoxB4, 89e-4
- Hox genes, 89e-4
- Hoyeraal-Hreidarsson syndrome (dyskeratosis congenita), 102e-3, 360-361, 375e-1, 664, 2109
- HPC1* gene mutations, 101e-6t
- HPRT (hypoxanthine phosphoribosyltransferase), 431e-2
- HPRT (hypoxanthine phosphoribosyltransferase) deficiency, 431e-3t, 431e-5
- HPT-JT syndrome. *See* Hyperparathyroidism-jaw tumor (HPT-JT) syndrome
- HPV. *See* Human papillomavirus (HPV)
- HPV infections. *See* Human papillomavirus (HPV) infections
- HRAS* gene mutations, 101e-2t, 680
- H-reflex studies, 442e-5
- HRPT2* gene mutations, 2471-2472
- HRSV infections. *See* Human respiratory syncytial virus (HRSV) infections
- HSAN (hereditary sensory and autonomic neuropathy), 2643, 2679-2680t, 2680
- hs-CRP (high-sensitivity C-reactive protein), in HIV infection, 1249t
- HSCT. *See* Hematopoietic stem cell transplantation (HSCT)
- HSD3B2* gene mutations, 2353, 2354t, 2356t, 2357
- HSD17B3* gene mutations, 2353, 2354t
- HSV. *See* Herpes simplex virus (HSV)
- HSV infections. *See* Herpes simplex virus (HSV) infections
- 5-HT. *See* Serotonin
- hTERT, 102e-3
- HTLV-1. *See* Human T-cell lymphotropic virus (HTLV-1)
- HTLV-2 (human T-cell lymphotropic virus-2) infections, 225e-5-6
- Human African trypanosomiasis. *See* Sleeping sickness (human African trypanosomiasis)
- Human bite-wounds, **167e-2**, 167e-3t, 830
- Human bocaviruses, 1197
- Human chorionic gonadotropin (hCG)
in carcinoma of unknown primary, 120e-1, 120e-2t
ectopic production of, 608, 609t, 612
in germ cell tumors, 594
in gestational trophoblastic disease, 124e-5, 597
in nonfunctional pNETs, 571
reference values for, 2760t
in testicular cancer, 588
as tumor marker, 473t
- Human chorionic gonadotropin (hCG) stimulation test, 2361
- Human chorionic gonadotropin (hCG) therapy, 7e-4, 2369
- Human evolution, 85e-3
- Human foamy virus, 91e-1t
- Human genome, **425**. *See also* Chromosome(s); Gene(s)
databases/websites, 426t
replication of, 427, 428f
structure of, 425-427, 427f
- Human Genome Project, 425-426, 445
- Human granulocytic anaplasmosis
clinical features of, 1152, 1155t, 1160
coinfection with *Babesia*, 1385, 1386
diagnosis of, 1159f, 1160
epidemiology of, 1160
pathogenesis of, 1155t
prevention of, 1161
treatment of, 1161
- Human growth hormone. *See* Growth hormone (GH)
- Human herpesvirus-6 (HHV-6), 145e-2t, 1194
- Human herpesvirus-6 (HHV-6) infection, **1194**
in cancer patient, 492
clinical features of, 1194
diagnosis of, 1190t
- epidemiology of, 1194
exanthem subitum, 25e-2f, 127, 128t, 1194
reactivation in hypersensitivity syndrome, 382
in transplant recipient, 920t, 921t, 922, 1194
treatment of, 1194
- Human herpesvirus-7 (HHV-7) infection, **1194**
in cancer patient, 492
clinical features of, 1194
epidemiology of, 1194
in transplant recipient, 920t, 921t, 922, 928
treatment of, 1194
- Human herpesvirus-8 (HHV-8) infection, **1194**
in cancer patient, 492
clinical features of, 1194
epidemiology of, 1194, 1242
Kaposi's sarcoma and, 119e-1, 214e-7, 1194, 1242
lymphoid malignancies and, 214e-7, 697, 697t, 1194
primary effusion lymphoma and, 135e-2, 214e-7, 1194
in transplant recipient, 921t, 923, 928
treatment of, 1194
- Human immunodeficiency virus. *See* HIV
- Human leukocyte antigen (HLA) complex, **372e-1**
disease associations with, 372e-27t, 373e-6-7, 373e-8-9, 435
HLA class Ib genes, 373e-1
HLA class I genes, 373e-1-2, 373e-1f
biosynthesis of, 373e-3-4, 373e-4f
disease associations, 372e-27t, 373e-7, 373e-8t, 2109
functions of, 373e-4-5
HIV infection and, 1239, 1240t, 1241
NK cell recognition and, 373e-4-5
structure of, 373e-1f, 373e-3
HLA class II genes, 373e-1f, 373e-2
biosynthesis of, 373e-4f, 373e-5
disease associations, 372e-27t, 373e-7-8, 373e-8t, 2109
functions of, 373e-4f, 373e-5
HIV infection and, 1240t
structure of, 373e-1f, 373e-5, 373e-6f
linkage disequilibrium in, 373e-3
map of, 373e-1f
in transplantation, 139e-1, 373e-5-6, 1826-1827
- Human leukocyte surface antigens, 372e-1, 372e-3-4t
- Human menopausal gonadotropin (hMG), 337
- Human metapneumovirus (HMPV) infections, 1202t, 1206-1207
- Human microbiome, **86e-1**
alpha diversity in, 86e-4-5
Archaea in, 86e-4
bacteria in, 86e-2-3, 1094-1095, 1095t
beta diversity in, 86e-4
co-occurrence analysis of, 86e-5
culture-independent analysis of, 86e-2-4, 86e-3t
definition of, 86e-1t, 425
disease states and, 86e-7-9, 425
effects on host biology, 86e-9-10
enterotype analysis of, 86e-5-6
eukaryotes in, 86e-4
functional redundancy in, 86e-6, 86e-6f
interpersonal variation in, 86e-6f
neurologic disease and, 444e-6-7
research studies of, 86e-9-11
resiliency of, 86e-7
stability of, 86e-6-7
terminology related to, 86e-1t
viral dynamics in, 86e-4
xenobiotic metabolism in, 86e-9
- Human Microbiome Project, 145e-1
- Human monocytotropic ehrlichiosis, 129t, 1155t, 1159-1160
- Human papillomavirus (HPV)
cell membrane interactions of, 214e-3
gene expression and regulation in, 214e-4, 1197
structure of, 214e-2t, 214e-3f, 1197
types of, 595, 1197

- Human papillomavirus (HPV) infections, **1197**
 anal cancer and, 544, 1198
 cervical cancer and
 mechanisms of, 101e-7, 214e-7
 risk factors for, 351, 595
 types associated with, 595, 1198
 clinical features of, 1198–1199, 1199f
 counseling for, 1201
 diagnosis of, 150e-7
 epidemiology of, 1198
 global considerations in, 595, 1198
 in HIV infection, 1271–1272
 HIV infection and, 1198
 immune response to, 1198
 incidence of, 869
 oropharyngeal cancer and, 477, 502, 505, 1198
 pathogenesis of, 1197–1198
 in pregnancy, 124e-2
 prevention of, 1200, 1201. *See also* Human papillomavirus (HPV) vaccine
 in transplant recipient, 926
 treatment of, 1200–1201, 1201t
 warts, 240t, 351, 829, 1198, 1199f
- Human papillomavirus (HPV) vaccine
 for anogenital wart prevention, 351
 bivalent, 1199–1200
 for cervical cancer prevention, 595
 contraindications to, 789t
 cross-protection from, 1200
 effectiveness of, 476–477
 precautions for, 789t
 quadrivalent, 1200
 recommended administration of
 in adults, 29t, 787f, 788f, 883
 in children and adolescents, 883, 1200
 in HIV infection, 1253t
 in transplant recipient, 929t
 second-generation, 1200
 virus-like particles in, 1199
- Human respiratory syncytial virus (HRSV), 1205
- Human respiratory syncytial virus (HRSV) infections, **1205**
 in cancer patient, 489, 492
 clinical features of, 225, 229, 234, 1202t, 1206
 diagnosis of, 1206
 epidemiology of, 1205
 health care–associated, 1205
 in immunocompromised patient, 1205
 pathogenesis of, 1205–1206
 prevention of, 1206
 in transplant recipient, 923
 treatment of, 215e-2t, 1206
- Human stem cell leukemic-lymphoma syndrome. *See* *FGFR1*-mutated eosinophilia
- Human T-cell lymphotropic virus (HTLV-1) carcinogenicity of, 477t
 discovery of, 225e-2–3
 gene expression and regulation in, 214e-5
 molecular biology of, 225e-3–4
 structure of, 225e-1–2, 225e-3f
- Human T-cell lymphotropic virus-1 (HTLV-1) infections
 adult T-cell leukemia/lymphoma. *See* Adult T-cell leukemia/lymphoma
 arthritis syndromes in, 837
 clinical features of, 152e-6t
 epidemiology of, 152e-6t, 225e-4, 225e-4f
 global considerations in, 225e-4, 225e-4f
 lymphoid malignancies, 697
 myelopathy, 163, 225e-5, 697, 707, 2657
 prevention of, 225e-5
 in returning war veterans, 152e-6t
 transfusion-transmitted, 138e-3t, 138e-6
 treatment of, 225e-5
- Human T-cell lymphotropic virus-2 (HTLV-2) infections, 225e-5–6
- Humidifier fever, 1681t
 “Hummingbird sign,” 2610
- Humoral hypercalcemia of malignancy, 2476.
See also Hypercalcemia, malignancy-associated
- Hunger, 415e-2
 Hunner’s lesion, 60e-1
 Hunter disease, 432e-2t, 432e-6
 Hunt-Hess scale, for subarachnoid hemorrhage, 1785t
- Huntingtin, 453e-6, 2621–2622
- Huntington’s disease, **2621**
 clinical features of, 163, 184, 2606, 2621
 etiology of, 2621–2622
 functional anatomy of, 170
 genetic considerations in, 434, 440, 440t, 444e-9, 2621–2622
 genetic testing for, 448
 imaging in, 441e-23f, 2622f
 neuropathology of, 444e-10, 2622
 prevalence of, 2621
 treatment of, 444e-7, 2622
- Huntington’s disease–like disorders, 2622
- Hu proteins, 372e-25t
- Hurler disease, 432e-2t, 432e-6, 436e-5f
 Hurler-Scheie syndrome, 432e-2t, 432e-6
- Hurricanes, 151e-3, 151e-8
- HUS. *See* Hemolytic-uremic syndrome (HUS)
- Hutchinson-Gilford progeria syndrome, 94e-4, 435, 436t
- Hutchinson’s sign, 200
- Hutchinson’s teeth (incisors), 236, 1137
- HV interval, 1474
- Hyaline casts, 62e-11f
- Hyaluronan, 2140
- Hyaluronate lyase, 947f
- Hyaluronic acid, 2226, 2232, 2232t
- Hybrid capture, 150e-6, 150e-7f
- Hybridization, for pathogen identification, 772
- Hydatid disease. *See* Echinococcosis (hydatid disease)
- Hydatidiform mole, 124e-5, 372e-8t, 439.
See also Gestational trophoblastic disease
- Hydralazine
 adverse effects of, 40, 44, 253t, 363, 1625, 2218t
 genetic variations in response to, 39t, 40
 for heart failure, 1512t
 for hypertension, 1624t, 1625, 2644
 for hypertensive emergencies, 1627t
 metabolism of, 34t
 for preeclampsia, 46
- Hydrocele, 1418, 1418f
- Hydrocephalus
 adipsic diabetes insipidus in, 304
 after subarachnoid hemorrhage, 1785
 coma in, 1776
 communicating, 174, 175f
 diagnosis of, 163
 gait disorders in, 162t, 163
 in intracranial meningitis, 906
 normal-pressure, 175f, 2606–2607
 obstructive
 in neurocysticercosis, 1430–1431, 1431f, 1432
- Hydrochloric acid therapy, for metabolic alkalosis, 322
- Hydrochlorothiazide, 727t, 1623, 1624t, 2640t
- Hydrocodone, for dyspnea, 62t
- Hydrocortisone
 for ACTH deficiency, 2260
 for adrenal insufficiency, 2325–2326
 for adult hypopituitarism, 2259t
 for CAH, 2328–2329, 2356
 for candidiasis, 233
 for hypercalcemia, 314
 for hypoadrenal shock, 1751
 for IBD, 1960
 for myxedema coma, 2293
 for septic shock, 1755
 for thyroid storm, 2297
- Hydrogen chloride, as chemical terrorism agent, 262e-1t
- Hydrogen cyanide. *See* Cyanide poisoning
- Hydrogenotrophic species, in human microbiome, 86e-4
- Hydrogen peroxide (H₂O₂), in endothelium, 265e-1
- Hydrogen sulfide exposure/poisoning, 473e-11t, 1692t
- Hydroids, 2738
- Hydromorphone, 62t, 91t, 92
- Hydronephrosis, 1871
- Hydrophobia, 896, 1301, 1302f
- Hydros, 2081
- Hydrops fetalis
 clinical features of, 638, 1196
 diagnosis of, 1196f
 hemoglobin levels in, 638, 638t
 IgG antibodies in, 138e-1
- Hydrothorax, 251, 288, 1717
- Hydroureter, 1871
- 5-Hydroxytryptophan, 2623
- Hydroxocobalamin (vitamin B_{12A})
 for cobalamin deficiency, 648
 as cyanide antidote, 262e-7–8, 262e-8t, 473e-11t
- β-Hydroxybutyrate, 318, 2434
- 4-Hydroxybutyric aciduria, 434e-2t
- Hydroxychloroquine
 adverse effects of, 2130t
 common, 2146f
 cutaneous, 353, 380
 myopathy, 462e-20, 462e-20t, 2218t
 neuropathy, 2686, 2687t
 ocular, 206
 serious, 2146t
 for calcium pyrophosphate deposition disease, 2236
 for hypercalcemia, 314
 for infective endocarditis, 823t
 for malaria prophylaxis, 1383t
 monitoring during treatment with, 2146t
 overdose/poisoning with, 473e-13t
 for porphyria cutanea tarda, 390, 2530
 for Q fever, 1162
 for rheumatoid arthritis, 2145, 2146t
 for sarcoidosis, 2211, 2212t
 for SLE, 2130t, 2132
 for Whipple’s disease, 1094
- 2-Hydroxyglutarate (2-HG), 102e-13
- Hydroxykynureninuria, 434e-2t
- 7α-Hydroxylase, 1934
- 11β-Hydroxylase, 434
- 11β-Hydroxylase deficiency
 clinical features of, 306, 1620, 1620t, 2313, 2357
 diagnostic markers of, 2328t
 genetic considerations in, 1620t
 pathophysiology of, 1621f
- 17α-Hydroxylase (CYP17), 372e-24t
- 17α-Hydroxylase deficiency
 clinical features of, 306, 1620t, 1621, 2353, 2354t
 diagnostic markers of, 2328t
 genetic considerations in, 2353, 2354t
 genetics of, 1620t
 hypergonadotropic hypogonadism in, 337
 hypertension in, 1621
 pathophysiology of, 1621f
- 21-Hydroxylase (CYP21), 372e-24t
- 21-Hydroxylase antibodies, 2346
- 21-Hydroxylase deficiency
 classic. *See* Congenital adrenal hyperplasia (CAH)
 classic simple virilizing, 2355
 diagnosis of, 333
 nonclassic, 2355
 salt-wasting, 2355–2356
- 21-Hydroxylase gene mutations. *See* CYP21A2 (21-hydroxylase gene) mutations
- Hydroxyl radicals, 103e-4
- 3-Hydroxy-3-methylglutaric aciduria, 434e-3t
- 17-Hydroxyprogesterone, 2329, 2760t
- 3β-Hydroxysteroid dehydrogenase deficiency, 2328t, 2357
- 11β-Hydroxysteroid dehydrogenase, 64e-8, 64e-8f, 611, 2312
- 11β-Hydroxysteroid dehydrogenase deficiency. *See* Syndrome of apparent mineralocorticoid excess (SAME)
- 17β-Hydroxysteroid dehydrogenase deficiency, 2353

- Hydroxyurea
 action of, 103e-4, 103e-10t, 103e-14, 632
 adverse effects of, 103e-10t, 379, 649
 for chronic myelomonocytic leukemia, 135e-6
 for CML, 694
 for hyper eosinophilic syndromes, 423
 for sickle cell anemia, 636
- Hydroxyzine
 for angioedema, 2119
 for ciguatera poisoning, 2742
 for pruritus, 345
 for urticaria, 2119
- Hygiene hypothesis, 1670
- Hymen, imperforate, 335
- Hymenolepiasis diminuta*, 1434
- Hymenolepiasis nana* (dwarf tapeworm), 245e-2t, 246e-11, 765t, 1434
- Hymenoptera venom/sting, 2116, 2748–2749
- Hyoscyamine, 473e-8t, 2672
- Hypalgesia, 158
- Hyperabduction maneuver, 1647
- Hyperacute rejection, 1827
- Hyperaldosteronism
 diagnosis of, 307, 1618–1619
 familial type I. *See* Glucocorticoid-remediable aldosteronism (GRA)
 familial types II and III, 306
 genetic considerations in, 306, 307
 hyperreninemic, secondary, 321–322
 hypertension in, 1614, 1618
 hypokalemia in, 306
 idiopathic, 306
 metabolic acidosis in, 320–321
 mineralocorticoid excess in. *See* Mineralocorticoid excess
 primary, 306, 307, 1614, 1618–1619
 secondary, 306, 1614
 treatment of, 1619
 weakness in, 462e-119
- Hyperalgesia, 87, 158
- Hyperammonemia, 434e-3t, 434e-5, 435e-1t, 435e-2, 1996
- Hyperamylasemia, 2089t
- Hyperamylasuria, 2089t
- Hyperbaric medicine, 477e-1, 477e-1f
- Hyperbaric oxygen therapy, 477e-1
 for acute coronary syndrome, 477e-4t
 adverse effects of, 477e-2
 for altitude illness, 476e-2t, 476e-3, 476e-4
 for carbon monoxide poisoning, 477e-5–6
 for chronic wounds, 477e-3, 477e-4t, 477e-5, 477e-5f
 contraindications to, 477e-2
 for decompression sickness, 477e-4t, 477e-8
 for gas gangrene, 833, 994
 for idiopathic sudden sensorineural hearing loss, 477e-4t
 indications for, 477e-3, 477e-3t
 for late radiation tissue injury, 477e-3, 477e-4t
 mechanisms of, 477e-1, 477e-2f
 for radiation therapy enhancement, 477e-4t
 for soft tissue infections, 1102
 for traumatic brain injury, 477e-4t
- Hyperbilirubinemia, 2000
 in alcoholic liver disease, 2053, 2053t
 approach to the patient, 281–282, 281f
 conjugated
 in benign recurrent intrahepatic cholestasis, 284, 2003t, 2004
 differential diagnosis of, 282
 in Dubin-Johnson syndrome, 282, 2003–2004, 2003t
 with other liver abnormalities
 approach to the patient, 281f, 282–283
 in cholestatic disorders, 283–285, 284t
 in hepatocellular disease, 283, 283t
 in progressive familial intrahepatic cholestasis, 284, 2003t, 2004
 in Rotor syndrome, 282, 2003t, 2004
 in sepsis/septic shock, 1756
 isolated, 281–282, 282t
 in pancreatitis, 2093
 pathophysiology of, 280
 unconjugated
 from decreased hepatic bilirubin clearance, 2001
 differential diagnosis of, 281–282, 282t
 from hereditary defects. *See* Crigler-Najjar syndrome; Gilbert's syndrome
 from increased bilirubin production, 2000
- Hypercalcemia, 313, 2469
 in adult T-cell leukemia/lymphoma, 225e-4
 after kidney transplantation, 1831
 aluminum intoxication and, 2479
 asymptomatic, 2479
 bone turnover-associated, 2478
 chronic, 1862, 2479, 2479f
 in chronic kidney disease, 1815, 2478
 clinical features of, 96e-9t, 152, 313, 463e-4, 2469–2470
 diagnosis of, 314, 2469–2470, 2479f
 differential diagnosis of, 2469–2470, 2479–2480, 2479f
 ECG in, 269e-13f, 269e-14f, 1457, 1458f
 etiology of, 313, 313t, 2469, 2470t, 2480
 familial hypocalciuric. *See* Familial hypocalciuric hypercalcemia (FHH)
 genetic considerations in, 2475, 2476f
 in hyperparathyroidism. *See* Hyperparathyroidism, primary
 hypertension in, 1620
 hyperthyroidism and, 2478
 idiopathic, of infancy, 83e-6, 2477–2478
 immobilization and, 2478
 inpatient neurologic consultation for, 463e-4
 lithium therapy and, 2475
 malignancy-associated, 609
 clinical features of, 610, 2479–2480
 clinical syndromes related to, 2475–2477
 diagnosis of, 610, 2476–2477
 etiology of, 609–610, 609t
 mechanisms of, 2476
 pathogenesis of, 609t
 treatment of, 610, 2477
 in metastatic bone disease, 119e-4
 in milk-alkali syndrome, 313, 2458, 2479
 in multiple myeloma, 714, 717
 in nephrogenic diabetes insipidus, 303
 in Paget's disease of bone, 426e-3
 with parenteral nutrition, 98e-7t
 pNET and, 559t, 571
 renal effects of, 1862
 in sarcoidosis, 2208, 2477
 thiazides and, 2478
 treatment of, 314, 610, 2480–2482, 2480t
 vitamin A intoxication and, 2478
 vitamin D–related, 2477–2478
- Hypercalciuria
 in Bartter's syndrome, 306, 2483
 hematuria and, 294
 hypocalcemic, 2483
 loop diuretics and, 306
 nephrolithiasis risk and, 1867
 in Paget's disease of bone, 426e-3
- Hypercapnia, 315
 metabolic alkalosis following, 322
 permissive, 322, 1741
 in respiratory acidosis, 323
- Hypercholesterolemia. *See also* Lipoprotein disorders
 autosomal dominant type 2, 2442
 autosomal dominant type 3, 2442
 autosomal recessive, 2442
 familial, 436e-4f, 2440t, 2441–2442, 2449
 severe, 2446
- Hypercortisolism, 2271. *See also* Cushing's syndrome
- HyperC-VAD regimen, 704
- Hyperemesis gravidarum, 49, 259, 2300
- Hyper eosinophilic syndromes (HES)
 clinical features of, 135e-8, 1567, 1686, 2683
 diagnosis of, 135e-8, 135e-t
 diseases associated with, 422
 idiopathic, 422–423
 pathophysiology of, 372e-8, 1567
 subtypes of, 135e-7, 135e-7t, 1686
 treatment of, 135e-8, 1686
- Hyperesthesia, 158
- Hyperfiltration, glomerular, 333e-1
- Hyperfiltration hypothesis, 333e-1
- Hypergammaglobulinemia, 2050
- Hypergastrinemia, 568, 1927
- Hyperglobulinemia, 64e-6
- Hyperglycemia
 β -agonists and, 312
 in cardiogenic shock, 1760
 clinical features of, 2406
 in diabetic ketoacidosis. *See* Diabetic ketoacidosis (DKA)
 hyponatremia in, 303
 hyponatremia in, 301
 in overdose/poisoning, 473e-3
 in pancreatitis, 2092–2093
 with parenteral nutrition, 98e-7
 perioperative, 55
 in sepsis/septic shock, 1756
 with specialized nutritional support, 98e-4
 in systemic inflammatory response syndrome, 98e-4
- Hyperglycemic hyperosmolar state (HHS), 2417t, 2420
- Hypergonadotropic hypogonadism, 336–337
- Hyperhidrosis, primary, 2643
- Hyperhomocysteinemia
 cardiac manifestations of, 290e-1t, 290e-2
 clinical features of, 434e-4
 pyridoxine deficiency in, 96e-4
 risk factors for, 434e-4
 treatment of, 434e-4
 venous thrombosis risk in, 744
- Hyperhomocystinurias, 434e-1, 434e-4
- Hyperhydroxyprolinemia, 434e-2t
- Hyper-IgD syndrome, 126
- Hyper-IgE syndrome (Job's syndrome), 423
 autosomal dominant, 375e-2, 2109
 autosomal recessive, 2109
 cellular/molecular defects in, 420t, 423
 clinical features of, 375e-2, 420t, 423, 830, 2109
 diagnosis of, 420t, 2109
 monocytosis in, 421
- Hyper-IgM syndrome, 2110, 2110f, 2111
- Hyperimmune equine immunoglobulin, 1328
- Hyperimmunoglobulinemia D with periodic fever syndrome (HIDS), 2213t, 2215
- Hyperinflation, 47e-1, 1702
- Hyperinsulinemia, 98e-7, 332, 2405
- Hyperinsulinism, endogenous, 2434
- Hyperkalemia, 308
 in acute kidney injury, 1808
 after adrenalectomy, 1619
 AV conduction block in, 1471t
 in chronic kidney disease, 333e-2, 1814, 1815
 clinical features of, 310–311
 definition of, 308
 diagnosis of, 311, 311f
 drug-induced, 43, 64e-2, 310
 ECG in, 310, 1457, 1458f
 etiology of, 304, 308–310, 309t
 in HIV infection, 310
 inpatient neurologic consultation for, 463e-4
 metabolic acidosis in, 64e-2–3
 in overdose/poisoning, 473e-3
 with parenteral nutrition, 98e-7t
 rebound, 307
 renal effects of, 1862
 transfusion-related, 138e-5
 treatment of, 64e-3, 312, 463e-4
 in tumor lysis syndrome, 1795
 urinary tract obstruction and, 1872
 ventricular tachycardia in, 1489
- Hyperkalemic periodic paralysis (HyperKPP)
 channelopathies in, 444e-2t
 clinical features of, 310, 462e-3, 462e-17–18, 462e-17t
 diagnosis of, 462e-18
 early interventions for, 450t
 genetic considerations in, 444e-2t, 462e-17f, 462e-18

- Hyperkalemic periodic paralysis (HyperKPP) (*Cont.*):
 genetic testing for, 450t
 secondary, 310
 treatment of, 462e-18
- Hyperkinetic movement disorders, **2618**, 2619t
 chorea. *See* Chorea(s)
 drug-induced, 2624
 dystonia. *See* Dystonia
 hemiballismus, 2623
 myoclonus. *See* Myoclonus
 paroxysmal dyskinesias, 2624–2625
 restless legs syndrome. *See* Restless legs syndrome
 tics, 2623
 Tourette's syndrome, 2623
 tremor. *See* Tremor
- Hyperleucine-isoleucinemia, 434e-3t
- Hyperlipidemia. *See* Lipoprotein disorders
- Hyperlipoproteinemia, type III (familial dysbetalipoproteinemia), 2440t, 2443
- Hyperlysinemia, 434e-3t
- Hypermagnesemia, **2462**
 AV conduction block in, 1471t
 clinical features of, 463e-4, 2463
 etiology of, 2462–2463, 2463t
 inpatient neurologic consultation for, 463e-4
 treatment of, 2463
- Hypermetabolic state, **460**, 460t
- Hypermethioninemia, 434e-2t
- Hypernatremia, **303**
 clinical features of, 303
 diagnosis of, 303–304, 303f
 etiology of, 64e-3–4, 303, 463e-3
 free water deficit in, 64e-4
 hypodipsic. *See* Hypodipsic hypernatremia
 inpatient neurologic consultation for, 463e-3
 with parenteral nutrition, 98e-7t
 treatment of, 302t, 304
- Hyperopia, 195
- Hyperornithinemia, 434e-3t, 435e-1t
- Hyperosmolality, 463e-3
- Hyperostosis corticalis generalisata (van Buchem's disease), 426e-5
- Hyperoxaluria, 434e-2t, 1868, 1943
- Hyperoxic preconditioning, 477e-1
- Hyperparathyroidism
 anemia in, 630
 familial isolated, 2340
 global considerations in, 315
 muscle weakness in, 462e-19
 prevalence of, 2254t
 primary, **2470**
 asymptomatic, 2470, 2472, 2473t
 clinical features of, 2472–2473
 diagnosis of, 2472, 2473, 2473f, 2479
 etiology of, 2470
 genetic considerations in, 2471–2472, 2471f
 incidence of, 2470
 in MEN syndromes. *See entries at* Multiple endocrine neoplasia
 natural history of, 2470
 pathology of, 2470–2471
 treatment of, 2473–2475, 2473t
 screening/testing for, 2254t
 secondary, 314t, 1815, 2478–2479
 tertiary, 2479
- Hyperparathyroidism-jaw tumor (HPT-JT) syndrome
 clinical features of, 2342
 genetic considerations in, 2337t, 2342, 2470, 2472
 screening tests in, 2343t
 treatment of, 2342
- Hyperparathyroid-like syndromes, 2475, 2476f
- Hyperpathia, 89–90, 158, 2643
- Hyperperfusion states, 463e-1–2, 463e-1t, 463e-2f
- Hyperphagia, 2264
- Hyperphenylalaninemia, 434e-1, 434e-2t
- Hyperphosphatemia, **2460**
 in acute kidney injury, 1808, 1810
 in chronic kidney disease, 1815, 2485
 clinical features of, 2461
- etiology of, 2460–2461, 2461t
 hypocalcemia and, 2487
 with parenteral nutrition, 98e-7t
 severe acute, 2487
 treatment of, 2461
- tumoral calcinosis in, 426e-8
 in tumor lysis syndrome, 1795
- Hyperpigmentation, **359**
 in adrenal insufficiency, 2324, 2326f
 diffuse, 360–361, 360t
 drug-induced, 361, 379
 localized, 359–360, 360t
 in Whipple's disease, 1093
- Hyperplastic polyp, 537
- Hyperprolactinemia, **2266**
 clinical features of, 2266–2267
 cocaine use and, 469e-1
 diagnosis of, 328, 2267
 erectile dysfunction in, 326
 etiology of, 2266, 2266t, 2339
 galactorrhea and, 2267
 hirsutism in, 332
 hypogonadotropic hypogonadism and, 2365
 laboratory investigation of, 2267
 prevalence of, 2254t
 screening/testing for, 2254t
 treatment of, 2267
- Hyperproliferinemia, 434e-2t
- Hyperpyrexia, 124, 126
- Hyperreactive malarial splenomegaly, 1374–1375
- Hypersensitivity myocarditis, 1562
- Hypersensitivity pneumonitis, **1681**
 clinical features of, 1682
 clinical prediction rule, 1683
 definition of, 1681
 diagnosis of, 1682–1683, 1682f, 1683f, 1711t
 differential diagnosis of, 1683
 etiology of, 1681, 1681t, 1691
 global considerations in, 1683
 pathophysiology of, 1682
 treatment of, 1683
- Hypersensitivity reactions. *See also* Anaphylactic reactions
 to contrast agents. *See* Contrast agents
 delayed-type, 372e-29–30, 377–378, 1107
 drug-related. *See* Drug-induced hypersensitivity syndrome (DIHS); Drug-induced illness
 to human antibodies, 1796
 immediate-type, 372e-29, 2115
 to insect stings, 2749
 intermediate-type, 2115
 pathophysiology of, 2115–2116
 to transfusions, 138e-5
- Hypersensitivity vasculitis, 2191
- Hypersplenism, 411, 2064
- Hypertension, **1611**
 after kidney transplantation, 1831
 after stroke, 1627
 approach to the patient, 1621–1622
 deaths from, by sex, 6e-1t
 definition of, 1616, 1616t
 in diabetes mellitus, 2426, 2428
 diagnosis of, 1621–1622, 1621t
 diet and, 95e-5
 ECG in, 269e-15f
 epidemiology of, 1611–1612
 genetic considerations in, 1612, 1620t
 global considerations in, 266e-4
 high-altitude travel and, 476e-5
 history in, 1622, 1622t
 hypokalemic, 64e-8, 307, 1618, 2319
 isolated systolic, 1616, 1616t
 laboratory evaluation of, 1622, 1622t
 malignant. *See* Malignant hypertension
 mechanisms of
 arterial pressure, 1612, 1612f
 autonomic nervous system, 1613
 intravascular volume, 1612–1613
 renin-angiotensin-aldosterone system, 1613–1614, 1614f
 vascular, 1614–1615
 in metabolic syndrome, 291e-6, 1617, 2451–2452, 2454
- mineralocorticoid-mediated, 64e-8, 1614
 monogenic, 1620–1621, 1620t, 1621f
 obesity and, 1617
 oral contraceptives and, 2391t
 in paraganglioma, 1620
 paraneoplastic, 609t
 pathologic consequences of, **1615**
 cardiovascular, 266e-4, 291e-6, 1587
 cognitive dysfunction, 1615
 erectile dysfunction, 326
 heart failure, 1501, 1615. *See also* Heart failure (HF)
 peripheral artery disease, 1616
 renal, 62e-9f, 289t, 1615–1616, 1835t, 1847–1848
 stroke, 1615, 1627, 2582–2583, 2583t. *See also* Intracranial hemorrhage
 physical examination in, 201, 201f, 1444–1445, 1616, 1621–1622
 portal. *See* Portal hypertension
 in pregnancy, 45–46, 1620t, 1621
 primary (essential), 46, 1616–1617
 progesterone-induced, 2318t
 pulmonary arterial. *See* Pulmonary arterial hypertension (PAH)
 racial differences in complications of, 1612
 resistant, 1626
 risk factors for, 1612
 screening for, 29t
 secondary
 in acromegaly, 1620
 in aldosteronism, 1614, 1618–1619
 in coarctation of the aorta, 1620
 in Cushing's syndrome, 1619
 drug-induced, 1617t
 etiology of, 1617t
 in pheochromocytoma, 1613, 1619, 2329
 in renal disease, 1617
 in renovascular disease, 1617–1618
 in sleep apnea, 1619–1620
 in thyroid disease, 1620
 sodium intake and, 1612
 stroke and, 2567t
 supine, 146
 systolic with wide pulse pressure, 1616
 treatment of, **1622**
 ACE inhibitors, 1623, 1623t, 1625
 aldosterone antagonists, 1623, 1623t
 alpha blockers, 1623t, 1624
 ARBs, 1623, 1623t, 1625
 beta blockers, 1623t, 1624–1625
 blood pressure goals in, 1626
 calcium channel blockers, 1624t, 1625
 carotid sinus stimulation, 1625–1626
 catheter-based renal denervation for, 296e-5
 in chronic kidney disease, 1817
 in diabetes mellitus, 2428
 diuretics, 1623, 1623t, 1625
 lifestyle modifications, 1622–1623, 1622t
 in metabolic syndrome, 2454
 in polycystic kidney disease, 1853
 renal denervation, 1625–1626
 selection of, 1625–1626
 sympatholytics, 1623t, 1625
 vasodilators, 1623t, 1625
 in urinary tract obstruction, 1873
 vasoconstrictor use in, 242
 white coat, 1445, 1616
 in women, 6e-1t, 6e-4
- Hypertensive emergencies, 1626–1627, 1626t, 1627t. *See also* Malignant hypertension
- Hypertensive encephalopathy, 1615, 1626–1627, 2584
- Hyperthermia, 124, 138. *See also* Heat-related illness
- Hyperthyroidism
 clinical features of, 152
 alopecia, 355
 cardiac, 290e-1t, 290e-2, 1563
 cutaneous, 360, 363
 diarrhea, 855t
 edema, 253
 hypercalcemia, 2478
 hypertension, 1620

- hypokalemia, 64e-7, 305
 muscle involvement, 462e-18
 definition of, 2293
 in gestational trophoblastic disease, 597
 in Graves's disease. *See* Graves' disease
 in HIV infection, 1259
 HLA gene association with, 373e-8t
 in myasthenia gravis, 2703
 paraneoplastic, 609t
 in pregnancy, 48
 primary, 2293
 secondary, 2293
- Hyperthyroxinemia, euthyroid, 2286, 2286t
 Hypertrichosis, 332, 380, 390
 Hypertriglyceridemia. *See also* Lipoprotein disorders
 in metabolic syndrome, 2453-2454
 primary, 2439-2440, 2440t
 severe, 2446-2447
- Hypertrophic cardiomyopathy, **1568**
 apical, 1569
 clinical features of, 1554t, 1569
 dyspnea, 47e-4
 heart sounds/murmurs, 51e-2f, 51e-3, 51e-7, 1448, 1450
 syncope. *See* Cardiac syncope
 ventricular tachycardia, 1495t
 diagnosis of, 1569
 CMR, 270e-21, 270e-21f, 271e-7f, 271e-7v
 ECG, 269e-10f, 1569
 echocardiography, 271e-6f, 271e-6v, 1568, 1568f
 initial approach to, 1554t
 differential diagnosis of, 433e-5
 early interventions for, 449t, 450
 epidemiology of, 1568
 in Fabry's disease, 432e-1
 genetic considerations in, 436t, 437, 1555t, 1568
 genetic testing for, 450t
 incidence of, 1498
 mitral regurgitation in, 1543
 natural history of, 1439
 pathology of, 1568f, 1569f
 pathophysiology of, 1568-1569
 prognosis of, 1570
 sudden death and, 1498, 1569-1570, 1570t, 1767
 treatment of, 1498, 1569-1570, 1570f
 ventricular tachycardia in, 1498
- Hypertrophic osteoarthropathy, **2244**
 clinical features of, 426e-7, 2244-2245
 clubbing in, 250, 426e-7, 2244, 2244f, 2245
 diagnosis of, 250, 2245
 differential diagnosis of, 426e-7
 disorders associated with, 2245, 2245t
 familial, 2244-2245
 pathogenesis of, 250
 pathophysiology of, 2245
 treatment of, 2246
- Hyperuricemia, **431e-1**
 in AML, 681-682
 asymptomatic, 431e-4
 complications of, 431e-1, 431e-4
 definition of, 431e-1
 drugs causing, 431e-2t, 431e-3
 etiology of, 431e-1-3, 431e-2t, 431e-3f
 evaluation of, 431e-4
 in gout. *See* Gout
 in inborn errors of purine metabolism, 431e-3t
 metabolic syndrome and, 431e-4, 2452
 renal effects of, 1862
 treatment of, 431e-4-5, 2234-2235
 in tumor lysis syndrome, 1795
- Hyperuricosuria, 294, 431e-3t
 Hypervalinemia, 434e-3t
 Hyperventilation
 clinical features of, 1722-1723
 diagnosis of, 1723
 in respiratory alkalosis, 324
 in sepsis/septic shock, 1756
 treatment of, 1723
- Hyperventilation tetany, 479e-1
- Hyperviscosity
 in congenital heart disease, 1523
 in polycythemia, 399
 Raynaud's phenomenon in, 1649
 treatment of, 1523
- Hypervolemia, 295, 296f, 1808
 Hypesthesia, 158
 Hyphae, 1329
 Hypnic headache, 2598
 Hypnosis, 14e-1t
 Hypnozoites, *Plasmodium*, 1369
 Hypoactive sexual desire, female, 330
 Hypoadrenal shock, 1748t, 1750-1751
 Hypoalbuminemia, 252, 294, 298, 1997-1998
 Hypoaldosteronism
 etiology of, 310
 hyperkalemia in, 310
 hyponatremia in, 299
 hyporeninemic, 64e-2, 309t, 310, 320-321
- Hypoalphalipoproteinemia, 2445-2446
 Hypoaminoacidemia, 569
 Hypobetalipoproteinemia, familial, 2444
 Hypocalcemia, **314, 2482**
 in acute kidney injury, 1808-1809
 after parathyroidectomy, 2474
 chronic, 2482, 2487
 classification of, 2482, 2482t
 clinical features of, 96e-9t, 315, 463e-4, 463t, 1563, 2474, 2482
 diagnosis of, 315
 differential diagnosis of, 2487-2488
 ECG in, 269e-14f, 1457, 1458f
 etiology of, 314-315, 314t, 2483-2484
 genetic syndromes associated with, 2483
 global considerations in, 315
 hyperphosphatemia and, 2460, 2487
 inherited disorders, 332e-7t, 332e-8
 inpatient neurologic consultation for, 463e-4
 in pancreatitis, 2093, 2482
 with parenteral nutrition, 98e-7t
 pathophysiology of, 2482, 2491
 PTH absent. *See* Hypoparathyroidism
 PTH ineffective, 2482t, 2484
 in chronic kidney disease, 2484-2485
 with defective vitamin D metabolism, 2484
 in pseudohypoparathyroidism. *See* Pseudohypoparathyroidism (PHP)
 in vitamin D deficiency, 2484
 PTH overwhelmed, 2482t, 2487
 transfusion-related, 138e-5
 treatment of, 315, 2484, 2488
- Hypocalciuria, 306
 Hypocapnia, 315, 323
 Hypoceruloplasminemia, 451e-1t
 Hypochondria, 409
 Hypochromia, 81e-1, 393
 Hypocretin (orexin), 185, 186f, 189, 215
 Hypocupric myelopathy, 2658f
 Hypodipsic hypernatremia, **2279, 2279f**
 Hypoesthesia, 158
 Hypogammaglobulinemia
 bronchiectasis in, 1694, 1694t
 clinical features of, 270
 in CLL, 485, 486t
 disorders associated with, 375e-2
 immunoglobulin replacement for, 492
 infectious diarrhea in, 855
 in multiple myeloma, 486t, 492, 714
 in thymoma, 123e-1
- Hypogeusia. *See* Taste disorders
 Hypoglossal nerve, 2538, 2649
 Hypoglycemia, **2430**
 accidental, 2434
 approach to the patient, 2435
 in cancer, 1794-1795
 clinical features of, 2430-2431, 2431t
 in critical illness, 2433
 in diabetes mellitus, **2431, 2432f**
 diagnosis of, 2435
 drug-induced, 2433
 in elderly, 82
 endogenous hyperinsulinism and, 2434
 etiology of, 2430, 2430t
 factitious, 2434
- in heatstroke, 479e-3
 hormonal defense against, 400e-5
 hormone deficiencies and, 2433
 hypothermia in, 478e-1
 inborn errors of metabolism and, 2434-2435
 insulin-induced, 2260
 in insulinoma, 569
 with insulin plus glucose, 312
 in malaria, 1373, 1381
 non-beta-cell tumors and, 2433-2434
 in overdose/poisoning, 473e-3
 pathophysiology of, 2430-2431, 2431t
 perioperative, 55
 recurrent, prevention of, 2435
 syncope and, 147
 treatment of, 2435
 tumor-induced, 609t, 611-612
- Hypogonadism
 acquired testicular causes of, 2365-2366
 in cryptorchidism, 2365
 diagnosis of, 2367f
 erectile dysfunction in, 326
 in hemochromatosis, 2517
 in HIV infection, 1260
 hypergonadotropic, 336-337
 hypogonadotropic. *See* Hypogonadotropic hypogonadism
 in Klinefelter's syndrome, 2365. *See also* Klinefelter's syndrome
 prevalence of, 2254t
 screening/testing for, 2254t
- Hypogonadotropic hypogonadism, **2363**. *See also* Androgen deficiency
 acquired, 2364-2365
 amenorrhea in, 336
 clinical features of, 2260-2261
 congenital disorders associated with, 2363-2364, 2364t
 delayed puberty in girls and, 2380, 2381t
 diagnosis of, 2261
 familial, 2363
 hemochromatosis and, 2365
 hyperprolactinemia and, 2365
 idiopathic, 2363
 isolated, 336, 2260
 obesity and, 415e-3, 2364-2365
 pathophysiology of, 2363
 sellar mass lesions and, 2365
 treatment of, 2261, 2369
- Hypokalemia, **305**
 in aldosteronism, 1618
 arrhythmia in, 273e-3
 in chronic kidney disease, 1814
 clinical features of, 307
 diagnosis of, 307, 308f
 in DKA, 2419
 ECG in, 43, 269e-13f, 307, 1457, 1458f
 etiology of, 305-307, 305t
 in hyperthyroidism, 64e-7
 hyponatremia and, 302
 inpatient neurologic consultation for, 463e-4
 magnesium deficiency in, 64e-8, 306-307
 in mineralocorticoid excess, 2319
 in overdose/poisoning, 473e-3
 with parenteral nutrition, 98e-7t
 renal effects of, 1862
 treatment of, 64e-8-9, 307-308, 463e-4
- Hypokalemic periodic paralysis (HypoKPP)
 channelopathies in, 444e-2t
 clinical features of, 305, 462e-16-17, 462e-17t
 etiology of, 305
 familial, 305
 genetic considerations in, 305, 444e-2t, 462e-17, 462e-17f
 thyrotoxicosis and, 64e-7, 444e-2, 462e-16, 462e-18, 2294
 treatment of, 307
- Hypomagnesemia, **2461**
 after parathyroidectomy, 2474
 alkalosis and, 322
 arrhythmia in, 273e-4
 clinical features of, 463e-4, 463t, 2462
 etiology of, 2461-2462, 2462t
 familial hypercalcemic, 332e-7t, 332e-8

- Hypomagnesemia (Cont.):**
 genetic considerations in, 2461–2462
 hypocalcemia and, 2484
 hypokalemia and, 64e-8, 305, 306–307
 hypoparathyroidism and, 2484
 inpatient neurologic consultation for, 463e-4
 with parenteral nutrition, 98e-7t
 primary, 332e-7t
 treatment of, 64e-8–9, 307, 2462, 2484
 vitamin D deficiency and, 2462
- Hypomelanosis, 357, 357t, 358t**
Hypomelanosis of Ito (linear nevoid hypopigmentation), 357, 357t
- Hypometabolic states, 460**
 anemia in, 630–631, 630t
 gluconeogenesis in, 461
 metabolic rate in, 461
 physiologic characteristics of, 460–461, 460t
 protein catabolism in, 461
- Hyponatremia, 298, 2280**
 in acute kidney injury, 1808
 in adrenal insufficiency, 2324
 after subarachnoid hemorrhage, 300, 1785–1786
 chronic, 64e-5–6, 301, 303
 in chronic kidney disease, 1814
 clinical features of, 300–301, 2280
 diagnosis of, 299f, 301
 differential diagnosis of, 2281, 2282t
 etiology of, 301t, 332e-10f, 2280–2281
 euvoletic, 299–300, 2280, 2282t. *See also* Syndrome of inappropriate antidiuretic hormone (SIADH)
 exercise-associated, 300–301
 global considerations in, 2283
 hypervolemic
 differential diagnosis of, 2282t
 disorders associated with, 299, 2280
 pathophysiology of, 299, 2281
 treatment of, 302, 2282–2283
 hypokalemia and, 302
 hypovolemic
 differential diagnosis of, 301, 2282t
 disorders associated with, 299, 2280
 pathophysiology of, 298–299, 2281
 treatment of, 64e-5–6, 302
 inpatient neurologic consultation for, 463e-3–4
 with low solute intake, 300
 in lung cancer, 511
 paraneoplastic, 610
 with parenteral nutrition, 98e-7t
 pathophysiology of, 2281
 treatment of, 64e-5–6, 302–303, 463e-3–4, 2281–2283
- Hypoparathyroidism**
 acquired, 2483–2484
 APS-1 and, 2345, 2346t
 APS-2 and, 2347t
 chronic, 2483–2484
 clinical features of, 2482–2483
 etiology of, 314t, 2482t
 genetic syndromes associated with, 2483
 hyperphosphatemia due to, 2460
 hypocalcemia and, 314–315, 314t, 2482–2483
 muscle involvement in, 462e-19
 transient, 2484
 treatment of, 2484
- Hypoparathyroidism, deafness, and renal dysplasia (HDR) syndrome, 2483**
- Hypophosphatasia, 237, 426e-1, 426e-5–6**
Hypophosphatemia, 2458
 clinical features of, 96e-9t, 463t, 1563, 2459–2460
 etiology of, 2458–2459, 2459t
 in hospitalized patients, 461
 hypokalemia and, 64e-7
 in oncogenic osteomalacia, 612
 osteomalacia and, 2465. *See also* Osteomalacia with parenteral nutrition, 98e-7t
 treatment of, 2460, 2460t
- Hypophysectomy, for breast cancer, 530t**
Hypophysitis, 2257, 2346t, 2347t
- Hypopigmentation, 357**
 diffuse, 357
 etiology of, 357, 357t, 358t, 359
 linear nevoid, 357, 357t
 localized, 357
 postinflammatory, 358t, 359
- Hypopituitarism, 2255**
 acquired causes of, 2256–2257
 clinical features of, 2257
 developmental and genetic causes of, 2256–2257
 etiology of, 2255–2257, 2255t
 hypothermia in, 478e-1
 laboratory investigation of, 2257, 2258t
 oral manifestations of, 236
 treatment of, 2257, 2259t
- Hypoproliferative anemia, 398, 625, 630, 631, 662. See also** Aplastic anemia; Iron-deficiency anemia
- Hypoproteinemia, 253**
Hyposensitization. See Immunotherapy
- Hyposmia. See** Olfactory disorders
- Hypospadias, isolated, 2355**
Hyposplenism, 413. See also Splenectomy
- Hypotension**
 in coma, 1773
 in gastrointestinal bleeding, 277
 in heatstroke, 479e-4
 in hemodialysis, 1824
 in hypovolemia, 298
 intracranial, 2607
 in malabsorption syndromes, 1946t
 in mechanically ventilated patient, 1743
 orthostatic. *See* Orthostatic hypotension
 in sepsis/septic shock, 1755
 in shock, 1729–1730, 1731f, 1732f, 1744–1745
 transfusion-related, 138e-5–6
- Hypothalamic amenorrhea, 336**
- Hypothalamic disease**
 body temperature in, 124
 hypopituitarism due to, 2256. *See also* Hypopituitarism
- Hypopituitarism**
 IgG4-related, 391e-1t
 metabolic effects of, 2264
 obesity in, 415e-4
 sellar masses, 2264–2265
- Hypothalamic gliomas, 2264**
Hypothalamic hamartomas, 2264
- Hypothalamic-pituitary-adrenal (HPA) axis, 2310, 2310f**
- Hypothalamic-pituitary-gonadal axis, 335, 336f**
Hypothalamic-pituitary-testis axis, 2358, 2358f
Hypothalamic-pituitary vasculature, 401e-2f
- Hypothalamus**
 age-related changes in, 76
 developmental dysfunction of, 2256
 in ovarian function regulation, 2376–2377, 2377f
 in sleep onset, 185–186
- Hypothermia, 478e-1**
 bradyarrhythmia in, 1467
 clinical features of, 478e-1–2, 478e-2t
 coma and, 1773
 diagnosis of, 478e-2
 drug-induced, 478e-1, 478e-1t
 ECG in, 1457, 1458f
 in elderly, 478e-1, 478e-1t, 478e-3
 etiology of, 478e-1
 in myxedema coma, 2293
 physiologic changes in, 478e-2t
 primary accidental, 478e-1
 risk factors for, 478e-1t
 SA node dysfunction in, 1467t
 secondary, 478e-1
 in shock, 1751
 therapeutic, after cardiac resuscitation, 1770, 1781
 treatment of, 478e-3, 1751
- Hypothyroidism, 2289**
 autoimmune
 classification of, 2290
 clinical features of, 2290t, 2291, 2291f
 differential diagnosis of, 2291
 laboratory evaluation of, 2291
- pathogenesis of, 372e-24t, 377e-4–5, 2290–2291
 prevalence of, 2290
 clinical, 2290, 2292
 clinical features of, 152, 2290t, 2291, 2291f
 alopecia, 355
 anemia, 630
 ascites, 287
 bleeding, 404
 cardiac, 290e-1t, 290e-2
 AV conduction block, 1471t
 heart failure, 1563
 SA node dysfunction, 1467, 1467t
 carotenemia, 96e-7
 cutaneous, 363
 depression, 2714
 edema, 253
 hirsutism, 332
 hypertension, 1620
 hyponatremia, 299, 301
 hypothermia, 478e-1, 478e-3
 lipoprotein disorders, 2441
 muscle involvement, 462e-18
 neuropathy, 2683
 obesity, 415e-4
 congenital, 2283, 2284t, 2289–2290
 diagnosis of, 2291, 2292f
 differential diagnosis of, 2291
 diseases associated with, 2291
 etiology of, 2289t, 2292
 in HIV infection, 1259–1260
 iatrogenic, 2292
 overt, 2290
 pathogenesis of, 2290–2291
 in pregnancy, 48, 49, 2293, 2300
 prevalence of, 2254t, 2290
 radiation therapy-related, 506, 623
 screening/testing for, 2254t
 secondary, 2292
 subclinical, 2290, 2292–2293
 treatment of, 2292–2293
- Hypotonic saline, 304. See also** Saline
- Hypouricemia, 332e-7t, 431e-3t, 431e-5, 435e-1t**
Hypoventilation. See also Obstructive sleep apnea
 central hypoventilation syndrome, 1722, 1782
 clinical features of, 1720–1721, 1721t
 diagnosis of, 1721–1722
 hypoxia in, 247–248
 in obesity, 415e-6, 1722
 pathophysiology of, 1721f
 respiratory acidosis in, 322
 treatment of, 1722
- Hypovolemia, 297**
 in acute blood loss, 393
 in acute kidney injury, 1808
 AVP levels and, 295, 296f
 clinical features of, 298
 diagnosis of, 298
 endogenous response to, 1745
 etiology of, 297–298
 in myocardial infarction, 1607
 treatment of, 298
- Hypovolemic shock, 1748**
 clinical features of, 21t, 298, 393, 1748
 diagnosis of, 1748
 hemodynamic patterns in, 1762t
 initial assessment of, 1729
 physiologic characteristics of, 1748t
 predictors of fluid responsiveness in, 1729, 1731f, 1732f
 treatment of, 1749–1750
- Hypoxanthine phosphoribosyltransferase (HPRT), 431e-2**
Hypoxanthine phosphoribosyltransferase (HPRT) deficiency, 431e-3t, 431e-5
- Hypoxemia, 47e-5, 392, 1755**
- Hypoxia, 247**
 in acute kidney injury, 333e-5
 adaptation to, 248
 in angiogenesis, 102e-14, 102e-16f
 effects of, 247
 etiology of, 247–248
 SA node dysfunction in, 1467, 1467t

- Hypoxia-inducible factor (HIF)1, 247
 Hypoxia-inducible factor (HIF) 1 α , 333e-4, 392
 Hypoxic-ischemic encephalopathy, 1773, 1781-1782, 1781f, 2559
 Hy's Law, 2025
 Hysterectomy, 593, 597
- I**
- I-309/TCA-3, 372e-12t
 IA-2 (ICA512), 372e-24t
 IABP (intraaortic balloon pump), 296e-5, 1763
 IADLs (instrumental activities of daily living), 72t, 78t
 IAP (inhibitor of apoptosis proteins), 102e-9, 102e-9f
 IAPP (islet amyloid polypeptide), 2402
 Iatrogenic polydipsia, 2276, 2279
 I band, of sarcomere, 265e-5
 Ibandronate, 314, 2383, 2499, 2500f
 IBD. *See* Inflammatory bowel disease (IBD)
 I blood group system, 138e-1, 138e-1t
 IBMPPFD (inclusion body myopathy with Paget's disease and frontotemporal dementia), 426e-1
 IBPN (immune-mediated brachial plexus neuropathy), 2692
 Ibritumomab tiuxetan, 103e-19, 103e-22
 Ibrutinib, 703
 IBS. *See* Irritable bowel syndrome (IBS)
 Ibuprofen
 for acute idiopathic pericarditis, 1573
 advantages of, 1923
 adverse effects of, 727t, 2232t
 drug interactions of, 2232, 2232t
 for dysmenorrhea, 338
 for fever, 126
 for frostbite, 478e-4t
 for gout, 2234
 for migraine, 2592t
 for osteoarthritis, 2232t
 for pain, 60, 91t
 for postprandial orthostatic hypotension, 2644
 therapeutic monitoring of, 2765t
 Ibutilide
 action of, 273e-6t
 adverse effects of, 1480t
 for arrhythmias, 1479t, 1483, 1483f
 for atrial fibrillation, 1487
 dosage of, 1479t
 ICAM-1 (intercellular adhesion molecule 1), 1242, 2170
 ICD. *See* Implantable cardioverter-defibrillator (ICD)
 ICF (immunodeficiency with centromere instability and facial anomalies), 375e-2, 2109
 Ichthyosis, in HIV infection, 1263
 ICMAs (immunochemiluminometric assays), 2288
 Icodextrin, 1825
 Icosapent ethyl, 2448t
 ICOS gene, 2111
 ICP. *See* Intracranial pressure (ICP)
 Icterus, 1992
 Ictotest, 280
 ICU patient. *See* Critically ill patient
 ICU psychosis. *See* Delirium
 Idarubicin, 103e-10t, 103e-13, 682
 IDDM (insulin-dependent diabetes mellitus). *See* Diabetes mellitus, type 1
 Ideational apraxia, 179
 Idelalisib, 703
 Ideogram, 83e-1, 83e-1f
 Ideomotor apraxia, 179
 IDH1/IDH2 gene mutations, 102e-13, 599, 680, 680t
 Idiopathic CD4+ T lymphocytopenia, 1272
 Idiopathic cutaneous vasculitis, 2191-2192
 Idiopathic growth hormone deficiency (IGHD), 2258
 Idiopathic hypercalcemia of infancy (Williams' syndrome), 83e-6, 2477-2478
 Idiopathic pulmonary fibrosis. *See also* Interstitial lung diseases (ILDs)
 clinical features of, 1711
 histologic findings in, 1711
 imaging of, 308e-7f
 lung transplantation for, 320e-1t
 pathogenesis of, 1709f
 treatment of, 1712
 Idiopathic torsion dystonia, 2620
 Idiotypes, of immunoglobulins, 710
 Idioventricular rhythms, 1489, 1490f, 1494.
 See also Premature ventricular contractions (PVCs)
 IDLs (intermediate-density lipoproteins), 1241t, 2436-2437, 2436f, 2436t
 IFN- α . *See* Interferon- α (IFN- α)
 IFN- α 2a. *See* Interferon- α 2a (IFN- α 2a)
 IFN- α 2b. *See* Interferon- α 2b (IFN- α 2b)
 IFN- γ -inducible protein-10 (IP-10), 372e-12t
 IFN- λ 4 gene, 2011, 2042
 Ifosfamide
 action of, 103e-8
 adverse effects of, 103e-9t
 cardiotoxicity, 621, 1563
 cystitis, 622-623, 1798
 hypokalemia, 306
 nausea and vomiting, 103e-25
 nephrogenic diabetes insipidus, 303
 nephrotoxicity, 1804
 neurologic, 103e-8, 607, 607t, 1793
 pulmonary, 621, 1797
 for Ewing's sarcoma, 119e-4
 for osteosarcoma, 119e-3
 for soft tissue sarcoma, 119e-2
 for testicular cancer, 591
 Ig. *See* Immunoglobulin(s)
 IgA nephropathy
 clinical features of, 1835t, 1839-1840
 epidemiology of, 1839
 hematuria in, 294
 vs. IgA vasculitis, 1839
 pathogenesis of, 372e-9, 1839, 1840f
 renal biopsy in, 62e-3f, 1839
 treatment of, 1840
 IgA vasculitis (Henoch-Schönlein), 2190
 clinical features of, 368, 732, 1835t, 2190
 diagnosis of, 2190
 vs. IgA nephropathy, 1839
 incidence and prevalence of, 2190
 pathogenesis of, 732, 2190
 treatment of, 2190
 IGF. *See* Insulin-like growth factor (IGF)
 Ig Fc receptors, 372e-1, 372e-6, 372e-7
 IgG4-related disease (IgG4-RD)
 clinical features of, 391e-1-2, 391e-1t, 391e-2f, 1638, 1858, 2085
 epidemiology of, 391e-2
 pathology of, 391e-2-3, 391e-3f
 pathophysiology of, 391e-3
 serologic findings in, 391e-2
 treatment of, 391e-3
 IGHD (idiopathic growth hormone deficiency), 2258
 IGRAs (interferon- γ release assays), 1115, 1131
 IKBKAP gene mutations, 2643
 IKK (I κ B kinase), 102e-6
 iKNRs (inhibitory NK cell receptors), in HIV infection, 1238
 IL2RG gene, 91e-1
 IL28B gene mutations, 2011, 2042
 ILDs. *See* Interstitial lung diseases (ILDs)
 Ileal resection, 645
 Ileal stricture, 1884f
 Ileocecal syndrome. *See* Enterocolitis, neutropenic
 Ilesha virus, 1308t
 Ileus
 abdominal swelling in, 286
 etiology of, 1982, 1982t, 1985
 gallstone, 1985, 2081-2082
 hypokalemia and, 307
 postoperative, 92
 treatment of, 1985
 Iliopsoas bursitis, 2222, 2222f, 2247
 Iliotibial band syndrome, 2248-2249
 Iloperidone, 2722t
 Iloprost, 1659, 1660t
 IMAge syndrome, 2324t
 Imatinib
 action and targets of, 102e-3, 102e-5t, 103e-16f, 103e-17t
 adverse effects of, 103e-17t
 cardiac, 103e-26, 621
 common, 692
 hepatic, 103e-26
 myopathy, 2218t
 pulmonary, 1797
 serious, 692
 for CML, 101e-3, 102e-3, 102e-5t, 103e-16, 688, 691-692, 691t, 693
 cost of, 694-695
 drug interactions of, 1793
 for gastrointestinal stromal tumors, 102e-3, 102e-5t, 103e-19, 536
 genetic variations in response to, 39t, 41
 for hypereosinophilic syndromes, 1686
 resistance to, 102e-3-4, 102e-16, 102e-19, 103e-23, 689
 for small intestinal leiomyosarcoma, 537
 for soft tissue sarcoma, 119e-2
 for thymoma, 123e-2
 Imerslund-Gräsbeck syndrome, 645
 Imerslund's syndrome, 645
 IMF (International Monetary Fund), 8
 Iminoglycinuria, 435e-1t
 Imipenem
 adverse effects of, 942
 for anaerobic bacterial infections, 1101, 1101t
 for bite-wound infections, 167e-3t
 for *Campylobacter* infections, 1061
 for community-acquired pneumonia, 808t
 for enterococcal infections, 975t
 for febrile neutropenia, 103e-23, 491
 for glanders, 1045t
 for health care-associated pneumonia, 767t, 812t
 indications for, 940t, 941
 for intraabdominal infections, 767t
 for melioidosis, 1045t
 for multidrug-resistant TB, 205e-7
 for *Nocardia* infections, 1087
 for osteomyelitis, 840t, 841t
 for *P. aeruginosa* infections, 808t, 1045t
 for peritonitis, 848
 renal excretion of, 935
 resistance to, 933, 940t
 for septic shock, 767t
 Imipenem-cilastin, 727t, 937t, 1091t, 1757t
 Imipramine
 adverse effects of, 284, 2710t
 for depression, 2710t, 2716
 for detrusor spasticity, 2660
 overdosage/poisoning with, 473e-9t, 473e-10t, 2765t
 for pain, 91t
 for sleep enuresis, 192
 therapeutic monitoring of, 2765t
 Imiquimod
 for basal cell carcinoma, 501, 502
 for extramammary Paget's disease, 502
 for genital warts, 351, 1201, 1201t
 for melanoma, 496
 Immersion cooling, 479e-3
 Immersion (trench) foot, 478e-4
 Immune-complex formation, 372e-28, 372e-29t
 Immune deficiency diseases, primary. *See* Primary immune deficiency diseases
 Immune-mediated brachial plexus neuropathy (IBPN), 2692
 Immune memory, 372e-2
 Immune reconstitution inflammatory syndrome (IRIS)
 in histoplasmosis, 1334
 in HIV infection, 205e-1, 900, 1112-1113, 1234, 1254, 1261, 1261t
 pathogenesis of, 1234, 1261
 in Whipple's disease, 1093

- Immune response/immune system, **372e-1**.
See also Cell-mediated immunity
 adaptive. *See* Adaptive immune system
 aging and, 94e-4
 antitumor effects, 103e-20
 in celiac disease, 1940–1941
 cellular interactions in regulation of, 372e-22–23
 cellular migration in, 372e-28f
 clinical evaluation of, 372e-30
 components of, 2103
 endogenous derangements of, 377e-2. *See also* Autoimmunity
 endothelium in, 265e-2
 exogenous derangements of, 377e-1–2
 of gut, dysfunction of, 1876
 gut microbiota and, 86e-8
 in HIV infection. *See* HIV infection, immune response to
 in IBD, 1949–1950
 innate. *See* Innate immune system
 laboratory evaluation of, 2757–2764t
 molecular defects of, 372e-27t
 at mucosal surfaces, 372e-23, 372e-25f
 pattern recognition in. *See* Pattern recognition receptors (PRRs)
 suppression by cancer cells, 102e-17–18, 102e-18f
 terminology related to, 372e-1–2
 in women, 6e-4
- Immune thrombocytopenic purpura, 413, 728–729
- Immune tolerance, 372e-23, 735
- Immunization, **785**. *See also* Vaccine(s)
 for adults, 29t, 786
 administration of, 791–792, 791f
 contraindications to, 786, 789–790t, 790
 determining need for, 786
 provider recommendation for, 792
 schedules for, 786, 787–788f
 system support for, 792–793
 for cancer patient, 485t, 492
 for children, 2021–2022, 2022t
 consumer access to and demand for, 792
 documentation of, 792
 for health care workers, 793, 918
 in HIV infection, 788f, 798
 in HSCT recipients, 139e-4
 in nonmedical settings, 793
 performance monitoring, 793
 in pregnancy, 50–51, 788f, 790
 for transplant recipient, 928–929, 929t
 for travelers, 794–796, 795t
- Immunoblastic lymphomas, 1270. *See also* Diffuse large B-cell lymphoma
- Immunochemiluminometric assays (ICMAs), 2288
- Immunocompromised patient. *See also* Transplant recipient
 abdominal pain in, 105
 actinomyces in, 1091
 adenovirus infection in, 1208
 antibacterial therapy in, 936
 bite-wound infection in, 167e-1
Candida infections in, 350
 CMV infections in, 1191t, 1192
 diarrhea in, 266, 270, 855
 HRSV infection in, 1205
 immunization in, 788f, 790
 primary CNS lymphoma in, 601–602
 rhinovirus infection in, 1203
 sexual practices of, 492
 sinusitis in, 226
 skin cancer in, 388
- Immunodeficiency viruses, 1216, 1216f, 1219–1220. *See also* HIV
- Immunodeficiency with centromere instability and facial anomalies (ICF), 375e-2, 2109
- Immunodysregulation polyendocrinopathy enteropathy X-linked syndrome (IPEX), 377e-2, 2113, 2347–2348
- Immunofluorescent antibody technique, 150e-2
- Immunoglobulin(s). *See also* Antibody(ies)
 in adaptive immunity, 372e-21–22
 classification of, 710
 electrophoresis of, 710–712, 712f
 gene rearrangements in, 372e-20–21, 710, 711f
 hyperimmune, for viral hemorrhagic fever, 261e-5t
 properties of, 372e-21t
 reference values for, 2761t
 replacement of, 2112
 structure of, 372e-20f, 372e-21, 710, 711f
 types of, 372e-15
- Immunoglobulin A (IgA), 372e-21t, 372e-22, 372e-25t
- Immunoglobulin A (IgA) deficiency, 2111–2112
- Immunoglobulin A (IgA) nephropathy. *See* IgA nephropathy
- Immunoglobulin D (IgD), 372e-21t, 372e-22
- Immunoglobulin E (IgE)
 allergies and, 1670
 antibody to, 1678, 2119
 assays for, 2122
 in asthma, 1672
 in atopic dermatitis, 344
 properties of, 372e-21t
- Immunoglobulin E (IgE) receptor, 372e-24t
- Immunoglobulin G (IgG), 138e-1, 2112
 properties of, 372e-21t, 372e-22
 receptors for, 372e-7
 in rubella, 230e-2
- Immunoglobulin M (IgM)
 in *Brucella* infections, 194e-3
 production of, 138e-1
 properties of, 372e-21t
 in rubella, 230e-2
- Immunoglobulin replacement, 492, 2112
- Immunologic memory, 372e-16
- Immunologic priming, 372e-16
- Immunologic synapse, 372e-19
- Immunomodulators, transfusion-related, 138e-6
- Immunoproliferative small intestinal disease, 537, 719
- Immunoreceptor tyrosine-based activation motifs (ITAMs), 372e-19, 2115
- Immunosuppression
 in burn patients, 166e-2
 by cancer cells, 102e-17–18, 102e-18f
 UV-induced, 388
- Immunosuppressive therapy
 after cardiac transplantation, 1517
 after kidney transplantation, 1828–1829, 1829t
 after liver transplantation, 2070–2071
 chronic meningitis associated with, 911
 immunization during, 790
 for myasthenia gravis, 2705
 osteoporosis associated with, 2492
- Immunotherapy
 for allergic rhinitis, 2122–2124, 2123f
 for asthma, 1678
 for autoimmune and inflammatory diseases, 372e-30–31
 for cancer, 102e-17–18
 contraindications to, 2124
- Immunotoxicity, 91e-2t
- Impaired physician, 17e-4
- Impella device, 296e-5
- Impetigo, **349**
 bullous, 349, 361t, 362, 829, 967
 burn-wound, 166e-1
 clinical features of, 341t, 349, 350t, 967, 967f
 differential diagnosis of, 967
 etiology of, 349, 350t, 966–967
S. aureus, 349, 350t, 967
 streptococcal, 966–967
 treatment of, 349, 350t, 965t, 967
- Impetigo contagiosa, 25e-4f, 76e-8f, 829
- Impingement syndrome, 2248
- Implantable cardioverter-defibrillator (ICD)
 CMR in patient with, 270e-9
 complications of, 1499
 endocarditis associated with. *See* Infective endocarditis, cardiovascular implantable device-associated
 for heart failure, 1514, 1515t, 1770
- for hypertrophic cardiomyopathy, 1569–1570
 indications for, 1499
 for myocardial infarction, 1609, 1609f, 1770
 travelers with, 799
 for ventricular arrhythmias, 273e-7, 1495, 1499, 1499f
 ventricular tachycardia and, 1491
- Implicit memory, 182
- Imprinting
 definition of, 83e-7, 444e-2
 disorders caused by, 430, 439
 genomic, 430, 432f
 penetrance and, 437
- Imprinting syndromes, 83e-7, 101e-4
- IMV (intermittent mandatory ventilation), 1741–1742, 1742t. *See also* Mechanical ventilation
- Incapacitating/behavior-altering agents, 262e-1t, 262e-2t
- Incidentalomas, adrenal, 2321, 2321t, 2322f
- Inclusion body myositis, **2194**
 clinical features of, 462e-1, 2194–2196, 2195t
 diagnosis of, 2199–2200, 2199t, 2201f
 differential diagnosis of, 2197–2199
 familial/hereditary, 2195–2196
 immunopathology of, 2196, 2198f
 nonimmune factors in, 2197
 prognosis of, 2202
 treatment of, 2200–2202
 viral infections and, 2197
- Inclusion cyst, epidermoid, 342f, 364, 364t, 366
- Incompetent patient
 advance directives for, 58
 assessing capacity to make medical decisions, 17e-1
 choice and role of surrogates, 17e-1–2
 end-of-life care for, 66
- Incomplete penetrance, 437
- Incontinence
 fecal. *See* Fecal incontinence
 urinary. *See* Urinary incontinence
- Incontinentia pigmenti, 359, 360t, 431t
- Index of independence in activities of daily living, 78t
- India, 13e-7
- India ink, 150e-1
- Indian hedgehog, 2455
- Indian tick typhus, 1157
- Indigestion, **261**
 approach to the patient, 262–263
 definition of, 258
 differential diagnosis of, 261–262, 1894
 in esophageal disease, 1901
 in GERD. *See* Gastroesophageal reflux disease (GERD)
 mechanisms of, 261
 treatment of, 263
- Indinavir
 adverse effects of, 380, 1258, 1259, 2030
 drug interactions of, 14e-4, 42t, 246e-3t
 for HIV infection, 1278
 metabolism of, 34t
 molecular structure of, 1276f
 resistance to, 1280f
- Indirect alloreactivity, 373e-5
- Indirect Coombs (indirect antiglobulin) test, 138e-4f
- Indirect fraction, 1995
- Indomethacin
 for acute idiopathic pericarditis, 1573
 for cough headache, 2597
 drug interactions of, 43
 for exertional headache, 2597
 for gout, 2234
 for hemicrania continua, 2596
 for medullary thyroid cancer, 272
 for nephrogenic diabetes insipidus, 2279
 for pain, 91t
 for paroxysmal hemicrania, 2596
 for postprandial orthostatic hypotension, 2644
 for reactive arthritis, 2175
 for sex headache, 2597
 for stabbing headache, 2598

- Indoor air pollution, 1693
 Indoor residual spraying, 14
 Indoor tanning, 387, 388
 Induced pluripotent stem cells, 90e-2, 92e-1, 444e-7, 453t, 454
 Ineffective osmoles, 295
 Inert gas dilution, 306e-6
 Infant botulism, 987, 988
 Infantile poikilocytosis, 657
 Infants. *See also* Neonate(s)
 atopic dermatitis in, 344
 C. trachomatis infections in, 1171
 cobalamin deficiency in, 644
 cradle cap in, 347
 microbiome in, 86e-5
 seizures in, 2545–2546, 2546t
 water requirement in, 95e-1
 Infection(s), 145e-1
 burn-wound. *See* Burn-wound infections
 surgical site. *See* Surgical-site infections
 Infection-control programs
 for chickenpox, 917
 in critical care setting, 1734
 employee health service issues in, 918
 for filovirus infections, 1329
 for fungal infections, 917
 genomics in, 776
 for influenza, 916–917
 isolation techniques in, 916
 for legionellosis, 917–918
 for nosocomial diarrhea, 917
 organization, responsibilities, and scrutiny of, 912–913, 912t, 913t
 for *Streptococcus* group A, 917
 surveillance by, 912–913
 for tuberculosis, 917
 Infection susceptibility syndromes, 421, 422f
 Infectious (septic) arthritis, 833. *See also specific types*
 approach to the patient, 833
 bacterial
 anaerobic bacterial, 1100, 1102
 Brucella, 194e-2
 gonococcal, 835, 1006, 1007t, 1009t
 microbiology of, 834
 mycobacterial, 836
 nongonococcal, 834, 834f
 nontyphoidal *Salmonella*, 1054
 pathogenesis of, 833–834
 S. aureus, 957, 963
 from bite-wound infections, 167e-1
 etiology of, 833, 833t
 fungal, 836
 global considerations in, 837
 in HIV infection, 1261
 Lyme arthritis, 372e-24t, 835–836
 parasitic, 837
 postinfectious, 837. *See also* Reactive arthritis (Reiter's syndrome)
 prevention of, 838
 in prosthetic joints. *See* Prosthetic joint infections
 in sickle cell disease, 2242
 syphilitic, 836
 viral, 836–837
 Infectious diseases, 761. *See also specific diseases and agents*
 approach to the patient, 762. *See also* Acute febrile illness
 climate change and
 arbovirus infections, 151e-5–6
 El Niño–related disease outbreaks, 151e-8, 151e-8f
 Lyme disease, 151e-5–6, 151e-6f
 vector-borne disease, 151e-4–6, 151e-5f, 151e-6f
 waterborne disease, 151e-6–8, 151e-7f
 emerging, 802, 916
 fatigue in, 152
 gene therapy for, 91e-2f
 genomics of, 768
 in antibiotic resistance detection, 773–775
 clinical applications, 768, 769t
 in drug and vaccine development, 776–777
 epidemic potential and, 777–778
 epidemiology methods and, 776–778
 global considerations in, 778
 host-based diagnostics and, 775
 limitations and progress of, 770–771
 in microbial diagnostics, 768, 770f
 in organism detection, 771–772
 in pathogen discovery, 772–773, 773f
 in therapeutics, 775
 global considerations in, 761, 762f
 historical perspective, 761
 history in, 762–763
 imaging in, 765
 immune status in, 763
 infection control for, 766
 laboratory diagnosis of, 150e-1, 764–765
 by culture, 150e-3, 764
 detection methods, 150e-1
 direct detection, 150e-1–2
 gas-liquid chromatography, 150e-3
 MALDI-TOF MS, 150e-3
 microbial detection in blood, automation of, 150e-3
 nucleic acid tests, 150e-7–8, 150e-7f
 phenotyping, 150e-3
 specimen collection and transport, 150e-2–3, 150e-4–5t
 specimen-processing algorithm, 150e-6f
 susceptibility testing, 150e-8
 microbial pathogenesis in. *See* Microbial pathogenesis
 perspective on, 766
 physical examination in, 763–764
 in returning war veterans. *See* War veterans, infections in
 sexually transmitted. *See* Sexually transmitted infections (STIs)
 skin signs of, 76e-7, 764
 specialist consultation for, 766
 treatment of, 765–766, 767–768t
 vaccine-preventable, 785, 785t, 786
 Infectious mononucleosis
 clinical features of, 128t, 152, 1187–1188, 1187t
 CMV, 1191–1192
 complications of, 664, 1188
 diagnosis of, 1188, 1188f, 1189, 1189f, 1190t
 differential diagnosis of, 1190t, 1403t
 lymphadenopathy in, 409
 oral manifestations of, 238t
 pharyngitis in, 231
 prevention of, 1190
 rash in, 127, 128t, 362, 1187, 1187f
 in returned traveler, 802t
 splenomegaly in, 411
 treatment of, 1189–1190
 Infective endocarditis, 816
 Abiotrophia, 821
 abscesses in, 818, 901
 actinomycotic, 1089
 acute, 816
 anaerobic bacterial, 1100
 Aspergillus, 1347
 AV conduction block in, 1471t
 Bartonella, 816, 817, 823t, 824, 1081t, 1082
 in burn patient, 166e-2
 in cancer patient, 489
 Candida, 817t, 824, 1345
 cardiovascular implantable device-associated
 antimicrobial therapy for, 824
 diagnosis of, 819–820
 etiology of, 816
 surgical treatment of, 825
 as cause of fever of unknown origin, 137–138
 clinical features of, 784, 817–819, 817t, 818f, 1443
 clubbing/hypertrophic osteoarthropathy in, 250
 complications of, 826
 Corynebacterium, 823–824
 Coxiella burnetii, 816, 823t
 culture-negative, 817t
 diagnosis of, 817t, 819–820, 819t
 CT-PET, 270e-24, 270e-25f
 echocardiography, 270e-24, 270e-24f, 820, 820f, 821f
 embolic events in, 818, 818f, 825
 Enterobacter, 823
 enterococcal
 clinical features of, 973–974
 incidence of, 817t
 treatment of, 821–823, 822t, 975t
 etiology of, 780t, 784, 816, 817t
 Gemella morbillorum, 821, 822t
 glomerulonephritis associated with, 1835t, 1837–1838
 Granulicatella, 821
 HACEK group
 clinical features of, 183e-1, 817
 epidemiology of, 183e-1
 etiology of, 183e-1
 incidence of, 817t
 pathogenesis of, 183e-1
 treatment of, 183e-1, 183e-2t, 822t, 824
 health care-associated, 819, 958–959
 heart failure in, 818, 825
 IgM antibodies in, 372e-22
 in injection drug users, 816, 817, 817t, 958
 murmur in, 51e-2
 mycotic aneurysm in, 819, 826
 Nocardia, 1085
 nontyphoidal *Salmonella*, 1054, 1054t
 osteomyelitis and, 839
 outcome of, 826
 P. aeruginosa, 823, 1045t, 1046
 pathogenesis of, 816–817
 perivalvular infection in, 825
 pneumococcal, 817t, 823
 polymicrobial, 817t
 prevention of, 826–827, 826t
 prophylaxis for dental care, 242, 826t, 945t
 prophylaxis in congenital heart disease, 1522
 in surgical patient, 55
 prosthetic valve. *See* Prosthetic heart valve, endocarditis in
 in Q fever, 1162
 retinal hemorrhage in, 200
 septic emboli and, 2564
 septic emboli in, 25e-6f, 76e-12f, 818, 818f
 skin manifestations of, 130t, 134, 784
 splenomegaly in, 411
 staphylococcal
 clinical features of, 130t, 817, 958
 diagnosis of, 958
 echocardiography in, 820f
 emboli in, 818, 818f, 958, 958f
 epidemiology of, 958
 incidence of, 817t
 pathogenesis of, 817
 treatment of, 822t, 823, 824, 963
 streptococcal
 clinical features of, 130t
 group C or D, 969
 group D, 971
 incidence of, 817t
 treatment of, 820, 822t, 823
 subacute, 816
 T. whipplei, 1092
 treatment of
 antimicrobial therapy, 780t, 820–824, 822–823t, 826, 975t
 empirical, 767t, 824
 monitoring, 824
 outpatient, 824
 surgical, 824–826, 824t, 825t
 vegetations in, 76e-12f
 Inferior petrosal venous sampling, in Cushing's syndrome, 2272
 Inferior pontine stroke syndrome, 2577f
 Inferior vena cava filter, 613, 1636
 Infertility, 2387
 after Hodgkin's lymphoma, 709
 after mumps infection, 231e-2
 after testicular cancer, 592

- Infertility (*Cont.*):
 approach to the patient, 2388
 chemotherapy-related, 103e-25–26, 623
 in congenital adrenal hyperplasia, 2356
 in cystic fibrosis, 1697
 definition of, 2387
 etiology of, 2387–2388, 2388f
 in Klinefelter's syndrome, 2350–2351
 male, 2360, 2388, 2388f, 2389, 2725
 menstrual disorders and, 2388. *See also*
 Menstrual disorders
 ovulatory dysfunction/failure in, 336–337,
 437, 2388, 2389
 prevalence of, 2254t, 2387
 psychological aspects of, 2388
 screening/testing for, 2254t
 STIs causing, 870t
 treatment of, 2388–2389
 tubal disease and, 2388, 2389
 in Turner's syndrome, 2352
- Infestations
 delusional, 2750–2751
 fly, 2746–2747
 leech, 2747
 louse, 2745–2746
 maggot, 2747
 scabies, 2744–2745
 tongue worms, 2747
- Inflamming, 94e-4
- Inflammasome(s)
 characteristics of, 372e-1, 372e-4
 diseases associated with, 372e-8t, 1845, 2215
 gut microbiota and, 86e-8
 in response to infection, 145e-7, 145e-8f, 1752
- Inflammation
 aging and, 75, 75f, 76, 94e-4
 anemia and, 398, 628, 628t
 in asthma, 1672–1673, 1672f
 atherosclerosis and, 291e-2, 291e-9, 291e-9f
 cardinal signs of, 414f
 chronic, 145e-10
 endothelium in, 265e-2
 immune cells at sites of, 372e-28f
 nociceptor-induced, 88
 phagocyte activation and, 145e-5
 trafficking molecules in, 372e-29t
 in vivo assessment of, 423
- Inflammatory bowel disease (IBD), **1947**. *See also*
Crohn's disease; *Ulcerative colitis*
 after infectious diarrhea, 855t
 in ankylosing spondylitis, 2171
 atypical colitides mimicking, 1957
 cancer risk in, 1965
 colorectal cancer in, 539, 1898t, 1964–1965,
 1964f
 commensal microbiota and, 1948f, 1949
 differential diagnosis of, 1061, 1956–1957
 disorders associated with, 1949t, 2178–2179
 dysplasia in, 1964f, 1965
 early-onset, 1949, 1949t
 epidemiology of, 1947, 1947t
 etiology of, 1947–1948, 1947f
 extraintestinal manifestations of
 aortitis, 1642
 cardiopulmonary, 1959
 dermatologic, 369, 1956–1957
 hepatobiliary, 1958
 metabolic bone disorders, 1958
 neuropathy, 2683
 ocular, 1958
 oral, 237
 rheumatologic, 1958
 thromboembolic disorders, 1958
 urologic, 1958
 genetic considerations in, 1947, 1948f, 1949,
 1949t, 1950t
 global considerations in, 1947
 gut microbiota and, 86e-8
 immune regulation in, 372e-27t, 1949–1950
 inflammatory cascade in, 1950–1951
 noninfectious diseases mimicking, 1957
 pathogenesis of, 1948–1950, 1948f
 in pregnancy, 49, 1964
 racial and ethnic differences in, 1947–1948
- smoking and, 1947
 in spondyloarthritis, 2178–2179
 treatment of, **1959**, 1963f
 antibiotics, 1960
 5-ASA agents, 1959–1960, 1959t
 azathioprine, 1960
 biologic therapies, 1961
 cyclosporine, 1960–1961
 glucocorticoids, 1960
 6-mercaptopurine, 1960
 methotrexate, 1960
 nutritional therapies, 1960
 during pregnancy, 1964
 surgery, 1962–1964, 1963t
 tacrolimus, 1961
 therapies in development, 1962
 tumor markers in, 473t
- Inflammatory diarrhea, 269–270. *See also*
 Diarrhea
- Inflammatory mediators, in sensitization, 87–88
- Inflammatory myopathies. *See also*
 Dermatomyositis; *Inclusion body*
 myositis; *Polymyositis*
 autoantibodies in, 2196
 clinical features of, 2194–2195, 2195t
 diagnosis of, 2199–2200, 2199t
 differential diagnosis of, 2195t, 2197–2199
 drug-induced, 462e-20, 2197, 2199
 global considerations in, 2197
 malignancies associated with, 2196
 overlap syndrome, 2196
 pathogenesis of, 2196–2197, 2197f, 2198f
 prognosis of, 2202
 treatment of, 2200–2202
- Inflammatory response, 1746–1747, 1747f,
 1950–1951
- Infliximab
 adverse effects of
 actinomycosis, 1088
 common, 2146t
 cutaneous, 377
 infections, 422, 1962
 septic arthritis, 834
 serious, 348t, 1962, 2146t
 for ankylosing spondylitis, 2172–2173, 2172f
 antibodies to, 1961
 for autoimmune and inflammatory diseases,
 372e-30
 for IBD, 1961, 2179
 monitoring during treatment with, 2146t
 for psoriatic arthritis, 348t, 2177
 for rheumatoid arthritis, 2145, 2146t
 for sarcoidosis, 2211–2212, 2212t
- Influenza, **1209**
 asthenia following, 1212
 in cancer patient, 489
 clinical features of, 784, 1202t, 1211–1212
 complications of, 1211, 1211t, 1212–1213
 control of, 916–917
 deaths from, 6e-1t, 56t
 definition of, 1209
 diagnosis of, 775, 1213
 epidemiology of, 778, 1209–1211
 extrapulmonary complications of, 1212–1213
 laryngitis in, 234
 monocyte dysfunction in, 422
 morbidity and mortality caused by, 1211
 myocarditis in, 1560
 otitis media in, 229
 pandemic and inter-pandemic disease patterns
 and features, 1209t, 1210
 pathogenesis of, 1211
 pathogen evolution of, 778, 1209–1210, 1209t
 pharyngitis in, 231
 pneumonia and, 1212
 in pregnancy, 50, 1212
 prevention of, 215e-2t, 215e-5, 1213t, 1214.
See also Influenza vaccine
 Reye's syndrome and, 1212–1213
 sinusitis in, 226
 in transplant recipient, 923
 treatment of, 214e-5–6, 215e-1, 215e-2t,
 215e-5–6, 1213–1214, 1213t
 in upper respiratory infections, 225
- Influenza A viruses, 1209–1211, 1209f, 1209t
 Influenza B viruses, 1209, 1211
 Influenza C viruses, 1209, 1211
 Influenza vaccine
 contraindications to, 789t, 1213
 coverage in U.S., 792
 Guillain-Barré syndrome and, 1214, 2695
 inactivated, 789t, 1214
 live attenuated, 789t, 1214
 precautions for, 789t
 recommended administration of
 in adults, 29t, 84, 787f, 788f, 809, 1214
 in cancer patients, 485t, 492
 in HIV infection, 798, 1252t
 in liver disease, 1995
 in pregnancy, 50
 in transplant recipients, 929t
 in travelers, 794
- Influenza virus
 adherence of, 145e-2t
 as bioterrorism agent, 261e-9
 cell membrane interactions of, 214e-3
 host range of, 214e-6
 mutations in, 214e-5
 subtypes of, 1209t
 viral assembly and egress in, 214e-5
 virology of, 1209
- Information technology, 3
- Informed consent, 4, 17e-1, 443, 449
- Infusate, contaminated, 915
- Inguinal syndrome, 1170
- INH. *See* Isoniazid (INH)
- Inhalational anthrax, 261e-2–3, 784
- Inhaled anesthetics, 54
- Inhibin, 401e-4, 2360, 2378, 2761t
- Inhibitor of apoptosis proteins (IAP), 102e-9,
 102e-9f
- Inhibitory NK cell receptors (iNKR), in HIV
 infection, 1238
- Injection drug users
Candida infections in, 836
 endocarditis in, 816, 817t, 819, 958
 HIV infection in, 1220t, 1222, 1285
 sepsis/septic shock in, 1757t
 tetanus in, 984
 wound botulism in, 988
- Inkoo virus, 1309t
- Innate immune system, 372e-1, **372e-2**. *See also*
 Immune response/immune system
 adaptive immune system and, 372e-2
 components of, 372e-2, 372e-4t
 definition of, 372e-1
 effector cells of, **372e-4**, 372e-7t
 basophils. *See* Basophil(s)
 dendritic cells. *See* Dendritic cells
 eosinophils. *See* Eosinophil(s)
 macrophages. *See* Macrophage(s)
 monocytes. *See* Monocyte(s)
 neutrophils. *See* Neutrophil(s)
 NK cells. *See* Natural killer (NK) cells
 primary deficiency of. *See* Primary immune
 deficiency diseases, of innate immune
 system
 PRRs of, 372e-2, 372e-4, 372e-5t
- Inorganic dusts, 1688t, 1691
- Inoue balloon technique, mitral valvotomy,
 1542f
- Inpatient care, 5
- Inpatient deaths, 55, 56f
- INR (International Normalized Ratio), 405,
 755
- Insect bites/stings, 341t, 364t, 365, 371, **2748**.
See also specific insects
- Insect-borne disease, prevention of, 796
- Insecticides, 41, 473e-10t
- Insecticide-treated bed nets, 14
- Insertion, gene, 434
- Insomnia
 assessment of, 65
 drug-induced, 190
 etiology of, 65
 evaluation of, 190
 fatal familial, 191
 in medical disorders, 190–191

- in neurologic disorders, 191
 psychiatric disorders and, 190
 psychophysiological factors in, 190
 sleep hygiene and, 190
 in terminally ill patient, 65
 treatment of, 65, 191–192, 191t
- Institute of Medicine (IOM), 16e-2, 16e-5–6
- Instrumental activities of daily living (IADLs), 72t, 78t
- Insulin. *See also* Insulin therapy
 action of, 2402–2403, 2430, 2431t
 biosynthesis of, 2402
 ectopic production of, 609t
 effectors of, 400e-1t
 reference values for, 2761t
 secretion of, 2402, 2402f, 2405
 in sepsis/septic shock, 1758
 signaling pathways of, 400e-1t
 structure of, 400e-1
- Insulin aspart, 2411, 2411t
- Insulin autoimmune syndrome (Hirata's syndrome), 2348
- Insulin-dependent diabetes mellitus (IDDM). *See* Diabetes mellitus, type 1
- Insulin detemir, 2411, 2411t
- Insulin glargine, 2411, 2411t
- Insulin glulisine, 2411, 2411t
- Insulin-like growth factor (IGF), in aging, 85, 94e-5
- Insulin-like growth factor I (IGF-I)
 action of, 401e-3, 401e-4
 adverse effects of, 401e-4
 factors affecting levels of, 401e-3
 in gastroduodenal mucosal defense and repair, 1913f
 injected, 401e-4
 pNET secreting, 559t, 571
 signal transduction pathway for, 400e-1t
 structure of, 400e-1
 in thyroid hormone synthesis, 2285
- Insulin-like growth factor II (IGF-II)
 action of, 401e-3
 ectopic production of, 609t, 611–612
 in sarcoma, 119e-1
- Insulin-like growth factor II gene, 439, 611
- Insulin lispro, 2411, 2411t
- Insulinoma, **569**
 clinical features of, 559t, 569, 2337
 diagnosis of, 569, 2337, 2434
 endogenous hyperinsulinism due to, 2434
 incidence of, 559t, 2434
 in MEN 1, 2337–2338
 obesity in, 415e-4
 screening tests for, 2337t
 treatment of, 569, 2338, 2434
- Insulinomatosis, 569
- Insulin receptor, 372e-24t
- Insulin receptor substrates, 2403
- Insulin resistance
 atherosclerosis and, 291e-6
 autoimmune, 372e-24t, 377e-4, 2348
 colorectal cancer and, 538
 fecal microbiota transplantation for, 86e-10
 free fatty acids in, 2450
 lipoprotein disorders and, 2438, 2439
 in metabolic syndrome. *See* Metabolic syndrome
 obesity and, 415e-6
 in pregnancy, 48
 specialized nutritional support and, 98e-4
 syndromes of, 2405–2406
 treatment of, 2454
- Insulin secretagogues
 incretins, 2414–2415, 2414t, 2416
 sulfonylureas. *See* Sulfonylureas
 for type 2 diabetes mellitus, 2414t
- Insulin therapy
 adverse effects of, 2392
 for diabetic ketoacidosis, 64e-1
 for DKA, 2419
 for hyperkalemia, 312
 hypokalemia and, 305
 with parenteral nutrition, 98e-7
 in pregnancy, 48
 for type 1 diabetes mellitus, 2411–2412, 2411t, 2412f
 for type 2 diabetes mellitus, 2414t, 2415–2416
- Insulin tolerance test, 2310–2311
- Integrase, 1217
- Integrase inhibitors. *See* Combination antiretroviral therapy (cART)
- Integrative medicine, 14e-1–2. *See also* Complementary and alternative medicine
- Integrin(s)
 of innate immune system, 372e-2
 in neutrophil adhesion, 416
 stem cell, 89e-1, 139e-1
 in tumor angiogenesis, 102e-14f, 102e-15, 102e-16f, 102e-17t
- Integrin $\alpha 4\beta 7$, 1234–1235
- Intensity-modulated radiation therapy, 583
- Intensivists, 5, 1729
- Interactome, 87e-7f
- Intercalated cells, 332e-5f, 332e-9
- Intercellular adhesion molecule 1 (ICAM-1), 1242, 2170
- Interdigitating dendritic cell sarcoma (reticulum cell sarcoma), 135e-9
- Interferon(s)
 genetic variations in response to, 39t
 pharmacology of, 215e-10
 structure of, 372e-16
 in viral infections, 214e-8, 215e-10
- Interferon- α (IFN- α)
 action and targets of, 103e-22
 for adult T-cell leukemia/lymphoma, 225e-5
 adverse effects of, 103e-22, 567, 574, 2033
 cardiac, 1563
 cutaneous, 378, 379, 381
 depression, 63
 for carcinoid syndrome, 567
 for chronic HBV infection, 2033
 for chronic HCV infection, 2041–2043, 2042f
 for chronic HDV infection, 215e-10, 2040
 for CML, 694
 for follicular lymphoma, 705
 for genital warts, 1201, 1201t
 for hypereosinophilic syndromes, 423
 for Kaposi's sarcoma, 1270, 1270t
 for NETs, 574
 for polycythemia vera, 674
 production of, 372e-6
 for renal cell carcinoma, 579
 source, target, and biologic activity of, 372e-11t
 for systemic mastocytosis, 2121
 thyroid dysfunction, 2299
 for viral hemorrhagic fever, 261e-8
- Interferon- $\alpha 2a$ (IFN- $\alpha 2a$)
 adverse effects of, 498
 for chronic HCV infection, 215e-4t, 215e-10
 for chronic HDV infection, 215e-5t
 for melanoma, 498
 pegylated. *See* Pegylated interferon- $\alpha 2a$ (PEG IFN- $\alpha 2a$)
- Interferon- $\alpha 2b$ (IFN- $\alpha 2b$)
 for chronic HBV infection, 215e-4t, 215e-10
 for chronic HCV infection, 215e-4t
 for chronic HDV infection, 215e-5t
 for condyloma acuminatum, 215e-4t
 pegylated. *See* Pegylated interferon- $\alpha 2b$ (PEG IFN- $\alpha 2b$)
- Interferon- $\alpha n3$ (IFN- $\alpha n3$), for condyloma acuminatum, 215e-4t
- Interferon- β (IFN- β)
 adverse effects of, 2667
 for MS, 2667, 2669t, 2671
 in pregnancy, 49
 source, target, and biologic activity of, 372e-11t
- Interferon- $\beta 1a$ (IFN- $\beta 1a$), 2667–2668, 2669t
- Interferon- $\beta 1b$ (IFN- $\beta 1b$), 2667–2668, 2669t
- Interferon- γ (IFN- γ)
 in *Brucella* infections, 194e-2
 for chronic granulomatous disease, 372e-30, 423
 in HIV infection, 422
 indications for, 423
 in nontuberculous mycobacterial infections, 1129, 1129f
 for Q fever, 1162
 source, target, and biologic activity of, 372e-11t
 suppression of erythropoietin production by, 630, 630f
 in T-cell lymphoma, 422
- Interferon- γ (IFN- γ) release assays (IGRAs), 138, 1115, 1131
- Interferon spray, 1203
- Interleukin(s), 372e-10–11t, 372e-14, 372e-14t, 413
- Interleukin-1 (IL-1)
 in anemia of inflammation, 630
 in familial Mediterranean fever, 422
 in fever induction, 124, 125f, 141
 in pneumonia, 804
 source, target, and biologic activity of, 372e-10t
- Interleukin-1 receptor antagonist (IL-1Ra)
 deficiency of, 2215
 recombinant. *See* Anakinra
- Interleukin-1 receptor-associated kinase (IRAK-4), 145e-5, 145e-6f, 950
- Interleukin-1 β (IL-1 β)
 in HIV infection, 1235
 in osteoarthritis, 2227, 2227f
 in sepsis/septic shock, 1752
- Interleukin-2 (IL-2), 372e-10t
- Interleukin-2 (IL-2) therapy
 action and targets of, 103e-22
 adverse effects of, 63, 103e-22, 253t, 379, 621, 2299
 for melanoma, 103e-22, 498
 for renal cell carcinoma, 103e-22, 579
- Interleukin-3 (IL-3), 372e-10t
- Interleukin-4 (IL-4), 372e-10t
- Interleukin-5 (IL-5), 372e-10t, 422, 1673
- Interleukin-6 (IL-6)
 age-related changes in levels of, 75, 75f
 in depression, 465e-4
 excess of, 612
 in fever induction, 124, 125f
 in HIV infection, 1240t, 1249t
 in sepsis/septic shock, 1753
 source, target, and biologic activity of, 372e-10t
- Interleukin-7 (IL-7), 372e-10t, 1237
- Interleukin-8 (IL-8), 372e-10t, 416–417, 1701
- Interleukin-9 (IL-9), 372e-10t
- Interleukin-10 (IL-10), 372e-10t, 1240t, 1753
- Interleukin-11 (IL-11), 103e-23, 372e-10t
- Interleukin-12 (IL-12), 372e-10t, 1129, 1129f
- Interleukin-13 (IL-13), 372e-10t
- Interleukin-14 (IL-14), 372e-10t
- Interleukin-15 (IL-15), 39t, 372e-10t
- Interleukin-16 (IL-16), 372e-10t
- Interleukin-17 (IL-17), 372e-10t, 2141, 2142f
- Interleukin-18 (IL-18), 372e-10t, 1752, 1808
- Interleukin-21 (IL-21), 372e-10t
- Interleukin-22 (IL-22), 372e-10t
- Interleukin-23 (IL-23), 372e-10t
- Interleukin-24 (IL-24), 372e-11t
- Interleukin-25 (IL-25; IL-17E), 372e-11t
- Interleukin-26 (IL-26), 372e-11t
- Interleukin-28B (IL-28B), 39t
- Interleukin-36 (IL-36) receptor antagonist, 2215
- Intermediate-density lipoproteins (IDLs), 1241t, 2436–2437, 2436f, 2436t
- Intermediate metabolizers, 40
- Intermittent mandatory ventilation (IMV), 1741–1742, 1742t. *See also* Mechanical ventilation
- Internal carotid artery, 2574, 2574f, 2575f
- International Cancer Genome Consortium, 101e-8
- International Classification of Functioning, Disability, and Health, 78–79

- International Health Regulations, 8
 International Monetary Fund (IMF), 8
 International Normalized Ratio (INR), 405, 755
 International Prognostic Index, for non-Hodgkin's lymphoma, 700, 701f, 701t
 International Sensitivity Index (ISI), 405, 755, 1998
 Internet resources
 on complementary and alternative medicine, 14e-4t
 on dietary supplement–drug interactions, 14e-4t
 on genomics and genetic disorders, 426t
 on HIV disease, 1273t
 for medical information, 6
 Internuclear ophthalmoplegia, 209–210, 210f
 Interstitial cystitis/bladder pain syndrome, **60e-3**
 Interstitial fluid, 250
 Interstitial lung diseases (ILDs)
 bronchoalveolar lavage in, 1710
 classification of, 1708, 1708t
 in connective tissue disorders, 1713–1714
 with diffuse alveolar hemorrhage, 1715
 drug-induced, 1714
 dyspnea in, 47e-2, 47e-2t
 eosinophilic. *See* Pulmonary infiltrates with eosinophilia
 global considerations in, 1716
 with granulomatous response, 1715
 eosinophilic granulomatosis with polyangiitis. *See* Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
 granulomatosis with polyangiitis. *See* Granulomatosis with polyangiitis (Wegener's)
 hypersensitivity pneumonitis. *See* Hypersensitivity pneumonitis
 sarcoidosis. *See* Sarcoidosis
 history in, 1709
 idiopathic pulmonary fibrosis. *See* Idiopathic pulmonary fibrosis
 imaging in, 308e-7–9f, 1710, 1710f
 inherited disorders associated with, 1715
 laboratory evaluation of, 1709–1711, 1711t
 lymphocytic, 1715–1716
 pathogenesis of, 1708, 1709f
 physical examination in, 1709
 pneumonia. *See* Interstitial pneumonia
 pulmonary alveolar proteinosis, 1714
 pulmonary hypertension in, 1659
 pulmonary Langerhans cell histiocytosis, 1713
 pulmonary lymphangioleiomyomatosis, 308e-5f, 1714–1715
 respiratory bronchiolitis–associated, 1713
 in rheumatoid arthritis, 2138
 smoking-associated, 1713
 in systemic sclerosis, 2160–2161, 2161f
 treatment of, 1711
 Interstitial nephritis. *See* Acute interstitial nephritis (AIN)
 Interstitial pneumonia
 acute, 1712
 desquamative, 1713
 in HIV infection, 1255
 in HSCT recipient, 139e-3
 lymphoid, 1255
 nonspecific, 1255, 1712, 1712f
 Interstitial pneumonitis, lymphocytic, 1716
 Interstitial space, 295
 Intertrigo, *Candida*, 1343
 Interventional cardiology, 296e-1. *See also* Percutaneous coronary interventions (PCI)
 Interventional neuroradiology, 440e-11
 Intervertebral disk
 anatomy of, 111, 112f
 herniation of, 113f, 114–115, 115f
 replacement, 120
 Intestinal flukes
 diagnosis of, 245e-2t
 epidemiology of, 245e-2t, 1423t, 1429
 life-cycle hosts of, 245e-2t
 treatment of, 1428t
 Intestinal ischemia, **1978**
 acute, 1979–1980, 1980t
 anatomy of, 1979, 1979f
 arterioocclusive, 1979, 1980t
 chronic, 1981
 clinical features of, 106t, 258, 259, 1979–1980
 diagnosis of, 1979–1980
 epidemiology of, 1978–1979
 incidence of, 1978
 mesenteric venous thrombosis, 1979, 1980t
 nonocclusive, 1980, 1980t
 pathophysiology of, 1979–1980
 treatment of, 1980–1981, 1980t
 Intestinal lymphangiectasia
 clinical features of, 269, 1654, 1945
 congenital, 269
 small-intestinal mucosal biopsies in, 1939, 1940t, 1941f, 1941t
 Intestinal necrosis, 312
 Intestinal obstruction, **1981**
 in cancer, 1789–1790
 clinical features of, 1982–1983
 abdominal pain, 103, 106t, 1983
 abdominal swelling, 285, 1983
 lactic acidosis, 318
 nausea and vomiting, 258, 259, 1983
 closed-loop, 1982
 in cystic fibrosis, 1697
 diagnosis of, 107, 1892, 1892f, 1983–1984
 differential diagnosis of, 103–104, 106
 epidemiology of, 1981–1982, 1982t
 etiology of, 1981–1982, 1981t, 1982t
 pathophysiology of, 1982
 physical examination in, 1983
 postoperative, 1985
 in systemic sclerosis, 2162, 2164
 treatment of, 346e-1v, 1892, 1895f, 1984–1985
 Intestinal pseudoobstruction (Ogilvie's disease)
 clinical features of, 258–259, 305, 1892
 diagnosis of, 1892, 1894f
 etiology of, 1982
 treatment of, 260t, 261, 1885, 1892, 1894f
 Intestinal resection, 268, 1943
 Intestinal stagnant loop syndrome, 645
 Intestinal stem cells, 453
 Intima, 265e-1
 Intimal dissection, 296e-1, 296e-2f
 Intimate partner violence. *See* Domestic abuse/violence
 Intraabdominal abscess/infections
 actinomycosis, 1089, 1090f
 anaerobic bacterial, 1099, 1102
 C. septicum, 1099
 enterococcal, 974
 intraoperative. *See* Intraoperative abscess
 liver. *See* Liver abscess
 nontyphoidal *Salmonella*, 1054
 perinephric, 851
 psoas, 852
 renal, 851
 splenic. *See* Splenic abscess
 in transplant recipient, 927
 treatment of, 767t, 1102
 Intraaortic balloon pump (IABP), 296e-5, 1763
 Intraarticular injections, 2232, 2232t
 Intracellular fluid, 295
 Intracranial atherosclerosis, 2572
 Intracranial hemorrhage, **2581**
 aphasia in, 178
 clinical features of, 2582–2583
 clinical outcomes of, 2585t
 coma in, 1775–1776
 diagnosis of, 2582, 2584
 emergency management of, 2582
 etiology of, 2559, 2582–2584, 2583t
 headache in, 108
 head injury and, 457e-2, 2584
 hypertension and, 1615, 1627, 2582–2583, 2583f
 lobar, 2583
 prevention of, 2585
 prognosis of, 2585t
 treatment of, 2584–2585
 vascular malformations and, 2585–2586
 Intracranial hypotension, 2607
 Intracranial infections, 780t, 784
 Intracranial meningitis, 906
 Intracranial pressure (ICP), increased
 in bacterial meningitis, 887, 890
 from brain metastases, 1792
 headache with, 110
 in intracranial meningitis, 906
 monitoring of, 1779, 1780f
 nausea and vomiting in, 259
 ocular manifestations of, 40e-4f, 203, 203f, 209
 pathophysiology of, 1778–1779, 1779f
 SA node dysfunction in, 1467, 1467t
 signs of, 1780
 in subarachnoid hemorrhage, 1785
 treatment of, 1779–1780, 1780t, 1786
 Intracytoplasmic sperm injection (ICSI), 2360, 2389
 Intrahepatic cholestasis of pregnancy, 49
 Intramedullary syndromes, 2652
 Intramuscular injection, of vaccines, 791, 791f
 Intraocular pressure, increased, 200
 Intraperitoneal abscess, **849**, 850f
 Intraperitoneal cavity, 846, 846f
 Intrathecal administration, opioids, 92
 Intrauterine devices (IUDs), 1089, 1090f, 2390, 2390t
 Intrauterine growth retardation, 333e-1–2, 2355
 Intrauterine infection, 50
 Intrauterine insemination, 2389
 Intravascular device–related infections. *See* Catheter-related bloodstream infections
 Intravascular large B-cell lymphoma, 135e-2
 Intravascular space, 295
 Intravenous immunoglobulin (IVIg)
 adverse effects of, 309, 2706
 for antiphospholipid syndrome, 2136
 for autoimmune and inflammatory diseases, 372e-31
 for *C. difficile* infections, 860t, 861
 for chronic inflammatory demyelinating polyneuropathy, 2699
 for CMV pneumonia, 922
 for dermatomyositis, 2201
 for epidermolysis bullosa acquisita, 374
 for Guillain-Barré syndrome, 2697
 for immune thrombocytopenic purpura, 728
 for Kawasaki disease, 2193
 for MS, 2672
 for mucous membrane pemphigoid, 374
 for myasthenia gravis, 2705–2706
 for parvovirus B19 infection, 669, 1197
 for pemphigus vulgaris, 371
 replacement therapy for cancer patient, 492
 for rheumatic fever, 2153
 for stiff-person syndrome, 618
 for streptococcal toxic shock syndrome, 965t, 969
 Intraventricular hemorrhage, primary, 2584
 Intravesical therapy, for bladder cancer, 576
 Intrinsic factor (IF), 350e-1, 372e-24t, 640, 645
 Intrinsic pathway, coagulation, 401, 401f
 Introns, 427
 Intussusception, 258, 274, 277, 1985
 Invariant chain, 373e-5
 Invasin, 1076
 Inverse psoriasis, 347. *See also* Psoriasis
 Inverse ratio ventilation (IRV), 1738, 1742. *See also* Mechanical ventilation
 Inversion, chromosome, 83e-6
 Inverted duplication 15 syndrome, 83e-6
Investing in Health, 9
 In vitro fertilization (IVF), 444, 2360, 2389
 Involuntary active euthanasia, 67t
 Involuntary weight loss, **274**
 assessment of, 275–276, 275t
 in cancer, 474
 in elderly, 81, 274–275
 in esophageal cancer, 533
 etiology of, 275, 275t
 in HIV infection, 1268
 in lung cancer, 511

- in malabsorption syndromes, 1946t
 significance of, 274
 in thyrotoxicosis, 2294
 treatment of, 275
- Iodides, adverse effects of, 356t, 366, 422
- Iodine, 96e-10
 deficiency of. *See* Iodine deficiency
 excess intake of, 2292
 for goiter, 2301
 metabolism and transport of, 2284–2285, 2285f
 parental supplementation of, 98e-6t
 recommended intake of, 95e-3t, 2285
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t
- Iodine deficiency
 clinical features of, 96e-9t, 2285
 global considerations in, 461, 2285, 2285f
 goiter and, 2301
 hypothyroidism and, 2292
 laboratory evaluation of, 2768–2769
 prevention of, 2292
- Iodine dermatitis, 246e-9
- ¹³¹Iodine-ethiodol, 263e-3t, 263e-6t, 263e-7, 550
- ¹³¹Iodine-MIGB, 2331
- Iodoquinol
 action of, 246e-9
 adverse effects of, 246e-4t, 246e-9
 for amebiasis, 246e-9, 1367t
 contraindications to, 246e-9
 for *D. fragilis* infection, 1409
 pharmacology of, 246e-9
 in pregnancy and lactation, 246e-4t
- Ion absorption, congenital defects in, 268
- Ion channels
 cardiac, 273e-1f, 273e-2, 273e-2f
 neurologic, 444e-2, 444e-2t
- Ion exchangers, for internal radionuclide contamination, 263e-6, 263e-6t
- Ionization, 103e-4
- Ionizing radiation
 AML and, 678
 aplastic anemia and, 663
 breast cancer and, 524
 from cardiac imaging, 270e-7–9
 lung cancer and, 506
 skin damage from, 357
- Iopanoic acid, 727t, 2286t, 2297
- IP-10 (IFN- γ -inducible protein-10), 372e-12t
- IPAF inflammasome, 145e-7
- Ipecac syrup
 adverse effects of chronic use of, 473e-6, 2199
 for internal radionuclide contamination, 263e-6
 for poisoning/drug overdose, 473e-6
- IPEX (immunodysregulation polyendocrinopathy enteropathy X-linked) syndrome, 377e-2, 1949t, 2113, 2347–2348
- IPF1* gene mutations, 431t, 2406
- Ipilimumab
 action of, 102e-5t
 adverse effects of, 359, 380, 499, 607
 indications for, 103e-21t
 for lung cancer, 521
 for melanoma, 103e-22, 498–499
- Ipodate, 2286t, 2297, 2300
- ipr1* gene, 1106
- Ipratropium bromide, 1676, 1705, 2122, 2123f
- Iquitos virus, 1309t
- IRAK1* gene mutations, 2155
- IRAK-4 (interleukin-1 receptor-associated kinase), 950
- IRAK-4 (interleukin-1 receptor-associated kinase) deficiency, 420t
- Iraqi Jewish population, factor XI deficiency in, 736
- Iridectomy, laser, 200
- Iridium exposure, 263e-3t
- Iridocyclitis. *See* Uveitis
- Irinotecan
 action of, 103e-13
 adverse effects of, 40, 103e-10t, 103e-13, 103e-25, 543, 1798
- for carcinoma of unknown primary, 120e-4
 for colorectal cancer, 543
 drug interactions of, 14e-4, 1793
 for Ewing's sarcoma, 119e-4
 for gastric cancer, 535
 genetic variations in response to, 39t
 for lung cancer, 520t, 522
 metabolism of, 34t, 35
 for pancreatic cancer, 557
 for soft tissue sarcoma, 119e-2
- IRIS. *See* Immune reconstitution inflammatory syndrome (IRIS)
- Iritis. *See* Uveitis
- Iron
 absorption of, 625–626, 625f
 accumulation in brain, 2625–2626
 average intake of, 626
 in bone marrow, 81e-8f
 deficiency of, 96e-9t, 98e-5, 192, 461, 463t.
 See also Iron-deficiency anemia;
 Iron-deficient erythropoiesis
 distribution in body, 625, 625t
 excess of. *See* Iron overload
 homeostasis, pathways of, 2515f
 metabolism of, 625–626, 625f
 nutritional balance of, 626
 parenchymal, in liver, 2516
 recommended intake of, 95e-3t, 626
 serum, 393, 397. *See also* Ferritin, serum
 in hypoproliferative anemia, 398, 629–630, 630t
 in iron-deficiency states, 627f
 levels by sex and age, 627–628, 627f
 stores of, 625f, 625t, 626, 627t, 628t, 2517t
 supplements. *See* Iron supplements
 toxicity of. *See* Iron overload
- Iron cycle, 625–626, 625f
- Iron-deficiency anemia, **626**
 causes of, 627, 627t
 clinical features of, 627
 diagnosis of, 627–628, 628t
 differential diagnosis of, 628, 628t
 from gastrointestinal bleeding, 279, 1896–1897
 gastrointestinal cancer and, 535, 541
 global considerations in, 626
 in hookworm infections, 1414
 pathophysiology of, 398
 peripheral blood smear in, 81e-2f, 394f
 stages of, 626–627, 627f
 surgery-related, 1927
 treatment of, 628–629, 629t
- Iron-deficient erythropoiesis, 627, 627f
- Iron dextran, 629
- Iron gluconate, 629
- Iron-hematoxylin stain, 245e-1
- Iron overload
 acute, 626
 chronic, 650
 clinical features of, 96e-9t
 CMR in, 270e-21, 270e-27v
 etiology of, 367e-1t, 2514t, 2516
 in hemochromatosis. *See* Hemochromatosis
 transfusion-related, 138e-5, 631, 639, 667
 treatment of, 639
 in Wilson's disease. *See* Wilson's disease
- Iron poisoning, 473e-11t
- Iron sucrose, 629
- Iron supplements
 adverse effects of, 629
 for iron-deficiency anemia, 628–629, 629t
 oral, 628–629, 629t
 parenteral, 629
- Iron supply studies, 397
- Iron tolerance test, 629
- Irritable bowel syndrome (IBS), **1965**
 approach to the patient, 1967–1968
 clinical features of, **1965**
 abdominal pain/pressure, 105, 106t, 285, 1965
 altered bowel habits, 266, 272t, 1965
 gas and flatulence, 1965–1966
 nausea and vomiting, 259
 upper gastrointestinal, 1966
- diagnostic criteria for, 1965, 1966t, 1967t
 differential diagnosis of, 1968
 incidence of, 266
 pathophysiology of, 1966–1967, 1966t, 1967f
 postinfectious, 797, 855t, 1966–1967
 psychological features in, 1966
 severity spectrum of, 1970t
 treatment of, 272, 1968–1970, 1968t, 1970t
- Irritant contact dermatitis. *See* Contact dermatitis
- Irukandji syndrome, 2739
- IRV (inverse ratio ventilation), 1738, 1742. *See also* Mechanical ventilation
- Isaacs' syndrome, 462e-3, 619
- Ischemia
 acute limb, 1646
 cardiac. *See* Ischemic heart disease
 cerebral, 2559, 2561f. *See also* Ischemic stroke
 intestinal. *See* Intestinal ischemia
- Ischemic cascade, 1777–1778
- Ischemic colitis, 267, 1882f, 1957, 1980
- Ischemic heart disease, **1578**
 angina pectoris in. *See* Angina pectoris
 arrhythmia in, 1580
 asymptomatic (silent), 1580, 1592–1593
 cardiomegaly in, 1580
 cardiomyopathy and, 1580
 in chronic kidney disease, 1816–1817
 diagnosis of
 approach to, 1583f
 cardiac imaging, 1582, 1584f, 1585
 coronary arteriography, 1585–1586
 ECG, 1455–1457, 1455f, 1456f, 1457t, 1582, 1583f
 history, 1580–1581
 laboratory tests, 1582
 physical examination, 1581–1582
 stress testing, 1582, 1583f, 1584f, 1585t
 dyslipidemia and, 1588
 epidemiology of, 1578
 functional classification of, 1439–1440, 1440t, 1501, 1501t, 1581t
 heart failure in, 1579, 1590
 neck pain in, 122
 pathogenesis of, 1578–1580, 1580f. *See also* Atherosclerosis; Coronary artery disease (CAD)
 prognosis of, 1586
 risk factors for, 1587
 symptomatic, 1580
 treatment of, **1586**
 ACE inhibitors, 1590
 activity adaptation, 1587, 1587t
 aggravating conditions, 1586–1587
 algorithm for, 1591f
 antiplatelet drugs, 1590
 beta blockers, 1588–1590, 1589t
 CABG. *See* Coronary artery bypass grafting (CABG)
 calcium channel blockers, 1589–1590, 1589t
 enhanced external counterpulsation, 1592
 explanation and reassurance, 1586
 lipid-lowering therapy, 1588
 nicorandil, 1590
 nitrates, 1588, 1588t
 PCI. *See* Percutaneous coronary interventions (PCI), for revascularization
 principles, 1441
 ranolazine, 1590
 risk factor management, 1587
 stem cell therapy, 90e-2
 in women, 1588
- Ischemic penumbra, 2560
- Ischemic stroke, **2560**
 after TIA, 2568, 2569t
 clinical features of, 2572, 2572f
 with anterior cerebral artery occlusion, 2573, 2574f
 with anterior choroidal artery occlusion, 2573–2574, 2575f
 with basilar artery occlusion, 2577–2579, 2577–2580f

- Ischemic stroke, clinical features of (*Cont.*):
 with common carotid artery occlusion, 2574
 inferior pontine syndrome, 2577f
 with internal carotid artery occlusion, 2574, 2574f, 2575f
 medullary syndrome, 2576f
 midbrain syndrome, 2580f
 with middle cerebral artery occlusion, 2572–2573, 2573f
 midpontine syndrome, 2578f
 with posterior cerebral artery occlusion, 2574–2575, 2574f, 2575f
 with posterior inferior cerebellar artery occlusion, 2576–2577
 superior pontine syndrome, 2579f
 with vertebral artery occlusion, 2575–2576, 2575f
- etiology of, 2563, 2565t
 artery-to-artery, 2564–2565
 CADASIL, 2568
 cardioembolic, 2563–2564
 CNS vasculitis, 2568
 CNS vasculopathy, 2568, 2568f
 drug-related, 2568
 Fabry's disease, 432e-1, 2568
 fibromuscular dysplasia, 2567
 giant cell arteritis, 2567
 hypercoagulable disorders, 2565–2566
 leukoaraiosis, 2568
 moyamoya disease, 2568
 necrotizing arteritis, 2567–2568
 sickle cell anemia, 636, 2567
 small-vessel, 2564, 2567f
 venous sinus thrombosis, 2566–2567
- imaging in, 2580–2581, 2581f, 2582f
 pathophysiology of, 2560–2561, 2561f, 2563, 2564f
- prevention of, 2569
 anticoagulation therapy for, 2570–2571
 antiplatelet agents for, 291e-9, 747, 2566t, 2569–2570
 risk factor management in, 2569
- treatment of, 2561
 algorithm for, 2560f
 anticoagulants, 2563
 antithrombotic agents, 2563
 endovascular techniques, 2562–2563
 intravenous thrombolysis, 2561–2562, 2562t
 medical support, 2561
 neuroprotection, 2563
 rehabilitation, 2563
 stroke centers in, 2563
- Ischial bursitis, 2247
 Isfahan virus, 1310t
 Ishihara color plates, 197
 ISI (International Sensitivity Index), 405, 755, 1998
 Islet-1, 265e-11
 Islet amyloid polypeptide (IAPP), 2402
 Islet cell(s)
 adenoma of, 536
 pancreatic, 2403–2404
 regeneration of, 90e-3
 tumors of. *See* Pancreatic neuroendocrine tumors (pNETs)
- Islet cell autoantibodies, 2404
 Isoagglutinins, 138e-1
 Isobutyl-CoA dehydrogenase deficiency, 434e-3t
 Isocarboxazid, 2710t
 Isocyanate exposure, 1692t
 Isoflavones, 96e-6
 Isoforms, 427
 Isolated stem cells, 453
 Isolation aphasia, 177t, 178
 Isolation techniques, 916
 Isoleucine metabolism disorders, 434e-3t
 Isometheptene, 2592, 2592t
 Isomorphic (Koebner's) phenomenon, 347
 Isoniazid (INH)
 action of, 205e-2–3
 adverse effects of, 205e-3
 acid-base disorders, 318
 cutaneous, 353, 360
 fever, 138
 hepatotoxicity, 40, 205e-3, 1121, 2023, 2025t, 2028
 lupus syndrome, 2218t
 neuropathy, 205e-3, 2687t, 2689
 optic neuropathy, 203
 pellagra, 96e-4
 pure red cell aplasia, 668t
 pyridoxine for prevention of, 96e-4, 205e-3, 1116
 seizure, 2547t
 dosage of, 205e-3
 drug interactions of, 205e-3, 246e-3t
 genetic variations in response to, 39t, 40
 metabolism of, 34t
 for nontuberculous mycobacterial infections, 1131–1132
 overdose/poisoning with, 473e-5, 473e-12t
 pharmacology of, 205e-3
 resistance to, 205e-3, 1118
 for TB
 active infection, 205e-2–3, 205e-2t, 1115t, 1116–1117, 1116t
 latent infection, 205e-1t, 205e-2t, 1121
 prophylaxis in HIV infection, 1251t
 for tuberculous meningitis, 898
- Isoprinosine, 900
 Isopropyl alcohol poisoning, 319
 8-Isoprostane, 1674
 Isoproterenol, 1469, 1473, 1474
 Isosexual precocity, 2361
 Isosorbide, 1512t, 1588t, 1608, 1763
Isospora belli/isosporiasis. *See* *Cystoisospora belli* cystoisosporiasis
- Isotretinoin
 for acne vulgaris, 352
 adverse effects of, 352, 354, 380, 380f, 2218t
 for oral leukoplakia, 477
 for pityriasis rubra pilaris, 354t
- Isotypes, of immunoglobulins, 710
 Isovaleric acidemia, 434e-3t
 Isradipine, 1589, 1589t, 1649
 Israeli spotted fever, 1157
 ITAMs (Immunoreceptor tyrosine-based activation motifs), 372e-19, 2115
- Itaqui virus, 1309t
ITPA gene, 39, 39t
- Itraconazole
 adverse effects of, 246e-3t, 1331
 for *Aspergillus* infections, 1348, 1349t, 1681
 for *Aspergillus* prophylaxis, 1349
 for *Blastomyces* infections, 1338, 1339t
 for *Candida* infections, 233, 1345
 for *Coccidioides* infections, 1336
 contraindications to, 350
 for dermatophyte infections, 1358, 1358t
 drug interactions of, 34t, 42t, 246e-3t, 328
 for *Histoplasma* infections, 899, 1334, 1334t
 indications for, 1331
 for *Malassezia* infections, 1357
 for onychomycosis, 350t, 1358, 1358t
 for paracoccidioidomycosis, 1354, 1354t
 for penicilliosis, 1354t
 in pregnancy and lactation, 246e-3t
 prophylactic use, in HIV infection, 1252t
 resistance to, 1348
 for *Sporothrix* infections, 1354, 1354t
 susceptibility testing for, 150e-8
 for thrush, 233
 for tinea versicolor, 350
- IUDs. *See* Intrauterine devices (IUDs)
- Ivabradine, 1477, 1512
 Ivacaftor, 1699
- Ivermectin
 action of, 246e-9
 adverse effects of, 246e-4t, 246e-9, 1420, 1421, 1422
 for ascariasis, 1413–1414
 for cutaneous larva migrans, 1411t
 for gnathostomiasis, 1411t
 indications for, 246e-9
 for loiasis, 1422
 for lymphatic filariasis, 1419
 for onchocerciasis, 1421
 pharmacology of, 246e-9
 in pregnancy and lactation, 246e-4t
 resistance to, 246e-9
 for scabies, 2745
 for strongyloidiasis, 1416
 for trichuriasis, 1416
- IVF (in vitro fertilization), 444, 2360, 2389
 IVIg. *See* Intravenous immunoglobulin (IVIg)
 "Ivory vertebra," 426e-2
 Ixabepilone, 103e-12t, 103e-15
Ixodes spp. ticks. *See also* Tick(s)
 in arborviral encephalitis, 1316
 in *Babesia* infections, 1384. *See also* *Babesia* spp. infections
 characteristics and life cycle of, 1149, 2745, 2746f
 climate change and habitat of, 151e-6, 151e-6f
 in human granulocytotropic anaplasmosis, 1160
 in Lyme borreliosis, 1149, 2745. *See also* Lyme borreliosis (Lyme disease)
 in relapsing fever transmission, 1146. *See also* Relapsing fever
 removal of, 2745
- Ixodid (hard) tick, 2745
 Izumi fever, 1077
- ## J
- Jackhammer esophagus, 1906f
 Jacksonian epilepsy, 1427
 "Jacksonian march," 2542
 Jacobsen syndrome, 375e-2
 Jaffe reaction, 480e-2
 JAK (Janus family of protein tyrosine kinases), 372e-16, 401e-3, 2143
- JAK2 gene mutations
 in IBD, 1949, 1950t
 in MDS/MPN, 135e-6
 in polycythemia vera, 400, 672–673, 674, 675
- JAK2 inhibitors, 102e-3, 674
 JAK3 deficiency, 372e-16
 JAK3 gene mutations, 135e-3
 Jamaica bush tea, 283
 Jamestown Canyon virus, 1309t, 1315
 Janeway lesions, 784, 818f, 1443
 Jansen's chondrodystrophy, 2458
 Jansen's disease, 2475, 2476f
 Japanese encephalitis virus, 1306t, 1315
 Japanese encephalitis virus vaccine, 795, 795t
 Japanese spotted fever, 1157
 Jargon aphasia, 177
 Jarisch-Herxheimer reaction
 in leptospirosis, 1145
 in Lyme disease, 1153
 in Q fever, 1162
 in relapsing fever, 1147, 1148
 in syphilis, 1140, 1259
 in Whipple's disease, 1094
- Jaundice, 279
 in acute viral hepatitis, 2015, 2016
 approach to the patient, 280–281, 281f
 with ascites, 283
 breast milk, 2001
 in cardiovascular disease, 1442
 in cholelithiasis, 2084
 in cholestatic disorders, 283–285, 284t, 2080
 in chronic hepatitis, 2033
 in clinical drug trials, 2025
 differential diagnosis of, 279–280, 1877, 1877t
 drug-induced, 283, 283t, 284
 global considerations in, 285
 in hemolytic anemia, 649
 in hepatocellular conditions, 283, 283t
 history in, 282
 laboratory evaluation of, 280, 283
 in liver disease, 1991
 neonatal, 653, 654, 2001
 in pancreatic cancer, 554–555
 pathophysiology of, 280
 physical examination in, 282–283
 in sepsis/septic shock, 1755
- Jaw, osteonecrosis of, 242, 623, 2481, 2501

- JC virus infection
 in cancer patient, 488t, 490
 in HIV infection, 1266
 in progressive multifocal leukoencephalopathy, 899, 1962, 2669
 in transplant recipient, 923, 925, 1831
 Jefferson fracture, 441e-32f
 Jejunal vascular ectasia, 1883f, 1897f
 Jejunostomy tube feeding, 98e-8t, 261
 Jellyfish sting, 2738-2739, 2741f
 Jerk nystagmus, 210
 Jervell and Lange-Nielsen syndrome, 220, 222t, 444e-2t
 Jet lag disorder, 194
 Jigger, 2750
 Jimson weed, 473e-9t
 JK bacillus infection, 486t
 JMML (juvenile myelomonocytic leukemia), 135e-6
 Job's syndrome. *See* Hyper-IgE syndrome
 Jod-Basedow effect, 2300, 2302
 Joint(s)
 disorders of. *See* Arthritis; Musculoskeletal disorders; Osteoarthritis
 failure of, 2226-2227, 2227f
 malalignment of, 2221, 2228, 2228f
 in nutritional deficiencies, 463t
 position testing, 159, 160t
 prosthetic, infections in. *See* Prosthetic joint infections
 protective mechanisms of, 2226-2227
 Joint Commission, infection-control guidance and oversight, 911-912
 Joint replacement, 2233. *See also* Knee replacement
 Joint United Nations Program on HIV/AIDS (UNAIDS), 8, 12
 Jones criteria, 2152, 2152t
 Josamycin, 1157
 Joubert's syndrome, 451e-4t, 1851f, 1855
JPH3 gene mutations, 2622
 Jugular septic thrombophlebitis. *See* Lemierre's disease/syndrome
 Jugular venous pressure
 in heart failure, 286, 1504
 in hypovolemia, 298
 measurement of, 1443-1444, 1444f
 in shock, 1729
 Jumper's knee, 2220-2221
 Junctional bradycardia, ECG in, 278e-3f
 Junctional tachycardia, 1480-1481
JUN gene mutations, 101e-2t
 Junín/Argentinian hemorrhagic fever, 1320
 Junin virus, 1306t
 Juquitiba virus, 1308t
 Justice, 17e-2
 Juvenile arthritis, 373e-7-8
 Juvenile arthritis, HLA gene association with, 373e-8t
 Juvenile goiter, 2301
 Juvenile granulosa tumors, 594
 Juvenile myelomonocytic leukemia (JMML), 135e-6
 Juvenile myotonic epilepsy, 2544
 Juvenile-onset spondyloarthritis, 2178
 Juvenile Paget's disease, 426e-1
 Juvenile polyposis, 101e-6t, 538t
 Juvenile polyps, 537
 Juxtaglomerular apparatus, 332e-2f
- K**
 K8/18 (keratins 8 and 18), 2056
KALI gene mutations, 2256, 2363, 2364t
 Kala-azar, 245e-2, 245e-4t. *See also* Visceral leishmaniasis
 Kallmann syndrome, 215, 2256, 2363
 Kanagawa phenomenon, 1065
 Kanamycin, 205e-6, 930, 937t, 942
 Kaolin-pectin, 41, 246e-1t
 Kaposi's sarcoma
 clinical features of, 76e-5f, 364t, 367, 1268-1269, 1269f
 diagnosis of, 1269
 epidemiology of, 1242
 etiology of, 502, 1242
 HHV-8 infection and, 119e-1, 214e-7, 1194
 in HIV infection, 1268-1269, 1269f, 1269t, 1270t
 oropharyngeal, 240t
 pathogenesis of, 1242
 pericardial, 1255
 staging of, 1269, 1269t
 in transplant recipient, 924, 925
 treatment of, 1269-1270, 1270t
 Kaposi's sarcoma-associated herpesvirus (KSHV) infection. *See* Human herpesvirus-8 (HHV-8) infection
 Kappa receptors, 468e-1t
 Karnofsky Performance Scale, 472, 472t
 Karyomegalic tubulointerstitial nephritis, 1855, 1861
 Karyorrhexis, 102e-8
 Katayama syndrome, 1425, 1427
katG gene, 1106
 Kava, 43, 283
 Kawasaki disease
 clinical features of, 130t, 134, 363
 epidemiology of, 2192
 prognosis of, 2193
 rash in, 25e-7f, 130t, 362t, 363
 treatment of, 2193
 vasculitis in, 2192-2193
 Kayser-Fleischer ring, 436e-8f, 2519, 2520f, 2520t, 2625
 KCl. *See* Potassium chloride (KCl)
KCNH2 gene mutations, 1496t
KCNJ5 gene mutations, 306
KCNJ18 gene, 462e-18
KCNQ1 gene mutations, 1496t
KCNQ2 gene mutations, 2545t
 Kearns-Sayre syndrome (KSS), 451e-3t
 clinical features of, 85e-6, 85e-6t, 462e-15, 2483
 AV conduction block, 1471, 1471t
 endocrine, 2324t, 2348
 ocular, 206, 207
 SA node dysfunction, 1467, 1467t
 diagnosis of, 462e-15
 genetic considerations in, 451e-3t, 462e-15, 2324t
 mitochondrial DNA mutation in, 85e-6, 85e-6t, 462e-15, 2348
 treatment of, 462e-15
 Kegel exercises (pelvic muscle exercises), 81
 Kell blood group system, 138e-1t, 138e-2
 Kelley-Seegmiller syndrome, 431e-5
 Kelling-Madlener procedure, 1926
 Keloid, 76e-4f, 341t
 Kemerovo virus, 1308t
 Kennedy's disease. *See* Spinobulbar muscular atrophy (Kennedy's disease)
 Kenney-Caffey syndrome, 2483
 Kenya tick typhus, 1157
 Keratic precipitates, 200
 Keratin, 372e-24t
 Keratinocyte(s), 386
 Keratinocyte growth factor therapy, for mucositis, 103e-25
 Keratins 8 and 18 (K8/18), 2056
 Keratitis, 200-201
Acanthamoeba, 1368, 1368f
Aspergillus, 1348
Fusarium, 1355-1356
 HSV, 200, 215e-9, 1178, 1182t
Nocardia, 1086, 1087, 1087t
Onchocerca, 1421
P. aeruginosa, 1045t, 1046
 pigmented mold, 1355
Scedosporium, 1356
 Keratoacanthoma, 76e-6f, 501, 501f
 Keratoconjunctivitis, 199-200, 1208
 Keratoconjunctivitis sicca (dry eye)
 clinical features of, 199
 diagnosis of, 2166
 differential diagnosis of, 2167t
 etiology of, 199
 in sarcoidosis, 2207
 in Sjögren's syndrome, 2166, 2167t
 in SLE, 2129t, 2131
 in systemic sclerosis, 2162
 treatment of, 199, 2168f, 2169
 Keratoderma blenorrhagica, 2174
 Keratolytic agents, 351
 Keratomalacia, 96e-6
 Keratosis pilaris, 341t, 342f
 Kerion, 350, 355t
 Kerley B lines, 476e-3, 1552
 Kernicterus, 2001t, 2002
 Kernig's sign, 883
 Kernohan-Woltman sign, 1772
 Keshan's disease, 96e-10, 1563
 Ketamine
 abuse of, 168, 469e-5
 adverse effects of, 1741
 antidepressant effects of, 465e-2, 2717
 overdosage/poisoning with, 473e-3
 for sedation during mechanical ventilation, 1741
 Ketanserin, 566
 Ketoacidosis
 alcoholic, 299, 318-319
 breath odor in, 242
 diabetic. *See* Diabetic ketoacidosis (DKA)
 nausea and vomiting in, 259
 α -Ketoacidic acidemia, 434e-3t
 Ketoconazole
 for adrenal blockade in breast and prostate cancer, 103e-16
 adverse effects of, 246e-3t, 310, 2273
 for *Blastomyces* infections, 1339t
 for candidiasis, 233
 for Cushing's syndrome, 2273, 2318
 for cutaneous leishmaniasis, 1393
 CYP3A and, 34t, 42t
 drug interactions of, 34t, 42t, 246e-3t, 328, 1590, 1921
 for ectopic ACTH syndrome, 611
 for hypercalcemia, 314
 for male precocious puberty, 2362
 in pregnancy and lactation, 246e-3t
 for seborrheic dermatitis, 347
 shampoo, for tinea, 355t
 for tinea versicolor, 350
 Ketolides, 931, 931t, 934. *See also* Telithromycin
 Ketoprofen, 338
 Ketorolac, 91, 91t, 199, 478e-4t, 635
 Khat leaves, 469e-3-4
 KI. *See* Potassium iodide (KI)
 Ki-67, 560, 2322
 Kidd blood group system, 138e-1t, 138e-2
 Kidney. *See also* Nephron(s)
 calcium absorption/excretion by, 2457
 congenital abnormalities of, 333e-1, 1855
 disease/failure
 acute. *See* Acute kidney injury (AKI)
 chronic. *See* Chronic kidney disease (CKD)
 glomerular. *See* Glomerular diseases
 nephrolithiasis. *See* Nephrolithiasis
 tubulointerstitial diseases. *See* Tubulointerstitial diseases
 vascular. *See* Renovascular disease
 drug clearance in, 935
 embryological development of, 332e-1-2, 332e-1f, 333e-1
 erythropoietin production in, 392, 392f
 hypoplastic, 333e-1
 in hypothermia, 478e-2t
 ischemic preconditioning in, 333e-3-4, 333e-4t
 phosphate absorption/excretion by, 2458
 potassium absorption/excretion by, 64e-2f, 304-305, 332e-5f, 332e-9
 renin-secreting tumors of, 1614
 sodium absorption/excretion in, 64e-2f, 296-297, 296f, 297f, 332e-3-6f, 332e-10-11, 332e-10f
 uric acid metabolism in, 431e-1, 431e-2
 vasculature of, 332e-2, 332e-2f, 1627
 in vitamin D metabolism, 2464, 2464f
 water absorption/excretion by, 64e-2f, 295-296, 296f, 332e-3-6f, 332e-9-10, 332e-10f

- Kidney biopsy. *See* Renal biopsy
- Kidney cancer. *See also* Renal cell carcinoma (RCC)
- deaths from, 467t, 471t, 578
 - genetic considerations in, 101e-2t
 - hereditary papillary, 101e-5, 101e-6t
 - incidence of, 467t, 471t, 577, 578
 - infections in, 484t
 - paraneoplastic syndromes in, 609t, 612t
- Kidney injury molecule-1 (KIM-1), 1808
- Kidney transplantation
- donor selection for, 1826–1827, 1826t
 - graft survival rates after, 1826, 1826t
 - HLA typing in, 1826–1827
 - for lupus nephritis, 2134
 - mortality rates after, 1826, 1826t
 - presensitization in, 1827
 - racial/ethnic disparities in referral for, 16e-1, 16e-2f
 - recipient management
 - chronic lesions of graft, 1831
 - complications, 1831
 - early care, 1829, 1830f
 - induction therapy, 1828
 - infections, 924–926, 924t, 1829–1831, 1830t
 - maintenance immunosuppressive therapy, 1828–1829, 1829t, 2764t, 2766t
 - malignancy, 1831
 - rejection episode, 1829
 - recipient selection, 1821, 1826
 - rejection in, 372e-30, 1827, 1828f
 - results of, 1825–1826, 1826t
 - in sickle cell anemia, 636
 - tissue typing and immunogenetics in, 1826–1827, 1828f
- Kikuchi's disease, 1109
- Killer cell immunoglobulin-like activating or inhibitory receptors (KIRs), 372e-7, 372e-15t, 373e-4–5
- Killian's triangle, 1903, 1903f
- KIM-1 (kidney injury molecule-1), 1808
- Kimmelstiel-Wilson nodules, 62e-8f, 1844
- Kimura's disease, 1109
- Kinase inhibitors, 102e-3–4, 102e-4f, 103e-18t, 103e-19, 2307. *See also specific drugs*
- Kinases, 102e-3
- Kindler's syndrome, 357, 2513
- Kingella kingae*, 183e-1, 183e-2t. *See also* Infective endocarditis, HACEK group
- Kir 2.1* gene, 462e-18
- Kir 2.6*, 305
- Kirby-Bauer method, 150e-9
- KIR* gene, 1241t
- KISS1* gene mutations, 401e-4, 2363, 2364t
- KISS1R* gene mutations, 2377
- Kissing bug bite, 2750
- Kisspeptin, 401e-4, 2357, 2363, 2377
- KIT* gene mutations, 135e-8, 680, 680t
- Kit ligand, 89e-2
- Klatskin tumor, 552
- Klebsiella* spp., 1025, 1031
- Klebsiella granulomatis*, 198e-1, 198e-2. *See also* Donovanosis
- Klebsiella* spp. infections, **1031**
- abdominal, 288, 1032, 1032f
 - bacteremia, 1033
 - burn-wound, 166e-1
 - in cancer patient, 486t
 - cellulitis, 1032–1033
 - clinical features of, 266t, 1031
 - diagnosis of, 1032f, 1033
 - diarrhea, 266t, 855
 - ecthyma gangrenosum, 369
 - epidemiology of, 1025
 - global considerations in, 1031–1032
 - health care–associated, 776, 1031
 - liver abscess, 850
 - pneumonia, 915, 1032
 - rhinoscleroma, 1352
 - sepsis and septic shock, 1752
 - sinusitis, 226
 - in transplant recipient, 919
 - treatment of, 940t, 1033
 - urinary tract, 862, 1032, 1871
- Klebsiella oxytoca*, 1031
- Klebsiella pneumoniae*
- antibiotic resistance in, 933
 - carbapenemase-producing (KPCs), 916, 918
 - epidemiology of, 1025, 1031
 - global considerations in, 1031–1032
 - hypervirulent, 1025, 1031–1032, 1032f
- Klf4* gene, 454
- Klinefelter's syndrome, **2350**
- clinical features of, 83e-5, 1654, 2350, 2351t, 2360, 2365, 2366
 - diseases associated with, 2126, 2365
 - genetic considerations in, 83e-5, 2365
 - mediastinal germ cell tumor in, 588
 - pathophysiology of, 2350
 - prevalence of, 2254t
 - screening/testing for, 2254t
 - treatment of, 2350–2351, 2365
- Klippel-Trénaunay syndrome, 1651, 1654
- KLLN* gene mutations, 2344
- Kluyvera*, 1036
- Knee(s)
- bursitis of, 2248
 - CPPD arthropathy of, 2235
 - gout tophi of, 436e-2f
 - iliotibial band syndrome, 2248–2249
 - jumper's, 2248
 - osteoarthritis of, 2226, 2228–2229, 2228f, 2230, 2230f
 - pain in, 2221–2222, 2230
- Knee replacement
- for osteoarthritis, 2233
 - racial/ethnic disparities in rates of, 16e-2, 16e-3f
- Knock-knees, 2221, 2228, 2228f
- Knock-out/knock-in mice, 432
- Knudson two-hit hypothesis/model, 101e-2, 431t, 440
- Koebner's (isomorphic) phenomenon, 347, 1269
- KOH (potassium hydroxide) wet-mount preparation, 150e-1, 341
- Koilonychia, 627
- Kokobera virus, 1306t, 1313
- Kommerell's diverticulum, 1637
- Koplik's spots, 25e-1f, 127, 362, 1297. *See also* Measles (rubeola)
- Korotkoff sounds, 1622
- Korsakoff's psychosis/syndrome, 1783, 2607, 2724
- Kostmann's syndrome, 418, 678
- KPCs (*Klebsiella pneumoniae*, carbapenemase-producing), 916, 918
- Krabbe's disease, 432e-4t, 441e-41f
- Kraits, 2734. *See also* Snakebites
- KRAS* gene mutations
- in AML, 680
 - cancers associated with, 101e-2t
 - in chronic myelomonocytic leukemia, 135e-6
 - in colorectal cancer, 101e-1f, 101e-8, 101e-9f, 537–538
 - drug response and, 39t, 41
 - in lung cancer, 507t, 508, 509, 509f, 521
 - in pancreatic cancer, 554
 - syndromes associated with, 436t
- Krukenberg's tumors, 534, 592, 593
- KSHV (Kaposi's sarcoma-associated herpesvirus) infection. *See* Human herpesvirus-8 (HHV-8) infection
- KSS. *See* Kearns-Sayre syndrome (KSS)
- Ku-DNA-protein kinase, 372e-24t
- Kugelberg-Welander disease, 2636
- Ku-nucleoprotein La phosphoprotein, 372e-24t
- Kupffer cells, 1990
- Kuru, 453e-1t, 453e-2
- Kussmaul respiration, 317, 1775, 2418, 2420
- Kussmaul's sign, 286, 1444, 1566, 1576
- Kveim-Siltzbach procedure, 2210
- Kwashiorkor (acute disease- or injury-related malnutrition), **460**
- cardiac manifestations of, 290e-1, 290e-1t
 - classification of, 98e-1, 98e-1t
 - clinical features of, 460, 460t, 463t
 - diagnosis of, 460, 460t
 - global issues, 461
 - physiologic characteristics of, 460t
 - prognosis of, 460t
 - small-intestinal mucosal biopsies in, 1940t
 - specialized nutritional support for. *See* Specialized nutritional support
- Kyasanur Forest disease virus, 1307t, 1322
- Kyphoplasty, 2503
- Kyphoscoliosis, 47e-2, 118
- ## L
- Labetalol
- for aortic dissection, 1641
 - for autonomic storm, 2643
 - for hypertension, 46, 1624, 1624t, 1787
 - for hypertensive emergencies, 1627, 1627t
 - for ischemic heart disease, 1589t
 - overdosage/poisoning with, 473e-9t
 - for preeclampsia, 46
 - for sympathomimetic overdosage/poisoning, 473e-8t
- Laboratory tests. *See* Clinical laboratory tests; Infectious diseases, laboratory diagnosis of
- Laboratory worker's lung, 1681t
- Labyrinthectomy, 219
- Labyrinthine infarction, 220
- Labyrinthitis, 62, 259, 260t
- Lacosamide, 2553t, 2558f
- Lacrimal glands, 199, 391e-2, 391e-2f
- Lacrimation, with headache, 107
- La Crosse virus, 1309t, 1314–1315
- Lactase, 1937
- Lactase deficiency
- diagnosis of, 272
 - diarrhea in, 269
 - disease, 263
 - epidemiology of, 1937t
 - indigestion in, 262
 - primary, 1937
 - secondary, 1937
 - treatment of, 263, 269, 272
- Lactate dehydrogenase (LDH)
- in ascitic fluid, 287–288
 - in melanoma, 498
 - in testicular cancer, 588
 - as tumor marker, 473t
- Lactate dehydrogenase (LDH) deficiency, 433e-2t, 462e-14
- Lactation. *See* Breast-feeding
- Lactic acidosis
- approach to the patient, 318
 - in cancer, 1794
 - etiology of, 318
 - in malaria, 1373–1374
 - treatment of, 318, 322
- Lactobacillus* spp., 86e-5, 874, 875
- Lactoferrin, 416
- Lactophenol cotton blue stain, 150e-1
- Lactose, 1937
- Lactose breath test, 272
- Lactose intolerance/malabsorption. *See* Lactase deficiency
- Lactulose, 62t, 274, 1970t, 2066
- Lacunar cells, 81e-11f
- Lacunar infarction, 441e-17f, 2565, 2573
- LAD. *See* Leukocyte adhesion deficiency (LAD)
- Lady Windermer's syndrome, 1130
- Laennec's cirrhosis, 282
- Lafora's disease (progressive myoclonus epilepsy), 2545t
- Laguna Negra virus, 1308t
- Laing's distal myopathy, 462e-12, 462e-13t
- Lambert-Eaton myasthenic syndrome (LEMS)
- vs. botulism, 989
 - channelopathies in, 444e-2t
 - clinical features of, 2703
 - diagnosis of, 2703
 - immunopathogenesis of, 372e-25t
 - in lung cancer, 511, 616, 2703
 - vs. myasthenia gravis, 2703
 - reflex responses in, 154
 - treatment of, 2703

- Lambrolizumab, 103e-21
 LAMB syndrome. *See* Carney's syndrome/complex (LAMB/NAME syndrome)
 Lamellar bone, 2466
 Laminar-flow rooms, for neutropenic patients, 492
 Laminectomy, for neck pain, 123
 Laminin, 462e-6
 Lamin proteins, 1554
 Lamivudine (3TC)
 adverse effects of, 380, 1274t, 2034
 for chronic HBV infection, 215e-4t, 215e-9, 2034–2035, 2037t, 2038f, 2074
 for HBV prophylaxis in transplant recipient, 927
 for HIV infection, 1274t, 1276–1277
 molecular structure of, 1273, 1276f
 resistance to, 215e-9, 1276, 1279f, 2034–2035, 2040t, 2074
 Lamotrigine
 adverse effects of
 common, 2718t
 cutaneous, 381, 2552
 fever, 138
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 neurologic, 2554t
 SJS or TEN, 383, 2718t. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
 systemic, 2554t
 drug interactions of, 2554t
 for epilepsy, 2552, 2554t, 2556
 for mood disorders, 2718, 2718t
 overdosage/poisoning with, 473e-10t
 pharmacology of, 2554t
 for SUNCT/SUNA prevention, 2596
 for trigeminal neuralgia, 2646
 LAMP2 deficiency (Danon's disease), 432e-4t, 433e-5, 1566
 LAMP2 gene mutations, 1555t, 1566
 Lancinations, 158
 Langerhans cell histiocytosis (histiocytosis X), 135e-9, 2257, 2264
 Langerhans cell sarcoma, 135e-9
 Language
 assessment of, 37e-1v, 177, 2537
 perisylvian network for, 177
 Language barriers, 16e-4
 Laninamivir, 215e-5
 Lanreotide
 for acromegaly, 2271
 adverse effects of, 567, 574
 for carcinoid syndrome, 567
 for gastrinoma, 2337
 for glucagonoma, 570
 for insulinoma, 569
 for NETs, 574
 structure of, 566f
 for VIPoma, 571
 Lansoprazole, 263, 1920, 1920t, 2337
 Lanthanum, 1816
 Laparoscopic adjustable gastric banding, 2398, 2398f
 Laparoscopic adrenalectomy, 2319
 Laparoscopic sleeve gastrectomy, 2398, 2398f
 Laparoscopic Ventral Rectopexy (LVR), for rectal prolapse, 1974, 1975f
 Laparoscopy, 103e-3, 877–878
 Laparotomy, 1980
 Lapatinib
 action of, 102e-5t, 103e-19
 adverse effects of, 103e-26, 621
 for breast cancer, 102e-5t, 529
 genetic variations in response to, 39t
 in pregnancy, 124e-4
 Laplace's law, 265e-8
 LA-PTT, 406
 LARC/MIP-3a/Exodus-1, 372e-12t
 Large granular lymphocytes, 81e-2, 372e-1, 372e-6. *See also* Natural killer (NK) cells
 Laron-type dwarfism, 94e-4, 375e-2
 Laropiprant, 96e-4
 Larva currens, 1415
 Larva migrans
 cutaneous
 clinical features of, 829, 1412
 diagnosis of, 245e-3t, 245e-5t
 pathogenesis of, 1412
 from pentastomes, 2747
 treatment of, 1411t
 ocular, 1411
 visceral
 clinical features of, 152e-6t, 1411
 diagnosis of, 152e-6t, 245e-3t, 245e-6t, 1411
 epidemiology of, 1411
 incubation period for, 152e-6t
 pathogenesis of, 1411
 in returning war veterans, 152e-6t
 treatment of, 152e-6t, 246e-8, 246e-9, 1411–1412, 1411t
 Laryngeal cancer, 503. *See also* Head and neck cancer
 Laryngeal nerve palsy, 2649
 Laryngitis, 234
 Laryngotracheal stricture, 2203
 Laser iridectomy, 200
 Laser photocoagulation, 2424
 Laser therapy
 for basal cell carcinoma, 501
 for hair removal, 334
 for hemoptysis, 247
 for neck pain, 123
 for sea urchin spine removal, 2740
 for warts, 1201, 1201t
 LASIK, 195
 Lasofofifene, 531
 Lassa fever, 215e-6, 782–783, 1306t, 1320–1321
 Lassaviruses, 1304. *See also* Rabies
 Latency, 214e-7
 Latency-associated transcripts (LATs), of HSV, 1175
 Latent tuberculosis infection (LTBI). *See* Tuberculosis (TB), latent infection
 Lateral epicondylitis, 2249
 Lateral femoral cutaneous neuropathy (meralgia paresthetica), 49, 463e-5, 463e-5f, 2692
 Lateral geniculate body, 195
 Lateral medullary stroke syndrome, 2576f, 2577
 Lateral sclerosis, 2632, 2659
 Latex agglutination assay, 150e-2, 888
 α -Latrotoxin, 2748
 LATs (latency-associated transcripts), of HSV, 1175
 Laugier-Hunziker syndrome, 359
 Laurence-Moon-Biedl syndrome, 415e-5t
 Laurence-Moon syndrome, 2364
 Laxatives
 abuse of, 268, 270, 305, 307
 adverse effects of, 267, 269, 307
 for constipation, 61, 62t, 273, 274
 for internal radionuclide contamination, 263e-6
 LCAT deficiency, familial, 2445
 LCK gene mutations, 101e-3t
 LCM (lymphocytic choriomeningitis), 891, 893, 908t, 1306t, 1317–1318
 LCZ696, 1507
 LDH. *See* Lactate dehydrogenase (LDH)
 LDL. *See* Low-density lipoprotein (LDL)
 Lead oxide exposure, 472e-3
 Lead poisoning
 vs. ALS, 2633
 clinical features of, 472e-2t, 2607–2608
 diagnosis of, 472e-2t, 2689, 2766t
 hemolytic anemia in, 659
 hypertension and, 472e-1
 metabolism of, 472e-2t
 nephropathy in, 1856t, 1862
 neuropathy in, 2688t, 2689
 optic neuropathy in, 203
 oral manifestations of, 240t
 peripheral blood smear in, 81e-5f
 pregnancy-associated risks, 472e-1
 sources of, 472e-2t
 treatment of, 472e-2t, 2608
 Lead-time bias, 479, 509
 Leaky epithelia, 332e-3
 Leber's hereditary optic neuropathy (congenital amaurosis)
 clinical features of, 85e-5, 85e-6t, 202–203, 1850t
 diagnosis of, 85e-5, 85e-6–7
 gene therapy for, 91e-3
 genetic considerations in, 202–203, 1850t
 genetic counseling in, 85e-9
 mitochondrial DNA mutation in, 85e-5, 85e-6–7, 85e-6t
 treatment of, 206
 visual fields in, 197
 Leboombo virus, 1308t
 Lechiguanas virus, 1308t
 Lectins, 89e-1
 Ledipasvir, 2049
 Leech infestations, 2747
 Leflunomide
 adverse effects of, 2146t, 2687t
 for CMV infection, 922
 monitoring during treatment with, 2146t
 for psoriatic arthritis, 2177
 pyrimidine metabolism and, 431e-6
 for rheumatoid arthritis, 2145, 2146t
 for SLE, 2133
 Left bundle branch, 1471
 Left bundle branch block
 conditions associated with, 1454–1455
 ECG in, 269e-16f, 278e-9f, 1454, 1455f
 heart sounds in, 1447f
 Left coronary artery, anomalous origin of, 1523
 Left ventricular aneurysm, 1610
 Left ventricular assist devices (LVADs)
 as bridge to cardiac transplantation, 1518
 components of, 1519f
 for heart failure, 1518
 for pulmonary edema, 1763
 results of, 1519
 types of, 1517–1518
 Left ventricular dysfunction
 cardiogenic shock in, 1759
 diagnosis of
 CMR, 270e-6–7, 270e-26v
 echocardiography, 270e-6–7, 270e-8f
 ventriculography, 1463–1464, 1464f
 in heart failure, 1502, 1502f. *See also* Heart failure (HF)
 in mitral regurgitation, 1543
 in myocardial infarction, 1607
 Left ventricular hypertrophy
 in chronic kidney disease, 1817
 ECG in, 269e-15f, 1454, 1454f
 hypertension and, 269e-15f, 1615
 with myocardial ischemia, 1579
 Left ventricular noncompaction, 1555t, 1564
 Left ventricular pressure-volume relationship, 265e-10, 265e-10f
 Left ventricular remodeling, 1502–1503, 1503t, 1607. *See also* Heart failure (HF)
 Left ventricular stroke work, 1748t
 Legal issues
 advance care planning, 58–59
 ethanasia and physician-assisted suicide, 67–68
 incompetent patient, 17e-1–2
 mandated reporting, 17e-1
 Leg elevation, for stasis dermatitis/ulceration, 346, 354t
 Legionella spp.
 adherence of, 145e-2t
 ecology of, 1014–1015
 genome of, 1016
 host response to, 1015–1016
 laboratory identification of, 150e-5t, 150e-8, 1017, 1020t
 microbiology of, 1014
 Naegleria and, 1367
 serogroups of, 1014
 virulence of, 1016
 Legionella spp. infections, 1014
 in cancer patient, 489, 489t
 clinical features of, 1016

- Legionella* spp. infections (Cont.):
 community-acquired, 1015
 diagnosis of
 chest radiograph, 1016, 1017f
 clinical cues, 1016f
 collection and transport of specimens, 150e-5t
 CT, 1016, 1018–1019f
 laboratory tests, 1017, 1020t
 nucleic acid tests, 150e-8, 1017
 urinary antigen tests, 806, 1017
 epidemiology of, 1015
 etiology of, 1014–1015
 extrapulmonary, 1016, 2198
 hospital-acquired, 1015, 1020
 pathogenesis of, 1015–1016
 prevention of, 917–918, 1020
 prognosis of, 1020
 risk factors for, 805
 in transplant recipients, 1015
 treatment of, 1017, 1020t
- Legionnaire's disease. *See* *Legionella* spp. infections
- Leigh's syndrome, 85e-6, 85e-6t, 441e-40f, 462e-16
- Leiomyoma
 small intestinal, 536
 uterine, 596
- Leiomyosarcoma. *See also* Soft tissue sarcoma
 gastric, 536
 pathology of, 119e-1
 small intestinal, 537
 treatment of, 119e-2
- Leishmania* spp.
 adherence of, 145e-3
 characteristics of, 1387, 1388t
 entry into host, 145e-1, 372e-22
 epidemiology of, 1388t
 geographic distribution of, 1388t
L. aethiops, 1388t, 1393
L. braziliensis, 245e-4t, 1388t, 1393
L. donovani, 245e-4t, 1388t. *See also* Visceral leishmaniasis
L. guyanensis, 1393
L. infantum, 1388, 1388t
L. major, 1388t, 1392, 1393
L. mexicana, 1388t, 1392, 1393
L. tropica, 245e-4t, 1388t, 1392
 laboratory identification of, 245e-2, 245e-4t, 245e-6t
 life-cycle hosts of, 245e-4t
 life cycle of, 1387, 1388t
- Leishmania* spp. infections, **1387**
 cutaneous
 clinical features of, 152e-3t, 1392, 1392f
 diagnosis of, 152e-3t, 245e-4t, 1392
 differential diagnosis of, 1392
 epidemiology of, 152e-3t, 1388t, 1389f, 1392
 immunopathogenesis of, 1392
 incubation period for, 152e-3t
 prevention of, 152e-3t
 in returning war veterans, 152e-3t, 152e-8
 treatment of, 152e-3t, 152e-7, 1393
 diffuse cutaneous, 1388t, 1393
 epidemiology of, 152e-8, 245e-4t, 1387, 1388t, 1389f
 mucosal, 1388t, 1393–1394, 1393f
 prevention of, 1394
 visceral. *See* Visceral leishmaniasis
- Lemierre's disease/syndrome
 clinical features of, 233–234, 784, 1097
 mortality from, 784
 pathogenesis of, 231, 814, 1097
 treatment of, 234
- LEMS. *See* Lambert-Eaton myasthenic syndrome (LEMS)
- Lenalidomide
 action and targets of, 103e-19
 adverse effects of, 671, 716–717
 for amyloidosis, 722–723
 for multiple myeloma, 716–717
 for myelodysplasia, 671
- Lenègre disease, 1471t
- Length-biased sampling, 479, 509
- Lennox-Gastaut syndrome, 2544
- Lenticulus, 2514
- Lenticulostriate arteries, 2572
- Lentigo, 359, 360t
- Lentiviruses, 91e-1t, 91e-2, 225e-2, 1216, 1216f.
See also HIV
- LEOPARD syndrome, 359, 360t
- LEP gene mutations, 415e-4t, 2364t
- Lepirudin, 728, 753–754, 753t, 754
- LEPr gene mutations, 415e-4t, 2364t
- Lepromatosis, diffuse, 1124
- Leprosy, **1122**
 borderline, 1123t
 clinical features of, 152e-6t, 1124–1125
 cutaneous, 76e-11f, 357t, 829, 1123t, 1124f, 1125f
 neuropathy, 2648, 2684
 renal, 1836t, 1849
 complications of, 228, 1125–1126
 diagnosis of, 1126
 differential diagnosis of, 1126
 epidemiology of, 1123
 global considerations in, 1123, 1124, 2684
 incubation period for, 152e-6t, 1124
 lepromatous, 76e-11f, 829, 1123t, 1124–1125, 1124f, 2684
 multibacillary, 1127
 paucibacillary, 1127
 prevention and control of, 1128
 reactional states in, 1125, 1125f, 1127–1128
 in returning war veterans, 152e-6t
 transmission of, 1123–1124
 treatment of, 152e-6t, 2684
 antimicrobial agents, 1126
 reactions to, 1127–1128, 2684
 regimens, 1126–1127, 1127t
 tuberculoid, 357t, 359, 1123t, 1124, 1124f, 2684
- Leptin
 actions of, 415e-3, 415e-3f
 deficiency of, 2256
 in hypogonadotropic hypogonadism, 336, 2364
 in male pubertal development, 2357–2358
 in obesity, 415e-4t, 415e-5
 resistance to, 2450
- Leptin receptors, 415e-3, 415e-4f, 415e-4t
- Leptomeningeal metastases, 605, 606f
- Leptospira*/leptospirosis (Weil's syndrome), **1140**
 clinical features of, 129t, 285, 908t, 1144
 diagnosis of, 150e-1, 152e-2t, 772, 1144–1145
 differential diagnosis of, 1145
 epidemiology of, 1141–1142
 etiologic agent of, 1140–1141, 1141f, 1142f
 global considerations in, 1141–1142
 incubation period for, 152e-2t, 1144
 pathogenesis of, 1142–1143, 1143f, 1144f
 prevention of, 152e-2t, 1145, 1145t
 prognosis of, 1145
 in returning war veterans, 152e-2t
 treatment of, 152e-2t, 1145, 1145t
- Lerich's syndrome, 1642
- LES. *See* Lower esophageal sphincter (LES)
- Lesch-Nyhan syndrome, 431e-5
- Leser-Trelat sign, 359, 535
- Letermovir, 215e-9
- Lethal midline granuloma. *See* Extranodal NK/T-cell lymphoma, nasal type
- Letrozole, 103e-15, 337, 1795
- Leucine metabolism disorders, 434e-3t
- Leucovorin, 103e-11t, 103e-14
 for colorectal cancer, 543
 for gastric cancer, 535
 for gestational trophoblastic disease, 124e-5, 597
 for osteosarcoma, 119e-3
 for pancreatic cancer, 556t
 for PCP prophylaxis, 1251t, 1362t
 for toxoplasmosis prophylaxis, 1251–1252t
- Leukapheresis, for CML, 694
- Leukemia
 acute lymphoid. *See* Acute lymphoid leukemia (ALL)
 acute myeloid. *See* Acute myeloid leukemia (AML)
 acute promyelocytic. *See* Acute promyelocytic leukemia (APL)
 adult T cell. *See* Adult T-cell leukemia/lymphoma
 aggressive NK cell, 135e-3
 chemotherapy-associated, 103e-8
 chromosome abnormalities in, 102e-7
 chronic eosinophilic, 135e-7–8, 135e-7t, 422, 672
 chronic lymphoid. *See* B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma
 chronic myeloid. *See* Chronic myeloid leukemia (CML)
 chronic neutrophilic, 135e-4–5, 135e-5t, 672
 deaths from, 467t, 469f, 470t
 definition of, 102e-1, 103e-1
 gene therapy for, 91e-4
 hairy cell. *See* Hairy cell leukemia
 incidence of, 467t
 infections in, 486
 joint involvement in, 2247
 late consequences of treatment, 623, 624t
 mast cell, 2120
 ocular invasion by, 206
 oral manifestations of, 236, 237
 paraneoplastic syndromes in, 612t
 polycythemia vera and, 674
 precursor B-cell lymphoblastic, 91e-4, 701
 precursor T-cell lymphoblastic, 702t
 splenomegaly in, 411
 T-cell large granular lymphocytic, 135e-3
 T-cell prolymphocytic, 135e-3
 viruses and, 101e-7
- Leukemia cutis, 364t, 366
- Leukemia inhibitory factor, 372e-11t, 455
- Leukemic cell lysis pneumopathy, 1793
- Leukemoid reaction, 418–419
- Leukoarthritis (diffuse white matter disease), 174f, 2568, 2602
- Leukocidins, 145e-7
- Leukocyte(s), 62e-12f, 81e-1–2, 413. *See also* Basophil(s); Eosinophil(s); Lymphocyte(s); Monocyte(s); Neutrophil(s)
- Leukocyte adhesion deficiency (LAD)
 cellular/molecular defects in, 419, 420t, 2106
 clinical features of, 375e-1, 419, 420t
 diagnosis of, 420t, 2106
 subtypes of, 419, 420t
 treatment of, 2106
- Leukocyte common antigen, 120e-2t
- Leukocyte esterase test, 865
- Leukocyte function-associated antigen (LFA-1), 372e-24t
- Leukocyte Ig-like receptor (LIR), 373e-4
- Leukocyte telomere length (LTL), 94e-3
- Leukocytoclastic (cutaneous small-vessel) vasculitis, 25e-11f, 76e-15f, 133t, 135, 364t, 368
- Leukoderma, 357, 357t, 358t, 359
- Leukodystrophy(ies), 432e-4t, 2608
- Leukoencephalopathy
 progressive multifocal. *See* Progressive multifocal leukoencephalopathy (PML)
 radiation therapy-related, 607
 reversible posterior, 1793, 2568, 2584
- Leukoerythroblastosis, 671. *See also* Primary myelofibrosis
- Leukopenia. *See* Neutropenia
- Leukoplakia, 342f
 hairy, 76e-9f, 240t
 oral, 46e-3f, 76e-9f, 237, 240t, 342f, 477, 503
 sublingual, 46e-2f
- Leukostasis, 1793
- Leukotrienes, 87, 372e-15t, 1746
- Leuprolide, 103e-15, 585, 2132

- Levamisole, 727t, 2184, 2218t
 Lev disease, 1471t
 Levetiracetam
 adverse effects of, 2554t, 2556
 for cyclic vomiting syndrome, 261
 for dementia, 175
 for epilepsy, 2554t
 for myoclonus, 2623
 overdosage/poisoning with, 473e-10t, 2765t
 for status epilepticus, 2558f
 therapeutic monitoring of, 2765t
 Levine's sign, 100, 1580
 Levocardia, isolated, 1528
 Levodopa
 action of, 2614, 2614f
 adverse effects of, 96e-4, 2614, 2618
 for dystonia, 2621
 for PD, 2613–2615, 2614f, 2616t, 2618
 Levofloxacin
 action of, 931t, 932–933, 943
 adverse effects of, 2023
 for *C. trachomatis* infection, 1171
 for community-acquired pneumonia, 767t, 808t
 for epididymitis, 872
 in *H. pylori* eradication, 1042t
 for HACEK endocarditis, 183e-1, 183e-2t
 for health care-associated pneumonia, 767t, 812t
 indications for, 941t, 943
 for infections in cancer patient, 491
 for infectious diarrhea, 857t
 for intraabdominal infections, 767t
 for *Legionella* infections, 1020t
 for multidrug-resistant TB, 205e-6
 for nongonococcal bacterial arthritis, 835
 for osteomyelitis, 840t, 841t
 for *P. aeruginosa* infections, 808t, 1045t
 for pelvic inflammatory disease, 878
 for peritonitis, 848
 for plague prophylaxis, 1075t
 for plague treatment, 261e-5, 261e-6, 1075
 for pneumonia, 951–952
 resistance to, 807, 931t, 941t
 for sepsis/septic shock, 1757t
 in traveler's medical kit, 797
 Levomilnacipran, 2710t, 2717
 Levonorgestrel, 2391
 Levorphanol, 91t, 94
 Levosimendan, 1510, 1510t
 Levothyroxine
 adverse effects of, 2292
 in elderly patients, 2293
 for hypothyroidism, 2292–2293
 for myxedema coma, 2293
 for nontoxic multinodular goiter, 2302
 in pregnancy, 2293
 for subacute thyroiditis, 2299
 for thyroid cancer, 2295
 Lewis blood group system, 138e-1, 138e-1t
 Lewisite, 262e-1t
 Lewis-Sumner syndrome, 2699
 Lewy body(ies)
 characteristics of, 2606
 dementia with. *See* Dementia with Lewy bodies
 prions and, 444e-8f, 453e-6
 Leydig cell, 2358, 2358f
 LFA-1 (leukocyte function-associated antigen), 372e-24t
 LGI1 gene mutations, 2545t
 LGMD (limb-girdle muscular dystrophy), 462e-6t, 462e-7t, 462e-8
 LGV. *See* Lymphogranuloma venereum (LGV)
 LH. *See* Luteinizing hormone (LH)
 LH β gene mutations, 2364, 2364t
 Lhermitte's sign/symptom
 in cervical spine lesions, 121, 161, 2535
 in MS, 2535, 2663
 radiation therapy-related, 622, 709
 in spinal cord compression, 1790
 LHR gene mutations, 2354t
 LHX3 gene mutations, 2364, 2364t
 Lialda, 1963
 Libido
 decreased, in malabsorption syndromes, 1946t
 female, 330
 male, 324
 Libman-Sacks endocarditis, 2136
 Lice
 body (*Pediculus humanus*), 1146, 2745–2746
 diseases carried by
 epidemic typhus. *See* Epidemic (louse-borne) typhus
 relapsing fever. *See* Relapsing fever
 head (*Pediculus capitis*), 2745–2746, 2746f
 Lichenification, 344f
 Lichen myxedematosus, generalized (papular mucinosis), 364t, 365
 Lichenoid lesions, 340t
 Lichen planopilaris, 349
 Lichen planus, 349
 clinical features of, 347t
 alopecia, 355, 355t
 distribution, 341t, 342f, 349
 esophageal, 257
 morphology, 341t
 oral manifestations, 46e-1f, 237, 239t, 240t
 papules and plaques, 76e-3f, 349
 papulosquamous lesions, 353, 353t
 and plaques, 349
 drug-related, 349
 erosive, 46e-1f
 HCV infection and, 349, 353, 2041
 histologic features of, 347t
 treatment of, 349
 Lichen simplex chronicus, 342f, 345
 Lichen tropicus, 479e-1
 Licorice, 64e-8, 306, 307
 Liddle's syndrome (pseudoaldosteronism)
 clinical features of, 306, 322, 332e-9, 1620t
 diagnosis of, 307, 2319
 genetic considerations in, 306, 332e-7t, 1620t, 2318t, 2319
 pathogenesis of, 322
 pathophysiology of, 332e-9, 1621
 treatment of, 307, 2319
 Lidocaine
 adverse effects of, 1480t
 for arrhythmias, 1479t, 1498
 for cardiac arrest, 1769f, 1770
 dosage of, 1479t
 for interstitial cystitis/bladder pain syndrome, 60e-4
 for jellyfish stings, 2739
 for marine envenomations, 2739, 2741
 metabolism of, 34t
 for neuropathy, 94, 2682t
 for SUNCT/SUNA, 2596
 therapeutic monitoring of, 2765t
 for topical anesthesia in lumbar puncture, 443e-2
 Lidoderm patch, 2682t
 Life expectancy
 at age 60, 93e-1t
 at birth, 93e-1t, 93e-2
 genetic considerations in, 85e-8
 global burden of disease and, 9
 global differences in, 13e-1–2, 13e-2f, 13e-3f, 93e-1t, 93e-2
 screening procedures and increase in, 28t
 trends in, 93e-3–4, 93e-3f
 of women, 6e-1
 Life history theory of aging, 94e-2
 Lifestyle modifications
 for atherosclerosis prevention, 291e-8t, 291e-9
 for diabetes mellitus, 82
 for GERD, 1908
 for hypertension, 1622–1623, 1622t
 for metabolic syndrome, 2452–2453
 for NAFLD, 2056–2057
 for obesity, 2394
 Li-Fraumeni syndrome
 clinical features of, 523, 599t
 genetic considerations in, 101e-6t, 102e-2, 119e-1, 431t, 523, 599t
 lung cancer risk in, 507
 Light chain deposition disease/nephropathy
 clinical features of, 1835t, 1845, 1859
 diagnosis of, 1859
 etiology of, 1859
 pathophysiology of, 1859
 renal biopsy in, 62e-7f, 1859f
 treatment of, 1845, 1859
 Light-headedness, 148
 Likelihood ratio, 22, 23f
 Lille score, 2053
LILRB2 gene, 1241t
 Limaprost, 478e-5
 Limb apraxia, 179
 Limb dystonia, 2620
 Limb-girdle muscular dystrophy (LGMD), 436t, 462e-6t, 462e-7t, 462e-8
 Limbic encephalitis, 441e-1f, 444e-2t, 616, 616f, 2607
 Limbic network/system, 182
 Limb ischemia
 acute, 1646
 in peripheral artery disease. *See* Peripheral artery disease
 Limb-kinetic apraxia, 179
 Limb-sparing surgery, 119e-2, 119e-3
 Limey bile, 2082
 Limit dextrinosis (type III GSD), 432e-2t, 433e-2t, 433e-3, 436e-6f, 462e-14, 1565
 Limulus amebocyte lysate assay, 888
 Linaclotide, 273, 274, 1970, 1970t
 Linagliptin, 2414t
 Lincosamides. *See* Clindamycin
 Lindane, 2744, 2746
 Linear IgA disease, 370t, 373, 379
 Linear nevoid hypopigmentation (hypermelanosis of Ito), 357t, 359
 Linear scleroderma, 376
 Linezolid
 action of, 205e-7, 931t, 932, 961
 adverse effects of, 205e-7, 727t, 938t, 944, 961, 976
 for burn-wound infection, 166e-3
 for community-acquired pneumonia, 767t, 808t
 drug interactions of, 205e-7, 939t, 944
 for enterococcal infections, 975t, 976
 for health care-associated pneumonia, 812t
 indications for, 941t, 944
 for MRSA infections, 767t, 808t, 962t
 for *Nocardia* infections, 1087
 for osteomyelitis, 841t
 in pregnancy and lactation, 937t
 resistance to, 931t, 934, 941t, 961
 for staphylococcal infections, 832t, 961, 962t
 for TB, 205e-7
 for toxic shock syndrome, 963
 LINGO-1, 444e-5
LINGO1 gene mutations, 2619
 Linitis plastica, 534
 Linkage analysis, 441–442, 442t
 Linkage disequilibrium, 373e-3, 442, 442t
 Lionfish sting, 2740
 Liothyronine, 2292, 2306
 Lipase, 2087t, 2088, 2100–2101
 Lipedema, 1654
 Lipemia retinalis, 2439
 Lipid(s)
 defects in metabolism of, 462e-15
 deposits of, in familial hypercholesterolemia, 436e-4f
 digestion and absorption of, 1935–1936, 1935t, 1936f
 as energy source, 462e-14–15
 neutral, disorders of, 432e-3t
 production of, 2405
 transport by lipoproteins, 2437, 2437f
 Lipid A, 1752
 Lipid mediator generation, 2115, 2115f
 Lipid membrane microdomain (lipid raft), 372e-19
 Lipid-modifying therapy. *See* Lipoprotein disorders, treatment of
 Lipid polyene, 1352t
 Lipid storage diseases, 367e-2–3, 367e-3t

- Lipoarabinomannan, 1106
- Lipoatrophy, 2430
- Lipodermatosclerosis, 366, 1651
- Lipodystrophy
- in diabetes mellitus, 2422
 - familial, 435, 436t, 2450
 - forms of, 436e-8f, 2439
 - in HIV infection, 1259, 1260f, 1273, 2422, 2450
 - lipoprotein disorders in, 2439
 - metabolic syndrome and, 2450
 - protease inhibitors and, 2422
 - treatment of, 2439
- Lipofuscin, 94e-3
- Lipohypertrophy, 2430
- Lipoic acid. *See* α -lipoic acid
- Lipolysis, 1935
- Lipolytic enzymes, 2090
- Lipomas
- cardiac, 289e-2
 - clinical features of, 364, 364t
 - in MEN 1, 2336t, 2340
 - small intestinal, 536
- Lipooligosaccharide (LOS), *N. gonorrhoeae*, 1004
- Lipopeptides. *See* Daptomycin
- Lipopolysaccharide (LPS)
- A. baumannii*, 1037
 - B. cepacia*, 1048
 - Brucella*, 194e-2, 194e-3
 - in inflammatory response, 145e-5, 145e-6f, 1752
 - P. aeruginosa*, 1043, 1043t
 - in septic shock, 1025
- Lipoprotein(a), 1241t, 2436t
- Lipoprotein(s)
- classification of, 1241t, 2435–2437, 2436f
 - composition of, 1241t, 2435–2437. *See also*
 - High-density lipoprotein (HDL);
 - Low-density lipoprotein (LDL);
 - Very-low-density lipoproteins (VLDLs)
 - metabolism of, 292e-1v
 - oxidized, 291e-2
 - proteins associated with, 2436–2437
 - in transport of dietary lipids, 2437, 2437f
- Lipoprotein disorders
- approach to the patient, **2446**
 - arthritis in, 2243
 - atherosclerosis and, 291e-4–6, 291e-6t
 - cardiovascular disease risk and, 266e-4, 1588
 - diagnosis of, 2446
 - erectile dysfunction in, 326
 - global considerations in, 266e-4
 - metabolic causes of
 - impaired lipolysis of triglyceride-rich lipoproteins, 2439–2440
 - primary (genetic), 2439–2440, 2440t
 - secondary, 2439
 - impaired uptake of LDL, 2440–2441
 - primary (genetic), 2441–2443
 - secondary, 2441
 - low ApoB, 2444–2445. *See also*
 - Abetalipoproteinemia (acanthocytosis)
 - low HDL-C, 2445–2446, 2451
 - very high HDL-C, 2446
 - VLDL overproduction, 2438–2439, 2451
- musculoskeletal disorders associated with, 2242–2243
- screening for
- calculations for, 2446
 - classification of results, 480e-3t, 2446
 - in elderly, 84
 - recommendations, 29t, 2254t
 - reference ranges, 480e-3
- secondary causes of, 2444t
- diabetes mellitus, 291e-6, 2428
 - drug-related, 2443–2444
 - hypothyroidism, 2441
 - insulin resistance, 2438, 2450
 - kidney disease, 2438–2439, 2441
 - liver disease, 2443
 - metabolic syndrome, 2451
 - obesity, 2438
 - pancreatitis, 2091, 2093, 2096
 - skin manifestations of, 365
- treatment of, **2446**
- ACC/AHA guidelines for, 291e-4–6, 291e-6t, 291e-7f
 - bile acid sequestrants. *See* Bile acid sequestrants
 - for cardiovascular disease risk reduction, 2447
 - cholesterol absorption inhibitors, 462e-19, 2449–2450, 2449t
 - fibric acid derivatives. *See* Fibric acid derivatives
 - high LDL management, 291e-6–7, 291e-8f, 291e-9f, 2447–2449, 2448t
 - LDL apheresis, 2449
 - lifestyle modifications, 291e-6, 291e-8t, 291e-9, 2447
 - in metabolic syndrome, 2453–2454
 - nicotinic acid. *See* Nicotinic acid
 - omega-3 fatty acids. *See* Omega-3 fatty acids
 - severe hypertriglyceridemia management, 2446–2447
 - statins. *See* Statins
- Lipoprotein glomerulopathy, 1847
- Lipoprotein lipase (LPL), 2450
- Lipoprotein lipase (LPL) deficiency, 91e-3, 2439–2440, 2440t
- Liposarcoma, 119e-1. *See also* Soft tissue sarcoma
- Lipoteichoic acid, 145e-5
- 5-Lipoxygenase, 40
- Lip reading, 224
- Liquid nitrogen therapy, 351
- LIR (leukocyte Ig-like receptor), 373e-4
- Liraglutide, 2397, 2414–2415, 2414t
- Lisch nodules, 2343
- Lisfranc fracture–dislocation, 2244
- Lisinopril, 1512t, 1624t
- Lispro, 2411, 2411t
- Listeria monocytogenes*, 145e-7, 150e-2t, 982, 982f
- Listeria monocytogenes* infections
- bacteremia, 983
 - in cancer patient, 484t, 486t
 - CNS, 983
 - complications of, 984
 - diagnosis of, 983
 - epidemiology of, 982–983
 - food-related, 266, 983
 - meningitis, 886, 889–890, 889t, 983
 - neonatal, 983
 - pathogenesis of, 982
 - in pregnancy, 983
 - prevention of, 984
 - prognosis of, 984
 - in transplant recipient, 924t, 925, 926
 - treatment of, 983–984
- Listeriolysin (LLO), 982
- Lithium
- action of, 465e-2
 - adverse effects of
 - acid-base disorders, 317
 - alopecia, 354
 - AV conduction block, 1471t
 - common, 2718, 2718t
 - cutaneous, 352, 353, 356, 356t, 378
 - hypercalcemia, 2475
 - lupus syndrome, 2218t
 - nail disorders, 380
 - nephrogenic diabetes insipidus, 64e-3–4, 303
 - nephropathy, 1856t, 1861–1862
 - neuropathy, 2687t
 - nystagmus, 211
 - obesity, 2392
 - psoriasis, 347
 - rare, 2718t
 - SA node dysfunction, 1467t
 - secondary parkinsonism, 2611
 - seizures, 2547t
 - for bipolar disorder, 2718, 2718t
 - for cluster headache prevention, 2595, 2595t
 - drug interactions of, 246e-5t
 - for hypnic headache, 2598
 - overdosage/poisoning with, 473e-12t, 2765t
 - pharmacology of, 2718t
 - therapeutic monitoring of, 2765t
- Livedoid vasculopathy, 369, 369t
- Livedo reticularis, 68, 369, 1648f, **1649**, 2135
- Liver
- age-related changes in, 94e-4
 - alcohol effects on. *See* Alcoholic liver disease
 - anatomy of, 366e-1, 366e-1f, 1989–1990
 - drug metabolism in, 33–34, 33f
 - functions of, 1989–1990
 - metastatic tumors to, 553
 - ascites with, 287
 - colorectal cancer, 542
 - NETs, 560, 572–573, 572f
 - treatment of, 574
 - palpation of, 286
 - recellularization of, 92e-2, 92e-3v
- Liver abscess, **850**
- in actinomycosis, 1089, 1090f
 - amebic
 - clinical features of, 1365
 - complications of, 1365
 - diagnosis of, 245e-1, 850, 1366, 1366f
 - pathogenesis of, 1365
 - treatment of, 1367, 1367t
 - Candida*, 850
 - clinical features of, 850
 - diagnosis of, 850
 - etiology of, 850
 - global considerations in, 850
 - K. pneumoniae*, 850, 1032, 1032f
 - treatment of, 850–851
- Liver biopsy, **366e-1**, 1994
- in acetaminophen hepatotoxicity and alcoholic liver injury, 366e-4f
 - in acute hepatitis, 366e-1f, 366e-6f
 - in α_1 -antitrypsin deficiency, 366e-4f
 - in autoimmune hepatitis, 366e-2–3f
 - in *Brucella* infections, 194e-3, 194e-3f
 - in chronic HBV infection, 366e-2f
 - in chronic HCV infection, 366e-1–2f, 366e-3f
 - in cirrhosis, 366e-3f
 - in cirrhosis secondary to hemochromatosis with hepatocellular carcinoma, 366e-4f
 - in CMV infection with liver allograft, 366e-3f
 - contraindications to, 1998
 - in hemochromatosis, 366e-4f, 366e-5f, 2517
 - in hepatocellular carcinoma, 366e-4f
 - indications for, 1996f, 1998
 - in NAFLD, 2056
 - in nonalcoholic steatohepatitis, 366e-5f
 - in primary biliary cirrhosis, 366e-3f
 - in primary sclerosing cholangitis, 366e-5f
 - in rejection of liver allograft, 366e-3f
 - in Wilson's disease, 366e-6f, 2519
- Liver cancer. *See* Hepatocellular carcinoma (HCC)
- Liver disease/failure, **1989**. *See also* Cirrhosis in alcohol abuse. *See* Alcoholic liver disease in α_1 -antitrypsin deficiency, 367e-2, 1992
- in amyloidosis, 367e-3
 - in cancer patient, 488, 622
 - cholestatic, 283–284, 1990, 1990t, 2060. *See also* Biliary cirrhosis
 - clinical features of, 286, 1990–1991
 - anemia, 631
 - ascites, 286, 286f, 1992
 - bleeding, 404
 - coagulation disorders, 739, 739f, 739t
 - cutaneous, 1992–1993
 - fatigue, 1991
 - halitosis, 242
 - hepatomegaly, 1992
 - icterus, 1992
 - itching, 1991
 - jaundice, 283, 1991
 - lactic acidosis, 318
 - lipoprotein disorders, 2443
 - nausea and vomiting, 259, 1991
 - neuropathy, 2684

- oral, 236
 pain, 1991
 respiratory, 324
 splenomegaly, 1992
 thromboembolism, 739–740, 739f
 clinical history, 1991–1992, 1992t
 in cystic fibrosis, 367e–2, 1697
 deaths from, by sex, 6e–1t
 diagnosis of
 algorithm for, 1994f
 biopsy in, 1994. *See also* Liver biopsy
 vs. DIC, 738
 imaging in, 1993–1994, 1999
 in jaundiced patient, 282–283
 laboratory tests in, 1993, 1993t, 1999
 liver function tests in. *See* Liver function tests
 drug-induced, **2023**, 2056t
 acetaminophen in, 43–44, 283, 366e–4f, 2025t, 2027, 2028f
 amiodarone in, 2029
 amoxicillin-clavulanate in, 2029
 anabolic steroids in, 2029
 antiretrovirals in, 2030
 clinical features of, 2025t
 erythromycin in, 2029
 in exacerbation of preexisting liver disease, 1995
 herbal medicines in, 2030
 isoniazid in, 2025t, 2029
 mechanisms of, 2023–2025, 2024f
 nitrofurantoin in, 2029
 oral contraceptives in, 2025t, 2029
 parenteral nutrition in, 2030
 phenytoin in, 2029
 statins in, 2030
 treatment of, 2025
 trimethoprim-sulfamethoxazole, 2029–2030
 in tuberculosis, 1117
 valproate in, 2028
 effect on drug concentration and response, 37
 etiology of, 1990, 1990t
 family history in, 1992
 folate deficiency in, 647
 genetic considerations in, 367e–1–2, 367e–1t, 1992, 2515, 2518
 grading of, 1994–1995
 granuloma, 367e–3, 2026t
 in hemochromatosis, 367e–1–2, 367e–1t, 367e–2t, 2516, 2516f
 in hepatitis. *See* Hepatitis
 hepatocellular, 1990, 1990t
 in HSCT recipients, 139e–3
 in IBD, 1958
 immunization recommendations for, 788f
 infiltrative, 367e–1t, 367e–3
 in leptospirosis, 1144
 in lipid storage disorders, 367e–2–3
 lymphoma, 367e–3
 metabolic, 367e–1t, 367e–2–3
 in NAFLD. *See* Nonalcoholic fatty liver disease (NAFLD)
 obesity and, 415e–6
 patterns of, 1990, 1990t
 peripheral blood smear in, 81e–5f
 physical examination in, 282–283, 1992–1993
 in porphyrias, 367e–3
 in pregnancy, 49
 risk factors for, 1991–1992
 in sarcoidosis, 2207–2208
 in schistosomiasis, 1426
 specialized nutritional support in, 98e–4
 staging of, 1994–1995
 thyroid hormone levels in, 2300
 toxic, 2023, 2025t, 2026t
 treatment of
 liver transplantation. *See* Liver transplantation
 stem cell therapy, 90e–3
 tumors. *See* Liver tumors
 in type 2 diabetes mellitus, 2405
 in Wilson's disease, 1992, 2519
 in women, 6e–1t
 Liver flukes. *See* Biliary (hepatic) flukes
 Liver function tests, **1995**
 abnormal, evaluation of, 1993, 1994f, 1995, 1996f
 blood ammonia, 1996
 coagulation factors, 1998
 serum albumin. *See* Albumin, serum
 serum bilirubin. *See* Bilirubin, serum
 serum enzymes, 1997. *See also* Alanine aminotransferase (ALT); Alkaline phosphatase (ALP); Aspartate aminotransferase (AST); γ -glutamyl transpeptidase (GGTP)
 serum globulins, 1998
 shortcomings of, 1995
 urine bilirubin. *See* Bilirubin, urine
 use of, 1998t, 1999
 Liver pain, 1991
 Liver phosphorylase deficiency (type VI GSD; Hers disease), 433e–2t, 433e–4
 Liver phosphorylase kinase deficiency (type IX GSD), 433e–2t, 433e–4
 LiverTox website, 2027
 Liver transplantation, **2067**
 for acute intermittent porphyria, 2529–2530
 for alcoholic liver disease, 2053–2054, 2075
 alternatives to, 2070–2071
 for ATTR amyloidosis, 724
 cadaver donor selection for, 2069–2070
 candidate selection for, 2069–2070, 2069t
 in children, 2068, 2068t
 for cholangiocarcinoma, 553
 complications of
 anemia, 2072
 graft rejection, 366e–3f, 2073
 hepatic, 2072t, 2073
 infections, 924t, 927–928, 1341f, 2072–2073
 nonhepatic, 2072–2073, 2072t
 recurrence of primary disease, 2068, 2073–2075
 renal, 2072
 contraindications to, 2068–2069, 2069t
 for erythropoietic protoporphyria, 2533
 frequency of, 2067
 for hemochromatosis, 2518
 for hepatocellular carcinoma, 549–550
 history of, 2067
 immunosuppressive therapy for, 2071–2072, 2764t, 2766t
 indications for, 2067–2068, 2068t
 living donor, 2070
 MELD score and, 1995
 for metastatic NETs, 575
 for NALFD, 2057
 orthotopic, 2067
 outcome of, 2067, 2073
 PELD system in, 1995
 quality of life after, 2075
 surgical technique for, 2070, 2070f
 survival after, 2073
 timing of, 2068
 Liver tumors, **544**
 benign, 553–554
 cholangiocarcinoma. *See* Cholangiocarcinoma
 epithelioid hemangioendothelioma, 552
 hepatocellular carcinoma. *See* Hepatocellular carcinoma (HCC)
 metastatic from other sites, 553
 Liver X receptor, 2076
 Living will, 58–59
 LL3858, 205e–7
 LLO (listeriolysin), 982
 LMNA gene mutations, 94e–4, 435, 436t
 LMO-2 gene, 91e–2
 LMWH. *See* Low-molecular-weight heparin (LMWH)
 LMX1B gene mutations, 1847
 LMYC gene mutations, 101e–2t, 101e–3
 Loading dose, 35, 35f
 Loa loa (African eye worm)/loiasis
 characteristics of parasite, 1418t, 1421
 clinical features of, 1421, 1422f
 diagnosis of, 245e–3t, 245e–6t, 765t, 1422
 epidemiology of, 765t, 1418t, 1421
 pathogenesis of, 1421
 prophylaxis for, 1422
 treatment of, 246e–8, 1422
 Lobectomy, for lung cancer, 514, 516
 Lobectomy, temporal, 2557
 Lobeline, overdose/poisoning with, 473e–10t
 Lobular neoplasia, 531–532
 Local anesthetics, 242, 473e–13t
 Local heat urticaria, 2118
 Locked-in state, 41e–1v, 442e–2, 1771, 2578
 Lockjaw (trismus), 985
 Locus coeruleus, 465e–3, 465e–3f
 Locus control region, 428
 Locus (nonallelic) heterogeneity, 435–436, 436t, 447
 Lod score, 442
 Loeffler's syndrome, 422
 Loeyes-Dietz syndrome, 1443, 1638, 1641, 2513
 Lofexidine, 468e–3
 Löffler's endocarditis, 1567
 Löffler's syndrome, 1686–1687, 1686t
 Löfgren's syndrome, 2206, 2207
 LOH. *See* Loss of heterozygosity (LOH)
 Lomitapide, 2448t, 2449
 Lomustine (CCNU), 103e–9t, 601
 Lone Star tick, 1160
 Long-chain fatty acids/triglycerides, 1935, 1935t, 1936
 Long-Distance Corridor Walk, 78t
 Longevity. *See* Life expectancy
 Longevity dividend, 94e–1
 Long QT syndrome
 arrhythmia in, 273e–4
 clinical features of, 147, 1497
 congenital, 41, 273e–4, 278e–12f, 1497
 ECG in, 278e–12f, 1497f
 genetic considerations in, 147, 1496t, 1497
 genetic testing for, 450t
 treatment of, 273e–4, 450, 450t
 Long-term care, 83–84, 83t
 Long terminal repeats (LTRs), 225e–1, 1218–1219, 1219f
 Loop diuretics
 action of, 299, 332e–4f, 332e–8
 adverse effects of, 306
 for chronic kidney disease, 1814, 1815
 for hypercalcemia, 2480t
 for hypercalcemia of malignancy, 610
 for hyperkalemia, 312
 for hypokalemia, 308
 for hyponatremia, 302
 for pulmonary edema, 1763
 Loop of Henle
 disorders involving, 332e–7t
 embryologic development of, 332e–1–2
 functions of, 332e–4f, 332e–8
 in water absorption/excretion, 296, 296f
 Looser's zones, 2466
 Loperamide
 for diarrhea, 103e–25, 267–268, 272
 for irritable bowel syndrome, 1969, 1970t
 for microscopic colitis, 270
 for travelers' diarrhea, 797, 857t
 Lopinavir, 1276f, 1278, 1280f
 LOPS (loss of protective sensation), 2407
 Lorazepam
 for acute alcohol intoxication, 2727
 for anxiety disorders, 2711t
 for delirium, 64t, 65
 for dyspnea, 62t
 for insomnia, 65
 for lumbar puncture analgesia, 443e–1
 for nausea and vomiting, 62, 103e–25, 260t, 261
 overdose/poisoning with, 473e–10t
 pharmacology of, 2711t
 for sedation after myocardial infarction, 1606
 for status epilepticus, 1735, 2558f
 during terminal weaning, 67
 Lorcaserin, 2396–2397, 2397t, 2453
 Lordotic posture, 462e–1, 462e–4f
 Losartan
 adverse effects of, 431e–5
 genetic variations in response to, 39t, 40
 for heart failure, 1512t

- Losartan (*Cont.*):
 for hypertension, 1624t
 metabolism of, 34t, 35, 42
 for Raynaud's phenomenon, 2164
- Loss of heterozygosity (LOH), 101e-4, 101e-5f, 2305t
- Loss of protective sensation (LOPS), 2407
- Louse-borne diseases
 epidemic typhus. *See* Epidemic (louse-borne) typhus
 relapsing fever. *See* Relapsing fever
- Lovastatin
 adverse effects of, 2199, 2218t, 2447–2448, 2448t
 drug interactions of, 42, 246e-3t, 2447
 for lipoprotein disorders, 2448t
- Low- and middle-income countries
 demographics of
 life expectancy, 93e-1t
 population aged 60+, 93e-2, 93e-2f
 population aged 80+, 93e-2–3, 93e-3t
 population aging indicators, 93e-1t
 health challenges in
 brain drain, 13–14
 burden of disease, 9, 11t, 13e-1–2, 13e-2f, 93e-5, 93e-5f, 93e-5t
 cancer, 16, 16f
 cardiovascular disease, 15–16, 266e-2–3, 266e-2f, 266e-3f
 child mortality, 13e-3, 13e-3f
 diabetes, 15
 health system performance, 13e-3–4, 13e-4f
 HIV/AIDS, 12, 12f, 13f, 1223–1225, 1224f
 life expectancy, 13e-1–2, 13e-2f, 13e-3f
 malaria, 14–15
 skews in research, 13e-4
 social stratification, 13e-4
 tuberculosis, 12–13
 primary care in, **13e-1**
 experiences with, 13e-6–7, 13e-6f
 opportunities to build, 13e-7–8
 revitalization of, 13e-4–6
 smoking patterns in, 266e-4
 travel to
 health problems during, 793–794, 794f
 health problems following, 800–801f, 802, 802t
- Low-density lipoprotein (LDL)
 atherosclerosis and, 291e-5–6
 composition of, 2436–2437, 2436f, 2436t
 elevated levels of. *See also* Lipoprotein disorders
 in metabolic syndrome, 2451, 2453
 pathophysiology of, 2440–2441
 primary (genetic) causes of, 2440t, 2441–2443
 secondary causes of, 2441
 treatment of, 291e-6–7, 291e-8f, 291e-9f, 2453
 estimate of, 2446
 inherited causes of low levels of, 2444–2445. *See also* Abetalipoproteinemia (acanthocytosis)
 in transport of hepatic lipids, 2437, 2437f
- Low-density lipoprotein (LDL) apheresis, 2441, 2449
- Lower esophageal sphincter (LES), 255, 1904, 1905f, 1906
- Lower motor neuron diseases, 2634t, 2636. *See also* Amyotrophic lateral sclerosis (ALS)
- Lower motor neuron pathways, 155f
- Lower motor neuron (neuropathic) weakness, 154, 154t, 157
- Lower urinary tract symptoms (LUTS), 7e-4, 587, 587t
- Low-fat diet, 476
- Low FODMAP diet, 263
- Low-molecular-weight heparin (LMWH), 752
 action of, 750f, 752
 adverse effects of, 753
 assessment of, 407
 discontinuation before lumbar puncture, 443e-1
 dosage of, 752–753
 for DVT/PE treatment, 47, 1635, 1635t
 vs. fondaparinux, 753t
 vs. heparin, 752t
 monitoring treatment with, 752
 for myocardial infarction, 1597, 1598t, 1606
 pharmacology of, 752
 thrombocytopenia induced by. *See* Heparin-induced thrombocytopenia
 for thrombophlebitis, 613
 for VTE prophylaxis, 55, 1637t
- Loxapine, 2722t
- LPL (lipoprotein lipase), 2450
- LPL (lipoprotein lipase) deficiency, 91e-3, 2439–2440, 2440t
- LPS. *See* Lipopolysaccharide (LPS)
- LPS-binding protein, 372e-2
- LPS receptor. *See* CD14 (LPS-receptor)
- LP-X, 2443
- Lrp4 antibodies, 2702
- LRP5 gene mutations, 2489
- LRKK2 gene mutations, 444e-9, 444e-10, 2609, 2613
- LSD (lysergic acid diethylamide), 469e-4
- LTA4H gene, 775
- LTBI (latent tuberculosis infection). *See* Tuberculosis (TB), latent infection
- LTRs (long terminal repeats), 225e-1, 1218–1219, 1219f
- L-type fatty acid binding protein, 1808
- LT- β , 372e-11t
- Lubiprostone
 for constipation, 273, 274
 for irritable bowel syndrome, 1970, 1970t
- Lubricin, 2140, 2226
- Lucio's phenomenon, 829, 1124, 1128
- Ludwig's angina, 233, 236, 1097
- Lues maligna, 1135, 1259
- Lujo virus, 1306t
- Lumbar adhesive arachnoiditis, 117
- Lumbar fracture, 2489
- Lumbar fusion, 120
- Lumbar lordosis, 285
- Lumbar plexus, 2693, 2693f
- Lumbar puncture, **443e-1**
 analgesia for, 443e-1–2
 in coma, 1775
 headache following, 111, 443e-3, 443e-3f, 443e-3t
 imaging and laboratory studies prior to, 443e-1
 in meningitis, 887–888
 needles for, 443e-3, 443e-3f
 in neurologic critical illness, 1779
 normal values, 443e-3t, 443e-4
 patient positioning for, 443e-2, 443e-2f
 in patient receiving anticoagulants or antiplatelet agents, 443e-1
 specimen collection and transport, 150e-4t
 in subarachnoid hemorrhage, 1786
 technique, 443e-2–3, 486e-1v
- Lumbar spinal lesions, 2651
- Lumbar spinal stenosis, 115–116, 116f
- Lumbosacral plexopathies, 462e-19, 2693–2694
- Lumbosacral plexus, 2693, 2694f
- Lumbo virus, 1309t
- Lumefantrine, 246e-2t, 246e-9, 798
- Luminal A breast cancer, 526
- Luminal B breast cancer, 526
- Lumpectomy, 526–528, 531
- Lung
 airways resistance in, 306e-6
 blood flow through, 306e-1
 compliance of, 306e-1–2, 1733
 diffusing capacity of, 306e-6
 elastic recoil pressure of, 306e-1
 flow-related properties of, 306e-2, 306e-2f, 306e-3f, 306e-6
 shunt, 306e-4, 306e-4f
 volume-related properties of, 306e-1–2, 306e-1f, 306e-2f, 306e-6
- Lung abscess, **813**
 anaerobic bacterial, 1098, 1098f
 approach to the patient, 815
 clinical features of, 814
 complications of, 815, 901
 definition of, 813
 diagnosis of, 815
 differential diagnosis of, 815
 epidemiology of, 813–814
 etiology of, 813, 814t, 1098
 hemoptysis in, 245
 pathogenesis of, 814, 814f
 pathology and microbiology of, 814
 prevention of, 815
 prognosis of, 815
 treatment of, 815, 1102
- Lung biopsy
 in granulomatosis with polyangiitis, 386e-1, 386e-5f
 in hypersensitivity pneumonitis, 1683, 1683f
 in interstitial lung diseases, 1710
 in microscopic polyangiitis, 386e-5f
 in vasculitic syndromes, 386e-1
- Lung cancer, **506**
 adenocarcinoma, 507, 507f
 asbestos-related, 1689
 carcinoma of unknown primary and, 120e-2, 120e-2t
 classification of, 507–508, 507f
 clinical features of, 510–511, 510t
 clubbing in, 250
 deaths from, 467t, 468f, 470t, 471t, 506
 diagnosis of, 308e-15–16f, 511–512
 epidemiology of, 506
 genetic considerations in, 101e-2t, 507–509, 507t, 509f
 hemoptysis in, 245–246
 immunohistochemistry of, 508
 incidence of, 467–468, 467t, 468f, 471t, 506
 large cell, 507, 507f
 lymphadenopathy in, 409
 metastatic
 to bone, 119e-4
 clinical features of, 511t
 to CNS, 604f, 604t
 to eye, 206
 to heart, 271e-9f, 271e-9v, 289e-2
 incidence of, 511
 treatment of, 519–521, 520t, 523f
 non-small cell
 metastatic, 519–521, 520t, 523f, 604f
 pathology of, 507–508, 507f
 prognosis of, 516t
 staging of, 512–513, 514f, 516t
 treatment of, 102e-3, 102e-5t, 512, 512f, 515–519, 523f
 occult, 515
 occupational exposures and, 1692–1693
 paraneoplastic syndromes in, 511, 609t, 612t
 pathogenesis of, 508–509
 pathology of, 507–508, 507f
 prevention of, 477
 prognosis of, 516t
 racial/ethnic disparities in treatment of, 16e-1
 radon exposure and, 1693
 risk factors for, 506–507
 screening for, 482, 509–510, 510t
 second malignancies in survivors of, 623
 small cell
 hypokalemia in, 64e-7–8
 immunohistochemistry of, 508
 intestinal pseudoobstruction in, 259–260
 Lambert-Eaton myasthenic syndrome in, 511, 616, 2703
 paraneoplastic syndromes in, 608–609, 609t, 616, 617
 pathology of, 507–508, 507f
 SIADH in, 64e-4–5, 300
 staging of, 513–514
 treatment of, 515f, 522, 522f
 smoking and, 475–476, 506–507, 2730t
 squamous cell, 507, 507f
 staging of, 512–515, 514f, 516t, 1664
 stem cells in, 508
 superior vena cava syndrome in, 510t, 511, 1788, 1789f

- treatment of
 in absent or nonbulky mediastinal lymph node disease, 518–519
 biologic therapy, 520, 521
 chemotherapy, 517–519, 518t, 522
 EGFR inhibitors, 521, 521t
 gene therapy, 91e-4
 maintenance chemotherapy, 520–521, 520t
 in mediastinal lymph node disease, 519
 in metastatic disease, 519–520, 520t
 neoadjuvant chemotherapy, 518
 in Pancoast tumors, 519
 prophylactic cranial irradiation, 522
 radiation therapy, 517, 522
 supportive care, 521
 surgical resection, 514, 516–517, 522
 targeted chemotherapy, 102e-3, 102e-5t, 102e-16, 102e-19, 103e-17t, 521
 tumor markers in, 473t
- Lung disease. *See also* Respiratory disorders; *specific disorders*
 air pollution and, 1693
 barotrauma-related, 477e-7
 biomass smoke-induced, 1693–1694, 1694f
 cancer. *See* Lung cancer
 cyanosis in, 249
 in cystic fibrosis, 1697. *See also* Cystic fibrosis (CF)
 deaths from, 56t
 global considerations in, 1693–1694
 high-altitude travel and, 476e-5
 IgG4-related, 391e-1t
 indoor exposures and, 1693
 infections in cancer patient, 488–489, 489t
 as late consequences of cancer/cancer therapy, 621–622
 in leptospirosis, 1143f
 obesity and, 415e-6
 occupational. *See* Occupational illness, lung disease
 pulmonary hypertension in, 1658
 respiratory acidosis in, 323
 in rheumatoid arthritis, 2138
 in sarcoidosis, 2206–2207, 2206f, 2206t
 in SLE, 1714, 2129, 2129t
 in systemic sclerosis, 2157–2158, 2158f, 2160–2161, 2161f
 in tularemia, 1068
- Lung flukes (*Paragonimus* spp.)
 clinical features of, 1429
 diagnosis of, 245e-2t, 245e-5t
 eosinophilia caused by, 765t
 epidemiology of, 245, 245e-2t, 765t, 1423t, 1429
 hemoptysis caused by, 245
 life-cycle hosts of, 245e-2t
 treatment of, 246e-13, 1428t, 1429
- Lung injury, acute. *See* Acute lung injury
- Lung lavage, 263e-7
- Lung stem cells, 453, 453t
- Lung transplantation
 complications of, 320e-3–4, 320e-3t
 for COPD, 1706
 cost of, 320e-3
 for cystic fibrosis, 320e-1t, 320e-2, 1699
 indications for, 320e-1
 infections in recipient, 924t, 926–927
 MRA of anastomosis following, 1667f
 organ allocation for, 320e-1–2
 outcomes of, 320e-2–3, 320e-2t
 post-transplantation management, 320e-2, 2764t
 procedure, 320e-2
 for pulmonary lymphangiomyomatosis, 1715
 recipient selection, 320e-1, 320e-1t
- Lung volume(s), 306e-1–2, 306e-2f, 306e-6
- Lung volume reduction surgery, 1706
- Lungworm, rat. *See* *Angiostrongylus* spp. infections
- Lupus anticoagulants
 in antiphospholipid syndrome, 2134
 laboratory evaluation of, 406, 2135t
 in SLE, 2131
- in thrombotic disease, 740
 warfarin therapy and, 756
- Lupus erythematosus
 acute cutaneous, 375, 375f
 discoid (chronic cutaneous). *See* Discoid (chronic cutaneous) lupus erythematosus
 drug-induced, 40, 44, 2218t
 subacute, 25e-3f, 2128
 subacute cutaneous, 356, 356t, 364t, 366, 367, 375, 2218t
 systemic. *See* Systemic lupus erythematosus (SLE)
- Lupus hairs, 354
- Lupus nephritis
 classification of, 1838, 1838t, 2126, 2127t
 clinical features of, 1838, 1858, 2128, 2129t
 crescentic, 2133
 membranous, 2133
 pathogenesis of, 1833, 1838
 renal biopsy in, 62e-5f, 1838
 treatment of, 1838–1839, 2128, 2130t, 2133
- Lupus pernio, 364t, 367, 2207, 2207f
- Lupus profundus, 2128
- Lupus vulgaris, 364t, 367
- Lurasidone, 2718, 2722t
- Lusitropic effect, 265e-7f
- Lutein, 96e-7
- Luteinizing hormone (LH), **401e-4**
 action of, 400e-2, 401e-4, 2377
 deficiency of, 2259t
 ectopic production of, 609t
 expression and regulation of, 401e-1t, 401e-2f
 feedback control of, 400e-6
 in females, age-related changes in levels of, 2376–2377, 2377f
 laboratory evaluation of, 2258t
 in male pubertal development, 2358
 in menstrual cycle regulation, 2378–2379, 2379f
 in ovarian follicle development, 2375f, 2376
 in perimenopause, 2381, 2381f
 pNET secreting, 559t, 571
 reference values for, 2761t
 secretion of, 401e-4, 2376
 synthesis of, 401e-4
- Luteinizing hormone (LH) receptor, 437, 2252
- Luteinizing hormone–releasing hormone (LHRH) agonists, 103e-15
- ¹⁷⁷Lutetium, 575
- LUTS (lower urinary tract symptoms), 7e-4, 587, 587t
- LVADs. *See* Left ventricular assist devices (LVADs)
- LVR (Laparoscopic Ventral Rectopexy), for rectal prolapse, 1974, 1975f
- Lycopene, 96e-7
- Lying to patient, 17e-1
- Lyme borreliosis (Lyme disease), **1149**
 chronic, 1151, 1153
 climate change and, 151e-5–6, 151e-6f
 clinical features of, 129t, 356, 1150–1151
 arthritis, 372e-24t, 835–836
 AV conduction block, 278e-4f, 1151, 1471t, 1472
 chronic meningitis, 907t
 disseminated infection, 1150–1151
 erythema migrans, 25e-2f, 129t, 134, 355–356, 1150, 1150f
 evolution of, 129t, 134, 356, 1150
 facial palsy, 2648
 localized infection, 1150
 myocarditis, 1561
 myositis, 2198
 neuropathy, 2633, 2684
 persistent infection, 1151
 SA node dysfunction, 1467t
 co-infection with *Babesia*, 1152, 1385, 1386
 diagnosis of, 770, 1151–1152, 1152t
 differential diagnosis of, 134, 1152
 epidemiology of, 1149
 etiology of, 1149
 pathogenesis of, 1150
 prevention of, 1153, 2745
- prognosis for, 1153
 prophylaxis for, 1153, 2745
 regional variations in, 1151
 reinfection in, 1153
 ticks in, 1149, 2745
 treatment of, 836, 1152–1153, 1152t, 1153f
- Lymphadenitis, tuberculous, 1109, 1109f
- Lymphadenopathy, **407**
 approach to the patient, 407–409
 benign, 407
 diseases associated with, 408t
 in fever of unknown origin, 141
 IgG4-related, 391e-1t, 391e-3
 infections associated with, 408, 408t, 763–764
Brucella, 194e-3
 cat-scratch disease, 1079, 1080f
 HIV infection, 1232–1233, 1262. *See also* HIV infection
 nontuberculous mycobacteria, 1130
 plague, 1072, 1072f
 tularemia, 1068, 1068f
 laboratory evaluation of, 409
 localized/regional, 408
 in Sjögren's syndrome, 2167, 2167t
 with splenomegaly, 408
 thoracic/abdominal, 408–409
- Lymphangiectasia, intestinal. *See* Intestinal lymphangiectasia
- Lymphangiography, 1654
- Lymphangiomyomatosis, pulmonary, 308e-5f, 1714–1715
- Lymphangitis, 968, 1654
- Lymphatic filariasis, **1417**
 clinical features of, 152e-6t, 837, 1417–1418, 1418f, 1419f, 1654
 diagnosis of, 152e-6t, 245e-3t, 245e-6, 245e-6t, 1418–1419
 differential diagnosis of, 1419
 epidemiology of, 1417, 1418t
 incubation period for, 152e-6t
 pathogenesis of, 1417
 prevention and control of, 1420
 in returning war veterans, 152e-6t
 treatment of, 152e-6t, 246e-8, 837, 1419–1420
- Lymphatic obstruction, 253, 269
- Lymphedema, **1653**
 anatomic considerations in, 1653
 clinical features of, 1654, 1654f
 congenital, 1653
 diagnosis of, 1654–1655
 differential diagnosis of, 1654
 etiology of, 1653–1654, 1653t
 in lymphatic filariasis, 1418, 1418f, 1419f, 1654
 pathophysiology of, 253
 stages of, 1654t
 treatment of, 1655
- Lymphedema praecox (Meigs's disease), 594, 1653–1654, 2620
- Lymphedema tarda, 1650
- Lymph node biopsy, 409
- Lymphocyte(s)
 B. *See* B cell(s)
 endothelial cell interactions of, 372e-26–28, 372e-28f, 372e-29t
 large granular, 81e-2, 372e-1, 372e-6
 in peripheral blood smear, 81e-2
 reactive, 81e-2
 T. *See* T cell(s)
- Lymphocytic choriomeningitis (LCM), 891, 893, 908t, 1306t, 1317–1318
- Lymphocytic colitis, 1957
- Lymphocytic gastritis, 1931
- Lymphocytic hypophysitis, 2257
- Lymphocytic interstitial pneumonitis, 1716
- Lymphocytic meningitis, 909t
- Lymphocytic variant hypereosinophilia, 135e-7, 135e-7t
- Lymphocytoma cutis, 364t, 367
- Lymphogranuloma venereum (LGV)
 clinical features of, 880f, 880t, 1169–1170, 1654
 complications of, 1170
 diagnosis of, 1170t
 epidemiology of, 879, 1169
 lymphadenopathy in, 408

- Lymphohistiocytosis, hemophagocytic, 375e-2, 2112
- Lymphoid hyperplasia
atypical, 710
cutaneous (pseudolymphoma), 353–354, 364t, 366
- Lymphoid malignancies, **135e-1, 695**. *See also* Leukemia
anemia in, 393
approach to the patient, 699–701
AV conduction block in, 1471t
B cell
 B-cell prolymphocytic lymphoma, 135e-2t, 706
 Burkitt's lymphoma. *See* Burkitt's lymphoma
 chronic lymphoid leukemia. *See* B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma
 diffuse large B-cell lymphoma. *See* Diffuse large B-cell lymphoma
 follicular lymphoma. *See* Follicular lymphoma
 hairy cell leukemia. *See* Hairy cell leukemia
 intravascular large B-cell lymphoma, 135e-2
 lymphomatoid granulomatosis, 135e-3
 MALT lymphoma. *See* Mucosa-associated lymphoid tissue (MALT) lymphoma
 mantle cell lymphoma. *See* Mantle cell lymphoma
 mediastinal (thymic) large B-cell lymphoma, 135e-2
 nodal marginal zone B-cell lymphoma, 135e-2, 707
 precursor B-cell lymphoblastic, 701
 primary effusion lymphoma, 135e-2, 214e-7, 1194, 1270
 splenic marginal zone lymphoma, 135e-1, 135e-2t, 706
 carcinoma of unknown primary and, 120e-2t
 classification of, 135e-1t, 695, 695t, 696t
 CNS, primary. *See* Primary central nervous system lymphoma (PCNSL)
 deaths from, 467t, 470t
 definition of, 102e-1
 diseases or exposures associated with, 697t
 EBV infection and, 135e-3, 135e-4, 697, 697t, 1188–1189
 epidemiology of, 695–697, 695f
 erythrophagocytosis in, 81e-11f
 etiology of, 695–697, 697t, 1188–1189
 fever in, 126
 gene profiling in, 699
 genetic considerations in, 101e-2t, 101e-3t, 698–699, 699t
 in HIV infection, 1270–1271, 1271f
 Hodgkin's lymphoma. *See* Hodgkin's lymphoma
 immunology of, 697–699, 698f
 incidence of, 467t
 infections in, 488
 liver disease in, 367e-3
 lymphadenopathy in, 408
 neuropathy in, 2685
 NK cell
 blastic, 135e-4
 extranodal NK/T cell, nasal type, 135e-3, 697, 708
 nodular follicular, 81e-7f
 ocular invasion by, 206
 oral manifestations of, 239t, 1271f
 paraneoplastic syndromes in, 608, 609t, 612t, 618–619
 pleural effusion in, 474
 proteinuria in, 293
 in rheumatoid arthritis, 2138
 in Sjögren's syndrome, 2167, 2167t
 skin manifestations of, 76e-5f, 353, 353t, 2193
 small intestinal, 537
 splenomegaly in, 411
 T cell
 adult. *See* Adult T-cell leukemia/lymphoma
 anaplastic large T/null cell lymphoma, 102e-5t, 699t, 702t, 707–708
 angiimmunoblastic T-cell lymphoma, 135e-4 to 135e-5, 708
 enteropathy-type lymphoma, 135e-3, 708
 extranodal T/NK cell lymphoma, nasal type, 135e-3, 697, 708, 2184
 hepatosplenic $\gamma\delta$ T-cell lymphoma, 135e-3, 708
 mycosis fungoides. *See* Mycosis fungoides; Mycosis fungoides (cutaneous T-cell lymphoma)
 precursor T-cell lymphoblastic lymphoma/leukemia, 702t, 707
 primary cutaneous CD30+ T-cell lymphoma, 135e-4
 subcutaneous panniculitis-like T-cell lymphoma, 135e-4, 708
 in thymus, 123e-1. *See also* Thymoma
 in thyroid, 2307
 treatment of, 102e-3, 102e-5t
 tumor markers in, 473t
 urticaria in, 135
- Lymphoid tissue, mucosa-associated. *See* Mucosa-associated lymphoid tissue (MALT)
- Lymphoma. *See* Lymphoid malignancies
- Lymphoma cutis, 364t, 366, 367
- Lymphoma-like disorders, 710
- Lymphomatoid granulomatosis, 135e-3, 1716, 2184
- Lymphomatoid papulosis, 710
- Lymphoocclusive bandage, for snakebite, 2735
- Lymphoplasmacytic lymphoma, 706–707
- Lymphoplasmacytic sclerosing pancreatitis. *See* IgG4-related disease (IgG4-RD)
- Lymphoscintigraphy, 1654
- Lymphotactin/SCM-1, 372e-11t
- Lyn, 2115
- Lynch's syndrome. *See* Hereditary nonpolyposis colon cancer
- Lyonization, 735
- Lysergic acid diethylamide (LSD), 469e-4
- Lysine metabolism disorders, 434e-3t
- Lysinuric protein intolerance (dibasic aminoaciduria), 332e-7t, 435e-1t, 435e-2
- Lysosomal acid lipase deficiency (cholesteryl ester storage disease), 432e-4t, 432e-5, 2443
- Lysosomal storage diseases, **432e-1**
 genetic considerations in, 432e-1
 glycogen disorders, 432e-4t. *See also* Pompe disease (acid α -1,4 glucosidase deficiency)
 glycoproteinoses, 432e-3t
 GM₂ gangliosidosis, 432e-3t. *See also* Tay-Sachs disease
 leukodystrophies, 432e-4t
 mucopolisaccharidoses, 432e-3t
 mucopolysaccharidoses, 432e-2t, 432e-5–6, 436e-5f
 neutral glycosphingolipidoses. *See* Fabry's disease; Gaucher's disease; Niemann-Pick disease
 neutral lipid disorders, 432e-4t
 pathogenesis of, 432e-1
- Lysosome(s), 94e-3, 432e-1
- Lysaviruses, 1299, 1304. *See also* Rabies
- M**
- Machado-Joseph disease (SCA3), 440t, 451e-1t, 2628–2629
- Machine operator's lung, 1681t
- Machupo/Bolivian hemorrhagic fever, 1320
- Machupo virus, 1306t
- Maciel virus, 1308t
- MAC infections. *See* *Mycobacterium avium* complex (MAC) infections
- Macitentan, 1659–1660, 1660t
- MACOP-B regimen, 135e-2
- Macroalbuminuria, 1616, 2425, 2425f
- Macroamylasemia, 2102
- Macrocytosis, 81e-3f, 393, 395f, 2067
- Macroglossia, 241t, 721, 721f
- Macrolides
 action of, 931, 931t, 942
 adverse effects of, 938t, 943
 drug interactions of, 939t
 indications for, 943
 resistance to
 in *M. pneumoniae*, 1164
 manifestations of, 807
 mechanisms of, 931t, 934
 in pneumococci, 807
 prevalence of, 966
 prevention of, 230
 in *S. pyogenes*, 233
 in *T. pallidum*, 1138
- Macroovalocytes, 81e-3f
- Macrophage(s)
 in adaptive immunity, 372e-7t, 372e-9f
 in asthma, 1672
 functions of, 421
 in HIV infection, 1238
 in innate immunity, 372e-4, 372e-6, 372e-7t, 372e-14f
 in pneumonia, 804
 in shock, 1747
 in smokers, 1701
 in tuberculosis, 1106–1107
- Macrophage colony-stimulating factor (M-CSF), 372e-11t
- Macrophage-derived chemokine. *See* MDC (CCL22)
- Macrophage inflammatory protein, 416
- Macrophage inflammatory protein-1 α (MIP-1 α), 372e-12t
- Macrophage inflammatory protein-1 β (MIP-1 β), 372e-12t
- Macrothrombocytopenia, 726f
- Macula
 anatomy of, 195
 cherry red spot, 40e-4, 40e-4f
 damage to, 197
 normal, 40e-1, 40e-1f
- Macula densa, 332e-2, 332e-2f
- Maculae ceruleae, 2746
- Macular degeneration
 diagnosis of, 205, 205f
 dry (nonexudative), 205
 gene therapy for, 91e-3
 pathogenesis of, 205
 prevention of, 96e-7
 treatment of, 205
 visual loss in, 205
 wet (exudative), 205
- Macular hole, 206
- Maculatum disease, 1155t
- Macules, 127, 240t, 339t, 340f
- Madrid virus, 1309t
- Madurella, 1355
- Mafenide acetate, 166e-3, 262e-4
- Maffucci's syndrome, 367, 426e-7, 594
- MAG (myelin-associated glycoprotein), 372e-24t, 444e-8, 718
- Maggots, 2747
- Magic mint (*Salvia divinorum*), 469e-4
- MAGIC syndrome, 2204
- Magnesium
 deficiency of. *See* Hypomagnesemia
 excess of. *See* Hypermagnesemia
 metabolism of, 2461
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-3t
 reference values for, 2761t
 renal transport of, 332e-4f, 332e-8
 supplements, 216, 2497
- Magnesium citrate, 62t
- Magnesium hydroxide, 62t, 1920, 1970t
- Magnesium sulfate
 for asthma exacerbations, 1679
 for cardiac arrest, 1769f
 for eclamptic seizures, 46
 for sedation in tetanus, 986
- Magnesium-wasting syndromes, 2461–2462, 2462t
- Magnetic resonance angiography (MRA)
 in frostbite, 478e-5
 in neurologic disease, 440e-6–8, 441e-16f
 in peripheral artery disease, 1643, 1644f
 phase-contrast, 440e-7

- of renal arteries, 1628, 1628t
in respiratory disease evaluation, 1666, 1667f
of thoracic aneurysm, 1639f
time-of-flight, 440e-3f, 440e-7
- Magnetic resonance cholangiography, 283
- Magnetic resonance cholangiopancreatography (MRCP)
of bile duct, 1892, 1895f, 2085t
in chronic pancreatitis, 2099, 2100f
in liver disease, 1993
in pancreatic cancer, 555
in pancreatic disease evaluation, 2087t, 2089
in primary sclerosing cholangitis, 139e-3, 2061–2062
- Magnetic resonance elastography, 1993, 1999
- Magnetic resonance imaging (MRI)
in adrenocortical carcinoma, 2323f
in ankylosing spondylitis, 2171, 2172f
for breast cancer screening, 480, 481t
in CAH, 2328f
in cardiac disease. *See* Cardiac magnetic resonance imaging (CMR)
contraindications to, 440e-8t
in Crohn's disease, 1954, 1955f
in Cushing's syndrome, 2317, 2317f
FLAIR. *See* Fluid-attenuated inversion recovery (FLAIR) MRI
in hearing loss evaluation, 223
in liver disease, 1993
in Ménière's disease, 219
in musculoskeletal disease evaluation, 2224t, 2225, 2225f
in NETs, 572
in neurologic disease. *See also* Neuroimaging, MRI
acute disseminated encephalomyelitis, 441e-25f
acute transverse myelitis, 441e-25f
adrenoleukodystrophy, 441e-41f
amyotrophic lateral sclerosis, 2632f
Bell's palsy, 441e-23f, 2648f
brachial plexopathy, 441e-30–31f
brain abscess, 901, 902f
brain metastases, 604f
brain tumors, 441e-39f, 598, 600f, 601f
CADASIL, 441e-18f
candidal brain infection, 441e-8f
cavernomas, 441e-38f
cavernous malformation, 440e-7f
central pontine myelinolysis, 1782, 1782f
cerebral abscess, 440e-5f
CNS aspergillosis, 441e-9–10f
CNS tuberculosis, 441e-2f
coccioidomycosis meningitis, 441e-7f
cranial epidural abscess, 905
dementia, 174, 174f
epidural metastases, 606f, 2653f
FTD, 2603f
Guillain-Barré syndrome, 441e-42f
Hashimoto's encephalopathy, 441e-29f
hemiplegic migraine, 441e-43f
hepatic encephalopathy, 441e-42f
histiocytosis, 441e-16f
histoplasmosis of the pons, 441e-6f
Huntington's disease, 441e-23f, 2622f
hypoxic-ischemic encephalopathy, 1781f
intracerebral hemorrhage, 2584
intramedullary astrocytoma, 2654f
Krabbe's disease, 441e-41f
Leigh's disease, 441e-40f
leptomeningeal metastases, 606f
limbic encephalitis, 441e-1f
MELAS, 441e-40f
meningioma, 603f
meningitis, 888
mesial temporal lobe epilepsy, 2546f
moyamoya disease, 441e-41f
MS, 441e-27–28f, 441e-35f, 2664–2665, 2666f
multiple system atrophy, 441e-22, 2641, 2641f
neurocutaneous melanosis, 441e-37f
neurofibromatosis type 2, 441e-36f
neurosyphilis, 441e-3–5f
primary central nervous system lymphoma, 601, 602f
progressive multifocal leukoencephalopathy, 899
retropharyngeal soft tissue mass, 441e-32f
seizures, 2550
sinonasal aspergillosis, 441e-10–11f
spinal arteriovenous malformation, 2658f
spinal cord infarction, 441e-25f
spinal epidural abscess, 2655
stroke, 2580, 2582f
Sturge-Weber syndrome, 441e-38f
subdural hematoma, 441e-34f
superior sagittal sinus thrombosis, 441e-20–22f
syringomyelia, 2658f
thoracic meningioma, 2654f
tuberous sclerosis, 441e-36f
vertebral fractures, 441e-31–34f
von Hippel-Lindau disease/syndrome, 441e-37f
in osteomyelitis, 840
of pituitary tumors, 2262, 2262f, 2278
in polydipsia, 2278
in respiratory disease evaluation, 1666, 1667f
in rheumatoid arthritis, 2144, 2144f
in sarcoidosis, 2209
of sellar masses, 2262
in spinal cord compression, 1791
in spinal disorders, 114f, 116f
in vestibular schwannoma, 603
in viral encephalitis, 440e-6f, 895, 895f, 895t
in Wernicke's disease, 1783f
- Magnetic resonance neurography, 440e-8
- Magnetoencephalography (MEG), 442e-3, 2550
- Mahaim fibers, 1481
- Ma huang (*Ephedra sinica*), 14e-3–4, 2547t
- Mahvash disease, 562, 569
- Maintenance dose, 36
- Majocchi's granuloma, 356
- Major depression, 2715, 2715t, 2716f. *See also* Depression/depressive disorders
- Major histocompatibility complex (MHC). *See* Human leukocyte antigen (HLA) complex
- MAL (methyl aminolevulinate), 386
- Malabsorption, surgery-related, 1927
- Malabsorption syndromes, 1932
approach to the patient, 1937
ataxia due to, 2630
bacterial overgrowth syndromes. *See* Bacterial overgrowth syndromes
celiac disease. *See* Celiac disease (gluten-induced enteropathy)
classification of, 1946t
clinical features of, 1937–1938, 1946t
diagnosis of
imaging, 1938, 1939f
Schilling test, 350e-1t, 350e-1t, 1938
small-intestinal mucosal biopsy, 1938–1940, 1940t, 1941f
urinary D-xylose test, 1938
pathophysiology of, 1946t
protein-losing enteropathy, 1945
short bowel syndrome. *See* Short bowel syndrome
tropical sprue. *See* Tropical sprue
Whipple's disease. *See* Whipple's disease
- Maladie de Roger, 51e-5
- Malaria, 1368
in children, 1374
chronic, 1374–1375
climate change and, 151e-4, 151e-8, 151e-8f
clinical features of, 152e-2t, 1372–1374, 1372t
acidosis, 318, 1373
encephalopathy, 780t, 783–784, 1373, 1373f, 1374t
fever, 1372
hematologic abnormalities, 1374
hypoglycemia, 1373, 1381
jaundice, 285
liver dysfunction, 1374
pulmonary edema, 1374, 1381
renal failure, 1374, 1381, 1836t, 1849
retinal hemorrhage, 1373, 1373f
complications of, 1374, 1374t, 1381
diagnosis of, 152e-2t, 245e-4t
alternative procedures, 1377t
blood films of parasites, 245e-2, 245e-6t, 250e-2–4f, 1375–1377, 1375–1377f, 1377t
vs. *Brucella* infection, 194e-2
laboratory findings, 1378
serologic and molecular tests, 245e-6t, 1377–1378, 1377t
drug research for, 13e-4
epidemiology of, 151e-8, 151e-8f, 152e-2t, 245e-4t, 1370–1371, 1370f
erythrocyte changes in, 1371
etiology of. *See Plasmodium* spp.
global considerations in, 14, 151e-4, 1370–1371, 1370f
in HIV infection, 798, 1268, 1374
host defense mechanisms in, 1371–1672
incubation period for, 152e-2t
pathogenesis of, 1368–1370, 1369f
in pregnancy, 1374
prevention of, 152e-2t
chemoprophylaxis, 796, 796t, 1382–1384, 1383t. *See also specific drugs*
in children, 1383t
indoor residual spraying, 14
insecticide-treated bed nets, 14
organizational efforts, 14–15, 1382
personal protection, 1382
in pregnancy, 1382
in travelers, 796, 796t, 1382–1384, 1383t
prognosis of, 1373t
resistance to
G6PD deficiency and, 656, 1371
hemoglobinopathies and, 633
ovalocytosis and, 1371
sickle cell anemia and, 442, 1371
thalassemias and, 1371
in returned traveler, 801–802, 802t
transfusion-transmitted, 138e-3t, 138e-6, 1374
transmission of, 1368–1370, 1369f
treatment of
antimalarial drugs, 246e-1–2t, 1380t. *See also specific drugs*
drug resistance and, 14–15, 1379, 1381f
Medicines for Malaria venture, 13e-4
regimens for, 14–15, 1378–1379, 1378t, 1379, 1381
in severe disease, 1378t, 1379
in uncomplicated disease, 1378t, 1379, 1381
in war veterans, 152e-1, 152e-2t, 152e-7
- Malassezia* spp., 1357
- Malassezia* spp. infections
clinical features of, 1357
diagnosis of, 1357
epidemiology of, 1357
folliculitis, 356, 1357
tinea versicolor. *See* Tinea versicolor
in transplant recipient, 921
treatment of, 1357
- Malathion, 2746
- Mal del pinto. *See* Pinta
- Maldonado virus, 1309t
- Male feminization, paraneoplastic, 609t
- Male pseudohermaphroditism. *See* 46,XY disorders of sex development
- Male reproductive system disorders, 2360
in adulthood, 2363
aging-related. *See* Androgen deficiency androgen insensitivity syndromes. *See* Androgen insensitivity syndrome (AIS)
diagnosis of, 2360–2361
erectile dysfunction. *See* Erectile dysfunction (ED)
gynecomastia. *See* Gynecomastia
hypogonadism. *See* Hypogonadism
hypogonadotropic hypogonadism. *See* Hypogonadotropic hypogonadism
infertility, 2360, 2388, 2388f, 2389
obesity and, 415e-6

- Male reproductive system disorders (*Cont.*):
 cancer. *See specific organs*
 in puberty
 delayed puberty, 2361t, 2362–2363
 precocious puberty. *See Precocious puberty, male*
 sex chromosome disorders. *See Sex chromosome disorders*
 testicular. *See Testes/testicles, disorders of 46,XY disorders. See 46,XY disorders of sex development*
- MALFDI-TOF MS (matrix-assisted laser desorption/ionization/time-of-flight mass spectrometry), 150e-3, 183e-1, 194e-3
- Malic dehydrogenase, 246e-10
- Malignant conversion, 387
- Malignant fibrous histiocytoma, 119e-1. *See also Soft tissue sarcoma*
- Malignant hypertension
 cerebral effects of, 1615, 1626
 epidemiology of, 1630
 genetic considerations in, 1630
 ocular effects of, 40e-5f, 1626
 renal effects of, 1615–1616, 1626, 1630, 1832
 treatment of, 1626–1627, 1626t, 1627t
- Malignant hyperthermia, 40–41, 450t, 462e-12
- Malignant pheochromocytoma, 2331
- Malignant transformation, 591
- Malingering, 2719
- Mallenkot catheter, 1978
- Mallory-Weiss tears
 bleeding in, 276t, 277, 1910
 diagnosis of, 1891, 1892f
 hematemesis in, 259
 treatment of, 277, 346e-1v, 1891, 1910
- Malnutrition, 459
 in acute kidney injury, 1810
 body mass index in, 98e-1t
 in chronic kidney disease, 1819
 in cirrhosis, 2066
 in elderly, 81
 fatigue in, 152
 global considerations in, 461
 gonadotropin deficiency due to, 2364
 hypothermia in, 478e-1
 in malabsorption syndromes, 1946t
 micronutrient, 461. *See also specific micronutrients*
 primary vs. secondary, 459
 protein-energy
 cachexia, 459–460, 1504
 kwashiorkor. *See Kwashiorkor*
 marasmus. *See Marasmus*
 risk of, 95e-5
- Malocclusion, 236
- Malpractice suits, 20
- MALT. *See Mucosa-associated lymphoid tissue (MALT)*
- Maltese cross formation, 62e-12f, 1846
- MALT lymphoma. *See Mucosa-associated lymphoid tissue (MALT) lymphoma*
- Malt worker's lung, 1681t
- MAMLD1 gene mutations, 2354t
- Mammalian target of rapamycin. *See mTOR*
- Mammary soufflé, 51e-6, 1449
- Mammography
 abnormalities detected on, 526, 526f
 diagnostic, 525
 screening, 28t, 29t, 480, 481t, 525. *See also Breast cancer, screening*
- Managed care systems, 5
- Manganese
 deficiency of, 96e-9t, 96e-10
 parental supplementation of, 98e-6t
 recommended intake of, 95e-3t
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t, 472e-3
- Manganese-56 exposure, 263e-3t
- Mania, 190, 2717, 2717t. *See also Bipolar disorder*
- Mannitol
 adverse effects of, 295, 297, 303, 309
 for ciguatera poisoning, 2742
- for increased intracranial pressure, 1780t
 for IV potassium delivery, 462e-17
- Mannose-binding lectin(s), 1752
- Mannose receptor, 372e-6t
- Mannosidosis, 432e-3t
- Manometry
 anorectal, 274
 esophageal. *See Esophageal manometry*
 intestinal, 260
- Mansonella ozzardi, 1418t, 1422
- Mansonella perstans, 1418t, 1422
- Mansonella streptocerca/streptocerciasis, 245e-6t, 1418t, 1422
- Mantle cell lymphoma, 704
 clinical features of, 702t, 704
 diagnosis of, 704
 genetic considerations in, 101e-3t, 699, 699t, 704
 staging of, 704
 treatment of, 103e-19, 704
 tumor markers in, 135e-2t
- MAO-B (monoamine oxidase type B) inhibitors, 2615, 2616t
- MAOIs. *See Monoamine oxidase inhibitors (MAOIs)*
- MAP3K1 gene mutations, 2354t
- MAP kinase pathway, in melanoma, 496, 496f
- MAP kinase phosphatase-1, 1677
- Maple syrup urine disease, 434e-3t
- Maprotiline, 2716
- MAPT gene mutations, 172t, 444e-1, 2603, 2604f
- Marantic endocarditis, 138, 367, 816
- Marasmus (starvation-related malnutrition)
 anemia in, 630–631
 cardiac manifestations of, 290e-1
 clinical features of, 459–460
 diagnosis of, 459–460
 global considerations in, 461
 vs. kwashiorkor, 460t
 physiologic characteristics of, 460t
- Maraviroc
 action of, 214e-3
 adverse effects of, 1275t, 1281
 drug interactions of, 205e-3, 1281
 genetic variations in response to, 39t, 1239, 1248–1249, 1280f
 for HIV infection, 1275t, 1281
 molecular structure of, 1277f
 resistance to, 1280f
- Marble bone disease (osteopetrosis), 426e-4
- Marburg's variant, MS, 2673
- Marburg virus, 1323, 1324f
- Marburg virus disease
 clinical features of, 783, 1327–1328, 1327t
 complications of, 1328
 control and prevention of, 1328–1329
 diagnosis of, 1328
 epidemiology of, 782–783, 1324, 1325f, 1326f
 pathogenesis of, 1326–1327
 prognosis of, 1328
 treatment of, 1328
- March hemoglobinuria, 657, 657t
- Marchiafava-Bignami disease, 2607
- Marcus Gunn pupil, 196, 196f
- Marfan's syndrome, 2512
 cardiac manifestations of, 100, 290e-1t, 2512
 aortic aneurysm, 1638, 1639
 aortic dissection, 1641
 murmur, 51e-5
 classification of, 2512
 diagnosis of, 2513
 genetic considerations in, 2513
 genetic testing for, 450t
 hand deformities in, 1443
 incidence of, 2512
 ocular features in, 2513
 oral deformities in, 1443
 paternal age and, 433, 438
 in pregnancy, 47
 skeletal manifestations of, 436e-6f, 2512
 spontaneous pneumothorax in, 2513
 treatment of, 445, 450–451, 450t, 2513
 vessel wall disorders in, 732
- Mariana Pastora (*Salvia divinorum*), 469e-4
- Maribavir, 215e-9, 922
- Marijuana, 469e-2
 acute and chronic intoxication, 469e-2
 dependence on, 469e-3
 dopamine levels and, 444e-4
 head and neck cancer risk and, 502
 hypogonadism and, 2364
 initial actions of, 465e-5t
 physical effects of, 469e-2–3
 prevalence of, 469e-2
 THC in, 469e-2
 therapeutic use of, 260t, 469e-3
 tolerance to, 469e-3
 withdrawal syndrome, 469e-3
- Marine envenomations, 2738
 approach to the patient, 2741–2742
 invertebrate, 2738–2740, 2738f, 2739f, 2741f
 treatment of, 2739, 2741–2742
 vertebrate, 2740–2741
- Marine poisoning, 2742
 ciguatera, 267, 2742, 2742t
 domoic acid intoxication, 2547–2548, 2743
 scombroid, 267, 2743–2744
 shellfish, 2743
- Marinesco-Sjögren syndrome, 451e-3t
- Maripa virus, 1308t
- Markesbery-Griggs distal myopathy, 462e-12, 462e-13t
- Market failure, 15e-1
- Maroteaux-Lamy disease, 432e-2t, 432e-6
- Marrara, 2747
- Marshall's syndrome, 2511
- Masaoka staging, of thymoma, 123e-1, 123e-1t
- Massage therapy, 14e-1t, 14e-3
- Massively parallel sequencing. *See Next-generation sequencing*
- Mass movements, intestinal, 265
- Mastadenovirus, 1208. *See also Adenovirus infections*
- Mast cell(s)
 activation of, 2114, 2115f
 in adaptive immunity, 372e-7t
 in allergy, 2115–2117, 2115f
 in asthma, 1672
 distribution of, 2115
 hematopoietic differentiation of, 89e-3f
 in innate immunity, 372e-7t
 membrane-derived lipid mediators from, 2115, 2115f
 proliferation of. *See Mastocytosis*
 secretory granules of, 2114
- Mast cell leukemia, 2120
- Mastectomy
 for breast cancer, 103e-3, 527–528
 complications of, 484
 male, 532
 prophylactic, 479
- “Master gland.” *See Pituitary gland*
- Master's disease (Southern tick-associated rash illness), 129t, 134, 1152
- Mastitis, 957, 1112
- Mastocytosis
 cutaneous, 135e-8, 357, 363, 367, 2119
 small-intestinal mucosal biopsies in, 1940t
 systemic, 2120
 aggressive, 135e-8, 2120
 associated with clonal hematologic nonmast cell lineage disease, 2120
 classification of, 18t, 2120
 clinical features of, 135e-8, 254, 269, 2119, 2120
 definition of, 2120
 diagnosis of, 135e-8, 2120–2121, 2121t
 differential diagnosis of, 2121
 indolent, 2120
 pathophysiology of, 2120
 prognosis of, 135e-8
 treatment of, 135e-8, 2121
- Mastoidectomy, 230, 231
- Mastoiditis, 230–231, 231f, 900
- Maternal age, trisomy and, 83e-4
- Maternal inheritance, 85e-1, 85e-2f. *See also Mitochondrial DNA diseases; X-linked disorders*

- Maternal mortality, 51
 Maternal serum screening, 83e-4
 Matrilin-1, 2202
 Matrix-assisted laser desorption/ionization/time-of-flight mass spectrometry (MALFDI-TOF MS), 150e-3, 183e-1, 194e-3
 Matrix metalloproteinase 3, 2170
 Matrix metalloproteinase 9, 1106
 Matrix metalloproteinase 12, 1701
 Matrix metalloproteinases (MMPs)
 in cancer cell signaling, 102e-11f
 in metastasis, 102e-10
 in photoaging of skin, 387
 in rheumatoid arthritis, 2143
 in tumor angiogenesis, 102e-15, 102e-17f
 Mat telangiectasias, 356f, 357
 Maturity-onset diabetes of the young (MODY), 431t, 441t, 2400, 2406
 Maximum-tolerated dose (MTD), 103e-5
 Mayaro virus, 837, 1306t
 Mayer-Rokitansky-Kuster-Hauser syndrome, 335, 2357
 May-Hegglin anomaly, 729
 Mazindol, 2396
 Mazzotti reaction, 246e-8
 MBC (minimal bactericidal concentration), 150e-9
 MBL gene, 1240t
 MBP (myelin basic protein), 372e-24t, 444e-4f
 McArdle's disease (muscle phosphorylase deficiency; type V GSD), 433e-2t, 433e-4, 462e-14
 McBurney's point, 1987
 McConnell's sign, 270e-7, 1634
 McCune-Albright syndrome, **426e-6**
 clinical features of
 adrenal tumors, 2321
 Cushing's syndrome, 2314-2315
 cutaneous, 359, 360t
 endocrinologic, 2266, 2344
 fibrous dysplasia, 426e-6, 2265t
 precocious puberty, 2362
 diagnosis of, 426e-6, 426e-6f
 genetic considerations in, 439, 2252, 2266, 2314-2315, 2344, 2362, 2476f
 treatment of, 426e-7
 MCH (mean cell hemoglobin), 393, 393t, 394t, 2755t
 MCH (melanin-concentrating hormone), 415e-2
 MCH/MCV (mean cell hemoglobin concentration), 393, 393t
 MCI (mild cognitive impairment), 75, 76f, 171
 MCIIR gene mutations, 386, 494
 McLeod blood group, 81e-4f
 McLeod phenotype, 138e-2, 375e-1
 McMurray test, 2222
 M component
 in heavy chain diseases, 718
 of immunoglobulin, 710-712
 in multiple myeloma, 715
 in plasma cell disorders, 712
 in Waldenström's macroglobulinemia, 718
 MCP-1 (monocyte chemoattractant protein-1), 372e-11t, 465e-4
 MCP-2 (monocyte chemoattractant protein-2), 372e-11t
 MCP-3 (monocyte chemoattractant protein-3), 372e-12t
 MCP-4 (monocyte chemoattractant protein-4), 372e-12t
 MCPs (monocyte chemoattractant proteins), 1242
 M-CSF (macrophage colony-stimulating factor), 372e-11t
 MCT8 gene mutations, 2286
 MCTD. *See* Mixed connective tissue disease (MCTD)
 MCV (mean cell volume), 393, 394t
 MD-2, 145e-5, 145e-6f
 MDC (CCL22), 372e-12t, 1674, 1674f
 MDCT (multidetector computed tomography), 440e-1, 1664-1666. *See also* Computed tomography (CT)
 MDM2 gene mutations, 562
 MDM3 gene mutations, 2077
 MDM4 gene mutations, 562
 MDMA (ecstasy), 168, 301, 469e-3
 MDRD equation, 291
mdr (multidrug resistance) gene/phenotype, 102e-10, 103e-23
 MDS (myelodysplastic syndromes). *See* Myelodysplasia
 Meadowsweet, 404t
 Mean arterial pressure, in shock, 1729, 1744, 1757
 Mean cell hemoglobin (MCH), 393, 394t, 2755t
 Mean cell hemoglobin concentration (MCH/MCV), 393, 394t, 2755t
 Mean cell volume (MCV), 393, 394t, 2755t
 Means-Lerman scratch, 290e-2
 Measles (rubeola), **1295**. *See also* Rubella (German measles)
 approach to the patient, 1297
 atypical, 25e-5f, 362
 clinical features of, 128t, 1296, 1296f, 1297
 in impaired immunity, 1297
 Koplik's spots, 25e-1f, 127, 362, 1297
 rash
 atypical, 25e-5f
 characteristics of, 25e-1f, 128t, 362, 362t, 1297
 stages of, 127, 362, 1297
 complications of, 900, 1298, 2673
 diagnosis of, 1297-1298
 differential diagnosis of, 1297
 epidemiology of, 1295-1296
 eradication of, 1299
 etiology of, 1295
 global considerations in, 786, 1295
 immune response to, 1296f, 1297
 incidence pre- and postvaccine, U.S., 785t
 pathogenesis of, 1296, 1296f
 postexposure prophylaxis, 1298
 prevention of, 1298-1299. *See also* Measles-mumps-rubella vaccine (MMR)
 prognosis of, 1298
 transmission of, 1296
 treatment of, 1299
 vitamin A deficiency in, 96e-7
 Measles-mumps-rubella vaccine (MMR)
 adverse effects of, 231e-4, 1299
 contraindications for, 789t
 coverage with, in U.S., 231e-1
 precautions for, 789t
 recommended administration of
 in adults, 29t, 50-51, 230e-3, 787-788f
 in cancer patient, 485t
 in children, 230e-3, 230e-3f, 231e-4, 1298-1299
 in health care personnel, 231e-4
 in HIV infection, 798
 in transplant recipient, 929, 929t
 storage and handling of, 791
 Measles vaccine, 794, 1298-1299
 Measles virus, 145e-1, 145e-2t, 1295. *See also* Measles (rubeola)
 Mebendazole
 action of, 246e-10
 adverse effects of, 246e-3t, 246e-10
 for ascariasis, 1413
 drug interactions of, 246e-3t
 for enterobiasis, 1416
 for hookworm infections, 1415
 indications for, 246e-3t
 pharmacology of, 246e-10
 in pregnancy and lactation, 246e-3t
 for trichinellosis, 1411t
 for trichuriasis, 1416
meA gene, 774, 807, 961
 Mecamylamine, 2660
meC gene, 774
 Mechanical heart valve. *See* Prosthetic heart valve
 Mechanical ptosis, 207
 Mechanical ventilation, **1740**
 for ARDS, 1738-1739, 1739t
 in coma, 1776
 complications of, 1743
 alveolar collapse, 1738
 lung injury, 1731, 1738, 1743
 pneumonia. *See* Ventilator-associated pneumonia (VAP)
 respiratory acidosis, 322
 volutrauma, 1731, 1743
 continuous positive airway pressure. *See* Continuous positive airway pressure (CPAP)
 for COPD, 1707
 for hemoptysis, 247
 indications for, 1740
 modes of, 1741-1742, 1742t
 nonconventional strategies for, 1742-1743
 noninvasive, 1740-1741, 1741t
 patient management during, 1732-1733, 1739, 1741, 1743
 physiology of, 1741, 1741f
 principles of, 1741, 1741f
 protective strategies in, 1743
 for pulmonary edema, 1763
 for respiratory acidosis, 323
 for sepsis/septic shock, 1755
 for shock, 1731
 tracheostomy in, 1744
 weaning from, 66-67, 1743-1744, 1744f
 Mechanic's hands, 2195
 Mechanoreceptors, in lung, 47e-1
 Mechanotransduction, 217
 Mechlorethamine (nitrogen mustard)
 action of, 103e-8
 adverse effects of, 103e-8, 103e-9t, 103e-25, 474, 607t
 carcinogenicity of, 477t
 as terrorism agent, 262e-1t
 Meckel-Gruber syndrome, 1850t, 1851f
 Meckel's diverticulum, 277, 279
 Mecizine, 62, 260, 260t, 261
 Meclofenamate, 727t
 Meconium ileus, 1697
MECP2 gene mutations, 431, 444e-4, 465e-2, 465e-2t
 Médecins Sans Frontières, 12
 Medial epicondylitis, 2249
 Medial medullary stroke syndrome, 2576f, 2577
 Median neuropathy, 2683, 2691
 Mediastinal (thymic) large B-cell lymphoma, 135e-2
 Mediastinal mass
 CT in, 1664, 1665f, 1719
 diagnosis of, 1719
 differential diagnosis of, 123e-1
 thymoma. *See* Thymoma
 Mediastinitis
 actinomycotic, 1089
 acute, 1719
 anaerobic bacterial, 1098
 chronic, 1719-1720
 fibrosing, in histoplasmosis, 1333, 1334
 nocardial, 1085
 sternal osteomyelitis and, 844
 in transplant recipient, 926, 927
 Mediastinoscopy, 1668
 Mediastinotomy, 1668
 Mediastinum, 1719, 1720t
 Medicaid, 15e-2, 82-83, 83t
 Medical decision-making, 3, 16e-4
 Medical education, 7, 17e-4
 Medical errors, 4, 17e-4
 Medical kit, traveler's, 797
 Medical practice, 1, 2f
 accountability in, 6-7
 clinical skills and, 1-3
 ethical issues in. *See* Ethical issues
 evaluation of outcomes, 3
 expanding frontiers of, 5-7
 globalization of, 6. *See also* Global health
 humane aspects of, 5
 Internet and, 6
 legal issues in. *See* Legal issues
 practice guidelines, 3
 public expectations of, 6-7

- Medical practice (*Cont.*):
 research and, 7
 trends in health care delivery, 5
- Medical records, electronic, 3
- Medical tourism, 799
- Medicare
 fee schedule, 15e-3
 health care–acquired conditions not eligible for coverage by, 912t
 hospice benefits, 70
 service covered by, 82, 83t
- Medication-overuse headache, **109**, 2593
- Medicinal leech, 274f
- Medicine, **1**
 art of, **1**
 economic issues in. *See* Economic issues
 evidence-based. *See* Evidence-based medicine
 practice of. *See* Medical practice
 science of, **1**
- Meditation, 14e-1t
- Mediterranean fever, familial. *See* Familial Mediterranean fever
- Mediterranean lymphoma, 537, 718–719
- Mediterranean spotted (boutonneuse) fever, 128t, 782, 1155t, 1157
- Medium-chain fatty acids, 1935, 1935t
- Medium-chain triglycerides, 98e-5, 1936
- MEDLINE, 25, 25t
- MEDNIK syndrome, 2519
- Medroxyprogesterone
 adverse effects of, 7e-6
 for breast cancer, 103e-15
 for contraception, 2391
 for hot flushes with androgen deprivation therapy, 7e-6
 for menorrhagia in perimenopause, 2382
 for polycystic ovarian syndrome, 337
- Medscape, 14e-4t
- Medulla, 2577f
- Medullary cystic kidney disease, 1850t, 1854
- Medullary sponge kidney, 1850t, 1855, 1866
- Medullary stroke syndrome, 2576f
- Medullary thyroid carcinoma (MTC), **2341**
 clinical features of, 269
 diagnosis of, 2341, 2341f
 familial, 2307, 2340–2341, 2342
 genetic considerations in, 2341, 2342
 in MEN 2, 2307, 2332, 2332f, 2340, 2341
 paraneoplastic syndromes in, 609t
 prevalence of, 2303t
 staging of, 2304t
 treatment of, 102e-5t, 272, 2341–2342
 tumor markers in, 473t
- Medulloblastomas, 599, 602
- Mees' lines, 2608, 2689
- Mefenamic acid, 338, 2382
- mef* gene, 807
- Mefloquine
 action of, 246e-10
 adverse effects of, 246e-5t, 246e-10, 1380t, 1381, 1383
 contraindications to, 1383
 drug interactions of, 246e-2t, 246e-5t, 246e-10, 798
 for malaria, 1378t, 1380t
 for malaria prophylaxis, 796t, 1382, 1383t
 pharmacology of, 246e-10, 1380t
 in pregnancy, 246e-5t, 246e-10, 1382
 resistance to, 1381f
- MEG (magnetoencephalography), 442e-3, 2550
- Megacolon, 273, 274, 1395. *See also* Toxic megacolon
- Megakaryocytes, 677
- Megalin, 400e-3
- Megaloblastic anemia, **641**
 autosomal recessive, 645
 biochemical basis of, 642f
 clinical features of, 642–643
 drug-induced, 647
 etiology of, 640t, 644t
 hematologic findings in, 643–644, 644f
 in orotic aciduria, 649
 peripheral blood smear in, 644f
- thiamine-responsive, 435e-1t, 649
 treatment of, 648
- Megarectum, 274
- Megestrol, 81, 310, 1795
- Meglumine antimonate, 246e-2t, 246e-7, 1390
- Meibomian gland carcinoma, 199
- Meibomitis, 199
- Meigs' disease/syndrome (lymphedema praecox), 594, 1653–1654, 2620
- Meiosis, 427–428
- MEK1 gene mutations, 507t, 509f
- MEK inhibitors, 102e-3, 102e-4f
- Melamine, 1804
- Melanin-concentrating hormone (MCH), 415e-2
- Melanocortin-2 receptor, 401e-4, 2311, 2312f
- Melanocyte(s), 386
- α -Melanocyte-stimulating hormone (MSH)
 in appetite control, 415e-3, 415e-4f, 415e-4t
 ectopic ATCH production and, 611–612
 overproduction of, 360
 sun response in skin and, 386
- Melanocytic nevus, 359
- Melanoma, 76e-6, **493**
 acral lentiginous, 76e-7f, 493f, 495, 495t
 amelanotic, 495
 AV conduction block in, 1471t
 carcinoma of unknown primary and, 120e-2t
 clinical classification of, 495, 495t
 clinical features of, 493f, 494–495
 deaths from, 467t
 desmoplastic, 495
 diagnosis of, 76e-6, 494–495
 dysplastic nevi and, 493f
 epidemiology of, 388, 493
 of eye, 40e-5f, 206, 206f
 familial, 101e-6t, 554
 genetic considerations in, 32, 101e-2t, 101e-3t, 494, 496, 496f
 incidence of, 467t, 468f
 KIRs with, 372e-15t
 lentigo maligna, 76e-7f, 493f, 495, 495t
 leukoderma in, 359
 lymphadenopathy in, 408
 metastatic, 289e-2, 360, 360t, 498–499, 553, 604t
 molecular classification of, 495–496, 496f
 nevi and, 493–494
 nodular, 76e-7f, 495, 495t
 oral, 240t
 pathogenesis of, 372e-24t, 388, 495–496, 496f
 during pregnancy, 124e-1t, 124e-4
 prevention of, 494
 prognosis of, 496, 497t
 retinopathy in, 614t, 619
 risk factors for, 493–494, 493t, 1962
 screening for, 481
 sentinel node biopsy in, 496–497
 staging of, 496, 497t
 sun/UV exposure and, 388, 476, 477t, 493, 494
 superficial spreading, 76e-7f, 339f, 495, 495t
 treatment of
 biologic therapy, 102e-17, 103e-22, 372e-30, 498–499
 chemotherapy, 499
 clinically localized disease, 496–497
 follow-up, 499–500
 metastatic disease, 497–499, 498t
 targeted chemotherapy, 102e-3, 102e-5t, 103e-17t, 103e-19, 499
- Melanopsin, 195
- Melanosis, neurocutaneous, 441e-37f
- Melanosis coli, 273
- Melanosomes, 386
- Melarsoprol, 246e-5t, 246e-10, 1398, 1398t
- Melasma, 341t, 353, 379
- MELAS (mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes) syndrome
 clinical features of, 85e-5, 85e-6t, 451e-3t, 462e-16, 2483
 diagnosis of, 85e-5, 441e-40f
 mitochondrial DNA mutation in, 85e-5, 85e-6t, 451e-3t, 462e-16
- Melatonin, 187, 193, 2595t
- MELD (Model for End-Stage Liver Disease) score, 1995, 1996, 2069, 2069t
- Melena, 276
- Meleney's gangrene, 1099
- Meleney's ulcer, 369
- Melioidosis, 152e-4t, 1045t, 1048
- Melkersson-Rosenthal syndrome, 2648
- Melodic intonation therapy, 178
- Melorheostosis, 426e-5
- Melphalan
 action of, 103e-8
 adverse effects of, 103e-8, 103e-9t, 621, 2025
 limb perfusion for melanoma, 498
 for multiple myeloma, 716–717
- Memantine, 176, 2601
- Membrane receptor(s), 400e-3–4, 400e-3f
- Membrane receptor families, 400e-1t
- Membrane transport, defects of, 435e-1–2, 435e-1t
- Membranoproliferative glomerulonephritis
 autoantibodies in, 372e-25t
 classification of, 1841t
 clinical features of, 1835t, 1841
 pathogenesis of, 1832, 1840–1841, 1841f
 renal biopsy in, 62e-3f, 62e-4f, 1841
 treatment of, 1841
- Membranous glomerulonephritis
 clinical features of, 1835t, 1844
 etiology of, 1843, 1843t
 pathogenesis of, 1843–1844, 1844f
 renal biopsy in, 62e-3f, 1843
 treatment of, 1844
- Memory
 assessment of, 183, 2537
 immune, 372e-2
 immunologic, 372e-16
 impairment of, rates by age group, 72t
 implicit, 182
 limbic network for, 182
 loss of. *See* Amnesia
 retentive, 183
 working, 183
- MEN1/MENIN gene mutations
 genetic testing for, 2340
 in GI-NETs, 562–563, 562t
 in MEN syndromes, 599t, 1928, 2251, 2265–2266, 2336t, 2340, 2471, 2471f
 tumors associated with, 101e-6t
 variable expressivity of, 437
- Menaquinone, 96e-8. *See also* Vitamin K
- Menarche, 335, 524, 2379, 2379t
- Mendelian disorders, 437, 437f
 autosomal dominant, 437–438
 autosomal recessive, 438
Online Mendelian Inheritance in Man, 425, 426t
 X-linked. *See* X-linked disorders
 Y-linked, 439
- Mendelian susceptibility to mycobacterial disease (MSMD), 2106
- Mendelson's syndrome, 1098
- Ménétrier's disease, 534, **1932**, 1945
- Ménière's disease
 clinical features of, 149, 150, 219, 220
 diagnosis of, 149, 150, 223
 epidemiology of, 219
 hearing loss in, 219, 222
 nystagmus in, 210
 pathophysiology of, 150, 219
 treatment of, 150, 219
- Meningeal biopsy, 910
- Meningeal syphilis, 1136
- Meningiomas
 clinical features of, 602
 imaging in, 602, 603f, 2262
 in MEN 1, 2340
 ocular signs in, 198
 pathophysiology of, 602
 sellar, 2263
 thoracic, 2654, 2654f
 treatment of, 602

- Meningitis**
 approach to the patient, 883–884, 884–885f
 bacterial, **885**
 A. baumannii, 1037
 clinical features of, 783, 887, 998–999
 complications of, 952
 deaths from, 890
 definition of, 885
 diagnosis of, **887**
 blood culture, 887, 951
 CSF analysis, 764, 766t, 888t, 951
 signs and symptoms, 998–999
 differential diagnosis of, 888, 951
 E. coli, 1028
 enterococcal, 974, 975t
 epidemiology of, 885
 etiology of, 780t, 783, 885–886
 gram-negative, 889t, 890
 H. influenzae, 886, 1011
 incidence of, 885
 increased ICP in, 890
 in infants, 970
 L. monocytogenes, 886, 889–890, 889t, 983
 meningococcal, 780t, 885, 889, 889t, 998–999
 nontyphoidal *Salmonella*, 1054, 1054t
 pathophysiology of, 886–887, 886f
 plague, 1072
 pneumococcal, 780t, 885, 889, 889t, 951
 prognosis of, 890
 prophylaxis for exposure to, 945t
 risk factors for, 783
 staphylococcal, 886, 889t, 890
 streptococcal, 970
 T. pallidum, 1135
 treatment of
 adjunctive, 890
 algorithm for, 884–885f
 antimicrobial therapy based on pathogen, 889–890, 889t, 975t, 1011
 empirical antimicrobial therapy, 767t, 780t, 887t, 889, 1000
 ICP-related, 890
 in cancer patient, 488, 1792–1793
 chronic, **906**
 amebic, 1367–1368
 approach to the patient, 907, 909
 bacterial, 907–908t
 basal, 906
 clinical features of, 906, 906t
 diagnosis of, 907–909t, 909–910, 910f
 epidemiology of, 907
 fungal, 908t
 helminthic, 908t
 in HIV infection, 907, 911
 in immunosuppressed patients, 911
 intracranial, 906
 Mollaret's, 909t
 noninfectious causes of, 909t
 pathophysiology of, 906
 protozoal, 908t
 spinal, 906–907
 systemic manifestations of, 907
 treatment of, 910–911, 945t
 viral, 908t
Coccidioides, 441e–7f, 1336, 1337
 complications of, 974
Cryptococcus, 1264, 1340, 1341
 delirium in, 169
 fungal, 898
 global considerations in, 898
 headache in, 108
 hyponatremia in, 299
 Mollaret's, 1179
 neoplastic, 1792–1793
 in neurosyphilis, 899, 1136
 pathophysiology of, 145e–8
 in secondary syphilis, 1135
 subacute, 898–899
 tuberculous, 766t, 775, 898–899, 907t, 1111
 viral, **890**
 arbovirus, 892
 chronic, 908t
 clinical features of, 890
 diagnosis of, 764, 766t, 891
 differential diagnosis of, 892
 EBV, 892
 enteroviral, 892, 1291, 1291t
 epidemiology of, 890
 etiology of, 890, 891t, 892–893
 in HIV infection, 892–893, 1264
 HSV, 892, 908t, 1179, 1182t
 lymphocytic choriomeningitis, 891, 893, 908t, 1306t, 1317–1318
 in mumps infection, 231e–2–3, 893
 prognosis of, 893
 treatment of, 893
 VZV, 892
 weakness in, 156
Meningococcal infections, 995
 in cancer patient, 484, 484t, 486t, 488
 clinical features of, 998–999
 cutaneous, 362, 780t, 998, 999t
 meningitis. *See* Meningitis
 meningococcemia/septicemia. *See* Meningococcemia
 postmeningococcal reactive disease, 999
 postsplenectomy sepsis, 780t, 781
 complications of, 1000
 diagnosis of, 999–1000
 epidemiology of, 996–997, 996f, 997f
 etiology of. *See* *Neisseria* spp. (meningococci); *Neisseria meningitidis*
 genetic considerations in, 997
 global considerations in, 995–996
 pathogenesis of, 998
 prevention of. *See* Meningococcal vaccine
 prognosis of, 1000
 prophylaxis for contacts, 1002–1003
 risk factors for, 997
 treatment of, 940t, 1000
Meningococcal vaccine
 based on subcapsular antigens, 1002, 1002f
 conjugate, 1001–1002
 contraindications to, 790t
 global considerations in, 1001, 1002
 polysaccharide, 1001, 1001f
 recommended administration of
 in adults, 787–788f
 in cancer patient, 485t, 492
 in splenectomized patient, 413
 in transplant recipient, 920t, 929, 929t
 in travelers, 795, 795t
 storage and handling of, 791
Meningococcemia
 acute, 133t, 135
 chronic, 25e–11f, 133t, 135, 999
 clinical features of, 781, 999
 complications of, 781–782, 1000
 epidemiology of, 781
 etiology of, 780t, 781
 fulminant, 25e–11f, 1754, 1755
 pathogenesis of, 998
 prognosis of, 1000
 shock in, 998
 skin manifestations of
 papular lesions, 25e–11f
 petechiae, 781
 purpura, 25e–11f, 76e–9f, 133t, 342f, 369, 998
 rash, 362, 362t
 treatment of, 780t, 1000
Meningoencephalitis
Balamuthia, 1368, 1368f
Cryptococcus, 1340, 1340f
L. monocytogenes, 983
Naegleria, 1367
 Meningovascular syphilis, 1136
 Menkes' kinky hair syndrome, 96e–10
Menopause, 2381
 age at, breast cancer risk and, 524
 bone remodeling and, 2491–2492, 2492f
 cardiovascular disease risk and, 6e–3
 clinical features of, 2382, 2382f
 definition of, 2381
 hirsutism after, 332
 hormone therapy in. *See* Postmenopausal hormone therapy
 physiology of, 2381, 2381f
 premature, after cancer treatment, 103e–25, 623
 prevalence of, 2254t
 screening/testing for, 2254t, 2381–2382, 2382f
 transition to, 2382, 2382f
 Menorrhagia, 403, 730, 2382
 Men's health, **7e–1**
 anabolic-androgen steroid abuse. *See* Anabolic-androgenic steroid (AAS) abuse
 androgen deprivation therapy. *See* Androgen deprivation therapy (ADT)
 breast cancer, 523, 532
 lower urinary tract symptoms, 7e–4, 60e–1.
 See also Benign prostatic hypertrophy (BPH)
 muscle dysphoria syndrome, 7e–2
 prostate cancer. *See* Prostate cancer
 reproductive function, 7e–1–2. *See also* Erectile dysfunction (ED); Male reproductive system disorders
 testosterone and, 7e–1, 7e–1f, 7e–2f. *See also* Testosterone; Testosterone therapy
 Menstrual cycle
 duration of, 400e–6
 follicular phase of, 2378–2379, 2379f
 hormonal integration of, 400e–5, **2378**, 2378f
 luteal phase of, 2379, 2379f
Menstrual disorders, 335
 amenorrhea. *See* Amenorrhea
 definition of, 335
 diagnosis of, 335, 337f
 dysmenorrhea, 338–339
 etiology of, 335–336
 hypothalamic-pituitary-gonadal axis in, 335, 336f
 menorrhagia, 403, 730
 obesity and, 415e–6
 prevalence of, 335
 sacral pain in, 118
 treatment of, 335–336
MEN syndromes. See under Multiple endocrine neoplasia
 Mental disorders. *See* Psychiatric/psychological disorders
 Mental status assessment, 2537. *See also* Mini-Mental Status Examination (MMSE)
 Men who have sex with men (MSM)
 gonococcal infections in, 1005–1006
 HPV infection in, 1198
 intestinal infections in, 881
 sexually acquired proctitis in, 881–882
 MEONs. *See* Multiple endocrine and other organ neoplasias (MEONs)
 MEOS (microsomal ethanol oxidizing system), 2058
 Meperidine
 adverse effects of, 462e–20, 462e–20t, 2547t
 dosage in kidney disease, 37, 1821
 dosage in liver disease, 37
 overdosage/poisoning with, 473e–13t
 for pain, 91t
 Mephenytoin, 34t
 Mepolizumab, 135e–8, 422, 1686, 2187
 Meproamate, 473e–11t
 Meralgia paresthetica (lateral femoral cutaneous neuropathy), 49, 463e–5, 463e–5f, 2692
 6-Mercaptopurine (6-MP)
 action of, 103e–14
 adverse effects of, 103e–10t, 103e–14, 417, 649, 1960, 1962
 drug interactions of, 42, 42t, 103e–10t
 genetic variations in response to, 39t, 40, 445, 1960
 for IBD, 1960
 metabolism of, 34t, 40, 42
 Mercury exposure/poisoning
 clinical features of, 472e–3t, 2608
 diagnosis of, 472e–3t
 global considerations in, 472e–3
 neuropathy due to, 2688t, 2689
 in pregnancy, 472e–1

- Mercury exposure/poisoning (*Cont.*):
 reference range for, 2766t
 skin manifestations of, 240t, 382
 treatment of, 472e-3t
- Merkel cell carcinoma, 502, 926, 928
- Merogony, 1369
- Meropenem
 for *Acinetobacter* infections, 1037t
 for anaerobic bacterial infections, 1101, 1101t
 for *B. cepacia* infection, 1045t
 for community-acquired pneumonia, 767t, 808t
 for febrile neutropenia, 491
 for glanders, 1045t
 for health care-associated pneumonia, 812t
 indications for, 940t, 941
 for intraabdominal infections, 767t
 for melioidosis, 1045t
 for meningitis, 887t, 889, 889t
 for multidrug-resistant TB, 205e-7
 for osteomyelitis, 841t
 for *P. aeruginosa* infections, 808t, 1045t
 in pregnancy and lactation, 937t
 resistance to, 933, 940t
 for sepsis/septic shock, 767t, 1757t
 for Whipple's disease, 1093-1094
- Merosin deficiency, 462e-9, 462e-10t
- Merozoites, *Plasmodium*, 1369
- MERRF (myoclonic epilepsy with ragged red fibers) syndrome
 clinical features of, 85e-5, 85e-6t, 451e-3t, 462e-16
 diagnosis of, 85e-7, 462e-16
 mitochondrial DNA mutation in, 85e-5, 85e-6t, 85e-7, 451e-3t, 462e-16
- MERS (Middle East respiratory syndrome), 261e-9, 1204-1205
- Mesalamine, 1959-1960, 1959t
- Mesangial cells, 332e-1
- Mesangioproliferative glomerulonephritis, 1835t, 1841
- Mesenchymal stem cells, 90e-2, 453t
- Mesenchymal tumor, 609t
- Mesenteric ischemia/vascular insufficiency. *See* Intestinal ischemia
- Mesenteric lymphadenitis, 106t
- Mesenteric venous thrombosis, 1979, 1980t
- Mesial temporal lobe epilepsy, 2544, 2546f, 2546t
- MESNA, 103e-8, 139e-3, 623, 1798
- Mesoamerican nephropathy, 1855-1856
- Mesocardia, 1528
- Mesothelin, 120e-2t
- Mesothelioma
 asbestos-related, 1689
 AV conduction block in, 289e-2, 1471t
 carcinoma of unknown primary and, 120e-2t, 120e-4
 clinical features of, 1577
 immunohistochemistry of, 508
 intramyocardial, 289e-2
 pericardial effusion in, 1577
 pleural effusion in, 1717
- Meta-analysis, 26
- Metabisulfite, as asthma trigger, 1671
- Metabolic acidosis, 317
 acid-base nomogram, 316f
 in acute kidney injury, 1808, 1810
 alcoholic ketoacidosis, 299, 318-319
 in chronic kidney disease, 1814
 clinical features of, 317
 compensatory responses in, 316t
 diabetic ketoacidosis. *See* Diabetic ketoacidosis (DKA)
 drug- or toxin-induced, 64e-2, 64e-9, 319-330. *See also* Ethylene glycol poisoning; Methanol poisoning
 high-anion gap, 64e-9, 318, 318t
 hyperkalemia in, 64e-2-3, 309, 311
 lactic acid. *See* Lactic acidosis
 in mixed acid-base disorders, 316-317, 316t
 non-anion gap, 64e-6, 320-321, 320t
 in overdose/poisoning, 473e-3
 with parenteral nutrition, 98e-7
 treatment of, 317-318
- Metabolic alkalosis, 321
 acid-base nomogram, 316f
 clinical features of, 321-322
 compensatory responses in, 316t
 differential diagnosis of, 321
 etiology of, 321-322, 321t
 hypokalemia in, 64e-8, 307
 mixed acid-base disorders, 316t
 in mixed acid-base disorders, 317
 pathogenesis of, 321
 treatment of, 322
- Metabolic disease. *See also specific diseases*
 classification of, 436e-1
 clinical features of, 436e-1, 436e-1-8f
 coma due to, 1772-1773
 etiology of, 436e-1
 myopathy in, 462e-19
- Metabolic encephalopathy, 178, 1782
- Metabolic equivalent tasks (METs), 1585t, 1587, 1587t
- Metabolic rate. *See* Resting (basal) metabolic rate
- Metabolic syndrome, 2449
 adiponectin in, 2452
 age and, 2449
 cardiovascular disease and, 1439, 2450, 2452
 clinical features of, 291e-6, 2452
 diabetes mellitus and, 2405, 2449, 2452, 2454
 diagnosis of, 2450t, 2452
 epidemiology of, 2449, 2450f
 HCV infection and, 2041
 in HIV infection, 1259
 hypertension in, 291e-6, 1617, 2451-2452, 2454
 hyperuricemia/uric acid stones in, 431e-4, 2452
 insulin resistance in, 2450, 2454
 lipodystrophy and, 2450
 lipoprotein disorders in, 2451, 2453-2454
 nonalcoholic fatty liver disease and, 2452
 obesity/overweight and, 2449
 obstructive sleep apnea and, 2452
 pathophysiology of, 292e-1v, 2450-2452, 2451f
 in polycystic ovarian syndrome, 337, 2452
 prevalence of, 2254t
 proinflammatory cytokines in, 2452
 in psoriasis, 348
 risk factors for, 2449-2450
 screening/testing for, 2254t
 thrombosis risk in, 744
 treatment of, 291e-7, 2452-2454
 waist circumference in, 2450t, 2451
- Metabolomics, 6, 436e-1
- Metaboreceptors, 47e-1
- Metachromatic leukodystrophy (MLD), 91e-2-3, 432e-4t, 2608
- Metagenome, 86e-11
- Metagenomics. *See also* Human microbiome
 definition of, 6, 86e-1t, 86e-2, 769t
 impact on clinical medicine, 86e-2
 in infectious disease diagnosis, 769t
 in pathogen discovery, 772-773, 773f
- Metagonimus yokogawai*, 245e-2t. *See also* Intestinal flukes
- Metal fume fever, 1692
- Metamorphopsia, 205, 206
- Metamyelocytes, 414f, 415
- Metanephrines, 2330t
- Metapneumovirus infections. *See* Human metapneumovirus (HMPV) infections
- Metastasis-suppressor genes, 102e-10
- Metastatic calcification, 426e-7, 426e-7t
- Metastatic disease, 102e-10
 in bladder cancer, 577
 to bone, 102e-10-11, 119e-4
 bone marrow examination in, 81e-7f
 to brain. *See* Brain tumors, metastatic characteristics of, 102e-1t, 102e-10-11, 102e-11f
 in colorectal cancer, 542
 to eye, 206, 207
 gene therapy for, 91e-4
 to heart, 270e-24, 271e-8f, 271e-8v, 289e-2
 to liver, 542, 553
 to lung, 246
 in lung cancer, 511, 511t
 lymphadenopathy in, 408
 to pericardium, 1788
 in renal cell carcinoma, 579
 to spine, 117, 605, 606, 606f
 splenomegaly in, 411
 surgery for, 103e-3
- Metastatic potential, 102e-10, 102e-11f
- Metaxalone, 473e-10t
- Metformin
 action of, 102e-13, 2413, 2414t
 adverse effects of, 644t, 1461, 1821, 2413, 2414t
 contraindications to, 2414t
 in elderly, 82
 for lifespan enhancement, 94e-7
 for metabolic syndrome, 2454
 for NALFD, 2057
 for prediabetes, 2406
 structure of, 94e-6f
 tumor-related effects of, 102e-13
 for type 2 diabetes mellitus, 2413, 2414t, 2416-2417, 2416f
- MET* gene mutations
 cancers associated with, 101e-2t
 in hereditary papillary renal carcinoma, 101e-5, 101e-6t
 in lung cancer, 507t
 in thyroid cancer, 2305t
- Methadone
 adverse effects of, 1467t
 in opioid use disorders, 468e-2, 468e-3
 for pain, 61, 91t, 94
 therapeutic monitoring of, 2765t
- Methamphetamine
 actions of, 465e-5t, 469e-3
 intracerebral hemorrhage due to, 2568, 2584
 overdose/poisoning with, 469e-3, 473e-6, 2765t
- Methane, 151e-2t, 151e-3f
- Methanogens, 86e-4
- Methanol poisoning
 acidosis in, 319
 clinical features of, 473e-12t
 laboratory evaluation of, 2765t
 mechanism of, 473e-12t
 optic neuropathy in, 203
 treatment of, 64e-9, 319, 473e-12t
- Methaqualone, 473e-11t
- Methemoglobin, 249, 636, 637, 2761t
- Methemoglobinemia, 636
- Methemoglobin inducers, 473e-11t
- Methicillin, 727t, 807, 961
- Methicillin-resistant *Staphylococcus aureus* (MRSA) infections
 in atopic dermatitis, 345
 in burn wounds, 166e-3
 cellulitis, 830
 community-acquired, 805, 807, 808t, 955, 956, 958
 control of, 918
 diagnosis of, 150e-8, 774
 emergence of, 918
 endocarditis, 822t, 823
 genomics of, 769t, 774, 776, 807
 health care-associated, 803t, 918
 meningitis. *See* Meningitis osteomyelitis, 840t
 prevalence of, 961
 risk factors for, 805
 sinusitis, 226
 treatment of, 345, 808t, 832t, 961-963, 962t
- Methimazole
 adverse effects of, 2296, 2348
 for Graves' disease, 2296
 for hyperthyroidism in pregnancy, 48
 for radioactive iodine exposure, 263e-7
 for thyroid storm, 2297
- Methionine aminopeptidase, 246e-9
- Methionine metabolism disorders, 434e-2t, 434e-3t
- Methionine synthase, 642f, 643

- Methocarbamol, 119, 473e-10t
Methotrexate
 action of, 103e-14, 641, 1960
 adverse effects of, 103e-11t, 103e-14, 1960
 anemia, 398
 common, 2146t
 cutaneous, 379, 380
 fetal, 124e-4
 folate deficiency, 647
 hepatic, 622, 2025
 in HIV infection, 1261
 late, 620t, 622
 myelopathy, 607t
 neurologic, 622
 prevention of, 2182
 pulmonary, 1797
 renal, 103e-11t, 103e-14
 seizures, 607t, 1793
 serious, 348t, 2130t, 2146t, 2182t
 for bladder cancer, 577
 for breast cancer, 530
 contraindications to, 1964
 drug interactions of, 42, 42t, 2130t
 asparaginase, 103e-11t
 NSAIDs, 103e-11t, 103e-14
 probenecid, 103e-14
 salicylates, 103e-14
 sulfonamides, 246e-4t
 trimethoprim, 246e-4t, 939t
 for ectopic pregnancy, 338
 in elderly patients, 2148–2149
 for gestational trophoblastic disease, 124e-5, 597
 for giant cell arteritis, 2189
 for granulomatosis with polyangiitis, 2185
 for head and neck cancer, 505
 for IBD, 1960
 for inflammatory myopathies, 2201
 for morphea, 376
 for MS, 2672
 for osteosarcoma, 119e-3
 for polymyalgia rheumatica, 2189
 for primary central nervous system lymphoma, 601–602
 for psoriasis, 348, 348t
 for psoriatic arthritis, 2177
 for reactive arthritis, 2175
 for rheumatoid arthritis, 2145, 2146t, 2148
 for sarcoidosis, 2211, 2212t
 for SLE, 2130t, 2133
 for systemic sclerosis, 2164, 2165
 for Takayasu's arteritis, 2190
 therapeutic monitoring of, 2146t, 2212t, 2765t
 for vasculitis, 2182
2-Methoxy estradiol, 102e-17f
8-Methoxypsoralen, 391, 678
Methyl aminolevulinate (MAL), 386
Methylation, of DNA, 101e-4
Methylation cycle, 642f
2-Methylbutyryl-glycinuria, 434e-3t
Methylcellulose, 1970t
6-Methylcoumarin, 389t
3-Methylcrotonyl glycinuria, 434e-3t
α-Methyl dopa
 adverse effects of, 253t, 727t, 2023, 2218t
 for carcinoid syndrome, 567
 for hypertension, 1624t
 for hypertension in pregnancy, 46
Methylene blue, 637, 656t
5,10-Methylenetetrahydrofolate (5-MTHF), 641, 641t
Methylenetetrahydrofolate reductase (MTHFR), 47, 641t, 642f, 643, 744
Methylfolate trap, 641
3-Methylglutaconic aciduria, 434e-3t
2-Methyl-3-hydroxybutyryl-CoA dehydrogenase deficiency, 434e-3t
Methylmalonate, serum, 647
Methylmalonic acidemia, 434e-2t, 434e-3t
Methylmalonic acidemia/aciduria, congenital, 645
Methylmalonyl CoA mutase deficiency, 645
Methylnaltrexone, 61, 92
7α-Methyl-19-nortestosterone, 2371
Methylphenidate
 for depression, 63
 for fatigue, 63, 2672
 for narcolepsy, 188t, 190
 for opioid-induced drowsiness, 61
Methylprednisolone
 adverse effects of, 2130t
 drug interactions of, 246e-3t
 for giant cell arteritis, 2189
 for gout, 2234
 for Graves' ophthalmopathy, 2297
 for IBD, 1960
 for interstitial lung disease with diffuse alveolar hemorrhage, 1715
 for MS, 2667, 2672
 for nausea and vomiting, 260t
 for optic neuritis, 202
 for PCP, 1361t
 for serum sickness, 667
 for SLE, 2130t, 2132
 for transplant rejection, 1828, 1829
 for vertigo, 151t
17-α-Methyl testosterone, 2370t, 2371
5-Methyltetrahydrofolate, 641, 641t
Methyl violet, 655
Methylxanthines, 324, 473e-8t
Methyprylon, 473e-11t
Methysergide
 adverse effects of, 1649, 2593, 2594t
 for carcinoid syndrome, 566
 for cluster headache prevention, 2595, 2595t
 for exertional headache, 2597
 for hypnic headache, 2598
 for migraine prevention, 2593, 2594t
 overdosage/poisoning with, 473e-8t
Metoclopramide
 adverse effects of, 260, 261, 2611
 for constipation, 61
 drug interactions of, 246e-2t
 for gastroparesis, 260, 260t, 2427
 for migraine, 2591, 2592t
 for nausea and vomiting, 62, 103e-25, 474
 parenteral, 2592t
Metoprolol
 for aortic dissection, 1641
 for arrhythmias, 1479t, 1480t
 dosage of, 1479t, 1480t
 drug interactions of, 246e-12
 genetic variations in response to, 39t, 40
 for heart failure, 1512t
 for hypertension, 1624t
 for ischemic heart disease, 1589t
 metabolism of, 34t
 for migraine prevention, 2594t
 for myocardial infarction, 1604
 overdosage/poisoning with, 473e-9t
 pharmacology of, 1479t, 1480t
Metronidazole, 246e-5t, 246e-10
Metronidazole
 for acne rosacea, 352
 action of, 931t, 932f, 933
 adverse effects of, 246e-5t, 353, 938t, 944, 2687t
 for amebiasis, 1367, 1367t
 for anaerobic bacterial infections, 1101, 1101t
 for bacterial vaginosis, 873t, 875
 for *C. difficile* infection, 860, 860t
 for clostridial infections, 991t, 994
 for *D. fragilis* infection, 1409
 drug interactions of, 246e-5t, 939t, 944
 for epidural abscess, 767t
 for febrile neutropenia, 103e-23
 for *Giardia* infections, 268, 1406
 in *H. pylori* eradication, 1042t, 1922, 1922t
 for hepatic encephalopathy, 2066
 indications for, 246e-5t, 941t, 944, 945t
 for intraabdominal infections, 767t
 for intracranial infections, 780t
 for Ludwig's angina, 233
 for meningitis, 887t, 889, 889t
 for necrotizing fasciitis, 832t
 for osteomyelitis, 841t
 for pelvic inflammatory disease, 878, 878t
 for peritonitis, 848
 for postpartum endomyometritis, 50
 for pouchitis in IBD, 1960
 in pregnancy and lactation, 246e-5t, 937t
 resistance to
 in anaerobic bacteria, 1101
 in *H. pylori*, 1042
 mechanisms of, 931t, 934
 prevalence of, 941t, 944
 for *T. vaginalis* infection, 871, 873t, 874, 1409
 for tetanus, 986
 for Vincent's angina, 233
METs (metabolic equivalent tasks), 1585t, 1587, 1587t
Metyrapone, 611–612, 2273, 2318
Mevalonate kinase, 372e-8t
Mevalonate kinase deficiency, 2215
Mexiletine
 adverse effects of, 1480t, 2682t
 for arrhythmias, 1480t, 1498
 dosage of, 1480t
 for hyperkalemic periodic paralysis, 462e-18
 metabolism of, 34t
 for myotonia, 462e-11
 for neuropathy, 2682t
 for pain, 95, 2672
 for paramyotonia congenita, 462e-18
 pharmacology of, 1480t
Meyer's loop, 199
Mezlocillin, 166e-3, 835
MFC (minimal fungicidal concentration), 150e-9
MGMT, 600
MGUS. *See* Monoclonal gammopathy of undetermined significance (MGUS)
MHC (major histocompatibility complex). *See* Human leukocyte antigen (HLA) complex
MHC class-restricted T cells, 372e-18, 373e-3, 373e-4
MHC-peptide complex, 373e-3
MI. *See* Myocardial infarction (MI)
Miasma, 761
Mibefradil, 328
MIC (minimal inhibitory concentration), 150e-9, 935
mic-2 gene, 119e-4
MIC-A5.1 allele, 2346
Micafungin, 233, 1331, 1344t, 1352t
MICA gene, 1240t, 1241
Mice. *See also* Rodent-borne viruses
 bite-wounds, 167e-2
 embryonic stem cells of, 454
 gnotobiotic, 86e-7. *See also* Gut microbiota
 knock-out/knock-in, 432
 in rickettsialpox. *See* Rickettsialpox
 transgenic, as models of genetic disease, 432
Micelles, 1935, 2076
Miconazole, 349, 873t, 875
Microaerophilic bacteria, 1094
Microalbuminuria
 in chronic kidney disease, 1813
 definition of, 1834t
 in diabetic nephropathy, 1844, 2425, 2425f
 in glomerular disease, 1834t
 in HIV infection, 1258
 in hypertension, 1616
Microaneurysms, retinal, 40e-3
Microangiopathic hemolytic anemia, 81e-4f, 282, 657, 657t, 726f. *See also* Thrombotic microangiopathy
Microarray technology
 in carcinoma of unknown primary, 120e-3
 definition of, 769t
 for gene expression in cancer, 101e-7, 101e-8f
 in infectious disease diagnosis, 769t
Microbial bioterrorism. *See* Bioterrorism
Microbial pathogenesis, **145e-1**
 avoidance of host defenses, 145e-4–7, 145e-4f, 145e-6f
 entry and adherence, 145e-1–3
 growth after entry, 145e-4
 host receptors, 145e-2t, 145e-3
 host response, 145e-10, 372e-23, 372e-26

- Microbial pathogenesis (*Cont.*):
 tissue damage and disease, 145e-9–10
 tissue invasion, 145e-7–8
 tissue tropism, 145e-8–9
 transmission to new hosts, 145e-10
 Microbial source tracking, 86e-1t
 Microbiome. *See also* Human microbiome
 analysis of, 145e-1, 145e-4
 definition of, 86e-1, 425
 in infectious disease evaluation, 761
 Microbiomics, 6
 Microbiota, 86e-1, 86e-1t. *See also* Gut microbiota
 Microcytosis, 393, 398
 Microfilaria, 245e-1, 245e-6t
 β_2 -Microglobulin
 in amyloidosis, 719, 720t, 724
 laboratory evaluation of, 406, 2758t
 in multiple myeloma, 715–716, 716t
 Microimmunofluorescence test, 1171
 Micronucleus assay, 263e-7
 Micronutrient malnutrition, **461**. *See also specific micronutrients*
 MicroRNAs (miRNAs)
 in aging, 94e-3
 in AML, 681
 in cancer, 101e-7
 gene expression silencing by, 102e-8
 in stem cells, 455
 in viruses, 214e-6
 Microsatellite instability, 101e-1f, 101e-5
 Microscopic polyangiitis, **2186**
 clinical features of, 1835t, 1840, 2186
 definition of, 2186
 diagnosis of, 1840, 2186
 incidence and prevalence of, 2186
 lung biopsy in, 386e-5f, 1840
 pathogenesis of, 2186
 treatment of, 2186
 Microsomal ethanol oxidizing system (MEOS), 2058
 Microsporidia/microsporidiosis, **1408**
 clinical features of, 1408
 diagnosis of, 245e-1, 245e-4t, 245e-5t, 245e-6t, 1408
 epidemiology of, 245e-4t
 in HIV infection, 1257, 1408
 life cycle of, 245e-4t, 1408, 1408f
 treatment of, 1408
 Microtubules, 103e-14
 Microvascular angina, 1579
 Mid-arm muscle circumference, 462
 MIDAS (Migraine Disability Assessment Score), 2590, 2591f
 Midazolam
 for conscious sedation, 35
 for delirium, 64t, 65
 dosage in liver disease, 37
 drug interactions of, 246e-3t
 for dyspnea, 62t
 for nerve agent exposure, 262e-7
 overdose/poisoning with, 473e-10t
 for scorpion stings, 2748
 for status epilepticus, 2558f
 during terminal weaning, 67
 Midbrain stroke syndrome, 2580f
 Middle cerebral artery
 lesion in territory of, 178
 occlusion of, 440e-3f, 2572–2573, 2573f, 2581
 stenosis of, 441e-16f
 territory of, 2572f, 2573f
 Middle East respiratory syndrome (MERS), 261e-9, 1204–1205
 Middleton maneuver, 411
 Midline destructive disease, 2184
 Midline granuloma, 237
 Midodrine
 adverse effects of, 2644
 for ascites, 288
 for dialysis-related hypotension, 1824
 for hepatorenal syndrome, 1810
 for orthostatic hypotension, 146, 2644
 for postural orthostatic tachycardia syndrome, 1477
 Midpontine stroke syndrome, 2578f
 Mifepristone
 adverse effects of, 2273
 for Cushing's syndrome, 2273
 for postcoital contraception, 2391
 MIG (monokine induced by IFN- γ), 372e-12t
 Miglitol, 2414t
 Migraine, **2586**
 acephalgic, 2560, 2589
 autonomic symptoms in, 107
 channelopathies in, 444e-2t
 chronic, 2589
 classification of, 2587t
 clinical features of, 2588t, 2589
 cyclic vomiting in, 259
 diagnosis of, 2589, 2590t
 familial hemiplegic, 2589
 fatigue in, 152
 genetic considerations in, 444e-2t, 2589
 imaging in, 441e-43f, 2589, 2589f, 2590f
 ocular manifestations of, 204
 pathogenesis of, 89, 2586, 2588–2589, 2588f
 in pregnancy, 49
 prevalence of, 107t, 2586
 prevention of, 2593, 2593t, 2594t
 Raynaud's phenomenon and, 1649
 retinal, 201
 vs. stroke, 2560
 taste disorders in, 215
 treatment of, **2590**
 acute attack therapies, 2590, 2592t, 2593t
 5-HT₁ agonists, 2590–2591, 2592t
 disability assessment in, 2590, 2591f
 dopamine antagonists, 2591–2592, 2592t
 drug combinations for, 2592–2593
 nonpharmacologic, 2590
 NSAIDs, 2590, 2592t
 opioids, 2592t, 2593
 triggers, 2586
 vertigo in, 148
 vestibular, 150
 Migraine aura, 2589
 Migraine Disability Assessment Score (MIDAS), 2590, 2591f
 Migrainous neuralgia, 237
 Migrating motor complex, 264–265
 Mikulicz's disease, 391e-1. *See also* IgG4-related disease (IgG4-RD)
 Mild cognitive impairment (MCI), 75, 76f, 171, 175f, 2598
 Miliaria rubra, 479e-1
 Miliary tuberculosis, 138, 1108f, 1112, 1119
 Military recruits, adenovirus infections in, 1208
 Milk-alkali syndrome, 313, 2458, 2479
 Milk of calcium bile, 2082
 Milk of magnesia, 62t
 Millard-Gubler syndrome, 209
 Miller Fisher syndrome, 989, 2695, 2696t, 2698t.
See also Guillain-Barré syndrome (GBS)
 Miller's lung, 1681t
 Millipede dermatitis, 2750
 Milnacipran, 2240, 2240t
 Milrinone, 1509–1510, 1510t, 1763
 Milroy's disease (congenital lymphedema), 1653
 Miltefosine
 action of, 246e-10–11
 adverse effects of, 246e-5t, 1391
 for cutaneous leishmaniasis, 1393
 for free-living ameba infections, 1367, 1368
 pharmacology of, 246e-10–11
 for visceral leishmaniasis, 1391
 Milwaukee shoulder, 2236
 Mind-body interventions, 14e-3
 Mineral(s), **96e-9**. *See also specific minerals*
 absorption of, 1933
 assessment of nutritional status, 462
 body stores of, 96e-1
 deficiencies of, 96e-9t
 parental supplementation of, 98e-6t
 recommended intake of, 95e-3t
 reference ranges for, 2766t
 in specialized nutritional support formulas, 98e-5
 tolerable upper intake levels of, 96e-9t
 toxicities of, 96e-9t
 Mineralocorticoid(s)
 for adrenal insufficiency, 2326
 resistance to, 297
 synthesis of, 2309f, 2311, 2311f
 Mineralocorticoid excess, **2318**. *See also* Syndrome of apparent mineralocorticoid excess (SAME)
 clinical features of, 2319
 diagnosis of, 2319, 2320f
 differential diagnosis of, 2319, 2320f
 epidemiology of, 2318
 etiology of, 2318–2319, 2318t, 2320t
 pathophysiology of, 64e-8, 64e-8f
 treatment of, 2319–2320, 2320f
 Mineralocorticoid receptors, 297, 306, 310
 Mineral oil, vitamin A absorption and, 96e-6
 Minimal bactericidal concentration (MBC), 150e-9
 Minimal change disease
 clinical features of, 1835t, 1841–1842
 pathogenesis of, 1841–1842, 1842f
 proteinuria in, 293, 1805, 1834, 1842
 renal biopsy in, 62e-1f, 1841–1842
 treatment of, 1842
 Minimal erythema dose, 389
 Minimal fungicidal concentration (MFC), 150e-9
 Minimal inhibitory concentration (MIC), 150e-9, 935
 Minimally conscious state, 1771
 Minimally invasive adenocarcinoma, 515–516
 Mini-Mental Status Examination (MMSE)
 in delirium evaluation, 64, 168
 in dementia evaluation, 173
 in neurologic examination, 2537
 Mini Nutritional Assessment, 81, 276
 Minitransplant, 103e-20, 636, 703
 Minocycline
 for acne rosacea, 352
 for acne vulgaris, 352
 for actinomycosis, 1091t
 action of, 930, 931t, 943
 adverse effects of, 943
 cutaneous, 379
 fever, 138
 hyperpigmentation, 360, 361
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 oral lesions, 240t
 vasculitis, 363, 2218t
 indications for, 940t, 943
 for *Legionella* infections, 1020t
 for leprosy, 1126, 1127
 for MRSA infections, 345, 962t
 for *Nocardia* infections, 1087
 for osteomyelitis, 841t
 resistance to, 931t, 940t
 for sarcoidosis, 2211
 for staphylococcal skin/soft tissue infections, 962t
 for Whipple's disease, 1094
 Minor histocompatibility antigens, 373e-6
 Minor histocompatibility loci, 373e-6
 Minoxidil
 adverse effects of, 253t, 332, 380, 1625
 for androgenetic alopecia, 355t
 development of, 43
 for hypertension, 1624t
 Miosis, 473e-3, 1774
 MIP-1 α (macrophage inflammatory protein-1 α), 372e-12t, 1241
 MIP-1 β (macrophage inflammatory protein-1 β), 372e-12t
 Mipomersen, 2448t, 2449
 Mirizzi's syndrome, 2081
 miRNAs. *See* MicroRNAs (miRNAs)
 Mirror neuron systems, 444e-12
 Mirtazapine
 adverse effects of, 2392, 2710t, 2717
 for depression, 63–64, 2710t, 2717

- for insomnia, 65
- for involuntary weight loss, 276
- for progressive multifocal leukoencephalopathy, 900
- Misonidazole, 2687t
- Misoprostol, 49, 1920t, 1921, 1924t
- Misense mutation, 432, 433f
- Mistake (error), 12e-1
- Mites
 - biting, 2745
 - in rickettsialpox. *See* Rickettsialpox
- MITF, 359, 386, 431t, 494
- Mitiglinide, 2413, 2414t
- Mitochondria, 85e-1
 - in apoptosis, 372e-26f, 444e-5f, 444e-9
 - dysfunction, in neurodegenerative disease, 444e-9
 - in energy production, 462e-15
 - functions of, 85e-1, 85e-1t
 - genetic defects of, 1555t
- Mitochondrial ataxias, 2630
- Mitochondrial cytochrome c, 372e-23
- Mitochondrial DNA (mtDNA), 85e-1. *See also* Mitochondrial DNA diseases
 - aging and, 85, 85e-8, 85e-8f, 94e-3, 94e-5
 - functions of, 85e-1-3, 439
 - heteroplasmy in, 85e-2, 85e-3f
 - homoplasmic variants of, 85e-3-4
 - inheritance of, 85e-1, 85e-2f, 85e-3f, 462e-15
 - mitotic segregation of, 85e-2
 - nuclear cross-talk perturbations with, 85e-7, 85e-7f
 - polyploidy in, 85e-2-3
 - replication of, 85e-1
 - structure of, 85e-1-3, 439
- Mitochondrial DNA diseases, 85e-1
 - age-associated, 85e-8, 85e-8f
 - cardiomyopathy in, 1554, 1555t
 - clinical features of, 85e-4-6, 85e-5t, 220, 2602
 - depletion syndromes, 462e-16
 - diagnosis of, 85e-6-7, 85e-7f
 - examples of, 85e-4f, 85e-5-7, 85e-6t
 - genetic control of, 85e-2f
 - genetic counseling in, 85e-9
 - genetic mutations causing, 85e-1, 85e-3-4, 85e-4f, 85e-6-7
 - heteroplasmy in, 85e-3-4, 85e-3f
 - homoplasmic sequence variation in, 85e-2-3, 85e-7-8
 - inheritance of, 85e-1, 85e-2f, 438f, 439
 - with large-scale DNA rearrangements, 85e-6t
 - organ manifestations of, 85e-2f
 - pathologic features of, 85e-3-4, 85e-5f
 - phenotypic heterogeneity of, 85e-4
 - with point mutations, 85e-6t
 - prevention of, 85e-9, 85e-10f
 - renal, 1855
 - sporadic, 85e-9
 - treatment of, 85e-9
- Mitochondrial myopathy(ies), 462e-15
 - drug-induced, 462e-20
 - Kearns-Sayre syndrome. *See* Kearns-Sayre syndrome
 - pathogenesis of, 462e-15
 - progressive external ophthalmoplegia. *See* Progressive external ophthalmoplegia
 - pure myopathy syndromes, 462e-16
 - skeletal muscle-CNS syndromes
 - MELAS syndrome. *See* MELAS (mitochondrial myopathy, encephalopathy, lactic acidosis, and strokelike episodes) syndrome
 - MERRF syndrome. *See* MERRF (myoclonic epilepsy with ragged red fibers) syndrome
- Mitomycin C
 - adverse effects of, 621, 729, 1796, 1797, 1864
 - for bladder cancer, 576
 - for hepatocellular carcinoma, 550
- Mitosis, 427
- Mitotane
 - for adrenocortical carcinoma, 2322-2323
 - adverse effects of, 2273, 2323
 - in cancer chemotherapy, 103e-16
 - for Cushing's syndrome, 2273, 2318
 - for ectopic ACTH syndrome, 611-612
- Mitotic segregation, 85e-2
- Mitotic spindle inhibitors, 103e-14-15
- Mitoxantrone
 - action of, 103e-13, 2670
 - adverse effects of, 103e-10t, 103e-13, 2670
 - for AML, 683
 - drug interactions of, 103e-10t
 - for MS, 2669t, 2670
 - for prostate cancer, 586
- Mitral regurgitation, 1543
 - cardiogenic shock in, 1762
 - clinical features of, 1545
 - diagnosis of
 - chest radiograph, 1545
 - CMR, 270e-19
 - ECG, 269e-9f, 1545
 - echocardiography, 270e-18-19, 270e-18f, 270e-19f, 1545, 2154v
 - hemodynamic patterns, 1762t
 - in dilated cardiomyopathy, 1556
 - dyspnea in, 47e-2
 - etiology of, 1539t, 1543
 - in heart failure, 1515
 - heart sounds/murmur in, 51e-1-5, 51e-1f, 51e-2f, 51e-5f, 1448, 1448f, 1545
 - hemoptysis in, 246
 - in hypertrophic cardiomyopathy, 1569
 - left atrial compliance in, 1543-1544
 - with mitral stenosis, 1552
 - pathophysiology of, 1543-1544
 - in pregnancy, 46-47
 - radiation-associated, 621
 - in rheumatic heart disease, 2151, 2154v
 - in rheumatoid arthritis, 2138
 - treatment of
 - medical, 1544
 - strategy for, 1544f
 - surgical, 1544-1545
 - transcatheter valve repair, 296e-4, 1546, 1546f
- Mitral stenosis, 1539
 - associated lesions, 1540, 1551
 - clinical features of, 1539-1540
 - diagnosis of
 - cardiac catheterization, 1541
 - chest radiograph, 1540
 - ECG, 269e-8f, 269e-9f, 1540
 - echocardiography, 270e-19, 270e-19f, 1540, 2154v
 - hemodynamic measurements, 1462, 1462f, 1539
 - valve area, 1463
 - differential diagnosis of, 1540-1541
 - etiology of, 1539, 1539t
 - heart sounds/murmur in, 51e-1, 51e-1f, 51e-6, 51e-6f, 1448f, 1449, 1540
 - with mitral regurgitation, 1552
 - pathology of, 1539
 - pathophysiology of, 1539
 - in pregnancy, 46, 1442
 - in rheumatic heart disease, 51e-6, 1539, 2151, 2154v
 - treatment of
 - medical therapy, 1541-1542
 - strategy for, 1541f
 - transcatheter, 296e-4
 - valve replacement, 1542-1543, 1543t
 - valvotomy, 1542-1543, 1542f
 - valvuloplasty, 296e-4, 1542, 1542f
- Mitral valve prolapse, 1546
 - clinical features of, 254, 1443, 1546
 - ECG in, 1547-1548
 - echocardiography in, 270e-16
 - etiology of, 1546
 - heart sounds/murmur in, 51e-2, 51e-3f, 51e-7, 1449, 1449f, 1546
 - imaging in, 1547-1548
 - pathophysiology of, 1546
 - in polycystic kidney disease, 1852
 - stroke prevention in, 2566t
 - treatment of, 1547
- Mitral valve repair, 1544, 1546, 1546f, 1552
- Mitral valve replacement, 1545, 1552
- Mittelschmerz, 338, 2379
- Mivazerol, 54
- Mixed connective tissue disease (MCTD)
 - AV conduction block in, 1471, 1471t
 - clinical features of, 2165
 - diagnosis of, 2165
 - lymphadenopathy in, 408
 - neuropathy in, 2683
 - SA node dysfunction in, 1467
- Mixed gonadal dysgenesis (45,X/46,XY mosaicism), 2351t, 2352-2353
- Mixed lineage leukemia (MLL) translocations, 643
- Mixed mania, 2717
- Mixed venous oxygen saturation, 1733-1734
- Mixing studies, in coagulation evaluation, 406
- Miyoshi's myopathy, 462e-12, 462e-13t
- MK3475, 521
- MLD (metachromatic leukodystrophy), 91e-2-3, 432e-4t, 2608
- MLH1 gene mutations, 101e-6t, 102e-7, 592, 596
- MLH6 gene mutations, 592
- MLL gene mutations, 699
- MLL (mixed lineage leukemia) translocations, 643
- MM12 gene mutations, 1703
- M-mode echocardiography, 270e-1, 270e-1f
- MMPs. *See* Matrix metalloproteinases (MMPs)
- MMR. *See* Measles-mumps-rubella vaccine (MMR)
- MMSE. *See* Mini-Mental Status Examination (MMSE)
- MNI overexpression, 680, 680t
- MNSsU blood group system, 138e-1t, 138e-2
- Mobile genetic element, 769t
- Mobility Questionnaire, 78t
- Modafinil
 - for depression, 63
 - for fatigue, 63, 153, 2672
 - for narcolepsy, 188t, 190
 - for opioid-induced drowsiness, 61
 - for shift-work disorder, 194
- Model for End-Stage Liver Disease (MELD) score, 1995, 1996, 2069, 2069t
- Modification of Diet in Renal Disease (MDRD) equation, 480e-4, 480e-4f, 1813t
- Modular scale-free topology, 87e-2
- MODY (maturity-onset diabetes of the young), 431t, 441t, 2400, 2406
- Moffett's position, 216
- MOG (myelin oligodendrocyte glycoprotein), 372e-24t, 444e-4f
- Mogamulizumab, 225e-5
- Mohs micrographic surgery, 501-502
- Mokola virus, 1304
- Molar pregnancy. *See* Gestational trophoblastic disease
- Molds, 1329
- Mole, atypical, 493
- Molecular mimicry, 377e-1, 444e-6-7
- Molecular phylogenetics, 86e-2
- Molindone, 2722t
- Mollaret's meningitis, 909t, 1179
- Molluscum contagiosum, 220e-1
 - diagnosis of, 220e-1
 - epidemiology of, 220e-1, 220e-1t
 - in HIV infection, 220e-1, 1263
 - persistent infections, mechanism of, 214e-7
 - skin manifestations of, 76e-9f, 220e-1, 220e-1f
 - treatment of, 220e-1
- Molly (MDMA), 168, 301, 469e-3
- Molybdenum
 - deficiency of, 96e-9t, 96e-10, 533
 - functions of, 96e-10
 - parental supplementation of, 98e-6t
 - recommended intake of, 95e-3t
 - tolerable upper intake level of, 96e-9t
 - toxicity of, 96e-9t, 263e-3t
 - "Monday chest tightness," 1691
- Monge's disease (chronic mountain sickness), 248, 476e-5
- Monkey, bite-wounds from, 167e-1, 167e-3t
- Monkeypox, 220e-1-2, 220e-1t

- Monoamine oxidase inhibitors (MAOIs)
 action of, 465e-2
 adverse effects of, 1617t, 2392, 2710t, 2717
 for depression, 2710t, 2717
 drug interactions of, 246e-4t, 246e-9, 2711t, 2717
 overdose/poisoning with, 473e-8t, 473e-13t
 for panic disorder, 2709
- Monoamine oxidase type B (MAO-B) inhibitors, 2615, 2616t
- Monobactams. *See* Aztreonam
- Monoblasts, 421
- Monoclonal cryoglobulinemia, 368
- Monoclonal gammopathy, malignant, 618–619, 712. *See also* B-cell chronic lymphoid leukemia (CLL)/small lymphocytic lymphoma; Multiple myeloma; Waldenström's macroglobulinemia
- Monoclonal gammopathy of undetermined significance (MGUS)
 amyloidosis in, 721
 diagnosis of, 714–715, 715t
 features of, 103e-2
 in HIV infection, 1262
 polyneuropathy in, 2696t, 2699, 2700
 tumor markers in, 473t
 von Willebrand disease in, 731
- Monoclonal immunoglobulin, 473t
- Monocyte(s), 421
 disorders of, 419t, 421–422
 hematopoietic differentiation of, 89e-3f
 in HIV infection, 1238
 in peripheral blood smear, 81e-2, 81e-6f
 in tuberculosis, 1107
- Monocyte chemotactic protein-1 (MCP-1), 372e-11t
- Monocyte chemotactic protein-2 (MCP-2), 372e-11t
- Monocyte chemotactic protein-3 (MCP-3), 372e-12t
- Monocyte chemotactic protein-4 (MCP-4), 372e-12t
- Monocyte chemotactic proteins (MCPs), 1242
- Monocytoid B-cell lymphoma, 707
- Monocytosis, 421
- Monogenic disorders. *See* Mendelian disorders
- Monokine induced by IFN- γ (MIG), 372e-12t
- Mononeuropathies, 463e-4–5, 463e-5f, 1124, 2691–2692
- Monongahela virus, 1308t
- Monoparesis, 157
- Monopotent cells, 454
- Monosodium glutamate, 216
- Montelukast
 adverse effects of, 2218t
 for allergic rhinitis, 2122, 2123f, 2124
 for angioedema, 2119
 for asthma, 1678
 for urticaria, 2119
- Mood disorders, 2714
 alcohol-induced, 2725
 bipolar disorder. *See* Bipolar disorder
 depression. *See* Depression/depressive disorders
 in fibromyalgia, 2239
 in the homeless, 2723
 neuropathology of, 465e-3–4, 465e-4f
- Mood stabilizers, 2718t. *See also* specific drugs
- Moral distress, 17e-2–3
- Moral hazard, 15e-2–3
- Moraxella catarrhalis*, 150e-2t, 1013, 1014
- Moraxella catarrhalis* infections
 clinical features of, 1013
 hemoptysis, 245
 laryngitis, 234
 mastoiditis, 231
 otitis media, 229–230, 1013
 pneumonia, 1013
 sinusitis, 226, 1013
 in COPD, 1013, 1014f
 diagnosis of, 1013–1014
 epidemiology of, 1013
 pathogenesis of, 1013
 treatment of, 230, 1014
- Morbilloform eruptions. *See also* Rash
 definition of, 340t
 drug-induced, 377–378, 380–381, 381f
 etiology of, 362, 362t
- Morganella* infections, 862, 1035
- Morgellons syndrome, 2750–2751
- Moricizine, 1479t
- Morphea, 355, 355t, 376
- Morphine
 adverse effects of, 1603, 1741, 2640t
 for autonomic storm, 2643
 dosage in liver disease, 37
 for dyspnea, 47e-4, 62t
 in mechanical ventilation, 1741
 for myocardial infarction, 1597t, 1603
 for pain, 60, 91t, 92
 for pulmonary edema, 1761f, 1763
 routes of administration of, 92
 for sickle cell painful crisis, 635
 sustained-release, 91t, 94
 during terminal weaning, 67
 therapeutic monitoring of, 2765t
- Morquio syndromes, 432e-2t
- Mortality. *See* Death
- Mortality rates, 93e-2. *See also* specific diseases
- Morumbi virus, 1309t
- Morvan's syndrome, 617, 619
- Mosaicism, 432, 439
- Mosquito(s)
Aedes. *See Aedes* mosquitos
 in arboviral encephalitis, 1314–1317
 bite, 2749–2750
Culex. *See Culex* mosquitos
 in malaria transmission, 1368–1370, 1369f, 1371
- Mother's curse, in aging, 94e-3
- Motherworth, 404t
- Motilin agonists, 260t
- Motion sickness, 62, 259, 260, 260t
- Motor cortex, 176f
- Motor efferents, 47e-1
- Motor evoked potentials, 442e-3–4
- Motor examination, 2538
- Motor neuron disorders. *See also* Amyotrophic lateral sclerosis (ALS)
 differential diagnosis of, 2633
 etiology of, 2631t
 genetic, 2634–2635t
 lower, 2632t, 2634t, 2635–2636
 sporadic, 2632t
 upper, 2632t, 2634–2635t, 2636
- Motor neurons, age-related changes in, 76
- Motor neuropathy, 618, 2699
- Motor tic, 2623
- Mounier-Kuhn syndrome, 1695
- Mountain sickness, chronic, 248, 476e-5
- Mouth
 dry. *See* Xerostomia
 healthy, 46e-3f
- Mouth-to-mouth resuscitation, 1769
- Movement disorders
 choreas. *See* Chorea(s)
 drug-induced, 2624
 dystonia. *See* Dystonia
 Huntington's disease. *See* Huntington's disease
 hyperkinetic. *See* Hyperkinetic movement disorders
 myoclonus. *See* Myoclonus
 in neurodegeneration with brain iron accumulation, 2625–2626
 in Parkinson's disease. *See* Parkinson's disease
 in pregnancy, 49
 psychogenic, 2626
 restless legs syndrome. *See* Restless legs syndrome
 Tourette's syndrome, 2623
 in Wilson's disease. *See* Wilson's disease
- Moxifloxacin
 action of, 931t, 932–933
 adverse effects of, 943
 for anaerobic bacterial infections, 1101
 for community-acquired pneumonia, 767t, 808t
 for febrile neutropenia, 103e-23
- for health care-associated pneumonia, 812t
 indications for, 941t, 943
 for *Legionella* infections, 1020t
 for leprosy, 1126
 for lung abscess, 815
 for mucopurulent cervicitis, 876
 for multidrug-resistant TB, 205e-6
 for pelvic inflammatory disease, 878
 for plague, 1075
 for pneumonia, 953
 resistance to, 931t, 941t, 1101
 for sinusitis, 227t
- Moxonidine, 1513
- Moyamoya disease, 441e-41f, 2568, 2584
- 6-MP. *See* 6-Mercaptopurine (6-MP)
- MPN-U (myeloproliferative neoplasm, unclassifiable), 135e-6–7
- M protein, 964
- MPTP, 444e-3, 2611
- MRA. *See* Magnetic resonance angiography (MRA)
- MRCP. *See* Magnetic resonance cholangiopancreatography (MRCP)
- MRE11* gene, 375e-1, 2109
- MRF (myelin gene regulatory factor), 444e-5
- MRI. *See* Magnetic resonance imaging (MRI)
- MRP2 (multidrug resistance-associated protein 2), 2000
- MRP3 (multidrug resistance-associated protein 3), 2000
- MRSA infections. *See* Methicillin-resistant *Staphylococcus aureus* (MRSA) infections
- MS. *See* Multiple sclerosis (MS)
- MSA. *See* Multiple system atrophy (MSA)
- MSCRAMMs, 145e-7, 817, 955–956, 972
- MSH. *See* α -Melanocyte-stimulating hormone (MSH)
- MSH2* gene mutations, 101e-6t, 538t, 592, 596
- MSH6* gene mutations, 101e-6t
- MSM. *See* Men who have sex with men (MSM)
- MSMD (Mendelian susceptibility to mycobacterial disease), 2106
- MTC. *See* Medullary thyroid carcinoma (MTC)
- MTD (maximum-tolerated dose), 103e-5
- mtDNA. *See* Mitochondrial DNA (mtDNA)
- 5-MTHF (5,10-methylenetetrahydrofolate), 641, 641t
- MTHFR (methylenetetrahydrofolate reductase), 47, 641t, 642f, 643, 744
- mTOR, 94e-5, 103e-16f, 562, 1701
- mTOR inhibitors, 102e-3, 103e-19, 119e-4, 562, 1853. *See also* Everolimus; Temsirolimus
- Mucambo virus, 1306t
- Muckle-Wells syndrome (MWS)
 amyloidosis in, 1845
 clinical features of, 372e-8t, 2213t, 2215
 genetic considerations in, 2213t
 pathogenesis of, 372e-8t
 treatment of, 2213t
- Mucociliary elevator, 949
- Mucopolidoses, 432e-3t
- Mucopolysaccharidoses, 432e-2t, 432e-5–6, 436e-5f, 1565
- Mucorales, 1350, 1351. *See also* Mucormycosis
- Mucormycosis, 1350
 clinical features of, 135, 226, 784, 1350–1351
 diagnosis of, 1351, 1351f
 differential diagnosis of, 1351–1352
 epidemiology of, 1350
 etiology of, 1350, 1350t
 in HIV infection, 1250
 in immunocompromised patient, 76e-12f, 489, 489t
 pathogenesis of, 1350
 in transplant recipient, 925, 926
 treatment of, 1352–1353, 1352t
- Mucosa-associated lymphoid tissue (MALT), 372e-23, 372e-25f
- Mucosa-associated lymphoid tissue (MALT) lymphoma, 703
 clinical features of, 535–536, 702t, 703
 diagnosis of, 703

- genetic considerations in, 699t, 703
H. pylori and, 536, 697, 703, 1038
 in thymus, 123e-1
 treatment of, 536, 703–704
- Mucosal bleeding, 404
- Mucosal malabsorption, 269
- Mucositis
 acute necrotizing ulcerative (noma), 236, 487, 1097
 chemotherapy-related, 103e-25
 radiation therapy-related, 505
 in transplant recipient, 139e-3
- Mucous retention cyst, oral, 240t
- Mu heavy chain disease, 719
- Muir-Torre syndrome, 365
- Muju virus, 1307t
- Mulberry molars, 236, 1137
- Muleshoe virus, 1308t
- Müllerian agenesis, 335, 336, 2357
- Müllerian epithelium, 592
- Müllerian inhibiting substance. *See* Antimüllerian hormone (AMH)
- Multicentric reticulohistiocytosis, 365
- Multidetector computed tomography (MDCT), 440e-1, 1664–1666. *See also* Computed tomography (CT)
- Multidrug resistance-associated protein 2 (MRP2), 2000
- Multidrug resistance-associated protein 3 (MRP3), 2000
- Multidrug resistance (MDR) phenotype, 102e-10
- Multifactorial disorders, 425, 440
- Multifactorial inheritance, 446
- Multifocal atrial tachycardia, 278e-5f, 1485, 1486f
- Multifocal motor neuropathy, 2696t, 2699
- Multifocal motor neuropathy with conduction block, 2633, 2636
- Multifocal visual evoked potentials, 442e-3
- Multi-infarct dementia, 2602
- Multikinase inhibitors, 102e-3–4, 102e-4f, 103e-18t, 103e-19
- Multilineage-differentiating stress-enduring cells, 453t
- Multilocus sequence typing, 769t
- Multinodular goiter, 2301–2302
- Multiorgan system failure, 1733
- Multiple carboxylase/biotinidase deficiency, 434e-3t
- Multiple endocrine and other organ neoplasias (MEONs), 2335
 Carney's syndrome. *See* Carney's syndrome/complex
 Cowden's syndrome. *See* Cowden's syndrome (multiple hamartoma syndrome)
 McCune-Albright syndrome. *See* McCune-Albright syndrome
 neurofibromatosis type 1. *See* Neurofibromatosis type 1 (NF1)
 neurofibromatosis type 2. *See* Neurofibromatosis type 2 (NF2)
 von Hippel-Lindau disease. *See* von Hippel-Lindau disease/syndrome
- Multiple endocrine neoplasia type 1 (MEN 1; Wermer's syndrome), 2335
 clinical features of, 2335, 2336t
 adrenocortical tumors, 2339–2340
 angiofibromas, 364, 2340
 collagenomas, 2340
 gastrinoma, 1927–1928
 glucagonoma, 2338
 hyperparathyroidism, 2470
 insulinomas, 2337–2338
 lipomas, 2340
 meningiomas, 2340
 NETs, 562, 562t, 2339
 neuropathy, 2685–2686
 parathyroid tumors, 2335–2336, 2470
 pituitary tumors, 2265–2266, 2265t, 2339
 pNETs, 571, 2336, 2338–2339, 2338f
 thyroid tumors, 2340
 VIPomas, 2338
 ZES, 562, 568, 2336–2337
 epidemiology of, 2335
- genetic considerations in, 2340
 chromosomes involved, 101e-6t, 562t, 2340, 2471, 2471f
MEN1/Menin gene mutations, 599t, 2265–2266, 2265t
 variable expressivity, 437
 genetic testing in, 2340
 pathogenesis of, 2251
 screening tests in, 2337t, 2340
- Multiple endocrine neoplasia type 2 (MEN 2)
 clinical features of, 2340
 early interventions for, 450t
 genetic considerations in, 101e-5, 2332, 2341t, 2342
 genetic testing for, 449, 450t, 2342
 medullary thyroid carcinoma in, 2332, 2332f, 2336t, 2340–2342. *See also* Medullary thyroid carcinoma (MTC)
 medullary thyroid carcinoma only, 2307, 2336t, 2340–2342
 parathyroid tumors in, 2342
 pathogenesis of, 2251–2252
 pheochromocytoma in, 2332, 2332f, 2342
 screening tests in, 2341t, 2342
- type 2A
 clinical features of, 365, 1619, 1620t, 2332, 2336t, 2340
 genetic considerations in, 101e-6t, 2336t
 hyperparathyroidism, 2470
- type 2B (MEN 3)
 clinical features of, 436e-3f, 1619, 1620t, 2332, 2336t, 2340, 2470
 genetic considerations in, 2336t
- Multiple endocrine neoplasia type 4 (MEN 4), 2265t, 2336t, 2342
- Multiple exostoses (diaphyseal aclasis), 426e-7
- Multiple hamartoma syndrome. *See* Cowden's syndrome (multiple hamartoma syndrome)
- Multiple/mixed valvular heart disease
 clinical features of, 1551–1552
 diagnosis of, 1552
 pathophysiology of, 1550–1551
 treatment of, 1552
- Multiple myeloma, 712
 clinical features of, 713–714
 amyloidosis, 721
 bone lesions/pain, 102e-11, 119e-4, 713, 714f, 717
 cardiac, 271e-6f, 271e-6v
 cryoglobulinemia, 386e-5f
 erythropoietin deficiency, 398
 hypercalcemia, 610, 714
 hypogammaglobulinemia, 486t, 492, 714
 infections, 76e-12f, 484t, 486t, 713–714
 neuropathy, 2685–2686, 2699–2700
 paraneoplastic syndromes, 610, 618–619
 proteinuria, 293
 renal
 contrast nephropathy, 1803, 1806t
 light chain cast nephropathy, 62e-7f, 1805, 1859, 1859f
 tubular damage, 714
 diagnosis of, 81e-11f, 712f, 713f, 714–715, 715t, 2494
 epidemiology of, 712–713
 epigenetic changes in, 102e-7
 etiology of, 712
 genetic considerations in, 101e-2t, 712, 716
 global considerations in, 713
 immunology of, 698f
 M component in, 715
 nonsecretory, 715t
 pathogenesis of, 102e-11, 713, 713f
 prognosis of, 715–716, 716t
 renal biopsy in, 62e-7f
 as second malignancy in breast cancer, 529
 smoldering, 715t
 staging of, 715–716, 716t
 treatment of
 bortezomib, 102e-5t, 102e-6, 716–717
 carfilzomib, 102e-5t
 chemotherapy, 716–717
 denosumab, 102e-11
- HSCT, 139e-5t, 139e-6, 717
 lenalidomide, 716–717
 relapsed disease, 717
 supportive care, 717
 thalidomide, 716–717
 tumor markers in, 473t
 von Willebrand disease in, 731
- Multiple sclerosis (MS), 2661
 clinical features of, 41e-1v, 2663–2664, 2663t
 constipation, 274
 cranial nerve palsies, 2664
 erectile dysfunction, 326
 fatigue, 152
 frontal lobe syndrome, 184
 gait disorders, 163
 hearing loss, 219
 myelitis, 2656
 myelopathy, 2659
 ocular, 200, 202, 210, 2663
 olfactory dysfunction, 214
 paroxysmal, 2664, 2672
 weakness, 156, 2663
 clinical variants of, 2673
 course of, 2664, 2664f
 diagnosis of, 2664–2665, 2665t
 differential diagnosis of, 2665, 2667t
 epidemiology of, 2661–2662
 genetic considerations in, 373e-8t, 444e-1, 444e-2, 2662
 HLA gene association with, 373e-5, 373e-8t
 imaging in, 441e-27–28f, 441e-35f, 2664–2665, 2666f
 immunopathology of
 autoimmunity, 372e-24t, 2662
 autoreactive T cells, 2662
 gut microbiota, 86e-8
 inflammasome activity, 372e-8t
 inflammatory mediators, 372e-29t
 Marburg's variant, 2673
 neurodegeneration in, 2662–2663
 pathogenesis of, 444e-4f, 2661–2663, 2661f
 pregnancy and, 49, 2666–2667
 primary progressive, 2664, 2664f
 prognosis of, 2665–2666
 progressive/relapsing, 2664, 2664f
 relapsing/remitting, 2664, 2664f
 vs. sarcoidosis, 2208
 scoring systems for, 2667, 2668t
 secondary progressive, 2664, 2664f
 treatment of, 2667
 for acute attacks, 2667
 alemtuzumab, 2669
 dietary, 96e-7, 2672
 dimethyl fumarate, 2669
 experimental therapies, 444e-5, 2672–2673
 fingolimod, 2669–2670
 glatiramer acetate, 2668–2669
 glucocorticoids, 2667
 INF- β , 2667–2668
 for initial demyelinating episodes, 2667
 initiation and changing of, 2670–2671, 2671f
 mitoxantrone hydrochloride, 2669
 natalizumab, 2669
 off-label options in, 2672
 outcomes of, 2669t
 for progressive forms, 2671–2672, 2671f
 for relapsing forms, 2667–2671, 2671f
 symptomatic, 2672
 teriflunomide, 2669
 vitamin D supplements, 2671
 triggers of, 2662
- Multiple sleep latency test, 188
- Multiple sulfatase deficiency, 432e-4t
- Multiple system atrophy (MSA)
 clinical features of, 2612t, 2641
 diagnosis of, 2610, 2612t
 epidemiology of, 2641
 imaging in, 441e-22f, 2641, 2641f
 ocular involvement in, 196
 parkinsonism in, 163, 2610, 2612t
 prions in, 444e-10, 453e-6
 treatment of, 2641
- Multiplex panels, 447

- Multiplex sequencing, 86e-2, 86e-3f
 Multipotent stem cells, 90e-2, 453t, 454, 454f
 Multivariate statistical models, 3
 Mumps, **231e-1**
 clinical features of, 231e-2-3, 231e-2f
 complications of, 231e-2-3, 893
 diagnosis of, 231e-3
 differential diagnosis of, 231e-3
 epidemiology of, 231e-1-2
 etiology of, 231e-1, 231e-1f
 global considerations in, 231e-1, 231e-1f
 immunity to, 231e-4
 incidence pre- and postvaccine, U.S., 785t
 meningitis in, 908t
 orchitis in, 2365
 pathogenesis of, 231e-2
 prevention of, 231e-3-4. *See also* Measles-mumps-rubella vaccine (MMR)
 transmission of, 231e-2
 treatment of, 231e-3
 Munchausen syndrome, 2719
 Mupirocin
 action of, 931t, 932, 932f
 in burn patients, 166e-3
 for impetigo, 967
 indications for, 941t
 nasal, in staphylococcal skin infections, 345
 prophylactic, 945t
 resistance to, 931t, 934, 941t
 Mu receptors, 468e-1, 468e-1t, 468e-2f
 Murine (endemic) typhus. *See* Endemic (murine) typhus
 Murmurs. *See* Heart murmurs
 Murphy's sign, 283, 2080
 Murray valley encephalitis virus, 1306t
 Murutucú virus, 1309t
 Muscarinic agonists, 473e-10t
 Muscle(s)
 atrophy of, 462e-3-4
 electrophysiologic studies of. *See* Electromyography (EMG)
 enlargement of, 462e-3
 fatigability of, 152, 154
 response to repetitive nerve stimulation, 442e-5-6
 Muscle biopsy
 in central core disease, 462e-12
 in dermatomyositis, 2200f
 in inclusion body myositis, 2201f
 in mitochondrial DNA diseases, 85e-5f, 85e-7
 in muscular dystrophy, 462e-6
 in myopathies, 462e-5
 in polymyositis, 2200f
 Muscle bulk, 154
 Muscle compartment syndrome, 2738
 Muscle contracture, 462e-3
 Muscle cramps, 462e-2-3, 479e-2, 1824
 Muscle disorders
 calcium channel, 462e-16-17, 462e-17f, 462e-17t
 chloride channel, 462e-18
 energy metabolism, 462e-14-15. *See also* Glycogen storage diseases
 muscle fiber. *See* Myopathy(ies)
 muscular dystrophy. *See* Muscular dystrophy
 potassium channel, 462e-17t, 462e-18
 sodium channel, 462e-17-18, 462e-17f, 462e-17t
 Muscle dysphoria syndrome, 7e-2
 Muscle-eye-brain disease, 462e-9, 462e-10t
 Muscle pain, 462e-2, 462e-4t
 Muscle phosphorylase deficiency (McArdle's disease; type V GSD), 433e-2t, 433e-4, 462e-14
 Muscle phosphorylase kinase deficiency, 433e-2t
 Muscle relaxants, 119, 377, 473e-9t, 473e-10t
 Muscle spindle, 155f
 Muscle stiffness, 462e-3
 Muscle strength, 71, 74f, 2220, 2538
 Muscle stretch (tendon) reflexes, 154, 154t, 155f, 2538
 Muscle tone, 154, 154t, 2538
 Muscle wasting, 81, 1268, 1704
 Muscle weakness. *See* Weakness
- Muscular dystrophy, **462e-1**
 AV conduction block in, 1471, 1471t
 Becker's. *See* Becker's muscular dystrophy
 clinical features of, 462e-1-4, 462e-3f
 congenital, 462e-9, 462e-10t
 diagnosis of, 462e-1f, 462e-2f, 462e-3t, 462e-4-5
 Duchenne's. *See* Duchenne's muscular dystrophy
 Emery-Dreifuss. *See* Emery-Dreifuss muscular dystrophy
 facioscapulohumeral, 462e-1, 462e-2f, 462e-5t, 462e-11
 gait disorders in, 163
 genetic considerations in, 434, 436, 436t
 limb-girdle, 462e-6, 462e-6t, 462e-7t
 myotonic. *See* Myotonic dystrophy
 oculopharyngeal, 208, 462e-5t, 462e-11
 progressive, 462e-5t
 proteins associated with, 462e-6, 462e-8, 462e-8f, 462e-9f
- Musculoskeletal disorders, **2216**
 approach to the patient, **2216**, 2216t, 2217f, 2218f
 hand pain, 2220-2221, 2220f
 hip pain, 2222, 2222f, 2230
 knee pain, 2221-2222, 2230
 shoulder pain, 2221, 2221f
 arthritis. *See* Arthritis
 articular vs. nonarticular, 2216
 chest discomfort/pain in, 96t, 98
 drug-induced, 2218t
 in the elderly, 2219
 in hemochromatosis, 2516-2517
 history in, 2218-2219, 2218f
 in hospitalized patient, 2219
 hyperlipidemia and, 2242-2243
 imaging in, 2223-2225, 2224t, 2225f
 infections
 actinomycotic, 1089-1090
 in cancer patient, 489-490
 in joints. *See* Infectious (septic) arthritis
 in prosthetic joints. *See* Prosthetic joint infections
 inflammatory vs. noninflammatory, 2215-2217
 laboratory studies in, 2222-2223, 2223t, 2224f
 obesity and, 415e-6
 osteoarthritis
 physical examination in, 2219-2220
 prevalence of, 2216
 in rheumatic fever, 2151
 in sarcoidosis, 2209, 2209f
 SLE and, 2127
 in systemic sclerosis, 2162
 terminology related to, 2219t
 Mushrooms, poisonous, 267, 283, 473e-9t, 473e-10t
 Mushroom worker's lung, 1681t
 Musk ambrette, 388, 389t
 MuSK antibodies, 2702
 Mussels, contaminated, 2743
 Mustard gas, 262e-1t, 506
 MUT4 gene mutations, 539
 Mutation(s), **432**. *See also* specific genes
 alleles, 435
 allelic heterogeneity in, 435
 in cancer cells, 102e-1. *See also* Cancer genetics
 chromosomal disorders. *See* Chromosome abnormalities/disorders
 definition of, 432
 dynamic, 434
 functional consequences of, 434-435
 genotypes, 435
 identifying, 442-445, 443t
 imprinting of, 439
 locus heterogeneity in, 435-436, 436t
 in Mendelian disorders. *See* Mendelian disorders
 mitochondrial. *See* Mitochondrial DNA diseases
 mosaicism in, 439
 nucleotide repeat disorders, 440, 440t
- origins and types of, 432-434, 433f
 penetrance of, 437
 phenocopies, 436
 phenotypes, 435
 phenotypic heterogeneity in, 435, 436t
 rate of, 433
 sex-influenced, 437
 somatic, 440
 uniparental disomy in, 439
 variable expressivity of, 436-437
 in viruses, 214e-5
 X-inactivation of, 439
 Mutation accumulation theory of aging, 94e-2
 Mutator phenotype, 101e-2
 Mutism, 179
 Mutualist, 86e-1
 MVAC regimen, 577
 MWS. *See* Muckle-Wells syndrome (MWS)
 Myalgia, 462e-2, 462e-4t
 Myasthenia gravis, **2701**
 APS-2 and, 2346t, 2347t
 autoantibodies in, 372e-24t, 377e-4, 2701
 vs. botulism, 989
 clinical features of, 157, 2701-2702
 diplopia, 208
 dysphagia, 257
 dyspnea, 47e-2
 nystagmus, 210
 ocular, 2701-2702
 ptosis, 41e-1v, 207
 weakness, 156, 157, 2701-2702
 congenital syndromes of, 2702, 2703t
 diagnosis of, 2702-2703, 2702t, 2703t, 2704t
 differential diagnosis of, 2702-2703
 disorders associated with, 2703-2704, 2704t
 drug-induced, 462e-20
 drugs to avoid in, 2706, 2706t
 HLA gene association with, 373e-8t
 muscle response to repetitive nerve stimulation in, 442e-6
 pathophysiology of, 2701, 2701f
 pemphigus and, 371
 prevalence of, 2701
 pulmonary function values in, 306e-5, 306e-5f
 reflex responses in, 154
 thymoma and, 123e-2, 2348
 treatment of, **2704**
 algorithm for, 2704f
 assessment of response to, 2706, 2706f
 cholinesterase inhibitors, 2704
 immunosuppressants, 2705
 IVIg, 2705-2706
 plasmapheresis, 2705
 thymectomy, 123e-2, 2704
 Myasthenic crisis, 2706
 Myasthenic fatigue, 2701
 MYB gene mutations, 101e-2t
 Mycetoma, 228, 308e-17f, 1090
 MYC gene, 454
 MYC gene mutations, 101e-3t, 508
Mycobacterium abscessus, 205e-9, 776-777, 1132
Mycobacterium africanum, 1103
Mycobacterium avium complex (MAC)
 infections
 bronchiectasis, 1694, 1696. *See also* Bronchiectasis
 diagnosis of, 1131
 disseminated, 1129-1130
 in HIV infection, 269, 1131, 1251t, 1254
 vs. IBD, 1956
 lymphadenopathy, 1130
 prophylaxis for, 1251t
 pulmonary, 1130
 treatment of, 205e-8, 1131-1132
Mycobacterium bovis, 1103
Mycobacterium bovis bacille Calmette-Guérin. *See* Bacille Calmette-Guérin (BCG) vaccine
Mycobacterium canettii, 1103
Mycobacterium chelonae infections, 205e-9, 829
Mycobacterium fortuitum infections, 205e-9
Mycobacterium spp. infections
 arthritis, 836
 chronic meningitis, 907t

- diagnosis of, 150e-2, 150e-2t, 150e-5t, 150e-6
 inherited susceptibility to, 420t, 421, 422f
 laryngitis, 234
 leprosy. *See* Leprosy
 nontuberculous. *See* Nontuberculous
 mycobacterial infections
 osteomyelitis, 836, 839, 844
 subacute meningitis, 766t, 775, 898–899,
 1111
 tuberculosis. *See* Tuberculosis (TB)
- Mycobacterium intracellulare* infections. *See*
Mycobacterium avium complex (MAC)
 infections
- Mycobacterium kansasii* infections, 205e-8, 1130,
 1131–1132
- Mycobacterium leprae*, 145e-4, 1123. *See also*
 Leprosy
- Mycobacterium marinum*, 1106
- Mycobacterium marinum* infections
 cellulitis, 830
 skin, 366, 829, 1130
 in transplant recipient, 926
 treatment of, 205e-8, 830, 1132
- Mycobacterium oryzae*, 1103
- Mycobacterium pinnipedii*, 1103
- Mycobacterium tuberculosis*. *See also*
 Tuberculosis (TB)
 adherence of, 145e-2t
 host response to, 372e-26, 1106–1107
 innate resistance to, 1106
 microbiology of, 1102–1103, 1103f
 phagosome of, 1105–1106
 rifampin-resistant, 774, 778
 transmission of, 1105
 virulence of, 1106
- Mycobacterium ulcerans*, 829, 1130
- Mycophenolate mofetil/sodium
 action of, 1829t
 adverse effects of, 269, 1829t, 1831, 2071,
 2130t
 for bullous pemphigoid, 372
 for drug-induced hypersensitivity syndrome,
 382
 drug interactions of, 2130t
 for immunosuppression
 after kidney transplantation, 1828, 1829t
 after liver transplantation, 2071
 for inflammatory myopathies, 2201
 for interstitial lung diseases, 1711, 1712
 for lupus nephritis, 1838
 for mucous membrane pemphigoid, 374
 for myasthenia gravis, 2705
 for neuromyelitis optica, 2673
 for pemphigus vulgaris, 371
 for photoallergy, 390
 for SLE, 2130t, 2132
 for systemic sclerosis, 2164
- Mycoplasma* spp., 1163
- Mycoplasma genitalium* infections, 870–871,
 872t, 876, 1164t, 1165
- Mycoplasma hominis* infections
 in mediastinitis, 926
 nonurogenital, 1165
 postpartum/postabortal, 1165
 in transplant recipient, 926
 treatment of, 1164t, 1165
 urogenital, 1165
- Mycoplasma* spp. infections
 AGEP and, 382
 in animal bite-wounds, 167e-1
- Mycoplasma pneumoniae*, 1163, 1164
- Mycoplasma pneumoniae* infections, **1163**
 diagnosis of, 1164, 1164t
 epidemiology of, 1163
 extrapulmonary, 1163–1164
 pathogenesis of, 1163
 treatment of, 1164, 1164t
 upper respiratory tract, 226, 231, 1163
- Mycosis fungoides (cutaneous T-cell lymphoma)
 clinical features of, 501f, 707
 peripheral blood smear in, 81e-10f
 skin manifestations of, 76e-5f, 353, 353t, 354,
 357t, 707
 treatment of, 102e-5t, 103e-20, 391, 707
- Mycotic aneurysm
 circle of Willis, 1784–1785. *See also*
 Subarachnoid hemorrhage
 clinical features of, 1639
 etiology of, 1638
 in infective endocarditis, 819, 826
 treatment of, 1643
- Mycotoxins, hepatocellular carcinoma and, 544,
 545t
- Myc transcription factor, 500
- MyD88, 86e-8, 145e-5, 145e-6f, 372e-5f
- MyD88 deficiency, 420t
- Mydriasis, 196
- Myelin
 defects of, 444e-3f, 444e-5–6, 444e-5f
 structure of, 444e-4, 444e-4f
- Myelin-associated glycoprotein (MAG),
 372e-24t, 444e-8, 718
- Myelin basic protein (MBP), 372e-24t, 444e-4f
- Myelin gene regulatory factor (MRF), 444e-5
- Myelin oligodendrocyte glycoprotein (MOG),
 372e-24t, 444e-4f
- Myelin protein zero (MPZ), 2678
- Myelitis. *See* Myelopathy, inflammatory and
 immune
- Myeloblasts, 414f, 415
- Myelocytes, 414f, 415
- Myelodysplasia, 415, **669**
 classification of, 669–670, 669t
 clinical features of, 670
 diagnosis of, 668f, 670–671
 differential diagnosis of, 628
 epidemiology of, 670
 etiology of, 670
 genetic considerations in, 670
 hemoglobinopathy in, 639
 pathophysiology of, 670
 prognosis of, 671, 671t
 as second malignancy in cancer survivors, 623
 treatment of, 102e-8, 139e-6, 671
- Myelodysplastic syndromes (MDS). *See*
 Myelodysplasia
- Myelofibrosis
 primary. *See* Primary myelofibrosis
 secondary, 671–672
- Myelography, 440e-2, 440e-9–10
- Myeloid dendritic cells, 372e-6, 372e-7t. *See also*
 Dendritic cells
- Myeloid/erythroid ratio, 397
- Myeloid malignancies, **135e-4**
 atypical chronic myeloid leukemia, 135e-5–6,
 135e-5t
 chronic myelomonocytic leukemia, 102e-3,
 135e-5t, 135e-6, 669
 chronic neutrophilic leukemia, 135e-5–6,
 135e-6t, 672
 classification of, 135e-4, 135e-4t
 juvenile myelomonocytic leukemia (JMML),
 135e-6
 MDS/MPN, unclassifiable, 135e-6
 MDS/MPN overlap, 135e-6
 myelodysplastic syndromes. *See*
 Myelodysplasia
 myeloproliferative neoplasm, unclassifiable,
 135e-6–7
 transient myeloproliferative disorder (TMD),
 135e-7
- Myeloid metaplasia. *See* Primary myelofibrosis
- Myeloma
 multiple. *See* Multiple myeloma
 sclerotic, 614, 618–619
- Myeloma kidney, 62e-7f, 1859, 1859f
- Myeloma light chains, 1804, 1805, 1859
- Myelopathy
 chemotherapy-induced, 607t
 chronic, **2657**
 in adrenomyeloneuropathy, 2659
 in cobalamin deficiency, 2659
 familial spastic paraplegia. *See* Familial
 spastic paraplegia (FSP)
 HTLV-1 associated, 163, 225e-5, 697, 707,
 2657–2658
 hypocupric, 2659
 in MS, 2659
- primary lateral sclerosis, 2636, 2659
 spondylitic, 2657
 in syringomyelia, 2658, 2658f
 in tabes dorsalis, 1136, 2659
 in tethered cord syndrome, 118, 2659
 toxic, 2659
 in vascular malformations of cord and dura,
 2657, 2658f
 compressive. *See* Spinal cord compression
 compressive vs. noncompressive, 2653
 evaluation of, 2655t
 gait disorders in, 162t
 in high-voltage electrical injury, 2657
 in HIV infection, 1266–1267
 inflammatory and immune (myelitis), **2656**
 acute infectious, 441e-25f, 2656–2657
 in multiple sclerosis, 2656. *See also* Multiple
 sclerosis (MS)
 in neuromyelitis optica, 2656, 2673
 paraneoplastic, 616
 postinfectious, 1212, 2656
 in systemic immune-mediated disorders,
 2656
 noncompressive acute transverse, 2655, 2655t
 paraneoplastic, 618, 2659
 radiation-associated, 622
 in spinal cord infarction, 441e-25f, 2656–2657
- Myeloperoxidase, 372e-24t, 416
- Myeloperoxidase deficiency, 419, 420t
- Myelophthisic anemias, 671–672
- Myeloproliferative disorders, **672**
 classification of, 672, 672t
 cutaneous ulcers in, 369
 evolution into AML, 678
 janus kinase inhibitors for, 102e-3
 splenomegaly in, 411
- Myeloproliferative neoplasm, unclassifiable
 (MPN-U), 135e-6–7
- Myelosuppression, chemotherapy-induced, 103e-
 23–25, 103e-24t. *See also* Neutropenia
- Myenteric plexus, 264
- MYH7 gene mutations, 1555t, 1568
- MYH9 gene, 1241, 1241t
- MYH-associated polyposis, 538t, 539
- Myiasis, 2746–2747
- Myoadenylate deaminase deficiency, 431e-3t,
 431e-6, 462e-15
- Myocardial contusion, 289e-3
- Myocardial fibrosis, radiation-associated, 621
- Myocardial infarction (MI)
 abdominal pain in, 105, 106t
 vs. acute pericarditis, 1571, 1575
 arrhythmias in
 AV conduction block, 1471t, 1472, 1609
 SA node dysfunction, 1467, 1467t
 sinus bradycardia, 1609
 supraventricular, 1609
 ventricular, 1496, 1608–1609, 1770
 atherosclerosis and, 291e-4
 cardiac arrest risk factors following, 1767,
 1767f
 cardiac imaging following, 270e-19–20,
 270e-25f, 1464f
 cardiogenic shock in. *See* Cardiogenic shock
 chest pain in, 99f, 1594, 1599–1600,
 1609–1610
 circadian variations in, 194, 1599, 1616
 classification of, 1602t
 definition of, 1602t
 dental care after, 242
 diagnosis of
 CMR, 271e-4f, 271e-4v
 ECG, 269e-3–7f
 echocardiography, 271e-1f, 271e-1v
 history in, 99, 99f
 myocardial perfusion imaging, 271e-5f
 erectile dysfunction and, 328
 hemodynamic patterns in, 1762t
 in HIV infection, 1255
 hypokalemia in, 64e-7, 305
 hypothermia in, 478e-1
 incidence of, 1593, 1594f, 1599
 inflammatory mediators in, 372e-29t
 in kidney transplant recipients, 1831

- Myocardial infarction (MI) (*Cont.*):
 mitral regurgitation in, 1543
 murmur after, 51e-2, 1600
 natural history of, 1439
 nausea and vomiting in, 259
 non-ST-segment elevation, 1593, 1594, 1595f.
 See also Non-ST-segment elevation
 acute coronary syndrome (NSTEMI-ACS)
 during PCI, 296e-2, 1602t
 postmenopausal hormone therapy and, 2384t
 prevention of, 84, 747
 Q wave abnormalities in, 269e-4f, 269e-5f,
 269e-8f, 1456, 1456f
 radiation-associated, 103e-4, 621
 right ventricular. *See* Right ventricular failure/
 infarction
 ST-segment elevation. *See* ST-segment
 elevation myocardial infarction
 (STEMI)
 systems-based analysis of, 87e-4
 treatment of
 fibrinolytic therapy, 758
 implantable cardiac defibrillator, 1770
 pacemaker, 1475-1476, 1475t
 racial/ethnic disparities in, 16e-2, 16e-3f
 in women, 6e-3, 6e-3f
- Myocardial ischemia. *See also* Ischemic heart
 disease
 anterior wall, 17, 269e-1f, 1455f
 anterolateral wall, 269e-2f
 in aortic stenosis, 1579
 clinical features of, 1439
 diagnosis of, 1440
 dyspnea in, 47e-1t, 47e-3
 ECG in, 269e-1-3f, 1455-1457, 1455f, 1456f,
 1457f, 1580
 effects of, 1579-1580
 lateral, 269e-2f
 with left ventricular hypertrophy, 1579
 pathophysiology of, 1578-1579, 1580f
 silent, 1580, 1592-1593, 1598
 transmural, 17, 1455f
 ventricular tachycardia in, 1496
- Myocardial perfusion imaging
 after myocardial infarction, 270e-20, 270e-20f
 in CAD evaluation, 270e-10, 270e-10f,
 270e-11f, 271e-2f, 271e-3f
 in chest pain evaluation, 271e-3f, 271e-3v
 indications for, 270e-3
 in ischemic heart disease, 1582, 1584f
 multidimensional, 270e-5f
 in myocardial infarction, 1601
 in myocardial infarction evaluation, 271e-5f
 PET vs. SPECT, 270e-3
 protocols for, 270e-3
 radiation exposure from, 270e-8
 radiopharmaceuticals for, 270e-3, 270e-3t
 in wall motion abnormalities, 270e-4f
- Myocarditis, 1557
 bacterial, 1561
 cardiogenic shock in, 1762. *See also*
 Cardiogenic shock
 CMR in, 270e-15, 270e-15f
 in dermatomyositis, 1562
 dilated cardiomyopathy and, 1558t. *See also*
 Dilated cardiomyopathy
 in diphtheria, 979
 eosinophilic, 1562
 fungal, 1561
 giant cell, 1561-1562
 granulomatous, 1561
 in HIV infection, 1560
 hypersensitivity, 1562
 in influenza, 1212
 parasitic, 1560-1561
 pathophysiology of, 372e-24t
 in relapsing fever, 1147
 rickettsial, 1561
 SA node dysfunction in, 1467, 1467t
 in sarcoidosis, 1561, 1562
 spirochetal, 1561
 viral
 acute, 1558, 1762
 chronic, 1558
 clinical features of, 1558
 diagnosis of, 1558-1560, 1559f
 enteroviral, 1282
 pathophysiology of, 1557-1558
 treatment of, 1560
 viruses causing, 1558
- Myocardium
 oxygen supply and demand, 1578-1579
 regenerative potential of, 265e-12
 rupture of, 289e-3, 1762
- Myoclonic epilepsy and ragged red fiber
 syndrome. *See* MERRF
- Myoclonic seizures, 2544
- Myoclonus, 2623
 Baltic, 451e-4t
 clinical features of, 2619t, 2623
 in Creutzfeldt-Jakob disease, 453e-4
 posthypoxic, 1782
 treatment of, 2623
- Myocytes, 265e-5
- Myofascial pain syndrome, 93, 237, 2246
- Myofasciitis, 2199
- Myofibrillar myopathies, 462e-12, 462e-13t
- Myogenic ptosis, 207
- Myogenic reflex, 332e-2, 1800
- Myoglobin, 625, 625t, 1804, 2762t
- Myoglobinuria, 462e-15, 1212
- Myokymia, 462e-3, 2649, 2664
- Myonecrosis
 anaerobic bacterial, 1099
 clostridial. *See* Gas gangrene (clostridial
 myonecrosis)
 etiology of, 828t
 with necrotizing fasciitis, 831
 synergistic nonclostridial anaerobic, 831
- Myopathy(ies)
 acute necrotizing, 619, 1784
 alcoholic, 2725
 clinical features of, 462e-1-4, 462e-2t, 462e-3t
 congenital, 462e-12, 462e-14
 in critical/systemic illness, 462e-19, 1784
 in diabetes mellitus, 462e-19
 diagnosis of, 462e-1f, 462e-2f, 462e-4-5
 distal, 462e-12, 462e-13t
 drug-induced, 462e-19-20, 462e-20t, 2199,
 2218t, 2447-2448
 gait disorders in, 163
 hereditary, 462e-5, 462e-5t. *See also* Muscular
 dystrophy
 in HIV infection, 1266-1267
 inflammatory. *See* Inflammatory myopathies
 in lung cancer, 511
 mitochondrial. *See* Mitochondrial
 myopathy(ies)
 in parathyroid disorders, 462e-19
 in pituitary disorders, 462e-19
 progressive, in type IIIa glycogen storage
 disease, 436e-6f
 thick-filament, 1784
 in thyroid disorders, 462e-18
 weakness in, 155-156, 2197-2199
- Myopericarditis, vaccinia vaccine and, 261e-7
- Myophosphorylase deficiency (McArdle's
 disease; type V GSD), 433e-2t, 433e-4,
 462e-14
- Myopia, 195
- Myosin
 in cardiac contraction, 265e-5, 265e-5f
 defects in, 1556f
 in rheumatic fever, 372e-24t
- Myosin-binding protein C gene, 437
- Myosin light chain, 265e-4
- Myositis
 clinical features of, 831
 etiology of, 828t, 831
 inclusion body. *See* Inclusion body myositis
 in influenza, 1212
 necrotizing autoimmune, 2198
 in SLE, 2127
 streptococcal, 831, 968
 treatment of, 832-833
 in trichinellosis, 1410
- Myositis ossificans, 426e-7t, 426e-8
 Myositis ossificans progressiva, 426e-7, 426e-7t
- Myotonia
 channelopathies in, 444e-2t
 in chloride channel disorders, 462e-18
 conditions associated with, 462e-3
 definition of, 462e-3
 diagnosis of, 462e-4, 462e-4t
 in muscular dystrophy, 462e-10
 potassium-aggravated, 462e-18
- Myotonia congenita, 444e-2t
- Myotonic discharges, EMG, 442e-4
- Myotonic dystrophy, 462e-9
 AV conduction block in, 1471, 1471t
 clinical features of, 462e-5t, 462e-10
 congenital, 462e-10
 diagnosis of, 462e-10-11
 genetic considerations in, 434, 440t, 462e-5t,
 462e-10-11
 ocular involvement in, 208
 SA node dysfunction in, 1467, 1467t
 testicular dysfunction in, 2366
 treatment of, 462e-11
- Myotubular (centronuclear) myopathy, 462e-12,
 462e-14
- MyPlate Food Guide, 95e-3
- MYPPC3 gene mutations, 1555t, 1568
- Myringoplasty, 230
- Myringotomy, 230
- Mystic sage (*Salvia divinorum*), 469e-4
- Myxedema
 clinical features of, 2291
 edema in, 253
 etiology of, 1651
 hypothermia in, 478e-3
 pericardial effusion in, 1575
 pretibial, 76e-16f, 1654
- Myxedema coma, 2293
- Myxofibrosarcoma, 119e-1
- Myxoma, 289e-1
 clinical features of, 289e-1
 CMR in, 270e-24, 289e-2f
 dyspnea in, 47e-3
 echocardiography in, 289e-1, 289e-1f
 epidemiology of, 289e-1
 heart sounds/murmur in, 51e-6, 289e-1
 vs. mitral stenosis, 1541
 palpitations in, 254
 pathology of, 289e-1
 skin manifestations of, 359
 syncope in. *See* Cardiac syncope
 treatment of, 289e-2
- N
- NAAT (nucleic acid amplification test), 214e-9,
 769t, 1247, 1247t
- Nab-paclitaxel, 103e-12t, 103e-15, 557, 557t
- Nabumetone, 2232
- N-acetyl-benzoquinone-imine (NAPQI), 2027
- N-acetylcysteine (NAC), 43
 for acetaminophen overdose, 2027
 for COPD, 1706
 for idiopathic pulmonary fibrosis, 1712
 for prevention of contrast-induced
 nephropathy, 1461
 for sulfur mustard exposure, 262e-4
- N-acetylprocainamide (NAPA), 35
- Na⁺-Cl⁻ co-transporter, thiazide-sensitive, 296,
 310
- Nadolol
 for arrhythmias, 1480t
 dosage of, 1480t
 for ischemic heart disease, 1589t
 for long QT syndrome, 1497
 overdosage/poisoning with, 473e-9t
 pharmacology of, 1480t
- Naegleria, 1367
- Naegleria infections
 clinical features of, 1367
 encephalitis, 896
 epidemiology of, 245e-4t, 1367
 laboratory evaluation of, 245e-4t, 245e-6t,
 1367
 pathogenesis of, 896, 1367
 treatment of, 246e-11, 1367
- Nafamostat, 310

- Nafcillin
adverse effects of, 306
for cellulitis, 228, 832t
drug interactions of, 939t
indications for, 937, 940t
for infective endocarditis, 823
for meningitis, 887t, 889t
for nongonococcal bacterial arthritis, 835
for osteomyelitis, 840t, 841t
for perichondritis, 228
resistance to, 933, 940t
for staphylococcal infections, 961, 962t
- Nafcillin/oxacillin, 768t
- NAFLD. *See* Nonalcoholic fatty liver disease (NAFLD)
- Nail(s)
in amyloidosis, 721, 723f
drug-induced disorders of, 380, 380f
in HIV infection, 1263
in lichen planus, 76e-3f, 349, 349f
in nutritional deficiencies, 463t
onychomycosis, 350
in psoriasis, 348
in psoriatic arthritis, 2176, 2176f
in systemic sclerosis, 376
telangiectasia, 356t, 357
- Nail-patella syndrome, 1835t, 1847
- Nairobi fly, 2750
- Nairobi sheep disease virus, 1308t
- Nairoviruses, 1305, 1308t
- Na⁺-K⁺-2Cl⁻ cotransporter, 296, 332e-8
- Na⁺/K⁺-ATPase, 296, 332e-8
- Nalidixic acid, 389t, 727t, 932–933
- Nalmefene, 2728
- Naloxone, 92, 468e-3, 468e-3t
- NALP1b inflammasome, 145e-7
- NALP3 gene, 372e-8t
- NALP7 gene, 372e-8t
- NALP12 gene, 372e-8t
- Naltrexone
in alcoholism management, 2728
with bupropion, for obesity, 2397
in opioid use disorders, 468e-3, 468e-4
- NAME syndrome. *See* Carney's syndrome/complex (LAMB/NAME syndrome)
- Naming deficit, 177
- ¹⁵N-ammonia, 270e-3, 270e-3t
- Nanocrystalline silver dressing, 166e-3
- Nanog gene, 455
- NAP-2 (neutrophil-activating protein-2), 372e-12t
- NAPA (N-acetylprocainamide), 35
- Naphthalene, 655, 656t
- NAPQI (N-acetyl-benzoquinone-imine), 2027
- Naproxen
adverse effects of, 362, 727t, 2232, 2232t
for dysmenorrhea, 338
for gout, 2234
for medication-overuse headache, 109
for migraine, 2592t
for osteoarthritis, 2232t
for pain, 91t
for rheumatic fever, 2153
- Napsin-A (Nap-A), 508
- Naratriptan, 2591, 2592t, 2593t
- Narcolepsy, 189
clinical features of, 188t, 189
diagnosis of, 188t, 189–190, 189f
genetic considerations in, 189, 373e-8t
HLA gene association with, 373e-8t
olfactory dysfunction in, 215
treatment of, 188t, 190
- Narcotics. *See* Opioid(s)
- NARES (perennial nonallergic rhinitis with eosinophilia syndrome), 2122
- NARP (neuropathy, ataxia, retinitis pigmentosa) syndrome, 85e-6, 85e-6t, 451e-3t
- Narrow band imaging (NBI), 1668
- Nasal congestion. *See* Allergic rhinitis; Common cold
- NASBA (nucleic acid sequence–based amplification), 1247, 1247t
- NASH. *See* Nonalcoholic steatohepatitis (NASH)
- Nasoduodenal tube, 98e-8t
- Nasogastric tube, 98e-8t, 257
- Nasopharyngeal cancer. *See also* Head and neck cancer
EBV infection and, 214e-7, 502, 1188–1189
formaldehyde exposure and, 1692t
histopathology of, 503
- Nasotracheal intubation, 226
- NAT-2, 39t, 40
- Natalizumab
adverse effects of, 372e-31, 1962, 2669
for IBD, 1962
for MS, 372e-31, 2669, 2669t, 2671
- Nateglinide, 2413, 2414t
- National Childhood Vaccine Injury Act (NCVIA), 790
- National Cholesterol Education Project Adult Treatment Panel III (ATP III), 480e-3t
- National Family Caregivers Association, 66
- National Health Care Disparities Report, 16e-2
- National Healthcare Safety Network, 913
- National Institutes of Health National Center for Complementary and Alternative Medicine (NCCAM), 14e-4t
- National Library of Medicine (NLM), 14e-4t
- National Lung Cancer Screening Trial (NLST), 509–510, 510t
- National Quality Forum, 12e-2
- Native American medicine, 14e-1t
- Natural cytotoxicity receptors (NCRs), 372e-7, 1238
- Natural helper cells, 2116
- Natural killer (NK) cells
functions of, 372e-1, 372e-6–7, 372e-7t
signaling of, 372e-7, 372e-15f
surface receptors of, 372e-7
- Natural killer (NK) T cells
in adaptive immunity, 372e-7t
in asthma, 1673
definition of, 372e-2
functions of, 372e-7
in HBV infection, 2010
hematopoietic differentiation of, 89e-3f
in HIV infection, 1238, 1245
HLA class I receptors and recognition of, 373e-4–5
in innate immunity, 372e-7, 372e-7t
neoplasms of, 135e-3–4, 697, 708
receptors of, 372e-7
signaling of, 372e-7
- Natural Medicines Comprehensive Database, 14e-4t
- Natural orifice transluminal endoscopic (NOTES) surgery, 346e-1v, 1881, 1883, 1886f
- Natural Standard, 14e-4t
- Naturopathy, 14e-1t, 14e-3
- Nausea, 258, 259
- Nausea and vomiting, 258
in appendicitis, 1987
approach to the patient, 259–260
chemotherapy-induced, 103e-25, 474
acute emesis, 259, 474
anticipatory, 62, 103e-25, 259, 474
delayed emesis, 259, 474
treatment of, 103e-25, 261, 474
differential diagnosis of, 258–259, 258t, 1877t
in gastric cancer, 534
gastrointestinal causes of, 1876
in liver disease, 1991
in myocardial infarction, 100
in pregnancy, 49, 259, 261
radiation-induced, 103e-4
in sepsis/septic shock, 1755
in terminally ill patient, 61–62
treatment of, 260–261, 260t
- Naxos syndrome, 1555t
- Nazer prognostic index, for Wilson's disease, 2520, 2521t
- NBI (narrow band imaging), 1668
- NBIA (neurodegeneration with brain iron accumulation), 2625–2626
- NBTE (nonbacterial thrombotic endocarditis), 817
- NCCAM (National Institutes of Health National Center for Complementary and Alternative Medicine), 14e-4t
- NCF (neutrophil chemotactic factor), 372e-15t
- NCRs (natural cytotoxicity receptors), 372e-7, 1238
- NCVIA (National Childhood Vaccine Injury Act), 790
- NEB gene, 462e-12
- Nebivolol, 1507, 1589t, 1624
- Necator americanus*, 245e-3t, 1414, 1414t. *See also* Hookworm infections
- necdin* gene, 439, 2256
- Neck pain, 121
in ankylosing spondylitis, 122
in brachial plexus injury, 122
in cancer, 122
in carpal tunnel syndrome, 122
in cervical disk disease, 121t, 122
in cervical spine trauma, 121–122
in cervical spondylosis, 122
in herpes zoster, 122
in orthostatic hypotension, 146
in peripheral nerve disorders, 122–123
physical examination in, 121t
in rheumatoid arthritis, 122
in thoracic outlet syndromes, 122
treatment of, 123
- Neck weakness, 2649
- Necrobiosis lipoidica, 364t, 365
- Necrobiosis lipoidica diabetorum, 436e-2f, 2429–2430
- Necrolytic migratory erythema, 355
- Necrosis
in cancer cells, 102e-10, 103e-6
intestinal, 312
- Necrotic arachnidism, 2747
- Necrotizing arteritis, 2567–2568
- Necrotizing autoimmune myositis, 2198
- Necrotizing encephalopathy, 622
- Necrotizing enterocolitis, 991
- Necrotizing fasciitis
clinical features of, 783, 831, 968, 2199
etiology of, 780t, 783, 828t, 830, 968, 1099, 2199
imaging of, 831f
pathophysiology of, 830–831, 968
treatment of, 780t, 832–833, 832t, 965t, 968
- Necrotizing myelopathy, paraneoplastic, 614t
- Necrotizing otitis externa, 229
- Necrotizing pancreatitis. *See also* Acute pancreatitis
diagnosis of, 2093t, 2094f, 2095f
intraabdominal abscess in, 849
pathogenesis of, 2092
treatment of, 346e-1v
- Necrotizing vasculitis
clinical features of, 340, 342, 342f, 367
differential diagnosis of, 369
pathogenesis of, 372e-8
- Nedocromil sodium, 1678
- Neer test, 2221
- nef* gene, 225e-3f, 1219f
- Nef protein, 1234
- Negative predictive value, 479, 479t
- Negative-strand genome, 214e-1
- Neglect (sensory phenomenon)
caring for patient with, 184
definition of, 160, 180
hemispatial, 180–182, 180f, 181f
- Negligible senescence, 94e-2, 94e-2f
- Negri body, 1301, 1302f
- Neisseria* spp. (meningococci). *See also* Meningococcal infections
adherence of, 145e-2, 145e-2t
laboratory identification of, 150e-2t
serogroups of, 995–996, 995t
tissue invasion by, 145e-7
tissue tropisms of, 145e-8–9
- Neisseria gonorrhoeae*. *See also* Gonococcal infections
antibiotic resistance in, 872f, 933, 1004–1005
host response to, 1004

- Neisseria gonorrhoeae* (Cont.):
laboratory identification of, 150e-2t, 150e-6, 1007-1008
lipooligosaccharide of, 1004
microbiology of, 1003
outer-membrane proteins of, 1003-1004
tissue tropisms of, 145e-8-9
- Neisseria meningitidis*. See also Meningococcal infections
antibiotic resistance in, 933
asymptomatic carriage of, 998
colonization by, 998
host response to, 998
laboratory identification of, 150e-2t
outer-membrane vesicles, 1002, 1002f
structure of, 995-996, 995f, 995t, 996f
tissue tropisms of, 145e-8-9
- Neisseria meningitidis* vaccine. See Meningococcal vaccine
- Nelarabine, 103e-11t
- NELF gene mutations, 2364t
- Nelfinavir, 246e-4t, 1276f, 1280f
- Nelson's syndrome, 360, 2273
- Nemaline myopathy, 462e-12
- Nematodes/nematode infections, 255e-1
- filarial
Brugia spp., 245e-3t, 1417, 1418t. See also Lymphatic filariasis
Dracunculus medinensis. See *Dracunculus medinensis* (guinea worm)/dracunculiasis
Loa loa. See *Loa loa* (African eye worm)/loiasis
Mansonella spp., 245e-6t, 1418t, 1422
Onchocerca volvulus. See *Onchocerca volvulus*/onchocerciasis
tropical pulmonary eosinophilia, 422, 1420
Wuchereria bancrofti, 245e-3t, 765t, 1417, 1418t. See also Lymphatic filariasis
zoonotic, 1423
- intestinal, 1404t, **1413**
anisakiasis, 1416-1417
Ascaris lumbricoides. See *Ascaris lumbricoides*/ascariasis
Capillaria philippinensis/capillariasis, 245e-3t, 765t, 1417
Enterobius vermicularis/enterobiasis (pinworm), 245e-3t, 245e-5t, 1414t, 1416
hookworm. See Hookworm infections
Strongyloides stercoralis. See *Strongyloides stercoralis*/strongyloidiasis
trichostrongyliasis, 1416
Trichuris trichiura. See *Trichuris trichiura*/trichuriasis (whipworm)
- tissue, **1410**
Ancylostoma braziliense, 245e-3t, 1415
Ancylostoma caninum, 1415
Angiostrongylus. See *Angiostrongylus* spp. infections
Gnathostoma. See *Gnathostoma spinigerum*/gnathostomiasis
larva migrans syndromes. See also Larva migrans
Ancylostoma braziliense, 245e-3t, 1412
Baylisascaris procyonis. See *Baylisascaris procyonis* infections
Toxocara canis, 245e-3t, 765t, 1411
Toxocara cati, 245e-3, 765t, 1411
Trichinella. See *Trichinella*/trichinellosis
- NEMO (NF- κ B essential modulator)
in cellular signaling pathway, 145e-6f
deficiency of, 420t, 421, 2106, 2111
in nontuberculous mycobacterial infections, 1129, 1129f
- Neoadjuvant therapy, 103e-3
- Neobladder, orthotopic, 577
- Neocarzinostatin, 550
- Neomycin
action of, 942
adverse effects of, 269, 381, 644t, 942
for hepatic encephalopathy, 2066
for irritable bowel syndrome, 1969
for otitis externa, 228
prophylactic, 945t
vitamin A absorption and, 96e-6
- Neonatal-onset multisystem inflammatory disease (NOMID), 2213t, 2215
- Neonate(s)
diabetes in, 2406
infections in
C. perfringens, 991
C. trachomatis, 1169, 1170t, 1171
Candida, 441e-8f
CMV, 50, 1191, 1191t
enterococcal, 974
enterovirus (generalized disease of the newborn), 1291, 1291t
gonococcal, 1006, 1009t
group B *Streptococcus*, 970
herpesvirus, 50
HIV, 1226
HSV, 215e-3t, 1180
L. monocytogenes, 983
rubella. See Rubella, congenital
syphilis. See Syphilis, congenital
T. gondii, 1398, 1399-1400, 1402, 1404
tetanus, 984
varicella, 1184
jaundice in, 653, 654, 2001
seizures in, 2545, 2546t
vitamin K deficiency in, 96e-8
- Neorickettsia*, 1159
- Neostigmine
for anticholinesterase testing, 2702
for intestinal pseudoobstruction, 1985
overdosage/poisoning with, 473e-10t
for snakebite treatment, 2737, 2737t
- Nephrectomy, 578-579
- Nephritis. See Acute interstitial nephritis (AIN); Acute kidney injury (AKI); Glomerulonephritis; Lupus nephritis; Pyelonephritis
- Nephrocalcinosis, 1862
- Nephrogenic fibrosing dermatopathy, 1819
- Nephrogenic systemic fibrosis, 270e-9, 440e-5
- Nephrolithiasis, **1866**
abdominal pain in, 106t
approach to the patient, **1868**
back pain in, 118
clinical features of, 289t, 1868
conditions associated with, 1866
diagnosis of, 289t, 1868-1870, 1868f, 1869f
differential diagnosis of, 1868
epidemiology of, 1866
genetic considerations in, 1867-1868
history in, 1869
in hyperparathyroidism, 2472
hyperuricemia and, 431e-4f
inherited, 332e-7t
pathogenesis of, 1866-1867
prevention of, 1869, 1870
risk factors for, 1867
treatment of, 431e-4-5, 1870-1871
in Wilson's disease, 2519
- Nephron(s)
embryologic development of, 332e-1-2, 332e-1f, 333e-1
functions of, **332e-3**
cellular transport, 332e-3
in collecting duct, 332e-5f, 332e-6f, 332e-8-9
in distal convoluted tubule, 332e-4f, 332e-8
epithelial solute transport, 332e-3
in loop of Henle, 332e-4f, 332e-8
membrane transport, 332e-6
in proximal tubule, 332e-3f, 332e-6-8
inherited disorders involving, 332e-7t
reduced number of, 333e-1-2, 333e-1t
- Nephronophthisis, 1850t, 1851f, 1855
- Nephropathia epidemica, 1321
- Nephropathy
analgesic, 1861, 1861f
aristolochic acid, 577, 1804, 1856t, 1861
atherosclerotic, 1835t, 1847
calcineurin-inhibitor, 1829, 1862
contrast-induced, 440e-2, 1461, 1803, 1806t, 1820
diabetic. See Diabetic nephropathy
heavy metal, 1862
HIV-associated, 62e-1f, 1258-1259, 1849
hypercalcemic, 1862
hypokalemic, 1862
IgA. See IgA nephropathy
light chain cast. See Light chain deposition disease/nephropathy
lithium-associated, 1861-1862
membranous. See Membranous glomerulonephritis
phosphate, 62e-11f, 1859
quartan malarial, 1375
radiation-induced, 1864
reflux, 1859-1860, 1860f
sickle cell, 1835t, 1848, 1860, 1865-1866
urate, 431e-4, 1858
uric acid. See Uric acid nephropathy
- Nephrosclerosis, hypertensive, 62e-9f, 1617, 1630, 1835t, 1847-1848
- Nephrosis, urate, 431e-4
- Nephrotic syndromes, **1841**
clinical features of, 289t, 1835t, 1836, 1841
acid-base disorders, 317
ascites, 287
edema, 252, 252t
hypoalbuminemia, 294
hyponatremia, 299
diabetic nephropathy. See Diabetic nephropathy
diagnosis of, 289t
in Fabry's disease. See Fabry's disease
focal segmental glomerulosclerosis. See Focal segmental glomerulosclerosis
glomerular deposition diseases, 1845
fibrillary-immunotactoid glomerulopathy, 1835t, 1845
light chain deposition disease. See Light chain deposition disease
renal amyloidosis, 62e-6f, 1835t, 1845
lipoprotein disorders in, 2438-2439
membranous glomerulonephritis. See Membranous glomerulonephritis
minimal change disease. See Minimal change disease
treatment of, 1841
- Nephroureterectomy, 577
- Nepuyo virus, 1309t
- Nerve abscess, in leprosy, 1126
- Nerve agents, **262e-4**
action of, 262e-4, 262e-4f, 473e-10t
clinical features of exposure to, 262e-4-5, 473e-10t
treatment of exposure to, 262e-5-7, 262e-5t, 262e-6f, 473e-10t
- Nerve biopsy, indications for, 2677
- Nerve conduction studies
in back pain, 114
in peripheral neuropathy, 2676-2677, 2677t
principles of, 442e-4-5, 442e-5f
- Nerve growth factor (NGF), 444e-7t
- Nesidioblastosis, 570, 572, 2434
- Nesiritide, 1509, 1510t, 1763
- Netilmicin, 930
- NETs. See Neuroendocrine tumors (NETs)
- NETs (neutrophil extracellular traps), 1753
- Networks
disease, 87e-3-4, 87e-5f, 87e-6f
gene, 87e-3, 87e-6f, 87e-7f
neural, 177
organization of, 87e-2, 87e-2f
- Neural foraminal narrowing with radiculopathy, 116, 116f
- Neuralgia(s)
glossopharyngeal, 237, 2649
migrainous, 237
trigeminal. See Trigeminal neuralgia
- Neurally adjusted ventilatory assist ventilation, 1743
- Neural network, 177
- Neural stem cells, 90e-3, 453t
- Neural tube defects, 29t, 47, 642-643
- Neuraminidase, 214e-5, 947, 947f
- Neurasthenia, 2703

- Neuritic plaque, in AD, 2599, 2600f
- Neuritis
 optic. *See* Optic neuritis
 oral, 237
- Neuroanthocytosis, 2622
- Neuroblastoma
 genetic considerations in, 101e-2t, 101e-3
 paraneoplastic syndromes in, 618
 tumor markers in, 473t
- Neurocutaneous melanosis, 441e-37f
- Neurocysticercosis
 brain abscess in, 900
 clinical features of, 902-903, 1430-1431, 1431f
 diagnosis of, 903, 1431, 1431t
 etiology of, 902
 treatment of, 903, 1432
- Neurodegeneration with brain iron accumulation (NBIA), 2625-2626
- Neuroendocrine tumors (NETs)
 appendiceal, 561t, 563, 567
 bronchial, 560t, 563-564, 2339
 carcinoma of unknown primary and, 120e-2t, 120e-4
 classification of, 560, 561t
 gastrointestinal. *See* Gastrointestinal neuroendocrine tumors (GI-NETS)
 general characteristics of, 557, 558t
 genetic syndromes associated with, 562, 562t
 in MEN 1, 2336t, 2339
 ovarian, 564
 pancreatic. *See* Pancreatic neuroendocrine tumors (pNETs)
 pathology of, 557-558, 558t
 prognostic factors in, 561t
 testicular, 564
 thymic, 564, 2339
 well-differentiated, 560
- Neuroepithelioma, peripheral, 119e-4
- Neurofibrillary tangles, 76
- Neurofibroma, 364
- Neurofibromatosis type 1 (NF1)
 clinical features of, **2343**
 hypertension, 1620t
 pheochromocytoma, 2331-2332, 2331f
 primary CNS tumors, 599t, 603-604
 genetic considerations in, **603, 2343**
 chromosomal location, 101e-6t, 562t, 2337t
NF1 gene mutations, 101e-6t, 599t, 1620t, 2331
 imaging in, 441e-35f, 2331f
 NETs in, 562, 562t
 pheochromocytoma in, 2330-2331, 2331f
 skin manifestations of, **2331**
 axillary freckling, 2331f
 hyperpigmentation, 359, 360t
 Lisch nodules of the iris, 2331f
 neurofibromas, 76e-16f, 604, 2331f
 papulonodular lesions, 364
 soft tissue sarcomas in, 119e-1
- Neurofibromatosis type 2 (NF2)
 clinical features of, 220, 599t, 604, 2343
 CNS tumors in, 599t, 604
 genetic considerations in, 101e-6t, 599t, 604
 imaging in, 441e-36f
- Neurofibromin, 603, 2343
- Neurogenic ptosis, 208
- Neurogenic shock, 1748t, **1750**
- Neurohypophysis. *See* Posterior pituitary
- Neuroimaging, **440e-1**. *See also specific disorders*
 atlas of, 441e-1-43f
 CT, **440e-1**. *See also* Computed tomography (CT), in neurologic disease
 complications of, 440e-2, 440e-4, 440e-4t
 guidelines for, 440e-1t
 indications for, 440e-1t, 440e-2
 technique, 440e-1, 440e-2f
 interventional neuroradiology, 440e-11
 magnetic resonance angiography, 440e-6-8
 MRI, **440e-4**. *See also* Magnetic resonance imaging (MRI), in neurologic disease
 arterial spin labeling, 440e-8
 complications of, 440e-5-6
 contrast agents, 440e-5
 diffusion tensor, 440e-8, 440e-9f
 diffusion-weighted, 440e-3f, 440e-5f, 440e-8
 echo-planar, 440e-8
 FLAIR. *See* Fluid-attenuated inversion recovery (FLAIR) MRI
 functional, 90f, 440e-8
 guidelines for, 440e-1t
 magnetic resonance neurography, 440e-8
 perfusion, 440e-8
 technique, 440e-4-5, 440e-4t
 myelography, 440e-9-10
 positron emission tomography, 440e-8-9. *See also* Positron emission tomography (PET), in neurologic disease
 in spine interventions, 440e-10
 Neurokinin (NK₁), 258
 Neurokinin (NK₁) receptor antagonists, 103e-25
 Neuroleptic malignant syndrome (NMS), 2624, 2643
 Neuroleptics. *See also* Antipsychotics
 adverse effects of, 65, 326, 2624
 for delirium, 64, 64t
 for Tourette's syndrome, 2623
 Neurologic disease. *See also specific diseases*
 approach to the patient. *See* Neurologic examination/consultation
 in Behçet's syndrome, 2194
 in critically ill patient. *See* Critically ill patient, neurologic disorders
 depression in, 2714
 electrodiagnostic studies of. *See specific techniques and* Electrodiagnostic studies
 fatigue in, 152
 gene therapy for, 91e-2f
 genome-wide association studies of, 444e-1
 global burden of, 2535, 2535t
 in HIV infection. *See* HIV infection, neurologic disease in
 in hypothermia, 478e-2t
 hypoventilation in, 1721, 1721f
 imaging studies. *See* Neuroimaging
 insomnia in, 191
 involuntary weight loss in, 275
 as late consequence of cancer/cancer therapy, 622
 localization of, 2535, 2540t
 mechanisms of, **444e-1**
 cell death, 444e-8-9, 444e-8f
 channelopathies, 444e-2, 444e-2t
 genetic, 444e-1-2, 444e-2t
 gene transcription disorders, 444e-4
 microbiota, 444e-6-7
 myelin disorders, 444e-4-5, 444e-4f
 neurotransmitter/neurotransmitter receptor disorders, 444e-2-4, 444e-3t
 neurotrophic factors, 444e-7, 444e-7t
 paraneoplastic. *See* Paraneoplastic syndromes, neurologic
 prions, 444e-10-12, 444e-12t
 protein aggregation, 444e-9-10
 signaling pathway defects, 444e-4
 in megaloblastic anemia, 643
 nutritional deficiencies and, 463t
 in pregnancy, 49
 prevalence of, 2535
 research in, 444e-12-13
 in sarcoidosis, 2206t, 2208
 in SLE, 2128-2129, 2129t
 stem cell therapy in, 90e-3, 444e-7-8
 in syphilis. *See* Neurosyphilis
 testicular dysfunction in, 2366
 in Wilson's disease, 2519
 Neurologic examination/consultation, **463e-1, 2535**
 after cardiac bypass, 463e-2-3, 463e-2f
 after solid organ transplantation, 463e-3
 in back pain, 113, 114t
 in coma, 1773-1775, 1774f
 coordination examination, 2539
 cranial nerve examination, 2537-2538
 in delirium, 168
 detailed, 439e-1v
 in dizziness and vertigo, 149
 in electrolyte disturbances, 463e-3-4
 gait examination, **2539**
 in headache, 107
 in hyperperfusion states, 463e-1-2, 463e-1t, 463e-2f
 imaging studies. *See* Neuroimaging
 localization of findings in nervous system, 2539-2540, 2540t
 mental status examination, 2537
 motor examination, 2538
 in neck pain, 121t
 neurologic history in, 2535-2536
 neurologic method in, 2535
 in olfactory/taste dysfunction, 216
 overview of, 438e-1v, 439e-1v, 2536-2537
 in peripheral nervous system dysfunction, 463e-4-5, 463e-5f
 reflex examination, 2538-2539
 screening, 438e-1v
 sensory examination, **2539**
 Neurologic-myopathic syndromes, in lung cancer, 511
 Neuroma, 364t, 2340. *See also* Vestibular schwannoma (acoustic neuroma)
 Neuromuscular blocking agents, 462e-20t, 1732-1733, 1741, 1784
 Neuromuscular jitter, 442e-6
 Neuromuscular junction disorders, **2701**
 autonomic dysfunction in, 2642
 botulism. *See* Botulism
 clinical features of, 2540t
 congenital myasthenic syndrome, 2702, 2703t
 in critically ill patient, 1784
 Lambert-Eaton myasthenic syndrome. *See* Lambert-Eaton myasthenic syndrome (LEMS)
 myasthenia gravis. *See* Myasthenia gravis
 progressive external ophthalmoplegia. *See* Progressive external ophthalmoplegia
 weakness in, 155, 157
 Neuromyelitis optica (NMO), 2656, 2673, 2673t
 Neuromyotonia, 444e-2t, 462e-3, 619
 Neuronal nicotinic acetylcholine receptor, 372e-25t
 Neuron-specific enolase (NSE)
 as brain damage indicator, 1781
 in carcinoma of unknown primary, 120e-2t
 in nonfunctional pNETs, 571
 as tumor marker, 473t
 Neuro-ophthalmology, 41e-1v
 Neuropathic joint (Charcot's joint) disease, 1136, 2243-2244, 2243f, 2243t
 Neuropathic (lower motor neuron) weakness, 154, 154t, 157
 Neuropathy. *See also* Peripheral neuropathy;
 Polyneuropathy
 acute quadriplegic, 462e-20
 anti-Hu paraneoplastic, 2700
 diabetic. *See* Diabetic neuropathy
 entrapment, 49, 463e-4
 femoral, 2692
 gait disorders in, 163-164
 in HIV infection. *See* HIV infection, neurologic disease in
 immune-mediated, 2696t
 acute motor axonal neuropathy, 2695, 2695t
 acute motor sensory axonal neuropathy, 2695, 2695t
 anti-Hu paraneoplastic neuropathy, 2700
 chronic inflammatory demyelinating polyneuropathy, 2698-2699
 Guillain-Barré syndrome. *See* Guillain-Barré syndrome (GBS)
 Miller-Fisher syndrome, 2695, 2695t
 in monoclonal gammopathy of undetermined significance, 2700
 multifocal motor neuropathy, 2699
 in multiple myeloma, 2699-2700
 vasculitic neuropathy, 2700
 lateral femoral cutaneous, 49, 463e-5, 463e-5f, 2692
 median, 2683, 2691
 multifocal motor, 2699
 obstetric, 463e-5
 optic. *See* Optic neuropathy

- Neuropathy (*Cont.*):
 pain in, 60, 89–90, 94–95
 peroneal, 463e-4, 463e-5f, 2683, 2692
 proximal femoral, 463e-5, 463e-5f
 radial, 463e-4, 463e-5f, 2691
 sacral, 274
 sciatic, 2692
 sensory deficits in, 161
 trigeminal, 2646–2647
 ulnar, 463e-4, 463e-5f, 1124, 2683, 2691
 Neuropathy, ataxia, retinitis pigmentosa (NARP) syndrome, 85e-6, 85e-6t, 451e-3t
 Neuropeptide Y, 415e-2
 Neuroprotective agents, 2563, 2616
 Neurosarcoidosis, 40e-4f, 441e-11–15f, 909t, 2208. *See also* Sarcoidosis
 Neurostimulation therapy, 2596
 Neurosyphilis
 clinical features of, 105, 163, 898–899, 907t, 1135–1136, 2607
 in congenital syphilis, 1137
 diagnosis of, 1135–1136, 1138t
 in HIV infection, 1259
 imaging in, 441e-3–5f
 treatment of, 1139, 1139t
 vs. viral meningitis, 892
 Neurotensin, 559t
 Neurotransmitter(s), 265e-4, 444e-2–4, 444e-3t
 Neurotrophic factors, 444e-7, 444e-7t
 Neutral protamine Hagedorn (NPH) insulin, 2411, 2411t
 Neutral theory of community assembly, 86e-6
 Neutron (η) particles, 263e-1
 Neutropenia, 413, 417
 acute, 417–418
 in cancer patient, 103e-3t, 103e-23–25. *See also* Cancer, infections in
 chronic, 423
 cyclic, 126, 237, 418
 diagnosis of, 423
 drug-induced, 417–418, 418t
 enterocolitis associated with, 488, 1099, 1797–1798, 1798f
 etiology of, 418t
 febrile, 103e-23, 490, 490f, 1047
 hereditary, 418
 in HIV infection, 1262
 in HSCT recipient. *See* Hematopoietic stem cell transplantation (HSCT), complications of
 refractory, 669t. *See also* Myelodysplasia in sepsis/septic shock, 1756, 1757t
 severe congenital, 2104, 2105f
 skin manifestations of, 76e-12f
 in SLE, 2129t, 2131
 treatment of, 423
 in viral hepatitis, 2016
 Neutrophil(s), 413
 abnormalities of
 chronic granulomatous diseases. *See* Chronic granulomatous diseases (CGDs)
 granule disorders, 419, 420t
 inherited disorders, 420t
 leukocyte adhesion disorders, 419, 420t
 neutropenia. *See* Neutropenia
 phagocyte activation disorders, 420t, 421
 in adaptive immunity, 372e-7t
 apoptosis of, 416
 in asthma, 1673
 circulating pools of, 415–417, 416f
 hypersegmented, 81e-3f
 in innate immunity, 372e-7t, 372e-8, 372e-15t
 marginated, 415–416, 416f
 marrow release of, 415–417
 maturation of, 413–415, 414f
 morphology of, 414f
 in peripheral blood smear, 81e-1–2
 senescent, 416
 shift to the left, 413
 in shock, 1747
 travel through pulmonary capillaries, 417f
 Neutrophil-activating protein-2 (NAP-2), 372e-12t
 Neutrophil bands, 415, 415f
 Neutrophil chemotactic factor (NCF), 372e-15t
 Neutrophil count, 417, 2754t, 2755t
 Neutrophil extracellular traps (NETs), 1753
 Neutrophil gelatinase associated lipocalin (NGAL), 1808
 Neutrophil granules
 disorders of, 419, 419t
 primary (azurophil), 415
 secondary (specific), 415
 toxic granulations, 415
 Neutrophilia, 413, 418–419, 418t
 Nevirapine
 action of, 1278
 adverse effects of
 cutaneous, 362, 381, 1278
 fever, 138
 hepatotoxicity, 1258, 2030
 in HIV infection, 1274t
 SJS or TEN, 46e-1f, 383, 1278. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
 drug interactions of, 42t
 for HIV infection, 1274t, 1278
 molecular structure of, 1276f
 resistance to, 1278, 1279f
 Nevoid basal cell carcinoma syndrome. *See* Gorlin's syndrome (nevoid basal cell carcinoma syndrome)
 Nevomelanocytic nevus, 339f
 Nevus
 atypical, melanoma and, 493
 dysplastic, 76e-7f, 493–494, 493f
 melanocytic, 493–494
 nevomelanocytic, 76e-6f, 339f
 oral, 240t
 Nevus depigmentosus, 359
 Newborn(s). *See* Neonate(s)
 New daily persistent headache. *See* Headache, new daily persistent
 New York Heart Association (NYHA), classification of cardiovascular disease, 1439–1440, 1440t, 1501, 1501t, 1581t
 New York virus, 1308t
 Next-generation sequencing, 83e-3, 85e-4, 447, 769t
 NF1. *See* Neurofibromatosis type 1 (NF1)
 NF1 gene mutations
 action of, 603, 2343
 in CNS tumors, 599t, 603–604
 in juvenile myelomonocytic leukemia, 135e-6
 in melanoma, 496
 in neurofibromatosis 1, 2331
 in pheochromocytoma/paraganglioma, 2334, 2334f
 in pNETS, 562, 562t
 in soft tissue sarcomas, 119e-1
 tumors associated with, 101e-6t
 NF2. *See* Neurofibromatosis type 2 (NF2)
 NF2 gene mutations, 101e-6t, 599t, 604
 NF- κ B. *See* Nuclear factor- κ B (NF- κ B)
 NF- κ B essential modulator. *See* NEMO (NF- κ B essential modulator)
 NGAL (neutrophil gelatinase associated lipocalin), 1808
 Ngari virus, 1308t
 NGF (nerve growth factor), 444e-7t
 Niacin (vitamin B₃), 96e-4. *See also* Nicotinic acid
 deficiency of. *See* Pellagra
 functions of, 96e-2f, 96e-4
 metabolism of, 96e-4
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 structure of, 96e-2f
 toxicity of, 96e-4
 Nicardipine
 adverse effects of, 1589t, 2218t
 for hypertension in subarachnoid hemorrhage, 1787
 for hypertensive emergencies, 1627, 1627t
 for ischemic heart disease, 1589, 1589t
 Niche site, stem cells, 89e-2
 Nickel allergy, 76e-2f, 345
 Nickel exposure, 472e-3, 477t, 502, 506
 Niclosamide, 246e-5t, 246e-11
 Nicoladoni-Branham sign, 1647
 Nicorandil, 1590
 Nicotinamide adenine dinucleotide phosphate (NADPH) oxidase, 416
 Nicotine, 444e-4, 465e-5t, 473e-10t, 2729. *See also* Smoking
 Nicotine replacement therapy, 507, 1705
 Nicotinic acid. *See also* Niacin (vitamin B₃)
 adverse effects of, 462e-19, 462e-20t, 2447, 2448t
 for lipoprotein disorders, 291e-6, 2447, 2448t
 for metabolic syndrome, 2454
 Nicotinic agonists, 473e-10t
 NIDDM (non–insulin-dependent diabetes mellitus). *See* Diabetes mellitus (DM), type 2
 Niemann-Pick disease, 432e-3t, 432e-5, 436e-7f, 451e-4t
 Nifedipine
 adverse effects of, 237, 253, 1589t
 for altitude illness, 476e-2t, 476e-4
 for chilblain, 478e-5
 for ciguatera poisoning, 2742
 dosage in liver disease, 37
 drug interactions of, 246e-6t
 for ergot alkaloid overdose/poisoning, 473e-8t
 for hypertension, 1624t, 1625, 2644
 for hypertension in pregnancy, 46
 for ischemic heart disease, 1589, 1589t
 overdose/poisoning with, 473e-9t
 for Raynaud's phenomenon, 1649, 2168f
 Nifurtimox
 action of, 246e-11
 adverse effects of, 246e-5t, 246e-11, 1396, 1560
 for Chagas disease, 1266, 1395–1396, 1560
 pharmacology of, 246e-11
 Night blindness, 96e-1t, 96e-6, 1946t
 Nightshade, 473e-9t
 Nijmegen breakage syndrome, 375e-1, 2109
 Nikolsky's sign, 134, 370, 959, 960f
 Nil lesion. *See* Minimal change disease
 Nilotinib
 action and targets of, 102e-3, 102e-5t, 103e-16, 103e-17t
 adverse effects of, 103e-17t, 692
 for CML, 691–692, 691t
 genetic variations in response to, 39t, 102e-16
 Nilutamide, 585
 Nimesulide, 338
 Nimodipine, 1787, 2597
 Nintedanib, 2164
 Nipah virus, 894
Nippostrongylus brasiliensis infection, 372e-8
 NIS (sodium iodide symporter), 372e-24t, 2284, 2284t
 NIS gene mutations, 2284
 Nisoldipine, 1589t
 Nissen fundoplication, 1908
 Nit(s), 2745–2746
 Nitazoxanide
 action of, 246e-11
 adverse effects of, 246e-5t
 for *C. difficile* infection, 860t, 861
 for *Cryptosporidium* infections, 1407
 for *Giardia* infections, 1406
 for *H. nana* infections, 1434
 pharmacology of, 246e-11
 in pregnancy and lactation, 246e-5t
 Nitisinone, 434e-5
 Nitrates
 action of, 1588
 adverse effects of, 1603
 dietary
 esophageal cancer and, 532
 gastrointestinal cancer and, 534, 534t
 for heart failure, 1509, 1608
 interaction with phosphodiesterase type 5 inhibitors, 43, 328, 1596, 1603
 for ischemic heart disease, 1588, 1588t
 long-acting, 1588

- for myocardial infarction, 1595–1596, 1597t, 1603
 overdose/poisoning with, 473e-11t
 for Prinzmetal's variant angina, 1598
 for pulmonary edema, 1763
 tolerance to, 1588
- Nitric oxide
 in asthma, 1674, 1676
 in blood vessels, 265e-2, 265e-4
 in cirrhosis, 286
 in clitoral engorgement, 330
 in erection, 324, 325f
 in gastric mucosal integrity, 1912
 in shock, 1745, 1747
 in swallowing, 255
- Nitric oxide synthase. *See* Endothelial nitric oxide synthase (eNOS)
- Nitric oxide therapy, 1739, 1739t, 1743
- Nitrites, 473e-11t, 637
- Nitroblue tetrazolium dye test, 423
- Nitrofurantoin
 action of, 931t, 932f, 933, 944
 adverse effects of, 938t, 944
 common, 867t
 eosinophilia, 422, 1687
 folate deficiency, 646t
 in G6PD deficiency, 656t
 hepatotoxicity, 2028
 neuropathy, 2687t
 for cystitis, 866, 867, 867t
 indications for, 941t, 944
 in pregnancy and lactation, 937t
 prophylactic, 945t
 resistance to, 931t, 934, 941t, 944
 for urinary tract infections, 975t, 1029
- Nitrogen dioxide exposure, 1692t
- Nitrogen mustard. *See* Mechlorethamine (nitrogen mustard)
- Nitrogen oxides, as chemical terrorism agents, 262e-1t
- Nitroglycerin
 adverse effects of, 1588, 1603
 drug interactions of, 43
 for ergot alkaloid overdose/poisoning, 473e-8t
 for heart failure, 1510t, 1608
 for hypertensive emergencies, 1627t
 for hypothermia, 478e-3
 intravenous, 1607
 for ischemic heart disease, 1588, 1588t
 metabolism of, 34
 for myocardial infarction, 1603, 1607
 for Prinzmetal's variant angina, 1598
 for pulmonary edema, 1761f, 1763
- Nitroimidazoles, 205e-7
- Nitroprusside
 for adrenergic crisis, 1627
 for aortic dissection, 1641
 for ergot alkaloid overdose/poisoning, 473e-8t
 for heart failure, 1510t
 for hypertensive emergencies, 1626t, 1627
 for pulmonary edema, 1763
 therapeutic monitoring of, 2765t
- Nitroprusside ketone reaction, 318
- Nitrosoureas, 103e-8, 621, 1797
- Nitrous oxide
 adverse effects of, 635
 as greenhouse gas, 151e-2t, 151e-3f
 inhalation of, cobalamin deficiency in, 646
 for sickle cell painful crisis, 635
- NIV. *See* Noninvasive ventilation (NIV)
- Nivolumab, 103e-21, 521
- Nixon's method, splenic percussion, 411
- Nizatidine, 263, 1920, 1920t
- Njovera. *See* Endemic syphilis
- NK cells. *See* Natural killer (NK) cells
- NK T cells. *See* Natural killer (NK) T cells
- NLM (National Library of Medicine), 14e-4t
- NLRC4/IPAF inflammasome, 145e-8f
- NLRP1 gene, 372e-8t
- NLRP1/NALP1b inflammasome, 145e-8f
- NLRP2/NALP3 inflammasome, 145e-8f
- NLRP3 gene, 372e-8t, 2215
- NLRs (NOD-like receptors), 145e-7, 145e-8f, 372e-2, 372e-5t
- NLST (National Lung Cancer Screening Trial), 509–510, 510t
- NMDA receptor antibodies, 444e-9, 614t, 615f, 617, 2607
- NMS (neuroleptic malignant syndrome), 2624, 2643
- NMYC gene mutations, 101e-2t, 101e-3
- NO-1, 167e-2
- Nocardia* spp., 1084, 1084t, 1086
- Nocardia* spp. infections, **1084**
 actinomycetoma, 1085–1086, 1086f, 1087t
 antimicrobial resistance in, 1084t
 in cancer patient, 488, 488t
 chronic meningitis, 907t
 clinical features of, 308e-6f, 1084–1086, 1085f
 diagnosis of, 150e-1, 150e-2t, 1086–1087, 1086f
 epidemiology of, 1084
 in HIV infection, 1084
 pathogenesis of, 1084
 in transplant recipient, 926
 treatment of, 1084t, 1087–1088, 1087t
- Nocebo effect, 89
- Nociceptin/orphanin receptors, 468e-1t
- Nociceptive pain, 60
- Nociceptor(s), 87
- Nociceptor(s), primary afferent, 87f, 88f, 89f
- Nociceptor-induced inflammation, 88, 88f
- Nocturia, 1872
- NOD2 gene mutations, 1949, 1950t, 2215
- Nodal marginal zone B-cell lymphoma, 135e-2, 707
- NOD-like receptors (NLRs), 145e-7, 145e-8f, 372e-2, 372e-5t
- NOD proteins, 145e-7, 372e-6t, 1752
- Nodular vasculitis, 366
- Nodule, 127, 339t, 340f. *See also* Papulonodular skin lesions
- Nogo, 444e-8
- Noise exposure, 224–225, 225t
- Noma (cancrum oris), 236, 487, 1097
- NOMID (neonatal-onset multisystem inflammatory disease), 2213t, 2215
- Non-24 hour sleep-wake rhythm disorder, 193
- Nonaka's distal myopathy, 462e-12, 462e-13t
- Nonalcoholic fatty liver disease (NAFLD), **2054**
 cirrhosis due to, 2060
 clinical features of, 2056
 diagnosis of, 2055–2056, 2056t
 epidemiology of, 2054
 etiology of, 2054, 2055t, 2056t
 global considerations in, 2057
 metabolic syndrome and, 2452
 obesity and, 415e-6
 pathogenesis of, 2055
 treatment of, 2056–2057
- Nonalcoholic steatohepatitis (NASH)
 cirrhosis due to, 2060
 diagnosis of, 366e-5f, 1993t
 etiology of, 2055t, 2056t
 genetic considerations in, 2518
 hepatocellular carcinoma and, 545t
 natural history of, 2054
 pathogenesis of, 2055
- Nonallelic (locus) heterogeneity, 435–436, 436t
- Nonbacterial thrombotic endocarditis (NBTE), 817
- Noncompliance, 36
- Nondisclosure of information, 17e-1
- Nonexercise activity thermogenesis, 415e-5
- Nonhistone proteins, 427f
- Non-Hodgkin's lymphoma. *See also* Lymphoid malignancies
 approach to the patient, 700–701
 classification of, 695, 696t
 clinical features of, 702t
 epidemiology of, 697
 etiology of, 697, 697t
 genetic considerations in, 101e-3t
 HSCT for, 139e-5t, 139e-6
 HTLV-1 and, 697, 707
 infections in, 484, 484t, 486t
- International Prognostic Index for, 700–701, 701f, 701t
 late consequences of treatment of, 623, 624t
 during pregnancy, 124e-4
 primary central nervous system. *See* Primary central nervous system lymphoma (PCNSL)
 renal involvement in, 1859
 rituximab for, 372e-30
 skin manifestations of, 76e-5f, 366, 372, 501f
 splenomegaly in, 411
 staging of, 700t, 702t
 thiopurine-associated, 1962
- Nonhuman primates, bite-wounds from, 167e-2, 167e-3t
- Non-insulin-dependent diabetes mellitus (NIDDM). *See* Diabetes mellitus, type 2
- Noninvasive ventilation (NIV)
 advantages and disadvantages of, 1742t
 contraindications for, 1707, 1741t
 for COPD, 1707, 1740
 for hypoventilation, 1722
 for pulmonary edema, 1763
 variables in, 1742t
- Nonocclusive mesenteric ischemia, 1979–1980, 1980t
- Nonoliguria, 293
- Nonoxynol 9, 863, 1009
- Nonreactive military tuberculosis, 1112
- Nonseminoma
 mediastinal, 591
 testicular. *See* Testicular cancer
- Nonsense mutation, 433f
- Nonsteroidal anti-inflammatory drugs (NSAIDs)
 action of, 1912
 adverse effects of
 bleeding, 404
 colitis, vs. IBD, 1957
 cross-sensitivity and, 385
 cutaneous, 360, 361, 377, 378, 381
 diarrhea, 267
 edema, 252, 253t
 gastrointestinal bleeding, 277, 2232, 2232t
 gastrointestinal ulcers, 262. *See also* Peptic ulcer disease, NSAID-induced
 hyperkalemia, 64e-2, 310
 hypertension, 1617t
 immunologic factors in, 377
 in ischemic heart disease, 1590
 in myocardial infarction, 1604
 nail disorders, 380
 proximal ileal stricture, 1884f
 renal, 90–91, 291, 1800, 1801f, 1821, 1857, 2232
 SJS or TEN, 383. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
 in SLE, 2130t, 2132
 vasculitis, 382
 for ankylosing spondylitis, 2172
 for back pain, 119
 for calcium pyrophosphate deposition disease, 2236
 for colorectal cancer prevention, 539
 for dementia prevention, 2602
 drug interactions of, 43, 103e-11t, 103e-14, 2130t
 for dysmenorrhea, 338–339
 for episcleritis and scleritis, 200
 for fever, 126
 for fever of unknown origin, 141
 gastric protection with, 1923–1924, 1924t
 for gout, 2234
 for menorrhagia in perimenopause, 2382
 for migraine, 2590, 2592t
 for nephrogenic diabetes insipidus, 304
 for osteoarthritis, 2231–2232, 2232t
 for pain, 60, 90–91, 91t
 for phototoxicity, 390
 for polyposis syndromes, 539
 for reactive arthritis, 2175
 for rheumatic fever, 2152–2153
 for rheumatoid arthritis, 2145

- Nonsteroidal anti-inflammatory drugs (NSAIDs)
(*Cont.*):
for rhinovirus infections, 1203
for sickle cell arthropathy, 635
for SLE, 2130t, 2132
for spinal stenosis, 116
for subacute thyroiditis, 2299
for sunburn, 387
for tension-type headache, 2594
topical, 199, 2232, 2232t
for upper respiratory infections, 226
- Non-ST-segment elevation acute coronary syndrome (NSTE-ACS), **1593**
chest pain in, 96, 96t
clinical features of, 1594
diagnosis of, 1594–1595, 1594f, 1595f
ECG in, 297e-4f, 1457f, 1594
incidence of, 1593, 1594f
pathophysiology of, 1593–1594, 1594f
prognosis of, 1596t
risk stratification in, 1595, 1596f
treatment of, **1595**
ACE inhibitors, 1596
antithrombotic therapy, 747, 1596–1597, 1598t
beta blockers, 1596, 1597t
calcium channel blockers, 1596, 1597t
invasive vs. conservative strategy, 1597, 1598t
long-term management, 1597–1598
monitoring, 1595
morphine sulfate, 1597t
nitrates, 1595–1596, 1597t
PCI, 297e-3–4, 297e-3–4v, 1597. *See also* Percutaneous coronary interventions (PCI)
statins, 1596
- Nontropical sprue. *See* Celiac disease (gluten-induced enteropathy)
- Nontuberculous mycobacterial (NTM) infections, **1128**
bronchiectasis, 1128, 1694, 1696. *See also* Bronchiectasis
in children, 1130
diagnosis of, 205e-7–8, 1130–1131, 1131f
disseminated disease, 1129–1130
epidemiology of, 205e-7, 1128–1129
global considerations in, 1128–1129, 1132
lymphadenopathy, 1130
M. avium complex. *See* *Mycobacterium avium* complex (MAC) infections
M. marinum. *See* *Mycobacterium marinum* infections
pathobiology of, 1129, 1129f
prevention of, 1131
prognosis of, 1132
pulmonary disease, 1130
skin and soft tissue, 1130
treatment of, 205e-3–4, 205e-8, 1131–1132
- Noonan's syndrome, 436t, 1654
- Norepinephrine
action of, 444e-3t, 2637
in cardiac contraction, 265e-6
for cardiogenic shock, 1760–1761, 1761f
in cardiovascular regulation, 1613
clinical aspects of, 444e-3t
for neurogenic shock, 1750
for pulmonary edema, 1761f
reference values for, 2762t
for shock, 1750, 1751, 1757
structure of, 444e-3t
in vascular smooth-muscle cells, 265e-4
- Norethindrone, 2382
- Norfloxacin
action of, 931t, 932–933
for bacterial peritonitis prophylaxis, 288
for donovanosis, 198e-2
for infectious diarrhea, 857t
for peritonitis prophylaxis, 847
resistance to, 931t
- Norgestimate, 334
- Normal-pressure hydrocephalus, 175f, 2606–2607
- Normal saline. *See* Saline
- Normative economics, 15e-3
- Normeperidine, 92
- Normochromia, 81e-1
- Noroviruses, 1285, 1286f
- Norovirus (Norwalk virus) infections
clinical features of, 258, 266, 266t, 1286, 1286t
diagnosis of, 1286–1017
epidemiology of, 266, 855, 855t, 1285–1286
etiology of, 1285, 1286t
health care–associated, 917
immunity to, 1286
pathogenesis of, 1286
prevention of, 917, 1287
treatment of, 1287
- Norrie's disease, 222t
- North American Chest Pain Rule, 102f
- North Queensland tick typhus, 128t
- Nortriptyline
adverse effects of, 2710t
for chronic daily headache, 109
for depression, 2672, 2710t
for migraine prevention, 2594t
for nausea and vomiting, 260t
for neuropathic pain, 61, 93
for pain, 91t, 94, 2672
for smoking cessation therapy, 507, 2732
therapeutic monitoring of, 2765t
- Norwegian (crusted) scabies, 1263, 2744
- Nose
in granulomatosis with polyangiitis, 1443, 2183
in lepromatous leprosy, 1126
in relapsing polychondritis, 1443, 2203, 2203f
swab, specimen collection and transport, 150e-4t
- No-see-um midges, 2749
- Nosema* spp., 1408
- Nosocomial infections. *See* Health care–associated infections
- NO SPECS mnemonic, in Graves' disease, 2295
- NOTCH1* gene mutations, 1529
- NOTCH2* gene mutations, 135e-1
- NOTCH3* gene mutations, 2568
- NOTCH4* gene mutations, 2155
- NOTES (natural orifice transluminal endoscopic) surgery, 346e-1v, 1881, 1883, 1886f
- Nothnagel's syndrome, 208
- NPH insulin, 2411, 2411t
- NPHP2* gene mutations, 1855
- NPHS1/2* gene mutations, 1832
- NPM1* gene mutations, 680, 680t
- NRAMP1, 1106
- NRAS* gene mutations
cancers associated with, 101e-2t
in chronic myelomonocytic leukemia, 135e-6
in lung cancer, 507t, 509f
in melanoma, 496
- NREM sleep, 184–185, 185f
- NRG1* gene mutations, 465e-4
- NRL* gene mutations, 431t
- NRXN1* gene mutations, 465e-2
- NSAIDs. *See* Nonsteroidal anti-inflammatory drugs (NSAIDs)
- NSE. *See* Neuron-specific enolase (NSE)
- NSTE-ACS. *See* Non-ST-segment elevation acute coronary syndrome (NSTE-ACS)
- N-terminal pro-BNP (NT-proBNP), 1505, 2762t
- NTM infections. *See* Nontuberculous mycobacterial (NTM) infections
- NTS (nucleus of the solitary tract), 213–214
- Nuclear factor- κ B (NF- κ B)
activation defects of, 420t
in asthma, 1674
essential modulator. *See* NEMO (NF- κ B essential modulator)
increased levels of, 102e-10
inhibition of, 102e-6, 102e-9f
nuclear translocation of, 145e-5, 145e-6f
receptor activator of. *See* RANK (receptor activator of NF κ B)
systems biology analysis of, 87e-2
- Nuclear membrane, inherited defects of, 1555t
- Nuclear-mitochondrial genomic cross-talk, 85e-7, 85e-7f
- Nuclear receptor(s), 400e-3, 400e-4, 400e-4f, 431t
- Nuclear receptor family, 400e-1
- Nuclear reprogramming, 454
- Nuclear transfer, 85e-9, 85e-10f
- Nuclear transplantation, 454
- Nuclear weapons, 263e-1, 263e-2
- Nucleic acid sequence–based amplification (NASBA), 1247, 1247t
- Nucleic acid tests
application of, 150e-8
nucleic acid amplification, 214e-9, 769t, 1247, 1247t
nucleic acid amplification strategies, 150e-7
probes for direct detection of pathogens, 150e-6–7
quantitative strategies, 150e-7–8
- Nucleocapsid, viral, 214e-1
- Nucleoside reverse transcriptase inhibitors. *See* Combination antiretroviral therapy (cART)
- Nucleosome, 427f, 431f
- 5'-Nucleotidase, in liver function evaluation, 1997
- Nucleotide repeat expansion disorders, 440, 440t
- Nucleotide reverse transcriptase inhibitors. *See* Combination antiretroviral therapy (cART)
- Nucleus of the solitary tract (NTS), 213–214, 213f
- Nucleus pulposus, herniated, 105
- Number needed to treat, 26
- Numbness, 158. *See also* Sensation
- Nummular eczema, 346
- Nummular headache, 2598
- Nummular skin lesions, 340t
- Nuocytes, 2116
- Nutrient(s)
definition of, 95e-1
digestion and absorption of, **1933**. *See also* *specific nutrients*
bile acids in, 1934–1935, 1934f, 1934t
gut microbiota and, 86e-6, 1875
in intestinal epithelia, 1933–1934
- Nutrient requirements, **95e-1**
in critically ill patient, 1734
factors altering, 95e-4–5
recommended intakes, **95e-1**
- Nutrient-sensing pathways, 94e-4–5
- Nutritional assessment, 95e-5, **461**
in acute care setting, 95e-5
in ambulatory setting, 95e-5–6
anthropometrics, 462
bedside, 462
of cancer patient, 474
in elderly, 81
global considerations in, 95e-6
in high-risk patient, 95e-6, 462t
laboratory studies in, 462, 464t
nutritional history in, 462
physical examination in, 462
for specialized nutritional support, 98e-1–2
- Nutritional monitoring, 95e-5
- Nutritional Screening Initiative, 276
- Nutritional short stature, 2258
- Nutritional supplements, 14e-3–4, 14e-4t, 95e-3
- Nutritional support/therapy
after bariatric surgery, 2398
for diabetes mellitus, 2408–2409, 2409t
for gastrointestinal disease, 1879
for hypertension, 1622t, 1623
for IBD, 2147
for irritable bowel syndrome, 1968–1969, 1968t
for metabolic syndrome, 2453
for MS, 2672
for myocardial infarction, 1606
for obesity, 2394–2396
for sepsis/septic shock, 1757
- Nutrition transition, 461
- Nyando virus, 1309t

- NYHA (New York Heart Association), classification of cardiovascular disease, 1439–1440, 1440t, 1501t, 1581t
- Nylon flock exposure, 1692
- Nystagmus, 41e-1v, **210**
congenital, 210
etiology of, 210
gaze-evoked, 210
jerk, 210
patterns of, 149, 149t
rotary, in stimulant overdose, 473e-3
vestibular, 210
- Nystatin
for burn-wound infection, 166e-3
for *Candida* infections, 350t, 351, 1332, 1344t
for thrush, 233, 1344t
- O**
- OA. *See* Osteoarthritis (OA)
- OATB1B1 (organic anion transport protein 1B1), 38, 2000
- OATs (organic anion transporters), 431e-1, 431e-2f
- Obesity, **2392**
body mass index in, 98e-1t, 2392
breast cancer risk and, 524
cancer and, 415e-6, 476
central adiposity in, 6e-4, 2449
comorbid conditions, 2392, 2394t
asthma, 1670–1671
bone and joint disorders, 415e-6, 2229
cardiovascular disease, 266e-4, 290e-1t, 290e-2, 291e-9, 415e-6, 1563, 1587
cutaneous disorders, 415e-6
diabetes mellitus, 82, 415e-6, 2401, 2401t, 2404–2405
erectile dysfunction, 326
gallstones, 415e-6
hypertension, 415e-6, 1612, 1622
hypogonadotropic hypogonadism, 2364–2365
hypothalamic disorders, 2264
lipoprotein disorders, 2438, 2439
liver disease, 415e-6, 2054, 2055, 2057
pulmonary disease, 415e-6
reproductive disorders, 415e-6
sleep apnea and, 415e-6
definition of, 98e-1t, 415e-1
drug-induced, 2392
dyspnea in, 47e-2
etiology of, **415e-3**
craniopharyngioma in, 415e-4
Cushing's syndrome in, 415e-4
energy expenditure in, 415e-5
food intake in, 415e-5
genetic syndromes, 415e-3–4, 415e-4t
genetic vs. environmental factors in, 415e-3
hypothalamic disorders in, 415e-4
hypothyroidism in, 415e-4
insulinoma in, 415e-4
leptin in, 415e-4t, 415e-5
evaluation of, 415e-1, 415e-1f, 2392–2393, 2393t, 2394t
fatigue and, 152
global burden of, 17, 266e-4
gut microbiota and, 86e-8–9
high-altitude travel and, 476e-5
history focused on, 2392
in metabolic syndrome, 2450, 2450t, 2453
oral contraceptives use in, 2391
physical fitness and, 2392
physiology of, 415e-2, 415e-2f
pregnancy and, 48
prevalence of, 415e-1–2, 2254t
pulmonary function values in, 306e-5, 306e-5f
screening/testing for, 29t, 2254t
treatment of, **2393**
bariatric surgery, 2398, 2398f
behavioral therapy, 2396
diet therapy, 2394–2396
exercise program, 2396
goal of, 2393–2394
in metabolic syndrome, 2453
pharmacotherapy, 2396–2397
readiness to change and, 2392–2393
selection of, 2395f, 2395t
upper body, 6e-4
in women, 6e-4
- Obesity hypoventilation syndrome, 415e-6, 1722
ob gene, 415e-4, 415e-5
- Obinutuzumab, 103e-21, 103e-21t, 703
- Object recognition, 182
- Observational data, 25
- Obsessive-compulsive disorder (OCD), **2713**
- Obstetric neuropathies, 463e-5
- Obstructive sleep apnea/hypopnea syndrome (OSAHS)
cardiovascular effects of, 1726
clinical features of, 1724–1725
daytime sleepiness in, 188, 1723, 1726
definition of, 1723
diagnosis of, 188t, 1721, 1721t, 1725, 1725t
epidemiology of, 1724
fatigue and, 152
in heart failure, 1514
hypertension and, 1619, 1725
in metabolic syndrome, 2452
obesity and, 415e-6, 1620, 1724
pathophysiology of, 306e-5, 1723–1724, 1723f
polysomnography in, 1725, 1726f
pulmonary hypertension in, 1657
quality of life in, 1726
risk factors for, 1724
severity scale for, 1725t
treatment of, 188t, 1620t, 1727, 1727t
- Occam's razor, 19
- Occipital lobe stroke, 204
- Occipital nerve stimulation, 2596
- Occipitotemporal network, 182
- Occupational illness. *See also* Health care workers
animal bite-wounds, 167e-1–2
asthma, 1670, 1671
contact dermatitis, 345
hand eczema, 346
hearing loss, 224–225, 225t
lung disease, 1662
asbestos-related. *See* Asbestos-related disease
categories of, 1688t
characteristics of agents causing, 1688
chest imaging in, 1687–1688
chronic beryllium disease, 1691
coal worker's pneumoconiosis, 1690
COPD, 1703
disability assessment in, 1693
exposure assessment in, 1687
history in, 1687
hypersensitivity pneumonitis. *See* Hypersensitivity pneumonitis
incidence of, 1687
inorganic dusts, 1688t, 1691
organic dusts, 1688t, 1691
pulmonary function tests in, 1687
silicosis, 1689–1690, 1690f
toxic chemicals, 1691–1692, 1692t
in World Trade Center rescuers, 1692
osteoarthritis, 2229
shift-work disorder, 194
- OCD (obsessive-compulsive disorder), **2713**
- OCG (oral cholecystography), 2078
- Ochrobactrum* spp., 183e-3, 194e-1
- Ochronosis, 434e-4
- Ochronotic arthritis, 434e-4
- Ochronotic pigmentation, 436e-7f
- Ocrelizumab, 2672
- OCT (optical coherence tomography), 1668
- Octopus envenomation, 2740
- Ocreotide
for acromegaly, 2271
adverse effects of, 567, 574, 2271
for carcinoid syndrome, 272, 565, 567
for chemotherapy-related diarrhea, 103e-16, 103e-25
for dumping syndrome, 1926
in end-of-life care, 61
for gastrinoma, 2337
for glucagonoma, 570
for hepatorenal syndrome, 1810
for hypomotility of small bowel, 2164
for insulinoma, 569, 2434
for intestinal pseudoobstruction, 260t, 261
for NETs, 574
for oncogenic osteomalacia, 612
for postprandial orthostatic hypotension, 2644
radiolabeled, 575
for somatostatinoma, 570
structure of, 566f
for thymoma, 123e-2
for thyroid dermopathy, 2298
for variceal bleeding, 277, 2064
for vascular ectasias, 277
for VIPoma, 571
for ZES, 1929
- Ocular alignment, 197
- Ocular disorders, **195**. *See also* Optic neuritis; Visual loss
acne rosacea and, 352
in ankylosing spondylitis, 2171
in Behçet's syndrome, 200, 2194
in cat-scratch disease, 1079, 1081f, 1081t
in diabetes mellitus. *See* Diabetes mellitus (DM), complications of; Diabetic retinopathy
in granulomatosis with polyangiitis, 2184
at high altitudes, 476e-4
in IBD, 1958
infections. *See also* Conjunctivitis; Keratitis; Retinitis
Candida, 1343, 1343f, 1345
gonococcal, 1006, 1009t
L. loa. *See* *Loa loa* (African eye worm)/ loiasis
Nocardia, 1086
Onchocerca, 1421
P. aeruginosa, 1045t, 1046
T. gondii, 1402, 1404
Toxocara, 1411
in lepromatous leprosy, 1125
in Marfan syndrome, 2513
in MS, 2663
in myasthenia gravis, 2701–2702
in osteogenesis imperfecta, 2508
in relapsing fever, 1147
in relapsing polychondritis, 2203
in sarcoidosis, 2207, 2656
in Sjögren's syndrome, 2166
SLE and, 2129t, 2131
trachoma, 1172
in tularemia, 1068
- Ocular flutter, 41e-1v, 211
- Ocular lubricants, 199
- Ocular movements, 41e-1v, 195
assessment of, 197
“bobbing,” 1775
in coma, 1774–1775, 1774f
“dipping,” 41e-1v, 1775
in dizziness evaluation, 149
- Oculocephalic reflexes, 1775, 2604
- Oculomotor ataxia, 181
- Oculomotor nerve, 2537
- Oculomotor nerve palsy, 195, 208
- Oculopharyngeal dystrophy, 208, 462e-5t, 462e-11
- Odanacatib, 2503
- Odontoid fracture, 441e-33f
- Odynophagia, 254, 262, 533, 1901, 1909
- Ofatumumab, 103e-21, 103e-21t, 703
- Office of Research on Women's Health (NIH), 6e-1
- Ofloxacin
for *C. trachomatis* infection, 1171
for enteric fever, 1052
for leprosy, 1127
for pelvic inflammatory disease, 878
for plague, 1075
- Ogilvie's disease. *See* Intestinal pseudoobstruction (Ogilvie's disease)
- Oka. *See* Varicella vaccine
- OKT3, 253t, 372e-30, 2071

- Okuda staging, of hepatocellular carcinoma, 546, 546t
- Olanzapine
for acute alcohol intoxication, 2727
adverse effects of, 2721, 2722t
for bipolar disorder, 2718
for delirium, 64, 64t, 65
for nausea and vomiting, 61, 261
overdosage/poisoning with, 473e-8t
for schizophrenia, 2721, 2722t
- Olecranon bursitis, 2247
- Olfactory bulb, 211, 212f
- Olfactory disorders, 214
diseases and conditions associated with, 214–215, 214t
epidemiology of, 214, 214f
etiology of, 214, 214t
evaluation of, 215–216
in gonadotropin deficiency, 2363, 2364t
head injury and, 457e-2
taste disorders and, 215
treatment of, 216–217
- Olfactory nerve, 2537
- Olfactory system, 211–212, 212f
- Olig 2* gene, 444e-5
- Oligoastrocytomas, 601
- Oligodendrocyte myelin glycoprotein (OMGP), 444e-8
- Oligodendroglioma, 600–601
- Oligomenorrhea, 335, 337f, 2355
- Oligomer(s), 444e-9
- Oligopotent stem cells, 454, 454f
- Oliguria, 292, 1729, 1755
- Olivopontocerebellar atrophy. *See* Spinocerebellar ataxia (SCA), SCA1
- Ollier's disease (enchondromatosis), 426e-7, 594
- Olmesartan, 269
- Olsalazine, 1959, 1959t
- Omacetaxine, 691, 691t
- Omalizumab
adverse effects of, 372e-30, 377
for allergic rhinitis, 372e-30, 2124
for asthma, 1678, 1680
for urticaria, 2119, 2120
- Omapatrilat, 1513
- Omecamtiv mecarbil, 1510, 1510t
- Omega-3 fatty acids (fish oils)
adverse effects of, 2447, 2448t
antiplatelet effects of, 404
in enteral nutrition formulas, 98e-5
for heart failure, 1513
for hypertriglyceridemia, 2447, 2448t
for metabolic syndrome, 2454
- Omenn syndrome, 2108–2109
- Omentectomy, 593
- Omeprazole
action of, 1920
formulations of, 1920
for gastrinoma, 2337
genetic variations in response to, 39t, 40
for GERD, 263
in *H. pylori* eradication, 1042t, 1922, 1922t
for high-altitude cough, 476e-4
metabolism of, 34t, 40
for peptic ulcer disease, 1920–1921, 1920t
for ZES, 1929
- OMGP (oligodendrocyte myelin glycoprotein), 444e-8
- OmpA protein, 1036
- Omsk hemorrhagic fever virus, 1307t, 1322
- OMVs (outer-membrane vesicles), 1002, 1002f
- Onchocerca volvulus*/onchocerciasis
characteristics of parasite, 1418t, 1420
clinical features of
lymph nodes, 1421
ocular, 1421
onchocercosoma, 1420–1421
skin, 357, 357t, 829, 1420, 1421f
systemic, 1421
diagnosis of, 245e-3t, 245e-6t, 765t, 1421
epidemiology of, 765t, 1418t
etiology of, 1420
pathogenesis of, 1420
prevention of, 1421
treatment of, 246e-9, 1421
- Oncocytic tumors, renal, 578, 578t
- Oncogene(s)
activation of, 101e-2–3, 101e-7f
definition of, 101e-1–2, 102e-1
in human cancers, 101e-1–2, 101e-2t. *See also* Cancer genetics
in lymphoid malignancies, 699t
signaling pathways, 102e-11f
- Oncogene addiction, 102e-6, 102e-7
- Oncogene checkpoint, 102e-2f
- Oncologic emergencies. *See* Cancer, emergencies in
- Oncology, 103e-2
- Oncolytic viruses, 91e-4
- Oncostatin M (OSM), 372e-11t
- Ondansetron
for ataxia/tremor, 2672
for carcinoid syndrome, 566
for ciguatera poisoning, 2742
for nausea and vomiting, 62, 103e-25, 109, 259, 260t, 474
for scombroid poisoning, 2744
- Ondine's curse, 1722
- One-and-a-half syndrome, 210
- "Onion skinning," 1865
- Online Mendelian Inheritance in Man*, 425, 426t
- Onycholysis, 348, 351, 380
- Onychomadesis, 380
- Onychomycosis, 350, 1343, 1357, 1358t
- O'nyong-nyong virus infection, 837, 1306t, 1313
- Oocyte donation, 85e-9, 85e-10f, 337
- Oogonia, 2375, 2375f
- Oophorectomy, 103e-3, 530t, 592, 594
- Opa protein, 1003
- OPC-31260, 1853
- OPC-67683 (delamid), 205e-7
- Open-access endoscopy, 1900
- Operational taxonomic units (OTUs), 86e-2
- OPG. *See* Osteoprotegerin (OPG)
- Ophthalmia neonatorum, 1006, 1009t
- Ophthalmomyiasis, 2747
- Ophthalmopathy
diabetic. *See* Diabetic retinopathy
Graves'. *See* Graves' ophthalmopathy
thyroid-associated, 2294
- Ophthalmoplegia
internuclear, 209–210, 210f
neuromuscular causes of, 462e-2t
progressive external. *See* Progressive external ophthalmoplegia
video library, 41e-1v
- Ophthalmoscopy, 40e-1, 40e-2, 40e-2t
- Opioid(s)
action of, 92, 444e-4, 465e-3, 465e-3f, 465e-5t
adverse effects of, 61, 468e-1, 2232t
delirium, 168
hypogonadism, 2364
management of, 92
pruritus, 381
respiratory depression, 92
for back pain, 119
COX inhibitor combinations, 92–93
cross-tolerance among, 60
dosage of, 60
for dyspnea, 62
endogenous, 89
for frostbite, 478e-4
in mechanical ventilation, 1741
for migraine, 2592t, 2593
most commonly abused, 468e-1. *See also* Opioid use disorders
neurobiology of, 468e-1, 468e-1t, 468e-2f
for osteoarthritis, 2232, 2232t
overdosage/poisoning with, 468e-3, 468e-3t, 473e-3, 473e-13t, 1773
for pain, 60–61, 91t, 92, 94, 94t
pharmacology of, 468e-1–3
for restless legs syndrome, 192
routes of administration of, 92
systems effects of, 468e-2
withdrawal from, 468e-3
- Opioid receptors, 89, 468e-1, 468e-1t, 468e-2f
- Opioid use disorders, 468e-1
clinical features of, 468e-2
complications of, 468e-2
diagnosis of, 468e-1
epidemiology of, 468e-1
pathogenesis of, 465e-3, 465e-3f
prevention of, 468e-4
in terminally ill patient, 61
treatment of, 468e-3–4, 468e-3t
- Opisthorchis*/opisthorchiasis, 552, 765t, 1428, 1428t. *See also* Biliary (hepatic) flukes
- Oppenheim's dystonia, 2620
- Oprelvekin (IL-11), 103e-24
- Opsoclonus, 211
- Opsoclonus-mycoclonus syndrome, paraneoplastic, 372e-25t, 614t, 618
- Opsonization, 372e-3
- Optical coherence tomography (OCT), 1465, 1465f, 1668
- Optic ataxia, 181
- Optic atrophy, 203, 203f, 1136
- Optic chiasm, 197
- Optic cup, 205, 205f
- Optic disc
anatomy of, 40e-1
damage to, 197
drusen, 203–204, 203f
edema, 203
normal, 40e-1, 40e-1f
swelling of, 40e-5, 40e-5f
vascular sheathing over, 40e-4, 40e-4f
- Optic gliomas, 2264
- Optic nerve, 195
disorders of, 197–199. *See also* Optic neuritis;
Optic neuropathy
examination of, 2537
injuries of, 457e-2
tumors of, 206–207
- Optic neuritis, 196
color desaturation in, 197
epidemiology of, 202
ethambutol-related, 205e-4
in MS, 202, 2663
in neuromyelitis optica, 2656
paraneoplastic, 614t, 619
recurrent, 202
retrobulbar, 196, 202, 202f
in sarcoidosis, 2208
treatment of, 202
visual field defects in, 197
- Optic neuropathy
anterior ischemic, 202, 202f
drug-induced, 203
Leber's. *See* Leber's hereditary optic neuropathy (congenital amaurosis)
posterior ischemic, 202
toxic, 203, 203f
- Optokinetic reflex, 195
- Oral cancer
clinical features of, 46e-3f, 46e-4f, 239t, 503
diagnosis of, 237
incidence of and deaths from, 467t
prevention of, 476
risk factors for, 237, 2725
smoking and, 476
- Oral cavity. *See also* Oral mucosa
anaerobic bacterial infections of, 1095, 1097, 1102
anaerobic flora of, 1094–1095, 1095t
cancer treatment effects in, 623
F. tularensis infection of, 1068
infections of, 233–234, 487
in nutritional deficiencies, 463t
systemic disease signs in, 46e-1, 235
- Oral cholecystography (OCG), 2078
- Oral contraceptives, 2390
for acne vulgaris, 352
adverse effects of, 2391t
cholestasis, 284, 2025t, 2029
cutaneous, 132t, 352, 353, 366, 379
hair loss, 380
hepatotoxicity, 2025, 2027
hirsutism, 380

- hyperkalemia, 310
hypertension, 6e-4
lipoprotein disorders, 2443–2444
oral pigmentation, 240t
venous thrombosis, 744
for amenorrhea, 337
breast cancer risk and, 524
contraindications to, 334, 2391, 2391t
drug interactions of, 42t, 44, 2559
for dysmenorrhea, 339
formulations of, 2390–2391
for hirsutism, 334
HIV infection risk and, 1221
in perimenopause, 2382
for polycystic ovarian syndrome, 337
- Oral-facial-digital syndrome type I, 1850t
- Oral glucose tolerance test, 2401, 2401t
- Oral hairy leukoplakia, 1188, 1188f, 1190, 1255, 1256f
- Oral mucosa
age-related changes in, 242
diseases of
pigmented lesions, 240t
ulcers, 237
vesicular, bullous, or ulcerative lesions, 238–239t
white lesions, 240t
- Oral pain, nondental, 237, 239
- Oral rehydration solution (ORS)
for cholera, 1063–1064, 1064t
composition of, 856, 1058, 1063, 1064t
for infectious diarrhea, 267, 856, 857t
for *Shigella* infections, 1058
- Oral sex, HIV transmission and, 1220f, 1221, 1285
- Oral ulcers, 46e-4f, 237, 239t, 2129t
- Orán virus, 1308t
- Orbit, disorders of, 207
- Orbital pseudotumor, 207, 391e-1t
- Orbitofrontal cortex, 211
- Orbiviruses, 1305, 1308t, 1319
- Orchiectomy
for prostate cancer, 103e-3, 103e-15, 585, 586
for testicular cancer, 590
- Orchiopexy, 588
- Orchitis
in lepromatous leprosy, 1126
in mumps infection, 231e-2, 231e-3
viral, 2365
- Orexin (hypocretin), 185, 186f, 189, 215
- Orf, 220e-1t, 829
- Organic acid(s), renal transport of, 332e-8
- Organic anion transporters (OATs), 431e-1, 431e-2f
- Organic anion transport protein 1B1 (OATP1B1), 38, 2000
- Organic dusts, 1688t, 1691
- Organ of Corti, 217, 218f
- Organophosphates, 267, 473e-10t, 2688t
- Organ transplantation. *See* Solid organ transplantation
- Oriboca virus, 1309t
- Orientia tsutsugamushi*, 782, 1159. *See also* Scrub typhus
- Oritavancin
action of, 930, 931t
adverse effects of, 938t, 942
drug interactions of, 939t
indications for, 940t, 942, 976
in pregnancy and lactation, 937t
resistance to, 931t, 933–934
- Orlistat, 96e-8, 2397, 2453
- Ornithine decarboxylase, 246e-8
- Ornithine metabolism disorders, 434e-3t
- Ornithine transcarbamylase deficiency, 434e-3t
- Ornithodoros* spp. tick, 1146, 1147f
- Oromandibular dystonia, 2620
- Oropharyngeal disease
cancer, 477, 502, 503, 505, 1198. *See also* Head and neck cancer
dysphagia in, 256
in tularemia, 1068
- Oropouche virus, 1309t, 1318
- Orotic aciduria
hepatocellular carcinoma and, 545
megaloblastic anemia and, 649
in pyrimidine metabolism disorders, 431e-5t, 431e-6, 431e-6f
- Oroya fever, 1081t, 1083
- Orphenadrine, 473e-10t, 473e-13t
- ORS. *See* Oral rehydration solution (ORS)
- Orthobunyaviruses, 1305, 1308–1309t
- Orthobunyavirus infections, 1308–1309t, 1318
- Orthodoxia, 1992
- Orthodromic AV reentry, 1481, 1482f
- Orthomyxoviruses, 214e-2t, 214e-3f, 214e-4, 1305. *See also* Influenza virus
- Orthomyxovirus infections, 1311–1312t
- Orthopnea, 47e-3, 1503
- Orthopoxvirus, 220e-1t
- Orthostatic hypotension
approach to the patient, 2639–2640
clinical features of, 145–146, 2639, 2639t
definition of, 1445
diagnosis of, 2639–2640
dizziness in, 148
drug-related, 2640t, 2644
etiology of, 145t, 2639t
falls and, 165
in hypovolemia, 298
neurogenic, 146
in opioid use disorders, 468e-2
pathophysiology of, 142, 145, 146f, 1445
in PD, 2617
in pheochromocytoma, 1613
prevalence of, 2639t
treatment of, 146, 2644, 2644t
- Orthostatic proteinuria, 1834
- Orthostatic tachycardia, 298
- Orungo virus, 1308t
- OSAHS. *See* Obstructive sleep apnea/hypopnea syndrome (OSAHS)
- Osborn wave, 1457
- Oscillopsia, 148, 149
- Oseltamivir
action of, 214e-5, 214e-9, 215e-1
adverse effects of, 215e-1, 1213
for influenza prophylaxis, 215e-2t, 1213t, 1214, 1252t
for influenza treatment, 215e-2t, 215e-5, 233, 1213, 1213t
pharmacology of, 215e-1, 215e-2t
resistance to, 215e-1, 215e-2t
- Osler maneuver, 1626
- Osler's nodes, 1443
- Osler-Weber-Rendu syndrome (hereditary hemorrhagic telangiectasia)
bleeding in, 403
clinical features of, 357
gastrointestinal bleeding in, 277
skin manifestations of, 356t, 357, 1442
small intestine angioma in, 536
vascular malformations in, 732, 2585
- Oslo Study, 1134
- OSM (ocostatin M), 372e-11t
- Osmolality, 295
- Osmolar gap, 319
- Osmolytes, 301
- Osmoreceptors, 295, 332e-9–10, 332e-10f, 2274–2275
- Osmoregulation, 332e-9–10, 332e-10f, 2279f
- Osmotic demyelination syndrome, 64e-5, 301, 302
- Osmotic diarrhea, 269, 303, 1933. *See also* Diarrhea
- Osmotic diuresis, 297
- Osmotic equilibrium, 296
- Osmotic fragility, 651
- Osmotic laxatives. *See* Laxatives
- OspA, 1150
- OspC, 1150
- Ospemifene, 2383
- Ossa virus, 1309t
- Ossification, extraskelatal, 426e-7–8, 426e-7t
- Osteitis fibrosa cystica, 1815, 2472, 2487
- Osteoarthritis (OA), 2226
in acromegaly, 2240–2241
back pain in, 117
clinical features of, 2230
definition of, 2226
diagnosis of, 2226, 2230, 2230f
global considerations in, 2228
joints affected by, 2226, 2226f
hand, 2226, 2226f
hip, 2228, 2230
knee, 2226, 2228–2229, 2228f, 2230, 2230f
toe, 2229f
neck pain in, 122
oral pain in, 237
pain sources in, 2230
pathogenesis of, 2226–2227, 2227f
pathology of, 2229–2230, 2229f
prevalence of, 2226
risk factors for, 2227–2229, 2228f
treatment of, 2230, 2232t
acetaminophen, 2231
cartilage regeneration, 2233
COX-2 inhibitors, 2232
exercise, 2231
intraarticular injections, 2232
malalignment correction, 2231
nonpharmacotherapy, 2230–2231
NSAIDs, 2231–2232
opioid analgesics, 2232
surgical, 2233
- Osteoarthropathy, hypertrophic. *See* Hypertrophic osteoarthropathy
- Osteoblast(s), 426e-1f, 2455f, 2456f, 2467
- Osteoblastic lesion, 119e-4
- Osteoblastoma, 609t
- Osteocarcinoma, 101e-2t
- Osteochondrodysplasias, 426e-7
- Osteochondroma, 119e-3
- Osteochondromatosis (multiple exostoses), 426e-7
- Osteoclast(s), 426e-1f, 2455–2456, 2455f, 2456f, 2467
- Osteoclast activating factor, 713
- Osteocyte(s), 2455
- Osteodystrophy, renal, 291, 2478
- Osteogenesis, 119e-4
- Osteogenesis imperfecta, 2507
classification of, 2507–2508, 2508t
clinical features of, 219, 1443, 2508–2509
diagnosis of, 2509
genetic considerations in, 435, 2509
incidence of, 2508
treatment of, 96e-5, 2509–2510
- Osteolytic lesion, 119e-4
- Osteoma cutis, 363–364
- Osteomalacia
axial, 426e-6
diagnosis of, 2465–2466
in hypophosphatemia, 426e-5–6
hypophosphatemia and, 2465
oncogenic (tumor-induced), 609t, 612
pathogenesis of, 462e-19, 2465
postgastrectomy, 1927
secondary hyperparathyroidism and, 2478
tumor-induced, 2459, 2460
- Osteomyelitis, 838
actinomycotic, 1088, 1089f
anaerobic bacterial, 1099–1100, 1102
B. dermatitidis, 1338
from bite-wound infections, 167e-1
classification of, 838–839
clinical features of, 152e-6t
E. coli, 1028
enterococcal, 974
foot, 845, 845f, 1046
in long bones
clinical features of, 842, 842f
complications of, 843
diagnosis of, 842
epidemiology of, 842
pathogenesis of, 841–842
treatment of, 840t, 841t, 842–843
Nocardia, 1087t
nontyphoidal *Salmonella*, 1054

- Osteomyelitis (*Cont.*):
P. aeruginosa, 1045t, 1046
 pathogenesis of, 194e-2, 838-839, 844
 in prosthetic joints. *See* Prosthetic joint infections
 psoas abscess in, 852
 in returning war veterans, 152e-6t, 152e-9
S. aureus, 957, 957f
 skull-base, 229
 sternal, 844-845, 844f
 treatment of, 840t, 841t, 963, 1102, 2429
 tuberculous, 836, 839, 844
 vertebral, 957, 957f, 1046
 clinical features of, 117, 839
 complications of, 841
 diagnosis of, 117, 839-840, 842f
 epidemiology of, 839
 microbiology of, 839
 pathogenesis of, 839
 treatment of, 840-841, 840t, 841t
- Osteonecrosis
 drug-induced, 2218t
 in HIV infection, 1261
 in IBD, 1958
 of jaw, 242, 623, 2481, 2501
 oral, 46e-4f
- Osteopathic medicine, 14e-1t, 14e-2
- Osteopenia, 2218t
- Osteopetrosis, 426e-4
- Osteopoikilosis, 426e-5
- Osteopontin, 89e-2
- Osteoporosis, **2488**
 approach to the patient, 2494-2495, 2495t
 back pain in, 117
 biochemical markers in, 2494-2495, 2494t
 bone biopsy in, 2494
 bone mass measurement in, 2493, 2493f, 2493t
 definition of, 2488
 diet and, 95e-5
 differential diagnosis of, 2494
 epidemiology of, 2488
 fractures in
 epidemiology of, 2488-2489, 2488f
 management of, 2495
 prevention of, 2499-2502, 2500f, 2502f
 risk factors for, 2489, 2490t
 glucocorticoid-induced, 2503-2504
 heparin-induced, 752
 in HIV infection, 1261
 in liver disease, 2067
 pathophysiology of, 2489-2490, 2491f
 prednisone and, 2145
 prevalence of, 2254t
 prevention of, 84, 2383, 2384t
 in rheumatoid arthritis, 2138
 risk factors for, 2489, 2490t
 assessment of, 2495-2496, 2496f
 chronic disease, 2492
 cigarette smoking, 2493
 estrogen status, 2491-2492, 2492f
 low calcium intake, 2491
 medications, 2492-2493
 physical inactivity, 2492
 vitamin D deficiency in, 2491
 screening for, 28, 29t, 84, 2254t
 treatment of, 117, **2495**
 anabolic agents, 2503
 bisphosphonates, 2499-2501, 2500f
 calcitonin, 2501
 calcium supplementation, 2496-2497, 2496t, 2497t
 denosumab, 2501, 2502f
 estrogens, 2497-2498, 2498f
 exercise, 2497
 fluoride, 2503
 GH, 2503
 monitoring of, 2503
 nonpharmacologic, 2503
 nutritional recommendations, 2496
 progestins, 2498
 PTH, 2502-2503, 2503f
 risk factor reduction, 2495-2496, 2496f
 SERMs, 2498-2499
 strontium ranelate, 2503
 vitamin D, 2497
 vertebral fracture in, 117
 in women, 6e-4
- Osteoprotegerin (OPG)
 in bone metastasis, 102e-10-11
 in bone remodeling, 2490, 2492f
 in osteopetrosis, 426e-4
 in primary myelofibrosis, 675
- Osteosarcoma, 101e-2t, **119e-3**, 236
- Osteosclerosis, 118, 426e-5, 675. *See also* Paget's disease of bone
- Ostium primum, 51e-4, 1520
- Ostium secundum, 51e-4, 1520
- Otitis externa, 228-229, 1045t, 1046-1047, 1348
- Otitis media, **229**
 acute, 229-230, 229t, 952, 953
 anaerobic bacterial, 1097
 brain abscess associated with, 901
C. auris, 981
 chronic, 230, 1097
 clinical features of, 230, 952
H. influenzae, 229-230, 1011
 hearing loss in, 217
 in influenza, 1212
M. catarrhalis, 229-230, 1013
 in nasopharyngeal cancer, 503
 recurrent, 230
 serous, 230
 treatment of, 229t, 230, 953
- Otoacoustic emissions, 219
- Otosclerosis, 219
- Otology, 217
- OTUs (operational taxonomic units), 86e-2
- Outcome (quality theory), 12e-2
- Outcome evaluation, 3
- Outer-membrane vesicles (OMVs), 1002, 1002f
- Outpatient care, 5
- Ova and parasite examination, 267
- Ovalocytosis, malaria resistance and, 1371
- Ovarian cancer, **592**
 breast cancer and, 592
 carcinoma of unknown primary and, 120e-3
 clinical features of, 592-593
 deaths from, 467t, 469f, 470t
 epidemiology of, 592
 epithelial, 592-593
 genetic considerations in, 101e-2t, 101e-6t, 523, 592
 germ cell tumors, 594-595
 incidence of, 467t, 592, 593
 infections in, 484t
 metastatic, 408
 oral contraceptives and, 2391t
 paraneoplastic syndromes in
 endocrinologic, 609t
 hematologic, 612, 612t
 neurologic, 614t, 615f, 617
 pathogenesis of, 592
 pathology of, 592
 postmenopausal hormone therapy and, 2384t, 2386
 during pregnancy, 124e-1t
 prognosis of, 593t, 594
 protective factors for, 592
 recurrent, 593-594
 risk factors for, 592
 screening for, 482, 593
 sex cord tumors, 594
 staging of, 593, 593t
 surgical prevention of, 479
 treatment of, 593-594
 tumor markers in, 473t, 593
- Ovarian follicles, 2376, 2376f
- Ovarian neuroendocrine tumors (carcinoids), 564
- Ovarian reserve, 2388
- Ovary
 development of, 2375-2376, 2375f, 2376f
 disorders of
 cancer. *See* Ovarian cancer
 cyst, 106, 404
 drug-related, 2132
 dysfunction/failure, 336-337, 437, 2388, 2389
 sex cord tumors, 594
 teratoma. *See* Teratoma, ovarian
 function of, **2376**
 assessment of, 2379
 hypothalamic secretion in, 2376-2377, 2377f
 ovarian peptides in, 2378
 ovarian steroids in, 2377-2378, 2377f. *See also* Estrogen(s); Progesterone
 pituitary secretion in, 2376-2377, 2377f
 metastatic disease to, 593
- Overactive bladder, 7e-4, 60e-2
- Overdiagnosis, 479, 509
- Overdosage. *See* Poisoning/drug overdose
- Overflow diarrhea, 265
- Overflow incontinence, 80-81
- Overlapping sphincteroplasty, 1976
- Overweight, 98e-1t, 415e-1, 2393t. *See also* Obesity
- Ovotesticular disorders of sexual development, 2351t, 2353
- Oxacillin
 adverse effects of, 306, 2023
 for burn-wound infections, 166e-3
 for cellulitis, 832t
 indications for, 937, 940t
 for infective endocarditis, 823
 for nongonococcal bacterial arthritis, 835
 for osteomyelitis, 840t, 841t
 resistance to, 933, 940t
 for staphylococcal infections, 961, 962t
- Oxalate, 318, 319, 1867
- Oxaliplatin
 action of, 103e-12
 adverse effects of, 103e-9t, 103e-12, 543, 1793
 for colorectal cancer, 543
 for pancreatic cancer, 557
- Oxalosis, 62e-10f, 2237, 2237f
- Oxamniquine, 246e-5t, 246e-11
- Oxandrolone, 2371
- Oxazepam, 473e-10t, 1606, 2711t
- Oxazolidinones, 931t, 932, 932f. *See also* Linezolid; Tedizolid
- Oxcarbazepine
 adverse effects of, 2554t, 2718t
 for anxiety disorders, 2711
 for bipolar disorder, 2718, 2718t
 drug interactions of, 2554t
 for epilepsy, 2552, 2554t
 overdose/poisoning with, 473e-10t
 for pain, 61, 91t
 pharmacology of, 2554t, 2718
 for trigeminal neuralgia, 2646
- Oxicam, 383
- Oxidative phosphorylation, 462e-15
- Oxidative stress
 aging and, 75, 94e-3, 94e-7, 2450
 in asthma, 1674
 metabolic syndrome and, 2450
- P450 Oxidoreductase deficiency, 2328t
- Oxime therapy, 262e-5t, 262e-6, 262e-6f
- 5-Oxoproline, 320
- 3-Oxothiolase deficiency, 434e-3t
- Oxybutynin, 192, 2660, 2672
- Oxycodone
 abuse of. *See* Opioid use disorders
 for dyspnea, 62t
 for pain, 91t
- Oxygen
 arterial concentration of, 1748t, 2762t
 plasma concentration of, 1748t
 toxicity of, 477e-2
 transport in blood, 631
 venous concentration of, 1748t, 2762t
- Oxygen-carrying capacity of hemoglobin, 632, 1748t
- Oxygen delivery, 1733, 1748t
- Oxygen delivery index, 1748t
- Oxygen extraction ratio, 1748t
- Oxygen-hemoglobin dissociation curve, 247, 393, 632, 632f

- Oxygen therapy
 for asthma exacerbation, 1679
 for cluster headache, 2595
 for COPD, 1706, 1707
 for dyspnea, 47e-4, 62
 for high-altitude pulmonary edema, 476e-4
 hyperbaric. *See* Hyperbaric oxygen therapy
 for hyponatremia, 302
 for hypothermia, 478e-2
 for hypovolemic shock, 1748
 for myocardial infarction, 1603
 for pulmonary edema, 1763
 for respiratory acidosis, 324
 Oxygen uptake, 306e-3, 1748t
 Oxygen uptake index, 1748t
 Oxyhemoglobin, 632, 632f
 Oxytazoline, 2122
 Oxyntic cell, 1911, 1911f
 Oxyntic gland, 1911, 1911f
 Oxyphenbutazone, 727t
 Oxytocin, 2274, 2275, 2275f
 Ozone exposure, 1692t
- P**
 P1 syndromes, 2575
 P2 syndromes, 2575
 P2Y₁₂ antagonists, reversible, 749
 P5N (pyrimidine-5'-nucleotidase) deficiency, 431e-5t, 431e-6f
 P5N (pyrimidine-5'-nucleotidase) superactivity, 431e-5t
 p14^{ARF}, 102e-2, 494
 p16/*CDKN2A* gene mutations, 554
 p16 gene mutations, 554, 2305t
 p16^{INK4a}, 89e-3, 101e-6t, 102e-1, 102e-7
 p16 protein, 494
 p21 gene mutations, 576
 p21 protein, 102e-1
 p21/*WAF* gene mutations, 2305t
 p24 antigen capture assay, 770, 771f, 1243f, 1246-1247, 1247t
 p27 protein, 102e-1
 p53 gene, adenoviral-mediated expression of, 91e-4
 p53 gene mutations
 in bladder cancer, 576
 in breast cancer, 523, 528
 in cancer, 101e-8, 102e-2, 102e-2f, 431t
 in colorectal cancer, 101e-9f
 in esophageal cancer, 533
 in head and neck cancer, 503
 in Li-Fraumeni syndrome, 119e-1, 599t
 in lung cancer, 507
 in pancreatic cancer, 554
 in skin cancer, 387
 in skin response to sun, 386
 in thyroid cancer, 2304, 2305t
 p53 protein
 activation of, 102e-2, 102e-2f
 in cancer cells, 102e-2, 102e-15, 372e-23, 372e-25t
 in colon cancer, 101e-1f
 in melanoma, 494
 in SLE, 372e-25t
 VEGF inhibition and, 102e-16f
 p62, 372e-25t
 p63, 508
 p70S6K, 102e-4f
 p-80-Collin, 372e-24t
 P450 oxidoreductase deficiency, 2328t
 PA 824, 205e-7
 PACE (Program for All Inclusive Care of the Elderly), 82-83
 Pacemaker, **1469**
 for AV conduction block, 1474-1476, 1475t, 1476t
 for bradyarrhythmia, 273e-7, 1466
 for carotid sinus hypersensitivity, 1470
 CMR in patient with, 270e-9
 complications of, 816, 1469
 five-letter codes for, 1469
 indications for, 273e-7
 for myocardial infarction, 1475-1476, 1475t, 1609
 nomenclature of, 1469
 pacing mode, 1469, 1476
 for pulmonary edema, 1763
 for SA node dysfunction, 1470, 1470t
 travelers with, 799
 for vasovagal syncope, 1470
 Pacemaker cells, 273e-1, 273e-3, 1466
 Pacemaker syndrome, 1469
 Pachydermoperiostosis, 2244. *See also* Hypertrophic osteoarthropathy
 Packed red blood cells, 138e-2, 138e-2t
 Paclitaxel
 action of, 103e-14-15
 adverse effects of, 103e-12t, 103e-15
 hypersensitivity reaction, 1798
 lupus syndrome, 2218t
 myopathy, 2218t
 nail disorders, 380
 neurologic, 1793, 2686t
 for bladder cancer, 577
 for breast cancer, 529, 530
 for carcinoma of unknown primary, 120e-4
 in drug-eluting stents, 296e-2
 for head and neck cancer, 505
 for lung cancer, 518t, 520t, 521t
 for melanoma, 499
 for ovarian cancer, 593
 premedication for, 103e-12t, 103e-15
 for testicular cancer, 591
PADI4 gene mutations, 2140
 PAF (pure autonomic failure), 2642
 Paget disease, extramammary, 502
 Paget-Schroetter syndrome, 1647
 Paget's disease of bone, **426e-1**
 back pain in, 118
 clinical features of, 426e-2
 complications of, 426e-2
 diagnosis of, 426e-2-3, 426e-2f, 426e-3f
 epidemiology of, 426e-1
 etiology of, 426e-1, 426e-1f
 genetic considerations in, 426e-1
 global considerations in, 426e-1
 juvenile, 426e-1
 multiple-site, 436e-3f
 oral manifestations of, 236
 pathophysiology of, 426e-1-2
 of tibia, 436e-3f
 treatment of, 426e-3-4, 426e-3t
 PAH. *See* Pulmonary arterial hypertension (PAH)
 PAI-1. *See* Plasminogen activator inhibitor I (PAI-1)
 Pain
 acute, treatment of, **90**
 acetaminophen for. *See* Acetaminophen
 aspirin for. *See* Aspirin
 nonsteroidal anti-inflammatory drugs for. *See* Nonsteroidal anti-inflammatory drugs
 opioids for. *See* Opioid(s)
 approach to the patient, 111-114
 assessment of, 60, 2539
 cancer, 473-474
 chronic, 80, **93**, 93t. *See also* Opioid(s)
 depression and, 93
 etiology of, 60
 frequency of, 60
 in MS, 2672
 neuropathic, 60, 89-90, 94-95. *See also* Neuropathy(ies)
 nociceptive, 60
 pathophysiology of, **87**
 central mechanisms, 88-89, 89f
 neuropathic pain, 89-90
 pain modulation, 89, 90f, 91f
 peripheral mechanisms, 87-88, 87f, 88f
 perception of, 107
 psychological variables, 93
 racial/ethnic disparities in treatment of, 16e-1
 referred. *See* Referred pain
 sympathetically maintained, 90
 in terminally ill patient, 60-61
 testing sense of, 159, 160t
 visceral, 60
 Painful articular syndrome, 1261
 Painless (silent) thyroiditis, 2299
 Palatal tremor, 41e-1v
PALB2 gene mutations, 554
 Palifermin, for mucositis, 103e-25, 139e-3
 Palilalia, 177, 2623
 Paliperidone, 2721, 2722t
 Palivizumab, 1206
 Palla's sign, 1633
 Palliative care, **56**. *See also* End-of-life care
 for cancer, 103e-3, 103e-7, 472, **473**
 insurance coverage of, 70
 for non-hospice patient, 70
 Pallister-Hall syndrome, 2264
 Pallister-Killian syndrome, 83e-6
 Palmar crease, 393, 1442
 Palmar erythema, 1992, 2058, 2059f
 Palmar fasciitis, 2247
 Palmonal response, 2539
 Palonosetron, 62, 103e-25, 261
 Palpation
 of abdomen, 106
 of chest, 1662
 of heart, 1447
 of spleen, 411
 Palpitations, **254**
 approach to the patient, 254
 in arrhythmia, 254, 273e-5
 in atrial fibrillation, 254
 etiology of, 254
 treatment of, 254
 Palsy(ies)
 Bell's. *See* Bell's palsy
 cranial nerve. *See* Cranial nerve disorders
 2-PAM Cl (2-pralidoxime chloride), for nerve agent exposure, 262e-5t, 262e-6, 262e-6f
 Pamidronate
 for bone pain, 61
 for hypercalcemia, 314, 610, 2480t, 2481
 for metastatic bone disease, 119e-4
 for multiple myeloma, 717
 for Paget's disease of bone, 426e-3-4, 426e-3t
 PAMPs (pathogen-associated molecular patterns), 145e-5, 145e-8f, 372e-2, 1747
 Pan American Sanitary Bureau, 8
 pANCA (perinuclear ANCA), 2180
 Pancoast tumor
 in lung cancer, 510
 neck pain in, 123
 shoulder pain in, 2221
 treatment of, 519
 Pancreas
 annular, 2102
 beta cells of, 2403-2404
 biopsy of, 2087t, 2089
 disorders of. *See* Pancreatic disease
 exocrine secretion in, **2090**
 in autoprotection of pancreas, 2091
 enteropancreatic axis and feedback inhibition in, 2091
 enzymes in, 2090-2091
 insufficient, 2098, 2098t. *See also* Chronic pancreatitis
 laboratory evaluation of, 2087t, 2090
 regulation of, 2090
 in steatorrhea, 1940
 water and electrolyte content of, 2090
 fluid volumes and electrolyte content of, 98e-4t
 transplantation, 928, 2417, 2433
 Pancreas divisum, 2102
 Pancreatotomy, 2101
 Pancreatic agenesis, 431t
 Pancreatic cancer, **554**
 clinical features of, 366, 554-555
 deaths from, 467t, 468f, 470t
 diagnosis of, 555, 555f
 epidemiology of, 554
 extrahepatic cholestasis in, 285
 gastroparesis in, 259
 genetic considerations in, 101e-2t, 102e-7, 554
 global considerations in, 554
 incidence of, 467t

- Pancreatic cancer (*Cont.*):
 obstruction due to, 1892, 1894f
 paraneoplastic syndromes in, 609t, 612t, 619
 precursor lesions in, 554
 prognosis of, 556f
 risk factors for, 554
 screening for, 554
 staging of, 555, 556f
 treatment of, 102e-5t, 555, 556t, 557, 557t
 tumor markers in, 473t, 555
- Pancreatic cholera. *See* VIPoma (vasoactive intestinal peptidoma)
- Pancreatic disease. *See also* Pancreatitis
 abdominal pain in, 103-104
 annular pancreas, 2102
 approach to the patient, **2086**
 ascites in, 288
 back pain in, 118
 cancer. *See* Pancreatic cancer
 in cystic fibrosis, 1697. *See also* Cystic fibrosis (CF)
 diagnosis of
 biopsy, 2089
 imaging, 2087t, 2088-2089, 2089t
 pancreatic enzymes, 2087-2088, 2087t
 pancreatic function tests, 2087t, 2090
 endocrine tumors. *See* Pancreatic neuroendocrine tumors (pNETs)
 etiology of, 2086
 macroamylasemia, 2102
 pancreas divisum, 2102
 skin manifestations of, 366
- Pancreatic enzyme(s)
 activation of, in pancreatitis, 2090
 for irritable bowel syndrome, 1969
 in pancreatic disease diagnosis, 2087-2088, 2087t, 2089t
- Pancreatic enzyme preparations, 272, 2100-2101, 2101t
- Pancreatic intraepithelial neoplasia (PanIN), 554
- Pancreatic necrosis, 346e-1v, 1881, 2095-2096
- Pancreatic neuroendocrine tumors (pNETs), **567**
 classification of, 558, 559t, 560
 clinical features of, 559t, 567
 Cushing's syndrome and, 559t, 571
 general characteristics of, 558t
 genetic syndromes associated with, 562, 562t
 incidence of, 559t, 560-561, 561t
 malignant behavior of, 559t, 561-562
 nonfunctional, **571**, 2338, 2338f
 paraneoplastic syndromes in, 609t
 prognosis of, 561t, 573f
 rare syndromes, 559t, 571
 screening tests for, 2337t
 syndromes of, 559t
 glucagonoma. *See* Glucagonoma
 GRFoma, 559t, **571**
 insulinoma. *See* Insulinoma
 in MEN 1, 2336, 2336t
 somatostatinoma, 269, 559t, 561, **570**
 VIPoma. *See* VIPoma (vasoactive intestinal peptidoma)
 Zollinger-Ellison syndrome. *See* Zollinger-Ellison syndrome (ZES)
 treatment of, 102e-3, 103e-16, 568, 572-573
 tumor localization in, 571-572, 572f
- Pancreaticoduodenectomy, 553, 555
- Pancreaticopleural fistula, 2095f
- Pancreatic polypeptide (PP), 571
- Pancreatic polypeptide-secreting tumors (PPomas), 559t, 2338
- Pancreatitis. *See also* Acute pancreatitis; Chronic pancreatitis
 ascites in, 287
 autoimmune, 391e-1, 391e-1t, 2099, 2099t
 choledocholithiasis and, 2084
 cobalamin malabsorption in, 645
 diagnosis of, 107
 differential diagnosis of, 2093
 endoscopic findings in, 1899-1900
 gallstone, 1893-1894, 2096
 hereditary, 554, 2101
 hypocalcemia in, 315
 hypovolemia in, 298
 in mumps, 231e-3
 nausea and vomiting in, 259
 pleural effusions in, 2092, 2095f
 post-ERCP, 1891
 tumor markers in, 473t
- Pancuronium, 1784
- Pancytopenia, 662, 663t
- Panencephalitis, 900
- Paneth cells, 86e-8
- Pan-genome, 86e-1t, 86e-2
- Panic attack, 254
- Panic disorder, 96t, 101, **2708**
- PanIN (pancreatic intraepithelial neoplasia), 554
- Panitumumab
 action and targets of, 102e-5t
 adverse effects of, 103e-26, 543
 for colorectal cancer, 102e-5t, 543
 genetic variations in response to, 39t, 41
- Panniculitis, 364t, 366
- PanOptic ophthalmoscopy, 40e-2, 40e-2t
- Panton-Valentine leukocidin (PVL) toxin, 807, 830, 956
- Pantoprazole, 263, 1920, 1920t, 1930
- Pantothenate kinase-associated neurodegeneration (PKAN), 2611, 2626
- Pantothenic acid (vitamin B₅), 95e-2t, **96e-5**, 98e-5t
- Panuveitis, 200
- PAP (pulmonary alveolar proteinosis), 1714
- Papain, 2739, 2740
- PAPA (pyogenic arthritis, pyoderma gangrenosum, and acne) syndrome, 372e-8t, 422
- Papillary fibroelastoma, 289e-2
- Papillary necrosis, 1860, 1860t
- Papillary renal carcinoma, 101e-5, 101e-6t, 578, 578t. *See also* Renal cell carcinoma (RCC)
- Papillary stenosis, 2082
- Papilledema, 40e-5, 40e-5f, 203, 203f
- Papillomaviruses. *See* Human papillomavirus (HPV)
- Papillon-Lefevre syndrome, 236
- PA/PRA. *See* Aldosterone-renin-ratio (ARR)
- Pap smear, 28t, 29t, 480, 481t, 595
- Papular eruptive xanthomas, 436e-4f
- Papular mucinosis, 364t, 365
- Papule, 127, 339t, 340f. *See also* Papulonodular skin lesions
- Papulonodular skin lesions, **363**
 blue, 364t, 367
 etiology of, 828t, 829
 pink, 364t, 365
 purple, 364t, 367
 red, 364t, 365-367
 red-brown, 364t, 367
 skin-colored, 364-365, 364t
 violaceous, 364t, 367
 white, 363-364, 364t
 yellow, 364t, 365
- Papulosis, lymphomatoid, 710
- Papulosquamous lesions, **353**, 353t
- PAR1 (protease activation receptor 1), 276, 742
- Para-aminosalicylic acid, 205e-6-7, 644t
- Parabacteroides distasonis*, 1094
- Paracellular transport, 332e-3
- Paracentesis, abdominal, 287, 288, 474, 483e-1v
- Parachlorophenylamine, 567
- Paracoccidiodomycosis, 1354, 1354t
- Paracrine regulation, 400e-6
- Paradoxical pulse, 1574
- Paradoxical undressing, 478e-1
- Paraesophageal hernia, 1902
- Parafacial respiratory group, 1720
- Parafibromin, 2337t, 2342
- Paraganglioma
 definition of, 2329
 diagnosis of, 2329-2330, 2330t
 genetic considerations in, 1619, 2329, 2334-2335, 2334f
 hypertension in, 1619
 pathogenesis of, 2329, 2330f
 treatment of, 2330-2331
- Paraganglioma syndromes, 2332, 2333f
- Paragonimus* spp. *See* Lung flukes (*Paragonimus* spp.)
- Parainfluenza virus, 1207
- Parainfluenza virus infections, **1207**
 in cancer patient, 489, 1207
 clinical features of, 225, 226, 231, 234, 1202t, 1207
 diagnosis of, 1207
 epidemiology of, 1207
 pathogenesis of, 1207
 prevention of, 1208
 in transplant recipient, 923, 1207
 treatment of, 214e-5, 1207-1208
- Paralysis
 definition of, 154
 periodic. *See* Periodic paralysis
 in rabies, 1301-1302
 sleep, 188t, 189
 tick-bite, 989, 2745
 Todd's, 2542
 tongue, 2649
- Paralytic poliomyelitis, 1290. *See also* Poliovirus infection
- Paralytic shellfish poisoning, 2742
- Paramyotonia congenita, 444e-2t, 462e-3, 462e-17t, 462e-18
- Paramyxoviruses
 gene expression and replication in, 214e-4
 human metapneumovirus, 1206
 measles. *See* Measles (rubeola)
 mumps. *See* Mumps
 structure of, 214e-2t
- Paranasal sinus cancer. *See* Head and neck cancer
- Paraneoplastic syndromes
 ataxia in, 2626
 definition of, 608
 endocrinologic, **608**
 ectopic ATCH production. *See* Adrenocorticotropic hormone (ACTH), ectopic production of
 ectopic hCG production, 608, 612
 etiology of, 608-609, 609t
 hypercalcemia. *See* Hypercalcemia, malignancy-associated
 oncogenic osteomalacia, 612
 tumor-associated SIADH, 610. *See also* Syndrome of inappropriate antidiuretic hormone (SIADH)
 tumor-induced hypoglycemia, 611-612
- eye movements in, 41e-1v
- hematologic, **612**, 612t
- in hepatocellular carcinoma, 545-546
- immunopathology of, 372e-25t
- joint involvement in, 2247
- in lung cancer, 511
- neurologic, **614**, 614t
 acute necrotizing myopathy, 619
 anti-Hu neuropathy, 2700
 approach to the patient, 615-616
 autoimmune autonomic ganglionopathy, 2642
 cerebellar degeneration/atrophy, 372e-25t, 614t, 617-618
 channelopathies in, 444e-2t
 vs. Creutzfeldt-Jakob disease, 453e-5
 diagnosis of, 615-616, 615f
 dorsal root ganglionopathy, 618
 encephalitis with antibodies to cell-surface or synaptic proteins, 614t, 617, 617f, 2607
 encephalomyelitis and focal encephalitis, 372e-25t, 616-617, 616f
- Lambert-Eaton myasthenic syndrome. *See* Lambert-Eaton myasthenic syndrome (LEMS)
- opsoclonus-myoclonus syndrome, 372e-25t, 614t, 618
- pathogenesis of, 614, 614t
- peripheral neuropathies, 618-619
- sensory neuroneuropathy/ganglionopathy, 618, 2685
- spinal cord, 618

- stiff-person syndrome, 618
visual syndromes, 619
pemphigus, 361, 361t, 370t, 372
in renal cell carcinoma, 578
- Paranoá virus, 1308t
- Paraparesis, 156–157, 156f
- Parapharyngeal space infections, 235
- Paraphasia, 177
- Paraplegin* gene, 2634t, 2636
- Parapneumonic effusion, 951, 1717
- Parapoxviruses, 220e-1t, 220e-2
- Paraproteinemia, 404. *See also* Plasma cell disorders
- Parapsoriasis, 353, 353t
- Parasite(s)
adherence of, 145e-2t, 145e-3
entry into host, 145e-1
laboratory identification of, 245e-1
 alternative procedures, 245e-5t
 blood and tissue parasites, 245e-1–4
 flatworm infections, 245e-2t. *See also*
 Cestodes/cestode infections;
 Trematodes/trematode infections
intestinal parasites, 245e-1
minor procedures, 245e-5t
molecular methods, 245e-6, 245e-6t
nonspecific tests, 245e-4, 245e-6
protozoal infections, 245e-4t. *See also*
 Protozoa
 roundworm infections, 245e-3t. *See also*
 Nematodes/nematode infections
tissue invasion by, 145e-8
- Parasitic infections. *See also specific infections*
arthritis in, 837–838
as cause of fever of unknown origin, 137t
hepatobiliary, 2084
myocarditis in, 1560–1561
polymyositis, 2198
renal involvement in, 1849–1850
transfusion-transmitted, 138e-6
in transplant recipient, 921
treatment of, 246e-1
- Parasomnias, 192–193
- Parathyroid disorders
adenoma, 2470
carcinoma, 2471, 2473
genetic considerations in, 2471–2472, 2471f
hyperparathyroidism. *See*
 Hyperparathyroidism
hyperparathyroidism-jaw tumor syndrome.
 See Hyperparathyroidism-jaw tumor
 (HPT-JT) syndrome
hyperplasia, 2470
hypoparathyroidism. *See* Hypoparathyroidism
in MEN syndromes. *See entries at* Multiple
 endocrine neoplasia
- Parathyroidectomy, 2473–2474, 2487
- Parathyroid hormone (PTH), 2466
actions of, 313, 313f, 2466–2467
assessment of, 400e-6
in calcium homeostasis, 2457, 2476F
in chronic kidney disease, 333e-3, 1815, 1816
deficiency of, 314–315
ectopic production of, 314, 511, 608, 609t
excess of, 313
exogenous injection of. *See* Teriparatide
half-life of, 400e-2
in hypercalcemia, 2480
for hypoparathyroidism, 2484
metabolism of, 2467–2468
in primary hyperparathyroidism, 2473, 2473f
vs. PTHrP, 2468, 2468f
receptor cross-talk of, 400e-1
reference values for, 2762t
regulation of, 2467
resistance to, 2486F, 2486f
structure of, 2467, 2468f
synthesis of, 2467
- Parathyroid hormone (PTH)/PTHrP receptor,
400e-1, 2469, 2469f, 2476f
- Parathyroid hormone-related protein (PTHrP),
2468
actions of, 2468–2469, 2469f
ectopic production of, 609–610, 609t
- vs. PTH, 400e-1, 2468, 2468f
structure of, 2468, 2468f
- Paratonia (gegenhalten), 154
- Parcopa, for PD, 2616t
- PARD3B* gene, 1241t
- Parachoviruses, 894, 1294–1295
- Parental disomy, 597
- Parental imprinting, 101e-4
- Parenteral nutrition, 98e-1. *See also* Specialized
 nutritional support
 catheter placement for, 98e-3–4, 98e-6
 cholestasis in, 284, 2030
 combined with enteral feeding, 98e-3
 complications of, 98e-6–8, 98e-7t, 295
 definition of, 98e-1
 diabetes mellitus and, 2421
 electrolyte additions to, 98e-5, 98e-5t
 global considerations in, 98e-9
 home, 98e-4
 hypercalcemia in, 313
 indications for, 98e-3
 metabolic alkalosis in, 322
 mineral supplementation in, 98e-6t
 monitoring of, 98e-6, 98e-6t
 nutrient composition of, 95e-4
 risks of, 98e-3, 431e-5
 vitamin requirements in, 98e-5, 98e-5t
- Paresis, 1136. *See also* Periodic paralysis
- Paresthesia, 158, 1946t
- Parietal cell, 1911, 1912f, 1913
- Parietal cell antibodies, 645
- Parietal cortex, posterior, 180, 180f
- Parietal lesions, 209
- Parietofrontal network, 180, 180f
- Parinaud's syndrome/complex
in *Bartonella* infections, 1079
clinical features of, 195, 210
etiology of, 210
in pineal region tumors, 602
in tularemia, 1067
- Parkes-Weber syndrome, 1651, 1654
- parkin* gene mutations, 2609, 2613
- Parkinsonism
ALS and, 2633, 2635t
atypical, 2609–2611, 2612t
differential diagnosis of, 2610t
drug-induced, 2611, 2624
gait disorders in, 162t, 163
lower body, 163
in multiple system atrophy, 2641
secondary, 2611, 2612t
in Wilson's disease. *See* Wilson's disease
- Parkinson's disease (PD), 2609
AD and, 2599
clinical features of, 2609, 2609t
autonomic disturbances, 2617–2618
cardinal, 2609, 2609t
constipation, 274, 2618
delirium, 166
dementia, 171, 2599, 2605, 2617
dysphagia, 256, 257
frontal lobe syndrome, 184
motor, 33e-1v, 163, 2609, 2609t, 2618
nonmotor, 2609t, 2617–2618
olfactory dysfunction, 211, 214–215
psychoses, 2617
seborrheic dermatitis, 347
sleep disorders, 191, 192, 2618
deaths from, 6e-1t
differential diagnosis of, 2609–2611, 2610t,
 2612t
ECG in, 278e-5f
in elderly, 82
etiology of, 2611–2613
falls in, 163
familial, 444e-9, 2611–2612
frailty and, 77, 82
genetic considerations in, 444e-9–10, 2611–
 2612, 2612t
imaging in, 2609, 2611f
pathophysiology of, 2609, 2610f, 2611f, 2612f,
 2613, 2613f
prevalence of, 2609
prions in, 453e-1, 2612
- vs. progressive supranuclear palsy, 2605
treatment of, 2613
 amantadine, 2616
 anticholinergics, 2615
 approach to, 2618, 2619f
 COMT inhibitors, 2615, 2616t
 deep brain stimulation, 2616, 2618
 dopamine agonists, 2615, 2616t, 2618
 experimental, 2617
 gene delivery, 2617
 levodopa, 2613–2614, 2614f, 2616t, 2618
 MAO-B inhibitors, 2615, 2616t, 2618
 neuroprotective agents, 2616
 for nonmotor and nondopaminergic
 features, 2617–2618
 nonpharmacologic, 2618
 stem cell therapy, 90e-3, 444e-7, 2617
 surgical, 2616–2617
- Parkinson's disease dementia (PDD), 171, 2605–
 2606, 2617
- Paromomycin
action of, 246e-11
adverse effects of, 246e-2t, 246e-11, 942, 1391
for amebiasis, 246e-9, 1366
for cutaneous leishmaniasis, 1393
for *D. fragilis* infections, 1409
for *Giardia* infections, 1406
indications for, 246e-2t
pharmacology of, 246e-11
in pregnancy and lactation, 246e-2t
for visceral leishmaniasis, 1391
- Paronychia, 351
Candida, 1343
drug-induced, 380, 380f
P. aeruginosa, 1047
- Parotid gland
bilateral enlargement of, 2167t
tumors, 503
- Parotitis
acute bacterial, 241
differential diagnosis of, 231e-3
in mumps infection, 231e-2, 231e-2f
- Paroxetine
adverse effects of, 2392, 2710t, 2716
for anxiety disorders, 2711
for depression, 64, 473, 2710t, 2716
drug interactions of, 34t, 40
for hot flashes, 623
for irritable bowel syndrome, 1969
for menopausal symptoms, 2383
metabolism of, 34t
for panic disorder, 2709
pharmacology of, 2717
for PTSD, 2712
- Paroxyssmal AV block, 1473, 1473f
- Paroxyssmal cold hemoglobinuria, 657t, 659
- Paroxyssmal dyskinesias, 2624–2625
- Paroxyssmal hemiparesis, 107, 109t, 2595t, 2596
- Paroxyssmal junctional reciprocating tachycardia,
1481
- Paroxyssmal kinesigenic dyskinesia (PKD),
2624–2625
- Paroxyssmal nocturnal dyspnea, 1503–1504
- Paroxyssmal nocturnal hemoglobinuria, 660
clinical features of, 282, 399, 660, 660f, 664
diagnosis of, 660, 660f
natural history of, 660
pathophysiology of, 660, 661f
treatment of, 661–662, 661f
- Paroxyssmal nonkinesigenic dyskinesia (PNKD),
2625
- Paroxyssmal supraventricular tachycardia
(PSVT). *See also* Supraventricular
 tachyarrhythmias
accessory pathways and, 1481
AV nodal reentry tachycardia, 278e-7f, 1478f,
 1479–1480, 1481f
AV reentry tachycardia, 1478f, 1481
mechanisms of, 1478f
orthodromic AV reentry tachycardia, 1481
preexcited tachycardias, 1482, 1483f
treatment of, 1483–1484, 1483f
vs. ventricular tachycardia, 1494
PARP inhibition, 102e-6f, 102e-7, 529

- Parrotfish poisoning, 2742
 Parrot's pseudoparalysis, 836
 Parsonage-Turner syndrome, 2692
 Pars planitis, 200
 Partial liquid ventilation, 1738–1739, 1742
 Partial response, to chemotherapy, 103e-5
 Partner notification, 883
 Parulis, 237
 PARV4/5 viral sequences, 1197
 Parvovirus(es), 214e-2t, 214e-3f, 214e-4, 1195
 Parvovirus B19 infections, **1195**
 aplastic anemia and, 664
 clinical features of, 1195f, 1196
 diagnosis of, 1196–1197, 1196t
 epidemiology of, 1195
 erythema infectiosum. *See* Erythema infectiosum
 hydrops fetalis. *See* Hydrops fetalis
 myocarditis, 1560
 pathogenesis of, 1195–1196, 1195f
 in pregnancy, 50, 1196
 prevention of, 1197
 pure red cell aplasia, 668, 668f, 1196. *See also* Pure red cell aplasia
 transfusion-transmitted, 138e-6
 in transplant recipient, 923
 treatment of, 669, 1197
 Parvus et tardus pulse, 51e-3, 1448
 Pasireotide, 567, 2272–2273
 Passive euthanasia, 67t
 Passive transport, 332e-6
 Pasteur effect, 247
Pasteurella spp. infections
 in cat-bite wound, 167e-1, 183e-3, 830
 cellulitis, 830
 clinical features of, 183e-3, 834
 in dog-bite wound, 167e-1, 183e-3
 treatment of, 167e-3, 167e-3t, 183e-2t, 183e-3, 830
 Pastia's lines, 134, 966, 966f
 Patch (skin lesion), 339t
 Patched gene, 387
 Patch testing, 344
 Patellar tendinitis, 2220–2221
 Patent ductus arteriosus (PDA)
 clinical features of, 1523, 1524f
 heart sounds/murmur in, 1449
 location of, 1521f, 1524f
 pathogenesis of, 1523
 treatment of, 1523
 Patent foramen ovale, 296e-4, 476e-3, 2566t
 Paternal age, Marfan syndrome and, 433, 438
 Paterson-Kelly syndrome, 532
 Pathogen-associated molecular patterns (PAMPs), 145e-5, 145e-8f, 372e-2, 377e-1, 1747
 Patient
 expectations of, 6–7
 hospital experience of, 5
 Internet use by, 6
 principles of patient care, 3–4
 Patient autonomy, 17e-2
 Patient-centeredness, 16e-6
 Patient-controlled analgesia, 92
 Patient Health Questionnaire-2 (PHQ-2)
 Depression Screen, 471e-5t
 Patient-physician relationship. *See* Physician-patient relationship
 Patient satisfaction, 16e-4–5
 Patient-ventilator synchrony, 1743
 Pattern recognition, 18–19
 Pattern-recognition molecules (collectins), 145e-5
 Pattern recognition receptors (PRRs). *See also* Toll-like receptors (TLRs)
 in adaptive immune response modulation, 372e-6t
 in autoimmunity, 377e-1–2
 definition of, 372e-2
 in innate immune system, 372e-2, 372e-4, 372e-4t
 ligands of, 372e-5t
 signaling pathways of, 372e-5f
 PAX3 gene mutations, 101e-3t, 359, 431t
 PAX7 gene mutations, 101e-3t
 PAX8 gene mutations, 2283, 2284t, 2304, 2305t
 Pay-for-performance system, 12e-4
 Pazopanib
 action and targets of, 102e-17, 102e-18t, 103e-19
 adverse effects of, 102e-18t, 103e-26
 for renal cell carcinoma, 579
 for soft tissue sarcoma, 119e-2
 P blood group system, 138e-2
 PC-1 gene mutations, 415e-4t, 2364t
 PCI. *See* Percutaneous coronary interventions (PCI)
 PCNSL. *See* Primary central nervous system lymphoma (PCNSL)
 PCOS. *See* Polycystic ovarian syndrome (PCOS)
 PCP. *See* Phencyclidine (PCP); *Pneumocystis pneumonia* (PCP)
 PC-PTSD (Primary Care PTSD) Screen, 471e-4
 PCR. *See* Polymerase chain reaction (PCR)
 PCSK9 deficiency, 2445
 PCSK9 inhibitors, 2442
 PCSK9 mutations, 2442, 2445
 PCT (procalcitonin), 806
 PCV (pressure-control ventilation), 1742. *See also* Mechanical ventilation
 PD. *See* Parkinson's disease (PD)
 PD-1 antibodies, 498, 499, 499f, 521
 PDA. *See* Patent ductus arteriosus (PDA)
 PDC (pyruvate dehydrogenase complex), 372e-24t, 2060
 PDD (Parkinson's disease dementia), 171, 2605–2606
 PDGF. *See* Platelet-derived growth factor (PDGF)
 PDGFR gene mutations, 135e-7, 135e-7t
 PDGFR-mutated eosinophilia, 135e-8
 PDK (pyruvate dehydrogenase kinase), 102e-13
 PD-L1, 103e-21, 103e-22f
 PE. *See* Pulmonary embolism (PE)
 Peak oxygen consumption, 73, 74f
 Pearson's syndrome, 85e-6, 85e-6t
 Peau d'orange skin, 967, 1654
 Pectoriloquy, 1662
 Pectus carinatum (pigeon chest), 1443, 2512
 Pectus excavatum (funnel chest), 1443, 2512
 Pedal edema, 1662
 Pediculiasis, 2745–2746
 Pediculicidosis, 2746
Pediculus capitatus (head lice), 2745–2746, 2746f
Pediculus humanus (body lice), 1146, 2745–2746. *See also* Epidemic (louse-borne) typhus
 Pedigree analysis
 in family history, 443
 in genetic counseling, 446, 447f
 in Mendelian disorders, 437–438, 438f
 in mitochondrial DNA diseases, 85e-1, 85e-2f
 standard symbols for, 438f
 Peduncular hallucinosis, 2575
 PEEP (positive end-expiratory pressure), 1738. *See also* Mechanical ventilation
 Pefloxacin, 379
 PEG-ADA, 91e-2
 Pegfilgrastim, 103e-24, 103e-24t
 PEG IFN- α 2a. *See* Pegylated interferon- α 2a (PEG IFN- α 2a)
 PEG IFN- α 2b. *See* Pegylated interferon- α 2b (PEG IFN- α 2b)
 Pegloticase, 2235
 Pegvisomant, 2271
 Pegylated interferon- α 2a (PEG IFN- α 2a)
 for chronic HBV infection, 215e-4t, 215e-10, 2035–2036, 2037t, 2040t
 for chronic HCV infection, 215e-4t, 2043–2044, 2043t, 2045–2046t, 2048
 for chronic HDV infection, 215e-5t, 2040
 Pegylated interferon- α 2b (PEG IFN- α 2b)
 adverse effects of, 498, 2060
 for chronic HBV infection, 2035–2036, 2037t, 2038f, 2039, 2040t
 for chronic HCV infection, 215e-4t, 2043–2044, 2043t, 2045–2046t, 2048
 with ribavirin, 215e-4t, 215e-10, 2043–2044
 for chronic HDV infection, 215e-5t, 215e-10, 2040
 for melanoma, 498
 PELD system, in liver transplantation, 1995
 Pel-Ebstein fever, 126, 708
 Pelger-Huet anomaly, 81e-6f, 415, 415f
 Peliosis hepatitis, 2027
 Pelizaeus-Merzbacher disease, 2636
 Pellagra
 clinical features of, 96e-1t, 96e-4, 360, 463t, 2607, 2690
 “gauntlet” of, 436e-1f
 global considerations in, 2690
 vs. Hartnup disease, 435e-2
 risk factors for, 96e-4
 treatment of, 96e-4, 2690
 Pelvic abscess, 1099
 Pelvic floor dysfunction, 273–274
 Pelvic floor relaxation, 60e-3
 Pelvic floor retraining, 274
 Pelvic fractures, 2489
 Pelvic inflammatory disease (PID), **876**
 abdominal pain in, 103, 106
 anaerobic, 1099
 C. trachomatis, 1169
 clinical features of, 877
 diagnosis of, 877–878
 epidemiology of, 877
 etiology of, 870t, 877
 follow-up for, 879
 M. hominis, 1165
 pelvic pain in, 338, 338t
 prevention of, 879
 prognosis of, 879
 treatment of, 878–879, 878t
 Pelvic muscle exercises (Kegel exercises), 81
 Pelvic pain, **337**
 acute, 338
 approach to the patient, 338
 in cervical cancer, 595
 chronic, 338
 etiology of, 337–338, 338t
 in interstitial cystitis/bladder pain syndrome, 60e-1–2
 in pelvic inflammatory disease. *See* Pelvic inflammatory disease (PID)
 Pelvic thrombophlebitis, septic, 50
 Pemberton's sign, 2301, 2302
 Pembrolizumab, 102e-5t, 103e-21t, 103e-22
 Pemetrexed
 action of, 103e-14
 adverse effects of, 103e-11t, 103e-14
 for lung cancer, 520, 520t, 521
 for ovarian cancer, 594
 Pemoline, 63
 Pemphigoid
 bullous, **372**
 clinical features of, 76e-15f, 370t, 372, 372d
 histology of, 370t, 372
 immunopathology of, 370t, 372, 372e-24t
 treatment of, 372
 esophageal involvement/dysphagia in, 257
 gestational, 361t, 370t, 372–373
 mucous membrane, 237, 239t, 370t, 372e-24t, 374
 Pemphigus
 drug-induced, 379
 paraneoplastic, 361, 361t, 370t, **372**
 Pemphigus foliaceus, **371**
 clinical features of, 370t, 371
 drug-induced, 371–372
 histology of, 370t
 immunopathology of, 370t, 371, 372e-24t
 treatment of, 372
 Pemphigus vulgaris, **370**
 clinical features of, 76e-15f, 370, 370t, 371f
 diagnosis of, 371
 histology of, 370t
 HLA gene association with, 373e-7, 373e-8t
 immunopathology of, 370–371, 370t, 372e-24t, 377e-4

- oral manifestations of, 237, 239t
treatment of, 371
- Penciclovir, 215e-3t, 215e-8, 233, 1182t
- Pendred's syndrome, 220, 222t, 2284, 2284t
- Pendrin, 2284, 2284t
- Penetrance, 437, 446
- Penetrating radiation, 263e-1
- D-Penicillamine
adverse effects of
 autoimmune myasthenia gravis, 2702–2703
 cutaneous, 349, 371, 379
 lupus syndrome, 2218t
 myopathy, 462e-20, 462e-20t, 2199, 2218t
 optic neuropathy, 203
 pyridoxine and, 96e-4
 vasculitis, 2218t
 zinc deficiency, 96e-10
for cystinuria, 435e-2
for mercury poisoning, 472e-3t
for systemic sclerosis, 2164
for Wilson's disease, 367e-2, 2519, 2520t, 2625
- Penicillin(s)
for actinomycosis, 1090–1091, 1091t
action of, 930, 931t, 937
adverse effects of
 acid-base disorders, 322
 cross-sensitivity and, 385
 cutaneous immunologic, 377
 desensitization to, 385
 erythema nodosum, 366
 erythroderma, 353
 hemolytic anemia, 44
 hypokalemia, 306
 neutropenia, 418
 pemphigus, 371
 platelet dysfunction, 730
 rash, 362
 seizure, 2547t, 2548
 toxic epidermal necrolysis, 361
 vasculitis, 2193
for anaerobic bacterial infections, 1101
for anthrax, 261e-3
for asymptomatic pharyngeal GAS carriage, 966
for bite-wound infections, 167e-3, 167e-3t
for *Capnocytophaga* infections, 183e-2t, 183e-3
for clostridial myonecrosis, 780t
drug interactions of, 42–43
for enterococcal infections, 975t
for erysipelas, 965t
for gas gangrene, 832t
for GAS infections, 965t
for group B *Streptococcus* infections, 970
for group C or G streptococcal infections, 969
for HACEK infections, 183e-2t
for infective endocarditis, 821, 822t, 823
for laryngitis, 234
for leptospirosis, 1145t
for meningococcemia, 780t
for necrotizing fasciitis, 780t, 965t
for pharyngitis, 232f, 233, 965t
for pneumonia, 965t
in pregnancy and lactation, 937t
resistance to
 in anaerobic bacteria, 1101
 mechanisms of, 931t, 933
 in meningococci, 1000
 in *N. gonorrhoeae*, 1004
 in pneumococci, 807
for rheumatic fever, 2152
for tetanus, 986
for toxic shock syndrome, 965t
- Penicillin-binding proteins, 930, 947f, 976
- Penicillin G
for diphtheria, 979
for group B *Streptococcus* infections, 50, 970
indications for, 937, 940t
for Lyme disease, 1153, 1153f
for meningitis, 887t, 889, 889t
for neurosyphilis, 1139, 1139t
for nongonococcal bacterial arthritis, 835
for osteomyelitis, 840t, 841t
for pharyngitis, 965t
for pneumococcal infections, 952
for relapsing fever, 1148, 1148f
resistance to, 940t
for staphylococcal infections, 962t
for syphilis, 1138–1139, 1139t
for syphilitic meningitis, 899
- Penicillin V, for diphtheria, 979
- Penicillium marneffeii* infection (penicilliosis), 1252t, 1268, 1354–1355, 1354t
- Penile cancer, 351, 1198
- Penile prosthesis, 329
- Penile tumescence, 324
- Penile warts, 1198, 1199f
- Pentamidine
action of, 246e-11, 332e-9
adverse effects of
 acid-base disorders, 321
 bronchospasm, 1362t
 frequent, 246e-6t, 1398
 hyperkalemia, 64e-2, 310
 hypotension, 1361
 hypovolemia, 297
 nephrotoxicity, 1804
 occasional, 246e-6t, 1398
 pancreatic, 1258
 rare, 246e-6t
 SA node dysfunction, 1467t
for PCP, 1361t
for PCP prophylaxis, 1251t, 1253, 1362t
pharmacology of, 246e-11
in pregnancy and lactation, 246e-6t
for sleeping sickness, 1398, 1398t
- Pentasa, 1959
- Pentastomiasis, 2747
- Pentavalent antimonials, 246e-2t, 246e-7, 1390, 1393
- Pentazocine, 462e-20, 2218t
- Pentobarbital, 473e-10t, 2558f
- Pentosan polysulfate, 60e-3–4
- Pentoxifylline
for alcoholic liver disease, 2053, 2054f, 2059
for intermittent claudication, 1645
- Penumbra, 1777
- PEP. See Postexposure prophylaxis (PEP)
- PEPFAR (U.S. President's Emergency Plan for AIDS Relief), 8
- Pepsin, 1914
- Peptic ulcer disease, **1911**
clinical features of, 96t, 100, 106t, 262, 1918
complications of
 bleeding
 endoscopic treatment of, 1886, 1890, 1890f
 incidence of, 1918
 recurrent, prevention of, 276–277, 346e-1v, 1890, 1890f
 gastric outlet obstruction, 1918
 perforation, 1918
 conditions related to
 gastritis. See Gastritis
 Ménétrier's disease, 534, **1932**
 stress-related mucosal injury, 277, 1734, 1743, 1929–1930
 Zollinger-Ellison syndrome. See Zollinger-Ellison syndrome (ZES)
deaths from, 1911
diagnosis of, 1918–1919
 barium studies, 1919, 1919f
 endoscopic, 1881, 1881f, 1886, 1895, 1919, 1919f
 H. pylori detection, 1919, 1920t
differential diagnosis of, 1918
duodenal ulcers in. See Duodenal ulcers
epidemiology of, 1911, 1914
etiology of, 262, 1917–1918, 1918t
gastroduodenal mucosal defense and, 1912–1913, 1912–1914
genetic considerations in, 1917
H. pylori-related. See *Helicobacter pylori* infections, in peptic ulcer disease
NSAID-induced, **1916**
 epidemiology of, 1916–1917
 etiology of, 1917, 1917f
 prevention of, 1923–1924, 1924t
- pathology of, 1915
pathophysiology of, 1915
in polycythemia vera, 673
recurrence of, 1926
risk factors for, 1914
stomach cancer and, 534
treatment of, **1919**, 1920t
 antacids, 1920–1921
 approach to, 1924, 1924f
 bismuth-containing preparations, 1921
 H. pylori eradication. See *Helicobacter pylori* infections, treatment of
 H₂ receptor antagonists, 1920
 NSAID-induced, 277, 1923–1924, 1923t
 PPIs, 1920–1921
 prostaglandin analogues, 1921
 in refractory disease, 1924
 sucralfate, 1921
 surgical, 1925–1927, 1925f
 ZES and, 568, 1927. See also Zollinger-Ellison syndrome (ZES)
- Peptide(s)
antimicrobial, 372e-1, 372e-2, 372e-4t
ovarian, 2378
- Peptide antigen, in HLA class I molecule, 373e-4
- Peptide nanostructures, 92e-1
- Peptide radioreceptor therapy (PRRT), 575
- Peptide YY, 415e-2
- Peptidoglycan, 145e-5, 145e-6f, 930, 1752
- Peptostreptococcus* spp. infections, 167e-2, 1095, 1099, 1100. See also Anaerobic bacterial infections, mixed
- Peramivir, 215e-5, 923, 1213
- Percent transferrin saturation, 627, 627f, 630t
- Perchlorate, 2300
- Percussion
of chest, 47e-3, 1662
of spleen, 411
- Percutaneous coronary interventions (PCI), **296e-1**, **297e-1**
for adult congenital heart disease, 296e-4
antiplatelet agents in, 296e-1, 296e-2, 1596
for aortic valve replacement. See Aortic valve replacement, transcatheter
aortic valvuloplasty, 1533
atherectomy, 296e-2
for atrial septic defect repair. See Atrial septal defect (ASD), treatment of
balloon angioplasty and stenting, 296e-1–2, 296e-1f
for cardiogenic shock, 1762
circulatory support during, 296e-5
complications of, 296e-2–3, 1602t
distal protection device for, 296e-2, 297e-3, 297e-3f, 297e-3v
embolic protection device for, 296e-2
Impella device for, 296e-5
indications and patient selection for, 296e-3–4
intimal dissection in, 296e-1, 296e-2f
intraaortic balloon for circulatory support in, 296e-5
vs. medical therapy in acute coronary syndromes, 296e-4
mitral balloon valvotomy, 1542, 1542f
mitral valve repair, 1546, 1546f
for myocardial infarction
 NSTEMI-ACS
 left main in high-risk patient, 297e-3, 297e-3v
 multivessel in diabetic patient, 297e-4, 297e-4v
 patient selection, 296e-3f, 296e-4
 rescue, 1605
 STEMI. See ST-segment elevation myocardial infarction (STEMI), treatment of
 time from symptom onset to, 1603, 1603f
 urgent, 1605
myocardial infarction related to, 296e-2
perioperative, 52f, 53
racial/ethnic disparities in referral for, 16e-2

- Percutaneous coronary interventions (PCI)
(*Cont.*):
for revascularization in stable ischemic heart disease, **1590**
of bifurcation lesions, 297e-1–2, 297e-1f, 297e-2–3v
vs. CABG, 296e-3–4, 296e-3f, 1592, 1593f
of chronic total occlusion, 297e-1, 297e-1v
efficacy of, 1591
indications and patient selection for, 1590–1591
restenosis after, 1591
risks of, 1591
saphenous vein graft with distal protection, 296e-2, 297e-3, 297e-3f, 297e-3v
stent thrombosis in proximal LAD lesion following, 297e-5, 297e-5f, 297e-5v
steerable guidewire in, 296e-1
success of, 296e-2–3, 296e-3f
techniques for, 296e-1–2
transluminal coronary angioplasty, 296e-1
for valvular heart disease, 296e-4–5
- Percutaneous interventions. *See also*
Thrombectomy, percutaneous
mechanical
for carotid atherosclerosis, 2571
coronary. *See* Percutaneous coronary interventions (PCI)
peripheral artery, 296e-5, 296e-5f, 296e-6f, 1645
for pulmonary embolism, 296e-5
for refractory hypertension, 296e-5
transluminal renal angioplasty, 1618
- Percutaneous needle aspiration, in pulmonary lesion, 1667
- Percutaneous transhepatic cholangiography, 2085t
- Percutaneous umbilical blood sampling (PUBS), 83e-4, 444
- Perennial nonallergic rhinitis, 2122
- Perennial nonallergic rhinitis with eosinophilia syndrome (NARES), 2122
- Perfringolysin, 993
- Perfusion magnetic resonance imaging, 440e-8
- Pergamino virus, 1308t
- Pergolide, 473e-8t, 565, 1650
- Perianal disease. *See* Anorectal disorders
- Periapical disease, 236, 1097
- Periappendicitis, 877
- Pericardial cysts, 1577
- Pericardial effusion
after cardiac injury, 289e-3
chronic, 1575, 1577
in Churg-Strauss syndrome, 386e-3f
ECG in, 1458, 1459f
echocardiography in, 270e-22–23, 270e-22f, 271e-8f, 271e-8v, 1572f
etiology of, 270e-22
malignant, 474, 1788–1789
in pericarditis, 1571–1572
physical examination in, 1450
tuberculous, 1577
- Pericardial friction rub
in myocardial infarction, 1600, 1610
in pericarditis, 1450, 1571
- Pericardial knock, 286
- Pericardial tamponade, 1450, 1750, 1788–1789
- Pericardiocentesis, 1574
- Pericarditis, **1571**
abdominal pain in, 105, 106t
acute
clinical features of, 1571–1572, 1580, 1600, 1610
CMR in, 271e-8f
differential diagnosis of, 1575
ECG in, 101, 269e-7f, 269e-8f, 1571, 1572f
echocardiography in, 1572, 1572f
postcardiac injury, 1575
treatment of, 1573
viral/idiopathic, 1574–1575
cardiac tamponade in. *See* Cardiac tamponade
chest pain in, 96t, 97, 99, 1571
in chronic kidney disease, 1575, 1817–1818
classification of, 1571t
in collagen vascular disease, 1575
constrictive
clinical features of, 1573t, 1576
CMR in, 270e-23f, 270e-27v, 271e-8f, 1576f
differential diagnosis of, 1462, 1463t, 1573t, 1574, 1576
dyspnea in, 47e-2
ECG in, 1573t, 1576
echocardiography in, 1573t, 1576
hemodynamic profile in, 1576
pathophysiology of, 1574f, 1575–1576
pulse pressure in, 1576
treatment of, 1576–1577
differential diagnosis of, 1575
etiology of, 1571t
infectious causes of, 1085, 1100, 1212, 1292, 1575
in myocardial infarction, 1610
in neoplastic disease, 1575
pericardial effusion in, 1571–1572, 1572f
pericardial friction rub in, 1450, 1571
radiation-induced, 103e-4, 621, 1575, 1788
in rheumatoid arthritis, 2138
SA node dysfunction in, 1467, 1467t
in SLE, 1575, 2129, 2129t
subacute effusive-constrictive, 1577
tuberculous, 1111–1112
- Pericardium
congenital defects of, 1571
cysts of, 1577
metastases to, 1577, 1788
structure and function of, 1571
tumors of, 1577
- Perichondritis, 228
- Pericytes, 265e-1, 265e-1f
- Perihepatitis (Fitz-Hugh-Curtis syndrome), 877, 1169
- Perilymphatic fistula, 220
- Perimenopause, 2381–2382, 2381f
- Perineal descent, 265f, 273–274
- Perinephric abscess, 851
- Perinuclear ANCA (pANCA), 2180
- Periodic acid–Schiff stain, 350
- Periodic hypothermia syndrome, 2264
- Periodic limb movement disorder, 192
- Periodic paralysis
channelopathies in, 444e-2t
hyperkalemic. *See* Hyperkalemic periodic paralysis (HyperKPP)
hypokalemic. *See* Hypokalemic periodic paralysis (HypoKPP)
thyrototoxic, 462e-16, 462e-18
- Periodontal disease (periodontitis)
abscess in, 1097
AIDS-related, 236
clinical features of, 46e-2f, 46e-5f, 236
in elderly, 242
etiology of, 236, 1097
localized juvenile, 236
prevention of, 236, 423
risk factors for, 236
- Periodontium, 236
- Periostitis, in lung cancer, 511
- Peripartum cardiomyopathy, 47, 1562
- Peripharyngeal space infections, 1097
- Peripheral artery disease, **1643**
cardiovascular disease and, 747, 1644
clinical features of, 1643
cyanosis in, 249
diagnosis of, 1446, 1446f, 1643, 1644f
hypertension and, 1616
pathophysiology of, 1643
preoperative risk assessment in, 1645
prognosis of, 1644
treatment of
exercise program, 1644
pharmacologic, 1644–1645
revascularization, 296e-5, 296e-6f, 1645
- Peripheral artery pulses, 1445
- Peripheral blood smear, **81e-1**. *See also specific disorders*
Howell-Jolly bodies in. *See* Howell-Jolly bodies
normal, 81e-2f, 81e-6f, 394f, 726f
red cell defects on, 394
red cell fragmentation in, 395f
reticulocytes in, 81e-2f, 396f
spur cells in, 396f
target cells in, 81e-4f, 395f
- Peripheral cyanosis, 249, 249t. *See also* Cyanosis
- Peripherally inserted central catheter, 98e-4, 98e-6
- Peripheral nerve(s)
components of, 87f
cutaneous fields of, 160f
pain mechanisms, 89–90
- Peripheral neuropathy, **2674**. *See also*
Plexopathies; Polyneuropathy
in alcohol use disorders, 2724
in amyloidosis, 2681–2682. *See also*
Amyloidosis
approach to, **2674**, 2675f
electrodiagnostic studies in, 2676–2677, 2677t
key questions in, 2674–2676, 2675t
laboratory studies in, 2677
nerve biopsy in, 2677
pattern recognition in, 2676t
skin biopsy in, 2677–2678
autonomic dysfunction and, 2642
in celiac disease, 2683
in chronic kidney disease, 1819, 2683–2684
in chronic liver disease, 2684
in CMV infections, 2685
critical illness polyneuropathy, 1755, 1783–1784, 2684
cryptogenic sensory and sensorimotor polyneuropathy, 2691
in diabetes mellitus. *See* Diabetic neuropathy
in diphtheria, 979, 2684
drug-induced, **2686**, 2687–2688t
amiodarone, 2687t, 2688
antiretrovirals, 2687t, 2689
cancer chemotherapy, 607, 607t, 622, 2686t
chloroquine, 2686, 2687t
colchicine, 2687t, 2688
hydroxychloroquine, 2686, 2687t
isoniazid, 2687t, 2689
pyridoxine, 2687t, 2688–2989
thalidomide, 2687t, 2688
in EBV infections, 2685
in hepatitis, 2685
hereditary, **2678**, 2679–2680t
adrenoleukodystrophy. *See*
Adrenoleukodystrophy
adrenomyeloneuropathy, 2680–2681
Charcot-Marie-Tooth disease. *See* Charcot-Marie-Tooth (CMT) disease
in Fabry's disease, 2680. *See also* Fabry's disease
familial amyloid polyneuropathy, 2681
hereditary neuralgic amyotrophy, 2678
hereditary neuropathy with liability to pressure palsies, 2678
hereditary sensory and autonomic neuropathy, 2680
in porphyria, 2681. *See also* Porphyria(s)
Refsum's disease, 206, 451e-4t, 2681
Tangier disease, 1443, 2445, 2681
in herpes zoster, 2685
in HIV infection, 1267, 2684–2685
in hypereosinophilic syndrome, 2683
in hypothyroidism, 2683
in IBD, 2683
in leprosy, 1124, 2684
in Lyme disease, 2633, 2684
in malabsorption syndromes, 1946t
malignancy-related, 614t, 618–619, 2685–2686
in mixed connective tissue disease, 2683
mononeuropathies, 463e-4–5, 463e-5f, 1124, 2691–2692
in nutritional deficiencies, 463t, 643, 2689–2691
- paraneoplastic, 614t, 618–619
in pregnancy, 49
in rheumatoid arthritis, 2683
in sarcoidosis, 2683
in Sjögren's syndrome, 2683
in SLE, 2683

- in systemic sclerosis, 2683
toxic, 2688t, 2689
weakness in, 156, 157, 2675, 2675t
- Peripheral node addressin (PNad), 372e-26
- Peripheral sensitization, 87
- Peripheral synovitis, 2169
- Peripheral T-cell lymphomas, 708
- Periplasmic space, 930
- Peristalsis, 255
- Peristaltic sounds, 106
- Perisylian network, 177
- Peritoneal carcinomatosis, 286, 287f, 288
- Peritoneal dialysis
access for, 1824-1825
complications of, 1825
continuous ambulatory, 848, 1811, 1824
continuous cyclic, 1824
dose for, 1825
for hypercalcemia, 2481
principles of, 1824
solutions for, 1824
- Peritoneal equilibrium test, 1825
- Peritoneal inflammation, 103
- Peritoneal lavage, as reawarming strategy, 478e-3
- Peritoneal papillary serous carcinoma, 120e-3-4
- Peritoneovenous shunt, 474
- Peritonitis, **846, 1989**
abdominal pain in, 103, 106, 106t, 846
acute, 1989
ascites and, 287-288, 2065-2066
aseptic, 1989
bacterial, 288, 846-847, 1099, 2065-2066
clinical features of, 288, 846-847, 1989
diagnosis of, 288, 847
etiology of, 846, 1028, 1989, 1989t
hypovolemia in, 298
pathophysiology of, 846, 846f
in peritoneal dialysis, 848, 1825
postoperative, 848
prevention of, 847
primary, 846-847, 1989
prognosis of, 1989
recurrence of, 847, 945t
secondary, 847-848, 1989
in transplant recipient, 924t, 927
treatment of, 288, 847, 848, 1989
tuberculous, 1111
tumor markers in, 473t
- Peritonsillar abscess, 233, 235, 1097
- Periventricular white matter disease, 2568
- Perkinsus marinus*, 151e-4
- Perleche, 342f, 351
- Permeability transition pore, 444e-9
- Permethrin, 2744-2745, 2746
- Pernicious anemia
APS-1 and, 2346t
APS-2 and, 2346t, 2347t
autoantibodies in, 372e-24t, 2347t
cobalamin absorption in, 350e-1, 350e-1t, 644-645
diagnosis of, 645
juvenile, 645
oral manifestations of, 237
- Pernio (chilblain), 478e-4, 1648f, **1649**
- Peroneal neuropathy, 463e-4, 463e-5f, 2683, 2692
- Peroxisomal catalase, 2058
- Peroxisome proliferator-activated receptor γ (PPAR γ), 96e-6, 415e-2, 1959, 2415
- Perphenazine, 2721, 2722t
- PERRLA, 195
- Persistent light reaction, 389
- Persistent müllerian duct syndrome, 2355
- Persistent vegetative state, 1771
- Personality disorders, **2720**
- Personalized medicine, 425
- Pertactin, 1021, 1022
- Perthes test, 1651f
- Pertussis, **1021**
clinical features of, 244, 1022-1023, 1023t
complications of, 1023
diagnosis of, 1023
differential diagnosis of, 1023-1024
epidemiology of, 1021-1022, 1021f, 1022f
global considerations in, 1021, 1021f
immune response in, 1022
infection control measures, 1024
pathogenesis of, 1022
prevention of, 1024
treatment of, 1024, 1024t
- Pertussis toxin, 1021, 1022
- Pertussis vaccine, 1022, 1024. *See also* Tetanus, diphtheria, acellular pertussis (Tdap) vaccine
- Pertuzumab
action and targets of, 102e-5t, 102e-6, 103e-21, 103e-21t
for breast cancer, 529
Perverse effects, 18
- PET. *See* Positron emission tomography (PET)
- Petechiae. *See also* Purpura
in anemia, 393
etiology of, 999t
in meningococemia, 781, 998
pathogenesis of, 368
in Rocky Mountain spotted fever, 1155, 1156f
in vitamin C deficiency, 463t
- Petrous bone fractures, 457e-2
- Peutz-Jeghers syndrome
lesions associated with, 538t
oral manifestations of, 240t
ovarian sex cord tumors in, 594
pancreatic cancer risk in, 554
polyp distribution in, 536, 538t
skin manifestations of, 359, 360t
- Peyer's patches, 372e-26-27, 1049, 2696f
- PF-4 (platelet factor 4), 372e-13t, 727, 750
- PFA-100, 405, 407
- Pfeiffer syndrome (acrocephalopolysyndactyly), 435
- PGA (polyglandular autoimmune) insufficiency, 2366, 2483
- PGC-1 α , 444e-10
- P gene mutations, 357
- PGL-1 (phenolic glycolipid), 1122, 1126
- P-glycoprotein (PGP), 33, 34t, 37f, 102e-10, 284
- pH
arterial, 315
esophageal, 262
urine, 1867
- Phaeohyphomycoses, 1355
- Phages, 86e-4
- Phagocyte(s). *See also* Neutrophil(s)
microbial survival inside, 145e-7
mononuclear. *See* Monocyte(s)
pathogen encounters with, 145e-5, 145e-6f
in primary immune deficiency diseases, 2104, 2105f
in vitro tests of, 423
- Phagocyte activation defects, 419t, 421-422
- Phagocytosis, 145e-5, 1105-1106
- Phagophobia, 254
- Phagosome, 145e-7
- Phalen's sign, 2221
- Phantom pain, oral, 237
- Pharmaceutical companies, gifts from, 17e-3-4
- Pharmacodynamics
of antibacterial drugs, 935, 935f
definition of, 31
principles of, 36
- Pharmacogenetics, 31, 425
candidate gene studies in, 38, 38f
in clinical practice, 41
of platelet inhibitors, 277, 277t, 742-743, 743t
- Pharmacogenomics
in clinical practice, 445, 451
definition of, 32, 427
genome-wide association studies in, 38-39, 39t
vascular, 265e-4-5
- Pharmacokinetics, 31, **33**
absorption of drugs, 33-34
of antibacterial drugs, 934-935, 935f
concepts, 34
drug distribution, 34-35
drug elimination, 35
drug metabolites, 35
high-risk, 35-36
interactions in, 41-43, 42t
- Pharmacology, **31**
clinical, 31
diseases affecting drug concentration and response, 37
genetic determinants of response to drugs, 38
global considerations in, 32
- Pharyngitis, **231**
arcanobacterial, 129t, 981
C. immitans, 981
C. ulcerans, 980
diagnosis of, 232-233, 232f
differential diagnosis of, 978
in diphtheria, 978, 978f
etiology of, 231, 965t, 1097
Fusobacterium, 231
in GERD, 262
gonococcal, 1006, 1008, 1009t
in HIV infection, 231
in infectious mononucleosis, 1187, 1187t
necrotizing, 1097
plague, 1072-1073
streptococcal
clinical features of, 231-232, 965
complications of, 233, 966
diagnosis of, 232-233, 232f, 965, 1190t
epidemiology of, 965
etiology of, 231
treatment of, 233, 965-966, 965t
viral, 965t
clinical features of, 231
coxsackievirus, 1292
etiology of, 231, 1202t
treatment of, 233
- Pharyngoconjunctival fever, 231, 1208
- Phenacetin, 477t, 575, 577, 1856t, 1861
- Phenazopyridine, 473e-11t, 867
- Phencyclidine (PCP)
actions of, 465e-5t, 469e-4
adverse effects of, 469e-4, 2199
overdosage/poisoning with, 469e-4, 473e-3, 473e-6
- Phendimetrazine, 2396
- Phenelzine
adverse effects of, 2593, 2710t
for depression, 2710t
for migraine prevention, 2593
for new daily persistent headache, 111
overdosage/poisoning with, 473e-8t
- Phenformin, 34t, 644t
- Phenobarbital
adverse effects of, 352, 2554t
drug interactions of, 43, 246e-3t, 246e-5t, 246e-6t, 2554t
for epilepsy, 2554t
overdosage/poisoning with, 473e-10t, 2765t
therapeutic monitoring of, 2765t
- Phenocopy, 40, 436
- Phenol exposure, 279
- Phenolic glycolipid (PGL-1), 1122, 1126
- Phenothiazines
adverse effects of
cardiac, 273e-4, 1467t
cutaneous, 349, 353, 379, 389t
heat-related illness, 479e-1
hypothermia, 478e-1
ocular, 206
orthostatic hypotension, 146
for dyspnea, 62
- Phenotoxicity, 91e-2t
- Phenotype, 435, 437
- Phenotypic heterogeneity, 435, 436t
- Phenotypic sex, 2349, 2352f
- Phenoxybenzamine
for bladder dysfunction, 2672
for frostbite, 478e-4t
for hypertension, 1624t, 2330-2331
- Phentermine, 2396, 2453
- Phentermine/topiramate (PHEN/TPM), 2396, 2397t, 2453
- Phentolamine
for adrenergic crisis, 1627
for high-altitude pulmonary edema, 476e-3
for hypertensive emergencies, 1627t
for Irukandji syndrome, 2739

- Phentolamine (*Cont.*):
 for nerve agent exposure, 262e-5t
 for sympathomimetic overdose/poisoning, 473e-8t
- Phenylacetate, 434e-5
- Phenylalanine metabolism disorders, 434e-2t
- Phenylbutazones, 42t, 678
- Phenylephrine
 for allergic rhinitis, 2122
 for neurogenic shock, 1750
 overdose/poisoning with, 473e-8t
- Phenylketonuria
 clinical features of, 357t, 434e-1, 434e-2t
 environmental factors in, 437
 maternal, 434e-1
 prevalence of, 434e-1
 screening for, 434e-1
 treatment of, 434e-1
- Phenylpropranolamine, 3, 473e-8t
- Phenytol
 for acroparesthesia, 432e-1
 adverse effects of
 cross-sensitivity and, 385
 cutaneous, 352, 353, 360, 361
 fever, 138
 folate deficiency, 646t, 647
 gingival hyperplasia, 236
 hepatotoxicity, 2023, 2029, 2682t
 hirsutism, 332
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 hypertrichosis, 380
 in kidney disease, 37
 long-term, 2552
 lupus syndrome, 2218t
 myopathy, 462e-20
 neurologic, 2554t, 2682t, 2687t
 osteopenia, 2218t
 systemic, 2554t
 thrombocytopenia, 727t
 vasculitis, 2193
- drug interactions of
 acetaminophen, 43
 azoles, 246e-3t
 cyclosporine, 42t
 fluoxetine, 2554t
 glipizide, 34t
 isoniazid, 205e-3
 losartan, 42, 42t
 methadone, 42t
 oral contraceptives, 42t
 PPIs, 1921
 quinidine, 42t, 246e-6t
 rufinamide, 2555t
 sulfonamides, 246e-4t, 2554t
 TMP-SMX, 939t
 warfarin, 42t
- for epilepsy, 2554t
 metabolism of, 34t, 40
 for myotonia, 462e-11
 for neuromyotonia, 619
 for neuropathy, 2682t
 overdose/poisoning with, 473e-10t, 2765t
- for pain, 91t, 94, 2672
 for paroxysmal symptoms in MS, 2672
 pharmacology of, 2552, 2554t
 for status epilepticus, 1735, 2558f
 therapeutic monitoring of, 2765t
 for trigeminal neuralgia, 2646
- Pheochromocytoma, **2329**
 clinical features of, **2329**
 cardiac, 254, 290e-1t, 290e-3, 1563
 by frequency of occurrence, 2330t
 hypertension, 1613, 1620t
 paraneoplastic syndromes, 610
 sympathetic overactivity, 2342, 2643
 definition of, 2329
 diagnosis of, 2329–2330, 2330t, 2341f, 2342
 differential diagnosis of, 2330
 epidemiology of, 2329
 etiology of, 2329
 familial, 1619, 2334
- genetic considerations in, **2329**
 mutation distribution, 2334–2335, 2334f
RET gene mutations, 1619, 1620t
 malignant, 2331
 in MEN syndromes, 1619, 1620t, 2332, 2332f, 2342
 in neurofibromatosis, 1619, 1620t, 2331–2332, 2331f, 2343
 in paraganglioma syndrome, 2332, 2333f
 pathogenesis of, 2329, 2330f
 during pregnancy, 2331
 renal cell carcinoma and, 578
 treatment of, 1619, 2330–2331
 tumor markers in, 473t
 in von Hippel-Lindau syndrome, 2332, 2333f
- Pheomelanin, 388
- PHEX* gene mutations, 2458
- Philadelphia chromosome
 in AML, 83e-1
 in CML, 101e-3, 101e-4f, 687–688, 688f, 689
 drug response and, 39t, 41
- Phlebectasia corona, 1651
- Phlebotomy
 for cyanotic CHD, 1523
 for erythrocytosis, 612
 for hemochromatosis, 367e-2, 2518
 for hemoglobinopathies, 637
 for polycythemia, 400, 674
 for porphyria cutanea tarda, 367e-3, 390, 2530
- Phleboviruses, 1305, 1309–1310t
- Phlebovirus infections
 Punta Toro virus, 1318
 Rift Valley fever. *See* Rift Valley fever
 sandfly fever, 1310t, 1318
 severe fever with thrombocytopenia syndrome, 1322
 Toscana virus, 894
- Phobic disorders, **2712**
- Phonemic paraphasia, 177
- Phoneutria* spp., 2748
- Phonic tics, 2623
- PhoP/PhoQ* genes, 145e-7
- Phosgene, 262e-1t
- Phosgene exposure, 1692t
- Phosgene oxime, 262e-1, 262e-1t
- Phosphate crystals, in urine, 62e-12f
- Phosphate nephropathy, 62e-11f, 1859
- Phosphate/phosphorus
 in chronic kidney disease, 333e-3, 333e-3f
 deficiency of. *See* Hypophosphatemia
 for diabetic ketoacidosis, 64e-2
 excess of. *See* Hyperphosphatemia
 for hypercalcemia, 610, 2480t, 2481
 metabolism of, 2458
 parenteral requirement for, 98e-5t
 phosphorus-32 exposure, 263e-3t
 recommended intake of, 95e-3t
 reference values for, 2762t
 renal tubular transport of, 332e-3f, 332e-7
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t
- Phosphatidylinositol-3-kinase (PI3K), 102e-6, 2403
- Phosphatonin, ectopic production of, 609t, 612
- Phosphodiesterase type 5 inhibitors
 action of, 324, 325f, 326f
 adverse effects of, 329t
 characteristics of, 329t
 contraindications to, 329t
 for erectile dysfunction, 324, 329t
 for female sexual dysfunction, 331
 for high-altitude pulmonary edema, 476e-3, 476e-4
 interaction with nitrates, 43, 328, 1596, 1603
 for lower urinary tract symptoms, 7e-4, 587
 for pulmonary arterial hypertension, 1659, 1660, 1660t, 2164
 for Raynaud's phenomenon, 1649, 2164
- Phosphofruktokinase deficiency (type VII
 glycogenosis; Tarui's disease), 433e-2t, 462e-14, 654t
- Phosphoglycerate dehydrogenase deficiency, 434e-2t
- Phosphoglycerate kinase deficiency, 433e-2t, 462e-14, 654t
- Phosphoglycerate mutase deficiency (type X
 GSD), 433e-2t, 462e-14
- Phospholamban, 265e-4, 265e-7, 265e-7f
- Phospholipase A₂, 126, 386
- Phospholipid transfer protein (PLTP), 2438
- Phosphoribosylpyrophosphate (PRPP), 431e-2
- Phosphoribosylpyrophosphate (PRPP)
 synthetase overactivity, 431e-3t, 431e-5
- Phosphorylate kinase deficiency (type IX
 glycogenosis), 332e-7t, 433e-2t, 433e-4, 462e-14
- Photoaging, 387
- Photoallergy, 389–390, 389t
- Photochemotherapy, 391
- Photodynamic therapy
 action of, 386
 for basal cell carcinoma, 501
 for cholangiocarcinoma, 553
- Photon, 103e-4
- Photopatch testing, 389
- Photoproducts, 386
- Photoprotection, 391, 391t. *See also* Sunscreen(s)
- Photoreceptors, 195
- Photosensitivity diseases, **388**
 classification of, 389t
 diagnosis of, 390f
 drug-induced, 379, 379f, 389–390, 389t
 management of, 390
 polymorphous light eruption, 364t, 366, 388–389
 porphyrias. *See* Porphyria(s)
 in SLE, 2129t
- Photosensitizer, 389
- Phototherapy. *See also* PUVA
 photochemotherapy
 action of, 391
 adverse effects of, 391
 for atopic dermatitis, 354t
 for cancer treatment of, 103e-5
 for morphea, 376
 for pityriasis rosea, 349
 for psoriasis, 348, 354t
 for psoriatic arthritis, 2177
 for shift-work disorder, 194
 for vitiligo, 358t
- Phototoxic eruptions, 361
- Phototoxicity, 389–390, 389t. *See also* Photosensitivity diseases
- PHP. *See* Pseudohypoparathyroidism (PHP)
- PHQ-2 (Patient Health Questionnaire-2)
 Depression Screen, 471e-5t
- Phrygian cap, 2076
- Phthalocyanines, 103e-5
- Phthisis. *See* Tuberculosis (TB)
- Phylloquinone, 96e-8. *See also* Vitamin K
- Phylogenetic analysis, 86e-1t
- Phylogenetic diversity, 86e-4–5
- Phylogenetic tree, 86e-1t
- Phylotype, 86e-1t, 86e-2
- Physical activity
 energy required for, 95e-1. *See also* Exercise
 for lifespan enhancement, 94e-7
- Physical examination
 clinical skills, 1–2
 screening tests, 26–27, 29t
- Physical restraints, in delirium, 65, 166, 170
- Physical therapy
 for back pain, 120
 for vertigo, 151t
- Physician
 brain drain, 13–14
 as citizen, 7
 claims of conscience by, 17e-2
 conflicts of interest for, 17e-3–4
 continuing education for, 7
 errors of, 4, 12e-1, 17e-4
 humanistic qualities of, 5
 impaired, 17e-4
 moral distress of, 17e-2–3
 oaths and codes of, 17e-2
 occupational risks of, 17e-4
 personal values of, 17e-2–3

- research and teaching by, 7
social media use by, 17e-3
as student, 7
trainee, ethical issues for, 17e-4
virtue ethics and, 17e-2
working hours of, 7, 12e-1, 17e-3
- Physician-assisted suicide, 67–68, 67t
- Physician-induced demand, 21
- Physician Orders for Life-Sustaining Treatment (POLST), 58
- Physician-patient relationship, 1, 4–5
racial/ethnic factors in communication, 16e-3–4, 16e-4f
trust within, 16e-4–5
- Physiology, 87e-1
- Physostigmine
for anticholinergic overdose, 473e-5, 473e-8t, 1776
for belladonna alkaloid overdose, 473e-8t
for mushroom/plant poisoning, 473e-9t
overdosage/poisoning with, 473e-10t
- Phytohemagglutinin (PHA), 83e-1
- PI3K (phosphatidylinositol-3-kinase), 102e-6
- PI3K/AKT pathway
in cancer, 102e-10
in endometrial cancer, 596
in head and neck cancer, 503
in melanoma, 496, 496f
- PI3P, 1106
- Pian. *See* Yaws
- Piano-key movement, ulnar styloid, 35
- PIC3CD gene, 2111
- Pica, 2719
- Pick's disease, 444e-8f, 2604, 2604f, 2605f
- Picornaviruses, 1286f, 1289
- Picornaviruses
gene expression and replication in, 214e-4
human respiratory syncytial virus, 1205. *See also* Human respiratory syncytial virus (HRSV) infections
rhinoviruses, 1202–1203. *See also* Rhinovirus infections
structure of, 214e-2t, 214e-3f
- “Picture frame” vertebra, 426e-2
- PID. *See* Pelvic inflammatory disease (PID)
- Piebaldism, 357t, 358t, 359
- PIEZO1 gene mutations, 653, 653t
- PIG-A gene mutations, 664
- Pigbel (enteritis necroticans), 991
- Pigeon chest (pectus carinatum), 1443, 2512
- Pigmented pretibial papules, 2429
- Pigmented villonodular synovitis (PVNS), 2246, 2247f
- Pigment stones, 2078, 2078t
- PIK3A gene mutations, 101e-2t
- PIK3CA gene mutations
in Cowden's syndrome, 2344
in lung cancer, 507t, 509, 509t
- Pili (fimbriae), bacterial, 145e-2, 145e-2t
of *B. Pertussis*, 1021
of *E. coli*, 145e-2, 145e-2f
of enterococci, 972
of *N. gonorrhoeae*, 1003
of *P. aeruginosa*, 1043, 1043t
of *S. pneumoniae*, 947, 947f
type IV, 145e-2, 145e-8
- Pill esophagitis, 257, 1910
- Pilocarpine
for acute angle-closure glaucoma, 200
in diagnosis of tonic pupil, 197
overdosage/poisoning with, 473e-10t
for xerostomia, 241
- Pilosebaceous unit, 331
- Pimecrolimus, 345, 354t
- Pimozide, 246e-3t, 2751
- Pinch purpura, 365
- Pindolol, 473e-9t, 1589t
- Pineal region tumors, 602
- Pinguecula, 199
- PINK1 gene mutations, 444e-9, 2092, 2613
- “Pink puffers,” 1704
- Pinna. *See* Ear
- Pinta, 207e-2–3, 207e-2f, 207e-2t
- Pintides, 207e-3
- Pinworm. *See* *Baylisascaris procyonis* infections; *Enterobius vermicularis*/enterobiasis (pinworm)
- Pioglitazone, 2057, 2414t, 2415
- Piperacillin
adverse effects of, 727t
for burn-wound infections, 166e-3
indications for, 938
for otitis externa, 229
resistance to, 938
- Piperacillin-tazobactam
for actinomycosis, 1091t
for anaerobic bacterial infections, 1101t
for community-acquired pneumonia, 808t
for febrile neutropenia, 491
for gram-negative bacterial infection, 1026
for health care-associated pneumonia, 767t, 812t
indications for, 938–939, 940t
for intraabdominal infections, 767t
for osteomyelitis, 840t, 841t
for *P. aeruginosa* infections, 808t, 1045t
for peritonitis, 847
resistance to, 940t, 1101
for sepsis/septic shock, 1757t
for septic shock, 767t, 780t
- Piperazine, 246e-6t, 246e-11–12
- Piracetam, 2623
- Piranha bite-wound, 167e-1
- Pirfenidone, 2164
- Piroxicam
adverse effects of, 356, 389t
for colorectal cancer prevention, 477
drug interactions of, 43
for paroxysmal hemichorea, 2596
- Piry virus, 1310t
- Pit-1
in anterior pituitary development, 401e-1, 401e-1t, 2256
in congenital hypothyroidism, 2284t
in growth hormone deficiency, 2258
- Pitavastatin, 2448t
- Pitting, 410
- Pitting edema, 251. *See also* Edema
- Pituitary apoplexy, 207, 2257, 2325t
- Pituitary axes, 401e-2f
- Pituitary disorders
anterior
insufficiency. *See* Hypopituitarism
tumors. *See* Pituitary tumors (adenomas)
- IgG4-related, 391e-1t
- posterior
diabetes insipidus. *See* Diabetes insipidus (DI)
hypodipsic hypernatremia. *See* Hypodipsic hypernatremia
- Pituitary gland
anatomy of, 401e-1, 401e-2
development of, 401e-1, 401e-1t
dysplasia, 2255–2256
hormones produced by. *See* Pituitary hormones; *specific hormones*
insufficiency of. *See* Hypopituitarism
tests of function of, 2258t
- Pituitary hormones, 401e-1. *See also specific types*
expression and regulation of, 401e-1t, 2274
in ovarian function regulation, 2376–2377, 2377f
secretion of, 401e-1, 401e-2f
- Pituitary snuff taker's lung, 1681t
- Pituitary tumors (adenomas), 2265
ACTH-producing. *See* Cushing's syndrome
classification of, 603, 2265, 2265t
clinical features of, 2261–2262, 2261t, 2339
etiology of, 2265
familial syndromes of, 2265–2266, 2265t, 2340
gonadotropin-producing, 2273–2274
histologic evaluation in, 2262
hormone-excess syndromes due to, 2265
hypogonadotropic hypogonadism and, 2365
laboratory investigation of, 2262, 2262t
local mass effects of, 2261–2263
- in MEN 1, 2265–2266, 2265t, 2336t, 2339
MRI in, 2262, 2262f
nonfunctioning, 603, 2273–2274, 2274f
ocular signs in, 198, 207
ophthalmologic evaluation in, 2262t
pathogenesis of, 2265
secondary adrenal insufficiency and, 2325t
treatment of, 603, 2262–2264, 2263f, 2339
TSH-secreting, 2274, 2298
- Pit viper bite, 2734, 2734f, 2736t, 2738
- Pityriasis rosea, 349
clinical features of, 347t
distribution, 341t, 342f, 347t
erythroderma, 353t
lesion characteristics, 349
morphology, 341t
papulosquamous lesions, 353
patches on trunk, 76e-3f, 349f
differential diagnosis of, 134
histologic features of, 347t
treatment of, 349
- Pityriasis rubra pilaris, 354t
- Pivmecillinam, 866, 867t, 1058t
- Pizotifen, 2594t
- PKAN (pantothenate kinase-associated neurodegeneration), 2626
- PKD (paroxysmal kinesigenic dyskinesia), 2624–2625
- PKD-1 gene mutations, 1850, 1851
- PKD-2 gene mutations, 1850, 1851
- PKHD1 gene mutations, 1853
- Placebo effect, 89, 90f
- Placental stem cells, 92e-1
- Plague, 1070. *See also* *Yersinia pestis*
in bioterrorism, 261e-3–4
bubonic, 261e-4, 1072, 1072f
case definitions of, 1073t
clinical features of, 261e-4–5, 261e-5t
diagnosis of, 150e-8, 261e-5t, 1073–1074, 1074f
epidemiology of, 151e-8f, 1070–1071, 1071f
global considerations in, 1070–1071, 1071f, 1073
meningeal, 1072
multidrug-resistant, 261e-6
pathogenesis of, 1071–1072
pharyngitis, 1072–1073
pneumonic, 261e-4, 261e-5t, 1072, 1073f
prevention of, 261e-5t, 261e-6, 1075–1076, 1075t
septicemic, 261e-4–5, 781, 1072
treatment of, 261e-5–6, 261e-5t, 1074–1075, 1074t
- Plague vaccine, 261e-6, 1075–1076
- PLAID (PLCγ-associated antibody deficiency and immune dysregulation), 2113
- Plan-Do-Check-Act cycle, 12e-3f
- Planktonic cells, 145e-4
- Plantar fasciitis, 2249–2250
- Plantar reflex, 2538
- Plantar warts, 351
- Plaque
atherosclerotic. *See* Atherosclerosis, pathogenesis of
dental, 46e-4f, 236, 1097
neuritic, in AD, 2599, 2600f
- Plaque (skin lesion), 127, 339t
- Plaques jaunes, 457e-1
- Plasma
derivatives, 138e-2, 138e-3
fresh-frozen. *See* Fresh-frozen plasma
O₂ concentration, 1748t
platelet-rich, 138e-2
- Plasma cell(s), 81e-11f
- Plasma cell disorders, 710
acid-base disorders in, 317
amyloidosis. *See* Amyloidosis
genetic considerations in, 710, 711f
heavy chain disease, 718–719
multiple myeloma. *See* Multiple myeloma
paraproteinemia in, 404
pathophysiology of, 710–712, 712f
proteinuria in, 293
Waldenström's macroglobulinemia. *See* Waldenström's macroglobulinemia

- Plasma cell leukemia, 715
 Plasmacytic lymphoma of the oral cavity, 1270
 Plasmacytoid T dendritic cells, 372e-6, 372e-7t.
See also Dendritic cells
 Plasmacytoma, 715, 715t
 Plasma exchange/plasmapheresis
 for cerebellar degeneration, 618
 for chronic inflammatory demyelinating polyneuropathy, 2699
 for Guillain-Barré syndrome, 2697
 for hemolytic-uremic syndrome, 1864
 for monoclonal gammopathy of undetermined significance, 716
 for MS, 2667
 for myasthenia gravis, 2705
 for neuromyelitis optica, 2673
 for neuromyotonia, 619
 for pemphigus vulgaris, 371
 for thrombotic thrombocytopenic purpura, 729-730, 1864
 for Waldenström's macroglobulinemia, 718
 Plasma protein binding, 35
 Plasma renin activity, 306
 Plasma vasopressin, in diabetes insipidus, 295
 Plasmin, 402f, 403, 758, 758f
 Plasminogen, 402, 402f, 758, 758f
 Plasminogen activator, 758, 758f
 Plasminogen activator inhibitor I (PAI-1)
 arterial thrombotic disease and, 744
 in endothelial cells, 265e-2
 in fibrinolysis, 758, 758f
 in metabolic syndrome, 744
 Plasminogen activator inhibitors, 402f, 403
Plasmodium spp. *See also* Malaria
 characteristics of, 250e-1, 250e-1t, 1369t
 drug resistance in, 1370f, 1379, 1381f, 1384
 entry into host, 145e-1
 epidemiology of, 245e-4t
 host response to, 1371-1672
 laboratory identification of
 blood findings, 1375-1377, 1377t
 demonstration of parasite, 245e-4t, 245e-6t, 250e-2-4f, 1375-1377f, 1375-1378, 1377t
 PCR, 245e-3
 life-cycle hosts of, 245e-4t
P. falciparum
 adherence of, 145e-2t
 characteristics of, 250e-1t, 1369, 1369t
 erythrocyte changes caused by, 1371
 geographic distribution of, 1370-1371
 laboratory identification of, 250e-2f, 250e-3f, 1375f, 1376, 1376f, 1377t
 ligand-receptor interactions of, 1371
 temperature sensitivity of, 1371
P. knowlesi
 characteristics of, 1369
 geographic distribution of, 1370-1371
 laboratory identification of, 245e-3
P. malariae
 characteristics of, 250e-1t, 1369, 1369t
 geographic distribution of, 1370-1371
 laboratory identification of, 250e-3f, 250e-4f
P. ovale
 characteristics of, 250e-1t, 1369t
 geographic distribution of, 1370
 laboratory identification of, 250e-3f, 250e-4f, 1377f
P. vivax
 adherence of, 145e-2t, 145e-3
 characteristics of, 250e-1t, 1369t
 geographic distribution of, 1370-1371
 host receptors for, 138e-2, 1369
 laboratory identification of, 250e-2f, 250e-3f, 1376f
 resistance to, 145e-3
 temperature sensitivity of, 1371
 transmission cycle, 145e-10, 1369f
 Plasticity, of cancer cells, 102e-12
 Plateau pressure, 1733
 Platelet(s), 725
 abnormal function of, 368
 activation of, 275f, 276, 400-401, 725, 741f, 742, 746f
 adhesion of, 275-276, 400-401, 725, 741-742, 745, 746f
 aggregation of, 276, 725, 742, 746f
 in AML, 681, 685
 in arterial thrombosis, 275-276, 741-742
 characteristics of, 275, 725, 741
 disorders of, 403t, 725
 acquired, 730
 hemolytic-uremic syndrome. *See* Hemolytic-uremic syndrome (HUS)
 inherited, 730
 thrombocytopenia. *See* Thrombocytopenia
 thrombocytosis. *See* Thrombocytosis
 von Willebrand disease. *See* von Willebrand disease
 elevated levels of, 677, 678
 giant, 81e-1, 81e-6f
 hematopoietic differentiation of, 89e-3f
 in hemostasis, 400-401, 725
 in inflammation, 276-277, 742
 in peripheral blood smear, 81e-1
 reference values for, 2756t
 in shock, 1747
 transfusion of. *See* Platelet transfusion
 Platelet-activating factor, 372e-15t, 1746
 Platelet aggregometry, 407
 Platelet count
 in DIC, 737
 normal, 404, 725
 in preoperative evaluation, 404
 Platelet-derived growth factor (PDGF), 102e-14, 102e-15f, 372e-14, 372e-14t
 Platelet-derived growth factor (PDGF) receptors, 102e-15f, 119e-1, 599
 Platelet emboli, retinal, 40e-4
 Platelet factor 4 (PF-4), 372e-13t, 727, 750
 Platelet function tests, 407
 Platelet inhibitors. *See* Antiplatelet agents
 Platelet plug, 400-401, 725
 Platelet-rich plasma, 138e-2
 Platelet transfusion
 for aplastic anemia, 667
 characteristics of, 138e-2t, 138e-3
 for coagulation disorders in liver disease, 739
 corrected count increment, 138e-3
 for DIC, 738
 refractoriness to, 138e-3
 response to, 138e-2t, 138e-3
 single-donor apheresis for, 138e-2
 for thrombocytopenia, 103e-24
 Platybasia, 426e-2
 Platypnea, 47e-3, 1992
 Platypus, 2740-2741
 PLC γ -associated antibody deficiency and immune dysregulation (PLAID), 2113
 Pleconaril, 1203, 1293
Pleistophora spp., 1408
 Pleomorphic adenoma, 241
 Pleomorphic xanthoastrocytomas, 601
 Plerixafor, 89e-1, 139e-2
 Pleural effusion, 1716
 in chylothorax, 1718
 in cirrhosis, 1717
 diagnosis of, 308e-14f, 1717f, 1725
 differential diagnosis of, 1718-1719, 1718t
 dyspnea in, 47e-2
 etiology of, 47e-4, 1716
 exudative, 1716
 in heart failure, 1725-1726
 in hemothorax, 1718
 malignant, 474, 510, 512, 1717
 in mesothelioma, 1717
 in pancreatitis, 2092, 2095f
 parapneumonic, 1717
 in pulmonary embolism, 1717
 transudative, 1716
 tuberculous, 1109-1110, 1718
 in viral infection, 1718
 Pleural (primary) effusion lymphoma, 214e-7, 1194
 Pleuritis
 abdominal pain in, 105
 chest pain in, 96t, 98, 99
 in rheumatoid arthritis, 2138
 in SLE, 2129, 2129t
 tuberculous, 1718
 Pleurodynia (Bornholm disease), 1291
 Plexopathies
 brachial. *See* Brachial plexopathy
 evaluation and treatment of, 2694
 lumbosacral, 2693-2694, 2694t
 perioperative, 2693
 radiation-induced, 2694
 PLP (pyridoxal 5'-phosphate), 426e-5
 PLTP (phospholipid transfer protein), 2438
 Plummer-Vinson syndrome, 237, 532
 Pluripotent stem cells, 454, 454f
 Plutonium exposure, 263e-3t, 263e-6t, 263e-7
 PML. *See* Progressive multifocal leukoencephalopathy (PML)
 PML gene, 679
 PML-RARA, 679, 687
 PML-RAR gene mutations, 102e-7, 431t
 PMNs (polymorphonuclear leukocytes), 150e-1, 1601
 PMP22 gene mutations, 434
 PMS1 gene mutations, 101e-6t, 596
 PMS2 gene mutations, 101e-6t, 592, 596
 PNad (peripheral node addressin), 372e-26
 PNAG, 145e-10
 pNETs. *See* Pancreatic neuroendocrine tumors (pNETs)
 pNETs (primitive neuroectodermal tumors), 119e-4. *See also* Ewing's sarcoma
 Pneumatic compression device, for VTE prophylaxis, 55
 Pneumatosis intestinalis, 1979, 2162
 Pneumococcal H inhibitor (hic), 947
 Pneumococcal infections, 946
 approach to the patient, 950
 in cancer patient, 484, 484t, 486t
 case-fatality ratios for, 949-950
 endocarditis, 817t, 823. *See also* Infective endocarditis
 epidemiology of, 947-949, 948f, 949f
 epiglottitis, 234-235
 etiology of, 946-947. *See also* *Streptococcus pneumoniae*
 global considerations in, 949
 in HIV infection, 1250
 host defense mechanisms, 950
 incidence pre- and postvaccine, U.S., 785t, 949f, 953, 953f
 mastoiditis, 231
 meningitis. *See* Meningitis
 otitis media. *See* Otitis media
 pathogenesis of, 949-950
 pneumonia. *See* Pneumonia, bacterial
 postsplenectomy, 413
 prevention of, 952-953. *See also* Pneumococcal vaccine
 risk factors for, 949, 949t
 seasonal variations in, 949
 sinusitis, 226, 952
 treatment of, 940t, 952-953
 in ventilator-associated pneumonia, 915
 Pneumococcal surface adhesion A, 947
 Pneumococcal surface antigen A, 947f
 Pneumococcal surface protein A, 947f
 Pneumococcal surface protein C, 947, 947f
 Pneumococcal vaccine
 capsular polysaccharide, 953-954
 genomics and, 778
 global considerations in, 947
 polysaccharide-protein conjugate, 949f, 953, 953f
 recommended administration of
 in adults, 29t, 787-788f, 809
 after splenectomy, 413, 635
 in cancer patients, 485t, 492
 in elderly, 84
 in HIV infection, 798, 1252-1253t
 in liver disease, 1995
 in transplant recipients, 929, 929t
 in travelers, 794
Pneumocystis jiroveci, 372e-26, 765t, 1358. *See also* *Pneumocystis pneumonia* (PCP)

- Pneumocystis pneumonia* (PCP)
 in cancer patient, 485, 489, 489t
 clinical features of, 1359, 1360f
 diagnosis of, 489, 1359–1360, 1360f
 ectopic ACTH production and, 611–612
 epidemiology of, 1358
 in HIV infection, 1250, 1251t, 1253, 1358, 1361
 pathogenesis of, 1358–1359, 1359f
 prevention of, 492
 prognosis of, 1360–1361
 prophylaxis for, 1361–1362, 1362t
 in HIV infection, 1251t
 pentamidine, 667
 trimethoprim-sulfamethoxazole, 139e-4t, 423
 sputum collection in, 1667
 in transplant recipient
 after HSCT, 921, 1358
 after solid organ transplant, 924t, 926, 927, 1358
 prophylaxis for, 139e-4, 139e-4t, 1361–1362, 1830
 treatment of, 1361, 1361t
 Pneumolysin, 947, 947f
 Pneumomediastinum, 1720
 Pneumonectomy, for lung cancer, 514, 516–517
 Pneumonia, **803**
 abdominal pain in, 105, 106t
 atypical, 804
 bacterial. *See also* Pneumonia, community-acquired
 actinomycotic, 1088
 after influenza, 958, 1212
 case-fatality ratios for, 949–950
 clinical features of, 950–951, 968
 complications of, 951, 968
 diagnosis of, 308e-10f, 951, 951f
 differential diagnosis of, 951
 E. coli, 1028
 F. tularensis, 1068
 in granulomatosis with polyangiitis, 386e-2f
 H. influenzae, 1011
 in HIV infection, 1250. *See also* *Pneumocystis pneumonia* (PCP)
 incidence pre- and postvaccine, U.S., 785t, 949f, 953, 953f
 K. pneumoniae, 1032
 Legionella. *See Legionella* spp. infections
 M. catarrhalis, 1013
 Nocardia, 308e-6f, 1084–1086, 1085f, 1086f
 nontyphoidal *Salmonella*, 1054
 P. aeruginosa, 1044–1045, 1045t, 1047
 parapneumonic effusion in, 1717
 T. whipplei, 1092
 treatment of, 952–953, 965t
 in burn patient, 166e-2
 C. pneumoniae, 1173–1174
 C. psittaci, 1173
 in cancer patient, 488–489, 489t, 1797
 chest pain in, 96t, 98
 Coccidioides, 1335
 community-acquired
 clinical features of, 805
 complications of, 808–809
 diagnosis of, 805–806
 epidemiology of, 805, 805t
 etiology of, 804–805, 804t, 806, 1015
 follow-up, 809
 PPIs and, 1921
 prevention of, 809
 prognosis of, 809
 risk factors for, 805, 805t
 severity indicators, 806–807, 807t
 treatment of
 antibiotic resistance in, 807
 empirical antibiotics, 767t, 807–808, 808t
 failure to respond to, 808
 general considerations, 808–809
 racial/ethnic disparities in, 16e-1, 16e-2f
 site of care, 806–807
 cryptogenic organizing, 1712–1713
 cyanosis in, 249
 deaths from, 6e-1t, 56t, 804
 definition of, 803
 eosinophilic. *See* Pulmonary infiltrates with eosinophilia
 health care–associated. *See also* Ventilator-associated pneumonia (VAP)
 conditions associated with, 803, 803t
 diagnosis of, 958
 prevention of, 914–915
 risk factors for, 809, 914
 treatment of, 767t, 914
 hemoptysis in, 245
 hospital-acquired, 767t, 813, 1015, 1020
 incidence of, 804
 infant (*C. trachomatis*), 1170t, 1171
 interstitial. *See* Interstitial pneumonia
 M. pneumoniae, 1163
 mixed viral and bacterial, 1212
 necrotizing, 812
 pathology of, 804
 pathophysiology of, 306e-3, 804
 Pneumocystis. *See Pneumocystis pneumonia* (PCP)
 Q fever, 1161
 R. equi, 981
 in transplant recipient, 923, 925–926
 typical, 804
 ventilator-associated. *See* Ventilator-associated pneumonia (VAP)
 viral
 adenovirus, 1208
 CMV, 1192
 enterovirus, 1293
 HMPV, 1207
 HRSV, 1206
 parainfluenza, 1207
 primary influenza, 1212
 varicella, 1184
 Pneumonia Severity Index (PSI), 806
 Pneumonic plague, 261e-3, 261e-5t, 1072, 1073f.
 See also Plague
 Pneumonitis
 aspiration, anaerobic, 1098
 in cancer patient, 489, 489t, 621–622, 1797
 deaths from, by sex, 6e-1t
 giant-cell, 1297, 1298
 HSV, 1179–1180
 hypersensitivity. *See* Hypersensitivity pneumonitis
 lymphocytic interstitial, 1716
 necrotizing, anaerobic, 1098
 radiation-induced, 1797
 VZV, 1183
 Pneumoperitoneum, 847, 847f
 Pneumothorax
 chest pain in, 96t, 98
 as contraindication to hyperbaric oxygen therapy, 477e-2
 imaging of, 308e-13–14f
 in mechanically ventilated patient, 1743
 spontaneous, 98, 1719
 tension, 98, 1719, 1750
 traumatic, 1719
 PNKD (paroxysmal nonkinesigenic dyskinesia), 2625
 Podagra, with gouty inflammation, 436e-2f
 Podophyllotoxin
 adverse effects of, 2687t
 for warts, 351, 1201, 1201t
 POEMS syndrome (Crow-Fukase syndrome), **718**
 clinical features of, 361, 715t, 718, 2348, 2700
 diagnosis of, 715t
 paraneoplastic, 619
 pathogenesis of, 718, 2348
 treatment of, 718, 2348
 Poikilocytes, 81e-1, 627
 Poikilocytosis, 81e-1, 394, 413, 637, 657
 Poikiloderma, 340t, 356t, 357, 374–375
 Poikilothermia, 478e-3
 Point mutation, 101e-2–3, 101e-4, 432–433, 433f
 Poisoning/drug overdose. *See also specific substances*
 epidemiology of, 473e-1
 history in, 473e-1
 hypothermia in, 478e-1
 laboratory evaluation in, 473e-3–4, 2764–2766t
 pattern recognition in, 473e-1, 473e-2t
 phases of, 473e-4
 physical examination in, 473e-1
 severity assessment in, 473e-1–3, 473e-2t
 treatment of
 antidote administration, 473e-7
 cardiovascular therapy, 473e-5
 CNS therapy, 473e-5–6
 decontamination at other sites, 473e-6–7
 enhancement of poison elimination, 473e-7
 gastrointestinal decontamination, 473e-6
 general principles, 473e-4–5, 473e-4t
 prevention of reexposure, 473e-7
 respiratory care, 473e-5
 supportive care, 473e-5
 Poison ivy, 345
 Poison oak, 345
 Poison sumac, 345
 Poisson distribution, 87e-1, 87e-2f
 Polar bear bite-wound, 167e-1
 Polarized cells, 332e-3
pol gene, 225e-1, 225e-2, 225e-3f, 1218, 1219f
POLG gene, 462e-16
 Poliomyelitis. *See* Poliovirus infection
 Poliovirus, 214e-6, 1289–1290
 Poliovirus infection
 clinical features of, 1290–1291
 elimination and eradication of, 785
 epidemiology of, 1290
 global considerations in, 786, 896, 1294, 1294t
 incidence pre- and postvaccine, U.S., 785t
 paralytic poliomyelitis, 1290
 pathogenesis of, 1290
 postpolio syndrome, 1291, 2633
 prevention and eradication of, 1294, 1294t
 vaccine-associated, 1291
 weakness in, 156
 Poliovirus vaccine
 effectiveness of, 1294
 poliomyelitis associated with, 1291, 1294
 recommended administration of
 in adults, 1295t
 in cancer patient, 485t
 in transplant recipient, 929, 929t
 in travelers, 794
 types of, 1294
 Polonium-210 exposure, 263e-3t
 POLST (Physician Orders for Life-Sustaining Treatment), 58–59
 Polyangiitis
 with granulomatosis. *See* Granulomatosis with polyangiitis (Wegener's)
 microscopic. *See* Microscopic polyangiitis
 Polyangiitis overlap syndromes, 2193
 Polyarteritis nodosa, **2187**
 clinical features of, 2187, 2187t
 definition of, 2187
 diagnosis of, 386e-3f, 386e-4f, 386e-6f, 2188–2189
 incidence and prevalence of, 2187
 neuropathy in, 2700
 pathology and pathogenesis of, 2180, 2187
 skin manifestations of, 364t, 366, 368–369
 treatment of, 2188
 Polyarthralgia syndrome, 1196, 1196t
 Polychondritis, relapsing. *See* Relapsing polychondritis
 Polychromasia, 394, 729
 Polychromatophilia, 81e-1, 81e-3f
 Polychromotropism, 575
 Polycarb family, 89e-4
 Polycyclic hydrocarbons, 119e-1, 477t
 Polycystic kidney disease
 autosomal dominant, **1850**
 clinical features of, 1621t, 1851–1852, 1852f
 diagnosis of, 1622, 1852–1853
 early interventions for, 450t
 genetic considerations in, 436t, 1621t, 1850–1851, 1850t
 genetic testing for, 450t

- Polycystic kidney disease, autosomal dominant
(*Cont.*):
pathogenesis of, 1850–1851, 1851f
treatment of, 1853
autosomal recessive, 1850t, **1853**
Polycystic ovarian syndrome (PCOS)
in CAH, 2356
hirsutism in, 331, 332
infertility in, 2389
insulin resistance in, 2406
menstrual disorders in, 337
metabolic syndrome and, 337, 2452
obesity and, 415e-6
prevalence of, 2254t
screening/testing for, 2254t
skin manifestations of, 352, 356t, 359
sleep apnea in, 6e-5
treatment of, 337
- Polycystin-1/2, 1850, 1851f
- Polycythemia, **399**
as adaptation to hypoxia, 248
approach to, 400f
clinical features of, 399–400
erythromelalgia in, 1650
secondary, 249
smoker's, 400
spurious, 400
- Polycythemia vera, **672**
classification of, 672, 672t
clinical features of, 399, 673
complications of, 673–674
vs. cyanosis, 248
diagnosis of, 400, 673, 673t
epidemiology of, 672
etiology of, 672–673
genetic considerations in, 672–673
splenomegaly in, 411, 673–674
treatment of, 674
- Polydipsia
dipsogenic, 2276, 2279
etiology of, 295
hyponatremia in, 301
iatrogenic, 2276, 2279
primary, 2276, 2277, 2279
psychogenic, 2276, 2279
- Polydrug abuse, 469e-5. *See also* Substance use disorders
- Polyethylene glycol, 274, 1970t
- Polygenic disorders, 425, 438t, 440
- Polyglandular autoimmune (PGA) insufficiency, 2366, 2483
- Polyglycolic acid, 92e-1
- Polymerase chain reaction (PCR)
definition of, 769t
for genetic testing, 444, 447
in infectious disease diagnosis, 769t
in pneumonia diagnosis, 806
in viral encephalitis diagnosis, 894
in viral meningitis diagnosis, 891
- Polymer fume fever, 1692
- Polymorphic microsatellite markers, 441
- Polymorphic reticulosis. *See* Extranodal NK/T-cell lymphoma, nasal type
- Polymorphism, 433. *See also* Single-nucleotide polymorphisms (SNPs)
- Polymorphonuclear leukocytes (PMNs), 150e-1, 1601
- Polymorphous light eruption, 364t, 366, 388–389
- Polymyalgia rheumatica, **2188**
as cause of fever of unknown origin, 138
clinical features of, 202, 462e-2, 2188, 2199
definition of, 2188
diagnosis of, 2189
treatment of, 141, 2189
- Polymyositis, **2194**
clinical features of, 157, 2194–2196, 2195t
diagnosis of, 2199–2200, 2199t, 2200f
differential diagnosis of, 2197–2199
dysphagia in, 257
immunopathology of, 372e-24t, 2196, 2197f
interstitial lung disease in, 1714
parasitic, 2198
prognosis of, 2202f
Raynaud's phenomenon in, 1649
treatment of, 2200–2202
tropical, 2198
viral infections and, 2197
- Polymyxin B
for *Acinetobacter* infections, 1037t
action of, 931t, 933, 934, 944
adverse effects of, 381, 938t, 944, 1037
for gram-negative bacterial infection, 1026, 1027
indications for, 944
for otitis externa, 228
resistance to, 931t, 934, 941t
- Polymyxin-bacitracin, 199
- Polymyxin E (colistin)
for *Acinetobacter* infections, 1037t
action of, 931t, 933, 934, 944
adverse effects of, 938t, 944, 1027
for gram-negative bacterial infection, 1026, 1027
indications for, 941t, 944
for *Klebsiella* infections, 1033
for *P. aeruginosa* infection, 1045t, 1047
resistance to, 931t, 934
- Polyneuropathy
acute inflammatory demyelinating, 2695, 2695t, 2696t
acute toxic, 157
autoantibodies in, 372e-24t
chronic inflammatory demyelinating, 2696t, **2698**
critical illness, 1755, 1783–1784, 2684
cryptogenic sensory and sensorimotor, 2691
diabetic. *See* Diabetic neuropathy
familial amyloid, 2681
in Guillain-Barré syndrome. *See* Guillain-Barré syndrome (GBS)
hereditary sensory and autonomic, 2680
in HIV infection, neurologic disease in
large-fiber, 161
in monoclonal gammopathy of undetermined significance, 2700
in multiple myeloma, 2699–2700
sensory deficits in, 161
small-fiber, 161
weakness in, 157
- Polyomavirus(es). *See also* BK virus infection; JC virus infection
gene expression and replication in, 214e-4
Merkel cell, 926
structure of, 214e-2t, 214e-3f
in transplant recipient, 926
- Polyp(s)
colorectal
bleeding from, 277
endoscopic findings in, 1881, 1882f, 1883f
endoscopic treatment of, 346e-1v
in familial adenomatous polyposis, 1897f. *See also* Familial adenomatous polyposis
flat serrated, 1881, 1883f
pathogenesis of cancer in, 537–538. *See also* Colorectal cancer
gallbladder, 2083
- Polypharmacy, in elderly, 77–78
- Polypill, 266e-6
- Polyploidy, of mitochondrial DNA, 85e-1–2
- Polyposis, juvenile, 101e-6t, 538t
- Polyposis coli. *See* Familial adenomatous polyposis
- Polyradiculopathy
diabetic. *See* Diabetic neuropathy
in HIV infection. *See* HIV infection, neurologic disease in
- Polyreactive natural antibodies, 372e-2
- Polysaccharide A, in abscess formation, 849
- Polysaccharide antigens, selective antibody deficiency to, 2112
- Polysaccharide capsule, pneumococcal, 947, 947f
- Polysaccharide iron, for iron-deficiency anemia, 629t
- Polysomnography, 184, 189f
- Polyunsaturated long-chain triglycerides, 98e-4–5
- Polyuria
approach to the patient, 294, 294f
hypokalemic, 307
pathophysiology of, 294–295
in sepsis/septic shock, 1755
treatment of, 304
in urinary tract obstruction, 1872
in water diuresis, 64e-3
- Pomalidomide, 103e-19, 717
- POMC. *See* Proopiomelanocortin (POMC)
- POMC gene, 401e-1, 401e-4, 611
- POMC gene mutations, 415e-3, 2325t
- Pompe disease (acid α -1,4 glucosidase deficiency)
clinical features of, 432e-4t, 432e-6, 433e-2t, 433e-4–5, 462e-14
diagnosis of, 433e-5, 462e-14
enzyme deficiency in, 432e-4t
genetic considerations in, 462e-14
late-onset, 432e-4t
prevalence of, 432e-6
treatment of, 432e-6, 462e-14
- Ponatinib
action and targets of, 102e-3, 102e-5t, 103e-16, 103e-17t
adverse effects of, 103e-16, 103e-17t, 621, 692
for CML, 691–692, 691t
- Poncet's disease, 836
- Pongola virus, 1308t
- Pons, 441e-6f, 2577–2578f
- Pontiac fever, 1015, 1016
- Pontine hemorrhage, 41e-1v
- Poor metabolizers, 38f, 40
- Popliteal artery
aneurysm, 1647
entrapment, 1647
pulse, 1446f
- Popliteal (Baker's) cyst, 2222
- Population genetics, 442
- Population migration, climate change and, 151e-3–4
- Porcelain gallbladder, 2082
- POR gene mutations, 2354t, 2356t
- Porins, 930, 1004
- Pork tapeworm infection. *See* *Taenia solium* (pork tapeworm) infection
- Porphyria(s), **2521**
acute, 367e-3, 2523t, 2525
classification of, 2521, 2522t, 2525
clinical features of, 2521, 2522t, 2525
abdominal pain, 105, 2681
autonomic dysfunction, 2642
cutaneous, 362
liver disease, 367e-3
neurologic, 2681
oral, 236
photosensitivity, 386, 390
cutaneous (chronic), 367e-3, 2523t, 2525–2526
diagnosis of, 2521, 2522t, 2523t
differential diagnosis of, 2525–2526
erythropoietic
clinical features of, 2521, 2522t, 2525
congenital, 2521, 2532
diagnosis of, 2522t
pathophysiology of, 2522f
protoporphyrin. *See* Erythropoietic protoporphyrin (EPP)
X-linked sideroblastic anemia, 2531–2532
global considerations in, 2521
hepatic
acute intermittent porphyria. *See* Acute intermittent porphyria (AIP)
ALA-dehydratase deficient, 2521, 2526
clinical features of, 2521, 2522t, 2526
diagnosis of, 2522t
hereditary coproporphyrin. *See* Hereditary coproporphyrin (HCP)
pathophysiology of, 2522f
porphyria cutanea tarda. *See* Porphyria cutanea tarda (PCT)
unsafe drugs in, 2526, 2527–2529t
variegate porphyria. *See* Variegate porphyria (VP)

- Porphyria cutanea tarda (PCT), 2530
 clinical features of, 436e-5f, 2523t, 2530, 2530f
 diagnosis of, 2523t, 2530
 differential diagnosis of, 362
 genetic considerations in, 367e-7, 390, 2530
 HCV infection and, 367e-3, 2041
 hepatocellular carcinoma and, 545, 545t, 2530
 pathogenesis of, 2522f
 photosensitivity in, 390
 risk factors for, 2530
 skin manifestations of
 blistering lesions, 2530, 2530f
 hyperpigmentation, 360, 360t, 390, 436e-5f
 vesicles/bullae, 361t, 362, 367e-3, 390, 436e-5f
 treatment of, 367e-3, 390, 2530–2531
- Porphyrinogens, 2524
- Porphyrins, 103e-5, 390
- Porphyromonas* spp., 1094
- Porphyromonas* spp. infections. *See also*
 Anaerobic bacterial infections, mixed
 of bone, 1099
 in dog-bite wound, 167e-1
 in human bite-wound, 167e-2
- Portal hypertension
 ascites in, 287, 2063, 2064. *See also* Ascites
 in cirrhosis, 2058
 classification of, 2063t
 clinical features of, 2063
 definition of, 2062–2063
 diagnosis of, 2063
 esophageal varices in. *See* Esophageal varices
 etiology of, 2063
 gastrointestinal bleeding in, 277, 2063. *See also*
 Esophageal varices
 in hepatocellular carcinoma, 545
 pulmonary hypertension and, 1658
 splenomegaly and hypersplenism in, 2064
- Portuguese man-of-war envenomation, 2738
- Posaconazole
 for *Aspergillus* infections, 1349t
 for *Blastomyces* infections, 1339
 for *Candida* prophylaxis, 1253t, 1345
 for *Coccidioides* infections, 1336
 for fungal infections in cancer patient, 491
 for *Fusarium* infection, 1356
 indications for, 1331
 for mucormycosis, 1353
- Positional cloning, 441
- Positive economics, 15e-3
- Positive end-expiratory pressure (PEEP), 1738.
See also Mechanical ventilation
- Positive predictive value (PPV), 479, 479t
- Positive-strand genome, 214e-1
- Positron emission tomography (PET)
 in adrenocortical carcinoma, 2322, 2323f
 in cardiac disease
 infective endocarditis, 270e-24, 270e-25f
 ischemic heart disease, 1582, 1584f
 for myocardial perfusion imaging, 270e-3.
See also Myocardial perfusion imaging
 in fever of unknown origin, 140–141
 in lung cancer, 513, 1666
 in medullary thyroid carcinoma, 2341f
 in MEN 2A, 2341f
 in musculoskeletal disease evaluation, 2225
 in neurologic disease, 440e-8–9
 cluster headache, 2590f
 dementia, 174, 175f
 migraine, 2589, 2589f, 2590f
 PD, 2609, 2611f
 stroke, 2581
 in osteomyelitis, 840
 in pancreatic cancer, 555
 principles of, 440e-8–9
 in respiratory disease, 1666
 in sarcoidosis, 2209, 2209f
 in solitary pulmonary nodule, 1666
 in tumor detection, 102e-12
- Postal workers, anthrax exposure in, 261e-2
- Postganglionic septicaemia. *See* Lemierre's disease/
 syndrome
- Postantibiotic effect, 935
- Postcardiac injury syndrome, 1575
- Postcoital contraception, 2391
- Postconcussion syndrome, 457e-5, 471e-3,
 471e-5–6
- Posterior cerebral artery
 fetal, 2574
 lesion in territory of, 179, 183
 occlusion of, 199, 2574–2575, 2575f, 2580f
 territory of, 2572f, 2575f
- Posterior column-medial lemniscal pathway,
 158, 159f
- Posterior cortical atrophy syndrome, 182
- Posterior fossa brain tumor, 108
- Posterior pituitary
 disorders of
 diabetes insipidus. *See* Diabetes insipidus
 (DI)
 hypodipsic hypernatremia. *See* Hypodipsic
 hypernatremia
 hormone production in, 2274. *See also*
 Arginine vasopressin (AVP); Oxytocin
- Posterior reversible encephalopathy syndrome
 (PRES), 1793, 2568, 2584
- Postexposure prophylaxis (PEP)
 HBV, 2021–2022
 HIV, 918, 1222, 1283
 measles, 1298
 rabies, 1303–1304, 1304f
 VZV, 917, 1186, 1186t
- Postextrasystolic beat, 254
- Postextrasystolic potentiation, 254
- Postherpetic neuralgia, 215e-8, 237, 1185, 1186,
 2685
- Postictal state, 1773
- Postmenopausal hormone therapy, 2382
 adverse effects of, 744, 2498, 2498f
 approach to the patient, 2386, 2387f
 benefits of, 2383, 2383t
 breast cancer risk and, 524, 2383, 2498f
 cognitive function and, 6e-2, 2386–2387
 colorectal cancer risk and, 478, 539, 2385
 coronary heart disease and, 6e-3, 291e-7–8,
 2385, 2498f
 endometrial cancer and, 596, 2383, 2386
 fracture reduction due to, 2498, 2498f
 gallbladder disease and, 2385
 health status changes after discontinuation
 of, 2386
 lipoprotein disorders and, 2443–2444
 ovarian cancer and, 2386
 risk-benefit profile, 2384t, 2386
 short-term vs. long-term use of, 2386
 stroke risk and, 2498f
 VTE and, 2383, 2498f
- Postnasal drip, 244
- Postoperative period
 acute kidney injury in, 1802–1803
 nausea and vomiting in, 259
- Postparalytic syndrome, 1733
- Postpartum period
 depression in, 6e-5
 DVT in, 47
 hemorrhage in, 403–404
 infections in, 50, 1165
 thyroiditis in, 2299
- Postpolio syndrome, 1291
- Post-premature ventricular contraction, 51e-8
- Poststreptococcal glomerulonephritis
 clinical features of, 233, 1835t, 1836t, 1837
 epidemiology of, 1837
 etiology of, 1837
 pathophysiology of, 1837, 1837f
 renal biopsy in, 62e-2f, 1837
 treatment of, 1837
- Posttest probability of disease, 22, 23f
- Postthrombotic syndrome, 1631, 1631f, 1634
- Posttranslational processing, 427, 428f
- Posttraumatic stress disorder (PTSD)
 clinical features of, 2712
 definition of, 471e-2
 diagnosis of, 471e-2–3, 2713t
 pathophysiology of, 471e-2, 2712
 treatment of, 471e-4–5, 2712–2713
 in war veterans, 471e-2–3
- Postural control, 162
- Postural hypotension. *See* Orthostatic
 hypotension
- Postural orthostatic tachycardia syndrome
 (POTS), 1477, 2642–2643
- Postwar symptoms. *See* War veterans
- Potassium, 304
 in cardiac contraction, 265e-6
 in chronic kidney disease, 333e-2–3, 1814
 depletion of, 305–306, 322
 excess intake of, 35, 310
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 renal transport of, 332e-5f, 332e-9
 in specialized nutritional support formulas,
 98e-5
 transport and reabsorption of, 296–297, 297f,
 304–305
- Potassium balance, 304–305
- Potassium bicarbonate, for hypokalemia, 307
- Potassium-binding resin, for hyperkalemia,
 64e-3, 1815
- Potassium channels
 ATP-sensitive, 2402, 2402f
 cardiac, 273e-1f, 273e-2, 273e-2f
 muscle, disorders of, 462e-17t, 462e-18
 in seizure initiation, 2547
- Potassium chloride (KCl)
 administration guidelines, 308
 adverse effects of, 308, 310, 644t
 for diabetic ketoacidosis, 64e-1
 for hypokalemia, 64e-7, 307–308
 for hypokalemic periodic paralysis, 462e-17
- Potassium citrate, 307, 431e-5
- Potassium-competitive acid pump antagonists,
 1921
- Potassium hydroxide (KOH) wet-mount
 preparation, 150e-1, 341
- Potassium iodide (KI)
 adverse effects of, 1354
 for radioactive iodine exposure, 263e-6t,
 263e-7
 for *Sporothrix* infections, 1354, 1354t
 for thyroid storm, 2297
- Potassium perchlorate, 2300
- Potassium phosphate, 64e-2
- Potassium-sparing diuretics, 332e-9, 1815
- Potato riddler's lung, 1681t
- Potency, of stem cells, 453
- POTS (postural orthostatic tachycardia
 syndrome), 1477, 2642–2643
- Pott's disease, 117, 836, 852, 1110, 1111f
- Pott's puffy tumor, 227
- Pou5f1* (*Oct4*) gene, 454, 455
- Pouchitis, 1960, 1962–1963
- Poultry worker's lung, 1681, 1681t
- Poverty, health and, 9. *See also* Low- and
 middle-income countries
- Powassan virus, 1307t, 1315
- Powassan virus encephalitis, 1152
- Power law, 86e-6–7
- Power law distribution, 87e-1–2, 87e-2f
- Power of attorney for health care, 59
- Poxviruses. *See also specific viruses*
 gene expression and replication in, 214e-4–5
 host defense inhibition by, 214e-8
 human infections caused by, 220e-1–2,
 220e-1t
 protein functions in, 214e-6
 structure of, 214e-2t, 214e-3f
- PPAR-alpha agonists, 291e-6
- PPARγ (peroxisome proliferator-activated
 receptor γ), 96e-6, 415e-2, 1959
- PPD. *See* Purified protein derivative (PPD)/
 tuberculin skin test (TST)
- PPHP (pseudopseudohypoparathyroidism), 430,
 2485, 2486t
- PPIs. *See* Proton pump inhibitors (PPIs)
- PPKARIA gene mutations, 2344
- PPNAD (primary pigmented nodular adrenal
 disease), 2314, 2343
- PPomas (pancreatic polypeptide-secreting
 tumors), 559t, 2338
- PPOX gene, 2523t
- PPV (positive predictive value), 479, 479t

- Practice guidelines, 3, 28
 PRAD 1, 2471f
 Prader-Willi syndrome
 clinical features of, 415e-4, 415e-5t, 439, 2256, 2364
 genetic considerations in
 imprinting, 83e-7, 415e-5t, 439, 444e-2, 2256, 2364
 uniparental disomy, 439
 X-inactivation, 439
 Pragmatics, 179
 Pralatrexate, 103e-11t
 2-Pralidoxime chloride (2-PAM Cl), 262e-5t, 262e-6, 262e-6f, 473e-10t
 Pramipexole
 adverse effects of, 2615
 for PD, 2615, 2616t
 for restless legs syndrome, 188t, 192, 2625
 Pramlintide, 2412–2413, 2414t
 Prasugrel
 action of, 747
 adverse effects of, 404, 747
 for cardiac catheterization pretreatment, 1461
 vs. clopidogrel, 747
 contraindications to, 1596
 dosage of, 747
 genetic variation in response to, 277t, 743t
 for ischemic heart disease, 1590
 management before endoscopic procedures, 1889t
 for myocardial infarction, 1596, 1598, 1598t, 1606
 in PCI, 296e-1, 296e-2, 1596
 resistance to, 747
 Prausnitz-Küstner reaction, 2116
 Pravastatin, 2199, 2218t, 2448t
 Praziquantel
 action of, 246e-12
 adverse effects of, 246e-6t, 246e-12
 for biliary flukes, 1428t
 for cysticercosis, 1432
 for diphyllorhynchiasis, 1434
 for echinococcosis, 1434
 for *H. nana* infections, 1434
 indications for, 246e-6t
 for intestinal flukes, 1428t
 for lung flukes, 1428t
 for neurocysticercosis, 903
 pharmacology of, 246e-12
 in pregnancy and lactation, 246e-6t
 for schistosomiasis, 246e-12, 1428, 1428t
 for *T. asiatica* infection, 1430
 for *T. saginata* infection, 1430
 for *T. solium* infection, 1432
 Prazosin
 adverse effects of, 2640t
 for ergot alkaloid overdose/poisoning, 473e-8t
 for hypertension, 1624t
 for PTSD, 471e-5, 2712
 for Raynaud's phenomenon, 1649, 2164
 Prealbumin. *See* Transthyretin (TTR)
 Pre-Bötzinger complex, 1720
 Precipitation, patterns of, 151e-3–4
 Precocious puberty
 female
 definition of, 2380
 diagnosis of, 2380t
 differential diagnosis of, 2380t
 etiology of, 2380
 ovarian germ cell tumors and, 594
 male, 2361
 approach to the patient, 2362
 breast enlargement in, 2362
 in congenital adrenal hyperplasia, 2362
 definition of, 2361
 etiology of, 2361t
 familial male-limited, 2362
 genetic considerations in, 437, 2252
 gonadotropin-dependent, 2361
 gonadotropin-independent, 2361
 in McCune-Albright syndrome, 2362. *See also* McCune-Albright syndrome
 treatment of, 2362
 Preconceptual diagnosis, 444
 Preconditioning, kidney, 333e-3–4, 333e-4t
 Precordial thump, 1768
 Precursor B-cell lymphoblastic leukemia/
 lymphoma, 91e-4, 701
 Precursor cells, 454
 Precursor T-cell lymphoblastic leukemia/
 lymphoma, 702t, 707
 Prediction models, 23–24, 24t
 Predictive value of test, 22
 Predisposition testing, 448
 Prednisolone
 for alcoholic liver disease, 2053, 2054f
 for asthma, 1677
 for chorea, 2153
 for COPD exacerbation, 1707
 for optic neuritis, 202
 for rheumatic fever, 2153
 Prednisone
 for ACTH deficiency, 2260
 for acute idiopathic pericarditis, 1573
 for adult hypopituitarism, 2259t
 adverse effects of, 462e-19, 2129, 2145, 2200–2201. *See also* Glucocorticoid(s), adverse effects of
 for alcoholic liver disease, 2053, 2054f
 for allergic rhinitis, 2124
 for asthma, 1677
 for autoimmune hemolytic anemia, 658
 for autoimmune hepatitis, 2051–2052
 for autoimmune pancreatitis, 2099
 for Behçet's syndrome, 2194
 for Bell's palsy, 2648
 for bullous pemphigoid, 372
 for CAH, 334
 for cancer treatment-induced pneumonitis, 622
 for chorea, 2153
 for chronic inflammatory demyelinating polyneuropathy, 2699
 for cluster headache prevention, 2595, 2595t
 for contact dermatitis, 345
 for drug-induced hypersensitivity syndrome, 382
 for Duchenne's muscular dystrophy, 462e-8
 for eosinophilic fasciitis, 376
 for gastric lymphoma, 536
 for giant cell arteritis, 2189
 for gout, 2234
 for Graves' ophthalmopathy, 207
 for hypercalcemia, 314
 for hypercalcemia of malignancy, 610
 for hyper eosinophilic syndrome, 135e-8
 for hypersensitivity pneumonitis, 1683
 for IBD, 1960
 for idiopathic cutaneous vasculitis, 2192
 for idiopathic pulmonary fibrosis, 1712
 for IgA vasculitis, 2190
 for IgG4-related disease, 391e-3
 for immune thrombocytopenic purpura, 728
 for immunosuppression, 1828
 for inflammatory myopathies, 2200
 for interstitial lung diseases, 1711
 for lepra reactions, 1127
 for loiasis, 1422
 for minimal change disease, 1842
 for MS, 2667
 for mucous membrane pemphigoid, 374
 for multiple myeloma, 716–717
 for myasthenia gravis, 2705
 for neurocysticercosis, 903
 for PCP, 1361t
 for pemphigoid gestationis, 373
 for pemphigus vulgaris, 371
 for polymyalgia rheumatica, 2189
 for postherpetic neuralgia, 1186
 for Q fever, 1162
 for relapsing polychondritis, 2204
 for rheumatic fever, 2153
 for rheumatoid arthritis, 2145
 for sarcoidosis, 2211, 2212t
 for serum sickness, 2737
 for sinusitis with hyposmia, 216
 for SJS and TEN, 383
 for SLE, 2132
 for subacute thyroiditis, 2299
 for thymoma, 123e-2
 for tonsillar hypertrophy, 1189
 for trichinellosis, 1411t
 for vasculitic neuropathy, 2700
 for VIPoma, 571
 Preeclampsia, 45
 clinical features of, 46
 definition of, 45
 diagnosis of, 46
 in gestational trophoblastic disease, 597
 KIRs with, 372e-15t
 pathophysiology of, 46
 risk factors for, 45
 treatment of, 46, 1626t
 Prefrontal network, 183
 Pregabalin
 adverse effects of, 2682t
 for anxiety disorders, 2711
 for fibromyalgia, 2240, 2240t
 for hot flashes, 623
 for neuropathy, 2427, 2682t
 for pain, 61, 91t, 94, 2101, 2672
 for status epilepticus, 2558f
 Pregnancy, 45
 abdominal swelling in, 285
 acute fatty liver of, 49
 age of first full-term, breast cancer risk and, 524
 alcohol use in, 2725
 antibacterial therapy in, 936, 937t
 antiparasitic agents in, 246e-1–7t
 antiphospholipid syndrome in, 2135t, 2136
 aortic regurgitation in, 46–47
 aortic stenosis in, 46–47
 aplastic anemia in, 664
 asthma in, 1680
 asymptomatic bacteriuria in, 49–50, 862
 AVP assays in, 304
 bacterial infections in, 49–50
 breast cancer in, 124e-1t, 124e-3–4, 124e-3t
 breast mass in, 124e-3, 525–526
 C. trachomatis infection in, 1169, 1171
 cancer chemotherapy during, 103e-26, 124e-1, 124e-2
 cardiomyopathy in, 47, 1562
 cervical cancer during, 124e-1t, 124e-2–3
 cholestasis in, 49, 284, 2029, 2078
 chronic kidney disease in, 46, 1819
 CML treatment during, 694
 CMV infection in, 50, 1191
 cobalamin deficiency in, 642
 congenital heart disease and, 47, 1519, 1522, 1528
 depression in, 6e-5
 diabetes insipidus in, 303
 diabetes mellitus in, 6e-3–4, 47–48, 2400, 2422
 DIC in, 737
 DVT in, 47
 ectopic, 338
 edema in, 253
 epilepsy and, 49, 2559
 erythema infectiosum in, 127
 fatigue in, 152
 filovirus infections in, 1328
 folate deficiency in, 642–643, 646
 folate requirement in, 47, 265e-12, 643, 646, 648
 gallbladder disease in, 49, 2078
 gastrointestinal disease in, 49
 GBS infection in, 970
 genital herpes in, 50
 gonococcal infections in, 1006, 1008
 Graves' disease in, 2297
 HBV infection in, 49
 headache in, 49
 heart disease in, 46–47
 heart murmurs in, 51e-6
 HELLP syndrome in, 46, 49, 1865, 2019
 hematologic disease in, 49
 hemoglobin/hematocrit in, 394t
 hemoglobinopathy in, 49

- in hemophilia carriers, 735
 herpesvirus in, 50
 high-altitude travel during, 476e-5
 HIV infection in, 50, 1223
 Hodgkin's lymphoma in, 124e-4
 hormonal integration of, 400e-5
 HSV infection in, 1180
 hypertension in, 45-46, 1620t, 1621
 hyperthyroidism in, 48
 hypothyroidism in, 48, 49, 2293, 2300
 IBD in, 49, 1964
 immunization during, 50-51, 788f, 790
 influenza in, 1212
 iron requirement in, 626
L. monocytogenes infections in, 983
 lead poisoning in, 472e-1
 liver disease in, 49
 lymphocytic choriomeningitis in, 1318
 malaria in, 1374
 malaria prevention in, 1382
 Marfan syndrome in, 47
 maternal mortality, 51
 melanoma during, 124e-4
 mercury consumption in, 472e-1
 migraine in, 49
 mitral regurgitation in, 46-47
 mitral stenosis in, 46, 1542
 molar. *See* Gestational trophoblastic disease
 in MS, 49, 2666-2667
 nausea and vomiting in, 49, 259, 261
 neurologic disease in, 49, 463e-5
 non-Hodgkin's lymphoma in, 124e-4
 obesity and, 48
 oral disease in, 237
 parvovirus B19 infection in, 50, 1196
 pemphigoid gestationis in, 370t, 372-373
 peripheral nerve disorders in, 49
 pheochromocytoma in, 2331
 prenatal diagnosis in, 83e-3-4, 83e-4t
 prolactinoma in, 2268-2269
 psoriasis in, 347
 pulmonary embolism in, 47
 pulmonary hypertension in, 47
 Q fever in, 1161
 radiation therapy during, 124e-1-2, 124e-1t
 respiratory alkalosis in, 324
 retinoic acid in, 96e-7
 rheumatoid arthritis in, 2148
 rubella in, 50, 127, 230e-1. *See also* Rubella
 (German measles), congenital
 sexually transmitted infections in, 50
 SLE in, 2133
 smoking effects in, 2730, 2730t
 stroke in, 49
 syphilis in, 1136, 1139, 1139t
 in systemic sclerosis, 2162
T. gondii infection in, 1399-1400, 1404
 taste function in, 215
 thrombocytopenia in, 49
 thyroid cancer in, 124e-4
 thyroid function during, 48, 124e-4, 124e-5t,
 2286, 2300
 travel in, 797-798
 tuberculosis treatment in, 1116t, 1120
 tumor markers in, 473t
 urinary tract infections in, 867
 varicella in, 1184
 viral infections in, 50
 vitamin A in, 96e-7
 VTE risk in, 744
 warfarin in, 756
 water requirement in, 95e-1
 Pregnenolone, 2309f, 2311
 Prehypertension, 1616, 1616t, 1622, 2454
 Prejudice, 16e-4
 Prekallikrein deficiency, 733t
 Preload, 265e-7, 265e-9t, 265e-10f
 Premalignant changes, 102e-1
 Premature atrial contractions, palpitations in,
 254
 Premature closure, in clinical reasoning, 19
 Premature ejaculation, 325
 Premature menopause. *See* Primary ovarian
 insufficiency
 Premature ovarian failure. *See* Primary ovarian
 insufficiency
 Premature ventricular contractions (PVCs)
 in acute coronary syndromes, 1493
 characteristics of, 1491-1492
 in congenital heart disease, 1494
 definition of, 1489
 in depressed ventricular function and heart
 failure, 1493-1494
 diagnosis of, 1492
 etiology of, 1492
 in heart failure, 1493-1494
 in hypertrophic cardiomyopathy, 1494
 idiopathic, 1492-1493
 idioventricular rhythms, 1489, 1490f, 1494
 multifocal, 1489, 1490f
 palpitations in, 254
 unifocal, 1489, 1490f
 ventricular dysfunction and, 1494
 Premunition, 1372
 Prenatal diagnosis, 83e-3-4, 83e-4t, 85e-9, 444
 Preoperative questionnaire, 51, 51t
 Preosteoclasts, 2143
 Prepatellar bursitis, 2248
 Presacral edema, 1504
 Presbycusis, 219, 219f
 Presbyopia, 195
 Presbyosmia, 214
 Presenilins, 2601
 Pre-S genes, of HBV, 2005f, 2006
 Pressure-control ventilation (PCV), 1742. *See*
also Mechanical ventilation
 Pressure-immobilization, for snakebite, 2735
 Pressure-natriuresis phenomenon, 1613
 Pressure-support ventilation (PSV), 1742, 1743t.
See also Mechanical ventilation
 Pressure (decubitus) ulcers, 830, 1743, 2660
 Pressure urticaria, 2118
 Pressure-volume curve
 of chest wall, 306e-1f
 of lung, 306e-1, 306e-1f
 Presymptomatic testing, 448
 Presynaptic rundown, 2701
 Presyncope, 142
 Presystemic elimination, 33, 33f
 Presystolic accentuation, 51e-6
 Pretest probability of disease, 21-22, 23f
 Preventive care. *See also specific diseases*
 in adult-onset genetic disorders, 450-451, 450t
 counseling in, 30, 30-31t, 30t
 screening tests in, 26-27, 29t
Prevotella spp., 1094
Prevotella spp. infections. *See also* Anaerobic
 bacterial infections, mixed
 of bone, 1099
 in dog-bite wound, 167e-1
 of head, neck, and mouth, 1097
 in human bite-wound, 167e-2
 Priapism, 325, 634
 Prickly heat, 479e-1
 Primaquine
 action of, 246e-12
 adverse effects of, 40, 246e-1t, 246e-12, 1380t,
 1384
 contraindications to, 246e-1t, 1378t, 1381,
 1384
 drug interactions of, 246e-1t, 246e-6t
 genetic variations in response to, 39t, 40
 for malaria, 1378t, 1380t, 1381
 for malaria prophylaxis, 1383-1384, 1383t
 overdose/poisoning with, 473e-11t
 for PCP, 1361t
 pharmacology of, 246e-12, 1380t
 Primary afferent(s), 87, 87f
 Primary afferent nociceptors, 87, 87f, 88f, 89f
 Primary alveolar proteinosis, 375e-2
 Primary biliary cirrhosis. *See* Biliary cirrhosis
 Primary care
 definition of, 13e-1
 revitalization of, 13e-4-6, 13e-5f
 worldwide access to, 13e-1. *See also* Low- and
 middle-income countries
 Primary Care PTSD (PC-PTSD) Screen,
 471e-5t
 Primary central nervous system lymphoma
 (PCNSL)
 diagnosis of, 601, 602f, 910f
 in HIV infection, 1270-1271, 1271f
 incidence of, 601
 pathogenesis of, 601
 treatment of, 601-602
 Primary cough headache, 2597
 Primary cutaneous CD30+ T-cell lymphoma,
 135e-4
 Primary (pleural) effusion lymphoma, 135e-2,
 214e-7, 1194, 1270
 Primary exertional headache, 2597
 Primary hyperhidrosis, 2643
 Primary immune deficiency diseases, **2103**
 of adaptive immune system, 2104t, **2107**
 B lymphocyte deficiencies. *See* B cell(s),
 deficiencies of
 T lymphocyte deficiencies. *See* T cell(s),
 deficiencies of
 B cell differentiation in, 2110f, 2111
 classification of, 2104t
 colitis and, 2113
 consequences of, 2103-2104
 definition of, 2103
 diagnosis of, 2104, 2105t, 2111
 diseases associated with, 375e-1-2
 effector pathways in, 2107, 2107f
 gene therapy for, 91e-1-2
 genetic considerations in, 2103
 HSCT for, 139e-5
 of innate immune system, **2104**
 asplenia, 375e-1, 413, 788f, 2104-2105
 chronic granulomatous diseases. *See*
 Chronic granulomatous diseases
 (CGDs)
 complement deficiency, 372e-28, 372e-29t,
 2106-2107
 diseases associated with, 375e-1
 leukocyte adhesion deficiency. *See*
 Leukocyte adhesion deficiency (LAD)
 Mendelian susceptibility to mycobacterial
 disease, 2106
 severe congenital neutropenia, 375e-1,
 2104, 2105f
 toll-like receptor pathway deficiencies, 2106
 phagocytic cells in, 2104, 2105f
 regulatory defects and, 2104t, 2112
 in autoimmune lymphoproliferative
 syndrome, 2113
 in autoimmune polyendocrinopathy-
 candidiasis-ectodermal dysplasia
 syndrome. *See* Autoimmune
 polyendocrine (polyglandular)
 syndrome-1 (APS-1)
 in hemophagocytic lymphohistiocytosis,
 2112
 in immunodysregulation
 polyendocrinopathy X-linked
 syndrome, 2113
 in PLC γ -associated antibody deficiency and
 immune dysregulation, 2113
 T cell differentiation in, 2107, 2107f, 2108f
 variable expression of, 2113
 Primary intraventricular hemorrhage, 2582-
 2584
 Primary lateral sclerosis, 2636, 2659
 Primary myofibrosis, **674**
 clinical features of, 675
 complications of, 675
 diagnosis of, 675
 bone marrow examination in, 81e-5f, 397f,
 675f
 peripheral blood smear in, 81e-5f, 395f, 675f
 diseases associated with, 674t
 genetic considerations in, 672, 674-675
 prognosis of, 675-676, 676t
 skin manifestations of, 369
 splenomegaly in, 411
 treatment of, 102e-5t, 676
 Primary olfactory cortex, 211, 212f
 Primary ovarian failure, 437
 Primary ovarian insufficiency, 336
 Primary peritoneal serous carcinoma, 120e-3-4

- Primary pigmented nodular adrenal disease (PPNAD), 2314, 2343
- Primary polydipsia, 2276, 2277, 2279
- Primary progressive aphasia, 2603. *See also* Frontotemporal dementia (FTD)
- Primary sclerosing cholangitis (PSC), **2061**
 cholangiocarcinoma and, 552
 clinical features of, 284, 1958, 2061, 2085–2086
 diagnosis of
 cholangiography in, 284, 1958, 2061–2062, 2079f, 2085–2086
 laboratory tests in, 1937, 2061
 liver biopsy in, 366e-5f
 diseases associated with, 2084–2085
 drug-induced, 2023
 in HIV infection, 2086
 in IBD, 1958
 IgG4-related, 391e-1, 391e-2. *See also* IgG4-related disease (IgG4-RD)
 natural history of, 2086
 small duct, 1958, 2086
 treatment of, 2062, 2068, 2086
- Primary sex headache, 2597
- Primary stabbing headache, 2597–2598
- Primary taste cortex (PTC), 214
- Primary thunderclap headache, 2597
- Primate immunodeficiency viruses, 1216, 1216f, 1219. *See also* HIV
- Primidone
 adverse effects of, 646t, 2554t
 for ataxia/tremor, 2620
 drug interactions of, 2554t
 for epilepsy, 2554t
 for myoclonus, 2623
 overdose/poisoning with, 473e-10t, 2765t
 therapeutic monitoring of, 2765t
- Priming, T cells, 103e-20
- Primitive neuroectodermal tumors (PNETs), 119e-4. *See also* Ewing's sarcoma
- Primitive reflexes, 2539
- Principal cells, 64e-2f, 304, 332e-5f, 332e-9
- Principal coordinates analysis, 86e-1t, 86e-3f, 86e-5
- PR interval, 273e-1
- Prinzmetal's variant angina, 1456, 1579, 1588, **1598**
- Prion(s)
 characteristics of, 214e-1
 decontamination of, 453e-5–6
 definition of, 453e-1, 453e-1t
 diseases due to. *See* Prion disease(s)
 strains of, 453e-2–3, 453e-2t
 structure of, 453e-1f
 terminology related to, 453e-1t
 transmissibility of, 453e-1–2
- Prion disease(s), **453e-1**
 classification of, 444e-10, 444e-12t
 clinical features of, 451e-2t
 Creutzfeldt-Jakob disease. *See* Creutzfeldt-Jakob disease
 epidemiology of, 453e-1t, 453e-2
 genetic considerations in, 451e-2t
 global considerations in, 453e-1–2, 453e-1t
 Huntington disease–like disorders, 2622
 inherited, 453e-3
 neurodegenerative, 444e-12t, 453e-6
 pathogenesis of, 444e-8f, 444e-10–11, 453e-1t, 453e-2–3, 453e-2f
 research in, 453e-6
 species barrier in, 453e-3
 sporadic, 453e-3
 types of, 453e-1–2, 453e-1t
- Prion proteins, 453e-1, 453e-1f, 453e-2t
- Prion rod, 453e-1t
- Pritelivir, 215e-9
- PRKAG2 deficiency, 433e-2t, 433e-5
- PRKAG2 gene mutations, 1566
- PRKARIA gene, 289e-1, 2266, 2314
- PRNP gene, 172t, 453e-1t, 453e-3, 2622
- Probenecid
 action of, 332e-8
 adverse effects of, 282
 for CMV infection prophylaxis in HIV infection, 1252t
- drug interactions of, 3, 42t, 103e-14, 246e-6t
 for gonococcal infections, 1009t
 for gout, 2234
 for neurosyphilis, 1139t
 for syphilitic meningitis, 899
- Probiotics, 263, 1970
- Procainamide
 action of, 273e-6t
 adverse effects of
 lupus syndrome, 40, 44, 2218t
 in myasthenia gravis, 2703
 myopathy, 462e-20, 2199
 nonarrhythmic, 1480t
 proarrhythmic, 1480t
 pure red cell aplasia, 668t
 thrombocytopenia, 727t
 for arrhythmias, 1479t
 for cardiac arrest, 1769f, 1770
 dosage of, 1479t
 genetic variations in response to, 39t, 40
 metabolism of, 34t, 35
 pharmacology of, 1479t
 for preexcited tachycardias, 1482
 for PSVT, 1483f
 therapeutic monitoring of, 2765t
 for ventricular arrhythmias, 1498
 for ventricular tachycardia, 1608
- Procalcitonin, 806
- Procarbazine
 action of, 103e-8, 103e-9t
 adverse effects of, 103e-9t
 gonadal dysfunction, 103e-25, 2365
 myelosuppression, 103e-23
 neurologic, 607t
 drug interactions of, 103e-9t
 for oligodendroglioma, 601
- Procaspase-9, 102e-9f
- Process (quality theory), 12e-2
- Prochlorperazine
 adverse effects of, 284
 for migraine, 2592, 2592t
 for nausea and vomiting, 61–62, 103e-25, 260t, 261, 474
 parenteral, 2592t
 for scambroid poisoning, 2744
- Procidencia. *See* Rectal prolapse
- Proctitis
 C. trachomatis, 1168, 1169, 1170t
 HSV, 1178, 1182t, 1956
 sexually acquired, 870t, 881–882, 1005, 1168, 1169
 ulcerative colitis and, 1952
- Proctocolitis, sexually acquired, 881–882
- Proctopathy, radiation, 346e-1v
- Proctosigmoidoscopy. *See* Sigmoidoscopy
- Prodrugs, 35
- Progenitor cells, 89e-4f, 454
- Progeria, 435, 436t
- Progeroid syndromes, 94e-4
- Progesterone, 324, 2378, 2762t
- Progesterone receptors, 120e-2t, 528
- Progestins
 adverse effects of, 253t
 with estrogens, 2498
 in oral contraceptives, 2390–2391
 in perimenopause, 2382
 for polycystic ovarian syndrome, 337
 in postmenopausal hormone therapy. *See* Postmenopausal hormone therapy
- Progestogens, 530t
- Proglottids, 255e-1
- Prognathism, 237
- Program for All Inclusive Care of the Elderly (PACE), 82–83
- Programmed cell death. *See* Apoptosis
- Programmed death theory of aging, 94e-2
- Progranulin, 2603
- Progressive diaphyseal dysplasia (Camurati-Engelmann disease), 426e-5
- Progressive external ophthalmoplegia
 clinical features of, 41e-1v, 85e-6, 85e-6t, 207, 462e-15–16
 diagnosis of, 462e-15–16
- differential diagnosis of, 2703
 genetic considerations in, 85e-6, 85e-6t, 85e-7f, 462e-16
- Progressive familial intrahepatic cholestasis, 284, 2003t, 2004
- Progressive massive fibrosis, 1690
- Progressive multifocal leukoencephalopathy (PML), **899**
 in cancer patient, 488, 488t, 490
 clinical features of, 899
 diagnosis of, 899–900
 in HIV infection, 899–900, 1266
 monoclonal antibodies and, 372e-31, 1962, 2147, 2669
 pathology of, 899
 TNF- α inhibitors and, 348
 treatment of, 899–900
- Progressive muscular atrophy. *See* Spinal muscle atrophy (SMA)
- Progressive myoclonus epilepsy, 2545t
- Progressive myopathy, 436e-6f
- Progressive rubella panencephalitis, 900
- Progressive supranuclear palsy syndrome
 clinical features of, 173, 2604, 2612t
 eye movements in, 41e-1v, 2604, 2610
 falls in, 163, 165
 gait disorders in, 163, 2610
 neuropathology of, 2604–2605, 2604f, 2605f, 2610–2611
- Proguanil
 action of, 246e-12
 adverse effects of, 246e-4t, 1380t, 1384
 drug interactions of, 246e-4t
 for malaria, 1380t
 for malaria prophylaxis, 1382, 1383, 1383t, 1384
 pharmacology of, 246e-12, 1380t
 in pregnancy and lactation, 246e-4t, 1384
 resistance to, 1384
- Proinsulin, 2434
- Project BioShield, 261e-10
- PROK2 gene mutations, 2363, 2364t
- PROK2R gene mutations, 2363, 2364t
- Prokinetic agents, 260t, 263
- Prolactin, **401e-1**
 action of, 401e-3
 expression and regulation of, 401e-1t
 laboratory evaluation of, 2258t, 2762t
 secretion of, 401e-2
 synthesis of, 401e-1–2
- Prolactinoma, **2267**
 clinical features of, 2267
 diagnosis of, 2267
 etiology of, 2267
 in MEN 1, 2336t, 2339
 during pregnancy, 2268–2269
 prevalence of, 2267
 screening tests for, 2262t
 treatment of, 2267–2268, 2268f
- Prolidase deficiency, 434e-2t
- Proliferative inflammatory atrophy, of prostate, 580
- Proline metabolism disorders, 434e-2t, 434e-3t
- Polymorphocytic leukemia, 412
- Promethazine, 151t, 389t
- Prometrium, 337
- PROMM (proximal myotonic myopathy), 462e-3, 462e-9, 462e-10
- Promonocytes, 421
- Promoter (carcinogenesis), 476
- Promyelocytes, 414f, 415
- Promyelocytic leukemia. *See* Acute promyelocytic leukemia
- Pronator drift, 2538
- Pronormoblast, 392
- Pronormoblasts, giant, 668, 668f
- Proopiomelanocortin (POMC)
 deficiency, 2325t
 in obesity, 415e-3, 415e-4f, 415e-4t
- PROP-1
 in anterior pituitary development, 401e-1, 401e-1t, 2256
 in congenital hypothyroidism, 2284t
 in GH deficiency, 2258

- Proteus* spp., 1033
Proteus spp. infections, **1033**
 antibiotic resistance in, 934, 1033–1034
 clinical features of, 1033
 diagnosis of, 1033
 epidemiology of, 1025
 renal abscess, 851
 treatment of, 1033–1034
 urinary tract, 862, 1033, 1871
- Prothrombin, 402f
 Prothrombin complex concentrate, 735, 739
 Prothrombin deficiency, 404, 733t
 Prothrombin gene mutations, 47, 404–405, 745
 Prothrombin inhibitors, 740
 Prothrombin time (PT), 732
 in acute viral hepatitis, 2016
 causes of abnormal values, 406t, 464t
 in coagulation disorders, 733t
 coagulation factors measured in, 405, 406f, 733f
 in hepatobiliary diseases, 1998, 1998t
 for monitoring warfarin therapy, 755
 in nutritional assessment, 464t
 prolonged, 283, 406t
 sensitivity of, 405
- Proton pump inhibitors (PPIs)
 adverse effects of, 263, 568, 644t, 1857, 1920–1921, 2218t
 for diarrhea, 272
 drug interactions of, 42t, 246e–3t, 1921
 for GERD, 263, 1908
 in *H. pylori* eradication, 263, 1041, 1042t
 hypergastrinemia and, 568
 for metabolic alkalosis, 322
 for NSAID-induced ulcer prophylaxis, 1924, 1924t
 for peptic ulcer disease, 276–277, 1920–1921, 1920t, 1924
 pharmacology of, 1921
 for stress-related mucosal injury, 1930
 for systemic mastocytosis, 2121
 for ZES, 568, 1929
- Protooncogenes, 101e–2
 Protoporphyrin, erythropoietic. *See* Erythropoietic protoporphyria
 Protoporphyrins, 390
 free erythrocyte, 2525
 red cell, 627f, 628
- Prototheca wickerhamii* infection, 926
- Protozoa
 adherence of, 145e–2t, 145e–3
 blood and tissue
Babesia. *See Babesia* spp.
Leishmania. *See Leishmania* spp.
Plasmodium. *See Plasmodium* spp.
Toxoplasma gondii. *See Toxoplasma gondii*
Trypanosoma. *See Trypanosoma* spp.
 entry into host, 145e–1
 free-living amebas
Acanthamoeba. *See Acanthamoeba* spp.
Balamuthia. *See Balamuthia* spp.
Naegleria. *See Naegleria* infection
 vs. helminthic parasites, 255e–1
 intestinal
Cryptosporidium. *See Cryptosporidium*/
 cryptosporidiosis
Cyclospora cayatanensis. *See Cyclospora*
cayatanensis/cyclosporiasis
Cystoisospora belli. *See Cystoisospora belli*/
 cystoisosporiasis
Entamoeba histolytica. *See Entamoeba*
histolytica
Giardia. *See Giardia* spp.
 Microsporida. *See Microsporida*/
 microsporidiosis
 laboratory identification of, 245e–4t, 245e–6t
 tissue invasion by, 145e–8
- Protriptyline, 190
Providencia infections, 1035, 1871
 Provirus, 225e–1
 PROX1 gene, 1241t
 Proximal femoral neuropathy, 463e–5, 463e–5f
 Proximal interphalangeal (PIP) joint,
 rheumatoid arthritis in, 2136, 2137f
- Proximal myotonic myopathy (PROMM),
 462e–3, 462e–9, 462e–10
- Proximal tubule
 disorders involving, 332e–7t
 functions of, 332e–3f, 332e–6–8
 toxic injury to, 333e–4
- Prozone phenomenon, 391e–2, 1137
- PrP amyloid, 453e–1t
 PrP^{sc}, 453e–1, 453e–1f, 453e–1t, 453e–2f. *See also*
 Prion disease(s)
- PRPP (phosphoribosylpyrophosphate), 431e–2
 PRPP (phosphoribosylpyrophosphate)
 synthetase overactivity, 431e–3t,
 431e–5
- PrP^{sc}, 172t, 453e–1, 453e–1f, 453e–1t, 453e–2f.
See also Prion disease(s)
- PRRs. *See* Pattern recognition receptors (PRRs)
 PRRT (peptide radioreceptor therapy), 575
 PRSSI gene mutations, 554, 2092, 2098
- Prucalopride, 274
 Prune juice, 62t
- Pruritus
 in atopic dermatitis, 345
 in biliary cirrhosis, 2061
 definition of, 340t
 drug-induced, 381
 in liver disease, 1991
 in scabies, 2744
 treatment of, 345
- Prussian blue, 81e–8f, 263e–6, 263e–6t, 472e–3
 PS-1 gene mutations, 2601
 PS-2 gene mutations, 2601
- PSA. *See* Prostate-specific antigen (PSA)
- PSC. *See* Primary sclerosing cholangitis (PSC)
- P-selectin, 276, 742
- Pseudallescheria boydii*, 491
 Pseudoachalasia, 1904
 Pseudoaldosteronism. *See* Liddle's syndrome
 (pseudoaldosteronism)
- Pseudoallergic fish poisoning, 2743
- Pseudoanemia, 636
- Pseudoaneurysm(s)
 aortic, 289e–3, 1637
 pancreatic, 2095f, 2097
 pericardial, 1610
- Pseudoappendicitis, 1077
- Pseudoarthrosis, 2171
- Pseudoathetosis, 158
- Pseudobulbar palsy, 154
- Pseudocholinesterase, 34t
- Pseudocowpox, 220e–1t
- Pseudocyst(s), pancreatic, 2093t, 2097
- Pseudodementia, 2608, 2715
- Pseudodiarrhea, 265
- Pseudodiverticulosis, esophageal, 1903, 1903f
- Pseudodominance, 438
- Pseudoephedrine
 adverse effects of, 305
 for allergic rhinitis, 2122, 2123f
 for bladder dysfunction, 2672
 contraindications to, 2122
 for orthostatic hypotension, 146
- Pseudogout, 372e–8t, 2235
- Pseudohalutosis, 242
- Pseudohermaphroditism
 female. *See* 46,XX disorders of sex
 development
 male. *See* 44,XY disorders of sex
 development
- Pseudo-Hurler polydystrophy, 432e–3t
- Pseudohyperkalemia, 309, 309t, 653, 1795
- Pseudohypertension, 1626
- Pseudohypertrophic muscular dystrophy. *See*
 Duchenne's muscular dystrophy
- Pseudohypoaldosteronism type I, 310, 332e–7t
- Pseudohypoaldosteronism type II (Gordon's
 syndrome)
 clinical features of, 310, 1620t
 genetic considerations in, 310, 332e–7t,
 332e–8, 1620t
 treatment of, 310
- Pseudohypoglycemia, 2430
- Pseudohypokalemia, 305
- Pseudohyponatremia, 301
- Pseudohypoparathyroidism (PHP)
 classification of, 2485, 2486t
 genetic considerations in, 430–431, 2476f,
 2486–2487, 2486f
 hypocalcemia and, 2485
 treatment of, 2487
- Pseudo-Kaposi's sarcoma, 367
- Pseudolymphoma (cutaneous lymphoid
 hyperplasia), 353, 364t, 365, 366
- Pseudomembrane(s)
 in diphtheria, 978, 978f
 in sulfur mustard exposure, 262e–3, 262e–3f
 in tularemia, 1068
- Pseudomembranous colitis, 855, 858, 858f, 1882f.
See also Clostridium difficile-associated
 disease
- Pseudomonas aeruginosa*, **1042**
 adherence of, 145e–2–3, 145e–2t
 antibiotic resistance in, 811, 934, 941t, 1047
 avoidance of host defenses by, 145e–3,
 145e–4f, 1043
 CTFR protein and, 145e–4f, 145e–5, 1043
 epidemiology of, 1043
 global considerations in, 1043
 growth after entry, 145e–4
 inflammatory response to, 1043
 laboratory identification of, 150e–2t, 1043
 motility of, 1043
 surface structures of, 145e–2f, 1043
 tissue injury by, 1043
 virulence factors of, 1043, 1043t
- Pseudomonas aeruginosa* infections, **1044**
 arthritis, 834, 835
 bacteremia, 1044, 1045t
 bone and joint, 1045t, 1046
 bronchiectasis, 1695
 burn-wound, 166e–1, 166e–2f, 166e–3, 1043
 in cancer patient, 486t, 487, 492
 cellulitis, 830
 chronic respiratory, 1044, 1045–1046
 CNS, 1045t, 1046
 in cystic fibrosis, 145e–4f, 145e–5, 1044, 1697
 diagnosis of, 343, 1043
 ear, 228–229, 1045t, 1046–1047
 ecthyma gangrenosum, 76e–12f, 135, 369, 830
 endovascular, 1045t, 1046
 epidemiology of, 1043
 in febrile neutropenia, 1045t, 1047
 folliculitis, 131t, 134, 356, 829, 1047
 global considerations in, 1045
 health care-associated pneumonia, 803t
 in HIV infection, 1047
 in lung abscess, 814
 mastoiditis, 231
 multidrug-resistant, 1045t, 1047
 nail, 1047
 ocular, 1045t, 1046
 osteomyelitis, 839, 840t, 841t, 844
 pathogenesis of, 1043–1044, 1043t
 perichondritis, 228
 pneumonia, 1044–1045, 1045t
 prevention of, 492
 risk factors for, 805
 sinusitis, 226
 in transplant recipient, 926
 treatment of, 808t, 830, 940t, 1045t
 urinary tract, 862, 1045t, 1047
 ventilator-associated pneumonia, 811, 812,
 915
- Pseudomonic acid. *See* Mupirocin
- Pseudoobstruction, intestinal. *See* Intestinal
 pseudoobstruction (Ogilvie's disease)
- Pseudoparalleloma, 203, 203f
- Pseudoporphyria, 361t, 362, 379, 379f
- Pseudopseudohypoparathyroidism (PPHP), 430,
 2486f, 2486t
- Pseudoterranova decipiens*, 1416
- Pseudothrombocytopenia, 725, 726f
- Pseudotumor(s)
 in hemophilic arthropathy, 2241
 in IgG4-related disease, 391e–2
 orbital, 207
- Pseudotumor cerebri, 49, 203, 2293
- Pseudotumor syndrome, 733

- Pseudoxanthoma elasticum, 364t, 365, 732, 1442
 PSI (Pneumonia Severity Index), 806
 Psittacosis, 1172–1173
 Psoas abscess, 852
 Psoralens, 361, 380, 389t, 391, 2177
 Psoriasis, **347**
 in ankylosing spondylitis, 2171
 clinical features of, 354t
 distribution of skin lesions, 340, 341t, 342f, 347t, 354t
 erythroderma, 353t, 354t
 morphology of skin lesions, 341t, 354t
 papules and plaques, 76e-1, 76e-1f, 343f, 347t, 353t
 diagnosis of, 354t
 drugs exacerbating, 378
 etiology of, 347
 genetic considerations in, 348
 guttate, 347
 histologic features of, 347t
 in HIV infection, 1263
 HLA gene association with, 373e-8t
 hypothermia in, 478e-1
 in IBD, 1958
 inflammatory mediators in, 372e-29t
 inverse, 347
 KIRs with, 372e-15t
 plaque-type, 347
 pustular
 vs. AGEP, 382
 clinical features of, 76e-1f, 347, 353, 356
 drugs exacerbating, 378
 treatment of, 96e-7, **348**, 348t, 354t, 387, 391
 Psoriatic arthritis, **2175**
 aortic aneurysm in, 1638
 aortitis in, 290e-3, 1638
 CASPAR criteria for, 2177, 2177t
 clinical features of, 348, 353, 2176–2177, 2176f
 definition of, 2175
 diagnosis of, 2177, 2177t
 epidemiology of, 2175–2176
 historic background of, 2175
 inflammasome activity in, 372e-8t
 KIRs with, 372e-15t
 pathogenesis of, 2176
 pathology of, 2176
 treatment of, 348t, 2177–2178
 Psoriatic spondylitis, 373e-8t
 PSORS1C3 gene, 1240t, 1241
 P-SPIKES, for communicating bad news, 57–58, 57t
 PSTP11 gene, 372e-8t, 2215
 PSV (pressure-support ventilation), 1742, 1743t.
 See also Mechanical ventilation
 PSVT. See Paroxysmal supraventricular tachycardia (PSVT)
 Psychiatric/psychological disorders, **2708**
 abdominal pain in, 106t
 in alcohol use disorder, 2724–2725
 anxiety disorders. See Anxiety disorders
 biology of, **465e-1**
 bipolar disorder. See Bipolar disorder
 classification of, 2708
 in cobalamin deficiency, 643
 vs. dementia, 2608
 depression. See Depression/depressive disorders
 eating and feeding disorders, **2719**
 fatigue in, 152
 in fibromyalgia, 2239
 in folate deficiency, 643
 gait disorders in, 164
 genetic considerations in, 465e-1–2, 465e-2t
 global considerations in, 17, 2708
 at high altitudes, 476e-5
 in the homeless, 2723
 inflammatory mechanisms in, 465e-4–5
 insomnia in, 190
 irritable bowel syndrome and, 1966
 mood disorders. See Mood disorders
 palpitations in, 254
 personality disorders, **2720**
 prevalence of, 2708
 primary care physician's responsibility in, 2708
 schizophrenia. See Schizophrenia
 signal transduction in, 465e-2–3, 465e-3f
 somatic symptom disorder, **2718**
 syncope in, 147, 148
 systems neuroscience of, 465e-3–4, 465e-4f, 465e-5t
 taste disorders in, 215
 in war veterans. See War veterans, neuropsychiatric illnesses in
 in Wilson's disease, 2519
 in women, 6e-5
 Psychogenic amnesia, 2608
 Psychogenic polydipsia, 2276, 2279
 Psychogenic seizures, 2551
 Psychosis(es). See also Schizophrenia
 alcohol-induced, 2725
 in Huntington's disease, 2622
 PCP-induced, 469e-4
 in PD, 2617
 postictal, 2558
 Psychosocial short stature, 2258
 Psychostimulants, 63, 65, 465e-5t
 Psychotherapy
 for anxiety disorders, 2709–2710
 for depression, 2716
 for panic disorder, 2709
 for phobic disorders, 2712
 for PTSD, 2713
 referral for, 2708
 for stress disease, 2712
 Psyllium, 1969, 1970t
 PT. See Prothrombin time (PT)
 PTCH gene mutations, 101e-6t, 500, 500f, 602
 PTEN, 89e-4
 PTEN gene mutations
 in autism spectrum disorders, 465e-1, 465e-2t
 in brain tumors, 599, 599t
 in breast cancer, 523
 cancers associated with, 101e-6t
 in Cowden's syndrome, 365, 2344
 in endometrial cancer, 596
 in lung cancer, 507t
 in melanoma, 496
 in thyroid cancer, 2305t
 Pterygium, 199
 PTH. See Parathyroid hormone (PTH)
 Pthiriasis palpebrum, 2746
 Pthirus pubis (pubic lice), 2745–2746
 PTH-related peptide, 313
 PTHrP. See Parathyroid hormone-related protein (PTHrP)
 PTNP22 gene mutations, 377e-3, 2140
 Ptoxis, 41e-1v, 207–208, 462e-2t
 PTPS (6-pyruvoyl-tetrahydropterin synthase) deficiency, 434e-2t
 PTSD. See Posttraumatic stress disorder (PTSD)
 Puberty
 definition of, 2357
 delayed. See Delayed puberty
 normal development during
 female, 2379, 2379t
 male, 2357–2358, 2357f
 precocious. See Precocious puberty
 Pubic lice (*Pthirus pubis*), 2745–2746
 Puborectal muscle, 265f
 PUBS (percutaneous umbilical blood sampling), 83e-4, 444
 Pudendal nerve terminal motor latency, in fecal incontinence, 1975
 "Puff of smoke" lesion, 2568
 Pulmonary alveolar proteinosis (PAP), 1714
 Pulmonary angiography, 1634, 1666–1667
 Pulmonary arterial hypertension (PAH), **1658**.
 See also Pulmonary hypertension
 conditions associated with, 1658
 in connective tissue disease, 1658
 diagnosis of, 47e-4, 308e-18f
 familial, 87e-3, 87e-3t
 in HIV infection, 1255, 1658
 idiopathic (primary), 271e-9f, 271e-9v, 1658
 pathogenesis of, 1655, 1656f, 1657t
 portal hypertension and, 1658
 in sarcoidosis, 2207
 in systemic sclerosis, 1658, 2161, 2164
 systems biology applied to, 87e-3, 87e-3t
 treatment of, 320e-1t, 1659–1660, 1660t
 Pulmonary artery, response to hypoxia, 248
 Pulmonary artery catheter
 in cardiogenic shock, 1760
 complications of, 1734
 in pulmonary edema, 1763
 in shock patient, 1747
 Pulmonary artery wedge pressure, 1444
 Pulmonary capillary wedge pressure
 in cardiogenic shock, 1759
 in pulmonary hypertension, 1657
 in sepsis/septic shock, 1755
 in shock, 1746, 1748t
 Pulmonary edema, **47e-4**
 cardiogenic, 47e-4, 1763
 chest radiography in, 47e-5, 308e-7f
 cyanosis in, 249
 diagnosis of, 1763
 differential diagnosis of, 47e-5
 dyspnea in, 47e-2t, 47e-4
 etiology of, 47e-5t
 hemodynamic patterns in, 1762t
 high-altitude. See High-altitude pulmonary edema (HAPE)
 in malaria, 1374, 1381
 noncardiogenic, 47e-4, 47e-5t, 1763
 pathophysiology of, 47e-4, 306e-3
 physical examination in, 47e-5, 1662
 reexpansion, 1764
 treatment of, 1761f, 1763
 in upper airway obstruction, 1764
 Pulmonary embolism (PE), **1631**
 adequacy of ventilation in, 306e-3
 after subarachnoid hemorrhage, 1787
 in burn patient, 166e-2
 catheter-based therapies for, 296e-5
 chest pain in, 96t, 97–98
 classification of, 1632
 diagnosis of
 blood tests, 1632
 cardiac biomarkers, 1594, 1633
 chest CT, 1633, 1633f
 chest radiograph, 101, 1633
 clinical decision rules, 1632t
 clinical evaluation, 1632
 clinical pearls, 1632
 CT, 308e-17f, 1664
 ECG, 101, 269e-11f, 1633
 echocardiography, 1634
 imaging algorithms, 1632f, 1634f
 lung scan, 1633–1634
 MRI, 1634
 pulmonary angiography, 1634, 1667–1668
 scintigraphic imaging, 1664
 Wells clinical prediction rule, 24t
 dyspnea in, 47e-2t, 47e-4, 1632
 epidemiology of, 743
 hemoptysis in, 100, 246
 in HIV infection, 1262
 incidence of, 274, 740
 in mitral stenosis, 1540
 nonthrombotic, 1632
 paraneoplastic, 613
 pathophysiology of, 1631–1632
 pleural effusion in, 1717
 postmenopausal hormone therapy and, 2384t
 in pregnancy, 47
 prevention of, 1636–1637, 1637t
 right ventricular dysfunction in, 270e-7, 1635
 risk stratification in, 1634
 septic, 958, 958f
 treatment of
 algorithm for, 1634f
 anticoagulants, 1634–1635, 1635t
 catheter-directed therapy, 1636
 duration of anticoagulation, 1635–1636
 emotional support, 1636
 fibrinolysis, 1636
 inferior vena cava filter, 1636
 in massive PE, 1636
 pulmonary embolectomy, 1636
 pulmonary thromboendarterectomy, 1636
 ventilation-perfusion heterogeneity in, 306e-4

- Pulmonary fibrosis
bleomycin-related, 621
bronchiectasis in, 1694
chemotherapy-associated, 103e-8, 103e-13
dyspnea in, 47e-1t, 47e-4
idiopathic. *See* Idiopathic pulmonary fibrosis
as late consequence of cancer/cancer therapy, 621
pathophysiology of, 306e-3, 1709f
pulmonary function values in, 306e-5, 306e-5f
radiation-induced, 103e-4
in systemic sclerosis, 2157–2158, 2158f
Pulmonary function tests
in asthma, 1675, 1675f
common abnormalities in, 306e-5f
in COPD, 1704, 1704t
in hypersensitivity pneumonitis, 1682
in interstitial lung diseases, 1710
in occupational lung disease, 1687
in pulmonary hypertension, 1656–1657
in respiratory disease evaluation, 1662, 1693
Pulmonary heart disease. *See* Cor pulmonale
Pulmonary hypertension, **1655**
arterial. *See* Pulmonary arterial hypertension
clinical features of, 1655
chest pain, 96t, 98
dyspnea, 47e-2, 47e-2t, 1655
tricuspid regurgitation, 1655
diagnosis of, 1655–1657
chest radiography, 47e-4, 1658f
CT, 1656, 1658f
ECG, 269e-11f
echocardiography, 271e-9f, 271e-9v, 1655–1656, 1657f
heart sounds, 1447, 1655
pulmonary function tests, 1656–1657
high-altitude, 476e-6
in left heart disease, 1658–1659
in lung disease, 1659
in mitral stenosis, 1539
pathogenesis of, 1655, 1657t
in pregnancy, 47
Raynaud's phenomenon in, 1649
in sarcoidosis, 1659
in schistosomiasis, 1659
in sickle cell disease, 1659
in sleep-disordered breathing, 1657
thromboembolic, 1659
treatment of, 1660
Pulmonary infarction, 105, 1632
Pulmonary infiltrates with eosinophilia
acute eosinophilic pneumonia, 1684, 1684t
allergic bronchopulmonary aspergillosis.
See Allergic bronchopulmonary aspergillosis (ABPA)
C. pseudotuberculosis, 980
chronic eosinophilic pneumonia, 1685
classification of, 1683–1684
disorders associated with, 1684t
drug- and toxin-induced, 1687
eosinophilic granulomatosis with polyangiitis.
See Eosinophilic granulomatosis with polyangiitis (Churg-Strauss syndrome)
global considerations in, 1687
hypereosinophilic syndromes. *See* Hypereosinophilic syndromes (HES)
infectious causes of, 1685–1686, 1685t
pathophysiology of, 1683
Pulmonary Langerhans cell histiocytosis, 1713
Pulmonary lymphangioleiomyomatosis, 308e-5f, 1714–1715
Pulmonary mass/nodule. *See also* Lung cancer
hamartoma, 515
solitary, 515–516, 516t, 517f, 1666
Pulmonary rehabilitation, 47e-4, 1706
Pulmonary risk assessment, preoperative, 54–55, 54t
Pulmonary vascular resistance, 1746, 1748t
Pulmonary veins, anomalous drainage of, 271e-10f
Pulmonic regurgitation, **1550**
clinical features of, 1550
etiology of, 1547t, 1550
heart sounds/murmur in, 51e-6–7, 1449, 1550
treatment of, 1550
Pulmonic stenosis, **1549**
in carcinoid syndrome, 564
clinical features of, 1549
congenital, 1525
diagnosis of, 1524–1525, 1525f, 1549–1550
etiology of, 1547t, 1549
heart sounds/murmur in
auscultation of, 1447, 1448, 1525, 1549–1550
duration of, 51e-3, 51e-4f
intensity and radiation of, 51e-2f
timing of, 51e-1f
pathophysiology of, 1549
treatment of, 296e-5, 1525f, 1550
Pulp, dental, 236
Pulpitis, 236, 1097
“Pulsatile tinnitus,” 2586
Pulse
assessment of, 1445–1446, 1445f, 1446f
in peripheral artery disease, 1447, 1447f, 1643
Pulsed-field gel electrophoresis, 150e-8
Pulsed wave Doppler, 270e-2, 270e-2f. *See also* Echocardiography
Pulseless disease. *See* Takayasu's arteritis
Pulseless electrical activity, 1764, 1769–1770, 1769f
Pulse oximetry, 1662, 1733
Pulse pressure, 292e-1v, 1611, 1729, 1732f
Pulsus alternans, 1446
Pulsus paradoxus
in cardiogenic shock, 1750
in dyspnea evaluation, 47e-3
echocardiography in, 271e-8f, 271e-8v
etiology of, 1445
measurement of, 1445
in obstructive lung disease, 1662
in pericardial tamponade, 1450
Pulsus parvus et tardus, 1445, 1531
Punch biopsy, 340
“Punch drunk” syndrome. *See* Chronic traumatic encephalopathy
Punding, 2615
Punta Toro virus, 1309t, 1318
Pupil(s)
assessment of, 195–197, 196f
in coma, 1774, 1774f
inequality in size of, 196
Pure alexia without agraphia, 177t, 178–179
Pure autonomic failure (PAF), 2642
Pure red cell aplasia, **667**
classification of, 668t
clinical associations of, 669
congenital (Diamond-Blackfan anemia), 669, 669t
definition of, 668
differential diagnosis of, 668
drug-induced, 44, 668t
etiology of, 668
parvovirus B19 infection and, 138e-6, 668, 668f, 1196
thymoma and, 123e-2, 668
treatment of, 668–669
Pure tone audiometry, 222
Pure white cell aplasia, 667
Pure word deafness, 177t, 178
Purified protein derivative (PPD)/tuberculin skin test (TST)
in fever of unknown origin, 138
in latent TB infection, 1114, 1121
limitations of, 770
in nontuberculous mycobacterial infections, 1131
in TB diagnosis, 1107
Purine metabolism, 431e-2, 431e-3
Purine metabolism disorders, **431e-5**
inborn, 431e-3t, 431e-5–6
in SCID, 375e-1, 2108
Purine nucleoside phosphorylase deficiency, 431e-3t, 2108
Purkinje fibers, 265e-12, 273e-1
Purple sticky (*Salvia divinorum*), 469e-4
Purpura, 127, 135, **367**
nonpalpable, 127, 367–368
palpable, 127, 364t, 368–369
posttransfusion, 138e-5
senile, 403–404
solar, 367
wet, 727, 728
Purpura fulminans
clinical features of, 133t
epidemiology of, 133t
etiology of, 780t, 782
genetic considerations in, 368, 737, 744
meningococcal, 998
treatment of, 738, 780t, 782
Purtscher's like retinopathy, 40e-4
Purupuru. *See* Pinta
Pus, 416
Push enteroscopy, 279, 1881
Push packages, 261e-9
Pustular lesions, 339t, **356**, 382
Pustular psoriasis. *See* Psoriasis, pustular
“Put Prevention into Practice” program, 30
Puumala virus, 1307t
PUVA photochemotherapy
action of, 391
adverse effects of, 348
for atopic dermatitis, 354t
for cutaneous T-cell lymphoma, 391
for morphea, 376
for psoriasis, 348, 354t, 391
for vitiligo, 358t, 391
PVCs. *See* Premature ventricular contractions (PVCs)
pVHL (elongin), 2343
PVL (Panton-Valentine leukocidin) toxin, 807, 830, 956
PVNS (pigmented villonodular synovitis), 2246, 2247f
P wave, 273e-1
Pyelonephritis. *See also* Urinary tract infections (UTIs)
actinomycotic, 1089
clinical features of, 863, 864f
definition of, 861
emphysematous, 863, 864f
M. hominis, 1165
in pregnancy, 49–50
prognosis of, 868
renal biopsy in, 62e-10f
risk factors for, 862
treatment of, 867
xanthogranulomatous, 863, 864f
Pyknodysostosis, 426e-4–5
Pyoderma. *See* Impetigo
Pyoderma gangrenosum
clinical features of, 76e-17f, 369, 369t
differential diagnosis of, 366
diseases associated with, 369
drug-induced, 379
in IBD, 1957
Pyoderma vegetans, in IBD, 1958
Pyogenic arthritis, pyoderma gangrenosum, and acne (PAPA) syndrome, 372e-8t, 422, 2215
Pyogenic granuloma, 237, 380, 380f
Pyomyositis, 831, 958, 2198
Pyostomatitis vegetans, in IBD, 1958
Pyramidal (corticospinal) system, 155f
Pyrantel pamoate, 246e-6t, 246e-12
Pyrazinamide
action of, 205e-4
adverse effects of, 205e-4–5, 1117
dosage of, 205e-4
genetic variations in response to, 39t
pharmacology of, 205e-4
in pregnancy, 205e-5
resistance to, 205e-5, 1118
for TB, 205e-2t, 205e-4–5, 1115–1117, 1115t
for tuberculous meningitis, 898
Pyrethrins, 2746
Pyridostigmine
adverse effects of, 2704
for gastroparesis, 260t, 261

- for Lambert-Eaton myasthenic syndrome, 2703
 for myasthenia gravis, 2704
 for orthostatic hypotension, 146, 2644
 Pyridoxal 5'-phosphate (PLP), 426e-5
 Pyridoxine (vitamin B₆), **96e-4**
 for acral erythema, 380
 for alcohol-induced acidosis, 64e-9, 319
 deficiency of. *See* Pyridoxine (vitamin B₆)
 deficiency
 dietary sources of, 96e-4
 drug interactions of, 205e-3
 functions of, 96e-2f, 96e-4
 for isoniazid toxicity, 473e-12t, 1116, 2690
 for nausea and vomiting in pregnancy, 261
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference range for, 2766t
 for sideroblastic anemia, 2531
 structure of, 96e-2f
 for TB prophylaxis in HIV infection, 1251t
 toxicity of, 96e-5, 463t, 2687-2688, 2687t
 for tuberculous meningitis, 899
 Pyridoxine (vitamin B₆) deficiency
 atherosclerosis in, 290e-2
 clinical features of, 96e-1t, 360
 dietary level associated with, 96e-1t, 96e-4
 neuropathy in, 2690
 Pyridoxine-dependent seizures, 434e-3t
 Pyrilamine, 473e-8t
 Pyrimethamine
 action of, 246e-12
 adverse effects of, 246e-4t, 246e-12, 647, 1380t, 1384
 for *C. belli* infections, 1407
 for CNS toxoplasmosis, 903
 drug interactions of, 246e-4t
 for malaria, 1378t, 1380t
 for PCP prophylaxis, 1251t, 1362t
 pharmacology of, 246e-12, 1380t
 in pregnancy and lactation, 246e-4t
 resistance to, 1384
 for toxoplasmosis, 1404
 for toxoplasmosis prophylaxis, 1251-1252t, 1404
 Pyrimidine-5'-nucleotidase (P5N) deficiency, 431e-5t, 431e-6f, 654t, 657
 Pyrimidine-5'-nucleotidase (P5N) superactivity, 431e-5t
 Pyrimidine metabolism
 inborn errors of, 431e-5t, 431e-6
 medication effects on, 431e-6
 schematic of, 431e-6f
 PYRIN domains, 422, 2212-2213
 Pyrin protein, 372e-8t, 2212
 Pyrogens, 124
 Pyroglutamic acid, 320
 Pyronaridine, 246e-6t, 246e-12
 Pyroptosis, 1234
 Pyrosis. *See* Heartburn
 Pyrroles, 205e-7
 Δ-Pyrroline-5-carboxylate synthase deficiency, 434e-3t
 Pyruvate dehydrogenase complex (PDC), 372e-24t, 2060
 Pyruvate dehydrogenase kinase (PDK), 102e-13
 Pyruvate kinase deficiency, 281, 433e-2t, 650f, 653-654, 654t
 6-Pyruvoyl-tetrahydropterin synthase (PTPS) deficiency, 434e-2t
 Pyuria, 294
 Q
 Q fever
 acute, 1155t, 1161
 chronic, 1162
 clinical features of, 137, 152e-3t, 1561
 diagnosis of, 1162
 epidemiology of, 1155t, 1161
 etiology of, 1155t, 1161
 fatigue syndrome following, 1161
 global considerations in, 1161
 incubation period for, 152e-3t
 prevention of, 152e-3t, 1162
 in returning war veterans, 152e-3t, 152e-8
 treatment of, 152e-3t, 1162
 Q fever vaccine, 1162
 QRS complex, 273e-1, 1451, 1452-1453, 1453f
 QRS voltage, low, 1457, 1459f
 QT interval
 prolonged, 41, 307, 313, 315, 1457, 1458f
 shortened, 1457
 variability, 273e-5
 Quadrantanopia, 198f
 Quadripareisis, 157, 157t
 Quadruplegic fever, 2660
 Quad screen, 83e-4
 Quality in health care, **12e-2**
 data about current state of, 12e-3
 factors relating to, 12e-3
 national state of measurement of, 12e-3-4
 pay-for-performance and, 12e-4
 public reporting of, 12e-4
 quality theory, 12e-2-3
 racial/ethnic disparities and, 16e-1
 strategies for improving, 12e-3
 "Quality of life" issues, 5
 Quality theory, 12e-2-3
 Quantitative sudomotor axon reflex test, 2640, 2641t
 Quarantivirus, 1310t
 Quarantaviruses, 1305, 1310t
 Quartan malarial nephropathy, 1375
 Queensland tick typhus, 1157
 Quetiapine
 adverse effects of, 2722t
 for bipolar disorder, 2718
 for delirium, 64, 64t
 for dementia, 175-176
 overdose/poisoning with, 473e-8t
 for psychosis in Huntington's disease, 2622
 for psychosis in PD, 2617
 for schizophrenia, 2721, 2722t
 Quinacrine
 action of, 246e-12
 adverse effects of, 246e-6t, 279, 379
 drug interactions of, 246e-1t, 246e-6t
 pharmacology of, 246e-12
 in pregnancy and lactation, 246e-6t
 for SLE, 2132
 Quincke's pulsations, of nailbed, 51e-5, 1535
 Quinidine
 action of, 273e-6t
 adverse effects of
 arthralgias, 2218t
 cardiac, 246e-13
 common/frequent, 246e-6t, 1379, 1380t
 cutaneous, 353
 fever, 138
 lupus syndrome, 2218t
 nonarrhythmic, 1480t
 occasional, 246e-6t
 proarrhythmic, 1379, 1480t
 rare, 1380t
 renal, 246e-13
 thrombocytopenia, 44, 727t
 for arrhythmias, 1479t, 1480t, 1498
 dosage of, 1479t, 1480t
 drug interactions of, 34t, 42, 42t, 43, 246e-6t
 azoles, 246e-3t
 mefloquine, 246e-10
 SSRIs, 2711t
 for malaria, 780t, 1378t, 1379, 1380t
 metabolism of, 34t
 pharmacology of, 246e-13, 1380t, 1479t, 1480t
 in pregnancy and lactation, 246e-6t
 therapeutic monitoring of, 2765t
 Quinine
 action of, 246e-13
 adverse effects of
 common/frequent, 246e-6t, 1380t
 occasional, 246e-6t
 oral pigmentation, 240t
 rare, 1380t
 renal, 246e-13
 thrombocytopenia, 44, 727, 727t, 729
 thrombotic thrombocytopenic purpura, 1864
 for *Babesia* infection, 780t, 1386-1387, 1386t
 drug interactions of, 246e-6t, 246e-10, 246e-13
 for malaria, 780t, 1380t
 overdose/poisoning with, 473e-13t
 pharmacology of, 246e-13
 in pregnancy and lactation, 246e-6t
 Quinolones
 action of, 931t, 932, 932f
 adverse effects of, 2218t
 overdose/poisoning with, 473e-13t
 resistance to, 931t, 934, 962, 1004
 for travelers' diarrhea, 796-797
 Quinsy. *See* Peritonsillar abscess
 Quintan fever. *See* Trench fever
 3-Quinuclidinyl benzilate, 262e-1t
 Quinupristin-dalfopristin
 action of, 931-932, 931t, 944, 961
 adverse effects of, 938t, 944, 976
 drug interactions of, 939t, 944
 for enterococcal infections, 975t, 976
 indications for, 941t, 944
 metabolism of, 935
 for MRSA infections, 961, 962t
 in pregnancy and lactation, 937t
 resistance to, 931t, 934, 941t
 Quorum sensing, 1062
 Q wave
 characteristics of, 1451
 in myocardial infarction, 1456
 anterior, 269e-4f, 269e-7f, 1456f
 anteroseptal, 269e-5f
 inferior, 269e-8f, 1456f
 variability, 1457f, 1600
 R
 R0 (reproduction number), 778
 R5 viruses, 1235
 R5X4 viruses, 1235
 R207910 (bedaquiline), 205e-7
 RA. *See* Rheumatoid arthritis (RA)
 RAA system. *See* Renin-angiotensin-aldosterone (RAA) system
 Rabeprazole
 adverse effects of, 1921
 for GERD, 263
 for peptic ulcer disease, 1920, 1920t
 for ZES, 1929
 Rabies, **1299**
 clinical features of, 896, 1301-1302, 1302f, 1302t
 diagnosis of, 152e-2t, 896, 1302, 1302f
 differential diagnosis of, 896, 1303
 epidemiology of, 1300, 1300f
 global considerations in, 1300
 incubation period for, 152e-2t
 pathogenesis of, 145e-1, 1300-1301, 1301f
 postexposure prophylaxis for, 167e-3, 167e-3t, 1303-1304, 1303f
 preexposure vaccination for, 1304
 prevention of, 152e-2t
 prognosis of, 1303
 in returning war veterans, 152e-2t, 152e-7
 treatment of, 1303
 Rabies immune globulin, 167e-3, 1303
 Rabies vaccine
 after bite-wounds, 167e-3
 Guillain-Barré syndrome and, 2695
 postexposure, 1304
 preexposure, 1304
 for travelers, 795t, 796, 1304
 Rabies virus, 1299
 Raccoon pinworm. *See* *Baylisascaris procyonis* infections
 Raccoon sign, 457e-2, 721, 721f
 "Rachitic rosary," 2466
 Racial/ethnic disparities in health care, **16e-1**
 clinical decision-making and, 16e-4
 health system interventions for, 16e-5
 implications for clinical practice, 16e-6-7
 mistrust and, 16e-4-5, 16e-6-7
 nature and extent of, 16e-1-2
 patient interventions for, 16e-6
 provider interventions for, 16e-5-6
 provider-patient communication and, 16e-3-4, 16e-4f

- Racial/ethnic disparities in health care (*Cont.*):
 recommendations to address, 16e-5-6
 root causes for, 16e-2-5
- Racism, 16e-4
- Rad, 103e-4, 263e-1
- Radial artery, for PCI access, 296e-1
- Radial neuropathy, 463e-4, 463e-5f, 2691
- Radiation, 263e-1. *See also* Ionizing radiation
- Radiation dose, 103e-4
- Radiation enteritis, 1940t
- Radiation esophagitis, 1910
- Radiation sensitizers, 103e-4
- Radiation syndrome, acute, 263e-2-4, 263e-4f
- Radiation terrorism, **263e-1**
 acute radiation syndrome, 263e-2-4, 263e-4f
 follow-up of exposed patient, 263e-7
 hospital management of, 263e-5-6, 263e-5f
 initial management of, 263e-4-5, 263e-4f, 263e-5f
 medical assay of radiation-exposed patient, 263e-7
 nuclear weapons, 263e-2
 radiologic dispersal events, 263e-2
 radionuclide properties and treatment, 263e-3t
 types of exposure, 263e-1-2
- Radiation therapy, **103e-4**
 for acromegaly, 2271
 adverse effects of, 103e-4
 bowel complications, 270, 583
 breast cancer, 103e-4, 524
 cardiovascular, 621, 1567, 1575
 cataract, 204
 cobalamin malabsorption, 646
 dysphagia, 256
 gastrointestinal, 1957
 head and neck, 505
 hepatic, 622
 in HIV infection, 1270
 hyperbaric oxygen therapy for, 477e-3, 477e-4t
 hypopituitarism, 2256, 2263-2264
 late, 620t, 621-623
 lymphedema, 1654
 myelofibrosis, 672
 nephropathy, 1864
 neurologic, 606-607, 622
 ocular, 623
 oral, 623
 pericarditis, 621, 1575, 1788
 pituitary, 2324t
 plexopathy, 2694
 pneumonitis, 489
 proctopathy, 346e-1v
 pulmonary, 621, 1797
 pulmonary eosinophilia, 1687
 soft tissue sarcoma, 119e-1
 testicular dysfunction, 2365
 thyroid cancer, 2304
 vascular, 103e-4
 for anal cancer, 544
 biologic effects of, 103e-4
 for bladder cancer, 576
 for bone pain, 61
 brachytherapy. *See* Brachytherapy
 for brain metastases, 605
 for breast cancer, 527-528
 for cholangiocarcinoma, 553
 for colorectal cancer, 543
 cranial, prophylactic in lung cancer, 522
 curative, 103e-4
 dosage of, 103e-4
 for ductal carcinoma in situ, 531
 for epidural metastases, 606
 for esophageal cancer, 533
 external beam. *See* External beam radiation therapy
 for gastric cancer, 535
 for head and neck cancer, 505
 for hepatocellular carcinoma, 551
 for Hodgkin's lymphoma, 709
 hyperbaric oxygen therapy for enhancement of, 477e-4t
 for lung cancer, 517, 518-519, 522
- modalities of, 103e-4
 physical properties of, 103e-4
 for pituitary tumors, 2263
 in pregnancy, 124e-1-2, 124e-1t
 for primary central nervous system lymphoma, 602
 for prostate cancer, 583
 quantitation of, 103e-4
 second malignancy and, 103e-4, 623, 709, 2264
 selective internal, 574
 for skin cancer, 501-502
 for soft tissue sarcoma, 119e-2
 systemic, 103e-22-23
 for testicular cancer, 590
- Radiculopathy, 111, 2692, 2692t
- Radiocontrast media. *See* Contrast agents
- Radiofrequency ablation
 for cardiac arrhythmias, 103e-6, 103e-6f
 characteristics of, 103e-5
 for chronic venous disease, 1652
 for hepatocellular carcinoma, 549
 for metastatic lesions, 103e-5
 for NETs, 574
- Radiofrequency denervation, 120, 1625
- Radiofrequency thermal rhizotomy, 2646
- Radioiodine therapy
 for Graves' disease, 2296-2297
 for hyperfunctioning solitary thyroid nodule, 2303
 for hyperthyroidism, 48
 for nontoxic multinodular goiter, 2302
 safety precautions for, 2297
 for thyroid cancer, 103e-22, 2306-2307
 for toxic multinodular goiter, 2302
- Radioiodine uptake, in thyroid evaluation, 2288
- Radionuclide contamination, 263e-6-7, 263e-6t
- Radionuclide scans/scintigraphy
 for colonic transit measurement, 273
 in gallbladder disease, 2078-2079, 2079t
 in gastrointestinal bleeding, 279
 in musculoskeletal disease evaluation, 2224, 2224t
 in NETs, 572, 572f
- Radium-233 chloride (alharadin), 119e-4, 586, 587
- Radium exposure, 263e-3t
- Radon exposure, 506, 1693
- Raf kinase inhibitors, 102e-4f, 102e-17
- Ragged red fibers, 462e-15
- Rai staging system, for lymphoid malignancies, 700t
- Rales, 1504, 1662
- Raloxifene
 for breast cancer prevention, 478, 531
 for gynecomastia, 2367
 for osteoporosis prevention/management, 2383, 2499
- Raltegravir
 action of, 1281
 adverse effects of, 1275t
 for HIV infection, 214e-9, 1275t, 1281
 resistance to, 1280f, 1281
 structure of, 1276f
- Ramipril, 1624t
- Ramsay Hunt syndrome, 149, 237, 1185, 2648
- Ramucirumab, 535
- Randall's plaque, 1866
- RAND Health Insurance Experiment, 15e-2
- Random Forests analysis/machine learning, 86e-1t, 86e-4
- Range of motion, 2219t, 2220
- Ranibizumab, 205
- Ranitidine
 for carcinoid syndrome, 566
 for gastrinoma, 2337
 for GERD, 263
 for peptic ulcer disease, 1920, 1920t
- Ranitidine bismuth, 1922, 1922t
- RANK (receptor activator of NF κ B)
 in bone metastasis, 102e-10-11
 in bone remodeling, 2490, 2492f
 in osteoclast and osteoblast differentiation, 426e-1f
- in Paget's disease of bone, 426e-1
 in rheumatoid arthritis, 2142f
- RANKL (RANK ligand), 102e-10
 in bone remodeling, 2455, 2490, 2492f
 in rheumatoid arthritis, 2142f, 2143
- Ranolazine, 1590
- RANTES, 372e-12t
- Rapamycin
 action of, 102e-4f, 465e-1
 for insulinoma, 569
 for Kaposi's sarcoma, 925
 for lifespan enhancement, 85, 94e-7
 for neurodegenerative diseases, 444e-9
 structure of, 94e-6f
- Rapeseed oil, 2218t
- Rapid acetylators, 40
- Rapid cycling, in bipolar disorder, 2717
- Rapid detection tests
 for group A streptococcal infections, 232
 for parasites, 245e-3
- Rapid reagent (RPR) test, 1137, 1137f
- RARA gene, 679
- Rarefaction, 86e-1t
- RAS (reticular activating system), 1771
- Rasagiline, 2615, 2616t
- Rasburicase
 adverse effects of, 1795
 genetic variations in response to, 39t, 40
 for tumor lysis syndrome treatment/prophylaxis, 431e-5, 1795
 for uric acid nephropathy, 681-682
- RAS gene mutations
 cancers associated with, 101e-2
 in hypercalcemia of malignancy, 609
 in juvenile myelomonocytic leukemia, 135e-6
 in melanoma, 496
 in thyroid cancer, 2304, 2305t
- Rash. *See also* Skin disorders/lesions; *specific diseases*
 centrally distributed maculopapular eruptions, 127, 128-130t, 134
 confluent desquamative erythemas, 130-131t, 134
 drug-induced, 25e-2f, 103e-26, 127, 380-381, 381f. *See also* Drug-induced illness, cutaneous
 with fever, **25e-1**, **127**, 128-134t, 369-370, 998
 in glucagonoma, 569
 morbilliform, 340t, 362, 362t
 nodular eruptions, 132-133t, 135
 peripheral eruptions, 130t, 134
 petechial, 999t, 1155, 1156f
 purpuric eruptions, 133t, 135, 999t
 pustular eruptions, 131-132t, 134-135
 scarlatiniform, 362-363, 362t
 ulcers and eschars, 133-134t, 135
 urticaria-like eruptions, 132t, 135
 vesiculobullous eruptions, 131-132t, 134-135
- Rasmussen's aneurysm, 245, 1108
- Rasmussen's encephalitis, 372e-24t
- Raspberry tongue, 241t
- Ras protooncogene, 102e-4f
- Rat-bite fever (Haverhill fever), 130t, 167e-1-2
- Rat-bite fever (sodoku), 129t, 167e-1-2
- Rat bite-wound, 167e-1-2
- Rat flea, 2750
- Rathke's cysts, 2264
- Rationing, 12e-3
- Rat lungworm. *See* *Angiostrongylus* spp. infections
- Rattlesnake bite, 2734, 2734f, 2736t, 2738. *See also* Snakebites
- Ravn virus disease, 1323, 1325f, 1326f
- R_{aw}. *See* Airway resistance (R_{aw})
- Raxibacumab, 261e-3
- Raynaud's phenomenon, **1647**
 chilblain in, 478e-4
 clinical features of, 1647-1649, 1648f
 cyanosis in, 249
 drug-induced, 590, 623, 1649
 in inflammatory myopathies, 2196
 localized hypoxia in, 248
 pathophysiology of, 372e-24t, 1648

- primary (Raynaud's disease), 47e-3, 1648-1649
 secondary causes of, 1648t, 1649
 in Sjögren's syndrome, 2167, 2167t
 in systemic sclerosis, 376, 1649, 2155, 2155t, 2159, 2160f
 treatment of, 1649, 2164, 2168f
- Rb. *See* Retinoblastoma protein (Rb)
- Rb gene mutations
 in bladder cancer, 576
 cancers associated with, 101e-6t
 in lung cancer, 507, 509
 in parathyroid carcinoma, 2472
 in retinoblastoma, 440
 in soft tissue sarcoma, 119e-1
- R-binder protein, 350e-1
- RCC. *See* Renal cell carcinoma (RCC)
- RCRI (Revised Cardiac Risk Index), 52, 53f, 53t
- RDW (red blood cell distribution width), 81e-1, 393
- Reactive arthritis (Reiter's syndrome), 2173
 after nontyphoidal *Salmonella* infections, 1054
 after *Shigella* infections, 1057
 after *Yersinia* infections, 1077
C. trachomatis, 1168, 1170t
 clinical features of, 837, 2174
 aortic aneurysm, 1638
 aortitis, 290e-3, 1638
 cutaneous, 353, 353t
 diarrhea, 266-267
 oral, 237, 2174
 diagnosis of, 2175
 epidemiology of, 2173-2174
 etiology of, 2174
 vs. gonococcal arthritis, 1006, 1007f, 2175
 historic background of, 2173
 in HIV infection, 1261
 HLA gene association with, 373e-7, 373e-8t
 vs. Lyme arthritis, 1152
 pathogenesis of, 2174
 pathology of, 2174
 vs. relapsing polychondritis, 2204
 treatment of, 1078, 2175
- Reactive hyperemia, 1648
- Reactive lymphocytes, 81e-2
- Reactive oxygen species (ROS)
 in aging, 85e-8
 damage to mitochondrial DNA by, 85e-8-9, 85e-8f
 in endothelium, 265e-2
 generation of, 386
 in hyperbaric oxygen therapy, 477e-1, 477e-2f
 skin damage caused by, 386
- Reactive skin decontamination lotion (RSDL), 262e-6
- "Readback," 12e-2
- Reading, assessment of, 177
- Rebound tenderness, 106
- Rebuck skin window test, 423
- Receiver operating characteristic (ROC) curve, 21, 22f
- Receptor tyrosine kinase, 102e-3, 102e-4f, 400e-1t
- Receptor tyrosine kinase receptors, 102e-9f
- Recessive disorders, 438-439, 438f
- Reciprocal translocation, 83e-5, 83e-5f
- RECIST (Response Evaluation Criteria in Solid Tumors), 103e-5, 472, 552
- Recluse spider bite, 2747
- Recombinant tissue plasminogen activator (rtPA). *See* Alteplase (rtPA)
- Recombination, genetic, 428, 429f
- Recommended dietary allowance, 95e-1
- Recoverin, 372e-25t, 614t
- Rectal bleeding
 in colorectal cancer. *See* Colorectal cancer
 in hemorrhoidal disease. *See* Hemorrhoidal disease
 in rectal prolapse, 1974
 in ulcerative colitis, 1952, 1952t
- Rectal cancer, 543. *See also* Colorectal cancer
- Rectal neuroendocrine tumors (carcinoids), 560t, 563, 564, 567. *See also* Gastrointestinal neuroendocrine tumors (GI-NETs)
- Rectal prolapse, 1973
 clinical features of, 1974, 1974f
 epidemiology of, 1973-1974
 pathophysiology of, 1974
 treatment of, 1974-1975, 1975f
- Rectal snip, 245e-3-4
- Rectal temperatures, 124
- Rectoanal (anorectal) angle, 265, 265f
- Rectocele, 274
- Rectopexy, 1974-1975, 1975f
- Rectus sheath, hematoma of, 105
- Red blood cell(s)
 agglutination of, 81e-1, 81e-3f
 in anemia, 392-393. *See also* Anemia
 antigens and antibodies, 138e-1-2, 138e-1t
 complement activation and, 661f
 excess of. *See* Polycythemia
 folate in, 648
 fragmented, 81e-4f
 glycolysis in, 650, 650f
 hematopoietic differentiation of, 89e-3f
 immune destruction of, 658, 659f
 iron content in, 626
 life cycle of, 626, 650
 mechanical destruction of, 657, 657t
 membrane-cytoskeleton complex of, 651, 651f
 nucleated, 81e-5f
 osmotic fragility of, 651
 in peripheral blood smear, 81e-1, 81e-2-7f
 production of, 392, 392f
 protoporphyrin levels in, 627f, 628
 redox metabolism in, 654f
 transfusions. *See* Transfusion(s)
 urinary casts, 62e-11f
- Red blood cell distribution width (RDW), 81e-1, 393
- Red blood cell indices, 393, 394t
- Red blood cell scintigraphy, 279
- Red blood cell survival study, 650
- Red cell anion exchanger (AE1), 309
- Red clover, 404t
- Red hair color (RHC) phenotype, 494
- Red hepatization, 804
- Red light, for basal cell carcinoma, 386
- Red man syndrome, 381, 938t, 942
- Red pulp, of spleen, 409, 410f
- "Red tide," 2743
- 5 α -Reductase, 332
- 5 α -Reductase inhibitors, 478, 587
- 5 α -Reductase type 2 deficiency, 2353
- Reductionism, 87e-1, 87e-3
- Reduviid bugs, 1560, 2750
- REE (resting energy expenditure), 98e-4
- Reed-Sternberg cells, 81e-11f, 709f
- Reentry, 273e-4, 273e-4f
- Refeeding edema, 253
- Refeeding syndrome, 460
- Reference values. *See* Clinical laboratory tests, reference values for
- Referred pain, 88
 in abdominal diseases, 104t, 105
 to back, 111, 118
 convergence-projection hypothesis of, 89f
- Reflex(es), testing of, 2538-2539. *See also specific reflexes*
- Reflexology, 14e-1t
- Reflex sympathetic dystrophy. *See* Complex regional pain syndrome (CRPS)
- Reflex syncope. *See* Syncope, neurally mediated
- Reflux nephropathy, 1859-1860, 1860f
- Refractive state, 195
- Refractory anemias, 669, 669t. *See also* Myelodysplasia
- Refractory cytopenia with multilineage dysplasia, 669t
- Refractory neutropenia, 669t
- Refractory thrombocytopenia, 669t
- Refsum's disease, 206, 451e-4t, 2681
- Refusal of care, 17e-2, 66
- Regenerative medicine, 453
- Registries, 25
- Regorafenib
 action and targets of, 102e-5t, 103e-18t
 adverse effects of, 103e-18t
 for colorectal cancer, 103e-19
 for gastrointestinal stromal tumor, 536
- Regulatory failure, 15e-1
- Regurgitation
 in achalasia, 1904
 definition of, 258
 in esophageal disease, 1901
 in GERD, 1907
 nasal, 256
- Rehabilitation
 in alcohol use disorder, 2728
 in ischemic stroke management, 2563
 in opiate use disorders, 468e-3-4
 pulmonary, 47e-4, 1706
 in spinal cord disorders, 2660-2661, 2660t
- Reifenstein's syndrome (partial androgen insensitivity syndrome), 2355
- Reiter's syndrome. *See* Reactive arthritis (Reiter's syndrome)
- Rejection
 HLA in, 373e-5-6
 inflammatory mediators in, 372e-29t
 in kidney transplantation, 1827, 1829
 in liver transplantation, 366e-3f, 2071-2072, 2073
 in lung transplantation, 320e-3
 treatment of, 372e-30, 372e-31
- Relapsing fever
 clinical features of, 126, 152e-5t, 167e-1-2, 1147
 diagnosis of, 152e-5t, 1147-1148, 1148f
 epidemiology of, 1146
 etiology of, 1146, 1146t
 global considerations in, 1146
 incubation period for, 152e-5t
 louse-borne, 1146
 pathogenesis of, 1146-1147
 prevention of, 1149
 prognosis of, 1148-1149
 in returning war veterans, 152e-5t
 treatment of, 152e-5t, 1148, 1148f
- Relapsing polychondritis, 2202
 aortic aneurysm in, 1638
 aortitis in, 1638
 clinical features of, 1443, 2193, 2203-2204, 2203f, 2203t
 diagnosis of, 2204
 differential diagnosis of, 2204
 disorders associated with, 2202, 2202t
 incidence of, 2202
 pathogenesis of, 2202-2203
 pathology of, 2202
 prognosis of, 2204
 treatment of, 2204
- Relative afferent pupil defect, 196, 196f
- Relative bradycardia (Faget's sign), 763, 763t
- Relative risk, 373e-7
- Relaxin, 2378
- REL gene mutations, 101e-2t, 101e-3t
- Reminder systems, 30
- REM sleep, 184-185, 185f
- REM sleep behavior disorder, 192-193, 194v, 214-215, 2618
- Renal abscess, 851
- Renal agenesis/dysgenesis, 48, 333e-1
- Renal amyloidosis, 62e-6f, 1845
- Renal artery stenosis, 1627
 azotemia in, 292
 diagnosis of, 1618, 1628, 1628t
 hyperaldosteronism and, 306
 incidence of, 1627
 pathophysiology of, 1627-1628
 treatment of, 1618, 1628-1629, 1629t
- Renal artery thrombosis, 292
- Renal biopsy
 in acute interstitial nephritis, 62e-10f
 in acute kidney injury, 62e-10f, 1808
 in acute nephritic syndromes, 62e-1-6f
 in Alport's syndrome, 62e-8f, 1847
 in antiglomerular basement membrane disease, 62e-6f, 1839

- Renal biopsy (*Cont.*):
 assessment of, 1836–1837
 atlas of, 62e-1–10
 in chronic kidney disease, 1820
 in diabetic nephropathy, 62e-8f, 1844
 in Fabry's disease, 62e-7f, 1846
 in focal segmental glomerulosclerosis, 62e-1f, 62e-2f
 in glomerular diseases, 62e-1–6f, 1836–1837
 in granulomatosis with polyangiitis, 62e-5f, 386e-1, 386e-6f, 1840
 in hemolytic-uremic syndrome, 62e-9f, 1848
 in hypertensive nephrosclerosis, 62e-9f, 1848
 in IgA nephropathy, 62e-3f, 1839
 in lupus nephritis, 62e-5f, 1838
 in membranoproliferative glomerulonephritis, 62e-3f, 62e-4f, 1841
 in membranous glomerulonephritis, 62e-3f, 1843
 in minimal change disease, 62e-1f, 1841–1842
 in multiple myeloma, 62e-7f, 1859f
 in oxalosis, 62e-10f
 in poststreptococcal glomerulonephritis, 62e-2f, 1837
 in pyelonephritis, 62e-10f
 stains for, 1836
 in vasculitic syndromes, 386e-1
- Renal cell carcinoma (RCC), 578
 clinical features of, 578
 epidemiology of, 578
 epigenetic silencing in, 102e-7
 genetic considerations in, 578, 578t
 intrahepatic cholestasis in, 284
 metastatic, 604t
 paraneoplastic syndromes in, 578, 612, 612t
 pathology of, 578, 578t
 prognosis of, 578, 579, 579f
 staging of, 578, 579f
 treatment of, 102e-5t, 102e-17, 103e-19, 103e-22, 578–579
 tuberous sclerosis and, 578, 1854
 von Hippel-Lindau syndrome and, 578, 1854, 2343
- Renal colic, 1866, 1868, 1872. *See also* Nephrolithiasis
- Renal coloboma syndrome, 1855
- Renal cysts. *See* Polycystic kidney disease
- Renal disease/failure
 acute. *See* Acute kidney injury (AKI)
 chronic. *See* Chronic kidney disease (CKD)
 glomerular. *See* Glomerular diseases
 nephrolithiasis. *See* Nephrolithiasis
 tubulointerstitial diseases. *See* Tubulointerstitial diseases
 vascular. *See* Renovascular disease
- Renal natriuretic peptide, 332e-9
- Renal osteodystrophy, 291, 2478
- Renal pelvis, carcinoma of. *See* Kidney cancer; Renal cell carcinoma (RCC)
- Renal stem cells, 453t
- Renal transplantation. *See* Kidney transplantation
- Renal tubular acidosis (RTA)
 distal (type I)
 genetic considerations in, 332e-7t
 hypokalemia in, 307
 metabolic acidosis in, 320
 in Sjögren's syndrome, 64e-6
 urinary tract obstruction and, 1872
 proximal (type II)
 genetic considerations in, 332e-7t
 metabolic acidosis in, 320
 type IV, 320, 2425
- Renal tubular defects, 289t
- Renal vein, renin ratio, 1618
- Renal vein thrombosis, 1866
- Renin
 ectopic production of, 609t
 pNET secreting, 559t, 571
 renal vein renin ratio, 1618
 in shock, 1745
 synthesis and secretion of, 1614
- Renin-angiotensin-aldosterone (RAA) system
 assessment of, 2311
 in blood pressure regulation, 1613–1614, 1614f
 in edema, 250, 252
 in glomerular filtration rate regulation, 332e-2–3, 332e-2f
 in heart failure pathogenesis, 1502, 1502f
 in mineralocorticoid synthesis, 2311, 2311f
 in potassium regulation, 304–305, 310
 in sodium regulation, 332e-10–11
- Renovascular disease, 1627
 atheroembolic, 1629, 1806t
 diagnosis of, 1628, 1628f, 1628t
 hypertension, 1614, 1617–1618, 1622, 1630
 hypertensive nephrosclerosis, 62e-9f, 1617, 1630, 1835t, 1847–1848
 renal artery stenosis. *See* Renal artery stenosis
 thromboembolic, 1629–1630, 1630f, 1866
 thrombotic microangiopathic. *See* Thrombotic microangiopathy, renal
- Reoviruses
 gene expression and replication in, 214e-4
 reservoirs and vectors of, 1305
 structure of, 214e-2t, 214e-3f, 1295
- Reovirus infections, 1311–1312t, 1319
- Repaglinide, 2413, 2414t
- Repetition, assessment of, 177
- Representativeness heuristic, 19
- Reproduction number (R0), 778
- Reproductive system disorders. *See* Female reproductive system disorders; Male reproductive system disorders
- Reptilase time, 407
- Reptile bites. *See* Snakebites
- Research
 ethical issues in, 17e-4
 physician engaged in, 7
 skews in, global effects of, 13e-4
- Reserpine, 1624t
- Resident physicians, working hours of, 7
- Resident stem cells, 453
- Residual volume (RV), 306e-1f, 306e-2, 306e-2f, 306e-5f
- Resilience, 86e-1t, 86e-3, 86e-7
- Resistance strength training, 165
- Resistance to thyroid hormone (RTH), 435, 2286t, 2287
- Respiration
 in hypothermia, 478e-2t
 respiratory rate, 47e-3
 in shock, 1730t, 1746
- Respiratory acidosis, 322
 acid-base nomogram, 316f
 clinical features of, 322
 compensatory responses in, 316t, 322
 drug-induced, 322
 etiology of, 322–323, 323t
 hyperkalemia in, 309
 in mixed acid-base disorders, 316t, 317
 treatment of, 323
- Respiratory alkalosis, 323
 acid-base nomogram, 316f
 compensatory responses in, 316t
 diagnosis of, 324
 etiology of, 323–324, 323t
 in hypoxia, 248
 in mixed acid-base disorders, 316t, 317
 in sepsis/septic shock, 1756
 treatment of, 324
- Respiratory bronchiolitis-associated ILD, 1713
- Respiratory disorders. *See also* Lung disease; Respiratory tract infections
 approach to the patient, 1661
 categories of, 1661, 1661t
 chest pain in, 1661
 clinical features of, 1661–1662
 clubbing in, 1662
 cough in, 1661
 deaths from, by sex, 6e-1t
 diagnosis of, 1663
 bronchoscopy, 1663, 1667, 1668
 chest radiography, 1663, 1664f
 CT, 1663, 1664–1666, 1664f, 1665f
 emerging techniques, 1668
 endobronchial ultrasound, 1668
 flexible fiberoptic bronchoscopy, 1667
 gas exchange measurement, 306e-6
 mediastinoscopy, 1668
 mediastinotomy, 1668
 MRI/MRA, 1666, 1667f
 nuclear medicine techniques, 1664
 percutaneous needle aspiration, 1667
 PET, 1667
 pulmonary angiography, 1667–1668
 pulmonary function testing, 1662–1663
 sputum analysis. *See* Sputum analysis
 thoracentesis, 1667
 thoracoscopy, 1668
 thoracotomy, 1669
 transbronchial needle aspiration, 1669
 ultrasound, 1663–1664
 ventilatory function measurement, 306e-6
 video-assisted thoracic surgery, 1668–1669
 virtual bronchoscopy, 1666, 1666f
- dyspnea in, 47e-1t, 47e-2, 1661
 hemoptysis in, 1661
 history in, 1661–1662
 inhaled agents associated with, 1662
 obesity and, 415e-6
 obstructive patterns, 306e-5–6
 pathophysiology of, 306e-1
 physical examination in, 1662
 pulmonary function values in, 306e-5–6, 306e-5f
 restrictive patterns, 306e-5–6, 306e-5f
 risk factors for, 1662
 smoking and, 1662, 2729, 2730t
 in women, 6e-2f
- Respiratory failure
 in ARDS. *See* Acute respiratory distress syndrome (ARDS)
 in critically ill patient, 1731–1732
 hypercarbic, 1740
 in hyponatremia, 300
 hypoxemic, 1740
 hypoxia in, 247
 pathophysiology of, 1731–1732
 perioperative, 1732
 in shock, 1731
- Respiratory function
 disturbances of. *See* Respiratory disorders
 flow-related properties of, 306e-2, 306e-2f, 306e-3f, 1733, 1734f
 gas exchange, 306e-3–4
 during sleep, 187
 ventilation-perfusion heterogeneity, 306e-4, 306e-4f
 ventilatory adequacy, 306e-3
 volume-related properties of, 306e-1–2, 306e-1f, 1733
 work of breathing, 306e-2–3
 “Respiratory hygiene and cough etiquette,” 917
- Respiratory muscles
 assessment of strength of, 306e-6
 fatigue/weakness of, 306e-5, 1731–1732
 hypoperfusion of, 1732
- Respiratory quotient, 306e-3
- Respiratory sinus arrhythmia, 278e-1f
- Respiratory syncytial virus infections. *See* Human respiratory syncytial virus (HRSV) infections
- Respiratory tract infections
 adenovirus. *See* Adenovirus infections
Aspergillus. *See* *Aspergillus* infections, respiratory tract
 asthma exacerbations and, 1671
C. pneumoniae, 1173–1174
 chronic *P. aeruginosa*, 1045–1046
 COPD exacerbations and. *See* Chronic obstructive pulmonary disease (COPD), exacerbations of
 coronavirus. *See* Coronavirus infections
 in cystic fibrosis, 1697
 enterovirus, 965t
 epiglottitis. *See* Epiglottitis
 hemoptysis in, 245
 HMPV, 965t, 1206

- HRSV. *See* Human respiratory syncytial virus (HRSV) infections
- HSV, 965t
- influenza virus. *See* Influenza
- laryngitis, 234
- lymphadenopathy in, 408
- M. pneumoniae*, 1163, 1164t
- mastoiditis, 230–231, 231f, 900
- in mitral stenosis, 1540
- mucormycosis, 1351
- Nocardia*, 1084–1085, 1085f, 1087t
- nonspecific, 225–226
- nontuberculous mycobacterial, 1130
- oral cavity, 233–234
- parainfluenza virus. *See* Parainfluenza virus infections
- pharyngitis. *See* Pharyngitis
- pneumonia. *See* Pneumonia
- rhinovirus. *See* Rhinovirus infections
- sinusitis. *See* Sinusitis
- sporotrichosis, 1354, 1354t
- Respite care, 65
- Response, to drug, 103e-5
- Response Evaluation Criteria in Solid Tumors (RECIST), 103e-5, 472, 552
- REST, 444e-4
- Restan virus, 1309t
- Restenosis, 296e-1–3, 296e-2f, 1591
- Resting energy expenditure (REE), 95e-1, 98e-4
- Resting (basal) metabolic rate, 73, 74f, 415e-2, 461
- Restitution, 1912
- Restless legs syndrome
- in chronic kidney disease, 1819
 - clinical features of, 188t, 192, 2625
 - diagnosis of, 188t, 192
 - genetic considerations in, 2625
 - incidence of, 2625
 - in pregnancy, 49
 - treatment of, 188t, 192, 2625
- Restriction element, 373e-4
- Restrictive cardiomyopathy
- in amyloidosis, 1566–1567, 1566f, 1567f
 - clinical features of, 1554t, 1573t
 - diagnosis of, 1554t, 1566f, 1573t
 - differential diagnosis of, 1462, 1463t, 1573t, 1576
 - drug-induced, 1567
 - endomyocardial, 1567–1568
 - etiology of, 1565t, 1566
 - fibrotic, 1567
 - genetic considerations in, 1555t
 - in metabolic disorders, 1565–1566
 - overlap with other cardiomyopathies, 1565t
 - pathogenesis of, 1566
 - radiation-related, 1567
 - in systemic sclerosis, 1567
- Resuscitation, ABCs of, 1750
- Resveratrol, 85, 94e-6–7, 94e-6f
- Reptilase
- action of, 759
 - for acute limb ischemia, 1646
 - contraindications to, 1605
 - domain structure of, 759, 759f
 - indications for, 759
 - for myocardial infarction, 1605
- RET gene mutations
- action of, 2329
 - cancers associated with, 101e-3t
 - as drug target, 101e-8
 - genetic testing for, 2342
 - in lung cancer, 508, 509f, 521
 - in MEN 2, 101e-6t, 2251–2252, 2307, 2332, 2336t, 2342, 2472
 - in pheochromocytoma/paraganglioma, 2334f, 2335, 2342
 - prophylactic thyroidectomy in, 2341, 2341t
 - screening tests in, 2341t, 2342
 - in thyroid cancer, 2304, 2305t, 2341
- Reticular activating system (RAS), 1771
- Reticular dysgenesis, 375e-1, 2108
- Reticulate body, 1166, 1166f
- Reticulin stain, 81e-5f
- Reticulocyte(s), 396f
- Reticulocyte count, 81e-2f
- in anemia, 394, 396–397
 - correction of, 396f, 397
 - in hemolytic anemia, 649t, 650
 - reference values for, 2756t
- Reticulocyte production index, 396–397, 396f, 396t, 397t
- Reticuloendothelial system, 626
- Reticulohistiocytosis, multicentric, 365
- Reticulospinal pathway, 155f
- Reticulum cell sarcoma (interdigitating dendritic cell sarcoma), 135e-9
- Retina, 195
- age-related changes in, 40e-2
 - cotton-wool spots in, 40e-3, 40e-3f
 - crystal deposition in, 40e-4
 - detached, 197, 204, 204f
 - detachment of, 40e-5
 - emboli in, 40e-4, 41e-1v
 - examination of, 40e-1, 40e-2, 40e-2t
 - gyrate atrophy of, 434e-3t
 - hard exudates in, 40e-3
 - hemorrhages in, 40e-2–3
 - deep, 40e-3f
 - diseases associated with, 40e-3t
 - in high-altitude illness, 476e-1
 - in infective endocarditis, 200, 201f
 - in malaria, 1373, 1373f
 - superficial, 40e-2f
 - in hypertension, 1622
 - lesions of, 197
 - microaneurysms of, 40e-3
 - neovascularization of, 40e-3–4, 40e-4f
 - radiation-induced damage to, 103e-4
 - vascular sheathing in, 40e-4–5, 40e-4f5
- Retinal artery occlusion, 201, 201f
- Retinaldehyde, 96e-5
- Retinal pigment epithelium, 40e-1
- Retinal vein occlusion, 201, 202f
- Retinitis
- CMV
 - clinical features of, 1192, 1192f
 - in HIV infection, 1267
 - treatment of, 215e-7–8, 1193–1194
 - T. gondii*, 1402, 1403t
- Retinitis pigmentosa, 206, 206f, 431t, 1855
- Retinoblastoma
- hereditary (familial), 101e-2t, 101e-5f, 101e-6, 101e-6t, 440
 - sarcomas and, 119e-1
 - sporadic, 440
- Retinoblastoma protein (Rb), 102e-1
- Retinoblastoma (RB) tumor suppressor protein, 494
- Retinoic acid, 96e-5, 96e-6. *See also* All-*trans*-retinoic acid (ATRA)
- for acne vulgaris, 352
 - in pregnancy, 96e-7
 - for promyelocytic leukemia, 96e-6, 400e-4
 - for squamous cell carcinoma, 502
- Retinoic acid-inducible gene-1-like receptors (RLRs), 372e-2, 372e-4, 372e-4t
- Retinoic acid receptors, 96e-6, 444e-5, 477
- Retinoid acid X receptors (RXRs), 2287, 2287f
- Retinoids, 96e-5, 216, 348
- Retinol, 477
- Retinol-binding protein, 462
- Retinopathy
- cancer-associated (paraneoplastic), 614t, 619
 - CMV, 1192, 1192f
 - diabetic. *See* Diabetic retinopathy
 - diseases associated with, 40e-2–3, 40e-3t
 - hypertensive, 201, 201f, 1622
 - Purtscher's like, 40e-4
- Retrolbulbar tumor, of optic nerve, 206
- Retrocalcaneal bursitis, 2247
- Retrocollis, 2620
- Retrograde amnesia, 182
- Retrograde ejaculation, 325, 2427
- Retromer complex, 444e-10
- Retropertoneal fibrosis, 391e-1t, 391e-2
- Retropertoneal hemorrhage, 733
- Retropharyngeal abscess, 235
- Retropharyngeal soft tissue mass, 441e-32f
- Retropharyngeal space infections, 235
- Retrovirus(es), 225e-1
- cancer development and, 101e-2, 101e-7
 - classification of, 225e-1, 225e-1t
 - gene expression and replication in, 214e-5, 225e-1–2, 225e-2f, 225e-3f
 - for gene therapy delivery, 91e-1t, 214e-9, 225e-2
- HIV. *See* HIV
- HLTV-1. *See* Human T-cell lymphotropic virus (HTLV-1)
- inflammatory myopathies and, 2197
 - life cycle of, 225e-1, 225e-2f
 - myopathies associated with, 2657
 - structure of, 214e-2t, 214e-3f, 225e-1, 225e-3f
- Rett's syndrome, 431, 444e-1, 444e-4
- Reverse cholesterol transport, 2437–2438
- Reverse isolation, 492
- Reverse straight-leg raising sign, 113
- Reverse T₃ (rT₃), 2283f
- Reverse transcriptase enzyme, HIV, 1216, 1219
- Reverse transcriptase inhibitors. *See* Combination antiretroviral therapy (cART)
- Reverse transcriptase polymerase chain reaction (RT-PCR), 1247, 1247t
- Reverse transcription, 214e-5
- "Reverse type", 138e-2
- Reverse vaccinology, 776
- Reversible cerebral vasoconstriction syndrome, 2568
- Reversible posterior leukoencephalopathy, 1793, 2568, 2584
- rev* gene, 225e-3f, 1219f
- Revised Cardiac Risk Index (RCRI), 52, 53f, 53t
- Rev protein, 225e-2
- Rewarming
- for frostbite, 478e-4, 478e-4t
 - for hypothermia, 478e-3, 478e-3t
 - for shock, 1751
- Rex proteins, 214e-5, 214e-7, 225e-2
- Reye's syndrome, 126, 1184, 1185, 1212–1213
- Rh0(D) immune globulin, for immune thrombocytopenic purpura, 728
- Rhabdomyolysis
- acute kidney injury in, 1804, 1805, 1806t, 1810
 - hyperkalemia in, 310
 - hypernatremic, 303
 - hypocalcemia in, 315
 - hypokalemia and, 307
 - hypophosphatemia-induced, 2459–2460
 - in influenza, 1212
 - treatment of, 1810
- Rhabdomyoma, cardiac, 289e-2
- Rhabdomyosarcoma, 101e-3t, 119e-1, 119e-2. *See also* Soft tissue sarcoma
- Rhabdoviruses
- gene expression and replication in, 214e-4
 - lyssaviruses, 1299, 1304
 - structure of, 214e-2t, 214e-3f, 1305
- Rhabdovirus infections
- Chandipura virus infection, 1316
 - clinical syndromes of, 1312t
 - geographic distribution of, 1311–1312t
 - rabies. *See* Rabies
 - vesicular stomatitis virus infection, 1299, 1304
- RHAG gene mutations, 653, 653t
- Rh blood group system, 138e-1, 138e-1t
- RHC (red hair color) phenotype, 494
- Rheumatic fever, 2149
- cardiovascular complications of
 - antithrombotic prophylaxis for, 2566t
 - aortic valve disease, 1529
 - diagnosis of, 2151, 2151f, 2151t
 - echocardiography in, 2151, 2151f, 2151t, 2154v
 - global considerations in, 16, 964, 964f, 1501, 2149
 - heart failure, 2153
 - mitral stenosis, 51e-6, 1539, 2151, 2154v. *See also* Mitral stenosis
 - pathogenesis of, 2149–2150, 2150f
 - pericarditis, 1575
 - SA node dysfunction, 1467, 1467t

- Rheumatic fever (*Cont.*):
 clinical features of, 129t, 2150
 chorea, 2151–2152, 2153
 erythema marginatum, 129t, 134, 356, 2152
 fever, 2152
 joint involvement, 837, 2151
 nodules, 364, 2152
 diagnosis of, 2151t, 2152, 2152t, 2153t
 epidemiology of, 2149, 2150f
 etiology of, 129t
 genetic considerations in, 2150
 global considerations in, 1528–1529, 2149
 group A streptococcal infection and, 233, 2152
 immunopathology of, 372e–24t, 377e–1, 2150
 pathogenesis of, 2149–2150, 2150f
 prevention of, 2153–2154, 2154t
 prognosis of, 2153
 treatment of, 2152–2153
- Rheumatic heart disease. *See* Rheumatic fever, clinical features of
- Rheumatoid arthritis (RA), **2136**
 approach to the patient, 2147–2148
 clinical features of, 2136–2139
 cardiovascular, 290e–1t, 290e–3, 2138
 aortic aneurysm, 1638
 aortitis, 1638
 AV conduction block, 1471, 1471t
 SA node dysfunction, 1467
 constitutional, 2138
 cutaneous, 76e–16f, 364, 364t, 2138
 extraarticular, 2137–2138, 2137f
 Felty's syndrome, 372e–15t
 hematologic, 372e–15t, 393, 630, 630f
 inflammation/swelling, 2137f, 2142, 2142f
 lymphoma, 2138
 neck pain, 122
 neuropathy, 2683
 oral pain, 237
 pulmonary, 1713–1714, 2138
 Raynaud's phenomenon, 1649
 Sjögren's syndrome, 2138
 T-cell large granular lymphocytic leukemia, 2138
 vasculitis, 372e–15t, 2138, 2193
 conditions associated with, 2137f, 2138–2139
 diagnosis of, 41t, 2143–2144, 2144f
 disability in, 2144
 EBV and, 2140
 environmental factors in, 2140
 epidemiology of, 2139, 2139f
 genetic considerations in, 373e–8, 373e–8t, 2139–2140
 global considerations in, 2139, 2139f, 2149
 natural history of, 2144
 pathogenesis of, 372e–22, 372e–24–25t, 372e–29t, 2141–2143, 2142f
 pathology of, 2140–2141
 prognosis of, 2144
 vs. relapsing polychondritis, 2204
 remission in, 2148, 2148t
 research progress in, 2136
 treatment of, **2144**
 approach to, 2144–2145, 2147–2148
 assistive devices, 2148
 biologic agents, 372e–30, 377e–5, 2145, 2146–2147t, 2147
 DMARDs, 2145, 2146t
 in the elderly, 2148–2149
 global considerations in, 2149
 glucocorticoids, 2145
 NSAIDs, 2145
 physical therapy, 2148
 in pregnancy, 2148
 surgery, 2148
 tofacitinib, 2147, 2147t
- Rheumatoid vasculitis, 372e–15t, 2138
- Rheumatologic diseases. *See* Musculoskeletal disorders
- Rhinitis
 allergic. *See* Allergic rhinitis
 chronic atrophic, 1032
 perennial nonallergic, 2122
 vasomotor, 2122
- Rhinophyma, 352
- Rhinorrhea, CSF, 457e–2
- Rhin scleroma, 1032
- Rhinosinusitis. *See* Sinusitis
- Rhinovirus, 1202–1203
- Rhinovirus infections, 145e–4, 225–226, 923, 1202t, **1203**
- Rhinovirus vaccine, 1203
- Rhizobium radiobacter*, 183e–2t, 183e–3
- Rhizopus* spp., 1350, 1351f. *See also* Mucormycosis
- Rhodococcus* spp. infections, 924, 981, 1254
- Rho kinase, 324, 325f
- Rhonchi, 1662
- Ribavirin
 action of, 215e–6
 for adenovirus, 1208
 adverse effects of, 215e–6, 381, 897, 2042–2043
 for chronic HCV infection, 215e–4t, 215e–6, 215e–10, 927, 2041–2042
 drug interactions of, 2030
 genetic variations in response to, 39t
 for HRSV infection, 215e–2t, 215e–6
 for influenza, 1214
 for parainfluenza virus infection, 1208
 for viral encephalitis, 897
 for viral hemorrhagic fevers, 215e–6, 261e–5t, 261e–8, 783, 1320
- Riboflavin (vitamin B₂), **96e–3**
 deficiency of, 96e–1t, 96e–4, 463t
 functions of, 96e–2f, 96e–3–4
 parenteral requirement for, 98e–5t
 recommended intake of, 95e–2t
 reference range for, 2766t
 structure of, 96e–2f
 supplements, adverse effects of, 216
- Rice water stool, 1063, 1063f
- Rickets
 clinical features of, 96e–1t
 diagnosis of, 2465–2466
 hereditary vitamin D–resistant, 2465
 in hypophosphatasia, 426e–5
 hypophosphatemic, 332e–7t, 2458
 pathogenesis of, 96e–1t, 2465
 vitamin D–dependent type I (pseudovitamin D–deficiency), 2465, 2485
 vitamin D–dependent type II, 2485
- Rickettsia africana*, 1157
- Rickettsia akari*. *See* Rickettsialpox
- Rickettsia australis*, 128t, 1157
- Rickettsia conorii*, 782. *See also* Mediterranean spotted fever
- Rickettsia felis*, 1158
- Rickettsia honei*, 1157
- Rickettsia japonica*, 1157
- Rickettsial diseases, **1154**
 clinical features of, 127, 128t, 1154, 1155t, 2193
 diagnosis of, 150e–8
 ehrlichioses, 1159–1160, 1159f
 endemic typhus. *See* Endemic (murine) typhus
 epidemic typhus. *See* Epidemic (louse-borne) typhus
 epidemiology of, 1154, 1155t
 etiology of, 1154, 1155t
 flea-borne spotted fever, 1158
 global considerations in, 782
 human granulocytic anaplasmosis, 1152, 1159f, 1160–1161
 Mediterranean spotted (boutonneuse) fever, 128t, 782, 1157
 Q fever. *See* Q fever
 rickettsialpox. *See* Rickettsialpox
 Rocky Mountain spotted fever. *See* Rocky Mountain spotted fever (RMSF)
 scrub typhus. *See* Scrub typhus
 tick-borne spotted fevers, 1157
 in travelers, 802t
- Rickettsialpox
 clinical features of, 132t, 1155t, 1157–1158
 diagnosis of, 1158
 epidemiology of, 132t, 1155t, 1157
 etiology of, 132t
- rash in
 evolution of, 132t, 829, 1157–1158, 1157f, 1158f
 papules and plaques, 25e–9f, 132t, 134–135, 362
 transmission of, 1155t
 treatment of, 1158
 vs. varicella, 135, 1185
- Rickettsia prowazekii*, 782, 1154. *See also* Epidemic (louse-borne) typhus
- Rickettsia rickettsii*, 782. *See also* Rocky Mountain spotted fever (RMSF)
- Rickettsia sibirica*, 128t
- Rickettsia typhi*. *See* Endemic (murine) typhus
- Riedel's thyroiditis, 2299
- Rifabutin
 action of, 205e–5, 931t, 932, 936
 adverse effects of, 205e–5
 drug interactions of, 246e–3t
 for MAC infection prophylaxis in HIV infection, 1251t
 for nontuberculous mycobacterial infections, 205e–8
 pharmacology of, 205e–5
 resistance to, 931t, 934
 for TB, 205e–5
 for TB prophylaxis in HIV infection, 1251t
- Rifampin
 action of, 205e–3–4, 931t, 932f, 933
 adverse effects of, 205e–4
 allergic interstitial nephritis, 1857
 gastrointestinal, 944
 hepatotoxicity, 938t, 944
 hyperbilirubinemia, 282
 thrombocytopenia, 727t
 for anthrax, 261e–3, 261e–5t
 for asymptomatic GAS carriage, 966
 for *Brucella* infections, 194e–4
 for cat-scratch disease, 1081t
 dosage of, 205e–4
 drug interactions of, 42t, 205e–4, 939t, 1126
 antiretroviral therapy, 205e–1
 atovaquone, 246e–2t
 dapsone, 246e–4t
 quinidine, 246e–6t
 genetic variations in response to, 39, 39t
 indications for, 941t, 943–944
 for infective endocarditis, 822t, 823, 963
 for *Legionella* infections, 1020, 1020t
 for leprosy, 1126–1127, 1127t
 for meningitis, 889, 952
 metabolism of, 935
 for nontuberculous mycobacterial infections, 205e–8, 1131–1132, 1696
 for osteomyelitis, 840t, 841t, 843
 pharmacology of, 205e–4
 prophylactic
 for *H. influenzae*, 235
 for meningitis, 889, 945t
 for TB in HIV infection, 1251t
 for prosthetic joint infections, 963
 for Q fever, 1162
 for reactive arthritis, 2175
 resistance to
 detection of, 778
 mechanisms of, 931t, 934
 in mycobacteria, 205e–4, 774, 1118
 prevalence of, 941t
 for TB
 active infection, 205e–2t, 205e–3–4, 1115–1117, 1115t
 latent infection, 205e–1t, 205e–2t, 205e–3, 1121
 for tuberculous meningitis, 898
 for verruga peruana, 1081t
- Rifamycins, 931t, 933, 943
- Rifapentine
 action of, 931t, 933
 adverse effects of, 205e–5
 for latent TB, 205e–1t, 205e–5, 1121
 for leprosy, 1126
 pharmacology of, 205e–5
 for pulmonary TB, 205e–5
 resistance to, 205e–5, 931t

- Rifaximin
for *C. difficile* infections, 861
for hepatic encephalopathy, 2066
for irritable bowel syndrome, 1969, 1970t
for travelers' diarrhea, 268, 797, 857, 857t
- Rift Valley fever
clinical features of, 1322
epidemiology of, 151e-8, 151e-8f, 782, 1321-1322
vaccine for, 1322
- Right atrial overload, ECG in, 1454f
- Right atrial pressure, in shock, 1729, 1731f
- Right atrium, 1476, 1478f
- Right bundle branch, 1471
- Right bundle branch block
in Chagas disease, 1395
conditions associated with, 1453-1454
ECG in, 269e-4f, 269e-5f, 269e-7f, 278e-3f, 1454, 1455f
heart sounds in, 1447f
- Right-left confusion, 179
- Right-to-left shunt
intracardiac, 1463
intrapulmonary, 248, 249
- Right ventricular dysfunction
cardiogenic shock in, 1762
diagnosis of, 269e-8f, 270e-7, 270e-26v
etiology of, 270e-7
- Right ventricular failure/infarction
clinical features of, 1573t
differential diagnosis of, 1573t
in STEMI, 1608
- Right ventricular outflow tract obstruction
in pulmonic stenosis, 1525
in tetralogy of Fallot, 1526
in ventricular septal defect, 1522
- Right ventricular stroke work, 1748t
- Rigidity, 154
- Riley-Day syndrome, 2643
- Riloncept, 2214, 2234
- Rilpivirine, 1275t, 1278
- Riluzole, 2635
- Rimantadine
action of, 214e-9, 215e-5-6
adverse effects of, 215e-7
for influenza prophylaxis, 215e-2t, 215e-6, 1213t, 1214, 1252t
for influenza treatment, 215e-2t, 1213-1214, 1213t
pharmacology of, 215e-6-7
resistance to, 215e-1, 215e-5
- Ringer's lactate, 1749
- Ring(s) esophageal, 1902, 1902f
- Ringworm. *See* Dermatophytosis
- Rinne test, 221, 2538
- Riociguat, 1660, 1660t
- Rio Mamoré virus, 1308t
- Ri protein, 372e-25t
- Risedronate
adverse effects of, 2499
for osteoporosis management/prevention, 2383, 2499, 2500f, 2504
for Paget's disease of bone, 426e-3, 426e-3t
- Risperidone
adverse effects of, 2722t
for bipolar disorder, 2718
for delirium, 64, 64t
overdosage/poisoning with, 473e-9t
for psychosis in Huntington's disease, 2622
for schizophrenia, 2721, 2722t
- Ritodrine, 305
- Ritonavir
action of, 1278
adverse effects of, 1275t, 1278, 2030
drug interactions of, 34t, 42t, 246e-3t, 798, 1278
for HIV infection, 1275t, 1278
metabolism of, 34t, 42
molecular structure of, 1276f
resistance to, 1280f
- Rituximab
action and targets of, 102e-5t, 103e-21, 103e-21t
action of, 2147
- adverse effects of, 2130t
common, 2146t, 2147
cutaneous, 377
cytokine release syndrome, 1796
hypersensitivity reaction, 1798
infections, 485, 488, 490, 2149
paraneoplastic pemphigus, 372
progressive multifocal leukoencephalopathy, 372e-31, 2147
serious, 2146t, 2182t, 2187
tumor lysis syndrome, 1795
- for autoimmune and inflammatory diseases, 372e-30
for autoimmune hemolytic anemia, 658
for cerebellar degeneration, 618
for CLL, 102e-5t, 102e-6, 103e-21, 703
for coagulation factor inhibitors, 735
for cold agglutinin disease, 658
for diffuse large B-cell lymphoma, 705-706
for follicular lymphoma, 705
for gastric lymphoma, 536
for granulomatosis with polyangiitis, 2185
for hemolytic-uremic syndrome, 1797
for hepatitis C-associated cryoglobulinemic vasculitis, 2191
for IgG4-related disease, 391e-3
for immune thrombocytopenic purpura, 728
for inflammatory myopathies, 2201
for kidney transplant rejection, 372e-30, 1829
for leukemia, 102e-5t
for mantle cell lymphoma, 704
for membranous glomerulonephritis, 1844
monitoring during treatment with, 2146t
for MS, 2671-2672
for mucous membrane pemphigoid, 374
for myasthenia gravis, 2705
for neuromyelitis optica, 2673
for non-Hodgkin's lymphoma, 372e-30
for paraneoplastic encephalitis, 617
for pemphigus vulgaris, 371
during pregnancy, 124e-4
for primary central nervous system lymphoma, 602
for rheumatoid arthritis, 2146t, 2147
for Sjögren's syndrome, 64e-7
for SLE, 2130t, 2133
for systemic sclerosis, 2164
for thrombotic thrombocytopenic purpura, 730
for Waldenström's macroglobulinemia, 718
- Rivaroxaban
adverse effects of, 757, 1488
for antiphospholipid antibody syndrome, 2136
dosage of, 757
for DVT/PE, 1635
indications for, 756-757
management of
before endoscopic procedures, 1889t
before lumbar puncture, 443e-1
monitoring treatment with, 757
pharmacology of, 756t
in pregnancy, 757
for stroke prevention in atrial fibrillation/atrial flutter, 1485t, 1488, 2570
for VTE prophylaxis, 1637t
- Rivastigmine, 176, 2601, 2617
- River blindness. *See* *Onchocerca volvulus*/onchocerciasis
- Rizatriptan, 2591, 2592t, 2593t
- RLRs (retinoic acid-inducible gene-1-like receptors), 372e-2, 372e-4, 372e-4t
- Rmp protein, 1004
- RMRP gene, 375e-2
- RMSF. *See* Rocky Mountain spotted fever (RMSF)
- RNA polymerase (RNP), 372e-24t, 2157t
- RNA virus(es), 214e-2t, 214e-3f, 214e-4. *See also* specific viruses
- Robertsonian translocation, 83e-5-6
- ROC (receiver operative characteristic curve), 21, 22f
- Rocio virus, 1307t
- "Rocker foot," 2244
- Rocky Mountain spotted fever (RMSF)
clinical features of, 1155, 1155t
cutaneous
characteristics of, 130t
cutaneous, 888
evolution of, 134, 782, 888, 1155
petechial rash, 25e-4f, 76e-9f, 782, 1155, 1156f
purpura, 369
ECG changes, 1561
sepsis, 1754
diagnosis of, 1156-1157
differential diagnosis of, 888, 1156
epidemiology of, 1154, 1155t
etiology of, 780t, 782
fulminant, 1156
pathogenesis of, 1154-1155
prevention of, 1157
treatment of, 780t, 782, 1157
- Rod(s), 195
- Rodent(s), in plague, 261e-4
- Rodent bite-wound, 167e-1-2, 167e-3t
- Rodent-borne viruses, 1304. *See also* specific viruses
diagnosis of, 1310
disease syndromes of, 1306-1310t, 1310, 1313. *See also* specific diseases
arthritis and rash, 1312t, 1313-1314
encephalitis, 1312t, 1314-1317
fever and myalgia, 1312t, 1317-1319
hemorrhagic fevers. *See* Viral hemorrhagic fevers
pulmonary disease, 1312t
epidemiology of, 1304
geographic distribution of, 1311-1312t
transmission of, 1305
- Rofecoxib, 43, 2232
- Roflumilast, 1706
- Rolfing/structural integration, 14e-1t
- Roll Back Malaria campaign, 14
- Romaña sign, 1395
- Romberg maneuver, 2539
- Romberg's sign, 158, 164
- Rombo syndrome, 500
- Romidepsin, 102e-5t, 103e-18t, 103e-20
- Romiplostim, 729
- Root canal therapy, 237
- Root cavity, 46e-4f
- Ropinirole
adverse effects of, 2615
for PD, 2615, 2616t
for restless legs syndrome, 188t, 192, 2625
- ROS. *See* Reactive oxygen species (ROS)
- ROS1 gene mutations, 508, 508f
- Rosacea. *See* Acne rosacea
- Rosai-Dorfman disease (sinus histiocytosis with massive lymphadenopathy), 710
- Rosenbaum card, 195, 196f
- Roseola (exanthem subitum), 25e-2f, 127, 128t, 1194
- Rose spots, 25e-3f, 1051f
- Rosiglitazone
action of, 96e-6, 2415
adverse effects of, 2414t, 2415
contraindications to, 2414t
for NALFD, 2057
for type 2 diabetes mellitus, 2414t, 2415
- Ross procedure, 1532
- Ross River virus infection (epidemic polyarthritis), 837, 1306t, 1313
- Rosuvastatin, 2448t
- Rotational atherectomy, 296e-2
- Rotator cuff tendinitis, 2221, 2248, 2248f
- Rotavirus, 1286f, 1286t, 1287
- Rotavirus infections
clinical features of, 258, 266t, 855t, 1288
diagnosis of, 267, 1286t, 1289
epidemiology of, 855, 1287-1288, 1288f
global considerations in, 1287, 1288f
immunity to, 1288-1289
pathogenesis of, 1288
prevention of, 1289
in transplant recipient, 923
treatment of, 1289

- Rotavirus vaccine, 1289
 Roth's spots, 40e-4, 200, 201f
 Rotigotine, 2615, 2616t
 Rotor syndrome, 282, 2003t, 2004
 Rouleaux formation, 81e-1, 81e-3f
 Round pneumonia, 951
 Roundworms, 255e-1. *See also* Nematodes/
 nematode infections
Rousettus aegyptiacus, 1324
 Rous sarcoma virus, 101e-2, 225e-1t
 Roux-en-Y gastric bypass, 2398, 2398f, 2434
 Rowasa enemas, 1960
 RPE65 gene, 91e-3
 RPR (rapid plasma reagin) test, 1137, 1137f
 16S rRNA genes, bacterial, 86e-2-4
 18S rRNA genes, eukaryotic, 86e-4
 RSDL (reactive skin decontamination lotion),
 262e-6
 RSP01 gene mutations, 2356t
 RSV infections. *See* Human respiratory syncytial
 virus (HRSV) infections
 rT₃ (reverse T₃), 2283f
 RTA. *See* Renal tubular acidosis (RTA)
 RTH (resistance to thyroid hormone), 435,
 2286t, 2287
 rtPA (recombinant tissue plasminogen activator).
See Alteplase (rtPA)
 RT-PCR (reverse transcriptase polymerase chain
 reaction), 1247, 1247t
 "Rubber person" syndrome, 2510
 Rubella (German measles), 230e-1
 clinical features of, 128t, 230e-1, 362
 oral, 236
 skin, 25e-2f, 127, 128t, 230e-1f, 362, 362t
 congenital, 50, 230e-1, 230e-2, 230e-2t, 785t,
 900, 1525
 diagnosis of, 230e-1-2, 1190t
 epidemiology of, 230e-1
 global considerations in, 230e-1, 230e-2-3,
 230e-3f
 incidence pre- and postvaccine, U.S., 785t
 pathogenesis of, 230e-1
 in pregnancy, 50, 127, 230e-2
 prevention of, 230e-2-3
 treatment of, 230e-2
 Rubella vaccine, 50, 230e-2-3, 230e-3f. *See*
also Measles-mumps-rubella vaccine
 (MMR)
 Rubella virus, 230e-1
 Rubeola. *See* Measles (rubeola)
 Rubidium-82, 270e-3, 270e-3t
 Rubinstein-Taybi syndrome, 431t
 Rubivirus, 230e-1. *See also* Rubella (German
 measles)
 Rubor, 414f, 1643, 1753
 Rubrospinal pathway, 155f
 Rufinamide, 2555t
 Rumination, 256, 1901
 Rumination disorder/syndrome, 259, 2719
 RUNX1 gene mutations, 135e-6, 680, 680t
 Runx2, 2455
 Russell body gastritis, 1932
 Russell-Silver syndrome, 83e-7
 Russell's viper bite, 2734, 2734f, 2738. *See also*
 Snakebites
 Russell venom viper test, 2131
 Ruxolitinib
 action of, 102e-5t, 103e-19
 adverse effects of, 674, 676
 for polycythemia vera, 674
 for primary myelofibrosis, 676
 RV (residual volume), 306e-1f, 306e-2, 306e-2f,
 306e-5f
 R value, 2023
 R wave, 269e-5f, 1451
 Ryanodine receptor, 40-41
 RYR1 gene, 462e-12
 RYR3 gene, 1241, 1241t
- S**
 S-100 protein, in carcinoma of unknown
 primary, 120e-2t
 Saaremaa virus, 1307t
 Sabia virus, 1306t
 Saccades, 197
 Saccadomania, 211
 Saccharomyces boulardii, 860, 860t
 Saccharopinuria, 434e-3t
 Sacular aneurysm
 aortic, 1637, 1638. *See also* Aortic aneurysm
 cerebral, 1784-1785. *See also* Subarachnoid
 hemorrhage
 Sacral agenesis, 48
 Sacral insufficiency fracture, 441e-33f
 Sacral pain, 118
 Sacral spinal lesions, 2651-2652
 Sacroiliitis
 in ankylosing spondylitis, 2169, 2171, 2172f
 in IBD, 1958
 in reactive arthritis, 2174, 2175
 SACT (sinoatrial conduction time), 1469
 Saddle-nose deformity
 in granulomatosis with polyangiitis, 1443,
 2183
 in relapsing polychondritis, 1443, 2203, 2203f
 Safety in health care, 12e-1
 in developing countries, 12e-1
 factors that increase likelihood of errors,
 12e-1
 frequency of adverse events, 12e-1-2
 measurement of, 12e-2
 national picture, 12e-2
 prevention of adverse events, 12e-2
 systems theory and, 12e-1, 12e-1f
 Safety theory, 12e-1
 Sagging brain syndrome, 2607
 St. John's wort, 14e-4, 42, 42t, 44, 389t
 St. Louis encephalitis virus, 1307t, 1315
 St. Vitus' dance. *See* Sydenham's chorea
 Salary compensation plan, 21
 Salicylates
 adverse effects of, 319, 324, 2130t
 drug interactions of, 42t, 103e-14
 overdose/poisoning with, 319, 473e-12t
 for rheumatic fever, 2152-2153
 for SLE, 2130t
 therapeutic monitoring of, 319
 Salicylic acid
 for acne vulgaris, 352
 for prickly heat, 479e-1
 for scabies, 2745
 for tinea versicolor, 350
 for warts, 351
 Saline
 half-normal, for hypovolemia, 298
 hypertonic, 302, 303, 309, 1780t
 hypotonic, 304
 isotonic, 298, 302, 2735
 normal
 for hyperkalemia, 312
 for hyponatremia, 301
 for hypovolemia, 298
 for sepsis/septic shock, 1757
 for SIADH, 610
 Saline infusion test, 1749, 2319
 Saliva
 artificial, 216
 characteristics of, 239
 HIV transmission via, 1223
 volume and electrolyte content of, 98e-4t
 Salivary duct obstruction, 241
 Salivary gland(s)
 benign tumors of, 503, 506
 diseases of, 239, 241
 enlargement of, 2167, 2167t
 IgG-4 related lesions of, 391e-1t, 391e-2,
 391e-2f
 malignant tumors of, 241, 503, 506
 Salivary stone (sialolithiasis), 46e-5f, 241
 Saliva substitutes, 241
 Salmeterol, 205e-3, 1705
 Salmonella spp.
 antibiotic resistance in, 941t, 1052, 1053, 1054
 chronic carriage of, 1052
 intercellular survival of, 145e-7
 laboratory identification of, 150e-2t
 subspecies/serotypes, 1049
Salmonella enteritidis, 1053
Salmonella spp. infections, 1049. *See also* Enteric
 (typhoid) fever
 antibiotic resistance in, 1053, 1054
 aortitis, 1642
 bacteremia, 1054
 bone, joint, and soft tissue, 1054
 in cancer patient, 486t
 CNS, 1054
 diagnosis of, 150e-2, 856, 1054
 endovascular, 1054
 epidemiology of, 1053
 food-related, 266
 gastroenteritis
 vs. appendicitis, 1987
 clinical features of, 266, 266t, 856t, 1053
 vs. IBD, 1956
 pathogenesis of, 1050
 treatment of, 857, 857t, 1054-1055, 1054t
 genital, 1054
 in HIV infection, 1252t, 1256-1257
 intraabdominal, 1054
 prevention and control of, 1055
 pulmonary, 1054
 in returned traveler, 802t
 splenic abscess, 851
 treatment of, 940t, 1054-1055, 1054t
 urinary tract, 1054
Salmonella typhi. *See also* Enteric (typhoid)
 fever
 as bioterrorism agent, 261e-9
 host receptors for, 145e-3
 multidrug-resistance in, 1050
 pathogenesis of, 1049
 Salmonellosis, nontyphoidal. *See* *Salmonella* spp.
 infections
 Salmon River virus, 1306t
 Salp15, 1150
 Salpingitis
 abdominal pain in, 106t, 107, 877
C. trachomatis, 1169, 1170t
 clinical features of, 877
 diagnosis of, 877
 Salpingo-oophorectomy, 593, 594, 597
 Salsalate, 2232t
 Salt-depletion heat exhaustion, 479e-2
 Salt therapy
 for metabolic alkalosis, 322
 for SIAD, 302
 for syncope, 145
 Salt-wasting disorders, 295, 299, 1612-1613
 Salt-wasting 21-hydroxylase deficiency,
 2355-2356
 Salty taste sensation, 212
 Salvinorin A (*Salvia divinorum*), 469e-4
¹⁵³Samarium therapy, 61, 103e-22, 119e-4, 474
 SAME. *See* Syndrome of apparent
 mineralocorticoid excess (SAME)
 Sand flea, 2750
 Sandfly
 in Carrión's disease, 1083
 in *Leishmania* infections, 1387, 1388t
 in pemphigus foliaceus, 371
 in Punta Toro virus disease, 1318
 in Toscana virus disease, 1318
 Sandfly fever, 152e-5t, 1310t, 1318
 Sandhoff's disease, 432e-1, 432e-3t, 2634t
 SANDO (sensory ataxic neuropathy and
 ophthalmoparesis), 451e-2t
 Sandostatin-LAR, 2271
 Sanfilippo syndromes, 432e-2t
 Sanjad-Sakati syndrome, 2483
 SA (sinoatrial) node, 273e-1, 1451, 1451f,
 1466-1467, 1466f
 SA (sinoatrial) node disease/dysfunction.
See Sinoatrial (SA) node disease/
 dysfunction
 SAPE (sentinel acute pancreatic event)
 hypothesis, 2098
 Saphenous vein bypass graft, 296e-2, 297e-3,
 297e-3f, 297e-3v, 1645
 SAPHO syndrome, 844, 2179
 SAPIEN valve, 296e-5, 297e-6f
 Sapovirus, 1286f, 1286t
 SAPS scoring systems, 1729

- Saquinavir
 drug interactions of, 246e-3t
 metabolism of, 34t
 molecular structure of, 1276f
 resistance to, 1280f
- Sarcoglycans, 462e-8, 462e-8f
- Sarcoidosis, **2205**
 clinical features of, **2206**, 2206t
 alopecia, 355, 355t
 azotemia, 292
 bone marrow involvement, 2208
 breast lesions, 2209
 cardiac, 290e-1t, 2208
 AV conduction block, 271e-7f, 1471, 1471t
 myocarditis, 1561, 1562f
 ventricular tachycardia, 1495
 cough, 244
 cutaneous
 characteristics of, 367, 1443, 2207
 figurate lesions, 356, 356t
 hypopigmentation, 357t
 lupus pernio, 364t, 367, 2207, 2207f
 maculopapular lesions, 76e-17f, 2207, 2207f
 extrahepatic cholestasis, 284
 facial palsy, 2648
 hepatic, 2207-2208
 hypercalcemia, 313, 2208, 2477
 hypocalcemia, 314
 hypopituitarism, 2256
 lymphadenopathy, 409
 musculoskeletal, 2209, 2209f
 neurologic, **2208**
 meningitis, 909t
 myelitis, 2656
 neuropathy, 2683
 ocular, 40e-4f, 199, 2207
 oral, 241
 pulmonary, 308e-8-9f, 308e-9f, 2206-2207, 2206f
 pulmonary hypertension, 1659
 renal, 2208
 spleen, 2208, 2208f
 weakness, 156
 complications of, 2209
 definition of, 2205
 diagnosis of
 approach to, 2209-2210, 2210f
 bronchoalveolar lavage, 1711t
 chest radiograph, 308e-8f, 308e-9f, 2206f, 2209
 CT, 308e-8f, 308e-9f, 2206f
 Kveim-Siltzbach procedures, 2210
 laboratory findings, 2209-2210
 MRI, 441e-11-15f, 2209
 PET, 2209, 2209f
 urinalysis, 62e-11f
 differential diagnosis of, 367, 2167t, 2208
 etiology of, 2205
 global considerations in, 2205
 immunopathogenesis of, 2205-2206, 2205f
 incidence of, 2205
 vs. occupational lung disease, 1691
 prognosis of, 2210
 treatment of, 1560, 2210-2212, 2211f, 2212t
- Sarcoma, **119e-1**
 bone. *See* Bone sarcoma
 cardiac, 289e-2
 deaths from, 470t
 definition of, 102e-1, 103e-1
 follicular dendritic cell, 135e-9
 gastric, 536
 genetic considerations in, 101e-3
 granulocytic, 366
 histiocytic, 135e-9
 interdigitating dendritic cell, 135e-9
 Langerhans cell, 135e-9
 metastatic, 308e-16f, 604t
 paraneoplastic syndromes in, 609t
 soft tissue. *See* Soft tissue sarcoma
 synovial, 2247
 uterine, 596, 597
- Sarcoma virus, for gene therapy delivery, 91e-1t
- Sarcomere, 265e-4
- Sarcomeric proteins, 1553, 1555t, 1556f
- Sarcopenia, 82, 94e-4
- Sarcoplasmic reticulum, 265e-6, 265e-8f, 1503
- Sarcoptes scabiei*, 2744. *See also* Scabies
- Sarcosinemia, 434e-2t
- sar gene, 955
- Sarin. *See* Nerve agents
- SARS. *See* Severe acute respiratory syndrome (SARS)
- Saturated solution of potassium iodide (SSKI), 1354, 1354t
- Sauna taker's lung, 1681t
- Saxagliptin, 291e-7, 2414t
- Saxitoxin, 2743
- SCA. *See* Spinocerebellar ataxia (SCA)
- Scabicides, 2743-2744
- Scabies, **2744**
 clinical features of, 76e-10f, 341t, 2744
 diagnosis of, 2744
 epidemiology of, 2744
 treatment of, 246e-9, 2744-2745
- Scaffolds, for tissue engineering, 92e-1
- Scale-free network, 87e-1, 87e-2f
- Scalene maneuver, 1647
- SCAMPS, 1914
- SCAN1 (spinocerebellar ataxia with neuropathy), 451e-4t
- Scapular winging, 462e-2f, 462e-11
- Scaritoxin, 2742
- Scarlatiniform eruptions, 362-363, 362t. *See also* Rash
- Scarlet fever (second disease)
 clinical features of, 130t, 232, 362-363, 966
 differential diagnosis of, 134, 966
 epidemiology of, 130t
 etiology of, 130t, 232, 966
 rash in
 differential diagnosis of, 127
 erythematous, 25e-6f, 134
 evolution of, 130t
 scarlatiniform, 362-363, 362t, 966f
- Scavenger receptors, 291e-1f, 291e-2
- SCCmec, 961
- Scedosporium* spp., 1355, 1356
- Scedosporium* spp. infections, 1356
- SCF (stem cell factor), 372e-11t
- Schatzki's ring, 256, 1895, 1896f, 1902, 1910f
- Scheie disease, 432e-2t, 432e-6
- Schilling test, 350e-1, 350e-1t, 1938
- Schimke immunosseous dysplasia, 375e-2
- Schinz-Giedion syndrome, 135e-5
- Schirmer strip, 199
- Schistocytes, 81e-1, 726f, 729, 737
- Schistosoma* spp. *See also* Schistosomiasis
 adherence of, 145e-3
 geographic distribution of, 245e-2t, 765t, 1423t, 1425f
 laboratory identification of, 245e-1, 245e-2t, 245e-5t, 245e-6t
 life cycle of, 245e-2t, 1424, 1424f
 transmission of, 1423t, 1425
- Schistosomiasis, **1423**
 bladder cancer and, 477t, 575, 1427
 clinical features of, 152e-4t, 1426-1427
 CNS, 1427
 eosinophilia, 245e-6t
 hepatosplenic, 1426f, 1427
 intestinal, 1426-1427
 myelitis, 2657
 pulmonary, 1427, 1659
 renal, 1836t, 1849
 skin, 829, 1426
 swimmer's itch, 1426
 diagnosis of, 152e-4t, 245e-2t, 245e-3, 245e-6t, 765t, 1427
 epidemiology of, 245e-2t, 765t, 1425, 1425f
 etiology of. *See* *Schistosoma* spp.
 immunity to, 1426
 incubation period for, 152e-4t
 pathogenesis of, 1425-1426
 prevention and control of, 1428
 in returning war veterans, 152e-4t, 152e-8
- in travelers, 245e-3, 797
 treatment of, 152e-4t, 1428, 1428t
 metrifonate, 246e-10
 oxamniquine, 246e-11
 praziquantel, 246e-12, 1428t
- Schizoaffective disorder, 2721
- Schizogony, 1369
- Schizoid personality disorders, 2721
- Schizont, *Plasmodium*, 1369
- Schizophrenia, **2720**
 CATIE study of, 2722
 chemosensory alterations in, 215
 clinical features of, 2720-2721
 vs. dementia, 2608
 diabetes mellitus type 2 in, 2722
 differential diagnosis of, 2721
 epidemiology of, 2721
 genetic considerations in, 444e-1, 465e-2, 465e-4, 2721
 insomnia in, 190
 neuropathology of, 465e-3-4
 pathophysiology of, 2721
 treatment of, 2721-2722, 2722t
- Schizophreniform disorder, 2721
- Schizotypal personality disorders, 2721
- Schmid metaphyseal chondrodysplasia, 2512
- Schnitzler's syndrome, 141, 372e-8t, 2215
- Schober test, modified, 2170
- Schwann cell tumor, benign (neurofibroma), 364
- Schwannomas, 364t, 602-603. *See also* Vestibular schwannoma (acoustic neuroma)
- Schwannomatosis, familial, 599t
- Sciatica, 121, 2222, 2222f
- Sciatic neuropathy, 2692
- SCID. *See* Severe combined immunodeficiency (SCID)
- Science of medicine, 1
- Scintigraphy
 adrenal, 1619
 in fever of unknown origin, 140
 in respiratory disease, 1664
- Scl-70 (topoisomerase I), 103e-13, 372e-24t
- Sclera, blue, 1443
- Scleral icterus, 279
- Scleredema, diabetic, 2430
- Scleritis, 200
- Sclerodactyly, 375, 1649, 2159, 2159f
- Scleroderma. *See* Systemic sclerosis (scleroderma)
- Scleroderma esophagus, 1911
- Scleroderma renal crisis, 62e-9f, 1864-1865, 2158, 2162, 2164. *See also* Systemic sclerosis (scleroderma)
- Scleromyxedema (papular mucinosis), 364t, 365
- Sclerosing cholangitis. *See* Primary sclerosing cholangitis (PSC)
- "Sclerosis of the left cardioskeleton," 1471
- Sclerosteosis, 426e-5
- Sclerostin antibodies, 2503
- Sclerotherapy, for chronic venous disease, 1652
- SCN (severe congenital neutropenia), 375e-1, 2104, 2105f
- SCN1A gene mutations, 2545t, 2589
- SCN4A gene mutations, 310, 462e-18
- SCN5A gene mutations, 1472, 1496t, 1498, 1554
- SCN9A gene mutations, 1650
- Scolex, 255e-1
- Scoliosis, 118, 462e-6
- Scombroid poisoning, 267, 2743-2744
- Scopolamine
 for excess secretions, 62, 68, 69t
 for nausea and vomiting, 62, 260, 260t
 overdose/poisoning with, 473e-8t
 in terminal weaning, 67
 for vertigo, 151t
- Scorpionfish envenomation, 2740, 2742
- Scorpion sting, 2748
- Scotoma, 197, 198f, 204
- Scrapie, 453e-1t, 453e-2

- Screening tests, 2, 26. *See also specific diseases and tests*
 adverse outcomes of, 28
 age-specific recommendations, 30–31t
 evidence-based recommendations for, 28–30, 29t
 health benefits of, 27–28, 28t
 principles of, 27t, 29
 Screening trials, 480
 Screwworm fly, 2747
 Scrub typhus
 clinical features of, 128t, 152e–3t, 1155t, 1159, 1561
 diagnosis of, 152e–3t, 1159
 epidemiology of, 128t, 1155t, 1159
 etiology of, 782, 1155t, 1159
 incubation period for, 152e–3t
 prevention of, 152e–3t
 rash in, 128t, 135, 782, 1159
 in returning war veterans, 152e–3t
 treatment of, 152e–3t, 1159
 Scurvy
 clinical features of, 96e–1t, 96e–5, 461, 732
 etiology of, 96e–5
 oral manifestations of, 237
 skin manifestations of, 367, 436e–1f, 463t
 treatment of, 96e–5
 SDF (stromal cell–derived factor), 372e–12t, 1235
 SDHB gene mutations, 2332, 2334f, 2335, 2344
 SDHC gene mutations, 2334f
 SDHD gene mutations, 2332, 2334–2335, 2334f
 SDH (succinate dehydrogenase) enzyme, 2329, 2332
 Sea anemone envenomation, 2738
 Seabather's eruption, 2741, 2741f
 Sea cucumber envenomation, 2740
 Seadornaviruses, 1305, 1310t
 Seal bite-wound, 167e–1, 167e–3
 Sea levels, rise in, 151e–3
 Seal finger, 167e–1, 167e–3
 Sealpox, 220e–1t
 Sea snake envenomation, 2734
 Seasonal affective disorder, 2715
 Sea sponge, 2739
 Sea urchins, 2739–2740, 2741f
 Sebaceous adenoma, 364t, 365
 Sebaceous carcinoma, 365
 Sebaceous hyperplasia, 365
 Sebastian's syndrome, 729
 Sebopsoriasis, 347
 Seborrheic dermatitis, 347
 clinical features of
 distribution and morphology of skin lesions, 341t, 342f, 347, 354t
 erythroderma, 354t
 facial, 76e–2f, 347f
 in HIV infection, 1262
Malassezia infection in, 1357
 treatment of, 347, 354t
 Seborrheic keratosis, 76e–3f, 341t, 342f, 359, 360t
 Secobarbital, 473e–10t
 Secondary taste cortex, 214
 Second disease. *See* Scarlet fever (second disease)
 Second-hand smoke. *See* Smoking, passive exposure to
 Second impact syndrome, 457e–4, 471e–5–6
 Secretin test, 2087t, 2100
 Secretion defects, 730
 Secretory diarrhea, 268–269, 268t, 303, 1933. *See also* Diarrhea
 Secretory leukocyte protease inhibitor (SLPI), 1223
 Secretory protein, 372e–22
 Secukinumab, 2173, 2177
 Sedation
 of critically ill patient, 1782
 of mechanically ventilated patient, 1732–1733, 1741, 1743
 for myocardial infarction, 1606
 Sedative-hypnotics, 473e–3, 473e–10t. *See also specific drugs*
 Segawa variant (dopa-responsive) dystonia, 434e–2t, 2620
 Seizure(s), 2542. *See also* Epilepsy
 absence, 2543
 after head injury, 457e–2–3
 in alcohol withdrawal, 2728
 approach to the patient, 2548, 2549f
 atonic, 2543–2544
 in brain abscess, 901
 in brain tumor, 598
 in cancer patient, 1793
 chemotherapy-induced, 607t
 classification of, 2542–2543, 2542t
 coma in, 1773
 continuous. *See* Status epilepticus
 definition of, 2542
 delirium and, 169, 2607
 differential diagnosis of, 2550–2551, 2551t
 eclamptic, 46
 EEG in, 442e–1–2, 442e–2f, 2550
 etiology of, 2544–2547, 2546t, 2547t
 evolution from focal seizures, 2543
 febrile, 126, 2545–2546
 focal, 2542–2543, 2586
 focal sensory, 162
 generalized, 2543
 in head injury, 2547
 in heatstroke, 479e–3
 history and examination in, 2548–2550
 in HIV infection, 1265, 1265t
 in hyperperfusion syndrome, 463e–1
 imaging in, 2550
 laboratory studies in, 2550
 lactic acidosis in, 318
 magnetoencephalography in, 442e–3, 2550
 in malaria, 1373
 mechanisms of, 2547–2548
 in meningitis, 887
 myoclonic, 2544
 in neurocysticercosis, 1430–1431, 1432
 in overdose/poisoning, 473e–3, 473e–5
 in pregnancy, 46, 49
 psychogenic, 2551
 pyridoxine-dependent, 434e–3t
 vs. syncope, 147, 2550–2551, 2551t
 temporal lobe, 183
 tonic-clonic, 2543
 treatment of, 2551. *See also* Epilepsy, treatment of
 unclassifiable, 2544
 Selectins, 139e–1, 416, 417f
 Selection bias, 480
 Selective AR modulators, 2371
 Selective digestive decontamination, in burn patients, 166e–3
 Selective estrogen response modulators (SERMs)
 action of, 102e–4f, 102e–7, 2499
 in osteoporosis management/prevention, 2498–2499
 Selective IgA deficiency, 2112
 Selective serotonin reuptake inhibitors (SSRIs)
 adverse effects of, 2710t, 2716
 serotonin syndrome. *See* Serotonin syndrome
 sexual dysfunction, 327, 2716
 SIADH, 300
 for dementia, 175
 for depression, 2710t, 2716, 2716f
 for depression in AD, 2602
 for depression in terminally ill patient, 64
 drug interactions of, 40, 2711t, 2717
 genetic variations in response to, 40
 for inappropriate sinus tachycardia, 1477
 for irritable bowel syndrome, 1969, 1970t
 overdose/poisoning with, 473e–13t
 for pain, 94
 pharmacology of, 2716–2717
 for premature ejaculation, 325
 for PTSD, 471e–4–5, 2712
 for vertigo, 151, 151t
 Selegiline
 adverse effects of, 2710t
 for depression, 2710t, 2717
 overdose/poisoning with, 473e–8t
 for PD, 2615, 2616t
 Selenium, 96e–10
 deficiency of, 96e–9t, 96e–10, 533, 1563
 dietary sources of, 96e–10
 parental supplementation of, 98e–6t
 for prostate cancer prevention, 478
 recommended intake of, 95e–3t
 reference range for, 2766t
 tolerable upper intake level of, 96e–9t
 toxicity of, 96e–9t, 96e–10
 Selenium sulfide
 for tinea capitis, 355t
 for tinea versicolor, 350, 350t, 358t
 Selenocysteine, 96e–10
 Selenosis, 96e–10
 Self-renewal, of stem cells, 453
 Seligman's disease (alpha heavy chain disease), 537, 718–719
 Sella chordomas, 2264
 Sellar fractures, 457e–2
 Sellar mass lesions, 2261. *See also specific lesions*
 clinical features of, 2261–2262, 2261t
 histologic investigation of, 2262
 hypogonadotropic hypogonadism and, 2365
 laboratory investigation of, 2262
 MRI in, 2262, 2262f
 ophthalmologic evaluation in, 2262
 Semantic dementia, 182
 Semantic paraphasia, 177
 Semen analysis, 2361, 2388
 Seminiferous tubules, 2358f, 2359–2360
 Seminoma. *See* Testicular cancer
 Semliki Forest virus, 1306t
 Senescence. *See also* Aging
 carcinogenesis and, 102e–1t, 102e–3
 definition of, 102e–8
 negligible, 94e–2, 94e–2f
 Senile purpura, 732
 Senior-Löken syndrome, 1850t, 1855
 Senna, 61, 62t, 273
 Sennetsu fever, 1159
 Sensation
 abnormalities of, 158
 in brainstem lesions, 161
 in cortical lesions, 161–162
 falls and, 165
 in focal seizures, 162
 localization of, 160–161
 in nerve and root lesions, 161
 in spinal cord lesions, 161
 in thalamic lesions, 161
 anatomy of, 158, 159f, 160f, 161f
 examination of, 158
 cortical, 159–160
 primary, 159, 160t
 quantitative sensory, 159
 positive and negative phenomena, 158
 terminology in, 158
 Sensenbrenner's syndrome (cranioectodermal dysplasia), 1850t
 Sensitivity
 analytical, 480e–2
 of tests, 21, 22, 479, 479t
 Sensitization, 87, 2114
 Sensory afferents, 47e–1
 Sensory ataxia, 158, 163, 164t
 Sensory ataxic neuropathy and ophthalmoparesis (SANDO), 451e–2t
 Sensory balance training, 163
 Sensory dissociation, 161
 Sensory examination, 2539
 Sensory neuropathy. *See* Peripheral neuropathy
 Sentinel acute pancreatitis event (SAPE) hypothesis, 2098
 Sentinel bleeds, subarachnoid, 1785
 Sentinel lymph node biopsy, 103e–3, 496–497, 528
 Seoul virus, 1307t
 Sepsis/septic shock, 1751
 anaerobic bacteremia and, 1100
 clinical features of, 780t, 781–783, 1754–1755
 clostridial, 991t, 994

- complications of
 adrenal insufficiency, 1755
 cardiopulmonary, 1746, 1755
 coagulopathy, 1755
 encephalopathy, 1782
 hypoglycemia, 2433
 hypothermia, 478e-1
 hypovolemia, 298
 immunosuppression, 1755
 intracranial hemorrhage, 2584
 neurologic, 169, 1755
 renal, 1755, 1801, 1806t
 in critically ill patient, 1734
 definition of, 1751, 1751t
 diagnosis of, 1755–1756
 epidemiology of, 1752
 etiology of, 780t, 781, 1751, 1752t, 1756
 hemodynamic patterns in, 1762t
 in HIV infection, 1757t
 in IV drug user, 1757t
 pathophysiology of, **1753**
 coagulation factors in, 1753
 control mechanisms in, 1753–1754
 cytokines in, 1753
 endothelial injury in, 1754
 host mechanisms and, 1752–1753
 organ dysfunction in, 1754
 physiologic characteristics of, **1748t**
 postsplenectomy, 484, 1757t
 prevention of, 1759
 prognosis of, 1758–1759
 puerperal, 968
 refractory, 1751t
 severe, 1754
 treatment of, 780t, **1756**, 1757t
 antimicrobial agents, 1756, 1757t
 empirical, 780t
 general support, 1757–1758
 hemodynamic, respiratory, and metabolic support, 1757
 investigational, 1758
 removal of infection source, 1756–1757
 without clear focus, 780t, 781
- Septal panniculitis. *See* Erythema nodosum
 Septic arthritis. *See* Infectious (septic) arthritis
 Septic diskitis. *See* Osteomyelitis, vertebral
 Septic emboli
 in infective endocarditis, 76e-12f, 818, 818f
 retinal, 40e-4
- Septicemia
 deaths from, 6e-1t
 hemolysis in, 657t
 meningococcal. *See* Meningococemia
 plague, 1072
 respiratory alkalosis in, 324
- Septic thrombosis
 of cavernous sinus, 783, 906
 of superior sagittal sinus, 783, 905–906
- Septo-optic dysplasia, 2256
 Septum primum, 1520
 Septum secundum, 1520
 SERCA2a, 91e-4, 1503, 1515
 Serelaxin, 1509, 1510t
 Serine kinase receptors, 400e-1t, 400e-3f, 400e-4
 Serine metabolism disorders, 434e-2t
 SERMs. *See* Selective estrogen response modulators (SERMs)
- Serotonin (5-HT)
 action of, 444e-3t
 in carcinoid syndrome, 565
 clinical aspects of, 444e-3t
 in coordination of emesis, 103e-25
 laboratory evaluation of, 566
 in migraine, 2588
 pain and, 88f
 structure of, 444e-3t, 566f
 synthesis, secretion, and metabolism of, 560f, 561
- Serotonin (5-HT_{1B/1D}) agonists
 adverse effects of, 2591
 for migraine, 2590–2591, 2592t
 nasal, 2591, 2592t
 parenteral, 2591, 2592t
- Serotonin (5-HT₄) agonists
 for gastroparesis, 260, 260t
 for irritable bowel syndrome, 1970
- Serotonin (5-HT₃) antagonists
 adverse effects of, 261
 for irritable bowel syndrome, 1970
 for nausea and vomiting, 109, 260, 260t
- Serotonin norepinephrine reuptake inhibitors (SNRIs)
 adverse effects of, 2710t
 for dementia, 175
 for depression, 64, 2710t
 for PTSD, 471e-5
- Serotonin (5-HT₃) receptor antagonists
 for carcinoid syndrome, 565, 566–567
 for nausea and vomiting, 62, 103e-25
 structure of, 566f
- Serotonin release assay, 752
- Serotonin syndrome
 clinical features of, 473e-13t
 etiology of, 473e-13t, 2624
 mechanism of, 473e-13t
 with SSRIs, 2716
 treatment of, 473e-6, 473e-13t
- Serra Norte virus, 1309t
- Serratia* spp. infections, **1034**
 antibiotic resistance in, 1034
 in cancer patient, 486t
 clinical features of, 1034
 diagnosis of, 1034
 ecthyma gangrenosum, 369
 epidemiology of, 1025, 1034
 sinusitis, 226
 treatment of, 1034
- Sertoli-Leydig tumor, 594
- Sertraline
 adverse effects of, 2710t, 2716
 for depression, 64, 473, 2672, 2710t, 2712, 2716
 for obsessive-compulsive disorder, 2714
 for panic disorder, 2709
 for PTSD, 2712
- Serum albumin. *See* Albumin, serum
 Serum ascites-albumin gradient, 287
 Serum bilirubin. *See* Bilirubin, serum
 Serum-binding proteins, 400e-2
 Serum ferritin. *See* Ferritin, serum
 Serum proteins, electrophoresis of, 710–712, 712f
- Serum sickness
 with antivenom, 2737
 with ATG treatment, 667
 clinical features of, 377, 384t, 2193, 2737
 immunologic mechanisms of, 44, 2180
 pathogenesis of, 377
 treatment of, 667, 2737
 urticaria/angioedema in, 363, 363t, 2118
 urticaria-like eruptions in, 132t, 135
- Serum sickness–like illness, 363, 377, 2193
 Setting sun sign, 210
 Sevelamer, 1816, 2458
- Seven transmembrane GPCR family, 372e-16, 400e-3, 400e-3f
- Severe acute respiratory syndrome (SARS)
 as bioterrorism agent, 261e-9
 clinical features of, 1204
 diagnosis of, 772, 1204
 epidemiology of, 916, 1204
 genome sequence of, 772
 infection-control practices for, 1205
 pathogenesis of, 1204
 super-spreaders, 1204
 surveillance for, 1205
- Severe combined immunodeficiency (SCID)
 clinical features of, 375e-1, 2107
 diagnosis of, 2107–2108
 etiology of, 2108
 genetic considerations in, 372e-16
 HSCT in, 139e-2, 139e-5, 139e-5t
 T cell differentiation and, 2108f
 treatment of, 91e-1–2, 2108
- Severe congenital neutropenia (SCN), 375e-1, 2104, 2105f
- Severe fever with thrombocytopenia syndrome, 1309t, 1322
- Severity-of-illness (SOI) scoring systems, 1729, 1730t
- Sex chromosome(s), 426, 437
 Sex chromosome disorders, 2350t
 Klinefelter's syndrome. *See* Klinefelter's syndrome
 ovotesticular DSD, 2351t, 2353
 Turner's syndrome. *See* Turner's syndrome
 45,X/46,XY mosaicism (mixed gonadal dysgenesis), 2351t, 2352–2353
- Sex cord tumors, ovarian, 594
- Sex development
 components of, 2349, 2349f, 2352f
 disorders of. *See* Disorders of sex development (DSDs)
 genetic regulation of, 2350f
 Sex differences, 6e-1, 7e-1. *See also* Men's health; Women's health
- Sex headache, primary, 2597
 Sex hormone-binding globulin, 332, 2378, 2763t
- Sex-influenced trait, 437
 Sex-limited trait, 437
 Sex-reversal, 431t, 435, 437
 Sex therapy, 329–330
- Sexual abuse
 gonococcal infections in, 1006, 1008
 sexual dysfunction in adulthood following, 326, 330t
- Sexual arousal disorder, female, 330
 Sexual assault, 6e-5
 Sexual dysfunction
 alcohol use and, 2725
 cocaine use and, 469e-1
 female, 330–331, 330t, 2427
 incidence of, 324
 male, 7e-1–2. *See also* Erectile dysfunction (ED)
 in MS, 2663, 2672
 SSRI-related, 2716
- Sexually transmitted infections (STIs), **869**. *See also specific infections*
 cervical cancer and, 594
 in children, 1006
 classification of, 869–870
 dysuria in, 60e-1
 epidemiology of, 869–870, 869t
 etiology of, 869t, 870t
 global considerations in, 869, 879
 HIV transmission and, 1221
 partner notification in, 883
 in pregnancy, 50
 prevention and control of, 882–883, 882f
 risk assessment for, 870, 871t
 screening for, 882–883
 in travelers, 797
- Sexual pain disorder, 330
 Sexual response, physiology of
 female, 330
 male, 324–325, 325f
- Sézary's syndrome, 81e-10f, 354, 391, 707
 SF1 (steroidogenic factor 1), 2310, 2349
 SF1 gene mutations
 in disorders of sex development, 2349, 2353, 2354t
 in hypogonadotropic hypogonadism, 2364t
- SF3B1 gene mutations, 135e-6
 S genes, of HBV, 2005–2006, 2005f
 SH3TC2 gene mutations, 444e-1
 Shah-Waardenburg syndrome, 359, 431t
 Shaking rigors, 478e-3
 Shannon index, 86e-4
 Shapiro syndrome, 478e-1
 Shared epitope, 2139
 Shawl sign, 2195
 SHBG gene mutations, 2360
 Sheehan's syndrome, 2257
- Shellfish poisoning
 amnestic (domoic acid intoxication), 2547–2548, 2743
 diarrhetic, 2741–2742
 paralytic, 2742
- Shewanella* spp. infections, 183e-2t, 183e-3
 Shift cells, 396, 396f

- Shift-work disorder, 193–194
- Shiga toxin–producing *E. coli* (STEC). *See also* *Escherichia coli* infections, intestinal characteristics of, 1028t, 1030 epidemiology of, 853, 855t, 857, 1028t, 1030 global considerations in, 1030
- Shigella* spp.
antibiotic resistance in, 1057 characteristics of, 1055 genomic signatures of, 1055 infectivity of, 145e-4, 145e-7, 1056, 1056f laboratory identification of, 150e-2t, 1057
- Shigella* spp. infections, **1055**
clinical features of, 266t, 855t, 856t, 1056–1057 complications of, 267, 1056–1057, 1058 diagnosis of, 150e-2, 856, 1057 epidemiology of, 1055 etiology of, 856t global considerations in, 1055, 1057 in HIV infection, 1257 vs. IBD, 1057 pathogenesis of, 1056, 1056f prevention of, 1058 treatment of, 1057–1058, 1058t
- Shingles. *See* Herpes zoster (shingles)
- Shivering, 124, 478e-1
- Shock
approach to the patient, 1729–1731, 1731f, 1744
cardiogenic. *See* Cardiogenic shock classification of, 1744, 1745t cyanosis in, 249 definition of, 1744 effect on drug concentration and response, 37
hypoadrenal, 1748t, 1750–1751 hypotension in, 1729–1730, 1744–1745 hypovolemic. *See* Hypovolemic shock hypoxia in, 248 lactic acidosis in, 318 neurogenic, 1748t, **1750** pathogenesis and organ response in, **1744**, 1745f
cardiovascular, 1746 cellular, 1745 inflammatory, 1746–1747, 1747f metabolic derangements, 1746 microcirculation, 1744–1745 neuroendocrine, 1745–1746 pulmonary, 1746 renal, 1746 respiratory failure in, 1732 septic. *See* Sepsis/septic shock traumatic, 1748t, **1750** treatment of
adjunctive therapies, 1751 algorithm for, 1749f mechanical ventilation, 1731 monitoring in, 1747–1748, 1748t rewarming, 1751
- Shohl's (sodium citrate) solution, 64e-6, 318, 320
- Shokwe virus, 1308t
- Shone's complex, 1529
- Short-bowel syndrome, **1943**
clinical features of, 1943 lactic acidosis in, 318 treatment of, 1944 vitamin A deficiency in, 96e-7
- Short-chain fatty acids, 1935, 1935t, 1936
- Shortened Fatigue Questionnaire, 464e-2, 464e-2f
- Short Physical Performance Battery, 78t
- Short QT syndrome, 1497
- Short stature
clinical features of, 2259–2260 etiology of, 2257 laboratory investigation of, 2259 nutritional, 2258 psychosocial, 2258 treatment of, 2259 in Turner's syndrome, 2352
- Short tandem repeats, 441
- Shotgun sequencing, 86e-1t, 86e-4, 772
- Shoulder
anatomy of, 2248f bursitis of, 2247 “frozen,” 2249 pain in, 123, 2221, 2221f tendinitis of, 2248
- Shunt
intracardiac, 1463 pulmonary, 248, 249, 306e-4, 306e-4f
- Shwachman-Diamond syndrome, 418, 664
- Shwachman syndrome, 375e-1
- Shy-Drager syndrome. *See* Multiple system atrophy (MSA)
- SIADH. *See* Syndrome of inappropriate antidiuretic hormone (SIADH)
- Sialadenitis, 215, 231e-2, 241
- Sialic acid, 145e-8
- Sialidosis, 432e-3t
- Sialolithiasis, 46e-5f, 241
- Siberian tick typhus, 128t
- Sib pair analysis, 442t
- Sick euthyroid syndrome, 478e-3, 2286, **2299**
- Sickle cell anemia, **634**
clinical features of, 634–635, 634t
acute chest syndrome, 634, 635–636 autosplenectomy, 413 hand-foot syndrome, 103e-26, 634, 2242 hyperkalemia, 307 jaundice, 281 musculoskeletal, 2242, 2242t nephropathy, 1835t, 1848, 1860, 1865–1866 painful crises, 634, 635 priapism, 325 pulmonary hypertension, 1659 splenic abscess, 851 splenic sequestration crisis, 634 stroke, 636, 2567 testicular atrophy, 2366 diagnosis of, 441e-41f, 634t, 635, 635f genetic considerations in, 87e-3, 435, 438 genetic testing for, 443, 445 high-altitude travel and, 476e-5 malaria resistance and, 442, 1371 pathophysiology of, 634, 634f peripheral blood smear in, 81e-4f, 635f reticulocyte index in, 399 systems biology applied to, 87e-5f treatment of, 139e-2, 139e-5, 635–636, 639
- Sickle cell crisis
clinical features of, 2242 hyperbilirubinemia in, 281 pathophysiology of, 634f splenic infarction in, 410
- Sickle cell trait, 634t, 635
- Sickle thalassemia syndrome, 634t, 635
- Sick sinus syndrome, 278e-3f, 1467, 1467t
- Siddha medicine, 14e-1t
- Sideroblast(s), 397
marrow, 627f ringed, 81e-9f, 399, 628, 668f
- Sideroblastic anemia
diagnosis of, 628t in myelodysplasia, 399 X-linked, 451e-4t, 2522f, 2522t, 2531–2532
- Sievert (Sv), 263e-1
- Sigmoidoscopy
for colorectal cancer screening, 29t, 30, 481t, 482, 540–541
in constipation, 273
in diarrhea, 267
in gastrointestinal bleeding, 279
indications for, 1881
in ulcerative colitis, 1952
- Sigmoid resection, 1973, 1973f
- Sigmoid volvulus, 1892, 1894f, 1982, 1985
- Signaling proteins, 372e-27t
- Signal recognition protein (SRP54), 372e-24t
- Signal transduction pathways
in cancer cells, 102e-3–7, 102e-4f, 102e-9f defects in neurologic disease, 444e-4 membrane receptor families and, 400e-1t in stem cells, 455
- “Signet-ring” sign, 1695
- Sign of Leser-Trelat, 359
- Sildenafil
action of, 43, 324, 325f, 326f adverse effects of, 328, 329t characteristics of, 329t contraindications to, 328, 329t development of, 43 for erectile dysfunction, 328, 2672 for heart failure, 1507 interaction with nitrates, 43, 1596, 1603 for lower urinary tract symptoms, 587 for pulmonary arterial hypertension, 1659, 1660, 1660t, 2164
for Raynaud's phenomenon, 1649, 2164
- Silent mutation, 433f
- Silent nociceptors, 88
- Silent (painless) thyroiditis, 2299
- “Silhouette sign,” 308e-10f
- Silibinin, 2025
- Silica, 1688t, 1689
- Silica exposure, 1689–1690, 1690f, 1840, 2126
- Silver, skin deposits, 361
- Silver nitrate, 166e-3
- Silver sulfadiazine, 166e-3, 262e-4
- Simeprevir, 215e-5t, 215e-11, 2047
- Simethicone, 263, 1969
- Simian immunodeficiency virus (SIV), 1216f
- Simple goiter, 2301
- Simple systems, 87e-1
- Simplicity heuristic, 19
- Simultaneous bilateral stimulation test, 180
- SIMV (synchronized intermittent mandatory ventilation), 1742. *See also* Mechanical ventilation
- Simvastatin
adverse effects of, 727t, 2199, 2218t, 2448t drug interactions of, 42, 246e-3t, 2447 genetic variations in response to, 38, 39t for lipoprotein disorders, 2448t
- Sindbis virus infection, 1306t, 1314
- Sinecatechins, 351, 1201, 1201t
- Single-balloon enteroscopy, 1881
- Single-fiber electromyography, 442e-6
- Single-nucleotide polymorphisms (SNPs)
analysis of, 83e-2f, 83e-3, 83e-3t in *CTFR* gene, 429f effects of, 441 frequency of, 427
- Single photon emission computed tomography (SPECT). *See* SPECT (single photon emission computed tomography)
- Single ventricle, 1527
- Sin Nombre virus, 1308t, 1319
- Sinoatrial (SA) conduction time (SACT), 1469
- Sinoatrial (SA) node, 273e-1, 1451, 1451f, 1466–1467, 1466f
- Sinoatrial (SA) node disease/dysfunction, **1467**
clinical features of, 1467–1468 diagnostic testing in, 1468–1469 ECG in, 1469, 1469f epidemiology of, 1466 etiology of, 1467, 1467t inherited forms of, 1467 pacemaker for, 1469–1470, 1470t treatment of, 1469
- Sinsemilla, 469e-2. *See also* Marijuana
- Sinus arrest, 1468
- Sinus bradycardia, 269e-17f, 273e-3t, 1609
- Sinus exit block, 1468
- Sinus histiocytosis with massive lymphadenopathy, 710
- Sinusitis, **226**
actinomycotic, 1088 acute, 226–227, 227t allergic fungal, 1347 anaerobic bacterial, 1097 *Aspergillus*, 489, 1346t, 1347 brain abscess and, 900–901 chronic, 227–228, 1097 in cystic fibrosis, 226, 1697 fungal, 226, 228 *H. influenzae*, 1011 in HIV infection, 1250 hospital-associated, 226

- in immunocompromised patient, 226
in influenza, 1212
M. catarrhalis, 226, 1013
maxillary, 239
Nocardia, 1087t
olfactory dysfunction in, 216
pneumococcal, 226, 952
rhinovirus, 1203
in stroke, 783
subdural empyema and, 903
viral, 226
- Sinus node, 1476, 1478f
Sinus node recovery time (SNRT), 1469
Sinus of Valsalva aneurysm, 1523
Sinusoidal obstructive syndrome, 139e-3, 286f
Sinus pause, 1468, 1468f
Sinus rhythm, 1476
Sinus tachycardia, **1476**
ECG in, 269e-3f, 269e-11f, 269e-17f, 278e-2f
inappropriate, 1476–1477
mechanism of, 3, 273e-3t
in myocardial infarction, 1609
physiologic, 1476, 1478t
in pulmonary edema, 1763
in pulmonary embolism, 100
in thyrotoxicosis, 2294
- Sinus thrombosis
cavernous, 783
superior sagittal, 441e-20–22f, 783
Sinus venosus, 1520
Sipuleucel-T, 103e-20, 586, 587
Sirolimus
action of, 1829, 1829t
adverse effects of, 1715, 1829t, 1831, 1864, 2071
in drug-eluting stents, 296e-2, 296e-3f
drug interactions of, 246e-3t
for immunosuppression, 1829, 1829t, 2071
for polycystic kidney disease, 1853
for pulmonary lymphangioleiomyomatosis, 1715
for reducing skin cancer risk in transplant recipients, 388
therapeutic monitoring of, 2765t
- SIRS (systemic inflammatory response syndrome), 98e-3, 98e-4
SIRT1, 94e-3, 94e-5
Sister Mary Joseph's node, 282, 534, 555
Sister of p-glycoprotein (bile salt excretory protein), 2004
Sitagliptin, 2414t
Siti. *See* Endemic syphilis
Sitosterolemia, 2440t, 2442
Situs inversus, 1528
SIV (simian immunodeficiency virus), 1216f
- Six-minute walk test, 78t, 1710
Sixth disease. *See* Exanthem subitum
Sixth nerve palsy, 1785
Sjögren's syndrome, **2166**
autoimmune diseases associated with, 2166, 2166t
classification of, 2168t
clinical features of, 2166–2167, 2167t
cutaneous, 363
extraglandular, 2167t
lymphoma, 2167
myelopathy, 2656
neuropathy, 2683
ocular, 199, 2166
pulmonary, 1714
renal, 64e-6, 1857, 1857t
trigeminal neuropathy, 2646
vasculitis, 2167, 2193
xerostomia, 241, 2166
definition of, 2166
diagnosis of, 2167, 2167t
differential diagnosis of, 2167, 2167t
epidemiology of, 2166
in HCV infection, 2041
in HIV infection, 1260–1261
HLA gene association with, 373e-8t
pathogenesis of, 372e-24–25t, 2166
in rheumatoid arthritis, 2138
in SLE, 2131
treatment of, 2167–2169, 2168f
- SJS. *See* Stevens-Johnson syndrome (SJS)
Skeletal dysplasias. *See* Chondrodysplasia(s)
Skeletal maturation, somatic growth and, 2257
- Skerljevo. *See* Endemic syphilis
Skew deviation, 210
Skin
actinically damaged, 356t, 357
age-related changes in, 387
in cardiovascular system examination, 1442–1443
color of, 386
complete examination of, 481, 482t
in cyanosis, 248
disorders/lesions. *See* Skin disorders/lesions
engineered tissue for replacement of, 92e-1, 92e-3t
Fitzpatrick classification of, 386, 386t
structural components of, 827, 827f
ultraviolet radiation effects on. *See* Sun exposure
- Skin biopsy
in lepromatous leprosy, 1126
in peripheral neuropathy, 2677–2678
technique for, 340
in vasculitic syndromes, 386e-1, 386e-6f
- Skin blister assay, in vivo, 423
Skin cancer, **493**
extramammary Paget disease, 502
genetic considerations in, 101e-2t
melanoma. *See* Melanoma
nonmelanoma, 76e-1, **500**, 502
basal cell carcinoma. *See* Basal cell carcinoma
Kaposi's sarcoma. *See* Kaposi's sarcoma
Merkel cell carcinoma, 502, 926, 928
squamous cell carcinoma. *See* Squamous cell carcinoma, skin
in transplant recipient, 388, 928
paraneoplastic syndromes in, 609t
prevention of, 502
screening for, 481, 482t
self-examination for, 476
sun/UV exposure and, 387–388, 476, 477t, 500
UV light therapy and, 348
- Skin disorders/lesions. *See also* Rash; *specific diseases*
approach to the patient, **339**
burrowing, 829
cancer. *See* Skin cancer
crusts, 827f, 828t, 829
descriptions of, 339t
diagnosis of, 340–344
distribution of, 341t, 342f
drug-induced. *See* Drug-induced illness; *specific drugs*
eschars, 828t, 829
esophageal manifestations of, 1911
history in, 340
immunologically mediated, **76e-7, 370**
induration, 2154, 2154t
infections, **827**. *See also specific infections*
anaerobic bacterial, 1099
in cancer patient, 484
diagnosis of, 831
pathophysiology of, 827, 827f
treatment of, 768t, 832–833, 832t
light-induced. *See* Photosensitivity diseases
as manifestations of internal/systemic disease, **76e-1, 76e-15, 353**. *See also specific diseases*
acneiform eruptions, 356, 356t
alopecia, 354–355, 355t. *See also* Alopecia
erythroderma, 353–354, 353t, 354t
exanthems, 362–363, 362t. *See also* Rash
figurate lesions, 355–356, 356t
hyperpigmentation, 359–361, 360t
hypopigmentation, 357–359, 357t, 358t
papulosquamous lesions, 353, 353t
purpura, 127, 367–369
pustular lesions, 356
in returned traveler, 802
telangiectasias, 356–357, 356t
urticaria, 363, 363t
morphology of, 341t
nodules, 828t, 829
papules, 828t
papulonodular lesions. *See* Papulonodular skin lesions
physical examination in, 340
purpura. *See* Purpura
shape of, 340
smallpox vaccination and, 352
telangiectasias. *See* Telangiectasia(s)
ulcers. *See* Cutaneous ulcers
vesicles/bullae. *See* Vesicles/bullae
- Skin snip, 245e-3
Skin tag (acrochordon), 341t, 342f, 359, 1958
Skull fractures, 457e-2
Slapped-check disease. *See* Erythema infectiosum
SLC01B1 gene, 38, 39t
SLC2A1 gene mutations, 653
SLC3A1 gene mutations, 332e-8
SLC4A1 gene mutations, 309, 653, 653t
SLC6A19 gene, 435e-2
SLC7A7 gene, 435e-2
SLC7A9 gene mutations, 332e-8
SLC12A3 gene mutations, 332e-8
SLC22A12 gene, 431e-5
SLC25A13 gene, 435e-2
SLC35C1 gene mutations, 419
SLC/TCA-4/Exodus 2, 372e-12t
SLE. *See* Systemic lupus erythematosus (SLE)
SLED (slow low-efficiency dialysis), 1811
Sleep, **184**
age-related changes in, 185, 185f
behavioral changes in, 187
insufficient, 188–189, 415e-3
neuroanatomy of, 185–186
neurochemistry of, 185–186
NREM, 184–185
organization of, 185
physiology of, 184–187
REM, 184–185, 185f
states and stages of, 184–185, 187
- Sleep apnea, **189**. *See also* Obstructive sleep apnea/hypopnea syndrome (OSAHS)
central, 189, 1727
complex, 1727
daytime sleepiness and, 188
mixed, 189
SA node dysfunction in, 1467, 1467t
sex differences in, 6e-5
- Sleep bruxism, 192
Sleep disorder(s), **187**
approach to the patient, 187–187
circadian rhythm, 193–194
daytime sleepiness in, 188, 188t
driving rights and, 188
fatigue and, 152
in fibromyalgia, 2239
at high altitudes, 476e-4
insomnia. *See* Insomnia
narcolepsy. *See* Narcolepsy
parasomnias, 192–193
in PD, 2618
periodic limb movement disorder, 192
restless legs syndrome, 192
in war veterans, 471e-5
in women, 6e-5
- Sleep enuresis, 192
Sleep hygiene, 65, 191, 191t
Sleeping sickness (human African trypanosomiasis), **1396**
cardiac manifestations of, 1560
clinical features of, 129t, 1397
diagnosis of, 245e-2, 245e-4t, 245e-6t, 1397–1398
epidemiology of, 245e-4t, 1397
parasites causing, 245e-4t, 1396, 1396t, 1397f
pathogenesis of, 1398
prevention of, 1398
skin manifestations of, 25e-12f, 129t, 1397

- Sleeping sickness (human African trypanosomiasis) (*Cont.*):
 treatment of, 1398, 1398t
 eflornithine, 246e-8-9
 melarsoprol, 246e-10
 suramin, 246e-13
- Sleep paralysis, 188t, 189
- Sleep terrors, 192
- Sleep-wake cycle, 185-186, 185f
- Sleepwalking, 192
- SLGT2 (sodium-glucose co-transporter 2)
 inhibitors, 2414t, 2415, 2416f
- Sliding filament model, for muscle contraction, 265e-5-6
- "Slip" (error), 12e-1
- Slipping clutch syndrome, 163
- Slow acetylators, 40
- Slow low-efficiency dialysis (SLED), 1811
- Slow reacting substance of anaphylaxis (SRSA), 372e-15t
- SLPI (secretory leukocyte protease inhibitor), 1223
- SMA (spinal muscle atrophy), 2631, 2634t, 2636
- SMAC, 102e-8-9
- SMAD4 gene mutations, 101e-1f, 101e-6t, 533
- SM-AHNMD (systemic mastocytosis associated with clonal hematologic non-mast cell lineage disease), 2120, 2120t
- Small airways disease. *See* Chronic obstructive pulmonary disease (COPD)
- Small intestine
 absorption and secretion in, 626
 benign tumors of, 536
 bleeding from, 277
 blood supply to, 1979, 1979f
 double-balloon enteroscopy of, 1881, 1883f
 endoscopy of, 1881. *See also* Capsule endoscopy
 fluid volumes and electrolyte content of, 98e-4t
 functions of, 1875, 1933
 ischemia of. *See* Intestinal ischemia
 length of, 1933
 malignant tumors of, 536-537
 motility of, 264-265
 mucosal biopsy of, 1938-1940, 1940t, 1941f
 neuroendocrine tumors of, 560t, 563, 567. *See also* Gastrointestinal neuroendocrine tumors (GI-NETs)
 nutrient digestion and absorption in, 1933-1934
 obstruction of, 103, 259, 1981. *See also* Intestinal obstruction
 physiology of, 264-265, 264t
 resection of, 96e-8
- Small-molecule inhibitors, 102e-9f, 102e-17
- Small noncleaved cell lymphoma. *See* Burkitt's lymphoma
- Smallpox
 in bioterrorism, 261e-6
 clinical features of, 131t, 261e-5t, 261e-6
 diagnosis of, 261e-5t, 1185
 differential diagnosis of, 134
 eradication of, 8, 785
 hemorrhagic, 261e-6
 incidence pre- and postvaccine, U.S., 785t
 malignant, 261e-6
 prevention of, 261e-5t, 261e-7, 261e-7t
 skin manifestations of, 25e-13f, 131t, 134, 261e-6, 829
 treatment of, 261e-5t, 261e-6
- Smallpox vaccine, 352
- Small round cell tumor, desmoplastic, 101e-3t
- Small subunit rRNA (SSU rRNA) genes, 86e-2
- Small-vessel stroke, 2565, 2567f. *See also* Ischemic stroke
- "Smart pumps," 12e-2
- Smell. *See* Olfactory disorders; Olfactory system
- Smith-Lemli-Opitz syndrome, 2324t, 2354t
- SMO gene mutations, 500, 500f
- Smoke inhalation, 1692
- Smokeless tobacco, 237, 240t, 476, 502
- Smoking. *See also* Smoking cessation
 cancer and, 2730, 2730t
 bladder, 575
 cervical, 594
 colorectal, 539
 esophageal, 475, 532-533
 head and neck, 502
 lung, 475, 506-507
 oral, 237
 pancreatic, 554
 renal cell carcinoma, 578
 skin, 500
- cardiovascular disease and, 2729-2730, 2730t
 global considerations in, 266e-4
 ischemic heart disease, 1587
 peripheral artery disease, 1644
 stroke risk, 2567t
- deaths related to, 2729
- drug interactions with, 2730, 2731t
- erectile dysfunction and, 326
- global considerations in, 17, 2729
- hemoglobin elevation and, 393
- IBD and, 1947
- in Leber hereditary optic neuropathy, 85e-9
- leukoplakia and, 240t
- of lower tar and nicotine cigarettes, 2731
- meningococcal infections and, 997
- neutrophil counts in, 418
- osteoporosis and, 2493
- passive exposure to
 in children, 2730
 COPD and, 1703
 lung cancer and, 475, 506, 2730
 respiratory disease and, 1662, 1693
- peptic ulcer disease and, 1917
- polycythemia in, 400
- in pregnancy, 2730, 2730t
- prevalence of, 2729
- prevention of, 2732
- respiratory disease and, 1662, 2730, 2730t
 asthma, 1680
 COPD, 1701, 1702, 1702f
 interstitial lung diseases, 1713
- rheumatoid arthritis and, 2140, 2141
- second malignancy and, 477
- in women, 6e-5
- Smoking cessation, 2731
 after lung cancer diagnosis, 506-507
 in COPD, 1705
 global considerations in, 17, 266e-4
 in ischemic heart disease treatment, 1587
 life expectancy increase following, 28t
 in peripheral artery disease, 1644
 pharmacologic therapy for, 507, 1705
 physician intervention in, 2731-2732, 2731t
 preoperative, 55t
 stages of, 476
 success rates of, 475
 in thromboangiitis obliterans, 1646
- Smoothed gene, 387
- Smooth muscle, vascular
 in atherosclerosis, 291e-2-3
 calcium in, 265e-3-4, 265e-3f
 contraction of, 265e-3-4, 265e-3f
 control of cell tone, 265e-4
 functions of, 265e-3-4
 structure of, 265e-1f
- Smudge cells, 81e-2, 702, 702f
- Snakebites, 2733
 clinical features of, 2733-2734, 2734f
 complications of, 2738
 epidemiology of, 2733
 field management of, 2734-2735
 global considerations in, 2738
 hospital management of, 2735-2738, 2736-2737t
 morbidity and mortality from, 2738
 snake identification, 2733
 wound infections following, 167e-2, 167e-3t
- Snellen chart, 195, 196f
- Snowshoe hare virus, 1309t
- SNPs. *See* Single-nucleotide polymorphisms (SNPs)
- SNRIs. *See* Serotonin norepinephrine reuptake inhibitors (SNRIs)
- SNRPN gene, 439, 2256
- SNRT (sinus node recovery time), 1469
- S_{obs} , 86e-4
- Sochi virus, 1307t
- Social distancing, 917
- Social history, 1
- Social jet lag, 194
- Social justice, 5
- Social media, 17e-3
- Social phobia, 2712
- SOD1, 2633, 2635
- Sodium
 blood pressure regulation and, 1612-1613
 in cardiac contraction, 265e-6
 in chronic kidney disease, 333e-3, 1814
 fractional excretion of, 291t
 hormonal regulation of, 332e-10-11, 332e-10f
 intake, hypertension and, 1612
 intestinal absorption of, 88
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 renal absorption/excretion of, 64e-2f, 296-297, 296f, 297f, 332e-3-6f, 332e-10-11, 332e-10f
 in specialized nutritional support formulas, 98e-5
 urine levels, in acute kidney injury, 291, 291t
- Sodium bicarbonate
 for acidosis after cardiac arrest, 1769-1770, 1769f
 adverse effects of, 1920
 for alcohol-induced acidosis, 64e-9
 for hyperkalemia, 312
 for prevention of contrast-induced nephropathy, 1461
 for uric acid nephropathy, 431e-5
 for ventricular arrhythmias in overdose/ poisoning, 473e-8t, 473e-9t
- Sodium channels
 cardiac, 273e-2, 273e-2f
 muscle, disorders of, 462e-17-18, 462e-17f, 462e-17t
 in seizure initiation, 2547
- Sodium citrate (Shohl's) solution, 64e-6, 318, 320
- Sodium current, in cardiac action potential, 273e-1f
- Sodium diarrhea, congenital, 268
- Sodium docusate, 62t
- Sodium ferric gluconate, 629
- Sodium-glucose co-transporter 2 (SLGT2)
 inhibitors, 2414t, 2415, 2416f
- Sodium hippurate, 969
- Sodium hypochlorite
 RSDL and, 262e-6
 for staphylococcal skin infections, 345
 for sulfur mustard exposure, 262e-4
- Sodium iodide, 2297
- Sodium iodide symporter (NIS), 372e-24t, 2284, 2284t
- Sodium ipodate, 2286t, 2297, 2300
- Sodium loading test, 2319
- Sodium modeling, in dialysis, 1822
- Sodium nitrite, 262e-7, 262e-8t
- Sodium phosphate enema, 62t
- Sodium polystyrene sulfonate (SPS), 312
- Sodium restriction
 for ascites, 288
 for hypertension, 1622t, 1623
- Sodium retention, 250, 251f
- Sodium stibogluconate, 246e-7, 1390, 1393
- Sodium sulfacetamide, 352
- Sodium thiosulfate, 262e-7, 262e-8t
- Sodium tyropanoate, 2286t
- Sodium valproate. *See* Valproate/valproic acid
- Sodoku (rat-bite fever), 129t, 167e-1-2
- Sofosbuvir, 215e-5t, 215e-6, 215e-10, 2047, 2049
- Soft (argasid) tick, 2745. *See also* Tick(s)
- Soft tissue, 827f
- Soft tissue infections. *See also* Cellulitis
 actinomycotic, 1090
 anaerobic bacterial, 1099
 deep. *See* Myositis; Necrotizing fasciitis
 enterococcal, 974

- in returning war veterans, 152e-9
treatment of, 832-833, 832t
- Soft tissue sarcoma, **119e-1**
classification of, 119e-1
deaths from, 470t
diagnosis of, 119e-1-2
epidemiology of, 119e-1
genetic considerations in, 119e-1
incidence of, 119e-1
metastatic, 119e-2
prognosis of, 119e-2
staging of, 119e-2, 119e-2t
treatment of, 103e-19, 119e-2
- SOI (severity-of-illness) scoring systems, 1729, 1730t
- SOJIA (systemic onset juvenile idiopathic arthritis), 372e-8t
- Solar elastosis, 386, 387
Solar purpura, 367
Solar radiation, 385
Solar urticaria, 363, 363t, 2118
Solid organ transplantation. *See also* Transplant recipient
brain injury after, 463e-3
heart. *See* Cardiac transplantation
infections after, 923-924, 924t, 925t
kidney. *See* Kidney transplantation
liver. *See* Liver transplantation
lung. *See* Lung transplantation
- Solifenacin, 81
Solitary rectal ulcer syndrome, 277, 1957
Soluble defense collagens (collectins), 145e-5
Solute diuresis, 295
Somali famine, 151e-3
Soman. *See* Nerve agents
Somatic hypermutation, 372e-20, 372e-21
Somatic mosaicism, 439
Somatic mutations
cancer development and, 101e-1, 101e-1f, 440
in mitochondrial DNA, 439
Somatic symptom disorder, **2718**
Somatization, 254
Somatosensory cortex, 88, 89f, 176f
Somatosensory evoked potentials (SSEPs), 442e-3-4, 1781
Somatosensory pathways, 159f
Somatostatin
actions of, 401e-2, 570, 1912
half-life of, 400e-2
structure of, 566f
synthesis of, 401e-3
Somatostatin analogues
for acromegaly, 2270f, 2271
adverse effects of, 574, 1853, 2271
for carcinoid syndrome, 272, 567
for Cushing's syndrome, 2272-2273
for nausea and vomiting, 260t
for polycystic kidney disease, 1853
for TSH-secreting adenomas, 2274
for VIPoma, 571
Somatostatinoma, 269, 559t, 561, **570**
Somatostatin receptor scintigraphy (SRS), 571-572, 572f
Somatotropin, 2258t
Somnambulation, 192
Sorafenib
action and targets of, 101e-8, 102e-3, 102e-5t, 102e-17, 103e-16f, 103e-18t, 103e-19
adverse effects of, 103e-18t, 103e-19, 103e-26, 380, 381f, 551
for cholangiocarcinoma, 553
for hepatocellular carcinoma, 102e-3, 102e-17, 550, 551, 551t, 552
for renal cell carcinoma, 102e-3, 102e-5t, 102e-17, 579
for thyroid cancer, 2307
Sorbitol, 1970t, 2743
Sotalol
action of, 273e-6t
adverse effects of, 1480t, 1498
for arrhythmias, 1480t
for atrial fibrillation, 1489
for atrial flutter, 1485
dosage in kidney disease, 37
dosage monitoring, 36
drug interactions of, 43
for focal atrial tachycardia, 1479
overdosage/poisoning with, 473e-9t
pharmacology of, 1480t
for ventricular arrhythmias, 1498
Sour taste sensation, 212
Southeast Asia ovalocytosis, 653, 653t
Southern tick-associated rash illness (Master's disease), 129t, 134, 1152
Sowdah, 1420
Sox2 gene, 454, 455
SOX3 gene mutations, 2325t
SOX9 gene, 2349, 2351f
SOX9 gene mutations
in disorders of testis development, 2353, 2354t
in testicular/ovotesticular disorders of sex development, 2355, 2356t
SOX-10, 372e-24t
SOX-10 gene mutations, 359
SOX18 gene mutations, 1654
Sp110 deficiency, 375e-2, 2110
SPARC, 554
Sparganosis, 1435
Spasmodic dysphonia, 2620
Spasms
hemifacial, 2648
in MS, 2663, 2672
muscle, 462e-2-3
in tetanus, 985, 986
Spasticity, 154, 163, 2660
Spastin gene, 2634t, 2636
Spatial orientation, 180-182, 180f
Specialized nutritional support, **98e-1**
approach to the patient, 98e-1-4
benefits of, 98e-3-4
decision-making for implementation of, 98e-2f
design of regimens for
amino acids, 98e-5
energy requirements, 98e-4-5
fluid requirements, 98e-4, 98e-4t
minerals and vitamins, 98e-5, 98e-5t, 98e-6t
monomeric formulas, 98e-5
proteins, 98e-5
disease-specific, 98e-4
enteral. *See* Enteral nutrition
home, 98e-4
hyperglycemia and, 98e-4
indications for, 98e-1-3, 98e-2f
parenteral. *See* Parenteral nutrition
risks of, 98e-3-4
Specific granule deficiency, 419, 419t
Specificity
analytical, 480e-2
in immune response, 372e-2
of tests, 21, 22, 479, 479t
SPECT (single photon emission computed tomography)
for myocardial perfusion imaging. *See* Myocardial perfusion imaging
in parathyroid surgery, 2474
radiopharmaceuticals for, 270e-3, 270e-3t
Spectinomycin
action of, 930
for gonococcal infections, 1008, 1009t
resistance to, 1004-1005
Speech
assessment of, 37e-1v, 2537
spontaneous, 177
Speech audiometry, 222
Speech reception threshold, 222
Sperm, mutations in, 433
Spermatogenesis
abnormalities of, genetic considerations in, 433, 437
process of, 2358f, 2359-2360
radiation-induced effects on, 623
Spermatogonial stem cells, 453t, 454
Sperm banking, 103e-25, 592, 623
Spermidine, 94e-6f, 94e-7
S-phase analysis, in breast cancer, 528
Spherocytosis, 81e-3f, 281
Spherules, 1335, 1335f
Sphincter of Oddi, 2076, 2082-2083
Sphincteroplasty, 1976
Sphincterotomy, endoscopic, 1881, 1884f
Spider angioma, 286, 1992, 2058, 2059f
Spider bites, 365, 2747
Spina bifida occulta, 118
Spinal accessory nerve, 2538
Spinal anesthesia, 54
Spinal angiography, 440e-10-11
Spinal cord
age-related changes in, 76
anatomy of, 2650-2651, 2652f
compression of. *See* Spinal cord compression disorders of. *See* Spinal cord disease/injury
pain mechanisms of, 88, 89f
Spinal cord compression
in hematomyelia, 2655
IgG4-related, 391e-1t
neoplastic
clinical features of, 1790-1791, 2653
diagnosis of, 1791, 1791f, 2653, 2653f
in intradural mass lesions, 2654, 2654f
in lung cancer, 513
metastatic, 1790, 2653
pathogenesis of, 2653
in primary intramedullary tumors, 2654, 2654f
treatment of, 1791-1792, 1791f, 2653-2654
in spinal epidural abscess, 780t, 784, 2654-2655, 2655f
in spinal epidural hematoma, 2655
Spinal cord disease/injury, **2650**
acute, 2653
anatomic considerations in, 2650-2651, 2651t, 2652f
approach to the patient, 2650-2653, 2651t
autonomic dysfunction in, 2641-2642, 2660
cervical. *See* Cervical spine
compressive. *See* Spinal cord compression
constipation in, 274
erectile dysfunction in, 326
in HIV infection, 1266-1267
infarction, 441e-25f, 2655-2656
infectious. *See* Meningitis
level of lesion in, 2540t, 2651-2652, 2651t
myelopathies. *See* Myelopathy
neurogenic shock in, 1750
paraneoplastic syndromes, 618
patterns of, 2653
prognosis of, 2660t
radiation-induced, 103e-4, 622
referred abdominal pain from, 105
remyelination after, 90e-3
sensory abnormalities in, 161
stem cell therapy for, 90e-3
subacute, 2653
treatment of, 2659-2660
vascular malformations, 2657, 2658f
Spinal epidural abscess
clinical features of, 117, 784, 2654
diagnosis of, 2654, 2655f
etiology of, 784, 2654
risk factors for, 2654
treatment of, 780t, 2655
Spinal epidural hematoma, 2655
Spinal fusion, 120
Spinal manipulative therapies, 14e-1t, 14e-2-3, 119
Spinal muscle atrophy (SMA), 2631, 2634t, 2636
Spinal osteomyelitis. *See* Osteomyelitis, vertebral
Spinal stenosis, 115-116, 116f
Spindle cell tumor, 372
Spine
anatomy of, 111, 112f
cervical. *See* Cervical spine
lumbar
congenital anomalies of, 118
disk disease in, 113f, 114-115, 115f
disk replacement in, 120
fusion surgery in, 120
pain referred to, 118
pain sensitive structures in, 111
sacral, 118
SPINK1 gene mutations, 2092, 2099, 2101

- Spinobulbar muscular atrophy (Kennedy's disease)
vs. ALS, 2636
clinical features of, 2366, 2636
genetic considerations in, 431t, 434, 440t, 2634t, 2636
- Spinocerebellar ataxia (SCA)
channelopathies in, 444e-2t
classification of, 451e-1, 451e-1-4t, 2627
clinical features of, 2627-2629
genetic considerations in, 431t, 451e-1-2t, 2627-2629
infantile-onset, 451e-4t
with neuropathy (SCAN1), 451e-4t
SCA1, 206, 440t, 451e-1t, 2627-2628, 2628f
SCA2, 440t, 451e-1t, 2628
SCA3 (Machado-Joseph disease), 440t, 451e-1t, 2628-2629
SCA4, 451e-1t
SCA5, 451e-1t
SCA6, 440t, 451e-1t, 2629
SCA7, 440t, 451e-1t, 2629
SCA8, 451e-1t, 2629
SCA10, 451e-1t
SCA11, 451e-1t
SCA12, 440t, 451e-1t
SCA13-36, 451e-1t
treatment of, 2630
- Spinothalamic tract, 88, 89f, 158, 159f
- Spiny sea urchins, 2739-2740, 2741f
- Spiral enteroscopy, 1881, 1883f
- Spiramycin, 246e-5t, 246e-13, 1404
- Spirillum* infection, 167e-1, 167e-2
- Spirillum minus*, 129t
- Spirometra*, 1435
- Spirometry
in asthma, 1675, 1675f
in COPD, 1701, 1704, 1704t
in interstitial lung diseases, 1710
preoperative, 55, 55t
in respiratory disease evaluation, 306e-2f, 306e-6, 1662
- Spirolactone
action of, 332e-9
adverse effects of, 1623
erectile dysfunction, 327
gynecomastia, 288
hyperkalemia, 43, 64e-2, 288, 335
hyponatremia, 288
hypotension, 335
for ascites, 288, 2065
drug interactions of, 43
for heart failure, 1511, 1512t
for hirsutism, 335
for hypertension, 1617, 1623, 1624t
for hypokalemic periodic paralysis, 462e-17
for male precocious puberty, 2362
for mineralocorticoid excess, 2319
for polycystic ovarian syndrome, 337
for syndrome of apparent mineralocorticoid excess, 307
- Spleen
accessory, 409
congenital absence of. *See* Asplenia
infarction of, 106t, 410
palpable, 410-411
percussion of, 411
rupture of, 106t, 410, 413, 1188
structure and function of, 409-410, 410f
- Splenectomy, 411
for autoimmune hemolytic anemia, 658
for CML, 694
for congenital erythropoietic porphyria, 2532
for hemoglobinopathies, 637
for hereditary spherocytosis, 652
for Hodgkin's lymphoma staging, 412
for immune thrombocytopenic purpura, 728
immunizations after, 635, 652, 788f, 929
infections after, 484, 652, 780t, 781
for leukemia, 412
long-term effects of, 413
for lymphoma, 412
for myelofibrosis, 676
for polycythemia vera, 674
for sickle cell anemia, 635
for splenic abscess, 851
for thalassemia, 638
for thrombotic thrombocytopenic purpura, 730
- Splenic abscess
abdominal pain in, 105, 106t, 851
actinomycotic, 1089, 1090f
clinical features of, 851
etiology of, 851
in infective endocarditis, 826
treatment of, 851
- Splenic marginal zone lymphoma, 135e-1, 135e-2t, 706
- Splenic sequestration crisis, in sickle cell anemia, 634
- Splenic vein thrombosis, 2097
- Splenomegaly, 409
abdominal swelling in, 286
approach to the patient, 410-411
clinical features of, 410-411
in CML, 689
congestive, 411
differential diagnosis of, 411
diseases associated with, 412t
in Gaucher disease, 432e-5
in hemolytic anemia, 399
in hepatocellular carcinoma, 545
in liver disease, 1992
with lymphadenopathy, 408
massive, 411, 412t
in myelodysplasia, 670
in myelofibrosis, 676
in polycythemia vera, 399, 673-674
in portal hypertension, 2064
in sarcoidosis, 2208, 2208f
in schistosomiasis, 1426f, 1427
tropical, 1374-1375
in visceral leishmaniasis, 1389, 1390f
- Splenosis, 413
- Splinter hemorrhages, 1443
- Spondylitic myelopathy, 2657
- Spondylitis
ankylosing. *See* Ankylosing spondylitis
tuberculous. *See* Pott's disease
- Spondyloarthritides, 372e-15t, 2169. *See also specific disorders*
- Spondyloarthritis
axial, 2169, 2169t
criteria for, 2178, 2178t
juvenile-onset, 2178
undifferentiated, 2178
- Spondylodiskitis. *See* Osteomyelitis, vertebral
- Spondylolisthesis, 117
- Spondylolysis, 118
- Spondylosis, 117, 122, 163
- Sponge diver's disease, 2739
- Spontaneous breathing trial, 1743-1744, 1744f
- Spontaneous speech, 177
- Sporadic goiter, 2301
- Spore, anthrax, 261e-2
- Sporogony, 1371
- Sporothrix* infections, 1353
clinical features of, 1353-1354
articular, 836
chronic meningitis, 908t
cutaneous, 364t, 366, 829, 1353, 1353f
diagnosis of, 1354
etiology of, 1353
pathogenesis of, 1353
treatment of, 1354, 1354t
- Sporothrix schenckii*, 1353, 1354
- Sporozoites, *Plasmodium*, 1369, 1369f
- Sports. *See* Athlete(s)
- Spousal abuse. *See* Domestic abuse/violence
- Sprouting, 102e-13-14. *See also* Angiogenesis
- SPS (sodium polystyrene sulfonate), 312
- SPTA1* gene mutations, 653t
- SPTB* gene mutations, 653t
- SPTLC1* gene mutations, 2643
- Spur cell(s), 396f
- Spur cell anemia, 282
- Spurling's sign, 122
- Sputum analysis
in cough, 244
Gram's stain, 806, 1085f
in lung cancer, 512
in pneumonia, 806
in respiratory disease evaluation, 1663, 1667
specimen collection and transport, 150e-4t, 1667
in tuberculosis, 1114
SQ109, 205e-7
- SQSTM1* gene, 426e-1
- Squamous cell carcinoma, 76e-6f
bladder, 575. *See also* Bladder cancer
cervix. *See* Cervical cancer
endometrium. *See* Endometrial cancer
esophagus, 532-533, 532t. *See also* Esophageal cancer
head and neck region, 503. *See also* Head and neck cancer
lung, 507, 507f. *See also* Lung cancer
nasal and sinus, 226
oral, 237, 239t
skin
clinical features of, 76e-6f, 341t, 500-501, 501f
distribution of, 342f
epidemiology of, 500
genetic considerations in, 500
infections in, 484t
metastatic, 35
natural history of, 501
pathophysiology of, 500
pre-malignant forms, 35
prevention of, 502
risk factors for, 500
screening for, 481
sun exposure and, 387, 500
in transplant recipient, 500
treatment of, 102e-5t, 502
of unknown primary, 120e-3f, 120e-4
- Square root sign, 1576
- SQUID (super-conducting quantum-interference device), 639
- SRD5A2* gene mutations, 2353, 2354t, 2366
- SRSA (slow reacting substance of anaphylaxis), 372e-15t
- SRY* gene mutations
in disorders of testis development, 437, 439, 2349, 2353, 2354t
in testicular/ovotesticular disorders of sex development, 2355, 2356t
transcription factor in, 431t
- SSc sine scleroderma, 2155. *See also* Systemic sclerosis (scleroderma)
- SSEPs (somatosensory evoked potentials), 442e-3-4
- SSKI (saturated solution of potassium iodide), 1354, 1354t
- SSPE (subacute sclerosing panencephalitis), 900, 1298
- SSRIs. *See* Selective serotonin reuptake inhibitors (SSRIs)
- SSU rRNA (small subunit rRNA) genes, 86e-2
- Stabbing headache, primary, 2597-2598
- Stab phlebectomy, 1653
- Staghorn calculi, 1871
- Stagnant bowel syndrome. *See* Bacterial overgrowth syndrome
- Staining, 150e-1
- Standard precautions, 916
- Stanford V regimen, for Hodgkin's lymphoma, 709
- Stapedectomy, 217, 223
- Stapes, 217, 218f
- Staphylococcal infections, 954
aortitis, 1642
in cancer patient, 484t, 486, 486t, 487t
catheter-related bloodstream, 915
cellulitis, 830
clinical syndromes, 960
diagnosis of, 956, 960
in dog-bite wound, 167e-1
endocarditis. *See* Infective endocarditis, staphylococcal
food-borne, 856, 856t, 956, 959

- meningitis. *See* Meningitis
- osteomyelitis, 839, 840t, 841t, 844
- pathogenesis of, 960
- of prosthetic devices, 960
- S. aureus*. *See* *Staphylococcus aureus* infections
- surgical-site, 915
- toxic shock syndrome. *See* Toxic shock syndrome (TSS), staphylococcal
- in transplant recipient, 919, 926
- treatment of, 940t, 960–963, 962t
- Staphylococcal scalded skin syndrome
- clinical features of, 131t, 959
- epidemiology of, 959
- pathogenesis of, 956
- vs. scarlet fever, 134
- skin manifestations of
- generalized desquamation, 25e-7f, 959
- Nikolsky's sign, 134, 959, 960f
- progression, 362
- rash, 131t, 134, 362t, 363
- vesicles/bullae, 361t, 362
- vs. toxic epidermal necrolysis, 134, 362, 829
- Staphylococcal α toxin, 956
- Staphylococcus aureus*
- adherence of, 145e-3
- antibiotic resistance in, 941t
- antimicrobial susceptibility testing, 150e-8
- β -lactam-resistant, 933
- daptomycin-resistant, 934
- evasion of host defenses by, 956
- genome of, 955
- laboratory identification of, 954, 954f, 956
- pathologic response to, 954
- phagocyte interaction with, 145e-7
- tissue invasion by, 145e-7
- vancomycin-resistant, 933, 961
- virulence regulation in, 955
- Staphylococcus aureus* infections
- in atopic dermatitis, 345
- burn-wound, 166e-1
- in cancer patient, 486, 486t, 487t
- CAPD peritonitis, 848
- catheter-related bloodstream, 915–916
- clinical syndromes, 957t
- deep neck, 235
- in diabetes mellitus, 2429
- diagnosis of, 150e-8
- endocarditis. *See* Infective endocarditis, staphylococcal
- epidemiology of, 954–955
- epiglottitis, 234
- food-borne
- clinical features of, 266t, 856, 856t
- diagnosis of, 959
- differential diagnosis of, 266t, 959
- pathogenesis of, 956
- sources of, 266, 856t, 959
- host response to, 956
- in human bite-wound, 167e-2
- in leukocyte adhesion deficiency, 419
- mastoiditis, 231
- meningitis. *See* Meningitis, bacterial
- methicillin-resistant. *See* Methicillin-resistant *Staphylococcus aureus*
- musculoskeletal, 957–958
- osteomyelitis, 839, 844, 845, 957, 957f
- pyomyositis, 831, 958
- septic arthritis, 834, 957–958. *See also* Infectious (septic) arthritis
- nausea and vomiting in, 258
- otitis externa, 228
- parotitis, 241
- pathogenesis of, 955–956
- perichondritis, 228
- prevention of, 959–960
- prosthetic device-related, 959
- psoas abscess, 852
- respiratory tract
- after influenza, 1212
- community-acquired, 958
- health care-associated, 958
- hemoptysis in, 245
- lung abscess, 814
- risk factors for, 955
- sinusitis, 226
- skin and soft tissue, 956–957
- cellulitis, 830
- folliculitis, 356, 829, 957
- furunculosis, 349, 957
- impetigo, 349, 350t, 967. *See also* Impetigo mastitis, 957
- staphylococcal scalded skin syndrome. *See* Staphylococcal scalded skin syndrome
- splenic abscess, 851
- surgical-site, 915
- toxic shock syndrome. *See* Toxic shock syndrome, staphylococcal
- in transplant recipient, 919, 926
- treatment of
- antimicrobial agent selection, 940t, 961–963, 962t
- empirical, 963
- endocarditis, 962
- osteomyelitis/septic arthritis, 962
- principles, 960
- therapy duration, 961
- toxic shock syndrome, 962
- urinary tract, 914, 959
- ventilator-associated pneumonia, 915
- Staphylococcus epidermidis*, 960
- Staphylococcus epidermidis* infections, 98e-7–8, 229, 960
- Staphylococcus lugdunensis* infections, 817, 960
- Staphylococcus saprophyticus* infections, 862, 960
- Staphylococcus schleiferi* infections, 960
- Staphylococcus suis*, 971
- STAR (steroidogenic acute regulatory protein), 400e-2, 2353, 2354t
- Starfish envenomation, 2740, 2740f
- STARI (Southern tick-associated rash illness), 129t, 134, 1152
- Starling forces, 250, 295
- STARR (stapled transanal rectal resection) procedure, 1974, 1975f
- Starvation-related malnutrition. *See* Marasmus (starvation-related malnutrition)
- Stasis dermatitis, 346
- clinical features of
- distribution and morphology, 341t, 342f
- erythema and crusting, 76e-2f, 346, 354t
- evolution of, 346f
- treatment of, 346–347, 354t
- Stasis ulcer, 342f, 346–347, 346f
- STAT1 gene mutations, 1129
- (Lif)/Stat3 (leukemia inhibitory factor), 455
- STAT3 deficiency. *See* Hyper-IgE syndrome
- STAT4 gene mutations, 2125
- STAT5, 89e-4
- STAT5b gene, 375e-2
- STAT factors, 102e-4f, 145e-7, 372e-16, 401e-3
- Statins
- adverse effects of, 2448t
- hepatotoxicity, 2023, 2030, 2448, 2453
- myopathy, 462e-19, 462e-20t, 2199, 2218t, 2243, 2447–2448
- tendinitis, 2218t
- anti-inflammatory effects of, 291e-9, 291e-9f
- for colorectal cancer prevention, 478
- for dementia prevention, 2602
- drug interactions of, 42, 96e-4, 246e-3t, 2199, 2447–2448
- genetic variations in response to, 42
- for intermittent claudication, 1644
- LDL-lowering effect of, 291e-6, 291e-7f, 291e-9, 291e-9f
- for lipoprotein disorders
- ACC/AHA guidelines, 291e-5–6, 291e-6t, 291e-7f, 291e-8f
- adverse effects of, 2448t
- approach to the patient, 2447–2448
- dosage of, 2448t
- in ischemic heart disease, 1588
- in metabolic syndrome, 2453
- metabolism of, 34t
- for NSTEMI-ACS, 1596, 1598
- perioperative prophylactic, 54
- for prostate cancer prevention, 580
- for stroke prevention, 2569
- Status epilepticus, 2557
- complex partial, 2607
- in critically ill patient, 1735, 1782
- definition of, 2557
- delirium in, 169
- EEG in, 442e-2f
- generalized tonic-clonic, 2557, 2558f
- in nerve agent exposure, 262e-5, 262e-6–7
- nonconvulsive, 2557
- treatment of, 2558f
- Staufer's syndrome, 284, 578
- Stavudine (d4T), 1276f, 1279f, 2030
- Steady state, 34, 35f, 36
- Steatohepatitis, nonalcoholic. *See* Nonalcoholic steatohepatitis (NASH)
- Steatorrhea
- in bacterial overgrowth syndrome, 1945
- in chronic pancreatitis, 2100–2101
- detection of, 1938
- etiology of, 269, 1932–1933, 1935–1936, 1941t
- lipid digestion and absorption defects in, 1935, 1935t
- pancreatic exocrine function assessment in, 1940
- in protein-losing enteropathy, 1945
- in short-bowel syndrome, 1943
- in Whipple's disease, 269, 1945
- STEBP1 gene mutations, 135e-5
- STEC. *See* Shiga toxin-producing *E. coli* (STEC)
- Steele-Richardson-Olszewski syndrome. *See* Progressive supranuclear palsy syndrome
- Steiner's disease, 375e-2
- Stem cell(s)
- biology of, 453
- cancer. *See* Cancer stem cells
- cell division in, 89e-1, 454
- cultured, 453t
- directed differentiation of, 455
- facultative, 454–455
- functions of, 89e-1, 89e-1f
- gene regulation of, 454
- genomics and proteomics of, 455
- genotype/phenotype stability of, 454
- hematopoietic. *See* Hematopoietic stem cells
- imaging of, 90e-3
- immortality of, 454
- isolated, 453t
- mitochondrial DNA mutations, 85e-10
- neurologic, 444e-7
- niche site, 89e-2
- nuclear reprogramming of, 454
- plasticity of, 454–455
- pluripotent, 413
- potency of, 453, 454f
- proliferation of, 454
- quiescence of, 89e-2
- relationship to cancer, 89e-4
- resident, 453
- self-renewal of, 89e-2–4, 89e-4f, 453–454
- source developmental stage of, 454f
- in tissue engineering, 92e-1
- totipotent, 454
- transdifferentiation of, 90e-2, 454–455
- unipotent, 454
- Stem cell factor (SCF), 372e-11t
- Stem cell therapy, 90e-1
- for diabetes mellitus, 90e-2–3
- ethical issues in, 90e-4
- for Huntington's disease, 444e-7–8
- for ischemic heart disease, 90e-2
- for liver disease, 90e-3
- for neurologic disorders, 90e-3
- for PD, 444e-7, 2617
- sources of cells for, 90e-1–2, 444e-7
- strategies for, 90e-1, 90e-1f
- STEMI. *See* ST-segment elevation myocardial infarction (STEMI)
- Stemmer's sign, 1654
- Stenotrophomonas maltophilia* infections
- antibiotic resistance in, 811
- in catheter-related infections, 487, 1048
- clinical features of, 1048
- diagnosis of, 150e-2t

- Stenotrophomonas maltophilia* infections (Cont.):
 pathogenesis of, 1042, 1048
 treatment of, 1045t, 1048
 in ventilator-associated infections, 1048
- Stent(s)
 for aortic aneurysm, 1640, 1640f
 coronary artery. *See also* Percutaneous coronary interventions (PCI)
 characteristics of, 296e-1-2, 296e-1f
 drug-eluting, 296e-2, 296e-3f, 747, 1591
 restenosis in, 296e-2-3, 297e-5, 297e-5f, 297e-5v, 1591
 for pancreatic cancer-related obstruction, 1892, 1894f
 for peripheral artery disease, 1645
 for renal artery stenosis, 1618, 1629
- Stereognosis, 160, 2539
- Stereopsis, 197
- Stereotactic radiation therapy, 517
- Stereotactic radiosurgery, 1792, 2263
- Stereotyping, 16e-4, 16e-6
- Sterilization, 2390t
- Sternal osteomyelitis, 844-845, 844f
- Sternoclavicular joint, infectious arthritis of, 834, 834f
- Steroid(s). *See specific hormones*
- Steroidogenesis. *See specific hormones and glands*
- Steroidogenic acute regulatory protein (StAR), 400e-2, 2353, 2354t
- Steroidogenic enzyme pathways, disorders of, 2353, 2354t, 2355f
- Steroidogenic factor 1 (SF1), 2310, 2349
- Steroid purpura, 367
- Stevens-Johnson syndrome (SJS)
 in cancer patient, 487
 clinical features of, 131t, 361t, 362, 382-383, 384t
 differential diagnosis of, 382-383
 drugs associated with, 383
 epidemiology of, 131t
 etiology of, 131t, 362
 genetic considerations in, 378
 in HIV infection, 1263
 immune pathways of, 378t
 in *M. pneumoniae* infection, 1164
 nondrug causes of, 384t
 oral manifestations of, 46e-1f, 239t
 prevention of future episodes, 384
 treatment of, 383
- Stevens-Johnson syndrome/toxic epidermal necrolysis overlap, 382
- STG11 genes, 102e-7
- Stickler's syndrome, 2511
- Stiff-person syndrome
 clinical features of, 462e-3
 gait disorders in, 163
 paraneoplastic, 614t, 618
 pathogenesis of, 372e-25t, 462e-3
- Still's disease
 adult-onset, 372e-8t
 clinical features of, 25e-3f, 129t, 134, 364
 treatment of, 141
- Still's murmur, 51e-4
- Stinging ants, 2749
- Stingray injury, 2740
- STIs. *See Sexually transmitted infections (STIs)*
- STK11 gene mutations, 554
- Stokes-Adams attack, 146
- Stomach
 acid and hormone secretion in, 1912-1914, 1914f
 biopsy in cobalamin deficiency, 645
 epithelial anatomy of, 1911-1912, 1911f, 1912f
 fluid volumes and electrolyte content of, 98e-4t
 functions of, 1875
 mucosal defense and repair in, 1912, 1913f
- Stomach tumors
 adenocarcinoma. *See* Gastric adenocarcinoma
 lymphoma, 535-536
 sarcoma, 536. *See also* Gastrointestinal stromal tumor
- Stomatitis, 342f, 1946t
- Stomatocytes, 81e-1
- Stomatocytosis, 81e-4f, 653f, 653t
- Stonefish sting, 2741-2742
- Stony coral injury, 2741
- Stool
 anion concentrations in, 1933
 electrolytes in, 1933
 fasting effects on, 1933
 microbiome of, 86e-6, 86e-6f, 86e-9
 osmolality of, 1933
 retained, abdominal swelling and, 285
 vomiting of, 259
- Stool analysis
 antigen test, for *H. pylori*, 1040, 1040t
 in chronic diarrhea, 272
 for intestinal parasite detection, 245e-1
 for occult blood. *See* Fecal occult blood in pancreatic disease evaluation, 2087t, 2090
 specimen collection and transport for, 150e-4t
- Stool-bulking agents, 1969
- Stool culture, 267
- Stool softeners, 61, 62t
- STOP-Bang survey, 1721
- Strategic National Stockpile, 261e-9
- Stop TB Strategy, 1122
- Storage pool disorder, platelet, 730
- Straight back syndrome, 1443
- Straight-leg raising sign, 112-113
- Strategic National Stockpile, 261e-9
- Strawberry gallbladder, 2083
- Strawberry gums, 237
- Strawberry tongue, 241t, 362, 966
- Strength testing, 2538
- Streptobacillus moniliformis*, 130t, 167e-1-2
- Streptocerciasis/*Mansonella streptocerca*, 245e-6t, 1418t, 1422
- Streptococcal infections. *See also* Group A *Streptococcus* (GAS) infections; Group B *Streptococcus* (GBS) infections
 aortitis, 1642
 brain abscess. *See* Brain abscess
 in burn-wounds, 166e-1
 in cancer patient, 484, 484t, 486t
 deep neck, 235
 in dog-bite wound, 167e-1
 endocarditis. *See* Infective endocarditis, streptococcal
 global considerations in, 1529
 glomerulonephritis following. *See* Poststreptococcal glomerulonephritis
 in human bite-wound, 167e-2
 meningitis. *See* Meningitis
 osteomyelitis, 839, 840t, 841t
 skin infections, 829
 toxic shock syndrome. *See* Toxic shock syndrome (TSS), streptococcal
 in transplant recipient, 919, 920t
- Streptococci
 anaerobic, 971. *See also* Anaerobic bacteria
 characteristics of, 963, 964t
 classification of, 963, 964t
 laboratory identification of, 150e-2t
 nutritionally variant, 971
 viridans, 940t, 963, 964t, 971
- Streptococcus agalactiae*, 963, 964t, 969. *See also* Group B *Streptococcus* (GBS) infections
- Streptococcus bovis* infections. *See* Group D *Streptococcus* infections
- Streptococcus iniae*, 971
- Streptococcus milleri*, 971
- Streptococcus mutans*, 236
- Streptococcus pneumoniae*. *See also* Pneumococcal infections
 antibiotic resistance in, 807, 933, 941t, 949, 952
 immune response to, 949-950
 laboratory identification of, 946, 946f
 molecular typing of, 947
 nasopharyngeal carriage of, 948, 948f
 serotype distribution of, 947-948, 948f
 tissue invasion by, 145e-9
 virulence factors in, 947, 947f
- Streptococcus pyogenes*
 cell-surface components, 964
 extracellular products of, 965
- M protein types of, 964
 phagocyte interaction with, 145e-7, 965
- Streptococcus pyogenes* infections. *See* Group A *Streptococcus* infections
- Streptococcus suis*, 971
- Streptogramins. *See* Quinupristin-dalfopristin
- Streptokinase, 758, 758f, 1605
- Streptolysin O, 145e-7
- Streptomycin
 action of, 205e-5, 930
 adverse effects of, 203, 205e-5, 821, 942
 for *Brucella* infections, 194e-4
 in developing countries, 205e-5
 dosage of, 205e-5
 indications for, 942
 for infective endocarditis, 821
 pharmacology of, 205e-5
 for plague, 261e-5t, 1074, 1074t
 in pregnancy and lactation, 937t
 resistance to, 205e-5, 934, 976
 for TB, 205e-2t, 205e-5, 1115
 for tularemia, 261e-5t, 261e-7, 1070
 for verruga peruana, 1081t
- Streptozotocin, 103e-25, 474, 574, 2337
- Stress disorders, 2712. *See also* Posttraumatic stress disorder (PTSD)
- Stress incontinence, 80-81
- Stress testing
 ECG. *See* Electrocardiography (ECG), stress
 echocardiography. *See* Echocardiography, stress
 myocardial perfusion imaging. *See* Myocardial perfusion imaging
 in peripheral arterial disease, 1643
 risk stratification using, 270e-14, 270e-14f
- Stress ulcers
 in critically ill patient, 1734
 in mechanically ventilated patient, 1743
 in sepsis/septic shock, 1755
- Stridor, 1662
- "String sign," 1953, 1954
- Stroke, 2559. *See also* Cerebrovascular disease; Intracranial hemorrhage; Ischemic stroke
 approach to the patient, 2560, 2560f
 atrial fibrillation and, 1487-1488
 carotid radiation and, 621
 circadian variations in, 194, 1616
 clinical features of, 2559
 aphasia, 178, 179-180
 coma, 1775-1776
 delirium, 169
 depression, 2714
 dysphagia, 256
 gait disorders, 163
 ocular, 204, 211
 seborrheic dermatitis, 347
 weakness, 156
 in critically ill patient, 1735
 deaths from, 56t, 266e-3t
 definition of, 2559
 dental care after, 242
 etiology of, 2560
 fatigue following, 152
 global considerations in, 266e-3
 at high altitudes, 476e-4
 in HIV infection, 1266
 hypokalemia and progression of, 307
 hypothermia in, 478e-1
 imaging in, 2580-2581
 inflammatory mediators in, 372e-29t
 postmenopausal hormone therapy and, 2384t, 2385
 in pregnancy, 49
 risk factors for, 2567t, 2569
 treatment of, 758, 1627, 2560, 2560f
- Stroke centers, 2563
- Stroke volume
 calculation and normal values, 1748t
 determinants of, 265e-8, 265e-9f, 265e-9t
 in shock, 1746
- Stromal cell-derived factor (SDF), 372e-12t, 1235
- Stromal tumors, ovarian, 594

- Strongyloides stercoralis*/strongyloidiasis
 adult T-cell leukemia/lymphoma and, 225e-4
 clinical features of, 152e-5t, 1414t, 1415
 diagnosis of, 152e-5t, 245e-1, 245e-3t, 245e-5t, 245e-6t, 1416
 eosinophilia in, 245e-6t, 765t, 1687
 epidemiology of, 152e-8, 245e-3t, 765t, 1415
 incubation period for, 152e-5t
 life cycle of parasite, 245e-3t, 1414t, 1415, 1415f
 in returning war veterans, 152e-5t, 152e-8
 in travelers, 797
 treatment of, 152e-5t, 246e-9, 1414t, 1416
⁹⁰Strontium exposure, 263e-3t, 263e-6t
 Strontium ranelate, 2503
⁸⁹Strontium therapy
 for bone pain, 61, 474
 for metastatic bone disease, 119e-4
 for metastatic breast cancer, 530
 Structure (quality theory), 12e-2
 Struma ovarii. *See* Teratoma, ovarian
 Struvite stones, 1871. *See also* Nephrolithiasis
 ST segment elevation, 1456, 1457, 1457f, 1457t
 ST-segment elevation myocardial infarction (STEMI), 1599. *See also* Myocardial infarction
 circadian variations in, 1599
 classification of, 1602t
 clinical features of, 96, 96t, 1599–1600
 complications of, 1602, 1607
 arrhythmias, 1608–1609, 1609f
 cardiogenic shock, 1608
 heart failure, 1607–1608
 hypovolemia, 1607–1608
 left ventricular aneurysm, 1610
 pericarditis, 1610
 recurrent chest pain, 1609–1610
 right ventricular infarction, 1608
 thromboembolism, 1610
 ventricular dysfunction, 1607
 definition of, 1602t
 diagnosis of
 biomarkers, 1600–1601, 1601f
 CMR, 1601–1602
 ECG, 269e-3f, 269e-4f, 297e-2f, 1456–1457, 1457f, 1457t, 1599f, 1600
 echocardiography, 271e-1f, 271e-1v, 1601
 physical examination, 1500
 radionuclide imaging, 1601
 differential diagnosis of, 1600
 epidemiology of, 1599
 hemodynamic assessment in, 1607
 incidence of, 1593, 1594f, 1599
 nonspecific reaction to, 1601
 painless, 1600
 pathophysiology of, 1599. *See also* Atherosclerosis; Coronary artery disease (CAD)
 pulmonary edema in, 1764
 recurrent, 1610
 risk factors for, 1599
 risk stratification and management after, 1610
 secondary prevention after, 1610–1611
 stages of, 1600
 TIMI grading system, 1605
 treatment of
 ACE inhibitors, 1607
 activity restriction, 1606
 antiplatelet agents, 1603, 1606
 antithrombotic therapy, 1606
 ARBs, 1607
 beta blockers, 1603–1604, 1607
 bowel management, 1606
 diet therapy, 1606
 in emergency department, 1603, 1604f
 fibrinolytic therapy, 1605
 hospital/coronary care unit, 1605–1606
 implantable cardioverter-defibrillator, 1609, 1609f
 integrated approach to, 1604f, 1605
 limitation of infarct size, 1604
 morphine, 1603
 nitrates, 1603, 1607
 oxygen therapy, 1603
 pacemaker, 1609
 pain control, 1603–1604
 PCI
 indications for, 1604–1605, 1604f
 left main, 297e-3–4, 297e-3–4v, 297e-4f
 with manual thrombectomy, 297e-2, 297e-2f, 297e-2v
 rescue, 1605
 time from symptom onset to, 1603, 1603f
 urgent, 1605
 prehospital care, 1602, 1603f
 reperfusion therapy, 1604, 1604f, 1605
 sedation, 1606
 Stupor, 1771
 Sturge-Weber syndrome, 441e-38f
 STW 5, 263
 Sty (hordeolum), 199
 Stylostome, 2745
 Subacromial bursitis, 2247
 Subacute combined degeneration, 2659
 Subacute necrotizing myelopathy, paraneoplastic, 614t
 Subacute sclerosing panencephalitis (SSPE), 900, 1298
 Subaortic stenosis, 1525
 Subarachnoid hemorrhage, 1784
 vs. bacterial meningitis, 888
 clinical features of, 1785
 in critically ill patient, 1735, 1784
 diagnosis of, 1786, 1786f
 DVT prophylaxis after, 1787
 ECG in, 1457, 1458f
 epidemiology of, 1784
 etiology of, 1784–1785
 fundus examination in, 1773
 grading of, 1785t
 headache in, 107t, 108, 1785
 at high altitudes, 476e-4
 hyponatremia in, 299, 1785–1786
 hypothermia in, 478e-1
 neurologic deficits after, 1785–1786
 pathophysiology of, 1785
 in polycystic kidney disease, 1852
 treatment of, 1786–1787
 Subclavian artery, occlusion of, 2576
 Subconjunctival hemorrhage, 199
 Subcortical aphasia, 179
 Subcutaneous panniculitis-like T-cell lymphoma, 135e-4, 708
 Subdural empyema, 783, 903
 approach to the patient, 883–884, 884–885f
 clinical features of, 903–904
 definition of, 903, 903f
 diagnosis of, 904, 904f
 differential diagnosis of, 904
 epidemiology of, 903
 etiology of, 903, 1097
 pathophysiology of, 903
 prognosis of, 904
 treatment of, 904
 Subdural hematoma
 acute, 457e-3, 457e-3f
 chronic, 457e-3–4, 457e-3f
 coma in, 1772
 dementia in, 2608
 imaging in, 441e-34f, 1772f
 weakness in, 156
 Suberosis, 1681t
 Subeschar clysis, 166e-3
 Subfascial endoscopic perforator surgery, 1653
 Sublingual space infections, 235
 Subluxation, 2219, 2219t
 Submandibular space infections, 233, 235, 236, 1097
 Submucosal plexus, 264
 Subperiosteal abscess, 231
 Sub-Saharan Africa
 health care delivery in, 13e-6–7
 health workforce and disease burden in, 13e-3, 13e-4f
 life expectancy in, 13e-3f
 Substance P
 in neural control of gastrointestinal system, 264
 pain and, 88, 88f
 in swallowing, 255
 Substance use disorders. *See also* Injection drug users; *specific substances*
 delirium in, 168–169
 dementia in, 2608
 genetic considerations in, 465e-1
 neuropathology of, 444e-4, 465e-4, 465e-5t
 pathogenesis of, 465e-3, 465e-3f
 polydrug, 469e-5
 prevalence of, 2708
 in women, 6e-5
 Substernal goiter, 2301
 Substrate reduction therapy, 432e-5
 Succussion, 86e-1t
 Succimer (DMSA), 472e-2–3t
 Succinate dehydrogenase (SDH) enzyme, 2329, 2332
 Succinylcholine, 34t, 310
 Suck response, 2539
 Sucralfat, 263, 1920t, 1921, 1930
 Sudan virus disease, 1323, 1325f, 1326f
 Sudden cardiac death, 1764
 circadian variations in, 194
 definition of, 1764, 1765t
 epidemiology of, 1764–1765, 1766f
 etiology of, 1764–1765, 1765t
 genetic considerations in, 1765
 in heart failure, 1514
 in hypertrophic cardiomyopathy, 1569–1570, 1570t
 incidence of, 1764
 pathology of, 1766
 prediction of, 1766–1767, 1767f
 prevention of, 1766–1767, 1767f, 1770–1771
 in young adults and adolescents, 1764, 1767
 Sudeck's point, 1979, 1979f
 Sudomotor function, 2640–2641, 2641t
 Sufentanil, 1741
 Sugars, malabsorption of, 269
 Suicide
 deaths from, 6e-1t, 56t
 physician-assisted, 67–68, 67t
 worldwide, 17
 Sulbactam, 933, 938, 1037, 1037t. *See also* Ampicillin-sulbactam
 Sulfacetamide, 199
 Sulfadiazine
 action of, 931t, 932
 adverse effects of, 246e-4t
 drug interactions of, 246e-4t
 in pregnancy and lactation, 246e-4t
 resistance to, 931t
 for toxoplasmosis, 903, 1404
 for toxoplasmosis prophylaxis, 1251–1252t
 for Whipple's disease, 1094
 Sulfadoxine, 246e-4t, 1378t, 1384
 Sulfamethoxazole
 action of, 931t, 932
 adverse effects of, 246e-4t, 377, 382, 656t
 drug interactions of, 246e-4t
 for *Nocardia* infections, 1087
 in pregnancy and lactation, 246e-4t
 resistance to, 931t
 Sulfasalazine
 adverse effects of, 138, 646t, 1857, 1959, 2146t
 for ankylosing spondylitis, 2173
 for IBD, 1959, 1959t
 monitoring during treatment with, 2146t
 for psoriatic arthritis, 2177
 for reactive arthritis, 2175
 for rheumatoid arthritis, 2145, 2146t
 Sulfhemoglobin, 249
 Sulfisoxazole, 931t, 932
 Sulfocysteinuria, 434e-2t
 S-Sulfo-L-cysteine metabolism disorders, 434e-2t
 Sulfonamides
 action of, 931t, 932, 932f
 adverse effects of
 cross-sensitivity and, 385
 cutaneous, 25e-6f, 353, 360, 361, 366

- Sulfonamides, adverse effects of (*Cont.*):
 desensitization to, 384, 385
 eosinophilia, 422
 fever, 138
 in G6PD deficiency, 656t
 genetic considerations in, 385
 hepatotoxicity, 2023
 hypersensitivity, 382. *See also* Drug-induced hypersensitivity syndrome (DIHS)
 hypersensitivity syndrome, 382
 nephrotoxicity, 1857
 optic neuropathy, 203
 phototoxicity, 389t
 SJS or TEN, 383. *See also* Stevens-Johnson syndrome (SJS); Toxic epidermal necrolysis (TEN)
 thrombocytopenia, 727, 727t
 vasculitis, 382, 2193
 drug interactions of, 246e-4t
 indications for, 941t
 metabolism of, 34t
 for *Nocardia* infections, 1087
 overdose/poisoning with, 473e-11t
 in pregnancy and lactation, 937t
 resistance to, 931t, 934, 941t
- Sulfonylureas
 action of, 2413, 2414t
 adverse effects of, 389t, 2392, 2413
 contraindications to, 2414t
 drug interactions of, 2413
 for type 2 diabetes mellitus, 2413, 2414t
- Sulfuraphane, 580
- Sulfur-containing shampoo, for tinea versicolor, 350t
- Sulfur dioxide exposure, 1692t
- Sulfur granules, 1090
- Sulfur mustard
 action of, 262e-1-2
 characteristics and health effects of, 262e-1t, 262e-2-3, 262e-3f
 as chemical terrorism agent, 262e-1
 decontamination and treatment of exposure to, 262e-2t, 262e-3-4
- Sulindac, 284, 477, 539
- Sumatriptan
 for cluster headache, 2595
 for cyclic vomiting syndrome, 261
 for migraine, 2591, 2592t, 2593t
 nasal spray, 2591, 2592t
 parenteral, 2591, 2592t
- Summer gripe, 1291
- Summer-type pneumonitis, 1681t
- Sunburn, 386, 386t, 476
- SUNCT/SUNA, 109t, 2595t, 2596
- Sundowning, 167, 168, 170
- Sun exposure, 30t, 385
 melanoma and, 387-388, 476, 477t, 493, 494
 nonmelanoma skin cancer and, 387, 476, 477t, 500
 skin effects of
 acute, 386-387, 386t
 chronic malignant, 387-388, 477t
 chronic nonmalignant, 387
 immunologic, 388
 immunosuppressive, 388
 molecular targets of, 386
 SLE skin flares and, 2126
 vitamin D synthesis and, 387
- Sunflower cataracts, 2519
- Sunitinib
 action and targets of, 101e-8, 102e-3, 102e-5t, 102e-17, 103e-18t, 103e-19
 adverse effects of, 103e-18t, 103e-19, 103e-26, 575, 579, 621
 for gastrointestinal stromal tumor, 102e-3, 102e-5t, 102e-17, 536
 for hepatocellular carcinoma, 102e-3
 for malignant pheochromocytoma, 2331
 for pNETs, 102e-3, 575
 for renal cell carcinoma, 102e-3, 102e-5t, 102e-17, 103e-19, 579
 for small intestinal leiomyosarcoma, 537
- Sunscreen(s)
 ingredients in, 391t
 jellyfish-safe, 2739
 for melanoma prevention, 494
 for polymorphous light eruptions, 389
 for skin cancer prevention, 476
 in SLE, 2130t
 SPF of, 391
 water resistance of, 391
- Superantigens, 124, 145e-9, 372e-19
- Super-conducting quantum-interference device (SQUID), 639
- Superior mesenteric artery occlusion, 104, 258
- Superior pontine stroke syndrome, 2579f
- Superior sagittal sinus thrombosis, 441e-20-22f, 783, 905-906
- Superior vena cava syndrome, 1787
 clinical features of, 1443, 1788
 diagnosis of, 1788, 1789f
 etiology of, 1787-1788
 in lung cancer, 510t, 511
 nocardial, 1085
 treatment of, 1788, 1789f
- Superoxide dismutase, 291e-2
- Supersaturation, 1866
- Supine hypertension, 146
- Suppurative thrombophlebitis, intracranial, 783, 905-906
- Suprachiasmatic nucleus, 195
- Supraglottis. *See* Epiglottitis
- Suprascapular neuropathy, 122
- Supraspinatus tendon, 2248, 2248f
- Supraventricular tachyarrhythmias, 1476
 atrial fibrillation. *See* Atrial fibrillation
 atrial flutter. *See* Atrial flutter
 atrial tachycardia. *See* Atrial tachycardia
 mechanisms of, 1476, 1477t
 in myocardial infarction, 1609
 in overdose/poisoning, 473e-5
 paroxysmal. *See* Paroxysmal supraventricular tachycardia (PSVT)
 in pregnancy, 47
 sinus tachycardia. *See* Sinus tachycardia
- Sural nerve biopsy, 386e-6f
- Suramin
 action of, 246e-13
 adverse effects of, 246e-7t, 607t, 1398, 2686t
 pharmacology of, 246e-13
 for sleeping sickness, 1398, 1398t
- Surfactant replacement therapy, 1739, 1739t
- "Surge-event" preparedness, 918
- Surgical ablation, for atrial fibrillation, 1489
- Surgical myectomy, 1569
- Surgical patient
 asthma in, 1680-1681
 blood loss in, 662
 diabetes mellitus in, 55
 infective endocarditis prophylaxis in, 55
 perioperative risk modification strategies
 ACE inhibitors, 54
 α_1 -adrenergic agonists, 54
 β -adrenergic antagonists, 53-54
 calcium channel blockers, 54
 coronary revascularization, 53
 oral antiplatelet agents, 54
 pulmonary, 54-55, 55t
 statins, 54
 preoperative evaluation of, 51
 for anesthesia, 54
 cardiac risk assessment, 51-53, 52f, 53f, 53t
 functional status, 52t
 intermediate- to high-risk patient, 51
 preoperative questionnaire, 51t
 pulmonary risk assessment, 54-55, 54t
 VTE prophylaxis in, 55, 1636-1637, 1637t
- Surgical-site infections
 diagnosis of, 915
 prevention of, 914t, 915, 945, 945t
 risk factors for, 915
 sternal, 844-845, 926
 streptococcal, 968
 treatment of, 915
- Surgical ventricular restoration, 1515
- Surrogates, for health care decisions, 17e-1-2
- Susceptibility testing, 150e-8
- Sutama. *See* Epidemic (louse-borne) typhus
- Sv (sievert), 263e-1
- Swallowing
 difficulty with. *See* Dysphagia
 physiology of, 255, 255f
 refusal to, 254
 tests of, 257
- Swan-neck deformity, 2136
- S wave, 1451
- Sweat/sweating
 in cystic fibrosis, 1699
 excess, 2643
 initiation of, 479e-1
 in pheochromocytoma, 2329
 potassium loss in, 305
 sodium loss in, 298
 water loss in, 95e-1, 298
- Sweet syndrome
 in cancer patient, 487
 clinical features of, 132t, 135, 366, 487
 differential diagnosis of, 366
 disorders associated with, 366
 drug-induced, 379
 epidemiology of, 132t
 etiology of, 132t
 in IBD, 1958
 myelodysplasia and, 670
 skin manifestations of, 135
 mucocutaneous ulcers, 369
 papulonodular lesions, 132t, 364t, 366
 plaque, 25e-10f, 76e-16f
- Sweet taste sensation, 212
- Swimmer's ear (otitis externa), 228-229, 1045t, 1046
- Swimmer's itch, 829, 1426
- Swine influenza A viruses, 778, 784, 917. *See also* Influenza
- Swinging flashlight test, 196, 196f
- Swyer's syndrome (pure gonadal dysgenesis), 2353
- Sydenham's chorea, 49, 377e-1, 2151, 2622
- Syk, 2115
- Symbiosis, 86e-9
- Sympathectomy, 1649
- Sympathetic apraxia, 179
- Sympathetic postganglionic neuron, 87f
- Sympatholytic agents, 473e-3, 473e-9t, 1624t, 1625
- Sympathomimetics, 473e-8t, 2320t
- Symplecty device, 296e-5
- Symporters (cotransporters), 332e-6
- Synaptobrevin (VAMP2), 985
- Synaptophysin, 120e-2t
- Synaptotagmin, 372e-25t
- Synbiotics, 86e-9
- Synchronized intermittent mandatory ventilation (SIMV), 1741. *See also* Mechanical ventilation
- Syncope, 142
 after vaccination, 791
 in aortic stenosis, 1531
 approach to the patient, 146-147
 cardiac. *See* Cardiac syncope
 definition of, 142
 diagnosis of, 147-148
 differential diagnosis of, 147
 ECG in, 148
 epidemiology of, 142
 heat-related, 479e-1
 high-risk features of, 142t
 natural history of, 142
 neurally mediated (reflex; vasovagal)
 AV conduction block in, 1471
 causes of, 145t
 classification of, 144-145
 clinical features of, 145
 pacemaker for, 1470
 pathophysiology of, 143-144, 144f
 treatment of, 145
 in orthostatic hypotension. *See* Orthostatic hypotension
 pathophysiology of, 142-143
 in psychiatric disorders, 148

- vs. seizure, 147, 2550–2551, 2551t
in ventricular tachycardia, 1491
- Syndrome of apparent mineralocorticoid excess (SAME; 11 β -hydroxysteroid dehydrogenase deficiency)
clinical features of, 306, 1620t, 2319
diagnosis of, 307
genetic considerations in, 306, 1620t, 2318t, 2319
pathogenesis of, 1621, 1621f
treatment of, 307
- Syndrome of inappropriate antidiuretic hormone (SIADH)
vs. cerebral salt wasting, 299
clinical characteristics of, 2280
differential diagnosis of, 301, 2281, 2282t
etiology of, 300, 300t, 2280–2281, 2281t
in HIV infection, 1259
hyponatremia in, 64e–4–5, 299–300, 2280
in lung cancer, 511
pathophysiology of, 299, 2279f, 2281
subtypes of, 299–300
treatment of, 64e–5, 302, 2281–2283, 2282f
tumor-associated, 64e–4–5, 610
- Syndrome X. *See* Metabolic syndrome
- Synkinesis, 2647
- Synovectomy, 2241–2242, 2246
- Synovial chondromatosis, 2246–2247
- Synovial effusion, 2221–2222
- Synovial fluid analysis
in calcium oxalate deposition disease, 2237, 2237f
in calcium pyrophosphate deposition disease, 2235, 2236f
interpretation of, 2223, 2224f
in rheumatoid arthritis, 2144
- Synovial sarcoma, 2247. *See also* Soft tissue sarcoma
- Synovitis
pathogenesis of, 2227, 2227f
pigmented villonodular, 2246, 2247f
- Synthetic cathinones (bath salts), 469e–3–4
- Synthetic lethality, 102e–6f, 102e–7, 103e–23
- Syntrophic relationships, 86e–5
- α -Synuclein, 172t, 444e–10, 453e–6
- Syphilis, **1132**
CNS involvement in. *See* Neurosyphilis
congenital
clinical features of, 836, 1136–1137
global considerations in, 1136
incidence of, 1133, 1136
oral manifestations of, 236, 238t, 1137
treatment of, 1139
diagnosis of
in HIV infection, 1138, 1138t
neurosyphilis, 1138, 1138t
organism identification, 1137
serologic tests, 770, 1137–1138, 1137f
endemic, 207e–2, 207e–2f, 207e–2t, 207e–3
epidemiology of, 879, 1133, 1133f
etiology of, 1132–1133
global considerations in, 1133
in HIV infection, 1259
immunity to, 1140
incidence of, 869–870, 1133f
latent, 1135
pathogenesis of, 1134
prevention of, 1138
primary
clinical features of, 1134–1135
differential diagnosis of, 1135
natural course of, 1134
skin manifestations of, 25e–4f, 76e–9f, 880t, 881, 1134, 1134f
treatment of, 1138–1139, 1139t
secondary, 1136
articular manifestations of, 836
cardiovascular manifestations of
aortic aneurysm, 1638
aortitis, 1134, 1136, 1643
AV conduction block, 1471, 1471t
epidemiology of, 1133, 1133f
hepatitis in, 1135
lymphadenopathy in, 408
natural course of, 1134
neurologic manifestations of. *See* Neurosyphilis
ocular involvement in, 1135
oral manifestations of, 237, 238t
pharyngitis in, 231
renal involvement in, 1836t, 1849
skin manifestations of, 130t, 134, 353, 362, 1135
alopecia, 354–355, 355t
condylomata lata, 25e–5f
figurate lesions, 356, 356f
histopathology of, 1134
mucous patch, 1135
on palms and soles, 25e–5f, 76e–10f, 1135f
papulosquamous eruption, 25e–5f, 76e–10f, 1135f
vs. pityriasis rosea, 134, 347t, 349
on tongue, 25e–5f
treatment of, 1139, 1139t
tertiary
clinical features of, 238t, 1134, 1136
treatment of, 1139, 1139t
transmission of, 1133
treatment of, 1138–1140, 1139t
untreated, natural course of, 1134
- Syngomyelia, 2658, 2658f
- Syrinx cavities, 2658f
- Systematic reviews, 26
- Systemic inflammatory response syndrome (SIRS), 98e–3, 98e–4
- Systemic lupus erythematosus (SLE)
antiphospholipid syndrome and, 2133
autoantibodies in, 377e–5, 2125, 2126t, 2127t, 2131
clinical features of, 129t, 377e–5, 2127, 2129t
cardiovascular, 290e–1t, 290e–3, 2129
AV conduction block, 1471, 1471t
pericarditis, 1575, 2129, 2129t
SA node dysfunction, 1467
cutaneous, **366**, 2127–2128, 2129t
alopecia, 354, 355t
characteristics of, 129t, 134
discoid (chronic), 25e–3f, 76e–13f, 2127–2128, 2129t
distribution of, 366
malar erythema, 25e–3f, 76e–13f, 2128
panniculitis, 366
papulonodular red lesions, 366
papulosquamous lesions, 76e–13f, 353t
subacute, 25e–3f, 2128
telangiectasias, 356t, 357
treatment of, 2132, 2134
vasculitis, 363, 2193
gastrointestinal, 2129t, 2131
hematologic, 664, 2129t, 2131
hemoptysis, 245
lymphadenopathy, 408
musculoskeletal, 2127, 2129t
neurologic, 909t, 2128–2129, 2129t, 2623, 2656, 2683
ocular, 2129t, 2131
oral, 237
pulmonary, 1714, 2129, 2129t
Raynaud's phenomenon, 1649
renal, 1858. *See also* Lupus nephritis
splenomegaly, 411
systemic, 2127, 2129t
course of, 2131
definition of, 2124
diagnosis of, 2126–2127, 2127t, 2131, 2328f
drug-induced, 379, 2134
EBV and, 2126
environmental factors in, 2125f, 2126
epidemiology of, 2124
genetic considerations in, 373e–8t, 2125–2126, 2125f
global considerations in, 2134
HLA gene association with, 373e–8t
incidence and prevalence of, 2124
inflammatory mediators in, 372e–29t
microvascular thrombotic crisis in, 2133–2134
pathogenesis of, 2124–2126, 2125f, 2126t
pathoimmunology of, 372e–24–25t, 372e–27t, 377e–2
pathology of, 2124–2126, 2125f, 2126
prognosis of, 2134
sex-related factors in, 2125–2126
survival rates, 2134
treatment of, 2128f, **2131**
cytokine blockers, 377e–5
experimental therapies, 2134
life-threatening disease, 2132–2133
non-life-threatening disease, 2132
outcomes, 2134
pharmacologic, 2130t, 2132–2134
during pregnancy, 2133
for prevention of complications, 2134
special conditions, 2133–2134
- Systemic lytic state, 758
- Systemic mastocytosis. *See* Mastocytosis, systemic
- Systemic mastocytosis associated with clonal hematologic non-mast cell lineage disease (SM-AHNMD), 2120, 2120t
- Systemic necrotizing vasculitis, 372e–8
- Systemic onset juvenile idiopathic arthritis (SOJIA), 372e–8t
- Systemic sclerosis (scleroderma), **2154**
cancer associated with, 2162–2163
clinical features of, 2158, 2158t
acro-osteolysis, 2160, 2160f
cardiac, 1471, 1471t, 1567, 2158, 2162
cutaneous, **375**, 2159–2160
calcinosis, 376, 2160, 2160f
vs. dermatomyositis, 375
digital necrosis, 376f, 2159f
dystrophic calcification, 363
facial, 76e–14f, 376f
hyperpigmentation, 360, 360t
hypopigmentation, 357t, 359
limited vs. diffuse, 2155t
pathology of, 2157, 2157f
sclerodactyly, 2159f
telangiectasias, 356t, 357, 376, 2160f
vascular, 2160, 2160f
esophageal involvement/dysphagia, 256, 257, 1911
facial hemiatrophy, 2649
gastrointestinal, 259, 2158, 2161–2162, 2164
initial presentation, 2159
musculoskeletal, 2162, 2218t
neuropathy, 2683
ocular, 2162
pulmonary
interstitial lung disease, 1713, 2160–2161, 2161f
pathology of, 2157–2158, 2158f
pulmonary arterial hypertension, 1658, 2161, 2164
Raynaud's phenomenon, 1649, 2156, 2159, 2159f, 2160f
renal, 62e–9f, 1864–1865, 2158, 2162, 2164
vasculopathy, 2156, 2162
definition of, 2154
diagnosis of, 2163
diffuse cutaneous, 2155t
environmental factors in, 2155
epidemiology of, 2155
genetic considerations in, 2155
limited cutaneous, 2155t
linear, 376
localized, 2165
natural history of, 2165
pathogenesis of, 372e–15t, 372e–24–25t, 2156–2157, 2157t
pathology of, 2157–2158, 2158f
pregnancy in, 2162
prognosis of, 2165
risk factors for, 2155–2156
treatment of, **2163**
antifibrotic therapy, 2164
for gastrointestinal complications, 2164
immunosuppressive agents, 2163–2164
for musculoskeletal complications, 2165
principles of, 2163, 2163t
for pulmonary complications, 1713, 2164

- Systemic sclerosis (scleroderma), treatment of
(*Cont.*):
for renal complications, 1865, 2164
skin care, 2164–2165
vascular therapy, 2164
- Systemic vascular resistance, 1729
calculation and normal values, 1463, 1748t
effect on murmurs, 51e-7
in pregnancy, 45
in sepsis/septic shock, 1755
in shock, 1744, 1748t
- Systems biology, **87e-1**
human disease classification and, 87e-3–4,
87e-4f, 87e-5f, 87e-6f
in medical practice, 6
pathobiology applications of, 87e-3–4
properties of biologic systems, 87e-1–2, 87e-2f
- Systems neuroscience, 444e-12, 465e-3–4
- Systems theory, 12e-1, 12e-1f
- Systolic blood pressure, 1616t, 1622
- Systolic click-murmur syndrome. *See* Mitral valve prolapse
- Systolic failure. *See* Heart failure (HF)
- Systolic murmurs. *See* Heart murmurs
- SYT gene mutations, 119e-1
- T**
- T₃. *See* Triiodothyronine (T₃)
- T₃-resin uptake test, 2288
- T₄. *See* Thyroxine (T₄)
- TABI/2, 145e-6f
- Tabes dorsalis, 1136, 2658f
- Tabun. *See* Nerve agents
- TAC3 gene mutations, 2364t, 2377
- TAC3R gene mutations, 2364t
- Tacaiuma virus, 1308t
- Tâche noire, 1157, 2745
- Tachycardia
atrioventricular nodal reentry, 1479–1480,
1481f
in hypovolemia, 298
in overdose/poisoning, 473e-3
preexcited, 1482, 1483f
supraventricular. *See* Supraventricular tachyarrhythmias
syncope in. *See* Cardiac syncope
treatment of, 1479t
ventricular. *See* Ventricular tachycardia
- Tachycardia-bradycardia syndrome, 146, 1468,
1469
- Tachykinins, 565
- Tachypnea, 1746, 1757
- Tacrine, 473e-10t, 2601
- Tacrolimus
action of, 1829t, 2071
adverse effects of, 1829, 1829t, 2071
hyperkalemia, 310
myopathy, 2218t
nephrotoxicity, 1856t, 1862, 1864, 2071
osteoporosis, 2492
seizures, 1793, 2547t
skin cancer, 388
thrombocytopenic purpura, 729
for atopic dermatitis, 345, 354t
drug interactions of, 246e-3t, 2071
genetic variations in response to, 39t
for IBD, 1961
for immunosuppression
after kidney transplantation, 1829, 1829t
after liver transplantation, 2071
for lupus dermatitis, 2134
metabolism of, 34t, 2071
for myasthenia gravis, 2705
for polymyositis, 2201
for SLE, 2133
therapeutic monitoring of, 2766t
- Tactile fremitus, 1662
- Tadalafil
action of, 324, 325f, 326f
adverse effects of, 328, 329t
characteristics of, 329t
contraindications to, 328, 329t
for erectile dysfunction, 328, 2672
for high-altitude pulmonary edema, 476e-3
for lower urinary tract symptoms, 587
for pulmonary arterial hypertension, 1660,
1660t
for Raynaud's phenomenon, 1649
- Taenia asiatica*, 1430
- Taenia multiceps*, 1435
- Taenia saginata* (beef tapeworm) infection
clinical features of, 1430
diagnosis of, 245e-1, 245e-2t, 245e-5t, 245e-6t,
1430
geographic distribution of, 245e-2t, 1430
life cycle of, 245e-2t, 1430
pathogenesis of, 1430
prevention of, 1430
treatment of, 1430
- Taenia serialis*, 1435
- Taenia solium* (pork tapeworm) infection
clinical features of, 1430–1431. *See also* Cysticercosis
diagnosis of, 245e-1, 245e-2t, 245e-5t, 245e-6t,
1431–1432, 1431t
geographic distribution of, 245e-2t, 1430
life cycle of, 245e-2t, 1430
pathogenesis of, 1430
prevention of, 1432
treatment of, 1432
- Tafamidis, 724
- Tafenoquine, 246e-1t, 246e-13
- TAFI (thrombin-activatable fibrinolysis inhibitor), 744
- Tag SNPs, 442
- Tahyña virus, 1309t, 1315
- Tai chi, 14e-1t
- Tai Forest virus disease, 1323, 1325f, 1326f
- TAK-1, 145e-6f
- Takayasu's arteritis, **2189**
aortic aneurysm in, 1638
aortitis in, 1642
arterial involvement in, 386e-5f, 2189, 2189t
carotid or vertebral artery occlusion in, 2567,
2574
clinical features of, 1642, 2189–2190
definition of, 2189
diagnosis of, 386e-5f, 1642, 2190
epidemiology of, 1642
incidence and prevalence of, 2189
pathogenesis of, 1642, 2189
treatment of, 1642, 2190
- Takotsubo (stress-related) cardiomyopathy, 97,
289e-3
- TALI* gene mutations, 101e-3t
- Talc, intrapleural, 474
- Talc dust, 1690
- Talc embolism, 40e-4
- Tamm-Horsfall protein, 293, 294, 1804, 1854,
1859
- Tamoxifen
action of, 102e-4f, 102e-7, 103e-15
adverse effects of
cataracts, 531, 623
depression, 63
endometrial cancer, 103e-15
hepatotoxicity, 2025
thrombocytopenia, 727t
thromboembolism, 103e-15
tumor lysis syndrome, 1795
uterine cancer, 531, 596, 623
visual loss, 607t
for breast cancer prevention, 29t, 478, 531,
2498–2499
for breast cancer treatment, 102e-7, 103e-15,
528–529, 529t, 530t
drug interactions of, 40, 205e-3
for ductal carcinoma in situ, 531
genetic variations in response to, 39t, 40
for gynec mastia, 585, 2367
metabolism of, 35
for osteoporosis management/prevention,
2498
- Tanapox, 220e-1t, 220e-2
- TandemHeart, 296e-5
- Tangier disease (ABCA1 deficiency), 1443, 2445,
2681
- Tanning, 386, 387, 498, 500
- Tapeworms/tapeworm infections. *See* Cestodes/cestode infections
- TAP (transporters associated with antigen processing) proteins, 372e-19, 373e-3–4
- Tap test, 2748
- Tarantulas, 2748
- TARC (CCL17), 372e-12t, 1674, 1674f
- Tardive dyskinesia, 163, 261, 2624, 2722
- Tardive dystonia, 2624
- Tardive syndromes, 2624
- TARDP* gene mutations, 444e-1, 2603
- Target cells, 81e-1, 81e-4f, 395f
- Target lesion revascularization, 296e-3
- Target of rapamycin (TOR), 94e-5. *See also* mTOR
- Target vessel revascularization, 296e-3
- Target volume, in radiation therapy, 103e-4
- Tartrate-resistant acid phosphatase (TRAP), 426e-4
- Tartrazine, 1671
- Tarui's disease (phosphofructokinase deficiency), 433e-2t, 462e-14, 654t
- Taste buds, 212, 213f, 215
- Taste disorders
diseases and conditions associated with, 215
drug-related, 215
etiology of, 215
evaluation of, 215–216
head injury and, 457e-2
olfactory dysfunction and, 215
treatment of, 216–217
in zinc deficiency, 96e-9
- Taste pore, 212, 213f
- Taste system, 212–214, 213f
- TATA-binding protein, 431t
- Tatagaine virus, 1306t
- tat* gene, 225e-3f, 1219f
- Tat protein, 225e-2
- Tau protein, 172t, 175, 453e-6
- TAVR (transcatheter aortic valve replacement).
See Aortic valve replacement, transcatheter
- Taxanes. *See also specific drugs*
adverse effects of, 620t, 1798, 2686t
for breast cancer, 529, 530
characteristics of, 103e-12t, 103e-14–15
drug interactions of, 1793
for melanoma, 499
for ovarian cancer, 593
resistance to, 103e-15
for soft tissue sarcoma, 119e-2
- Tax proteins, 214e-5, 214e-7, 225e-2, 225e-3–4
- Tay-Sachs disease, **432e-1**
adult, 432e-1, 432e-3t, 2634t, 2636
“cherry red” spot in eye in, 436e-7f
enzyme deficiency in, 432e-3t
genetic screening for, 432e-1, 443
infantile, 432e-1
juvenile, 432e-1, 432e-3t
- Tazarotene, 348
- TAZ* gene mutations, 1564
- Tazobactam, 933
- TB. *See* Tuberculosis (TB)
- TBG (thyroxine-binding globulin), 2285, 2286,
2286t, 2288
- TBI (traumatic brain injury). *See* Head injury
- TBII (thyrotropin-binding inhibitory immunoglobulin), 2290, 2294
- TBX* gene mutations, 431t
- TCAs. *See* Tricyclic antidepressants (TCAs)
- T cell(s)
age-related reduction in, 94e-4
allogenic, transfer of, 103e-20
antigen recognition by, 372e-18–19,
372e-18f
in asthma, 1673, 1674f
in autoimmune diseases, 372e-23, 377e-1t,
377e-2, 377e-4
autologous, manipulation of, 103e-20
autoreactive, in MS, 2662
central memory, 372e-16
in cord blood, 89e-3–4
cytotoxic. *See* CD8+ T cells

- deficiencies of, 375e-1, 2104t, 2107, 2107f, 2108f
 cartilage hair hypoplasia, 418, 2109
 with DNA repair defects, 375e-1–2, 2109.
See also Ataxia telangiectasia (AT)
 functional defects, 2109
 with hyper IgE. *See* Hyper-IgE syndrome
 in hyper-IgM syndrome, 2110
 in Omenn syndrome, 2108–2109
 in severe combined immunodeficiency. *See*
 Severe combined immunodeficiency
 (SCID)
 in thymic defects, 2108
 in Wiskott-Aldrich syndrome. *See* Wiskott-
 Aldrich syndrome
 definition of, 372e-2
 development of, 372e-16–18, 372e-17f
 differentiation of, 697–698, 699f
 hematopoietic, 89e-3f
 in primary immune deficiency diseases,
 2107, 2107f, 2108f
 effector memory, 372e-15–16
 helper. *See* CD4+ T cells
 inactivation by tumor cells, 102e-17, 102e-18f
 locations of, 372e-16
 MHC class-restricted, 372e-18, 373e-3, 373e-4
 monoclonal antibodies to, for autoimmune
 and inflammatory diseases, 372e-
 30–31
 negative selection, 372e-17
 NK. *See* Natural killer (NK) T cells
 positive selection, 372e-17
 priming against tumor-associated peptides,
 103e-20
 in psoriasis, 348
 regulatory, 1673
 in response to *S. aureus*, 956
 in sarcoidosis, 2205f
 in vasculitis, 2180–2181
 T-cell exhaustion, 372e-2, 372e-19
 T cell large granular lymphocytic leukemia,
 135e-3, 2138
 T-cell prolymphocytic leukemia, 135e-3
 T cell receptors (TCRs)
 for antigens, 372e-2, 372e-18–19
 defective rearrangements of, 2108
 diversity of, 372e-18–19
 HLA genes and, 373e-1, 373e-3
 signaling through, 372e-18f
 trimolecular complex of, 373e-2f
 T cell superantigens, 372e-19
TCIRG1 gene, 426e-4
TCL1 gene mutations, 135e-3
 Tdap vaccine. *See* Tetanus, diphtheria, acellular
 pertussis (Tdap) vaccine
TDF (testis-determining factor), 439
TDP-43 gene mutations, 444e-9, 2633
TDP-43 protein, 172t, 2633
 Td vaccine. *See* Tetanus-diphtheria (Td) vaccine
 Teaching, physician engaged in, 7
 Teardrop cells, 81e-5f
 Teardrop fracture, 441e-34f
 Technetium-99m, 263e-3t, 270e-3, 270e-3t
 TECK (thymus expressed chemokine), 372e-12t
 Tecovirimat, 261e-6
 Tedizolid
 action of, 931t, 932, 961
 adverse effects of, 938t
 drug interactions of, 939t
 indications for, 941t, 944
 for MRSA infections, 962t
 in pregnancy and lactation, 937t
 resistance to, 931t, 934, 941t
 for staphylococcal infections, 961, 962t
 Teduglutide, 1944
 TEE. *See* Transesophageal echocardiography
 (TEE)
 Teeth. *See also* Dental entries
 age-related changes in, 242
 development of, 235–236
 discolored, 236
 diseases of, 46e-1, 235
 Hutchinson's, 237
 loss of, 236–237, 242
 in osteogenesis imperfecta, 2508
 structure of, 235–236
 tooth extraction, 242
 TEF, 401e-1, 401e-1t
 Tegaserod, 1970
 Teicoplanin, 930, 931t
 Telangiectasia(s), 356. *See also* Ataxia
 telangiectasia (AT)
 capillary, 2586
 in cardiovascular disease, 1442
 definition of, 339t, 1650
 etiology of, 356t
 hereditary hemorrhagic. *See* Osler-Weber-
 Rendu syndrome
 intestinal, 536
 linear, 356–357
 mat, 357
 periungual, 357
 in systemic sclerosis, 2160, 2160f
 treatment of, 2165
 Telaprevir
 action of, 215e-11
 adverse effects of, 381, 2046–2047
 for chronic HCV infection, 2044, 2045–2046t,
 2046–2047
 drug interactions of, 2047
 Telavancin
 action of, 930, 931t, 961
 adverse effects of, 938t, 942
 for enterococcal infections, 976
 indications for, 940t, 942
 in pregnancy and lactation, 937t
 resistance to, 931t, 933–934
 for staphylococcal infections, 961
 Telbivudine
 action of, 215e-10
 adverse effects of, 2036
 for chronic HBV infection, 215e-4t, 215e-10,
 2036, 2037t, 2038f
 pharmacology of, 215e-10
 resistance to, 215e-10, 2040t
 Teletherapy, 103e-4
 Telithromycin
 action of, 931, 931t, 943
 adverse effects of, 943
 drug interactions of, 943
 indications for, 943
 for *Legionella* infections, 1020t
 resistance to, 931t, 934
 Telmisartan, 2570
 Telogen, 331
 Telogen effluvium, 354, 355t, 379
 Telomerase, in cancer cells, 102e-3, 440
 Telomere, 83e-1, 94e-3, 102e-3, 440
 Telotristat eprate, 566
 Temazepam
 for anxiety disorders, 2711t
 for insomnia, 191
 overdosage/poisoning with, 473e-10t
 pharmacology of, 2711t
 for sleep impairment at high altitudes, 476e-4
 Temozolomide
 action of, 103e-12
 adverse effects of, 103e-9t, 574
 for astrocytoma, 600
 for glioblastoma, 600
 for malignant pheochromocytoma, 2331
 for melanoma, 499
 for oligodendroglioma, 601
 for pNETS, 574
 Temperature sense, 159, 160t
 Temporal arteritis. *See* Giant cell arteritis
 (temporal arteritis)
 Temporal bone, fracture of, 220
 Temporal lobectomy, for refractory epilepsy,
 2557
 Temporal lobe epilepsy, 2546
 Temporomandibular joint disorder, 237
 Temsirolimus
 action and targets of, 102e-5t, 103e-16f,
 103e-19, 103e-19t
 adverse effects of, 102e-5t, 103e-19, 103e-19t,
 103e-26, 1797
 for mantle cell lymphoma, 704
 for NETs, 574
 for renal cell carcinoma, 102e-5t, 579
 TEN. *See* Toxic epidermal necrolysis (TEN)
 Tenatoprazole, 1921
 Tender point assessment, in fibromyalgia, 2238,
 2238f
 Tendinitis
 bicipital, 2248
 calcific, 2248
 drug-induced, 2218t
 patellar, 2248
 rotator cuff, 2221, 2248
 Tendinous xanthomas, 436e-3f
 Tenecteplase
 action of, 759
 for acute limb ischemia, 1646
 contraindications to, 1605
 domain structure of, 759, 759f
 indications for, 759
 for myocardial infarction, 1605
 for pulmonary embolism, 1636
 Teneligliptin, 2414t
 Tennis elbow, 2249
 Tenofovir
 action of, 215e-9, 1278
 adverse effects of, 215e-9, 1274t, 1278
 for chronic HBV infection, 2037
 comparison to other antivirals, 2037t, 2038f
 dosage for, 215e-4t, 215e-9
 therapeutic monitoring of, 215e-4t, 2039,
 2074
 drug interactions of, 1278
 for HIV infection, 215e-9, 1274t, 1278
 molecular structure of, 1276f, 1278
 resistance to, 215e-9, 1279f, 2040t
 Tenosynovitis, 2248
 Tension pneumothorax, 98, 1719
 Tension-type headache, 107t, 2587t, 2593–2594
 Teratoma
 ovarian
 anti-NDMA encephalitis and, 615f, 617,
 617f
 benign, 594
 clinical features of, 594
 genetic considerations in, 439
 paraneoplastic opsoclonus-myoclonus
 syndrome and, 618
 thyrotoxicosis in, 2298
 parasellar, 2264
 pineal, 603
 testicular, 588, 591
 Terazosin
 for hypertension, 1624t
 for Raynaud's phenomenon, 1649
 for urinary dyssynergia, 2660, 2672
 Terbinafine
 adverse effects of, 215, 2218t
 for onychomycosis, 350, 1331, 1358, 1358t
 for *Sporothrix* infections, 1354t
 for tinea, 1331, 1358, 1358t
 for tinea capitis, 355t
 Terbutaline, 473e-8t
TERC gene mutations, 664–665
 Terfenadine, 42t
 Teriflunomide, 2669t, 2670, 2671
 Teriparatide, 2383, 2502–2503, 2502f, 2503f
 Terlipressin, 278, 1810
 Terminal extubation, 66–67
 Terminal hair, 331
 Terminally ill patient, 55
 advance care planning for, 58–59, 59t
 assessment of, 56–57
 with cancer, 475
 care during last hours, 68–69, 69t
 clinical courses of, 68f
 communicating bad news to, 57–58, 57t
 continuous goal assessment, 48
 delirium in, 169
 depression in, 64–65
 euthanasia and physician-assisted suicide for,
 67–68, 67t
 existential needs of, 66
 financial burdens of, 65
 futile care of, 67

- Terminally ill patient (*Cont.*):
 managing last stages for, 66–67
 physical symptoms of, 59, 60t
 anorexia, 69t
 apnea, 69t
 constipation, 61, 62t
 dehydration, 69t
 delirium, 69t
 dysphagia, 69t
 dyspnea, 62, 62t
 fatigue, 63–64, 69t
 incontinence, 69t
 nausea and vomiting, 61–62
 pain, 60–61
 psychological symptoms of, 60t, 63
 delirium, 64–65, 65t
 depression, 64–65
 insomnia, 65
 social needs of, 65–66
 withholding/withdrawal of care of, 66–67
- Terminal restlessness, 169
- Terminal weaning, 66–67
- Terrorism
 chemical. *See* Chemical terrorism
 microbial/biological. *See* Bioterrorism
 radiation. *See* Radiation terrorism
- Terson's syndrome, 40e-3
- TERT gene mutations, 664–665
- Testes/testicles, 2357
 biopsy of, 2361
 development of, 2357–2358, 2357f
 disorders of. *See also* Male reproductive system disorders
 acquired defects, 2365–2366
 adrenal rest tumors, 2328, 2328f, 2356
 alcohol use disorder and, 2725
 cancer. *See* Testicular cancer
 in Carney's complex, 2343
 cryptorchidism, 588, 2355, 2365
 in disorders of sex development. *See* 46,XY disorders of sex development
 in Klinefelter's syndrome, 2365. *See also* Klinefelter's syndrome
 mass, 588
 neuroendocrine tumors, 564
 torsion, 871–872, 2365
 dysgenetic, 2353, 2354t
 in lepromatous leprosy, 1125
 in mumps infection, 231e-2, 231e-3
 physical examination of, 2360
 referred pain from, 105
 regulation of function of
 androgen in, 2358–2359, 2358f, 2359f, 2359t
 hypothalamic-pituitary-testis axis in, 2358, 2358f
 seminiferous tubules in, 2359–2360
 structure of, 2357–2358
- Testicular cancer, 588
 clinical features of, 588
 epidemiology of, 588
 etiology of, 588
 genetic considerations in, 588
 global considerations in, 588
 incidence of, 588
 metastatic, 408
 paraneoplastic syndromes in, 609t, 614t, 619
 pathology of, 588
 risk classification in, 590–591, 591t
 staging of, 588, 589f
 treatment of
 chemotherapy for advanced disease, 590
 HSCT, 139e-6
 infertility after, 592
 late consequences of, 623, 624t
 nonseminoma, 589f, 590
 postchemotherapy surgery, 591
 risk-directed chemotherapy, 590–591, 591t
 salvage chemotherapy, 591
 seminoma, 589f, 590
 tumor markers in, 588
- Testicular dysgenesis, 2353, 2354t
- Testicular feminization syndrome. *See* Androgen insensitivity syndrome (AIS)
- Testis-determining factor (*TDF*), 439
- Testosterone
 age-related changes in levels of, 7e-1, 7e-1f, 75, 75f
 assays of, 2360–2361
 bioavailable, 2361
 in hirsutism, 332–333
 in men with prostate cancer, 7e-4–5
 reference values for, 2763t
 synthesis of, 2309f, 2359f
 total, tests of, 2360
 transport and metabolism of, 2358–2359, 2359f
 unbound, measurement of, 2361
- Testosterone cypionate, 2370t, 2371
- Testosterone enanthate, 2370t, 2371
- Testosterone-lowering drugs. *See* Androgen deprivation therapy (ADT)
- Testosterone therapy
 abuse of. *See* Anabolic-androgenic steroid (AAS) abuse
 for adult hypopituitarism, 2259t
 adverse effects of, 2371, 2372, 2372t
 for androgen deficiency, 2370–2371
 contraindications to, 2372, 2372t
 for delayed puberty, 2363
 efficacy of, 2372
 for erectile dysfunction, 328
 formulations of, 2370t
 buccal adhesive, 2371
 in development, 2371
 gel, 2370t, 2371
 implants, 2371
 injectable, 2370t, 2371
 not available in U.S., 2371
 oral, 2370t, 2371
 transdermal patch, 2370t, 2371
 gonadotropin inhibitors and, 2372
 for hypogonadism, 2261
 indications for, 2371–2372
 for male contraception, 2372
 in men with history of prostate cancer, 7e-5, 2372
 monitoring of, 329, 2372–2374, 2373t
 in older men, 7e-1, 7e-2f, 7e-2t
 regimens for, 2372
- Testosterone undecanoate, 2370t, 2371
- Testotoxicosis, 2362
- TET2 gene mutations, 135e-6, 670, 680, 680t
- Tetanospasm, 984
- Tetanus, 984
 approach to the patient, 985, 985f
 clinical features of, 985, 985f
 complications of, 986
 definition of, 984
 diagnosis of, 985–986
 epidemiology of, 984
 etiology of, 984
 global considerations in, 984, 986–987
 incidence pre- and postvaccine, U.S., 785t, 984
 neonatal, 984, 986–987
 pathogenesis of, 984–985
 prevention of, 29t, 787–788f, 986–987
 prognosis of, 986, 986t
 prophylaxis, for bite-wounds, 167e-3
 treatment of, 986
- Tetanus, diphtheria, acellular pertussis (Tdap) vaccine
 contraindications to, 789t
 precautions for, 789t
 recommended administration of
 in adults, 787–788f, 980
 in children, 980
 in pregnancy, 51, 788f, 987
 in travelers, 794
- Tetanus booster, for bite-wounds, 167e-3
- Tetanus-diphtheria (Td) vaccine
 contraindications to, 789t
 recommended administration of
 in adults, 29t, 787–788t, 980
 in cancer patient, 485t
- Tetanus immune globulin (TIG), 167e-3, 986
- Tetanus vaccine
 for burn patient, 166e-3
 for travelers, 794
- Tetany, in malabsorption syndromes, 1946t
- Tethered cord syndrome, 118, 2659
- Tetherin, 1218
- Tetrabenazine, 2611, 2621, 2622, 2624
- Tetracycline(s)
 for acne rosacea, 352
 for acne vulgaris, 352
 for actinomycosis, 1091t
 action of, 930–931, 931t, 932f, 943
 adverse effects of, 938t, 943
 cutaneous, 246e-7t, 360, 362, 379
 folate deficiency, 646t
 frequent, 246e-7t, 1380t
 lupus syndrome, 2218t
 nail disorders, 380
 papilledema, 203f
 phototoxicity, 389t
 rare, 246e-7t
 renal failure, 1380t
 teeth discoloration, 236
- for balantidiasis, 1409
 for *C. pneumoniae* infection, 1174
 for *C. trachomatis* infection, 1171
 for cholera, 1064
 contraindications to, 1148
 for donovanosis, 198e-2t
 drug interactions of, 42t, 246e-2t, 246e-7t, 939t
 for ehrlichiosis, 1160
 for endemic treponematoses, 207e-2
 in *H. pylori* eradication, 1042t, 1922, 1922t
 indications for, 943
 for *Legionella* infection, 1020t
 for *M. hominis* infection, 926
 for *M. pneumoniae* infections, 1164
 for malaria, 1378t, 1380t
 for meibomitis, 199
 for parasitic infections, 246e-7t
 pharmacology of, 1380t
 for plague prophylaxis, 1075t
 for plague treatment, 1074t
 in pregnancy and lactation, 246e-7t, 937t
 for psittacosis, 1173
 for relapsing fever, 1148, 1148f
 resistance to, 931t, 934, 1064
 for Rocky Mountain spotted fever, 1157
 for syphilis, 1139, 1139t
 for tropical sprue, 1943
- Tetrahydrocannabinol (THC), 103e-25, 260t, 444e-4, 469e-2
- Tetrahydrofolate, 641, 641t
- Tetrahydrozoline, 473e-9t
- Tetralogy of Fallot
 clinical features of, 400, 1526–1527
 diagnosis of, 1527
 hypoxia in, 248
 murmur in, 51e-4f
 pathophysiology of, 1526
 repaired, ventricular arrhythmias in, 1495, 1495t
 treatment of, 1527
- Tetrathiomolybdate, 2520, 2520t, 2625
- TF (tissue factor), 401, 401f
- Tg. *See* Thyroglobulin (Tg)
- TGF- α . *See* Transforming growth factor- α (TGF- α)
- TGF- β . *See* Transforming growth factor- β (TGF- β)
- Thalamic Déjérine-Roussy syndrome, 2575
- Thalamus, 161, 211
- Thalassemia syndromes, 633, 633t, 637
 α thalassemia, 633, 637–638, 638t
 α thalassemia-1 trait, 638, 638t
 α thalassemia-2 trait, 638, 638t
 arthropathy in, 2242
 β thalassemia
 arthropathy in, 2242
 clinical features of, 637
 genetic considerations in, 433f, 435, 633
 β thalassemia intermedia, 637, 637f
 β thalassemia major, 637

- β thalassemia minor, 637, 2242
 β thalassemia trait, 637
 diagnosis and management of, 638
 differential diagnosis of, 628, 628t
 epidemiology of, 633
 genetic considerations in, 633
 HSCt for, 139e-2, 139e-5, 139e-5t
 jaundice in, 281
 malaria resistance and, 633, 1372
 peripheral blood smear in, 81e-1, 81e-4f
 prevention of, 638
 reticulocyte index in, 399
 silent, 638t
 splenomegaly in, 411
 treatment of, 638
 Thalassaemic structural variants, 638–639
Thalidomide
 action and targets of, 103e-19
 adverse effects of, 103e-19, 716–717
 neuropathy, 607, 2687t, 2688
 in pregnancy, 1128
 for ankylosing spondylitis, 2173
 for Behçet's syndrome, 2194
 for cough in idiopathic pulmonary fibrosis, 1712
 for erythema nodosum leprosum, 1128
 for lupus dermatitis, 2134
 for malignant pheochromocytoma, 2331
 for multiple myeloma, 103e-19, 716–717
 for primary myelofibrosis, 676
 for vascular ectasias, 277
 Thallium-201, 270e-3, 270e-3t
 Thallium poisoning, 203, 472e-3, 2688t, 2689
THAP1 gene mutations, 2620
 Thawing, for frostbite, 478e-4, 478e-4t
 THBR (thyroid hormone binding ratio), 2288
 THC (tetrahydrocannabinol), 103e-25, 260t, 444e-4, 469e-2
 Thecoma, of ovary, 594
 Thelarche, 2379, 2379t
Theophylline
 adverse effects of, 305, 324, 1677, 1706, 2547t
 for asthma, 1677, 1677t
 for COPD, 1706
 dosage of, 36
 drug interactions of, 42t, 1677, 1677t
 carbapenems, 939t
 ciprofloxacin, 246e-6t
 fluoroquinolones, 939t
 PPIs, 1921
 thiabendazole, 246e-3t, 246e-13
 metabolism of, 1677, 1677t
 overdose/poisoning with, 473e-8t
 for SA node dysfunction, 1469
 therapeutic monitoring of, 2766t
 Therapeutic diet, 95e-5
 Therapeutic index, 103e-5
 Therapeutic index/ratio/window, 32, 32f
 Therapeutic massage, 14e-1t, 14e-3
 Therapeutic touch, 14e-1t
 Thermoregulation, 124, 187, 478e-1, 479e-1
 THF starvation, 641
Thiabendazole
 action of, 246e-13
 adverse effects of, 246e-3t
 drug interactions of, 246e-3t, 246e-13
 indications for, 246e-3t
 pharmacology of, 246e-13
Thiamine (vitamin B₁), **96e-1**
 for alcohol-induced acidosis, 64e-9
 deficiency of. *See* Thiamine (vitamin B₁) deficiency
 food sources of, 96e-1
 functions of, 96e-1, 96e-2f
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference range for, 2766t
 structure of, 96e-2f
 supplements. *See* Thiamine (vitamin B₁) supplements
 toxicity of, 96e-3
Thiamine (vitamin B₁) deficiency. *See also* Beriberi
 acid-base disorders in, 318
 in alcoholism, 96e-1, 461
 cardiac manifestations of, 290e-1t, 290e-2
 clinical features of, 96e-1t, 96e-2–3, 463t, 2690. *See also* Wernicke's disease/encephalopathy
 delirium in, 169
 dementia in, 2607
 diagnosis of, 2690
 neuropathies in, 2690
 optic neuropathy in, 203
 risk factors for, 96e-1–2, 96e-1t
 treatment of, 96e-3, 2690
Thiamine-responsive megaloblastic anemia, 435e-1t
Thiamine (vitamin B₁) supplements, 96e-3
 for alcohol-induced acidosis, 319
 for alcohol withdrawal, 2727
 for dysmenorrhea, 338
Thiazide diuretics
 action of, 332e-4f, 332e-9
 adverse effects of, 326
 cutaneous, 349, 353, 379
 hypercalcemia, 2478
 hypokalemia, 306
 hyponatremia, 299, 301
 phototoxicity, 389t
 vasculitis, 382, 2193
 for chronic kidney disease, 1814
 drug interactions of, 246e-4t, 246e-6t
 for hyperkalemia, 312
 for hypertension, 1623
 for hypoparathyroidism, 2484
 for nephrogenic diabetes insipidus, 304, 2279
 for nephrolithiasis, 1870
 for paramyotonia congenita, 462e-18
 for pseudohypoadosteronism, 310
Thiazolidinediones
 action of, 2414t, 2415
 adverse effects of, 253t, 2392, 2414t
 contraindications to, 2414t
 for metabolic syndrome, 2454
 for NALFD, 2057
 for type 2 diabetes mellitus, 2414t, 2415, 2416
Thick-filament myopathy, 1784
Thienopyridines, 747–748. *See also* Clopidogrel; Prasugrel; Ticlopidine
Thiethylperazine, 103e-25, 260t
Thimerosal, 345
Thin basement membrane disease, 294, 1835t, 1847
6-Thioguanine, 103e-10t, 103e-14
Thiopental, 2558f
Thiopurine S-methyltransferase (TPMT), 34t, 39t, 40, 445
Thiopurine S-methyltransferase (TPMT) deficiency, 451
Thioridazine, 64t, 473e-8t, 473e-9t, 2721, 2722t
Thiotepa, 103e-12, 607t
Thiothixene, 2721, 2722t
Third disease. *See* Rubella
Third nerve palsy, 41e-1v, 1785
Third spacing, 298, 462
Third window, inner ear, 219
Thirst
 activation of, 295
 defects in regulation of, 2280
 in elderly, 303
 mechanism of, 332e-9, 332e-10f, 2275
Thogotoviruses, 1305, 1310t
Thomsen's disease, 462e-18
Thoracentesis, 474, 482e-1v, 1667
Thoracic aortic aneurysm, 1638–1639, 1639f. *See also* Aortic aneurysm
Thoracic lavage, as rewarming strategy, 478e-3
Thoracic meningioma, 2654f
Thoracic outlet, 122
Thoracic outlet syndromes, 123, 1646–1647
Thoracic spinal lesions, 2651
Thoracoscopy, medical, 1668
Thoracotomy, 1669
Thoratec LVAD, 1518
Thorium-232, in radiation bioterrorism, 263e-3t
3TC. *See* Lamivudine (3TC)
Three-dimensional printing, 92e-2, 92e-3v
Threonine kinase gene mutations, 359
Threonine metabolism disorders, 434e-3t
Throat swab, 150e-4t, 232
Thrombectomy, percutaneous mechanical. *See also* Percutaneous interventions
 for acute limb ischemia, 1646
 for ischemic stroke, 2561
 in STEMI, 297e-2, 297e-2f, 297e-2v
Thrombin
 in fibrin clot formation, 401, 401f, 746f
 inhibitors to, 732
 plasma-derived bovine and human, as hemostatic sealants, 740
 production of, 744
Thrombin-activatable fibrinolysis inhibitor (TAFI), 744
Thrombin generation test, 407
Thrombin time, 406t, 407, 733f, 733t
Thromboangiitis obliterans, 1645–1646, 1649
Thrombocytopenia, **725**
 approach to the patient, 726–727, 726f
 bleeding in, 404
 chemotherapy-related, 103e-24
 drug-induced, 727, 727t
 familial, with propensity to acute AML, 431t
 heparin-induced. *See* Heparin-induced thrombocytopenia
 history in, 726–727
 in HIV infection, 727, 728, 1262
 infection-induced, 727
 inherited, 726, 729
 in leptospirosis, 1142, 1144
 peripheral blood smear in, 725, 726f
 physical examination in, 726–727
 in pregnancy, 49
 refractory, 669t. *See also* Myelodysplasia
 in sepsis/septic shock, 1756
 skin manifestations of, 367
 in SLE, 2129t, 2131
 treatment of, 103e-24
Thrombocytopenia with absent radii, 729
Thrombocytopenic purpura
 autoimmune, 377e-4
H. pylori infection and, 1040
 immune, 413, 728–729
 thrombotic. *See* Thrombotic thrombocytopenic purpura (TTP)
Thrombocytosis
 erythromelalgia in, 1650
 essential. *See* Essential thrombocytosis (ET)
 etiology of, 676t, 730
 paraneoplastic, 612–613, 612t
Thromboembolism, 1646
Thromboembolic disease
 acute limb ischemia in, 1646
 in IBD, 1958
 in myocardial infarction, 1610
 pulmonary hypertension in, 1659
 renal, 1629–1630, 1630f
Thromboendarterectomy, 1636, 1645
Thrombogenesis, 745, 746f
Thrombolysis in Myocardial Infarction trial. *See* TIMI (Thrombolysis in Myocardial Infarction) trial
Thrombolytic therapy
 for acute limb ischemia, 1646
 for ischemic stroke, 2561–2562, 2562t
Thrombomodulin, 265e-2, 402
Thrombophilia, 407
Thrombophlebitis. *See* Venous thromboembolism (VTE)
Thromboplastin, 405
Thrombopoietin, 89e-3f, 275, 725, 741
Thrombopoietin receptor agonists, 728–729
Thrombopoietin therapy, 103e-24
Thrombosis, **403**
 approach to the patient, 404–405
 arterial. *See* Arterial thrombosis
 in central venous catheter, 98e-6–7
 dural sinus, 2572
 endothelium in, 265e-2
 epidemiology of, 274, 740
 in heparin-induced thrombocytopenia, 727
 history in, 404–405

- Thrombosis (*Cont.*):
 in inflammation, 276–277, 742
 laboratory evaluation of, 405–407
 in liver disease, 739–740, 739f, 739t
 pathophysiology of, 274–275, 275f, 740–741, 741f
 in polycythemia, 399
 renal vein, 1866
 risk factors for, 404, 405f, 405t
 venous. *See* Venous thromboembolism (VTE)
- Thrombotic microangiopathy, renal, **1863**
 in antiphospholipid antibody syndrome, 1865
 drug-related, 1864
 in HELLP syndrome, 1865
 in hemolytic-uremic syndrome. *See*
 Hemolytic-uremic syndrome (HUS)
 HIV-related, 1864
 HSCT-related, 1864, 1864t
 pathophysiology of, 1863
 radiation-related, 1864
 in sickle cell disease, 1865–1866. *See also*
 Sickle cell anemia
 in systemic sclerosis, 1864–1865. *See also*
 Systemic sclerosis (scleroderma)
- in thrombotic thrombocytopenic purpura.
See Thrombotic thrombocytopenic
 purpura
- Thrombotic thrombocytopenic purpura (TTP),
729
 diagnosis of, 729
 drug-induced, 1864
 epidemiology of, 133t
 genetic considerations in, 1848
 vs. hemolytic-uremic syndrome, 729, 1863
 in HIV infection, 1262
 idiopathic, 729, 1863–1864
 inherited, 729
 pathology and pathogenesis of, 265e-2, 729,
 729f, 1848, 1863
 peripheral blood smear in, 729
 renal involvement in, 1806t, 1848
 risk factors for, 1848
 skin manifestations of, 133t, 135, 368
 in SLE, 2133–2134
 treatment of, 729–730, 1848, 1864
- Thromboxane A₂
 in arterial thrombosis, 745, 746f
 in myocardial infarction, 1599
 in shock, 1745, 1746
- Thrush
 in cancer patient, 487
 clinical features of, 46e-1f, 46e-5f, 233, 240t,
 351, 1342
 etiology of, 233
 in HIV infection, 1255, 1256f
 treatment of, 233, 240t, 1344t
- “Thumbprint sign,” 234, 235f
- Thunderclap headache, 1785, 2597
- Thunderstorm asthma, 1671
- Thymectomy
 for myasthenia gravis, 123e-2, 2704
 for pure red cell aplasia, 123e-2
 for thymoma, 123e-2, 2704
- Thymic neuroendocrine tumors (carcinoids),
 564
- Thymic stromal lymphopoietin (TSLP), in
 asthma, 1672
- Thymidine, 426
- Thymidine phosphorylase deficiency, 431e-5t
- Thymidylate synthase, in lung cancer, 519–520
- Thymoma, **123e-1**
 in autoimmune diseases, 2348
 clinical features of, 123e-1, 664
 diagnosis of, 123e-1, 2703
 EBV infection and, 123e-2
 etiology of, 123e-2
 hypogammaglobulinemia in, 2111
 in myasthenia gravis, 123e-2, 2701, 2703
 paraneoplastic syndromes in, 372, 614t, 619
 pathology of, 123e-1–2, 123e-2t
 pemphigus and, 371, 372
 pure red cell aplasia and, 123e-2, 668
 staging of, 123e-1, 123e-1t
 treatment of, 123e-2
- Thymus
 cancer, paraneoplastic syndromes in, 609t
 in HIV infection, 1235, 1236
 in myasthenia gravis, 2701, 2703, 2704t
 primary defects of, 375e-1, 2108
- Thymus- and activation-regulated chemokine.
See TARC (CCL17)
- Thymus expressed chemokine (TECK), 372e-12t
- Thyroglobulin (Tg)
 in autoimmune thyroiditis, 372e-24t
 in carcinoma of unknown primary, 120e-2
 in congenital hypothyroidism, 2283, 2284t
 reference values for, 2763t
 in thyroid cancer follow-up, 2306–2307
 in thyrotoxicosis, 2288
- Thyroid acropachy, 2245, 2295, 2295f
- Thyroid and eye muscle shared protein, 372e-24t
- Thyroid cancer, **2303**
 anaplastic, 2304t, 2307
 carcinoma of unknown primary and, 120e-2t
 classification of, 2303t, 2304, 2304t
 deaths from, 467t
 diagnosis of, 2254t, 2289, 2289t
 follicular, 2304t, 2305–2307, 2305t
 genetic considerations in, 101e-3t, 2304–2305,
 2305t
 incidence of, 467t, 468f, 2254t, 2303, 2303t
 medullary. *See* Medullary thyroid carcinoma
 (MTC)
 papillary, 2303t, 2304t, 2305–2307, 2306f
 pathogenesis of, 2304–2305
 in patients with thyroid nodule, 2303, 2303t
 during pregnancy, 124e-1t, 124e-4
 radiation therapy-related, 623
 risk factors for, 2303, 2303t
 staging of, 2304, 2304t
 survival rates, 2306f
 treatment of, 2306–2307
- Thyroid cysts, 2303
- Thyroid dermatopathy, 2295, 2295f, 2297–2298
- Thyroidectomy
 for diffuse nontoxic goiter, 2301
 for Graves’ disease, 2297
 prophylactic, in *RET* mutations, 2341, 2341t
 for thyroid cancer, 2306, 2342
- Thyroid gland, **2283**
 amiodarone effects on, 2300
 anatomy and development of, 2283
 disorders of
 cancer. *See* Thyroid cancer
Candida infection, 489
 in HIV infection, 1259–1260
 hypertension in, 1620
 hyperthyroidism. *See* Hyperthyroidism
 hypothyroidism. *See* Hypothyroidism
 lymphoma, 2307
 nodular. *See* Thyroid nodule
 during pregnancy, 48–49, 124e-4, 124e-5t
 radiation-induced, 623
 thyroiditis. *See* Thyroiditis
 thyrotoxicosis. *See* Thyrotoxicosis
- hormones of. *See* Thyroid hormone(s)
- laboratory studies of
 radioiodine uptake and thyroid scanning,
 2288
 Tg levels, 2288
 thyroid hormone measurement, 2288
 TPO antibodies, 2288
 ultrasonography, 2289, 2289f, 2289t
- pain in, differential diagnosis of, 2298
- physical examination of, 2287
- regulation of, 2283–2284, 2284f
- Thyroid hormone(s). *See also* Thyroxine (T₄);
 Triiodothyronine (T₃)
 action of, 2286–2287, 2287f
 half-lives of, 400e-2
 laboratory evaluation of, 2288, 2763t
 receptor binding of, 2286–2287, 2286f
 resistance to, 435, 2286t, 2287
 structure of, 2283f
 synthesis of, 2284–2285, 2284f, 2285f
 transport and metabolism of, 2285–2286,
 2286t
 unbound, 2288
- Thyroid hormone binding proteins, 2286
- Thyroid hormone binding ratio (THBR), 2288
- Thyroid hormone receptors (TRs), 2286–2287,
 2286f
- Thyroiditis, **2298**
 acute, 2298, 2298t
 atrophic, 2290
 autoimmune. *See* Hashimoto’s thyroiditis
 (autoimmune hypothyroidism)
 chronic, 2298t, 2299
 destructive, 2298
 drug-induced, 2299
 etiology of, 2298t
 postpartum, 2299
 radiation therapy-related, 623
 Riedel’s, 391e-1t, 2299
 silent (painless), 2299
 subacute (de Quervain’s), 239, 2298–2299,
 2298f, 2298t
- Thyroid nodule
 approach to the patient, 2307–2308, 2308f
 benign, 2303, 2303t
 evaluation of, 2254t, 2288–2289, 2289f,
 2289t
 prevalence of, 2254t
 solitary, hyperfunctioning, 2302–2303,
 2303f
 thyroid cancer and, 2303, 2303t. *See also*
 Thyroid cancer
- Thyroid peroxidase (TPO)
 in autoimmune hypothyroidism, 372e-24t
 in thyroid development, 2283, 2284t
 in thyroid hormone synthesis, 2285
- Thyroid peroxidase (TPO) antibodies, 2288,
 2294
- Thyroid response elements (TREs), 2287, 2287f
- Thyroid scintigraphy, 2288
- Thyroid-stimulating hormone (TSH), **401e-4**
 action of, 401e-4
 assessment of, 400e-6
 autoantibodies to, 377e-4
 deficiency of, 2259t
 ectopic production of, 609t
 expression and regulation of, 401e-1t
 laboratory evaluation of, 2258t, 2288, 2763t
 pituitary adenoma secretion of, 2274, 2298
 in pregnancy, 124e-4, 124e-5t
 secretion of, 401e-4
 suppression therapy, 2306
 synthesis of, 401e-4
- Thyroid-stimulating hormone receptor (TSH-R)
 in congenital hypothyroidism, 2284t
 in hyperfunctioning thyroid solitary nodule,
 2302–2303, 2302f
 mutations of, 2284t, 2302–2303, 2302f
 in thyroid cancer, 2305t
 in thyroid development, 2283
- Thyroid-stimulating immunoglobulins (TSI),
 2288, 2294
- Thyroid storm (thyrotoxic crisis), 48, 2297
- Thyroid transcription factor-1 (TTF-1)
 in carcinoma of unknown primary, 120e-2
 in congenital hypothyroidism, 2283, 2284t
 in lung cancer, 120e-2, 508
- Thyrotoxicosis, **2293**. *See also* Graves’ disease
 amiodarone-induced, 2300
 apathetic, 2294
 clinical features of, 2294–2295, 2294t, 2295f
 definition of, 2293
 diagnosis of, 2295–2296, 2295f
 etiology of, 2293t, 2298
 fetal/neonatal, 2297
 hypoxia in, 248
 palpitations in, 254
 primary hyperthyroidism and, 2293
 subclinical, 2301
 without hyperthyroidism, 2293
- Thyrotoxicosis factitia, 2288, 2298
- Thyrotoxic periodic paralysis (TPP). *See*
 Hypokalemic periodic paralysis
 (HypoKPP)
- Thyrotropin-binding inhibitory immunoglobulin
 (TBII), 2290, 2294
- Thyrotropin receptor, 372e-24t

- Thyrotropin-releasing hormone (TRH), 401e-2, 401e-4
- Thyroxine (T₄)
 characteristics of, 2286f
 free, 2286, 2288
 half-life of, 400e-2
 laboratory evaluation of, 2286f, 2288, 2763t
 in pregnancy, 124e-4, 124e-5t
 receptor binding of, 2287, 2287f
 in sick euthyroid syndrome, 2299
 structure of, 2283f
 in subacute thyroiditis, 2298–2299, 2298f
 total, 2288
 transport and metabolism of, 2285–2286
- Thyroxine-binding globulin (TBG), 2285, 2286, 2286t, 2288
- Thyroxine-binding prealbumin. *See* Transthyretin
- Thyroxine therapy
 for adult hypopituitarism, 2259t
 adverse effects of, 254
 for hypothyroidism in pregnancy, 49
- TIA. *See* Transient ischemic attack (TIA)
- Tiagabine
 adverse effects of, 2555t
 for anxiety disorders, 2711
 for epilepsy, 2555t
 overdosage/poisoning with, 473e-10t
 for stiff-person syndrome, 618
- Tibetan medicine, 14e-1t
- Tibia, Paget's disease of, 436e-3f
- Tibial muscular dystrophy (Udd's distal myopathy), 462e-12, 462e-13t
- Tic(s), 2619t, 2623
- Ticagrelor
 action of, 747
 adverse effects of, 748
 for cardiac catheterization pretreatment, 1461
 dosage of, 748
 indications for, 747–748
 for ischemic heart disease, 1590
 management before endoscopic procedures, 1889t
 for myocardial infarction, 1596, 1598, 1598t, 1606
 in PCI, 296e-1, 296e-2
- Ticarcillin, 306
- Ticarcillin-clavulanate
 for anaerobic bacterial infections, 1101t
 for bite-wound infections, 167e-2
 indications for, 938
 for peritonitis, 848
 resistance to, 1101
 for *S. maltophilia* infection, 1045t
- Tic douloureux. *See* Trigeminal neuralgia
- Tick(s)
Amblyomma spp., 1152, 1154
 bite characteristics of, 2745
Ixodes. *See* *Ixodes* spp. ticks
 Lyme disease prophylaxis following bite, 1153, 2745
Ornithodoros spp., 1146, 1147f
 removal of, 2745
 in Rocky Mountain spotted fever, 1154
- Tick-borne diseases
Babesia infections. *See* *Babesia* spp. infections
 Colorado tick fever, 1306t, 1319
 Crimean-Congo hemorrhagic fever, 1308t, 1321
 ehrlichioses, 1155t, 1159–1160, 1159f
 encephalitis, 1307t, 1315–1316
 human granulocytic anaplasmosis, 1152, 1155t, 1159f, 1160–1161
 Lyme borreliosis. *See* Lyme borreliosis (Lyme disease)
 relapsing fever. *See* Relapsing fever rickettsial. *See* Rickettsial diseases
 Rocky Mountain spotted fever. *See* Rocky Mountain spotted fever (RMSF)
 Southern tick-associated rash illness, 129t, 134, 1152
 tick-borne lymphadenopathy, 1155t
 tick granuloma, 2745
 tick paralysis, 989, 2745
- Ticlopidine
 action of, 746f
 adverse effects of, 729, 1864, 2218t, 2569
 management of
 before endoscopic procedures, 1889t
 before lumbar puncture, 443e-1
 for stroke/TIA prevention, 2569–2570
- Tie2, 102e-14f, 102e-15
- Tietze syndrome, 2246
- TIG (tetanus immune globulin), 167e-3, 986
- Tigecycline
 for *Acinetobacter* infections, 1037, 1037t
 action of, 930, 931t, 943, 1027
 adverse effects of, 943
 for anaerobic bacterial infections, 1101
 distribution of, 935
 for *Enterobacter* infections, 1034
 indications for, 940t, 943
 for *Klebsiella* infections, 1033
 for *Legionella* infections, 1020, 1020t
 for *Nocardia* infections, 1087
 in pregnancy and lactation, 937t
 resistance to
 in *Acinetobacter*, 1037
 in anaerobic bacteria, 1101
 in gram-negative bacteria, 1026, 1027
 mechanisms of, 931t, 934
 rates of, 940t
 for staphylococcal infections, 962
- Tiger snakes, 2734. *See also* Snakebites
- "Tight-building" syndrome, 1693
- Tight epithelia, 332e-3
- Tight junction, 332e-3
- Tilt table test, 148, 2641, 2641t
- Tiludronate, 426e-3, 426e-3t
- Time-of-flight (TOF) imaging, 440e-7
- TIMI (Thrombolysis in Myocardial Infarction) trial
 frame count, 1605
 grading system, 1605
 risk score, 1595, 1596f
- Timolol
 adverse effects of, 1671
 genetic variations in response to, 39t, 40
 for ischemic heart disease, 1589t
 metabolism of, 34t
 overdosage/poisoning with, 473e-9t
- Tincture of opium, 272
- Tinea, 353, 353t, 355, 1357, 1358t. *See also* Dermatophytosis
- Tinea capitis
 alopecia in, 354, 355t
 clinical features of, 355t, 1357
 diagnosis of, 343
- Tinea corporis, 76e-10f, 349, 1357
- Tinea cruris, 342f, 350, 1357
- Tinea manuum, 76e-10f
- Tinea pedis, 76e-10f, 342f, 350, 1357
- Tinea unguium, 76e-10f, 350
- Tinea versicolor, 350
 clinical features of, 341t, 350, 350t, 357, 357t, 358t, 1357
 diagnosis of, 341, 344, 358t
 etiology of, 350, 350t
 treatment of, 350, 350t, 358t, 1357
- Tinel's sign, 2221
- Tin exposure, 472e-3
- Tingling, 158. *See also* Sensation
- Tinidazole
 action of, 246e-13
 adverse effects of, 246e-5t
 for amebiasis, 1367, 1367t
 for bacterial vaginosis, 875
 drug interactions of, 246e-5t
 for *Giardia* infections, 1406
 in *H. pylori* eradication, 1042t
 pharmacology of, 246e-13
 in pregnancy and lactation, 246e-5t
 for *T. vaginalis* infections, 871, 873t, 1409
- Tinnitus, 220, 224, 2586
- Tinnitus masker, 224
- TINU (tubulointerstitial nephritis with uveitis), 1857–1858, 1858f
- Tinzaparin, 1635t
- Tiopronin, 435e-2, 1871
- Tiotropium, 1676, 1705
- TIP39, 2468
- Tipranavir, 1276f, 1280f
- TIPS. *See* Transjugular intrahepatic portosystemic shunt (TIPS)
- TIRAP, 145e-5, 145e-6f
- TIR domains, 145e-5, 145e-6f
- Tirofiban
 action of, 746f, 748–749
 adverse effects of, 727t, 749
 dosage of, 749
 genetic variation in response to, 277t, 743t
 indications for, 749
 for myocardial infarction, 1598t
 in PCI, 296e-1
 pharmacology of, 749t
- Tissue engineering, 92e-1
 future directions in, 92e-3–4
 isolation and growth of cells in, 92e-1
 level of complexity in, 92e-1–2, 92e-2f
 scaffolds for, 92e-1
 strategies for, 92e-2, 92e-3v
 vascularization and innervation in, 92e-1
- Tissue factor, 745
- Tissue factor (TF), 401, 401f
- Tissue factor pathway inhibitor
 antithrombotic actions of, 402, 402f, 750
 for sepsis/septic shock, 1758
- Tissue invasion, by pathogens, 145e-7–8
- Tissue nonspecific alkaline phosphatase (TNSALP), 426e-5
- Tissue plasminogen activator (tPA)
 for acute limb ischemia, 1646
 arterial thrombotic disease and, 744
 contraindications to, 1605
 in fibrinolysis, 402–403, 402f
 for myocardial infarction, 1605
 recombinant. *See* Alteplase (rtPA)
- Tissue stem cells, 453
- Tissue thromboplastin inhibition test, 406
- Tissue transglutaminase, 372e-24t, 2763t
- Tissue tropism, of pathogens, 145e-8–9
- Titanium dioxide, 1691
- Titin, 265e-5
- Tivantinib, 551
- Tizanidine, 473e-9t, 473e-10t, 2660, 2672
- TK gene, 91e-4
- TLC (total lung capacity), 306e-1f, 306e-2, 306e-2f, 306e-5f
- TLR7 gene, in HIV infection, 1240t
- TLR (toll-like receptor) pathway deficiencies, 2106
- TLRs. *See* Toll-like receptors (TLRs)
- T lymphocytes. *See* T cell(s)
- TMC207 (bedaquiline), 205e-7
- TMP-SMZ. *See* Trimethoprim-sulfamethoxazole (TMP-SMZ)
- TNF. *See* Tumor necrosis factor (TNF)
- TNF-R1 gene, 372e-8t
- TNFRSF11A/B genes, 426e-1, 2215
- TNM staging, 470
 of appendiceal NETs, 560, 561t
 of bladder cancer, 575, 576f
 of bone sarcoma, 119e-3, 119e-3t
 of breast cancer, 526, 527t
 of colorectal cancer, 541–542, 542f
 of head and neck cancer, 504, 504f
 of hepatocellular carcinoma, 546
 of lung cancer, 513–514, 514f, 516t
 of pancreatic cancer, 555, 556f
 of pNETs, 560, 560t
 of prostate cancer, 582, 582f, 582t
 of soft tissue sarcoma, 119e-2, 119e-2t
- TNSALP (tissue nonspecific alkaline phosphatase), 426e-5
- Tobacco grower's lung, 1681t
- Tobacco use. *See also* Smoking
 global burden of disease caused by, 17
 types of, 2730–2731
 in women, 6e-5
- Tobramycin
 action of, 930, 931t
 adverse effects of, 942
 for community-acquired pneumonia, 808t
 for health care-associated pneumonia, 767t

- Tobramycin (*Cont.*):
 indications for, 940t
 for infective endocarditis, 823
 inhalation solution, for bronchiectasis prophylaxis, 1696
 for *P. aeruginosa* infections, 808t
 resistance to, 931t, 934, 940t
 for sepsis/septic shock, 1757t
 therapeutic monitoring of, 2766t
- Tocilizumab
 action of, 372e-30, 2147
 adverse effects of, 2147, 2147t
 for giant cell arteritis, 2189
 for rheumatoid arthritis, 2147, 2147t
- α -Tocopherol. *See also* Vitamin E
 for lung cancer prevention, 477
 for prostate cancer prevention, 478
- Todd's paralysis, 2542
- Toe(s)
 gout of, 436e-2f
 osteoarthritis of, 2229f
- Tofacitinib
 action of, 2147
 adverse effects of, 2147, 2147t
 for IBD, 1962
 for psoriatic arthritis, 2177
 for rheumatoid arthritis, 2147, 2147t
- TOF (time-of-flight) imaging, 440e-7
- To-fro murmur, 51e-6, 51e-7f
- Togaviruses, 214e-2t, 214e-3f, 214e-4, 1305. *See also* Alphaviruses
- Togavirus infections. *See* Alphavirus (togavirus) infections
- Tolazamide, 2413
- Tolbutamide, 284, 2413
- Tolcapone, 2615, 2616t
- Tolerable upper levels, 95e-3
- Tolerance, definition of, 61, 372e-2
- Tolfenamic acid, 2592t
- Toll-like receptor (TLR) pathway deficiencies, 2106
- Toll-like receptors (TLRs)
 in adaptive immune response, 372e-2, 372e-6t
 in autoimmunity, 377e-1-2
 fever and, 125
 in glomerular disease, 1833
 in immune response, 950
 in inflammatory response, 145e-5, 145e-6f
 ligands and functions of, 372e-5t
 in platelets, 276, 742
 in rheumatoid arthritis, 2141, 2142f
 in sepsis and septic shock, 1752
 signaling pathways of, 372e-5f
- Tolosa-Hunt syndrome, 207, 1352, 2650
- Tolterodine, 81, 2672
- Tolvaptan
 adverse effects of, 302, 511
 for hypervolemic hyponatremia, 2283
 for polycystic kidney disease, 1853
 for SIADH, 64e-5, 302, 511, 610, 2282
- Tonate virus, 1306t
- Tone, 154
- Tongue
 alterations of, 241t
 bald, 241t
 black hairy, 240t
 fissured (scrotal), 46e-5f, 241t
 geographic, 46e-3f, 241t, 342f
 microbiome of, 86e-5, 86e-6f
 paralysis of, 2649
 raspberry, 241t
 strawberry, 241t
 ulcer on, 46e-4f
 white coated, 46e-5f. *See also* Thrush
- Tongue worms, 2747
- Tonic-clonic seizures, 2543, 2550-2551, 2551t.
See also Seizure(s)
- Tonicity, 332e-9, 332e-10f
- Tonic pupil, 196
- Tonsillectomy, 230, 404
- Tonsillitis, 235
- Tonsils, orange, 1443
- Tophi, 364t, 365, 2225f
- Topiramate
 adverse effects of, 215, 2396, 2555t, 2556, 2594t
 for cluster headache prevention, 2595t
 for epilepsy, 2555t, 2556
 for hemicrania continua, 2596
 for migraine prevention, 2594t
 overdose/poisoning with, 473e-10t
 for pain, 61
 pharmacology of, 2555t
 with phentermine, for obesity, 2396-2397, 2397t
 for posttraumatic headache, 111
 for PTSD, 2712
 for raised CSF pressure headache, 110
 for SUNCT/SUNA prevention, 2596
 "Top of the basilar" syndromes, 2578
- Topoisomerase I (Scl-70), 103e-13, 372e-24t, 2157t
- Topoisomerase II, 103e-13
- Topoisomerase inhibitors, 103e-9-10t, 103e-13, 623, 670, 678. *See also* specific drugs
- Topoisomerase IV, 932
- Topology, of biologic system, 87e-1-2, 87e-2f
- Topotecan
 action of, 103e-13
 adverse effects of, 103e-10t, 103e-13
 for Ewing's sarcoma, 119e-4
 for lung cancer, 522
 for ovarian cancer, 594
- TOR (target of rapamycin), 94e-5
- Toroviruses, 1286f, 1289
- Torsades de pointes. *See* Ventricular tachycardia, polymorphic
- Torsemide, 322, 1510t, 1763
- Torsin A, 2620
- Torticollis, 2620
- TOS (true neurogenic thoracic outlet syndrome), 122
- Toscana virus, 894, 1310t, 1318
- Total artificial heart (TAH), 1518
- Total-body water, 64e-4, 332e-10, 332e-10f
- Total iron-binding capacity
 in anemia evaluation, 393
 causes of abnormal values, 464t
 in hypoproliferative anemia, 398, 630t
 in iron-deficiency anemia, 627f
 in microcytic anemia, 628t
 normal, 397, 627
 in nutritional assessment, 462, 464t
- Total lung capacity (TLC), 306e-1f, 306e-2, 306e-2f, 306e-5f
- Total parenteral nutrition. *See* Parenteral nutrition
- Totipotent stem cells, 454, 454f
- Touch localization test, 159
- Touch sensation, 159, 160t, 2539
- Touraine-Solente-Golé syndrome, 2244. *See also* Hypertrophic osteoarthropathy
- Tourette's syndrome, 2623
- Tourniquet, for snakebite, 2735
- Toxic adenoma, 2302-2303, 2302f
- Toxic epidermal necrolysis (TEN)
 vs. AGEF, 382
 clinical features of, 361, 361t, 382-383, 384t
 drugs associated with, 361
 drugs causing, 383
 epidemiology of, 131t
 etiology of, 131t
 genetic considerations in, 378
 immune pathways of, 378, 378t
 nondrug causes of, 382-383, 384t
 prevention of future episodes, 384
 skin manifestations of
 blistering eruptions, 382-383, 383f, 384t
 epidermal sloughing, 25e-7f, 131t, 134
 vs. staphylococcal scalded skin syndrome, 134, 362, 829
 treatment of, 383
- Toxic erythema of chemotherapy, 380
- Toxic granulations, neutrophils, 415
- Toxic megacolon
 in *Shigella* infections, 1056-1057
 treatment of, 1058, 1953
 in ulcerative colitis, 1953
- Toxic multinodular goiter, 2302
- Toxic oil syndrome, 2155
- Toxic shock syndrome (TSS)
 clostridial, 991t, 994-995
 diarrhea in, 267
 etiology of, 782
 focal infections and, 784
 skin manifestations of, 362t, 363, 782, 1755
 staphylococcal
 clinical features of, 130t, 363, 782, 959, 959t
 diagnosis of, 959
 pathogenesis of, 124, 372e-19, 955, 956
 rash in, 130t, 134
 tampon-associated, 959
 treatment of, 963
- streptococcal
 case definition of, 968, 968t
 clinical features of, 130t, 134, 363, 782, 968, 968t
 diagnosis of, 968-969
 epidemiology of, 130t
 etiology of, 130t, 968-969
 rash in, 130t, 134
 treatment of, 965t, 969t
- Toxic sponges, 2741
- Toxidrome, 473e-1
- Toxins. *See also* specific toxins
 bacterial, 145e-9
 hepatic effects of, 2023, 2025t, 2026t
 neurologic effects of, 2688t, 2689
 pulmonary effects of, 1688t, 1691-1692, 1692t
- Toxocara canis*, 245e-3t, 765t, 1411. *See also* Larva migrans, visceral
- Toxocara cati*, 245e-3t, 765t, 1411. *See also* Larva migrans, visceral
- Toxoplasma gondii*, 245e-4t, 245e-5t, 1398-1399, 1399f
- Toxoplasma gondii* infection (toxoplasmosis), 1398
 asymptomatic, 1401
 in cancer patient, 488, 488t
 clinical features of
 cardiac, 1400, 1471t, 1561
 chronic meningitis, 908t
 CNS, 903, 1265-1266, 1266f, 1401-1402, 1402f
 gastrointestinal, 1400
 in immunocompetent patient, 1401
 in immunocompromised patient, 1401-1402, 1402f, 1561
 lymphadenopathy, 409, 1400, 1401
 myelopathy, 2657
 ocular, 40e-6f, 1400, 1402, 1404
 pulmonary, 1400
 renal, 1849-1850
 weakness, 156
 congenital, 1398, 1399-1400, 1402, 1404
 diagnosis of, 245e-4t, 245e-5t, 245e-6, 245e-6t, 1190t, 1402-1404
 differential diagnosis of, 1402, 1403t
 epidemiology of, 245e-4t, 1399
 etiology of, 1398-1399, 1399f
 in HIV infection, 1251-1252t, 1265-1266, 1266f, 1401-1402, 1402f, 1561
 immune response to, 1401
 pathogenesis of, 1400
 pathology of, 1400-1401
 in pregnancy, 1399-1400, 1401, 1404
 prevention of, 1404-1405
 prophylaxis for, 925t, 1404
 transmission of, 1399-1400
 in transplant recipient, 920t, 921, 925t, 926
 treatment of, 246e-12, 246e-13, 903, 1404
- TP53 gene mutations
 in bladder cancer, 576
 genetic testing for, 448
 in lung cancer, 509
 tumors associated with, 101e-6t
- tPA. *See* Alteplase (rtPA); Tissue plasminogen activator (tPA)
- TPIT gene, 2256

- TPMT (thiopurine S-methyltransferase), 34t, 39t, 40, 445
- TPMT (thiopurine S-methyltransferase) deficiency, 451
- TPN (total parenteral nutrition). *See* Parenteral nutrition
- TPO. *See* Thyroid peroxidase (TPO)
- TPP (thyrotoxic periodic paralysis). *See* Hypokalemic periodic paralysis (HypoKPP)
- TPPA (*T. pallidum* particle agglutination) test, 1137, 1137f
- TR (thyroid hormone receptor), 2286–2287, 2286f
- TR α - gene mutations, 2287
- TR β - gene mutations, 2287
- Trabeculectomy, 205
- Trace minerals. *See* Mineral(s)
- Trachea, engineered tissue for replacement of, 92e-3t
- Tracheal intubation, 324, 1776
- Tracheobronchitis
 - Aspergillus*, 1346
 - HRSV, 1206
 - parainfluenza virus, 1207
- Tracheoesophageal fistula, 533
- Tracheomalacia, 306e-2, 308e-13f
- Tracheostomy, 1744
- Trachoma, 199, 1172
- Trade-off hypothesis, 333e-3, 333e-3f
- Traditional Chinese medicine, 14e-1t, 14e-3
- TRAF6, 145e-6f
- TRAIL, 102e-8, 102e-9f, 102e-10
- TRALI (transfusion-related acute lung injury), 138e-5
- TRAM, 145e-5
- Tramadol
 - adverse effects of, 2547t, 2682t
 - for back pain, 119
 - for neuropathy, 2682t
 - overdosage/poisoning with, 473e-13t
 - for pain, 91t
- Trametinib
 - action and targets of, 102e-5t, 103e-16f, 103e-17t, 103e-19
 - adverse effects of, 103e-17t
 - for melanoma, 103e-19, 498, 499
- “Tram tracks” sign, 62e-3f, 1695, 1841
- Trandolapril, 1512t
- Tranexamic acid, 731–732, 734, 738
- Transaminases, 752, 1997, 1998
- Transbronchial biopsy, 1667
- Transbronchial needle aspiration, 1668
- Transcatheter aortic valve replacement (TAVR). *See* Aortic valve replacement, transcatheter
- Transcobalamin deficiency, 375e-2
- Transcobalamin I, 640
- Transcobalamin II, 640
- Transcortical aphasia, 177t, 178
- Transcranial magnetic stimulation, 2717
- Transcription coactivator p75, 372e-24t
- Transcription factors
 - in asthma, 1674
 - defects in, diseases caused by, 431t
 - gene expression and, 428–429, 428f
 - in signal transduction in cancer cells, 102e-3–7
 - in stem cells, 89e-4, 455
- Transcriptome, 427, 769t
- Transcriptomics, 428–429
- Transdifferentiation, of stem cells, 90e-2, 454–455
- Transduction, 91e-1
- Transesophageal echocardiography (TEE)
 - in aortic dissection, 1641
 - in aortic stenosis, 270e-17, 1531
 - in atrial fibrillation, 1487
 - in atrial septal defect, 1521, 1522f
 - indications for, 270e-2
 - in infective endocarditis, 820, 820f, 821f
 - in mitral regurgitation, 1545
 - in mitral valve prolapse, 1547
 - in murmur evaluation, 51e-8
- principles of, 270e-2
- in pulmonary stenosis, 1526
- in valvular heart disease, 1540
- Transfalcial herniation, 1772, 1772f
- Transferase variants, 40
- Transfer dysphagia, 254, 1895. *See also* Dysphagia
- Transferrin
 - diferric, 625
 - iron content of, 625t
 - in iron metabolism, 625, 625f
 - laboratory measurement of. *See* Total iron-binding capacity
 - monoferric, 625
- Transferrin-binding proteins, 1004
- Transferrin receptor protein, 628
- Transferrin receptors, 625, 2515
- Transferrin saturation, 397, 398, 625f
- Transformation, bacterial, 933
- Transforming growth factor- α (TGF- α)
 - in gastroduodenal mucosal defense and repair, 1912, 1913f
 - in Ménétrier’s disease, 1932
 - in tumor angiogenesis, 102e-15
- Transforming growth factor- β (TGF- β), 87e-3
 - in ankylosing spondylitis, 2170
 - in cardiac development, 265e-11
 - in COPD, 1700
 - cytokine family, 372e-14t
 - in diabetic nephropathy, 2424
 - in familial pulmonary arterial hypertension, 87e-3
 - in hypercalcemia of malignancy, 609
 - mutations in, 87e-3
 - in oncogene signaling pathway, 102e-11f
 - in preeclampsia, 45
 - in radiation-induced fibrosis, 622
 - source, target, and biologic activity of, 372e-11t
 - in thyroid hormone synthesis, 2285
- Transfusion(s), 138e-2
 - adverse reactions to, 138e-3
 - allergic, 138e-5
 - alloimmunization, 138e-5
 - anaphylactic, 138e-5
 - delayed hemolytic and serological, 138e-4
 - electrolyte toxicity, 138e-5
 - febrile nonhemolytic, 138e-4–5
 - fluid overload, 138e-5
 - graft-versus-host disease, 138e-5
 - hemosiderosis, 639
 - hypotension, 138e-5–6
 - hypothermia, 138e-5
 - immunomodulation, 138e-6
 - incidence of, 138e-3t
 - infectious, 138e-3t, 138e-6, 1077
 - iron overload, 138e-5, 631, 667
 - lung injury, 138e-5
 - purpura, 138e-5
 - alternatives to, 138e-6
 - for AML, 685
 - for anemia, 399
 - for aplastic anemia, 667
 - autologous, 138e-6
 - blood components for, 138e-2–3, 138e-2t
 - for chemotherapy-associated anemia/thrombocytopenia, 103e-24
 - for DIC, 738
 - for hemoglobinopathies, 637
 - for hemophilia, 734
 - for hypovolemic shock, 1749–1750
 - for iron-deficiency anemia, 628
 - massive, hyperkalemia in, 310
 - metabolic alkalosis in, 322
 - packed red blood cell, 138e-2–3, 138e-2t
 - for AML, 685
 - for aplastic anemia, 667
 - for hypoproliferative anemia, 631
 - for iron-deficiency anemia, 628
 - platelet. *See* Platelet transfusion
 - pretransfusion testing, 138e-2
 - reducing need for, 138e-6
 - risks of, 631
 - for sepsis/septic shock, 1757- for sickle cell anemia, 636
- for thalassemia, 638
- whole blood, 138e-2
- Transfusion-related acute lung injury (TRALI), 138e-5
- Transgenic animal models, 432
- Transhepatic artery chemoembolization (TACE), 549, 550
- Transient acantholytic dermatosis, 341t
- Transient aplastic crisis, 1196, 1196t
- Transient elastography, 1999
- Transient global amnesia, 183, 2608
- Transient ischemic attack (TIA), 2568
 - in basilar artery distribution, 2578
 - definition of, 2559
 - at high altitudes, 476e-4
 - prevention of, 2569–2570
 - in SLE, 2129, 2129t
 - small-vessel, 2565
 - stroke risk following, 2568, 2569t
 - treatment of, 2560, 2560f
 - visual symptoms of, 204
- Transient myeloproliferative disorder (TMD), 135e-7
- Transient receptor potentials (TRPs), 444e-9
- Transition (mutation), 432
- Transit volume, in radiation therapy, 103e-4
- Transjugular intrahepatic portosystemic shunt (TIPS)
 - for ascites, 288, 2065
 - for variceal bleeding, 277, 2064, 2064f
- Translocation, chromosome, 83e-5–6, 83e-5f, 101e-3, 101e-3t
- Transmissibility, of pathogens, 145e-10
- Transmission disequilibrium test, 442t
- Transplantation. *See specific types*
- Transplant elbow, 926
- Transplant recipient. *See also* Immunocompromised patient
 - abdominal pain in, 105
 - immunization of, 928–929, 929t
 - infections in, 919. *See also under specific transplant procedures*
 - adenovirus, 1208
 - CMV, 1192
 - CNS, 463e-3
 - HRSV, 1206
 - Legionella*, 1015
 - parainfluenza, 489, 1207
 - rhinovirus, 1203
 - lipoprotein disorders in, 2441
 - pretransplantation evaluation of, 919
 - skin cancer in, 351, 388, 500, 1831
 - travel by, 929- Transporters associated with antigen processing (TAP) proteins, 372e-19, 373e-3–4
- Transposition of the great arteries
 - AV conduction block in, 1473
 - clinical features of, 1527
 - hypoxia in, 248
 - pathophysiology of, 1527
 - treatment of, 1527
- Transrectal ultrasonography, 582
- Transsphenoidal surgery
 - for acromegaly, 2270
 - complications of, 2263
 - for Cushing’s syndrome, 2272, 2318
 - for pituitary tumors, 2263, 2263f, 2272–2273
- Transtentorial herniation, 1772, 1772f
- Transthoracic (transtracheal) echocardiography, 820, 821f
- Transthyretin (TTR)
 - in amyloidosis, 271e-6f, 719, 720t
 - causes of abnormal values, 464t, 2286t
 - in nutritional assessment, 462, 464t
- Transtubular K⁺ concentration gradient (TTKG), 64e-7, 307, 311, 311f
- Transvaginal ultrasound, 482, 593
- Transverse sinus thrombosis, 906
- Transversion (genetic), 432
- Tranylcypromine, 473e-8t
- TRAP (tartrate-resistant acid phosphatase), 426e-4

- TRAPS (tumor necrosis factor- α receptor-associated periodic syndrome), 141, 422, 2213t, 2214–2215
- Trastuzumab
action and targets of, 102e-4, 102e-5t, 102e-10, 103e-21, 103e-21t
adverse effects of
cardiotoxicity, 103e-13, 103e-21, 103e-26, 620t, 621, 1563
hypersensitivity reaction, 1796, 1798
for breast cancer, 102e-4, 102e-5t, 102e-10, 528, 529, 529t, 530
drug interactions of, 103e-13
for esophageal cancer, 533
genetic variations in response to, 39t, 41
for ovarian cancer, 594
- Traube's sign, 411, 1536
- Trauma
aortic, 289e-3, 1637, 1638
back pain following, 117
bile duct, 2084
cardiac. *See* Cardiac trauma
cervical spine, 121–122
chest wall, 51e-2
head-related. *See* Head injury
hypothermia in, 478e-1
pneumothorax in, 1719
testicular, 2365
travel-associated, 797
- Traumatic brain injury (TBI). *See* Head injury
- Traumatic shock, 1748t, 1750
- Travel
to developing countries, health problems during, 793–794, 794f
by diabetic patient, 799
by disabled person, 799
fitness to fly, 794
food and water precautions during, 796
gastrointestinal illness during. *See* Travelers' diarrhea
health advice for, 794
for health care purposes, 799
by heart disease patient, 799
to high altitudes, 476e-5. *See also* High-altitude illness
by HIV-infected person, 798
immunizations for, 794–796, 795t
infectious disease exposure and, 763
injury during, 797
by lung disease patient, 799
malaria prevention during, 796, 796t, 1382–1384, 1383t
medical kit for, 797
in pregnancy, 797–798
problems after return, 800–801f, 800–802, 802t
sexually transmitted disease and, 797
by transplant recipient, 929
- Travelers' diarrhea, 796
after return, 797
epidemiology of, 266, 855
etiology of, 266, 797, 855, 855t
in HIV infection, 798
in pregnancy, 797
prevention of, 268, 797, 857
self-treatment of, 796–797
treatment of, 856–857, 857t
- Trazodone
adverse effects of, 64, 2710t
for depression, 64, 2710t
for insomnia, 65, 191
for PTSD, 2712–2713
- Treacher Collins syndrome, 222t
- "Tree-in-bud" pattern, 308e-12f, 1695
- T regulatory cells (T-regs), 377e-2, 1237. *See also* CD4+ T cells
- TREM2 gene mutations, 2601
- Trematodes/trematode infections, 255e-2, 1423
approach to the patient, 1424
biliary (hepatic) flukes. *See* Biliary (hepatic) flukes
blood flukes. *See* Schistosomiasis
epidemiology of, 1423t
intestinal flukes. *See* Intestinal flukes
lung flukes. *See* Lung flukes (*Paragonimus* spp.)
prevention and control of, 1429
treatment of, 1428t
- Tremor, 2618
clinical features of, 2618, 2619t
essential, 2619
etiology of, 2619
in PD, 2618
treatment of, 2619–2620
in Wilson's disease, 2625
- Trench fever, 1080–1082, 1081t
- Trench foot (immersion foot), 478e-4
- Trench mouth (Vincent's angina), 231, 233, 236, 238t
- Trypanosoma* spp., 207e-1, 207e-2t, 1132–1133.
See also Trypanematoses
- Trypanosoma pallidum*, 145e-3, 1132–1133, 1137.
See also Syphilis
- Trypanosoma pallidum* particle agglutination (TPPA) test, 1137
- Trypanematoses, 207e-1
clinical features of, 207e-2f, 207e-1–3
comparison of, 207e-2t
control of, 207e-3
diagnosis of, 207e-3
epidemiology of, 207e-1, 207e-1f
treatment of, 207e-3
- Treprostinil, 1659, 1660t, 2164
- TREs (thyroid response elements), 2287, 2287f
- Tretinoin. *See* All-trans-retinoic acid (ATRA)
- TREX1 gene mutations, 2125
- TRH (thyrotropin-releasing hormone), 401e-2, 401e-4
- Triamcinolone, 462e-19, 2234
- Triamterene
action of, 332e-9
drug interactions of, 43
for hypertension, 1623, 1624t
for hypokalemic periodic paralysis, 462e-17
- Triazolam
for anxiety disorders, 2711t
drug interactions of, 246e-3t
for insomnia, 191
overdosage/poisoning with, 473e-10t
pharmacology of, 2711t
- Triazoles, 350
- Tribeč virus, 1308t
- Triceps skinfold thickness, 462
- Trichiasis, 1172
- Trichinella*/trichinellosis, 1410
clinical features of, 908t, 1410, 1561
diagnosis of, 245e-3t, 245e-6t, 1410–1411, 1411f
eosinophilia in, 245e-6t, 765t
epidemiology of, 765t, 1410
life cycle of parasite, 1410, 1410f
pathogenesis of, 1410
prevention of, 1411
treatment of, 246e-13, 1411, 1411t
- Trichloroethylene, 2023
- Tricholemmoma, 364t
- Trichomonas vaginalis*, 1409
- Trichomonas vaginalis* infections, 1409
clinical features of, 1409
diagnosis of, 150e-6, 873t, 1409
screening for, 882
treatment of, 246e-13, 871, 1409
vaginal, 873t, 874
- Trichophyton*, 349, 350, 1357. *See also* Dermatophytosis
- Trichosporon*, 1356–1357
- Trichostrongylus*/trichostrongyliasis, 1416
- Trichotillomania, 355t
- Trichromat, 197
- Trichrome stain, 245e-1
- Trichuris trichiura*/trichuriasis (whipworm)
clinical features of, 1414t, 1416
diagnosis of, 245e-3t, 245e-5t, 765t, 1416
epidemiology of, 765t
life cycle of parasite, 1414t, 1416
treatment of, 1414t, 1416
- Triclabendazole, 246e-3t, 246e-13
- Tricuspid atresia, 1527
- Tricuspid regurgitation, 1548
clinical features of, 1548
diagnosis of, 1548
etiology of, 1547t, 1548
in mitral valve disease, 1551
murmur in, 51e-1f, 51e-2, 51e-4–5, 51e-5f, 1448
in pulmonary hypertension, 1655
treatment of, 1548–1549, 1549f
- Tricuspid stenosis, 1547
in carcinoid syndrome, 564
clinical features of, 1443, 1547
vs. constrictive pericarditis, 1576
diagnosis of, 1547–1548
etiology of, 1547, 1547t
heart sounds/murmur in, 51e-1f, 51e-6, 1449, 1547
pathophysiology of, 1547
treatment of, 1548
- Tricuspid valve, Ebstein's anomaly of, 1527
- Tricuspid valve repair, 1552
- Tricyclic antidepressants (TCAs)
action of, 465e-2
adverse effects of, 93–94, 2682t, 2710t, 2716
autonomic overactivity, 2643
erectile dysfunction, 327
hypertension, 1617t
hypothermia, 478e-1
obesity, 2392
orthostatic hypotension, 146
contraindications to, 2716
for depression, 2710t, 2716, 2716f
distribution of, 34
drug interactions of, 34t, 42t, 246e-2t, 2711t
genetic variations in response to, 39t, 40
for headache, 109, 111
for irritable bowel syndrome, 1969, 1970t
metabolism of, 34t
for migraine prevention, 2594t
for narcolepsy, 190
for nausea and vomiting, 260t, 261
for neuropathy, 2682t
overdosage/poisoning with, 269e-17f, 473e-9t, 473e-10t, 473e-13t, 1458f
for pain, 91t, 93–94, 93t
pharmacology of, 2716
- Trientine, 2519, 2520, 2520t, 2625
- TRIF, 145e-5
- Trifascicular block, 1455, 1476, 1476t. *See also* Atrioventricular conduction block
- Trifluoperazine, 2722t
- Trifluridine, 215e-3t, 215e-9
- Trigeminal autonomic cephalalgias, 107, 2587t, 2594, 2595t
- Trigeminal nerve
anatomy of, 2645–2646, 2645f
disorders of, 215, 2646–2647, 2647t
examination of, 2538
- Trigeminal neuralgia, 2646
clinical features of, 2646
differential diagnosis of, 2646
incidence of, 2646
in MS, 2664
oral pain in, 237
with paroxysmal hemicrania, 2596
pathophysiology of, 2646
treatment of, 94, 2646
- Trigeminal neuropathy, 2646–2647
- Trigeminovascular system, 107
- Trigger(s), migraine, 2586
- Trigger point injections, 120
- Triglycerides
elevated levels of, 2439–2440, 2440t, 2453–2454
impaired lipolysis of. *See* Lipoprotein disorders
long-chain, 1935t, 1936
medium-chain, 1936
in NALFD, 2055
in specialized nutritional support formulas, 98e-5
- Trihexyphenidyl, 2615, 2621, 2722

- Triiodothyronine (T₃)
 characteristics of, 2286f
 free, 2286, 2288
 half-life of, 400e-2
 laboratory evaluation of, 2286t, 2288, 2763t
 receptor binding of, 2287, 2287f
 in sick euthyroid syndrome, 2299
 structure of, 2283f
 in subacute thyroiditis, 2298–2299, 2298f
 total, 2288
 transport and metabolism of, 2285–2286
- Trilostane, 2273
- TRIM5- α protein, 1217
- Trimethoprim
 action of, 64e-2, 332e-8, 641, 931t, 932
 adverse effects of, 64e-2, 246e-4t, 297, 310, 321, 647
 drug interactions of, 246e-4t
 indications for, 246e-4t
 for PCP, 1361t
 resistance to, 931t, 934
- Trimethoprim-polymyxin, 199
- Trimethoprim-sulfamethoxazole (TMP-SMZ)
 for *Acinetobacter* infections, 1037t
 action of, 943
 adverse effects of, 867t, 938t, 943, 1361, 1361t
 cholestatic hepatitis, 284
 in G6PD deficiency, 656t
 hepatotoxicity, 2029–2030
 in HIV infection, 1259, 1260
 neutropenia, 417
 thrombocytopenia, 727t
 vasculitis, 2218t
 for *B. cepacia* infections, 1045t
 for bite-wound infections, 167e-2, 167e-3t
 for blastocystosis, 1409
 for *Brucella* spp. infections, 194e-4
 for *C. belli* infections, 1407
 for chronic granulomatous disease, 423
 for *Cyclospora* infections, 1408
 for cystitis, 866, 867t
 for donovanosis, 198e-2t
 drug interactions of, 939t
 for enteric fever, 1052t
 for glanders, 1045t
 for granulomatosis with polyangiitis, 2185
 indications for, 941t, 943
 for *L. monocytogenes* infections, 983
 for *Legionella* infections, 1020t
 for melioidosis, 1045t
 for MRSA infections, 345, 962t
 for *Nocardia* infections, 1087
 for nontyphoidal *Salmonella* infections, 1054t
 for osteomyelitis, 840t, 841t
 for otitis media, 230
 for paracoccidioidomycosis, 1354t
 for PCP, 1361, 1361t
 for PCP in HIV infection, 1251t, 1253–1254, 1830
 for pertussis, 1024, 1024t
 for plague treatment, 261e-5
 prophylactic
 in chronic neutropenia, 423
 for cystitis, 889
 in HIV infection, 1251t
 for PCP, 1251t, 1362t
 for peritonitis, 847
 for plague, 1075t
 for toxoplasmosis, 1251t, 1404
 in transplant recipient, 139e-4t, 921, 925–926, 925t
 for pyelonephritis, 867
 resistance to, 943
E. coli, 1029
 in *E. coli*, 862
 mechanisms of, 934
 rates of, 941t
 for *S. maltophilia* infections, 1045t
 for staphylococcal soft tissue infections, 962t
 for vasculitis, 2182
 for Whipple's disease, 1093–1094, 1945
- Trimethylaminuria (fish odor syndrome), 96e-6
- Trimetrexate, 246e-3t
- Trinucleotide repeat disorders, 434, 440, 440t
- Triose phosphate isomerase deficiency, 654t
- Triple A syndrome, 2324t
- Triple-negative breast cancer, 526
- Triptans, 473e-13t, 2591, 2592t
- Trismus (lockjaw), 985, 2647
- Trisomy, 83e-4
- Trisomy 6, 83e-4
- Trisomy 8, 135e-6
- Trisomy 13, 83e-4, 1654
- Trisomy 18, 83e-4, 1654
- Trisomy 21. *See* Down's syndrome (trisomy 21)
- Tritium exposure, 263e-3t, 263e-6t, 263e-7
- TrkB* gene, 415e-4t
- TRK* gene mutations, 101e-3t, 2304, 2305t
- Trochanteric bursitis, 2222, 2222f, 2247
- Trochlear nerve, 2537
 injuries of, 457e-2
 palsy, 209
- Troglitazone, 96e-6, 2025, 2415
- Tropheryma whippelii*. *See also* Whipple's disease
 asymptomatic colonization of, 1092
 characteristics of, 840, 1091, 1945
 isolated infection, 1093
- Trophoblast stem cells, 453, 453t
- Trophozoites, *Plasmodium*, 1369
- Tropical pulmonary eosinophilia, 422, 1420
- Tropical spastic paraparesis. *See* Human T-cell lymphotropic virus-1 (HTLV-1) infections, myelopathy
- Tropical splenomegaly, 1374–1375
- Tropical sprue, 1942
 clinical features of, 269, 1942
 clinical pattern of, 1943
 cobalamin deficiency in, 645
 diagnosis of, 1943
 etiology of, 1943
 small-intestinal mucosal biopsies in, 1939, 1940t, 1941f, 1942
 treatment of, 1943
- Tropisetron, 566
- Tropomyosin, 265e-6, 1556f
- Troponins
 in cardiac contraction, 265e-4–5, 265e-5f
 in cardiac trauma, 289e-3
 in chest pain evaluation, 101
 in chronic kidney disease, 1817
 in heart failure, 1505
 in myocardial infarction, 1594, 1596t, 1600–1601, 1601f
 reference values for, 2763t
- Trousseau's sign, 315
- Trousseau's syndrome, 511, 613
- TRPM6* gene mutations, 332e-8
- TRP proteins, 212
- TRPs (transient receptor potentials), 444e-9
- Trubanam virus, 1306t, 1313
- True bug bites, 2750
- True hermaphroditism. *See* Ovotesticular disorders of sexual development
- True-negative tests, 21, 22
- True neurogenic thoracic outlet syndrome (TOS), 122
- True-positive tests, 21, 22
- Trust, between patient and health care provider, 16e-4–5, 16e-6–7
- Trypanosoma* spp.
 entry into host, 145e-1
T. brucei gambiense/rhodesiense, 245e-4t, 245e-6t, 908t, 1396, 1396t. *See also* Sleeping sickness (human African trypanosomiasis)
T. cruzi, 245e-2, 245e-4t, 245e-6t, 1394. *See also* Chagas disease
- Trypanothione reductase, 246e-7
- L-Tryptophan
 contamination of, 14e-3, 376, 423, 462e-20, 2155, 2199
 drug interactions of, 2711t
 overdose/poisoning with, 473e-13t
 Tryptophan hydroxylase, 372e-24t
 Tryptophan metabolism disorders, 434e-2t
 Tryptophanuria, 434e-2t
TSC1/2 gene mutations, 101e-6t, 465e-1, 465e-2t, 1854
- T-scores, 2488, 2493, 2493f
- Tsetse flies, in sleeping sickness, 1396
- TSH. *See* Thyroid-stimulating hormone (TSH)
- TSI (thyroid-stimulating immunoglobulins), 2288, 2294
- TSLP (thymic stromal lymphopoietin), in asthma, 1672
- T-SPOT TB assay, 1115
- TSS. *See* Toxic shock syndrome (TSS)
- TTF-1. *See* Thyroid transcription factor-1 (TTF-1)
- TTF-2, 2283, 2284t
- TTKG (transtubular K⁺ concentration gradient), 64e-7, 307, 311, 311f
- TTN* gene mutations, 1564
- TTP. *See* Thrombotic thrombocytopenic purpura (TTP)
- TTR. *See* Transthyretin (TTR)
- Tubal pregnancy. *See* Ectopic pregnancy
- Tuberculin skin test (TST). *See* Purified protein derivative (PPD)/tuberculin skin test (TST)
- Tuberculoma, 1111
- Tuberculosis (TB), 1102
 anemia in, 630
 AV conduction block in, 1471t
 bronchiectasis in, 1695
 in cancer patient, 484t, 485, 486t
 as cause of fever of unknown origin, 138, 141
 in children, 1108, 1108f
 as chronic disease, 13–14
 diagnosis of
 culture, 1113
 drug susceptibility testing, 1113–1114
 genomics in, 769t
 imaging, 308e-3f, 1108f, 1109f, 1114
 invasive procedures, 1114
 limitations of, 770
 microscopy, 1113
 nucleic acid amplification, 150e-8, 1113
 serologic tests, 1114
 sputum analysis, 1114
 tuberculin skin testing, 1107, 1114
 epidemiology of, 12–13, 205e-1, 1103–1105, 1104f
 etiology of. *See* *Mycobacterium tuberculosis* extensively drug-resistant, 13, 1116t, 1119, 1119t
 extrapulmonary
 adrenal, 1112
 aortic aneurysm, 1638
 congenital, 1112
 gastrointestinal, 287, 1111
 genital, 335
 genitourinary, 1110, 1110f
 in HIV infection, 1112–1113
 lymphadenitis, 408, 1109, 1109f
 lymphedema, 1654
 mastitis, 1112
 meningitis, 488, 766t, 775, 907t, 1111
 military, 138, 1108f, 1112, 1119
 myocarditis, 992
 ocular manifestations, 1112
 oral manifestations, 237, 238t, 241
 otitis, 1112
 pericardial, 1574, 1577
 pericarditis, 1111–1112
 pleural, 1109–1110
 skin manifestations, 366, 367, 829, 1112
 spinal/skeletal
 vs. *Brucella* infection, 194e-2, 194e-2t
 clinical features of, 836, 1110–1111
 diagnosis of, 138
 MRI, 441e-2f
 osteomyelitis, 836
 Pott's disease, 836, 1110, 1111f
 treatment of, 1119–1120
 tuberculoma, 1111
 upper airway, 234, 1110
 global burden of, 12–13, 205e-1, 1103–1105, 1104f
 in HIV infection
 antiretroviral therapy and, 205e-1, 1112–1113
 clinical features of, 1112–1113, 1254

- Tuberculosis (TB), in HIV infection (*Cont.*):
 in developing countries, 12, 12f, 1112–1113, 1122
 epidemiology of, 1103, 1254
 immune reconstitution inflammatory syndrome in, 205e-1, 1112–3532110, 1254
 interaction of, 1233
 latent infection, 1120t, 1121
 prophylaxis for, 1251t
 risk of, 1105, 1105t
 treatment of, 205e-5, 1119
 incubation period for, 152e-3t
 infection control measures, 917, 1284
 innate resistance to, 1106
 latent infection
 diagnosis of, 775
 screening for, 2147
 treatment of, 205e-1t, 1120–1121, 1120t
 multidrug-resistant
 detection of, 769t, 773, 774
 epidemiology of, 1103–1105, 1118, 1118f
 treatment of, 13, 205e-1–2, 1116t, 1118–1119
 natural history of, 1105
 pathogenesis of, 1105–1107
 prevention and control of, 13, 1121–1122
 pulmonary
 cough in, 1108
 hemoptysis, 245
 hemoptysis in, 1108
 pleuritis in, 1718
 postprimary disease, 1105, 1108–1109, 1109f
 primary disease, 1105, 1108, 1108f
 in returning war veterans, 152e-3t, 152e-8
 risk factors for, 1105, 1105t
 silica exposure and, 1690
 transmission of, 1105
 in transplant recipient, 920, 925t, 928
 treatment of, **205e-1**
 adherence to, 1116–1117
 adverse effects of, 205e-1, 1117
 DOTS/DOts plus strategy, 13, 205e-1
 for drug-resistant strains, 205e-1–2, 1118–1119, 1119t
 drug susceptibility testing, 1113–1114
 in extrapulmonary disease, 1119–1120
 failure and relapse, 1117–1118
 first-line antituberculosis drugs, 205e-2–5, 205e-2t, 1115–1116, 1115t
 ethambutol. *See* Ethambutol
 isoniazid. *See* Isoniazid
 pyrazinamide. *See* Pyrazinamide
 rifabutin, 205e-5
 rifampin. *See* Rifampin
 rifapentine, 205e-5
 streptomycin, 205e-5, 1115
 latent infection, 205e-1t, 1120–1121, 1120t
 monitoring response to, 205e-2t, 1117
 newer drugs, 205e-7
 in pregnancy, 1116t, 1120
 principles of, 205e-1–2, 205e-2t
 regimens, 205e-2t, 1113–1117, 1116t
 second-line antituberculosis drugs, 205e-6–7
 simplified approach to, 205e-2t
 Tuberin, 562
 Tuberos sclerosis (Bourneville's disease)
 brain tumors in, 600
 clinical features of, 599t, 604, 1850t, 1854
 genetic considerations in, 101e-6t, 562, 562t, 599t, 604, 1850t, 1854
 imaging in, 441e-36f
 pathogenesis of, 604, 1854
 pNETs in, 562, 562t
 renal cell carcinoma and, 578, 1854
 skin manifestations of, 344, 359, 364
 treatment of, 604
 Tuberos xanthoma, 436e-4f
tub gene, 415e-4, 415e-4t
 Tublin, 372e-24t
 Tuboovarian abscess, 1089, 1099
 Tubuloglomerular feedback, 332e-2, 332e-10f, 333e-3, 333e-5, 1800
 Tubulointerstitial diseases, **1856**
 acute interstitial nephritis. *See* Acute interstitial nephritis (AIN)
 chronic, 1856t, **1859**
 analgesic nephropathy, 1861, 1861f
 aristolochic acid nephropathy, 577, 1804, 1861
 calcineurin-inhibitor nephropathy, 1829, 1862
 clinical features of, 1856
 in glomerulonephritis, 1860–1861
 heavy metal nephropathy, 1862
 karyomegalic tubulointerstitial nephritis, 1855, 1861
 lithium-associated nephropathy, 1861–1862
 sickle cell nephropathy, 1835t, 1848, 1860
 vesicoureteral reflux and reflux nephropathy, 1859–1860, 1860f
 classification of, 1867t
 cystic and hereditary
 genetic considerations in, 332e-7t
 global considerations in, 1855–1856
 medullary cystic kidney disease, 1850t, 1854
 medullary sponge kidney, 1850t, 1855, 1866
 nephronophthisis, 1850t, 1855
 polycystic kidney disease. *See* Polycystic kidney disease
 tuberous sclerosis. *See* Tuberos sclerosis
 von Hippel-Lindau disease. *See* von Hippel-Lindau disease
 global considerations in, 1862–1863
 metabolic, 1856t
 hyperkalemic nephropathy, 307, 1862
 hypokalemic nephropathy, 1862
 uric acid nephropathy. *See* Uric acid nephropathy
 Tubulointerstitial nephritis with uveitis (TINU), 1857–1858, 1857t, 1858f
 Tularemia, **1066**
 animal bite-related, 167e-1
 in bioterrorism, 261e-7
 clinical features of, 261e-5t, 261e-7, **1067**, 1068t
 oropharyngeal/gastrointestinal, 1068
 pulmonary, 1068
 skin manifestations, 25e-11f, 134t
 typhoidal, 1067, 1069
 ulceroglandular/glandular, 134t, 829, 1068, 1068f
 diagnosis of, 261e-5t, 1069
 differential diagnosis of, 1069, 1069t
 epidemiology of, 1067
 etiology of, 1067
 global considerations in, 1067
 pathogenesis of, 1067
 prevention of, 261e-5t, 261e-8, 1070
 prognosis of, 1070
 septic shock in, 781
 treatment of, 261e-5t, 261e-7–8, 1069–1070
 Tula virus, 1307t
 Tullio phenomenon, 219
 Tumbu fly, 2747
 Tumeric, 404t
 Tumor, 103e-1. *See also specific types and sites*
 Tumor (sign of inflammation), 414f, 1753
 Tumor (skin lesion), 339t
 Tumoral calcinosis, 426e-7t, 426e-8, 1815, 1815f, 2460
 Tumor burden, 470
 Tumor-induced osteomalacia, 2459, 2460
 Tumor lysis syndrome
 acute kidney injury in, 1804, 1806t
 clinical features of, 1795, 1804
 hyperkalemia in, 310
 hypocalcemia in, 315
 treatment of, 431e-5, 1795–1796, 1796f
 Tumor markers, 473, 473t
 in carcinoma of unknown primary, 120e-1, 120e-2t
 in ovarian cancer, 592
 in pancreatic cancer, 554, 555
 in testicular cancer, 588
 Tumor necrosis factor (TNF)
 in anemia of inflammation, 630, 630f
 in fever induction, 124, 125f
 in sarcoidosis, 2205, 2205f
 Tumor necrosis factor receptor(s), 102e-8
 Tumor necrosis factor receptor-1–associated syndrome, 372e-8t, 1845
 Tumor necrosis factor (type III) receptor family, 372e-16
 Tumor necrosis factor- α (TNF- α)
 in ankylosing spondylitis, 2170
 in *Brucella* infections, 194e-2
 in depression, 465e-4
 functions of, 422
 in HIV infection, 1235
 in nontuberculous mycobacterial infections, 1129
 in rheumatic fever, 2150
 in rheumatoid arthritis, 2141, 2142f, 2143
 in sepsis/septic shock, 1753
 in shock, 1746–1747
 source, target, and biologic activity of, 372e-11t
 in tumor angiogenesis, 102e-15
 Tumor necrosis factor- α (TNF- α) inhibitors. *See also specific drugs*
 adverse effects of, 1962, 2146t, 2173
 antibody development, 1961
 cutaneous, 378–379, 1962
 hepatosplenic T-cell lymphoma, 1962
 infections, 1962, 2145, 2147, 2173
 latent tuberculosis reactivation, 2147
 non-Hodgkin's lymphoma, 1961–1962
 periodic syndrome (TRAPS), 141, 422, 2213t, 2214–2215
 serious, 348, 348t, 2146t
 for alcoholic liver disease, 2053, 2054f, 2059
 for ankylosing spondylitis, 2172–2173
 for autoimmune and inflammatory diseases, 372e-30
 contraindications to, 348, 2173
 for IBD, 1961, 2179
 monitoring during treatment with, 2146t
 for psoriasis, 348, 348t
 for psoriatic arthritis, 2177
 for rheumatoid arthritis, 2145, 2146t, 2147, 2148
 Tumor necrosis factor- α receptor-associated periodic syndrome (TRAPS), 141, 422, 2213t, 2214–2215
 Tumor necrosis factor- β (TNF- β), 372e-11t
 Tumor-suppressor genes
 caretaker genes, 101e-2, 101e-5
 definition of, 101e-2, 101e-4
 epigenetic silencing of, in cancer, 102e-7–8, 102e-8f
 inactivation of, 101e-2, 101e-4
 in lung cancer, 508
 in thyroid cancer, 2304–2305, 2305t
 Tumor vaccines, 103e-20
 Tumstatin, 102e-17f
 Tunari virus, 1319
Tunga penetrans, 2750
 Tungiasis, 2750
 Tungsten carbide exposure, 1691
 Tunica media, 265e-1, 265e-1f
 Tunneled catheters, 1823
 Tunnel infections, 1825
 Turcot's syndrome, 538t, 539, 599t, 602
 Turkey handling disease, 1681t
 Turner's sign, 2092
 Turner's syndrome, **2351**
 aortic valve disease in, 1529
 autoimmune polyendocrine deficiencies in, 2347t, 2348
 clinical features of, 2351, 2351t
 coarctation of the aorta in, 1620
 genetic considerations in, 83e-4, 439
 IBD and, 1949t
 lymphedema in, 1654
 pathophysiology of, 2351
 prevalence of, 2254t
 primary ovarian insufficiency in, 336, 337
 screening/testing for, 2254t
 treatment of, 2351–2352

- Tuskegee study, 16e-5, 1134
 T wave, 273e-1, 1453
 Twiddler's syndrome, 1469
Twinkle, 462e-16
 Two-point discrimination test, 159, 2539
 Tylosis palmaris et plantaris, 533
 Tympanic membrane, 217, 218f
 assessment of, 220
 perforation of, 217, 223
 Tympanic membrane thermometer, 124
 Tympanocentesis, 229t
 Tympanogram, 222
 Tympanometry, 222
 Tympanoplasty, 217, 223, 230
 Tympanostomy tubes, 223, 230
 "Type and screen," 138e-2
 Typhoidal tularemia, 1067, 1068t, 1069
 Typhoid fever. *See* Enteric (typhoid) fever
 Typhoid vaccine
 for transplant recipient, 929
 for travelers, 795, 795t, 1052-1053
 Vi CPS, 1052
 Vi-rEPA, 1052
 whole-cell, 1052
 Typhus
 endemic. *See* Endemic (murine) typhus
 epidemic. *See* Epidemic (louse-borne) typhus
 scrub. *See* Scrub typhus
 Tyramine, 2615
 Tyrosinase, 372e-24t, 386
 Tyrosinase gene mutations, 357
 Tyrosine hydroxylase, 372e-24t
 Tyrosine kinase inhibitors
 action and targets of, 102e-3, 102e-4, 103e-18t, 103e-19
 adverse effects of, 505, 692, 694, 1563
 for AML, 685t
 for CML, 691-692, 691t
 cost of, 694-695
 global considerations in, 694-695, 694f
 for head and neck cancer, 505
 in pregnancy, 694
 resistance to, 689
 Tyrosine kinase receptors, 400e-3, 400e-3f
 Tyrosine metabolism disorders, 434e-2t
 Tyrosinemia, 434e-2t, 545, 545t
 Tzanck smear, 341-342
- U**
 Ubiquitin, 2632
 UDCA. *See* Ursodeoxycholic acid (UDCA)
 Udd's distal myopathy (tibial muscular dystrophy), 462e-12, 462e-13t
 UDPGT (uridine diphosphate-glucuronosyl transferase), 280, 282, 1999, 1999f
 UDPGT gene mutations, 282
 UGT1A1, 34t, 39t, 40, 2001
 UGT1A1 gene mutations, 655, 2001-2002, 2077
 UGT1 gene complex, 2000, 2000f
 Uhthoff's symptom, 2663
 Ulcer(s). *See also* specific types
 cutaneous. *See* Cutaneous ulcers
 definition of, 1911, 1914
 foot, 2428-2429. *See also* Diabetes mellitus, complications of
 genital. *See* Genital ulcer(s)
 oral, 46e-4f, 237, 239t, 2129t
 pressure (decubitus), 830, 1743, 2660
 Ulcerative colitis. *See also* Inflammatory bowel disease (IBD)
 clinical features of, 1952, 1952t
 colorectal cancer in, 1964, 1964f
 complications of, 1953
 diagnosis of, 1952-1953, 1952t, 1953f
 differential diagnosis of, 1061, 1956-1957, 1956t
 epidemiology of, 1947, 1947t
 etiology of, 1947
 genetic considerations in, 1949, 1950t
 immune regulation in, 1949-1950
 pathogenesis of, 372e-23, 1948
 pathology of, 1951, 1951f
 in pregnancy, 49, 1964
 racial and ethnic differences in, 1947-1948
 serologic markers in, 1954-1955
 smoking effects on, 1947
 spondyloarthritis and, 2178-2179
 treatment of, 1962-1963, 1963f, 1963t
 tumor markers in, 473t
 Ulipristal acetate, 2391
 Ulnar nerve, 442e-5f
 Ulnar neuropathy, 463e-4, 463e-5f, 1124, 2683, 2691
 Ultrafiltration, 1508-1509, 1823
 Ultra-rapid metabolizers, 38f, 40
 Ultrasonography
 in appendicitis, 1987-1988
 in bile duct evaluation, 2085t
 in bone mass measurement, 2493
 in chronic venous disease, 1651
 of deep leg veins, 1633, 1633f
 endobronchial, 512, 1664, 1668
 endoscopic. *See* Endoscopic ultrasound (EUS)
 in fever of unknown origin, 138
 in gallbladder disease, 2078, 2079f, 2079t
 intravascular coronary artery, 1465, 1465f
 in liver disease, 1993
 in liver disease evaluation, 1999
 in musculoskeletal disease evaluation, 2223-2224, 2224t
 in pancreatic disease evaluation, 2087t, 2088
 in peripheral arterial disease, 1643
 pleural, 1663
 prenatal, 83e-4
 of renal arteries, 1628, 1628t
 in rheumatoid arthritis, 2144
 in stroke, 2581
 in thyroid dysfunction evaluation, 2289, 2289f
 of thyroid nodule, 2289, 2289f, 2289t
 Ultratrace elements, **96e-10**
 Ultraviolet light, **385**. *See also* Sun exposure
 skin effects of, 386
 UV-A, 385-386
 UV-B, 385-386
 Ultraviolet light therapy. *See* Phototherapy
 Umami taste sensation, 212
 Umbilical cord stem cells, 89e-3-4, 90e-2, 139e-2, 453t
 Umbilical venous hum, 286
 UMOD gene mutations, 1854
 UMP (uridine-5'-monophosphate synthetase) deficiency, 431e-5t, 431e-6
 UNAIDS (Joint United Nations Program on HIV/AIDS), 8, 12
 Unani medicine, 14e-1t
 Una virus, 1306t
 Uncal transtentorial herniation, 1772, 1772f
 Uncoupling protein, 415e-2
 Underweight, 2393t
 Undulant fever. *See* *Brucella* spp. infections
Unequal Treatment (IOM report), 16e-2, 16e-5-6
 Unfractionated heparin. *See* Heparin
 UniFrac analysis, 86e-1t, 86e-3f, 86e-5
 Uniparental disomy, 83e-6-7, 83e-6f, 439
 Unipolar depressive disorders, 2715. *See also* Depression/depressive disorders
 Uniporters, 332e-6
 Unipotent cells, 454, 454f
 United Nations Children's Fund (UNICEF), 8
 United Network of Organ Sharing (UNOS)
 in cardiac transplantation cadaver donor selection, 1516
 liver transplantation waiting list criteria, 2069-2070, 2069t
 Universal coverage reforms, 13e-4-5, 13e-5f
 University of Pennsylvania Smell Identification Test, 214f, 216
 Unrestricted somatic stem cells, 453t
 Unstable angina, 96, 96t, 97, 1593. *See also* Non-ST-segment elevation acute coronary syndrome (NSTEMI-ACS)
 Unverricht-Lundborg disease (Baltic myoclonus), 451e-3t
 Upbeat nystagmus, 211
 Upper esophageal sphincter, 255
 Upper motor neuron diseases, 2634-2635t, 2636
 Upper motor neuron pathways, 155f
 Upper motor neuron weakness, 154, 154t
 Upper respiratory tract infections. *See* Respiratory tract infections
 Upshaw-Schulman syndrome, 729, 1864
 Uranium-235 exposure, 263e-3t, 263e-6t
 Urate(s), 431e-1, 435e-1t
 Urate nephropathy (urate nephrosis), 431e-4, 1858
 Urate oxidase. *See* Rasburicase
 Urate transporter 1 (URAT1), 431e-1, 431e-2f
 Urea breath test, 1040, 1040t
 Urea cycle defects, 434e-3t, 434e-5
Ureaplasma infections, 871, 1164t, 1165
 Ureidopropionase deficiency, 431e-5t
 Uremia
 abdominal pain in, 105
 in acute kidney injury, 1808
 peripheral blood smear in, 81e-4f, 395f
 Uremic acidosis, 320
 Uremic fetor, 1819
 Uremic syndrome, 1811, 1813. *See also* Chronic kidney disease (CKD)
 Ureter
 cancer of, 577
 infections in cancer patient, 490
 obstruction of, 104
 Ureteric bud, 332e-1, 332e-1f
 Urethra, 92e-3t
 Urethritis
 in men
 C. glaucuronolyticum, 981
 chlamydial, 1167, 1170t
 clinical features of, 870, 1005
 diagnosis of, 871
 etiology of, 870-871, 870t, 1005
 gonococcal, 1005, 1005f
 nongonococcal, 1167-1168, 1170t
 postgonococcal, 1167, 1170t
 treatment of, 872t
 Mycoplasma, 1165
 Ureaplasma, 1165
 in women, 294, 872, 1005, 1169, 1170t
 Urge incontinence, 80
 Uric acid
 decreased excretion of. *See* Hypouricemia
 increased production of. *See* Hyperuricemia
 laboratory evaluation of, 2222-2223, 2764t
 metabolism of, 431e-1, 431e-1f, 431e-2f
 stones, renal, 431e-4-5, 1870-1871. *See also* Nephrolithiasis
 tissue deposition of. *See* Tophi
 urine, 62e-12f, 1867
 Uric acid nephropathy
 in AML, 681-682
 hyperuricemia and, 431e-4, 1862
 pathophysiology of, 1862
 in polycystic kidney disease, 1852
 treatment of, 431e-5, 1862
 Uric acid stones, renal, 1866, 1870-1871. *See also* Nephrolithiasis
 Uricosuric compounds, 431e-1, 431e-2t
 Uridine-5'-monophosphate synthetase (UMP) deficiency, 431e-5t, 431e-6
 Uridine diphosphate galactose 4-epimerase deficiency, 433e-3t, 433e-5
 Uridine diphosphate-glucuronosyl transferase (UDPGT), 280, 282, 1999, 1999f
 Urinalysis
 in abdominal pain evaluation, 106
 in acute kidney injury, 1805, 1807f
 atlas of, 62e-11-12f
 in hematuria, 294. *See also* Hematuria
 in nephrolithiasis, 1868f, 1869
 parasites in, 245e-6t
 in proteinuria, 293-294. *See also* Proteinuria
 reference ranges for, 2768-2769
 specimen collection and transport, 150e-4t
 Urinary alkalization, for overdosage/poisoning, 473e-7
 Urinary analgesics, 867
 Urinary diversion, 577

- Urinary incontinence
 after prostatectomy, 583
 in elderly, 80–81, 81f
 postmenopausal hormone therapy and, 2384t
 in terminally ill patient, 69t
 treatment of, 81
- Urinary tract
 bleeding from. *See* Hematuria
 congenital abnormalities of, 1855, 1871, 1871t
- Urinary tract infections (UTIs), **861**
 approach to the patient, 863–864, 865f
 in burn patient, 166e-2
 in cancer patient, 490
 clinical features of, 863–864, 865f
 complicated, 861, 864, 867–868
 definitions in, 861
 delirium in, 169
 in diabetes mellitus, 2427
 diagnosis of, 864–866, 865f
 differential diagnosis of, 864
 epidemiology of, 861–862
 etiology of, 862
 A. baumannii, 1037
 Citrobacter, 1035
 E. coli, 1028
 enterococcal, 973, 975t
 K. pneumoniae, 1032, 1871
 Morganella, 1035
 nontyphoidal *Salmonella*, 1054
 P. aeruginosa, 1045t, 1047
 Proteus, 1033, 1871
 Providencia, 1035, 1871
 S. aureus, 959
 S. saprophyticus, 960
 tuberculous, 1110
 genetic considerations in, 863
 global considerations in, 862, 866
 health care–associated, 868, 913–914, 914t
 in MS, 2672
 nephrolithiasis and, 1871
 pathogenesis of, 862–863, 862f
 in pregnancy, 49–50, 867
 prevention of, 868
 prognosis of, 868
 pyelonephritis. *See* Pyelonephritis
 recurrent, 861
 renal abscess and, 851
 risk factors for, 861–862
 in transplant recipient, 924
 treatment of
 acute cystitis in women, 866–867, 867t
 asymptomatic bacteriuria, 868
 Candida infection, 868
 catheter-associated, 868, 914
 complicated infection, 867–868
 drug resistance and, 862, 866
 E. coli infection, 1029
 in men, 867
 in pregnancy, 867
 urinary tract obstruction in, 1873
- Urinary tract obstruction, **1871**
 acute kidney injury and, 291–292, 1804–1805, 1804f
 in cancer, 1790
 clinical features of, 289t, 1872–1873
 diagnosis of, 289t, 1873, 1873f
 diuresis following, 1874
 etiology of, 1871–1872, 1871t
 hyperkalemia in, 307
 hyponatremia in, 299
 hypovolemia in, 297
 pathophysiology of, 1872–1873, 1872t
 prognosis of, 1874
 treatment of, 1873–1874
- Urinary urea nitrogen, 462–463, 464t
- Urinary D-xylose test, 1938
- Urination, painful
 in cystitis. *See* Cystitis
 in interstitial cystitis bladder pain syndrome.
 See Interstitial cystitis/bladder pain syndrome
 in men, 60e-1
 in urinary tract infections. *See* Urinary tract infections (UTIs)
 in women, 60e-1
- Urine
 abnormalities of, 289t, **293**. *See also* Urinalysis
 casts, 62e-11–12f, 294
 crystals, 62e-12f, 435e-2
 hematuria. *See* Hematuria
 polyuria. *See* Polyuria
 proteinuria. *See* Proteinuria
 pyuria, 294
 black, 434e-4
 nephrolithiasis risk and, 1867
 osmolality of, 291t, 294–295, 301, 304
 sodium loss in, 298, 301
 tea/cola colored, 279, 462e-1
 Urine bilirubin, 280, 1996, 1998t, 2000
 Urine-derived stem cells, 453t
 Urine dipstick test, 280, 864
 Urine output, 95e-1
 Urobilinogens, 280, 2000
 Urobilins, 280
 Urocanic acid, 388
 Urocanic aciduria, 434e-2t
 Urochromes, 1819
 UROD gene mutations, 2530–2531
 Urogenital secretions, specimen collection and transport, 150e-4t
 Urography, 1873
 Urokinase, 758–759
 Urokinase-type plasminogen activator (u-PA), 402, 402f, 758
 Uromodulin. *See* Tamm-Horsfall protein
 Uroplakins, 145e-2, 145e-5
 Uroporphyrinogen (URO) decarboxylase, 2523t, 2525
 Uroporphyrinogen III (URO) synthase, 2523t, 2524–2525
 UROS gene, 2523t
 Ursodeoxycholic acid (UDCA)
 for gallstone dissolution, 2080
 for gallstone prevention during rapid weight loss, 2078
 for primary biliary cirrhosis, 2061
 for primary sclerosing cholangitis, 2062, 2086
- Urticaria, **363, 2117**
 aquagenic, 2118
 cholinergic, 363, 363t, 2119
 classification of, 2117, 2118t
 clinical features of, 135, 341t, 2118–2119
 distribution, 341t
 morphology, 355
 papules and plaques, 25e-9f, 76e-4f, 342, 343f
 cold, **363, 363t, 422, 2118–2119**
 contact, 2118
 definition of, 2117
 diagnosis of, 342, 2119
 drug-induced, 377, 381
 etiology of, 135, 363t, 2117–2118, 2118t
 local heat, 2118
 pathophysiology of, 2118–2119
 physical, **363, 363t**
 predisposing factors for, 2117–2118, 2118t
 pressure, 2118
 solar, **363, 363t, 2118**
 in systemic mastocytosis, 2120. *See also* Mastocytosis
 treatment of, 381, 2119–2120
- Urticarial vasculitis, 132t, 135, **363, 363t**
- Urticaria pigmentosa, 363, 363t, 367
- Urushiol, 345
- U.S. Department of Health and Human Services, Action Plan to Prevent Healthcare-Associated Infections, 912
- U.S. President's Emergency Plan for AIDS Relief (PEPFAR), 8
- U.S. Preventive Services Task Force, 28–30, 29t
- Usher's syndrome, 220, 222t
- Ustekinumab, 348t, 1962, 2173, 2177
- Usutu virus, 1307t, 1315
- Uterus, disorders of
 amenorrhea in, 336, 337f
 endometrial cancer. *See* Endometrial cancer
 endometriosis. *See* Endometriosis
 gas gangrene, 831
 pelvic pain in, 338
 sarcoma, 596, 597
 treatment of, 336
- UTIs. *See* Urinary tract infections (UTIs)
- Uukuniemi virus, 1310t
- Uveitis
 in ankylosing spondylitis, 2171
 anterior, 200, 373e-7, 373e-8t, 2171
 in Behçet's syndrome, 2194
 in IBD, 1958
 paraneoplastic, 614t, 619
 posterior, 200
 in secondary syphilis, 1135
 treatment of, 2173
 with tubulointerstitial nephritis, 1857–1858
 in Whipple's disease, 1093, 1094
- Uvula, bifid, 1443
- Uvulopalatopharyngoplasty, 1727
- U wave, 1453
- V**
- Vaccine(s), **785**. *See also* Immunization; *specific vaccines*
 administration of, 791–792, 791f
 adverse event reporting, 792
 cancer prevention and, 476–477
 contraindications to, 786, 789–790t
 development of, genomics in, 775–776, 778
 documentation of, 792
 for elderly, 84
 future trends in, 793
 global considerations in, 785
 hypersensitivity to, 790
 impact of, 785–786, 785t
 information statements, 790–791
 precautions for, 789–790t
 routinely administered in U.S., 785t
 safety monitoring, 792
 storage and handling of, 791
 tumor, 103e-20
 for viral infections, 214e-9
 viral vectors for, 214e-9
- Vaccine Adverse Event Reporting System (VAERS), 792
- Vaccine Information Sheet (VIS), 790–791
- Vaccine-preventable disease, 785, 785t, 786
- Vaccine Safety Datalink Project, 792
- Vaccinia vaccine, 261e-7, 261e-7t
- Vaccinia virus, 214e-8. *See also* Smallpox
- Vacuum constriction device, 329
- VAERS (Vaccine Adverse Event Reporting System), 792
- Vagabonds' disease, 2746
- Vagina
 blind, 335
 cancer, 1198
 congenital absence of, 2357
 dilatation of, 336
 discharge from, 872–873, 873t, 875–876.
 See also Vulvovaginal infections
 microbiota of, 86e-5, 863
- Vaginal ring, monthly, 2391
- Vaginal septum, 335
- Vagotomy, 1925–1927, 1925f
- Vagus nerve
 anatomy of, 213f
 disorders of, 2649
 examination of, 2538
 stimulation, for depression, 2717
 in taste sensation, 213
- Valacyclovir
 action of, 215e-6
 adverse effects of, 215e-7
 for Bell's palsy, 2648
 for herpes zoster, 215e-3–4t, 215e-7, 832t, 1186, 2657
 for HSV infections, **1181, 1182t**
 esophagitis, 1910
 genital, 215e-3t, 215e-7, 832t
 mucocutaneous, 215e-3t
 for HSV prophylaxis in HIV infection, 1181, 1252t

- pharmacology of, 215e-6, 215e-7
for varicella, 1185–1186
for viral encephalitis, 897
for viral meningitis, 893
- Valdecoxib, 91t
- Valganciclovir
for CMV infection prophylaxis, 215e-8, 922, 1252t
for CMV infection treatment, 215e-2t, 1193, 1831
pharmacology of, 215e-8
resistance to, 215e-8
- Valgus, 2221, 2228, 2228f
- Valine metabolism disorders, 434e-3t
- Valley fever. *See* *Coccidioides* spp. infections
- Valproate/valproic acid
adverse effects of, 2594t
bone marrow suppression, 2556
common, 2718t
hair loss, 380
hepatotoxicity, 2023, 2028, 2556
mitochondrial defects and, 85e-9
neurologic, 2555t
obesity, 2392
rare, 2718t
systemic, 2555t
thrombocytopenia, 727t
zinc deficiency, 96e-9
- for bipolar disorder, 2718, 2718t
drug interactions of, 939t, 2555t
for epilepsy, 2552, 2555t, 2556
for migraine prevention, 2594t
for myoclonus, 2623
for new daily persistent headache, 111
overdosage/poisoning with, 473e-10t, 2766t
pharmacology of, 2555t, 2718t
for posthypoxic myoclonus, 1782
for status epilepticus, 2558f
for stiff-person syndrome, 618
for tardive dyskinesia, 2624
therapeutic monitoring of, 2766t
- Valrubicin, 576
- Valsalva maneuver
in autonomic dysfunction evaluation, 2640, 2640t, 2641t
for focal atrial tachycardia, 1477
heart murmurs and, 51e-7, 1450, 1450t
for PSVT, 1484
- Valsalva ratio, 2640, 2641t
- Valsartan, 1511–1512, 1512t, 1624t
- Valve area, measurement of, 1463
- Valvular heart disease. *See also specific conditions* in carcinoid syndrome, 564, 565
cardiac imaging in, 270e-15–16
drug-induced, 565
dyspnea in, 47e-3
global burden of, 1528–1529
multiple and mixed. *See* Multiple/mixed valvular heart disease
murmurs in, **51e-1**
natural history of, 1439
in polycystic kidney disease, 1852
in pregnancy, 47–48
radiation-induced, 621
in relapsing polycondritis, 2203–2204
- VAMP2 (synaptobrevin), 985
- vanA* gene, 961
- van Buchem's disease (hyperostosis corticalis generalisata), 426e-5
- Vancomycin
action of, 930, 931t
adverse effects of, 491, 727t, 938t, 942
cutaneous, 379, 381
fever, 138
nephrotoxicity, 1803
for asymptomatic GAS carriage, 966
for burn-wound infections, 166e-3
for *C. difficile* infections, 860, 860t, 861
for catheter-related bloodstream infections, 487, 487t
for clostridial infections, 991t, 994
for community-acquired pneumonia, 808t
dosage monitoring, 36
empirical uses of, 767t
for enterococcal infections, 975t
for epidural abscess, 767t, 780t
for febrile neutropenia, 103e-23, 491
for health care–associated pneumonia, 812t
indications for, 940t, 942
for infective endocarditis
as empirical therapy, 767t, 780t, 824
enterococcal, 821, 822t, 975t
staphylococcal, 822t, 823, 963
streptococcal, 822t
for intraabdominal infection, 767t
for intracranial infections, 780t
for lung abscess, 815
for meningitis
as empirical therapy, 767t, 780t, 887t
enterococcal, 975t
MRSA, 889t, 890
pneumococcal, 889, 952
for MRSA infections, 961, 962t, 963
community-acquired pneumonia, 808t
as empirical therapy, 767t
endocarditis, 823
meningitis, 889t, 890
skin, 832t
for necrotizing fasciitis, 832t
for nongonococcal bacterial arthritis, 835
for osteomyelitis, 840t, 841t
for postsplenectomy sepsis, 780t
in pregnancy and lactation, 937t
prophylactic, 945t
for purpura fulminans, 780t
resistance to. *See also* Vancomycin-resistant enterococci (VRE)
mechanisms of, 931t, 933–934, 961, 976
rates of, 940t, 942
for sepsis/septic shock, 767t, 780t, 1757t
for staphylococcal infections, 962t
therapeutic monitoring of, 2766t
for toxic shock syndrome, 780t, 963
- Vancomycin-intermediate *S. aureus* (VISA), 933, 961
- Vancomycin-resistant enterococci (VRE)
diagnosis of, 774
epidemiology of, 973
genomics of, 769t, 774
global considerations in, 973
in health care–associated infections, 973
laboratory identification of, 150e-8
mechanism of, 933–934, 976
treatment of, 975t
- van den Bergh reaction, 280
- Vandetanib, 102e-5t, 103e-18t, 2342
- van* gene, 774
- Vanilloids, 262e-4
- Vanishing bile duct syndrome, 284, 2027–2028
- VAP. *See* Ventilator-associated pneumonia (VAP)
- Vaptans (vasopressin antagonists)
for hyponatremia, 64e-5, 302, 2281–2282
for SIADH, 2282f, 2283
- Vardenafil
action of, 324, 325f, 326f
adverse effects of, 329t
characteristics of, 329t
contraindications to, 328, 329t
for erectile dysfunction, 328, 2672
for lower urinary tract symptoms, 587
- Varenicline, 507, 1705, 2732
- Variable expressivity, 436–437
- Variable number of tandem repeats (VNTRs), 441
- Variant of unknown significance (VUS), 433, 448
- Variceal bleeding
esophageal. *See* Esophageal varices
gastric, 346e-1v, 1891, 1891f, 2097
- Varicella (chickenpox), **1183**
clinical features of, 131t, 341t, 1184
extracutaneous, 1184
oral, 238t
skin
characteristics of, 131t, 134
evolution of, 25e-8f, 1184, 1184f
rash, 361t, 362
vesicles, 827
complications of, 1184, 2673
congenital, 1184
differential diagnosis of, 134, 1185
epidemiology of, 131t, 1183
health care–associated, 913
incidence pre- and postvaccine, U.S., 785t
infection control measures, 917
pathogenesis of, 1183
perinatal, 1184
prevention of, 1186, 1186t. *See also* Varicella vaccine
treatment of, 215e-7, 1185–1186
- Varicella vaccine
contraindications for, 789t
precautions for, 789t
recommended administration of
in adults, 29t, 787–788f
in children, 1186
for health care–associated exposure, 917
in pregnancy, 788f
storage and handling of, 791
- Varicella-zoster immune globulin (VZIG), 917, 1186, 1186t, 1252t
- Varicella-zoster virus (VZV), 342, 1183, 1185
- Varicella-zoster virus (VZV) infections. *See also* Herpes zoster (shingles); Varicella (chickenpox)
in cancer patient, 488, 492
encephalitis, 488, 894, 895t
etiology of, 1183
meningitis, 892
neuropathy in, 2685
pathogenesis of, 1183
prevention of, 1186, 1186t
in transplant recipient, 139e-4, 921, 921t
treatment of, 1185–1186
- Varicella-zoster virus (VZV) vaccine, 485t, 893, 921, 929, 1186
- Varicocele, 2360
- Varicose veins, 1650. *See also* Chronic venous disease
- Variegate porphyria (VP)
clinical features of, 361t, 362, 2522t, 2531, 2681
diagnosis of, 2522t, 2531
global considerations in, 2521, 2531, 2681
pathophysiology of, 2522f, 2681
vs. porphyria cutanea tarda, 362
treatment of, 2531, 2681
unsafe drugs in, 2526, 2527–2529t
- Variola virus, 261e-6. *See also* Smallpox
- Varioliform gastritis, 1931
- Varus, 2221, 2228, 2228f
- Vasa recta, 332e-2
- Vascular access-related infections. *See* Catheter-related bloodstream infections
- Vascular addresses, 102e-17f
- Vascular cell adhesion molecule-1 (VCAM-1), 89e-1, 1242, 2150
- Vascular dementia, 170, 172, 173t, 2602
- Vascular disease. *See also specific diseases*
aging and, 94e-4
gene therapy in, 91e-4
ophthalmological manifestations of, 41e-1v
- Vascular ectasias
bleeding in, 277
diagnosis of, 1891, 1892f
treatment of, 1897, 1897f
- Vascular endothelial growth factor (VEGF)
in diabetic retinopathy, 2424
in gene therapy, 91e-4
in high-altitude illness, 476e-1
in kidney development, 332e-1
in preeclampsia, 45
in stem cell therapy, 90e-3
in systemic sclerosis, 2156
in tumor angiogenesis, 102e-14–15, 102e-14f, 102e-15f, 102e-16f
- Vascular endothelial growth factor (VEGF) antibodies, 103e-21, 103e-21t
- Vascular endothelial growth factor (VEGF) inhibitors, 91e-4
- Vascularization, in tissue engineering, 92e-1
- Vascular mimicry, 102e-13
- Vascular resistance, 1463
- Vascular tone, 265e-4

- Vasculitic neuropathy, 2700
- Vasculitis. *See also* Vasculitis syndromes
 anemia and, 630f
 as cause of fever of unknown origin, 137t, 141
 CNS
 clinical features of, 2192
 dementia in, 2607
 differential diagnosis of, 2192, 2568
 imaging in, 386e-4f, 386e-6f, 441e-19f, 2192f
 definition of, 2179
 diagnosis of, 2181
 differential diagnosis of, 2181, 2181t
 drug-induced, 382, 2193, 2218t
 in familial Mediterranean fever, 2214
 leukocytoclastic (cutaneous small-vessel), 25e-11f, 76e-15f, 133t, 135, 364t, 368
 necrotizing. *See* Necrotizing vasculitis
 of nerve and muscle, 619
 paraneoplastic, 614t, 619
 in relapsing polychondritis, 2204
 rheumatoid, 2138
 secondary, 2179t, 2193
 in serum sickness, 2193
 in serum sickness–like reactions, 2193
 single-organ, 2191
 in Sjögren's syndrome, 2167, 2167t
 skin manifestations of, 76e-15f, 364t, 366
 treatment of, 2181–2182
 urticarial, 132t, 135
- Vasculitis syndromes, **2179**
 approach to the patient, 2181–2182, 2181f
 Behçet's syndrome. *See* Behçet's syndrome
 classification of, 2179, 2179t
 Cogan's syndrome. *See* Cogan's syndrome
 cryoglobulinemic. *See* Cryoglobulinemia
 cutaneous, 386e-6f
 definition of, 2179
 diagnosis of, 386e-1, 386e-1–6f, 2181, 2181f
 differential diagnosis of, 2181, 2181t
 diseases associated with, 2193
 eosinophilic granulomatosis with polyangiitis.
 See Eosinophilic granulomatosis with
 polyangiitis (Churg-Strauss syndrome)
 giant cell arteritis. *See* Giant cell arteritis
 (temporal arteritis)
 granulomatosis with polyangiitis. *See*
 Granulomatosis with polyangiitis
 (Wegener's)
 idiopathic cutaneous, 2191–2192
 IgA vasculitis. *See* IgA vasculitis (Henoch-
 Schönlein)
 Kawasaki disease. *See* Kawasaki disease
 microscopic polyangiitis. *See* Microscopic
 polyangiitis
 pathogenesis of, 2179–2181, 2180t
 polyangiitis overlap syndromes, 2193
 polyarteritis nodosa. *See* Polyarteritis nodosa
 polymyalgia rheumatica. *See* Polymyalgia
 rheumatica
 primary CNS, 386e-4f, 386e-6f, 2192, 2192f
 Takayasu's arteritis. *See* Takayasu's arteritis
 treatment of, 2181–2182, 2181f, 2182t
 vessel damage in, 2179–2180
- Vasculogenesis, 102e-14
- Vasculopathy. *See specific types*
- Vasoactive intestinal peptide (VIP)
 actions of, 570
 ectopic production of, 609t
 in prolactin secretion, 401e-2
 reference values for, 2764t
 regulation of, 2090
 in swallowing, 255
- Vasoactive intestinal peptidoma (VIPoma).
 See VIPoma (vasoactive intestinal
 peptidoma)
- Vasoconstriction, 725
 cold-induced, 478e-1
 cyanosis in, 249
 heat conservation by, 124
 localized hypoxia and, 248
- Vasodilation, 725
 as adaptation to hypoxia, 248
 endothelium-dependent, 1615
 in exercising muscle, 265e-8
 heat loss by, 124
- Vasodilators
 circulating, in shock, 1746
 for heart failure, 1509, 1509f
 for hypertension, 1624t, 1625
- Vasodilator testing, 1657
- Vasogenic edema, 1777
- Vasointestinal polypeptide, 330
- Vasomotor rhinitis, 2122
- Vasopressin. *See also* Arginine vasopressin (AVP)
 action of, 332e-5f, 332e-6f
 in coordination of emesis, 103e-25
 deficiency, 2259t
 ectopic production of. *See* Syndrome of
 inappropriate antidiuretic hormone
 (SIADH)
 in shock, 1745
- Vasopressin antagonists. *See* Vaptans
 (vasopressin antagonists)
- Vasospasm, after subarachnoid hemorrhage,
 1785, 1787
- Vasovagal syncope. *See* Syncope, neurally
 mediated
- Vaughan-Williams classification, of antiarrhythmic
 drugs, 273e-5, 273e-6t
- VCAM-1 (vascular cell adhesion molecule-1),
 89e-1, 1242, 2150
- Vd (volume of distribution), 34, 935
- VDR (vitamin D receptor), 2464–2465
- VDRL (Venereal Disease Research Laboratory)
 test, 1137, 1137f, 1259
- Vedolizumab, 1962
- Vegetarian diet, 300, 626, 644
- Vegetations, valvular, 76e-12f, 816, 816f
- Vegetative state, 1771
- VEGF. *See* Vascular endothelial growth factor
 (VEGF)
- Veins. *See* Blood vessels
- VeIP regimen, for testicular cancer, 591
- “Velcro” crackles, 2161
- Vellus hair, 331
- Velocardiofacial (DiGeorge) syndrome, 83e-6,
 375e-1, 465e-2, 2113, 2483
- Vemurafenib
 action and targets of, 41, 101e-8, 102e-5t,
 103e-16f, 103e-17t, 103e-19
 adverse effects of, 103e-17t
 for hairy cell leukemia, 135e-2
 for lung cancer, 521
 for melanoma, 102e-5t, 103e-17t, 103e-19,
 498, 499
 relapse following treatment with, 32
 resistance to, 103e-23
- Venereal Disease Research Laboratory (VDRL)
 test, 1137, 1137f, 1259
- Venezuelan equine encephalitis virus, 1306t, 1317
- Venlafaxine
 adverse effects of, 7e-6, 2640t, 2682t, 2710t,
 2716–2717
 for anxiety disorders, 2711
 for cataplexy, 190
 for depression, 64, 2672, 2710t, 2716
 for hot flushes with androgen deprivation
 therapy, 7e-6
 for menopausal symptoms, 2383
 for migraine prevention, 2594t
 for neuropathic pain, 94–95, 2682t
 for pain, 91t, 94, 2672
 for panic disorder, 2709
 pharmacology of, 2716
 for PTSD, 2712
- Venomous snakebite. *See* Snakebites
- Venoocclusive disease, of liver. *See* Sinusoidal
 obstructive syndrome
- Venoocclusive disease with immunodeficiency
 (VODI) syndrome (Sp110 deficiency),
 375e-2, 2110
- Venous disorders, of extremities
 anatomic considerations, 1650
 chronic venous disease. *See* Chronic venous
 disease
 venous thrombosis. *See* Venous
 thromboembolism (VTE)
- Venous hum, 51e-6
- Venous hypertension, 357, 367
- Venous lake, 364t, 367
- Venous malformation, 364t, 367
- Venous O₂ concentration, 1748t
- Venous pressure, 253
- Venous return, 51e-7
- Venous sinus thrombosis, 2566
- Venous thoracic outlet syndrome, 122
- Venous thromboembolism (VTE), 277, **743**.
 See also Deep vein thrombosis (DVT);
 Pulmonary embolism (PE)
 in antiphospholipid syndrome, 2135, 2135t
 vs. arterial thrombosis, 744–745
 with brain tumor, 598
 in burn patient, 166e-2
 coagulation cascade in, 743–744
 cyanosis in, 249
 edema in, 253
 epidemiology of, 743
 etiology of, 277t, 743–744, 743t
 genetic considerations in, 277t, 743t, 744
 history in, 404–405
 in HIV infection, 1262
 incidence of, 274, 740
 in liver disease, 739–740, 739f, 739t
 lupus anticoagulant in, 740
 mesenteric, 1979, 1980t
 oral contraceptives and, 2391, 2391t
 paraneoplastic, 612t, 613
 pathophysiology of, 1631–1632, 1631f
 postmenopausal hormone therapy and, 2383,
 2384t
 postthrombotic syndrome, 1631, 1631f,
 1634
 prevention of
 fondaparinux for, 753
 in hospitalized patients, 1636, 1637t
 in mechanically ventilated patient, 1743
 in surgical patient, 55
 prothrombotic states and, 1631
 renal, 1866
 risk factors for, 405t, 1631
 acquired, 277t, 743t, 744
 age, 405f
 history of thrombosis, 404–405
 inherited, 277t, 743t, 744
 suppurative intracranial, 783, 905–906, 905f
 Ventilation-perfusion mismatch/heterogeneity,
 306e-4, 306e-4f
 in asthma, 47e-2
 in cardiogenic pulmonary edema, 47e-5
 in COPD, 47e-2
 in hypoxia, 247–248
 in interstitial lung diseases, 1710
 in pulmonary hypertension, 1656
 in sepsis/septic shock, 1755
- Ventilation-perfusion scan, 1664
- Ventilator-associated pneumonia (VAP)
 A. *baumannii*, 1037
 in burn patient, 166e-2
 clinical features of, 810
 complications of, 812
 diagnosis of, 810–811, 811t, 915
 epidemiology of, 810
 etiology of, 809, 809t
 failure to improve, 812
 follow-up, 812
 MRSA, 811, 812
 P. *aeruginosa*, 811, 812, 1044
 pathophysiology of, 810, 810t, 915, 1743
 prevention of, 12e-2, 12e-3, 813, 914–915,
 914t
 prognosis of, 812–813
 risk factors for, 914
 S. *maltophilia*, 1048
 treatment of, 811–812, 812t
- Ventilatory function
 adequacy of, 306e-3
 disturbances of. *See* Hyperventilation;
 Hypoventilation; Respiratory disorders
 flow-related properties of, 306e-1–2, 306e-2f,
 306e-3f
 measurement of, 306e-6

- physiology of, 1720, 1721f
 volume-related properties of, 306e-1-2, 306e-1f
 work of breathing, 306e-2-3
- Ventral respiratory column, 1720
- Ventricular afterload, 265e-8, 265e-9f
- Ventricular aneurysm, 1610
- Ventricular arrhythmias, **1489**
 approach to the patient, 1491, 1500
 characteristics of, 1489-1491, 1490f
 clinical features of, 1491
 diagnosis of, 1491, 1492t
 diseases associated with, 1495t
 in myocardial infarction, 1608-1609
 premature ventricular beats. *See* Premature ventricular contractions (PVCs)
 treatment of, 1498-1500, 1499f
 ventricular fibrillation. *See* Ventricular fibrillation (VF)
 ventricular tachycardia. *See* Ventricular tachycardia (VT)
- Ventricular couplets, 1489
- Ventricular dysplasia, arrhythmogenic. *See* Arrhythmogenic right ventricular cardiomyopathy (ARVC)
- Ventricular end-diastolic pressure, 265e-7
- Ventricular end-diastolic volume, 265e-7, 265e-9f
- Ventricular fibrillation (VF)
 cardiac arrest/sudden cardiac death in, 1764
 ECG in, 1491f
 etiology of, 1498
 in hypothermia, 478e-2, 478e-3
 in ischemic heart disease, 1580
 mechanism of, 273e-2, 273e-3t, 273e-4
 in myocardial infarction, 1608-1609
 postresuscitation care, 1770
 treatment of, 1498, 1769, 1769f
- Ventricular filling, assessment of, 265e-10
- Ventricular flutter, 1489. *See also* Ventricular tachycardia
- Ventricular function curve, 265e-7
- Ventricular premature beats, 1580, 1608
- Ventricular remodeling. *See* Left ventricular remodeling
- Ventricular septal defect (VSD)
 AV conduction block in, 1473
 cardiac catheterization in, 1463
 echocardiography in, 270e-25
 erythrocytosis in, 1523
 heart sounds/murmur in, 51e-1f, 51e-2-5, 51e-2f, 1449
 location of, 1521f
 natural history of, 1522
 in pregnancy, 47
 pregnancy and, 1522
 pulmonary hypertension and, 1522
 RV outflow tract obstruction in, 1522-1523
 treatment of, 1523
- Ventricular septal rupture, 1762, 1762t
- Ventricular tachycardia (VT)
 cardiac arrest/sudden cardiac death in, 1764
 definition of, 1489
 implantable cardioverter-defibrillator for, 273e-7
 incessant, 1498
 in ischemic heart disease, 1580
 LV intrafascicular, 1496
 mechanism of, 273e-3, 273e-3t, 273e-4
 monomorphic, 278e-13f, 1489, 1490f, 1491f
 in myocardial infarction, 1608
 nonsustained
 characteristics of, 1489, 1490f
 diagnosis of, 1492
 diseases associated with, 1493-1494
 prognosis of, 1491-1492
 treatment of, 1492-1494
 outflow tract, 1495-1496
 polymorphic
 in Brugada syndrome, 1497-1498. *See also* Brugada syndrome
 catecholaminergic, 147, 1498
 diseases associated with, 1496t
 drug-induced, 43, 1496t
 in early repolarization syndrome, 1498
 ECG in, 1491, 1491f, 1497f
 in genetic dilated cardiomyopathy, 1498
 in hypertrophic cardiomyopathy, 1498
 in long QT syndrome, 147, 1497, 1497f. *See also* Long QT syndrome
 mechanism of, 273e-3t, 273e-4
 in myocardial infarction/ischemia, 1496, 1608
 in short QT syndrome, 1497
 sinusoidal, 1489, 1491f
 sustained monomorphic
 characteristics of, 1489, 1491f, 1494
 diagnosis of, 1494
 diseases associated with, 1494-1496, 1496t
 idiopathic, 1495-1496
 in myocardial infarction, 1608
 prognosis of, 1494
 vs. supraventricular tachycardia, 1494, 1494f
 treatment of, 1494-1496
 treatment of
 antiarrhythmic drugs, 1498-1499
 catheter ablation, 1499-1500
 implantable cardioverter-defibrillators, 1499, 1499f
 ventricular fibrillation. *See* Ventricular fibrillation (VF)
 ventricular flutter, 1489, 1491f
- Ventriculography, 1463-1464, 1464f
- Ventriculostomy, 1779, 1780f
- VEPs (visual evoked potentials), 442e-3
- Verapamil
 for accessory pathway-mediated tachycardia, 1483
 adverse effects of, 1589t, 2595-2596, 2640t
 for aortic dissection, 1641
 for arrhythmias, 1479t, 1480t
 for atrial fibrillation, 1487
 for AV nodal reentrant tachycardia, 1479
 for cluster headache prevention, 2595, 2595t
 dosage of, 1479t, 1480t
 drug interactions of, 34t
 azoles, 246e-3t
 digoxin, 42t
 quinidine, 246e-6t
 ranolazine, 1590
 for hypertension, 1624t, 1625, 1821
 for hypnic headache, 2598
 for insulinoma, 569
 for ischemic heart disease, 1589t
 metabolism of, 34, 34t
 for NSTEMI-ACS, 1596
 overdose/poisoning with, 473e-9t
 pharmacology of, 1479t, 1480t
 for supraventricular arrhythmia, 1609
- Vermiculite dust, 1690
- Verner-Morrison syndrome. *See* VIPoma (vasoactive intestinal peptidoma)
- Verruca plana, 342f
- Verruca vulgaris, 342f, 351
- Verruga peruana, 829, 1081t, 1083
- Vertebrae
 anatomy of, 111, 112f
 "ivory," 426e-2
 ligament injury, 441e-33f
 "picture frame," 426e-2
 "squaring" of, 2169
- Vertebral artery, occlusion of, 2575-2577
- Vertebral fractures
 epidemiology of, 2488-2489, 2488f, 2489f
 estrogen deficiency and, 2492
 imaging in, 441e-31-34f
 in osteoporosis, 117
 pain in, 117, 2495
 prevention of
 bisphosphonates for, 2499-2500, 2500f
 denosumab for, 2501, 2502f
 teriparatide for, 2502, 2502f
 treatment of, 2495
- Vertebral osteomyelitis. *See* Osteomyelitis, vertebral
- Vertebroplasty, 2503
- Vertigo, **148**
 acute prolonged (vestibular neuritis), 149
 approach to the patient, 148-149
 benign paroxysmal positional, 149t, 150, 150f
 central, 149t
 evoked by loud sounds, 219
 in Ménière's disease. *See* Ménière's disease in migraine, 150
 in MS, 2663
 peripheral, 149t
 prevention of, 151
 psychosomatic, 151
 treatment of, 151, 151t
 in vestibular dysfunction, 149, 150-151
- Very-low-calorie diet, 2396
- Very-low-density lipoproteins (VLDLs)
 characteristics of, 1241t, 2436-2437, 2436f
 estimate of, 2446
 overproduction of, 2438-2439, 2451. *See also* Lipoprotein disorders
 in transport of hepatic lipids, 2437, 2437f
- Vesicants, **262e-1**, 262e-1t
- Vesicles/bullae, 339t, **361**
 clinical features of, 341t
 description of, 127, 339t
 drug-induced, 361
 etiology of, 361-362, 361t, 827, 828t, 829
 with fever, 131-132t, 134-135
 in herpes simplex virus infections. *See* Herpes simplex virus (HSV) infections
 infections associated with, 827, 828t, 829
 in smallpox. *See* Smallpox
 in varicella. *See* Varicella (chickenpox)
- Vesicoureteral reflux, 1859-1860, 1860f, 1871
- Vesiculoviruses, 1305, 1310t
- Vesiculovirus infections
 Chandipura virus, 1316
 vesicular stomatitis viruses, 1299, 1304, 1310t
- Vessel wall, 725, 732
- Vestibular dysfunction, 149, 149t, 151, 164
- Vestibular-evoked myogenic potential test, 223
- Vestibular migraine, 150
- Vestibular neuritis, 149, 149t. *See also* Vertigo
- Vestibular nystagmus, 210
- Vestibular rehabilitation therapy, 151
- Vestibular schwannoma (acoustic neuroma)
 diagnosis of, 149, 150, 220, 603
 facial nerve involvement in, 2648
 hearing loss in, 150, 220, 222
 imaging in, 223
 in neurofibromatosis, 603, 603f, 604, 2343
- Vestibulocochlear nerve, 2538
- Vestibuloocular reflex, 149
- Vestibulopathy, acute peripheral, 149
- Vestibulospinal pathway, 155f
- Veterans. *See* War veterans
- VHL gene mutations
 action of, 2329, 2332, 2343
 epigenetic silencing of, 102e-7
 in pheochromocytoma/paraganglioma, 2334f, 2335
 in renal cell carcinoma, 578
 in von Hippel-Lindau disease/syndrome. *See* von Hippel-Lindau disease/syndrome, genetic considerations in
- VHL protein, 2329
- Vibration injury, 1649
- Vibration sense, 159, 160t, 2539
- Vibratory angioedema, 2118
- Vibrio alginolyticus* infections, 1065t, 1066
- Vibrio cholerae*. *See also* Cholera
 adherence of, 145e-3, 852
 epidemiology of, 1061-1062, 1062f
 genome of, 777, 777f, 1063
 laboratory identification of, 150e-2t, 1063
 microbiology of, 1061
 non-O1/O139, 1065-1066, 1065t
 serotypes of, 1061-1062
 toxin production by, 1062-1063
- Vibrio parahaemolyticus* infections, 151e-8, 856t, 1065, 1065t
- Vibrio vulnificus* infections, **1066**
 clinical features of, 132t, 135, 782, 1066, 1755
 epidemiology of, 1065t, 1066
 treatment of, 1066

- Vici syndrome, 375e-1
 Vidarabine, 215e-3t, 215e-9, 417
 Video-assisted thoracic surgery, 1668–1669
 Video capsule endoscopy, 279
 Video game training, for cognitive training, 444e-11f, 444e-12
vif gene, 225e-3f, 1219f
 Vif protein, 225e-2
 Vigabatrin, 618
 Vilazodone, 2710t
 Vildagliptin, 2414t
 Villous adenoma, 254, 305
 Vimentin, 102e-11f, 372e-24t
 Vinblastine
 action of, 103e-14
 adverse effects of, 103e-12t
 alopecia, 380
 gonadal dysfunction, 103e-25
 myelotoxicity, 103e-14
 neurologic, 1793, 2686t
 phototoxicity, 389t
 Raynaud's phenomenon, 1649
 for bladder cancer, 577
 drug interactions of, 103e-12t
 for testicular cancer, 591
 Vinca alkaloids. *See also specific drugs*
 adverse effects of, 380, 622, 2686t
 for breast cancer, 531
 characteristics of, 103e-11–12t, 103e-14–15
 drug interactions of, 246e-3t
 Vincent's angina (trench mouth), 231, 233, 236, 238t, 1097
 Vincristine
 action of, 103e-14
 adverse effects of, 103e-11t, 103e-14
 depression, 63
 late, 620t
 nail disorders, 380
 neurologic, 607, 607t, 1793
 neuropathy, 103e-14, 622, 2686t
 for Ewing's sarcoma, 119e-4
 for gastric lymphoma, 536
 genetic variations in response to, 39t
 for gestational trophoblastic disease, 124e-5
 for lung cancer, 522
 for malignant pheochromocytoma, 2331
 for oligodendroglioma, 601
 for soft tissue sarcoma, 119e-2
 for thrombotic thrombocytopenic purpura, 730
 Vinorelbine
 action of, 103e-14
 adverse effects of, 103e-12t, 1798, 2686t
 for lung cancer, 518t, 520t, 522
 resistance to, 103e-14
 Vinyl chloride, 283, 477t, 2218t
 Violence
 assessment of, 2722–2723
 domestic. *See* Domestic abuse/violence
 VIP. *See* Vasoactive intestinal peptide (VIP)
 Vipers, 2734, 2737, 2738. *See also* Snakebites
 VIPoma (vasoactive intestinal peptidoma), 570
 clinical features of, 269, 305, 559t, 570, 2338
 diagnosis of, 570
 incidence of, 561
 locations of, 559t, 570
 in MEN 1, 2338
 treatment of, 570–571, 2338
 Viral encephalitis. *See* Encephalitis, viral
 Viral hemorrhagic fevers, 1320. *See also specific diseases*
 in bioterrorism, 261e-8
 clinical features of, 133t, 261e-5t, 261e-8, 782, 1320
 rash, 135
 diagnosis of, 261e-5t, 1320
 epidemiology of, 133t, 783
 etiology of, 782–783, 1320–1323
 pathogenesis of, 1320
 prevention of, 261e-5t, 261e-8
 treatment of, 215e-6, 261e-5t, 261e-8
 Viral hepatitis. *See* Hepatitis
 Viral infections. *See also specific diseases*
 arthropod-borne. *See* Arthropod-borne viruses
 cancer development and, 101e-7, 214e-7
 as cause of fever of unknown origin, 137t
 diagnosis of, 214e-8–9. *See also* Virus(es), laboratory identification of
 interferon production in, 214e-8
 persistent and latent, 214e-7
 in pregnancy, 50
 prevention of, 214e-9
 primary, 214e-6
 renal involvement in, 1849
 resistance to, 214e-8
 rodent-borne. *See* Rodent-borne viruses
 transmission of, 214e-6
 in transplant recipient
 after HSCT, 920t, 921–923, 921t
 after kidney transplantation, 924–926, 924t, 1829–1831, 1830t
 after liver transplantation, 924t, 927–928, 1341f, 2072–2073
 malignancies and, 928
 treatment of, 214e-9
 upper respiratory tract, 225–226
 Viral load, 150e-8
 Viral meningitis. *See* Meningitis, viral
 Viral orchitis, 2365
 Viral protein U (Vpu), 145e-9
 Viral (subacute) thyroiditis, 2298–2299, 2298f
 Virchow-Robin spaces, 906
 Virchow's node, 282, 286, 408, 555
 Viridans streptococci, 940t, 963, 964t, 971
 Virilization, 331
 Viroids, 214e-1
 Virtual bronchoscopy, 1666, 1666f
 Virtual colonoscopy, 540, 1898–1899, 1899f
 Virtue ethics, 17e-2
 Virulence, 145e-1
 Virus(es). *See also specific viruses*
 adherence of, 145e-1–2, 145e-2t
 bacterial hosts of, 86e-4
 cell entry by, 214e-1–2
 cell/tissue invasion by, 145e-7, 214e-13
 cytopathic effects and apoptosis inhibition, 214e-6
 definition of, 214e-1
 egress of, 214e-5
 gene expression and replication in, 214e-4–5
 growth after entry, 145e-4, 214e-5
 host range of, 214e-6
 host receptors of, 145e-3
 in human microbiome, 86e-4
 laboratory identification of
 ELISA, 214e-8
 genomics in, 769t, 772
 nucleic acid amplification, 214e-9
 specimen collection and transport, 150e-5t
 specimen-processing algorithm, 150e-6f
 tissue culture, 214e-9
 Western blot, 214e-8
 mutations of, 214e-5
 oncolytic, 91e-4
 protein activity in, 214e-5–6
 structure of, 214e-1, 214e-1f
 taxonomy of, 214e-1, 214e-2t, 214e-3f
 as therapeutic tool or agents, 91e-1t, 214e-9
 vaccines for, 214e-9
 virulence mechanisms of, 145e-9
 Virus-like particle(s), 86e-4
 Virus-like particle vaccines, 929
 Virusoids, 214e-1
 VISA (vancomycin-intermediate *S. aureus*), 933, 961
 Visceral afferent hypersensitivity, 261
 Visceral larva migrans. *See* Larva migrans, visceral
 Visceral leishmaniasis, 1387. *See also* Leishmania spp. infections
 clinical features of, 152e-3t, 1389, 1390f
 diagnosis of, 152e-3t, 245e-4t, 245e-6t, 1389, 1389f
 differential diagnosis of, 1390
 epidemiology of, 1387–1388, 1388t, 1389f
 in HIV infection, 798, 1268, 1389, 1391
 immunopathogenesis of, 1389
 incubation period for, 152e-3t
 prognosis of, 1391
 in returning war veterans, 152e-3t, 152e-8
 skin lesions after, 1391–1392, 1391t, 1392f
 treatment of, 152e-3t, 152e-7, 246e-10–11, 1390–1391
 Visceral pain, 60
 Vismodegib, 102e-5t, 103e-19t, 500, 500f
 Visual acuity, 149, 195, 196f, 2537
 Visual aura, 204
 Visual cortex, 176f
 Visual evoked potentials (VEPs), 442e-3
 Visual function, assessment of
 color vision, 197
 eye movements and alignment, 197
 pupils, 195–197, 196f
 refractive state, 195
 stereopsis, 197
 visual acuity, 195, 196f
 visual fields, 197–199, 198f, 2537
 Visual loss
 chemotherapy-induced, 607t
 chronic/permanent
 in cataract. *See* Cataract
 in central serous chorioretinopathy, 205
 in diabetic retinopathy. *See* Diabetic retinopathy
 in epiretinal membrane, 206
 in glaucoma. *See* Glaucoma
 in macular degeneration. *See* Macular degeneration
 in retinitis pigmentosa, 206, 206f, 431t
 trachoma and, 1172
 tumor-related, 206–207, 206f
 CMV retinitis and, 1192
 cortical, 199
 factitious, 204
 in giant cell arteritis, 108
 HSV keratitis and, 1178
 in legal blindness, 195
 in lepromatous leprosy, 1125
 with sellar masses, 2261
 transient or sudden, 200–202, 201f
 Visual object agnosia, 182
 Visual target cancellation test, 180
 Visual vertigo, 151
 Vital capacity, 306e-2, 306e-2f
 Vitamin(s), 96e-1. *See also specific vitamins*
 absorption of, 1933
 assessment of nutritional status, 462
 body stores of, 96e-1
 deficiencies of, 96e-1t, 2398
 functions of, 96e-2–3f
 genetic disorders related to, 435e-1t
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference ranges for, 2766t
 in specialized nutritional support formulas, 98e-5
 structures of, 96e-2–3f
 Vitamin A, 96e-6
 deficiency of. *See* Vitamin A deficiency
 dietary sources of, 96e-6
 functions of, 96e-3f, 96e-6
 metabolism of, 96e-6
 parenteral requirement for, 98e-5t
 in pregnancy, 96e-7
 recommended intake of, 95e-2t
 reference range for, 2766t
 for retinitis pigmentosa, 206
 structure of, 96e-3f
 supplements. *See* Vitamin A supplements
 toxicity of, 96e-7, 463t, 2478
 Vitamin A deficiency
 clinical features of, 96e-1t, 96e-6, 463t
 esophageal cancer risk and, 533
 global considerations in, 96e-6, 461
 keratitis in, 199
 oral manifestations of, 236
 in pregnancy, 265e-12
 risk factors for, 96e-1t

- Vitamin A supplements, 96e-7
 for bladder cancer prevention, 575
 for measles treatment, 1298
 for olfactory dysfunction, 216
- Vitamin B₁. *See* Thiamine
- Vitamin B₂. *See* Riboflavin
- Vitamin B₃. *See* Niacin
- Vitamin B₅. *See* Pantothenic acid
- Vitamin B₆. *See* Pyridoxine
- Vitamin B₁₂. *See* Cobalamin
- Vitamin B_{12A}. *See* Hydroxocobalamin (vitamin B_{12A})
- Vitamin C, **96e-5**
 absorption of, 96e-5
 deficiency of, 96e-1t, 96e-5, 236, 461, 463t. *See also* Scurvy
 dietary sources of, 96e-5
 functions of, 96e-3f, 96e-5, 367
 for macular degeneration, 205
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference range for, 2766t
 structure of, 96e-3f
 supplements, 96e-5, 1867, 2237
 toxicity of, 96e-5
- Vitamin D, **96e-7, 2463**
 beneficial effects of, 387
 deficiency of. *See* Vitamin D deficiency
 functions of, 96e-3f, 96e-7
 for hypocalcemia, 2488
 for hypoparathyroidism, 2484
 malabsorption of, surgery-related, 1927
 metabolism of, 2463-2464, 2464f
 parenteral requirement for, 98e-5t
 photochemistry of, 387
 recommended intake of, 95e-2t
 reference range for, 2766t
 resistance to, 315, 2485
 sources of, 96e-7
 structure of, 96e-3f
 supplements. *See* Vitamin D supplements
 synthesis of, 387, 2463-2464, 2463f
 toxicity of, 96e-8, 2477
- Vitamin D deficiency
 anticonvulsants and, 2485
 asthma due to, 1670
 clinical features of, 96e-1t, 463t, 2465
 diagnosis of, 315, 2465-2466
 in elderly, 75, 2465
 etiology of, 2465, 2465t
 genetic considerations in, 2465
 global considerations in, 315
 hypocalcemia and, 314, 2485
 hypomagnesemia and, 2462
 intestinal disease and, 2465
 muscle weakness in, 462e-19
 nonskeletal disease and, 96e-7
 oral manifestations of, 236
 osteomalacia due to. *See* Osteomalacia
 osteoporosis and, 2491
 prevalence of, 2254t
 rickets due to. *See* Rickets
 risk factors for, 96e-1t, 96e-7
 screening/testing for, 2254t
 treatment of, 96e-8, 315, 461, 2462, 2466
 in women, 6e-4
- Vitamin D receptor (VDR), 2464-2465
- Vitamin D supplements, 96e-8
 in elderly, 29t, 75, 80, 84
 for hypocalcemia, 315
 for MS, 2671
 for osteoporosis prevention/management, 2383, 2497
 for psoriasis, 354t
 for vitamin D deficiency, 2466
- Vitamin E, **96e-8**
 absorption of, 96e-8
 deficiency of. *See* Vitamin E deficiency
 dietary sources of, 96e-8
 functions of, 96e-3f, 96e-8
 for macular degeneration, 205
 metabolism of, 96e-8
 for NALFD, 2057
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t, 96e-8
 reference range for, 2766t
 structure of, 96e-3f
 supplements. *See* Vitamin E supplements
 toxicity of, 96e-8
- Vitamin E deficiency
 ataxic disorders in, 2630
 clinical features of, 96e-1t, 96e-8, 462e-19
 diagnosis of, 2690
 familial, 96e-8, 451e-2t, 2690
 neuropathies in, 462e-19, 2690
 risk factors for, 96e-1t, 96e-8
 treatment of, 96e-8, 2690
- Vitamin E supplements
 adverse effects of, 403
 for dysmenorrhea, 338
 for menopausal symptoms, 2383
- Vitamin K, **96e-8**
 deficiency of. *See* Vitamin K deficiency
 dietary sources of, 96e-8
 functions of, 96e-3f, 96e-8, 738
 high-dose, for hepatocellular carcinoma, 551
 metabolism of, 736, 736f
 parenteral requirement for, 98e-5t
 recommended intake of, 95e-2t
 reference range for, 2766t
 in specialized nutritional support formulas, 98e-6-7
 structure of, 96e-3f
 supplements. *See* Vitamin K supplements
 toxicity of, 96e-9
 for warfarin necrosis, 379
- Vitamin K antagonists, 754. *See also* Warfarin
- Vitamin K cycle, 736f, 754
- Vitamin K deficiency, **738**
 bleeding in, 403
 clinical features of, 96e-1t, 96e-8, 463t
 etiology of, 738
 risk factors for, 96e-1t, 96e-8
 treatment of, 738-739
- Vitamin K epoxide reductase (VKOR), 736, 754
- Vitamin K supplements, 96e-8-9
 for coagulation disorders in liver disease, 739, 2067
 for neonates, 738
 for osteoporosis management/prevention, 2497
 for warfarin-induced bleeding, 756
- Vitiligo
 APS-2 and, 2347t
 clinical features of, 76e-3f, 341t, 344f, 357t, 358t
 diagnosis of, 343-344, 358t
 disorders associated with, 357, 2430
 inflammasome activity in, 372e-8t
 pathophysiology of, 358t, 372e-24t
 treatment of, 358t, 391
- Vitreous disorders, 204
- Vittiforma* spp., 1408
- VKORC1 polymorphisms, warfarin dose requirements and, 39t, 41, 755, 755t
- VLA-4, 89e-1
- VLDLs. *See* Very-low-density lipoproteins (VLDLs)
- VNTRs (Variable number of tandem repeats), 441
- Vocal tic, 2623
- VODI (venoocclusive disease with immunodeficiency) syndrome (Sp110 deficiency), 375e-2, 2110
- Voglibose, 2414t
- Vogt-Koyanagi-Harada syndrome, 357, 357t, 359, 909t
- Voiding cystourethrography, 1872, 1873
- Volume of distribution (V_d), 34, 935
- Voluntary active euthanasia, 67t
- Volutrauma, ventilator-induced, 1731
- Volvulus, 1892, 1894f, 1982, 1985
- Vomiting. *See also* Nausea and vomiting
 in acute cholecystitis, 2080
 after head injury, 457e-4
 in alcohol use disorder, 2091
 definition of, 258
 in gastroenteritis. *See* Gastroenteritis
 hypokalemia in, 307
- hyponatremia in, 298
 hypovolemia in, 298
 in intestinal obstruction, 1983
 Mallory-Weiss tear following. *See* Mallory-Weiss tears
 mechanisms of, 258
 metabolic alkalosis in, 321
 water loss in, 95e-1
- von Gierke's disease (glucose-6-phosphatase deficiency; type I GSD), 433e-1, 433e-2t, 433e-3, 436e-5f
- von Hippel-Lindau disease/syndrome
 clinical features of, **2342**
 CNS tumors, 451e-2t, 599t, 1850t
 hypertension, 1620t
 neuroendocrine tumors, 562
 pheochromocytoma, 2332, 2333f
 pNETs, 562, 562t
 renal, 578, 1850t, 1854, 2343
 diagnosis of, 441e-37f, 1854, 2333f
 genetic considerations in
 genotype-phenotype correlation, 435
 locus and phenotype, 101e-6t, 451e-2t, 562t
 transcription factor mutations, 431t
 VHL gene mutations, 578, 599t, 1620t, 1850t, 1854, 2332, 2337t, 2343
 treatment of, 1854
- von Recklinghausen's disease. *See* Neurofibromatosis type 1 (NF1)
- von Willebrand disease, **730**
 acquired, 677, 730
 bleeding in, 403
 clinical features of, 730-731
 diagnosis of, 731f, 731t
 inheritance patterns in, 730-731, 731f
 pathophysiology of, 403
 subtypes of, 730, 731t
 treatment of, 731-732
- von Willebrand factor
 in platelet activation, 275-276, 275f, 400, 741-742, 741f
 reference values for, 2756t
 in thrombotic thrombocytopenia purpura, 729, 729f
 in von Willebrand disease, 730. *See also* von Willebrand disease
- von Willebrand factor concentrate, 731
- Vorapaxar, 749
- Voriconazole
 adverse effects of, 362, 379, 1331
 for *Aspergillus* infections, 1348, 1349t
 for *Blastomyces* infections, 1338-1339
 for *Candida* infections, 487, 1344t
 for *Coccidioides* infections, 1336
 contraindications to, 1331
 for fungal infections in cancer patient, 491
 for *Fusarium* infections, 1356
 genetic variations in response to, 39t, 1331
 indications for, 1331
 resistance to, 1348
 susceptibility testing for, 150e-8
 for *Trichosporon* infections, 1357
- Vorinostat
 action and targets of, 102e-5t, 102e-8, 103e-18t, 103e-19-20
 adverse effects of, 103e-18t
 for mycosis fungoides, 102e-5t, 103e-20
- Vortioxetine, 2710t
- vpr gene, 225e-3f, 1219f
- Vpr protein, 225e-2
- Vpu (viral protein U), 145e-9
- vpu gene, 225e-3f, 1219f
- Vpu protein, 225e-2
- Vpx protein, 225e-2
- VR, 262e-1t
- VRE. *See* Vancomycin-resistant enterococci (VRE)
- VSD. *See* Ventricular septal defect (VSD)
- V sign, 2195
- v-src gene mutations, 101e-2
- VT. *See* Ventricular tachycardia (VT)
- VTE. *See* Venous thromboembolism (VTE)
- "VT storm," 1498
- Vulvar cancer, 351, 1198

- Vulvar warts, 1198, 1199f
- Vulvovaginal infections
- bacterial vaginosis. *See* Bacterial vaginosis
 - candidiasis. *See* *Candida* spp. infections, vulvovaginal
 - clinical features of, 60e-1, 872-874, 873t, 875
 - etiology of, 870t, 875-876
 - gonococcal, 1005
 - Trichomonas vaginalis*. *See* *Trichomonas vaginalis* infections
- VUS (variant of unknown significance), 433, 448
- VX. *See* Nerve agents
- VZIG. *See* Varicella-zoster immune globulin (VZIG)
- VZV. *See* Varicella-zoster virus (VZV)
- W**
- Waardenburg's syndrome, 220, 222t, 359, 431t
- Wada test, 2557
- WAGR syndrome, 431t
- Waist circumference
- in metabolic syndrome, 2450t, 2451
 - in obesity evaluation, 2393, 2394t
 - over lifespan, 74f
- Waist-to-hip ratio, 2392
- Wakefulness, 185
- Waldenström's hypergammaglobulinemia purpura, 368
- Waldenström's macroglobulinemia, 717
- clinical features of, 717-718
 - genetic considerations in, 717-718
 - immunology of, 698f
 - paraneoplastic syndromes in, 372, 618-619
 - purpura in, 368
 - Raynaud's phenomenon in, 1649
 - treatment of, 718
 - vascular purpura in, 732
 - von Willebrand disease in, 731
- Walker-Warburg syndrome, 462e-9, 462e-10t
- Walking speed, 78t
- Walled-off necrosis, pancreatic, 2093t, 2095f, 2097
- Wallenberg's syndrome, 2576f, 2577
- Walrus bite-wound, 167e-1
- Warburg effect, 102e-12, 102e-13f
- Warfarin, 754
- action of, 754
 - adverse effects of
 - alopecia, 354
 - bleeding, 756, 1635
 - calciophylaxis, 1816
 - fetal, 47
 - in pregnancy, 756
 - skin necrosis
 - clinical features of, 76e-16f, 368, 379, 379f, 384t, 756
 - genetic considerations in, 756
 - pathogenesis of, 379
 - treatment of, 756
 - for antiphospholipid syndrome, 2136
 - distribution of, 34
 - dosage of, 40, 755, 755t
 - drug interactions of
 - amiodarone, 34t, 42t
 - azoles, 246e-3t
 - ciprofloxacin, 246e-8
 - isoniazid, 205e-3
 - metronidazole, 246e-5t, 939t
 - nafcillin, 939t
 - NSAIDs, 43
 - orlistat, 96e-8
 - PPIs, 1921
 - SSRIs, 2711t
 - sulfonamides, 246e-4t
 - tetracyclines, 246e-7t
 - trimethoprim, 246e-4t, 939t
 - for DVT/PE treatment, 1635, 1635t
 - genetic variations in response to, 39t, 41, 451, 754, 755, 755t
 - management of
 - before cardiac catheterization, 1461
 - before endoscopic procedures, 1889t
 - preoperative, 756
 - treatment monitoring, 755
 - metabolism of, 34t
 - in patient with heparin-induced thrombocytopenia, 728
 - pharmacology of, 754-755
 - in pregnancy, 756
 - for secondary prevention of myocardial infarction, 1610-1611
 - for stroke prevention in atrial fibrillation/atrial flutter, 1488, 2570
 - for thrombophlebitis, 613
 - for VTE prophylaxis, 1637t
- Wart(s), 351
- etiology of, 351, 829
 - flat, 351
 - genital, 351, 829, 1198, 1199f, 1200-1201. *See also* Condyloma acuminata (anogenital warts)
 - oral, 240t
 - plantar, 351
 - treatment of, 351
- Warts, hypogammaglobulinemia, infections, and myelokathexis (WHIM) syndrome, 415, 2111
- War veterans
- infections in, 152e-1, 152e-1t, 152e-2-6t
 - chronic, 152e-8-9
 - diarrheal disease, 152e-8
 - differential diagnosis of, 152e-7f
 - enteric pathogens, 152e-8
 - head injury-related, 152e-9
 - leishmaniasis, 152e-7-8
 - malaria, 152e-1, 152e-7
 - osteomyelitis, 152e-9
 - rabies, 152e-7
 - soft tissue/wound, 152e-9
 - viral hepatitis, 152e-7
 - neuropsychiatric illnesses in, 471e-1
 - approach to the patient, 471e-3-4, 471e-4t, 471e-5t
 - concussion, 471e-2, 471e-2t
 - epidemiology of, 471e-1-3
 - postwar symptoms, 471e-1
 - PTSD, 471e-2-3, 471e-4-5
 - stigma and barriers to care for, 471e-3
 - traumatic brain injury, 471e-2-3, 471e-2t, 471e-5-6
 - treatment of, 471e-4-6
- WASP gene, 2104, 2110
- Wasp sting, 2116, 2748-2749
- Wasting syndrome, 81, 1268, 1704
- Water
- renal transport of, 332e-9-10, 332e-10f
 - requirement for, 95e-1
- Water balance
- in chronic kidney disease, 1814
 - regulation of, 295-296, 332e-9-10, 332e-10f
- Waterborne disease, climate change and, 151e-6-8, 151e-7f
- Water brash, 262, 1901
- Water-depletion heat exhaustion, 479e-2
- Water deprivation, for hyponatremia, 64e-5, 302
- Water deprivation test, 64e-3-4, 295
- Water diuresis, 64e-3, 2275, 2276f
- Water-hammer (Corrigan's) pulse, 51e-5, 1445, 1535
- Water intoxication, 332e-9, 332e-10f
- Water loss, 95e-1
- insensible, 298
 - renal, 294-295
- Watermelon stomach, 277, 2161
- Waxy casts, urinary, 294
- WDHA syndrome. *See* VIPoma (vasoactive intestinal peptidoma)
- Weakness, 154
- abnormalities associated with, 154
 - acute, 2199
 - in botulism, 2703
 - in congenital muscular dystrophies, 462e-10, 462e-10t
 - in critically ill patient, 1734-1735, 1784
 - definition of, 154
 - distal, 157
 - in distal myopathies, 462e-12, 462e-13t
 - distribution of, 154
 - in endocrine disorders, 462e-18-19
 - evaluation of, 156f, 462e-1-2, 462e-1f, 462e-2f, 462e-3t
 - facial. *See* Facial weakness/paralysis falls and, 163
 - vs. fatigue, 151
 - generalized (quadriparesis), 157, 157t
 - in glycogen storage disorders, 462e-14
 - in Guillain-Barré syndrome, 2694
 - in hemiparesis, 156
 - in hypokalemia, 64e-7, 64e-8, 298, 307
 - in hypokalemic periodic paralysis, 462e-16
 - in limb-girdle muscular dystrophies, 462e-6t, 462e-7t, 462e-8
 - lower motor neuron (neuropathic), 154, 154t, 155f
 - in malabsorption syndromes, 1946t
 - in monoparesis, 157
 - in MS, 2663, 2672
 - in myasthenia gravis, 2701-2702
 - myopathic, 154t, 155-156
 - in nemaline myopathy, 462e-12
 - neuromuscular junction, 155
 - in paraparesis, 156-157
 - in parathyroid disorders, 462e-19
 - patterns of, 462e-1
 - in peripheral neuropathies, 2675, 2675t
 - in progressive diaphyseal dysplasia, 426e-5
 - in progressive muscular dystrophies, 462e-5-6, 462e-5t, 462e-8, 462e-9, 462e-11
 - proximal, 157
 - psychogenic, 154t, 156
 - in restricted distribution, 157
 - subacute or chronic progressive, 2197-2198
 - upper motor neuron, 154, 154t, 155f
- Weaning, from mechanical ventilation, 66-67, 1743-1744, 1744f
- Weaponization, of microbes for bioterrorism, 261e-1
- Web(s), esophageal, 1902-1903
- Weber's syndrome, 208, 2575, 2575f
- Weber test, 221, 2538
- Wegener's syndrome. *See* Granulomatosis with polyangiitis (Wegener's)
- Weight
- over lifespan, 74f
 - regulation of, 275, 415e-2
 - set point of, 415e-5
- Weight gain. *See also* Obesity
- abdominal fat from, 285
 - in edema, 251
 - in hypothyroidism, 2291
 - physiology of, 415e-2
- Weight loss
- biliary sludge formation and, 2078
 - cholesterol stone formation and, 2078
 - for hypertension, 1622t
 - involuntary. *See* Involuntary weight loss
 - for metabolic syndrome, 2453
 - for obesity treatment. *See* Obesity, treatment of
- Weil's syndrome. *See* *Leptospira/leptospirosis* (Weil's syndrome)
- Weiss score, adrenocortical carcinoma, 2322
- Welander's distal myopathy, 462e-12, 462e-13t
- Wells criteria, for pulmonary embolism, 24t
- Werdnig-Hoffmann disease, 444e-9, 2636
- Wermer's syndrome. *See* Multiple endocrine neoplasia type 1 (MEN 1)
- Werner's syndrome, 94e-4, 436t
- Wernicke-Korsakoff syndrome, 96e-3, 182
- Wernicke's aphasia, 177-178, 177t
- Wernicke's area, 177, 180f
- Wernicke's disease/encephalopathy
- in alcohol use disorder, 2724
 - clinical features of, 41e-1v, 96e-3, 209, 1783, 2607
 - imaging in, 1783, 1783f
 - pathogenesis of, 1783
 - treatment of, 1783, 2607
- Westermark's sign, 101, 1633
- Western blot, 214e-8, 1245, 1246g
- Western equine encephalitis virus, 1306t, 1317

- West Nile virus infections
 climate change and, 151e-5
 clinical features of, 129t, 362, 1307t, 1316
 fever-myalgia, 1316
 meningoencephalitis, 892
 weakness, 156
 diagnosis of, 891, 895t
 epidemiology of, 151e-5, 894, 1316
 transfusion-transmitted, 138e-6, 1316
 in transplant recipient, 923, 1316
 treatment of, 897
- Westphal variant, Huntington's disease, 2621
- Wet beriberi, 96e-2-3, 463t, 2690
- Wet-bulb globe temperature, 479e-1
- Wet mount, 150e-1, 245e-1
- Wheal, 127, 339t
- Wheel bug bite, 2750
- Wheezing, 1661, 1676
- WHIM (warts, hypogammaglobulinemia, infections, and myelokathexis) syndrome, 415, 2111
- Whiplash injury, 121-122
- Whipple procedure, 555, 2101
- Whipple's disease, **1091**, **1944**
 clinical features of, 1945
 in acute infection, 1092
 cardiac, 1092, 1111-1112
 chronic meningitis, 907t
 "classic," 269, 1092, 1092f
 cutaneous, 360, 360t
 dermatologic, 1093
 gastrointestinal, 269, 1090, 1092f
 lymphatic, 1092-1093
 neurologic, 1092
 ocular, 1093
 pulmonary, 1092
 in reinfection/relapsing disease, 1093
 diagnosis of, 137, 1093, 1945
 epidemiology of, 1091
 etiology of, 1091, 1945
 global considerations in, 1091
 pathogenesis of, 1091-1092
 small-intestinal mucosal biopsies in, 1939, 1940t, 1941f
 treatment of, 1093-1094, 1945
- Whipple's triad, 2430
- Whipworm. *See Trichuris trichiura/trichuriasis* (whipworm)
- Whispered pectoriloquy, 1662
- White blood cells, 62e-12f, 81e-1-2, 413. *See also* Basophil(s); Eosinophil(s); Lymphocyte(s); Monocyte(s); Neutrophil(s)
- White coat hypertension, 1445, 1616
- Whitehead, 351
- White matter disease, diffuse (leukoaraiosis), 174f, 2568, 2602
- White pulp, of spleen, 409, 410f
- White sponge nevus, 240t
- Whitewater Arroyo virus, 1306t
- Whitlow, herpetic, 829
- WHO (World Health Organization), 8, 12, 12e-1, 13e-7
- Whole blood, 138e-2
- Whole-bowel irrigation, 473e-6
- Whole-exome sequencing, 448-449
- Whole-genome sequencing, 449, 769t
- Whooping cough. *See Pertussis*
- Wickham's striae, 349
- Widow spider bite, 2747-2748
- Wild type allele, 435
- Williams-Beuren syndrome, 1525
- Williams-Campbell syndrome, 1695
- Williams' myopathy, 462e-12, 462e-13t
- Williams' syndrome, 83e-6, 2477-2478
- Willow, 404t
- Wilms' tumor, 101e-6t
- Wilms' tumor gene-1 mutations. *See WT1 gene* mutations
- Wilson's disease, **2519**
 vs. autoimmune hepatitis, 2051
 clinical features of, 367e-2, 2519
 anemia, 96e-10
 hepatic, 283, 367e-2, 1992, 2062, 2519, 2625
- Kayser-Fleischer ring, 436e-8f, 2519, 2520f, 2625
 movement disorders, 2625
 neurologic, 2519
 psychiatric, 2519
 diagnosis of, 366e-6f, 367e-2, 1993t, 2062, 2520, 2520t
 epidemiology of, 2062, 2519
 genetic testing for, 450t
 global considerations in, 2521
 hepatocellular carcinoma and, 545, 545t
 pathogenesis of, 96e-10, 2062, 2519
 prevalence of, 2625
 prognostic index in, 2520, 2521t
 treatment of, 367e-2, 450t, 2519-2521, 2520t, 2625
 zinc deficiency in, 96e-9
- Wine maker's lung, 1681t
- "Winking owl" sign, 1791
- Winterbottom's sign, 1397
- WIPI gene mutations, 562
- Wiskott-Aldrich syndrome, **2110**
 clinical features of, 2110
 genetic considerations in, 2110
 IBD and, 1949t
 incidence of, 2110
 neutropenia in, 418
 thrombocytopenia in, 729, 2110
 treatment of, 2110
- Withdrawal syndrome
 alcohol. *See Alcohol withdrawal syndrome*
 benzodiazepine, 169, 2710-2711
 marijuana, 469e-3
 opioid, 468e-2, 468e-3
- Withdrawal/withholding of care
 from critically ill patient, 1735-1736
 enteral/parenteral nutrition, 98e-2
 legal aspects of, 6
 from terminally ill patient, 66-67
- WNT1 gene mutations, 101e-2t
- WNT4 gene mutations, 2354t, 2356t, 2357
- Wnt ligands, 89e-4
- Wnt proteins, 2143
- Wolbachia, 1159
- Wolf-Chaikoff effect, 2285, 2297, 2300
- Wolff-Parkinson-White syndrome
 accessory pathways in, 1481, 1482f
 developmental defects in, 265e-12
 ECG in, 273e-5, 278e-11f, 1482f
 treatment of, 1483
- Wolf-Hirschhorn syndrome, 83e-6
- Wolfram's (DIDMOAD) syndrome, 2276, 2348
- Wolman disease, 432e-4t, 432e-5
- Women's health, **6e-1**
 AD, 6e-1-2, 6e-2f
 atherosclerosis, 291e-7-8
 autoimmune disorders, 6e-4
 body temperature variations, 124
 breast cancer. *See Breast cancer*
 cancer, 6e-2f
 cardiovascular disease, 6e-2-3, 6e-3f, 1588
 diabetes mellitus, 6e-3-4
 disease risk, 6e-1
 domestic violence. *See Domestic abuse/violence*
 dysuria, 60e-1
 epilepsy, 2559
 hirsutism. *See Hirsutism*
 HIV infection, 6e-4
 hypertension, 6e-4
 leading causes of death, 6e-1, 6e-1t, 6e-2f
 life expectancy, 6e-1
 obesity, 6e-4
 osteoporosis, 6e-4
 perceived greatest health problems, by year, 6e-2f
 pharmacology, 6e-4-5
 principles of, 4
 psychological disorders, 6e-5
 sleep disorders, 6e-5
 substance abuse, 6e-5
 tobacco use, 6e-5
- Wood's lamp examination, 342-344, 359
- Woodworker's lung, 1681t
- "Work hardening" regimes, 120
- World Alliance for Patient Safety, 12e-1
- World Bank, 8
- World Federation of Neurosurgical Societies
 scale, for subarachnoid hemorrhage, 1785t
- World Health Organization (WHO), 8, 12, 12e-1, 13e-7
- World Mental Health: Problems and Priorities in Low-Income Countries*, 17
- World Trade Center disaster, 1692
- Wound(s)
 chronic. *See Cutaneous ulcers, chronic infections. See also Surgical-site infections*
 anaerobic bacterial, 1099
 botulism, 987, 988, 1784
 clostridial, 990
 in returning war veterans, 152e-5t, 152e-9
 specimen collection and transport, 150e-5t
- Woven bone, 2466
- Wrap-around services, 14
- Wright's stain, 245e-2, 342, 422
- Wrist drop, 463e-4
- Wrist fractures, 2489
- Writing, assessment of, 177
- WRN gene, 94e-4
- "Wrong-way" eyes, 1774
- WT1 gene mutations
 in AML, 680, 680t
 cancers associated with, 101e-3t, 101e-6t
 in disorders of sex development, 2349
 in mesothelioma, 508
 in testis development disorders, 2353, 2354t
 in WAGR syndrome, 431t
- Wuchereria bancrofti*, 245e-3t, 765t, 1417, 1418t.
See also Lymphatic filariasis
- Wyeomyia virus, 1309t
- X**
- X4 viruses, 1235
- Xanthelasma, 342f, 365
- Xanthine oxidase deficiency (hereditary xanthinuria), 431e-3t, 431e-6
- Xanthine oxidase inhibitors, 42, 42t, 431e-4-5
- Xanthoastrocytomas, pleomorphic, 601
- Xanthoma, 364t, 365
 in cardiovascular disease, 1442
 in cholestasis, 2443
 eruptive, 365, 436e-4f
 in familial chylomicronemia syndromes, 2439
 in familial dysbetalipoproteinemia, 2443
 in familial hypercholesterolemia, 436e-4f, 2441
 in lipoprotein disorders, 436e-4f, 2243
 normolipemic plane, 365
 palmar crease, 1442, 2443
 plane, 365
 tendinous, 436e-3f
 tendon, 365, 2441
 tuberoeruptive, 2443
 tuberous, 365
- Xanthomatosis, cerebrotendinous, 451e-4t
- X chromosome. *See also X-linked disorders*
 G-banded, 83e-1f
 ideogram, 83e-1f
 inactivation of, 101e-4, 430, 439
- XCR1, 372e-13t
- Xerocytosis, 653
- Xeroderma pigmentosum, 386, 434, 500
- Xerophthalmia, 96e-6, 1946t
- Xerosis, 341t, 1172
- Xerostomia
 dental disease and, 237
 differential diagnosis of, 2167t
 drug-induced, 241
 etiology of, 241
 radiation therapy-related, 505
 in Sjögren's syndrome, 2166
 in systemic sclerosis, 2162
 treatment of, 2168f, 2169
- Xerotic eczema (asteatotic eczema), 342f, 346.
See also Eczema
- Xingu virus, 1308t
- Xist gene, 430

- X-linked disorders
 adrenoleukodystrophy. *See*
 Adrenoleukodystrophy
 agammaglobulinemia, 377e-2
 chronic granulomatous disease, 375e-1, 2106
 color blindness, 197
 incidence of, 437
 inheritance of, 438–439, 438f
 mental retardation, 444e-4
 NEMO deficiency, 375e-1, 420t, 421, 2106
 spinobulbar muscular atrophy. *See*
 Spinobulbar muscular atrophy
 (Kennedy's disease)
 X-linked lymphoproliferative disease, 1188
 X-linked proliferative syndrome (XLP), 2112
 X-linked sideroblastic anemia (protoporphyrin),
 2522f, 2522t, 2531
 X-rays, 103e-4, 263e-1
 46,XX disorders of sex development, 2350t,
 2356t
 androgen excess, 2356t, 2357. *See also*
 Congenital adrenal hyperplasia
 (CAH); 21-Hydroxylase deficiency
 congenital absence of the vagina, 2357
 testicular /ovotesticular disorders of sex
 development, 2355, 2356t
 45,X/46,XY mosaicism (mixed gonadal
 dysgenesis), 2351t, 2352–2353
 46,XY disorders of sex development, 2350t,
 2354t
 androgen action disorders, 2353–2355
 androgen synthesis disorders, 2353
 cryptorchidism, 2355
 isolated hypospadias, 2355
 persistent müllerian duct syndrome, 2355
 testis development disorders, 2353
Xylohypha infections, 908t
 D-Xylose test, in malabsorption syndromes, 1938
- Y**
 Yamaguchi's syndrome, ECG in, 269e-10f
 Yatapoxvirus, 220e-1t
 Yaws, 207e-2, 207e-2f, 207e-2t, 207e-3
 Y chromosome, 426, 439
 Yeasts, 1329
 Yellow fever, 8, 1307t, 1322–1323
 Yellow fever vaccine
 adverse effects of, 795
 contraindications to, 798
 for transplant recipient, 929
 for travelers, 794–795, 795t
 Yellow nail syndrome, 1654
 Yellow oleander, 310
 Yellow sac spider, 2748
 Yergason's supination sign, 2248
Yersinia spp.
 adherence of, 145e-2t
 laboratory identification of, 1073–1074
 life cycle of, 1071
 phenotypes of, 1070
 tissue invasion by, 145e-7, 1071–1072, 1076
 Yersiniabactin, 1076
Yersinia enterocolitica, 1076, 1077. *See also*
Yersinia spp. infections (yersiniosis)
Yersinia spp. infections (yersiniosis)
 antithyroid antibodies in, 1077
 clinical features of, 266, 266t, 1077
 complications of, 267
 diagnosis of, 1077–1078
 enteric fever, 852t, 853
 epidemiology of, 1076
 etiology of, 1076
 global considerations in, 1076, 1077, 1078
 vs. IBD, 1956
 pathogenesis of, 1076
 pharyngitis, 231
 postinfective phenomena of, 1077
 prevention and control of, 1078
 treatment of, 1078
Yersinia pestis, 261e-3. *See also* Plague
 as bioterrorism agent, 261e-3–4, 1070, 1073
 laboratory identification of, 261e-4, 1073–
 1074, 1074f
 pathogenesis of, 1071–1072
Yersinia pseudotuberculosis, 1076, 1077. *See also*
Yersinia spp. infections (yersiniosis)
Yin Yan 1 gene, 444e-5
 Y-linked disorders, 439
 Yoga, 14e-1t, 338, 2240
 Yohimbine, 146, 2644
 Yolk sac tumor, of ovary, 594–595
 Yo protein, 372e-25t
⁹⁰Yttrium
 for adult T-cell leukemia/lymphoma, 225e-5
 for hepatocellular carcinoma, 551
 for NETs, 575
 Yttrium exposure, 263e-6t
- Z**
 Zafirlukast, 2119, 2124
 Zalcitabine (ddC), 1276f
 Zaleplon, 191, 473e-10t
 Zanamivir
 action of, 214e-5, 214e-9, 215e-1
 adverse effects of, 1213
 for influenza prophylaxis, 215e-2t, 1213t, 1214
 for influenza treatment, 215e-2t, 215e-5, 233,
 1213, 1213t
 pharmacology of, 215e-1
 resistance to, 215e-1, 215e-2t
 Z-disk, genetic defects of, 1555t
 Zeaxanthin, 96e-7
 Zenker's diverticulum, 256, 259, 1903, 1903f
 ZES. *See* Zollinger-Ellison syndrome (ZES)
 Zeta-associated protein 70 (ZAP70) deficiency,
 2109
 Zicam, 14e-3
 Zidovudine (AZT)
 action of, 1273
 for adult T-cell leukemia/lymphoma, 225e-5
 adverse effects of, 1273, 1274t
 anemia, 1262
 cutaneous, 379
 hepatotoxicity, 2023, 2030
 hypertrichosis, 380
 myopathy, 439, 462e-20, 462e-20t, 1267,
 2197, 2199, 2218t
 nail disorders, 380, 1263
 neutropenia, 417
 oral pigmentation, 240t
 drug interactions of, 246e-2t, 246e-4t, 798
 for HIV infection, 50, 1223, 1273, 1274t
 molecular structure of, 1276f
 resistance to, 1279f
 Zieve's syndrome, 2059
 Zika virus infection, 1307t, 1314
 Zileuton, 40
 Zimbabwe, 13e-6–7
- Zinc, 96e-9
 absorption of, 96e-9
 deficiency of. *See* Zinc deficiency
 functions of, 96e-9
 for macular degeneration, 205
 recommended intake of, 95e-3t
 reference range for, 2766t
 supplements. *See* Zinc supplements
 tolerable upper intake level of, 96e-9t
 toxicity of, 96e-9t, 96e-10
 for Wilson's disease, 2520–2521, 2520t, 2625
 Zinc deficiency
 clinical features of, 96e-9, 96e-9t, 461
 esophageal cancer risk and, 533
 global considerations in, 96e-9, 461
 treatment of, 461
 with vitamin A deficiency, 96e-6
 Zinc gluconate lozenges, 96e-9
 Zinc oxide exposure, 1692
 Zinc sulfate flotation, 152e-1
 Zinc supplements
 adverse effects of, 215
 for hepatic encephalopathy, 2066
 for olfactory dysfunction, 216
 parenteral, 98e-6t
 for zinc deficiency, 96e-9
 Ziprasidone, 2721, 2722t
 Zirconium, 367
 Ziv-aflibercept, 102e-5t, 103e-21
 Z-line deformity, 2136
 Z lines, of sarcomere, 265e-5
 Zn-DTPA, 263e-6t, 263e-7
 ZNRD1 gene, in HIV infection, 1240t
 Zoledronate (zoledronic acid)
 adverse effects of, 426e-4, 2499–2500
 for bone loss prevention in prostate cancer,
 586
 for hypercalcemia, 314, 610, 2480t, 2481
 for multiple myeloma, 717
 for osteoporosis management/prevention,
 2499–2500, 2500f, 2504
 for Paget's disease of bone, 426e-3–4,
 426e-3t
 Zollinger-Ellison syndrome (ZES), 568, 1927
 clinical features of, 559t, 1927–1928
 cobalamin malabsorption in, 646
 diagnosis of, 568, 1928, 1928t, 1929t, 1940t
 differential diagnosis of, 1924, 1927–1928
 epidemiology of, 1927
 gastroduodenal ulcer in, 262
 incidence of, 559t
 MEN 1 and, 562, 568, 1927–1928,
 2336–2337
 pathophysiology of, 559t, 1927
 treatment of, 568–569, 1929
 tumor distribution in, 559t, 1927
 tumor localization in, 1928–1929, 1929t
 Zolmitriptan, 2591, 2592t, 2593t, 2595
 Zolpidem, 65, 191, 473e-10t, 476e-4
 Zonisamide, 261, 473e-10t, 2555t, 2766t
 Zoster. *See* Herpes zoster (shingles)
 Zoster ophthalmicus. *See* Herpes zoster
 ophthalmicus
 Zoster sine herpetica, 1185
 Zotarolimus, 296e-2
 Z-scores, 2493, 2493f
 Zungarococha virus, 1309t
 Zwitterionic characteristics, 849
 Zygomycosis. *See* Mucormycosis